

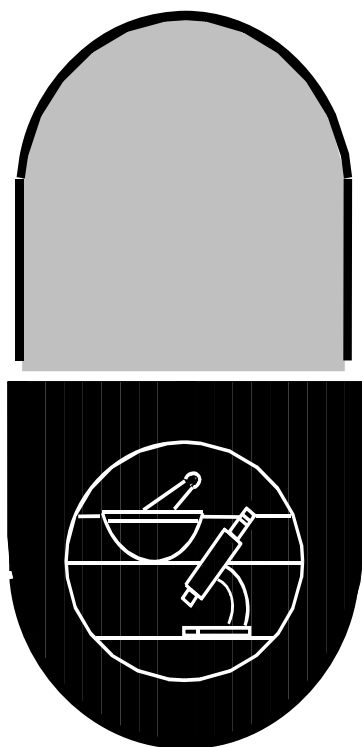
APPROVED DRUG PRODUCTS

With Therapeutic Equivalence Evaluations



The "Orange Book"

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APPROVED DRUG PRODUCTS

WITH

**THERAPEUTIC
EQUIVALENCE
EVALUATIONS**

27th EDITION

**THE PRODUCTS IN THIS LIST HAVE BEEN APPROVED UNDER
SECTION 505 OF THE FEDERAL FOOD, DRUG, AND COSMETIC ACT.**

**U.S. DEPARTMENT OF HEALTH AND HUMAN SERVICES
FOOD AND DRUG ADMINISTRATION
CENTER FOR DRUG EVALUATION AND RESEARCH
OFFICE OF PHARMACEUTICAL SCIENCE
OFFICE OF GENERIC DRUGS**

2007

APPROVED DRUG PRODUCTS with THERAPEUTIC EQUIVALENCE EVALUATIONS

The products in this list have been approved under section 505 of the Federal Food, Drug, and Cosmetic Act. This volume is current through December 31, 2006.

27th EDITION



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APPROVED DRUG PRODUCTS
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CONTENTS

	<i>PAGE</i>
PREFACE TO TWENTY SEVENTH EDITION.....	iv
1 INTRODUCTION.....	v
1.1 Content and Exclusion.....	v
1.2 Therapeutic Equivalence-Related Terms	v
1.3 Statistical Criteria for Bioequivalence	vii
1.4 Reference Listed Drug.....	ix
1.5 General Policies and Legal Status	ix
1.6 Practitioner/User Responsibilities.....	x
1.7 Therapeutic Equivalence Evaluations Codes	xii
1.8 Description of Special Situations.....	xix
1.9 Therapeutic Equivalence Code Change for a Drug Entity.....	xxi
1.10 Change of the Therapeutic Equivalence Evaluation for a Single Product	xxi
1.11 Discontinued Section.....	xxii
1.12 Changes to the Orange Book	xxii
1.13 Availability of the Edition.....	xxii
2 HOW TO USE THE DRUG PRODUCTS LISTS	2-1
2.1 Key Sections for Using the Drug Product Lists	2-1
2.2 Drug Product Illustration	2-3
2.3 Therapeutic Equivalence Evaluations Illustration	2-4
 DRUG PRODUCT LISTS	
Prescription Drug Product List	3-1
OTC Drug Product List	4-1
Drug Products with Approval under Section 505 of the Act Administered by the Center for Biologics Evaluation and Research List	5-1
Discontinued Drug Product List	6-1
Orphan Products Designations and Approvals List	7-1
Drug Products Which Must Demonstrate in vivo Bioavailability Only if Product Fails to Achieve Adequate Dissolution	8-1
 APPENDICES	
A. Product Name Index	A-1
B. Product Name Index Listed by Applicant	B-1
C. Uniform Terms	C-1
 PATENT AND EXCLUSIVITY INFORMATION ADDENDUM	
A. Patent and Exclusivity Lists	ADA1
B. Patent and Exclusivity Terms	ADB1

**FOOD AND DRUG ADMINISTRATION
CENTER FOR DRUG EVALUATION AND RESEARCH
APPROVED DRUG PRODUCTS
with
Therapeutic Equivalence Evaluations**

PREFACE TO TWENTY SEVENTH EDITION

The publication, *Approved Drug Products with Therapeutic Equivalence Evaluations* (the List, commonly known as the Orange Book), identifies drug products approved on the basis of safety and effectiveness by the Food and Drug Administration (FDA) under the Federal Food, Drug, and Cosmetic Act (the Act). Drugs on the market approved only on the basis of safety (covered by the ongoing Drug Efficacy Study Implementation [DESI] review [e.g., Donnatal® Tablets and Librax® Capsules] or pre-1938 drugs [e.g., Phenobarbital Tablets]) are not included in this publication. The main criterion for the inclusion of any product is that the product is the subject of an application with an effective approval that has not been withdrawn for safety or efficacy reasons. Inclusion of products on the List is independent of any current regulatory action through administrative or judicial means against a drug product. In addition, the List contains therapeutic equivalence evaluations for approved multisource prescription drug products. These evaluations have been prepared to serve as public information and advice to state health agencies, prescribers, and pharmacists to promote public education in the area of drug product selection and to foster containment of health care costs. Therapeutic equivalence evaluations in this publication are not official FDA actions affecting the legal status of products under the Act.

Background of the Publication. To contain drug costs, virtually every state has adopted laws and/or regulations that encourage the substitution of drug products. These state laws generally require either that substitution be limited to drugs on a specific list (the positive formulary approach) or that it be permitted for all drugs except those prohibited by a particular list (the negative formulary approach). Because of the number of requests in the late 1970s for FDA assistance in preparing both positive and negative formularies, it became apparent that FDA could not serve the needs of each state on an individual basis. The Agency also recognized that providing a single list based on common criteria would be preferable to evaluating drug products on the basis of differing definitions and criteria in various state laws. As a result, on May 31, 1978, the Commissioner of the Food and Drug Administration sent a letter to officials of each state stating FDA's intent to provide a list of all prescription drug products that are approved by FDA for safety and effectiveness, along with therapeutic equivalence determinations for multisource prescription products.

The List was distributed as a proposal in January 1979. It included only currently marketed prescription drug products approved by FDA through new drug applications (NDAs) and abbreviated new drug applications (ANDAs) under the provisions of Section 505 of the Act.

The therapeutic equivalence evaluations in the List reflect FDA's application of specific criteria to the multisource prescription drug products on the List approved under Section 505 of the Act. These evaluations are presented in the form of code letters that indicate the basis for the evaluation made. An explanation of the code appears in the *Introduction*.

A complete discussion of the background and basis of FDA's therapeutic equivalence evaluation policy was published in the *Federal Register* on January 12, 1979 (44 FR 2932). The final rule, which includes FDA's responses to the public comments on the proposal, was published in the *Federal Register* on October 31, 1980 (45 FR 72582). The first publication, October 1980, of

the final version of the List incorporated appropriate corrections and additions. Each subsequent edition has included the new approvals and made appropriate changes in data.

On September 24, 1984, the President signed into law the Drug Price Competition and Patent Term Restoration Act (1984 Amendments). The 1984 Amendments require that FDA, among other things, make publicly available a list of approved drug products with monthly supplements. The *Approved Drug Products with Therapeutic Equivalence Evaluations* publication and its monthly Cumulative Supplements satisfy this requirement. The *Addendum* to this publication identifies drugs that qualify under the 1984 Amendments for periods of exclusivity (during which ANDAs or applications described in Section 505(b)(2) of the Act for those drugs may not be submitted for a specified period of time and, if allowed to be submitted, would be tentatively approved) and provides patent information concerning the listed drugs which also may delay the approval of ANDAs or Section 505(b)(2) applications. The *Addendum* also provides additional information that may be helpful to those submitting a new drug application to the Agency.

The Agency intends to use this publication to further its objective of obtaining input and comment on the publication itself and related Agency procedures. Therefore, if you have comments on how the publication can be improved, please send them to the Director, Division of Labeling and Program Support, HFD-610, Office of Generic Drugs, Center for Drug and Evaluation and Research, 7500 Standish Place, Rockville, MD 20855. Comments received are publicly available to the extent allowable under the Freedom of Information regulations.

1. INTRODUCTION

1.1 Content and Exclusion

The List is composed of four parts: (1) approved prescription drug products with therapeutic equivalence evaluations; (2) approved over-the-counter (OTC) drug products for those drugs that may not be marketed without NDAs or ANDAs because they are not covered under existing OTC monographs; (3) drug products with approval under Section 505 of the Act administered by the Center for Biologics Evaluation and Research; and (4) a cumulative list of approved products that have never been marketed, are for exportation, are for military use, have been discontinued from marketing, or have had their approvals withdrawn for other than safety or efficacy reasons subsequent to being discontinued from marketing.¹ This publication also includes indices of prescription and OTC drug products by trade or established name (if no trade name exists) and by applicant name (holder of the approved application). All established names for active ingredients generally conform to official compendial names or *United States Adopted Names* (USAN) as prescribed in (21 CFR 299.4(e)). The latter list includes applicants' names as abbreviated in this publication; in addition, a list of uniform terms is provided. An *Addendum* contains drug patent and exclusivity information for the Prescription and OTC Drug Product Lists, and for the Drug Products with Approval under Section 505 of the Act Administered by the Center for Biologics Evaluation and Research. The publication may include additional information that the Agency deems appropriate to disseminate.

Prior to the 6th Edition, the publication had excluded OTC drug products and drug products with approval under Section 505 of the Act administered by the Center for Biologics Evaluation and Research because the main purpose of the publication was to provide information to states regarding FDA's recommendation as to which generic prescription drug products were acceptable candidates for drug product selection. The 1984 Amendments required the Agency to begin publishing an up-to-date list of all marketed drug products, OTC as well as prescription, that have been approved for safety and efficacy and for which new drug applications are required.

Under the 1984 Amendments, some drug products were given tentative approvals. Prior to the effective date, the Agency will not include drug products with tentative approval in the List; however, they are available at <http://www.fda.gov/cder/ogd/approvals/default.htm>. When the tentative approval becomes a full approval through a subsequent action letter to the application holder, the Agency will list the drug product and the final, effective approval date in the appropriate approved drug product list.

Distributors or repackagers of products on the List are not identified. Because distributors or repackagers are not required to notify FDA when they shift their sources of supply from one approved manufacturer to another, it is not possible to maintain complete information linking product approval with the distributor or repackager handling the products.

1.2 Therapeutic Equivalence-Related Terms

Pharmaceutical Equivalents. Drug products are considered pharmaceutical equivalents if they contain the same active ingredient(s), are of the same dosage form, route of administration and are identical in strength or concentration (e.g., chlorthalidone hydrochloride, 5mg capsules). Pharmaceutically equivalent drug products are formulated to contain the same amount of active ingredient in the same dosage form and to meet the same or

¹ Newly approved products are added to parts 1, 2, or 3, of the List, depending on the dispensing requirements (prescription or OTC) or approval authority, unless the Orange Book staff is otherwise notified before publication.

compendial or other applicable standards (i.e., strength, quality, purity, and identity), but they may differ in characteristics such as shape, scoring configuration, release mechanisms, packaging, excipients (including colors, flavors, preservatives), expiration time, and, within certain limits, labeling.

Pharmaceutical Alternatives. Drug products are considered pharmaceutical alternatives if they contain the same therapeutic moiety, but are different salts, esters, or complexes of that moiety, or are different dosage forms or strengths (e.g., tetracycline hydrochloride, 250mg capsules vs. tetracycline phosphate complex, 250mg capsules; quinidine sulfate, 200mg tablets vs. quinidine sulfate, 200mg capsules). Data are generally not available for FDA to make the determination of tablet to capsule bioequivalence. Different dosage forms and strengths within a product line by a single manufacturer are thus pharmaceutical alternatives, as are extended-release products when compared with immediate-release or standard-release formulations of the same active ingredient.

Therapeutic Equivalents. Drug products are considered to be therapeutic equivalents only if they are pharmaceutical equivalents and if they can be expected to have the same clinical effect and safety profile when administered to patients under the conditions specified in the labeling.

FDA classifies as therapeutically equivalent those products that meet the following general criteria: (1) they are approved as safe and effective; (2) they are pharmaceutical equivalents in that they (a) contain identical amounts of the same active drug ingredient in the same dosage form and route of administration, and (b) meet compendial or other applicable standards of strength, quality, purity, and identity; (3) they are bioequivalent in that (a) they do not present a known or potential bioequivalence problem, and they meet an acceptable *in vitro* standard, or (b) if they do present such a known or potential problem, they are shown to meet an appropriate bioequivalence standard; (4) they are adequately labeled; (5) they are manufactured in compliance with Current Good Manufacturing Practice regulations. *The concept of therapeutic equivalence, as used to develop the List, applies only to drug products containing the same active ingredient(s) and does not encompass a comparison of different therapeutic agents used for the same condition (e.g., propoxyphene hydrochloride vs. pentazocine hydrochloride for the treatment of pain).* Any drug product in the List repackaged and/or distributed by other than the application holder is considered to be therapeutically equivalent to the application holder's drug product even if the application holder's drug product is single source or coded as non-equivalent (e.g., **BN**). Also, distributors or repackagers of an application holder's drug product are considered to have the same code as the application holder. Therapeutic equivalence determinations are not made for unapproved, off-label indications.

FDA considers drug products to be therapeutically equivalent if they meet the criteria outlined above, even though they may differ in certain other characteristics such as shape, scoring configuration, release mechanisms, packaging, excipients (including colors, flavors, preservatives), expiration date/time and minor aspects of labeling (e.g., the presence of specific pharmacokinetic information) and storage conditions. When such differences are important in the care of a particular patient, it may be appropriate for the prescribing physician to require that a particular brand be dispensed as a medical necessity. With this limitation, however, FDA believes that products classified as therapeutically equivalent can be substituted with the full expectation that the substituted product will produce the same clinical effect and safety profile as the prescribed product.

Bioavailability. This term means the rate and extent to which the active ingredient or active moiety is absorbed from a drug product and becomes available at the site of action. For drug products that are not intended to be absorbed into the bloodstream, bioavailability may be assessed by measurements intended to reflect the rate and extent to which the active ingredient or active moiety becomes available at the site of action.

Bioequivalent Drug Products. This term describes pharmaceutical equivalent or pharmaceutical alternative products that display comparable bioavailability when studied under similar experimental conditions. Section 505 (j)(7)(B) of the Act describes one set of conditions under which a test and reference listed drug (see Section 1.4) shall be considered bioequivalent:

the rate and extent of absorption of the test drug do not show a significant difference from the rate and extent of absorption of the reference drug when administered at the same molar dose of the therapeutic ingredient under similar experimental conditions in either a single dose or multiple doses; or

the extent of absorption of the test drug does not show a significant difference from the extent of absorption of the reference drug when administered at the same molar dose of the therapeutic ingredient under similar experimental conditions in either a single dose or multiple doses and the difference from the reference drug in the rate of absorption of the drug is intentional, is reflected in its proposed labeling, is not essential to the attainment of effective body drug concentrations on chronic use, and is considered medically insignificant for the drug.

Where these above methods are not applicable (e.g., for drug products that are not intended to be absorbed into the bloodstream), other *in vivo* or *in vitro* test methods to demonstrate bioequivalence may be appropriate.

Bioequivalence may sometimes be demonstrated using an *in vitro* bioequivalence standard, especially when such an *in vitro* test has been correlated with human *in vivo* bioavailability data. In other situations, bioequivalence may sometimes be demonstrated through comparative clinical trials or pharmacodynamic studies.

1.3 Statistical Criteria for Bioequivalence

Under the Drug Price Competition and Patent Term Restoration Act of 1984, manufacturers seeking approval to market a generic drug product must submit data demonstrating that the drug product is bioequivalent to the pioneer (innovator) drug product. A major premise underlying the 1984 law is that bioequivalent drug products are therapeutically equivalent, and therefore, interchangeable.

Bioavailability refers to the rate and extent to which the active ingredient or therapeutic ingredient is absorbed from a drug product and becomes available at the site of drug action (Federal Food, Drug and Cosmetic Act, section 505(j)(8)). Bioequivalence refers to equivalent release of the same drug substance from two or more drug products or formulations. This leads to an equivalent rate and extent of absorption from these formulations. Underlying the concept of bioequivalence is the thesis that, if a drug product contains a drug substance that is chemically identical and is delivered to the site of action at the same rate and extent as another drug product, then it is equivalent and can be substituted for that drug product. Methods used to define bioequivalence can be found in 21 CFR 320.24, and include (1) pharmacokinetic (PK) studies, (2) pharmacodynamic (PD) studies, (3) comparative clinical trials, and (4) in-vitro studies. The choice of study used is based on the site of action of the drug and the ability of the study design to compare drug delivered to that site by the two products.

The standard bioequivalence (PK) study is conducted using a two-treatment crossover study design in a limited number of volunteers, usually 24 to 36 adults. Alternately, a four-period, replicate design crossover study may also be used. Single doses of the test and reference drug products are administered and blood or plasma levels of the drug are measured over time. Pharmacokinetic parameters characterizing rate and extent of drug absorption are evaluated statistically. The PK parameters of interest are the resulting area under the plasma concentration-time curve (AUC), calculated to the last

measured concentration ($AUC_{(0-t)}$) and extrapolated to infinity ($AUC_{(0-inf)}$), for extent of absorption; and the maximum or peak drug concentrations (C_{max}), for rate of absorption. Crossover studies may not be practical in drugs with a long half-life in the body, and a parallel study design may be used instead. Alternate study methods, such as in-vitro studies or equivalence studies with clinical or pharmacodynamic endpoints, are used for drug products where plasma concentrations are not useful to determine delivery of the drug substance to the site of activity (such as inhalers, nasal sprays and topical products applied to the skin).

The statistical methodology for analyzing these bioequivalence studies is called the two one-sided test procedure. Two situations are tested with this statistical methodology. The first of the two one-sided tests determines whether a generic product (test), when substituted for a brand-name product (reference) is significantly less bioavailable. The second of the two one-sided tests determines whether a brand-name product when substituted for a generic product is significantly less bioavailable. Based on the opinions of FDA medical experts, a difference of greater than 20% for each of the above tests was determined to be significant, and therefore, undesirable for all drug products. Numerically, this is expressed as a limit of test-product average/reference-product average of 80% for the first statistical test and a limit of reference-product average/test-product average of 80% for the second statistical test. By convention, all data is expressed as a ratio of the average response (AUC and C_{max}) for test/reference, so the limit expressed in the second statistical test is 125% (reciprocal of 80%).

For statistical reasons, all data is log-transformed prior to conducting statistical testing. In practice, these statistical tests are carried out using an analysis of variance procedure (ANOVA) and calculating a 90% confidence interval for each pharmacokinetic parameter (C_{max} and AUC). The confidence interval for both pharmacokinetic parameters, AUC and C_{max} , must be entirely within the 80% to 125% boundaries cited above. Because the mean of the study data lies in the center of the 90% confidence interval, the mean of the data is usually close to 100% (a test/reference ratio of 1). Different statistical criteria are sometimes used when bioequivalence is demonstrated through comparative clinical trials pharmacodynamic studies, or comparative in-vitro methodology.

The bioequivalence methodology and criteria described above simultaneously control for both, differences in the average response between test and reference, as well as the precision with which the average response in the population is estimated. This precision depends on the within-subject (normal volunteer or patient) variability in the pharmacokinetic parameters (AUC and C_{max}) of the two products and on the number of subjects in the study. The width of the 90% confidence interval is a reflection in part of the within-subject variability of the test and reference products in the bioequivalence study. A test product with no differences in the average response when compared to the reference might still fail to pass the bioequivalence criteria if the variability of one or both products is high and the bioequivalence study has insufficient statistical power (i.e., insufficient number of subjects). Likewise, a test product with low variability may pass the bioequivalence criteria, when there are somewhat larger differences in the average response.

This system of assessing bioequivalence of generic products assures that these substitutable products do not deviate substantially in in-vivo performance from the reference product. The Office of Generic Drugs has conducted two surveys to quantify the differences between generic and brand name products. The first survey included 224 bioequivalence studies submitted in approved applications during 1985 and 1986. The observed average differences between reference and generic products for AUC was 3.5% (JAMA, Sept. 4, 1987, Vol. 258, No. 9). The second survey included 127 bioequivalence studies submitted to the agency in 273 ANDAs approved in 1997.

The three measures reviewed include $AUC_{(0-t)}$, $AUC_{(0-inf)}$, and C_{max} . The observed average differences between the reference and generic products were $+ 3.47\%$ (SD 2.84) for $AUC_{(0-t)}$, $\pm 3.25\%$ (SD 2.97) for $AUC_{(0-inf)}$, and $\pm 4.29\%$ (SD 3.72)

The primary concern from the regulatory point of view is the protection of the patient against approval of products that are not bioequivalent. The current practice of carrying out two one-sided tests at the 0.05 level of significance ensures that there is no more than a 5% chance that a generic product that is not truly equivalent to the reference will be approved.

1.4 Reference Listed Drug

A reference listed drug (21 CFR 314.94(a)(3)) means the listed drug identified by FDA as the drug product upon which an applicant relies in seeking approval of its ANDA.

FDA has identified in the Prescription Drug Product and OTC Drug Product Lists those reference listed drugs to which the *in vivo* bioequivalence (reference standard) and, in some instances, the *in vitro* bioequivalence of the applicant's product is compared. By designating a single reference listed drug as the standard to which all generic versions must be shown to be bioequivalent, FDA hopes to avoid possible significant variations among generic drugs and their brand name counterpart. Such variations could result if generic drugs were compared to different reference listed drugs. However, in some instances when listed drugs are approved for a single drug product, a product not designated as the reference listed drug and not shown to be bioequivalent to the reference listed drug may be shielded from generic competition. A firm wishing to market a generic version of a listed drug that is not designated as the reference listed drug may petition the Agency through the Citizen Petition procedure (see 21 CFR 10.25(a) and CFR 10.30). When the Citizen Petition is approved, the second listed drug will be designated as an additional reference listed drug and the petitioner may submit an Abbreviated New Drug Application citing the designated reference listed drug. Section 1.7, *Therapeutic Equivalence Evaluations Codes products meeting necessary bioequivalence requirements* explains the (AB, AB1, AB2, AB3... coding system for multisource drug products listed under the same heading with two reference listed drugs.

In addition, there are two situations in which two listed drugs that have been shown to be bioequivalent to each other may both be designated as reference listed drugs. The first situation occurs when the *in vivo* determination of bioequivalence is self-evident and a waiver of the *in vivo* bioequivalence may be granted. The second situation occurs when the bioequivalence of two listed products may be determined through *in vitro* methodology. The reference listed drug is identified by the symbol "+" in the Prescription and Over-the-Counter (OTC) Drug Product Lists. These identified reference listed drugs represent the best judgment of the Division of Bioequivalence at this time. The Prescription and OTC Drug Product Lists identify reference drugs for oral dosage forms, injectables, ophthalmics, otics, and topical products. It is recommended that a firm planning to conduct an *in vivo* bioequivalence study, or planning to manufacture a batch of a drug product for which an *in vivo* waiver of bioequivalence will be requested, contact the Division of Bioequivalence, Office of Generic Drugs, to confirm the appropriate reference listed drug.

1.5 General Policies and Legal Status

The List contains public information and advice. It does not mandate the drug products which may be purchased, prescribed, dispensed, or substituted for one another, nor does it, conversely, mandate the products that should be avoided. To the extent that the List sets forth FDA's evaluations of the therapeutic equivalence of drug products that have been approved, it contains FDA's advice to the public, to practitioners and to the states regarding drug product selection. These evaluations do not constitute determinations that any product is in violation of the Act or that any product is preferable to any other. Therapeutic equivalence evaluations are a scientific judgment

based upon evidence, while generic substitution may involve social and economic policy administered by the states, intended to reduce the cost of drugs to consumers. To the extent that the List identifies drug products approved under Section 505 of the Act, it sets forth information that the Agency is required to publish and that the public is entitled to under the Freedom of Information Act. Exclusion of a drug product from the List does not necessarily mean that the drug product is either in violation of Section 505 of the Act, or that such a product is not safe or effective, or that such a product is not therapeutically equivalent to other drug products. Rather, the exclusion is based on the fact that FDA has not evaluated the safety, effectiveness, and quality of the drug product.

1.6 Practitioner/User Responsibilities

Professional care and judgment should be exercised in using the List. Evaluations of therapeutic equivalence for prescription drugs are based on scientific and medical evaluations by FDA. Products evaluated as therapeutically equivalent can be expected, in the judgment of FDA, to have equivalent clinical effect and no difference in their potential for adverse effects when used under the conditions of their labeling. However, these products may differ in other characteristics such as shape, scoring configuration, release mechanisms, packaging, excipients (including colors, flavors, preservatives), expiration date/time, and, in some instances, labeling. If products with such differences are substituted for each other, there is a potential for patient confusion due to differences in color or shape of tablets, inability to provide a given dose using a partial tablet if the proper scoring configuration is not available, or decreased patient acceptance of certain products because of flavor. There may also be better stability of one product over another under adverse storage conditions, or allergic reactions in rare cases due to a coloring or a preservative ingredient, as well as differences in cost to the patient.

FDA evaluation of therapeutic equivalence in no way relieves practitioners of their professional responsibilities in prescribing and dispensing such products with due care and with appropriate information to individual patients. In those circumstances where the characteristics of a specific product, other than its active ingredient, are important in the therapy of a particular patient, the physician's specification of that product is appropriate. Pharmacists must also be familiar with the expiration dates/times and labeling directions for storage of the different products, particularly for reconstituted products, to assure that patients are properly advised when one product is substituted for another.

Multisource and single-source drug products. FDA has evaluated for therapeutic equivalence only multisource prescription drug products approved under Section 505 of the Act, which in most instances means those pharmaceutical equivalents available from more than one manufacturer. For such products, a therapeutic equivalence code is included and, in addition, product information is highlighted in bold face and underlined. Those products with approved applications that are single-source (i.e., there is only one approved product available for that active ingredient, dosage form, route of administration, and strength) are also included on the List, but no therapeutic equivalence code is included with such products. Any drug product in the List repackaged and/or distributed by other than the application holder is considered to be therapeutically equivalent to the application holder's drug product even if the application holder's drug product is single source or coded as non-equivalent (e.g., **BN**). Also, although not identified in the List, distributors or repackagers of an application holder's drug product are considered to have the same code as the application holder. The details of these codes and the policies underlying them are discussed in Section 1.7, *Therapeutic Equivalence Evaluations Codes*.

Products on the List are identified by the names of the holders of approved applications (applicants) who may not necessarily be the manufacturer of the product. The applicant may have had its product manufactured by a

contract manufacturer and may simply be distributing the product for which it has obtained approval. In most instances, however, the manufacturer of the product is also the applicant. The name of the manufacturer is permitted by regulation to appear on the label, even when the manufacturer is not the marketer.

Although the products on the List are identified by the names of the applicants, circumstances, such as changing corporate ownership, have sometimes made identification of the applicant difficult. The Agency believes, based on continuing document review and communication with firms, that the applicant designations on the List are, in most cases, correct.

To relate firm name information on a product label to that on the List, the following should be noted: the applicant's name always appears on the List. This applies whether the applicant (firm name on the Form FDA 356h in the application) is the marketer (firm name in largest letters on the label) or not. However, the applicant's name may not always appear on the label of the product.

If the applicant is the marketer, its name appears on the List and on the label; if the applicant is not the marketer, and the Agency is aware of a corporate relationship (e.g., parent and subsidiary) between the applicant and the marketer, the name of the applicant appears on the List and both firm names may appear on the label. Firms with known corporate relationships are displayed in Appendix B. If there is no known corporate relationship between the applicant and the marketer, the applicant's name appears on the List; however, unless the applicant is the manufacturer, packager, or distributor, the applicant's name may not appear on the label. In this case, the practitioner, from labeling alone, will not be able to relate the marketed product to an applicant cited in the List, and hence to a specific approved drug product. In such cases, to assure that the product in question is the subject of an approved application, the firm named on the label should be contacted.

To relate trade name (proprietary name) information on a product label to that on the List, the following should be noted: if the applicant is the marketer, its name appears on the List and on the label; if the Agency is aware of a corporate relationship between the applicant and the marketer, the trade name (proprietary name) of the drug product (established drug name if no trade name exists) appears on the List. If a corporate relationship exists between an application holder and a marketer and both firms are distributing the drug product, the FDA reserves the right to select the trade name of either the marketer or the application holder to appear on the List. If there is no known corporate relationship between the applicant and the marketer, the established drug name appears on the List.

Every product on the List is subject at all times to regulatory action.

From time to time, approved products may be found in violation of one or more provisions of the Act. In such circumstances, the Agency will commence appropriate enforcement action to correct the violation, if necessary, by securing removal of the product from the market by voluntary recall, seizure, or other enforcement actions. Such regulatory actions are, however, independent of the inclusion of a product on the List. The main criterion for inclusion of a product is that it has an application with an effective approval that has not been withdrawn for safety or efficacy reasons. FDA believes that retention of a violative product on the List will not have any significant adverse health consequences, because other legal mechanisms are available to the Agency to prevent the product's actual marketing. FDA may however, change a product's therapeutic equivalence rating if the circumstances giving rise to the violation change or otherwise call into question the data upon which the Agency's assessment of whether a product meets the criteria for therapeutic equivalence was made.

1.7 Therapeutic Equivalence Evaluations Codes

The coding system for therapeutic equivalence evaluations is constructed to allow users to determine quickly whether the Agency has evaluated a particular approved product as therapeutically equivalent to other pharmaceutically equivalent products (first letter) and to provide additional information on the basis of FDA's evaluations (second letter). With few exceptions, the therapeutic equivalence evaluation date is the same as the approval date.

The two basic categories into which multisource drugs have been placed are indicated by the first letter as follows:

A Drug products that FDA considers to be therapeutically equivalent to other pharmaceutically equivalent products, i.e., drug products for which:

- (1) there are no known or suspected bioequivalence problems. These are designated **AA, AN, AO, AP, or AT**, depending on the dosage form; or
- (2) actual or potential bioequivalence problems have been resolved with adequate *in vivo* and/or *in vitro* evidence supporting bioequivalence. These are designated **AB**.

B Drug products that FDA at this time, considers not to be therapeutically equivalent to other pharmaceutically equivalent products, i.e.,

drug products for which actual or potential bioequivalence problems have not been resolved by adequate evidence of bioequivalence. Often the problem is with specific dosage forms rather than with the active ingredients. These are designated **BC, BD, BE, BN, BP, BR, BS, BT, BX, or B***.

Individual drug products have been evaluated as therapeutically equivalent to the reference product in accordance with the definitions and policies outlined below:

"A" CODES

Drug products that are considered to be therapeutically equivalent to other pharmaceutically equivalent products.

"A" products are those for which actual or potential bioequivalence problems have been resolved with adequate *in vivo* and/or *in vitro* evidence supporting bio-equivalence. Drug products designated with an "A" code fall under one of two main policies:

- (1) for those active ingredients or dosage forms for which no *in vivo* bioequivalence issue is known or suspected, the information necessary to show bioequivalence between pharmaceutically equivalent products is presumed and considered self-evident based on other data in the application for some dosage forms (e.g., solutions) or satisfied for solid oral dosage forms by a showing that an acceptable *in vitro* dissolution standard is met. A therapeutically equivalent rating is assigned such products so long as they are manufactured in accordance with Current Good Manufacturing Practice regulations and meet the other requirements of their approved applications (these are designated **AA, AN, AO, AP, or AT**, depending on the dosage form, as described below); or
- (2) for those DESI drug products containing active ingredients or dosage forms that have been identified by FDA as having actual or potential bioequivalence problems, and for post-1962 drug products in a dosage form presenting a potential bioequivalence problem, an evaluation of

therapeutic equivalence is assigned to pharmaceutical equivalents only if the approved application contains adequate scientific evidence establishing through *in vivo* and/or *in vitro* studies the bioequivalence of the product to a selected reference product (these products are designated as **AB**).

There are some general principles that may affect the substitution of pharmaceutically equivalent products in specific cases. Prescribers and dispensers of drugs should be alert to these principles so as to deal appropriately with situations that require professional judgment and discretion.

There may be labeling differences among pharmaceutically equivalent products that require attention on the part of the health professional. For example, pharmaceutically equivalent powders to be reconstituted for administration as oral or injectable liquids may vary with respect to their expiration time or storage conditions after reconstitution. An FDA evaluation that such products are therapeutically equivalent is applicable only when each product is reconstituted, stored, and used under the conditions specified in the labeling of that product.

The Agency will use notes in this publication to point out special situations such as potential differences between two drug products that have been evaluated as bioequivalent and otherwise therapeutically equivalent, when they should be brought to the attention of health professionals. These notes are contained in Section 1.8, *Description of Special Situations*.

For example, in rare instances, there may be variations among therapeutically equivalent products in their use or in conditions of administration. Such differences may be due to patent or exclusivity rights associated with such use. When such variations may, in the Agency's opinion, affect prescribing or substitution decisions by health professionals, a note will be added to Section 1.8.

Also, occasionally a situation may arise in which changes in a listed drug product after its approval (for example, a change in dosing interval) may have an impact on the substitutability of already approved generic versions of that product that were rated by the Agency as therapeutically equivalent to the listed product. When such changes in the listed drug product are considered by the Agency to have a significant impact on therapeutic equivalence, the Agency will change the therapeutic equivalence ratings for other versions of the drug product unless the manufacturers of those other versions of the product provide additional information to assure equivalence under the changed conditions. Pending receipt of the additional data, the Agency may add a note to Section 1.8, or, in rare cases, may even change the therapeutic equivalence rating.

In some cases (e.g., Isolyte® S w/ Dextrose 5% in Plastic Container and Plasma-Lyte® 148 and Dextrose 5% in Plastic Container), closely related products are listed as containing the same active ingredients, but in somewhat different amounts. In determining which of these products are pharmaceutically equivalent, the Agency has considered products to be pharmaceutically equivalent with labeled strengths of an ingredient that do not vary by more than 1%.

Different salts and esters of the same therapeutic moiety are regarded as pharmaceutical alternatives. For the purpose of this publication, such products are not considered to be therapeutically equivalent. There are no instances in this List where pharmaceutical alternatives are evaluated or coded with regard to therapeutic equivalence. Anhydrous and hydrated entities, as well as different polymorphs, are considered pharmaceutical equivalents and must meet the same standards and, where necessary, as in the case of ampicillin/ampicillin trihydrate, their equivalence is supported by appropriate bioavailability/bioequivalence studies.

The codes in this book are not intended to preclude health care professionals from converting pharmaceutically different concentrations into pharmaceutical equivalents using accepted professional practice.

Where package size variations have therapeutic implications, products so packaged have not been considered pharmaceutically equivalent. For example, some oral contraceptives are supplied in 21-tablet and 28-tablet packets; the 28-tablet packets contain 7 placebo or iron tablets. These two packaging configurations are not regarded as pharmaceutically equivalent; thus, they are not designated as therapeutically equivalent.

Preservatives may differ among some therapeutically equivalent drug products. Differences in preservatives and other inactive ingredients do not affect FDA's evaluation of therapeutic equivalence except in cases where these components may influence bioequivalence or routes of administration.

The specific sub-codes for those drugs evaluated as therapeutically equivalent and the policies underlying these sub-codes follow:

AA Products in conventional dosage forms not presenting bioequivalence problems

Products coded as **AA** contain active ingredients and dosage forms that are not regarded as presenting either actual or potential bioequivalence problems or drug quality or standards issues. However, all oral dosage forms must, nonetheless, meet an appropriate *in vitro* bioequivalence standard that is acceptable to the Agency in order to be approved.

AB, AB1, AB2, AB3... Products meeting necessary bioequivalence requirements

Multisource drug products listed under the same heading (i.e., identical active ingredient(s), dosage form, and route(s) of administration) and having the same strength (see Section 1.2, *Therapeutic Equivalence-Related Terms, Pharmaceutical Equivalents*) generally will be coded **AB** if a study is submitted demonstrating bioequivalence.

In certain instances, a number is added to the end of the **AB** code to make a three character code (i.e., **AB1**, **AB2**, **AB3**, etc.). Three-character codes are assigned only in situations when more than one reference listed drug of the same strength has been designated under the same heading. Two or more reference listed drugs are generally selected only when there are at least two potential reference drug products which are not bioequivalent to each other. If a study is submitted that demonstrates bioequivalence to a specific listed drug product, the generic product will be given the same three-character code as the reference listed drug it was compared against.

For example, Adalat® CC (Miles) and Procardia XL® (Pfizer), extended-release tablets, are listed under the active ingredient nifedipine. These drug products, listed under the same heading, are not bioequivalent to each other. Generic drug products deemed by FDA to be bioequivalent to Adalat® CC and Procardia XL® have been approved, Adalat® CC and Procardia XL® have been assigned ratings of **AB1** and **AB2**, respectively. The generic drug products bioequivalent to Adalat® CC would be assigned a rating of **AB1** and those bioequivalent to Procardia XL® would be assigned a rating of **AB2**. (The assignment of an **AB1** or **AB2** rating to a specific product does not imply product preference.) Even though drug products of distributors and/or repackagers are not included in the List, they are considered therapeutically equivalent to the application holder's drug product if the application holder's drug product is rated either with an **AB** or three-character code or is single source in the List. Drugs coded as **AB** under a heading are considered therapeutically equivalent only to other drugs coded as **AB** under that heading. Drugs coded with a three-character code under a heading are considered therapeutically equivalent only to other drugs coded with the same three-character code under that heading.

AN Solutions and powders for aerosolization

Uncertainty regarding the therapeutic equivalence of aerosolized products arises primarily because of differences in the drug delivery system. Solutions and powders intended for aerosolization that are marketed for use in any of several delivery systems are considered to be pharmaceutically and therapeutically equivalent and are coded **AN**. Those products that are compatible only with a specific delivery system or those products that are packaged in and with a specific delivery system are coded **BN**, unless they have met an appropriate bioequivalence standard. Solutions or suspensions in a specific delivery system will be coded **AN** if the bioequivalence standard is based upon *in vitro* methodology, if bioequivalence needs to be demonstrated by *in vivo* methodology then the drug products will be coded **AB**.

AO Injectable oil solutions

The absorption of drugs in injectable (parenteral) oil solutions may vary substantially with the type of oil employed as a vehicle and the concentration of the active ingredient. Injectable oil solutions are therefore considered to be pharmaceutically and therapeutically equivalent only when the active ingredient, its concentration, and the type of oil used as a vehicle are all identical.

AP Injectable aqueous solutions and, in certain instances, intravenous non-aqueous solutions

It should be noted that even though injectable (parenteral) products under a specific listing may be evaluated as therapeutically equivalent, there may be important differences among the products in the general category, *Injectable; Injection*. For example, some injectable products that are rated therapeutically equivalent are labeled for different routes of administration. In addition, some products evaluated as therapeutically equivalent may have different preservatives or no preservatives at all. Injectable products available as dry powders for reconstitution, concentrated sterile solutions for dilution, or sterile solutions ready for injection are all considered to be pharmaceutically and therapeutically equivalent provided they are designed to produce the same concentration prior to injection and are similarly labeled. Consistent with accepted professional practice, it is the responsibility of the prescriber, dispenser, or individual administering the product to be familiar with a product's labeling to assure that it is given only by the route(s) of administration stated in the labeling.

Certain commonly used large volume intravenous products in glass containers are not included on the List (e.g., dextrose injection 5%, dextrose injection 10%, sodium chloride injection 0.9%) since these products are on the market without FDA approval and the FDA has not published conditions for marketing such parenteral products under approved NDAs. When packaged in plastic containers, however, FDA regulations require approved applications prior to marketing. Approval then depends on, among other things, the extent of the available safety data involving the specific plastic component of the product. All large volume parenteral products are manufactured under similar standards, regardless of whether they are packaged in glass or plastic. Thus, FDA has no reason to believe that the packaging container of large volume parenteral drug products that are pharmaceutically equivalent would have any effect on their therapeutic equivalence.

The strength of parenteral drugs products is defined as the total drug content of the container. Until recently the strength of liquid parenteral drug products in the Orange Book have not been displayed. The concentration of the liquid parenteral drug product in the Orange Book has

been shown as xmg/ml. The amount of dry powder or freeze dried powder in a container has always been identified as the strength.

With the finalization of the Waxman-Hatch amendments that characterized each strength of a drug product as a listed drug, it became evident that the format of the Orange Book should be changed to reflect each strength of a parenteral solution. To this end the OGD has started to display the strength of all new approvals of parenteral solutions. Previously we would have displayed only the concentration of an approved parenteral solution, e.g. 50mg/ml. If this drug product had a 20 ml and 60 ml container approved the two products would be shown as 1Gm / 20ml (50mg/ml) and 3Gm / 60ml (50mg/ml).

AT Topical products

There are a variety of topical dosage forms available for dermatologic, ophthalmic, otic, rectal, and vaginal administration, including creams, gels, lotions, oils, ointments, pastes, solutions, sprays and suppositories. Even though different topical dosage forms may contain the same active ingredient and potency, these dosage forms are not considered pharmaceutically equivalent. Therefore, they are not considered therapeutically equivalent. All solutions and DESI drug products containing the same active ingredient in the same topical dosage form for which a waiver of *in vivo* bioequivalence has been granted and for which chemistry and manufacturing processes are adequate to demonstrate bioequivalence, are considered therapeutically equivalent and coded **AT**. Pharmaceutically equivalent topical products that raise questions of bioequivalence, including all post-1962 non-solution topical drug products, are coded **AB** when supported by adequate bioequivalence data, and **BT** in the absence of such data.

"B" CODES

Drug products that FDA, at this time, considers not to be therapeutically equivalent to other pharmaceutically equivalent products.

"B" products, for which actual or potential bioequivalence problems have not been resolved by adequate evidence of bioequivalence, often have a problem with specific dosage forms rather than with the active ingredients. Drug products designated with a "B" code fall under one of three main policies:

- (1) the drug products contain active ingredients or are manufactured in dosage forms that have been identified by the Agency as having documented bio-equivalence problems or a significant potential for such problems and for which no adequate studies demonstrating bioequivalence have been submitted to FDA; or
- (2) the quality standards are inadequate or FDA has an insufficient basis to determine therapeutic equivalence; or
- (3) the drug products are under regulatory review.

The specific coding definitions and policies for the "B" sub-codes are as follows:

B* Drug products requiring further FDA investigation and review to determine therapeutic equivalence

The code **B*** is assigned to products previously assigned an **A** or **B** code when FDA receives new information that raises a significant question regarding therapeutic equivalence that can be resolved only through further Agency investigation and/or review of data and information

FDA data supplied by DrugPatentWatch.com
submitted by the applicant. The **B*** code signifies that the Agency will take no position regarding the therapeutic equivalence of the product until the Agency completes its investigation and review.

BC Extended-release dosage forms (capsules, injectables and tablets)

Extended-release tablets are formulated in such a manner as to make the contained medicament available over an extended period of time following ingestion.

Although bioavailability studies have been conducted on these dosage forms, they may be subject to bioavailability differences, primarily because firms developing extended-release products for the same active ingredient rarely employ the same formulation approach. FDA, therefore, does not consider different extended-release dosage forms containing the same active ingredient in equal strength to be therapeutically equivalent unless equivalence between individual products in both rate and extent has been specifically demonstrated through appropriate bioequivalence studies. Extended-release products for which such bioequivalence data have not been submitted are coded **BC**, while those for which such data are available have been coded **AB**.

BD Active ingredients and dosage forms with documented bioequivalence problems

The **BD** code denotes products containing active ingredients with known bioequivalence problems and for which adequate studies have not been submitted to FDA demonstrating bioequivalence. Where studies showing bioequivalence have been submitted, the product has been coded **AB**.

BE Delayed-release oral dosage forms

Where the drug may be destroyed or inactivated by the gastric juice or where it may irritate the gastric mucosa, the use of "enteric" coatings is indicated. Such coatings are intended to delay the release of the medication until the tablet has passed through the stomach. Drug products in delayed-release dosage forms containing the same active ingredients are subject to significant differences in absorption. Unless otherwise specifically noted, the Agency considers different delayed-release products containing the same active ingredients as presenting a potential bioequivalence problem and codes these products **BE** in the absence of *in vivo* studies showing bioequivalence. If adequate *in vivo* studies have demonstrated the bioequivalence of specific delayed-release products, such products are coded **AB**.

BN Products in aerosol-nebulizer drug delivery systems

This code applies to drug solutions or powders that are marketed only as a component of, or as compatible with, a specific drug delivery system. There may, for example, be significant differences in the dose of drug and particle size delivered by different products of this type. Therefore, the Agency does not consider different metered aerosol dosage forms containing the same active ingredient(s) in equal strengths to be therapeutically equivalent unless the drug products meet an appropriate bioequivalence standard, such products are coded **AB**.

BP Active ingredients and dosage forms with potential bioequivalence problems

FDA's bioequivalence regulations (21 CFR 320.33) contain criteria and procedures for determining whether a specific active ingredient in a

specific dosage form has a potential for causing a bioequivalence problem.

It is FDA's policy to consider an ingredient meeting these criteria as having a potential bioequivalence problem even in the absence of positive data demonstrating inequivalence. Pharmaceutically equivalent products containing these ingredients in oral dosage forms are coded **BP** until adequate *in vivo* bioequivalence data are submitted, such products are coded **AB**. Injectable suspensions containing an active ingredient suspended in an aqueous or oleaginous vehicle have also been coded **BP**. Injectable suspensions are subject to bioequivalence problems because differences in particle size, polymorphic structure of the suspended active ingredient, or the suspension formulation can significantly affect the rate of release and absorption. FDA does not consider pharmaceutical equivalents of these products bioequivalent without adequate evidence of bioequivalence, such products would be coded **AB**.

BR Suppositories or enemas that deliver drugs for systemic absorption

The absorption of active ingredients from suppositories or enemas that are intended to have a systemic effect (as distinct from suppositories administered for local effect) can vary significantly from product to product. Therefore, FDA considers pharmaceutically equivalent systemic suppositories or enemas bio-equivalent only if *in vivo* evidence of bioequivalence is available. In those cases where *in vivo* evidence is available, the product is coded **AB**. If such evidence is not available, the products are coded **BR**.

BS Products having drug standard deficiencies

If the drug standards for an active ingredient in a particular dosage form are found by FDA to be deficient so as to prevent an FDA evaluation of either pharmaceutical or therapeutic equivalence, all drug products containing that active ingredient in that dosage form are coded **BS**. For example, if the standards permit a wide variation in pharmacologically active components of the active ingredient such that pharmaceutical equivalence is in question, all products containing that active ingredient in that dosage form are coded **BS**.

BT Topical products with bioequivalence issues

This code applies mainly to post-1962 dermatologic, ophthalmic, otic, rectal, and vaginal products for topical administration, including creams, ointments, gels, lotions, pastes, and sprays, as well as suppositories not intended for systemic drug absorption. Topical products evaluated as having acceptable clinical performance, but that are not bioequivalent to other pharmaceutically equivalent products or that lack sufficient evidence of bioequivalence, will be coded **BT**.

BX Drug products for which the data are insufficient to determine therapeutic equivalence

The code **BX** is assigned to specific drug products for which the data that have been reviewed by the Agency are insufficient to determine therapeutic equivalence under the policies stated in this document. In these situations, the drug products are presumed to be therapeutically inequivalent until the Agency has determined that there is adequate information to make a full evaluation of therapeutic equivalence.

1.8 Description of Special Situations

Certain drugs listed in the Orange Book present special situations that merit further discussion. Following is a description of those special situations:

Amino Acid and Protein Hydrolysate Injections. These products differ in the amount and kinds of amino acids they contain and, therefore, are not considered pharmaceutical equivalents. For this reason, these products are not considered therapeutically equivalent. At the same time, the Agency believes that it is appropriate to point out that where nitrogen balance is the sole therapeutic objective and individual amino acid content is not a consideration, pharmaceutical alternatives with the same total amount of nitrogen content may be considered therapeutically equivalent.

Follitropin Alfa and Beta. Based on available data derived from physico-chemical tests and bioassay, follitropin alfa and follitropin beta are indistinguishable.

Gaviscon®. Gaviscon® is an OTC product which has been marketed since September 1970. The active ingredients in this product, aluminum hydroxide and magnesium trisilicate, were reviewed by the Agency's OTC Antacid Panel and were considered to be safe and effective ingredients (Category I) by that Panel. However, the tablet failed to pass the antacid test which is required of all antacid products. The Agency, therefore, placed the tablet in Category III for lack of effectiveness. A full NDA with clinical studies was submitted by Marion Laboratories, Inc., and approved by FDA on December 9, 1983. Gaviscon®'s activity in treating reflux acidity is made possible by the physical-chemical properties of the inactive ingredients, sodium bicarbonate and alginic acid. Therefore, *all ANDAs which cite Gaviscon® tablets as the listed drug must contain the inactive ingredients sodium bicarbonate and alginic acid.* A full NDA will be required to support the effectiveness of the drug product if different inactive ingredients are to be substituted for sodium bicarbonate or alginic acid or if different proportions of these ingredients are to be used.

Levothyroxine Sodium. Because there are multiple reference listed drugs of levothyroxine sodium tablets and some reference listed drugs' sponsors have conducted studies to establish their drugs' therapeutic equivalence to other reference listed drugs, FDA has determined that its usual practice of assigning two or three character TE codes may be potentially confusing and inadequate for these drug products. Accordingly, FDA provides the following explanation and chart of therapeutic equivalence evaluations for levothyroxine sodium drug products.

Levothyroxine Sodium (Mylan ANDA 76187) tablets have been determined to be therapeutically equivalent to corresponding strengths of Unithroid (Jerome Stevens NDA 021210) tablets.

Levo-T (Alara NDA 021342), Levothyroxine Sodium (Mylan ANDA 76187), Unithroid (Jerome Stevens NDA 021210) and Levothyroxine Sodium (Genpharm ANDA 76752) tablets have been determined to be therapeutically equivalent to corresponding strengths of Synthroid (Abbott NDA 021402) tablets.

Levo-T (Alara NDA 021342), Unithroid (Jerome Stevens NDA 021210), Levothyroxine Sodium (Mylan ANDA 076187) and Levothyroxine Sodium (Genpharm ANDA 76752) tablets have been determined to be therapeutically equivalent to corresponding strengths of Levoxyl (King/Jones Pharma NDA 021301) tablets.

Levothyroxine Sodium (Mylan ANDA 76187) tablets have been determined to be therapeutically equivalent to corresponding strengths of Levotheroid (Lloyd NDA 021116) tablets.

Novothyrox (Genpharm NDA 021292) requires further investigation and review to establish therapeutic equivalence to corresponding strengths of any other

Levothyroxine Sodium drug products and is rated BX.

Levolet (Vintage NDA 021137) requires further investigation and review to establish therapeutic equivalence to corresponding strengths of any other Levothyroxine Sodium drug products and is rated BX.

The chart outlines TE codes for all 0.025mg products with other products being similar. Therapeutic equivalence has been established between products that have the same AB+number TE code. More than one TE code may apply to some products. One common TE code indicates therapeutic equivalence between products.

Trade Name	Applicant	Potency	TE Code	Appl No	Product No
UNITHROID	STEVENS J	0.025MG	AB1	21210	001
LEVOTHYROXINE SODIUM	MYLAN	0.025MG	AB1	76187	001
LEVOXYL	JONES PHARMA	0.025MG	AB1	21301	001
SYNTHROID	ABBOTT	0.025MG	AB1	21402	001
SYNTHROID	ABBOTT	0.025MG	AB2	21402	001
LEVOTHYROXINE SODIUM	MYLAN	0.025MG	AB2	76187	001
LEVO-T	ALARA PHARM	0.025MG	AB2	21342	001
UNITHROID	STEVENS J	0.025MG	AB2	21210	001
LEVOTHYROXINE SODIUM	GENPHARM	0.025MG	AB2	76752	001
LEVOXYL	JONES PHARMA	0.025MG	AB3	21301	001
LEVO-T	ALARA PHARM	0.025MG	AB3	21342	001
UNITHROID	STEVENS J	0.025MG	AB3	21210	001
LEVOTHYROXINE SODIUM	MYLAN	0.025MG	AB3	76187	001
LEVOTHYROXINE SODIUM	GENPHARM	0.025MG	AB3	76752	001
LEVOTHROID	LLOYD	0.025MG	AB4	21116	001
LEVOTHYROXINE SODIUM	MYLAN	0.025MG	AB4	76187	001
NOVOTHYROX	GENPHARM	0.025MG	BX	21292	001
LEVOLET	VINTAGE PHARMS	0.025MG	BX	21137	001

Patent Certification(s) Reference Listed Drug based upon a suitability petition. An abbreviated new drug application that refers to a Reference Listed Drug (RLD) approved pursuant to a suitability petition must demonstrate that the proposed product is bioequivalent to the RLD, and it must include appropriate patent certification(s) and an exclusivity statement with respect to the listed drug which served as the basis for the approved suitability petition.

Waived exclusivity. If a new drug application (NDA) submitted under section 505(b) of the Federal Food, Drug, and Cosmetic Act (Act) qualifies for exclusivity under sections 505(c)(3)(D) and 505(j)(5)(D), the exclusivity is listed in the Patent and Exclusivity Section of the Orange Book. If a drug product has received this exclusivity, the FDA will delay the approval of a 505(b)(2) application or an abbreviated new drug application (ANDAs) under section 505(j) of the Act until the expiration of the exclusivity. If the listed drug is also protected by one or more patents, the approval date for the 505(b)(2) application or ANDA will be determined by the latest expiring patent or exclusivity listed in the Orange Book. However, the holder of the NDA may waive its exclusivity as to any or all 505(b)(2) and ANDA applications referencing the protected drug product. If an NDA sponsor

waivers its right to the exclusivity protection, qualified 505(b)(2) or ANDA applications may be approved without regard to the NDA holder's exclusivity. An NDA for which the holder has waived its exclusivity as to all 505(b)(2) and ANDA applications will be coded with a W in the Patent and Exclusivity Section of the Orange Book and be referred to this section. The applicant referencing this listed drug should indicate in the exclusivity statement that the holder of the listed drug has waived its exclusivity.

Zocor (simvastatin) patent relisting. U.S. Patent Nos. RE 36481 and RE 36520 were relisted for Zocor (NDA 19-766) pursuant to the decision and related order in *Ranbaxy Labs. v. Leavitt*, No. 05-1838 (D.D.C. April 30, 2006). The '481 and '520 patents remained listed in *Approved Drug Products with Therapeutic Equivalence Evaluations* until any applicable periods of exclusivity pursuant to section 505(j)(5)(B)(iv) of the Federal Food, Drug, and Cosmetic Act were triggered and run. For additional information on this matter, please refer to Docket Nos. 2005P-0008 and 2005P-0046. Patents were subsequently delisted in the December 2006 Orange Book update as the exclusivity periods have triggered and run to expiration.

1.9 Therapeutic Equivalence Code Change for a Drug Entity

The Agency will use the following procedures when, in response to a petition or on its own initiative, it is considering a change in the therapeutic equivalence code for approved multi-source drug products. Such changes will generally occur when the Agency becomes aware of new scientific information affecting the therapeutic equivalence of an entire category of drug products in the List (e.g., information concerning the active ingredient or the dosage form), rather than information concerning a single drug product within the category. These procedures will be used when a change in therapeutic equivalence code is under consideration for all drug products found in the Prescription Drug Product List under a specific drug entity and dosage form. The change may be from the code signifying that the drug does not present a bioequivalence problem (e.g., **AA**) to a code signifying a bioequivalence problem (e.g., **BP**), or vice versa. This procedure does not apply to a change of a particular product code (e.g., a change from **BP** to **AB** or from **AB** to **BX**).

Before making a change in a therapeutic equivalence code for an entire category of drugs, the Agency will announce in the *Introduction* to the Cumulative Supplement that it is considering the change and will invite comment. Comments, along with scientific data, may be sent to the Director, Division of Bioequivalence, Office of Generic Drugs, Center for Drug Evaluation and Research, HFD-650, 7500 Standish Place, Rockville, MD 20855. The comment period will generally be 60 days in length, and the closing date for comments will be listed in the description of the proposed change for each drug entity.

The most useful type of scientific data submission is an *in vivo* bioavailability/bioequivalence study conducted on batches of the subject drug products. These submissions should present a full description of the analytical procedures and equipment used, a validation of the analytical methodology, including the standard curve, a description of the method of calculating results, and a description of the pharmacokinetic and statistical models used in analyzing the data. Anecdotal or testimonial information is the least useful to the Agency, and such submissions are discouraged. Copies of supporting reports published in the scientific literature or unpublished material, however, are welcome.

1.10 Change of the Therapeutic Equivalence Evaluation for a Single Product

The aforementioned procedure does not apply to a change in a single drug product code. For example, a change in a single drug product's code from **BP**

to **AB** as a result of the submission of an acceptable bioequivalence study ordinarily will not be the subject of notice and comment. Likewise, a change in a single drug product's code from **AB** to **BX** (e.g., as a result of new information raising a significant question as to bioequivalence) does not require notice and comment. The Agency's responsibility to provide the public with the Agency's most current information related to therapeutic equivalence may require a change in a drug product's code prior to any formal notice and opportunity for the applicant to be heard. The publication in the *Federal Register* of a proposal to withdraw approval of a drug product will ordinarily result in a change in a product's code from **AB** to **BX** if this action has not already been taken.

1.11 Discontinued Section

Those drug products in the Discontinued Section of the Orange Book in which a determination has already been made that the products were not withdrawn for safety or efficacy reasons have "***Federal Register determination that product was not discontinued or withdrawn for safety or efficacy reasons***" following the product strength. Those drug products are only reflective of citizen petitions approved since 1995. The identification of these drug products in the Discontinued Section of the Orange Book should avoid the submission of multiple citizen petitions for the same drug product

Generally, approved products are added to the Discontinued Section of the Orange Book when the applicant holder notifies the Orange Book staff of the products' not marketed status. Products may also be added if annual reports indicate the product is no longer marketed or other Agency administrative action (e.g., Withdrawal of an Application). Changes to the Orange Book are not affected by the drug registration and listing requirements of Section 510 of the Act.

1.12 Changes to the Orange Book

Every effort is made to ensure the Annual Edition is current and accurate. Applicant holders are requested to inform the FDA Orange Book Staff (OBS) of any changes or corrections. Please inform the OBS when products are no longer marketed. Notification of the Orange Book staff to include the newly approved product in the Discontinued Drug Product List rather than parts 1, 2 or 3 of the List (as discussed in Section 1.1) must occur by the end of the month in which the product is approved to ensure that the product is not included in the "active" portions of the next published Orange Book update

We can be contacted by email at drugproducts@cderr.fda.gov. Send Changes by FAX: 301-827-7337; mail to:

FDA/CDER Orange Book Staff
Office of Generic Drugs, HFD-610
7500 Standish Place
Rockville, MD 20855-2773

1.13 Availability of the Edition

Commencing with the 25th edition, the Annual Edition and current monthly Cumulative Supplements are available in a Portable Document Format (PDF) at the EOB home page, <http://www.fda.gov/cder/ob/default.htm>, by clicking on the [Annual Edition](#). The PDF annual format will duplicate previous paper versions except for the Orphan Products Designations and Approvals List. An annual subscription of the PDF format may be obtained from the U.S. Government Printing Office, 866-512-1800.

2. HOW TO USE THE DRUG PRODUCT LISTS

2.1 Key Sections for Using the Drug Product Lists

This publication contains illustrations, along with Drug Product Lists, indices, and lists of abbreviations and terms which facilitate their use.

Illustrations. The annotated *Drug Product Illustration*, see Section 2.2, and the *Therapeutic Equivalence Evaluations Illustration*, see Section 2.3, are offered to provide further clarification. These depict the format found in the Prescription Drug Product List (the only list in which therapeutic equivalence evaluation codes are displayed).

Drug Product Lists. The Prescription and OTC Drug Product Lists, arranged alphabetically by active ingredient(s), contain product identification information (active ingredients, dosage forms, routes of administration, product names, application holders, strengths) for single and multiple ingredient drug products. Also shown are the application number and drug product number (FDA internal computer data use only) and approval dates for those drug products approved on or after January 1, 1982.

The Discontinued Product List, arranged alphabetically by active ingredient(s), contain product identification information (dosage form, product name, strength, and application number).

If a prescription drug product is available from more than one source (multisource), a therapeutic equivalence code will appear in front of the applicant's name. If a product is therapeutically equivalent to one or more products or to an appropriate reference, it will be designated with a code beginning with "A" and the entry will be underlined and printed in bold font for emphasis.

Active ingredient headings for multiple ingredient (combination) drug products are arranged alphabetically. For purposes of this publication, this alphabetical sort takes precedence over United States Pharmacopeia official monograph order (i.e., Reserpine, Hydralazine Hydrochloride, Hydrochlorothiazide). For example, product information labeled as Reserpine, Hydrochlorothiazide and Hydralazine Hydrochloride appears under the active ingredient heading *Hydralazine Hydrochloride; Hydrochlorothiazide; Reserpine*. A cross-reference to the product information (for prescription and OTC products) appears for each additional active ingredient in the product. For combination drug products, the ingredient strengths are separated by semicolons and appear in the same relative sequence as the ingredients in the heading. Available strengths of the dosage form from an applicant appear on separate lines.

To use the Drug Product Lists, determine by alphabetical order the ingredient under which the product information is listed, using the Product Name Index, if necessary. Then, find the ingredient in the applicable Drug Product List. Proceed to the dosage form and route of administration and compare products within that ingredient heading only. Therapeutic equivalence or inequivalence for prescription products is determined on the basis of the therapeutic equivalence codes provided within that specific dosage form and route heading. The OTC Drug Product List, Discontinued Drug Product List, and Drug Products with Approval under Section 505 of the Act Administered by the Center for Biologics Evaluation and Research List have their data arranged similarly.

The Discontinued Drug Product List contains approved products that have never been marketed, have been discontinued from marketing, are for military use, or have had their approvals withdrawn for other than safety or efficacy reasons subsequent to being discontinued from marketing. All products having a "@" in the 12th Cumulative Supplement of the 26th Edition List have been added to the Discontinued Drug Product List appearing in the 27th Edition. In addition, approved drug products that are not in the commercial distribution channel e.g., approved drug products in applications for export only are also listed in the Discontinued Section of the Orange Book.

Product Name Index (*Prescription and OTC Drug Product Lists*). This is an index of drug products by established or trade name. The second term of each entry indicates the active ingredient name under which product information can be found in the appropriate Drug Product List. For those drug products with multiple active ingredients, only the first active ingredient (in alphabetical order) will appear. OTC products are so designated.

Product Name Index Listed by Applicant (*Prescription and OTC Drug Product Lists*). This is an index that cross-references applicants to drug products. The bolded and underlined entry represents the applicant name abbreviation used in this publication. Each complete applicant name that is represented by the abbreviated name is marked with an asterisk (*). Listed under each complete applicant name is the first alphabetically arranged ingredient under which product information can be found in the appropriate Drug Product List. OTC products are so designated. To use the Drug Product Lists, determine by alphabetical order the ingredient under which the product information is listed, using the Product Name Index, if appropriate.

Uniform Terms. To improve readability, uniform terms are used to designate dosage forms, routes of administration, and abbreviations used to express strengths. These terms are listed in Appendix C. In some cases, the terms used may differ from those used in product labels and other labeling.

2.2 DRUG PRODUCT ILLUSTRATION

SINGLE INGREDIENT

ACTIVE INGREDIENT	MEPERIDINE HYDROCHLORIDE			
DOSAGE FORM; ROUTE OF ADMINISTRATION	INJECTABLE; INJECTION			
TRADE OR GENERIC NAMES	HEXANON			
REFERENCE LISTED DRUG	AP + METRO-PHYS	25MG/ML	N13111 001	
	AP +	50MG/ML	N13111 002	
	AP +	75MG/ML	N13111 003	
	AP +	100MG/ML	N13111 004	AUG 22, 1983
	MEPERIDINE HCL			JAN 04, 1985
	DONHARE PHARM			
THERAPEUTIC EQUIVALENCE (TE)	AP	25MG/ML	N42242 001	
CODE FOR MULTISOURCE PRODUCT	AP	50MG/ML	N42296 001	
	AP	75MG/ML	N42301 002	
	AP	100MG/ML	N42301 003	AUG 27, 1987
FINAL APPROVAL DATE				AUG 27, 1987
SINGLE SOURCE PRODUCT (NO TE CODE)	AP	JOHNSON MED	10MG/ML	N40000 001
		HOOVAC LLC	25MG/ML	N47222 001
		OBS PHARM	150MG/ML	N47100 001
APPLICANT				
AVAILABLE STRENGTH(S) OF A PRODUCT				
APPLICATION NUMBER AND PRODUCT NUMBER				
PRODUCT NUMBER IS FOR FDA INTERNAL COMPUTER DATA USE ONLY				

MULTIPLE INGREDIENTS WITH PRODUCT INFORMATION

ALPHABETICALLY SORTED BY ACTIVE INGREDIENT	HYDRALAZINE HYDROCHLORIDE; HYDROCHLOROTHIAZIDE; RESERPINE
PRODUCT INFORMATION	TABLET; ORAL HYDROCHLOROTHIAZIDE, RESERPINE AND HYDRALAZINE HCL MADISON 25MG;15MG;0.1MG N41290 001 JAN 18, 1982

THIS EXAMPLE IS FOR PURPOSES OF ILLUSTRATION ONLY. IT DOES NOT REPRESENT ACTUAL PRODUCTS FROM THE PRESCRIPTION DRUG PRODUCT LIST.

2.3 THERAPEUTIC EQUIVALENCE EVALUATIONS ILLUSTRATION

DRUG PRODUCTS CODED **AB** (OR ANY CODE BEGINNING WITH AN "A") UNDER AN INGREDIENT AND DOSAGE FORM HEADING ARE CONSIDERED THERAPEUTICALLY EQUIVALENT ONLY TO OTHER PRODUCTS CODED **AB** (OR ANY CODE BEGINNING WITH AN "A") AND **NOT** TO THOSE CODED **BP** (OR ANY CODE BEGINNING WITH "B") AND ANY PRODUCTS NOT LISTED. DRUG PRODUCTS CODED **BP** (OR ANY CODE BEGINNING WITH A "B") ARE **NOT** CONSIDERED THERAPEUTICALLY EQUIVALENT TO ANY OTHER PRODUCT. FOR A COMPLETE EXPLANATION OF THE **TE** CODES REFER TO SECTION 1.7 OF THE *INTRODUCTION*.

		<u>SULFASALAZINE</u>			
		TABLET; ORAL			
		<u>FAZINE</u>			
PRODUCTS CONSIDERED THERAPEUTICALLY EQUIVALENT TO EACH OTHER	→	<u>AB</u>	PARKLAND	<u>500MG</u>	N42999 001
	→	<u>AB</u>	URSA	<u>500MG</u>	N40222 001
		<u>SULFASALAZINE</u>			
PRODUCTS CONSIDERED NOT THERAPEUTICALLY EQUIVALENT TO ANY OTHER PRODUCTS LISTED	→	BP	BROWN	500MG	N41297 001
		<u>SULFASALAZINE</u>			
		TABLET; ORAL			
		<u>FAZINE</u>			
PRODUCTS CONSIDERED NOT THERAPEUTICALLY EQUIVALENT TO EACH OTHER	→	<u>AB</u>	PARKLAND	<u>500MG</u>	N42999 001
	→		SULFASALAZINE		
	→	BP	BROWN	500MG	N41297 001
			SOUTH	500MG	N40627 001

NOTE: BOLD FONT AND UNDERLINING DENOTES MULTISOURCE PRODUCTS WHICH ARE CONSIDERED THERAPEUTICALLY EQUIVALENT.

THIS EXAMPLE IS FOR PURPOSES OF ILLUSTRATION ONLY. IT DOES NOT REPRESENT ACTUAL PRODUCTS FROM THE PRESCRIPTION DRUG PRODUCT LIST.

PRESCRIPTION DRUG PRODUCT LIST

3 - 1 (of 370)

ABACAVIR SULFATE

SOLUTION; ORAL					
ZIAGEN					
+ GLAXOSMITHKLINE	EQ 20MG BASE/ML		N20978	001	Dec 17, 1998
TABLET; ORAL					
ZIAGEN					
+ GLAXOSMITHKLINE	EQ 300MG BASE		N20977	001	Dec 17, 1998

ABACAVIR SULFATE; LAMIVUDINE

TABLET; ORAL					
EPZICOM					
+ SMITHKLINE BEECHAM	EQ 600MG BASE;300MG		N21652	001	Aug 02, 2004

ABACAVIR SULFATE; LAMIVUDINE; ZIDOVUDINE

TABLET; ORAL					
TRIZIVIR					
+ GLAXOSMITHKLINE	EQ 300MG BASE;150MG;300MG		N21205	001	Nov 14, 2000

ACAMPROSATE CALCIUM

TABLET, DELAYED RELEASE; ORAL					
CAMPRAL					
+ FOREST LABS	333MG		N21431	001	Jul 29, 2004

ACARBOSE

TABLET; ORAL					
PRECOSE					
BAYER PHARMS	25MG		N20482	004	May 29, 1997
	50MG		N20482	001	Sep 06, 1995
+	100MG		N20482	002	Sep 06, 1995

ACEBUTOLOL HYDROCHLORIDE

CAPSULE; ORAL					
<u>ACEBUTOLOL HYDROCHLORIDE</u>					
<u>AB</u>	ALPHAPHARM	<u>EQ 200MG BASE</u>	<u>N75047</u>	<u>001</u>	Dec 30, 1999
<u>AB</u>		<u>EQ 400MG BASE</u>	<u>N75047</u>	<u>002</u>	Dec 30, 1999
<u>AB</u>	MYLAN	<u>EQ 200MG BASE</u>	<u>N74288</u>	<u>001</u>	Apr 24, 1995
<u>AB</u>		<u>EQ 400MG BASE</u>	<u>N74288</u>	<u>002</u>	Apr 24, 1995
<u>AB</u>	WATSON LABS	<u>EQ 200MG BASE</u>	<u>N74007</u>	<u>001</u>	Oct 18, 1995
<u>AB</u>		<u>EQ 400MG BASE</u>	<u>N74007</u>	<u>002</u>	Oct 18, 1995
<u>SECTRAL</u>					
<u>AB</u>	DR REDDYS LABS INC	<u>EQ 200MG BASE</u>	<u>N18917</u>	<u>001</u>	Dec 28, 1984
<u>AB</u>	+	<u>EQ 400MG BASE</u>	<u>N18917</u>	<u>003</u>	Dec 28, 1984

ACETAMINOPHEN; ASPIRIN; CODEINE PHOSPHATE

CAPSULE; ORAL					
ACETAMINOPHEN, ASPIRIN, AND CODEINE PHOSPHATE					
+ MIKART	150MG;180MG;30MG		N81096	001	Oct 26, 1990

ACETAMINOPHEN; BUTALBITAL

CAPSULE; ORAL					
<u>BUCET</u>					
<u>AB</u>	MALLINCKRODT	<u>650MG;50MG</u>	<u>N88991</u>	<u>001</u>	Jun 28, 1985
<u>PHRENILIN FORTE</u>					
<u>AB</u>	+ VALEANT	<u>650MG;50MG</u>	<u>N88831</u>	<u>001</u>	Jun 19, 1985
<u>TENCON</u>					
<u>AB</u>	MALLINCKRODT	<u>650MG;50MG</u>	<u>N89405</u>	<u>001</u>	May 15, 1990
TABLET; ORAL					
<u>BUTAPAP</u>					
<u>AB</u>	MIKART	<u>325MG;50MG</u>	<u>N89987</u>	<u>001</u>	Oct 26, 1992

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 2 (of 370)

ACETAMINOPHEN; BUTALBITAL

TABLET; ORAL

BUTAPAP

<u>AB</u>	+	MIKART	<u>650MG;50MG</u>	<u>N89988</u>	<u>001</u>	Oct 26, 1992
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PHRENILIN

<u>AB</u>	+	VALEANT	<u>325MG;50MG</u>	<u>N87811</u>	<u>001</u>	Jun 19, 1985
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ACETAMINOPHEN; BUTALBITAL; CAFFEINE

CAPSULE; ORAL

ACETAMINOPHEN, BUTALBITAL, AND CAFFEINE

<u>AB</u>	+	MIKART	<u>325MG;50MG;40MG</u>	<u>N89007</u>	<u>001</u>	Mar 17, 1986
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BUTALBITAL, ACETAMINOPHEN AND CAFFEINE

<u>AB</u>		WEST WARD	<u>500MG;50MG;40MG</u>	<u>N40261</u>	<u>001</u>	Oct 28, 1998
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ESGIC-PLUS

<u>AB</u>	+	MIKART	<u>500MG;50MG;40MG</u>	<u>N40085</u>	<u>001</u>	Mar 28, 1996
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SOLUTION; ORAL

BUTALBITAL, ACETAMINOPHEN AND CAFFEINE

+	MIKART	325MG/15ML;50MG/15ML;40MG/15ML	N40387	001	Jan 31, 2003
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TABLET; ORAL

BUTALBITAL, ACETAMINOPHEN AND CAFFEINE

<u>AB</u>		MALLINCKRODT	<u>325MG;50MG;40MG</u>	<u>N87804</u>	<u>001</u>	Jan 24, 1985
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<u>AB</u>		MIKART	<u>325MG;50MG;40MG</u>	<u>N89175</u>	<u>001</u>	Jan 21, 1987
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+			750MG;50MG;40MG	N40496	001	Dec 23, 2003
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<u>AB</u>		MUTUAL PHARM	<u>325MG;50MG;40MG</u>	<u>N40601</u>	<u>001</u>	Jul 29, 2005
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<u>AB</u>		VINTAGE PHARMS	<u>325MG;50MG;40MG</u>	<u>N40511</u>	<u>001</u>	Aug 27, 2003
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<u>AB</u>			<u>500MG;50MG;40MG</u>	<u>N40513</u>	<u>001</u>	Aug 25, 2003
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<u>AB</u>		WATSON LABS	<u>500MG;50MG;40MG</u>	<u>N40267</u>	<u>001</u>	Jul 30, 1998
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<u>AB</u>		WEST WARD	<u>325MG;50MG;40MG</u>	<u>N89718</u>	<u>001</u>	Jun 12, 1995
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<u>AB</u>			<u>500MG;50MG;40MG</u>	<u>N40336</u>	<u>001</u>	Aug 18, 1999
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BUTALBITAL, APAP, AND CAFFEINE

<u>AB</u>		WATSON LABS	<u>325MG;50MG;40MG</u>	<u>N89536</u>	<u>001</u>	Feb 16, 1988
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ESGIC-PLUS

<u>AB</u>	+	MIKART	<u>500MG;50MG;40MG</u>	<u>N89451</u>	<u>001</u>	May 23, 1988
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FIORICET

<u>AB</u>	+	WATSON PHARMS	<u>325MG;50MG;40MG</u>	<u>N88616</u>	<u>001</u>	Nov 09, 1984
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ACETAMINOPHEN; BUTALBITAL; CAFFEINE; CODEINE PHOSPHATE

CAPSULE; ORAL

ACETAMINOPHEN, BUTALBITAL, CAFFEINE, AND CODEINE PHOSPHATE

<u>AB</u>		VINTAGE PHARMS	<u>325MG;50MG;40MG;30MG</u>	<u>N75929</u>	<u>001</u>	Apr 22, 2002
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BUTALBITAL, ACETAMINOPHEN, AND CAFFEINE WITH CODEINE PHOSPHATE

<u>AB</u>		WEST WARD	<u>325MG;50MG;40MG;30MG</u>	<u>N75618</u>	<u>001</u>	Mar 23, 2001
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BUTALBITAL, ACETAMINOPHEN, CAFFEINE AND CODEINE PHOSPHATE

<u>AB</u>		ANABOLIC LABS	<u>325MG;50MG;40MG;30MG</u>	<u>N76560</u>	<u>001</u>	Jun 10, 2004
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FIORICET W/ CODEINE

<u>AB</u>	+	WATSON PHARMS	<u>325MG;50MG;40MG;30MG</u>	<u>N20232</u>	<u>001</u>	Jul 30, 1992
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PHRENILIN WITH CAFFEINE AND CODEINE

<u>AB</u>		VALEANT	<u>325MG;50MG;40MG;30MG</u>	<u>N74911</u>	<u>001</u>	Aug 22, 2001
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ACETAMINOPHEN; CAFFEINE; DIHYDROCODEINE BITARTRATE

CAPSULE; ORAL

ACETAMINOPHEN, CAFFEINE, AND DIHYDROCODEINE BITARTRATE

+	MIKART	356.4MG;30MG;16MG	N40109	001	Aug 26, 1997
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TABLET; ORAL

ACETAMINOPHEN, CAFFEINE, AND DIHYDROCODEINE BITARTRATE

<u>AB</u>	+	MIKART	<u>712.8MG;60MG;32MG</u>	<u>N40316</u>	<u>001</u>	Apr 28, 1999
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ACETAMINOPHEN, CAFFEINE, AND DIHYDROCODEINE BITATRATE

<u>AB</u>		WEST WARD	<u>712.8MG;60MG;32MG</u>	<u>N40637</u>	<u>001</u>	Sep 22, 2006
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27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 3 (of 370)

ACETAMINOPHEN; CODEINE PHOSPHATE

SOLUTION; ORAL

ACETAMINOPHEN AND CODEINE PHOSPHATE

<u>AA</u>	+	ACTAVIS MID ATLANTIC	<u>120MG/5ML;12MG/5ML</u>	<u>N85861</u>	<u>001</u>	
<u>AA</u>		HI TECH PHARMA	<u>120MG/5ML;12MG/5ML</u>	<u>N40119</u>	<u>001</u>	Apr 26, 1996
<u>AA</u>		MIKART	<u>120MG/5ML;12MG/5ML</u>	<u>N89450</u>	<u>001</u>	Oct 27, 1992
<u>AA</u>		MORTON GROVE	<u>120MG/5ML;12MG/5ML</u>	<u>N87006</u>	<u>001</u>	
<u>AA</u>		PHARM ASSOC	<u>120MG/5ML;12MG/5ML</u>	<u>N87508</u>	<u>001</u>	

ACETAMINOPHEN W/ CODEINE

<u>AA</u>		ROXANE	<u>120MG/5ML;12MG/5ML</u>	<u>N86366</u>	<u>001</u>	
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SUSPENSION; ORAL

ACETAMINOPHEN AND CODEINE PHOSPHATE

<u>AA</u>	+	VALEANT	<u>120MG/5ML;12MG/5ML</u>	<u>N86024</u>	<u>001</u>	
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CAPITAL AND CODEINE

<u>AA</u>		ACTAVIS MID ATLANTIC	<u>120MG/5ML;12MG/5ML</u>	<u>N85883</u>	<u>001</u>	
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TABLET; ORAL

ACETAMINOPHEN AND CODEINE PHOSPHATE

<u>AA</u>		ANDRX PHARMS	<u>300MG;15MG</u>	<u>N40443</u>	<u>001</u>	Jan 22, 2003
<u>AA</u>			<u>300MG;30MG</u>	<u>N40443</u>	<u>002</u>	Jan 22, 2003
<u>AA</u>			<u>300MG;60MG</u>	<u>N40443</u>	<u>003</u>	Jan 22, 2003
<u>AA</u>	+	DURAMED PHARMS BARR	<u>300MG;15MG</u>	<u>N40223</u>	<u>001</u>	Nov 18, 1997
<u>AA</u>			<u>300MG;30MG</u>	<u>N40223</u>	<u>002</u>	Nov 18, 1997
<u>AA</u>			<u>300MG;60MG</u>	<u>N40223</u>	<u>003</u>	Nov 18, 1997
<u>AA</u>		MALLINCKRODT	<u>300MG;15MG</u>	<u>N40419</u>	<u>001</u>	May 31, 2001
<u>AA</u>			<u>300MG;30MG</u>	<u>N40419</u>	<u>002</u>	May 31, 2001
<u>AA</u>			<u>300MG;60MG</u>	<u>N40419</u>	<u>003</u>	May 31, 2001
<u>AA</u>		MIKART	<u>300MG;30MG</u>	<u>N89238</u>	<u>001</u>	Feb 25, 1986
	+		650MG;30MG	N89231	001	Mar 03, 1986
	+		650MG;60MG	N89363	001	Sep 09, 1991
<u>AA</u>		MUTUAL PHARM	<u>300MG;15MG</u>	<u>N89671</u>	<u>001</u>	Feb 10, 1988
<u>AA</u>			<u>300MG;30MG</u>	<u>N89672</u>	<u>001</u>	Feb 10, 1988
<u>AA</u>		PHARMERAL	<u>300MG;30MG</u>	<u>N87762</u>	<u>001</u>	Dec 10, 1982
<u>AA</u>		RANBAXY	<u>300MG;60MG</u>	<u>N87083</u>	<u>001</u>	
<u>AA</u>		SANDOZ	<u>300MG;30MG</u>	<u>N81250</u>	<u>001</u>	Jul 16, 1992
<u>AA</u>			<u>300MG;60MG</u>	<u>N81249</u>	<u>001</u>	Jul 16, 1992
<u>AA</u>		TEVA	<u>300MG;15MG</u>	<u>N88627</u>	<u>001</u>	Mar 06, 1985
<u>AA</u>			<u>300MG;30MG</u>	<u>N88628</u>	<u>001</u>	Mar 06, 1985
<u>AA</u>	+		<u>300MG;60MG</u>	<u>N88629</u>	<u>001</u>	Mar 06, 1985
<u>AA</u>		VINTAGE	<u>300MG;15MG</u>	<u>N89990</u>	<u>001</u>	Sep 30, 1988
<u>AA</u>		VINTAGE PHARMS	<u>300MG;30MG</u>	<u>N89805</u>	<u>001</u>	Sep 30, 1988
<u>AA</u>			<u>300MG;60MG</u>	<u>N89828</u>	<u>001</u>	Sep 30, 1988
<u>AA</u>		WATSON LABS	<u>300MG;15MG</u>	<u>N89997</u>	<u>001</u>	Dec 28, 1994
<u>AA</u>			<u>300MG;30MG</u>	<u>N89998</u>	<u>001</u>	Dec 28, 1994
<u>AA</u>			<u>300MG;60MG</u>	<u>N89999</u>	<u>001</u>	Dec 28, 1994

ACETAMINOPHEN W/ CODEINE PHOSPHATE #3

<u>AA</u>		RANBAXY	<u>300MG;30MG</u>	<u>N85868</u>	<u>001</u>	
		CODRIX				
	+	ANDRX PHARMS	500MG;15MG	N40447	001	Feb 26, 2003
	+		500MG;30MG	N40441	001	Mar 27, 2003
	+		500MG;60MG	N40488	001	Mar 28, 2003

TYLENOL W/ CODEINE NO. 3

<u>AA</u>	+	ORTHO MCNEIL PHARM	<u>300MG;30MG</u>	<u>N85055</u>	<u>003</u>	
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ACETAMINOPHEN; HYDROCODONE BITARTRATE

CAPSULE; ORAL

ALLAY

<u>AA</u>		IVAX PHARMS	<u>500MG;5MG</u>	<u>N89907</u>	<u>001</u>	Jan 13, 1989
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HYDROCET

<u>AA</u>		MALLINCKRODT	<u>500MG;5MG</u>	<u>N89006</u>	<u>001</u>	Aug 09, 1985
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PRESCRIPTION DRUG PRODUCT LIST

3 - 4 (of 370)

ACETAMINOPHEN; HYDROCODONE BITARTRATE

CAPSULE; ORAL

HYDROCODONE BITARTRATE AND ACETAMINOPHEN

<u>AA</u>	MALLINCKRODT	<u>500MG;5MG</u>	<u>N88956</u>	<u>001</u>	Jul 19, 1985
<u>AA</u>	MIKART	<u>500MG;5MG</u>	<u>N81067</u>	<u>001</u>	Nov 30, 1989
<u>AA</u>		<u>500MG;5MG</u>	<u>N81068</u>	<u>001</u>	Nov 30, 1989
<u>AA</u>		<u>500MG;5MG</u>	<u>N81069</u>	<u>001</u>	Nov 30, 1989
<u>AA</u>		<u>500MG;5MG</u>	<u>N81070</u>	<u>001</u>	Nov 30, 1989
<u>AA</u>		<u>500MG;5MG</u>	<u>N89008</u>	<u>001</u>	Feb 21, 1986

LORCET-HD

<u>AA</u>	+ MALLINCKRODT	<u>500MG;5MG</u>	<u>N87336</u>	<u>001</u>	Jul 08, 1982
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SOLUTION; ORAL

HYDROCODONE BITARTRATE AND ACETAMINOPHEN

<u>AA</u>	KV PHARM	<u>500MG/15ML;7.5MG/15ML</u>	<u>N40366</u>	<u>001</u>	Jan 23, 2002
<u>AA</u>	MALLINCKRODT	<u>500MG/15ML;7.5MG/15ML</u>	<u>N40418</u>	<u>001</u>	Jun 27, 2001
<u>AA</u>	+	<u>500MG/15ML;10MG/15ML</u>	<u>N40508</u>	<u>001</u>	Aug 29, 2003
<u>AA</u>	MIKART	<u>500MG/15ML;5MG/15ML</u>	<u>N81226</u>	<u>001</u>	Oct 27, 1992
<u>AA</u>	+	<u>500MG/15ML;5MG/15ML</u>	<u>N89557</u>	<u>001</u>	Apr 29, 1992
<u>AA</u>	+	<u>500MG/15ML;7.5MG/15ML</u>	<u>N81051</u>	<u>001</u>	Aug 28, 1992
<u>AA</u>	+	325MG/15ML;7.5MG/15ML	N40482	001	Sep 25, 2003
<u>AA</u>	PHARM ASSOC	<u>500MG/15ML;7.5MG/15ML</u>	<u>N40182</u>	<u>001</u>	Mar 13, 1998
<u>AA</u>	VINTAGE PHARMS	<u>500MG/15ML;7.5MG/15ML</u>	<u>N40520</u>	<u>001</u>	Oct 30, 2003

TABLET; ORAL

ANEXSIA

<u>AA</u>	MALLINCKRODT	<u>500MG;5MG</u>	<u>N89160</u>	<u>001</u>	Apr 23, 1987
<u>AA</u>		<u>750MG;10MG</u>	<u>N40468</u>	<u>001</u>	Oct 31, 2002

ANEXSIA 5/325

<u>AA</u>	MALLINCKRODT	<u>325MG;5MG</u>	<u>N40409</u>	<u>001</u>	Oct 20, 2000
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ANEXSIA 7.5/325

<u>AA</u>	MALLINCKRODT	<u>325MG;7.5MG</u>	<u>N40405</u>	<u>001</u>	Sep 08, 2000
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ANEXSIA 7.5/650

<u>AA</u>	MALLINCKRODT	<u>650MG;7.5MG</u>	<u>N89725</u>	<u>001</u>	Sep 30, 1987
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CO-GESIC

<u>AA</u>	SCHWARZ PHARMA	<u>500MG;5MG</u>	<u>N87757</u>	<u>001</u>	May 03, 1982
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HYDROCODONE BITARTRATE AND ACETAMINOPHEN

<u>AA</u>	ANDRX PHARMS	<u>500MG;5MG</u>	<u>N40493</u>	<u>001</u>	May 28, 2003
<u>AA</u>		<u>660MG;10MG</u>	<u>N40495</u>	<u>001</u>	May 28, 2003
<u>AA</u>		<u>750MG;7.5MG</u>	<u>N40494</u>	<u>001</u>	May 28, 2003
<u>AA</u>	BARR	<u>500MG;2.5MG</u>	<u>N40307</u>	<u>001</u>	Jul 26, 2000
<u>AA</u>		<u>500MG;5MG</u>	<u>N40308</u>	<u>001</u>	Jul 26, 2000
<u>AA</u>		<u>500MG;7.5MG</u>	<u>N40307</u>	<u>002</u>	Jul 26, 2000
<u>AA</u>		<u>500MG;10MG</u>	<u>N40309</u>	<u>001</u>	Jul 26, 2000
<u>AA</u>		<u>650MG;7.5MG</u>	<u>N40307</u>	<u>003</u>	Jul 26, 2000
<u>AA</u>		<u>650MG;10MG</u>	<u>N40307</u>	<u>004</u>	Jul 26, 2000
<u>AA</u>		<u>750MG;7.5MG</u>	<u>N40308</u>	<u>002</u>	Jul 26, 2000
<u>AA</u>	INTERPHARM	<u>325MG;5MG</u>	<u>N40736</u>	<u>001</u>	Aug 25, 2006
<u>AA</u>		<u>325MG;10MG</u>	<u>N40746</u>	<u>001</u>	Aug 25, 2006
<u>AA</u>		<u>500MG;5MG</u>	<u>N40729</u>	<u>001</u>	Aug 25, 2006
<u>AA</u>		<u>500MG;7.5MG</u>	<u>N40748</u>	<u>001</u>	Aug 25, 2006
<u>AA</u>		<u>650MG;7.5MG</u>	<u>N40754</u>	<u>001</u>	Aug 25, 2006
<u>AA</u>		<u>650MG;10MG</u>	<u>N40757</u>	<u>001</u>	Aug 25, 2006
<u>AA</u>		<u>750MG;7.5MG</u>	<u>N40769</u>	<u>001</u>	Aug 28, 2006
<u>AA</u>	IVAX PHARMS	<u>500MG;5MG</u>	<u>N89696</u>	<u>001</u>	Apr 21, 1988
<u>AA</u>	MALLINCKRODT	<u>325MG;10MG</u>	<u>N40400</u>	<u>001</u>	Jul 26, 2000
<u>AA</u>		<u>500MG;5MG</u>	<u>N40084</u>	<u>002</u>	Jun 01, 1995
<u>AA</u>		<u>500MG;7.5MG</u>	<u>N40201</u>	<u>001</u>	Feb 27, 1998
<u>AA</u>		<u>500MG;10MG</u>	<u>N40201</u>	<u>002</u>	Feb 27, 1998
<u>AA</u>		<u>650MG;10MG</u>	<u>N40084</u>	<u>004</u>	Oct 16, 1996
<u>AA</u>	+	<u>660MG;10MG</u>	<u>N40084</u>	<u>003</u>	Jul 29, 1996
<u>AA</u>		<u>750MG;7.5MG</u>	<u>N40084</u>	<u>001</u>	Jun 01, 1995

PRESCRIPTION DRUG PRODUCT LIST

3 - 5 (of 370)

ACETAMINOPHEN; HYDROCODONE BITARTRATE

TABLET; ORAL			
HYDROCODONE BITARTRATE AND ACETAMINOPHEN			
<u>AA</u>	MIKART	<u>325MG; 7.5MG</u>	<u>N40432 001</u> Jan 22, 2003
<u>AA</u>	+	<u>500MG; 2.5MG</u>	<u>N89698 001</u> Aug 25, 1989
<u>AA</u>		<u>500MG; 5MG</u>	<u>N89271 001</u> Jul 16, 1986
<u>AA</u>		<u>500MG; 5MG</u>	<u>N89697 001</u> Jan 28, 1992
<u>AA</u>	+	<u>500MG; 7.5MG</u>	<u>N89699 001</u> Aug 25, 1989
<u>AA</u>	+	<u>650MG; 7.5MG</u>	<u>N89689 001</u> Jun 29, 1988
<u>AA</u>	+	<u>650MG; 10MG</u>	<u>N81223 001</u> May 29, 1992
		300MG; 5MG	N40658 001 Jan 19, 2006
	+	300MG; 7.5MG	N40556 002 Mar 24, 2006
	+	300MG; 10MG	N40556 001 Jun 23, 2004
<u>AA</u>	MUTUAL PHARM	<u>500MG; 5MG</u>	<u>N40236 001</u> Sep 25, 1997
<u>AA</u>		<u>650MG; 7.5MG</u>	<u>N40240 002</u> Nov 26, 1997
<u>AA</u>		<u>650MG; 10MG</u>	<u>N40240 001</u> Nov 26, 1997
<u>AA</u>		<u>750MG; 7.5MG</u>	<u>N40236 002</u> Sep 25, 1997
<u>AA</u>	SANDOZ	<u>500MG; 5MG</u>	<u>N40149 001</u> Jan 27, 1997
<u>AA</u>		<u>750MG; 7.5MG</u>	<u>N40149 002</u> Jan 27, 1997
<u>AA</u>	UCB INC	<u>500MG; 10MG</u>	<u>N40210 001</u> Aug 13, 1997
<u>AA</u>		<u>650MG; 7.5MG</u>	<u>N40134 001</u> Nov 21, 1996
<u>AA</u>	VINTAGE PHARMS	<u>325MG; 5MG</u>	<u>N40655 001</u> Jan 19, 2006
<u>AA</u>		<u>325MG; 7.5MG</u>	<u>N40656 001</u> Jan 19, 2006
<u>AA</u>		<u>325MG; 10MG</u>	<u>N40355 001</u> May 31, 2000
<u>AA</u>		<u>500MG; 2.5MG</u>	<u>N40144 002</u> Apr 25, 1997
<u>AA</u>		<u>500MG; 5MG</u>	<u>N89831 001</u> Sep 07, 1988
<u>AA</u>		<u>500MG; 5MG</u>	<u>N89971 001</u> Dec 02, 1988
<u>AA</u>		<u>500MG; 7.5MG</u>	<u>N40144 001</u> Feb 22, 1996
<u>AA</u>		<u>500MG; 10MG</u>	<u>N40356 001</u> May 31, 2000
<u>AA</u>		<u>650MG; 7.5MG</u>	<u>N40155 001</u> Apr 14, 1997
<u>AA</u>		<u>650MG; 10MG</u>	<u>N40143 001</u> Feb 22, 1996
<u>AA</u>		<u>660MG; 10MG</u>	<u>N40358 001</u> May 31, 2000
<u>AA</u>		<u>750MG; 7.5MG</u>	<u>N40157 001</u> Apr 12, 1996
<u>AA</u>	WATSON LABS	<u>325MG; 10MG</u>	<u>N40248 002</u> Apr 28, 2000
<u>AA</u>		<u>500MG; 2.5MG</u>	<u>N40123 003</u> Mar 04, 1996
<u>AA</u>		<u>500MG; 2.5MG</u>	<u>N81079 001</u> Aug 30, 1991
<u>AA</u>		<u>500MG; 5MG</u>	<u>N40122 001</u> Mar 04, 1996
<u>AA</u>		<u>500MG; 5MG</u>	<u>N89883 001</u> Dec 01, 1988
<u>AA</u>		<u>500MG; 7.5MG</u>	<u>N40123 004</u> Mar 04, 1996
<u>AA</u>		<u>500MG; 7.5MG</u>	<u>N81080 001</u> Aug 30, 1991
<u>AA</u>		<u>500MG; 10MG</u>	<u>N40148 002</u> Feb 14, 1997
<u>AA</u>		<u>650MG; 7.5MG</u>	<u>N40094 001</u> Sep 29, 1995
<u>AA</u>		<u>650MG; 7.5MG</u>	<u>N40123 001</u> Mar 04, 1996
<u>AA</u>		<u>650MG; 10MG</u>	<u>N40094 002</u> Sep 29, 1995
<u>AA</u>		<u>650MG; 10MG</u>	<u>N40123 002</u> Mar 04, 1996
<u>AA</u>		<u>660MG; 10MG</u>	<u>N40094 003</u> Aug 08, 2000
<u>AA</u>		<u>750MG; 7.5MG</u>	<u>N40122 002</u> Mar 04, 1996
<u>AA</u>		<u>750MG; 7.5MG</u>	<u>N81083 001</u> Aug 30, 1991
<u>AA</u>	+	<u>750MG; 10MG</u>	<u>N40094 004</u> Mar 22, 1999
<u>LORTAB</u>			
<u>AA</u>	MALLINCKRODT	<u>500MG; 5MG</u>	<u>N87722 001</u> Jul 09, 1982
<u>AA</u>	+	<u>500MG; 10MG</u>	<u>N40100 001</u> Jan 26, 1996
<u>NORCO</u>			
<u>AA</u>	+	<u>325MG; 5MG</u>	<u>N40099 001</u> Jun 25, 1997
<u>AA</u>	+	<u>325MG; 7.5MG</u>	<u>N40148 003</u> Sep 12, 2000
<u>AA</u>	+	<u>325MG; 10MG</u>	<u>N40148 001</u> Feb 14, 1997
<u>VICODIN</u>			
<u>AA</u>	+	<u>500MG; 5MG</u>	<u>N88058 001</u> Jan 07, 1983
<u>VICODIN ES</u>			
<u>AA</u>	+	<u>750MG; 7.5MG</u>	<u>N89736 001</u> Dec 09, 1988

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 6 (of 370)

ACETAMINOPHEN; HYDROCODONE BITARTRATE

TABLET; ORAL

VICODIN HP

<u>AA</u>	ABBOTT	<u>660MG;10MG</u>	<u>N40117</u>	<u>001</u>	Sep 23, 1996
	ZYDONE				
	+ ENDO PHARMS	400MG;5MG	N40288	001	Nov 27, 1998
	+	400MG;7.5MG	N40288	002	Nov 27, 1998
	+	400MG;10MG	N40288	003	Nov 27, 1998

ACETAMINOPHEN; OXYCODONE HYDROCHLORIDE

CAPSULE; ORAL

OXYCODONE AND ACETAMINOPHEN

<u>AA</u>	ACTAVIS TOTOWA	<u>500MG;5MG</u>	<u>N40199</u>	<u>001</u>	Dec 30, 1998
<u>AA</u>	BARR	<u>500MG;5MG</u>	<u>N40304</u>	<u>001</u>	Oct 02, 2000
<u>AA</u>	DURAMED PHARMS BARR	<u>500MG;5MG</u>	<u>N40289</u>	<u>001</u>	Mar 16, 1999
<u>AA</u>	ENDO PHARMS	<u>500MG;5MG</u>	<u>N40303</u>	<u>001</u>	Dec 30, 1999
<u>AA</u>	MALLINCKRODT	<u>500MG;5MG</u>	<u>N40257</u>	<u>001</u>	Aug 04, 1998
<u>AA</u>	MUTUAL PHARM	<u>500MG;5MG</u>	<u>N40219</u>	<u>001</u>	Jan 22, 1998
<u>AA</u>	VINTAGE PHARMS	<u>500MG;5MG</u>	<u>N40106</u>	<u>001</u>	Jul 30, 1996
<u>AA</u>	WATSON LABS	<u>500MG;5MG</u>	<u>N40234</u>	<u>001</u>	Oct 30, 1997

ROXILOX

<u>AA</u>	ROXANE	<u>500MG;5MG</u>	<u>N40061</u>	<u>001</u>	Jul 03, 1995
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TYLOX

<u>AA</u>	+ ORTHO MCNEIL PHARM	<u>500MG;5MG</u>	<u>N88790</u>	<u>001</u>	Dec 12, 1984
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SOLUTION; ORAL

OXYCODONE AND ACETAMINOPHEN

<u>AA</u>	MALLINCKRODT	<u>325MG/5ML;5MG/5ML</u>	<u>N40680</u>	<u>001</u>	Sep 29, 2006
	<u>ROXICET</u>				
<u>AA</u>	+ ROXANE	<u>325MG/5ML;5MG/5ML</u>	<u>N89351</u>	<u>001</u>	Dec 03, 1986

TABLET; ORAL

OXYCET

<u>AA</u>	MALLINCKRODT	<u>325MG;5MG</u>	<u>N87463</u>	<u>001</u>	Dec 07, 1983
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OXYCODONE AND ACETAMINOPHEN

<u>AA</u>	ACTAVIS TOTOWA	<u>325MG;5MG</u>	<u>N40203</u>	<u>001</u>	Mar 15, 1999
<u>AA</u>	BARR	<u>325MG;5MG</u>	<u>N87406</u>	<u>001</u>	
<u>AA</u>	DURAMED PHARMS BARR	<u>325MG;5MG</u>	<u>N40272</u>	<u>001</u>	Jun 30, 1998
<u>AA</u>	MALLINCKRODT	<u>325MG;7.5MG</u>	<u>N40545</u>	<u>001</u>	Jun 30, 2004
<u>AA</u>		<u>325MG;10MG</u>	<u>N40545</u>	<u>002</u>	Jun 30, 2004
<u>AA</u>		<u>500MG;7.5MG</u>	<u>N40550</u>	<u>001</u>	Jun 30, 2004
<u>AA</u>		<u>650MG;10MG</u>	<u>N40550</u>	<u>002</u>	Jun 30, 2004
	+ MIKART	300MG;2.5MG	N40608	001	Dec 30, 2005
	+	300MG;5MG	N40608	002	Dec 30, 2005
	+	300MG;7.5MG	N40608	003	Dec 30, 2005
	+	300MG;10MG	N40608	004	Dec 30, 2005
	+	400MG;2.5MG	N40679	001	May 16, 2006
	+	400MG;5MG	N40687	001	Apr 27, 2006
	+	400MG;7.5MG	N40698	001	Apr 27, 2006
	+	400MG;10MG	N40692	001	Apr 27, 2006
		500MG;10MG	N40676	001	Apr 19, 2006
<u>AA</u>	VINTAGE PHARMS	<u>325MG;5MG</u>	<u>N40105</u>	<u>001</u>	Jul 30, 1996
<u>AA</u>	WATSON LABS	<u>325MG;5MG</u>	<u>N40171</u>	<u>001</u>	Oct 30, 1997
<u>AA</u>		<u>325MG;7.5MG</u>	<u>N40535</u>	<u>001</u>	Sep 05, 2003
<u>AA</u>		<u>325MG;10MG</u>	<u>N40535</u>	<u>002</u>	Sep 05, 2003
<u>AA</u>		<u>500MG;7.5MG</u>	<u>N40371</u>	<u>001</u>	Dec 29, 2000
<u>AA</u>		<u>650MG;10MG</u>	<u>N40371</u>	<u>002</u>	Dec 29, 2000
	<u>PERCOCET</u>				
<u>AA</u>	ENDO PHARMS	<u>325MG;5MG</u>	<u>N40330</u>	<u>002</u>	Jun 25, 1999
<u>AA</u>	+	<u>325MG;5MG</u>	<u>N85106</u>	<u>002</u>	
<u>AA</u>	+	<u>325MG;7.5MG</u>	<u>N40434</u>	<u>001</u>	Nov 23, 2001
<u>AA</u>	+	<u>325MG;10MG</u>	<u>N40434</u>	<u>002</u>	Nov 23, 2001

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 7 (of 370)

ACETAMINOPHEN; OXYCODONE HYDROCHLORIDE

TABLET; ORAL

PERCOCET

<u>AA</u>	+	ENDO PHARMS	<u>500MG;7.5MG</u>	<u>N40341</u>	<u>001</u>	Jul 26, 1999
<u>AA</u>	+		<u>650MG;10MG</u>	<u>N40341</u>	<u>002</u>	Jul 26, 1999
	+		325MG;2.5MG	N40330	001	Jun 25, 1999

ROXICET

<u>AA</u>		ROXANE	<u>325MG;5MG</u>	<u>N87003</u>	<u>001</u>	
		ROXICET 5/500				
	+	ROXANE	500MG;5MG	N89775	001	Jan 12, 1989

ACETAMINOPHEN; PENTAZOCINE HYDROCHLORIDE

TABLET; ORAL

ACETAMINOPHEN AND PENTAZOCINE HYDROCHLORIDE

<u>AB</u>		ACTAVIS TOTOWA	<u>650MG;EQ 25MG BASE</u>	<u>N76202</u>	<u>001</u>	Aug 02, 2002
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PENTAZOCINE HYDROCHLORIDE AND ACETAMINOPHEN

<u>AB</u>		WATSON LABS	<u>650MG;EQ 25MG BASE</u>	<u>N74699</u>	<u>001</u>	Mar 24, 2000
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TALACEN

<u>AB</u>	+	SANOFI AVENTIS US	<u>650MG;EQ 25MG BASE</u>	<u>N18458</u>	<u>001</u>	Sep 23, 1982
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ACETAMINOPHEN; PROPOXYPHENE HYDROCHLORIDE

TABLET; ORAL

PROPOXYPHENE HYDROCHLORIDE AND ACETAMINOPHEN

<u>AA</u>		MYLAN	<u>650MG;65MG</u>	<u>N83978</u>	<u>001</u>	
<u>AA</u>		SANDOZ	<u>650MG;65MG</u>	<u>N89959</u>	<u>001</u>	Jul 18, 1989
<u>AA</u>		VINTAGE PHARMS	<u>650MG;65MG</u>	<u>N40507</u>	<u>001</u>	Jul 30, 2003
<u>AA</u>		WATSON LABS	<u>650MG;65MG</u>	<u>N40139</u>	<u>001</u>	Dec 16, 1996

WYGESIC

<u>AA</u>	+	LEITNER PHARMS	<u>650MG;65MG</u>	<u>N84999</u>	<u>001</u>	
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ACETAMINOPHEN; PROPOXYPHENE NAPSYLATE

TABLET; ORAL

DARVOCET A500

<u>AB</u>		XANODYNE PHARM	<u>500MG;100MG</u>	<u>N76429</u>	<u>001</u>	Sep 10, 2003
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DARVOCET-N 100

<u>AB</u>	+	XANODYNE PHARM	<u>650MG;100MG</u>	<u>N17122</u>	<u>002</u>	
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DARVOCET-N 50

<u>AB</u>		XANODYNE PHARM	<u>325MG;50MG</u>	<u>N17122</u>	<u>001</u>	
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PROPOXYPHENE NAPSYLATE AND ACETAMINOPHEN

<u>AB</u>		ACTAVIS ELIZABETH	<u>650MG;100MG</u>	<u>N70910</u>	<u>001</u>	Jan 02, 1987
<u>AB</u>		ANDRX PHARMS	<u>500MG;100MG</u>	<u>N77196</u>	<u>001</u>	Jun 28, 2005
<u>AB</u>			<u>650MG;100MG</u>	<u>N76609</u>	<u>001</u>	Nov 16, 2004
<u>AB</u>		CORNERSTONE	<u>500MG;100MG</u>	<u>N76750</u>	<u>001</u>	Jun 28, 2004
			325MG;100MG	N76743	001	May 07, 2004
<u>AB</u>		IVAX PHARMS	<u>650MG;100MG</u>	<u>N70146</u>	<u>001</u>	Aug 02, 1985
<u>AB</u>		MALLINCKRODT	<u>650MG;100MG</u>	<u>N75738</u>	<u>001</u>	Feb 02, 2001
<u>AB</u>		MUTUAL PHARM	<u>650MG;100MG</u>	<u>N70615</u>	<u>001</u>	Mar 21, 1986
<u>AB</u>			<u>650MG;100MG</u>	<u>N70771</u>	<u>001</u>	Mar 21, 1986
<u>AB</u>			<u>650MG;100MG</u>	<u>N70775</u>	<u>001</u>	Mar 21, 1986
<u>AB</u>		MYLAN	<u>650MG;100MG</u>	<u>N70145</u>	<u>001</u>	Jun 12, 1985
<u>AB</u>			<u>650MG;100MG</u>	<u>N72195</u>	<u>001</u>	Feb 16, 1988
<u>AB</u>		SANDOZ	<u>650MG;100MG</u>	<u>N70443</u>	<u>001</u>	Jan 23, 1986
<u>AB</u>		TEVA	<u>650MG;100MG</u>	<u>N74119</u>	<u>001</u>	Dec 19, 1994
<u>AB</u>		VINTAGE PHARMS	<u>325MG;50MG</u>	<u>N74843</u>	<u>002</u>	Feb 15, 2001
<u>AB</u>			<u>650MG;100MG</u>	<u>N74843</u>	<u>001</u>	Feb 12, 1997

ACETAMINOPHEN; TRAMADOL HYDROCHLORIDE

TABLET; ORAL

TRAMADOL HYDROCHLORIDE AND ACETAMINOPHEN

<u>AB</u>		BARR	<u>325MG;37.5MG</u>	<u>N76914</u>	<u>001</u>	Jul 26, 2006
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27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 8 (of 370)

ACETAMINOPHEN; TRAMADOL HYDROCHLORIDE

TABLET; ORAL

TRAMADOL HYDROCHLORIDE AND ACETAMINOPHEN

<u>AB</u>	CARACO	<u>325MG; 37.5MG</u>	<u>N77184</u>	<u>001</u>	Dec 16, 2005
<u>AB</u>	KALI LABS	<u>325MG; 37.5MG</u>	<u>N76475</u>	<u>001</u>	Apr 21, 2005
<u>ULTRACET</u>					
<u>AB</u>	+ ORTHO MCNEIL PHARM	<u>325MG; 37.5MG</u>	<u>N21123</u>	<u>001</u>	Aug 15, 2001

ACETAZOLAMIDE

CAPSULE, EXTENDED RELEASE; ORAL

DIAMOX

	+ DURAMED PHARMS BARR	500MG	N12945	003	Apr 25, 1995
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TABLET; ORAL

ACETAZOLAMIDE

<u>AB</u>	LANNETT	<u>250MG</u>	<u>N84840</u>	<u>001</u>	
<u>AB</u>	MUTUAL PHARM	<u>125MG</u>	<u>N89752</u>	<u>001</u>	Jun 22, 1988
<u>AB</u>		<u>250MG</u>	<u>N89753</u>	<u>001</u>	Jun 22, 1988
<u>AB</u>	TARO	<u>125MG</u>	<u>N40195</u>	<u>001</u>	May 28, 1997
<u>AB</u>	+	<u>250MG</u>	<u>N40195</u>	<u>002</u>	May 28, 1997
<u>AB</u>	WATSON LABS	<u>250MG</u>	<u>N88882</u>	<u>001</u>	Oct 22, 1985

ACETAZOLAMIDE SODIUM

INJECTABLE; INJECTION

ACETAZOLAMIDE SODIUM

<u>AP</u>	BEDFORD	<u>EQ 500MG BASE/VIAL</u>	<u>N40089</u>	<u>001</u>	Feb 28, 1995
<u>AP</u>	HOSPIRA	<u>EQ 500MG BASE/VIAL</u>	<u>N40108</u>	<u>001</u>	Oct 30, 1995
<u>DIAMOX</u>					
<u>AP</u>	+ DURAMED PHARMS BARR	<u>EQ 500MG BASE/VIAL</u>	<u>N09388</u>	<u>001</u>	Dec 05, 1990

ACETIC ACID, GLACIAL

SOLUTION; IRRIGATION, URETHRAL

ACETIC ACID 0.25% IN PLASTIC CONTAINER

<u>AT</u>	B BRAUN	<u>250MG/100ML</u>	<u>N18161</u>	<u>001</u>	
<u>AT</u>	BAXTER HLTHCARE	<u>250MG/100ML</u>	<u>N18523</u>	<u>001</u>	Feb 19, 1982
<u>AT</u>	HOSPIRA	<u>250MG/100ML</u>	<u>N17656</u>	<u>001</u>	

SOLUTION/DROPS; OTIC

ACETASOL

<u>AT</u>	ACTAVIS MID ATLANTIC	<u>2%</u>	<u>N87146</u>	<u>001</u>	
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ACETIC ACID

<u>AT</u>	+ MORTON GROVE	<u>2%</u>	<u>N40166</u>	<u>001</u>	Jul 26, 1996
<u>AT</u>	TARO	<u>2%</u>	<u>N88638</u>	<u>001</u>	Sep 06, 1984
<u>AT</u>	VINTAGE	<u>2%</u>	<u>N40607</u>	<u>001</u>	Feb 24, 2005

ACETIC ACID, GLACIAL; ALUMINUM ACETATE

SOLUTION/DROPS; OTIC

	+ BAUSCH AND LOMB	2%; 0.79%	N40063	001	Feb 25, 1994
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ACETIC ACID, GLACIAL; HYDROCORTISONE

SOLUTION/DROPS; OTIC

ACETASOL HC

<u>AT</u>	ACTAVIS MID ATLANTIC	<u>2%; 1%</u>	<u>N87143</u>	<u>001</u>	Jan 13, 1982
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HYDROCORTISONE AND ACETIC ACID

<u>AT</u>	MORTON GROVE	<u>2%; 1%</u>	<u>N40168</u>	<u>001</u>	Aug 30, 1996
<u>AT</u>	TARO	<u>2%; 1%</u>	<u>N88759</u>	<u>001</u>	Mar 04, 1985
<u>AT</u>	VINTAGE	<u>2%; 1%</u>	<u>N40609</u>	<u>001</u>	Feb 06, 2006

VOSOL HC

<u>AT</u>	+ MEDPOINTE PHARM HLC	<u>2%; 1%</u>	<u>N12770</u>	<u>001</u>	
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27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 9 (of 370)

ACETOHEXAMIDE

TABLET; ORAL

ACETOHEXAMIDE

<u>AB</u>	BARR	<u>250MG</u>	<u>N70869</u>	<u>001</u>	Feb 09, 1987
<u>AB</u>	+	<u>500MG</u>	<u>N70870</u>	<u>001</u>	Feb 09, 1987
<u>AB</u>	WATSON LABS	<u>250MG</u>	<u>N71893</u>	<u>001</u>	Nov 25, 1987
<u>AB</u>		<u>500MG</u>	<u>N71894</u>	<u>001</u>	Nov 25, 1987

ACETOHYDROXAMIC ACID

TABLET; ORAL

LITHOSTAT

+	MISSION PHARMA	250MG	N18749	001	May 31, 1983
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ACETYLCHOLINE CHLORIDE

FOR SOLUTION; OPHTHALMIC

MIOCHOL-E

+	NOVARTIS	20MG/VIAL	N20213	001	Sep 22, 1993
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ACETYLCYSTEINE

INJECTABLE; INTRAVENOUS

ACETADOTE

+	CUMBERLAND PHARMS	6GM/30ML (200MG/ML)	N21539	001	Jan 23, 2004
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SOLUTION; INHALATION, ORAL

ACETYLCYSTEINE

<u>AN</u>	BEDFORD	<u>10%</u>	<u>N72323</u>	<u>001</u>	Apr 30, 1992
<u>AN</u>		<u>20%</u>	<u>N72324</u>	<u>001</u>	Apr 30, 1992
<u>AN</u>	HOSPIRA	<u>10%</u>	<u>N73664</u>	<u>001</u>	Aug 30, 1994
<u>AN</u>		<u>20%</u>	<u>N74037</u>	<u>001</u>	Aug 30, 1994
<u>AN</u>	LUITPOLD	<u>10%</u>	<u>N72489</u>	<u>001</u>	Jul 28, 1995
<u>AN</u>		<u>20%</u>	<u>N72547</u>	<u>001</u>	Jul 28, 1995
<u>AN</u>	MAYNE PHARMA USA	<u>10%</u>	<u>N71364</u>	<u>001</u>	May 01, 1989
<u>AN</u>		<u>10%</u>	<u>N71365</u>	<u>002</u>	Jan 18, 2002
<u>AN</u>		<u>20%</u>	<u>N71364</u>	<u>002</u>	Jan 18, 2002
<u>AN</u>		<u>20%</u>	<u>N71365</u>	<u>001</u>	May 01, 1989
	<u>MUCOMYST</u>				
<u>AN</u>	+	<u>10%</u>	<u>N13601</u>	<u>002</u>	
<u>AN</u>	+	<u>20%</u>	<u>N13601</u>	<u>001</u>	

ACITRETIN

CAPSULE; ORAL

SORIATANE

	CONNETICS	10MG	N19821	001	Oct 28, 1996
+		25MG	N19821	002	Oct 28, 1996

ACRIVASTINE; PSEUDOEPHEDRINE HYDROCHLORIDE

CAPSULE; ORAL

SEMPREX-D

+	UCB INC	8MG; 60MG	N19806	001	Mar 25, 1994
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ACYCLOVIR

CAPSULE; ORAL

ACYCLOVIR

<u>AB</u>	ACTAVIS ELIZABETH	<u>200MG</u>	<u>N74906</u>	<u>001</u>	Aug 26, 1997
<u>AB</u>	APOTEX INC	<u>200MG</u>	<u>N75677</u>	<u>001</u>	Sep 28, 2005
<u>AB</u>	APOTHECON	<u>200MG</u>	<u>N74889</u>	<u>001</u>	Oct 31, 1997
<u>AB</u>	CLONMEL HLTHCARE	<u>200MG</u>	<u>N74833</u>	<u>001</u>	Apr 22, 1997
<u>AB</u>		<u>200MG</u>	<u>N74872</u>	<u>001</u>	Apr 22, 1997
<u>AB</u>	GENPHARM	<u>200MG</u>	<u>N74977</u>	<u>001</u>	Apr 13, 1998
<u>AB</u>	IVAX PHARMS	<u>200MG</u>	<u>N74674</u>	<u>001</u>	Apr 22, 1997

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 10 (of 370)

ACYCLOVIR

CAPSULE; ORAL

ACYCLOVIR

<u>AB</u>	MYLAN	<u>200MG</u>	<u>N74727</u>	<u>001</u>	Apr 22, 1997
<u>AB</u>	RANBAXY	<u>200MG</u>	<u>N74975</u>	<u>001</u>	Sep 30, 1998
<u>AB</u>	STASON	<u>200MG</u>	<u>N75090</u>	<u>001</u>	Jan 26, 1999
<u>AB</u>	TEVA	<u>200MG</u>	<u>N74578</u>	<u>001</u>	Apr 22, 1997
<u>AB</u>		<u>200MG</u>	<u>N74828</u>	<u>001</u>	Apr 22, 1997
<u>AB</u>	TEVA PHARMS	<u>200MG</u>	<u>N74914</u>	<u>001</u>	Nov 26, 1997
<u>AB</u>	WATSON LABS	<u>200MG</u>	<u>N75101</u>	<u>001</u>	Apr 15, 1998

ZOVIRAX

<u>AB</u>	+ GLAXOSMITHKLINE	<u>200MG</u>	<u>N18828</u>	<u>001</u>	Jan 25, 1985
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CREAM; TOPICAL

ZOVIRAX

	+ GLAXOSMITHKLINE	5%	N21478	001	Dec 30, 2002
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OINTMENT; TOPICAL

ZOVIRAX

	+ GLAXOSMITHKLINE	5%	N18604	001	Mar 29, 1982
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SUSPENSION; ORAL

ACYCLOVIR

<u>AB</u>	ACTAVIS MID ATLANTIC	<u>200MG/5ML</u>	<u>N74738</u>	<u>001</u>	Apr 28, 1997
<u>AB</u>	HI TECH PHARMA	<u>200MG/5ML</u>	<u>N77026</u>	<u>001</u>	Jun 07, 2005

ZOVIRAX

<u>AB</u>	+ GLAXOSMITHKLINE	<u>200MG/5ML</u>	<u>N19909</u>	<u>001</u>	Dec 22, 1989
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TABLET; ORAL

ACYCLOVIR

<u>AB</u>	ACTAVIS ELIZABETH	<u>400MG</u>	<u>N74870</u>	<u>001</u>	Jun 05, 1997
<u>AB</u>		<u>800MG</u>	<u>N74870</u>	<u>002</u>	Jun 05, 1997
<u>AB</u>	APOTEX INC	<u>400MG</u>	<u>N77309</u>	<u>001</u>	Sep 29, 2005
<u>AB</u>		<u>800MG</u>	<u>N77309</u>	<u>002</u>	Sep 29, 2005
<u>AB</u>	APOTHECON	<u>400MG</u>	<u>N74891</u>	<u>001</u>	Oct 31, 1997
<u>AB</u>		<u>800MG</u>	<u>N74891</u>	<u>002</u>	Oct 31, 1997
<u>AB</u>	CARLSBAD	<u>400MG</u>	<u>N75382</u>	<u>001</u>	Apr 30, 1999
<u>AB</u>		<u>800MG</u>	<u>N75382</u>	<u>002</u>	Apr 30, 1999
<u>AB</u>	CLONMEL HLTHCARE	<u>400MG</u>	<u>N74946</u>	<u>001</u>	Nov 19, 1997
<u>AB</u>		<u>800MG</u>	<u>N74946</u>	<u>002</u>	Nov 19, 1997
<u>AB</u>	GENPHARM	<u>400MG</u>	<u>N74976</u>	<u>001</u>	Apr 13, 1998
<u>AB</u>		<u>800MG</u>	<u>N74976</u>	<u>002</u>	Apr 13, 1998
<u>AB</u>	IVAX PHARMS	<u>400MG</u>	<u>N74836</u>	<u>001</u>	Apr 22, 1997
<u>AB</u>		<u>800MG</u>	<u>N74836</u>	<u>002</u>	Apr 22, 1997
<u>AB</u>	MYLAN	<u>400MG</u>	<u>N75211</u>	<u>001</u>	Sep 28, 1998
<u>AB</u>		<u>800MG</u>	<u>N75211</u>	<u>002</u>	Sep 28, 1998
<u>AB</u>	RANBAXY	<u>400MG</u>	<u>N74980</u>	<u>001</u>	Sep 30, 1998
<u>AB</u>		<u>800MG</u>	<u>N74980</u>	<u>002</u>	Sep 30, 1998
<u>AB</u>	TEVA	<u>400MG</u>	<u>N74556</u>	<u>002</u>	Apr 22, 1997
<u>AB</u>		<u>800MG</u>	<u>N74556</u>	<u>003</u>	Apr 22, 1997
<u>AB</u>	TEVA PHARMS	<u>400MG</u>	<u>N75021</u>	<u>001</u>	Mar 18, 1998
<u>AB</u>		<u>800MG</u>	<u>N75021</u>	<u>002</u>	Mar 18, 1998
	<u>ZOVIRAX</u>				
<u>AB</u>	GLAXOSMITHKLINE	<u>400MG</u>	<u>N20089</u>	<u>001</u>	Apr 30, 1991
<u>AB</u>	+	<u>800MG</u>	<u>N20089</u>	<u>002</u>	Apr 30, 1991

ACYCLOVIR SODIUM

INJECTABLE; INJECTION

ACYCLOVIR

<u>AP</u>	SICOR PHARMS	<u>EQ 50MG BASE/ML</u>	<u>N75627</u>	<u>001</u>	Mar 28, 2001
	ACYCLOVIR IN SODIUM CHLORIDE 0.9% PRESERVATIVE FREE				
	+ BAXTER HLTHCARE	EQ 500MG BASE/VIAL	N74885	001	Dec 19, 1997
		EQ 1GM BASE/VIAL	N74885	002	Dec 19, 1997

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 11 (of 370)

ACYCLOVIR SODIUM

INJECTABLE; INJECTION

ACYCLOVIR SODIUM

<u>AP</u>	+	ABRAXIS PHARM	<u>EQ 50MG BASE/ML</u>	<u>N74930</u>	<u>001</u>	May 13, 1998
<u>AP</u>			<u>EQ 500MG BASE/VIAL</u>	<u>N75015</u>	<u>001</u>	Apr 30, 1998
<u>AP</u>		BAXTER HLTHCARE	<u>EQ 500MG BASE/VIAL</u>	<u>N74913</u>	<u>001</u>	Oct 15, 1997
<u>AP</u>			<u>EQ 1GM BASE/VIAL</u>	<u>N74913</u>	<u>002</u>	Oct 15, 1997
<u>AP</u>		BEDFORD	<u>EQ 500MG BASE/VIAL</u>	<u>N74596</u>	<u>002</u>	Apr 22, 1997
<u>AP</u>			<u>EQ 1GM BASE/VIAL</u>	<u>N74596</u>	<u>001</u>	Apr 22, 1997
<u>AP</u>		HOSPIRA	<u>EQ 500MG BASE/VIAL</u>	<u>N74758</u>	<u>001</u>	Apr 22, 1997
<u>AP</u>			<u>EQ 1GM BASE/VIAL</u>	<u>N74758</u>	<u>002</u>	Apr 22, 1997
<u>AP</u>		MAYNE PHARMA USA	<u>EQ 50MG BASE/ML</u>	<u>N75065</u>	<u>001</u>	Feb 25, 1999
	+		EQ 25MG BASE/ML	N74720	001	Apr 22, 1997
<u>AP</u>		SICOR PHARMS	<u>EQ 500MG BASE/VIAL</u>	<u>N74969</u>	<u>001</u>	Aug 26, 1997
<u>AP</u>			<u>EQ 1GM BASE/VIAL</u>	<u>N74969</u>	<u>002</u>	Aug 26, 1997
<u>ZOVIRAX</u>						
<u>AP</u>	+	GLAXOSMITHKLINE	<u>EQ 500MG BASE/VIAL</u>	<u>N18603</u>	<u>001</u>	Oct 22, 1982
<u>AP</u>	+		<u>EQ 1GM BASE/VIAL</u>	<u>N18603</u>	<u>002</u>	Jun 29, 1989

ADAPALENECREAM; TOPICAL
DIFFERIN

+ GALDERMA LABS LP 0.1% N20748 001 May 26, 2000

GEL; TOPICAL
DIFFERIN

+ GALDERMA LABS LP 0.1% N20380 001 May 31, 1996

ADEFOVIR DIPIVOXILTABLET; ORAL
HEPSERA

+ GILEAD 10MG N21449 001 Sep 20, 2002

ADENOSINE

INJECTABLE; INJECTION

ADENOCARD

<u>AP</u>	+	ASTELLAS	<u>3MG/ML</u>	<u>N19937</u>	<u>002</u>	Oct 30, 1989
		ADENOSCAN				
	+	ASTELLAS	3MG/ML	N20059	001	May 18, 1995

ADENOSINE

<u>AP</u>		ABRAXIS PHARM	<u>3MG/ML</u>	<u>N77133</u>	<u>001</u>	Apr 27, 2005
<u>AP</u>		BAXTER HLTHCARE	<u>3MG/ML</u>	<u>N76500</u>	<u>001</u>	Jun 16, 2004
<u>AP</u>			<u>3MG/ML</u>	<u>N76501</u>	<u>001</u>	Jun 16, 2004
<u>AP</u>		BEDFORD	<u>3MG/ML</u>	<u>N76404</u>	<u>001</u>	Jun 16, 2004
<u>AP</u>		SICOR PHARMS	<u>3MG/ML</u>	<u>N76564</u>	<u>001</u>	Jun 16, 2004

ALBENDAZOLETABLET; ORAL
ALBENZA

+ GLAXOSMITHKLINE 200MG N20666 001 Jun 11, 1996

ALBUMIN HUMANINJECTABLE; INJECTION
OPTISON

+ GE HEALTHCARE 10MG/ML N20899 001 Dec 31, 1997

ALBUMIN IODINATED I-125 SERUMINJECTABLE; INJECTION
JEANATOPE

ISO TEX 100UCI/10ML(10UCI/ML) N17836 003 Jun 08, 2004

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 12 (of 370)

ALBUMIN IODINATED I-125 SERUM

INJECTABLE; INJECTION

JEANATOPE

ISO TEX	500uCi/0.5ML	N17836	001
+	1,000uCi/ML	N17836	002

ALBUMIN IODINATED I-131 SERUM

INJECTABLE; INJECTION

MEGATOPE

+ ISO TEX	0.5mCi/VIAL	N17837	001
+	1mCi/VIAL	N17837	002

ALBUTEROL

AEROSOL, METERED; INHALATION

ALBUTEROL

<u>AB</u>	ARMSTRONG PHARMS	<u>0.09MG/INH</u>	<u>N72273</u>	<u>001</u>	Aug 14, 1996
<u>AB</u>	IVAX PHARMS	<u>0.09MG/INH</u>	<u>N73272</u>	<u>001</u>	Dec 28, 1995

PROVENTIL

BN + SCHERING	0.09MG/INH	N17559	001
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VENTOLIN

<u>AB</u> + GLAXOSMITHKLINE	<u>0.09MG/INH</u>	<u>N18473</u>	<u>001</u>
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ALBUTEROL SULFATE

AEROSOL, METERED; INHALATION

PROAIR HFA

BX + IVAX RES	EQ 0.09MG BASE/INH	N21457	001	Oct 29, 2004
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PROVENTIL-HFA

BX + 3M	EQ 0.09MG BASE/INH	N20503	001	Aug 15, 1996
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VENTOLIN HFA

BX + GLAXOSMITHKLINE	EQ 0.09MG BASE/INH	N20983	001	Apr 19, 2001
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SOLUTION; INHALATION

ACCUNE

<u>AN</u> + DEY	<u>EQ 0.042% BASE</u>	<u>N20949</u>	<u>001</u>	Apr 30, 2001
+	EQ 0.021% BASE	N20949	002	Apr 30, 2001

ALBUTEROL SULFATE

<u>AN</u>	ACTAVIS MID ATLANTIC	<u>EQ 0.083% BASE</u>	<u>N73533</u>	<u>001</u>	Sep 26, 1995
<u>AN</u> + BAUSCH AND LOMB	<u>EQ 0.083% BASE</u>	<u>N75358</u>	<u>001</u>	Mar 29, 2000	
<u>AN</u> +	<u>EQ 0.5% BASE</u>	<u>N75050</u>	<u>001</u>	Jun 18, 1998	
<u>AN</u> + DEY	<u>EQ 0.083% BASE</u>	<u>N72652</u>	<u>001</u>	Feb 21, 1992	
<u>AN</u> HI TECH PHARMA	<u>EQ 0.083% BASE</u>	<u>N75063</u>	<u>001</u>	Feb 09, 1999	
<u>AN</u>	<u>EQ 0.5% BASE</u>	<u>N74543</u>	<u>001</u>	Jan 15, 1998	
<u>AN</u> IVAX PHARMS	<u>EQ 0.083% BASE</u>	<u>N75343</u>	<u>001</u>	Nov 09, 1999	
<u>AN</u> MORTON GROVE	<u>EQ 0.083% BASE</u>	<u>N75394</u>	<u>001</u>	Nov 22, 1999	
<u>AN</u> NEPHRON	<u>EQ 0.042% BASE</u>	<u>N76355</u>	<u>001</u>	Jun 28, 2004	
<u>AN</u>	<u>EQ 0.083% BASE</u>	<u>N74880</u>	<u>001</u>	Sep 17, 1997	
<u>AN</u>	<u>EQ 0.5% BASE</u>	<u>N75664</u>	<u>001</u>	Jun 26, 2001	
<u>AN</u> NOVEX	<u>EQ 0.5% BASE</u>	<u>N76391</u>	<u>001</u>	Apr 01, 2003	
<u>AN</u> RESPIRARE	<u>EQ 0.083% BASE</u>	<u>N76370</u>	<u>001</u>	Nov 24, 2003	
<u>AN</u> ROXANE	<u>EQ 0.083% BASE</u>	<u>N75129</u>	<u>001</u>	Feb 13, 2001	
<u>AN</u> RXELITE	<u>EQ 0.083% BASE</u>	<u>N77569</u>	<u>001</u>	Apr 04, 2006	

SYRUP; ORAL

ALBUTEROL SULFATE

<u>AA</u>	ACTAVIS MID ATLANTIC	<u>EQ 2MG BASE/5ML</u>	<u>N74454</u>	<u>001</u>	Sep 25, 1995
<u>AA</u>		<u>EQ 2MG BASE/5ML</u>	<u>N75262</u>	<u>001</u>	Mar 30, 1999
<u>AA</u> HI TECH PHARMA	<u>EQ 2MG BASE/5ML</u>	<u>N74749</u>	<u>001</u>	Jan 30, 1998	
<u>AA</u> + TEVA	<u>EQ 2MG BASE/5ML</u>	<u>N73419</u>	<u>001</u>	Mar 30, 1992	
<u>AA</u> VINTAGE	<u>EQ 2MG BASE/5ML</u>	<u>N78105</u>	<u>001</u>	Dec 27, 2006	

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 13 (of 370)

ALBUTEROL SULFATE

TABLET; ORAL

ALBUTEROL SULFATE

<u>AB</u>	MUTUAL PHARM	<u>EQ 2MG BASE</u>	<u>N72636</u>	<u>001</u>	Dec 05, 1989
<u>AB</u>		<u>EQ 4MG BASE</u>	<u>N72637</u>	<u>001</u>	Dec 05, 1989
<u>AB</u>	+ MYLAN	<u>EQ 2MG BASE</u>	<u>N72894</u>	<u>002</u>	Jan 17, 1991
<u>AB</u>	+	<u>EQ 4MG BASE</u>	<u>N72894</u>	<u>001</u>	Jan 17, 1991
<u>AB</u>	PLIVA	<u>EQ 2MG BASE</u>	<u>N72316</u>	<u>001</u>	Dec 05, 1989
<u>AB</u>		<u>EQ 4MG BASE</u>	<u>N72317</u>	<u>001</u>	Dec 05, 1989
<u>AB</u>	SANDOZ	<u>EQ 2MG BASE</u>	<u>N72151</u>	<u>001</u>	Dec 05, 1989
<u>AB</u>		<u>EQ 4MG BASE</u>	<u>N72152</u>	<u>001</u>	Dec 05, 1989
<u>AB</u>	TEVA	<u>EQ 2MG BASE</u>	<u>N72619</u>	<u>001</u>	Dec 05, 1989
<u>AB</u>		<u>EQ 2MG BASE</u>	<u>N72779</u>	<u>001</u>	Jun 25, 1993
<u>AB</u>		<u>EQ 2MG BASE</u>	<u>N72938</u>	<u>001</u>	Mar 30, 1990
<u>AB</u>		<u>EQ 4MG BASE</u>	<u>N72620</u>	<u>001</u>	Dec 05, 1989
<u>AB</u>		<u>EQ 4MG BASE</u>	<u>N72780</u>	<u>001</u>	Jun 25, 1993
<u>AB</u>		<u>EQ 4MG BASE</u>	<u>N72939</u>	<u>001</u>	Mar 30, 1990
<u>AB</u>	WATSON LABS	<u>EQ 2MG BASE</u>	<u>N72629</u>	<u>001</u>	Jan 31, 1991
<u>AB</u>		<u>EQ 2MG BASE</u>	<u>N72764</u>	<u>001</u>	Aug 28, 1991
<u>AB</u>		<u>EQ 4MG BASE</u>	<u>N72630</u>	<u>001</u>	Jan 31, 1991
<u>AB</u>		<u>EQ 4MG BASE</u>	<u>N72765</u>	<u>001</u>	Aug 28, 1991

TABLET, EXTENDED RELEASE; ORAL

VOSPIRE ER

	DAVA PHARMS INC	EQ 4MG BASE	N76130	002	Sep 26, 2002
+		EQ 8MG BASE	N76130	003	Sep 26, 2002

ALBUTEROL SULFATE; IPRATROPIUM BROMIDE

AEROSOL, METERED; INHALATION

COMBIVENT

+	BOEHRINGER INGELHEIM	EQ 0.09MG BASE/INH;0.018MG/INH	N20291	001	Oct 24, 1996
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SOLUTION; INHALATION

ALBUTEROL SULFATE AND IPRATROPIUM BROMIDE

<u>AN</u>	SANDOZ	<u>EQ 0.083% BASE;0.017%</u>	<u>N76867</u>	<u>001</u>	Dec 21, 2006
	<u>DUONEB</u>				
<u>AN</u>	+ DEY	<u>EQ 0.083% BASE;0.017%</u>	<u>N20950</u>	<u>001</u>	Mar 21, 2001

ALCLOMETASONE DIPROPIONATE

CREAM; TOPICAL

ACLOVATE

<u>AB</u>	+ GLAXOSMITHKLINE	<u>0.05%</u>	<u>N18707</u>	<u>001</u>	Dec 14, 1982
	<u>ALCLOMETASONE DIPROPIONATE</u>				
<u>AB</u>	ALTANA	<u>0.05%</u>	<u>N76973</u>	<u>001</u>	Jul 12, 2005
<u>AB</u>	TARO	<u>0.05%</u>	<u>N76587</u>	<u>001</u>	Sep 15, 2005

OINTMENT; TOPICAL

ACLOVATE

<u>AB</u>	+ GLAXOSMITHKLINE	<u>0.05%</u>	<u>N18702</u>	<u>001</u>	Dec 14, 1982
	<u>ALCLOMETASONE DIPROPIONATE</u>				
<u>AB</u>	ALTANA	<u>0.05%</u>	<u>N76884</u>	<u>001</u>	Jul 18, 2005
<u>AB</u>	TARO	<u>0.05%</u>	<u>N76730</u>	<u>001</u>	Jul 29, 2004

ALCOHOL; DEXTROSE

INJECTABLE; INJECTION

ALCOHOL 10% AND DEXTROSE 5%

+	B BRAUN	10ML/100ML;5GM/100ML	N04589	006	
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ALCOHOL 5% AND DEXTROSE 5%

<u>AP</u>	+ B BRAUN	<u>5ML/100ML;5GM/100ML</u>	<u>N04589</u>	<u>004</u>	
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ALCOHOL 5% IN D5-W

<u>AP</u>	HOSPIRA	<u>5ML/100ML;5GM/100ML</u>	<u>N83263</u>	<u>001</u>	
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27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 14 (of 370)

ALENDRONATE SODIUM

SOLUTION; ORAL

FOSAMAX

+ MERCK

EQ 70MG BASE/75ML

N21575 001

Sep 17, 2003

TABLET; ORAL

FOSAMAX

MERCK

EQ 5MG BASE

N20560 003

Apr 25, 1997

EQ 10MG BASE

N20560 001

Sep 29, 1995

EQ 35MG BASE

N20560 004

Oct 20, 2000

EQ 40MG BASE

N20560 002

Sep 29, 1995

+

EQ 70MG BASE

N20560 005

Oct 20, 2000

ALENDRONATE SODIUM; CHOLECALCIFEROL

TABLET; ORAL

FOSAMAX PLUS D

+ MERCK

EQ 70MG BASE;2,800 IU

N21762 001

Apr 07, 2005

ALFENTANIL HYDROCHLORIDE

INJECTABLE; INJECTION

ALFENTAAP + AKORNEQ 0.5MG BASE/MLN19353 001

Dec 29, 1986

ALFENTANILAP HOSPIRAEQ 0.5MG BASE/MLN75221 001

Oct 28, 1999

ALFUZOSIN HYDROCHLORIDE

TABLET, EXTENDED RELEASE; ORAL

UROXATRAL

+ SANOFI AVENTIS US

10MG

N21287 001

Jun 12, 2003

ALGLUCERASE

INJECTABLE; INJECTION

CEREDASE

+ GENZYME

80 UNITS/ML

N20057 003

Apr 05, 1991

ALITRETINOIN

GEL; TOPICAL

PANRETIN

+ EISAI MEDCL RES

EQ 0.1% BASE

N20886 001

Feb 02, 1999

ALLOPURINOL

TABLET; ORAL

ALLOPURINOLAB APOTEX INC100MGN77353 001

Sep 08, 2005

AB300MGN77353 002

Sep 08, 2005

AB

MUTUAL PHARM

100MGN71449 001

Jan 09, 1987

AB300MGN71450 001

Jan 09, 1987

AB

MYLAN

100MGN18659 001

Oct 24, 1986

AB300MGN18659 002

Oct 24, 1986

AB

PAR PHARM

100MGN70150 001

Dec 10, 1985

AB300MGN70147 001

Dec 10, 1985

AB

SANDOZ

100MGN70268 001

Dec 31, 1985

AB300MGN70269 001

Dec 31, 1985

AB

VINTAGE PHARMS

100MGN75798 001

Jun 27, 2003

AB300MGN75798 002

Jun 27, 2003

AB

WATSON LABS

100MGN18832 002

Sep 28, 1984

AB300MGN18877 001

Sep 28, 1984

LOPURINAB BASF100MGN71586 001

Apr 02, 1987

AB300MGN71587 001

Apr 02, 1987

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 15 (of 370)

ALLOPURINOL

TABLET; ORAL

ZYLOPRIM

<u>AB</u>	PROMETHEUS LABS	<u>100MG</u>	<u>N16084</u>	<u>001</u>	
<u>AB</u>	+	<u>300MG</u>	<u>N16084</u>	<u>002</u>	

ALLOPURINOL SODIUM

INJECTABLE; INJECTION

ALLOPURINOL SODIUM

<u>AP</u>	BEDFORD LABS	<u>EQ 500MG BASE/VIAL</u>	<u>N76870</u>	<u>001</u>	Aug 26, 2004
<u>AP</u>	+	<u>EQ 500MG BASE/VIAL</u>	<u>N20298</u>	<u>001</u>	May 17, 1996

ALMOTRIPTAN MALATE

TABLET; ORAL

AXERT

	ORTHO MCNEIL PHARM	EQ 6.25MG BASE	N21001	001	May 07, 2001
+		EQ 12.5MG BASE	N21001	002	May 07, 2001

ALOSETRON HYDROCHLORIDE

TABLET; ORAL

LOTRONEX

	GLAXOSMITHKLINE	EQ 0.5MG BASE	N21107	002	Dec 23, 2003
+		EQ 1MG BASE	N21107	001	Feb 09, 2000

ALPHA-TOCOPHEROL ACETATE; ASCORBIC ACID; BIOTIN; CHOLECALCIFEROL; CYANOCOBALAMIN;
DEXPANTHENOL; FOLIC ACID; NIACINAMIDE; PYRIDOXINE HYDROCHLORIDE; RIBOFLAVIN
PHOSPHATE SODIUM; THIAMINE HYDROCHLORIDE; VITAMIN A PALMITATE; VITAMIN K

INJECTABLE; INJECTION

INFUVITE ADULT

+	SANDOZ	2 IU/ML; 40MG/ML; 12UGM/ML; 40 IU/ML; 1UGM/ML; 3MG/ML; 120UGM/ML; 8MG/ML; 1.2MG/ML; 0.72MG/ML; 1.2MG/ML; 660 IU/ML; 0.03MG/ML	N21163	001	May 18, 2000
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INJECTABLE; IV (INFUSION)

+	SANDOZ	2 IU/ML; 40MG/ML; 12UGM/ML; 40 IU/ML; 1UGM/ML; 3MG/ML; 120UGM/ML; 8MG/ML; 1.2MG/ML; 0.72MG/ML; 1.2MG/ML; 660 IU/ML; 30UGM/ML	N21559	001	Jun 16, 2003
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ALPRAZOLAM

CONCENTRATE; ORAL

ALPRAZOLAM

+	ROXANE	1MG/ML	N74312	001	Oct 31, 1993
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TABLET; ORAL

ALPRAZOLAM

<u>AB</u>	ACTAVIS ELIZABETH	<u>0.25MG</u>	<u>N74342</u>	<u>001</u>	Oct 31, 1993
<u>AB</u>		<u>0.5MG</u>	<u>N74342</u>	<u>002</u>	Oct 31, 1993
<u>AB</u>		<u>1MG</u>	<u>N74342</u>	<u>003</u>	Oct 31, 1993
<u>AB</u>		<u>2MG</u>	<u>N74342</u>	<u>004</u>	Oct 31, 1993
<u>AB</u>	ALPHAPHARM	<u>0.25MG</u>	<u>N74046</u>	<u>001</u>	Oct 19, 1993
<u>AB</u>		<u>0.5MG</u>	<u>N74046</u>	<u>002</u>	Oct 19, 1993
<u>AB</u>		<u>1MG</u>	<u>N74046</u>	<u>003</u>	Oct 19, 1993
<u>AB</u>		<u>2MG</u>	<u>N74046</u>	<u>004</u>	May 07, 1997
<u>AB</u>	IVAX PHARMS	<u>0.25MG</u>	<u>N74294</u>	<u>001</u>	Jul 29, 1994
<u>AB</u>		<u>0.5MG</u>	<u>N74294</u>	<u>002</u>	Jul 29, 1994
<u>AB</u>		<u>1MG</u>	<u>N74294</u>	<u>003</u>	Jul 29, 1994
<u>AB</u>		<u>2MG</u>	<u>N74294</u>	<u>004</u>	Jul 29, 1994
<u>AB</u>	MYLAN	<u>0.25MG</u>	<u>N74215</u>	<u>001</u>	Jan 27, 1994

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 16 (of 370)

ALPRAZOLAM

TABLET; ORAL

ALPRAZOLAM

<u>AB</u>	MYLAN	<u>0.5MG</u>	<u>N74215</u>	<u>002</u>	Jan 27, 1994
<u>AB</u>		<u>1MG</u>	<u>N74215</u>	<u>003</u>	Jan 27, 1994
<u>AB</u>		<u>2MG</u>	<u>N74215</u>	<u>004</u>	Jan 27, 1994
<u>AB</u>	SANDOZ	<u>0.25MG</u>	<u>N74112</u>	<u>001</u>	Dec 29, 1995
<u>AB</u>		<u>0.5MG</u>	<u>N74112</u>	<u>002</u>	Dec 29, 1995
<u>AB</u>		<u>1MG</u>	<u>N74112</u>	<u>003</u>	Dec 29, 1995
<u>AB</u>		<u>2MG</u>	<u>N74909</u>	<u>001</u>	Mar 25, 1998
<u>AB</u>	TEVA	<u>0.25MG</u>	<u>N74085</u>	<u>001</u>	Feb 16, 1994
<u>AB</u>		<u>0.5MG</u>	<u>N74085</u>	<u>002</u>	Feb 16, 1994
<u>AB</u>		<u>1MG</u>	<u>N74085</u>	<u>003</u>	Feb 16, 1994
<u>AB</u>		<u>2MG</u>	<u>N74085</u>	<u>004</u>	Feb 26, 1996
<u>AB</u>	WATSON LABS	<u>0.25MG</u>	<u>N74456</u>	<u>001</u>	Aug 31, 1995
<u>AB</u>		<u>0.25MG</u>	<u>N74479</u>	<u>001</u>	Jan 21, 1997
<u>AB</u>		<u>0.5MG</u>	<u>N74456</u>	<u>002</u>	Aug 31, 1995
<u>AB</u>		<u>0.5MG</u>	<u>N74479</u>	<u>002</u>	Jan 21, 1997
<u>AB</u>		<u>1MG</u>	<u>N74456</u>	<u>003</u>	Aug 31, 1995
<u>AB</u>		<u>1MG</u>	<u>N74479</u>	<u>003</u>	Jan 21, 1997

XANAX

<u>AB</u>	PHARMACIA AND UPJOHN	<u>0.25MG</u>	<u>N18276</u>	<u>001</u>	
<u>AB</u>		<u>0.5MG</u>	<u>N18276</u>	<u>002</u>	
<u>AB</u>	+	<u>1MG</u>	<u>N18276</u>	<u>003</u>	
<u>AB</u>		<u>2MG</u>	<u>N18276</u>	<u>004</u>	Nov 27, 1985

TABLET, EXTENDED RELEASE; ORAL

ALPRAZOLAM

<u>AB</u>	BARR	<u>0.5MG</u>	<u>N77725</u>	<u>001</u>	Jul 31, 2006
<u>AB</u>		<u>1MG</u>	<u>N77725</u>	<u>002</u>	Jul 31, 2006
<u>AB</u>		<u>2MG</u>	<u>N77725</u>	<u>004</u>	Jul 31, 2006
<u>AB</u>		<u>3MG</u>	<u>N77725</u>	<u>003</u>	Jul 31, 2006
<u>AB</u>	MYLAN	<u>0.5MG</u>	<u>N77391</u>	<u>002</u>	Jan 26, 2006
<u>AB</u>		<u>1MG</u>	<u>N77391</u>	<u>003</u>	Jan 26, 2006
<u>AB</u>		<u>2MG</u>	<u>N77391</u>	<u>004</u>	Jan 26, 2006
<u>AB</u>		<u>3MG</u>	<u>N77391</u>	<u>001</u>	Jan 26, 2006
<u>AB</u>	SANDOZ	<u>0.5MG</u>	<u>N77777</u>	<u>001</u>	Jun 30, 2006
<u>AB</u>		<u>1MG</u>	<u>N77777</u>	<u>002</u>	Jun 30, 2006
<u>AB</u>		<u>2MG</u>	<u>N77777</u>	<u>003</u>	Jun 30, 2006
<u>AB</u>		<u>3MG</u>	<u>N77777</u>	<u>004</u>	Jun 30, 2006

XANAX XR

<u>AB</u>	PHARMACIA AND UPJOHN	<u>0.5MG</u>	<u>N21434</u>	<u>001</u>	Jan 17, 2003
<u>AB</u>		<u>1MG</u>	<u>N21434</u>	<u>002</u>	Jan 17, 2003
<u>AB</u>		<u>2MG</u>	<u>N21434</u>	<u>003</u>	Jan 17, 2003
<u>AB</u>	+	<u>3MG</u>	<u>N21434</u>	<u>004</u>	Jan 17, 2003

TABLET, ORALLY DISINTEGRATING; ORAL

NIRAVAM

	SCHWARZ PHARMA	0.25MG	N21726	001	Jan 19, 2005
		0.5MG	N21726	002	Jan 19, 2005
		1MG	N21726	003	Jan 19, 2005
	+	2MG	N21726	004	Jan 19, 2005

ALPROSTADIL

INJECTABLE; INJECTION

ALPROSTADIL

<u>AP</u>	BEDFORD	<u>0.5MG/ML</u>	<u>N74815</u>	<u>001</u>	Jan 20, 1998
<u>AP</u>	SICOR PHARMS	<u>0.5MG/ML</u>	<u>N75196</u>	<u>001</u>	Apr 30, 1999
	<u>CAVERJECT</u>				
<u>AP</u>	PHARMACIA AND UPJOHN	<u>0.01MG/VIAL</u>	<u>N20379</u>	<u>001</u>	Jul 06, 1995
<u>AP</u>	+	<u>0.02MG/VIAL</u>	<u>N20379</u>	<u>002</u>	Jul 06, 1995
<u>AP</u>	+	<u>0.04MG/VIAL</u>	<u>N20379</u>	<u>004</u>	May 19, 1997

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 17 (of 370)

ALPROSTADIL

INJECTABLE; INJECTION

CAVERJECT

PHARMACIA AND UPJOHN 0.005MG/VIAL
 0.01MG/VIAL
 0.02MG/VIAL

N20379 003 Jun 27, 1996
 N21212 001 Jun 11, 2002
 N21212 002 Jun 11, 2002

EDEX

AP SCHWARZ PHARMA 0.01MG/VIAL
AP 0.02MG/VIAL
AP + 0.04MG/VIAL
 + 0.01MG/VIAL
 + 0.02MG/VIAL
 + 0.04MG/VIAL

N20649 002 Jun 12, 1997
N20649 003 Jun 12, 1997
N20649 004 Jun 12, 1997
 N20649 005 Jul 30, 1998
 N20649 006 Jul 30, 1998
 N20649 007 Jul 30, 1998

PROSTIN VR PEDIATRIC

AP + PHARMACIA AND UPJOHN 0.5MG/ML

N18484 001

SUPPOSITORY; URETHRAL

MUSE

VIVUS 0.125MG
 0.25MG
 0.5MG
 + 1MG

N20700 001 Nov 19, 1996
 N20700 002 Nov 19, 1996
 N20700 003 Nov 19, 1996
 N20700 004 Nov 19, 1996

ALTRETAMINE

CAPSULE; ORAL

HEXALEN

+ MGI PHARMA INC 50MG

N19926 001 Dec 26, 1990

AMANTADINE HYDROCHLORIDE

CAPSULE; ORAL

AMANTADINE HYDROCHLORIDE

AB ACTAVIS TOTOWA 100MG
AB SANDOZ 100MG
AB + USL PHARMA 100MG

N77659 001 Feb 23, 2006
N71293 001 Feb 18, 1987
N70589 001 Aug 05, 1986

SYRUP; ORAL

AMANTADINE HYDROCHLORIDE

AA + ACTAVIS MID ATLANTIC 50MG/5ML
AA + CAROLINA MEDCL 50MG/5ML
AA + HI TECH PHARMA 50MG/5ML
AA + MIKART 50MG/5ML
AA + MORTON GROVE 50MG/5ML
AA + PHARM ASSOC 50MG/5ML
AA + SILARX 50MG/5ML
AA + TEVA PHARMS 50MG/5ML
AA + VINTAGE 50MG/5ML

N72655 001 Oct 30, 1990
N75819 001 Sep 11, 2002
N74170 001 Oct 28, 1994
N74028 001 Jun 28, 1993
N75060 001 Dec 24, 1998
N74509 001 Jul 17, 1995
N76352 001 Sep 10, 2004
N73115 001 Aug 23, 1991
N77992 001 Dec 12, 2006

TABLET; ORAL

AMANTADINE HYDROCHLORIDE

AB USL PHARMA 100MG

N76186 001 Dec 16, 2002

SYMMETREL

AB + ENDO PHARMS 100MG

N18101 001

AMBENONIUM CHLORIDE

TABLET; ORAL

MYTELASE

+ SANOFI AVENTIS US 10MG

N10155 002

AMCINONIDE

CREAM; TOPICAL

AMCINONIDE

AB + ALTANA 0.1%

N76065 001 May 15, 2003

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 18 (of 370)

AMCINONIDE

CREAM; TOPICAL

AMCINONIDE

<u>AB</u>	TARO PHARM INDS	<u>0.1%</u>	<u>N76229</u>	<u>001</u>	May 31, 2002
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LOTION; TOPICAL

AMCINONIDE

	+ ALTANA	0.1%	N76329	001	Nov 06, 2002
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OINTMENT; TOPICAL

AMCINONIDE

<u>AB</u>	+ ALTANA	<u>0.1%</u>	<u>N76096</u>	<u>001</u>	Nov 19, 2002
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<u>AB</u>	TARO PHARM INDS	<u>0.1%</u>	<u>N76367</u>	<u>001</u>	Mar 19, 2003
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AMIFOSTINE

INJECTABLE; INJECTION

ETHYOL

	+ MEDIMMUNE ONCOLOGY	500MG/VIAL	N20221	001	Dec 08, 1995
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AMIKACIN SULFATE

INJECTABLE; INJECTION

AMIKACIN SULFATE

<u>AP</u>	BEDFORD	<u>EQ 50MG BASE/ML</u>	<u>N63313</u>	<u>001</u>	Apr 11, 1994
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<u>AP</u>		<u>EQ 250MG BASE/ML</u>	<u>N63315</u>	<u>001</u>	Apr 11, 1994
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<u>AP</u>	HOSPIRA	<u>EQ 50MG BASE/ML</u>	<u>N63263</u>	<u>001</u>	Nov 30, 1994
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<u>AP</u>		<u>EQ 250MG BASE/ML</u>	<u>N63264</u>	<u>001</u>	Nov 30, 1994
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<u>AP</u>		<u>EQ 250MG BASE/ML</u>	<u>N64098</u>	<u>001</u>	Jun 26, 1995
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		EQ 62.5MG BASE/ML	N63283	001	Oct 31, 1994
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<u>AP</u>	SICOR PHARMS	<u>EQ 50MG BASE/ML</u>	<u>N64045</u>	<u>001</u>	Sep 28, 1993
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<u>AP</u>		<u>EQ 250MG BASE/ML</u>	<u>N64045</u>	<u>002</u>	Sep 28, 1993
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AMIKACIN SULFATE IN SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER

	HOSPIRA	EQ 500MG BASE/100ML	N64146	001	Apr 02, 1997
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AMIKIN

<u>AP</u>	+ APOTHECON	<u>EQ 50MG BASE/ML</u>	<u>N62311</u>	<u>001</u>	
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<u>AP</u>	+	<u>EQ 250MG BASE/ML</u>	<u>N62311</u>	<u>002</u>	
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AMILORIDE HYDROCHLORIDE

TABLET; ORAL

AMILORIDE HYDROCHLORIDE

	+ PAR PHARM	5MG	N70346	001	Jan 22, 1986
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AMILORIDE HYDROCHLORIDE; HYDROCHLOROTHIAZIDE

TABLET; ORAL

AMILORIDE HYDROCHLORIDE AND HYDROCHLOROTHIAZIDE

<u>AB</u>	BARR	<u>EQ 5MG ANHYDROUS;50MG</u>	<u>N71111</u>	<u>001</u>	May 10, 1988
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<u>AB</u>	+ MYLAN	<u>EQ 5MG ANHYDROUS;50MG</u>	<u>N73209</u>	<u>001</u>	Oct 31, 1991
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<u>AB</u>	SANDOZ	<u>EQ 5MG ANHYDROUS;50MG</u>	<u>N73357</u>	<u>001</u>	Nov 27, 1991
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<u>AB</u>	TEVA	<u>EQ 5MG ANHYDROUS;50MG</u>	<u>N70795</u>	<u>001</u>	Apr 17, 1988
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<u>AB</u>	WATSON LABS	<u>EQ 5MG ANHYDROUS;50MG</u>	<u>N73334</u>	<u>001</u>	Jul 19, 1991
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HYDRO-RIDE

<u>AB</u>	PAR PHARM	<u>EQ 5MG ANHYDROUS;50MG</u>	<u>N70347</u>	<u>001</u>	Dec 25, 1990
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AMINO ACIDS

INJECTABLE; INJECTION

AMINOSYN 10%

	HOSPIRA	10% (10GM/100ML)	N17673	003	
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AMINOSYN 10% (PH6)

	HOSPIRA	10% (10GM/100ML)	N17673	008	Nov 18, 1985
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AMINOSYN 3.5%

	HOSPIRA	3.5% (3.5GM/100ML)	N17789	004	
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27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 19 (of 370)

AMINO ACIDS

INJECTABLE; INJECTION

AMINOSYN 5%				
HOSPIRA	5% (5GM/100ML)	N17673	001	
AMINOSYN 7%				
HOSPIRA	7% (7GM/100ML)	N17673	002	
AMINOSYN 7% (PH6)				
HOSPIRA	7% (7GM/100ML)	N17673	006	Nov 18, 1985
AMINOSYN 8.5%				
HOSPIRA	8.5% (8.5GM/100ML)	N17673	004	
AMINOSYN 8.5% (PH6)				
HOSPIRA	8.5% (8.5GM/100ML)	N17673	007	Nov 18, 1985
AMINOSYN II 10%				
HOSPIRA	10% (10GM/100ML)	N19438	005	Apr 03, 1986
AMINOSYN II 10% IN PLASTIC CONTAINER				
HOSPIRA	10% (10GM/100ML)	N20015	001	Dec 19, 1991
AMINOSYN II 15% IN PLASTIC CONTAINER				
HOSPIRA	15% (15GM/100ML)	N20041	001	Dec 19, 1991
AMINOSYN II 7%				
HOSPIRA	7% (7GM/100ML)	N19438	003	Apr 03, 1986
AMINOSYN II 8.5%				
HOSPIRA	8.5% (8.5GM/100ML)	N19438	004	Apr 03, 1986
AMINOSYN-HBC 7%				
HOSPIRA	7% (7GM/100ML)	N19374	001	Jul 12, 1985
AMINOSYN-HF 8%				
HOSPIRA	8% (8GM/100ML)	N20345	001	Apr 04, 1996
AMINOSYN-PF 10%				
HOSPIRA	10% (10GM/100ML)	N19492	002	Oct 17, 1986
AMINOSYN-PF 7%				
HOSPIRA	7% (7GM/100ML)	N19398	001	Sep 06, 1985
AMINOSYN-RF 5.2%				
HOSPIRA	5.2% (5.2GM/100ML)	N18429	001	
BRANCHAMIN 4% IN PLASTIC CONTAINER				
BAXTER HLTHCARE	4% (4GM/100ML)	N18684	001	Sep 28, 1984
CLINISOL 15% SULFITE FREE IN PLASTIC CONTAINER				
CLINTEC NUTR	15% (15GM/100ML)	N20512	001	Aug 30, 1996
FREAMINE HBC 6.9%				
B BRAUN	6.9% (6.9GM/100ML)	N16822	006	May 17, 1983
FREAMINE III 10%				
B BRAUN	10% (10GM/100ML)	N16822	005	
FREAMINE III 8.5%				
B BRAUN	8.5% (8.5GM/100ML)	N16822	004	
HEPATAMINE 8%				
B BRAUN	8% (8GM/100ML)	N18676	001	Aug 03, 1982
HEPATASOL 8%				
BAXTER HLTHCARE	8% (8GM/100ML)	N20360	001	Apr 04, 1996
NEPHRAMINE 5.4%				
B BRAUN	5.4% (5.4GM/100ML)	N17766	001	
NOVAMINE 11.4%				
HOSPIRA	11.4% (11.4GM/100ML)	N17957	003	Aug 09, 1982
NOVAMINE 15%				
HOSPIRA	15% (15GM/100ML)	N17957	004	Nov 28, 1986
PREMASOL 10% IN PLASTIC CONTAINER				
BAXTER HLTHCARE	10% (10GM/100ML)	N75880	002	Jun 19, 2003
PREMASOL 6% IN PLASTIC CONTAINER				
BAXTER HLTHCARE	6% (6GM/100ML)	N75880	001	Jun 19, 2003
PROSOL 20% SULFITE FREE IN PLASTIC CONTAINER				
+ BAXTER HLTHCARE	20% (20GM/100ML)	N20849	001	Aug 26, 1998
RENAMIN W/O ELECTROLYTES				
BAXTER HLTHCARE	6.5% (6.5GM/100ML)	N17493	007	Oct 15, 1982
TRAVASOL 10% IN PLASTIC CONTAINER				
BAXTER HLTHCARE	10% (10MG/100ML)	N18931	003	Aug 23, 1984

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 20 (of 370)

AMINO ACIDS

INJECTABLE; INJECTION

TRAVASOL 10% W/O ELECTROLYTES

BAXTER HLTHCARE 10% (10GM/100ML) N17493 006

TRAVASOL 5.5% IN PLASTIC CONTAINER

BAXTER HLTHCARE 5.5% (5.5GM/100ML) N18931 001 Aug 23, 1984

TRAVASOL 5.5% W/O ELECTROLYTES

BAXTER HLTHCARE 5.5% (5.5GM/100ML) N17493 004

TRAVASOL 8.5% IN PLASTIC CONTAINER

BAXTER HLTHCARE 8.5% (8.5GM/100ML) N18931 002 Aug 23, 1984

TRAVASOL 8.5% W/O ELECTROLYTES

BAXTER HLTHCARE 8.5% (8.5GM/100ML) N17493 005

TROPHAMINE

+ B BRAUN 6% (6GM/100ML) N19018 001 Jul 20, 1984

TROPHAMINE 10%

+ B BRAUN 10% (10GM/100ML) N19018 003 Sep 07, 1988

AMINO ACIDS; CALCIUM ACETATE; GLYCERIN; MAGNESIUM ACETATE; PHOSPHORIC ACID; POTASSIUM CHLORIDE; SODIUM ACETATE; SODIUM CHLORIDE

INJECTABLE; INJECTION

PROCALAMINE

B BRAUN 3%;26MG/100ML;3GM/100ML;54MG/100ML;41MG
/100ML;150MG/100ML;200MG/100ML;120MG/10
0ML N18582 001 May 08, 1982AMINO ACIDS; CALCIUM CHLORIDE; DEXTROSE; MAGNESIUM CHLORIDE; POTASSIUM CHLORIDE; POTASSIUM PHOSPHATE, DIBASIC; SODIUM CHLORIDE

INJECTABLE; INJECTION

AMINOSYN II 3.5% W/ ELECTROLYTES IN DEXTROSE 25% W/ CALCIUM IN PLASTIC CONTAINER

HOSPIRA 3.5%;36.8MG/100ML;25GM/100ML;51MG/100ML
;22.4MG/100ML;261MG/100ML;205MG/100ML N19683 001 Nov 07, 1988

AMINOSYN II 4.25% W/ ELECTROLYTES IN DEXTROSE 20% W/ CALCIUM IN PLASTIC CONTAINER

HOSPIRA 4.25%;36.8MG/100ML;20GM/100ML;51MG/100M
L;22.4MG/100ML;261MG/100ML;205MG/100ML N19683 002 Nov 07, 1988

AMINOSYN II 4.25% W/ ELECTROLYTES IN DEXTROSE 25% W/ CALCIUM IN PLASTIC CONTAINER

HOSPIRA 4.25%;36.8MG/100ML;25GM/100ML;51MG/100M
L;22.4MG/100ML;261MG/100ML;205MG/100ML N19683 003 Nov 07, 1988AMINO ACIDS; CALCIUM CHLORIDE; DEXTROSE; MAGNESIUM CHLORIDE; POTASSIUM PHOSPHATE, DIBASIC; SODIUM ACETATE; SODIUM CHLORIDE

INJECTABLE; INJECTION

CLINIMIX E 2.75/10 SULFITE-FREE W/ ELECT IN DEXTROSE 10% W/ CALCIUM IN PLASTIC CONTAINER

+ BAXTER HLTHCARE 2.75%;33MG/100ML;10GM/100ML;51MG/100ML;
261MG/100ML;217MG/100ML;112MG/100ML N20678 002 Mar 26, 1997

CLINIMIX E 2.75/25 SULFITE-FREE W/ ELECT IN DEXTROSE 25% W/ CALCIUM IN PLASTIC CONTAINER

+ BAXTER HLTHCARE 2.75%;33MG/100ML;25GM/100ML;51MG/100ML;
261MG/100ML;217MG/100ML;112MG/100ML N20678 005 Mar 26, 1997

CLINIMIX E 2.75/5 SULFITE-FREE W/ ELECT IN DEXTROSE 5% W/ CALCIUM IN PLASTIC CONTAINER

+ BAXTER HLTHCARE 2.75%;33MG/100ML;5GM/100ML;51MG/100ML;2
61MG/100ML;217MG/100ML;112MG/100ML N20678 001 Mar 26, 1997

CLINIMIX E 4.25/10 SULFITE-FREE W/ ELECT IN DEXTROSE 10% W/ CALCIUM IN PLASTIC CONTAINER

+ BAXTER HLTHCARE 4.25%;33MG/100ML;10GM/100ML;51MG/100ML;
261MG/100ML;297MG/100ML;77MG/100ML N20678 009 Mar 26, 1997

CLINIMIX E 4.25/20 SULFITE-FREE W/ ELECT IN DEXTROSE 20% W/ CALCIUM IN PLASTIC CONTAINER

+ BAXTER HLTHCARE 4.25%;33MG/100ML;20GM/100ML;51MG/100ML;
261MG/100ML;297MG/100ML;77MG/100ML N20678 011 Mar 26, 1997

CLINIMIX E 4.25/25 SULFITE-FREE W/ ELECT IN DEXTROSE 25% W/ CALCIUM IN PLASTIC CONTAINER

+ BAXTER HLTHCARE 4.25%;33MG/100ML;25GM/100ML;51MG/100ML;
261MG/100ML;297MG/100ML;77MG/100ML N20678 012 Mar 26, 1997

CLINIMIX E 4.25/5 SULFITE-FREE W/ ELECT IN DEXTROSE 5% W/ CALCIUM IN PLASTIC CONTAINER

+ BAXTER HLTHCARE 4.25%;33MG/100ML;5GM/100ML;51MG/100ML;2
61MG/100ML;297MG/100ML;77MG/100ML N20678 008 Mar 26, 1997

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 21 (of 370)

AMINO ACIDS; CALCIUM CHLORIDE; DEXTROSE; MAGNESIUM CHLORIDE; POTASSIUM PHOSPHATE, DIBASIC; SODIUM ACETATE; SODIUM CHLORIDEINJECTABLE; INJECTION

CLINIMIX E 5/10 SULFITE-FREE W/ ELECT IN DEXTROSE 10% W/ CALCIUM IN PLASTIC CONTAINER			
+ BAXTER HLTHCARE	5%;33MG/100ML;10GM/100ML;51MG/100ML;261 MG/100ML;340MG/100ML;59MG/100ML	N20678 016	Mar 26, 1997
CLINIMIX E 5/15 SULFITE-FREE W/ ELECT IN DEXTROSE 15% W/ CALCIUM IN PLASTIC CONTAINER			
+ BAXTER HLTHCARE	5%;33MG/100ML;15GM/100ML;51MG/100ML;261 MG/100ML;340MG/100ML;59MG/100ML	N20678 017	Mar 26, 1997
CLINIMIX E 5/20 SULFITE-FREE W/ ELECT IN DEXTROSE 20% W/ CALCIUM IN PLASTIC CONTAINER			
+ BAXTER HLTHCARE	5%;33MG/100ML;20GM/100ML;51MG/100ML;261 MG/100ML;340MG/100ML;59MG/100ML	N20678 018	Mar 26, 1997
CLINIMIX E 5/25 SULFITE-FREE W/ ELECT IN DEXTROSE 25% W/ CALCIUM IN PLASTIC CONTAINER			
+ BAXTER HLTHCARE	5%;33MG/100ML;25GM/100ML;51MG/100ML;261 MG/100ML;340MG/100ML;59MG/100ML	N20678 019	Mar 26, 1997
CLINIMIX E 5/35 SULFITE-FREE W/ ELECT IN DEXTROSE 35% W/ CALCIUM IN PLASTIC CONTAINER			
+ BAXTER HLTHCARE	5%;33MG/100ML;35GM/100ML;51MG/100ML;261 MG/100ML;340MG/100ML;59MG/100ML	N20678 021	Mar 26, 1997

AMINO ACIDS; DEXTROSEINJECTABLE; INJECTION

AMINOSYN II 3.5% IN DEXTROSE 25% IN PLASTIC CONTAINER			
HOSPIRA	3.5%;25GM/100ML	N19681 001	Nov 01, 1988
AMINOSYN II 3.5% IN DEXTROSE 5% IN PLASTIC CONTAINER			
HOSPIRA	3.5%;5GM/100ML	N19681 002	Nov 01, 1988
AMINOSYN II 4.25% IN DEXTROSE 10% IN PLASTIC CONTAINER			
HOSPIRA	4.25%;10GM/100ML	N19681 004	Nov 01, 1988
AMINOSYN II 4.25% IN DEXTROSE 20% IN PLASTIC CONTAINER			
HOSPIRA	4.25%;20GM/100ML	N19681 005	Nov 01, 1988
AMINOSYN II 4.25% IN DEXTROSE 25% IN PLASTIC CONTAINER			
HOSPIRA	4.25%;25GM/100ML	N19681 003	Nov 01, 1988
AMINOSYN II 5% IN DEXTROSE 25% IN PLASTIC CONTAINER			
HOSPIRA	5%;25GM/100ML	N19681 006	Nov 01, 1988
CLINIMIX 2.75/10 SULFITE FREE IN DEXTROSE 10% IN PLASTIC CONTAINER			
BAXTER HLTHCARE	2.75%;10GM/100ML	N20734 002	Sep 29, 1997
CLINIMIX 2.75/25 SULFITE FREE IN DEXTROSE 25% IN PLASTIC CONTAINER			
BAXTER HLTHCARE	2.75%;25GM/100ML	N20734 005	Sep 29, 1997
CLINIMIX 2.75/5 SULFITE FREE IN DEXTROSE 5% IN PLASTIC CONTAINER			
BAXTER HLTHCARE	2.75%;5GM/100ML	N20734 001	Sep 29, 1997
CLINIMIX 4.25/10 SULFITE FREE IN DEXTROSE 10% IN PLASTIC CONTAINER			
BAXTER HLTHCARE	4.25%;10GM/100ML	N20734 008	Sep 29, 1997
CLINIMIX 4.25/20 SULFITE FREE IN DEXTROSE 20% IN PLASTIC CONTAINER			
BAXTER HLTHCARE	4.25%;20GM/100ML	N20734 010	Sep 29, 1997
CLINIMIX 4.25/25 SULFITE FREE IN DEXTROSE 25% IN PLASTIC CONTAINER			
BAXTER HLTHCARE	4.25%;25GM/100ML	N20734 011	Sep 29, 1997
CLINIMIX 4.25/5 SULFITE FREE IN DEXTROSE 5% IN PLASTIC CONTAINER			
BAXTER HLTHCARE	4.25%;5GM/100ML	N20734 007	Sep 29, 1997
CLINIMIX 5/10 SULFITE FREE IN DEXTROSE 10% IN PLASTIC CONTAINER			
BAXTER HLTHCARE	5%;10GM/100ML	N20734 014	Sep 29, 1997
CLINIMIX 5/15 SULFITE FREE IN DEXTROSE 15% IN PLASTIC CONTAINER			
BAXTER HLTHCARE	5%;15GM/100ML	N20734 015	Sep 29, 1997
CLINIMIX 5/20 SULFITE FREE IN DEXTROSE 20% IN PLASTIC CONTAINER			
BAXTER HLTHCARE	5%;20GM/100ML	N20734 016	Sep 29, 1997
CLINIMIX 5/25 SULFITE FREE IN DEXTROSE 25% IN PLASTIC CONTAINER			
BAXTER HLTHCARE	5%;25GM/100ML	N20734 017	Sep 29, 1997
CLINIMIX 5/35 SULFITE FREE IN DEXTROSE 35% IN PLASTIC CONTAINER			
BAXTER HLTHCARE	5%;35GM/100ML	N20734 018	Sep 29, 1997

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 22 (of 370)

AMINO ACIDS; DEXTROSE; MAGNESIUM CHLORIDE; POTASSIUM CHLORIDE; SODIUM CHLORIDE; SODIUM PHOSPHATE, DIBASIC, HEPTAHYDRATE

INJECTABLE; INJECTION

AMINOSYN II 3.5% M IN DEXTROSE 5% IN PLASTIC CONTAINER

HOSPIRA	3.5%;5GM/100ML;30MG/100ML;97MG/100ML;120MG/100ML;49.3MG/100ML	N19682 001	Nov 01, 1988
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AMINOSYN II 4.25% M IN DEXTROSE 10% IN PLASTIC CONTAINER

HOSPIRA	4.25%;5GM/100ML;30MG/100ML;97MG/100ML;120MG/100ML;49.3MG/100ML	N19682 002	Nov 01, 1988
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AMINO ACIDS; MAGNESIUM ACETATE; PHOSPHORIC ACID; POTASSIUM ACETATE; POTASSIUM CHLORIDE; SODIUM ACETATE

INJECTABLE; INJECTION

FREAMINE III 8.5% W/ ELECTROLYTES

B BRAUN	8.5%;110MG/100ML;230MG/100ML;10MG/100ML;440MG/100ML;690MG/100ML	N16822 007	Jul 01, 1988
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AMINO ACIDS; MAGNESIUM ACETATE; PHOSPHORIC ACID; POTASSIUM ACETATE; SODIUM CHLORIDE

INJECTABLE; INJECTION

AMINOSYN 3.5% M

HOSPIRA	3.5%;21MG/100ML;40MG/100ML;128MG/100ML;234MG/100ML	N17789 003	
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AMINO ACIDS; MAGNESIUM ACETATE; PHOSPHORIC ACID; POTASSIUM CHLORIDE; SODIUM ACETATE; SODIUM CHLORIDE

INJECTABLE; INJECTION

FREAMINE III 3% W/ ELECTROLYTES

B BRAUN	3%;54MG/100ML;40MG/100ML;150MG/100ML;200MG/100ML;120MG/100ML	N16822 003	
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AMINO ACIDS; MAGNESIUM CHLORIDE; POTASSIUM CHLORIDE; POTASSIUM PHOSPHATE, DIBASIC; SODIUM CHLORIDE

INJECTABLE; INJECTION

AMINOSYN II 10% W/ ELECTROLYTES

HOSPIRA	10%;102MG/100ML;45MG/100ML;522MG/100ML;410MG/100ML	N19437 004	Apr 03, 1986
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AMINOSYN II 8.5% W/ ELECTROLYTES

HOSPIRA	8.5%;102MG/100ML;45MG/100ML;522MG/100ML;410MG/100ML	N19437 005	Apr 03, 1986
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AMINO ACIDS; MAGNESIUM CHLORIDE; POTASSIUM PHOSPHATE, DIBASIC; SODIUM ACETATE; SODIUM CHLORIDE

INJECTABLE; INJECTION

TRAVASOL 3.5% SULFITE FREE W/ ELECTROLYTES IN PLASTIC CONTAINER

BAXTER HLTHCARE	3.5%;51MG/100ML;131MG/100ML;218MG/100ML;35MG/100ML	N20177 001	Oct 23, 1995
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TRAVASOL 3.5% W/ ELECTROLYTES

BAXTER HLTHCARE	3.5%;51MG/100ML;131MG/100ML;218MG/100ML;35MG/100ML	N17493 003	
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TRAVASOL 5.5% SULFITE FREE W/ ELECTROLYTES IN PLASTIC CONTAINER

BAXTER HLTHCARE	5.5%;102MG/100ML;522MG/100ML;431MG/100ML;224MG/100ML	N20173 001	Oct 27, 1995
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TRAVASOL 5.5% W/ ELECTROLYTES

BAXTER HLTHCARE	5.5%;102MG/100ML;522MG/100ML;431MG/100ML;224MG/100ML	N17493 001	
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TRAVASOL 8.5% SULFITE FREE W/ ELECTROLYTES IN PLASTIC CONTAINER

BAXTER HLTHCARE	8.5%;102MG/100ML;522MG/100ML;594MG/100ML;154MG/100ML	N20173 002	Oct 27, 1995
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TRAVASOL 8.5% W/ ELECTROLYTES

BAXTER HLTHCARE	8.5%;102MG/100ML;522MG/100ML;594MG/100ML;154MG/100ML	N17493 002	
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27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 23 (of 370)

AMINO ACIDS; MAGNESIUM CHLORIDE; POTASSIUM PHOSPHATE, DIBASIC; SODIUM CHLORIDE

INJECTABLE; INJECTION

AMINOSYN 7% W/ ELECTROLYTES

HOSPIRA 7%;102MG/100ML;522MG/100ML;410MG/100ML N17789 002

AMINOSYN 8.5% W/ ELECTROLYTES

HOSPIRA 8.5%;102MG/100ML;522MG/100ML;410MG/100ML N17673 005
LAMINOCAPROIC ACID

INJECTABLE; INJECTION

AMICARAP + XANODYNE PHARM 250MG/ML N15229 002AMINOCAPROIC ACIDAP HOSPIRA 250MG/ML N70888 001 Jun 16, 1988AP LUITPOLD 250MG/ML N71192 001 Dec 01, 1987AMINOCAPROIC ACID IN PLASTIC CONTAINERAP HOSPIRA 250MG/ML N70010 001 Mar 09, 1987

SYRUP; ORAL

AMICARAA + XANODYNE PHARM 1.25GM/5ML N15230 002AMINOCAPROIC ACIDAA MIKART 1.25GM/5ML N74759 001 Sep 02, 1998

TABLET; ORAL

AMICARAB + XANODYNE PHARM 500MG N15197 001AMINOCAPROICAB MIKART 500MG N75602 001 May 24, 2001AMINOGLUTETHIMIDE

TABLET; ORAL

CYTADREN

+ NOVARTIS 250MG N18202 001

AMINOHIPPURATE SODIUM

INJECTABLE; INJECTION

AMINOHIPPURATE SODIUM

+ MERCK 20% N05619 001

AMINOLEVULINIC ACID HYDROCHLORIDE

SOLUTION; TOPICAL

LEVULAN

+ DUSA 20% N20965 001 Dec 03, 1999

AMINOPHYLLINE

INJECTABLE; INJECTION

AMINOPHYLLINEAP HOSPIRA 25MG/ML N87242 001 Oct 26, 1983AP + 25MG/ML N87601 001 Jul 23, 1982AP INTL MEDICATION 25MG/ML N87209 001 Feb 01, 1982AP LUITPOLD 25MG/ML N87240 001AP 25MG/ML N87600 001AP PHARMA SERVE NY 25MG/ML N87392 001 Dec 15, 1983AP SICOR PHARMS 25MG/ML N81142 001 Sep 25, 1991

AMINOPHYLLINE IN SODIUM CHLORIDE 0.45%

+ HOSPIRA 100MG/100ML N88147 002 May 03, 1983

+ 200MG/100ML N88147 003 May 03, 1983

SOLUTION; ORAL

AMINOPHYLLINEAA + ROXANE 105MG/5ML N88126 001 Aug 19, 1983

PRESCRIPTION DRUG PRODUCT LIST

3 - 24 (of 370)

AMINOPHYLLINE

SOLUTION; ORAL

AMINOPHYLLINE DYE FREE

<u>AA</u>	+	ACTAVIS MID ATLANTIC	<u>105MG/5ML</u>	<u>N87727</u>	<u>001</u>	Apr 16, 1982
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SUPPOSITORY; RECTAL

TRUPHYLLINE

		G AND W LABS	250MG	N85498	001	Mar 23, 1983
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	+		500MG	N85498	002	Jan 03, 1983
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TABLET; ORAL

AMINOPHYLLINE

	+	WEST WARD	100MG	N84540	001	
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	+		200MG	N85003	001	
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AMINOSALICYLATE SODIUM

TABLET; ORAL

SODIUM P.A.S.

	+	LANNETT	500MG	N80138	002	
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AMINOSALICYLIC ACID

GRANULE, DELAYED RELEASE; ORAL

PASER

	+	JACOBUS	4GM/PACKET	N74346	001	Jun 30, 1994
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AMIODARONE HYDROCHLORIDE

INJECTABLE; INJECTION

AMIODARONE HYDROCHLORIDE

<u>AP</u>	+	ABRAXIS PHARM	<u>50MG/ML</u>	<u>N75761</u>	<u>001</u>	Oct 15, 2002
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<u>AP</u>	+	AKORN	<u>50MG/ML</u>	<u>N76232</u>	<u>001</u>	Jul 05, 2006
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<u>AP</u>	+	APOTEX INC	<u>50MG/ML</u>	<u>N76394</u>	<u>001</u>	Apr 25, 2003
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<u>AP</u>	+	BEDFORD	<u>50MG/ML</u>	<u>N76018</u>	<u>001</u>	Oct 15, 2002
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<u>AP</u>	+	BEDFORD LABS	<u>50MG/ML</u>	<u>N76299</u>	<u>001</u>	Oct 24, 2002
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<u>AP</u>	+	BEN VENUE	<u>50MG/ML</u>	<u>N76088</u>	<u>001</u>	Oct 15, 2002
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<u>AP</u>	+	BIONICHE (CANADA)	<u>50MG/ML</u>	<u>N76217</u>	<u>001</u>	Oct 15, 2002
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<u>AP</u>	+	GLAND PHARMA LTD	<u>50MG/ML</u>	<u>N77161</u>	<u>001</u>	Apr 20, 2005
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<u>AP</u>	+	HOSPIRA	<u>50MG/ML</u>	<u>N75955</u>	<u>001</u>	Oct 18, 2002
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<u>AP</u>	+	INTL MEDICATION SYS	<u>50MG/ML</u>	<u>N21594</u>	<u>001</u>	Feb 04, 2004
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<u>AP</u>	+	MAYNE PHARMA USA	<u>50MG/ML</u>	<u>N76108</u>	<u>001</u>	Oct 15, 2002
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<u>AP</u>	+	SICOR PHARMS	<u>50MG/ML</u>	<u>N76163</u>	<u>001</u>	Sep 05, 2003
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CORDARONE

<u>AP</u>	+	WYETH PHARMS INC	<u>50MG/ML</u>	<u>N20377</u>	<u>001</u>	Aug 03, 1995
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TABLET; ORAL

AMIODARONE HYDROCHLORIDE

<u>AB</u>		ALPHAPHARM	<u>200MG</u>	<u>N75188</u>	<u>001</u>	Feb 24, 1999
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<u>AB</u>		AUROSAL PHARMS	<u>200MG</u>	<u>N77069</u>	<u>001</u>	Apr 08, 2005
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<u>AB</u>			<u>400MG</u>	<u>N77069</u>	<u>002</u>	Apr 08, 2005
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<u>AB</u>		BARR	<u>200MG</u>	<u>N75389</u>	<u>001</u>	Jan 25, 2001
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<u>AB</u>		SANDOZ	<u>200MG</u>	<u>N75315</u>	<u>001</u>	Dec 23, 1998
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<u>AB</u>			<u>400MG</u>	<u>N75315</u>	<u>002</u>	Jun 30, 2000
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<u>AB</u>		TARO	<u>100MG</u>	<u>N75424</u>	<u>002</u>	Dec 18, 2002
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<u>AB</u>			<u>200MG</u>	<u>N75424</u>	<u>001</u>	Mar 30, 2001
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<u>AB</u>			<u>400MG</u>	<u>N76362</u>	<u>001</u>	Nov 29, 2002
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			<u>300MG</u>	<u>N76362</u>	<u>002</u>	Dec 02, 2003
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<u>AB</u>		TEVA	<u>200MG</u>	<u>N74895</u>	<u>001</u>	Apr 16, 1999
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<u>AB</u>		TEVA PHARMS	<u>200MG</u>	<u>N74739</u>	<u>001</u>	Nov 30, 1998
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CORDARONE

<u>AB</u>	+	WYETH PHARMS INC	<u>200MG</u>	<u>N18972</u>	<u>001</u>	Dec 24, 1985
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PACERONE

<u>AB</u>		UPSHER SMITH	<u>100MG</u>	<u>N75135</u>	<u>002</u>	Apr 12, 2005
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<u>AB</u>			<u>200MG</u>	<u>N75135</u>	<u>001</u>	Apr 30, 1998
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27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 25 (of 370)

AMITRIPTYLINE HYDROCHLORIDE

TABLET; ORAL

AMITRIPTYLINE HYDROCHLORIDE

<u>AB</u>	MUTUAL PHARM	<u>10MG</u>	<u>N89398</u>	<u>001</u>	Jul 14, 1987
<u>AB</u>		<u>25MG</u>	<u>N89399</u>	<u>001</u>	Jul 14, 1987
<u>AB</u>		<u>50MG</u>	<u>N89400</u>	<u>001</u>	Jul 14, 1987
<u>AB</u>		<u>75MG</u>	<u>N89401</u>	<u>001</u>	Jul 14, 1987
<u>AB</u>		<u>100MG</u>	<u>N89402</u>	<u>001</u>	Jul 14, 1987
<u>AB</u>		<u>150MG</u>	<u>N89403</u>	<u>001</u>	Jul 14, 1987
<u>AB</u>	MYLAN	<u>10MG</u>	<u>N86157</u>	<u>001</u>	
<u>AB</u>		<u>25MG</u>	<u>N86010</u>	<u>001</u>	
<u>AB</u>		<u>50MG</u>	<u>N86009</u>	<u>001</u>	
<u>AB</u>		<u>75MG</u>	<u>N86011</u>	<u>001</u>	
<u>AB</u>		<u>100MG</u>	<u>N86158</u>	<u>001</u>	
<u>AB</u>		<u>150MG</u>	<u>N86153</u>	<u>001</u>	
<u>AB</u>	PLIVA	<u>10MG</u>	<u>N88883</u>	<u>001</u>	Sep 26, 1984
<u>AB</u>		<u>25MG</u>	<u>N88884</u>	<u>001</u>	Sep 26, 1984
<u>AB</u>		<u>50MG</u>	<u>N88885</u>	<u>001</u>	Sep 26, 1984
<u>AB</u>		<u>75MG</u>	<u>N88886</u>	<u>001</u>	Sep 26, 1984
<u>AB</u>		<u>100MG</u>	<u>N88887</u>	<u>001</u>	Sep 26, 1984
<u>AB</u>		<u>150MG</u>	<u>N88888</u>	<u>001</u>	Sep 26, 1984
<u>AB</u>	SANDOZ	<u>10MG</u>	<u>N85969</u>	<u>001</u>	
<u>AB</u>	+	<u>25MG</u>	<u>N85966</u>	<u>001</u>	
<u>AB</u>		<u>50MG</u>	<u>N85968</u>	<u>001</u>	
<u>AB</u>		<u>75MG</u>	<u>N85971</u>	<u>001</u>	
<u>AB</u>		<u>100MG</u>	<u>N85967</u>	<u>001</u>	
<u>AB</u>		<u>150MG</u>	<u>N85970</u>	<u>001</u>	
<u>AB</u>	TEVA	<u>10MG</u>	<u>N84910</u>	<u>003</u>	
<u>AB</u>		<u>25MG</u>	<u>N85031</u>	<u>001</u>	
<u>AB</u>		<u>100MG</u>	<u>N85836</u>	<u>001</u>	
<u>AB</u>	VINTAGE PHARMS	<u>10MG</u>	<u>N40218</u>	<u>001</u>	Sep 11, 1997
<u>AB</u>		<u>25MG</u>	<u>N40218</u>	<u>002</u>	Sep 11, 1997
<u>AB</u>		<u>50MG</u>	<u>N40218</u>	<u>003</u>	Sep 11, 1997
<u>AB</u>		<u>75MG</u>	<u>N40218</u>	<u>004</u>	Sep 11, 1997
<u>AB</u>		<u>100MG</u>	<u>N40218</u>	<u>005</u>	Sep 11, 1997
<u>AB</u>		<u>150MG</u>	<u>N40218</u>	<u>006</u>	Sep 11, 1997

AMITRIPTYLINE HYDROCHLORIDE; CHLORDIAZEPOXIDE

TABLET; ORAL

CHLORDIAZEPOXIDE AND AMITRIPTYLINE HYDROCHLORIDE

<u>AB</u>	MYLAN	<u>EQ 12.5MG BASE;5MG</u>	<u>N71296</u>	<u>001</u>	Dec 10, 1986
<u>AB</u>		<u>EQ 25MG BASE;10MG</u>	<u>N71297</u>	<u>001</u>	Dec 10, 1986
<u>AB</u>	PAR PHARM	<u>EQ 12.5MG BASE;5MG</u>	<u>N72277</u>	<u>001</u>	May 09, 1988
<u>AB</u>		<u>EQ 25MG BASE;10MG</u>	<u>N72278</u>	<u>001</u>	May 09, 1988
<u>AB</u>	WATSON LABS	<u>EQ 12.5MG BASE;5MG</u>	<u>N72052</u>	<u>001</u>	Dec 16, 1988
<u>AB</u>		<u>EQ 25MG BASE;10MG</u>	<u>N72053</u>	<u>001</u>	Dec 16, 1988
	<u>LIMBITROL</u>				
<u>AB</u>	VALEANT PHARM INTL	<u>EQ 12.5MG BASE;5MG</u>	<u>N16949</u>	<u>001</u>	
	<u>LIMBITROL DS</u>				
<u>AB</u>	+	<u>EQ 25MG BASE;10MG</u>	<u>N16949</u>	<u>002</u>	

AMITRIPTYLINE HYDROCHLORIDE; PERPHENAZINE

TABLET; ORAL

PERPHENAZINE AND AMITRIPTYLINE HYDROCHLORIDE

	MYLAN	10MG;2MG	N70336	001	Nov 10, 1988
		10MG;4MG	N71442	001	Nov 10, 1988
		25MG;2MG	N70337	001	Nov 10, 1988
	+	25MG;4MG	N70338	001	Nov 10, 1988
	+	50MG;4MG	N71443	001	Nov 10, 1988

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 26 (of 370)

AMLEXANOX

PASTE; DENTAL					
APHTHASOL					
+	ULURU	5%	N20511	001	Dec 17, 1996
PATCH; TOPICAL					
AMLEXANOX					
+	ULURU	2MG	N21727	001	Sep 29, 2004

AMLODIPINE BESYLATE

TABLET; ORAL					
<u>AMLODIPINE BESYLATE</u>					
<u>AB</u>	MYLAN	<u>EQ 2.5MG BASE</u>	<u>N76418</u>	<u>001</u>	Oct 03, 2005
<u>AB</u>		<u>EQ 5MG BASE</u>	<u>N76418</u>	<u>002</u>	Oct 03, 2005
<u>AB</u>		<u>EQ 10MG BASE</u>	<u>N76418</u>	<u>003</u>	Oct 03, 2005
<u>NORVASC</u>					
<u>AB</u>	PFIZER	<u>EQ 2.5MG BASE</u>	<u>N19787</u>	<u>001</u>	Jul 31, 1992
<u>AB</u>		<u>EQ 5MG BASE</u>	<u>N19787</u>	<u>002</u>	Jul 31, 1992
<u>AB</u>	+	<u>EQ 10MG BASE</u>	<u>N19787</u>	<u>003</u>	Jul 31, 1992

AMLODIPINE BESYLATE; ATORVASTATIN CALCIUM

TABLET; ORAL					
CADUET					
PFIZER					
		EQ 2.5MG BASE;EQ 10MG BASE	N21540	009	Jul 29, 2004
		EQ 2.5MG BASE;EQ 20MG BASE	N21540	010	Jul 29, 2004
		EQ 2.5MG BASE;EQ 40MG BASE	N21540	011	Jul 29, 2004
		EQ 5MG BASE;EQ 10MG BASE	N21540	001	Jan 30, 2004
		EQ 5MG BASE;EQ 20MG BASE	N21540	002	Jan 30, 2004
		EQ 5MG BASE;EQ 40MG BASE	N21540	003	Jan 30, 2004
		EQ 5MG BASE;EQ 80MG BASE	N21540	004	Jan 30, 2004
		EQ 10MG BASE;EQ 10MG BASE	N21540	005	Jan 30, 2004
		EQ 10MG BASE;EQ 20MG BASE	N21540	006	Jan 30, 2004
		EQ 10MG BASE;EQ 40MG BASE	N21540	007	Jan 30, 2004
+		EQ 10MG BASE;EQ 80MG BASE	N21540	008	Jan 30, 2004

AMLODIPINE BESYLATE; BENAZEPRIL HYDROCHLORIDE

CAPSULE; ORAL					
LOTREL					
NOVARTIS					
		EQ 2.5MG BASE;10MG	N20364	002	Mar 03, 1995
		EQ 5MG BASE;10MG	N20364	003	Mar 03, 1995
		EQ 5MG BASE;20MG	N20364	004	Mar 03, 1995
		EQ 5MG BASE;40MG	N20364	007	Apr 11, 2006
		EQ 10MG BASE;20MG	N20364	005	Jun 20, 2002
+		EQ 10MG BASE;40MG	N20364	006	Apr 11, 2006

AMMONIUM CHLORIDE

INJECTABLE; INJECTION					
AMMONIUM CHLORIDE IN PLASTIC CONTAINER					
+	HOSPIRA	5MEQ/ML	N88366	001	Jun 13, 1984

AMMONIUM LACTATE

CREAM; TOPICAL					
<u>AMMONIUM LACTATE</u>					
<u>AB</u>	PADDOCK	<u>EQ 12% BASE</u>	<u>N76829</u>	<u>001</u>	Feb 07, 2006
<u>AB</u>	PERRIGO NEW YORK	<u>EQ 12% BASE</u>	<u>N75774</u>	<u>001</u>	May 01, 2002
<u>AB</u>	TARO	<u>EQ 12% BASE</u>	<u>N75883</u>	<u>001</u>	Apr 10, 2003
<u>LAC-HYDRIN</u>					
<u>AB</u>	+	<u>EQ 12% BASE</u>	<u>N20508</u>	<u>001</u>	Aug 29, 1996

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 27 (of 370)

AMMONIUM LACTATE

LOTION; TOPICAL

AMMONIUM LACTATE

<u>AB</u>	PADDOCK	<u>EQ 12% BASE</u>	<u>N75575</u>	<u>001</u>	Jun 11, 2002
<u>AB</u>	PERRIGO NEW YORK	<u>EQ 12% BASE</u>	<u>N75570</u>	<u>001</u>	Jun 23, 2004
<u>AB</u>	TARO	<u>EQ 12% BASE</u>	<u>N76216</u>	<u>001</u>	May 28, 2004
<u>LAC-HYDRIN</u>					
<u>AB</u>	+ WESTWOOD SQUIBB	<u>EQ 12% BASE</u>	<u>N19155</u>	<u>001</u>	Apr 24, 1985

AMOXAPINE

TABLET; ORAL

AMOXAPINE

<u>AB</u>	SANDOZ	<u>25MG</u>	<u>N72943</u>	<u>001</u>	Jun 28, 1991
<u>AB</u>		<u>50MG</u>	<u>N72944</u>	<u>001</u>	Jun 28, 1991
<u>AB</u>		<u>100MG</u>	<u>N72878</u>	<u>001</u>	Jun 28, 1991
<u>AB</u>		<u>150MG</u>	<u>N72879</u>	<u>001</u>	Jun 28, 1991
<u>AB</u>	WATSON LABS	<u>25MG</u>	<u>N72418</u>	<u>001</u>	May 11, 1989
<u>AB</u>		<u>25MG</u>	<u>N72688</u>	<u>001</u>	Aug 28, 1992
<u>AB</u>		<u>50MG</u>	<u>N72419</u>	<u>001</u>	May 11, 1989
<u>AB</u>		<u>50MG</u>	<u>N72689</u>	<u>001</u>	Aug 28, 1992
<u>AB</u>		<u>100MG</u>	<u>N72420</u>	<u>001</u>	May 11, 1989
<u>AB</u>		<u>100MG</u>	<u>N72690</u>	<u>001</u>	Aug 28, 1992
<u>AB</u>	+	<u>150MG</u>	<u>N72421</u>	<u>001</u>	May 11, 1989
<u>AB</u>		<u>150MG</u>	<u>N72691</u>	<u>001</u>	Aug 28, 1992

AMOXICILLIN

CAPSULE; ORAL

AMOXICILLIN

<u>AB</u>	AM ANTIBIOTICS	<u>250MG</u>	<u>N62058</u>	<u>001</u>	
<u>AB</u>		<u>500MG</u>	<u>N62058</u>	<u>002</u>	
<u>AB</u>	AUROBINDO	<u>250MG</u>	<u>N65271</u>	<u>001</u>	Nov 09, 2005
<u>AB</u>		<u>500MG</u>	<u>N65271</u>	<u>002</u>	Nov 09, 2005
<u>AB</u>	BIOCHEMIE	<u>250MG</u>	<u>N64076</u>	<u>001</u>	Sep 30, 1994
<u>AB</u>		<u>500MG</u>	<u>N64076</u>	<u>002</u>	Sep 30, 1994
<u>AB</u>	CLONMEL	<u>250MG</u>	<u>N62884</u>	<u>001</u>	Feb 25, 1988
<u>AB</u>		<u>500MG</u>	<u>N62881</u>	<u>001</u>	Feb 25, 1988
<u>AB</u>	RANBAXY	<u>250MG</u>	<u>N65016</u>	<u>001</u>	Apr 08, 1999
<u>AB</u>		<u>500MG</u>	<u>N65016</u>	<u>002</u>	Apr 08, 1999
<u>AB</u>	TEVA	<u>250MG</u>	<u>N61926</u>	<u>001</u>	
<u>AB</u>		<u>250MG</u>	<u>N62853</u>	<u>001</u>	Dec 22, 1987
<u>AB</u>		<u>500MG</u>	<u>N61926</u>	<u>003</u>	
<u>AB</u>		<u>500MG</u>	<u>N62854</u>	<u>001</u>	Dec 22, 1987
<u>AMOXIL</u>					
<u>AB</u>	GLAXOSMITHKLINE	<u>250MG</u>	<u>N62216</u>	<u>001</u>	
<u>AB</u>	+	<u>500MG</u>	<u>N62216</u>	<u>004</u>	
<u>TRIMOX</u>					
<u>AB</u>	APOTHECON	<u>250MG</u>	<u>N61885</u>	<u>001</u>	
<u>AB</u>		<u>500MG</u>	<u>N61885</u>	<u>002</u>	

FOR SUSPENSION; ORAL

AMOXICILLIN

<u>AB</u>	AM ANTIBIOTICS	<u>125MG/5ML</u>	<u>N62059</u>	<u>001</u>	
<u>AB</u>		<u>250MG/5ML</u>	<u>N62059</u>	<u>002</u>	
<u>AB</u>	AUROBINDO	<u>200MG/5ML</u>	<u>N65334</u>	<u>001</u>	Dec 28, 2006
<u>AB</u>		<u>400MG/5ML</u>	<u>N65334</u>	<u>002</u>	Dec 28, 2006
<u>AB</u>	CLONMEL	<u>125MG/5ML</u>	<u>N62927</u>	<u>001</u>	Nov 25, 1988
<u>AB</u>		<u>250MG/5ML</u>	<u>N62927</u>	<u>002</u>	Nov 25, 1988
<u>AB</u>	HIKMA	<u>125MG/5ML</u>	<u>N65322</u>	<u>002</u>	Jun 19, 2006
<u>AB</u>		<u>200MG/5ML</u>	<u>N65325</u>	<u>002</u>	Jun 19, 2006
<u>AB</u>		<u>250MG/5ML</u>	<u>N65322</u>	<u>001</u>	Jun 19, 2006
<u>AB</u>		<u>400MG/5ML</u>	<u>N65325</u>	<u>001</u>	Jun 19, 2006

PRESCRIPTION DRUG PRODUCT LIST

3 - 28 (of 370)

AMOXICILLIN

FOR SUSPENSION; ORAL

<u>AMOXICILLIN</u>				
<u>AB</u>	RANBAXY	<u>200MG/5ML</u>	<u>N65113</u>	<u>001</u> Nov 29, 2002
<u>AB</u>		<u>400MG/5ML</u>	<u>N65113</u>	<u>002</u> Nov 29, 2002
<u>AB</u>	TEVA	<u>125MG/5ML</u>	<u>N61931</u>	<u>001</u>
<u>AB</u>		<u>125MG/5ML</u>	<u>N62946</u>	<u>001</u> Nov 01, 1988
<u>AB</u>		<u>200MG/5ML</u>	<u>N65119</u>	<u>001</u> Dec 04, 2002
<u>AB</u>		<u>250MG/5ML</u>	<u>N61931</u>	<u>002</u>
<u>AB</u>		<u>250MG/5ML</u>	<u>N63001</u>	<u>001</u> Jan 06, 1989
<u>AB</u>		<u>400MG/5ML</u>	<u>N65119</u>	<u>002</u> Dec 04, 2002
<u>AMOXICILLIN PEDIATRIC</u>				
<u>AB</u>	TEVA	<u>50MG/ML</u>	<u>N61931</u>	<u>003</u> Dec 01, 1982
<u>AMOXIL</u>				
<u>AB</u>	GLAXOSMITHKLINE	<u>50MG/ML</u>	<u>N62226</u>	<u>005</u>
<u>AB</u>		<u>125MG/5ML</u>	<u>N62226</u>	<u>001</u>
<u>AB</u>		<u>200MG/5ML</u>	<u>N50760</u>	<u>001</u> Apr 15, 1999
<u>AB</u>	+	<u>250MG/5ML</u>	<u>N62226</u>	<u>002</u>
<u>AB</u>	+	<u>400MG/5ML</u>	<u>N50760</u>	<u>002</u> Apr 15, 1999
<u>LAROTID</u>				
<u>AB</u>	GLAXOSMITHKLINE	<u>125MG/5ML</u>	<u>N62226</u>	<u>003</u>
<u>AB</u>		<u>250MG/5ML</u>	<u>N62226</u>	<u>004</u>
<u>TRIMOX</u>				
<u>AB</u>	APOTHECON	<u>50MG/ML</u>	<u>N61886</u>	<u>001</u>
<u>AB</u>		<u>125MG/5ML</u>	<u>N61886</u>	<u>002</u>
<u>AB</u>		<u>125MG/5ML</u>	<u>N62885</u>	<u>001</u> Mar 08, 1988
<u>AB</u>		<u>250MG/5ML</u>	<u>N61886</u>	<u>003</u>
<u>AB</u>		<u>250MG/5ML</u>	<u>N62885</u>	<u>002</u> Mar 08, 1988

TABLET; ORAL

<u>AMOXICILLIN</u>				
<u>AB</u>	AUROBINDO	<u>500MG</u>	<u>N65256</u>	<u>001</u> Nov 09, 2005
<u>AB</u>		<u>875MG</u>	<u>N65256</u>	<u>002</u> Nov 09, 2005
<u>AB</u>	HIKMA	<u>875MG</u>	<u>N65255</u>	<u>001</u> Mar 29, 2006
<u>AB</u>	RANBAXY	<u>500MG</u>	<u>N65059</u>	<u>001</u> Nov 24, 2000
<u>AB</u>		<u>875MG</u>	<u>N65059</u>	<u>002</u> Nov 24, 2000
<u>AB</u>	SANDOZ	<u>500MG</u>	<u>N65228</u>	<u>001</u> Jul 13, 2005
<u>AB</u>		<u>875MG</u>	<u>N65228</u>	<u>002</u> Jul 13, 2005
<u>AB</u>	TEVA	<u>500MG</u>	<u>N65056</u>	<u>001</u> Sep 18, 2000
<u>AB</u>		<u>875MG</u>	<u>N65056</u>	<u>002</u> Sep 18, 2000
<u>AMOXIL</u>				
<u>AB</u>	GLAXOSMITHKLINE	<u>500MG</u>	<u>N50754</u>	<u>002</u> Jul 10, 1998
<u>AB</u>	+	<u>875MG</u>	<u>N50754</u>	<u>001</u> Jul 10, 1998

TABLET, CHEWABLE; ORAL

<u>AMOXICILLIN</u>				
<u>AB</u>	CLONMEL	<u>125MG</u>	<u>N64139</u>	<u>001</u> Jan 29, 1996
<u>AB</u>		<u>250MG</u>	<u>N64139</u>	<u>002</u> Jan 29, 1996
<u>AB</u>	RANBAXY	<u>125MG</u>	<u>N65021</u>	<u>001</u> Dec 23, 1999
<u>AB</u>		<u>200MG</u>	<u>N65060</u>	<u>001</u> Nov 29, 2000
<u>AB</u>		<u>250MG</u>	<u>N65021</u>	<u>002</u> Dec 23, 1999
<u>AB</u>		<u>400MG</u>	<u>N65060</u>	<u>002</u> Nov 29, 2000
<u>AB</u>	TEVA	<u>125MG</u>	<u>N64013</u>	<u>002</u> Sep 11, 1995
<u>AB</u>		<u>125MG</u>	<u>N64031</u>	<u>001</u> Dec 19, 1996
<u>AB</u>		<u>250MG</u>	<u>N64013</u>	<u>001</u> Dec 22, 1992
<u>AB</u>		<u>250MG</u>	<u>N64031</u>	<u>002</u> Dec 19, 1996
<u>AMOXIL</u>				
<u>AB</u>	GLAXOSMITHKLINE	<u>200MG</u>	<u>N50761</u>	<u>001</u> Apr 15, 1999
<u>AB</u>	+	<u>400MG</u>	<u>N50761</u>	<u>002</u> Apr 15, 1999

TABLET, FOR SUSPENSION; ORAL

DISPERMOX				
	RANBAXY	200MG	N65080	002 Aug 11, 2003

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 29 (of 370)

AMOXICILLINTABLET, FOR SUSPENSION; ORAL
DISPERMOX

+	RANBAXY	400MG	N65080	001	Aug 11, 2003
+		600MG	N65159	001	Dec 04, 2003

AMOXICILLIN; CLARITHROMYCIN; LANSOPRAZOLECAPSULE, TABLET, CAPSULE, DELAYED REL PELLETS; ORAL
PREVPAC

+	TAP PHARM	500MG, N/A, N/A; N/A, 500MG, N/A; N/A, N/A, 30M G	N50757	001	Dec 02, 1997
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AMOXICILLIN; CLAVULANATE POTASSIUM

FOR SUSPENSION; ORAL

AMOXICILLIN AND CLAVULANATE POTASSIUM

<u>AB</u>	HIKMA PHARMS	<u>200MG/5ML;EQ 28.5MG BASE/5ML</u>	<u>N65191</u>	<u>002</u>	Jan 25, 2005
<u>AB</u>		<u>400MG/5ML;EQ 57MG BASE/5ML</u>	<u>N65191</u>	<u>001</u>	Jan 25, 2005
<u>AB</u>	LEK PHARMS	<u>200MG/5ML;EQ 28.5MG BASE/5ML</u>	<u>N65098</u>	<u>001</u>	Dec 16, 2002
<u>AB</u>		<u>400MG/5ML;EQ 57MG BASE/5ML</u>	<u>N65098</u>	<u>002</u>	Dec 16, 2002
<u>AB</u>	RANBAXY	<u>200MG/5ML;EQ 28.5MG BASE/5ML</u>	<u>N65132</u>	<u>001</u>	Mar 19, 2003
<u>AB</u>		<u>400MG/5ML;EQ 57MG BASE/5ML</u>	<u>N65132</u>	<u>002</u>	Mar 19, 2003
<u>AB</u>	SANDOZ	<u>200MG/5ML;EQ 28.5MG BASE/5ML</u>	<u>N65066</u>	<u>001</u>	Jun 05, 2002
<u>AB</u>		<u>400MG/5ML;EQ 57MG BASE/5ML</u>	<u>N65066</u>	<u>002</u>	Jun 05, 2002
<u>AB</u>	TEVA	<u>200MG/5ML;EQ 28.5MG BASE/5ML</u>	<u>N65089</u>	<u>001</u>	May 25, 2004
<u>AB</u>		<u>400MG/5ML;EQ 57MG BASE/5ML</u>	<u>N65089</u>	<u>002</u>	May 25, 2004
<u>AB</u>		<u>600MG/5ML;EQ 42.9MG BASE/5ML</u>	<u>N65162</u>	<u>001</u>	Mar 12, 2004
	AUGMENTIN '125'				
	GLAXOSMITHKLINE	125MG/5ML;EQ 31.25MG BASE/5ML	N50575	001	Aug 06, 1984
	<u>AUGMENTIN '200'</u>				
<u>AB</u>	GLAXOSMITHKLINE	<u>200MG/5ML;EQ 28.5MG BASE/5ML</u>	<u>N50725</u>	<u>001</u>	May 31, 1996
	AUGMENTIN '250'				
	+ GLAXOSMITHKLINE	250MG/5ML;EQ 62.5MG BASE/5ML	N50575	002	Aug 06, 1984
	<u>AUGMENTIN '400'</u>				
<u>AB</u>	+ GLAXOSMITHKLINE	<u>400MG/5ML;EQ 57MG BASE/5ML</u>	<u>N50725</u>	<u>002</u>	May 31, 1996
	<u>AUGMENTIN ES-600</u>				
<u>AB</u>	+ GLAXOSMITHKLINE	<u>600MG/5ML;EQ 42.9MG BASE/5ML</u>	<u>N50755</u>	<u>001</u>	Jun 22, 2001

TABLET; ORAL

AMOXICILLIN AND CLAVULANATE POTASSIUM

<u>AB</u>	LEK PHARMS	<u>500MG;EQ 125MG BASE</u>	<u>N65117</u>	<u>001</u>	Nov 27, 2002
<u>AB</u>		<u>875MG;EQ 125MG BASE</u>	<u>N65093</u>	<u>001</u>	Nov 21, 2002
<u>AB</u>	RANBAXY	<u>500MG;EQ 125MG BASE</u>	<u>N65109</u>	<u>001</u>	Nov 04, 2002
<u>AB</u>		<u>875MG;EQ 125MG BASE</u>	<u>N65102</u>	<u>001</u>	Sep 17, 2002
<u>AB</u>	SANDOZ	<u>250MG;EQ 125MG BASE</u>	<u>N65189</u>	<u>001</u>	Aug 23, 2005
<u>AB</u>		<u>500MG;EQ 125MG BASE</u>	<u>N65064</u>	<u>001</u>	Mar 15, 2002
<u>AB</u>		<u>875MG;EQ 125MG BASE</u>	<u>N65063</u>	<u>001</u>	Mar 14, 2002
<u>AB</u>	TEVA	<u>500MG;EQ 125MG BASE</u>	<u>N65101</u>	<u>001</u>	Oct 30, 2002
<u>AB</u>		<u>875MG;EQ 125MG BASE</u>	<u>N65096</u>	<u>001</u>	Oct 29, 2002
	<u>AUGMENTIN '250'</u>				
<u>AB</u>	GLAXOSMITHKLINE	<u>250MG;EQ 125MG BASE</u>	<u>N50564</u>	<u>001</u>	Aug 06, 1984
	<u>AUGMENTIN '500'</u>				
<u>AB</u>	GLAXOSMITHKLINE	<u>500MG;EQ 125MG BASE</u>	<u>N50564</u>	<u>002</u>	Aug 06, 1984
	<u>AUGMENTIN '875'</u>				
<u>AB</u>	+ GLAXOSMITHKLINE	<u>875MG;EQ 125MG BASE</u>	<u>N50720</u>	<u>001</u>	Feb 13, 1996

TABLET, CHEWABLE; ORAL

AMOXICILLIN AND CLAVULANATE POTASSIUM

<u>AB</u>	RANBAXY	<u>200MG;EQ 28.5MG BASE</u>	<u>N65161</u>	<u>001</u>	Dec 03, 2003
<u>AB</u>		<u>400MG;EQ 57MG BASE</u>	<u>N65161</u>	<u>002</u>	Dec 03, 2003
<u>AB</u>	SANDOZ	<u>200MG;EQ 28.5MG BASE</u>	<u>N65065</u>	<u>001</u>	Apr 18, 2002
<u>AB</u>		<u>400MG;EQ 57MG BASE</u>	<u>N65065</u>	<u>002</u>	Apr 18, 2002
<u>AB</u>	TEVA	<u>200MG;EQ 28.5MG BASE</u>	<u>N65205</u>	<u>001</u>	Feb 09, 2005

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 30 (of 370)

AMOXICILLIN; CLAVULANATE POTASSIUM

TABLET, CHEWABLE; ORAL

AMOXICILLIN AND CLAVULANATE POTASSIUM

<u>AB</u>	TEVA	<u>400MG;EQ 57MG BASE</u>	<u>N65205</u>	<u>002</u>	Feb 09, 2005
	AUGMENTIN '125'				
	GLAXOSMITHKLINE	125MG;EQ 31.25MG BASE	N50597	001	Jul 22, 1985
	<u>AUGMENTIN '200'</u>				
<u>AB</u>	GLAXOSMITHKLINE	<u>200MG;EQ 28.5MG BASE</u>	<u>N50726</u>	<u>001</u>	May 31, 1996
	AUGMENTIN '250'				
	+ GLAXOSMITHKLINE	250MG;EQ 62.5MG BASE	N50597	002	Jul 22, 1985
	<u>AUGMENTIN '400'</u>				
<u>AB</u>	+ GLAXOSMITHKLINE	<u>400MG;EQ 57MG BASE</u>	<u>N50726</u>	<u>002</u>	May 31, 1996
TABLET, EXTENDED RELEASE; ORAL					
	AUGMENTIN XR				
	+ GLAXOSMITHKLINE	1GM;EQ 62.5MG BASE	N50785	001	Sep 25, 2002

AMPHETAMINE ASPARTATE; AMPHETAMINE SULFATE; DEXTROAMPHETAMINE SACCHARATE; DEXTROAMPHETAMINE SULFATE

CAPSULE, EXTENDED RELEASE; ORAL

ADDERALL XR 10

	SHIRE	2.5MG;2.5MG;2.5MG;2.5MG	N21303	001	Oct 11, 2001
	ADDERALL XR 15				
	SHIRE	3.75MG;3.75MG;3.75MG;3.75MG	N21303	006	May 22, 2002
	ADDERALL XR 20				
	SHIRE	5MG;5MG;5MG;5MG	N21303	002	Oct 11, 2001
	ADDERALL XR 25				
	SHIRE	6.25MG;6.25MG;6.25MG;6.25MG	N21303	004	May 22, 2002
	ADDERALL XR 30				
	+ SHIRE	7.5MG;7.5MG;7.5MG;7.5MG	N21303	003	Oct 11, 2001
	ADDERALL XR 5				
	SHIRE	1.25MG;1.25MG;1.25MG;1.25MG	N21303	005	May 22, 2002

TABLET; ORAL

ADDERALL 10

<u>AB</u>	SHIRE	<u>2.5MG;2.5MG;2.5MG;2.5MG</u>	<u>N11522</u>	<u>007</u>	Feb 13, 1996
	<u>ADDERALL 12.5</u>				
<u>AB</u>	SHIRE	<u>3.125MG;3.125MG;3.125MG;3.125MG</u>	<u>N11522</u>	<u>012</u>	Aug 31, 2000
	<u>ADDERALL 15</u>				
<u>AB</u>	SHIRE	<u>3.75MG;3.75MG;3.75MG;3.75MG</u>	<u>N11522</u>	<u>013</u>	Aug 31, 2000
	<u>ADDERALL 20</u>				
<u>AB</u>	SHIRE	<u>5MG;5MG;5MG;5MG</u>	<u>N11522</u>	<u>008</u>	Feb 13, 1996
	<u>ADDERALL 30</u>				
<u>AB</u>	+ SHIRE	<u>7.5MG;7.5MG;7.5MG;7.5MG</u>	<u>N11522</u>	<u>010</u>	May 12, 1997
	<u>ADDERALL 5</u>				
<u>AB</u>	SHIRE	<u>1.25MG;1.25MG;1.25MG;1.25MG</u>	<u>N11522</u>	<u>009</u>	May 12, 1997
	<u>ADDERALL 7.5</u>				
<u>AB</u>	SHIRE	<u>1.875MG;1.875MG;1.875MG;1.875MG</u>	<u>N11522</u>	<u>011</u>	Aug 31, 2000
<u>DEXTROAMP SACCHARATE, AMP ASPARTATE, DEXTROAMP SULFATE AND AMP SULFATE</u>					
<u>AB</u>	BARR	<u>1.25MG;1.25MG;1.25MG;1.25MG</u>	<u>N40422</u>	<u>001</u>	Feb 11, 2002
<u>AB</u>		<u>1.875MG;1.875MG;1.875MG;1.875MG</u>	<u>N40422</u>	<u>005</u>	Mar 19, 2003
<u>AB</u>		<u>2.5MG;2.5MG;2.5MG;2.5MG</u>	<u>N40422</u>	<u>002</u>	Feb 11, 2002
<u>AB</u>		<u>3.125MG;3.125MG;3.125MG;3.125MG</u>	<u>N40422</u>	<u>006</u>	Mar 19, 2003
<u>AB</u>		<u>3.75MG;3.75MG;3.75MG;3.75MG</u>	<u>N40422</u>	<u>007</u>	Mar 19, 2003
<u>AB</u>		<u>5MG;5MG;5MG;5MG</u>	<u>N40422</u>	<u>003</u>	Feb 11, 2002
<u>AB</u>		<u>7.5MG;7.5MG;7.5MG;7.5MG</u>	<u>N40422</u>	<u>004</u>	Feb 11, 2002
<u>AB</u>	COREPHARMA	<u>1.25MG;1.25MG;1.25MG;1.25MG</u>	<u>N40444</u>	<u>001</u>	Jun 19, 2002
<u>AB</u>		<u>2.5MG;2.5MG;2.5MG;2.5MG</u>	<u>N40444</u>	<u>002</u>	Jun 19, 2002
<u>AB</u>		<u>5MG;5MG;5MG;5MG</u>	<u>N40444</u>	<u>003</u>	Jun 19, 2002
<u>AB</u>		<u>7.5MG;7.5MG;7.5MG;7.5MG</u>	<u>N40444</u>	<u>004</u>	Jun 19, 2002
<u>AB</u>	MALLINCKRODT	<u>1.25MG;1.25MG;1.25MG;1.25MG</u>	<u>N40440</u>	<u>001</u>	Oct 07, 2003
<u>AB</u>		<u>1.875MG;1.875MG;1.875MG;1.875MG</u>	<u>N40440</u>	<u>002</u>	Oct 07, 2003
<u>AB</u>		<u>2.5MG;2.5MG;2.5MG;2.5MG</u>	<u>N40440</u>	<u>003</u>	Oct 07, 2003

PRESCRIPTION DRUG PRODUCT LIST

3 - 31 (of 370)

AMPHETAMINE ASPARTATE; AMPHETAMINE SULFATE; DEXTROAMPHETAMINE SACCHARATE;
DEXTROAMPHETAMINE SULFATE

TABLET; ORAL

<u>DEXTROAMP SACCHARATE, AMP ASPARTATE, DEXTROAMP SULFATE AND AMP SULFATE</u>					
<u>AB</u>	MALLINCKRODT	<u>3.125MG;3.125MG;3.125MG;3.125MG</u>	<u>N40440</u>	<u>004</u>	Oct 07, 2003
<u>AB</u>		<u>3.75MG;3.75MG;3.75MG;3.75MG</u>	<u>N40440</u>	<u>005</u>	Oct 07, 2003
<u>AB</u>		<u>5MG;5MG;5MG;5MG</u>	<u>N40440</u>	<u>006</u>	Oct 07, 2003
<u>AB</u>		<u>7.5MG;7.5MG;7.5MG;7.5MG</u>	<u>N40440</u>	<u>007</u>	Oct 07, 2003
<u>AB</u>	MUTUAL PHARM	<u>1.25MG;1.25MG;1.25MG;1.25MG</u>	<u>N40480</u>	<u>001</u>	Sep 09, 2003
<u>AB</u>		<u>1.875MG;1.875MG;1.875MG;1.875MG</u>	<u>N40480</u>	<u>002</u>	Sep 09, 2003
<u>AB</u>		<u>2.5MG;2.5MG;2.5MG;2.5MG</u>	<u>N40480</u>	<u>003</u>	Sep 09, 2003
<u>AB</u>		<u>3.125MG;3.125MG;3.125MG;3.125MG</u>	<u>N40480</u>	<u>004</u>	Sep 09, 2003
<u>AB</u>		<u>3.75MG;3.75MG;3.75MG;3.75MG</u>	<u>N40480</u>	<u>005</u>	Sep 09, 2003
<u>AB</u>		<u>5MG;5MG;5MG;5MG</u>	<u>N40480</u>	<u>006</u>	Sep 09, 2003
<u>AB</u>		<u>7.5MG;7.5MG;7.5MG;7.5MG</u>	<u>N40480</u>	<u>007</u>	Sep 09, 2003
<u>AB</u>	SANDOZ	<u>1.25MG;1.25MG;1.25MG;1.25MG</u>	<u>N40470</u>	<u>001</u>	Sep 27, 2002
<u>AB</u>		<u>2.5MG;2.5MG;2.5MG;2.5MG</u>	<u>N40439</u>	<u>001</u>	Jun 14, 2002
<u>AB</u>		<u>5MG;5MG;5MG;5MG</u>	<u>N40439</u>	<u>002</u>	Jun 14, 2002
<u>AB</u>		<u>7.5MG;7.5MG;7.5MG;7.5MG</u>	<u>N40439</u>	<u>003</u>	Jun 14, 2002
<u>AB</u>	TEVA PHARMS	<u>1.25MG;1.25MG;1.25MG;1.25MG</u>	<u>N40472</u>	<u>001</u>	Sep 30, 2003
<u>AB</u>		<u>2.5MG;2.5MG;2.5MG;2.5MG</u>	<u>N40472</u>	<u>002</u>	Sep 30, 2003
<u>AB</u>		<u>5MG;5MG;5MG;5MG</u>	<u>N40472</u>	<u>003</u>	Sep 30, 2003
<u>AB</u>		<u>7.5MG;7.5MG;7.5MG;7.5MG</u>	<u>N40472</u>	<u>004</u>	Sep 30, 2003
<u>AB</u>	WATSON LABS	<u>1.25MG;1.25MG;1.25MG;1.25MG</u>	<u>N40456</u>	<u>001</u>	May 06, 2003
<u>AB</u>		<u>2.5MG;2.5MG;2.5MG;2.5MG</u>	<u>N40456</u>	<u>002</u>	May 06, 2003
<u>AB</u>		<u>5MG;5MG;5MG;5MG</u>	<u>N40456</u>	<u>003</u>	May 06, 2003
<u>AB</u>		<u>7.5MG;7.5MG;7.5MG;7.5MG</u>	<u>N40456</u>	<u>004</u>	May 06, 2003

AMPHOTERICIN B

CREAM; TOPICAL
FUNGIZONE

+	APOTHECON	3%	N50314	001	
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INJECTABLE; INJECTION

<u>AMPHOTERICIN B</u>					
<u>AP</u>	SICOR PHARMS	<u>50MG/VIAL</u>	<u>N64062</u>	<u>001</u>	Mar 31, 1995
<u>AP</u>	X GEN PHARMS	<u>50MG/VIAL</u>	<u>N63206</u>	<u>001</u>	Apr 29, 1992
<u>FUNGIZONE</u>					
<u>AP</u>	+ APOTHECON	<u>50MG/VIAL</u>	<u>N60517</u>	<u>001</u>	

INJECTABLE, LIPID COMPLEX; INJECTION

ABELCET					
+	ENZON	5MG/ML	N50724	001	Nov 20, 1995
AMPHOTEC					
+	THREE RIVERS PHARMS	50MG/VIAL	N50729	001	Nov 22, 1996
+		100MG/VIAL	N50729	002	Nov 22, 1996

INJECTABLE, LIPOSOMAL; INJECTION

AMBISOME					
+	ASTELLAS	50MG/VIAL	N50740	001	Aug 11, 1997

AMPICILLIN SODIUM

INJECTABLE; INJECTION

<u>AMPICILLIN SODIUM</u>					
<u>AP</u>	IBI	<u>EQ 250MG BASE/VIAL</u>	<u>N62719</u>	<u>001</u>	May 12, 1987
<u>AP</u>		<u>EQ 500MG BASE/VIAL</u>	<u>N62719</u>	<u>003</u>	May 12, 1987
<u>AP</u>		<u>EQ 1GM BASE/VIAL</u>	<u>N62719</u>	<u>002</u>	May 12, 1987
<u>AP</u>	INSTITUTO BIOCHEMICO	<u>EQ 125MG BASE/VIAL</u>	<u>N62797</u>	<u>001</u>	Jul 12, 1993
<u>AP</u>		<u>EQ 2GM BASE/VIAL</u>	<u>N62797</u>	<u>002</u>	Jul 12, 1993
<u>AP</u>	+ SANDOZ	<u>EQ 125MG BASE/VIAL</u>	<u>N61395</u>	<u>001</u>	
<u>AP</u>	+	<u>EQ 250MG BASE/VIAL</u>	<u>N61395</u>	<u>002</u>	
<u>AP</u>	+	<u>EQ 500MG BASE/VIAL</u>	<u>N61395</u>	<u>003</u>	

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 32 (of 370)

AMPICILLIN SODIUM

INJECTABLE; INJECTION

AMPICILLIN SODIUM

<u>AP</u>	+	SANDOZ	<u>EQ 1GM BASE/VIAL</u>	<u>N61395</u>	<u>004</u>	
<u>AP</u>			<u>EQ 1GM BASE/VIAL</u>	<u>N62738</u>	<u>001</u>	Feb 19, 1987
<u>AP</u>	+		<u>EQ 2GM BASE/VIAL</u>	<u>N61395</u>	<u>005</u>	
<u>AP</u>			<u>EQ 2GM BASE/VIAL</u>	<u>N62738</u>	<u>002</u>	Feb 19, 1987
<u>AP</u>	+		<u>EQ 10GM BASE/VIAL</u>	<u>N61395</u>	<u>006</u>	

AMPICILLIN SODIUM; SULBACTAM SODIUM

INJECTABLE; INJECTION

AMPICILLIN AND SULBACTAM

<u>AP</u>		BAXTER HLTHCARE	<u>EQ 1GM BASE/VIAL;EQ 500MG BASE/VIAL</u>	<u>N65074</u>	<u>001</u>	Mar 19, 2002
<u>AP</u>			<u>EQ 2GM BASE/VIAL;EQ 1GM BASE/VIAL</u>	<u>N65074</u>	<u>002</u>	Mar 19, 2002
<u>AP</u>			<u>EQ 10GM BASE/VIAL;EQ 5GM BASE/VIAL</u>	<u>N65076</u>	<u>001</u>	Mar 19, 2002
<u>AP</u>		HANFORD GC	<u>EQ 1GM BASE/VIAL;EQ 500MG BASE/VIAL</u>	<u>N65176</u>	<u>001</u>	Nov 30, 2005
<u>AP</u>			<u>EQ 2GM BASE/VIAL;EQ 1GM BASE/VIAL</u>	<u>N65176</u>	<u>002</u>	Nov 30, 2005
<u>AP</u>			<u>EQ 10GM BASE/VIAL;EQ 5GM BASE/VIAL</u>	<u>N65188</u>	<u>001</u>	Nov 25, 2005
<u>AP</u>		INSTITUTO BIOCHIMICO	<u>EQ 1GM BASE/VIAL;EQ 500MG BASE/VIAL</u>	<u>N65222</u>	<u>001</u>	Nov 29, 2005
<u>AP</u>			<u>EQ 2GM BASE/VIAL;EQ 1GM BASE/VIAL</u>	<u>N65222</u>	<u>002</u>	Nov 29, 2005
<u>AP</u>			<u>EQ 10GM BASE/VIAL;EQ 5GM BASE/VIAL</u>	<u>N65314</u>	<u>001</u>	Nov 27, 2006
<u>AP</u>		SANDOZ	<u>EQ 1GM BASE/VIAL;EQ 500MG BASE/VIAL</u>	<u>N65241</u>	<u>001</u>	Jul 25, 2006
<u>AP</u>			<u>EQ 1GM BASE/VIAL;EQ 500MG BASE/VIAL</u>	<u>N65310</u>	<u>001</u>	Jul 25, 2006
<u>AP</u>			<u>EQ 2GM BASE/VIAL;EQ 1GM BASE/VIAL</u>	<u>N65241</u>	<u>002</u>	Jul 25, 2006
<u>AP</u>			<u>EQ 2GM BASE/VIAL;EQ 1GM BASE/VIAL</u>	<u>N65310</u>	<u>002</u>	Jul 25, 2006
<u>AP</u>			<u>EQ 10GM BASE/VIAL;EQ 5GM BASE/VIAL</u>	<u>N65240</u>	<u>001</u>	Jul 25, 2006
<u>UNASYN</u>						
<u>AP</u>	+	PFIZER	<u>EQ 1GM BASE/VIAL;EQ 500MG BASE/VIAL</u>	<u>N50608</u>	<u>002</u>	Dec 31, 1986
<u>AP</u>	+		<u>EQ 2GM BASE/VIAL;EQ 1GM BASE/VIAL</u>	<u>N50608</u>	<u>001</u>	Dec 31, 1986
<u>AP</u>	+		<u>EQ 10GM BASE/VIAL;EQ 5GM BASE/VIAL</u>	<u>N50608</u>	<u>005</u>	Dec 10, 1993
			<u>EQ 1GM BASE/VIAL;EQ 500MG BASE/VIAL</u>	<u>N62901</u>	<u>001</u>	Nov 23, 1988

AMPICILLIN/AMPICILLIN TRIHYDRATE

CAPSULE; ORAL

AMPICILLIN TRIHYDRATE

		CLONMEL	EQ 250MG BASE	N62883	001	Feb 25, 1988
	+		EQ 500MG BASE	N62882	001	Feb 25, 1988

FOR SUSPENSION; ORAL

AMPICILLIN TRIHYDRATE

<u>AB</u>		CLONMEL	<u>EQ 125MG BASE/5ML</u>	<u>N62982</u>	<u>001</u>	Feb 10, 1989
<u>AB</u>			<u>EQ 250MG BASE/5ML</u>	<u>N62982</u>	<u>002</u>	Feb 10, 1989
<u>AB</u>		PUREPAC PHARM	<u>EQ 250MG BASE/5ML</u>	<u>N61980</u>	<u>002</u>	
<u>PRINCIPEN</u>						
<u>AB</u>		APOTHECON	<u>EQ 125MG BASE/5ML</u>	<u>N61394</u>	<u>002</u>	
<u>AB</u>	+		<u>EQ 250MG BASE/5ML</u>	<u>N61394</u>	<u>003</u>	

AMPRENAVIR

CAPSULE; ORAL

AGENERASE

	+	GLAXOSMITHKLINE	50MG	N21007	001	Apr 15, 1999
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SOLUTION; ORAL

AGENERASE

	+	GLAXOSMITHKLINE	15MG/ML	N21039	001	Apr 15, 1999
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ANAGRELIDE HYDROCHLORIDE

CAPSULE; ORAL

AGRYLIN

<u>AB</u>		SHIRE	<u>EQ 0.5MG BASE</u>	<u>N20333</u>	<u>001</u>	Mar 14, 1997
<u>AB</u>	+		<u>EQ 1MG BASE</u>	<u>N20333</u>	<u>002</u>	Mar 14, 1997

PRESCRIPTION DRUG PRODUCT LIST

3 - 33 (of 370)

ANAGRELIDE HYDROCHLORIDE

CAPSULE; ORAL

ANAGRELIDE HYDROCHLORIDE

<u>AB</u>	ALPHAPHARM	<u>EQ 0.5MG BASE</u>	<u>N77613</u>	<u>001</u>	Jun 27, 2006
<u>AB</u>		<u>EQ 1MG BASE</u>	<u>N77613</u>	<u>002</u>	Jun 27, 2006
<u>AB</u>	BARR	<u>EQ 0.5MG BASE</u>	<u>N76530</u>	<u>001</u>	Apr 18, 2005
<u>AB</u>		<u>EQ 1MG BASE</u>	<u>N76530</u>	<u>002</u>	Apr 18, 2005
<u>AB</u>	IMPAX LABS	<u>EQ 0.5MG BASE</u>	<u>N76910</u>	<u>001</u>	Apr 18, 2005
<u>AB</u>		<u>EQ 1MG BASE</u>	<u>N76910</u>	<u>002</u>	Apr 18, 2005
<u>AB</u>	IVAX PHARMS	<u>EQ 0.5MG BASE</u>	<u>N76468</u>	<u>001</u>	Apr 18, 2005
<u>AB</u>		<u>EQ 1MG BASE</u>	<u>N76468</u>	<u>002</u>	Apr 18, 2005
<u>AB</u>	MYLAN	<u>EQ 0.5MG BASE</u>	<u>N76811</u>	<u>001</u>	Apr 18, 2005
<u>AB</u>		<u>EQ 1MG BASE</u>	<u>N76811</u>	<u>002</u>	Apr 18, 2005
<u>AB</u>	ROXANE	<u>EQ 0.5MG BASE</u>	<u>N76489</u>	<u>001</u>	Apr 18, 2005
<u>AB</u>		<u>EQ 1MG BASE</u>	<u>N76489</u>	<u>002</u>	Apr 18, 2005
<u>AB</u>	SANDOZ	<u>EQ 0.5MG BASE</u>	<u>N76683</u>	<u>001</u>	Apr 18, 2005
<u>AB</u>		<u>EQ 1MG BASE</u>	<u>N76683</u>	<u>002</u>	Apr 18, 2005
<u>AB</u>	WATSON LABS	<u>EQ 0.5MG BASE</u>	<u>N76417</u>	<u>001</u>	Apr 18, 2005
<u>AB</u>		<u>EQ 1MG BASE</u>	<u>N76417</u>	<u>002</u>	Apr 18, 2005

ANASTROZOLE

TABLET; ORAL

ARIMIDEX

+	ASTRAZENECA	1MG	N20541	001	Dec 27, 1995
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ANIDULAFUNGIN

INJECTABLE; IV (INFUSION)

ERAXIS

+	VICURON	50MG/VIAL	N21632	001	Feb 17, 2006
+		100MG/VIAL	N21632	002	Nov 14, 2006

APOMORPHINE HYDROCHLORIDE

INJECTABLE; SUBCUTANEOUS

APOKYN

+	VERNALIS	30MG/3ML (10MG/ML)	N21264	002	Apr 20, 2004
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APRACLONIDINE HYDROCHLORIDE

SOLUTION/DROPS; OPHTHALMIC

+	ALCON	EQ 0.5% BASE	N20258	001	Jul 30, 1993
+		EQ 1% BASE	N19779	001	Dec 31, 1987

APREPITANT

CAPSULE; ORAL

EMEND

MERCK

		40MG	N21549	003	Jun 30, 2006
		80MG	N21549	001	Mar 26, 2003
+		125MG	N21549	002	Mar 26, 2003

APROTININ BOVINE

INJECTABLE; INJECTION

TRASYLOL

+	BAYER PHARMS	10,000KIU/ML	N20304	001	Dec 29, 1993
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ARFORMOTEROL TARTRATE

SOLUTION; INHALATION

BROVANA

+	SEPRACOR	EQ 0.015MG BASE/2ML	N21912	001	Oct 06, 2006
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27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 34 (of 370)

ARGATROBAN

INJECTABLE; INJECTION

ARGATROBAN

+	ENCYSIVE	100MG/ML	N20883	001	Jun 30, 2000
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ARGININE HYDROCHLORIDE

INJECTABLE; INJECTION

R-GENE 10

+	PHARMACIA AND UPJOHN	10GM/100ML	N16931	001	
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ARIPIPIRAZOLE

INJECTABLE; INTRAMUSCULAR

ABILIFY

+	OTSUKA	9.75MG/1.3ML (7.5MG/ML)	N21866	001	Sep 20, 2006
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SOLUTION; ORAL

ABILIFY

+	OTSUKA	1MG/ML	N21713	001	Dec 10, 2004
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TABLET; ORAL

ABILIFY

	OTSUKA	2MG	N21436	006	Nov 15, 2002
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+		5MG	N21436	005	Nov 15, 2002
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+		10MG	N21436	001	Nov 15, 2002
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+		15MG	N21436	002	Nov 15, 2002
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		20MG	N21436	003	Nov 15, 2002
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+		30MG	N21436	004	Nov 15, 2002
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TABLET, ORALLY DISINTEGRATING; ORAL

ABILIFY

	OTSUKA	10MG	N21729	002	Jun 07, 2006
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		15MG	N21729	003	Jun 07, 2006
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		20MG	N21729	004	Jun 07, 2006
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+		30MG	N21729	005	Jun 07, 2006
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ARSENIC TRIOXIDE

INJECTABLE; INJECTION

TRISENOX

+	CEPHALON	1MG/ML	N21248	001	Sep 25, 2000
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ARTICAINE HYDROCHLORIDE; EPINEPHRINE BITARTRATE

INJECTABLE; INJECTION

SEPTOCAINE

+	DEPROCO	4%;EQ 0.0085MG BASE/1.7ML(4%; EQ 0.005MG BASE/ML)	N22010	001	Mar 30, 2006
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+		4%; EQ 0.017MG BASE/1.7ML (4%; EQ 0.01MG BASE/ML)	N20971	001	Apr 03, 2000
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ASCORBIC ACID; BIOTIN; CHOLECALCIFEROL; CYANOCOBALAMIN; DEXPANTHENOL; FOLIC ACID; NIACINAMIDE; PYRIDOXINE; RIBOFLAVIN; THIAMINE; TOCOPHEROL ACETATE; VITAMIN A; VITAMIN K

INJECTABLE; IV (INFUSION)

+	SANDOZ	80MG/VIAL;0.02MG/VIAL;400 IU/VIAL;0.001MG/VIAL;5MG/VIAL;0.14MG/VIAL;17MG/VIAL;1MG/VIAL;1.4MG/VIAL;1.2MG/VIAL;7 IU/VIAL;2,300 IU/VIAL;0.2MG/VIAL	N21265	001	Feb 21, 2001
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INFUVITE PEDIATRIC (PHARMACY BULK PACKAGE)

+	SANDOZ	80MG/VIAL;0.02MG/VIAL;400 IU/VIAL;0.001MG/VIAL;5MG/VIAL;0.14MG/VIAL;17MG/VIAL;1MG/VIAL;1.4MG/VIAL;1.2MG/VIAL;7 IU/VIAL;2,300 IU/VIAL;0.2MG/VIAL	N21646	001	Jan 29, 2004
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27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 35 (of 370)

ASCORBIC ACID; BIOTIN; CYANOCOBALAMIN; DEXPANTHENOL; ERGOCALCIFEROL; FOLIC ACID; NIACINAMIDE; PHYTONADIONE; PYRIDOXINE HYDROCHLORIDE; RIBOFLAVIN PHOSPHATE SODIUM; THIAMINE HYDROCHLORIDE; VITAMIN A; VITAMIN E

FOR SOLUTION; IV (INFUSION)

M.V.I. PEDIATRIC

+ MAYNE PHARMA USA	80MG/VIAL;0.02MG/VIAL;0.001MG/VIAL;5MG/VIAL;0.01MG/VIAL;0.14MG/VIAL;17MG/VIAL;0.2MG/VIAL;1MG/VIAL;1.4MG/VIAL;EQ 1.2MG BASE/VIAL;0.7MG/VIAL;7MG/VIAL	N18920 001	Sep 21, 2000
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ASCORBIC ACID; BIOTIN; CYANOCOBALAMIN; DEXPANTHENOL; ERGOCALCIFEROL; FOLIC ACID; NIACINAMIDE; PYRIDOXINE HYDROCHLORIDE; RIBOFLAVIN PHOSPHATE SODIUM; THIAMINE HYDROCHLORIDE; VITAMIN A; VITAMIN E

INJECTABLE; INJECTION

M.V.I. -12 (WITHOUT VITAMIN K)

+ MAYNE PHARMA USA	20MG/ML;0.006MG/ML;0.05UGM/ML;1.5MG/ML;0.0005MG/ML;0.06MG/ML;4MG/ML;0.6MG/ML;0.36MG/ML;0.6MG/ML;0.1MG/ML;1MG/ML	N08809 006	Sep 09, 2004
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ASCORBIC ACID; BIOTIN; CYANOCOBALAMIN; DEXPANTHENOL; ERGOCALCIFEROL; FOLIC ACID; NIACINAMIDE; PYRIDOXINE HYDROCHLORIDE; RIBOFLAVIN PHOSPHATE SODIUM; THIAMINE HYDROCHLORIDE; VITAMIN A; VITAMIN E; VITAMIN K

INJECTABLE; IV (INFUSION)

M.V.I. ADULT

+ MAYNE PHARMA USA	200MG/VIAL;0.06MG/VIAL;0.005MG/VIAL;15MG/VIAL;0.005MG/VIAL;0.6MG/VIAL;40MG/VIAL;6MG/VIAL;3.6MG/VIAL;6MG/VIAL;1MG/VIAL;10MG/VIAL;0.15MG/VIAL	N21625 001	Jan 30, 2004
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M.V.I. ADULT (PHARMACY BULK PACKAGE)

+ MAYNE PHARMA USA	200MG/5ML;0.06MG/5ML;0.005MG/5ML;15MG/5ML;0.005MG/5ML;0.6MG/5ML;40MG/5ML;6MG/5ML;3.6MG/5ML;6MG/5ML;1MG/5ML;10MG/5ML;0.15MG/5ML	N21643 001	Feb 18, 2004
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ASCORBIC ACID; POLYETHYLENE GLYCOL 3350; POTASSIUM CHLORIDE; SODIUM ASCORBATE; SODIUM CHLORIDE; SODIUM SULFATE

FOR SOLUTION; ORAL

MOVIPREP

+ SALIX PHARMS	4.7GM;100GM;1.015GM;5.9GM;2.691GM;7.5GM	N21881 001	Aug 02, 2006
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ASPIRIN; BUTALBITAL; CAFFEINE

CAPSULE; ORAL

FIORINAL

<u>AB</u> + WATSON PHARMS	<u>325MG;50MG;40MG</u>	<u>N17534</u> <u>005</u>	Apr 16, 1986
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LANORINAL

<u>AB</u> LANNETT	<u>325MG;50MG;40MG</u>	<u>N86996</u> <u>002</u>	Oct 11, 1985
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TABLET; ORAL

ASPIRIN AND CAFFEINE W/ BUTALBITAL

<u>AB</u> ACTAVIS ELIZABETH	<u>325MG;50MG;40MG</u>	<u>N86710</u> <u>002</u>	Aug 23, 1983
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BUTAL COMPOUND

<u>AB</u> SANDOZ	<u>325MG;50MG;40MG</u>	<u>N86398</u> <u>002</u>	Apr 06, 1984
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BUTALBITAL, ASPIRIN AND CAFFEINE

<u>AB</u> + WEST WARD	<u>325MG;50MG;40MG</u>	<u>N86162</u> <u>002</u>	Feb 16, 1984
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ASPIRIN; BUTALBITAL; CAFFEINE; CODEINE PHOSPHATE

CAPSULE; ORAL

BUTALBITAL, ASPIRIN, CAFFEINE, AND CODEINE PHOSPHATE

<u>AB</u> ANABOLIC	<u>325MG;50MG;40MG;30MG</u>	<u>N75231</u> <u>001</u>	Nov 30, 2001
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<u>AB</u> STEVENS J	<u>325MG;50MG;40MG;30MG</u>	<u>N74951</u> <u>001</u>	Aug 31, 1998
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<u>AB</u> WATSON LABS	<u>325MG;50MG;40MG;30MG</u>	<u>N74359</u> <u>001</u>	Aug 31, 1995
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FIORINAL W/CODEINE

<u>AB</u> + WATSON PHARMS	<u>325MG;50MG;40MG;30MG</u>	<u>N19429</u> <u>003</u>	Oct 26, 1990
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27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 36 (of 370)

ASPIRIN; CAFFEINE; DIHYDROCODEINE BITARTRATE

CAPSULE; ORAL

SYNALGOS-DC

+	LEITNER PHARMS	356.4MG;30MG;16MG	N11483	004	Sep 06, 1983
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ASPIRIN; CAFFEINE; ORPHENADRINE CITRATE

TABLET; ORAL

INVAGESIC

<u>AB</u>	SANDOZ	<u>385MG;30MG;25MG</u>	<u>N74817</u>	<u>001</u>	Nov 27, 1996
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INVAGESIC FORTE

<u>AB</u>	SANDOZ	<u>770MG;60MG;50MG</u>	<u>N74817</u>	<u>002</u>	Nov 27, 1996
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NORGESIC

<u>AB</u>	3M	<u>385MG;30MG;25MG</u>	<u>N13416</u>	<u>003</u>	Oct 27, 1982
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NORGESIC FORTE

<u>AB</u>	+ 3M	<u>770MG;60MG;50MG</u>	<u>N13416</u>	<u>004</u>	Oct 27, 1982
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ORPHENADRINE CITRATE, ASPIRIN, AND CAFFEINE

<u>AB</u>	SANDOZ	<u>385MG;30MG;25MG</u>	<u>N74654</u>	<u>001</u>	Dec 31, 1996
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<u>AB</u>		<u>770MG;60MG;50MG</u>	<u>N74654</u>	<u>002</u>	Dec 31, 1996
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<u>AB</u>	STEVENS J	<u>385MG;30MG;25MG</u>	<u>N74988</u>	<u>001</u>	Apr 30, 1999
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<u>AB</u>		<u>770MG;60MG;50MG</u>	<u>N74988</u>	<u>002</u>	Apr 30, 1999
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ORPHENGESIC

<u>AB</u>	PAR PHARM	<u>385MG;30MG;25MG</u>	<u>N75141</u>	<u>001</u>	May 29, 1998
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ORPHENGESIC FORTE

<u>AB</u>	PAR PHARM	<u>770MG;60MG;50MG</u>	<u>N75141</u>	<u>002</u>	May 29, 1998
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ASPIRIN; CAFFEINE; PROPOXYPHENE HYDROCHLORIDE

CAPSULE; ORAL

DARVON COMPOUND-65

<u>AA</u>	+ XANODYNE PHARM	<u>389MG;32.4MG;65MG</u>	<u>N10996</u>	<u>007</u>	Mar 08, 1983
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PROPOXYPHENE COMPOUND 65

<u>AA</u>	TEVA	<u>389MG;32.4MG;65MG</u>	<u>N89025</u>	<u>001</u>	Mar 29, 1985
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ASPIRIN; CARISOPRODOL

TABLET; ORAL

CARISOPRODOL AND ASPIRIN

<u>AB</u>	ACTAVIS TOTOWA	<u>325MG;200MG</u>	<u>N40252</u>	<u>001</u>	Dec 10, 1997
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<u>AB</u>	PAR PHARM	<u>325MG;200MG</u>	<u>N89594</u>	<u>001</u>	Mar 31, 1989
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<u>AB</u>	SANDOZ	<u>325MG;200MG</u>	<u>N40116</u>	<u>001</u>	Apr 25, 1996
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SOMA COMPOUND

<u>AB</u>	+ MEDPOINTE PHARM HLC	<u>325MG;200MG</u>	<u>N12365</u>	<u>005</u>	Jul 11, 1983
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ASPIRIN; CARISOPRODOL; CODEINE PHOSPHATE

TABLET; ORAL

CARISOPRODOL, ASPIRIN AND CODEINE PHOSPHATE

<u>AB</u>	ACTAVIS TOTOWA	<u>325MG;200MG;16MG</u>	<u>N40283</u>	<u>001</u>	Dec 29, 1998
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<u>AB</u>	SANDOZ	<u>325MG;200MG;16MG</u>	<u>N40118</u>	<u>001</u>	Apr 16, 1996
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SOMA COMPOUND W/ CODEINE

<u>AB</u>	+ MEDPOINTE PHARM HLC	<u>325MG;200MG;16MG</u>	<u>N12366</u>	<u>002</u>	Jul 11, 1983
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ASPIRIN; DIPYRIDAMOLE

CAPSULE, EXTENDED RELEASE; ORAL

AGGRENOLX

+	BOEHRINGER INGELHEIM	25MG;200MG	N20884	001	Nov 22, 1999
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ASPIRIN; HYDROCODONE BITARTRATE

TABLET; ORAL

AZDONE

+	SCHWARZ PHARMA	500MG;5MG	N89420	001	Jan 25, 1988
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27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 37 (of 370)

ASPIRIN; MEPROBAMATETABLET; ORAL
EQUAGESIC

+ LEITNER PHARMS 325MG;200MG N11702 003 Dec 29, 1983

ASPIRIN; METHOCARBAMOL

TABLET; ORAL

METHOCARBAMOL AND ASPIRIN

<u>AB</u>	+	IVAX PHARMS	<u>325MG;400MG</u>	<u>N87211</u>	<u>001</u>	Dec 22, 1982
<u>AB</u>		PAR PHARM	<u>325MG;400MG</u>	<u>N89657</u>	<u>001</u>	Nov 04, 1988
<u>AB</u>		STEVENS J	<u>325MG;400MG</u>	<u>N81145</u>	<u>001</u>	Jan 31, 1995

ASPIRIN; OXYCODONE HYDROCHLORIDE

TABLET; ORAL

ASPIRIN AND OXYCODONE

+ ENDO PHARMS 325MG;4.8355MG N07337 007 Aug 05, 2005

ASPIRIN; OXYCODONE HYDROCHLORIDE; OXYCODONE TEREPHTHALATE

TABLET; ORAL

OXYCODONE AND ASPIRIN

<u>AA</u>		MUTUAL PHARM	<u>325MG;4.5MG;0.38MG</u>	<u>N40260</u>	<u>001</u>	Jul 17, 1998
<u>AA</u>			<u>325MG;4.5MG;0.38MG</u>	<u>N87794</u>	<u>001</u>	May 26, 1982
<u>AA</u>		WATSON LABS	<u>325MG;4.5MG;0.38MG</u>	<u>N40255</u>	<u>001</u>	Feb 27, 1998

PERCODAN

<u>AA</u>	+	ENDO PHARMS	<u>325MG;4.5MG;0.38MG</u>	<u>N07337</u>	<u>006</u>	
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ASPIRIN; PRAVASTATIN SODIUM

TABLET, TABLET; ORAL

PRAVIGARD PAC (COPACKAGED)

		BRISTOL MYERS SQUIBB	81MG,N/A;N/A,40MG	N21387	002	Jun 24, 2003
			81MG,N/A;N/A,20MG	N21387	001	Jun 24, 2003
			81MG,N/A;N/A,80MG	N21387	003	Jun 24, 2003
			325MG,N/A;N/A,40MG	N21387	005	Jun 24, 2003
	+		325MG,N/A;N/A,80MG	N21387	006	Jun 24, 2003
			325MG,N/A;N/A,20MG	N21387	004	Jun 24, 2003

ATAZANAVIR SULFATE

CAPSULE; ORAL

REYATAZ

		BRISTOL MYERS SQUIBB	EQ 100MG BASE	N21567	001	Jun 20, 2003
			EQ 150MG BASE	N21567	002	Jun 20, 2003
			EQ 200MG BASE	N21567	003	Jun 20, 2003
	+		EQ 300MG BASE	N21567	004	Oct 16, 2006

ATENOLOL

INJECTABLE; INJECTION

TENORMIN

+ ASTRAZENECA 0.5MG/ML N19058 001 Sep 13, 1989

TABLET; ORAL

ATENOLOL

<u>AB</u>		CLONMEL HLTHCARE	<u>25MG</u>	<u>N74099</u>	<u>001</u>	Apr 28, 1992
<u>AB</u>			<u>50MG</u>	<u>N73542</u>	<u>001</u>	Dec 19, 1991
<u>AB</u>			<u>100MG</u>	<u>N73543</u>	<u>001</u>	Dec 19, 1991
<u>AB</u>		GENPHARM	<u>25MG</u>	<u>N74126</u>	<u>003</u>	Aug 26, 1998
<u>AB</u>			<u>50MG</u>	<u>N74126</u>	<u>001</u>	Mar 23, 1994
<u>AB</u>			<u>100MG</u>	<u>N74126</u>	<u>002</u>	Mar 23, 1994
<u>AB</u>		IPR	<u>25MG</u>	<u>N73646</u>	<u>001</u>	Jul 31, 1992
<u>AB</u>			<u>50MG</u>	<u>N72303</u>	<u>001</u>	Jul 15, 1988
<u>AB</u>			<u>100MG</u>	<u>N72304</u>	<u>001</u>	Jul 15, 1988

PRESCRIPTION DRUG PRODUCT LIST

3 - 38 (of 370)

ATENOLOL

TABLET; ORAL

ATENOLOL

<u>AB</u>	MARTEC USA LLC	<u>50MG</u>	<u>N74127</u>	<u>001</u>	Feb 21, 1995
<u>AB</u>		<u>100MG</u>	<u>N74127</u>	<u>002</u>	Feb 21, 1995
<u>AB</u>	MUTUAL PHARM	<u>25MG</u>	<u>N74499</u>	<u>001</u>	Jul 30, 1997
<u>AB</u>		<u>50MG</u>	<u>N73475</u>	<u>001</u>	Mar 30, 1993
<u>AB</u>		<u>100MG</u>	<u>N73476</u>	<u>001</u>	Mar 30, 1993
<u>AB</u>	MYLAN	<u>25MG</u>	<u>N73457</u>	<u>002</u>	Apr 26, 1999
<u>AB</u>		<u>50MG</u>	<u>N73456</u>	<u>001</u>	Jan 24, 1992
<u>AB</u>		<u>100MG</u>	<u>N73457</u>	<u>001</u>	Jan 24, 1992
<u>AB</u>	OHM LABS	<u>25MG</u>	<u>N77877</u>	<u>001</u>	Dec 27, 2006
<u>AB</u>		<u>50MG</u>	<u>N77877</u>	<u>002</u>	Dec 27, 2006
<u>AB</u>		<u>100MG</u>	<u>N77877</u>	<u>003</u>	Dec 27, 2006
<u>AB</u>	PLIVA	<u>25MG</u>	<u>N74101</u>	<u>001</u>	Jul 17, 1997
<u>AB</u>		<u>50MG</u>	<u>N74101</u>	<u>002</u>	Jul 17, 1997
<u>AB</u>		<u>100MG</u>	<u>N74101</u>	<u>003</u>	Jul 17, 1997
<u>AB</u>	SANDOZ	<u>25MG</u>	<u>N74052</u>	<u>001</u>	May 01, 1992
<u>AB</u>		<u>25MG</u>	<u>N74265</u>	<u>001</u>	Feb 28, 1994
<u>AB</u>		<u>50MG</u>	<u>N73025</u>	<u>001</u>	Sep 17, 1991
<u>AB</u>		<u>50MG</u>	<u>N74265</u>	<u>002</u>	Feb 28, 1994
<u>AB</u>		<u>100MG</u>	<u>N73026</u>	<u>001</u>	Sep 17, 1991
<u>AB</u>		<u>100MG</u>	<u>N74265</u>	<u>003</u>	Feb 28, 1994
<u>AB</u>	SCS	<u>50MG</u>	<u>N73676</u>	<u>001</u>	Oct 30, 1992
<u>AB</u>		<u>100MG</u>	<u>N73676</u>	<u>002</u>	Oct 30, 1992
<u>AB</u>	TEVA	<u>25MG</u>	<u>N74056</u>	<u>003</u>	Jul 19, 2004
<u>AB</u>		<u>50MG</u>	<u>N73315</u>	<u>001</u>	May 28, 1993
<u>AB</u>		<u>50MG</u>	<u>N74056</u>	<u>001</u>	Jan 18, 1995
<u>AB</u>		<u>100MG</u>	<u>N73316</u>	<u>001</u>	May 28, 1993
<u>AB</u>		<u>100MG</u>	<u>N74056</u>	<u>002</u>	Jan 18, 1995
<u>AB</u>	TEVA PHARMS	<u>50MG</u>	<u>N74120</u>	<u>001</u>	Feb 24, 1995
<u>AB</u>		<u>100MG</u>	<u>N74120</u>	<u>002</u>	Feb 24, 1995
<u>AB</u>	UNIQUE PHARM LABS	<u>25MG</u>	<u>N77443</u>	<u>001</u>	Sep 13, 2006
<u>AB</u>		<u>50MG</u>	<u>N77443</u>	<u>002</u>	Sep 13, 2006
<u>AB</u>		<u>100MG</u>	<u>N77443</u>	<u>003</u>	Sep 13, 2006
<u>AB</u>	WATSON LABS	<u>50MG</u>	<u>N73352</u>	<u>001</u>	Dec 27, 1991
<u>AB</u>		<u>100MG</u>	<u>N73353</u>	<u>001</u>	Dec 27, 1991
<u>AB</u>	ZYDUS PHARMS USA	<u>25MG</u>	<u>N76900</u>	<u>001</u>	Jan 28, 2005
<u>AB</u>		<u>50MG</u>	<u>N76900</u>	<u>002</u>	Jan 28, 2005
<u>AB</u>		<u>100MG</u>	<u>N76900</u>	<u>003</u>	Jan 28, 2005
<u>TENORMIN</u>					
<u>AB</u>	ASTRAZENECA	<u>25MG</u>	<u>N18240</u>	<u>004</u>	Apr 09, 1990
<u>AB</u>		<u>50MG</u>	<u>N18240</u>	<u>001</u>	
<u>AB</u>	+	<u>100MG</u>	<u>N18240</u>	<u>002</u>	

ATENOLOL; CHLORTHALIDONE

TABLET; ORAL

ATENOLOL AND CHLORTHALIDONE

<u>AB</u>	IPR	<u>50MG; 25MG</u>	<u>N72301</u>	<u>001</u>	May 31, 1990
<u>AB</u>		<u>100MG; 25MG</u>	<u>N72302</u>	<u>001</u>	May 31, 1990
<u>AB</u>	MARTEC USA LLC	<u>50MG; 25MG</u>	<u>N74404</u>	<u>001</u>	May 14, 1998
<u>AB</u>		<u>100MG; 25MG</u>	<u>N74404</u>	<u>002</u>	May 14, 1998
<u>AB</u>	MUTUAL PHARM	<u>50MG; 25MG</u>	<u>N73581</u>	<u>001</u>	Apr 29, 1993
<u>AB</u>		<u>100MG; 25MG</u>	<u>N73582</u>	<u>001</u>	Apr 29, 1993
<u>AB</u>	MYLAN	<u>50MG; 25MG</u>	<u>N74203</u>	<u>001</u>	Oct 31, 1993
<u>AB</u>		<u>100MG; 25MG</u>	<u>N74203</u>	<u>002</u>	Oct 31, 1993
<u>AB</u>	PLIVA	<u>50MG; 25MG</u>	<u>N74107</u>	<u>001</u>	Sep 24, 1997
<u>AB</u>		<u>100MG; 25MG</u>	<u>N74107</u>	<u>002</u>	Sep 24, 1997
<u>AB</u>	WATSON LABS	<u>50MG; 25MG</u>	<u>N73665</u>	<u>001</u>	Jul 02, 1992
<u>AB</u>		<u>100MG; 25MG</u>	<u>N73665</u>	<u>002</u>	Jul 02, 1992

PRESCRIPTION DRUG PRODUCT LIST

3 - 39 (of 370)

ATENOLOL; CHLORTHALIDONE

TABLET; ORAL

<u>TENORETIC 100</u>					
<u>AB</u>	+ ASTRAZENECA	<u>100MG; 25MG</u>	<u>N18760</u>	<u>001</u>	Jun 08, 1984
<u>TENORETIC 50</u>					
<u>AB</u>	ASTRAZENECA	<u>50MG; 25MG</u>	<u>N18760</u>	<u>002</u>	Jun 08, 1984

ATOMOXETINE HYDROCHLORIDE

CAPSULE; ORAL

STRATTERA

LILLY

	10MG	N21411	002	Nov 26, 2002
	18MG	N21411	003	Nov 26, 2002
	25MG	N21411	004	Nov 26, 2002
	40MG	N21411	005	Nov 26, 2002
+	60MG	N21411	006	Nov 26, 2002
	80MG	N21411	007	Feb 14, 2005
	100MG	N21411	008	Feb 14, 2005

ATORVASTATIN CALCIUM

TABLET; ORAL

LIPITOR

PFIZER

	EQ 10MG BASE	N20702	001	Dec 17, 1996
	EQ 20MG BASE	N20702	002	Dec 17, 1996
	EQ 40MG BASE	N20702	003	Dec 17, 1996
+	EQ 80MG BASE	N20702	004	Apr 07, 2000

ATOVAQUONE

SUSPENSION; ORAL

MEPRON

+	GLAXOSMITHKLINE	750MG/5ML	N20500	001	Feb 08, 1995
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ATOVAQUONE; PROGUANIL HYDROCHLORIDE

TABLET; ORAL

MALARONE

+	GLAXOSMITHKLINE	250MG; 100MG	N21078	001	Jul 14, 2000
MALARONE PEDIATRIC					
+	GLAXOSMITHKLINE	62.5MG; 25MG	N21078	002	Jul 14, 2000

ATRACURIUM BESYLATE

INJECTABLE; INJECTION

ATRACURIUM BESYLATE

<u>AP</u>	BAXTER HLTHCARE	<u>10MG/ML</u>	<u>N74824</u>	<u>001</u>	Sep 30, 1997
<u>AP</u>	BAXTER HLTHCARE CORP	<u>10MG/ML</u>	<u>N74753</u>	<u>001</u>	Jan 23, 1997
<u>AP</u>	BEDFORD	<u>10MG/ML</u>	<u>N74901</u>	<u>001</u>	Jul 18, 1997
<u>AP</u>	HOSPIRA	<u>10MG/ML</u>	<u>N74632</u>	<u>001</u>	Dec 23, 1996
<u>AP</u>	MARSAM PHARMS LLC	<u>10MG/ML</u>	<u>N74945</u>	<u>001</u>	Jul 28, 1998
<u>AP</u>	MAYNE PHARMA USA	<u>10MG/ML</u>	<u>N74740</u>	<u>001</u>	Mar 28, 1997
<u>AP</u>	SICOR PHARMS	<u>10MG/ML</u>	<u>N74784</u>	<u>001</u>	Jun 11, 1997

ATRACURIUM BESYLATE PRESERVATIVE FREE

<u>AP</u>	BAXTER HLTHCARE CORP	<u>10MG/ML</u>	<u>N74768</u>	<u>001</u>	Jan 23, 1997
<u>AP</u>	BEDFORD	<u>10MG/ML</u>	<u>N74900</u>	<u>001</u>	Jul 18, 1997
<u>AP</u>	HOSPIRA	<u>10MG/ML</u>	<u>N74633</u>	<u>001</u>	Dec 23, 1996
<u>AP</u>		<u>10MG/ML</u>	<u>N74639</u>	<u>001</u>	Mar 25, 1997
<u>AP</u>	MARSAM PHARMS LLC	<u>10MG/ML</u>	<u>N74944</u>	<u>001</u>	Jul 28, 1998
<u>AP</u>	MAYNE PHARMA USA	<u>10MG/ML</u>	<u>N74741</u>	<u>001</u>	Mar 28, 1997

TRACRIUM

<u>AP</u>	+	HOSPIRA	<u>10MG/ML</u>	<u>N18831</u>	<u>002</u>	Jun 20, 1985
		<u>TRACRIUM PRESERVATIVE FREE</u>				
<u>AP</u>	+	HOSPIRA	<u>10MG/ML</u>	<u>N18831</u>	<u>001</u>	Nov 23, 1983

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 40 (of 370)

ATROPINE

INJECTABLE; INJECTION

ATROPEN

+	MERIDIAN MEDCL TECHN	EQ 0.25MG SULFATE/0.3ML	N17106	004	Sep 17, 2004
+		EQ 0.5MG SULFATE/0.7ML	N17106	003	Jun 19, 2003
+		EQ 1MG SULFATE/0.7ML	N17106	002	Jun 19, 2003
+		EQ 2MG SULFATE/0.7ML	N17106	001	

ATROPINE SULFATE

INJECTABLE; IM-IV-SC

ATROPINE SULFATE ANSYR PLASTIC SYRINGE

	HOSPIRA	0.05MG/ML	N21146	002	Jul 09, 2001
+		0.1MG/ML	N21146	001	Jul 09, 2001

ATROPINE SULFATE; DIFENOXIN HYDROCHLORIDE

TABLET; ORAL

MOTOFEN

+	VALEANT	0.025MG;1MG	N17744	002	
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ATROPINE SULFATE; DIPHENOXYLATE HYDROCHLORIDE

CAPSULE; ORAL

DIPHENOXYLATE HYDROCHLORIDE W/ ATROPINE SULFATE

	SCHERER RP	0.025MG;2.5MG	N86440	001	
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SOLUTION; ORAL

DIPHENOXYLATE HYDROCHLORIDE AND ATROPINE SULFATE

<u>AA</u>	ROXANE	<u>0.025MG/5ML;2.5MG/5ML</u>	<u>N87708</u>	<u>001</u>	May 03, 1982
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LOMOTIL

<u>AA</u>	+	GD SEARLE LLC	<u>0.025MG/5ML;2.5MG/5ML</u>	<u>N12699</u>	<u>001</u>
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TABLET; ORAL

DIPHENOXYLATE HYDROCHLORIDE AND ATROPINE SULFATE

<u>AA</u>	LANNETT	<u>0.025MG;2.5MG</u>	<u>N85372</u>	<u>001</u>	May 02, 2000
<u>AA</u>	MYLAN	<u>0.025MG;2.5MG</u>	<u>N85762</u>	<u>001</u>	
<u>AA</u>	PAR PHARM	<u>0.025MG;2.5MG</u>	<u>N40357</u>	<u>001</u>	

DIPHENOXYLATE HYDROCHLORIDE W/ ATROPINE SULFATE

<u>AA</u>	IVAX PHARMS	<u>0.025MG;2.5MG</u>	<u>N86727</u>	<u>001</u>	
<u>AA</u>	SANDOZ	<u>0.025MG;2.5MG</u>	<u>N86173</u>	<u>001</u>	

LOMOTIL

<u>AA</u>	+	GD SEARLE LLC	<u>0.025MG;2.5MG</u>	<u>N12462</u>	<u>001</u>
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LONOX

<u>AA</u>	SANDOZ	<u>0.025MG;2.5MG</u>	<u>N85311</u>	<u>002</u>	
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ATROPINE SULFATE; EDROPHONIUM CHLORIDE

INJECTABLE; INJECTION

ENLON-PLUS

+	BAXTER HLTHCARE CORP	0.14MG/ML;10MG/ML	N19677	001	Nov 06, 1991
+		0.14MG/ML;10MG/ML	N19678	001	Nov 06, 1991

ATROPINE; PRALIDOXIME CHLORIDE

INJECTABLE; INTRAMUSCULAR

DUODOTE

+	MERIDIAN MEDCL	2.1MG/0.7ML;600MG/2ML	N21983	001	Sep 28, 2006
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AURANOFIN

CAPSULE; ORAL

RIDAURA

+	PROMETHEUS LABS	3MG	N18689	001	May 24, 1985
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27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 41 (of 370)

AZACITIDINE

INJECTABLE; SUBCUTANEOUS

VIDAZA

+ PHARMION

100MG/VIAL

N50794 001

May 19, 2004

AZATHIOPRINE

TABLET; ORAL

AZASANAB AAIPHARMA LLC50MGN75252 001

Jun 07, 1999

25MG

N75252 002

Feb 03, 2003

75MG

N75252 003

Feb 03, 2003

100MG

N75252 004

Feb 03, 2003

AZATHIOPRINEAB GENPHARM50MGN75568 001

Dec 13, 1999

AB ROXANE50MGN74069 001

Feb 16, 1996

IMURANAB + PROMETHEUS LABS50MGN16324 001AZATHIOPRINE SODIUM

INJECTABLE; INJECTION

AZATHIOPRINE SODIUM

+ BEDFORD

EQ 100MG BASE/VIAL

N74419 001

Mar 31, 1995

AZELAIC ACID

CREAM; TOPICAL

AZELEX

+ ALLERGAN

20%

N20428 001

Sep 13, 1995

GEL; TOPICAL

FINACEA

+ INTENDIS

15%

N21470 001

Dec 24, 2002

AZELASTINE HYDROCHLORIDE

SOLUTION/DROPS; OPHTHALMIC

+ MEDPOINTE PHARM HLC

0.05%

N21127 001

May 22, 2000

SPRAY, METERED; NASAL

ASTELIN

+ MEDPOINTE PHARM HLC

EQ 0.125MG BASE/SPRAY

N20114 001

Nov 01, 1996

AZITHROMYCIN

FOR SUSPENSION; ORAL

AZITHROMYCINAB PLIVAEQ 100MG BASE/5MLN65246 002

Jul 05, 2006

ABEQ 200MG BASE/5MLN65246 001

Jul 05, 2006

AB SANDOZEQ 100MG BASE/5MLN65297 001

Sep 18, 2006

ABEQ 200MG BASE/5MLN65297 002

Sep 18, 2006

ZITHROMAXAB PFIZEREQ 100MG BASE/5MLN50710 001

Oct 19, 1995

AB +EQ 200MG BASE/5MLN50710 002

Oct 19, 1995

+

EQ 1GM BASE/PACKET

N50693 001

Sep 28, 1994

FOR SUSPENSION, EXTENDED RELEASE; ORAL

ZMAX

+ PFIZER GLOBAL

EQ 2GM BASE/BOT

N50797 001

Jun 10, 2005

INJECTABLE; INJECTION

AZITHROMYCINAP ABRAXIS PHARMEQ 500MG BASE/VIALN65179 001

Dec 13, 2005

ZITHROMAXAP + PFIZEREQ 500MG BASE/VIALN50733 001

Jan 30, 1997

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 42 (of 370)

AZITHROMYCIN

TABLET; ORAL

AZITHROMYCIN

<u>AB</u>	MYLAN	<u>EQ 600MG BASE</u>	<u>N65360</u>	<u>001</u>	Jan 08, 2007
<u>AB</u>	PLIVA	<u>EQ 250MG BASE</u>	<u>N65225</u>	<u>001</u>	Nov 14, 2005
<u>AB</u>		<u>EQ 500MG BASE</u>	<u>N65223</u>	<u>001</u>	Nov 14, 2005
<u>AB</u>		<u>EQ 600MG BASE</u>	<u>N65218</u>	<u>001</u>	Nov 14, 2005
<u>AB</u>	SANDOZ	<u>EQ 250MG BASE</u>	<u>N65211</u>	<u>001</u>	Nov 14, 2005
<u>AB</u>		<u>EQ 500MG BASE</u>	<u>N65212</u>	<u>001</u>	Nov 14, 2005
<u>AB</u>		<u>EQ 600MG BASE</u>	<u>N65209</u>	<u>001</u>	Nov 14, 2005
<u>AB</u>	TEVA	<u>EQ 250MG BASE</u>	<u>N65153</u>	<u>001</u>	Nov 14, 2005
<u>AB</u>		<u>EQ 500MG BASE</u>	<u>N65193</u>	<u>001</u>	Nov 14, 2005
<u>AB</u>		<u>EQ 600MG BASE</u>	<u>N65150</u>	<u>001</u>	Nov 14, 2005
<u>ZITHROMAX</u>					
<u>AB</u>	PFIZER	<u>EQ 250MG BASE</u>	<u>N50711</u>	<u>001</u>	Jul 18, 1996
<u>AB</u>		<u>EQ 500MG BASE</u>	<u>N50784</u>	<u>001</u>	May 24, 2002
<u>AB</u>	+	<u>EQ 600MG BASE</u>	<u>N50730</u>	<u>001</u>	Jun 12, 1996

AZITHROMYCIN HYDROGENCITRATE

INJECTABLE; INJECTION

AZITHROMYCIN

+	SICOR PHARMS	EQ 500MG BASE/VIAL	N50809	001	Dec 19, 2006
+		EQ 2.5GM BASE/VIAL	N50809	002	Dec 19, 2006

AZTREONAM

INJECTABLE; INJECTION

AZACTAM

+	BRISTOL MYERS SQUIBB	500MG/VIAL	N50580	001	Dec 31, 1986
+		1GM/VIAL	N50580	002	Dec 31, 1986
+		2GM/VIAL	N50580	003	Dec 31, 1986
AZACTAM IN PLASTIC CONTAINER					
+	BRISTOL MYERS SQUIBB	20MG/ML	N50632	002	May 24, 1989
+		40MG/ML	N50632	001	May 24, 1989

BACITRACIN

INJECTABLE; INJECTION

BACIIM

<u>AP</u>	X GEN PHARMS	<u>50,000 UNITS/VIAL</u>	<u>N64153</u>	<u>001</u>	May 09, 1997
<u>BACITRACIN</u>					
<u>AP</u>	ABRAXIS PHARM	<u>50,000 UNITS/VIAL</u>	<u>N65116</u>	<u>001</u>	Dec 03, 2002
<u>AP</u>	+	<u>PHARMACIA AND UPJOHN</u>	<u>50,000 UNITS/VIAL</u>	<u>N60733</u>	<u>002</u>
		10,000 UNITS/VIAL	N60733	001	

OINTMENT; OPHTHALMIC

BACITRACIN

+	ALTANA	500 UNITS/GM	N61212	001	
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POWDER; FOR RX COMPOUNDING

BACI-RX

<u>AA</u>	X GEN PHARMS	<u>5,000,000 UNITS/BOT</u>	<u>N61580</u>	<u>001</u>	
<u>BACITRACIN</u>					
<u>AA</u>	APOTHEKERNES	<u>5,000,000 UNITS/BOT</u>	<u>N61699</u>	<u>001</u>	

BACITRACIN ZINC; HYDROCORTISONE; NEOMYCIN SULFATE; POLYMYXIN B SULFATE

OINTMENT; OPHTHALMIC

CORTISPORIN

<u>AT</u>	MONARCH PHARMS	<u>400 UNITS/GM;1%;EQ 3.5MG</u>	<u>N50416</u>	<u>002</u>	
		<u>BASE/GM;10,000 UNITS/GM</u>			

NEOMYCIN AND POLYMYXIN B SULFATES, BACITRACIN ZINC AND HYDROCORTISONE

<u>AT</u>	+	<u>BAUSCH AND LOMB</u>	<u>400 UNITS/GM;1%;EQ 3.5MG</u>	<u>N64068</u>	<u>001</u>	Oct 30, 1995
			<u>BASE/GM;10,000 UNITS/GM</u>			

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 43 (of 370)

BACITRACIN ZINC; HYDROCORTISONE; NEOMYCIN SULFATE; POLYMYXIN B SULFATE

OINTMENT; TOPICAL

CORTISPORIN

+	MONARCH PHARMS	400 UNITS/GM;1%;EQ 3.5MG BASE/GM;5,000 UNITS/GM	N50168	002	May 04, 1984
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BACITRACIN ZINC; NEOMYCIN SULFATE; POLYMYXIN B SULFATE

OINTMENT; OPHTHALMIC

BACITRACIN-NEOMYCIN-POLYMYXIN

<u>AT</u>	ALTANA	<u>400 UNITS/GM;EQ 3.5MG BASE/GM;10,000 UNITS/GM</u>	<u>N60764</u>	<u>002</u>	
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NEOMYCIN AND POLYMYXIN B SULFATE AND BACITRACIN ZINC

<u>AT</u>	AKORN	<u>400 UNITS/GM;EQ 3.5MG BASE/GM;10,000 UNITS/GM</u>	<u>N65088</u>	<u>001</u>	Feb 06, 2004
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NEOMYCIN AND POLYMYXIN B SULFATES AND BACITRACIN ZINC

<u>AT</u>	+	BAUSCH AND LOMB	<u>400 UNITS/GM;EQ 3.5MG BASE/GM;10,000 UNITS/GM</u>	<u>N64064</u>	<u>001</u>	Oct 30, 1995
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NEOSPORIN

<u>AT</u>	MONARCH PHARMS	<u>400 UNITS/GM;EQ 3.5MG BASE/GM;10,000 UNITS/GM</u>	<u>N50417</u>	<u>001</u>	
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BACITRACIN ZINC; POLYMYXIN B SULFATE

OINTMENT; OPHTHALMIC

BACITRACIN ZINC AND POLYMYXIN B SULFATE

<u>AT</u>	AKORN	<u>500 UNITS/GM;10,000 UNITS/GM</u>	<u>N64028</u>	<u>001</u>	Jan 30, 1995
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<u>AT</u>	ALTANA	<u>500 UNITS/GM;10,000 UNITS/GM</u>	<u>N65022</u>	<u>001</u>	Feb 27, 2002
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<u>AT</u>	+	BAUSCH AND LOMB	<u>500 UNITS/GM;10,000 UNITS/GM</u>	<u>N64046</u>	<u>001</u>	Jan 26, 1995
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POLYSPORIN

<u>AT</u>	MONARCH PHARMS	<u>500 UNITS/GM;10,000 UNITS/GM</u>	<u>N61229</u>	<u>001</u>	
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BACITRACIN; HYDROCORTISONE ACETATE; NEOMYCIN SULFATE; POLYMYXIN B SULFATE

OINTMENT; OPHTHALMIC

BACITRACIN-NEOMYCIN-POLYMYXIN W/ HYDROCORTISONE ACETATE

+	PHARMADERM	400 UNITS/GM;1%;EQ 3.5MG BASE/GM;10,000 UNITS/GM	N62166	002	
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BACLOFEN

INJECTABLE; INTRATHECAL

LIORESAL

+	MEDTRONIC	0.05MG/ML	N20075	003	Nov 07, 1996
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+		0.5MG/ML	N20075	001	Jun 17, 1992
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+		2MG/ML	N20075	002	Jun 17, 1992
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TABLET; ORAL

BACLOFEN

<u>AB</u>	ALPHAPHARM	<u>10MG</u>	<u>N77181</u>	<u>001</u>	Jul 29, 2005
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<u>AB</u>		<u>20MG</u>	<u>N77121</u>	<u>002</u>	Jul 29, 2005
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<u>AB</u>	CARACO	<u>10MG</u>	<u>N77984</u>	<u>001</u>	Aug 14, 2006
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<u>AB</u>		<u>20MG</u>	<u>N77862</u>	<u>002</u>	Aug 14, 2006
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<u>AB</u>	IVAX PHARMS	<u>10MG</u>	<u>N72234</u>	<u>001</u>	Jul 21, 1988
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<u>AB</u>	+	<u>20MG</u>	<u>N72235</u>	<u>001</u>	Jul 21, 1988
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<u>AB</u>	LANNETT	<u>20MG</u>	<u>N77241</u>	<u>001</u>	Dec 20, 2005
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<u>AB</u>	USL PHARMA	<u>10MG</u>	<u>N74584</u>	<u>001</u>	Aug 19, 1996
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<u>AB</u>		<u>20MG</u>	<u>N74584</u>	<u>002</u>	Aug 19, 1996
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<u>AB</u>	VINTAGE PHARMS	<u>10MG</u>	<u>N77156</u>	<u>001</u>	Aug 30, 2005
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<u>AB</u>		<u>20MG</u>	<u>N77068</u>	<u>001</u>	Aug 30, 2005
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<u>AB</u>	WATSON LABS	<u>10MG</u>	<u>N72824</u>	<u>001</u>	Sep 18, 1991
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<u>AB</u>		<u>10MG</u>	<u>N73092</u>	<u>001</u>	Jan 28, 1994
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<u>AB</u>		<u>20MG</u>	<u>N72825</u>	<u>001</u>	Sep 18, 1991
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PRESCRIPTION DRUG PRODUCT LIST

3 - 44 (of 370)

BACLOFEN

TABLET, ORALLY DISINTEGRATING; ORAL

KEMSTRO

SCHWARZ PHARMA	10MG	N21589	001	Oct 30, 2003
+	20MG	N21589	002	Oct 30, 2003

BALSALAZIDE DISODIUM

CAPSULE; ORAL

COLAZAL

+ SALIX PHARMS	750MG	N20610	001	Jul 18, 2000
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BECLOMETHASONE DIPROPIONATE

AEROSOL, METERED; INHALATION

QVAR 40

+ 3M	0.04MG/INH	N20911	002	Sep 15, 2000
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QVAR 80

+ 3M	0.08MG/INH	N20911	001	Sep 15, 2000
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BECLOMETHASONE DIPROPIONATE MONOHYDRATE

SPRAY, METERED; NASAL

BECONASE AQ

+ GLAXOSMITHKLINE	EQ 0.042MG DIPROP/SPRAY	N19389	001	Jul 27, 1987
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BENZAEPRIIL HYDROCHLORIDE

TABLET; ORAL

BENZAEPRIIL HYDROCHLORIDE

<u>AB</u>	ANDRX PHARMS	<u>5MG</u>	<u>N76267</u>	<u>001</u>	Feb 11, 2004
<u>AB</u>		<u>10MG</u>	<u>N76267</u>	<u>002</u>	Feb 11, 2004
<u>AB</u>		<u>20MG</u>	<u>N76267</u>	<u>003</u>	Feb 11, 2004
<u>AB</u>		<u>40MG</u>	<u>N76267</u>	<u>004</u>	Feb 11, 2004
<u>AB</u>	APOTEX INC	<u>5MG</u>	<u>N77128</u>	<u>001</u>	Mar 08, 2006
<u>AB</u>		<u>10MG</u>	<u>N77128</u>	<u>002</u>	Mar 08, 2006
<u>AB</u>		<u>20MG</u>	<u>N77128</u>	<u>003</u>	Mar 08, 2006
<u>AB</u>		<u>40MG</u>	<u>N77128</u>	<u>004</u>	Mar 08, 2006
<u>AB</u>	BIOKEY	<u>5MG</u>	<u>N76820</u>	<u>001</u>	Feb 03, 2006
<u>AB</u>		<u>10MG</u>	<u>N76820</u>	<u>002</u>	Feb 03, 2006
<u>AB</u>		<u>20MG</u>	<u>N76820</u>	<u>003</u>	Feb 03, 2006
<u>AB</u>		<u>40MG</u>	<u>N76820</u>	<u>004</u>	Feb 03, 2006
<u>AB</u>	GENPHARM	<u>5MG</u>	<u>N76476</u>	<u>001</u>	Feb 11, 2004
<u>AB</u>		<u>10MG</u>	<u>N76476</u>	<u>002</u>	Feb 11, 2004
<u>AB</u>		<u>20MG</u>	<u>N76476</u>	<u>003</u>	Feb 11, 2004
<u>AB</u>		<u>40MG</u>	<u>N76476</u>	<u>004</u>	Feb 11, 2004
<u>AB</u>	IVAX PHARMS	<u>5MG</u>	<u>N76333</u>	<u>001</u>	Feb 11, 2004
<u>AB</u>		<u>10MG</u>	<u>N76333</u>	<u>002</u>	Feb 11, 2004
<u>AB</u>		<u>20MG</u>	<u>N76333</u>	<u>003</u>	Feb 11, 2004
<u>AB</u>		<u>40MG</u>	<u>N76333</u>	<u>004</u>	Feb 11, 2004
<u>AB</u>	KV PHARM	<u>5MG</u>	<u>N76118</u>	<u>001</u>	Feb 11, 2004
<u>AB</u>		<u>10MG</u>	<u>N76118</u>	<u>002</u>	Feb 11, 2004
<u>AB</u>		<u>20MG</u>	<u>N76118</u>	<u>003</u>	Feb 11, 2004
<u>AB</u>		<u>40MG</u>	<u>N76118</u>	<u>004</u>	Feb 11, 2004
<u>AB</u>	MYLAN	<u>5MG</u>	<u>N76430</u>	<u>001</u>	Feb 11, 2004
<u>AB</u>		<u>10MG</u>	<u>N76430</u>	<u>002</u>	Feb 11, 2004
<u>AB</u>		<u>20MG</u>	<u>N76430</u>	<u>003</u>	Feb 11, 2004
<u>AB</u>		<u>40MG</u>	<u>N76430</u>	<u>004</u>	Feb 11, 2004
<u>AB</u>	RANBAXY	<u>5MG</u>	<u>N76344</u>	<u>001</u>	Feb 11, 2004
<u>AB</u>		<u>10MG</u>	<u>N76344</u>	<u>002</u>	Feb 11, 2004
<u>AB</u>		<u>20MG</u>	<u>N76344</u>	<u>003</u>	Feb 11, 2004
<u>AB</u>		<u>40MG</u>	<u>N76344</u>	<u>004</u>	Feb 11, 2004
<u>AB</u>	SANDOZ	<u>5MG</u>	<u>N76402</u>	<u>001</u>	Feb 11, 2004

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 45 (of 370)

BENAZEPRIL HYDROCHLORIDE

TABLET; ORAL

BENAZEPRIL HYDROCHLORIDE

<u>AB</u>	SANDOZ	<u>10MG</u>	<u>N76402</u>	<u>002</u>	Feb 11, 2004
<u>AB</u>		<u>20MG</u>	<u>N76402</u>	<u>003</u>	Feb 11, 2004
<u>AB</u>		<u>40MG</u>	<u>N76402</u>	<u>004</u>	Feb 11, 2004
<u>AB</u>	TEVA	<u>5MG</u>	<u>N76211</u>	<u>001</u>	Feb 11, 2004
<u>AB</u>		<u>10MG</u>	<u>N76211</u>	<u>002</u>	Feb 11, 2004
<u>AB</u>		<u>20MG</u>	<u>N76211</u>	<u>003</u>	Feb 11, 2004
<u>AB</u>		<u>40MG</u>	<u>N76211</u>	<u>004</u>	Feb 11, 2004
<u>LOTENSIN</u>					
<u>AB</u>	NOVARTIS	<u>5MG</u>	<u>N19851</u>	<u>001</u>	Jun 25, 1991
<u>AB</u>		<u>10MG</u>	<u>N19851</u>	<u>002</u>	Jun 25, 1991
<u>AB</u>		<u>20MG</u>	<u>N19851</u>	<u>003</u>	Jun 25, 1991
<u>AB</u>	+	<u>40MG</u>	<u>N19851</u>	<u>004</u>	Jun 25, 1991

BENAZEPRIL HYDROCHLORIDE; HYDROCHLOROTHIAZIDE

TABLET; ORAL

BENAZEPRIL HYDROCHLORIDE AND HYDROCHLOROTHIAZIDE

<u>AB</u>	ANDRX PHARMS	<u>5MG; 6.25MG</u>	<u>N76342</u>	<u>001</u>	Feb 11, 2004
<u>AB</u>		<u>10MG; 12.5MG</u>	<u>N76342</u>	<u>002</u>	Feb 11, 2004
<u>AB</u>		<u>20MG; 12.5MG</u>	<u>N76342</u>	<u>003</u>	Feb 11, 2004
<u>AB</u>		<u>20MG; 25MG</u>	<u>N76342</u>	<u>004</u>	Feb 11, 2004
<u>AB</u>	GENPHARM	<u>5MG; 6.25MG</u>	<u>N76612</u>	<u>001</u>	Feb 11, 2004
<u>AB</u>		<u>10MG; 12.5MG</u>	<u>N76612</u>	<u>002</u>	Feb 11, 2004
<u>AB</u>		<u>20MG; 12.5MG</u>	<u>N76612</u>	<u>003</u>	Feb 11, 2004
<u>AB</u>		<u>20MG; 25MG</u>	<u>N76612</u>	<u>004</u>	Feb 11, 2004
<u>AB</u>	IVAX PHARMS	<u>5MG; 6.25MG</u>	<u>N76348</u>	<u>001</u>	Feb 11, 2004
<u>AB</u>		<u>10MG; 12.5MG</u>	<u>N76348</u>	<u>002</u>	Feb 11, 2004
<u>AB</u>		<u>20MG; 12.5MG</u>	<u>N76348</u>	<u>003</u>	Feb 11, 2004
<u>AB</u>		<u>20MG; 25MG</u>	<u>N76348</u>	<u>004</u>	Feb 11, 2004
<u>AB</u>	MYLAN	<u>5MG; 6.25MG</u>	<u>N76688</u>	<u>001</u>	Feb 11, 2004
<u>AB</u>		<u>10MG; 12.5MG</u>	<u>N76688</u>	<u>002</u>	Feb 11, 2004
<u>AB</u>		<u>20MG; 12.5MG</u>	<u>N76688</u>	<u>003</u>	Feb 11, 2004
<u>AB</u>		<u>20MG; 25MG</u>	<u>N76688</u>	<u>004</u>	Feb 11, 2004
<u>AB</u>	RANBAXY	<u>5MG; 6.25MG</u>	<u>N77483</u>	<u>001</u>	Sep 08, 2005
<u>AB</u>		<u>10MG; 12.5MG</u>	<u>N77483</u>	<u>002</u>	Sep 08, 2005
<u>AB</u>		<u>20MG; 12.5MG</u>	<u>N77483</u>	<u>003</u>	Sep 08, 2005
<u>AB</u>		<u>20MG; 25MG</u>	<u>N77483</u>	<u>004</u>	Sep 08, 2005
<u>AB</u>	SANDOZ	<u>5MG; 6.25MG</u>	<u>N76631</u>	<u>001</u>	Feb 11, 2004
<u>AB</u>		<u>10MG; 12.5MG</u>	<u>N76631</u>	<u>002</u>	Feb 11, 2004
<u>AB</u>		<u>20MG; 12.5MG</u>	<u>N76631</u>	<u>003</u>	Feb 11, 2004
<u>AB</u>		<u>20MG; 25MG</u>	<u>N76631</u>	<u>004</u>	Feb 11, 2004
<u>LOTENSIN HCT</u>					
<u>AB</u>	NOVARTIS	<u>5MG; 6.25MG</u>	<u>N20033</u>	<u>001</u>	May 19, 1992
<u>AB</u>		<u>10MG; 12.5MG</u>	<u>N20033</u>	<u>002</u>	May 19, 1992
<u>AB</u>		<u>20MG; 12.5MG</u>	<u>N20033</u>	<u>004</u>	May 19, 1992
<u>AB</u>	+	<u>20MG; 25MG</u>	<u>N20033</u>	<u>003</u>	May 19, 1992

BENDROFLUMETHIAZIDE; NADOLOL

TABLET; ORAL

CORZIDE

	KING PHARMS	5MG; 40MG	N18647	001	May 25, 1983
+		5MG; 80MG	N18647	002	May 25, 1983

BENZONATATE

CAPSULE; ORAL

BENZONATATE

<u>AA</u>	BANNER PHARMACAPS	<u>100MG</u>	<u>N81297</u>	<u>001</u>	Jan 29, 1993
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PRESCRIPTION DRUG PRODUCT LIST

3 - 46 (of 370)

BENZONATATE

CAPSULE; ORAL

TESSALON

<u>AA</u>	+ FOREST LABS	<u>100MG</u>	<u>N11210</u>	<u>001</u>	
	+	<u>200MG</u>	<u>N11210</u>	<u>003</u>	Jun 25, 1999

BENZOYL PEROXIDE; CLINDAMYCIN PHOSPHATE

GEL; TOPICAL

BENZACLIN

BT	+ SANOFI AVENTIS US	5%;EQ 1% BASE	N50756	001	Dec 21, 2000
	DUAC				
BT	+ STIEFEL	5%;EQ 1% BASE	N50741	001	Aug 26, 2002

BENZOYL PEROXIDE; ERYTHROMYCIN

GEL; TOPICAL

BENZAMYCIN

<u>AB</u>	+ SANOFI AVENTIS US	<u>5%;3%</u>	<u>N50557</u>	<u>001</u>	Oct 26, 1984
	BENZAMYCIN PAK				
	+ SANOFI AVENTIS US	5%;3%	N50769	001	Nov 27, 2000

ERYTHROMYCIN AND BENZOYL PEROXIDE

<u>AB</u>	QLT USA	<u>5%;3%</u>	<u>N65112</u>	<u>001</u>	Mar 29, 2004
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BENZPHETAMINE HYDROCHLORIDE

TABLET; ORAL

BENZPHETAMINE HYDROCHLORIDE

<u>AA</u>	PADDOCK	<u>50MG</u>	<u>N40578</u>	<u>001</u>	Apr 17, 2006
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DIDREX

<u>AA</u>	+ PHARMACIA AND UPJOHN	<u>50MG</u>	<u>N12427</u>	<u>002</u>	
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BENZTROPINE MESYLATE

INJECTABLE; INJECTION

COGENTIN

	+ OVATION PHARMS	1MG/ML	N12015	001	
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TABLET; ORAL

BENZTROPINE MESYLATE

<u>AA</u>	COREPHARMA	<u>0.5MG</u>	<u>N72264</u>	<u>001</u>	Feb 27, 1989
<u>AA</u>		<u>1MG</u>	<u>N72265</u>	<u>001</u>	Feb 27, 1989
<u>AA</u>		<u>2MG</u>	<u>N72266</u>	<u>001</u>	Feb 27, 1989
<u>AA</u>	MUTUAL PHARM	<u>1MG</u>	<u>N81264</u>	<u>001</u>	Jan 23, 1992
<u>AA</u>		<u>2MG</u>	<u>N81265</u>	<u>001</u>	Jan 23, 1992
<u>AA</u>	+ PAR PHARM	<u>0.5MG</u>	<u>N88877</u>	<u>001</u>	Apr 11, 1985
<u>AA</u>	+	<u>1MG</u>	<u>N88894</u>	<u>001</u>	Apr 11, 1985
<u>AA</u>	+	<u>2MG</u>	<u>N88895</u>	<u>001</u>	Apr 11, 1985
<u>AA</u>	PLIVA	<u>0.5MG</u>	<u>N89058</u>	<u>001</u>	Aug 10, 1988
<u>AA</u>		<u>1MG</u>	<u>N89059</u>	<u>001</u>	Aug 10, 1988
<u>AA</u>		<u>2MG</u>	<u>N89060</u>	<u>001</u>	Aug 10, 1988
<u>AA</u>	USL PHARMA	<u>0.5MG</u>	<u>N40103</u>	<u>001</u>	Dec 12, 1996
<u>AA</u>		<u>1MG</u>	<u>N40103</u>	<u>002</u>	Dec 12, 1996
<u>AA</u>		<u>2MG</u>	<u>N40103</u>	<u>003</u>	Dec 12, 1996

BERACTANT

SUSPENSION; INTRATRACHEAL

SURVANTA

	+ ROSS LABS	25MG/ML	N20032	001	Jul 01, 1991
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BETAINE HYDROCHLORIDE

FOR SOLUTION; ORAL

CYSTADANE

	+ JAZZ	1GM/SCOOPFUL	N20576	001	Oct 25, 1996
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27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 47 (of 370)

BETAMETHASONE

SYRUP; ORAL
CELESTONE
+ SCHERING

0.6MG/5ML

N14215 002

BETAMETHASONE ACETATE; BETAMETHASONE SODIUM PHOSPHATE

INJECTABLE; INJECTION
CELESTONE SOLUSPAN
+ SCHERING

3MG/ML;EQ 3MG BASE/ML

N14602 001

BETAMETHASONE DIPROPIONATE

CREAM; TOPICAL

BETAMETHASONE DIPROPIONATE

AB ACTAVIS MID ATLANTIC EQ 0.05% BASE
AB + FOUGERA EQ 0.05% BASE
AB TARO EQ 0.05% BASE
AB EQ 0.05% BASE
AB TEVA EQ 0.05% BASE

N70885 001 Feb 03, 1987
N19137 001 Jun 26, 1984
N71143 001 Jun 17, 1987
N73552 001 Apr 30, 1992
N71476 001 Aug 10, 1987

CREAM, AUGMENTED; TOPICAL

BETAMETHASONE DIPROPIONATE

AB ALTANA EQ 0.05% BASE
AB PERRIGO NEW YORK EQ 0.05% BASE
AB QLT USA EQ 0.05% BASE
AB TARO EQ 0.05% BASE

N76215 001 Dec 09, 2003
N76592 001 Dec 09, 2003
N76603 001 Jan 23, 2004
N76543 001 Dec 09, 2003

DIPROLENE AF

AB + SCHERING EQ 0.05% BASE

N19555 001 Apr 27, 1987

GEL, AUGMENTED; TOPICAL

BETAMETHASONE DIPROPIONATE

AB + ALTANA EQ 0.05% BASE
AB TARO EQ 0.05% BASE

N75276 001 May 13, 2003
N76508 001 Dec 02, 2003

LOTION; TOPICAL

BETAMETHASONE DIPROPIONATE

AB ACTAVIS MID ATLANTIC EQ 0.05% BASE
AB + FOUGERA EQ 0.05% BASE
AB PERRIGO NEW YORK EQ 0.05% BASE
AB TARO EQ 0.05% BASE
AB EQ 0.05% BASE
AB TEVA EQ 0.05% BASE
AB TEVA PHARMS EQ 0.05% BASE

N70281 001 Jul 31, 1985
N70275 001 Aug 12, 1985
N72538 001 Jan 31, 1990
N72276 001 Aug 24, 1988
N74272 001 Sep 30, 1994
N71467 001 Aug 10, 1987
N71882 001 Jun 06, 1988

LOTION, AUGMENTED; TOPICAL

DIPROLENE

+ SCHERING EQ 0.05% BASE

N19716 001 Aug 01, 1988

OINTMENT; TOPICAL

BETAMETHASONE DIPROPIONATE

AB ACTAVIS MID ATLANTIC EQ 0.05% BASE
AB + FOUGERA EQ 0.05% BASE
AB TARO EQ 0.05% BASE
AB TEVA EQ 0.05% BASE

N71012 001 Feb 03, 1987
N19141 001 Sep 04, 1984
N74271 001 Sep 15, 1994
N71477 001 Aug 10, 1987

OINTMENT, AUGMENTED; TOPICAL

BETAMETHASONE DIPROPIONATE

AB ACTAVIS MID ATLANTIC EQ 0.05% BASE
AB ALTANA EQ 0.05% BASE
AB TARO EQ 0.05% BASE

N74304 001 Aug 31, 1995
N75373 001 Jun 22, 1999
N76753 001 Oct 12, 2004

DIPROLENE

AB + SCHERING EQ 0.05% BASE

N18741 001 Jul 27, 1983

PRESCRIPTION DRUG PRODUCT LIST

3 - 48 (of 370)

BETAMETHASONE DIPROPIONATE; CALCIPOTRIENE HYDRATE

OINTMENT; TOPICAL

TACLONEX

+ LEO PHARM PRODS	0.064%;0.005%	N21852	001	Jan 09, 2006
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BETAMETHASONE DIPROPIONATE; CLOTRIMAZOLE

CREAM; TOPICAL

CLOTRIMAZOLE AND BETAMETHASONE DIPROPIONATE

<u>AB</u> ACTAVIS MID ATLANTIC	<u>EQ 0.05% BASE;1%</u>	<u>N76002</u>	<u>001</u>	Aug 02, 2002
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<u>AB</u> ALTANA	<u>EQ 0.05% BASE;1%</u>	<u>N75502</u>	<u>001</u>	Jun 05, 2001
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<u>AB</u> TARO	<u>EQ 0.05% BASE;1%</u>	<u>N75673</u>	<u>001</u>	May 29, 2001
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LOTTRISONE

<u>AB</u> + SCHERING	<u>EQ 0.05% BASE;1%</u>	<u>N18827</u>	<u>001</u>	Jul 10, 1984
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LOTION; TOPICAL

CLOTRIMAZOLE AND BETAMETHASONE DIPROPIONATE

<u>AB</u> ALTANA PHARMA	<u>EQ 0.05% BASE;1%</u>	<u>N76516</u>	<u>001</u>	Jun 16, 2005
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<u>AB</u> TARO	<u>EQ 0.05% BASE;1%</u>	<u>N76493</u>	<u>001</u>	Jul 28, 2004
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LOTTRISONE

<u>AB</u> + SCHERING PLOUGH RES	<u>EQ 0.05% BASE;1%</u>	<u>N20010</u>	<u>001</u>	Dec 08, 2000
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BETAMETHASONE VALERATE

AEROSOL, FOAM; TOPICAL

LUXIQ

+ CONNETICS	EQ 0.12% BASE	N20934	001	Feb 28, 1999
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CREAM; TOPICAL

BETAMETHASONE VALERATE

<u>AB</u> + FOUGERA	<u>EQ 0.1% BASE</u>	<u>N18861</u>	<u>001</u>	Aug 31, 1983
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<u>AB</u> TARO	<u>EQ 0.1% BASE</u>	<u>N70062</u>	<u>001</u>	May 14, 1985
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BETA-VAL

<u>AB</u> TEVA	<u>EQ 0.1% BASE</u>	<u>N18642</u>	<u>001</u>	Mar 24, 1983
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DERMABET

<u>AB</u> TARO	<u>EQ 0.1% BASE</u>	<u>N72041</u>	<u>001</u>	Jan 06, 1988
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VALNAC

<u>AB</u> ACTAVIS MID ATLANTIC	<u>EQ 0.1% BASE</u>	<u>N70050</u>	<u>001</u>	Oct 10, 1984
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LOTION; TOPICAL

BETAMETHASONE VALERATE

<u>AB</u> ACTAVIS MID ATLANTIC	<u>EQ 0.1% BASE</u>	<u>N70052</u>	<u>001</u>	Jul 31, 1985
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<u>AB</u> + FOUGERA	<u>EQ 0.1% BASE</u>	<u>N18866</u>	<u>001</u>	Aug 31, 1983
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<u>AB</u> TEVA PHARMS	<u>EQ 0.1% BASE</u>	<u>N71883</u>	<u>001</u>	Apr 22, 1988
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BETA-VAL

<u>AB</u> TEVA	<u>EQ 0.1% BASE</u>	<u>N70072</u>	<u>001</u>	Jun 27, 1985
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OINTMENT; TOPICAL

BETAMETHASONE VALERATE

<u>AB</u> ACTAVIS MID ATLANTIC	<u>EQ 0.1% BASE</u>	<u>N70051</u>	<u>001</u>	Oct 10, 1984
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<u>AB</u> + FOUGERA	<u>EQ 0.1% BASE</u>	<u>N18865</u>	<u>001</u>	Aug 31, 1983
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BETA-VAL

<u>AB</u> TEVA	<u>EQ 0.1% BASE</u>	<u>N70069</u>	<u>001</u>	Dec 19, 1985
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BETAXOLOL HYDROCHLORIDE

SOLUTION/DROPS; OPHTHALMIC

BETAXOLOL

<u>AT</u> AKORN	<u>EQ 0.5% BASE</u>	<u>N75386</u>	<u>001</u>	Jun 30, 2000
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<u>AT</u> NOVEX	<u>EQ 0.5% BASE</u>	<u>N75446</u>	<u>001</u>	Sep 28, 2000
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BETAXOLOL HYDROCHLORIDE

<u>AT</u> BAUSCH AND LOMB	<u>EQ 0.5% BASE</u>	<u>N75630</u>	<u>001</u>	Apr 12, 2001
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BETOPTIC

<u>AT</u> + ALCON	<u>EQ 0.5% BASE</u>	<u>N19270</u>	<u>001</u>	Aug 30, 1985
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27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 49 (of 370)

BETAXOLOL HYDROCHLORIDE

SUSPENSION/DROPS; OPHTHALMIC

BETOPTIC S

+ ALCON

EQ 0.25% BASE

N19845 001

Dec 29, 1989

TABLET; ORAL

BETAXOLOL HYDROCHLORIDEAB ACTAVIS TOTOWA10MGN75541 001

Oct 22, 1999

AB20MGN75541 002

Oct 22, 1999

KERLONEAB SANOFI AVENTIS US10MGN19507 001

Oct 27, 1989

AB +20MGN19507 002

Oct 27, 1989

BETHANECHOL CHLORIDE

TABLET; ORAL

BETHANECHOL CHLORIDEAA ACTAVIS TOTOWA5MGN40552 001

Oct 28, 2004

AA10MGN40553 001

Oct 28, 2004

AA25MGN40554 001

Oct 28, 2004

AA50MGN40551 001

Oct 28, 2004

AA IMPAX LABS5MGN40739 001

Nov 01, 2006

AA10MGN40741 001

Nov 01, 2006

AA25MGN40740 001

Nov 01, 2006

AA50MGN40721 004

Nov 01, 2006

AA UPSHER SMITH5MGN40633 001

Jun 01, 2005

AA10MGN40634 001

Jun 01, 2005

AA25MGN40635 001

Jun 01, 2005

AA50MGN40636 001

Jun 01, 2005

AA WOCKHARDT5MGN40532 001

Sep 29, 2003

AA10MGN40533 001

Sep 29, 2003

AA25MGN40534 001

Sep 29, 2003

AA50MGN40518 001

Sep 29, 2003

DUVOIDAA WELLSRING PHARM10MGN86262 001AA25MGN86263 001AA50MGN85882 003URECHOLINEAA + ODYSSEY PHARMS5MGN89095 001

Dec 19, 1985

AA +10MGN88440 001

May 29, 1984

AA +25MGN88441 001

May 29, 1984

AA +50MGN89096 001

Dec 19, 1985

BEXAROTENE

CAPSULE; ORAL

TARGRETIN

+ EISAI MEDCL RES

75MG

N21055 001

Dec 29, 1999

GEL; TOPICAL

TARGRETIN

+ EISAI MEDCL RES

1%

N21056 001

Jun 28, 2000

BICALUTAMIDE

TABLET; ORAL

CASODEX

+ ASTRAZENECA

50MG

N20498 001

Oct 04, 1995

BIMATOPROST

SOLUTION/DROPS; OPHTHALMIC

+ ALLERGAN

0.03%

N21275 001

Mar 16, 2001

PRESCRIPTION DRUG PRODUCT LIST

3 - 50 (of 370)

BIPERIDEN HYDROCHLORIDE

TABLET; ORAL
AKINETON

+ ABBOTT 2MG N12003 001

BISACODYL; POLYETHYLENE GLYCOL 3350; POTASSIUM CHLORIDE; SODIUM BICARBONATE; SODIUM CHLORIDE

TABLET, DELAYED RELEASE, FOR SOLUTION; ORAL
HALFLYTELY

+ BRAINTREE 5MG, N/A; N/A, 210GM; N/A, 0.74GM; N/A, 2.86GM N21551 001 May 10, 2004
; N/A, 5.6GM

BISKALCITRATE; METRONIDAZOLE; TETRACYCLINE

CAPSULE; ORAL
PYLERA

+ AXCAN SCANDIPHARM 140MG; 125MG; 125MG N50786 001 Sep 28, 2006

BISMUTH SUBSALICYLATE; METRONIDAZOLE; TETRACYCLINE HYDROCHLORIDE

TABLET, CHEWABLE, TABLET, CAPSULE; ORAL
HELIDAC

+ PROMETHEUS LABS 262.4MG, N/A, N/A; N/A, 250MG, N/A; N/A, N/A, 5 N50719 001 Aug 15, 1996
00MG

BISOPROLOL FUMARATE

TABLET; ORAL

BISOPROLOL FUMARATE

<u>AB</u>	AUROBINDO PHARMA	<u>5MG</u>	<u>N77910</u>	<u>001</u>	Dec 27, 2006
<u>AB</u>		<u>10MG</u>	<u>N77910</u>	<u>002</u>	Dec 27, 2006
<u>AB</u>	MUTUAL PHARM	<u>5MG</u>	<u>N75474</u>	<u>001</u>	Oct 25, 2002
<u>AB</u>		<u>10MG</u>	<u>N75474</u>	<u>002</u>	Oct 25, 2002
<u>AB</u>	MYLAN	<u>5MG</u>	<u>N75831</u>	<u>001</u>	Dec 14, 2005
<u>AB</u>		<u>10MG</u>	<u>N75831</u>	<u>002</u>	Dec 14, 2005
<u>AB</u>	SANDOZ	<u>5MG</u>	<u>N75643</u>	<u>001</u>	Nov 16, 2000
<u>AB</u>		<u>10MG</u>	<u>N75643</u>	<u>002</u>	Nov 16, 2000
<u>AB</u>	TEVA PHARMS	<u>5MG</u>	<u>N75644</u>	<u>001</u>	Jun 26, 2001
<u>AB</u>		<u>10MG</u>	<u>N75644</u>	<u>002</u>	Jun 26, 2001
	<u>ZEBETA</u>				
<u>AB</u>	DURAMED PHARMS BARR	<u>5MG</u>	<u>N19982</u>	<u>002</u>	Jul 31, 1992
<u>AB</u>	+	<u>10MG</u>	<u>N19982</u>	<u>001</u>	Jul 31, 1992

BISOPROLOL FUMARATE; HYDROCHLOROTHIAZIDE

TABLET; ORAL

BISOPROLOL FUMARATE AND HYDROCHLOROTHIAZIDE

<u>AB</u>	ACTAVIS ELIZABETH	<u>2.5MG; 6.25MG</u>	<u>N75672</u>	<u>001</u>	Sep 25, 2000
<u>AB</u>		<u>5MG; 6.25MG</u>	<u>N75672</u>	<u>002</u>	Sep 25, 2000
<u>AB</u>		<u>10MG; 6.25MG</u>	<u>N75672</u>	<u>003</u>	Sep 25, 2000
<u>AB</u>	MYLAN	<u>2.5MG; 6.25MG</u>	<u>N75768</u>	<u>001</u>	Sep 25, 2000
<u>AB</u>		<u>5MG; 6.25MG</u>	<u>N75768</u>	<u>002</u>	Sep 25, 2000
<u>AB</u>		<u>10MG; 6.25MG</u>	<u>N75768</u>	<u>003</u>	Sep 25, 2000
<u>AB</u>	SANDOZ	<u>2.5MG; 6.25MG</u>	<u>N75527</u>	<u>001</u>	Sep 25, 2000
<u>AB</u>		<u>2.5MG; 6.25MG</u>	<u>N75579</u>	<u>001</u>	Sep 25, 2000
<u>AB</u>		<u>5MG; 6.25MG</u>	<u>N75527</u>	<u>003</u>	Sep 25, 2000
<u>AB</u>		<u>5MG; 6.25MG</u>	<u>N75579</u>	<u>002</u>	Sep 25, 2000
<u>AB</u>		<u>10MG; 6.25MG</u>	<u>N75527</u>	<u>002</u>	Sep 25, 2000
<u>AB</u>		<u>10MG; 6.25MG</u>	<u>N75579</u>	<u>003</u>	Sep 25, 2000
<u>AB</u>	TEVA	<u>2.5MG; 6.25MG</u>	<u>N75686</u>	<u>001</u>	Jan 19, 2001
<u>AB</u>		<u>5MG; 6.25MG</u>	<u>N75686</u>	<u>002</u>	Jan 19, 2001
<u>AB</u>		<u>10MG; 6.25MG</u>	<u>N75686</u>	<u>003</u>	Jan 19, 2001
<u>AB</u>	WATSON LABS	<u>2.5MG; 6.25MG</u>	<u>N75469</u>	<u>001</u>	Sep 25, 2000
<u>AB</u>		<u>5MG; 6.25MG</u>	<u>N75469</u>	<u>002</u>	Sep 25, 2000

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 51 (of 370)

BISOPROLOL FUMARATE; HYDROCHLOROTHIAZIDE

TABLET; ORAL

BISOPROLOL FUMARATE AND HYDROCHLOROTHIAZIDE

<u>AB</u>	WATSON LABS	<u>10MG;6.25MG</u>	<u>N75469</u>	<u>003</u>	Sep 25, 2000
	<u>ZIAC</u>				
<u>AB</u>	DURAMED PHARMS BARR	<u>2.5MG;6.25MG</u>	<u>N20186</u>	<u>003</u>	Mar 26, 1993
<u>AB</u>		<u>5MG;6.25MG</u>	<u>N20186</u>	<u>001</u>	Mar 26, 1993
<u>AB</u>	+	<u>10MG;6.25MG</u>	<u>N20186</u>	<u>002</u>	Mar 26, 1993

BIVALIRUDININJECTABLE; INTRAVENOUS
ANGIOMAX

+	MEDICINES CO	250MG/VIAL	N20873	001	Dec 15, 2000
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BLEOMYCIN SULFATE

INJECTABLE; INJECTION

BLENOXANE

<u>AP</u>	+	BRISTOL MYERS SQUIBB	<u>EQ 15 UNITS BASE/VIAL</u>	<u>N50443</u>	<u>001</u>	
<u>AP</u>	+		<u>EQ 30 UNITS BASE/VIAL</u>	<u>N50443</u>	<u>002</u>	Sep 07, 1995

BLEOMYCIN

<u>AP</u>		BEDFORD	<u>EQ 15 UNITS BASE/VIAL</u>	<u>N65042</u>	<u>002</u>	Oct 17, 2001
<u>AP</u>			<u>EQ 30 UNITS BASE/VIAL</u>	<u>N65042</u>	<u>001</u>	Oct 17, 2001
<u>AP</u>		MAYNE PHARMA USA	<u>EQ 15 UNITS BASE/VIAL</u>	<u>N65031</u>	<u>001</u>	Mar 10, 2000
<u>AP</u>			<u>EQ 30 UNITS BASE/VIAL</u>	<u>N65031</u>	<u>002</u>	Mar 10, 2000
<u>AP</u>		SICOR PHARMS	<u>EQ 15 UNITS BASE/VIAL</u>	<u>N65033</u>	<u>001</u>	Jun 27, 2000
<u>AP</u>			<u>EQ 30 UNITS BASE/VIAL</u>	<u>N65033</u>	<u>002</u>	Jun 27, 2000

BORTEZOMIBINJECTABLE; INTRAVENOUS
VELCADE

+	MILLENNIUM PHARMS	3.5MG/VIAL	N21602	001	May 13, 2003
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BOSENTANTABLET; ORAL
TRACLEER

	ACTELION	62.5MG	N21290	001	Nov 20, 2001
+		125MG	N21290	002	Nov 20, 2001

BRETYLIUM TOSYLATE

INJECTABLE; INJECTION

BRETYLIUM TOSYLATE

<u>AP</u>	+	INTL MEDICATION	<u>50MG/ML</u>	<u>N70119</u>	<u>001</u>	Apr 29, 1986
<u>AP</u>		LUITPOLD	<u>50MG/ML</u>	<u>N70891</u>	<u>001</u>	Jul 26, 1988

BRETYLIUM TOSYLATE IN DEXTROSE 5% IN PLASTIC CONTAINER

<u>AP</u>	+	B BRAUN	<u>200MG/100ML</u>	<u>N19121</u>	<u>002</u>	Apr 29, 1986
<u>AP</u>	+		<u>400MG/100ML</u>	<u>N19121</u>	<u>003</u>	Apr 29, 1986
	+		100MG/100ML	N19121	001	Apr 29, 1986
<u>AP</u>	+	HOSPIRA	<u>200MG/100ML</u>	<u>N19008</u>	<u>002</u>	Apr 29, 1986
<u>AP</u>	+		<u>400MG/100ML</u>	<u>N19008</u>	<u>003</u>	Apr 29, 1986

BRETYLIUM TOSYLATE IN PLASTIC CONTAINER

<u>AP</u>	+	HOSPIRA	<u>50MG/ML</u>	<u>N19030</u>	<u>001</u>	Apr 29, 1986
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BRIMONIDINE TARTRATE

SOLUTION/DROPS; OPHTHALMIC

ALPHAGAN P

<u>AT</u>	+	ALLERGAN	<u>0.15%</u>	<u>N21262</u>	<u>001</u>	Mar 16, 2001
	+		0.1%	N21770	001	Aug 19, 2005

BRIMONIDINE TARTRATE

<u>AT</u>		AKORN	<u>0.2%</u>	<u>N76439</u>	<u>001</u>	Mar 14, 2006
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PRESCRIPTION DRUG PRODUCT LIST

3 - 52 (of 370)

BRIMONIDINE TARTRATE

SOLUTION/DROPS; OPHTHALMIC

BRIMONIDINE TARTRATE

<u>AT</u>	ALCON	<u>0.2%</u>	<u>N76254</u>	<u>001</u>	Sep 16, 2003
<u>AT</u>	ALCON RES	<u>0.15%</u>	<u>N21764</u>	<u>001</u>	May 22, 2006
<u>AT</u>	+ BAUSCH AND LOMB	<u>0.2%</u>	<u>N76260</u>	<u>001</u>	May 28, 2003
<u>AT</u>	IVAX PHARMS	<u>0.2%</u>	<u>N76372</u>	<u>001</u>	Sep 10, 2004

BRINZOLAMIDE

SUSPENSION/DROPS; OPHTHALMIC

AZOPT

	+ ALCON	1%	N20816	001	Apr 01, 1998
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BROMFENAC SODIUM

SOLUTION/DROPS; OPHTHALMIC

	+ ISTA PHARMS	0.09%	N21664	001	Mar 24, 2005
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BROMOCRIPTINE MESYLATE

CAPSULE; ORAL

BROMOCRIPTINE MESYLATE

<u>AB</u>	MYLAN	<u>EQ 5MG BASE</u>	<u>N77226</u>	<u>001</u>	Apr 04, 2005
<u>AB</u>	+ <u>PARLODEL</u> NOVARTIS	<u>EQ 5MG BASE</u>	<u>N17962</u>	<u>002</u>	Mar 01, 1982

TABLET; ORAL

BROMOCRIPTINE MESYLATE

<u>AB</u>	LEK PHARMS	<u>EQ 2.5MG BASE</u>	<u>N74631</u>	<u>001</u>	Jan 13, 1998
<u>AB</u>	MYLAN	<u>EQ 2.5MG BASE</u>	<u>N76962</u>	<u>001</u>	Sep 24, 2004
<u>AB</u>	+ <u>PARLODEL</u> NOVARTIS	<u>EQ 2.5MG BASE</u>	<u>N17962</u>	<u>001</u>	

BROMODIPHENHYDRAMINE HYDROCHLORIDE; CODEINE PHOSPHATE

SYRUP; ORAL

MYBANIL

	+ MORTON GROVE	12.5MG/5ML;10MG/5ML	N88626	001	Oct 12, 1984
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BROMPHENIRAMINE MALEATE

ELIXIR; ORAL

BROMPHENIRAMINE MALEATE

	+ KV PHARM	2MG/5ML	N85466	001	
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INJECTABLE; INJECTION

BROMPHENIRAMINE MALEATE

	+ WATSON LABS	10MG/ML	N83821	001	
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BROMPHENIRAMINE MALEATE; DEXTROMETHORPHAN HYDROBROMIDE; PSEUDOEPHEDRINE HYDROCHLORIDE

SYRUP; ORAL

BROMFED-DM

<u>AA</u>	BRIGHTON PHARMS INC	<u>2MG/5ML;10MG/5ML;30MG/5ML</u>	<u>N89681</u>	<u>001</u>	Dec 22, 1988
<u>AA</u>	+ <u>MYPHETANE DX</u> MORTON GROVE	<u>2MG/5ML;10MG/5ML;30MG/5ML</u>	<u>N88811</u>	<u>001</u>	Jun 07, 1985

BUDESONIDE

CAPSULE; ORAL

ENTOCORT EC

	+ ASTRAZENECA	3MG	N21324	001	Oct 02, 2001
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POWDER, METERED; INHALATION

BUDESONIDE

	ASTRAZENECA	0.08MG/INH	N21949	001	Jul 12, 2006
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PRESCRIPTION DRUG PRODUCT LIST

3 - 53 (of 370)

BUDESONIDE

POWDER, METERED; INHALATION

BUDESONIDE

+ ASTRAZENECA	0.16MG/INH	N21949	002	Jul 12, 2006
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PULMICORT

+ ASTRAZENECA	0.16MG/INH	N20441	002	Jun 24, 1997
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SPRAY, METERED; NASAL

RHINOCORT

+ ASTRAZENECA	0.032MG/INH	N20746	001	Oct 01, 1999
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SUSPENSION; INHALATION

PULMICORT RESPULES

ASTRAZENECA	0.25MG/2ML	N20929	001	Aug 08, 2000
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+	0.5MG/2ML	N20929	002	Aug 08, 2000
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BUDESONIDE; FORMOTEROL FUMARATE

SPRAY, METERED; INHALATION

SYMBICORT

ASTRAZENECA	0.08MG/INH;EQ 0.045MG BASE	N21929	001	Jul 21, 2006
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+	0.016MG/INH;EQ 0.045MG BASE	N21929	002	Jul 21, 2006
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BUMETANIDE

INJECTABLE; INJECTION

BUMETANIDE

<u>AP</u> + BEDFORD	<u>0.25MG/ML</u>	<u>N74441</u>	<u>001</u>	Jan 27, 1995
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<u>AP</u> HOSPIRA	<u>0.25MG/ML</u>	<u>N74160</u>	<u>001</u>	Oct 30, 1997
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<u>AP</u>	<u>0.25MG/ML</u>	<u>N74332</u>	<u>001</u>	Oct 31, 1994
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<u>AP</u> SICOR PHARMS	<u>0.25MG/ML</u>	<u>N74613</u>	<u>001</u>	Nov 18, 1997
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TABLET; ORAL

BUMETANIDE

<u>AB</u> IVAX PHARMS	<u>0.5MG</u>	<u>N74225</u>	<u>001</u>	Apr 24, 1995
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<u>AB</u>	<u>1MG</u>	<u>N74225</u>	<u>002</u>	Apr 24, 1995
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<u>AB</u>	<u>2MG</u>	<u>N74225</u>	<u>003</u>	Apr 24, 1995
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<u>AB</u> SANDOZ	<u>0.5MG</u>	<u>N74700</u>	<u>001</u>	Nov 21, 1996
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<u>AB</u>	<u>1MG</u>	<u>N74700</u>	<u>002</u>	Nov 21, 1996
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<u>AB</u>	<u>2MG</u>	<u>N74700</u>	<u>003</u>	Nov 21, 1996
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BUMEX

<u>AB</u> ROCHE	<u>0.5MG</u>	<u>N18225</u>	<u>002</u>	Feb 28, 1983
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<u>AB</u>	<u>1MG</u>	<u>N18225</u>	<u>001</u>	Feb 28, 1983
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<u>AB</u> +	<u>2MG</u>	<u>N18225</u>	<u>003</u>	Jun 14, 1985
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BUPIVACAINE HYDROCHLORIDE

INJECTABLE; INJECTION

BUPIVACAINE HYDROCHLORIDE

<u>AP</u> HOSPIRA	<u>0.25%</u>	<u>N18053</u>	<u>002</u>	
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<u>AP</u>	<u>0.25%</u>	<u>N70583</u>	<u>001</u>	Feb 17, 1987
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<u>AP</u>	<u>0.25%</u>	<u>N70586</u>	<u>001</u>	Mar 03, 1987
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<u>AP</u>	<u>0.25%</u>	<u>N70590</u>	<u>001</u>	Feb 17, 1987
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<u>AP</u>	<u>0.5%</u>	<u>N18053</u>	<u>001</u>	
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<u>AP</u>	<u>0.5%</u>	<u>N70584</u>	<u>001</u>	Feb 17, 1986
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<u>AP</u>	<u>0.5%</u>	<u>N70597</u>	<u>001</u>	Mar 03, 1987
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<u>AP</u>	<u>0.5%</u>	<u>N70609</u>	<u>001</u>	Mar 03, 1987
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<u>AP</u>	<u>0.75%</u>	<u>N18053</u>	<u>003</u>	
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<u>AP</u>	<u>0.75%</u>	<u>N70585</u>	<u>001</u>	Mar 03, 1987
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<u>AP</u>	<u>0.75%</u>	<u>N70587</u>	<u>001</u>	Mar 03, 1987
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BUPIVACAINE HYDROCHLORIDE PRESERVATIVE FREE

<u>AP</u> INTL MEDICATED	<u>0.25%</u>	<u>N76012</u>	<u>001</u>	Jan 09, 2002
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<u>AP</u>	<u>0.5%</u>	<u>N76012</u>	<u>002</u>	Jan 09, 2002
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<u>AP</u>	<u>0.75%</u>	<u>N76012</u>	<u>003</u>	Jan 09, 2002
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27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 54 (of 370)

BUPIVACAINE HYDROCHLORIDE

INJECTABLE; INJECTION

MARCAINE HYDROCHLORIDE

<u>AP</u>	+	HOSPIRA	<u>0.25%</u>	<u>N16964</u>	<u>001</u>	
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<u>AP</u>	+		<u>0.5%</u>	<u>N16964</u>	<u>006</u>	
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MARCAINE HYDROCHLORIDE PRESERVATIVE FREE

<u>AP</u>	+	HOSPIRA	<u>0.25%</u>	<u>N16964</u>	<u>012</u>	
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<u>AP</u>	+		<u>0.5%</u>	<u>N16964</u>	<u>005</u>	
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<u>AP</u>	+		<u>0.75%</u>	<u>N16964</u>	<u>009</u>	
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SENSORCAINE

<u>AP</u>		ASTRAZENECA	<u>0.25%</u>	<u>N18304</u>	<u>001</u>	
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<u>AP</u>			<u>0.25%</u>	<u>N70552</u>	<u>001</u>	May 21, 1986
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<u>AP</u>			<u>0.5%</u>	<u>N18304</u>	<u>002</u>	
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<u>AP</u>			<u>0.5%</u>	<u>N70553</u>	<u>001</u>	May 21, 1986
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<u>AP</u>			<u>0.75%</u>	<u>N18304</u>	<u>003</u>	
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<u>AP</u>			<u>0.75%</u>	<u>N70554</u>	<u>001</u>	May 21, 1986
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INJECTABLE; SPINAL

BUPIVACAINE

<u>AP</u>		HOSPIRA	<u>0.75%</u>	<u>N71810</u>	<u>001</u>	Dec 11, 1987
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MARCAINE

<u>AP</u>	+	HOSPIRA	<u>0.75%</u>	<u>N18692</u>	<u>001</u>	May 04, 1984
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SENSORCAINE

<u>AP</u>		ASTRAZENECA	<u>0.75%</u>	<u>N71202</u>	<u>001</u>	Apr 15, 1987
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BUPIVACAINE HYDROCHLORIDE; EPINEPHRINE

INJECTABLE; INJECTION

BUPIVACAINE HYDROCHLORIDE AND EPINEPHRINE

+	HOSPIRA	0.25%;0.005MG/ML	N71165	001	Jun 16, 1988
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		0.25%;0.005MG/ML	N71166	001	Jun 16, 1988
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		0.25%;0.005MG/ML	N71167	001	Jun 16, 1988
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+		0.5%;0.005MG/ML	N71168	001	Jun 16, 1988
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		0.5%;0.005MG/ML	N71169	001	Jun 16, 1988
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		0.5%;0.005MG/ML	N71170	001	Jun 16, 1988
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+		0.75%;0.005MG/ML	N71171	001	Jun 16, 1988
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BUPIVACAINE HYDROCHLORIDE; EPINEPHRINE BITARTRATE

INJECTABLE; INJECTION

BUPIVACAINE HYDROCHLORIDE AND EPINEPHRINE

<u>AP</u>		SEPTODONT	<u>0.5%;0.0091MG/ML</u>	<u>N77250</u>	<u>001</u>	Sep 27, 2006
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BUPIVACAINE HYDROCHLORIDE W/EPINEPHRINE

<u>AP</u>	+	HOSPIRA	<u>0.5%;0.0091MG/ML</u>	<u>N22046</u>	<u>001</u>	Jul 13, 1983
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MARCAINE HYDROCHLORIDE W/ EPINEPHRINE

<u>AP</u>	+	HOSPIRA	<u>0.25%;0.0091MG/ML</u>	<u>N16964</u>	<u>004</u>	
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<u>AP</u>	+		<u>0.5%;0.0091MG/ML</u>	<u>N16964</u>	<u>008</u>	
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MARCAINE HYDROCHLORIDE W/ EPINEPHRINE PRESERVATIVE FREE

<u>AP</u>	+	HOSPIRA	<u>0.25%;0.0091MG/ML</u>	<u>N16964</u>	<u>013</u>	
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<u>AP</u>	+		<u>0.5%;0.0091MG/ML</u>	<u>N16964</u>	<u>007</u>	
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<u>AP</u>	+		<u>0.75%;0.0091MG/ML</u>	<u>N16964</u>	<u>010</u>	
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SENSORCAINE

<u>AP</u>		ASTRAZENECA	<u>0.25%;0.0091MG/ML</u>	<u>N70966</u>	<u>001</u>	Oct 13, 1987
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<u>AP</u>			<u>0.25%;0.0091MG/ML</u>	<u>N70967</u>	<u>001</u>	Oct 13, 1987
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<u>AP</u>			<u>0.5%;0.0091MG/ML</u>	<u>N18304</u>	<u>004</u>	Sep 02, 1983
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<u>AP</u>			<u>0.5%;0.0091MG/ML</u>	<u>N70968</u>	<u>001</u>	Oct 13, 1987
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<u>AP</u>			<u>0.75%;0.0091MG/ML</u>	<u>N18304</u>	<u>005</u>	Sep 02, 1983
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BUPIVACAINE HYDROCHLORIDE; LIDOCAINE HYDROCHLORIDE

INJECTABLE; INJECTION

DUOCAINE

+	AMPHASTAR	EQ 0.375% (37.5MG/10ML);EQ 1% (100MG/10ML)	N21496	001	May 23, 2003
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27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 55 (of 370)

BUPRENORPHINE HYDROCHLORIDE

INJECTABLE; INJECTION

BUPRENEXAP + RECKITT BENCKISER EQ 0.3MG BASE/MLN18401 001BUPRENORPHINE HYDROCHLORIDEAP BEDFORD EQ 0.3MG BASE/MLN76931 001

Mar 02, 2005

AP HOSPIRA EQ 0.3MG BASE/MLN74137 001

Jun 03, 1996

TABLET; SUBLINGUAL

SUBUTEX

RECKITT BENCKISER EQ 2MG BASE

N20732 002

Oct 08, 2002

+ EQ 8MG BASE

N20732 003

Oct 08, 2002

BUPRENORPHINE HYDROCHLORIDE; NALOXONE HYDROCHLORIDE

TABLET; SUBLINGUAL

SUBOXONE

RECKITT BENCKISER 2MG;0.5MG

N20733 001

Oct 08, 2002

+ 8MG;2MG

N20733 002

Oct 08, 2002

BUPROPION HYDROCHLORIDE

TABLET; ORAL

BUPROPION HYDROCHLORIDEAB APOTEX INC 75MGN76143 001

Jan 17, 2006

AB 100MGN76143 002

Jan 17, 2006

AB MYLAN 75MGN75491 001

Apr 17, 2000

AB 100MGN75491 002

Apr 17, 2000

AB SANDOZ 75MGN75584 001

Feb 07, 2000

AB 75MGN75613 002

Oct 10, 2000

AB 100MGN75584 002

Feb 07, 2000

AB 100MGN75613 001

Oct 10, 2000

AB TEVA 75MGN75310 001

Nov 29, 1999

AB 100MGN75310 002

Nov 29, 1999

WELLBUTRINAB GLAXOSMITHKLINE 75MGN18644 002

Dec 30, 1985

AB + 100MGN18644 003

Dec 30, 1985

TABLET, EXTENDED RELEASE; ORAL

BUPROPION HYDROCHLORIDEAB3 ANCHEN PHARMS 150MGN77284 001

Dec 14, 2006

AB3 300MGN77284 002

Dec 14, 2006

AB1 IMPAX LABS 100MGN75913 001

Jan 28, 2004

AB1 150MGN75913 002

Mar 22, 2004

AB1 200MGN76711 001

Dec 03, 2004

AB2 150MGN75914 001

May 27, 2004

AB3 300MGN77415 002

Dec 15, 2006

AB1 SANDOZ 100MGN75932 001

Nov 25, 2003

AB1 100MGN76845 001

Jul 14, 2005

AB1 150MGN75932 002

Mar 22, 2004

AB1 150MGN76845 002

Jul 14, 2005

AB1 200MGN75932 003

Jun 22, 2005

AB2 150MGN76834 001

Jul 14, 2005

WELLBUTRIN SRAB1 GLAXOSMITHKLINE 100MGN20358 002

Oct 04, 1996

AB1 + 150MGN20358 003

Oct 04, 1996

AB1 200MGN20358 004

Jun 14, 2002

50MG

N20358 001

Oct 04, 1996

WELLBUTRIN XLAB3 + SMITHKLINE BEECHAM 150MGN21515 001

Aug 28, 2003

AB3 300MGN21515 002

Aug 28, 2003

ZYBANAB2 + GLAXOSMITHKLINE 150MGN20711 003

May 14, 1997

PRESCRIPTION DRUG PRODUCT LIST

3 - 56 (of 370)

BUSPIRONE HYDROCHLORIDE

TABLET; ORAL

BUSPAR

<u>AB</u>	BRISTOL MYERS SQUIBB	<u>5MG</u>	<u>N18731</u>	<u>001</u>	Sep 29, 1986
<u>AB</u>		<u>10MG</u>	<u>N18731</u>	<u>002</u>	Sep 29, 1986
<u>AB</u>		<u>15MG</u>	<u>N18731</u>	<u>003</u>	Apr 22, 1996
<u>AB</u>	+	<u>30MG</u>	<u>N18731</u>	<u>004</u>	Apr 22, 1996
<u>BUSPIRONE HYDROCHLORIDE</u>					
<u>AB</u>	ACTAVIS TOTOWA	<u>5MG</u>	<u>N75388</u>	<u>001</u>	May 09, 2002
<u>AB</u>		<u>10MG</u>	<u>N75388</u>	<u>002</u>	May 09, 2002
<u>AB</u>		<u>15MG</u>	<u>N75388</u>	<u>003</u>	May 09, 2002
<u>AB</u>	EGIS	<u>5MG</u>	<u>N75119</u>	<u>001</u>	Mar 14, 2002
<u>AB</u>		<u>10MG</u>	<u>N75119</u>	<u>002</u>	Mar 14, 2002
<u>AB</u>		<u>15MG</u>	<u>N75119</u>	<u>003</u>	Jan 23, 2003
<u>AB</u>	KV PHARM	<u>5MG</u>	<u>N75572</u>	<u>001</u>	Feb 27, 2002
<u>AB</u>		<u>10MG</u>	<u>N75572</u>	<u>002</u>	Feb 27, 2002
<u>AB</u>		<u>15MG</u>	<u>N75572</u>	<u>003</u>	Feb 27, 2002
<u>AB</u>	MYLAN	<u>5MG</u>	<u>N75272</u>	<u>001</u>	Mar 01, 2002
<u>AB</u>		<u>10MG</u>	<u>N75272</u>	<u>002</u>	Mar 01, 2002
<u>AB</u>		<u>15MG</u>	<u>N75272</u>	<u>003</u>	Mar 28, 2001
<u>AB</u>		<u>30MG</u>	<u>N76008</u>	<u>001</u>	Jun 28, 2001
<u>AB</u>	PAR PHARM	<u>5MG</u>	<u>N75467</u>	<u>001</u>	Feb 28, 2002
<u>AB</u>		<u>7.5MG</u>	<u>N75467</u>	<u>002</u>	Mar 28, 2001
<u>AB</u>		<u>10MG</u>	<u>N75467</u>	<u>003</u>	Feb 28, 2002
<u>AB</u>		<u>15MG</u>	<u>N75467</u>	<u>004</u>	Feb 28, 2002
<u>AB</u>	SANDOZ	<u>5MG</u>	<u>N75413</u>	<u>001</u>	Mar 19, 2002
<u>AB</u>		<u>10MG</u>	<u>N75413</u>	<u>002</u>	Mar 19, 2002
<u>AB</u>		<u>15MG</u>	<u>N75413</u>	<u>003</u>	Mar 19, 2002
<u>AB</u>	TEVA	<u>5MG</u>	<u>N75022</u>	<u>001</u>	Feb 28, 2002
<u>AB</u>		<u>10MG</u>	<u>N75022</u>	<u>002</u>	Feb 28, 2002
<u>AB</u>		<u>15MG</u>	<u>N75022</u>	<u>003</u>	Feb 28, 2002
<u>AB</u>		<u>30MG</u>	<u>N75022</u>	<u>004</u>	Mar 25, 2004
<u>AB</u>	TORPHARM	<u>5MG</u>	<u>N75521</u>	<u>001</u>	Apr 05, 2002
<u>AB</u>		<u>10MG</u>	<u>N75521</u>	<u>002</u>	Apr 05, 2002
<u>AB</u>		<u>15MG</u>	<u>N75521</u>	<u>003</u>	Apr 05, 2002
<u>AB</u>	WATSON LABS	<u>5MG</u>	<u>N74253</u>	<u>001</u>	Mar 28, 2001
<u>AB</u>		<u>10MG</u>	<u>N74253</u>	<u>002</u>	Mar 28, 2001
<u>AB</u>		<u>15MG</u>	<u>N74253</u>	<u>003</u>	Mar 13, 2002
<u>AB</u>	ZENITH GOLDLINE	<u>5MG</u>	<u>N75385</u>	<u>001</u>	Mar 01, 2002
<u>AB</u>		<u>10MG</u>	<u>N75385</u>	<u>002</u>	Mar 01, 2002
<u>AB</u>		<u>15MG</u>	<u>N75385</u>	<u>003</u>	Mar 01, 2002

BUSULFAN

INJECTABLE; INJECTION

BUSULFEX

+ PDL BIOPHARMA INC 6MG/ML N20954 001 Feb 04, 1999

TABLET; ORAL

MYLERAN

+ GLAXOSMITHKLINE 2MG N09386 001

BUTABARBITAL SODIUM

ELIXIR; ORAL

BUTISOL SODIUM

+ MEDPOINTE PHARM HLC 30MG/5ML N85380 001

TABLET; ORAL

BUTABARBITAL

AA BUNDY 30MG N85550 001

BUTISOL SODIUM

AA + MEDPOINTE PHARM HLC 30MG N00793 004

PRESCRIPTION DRUG PRODUCT LIST

3 - 57 (of 370)

BUTABARBITAL SODIUM

TABLET; ORAL

BUTISOL SODIUM

+	MEDPOINTE PHARM HLC	50MG	N00793	003	
	SODIUM BUTABARBITAL				
	MARSHALL PHARMA	16.2MG	N83524	001	
		32.4MG	N83858	001	

BUTENAFINE HYDROCHLORIDE

CREAM; TOPICAL

MENTAX

+	MYLAN BERTEK	1%	N20524	001	Oct 18, 1996
	MENTAX-TC				
+	MYLAN BERTEK	1%	N21408	001	Oct 17, 2002

BUTOCONAZOLE NITRATE

CREAM; VAGINAL

GYNAZOLE-1

+	KV PHARM	2%	N19881	001	Feb 07, 1997
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BUTORPHANOL TARTRATE

INJECTABLE; INJECTION

BUTORPHANOL TARTRATE

<u>AP</u>	APOTEX INC	<u>2MG/ML</u>	<u>N75697</u>	<u>001</u>	Oct 23, 2001
<u>AP</u>	BEDFORD	<u>2MG/ML</u>	<u>N75046</u>	<u>001</u>	Aug 12, 1998
<u>AP</u>	HOSPIRA	<u>1MG/ML</u>	<u>N75559</u>	<u>001</u>	Mar 20, 2000
<u>AP</u>		<u>2MG/ML</u>	<u>N75559</u>	<u>002</u>	Mar 20, 2000
<u>AP</u>	MAYNE PHARMA USA	<u>1MG/ML</u>	<u>N75342</u>	<u>001</u>	Nov 04, 1999
<u>AP</u>		<u>2MG/ML</u>	<u>N75342</u>	<u>002</u>	Nov 04, 1999

BUTORPHANOL TARTRATE PRESERVATIVE FREE

<u>AP</u>	APOTEX INC	<u>1MG/ML</u>	<u>N75695</u>	<u>001</u>	Oct 23, 2001
<u>AP</u>		<u>2MG/ML</u>	<u>N75695</u>	<u>002</u>	Oct 23, 2001
<u>AP</u>	BEDFORD	<u>1MG/ML</u>	<u>N75045</u>	<u>001</u>	Aug 12, 1998
<u>AP</u>		<u>2MG/ML</u>	<u>N75045</u>	<u>002</u>	Aug 12, 1998
<u>AP</u>	HOSPIRA	<u>1MG/ML</u>	<u>N74620</u>	<u>001</u>	Jan 22, 1997
<u>AP</u>		<u>1MG/ML</u>	<u>N74626</u>	<u>001</u>	Jan 23, 1997
<u>AP</u>		<u>2MG/ML</u>	<u>N74620</u>	<u>002</u>	Jan 22, 1997
<u>AP</u>		<u>2MG/ML</u>	<u>N74626</u>	<u>002</u>	Jan 23, 1997
<u>AP</u>	MAYNE PHARMA USA	<u>1MG/ML</u>	<u>N75170</u>	<u>001</u>	Sep 28, 1998
<u>AP</u>		<u>2MG/ML</u>	<u>N75170</u>	<u>002</u>	Sep 28, 1998

STADOL

<u>AP</u>	+	APOTHECON	<u>2MG/ML</u>	<u>N17857</u>	<u>004</u>
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STADOL PRESERVATIVE FREE

<u>AP</u>	+	APOTHECON	<u>1MG/ML</u>	<u>N17857</u>	<u>001</u>
<u>AP</u>	+		<u>2MG/ML</u>	<u>N17857</u>	<u>002</u>

SPRAY, METERED; NASAL

BUTORPHANOL TARTRATE

<u>AB</u>	+	MYLAN	<u>1MG/SPRAY</u>	<u>N75759</u>	<u>001</u>	Aug 08, 2001
<u>AB</u>		NOVEX	<u>1MG/SPRAY</u>	<u>N75499</u>	<u>001</u>	Dec 04, 2002
<u>AB</u>		ROXANE	<u>1MG/SPRAY</u>	<u>N75824</u>	<u>001</u>	Mar 12, 2002

CABERGOLINE

TABLET; ORAL

CABERGOLINE

<u>AB</u>		PAR PHARM	<u>0.5MG</u>	<u>N76310</u>	<u>001</u>	Dec 29, 2005
		<u>DOSTINEX</u>				
<u>AB</u>	+	PHARMACIA AND UPJOHN	<u>0.5MG</u>	<u>N20664</u>	<u>001</u>	Dec 23, 1996

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 58 (of 370)

CAFFEINE CITRATE

SOLUTION; INTRAVENOUS

CAFCIT

<u>AP</u>	+	MEAD JOHNSON	<u>EQ 30MG BASE/3ML (EQ 10MG BASE/ML)</u>	<u>N20793</u>	<u>001</u>	Sep 21, 1999
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CAFFEINE CITRATE

<u>AP</u>		PHARMAFORCE	<u>EQ 30MG BASE/3ML (EQ 10MG BASE/ML)</u>	<u>N77233</u>	<u>001</u>	Sep 21, 2006
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SOLUTION; ORAL

CAFCIT

<u>AA</u>	+	MEAD JOHNSON	<u>EQ 30MG BASE/3ML (EQ 10MG BASE/ML)</u>	<u>N20793</u>	<u>002</u>	Apr 12, 2000
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CAFFEINE CITRATE

<u>AA</u>		PHARMAFORCE	<u>EQ 30MG BASE/3ML (EQ 10MG BASE/ML)</u>	<u>N77304</u>	<u>001</u>	Sep 21, 2006
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CAFFEINE; ERGOTAMINE TARTRATE

SUPPOSITORY; RECTAL

MIGERGOT

+	G AND W LABS	100MG;2MG		N86557	001	Oct 04, 1983
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TABLET; ORAL

CAFERGOT

<u>AA</u>	+	SANDOZ	<u>100MG;1MG</u>	<u>N84294</u>	<u>001</u>	
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ERGOTAMINE TARTRATE AND CAFFEINE

<u>AA</u>		MIKART	<u>100MG;1MG</u>	<u>N40590</u>	<u>001</u>	Sep 16, 2005
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<u>AA</u>		WEST WARD	<u>100MG;1MG</u>	<u>N40510</u>	<u>001</u>	Sep 17, 2004
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CALCIPOTRIENE

CREAM; TOPICAL

DOVONEX

+	LEO PHARM	0.005%		N20554	001	Jul 22, 1996
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OINTMENT; TOPICAL

DOVONEX

+	LEO PHARM	0.005%		N20273	001	Dec 29, 1993
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SOLUTION; TOPICAL

DOVONEX

+	LEO PHARM	0.005%		N20611	001	Mar 03, 1997
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CALCITONIN SALMON RECOMBINANT

SPRAY, METERED; NASAL

FORTICAL

+	UPSHER SMITH	200 IU/SPRAY		N21406	001	Aug 12, 2005
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CALCITONIN, SALMON

INJECTABLE; INJECTION

MIACALCIN

+	NOVARTIS	200 IU/ML		N17808	002	Mar 29, 1991
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SPRAY, METERED; NASAL

MIACALCIN

+	NOVARTIS	200 IU/SPRAY		N20313	002	Aug 17, 1995
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CALCITRIOL

CAPSULE; ORAL

CALCITRIOL

<u>AB</u>		ROXANE	<u>0.25UGM</u>	<u>N76917</u>	<u>001</u>	Mar 27, 2006
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<u>AB</u>		TEVA	<u>0.25UGM</u>	<u>N75765</u>	<u>001</u>	Oct 12, 2001
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<u>AB</u>			<u>0.5UGM</u>	<u>N75765</u>	<u>002</u>	Oct 12, 2001
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ROCALTRIL

<u>AB</u>		ROCHE	<u>0.25UGM</u>	<u>N18044</u>	<u>001</u>	
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<u>AB</u>	+		<u>0.5UGM</u>	<u>N18044</u>	<u>002</u>	
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27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 59 (of 370)

CALCITRIOL

INJECTABLE; INJECTION

CALCIJEX

<u>AP</u>	+	ABBOTT	<u>0.001MG/ML</u>	<u>N18874</u>	<u>001</u>	Sep 25, 1986
<u>AP</u>	+		<u>0.002MG/ML</u>	<u>N18874</u>	<u>002</u>	Sep 25, 1986

CALCITRIOL

<u>AP</u>		ABRAXIS PHARM	<u>0.001MG/ML</u>	<u>N75836</u>	<u>001</u>	Dec 31, 2002
<u>AP</u>			<u>0.002MG/ML</u>	<u>N75836</u>	<u>002</u>	Dec 31, 2002
<u>AP</u>		FRESENIUS MEDCL	<u>0.001MG/ML</u>	<u>N75766</u>	<u>001</u>	Feb 20, 2003
<u>AP</u>			<u>0.002MG/ML</u>	<u>N75766</u>	<u>002</u>	Feb 20, 2003
<u>AP</u>		GENIX THERAP	<u>0.001MG/ML</u>	<u>N77102</u>	<u>001</u>	Feb 08, 2006
<u>AP</u>		LUITPOLD	<u>0.001MG/ML</u>	<u>N75746</u>	<u>001</u>	Sep 26, 2003
<u>AP</u>			<u>0.002MG/ML</u>	<u>N75746</u>	<u>002</u>	Sep 26, 2003
<u>AP</u>		LYNE	<u>0.001MG/ML</u>	<u>N76206</u>	<u>001</u>	Sep 17, 2003
<u>AP</u>		SICOR PHARMS	<u>0.001MG/ML</u>	<u>N75823</u>	<u>001</u>	Mar 31, 2003
<u>AP</u>			<u>0.002MG/ML</u>	<u>N75823</u>	<u>002</u>	Mar 31, 2003

SOLUTION; ORAL

CALCITRIOL

<u>AA</u>		ROXANE	<u>1UGM/ML</u>	<u>N76242</u>	<u>001</u>	Jul 18, 2003
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ROCALTROL

<u>AA</u>	+	ROCHE	<u>1UGM/ML</u>	<u>N21068</u>	<u>001</u>	Nov 20, 1998
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CALCIUM ACETATE

CAPSULE; ORAL

PHOSLO GELCAPS

+	FRESENIUS MEDCL	EQ 169MG CALCIUM	N21160	003	Apr 02, 2001
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CALCIUM CARBONATE; RISEDRONATE SODIUM

TABLET, TABLET; ORAL

ACTONEL WITH CALCIUM (COPACKAGED)

+	PROCTER AND GAMBLE	EQ 500MG BASE,N/A;N/A,35MG	N21823	001	Aug 12, 2005
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CALCIUM CHLORIDE

INJECTABLE; INJECTION

CALCIUM CHLORIDE 10% IN PLASTIC CONTAINER

+	HOSPIRA	100MG/ML	N21117	001	Jan 28, 2000
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CALCIUM CHLORIDE; DEXTROSE; GLUTATHIONE DISULFIDE; MAGNESIUM CHLORIDE; POTASSIUM CHLORIDE; SODIUM BICARBONATE; SODIUM CHLORIDE; SODIUM PHOSPHATE

SOLUTION; IRRIGATION

BSS PLUS

<u>AT</u>	+	ALCON	<u>0.154MG/ML;0.92MG/ML;0.184MG/ML;0.2MG/ML;0.38MG/ML;2.1MG/ML;7.14MG/ML;0.42MG/ML</u>	<u>N18469</u>	<u>001</u>	
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ENDOSOL EXTRA

<u>AT</u>	+	AKORN	<u>0.154MG/ML;0.92MG/ML;0.184MG/ML;0.2MG/ML;0.38MG/ML;2.1MG/ML;7.14MG/ML;0.42MG/ML</u>	<u>N20079</u>	<u>001</u>	Nov 27, 1991
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CALCIUM CHLORIDE; DEXTROSE; LACTIC ACID; MAGNESIUM CHLORIDE; POTASSIUM CHLORIDE; SODIUM BICARBONATE; SODIUM CHLORIDE

INJECTABLE; INJECTION

PRISMASOL BGK 0/2.5 IN PLASTIC CONTAINER

+	GAMBRO RENAL PRODS	3.68GM/1000ML;20GM/1000ML;5.4GM/1000ML;3.05GM/1000ML;N/A/1000ML;3.09GM/1000ML;6.46GM/1000ML	N21703	006	Oct 25, 2006
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PRISMASOL BGK 2/0 IN PLASTIC CONTAINER

+	GAMBRO RENAL PRODS	N/A/1000ML;20GM/1000ML;5.4GM/1000ML;2.03GM/1000ML;0.157GM/1000ML;3.09GM/1000ML;6.46GM/1000ML	N21703	002	Oct 25, 2006
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27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 60 (of 370)

CALCIUM CHLORIDE; DEXTROSE; LACTIC ACID; MAGNESIUM CHLORIDE; POTASSIUM CHLORIDE;
SODIUM BICARBONATE; SODIUM CHLORIDE

INJECTABLE; INJECTION

PRISMASOL BGK 2/3.5 IN PLASTIC CONTAINER

+ GAMBRO RENAL PRODS	5.15GM/1000ML;20GM/1000ML;5.4GM/1000ML; 2.03GM/1000ML;0.157GM/1000ML;3.09GM/100 0ML;6.46GM/1000ML	N21703 003	Oct 25, 2006
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PRISMASOL BGK 4/0 IN PLASTIC CONTAINER

+ GAMBRO RENAL PRODS	N/A/1000ML;20GM/1000ML;5.4GM/1000ML;3.0 5GM/1000ML;0.314GM/1000ML;3.09GM/1000ML ;6.46GM/1000ML	N21703 005	Oct 25, 2006
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PRISMASOL BGK 4/2.5 IN PLASTIC CONTAINER

+ GAMBRO RENAL PRODS	3.68GM/1000ML;20GM/1000ML;5.4GM/1000ML; 3.05GM/1000ML;0.314GM/1000ML;3.09GM/100 0ML;6.46GM/1000ML	N21703 004	Oct 25, 2006
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PRISMASOL BK 0/0 IN PLASTIC CONTAINER

+ GAMBRO RENAL PRODS	N/A/1000ML;N/A/1000ML;5.4GM/1000ML;3.05 GM/1000ML;N/A/1000ML;3.09GM/1000ML;6.46 GM/1000ML	N21703 007	Oct 25, 2006
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PRISMASOL BK 0/3.5 IN PLASTIC CONTAINER

+ GAMBRO RENAL PRODS	5.15GM/1000ML;N/A/1000ML;5.4GM/1000ML;2 .03GM/1000ML;N/A/1000ML;3.09GM/1000ML;6 .46GM/1000ML	N21703 001	Oct 25, 2006
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CALCIUM CHLORIDE; DEXTROSE; MAGNESIUM CHLORIDE; POTASSIUM CHLORIDE; SODIUM ACETATE;
SODIUM CHLORIDE

INJECTABLE; INJECTION

ISOLYTE R IN DEXTROSE 5% IN PLASTIC CONTAINER

+ B BRAUN	37MG/100ML;5GM/100ML;31MG/100ML;120MG/1 00ML;330MG/100ML;88MG/100ML	N19864 001	Jun 10, 1993
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CALCIUM CHLORIDE; DEXTROSE; MAGNESIUM CHLORIDE; POTASSIUM CHLORIDE; SODIUM ACETATE;
SODIUM CHLORIDE; SODIUM CITRATE

INJECTABLE; INJECTION

ISOLYTE E IN DEXTROSE 5% IN PLASTIC CONTAINER

+ B BRAUN	35MG/100ML;5GM/100ML;30MG/100ML;74MG/10 0ML;640MG/100ML;500MG/100ML;74MG/100ML	N19867 001	Dec 20, 1993
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CALCIUM CHLORIDE; DEXTROSE; MAGNESIUM CHLORIDE; POTASSIUM CHLORIDE; SODIUM ACETATE;
SODIUM CHLORIDE; SODIUM LACTATE

INJECTABLE; INJECTION

PLASMA-LYTE M AND DEXTROSE 5% IN PLASTIC CONTAINER

+ BAXTER HLTHCARE	37MG/100ML;5GM/100ML;30MG/100ML;119MG/1 00ML;161MG/100ML;94MG/100ML;138MG/100ML	N17390 001	
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CALCIUM CHLORIDE; DEXTROSE; MAGNESIUM CHLORIDE; SODIUM ACETATE; SODIUM CHLORIDE

SOLUTION; INTRAPERITONEAL

DIALYTE CONCENTRATE W/ DEXTROSE 30% IN PLASTIC CONTAINER

+ B BRAUN	510MG/100ML;30GM/100ML;200MG/100ML;9.4G M/100ML;11GM/100ML	N18807 003	Aug 26, 1983
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+	510MG/100ML;30GM/100ML;200MG/100ML;9.2G M/100ML;9.6GM/100ML	N18807 001	Aug 26, 1983
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DIALYTE CONCENTRATE W/ DEXTROSE 50% IN PLASTIC CONTAINER

+ B BRAUN	510MG/100ML;50GM/100ML;200MG/100ML;9.4G M/100ML;11GM/100ML	N18807 004	Aug 26, 1983
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+	510MG/100ML;50GM/100ML;200MG/100ML;9.2G M/100ML;9.6GM/100ML	N18807 002	Aug 26, 1983
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CALCIUM CHLORIDE; DEXTROSE; MAGNESIUM CHLORIDE; SODIUM CHLORIDE; SODIUM LACTATE

SOLUTION; INTRAPERITONEAL

DELFLX W/ DEXTROSE 1.5% IN PLASTIC CONTAINER

<u>AT</u>	FRESENIUS MEDCL	<u>25.7MG/100ML;1.5GM/100ML;15.2MG/100ML;5 67MG/100ML;392MG/100ML</u>	<u>N18379 002</u>
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27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 61 (of 370)

CALCIUM CHLORIDE; DEXTROSE; MAGNESIUM CHLORIDE; SODIUM CHLORIDE; SODIUM LACTATE

SOLUTION; INTRAPERITONEAL					
<u>DELFLX W/ DEXTROSE 1.5% IN PLASTIC CONTAINER</u>					
<u>AT</u>	FRESENIUS MEDCL	<u>25.7MG/100ML;1.5GM/100ML;15.2MG/100ML;5</u> <u>67MG/100ML;392MG/100ML</u>	<u>N18883</u>	<u>001</u>	Nov 30, 1984
<u>DELFLX W/ DEXTROSE 1.5% LOW MAGNESIUM IN PLASTIC CONTAINER</u>					
<u>AT</u>	FRESENIUS MEDCL	<u>25.7MG/100ML;1.5GM/100ML;5.08MG/100ML;5</u> <u>38MG/100ML;448MG/100ML</u>	<u>N18883</u>	<u>004</u>	Nov 30, 1984
<u>DELFLX W/ DEXTROSE 1.5% LOW MAGNESIUM LOW CALCIUM IN PLASTIC CONTAINER</u>					
<u>AT</u>	FRESENIUS MEDCL	<u>18.4MG/100ML;1.5GM/100ML;5.08MG/100ML;5</u> <u>38MG/100ML;448MG/100ML</u>	<u>N20171</u>	<u>001</u>	Aug 19, 1992
<u>DELFLX W/ DEXTROSE 2.5% IN PLASTIC CONTAINER</u>					
<u>AT</u>	FRESENIUS MEDCL	<u>25.7MG/100ML;2.5GM/100ML;15.2MG/100ML;5</u> <u>67MG/100ML;392MG/100ML</u>	<u>N18379</u>	<u>003</u>	
<u>AT</u>		<u>25.7MG/100ML;2.5GM/100ML;15.2MG/100ML;5</u> <u>67MG/100ML;392MG/100ML</u>	<u>N18883</u>	<u>002</u>	Nov 30, 1984
<u>DELFLX W/ DEXTROSE 2.5% LOW MAGNESIUM IN PLASTIC CONTAINER</u>					
<u>AT</u>	FRESENIUS MEDCL	<u>25.7MG/100ML;2.5GM/100ML;5.08MG/100ML;5</u> <u>38MG/100ML;448MG/100ML</u>	<u>N18883</u>	<u>005</u>	Nov 30, 1984
<u>DELFLX W/ DEXTROSE 2.5% LOW MAGNESIUM LOW CALCIUM IN PLASTIC CONTAINER</u>					
<u>AT</u>	FRESENIUS MEDCL	<u>18.4MG/100ML;2.5GM/100ML;5.08MG/100ML;5</u> <u>38MG/100ML;448MG/100ML</u>	<u>N20171</u>	<u>002</u>	Aug 19, 1992
<u>DELFLX W/ DEXTROSE 3.5% IN PLASTIC CONTAINER</u>					
<u>AT</u>	FRESENIUS MEDCL	<u>25.7MG/100ML;3.5GM/100ML;15.2MG/100ML;5</u> <u>67MG/100ML;392MG/100ML</u>	<u>N18379</u>	<u>007</u>	Jun 24, 1988
<u>DELFLX W/ DEXTROSE 4.25% IN PLASTIC CONTAINER</u>					
<u>AT</u>	FRESENIUS MEDCL	<u>25.7MG/100ML;4.25GM/100ML;15.2MG/100ML;</u> <u>567MG/100ML;392MG/100ML</u>	<u>N18379</u>	<u>001</u>	
<u>AT</u>		<u>25.7MG/100ML;4.25GM/100ML;15.2MG/100ML;</u> <u>567MG/100ML;392MG/100ML</u>	<u>N18883</u>	<u>003</u>	Nov 30, 1984
<u>DELFLX W/ DEXTROSE 4.25% LOW MAGNESIUM IN PLASTIC CONTAINER</u>					
<u>AT</u>	FRESENIUS MEDCL	<u>25.7MG/100ML;4.25GM/100ML;5.08MG/100ML;</u> <u>538MG/100ML;448MG/100ML</u>	<u>N18883</u>	<u>006</u>	Nov 30, 1984
<u>DELFLX W/ DEXTROSE 4.25% LOW MAGNESIUM LOW CALCIUM IN PLASTIC CONTAINER</u>					
<u>AT</u>	FRESENIUS MEDCL	<u>18.4MG/100ML;4.25GM/100ML;5.08MG/100ML;</u> <u>538MG/100ML;448MG/100ML</u>	<u>N20171</u>	<u>003</u>	Aug 19, 1992
<u>DELFLX-LM W/ DEXTROSE 1.5% IN PLASTIC CONTAINER</u>					
<u>AT</u>	FRESENIUS MEDCL	<u>25.7MG/100ML;1.5GM/100ML;5.08MG/100ML;5</u> <u>38MG/100ML;448MG/100ML</u>	<u>N18379</u>	<u>004</u>	Jul 07, 1982
<u>DELFLX-LM W/ DEXTROSE 2.5% IN PLASTIC CONTAINER</u>					
<u>AT</u>	FRESENIUS MEDCL	<u>25.7MG/100ML;2.5GM/100ML;5.08MG/100ML;5</u> <u>38MG/100ML;448MG/100ML</u>	<u>N18379</u>	<u>005</u>	Jul 07, 1982
<u>DELFLX-LM W/ DEXTROSE 3.5% IN PLASTIC CONTAINER</u>					
<u>AT</u>	FRESENIUS MEDCL	<u>25.7MG/100ML;3.5GM/100ML;5.08MG/100ML;5</u> <u>38MG/100ML;448MG/100ML</u>	<u>N18379</u>	<u>008</u>	Jun 24, 1988
<u>DELFLX-LM W/ DEXTROSE 4.25% IN PLASTIC CONTAINER</u>					
<u>AT</u>	FRESENIUS MEDCL	<u>25.7MG/100ML;4.25GM/100ML;5.08MG/100ML;</u> <u>538MG/100ML;448MG/100ML</u>	<u>N18379</u>	<u>006</u>	Jul 07, 1982
<u>DIALYTE LM/ DEXTROSE 1.5% IN PLASTIC CONTAINER</u>					
	B BRAUN	26MG/100ML;1.5GM/100ML;5MG/100ML;530MG/ 100ML;450MG/100ML	N18460	007	Jan 29, 1986
<u>DIALYTE LM/ DEXTROSE 2.5% IN PLASTIC CONTAINER</u>					
	B BRAUN	26MG/100ML;2.5GM/100ML;5MG/100ML;530MG/ 100ML;450MG/100ML	N18460	005	Nov 02, 1983
<u>DIALYTE LM/ DEXTROSE 4.25% IN PLASTIC CONTAINER</u>					
	B BRAUN	26MG/100ML;4.25GM/100ML;5MG/100ML;530MG /100ML;450MG/100ML	N18460	009	Jan 29, 1986
<u>DIANEAL 137 W/ DEXTROSE 1.5% IN PLASTIC CONTAINER</u>					
<u>AT</u>	BAXTER HLTHCARE	<u>25.7MG/100ML;1.5GM/100ML;15.2MG/100ML;5</u> <u>67MG/100ML;392MG/100ML</u>	<u>N17512</u>	<u>001</u>	
<u>DIANEAL 137 W/ DEXTROSE 2.5% IN PLASTIC CONTAINER</u>					
<u>AT</u>	BAXTER HLTHCARE	<u>25.7MG/100ML;2.5GM/100ML;15.2MG/100ML;5</u> <u>67MG/100ML;392MG/100ML</u>	<u>N17512</u>	<u>003</u>	

PRESCRIPTION DRUG PRODUCT LIST

3 - 62 (of 370)

CALCIUM CHLORIDE; DEXTROSE; MAGNESIUM CHLORIDE; SODIUM CHLORIDE; SODIUM LACTATE

SOLUTION; INTRAPERITONEAL

<u>DIANEAL 137 W/ DEXTROSE 4.25% IN PLASTIC CONTAINER</u>			
<u>AT</u>	BAXTER HLTHCARE	<u>25.7MG/100ML; 4.25GM/100ML; 15.2MG/100ML; 567MG/100ML; 392MG/100ML</u>	<u>N17512 002</u>
<u>DIANEAL LOW CALCIUM W/ DEXTROSE 1.5% IN PLASTIC CONTAINER</u>			
<u>AT</u>	BAXTER HLTHCARE	<u>18.3MG/100ML; 1.5GM/100ML; 5.08MG/100ML; 538MG/100ML; 448MG/100ML</u>	<u>N20183 001</u> Dec 04, 1992
<u>DIANEAL LOW CALCIUM W/ DEXTROSE 2.5% IN PLASTIC CONTAINER</u>			
	BAXTER HLTHCARE	18.3MG/100ML; 2.5GM/100ML; 5.08MG/100ML; 538MG/100ML; 448MG/100ML	N20183 002 Dec 04, 1992
<u>DIANEAL LOW CALCIUM W/ DEXTROSE 3.5% IN PLASTIC CONTAINER</u>			
	BAXTER HLTHCARE	18.3MG/100ML; 3.5GM/100ML; 5.08MG/100ML; 538MG/100ML; 448MG/100ML	N20183 003 Dec 04, 1992
<u>DIANEAL LOW CALCIUM W/ DEXTROSE 4.25% IN PLASTIC CONTAINER</u>			
	BAXTER HLTHCARE	18.3MG/100ML; 4.25GM/100ML; 5.08MG/100ML; 538MG/100ML; 448MG/100ML	N20183 004 Dec 04, 1992
<u>DIANEAL PD-1 W/ DEXTROSE 1.5% IN PLASTIC CONTAINER</u>			
<u>AT</u>	BAXTER HLTHCARE	<u>25.7MG/100ML; 1.5GM/100ML; 15.2MG/100ML; 567MG/100ML; 392MG/100ML</u>	<u>N17512 007</u> Jul 09, 1984
<u>DIANEAL PD-1 W/ DEXTROSE 2.5% IN PLASTIC CONTAINER</u>			
<u>AT</u>	BAXTER HLTHCARE	<u>25.7MG/100ML; 2.5GM/100ML; 15.2MG/100ML; 567MG/100ML; 392MG/100ML</u>	<u>N17512 008</u> Jul 09, 1984
<u>DIANEAL PD-1 W/ DEXTROSE 3.5% IN PLASTIC CONTAINER</u>			
<u>AT</u>	BAXTER HLTHCARE	<u>25.7MG/100ML; 3.5GM/100ML; 15.2MG/100ML; 567MG/100ML; 392MG/100ML</u>	<u>N17512 010</u> Nov 18, 1985
<u>DIANEAL PD-1 W/ DEXTROSE 4.25% IN PLASTIC CONTAINER</u>			
<u>AT</u>	BAXTER HLTHCARE	<u>25.7MG/100ML; 4.25GM/100ML; 15.2MG/100ML; 567MG/100ML; 392MG/100ML</u>	<u>N17512 009</u> Jul 09, 1984
<u>DIANEAL PD-2 W/ DEXTROSE 1.5% IN PLASTIC CONTAINER</u>			
<u>AT</u>	BAXTER HLTHCARE	<u>18.3MG/100ML; 1.5GM/100ML; 5.08MG/100ML; 538MG/100ML; 448MG/100ML</u>	<u>N17512 004</u>
<u>AT</u>		<u>25.7MG/100ML; 1.5GM/100ML; 5.08MG/100ML; 538MG/100ML; 448MG/100ML</u>	<u>N20163 001</u> Dec 04, 1992
<u>DIANEAL PD-2 W/ DEXTROSE 2.5% IN PLASTIC CONTAINER</u>			
<u>AT</u>	BAXTER HLTHCARE	<u>25.7MG/100ML; 2.5GM/100ML; 5.08MG/100ML; 538MG/100ML; 448MG/100ML</u>	<u>N17512 005</u>
<u>AT</u>		<u>25.7MG/100ML; 2.5GM/100ML; 5.08MG/100ML; 538MG/100ML; 448MG/100ML</u>	<u>N20163 002</u> Dec 04, 1992
<u>DIANEAL PD-2 W/ DEXTROSE 3.5% IN PLASTIC CONTAINER</u>			
<u>AT</u>	BAXTER HLTHCARE	<u>25.7MG/100ML; 3.5GM/100ML; 5.08MG/100ML; 538MG/100ML; 448MG/100ML</u>	<u>N17512 011</u> Nov 18, 1985
<u>DIANEAL PD-2 W/ DEXTROSE 4.25% IN PLASTIC CONTAINER</u>			
<u>AT</u>	BAXTER HLTHCARE	<u>25.7MG/100ML; 4.25GM/100ML; 5.08MG/100ML; 538MG/100ML; 448MG/100ML</u>	<u>N17512 006</u>
<u>AT</u>		<u>25.7MG/100ML; 4.25GM/100ML; 5.08MG/100ML; 538MG/100ML; 448MG/100ML</u>	<u>N20163 003</u> Dec 04, 1992
<u>INPERSOL-LC/LM W/ DEXTROSE 1.5% IN PLASTIC CONTAINER</u>			
<u>AT</u>	FRESENIUS	<u>18.4MG/100ML; 1.5GM/100ML; 5.08MG/100ML; 538MG/100ML; 448MG/100ML</u>	<u>N20374 001</u> Jun 13, 1994
<u>INPERSOL-LC/LM W/ DEXTROSE 2.5% IN PLASTIC CONTAINER</u>			
<u>AT</u>	FRESENIUS	<u>18.4MG/100ML; 2.5GM/100ML; 5.08MG/100ML; 538MG/100ML; 448MG/100ML</u>	<u>N20374 002</u> Jun 13, 1994
<u>INPERSOL-LC/LM W/ DEXTROSE 3.5% IN PLASTIC CONTAINER</u>			
	FRESENIUS	18.4MG/100ML; 3.5GM/100ML; 5.08MG/100ML; 538MG/100ML; 448MG/100ML	N20374 003 Jun 13, 1994
<u>INPERSOL-LC/LM W/ DEXTROSE 4.25% IN PLASTIC CONTAINER</u>			
<u>AT</u>	FRESENIUS	<u>18.4MG/100ML; 4.25GM/100ML; 5.08MG/100ML; 538MG/100ML; 448MG/100ML</u>	<u>N20374 004</u> Jun 13, 1994

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 63 (of 370)

CALCIUM CHLORIDE; DEXTROSE; MAGNESIUM SULFATE; POTASSIUM CHLORIDE; SODIUM BICARBONATE; SODIUM CHLORIDE; SODIUM PHOSPHATE, DIBASIC, HEPTAHYDRATE

INJECTABLE; INTRATHECAL
ELLIOTTS B SOLUTION
+ QOL MEDCL

0.2MG/ML;0.8MG/ML;0.3MG/ML;0.3MG/ML;1.9 MG/ML;7.3MG/ML;0.2MG/ML N20577 001 Sep 27, 1996

CALCIUM CHLORIDE; DEXTROSE; POTASSIUM CHLORIDE; SODIUM ACETATE; SODIUM CHLORIDE

INJECTABLE; INJECTION
DEXTROSE 5% IN ACETATED RINGER'S IN PLASTIC CONTAINER
+ B BRAUN

20MG/100ML;5GM/100ML;30MG/100ML;380MG/1 00ML;600MG/100ML N18258 001

CALCIUM CHLORIDE; DEXTROSE; POTASSIUM CHLORIDE; SODIUM CHLORIDE

INJECTABLE; INJECTION
DEXTROSE 5% AND RINGER'S IN PLASTIC CONTAINER

AP HOSPIRA 33MG/100ML;5GM/100ML;30MG/100ML;860MG/1 00ML N18254 001

DEXTROSE 5% IN RINGER'S IN PLASTIC CONTAINER

AP B BRAUN 33MG/100ML;5GM/100ML;30MG/100ML;860MG/1 00ML N20000 001 Apr 17, 1992

AP BAXTER HLTHCARE 33MG/100ML;5GM/100ML;30MG/100ML;860MG/1 00ML N16695 001

CALCIUM CHLORIDE; DEXTROSE; POTASSIUM CHLORIDE; SODIUM CHLORIDE; SODIUM LACTATE

INJECTABLE; INJECTION
DEXTROSE 2.5% IN HALF-STRENGTH LACTATED RINGER'S IN PLASTIC CONTAINER
B BRAUN

10MG/100ML;2.5GM/100ML;15MG/100ML;300MG /100ML;160MG/100ML N19634 001 Feb 24, 1988

DEXTROSE 5% AND LACTATED RINGER'S IN PLASTIC CONTAINER

AP HOSPIRA 20MG/100ML;5GM/100ML;30MG/100ML;600MG/1 00ML;310MG/100ML N17608 001

DEXTROSE 5% IN LACTATED RINGER'S IN PLASTIC CONTAINER

AP B BRAUN 20MG/100ML;5GM/100ML;30MG/100ML;600MG/1 00ML;310MG/100ML N19634 003 Feb 24, 1988

LACTATED RINGER'S AND DEXTROSE 5% IN PLASTIC CONTAINER

AP BAXTER HLTHCARE 20MG/100ML;5GM/100ML;30MG/100ML;600MG/1 00ML;310MG/100ML N16679 001

POTASSIUM CHLORIDE 10MEQ IN DEXTROSE 5% AND LACTATED RINGER'S IN PLASTIC CONTAINER

BAXTER HLTHCARE 20MG/100ML;5GM/100ML;105MG/100ML;600MG/ 100ML;310MG/100ML N19367 002 Apr 05, 1985

20MG/100ML;5GM/100ML;179MG/100ML;600MG/ 100ML;310MG/100ML N19367 003 Apr 05, 1985

POTASSIUM CHLORIDE 15MEQ IN DEXTROSE 5% AND LACTATED RINGER'S IN PLASTIC CONTAINER

AP BAXTER HLTHCARE 20MG/100ML;5GM/100ML;254MG/100ML;600MG/ 100ML;310MG/100ML N19367 006 Apr 05, 1985

POTASSIUM CHLORIDE 20MEQ IN DEXTROSE 5% AND LACTATED RINGER'S IN PLASTIC CONTAINER

AP BAXTER HLTHCARE 20MG/100ML;5GM/100ML;179MG/100ML;600MG/ 100ML;310MG/100ML N19367 004 Apr 05, 1985

AP 20MG/100ML;5GM/100ML;328MG/100ML;600MG/ 100ML;310MG/100ML N19367 005 Apr 05, 1985

AP HOSPIRA 20MG/100ML;5GM/100ML;179MG/100ML;600MG/ 100ML;310MG/100ML N19685 002 Oct 17, 1988

AP 20MG/100ML;5GM/100ML;328MG/100ML;600MG/ 100ML;310MG/100ML N19685 008 Oct 17, 1988

POTASSIUM CHLORIDE 30MEQ IN DEXTROSE 5% AND LACTATED RINGER'S IN PLASTIC CONTAINER

AP BAXTER HLTHCARE 20MG/100ML;5GM/100ML;254MG/100ML;600MG/ 100ML;310MG/100ML N19367 007 Apr 05, 1985

POTASSIUM CHLORIDE 40MEQ IN DEXTROSE 5% AND LACTATED RINGER'S IN PLASTIC CONTAINER

AP BAXTER HLTHCARE 20MG/100ML;5GM/100ML;328MG/100ML;600MG/ 100ML;310MG/100ML N19367 008 Apr 05, 1985

AP HOSPIRA 20MG/100ML;5GM/100ML;328MG/100ML;600MG/ 100ML;310MG/100ML N19685 004 Oct 17, 1988

PRESCRIPTION DRUG PRODUCT LIST

3 - 64 (of 370)

CALCIUM CHLORIDE; DEXTROSE; POTASSIUM CHLORIDE; SODIUM CHLORIDE; SODIUM LACTATE

INJECTABLE; INJECTION

POTASSIUM CHLORIDE 5MEQ IN DEXTROSE 5% AND LACTATED RINGER'S IN PLASTIC CONTAINER

BAXTER HLTHCARE 20MG/100ML; 5GM/100ML; 105MG/100ML; 600MG/ 100ML; 310MG/100ML N19367 001 Apr 05, 1985

CALCIUM CHLORIDE; MAGNESIUM CHLORIDE; POTASSIUM CHLORIDE; SODIUM ACETATE; SODIUM CHLORIDE

INJECTABLE; INJECTION

TPN ELECTROLYTES IN PLASTIC CONTAINER

+ HOSPIRA 16.5MG/ML; 25.4MG/ML; 74.6MG/ML; 121MG/ML; 16.1MG/ML N18895 001 Jul 20, 1984

CALCIUM CHLORIDE; MAGNESIUM CHLORIDE; POTASSIUM CHLORIDE; SODIUM ACETATE; SODIUM CHLORIDE; SODIUM CITRATE

INJECTABLE; INJECTION

ISOLYTE E IN PLASTIC CONTAINER

+ B BRAUN 35MG/100ML; 30MG/100ML; 74MG/100ML; 640MG/ 100ML; 500MG/100ML; 74MG/100ML N19718 001 Sep 29, 1989

SOLUTION; IRRIGATION

BALANCED SALT SOLUTION

AT AKORN 0.48MG/ML; 0.3MG/ML; 0.75MG/ML; 3.9MG/ML; 6.4MG/ML; 1.7MG/ML N75503 001 Sep 27, 2006

BSS

AT + ALCON 0.48MG/ML; 0.3MG/ML; 0.75MG/ML; 3.9MG/ML; 6.4MG/ML; 1.7MG/ML N20742 001 Dec 10, 1997

CALCIUM CHLORIDE; MAGNESIUM CHLORIDE; POTASSIUM CHLORIDE; SODIUM ACETATE; SODIUM CHLORIDE; SODIUM LACTATE

INJECTABLE; INJECTION

PLASMA-LYTE R IN PLASTIC CONTAINER

+ BAXTER HLTHCARE 36.8MG/100ML; 30.5MG/100ML; 74.6MG/100ML; 640MG/100ML; 496MG/100ML; 89.6MG/100ML N17438 001

CALCIUM CHLORIDE; MAGNESIUM CHLORIDE; POTASSIUM CHLORIDE; SODIUM CHLORIDE

SOLUTION; PERFUSION, CARDIAC

CARDIOPLEGIC IN PLASTIC CONTAINER

AT BAXTER HLTHCARE 17.6MG/100ML; 325.3MG/100ML; 119.3MG/100ML; 643MG/100ML N75323 001 Apr 21, 2000

PLEGISOL IN PLASTIC CONTAINER

AT + HOSPIRA 17.6MG/100ML; 325.3MG/100ML; 119.3MG/100ML; 643MG/100ML N18608 001 Feb 26, 1982

CALCIUM CHLORIDE; POTASSIUM CHLORIDE; SODIUM CHLORIDE

INJECTABLE; INJECTION

RINGER'S IN PLASTIC CONTAINER

AP B BRAUN 33MG/100ML; 30MG/100ML; 860MG/100ML N20002 001 Apr 17, 1992
AP BAXTER HLTHCARE 33MG/100ML; 30MG/100ML; 860MG/100ML N16693 001
AP HOSPIRA 33MG/100ML; 30MG/100ML; 860MG/100ML N18251 001

SOLUTION; IRRIGATION

RINGER'S IN PLASTIC CONTAINER

AT B BRAUN 33MG/100ML; 30MG/100ML; 860MG/100ML N18156 001
AT BAXTER HLTHCARE 33MG/100ML; 30MG/100ML; 860MG/100ML N18495 001 Feb 19, 1982
AT HOSPIRA 33MG/100ML; 30MG/100ML; 860MG/100ML N17635 001

CALCIUM CHLORIDE; POTASSIUM CHLORIDE; SODIUM CHLORIDE; SODIUM LACTATE

INJECTABLE; INJECTION

LACTATED RINGER'S IN PLASTIC CONTAINER

AP B BRAUN 20MG/100ML; 30MG/100ML; 600MG/100ML; 310MG/100ML N19632 001 Feb 29, 1988

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 65 (of 370)

CALCIUM CHLORIDE; POTASSIUM CHLORIDE; SODIUM CHLORIDE; SODIUM LACTATE

INJECTABLE; INJECTION

LACTATED RINGER'S IN PLASTIC CONTAINER

<u>AP</u>	BAXTER HLTHCARE	<u>20MG/100ML; 30MG/100ML; 600MG/100ML; 310MG/100ML</u>	<u>N16682</u>	<u>001</u>	
<u>AP</u>	HOSPIRA	<u>20MG/100ML; 30MG/100ML; 600MG/100ML; 310MG/100ML</u>	<u>N17641</u>	<u>001</u>	

SOLUTION; IRRIGATION

LACTATED RINGER'S IN PLASTIC CONTAINER

<u>AT</u>	B BRAUN	<u>20MG/100ML; 30MG/100ML; 600MG/100ML; 310MG/100ML</u>	<u>N18681</u>	<u>001</u>	Dec 27, 1982
<u>AT</u>	BAXTER HLTHCARE	<u>20MG/100ML; 30MG/100ML; 600MG/100ML; 310MG/100ML</u>	<u>N18494</u>	<u>001</u>	Feb 19, 1982
<u>AT</u>		<u>20MG/100ML; 30MG/100ML; 600MG/100ML; 310MG/100ML</u>	<u>N18921</u>	<u>001</u>	Apr 03, 1984
<u>AT</u>		<u>20MG/100ML; 30MG/100ML; 600MG/100ML; 310MG/100ML</u>	<u>N19933</u>	<u>001</u>	Aug 29, 1989
<u>AT</u>	HOSPIRA	<u>20MG/100ML; 30MG/100ML; 600MG/100ML; 310MG/100ML</u>	<u>N19416</u>	<u>001</u>	Jan 17, 1986

CALFACTANTSUSPENSION; INTRATRACHEAL
INFASURF PRESERVATIVE FREE

+	ONY	35MG/ML	N20521	001	Jul 01, 1998
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CANDESARTAN CILEXETIL

TABLET; ORAL

ATACAND

	ASTRAZENECA	4MG	N20838	001	Jun 04, 1998
		8MG	N20838	002	Jun 04, 1998
		16MG	N20838	003	Jun 04, 1998
+		32MG	N20838	004	Jun 04, 1998

CANDESARTAN CILEXETIL; HYDROCHLOROTHIAZIDE

TABLET; ORAL

ATACAND HCT

	ASTRAZENECA	16MG; 12.5MG	N21093	001	Sep 05, 2000
+		32MG; 12.5MG	N21093	002	Sep 05, 2000

CAPECITABINE

TABLET; ORAL

XELODA

	HLR	150MG	N20896	001	Apr 30, 1998
+		500MG	N20896	002	Apr 30, 1998

CAPREOMYCIN SULFATE

INJECTABLE; INJECTION

CAPASTAT SULFATE

+	LILLY	EQ 1GM BASE/VIAL	N50095	001	
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CAPTOPRIL

TABLET; ORAL

CAPOTEN

<u>AB</u>	PAR PHARM	<u>12.5MG</u>	<u>N18343</u>	<u>005</u>	Jan 17, 1985
<u>AB</u>		<u>25MG</u>	<u>N18343</u>	<u>002</u>	
<u>AB</u>		<u>50MG</u>	<u>N18343</u>	<u>001</u>	
<u>AB</u>	+	<u>100MG</u>	<u>N18343</u>	<u>003</u>	
	<u>CAPTOPRIL</u>				
<u>AB</u>	EGIS PHARMS	<u>12.5MG</u>	<u>N74748</u>	<u>004</u>	May 29, 1997
<u>AB</u>		<u>25MG</u>	<u>N74748</u>	<u>002</u>	May 29, 1997

PRESCRIPTION DRUG PRODUCT LIST

3 - 66 (of 370)

CAPTOPRIL

TABLET; ORAL

CAPTOPRIL

<u>AB</u>	EGIS PHARMS	<u>50MG</u>	<u>N74748</u>	<u>001</u>	May 29, 1997
<u>AB</u>		<u>100MG</u>	<u>N74748</u>	<u>003</u>	May 29, 1997
<u>AB</u>	IVAX PHARMS	<u>12.5MG</u>	<u>N74590</u>	<u>004</u>	Aug 30, 1996
<u>AB</u>		<u>25MG</u>	<u>N74590</u>	<u>002</u>	Aug 30, 1996
<u>AB</u>		<u>50MG</u>	<u>N74590</u>	<u>001</u>	Aug 30, 1996
<u>AB</u>		<u>100MG</u>	<u>N74590</u>	<u>003</u>	Aug 30, 1996
<u>AB</u>	KALI LABS	<u>12.5MG</u>	<u>N74477</u>	<u>001</u>	Feb 13, 1996
<u>AB</u>		<u>25MG</u>	<u>N74477</u>	<u>002</u>	Feb 13, 1996
<u>AB</u>		<u>50MG</u>	<u>N74477</u>	<u>003</u>	Feb 13, 1996
<u>AB</u>		<u>100MG</u>	<u>N74477</u>	<u>004</u>	Feb 13, 1996
<u>AB</u>	MYLAN	<u>12.5MG</u>	<u>N74434</u>	<u>001</u>	Feb 13, 1996
<u>AB</u>		<u>25MG</u>	<u>N74434</u>	<u>002</u>	Feb 13, 1996
<u>AB</u>		<u>50MG</u>	<u>N74434</u>	<u>003</u>	Feb 13, 1996
<u>AB</u>		<u>100MG</u>	<u>N74434</u>	<u>004</u>	Feb 13, 1996
<u>AB</u>	SANDOZ	<u>12.5MG</u>	<u>N74363</u>	<u>001</u>	Nov 09, 1995
<u>AB</u>		<u>12.5MG</u>	<u>N74481</u>	<u>001</u>	Feb 13, 1996
<u>AB</u>		<u>12.5MG</u>	<u>N74519</u>	<u>001</u>	Feb 13, 1996
<u>AB</u>		<u>25MG</u>	<u>N74363</u>	<u>002</u>	Nov 09, 1995
<u>AB</u>		<u>25MG</u>	<u>N74481</u>	<u>002</u>	Feb 13, 1996
<u>AB</u>		<u>25MG</u>	<u>N74519</u>	<u>002</u>	Feb 13, 1996
<u>AB</u>		<u>50MG</u>	<u>N74363</u>	<u>003</u>	Nov 09, 1995
<u>AB</u>		<u>50MG</u>	<u>N74481</u>	<u>003</u>	Feb 13, 1996
<u>AB</u>		<u>50MG</u>	<u>N74519</u>	<u>003</u>	Feb 13, 1996
<u>AB</u>		<u>100MG</u>	<u>N74363</u>	<u>004</u>	Nov 09, 1995
<u>AB</u>		<u>100MG</u>	<u>N74481</u>	<u>004</u>	Feb 13, 1996
<u>AB</u>		<u>100MG</u>	<u>N74519</u>	<u>004</u>	Feb 13, 1996
<u>AB</u>		<u>12.5MG</u>	<u>N74677</u>	<u>004</u>	May 30, 1997
<u>AB</u>		<u>25MG</u>	<u>N74677</u>	<u>002</u>	May 30, 1997
<u>AB</u>		<u>50MG</u>	<u>N74677</u>	<u>001</u>	May 30, 1997
<u>AB</u>		<u>100MG</u>	<u>N74677</u>	<u>003</u>	May 30, 1997
<u>AB</u>	TEVA	<u>12.5MG</u>	<u>N74322</u>	<u>001</u>	Feb 13, 1996
<u>AB</u>		<u>12.5MG</u>	<u>N74433</u>	<u>001</u>	Feb 13, 1996
<u>AB</u>		<u>12.5MG</u>	<u>N74483</u>	<u>001</u>	Feb 13, 1996
<u>AB</u>		<u>25MG</u>	<u>N74322</u>	<u>002</u>	Feb 13, 1996
<u>AB</u>		<u>25MG</u>	<u>N74433</u>	<u>002</u>	Feb 13, 1996
<u>AB</u>		<u>25MG</u>	<u>N74483</u>	<u>002</u>	Feb 13, 1996
<u>AB</u>		<u>50MG</u>	<u>N74322</u>	<u>003</u>	Feb 13, 1996
<u>AB</u>		<u>50MG</u>	<u>N74433</u>	<u>003</u>	Feb 13, 1996
<u>AB</u>		<u>50MG</u>	<u>N74483</u>	<u>003</u>	Feb 13, 1996
<u>AB</u>		<u>100MG</u>	<u>N74322</u>	<u>004</u>	Feb 13, 1996
<u>AB</u>		<u>100MG</u>	<u>N74433</u>	<u>004</u>	Feb 13, 1996
<u>AB</u>		<u>100MG</u>	<u>N74483</u>	<u>004</u>	Feb 13, 1996
<u>AB</u>	TEVA PHARMS	<u>12.5MG</u>	<u>N74462</u>	<u>001</u>	Feb 13, 1996
<u>AB</u>		<u>25MG</u>	<u>N74462</u>	<u>002</u>	Feb 13, 1996
<u>AB</u>		<u>50MG</u>	<u>N74462</u>	<u>003</u>	Feb 13, 1996
<u>AB</u>		<u>100MG</u>	<u>N74462</u>	<u>004</u>	Feb 13, 1996
<u>AB</u>	TORPHARM	<u>12.5MG</u>	<u>N74737</u>	<u>001</u>	Oct 28, 1998
<u>AB</u>		<u>25MG</u>	<u>N74737</u>	<u>002</u>	Oct 28, 1998
<u>AB</u>		<u>50MG</u>	<u>N74737</u>	<u>003</u>	Oct 28, 1998
<u>AB</u>		<u>100MG</u>	<u>N74737</u>	<u>004</u>	Oct 28, 1998
<u>AB</u>	WATSON LABS	<u>12.5MG</u>	<u>N74386</u>	<u>001</u>	May 23, 1996
<u>AB</u>		<u>12.5MG</u>	<u>N74451</u>	<u>001</u>	Feb 13, 1996
<u>AB</u>		<u>12.5MG</u>	<u>N74576</u>	<u>001</u>	Apr 23, 1996
<u>AB</u>		<u>25MG</u>	<u>N74386</u>	<u>002</u>	May 23, 1996
<u>AB</u>		<u>25MG</u>	<u>N74451</u>	<u>002</u>	Feb 13, 1996
<u>AB</u>		<u>25MG</u>	<u>N74576</u>	<u>002</u>	Apr 23, 1996
<u>AB</u>		<u>50MG</u>	<u>N74386</u>	<u>003</u>	May 23, 1996

PRESCRIPTION DRUG PRODUCT LIST

3 - 67 (of 370)

CAPTOPRIL

TABLET; ORAL

CAPTOPRIL

<u>AB</u>	WATSON LABS	<u>50MG</u>	<u>N74451</u>	<u>003</u>	Feb 13, 1996
<u>AB</u>		<u>50MG</u>	<u>N74576</u>	<u>003</u>	Apr 23, 1996
<u>AB</u>		<u>100MG</u>	<u>N74386</u>	<u>004</u>	May 23, 1996
<u>AB</u>		<u>100MG</u>	<u>N74451</u>	<u>004</u>	Feb 13, 1996
<u>AB</u>		<u>100MG</u>	<u>N74576</u>	<u>004</u>	Apr 23, 1996
<u>AB</u>	WEST WARD	<u>12.5MG</u>	<u>N74505</u>	<u>001</u>	Feb 13, 1996
<u>AB</u>		<u>25MG</u>	<u>N74505</u>	<u>002</u>	Feb 13, 1996
<u>AB</u>		<u>50MG</u>	<u>N74505</u>	<u>003</u>	Feb 13, 1996
<u>AB</u>		<u>100MG</u>	<u>N74505</u>	<u>004</u>	Feb 13, 1996
<u>AB</u>	WOCKHARDT	<u>12.5MG</u>	<u>N74532</u>	<u>001</u>	Mar 28, 1997
<u>AB</u>		<u>25MG</u>	<u>N74532</u>	<u>002</u>	Mar 28, 1997
<u>AB</u>		<u>50MG</u>	<u>N74532</u>	<u>003</u>	Mar 28, 1997
<u>AB</u>		<u>100MG</u>	<u>N74532</u>	<u>004</u>	Mar 28, 1997

CAPTOPRIL; HYDROCHLOROTHIAZIDE

TABLET; ORAL

<u>AB</u>	<u>CAPOZIDE 25/15</u> APOTHECON	<u>25MG;15MG</u>	<u>N18709</u>	<u>001</u>	Oct 12, 1984
<u>AB</u>	<u>CAPOZIDE 25/25</u> + APOTHECON	<u>25MG;25MG</u>	<u>N18709</u>	<u>002</u>	Oct 12, 1984
<u>AB</u>	<u>CAPOZIDE 50/15</u> + APOTHECON	<u>50MG;15MG</u>	<u>N18709</u>	<u>004</u>	Oct 12, 1984
<u>AB</u>	<u>CAPOZIDE 50/25</u> APOTHECON	<u>50MG;25MG</u>	<u>N18709</u>	<u>003</u>	Oct 12, 1984
<u>AB</u>	<u>CAPTOPRIL AND HYDROCHLOROTHIAZIDE</u> IVAX PHARMS	<u>25MG;15MG</u>	<u>N75055</u>	<u>001</u>	Jun 18, 1998
<u>AB</u>		<u>25MG;25MG</u>	<u>N75055</u>	<u>002</u>	Jun 18, 1998
<u>AB</u>		<u>50MG;15MG</u>	<u>N75055</u>	<u>004</u>	Jun 18, 1998
<u>AB</u>		<u>50MG;25MG</u>	<u>N75055</u>	<u>003</u>	Jun 18, 1998
<u>AB</u>	MYLAN	<u>25MG;15MG</u>	<u>N74896</u>	<u>001</u>	Dec 29, 1997
<u>AB</u>		<u>25MG;25MG</u>	<u>N74896</u>	<u>002</u>	Dec 29, 1997
<u>AB</u>		<u>50MG;15MG</u>	<u>N74896</u>	<u>004</u>	Dec 29, 1997
<u>AB</u>		<u>50MG;25MG</u>	<u>N74896</u>	<u>003</u>	Dec 29, 1997
<u>AB</u>	TEVA	<u>25MG;15MG</u>	<u>N74827</u>	<u>001</u>	Dec 29, 1997
<u>AB</u>		<u>25MG;25MG</u>	<u>N74827</u>	<u>002</u>	Dec 29, 1997
<u>AB</u>		<u>50MG;15MG</u>	<u>N74827</u>	<u>004</u>	Dec 29, 1997
<u>AB</u>		<u>50MG;25MG</u>	<u>N74827</u>	<u>003</u>	Dec 29, 1997

CARBACHOL

SOLUTION; INTRAOCULAR

<u>AT</u>	<u>CARBASTAT</u> NOVARTIS	<u>0.01%</u>	<u>N73677</u>	<u>001</u>	Apr 28, 1995
<u>AT</u>	<u>MIOSTAT</u> + ALCON	<u>0.01%</u>	<u>N16968</u>	<u>001</u>	

CARBAMAZEPINE

CAPSULE, EXTENDED RELEASE; ORAL

CARBATROL

	SHIRE	100MG	N20712	003	Sep 30, 1997
		200MG	N20712	001	Sep 30, 1997
+		300MG	N20712	002	Sep 30, 1997
	EQUETRO				
	SHIRE	100MG	N21710	001	Dec 10, 2004
		200MG	N21710	002	Dec 10, 2004
+		300MG	N21710	003	Dec 10, 2004

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 68 (of 370)

CARBAMAZEPINE

SUSPENSION; ORAL

CARBAMAZEPINE

<u>AB</u>	MORTON GROVE	<u>100MG/5ML</u>	<u>N75714</u>	<u>001</u>	Jun 05, 2002
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TEGRETOL

<u>AB</u>	+ NOVARTIS	<u>100MG/5ML</u>	<u>N18927</u>	<u>001</u>	Dec 18, 1987
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TERIL

<u>AB</u>	TARO	<u>100MG/5ML</u>	<u>N76729</u>	<u>001</u>	Sep 20, 2004
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TABLET; ORAL

CARBAMAZEPINE

<u>AB</u>	ACTAVIS ELIZABETH	<u>200MG</u>	<u>N71696</u>	<u>001</u>	Nov 09, 1987
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<u>AB</u>	APOTEX INC	<u>200MG</u>	<u>N75948</u>	<u>001</u>	Feb 27, 2002
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<u>AB</u>	CARACO	<u>200MG</u>	<u>N77272</u>	<u>002</u>	Dec 07, 2005
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		100MG	N77272	001	Dec 07, 2005
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		300MG	N77272	003	Dec 07, 2005
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		400MG	N77272	004	Dec 07, 2005
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<u>AB</u>	INWOOD LABS	<u>200MG</u>	<u>N70231</u>	<u>001</u>	Aug 14, 1986
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<u>AB</u>	PLIVA	<u>200MG</u>	<u>N71479</u>	<u>001</u>	Jul 24, 1987
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<u>AB</u>	TARO	<u>200MG</u>	<u>N74649</u>	<u>001</u>	Oct 03, 1996
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EPITOL

<u>AB</u>	TEVA	<u>200MG</u>	<u>N70541</u>	<u>001</u>	Sep 17, 1986
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TEGRETOL

<u>AB</u>	+ NOVARTIS	<u>200MG</u>	<u>N16608</u>	<u>001</u>	
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TERIL

<u>AB</u>	TARO	<u>200MG</u>	<u>N76525</u>	<u>001</u>	Sep 26, 2003
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TABLET, CHEWABLE; ORAL

CARBAMAZEPINE

<u>AB</u>	CARACO	<u>100MG</u>	<u>N75712</u>	<u>001</u>	Jul 05, 2001
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<u>AB</u>	JUBILANT PHARMS	<u>100MG</u>	<u>N71940</u>	<u>001</u>	Feb 01, 1988
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<u>AB</u>	TARO PHARM INDS	<u>100MG</u>	<u>N75687</u>	<u>001</u>	Oct 24, 2000
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	+	200MG	N75687	002	Jul 29, 2002
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EPITOL

<u>AB</u>	TEVA	<u>100MG</u>	<u>N73524</u>	<u>001</u>	Jul 29, 1992
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TEGRETOL

<u>AB</u>	+ NOVARTIS	<u>100MG</u>	<u>N18281</u>	<u>001</u>	
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TABLET, EXTENDED RELEASE; ORAL

TEGRETOL-XR

	NOVARTIS	100MG	N20234	001	Mar 25, 1996
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		200MG	N20234	002	Mar 25, 1996
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	+	400MG	N20234	003	Mar 25, 1996
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CARBENICILLIN INDANYL SODIUM

TABLET; ORAL

GEOCILLIN

	+ PFIZER	EQ 382MG BASE	N50435	001	
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CARBIDOPA

TABLET; ORAL

LODOSYN

	+ BRISTOL MYERS SQUIBB	25MG	N17830	001	
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CARBIDOPA; ENTACAPONE; LEVODOPA

TABLET; ORAL

STALEVO 100

	ORION PHARMA INC	25MG;200MG;100MG	N21485	002	Jun 11, 2003
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STALEVO 150

	+ ORION PHARMA INC	37.5MG;200MG;150MG	N21485	003	Jun 11, 2003
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STALEVO 50

	ORION PHARMA INC	12.5MG;200MG;50MG	N21485	001	Jun 11, 2003
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27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 69 (of 370)

CARBIDOPA; LEVODOPA

TABLET; ORAL

CARBIDOPA AND LEVODOPA

<u>AB</u>	ACTAVIS ELIZABETH	<u>10MG;100MG</u>	<u>N74260</u>	<u>001</u>	Sep 03, 1993
<u>AB</u>		<u>25MG;100MG</u>	<u>N74260</u>	<u>002</u>	Sep 03, 1993
<u>AB</u>		<u>25MG;250MG</u>	<u>N74260</u>	<u>003</u>	Sep 03, 1993
<u>AB</u>	SANDOZ	<u>10MG;100MG</u>	<u>N73586</u>	<u>001</u>	Jun 29, 1995
<u>AB</u>		<u>25MG;100MG</u>	<u>N73587</u>	<u>001</u>	Jun 29, 1995
<u>AB</u>		<u>25MG;250MG</u>	<u>N73620</u>	<u>001</u>	Jun 29, 1995
<u>AB</u>	TEVA	<u>10MG;100MG</u>	<u>N73618</u>	<u>001</u>	Aug 28, 1992
<u>AB</u>		<u>25MG;100MG</u>	<u>N73589</u>	<u>001</u>	Aug 28, 1992
<u>AB</u>		<u>25MG;250MG</u>	<u>N73607</u>	<u>001</u>	Aug 28, 1992
	<u>SINEMET</u>				
<u>AB</u>	BRISTOL MYERS SQUIBB	<u>10MG;100MG</u>	<u>N17555</u>	<u>001</u>	
<u>AB</u>		<u>25MG;100MG</u>	<u>N17555</u>	<u>003</u>	
<u>AB</u>	+	<u>25MG;250MG</u>	<u>N17555</u>	<u>002</u>	

TABLET, EXTENDED RELEASE; ORAL

CARBIDOPA AND LEVODOPA

<u>AB</u>	IMPAX LABS	<u>25MG;100MG</u>	<u>N76521</u>	<u>001</u>	May 14, 2004
<u>AB</u>		<u>50MG;200MG</u>	<u>N76521</u>	<u>002</u>	May 14, 2004
<u>AB</u>	KV PHARM	<u>50MG;200MG</u>	<u>N76663</u>	<u>001</u>	Jun 24, 2004
<u>AB</u>	MYLAN	<u>25MG;100MG</u>	<u>N75091</u>	<u>002</u>	Apr 21, 2000
<u>AB</u>		<u>50MG;200MG</u>	<u>N75091</u>	<u>001</u>	Sep 30, 1999
<u>AB</u>	TORPHARM	<u>25MG;100MG</u>	<u>N76212</u>	<u>001</u>	Jun 16, 2004
<u>AB</u>		<u>50MG;200MG</u>	<u>N76212</u>	<u>002</u>	Jun 16, 2004
	<u>SINEMET CR</u>				
<u>AB</u>	BRISTOL MYERS SQUIBB	<u>25MG;100MG</u>	<u>N19856</u>	<u>002</u>	Dec 24, 1992
<u>AB</u>	+	<u>50MG;200MG</u>	<u>N19856</u>	<u>001</u>	May 30, 1991

TABLET, FOR SUSPENSION; ORAL

CARBILEV

	RANBAXY	10MG;100MG	N76643	001	Jun 10, 2005
		25MG;100MG	N76643	002	Jun 10, 2005
	+	25MG;250MG	N76643	003	Jun 10, 2005

TABLET, ORALLY DISINTEGRATING; ORAL

PARCOPA

	SCHWARZ PHARMA	10MG;100MG	N76699	001	Aug 27, 2004
		25MG;100MG	N76699	002	Aug 27, 2004
	+	25MG;250MG	N76699	003	Aug 27, 2004

CARBINOXAMINE MALEATE

SOLUTION; ORAL

CARBINOXAMINE MALEATE

	+ MIKART	4MG/5ML	N40458	001	Apr 25, 2003
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TABLET; ORAL

CARBINOXAMINE MALEATE

	+ MIKART	4MG	N40442	001	Mar 19, 2003
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CARBOPLATIN

INJECTABLE; INJECTION

CARBOPLATIN

<u>AP</u>	ABRAXIS PHARM	<u>50MG/VIAL</u>	<u>N76235</u>	<u>001</u>	Oct 14, 2004
<u>AP</u>		<u>150MG/VIAL</u>	<u>N76235</u>	<u>002</u>	Oct 14, 2004
<u>AP</u>		<u>450MG/VIAL</u>	<u>N76235</u>	<u>003</u>	Oct 14, 2004
<u>AP</u>	BEDFORD	<u>50MG/VIAL</u>	<u>N76099</u>	<u>001</u>	Oct 14, 2004
<u>AP</u>		<u>150MG/VIAL</u>	<u>N76099</u>	<u>002</u>	Oct 14, 2004
<u>AP</u>		<u>450MG/VIAL</u>	<u>N76099</u>	<u>003</u>	Oct 14, 2004
<u>AP</u>	PHARMACHEMIE	<u>50MG/VIAL</u>	<u>N76162</u>	<u>001</u>	Oct 14, 2004
<u>AP</u>		<u>150MG/VIAL</u>	<u>N76162</u>	<u>002</u>	Oct 14, 2004
<u>AP</u>		<u>450MG/VIAL</u>	<u>N76162</u>	<u>003</u>	Oct 14, 2004

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 70 (of 370)

CARBOPLATIN

INJECTABLE; INJECTION

CARBOPLATIN

<u>AP</u>	<u>PLIVA</u>	<u>50MG/VIAL</u>	<u>N76602</u>	<u>001</u>	Nov 16, 2004
<u>AP</u>		<u>150MG/VIAL</u>	<u>N76602</u>	<u>002</u>	Nov 16, 2004
<u>AP</u>		<u>450MG/VIAL</u>	<u>N76602</u>	<u>003</u>	Nov 16, 2004
<u>AP</u>	<u>SANDOZ</u>	<u>50MG/VIAL</u>	<u>N76959</u>	<u>001</u>	Mar 18, 2005
<u>AP</u>		<u>150MG/VIAL</u>	<u>N76959</u>	<u>002</u>	Mar 18, 2005
<u>AP</u>		<u>450MG/VIAL</u>	<u>N76959</u>	<u>003</u>	Mar 18, 2005
<u>AP</u>	<u>WATSON LABS</u>	<u>50MG/VIAL</u>	<u>N77383</u>	<u>001</u>	Jan 27, 2006
<u>AP</u>		<u>150MG/VIAL</u>	<u>N77383</u>	<u>002</u>	Jan 27, 2006
<u>AP</u>		<u>450MG/VIAL</u>	<u>N77383</u>	<u>003</u>	Jan 27, 2006
<u>PARAPLATIN</u>					
<u>AP</u>	<u>+ BRISTOL MYERS SQUIBB</u>	<u>50MG/VIAL</u>	<u>N19880</u>	<u>001</u>	Mar 03, 1989
<u>AP</u>	<u>+</u>	<u>150MG/VIAL</u>	<u>N19880</u>	<u>002</u>	Mar 03, 1989
<u>AP</u>	<u>+</u>	<u>450MG/VIAL</u>	<u>N19880</u>	<u>003</u>	Mar 03, 1989

INJECTABLE; IV (INFUSION)

CARBOPLATIN

<u>AP</u>	<u>ABRAXIS PHARM</u>	<u>EQ 50MG/5ML (10MG/ML)</u>	<u>N77266</u>	<u>001</u>	Feb 15, 2006
<u>AP</u>		<u>EQ 150MG/15ML (10MG/ML)</u>	<u>N77266</u>	<u>002</u>	Feb 15, 2006
<u>AP</u>		<u>EQ 450MG/45ML (10MG/ML)</u>	<u>N77247</u>	<u>003</u>	Oct 21, 2004
<u>AP</u>		<u>EQ 450MG/45ML (10MG/ML)</u>	<u>N77266</u>	<u>003</u>	Feb 15, 2006
<u>AP</u>		<u>EQ 600MG/60ML (10MG/ML)</u>	<u>N77266</u>	<u>004</u>	Feb 15, 2006
<u>AP</u>	<u>BEDFORD LABS</u>	<u>EQ 50MG/5ML (10MG/ML)</u>	<u>N77244</u>	<u>001</u>	Oct 15, 2004
<u>AP</u>		<u>EQ 150MG/15ML (10MG/ML)</u>	<u>N77244</u>	<u>002</u>	Oct 15, 2004
<u>AP</u>		<u>EQ 450MG/45ML (10MG/ML)</u>	<u>N77244</u>	<u>003</u>	Oct 15, 2004
<u>AP</u>		<u>EQ 600MG/60ML (10MG/ML)</u>	<u>N77244</u>	<u>004</u>	Jan 20, 2006
<u>AP</u>	<u>DABUR ONCOLOGY PLC</u>	<u>EQ 50MG/5ML (10MG/ML)</u>	<u>N77432</u>	<u>001</u>	Sep 29, 2006
<u>AP</u>		<u>EQ 150MG/15ML (10MG/ML)</u>	<u>N77432</u>	<u>002</u>	Sep 29, 2006
<u>AP</u>		<u>EQ 450MG/45ML (10MG/ML)</u>	<u>N77432</u>	<u>003</u>	Sep 29, 2006
<u>AP</u>	<u>MAYNE PHARMA USA</u>	<u>EQ 50MG/5ML (10MG/ML)</u>	<u>N76517</u>	<u>001</u>	Oct 14, 2004
<u>AP</u>		<u>EQ 150MG/15ML (10MG/ML)</u>	<u>N76517</u>	<u>002</u>	Oct 14, 2004
<u>AP</u>		<u>EQ 450MG/45ML (10MG/ML)</u>	<u>N76517</u>	<u>003</u>	Oct 14, 2004
<u>AP</u>		<u>EQ 600MG/60ML (10MG/ML)</u>	<u>N77059</u>	<u>001</u>	Nov 23, 2004
<u>AP</u>	<u>PHARMACHEMIE</u>	<u>EQ 50MG/5ML (10MG/ML)</u>	<u>N77269</u>	<u>001</u>	Oct 14, 2004
<u>AP</u>		<u>EQ 150MG/15ML (10MG/ML)</u>	<u>N77269</u>	<u>002</u>	Oct 14, 2004
<u>AP</u>		<u>EQ 450MG/45ML (10MG/ML)</u>	<u>N77269</u>	<u>003</u>	Oct 14, 2004
<u>AP</u>	<u>SICOR PHARMS</u>	<u>EQ 50MG/5ML (10MG/ML)</u>	<u>N77139</u>	<u>001</u>	Sep 21, 2005
<u>AP</u>		<u>EQ 150MG/15ML (10MG/ML)</u>	<u>N77139</u>	<u>002</u>	Sep 21, 2005
<u>AP</u>		<u>EQ 450MG/45ML (10MG/ML)</u>	<u>N77139</u>	<u>003</u>	Sep 21, 2005
<u>AP</u>		<u>EQ 600MG/60ML (10MG/ML)</u>	<u>N77139</u>	<u>004</u>	Sep 21, 2005
<u>AP</u>	<u>SPECTRUM PHARMS</u>	<u>EQ 50MG/5ML (10MG/ML)</u>	<u>N77096</u>	<u>001</u>	Jun 14, 2005
<u>AP</u>		<u>EQ 150MG/15ML (10MG/ML)</u>	<u>N77096</u>	<u>002</u>	Jun 14, 2005
<u>AP</u>		<u>EQ 450MG/45ML (10MG/ML)</u>	<u>N77096</u>	<u>003</u>	Jun 14, 2005
<u>PARAPLATIN</u>					
<u>AP</u>	<u>+ BRISTOL MYERS SQUIBB</u>	<u>EQ 50MG /5ML (10MG/ML)</u>	<u>N20452</u>	<u>001</u>	Jul 14, 2003
<u>AP</u>	<u>+</u>	<u>EQ 150MG /15ML (10MG/ML)</u>	<u>N20452</u>	<u>002</u>	Jul 14, 2003
<u>AP</u>	<u>+</u>	<u>EQ 450MG /45ML (10MG/ML)</u>	<u>N20452</u>	<u>003</u>	Jul 14, 2003
<u>AP</u>	<u>+</u>	<u>EQ 600MG /60ML (10MG/ML)</u>	<u>N20452</u>	<u>004</u>	Jan 15, 2004

CARBOPROST TROMETHAMINE

INJECTABLE; INJECTION

HEMABATE

+ PHARMACIA AND UPJOHN EQ 0.25MG BASE/ML N17989 001

CARISOPRODOL

TABLET; ORAL

CARISOPRODOL

<u>AA</u>	<u>ACTAVIS TOTOWA</u>	<u>350MG</u>	<u>N40188</u>	<u>001</u>	Mar 07, 1997
<u>AA</u>	<u>COREPHARMA</u>	<u>350MG</u>	<u>N40397</u>	<u>001</u>	Sep 21, 2000

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 71 (of 370)

CARISOPRODOL

TABLET; ORAL

CARISOPRODOL

<u>AA</u>	MUTUAL PHARM	<u>350MG</u>	<u>N89346</u>	<u>001</u>	Oct 17, 1991
<u>AA</u>	NEIL	<u>350MG</u>	<u>N40576</u>	<u>001</u>	Jun 07, 2005
<u>AA</u>	SANDOZ	<u>350MG</u>	<u>N81025</u>	<u>001</u>	Apr 13, 1989
<u>AA</u>	VINTAGE PHARMS	<u>350MG</u>	<u>N40245</u>	<u>001</u>	Sep 08, 1997
<u>AA</u>	WATSON LABS	<u>350MG</u>	<u>N40152</u>	<u>001</u>	Dec 03, 1996
<u>AA</u>		<u>350MG</u>	<u>N86179</u>	<u>001</u>	
<u>AA</u>		<u>350MG</u>	<u>N87499</u>	<u>001</u>	Apr 20, 1982
<u>AA</u>	WEST WARD	<u>350MG</u>	<u>N40124</u>	<u>001</u>	Jan 24, 1996
<u>SOMA</u>					
<u>AA</u>	+ MEDPOINTE PHARM HLC	<u>350MG</u>	<u>N11792</u>	<u>001</u>	

CARMUSTINE

IMPLANT; INTRACRANIAL

GLIADEL

+ MGI PHARMA INC	7.7MG	N20637	001	Sep 23, 1996
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INJECTABLE; INJECTION

BICNU

+ BRISTOL	100MG/VIAL	N17422	001	
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CARTEOLOL HYDROCHLORIDE

SOLUTION/DROPS; OPHTHALMIC

CARTEOLOL HYDROCHLORIDE

<u>AT</u>	ALCON	<u>1%</u>	<u>N75476</u>	<u>001</u>	Jan 03, 2000
<u>AT</u>	BAUSCH AND LOMB	<u>1%</u>	<u>N75546</u>	<u>001</u>	Jan 20, 2000
<u>AT</u>	NOVEX	<u>1%</u>	<u>N76097</u>	<u>001</u>	Feb 06, 2002
<u>OCUPRESS</u>					
<u>AT</u>	+ NOVARTIS	<u>1%</u>	<u>N19972</u>	<u>001</u>	May 23, 1990

CARVEDILOL

TABLET; ORAL

COREG

SMITHKLINE BEECHAM	3.125MG	N20297	004	May 29, 1997
	6.25MG	N20297	003	Sep 14, 1995
+	12.5MG	N20297	002	Sep 14, 1995
	25MG	N20297	001	Sep 14, 1995

CARVEDILOL PHOSPHATE

CAPSULE, EXTENDED RELEASE; ORAL

COREG CR

SB PHARMC0	10MG	N22012	001	Oct 20, 2006
	20MG	N22012	002	Oct 20, 2006
	40MG	N22012	003	Oct 20, 2006
+	80MG	N22012	004	Oct 20, 2006

CASPOFUNGIN ACETATE

INJECTABLE; IV (INFUSION)

+ MERCK	50MG/VIAL	N21227	001	Jan 26, 2001
+	70MG/VIAL	N21227	002	Jan 26, 2001

CEFACLOR

CAPSULE; ORAL

CEFACLOR

<u>AB</u>	CARLSBAD	<u>EQ 250MG BASE</u>	<u>N65146</u>	<u>001</u>	Jan 22, 2004
<u>AB</u>		<u>EQ 500MG BASE</u>	<u>N65146</u>	<u>002</u>	Jan 22, 2004
<u>AB</u>	CEPH INTL	<u>EQ 250MG BASE</u>	<u>N62205</u>	<u>001</u>	

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 72 (of 370)

CEFACLOR

CAPSULE; ORAL

CEFACLOR

<u>AB</u>	CEPH INTL	<u>EQ 500MG BASE</u>	<u>N62205</u>	<u>002</u>	
<u>AB</u>	IVAX PHARMS	<u>EQ 250MG BASE</u>	<u>N64061</u>	<u>001</u>	Apr 27, 1995
<u>AB</u>		<u>EQ 500MG BASE</u>	<u>N64061</u>	<u>002</u>	Apr 27, 1995
<u>AB</u>	MARSAM PHARMS LLC	<u>EQ 250MG BASE</u>	<u>N64148</u>	<u>001</u>	May 23, 1996
<u>AB</u>		<u>EQ 500MG BASE</u>	<u>N64148</u>	<u>002</u>	May 23, 1996
<u>AB</u>	RANBAXY	<u>EQ 250MG BASE</u>	<u>N64156</u>	<u>001</u>	Aug 28, 1997
<u>AB</u>	+	<u>EQ 500MG BASE</u>	<u>N64156</u>	<u>002</u>	Aug 28, 1997
<u>AB</u>	TEVA	<u>EQ 250MG BASE</u>	<u>N64145</u>	<u>001</u>	Jun 24, 1996
<u>AB</u>		<u>EQ 500MG BASE</u>	<u>N64145</u>	<u>002</u>	Jun 24, 1996

FOR SUSPENSION; ORAL

CEFACLOR

<u>AB</u>	CEPH INTL	<u>EQ 125MG BASE/5ML</u>	<u>N62206</u>	<u>001</u>	
<u>AB</u>		<u>EQ 187MG BASE/5ML</u>	<u>N62206</u>	<u>003</u>	Apr 20, 1988
<u>AB</u>		<u>EQ 250MG BASE/5ML</u>	<u>N62206</u>	<u>002</u>	
<u>AB</u>		<u>EQ 375MG BASE/5ML</u>	<u>N62206</u>	<u>004</u>	Apr 20, 1988
<u>AB</u>	IVAX PHARMS	<u>EQ 375MG BASE/5ML</u>	<u>N64070</u>	<u>001</u>	Apr 28, 1995
<u>AB</u>	MARSAM PHARMS LLC	<u>EQ 125MG BASE/5ML</u>	<u>N64204</u>	<u>001</u>	Feb 18, 1998
<u>AB</u>		<u>EQ 187MG BASE/5ML</u>	<u>N64205</u>	<u>001</u>	Feb 18, 1998
<u>AB</u>		<u>EQ 250MG BASE/5ML</u>	<u>N64206</u>	<u>001</u>	Feb 18, 1998
<u>AB</u>		<u>EQ 375MG BASE/5ML</u>	<u>N64207</u>	<u>001</u>	Feb 18, 1998
<u>AB</u>	RANBAXY	<u>EQ 125MG BASE/5ML</u>	<u>N64166</u>	<u>001</u>	Oct 02, 1997
<u>AB</u>		<u>EQ 187MG BASE/5ML</u>	<u>N64165</u>	<u>001</u>	Oct 02, 1997
<u>AB</u>		<u>EQ 250MG BASE/5ML</u>	<u>N64164</u>	<u>001</u>	Oct 02, 1997
<u>AB</u>	+	<u>EQ 375MG BASE/5ML</u>	<u>N64155</u>	<u>001</u>	Oct 02, 1997

TABLET, CHEWABLE; ORAL

RANICLOR

	RANBAXY	EQ 125MG BASE	N65092	001	Dec 22, 2003
		EQ 187MG BASE	N65092	002	Dec 22, 2003
		EQ 250MG BASE	N65092	003	Dec 22, 2003
	+	EQ 375MG BASE	N65092	004	Dec 22, 2003

TABLET, EXTENDED RELEASE; ORAL

CEFACLOR

<u>AB</u>	+	PAR PHARM	<u>EQ 500MG BASE</u>	<u>N65057</u>	<u>001</u>	Jan 05, 2001
<u>AB</u>		TEVA	<u>EQ 500MG BASE</u>	<u>N65058</u>	<u>002</u>	Sep 04, 2002
			EQ 375MG BASE	N65058	001	Sep 04, 2002

CEFADROXIL/CEFADROXIL HEMIHYDRATE

CAPSULE; ORAL

CEFADROXIL

<u>AB</u>	+	IVAX PHARMS	<u>EQ 500MG BASE</u>	<u>N62766</u>	<u>001</u>	Mar 03, 1987
<u>AB</u>		ORCHID HLTHCARE	<u>EQ 500MG BASE</u>	<u>N65309</u>	<u>001</u>	Sep 18, 2006
<u>AB</u>		RANBAXY	<u>EQ 500MG BASE</u>	<u>N65015</u>	<u>001</u>	Jun 22, 1999
<u>AB</u>		SANDOZ	<u>EQ 500MG BASE</u>	<u>N62291</u>	<u>001</u>	
<u>AB</u>		TEVA PHARMS	<u>EQ 500MG BASE</u>	<u>N65282</u>	<u>001</u>	Jan 20, 2006
<u>AB</u>		WESTWARD	<u>EQ 500MG BASE</u>	<u>N65311</u>	<u>001</u>	Feb 07, 2006

FOR SUSPENSION; ORAL

CEFADROXIL

<u>AB</u>		ORCHID HLTHCARE	<u>EQ 250MG BASE/5ML</u>	<u>N65307</u>	<u>002</u>	Oct 16, 2006
<u>AB</u>			<u>EQ 500MG BASE/5ML</u>	<u>N65307</u>	<u>003</u>	Oct 16, 2006
<u>AB</u>		RANBAXY	<u>EQ 250MG BASE/5ML</u>	<u>N65115</u>	<u>002</u>	Mar 26, 2003
<u>AB</u>			<u>EQ 500MG BASE/5ML</u>	<u>N65115</u>	<u>003</u>	Mar 26, 2003
	+		EQ 125MG BASE/5ML	N65115	001	Mar 26, 2003
<u>AB</u>		TEVA PHARMS	<u>EQ 250MG BASE/5ML</u>	<u>N65278</u>	<u>001</u>	Jan 20, 2006
<u>AB</u>			<u>EQ 500MG BASE/5ML</u>	<u>N65278</u>	<u>002</u>	Jan 20, 2006
		<u>DURICEF</u>				
<u>AB</u>		WARNER CHILCOTT	<u>EQ 250MG BASE/5ML</u>	<u>N50527</u>	<u>003</u>	

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 73 (of 370)

CEFADROXIL/CEFADROXIL HEMIHYDRATE

FOR SUSPENSION; ORAL

DURICEF

<u>AB</u>	+	WARNER CHILCOTT	<u>EQ 500MG BASE/5ML</u>	<u>N50527</u>	<u>001</u>	
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TABLET; ORAL

CEFADROXIL

<u>AB</u>		HIKMA	<u>EQ 1GM BASE</u>	<u>N65260</u>	<u>001</u>	Mar 30, 2006
<u>AB</u>	+	IVAX PHARMS	<u>EQ 1GM BASE</u>	<u>N62774</u>	<u>001</u>	Apr 08, 1987
<u>AB</u>		ORCHID HLTHCARE	<u>EQ 1GM BASE</u>	<u>N65301</u>	<u>001</u>	Sep 18, 2006
<u>AB</u>		RANBAXY	<u>EQ 1GM BASE</u>	<u>N65018</u>	<u>001</u>	Apr 23, 1999

CEFAZOLIN SODIUM

INJECTABLE; INJECTION

ANCEF

<u>AP</u>	+	GLAXOSMITHKLINE	<u>EQ 1GM BASE/VIAL</u>	<u>N50461</u>	<u>003</u>	
<u>AP</u>	+		<u>EQ 10GM BASE/VIAL</u>	<u>N50461</u>	<u>005</u>	

ANCEF IN PLASTIC CONTAINER

+	BAXTER HLTHCARE	EQ 10MG BASE/ML	N63002	001	Mar 28, 1991
+		EQ 20MG BASE/ML	N63002	002	Mar 28, 1991

CEFAZOLIN

<u>AP</u>		ORCHID HLTHCARE	<u>EQ 1GM BASE/VIAL</u>	<u>N65244</u>	<u>001</u>	Aug 12, 2005
<u>AP</u>			<u>EQ 10GM BASE/VIAL</u>	<u>N65247</u>	<u>001</u>	Aug 12, 2005

CEFAZOLIN AND DEXTROSE

+	B BRAUN	EQ 1GM BASE/VIAL	N50779	002	Jul 27, 2000
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CEFAZOLIN SODIUM

<u>AP</u>	+	ABRAXIS PHARM	<u>EQ 500MG BASE/VIAL</u>	<u>N64169</u>	<u>001</u>	Aug 14, 1998
<u>AP</u>	+		<u>EQ 1GM BASE/VIAL</u>	<u>N64169</u>	<u>002</u>	Aug 14, 1998
<u>AP</u>	+		<u>EQ 10GM BASE/VIAL</u>	<u>N64170</u>	<u>001</u>	Mar 18, 1998
<u>AP</u>	+		<u>EQ 20GM BASE/VIAL</u>	<u>N64170</u>	<u>002</u>	Mar 18, 1998
<u>AP</u>		APOTHECON	<u>EQ 500MG BASE/VIAL</u>	<u>N62831</u>	<u>001</u>	Dec 09, 1988
<u>AP</u>			<u>EQ 1GM BASE/VIAL</u>	<u>N62831</u>	<u>002</u>	Dec 09, 1988
<u>AP</u>			<u>EQ 10GM BASE/VIAL</u>	<u>N62831</u>	<u>003</u>	Sep 25, 1992
<u>AP</u>		GLAXOSMITHKLINE	<u>EQ 1GM BASE/VIAL</u>	<u>N64033</u>	<u>001</u>	Oct 31, 1993
<u>AP</u>		HANFORD GC	<u>EQ 500MG BASE/VIAL</u>	<u>N63214</u>	<u>001</u>	Dec 27, 1991
<u>AP</u>			<u>EQ 500MG BASE/VIAL</u>	<u>N63216</u>	<u>001</u>	Dec 27, 1991
<u>AP</u>			<u>EQ 1GM BASE/VIAL</u>	<u>N63207</u>	<u>001</u>	Dec 27, 1991
<u>AP</u>			<u>EQ 1GM BASE/VIAL</u>	<u>N63208</u>	<u>001</u>	Dec 27, 1991
<u>AP</u>			<u>EQ 10GM BASE/VIAL</u>	<u>N63209</u>	<u>001</u>	Dec 27, 1991
<u>AP</u>		HIKMA FARMACEUTICA	<u>EQ 500MG BASE/VIAL</u>	<u>N65047</u>	<u>001</u>	Sep 18, 2001
<u>AP</u>			<u>EQ 1GM BASE/VIAL</u>	<u>N65047</u>	<u>002</u>	Sep 18, 2001
<u>AP</u>			<u>EQ 10GM BASE/VIAL</u>	<u>N65143</u>	<u>001</u>	Oct 18, 2004
<u>AP</u>		ORCHID HLTHCARE	<u>EQ 500MG BASE/VIAL</u>	<u>N65226</u>	<u>001</u>	Apr 21, 2005
<u>AP</u>			<u>EQ 1GM BASE/VIAL</u>	<u>N65226</u>	<u>002</u>	Apr 21, 2005
	+	SAMSON MEDCL	EQ 100GM BASE/VIAL	N65141	001	Nov 29, 2006
	+		EQ 300GM BASE/VIAL	N65141	002	Nov 29, 2006

CEFDINIR

CAPSULE; ORAL

CEFDINIR

<u>AB</u>		LUPIN	<u>300MG</u>	<u>N65264</u>	<u>001</u>	May 19, 2006
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OMNICEF

<u>AB</u>	+	ABBOTT	<u>300MG</u>	<u>N50739</u>	<u>001</u>	Dec 04, 1997
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FOR SUSPENSION; ORAL

CEFDINIR

<u>AB</u>		LUPIN	<u>125MG/5ML</u>	<u>N65259</u>	<u>001</u>	May 31, 2006
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OMNICEF

<u>AB</u>		ABBOTT	<u>125MG/5ML</u>	<u>N50749</u>	<u>001</u>	Dec 04, 1997
	+		250MG/5ML	N50749	002	Jul 29, 2004

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 74 (of 370)

CEFDITOREN PIVOXILTABLET; ORAL
SPECTRACEF

+ CORNERSTONE 200MG N21222 001 Aug 29, 2001

CEFEPIME HYDROCHLORIDE (ARGININE FORMULATION)INJECTABLE; INJECTION
MAXIPIME

+	BRISTOL MYERS SQUIBB	EQ 500MG BASE/VIAL	N50679	001	Jan 18, 1996
+		EQ 1GM BASE/VIAL	N50679	002	Jan 18, 1996
+		EQ 2GM BASE/VIAL	N50679	003	Jan 18, 1996

CEFIXIMESUSPENSION; ORAL
SUPRAX

+ LUPIN 100MG/5ML N65129 001 Feb 23, 2004

CEFOPERAZONE SODIUMINJECTABLE; INJECTION
CEFOBID

+	PFIZER	EQ 1GM BASE/VIAL	N50551	001	Nov 18, 1982
+		EQ 2GM BASE/VIAL	N50551	002	Nov 18, 1982
+		EQ 10GM BASE/VIAL	N50551	003	Mar 05, 1990

CEFOBID IN PLASTIC CONTAINER

+	PFIZER	EQ 20MG BASE/ML	N50613	002	Jul 31, 1987
+		EQ 40MG BASE/ML	N50613	001	Jul 23, 1986

CEFOTAXIME SODIUMINJECTABLE; INJECTION
CEFOTAXIME

<u>AP</u>	ABRAXIS PHARM	<u>EQ 500MG BASE/VIAL</u>	<u>N64200</u>	<u>001</u>	Mar 24, 2000
<u>AP</u>		<u>EQ 1GM BASE/VIAL</u>	<u>N64200</u>	<u>002</u>	Mar 24, 2000
<u>AP</u>		<u>EQ 2GM BASE/VIAL</u>	<u>N64200</u>	<u>003</u>	Mar 24, 2000
<u>AP</u>		<u>EQ 10GM BASE/VIAL</u>	<u>N64201</u>	<u>001</u>	Mar 24, 2000
	+	EQ 20GM BASE/VIAL	N64201	002	Mar 24, 2000
<u>AP</u>	HIKMA	<u>EQ 500MG BASE/VIAL</u>	<u>N65072</u>	<u>001</u>	Nov 20, 2002
<u>AP</u>		<u>EQ 1GM BASE/VIAL</u>	<u>N65072</u>	<u>002</u>	Nov 20, 2002
<u>AP</u>		<u>EQ 2GM BASE/VIAL</u>	<u>N65072</u>	<u>003</u>	Nov 20, 2002
<u>AP</u>		<u>EQ 10GM BASE/VIAL</u>	<u>N65071</u>	<u>001</u>	Nov 20, 2002
<u>AP</u>	WOCKHARDT	<u>EQ 1GM BASE/VIAL</u>	<u>N65197</u>	<u>001</u>	Aug 29, 2006

CEFOTAXIME SODIUM

<u>AP</u>	LUPIN	<u>EQ 500MG BASE/VIAL</u>	<u>N65124</u>	<u>001</u>	Sep 24, 2003
<u>AP</u>		<u>EQ 1GM BASE/VIAL</u>	<u>N65124</u>	<u>002</u>	Sep 24, 2003
<u>AP</u>		<u>EQ 2GM BASE/VIAL</u>	<u>N65124</u>	<u>003</u>	Sep 24, 2003
<u>AP</u>	ORCHID HLTHCARE	<u>EQ 500MG BASE/VIAL</u>	<u>N65290</u>	<u>001</u>	Aug 11, 2006
<u>AP</u>		<u>EQ 1GM BASE/VIAL</u>	<u>N65290</u>	<u>002</u>	Aug 11, 2006
<u>AP</u>		<u>EQ 1GM BASE/VIAL</u>	<u>N65293</u>	<u>001</u>	Aug 10, 2006
<u>AP</u>		<u>EQ 2GM BASE/VIAL</u>	<u>N65290</u>	<u>003</u>	Aug 11, 2006
<u>AP</u>		<u>EQ 2GM BASE/VIAL</u>	<u>N65293</u>	<u>002</u>	Aug 10, 2006
<u>AP</u>		<u>EQ 10GM BASE/VIAL</u>	<u>N65292</u>	<u>001</u>	Aug 10, 2006

CLAFORAN

<u>AP</u>	+	SANOFI AVENTIS US	<u>EQ 500MG BASE/VIAL</u>	<u>N50547</u>	<u>001</u>	
<u>AP</u>	+		<u>EQ 1GM BASE/VIAL</u>	<u>N50547</u>	<u>002</u>	
<u>AP</u>			<u>EQ 1GM BASE/VIAL</u>	<u>N62659</u>	<u>001</u>	Jan 13, 1987
<u>AP</u>	+		<u>EQ 2GM BASE/VIAL</u>	<u>N50547</u>	<u>003</u>	
<u>AP</u>			<u>EQ 2GM BASE/VIAL</u>	<u>N62659</u>	<u>002</u>	Jan 13, 1987
<u>AP</u>	+		<u>EQ 10GM BASE/VIAL</u>	<u>N50547</u>	<u>004</u>	Dec 29, 1983
		CLAFORAN IN DEXTROSE 5% IN PLASTIC CONTAINER				
	+	SANOFI AVENTIS US	EQ 20MG BASE/ML	N50596	002	May 20, 1985
	+		EQ 40MG BASE/ML	N50596	004	May 20, 1985

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 75 (of 370)

CEFOXITIN SODIUM

INJECTABLE; INJECTION

CEFOXITIN

<u>AP</u>	+	ABRAXIS PHARM	<u>EQ 1GM BASE/VIAL</u>	<u>N65012</u>	<u>001</u>	Jul 03, 2000
<u>AP</u>	+		<u>EQ 2GM BASE/VIAL</u>	<u>N65012</u>	<u>002</u>	Jul 03, 2000
<u>AP</u>	+		<u>EQ 10GM BASE/VIAL</u>	<u>N65011</u>	<u>001</u>	Jul 03, 2000
<u>AP</u>		BAXTER HLTHCARE	<u>EQ 1GM BASE/VIAL</u>	<u>N65051</u>	<u>001</u>	Sep 11, 2000
<u>AP</u>			<u>EQ 2GM BASE/VIAL</u>	<u>N65051</u>	<u>002</u>	Sep 11, 2000
<u>AP</u>			<u>EQ 10GM BASE/VIAL</u>	<u>N65050</u>	<u>001</u>	Sep 11, 2000
<u>AP</u>		ORCHID HLTHCARE	<u>EQ 1GM BASE/VIAL</u>	<u>N65313</u>	<u>001</u>	Jan 23, 2006
<u>AP</u>			<u>EQ 2GM BASE/VIAL</u>	<u>N65313</u>	<u>002</u>	Jan 23, 2006
<u>AP</u>			<u>EQ 10GM BASE/VIAL</u>	<u>N65312</u>	<u>001</u>	Feb 13, 2006

CEFOXITIN AND DEXTROSE IN DUPLEX CONTAINER

<u>AP</u>		B BRAUN	<u>EQ 1GM BASE/VIAL</u>	<u>N65214</u>	<u>001</u>	Mar 10, 2006
<u>AP</u>			<u>EQ 2GM BASE/VIAL</u>	<u>N65214</u>	<u>002</u>	Mar 10, 2006

MEFOXIN

<u>AP</u>		MERCK	<u>EQ 1GM BASE/VIAL</u>	<u>N62757</u>	<u>001</u>	Jan 08, 1987
<u>AP</u>			<u>EQ 2GM BASE/VIAL</u>	<u>N62757</u>	<u>002</u>	Jan 08, 1987
		MEFOXIN IN PLASTIC CONTAINER				
	+	MERCK	EQ 20MG BASE/ML	N63182	001	Jan 25, 1993
	+		EQ 40MG BASE/ML	N63182	002	Jan 25, 1993

CEFPODOXIME PROXETIL

FOR SUSPENSION; ORAL

CEFPODOXIME PROXETIL

<u>AB</u>		RANBAXY	<u>EQ 50MG BASE/5ML</u>	<u>N65082</u>	<u>001</u>	May 31, 2002
<u>AB</u>			<u>EQ 100MG BASE/5ML</u>	<u>N65082</u>	<u>002</u>	May 31, 2002
		<u>VANTIN</u>				
<u>AB</u>		PHARMACIA AND UPJOHN	<u>EQ 50MG BASE/5ML</u>	<u>N50675</u>	<u>001</u>	Aug 07, 1992
<u>AB</u>	+		<u>EQ 100MG BASE/5ML</u>	<u>N50675</u>	<u>002</u>	Aug 07, 1992

TABLET; ORAL

CEFPODOXIME PROXETIL

<u>AB</u>		RANBAXY	<u>EQ 100MG BASE</u>	<u>N65083</u>	<u>001</u>	Aug 20, 2003
<u>AB</u>			<u>EQ 200MG BASE</u>	<u>N65083</u>	<u>002</u>	Aug 20, 2003
		<u>VANTIN</u>				
<u>AB</u>		PHARMACIA AND UPJOHN	<u>EQ 100MG BASE</u>	<u>N50674</u>	<u>001</u>	Aug 07, 1992
<u>AB</u>	+		<u>EQ 200MG BASE</u>	<u>N50674</u>	<u>002</u>	Aug 07, 1992

CEFPROZIL

FOR SUSPENSION; ORAL

CEFPROZIL

<u>AB</u>		LUPIN	<u>125MG/5ML</u>	<u>N65261</u>	<u>001</u>	Dec 19, 2005
<u>AB</u>			<u>250MG/5ML</u>	<u>N65261</u>	<u>002</u>	Dec 19, 2005
<u>AB</u>		ORCHID HLTHCARE	<u>125MG/5ML</u>	<u>N65284</u>	<u>002</u>	Dec 30, 2005
<u>AB</u>			<u>250MG/5ML</u>	<u>N65284</u>	<u>001</u>	Dec 30, 2005
<u>AB</u>		RANBAXY	<u>125MG/5ML</u>	<u>N65202</u>	<u>001</u>	Jun 30, 2006
<u>AB</u>			<u>250MG/5ML</u>	<u>N65202</u>	<u>002</u>	Jun 30, 2006
<u>AB</u>		SANDOZ	<u>125MG/5ML</u>	<u>N65257</u>	<u>001</u>	Dec 08, 2005
<u>AB</u>			<u>250MG/5ML</u>	<u>N65257</u>	<u>002</u>	Dec 08, 2005
<u>AB</u>		TEVA PHARMS	<u>125MG/5ML</u>	<u>N65236</u>	<u>001</u>	Dec 08, 2005
<u>AB</u>			<u>250MG/5ML</u>	<u>N65236</u>	<u>002</u>	Dec 08, 2005

CEFZIL

<u>AB</u>		BRISTOL MYERS SQUIBB	<u>125MG/5ML</u>	<u>N50665</u>	<u>001</u>	Dec 23, 1991
<u>AB</u>	+		<u>250MG/5ML</u>	<u>N50665</u>	<u>002</u>	Dec 23, 1991

TABLET; ORAL

CEFPROZIL

<u>AB</u>		LUPIN	<u>250MG</u>	<u>N65276</u>	<u>001</u>	Dec 08, 2005
<u>AB</u>			<u>500MG</u>	<u>N65276</u>	<u>002</u>	Dec 08, 2005
<u>AB</u>		ORCHID HLTHCARE	<u>250MG</u>	<u>N65267</u>	<u>001</u>	Dec 19, 2005
<u>AB</u>			<u>500MG</u>	<u>N65267</u>	<u>002</u>	Dec 19, 2005

PRESCRIPTION DRUG PRODUCT LIST

3 - 76 (of 370)

CEFPROZIL

TABLET; ORAL

CEFPROZIL

<u>AB</u>	RANBAXY	<u>250MG</u>	<u>N65198</u>	<u>001</u>	Dec 13, 2006
<u>AB</u>		<u>500MG</u>	<u>N65198</u>	<u>002</u>	Dec 13, 2006
<u>AB</u>	SANDOZ	<u>250MG</u>	<u>N65235</u>	<u>001</u>	Nov 14, 2005
<u>AB</u>		<u>500MG</u>	<u>N65235</u>	<u>002</u>	Nov 14, 2005
<u>AB</u>	TEVA	<u>250MG</u>	<u>N65208</u>	<u>001</u>	Dec 06, 2005
<u>AB</u>		<u>500MG</u>	<u>N65208</u>	<u>002</u>	Dec 06, 2005

CEFZIL

<u>AB</u>	BRISTOL MYERS SQUIBB	<u>250MG</u>	<u>N50664</u>	<u>001</u>	Dec 23, 1991
<u>AB</u>	+	<u>500MG</u>	<u>N50664</u>	<u>002</u>	Dec 23, 1991

CEFTAZIDIME

INJECTABLE; INJECTION

CEFTAZIDIME

<u>AP</u>	ACS DOBFAR	<u>500MG/VIAL</u>	<u>N62640</u>	<u>001</u>	Nov 20, 1985
<u>AP</u>		<u>1GM/VIAL</u>	<u>N62640</u>	<u>002</u>	Nov 20, 1985
<u>AP</u>		<u>2GM/VIAL</u>	<u>N62640</u>	<u>003</u>	Nov 20, 1985
<u>AP</u>		<u>6GM/VIAL</u>	<u>N62640</u>	<u>004</u>	Feb 03, 1992

FORTAZ

<u>AP</u>	+	GLAXOSMITHKLINE	<u>500MG/VIAL</u>	<u>N50578</u>	<u>001</u>	Jul 19, 1985
<u>AP</u>	+		<u>1GM/VIAL</u>	<u>N50578</u>	<u>002</u>	Jul 19, 1985
<u>AP</u>	+		<u>2GM/VIAL</u>	<u>N50578</u>	<u>003</u>	Jul 19, 1985
<u>AP</u>	+		<u>6GM/VIAL</u>	<u>N50578</u>	<u>004</u>	Jul 19, 1985

TAZICEF

<u>AP</u>	HOSPIRA	<u>500MG/VIAL</u>	<u>N62662</u>	<u>001</u>	Mar 06, 1986
<u>AP</u>		<u>1GM/VIAL</u>	<u>N62662</u>	<u>002</u>	Mar 06, 1986
<u>AP</u>		<u>1GM/VIAL</u>	<u>N64032</u>	<u>001</u>	Oct 31, 1993
<u>AP</u>		<u>2GM/VIAL</u>	<u>N62662</u>	<u>003</u>	Mar 06, 1986
<u>AP</u>		<u>2GM/VIAL</u>	<u>N64032</u>	<u>002</u>	Oct 31, 1993
<u>AP</u>		<u>6GM/VIAL</u>	<u>N62662</u>	<u>004</u>	Mar 06, 1986

CEFTAZIDIME SODIUM

INJECTABLE; INJECTION

FORTAZ IN PLASTIC CONTAINER

+	GLAXOSMITHKLINE	EQ 20MG BASE/ML	N50634	002	Apr 28, 1989
+		EQ 40MG BASE/ML	N50634	003	Apr 28, 1989

CEFTIBUTEN DIHYDRATE

CAPSULE; ORAL

CEDAX

+	SHIONOGI	EQ 400MG BASE	N50685	002	Dec 20, 1995
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FOR SUSPENSION; ORAL

CEDAX

+	SHIONOGI	EQ 90MG BASE/5ML	N50686	001	Dec 20, 1995
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CEFTIZOXIME SODIUM

INJECTABLE; INJECTION

CEFIZOX

+	ASTELLAS	EQ 1GM BASE/VIAL	N50560	002	Sep 15, 1983
		EQ 1GM BASE/VIAL	N63294	002	Mar 31, 1994
+		EQ 2GM BASE/VIAL	N50560	003	Sep 15, 1983
		EQ 2GM BASE/VIAL	N63294	003	Mar 31, 1994
		EQ 10GM BASE/VIAL	N50560	005	Mar 19, 1993

CEFIZOX IN PLASTIC CONTAINER

+	ASTELLAS	EQ 20MG BASE/ML	N50589	003	Apr 13, 1995
+		EQ 40MG BASE/ML	N50589	004	Apr 13, 1995

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 77 (of 370)

CEFTRIAXONE SODIUM

INJECTABLE; IM-IV

CEFTRIAXONE

<u>AP</u>	ABRAXIS PHARM	<u>EQ 250MG BASE/VIAL</u>	<u>N65245</u>	<u>001</u>	Feb 15, 2006
<u>AP</u>		<u>EQ 500MG BASE/VIAL</u>	<u>N65245</u>	<u>002</u>	Feb 15, 2006
<u>AP</u>		<u>EQ 1GM BASE/VIAL</u>	<u>N65245</u>	<u>003</u>	Feb 15, 2006
<u>AP</u>		<u>EQ 2GM BASE/VIAL</u>	<u>N65245</u>	<u>004</u>	Feb 15, 2006
<u>AP</u>	LUPIN	<u>EQ 250MG BASE/VIAL</u>	<u>N65125</u>	<u>001</u>	Sep 30, 2003
<u>AP</u>		<u>EQ 500MG BASE/VIAL</u>	<u>N65125</u>	<u>002</u>	Sep 30, 2003
<u>AP</u>		<u>EQ 1GM BASE/VIAL</u>	<u>N65125</u>	<u>003</u>	Sep 30, 2003
<u>AP</u>		<u>EQ 2GM BASE/VIAL</u>	<u>N65125</u>	<u>004</u>	Sep 30, 2003
<u>AP</u>	ORCHID HLTHCARE	<u>EQ 250MG BASE/VIAL</u>	<u>N65230</u>	<u>001</u>	Aug 02, 2005
<u>AP</u>		<u>EQ 500MG BASE/VIAL</u>	<u>N65230</u>	<u>002</u>	Aug 02, 2005
<u>AP</u>		<u>EQ 1GM BASE/VIAL</u>	<u>N65230</u>	<u>003</u>	Aug 02, 2005
<u>AP</u>		<u>EQ 2GM BASE/VIAL</u>	<u>N65230</u>	<u>004</u>	Aug 02, 2005
<u>AP</u>	SANDOZ	<u>EQ 250MG BASE/VIAL</u>	<u>N65169</u>	<u>001</u>	May 09, 2005
<u>AP</u>		<u>EQ 500MG BASE/VIAL</u>	<u>N65169</u>	<u>002</u>	May 09, 2005
<u>AP</u>		<u>EQ 1GM BASE/VIAL</u>	<u>N65169</u>	<u>003</u>	May 09, 2005
<u>AP</u>		<u>EQ 2GM BASE/VIAL</u>	<u>N65169</u>	<u>004</u>	May 09, 2005
<u>CEFTRIAXONE SODIUM</u>					
<u>AP</u>	TEVA	<u>EQ 1GM BASE/VIAL</u>	<u>N65262</u>	<u>001</u>	Jun 29, 2006
<u>AP</u>		<u>EQ 2GM BASE/VIAL</u>	<u>N65262</u>	<u>002</u>	Jun 29, 2006
<u>ROCEPHIN</u>					
<u>AP</u>	+ HLR	<u>EQ 250MG BASE/VIAL</u>	<u>N50585</u>	<u>001</u>	Dec 21, 1984
<u>AP</u>	+	<u>EQ 500MG BASE/VIAL</u>	<u>N50585</u>	<u>002</u>	Dec 21, 1984
<u>AP</u>	+	<u>EQ 1GM BASE/VIAL</u>	<u>N50585</u>	<u>003</u>	Dec 21, 1984
<u>AP</u>	+	<u>EQ 2GM BASE/VIAL</u>	<u>N50585</u>	<u>004</u>	Dec 21, 1984

INJECTABLE; INJECTION

CEFTRIAXONE

<u>AP</u>	ABRAXIS PHARM	<u>EQ 10GM BASE/VIAL</u>	<u>N65252</u>	<u>001</u>	Feb 15, 2006
<u>AP</u>	LUPIN	<u>EQ 10GM BASE/VIAL</u>	<u>N65263</u>	<u>001</u>	Sep 12, 2006
<u>AP</u>	ORCHID HLTHCARE	<u>EQ 1GM BASE/VIAL</u>	<u>N65231</u>	<u>001</u>	Aug 02, 2005
<u>AP</u>		<u>EQ 2GM BASE/VIAL</u>	<u>N65231</u>	<u>002</u>	Aug 02, 2005
<u>AP</u>		<u>EQ 10GM BASE/VIAL</u>	<u>N65232</u>	<u>001</u>	Aug 02, 2005
<u>AP</u>	SANDOZ	<u>EQ 1GM BASE/VIAL</u>	<u>N65204</u>	<u>001</u>	May 03, 2005
<u>AP</u>		<u>EQ 2GM BASE/VIAL</u>	<u>N65204</u>	<u>002</u>	May 03, 2005
<u>AP</u>		<u>EQ 10GM BASE/VIAL</u>	<u>N65168</u>	<u>001</u>	May 17, 2005
<u>AP</u>	WOCKHARDT	<u>EQ 1GM BASE/VIAL</u>	<u>N65180</u>	<u>001</u>	May 12, 2006
<u>CEFTRIAXONE AND DEXTROSE IN DUPLEX CONTAINER</u>					
<u>AP</u>	+ B BRAUN	<u>EQ 1GM BASE/VIAL</u>	<u>N50796</u>	<u>001</u>	Apr 20, 2005
<u>AP</u>	+	<u>EQ 2GM BASE/VIAL</u>	<u>N50796</u>	<u>002</u>	Apr 20, 2005
<u>CEFTRIAXONE IN PLASTIC CONTAINER</u>					
<u>AP</u>	BAXTER HLTHCARE	<u>EQ 20MG BASE/ML</u>	<u>N65224</u>	<u>001</u>	Aug 23, 2005
<u>AP</u>		<u>EQ 40MG BASE/ML</u>	<u>N65224</u>	<u>002</u>	Aug 23, 2005
<u>CEFTRIAXONE SODIUM</u>					
<u>AP</u>	TEVA	<u>EQ 10GM BASE/VIAL</u>	<u>N65274</u>	<u>001</u>	May 01, 2006
<u>ROCEPHIN</u>					
<u>AP</u>	+ HLR	<u>EQ 1GM BASE/VIAL</u>	<u>N62654</u>	<u>002</u>	Apr 30, 1987
<u>AP</u>	+	<u>EQ 2GM BASE/VIAL</u>	<u>N62654</u>	<u>003</u>	Apr 30, 1987
<u>AP</u>	+	<u>EQ 10GM BASE/VIAL</u>	<u>N50585</u>	<u>005</u>	Dec 21, 1984
<u>ROCEPHIN W/ DEXTROSE IN PLASTIC CONTAINER</u>					
<u>AP</u>	+ HLR	<u>EQ 20MG BASE/ML</u>	<u>N50624</u>	<u>002</u>	Feb 11, 1987
<u>AP</u>	+	<u>EQ 40MG BASE/ML</u>	<u>N50624</u>	<u>003</u>	Feb 11, 1987

CEFUROXIME AXETIL

FOR SUSPENSION; ORAL

CEFTIN

	GLAXOSMITHKLINE	EQ 125MG BASE/5ML	N50672	001	Jun 30, 1994
	+	EQ 250MG BASE/5ML	N50672	002	Apr 29, 1997

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 78 (of 370)

CEFUROXIME AXETIL

TABLET; ORAL

CEFTIN

<u>AB</u>	GLAXOSMITHKLINE	<u>EQ 125MG BASE</u>	<u>N50605</u>	<u>001</u>	Dec 28, 1987
<u>AB</u>		<u>EQ 250MG BASE</u>	<u>N50605</u>	<u>002</u>	Dec 28, 1987
<u>AB</u>	+	<u>EQ 500MG BASE</u>	<u>N50605</u>	<u>003</u>	Dec 28, 1987
<u>CEFUROXIME AXETIL</u>					
<u>AB</u>	APOTEX INC	<u>EQ 250MG BASE</u>	<u>N65069</u>	<u>001</u>	Oct 02, 2002
<u>AB</u>		<u>EQ 500MG BASE</u>	<u>N65069</u>	<u>002</u>	Oct 02, 2002
<u>AB</u>	AUROBINDO PHARMA LTD	<u>EQ 125MG BASE</u>	<u>N65308</u>	<u>001</u>	Mar 29, 2006
<u>AB</u>		<u>EQ 250MG BASE</u>	<u>N65308</u>	<u>002</u>	Mar 29, 2006
<u>AB</u>		<u>EQ 500MG BASE</u>	<u>N65308</u>	<u>003</u>	Mar 29, 2006
<u>AB</u>	LUPIN	<u>EQ 250MG BASE</u>	<u>N65135</u>	<u>001</u>	Jul 25, 2003
<u>AB</u>		<u>EQ 500MG BASE</u>	<u>N65135</u>	<u>002</u>	Jul 25, 2003
<u>AB</u>	RANBAXY	<u>EQ 125MG BASE</u>	<u>N65043</u>	<u>003</u>	Feb 15, 2002
<u>AB</u>		<u>EQ 125MG BASE</u>	<u>N65118</u>	<u>001</u>	Apr 25, 2003
<u>AB</u>		<u>EQ 250MG BASE</u>	<u>N65043</u>	<u>002</u>	Feb 15, 2002
<u>AB</u>		<u>EQ 250MG BASE</u>	<u>N65118</u>	<u>002</u>	Apr 25, 2003
<u>AB</u>		<u>EQ 500MG BASE</u>	<u>N65043</u>	<u>001</u>	Feb 15, 2002
<u>AB</u>		<u>EQ 500MG BASE</u>	<u>N65118</u>	<u>003</u>	Apr 25, 2003
<u>AB</u>	SANDOZ	<u>EQ 250MG BASE</u>	<u>N65126</u>	<u>001</u>	Oct 28, 2003
<u>AB</u>		<u>EQ 500MG BASE</u>	<u>N65126</u>	<u>002</u>	Oct 28, 2003
<u>AB</u>	TEVA	<u>EQ 250MG BASE</u>	<u>N65190</u>	<u>001</u>	Oct 18, 2004
<u>AB</u>		<u>EQ 500MG BASE</u>	<u>N65190</u>	<u>002</u>	Oct 18, 2004
<u>AB</u>	WOCKHARDT	<u>EQ 125MG BASE</u>	<u>N65166</u>	<u>001</u>	Jul 29, 2005
<u>AB</u>		<u>EQ 250MG BASE</u>	<u>N65166</u>	<u>002</u>	Jul 29, 2005
<u>AB</u>		<u>EQ 500MG BASE</u>	<u>N65166</u>	<u>003</u>	Jul 29, 2005

CEFUROXIME SODIUM

INJECTABLE; IM-IV

CEFUROXIME

<u>AB</u>	ABRAXIS PHARM	<u>EQ 750MG BASE/VIAL</u>	<u>N65001</u>	<u>001</u>	May 30, 2001
<u>AB</u>	HIKMA FARMACEUTICA	<u>EQ 750MG BASE/VIAL</u>	<u>N65048</u>	<u>001</u>	Jan 09, 2004
<u>AB</u>	TEVA	<u>EQ 750MG BASE/VIAL</u>	<u>N64192</u>	<u>002</u>	Apr 16, 1998

CEFUROXIME SODIUM

<u>AB</u>	HANFORD GC	<u>EQ 750MG BASE/VIAL</u>	<u>N64125</u>	<u>001</u>	May 30, 1997
<u>AB</u>	MARSAM PHARMS LLC	<u>EQ 750MG BASE/VIAL</u>	<u>N64035</u>	<u>001</u>	Feb 26, 1993

ZINACEF

<u>AB</u>	+	GLAXOSMITHKLINE	<u>EQ 750MG BASE/VIAL</u>	<u>N50558</u>	<u>002</u>	Oct 19, 1983
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INJECTABLE; INJECTION

CEFUROXIME

<u>AP</u>	ABRAXIS PHARM	<u>EQ 1.5GM BASE/VIAL</u>	<u>N65001</u>	<u>002</u>	May 30, 2001
<u>AP</u>	HIKMA FARMACEUTICA	<u>EQ 1.5GM BASE/VIAL</u>	<u>N65048</u>	<u>002</u>	Jan 09, 2004
<u>AP</u>		<u>EQ 7.5GM BASE/VIAL</u>	<u>N65046</u>	<u>001</u>	Jan 09, 2004
<u>AP</u>	TEVA	<u>EQ 1.5GM BASE/VIAL</u>	<u>N64192</u>	<u>001</u>	Apr 16, 1998
<u>AP</u>		<u>EQ 7.5GM BASE/VIAL</u>	<u>N64191</u>	<u>001</u>	Apr 16, 1998

CEFUROXIME AND DEXTROSE IN DUPLEX CONTAINER

<u>AP</u>	+	B BRAUN	<u>EQ 750MG BASE/VIAL</u>	<u>N50780</u>	<u>001</u>	Feb 21, 2001
<u>AP</u>	+		<u>EQ 1.5GM BASE/VIAL</u>	<u>N50780</u>	<u>002</u>	Feb 21, 2001

CEFUROXIME SODIUM

<u>AP</u>	ABRAXIS PHARM	<u>EQ 7.5GM BASE/VIAL</u>	<u>N65002</u>	<u>001</u>	Sep 28, 1998
<u>AP</u>	HANFORD GC	<u>EQ 1.5GM BASE/VIAL</u>	<u>N64125</u>	<u>002</u>	May 30, 1997
<u>AP</u>		<u>EQ 7.5GM BASE/VIAL</u>	<u>N64124</u>	<u>001</u>	May 30, 1997
<u>AP</u>	MARSAM PHARMS LLC	<u>EQ 1.5GM BASE/VIAL</u>	<u>N64035</u>	<u>002</u>	Feb 26, 1993
<u>AP</u>		<u>EQ 7.5GM BASE/VIAL</u>	<u>N64036</u>	<u>001</u>	Feb 26, 1993

ZINACEF

<u>AP</u>	+	GLAXOSMITHKLINE	<u>EQ 1.5GM BASE/VIAL</u>	<u>N50558</u>	<u>003</u>	Oct 19, 1983
<u>AP</u>	+		<u>EQ 7.5GM BASE/VIAL</u>	<u>N50558</u>	<u>004</u>	Oct 23, 1986

ZINACEF IN PLASTIC CONTAINER

	+	GLAXOSMITHKLINE	EQ 15MG BASE/ML	N50643	001	Apr 28, 1989
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PRESCRIPTION DRUG PRODUCT LIST

3 - 79 (of 370)

CEFUROXIME SODIUM

INJECTABLE; INJECTION

ZINACEF IN PLASTIC CONTAINER

+ GLAXOSMITHKLINE	EQ 30MG BASE/ML	N50643	002	Apr 28, 1989
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CELECOXIB

CAPSULE; ORAL

CELEBREX

GD SEARLE	100MG	N20998	001	Dec 31, 1998
	200MG	N20998	002	Dec 31, 1998
+	400MG	N20998	003	Aug 29, 2002

CEPHALEXIN

CAPSULE; ORAL

CEPHALEXIN

<u>AB</u> APOTHECON	<u>EQ 250MG BASE</u>	<u>N63063</u>	<u>001</u>	Sep 29, 1989
<u>AB</u>	<u>EQ 500MG BASE</u>	<u>N63063</u>	<u>002</u>	Sep 29, 1989
<u>AB</u> AUROBINDO PHARMA LTD	<u>EQ 250MG BASE</u>	<u>N65253</u>	<u>001</u>	Nov 16, 2005
<u>AB</u>	<u>EQ 500MG BASE</u>	<u>N65253</u>	<u>002</u>	Nov 16, 2005
<u>AB</u> BARR	<u>EQ 250MG BASE</u>	<u>N62773</u>	<u>001</u>	Jun 26, 1987
<u>AB</u>	<u>EQ 500MG BASE</u>	<u>N62775</u>	<u>001</u>	Apr 22, 1987
<u>AB</u> BELCHER	<u>EQ 250MG BASE</u>	<u>N62713</u>	<u>001</u>	Jul 15, 1988
<u>AB</u>	<u>EQ 500MG BASE</u>	<u>N62713</u>	<u>002</u>	Jul 15, 1988
<u>AB</u> CEPH INTL	<u>EQ 250MG BASE</u>	<u>N62118</u>	<u>001</u>	
<u>AB</u>	<u>EQ 500MG BASE</u>	<u>N62118</u>	<u>002</u>	
<u>AB</u> HIKMA	<u>EQ 250MG BASE</u>	<u>N65215</u>	<u>001</u>	Jan 24, 2006
<u>AB</u>	<u>EQ 500MG BASE</u>	<u>N65215</u>	<u>002</u>	Jan 24, 2006
<u>AB</u> IVAX PHARMS	<u>EQ 250MG BASE</u>	<u>N61969</u>	<u>001</u>	
<u>AB</u>	<u>EQ 500MG BASE</u>	<u>N61969</u>	<u>002</u>	
<u>AB</u> LUPIN	<u>EQ 250MG BASE</u>	<u>N65229</u>	<u>001</u>	Nov 25, 2005
<u>AB</u>	<u>EQ 500MG BASE</u>	<u>N65229</u>	<u>002</u>	Nov 25, 2005
<u>AB</u> ORCHID HLTHCARE	<u>EQ 250MG BASE</u>	<u>N65248</u>	<u>001</u>	Jun 28, 2005
<u>AB</u>	<u>EQ 500MG BASE</u>	<u>N65248</u>	<u>002</u>	Jun 28, 2005
<u>AB</u> RANBAXY	<u>EQ 250MG BASE</u>	<u>N65007</u>	<u>001</u>	Sep 16, 1999
<u>AB</u>	<u>EQ 500MG BASE</u>	<u>N65007</u>	<u>002</u>	Sep 16, 1999
<u>AB</u> STEVENS J	<u>EQ 250MG BASE</u>	<u>N62870</u>	<u>001</u>	Mar 17, 1988
<u>AB</u> SUN PHARM INDS (IN)	<u>EQ 250MG BASE</u>	<u>N62791</u>	<u>001</u>	Jun 11, 1987
<u>AB</u>	<u>EQ 500MG BASE</u>	<u>N62791</u>	<u>002</u>	Jun 11, 1987
<u>AB</u> TEVA	<u>EQ 250MG BASE</u>	<u>N62702</u>	<u>001</u>	Feb 13, 1987
<u>AB</u>	<u>EQ 250MG BASE</u>	<u>N62760</u>	<u>001</u>	Apr 24, 1987
<u>AB</u>	<u>EQ 500MG BASE</u>	<u>N62702</u>	<u>002</u>	Feb 13, 1987
<u>AB</u>	<u>EQ 500MG BASE</u>	<u>N62761</u>	<u>001</u>	Apr 24, 1987
<u>AB</u> YUNG SHIN PHARM	<u>EQ 250MG BASE</u>	<u>N65152</u>	<u>001</u>	Feb 24, 2005
<u>AB</u>	<u>EQ 500MG BASE</u>	<u>N65152</u>	<u>002</u>	Feb 24, 2005
<u>KEFLEX</u>				
<u>AB</u> ADVANCIS PHARM	<u>EQ 250MG BASE</u>	<u>N50405</u>	<u>002</u>	
<u>AB</u>	<u>EQ 500MG BASE</u>	<u>N50405</u>	<u>003</u>	
	EQ 333MG BASE	N50405	004	May 12, 2006
+	EQ 750MG BASE	N50405	005	May 12, 2006

FOR SUSPENSION; ORAL

CEPHALEXIN

<u>AB</u> BARR	<u>EQ 250MG BASE/5ML</u>	<u>N62777</u>	<u>001</u>	Aug 06, 1987
<u>AB</u> CEPH INTL	<u>EQ 125MG BASE/5ML</u>	<u>N62117</u>	<u>002</u>	
<u>AB</u> +	<u>EQ 250MG BASE/5ML</u>	<u>N62117</u>	<u>003</u>	
+	EQ 100MG BASE/ML	N62117	001	
<u>AB</u> LUPIN	<u>EQ 125MG BASE/5ML</u>	<u>N65234</u>	<u>001</u>	Aug 17, 2005
<u>AB</u>	<u>EQ 250MG BASE/5ML</u>	<u>N65234</u>	<u>002</u>	Aug 17, 2005
<u>AB</u> ORCHID HLTHCARE	<u>EQ 125MG BASE/5ML</u>	<u>N65326</u>	<u>001</u>	Jul 10, 2006
<u>AB</u>	<u>EQ 250MG BASE/5ML</u>	<u>N65326</u>	<u>002</u>	Jul 10, 2006
<u>AB</u> RANBAXY	<u>EQ 125MG BASE/5ML</u>	<u>N65081</u>	<u>001</u>	Jul 27, 2001

PRESCRIPTION DRUG PRODUCT LIST

3 - 80 (of 370)

CEPHALEXIN

FOR SUSPENSION; ORAL

CEPHALEXIN

<u>AB</u>	RANBAXY	<u>EQ 250MG BASE/5ML</u>	<u>N65081</u>	<u>002</u>	Jul 27, 2001
<u>AB</u>	TEVA	<u>EQ 125MG BASE/5ML</u>	<u>N62703</u>	<u>001</u>	Feb 13, 1987
<u>AB</u>		<u>EQ 125MG BASE/5ML</u>	<u>N62767</u>	<u>001</u>	Jun 16, 1987
<u>AB</u>		<u>EQ 250MG BASE/5ML</u>	<u>N62703</u>	<u>002</u>	Feb 13, 1987
<u>AB</u>		<u>EQ 250MG BASE/5ML</u>	<u>N62768</u>	<u>001</u>	Jun 16, 1987

TABLET; ORAL
CEPHALEXIN

	TEVA	EQ 250MG BASE	N63023	001	Jan 12, 1989
+		EQ 500MG BASE	N63024	001	Jan 12, 1989

TABLET, FOR SUSPENSION; ORAL
PANIXINE DISPERDOSE

	RANBAXY	EQ 125MG BASE	N65100	002	Sep 11, 2003
+		EQ 250MG BASE	N65100	001	Sep 11, 2003

CEPHRADINE

CAPSULE; ORAL
CEPHRADINE

	TEVA	250MG	N62683	001	Jan 09, 1987
+		500MG	N62683	002	Jan 09, 1987

FOR SUSPENSION; ORAL

ANSPOR

<u>AB</u>	GLAXOSMITHKLINE	<u>125MG/5ML</u>	<u>N61866</u>	<u>001</u>	
<u>AB</u>		<u>250MG/5ML</u>	<u>N61866</u>	<u>002</u>	

CEPHRADINE

<u>AB</u>	TEVA	<u>125MG/5ML</u>	<u>N62693</u>	<u>001</u>	Jan 09, 1987
<u>AB</u>		<u>250MG/5ML</u>	<u>N62693</u>	<u>002</u>	Jan 09, 1987

VELOSEF '125'

<u>AB</u>	+	APOTHECON	<u>125MG/5ML</u>	<u>N61763</u>	<u>001</u>
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VELOSEF '250'

<u>AB</u>	+	APOTHECON	<u>250MG/5ML</u>	<u>N61763</u>	<u>002</u>
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CETIRIZINE HYDROCHLORIDE

SYRUP; ORAL
ZYRTEC

+	PFIZER	5MG/5ML	N20346	001	Sep 27, 1996
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TABLET; ORAL
ZYRTEC

	PFIZER	5MG	N19835	001	Dec 08, 1995
+		10MG	N19835	002	Dec 08, 1995

TABLET, CHEWABLE; ORAL
ZYRTEC

	PFIZER	5MG	N21621	001	Mar 16, 2004
+		10MG	N21621	002	Mar 16, 2004

CETIRIZINE HYDROCHLORIDE; PSEUDOEPHEDRINE HYDROCHLORIDE

TABLET, EXTENDED RELEASE; ORAL
ZYRTEC-D 12 HOUR

+	PFIZER	5MG;120MG	N21150	001	Aug 10, 2001
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CETRORELIX

INJECTABLE; INJECTION
CETROTIDE

+	SERONO INC	EQ 0.25MG BASE/ML	N21197	001	Aug 11, 2000
+		EQ 3MG BASE/ML	N21197	002	Aug 11, 2000

PRESCRIPTION DRUG PRODUCT LIST

3 - 81 (of 370)

CEVIMELINE HYDROCHLORIDE

CAPSULE; ORAL
EVOXAC

+ DAIICHI	EQ 30MG BASE	N20989	002	Jan 11, 2000
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CHLORAMBUCIL

TABLET; ORAL
LEUKERAN

+ SMITHKLINE BEECHAM	2MG	N10669	002	
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CHLORAMPHENICOL

OINTMENT; OPHTHALMIC
CHLORAMPHENICOL

+ ALTANA	1%	N60133	001	
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CHLORAMPHENICOL SODIUM SUCCINATE

INJECTABLE; INJECTION

CHLORAMPHENICOL SODIUM SUCCINATE

<u>AP</u>	ABRAXIS PHARM	EQ 1GM BASE/VIAL	N62365	001	Aug 25, 1982
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CHLOROMYCETIN

<u>AP</u>	+ PARKEDALE	EQ 1GM BASE/VIAL	N50155	001	
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CHLORDIAZEPOXIDE HYDROCHLORIDE

CAPSULE; ORAL

CHLORDIAZEPOXIDE HYDROCHLORIDE

<u>AB</u>	BARR	5MG	N84768	001	
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<u>AB</u>		10MG	N83116	001	
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<u>AB</u>		25MG	N84769	001	
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<u>AB</u>	USL PHARMA	10MG	N84623	001	
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<u>AB</u>	WATSON LABS	5MG	N86383	001	
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<u>AB</u>		10MG	N86294	001	
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<u>AB</u>		25MG	N86382	001	
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LIBRIUM

<u>AB</u>	VALEANT PHARM INTL	5MG	N85461	001	
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<u>AB</u>		10MG	N85472	001	
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<u>AB</u>	+	25MG	N85475	001	
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CHLORHEXIDINE GLUCONATE

SOLUTION; DENTAL

CHLORHEXIDINE GLUCONATE

<u>AT</u>	ACTAVIS MID ATLANTIC	0.12%	N74291	001	Dec 28, 1995
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<u>AT</u>	HI TECH PHARMA	0.12%	N74356	001	May 07, 1996
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<u>AT</u>	JOHN O BUTLER CO	0.12%	N76434	001	Nov 29, 2005
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<u>AT</u>	MORTON GROVE	0.12%	N75006	001	Mar 03, 2004
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<u>AT</u>	NOVEX	0.12%	N75561	001	Nov 14, 2000
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<u>AT</u>	TEVA	0.12%	N74522	001	Dec 15, 1995
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PERIDEX

<u>AT</u>	+ ZILA	0.12%	N19028	001	Aug 13, 1986
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PERIOGARD

<u>AT</u>	COLGATE	0.12%	N73695	001	Jan 14, 1994
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TABLET; DENTAL

PERIOCHIP

+ DEXCEL PHARMA	2.5MG	N20774	001	May 15, 1998
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CHLOROPROCAINE HYDROCHLORIDE

INJECTABLE; INJECTION

CHLOROPROCAINE HYDROCHLORIDE

<u>AP</u>	BEDFORD	2%	N40273	001	Sep 09, 1998
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<u>AP</u>		3%	N40273	002	Sep 09, 1998
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27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 82 (of 370)

CHLOROPROCAINE HYDROCHLORIDE

INJECTABLE; INJECTION

CHLOROPROCAINE HYDROCHLORIDE

<u>AP</u>	HOSPIRA	<u>2%</u>	<u>N87447</u>	<u>001</u>	Apr 16, 1982
<u>AP</u>		<u>3%</u>	<u>N87446</u>	<u>001</u>	Apr 16, 1982
	<u>NESACAINE</u>				
<u>AP</u>	ABRAXIS BIOSCIENCE	<u>2%</u>	<u>N09435</u>	<u>002</u>	
	+	<u>1%</u>	N09435	001	
	<u>NESACAINE-MPF</u>				
<u>AP</u>	+ ABRAXIS BIOSCIENCE	<u>2%</u>	<u>N09435</u>	<u>006</u>	May 02, 1996
<u>AP</u>	+	<u>3%</u>	<u>N09435</u>	<u>007</u>	May 02, 1996

CHLOROQUINE PHOSPHATE

TABLET; ORAL

ARALEN

<u>AA</u>	+ SANOFI AVENTIS US	<u>EQ 300MG BASE</u>	<u>N06002</u>	<u>001</u>	
	<u>CHLOROQUINE PHOSPHATE</u>				
<u>AA</u>	IMPAX LABS	<u>EQ 150MG BASE</u>	<u>N80880</u>	<u>001</u>	
<u>AA</u>		<u>EQ 300MG BASE</u>	<u>N40516</u>	<u>001</u>	Aug 29, 2003
<u>AA</u>	+ WEST WARD	<u>EQ 150MG BASE</u>	<u>N83082</u>	<u>001</u>	
<u>AA</u>		<u>EQ 300MG BASE</u>	<u>N83082</u>	<u>002</u>	Sep 17, 1999

CHLOROTHIAZIDE

SUSPENSION; ORAL

DIURIL

	+ MERCK	250MG/5ML	N11870	001	
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TABLET; ORAL

CHLOROTHIAZIDE

<u>AB</u>	+ MYLAN	<u>250MG</u>	<u>N84388</u>	<u>001</u>	
<u>AB</u>		<u>500MG</u>	<u>N84217</u>	<u>001</u>	
<u>AB</u>	WEST WARD	<u>250MG</u>	<u>N86028</u>	<u>001</u>	Jul 14, 1982
<u>AB</u>		<u>500MG</u>	<u>N87736</u>	<u>001</u>	Jul 14, 1982

CHLOROTHIAZIDE SODIUM

INJECTABLE; INJECTION

DIURIL

	+ OVATION PHARMS	EQ 500MG BASE/VIAL	N11145	005	
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CHLOROXINE

SHAMPOO; TOPICAL

CAPITROL

	+ WESTWOOD SQUIBB	2%	N17594	001	
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CHLORPHENIRAMINE POLISTIREX; CODEINE POLISTIREX

SUSPENSION, EXTENDED RELEASE; ORAL

CODEPREX

	+ UCB INC	EQ 4MG MALEATE/5ML;EQ 20MG BASE/5ML	N21369	001	Jun 21, 2004
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CHLORPHENIRAMINE POLISTIREX; HYDROCODONE POLISTIREX

SUSPENSION, EXTENDED RELEASE; ORAL

TUSSIONEX PENNKINETIC

	+ UCB INC	EQ 8MG MALEATE/5ML;EQ 10MG BITARTRATE/5ML	N19111	001	Dec 31, 1987
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CHLORPROMAZINE HYDROCHLORIDE

CONCENTRATE; ORAL

CHLORPROMAZINE HYDROCHLORIDE

<u>AA</u>	ACTAVIS MID ATLANTIC	<u>100MG/ML</u>	<u>N86863</u>	<u>001</u>	
<u>AA</u>	PHARM ASSOC	<u>30MG/ML</u>	<u>N40231</u>	<u>001</u>	Dec 30, 1999

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 83 (of 370)

CHLORPROMAZINE HYDROCHLORIDE

CONCENTRATE; ORAL

CHLORPROMAZINE HYDROCHLORIDE

<u>AA</u>	PHARM ASSOC	<u>100MG/ML</u>	<u>N40224</u>	<u>001</u>	Jan 26, 1999
	<u>CHLORPROMAZINE HYDROCHLORIDE INTENSOL</u>				
<u>AA</u>	ROXANE	<u>30MG/ML</u>	<u>N88157</u>	<u>001</u>	Apr 27, 1983
<u>AA</u>		<u>100MG/ML</u>	<u>N88158</u>	<u>001</u>	Apr 27, 1983
	<u>SONAZINE</u>				
<u>AA</u>	SANDOZ	<u>30MG/ML</u>	<u>N80983</u>	<u>004</u>	
<u>AA</u>		<u>100MG/ML</u>	<u>N80983</u>	<u>005</u>	
	<u>THORAZINE</u>				
<u>AA</u>	+ GLAXOSMITHKLINE	<u>30MG/ML</u>	<u>N09149</u>	<u>032</u>	
<u>AA</u>	+	<u>100MG/ML</u>	<u>N09149</u>	<u>043</u>	

INJECTABLE; INJECTION

CHLORPROMAZINE HYDROCHLORIDE

<u>AP</u>	BAXTER HLTHCARE	<u>25MG/ML</u>	<u>N83329</u>	<u>001</u>	
	<u>THORAZINE</u>				
<u>AP</u>	+ GLAXOSMITHKLINE	<u>25MG/ML</u>	<u>N09149</u>	<u>011</u>	

SYRUP; ORAL

SONAZINE

<u>AA</u>	SANDOZ	<u>10MG/5ML</u>	<u>N83040</u>	<u>001</u>	
	<u>THORAZINE</u>				
<u>AA</u>	+ GLAXOSMITHKLINE	<u>10MG/5ML</u>	<u>N09149</u>	<u>022</u>	

TABLET; ORAL

CHLORPROMAZINE HYDROCHLORIDE

BP	SANDOZ	10MG	N80439	001	
BP		25MG	N80439	002	
BP		50MG	N80439	003	
BP	+	100MG	N80439	004	
BP		200MG	N80439	005	
BP	USL PHARMA	10MG	N83386	001	
BP		25MG	N84112	001	
BP		50MG	N84113	001	
BP		100MG	N84114	001	
BP		200MG	N84115	001	

CHLORPROPAMIDE

TABLET; ORAL

CHLORPROPAMIDE

<u>AB</u>	MYLAN	<u>100MG</u>	<u>N88548</u>	<u>001</u>	Jun 01, 1984
<u>AB</u>		<u>250MG</u>	<u>N88549</u>	<u>001</u>	Jun 01, 1984
<u>AB</u>	PAR PHARM	<u>100MG</u>	<u>N88175</u>	<u>001</u>	Feb 27, 1984
<u>AB</u>		<u>250MG</u>	<u>N88176</u>	<u>001</u>	Feb 27, 1984
<u>AB</u>	PLIVA	<u>100MG</u>	<u>N88921</u>	<u>001</u>	Apr 12, 1985
<u>AB</u>		<u>250MG</u>	<u>N88922</u>	<u>001</u>	Apr 12, 1985
<u>AB</u>	SANDOZ	<u>100MG</u>	<u>N88725</u>	<u>001</u>	Aug 31, 1984
<u>AB</u>		<u>250MG</u>	<u>N88726</u>	<u>001</u>	Aug 31, 1984
<u>AB</u>	WATSON LABS	<u>100MG</u>	<u>N88852</u>	<u>001</u>	Sep 26, 1984
<u>AB</u>		<u>250MG</u>	<u>N88826</u>	<u>001</u>	Sep 26, 1984
	<u>DIABINESE</u>				
<u>AB</u>	PFIZER	<u>100MG</u>	<u>N11641</u>	<u>003</u>	
<u>AB</u>	+	<u>250MG</u>	<u>N11641</u>	<u>006</u>	

CHLORTHALIDONE

TABLET; ORAL

CHLORTHALIDONE

<u>AB</u>	MYLAN	<u>25MG</u>	<u>N87180</u>	<u>001</u>	
<u>AB</u>	+	<u>50MG</u>	<u>N86831</u>	<u>001</u>	
<u>AB</u>	PLIVA	<u>25MG</u>	<u>N88902</u>	<u>001</u>	Sep 19, 1985

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 84 (of 370)

CHLORTHALIDONE

TABLET; ORAL

CHLORTHALIDONE

<u>AB</u>	PLIVA	<u>50MG</u>	<u>N88903</u>	<u>001</u>	Sep 19, 1985
	THALITONE				
	+ MONARCH PHARMS	15MG	N19574	001	Dec 20, 1988

CHLORTHALIDONE; CLONIDINE HYDROCHLORIDE

TABLET; ORAL

CLORPRES

<u>AB</u>	MYLAN	<u>15MG;0.1MG</u>	<u>N71323</u>	<u>001</u>	Feb 09, 1987
<u>AB</u>		<u>15MG;0.2MG</u>	<u>N71324</u>	<u>001</u>	Feb 09, 1987
	+	15MG;0.3MG	N71325	001	Feb 09, 1987

CHLORZOXAZONE

TABLET; ORAL

CHLORZOXAZONE

<u>AA</u>	ACTAVIS TOTOWA	<u>250MG</u>	<u>N88928</u>	<u>001</u>	May 08, 1987
<u>AA</u>		<u>500MG</u>	<u>N40113</u>	<u>001</u>	Sep 29, 1995
<u>AA</u>	BARR	<u>500MG</u>	<u>N89895</u>	<u>001</u>	May 04, 1988
<u>AA</u>	MUTUAL PHARM	<u>500MG</u>	<u>N89970</u>	<u>001</u>	Sep 27, 1990
<u>AA</u>	OHM LABS	<u>250MG</u>	<u>N81298</u>	<u>001</u>	Dec 29, 1993
<u>AA</u>		<u>500MG</u>	<u>N81299</u>	<u>001</u>	Dec 29, 1993
<u>AA</u>	PAR PHARM	<u>250MG</u>	<u>N87981</u>	<u>001</u>	Sep 20, 1983
<u>AA</u>	SANDOZ	<u>250MG</u>	<u>N89852</u>	<u>001</u>	May 04, 1988
<u>AA</u>		<u>500MG</u>	<u>N89853</u>	<u>001</u>	May 04, 1988
<u>AA</u>	TEVA	<u>500MG</u>	<u>N89859</u>	<u>001</u>	May 04, 1988
<u>AA</u>	WATSON LABS	<u>500MG</u>	<u>N40137</u>	<u>001</u>	Aug 09, 1996
<u>AA</u>		<u>500MG</u>	<u>N81040</u>	<u>001</u>	Aug 22, 1989
	<u>PARAFON FORTE DSC</u>				
<u>AA</u>	+ ORTHO MCNEIL PHARM	<u>500MG</u>	<u>N11529</u>	<u>002</u>	Jun 15, 1987
	<u>STRIFON FORTE DSC</u>				
<u>AA</u>	FERNDAL LABS	<u>500MG</u>	<u>N81008</u>	<u>001</u>	Dec 23, 1988

CHOLESTYRAMINE

POWDER; ORAL

CHOLESTYRAMINE

<u>AB</u>	PAR PHARM	<u>EQ 4GM RESIN/SCOOPFUL</u>	<u>N77204</u>	<u>002</u>	Aug 26, 2005
<u>AB</u>		<u>EQ 4GM RESIN/PACKET</u>	<u>N77204</u>	<u>001</u>	Aug 26, 2005
<u>AB</u>	SANDOZ	<u>EQ 4GM RESIN/SCOOPFUL</u>	<u>N74557</u>	<u>002</u>	Aug 15, 1996
<u>AB</u>		<u>EQ 4GM RESIN/PACKET</u>	<u>N74557</u>	<u>001</u>	Aug 15, 1996
<u>AB</u>	TEVA	<u>EQ 4GM RESIN/PACKET</u>	<u>N74347</u>	<u>001</u>	May 28, 1998
<u>AB</u>		<u>EQ 4GM RESIN/SCOOPFUL</u>	<u>N74347</u>	<u>002</u>	May 28, 1998
<u>AB</u>	TEVA PHARMS	<u>EQ 4GM RESIN/PACKET</u>	<u>N74554</u>	<u>001</u>	Oct 02, 1996
<u>AB</u>		<u>EQ 4GM RESIN/SCOOPFUL</u>	<u>N74554</u>	<u>002</u>	Oct 02, 1996

CHOLESTYRAMINE LIGHT

<u>AB</u>	PAR PHARM	<u>EQ 4GM RESIN/PACKET</u>	<u>N77203</u>	<u>001</u>	Aug 26, 2005
<u>AB</u>		<u>EQ 4GM RESIN/SCOOPFUL</u>	<u>N77203</u>	<u>002</u>	Aug 26, 2005
<u>AB</u>	SANDOZ	<u>EQ 4GM RESIN/SCOOPFUL</u>	<u>N74558</u>	<u>002</u>	Aug 15, 1996
<u>AB</u>		<u>EQ 4GM RESIN/PACKET</u>	<u>N74558</u>	<u>001</u>	Aug 15, 1996
<u>AB</u>	TEVA	<u>EQ 4GM RESIN/PACKET</u>	<u>N74348</u>	<u>001</u>	May 28, 1998
<u>AB</u>		<u>EQ 4GM RESIN/SCOOPFUL</u>	<u>N74348</u>	<u>002</u>	May 28, 1998
<u>AB</u>	TEVA PHARMS	<u>EQ 4GM RESIN/SCOOPFUL</u>	<u>N74555</u>	<u>002</u>	Sep 30, 1998
<u>AB</u>		<u>EQ 4GM RESIN/PACKET</u>	<u>N74555</u>	<u>001</u>	Sep 30, 1998

LOCHOLEST

<u>AB</u>	SANDOZ	<u>EQ 4GM RESIN/SCOOPFUL</u>	<u>N74561</u>	<u>002</u>	Aug 15, 1996
<u>AB</u>		<u>EQ 4GM RESIN/PACKET</u>	<u>N74561</u>	<u>001</u>	Aug 15, 1996

LOCHOLEST LIGHT

<u>AB</u>	SANDOZ	<u>EQ 4GM RESIN/SCOOPFUL</u>	<u>N74562</u>	<u>002</u>	Aug 15, 1996
<u>AB</u>		<u>EQ 4GM RESIN/PACKET</u>	<u>N74562</u>	<u>001</u>	Aug 15, 1996

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 85 (of 370)

CHOLESTYRAMINE

POWDER; ORAL

PREVALITE

<u>AB</u>	UPSHER SMITH	<u>EQ 4GM RESIN/PACKET</u>	<u>N73263</u>	<u>001</u>	Feb 22, 1996
<u>AB</u>		<u>EQ 4GM RESIN/SCOOPFUL</u>	<u>N73263</u>	<u>002</u>	Oct 30, 1997

QUESTRAN

<u>AB</u>	+ BRISTOL MYERS	<u>EQ 4GM RESIN/PACKET</u>	<u>N16640</u>	<u>001</u>	
<u>AB</u>		<u>EQ 4GM RESIN/SCOOPFUL</u>	<u>N16640</u>	<u>003</u>	

QUESTRAN LIGHT

<u>AB</u>	BRISTOL MYERS	<u>EQ 4GM RESIN/PACKET</u>	<u>N19669</u>	<u>001</u>	Dec 05, 1988
<u>AB</u>		<u>EQ 4GM RESIN/SCOOPFUL</u>	<u>N19669</u>	<u>003</u>	Dec 05, 1988

CHORIOGONADOTROPIN ALFA

INJECTABLE; SUBCUTANEOUS

OVIDREL

+ SERONO INC	EQ 0.25MG /0.5ML	N21149	002	Oct 06, 2003
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CHROMIC CHLORIDE

INJECTABLE; INJECTION

CHROMIC CHLORIDE IN PLASTIC CONTAINER

+ HOSPIRA	EQ 0.004MG CHROMIUM/ML	N18961	001	Jun 26, 1986
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CHROMIC PHOSPHATE, P-32

INJECTABLE; INJECTION

PHOSPHOCOL P32

+ MALLINCKRODT	5mCi/ML	N17084	001	
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CICLESONIDE

SPRAY, METERED; NASAL

OMNARIS

+ ALTANA PHARMA	0.05MG/INH	N22004	001	Oct 20, 2006
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CICLOPIROX

CREAM; TOPICAL

CICLOPIROX

<u>AB</u>	ALTANA	<u>0.77%</u>	<u>N76435</u>	<u>001</u>	Dec 29, 2004
<u>AB</u>	PERRIGO	<u>0.77%</u>	<u>N77364</u>	<u>001</u>	Mar 03, 2006
<u>AB</u>	TARO	<u>0.77%</u>	<u>N76790</u>	<u>001</u>	Apr 12, 2005

LOPROX

<u>AB</u>	+ MEDICIS	<u>0.77%</u>	<u>N18748</u>	<u>001</u>	Dec 30, 1982
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GEL; TOPICAL

LOPROX

+ MEDICIS	0.77%	N20519	001	Jul 21, 1997
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SHAMPOO; TOPICAL

LOPROX

+ MEDICIS	1%	N21159	001	Feb 28, 2003
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SOLUTION; TOPICAL

PENLAC

+ SANOFI AVENTIS US	8%	N21022	001	Dec 17, 1999
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SUSPENSION; TOPICAL

CICLOPIROX

<u>AB</u>	ALTANA	<u>0.77%</u>	<u>N76422</u>	<u>001</u>	Aug 06, 2004
<u>AB</u>	PERRIGO NEW YORK	<u>0.77%</u>	<u>N77676</u>	<u>001</u>	Dec 15, 2006
<u>AB</u>	TARO	<u>0.77%</u>	<u>N77092</u>	<u>001</u>	Aug 10, 2005

LOPROX

<u>AB</u>	+ MEDICIS	<u>0.77%</u>	<u>N19824</u>	<u>001</u>	Dec 30, 1988
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PRESCRIPTION DRUG PRODUCT LIST

3 - 86 (of 370)

CIDOFOVIR

INJECTABLE; INJECTION
VISTIDE

+ GILEAD	EQ 75MG BASE/ML	N20638	001	Jun 26, 1996
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CILASTATIN SODIUM; IMIPENEM

INJECTABLE; INJECTION
PRIMAXIN

+ MERCK	EQ 250MG BASE/VIAL; 250MG/VIAL	N50587	001	Nov 26, 1985
	EQ 250MG BASE/VIAL; 250MG/VIAL	N62756	001	Jan 08, 1987
+	EQ 500MG BASE/VIAL; 500MG/VIAL	N50587	002	Nov 26, 1985
	EQ 500MG BASE/VIAL; 500MG/VIAL	N50630	001	Dec 14, 1990
	EQ 500MG BASE/VIAL; 500MG/VIAL	N62756	002	Jan 08, 1987

CILOSTAZOL

TABLET; ORAL

CILOSTAZOL

<u>AB</u>	ACTAVIS TOTOWA	<u>100MG</u>	<u>N77028</u>	<u>002</u>	Nov 26, 2004
<u>AB</u>	ALPHAPHARM	<u>50MG</u>	<u>N77019</u>	<u>001</u>	Nov 23, 2004
<u>AB</u>		<u>100MG</u>	<u>N77019</u>	<u>002</u>	Nov 23, 2004
<u>AB</u>	APOTEX INC	<u>50MG</u>	<u>N77030</u>	<u>001</u>	Dec 10, 2004
<u>AB</u>		<u>100MG</u>	<u>N77030</u>	<u>002</u>	Dec 10, 2004
<u>AB</u>	COREPHARMA	<u>50MG</u>	<u>N77150</u>	<u>001</u>	Mar 11, 2005
<u>AB</u>		<u>100MG</u>	<u>N77022</u>	<u>001</u>	Nov 23, 2004
<u>AB</u>	IVAX PHARMS	<u>100MG</u>	<u>N77020</u>	<u>002</u>	Mar 01, 2005
<u>AB</u>	MUTUAL PHARM	<u>50MG</u>	<u>N77208</u>	<u>002</u>	Mar 29, 2006
<u>AB</u>		<u>100MG</u>	<u>N77208</u>	<u>001</u>	Mar 29, 2006
<u>AB</u>	MYLAN	<u>50MG</u>	<u>N77323</u>	<u>002</u>	Apr 20, 2006
<u>AB</u>		<u>100MG</u>	<u>N77323</u>	<u>001</u>	Apr 20, 2006
<u>AB</u>	ROXANE	<u>50MG</u>	<u>N77024</u>	<u>001</u>	May 17, 2005
<u>AB</u>		<u>100MG</u>	<u>N77024</u>	<u>002</u>	May 17, 2005
<u>AB</u>	SANDOZ	<u>50MG</u>	<u>N77310</u>	<u>001</u>	Nov 08, 2005
<u>AB</u>		<u>100MG</u>	<u>N77021</u>	<u>001</u>	Nov 23, 2004
<u>AB</u>	TEVA	<u>50MG</u>	<u>N77027</u>	<u>001</u>	Nov 24, 2004
<u>AB</u>		<u>100MG</u>	<u>N77027</u>	<u>002</u>	Nov 24, 2004

PLETAL

<u>AB</u>	+ OTSUKA	<u>50MG</u>	<u>N20863</u>	<u>001</u>	Jan 15, 1999
<u>AB</u>	+	<u>100MG</u>	<u>N20863</u>	<u>002</u>	Jan 15, 1999

CIMETIDINE

TABLET; ORAL

CIMETIDINE

<u>AB</u>	CLONMEL HLTHCARE	<u>300MG</u>	<u>N74340</u>	<u>001</u>	Jun 23, 1995
<u>AB</u>		<u>400MG</u>	<u>N74340</u>	<u>002</u>	Jun 23, 1995
<u>AB</u>		<u>800MG</u>	<u>N74339</u>	<u>001</u>	Jun 23, 1995
<u>AB</u>	IVAX PHARMS	<u>200MG</u>	<u>N74424</u>	<u>001</u>	Jul 28, 1995
<u>AB</u>		<u>300MG</u>	<u>N74424</u>	<u>002</u>	Jul 28, 1995
<u>AB</u>		<u>400MG</u>	<u>N74424</u>	<u>003</u>	Jul 28, 1995
<u>AB</u>		<u>800MG</u>	<u>N74402</u>	<u>001</u>	May 30, 1995
<u>AB</u>		<u>800MG</u>	<u>N74424</u>	<u>004</u>	Jul 28, 1995
<u>AB</u>	LEK PHARMS	<u>300MG</u>	<u>N74250</u>	<u>002</u>	Jun 29, 1995
<u>AB</u>		<u>400MG</u>	<u>N74250</u>	<u>003</u>	Jun 29, 1995
<u>AB</u>		<u>800MG</u>	<u>N74250</u>	<u>004</u>	Jun 29, 1995
<u>AB</u>	MYLAN	<u>200MG</u>	<u>N74246</u>	<u>001</u>	May 17, 1994
<u>AB</u>		<u>300MG</u>	<u>N74246</u>	<u>002</u>	May 17, 1994
<u>AB</u>		<u>400MG</u>	<u>N74246</u>	<u>003</u>	May 17, 1994
<u>AB</u>		<u>800MG</u>	<u>N74246</u>	<u>004</u>	May 17, 1994
<u>AB</u>	PLIVA	<u>200MG</u>	<u>N74568</u>	<u>001</u>	Feb 27, 1997
<u>AB</u>		<u>300MG</u>	<u>N74568</u>	<u>002</u>	Feb 27, 1997
<u>AB</u>		<u>400MG</u>	<u>N74568</u>	<u>003</u>	Feb 27, 1997

PRESCRIPTION DRUG PRODUCT LIST

3 - 87 (of 370)

CIMETIDINE

TABLET; ORAL

CIMETIDINE

<u>AB</u>	PLIVA	<u>800MG</u>	<u>N74566</u>	<u>001</u>	Feb 27, 1997
<u>AB</u>	SANDOZ	<u>200MG</u>	<u>N74100</u>	<u>001</u>	Jan 31, 1995
<u>AB</u>		<u>200MG</u>	<u>N74506</u>	<u>001</u>	Jan 24, 1996
<u>AB</u>		<u>300MG</u>	<u>N74100</u>	<u>002</u>	Jan 31, 1995
<u>AB</u>		<u>300MG</u>	<u>N74506</u>	<u>002</u>	Jan 24, 1996
<u>AB</u>		<u>400MG</u>	<u>N74100</u>	<u>003</u>	Jan 31, 1995
<u>AB</u>		<u>400MG</u>	<u>N74506</u>	<u>003</u>	Jan 24, 1996
<u>AB</u>		<u>800MG</u>	<u>N74100</u>	<u>004</u>	Jan 31, 1995
<u>AB</u>		<u>800MG</u>	<u>N74506</u>	<u>004</u>	Jan 24, 1996
<u>AB</u>	TEVA	<u>200MG</u>	<u>N74151</u>	<u>001</u>	May 17, 1994
<u>AB</u>		<u>200MG</u>	<u>N74365</u>	<u>001</u>	Feb 28, 1995
<u>AB</u>		<u>300MG</u>	<u>N74151</u>	<u>002</u>	May 17, 1994
<u>AB</u>		<u>300MG</u>	<u>N74365</u>	<u>002</u>	Feb 28, 1995
<u>AB</u>		<u>400MG</u>	<u>N74151</u>	<u>003</u>	May 17, 1994
<u>AB</u>		<u>400MG</u>	<u>N74365</u>	<u>003</u>	Feb 28, 1995
<u>AB</u>		<u>800MG</u>	<u>N74365</u>	<u>004</u>	Feb 28, 1995
<u>AB</u>		<u>800MG</u>	<u>N74463</u>	<u>001</u>	May 17, 1994
<u>AB</u>	TORPHARM	<u>200MG</u>	<u>N74890</u>	<u>001</u>	Dec 18, 1998
<u>AB</u>		<u>300MG</u>	<u>N74890</u>	<u>002</u>	Dec 18, 1998
<u>AB</u>		<u>400MG</u>	<u>N74890</u>	<u>003</u>	Dec 18, 1998
<u>AB</u>		<u>800MG</u>	<u>N74890</u>	<u>004</u>	Dec 18, 1998
<u>AB</u>	WATSON LABS	<u>200MG</u>	<u>N74349</u>	<u>001</u>	Aug 30, 1996
<u>AB</u>		<u>300MG</u>	<u>N74349</u>	<u>002</u>	Aug 30, 1996
<u>AB</u>		<u>400MG</u>	<u>N74349</u>	<u>003</u>	Aug 30, 1996
<u>AB</u>		<u>800MG</u>	<u>N74316</u>	<u>001</u>	Feb 28, 1996
<u>TAGAMET</u>					
<u>AB</u>	GLAXOSMITHKLINE	<u>200MG</u>	<u>N17920</u>	<u>002</u>	
<u>AB</u>		<u>300MG</u>	<u>N17920</u>	<u>003</u>	
<u>AB</u>		<u>400MG</u>	<u>N17920</u>	<u>004</u>	Dec 14, 1983
<u>AB</u>	+	<u>800MG</u>	<u>N17920</u>	<u>005</u>	Apr 30, 1986

CIMETIDINE HYDROCHLORIDE

INJECTABLE; INJECTION

CIMETIDINE HYDROCHLORIDE

<u>AP</u>	CLONMEL HLTHCARE	<u>EQ 300MG BASE/2ML</u>	<u>N74428</u>	<u>001</u>	Apr 25, 1996
<u>AP</u>	HOSPIRA	<u>EQ 300MG BASE/2ML</u>	<u>N74344</u>	<u>001</u>	Jan 31, 1995
<u>AP</u>		<u>EQ 300MG BASE/2ML</u>	<u>N74345</u>	<u>001</u>	Jan 31, 1995
<u>AP</u>	SICOR PHARMS	<u>EQ 300MG BASE/2ML</u>	<u>N74252</u>	<u>001</u>	Nov 26, 1997
CIMETIDINE HYDROCHLORIDE IN SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER					
	+ HOSPIRA	EQ 6MG BASE/ML	N74269	001	Dec 27, 1994

SOLUTION; ORAL

CIMETIDINE HYDROCHLORIDE

<u>AA</u>	+ ACTAVIS MID ATLANTIC	<u>EQ 300MG BASE/5ML</u>	<u>N74176</u>	<u>001</u>	Jun 01, 1994
<u>AA</u>	DURAMED PHARMS BARR	<u>EQ 300MG BASE/5ML</u>	<u>N75110</u>	<u>001</u>	Jun 18, 1998
<u>AA</u>	HI TECH PHARMA	<u>EQ 300MG BASE/5ML</u>	<u>N74664</u>	<u>001</u>	Oct 28, 1997
<u>AA</u>	MORTON GROVE	<u>EQ 300MG BASE/5ML</u>	<u>N74757</u>	<u>001</u>	Oct 17, 1997
<u>AA</u>	NOVEX	<u>EQ 300MG BASE/5ML</u>	<u>N75560</u>	<u>001</u>	Mar 15, 2000
<u>AA</u>	PHARM ASSOC	<u>EQ 300MG BASE/5ML</u>	<u>N74553</u>	<u>001</u>	Jan 27, 1997
<u>AA</u>	TEVA	<u>EQ 300MG BASE/5ML</u>	<u>N74610</u>	<u>001</u>	Sep 26, 1996
<u>AA</u>	TEVA PHARMS	<u>EQ 300MG BASE/5ML</u>	<u>N74859</u>	<u>001</u>	Jul 09, 1998

CINACALCET HYDROCHLORIDE

TABLET; ORAL

SENSIPAR

	AMGEN	EQ 30MG BASE	N21688	001	Mar 08, 2004
		EQ 60MG BASE	N21688	002	Mar 08, 2004
	+	EQ 90MG BASE	N21688	003	Mar 08, 2004

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 88 (of 370)

CIPROFLOXACIN

FOR SUSPENSION; ORAL

CIPRO

BAYER PHARMS

250MG/5ML

N20780 001

Sep 26, 1997

+

500MG/5ML

N20780 002

Sep 26, 1997

INJECTABLE; INJECTION

CIPROAP + BAYER PHARMS400MG/40ML (10MG/ML)N19847 001

Dec 26, 1990

AP +200MG/20ML (10MG/ML)N19847 002

Dec 26, 1990

CIPRO IN DEXTROSE 5% IN PLASTIC CONTAINER

+ BAYER PHARMS

200MG/100ML

N19857 001

Dec 26, 1990

CIPROFLOXACINAP ABRAXIS PHARM200MG/20ML (10MG/ML)N76484 001

Aug 28, 2006

AP400MG/40ML (10MG/ML)N76484 002

Aug 28, 2006

AP

BEDFORD LABS

200MG/20ML (10MG/ML)N76992 001

Aug 28, 2006

AP400MG/40ML (10MG/ML)N76992 002

Aug 28, 2006

1200MG/120ML (10MG/ML)

N76993 001

Aug 28, 2006

AP

HOSPIRA

200MG/20ML (10MG/ML)N77245 001

Aug 28, 2006

AP400MG/40ML (10MG/ML)N77245 002

Aug 28, 2006

AP

SICOR PHARMS

200MG/20ML (10MG/ML)N77782 001

Aug 28, 2006

AP400MG/40ML (10MG/ML)N77782 002

Aug 28, 2006

SOLUTION/DROPS; OPHTHALMIC

CIPROFLOXACINAT NEXUS PHARMSEQ 0.3% BASEN77689 001

Dec 13, 2006

CIPROFLOXACIN HYDROCHLORIDE

OINTMENT; OPHTHALMIC

CILOXAN

+ ALCON

EQ 0.3% BASE

N20369 001

Mar 30, 1998

SOLUTION/DROPS; OPHTHALMIC

CILOXANAT + ALCONEQ 0.3% BASEN19992 001

Dec 31, 1990

CIPROFLOXACINAT BAUSCH AND LOMBEQ 0.3% BASEN76754 001

Jun 09, 2004

AT

HITECH PHARMA

EQ 0.3% BASEN76673 001

Jan 21, 2005

AT

NOVEX

EQ 0.3% BASEN75928 001

Jun 09, 2004

TABLET; ORAL

CIPROAB BAYER PHARMSEQ 100MG BASEN19537 001

Apr 08, 1996

ABEQ 250MG BASEN19537 002

Oct 22, 1987

ABEQ 500MG BASEN19537 003

Oct 22, 1987

AB +EQ 750MG BASEN19537 004

Oct 22, 1987

CIPROFLOXACINAB BARREQ 250MG BASEN74124 001

Jun 09, 2004

ABEQ 500MG BASEN74124 002

Jun 09, 2004

ABEQ 750MG BASEN74124 003

Jun 09, 2004

AB

CARLSBAD

EQ 250MG BASEN76126 002

Jun 09, 2004

ABEQ 500MG BASEN76126 003

Jun 09, 2004

ABEQ 750MG BASEN76126 004

Jun 09, 2004

AB

COBALT

EQ 100MG BASEN76794 001

Feb 10, 2005

ABEQ 250MG BASEN76794 002

Jun 09, 2004

ABEQ 500MG BASEN76794 003

Jun 09, 2004

ABEQ 750MG BASEN76794 004

Jun 09, 2004

AB

DR REDDYS LABS LTD

EQ 100MG BASEN75593 002

Jun 09, 2004

ABEQ 250MG BASEN75593 003

Jun 09, 2004

ABEQ 500MG BASEN75593 004

Jun 09, 2004

ABEQ 750MG BASEN75593 001

Jun 09, 2004

AB

GENPHARM

EQ 250MG BASEN75817 002

Jun 09, 2004

ABEQ 500MG BASEN75817 003

Jun 09, 2004

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 89 (of 370)

CIPROFLOXACIN HYDROCHLORIDE

TABLET; ORAL

CIPROFLOXACIN

<u>AB</u>	GENPHARM	<u>EQ 750MG BASE</u>	<u>N75817</u>	<u>004</u>	Jun 09, 2004
<u>AB</u>	HIKMA	<u>EQ 250MG BASE</u>	<u>N76558</u>	<u>002</u>	Jun 09, 2004
<u>AB</u>		<u>EQ 500MG BASE</u>	<u>N76558</u>	<u>003</u>	Jun 09, 2004
<u>AB</u>		<u>EQ 750MG BASE</u>	<u>N76558</u>	<u>004</u>	Jun 09, 2004
<u>AB</u>	IVAX PHARMS	<u>EQ 250MG BASE</u>	<u>N76089</u>	<u>002</u>	Jun 09, 2004
<u>AB</u>		<u>EQ 500MG BASE</u>	<u>N76089</u>	<u>003</u>	Jun 09, 2004
<u>AB</u>		<u>EQ 750MG BASE</u>	<u>N76089</u>	<u>004</u>	Jun 09, 2004
<u>AB</u>	MARTEC USA LLC	<u>EQ 250MG BASE</u>	<u>N76138</u>	<u>001</u>	Jun 09, 2004
<u>AB</u>		<u>EQ 500MG BASE</u>	<u>N76138</u>	<u>002</u>	Jun 09, 2004
<u>AB</u>		<u>EQ 750MG BASE</u>	<u>N76138</u>	<u>003</u>	Jun 09, 2004
<u>AB</u>	MYLAN	<u>EQ 250MG BASE</u>	<u>N75685</u>	<u>002</u>	Jun 09, 2004
<u>AB</u>		<u>EQ 500MG BASE</u>	<u>N75685</u>	<u>003</u>	Jun 09, 2004
<u>AB</u>		<u>EQ 750MG BASE</u>	<u>N75685</u>	<u>001</u>	Jun 09, 2004
<u>AB</u>	PLIVA	<u>EQ 100MG BASE</u>	<u>N76426</u>	<u>001</u>	Jun 15, 2005
<u>AB</u>		<u>EQ 250MG BASE</u>	<u>N76426</u>	<u>002</u>	Jun 15, 2005
<u>AB</u>		<u>EQ 500MG BASE</u>	<u>N76426</u>	<u>003</u>	Jun 15, 2005
<u>AB</u>		<u>EQ 750MG BASE</u>	<u>N76426</u>	<u>004</u>	Jun 15, 2005
<u>AB</u>	RANBAXY	<u>EQ 250MG BASE</u>	<u>N75747</u>	<u>001</u>	Jun 09, 2004
<u>AB</u>		<u>EQ 500MG BASE</u>	<u>N75747</u>	<u>002</u>	Jun 09, 2004
<u>AB</u>		<u>EQ 750MG BASE</u>	<u>N75747</u>	<u>003</u>	Jun 09, 2004
<u>AB</u>	SANDOZ	<u>EQ 100MG BASE</u>	<u>N75939</u>	<u>001</u>	Mar 03, 2005
<u>AB</u>		<u>EQ 250MG BASE</u>	<u>N75939</u>	<u>002</u>	Jun 09, 2004
<u>AB</u>		<u>EQ 250MG BASE</u>	<u>N76593</u>	<u>002</u>	Jun 09, 2004
<u>AB</u>		<u>EQ 500MG BASE</u>	<u>N75939</u>	<u>003</u>	Jun 09, 2004
<u>AB</u>		<u>EQ 500MG BASE</u>	<u>N76593</u>	<u>003</u>	Jun 09, 2004
<u>AB</u>		<u>EQ 750MG BASE</u>	<u>N75939</u>	<u>004</u>	Jun 09, 2004
<u>AB</u>		<u>EQ 750MG BASE</u>	<u>N76593</u>	<u>004</u>	Jun 09, 2004
<u>AB</u>	TARO	<u>EQ 100MG BASE</u>	<u>N76912</u>	<u>001</u>	Feb 18, 2005
<u>AB</u>		<u>EQ 250MG BASE</u>	<u>N76912</u>	<u>002</u>	Oct 06, 2004
<u>AB</u>		<u>EQ 500MG BASE</u>	<u>N76912</u>	<u>003</u>	Oct 06, 2004
<u>AB</u>		<u>EQ 750MG BASE</u>	<u>N76912</u>	<u>004</u>	Oct 06, 2004
<u>AB</u>	TEVA	<u>EQ 250MG BASE</u>	<u>N76136</u>	<u>001</u>	Jun 09, 2004
<u>AB</u>		<u>EQ 500MG BASE</u>	<u>N76136</u>	<u>002</u>	Jun 09, 2004
<u>AB</u>		<u>EQ 750MG BASE</u>	<u>N76136</u>	<u>003</u>	Jun 09, 2004
<u>AB</u>	TORPHARM	<u>EQ 250MG BASE</u>	<u>N76896</u>	<u>001</u>	Nov 04, 2004
<u>AB</u>		<u>EQ 500MG BASE</u>	<u>N76896</u>	<u>002</u>	Nov 04, 2004
<u>AB</u>		<u>EQ 750MG BASE</u>	<u>N76896</u>	<u>003</u>	Nov 04, 2004
<u>AB</u>	UNIQUE PHARM LABS	<u>EQ 250MG BASE</u>	<u>N76639</u>	<u>001</u>	Sep 10, 2004
<u>AB</u>		<u>EQ 500MG BASE</u>	<u>N76639</u>	<u>002</u>	Sep 10, 2004
<u>AB</u>		<u>EQ 750MG BASE</u>	<u>N76639</u>	<u>003</u>	Sep 10, 2004

TABLET, EXTENDED RELEASE; ORAL

PROQUIN XR

+	ESPRIT PHARMA	EQ 500MG BASE	N21744	001	May 19, 2005
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CIPROFLOXACIN HYDROCHLORIDE; HYDROCORTISONE

SUSPENSION/DROPS; OTIC

+	ALCON	EQ 0.2% BASE;1%	N20805	001	Feb 10, 1998
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CIPROFLOXACIN; CIPROFLOXACIN HYDROCHLORIDE

TABLET, EXTENDED RELEASE; ORAL

CIPRO XR

+	BAYER PHARMS	212.6MG;EQ 287.5MG BASE	N21473	001	Dec 13, 2002
+		425.2MG;EQ 574.9MG BASE	N21473	002	Aug 28, 2003

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 90 (of 370)

CIPROFLOXACIN; DEXAMETHASONE

SUSPENSION/DROPS; OTIC

+	ALCON	0.3%;0.1%	N21537	001	Jul 18, 2003
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CISATRACURIUM BESYLATE

INJECTABLE; INJECTION

NIMBEX

+	ABBOTT	EQ 2MG BASE/ML	N20551	001	Dec 15, 1995
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NIMBEX PRESERVATIVE FREE

+	ABBOTT	EQ 2MG BASE/ML	N20551	003	Dec 15, 1995
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+		EQ 10MG BASE/ML	N20551	002	Dec 15, 1995
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CISPLATIN

INJECTABLE; INJECTION

CISPLATIN

<u>AP</u>	ABRAXIS PHARM	<u>1MG/ML</u>	<u>N74735</u>	<u>001</u>	Jul 16, 1999
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<u>AP</u>	BEDFORD	<u>1MG/ML</u>	<u>N75036</u>	<u>001</u>	Nov 07, 2000
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<u>AP</u>		<u>10MG/VIAL</u>	<u>N74713</u>	<u>001</u>	Nov 14, 2000
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<u>AP</u>		<u>50MG/VIAL</u>	<u>N74713</u>	<u>002</u>	Nov 14, 2000
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<u>AP</u>	PHARMACHEMIE	<u>1MG/ML</u>	<u>N74656</u>	<u>001</u>	May 16, 2000
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<u>AP</u>	SICOR PHARMS	<u>1MG/ML</u>	<u>N74814</u>	<u>001</u>	May 16, 2000
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PLATINOL

<u>AP</u>	+	BRISTOL MYERS	<u>10MG/VIAL</u>	<u>N18057</u>	<u>001</u>	
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<u>AP</u>	+		<u>50MG/VIAL</u>	<u>N18057</u>	<u>002</u>	
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PLATINOL-AQ

<u>AP</u>	+	BRISTOL MYERS	<u>1MG/ML</u>	<u>N18057</u>	<u>004</u>	Nov 08, 1988
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CITALOPRAM HYDROBROMIDE

SOLUTION; ORAL

CELEXA

<u>AA</u>	+	FOREST LABS	<u>EQ 10MG BASE/5ML</u>	<u>N21046</u>	<u>001</u>	Dec 22, 1999
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CITALOPRAM HYDROBROMIDE

<u>AA</u>	APOTEX INC	<u>EQ 10MG BASE/5ML</u>	<u>N77601</u>	<u>001</u>	Nov 15, 2005
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<u>AA</u>	AUROBINDO PHARMA LTD	<u>EQ 10MG BASE/5ML</u>	<u>N77812</u>	<u>001</u>	Aug 28, 2006
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<u>AA</u>	ROXANE	<u>EQ 10MG BASE/5ML</u>	<u>N77043</u>	<u>001</u>	Dec 13, 2004
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<u>AA</u>	SILARX	<u>EQ 10MG BASE/5ML</u>	<u>N77629</u>	<u>001</u>	Jun 15, 2006
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TABLET; ORAL

CELEXA

<u>AB</u>	FOREST LABS	<u>EQ 10MG BASE</u>	<u>N20822</u>	<u>001</u>	Apr 27, 2000
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<u>AB</u>		<u>EQ 20MG BASE</u>	<u>N20822</u>	<u>002</u>	Jul 17, 1998
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<u>AB</u>	+		<u>EQ 40MG BASE</u>	<u>N20822</u>	<u>003</u>	Jul 17, 1998
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CITALOPRAM HYDROBROMIDE

<u>AB</u>	ACTAVIS ELIZABETH	<u>EQ 10MG BASE</u>	<u>N77033</u>	<u>001</u>	Oct 28, 2004
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<u>AB</u>		<u>EQ 20MG BASE</u>	<u>N77033</u>	<u>002</u>	Oct 28, 2004
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<u>AB</u>		<u>EQ 40MG BASE</u>	<u>N77033</u>	<u>003</u>	Oct 28, 2004
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<u>AB</u>	AKYMA PHARMS	<u>EQ 10MG BASE</u>	<u>N77045</u>	<u>003</u>	Apr 29, 2005
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<u>AB</u>		<u>EQ 20MG BASE</u>	<u>N77045</u>	<u>002</u>	Apr 29, 2005
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<u>AB</u>		<u>EQ 40MG BASE</u>	<u>N77045</u>	<u>001</u>	Apr 29, 2005
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<u>AB</u>	ALPHAPHARM	<u>EQ 10MG BASE</u>	<u>N77037</u>	<u>001</u>	Nov 05, 2004
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<u>AB</u>		<u>EQ 20MG BASE</u>	<u>N77037</u>	<u>002</u>	Nov 05, 2004
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<u>AB</u>		<u>EQ 40MG BASE</u>	<u>N77037</u>	<u>003</u>	Nov 05, 2004
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<u>AB</u>	APOTEX INC	<u>EQ 10MG BASE</u>	<u>N77046</u>	<u>001</u>	Nov 24, 2004
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<u>AB</u>		<u>EQ 20MG BASE</u>	<u>N77046</u>	<u>002</u>	Nov 24, 2004
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<u>AB</u>		<u>EQ 40MG BASE</u>	<u>N77046</u>	<u>003</u>	Nov 24, 2004
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<u>AB</u>	AUROBINDO	<u>EQ 10MG BASE</u>	<u>N77031</u>	<u>001</u>	Oct 28, 2004
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<u>AB</u>		<u>EQ 20MG BASE</u>	<u>N77031</u>	<u>002</u>	Oct 28, 2004
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<u>AB</u>		<u>EQ 40MG BASE</u>	<u>N77031</u>	<u>003</u>	Oct 28, 2004
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<u>AB</u>	CARACO	<u>EQ 10MG BASE</u>	<u>N77032</u>	<u>001</u>	Nov 12, 2004
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<u>AB</u>		<u>EQ 20MG BASE</u>	<u>N77032</u>	<u>002</u>	Nov 12, 2004
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27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 91 (of 370)

CITALOPRAM HYDROBROMIDE

TABLET; ORAL

CITALOPRAM HYDROBROMIDE

<u>AB</u>	CARACO	<u>EQ 40MG BASE</u>	<u>N77032</u>	<u>003</u>	Nov 12, 2004
<u>AB</u>	COBALT	<u>EQ 10MG BASE</u>	<u>N77034</u>	<u>001</u>	Jun 30, 2005
<u>AB</u>		<u>EQ 20MG BASE</u>	<u>N77034</u>	<u>002</u>	Jun 30, 2005
<u>AB</u>		<u>EQ 40MG BASE</u>	<u>N77034</u>	<u>003</u>	Jun 30, 2005
<u>AB</u>	COREPHARMA	<u>EQ 10MG BASE</u>	<u>N77036</u>	<u>001</u>	Oct 28, 2004
<u>AB</u>		<u>EQ 20MG BASE</u>	<u>N77036</u>	<u>002</u>	Oct 28, 2004
<u>AB</u>		<u>EQ 40MG BASE</u>	<u>N77036</u>	<u>003</u>	Oct 28, 2004
<u>AB</u>	DR REDDYS LABS LTD	<u>EQ 10MG BASE</u>	<u>N77038</u>	<u>001</u>	Oct 28, 2004
<u>AB</u>		<u>EQ 20MG BASE</u>	<u>N77038</u>	<u>002</u>	Oct 28, 2004
<u>AB</u>		<u>EQ 40MG BASE</u>	<u>N77038</u>	<u>003</u>	Oct 28, 2004
<u>AB</u>	INTERPHARM	<u>EQ 10MG BASE</u>	<u>N77289</u>	<u>001</u>	Nov 30, 2006
<u>AB</u>		<u>EQ 20MG BASE</u>	<u>N77289</u>	<u>002</u>	Nov 30, 2006
<u>AB</u>		<u>EQ 40MG BASE</u>	<u>N77289</u>	<u>003</u>	Nov 30, 2006
<u>AB</u>	INVAGEN PHARMS	<u>EQ 10MG BASE</u>	<u>N77534</u>	<u>001</u>	Oct 03, 2006
<u>AB</u>		<u>EQ 20MG BASE</u>	<u>N77534</u>	<u>002</u>	Oct 03, 2006
<u>AB</u>		<u>EQ 40MG BASE</u>	<u>N77534</u>	<u>003</u>	Oct 03, 2006
<u>AB</u>	IVAX PHARMS	<u>EQ 10MG BASE</u>	<u>N77048</u>	<u>001</u>	Nov 16, 2004
<u>AB</u>		<u>EQ 20MG BASE</u>	<u>N77048</u>	<u>002</u>	Nov 16, 2004
<u>AB</u>		<u>EQ 40MG BASE</u>	<u>N77048</u>	<u>003</u>	Nov 16, 2004
<u>AB</u>	KALI LABS	<u>EQ 10MG BASE</u>	<u>N77042</u>	<u>001</u>	Nov 05, 2004
<u>AB</u>		<u>EQ 20MG BASE</u>	<u>N77042</u>	<u>002</u>	Nov 05, 2004
<u>AB</u>		<u>EQ 40MG BASE</u>	<u>N77042</u>	<u>003</u>	Nov 05, 2004
<u>AB</u>	MUTUAL PHARM	<u>EQ 10MG BASE</u>	<u>N77052</u>	<u>001</u>	Jul 03, 2006
<u>AB</u>		<u>EQ 20MG BASE</u>	<u>N77052</u>	<u>002</u>	Jul 03, 2006
<u>AB</u>		<u>EQ 40MG BASE</u>	<u>N77052</u>	<u>003</u>	Jul 03, 2006
<u>AB</u>	MYLAN	<u>EQ 10MG BASE</u>	<u>N77039</u>	<u>001</u>	Feb 03, 2005
<u>AB</u>		<u>EQ 20MG BASE</u>	<u>N77039</u>	<u>002</u>	Feb 03, 2005
<u>AB</u>		<u>EQ 40MG BASE</u>	<u>N77039</u>	<u>003</u>	Feb 03, 2005
<u>AB</u>	PLIVA	<u>EQ 10MG BASE</u>	<u>N77232</u>	<u>001</u>	Oct 31, 2005
<u>AB</u>		<u>EQ 20MG BASE</u>	<u>N77232</u>	<u>002</u>	Oct 31, 2005
<u>AB</u>		<u>EQ 40MG BASE</u>	<u>N77232</u>	<u>003</u>	Oct 31, 2005
<u>AB</u>	ROXANE	<u>EQ 10MG BASE</u>	<u>N77041</u>	<u>001</u>	Nov 23, 2004
<u>AB</u>		<u>EQ 20MG BASE</u>	<u>N77041</u>	<u>002</u>	Nov 23, 2004
<u>AB</u>		<u>EQ 40MG BASE</u>	<u>N77041</u>	<u>003</u>	Nov 23, 2004
<u>AB</u>	SANDOZ	<u>EQ 10MG BASE</u>	<u>N77035</u>	<u>001</u>	Oct 28, 2004
<u>AB</u>		<u>EQ 10MG BASE</u>	<u>N77040</u>	<u>001</u>	Aug 17, 2005
<u>AB</u>		<u>EQ 20MG BASE</u>	<u>N77035</u>	<u>002</u>	Oct 28, 2004
<u>AB</u>		<u>EQ 20MG BASE</u>	<u>N77040</u>	<u>002</u>	Aug 17, 2005
<u>AB</u>		<u>EQ 40MG BASE</u>	<u>N77035</u>	<u>003</u>	Oct 28, 2004
<u>AB</u>		<u>EQ 40MG BASE</u>	<u>N77040</u>	<u>003</u>	Aug 17, 2005
<u>AB</u>	TARO	<u>EQ 10MG BASE</u>	<u>N77278</u>	<u>001</u>	Mar 22, 2006
<u>AB</u>		<u>EQ 20MG BASE</u>	<u>N77278</u>	<u>002</u>	Mar 22, 2006
<u>AB</u>		<u>EQ 40MG BASE</u>	<u>N77278</u>	<u>003</u>	Mar 22, 2006
<u>AB</u>	TEVA PHARMS	<u>EQ 10MG BASE</u>	<u>N77213</u>	<u>001</u>	Mar 31, 2006
<u>AB</u>		<u>EQ 20MG BASE</u>	<u>N77213</u>	<u>002</u>	Mar 31, 2006
<u>AB</u>		<u>EQ 40MG BASE</u>	<u>N77213</u>	<u>003</u>	Mar 31, 2006
<u>AB</u>	WATSON LABS	<u>EQ 10MG BASE</u>	<u>N77044</u>	<u>001</u>	Nov 05, 2004
<u>AB</u>		<u>EQ 20MG BASE</u>	<u>N77044</u>	<u>002</u>	Nov 05, 2004
<u>AB</u>		<u>EQ 40MG BASE</u>	<u>N77044</u>	<u>003</u>	Nov 05, 2004

CITRIC ACID; GLUCONOLACTONE; MAGNESIUM CARBONATE

SOLUTION; IRRIGATION

RENACIDIN

+ UNITED GUARDIAN 6.602GM/100ML;198MG/100ML;3.177GM/100ML N19481 001 Oct 02, 1990

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 92 (of 370)

CITRIC ACID; MAGNESIUM OXIDE; SODIUM CARBONATE

SOLUTION; IRRIGATION

UROLOGIC G IN PLASTIC CONTAINER

+	HOSPIRA	3.24GM/100ML; 380MG/100ML; 430MG/100ML	N18904	001	May 27, 1983
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CLADRIBINE

INJECTABLE; INJECTION

CLADRIBINE

<u>AP</u>	ABRAXIS PHARM	<u>1MG/ML</u>	<u>N76571</u>	<u>001</u>	Apr 22, 2004
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<u>AP</u>	BEDFORD	<u>1MG/ML</u>	<u>N75405</u>	<u>001</u>	Feb 28, 2000
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LEUSTATIN

<u>AP</u>	+ ORTHO BIOTECH	<u>1MG/ML</u>	<u>N20229</u>	<u>001</u>	Feb 26, 1993
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CLARITHROMYCIN

FOR SUSPENSION; ORAL

BIAXIN

	ABBOTT	125MG/5ML	N50698	001	Dec 23, 1993
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+		250MG/5ML	N50698	002	Dec 23, 1993
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TABLET; ORAL

BIAXIN

<u>AB</u>	+ ABBOTT	<u>250MG</u>	<u>N50662</u>	<u>001</u>	Oct 31, 1991
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<u>AB</u>	+	<u>500MG</u>	<u>N50662</u>	<u>002</u>	Oct 31, 1991
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CLARITHROMYCIN

<u>AB</u>	GENPHARM	<u>250MG</u>	<u>N65195</u>	<u>001</u>	Mar 11, 2005
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<u>AB</u>		<u>500MG</u>	<u>N65195</u>	<u>002</u>	Mar 11, 2005
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<u>AB</u>	IVAX PHARMS	<u>250MG</u>	<u>N65137</u>	<u>001</u>	May 31, 2005
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<u>AB</u>		<u>500MG</u>	<u>N65137</u>	<u>002</u>	May 31, 2005
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<u>AB</u>	RANBAXY	<u>250MG</u>	<u>N65174</u>	<u>001</u>	Sep 24, 2004
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<u>AB</u>		<u>500MG</u>	<u>N65174</u>	<u>002</u>	Sep 24, 2004
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<u>AB</u>	ROXANE	<u>250MG</u>	<u>N65178</u>	<u>002</u>	May 25, 2004
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<u>AB</u>		<u>500MG</u>	<u>N65178</u>	<u>001</u>	May 25, 2004
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<u>AB</u>	SANDOZ	<u>250MG</u>	<u>N65144</u>	<u>001</u>	Oct 18, 2005
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<u>AB</u>		<u>500MG</u>	<u>N65136</u>	<u>001</u>	Aug 25, 2005
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<u>AB</u>	TEVA	<u>250MG</u>	<u>N65155</u>	<u>001</u>	May 31, 2005
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<u>AB</u>		<u>500MG</u>	<u>N65155</u>	<u>002</u>	May 31, 2005
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<u>AB</u>	WOCKHARDT	<u>250MG</u>	<u>N65266</u>	<u>001</u>	May 31, 2006
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<u>AB</u>		<u>500MG</u>	<u>N65266</u>	<u>002</u>	May 31, 2006
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TABLET, EXTENDED RELEASE; ORAL

BIAXIN XL

<u>AB</u>	+ ABBOTT	<u>500MG</u>	<u>N50775</u>	<u>001</u>	Mar 03, 2000
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CLARITHROMYCIN

<u>AB</u>	ANDRX PHARMS	<u>500MG</u>	<u>N65145</u>	<u>001</u>	Jun 24, 2004
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+	RANBAXY	1GM	N65210	001	Jan 26, 2005
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<u>AB</u>	TEVA	<u>500MG</u>	<u>N65154</u>	<u>001</u>	May 18, 2005
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CLARITHROMYCIN EXTENDED RELEASE

<u>AB</u>	SANDOZ	<u>500MG</u>	<u>N65250</u>	<u>001</u>	Aug 25, 2005
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CLAVULANATE POTASSIUM; TICARCILLIN DISODIUM

INJECTABLE; INJECTION

TIMENTIN

+	GLAXOSMITHKLINE	EQ 1GM BASE/VIAL; EQ 30GM BASE/VIAL	N50590	003	Aug 18, 1987
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+		EQ 100MG BASE/VIAL; EQ 3GM BASE/VIAL	N50590	001	Apr 01, 1985
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		EQ 100MG BASE/VIAL; EQ 3GM BASE/VIAL	N62691	001	Dec 19, 1986
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+		EQ 200MG BASE/VIAL; EQ 3GM BASE/VIAL	N50590	002	Apr 01, 1985
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TIMENTIN IN PLASTIC CONTAINER

+	GLAXOSMITHKLINE	EQ 100MG BASE/100ML; EQ 3GM BASE/100ML	N50658	001	Dec 15, 1989
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PRESCRIPTION DRUG PRODUCT LIST

3 - 93 (of 370)

CLEMASTINE FUMARATE

SYRUP; ORAL

CLEMASTINE FUMARATE

<u>AA</u>	ACTAVIS MID ATLANTIC	<u>EQ 0.5MG BASE/5ML</u>	<u>N74075</u>	<u>001</u>	Oct 31, 1993
<u>AA</u>	MORTON GROVE	<u>EQ 0.5MG BASE/5ML</u>	<u>N74863</u>	<u>001</u>	Mar 13, 1998
<u>AA</u>	NOVEX	<u>EQ 0.5MG BASE/5ML</u>	<u>N75703</u>	<u>001</u>	Nov 27, 2000
<u>AA</u>	SILARX	<u>EQ 0.5MG BASE/5ML</u>	<u>N74884</u>	<u>001</u>	Dec 17, 1997
<u>AA</u> +	TEVA	<u>EQ 0.5MG BASE/5ML</u>	<u>N73399</u>	<u>001</u>	Jun 30, 1994
<u>AA</u>	TEVA PHARMS	<u>EQ 0.5MG BASE/5ML</u>	<u>N73095</u>	<u>001</u>	Apr 21, 1992

TABLET; ORAL

CLEMASTINE FUMARATE

<u>AB</u>	SANDOZ	<u>2.68MG</u>	<u>N73459</u>	<u>001</u>	Oct 31, 1993
<u>AB</u> +	TEVA	<u>2.68MG</u>	<u>N73283</u>	<u>001</u>	Jan 31, 1992

CLINDAMYCIN HYDROCHLORIDE

CAPSULE; ORAL

CLEOCIN HYDROCHLORIDE

<u>AB</u>	PHARMACIA AND UPJOHN	<u>EQ 75MG BASE</u>	<u>N50162</u>	<u>001</u>	
<u>AB</u>		<u>EQ 150MG BASE</u>	<u>N50162</u>	<u>002</u>	
<u>AB</u> +		<u>EQ 300MG BASE</u>	<u>N50162</u>	<u>003</u>	Apr 14, 1988

CLINDAMYCIN HYDROCHLORIDE

<u>AB</u>	COREPHARMA	<u>EQ 150MG BASE</u>	<u>N65194</u>	<u>001</u>	Mar 22, 2004
<u>AB</u>		<u>EQ 300MG BASE</u>	<u>N65194</u>	<u>002</u>	Mar 22, 2004
<u>AB</u>	LANNETT	<u>EQ 75MG BASE</u>	<u>N65242</u>	<u>001</u>	Aug 12, 2005
<u>AB</u>		<u>EQ 150MG BASE</u>	<u>N65242</u>	<u>002</u>	Aug 12, 2005
<u>AB</u>		<u>EQ 300MG BASE</u>	<u>N65243</u>	<u>001</u>	Aug 12, 2005
<u>AB</u>	RANBAXY	<u>EQ 150MG BASE</u>	<u>N65061</u>	<u>001</u>	Feb 02, 2001
<u>AB</u>		<u>EQ 300MG BASE</u>	<u>N65061</u>	<u>002</u>	Feb 02, 2001
<u>AB</u>	TEVA	<u>EQ 75MG BASE</u>	<u>N63027</u>	<u>001</u>	Sep 20, 1989
<u>AB</u>		<u>EQ 150MG BASE</u>	<u>N63029</u>	<u>001</u>	Sep 20, 1989
<u>AB</u>		<u>EQ 300MG BASE</u>	<u>N63029</u>	<u>002</u>	Aug 05, 2005
<u>AB</u>	WATSON LABS	<u>EQ 150MG BASE</u>	<u>N63083</u>	<u>001</u>	Jul 31, 1991
<u>AB</u>		<u>EQ 300MG BASE</u>	<u>N63083</u>	<u>002</u>	Mar 18, 2003
<u>AB</u>	ZYDUS PHARMS USA	<u>EQ 75MG BASE</u>	<u>N65217</u>	<u>001</u>	Jan 31, 2005
<u>AB</u>		<u>EQ 150MG BASE</u>	<u>N65217</u>	<u>002</u>	Jan 31, 2005
<u>AB</u>		<u>EQ 300MG BASE</u>	<u>N65217</u>	<u>003</u>	Jan 31, 2005

CLINDAMYCIN PALMITATE HYDROCHLORIDE

FOR SOLUTION; ORAL

CLEOCIN

+	PHARMACIA AND UPJOHN	EQ 75MG BASE/5ML	N62644	001	Apr 07, 1986
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CLINDAMYCIN PHOSPHATE

AEROSOL, FOAM; TOPICAL

EVOCLIN

+	CONNETICS	1%	N50801	001	Oct 22, 2004
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CREAM; VAGINAL

CLEOCIN

<u>AB</u> +	PHARMACIA AND UPJOHN	<u>EQ 2% BASE</u>	<u>N50680</u>	<u>002</u>	Mar 02, 1998
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CLINDAMYCIN PHOSPHATE

<u>AB</u>	ALTANA PHARMA	<u>EQ 2% BASE</u>	<u>N65139</u>	<u>001</u>	Dec 27, 2004
	CLINDESSE				
+	KV PHARM	EQ 2% BASE	N50793	001	Nov 30, 2004

GEL; TOPICAL

CLEOCIN T

<u>AB</u> +	PHARMACIA AND UPJOHN	<u>EQ 1% BASE</u>	<u>N50615</u>	<u>001</u>	Jan 07, 1987
	CLINDAGEL				
BT	GALDERMA LABS LP	EQ 1% BASE	N50782	001	Nov 27, 2000

PRESCRIPTION DRUG PRODUCT LIST

3 - 94 (of 370)

CLINDAMYCIN PHOSPHATE

GEL; TOPICAL

CLINDAMYCIN PHOSPHATE

<u>AB</u>	ALTANA	<u>EQ 1% BASE</u>	<u>N64160</u>	<u>001</u>	Jan 28, 2000
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INJECTABLE; INJECTION

CLEOCIN PHOSPHATE

<u>AP</u>	+	PHARMACIA AND UPJOHN	<u>EQ 150MG BASE/ML</u>	<u>N50441</u>	<u>001</u>	
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<u>AP</u>			<u>EQ 150MG BASE/ML</u>	<u>N62803</u>	<u>001</u>	Oct 16, 1987
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CLEOCIN PHOSPHATE IN DEXTROSE 5% IN PLASTIC CONTAINER

<u>AP</u>	+	PHARMACIA AND UPJOHN	<u>EQ 6MG BASE/ML</u>	<u>N50639</u>	<u>001</u>	Aug 30, 1989
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<u>AP</u>	+		<u>EQ 12MG BASE/ML</u>	<u>N50639</u>	<u>002</u>	Aug 30, 1989
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<u>AP</u>	+		<u>EQ 18MG BASE/ML</u>	<u>N50639</u>	<u>003</u>	Apr 10, 1991
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CLINDAMYCIN PHOSPHATE

<u>AP</u>		BAXTER HLTHCARE	<u>EQ 150MG BASE/ML</u>	<u>N62889</u>	<u>001</u>	Apr 25, 1988
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<u>AP</u>		BEDFORD	<u>EQ 150MG BASE/ML</u>	<u>N65206</u>	<u>001</u>	Sep 24, 2004
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<u>AP</u>		HOSPIRA	<u>EQ 150MG BASE/ML</u>	<u>N62800</u>	<u>001</u>	Jul 24, 1987
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<u>AP</u>			<u>EQ 150MG BASE/ML</u>	<u>N62801</u>	<u>001</u>	Jul 24, 1987
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<u>AP</u>			<u>EQ 150MG BASE/ML</u>	<u>N62943</u>	<u>001</u>	Sep 29, 1988
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CLINDAMYCIN PHOSPHATE IN DEXTROSE 5%

<u>AP</u>	+	ABRAXIS PHARM	<u>EQ 900MG BASE/100ML</u>	<u>N50635</u>	<u>001</u>	Dec 22, 1989
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LOTION; TOPICAL

CLEOCIN T

<u>AB</u>	+	PHARMACIA AND UPJOHN	<u>EQ 1% BASE</u>	<u>N50600</u>	<u>001</u>	May 31, 1989
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CLINDAMYCIN PHOSPHATE

<u>AB</u>		ALTANA	<u>EQ 1% BASE</u>	<u>N65067</u>	<u>001</u>	Jan 31, 2002
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SOLUTION; TOPICAL

CLEOCIN T

<u>AT</u>	+	PHARMACIA AND UPJOHN	<u>EQ 1% BASE</u>	<u>N50537</u>	<u>001</u>	
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CLINDA-DERM

<u>AT</u>		PADDOCK	<u>EQ 1% BASE</u>	<u>N63329</u>	<u>001</u>	Sep 30, 1992
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CLINDAMYCIN PHOSPHATE

<u>AT</u>		ACTAVIS MID ATLANTIC	<u>EQ 1% BASE</u>	<u>N62811</u>	<u>001</u>	Sep 01, 1988
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<u>AT</u>		ALTANA	<u>EQ 1% BASE</u>	<u>N65254</u>	<u>001</u>	Feb 14, 2006
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<u>AT</u>		FOUGERA	<u>EQ 1% BASE</u>	<u>N64159</u>	<u>001</u>	Jun 05, 1997
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<u>AT</u>		MORTON GROVE	<u>EQ 1% BASE</u>	<u>N63304</u>	<u>001</u>	Jul 15, 1997
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<u>AT</u>		PERRIGO NEW YORK	<u>EQ 1% BASE</u>	<u>N64050</u>	<u>001</u>	Nov 30, 1995
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<u>AT</u>		STIEFEL	<u>EQ 1% BASE</u>	<u>N64108</u>	<u>001</u>	Sep 27, 1996
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<u>AT</u>		TARO PHARM INDS	<u>EQ 1% BASE</u>	<u>N65184</u>	<u>001</u>	Mar 31, 2004
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SUPPOSITORY; VAGINAL

CLEOCIN

	+	PHARMACIA AND UPJOHN	100MG	N50767	001	Aug 13, 1999
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SWAB; TOPICAL

CLEOCIN

<u>AT</u>		PHARMACIA AND UPJOHN	<u>EQ 1% BASE</u>	<u>N50537</u>	<u>002</u>	Feb 22, 1994
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CLINDAMYCIN PHOSPHATE

<u>AT</u>		PERRIGO NEW YORK	<u>EQ 1% BASE</u>	<u>N65049</u>	<u>001</u>	May 25, 2000
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CLINDETS

<u>AT</u>		STIEFEL	<u>EQ 1% BASE</u>	<u>N64136</u>	<u>001</u>	Sep 30, 1996
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CLINDAMYCIN PHOSPHATE; TRETINOIN

GEL; TOPICAL

ZIANA

	+	MEDICIS	1.2%;0.025%	N50802	001	Nov 07, 2006
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CLOBETASOL PROPIONATE

AEROSOL, FOAM; TOPICAL

OLUX

	+	CONNETICS	0.05%	N21142	001	May 26, 2000
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27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 95 (of 370)

CLOBETASOL PROPIONATE

CREAM; TOPICAL

CLOBETASOL PROPIONATE

<u>AB1</u>	ACTAVIS MID ATLANTIC	<u>0.05%</u>	<u>N74139</u>	<u>001</u>	Aug 03, 1994
<u>AB1</u>	FOUGERA	<u>0.05%</u>	<u>N74392</u>	<u>001</u>	Sep 30, 1996
<u>AB1</u>	STIEFEL	<u>0.05%</u>	<u>N75338</u>	<u>001</u>	Feb 09, 2001
<u>AB1</u>	TARO	<u>0.05%</u>	<u>N74249</u>	<u>001</u>	Jul 08, 1996
<u>AB1</u>	TEVA PHARMS	<u>0.05%</u>	<u>N74087</u>	<u>001</u>	Feb 16, 1994

CLOBETASOL PROPIONATE (EMOLLIENT)

<u>AB2</u>	ALTANA	<u>0.05%</u>	<u>N75430</u>	<u>001</u>	May 26, 1999
<u>AB2</u>	STIEFEL	<u>0.05%</u>	<u>N75733</u>	<u>001</u>	Aug 22, 2001
<u>AB2</u>	TARO	<u>0.05%</u>	<u>N75633</u>	<u>001</u>	May 17, 2000

CORMAX

<u>AB1</u>	HEALTHPOINT	<u>0.05%</u>	<u>N74220</u>	<u>001</u>	May 16, 1997
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EMBELINE E

<u>AB2</u>	HEALTHPOINT	<u>0.05%</u>	<u>N75325</u>	<u>001</u>	Dec 24, 1998
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TEMOVATE

<u>AB1</u> +	ALTANA	<u>0.05%</u>	<u>N19322</u>	<u>001</u>	Dec 27, 1985
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TEMOVATE E

<u>AB2</u> +	ALTANA	<u>0.05%</u>	<u>N20340</u>	<u>001</u>	Jun 17, 1994
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GEL; TOPICAL

CLOBETASOL PROPIONATE

<u>AB</u>	ALTANA	<u>0.05%</u>	<u>N75368</u>	<u>001</u>	Feb 15, 2000
<u>AB</u>	STIEFEL	<u>0.05%</u>	<u>N75027</u>	<u>001</u>	Oct 31, 1997
<u>AB</u>	TARO	<u>0.05%</u>	<u>N75279</u>	<u>001</u>	May 28, 1999

EMBELINE

<u>AB</u>	HEALTHPOINT	<u>0.05%</u>	<u>N76141</u>	<u>001</u>	Apr 12, 2002
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TEMOVATE

<u>AB</u> +	ALTANA	<u>0.05%</u>	<u>N20337</u>	<u>001</u>	Apr 29, 1994
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LOTION; TOPICAL

CLOBEX

+	GALDERMA LABS LP	0.05%	N21535	001	Jul 24, 2003
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OINTMENT; TOPICAL

CLOBETASOL PROPIONATE

<u>AB</u>	ACTAVIS MID ATLANTIC	<u>0.05%</u>	<u>N74128</u>	<u>001</u>	Aug 03, 1994
<u>AB</u>	FOUGERA	<u>0.05%</u>	<u>N74407</u>	<u>001</u>	Feb 23, 1996
<u>AB</u>	STIEFEL	<u>0.05%</u>	<u>N75057</u>	<u>001</u>	Aug 12, 1998
<u>AB</u>	TARO	<u>0.05%</u>	<u>N74248</u>	<u>001</u>	Jul 12, 1996
<u>AB</u>	TEVA PHARMS	<u>0.05%</u>	<u>N74089</u>	<u>001</u>	Feb 16, 1994

EMBELINE

<u>AB</u>	HEALTHPOINT	<u>0.05%</u>	<u>N74221</u>	<u>001</u>	Mar 31, 1995
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TEMOVATE

<u>AB</u> +	ALTANA	<u>0.05%</u>	<u>N19323</u>	<u>001</u>	Dec 27, 1985
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SHAMPOO; TOPICAL

CLOBEX

+	GALDERMA LABS	0.05%	N21644	001	Feb 05, 2004
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SOLUTION; TOPICAL

CLOBETASOL PROPIONATE

<u>AT</u>	ACTAVIS MID ATLANTIC	<u>0.05%</u>	<u>N74331</u>	<u>001</u>	Dec 15, 1995
<u>AT</u>	MORTON GROVE	<u>0.05%</u>	<u>N75205</u>	<u>001</u>	Nov 13, 1998
<u>AT</u>	QLT USA	<u>0.05%</u>	<u>N76977</u>	<u>001</u>	Aug 05, 2005
<u>AT</u>	TARO	<u>0.05%</u>	<u>N75224</u>	<u>001</u>	Nov 16, 1998
<u>AT</u>		<u>0.05%</u>	<u>N75363</u>	<u>001</u>	Dec 29, 2000

EMBELINE

<u>AT</u>	DPT	<u>0.05%</u>	<u>N74222</u>	<u>001</u>	Dec 06, 1995
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TEMOVATE

<u>AT</u> +	ALTANA	<u>0.05%</u>	<u>N19966</u>	<u>001</u>	Feb 22, 1990
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27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 96 (of 370)

CLOBETASOL PROPIONATE

SPRAY; TOPICAL

CLOBEX

+ GALDERMA LABS LP 0.05%

N21835 001 Oct 27, 2005

CLOCORTOLONE PIVALATE

CREAM; TOPICAL

CLODERM

+ HEALTHPOINT 0.1%

N17765 001

CLOFARABINE

INJECTABLE; IV (INFUSION)

+ GENZYME 20MG/20ML (1MG/ML)

N21673 001 Dec 28, 2004

CLOFAZIMINE

CAPSULE; ORAL

LAMPRENE

+ NOVARTIS 50MG

N19500 002 Dec 15, 1986

CLOFIBRATE

CAPSULE; ORAL

CLOFIBRATEAB BANNER PHARMACAPS 500MGN73396 001 Mar 20, 1992AB TEVA 500MGN72600 001 Jul 25, 1991AB + USL PHARMA 500MGN70531 001 Jun 16, 1986CLOMIPHENE CITRATE

TABLET; ORAL

CLOMIDAB + SANOFI AVENTIS US 50MGN16131 002CLOMIPHENE CITRATEAB PAR PHARM 50MGN75528 001 Aug 30, 1999MILOPHENEAB MILEX 50MGN72196 001 Dec 20, 1988SEROPHENEAB SERONO 50MGN18361 001 Mar 22, 1982CLOMIPRAMINE HYDROCHLORIDE

CAPSULE; ORAL

ANAFRANILAB TYCO HLTHCARE 25MGN19906 001 Dec 29, 1989AB + 50MGN19906 002 Dec 29, 1989AB 75MGN19906 003 Dec 29, 1989CLOMIPRAMINE HYDROCHLORIDEAB MYLAN 25MGN74947 001 Apr 30, 1998AB 50MGN74947 002 Apr 30, 1998AB 75MGN74947 003 Apr 30, 1998AB SANDOZ 25MGN74364 001 Mar 29, 1996AB 25MGN74953 001 Jun 25, 1997AB 50MGN74364 002 Mar 29, 1996AB 50MGN74953 002 Jun 25, 1997AB 75MGN74364 003 Mar 29, 1996AB 75MGN74953 003 Jun 25, 1997AB TARO 25MGN74694 001 Dec 31, 1996AB 50MGN74694 002 Dec 31, 1996AB 75MGN74694 003 Dec 31, 1996AB TEVA 25MGN74849 001 Apr 04, 1997AB 25MGN74958 001 Aug 26, 1997

PRESCRIPTION DRUG PRODUCT LIST

3 - 97 (of 370)

CLOMIPRAMINE HYDROCHLORIDE

CAPSULE; ORAL

CLOMIPRAMINE HYDROCHLORIDE

<u>AB</u>	TEVA	<u>50MG</u>	<u>N74849</u>	<u>002</u>	Apr 04, 1997
<u>AB</u>		<u>50MG</u>	<u>N74958</u>	<u>002</u>	Aug 26, 1997
<u>AB</u>		<u>75MG</u>	<u>N74849</u>	<u>003</u>	Apr 04, 1997
<u>AB</u>		<u>75MG</u>	<u>N74958</u>	<u>003</u>	Aug 26, 1997
<u>AB</u>	WATSON LABS	<u>25MG</u>	<u>N74600</u>	<u>001</u>	Nov 27, 1996
<u>AB</u>		<u>25MG</u>	<u>N74751</u>	<u>001</u>	Sep 30, 1998
<u>AB</u>		<u>50MG</u>	<u>N74600</u>	<u>002</u>	Nov 27, 1996
<u>AB</u>		<u>50MG</u>	<u>N74751</u>	<u>002</u>	Sep 30, 1998
<u>AB</u>		<u>75MG</u>	<u>N74600</u>	<u>003</u>	Nov 27, 1996
<u>AB</u>		<u>75MG</u>	<u>N74751</u>	<u>003</u>	Sep 30, 1998

CLONAZEPAM

TABLET; ORAL

CLONAZEPAM

<u>AB</u>	ACTAVIS ELIZABETH	<u>0.5MG</u>	<u>N74869</u>	<u>001</u>	Oct 31, 1996
<u>AB</u>		<u>1MG</u>	<u>N74869</u>	<u>002</u>	Oct 31, 1996
<u>AB</u>		<u>2MG</u>	<u>N74869</u>	<u>003</u>	Oct 31, 1996
<u>AB</u>	ALPHAPHARM	<u>0.5MG</u>	<u>N74940</u>	<u>001</u>	Oct 30, 1997
<u>AB</u>		<u>1MG</u>	<u>N74940</u>	<u>002</u>	Oct 30, 1997
<u>AB</u>		<u>2MG</u>	<u>N74940</u>	<u>003</u>	Oct 30, 1997
<u>AB</u>	CARACO	<u>0.5MG</u>	<u>N75423</u>	<u>001</u>	Apr 27, 2001
<u>AB</u>		<u>1MG</u>	<u>N75423</u>	<u>002</u>	Apr 27, 2001
<u>AB</u>		<u>2MG</u>	<u>N75423</u>	<u>003</u>	Apr 27, 2001
<u>AB</u>	KALI LABS	<u>0.5MG</u>	<u>N77147</u>	<u>001</u>	May 02, 2005
<u>AB</u>		<u>1MG</u>	<u>N77147</u>	<u>002</u>	May 02, 2005
<u>AB</u>		<u>2MG</u>	<u>N77147</u>	<u>003</u>	May 02, 2005
<u>AB</u>	MYLAN	<u>0.5MG</u>	<u>N75150</u>	<u>001</u>	Oct 05, 1998
<u>AB</u>		<u>1MG</u>	<u>N75150</u>	<u>002</u>	Oct 05, 1998
<u>AB</u>		<u>2MG</u>	<u>N75150</u>	<u>003</u>	Oct 05, 1998
<u>AB</u>	SANDOZ	<u>0.5MG</u>	<u>N74925</u>	<u>001</u>	Sep 30, 1997
<u>AB</u>		<u>0.5MG</u>	<u>N74979</u>	<u>001</u>	Aug 29, 1997
<u>AB</u>		<u>1MG</u>	<u>N74925</u>	<u>002</u>	Sep 30, 1997
<u>AB</u>		<u>1MG</u>	<u>N74979</u>	<u>002</u>	Aug 29, 1997
<u>AB</u>		<u>2MG</u>	<u>N74925</u>	<u>003</u>	Sep 30, 1997
<u>AB</u>		<u>2MG</u>	<u>N74979</u>	<u>003</u>	Aug 29, 1997
<u>AB</u>	TEVA	<u>0.5MG</u>	<u>N74569</u>	<u>001</u>	Sep 10, 1996
<u>AB</u>		<u>0.5MG</u>	<u>N74920</u>	<u>001</u>	Aug 04, 1998
<u>AB</u>		<u>1MG</u>	<u>N74569</u>	<u>002</u>	Sep 10, 1996
<u>AB</u>		<u>1MG</u>	<u>N74920</u>	<u>002</u>	Aug 04, 1998
<u>AB</u>		<u>2MG</u>	<u>N74569</u>	<u>003</u>	Sep 10, 1996
<u>AB</u>		<u>2MG</u>	<u>N74920</u>	<u>003</u>	Aug 04, 1998
<u>AB</u>	TORPHARM	<u>0.5MG</u>	<u>N75468</u>	<u>001</u>	Oct 06, 2000
<u>AB</u>		<u>1MG</u>	<u>N75468</u>	<u>002</u>	Oct 06, 2000
<u>AB</u>		<u>2MG</u>	<u>N75468</u>	<u>003</u>	Oct 06, 2000
<u>AB</u>	VINTAGE PHARMS	<u>0.5MG</u>	<u>N77856</u>	<u>001</u>	Jun 28, 2006
<u>AB</u>		<u>1MG</u>	<u>N77856</u>	<u>002</u>	Jun 28, 2006
<u>AB</u>		<u>2MG</u>	<u>N77856</u>	<u>003</u>	Jun 28, 2006
<u>AB</u>	WATSON LABS	<u>0.5MG</u>	<u>N74964</u>	<u>001</u>	Dec 30, 1997
<u>AB</u>		<u>1MG</u>	<u>N74964</u>	<u>002</u>	Dec 30, 1997
<u>AB</u>		<u>2MG</u>	<u>N74964</u>	<u>003</u>	Dec 30, 1997
<u>KLONOPIN</u>					
<u>AB</u>	ROCHE	<u>0.5MG</u>	<u>N17533</u>	<u>001</u>	
<u>AB</u>	+	<u>1MG</u>	<u>N17533</u>	<u>002</u>	
<u>AB</u>		<u>2MG</u>	<u>N17533</u>	<u>003</u>	

TABLET, ORALLY DISINTEGRATING; ORAL

CLONAZEPAM

<u>AB</u>	BARR	<u>0.125MG</u>	<u>N77194</u>	<u>001</u>	Aug 10, 2005
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PRESCRIPTION DRUG PRODUCT LIST

3 - 98 (of 370)

CLONAZEPAM

TABLET, ORALLY DISINTEGRATING; ORAL

CLONAZEPAM

<u>AB</u>	BARR	<u>0.25MG</u>	<u>N77194</u>	<u>002</u>	Aug 10, 2005
<u>AB</u>		<u>0.5MG</u>	<u>N77194</u>	<u>003</u>	Aug 10, 2005
<u>AB</u>		<u>1MG</u>	<u>N77194</u>	<u>004</u>	Aug 10, 2005
<u>AB</u>		<u>2MG</u>	<u>N77194</u>	<u>005</u>	Aug 10, 2005
<u>AB</u>	KALI LABS	<u>0.125MG</u>	<u>N77171</u>	<u>001</u>	Aug 03, 2005
<u>AB</u>		<u>0.25MG</u>	<u>N77171</u>	<u>002</u>	Aug 03, 2005
<u>AB</u>		<u>0.5MG</u>	<u>N77171</u>	<u>003</u>	Aug 03, 2005
<u>AB</u>		<u>1MG</u>	<u>N77171</u>	<u>004</u>	Aug 03, 2005
<u>AB</u>		<u>2MG</u>	<u>N77171</u>	<u>005</u>	Aug 03, 2005
<u>KLONOPIN RAPIDLY DISINTEGRATING</u>					
<u>AB</u>	ROCHE	<u>0.125MG</u>	<u>N20813</u>	<u>001</u>	Dec 23, 1997
<u>AB</u>		<u>0.25MG</u>	<u>N20813</u>	<u>002</u>	Dec 23, 1997
<u>AB</u>		<u>0.5MG</u>	<u>N20813</u>	<u>003</u>	Dec 23, 1997
<u>AB</u>	+	<u>1MG</u>	<u>N20813</u>	<u>004</u>	Dec 23, 1997
<u>AB</u>		<u>2MG</u>	<u>N20813</u>	<u>005</u>	Dec 23, 1997

CLONIDINE

FILM, EXTENDED RELEASE; TRANSDERMAL

CATAPRES-TTS-1

BOEHRINGER INGELHEIM 0.1MG/24HR

N18891 001 Oct 10, 1984

CATAPRES-TTS-2

BOEHRINGER INGELHEIM 0.2MG/24HR

N18891 002 Oct 10, 1984

CATAPRES-TTS-3

+ BOEHRINGER INGELHEIM 0.3MG/24HR

N18891 003 Oct 10, 1984

CLONIDINE HYDROCHLORIDE

INJECTABLE; INJECTION

DURACLON

XANODYNE PHARM 0.1MG/ML

N20615 001 Oct 02, 1996

+ 0.5MG/ML

N20615 002 Apr 27, 1999

TABLET; ORAL

CATAPRES

<u>AB</u>	BOEHRINGER INGELHEIM	<u>0.1MG</u>	<u>N17407</u>	<u>001</u>	
<u>AB</u>		<u>0.2MG</u>	<u>N17407</u>	<u>002</u>	
<u>AB</u>	+	<u>0.3MG</u>	<u>N17407</u>	<u>003</u>	

CLONIDINE HYDROCHLORIDE

<u>AB</u>	ACTAVIS ELIZABETH	<u>0.1MG</u>	<u>N70974</u>	<u>001</u>	Dec 16, 1986
<u>AB</u>		<u>0.2MG</u>	<u>N70975</u>	<u>001</u>	Dec 16, 1986
<u>AB</u>		<u>0.3MG</u>	<u>N70976</u>	<u>001</u>	Dec 16, 1986
<u>AB</u>	CLONMEL HLTHCARE	<u>0.1MG</u>	<u>N71783</u>	<u>001</u>	Apr 05, 1988
<u>AB</u>		<u>0.2MG</u>	<u>N71784</u>	<u>001</u>	Apr 05, 1988
<u>AB</u>		<u>0.3MG</u>	<u>N71785</u>	<u>001</u>	Apr 05, 1988
<u>AB</u>	MUTUAL PHARM	<u>0.1MG</u>	<u>N70925</u>	<u>001</u>	Sep 04, 1987
<u>AB</u>		<u>0.2MG</u>	<u>N70924</u>	<u>001</u>	Sep 04, 1987
<u>AB</u>		<u>0.3MG</u>	<u>N70923</u>	<u>001</u>	Sep 04, 1987
<u>AB</u>	MYLAN	<u>0.1MG</u>	<u>N70315</u>	<u>001</u>	Jun 09, 1987
<u>AB</u>		<u>0.2MG</u>	<u>N70316</u>	<u>001</u>	Jun 09, 1987
<u>AB</u>		<u>0.3MG</u>	<u>N70317</u>	<u>001</u>	Jun 09, 1987
<u>AB</u>	SANDOZ	<u>0.1MG</u>	<u>N70887</u>	<u>001</u>	Aug 31, 1988
<u>AB</u>		<u>0.2MG</u>	<u>N70886</u>	<u>001</u>	Aug 31, 1988
<u>AB</u>		<u>0.3MG</u>	<u>N71294</u>	<u>001</u>	Aug 31, 1988
<u>AB</u>	WATSON LABS	<u>0.1MG</u>	<u>N70965</u>	<u>001</u>	Jul 08, 1986
<u>AB</u>		<u>0.2MG</u>	<u>N70964</u>	<u>001</u>	Jul 08, 1986
<u>AB</u>		<u>0.3MG</u>	<u>N70963</u>	<u>001</u>	Jul 08, 1986

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 99 (of 370)

CLOPIDOGREL BISULFATE

TABLET; ORAL

CLOPIDOGREL BISULFATE

<u>AB</u>	APOTEX INC	<u>EQ 75MG BASE</u>	<u>N76274</u>	<u>001</u>	Jan 20, 2006
	<u>PLAVIX</u>				
<u>AB</u>	+ SANOFI AVENTIS US	<u>EQ 75MG BASE</u>	<u>N20839</u>	<u>001</u>	Nov 17, 1997

CLORAZEPATE DIPOTASSIUM

CAPSULE; ORAL

CLORAZEPATE DIPOTASSIUM

<u>AB</u>	MYLAN	<u>3.75MG</u>	<u>N71509</u>	<u>001</u>	Oct 19, 1987
<u>AB</u>		<u>7.5MG</u>	<u>N71510</u>	<u>001</u>	Oct 19, 1987
<u>AB</u>	+	<u>15MG</u>	<u>N71511</u>	<u>001</u>	Oct 19, 1987
<u>AB</u>	SANDOZ	<u>3.75MG</u>	<u>N72219</u>	<u>001</u>	Aug 26, 1988
<u>AB</u>		<u>15MG</u>	<u>N72112</u>	<u>001</u>	Aug 26, 1988

TABLET; ORAL

CLORAZEPATE DIPOTASSIUM

<u>AB</u>	MYLAN	<u>3.75MG</u>	<u>N71856</u>	<u>001</u>	Jul 17, 1987
<u>AB</u>		<u>7.5MG</u>	<u>N71857</u>	<u>001</u>	Jul 17, 1987
<u>AB</u>		<u>15MG</u>	<u>N71858</u>	<u>001</u>	Jul 17, 1987
<u>AB</u>	RANBAXY	<u>3.75MG</u>	<u>N76911</u>	<u>001</u>	Sep 29, 2004
<u>AB</u>		<u>7.5MG</u>	<u>N76911</u>	<u>002</u>	Sep 29, 2004
<u>AB</u>		<u>15MG</u>	<u>N76911</u>	<u>003</u>	Sep 29, 2004
<u>AB</u>	SANDOZ	<u>7.5MG</u>	<u>N72513</u>	<u>001</u>	May 11, 1990
<u>AB</u>		<u>15MG</u>	<u>N72514</u>	<u>001</u>	May 11, 1990
<u>AB</u>	TARO	<u>3.75MG</u>	<u>N75731</u>	<u>003</u>	Apr 27, 2000
<u>AB</u>		<u>7.5MG</u>	<u>N75731</u>	<u>002</u>	Apr 27, 2000
<u>AB</u>		<u>15MG</u>	<u>N75731</u>	<u>001</u>	Apr 27, 2000
<u>AB</u>	WATSON LABS	<u>3.75MG</u>	<u>N71852</u>	<u>001</u>	Feb 09, 1988
<u>AB</u>		<u>7.5MG</u>	<u>N71853</u>	<u>001</u>	Feb 09, 1988
<u>AB</u>		<u>15MG</u>	<u>N71854</u>	<u>001</u>	Feb 09, 1988
	<u>GEN-XENE</u>				
<u>AB</u>	ALRA	<u>3.75MG</u>	<u>N71787</u>	<u>001</u>	Apr 26, 1988
<u>AB</u>		<u>7.5MG</u>	<u>N71788</u>	<u>001</u>	Apr 26, 1988
<u>AB</u>		<u>15MG</u>	<u>N71789</u>	<u>001</u>	Apr 26, 1988
	<u>TRANXENE</u>				
<u>AB</u>	OVATION PHARMS	<u>3.75MG</u>	<u>N17105</u>	<u>006</u>	
<u>AB</u>		<u>7.5MG</u>	<u>N17105</u>	<u>007</u>	
<u>AB</u>	+	<u>15MG</u>	<u>N17105</u>	<u>008</u>	
	TRANXENE SD				
	OVATION PHARMS	11.25MG	N17105	005	
	+	22.5MG	N17105	004	

CLOTRIMAZOLE

CREAM; TOPICAL

CLOTRIMAZOLE

	+ TARO	1%	N72640	001	Aug 31, 1993
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SOLUTION; TOPICAL

CLOTRIMAZOLE

<u>AT</u>	TARO	<u>1%</u>	<u>N74580</u>	<u>001</u>	Jul 29, 1996
<u>AT</u>	TEVA	<u>1%</u>	<u>N73306</u>	<u>001</u>	Feb 28, 1995
	<u>MYCELEX</u>				
<u>AT</u>	+ BAYER PHARMS	<u>1%</u>	<u>N18181</u>	<u>001</u>	

TROCHE/LOZENGE; ORAL

CLOTRIMAZOLE

<u>AB</u>	PADDOCK	<u>10MG</u>	<u>N76763</u>	<u>001</u>	Oct 28, 2005
<u>AB</u>	ROXANE	<u>10MG</u>	<u>N76387</u>	<u>001</u>	Jul 29, 2004
	<u>MYCELEX</u>				
<u>AB</u>	+ BAYER PHARMS	<u>10MG</u>	<u>N18713</u>	<u>001</u>	Jun 17, 1983

PRESCRIPTION DRUG PRODUCT LIST

3 - 100 (of 370)

CLOXACILLIN SODIUM

CAPSULE; ORAL

CLOXACILLIN SODIUM

<u>AB</u>	TEVA	<u>EQ 250MG BASE</u>	<u>N62240</u>	<u>001</u>	
<u>AB</u>		<u>EQ 500MG BASE</u>	<u>N62240</u>	<u>002</u>	
	<u>CLOXAPEN</u>				
<u>AB</u>	GLAXOSMITHKLINE	<u>EQ 250MG BASE</u>	<u>N61806</u>	<u>001</u>	
<u>AB</u>	+	<u>EQ 500MG BASE</u>	<u>N61806</u>	<u>002</u>	

FOR SOLUTION; ORAL
CLOXACILLIN SODIUM

	+	TEVA	EQ 125MG BASE/5ML	N62268	001
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CLOZAPINE

TABLET; ORAL

CLOZAPINE

<u>AB</u>	CARACO	<u>25MG</u>	<u>N75713</u>	<u>001</u>	Nov 15, 2002
<u>AB</u>		<u>50MG</u>	<u>N75713</u>	<u>003</u>	Aug 19, 2005
<u>AB</u>		<u>100MG</u>	<u>N75713</u>	<u>002</u>	Nov 15, 2002
<u>AB</u>	GOLDLINE	<u>50MG</u>	<u>N76809</u>	<u>003</u>	Dec 16, 2005
<u>AB</u>		<u>100MG</u>	<u>N76809</u>	<u>002</u>	Dec 16, 2005
		200MG	N76809	001	Dec 16, 2005
<u>AB</u>	IVAX PHARMS	<u>25MG</u>	<u>N74949</u>	<u>001</u>	Nov 26, 1997
<u>AB</u>		<u>50MG</u>	<u>N74949</u>	<u>004</u>	Apr 25, 2005
<u>AB</u>		<u>100MG</u>	<u>N74949</u>	<u>002</u>	Nov 26, 1997
		12.5MG	N74949	003	Jul 31, 2003
<u>AB</u>	MYLAN	<u>25MG</u>	<u>N75417</u>	<u>001</u>	May 27, 1999
<u>AB</u>		<u>100MG</u>	<u>N75417</u>	<u>002</u>	May 27, 1999
<u>AB</u>	TEVA	<u>25MG</u>	<u>N75162</u>	<u>001</u>	Apr 26, 2005
<u>AB</u>		<u>100MG</u>	<u>N75162</u>	<u>002</u>	Apr 26, 2005
	<u>CLOZARIL</u>				
<u>AB</u>	NOVARTIS	<u>25MG</u>	<u>N19758</u>	<u>001</u>	Sep 26, 1989
<u>AB</u>	+	<u>100MG</u>	<u>N19758</u>	<u>002</u>	Sep 26, 1989

TABLET, ORALLY DISINTEGRATING; ORAL

FAZACLO ODT

	AVANIR PHARMS	25MG	N21590	001	Feb 10, 2004
		50MG	N21590	003	Jun 03, 2005
	+	100MG	N21590	002	Feb 10, 2004

CODEINE PHOSPHATE; PHENYLEPHRINE HYDROCHLORIDE; PROMETHAZINE HYDROCHLORIDE

SYRUP; ORAL

PROMETH HYDROCHLORIDE, PHENYLEPHRINE HYDROCHLORIDE W/CODEINE PHOSPHATE

<u>AA</u>	VINTAGE	<u>10MG/5ML;5MG/5ML;6.25MG/5ML</u>	<u>N40660</u>	<u>001</u>	Dec 07, 2006
	<u>PROMETH VC W/ CODEINE</u>				
AA +	ACTAVIS MID ATLANTIC	10MG/5ML;5MG/5ML;6.25MG/5ML	N88764	001	Oct 31, 1984

CODEINE PHOSPHATE; PROMETHAZINE HYDROCHLORIDE

SYRUP; ORAL

PROMETH W/ CODEINE

<u>AA</u>	+	ACTAVIS MID ATLANTIC	<u>10MG/5ML; 6.25MG/5ML</u>	<u>N88763</u>	<u>001</u>	Oct 31, 1984
	<u>PROMETHAZINE HYDROCHLORIDE AND CODEINE PHOSPHATE</u>					
<u>AA</u>	HI TECH PHARMA	<u>10MG/5ML; 6.25MG/5ML</u>	<u>N40151</u>	<u>001</u>	Aug 26, 1997	
<u>AA</u>	PHARM ASSOC	<u>10MG/5ML; 6.25MG/5ML</u>	<u>N89647</u>	<u>001</u>	Dec 22, 1988	
	<u>PROMETHAZINE W/ CODEINE</u>					
<u>AA</u>	MORTON GROVE	<u>10MG/5ML; 6.25MG/5ML</u>	<u>N88875</u>	<u>001</u>	Dec 17, 1984	
	<u>PROMETHAZINE WITH CODEINE SYRUP</u>					
<u>AA</u>	VINTAGE	<u>10MG/5ML; 6.25MG/5ML</u>	<u>N40650</u>	<u>001</u>	Jan 31, 2006	

PRESCRIPTION DRUG PRODUCT LIST

3 - 101 (of 370)

CODEINE PHOSPHATE; PSEUDOEPHEDRINE HYDROCHLORIDE; TRIPROLIDINE HYDROCHLORIDE

SYRUP; ORAL

TRIACIN-C

<u>AA</u>	ACTAVIS MID ATLANTIC	<u>10MG/5ML;30MG/5ML;1.25MG/5ML</u>	<u>N88704</u>	<u>001</u>	Mar 22, 1985
<u>AA</u>	<u>TRIPROLIDINE HCL, PSEUDOEPHEDRINE HCL AND CODEINE PHOSPHATE</u>				
<u>AA</u>	+ MORTON GROVE	<u>10MG/5ML;30MG/5ML;1.25MG/5ML</u>	<u>N88833</u>	<u>001</u>	Nov 16, 1984

COLCHICINE; PROBENECID

TABLET; ORAL

COL-PROBENECID

BP	+ WATSON LABS	0.5MG;500MG	N84279	001	
	PROBENECID AND COLCHICINE				
BP	IVAX PHARMS	0.5MG;500MG	N83734	001	

COLESEVELAM HYDROCHLORIDE

TABLET; ORAL

WELCHOL

	+ DAIICHI SANKYO	625MG	N21176	001	May 26, 2000
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COLESTIPOL HYDROCHLORIDE

GRANULE; ORAL

COLESTID

<u>AB</u>	PHARMACIA AND UPJOHN	<u>5GM/SC00PFUL</u>	<u>N17563</u>	<u>003</u>	Sep 22, 1995
<u>AB</u>	<u>5GM/PACKET</u>		<u>N17563</u>	<u>004</u>	Sep 22, 1995
	<u>COLESTIPOL HYDROCHLORIDE</u>				
<u>AB</u>	IMPAX LABS	<u>5GM/SC00PFUL</u>	<u>N77277</u>	<u>001</u>	May 02, 2006
<u>AB</u>		<u>5GM/PACKET</u>	<u>N77277</u>	<u>002</u>	May 02, 2006
	FLAVORED COLESTID				
	PHARMACIA AND UPJOHN	5GM/PACKET	N17563	001	
		5GM/SC00PFUL	N17563	002	

TABLET; ORAL

COLESTID

<u>AB</u>	+ PHARMACIA AND UPJOHN	<u>1GM</u>	<u>N20222</u>	<u>001</u>	Jul 19, 1994
	<u>COLESTIPOL HYDROCHLORIDE</u>				
<u>AB</u>	IMPAX LABS	<u>1GM</u>	<u>N77510</u>	<u>001</u>	Oct 24, 2006

COLISTIMETHATE SODIUM

INJECTABLE; INJECTION

COLISTIMETHATE

<u>AP</u>	PADDOCK	<u>EQ 150MG BASE/VIAL</u>	<u>N65177</u>	<u>001</u>	Mar 19, 2004
<u>AP</u>	X GEN PHARMS	<u>EQ 150MG BASE/VIAL</u>	<u>N64216</u>	<u>001</u>	Feb 26, 1999
	<u>COLY-MYCIN M</u>				
<u>AP</u>	+ PARKEDALE	<u>EQ 150MG BASE/VIAL</u>	<u>N50108</u>	<u>002</u>	

COLISTIN SULFATE; HYDROCORTISONE ACETATE; NEOMYCIN SULFATE; THONZONIUM BROMIDE

SUSPENSION/DROPS; OTIC

COLY-MYCIN S

	+ PARKEDALE	EQ 3MG BASE/ML;10MG/ML;EQ 3.3MG BASE/ML;0.5MG/ML	N50356	001	
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CONIVAPTAN HYDROCHLORIDE

INJECTABLE; IV (INFUSION)

	+ ASTELLAS	20MG/4ML (5MG/ML)	N21697	001	Dec 29, 2005
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COPPER

INTRAUTERINE DEVICE; INTRAUTERINE

PARAGARD T 380A

	+ DURAMED PHARMS BARR	309MG/COPPER	N18680	001	Nov 15, 1984
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27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 102 (of 370)

CORTICORELIN OVINE TRIFLUTATE

INJECTABLE; INJECTION

ACTHREL

+ FERRING

EQ 0.1MG BASE/VIAL

N20162 001

May 23, 1996

CORTICOTROPIN

INJECTABLE; INJECTION

H.P. ACTHAR GEL

+ QUESTCOR PHARMS

80 UNITS/ML

N08372 008

CORTISONE ACETATE

TABLET; ORAL

CORTISONE ACETATE

BP + PHARMACIA AND UPJOHN

25MG

N08126 001

5MG

N08126 003

10MG

N08126 004

BP WEST WARD

25MG

N80776 002

COSYNTROPIN

INJECTABLE; INJECTION

CORTROSYN

+ AMPHASTAR

0.25MG/VIAL

N16750 001

CROMOLYN SODIUM

AEROSOL, METERED; INHALATION

INTAL

+ KING PHARMS

0.8MG/INH

N18887 001

Dec 05, 1985

CONCENTRATE; ORAL

GASTROCROM

+ AZUR PHARMA

100MG/5ML

N20479 001

Feb 29, 1996

SOLUTION; INHALATION

CROMOLYN SODIUMAN ACTAVIS MID ATLANTIC10MG/MLN75067 001

Jul 19, 1999

AN BAUSCH AND LOMB10MG/MLN75585 001

Dec 21, 2000

AN DEY10MG/MLN74209 001

Apr 26, 1994

AN IVAX PHARMS10MG/MLN75271 001

Jan 18, 2000

AN MORTON GROVE10MG/MLN75346 001

Oct 25, 1999

AN NOVEX10MG/MLN75333 001

Apr 30, 2002

AN RESPIRARE10MG/MLN76469 001

Jun 17, 2005

AN ROXANE10MG/MLN75175 001

Sep 30, 1999

AN WARRICK PHARMS10MG/MLN75437 001

Apr 21, 2000

INTALAN + KING PHARMS10MG/MLN18596 001

May 28, 1982

SOLUTION/DROPS; OPHTHALMIC

CROLOMAT BAUSCH AND LOMB4%N74443 001

Jan 30, 1995

CROMOLYN SODIUMAT AKORN4%N74706 001

Apr 29, 1998

AT ALCON4%N75282 001

Jun 16, 1999

AT NOVEX4%N75615 001

Jan 26, 2001

OPTICROMAT + ALLERGAN4%N18155 001

Oct 03, 1984

CROTAMITON

CREAM; TOPICAL

EURAX

+ WESTWOOD SQUIBB

10%

N06927 001

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 103 (of 370)

CROTAMITON

LOTION; TOPICAL

CROTAN

<u>AT</u>	SUMMERS	<u>10%</u>	<u>N87204</u>	<u>001</u>	
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EURAX

<u>AT</u>	+ WESTWOOD SQUIBB	<u>10%</u>	<u>N09112</u>	<u>003</u>	
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CUPRIC CHLORIDE

INJECTABLE; INJECTION

CUPRIC CHLORIDE IN PLASTIC CONTAINER

+	HOSPIRA	EQ 0.4MG COPPER/ML	N18960	001	Jun 26, 1986
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CYANOCOBALAMIN

INJECTABLE; INJECTION

CYANOCOBALAMIN

<u>AP</u>	+ BIONICHE PHARMA	<u>1MG/ML</u>	<u>N40451</u>	<u>001</u>	Sep 23, 2003
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<u>AP</u>	+ LUITPOLD	<u>1MG/ML</u>	<u>N80737</u>	<u>001</u>	
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VIBISONE

<u>AP</u>	+ ABRAXIS PHARM	<u>1MG/ML</u>	<u>N80557</u>	<u>003</u>	
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SPRAY, METERED; NASAL

NASCOBAL

+	QOL MEDCL	0.5MG/SPRAY	N21642	001	Jan 31, 2005
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CYANOCOBALAMIN, CO-57

CAPSULE; ORAL

RUBRATOPE-57

+	BRACCO	0.5-1uCi	N16089	002	
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CYCLOBENZAPRINE HYDROCHLORIDE

TABLET; ORAL

CYCLOBENZAPRINE HYDROCHLORIDE

<u>AB</u>	ACTAVIS TOTOWA	<u>5MG</u>	<u>N77291</u>	<u>001</u>	Feb 03, 2006
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<u>AB</u>		<u>10MG</u>	<u>N77209</u>	<u>001</u>	Oct 04, 2005
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<u>AB</u>	JUBILANT PHARMS	<u>5MG</u>	<u>N77563</u>	<u>001</u>	Apr 19, 2006
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<u>AB</u>		<u>10MG</u>	<u>N77563</u>	<u>002</u>	Apr 19, 2006
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<u>AB</u>	MUTUAL PHARM	<u>5MG</u>	<u>N73541</u>	<u>002</u>	Apr 06, 2006
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<u>AB</u>		<u>10MG</u>	<u>N73541</u>	<u>001</u>	May 23, 1995
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<u>AB</u>	MYLAN	<u>5MG</u>	<u>N73144</u>	<u>002</u>	Feb 03, 2006
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<u>AB</u>		<u>10MG</u>	<u>N73144</u>	<u>001</u>	May 30, 1991
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<u>AB</u>	PLIVA	<u>10MG</u>	<u>N74421</u>	<u>001</u>	Sep 29, 1995
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<u>AB</u>	SANDOZ	<u>5MG</u>	<u>N72854</u>	<u>002</u>	Feb 03, 2006
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<u>AB</u>		<u>10MG</u>	<u>N72854</u>	<u>001</u>	Nov 19, 1991
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<u>AB</u>		<u>10MG</u>	<u>N73683</u>	<u>001</u>	Feb 26, 1993
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<u>AB</u>	WATSON LABS	<u>5MG</u>	<u>N71611</u>	<u>002</u>	Feb 03, 2006
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<u>AB</u>		<u>10MG</u>	<u>N71611</u>	<u>001</u>	May 03, 1989
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<u>AB</u>		<u>10MG</u>	<u>N73143</u>	<u>001</u>	Nov 27, 1991
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<u>AB</u>		<u>10MG</u>	<u>N74436</u>	<u>001</u>	Nov 30, 1994
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		7.5MG	N71611	003	Feb 03, 2006
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FLEXERIL

<u>AB</u>	MCNEIL PED	<u>5MG</u>	<u>N17821</u>	<u>001</u>	
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<u>AB</u>	+	<u>10MG</u>	<u>N17821</u>	<u>002</u>	
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CYCLOPENTOLATE HYDROCHLORIDE

SOLUTION/DROPS; OPHTHALMIC

AKPENTOLATE

<u>AT</u>	AKORN	<u>1%</u>	<u>N40164</u>	<u>001</u>	Jan 13, 1997
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<u>AT</u>		<u>2%</u>	<u>N40165</u>	<u>001</u>	Jan 13, 1997
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CYCLOGYL

<u>AT</u>	+ ALCON	<u>0.5%</u>	<u>N84109</u>	<u>001</u>	
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PRESCRIPTION DRUG PRODUCT LIST

3 - 104 (of 370)

CYCLOPENTOLATE HYDROCHLORIDE

SOLUTION/DROPS; OPHTHALMIC

CYCLOGYL

<u>AT</u>	+	ALCON	<u>1%</u>	<u>N84110</u>	<u>001</u>	
<u>AT</u>	+		<u>2%</u>	<u>N84108</u>	<u>001</u>	
<u>CYCLOPENTOLATE HYDROCHLORIDE</u>						
<u>AT</u>		ALCON UNIVERSAL	<u>1%</u>	<u>N89162</u>	<u>001</u>	Jan 24, 1991
<u>PENTOLAIR</u>						
<u>AT</u>		BAUSCH AND LOMB	<u>1%</u>	<u>N40075</u>	<u>001</u>	Apr 29, 1994

CYCLOPENTOLATE HYDROCHLORIDE; PHENYLEPHRINE HYDROCHLORIDE

SOLUTION/DROPS; OPHTHALMIC

	+	ALCON	0.2%;1%	N84300	001	
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CYCLOPHOSPHAMIDE

INJECTABLE; INJECTION

CYCLOPHOSPHAMIDE

<u>AP</u>		BAXTER HLTHCARE	<u>100MG/VIAL</u>	<u>N88371</u>	<u>001</u>	Jul 03, 1986
<u>AP</u>			<u>200MG/VIAL</u>	<u>N88372</u>	<u>001</u>	Jul 03, 1986
<u>AP</u>			<u>500MG/VIAL</u>	<u>N88373</u>	<u>001</u>	Jul 03, 1986
<u>AP</u>			<u>1GM/VIAL</u>	<u>N88374</u>	<u>001</u>	Sep 24, 1986
<u>LYOPHILIZED CYTOXAN</u>						
<u>AP</u>	+	BRISTOL MYERS SQUIBB	<u>100MG/VIAL</u>	<u>N12142</u>	<u>006</u>	Dec 05, 1985
<u>AP</u>	+		<u>200MG/VIAL</u>	<u>N12142</u>	<u>007</u>	Dec 10, 1985
<u>AP</u>	+		<u>500MG/VIAL</u>	<u>N12142</u>	<u>008</u>	Jan 04, 1984
<u>AP</u>	+		<u>1GM/VIAL</u>	<u>N12142</u>	<u>010</u>	Sep 24, 1985
<u>AP</u>	+		<u>2GM/VIAL</u>	<u>N12142</u>	<u>009</u>	Dec 10, 1984
<u>NEOSAR</u>						
<u>AP</u>		SICOR PHARMS	<u>100MG/VIAL</u>	<u>N87442</u>	<u>001</u>	Feb 16, 1982
<u>AP</u>			<u>200MG/VIAL</u>	<u>N87442</u>	<u>002</u>	Feb 16, 1982
<u>AP</u>			<u>500MG/VIAL</u>	<u>N87442</u>	<u>003</u>	Feb 16, 1982
<u>AP</u>			<u>1GM/VIAL</u>	<u>N87442</u>	<u>004</u>	Jul 08, 1983
<u>AP</u>			<u>2GM/VIAL</u>	<u>N87442</u>	<u>005</u>	Mar 30, 1989

TABLET; ORAL

CYCLOPHOSPHAMIDE

<u>AB</u>		ROXANE	<u>25MG</u>	<u>N40032</u>	<u>001</u>	Aug 17, 1999
<u>AB</u>			<u>50MG</u>	<u>N40032</u>	<u>002</u>	Aug 17, 1999
<u>CYTOXAN</u>						
<u>AB</u>		BRISTOL MYERS SQUIBB	<u>25MG</u>	<u>N12141</u>	<u>002</u>	
<u>AB</u>	+		<u>50MG</u>	<u>N12141</u>	<u>001</u>	

CYCLOSERINE

CAPSULE; ORAL

SEROMYCIN

	+	LILLY	250MG	N60593	001	
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CYCLOSPORINE

CAPSULE; ORAL

CYCLOSPORINE

<u>AB1</u>		IVAX PHARMS	<u>25MG</u>	<u>N65110</u>	<u>003</u>	Mar 29, 2005
<u>AB1</u>			<u>50MG</u>	<u>N65110</u>	<u>001</u>	Mar 29, 2005
<u>AB1</u>			<u>100MG</u>	<u>N65110</u>	<u>002</u>	Mar 29, 2005
<u>AB1</u>		PLIVA	<u>25MG</u>	<u>N65044</u>	<u>002</u>	Dec 20, 2000
<u>AB1</u>			<u>100MG</u>	<u>N65044</u>	<u>001</u>	Dec 20, 2000
<u>AB1</u>		SANDOZ	<u>25MG</u>	<u>N65017</u>	<u>002</u>	Jan 13, 2000
<u>AB1</u>			<u>100MG</u>	<u>N65017</u>	<u>001</u>	Jan 13, 2000
<u>AB2</u>		TORPHARM	<u>25MG</u>	<u>N65040</u>	<u>001</u>	May 09, 2002
<u>AB2</u>			<u>100MG</u>	<u>N65040</u>	<u>002</u>	May 09, 2002

PRESCRIPTION DRUG PRODUCT LIST

3 - 105 (of 370)

CYCLOSPORINE

CAPSULE; ORAL

GENGRAF

<u>AB1</u>	ABBOTT	<u>25MG</u>	<u>N65003</u>	<u>001</u>	May 12, 2000
<u>AB1</u>		<u>50MG</u>	<u>N65003</u>	<u>002</u>	May 12, 2000
<u>AB1</u>		<u>100MG</u>	<u>N65003</u>	<u>003</u>	May 12, 2000

NEORAL

<u>AB1</u>	NOVARTIS	<u>25MG</u>	<u>N50715</u>	<u>001</u>	Jul 14, 1995
<u>AB1</u> +		<u>100MG</u>	<u>N50715</u>	<u>002</u>	Jul 14, 1995

SANDIMMUNE

<u>AB2</u>	NOVARTIS	<u>25MG</u>	<u>N50625</u>	<u>001</u>	Mar 02, 1990
<u>AB2</u> +		<u>100MG</u>	<u>N50625</u>	<u>002</u>	Mar 02, 1990
BX		50MG	N50625	003	Nov 23, 1992

EMULSION; OPHTHALMIC

RESTASIS

+	ALLERGAN	0.05%	N50790	001	Dec 23, 2002
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INJECTABLE; INJECTION

CYCLOSPORINE

<u>AP</u>	BEDFORD	<u>50MG/ML</u>	<u>N65004</u>	<u>001</u>	Oct 29, 1999
<u>AP</u>	PHARMAFORCE	<u>50MG/ML</u>	<u>N65151</u>	<u>001</u>	Oct 07, 2003

SANDIMMUNE

<u>AP</u> +	NOVARTIS	<u>50MG/ML</u>	<u>N50573</u>	<u>001</u>	Nov 14, 1983
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SOLUTION; ORAL

CYCLOSPORINE

<u>AB1</u>	ABBOTT	<u>100MG/ML</u>	<u>N65025</u>	<u>001</u>	Mar 03, 2000
<u>AB1</u>	IVAX PHARMS	<u>100MG/ML</u>	<u>N65078</u>	<u>001</u>	Mar 25, 2005
<u>AB2</u>	MORTON GROVE	<u>100MG/ML</u>	<u>N65133</u>	<u>001</u>	Sep 17, 2004
<u>AB1</u>	NOVEX	<u>100MG/ML</u>	<u>N65167</u>	<u>001</u>	Jan 05, 2005
<u>AB1</u>	PLIVA	<u>100MG/ML</u>	<u>N65054</u>	<u>001</u>	Dec 18, 2001

NEORAL

<u>AB1</u> +	NOVARTIS	<u>100MG/ML</u>	<u>N50716</u>	<u>001</u>	Jul 14, 1995
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SANDIMMUNE

<u>AB2</u> +	NOVARTIS	<u>100MG/ML</u>	<u>N50574</u>	<u>001</u>	Nov 14, 1983
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CYPROHEPTADINE HYDROCHLORIDE

SYRUP; ORAL

CYPROHEPTADINE HYDROCHLORIDE

<u>AA</u> +	ACTAVIS MID ATLANTIC	<u>2MG/5ML</u>	<u>N86833</u>	<u>001</u>	
<u>AA</u>	LYNE	<u>2MG/5ML</u>	<u>N40668</u>	<u>001</u>	Jun 28, 2006

TABLET; ORAL

CYPROHEPTADINE HYDROCHLORIDE

<u>AA</u>	COREPHARMA	<u>4MG</u>	<u>N40537</u>	<u>001</u>	Sep 30, 2003
<u>AA</u> +	IVAX PHARMS	<u>4MG</u>	<u>N87056</u>	<u>001</u>	
<u>AA</u>	PAR PHARM	<u>4MG</u>	<u>N87129</u>	<u>001</u>	
<u>AA</u>	PLIVA	<u>4MG</u>	<u>N88205</u>	<u>001</u>	Jul 26, 1983

CYPROHEPTADINE HYDROCHLORIDE

<u>AA</u>	STASON PHARMS	<u>4MG</u>	<u>N40644</u>	<u>001</u>	May 30, 2006
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CYSTEAMINE BITARTRATE

CAPSULE; ORAL

CYSTAGON

	MYLAN	EQ 50MG BASE	N20392	001	Aug 15, 1994
+		EQ 150MG BASE	N20392	002	Aug 15, 1994

CYTARABINE

INJECTABLE; INJECTION

CYTARABINE

<u>AP</u>	ABRAXIS PHARM	<u>100MG/ML</u>	<u>N76512</u>	<u>001</u>	Jan 15, 2004
<u>AP</u>	BEDFORD	<u>100MG/VIAL</u>	<u>N71471</u>	<u>001</u>	Aug 02, 1989

PRESCRIPTION DRUG PRODUCT LIST

3 - 106 (of 370)

CYTARABINE

INJECTABLE; INJECTION

CYTARABINE

<u>AP</u>	BEDFORD	<u>500MG/VIAL</u>	<u>N71472</u>	<u>001</u>	Aug 02, 1989
<u>AP</u>		<u>1GM/VIAL</u>	<u>N74245</u>	<u>001</u>	Aug 31, 1994
<u>AP</u>		<u>2GM/VIAL</u>	<u>N74245</u>	<u>002</u>	Aug 31, 1994
<u>AP</u>	+ MAYNE PHARMA USA	<u>100MG/ML</u>	<u>N75383</u>	<u>001</u>	Nov 22, 1999
	+	20MG/ML	N71868	001	Jun 04, 1990
	+	20MG/ML	N72168	001	Aug 31, 1990
	+	20MG/ML	N72945	001	Feb 28, 1994
	<u>CYTOSAR-U</u>				
<u>AP</u>	SICOR PHARMS	<u>100MG/VIAL</u>	<u>N75206</u>	<u>001</u>	Dec 30, 1998
<u>AP</u>	+	<u>500MG/VIAL</u>	<u>N75206</u>	<u>002</u>	Dec 30, 1998
<u>AP</u>	+	<u>1GM/VIAL</u>	<u>N75206</u>	<u>004</u>	Dec 30, 1998
<u>AP</u>	+	<u>2GM/VIAL</u>	<u>N75206</u>	<u>003</u>	Dec 30, 1998
	INJECTABLE, LIPOSOMAL; INJECTION				
	DEPOCYT				
	+ SKYEPHARMA	10MG/ML	N21041	001	Apr 01, 1999

DACARBAZINE

INJECTABLE; INJECTION

DACARBAZINE

<u>AP</u>	ABRAXIS PHARM	<u>100MG/VIAL</u>	<u>N75371</u>	<u>001</u>	Aug 27, 1999
<u>AP</u>		<u>200MG/VIAL</u>	<u>N75371</u>	<u>002</u>	Aug 27, 1999
<u>AP</u>	BEDFORD	<u>200MG/VIAL</u>	<u>N75812</u>	<u>001</u>	Jun 15, 2001
<u>AP</u>		<u>500MG/VIAL</u>	<u>N75812</u>	<u>002</u>	Oct 31, 2002
<u>AP</u>	MAYNE PHARMA USA	<u>200MG/VIAL</u>	<u>N75940</u>	<u>001</u>	Oct 18, 2001
<u>AP</u>	SICOR PHARMS	<u>200MG/VIAL</u>	<u>N75259</u>	<u>002</u>	Aug 27, 1998
<u>AP</u>	+	<u>500MG/VIAL</u>	<u>N75259</u>	<u>001</u>	Sep 22, 2000
	<u>DTIC-DOME</u>				
<u>AP</u>	+ BAYER PHARMS	<u>100MG/VIAL</u>	<u>N17575</u>	<u>001</u>	
<u>AP</u>	+	<u>200MG/VIAL</u>	<u>N17575</u>	<u>002</u>	

DACTINOMYCIN

INJECTABLE; INJECTION

COSMEGEN

+	OVATION PHARMS	0.5MG/VIAL	N50682	001	
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DALFOPRISTIN; QUINUPRISTIN

INJECTABLE; IV (INFUSION)

+	KING PHARMS	350MG/VIAL;150MG/VIAL	N50748	001	Sep 21, 1999
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DALTEPARIN SODIUM

INJECTABLE; INJECTION

FRAGMIN

+	PHARMACIA AND UPJOHN	2,500 IU/0.2ML	N20287	001	Dec 22, 1994
+		5,000 IU/0.2ML	N20287	003	Mar 18, 1996
+		7,500 IU/0.3ML	N20287	005	Apr 04, 2002
+		10,000 IU/ML	N20287	004	Jan 30, 1998
+		95,000IU/3.8ML (25,000IU/ML)	N20287	006	Apr 04, 2002
+		95,000IU/9.5ML (10,000IU/ML)	N20287	007	Apr 04, 2002

DANAZOL

CAPSULE; ORAL

DANAZOL

<u>AB</u>	+ BARR	<u>200MG</u>	<u>N74582</u>	<u>001</u>	Aug 09, 1996
		50MG	N74582	003	May 29, 1998
		100MG	N74582	002	May 29, 1998

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 107 (of 370)

DANAZOL

CAPSULE; ORAL

DANAZOL

<u>AB</u>	LANNETT	<u>200MG</u>	<u>N77246</u>	<u>001</u>	Sep 28, 2005
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DANTROLENE SODIUM

CAPSULE; ORAL

DANTRIUM

<u>AB</u>	PROCTER AND GAMBLE	<u>25MG</u>	<u>N17443</u>	<u>001</u>	
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<u>AB</u>		<u>50MG</u>	<u>N17443</u>	<u>003</u>	
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<u>AB</u>	+	<u>100MG</u>	<u>N17443</u>	<u>002</u>	
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DANTROLENE SODIUM

<u>AB</u>	ACTAVIS TOTOWA	<u>25MG</u>	<u>N76686</u>	<u>001</u>	Oct 24, 2005
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<u>AB</u>		<u>50MG</u>	<u>N76686</u>	<u>002</u>	Oct 24, 2005
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<u>AB</u>		<u>100MG</u>	<u>N76686</u>	<u>003</u>	Oct 24, 2005
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<u>AB</u>	IMPAX LABS	<u>25MG</u>	<u>N76856</u>	<u>001</u>	Mar 01, 2005
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<u>AB</u>		<u>50MG</u>	<u>N76856</u>	<u>002</u>	Mar 01, 2005
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<u>AB</u>		<u>100MG</u>	<u>N76856</u>	<u>003</u>	Mar 01, 2005
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INJECTABLE; INJECTION

DANTRIUM

+	PROCTER AND GAMBLE	20MG/VIAL	N18264	001	
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DAPIPRAZOLE HYDROCHLORIDE

SOLUTION/DROPS; OPHTHALMIC

REV-EYES

+	ANGELINI	0.5%	N19849	001	Dec 31, 1990
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DAPSONE

TABLET; ORAL

DAPSONE

	JACOBUS	25MG	N86841	001	
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+		100MG	N86842	001	
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DAPTOMYCIN

INJECTABLE; IV (INFUSION)

+	CUBIST	500MG/VIAL	N21572	002	Sep 12, 2003
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DARIFENACIN HYDROBROMIDE

TABLET, EXTENDED RELEASE; ORAL

ENABLEX

	NOVARTIS	EQ 7.5MG BASE	N21513	001	Dec 22, 2004
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+		EQ 15MG BASE	N21513	002	Dec 22, 2004
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DARUNAVIR ETHANOLATE

TABLET; ORAL

PREZISTA

+	TIBOTEC	EQ 300MG BASE	N21976	001	Jun 23, 2006
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DASATINIB

TABLET; ORAL

SPRYCEL

	BRISTOL MYERS SQUIBB	20MG	N21986	001	Jun 28, 2006
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		50MG	N21986	002	Jun 28, 2006
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+		70MG	N21986	003	Jun 28, 2006
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27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 108 (of 370)

DAUNORUBICIN CITRATE

INJECTABLE, LIPOSOMAL; INJECTION

DAUNOXOME

+ GILEAD

EQ 2MG BASE/ML

N50704 002

Apr 08, 1996

DAUNORUBICIN HYDROCHLORIDE

INJECTABLE; INJECTION

CERUBIDINEAP + BEDFORDEQ 20MG BASE/VIALN64103 001

Feb 03, 1995

DAUNORUBICIN HYDROCHLORIDEAP ABRAXIS PHARMEQ 5MG BASE/VIALN65034 001

Nov 20, 2001

APEQ 20MG BASE/VIALN65000 001

May 25, 1999

AP + BEDFORDEQ 5MG BASE/MLN50731 001

Jan 30, 1998

AP SICOR PHARMSEQ 5MG BASE/MLN65035 001

Jan 24, 2000

APEQ 20MG BASE/VIALN64212 001

Jun 23, 1998

+

EQ 50MG BASE/VIAL

N64212 002

May 03, 1999

DECITABINE

INJECTABLE; INTRAVENOUS

DACOGEN

+ MGI PHARMA INC

50MG/VIAL

N21790 001

May 02, 2006

DEFERASIROX

TABLET, FOR SUSPENSION; ORAL

EXJADE

NOVARTIS

125MG

N21882 001

Nov 02, 2005

250MG

N21882 002

Nov 02, 2005

+

500MG

N21882 003

Nov 02, 2005

DEFEROXAMINE MESYLATE

INJECTABLE; INJECTION

DEFEROXAMINE MESYLATEAP HOSPIRA500MG/VIALN76019 001

Mar 17, 2004

AP2GM/VIALN76019 002

Mar 17, 2004

AP SICOR PHARMS500MG/VIALN76806 001

Mar 31, 2006

AP2GM/VIALN76806 002

Mar 31, 2006

DESFERALAP + NOVARTIS500MG/VIALN16267 001AP +2GM/VIALN16267 002

May 25, 2000

DELAVIRDINE MESYLATE

TABLET; ORAL

RESCRIPTOR

AGOURON

100MG

N20705 001

Apr 04, 1997

+

200MG

N20705 002

Jul 14, 1999

DEMECLOCYCLINE HYDROCHLORIDE

TABLET; ORAL

DECLOMYCINAB GLADES PHARMS LLC150MGN50261 002AB +300MGN50261 003DEMECLOCYCLINE HYDROCHLORIDEAB BARR150MGN65171 001

Dec 13, 2004

AB300MGN65171 002

Dec 13, 2004

AB IMPAX LABS150MGN65094 001

Mar 22, 2004

AB300MGN65094 002

Mar 22, 2004

PRESCRIPTION DRUG PRODUCT LIST

3 - 109 (of 370)

DESFLURANE

LIQUID; INHALATION
SUPRANE

+ BAXTER HLTHCARE CORP 99.9%	N20118	001	Sep 18, 1992
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DESIPRAMINE HYDROCHLORIDE

TABLET; ORAL

DESIPRAMINE HYDROCHLORIDE

<u>AB</u>	PLIVA	<u>25MG</u>	<u>N71800</u>	<u>001</u>	Dec 08, 1987
<u>AB</u>		<u>50MG</u>	<u>N71801</u>	<u>001</u>	Dec 08, 1987
<u>AB</u>		<u>75MG</u>	<u>N71802</u>	<u>001</u>	Dec 08, 1987
<u>AB</u>		<u>100MG</u>	<u>N71803</u>	<u>001</u>	May 29, 1997
<u>AB</u>		<u>150MG</u>	<u>N71804</u>	<u>001</u>	May 29, 1997
<u>AB</u>	SANDOZ	<u>10MG</u>	<u>N72099</u>	<u>001</u>	May 24, 1988
<u>AB</u>		<u>10MG</u>	<u>N74430</u>	<u>001</u>	Feb 09, 1996
<u>AB</u>		<u>25MG</u>	<u>N71601</u>	<u>001</u>	Jun 05, 1987
<u>AB</u>		<u>25MG</u>	<u>N72100</u>	<u>001</u>	May 24, 1988
<u>AB</u>		<u>50MG</u>	<u>N71588</u>	<u>001</u>	Jun 05, 1987
<u>AB</u>		<u>50MG</u>	<u>N72101</u>	<u>001</u>	May 24, 1988
<u>AB</u>		<u>75MG</u>	<u>N71602</u>	<u>001</u>	Oct 05, 1987
<u>AB</u>		<u>75MG</u>	<u>N72102</u>	<u>001</u>	Jun 20, 1988
<u>AB</u>		<u>100MG</u>	<u>N71766</u>	<u>001</u>	Oct 05, 1987
<u>AB</u>		<u>100MG</u>	<u>N72103</u>	<u>001</u>	Jun 20, 1988
<u>AB</u>		<u>150MG</u>	<u>N72104</u>	<u>001</u>	Jun 20, 1988
<u>AB</u>		<u>150MG</u>	<u>N74430</u>	<u>002</u>	Feb 09, 1996

NORPRAMIN

<u>AB</u>	SANOFI AVENTIS US	<u>10MG</u>	<u>N14399</u>	<u>007</u>	Feb 11, 1982
<u>AB</u>		<u>25MG</u>	<u>N14399</u>	<u>001</u>	
<u>AB</u>	+	<u>50MG</u>	<u>N14399</u>	<u>003</u>	
<u>AB</u>		<u>75MG</u>	<u>N14399</u>	<u>004</u>	
<u>AB</u>	+	<u>100MG</u>	<u>N14399</u>	<u>005</u>	
<u>AB</u>		<u>150MG</u>	<u>N14399</u>	<u>006</u>	

DESIRUDIN RECOMBINANT

INJECTABLE; SUBCUTANEOUS
IPRIVASK

+ CANYON	15MG/VIAL	N21271	001	Apr 04, 2003
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DESLORATADINE

SYRUP; ORAL
CLARINEX

+ SCHERING	0.5MG/ML	N21300	001	Sep 01, 2004
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TABLET; ORAL
CLARINEX

+ SCHERING PLOUGH	5MG	N21165	001	Dec 21, 2001
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TABLET, ORALLY DISINTEGRATING; ORAL
CLARINEX

SCHERING	2.5MG	N21312	002	Jul 14, 2005
+	5MG	N21312	001	Jun 26, 2002

DESLORATADINE; PSEUDOEPHEDRINE SULFATE

TABLET, EXTENDED RELEASE; ORAL
CLARINEX D 24 HOUR

+ SCHERING	5MG;240MG	N21605	001	Mar 03, 2005
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CLARINEX-D 12 HOUR

+ SCHERING	2.5MG;120MG	N21313	001	Feb 01, 2006
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PRESCRIPTION DRUG PRODUCT LIST

3 - 110 (of 370)

DESMOPRESSIN ACETATE

INJECTABLE; INJECTION

DDAVP

<u>AP</u>	+ SANOFI AVENTIS US	<u>0.004MG/ML</u>	<u>N18938</u>	<u>001</u>	Mar 30, 1984
	+	0.015MG/ML	N18938	002	Apr 25, 1995

DESMOPRESSIN ACETATE

<u>AP</u>	HOSPIRA	<u>0.004MG/ML</u>	<u>N75220</u>	<u>001</u>	Aug 28, 2000
<u>AP</u>	SICOR PHARMS	<u>0.004MG/ML</u>	<u>N74888</u>	<u>001</u>	Oct 15, 1997

SOLUTION; NASAL

DDAVP

	+ SANOFI AVENTIS US	0.01%	N17922	001	
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SPRAY, METERED; NASAL

DDAVP (NEEDS NO REFRIGERATION)

<u>AB</u>	+ SANOFI AVENTIS US	<u>0.01MG/SPRAY</u>	<u>N17922</u>	<u>003</u>	Aug 07, 1996
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DESMOPRESSIN ACETATE

<u>AB</u>	+ BAUSCH AND LOMB	<u>0.01MG/SPRAY</u>	<u>N74830</u>	<u>001</u>	Jan 25, 1999
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DESMOPRESSIN ACETATE (NEEDS NO REFRIGERATION)

<u>AB</u>	APOTEX INC	<u>0.01MG/SPRAY</u>	<u>N76703</u>	<u>001</u>	Jan 27, 2005
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MINIRIN

<u>AB</u>	+ FERRING	<u>0.01MG/SPRAY</u>	<u>N21333</u>	<u>001</u>	Sep 16, 2002
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STIMATE

	+ ZLB BEHRING	0.15MG/SPRAY	N20355	001	Mar 07, 1994
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TABLET; ORAL

DDAVP

<u>AB</u>	SANOFI AVENTIS US	<u>0.1MG</u>	<u>N19955</u>	<u>001</u>	Sep 06, 1995
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<u>AB</u>	+	<u>0.2MG</u>	<u>N19955</u>	<u>002</u>	Sep 06, 1995
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DESMOPRESSIN ACETATE

<u>AB</u>	APOTEX INC	<u>0.1MG</u>	<u>N77414</u>	<u>001</u>	Mar 07, 2006
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<u>AB</u>		<u>0.2MG</u>	<u>N77414</u>	<u>002</u>	Mar 07, 2006
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<u>AB</u>	BARR	<u>0.1MG</u>	<u>N76470</u>	<u>001</u>	Jul 01, 2005
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<u>AB</u>		<u>0.2MG</u>	<u>N76470</u>	<u>002</u>	Jul 01, 2005
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<u>AB</u>	TEVA PHARMS	<u>0.1MG</u>	<u>N77122</u>	<u>001</u>	Jan 25, 2006
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<u>AB</u>		<u>0.2MG</u>	<u>N77122</u>	<u>002</u>	Jan 25, 2006
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DESOGESTREL; ETHINYL ESTRADIOL

TABLET; ORAL-21

DESOGESTREL AND ETHINYL ESTRADIOL

<u>AB</u>	DURAMED PHARMS BARR	<u>0.15MG;0.03MG</u>	<u>N75256</u>	<u>001</u>	Aug 12, 1999
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TABLET; ORAL-28

CYCLESSA

<u>AB</u>	+ ORGANON USA INC	<u>0.1MG,0.125MG,0.15MG;0.025MG,0.025MG,0.025MG</u>	<u>N21090</u>	<u>001</u>	Dec 20, 2000
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DESOGEN

<u>AB</u>	ORGANON USA INC	<u>0.15MG;0.03MG</u>	<u>N20071</u>	<u>002</u>	Dec 10, 1992
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DESOGESTREL AND ETHINYL ESTRADIOL

<u>AB</u>	DURAMED PHARMS BARR	<u>0.15MG;0.03MG</u>	<u>N75256</u>	<u>002</u>	Aug 12, 1999
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<u>AB</u>	WATSON LABS	<u>0.15MG;0.03MG</u>	<u>N76915</u>	<u>001</u>	Jul 29, 2005
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KARIVA

<u>AB</u>	BARR	<u>0.15MG,N/A;0.02MG,0.01MG</u>	<u>N75863</u>	<u>001</u>	Apr 05, 2002
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MIRCETTE

<u>AB</u>	+ DURAMED	<u>0.15MG,N/A;0.02MG,0.01MG</u>	<u>N20713</u>	<u>001</u>	Apr 22, 1998
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ORTHO-CEPT

<u>AB</u>	+ ORTHO MCNEIL PHARM	<u>0.15MG;0.03MG</u>	<u>N20301</u>	<u>002</u>	Dec 14, 1992
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VELIVET

<u>AB</u>	DURAMED PHARMS BARR	<u>0.1MG,0.125MG,0.15MG;0.025MG,0.025MG,0.025MG</u>	<u>N76455</u>	<u>001</u>	Feb 24, 2004
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27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 111 (of 370)

DESONIDE

AEROSOL, FOAM; TOPICAL					
VERDESO					
	+ CONNETICS	0.05%	N21978	001	Sep 19, 2006
CREAM; TOPICAL					
<u>DESONIDE</u>					
<u>AB</u>	TARO	<u>0.05%</u>	<u>N73548</u>	<u>001</u>	Jun 30, 1992
<u>AB</u>	TEVA PHARMS	<u>0.05%</u>	<u>N74027</u>	<u>001</u>	Sep 28, 1992
<u>DESOWEN</u>					
<u>AB</u>	GALDERMA LABS LP	<u>0.05%</u>	<u>N19048</u>	<u>001</u>	Dec 14, 1984
<u>TRIDESILON</u>					
<u>AB</u>	+ PERRIGO NEW YORK	<u>0.05%</u>	<u>N17010</u>	<u>001</u>	
GEL; TOPICAL					
DESONATE					
	+ SKINMEDICA	0.05%	N21844	001	Oct 20, 2006
LOTION; TOPICAL					
<u>DESONIDE</u>					
<u>AB</u>	ALTANA	<u>0.05%</u>	<u>N75860</u>	<u>001</u>	Mar 19, 2002
<u>DESOWEN</u>					
<u>AB</u>	+ GALDERMA LABS LP	<u>0.05%</u>	<u>N72354</u>	<u>001</u>	Jan 24, 1992
OINTMENT; TOPICAL					
<u>DESONIDE</u>					
<u>AB</u>	ALTANA	<u>0.05%</u>	<u>N75751</u>	<u>001</u>	Mar 12, 2001
<u>AB</u>	TARO	<u>0.05%</u>	<u>N74254</u>	<u>001</u>	Aug 03, 1994
<u>DESOWEN</u>					
<u>AB</u>	GALDERMA LABS LP	<u>0.05%</u>	<u>N71425</u>	<u>001</u>	Jun 15, 1988
<u>TRIDESILON</u>					
<u>AB</u>	+ PERRIGO NEW YORK	<u>0.05%</u>	<u>N17426</u>	<u>001</u>	

DESOXIMETASONE

CREAM; TOPICAL					
<u>DESOXIMETASONE</u>					
<u>AB</u>	PERRIGO NEW YORK	<u>0.25%</u>	<u>N76510</u>	<u>001</u>	Jul 01, 2003
<u>AB</u>	TARO	<u>0.05%</u>	<u>N73210</u>	<u>001</u>	Nov 30, 1990
<u>AB</u>		<u>0.25%</u>	<u>N73193</u>	<u>001</u>	Nov 30, 1990
<u>TOPICORT</u>					
<u>AB</u>	+ TARO PHARMS NORTH	<u>0.25%</u>	<u>N17856</u>	<u>001</u>	
<u>TOPICORT LP</u>					
<u>AB</u>	+ TARO PHARMS NORTH	<u>0.05%</u>	<u>N18309</u>	<u>001</u>	
GEL; TOPICAL					
<u>DESOXIMETASONE</u>					
<u>AB</u>	PERRIGO NEW YORK	<u>0.05%</u>	<u>N77552</u>	<u>001</u>	Jan 09, 2006
<u>AB</u>	TARO	<u>0.05%</u>	<u>N74904</u>	<u>001</u>	Jul 14, 1998
<u>TOPICORT</u>					
<u>AB</u>	+ TARO PHARMS NORTH	<u>0.05%</u>	<u>N18586</u>	<u>001</u>	Mar 29, 1982
OINTMENT; TOPICAL					
<u>DESOXIMETASONE</u>					
<u>AB</u>	TARO	<u>0.25%</u>	<u>N74286</u>	<u>001</u>	Jun 07, 1996
<u>TOPICORT</u>					
<u>AB</u>	+ TARO PHARMS NORTH	<u>0.25%</u>	<u>N18763</u>	<u>001</u>	Sep 30, 1983

DEXAMETHASONE

CONCENTRATE; ORAL					
DEXAMETHASONE INTENSOL					
	+ ROXANE	1MG/ML	N88252	001	Sep 01, 1983
ELIXIR; ORAL					
<u>DEXAMETHASONE</u>					
<u>AA</u>	+ ACTAVIS MID ATLANTIC	<u>0.5MG/5ML</u>	<u>N84754</u>	<u>001</u>	

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 112 (of 370)

DEXAMETHASONE

ELIXIR; ORAL					
<u>MYMETHASONE</u>					
<u>AA</u>	+	MORTON GROVE	<u>0.5MG/5ML</u>	<u>N88254</u>	<u>001</u> Jul 27, 1983
SOLUTION; ORAL					
DEXAMETHASONE					
	+	ROXANE	0.5MG/5ML	N88248	001 Sep 01, 1983
SUSPENSION/DROPS; OPHTHALMIC					
	+	ALCON	0.1%	N13422	001
TABLET; ORAL					
DEXAMETHASONE					
BP		PAR PHARM	0.5MG	N88148	001 Apr 28, 1983
BP			0.75MG	N88160	001 Apr 28, 1983
BP			1.5MG	N88237	001 Apr 28, 1983
BP			4MG	N88238	001 Apr 28, 1983
BP			6MG	N88481	001 Nov 28, 1983
			0.25MG	N88149	001 Apr 28, 1983
BP		ROXANE	0.5MG	N84611	001
BP			0.75MG	N84613	001
BP			1.5MG	N84610	001
BP			4MG	N84612	001
BP	+		6MG	N88316	001 Sep 15, 1983
			1MG	N88306	001 Sep 15, 1983
			2MG	N87916	001 Aug 26, 1982

DEXAMETHASONE SODIUM PHOSPHATE

INJECTABLE; INJECTION					
<u>DEXAMETHASONE</u>					
<u>AP</u>		BAXTER HLTHCARE	<u>EQ 4MG PHOSPHATE/ML</u>	<u>N84282</u>	<u>001</u>
<u>AP</u>	+		<u>EQ 10MG PHOSPHATE/ML</u>	<u>N87702</u>	<u>001</u> Sep 07, 1982
<u>DEXAMETHASONE SODIUM PHOSPHATE</u>					
<u>AP</u>		ABRAXIS PHARM	<u>EQ 4MG PHOSPHATE/ML</u>	<u>N84916</u>	<u>001</u>
<u>AP</u>			<u>EQ 10MG PHOSPHATE/ML</u>	<u>N40491</u>	<u>001</u> Apr 11, 2003
<u>AP</u>			<u>EQ 10MG PHOSPHATE/ML</u>	<u>N40572</u>	<u>001</u> Apr 22, 2005
<u>AP</u>	+	LUITPOLD	<u>EQ 4MG PHOSPHATE/ML</u>	<u>N87440</u>	<u>001</u> Jul 21, 1982
<u>AP</u>	+	SICOR PHARMS	<u>EQ 10MG PHOSPHATE/ML</u>	<u>N81126</u>	<u>001</u> Aug 31, 1990
SOLUTION/DROPS; OPHTHALMIC, OTIC					
<u>DEXAMETHASONE SODIUM PHOSPHATE</u>					
<u>AT</u>	+	ALCON UNIVERSAL	<u>EQ 0.1% PHOSPHATE</u>	<u>N88771</u>	<u>001</u> Jan 16, 1985
<u>AT</u>		BAUSCH AND LOMB	<u>EQ 0.1% PHOSPHATE</u>	<u>N40069</u>	<u>001</u> Jul 26, 1996

DEXAMETHASONE; NEOMYCIN SULFATE; POLYMYXIN B SULFATE

OINTMENT; OPHTHALMIC					
<u>MAXITROL</u>					
<u>AT</u>	+	FALCON PHARMS	<u>0.1%;EQ 3.5MG BASE/GM;10,000 UNITS/GM</u>	<u>N50065</u>	<u>002</u>
<u>NEOMYCIN AND POLYMYXIN B SULFATES AND DEXAMETHASONE</u>					
<u>AT</u>		BAUSCH AND LOMB	<u>0.1%;EQ 3.5MG BASE/GM;10,000 UNITS/GM</u>	<u>N64063</u>	<u>001</u> Jul 25, 1994
<u>AT</u>		FOUGERA	<u>0.1%;EQ 3.5MG BASE/GM;10,000 UNITS/GM</u>	<u>N62938</u>	<u>001</u> Jul 31, 1989
SUSPENSION/DROPS; OPHTHALMIC					
<u>DEXASPORIN</u>					
<u>AT</u>		BAUSCH AND LOMB	<u>0.1%;EQ 3.5MG BASE/ML;10,000 UNITS/ML</u>	<u>N64135</u>	<u>001</u> Sep 13, 1995
<u>MAXITROL</u>					
<u>AT</u>		ALCON	<u>0.1%;EQ 3.5MG BASE/ML;10,000 UNITS/ML</u>	<u>N62341</u>	<u>001</u> May 22, 1984
<u>AT</u>	+	FALCON PHARMS	<u>0.1%;EQ 3.5MG BASE/ML;10,000 UNITS/ML</u>	<u>N50023</u>	<u>002</u>
<u>NEOMYCIN AND POLYMYXIN B SULFATES AND DEXAMETHASONE</u>					
<u>AT</u>		ALCON UNIVERSAL	<u>0.1%;EQ 3.5MG BASE/ML;10,000 UNITS/ML</u>	<u>N62721</u>	<u>001</u> Nov 17, 1986

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 113 (of 370)

DEXAMETHASONE; TOBRAMYCIN

OINTMENT; OPHTHALMIC

TOBRADEX

+ ALCON 0.1%;0.3%

N50616 001 Sep 28, 1988

SUSPENSION/DROPS; OPHTHALMIC

TOBRADEXAB + ALCON 0.1%;0.3%N50592 001 Aug 18, 1988TOBRAMYCIN AND DEXAMETHASONEAB BAUSCH AND LOMB 0.1%;0.3%N64134 001 Oct 27, 1999DEXCHLORPHENIRAMINE MALEATE

SYRUP; ORAL

MYLARAMINE

+ MORTON GROVE 2MG/5ML

N88251 001 Mar 23, 1984

TABLET; ORAL

DEXCHLORPHENIRAMINE MALEATE

+ PLIVA 2MG

N88682 001 Jan 17, 1986

DEXMEDETOMIDINE

INJECTABLE; INJECTION

PRECEDEX

+ HOSPIRA EQ 100UGM BASE/ML

N21038 001 Dec 17, 1999

DEXMETHYLPHENIDATE HYDROCHLORIDE

CAPSULE, EXTENDED RELEASE; ORAL

FOCALIN XR

NOVARTIS 5MG

N21802 001 May 26, 2005

10MG

N21802 002 May 26, 2005

15MG

N21802 004 Aug 01, 2006

+ 20MG

N21802 003 May 26, 2005

TABLET; ORAL

FOCALIN

NOVARTIS 2.5MG

N21278 001 Nov 13, 2001

5MG

N21278 002 Nov 13, 2001

+ 10MG

N21278 003 Nov 13, 2001

DEXRAZOXANE HYDROCHLORIDE

INJECTABLE; INJECTION

DEXRAZOXANEAP BEDFORD EQ 250MG BASE/VIALN76068 001 Sep 28, 2004AP EQ 500MG BASE/VIALN76068 002 Sep 28, 2004ZINECARDAP + PHARMACIA AND UPJOHN EQ 250MG BASE/VIALN20212 001 May 26, 1995AP + EQ 500MG BASE/VIALN20212 002 May 26, 1995DEXTROAMPHETAMINE SULFATE

CAPSULE, EXTENDED RELEASE; ORAL

DEXEDRINEAB SMITHKLINE BEECHAM 5MGN17078 001AB 10MGN17078 002AB + 15MGN17078 003DEXTROAMPHETAMINE SULFATEAB BARR 5MGN76137 001 Jan 18, 2002AB 10MGN76137 002 Jan 18, 2002AB 15MGN76137 003 Jan 18, 2002AB MALLINCKRODT 5MGN76353 001 May 06, 2003AB 10MGN76353 002 May 06, 2003AB 15MGN76353 003 May 06, 2003

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 114 (of 370)

DEXTROAMPHETAMINE SULFATE

TABLET; ORAL

DEXEDRINE

<u>AA</u>	GLAXOSMITHKLINE	<u>5MG</u>	<u>N84935</u>	<u>001</u>	
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DEXTROAMPHETAMINE SULFATE

<u>AA</u>	BARR	<u>5MG</u>	<u>N40361</u>	<u>001</u>	Jan 31, 2001
<u>AA</u>		<u>10MG</u>	<u>N40361</u>	<u>002</u>	Jan 31, 2001
<u>AA</u>	KV PHARM	<u>5MG</u>	<u>N40365</u>	<u>001</u>	Oct 31, 2002
<u>AA</u>		<u>10MG</u>	<u>N40367</u>	<u>001</u>	Oct 31, 2002
<u>AA</u>	MALLINCKRODT	<u>5MG</u>	<u>N40436</u>	<u>001</u>	Jan 29, 2002
<u>AA</u>		<u>10MG</u>	<u>N40436</u>	<u>002</u>	Jan 29, 2002

DEXTROSTAT

<u>AA</u>	+ SHIRE RICHWOOD	<u>5MG</u>	<u>N84051</u>	<u>001</u>	
<u>AA</u>	+	<u>10MG</u>	<u>N84051</u>	<u>002</u>	

DEXTROMETHORPHAN HYDROBROMIDE; PROMETHAZINE HYDROCHLORIDE

SYRUP; ORAL

PROMETH W/ DEXTROMETHORPHAN

<u>AA</u>	+ ACTAVIS MID ATLANTIC	<u>15MG/5ML; 6.25MG/5ML</u>	<u>N88762</u>	<u>001</u>	Oct 31, 1984
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PROMETHAZINE DM

<u>AA</u>	VINTAGE	<u>15MG/5ML; 6.25MG/5ML</u>	<u>N40649</u>	<u>001</u>	Feb 14, 2006
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PROMETHAZINE HYDROCHLORIDE AND DEXTROMETHORPHAN HYDROBROMIDE

<u>AA</u>	HI TECH PHARMA	<u>15MG/5ML; 6.25MG/5ML</u>	<u>N40027</u>	<u>001</u>	Jul 31, 1996
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PROMETHAZINE W/ DEXTROMETHORPHAN

<u>AA</u>	MORTON GROVE	<u>15MG/5ML; 6.25MG/5ML</u>	<u>N88864</u>	<u>001</u>	Jan 04, 1985
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DEXTROSE

INJECTABLE; INJECTION

DEXTROSE 10% IN PLASTIC CONTAINER

<u>AP</u>	+ B BRAUN	<u>10GM/100ML</u>	<u>N19626</u>	<u>004</u>	Feb 02, 1988
<u>AP</u>	+ BAXTER HLTHCARE	<u>10GM/100ML</u>	<u>N16694</u>	<u>001</u>	
<u>AP</u>	+ HOSPIRA	<u>10GM/100ML</u>	<u>N18080</u>	<u>001</u>	

DEXTROSE 20% IN PLASTIC CONTAINER

<u>AP</u>	+ BAXTER HLTHCARE	<u>20GM/100ML</u>	<u>N17521</u>	<u>004</u>	
<u>AP</u>	+ HOSPIRA	<u>20GM/100ML</u>	<u>N18564</u>	<u>001</u>	Mar 23, 1982

DEXTROSE 25%

+	HOSPIRA	250MG/ML	N19445	002	Nov 23, 1998
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DEXTROSE 30% IN PLASTIC CONTAINER

<u>AP</u>	+ BAXTER HLTHCARE	<u>30GM/100ML</u>	<u>N17521</u>	<u>003</u>	
<u>AP</u>	+ HOSPIRA	<u>30GM/100ML</u>	<u>N19345</u>	<u>001</u>	Jan 26, 1985

DEXTROSE 40% IN PLASTIC CONTAINER

<u>AP</u>	+ BAXTER HLTHCARE	<u>40GM/100ML</u>	<u>N17521</u>	<u>002</u>	
<u>AP</u>	+ HOSPIRA	<u>40GM/100ML</u>	<u>N18562</u>	<u>001</u>	Mar 23, 1982

DEXTROSE 5% IN PLASTIC CONTAINER

<u>AP</u>	+ B BRAUN	<u>50MG/ML</u>	<u>N16730</u>	<u>002</u>	
<u>AP</u>	+	<u>5GM/100ML</u>	<u>N16730</u>	<u>001</u>	
<u>AP</u>	+	<u>5GM/100ML</u>	<u>N19626</u>	<u>002</u>	Feb 02, 1988
<u>AP</u>	+ BAXTER HLTHCARE	<u>50MG/ML</u>	<u>N16673</u>	<u>003</u>	Oct 30, 1985
<u>AP</u>	+	<u>50MG/ML</u>	<u>N20179</u>	<u>002</u>	Dec 07, 1992
<u>AP</u>	+	<u>5GM/100ML</u>	<u>N16673</u>	<u>001</u>	
<u>AP</u>	+	<u>5GM/100ML</u>	<u>N20179</u>	<u>001</u>	Dec 07, 1992
<u>AP</u>	+ DHL	<u>5GM/100ML</u>	<u>N19971</u>	<u>001</u>	Sep 28, 1995
<u>AP</u>	+ HOSPIRA	<u>50MG/ML</u>	<u>N16367</u>	<u>002</u>	
<u>AP</u>	+	<u>50MG/ML</u>	<u>N19222</u>	<u>001</u>	Jul 13, 1984
<u>AP</u>	+	<u>5GM/100ML</u>	<u>N19466</u>	<u>001</u>	Jul 15, 1985
<u>AP</u>	+	<u>5GM/100ML</u>	<u>N19479</u>	<u>001</u>	Sep 17, 1985

DEXTROSE 50% IN PLASTIC CONTAINER

<u>AP</u>	+ BAXTER HLTHCARE	<u>50GM/100ML</u>	<u>N17521</u>	<u>001</u>	
<u>AP</u>	+	<u>50GM/100ML</u>	<u>N20047</u>	<u>001</u>	Jul 02, 1991
<u>AP</u>	HOSPIRA	<u>500MG/ML</u>	<u>N19445</u>	<u>001</u>	Jun 03, 1986

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 115 (of 370)

DEXTROSE

INJECTABLE; INJECTION

DEXTROSE 50% IN PLASTIC CONTAINER

AP	+	HOSPIRA	50GM/100ML	N18563	001	Mar 23, 1982
AP	+		50GM/100ML	N19894	001	Dec 26, 1989

DEXTROSE 60% IN PLASTIC CONTAINER

AP	+	BAXTER HLTHCARE	60GM/100ML	N17521	005	Mar 26, 1982
AP	+	HOSPIRA	60GM/100ML	N19346	001	Jan 25, 1985

DEXTROSE 70% IN PLASTIC CONTAINER

AP	+	BAXTER HLTHCARE	70GM/100ML	N17521	006	Mar 26, 1982
AP	+		70GM/100ML	N20047	003	Jul 02, 1991
AP	+	HOSPIRA	70GM/100ML	N18561	001	Mar 23, 1982
AP	+		70GM/100ML	N19893	001	Dec 26, 1989

DEXTROSE; MAGNESIUM ACETATE TETRAHYDRATE; POTASSIUM ACETATE; SODIUM CHLORIDE

INJECTABLE; INJECTION

PLASMA-LYTE 56 AND DEXTROSE 5% IN PLASTIC CONTAINER

BAXTER HLTHCARE	5GM/100ML;32MG/100ML;128MG/100ML;234MG/100ML	N17385	001
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DEXTROSE; MAGNESIUM ACETATE; POTASSIUM ACETATE; SODIUM CHLORIDE

INJECTABLE; INJECTION

NORMOSOL-M AND DEXTROSE 5% IN PLASTIC CONTAINER

HOSPIRA	5GM/100ML;21MG/100ML;128MG/100ML;234MG/100ML	N17610	001
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DEXTROSE; MAGNESIUM CHLORIDE; POTASSIUM CHLORIDE; POTASSIUM PHOSPHATE, DIBASIC; SODIUM ACETATE

INJECTABLE; INJECTION

ISOLYTE P IN DEXTROSE 5% IN PLASTIC CONTAINER

B BRAUN	5GM/100ML;31MG/100ML;130MG/100ML;26MG/100ML;320MG/100ML	N19873	001	Jun 10, 1993
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DEXTROSE; MAGNESIUM CHLORIDE; POTASSIUM CHLORIDE; POTASSIUM PHOSPHATE, DIBASIC; SODIUM CHLORIDE; SODIUM LACTATE; SODIUM PHOSPHATE, MONOBASIC, ANHYDROUS

INJECTABLE; INJECTION

IONOSOL B AND DEXTROSE 5% IN PLASTIC CONTAINER

HOSPIRA	5GM/100ML;53MG/100ML;100MG/100ML;100MG/100ML;180MG/100ML;280MG/100ML;16MG/100ML	N19515	001	May 08, 1986
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DEXTROSE; MAGNESIUM CHLORIDE; POTASSIUM CHLORIDE; POTASSIUM PHOSPHATE, MONOBASIC; SODIUM CHLORIDE; SODIUM LACTATE

INJECTABLE; INJECTION

DEXTROSE 5% AND ELECTROLYTE NO.48 IN PLASTIC CONTAINER

BAXTER HLTHCARE	5GM/100ML;31MG/100ML;141MG/100ML;20MG/100ML;12MG/100ML;260MG/100ML	N17484	001
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DEXTROSE; MAGNESIUM CHLORIDE; POTASSIUM CHLORIDE; POTASSIUM PHOSPHATE, MONOBASIC; SODIUM LACTATE; SODIUM PHOSPHATE, MONOBASIC, ANHYDROUS

INJECTABLE; INJECTION

IONOSOL MB AND DEXTROSE 5% IN PLASTIC CONTAINER

HOSPIRA	5GM/100ML;30MG/100ML;141MG/100ML;15MG/100ML;260MG/100ML;25MG/100ML	N19513	001	May 08, 1986
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DEXTROSE; MAGNESIUM CHLORIDE; POTASSIUM CHLORIDE; SODIUM ACETATE; SODIUM CHLORIDE

INJECTABLE; INJECTION

ISOLYTE H IN DEXTROSE 5% IN PLASTIC CONTAINER

B BRAUN	5GM/100ML;30MG/100ML;97MG/100ML;220MG/100ML;140MG/100ML	N19844	001	Jun 10, 1993
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27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 116 (of 370)

DEXTROSE; MAGNESIUM CHLORIDE; POTASSIUM CHLORIDE; SODIUM ACETATE; SODIUM CHLORIDE; SODIUM GLUCONATE

INJECTABLE; INJECTION

ISOLYTE S IN DEXTROSE 5% IN PLASTIC CONTAINER				
B BRAUN	5GM/100ML;30MG/100ML;37MG/100ML;370MG/100ML;530MG/100ML;500MG/100ML	N19843	001	Aug 09, 1993
NORMOSOL-R AND DEXTROSE 5% IN PLASTIC CONTAINER				
HOSPIRA	5GM/100ML;30MG/100ML;37MG/100ML;222MG/100ML;526MG/100ML;502MG/100ML	N17609	001	
PLASMA-LYTE 148 AND DEXTROSE 5% IN PLASTIC CONTAINER				
BAXTER HLTHCARE	5GM/100ML;30MG/100ML;37MG/100ML;368MG/100ML;526MG/100ML;502MG/100ML	N17451	001	

DEXTROSE; POTASSIUM CHLORIDE

INJECTABLE; INJECTION

<u>DEXTROSE 5% AND POTASSIUM CHLORIDE 0.075% IN PLASTIC CONTAINER</u>				
<u>AP</u>	BRISTOL MYERS SQUIBB <u>5GM/100ML;75MG/100ML</u>	<u>N17634</u>	<u>004</u>	
<u>DEXTROSE 5% AND POTASSIUM CHLORIDE 0.15% IN PLASTIC CONTAINER</u>				
<u>AP</u>	BRISTOL MYERS SQUIBB <u>5GM/100ML;150MG/100ML</u>	<u>N17634</u>	<u>001</u>	
<u>DEXTROSE 5% AND POTASSIUM CHLORIDE 0.224% IN PLASTIC CONTAINER</u>				
<u>AP</u>	BRISTOL MYERS SQUIBB <u>5GM/100ML;224MG/100ML</u>	<u>N17634</u>	<u>003</u>	
<u>DEXTROSE 5% AND POTASSIUM CHLORIDE 0.3% IN PLASTIC CONTAINER</u>				
<u>AP</u>	BRISTOL MYERS SQUIBB <u>5GM/100ML;300MG/100ML</u>	<u>N17634</u>	<u>002</u>	
<u>POTASSIUM CHLORIDE 0.075% IN DEXTROSE 5% IN PLASTIC CONTAINER</u>				
<u>AP</u>	B BRAUN <u>5GM/100ML;75MG/100ML</u>	<u>N18744</u>	<u>001</u>	Nov 09, 1982
<u>POTASSIUM CHLORIDE 0.15% IN DEXTROSE 5% IN PLASTIC CONTAINER</u>				
<u>AP</u>	B BRAUN <u>5GM/100ML;150MG/100ML</u>	<u>N18744</u>	<u>002</u>	Nov 09, 1982
<u>AP</u>	<u>5GM/100ML;150MG/100ML</u>	<u>N19699</u>	<u>004</u>	Sep 29, 1989
POTASSIUM CHLORIDE 0.22% IN DEXTROSE 5% IN PLASTIC CONTAINER				
B BRAUN	5GM/100ML;220MG/100ML	N18744	003	Nov 09, 1982
<u>POTASSIUM CHLORIDE 0.3% IN DEXTROSE 5% IN PLASTIC CONTAINER</u>				
<u>AP</u>	B BRAUN <u>5GM/100ML;300MG/100ML</u>	<u>N18744</u>	<u>004</u>	Nov 09, 1982
<u>AP</u>	<u>5GM/100ML;300MG/100ML</u>	<u>N19699</u>	<u>006</u>	Sep 29, 1989
POTASSIUM CHLORIDE 20MEQ IN DEXTROSE 5% IN PLASTIC CONTAINER				
HOSPIRA	5GM/100ML;149MG/100ML	N18371	001	
<u>POTASSIUM CHLORIDE 30MEQ IN DEXTROSE 5% IN PLASTIC CONTAINER</u>				
<u>AP</u>	HOSPIRA <u>5GM/100ML;224MG/100ML</u>	<u>N18371</u>	<u>003</u>	
POTASSIUM CHLORIDE 40MEQ IN DEXTROSE 5% IN PLASTIC CONTAINER				
HOSPIRA	5GM/100ML;298MG/100ML	N18371	002	

DEXTROSE; POTASSIUM CHLORIDE; POTASSIUM LACTATE; SODIUM CHLORIDE; SODIUM PHOSPHATE, MONOBASIC, ANHYDROUS

INJECTABLE; INJECTION

IONOSOL T AND DEXTROSE 5% IN PLASTIC CONTAINER				
HOSPIRA	5GM/100ML;111MG/100ML;256MG/100ML;146MG/100ML;207MG/100ML	N19514	001	May 08, 1986

DEXTROSE; POTASSIUM CHLORIDE; POTASSIUM PHOSPHATE, DIBASIC; SODIUM ACETATE; SODIUM CHLORIDE

INJECTABLE; INJECTION

ISOLYTE M IN DEXTROSE 5% IN PLASTIC CONTAINER				
B BRAUN	5GM/100ML;150MG/100ML;130MG/100ML;280MG/100ML;91MG/100ML	N19870	001	Jun 10, 1993

DEXTROSE; POTASSIUM CHLORIDE; POTASSIUM PHOSPHATE, MONOBASIC; SODIUM CHLORIDE; SODIUM LACTATE

INJECTABLE; INJECTION

DEXTROSE 5% AND ELECTROLYTE NO 75 IN PLASTIC CONTAINER				
BAXTER HLTHCARE	5GM/100ML;205MG/100ML;100MG/100ML;120MG/100ML;220MG/100ML	N18840	001	Jun 29, 1983

PRESCRIPTION DRUG PRODUCT LIST

3 - 117 (of 370)

DEXTROSE; POTASSIUM CHLORIDE; SODIUM CHLORIDE

INJECTABLE; INJECTION

DEXTROSE 5%, SODIUM CHLORIDE 0.2% AND POTASSIUM CHLORIDE 10MEQ					
AP	BAXTER HLTHCARE	5GM/100ML;75MG/100ML;200MG/100ML	N18037	006	Apr 13, 1982
AP		5GM/100ML;150MG/100ML;200MG/100ML	N18037	007	Apr 13, 1982
DEXTROSE 5%, SODIUM CHLORIDE 0.2% AND POTASSIUM CHLORIDE 15MEQ (K)					
AP	BAXTER HLTHCARE	5GM/100ML;224MG/100ML;200MG/100ML	N18037	004	
AP	BAXTER HLTHCARE	5GM/100ML;150MG/100ML;200MG/100ML	N18037	008	Apr 13, 1982
DEXTROSE 5%, SODIUM CHLORIDE 0.2% AND POTASSIUM CHLORIDE 20MEQ (K)					
AP	BAXTER HLTHCARE	5GM/100ML;300MG/100ML;200MG/100ML	N18037	001	
DEXTROSE 5%, SODIUM CHLORIDE 0.2% AND POTASSIUM CHLORIDE 30MEQ					
AP	BAXTER HLTHCARE	5GM/100ML;224MG/100ML;200MG/100ML	N18037	005	Apr 13, 1982
DEXTROSE 5%, SODIUM CHLORIDE 0.2% AND POTASSIUM CHLORIDE 40MEQ					
AP	BAXTER HLTHCARE	5GM/100ML;300MG/100ML;200MG/100ML	N18037	009	Apr 13, 1982
DEXTROSE 5%, SODIUM CHLORIDE 0.2% AND POTASSIUM CHLORIDE 5MEQ					
AP	BAXTER HLTHCARE	5GM/100ML;75MG/100ML;200MG/100ML	N18037	002	
DEXTROSE 5%, SODIUM CHLORIDE 0.2% AND POTASSIUM CHLORIDE 5MEQ (K)					
AP	BAXTER HLTHCARE	5GM/100ML;150MG/100ML;200MG/100ML	N18037	003	
DEXTROSE 5%, SODIUM CHLORIDE 0.33% AND POTASSIUM CHLORIDE 10MEQ IN PLASTIC CONTAINER					
AP	BAXTER HLTHCARE	5GM/100ML;75MG/100ML;330MG/100ML	N18629	005	Mar 23, 1982
AP		5GM/100ML;150MG/100ML;330MG/100ML	N18629	002	Mar 23, 1982
DEXTROSE 5%, SODIUM CHLORIDE 0.33% AND POTASSIUM CHLORIDE 15MEQ IN PLASTIC CONTAINER					
AP	BAXTER HLTHCARE	5GM/100ML;224MG/100ML;330MG/100ML	N18629	003	Mar 23, 1982
DEXTROSE 5%, SODIUM CHLORIDE 0.33% AND POTASSIUM CHLORIDE 20MEQ IN PLASTIC CONTAINER					
AP	BAXTER HLTHCARE	5GM/100ML;150MG/100ML;330MG/100ML	N18629	004	Mar 23, 1982
AP		5GM/100ML;300MG/100ML;330MG/100ML	N18629	006	Mar 23, 1982
DEXTROSE 5%, SODIUM CHLORIDE 0.33% AND POTASSIUM CHLORIDE 30MEQ IN PLASTIC CONTAINER					
AP	BAXTER HLTHCARE	5GM/100ML;224MG/100ML;330MG/100ML	N18629	007	Mar 23, 1982
DEXTROSE 5%, SODIUM CHLORIDE 0.33% AND POTASSIUM CHLORIDE 40MEQ IN PLASTIC CONTAINER					
AP	BAXTER HLTHCARE	5GM/100ML;300MG/100ML;330MG/100ML	N18629	008	Mar 23, 1982
DEXTROSE 5%, SODIUM CHLORIDE 0.33% AND POTASSIUM CHLORIDE 5MEQ IN PLASTIC CONTAINER					
AP	BAXTER HLTHCARE	5GM/100ML;75MG/100ML;330MG/100ML	N18629	001	Mar 23, 1982
DEXTROSE 5%, SODIUM CHLORIDE 0.45% AND POTASSIUM CHLORIDE 20MEQ (K) IN PLASTIC CONTAINER					
AP	BAXTER HLTHCARE	5GM/100ML;300MG/100ML;450MG/100ML	N18008	010	
POTASSIUM CHLORIDE 0.037% IN DEXTROSE 10% AND SODIUM CHLORIDE 0.2% IN PLASTIC CONTAINER					
	B BRAUN	10GM/100ML;37MG/100ML;200MG/100ML	N19630	031	Feb 17, 1988
POTASSIUM CHLORIDE 0.037% IN DEXTROSE 10% AND SODIUM CHLORIDE 0.45% IN PLASTIC CONTAINER					
	B BRAUN	10GM/100ML;37MG/100ML;450MG/100ML	N19630	037	Feb 17, 1988
POTASSIUM CHLORIDE 0.037% IN DEXTROSE 10% AND SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER					
	B BRAUN	10GM/100ML;37MG/100ML;900MG/100ML	N19630	043	Feb 17, 1988
POTASSIUM CHLORIDE 0.037% IN DEXTROSE 5% AND SODIUM CHLORIDE 0.11% IN PLASTIC CONTAINER					
	B BRAUN	5GM/100ML;37MG/100ML;110MG/100ML	N19630	001	Feb 17, 1988
POTASSIUM CHLORIDE 0.037% IN DEXTROSE 5% AND SODIUM CHLORIDE 0.2% IN PLASTIC CONTAINER					
	B BRAUN	5GM/100ML;37MG/100ML;200MG/100ML	N19630	007	Feb 17, 1988
POTASSIUM CHLORIDE 0.037% IN DEXTROSE 5% AND SODIUM CHLORIDE 0.33% IN PLASTIC CONTAINER					
	B BRAUN	5GM/100ML;37MG/100ML;330MG/100ML	N19630	013	Feb 17, 1988
POTASSIUM CHLORIDE 0.037% IN DEXTROSE 5% AND SODIUM CHLORIDE 0.45% IN PLASTIC CONTAINER					
	B BRAUN	5GM/100ML;37MG/100ML;450MG/100ML	N19630	019	Feb 17, 1988
POTASSIUM CHLORIDE 0.037% IN DEXTROSE 5% AND SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER					
	B BRAUN	5GM/100ML;37MG/100ML;900MG/100ML	N19630	025	Feb 17, 1988
POTASSIUM CHLORIDE 0.075% IN DEXTROSE 10% AND SODIUM CHLORIDE 0.2% IN PLASTIC CONTAINER					
	B BRAUN	10GM/100ML;75MG/100ML;200MG/100ML	N19630	032	Feb 17, 1988
POTASSIUM CHLORIDE 0.075% IN DEXTROSE 10% AND SODIUM CHLORIDE 0.45% IN PLASTIC CONTAINER					
	B BRAUN	10GM/100ML;75MG/100ML;450MG/100ML	N19630	038	Feb 17, 1988
POTASSIUM CHLORIDE 0.075% IN DEXTROSE 10% AND SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER					
	B BRAUN	10GM/100ML;75MG/100ML;900MG/100ML	N19630	044	Feb 17, 1988
POTASSIUM CHLORIDE 0.075% IN DEXTROSE 3.3% AND SODIUM CHLORIDE 0.3% IN PLASTIC CONTAINER					
	B BRAUN	3.3GM/100ML;75MG/100ML;300MG/100ML	N19630	049	May 07, 1992
POTASSIUM CHLORIDE 0.075% IN DEXTROSE 5% AND SODIUM CHLORIDE 0.11% IN PLASTIC CONTAINER					
	B BRAUN	5GM/100ML;75MG/100ML;110MG/100ML	N19630	002	Feb 17, 1988

PRESCRIPTION DRUG PRODUCT LIST

3 - 118 (of 370)

DEXTROSE; POTASSIUM CHLORIDE; SODIUM CHLORIDE

INJECTABLE; INJECTION

	POTASSIUM CHLORIDE 0.075% IN DEXTROSE 5% AND SODIUM CHLORIDE 0.2% IN PLASTIC CONTAINER		
AP	B BRAUN	5GM/100ML;75MG/100ML;200MG/100ML	N19630 008 Feb 17, 1988
	POTASSIUM CHLORIDE 0.075% IN DEXTROSE 5% AND SODIUM CHLORIDE 0.33% IN PLASTIC CONTAINER		
AP	B BRAUN	5GM/100ML;75MG/100ML;330MG/100ML	N19630 014 Feb 17, 1988
	POTASSIUM CHLORIDE 0.075% IN DEXTROSE 5% AND SODIUM CHLORIDE 0.45% IN PLASTIC CONTAINER		
AP	B BRAUN	5GM/100ML;75MG/100ML;450MG/100ML	N19630 020 Feb 17, 1988
	POTASSIUM CHLORIDE 0.075% IN DEXTROSE 5% AND SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER		
AP	B BRAUN	5GM/100ML;75MG/100ML;900MG/100ML	N19630 026 Feb 17, 1988
	POTASSIUM CHLORIDE 0.11% IN DEXTROSE 10% AND SODIUM CHLORIDE 0.2% IN PLASTIC CONTAINER		
	B BRAUN	10GM/100ML;110MG/100ML;200MG/100ML	N19630 033 Feb 17, 1988
	POTASSIUM CHLORIDE 0.11% IN DEXTROSE 10% AND SODIUM CHLORIDE 0.45% IN PLASTIC CONTAINER		
	B BRAUN	10GM/100ML;110MG/100ML;450MG/100ML	N19630 039 Feb 17, 1988
	POTASSIUM CHLORIDE 0.11% IN DEXTROSE 10% AND SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER		
	B BRAUN	10GM/100ML;110MG/100ML;900MG/100ML	N19630 045 Feb 17, 1988
	POTASSIUM CHLORIDE 0.11% IN DEXTROSE 3.3% AND SODIUM CHLORIDE 0.3% IN PLASTIC CONTAINER		
	B BRAUN	3.3GM/100ML;110MG/100ML;300MG/100ML	N19630 050 May 07, 1992
	POTASSIUM CHLORIDE 0.11% IN DEXTROSE 5% AND SODIUM CHLORIDE 0.11% IN PLASTIC CONTAINER		
	B BRAUN	5GM/100ML;110MG/100ML;110MG/100ML	N19630 003 Feb 17, 1988
	POTASSIUM CHLORIDE 0.11% IN DEXTROSE 5% AND SODIUM CHLORIDE 0.2% IN PLASTIC CONTAINER		
	B BRAUN	5GM/100ML;110MG/100ML;200MG/100ML	N19630 009 Feb 17, 1988
	POTASSIUM CHLORIDE 0.11% IN DEXTROSE 5% AND SODIUM CHLORIDE 0.33% IN PLASTIC CONTAINER		
	B BRAUN	5GM/100ML;110MG/100ML;330MG/100ML	N19630 015 Feb 17, 1988
	POTASSIUM CHLORIDE 0.11% IN DEXTROSE 5% AND SODIUM CHLORIDE 0.45% IN PLASTIC CONTAINER		
	B BRAUN	5GM/100ML;110MG/100ML;450MG/100ML	N19630 021 Feb 17, 1988
	POTASSIUM CHLORIDE 0.11% IN DEXTROSE 5% AND SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER		
	B BRAUN	5GM/100ML;110MG/100ML;900MG/100ML	N19630 027 Feb 17, 1988
	POTASSIUM CHLORIDE 0.15% IN DEXTROSE 10% AND SODIUM CHLORIDE 0.2% IN PLASTIC CONTAINER		
	B BRAUN	10GM/100ML;150MG/100ML;200MG/100ML	N19630 034 Feb 17, 1988
	POTASSIUM CHLORIDE 0.15% IN DEXTROSE 10% AND SODIUM CHLORIDE 0.45% IN PLASTIC CONTAINER		
	B BRAUN	10GM/100ML;150MG/100ML;450MG/100ML	N19630 040 Feb 17, 1988
	POTASSIUM CHLORIDE 0.15% IN DEXTROSE 10% AND SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER		
	B BRAUN	10GM/100ML;150MG/100ML;900MG/100ML	N19630 046 Feb 17, 1988
	POTASSIUM CHLORIDE 0.15% IN DEXTROSE 3.3% AND SODIUM CHLORIDE 0.3% IN PLASTIC CONTAINER		
	B BRAUN	3.3GM/100ML;150MG/100ML;300MG/100ML	N19630 051 May 07, 1992
	POTASSIUM CHLORIDE 0.15% IN DEXTROSE 5% AND SODIUM CHLORIDE 0.11% IN PLASTIC CONTAINER		
	B BRAUN	5GM/100ML;150MG/100ML;110MG/100ML	N19630 004 Feb 17, 1988
	POTASSIUM CHLORIDE 0.15% IN DEXTROSE 5% AND SODIUM CHLORIDE 0.2% IN PLASTIC CONTAINER		
AP	B BRAUN	5GM/100ML;150MG/100ML;200MG/100ML	N19630 010 Feb 17, 1988
	POTASSIUM CHLORIDE 0.15% IN DEXTROSE 5% AND SODIUM CHLORIDE 0.33% IN PLASTIC CONTAINER		
AP	B BRAUN	5GM/100ML;150MG/100ML;330MG/100ML	N19630 016 Feb 17, 1988
	POTASSIUM CHLORIDE 0.15% IN DEXTROSE 5% AND SODIUM CHLORIDE 0.45% IN PLASTIC CONTAINER		
AP	B BRAUN	5GM/100ML;150MG/100ML;450MG/100ML	N19630 022 Feb 17, 1988
	POTASSIUM CHLORIDE 0.15% IN DEXTROSE 5% AND SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER		
AP	B BRAUN	5GM/100ML;150MG/100ML;900MG/100ML	N19630 028 Feb 17, 1988
	POTASSIUM CHLORIDE 0.22% IN DEXTROSE 10% AND SODIUM CHLORIDE 0.2% IN PLASTIC CONTAINER		
	B BRAUN	10GM/100ML;220MG/100ML;200MG/100ML	N19630 035 Feb 17, 1988
	POTASSIUM CHLORIDE 0.22% IN DEXTROSE 10% AND SODIUM CHLORIDE 0.45% IN PLASTIC CONTAINER		
	B BRAUN	10GM/100ML;220MG/100ML;450MG/100ML	N19630 041 Feb 17, 1988
	POTASSIUM CHLORIDE 0.22% IN DEXTROSE 10% AND SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER		
	B BRAUN	10GM/100ML;220MG/100ML;900MG/100ML	N19630 047 Feb 17, 1988
	POTASSIUM CHLORIDE 0.22% IN DEXTROSE 3.3% AND SODIUM CHLORIDE 0.3% IN PLASTIC CONTAINER		
	B BRAUN	3.3GM/100ML;220MG/100ML;300MG/100ML	N19630 052 May 07, 1992
	POTASSIUM CHLORIDE 0.22% IN DEXTROSE 5% AND SODIUM CHLORIDE 0.11% IN PLASTIC CONTAINER		
	B BRAUN	5GM/100ML;220MG/100ML;110MG/100ML	N19630 005 Feb 17, 1988
	POTASSIUM CHLORIDE 0.22% IN DEXTROSE 5% AND SODIUM CHLORIDE 0.2% IN PLASTIC CONTAINER		
	B BRAUN	5GM/100ML;220MG/100ML;200MG/100ML	N19630 011 Feb 17, 1988
	POTASSIUM CHLORIDE 0.22% IN DEXTROSE 5% AND SODIUM CHLORIDE 0.33% IN PLASTIC CONTAINER		
	B BRAUN	5GM/100ML;220MG/100ML;330MG/100ML	N19630 017 Feb 17, 1988
	POTASSIUM CHLORIDE 0.22% IN DEXTROSE 5% AND SODIUM CHLORIDE 0.45% IN PLASTIC CONTAINER		
	B BRAUN	5GM/100ML;220MG/100ML;450MG/100ML	N19630 023 Feb 17, 1988

PRESCRIPTION DRUG PRODUCT LIST

3 - 119 (of 370)

DEXTROSE; POTASSIUM CHLORIDE; SODIUM CHLORIDE

INJECTABLE; INJECTION			
	POTASSIUM CHLORIDE 0.22% IN DEXTROSE 5% AND SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER		
	B BRAUN 5GM/100ML;220MG/100ML;900MG/100ML	N19630 029	Feb 17, 1988
	POTASSIUM CHLORIDE 0.3% IN DEXTROSE 10% AND SODIUM CHLORIDE 0.2% IN PLASTIC CONTAINER		
	B BRAUN 10GM/100ML;300MG/100ML;200MG/100ML	N19630 036	Feb 17, 1988
	POTASSIUM CHLORIDE 0.3% IN DEXTROSE 10% AND SODIUM CHLORIDE 0.45% IN PLASTIC CONTAINER		
	B BRAUN 10GM/100ML;300MG/100ML;450MG/100ML	N19630 042	Feb 17, 1988
	POTASSIUM CHLORIDE 0.3% IN DEXTROSE 10% AND SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER		
	B BRAUN 10GM/100ML;300MG/100ML;900MG/100ML	N19630 048	Feb 17, 1988
	POTASSIUM CHLORIDE 0.3% IN DEXTROSE 3.3% AND SODIUM CHLORIDE 0.3% IN PLASTIC CONTAINER		
	B BRAUN 3.3GM/100ML;300MG/100ML;300MG/100ML	N19630 053	May 07, 1992
	POTASSIUM CHLORIDE 0.3% IN DEXTROSE 5% AND SODIUM CHLORIDE 0.11% IN PLASTIC CONTAINER		
	B BRAUN 5GM/100ML;300MG/100ML;110MG/100ML	N19630 006	Feb 17, 1988
	POTASSIUM CHLORIDE 0.3% IN DEXTROSE 5% AND SODIUM CHLORIDE 0.2% IN PLASTIC CONTAINER		
AP	B BRAUN 5GM/100ML;300MG/100ML;200MG/100ML	N19630 012	Feb 17, 1988
	POTASSIUM CHLORIDE 0.3% IN DEXTROSE 5% AND SODIUM CHLORIDE 0.33% IN PLASTIC CONTAINER		
AP	B BRAUN 5GM/100ML;300MG/100ML;330MG/100ML	N19630 018	Feb 17, 1988
	POTASSIUM CHLORIDE 0.3% IN DEXTROSE 5% AND SODIUM CHLORIDE 0.45% IN PLASTIC CONTAINER		
AP	B BRAUN 5GM/100ML;300MG/100ML;450MG/100ML	N19630 024	Feb 17, 1988
	POTASSIUM CHLORIDE 0.3% IN DEXTROSE 5% AND SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER		
AP	B BRAUN 5GM/100ML;300MG/100ML;900MG/100ML	N19630 030	Feb 17, 1988
	POTASSIUM CHLORIDE 10MEQ IN DEXTROSE 5% AND SODIUM CHLORIDE 0.225% IN PLASTIC CONTAINER		
	HOSPIRA 5GM/100ML;74.5MG/100ML;225MG/100ML	N18365 002	Jul 05, 1983
	5GM/100ML;149MG/100ML;225MG/100ML	N18365 006	Mar 28, 1988
	POTASSIUM CHLORIDE 10MEQ IN DEXTROSE 5% AND SODIUM CHLORIDE 0.3% IN PLASTIC CONTAINER		
	HOSPIRA 5GM/100ML;74.5MG/100ML;300MG/100ML	N18876 001	Jan 17, 1986
	5GM/100ML;149MG/100ML;300MG/100ML	N18876 006	Mar 28, 1988
	POTASSIUM CHLORIDE 10MEQ IN DEXTROSE 5% AND SODIUM CHLORIDE 0.45% IN PLASTIC CONTAINER		
AP	BAXTER HLTHCARE 5GM/100ML;75MG/100ML;450MG/100ML	N18008 005	Apr 28, 1982
AP	5GM/100ML;150MG/100ML;450MG/100ML	N18008 006	Apr 28, 1982
AP	HOSPIRA 5GM/100ML;74.5MG/100ML;450MG/100ML	N18362 009	Jul 05, 1983
AP	5GM/100ML;74.5MG/100ML;450MG/100ML	N18362 005	Mar 28, 1988
	POTASSIUM CHLORIDE 10MEQ IN DEXTROSE 5% AND SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER		
AP	BAXTER HLTHCARE 5GM/100ML;75MG/100ML;900MG/100ML	N19308 004	Apr 05, 1985
AP	5GM/100ML;150MG/100ML;900MG/100ML	N19308 002	Apr 05, 1985
AP	HOSPIRA 5GM/100ML;74.5MG/100ML;900MG/100ML	N19691 002	Mar 24, 1988
AP	5GM/100ML;149MG/100ML;900MG/100ML	N19691 004	Mar 24, 1988
	POTASSIUM CHLORIDE 15MEQ IN DEXTROSE 5% AND SODIUM CHLORIDE 0.225% IN PLASTIC CONTAINER		
	HOSPIRA 5GM/100ML;224MG/100ML;225MG/100ML	N18365 008	Mar 28, 1988
	POTASSIUM CHLORIDE 15MEQ IN DEXTROSE 5% AND SODIUM CHLORIDE 0.3% IN PLASTIC CONTAINER		
	HOSPIRA 5GM/100ML;224MG/100ML;300MG/100ML	N18876 007	Mar 28, 1988
	POTASSIUM CHLORIDE 15MEQ IN DEXTROSE 5% AND SODIUM CHLORIDE 0.45% IN PLASTIC CONTAINER		
AP	HOSPIRA 5GM/100ML;224MG/100ML;450MG/100ML	N18362 006	Mar 28, 1988
	POTASSIUM CHLORIDE 15MEQ IN DEXTROSE 5% AND SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER		
AP	HOSPIRA 5GM/100ML;224MG/100ML;900MG/100ML	N19691 006	Mar 24, 1988
	POTASSIUM CHLORIDE 20MEQ IN DEXTROSE 5% AND SODIUM CHLORIDE 0.225% IN PLASTIC CONTAINER		
	HOSPIRA 5GM/100ML;149MG/100ML;225MG/100ML	N18365 001	
	5GM/100ML;298MG/100ML;225MG/100ML	N18365 009	Mar 28, 1988
	POTASSIUM CHLORIDE 20MEQ IN DEXTROSE 5% AND SODIUM CHLORIDE 0.3% IN PLASTIC CONTAINER		
	HOSPIRA 5GM/100ML;298MG/100ML;300MG/100ML	N18876 008	Mar 28, 1988
	POTASSIUM CHLORIDE 20MEQ IN DEXTROSE 5% AND SODIUM CHLORIDE 0.45% IN PLASTIC CONTAINER		
AP	BAXTER HLTHCARE 5GM/100ML;150MG/100ML;450MG/100ML	N18008 007	Apr 28, 1982
AP	HOSPIRA 5GM/100ML;149MG/100ML;450MG/100ML	N18362 010	Jul 05, 1983
AP	5GM/100ML;298MG/100ML;450MG/100ML	N18362 007	Mar 28, 1988
	POTASSIUM CHLORIDE 20MEQ IN DEXTROSE 5% AND SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER		
AP	BAXTER HLTHCARE 5GM/100ML;150MG/100ML;900MG/100ML	N19308 005	Apr 05, 1985
AP	5GM/100ML;300MG/100ML;900MG/100ML	N19308 003	Apr 05, 1985
AP	HOSPIRA 5GM/100ML;149MG/100ML;900MG/100ML	N19691 005	Mar 24, 1988
AP	5GM/100ML;298MG/100ML;900MG/100ML	N19691 008	Mar 24, 1988

PRESCRIPTION DRUG PRODUCT LIST

3 - 120 (of 370)

DEXTROSE; POTASSIUM CHLORIDE; SODIUM CHLORIDE

INJECTABLE; INJECTION			
POTASSIUM CHLORIDE 20MEQ IN DEXTROSE 5% IN SODIUM CHLORIDE 0.3% IN PLASTIC CONTAINER			
	HOSPIRA	5GM/100ML;149MG/100ML;300MG/100ML	N18876 002 Jan 17, 1986
POTASSIUM CHLORIDE 30MEQ IN DEXTROSE 5% AND SODIUM CHLORIDE 0.225% IN PLASTIC CONTAINER			
	HOSPIRA	5GM/100ML;224MG/100ML;225MG/100ML	N18365 003 Jul 05, 1983
POTASSIUM CHLORIDE 30MEQ IN DEXTROSE 5% AND SODIUM CHLORIDE 0.3% IN PLASTIC CONTAINER			
	HOSPIRA	5GM/100ML;224MG/100ML;300MG/100ML	N18876 003 Jan 17, 1986
POTASSIUM CHLORIDE 30MEQ IN DEXTROSE 5% AND SODIUM CHLORIDE 0.45% IN PLASTIC CONTAINER			
<u>AP</u>	BAXTER HLTHCARE	<u>5GM/100ML;224MG/100ML;450MG/100ML</u>	<u>N18008 008</u> Apr 28, 1982
<u>AP</u>	HOSPIRA	<u>5GM/100ML;224MG/100ML;450MG/100ML</u>	<u>N18362 002</u>
POTASSIUM CHLORIDE 30MEQ IN DEXTROSE 5% AND SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER			
<u>AP</u>	BAXTER HLTHCARE	<u>5GM/100ML;224MG/100ML;900MG/100ML</u>	<u>N19308 006</u> Apr 05, 1985
<u>AP</u>	HOSPIRA	<u>5GM/100ML;224MG/100ML;900MG/100ML</u>	<u>N19691 007</u> Mar 24, 1988
POTASSIUM CHLORIDE 40MEQ IN DEXTROSE 5% AND SODIUM CHLORIDE 0.225% IN PLASTIC CONTAINER			
	HOSPIRA	5GM/100ML;298MG/100ML;225MG/100ML	N18365 004 Jul 05, 1983
POTASSIUM CHLORIDE 40MEQ IN DEXTROSE 5% AND SODIUM CHLORIDE 0.3% IN PLASTIC CONTAINER			
	HOSPIRA	5GM/100ML;298MG/100ML;300MG/100ML	N18876 004 Mar 28, 1988
POTASSIUM CHLORIDE 40MEQ IN DEXTROSE 5% AND SODIUM CHLORIDE 0.45% IN PLASTIC CONTAINER			
<u>AP</u>	BAXTER HLTHCARE	<u>5GM/100ML;300MG/100ML;450MG/100ML</u>	<u>N18008 009</u> Apr 28, 1982
<u>AP</u>	HOSPIRA	<u>5GM/100ML;298MG/100ML;450MG/100ML</u>	<u>N18362 003</u>
POTASSIUM CHLORIDE 40MEQ IN DEXTROSE 5% AND SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER			
<u>AP</u>	BAXTER HLTHCARE	<u>5GM/100ML;300MG/100ML;900MG/100ML</u>	<u>N19308 007</u> Apr 05, 1985
<u>AP</u>	HOSPIRA	<u>5GM/100ML;298MG/100ML;900MG/100ML</u>	<u>N19691 009</u> Mar 24, 1988
POTASSIUM CHLORIDE 5MEQ IN DEXTROSE 5% AND SODIUM CHLORIDE 0.225% IN PLASTIC CONTAINER			
	HOSPIRA	5GM/100ML;74.5MG/100ML;225MG/100ML	N18365 005 Mar 28, 1988
		5GM/100ML;149MG/100ML;225MG/100ML	N18365 007 Mar 28, 1988
POTASSIUM CHLORIDE 5MEQ IN DEXTROSE 5% AND SODIUM CHLORIDE 0.3% IN PLASTIC CONTAINER			
	HOSPIRA	5GM/100ML;74.5MG/100ML;300MG/100ML	N18876 005 Mar 28, 1988
		5GM/100ML;149MG/100ML;300MG/100ML	N18876 009 Mar 28, 1988
POTASSIUM CHLORIDE 5MEQ IN DEXTROSE 5% AND SODIUM CHLORIDE 0.45% IN PLASTIC CONTAINER			
<u>AP</u>	BAXTER HLTHCARE	<u>5GM/100ML;150MG/100ML;450MG/100ML</u>	<u>N18008 004</u>
<u>AP</u>	HOSPIRA	<u>5GM/100ML;74.5MG/100ML;450MG/100ML</u>	<u>N18362 008</u> Mar 28, 1988
<u>AP</u>		<u>5GM/100ML;149MG/100ML;450MG/100ML</u>	<u>N18362 004</u> Mar 28, 1988
POTASSIUM CHLORIDE 5MEQ IN DEXTROSE 5% AND SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER			
<u>AP</u>	BAXTER HLTHCARE	<u>5GM/100ML;150MG/100ML;900MG/100ML</u>	<u>N19308 001</u> Apr 05, 1985
<u>AP</u>	HOSPIRA	<u>5GM/100ML;74.5MG/100ML;900MG/100ML</u>	<u>N19691 001</u> Mar 24, 1988
<u>AP</u>		<u>5GM/100ML;149MG/100ML;900MG/100ML</u>	<u>N19691 003</u> Mar 24, 1988

DEXTROSE; SODIUM CHLORIDE

INJECTABLE; INJECTION			
DEXTROSE 10% AND SODIUM CHLORIDE 0.11% IN PLASTIC CONTAINER			
	B BRAUN	10GM/100ML;110MG/100ML	N19631 011 Feb 24, 1988
DEXTROSE 10% AND SODIUM CHLORIDE 0.2% IN PLASTIC CONTAINER			
	B BRAUN	10GM/100ML;200MG/100ML	N19631 012 Feb 24, 1988
DEXTROSE 10% AND SODIUM CHLORIDE 0.33% IN PLASTIC CONTAINER			
	B BRAUN	10GM/100ML;330MG/100ML	N19631 013 Feb 24, 1988
DEXTROSE 10% AND SODIUM CHLORIDE 0.45% IN PLASTIC CONTAINER			
	B BRAUN	10GM/100ML;450MG/100ML	N19631 014 Feb 24, 1988
DEXTROSE 10% AND SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER			
<u>AP</u>	B BRAUN	<u>10GM/100ML;900MG/100ML</u>	<u>N19631 015</u> Feb 24, 1988
<u>AP</u>	BAXTER HLTHCARE	<u>10GM/100ML;900MG/100ML</u>	<u>N16696 001</u>
DEXTROSE 2.5% AND SODIUM CHLORIDE 0.11% IN PLASTIC CONTAINER			
	B BRAUN	2.5GM/100ML;110MG/100ML	N19631 001 Feb 24, 1988
DEXTROSE 2.5% AND SODIUM CHLORIDE 0.2% IN PLASTIC CONTAINER			
	B BRAUN	2.5GM/100ML;200MG/100ML	N19631 002 Feb 24, 1988
DEXTROSE 2.5% AND SODIUM CHLORIDE 0.33% IN PLASTIC CONTAINER			
	B BRAUN	2.5GM/100ML;330MG/100ML	N19631 003 Feb 24, 1988
DEXTROSE 2.5% AND SODIUM CHLORIDE 0.45% IN PLASTIC CONTAINER			
<u>AP</u>	B BRAUN	<u>2.5GM/100ML;450MG/100ML</u>	<u>N19631 004</u> Feb 24, 1988
<u>AP</u>	BAXTER HLTHCARE	<u>2.5GM/100ML;450MG/100ML</u>	<u>N16697 001</u>

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 121 (of 370)

DEXTROSE; SODIUM CHLORIDE

INJECTABLE; INJECTION

<u>DEXTROSE 2.5% AND SODIUM CHLORIDE 0.45% IN PLASTIC CONTAINER</u>				
<u>AP</u>	HOSPIRA	<u>2.5GM/100ML;450MG/100ML</u>	<u>N18096</u>	<u>001</u>
	DEXTROSE 2.5% AND SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER			
	B BRAUN	2.5GM/100ML;900MG/100ML	N19631	005 Feb 24, 1988
	DEXTROSE 3.3% AND SODIUM CHLORIDE 0.3% IN PLASTIC CONTAINER			
	B BRAUN	3.3GM/100ML;300MG/100ML	N19631	016 Jan 19, 1990
	DEXTROSE 5% AND SODIUM CHLORIDE 0.11% IN PLASTIC CONTAINER			
	B BRAUN	5GM/100ML;110MG/100ML	N19631	006 Feb 24, 1988
<u>DEXTROSE 5% AND SODIUM CHLORIDE 0.2% IN PLASTIC CONTAINER</u>				
<u>AP</u>	B BRAUN	<u>5GM/100ML;200MG/100ML</u>	<u>N19631</u>	<u>007</u> Feb 24, 1988
	DEXTROSE 5% AND SODIUM CHLORIDE 0.225% IN PLASTIC CONTAINER			
	HOSPIRA	5GM/100ML;225MG/100ML	N17606	001
	DEXTROSE 5% AND SODIUM CHLORIDE 0.3% IN PLASTIC CONTAINER			
	HOSPIRA	5GM/100ML;300MG/100ML	N17799	001
<u>DEXTROSE 5% AND SODIUM CHLORIDE 0.33% IN PLASTIC CONTAINER</u>				
<u>AP</u>	B BRAUN	<u>5GM/100ML;330MG/100ML</u>	<u>N19631</u>	<u>008</u> Feb 24, 1988
<u>DEXTROSE 5% AND SODIUM CHLORIDE 0.45% IN PLASTIC CONTAINER</u>				
<u>AP</u>	B BRAUN	<u>5GM/100ML;450MG/100ML</u>	<u>N19631</u>	<u>009</u> Feb 24, 1988
<u>AP</u>	HOSPIRA	<u>5GM/100ML;450MG/100ML</u>	<u>N17607</u>	<u>001</u>
<u>DEXTROSE 5% AND SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER</u>				
<u>AP</u>	B BRAUN	<u>5GM/100ML;900MG/100ML</u>	<u>N19631</u>	<u>010</u> Feb 24, 1988
<u>AP</u>	HOSPIRA	<u>5GM/100ML;900MG/100ML</u>	<u>N17585</u>	<u>001</u>
<u>DEXTROSE 5% IN SODIUM CHLORIDE 0.2% IN PLASTIC CONTAINER</u>				
<u>AP</u>	BAXTER HLTHCARE	<u>5GM/100ML;200MG/100ML</u>	<u>N16689</u>	<u>001</u>
<u>DEXTROSE 5% IN SODIUM CHLORIDE 0.33% IN PLASTIC CONTAINER</u>				
<u>AP</u>	BAXTER HLTHCARE	<u>5GM/100ML;330MG/100ML</u>	<u>N16687</u>	<u>001</u>
<u>DEXTROSE 5% IN SODIUM CHLORIDE 0.45% IN PLASTIC CONTAINER</u>				
<u>AP</u>	BAXTER HLTHCARE	<u>5GM/100ML;450MG/100ML</u>	<u>N16683</u>	<u>001</u>
<u>DEXTROSE 5% IN SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER</u>				
<u>AP</u>	BAXTER HLTHCARE	<u>5GM/100ML;900MG/100ML</u>	<u>N16678</u>	<u>001</u>

DIATRIZOATE MEGLUMINE

INJECTABLE; INJECTION

<u>HYPaque</u>				
<u>AP</u>	+ GE HEALTHCARE	<u>60%</u>	<u>N16403</u>	<u>001</u>
<u>RENO-60</u>				
<u>AP</u>	+ BRACCO	<u>60%</u>	<u>N10040</u>	<u>016</u>
	RENO-DIP			
	+ BRACCO	30%	N10040	012
SOLUTION; URETERAL				
	RENO-30			
	BRACCO	30%	N10040	021
SOLUTION; URETHRAL				
<u>CYSTOGRAFIN</u>				
<u>AT</u>	BRACCO	<u>30%</u>	<u>N10040</u>	<u>018</u>
	CYSTOGRAFIN DILUTE			
	BRACCO	18%	N10040	022 Nov 09, 1982
<u>HYPaque-CYSTO</u>				
<u>AT</u>	GE HEALTHCARE	<u>30%</u>	<u>N16403</u>	<u>003</u>

DIATRIZOATE MEGLUMINE; DIATRIZOATE SODIUM

INJECTABLE; INJECTION

<u>HYPaque-76</u>				
<u>AP</u>	GE HEALTHCARE	<u>66%;10%</u>	<u>N86505</u>	<u>001</u>
<u>MD-76R</u>				
<u>AP</u>	+ MALLINCKRODT	<u>66%;10%</u>	<u>N19292</u>	<u>001</u> Sep 29, 1989
<u>RENOCAL-76</u>				
<u>AP</u>	BRACCO	<u>66%;10%</u>	<u>N89347</u>	<u>001</u> Jun 01, 1988

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 122 (of 370)

DIATRIZOATE MEGLUMINE; DIATRIZOATE SODIUM

INJECTABLE; INJECTION			
RENOGRAFIN-60			
	+ BRACCO	52%;8%	N10040 006
<u>RENOGRAFIN-76</u>			
<u>AP</u>	+ BRACCO	<u>66%;10%</u>	<u>N10040 001</u>
SOLUTION; ORAL, RECTAL			
<u>GASTROGRAFIN</u>			
<u>AA</u>	+ BRACCO	<u>66%;10%</u>	<u>N11245 003</u>
<u>MD-GASTROVIEW</u>			
<u>AA</u>	MALLINCKRODT	<u>66%;10%</u>	<u>N87388 001</u>

DIATRIZOATE MEGLUMINE; IODIPAMIDE MEGLUMINE

SOLUTION; INTRAUTERINE			
SINOGRAPHIN			
	+ BRACCO	52.7%;26.8%	N11324 002

DIATRIZOATE SODIUM

FOR SOLUTION; ORAL, RECTAL			
HYPAQUE			
	GE HEALTHCARE	100%	N11386 001
INJECTABLE; INJECTION			
HYPAQUE			
	+ GE HEALTHCARE	50%	N09561 001

DIAZEPAM

CONCENTRATE; ORAL			
DIAZEPAM INTENSOL			
	+ ROXANE	5MG/ML	N71415 001 Apr 03, 1987
GEL; RECTAL			
DIASTAT			
	VALEANT	2.5MG/0.5ML (5MG/ML)	N20648 001 Jul 29, 1997
DIASTAT ACUDIAL			
	VALEANT	10MG/2ML (5MG/ML)	N20648 007 Sep 15, 2005
	+	20MG/4ML (5MG/ML)	N20648 006 Sep 15, 2005
INJECTABLE; INJECTION			
<u>DIAZEPAM</u>			
<u>AP</u>	BAXTER HLTHCARE	<u>5MG/ML</u>	<u>N71309 001</u> Jul 17, 1987
<u>AP</u>		<u>5MG/ML</u>	<u>N71310 001</u> Jul 17, 1987
<u>AP</u>	+ HOSPIRA	<u>5MG/ML</u>	<u>N71583 001</u> Oct 13, 1987
<u>AP</u>		<u>5MG/ML</u>	<u>N71584 001</u> Oct 13, 1987
<u>AP</u>		<u>5MG/ML</u>	<u>N72079 001</u> Dec 20, 1988
<u>AP</u>	MARSAM PHARMS LLC	<u>5MG/ML</u>	<u>N72370 001</u> Jan 29, 1993
<u>AP</u>		<u>5MG/ML</u>	<u>N72397 001</u> Jan 29, 1993
<u>AP</u>	PARENTA PHARMS	<u>5MG/ML</u>	<u>N76815 001</u> Apr 15, 2004
<u>AP</u>	WATSON LABS	<u>5MG/ML</u>	<u>N70296 001</u> Feb 12, 1986
SOLUTION; ORAL			
DIAZEPAM			
	+ ROXANE	5MG/5ML	N70928 001 Apr 03, 1987
TABLET; ORAL			
<u>DIAZEPAM</u>			
<u>AB</u>	ACTAVIS ELIZABETH	<u>2MG</u>	<u>N70781 001</u> Mar 19, 1986
<u>AB</u>		<u>5MG</u>	<u>N70706 001</u> Mar 19, 1986
<u>AB</u>		<u>10MG</u>	<u>N70707 001</u> Mar 19, 1986
<u>AB</u>	BARR	<u>2MG</u>	<u>N70152 001</u> Nov 01, 1985
<u>AB</u>		<u>5MG</u>	<u>N70153 001</u> Nov 01, 1985
<u>AB</u>		<u>10MG</u>	<u>N70154 001</u> Nov 01, 1985
<u>AB</u>	CLONMEL HLTHCARE	<u>2MG</u>	<u>N70226 001</u> Sep 26, 1985
<u>AB</u>	IVAX PHARMS	<u>2MG</u>	<u>N71307 001</u> Dec 10, 1986

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 123 (of 370)

DIAZEPAM

TABLET; ORAL

DIAZEPAM

<u>AB</u>	IVAX PHARMS	<u>5MG</u>	<u>N71321</u>	<u>001</u>	Dec 10, 1986
<u>AB</u>		<u>10MG</u>	<u>N71322</u>	<u>001</u>	Dec 10, 1986
<u>AB</u>	MYLAN	<u>2MG</u>	<u>N70323</u>	<u>001</u>	Sep 04, 1985
<u>AB</u>		<u>5MG</u>	<u>N70324</u>	<u>001</u>	Sep 04, 1985
<u>AB</u>		<u>10MG</u>	<u>N70325</u>	<u>001</u>	Sep 04, 1985
<u>AB</u>	PAR PHARM	<u>2MG</u>	<u>N70462</u>	<u>001</u>	Feb 25, 1986
<u>AB</u>		<u>5MG</u>	<u>N70463</u>	<u>001</u>	Feb 25, 1986
<u>AB</u>	SANDOZ	<u>2MG</u>	<u>N70302</u>	<u>001</u>	Dec 20, 1985
<u>AB</u>		<u>5MG</u>	<u>N70303</u>	<u>001</u>	Dec 20, 1985
<u>AB</u>		<u>10MG</u>	<u>N70304</u>	<u>001</u>	Dec 20, 1985
<u>AB</u>	VINTAGE PHARMS	<u>2MG</u>	<u>N77749</u>	<u>001</u>	Mar 31, 2006
<u>AB</u>		<u>5MG</u>	<u>N77749</u>	<u>002</u>	Mar 31, 2006
<u>AB</u>		<u>10MG</u>	<u>N77749</u>	<u>003</u>	Mar 31, 2006
<u>AB</u>	WATSON LABS	<u>2MG</u>	<u>N71134</u>	<u>001</u>	Feb 03, 1987
<u>AB</u>		<u>5MG</u>	<u>N71135</u>	<u>001</u>	Feb 03, 1987
<u>AB</u>		<u>10MG</u>	<u>N71136</u>	<u>001</u>	Feb 03, 1987
<u>VALIUM</u>					
<u>AB</u>	ROCHE	<u>2MG</u>	<u>N13263</u>	<u>002</u>	
<u>AB</u>		<u>5MG</u>	<u>N13263</u>	<u>004</u>	
<u>AB</u>	+	<u>10MG</u>	<u>N13263</u>	<u>006</u>	

DIAZOXIDE

SUSPENSION; ORAL

PROGLYCEM

+ BAKER NORTON 50MG/ML N17453 001

DICLOFENAC POTASSIUM

TABLET; ORAL

CATAFLAM

<u>AB</u>	+	NOVARTIS	<u>50MG</u>	<u>N20142</u>	<u>002</u>	Nov 24, 1993
<u>DICLOFENAC POTASSIUM</u>						
<u>AB</u>		MUTUAL PHARM	<u>50MG</u>	<u>N75470</u>	<u>001</u>	Feb 21, 2002
<u>AB</u>		MYLAN	<u>50MG</u>	<u>N75463</u>	<u>001</u>	Jul 26, 1999
<u>AB</u>		SANDOZ	<u>50MG</u>	<u>N75229</u>	<u>001</u>	Nov 20, 1998
<u>AB</u>			<u>50MG</u>	<u>N75582</u>	<u>001</u>	Feb 23, 2001
<u>AB</u>		TEVA	<u>50MG</u>	<u>N75219</u>	<u>001</u>	Aug 06, 1998
<u>AB</u>		TORPHARM	<u>50MG</u>	<u>N76561</u>	<u>001</u>	Mar 18, 2004
<u>AB</u>		WATSON LABS	<u>50MG</u>	<u>N75152</u>	<u>001</u>	Nov 27, 1998

DICLOFENAC SODIUM

GEL; TOPICAL

SOLARAZE

+ BIOGLAN PHARMS CORP 3% N21005 001 Oct 16, 2000

SOLUTION/DROPS; OPHTHALMIC

+ NOVARTIS 0.1% N20037 001 Mar 28, 1991

TABLET, DELAYED RELEASE; ORAL

DICLOFENAC SODIUM

<u>AB</u>		ACTAVIS ELIZABETH	<u>50MG</u>	<u>N74514</u>	<u>001</u>	Mar 26, 1996
<u>AB</u>			<u>75MG</u>	<u>N74514</u>	<u>002</u>	Mar 26, 1996
<u>AB</u>		ALPHAPHARM	<u>50MG</u>	<u>N75281</u>	<u>002</u>	Feb 12, 2002
<u>AB</u>			<u>75MG</u>	<u>N75281</u>	<u>003</u>	Feb 12, 2002
<u>AB</u>		CARLSBAD	<u>25MG</u>	<u>N75185</u>	<u>002</u>	Nov 13, 1998
<u>AB</u>			<u>50MG</u>	<u>N75185</u>	<u>003</u>	Nov 13, 1998
<u>AB</u>			<u>75MG</u>	<u>N75185</u>	<u>001</u>	Nov 13, 1998
<u>AB</u>		MARTEC USA LLC	<u>50MG</u>	<u>N74986</u>	<u>001</u>	Feb 26, 1999

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 124 (of 370)

DICLOFENAC SODIUM

TABLET, DELAYED RELEASE; ORAL

DICLOFENAC SODIUM

<u>AB</u>	MARTEC USA LLC	<u>75MG</u>	<u>N74986</u>	<u>002</u>	Feb 26, 1999
<u>AB</u>	PLIVA	<u>50MG</u>	<u>N74432</u>	<u>002</u>	Jul 29, 1999
<u>AB</u>		<u>75MG</u>	<u>N74432</u>	<u>003</u>	Jul 29, 1999
<u>AB</u>	ROXANE	<u>25MG</u>	<u>N74391</u>	<u>001</u>	Jun 29, 1995
<u>AB</u>		<u>50MG</u>	<u>N74391</u>	<u>002</u>	Jun 29, 1995
<u>AB</u>		<u>75MG</u>	<u>N74391</u>	<u>003</u>	Jun 29, 1995
<u>AB</u>	+ SANDOZ	<u>25MG</u>	<u>N74376</u>	<u>001</u>	Sep 28, 1995
<u>AB</u>	+	<u>50MG</u>	<u>N74376</u>	<u>002</u>	Sep 28, 1995
<u>AB</u>		<u>75MG</u>	<u>N74394</u>	<u>001</u>	Nov 30, 1995
<u>AB</u>	TEVA	<u>50MG</u>	<u>N74723</u>	<u>001</u>	Mar 30, 1999
<u>AB</u>		<u>75MG</u>	<u>N74390</u>	<u>001</u>	Aug 15, 1996
<u>AB</u>	TEVA PHARMS	<u>25MG</u>	<u>N74459</u>	<u>001</u>	Jun 25, 1997
<u>AB</u>		<u>50MG</u>	<u>N74459</u>	<u>002</u>	Jun 25, 1997
<u>AB</u>		<u>75MG</u>	<u>N74459</u>	<u>003</u>	Jun 25, 1997

VOLTAREN

<u>AB</u>	+ NOVARTIS	<u>75MG</u>	<u>N19201</u>	<u>003</u>	Jul 28, 1988
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TABLET, EXTENDED RELEASE; ORAL

DICLOFENAC SODIUM

<u>AB</u>	ACTAVIS ELIZABETH	<u>100MG</u>	<u>N75910</u>	<u>001</u>	Jan 07, 2002
<u>AB</u>	BIOVAIL	<u>100MG</u>	<u>N75492</u>	<u>001</u>	Feb 11, 2000
<u>AB</u>	DEXCEL LTD	<u>100MG</u>	<u>N76201</u>	<u>001</u>	Nov 06, 2002
<u>AB</u>	MYLAN	<u>100MG</u>	<u>N76152</u>	<u>001</u>	Dec 13, 2001

VOLTAREN-XR

<u>AB</u>	+ NOVARTIS	<u>100MG</u>	<u>N20254</u>	<u>001</u>	Mar 08, 1996
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DICLOFENAC SODIUM; MISOPROSTOL

TABLET, DELAYED RELEASE; ORAL

ARTHROTEC

	GD SEARLE LLC	50MG;0.2MG	N20607	001	Dec 24, 1997
	+	75MG;0.2MG	N20607	002	Dec 24, 1997

DICLOXACILLIN SODIUM

CAPSULE; ORAL

DICLOXACILLIN SODIUM

<u>AB</u>	SANDOZ	<u>EQ 250MG BASE</u>	<u>N61454</u>	<u>001</u>	
<u>AB</u>	+	<u>EQ 500MG BASE</u>	<u>N61454</u>	<u>003</u>	
		EQ 125MG BASE	N61454	002	
<u>AB</u>	TEVA	<u>EQ 250MG BASE</u>	<u>N62286</u>	<u>001</u>	Jun 03, 1982
<u>AB</u>		<u>EQ 500MG BASE</u>	<u>N62286</u>	<u>002</u>	Jun 03, 1982

FOR SUSPENSION; ORAL

DICLOXACILLIN SODIUM

	+ APOTHECON	EQ 62.5MG BASE/5ML	N61455	001	
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DICYCLOMINE HYDROCHLORIDE

CAPSULE; ORAL

BENTYL

<u>AB</u>	+ AXCAN SCANDIPHARM	<u>10MG</u>	<u>N07409</u>	<u>003</u>	Oct 15, 1984
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DICYCLOMINE HYDROCHLORIDE

<u>AB</u>	LANNETT	<u>10MG</u>	<u>N84285</u>	<u>001</u>	
<u>AB</u>	MYLAN	<u>10MG</u>	<u>N40319</u>	<u>001</u>	Sep 07, 1999
<u>AB</u>	WATSON LABS	<u>10MG</u>	<u>N85082</u>	<u>001</u>	Jun 19, 1986
<u>AB</u>	WEST WARD	<u>10MG</u>	<u>N40204</u>	<u>001</u>	Feb 28, 1997

INJECTABLE; INJECTION

BENTYL

<u>AP</u>	+ AXCAN SCANDIPHARM	<u>10MG/ML</u>	<u>N08370</u>	<u>001</u>	Oct 15, 1984
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27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 125 (of 370)

DICYCLOMINE HYDROCHLORIDE

INJECTABLE; INJECTION

BENTYL PRESERVATIVE FREE

<u>AP</u>	+	AXCAN SCANDIPHARM	<u>10MG/ML</u>	<u>N08370</u>	<u>002</u>	Oct 15, 1984
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DICYCLOMINE HYDROCHLORIDE (PRESERVATIVE-FREE)

<u>AP</u>		BEDFORD	<u>10MG/ML</u>	<u>N40465</u>	<u>001</u>	Jun 30, 2003
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SYRUP; ORAL

BENTYL

<u>AA</u>	+	AXCAN SCANDIPHARM	<u>10MG/5ML</u>	<u>N07961</u>	<u>002</u>	Oct 15, 1984
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DICYCLOMINE HYDROCHLORIDE

<u>AA</u>		MIKART	<u>10MG/5ML</u>	<u>N40169</u>	<u>001</u>	Mar 24, 2005
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TABLET; ORAL

BENTYL

<u>AB</u>	+	AXCAN SCANDIPHARM	<u>20MG</u>	<u>N07409</u>	<u>001</u>	Oct 15, 1984
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DICYCLOMINE HYDROCHLORIDE

<u>AB</u>		LANNETT	<u>20MG</u>	<u>N40230</u>	<u>001</u>	Feb 26, 1999
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<u>AB</u>		MYLAN	<u>20MG</u>	<u>N40317</u>	<u>001</u>	Sep 07, 1999
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<u>AB</u>		WATSON LABS	<u>20MG</u>	<u>N85223</u>	<u>001</u>	Jul 30, 1986
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<u>AB</u>		WEST WARD	<u>20MG</u>	<u>N40161</u>	<u>001</u>	Oct 01, 1996
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DIDANOSINE

CAPSULE, DELAYED REL PELLETS; ORAL

DIDANOSINE

<u>AB</u>		BARR	<u>200MG</u>	<u>N77167</u>	<u>001</u>	Dec 03, 2004
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<u>AB</u>			<u>250MG</u>	<u>N77167</u>	<u>002</u>	Dec 03, 2004
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<u>AB</u>			<u>400MG</u>	<u>N77167</u>	<u>003</u>	Dec 03, 2004
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VIDEX EC

<u>AB</u>		BRISTOL MYERS SQUIBB	<u>200MG</u>	<u>N21183</u>	<u>002</u>	Oct 31, 2000
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<u>AB</u>			<u>250MG</u>	<u>N21183</u>	<u>003</u>	Oct 31, 2000
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<u>AB</u>	+		<u>400MG</u>	<u>N21183</u>	<u>004</u>	Oct 31, 2000
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			125MG	N21183	001	Oct 31, 2000
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FOR SOLUTION; ORAL

VIDEX

	+	BRISTOL MYERS SQUIBB	10MG/ML	N20156	001	Oct 09, 1991
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TABLET, CHEWABLE; ORAL

VIDEX

		BRISTOL MYERS SQUIBB	25MG	N20154	002	Oct 09, 1991
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			50MG	N20154	003	Oct 09, 1991
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			100MG	N20154	004	Oct 09, 1991
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			150MG	N20154	005	Oct 09, 1991
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	+		200MG	N20154	006	Oct 28, 1999
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DIETHYLPROPION HYDROCHLORIDE

TABLET; ORAL

TENUATE

	+	WATSON PHARMS	25MG	N11722	002	
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TABLET, EXTENDED RELEASE; ORAL

TENUATE DOSPAN

	+	WATSON PHARMS	75MG	N12546	001	
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DIFLORASONE DIACETATE

CREAM; TOPICAL

DIFLORASONE DIACETATE

<u>AB1</u>		ALTANA	<u>0.05%</u>	<u>N75187</u>	<u>001</u>	Mar 30, 1998
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<u>BX</u>	+		<u>0.05%</u>	N76263	001	Dec 20, 2002
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<u>AB1</u>		TARO	<u>0.05%</u>	<u>N75508</u>	<u>001</u>	Apr 24, 2000
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PSORCON

<u>AB1</u>	+	SANOFI AVENTIS US	<u>0.05%</u>	<u>N20205</u>	<u>001</u>	Nov 20, 1992
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27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 126 (of 370)

DIFLORASONE DIACETATE

OINTMENT; TOPICAL

DIFLORASONE DIACETATE

<u>AB</u>	ALTANA	<u>0.05%</u>	<u>N75374</u>	<u>001</u>	Apr 27, 1999
<u>AB</u>	+ TARO	<u>0.05%</u>	<u>N75331</u>	<u>001</u>	May 14, 1999

DIFLUNISAL

TABLET; ORAL

DIFLUNISAL

<u>AB</u>	SANDOZ	<u>500MG</u>	<u>N74604</u>	<u>001</u>	Jun 10, 1996
<u>AB</u>	TEVA	<u>250MG</u>	<u>N73679</u>	<u>001</u>	Jul 31, 1992
<u>AB</u>	+	<u>500MG</u>	<u>N73673</u>	<u>001</u>	Jul 31, 1992
<u>AB</u>	WATSON LABS	<u>250MG</u>	<u>N74400</u>	<u>001</u>	Jul 17, 1997
<u>AB</u>		<u>500MG</u>	<u>N74400</u>	<u>002</u>	Jul 17, 1997

DIGOXIN

CAPSULE; ORAL

LANOXICAPS

	SMITHKLINE BEECHAM	0.05MG	N18118	002	Jul 26, 1982
		0.1MG	N18118	003	Jul 26, 1982
+		0.2MG	N18118	001	Jul 26, 1982

ELIXIR; ORAL

DIGOXIN

+	ROXANE	0.05MG/ML	N21648	001	Aug 26, 2004
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INJECTABLE; INJECTION

DIGOXIN

<u>AP</u>	BAXTER HLTHCARE	<u>0.25MG/ML</u>	<u>N83391</u>	<u>001</u>	
<u>AP</u>	HOSPIRA	<u>0.25MG/ML</u>	<u>N40093</u>	<u>001</u>	May 16, 1996
<u>AP</u>		<u>0.25MG/ML</u>	<u>N40206</u>	<u>001</u>	Aug 28, 1998
<u>AP</u>	SANDOZ	<u>0.25MG/ML</u>	<u>N40481</u>	<u>001</u>	Aug 21, 2003

DIGOXIN PEDIATRIC

<u>AP</u>	HOSPIRA	<u>0.1MG/ML</u>	<u>N40092</u>	<u>001</u>	Apr 25, 1996
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LANOXIN

<u>AP</u>	+	GLAXOSMITHKLINE	<u>0.25MG/ML</u>	<u>N09330</u>	<u>002</u>
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LANOXIN PEDIATRIC

<u>AP</u>	+	GLAXOSMITHKLINE	<u>0.1MG/ML</u>	<u>N09330</u>	<u>004</u>
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TABLET; ORAL

DIGOXIN

<u>AB</u>	ACTAVIS TOTOWA	<u>0.125MG</u>	<u>N40282</u>	<u>001</u>	Dec 23, 1999
<u>AB</u>		<u>0.25MG</u>	<u>N40282</u>	<u>002</u>	Dec 23, 1999
<u>AB</u>	CARACO	<u>0.125MG</u>	<u>N76363</u>	<u>001</u>	Jan 31, 2003
<u>AB</u>		<u>0.25MG</u>	<u>N76363</u>	<u>002</u>	Jan 31, 2003
<u>AB</u>	STEVENS J	<u>0.125MG</u>	<u>N76268</u>	<u>001</u>	Jul 26, 2002
<u>AB</u>		<u>0.25MG</u>	<u>N76268</u>	<u>002</u>	Jul 26, 2002

LANOXIN

<u>AB</u>	SMITHKLINE BEECHAM	<u>0.125MG</u>	<u>N20405</u>	<u>002</u>	Sep 30, 1997
<u>AB</u>	+	<u>0.25MG</u>	<u>N20405</u>	<u>004</u>	Sep 30, 1997

DIHYDROERGOTAMINE MESYLATE

INJECTABLE; INJECTION

D.H.E. 45

<u>AP</u>	+	VALEANT	<u>1MG/ML</u>	<u>N05929</u>	<u>001</u>
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DIHYDROERGOTAMINE MESYLATE

<u>AP</u>	BEDFORD LABS	<u>1MG/ML</u>	<u>N40453</u>	<u>001</u>	Jun 09, 2003
<u>AP</u>	PADDOCK	<u>1MG/ML</u>	<u>N40475</u>	<u>001</u>	Apr 28, 2003

SPRAY, METERED; NASAL

MIGRANAL

+	VALEANT	0.5MG/INH	N20148	001	Dec 08, 1997
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PRESCRIPTION DRUG PRODUCT LIST

3 - 127 (of 370)

DILTIAZEM HYDROCHLORIDE

CAPSULE, EXTENDED RELEASE; ORAL

CARDIZEM CD

<u>AB3</u>	<u>BIOVAIL</u>	<u>120MG</u>	<u>N20062</u>	<u>001</u>	Aug 10, 1992
<u>AB3</u>		<u>180MG</u>	<u>N20062</u>	<u>002</u>	Dec 27, 1991
<u>AB3</u>		<u>240MG</u>	<u>N20062</u>	<u>003</u>	Dec 27, 1991
<u>AB3</u>		<u>300MG</u>	<u>N20062</u>	<u>004</u>	Dec 27, 1991
BC +		360MG	N20062	005	Aug 24, 1999

CARTIA XT

<u>AB3</u>	<u>ANDRX PHARMS</u>	<u>120MG</u>	<u>N74752</u>	<u>002</u>	Jul 09, 1998
<u>AB3</u>		<u>180MG</u>	<u>N74752</u>	<u>001</u>	Jul 09, 1998
<u>AB3</u>		<u>240MG</u>	<u>N74752</u>	<u>003</u>	Jul 09, 1998
<u>AB3</u>		<u>300MG</u>	<u>N74752</u>	<u>004</u>	Jul 09, 1998

DILACOR XR

<u>AB2</u>	WATSON LABS	<u>120MG</u>	<u>N20092</u>	<u>001</u>	May 29, 1992
<u>AB2</u>		<u>180MG</u>	<u>N20092</u>	<u>002</u>	May 29, 1992
<u>AB2</u> +		<u>240MG</u>	<u>N20092</u>	<u>003</u>	May 29, 1992

DILT-CD

AB3	TORPHARM	120MG	N76151	001	May 20, 2004
AB3		180MG	N76151	002	May 20, 2004
AB3		240MG	N76151	003	May 20, 2004
AB3		300MG	N76151	004	May 20, 2004

DILTIAZEM HYDROCHLORIDE

<u>AB3</u>	ACTAVIS ELIZABETH	<u>120MG</u>	<u>N74984</u>	<u>001</u>	Dec 20, 1999
<u>AB3</u>		<u>180MG</u>	<u>N74984</u>	<u>002</u>	Dec 20, 1999
<u>AB3</u>		<u>240MG</u>	<u>N74984</u>	<u>003</u>	Dec 20, 1999
<u>AB3</u>		<u>300MG</u>	<u>N74984</u>	<u>004</u>	Dec 20, 1999
<u>AB2</u>	ANDRX	<u>120MG</u>	<u>N74852</u>	<u>001</u>	Oct 10, 1997
<u>AB2</u>		<u>180MG</u>	<u>N74852</u>	<u>002</u>	Oct 10, 1997
<u>AB2</u>		<u>240MG</u>	<u>N74852</u>	<u>003</u>	Oct 10, 1997
<u>AB3</u>	BIOVAIL	<u>120MG</u>	<u>N20939</u>	<u>001</u>	Jan 28, 2000
<u>AB3</u>		<u>120MG</u>	<u>N75116</u>	<u>001</u>	Dec 23, 1999
<u>AB3</u>		<u>180MG</u>	<u>N20939</u>	<u>002</u>	Jan 28, 2000
<u>AB3</u>		<u>180MG</u>	<u>N75116</u>	<u>002</u>	Dec 23, 1999
<u>AB3</u>		<u>240MG</u>	<u>N20939</u>	<u>003</u>	Jan 28, 2000
<u>AB3</u>		<u>240MG</u>	<u>N75116</u>	<u>003</u>	Dec 23, 1999
<u>AB3</u>		<u>300MG</u>	<u>N20939</u>	<u>004</u>	Jan 28, 2000
<u>AB3</u>		<u>300MG</u>	<u>N75116</u>	<u>004</u>	Dec 23, 1999
<u>AB4</u>	KV PHARM	<u>120MG</u>	<u>N76563</u>	<u>002</u>	Sep 12, 2006
<u>AB4</u>		<u>180MG</u>	<u>N76563</u>	<u>003</u>	Sep 12, 2006
<u>AB4</u>		<u>240MG</u>	<u>N76563</u>	<u>004</u>	Sep 12, 2006
<u>AB4</u>		<u>300MG</u>	<u>N76563</u>	<u>005</u>	Sep 12, 2006
<u>AB4</u>		<u>360MG</u>	<u>N76563</u>	<u>006</u>	Sep 12, 2006
<u>AB4</u>		<u>420MG</u>	<u>N76563</u>	<u>001</u>	Sep 12, 2006
<u>AB2</u>	MYLAN	<u>120MG</u>	<u>N75124</u>	<u>002</u>	Mar 18, 1998
<u>AB2</u>		<u>180MG</u>	<u>N75124</u>	<u>003</u>	Mar 18, 1998
<u>AB2</u>		<u>240MG</u>	<u>N75124</u>	<u>001</u>	Mar 18, 1998
		60MG	N74910	001	May 02, 1997
		90MG	N74910	002	May 02, 1997
		120MG	N74910	003	May 02, 1997
<u>AB2</u>	TORPHARM	<u>120MG</u>	<u>N74943</u>	<u>003</u>	Dec 19, 2000
<u>AB2</u>		<u>180MG</u>	<u>N74943</u>	<u>002</u>	Dec 19, 2000
<u>AB2</u>		<u>240MG</u>	<u>N74943</u>	<u>001</u>	Aug 06, 1998
	<u>DILTZAC</u>				
<u>AB4</u>	APOTEX INC	<u>120MG</u>	<u>N76395</u>	<u>001</u>	Feb 01, 2006
<u>AB4</u>		<u>180MG</u>	<u>N76395</u>	<u>002</u>	Feb 01, 2006
<u>AB4</u>		<u>240MG</u>	<u>N76395</u>	<u>003</u>	Feb 01, 2006
<u>AB4</u>		<u>300MG</u>	<u>N76395</u>	<u>004</u>	Feb 01, 2006
<u>AB4</u>		<u>360MG</u>	<u>N76395</u>	<u>005</u>	Feb 01, 2006

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 128 (of 370)

DILTIAZEM HYDROCHLORIDE

CAPSULE, EXTENDED RELEASE; ORAL

TAZTIA XT

<u>AB4</u>	ANDRX PHARMS	<u>120MG</u>	<u>N75401</u>	<u>001</u>	Apr 10, 2003
<u>AB4</u>		<u>180MG</u>	<u>N75401</u>	<u>002</u>	Apr 10, 2003
<u>AB4</u>		<u>240MG</u>	<u>N75401</u>	<u>003</u>	Apr 10, 2003
<u>AB4</u>		<u>300MG</u>	<u>N75401</u>	<u>004</u>	Apr 10, 2003
<u>AB4</u>		<u>360MG</u>	<u>N75401</u>	<u>005</u>	Apr 10, 2003

TIAZAC

<u>AB4</u>	BIOVAIL	<u>120MG</u>	<u>N20401</u>	<u>001</u>	Sep 11, 1995
<u>AB4</u>		<u>180MG</u>	<u>N20401</u>	<u>002</u>	Sep 11, 1995
<u>AB4</u>		<u>240MG</u>	<u>N20401</u>	<u>003</u>	Sep 11, 1995
<u>AB4</u>		<u>300MG</u>	<u>N20401</u>	<u>004</u>	Sep 11, 1995
<u>AB4</u>		<u>360MG</u>	<u>N20401</u>	<u>005</u>	Sep 11, 1995
<u>AB4</u> +		<u>420MG</u>	<u>N20401</u>	<u>006</u>	Oct 16, 1998

INJECTABLE; INJECTION

CARDIZEM

<u>AP</u> +	BIOVAIL	<u>100MG/VIAL</u>	<u>N20792</u>	<u>001</u>	Sep 05, 1997
<u>AP</u> +	BIOVAIL LABS INTL	<u>5MG/ML</u>	<u>N20027</u>	<u>001</u>	Oct 24, 1991
<u>AP</u> +		25MG/VIAL	N20027	003	Aug 18, 1995

DILTIAZEM HYDROCHLORIDE

<u>AP</u>	APOTEX INC	<u>5MG/ML</u>	<u>N75375</u>	<u>001</u>	Sep 30, 1999
<u>AP</u> +	BEDFORD	<u>5MG/ML</u>	<u>N74617</u>	<u>001</u>	Feb 28, 1996
<u>AP</u>	HOSPIRA	<u>5MG/ML</u>	<u>N74941</u>	<u>001</u>	Apr 15, 1998
<u>AP</u>		<u>5MG/ML</u>	<u>N75004</u>	<u>001</u>	Feb 16, 2000
<u>AP</u>		<u>100MG/VIAL</u>	<u>N75853</u>	<u>001</u>	Dec 17, 2002
<u>AP</u>	INTL MEDICATION	<u>5MG/ML</u>	<u>N75749</u>	<u>001</u>	Nov 21, 2001
<u>AP</u>	MAYNE PHARMA USA	<u>5MG/ML</u>	<u>N75106</u>	<u>001</u>	Apr 29, 1999
<u>AP</u>	SICOR PHARMS	<u>5MG/ML</u>	<u>N74894</u>	<u>001</u>	Aug 26, 1997
<u>AP</u> +		10MG/ML	N74894	002	Apr 19, 2002
<u>AP</u>	TAYLOR PHARMA	<u>5MG/ML</u>	<u>N75086</u>	<u>001</u>	Apr 09, 1998

TABLET; ORAL

CARDIZEM

<u>AB</u>	BIOVAIL LABS INTL	<u>30MG</u>	<u>N18602</u>	<u>001</u>	Nov 05, 1982
<u>AB</u>		<u>60MG</u>	<u>N18602</u>	<u>002</u>	Nov 05, 1982
<u>AB</u>		<u>90MG</u>	<u>N18602</u>	<u>003</u>	Dec 08, 1986
<u>AB</u> +		<u>120MG</u>	<u>N18602</u>	<u>004</u>	Dec 08, 1986

DILTIAZEM HYDROCHLORIDE

<u>AB</u>	CLONMEL HLTHCARE	<u>30MG</u>	<u>N74093</u>	<u>001</u>	Nov 05, 1992
<u>AB</u>		<u>60MG</u>	<u>N74093</u>	<u>002</u>	Nov 05, 1992
<u>AB</u>		<u>90MG</u>	<u>N74093</u>	<u>003</u>	Nov 05, 1992
<u>AB</u>		<u>120MG</u>	<u>N74093</u>	<u>004</u>	Nov 05, 1992
<u>AB</u>	IVAX PHARMS	<u>30MG</u>	<u>N74168</u>	<u>001</u>	Mar 03, 1995
<u>AB</u>		<u>60MG</u>	<u>N74168</u>	<u>002</u>	Mar 03, 1995
<u>AB</u>		<u>90MG</u>	<u>N74168</u>	<u>003</u>	Mar 03, 1995
<u>AB</u>		<u>120MG</u>	<u>N74168</u>	<u>004</u>	Mar 03, 1995
<u>AB</u>	MYLAN	<u>30MG</u>	<u>N73185</u>	<u>001</u>	Nov 05, 1992
<u>AB</u>		<u>60MG</u>	<u>N73186</u>	<u>001</u>	Nov 05, 1992
<u>AB</u>		<u>90MG</u>	<u>N72837</u>	<u>001</u>	Nov 05, 1992
<u>AB</u>		<u>120MG</u>	<u>N72838</u>	<u>001</u>	Nov 05, 1992
<u>AB</u>	TEVA	<u>30MG</u>	<u>N74185</u>	<u>001</u>	May 31, 1995
<u>AB</u>		<u>60MG</u>	<u>N74185</u>	<u>002</u>	May 31, 1995
<u>AB</u>		<u>90MG</u>	<u>N74185</u>	<u>003</u>	May 31, 1995
<u>AB</u>		<u>120MG</u>	<u>N74185</u>	<u>004</u>	May 31, 1995

TABLET, EXTENDED RELEASE; ORAL

CARDIZEM LA

	BIOVAIL LABS INTL	120MG	N21392	001	Feb 06, 2003
		180MG	N21392	002	Feb 06, 2003
		240MG	N21392	003	Feb 06, 2003
		300MG	N21392	004	Feb 06, 2003

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 129 (of 370)

DILTIAZEM HYDROCHLORIDE

TABLET, EXTENDED RELEASE; ORAL

CARDIZEM LA

BIOVAIL LABS INTL 360MG

N21392 005 Feb 06, 2003

+ 420MG

N21392 006 Feb 06, 2003

DIMENHYDRINATE

INJECTABLE; INJECTION

DIMENHYDRINATEAP ABRAXIS PHARM 50MG/MLN40519 001 Jun 23, 2004AP + WATSON LABS 50MG/MLN80615 001DIMERCAPROL

INJECTABLE; INJECTION

BAL

+ AKORN 10%

N05939 001

DIMETHYL SULFOXIDE

SOLUTION; INTRAVESICAL

DIMETHYL SULFOXIDEAT BIONICHE (CANADA) 50%N76185 001 Nov 29, 2002RIMS0-50AT + BIONICHE PHARMA 50%N17788 001DINOPROSTONE

GEL; ENDOCERVICAL

PREPIDIL

+ PHARMACIA AND UPJOHN 0.5MG/3GM

N19617 001 Dec 09, 1992

INSERT, EXTENDED RELEASE; VAGINAL

CERVIDIL

+ CONTROLLED THERAP 10MG

N20411 001 Mar 30, 1995

SUPPOSITORY; VAGINAL

PROSTIN E2

+ PHARMACIA AND UPJOHN 20MG

N17810 001

DIPHENHYDRAMINE HYDROCHLORIDE

CAPSULE; ORAL

DIPHENHYDRAMINE HYDROCHLORIDEAA BARR 50MGN80738 001AA LNK 25MGN87977 001 Jan 27, 1983AA 50MGN87978 001 Jan 27, 1983AA MUTUAL PHARM 25MGN84506 001AA 25MGN89488 001 Jan 02, 1987AA 50MGN89489 001 Jan 02, 1987AA + SANDOZ 25MGN80832 001AA + 50MGN80832 002AA VALEANT PHARM INTL 50MGN80592 001AA WATSON LABS 25MGN80728 001AA 50MGN80727 001

ELIXIR; ORAL

DIPHENHYDRAMINE HYDROCHLORIDEAA + PHARM ASSOC 12.5MG/5MLN87513 001 Feb 10, 1982

INJECTABLE; INJECTION

BENADRYLAP + PARKE DAVIS 50MG/MLN06146 002BENADRYL PRESERVATIVE FREEAP + PARKE DAVIS 50MG/MLN09486 001DIPHENHYDRAMINE HYDROCHLORIDEAP ABRAXIS PHARM 50MG/MLN40466 001 May 28, 2002

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 130 (of 370)

DIPHENHYDRAMINE HYDROCHLORIDE

INJECTABLE; INJECTION

DIPHENHYDRAMINE HYDROCHLORIDE

<u>AP</u>	BAXTER HLTHCARE	<u>50MG/ML</u>	<u>N80817</u>	<u>002</u>	
<u>AP</u>	BIONICHE PHARMA	<u>50MG/ML</u>	<u>N40498</u>	<u>001</u>	Jul 12, 2005
<u>AP</u>	HOSPIRA	<u>50MG/ML</u>	<u>N40140</u>	<u>001</u>	Nov 20, 1998
<u>AP</u>	WATSON LABS	<u>50MG/ML</u>	<u>N80873</u>	<u>002</u>	
	+	10MG/ML	<u>N80873</u>	<u>001</u>	

DIPHENHYDRAMINE HYDROCHLORIDE PRESERVATIVE FREE

<u>AP</u>	INTL MEDICATION	<u>50MG/ML</u>	<u>N84094</u>	<u>001</u>	
<u>AP</u>	WATSON LABS	<u>50MG/ML</u>	<u>N80873</u>	<u>003</u>	

DIPIVEFRIN HYDROCHLORIDE

SOLUTION/DROPS; OPHTHALMIC

AKPRO

<u>AT</u>	AKORN	<u>0.1%</u>	<u>N74382</u>	<u>001</u>	Sep 29, 1995
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DIPIVEFRIN HYDROCHLORIDE

<u>AT</u>	BAUSCH AND LOMB	<u>0.1%</u>	<u>N74188</u>	<u>001</u>	May 19, 1995
<u>AT</u>	FALCON PHARMS	<u>0.1%</u>	<u>N73636</u>	<u>001</u>	Jun 30, 1994

PROPINE

<u>AT</u>	+	ALLERGAN	<u>0.1%</u>	<u>N18239</u>	<u>001</u>
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DIPYRIDAMOLE

INJECTABLE; INJECTION

DIPYRIDAMOLE

<u>AP</u>	ABRAXIS PHARM	<u>5MG/ML</u>	<u>N74956</u>	<u>001</u>	Sep 30, 1998
<u>AP</u>	APOTEX INC	<u>5MG/ML</u>	<u>N75769</u>	<u>001</u>	Nov 27, 2002
<u>AP</u>	BAXTER HLTHCARE	<u>5MG/ML</u>	<u>N74521</u>	<u>001</u>	Oct 18, 1996
<u>AP</u>	+	BEDFORD	<u>5MG/ML</u>	<u>N74939</u>	<u>001</u>
<u>AP</u>	HOSPIRA	<u>5MG/ML</u>	<u>N74601</u>	<u>001</u>	Dec 19, 1997
<u>AP</u>	SICOR PHARMS	<u>5MG/ML</u>	<u>N74952</u>	<u>001</u>	Nov 26, 1997

TABLET; ORAL

DIPYRIDAMOLE

<u>AB</u>	ACTAVIS TOTOWA	<u>25MG</u>	<u>N40542</u>	<u>001</u>	Apr 21, 2006
<u>AB</u>		<u>50MG</u>	<u>N40542</u>	<u>002</u>	Apr 21, 2006
<u>AB</u>		<u>75MG</u>	<u>N40542</u>	<u>003</u>	Apr 21, 2006
<u>AB</u>	BARR	<u>25MG</u>	<u>N87184</u>	<u>001</u>	Oct 03, 1990
<u>AB</u>		<u>50MG</u>	<u>N87716</u>	<u>001</u>	Oct 03, 1990
<u>AB</u>		<u>75MG</u>	<u>N87717</u>	<u>001</u>	Oct 03, 1990
<u>AB</u>	GLENMARK PHARMA	<u>25MG</u>	<u>N88999</u>	<u>001</u>	Feb 05, 1991
<u>AB</u>		<u>50MG</u>	<u>N89000</u>	<u>001</u>	Feb 05, 1991
<u>AB</u>		<u>75MG</u>	<u>N89001</u>	<u>001</u>	Feb 05, 1991
<u>AB</u>	PUREPAC PHARM	<u>25MG</u>	<u>N89425</u>	<u>001</u>	Jul 12, 1990
<u>AB</u>	SANDOZ	<u>25MG</u>	<u>N86944</u>	<u>002</u>	Apr 16, 1991
<u>AB</u>		<u>50MG</u>	<u>N87562</u>	<u>001</u>	Feb 25, 1992
<u>AB</u>		<u>75MG</u>	<u>N87561</u>	<u>001</u>	Feb 25, 1992
<u>AB</u>	WATSON LABS	<u>50MG</u>	<u>N87160</u>	<u>001</u>	Jun 07, 1996
	<u>PERSANTINE</u>				
<u>AB</u>	BOEHRINGER INGELHEIM	<u>25MG</u>	<u>N12836</u>	<u>003</u>	Dec 22, 1986
<u>AB</u>		<u>50MG</u>	<u>N12836</u>	<u>004</u>	Feb 06, 1987
<u>AB</u>	+	<u>75MG</u>	<u>N12836</u>	<u>005</u>	Feb 06, 1987

DISOPYRAMIDE PHOSPHATE

CAPSULE; ORAL

DISOPYRAMIDE PHOSPHATE

<u>AB</u>	TEVA	<u>EQ 100MG BASE</u>	<u>N70101</u>	<u>001</u>	Feb 22, 1985
<u>AB</u>		<u>EQ 150MG BASE</u>	<u>N70102</u>	<u>001</u>	Feb 22, 1985
<u>AB</u>	WATSON LABS	<u>EQ 100MG BASE</u>	<u>N70173</u>	<u>001</u>	May 31, 1985
<u>AB</u>		<u>EQ 150MG BASE</u>	<u>N70174</u>	<u>001</u>	May 31, 1985

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 131 (of 370)

DISOPYRAMIDE PHOSPHATE

CAPSULE; ORAL

NORPACE

<u>AB</u>	GD SEARLE LLC	<u>EQ 100MG BASE</u>	<u>N17447</u>	<u>001</u>	
<u>AB</u>	+	<u>EQ 150MG BASE</u>	<u>N17447</u>	<u>002</u>	

CAPSULE, EXTENDED RELEASE; ORAL

DISOPYRAMIDE PHOSPHATE

<u>AB</u>	KV PHARM	<u>EQ 150MG BASE</u>	<u>N71200</u>	<u>001</u>	Dec 15, 1987
	<u>NORPACE CR</u>				
<u>AB</u>	+	GD SEARLE LLC	<u>EQ 150MG BASE</u>	<u>N18655</u>	<u>002</u>
			<u>EQ 100MG BASE</u>	<u>N18655</u>	<u>001</u>

DISULFIRAM

TABLET; ORAL

ANTABUSE

	ODYSSEY PHARMS	250MG	N88482	001	Dec 08, 1983
+		500MG	N88483	001	Dec 08, 1983

DIVALPROEX SODIUM

CAPSULE, DELAYED REL PELLETS; ORAL

DEPAKOTE

+	ABBOTT	EQ 125MG VALPROIC ACID	N19680	001	Sep 12, 1989
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TABLET, DELAYED RELEASE; ORAL

DEPAKOTE

	ABBOTT	EQ 125MG VALPROIC ACID	N18723	003	Oct 26, 1984
		EQ 250MG VALPROIC ACID	N18723	001	Mar 10, 1983
+		EQ 500MG VALPROIC ACID	N18723	002	Mar 10, 1983

TABLET, EXTENDED RELEASE; ORAL

DEPAKOTE ER

	ABBOTT	EQ 250MG VALPROIC ACID	N21168	002	May 31, 2002
+		EQ 500MG VALPROIC ACID	N21168	001	Aug 04, 2000

DOBUTAMINE HYDROCHLORIDE

INJECTABLE; INJECTION

DOBUTAMINE HYDROCHLORIDE

<u>AP</u>	BEDFORD	<u>EQ 12.5MG BASE/ML</u>	<u>N74277</u>	<u>001</u>	Oct 31, 1994
<u>AP</u>	HOSPIRA	<u>EQ 12.5MG BASE/ML</u>	<u>N74086</u>	<u>001</u>	Nov 29, 1993
<u>AP</u>		<u>EQ 12.5MG BASE/ML</u>	<u>N74292</u>	<u>001</u>	Feb 16, 1995
<u>AP</u>		<u>EQ 1.25GM BASE/100ML</u>	<u>N74634</u>	<u>001</u>	Sep 27, 1996
<u>AP</u>	LUITPOLD	<u>EQ 12.5MG BASE/ML</u>	<u>N74545</u>	<u>001</u>	Jun 25, 1998
<u>AP</u>	MARSAM PHARMS LLC	<u>EQ 12.5MG BASE/ML</u>	<u>N74279</u>	<u>001</u>	Feb 18, 1998
<u>AP</u>		<u>EQ 12.5MG BASE/ML</u>	<u>N74995</u>	<u>001</u>	Mar 31, 1998
<u>AP</u>	SICOR PHARMS	<u>EQ 12.5MG BASE/ML</u>	<u>N74206</u>	<u>001</u>	Oct 19, 1993
<u>AP</u>	WATSON LABS	<u>EQ 12.5MG BASE/ML</u>	<u>N74114</u>	<u>001</u>	Nov 30, 1993

DOBUTAMINE HYDROCHLORIDE IN DEXTROSE 5%

<u>AP</u>	+	HOSPIRA	<u>EQ 50MG BASE/100ML</u>	<u>N20269</u>	<u>001</u>	Oct 19, 1993
<u>AP</u>	+		<u>EQ 100MG BASE/100ML</u>	<u>N20269</u>	<u>002</u>	Oct 19, 1993
<u>AP</u>	+		<u>EQ 200MG BASE/100ML</u>	<u>N20269</u>	<u>003</u>	Oct 19, 1993

DOBUTAMINE HYDROCHLORIDE IN DEXTROSE 5% IN PLASTIC CONTAINER

<u>AP</u>	+	BAXTER HLTHCARE	<u>EQ 50MG BASE/100ML</u>	<u>N20255</u>	<u>001</u>	Oct 19, 1993
<u>AP</u>	+		<u>EQ 100MG BASE/100ML</u>	<u>N20255</u>	<u>003</u>	Oct 19, 1993
<u>AP</u>	+		<u>EQ 200MG BASE/100ML</u>	<u>N20255</u>	<u>004</u>	Oct 19, 1993
<u>AP</u>	+		<u>EQ 400MG BASE/100ML</u>	<u>N20255</u>	<u>005</u>	Oct 19, 1993
<u>AP</u>	+	HOSPIRA	<u>EQ 50MG BASE/100ML</u>	<u>N20201</u>	<u>003</u>	Oct 19, 1993
<u>AP</u>	+		<u>EQ 100MG BASE/100ML</u>	<u>N20201</u>	<u>002</u>	Oct 19, 1993
<u>AP</u>	+		<u>EQ 200MG BASE/100ML</u>	<u>N20201</u>	<u>001</u>	Oct 19, 1993
<u>AP</u>	+		<u>EQ 400MG BASE/100ML</u>	<u>N20201</u>	<u>006</u>	Jul 07, 1994

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 132 (of 370)

DOCETAXELINJECTABLE; INJECTION
TAXOTERE

+	SANOFI AVENTIS US	EQ 40MG BASE/ML	N20449	001	May 14, 1996
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DOFETILIDECAPSULE; ORAL
TIKOSYN
PFIZER

	0.125MG	N20931	001	Oct 01, 1999
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	0.25MG	N20931	002	Oct 01, 1999
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+	0.5MG	N20931	003	Oct 01, 1999
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DOLASETRON MESYLATEINJECTABLE; INJECTION
ANZEMET

+	SANOFI AVENTIS US	EQ 12.5MG BASE/0.625ML (EQ 20MG BASE/ML)	N20624	002	Sep 11, 1997
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+		EQ 100MG BASE/5ML (EQ 20MG BASE/ML)	N20624	001	Sep 11, 1997
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+		EQ 500MG BASE/25ML (EQ 20MG BASE/ML)	N20624	003	Dec 11, 2001
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TABLET; ORAL
ANZEMET

	SANOFI AVENTIS US	EQ 50MG BASE	N20623	001	Sep 11, 1997
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+		EQ 100MG BASE	N20623	002	Sep 11, 1997
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DONEPEZIL HYDROCHLORIDETABLET; ORAL
ARICEPT

	EISAI MEDCL RES	5MG	N20690	002	Nov 25, 1996
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+		10MG	N20690	001	Nov 25, 1996
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TABLET, ORALLY DISINTEGRATING; ORAL
ARICEPT ODT

	EISAI MEDCL RES	5MG	N21720	001	Oct 18, 2004
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+		10MG	N21720	002	Oct 18, 2004
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DOPAMINE HYDROCHLORIDEINJECTABLE; INJECTION
DOPAMINE HYDROCHLORIDE

<u>AP</u>	+	HOSPIRA	<u>40MG/ML</u>	<u>N18132</u>	<u>001</u>	
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<u>AP</u>	+		<u>80MG/100ML</u>	<u>N18132</u>	<u>002</u>	Feb 04, 1982
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<u>AP</u>	+		<u>80MG/ML</u>	<u>N18132</u>	<u>004</u>	Jul 09, 1982
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<u>AP</u>	+		<u>160MG/100ML</u>	<u>N18132</u>	<u>003</u>	Feb 04, 1982
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<u>AP</u>	+	LUITPOLD	<u>40MG/ML</u>	<u>N70799</u>	<u>001</u>	Feb 11, 1987
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<u>AP</u>	+		<u>80MG/ML</u>	<u>N70820</u>	<u>001</u>	Feb 11, 1987
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<u>AP</u>	+		<u>160MG/ML</u>	<u>N70826</u>	<u>001</u>	Feb 11, 1987
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DOPAMINE HYDROCHLORIDE AND DEXTROSE 5%

<u>AP</u>	+	B BRAUN	<u>80MG/100ML</u>	<u>N19099</u>	<u>002</u>	Oct 15, 1986
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<u>AP</u>	+		<u>320MG/100ML</u>	<u>N19099</u>	<u>004</u>	Oct 15, 1986
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DOPAMINE HYDROCHLORIDE AND DEXTROSE 5% IN PLASTIC CONTAINER

<u>AP</u>	+	B BRAUN	<u>160MG/100ML</u>	<u>N19099</u>	<u>003</u>	Oct 15, 1986
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	+		40MG/100ML	N19099	001	Oct 15, 1986
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DOPAMINE HYDROCHLORIDE IN DEXTROSE 5% IN PLASTIC CONTAINER

<u>AP</u>	+	BAXTER HLTHCARE	<u>80MG/100ML</u>	<u>N19615</u>	<u>001</u>	Mar 27, 1987
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<u>AP</u>	+		<u>160MG/100ML</u>	<u>N19615</u>	<u>002</u>	Mar 27, 1987
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<u>AP</u>	+		<u>320MG/100ML</u>	<u>N19615</u>	<u>003</u>	Mar 27, 1987
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	+		640MG/100ML	N19615	004	Mar 27, 1987
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<u>AP</u>	+	HOSPIRA	<u>80MG/100ML</u>	<u>N18826</u>	<u>001</u>	Sep 30, 1983
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<u>AP</u>	+		<u>160MG/100ML</u>	<u>N18826</u>	<u>002</u>	Sep 30, 1983
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<u>AP</u>	+		<u>320MG/100ML</u>	<u>N18826</u>	<u>003</u>	Sep 30, 1983
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27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 133 (of 370)

DORZOLAMIDE HYDROCHLORIDESOLUTION/DROPS; OPHTHALMIC
TRUSOPT

+ MERCK EQ 2% BASE N20408 001 Dec 09, 1994

DORZOLAMIDE HYDROCHLORIDE; TIMOLOL MALEATE

SOLUTION/DROPS; OPHTHALMIC

+ MERCK EQ 2% BASE;EQ 0.5% BASE N20869 001 Apr 07, 1998

DOXAPRAM HYDROCHLORIDE

INJECTABLE; INJECTION

DOPRAMAP + BAXTER HLTHCARE 20MG/ML N14879 001DOXAPRAM HYDROCHLORIDEAP BEDFORD 20MG/ML N76266 001 Jan 10, 2003AP WATSON LABS 20MG/ML N73529 001 Jan 30, 1992DOXAZOSIN MESYLATE

TABLET; ORAL

CARDURAAB + PFIZER EQ 1MG BASE N19668 001 Nov 02, 1990AB EQ 2MG BASE N19668 002 Nov 02, 1990AB EQ 4MG BASE N19668 003 Nov 02, 1990AB EQ 8MG BASE N19668 004 Nov 02, 1990DOXAZOSIN MESYLATEAB ACTAVIS ELIZABETH EQ 1MG BASE N75574 001 Oct 18, 2000AB EQ 2MG BASE N75574 002 Oct 18, 2000AB EQ 4MG BASE N75574 003 Oct 18, 2000AB EQ 8MG BASE N75574 004 Oct 18, 2000AB CLONMEL HLTHCARE EQ 1MG BASE N76161 001 Jun 10, 2004AB EQ 2MG BASE N76161 002 Jun 10, 2004AB EQ 4MG BASE N76161 003 Jun 10, 2004AB EQ 8MG BASE N76161 004 Jun 10, 2004AB GENPHARM EQ 1MG BASE N75466 001 Oct 18, 2000AB EQ 2MG BASE N75466 002 Oct 18, 2000AB EQ 4MG BASE N75466 003 Oct 18, 2000AB EQ 8MG BASE N75466 004 Oct 18, 2000AB IVAX PHARMS EQ 1MG BASE N75453 001 Oct 18, 2000AB EQ 2MG BASE N75453 002 Oct 18, 2000AB EQ 4MG BASE N75453 003 Oct 18, 2000AB EQ 8MG BASE N75453 004 Oct 18, 2000AB KV PHARM EQ 1MG BASE N75609 001 Oct 18, 2000AB EQ 2MG BASE N75609 002 Oct 18, 2000AB EQ 4MG BASE N75609 003 Oct 18, 2000AB EQ 8MG BASE N75609 004 Oct 18, 2000AB MYLAN EQ 1MG BASE N75509 001 Oct 19, 2000AB EQ 2MG BASE N75509 002 Oct 19, 2000AB EQ 4MG BASE N75509 003 Oct 19, 2000AB EQ 8MG BASE N75509 004 Oct 19, 2000AB PLIVA EQ 1MG BASE N75750 001 Jun 08, 2001AB EQ 2MG BASE N75750 002 Jun 08, 2001AB EQ 4MG BASE N75750 003 Jun 08, 2001AB EQ 8MG BASE N75750 004 Jun 08, 2001AB SANDOZ EQ 1MG BASE N75432 001 Oct 18, 2000AB EQ 1MG BASE N75646 001 Oct 18, 2000AB EQ 2MG BASE N75432 002 Oct 18, 2000AB EQ 2MG BASE N75646 002 Oct 18, 2000AB EQ 4MG BASE N75432 003 Oct 18, 2000AB EQ 4MG BASE N75646 003 Oct 18, 2000

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 134 (of 370)

DOXAZOSIN MESYLATE

TABLET; ORAL

DOXAZOSIN MESYLATE

<u>AB</u>	<u>SANDOZ</u>	<u>EQ 8MG BASE</u>	<u>N75432</u>	<u>004</u>	Oct 18, 2000
<u>AB</u>		<u>EQ 8MG BASE</u>	<u>N75646</u>	<u>004</u>	Oct 18, 2000
<u>AB</u>	<u>TEVA</u>	<u>EQ 1MG BASE</u>	<u>N75353</u>	<u>001</u>	Jan 12, 2001
<u>AB</u>		<u>EQ 1MG BASE</u>	<u>N75536</u>	<u>001</u>	Oct 18, 2000
<u>AB</u>		<u>EQ 2MG BASE</u>	<u>N75353</u>	<u>002</u>	Jan 12, 2001
<u>AB</u>		<u>EQ 2MG BASE</u>	<u>N75536</u>	<u>002</u>	Oct 18, 2000
<u>AB</u>		<u>EQ 4MG BASE</u>	<u>N75353</u>	<u>003</u>	Jan 12, 2001
<u>AB</u>		<u>EQ 4MG BASE</u>	<u>N75536</u>	<u>003</u>	Oct 18, 2000
<u>AB</u>		<u>EQ 8MG BASE</u>	<u>N75353</u>	<u>004</u>	Jan 12, 2001
<u>AB</u>		<u>EQ 8MG BASE</u>	<u>N75536</u>	<u>004</u>	Oct 18, 2000
<u>AB</u>	<u>TORPHARM</u>	<u>EQ 1MG BASE</u>	<u>N75580</u>	<u>001</u>	Oct 18, 2000
<u>AB</u>		<u>EQ 2MG BASE</u>	<u>N75580</u>	<u>002</u>	Oct 18, 2000
<u>AB</u>		<u>EQ 4MG BASE</u>	<u>N75580</u>	<u>003</u>	Oct 18, 2000
<u>AB</u>		<u>EQ 8MG BASE</u>	<u>N75580</u>	<u>004</u>	Oct 18, 2000
<u>AB</u>	<u>WATSON LABS</u>	<u>EQ 1MG BASE</u>	<u>N75426</u>	<u>001</u>	Oct 18, 2000
<u>AB</u>		<u>EQ 2MG BASE</u>	<u>N75426</u>	<u>002</u>	Oct 18, 2000
<u>AB</u>		<u>EQ 4MG BASE</u>	<u>N75426</u>	<u>003</u>	Oct 18, 2000
<u>AB</u>		<u>EQ 8MG BASE</u>	<u>N75426</u>	<u>004</u>	Oct 18, 2000

TABLET, EXTENDED RELEASE; ORAL

CARDURA XL

PFIZER

+

EQ 4MG BASE

N21269 001

Feb 22, 2005

EQ 8MG BASE

N21269 002

Feb 22, 2005

DOXEPIN HYDROCHLORIDE

CAPSULE; ORAL

DOXEPIN HYDROCHLORIDE

<u>AB</u>	<u>MYLAN</u>	<u>EQ 10MG BASE</u>	<u>N70789</u>	<u>001</u>	May 13, 1986
<u>AB</u>		<u>EQ 25MG BASE</u>	<u>N70790</u>	<u>001</u>	May 13, 1986
<u>AB</u>		<u>EQ 50MG BASE</u>	<u>N70791</u>	<u>001</u>	May 13, 1986
<u>AB</u>		<u>EQ 75MG BASE</u>	<u>N70792</u>	<u>001</u>	May 13, 1986
<u>AB</u>		<u>EQ 100MG BASE</u>	<u>N70793</u>	<u>001</u>	May 13, 1986
<u>AB</u>	<u>PAR PHARM</u>	<u>EQ 10MG BASE</u>	<u>N71697</u>	<u>001</u>	Nov 09, 1987
<u>AB</u>		<u>EQ 25MG BASE</u>	<u>N71437</u>	<u>001</u>	Nov 09, 1987
<u>AB</u>		<u>EQ 50MG BASE</u>	<u>N71595</u>	<u>001</u>	Nov 09, 1987
<u>AB</u>		<u>EQ 75MG BASE</u>	<u>N71608</u>	<u>001</u>	Nov 09, 1987
<u>AB</u>		<u>EQ 100MG BASE</u>	<u>N71422</u>	<u>001</u>	Nov 09, 1987
<u>AB</u>		<u>EQ 150MG BASE</u>	<u>N71669</u>	<u>001</u>	Nov 09, 1987
<u>AB</u>	<u>SANDOZ</u>	<u>EQ 10MG BASE</u>	<u>N71487</u>	<u>001</u>	Mar 02, 1987
<u>AB</u>	<u>WATSON LABS</u>	<u>EQ 10MG BASE</u>	<u>N71485</u>	<u>001</u>	Apr 30, 1987
<u>AB</u>		<u>EQ 10MG BASE</u>	<u>N72985</u>	<u>001</u>	Mar 29, 1991
<u>AB</u>		<u>EQ 25MG BASE</u>	<u>N71486</u>	<u>001</u>	Apr 30, 1987
<u>AB</u>		<u>EQ 25MG BASE</u>	<u>N72986</u>	<u>001</u>	Mar 29, 1991
<u>AB</u>		<u>EQ 50MG BASE</u>	<u>N71238</u>	<u>001</u>	Apr 30, 1987
<u>AB</u>		<u>EQ 50MG BASE</u>	<u>N72987</u>	<u>001</u>	Mar 29, 1991
<u>AB</u>		<u>EQ 75MG BASE</u>	<u>N71326</u>	<u>001</u>	Apr 30, 1987
<u>AB</u>		<u>EQ 100MG BASE</u>	<u>N71239</u>	<u>001</u>	Apr 30, 1987
	<u>SINEQUAN</u>				
<u>AB</u>	<u>PFIZER</u>	<u>EQ 10MG BASE</u>	<u>N16798</u>	<u>003</u>	
<u>AB</u>	<u>+</u>	<u>EQ 25MG BASE</u>	<u>N16798</u>	<u>001</u>	
<u>AB</u>		<u>EQ 50MG BASE</u>	<u>N16798</u>	<u>002</u>	
<u>AB</u>		<u>EQ 75MG BASE</u>	<u>N16798</u>	<u>006</u>	
<u>AB</u>	<u>+</u>	<u>EQ 100MG BASE</u>	<u>N16798</u>	<u>005</u>	
<u>AB</u>		<u>EQ 150MG BASE</u>	<u>N16798</u>	<u>007</u>	

CONCENTRATE; ORAL

DOXEPIN HYDROCHLORIDE

<u>AA</u>	<u>MORTON GROVE</u>	<u>EQ 10MG BASE/ML</u>	<u>N71918</u>	<u>001</u>	Jul 20, 1988
<u>AA</u>	<u>PHARM ASSOC</u>	<u>EQ 10MG BASE/ML</u>	<u>N75924</u>	<u>001</u>	Jan 15, 2004

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 135 (of 370)

DOXEPIN HYDROCHLORIDE

CONCENTRATE; ORAL

DOXEPIN HYDROCHLORIDE

<u>AA</u>	SILARX	<u>EQ 10MG BASE/ML</u>	<u>N74721</u>	<u>001</u>	Dec 29, 1998
<u>AA</u>	TEVA PHARMS	<u>EQ 10MG BASE/ML</u>	<u>N71609</u>	<u>001</u>	Nov 09, 1987
<u>SINEQUAN</u>					
<u>AA</u>	+ PFIZER	<u>EQ 10MG BASE/ML</u>	<u>N17516</u>	<u>001</u>	
CREAM; TOPICAL					
ZONALON					
	+ BRADLEY PHARMS	5%	N20126	001	Apr 01, 1994

DOXERCALCIFEROL

CAPSULE; ORAL

HECTOROL

	GENZYME	0.5UGM	N20862	002	Apr 23, 2004
	+	2.5UGM	N20862	001	Jun 09, 1999

INJECTABLE; INJECTION

HECTOROL

	+ GENZYME	2UGM/ML	N21027	001	Apr 06, 2000
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DOXORUBICIN HYDROCHLORIDE

INJECTABLE; INJECTION

ADRIAMYCIN PFS

<u>AP</u>	+ PHARMACIA AND UPJOHN	<u>2MG/ML</u>	<u>N50629</u>	<u>001</u>	Dec 23, 1987
<u>AP</u>		<u>2MG/ML</u>	<u>N63165</u>	<u>001</u>	Jan 30, 1991
<u>AP</u>	+	<u>200MG/100ML</u>	<u>N50629</u>	<u>002</u>	May 03, 1988
<u>AP</u>		<u>200MG/100ML</u>	<u>N63165</u>	<u>002</u>	Jan 30, 1991

DOXORUBICIN HYDROCHLORIDE

<u>AP</u>	ABRAXIS PHARM	<u>2MG/ML</u>	<u>N63277</u>	<u>001</u>	Oct 26, 1995
<u>AP</u>	BEDFORD	<u>2MG/ML</u>	<u>N62975</u>	<u>001</u>	Mar 17, 1989
<u>AP</u>		<u>10MG/VIAL</u>	<u>N62921</u>	<u>001</u>	Mar 17, 1989
<u>AP</u>		<u>20MG/VIAL</u>	<u>N62921</u>	<u>002</u>	Mar 17, 1989
<u>AP</u>		<u>50MG/VIAL</u>	<u>N62921</u>	<u>003</u>	Mar 17, 1989
<u>AP</u>		<u>200MG/100ML</u>	<u>N64097</u>	<u>001</u>	Sep 13, 1994
<u>AP</u>	PHARMACHEMIE	<u>2MG/ML</u>	<u>N63336</u>	<u>001</u>	Feb 28, 1995
<u>AP</u>		<u>10MG/VIAL</u>	<u>N63097</u>	<u>001</u>	May 21, 1990
<u>AP</u>		<u>20MG/VIAL</u>	<u>N63097</u>	<u>002</u>	May 21, 1990
<u>AP</u>		<u>50MG/VIAL</u>	<u>N63097</u>	<u>003</u>	May 21, 1990
<u>AP</u>		<u>200MG/100ML</u>	<u>N63336</u>	<u>004</u>	Feb 28, 1995
<u>AP</u>	SICOR PHARMS	<u>2MG/ML</u>	<u>N64140</u>	<u>001</u>	Jul 28, 1995
<u>AP</u>		<u>200MG/100ML</u>	<u>N64140</u>	<u>002</u>	Jul 28, 1995
<u>RUBEX</u>					
<u>AP</u>	BRISTOL MYERS SQUIBB	<u>10MG/VIAL</u>	<u>N62926</u>	<u>001</u>	Apr 13, 1989
<u>AP</u>		<u>50MG/VIAL</u>	<u>N62926</u>	<u>002</u>	Apr 13, 1989
		100MG/VIAL	N62926	003	Apr 13, 1989

INJECTABLE, LIPOSOMAL; INJECTION

DOXIL

	+ ALZA	2MG/ML	N50718	001	Nov 17, 1995
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DOXYCYCLINE

CAPSULE; ORAL

DOXYCYCLINE

<u>AB</u>	PAR PHARM	<u>EQ 50MG BASE</u>	<u>N65055</u>	<u>001</u>	Dec 01, 2000
<u>AB</u>		<u>EQ 100MG BASE</u>	<u>N65055</u>	<u>002</u>	Dec 01, 2000
<u>AB</u>	RANBAXY	<u>EQ 50MG BASE</u>	<u>N65053</u>	<u>001</u>	Nov 22, 2000
<u>AB</u>		<u>EQ 100MG BASE</u>	<u>N65053</u>	<u>002</u>	Nov 22, 2000
		EQ 75MG BASE	N65053	003	Sep 10, 2003
<u>AB</u>	SANDOZ	<u>EQ 50MG BASE</u>	<u>N65032</u>	<u>001</u>	Jun 30, 2000
<u>AB</u>		<u>EQ 100MG BASE</u>	<u>N65032</u>	<u>002</u>	Jun 30, 2000

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 136 (of 370)

DOXYCYCLINE

CAPSULE; ORAL

DOXYCYCLINE

<u>AB</u>	WATSON LABS	<u>EQ 50MG BASE</u>	<u>N65041</u>	<u>001</u>	Apr 28, 2000
<u>AB</u>		<u>EQ 100MG BASE</u>	<u>N65041</u>	<u>002</u>	Apr 28, 2000

MONODOX

<u>AB</u>	OCLASSEN	<u>EQ 50MG BASE</u>	<u>N50641</u>	<u>002</u>	Feb 10, 1992
<u>AB</u>	+	<u>EQ 100MG BASE</u>	<u>N50641</u>	<u>001</u>	Dec 29, 1989

CAPSULE, DELAYED RELEASE; ORAL

ORACEA

+	COLLAGENEX PHARMS	40MG	N50805	001	May 26, 2006
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FOR SUSPENSION; ORAL

VIBRAMYCIN

+	PFIZER	EQ 25MG BASE/5ML	N50006	001	
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TABLET; ORAL

DOXYCYCLINE

<u>AB</u>	LANNETT	<u>EQ 50MG BASE</u>	<u>N65285</u>	<u>001</u>	Dec 08, 2005
<u>AB</u>		<u>EQ 100MG BASE</u>	<u>N65285</u>	<u>002</u>	Dec 08, 2005
<u>AB</u>	MYLAN	<u>EQ 50MG BASE</u>	<u>N65377</u>	<u>001</u>	Nov 07, 2006
<u>AB</u>		<u>EQ 75MG BASE</u>	<u>N65377</u>	<u>002</u>	Nov 07, 2006
<u>AB</u>		<u>EQ 100MG BASE</u>	<u>N65377</u>	<u>003</u>	Nov 07, 2006
<u>AB</u>	PAR PHARM	<u>EQ 50MG BASE</u>	<u>N65070</u>	<u>001</u>	Dec 15, 2000
<u>AB</u>		<u>EQ 75MG BASE</u>	<u>N65070</u>	<u>003</u>	Dec 30, 2002
<u>AB</u>		<u>EQ 100MG BASE</u>	<u>N65070</u>	<u>002</u>	Dec 15, 2000
<u>AB</u>	+	EQ 150MG BASE	N65070	004	Jul 14, 2005
<u>AB</u>	RANBAXY	<u>EQ 50MG BASE</u>	<u>N65356</u>	<u>001</u>	May 31, 2006
<u>AB</u>		<u>EQ 75MG BASE</u>	<u>N65356</u>	<u>002</u>	May 31, 2006
<u>AB</u>		<u>EQ 100MG BASE</u>	<u>N65356</u>	<u>003</u>	May 31, 2006
<u>AB</u>	SANDOZ	<u>EQ 50MG BASE</u>	<u>N65353</u>	<u>001</u>	Nov 27, 2006
<u>AB</u>		<u>EQ 75MG BASE</u>	<u>N65353</u>	<u>002</u>	Nov 27, 2006
<u>AB</u>		<u>EQ 100MG BASE</u>	<u>N65353</u>	<u>003</u>	Nov 27, 2006

DOXYCYCLINE CALCIUM

SUSPENSION; ORAL

VIBRAMYCIN

+	PFIZER	EQ 50MG BASE/5ML	N50480	001	
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DOXYCYCLINE HYCLATE

CAPSULE; ORAL

DOXYCYCLINE HYCLATE

<u>AB</u>	IVAX PHARMS	<u>EQ 50MG BASE</u>	<u>N62500</u>	<u>001</u>	Sep 11, 1984
<u>AB</u>		<u>EQ 100MG BASE</u>	<u>N62500</u>	<u>002</u>	Sep 11, 1984
<u>AB</u>	MUTUAL PHARM	<u>EQ 50MG BASE</u>	<u>N62675</u>	<u>001</u>	Jul 10, 1986
<u>AB</u>		<u>EQ 100MG BASE</u>	<u>N62676</u>	<u>001</u>	Jul 10, 1986
<u>AB</u>	MYLAN	<u>EQ 50MG BASE</u>	<u>N62337</u>	<u>001</u>	Mar 29, 1982
<u>AB</u>		<u>EQ 100MG BASE</u>	<u>N62337</u>	<u>002</u>	Mar 29, 1982
<u>AB</u>	WATSON LABS	<u>EQ 50MG BASE</u>	<u>N61717</u>	<u>001</u>	
<u>AB</u>		<u>EQ 50MG BASE</u>	<u>N62031</u>	<u>002</u>	Oct 13, 1982
<u>AB</u>		<u>EQ 100MG BASE</u>	<u>N61717</u>	<u>002</u>	
<u>AB</u>		<u>EQ 100MG BASE</u>	<u>N62031</u>	<u>001</u>	
<u>AB</u>	WEST WARD	<u>EQ 50MG BASE</u>	<u>N62396</u>	<u>002</u>	Nov 07, 1984
<u>AB</u>		<u>EQ 100MG BASE</u>	<u>N62396</u>	<u>001</u>	May 07, 1984
<u>AB</u>	+	EQ 20MG BASE	N65103	001	May 13, 2005
	<u>VIBRAMYCIN</u>				
<u>AB</u>	PFIZER	<u>EQ 50MG BASE</u>	<u>N50007</u>	<u>001</u>	
<u>AB</u>	+	<u>EQ 100MG BASE</u>	<u>N50007</u>	<u>002</u>	

CAPSULE, DELAYED RELEASE; ORAL

DOXYCYCLINE HYCLATE

	SANDOZ	EQ 75MG BASE	N65281	001	Dec 21, 2005
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27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 137 (of 370)

DOXYCYCLINE HYCLATE

CAPSULE, DELAYED RELEASE; ORAL

DOXYCYCLINE HYCLATE

+	SANDOZ	EQ 100MG BASE	N65281	002	Dec 21, 2005
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INJECTABLE; INJECTION

DOXY 100

<u>AP</u>	+	ABRAXIS PHARM	<u>EQ 100MG BASE/VIAL</u>	<u>N62475</u>	<u>001</u>	Dec 09, 1983
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DOXY 200

+	ABRAXIS PHARM	EQ 200MG BASE/VIAL	N62475	002	Dec 09, 1983
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DOXYCYCLINE

<u>AP</u>	+	BEDFORD	<u>EQ 100MG BASE/VIAL</u>	<u>N62569</u>	<u>001</u>	Mar 09, 1988
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LIQUID, EXTENDED RELEASE; PERIODONTAL

ATRIDOX

+	QLT USA	EQ 10% W/W	N50751	001	Sep 03, 1998
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TABLET; ORAL

DOXYCYCLINE HYCLATE

<u>AB</u>		COREPHARMA	<u>EQ 20MG BASE</u>	<u>N65182</u>	<u>001</u>	May 13, 2005
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<u>AB</u>		IVAX PHARMS	<u>EQ 20MG BASE</u>	<u>N65163</u>	<u>001</u>	May 13, 2005
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<u>AB</u>			<u>EQ 100MG BASE</u>	<u>N62505</u>	<u>001</u>	Sep 11, 1984
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<u>AB</u>		LANNETT	<u>EQ 20MG BASE</u>	<u>N65277</u>	<u>001</u>	Nov 10, 2005
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<u>AB</u>		MUTUAL PHARM	<u>EQ 100MG BASE</u>	<u>N62677</u>	<u>001</u>	Jul 10, 1986
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<u>AB</u>		MUTUAL PHARMA	<u>EQ 20MG BASE</u>	<u>N65134</u>	<u>001</u>	May 13, 2005
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<u>AB</u>		MYLAN	<u>EQ 100MG BASE</u>	<u>N62432</u>	<u>001</u>	Feb 15, 1983
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<u>AB</u>		PAR PHARM	<u>EQ 20MG BASE</u>	<u>N65287</u>	<u>001</u>	Feb 28, 2006
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<u>AB</u>		TRUXTON INC	<u>EQ 100MG BASE</u>	<u>N62269</u>	<u>002</u>	Nov 08, 1982
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<u>AB</u>		VINTAGE PHARMS	<u>EQ 100MG BASE</u>	<u>N62538</u>	<u>001</u>	Apr 07, 1986
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<u>AB</u>		WATSON LABS	<u>EQ 100MG BASE</u>	<u>N62421</u>	<u>001</u>	Feb 02, 1983
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<u>AB</u>		WEST WARD	<u>EQ 100MG BASE</u>	<u>N65095</u>	<u>001</u>	Jul 02, 2003
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PERIOSTAT

<u>AB</u>	+	COLLAGENEX PHARMS	<u>EQ 20MG BASE</u>	<u>N50783</u>	<u>001</u>	Feb 02, 2001
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VIBRA-TABS

<u>AB</u>	+	PFIZER	<u>EQ 100MG BASE</u>	<u>N50533</u>	<u>001</u>	
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TABLET, DELAYED RELEASE; ORAL

DORYX

	MAYNE PHARMA INTL	EQ 75MG BASE	N50795	001	May 06, 2005
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+		EQ 100MG BASE	N50795	002	May 06, 2005
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DRONABINOL

CAPSULE; ORAL

MARINOL

	UNIMED	2.5MG	N18651	001	May 31, 1985
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		5MG	N18651	002	May 31, 1985
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+		10MG	N18651	003	May 31, 1985
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DROPERIDOL

INJECTABLE; INJECTION

DROPERIDOL

<u>AP</u>		HOSPIRA	<u>2.5MG/ML</u>	<u>N71981</u>	<u>001</u>	Feb 29, 1988
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<u>AP</u>			<u>2.5MG/ML</u>	<u>N72272</u>	<u>001</u>	Aug 31, 1995
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<u>AP</u>		LUITPOLD	<u>2.5MG/ML</u>	<u>N72123</u>	<u>001</u>	Oct 24, 1988
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<u>AP</u>			<u>2.5MG/ML</u>	<u>N72335</u>	<u>001</u>	Oct 24, 1988
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INAPSINE

<u>AP</u>	+	AKORN	<u>2.5MG/ML</u>	<u>N16796</u>	<u>001</u>	
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DROSPIRENONE; ESTRADIOL

TABLET; ORAL

ANGELIQ

+	BERLEX	0.5MG;1MG	N21355	002	Sep 28, 2005
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PRESCRIPTION DRUG PRODUCT LIST

3 - 138 (of 370)

DROSPIRENONE; ETHINYL ESTRADIOL

TABLET; ORAL

YAZ

+ BERLEX	3MG;0.02MG	N21676 001	Mar 16, 2006
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TABLET; ORAL-28

YASMIN

+ BERLEX	3MG;0.03MG	N21098 001	May 11, 2001
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DULOXETINE HYDROCHLORIDE

CAPSULE, DELAYED REL PELLETS; ORAL

CYMBALTA

LILLY

EQ 20MG BASE

N21427 001 Aug 03, 2004

EQ 30MG BASE

N21427 002 Aug 03, 2004

+	EQ 60MG BASE	N21427 004	Aug 03, 2004
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DUTASTERIDE

CAPSULE; ORAL

AVODART

+ GLAXOSMITHKLINE	0.5MG	N21319 001	Nov 20, 2001
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DYPHYLLINE

TABLET; ORAL

LUFYLLIN

MEDPOINTE PHARM HLC 200MG

N84566 001

+	400MG	N84566 002	
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ECHOTHIOPHATE IODIDE

FOR SOLUTION; OPHTHALMIC

PHOSPHOLINE IODIDE

+ AYERST	0.125%	N11963 001	
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ECONAZOLE NITRATE

CREAM; TOPICAL

ECONAZOLE NITRATE

<u>AB</u>	ALTANA	1%	<u>N76075</u> <u>001</u>	Nov 26, 2002
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<u>AB</u>	HEALTHPOINT	1%	<u>N76574</u> <u>001</u>	Dec 17, 2004
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<u>AB</u>	PERRIGO NEW YORK	1%	<u>N76479</u> <u>001</u>	Jun 23, 2004
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<u>AB</u>	TARO	1%	<u>N76005</u> <u>001</u>	Nov 26, 2002
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SPECTAZOLE

<u>AB</u>	+ JOHNSON AND JOHNSON	1%	<u>N18751</u> <u>001</u>	Dec 23, 1982
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EDETATE CALCIUM DISODIUM

INJECTABLE; INJECTION

CALCIUM DISODIUM VERSENATE

+ 3M	200MG/ML	N08922 001	
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EDETATE DISODIUM

INJECTABLE; INJECTION

EDETATE DISODIUM

<u>AP</u>	APOTEX INC	150MG/ML	<u>N40376</u> <u>001</u>	Nov 04, 2002
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<u>AP</u>	BIONICHE (CANADA)	150MG/ML	<u>N40437</u> <u>001</u>	Jul 09, 2002
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ENDRATE

<u>AP</u>	+ HOSPIRA	150MG/ML	<u>N11355</u> <u>001</u>	
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EDROPHONIUM CHLORIDE

INJECTABLE; INJECTION

EDROPHONIUM CHLORIDE

<u>AP</u>	HOSPIRA	10MG/ML	<u>N40131</u> <u>001</u>	Feb 24, 1998
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27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 139 (of 370)

EDROPHONIUM CHLORIDE

INJECTABLE; INJECTION

ENLON

<u>AP</u>	BAXTER HLTHCARE CORP	<u>10MG/ML</u>	<u>N88873</u>	<u>001</u>	Aug 06, 1985
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TENSILON

<u>AP</u>	+ VALEANT PHARM INTL	<u>10MG/ML</u>	<u>N07959</u>	<u>001</u>	
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TENSILON PRESERVATIVE FREE

<u>AP</u>	+ VALEANT PHARM INTL	<u>10MG/ML</u>	<u>N07959</u>	<u>002</u>	
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EFAVIRENZ

CAPSULE; ORAL

SUSTIVA

	BRISTOL MYERS SQUIBB	50MG	N20972	001	Sep 17, 1998
		100MG	N20972	002	Sep 17, 1998
+		200MG	N20972	003	Sep 17, 1998

TABLET; ORAL

SUSTIVA

+	BRISTOL MYERS SQUIBB	600MG	N21360	002	Feb 01, 2002
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EFAVIRENZ; EMTRICITABINE; TENOFOVIR DISOPROXIL FUMARATE

TABLET; ORAL

ATRIPLA

+	GILEAD	600MG;200MG;300MG	N21937	001	Jul 12, 2006
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EFLORNITHINE HYDROCHLORIDE

CREAM; TOPICAL

VANIQA

+	SKINMEDICA	13.9%	N21145	001	Jul 27, 2000
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ELETRIPTAN HYDROBROMIDE

TABLET; ORAL

RELPAX

	PFIZER IRELAND	EQ 20MG BASE	N21016	001	Dec 26, 2002
+		EQ 40MG BASE	N21016	002	Dec 26, 2002

EMEDASTINE DIFUMARATE

SOLUTION/DROPS; OPHTHALMIC

+	ALCON	0.05%	N20706	001	Dec 29, 1997
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EMTRICITABINE

CAPSULE; ORAL

EMTRIVA

+	GILEAD	200MG	N21500	001	Jul 02, 2003
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SOLUTION; ORAL

EMTRIVA

+	GILEAD	10MG/ML	N21896	001	Sep 28, 2005
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EMTRICITABINE; TENOFOVIR DISOPROXIL FUMARATE

TABLET; ORAL

TRUVADA

+	GILEAD	200MG;300MG	N21752	001	Aug 02, 2004
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ENALAPRIL MALEATE

TABLET; ORAL

ENALAPRIL MALEATE

<u>AB</u>	GENPHARM	<u>2.5MG</u>	<u>N75472</u>	<u>001</u>	Aug 22, 2000
<u>AB</u>		<u>5MG</u>	<u>N75472</u>	<u>002</u>	Aug 22, 2000

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 140 (of 370)

ENALAPRIL MALEATE

TABLET; ORAL

ENALAPRIL MALEATE

<u>AB</u>	GENPHARM	<u>10MG</u>	<u>N75472</u>	<u>003</u>	Aug 22, 2000
<u>AB</u>		<u>20MG</u>	<u>N75472</u>	<u>004</u>	Aug 22, 2000
<u>AB</u>	KRKA DD NOVO MESTO	<u>2.5MG</u>	<u>N75370</u>	<u>001</u>	Aug 22, 2000
<u>AB</u>		<u>5MG</u>	<u>N75370</u>	<u>002</u>	Aug 22, 2000
<u>AB</u>		<u>10MG</u>	<u>N75369</u>	<u>001</u>	Aug 22, 2000
<u>AB</u>		<u>20MG</u>	<u>N75369</u>	<u>002</u>	Aug 22, 2000
<u>AB</u>	LEK PHARMS	<u>2.5MG</u>	<u>N75496</u>	<u>001</u>	Aug 22, 2000
<u>AB</u>		<u>5MG</u>	<u>N75496</u>	<u>002</u>	Aug 22, 2000
<u>AB</u>		<u>10MG</u>	<u>N75459</u>	<u>001</u>	Aug 22, 2000
<u>AB</u>		<u>20MG</u>	<u>N75459</u>	<u>002</u>	Aug 22, 2000
<u>AB</u>	MYLAN	<u>2.5MG</u>	<u>N75480</u>	<u>001</u>	Aug 22, 2000
<u>AB</u>		<u>5MG</u>	<u>N75480</u>	<u>002</u>	Aug 22, 2000
<u>AB</u>		<u>10MG</u>	<u>N75480</u>	<u>003</u>	Aug 22, 2000
<u>AB</u>		<u>20MG</u>	<u>N75480</u>	<u>004</u>	Aug 22, 2000
<u>AB</u>	RANBAXY	<u>2.5MG</u>	<u>N75556</u>	<u>001</u>	Aug 22, 2000
<u>AB</u>		<u>5MG</u>	<u>N75556</u>	<u>002</u>	Aug 22, 2000
<u>AB</u>		<u>10MG</u>	<u>N75556</u>	<u>003</u>	Aug 22, 2000
<u>AB</u>		<u>20MG</u>	<u>N75556</u>	<u>004</u>	Aug 22, 2000
<u>AB</u>	SANDOZ	<u>2.5MG</u>	<u>N75048</u>	<u>001</u>	Aug 22, 2000
<u>AB</u>		<u>2.5MG</u>	<u>N75621</u>	<u>001</u>	Aug 22, 2000
<u>AB</u>		<u>5MG</u>	<u>N75048</u>	<u>002</u>	Aug 22, 2000
<u>AB</u>		<u>5MG</u>	<u>N75621</u>	<u>002</u>	Aug 22, 2000
<u>AB</u>		<u>10MG</u>	<u>N75048</u>	<u>003</u>	Aug 22, 2000
<u>AB</u>		<u>10MG</u>	<u>N75621</u>	<u>003</u>	Aug 22, 2000
<u>AB</u>		<u>20MG</u>	<u>N75048</u>	<u>004</u>	Aug 22, 2000
<u>AB</u>		<u>20MG</u>	<u>N75621</u>	<u>004</u>	Aug 22, 2000
<u>AB</u>	TARO	<u>2.5MG</u>	<u>N75657</u>	<u>001</u>	Jan 23, 2001
<u>AB</u>		<u>5MG</u>	<u>N75657</u>	<u>002</u>	Jan 23, 2001
<u>AB</u>		<u>10MG</u>	<u>N75657</u>	<u>003</u>	Jan 23, 2001
<u>AB</u>		<u>20MG</u>	<u>N75657</u>	<u>004</u>	Jan 23, 2001
<u>AB</u>	TEVA	<u>2.5MG</u>	<u>N75479</u>	<u>001</u>	Aug 22, 2000
<u>AB</u>		<u>5MG</u>	<u>N75479</u>	<u>002</u>	Aug 22, 2000
<u>AB</u>		<u>10MG</u>	<u>N75479</u>	<u>003</u>	Aug 22, 2000
<u>AB</u>		<u>20MG</u>	<u>N75479</u>	<u>004</u>	Aug 22, 2000
<u>AB</u>	TORPHARM	<u>2.5MG</u>	<u>N75178</u>	<u>002</u>	Mar 23, 2001
<u>AB</u>		<u>5MG</u>	<u>N75178</u>	<u>001</u>	Mar 23, 2001
<u>AB</u>		<u>10MG</u>	<u>N75178</u>	<u>003</u>	Mar 23, 2001
<u>AB</u>		<u>20MG</u>	<u>N75178</u>	<u>004</u>	Mar 23, 2001
<u>AB</u>	WATSON LABS	<u>2.5MG</u>	<u>N75501</u>	<u>001</u>	Aug 22, 2000
<u>AB</u>		<u>5MG</u>	<u>N75501</u>	<u>002</u>	Aug 22, 2000
<u>AB</u>		<u>10MG</u>	<u>N75501</u>	<u>003</u>	Aug 22, 2000
<u>AB</u>		<u>20MG</u>	<u>N75501</u>	<u>004</u>	Aug 22, 2000
<u>AB</u>	WOCKHARDT	<u>2.5MG</u>	<u>N75483</u>	<u>001</u>	Aug 22, 2000
<u>AB</u>		<u>5MG</u>	<u>N75483</u>	<u>002</u>	Aug 22, 2000
<u>AB</u>		<u>10MG</u>	<u>N75483</u>	<u>003</u>	Aug 22, 2000
<u>AB</u>		<u>20MG</u>	<u>N75483</u>	<u>004</u>	Aug 22, 2000
<u>VASOTEC</u>					
<u>AB</u>	BIOVAIL LABS INTL	<u>2.5MG</u>	<u>N18998</u>	<u>005</u>	Jul 26, 1988
<u>AB</u>		<u>5MG</u>	<u>N18998</u>	<u>001</u>	Dec 24, 1985
<u>AB</u>		<u>10MG</u>	<u>N18998</u>	<u>002</u>	Dec 24, 1985
<u>AB</u>	+	<u>20MG</u>	<u>N18998</u>	<u>003</u>	Dec 24, 1985

ENALAPRIL MALEATE; FELODIPINE

TABLET, EXTENDED RELEASE; ORAL

LEXXEL

+ ASTRAZENECA 5MG; 2.5MG

N20668 002 Oct 28, 1998

+ 5MG; 5MG

N20668 001 Dec 27, 1996

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 141 (of 370)

ENALAPRIL MALEATE; HYDROCHLOROTHIAZIDE

TABLET; ORAL

ENALAPRIL MALEATE AND HYDROCHLOROTHIAZIDE

<u>AB</u>	APOTEX INC	<u>5MG;12.5MG</u>	<u>N76486</u>	<u>001</u>	Oct 27, 2004
<u>AB</u>		<u>10MG;25MG</u>	<u>N76486</u>	<u>002</u>	Oct 27, 2004
<u>AB</u>	DR REDDYS LABS LTD	<u>5MG;12.5MG</u>	<u>N75909</u>	<u>001</u>	Oct 15, 2001
<u>AB</u>		<u>10MG;25MG</u>	<u>N75909</u>	<u>002</u>	Oct 15, 2001
<u>AB</u>	MYLAN	<u>5MG;12.5MG</u>	<u>N75624</u>	<u>001</u>	Sep 18, 2001
<u>AB</u>		<u>10MG;25MG</u>	<u>N75624</u>	<u>002</u>	Sep 18, 2001
<u>AB</u>	SANDOZ	<u>5MG;12.5MG</u>	<u>N76116</u>	<u>001</u>	Sep 19, 2001
<u>AB</u>		<u>10MG;25MG</u>	<u>N76116</u>	<u>002</u>	Sep 19, 2001
<u>AB</u>	TARO PHARM INDS	<u>5MG;12.5MG</u>	<u>N75788</u>	<u>001</u>	Sep 18, 2001
<u>AB</u>		<u>10MG;25MG</u>	<u>N75788</u>	<u>002</u>	Sep 18, 2001
<u>AB</u>	TEVA	<u>5MG;12.5MG</u>	<u>N75727</u>	<u>001</u>	Sep 18, 2001
<u>AB</u>		<u>10MG;25MG</u>	<u>N75727</u>	<u>002</u>	Sep 18, 2001
<u>VASERETIC</u>					
<u>AB</u>	BIOVAIL LABS INTL	<u>5MG;12.5MG</u>	<u>N19221</u>	<u>003</u>	Jul 12, 1995
<u>AB</u>	+	<u>10MG;25MG</u>	<u>N19221</u>	<u>001</u>	Oct 31, 1986

ENALAPRILAT

INJECTABLE; INJECTION

ENALAPRILAT

<u>AP</u>	BEDFORD	<u>1.25MG/ML</u>	<u>N75634</u>	<u>001</u>	Aug 22, 2000
<u>AP</u>	HOSPIRA	<u>1.25MG/ML</u>	<u>N75456</u>	<u>001</u>	Aug 22, 2000
<u>AP</u>		<u>1.25MG/ML</u>	<u>N75458</u>	<u>001</u>	Aug 22, 2000
<u>AP</u>	MAYNE PHARMA USA	<u>1.25MG/ML</u>	<u>N75571</u>	<u>001</u>	Aug 22, 2000
<u>AP</u>	SICOR PHARMS	<u>1.25MG/ML</u>	<u>N75578</u>	<u>001</u>	Aug 22, 2000
<u>VASOTEC</u>					
<u>AP</u>	+	<u>1.25MG/ML</u>	<u>N19309</u>	<u>001</u>	Feb 09, 1988

ENFLURANE

LIQUID; INHALATION

ENFLURANE

<u>AN</u>	MINRAD	<u>99.9%</u>	<u>N74396</u>	<u>001</u>	Jul 29, 1994
<u>ETHRANE</u>					
<u>AN</u>	+	<u>99.9%</u>	<u>N17087</u>	<u>001</u>	

ENFUVIRTIDE

INJECTABLE; SUBCUTANEOUS

FUZEON

+	ROCHE	90MG/VIAL	N21481	001	Mar 13, 2003
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ENOXAPARIN SODIUM

INJECTABLE; SUBCUTANEOUS

LOVENOX

	SANOFI AVENTIS US	300MG/3ML (100MG/ML)	N20164	009	Jan 23, 2003
	LOVENOX (PRESERVATIVE FREE)				
	SANOFI AVENTIS US	30MG/0.3ML (100MG/ML)	N20164	001	Mar 29, 1993
		40MG/0.4ML (100MG/ML)	N20164	002	Jan 30, 1998
		60MG/0.6ML (100MG/ML)	N20164	003	Mar 27, 1998
		80MG/0.8ML (100MG/ML)	N20164	004	Mar 27, 1998
+		100MG/ML (100MG/ML)	N20164	005	Mar 27, 1998
		120MG/0.8ML (150MG/ML)	N20164	007	Jun 02, 2000
		150MG/ML (150MG/ML)	N20164	008	Jun 02, 2000

ENTACAPONE

TABLET; ORAL

COMTAN

+	ORION	200MG	N20796	001	Oct 19, 1999
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27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 142 (of 370)

ENTECAVIR

SOLUTION; ORAL

BARACLUDE

+	BRISTOL MYERS SQUIBB	0.05MG/ML	N21798	001	Mar 29, 2005
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TABLET; ORAL

BARACLUDE

	BRISTOL MYERS SQUIBB	0.5MG	N21797	001	Mar 29, 2005
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+		1MG	N21797	002	Mar 29, 2005
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EPINASTINE HYDROCHLORIDE

SOLUTION/DROPS; OPHTHALMIC

+	ALLERGAN	0.05%	N21565	001	Oct 16, 2003
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EPINEPHRINE

INJECTABLE; IM-SC

TWINJECT

+	VERUS PHARMS	EQ 0.15MG /DELIVERY	N20800	002	May 28, 2004
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TWINJECT 0.3

+	VERUS PHARMS	EQ 0.3MG /DELIVERY	N20800	001	May 30, 2003
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INJECTABLE; INTRAMUSCULAR

EPIPEN

+	MERIDIAN MEDCL TECHN	0.3MG/DELIVERY	N19430	001	Dec 22, 1987
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EPIPEN JR.

+	MERIDIAN MEDCL TECHN	0.15MG/DELIVERY	N19430	002	Dec 22, 1987
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EPINEPHRINE BITARTRATE; LIDOCAINE HYDROCHLORIDE

INJECTABLE; INJECTION

LIGNOSPAN FORTE

+	DEPROCO	EQ 0.02MG BASE/ML;2%	N88389	001	Jan 22, 1985
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LIGNOSPAN STANDARD

+	DEPROCO	EQ 0.01MG BASE/ML;2%	N88390	001	Jan 22, 1985
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EPINEPHRINE BITARTRATE; PRILOCAINE HYDROCHLORIDE

INJECTABLE; INJECTION

CITANEST FORTE

+	DENTSPLY PHARM	0.005MG/ML;4%	N21383	001	
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EPINEPHRINE; LIDOCAINE HYDROCHLORIDE

INJECTABLE; INJECTION

LIDOCAINE HYDROCHLORIDE AND EPINEPHRINE

<u>AP</u>	EASTMAN KODAK	<u>0.01MG/ML;2%</u>	<u>N40057</u>	<u>002</u>	Feb 26, 1993
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<u>AP</u>		<u>0.02MG/ML;2%</u>	<u>N40057</u>	<u>001</u>	Feb 26, 1993
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<u>AP</u>	GRAHAM CHEM	<u>0.01MG/ML;2%</u>	<u>N80504</u>	<u>004</u>	Oct 19, 1983
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<u>AP</u>		<u>0.02MG/ML;2%</u>	<u>N80504</u>	<u>005</u>	Oct 19, 1983
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<u>AP</u>	HOSPIRA	<u>0.005MG/ML;0.5%</u>	<u>N89635</u>	<u>001</u>	Jun 21, 1988
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<u>AP</u>		<u>0.005MG/ML;1%</u>	<u>N89649</u>	<u>001</u>	Jun 21, 1988
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<u>AP</u>		<u>0.005MG/ML;1.5%</u>	<u>N88571</u>	<u>001</u>	Sep 13, 1985
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<u>AP</u>		<u>0.005MG/ML;1.5%</u>	<u>N89645</u>	<u>001</u>	Jun 21, 1988
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<u>AP</u>		<u>0.005MG/ML;1.5%</u>	<u>N89650</u>	<u>001</u>	Jun 21, 1988
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<u>AP</u>		<u>0.005MG/ML;2%</u>	<u>N89651</u>	<u>001</u>	Jun 21, 1988
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<u>AP</u>		<u>0.01MG/ML;1%</u>	<u>N89644</u>	<u>001</u>	Jun 21, 1988
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<u>AP</u>		<u>0.01MG/ML;2%</u>	<u>N89646</u>	<u>001</u>	Jun 21, 1988
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LIDOCAINE HYDROCHLORIDE W/ EPINEPHRINE

<u>AP</u>	DELL LABS	<u>0.01MG/ML;1%</u>	<u>N83389</u>	<u>001</u>	
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<u>AP</u>		<u>0.01MG/ML;2%</u>	<u>N83390</u>	<u>001</u>	
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LIDOCATON

<u>AP</u>	PHARMATON	<u>0.01MG/ML;2%</u>	<u>N84729</u>	<u>001</u>	Aug 17, 1983
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PRESCRIPTION DRUG PRODUCT LIST

3 - 143 (of 370)

EPINEPHRINE; LIDOCAINE HYDROCHLORIDE

INJECTABLE; INJECTION

OCTOCAINE

<u>AP</u>	SEPTODONT	<u>0.01MG/ML;2%</u>	<u>N84048</u>	<u>001</u>	
<u>AP</u>	+	<u>0.02MG/ML;2%</u>	<u>N84048</u>	<u>002</u>	

XYLOCAINE W/ EPINEPHRINE

<u>AP</u>	+ ABRAXIS BIOSCIENCE	<u>0.005MG/ML;0.5%</u>	<u>N06488</u>	<u>012</u>	
<u>AP</u>	+	<u>0.005MG/ML;1%</u>	<u>N06488</u>	<u>018</u>	Nov 13, 1986
<u>AP</u>	+	<u>0.005MG/ML;1.5%</u>	<u>N06488</u>	<u>017</u>	
<u>AP</u>	+	<u>0.005MG/ML;2%</u>	<u>N06488</u>	<u>019</u>	Nov 13, 1986
<u>AP</u>	+	<u>0.01MG/ML;1%</u>	<u>N06488</u>	<u>004</u>	
	+ DENTSPLY PHARM	0.01MG/ML;2%	N21381	001	
	+	0.02MG/ML;2%	N21381	002	

PATCH; IONTOPHORESIS, TOPICAL

LIDOSITE TOPICAL SYSTEM KIT

	+ VYTERIS	1.05MG/PATCH;100MG/PATCH	N21504	001	May 06, 2004
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SOLUTION; IONTOPHORESIS, TOPICAL

LIDOCAINE HYDROCHLORIDE AND EPINEPHRINE

	+ EMPI	0.01MG/ML;2%	N21486	001	Oct 26, 2004
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EPIRUBICIN HYDROCHLORIDE

INJECTABLE; INJECTION

ELLECE

	+ PFIZER INC	2MG/ML	N50778	001	Sep 15, 1999
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INJECTABLE; IV (INFUSION)

	+ MAYNE PHARMA USA	50MG/VIAL	N50807	001	Sep 15, 2006
	+	200MG/VIAL	N50807	002	Sep 15, 2006

EPLERENONE

TABLET; ORAL

INSPRA

	GD SEARLE LLC	25MG	N21437	001	Sep 27, 2002
	+	50MG	N21437	002	Sep 27, 2002

EPOPROSTENOL SODIUM

INJECTABLE; INJECTION

FLOLAN

	+ GLAXOSMITHKLINE	EQ 0.5MG BASE/VIAL	N20444	001	Sep 20, 1995
	+	EQ 1.5MG BASE/VIAL	N20444	002	Sep 20, 1995

EPROSARTAN MESYLATE

TABLET; ORAL

TEVETEN

	KOS LIFE	EQ 400MG BASE	N20738	005	Dec 22, 1997
	+	EQ 600MG BASE	N20738	006	May 27, 1999

EPROSARTAN MESYLATE; HYDROCHLOROTHIAZIDE

TABLET; ORAL

TEVETEN HCT

	KOS LIFE	600MG;12.5MG	N21268	001	Nov 01, 2001
	+	600MG;25MG	N21268	002	Nov 01, 2001

EPTIFIBATIDE

INJECTABLE; INJECTION

INTEGRILIN

	+ SCHERING	2MG/ML	N20718	001	May 18, 1998
	+	75MG/100ML	N20718	002	May 18, 1998

PRESCRIPTION DRUG PRODUCT LIST

3 - 144 (of 370)

ERGOCALCIFEROL

CAPSULE; ORAL

DRISDOL

<u>AA</u>	+ SANOFI AVENTIS US	<u>50,000 IU</u>	<u>N03444</u>	<u>001</u>	
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VITAMIN D

<u>AA</u>	BANNER PHARMACAPS	<u>50,000 IU</u>	<u>N80704</u>	<u>001</u>	
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ERGOLOID MESYLATES

TABLET; ORAL

ERGOLOID MESYLATES

<u>AB</u>	MUTUAL PHARM	<u>1MG</u>	<u>N81113</u>	<u>001</u>	Oct 31, 1991
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HYDERGINE

<u>AB</u>	+ NOVARTIS	<u>1MG</u>	<u>N17993</u>	<u>001</u>	
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TABLET; SUBLINGUAL

ERGOLOID MESYLATES

<u>AA</u>	WATSON LABS	<u>0.5MG</u>	<u>N87233</u>	<u>001</u>	
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GERIMAL

<u>AA</u>	WATSON LABS	<u>0.5MG</u>	<u>N86189</u>	<u>001</u>	
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<u>AA</u>		<u>1MG</u>	<u>N86188</u>	<u>001</u>	
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HYDERGINE

<u>AA</u>	+ NOVARTIS	<u>0.5MG</u>	<u>N09087</u>	<u>002</u>	
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<u>AA</u>	+	<u>1MG</u>	<u>N09087</u>	<u>001</u>	
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HYDROGENATED ERGOT ALKALOIDS

<u>AA</u>	IVAX PHARMS	<u>1MG</u>	<u>N87185</u>	<u>001</u>	
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ERGOTAMINE TARTRATE

TABLET; SUBLINGUAL

ERGOMAR

	+ HARVEST PHARMS	2MG	N87693	001	Feb 24, 1983
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ERLOTINIB HYDROCHLORIDE

TABLET; ORAL

TARCEVA

	OSI PHARMS	EQ 25MG BASE	N21743	001	Nov 18, 2004
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		EQ 100MG BASE	N21743	002	Nov 18, 2004
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+		EQ 150MG BASE	N21743	003	Nov 18, 2004
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ERTAPENEM SODIUM

INJECTABLE; INTRAMUSCULAR, IV (INFUSION)

	+ MERCK	EQ 1GM BASE/VIAL	N21337	001	Nov 21, 2001
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ERYTHROMYCIN

CAPSULE, DELAYED REL PELLETS; ORAL

ERYC

<u>AB</u>	+ MAYNE PHARMA USA	<u>250MG</u>	<u>N50536</u>	<u>001</u>	
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<u>AB</u>	WARNER CHILCOTT	<u>250MG</u>	<u>N62338</u>	<u>001</u>	
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ERYTHROMYCIN

<u>AB</u>	ABBOTT	<u>250MG</u>	<u>N62746</u>	<u>001</u>	Dec 22, 1986
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<u>AB</u>	BARR	<u>250MG</u>	<u>N63098</u>	<u>001</u>	May 04, 1989
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GEL; TOPICAL

E-GLADES

<u>AT</u>	GLADES PHARMS	<u>2%</u>	<u>N65009</u>	<u>001</u>	Mar 18, 2002
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ERYGEL

<u>AT</u>	+ MERZ PHARMS	<u>2%</u>	<u>N50617</u>	<u>001</u>	Oct 21, 1987
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ERYTHROMYCIN

<u>AT</u>	ALTANA	<u>2%</u>	<u>N64184</u>	<u>001</u>	Sep 30, 1997
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<u>AT</u>	STIEFEL	<u>2%</u>	<u>N63211</u>	<u>001</u>	Jan 29, 1993
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27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 145 (of 370)

ERYTHROMYCIN

LOTION; TOPICAL					
E-SOLVE 2					
	+ SYOSSET	2%	N62467	001	Jul 03, 1985
OINTMENT; OPHTHALMIC					
<u>ERYTHROMYCIN</u>					
<u>AT</u>	AKORN	<u>0.5%</u>	<u>N64030</u>	<u>001</u>	Jul 18, 1996
<u>AT</u>	BAUSCH AND LOMB	<u>0.5%</u>	<u>N64067</u>	<u>001</u>	Jul 29, 1994
<u>AT</u>	+ FOUGERA	<u>0.5%</u>	<u>N62447</u>	<u>001</u>	Sep 26, 1983
OINTMENT; TOPICAL					
AKNE-MYCIN					
	+ HEALTHPOINT	2%	N50584	001	Jan 10, 1985
SOLUTION; TOPICAL					
<u>A/T/S</u>					
<u>AT</u>	TARO PHARMS NORTH	<u>2%</u>	<u>N62405</u>	<u>001</u>	Nov 18, 1982
<u>ERYDERM</u>					
<u>AT</u>	ABBOTT	<u>2%</u>	<u>N62290</u>	<u>001</u>	
<u>ERYTHRA-DERM</u>					
<u>AT</u>	PADDOCK	<u>2%</u>	<u>N62687</u>	<u>001</u>	Feb 05, 1988
<u>ERYTHROMYCIN</u>					
<u>AT</u>	+ ALTANA	<u>2%</u>	<u>N64187</u>	<u>001</u>	Sep 30, 1997
<u>AT</u>	MORTON GROVE	<u>2%</u>	<u>N62825</u>	<u>001</u>	Oct 23, 1987
<u>AT</u>	PERRIGO NEW YORK	<u>2%</u>	<u>N63038</u>	<u>001</u>	Jan 11, 1991
SWAB; TOPICAL					
<u>C-SOLVE-2</u>					
<u>AT</u>	IVAX PHARMS	<u>2%</u>	<u>N62751</u>	<u>001</u>	Jul 30, 1993
<u>ERYCETTE</u>					
<u>AT</u>	+ J AND J	<u>2%</u>	<u>N50594</u>	<u>001</u>	Feb 15, 1985
<u>ERYTHROMYCIN</u>					
<u>AT</u>	ALTANA	<u>2%</u>	<u>N65320</u>	<u>001</u>	Jul 25, 2006
<u>AT</u>	STIEFEL	<u>2%</u>	<u>N64126</u>	<u>001</u>	Jul 03, 1996
<u>AT</u>		<u>2%</u>	<u>N64128</u>	<u>001</u>	Jul 03, 1996
TABLET; ORAL					
ERYTHROMYCIN					
	ABBOTT	250MG	N61621	001	
	+	500MG	N61621	002	
TABLET, COATED PARTICLES; ORAL					
PCE					
	ABBOTT	333MG	N50611	001	Sep 09, 1986
	+	500MG	N50611	002	Aug 22, 1990
TABLET, DELAYED RELEASE; ORAL					
<u>E-BASE</u>					
<u>AB</u>	BARR	<u>333MG</u>	<u>N63086</u>	<u>001</u>	May 15, 1990
<u>AB</u>		<u>500MG</u>	<u>N62999</u>	<u>001</u>	Nov 25, 1988
<u>E-MYCIN</u>					
<u>AB</u>	ABBOTT	<u>250MG</u>	<u>N60272</u>	<u>001</u>	
<u>AB</u>		<u>333MG</u>	<u>N60272</u>	<u>002</u>	
<u>ERY-TAB</u>					
<u>AB</u>	+ ABBOTT	<u>250MG</u>	<u>N62298</u>	<u>001</u>	
<u>AB</u>	+	<u>333MG</u>	<u>N62298</u>	<u>003</u>	Mar 29, 1982
<u>AB</u>	+	<u>500MG</u>	<u>N62298</u>	<u>002</u>	
<u>ERYTHROMYCIN ESTOLATE</u>					
SUSPENSION; ORAL					
ERYTHROMYCIN ESTOLATE					
	ALPHARMA US PHARMS	EQ 125MG BASE/5ML	N62353	001	Nov 18, 1982
	+	EQ 250MG BASE/5ML	N62409	001	Dec 16, 1982

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 146 (of 370)

ERYTHROMYCIN ETHYLSUCCINATE

GRANULE; ORAL

E.E.S.

<u>AB</u>	ABBOTT	<u>EQ 200MG BASE/5ML</u>	<u>N50207</u>	<u>001</u>	
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ERYPED

<u>AB</u>	ABBOTT	<u>EQ 200MG BASE/5ML</u>	<u>N50207</u>	<u>003</u>	Mar 30, 1987
+		<u>EQ 400MG BASE/5ML</u>	<u>N50207</u>	<u>002</u>	

ERYTHROMYCIN ETHYLSUCCINATE

<u>AB</u>	BARR	<u>EQ 200MG BASE/5ML</u>	<u>N62055</u>	<u>001</u>	
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SUSPENSION; ORAL

E.E.S. 200

<u>AB</u>	ABBOTT	<u>EQ 200MG BASE/5ML</u>	<u>N61639</u>	<u>001</u>	
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E.E.S. 400

<u>AB</u>	+	ABBOTT	<u>EQ 400MG BASE/5ML</u>	<u>N61639</u>	<u>002</u>
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ERYTHROMYCIN ETHYLSUCCINATE

<u>AB</u>	ALPHARMA US PHARMS	<u>EQ 200MG BASE/5ML</u>	<u>N62200</u>	<u>001</u>	
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<u>AB</u>		<u>EQ 400MG BASE/5ML</u>	<u>N62200</u>	<u>002</u>	
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PEDIAMYCIN

<u>AB</u>	ROSS LABS	<u>EQ 200MG BASE/5ML</u>	<u>N62304</u>	<u>001</u>	
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PEDIAMYCIN 400

<u>AB</u>	ROSS LABS	<u>EQ 400MG BASE/5ML</u>	<u>N62304</u>	<u>002</u>	
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TABLET; ORAL

E.E.S. 400

<u>AB</u>	+	ABBOTT	<u>EQ 400MG BASE</u>	<u>N61905</u>	<u>002</u>	Aug 12, 1982
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ERYTHROMYCIN ETHYLSUCCINATE

<u>AB</u>	MYLAN	<u>EQ 400MG BASE</u>	<u>N62847</u>	<u>001</u>	Sep 14, 1988
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ERYTHROMYCIN ETHYLSUCCINATE; SULFISOXAZOLE ACETYL

GRANULE; ORAL

ERYTHROMYCIN ETHYLSUCCINATE AND SULFISOXAZOLE ACETYL

<u>AB</u>	BARR	<u>EQ 200MG BASE/5ML;EQ 600MG BASE/5ML</u>	<u>N62759</u>	<u>001</u>	May 20, 1988
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ERYZOLE

<u>AB</u>	ALRA	<u>EQ 200MG BASE/5ML;EQ 600MG BASE/5ML</u>	<u>N62758</u>	<u>001</u>	Jun 15, 1988
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PEDIAZOLE

<u>AB</u>	+	ROSS LABS	<u>EQ 200MG BASE/5ML;EQ 600MG BASE/5ML</u>	<u>N50529</u>	<u>001</u>
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ERYTHROMYCIN LACTOBIONATE

INJECTABLE; INJECTION

ERYTHROCIN

<u>AP</u>	+	HOSPIRA	<u>EQ 500MG BASE/VIAL</u>	<u>N50182</u>	<u>002</u>	
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<u>AP</u>			<u>EQ 500MG BASE/VIAL</u>	<u>N50609</u>	<u>001</u>	Sep 24, 1986
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<u>AP</u>			<u>EQ 500MG BASE/VIAL</u>	<u>N62638</u>	<u>001</u>	Oct 31, 1986
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<u>AP</u>	+		<u>EQ 1GM BASE/VIAL</u>	<u>N50182</u>	<u>003</u>	
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<u>AP</u>			<u>EQ 1GM BASE/VIAL</u>	<u>N50609</u>	<u>002</u>	Sep 24, 1986
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<u>AP</u>			<u>EQ 1GM BASE/VIAL</u>	<u>N62638</u>	<u>002</u>	Oct 31, 1986
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ERYTHROMYCIN LACTOBIONATE

<u>AP</u>	BAXTER HLTHCARE	<u>EQ 500MG BASE/VIAL</u>	<u>N62993</u>	<u>001</u>	May 09, 1989
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<u>AP</u>		<u>EQ 1GM BASE/VIAL</u>	<u>N62993</u>	<u>002</u>	May 09, 1989
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<u>AP</u>	SICOR PHARMS	<u>EQ 500MG BASE/VIAL</u>	<u>N63253</u>	<u>001</u>	Jul 30, 1993
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<u>AP</u>		<u>EQ 1GM BASE/VIAL</u>	<u>N63253</u>	<u>002</u>	Jul 30, 1993
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ERYTHROMYCIN STEARATE

TABLET; ORAL

ERYTHROCIN STEARATE

<u>AB</u>	ABBOTT	<u>EQ 250MG BASE</u>	<u>N60359</u>	<u>001</u>	
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<u>AB</u>	+		<u>EQ 500MG BASE</u>	<u>N60359</u>	<u>003</u>
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ERYTHROMYCIN STEARATE

<u>AB</u>	BARR	<u>EQ 250MG BASE</u>	<u>N61591</u>	<u>001</u>	
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<u>AB</u>	MYLAN	<u>EQ 250MG BASE</u>	<u>N61505</u>	<u>001</u>	
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<u>AB</u>		<u>EQ 500MG BASE</u>	<u>N61505</u>	<u>002</u>	
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27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 147 (of 370)

ESCITALOPRAM OXALATE

SOLUTION; ORAL

LEXAPRO

+ FOREST LABS

EQ 5MG BASE/5ML

N21365 001

Nov 27, 2002

TABLET; ORAL

ESCITALOPRAM OXALATEAB IVAX PHARMSEQ 5MG BASEN76765 001

May 22, 2006

ABEQ 10MG BASEN76765 002

May 22, 2006

ABEQ 20MG BASEN76765 003

May 22, 2006

LEXAPROAB FOREST LABSEQ 5MG BASEN21323 001

Aug 14, 2002

ABEQ 10MG BASEN21323 002

Aug 14, 2002

AB +EQ 20MG BASEN21323 003

Aug 14, 2002

ESMOLOL HYDROCHLORIDE

INJECTABLE; INJECTION

BREVIBLOCAP + BAXTER HLTHCARE CORP10MG/MLN19386 006

Feb 25, 2003

+

20MG/ML

N19386 007

May 28, 2003

+

250MG/ML

N19386 002

Dec 31, 1986

BREVIBLOC DOUBLE STRENGTH IN PLASTIC CONTAINER

+

BAXTER HLTHCARE CORP 2GM/100ML

N19386 005

Jan 27, 2003

BREVIBLOC IN PLASTIC CONTAINER

+

BAXTER HLTHCARE CORP 1GM/100ML

N19386 004

Feb 16, 2001

ESMOLOL HYDROCHLORIDEAP ABRAXIS PHARM10MG/MLN76573 001

May 02, 2005

AP BEDFORD LABS10MG/MLN76323 001

Aug 10, 2004

AP PHARMAFORCE10MG/MLN76474 001

May 02, 2005

ESOMEPRAZOLE MAGNESIUM

CAPSULE, DELAYED REL PELLETS; ORAL

NEXIUM

ASTRAZENECA

EQ 20MG BASE

N21153 001

Feb 20, 2001

+

EQ 40MG BASE

N21153 002

Feb 20, 2001

FOR SUSPENSION, DELAYED RELEASE; ORAL

NEXIUM

ASTRAZENECA

EQ 20MG BASE/PACKET

N21957 001

Oct 20, 2006

+

EQ 40MG BASE/PACKET

N21957 002

Oct 20, 2006

ESOMEPRAZOLE SODIUM

INJECTABLE; INTRAVENOUS

NEXIUM IV

+ ASTRAZENECA

EQ 20MG BASE/VIAL

N21689 001

Mar 31, 2005

+

EQ 40MG BASE/VIAL

N21689 002

Mar 31, 2005

ESTAZOLAM

TABLET; ORAL

ESTAZOLAMAB PAR PHARM1MGN74826 001

Jul 03, 1997

AB2MGN74826 002

Jul 03, 1997

AB TEVA1MGN74921 001

Jul 10, 1997

AB2MGN74921 002

Jul 10, 1997

AB WATSON LABS1MGN74818 001

Aug 19, 1997

AB2MGN74818 002

Aug 19, 1997

PROSOMAB ABBOTT1MGN19080 001

Dec 26, 1990

AB +2MGN19080 002

Dec 26, 1990

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 148 (of 370)

ESTRADIOL

CREAM; VAGINAL					
ESTRACE					
+	WARNER CHILCOTT	0.01%	N86069	001	Jan 31, 1984
FILM, EXTENDED RELEASE; TRANSDERMAL					
ALORA					
BX	WATSON LABS	0.025MG/24HR	N20655	004	Apr 05, 2002
BX		0.05MG/24HR	N20655	001	Dec 20, 1996
BX		0.075MG/24HR	N20655	002	Dec 20, 1996
BX		0.1MG/24HR	N20655	003	Dec 20, 1996
<u>CLIMARA</u>					
<u>AB</u>	BERLEX	<u>0.0375MG/24HR</u>	<u>N20375</u>	<u>005</u>	May 27, 2003
<u>AB</u>		<u>0.06MG/24HR</u>	<u>N20375</u>	<u>006</u>	May 27, 2003
<u>AB2</u>		<u>0.025MG/24HR</u>	<u>N20375</u>	<u>004</u>	Mar 05, 1999
<u>AB2</u>		<u>0.05MG/24HR</u>	<u>N20375</u>	<u>001</u>	Dec 22, 1994
<u>AB2</u>		<u>0.075MG/24HR</u>	<u>N20375</u>	<u>003</u>	Mar 23, 1998
<u>AB2</u> +		<u>0.1MG/24HR</u>	<u>N20375</u>	<u>002</u>	Dec 22, 1994
ESTRADERM					
BX	NOVARTIS	0.05MG/24HR	N19081	002	Sep 10, 1986
BX	+	0.1MG/24HR	N19081	003	Sep 10, 1986
<u>ESTRADIOL</u>					
<u>AB</u>	MYLAN TECHNOLOGIES	<u>0.0375MG/24HR</u>	<u>N75182</u>	<u>004</u>	Jul 20, 2006
<u>AB</u>		<u>0.06MG/24HR</u>	<u>N75182</u>	<u>005</u>	Jul 20, 2006
<u>AB2</u>		<u>0.025MG/24HR</u>	<u>N75182</u>	<u>003</u>	Jan 26, 2005
<u>AB2</u>		<u>0.05MG/24HR</u>	<u>N75233</u>	<u>001</u>	Feb 24, 2000
<u>AB2</u>		<u>0.075MG/24HR</u>	<u>N75182</u>	<u>002</u>	Jan 26, 2005
<u>AB2</u>		<u>0.1MG/24HR</u>	<u>N75182</u>	<u>001</u>	Feb 24, 2000
MENOSTAR					
+	BERLEX	0.014MG/24HR	N21674	001	Jun 08, 2004
<u>VIVELLE</u>					
<u>AB1</u>	NOVARTIS	<u>0.05MG/24HR</u>	<u>N20323</u>	<u>002</u>	Oct 28, 1994
<u>AB1</u>		<u>0.1MG/24HR</u>	<u>N20323</u>	<u>004</u>	Oct 28, 1994
<u>VIVELLE-DOT</u>					
<u>AB1</u>	NOVARTIS	<u>0.05MG/24HR</u>	<u>N20538</u>	<u>006</u>	Jan 08, 1999
<u>AB1</u> +		<u>0.1MG/24HR</u>	<u>N20538</u>	<u>008</u>	Jan 08, 1999
BX		0.025MG/24HR	N20538	009	May 03, 2002
BX		0.0375MG/24HR	N20538	005	Jan 08, 1999
BX		0.075MG/24HR	N20538	007	Jan 08, 1999
GEL, METERED; TRANSDERMAL					
ELESTRIN					
BX	+	BIOSANTE	N21813	001	Dec 15, 2006
ESTROGEL					
BX	+	ASCEND	N21166	002	Feb 09, 2004
INSERT, EXTENDED RELEASE; VAGINAL					
ESTRING					
+	PHARMACIA AND UPJOHN	0.0075MG/24HR	N20472	001	Apr 26, 1996
TABLET; ORAL					
<u>ESTRACE</u>					
<u>AB</u>	BRISTOL MYERS SQUIBB	<u>0.5MG</u>	<u>N81295</u>	<u>001</u>	Jun 30, 1993
<u>AB</u>		<u>1MG</u>	<u>N84499</u>	<u>001</u>	
<u>AB</u> +		<u>2MG</u>	<u>N84500</u>	<u>001</u>	
<u>ESTRADIOL</u>					
<u>AB</u>	APPLIED ANAL	<u>0.5MG</u>	<u>N40138</u>	<u>001</u>	Jan 30, 1998
<u>AB</u>		<u>1MG</u>	<u>N40138</u>	<u>002</u>	Jan 30, 1998
<u>AB</u>		<u>2MG</u>	<u>N40138</u>	<u>003</u>	Jan 30, 1998
<u>AB</u>	BARR	<u>0.5MG</u>	<u>N40197</u>	<u>001</u>	Oct 22, 1997
<u>AB</u>		<u>1MG</u>	<u>N40197</u>	<u>002</u>	Oct 22, 1997
<u>AB</u>		<u>2MG</u>	<u>N40197</u>	<u>003</u>	Oct 22, 1997
<u>AB</u>	MYLAN	<u>0.5MG</u>	<u>N40326</u>	<u>001</u>	Apr 21, 1999
<u>AB</u>		<u>1MG</u>	<u>N40326</u>	<u>002</u>	Apr 21, 1999

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 149 (of 370)

ESTRADIOL

TABLET; ORAL

ESTRADIOL

<u>AB</u>	MYLAN	<u>2MG</u>	<u>N40326</u>	<u>003</u>	Apr 21, 1999
<u>AB</u>	USL PHARMA	<u>0.5MG</u>	<u>N40297</u>	<u>001</u>	Apr 17, 2002
<u>AB</u>		<u>1MG</u>	<u>N40297</u>	<u>002</u>	Apr 17, 2002
<u>AB</u>		<u>2MG</u>	<u>N40297</u>	<u>003</u>	Apr 17, 2002
<u>AB</u>	WATSON LABS	<u>0.5MG</u>	<u>N40114</u>	<u>003</u>	Mar 14, 1996
<u>AB</u>		<u>1MG</u>	<u>N40114</u>	<u>001</u>	Mar 14, 1996
<u>AB</u>		<u>2MG</u>	<u>N40114</u>	<u>002</u>	Mar 14, 1996

GYNODIOL

<u>AB</u>	DURAMED PHARMS BARR	<u>0.5MG</u>	<u>N40212</u>	<u>001</u>	Dec 29, 1997
<u>AB</u>		<u>1MG</u>	<u>N40212</u>	<u>002</u>	Dec 29, 1997
<u>AB</u>		<u>2MG</u>	<u>N40212</u>	<u>004</u>	Dec 29, 1997
		1.5MG	N40212	003	Dec 29, 1997

INNOFEM

<u>AB</u>	NOVO NORDISK INC	<u>0.5MG</u>	<u>N40312</u>	<u>001</u>	Nov 19, 1999
<u>AB</u>		<u>1MG</u>	<u>N40312</u>	<u>002</u>	Nov 19, 1999
<u>AB</u>		<u>2MG</u>	<u>N40312</u>	<u>003</u>	Nov 19, 1999

TABLET; VAGINAL
VAGIFEM

+	NOVO NORDISK INC	25UGM	N20908	001	Mar 26, 1999
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ESTRADIOL ACETATEINSERT, EXTENDED RELEASE; VAGINAL
FEMRING

	GALEN LTD	EQ 0.05MG BASE/24HR	N21367	001	Mar 20, 2003
+		EQ 0.1MG BASE/24HR	N21367	002	Mar 20, 2003

TABLET; ORAL
FEMTRACE

	WARNER CHILCOTT	0.45MG	N21633	001	Aug 20, 2004
		0.9MG	N21633	002	Aug 20, 2004
+		1.8MG	N21633	003	Aug 20, 2004

ESTRADIOL CYPIONATE

INJECTABLE; INJECTION

DEPO-ESTRADIOL

<u>AO</u>	+	PHARMACIA AND UPJOHN	<u>5MG/ML</u>	<u>N85470</u>	<u>003</u>
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ESTRADIOL CYPIONATE

<u>AO</u>		WATSON LABS	<u>5MG/ML</u>	<u>N85620</u>	<u>001</u>
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ESTRADIOL HEMIHYDRATEEMULSION; TOPICAL
ESTRASORB

+	ESPRIT PHARMA	0.25%	N21371	001	Oct 09, 2003
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ESTRADIOL VALERATE

INJECTABLE; INJECTION

DELESTROGEN

<u>AO</u>	+	KING PHARMS	<u>20MG/ML</u>	<u>N09402</u>	<u>004</u>
<u>AO</u>	+		<u>40MG/ML</u>	<u>N09402</u>	<u>003</u>
	+		10MG/ML	N09402	002

ESTRADIOL VALERATE

<u>AO</u>		WATSON LABS	<u>20MG/ML</u>	<u>N83547</u>	<u>001</u>
<u>AO</u>			<u>40MG/ML</u>	<u>N83714</u>	<u>001</u>

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 150 (of 370)

ESTRADIOL; LEVONORGESTRELFILM, EXTENDED RELEASE; TRANSDERMAL
CLIMARA PRO

+ BERLEX 0.045MG/24HR;0.015MG/24HR N21258 001 Nov 21, 2003

ESTRADIOL; NORETHINDRONE ACETATEFILM, EXTENDED RELEASE; TRANSDERMAL
COMBIPATCH

NOVARTIS 0.05MG/24HR;0.14MG/24HR N20870 001 Aug 07, 1998

+ 0.05MG/24HR;0.25MG/24HR N20870 002 Aug 07, 1998

TABLET; ORAL
ACTIVELLA

NOVO NORDISK INC 0.5MG;0.1MG N20907 002 Dec 28, 2006

+ 1MG;0.5MG N20907 001 Nov 18, 1998

ESTRADIOL; NORGESTIMATE

TABLET; ORAL

ESTRADIOL AND NORGESTIMATEAB BARR 1MG, 1MG; N/A, 0.09MG N76812 001 Apr 29, 2005PREFESTAB + DURAMED 1MG, 1MG; N/A, 0.09MG N21040 001 Oct 22, 1999ESTRAMUSTINE PHOSPHATE SODIUMCAPSULE; ORAL
EMCYT

+ PHARMACIA AND UPJOHN EQ 140MG PHOSPHATE N18045 001

ESTROGENS, CONJUGATEDCREAM; TOPICAL, VAGINAL
PREMARIN

+ WYETH PHARMS INC 0.625MG/GM N20216 001

INJECTABLE; INJECTION
PREMARIN

+ WYETH PHARMS INC 25MG/VIAL N10402 001

TABLET; ORAL
PREMARIN

WYETH PHARMS INC 0.3MG N04782 003

0.45MG N04782 006 Jul 16, 2003

+ 0.625MG N04782 004

0.9MG N04782 005 Jan 26, 1984

+ 1.25MG N04782 001

ESTROGENS, CONJUGATED SYNTHETIC ATABLET; ORAL
CENESTIN

DURAMED 0.3MG N20992 001 Jun 21, 2002

0.45MG N20992 005 Feb 05, 2004

0.625MG N20992 002 Mar 24, 1999

0.9MG N20992 003 Mar 24, 1999

+ 1.25MG N20992 004 Mar 13, 2000

ESTROGENS, CONJUGATED SYNTHETIC BTABLET; ORAL
ENJUVIA

DURAMED 0.3MG N21443 001 Dec 20, 2004

0.45MG N21443 002 Dec 20, 2004

0.625MG N21443 003 May 10, 2004

+ 1.25MG N21443 004 May 10, 2004

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 151 (of 370)

ESTROGENS, CONJUGATED; MEDROXYPROGESTERONE ACETATE

TABLET; ORAL-28

PREMPHASE 14/14

+ WYETH PHARMS INC	0.625MG, 0.625MG; N/A, 5MG	N20527	002	Nov 17, 1995
PREMPRO				
+ WYETH PHARMS INC	0.3MG; 1.5MG	N20527	005	Jun 04, 2003
+	0.45MG; 1.5MG	N20527	004	Mar 12, 2003
+	0.625MG; 2.5MG	N20527	001	Nov 17, 1995
+	0.625MG; 5MG	N20527	003	Jan 09, 1998

ESTROGENS, ESTERIFIED

TABLET; ORAL

MENEST

MONARCH PHARMS	0.3MG	N84951	001	
	0.625MG	N84948	001	
	1.25MG	N84950	001	
+	2.5MG	N84949	001	

ESTRONE

INJECTABLE; INJECTION

ESTRONE

+ WATSON LABS	5MG/ML	N85239	001	
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ESTROPIPATE

CREAM; VAGINAL

OGEN

+ PHARMACIA AND UPJOHN	1.5MG/GM	N84710	001	
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TABLET; ORAL

ESTROPIPATE

<u>AB</u>	BARR	<u>0.75MG</u>	<u>N40135</u>	<u>001</u>	Nov 27, 1996
<u>AB</u>		<u>1.5MG</u>	<u>N40135</u>	<u>002</u>	Nov 27, 1996
<u>AB</u>		<u>3MG</u>	<u>N40135</u>	<u>003</u>	Nov 27, 1996
<u>AB</u>	DURAMED PHARMS BARR	<u>0.75MG</u>	<u>N40296</u>	<u>001</u>	Nov 01, 1999
<u>AB</u>		<u>1.5MG</u>	<u>N40296</u>	<u>002</u>	Nov 01, 1999
<u>AB</u>		<u>3MG</u>	<u>N40296</u>	<u>003</u>	Nov 01, 1999
<u>AB</u>	MYLAN	<u>0.75MG</u>	<u>N40359</u>	<u>001</u>	Aug 26, 1999
<u>AB</u>		<u>1.5MG</u>	<u>N40359</u>	<u>002</u>	Aug 26, 1999
<u>AB</u>		<u>3MG</u>	<u>N40359</u>	<u>003</u>	Aug 26, 1999
<u>AB</u>	WATSON LABS	<u>0.75MG</u>	<u>N81213</u>	<u>001</u>	Sep 23, 1993
<u>AB</u>		<u>1.5MG</u>	<u>N81214</u>	<u>001</u>	Sep 23, 1993
<u>AB</u>		<u>3MG</u>	<u>N81215</u>	<u>001</u>	Sep 23, 1993
<u>AB</u>		<u>6MG</u>	<u>N81216</u>	<u>001</u>	Sep 23, 1993
<u>OGEN .625</u>					
<u>AB</u>	PHARMACIA AND UPJOHN	<u>0.75MG</u>	<u>N83220</u>	<u>001</u>	
<u>OGEN 1.25</u>					
<u>AB</u>	PHARMACIA AND UPJOHN	<u>1.5MG</u>	<u>N83220</u>	<u>002</u>	
<u>OGEN 2.5</u>					
<u>AB</u>	+ PHARMACIA AND UPJOHN	<u>3MG</u>	<u>N83220</u>	<u>003</u>	
<u>OGEN 5</u>					
<u>AB</u>	PHARMACIA AND UPJOHN	<u>6MG</u>	<u>N83220</u>	<u>004</u>	
<u>ORTHO-EST</u>					
<u>AB</u>	SUN PHARM INDS (IN)	<u>0.75MG</u>	<u>N89567</u>	<u>001</u>	Feb 27, 1991
<u>AB</u>		<u>1.5MG</u>	<u>N89582</u>	<u>001</u>	Jul 17, 1991

ESZOPICLONE

TABLET; ORAL

LUNESTA

SEPRACOR	1MG	N21476	001	Dec 15, 2004
	2MG	N21476	002	Dec 15, 2004
+	3MG	N21476	003	Dec 15, 2004

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 152 (of 370)

ETHACRYNATE SODIUM

INJECTABLE; INJECTION

EDECIN

+ ATON

EQ 50MG BASE/VIAL

N16093 001

ETHACRYNIC ACID

TABLET; ORAL

EDECIN

+ ATON

25MG

N16092 001

ETHAMBUTOL HYDROCHLORIDE

TABLET; ORAL

ETHAMBUTOL HYDROCHLORIDEAB BARR 400MGN76057 001

Nov 26, 2001

AB WEST WARD 100MGN75095 001

Nov 30, 1999

AB + 400MGN75095 002

Nov 30, 1999

MYAMBUTOLAB STAT TRADE 100MGN16320 001AB 400MGN16320 003ETHANOLAMINE OLEATE

INJECTABLE; INJECTION

ETHAMOLIN

+ QOL MEDCL

50MG/ML

N19357 001

Dec 22, 1988

ETHINYL ESTRADIOL; ETHYNODIOL DIACETATE

TABLET; ORAL-21

DEMULEN 1/35-21AB + GD SEARLE LLC 0.035MG;1MGN18168 001DEMULEN 1/50-21AB + GD SEARLE LLC 0.05MG;1MGN16927 001ZOVIA 1/35E-21AB WATSON LABS 0.035MG;1MGN72720 001

Dec 30, 1991

ZOVIA 1/50E-21AB WATSON LABS 0.05MG;1MGN72722 001

Dec 30, 1991

TABLET; ORAL-28

DEMULEN 1/35-28AB + GD SEARLE LLC 0.035MG;1MGN18160 001DEMULEN 1/50-28AB GD SEARLE LLC 0.05MG;1MGN16936 001KELNORAB BARR 0.035MG;1MGN76785 001

May 23, 2005

ZOVIA 1/35E-28AB WATSON LABS 0.035MG;1MGN72721 001

Dec 30, 1991

ZOVIA 1/50E-28AB WATSON LABS 0.05MG;1MGN72723 001

Dec 30, 1991

ETHINYL ESTRADIOL; ETONOGESTREL

RING; VAGINAL

NUVARING

+ ORGANON USA INC

0.015MG;0.12MG

N21187 001

Oct 03, 2001

ETHINYL ESTRADIOL; LEVONORGESTREL

TABLET; ORAL

QUASENSEAB WATSON LABS 0.03MG;0.15MGN77101 001

Sep 06, 2006

SEASONALEAB + DURAMED 0.03MG;0.15MGN21544 001

Sep 05, 2003

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 153 (of 370)

ETHINYL ESTRADIOL; LEVONORGESTREL

TABLET; ORAL					
SEASONIQUE					
	+ DURAMED RES	0.03MG, 0.01MG; 0.15MG, N/A	N21840	001	May 25, 2006
TABLET; ORAL-28					
<u>ALESSE</u>					
<u>AB1</u>	+ WYETH PHARMS INC	<u>0.02MG; 0.1MG</u>	<u>N20683</u>	<u>002</u>	Mar 27, 1997
<u>AVIANE-28</u>					
<u>AB1</u>	DURAMED PHARMS BARR	<u>0.02MG; 0.1MG</u>	<u>N75796</u>	<u>001</u>	Apr 30, 2001
<u>ENPRESSE-28</u>					
<u>AB</u>	DURAMED PHARMS BARR	<u>0.03MG, 0.04MG, 0.03MG; 0.05MG, 0.075MG, 0.125MG</u>	<u>N75809</u>	<u>002</u>	Jul 16, 2001
<u>LESSINA-28</u>					
<u>AB2</u>	BARR	<u>0.02MG; 0.1MG</u>	<u>N75803</u>	<u>002</u>	Mar 20, 2002
<u>LEVLITE</u>					
<u>AB2</u>	+ BERLEX	<u>0.02MG; 0.1MG</u>	<u>N20860</u>	<u>002</u>	Jul 13, 1998
<u>LEVONORGESTREL AND ETHINYL ESTRADIOL</u>					
<u>AB1</u>	BARR	<u>0.02MG; 0.1MG</u>	<u>N75862</u>	<u>002</u>	Apr 29, 2003
<u>AB1</u>	WATSON LABS	<u>0.02MG; 0.1MG</u>	<u>N76625</u>	<u>001</u>	Nov 18, 2004
<u>AB2</u>		<u>0.02MG; 0.1MG</u>	<u>N77681</u>	<u>001</u>	May 31, 2006
<u>LEVORA 0.15/30-28</u>					
<u>AB</u>	WATSON LABS	<u>0.03MG; 0.15MG</u>	<u>N73594</u>	<u>001</u>	Dec 13, 1993
<u>NORDETTE-28</u>					
<u>AB</u>	+ DURAMED	<u>0.03MG; 0.15MG</u>	<u>N18782</u>	<u>001</u>	Jul 21, 1982
<u>PORTIA-28</u>					
<u>AB</u>	BARR	<u>0.03MG; 0.15MG</u>	<u>N75866</u>	<u>002</u>	May 23, 2002
<u>TRIPHASIL-28</u>					
<u>AB</u>	+ WYETH PHARMS INC	<u>0.03MG, 0.04MG, 0.03MG; 0.05MG, 0.075MG, 0.125MG</u>	<u>N19190</u>	<u>001</u>	Nov 01, 1984
<u>TRIVORA-28</u>					
<u>AB</u>	WATSON LABS	<u>0.03MG, 0.04MG, 0.03MG; 0.05MG, 0.075MG, 0.125MG</u>	<u>N74538</u>	<u>002</u>	Dec 18, 1997

ETHINYL ESTRADIOL; NORELGESTROMIN

FILM, EXTENDED RELEASE; TRANSDERMAL					
ORTHO EVRA					
	+ ORTHO MCNEIL PHARM	0.02MG/24HR; 0.15MG/24HR	N21180	001	Nov 20, 2001

ETHINYL ESTRADIOL; NORETHINDRONE

TABLET; ORAL-21					
BALZIVA-21					
	BARR	0.035MG; 0.4MG	N76198	001	Apr 22, 2004
<u>BREVICON 21-DAY</u>					
<u>AB</u>	+ WATSON LABS	<u>0.035MG; 0.5MG</u>	<u>N17566</u>	<u>001</u>	
<u>GENCEPT 10/11-21</u>					
<u>AB</u>	BARR	<u>0.035MG, 0.035MG; 0.5MG, 1MG</u>	<u>N72694</u>	<u>001</u>	Feb 28, 1992
<u>NORETHIN 1/35E-21</u>					
<u>AB</u>	WATSON LABS	<u>0.035MG; 1MG</u>	<u>N71480</u>	<u>001</u>	Apr 12, 1988
<u>NORETHINDRONE AND ETHINYL ESTRADIOL</u>					
<u>AB</u>	WATSON LABS	<u>0.035MG; 0.5MG</u>	<u>N70684</u>	<u>001</u>	Jan 29, 1987
<u>AB</u>		<u>0.035MG; 1MG</u>	<u>N70685</u>	<u>001</u>	Jan 29, 1987
<u>NORETHINDRONE AND ETHINYL ESTRADIOL (10/11)</u>					
<u>AB</u>	WATSON LABS	<u>0.035MG, 0.035MG; 0.5MG, 1MG</u>	<u>N71043</u>	<u>001</u>	Apr 01, 1988
<u>NORETHINDRONE AND ETHINYL ESTRADIOL (7/14)</u>					
	+ WATSON LABS	0.035MG, 0.035MG; 0.5MG, 1MG	N71041	001	Sep 24, 1991
<u>NORINYL 1+35 21-DAY</u>					
<u>AB</u>	WATSON LABS	<u>0.035MG; 1MG</u>	<u>N17565</u>	<u>001</u>	
<u>NORTREL 0.5/35-21</u>					
<u>AB</u>	BARR	<u>0.035MG; 0.5MG</u>	<u>N72692</u>	<u>001</u>	Feb 28, 1992
<u>NORTREL 1/35-21</u>					
<u>AB</u>	BARR	<u>0.035MG; 1MG</u>	<u>N72693</u>	<u>001</u>	Feb 28, 1992

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 154 (of 370)

ETHINYL ESTRADIOL; NORETHINDRONE

TABLET; ORAL-21

NORTREL 7/7/7

BARR

0.035MG, 0.035MG, 0.035MG; 0.5MG, 0.75MG, 1MG

N75478 001

Aug 30, 2002

TABLET; ORAL-28

ARANELLE

AB

BARR

0.035MG, 0.035MG, 0.035MG; 0.5MG, 1MG, 0.5MGN76783 001

Sep 29, 2004

BALZIVA-28

AB

BARR

0.035MG; 0.4MGN76238 001

Apr 22, 2004

BREVICON 28-DAY

AB

WATSON LABS

0.035MG; 0.5MGN17743 001GENCEPT 10/11-28

AB

BARR

0.035MG, 0.035MG; 0.5MG, 1MGN72697 001

Feb 28, 1992

MODICON 28

AB

ORTHO MCNEIL PHARM

0.035MG; 0.5MGN17735 001NORETHIN 1/35E-28

AB

WATSON LABS

0.035MG; 1MGN71481 001

Apr 12, 1988

NORETHINDRONE AND ETHINYL ESTRADIOL

AB

WATSON LABS

0.035MG; 0.5MGN70686 001

Jan 29, 1987

AB

0.035MG; 1MGN70687 001

Jan 29, 1987

NORETHINDRONE AND ETHINYL ESTRADIOL (10/11)

AB

WATSON LABS

0.035MG, 0.035MG; 0.5MG, 1MGN71044 001

Apr 01, 1988

NORETHINDRONE AND ETHINYL ESTRADIOL (7/14)

WATSON LABS

0.035MG, 0.035MG; 0.5MG, 1MG

N71042 001

Sep 24, 1991

NORINYL 1+35 28-DAY

AB

WATSON LABS

0.035MG; 1MGN17565 002NORTREL 0.5/35-28

AB

BARR

0.035MG; 0.5MGN72695 001

Feb 28, 1992

NORTREL 1/35-28

AB

BARR

0.035MG; 1MGN72696 001

Feb 28, 1992

NORTREL 7/7/7

AB

BARR

0.035MG, 0.035MG, 0.035MG; 0.5MG, 0.75MG, 1MGN75478 002

Aug 30, 2002

ORTHO-NOVUM 1/35-28

AB

+ ORTHO MCNEIL PHARM

0.035MG; 1MGN17919 002ORTHO-NOVUM 10/11-28

AB

+ ORTHO MCNEIL PHARM

0.035MG, 0.035MG; 0.5MG, 1MGN18354 002

Jan 11, 1982

ORTHO-NOVUM 7/7/7-28

AB

+ ORTHO MCNEIL PHARM

0.035MG, 0.035MG, 0.035MG; 0.5MG, 0.75MG, 1MGN18985 002

Apr 04, 1984

OVCON-35

AB

+ WARNER CHILCOTT

0.035MG; 0.4MGN17716 001

OVCON-50

+ WARNER CHILCOTT

0.05MG; 1MG

N17576 001

TRI-NORINYL 28-DAY

AB

+ WATSON LABS

0.035MG, 0.035MG, 0.035MG; 0.5MG, 1MG, 0.5MGN18977 002

Apr 13, 1984

TABLET, CHEWABLE; ORAL-28

OVCON-35

+ WARNER CHILCOTT

0.035MG; 0.4MG

N21490 001

Nov 14, 2003

ETHINYL ESTRADIOL; NORETHINDRONE ACETATE

TABLET; ORAL

FEMHRT

WARNER CHILCOTT

0.0025MG; 0.5MG

N21065 001

Jan 14, 2005

+

0.005MG; 1MG

N21065 002

Oct 15, 1999

LOESTRIN 24 FE

+ WARNER CHILCOTT

0.02MG; 1MG

N21871 001

Feb 17, 2006

TABLET; ORAL-21

JUNEL 1.5/30

AB

BARR

0.03MG; 1.5MGN76381 001

May 30, 2003

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 155 (of 370)

ETHINYL ESTRADIOL; NORETHINDRONE ACETATE

TABLET; ORAL-21					
<u>JUNEL 1/20</u>					
<u>AB</u>	BARR	<u>0.02MG;1MG</u>	<u>N76380</u>	<u>001</u>	May 30, 2003
<u>LOESTRIN 21 1.5/30</u>					
<u>AB</u>	+ WARNER CHILCOTT	<u>0.03MG;1.5MG</u>	<u>N17875</u>	<u>001</u>	
<u>LOESTRIN 21 1/20</u>					
<u>AB</u>	+ WARNER CHILCOTT	<u>0.02MG;1MG</u>	<u>N17876</u>	<u>001</u>	
TABLET; ORAL-28					
ESTROSTEP FE					
+ WARNER CHILCOTT					
0.02MG,0.03MG,0.035MG;1MG,1MG,1MG					
<u>JUNEL FE 1.5/30</u>					
<u>AB</u>	BARR	<u>0.03MG;1.5MG</u>	<u>N76064</u>	<u>001</u>	Sep 18, 2003
<u>JUNEL FE 1/20</u>					
<u>AB</u>	BARR	<u>0.02MG;1MG</u>	<u>N76081</u>	<u>001</u>	Sep 18, 2003
<u>LOESTRIN FE 1.5/30</u>					
<u>AB</u>	+ WARNER CHILCOTT	<u>0.03MG;1.5MG</u>	<u>N17355</u>	<u>001</u>	
<u>LOESTRIN FE 1/20</u>					
<u>AB</u>	+ WARNER CHILCOTT	<u>0.02MG;1MG</u>	<u>N17354</u>	<u>001</u>	
<u>MICROGESTIN FE 1.5/30</u>					
<u>AB</u>	WATSON LABS	<u>0.03MG;1.5MG</u>	<u>N75548</u>	<u>001</u>	Feb 05, 2001
<u>MICROGESTIN FE 1/20</u>					
<u>AB</u>	WATSON LABS	<u>0.02MG;1MG</u>	<u>N75647</u>	<u>001</u>	Feb 05, 2001
<u>NORETHINDRONE ACETATE AND ETHINYL ESTRADIOL AND FERROUS FUMARATE</u>					
<u>AB</u>	ANDRX PHARMS	<u>0.02MG;1MG</u>	<u>N77077</u>	<u>001</u>	May 20, 2005
<u>AB</u>		<u>0.03MG;1.5MG</u>	<u>N77075</u>	<u>001</u>	Apr 28, 2005

ETHINYL ESTRADIOL; NORGESTIMATE

TABLET; ORAL-28					
<u>NORGESTIMATE AND ETHINYL ESTRADIOL</u>					
<u>AB</u>	WATSON LABS	<u>0.035MG,0.035MG,0.035MG;0.18MG,0.215MG,0.25MG</u>	<u>N76626</u>	<u>001</u>	Aug 17, 2006
<u>AB</u>		<u>0.035MG;0.25MG</u>	<u>N76627</u>	<u>001</u>	Aug 17, 2006
<u>ORTHO CYCLEN-28</u>					
<u>AB</u>	+ JOHNSON RW	<u>0.035MG;0.25MG</u>	<u>N19653</u>	<u>002</u>	Dec 29, 1989
<u>ORTHO TRI-CYCLEN</u>					
<u>AB</u>	+ JOHNSON RW	<u>0.035MG,0.035MG,0.035MG;0.18MG,0.215MG,0.25MG</u>	<u>N19697</u>	<u>001</u>	Jul 03, 1992
ORTHO TRI-CYCLEN LO					
+ ORTHO MCNEIL PHARM					
0.025MG,0.025MG,0.25MG;0.18MG,0.215MG,0.25MG					
<u>PREVIFEM</u>					
<u>AB</u>	ANDRX PHARMS	<u>0.035MG;0.25MG</u>	<u>N76334</u>	<u>001</u>	Jan 09, 2004
<u>SPRINTEC</u>					
<u>AB</u>	BARR	<u>0.035MG;0.25MG</u>	<u>N75804</u>	<u>001</u>	Sep 25, 2002
<u>TRI-PREVIFEM</u>					
<u>AB</u>	ANDRX PHARMS	<u>0.035MG,0.035MG,0.035MG;0.18MG,0.215MG,0.25MG</u>	<u>N76335</u>	<u>001</u>	Mar 26, 2004
<u>TRI-SPRINTEC</u>					
<u>AB</u>	BARR	<u>0.035MG,0.035MG,0.035MG;0.18MG,0.215MG,0.25MG</u>	<u>N75808</u>	<u>001</u>	Dec 29, 2003

ETHINYL ESTRADIOL; NORGESTREL

TABLET; ORAL-21					
<u>CRYSELLE</u>					
<u>AB</u>	DURAMED PHARMS BARR	<u>0.03MG;0.3MG</u>	<u>N75840</u>	<u>001</u>	Nov 30, 2001
<u>LOW-OGESTREL-21</u>					
<u>AB</u>	+ WATSON LABS	<u>0.03MG;0.3MG</u>	<u>N75288</u>	<u>001</u>	Jul 28, 1999
OGESTREL 0.5/50-21					
+ WATSON LABS					
0.05MG;0.5MG					
N75406 001 Dec 15, 1999					

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 156 (of 370)

ETHINYL ESTRADIOL; NORGESTREL

TABLET; ORAL-28

CRYSELLE

<u>AB</u>	DURAMED PHARMS BARR	<u>0.03MG;0.3MG</u>	<u>N75840</u>	<u>002</u>	Nov 30, 2001
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LO/OVRAL-28

<u>AB</u>	WYETH PHARMS INC	<u>0.03MG;0.3MG</u>	<u>N17802</u>	<u>001</u>	
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LOW-OGESTREL-28

<u>AB</u>	WATSON LABS	<u>0.03MG;0.3MG</u>	<u>N75288</u>	<u>002</u>	Jul 28, 1999
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OGESTREL 0.5/50-28

+	WATSON LABS	0.05MG;0.5MG	N75406	002	Dec 15, 1999
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ETHIODIZED OIL

OIL; INTRALYMPHATIC

ETHIODOL

SAVAGE LABS	99%	N09190	001	
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ETHIONAMIDE

TABLET; ORAL

TRECATOR-SC

+	WYETH PHARMS INC	250MG	N13026	002	
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ETHOSUXIMIDE

CAPSULE; ORAL

ETHOSUXIMIDE

<u>AB</u>	BANNER PHARMACAPS	<u>250MG</u>	<u>N40430</u>	<u>001</u>	Oct 28, 2002
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ZARONTIN

<u>AB</u>	+	PARKE DAVIS	<u>250MG</u>	<u>N12380</u>	<u>001</u>
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SYRUP; ORAL

ETHOSUXIMIDE

<u>AA</u>	MIKART	<u>250MG/5ML</u>	<u>N40506</u>	<u>001</u>	Dec 22, 2003
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<u>AA</u>	PHARM ASSOC	<u>250MG/5ML</u>	<u>N40253</u>	<u>001</u>	Nov 22, 2000
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<u>AA</u>	TEVA PHARMS	<u>250MG/5ML</u>	<u>N81306</u>	<u>001</u>	Jul 30, 1993
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ZARONTIN

<u>AA</u>	+	PARKE DAVIS	<u>250MG/5ML</u>	<u>N80258</u>	<u>001</u>
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ETHOTOIN

TABLET; ORAL

PEGANONE

+	OVATION PHARMS	250MG	N10841	001	
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ETIDRONATE DISODIUM

TABLET; ORAL

DIDRONEL

<u>AB</u>	PROCTER AND GAMBLE	<u>200MG</u>	<u>N17831</u>	<u>001</u>	
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<u>AB</u>	+		<u>400MG</u>	<u>N17831</u>	<u>002</u>
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ETIDRONATE DISODIUM

<u>AB</u>	GENPHARM	<u>200MG</u>	<u>N75800</u>	<u>001</u>	Jan 24, 2003
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<u>AB</u>		<u>400MG</u>	<u>N75800</u>	<u>002</u>	Jan 24, 2003
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ETODOLAC

CAPSULE; ORAL

ETODOLAC

<u>AB</u>	AAIPHARMA LLC	<u>300MG</u>	<u>N74929</u>	<u>001</u>	Jan 30, 1998
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<u>AB</u>	GENPHARM	<u>200MG</u>	<u>N75071</u>	<u>001</u>	Sep 30, 1998
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<u>AB</u>		<u>300MG</u>	<u>N75071</u>	<u>002</u>	Sep 30, 1998
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<u>AB</u>	MYLAN	<u>200MG</u>	<u>N74932</u>	<u>001</u>	May 16, 1997
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<u>AB</u>		<u>300MG</u>	<u>N74932</u>	<u>002</u>	May 16, 1997
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<u>AB</u>	SANDOZ	<u>200MG</u>	<u>N74840</u>	<u>001</u>	Aug 29, 1997
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<u>AB</u>		<u>200MG</u>	<u>N74942</u>	<u>001</u>	Sep 30, 1997
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PRESCRIPTION DRUG PRODUCT LIST

3 - 157 (of 370)

ETODOLAC

CAPSULE; ORAL

<u>ETODOLAC</u>					
<u>AB</u>	SANDOZ	<u>300MG</u>	<u>N74840</u>	<u>002</u>	Aug 29, 1997
<u>AB</u>		<u>300MG</u>	<u>N74942</u>	<u>002</u>	Sep 30, 1997
<u>AB</u>	TARO	<u>200MG</u>	<u>N75078</u>	<u>001</u>	Apr 30, 1998
<u>AB</u>	+	<u>300MG</u>	<u>N75078</u>	<u>002</u>	Apr 30, 1998
<u>AB</u>	TEVA	<u>200MG</u>	<u>N75126</u>	<u>001</u>	Sep 16, 1999
<u>AB</u>		<u>300MG</u>	<u>N75126</u>	<u>002</u>	Sep 16, 1999
<u>AB</u>	TORPHARM	<u>200MG</u>	<u>N75419</u>	<u>001</u>	Jul 28, 2000
<u>AB</u>		<u>300MG</u>	<u>N75419</u>	<u>002</u>	Jul 28, 2000
<u>AB</u>	WATSON LABS	<u>200MG</u>	<u>N74844</u>	<u>001</u>	Dec 23, 1997
<u>AB</u>		<u>300MG</u>	<u>N74844</u>	<u>002</u>	Dec 23, 1997

TABLET; ORAL

<u>ETODOLAC</u>					
<u>AB</u>	AAIPHARMA LLC	<u>400MG</u>	<u>N74927</u>	<u>001</u>	Oct 30, 1997
<u>AB</u>	ACTAVIS ELIZABETH	<u>400MG</u>	<u>N74819</u>	<u>001</u>	Feb 28, 1997
<u>AB</u>	APOTEX INC	<u>400MG</u>	<u>N76004</u>	<u>001</u>	Dec 03, 2002
<u>AB</u>		<u>500MG</u>	<u>N76004</u>	<u>002</u>	Dec 03, 2002
<u>AB</u>	GENPHARM	<u>400MG</u>	<u>N75012</u>	<u>001</u>	Sep 30, 1998
<u>AB</u>		<u>500MG</u>	<u>N75012</u>	<u>002</u>	Sep 30, 1998
<u>AB</u>	IVAX PHARMS	<u>400MG</u>	<u>N74883</u>	<u>001</u>	Feb 28, 1997
<u>AB</u>		<u>500MG</u>	<u>N74883</u>	<u>002</u>	Nov 20, 1998
<u>AB</u>	MYLAN	<u>400MG</u>	<u>N75104</u>	<u>001</u>	Feb 06, 1998
<u>AB</u>		<u>500MG</u>	<u>N75104</u>	<u>002</u>	Nov 20, 1998
<u>AB</u>	RANBAXY	<u>400MG</u>	<u>N75226</u>	<u>001</u>	Nov 24, 1998
<u>AB</u>		<u>500MG</u>	<u>N75226</u>	<u>002</u>	Nov 24, 1998
<u>AB</u>	SANDOZ	<u>400MG</u>	<u>N74839</u>	<u>001</u>	Jul 11, 1997
<u>AB</u>		<u>400MG</u>	<u>N74846</u>	<u>001</u>	Feb 28, 1997
<u>AB</u>		<u>400MG</u>	<u>N74903</u>	<u>001</u>	Apr 11, 1997
<u>AB</u>		<u>500MG</u>	<u>N74903</u>	<u>002</u>	Apr 19, 1999
<u>AB</u>	TARO PHARM INDS	<u>400MG</u>	<u>N75074</u>	<u>001</u>	Mar 11, 1998
<u>AB</u>	+	<u>500MG</u>	<u>N75074</u>	<u>002</u>	Apr 25, 2000
<u>AB</u>	TEVA	<u>400MG</u>	<u>N74847</u>	<u>001</u>	Apr 23, 1999
<u>AB</u>		<u>400MG</u>	<u>N75009</u>	<u>001</u>	Nov 26, 1997
<u>AB</u>		<u>500MG</u>	<u>N74847</u>	<u>002</u>	Apr 23, 1999
<u>AB</u>		<u>500MG</u>	<u>N75009</u>	<u>002</u>	Dec 28, 1999
<u>AB</u>	WATSON LABS	<u>400MG</u>	<u>N74892</u>	<u>001</u>	Apr 16, 1997
<u>AB</u>		<u>400MG</u>	<u>N75069</u>	<u>001</u>	Apr 16, 1998
<u>AB</u>		<u>500MG</u>	<u>N74892</u>	<u>002</u>	Oct 29, 1998

TABLET, EXTENDED RELEASE; ORAL

<u>ETODOLAC</u>					
<u>AB</u>	ANDRX PHARMS	<u>400MG</u>	<u>N75829</u>	<u>001</u>	Nov 30, 2001
<u>AB</u>		<u>500MG</u>	<u>N75829</u>	<u>002</u>	Nov 30, 2001
<u>AB</u>	SANDOZ	<u>400MG</u>	<u>N75943</u>	<u>001</u>	Jul 26, 2002
<u>AB</u>		<u>500MG</u>	<u>N75943</u>	<u>002</u>	Jul 26, 2002
<u>AB</u>		<u>600MG</u>	<u>N75943</u>	<u>003</u>	Jul 26, 2002
<u>AB</u>	TARO	<u>400MG</u>	<u>N76174</u>	<u>001</u>	Mar 13, 2003
<u>AB</u>		<u>500MG</u>	<u>N76174</u>	<u>002</u>	Mar 13, 2003
<u>AB</u>		<u>600MG</u>	<u>N76174</u>	<u>003</u>	Mar 13, 2003
<u>AB</u>	TEVA	<u>400MG</u>	<u>N75665</u>	<u>003</u>	Feb 05, 2001
<u>AB</u>		<u>500MG</u>	<u>N75665</u>	<u>002</u>	Jul 31, 2000
<u>AB</u>	+	<u>600MG</u>	<u>N75665</u>	<u>001</u>	Jul 31, 2000

ETOMIDATE

INJECTABLE; INJECTION

<u>AMIDATE</u>					
<u>AP</u>	+	<u>2MG/ML</u>	<u>N18227</u>	<u>001</u>	Sep 07, 1982
<u>ETOMIDATE</u>					
<u>AP</u>	BEDFORD	<u>2MG/ML</u>	<u>N74593</u>	<u>001</u>	Nov 04, 1996

PRESCRIPTION DRUG PRODUCT LIST

3 - 158 (of 370)

ETONOGESTREL

IMPLANT; IMPLANTATION

IMPLANON

+ ORGANON USA INC	68MG/IMPLANT	N21529	001	Jul 17, 2006
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ETOPOSIDE

CAPSULE; ORAL

ETOPOSIDE

<u>AB</u> GENPHARM	<u>50MG</u>	<u>N75635</u>	<u>001</u>	Sep 19, 2001
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VEPESID

<u>AB</u> + BRISTOL MYERS SQUIBB	<u>50MG</u>	<u>N19557</u>	<u>001</u>	Dec 30, 1986
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INJECTABLE; INJECTION

ETOPOSIDE

<u>AP</u> ABRAXIS PHARM	<u>20MG/ML</u>	<u>N74983</u>	<u>001</u>	Sep 30, 1998
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<u>AP</u> BEDFORD	<u>20MG/ML</u>	<u>N74290</u>	<u>001</u>	Jul 17, 1995
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<u>AP</u> HOSPIRA	<u>20MG/ML</u>	<u>N74320</u>	<u>001</u>	Aug 30, 1995
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<u>AP</u>	<u>20MG/ML</u>	<u>N74351</u>	<u>001</u>	Aug 30, 1995
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<u>AP</u> MARSAM PHARMS LLC	<u>20MG/ML</u>	<u>N74968</u>	<u>001</u>	Jan 09, 1998
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<u>AP</u> PHARMACHEMIE	<u>20MG/ML</u>	<u>N74227</u>	<u>001</u>	Feb 22, 1996
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<u>AP</u> SICOR PHARMS	<u>20MG/ML</u>	<u>N74284</u>	<u>001</u>	Feb 10, 1994
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<u>AP</u>	<u>20MG/ML</u>	<u>N74529</u>	<u>001</u>	Jul 24, 1996
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<u>AP</u> SUPERGEN	<u>20MG/ML</u>	<u>N74513</u>	<u>001</u>	Mar 14, 1996
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VEPESID

<u>AP</u> + BRISTOL MYERS SQUIBB	<u>20MG/ML</u>	<u>N18768</u>	<u>001</u>	Nov 10, 1983
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ETOPOSIDE PHOSPHATE

INJECTABLE; INJECTION

ETOPHOS PRESERVATIVE FREE

+ BRISTOL MYERS SQUIBB	EQ 100MG BASE/VIAL	N20457	001	May 17, 1996
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EXEMESTANE

TABLET; ORAL

AROMASIN

+ PHARMACIA AND UPJOHN	25MG	N20753	001	Oct 21, 1999
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EXENATIDE SYNTHETIC

INJECTABLE; SUBCUTANEOUS

BYETTA

+ AMYLIN	300UGM/1.2ML (250UGM/ML)	N21773	001	Apr 28, 2005
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+	600UGM/2.4ML (250UGM/ML)	N21773	002	Apr 28, 2005
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EZETIMIBE

TABLET; ORAL

ZETIA

+ MSP SINGAPORE	10MG	N21445	001	Oct 25, 2002
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EZETIMIBE; SIMVASTATIN

TABLET; ORAL

VYTORIN

MSP SINGAPORE	10MG;10MG	N21687	001	Jul 23, 2004
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	10MG;20MG	N21687	002	Jul 23, 2004
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	10MG;40MG	N21687	003	Jul 23, 2004
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+	10MG;80MG	N21687	004	Jul 23, 2004
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FAMCICLOVIR

TABLET; ORAL

FAMVIR

NOVARTIS	125MG	N20363	003	Dec 11, 1995
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	250MG	N20363	001	Apr 26, 1996
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27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 159 (of 370)

FAMCICLOVIR

TABLET; ORAL

FAMVIR

+ NOVARTIS

500MG

N20363 002

Jun 29, 1994

FAMOTIDINE

FOR SUSPENSION; ORAL

PEPCID

+ MERCK

40MG/5ML

N19527 001

Feb 02, 1987

INJECTABLE; INJECTION

FAMOTIDINE

<u>AP</u>	ABRAXIS PHARM	<u>10MG/ML</u>	<u>N75709</u>	<u>001</u>	Apr 16, 2001
<u>AP</u>	APOTEX INC	<u>10MG/ML</u>	<u>N75942</u>	<u>001</u>	Aug 02, 2002
<u>AP</u>	BAXTER HLTHCARE	<u>10MG/ML</u>	<u>N75488</u>	<u>001</u>	Apr 16, 2001
<u>AP</u>		<u>10MG/ML</u>	<u>N75799</u>	<u>001</u>	Apr 30, 2002
<u>AP</u>	BEDFORD	<u>10MG/ML</u>	<u>N75651</u>	<u>001</u>	Apr 16, 2001
<u>AP</u>		<u>10MG/ML</u>	<u>N75684</u>	<u>001</u>	Apr 16, 2001
<u>AP</u>	HOSPIRA	<u>10MG/ML</u>	<u>N75870</u>	<u>001</u>	Nov 23, 2001
<u>AP</u>		<u>10MG/ML</u>	<u>N75905</u>	<u>001</u>	Nov 23, 2001
	<u>FAMOTIDINE PRESERVATIVE FREE</u>				
<u>AP</u>	ABRAXIS PHARM	<u>10MG/ML</u>	<u>N75813</u>	<u>001</u>	Apr 16, 2001
<u>AP</u>	APOTEX INC	<u>10MG/ML</u>	<u>N76324</u>	<u>001</u>	Nov 27, 2002
<u>AP</u>	BAXTER HLTHCARE	<u>10MG/ML</u>	<u>N75486</u>	<u>001</u>	Apr 16, 2001
<u>AP</u>		<u>10MG/ML</u>	<u>N75789</u>	<u>001</u>	Apr 30, 2002
<u>AP</u>	BEDFORD	<u>10MG/ML</u>	<u>N75622</u>	<u>001</u>	Apr 16, 2001
<u>AP</u>	BEN VENUE	<u>10MG/ML</u>	<u>N75825</u>	<u>001</u>	Apr 17, 2001
	<u>FAMOTIDINE PRESERVATIVE FREE (PHARMACY BULK)</u>				
<u>AP</u>	APOTEX INC	<u>10MG/ML</u>	<u>N76322</u>	<u>001</u>	Nov 27, 2002
	<u>FAMOTIDINE PRESERVATIVE FREE IN PLASTIC CONTAINER</u>				
<u>AP</u>	BAXTER HLTHCARE	<u>0.4MG/ML</u>	<u>N75591</u>	<u>001</u>	May 10, 2001
	<u>PEPCID</u>				
<u>AP</u>	+ MERCK	<u>10MG/ML</u>	<u>N19510</u>	<u>001</u>	Nov 04, 1986
	<u>PEPCID PRESERVATIVE FREE</u>				
<u>AP</u>	+ MERCK	<u>10MG/ML</u>	<u>N19510</u>	<u>004</u>	Nov 04, 1986
	<u>PEPCID PRESERVATIVE FREE IN PLASTIC CONTAINER</u>				
<u>AP</u>	+ MERCK	<u>0.4MG/ML</u>	<u>N20249</u>	<u>001</u>	Feb 18, 1994
	TABLET; ORAL				
	<u>FAMOTIDINE</u>				
<u>AB</u>	ACTAVIS ELIZABETH	<u>20MG</u>	<u>N75650</u>	<u>001</u>	Sep 14, 2001
<u>AB</u>		<u>40MG</u>	<u>N75650</u>	<u>002</u>	Sep 14, 2001
<u>AB</u>	CARLSBAD	<u>20MG</u>	<u>N75805</u>	<u>001</u>	Apr 16, 2001
<u>AB</u>		<u>40MG</u>	<u>N75805</u>	<u>002</u>	Apr 16, 2001
<u>AB</u>	DR REDDYS LABS LTD	<u>20MG</u>	<u>N75718</u>	<u>001</u>	Apr 16, 2001
<u>AB</u>		<u>40MG</u>	<u>N75718</u>	<u>002</u>	Apr 16, 2001
<u>AB</u>	GENPHARM	<u>20MG</u>	<u>N75457</u>	<u>001</u>	Apr 18, 2001
<u>AB</u>		<u>40MG</u>	<u>N75457</u>	<u>002</u>	Apr 18, 2001
<u>AB</u>	IVAX PHARMS	<u>20MG</u>	<u>N75511</u>	<u>001</u>	Apr 16, 2001
<u>AB</u>		<u>40MG</u>	<u>N75511</u>	<u>002</u>	Apr 16, 2001
<u>AB</u>	MUTUAL PHARM	<u>20MG</u>	<u>N75639</u>	<u>002</u>	Dec 12, 2001
<u>AB</u>		<u>40MG</u>	<u>N75639</u>	<u>001</u>	Dec 12, 2001
<u>AB</u>	MYLAN	<u>20MG</u>	<u>N75704</u>	<u>001</u>	Apr 16, 2001
<u>AB</u>		<u>40MG</u>	<u>N75704</u>	<u>002</u>	Apr 16, 2001
<u>AB</u>	PERRIGO	<u>20MG</u>	<u>N77352</u>	<u>002</u>	Jul 27, 2005
<u>AB</u>		<u>40MG</u>	<u>N77352</u>	<u>001</u>	Jul 27, 2005
<u>AB</u>	SANDOZ	<u>20MG</u>	<u>N75302</u>	<u>001</u>	Apr 16, 2001
<u>AB</u>		<u>20MG</u>	<u>N75607</u>	<u>001</u>	May 10, 2001
<u>AB</u>		<u>20MG</u>	<u>N75793</u>	<u>001</u>	Apr 16, 2001
<u>AB</u>		<u>40MG</u>	<u>N75302</u>	<u>002</u>	Apr 16, 2001
<u>AB</u>		<u>40MG</u>	<u>N75607</u>	<u>002</u>	May 10, 2001
<u>AB</u>		<u>40MG</u>	<u>N75793</u>	<u>002</u>	Apr 16, 2001

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 160 (of 370)

FAMOTIDINE

TABLET; ORAL

FAMOTIDINE

<u>AB</u>	TEVA	<u>20MG</u>	<u>N75311</u>	<u>001</u>	Apr 16, 2001
<u>AB</u>		<u>40MG</u>	<u>N75311</u>	<u>002</u>	Apr 16, 2001
<u>AB</u>	TORPHARM	<u>20MG</u>	<u>N75611</u>	<u>001</u>	Jul 23, 2001
<u>AB</u>		<u>40MG</u>	<u>N75611</u>	<u>002</u>	Jul 23, 2001
<u>AB</u>	WATSON LABS	<u>20MG</u>	<u>N75062</u>	<u>002</u>	Apr 16, 2001
<u>AB</u>		<u>40MG</u>	<u>N75062</u>	<u>001</u>	Apr 16, 2001
<u>AB</u>	WOCKHARDT	<u>20MG</u>	<u>N75786</u>	<u>001</u>	Apr 16, 2001
<u>AB</u>		<u>40MG</u>	<u>N75786</u>	<u>002</u>	Apr 16, 2001

PEPCID

<u>AB</u>	MERCK	<u>20MG</u>	<u>N19462</u>	<u>001</u>	Oct 15, 1986
<u>AB</u>	+	<u>40MG</u>	<u>N19462</u>	<u>002</u>	Oct 15, 1986

TABLET, ORALLY DISINTEGRATING; ORAL

FLUXID

	SCHWARZ PHARMA	20MG	N21712	001	Sep 24, 2004
+		40MG	N21712	002	Sep 24, 2004

FELBAMATE

SUSPENSION; ORAL

FELBATOL

+	MEDPOINTE	600MG/5ML	N20189	003	Jul 29, 1993
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TABLET; ORAL

FELBATOL

	MEDPOINTE	400MG	N20189	001	Jul 29, 1993
+		600MG	N20189	002	Jul 29, 1993

FELODIPINE

TABLET, EXTENDED RELEASE; ORAL

FELODIPINE

<u>AB</u>	MUTUAL PHARM	<u>2.5MG</u>	<u>N75896</u>	<u>001</u>	Nov 02, 2004
<u>AB</u>		<u>5MG</u>	<u>N75896</u>	<u>002</u>	Nov 02, 2004
<u>AB</u>		<u>10MG</u>	<u>N75896</u>	<u>003</u>	Nov 02, 2004

PLENDIL

<u>AB</u>	ASTRAZENECA	<u>2.5MG</u>	<u>N19834</u>	<u>004</u>	Sep 22, 1994
<u>AB</u>		<u>5MG</u>	<u>N19834</u>	<u>001</u>	Jul 25, 1991
<u>AB</u>	+	<u>10MG</u>	<u>N19834</u>	<u>002</u>	Jul 25, 1991

FENOFIBRATE

CAPSULE; ORAL

ANTARA (MICRONIZED)

	OSCIENT	43MG	N21695	001	Nov 30, 2004
+		130MG	N21695	003	Nov 30, 2004

FENOFIBRATE (MICRONIZED)

<u>AB</u>	IMPAX LABS	<u>67MG</u>	<u>N75868</u>	<u>001</u>	Oct 27, 2003
<u>AB</u>		<u>134MG</u>	<u>N75868</u>	<u>002</u>	Oct 27, 2003
<u>AB</u>		<u>200MG</u>	<u>N75868</u>	<u>003</u>	Oct 27, 2003
<u>AB</u>	TEVA	<u>67MG</u>	<u>N75753</u>	<u>001</u>	Sep 03, 2002
<u>AB</u>		<u>134MG</u>	<u>N75753</u>	<u>002</u>	Apr 09, 2002
<u>AB</u>	+	<u>200MG</u>	<u>N75753</u>	<u>003</u>	Apr 09, 2002
	LIPOFEN				
	CIPHER	50MG	N21612	001	Jan 11, 2006
		100MG	N21612	002	Jan 11, 2006
+		150MG	N21612	003	Jan 11, 2006

TABLET; ORAL

FENOFIBRATE

<u>AB</u>	PAR PHARM	<u>107MG</u>	<u>N76520</u>	<u>002</u>	Dec 29, 2005
<u>AB</u>	RANBAXY	<u>54MG</u>	<u>N76635</u>	<u>001</u>	Oct 31, 2005

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 161 (of 370)

FENOFIBRATE

TABLET; ORAL

FENOFIBRATE

<u>AB</u>	RANBAXY	<u>107MG</u>	<u>N76635</u>	<u>002</u>	Oct 31, 2005
<u>AB</u>		<u>160MG</u>	<u>N76635</u>	<u>003</u>	Oct 31, 2005
<u>AB</u>	TEVA	<u>54MG</u>	<u>N76433</u>	<u>001</u>	May 13, 2005
<u>AB</u>	+	<u>160MG</u>	<u>N76433</u>	<u>002</u>	May 13, 2005
	TRICOR				
	ABBOTT	48MG	N21656	001	Nov 05, 2004
	+	145MG	N21656	002	Nov 05, 2004
	TRIGLIDE				
BX	SKYEPHARMA	160MG	N21350	002	May 07, 2005
		50MG	N21350	001	May 07, 2005

FENOLDOPAM MESYLATE

INJECTABLE; INJECTION

CORLOPAM

<u>AP</u>	+	<u>HOSPIRA</u>	<u>EQ 10MG BASE/ML</u>	<u>N19922</u>	<u>001</u>	Sep 23, 1997
		<u>FENOLDOPAM MESYLATE</u>				
<u>AP</u>		BEDFORD LABS	<u>EQ 10MG BASE/ML</u>	<u>N76582</u>	<u>001</u>	Oct 12, 2004
<u>AP</u>		PHARMAFORCE	<u>EQ 10MG BASE/ML</u>	<u>N76656</u>	<u>001</u>	Dec 01, 2003
<u>AP</u>		SANDOZ	<u>EQ 10MG BASE/ML</u>	<u>N77155</u>	<u>001</u>	Feb 15, 2005

FENOPROFEN CALCIUM

CAPSULE; ORAL

FENOPROFEN CALCIUM

<u>AB</u>		PAR PHARM	<u>EQ 200MG BASE</u>	<u>N72437</u>	<u>001</u>	Aug 22, 1988
<u>AB</u>			<u>EQ 300MG BASE</u>	<u>N72438</u>	<u>001</u>	Aug 22, 1988
<u>AB</u>		SANDOZ	<u>EQ 200MG BASE</u>	<u>N72394</u>	<u>001</u>	Oct 17, 1988
<u>AB</u>			<u>EQ 300MG BASE</u>	<u>N72395</u>	<u>001</u>	Oct 17, 1988
		<u>NALFON</u>				
<u>AB</u>	+	PEDINOL	<u>EQ 300MG BASE</u>	<u>N17604</u>	<u>002</u>	
		<u>NALFON 200</u>				
<u>AB</u>		PEDINOL	<u>EQ 200MG BASE</u>	<u>N17604</u>	<u>003</u>	

TABLET; ORAL

FENOPROFEN CALCIUM

<u>AB</u>		ACTAVIS ELIZABETH	<u>EQ 600MG BASE</u>	<u>N72274</u>	<u>001</u>	May 02, 1988
<u>AB</u>		IVAX PHARMS	<u>EQ 600MG BASE</u>	<u>N72557</u>	<u>001</u>	Aug 29, 1988
<u>AB</u>		MUTUAL PHARM	<u>EQ 600MG BASE</u>	<u>N72902</u>	<u>001</u>	Dec 21, 1990
<u>AB</u>	+	MYLAN	<u>EQ 600MG BASE</u>	<u>N72267</u>	<u>001</u>	Aug 17, 1988
<u>AB</u>		PAR PHARM	<u>EQ 600MG BASE</u>	<u>N72429</u>	<u>001</u>	Aug 17, 1988
<u>AB</u>		SANDOZ	<u>EQ 600MG BASE</u>	<u>N72396</u>	<u>001</u>	Oct 17, 1988
<u>AB</u>		WATSON LABS	<u>EQ 600MG BASE</u>	<u>N72407</u>	<u>001</u>	Aug 17, 1988
<u>AB</u>			<u>EQ 600MG BASE</u>	<u>N72602</u>	<u>001</u>	Oct 11, 1988

FENTANYL

FILM, EXTENDED RELEASE; TRANSDERMAL

DURAGESIC-100

<u>AB</u>		ALZA	<u>100UGM/HR</u>	<u>N19813</u>	<u>001</u>	Aug 07, 1990
		DURAGESIC-12				
		ALZA	12.5UGM/HR	N19813	005	Feb 04, 2005
		<u>DURAGESIC-25</u>				
<u>AB</u>	+	ALZA	<u>25UGM/HR</u>	<u>N19813</u>	<u>004</u>	Aug 07, 1990
		<u>DURAGESIC-50</u>				
<u>AB</u>		ALZA	<u>50UGM/HR</u>	<u>N19813</u>	<u>003</u>	Aug 07, 1990
		<u>DURAGESIC-75</u>				
<u>AB</u>		ALZA	<u>75UGM/HR</u>	<u>N19813</u>	<u>002</u>	Aug 07, 1990
		<u>FENTANYL</u>				
<u>AB</u>		LAVIPHARM LABS	<u>25UGM/HR</u>	<u>N77051</u>	<u>001</u>	Aug 04, 2006
<u>AB</u>			<u>50UGM/HR</u>	<u>N77051</u>	<u>002</u>	Aug 04, 2006

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 162 (of 370)

FENTANYL

FILM, EXTENDED RELEASE; TRANSDERMAL

FENTANYL

<u>AB</u>	LAVIPHARM LABS	<u>75UGM/HR</u>	<u>N77051</u>	<u>003</u>	Aug 04, 2006
<u>AB</u>		<u>100UGM/HR</u>	<u>N77051</u>	<u>004</u>	Aug 04, 2006
<u>AB</u>	MYLAN TECHNOLOGIES	<u>25UGM/HR</u>	<u>N76258</u>	<u>001</u>	Jan 28, 2005
<u>AB</u>		<u>50UGM/HR</u>	<u>N76258</u>	<u>002</u>	Jan 28, 2005
<u>AB</u>		<u>75UGM/HR</u>	<u>N76258</u>	<u>003</u>	Jan 28, 2005
<u>AB</u>		<u>100UGM/HR</u>	<u>N76258</u>	<u>004</u>	Jan 28, 2005

FENTANYL CITRATE

INJECTABLE; INJECTION

FENTANYL CITRATE

<u>AP</u>	HOSPIRA	<u>EQ 0.05MG BASE/ML</u>	<u>N19115</u>	<u>001</u>	Jan 12, 1985
<u>FENTANYL CITRATE PRESERVATIVE FREE</u>					
<u>AP</u>	+ BAXTER HLTHCARE	<u>EQ 0.05MG BASE/ML</u>	<u>N19101</u>	<u>001</u>	Jul 11, 1984
<u>AP</u>	HOSPIRA	<u>EQ 0.05MG BASE/ML</u>	<u>N72786</u>	<u>001</u>	Sep 24, 1991
<u>SUBLIMAZE PRESERVATIVE FREE</u>					
<u>AP</u>	+ AKORN MFG	<u>EQ 0.05MG BASE/ML</u>	<u>N16619</u>	<u>001</u>	

TABLET; BUCCAL

FENTORA

	CEPHALON	EQ 0.1MG BASE	N21947	001	Sep 25, 2006
		EQ 0.2MG BASE	N21947	002	Sep 25, 2006
+		EQ 0.4MG BASE	N21947	003	Sep 25, 2006
		EQ 0.6MG BASE	N21947	004	Sep 25, 2006
		EQ 0.8MG BASE	N21947	005	Sep 25, 2006

TROCHE/LOZENGE; TRANSMUCOSAL

	CEPHALON	EQ 0.2MG BASE	N20747	001	Nov 04, 1998
+		EQ 0.4MG BASE	N20747	002	Nov 04, 1998
		EQ 0.6MG BASE	N20747	003	Nov 04, 1998
		EQ 0.8MG BASE	N20747	004	Nov 04, 1998
		EQ 1.2MG BASE	N20747	005	Nov 04, 1998
		EQ 1.6MG BASE	N20747	006	Nov 04, 1998

FENTANYL HYDROCHLORIDESYSTEM; IONTOPHORESIS, TRANSDERMAL
IONSYS

+	ALZA	10.8UGM	N21338	001	May 22, 2006
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FERRIC HEXACYANOFERRATE(II)

CAPSULE; ORAL

RADIOGARDASE (PRUSSIAN BLUE)

+	HEYL CHEMISCH	500MG	N21626	001	Oct 02, 2003
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FERUMOXIDES

INJECTABLE; INJECTION

FERIDEX I.V.

+	ADV MAGNETICS	EQ 11.2MG IRON/ML	N20416	001	Aug 30, 1996
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FERUMOXSIIL

SUSPENSION; ORAL

GASTROMARK

+	ADV MAGNETICS	EQ 0.175MG IRON/ML	N20410	001	Dec 06, 1996
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FEXOFENADINE HYDROCHLORIDE

CAPSULE; ORAL

ALLEGRA

<u>AB</u>	+ SANOFI AVENTIS US	<u>60MG</u>	<u>N20625</u>	<u>001</u>	Jul 25, 1996
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27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 163 (of 370)

FEXOFENADINE HYDROCHLORIDE

CAPSULE; ORAL

FEXOFENADINE HYDROCHLORIDE

<u>AB</u>	BARR	<u>60MG</u>	<u>N76169</u>	<u>001</u>	Jul 13, 2005
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SUSPENSION; ORAL

ALLEGRA

+	SANOFI AVENTIS US	30MG/5ML	N21963	001	Oct 16, 2006
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TABLET; ORAL

ALLEGRA

<u>AB</u>	SANOFI AVENTIS US	<u>30MG</u>	<u>N20872</u>	<u>001</u>	Feb 25, 2000
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<u>AB</u>		<u>60MG</u>	<u>N20872</u>	<u>002</u>	Feb 25, 2000
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<u>AB</u>	+	<u>180MG</u>	<u>N20872</u>	<u>004</u>	Feb 25, 2000
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FEXOFENADINE HYDROCHLORIDE

<u>AB</u>	BARR	<u>30MG</u>	<u>N76191</u>	<u>001</u>	Aug 31, 2005
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<u>AB</u>		<u>60MG</u>	<u>N76191</u>	<u>002</u>	Aug 31, 2005
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<u>AB</u>		<u>180MG</u>	<u>N76191</u>	<u>003</u>	Aug 31, 2005
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<u>AB</u>	DR REDDYS LABS LTD	<u>30MG</u>	<u>N76502</u>	<u>001</u>	Apr 11, 2006
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<u>AB</u>		<u>60MG</u>	<u>N76502</u>	<u>002</u>	Apr 11, 2006
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<u>AB</u>		<u>180MG</u>	<u>N76502</u>	<u>003</u>	Apr 11, 2006
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<u>AB</u>	TEVA	<u>30MG</u>	<u>N76447</u>	<u>001</u>	Sep 01, 2005
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<u>AB</u>		<u>60MG</u>	<u>N76447</u>	<u>002</u>	Sep 01, 2005
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<u>AB</u>		<u>180MG</u>	<u>N76447</u>	<u>003</u>	Sep 01, 2005
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FEXOFENADINE HYDROCHLORIDE; PSEUDOEPHEDRINE HYDROCHLORIDE

TABLET, EXTENDED RELEASE; ORAL

ALLEGRA D 24 HOUR

+	SANOFI AVENTIS US	180MG;240MG	N21704	001	Oct 19, 2004
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ALLEGRA-D 12 HOUR

<u>AB</u>	+	SANOFI AVENTIS US	<u>60MG;120MG</u>	<u>N20786</u>	<u>001</u>	Dec 24, 1997
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FEXOFENADINE HYDROCHLORIDE AND PSEUDOEPHEDRINE HYDROCHLORIDE

<u>AB</u>	BARR	<u>60MG;120MG</u>	<u>N76236</u>	<u>001</u>	Apr 14, 2005
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FINASTERIDE

TABLET; ORAL

FINASTERIDE

<u>AB</u>	DR REDDYS LABS INC	<u>1MG</u>	<u>N76436</u>	<u>001</u>	Jul 28, 2006
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<u>AB</u>	GEDEON RICHTER USA	<u>5MG</u>	<u>N77251</u>	<u>001</u>	Dec 22, 2006
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<u>AB</u>	IVAX PHARMS	<u>5MG</u>	<u>N76340</u>	<u>001</u>	Jun 19, 2006
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<u>AB</u>	MYLAN	<u>5MG</u>	<u>N77578</u>	<u>001</u>	Dec 18, 2006
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<u>AB</u>	TEVA	<u>5MG</u>	<u>N76511</u>	<u>001</u>	Dec 15, 2006
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PROPECIA

<u>AB</u>	+	MERCK	<u>1MG</u>	<u>N20788</u>	<u>001</u>	Dec 19, 1997
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PROSCAR

<u>AB</u>	+	MERCK	<u>5MG</u>	<u>N20180</u>	<u>001</u>	Jun 19, 1992
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FLAVOXATE HYDROCHLORIDE

TABLET; ORAL

FLAVOXATE HYDROCHLORIDE

<u>AB</u>	HAMPTON LAINE LLC	<u>100MG</u>	<u>N76835</u>	<u>001</u>	Nov 30, 2005
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<u>AB</u>	IMPAX PHARMS	<u>100MG</u>	<u>N76234</u>	<u>001</u>	Aug 28, 2003
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<u>AB</u>	PADDOCK	<u>100MG</u>	<u>N76831</u>	<u>001</u>	Dec 16, 2004
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URISPAS

<u>AB</u>	+	ORTHO MCNEIL PHARM	<u>100MG</u>	<u>N16769</u>	<u>001</u>
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FLECAINIDE ACETATE

TABLET; ORAL

FLECAINIDE ACETATE

<u>AB</u>	ALPHAPHARM	<u>50MG</u>	<u>N75442</u>	<u>001</u>	Jul 31, 2001
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<u>AB</u>		<u>100MG</u>	<u>N75442</u>	<u>002</u>	Jul 31, 2001
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PRESCRIPTION DRUG PRODUCT LIST

3 - 164 (of 370)

FLECAINIDE ACETATE

TABLET; ORAL

FLECAINIDE ACETATE

<u>AB</u>	ALPHAPHARM	<u>150MG</u>	<u>N75442</u>	<u>003</u>	Jul 31, 2001
<u>AB</u>	BARR	<u>50MG</u>	<u>N75882</u>	<u>001</u>	Oct 28, 2002
<u>AB</u>		<u>100MG</u>	<u>N75882</u>	<u>002</u>	Oct 28, 2002
<u>AB</u>		<u>150MG</u>	<u>N75882</u>	<u>003</u>	Oct 28, 2002
<u>AB</u>	RANBAXY	<u>50MG</u>	<u>N76421</u>	<u>001</u>	Mar 28, 2003
<u>AB</u>		<u>100MG</u>	<u>N76421</u>	<u>002</u>	Mar 28, 2003
<u>AB</u>		<u>150MG</u>	<u>N76421</u>	<u>003</u>	Mar 28, 2003
<u>AB</u>	ROXANE	<u>50MG</u>	<u>N76278</u>	<u>001</u>	Jan 14, 2003
<u>AB</u>		<u>100MG</u>	<u>N76278</u>	<u>002</u>	Jan 14, 2003
<u>AB</u>		<u>150MG</u>	<u>N76278</u>	<u>003</u>	Jan 14, 2003
<u>AB</u>	SANDOZ	<u>50MG</u>	<u>N76030</u>	<u>001</u>	Oct 28, 2002
<u>AB</u>		<u>100MG</u>	<u>N76030</u>	<u>002</u>	Oct 28, 2002
<u>AB</u>		<u>150MG</u>	<u>N76030</u>	<u>003</u>	Oct 28, 2002
	<u>TAMBOCOR</u>				
<u>AB</u>	3M	<u>50MG</u>	<u>N18830</u>	<u>004</u>	Aug 23, 1988
<u>AB</u>		<u>100MG</u>	<u>N18830</u>	<u>001</u>	Oct 31, 1985
<u>AB</u>	+	<u>150MG</u>	<u>N18830</u>	<u>003</u>	Jun 03, 1988

FLOXURIDINE

INJECTABLE; INJECTION

FLOXURIDINE

<u>AP</u>	ABRAXIS PHARM	<u>500MG/VIAL</u>	<u>N75837</u>	<u>001</u>	Feb 22, 2001
<u>AP</u>	BEDFORD	<u>500MG/VIAL</u>	<u>N75387</u>	<u>001</u>	Apr 16, 2000
	<u>FUDR</u>				
<u>AP</u>	+	<u>500MG/VIAL</u>	<u>N16929</u>	<u>001</u>	

FLUCONAZOLE

FOR SUSPENSION; ORAL

DIFLUCAN

<u>AB</u>	PFIZER	<u>50MG/5ML</u>	<u>N20090</u>	<u>001</u>	Dec 23, 1993
<u>AB</u>	+	<u>200MG/5ML</u>	<u>N20090</u>	<u>002</u>	Dec 23, 1993

FLUCONAZOLE

<u>AB</u>	RANBAXY	<u>50MG/5ML</u>	<u>N76332</u>	<u>001</u>	Jul 29, 2004
<u>AB</u>		<u>200MG/5ML</u>	<u>N76332</u>	<u>002</u>	Jul 29, 2004
<u>AB</u>	ROXANE	<u>50MG/5ML</u>	<u>N76246</u>	<u>001</u>	Jul 29, 2004
<u>AB</u>		<u>200MG/5ML</u>	<u>N76246</u>	<u>002</u>	Jul 29, 2004
<u>AB</u>	TARO PHARM INDS	<u>50MG/5ML</u>	<u>N76918</u>	<u>001</u>	Dec 18, 2006
<u>AB</u>		<u>200MG/5ML</u>	<u>N76918</u>	<u>002</u>	Dec 18, 2006

INJECTABLE; INJECTION

DIFLUCAN IN DEXTROSE 5% IN PLASTIC CONTAINER

<u>AP</u>	+	<u>PFIZER</u>	<u>200MG/100ML</u>	<u>N19950</u>	<u>003</u>	Sep 29, 1992
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DIFLUCAN IN SODIUM CHLORIDE 0.9%

<u>AP</u>	+	<u>PFIZER</u>	<u>200MG/100ML</u>	<u>N19950</u>	<u>001</u>	Jan 29, 1990
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DIFLUCAN IN SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER

<u>AP</u>	+	<u>PFIZER</u>	<u>200MG/100ML</u>	<u>N19950</u>	<u>002</u>	Jan 29, 1990
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FLUCONAZOLE IN DEXTROSE 5% IN PLASTIC CONTAINER

<u>AP</u>	APOTEX INC	<u>200MG/100ML</u>	<u>N76888</u>	<u>001</u>	Mar 25, 2005
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<u>AP</u>	HOSPIRA	<u>200MG/100ML</u>	<u>N76304</u>	<u>001</u>	Jul 29, 2004
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FLUCONAZOLE IN SODIUM CHLORIDE 0.9%

<u>AP</u>	ABRAXIS PHARM	<u>200MG/100ML</u>	<u>N76145</u>	<u>001</u>	Jul 29, 2004
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<u>AP</u>	BEDFORD	<u>200MG/100ML</u>	<u>N76087</u>	<u>001</u>	Jul 29, 2004
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<u>AP</u>	HIKMA FARMACEUTICA	<u>200MG/100ML</u>	<u>N76736</u>	<u>001</u>	Aug 23, 2005
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<u>AP</u>	SICOR PHARMS	<u>200MG/100ML</u>	<u>N76653</u>	<u>001</u>	Jul 29, 2004
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FLUCONAZOLE IN SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER

<u>AP</u>	APOTEX INC	<u>200MG/100ML</u>	<u>N76889</u>	<u>001</u>	Mar 25, 2005
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<u>AP</u>	BAXTER HLTHCARE	<u>200MG/100ML</u>	<u>N76766</u>	<u>001</u>	Jul 29, 2004
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<u>AP</u>	HOSPIRA	<u>200MG/100ML</u>	<u>N76303</u>	<u>001</u>	Jul 29, 2004
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27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 165 (of 370)

FLUCONAZOLE

INJECTABLE; INJECTION

FLUCONAZOLE IN SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER

<u>AP</u>	MAYNE PHARMA USA	<u>200MG/100ML</u>	<u>N76617</u>	<u>001</u>	Jul 29, 2004
<u>AP</u>	SICOR PHARMS	<u>200MG/100ML</u>	<u>N76837</u>	<u>001</u>	Jan 13, 2005

TABLET; ORAL

DIFLUCAN

<u>AB</u>	PFIZER	<u>50MG</u>	<u>N19949</u>	<u>001</u>	Jan 29, 1990
<u>AB</u>		<u>100MG</u>	<u>N19949</u>	<u>002</u>	Jan 29, 1990
<u>AB</u>		<u>150MG</u>	<u>N19949</u>	<u>004</u>	Jun 30, 1994
<u>AB</u>	+	<u>200MG</u>	<u>N19949</u>	<u>003</u>	Jan 29, 1990

FLUCONAZOLE

<u>AB</u>	DR REDDYS LABS INC	<u>50MG</u>	<u>N76658</u>	<u>001</u>	Jul 29, 2004
<u>AB</u>		<u>100MG</u>	<u>N76658</u>	<u>002</u>	Jul 29, 2004
<u>AB</u>		<u>150MG</u>	<u>N76658</u>	<u>003</u>	Jul 29, 2004
<u>AB</u>		<u>200MG</u>	<u>N76658</u>	<u>004</u>	Jul 29, 2004
<u>AB</u>	GENPHARM	<u>50MG</u>	<u>N76042</u>	<u>001</u>	Jul 29, 2004
<u>AB</u>		<u>100MG</u>	<u>N76042</u>	<u>002</u>	Jul 29, 2004
<u>AB</u>		<u>150MG</u>	<u>N76042</u>	<u>003</u>	Jul 29, 2004
<u>AB</u>		<u>200MG</u>	<u>N76042</u>	<u>004</u>	Jul 29, 2004
<u>AB</u>	GLENMARK PHARMA	<u>50MG</u>	<u>N77253</u>	<u>001</u>	Jan 25, 2006
<u>AB</u>		<u>100MG</u>	<u>N77253</u>	<u>002</u>	Jan 25, 2006
<u>AB</u>		<u>150MG</u>	<u>N77253</u>	<u>003</u>	Jan 25, 2006
<u>AB</u>		<u>200MG</u>	<u>N77253</u>	<u>004</u>	Jan 25, 2006
<u>AB</u>	IVAX PHARMS	<u>50MG</u>	<u>N76077</u>	<u>001</u>	Jul 29, 2004
<u>AB</u>		<u>100MG</u>	<u>N76077</u>	<u>002</u>	Jul 29, 2004
<u>AB</u>		<u>150MG</u>	<u>N76077</u>	<u>003</u>	Jul 29, 2004
<u>AB</u>		<u>200MG</u>	<u>N76077</u>	<u>004</u>	Jul 29, 2004
<u>AB</u>	MYLAN	<u>50MG</u>	<u>N76351</u>	<u>001</u>	Jul 29, 2004
<u>AB</u>		<u>100MG</u>	<u>N76351</u>	<u>002</u>	Jul 29, 2004
<u>AB</u>		<u>150MG</u>	<u>N76351</u>	<u>003</u>	Jul 29, 2004
<u>AB</u>		<u>200MG</u>	<u>N76351</u>	<u>004</u>	Jul 29, 2004
<u>AB</u>	PLIVA	<u>50MG</u>	<u>N76424</u>	<u>001</u>	Jul 29, 2004
<u>AB</u>		<u>100MG</u>	<u>N76424</u>	<u>002</u>	Jul 29, 2004
<u>AB</u>		<u>150MG</u>	<u>N76424</u>	<u>003</u>	Jul 29, 2004
<u>AB</u>		<u>200MG</u>	<u>N76424</u>	<u>004</u>	Jul 29, 2004
<u>AB</u>	RANBAXY	<u>50MG</u>	<u>N76386</u>	<u>001</u>	Jul 29, 2004
<u>AB</u>		<u>100MG</u>	<u>N76386</u>	<u>002</u>	Jul 29, 2004
<u>AB</u>		<u>150MG</u>	<u>N76386</u>	<u>003</u>	Jul 29, 2004
<u>AB</u>		<u>200MG</u>	<u>N76386</u>	<u>004</u>	Jul 29, 2004
<u>AB</u>	ROXANE	<u>50MG</u>	<u>N76213</u>	<u>001</u>	Jul 29, 2004
<u>AB</u>		<u>100MG</u>	<u>N76213</u>	<u>002</u>	Jul 29, 2004
<u>AB</u>		<u>150MG</u>	<u>N76213</u>	<u>003</u>	Jul 29, 2004
<u>AB</u>		<u>200MG</u>	<u>N76213</u>	<u>004</u>	Jul 29, 2004
<u>AB</u>	SANDOZ	<u>50MG</u>	<u>N76086</u>	<u>001</u>	Jul 29, 2004
<u>AB</u>		<u>100MG</u>	<u>N76086</u>	<u>002</u>	Jul 29, 2004
<u>AB</u>		<u>150MG</u>	<u>N76086</u>	<u>003</u>	Jul 29, 2004
<u>AB</u>		<u>200MG</u>	<u>N76086</u>	<u>004</u>	Jul 29, 2004
<u>AB</u>	TARO	<u>50MG</u>	<u>N76507</u>	<u>001</u>	Jul 29, 2004
<u>AB</u>		<u>100MG</u>	<u>N76507</u>	<u>002</u>	Jul 29, 2004
<u>AB</u>		<u>150MG</u>	<u>N76507</u>	<u>003</u>	Jul 29, 2004
<u>AB</u>		<u>200MG</u>	<u>N76507</u>	<u>004</u>	Jul 29, 2004
<u>AB</u>	TEVA	<u>50MG</u>	<u>N74681</u>	<u>001</u>	Jul 29, 2004
<u>AB</u>		<u>100MG</u>	<u>N74681</u>	<u>002</u>	Jul 29, 2004
<u>AB</u>		<u>150MG</u>	<u>N74681</u>	<u>003</u>	Jul 29, 2004
<u>AB</u>		<u>200MG</u>	<u>N74681</u>	<u>004</u>	Jul 29, 2004
<u>AB</u>	TORPHARM	<u>50MG</u>	<u>N76665</u>	<u>001</u>	Jul 29, 2004
<u>AB</u>		<u>100MG</u>	<u>N76665</u>	<u>002</u>	Jul 29, 2004
<u>AB</u>		<u>150MG</u>	<u>N76665</u>	<u>003</u>	Jul 29, 2004
<u>AB</u>		<u>200MG</u>	<u>N76665</u>	<u>004</u>	Jul 29, 2004

PRESCRIPTION DRUG PRODUCT LIST

3 - 166 (of 370)

FLUCONAZOLE

TABLET; ORAL

FLUCONAZOLE

<u>AB</u>	UNIQUE PHARM LABS	<u>50MG</u>	<u>N76957</u>	<u>001</u>	Sep 28, 2005
<u>AB</u>		<u>100MG</u>	<u>N76957</u>	<u>002</u>	Sep 28, 2005
<u>AB</u>		<u>200MG</u>	<u>N76957</u>	<u>003</u>	Sep 28, 2005

FLUCYTOSINE

CAPSULE; ORAL

ANCOBON

	VALEANT	250MG	N17001	001	
+		500MG	N17001	002	

FLUDARABINE PHOSPHATE

INJECTABLE; INJECTION

FLUDARA

<u>AP</u>	+ BERLEX	<u>50MG/VIAL</u>	<u>N20038</u>	<u>001</u>	Apr 18, 1991
	<u>FLUDARABINE PHOSPHATE</u>				
<u>AP</u>	SICOR PHARMS	<u>50MG/VIAL</u>	<u>N76349</u>	<u>001</u>	Aug 28, 2003
+		50MG/2ML (25MG/ML)	N76661	001	Apr 28, 2004

FLUDEOXYGLUCOSE F-18

INJECTABLE; INTRAVENOUS

FLUDEOXYGLUCOSE F 18

+	WEILL MEDCL COLL	10-100mCi/ML	N21768	001	Aug 05, 2004
	FLUDEOXYGLUCOSE F18				
+	NORTH SHORE LIJ	20-200mCi/ML	N21870	001	Aug 19, 2005

FLUDROCORTISONE ACETATE

TABLET; ORAL

FLORINEF

<u>AB</u>	+ KING PHARMS	<u>0.1MG</u>	<u>N10060</u>	<u>001</u>	
	<u>FLUDROCORTISONE ACETATE</u>				
<u>AB</u>	BARR	<u>0.1MG</u>	<u>N40425</u>	<u>001</u>	Jan 21, 2003
<u>AB</u>	IMPAX LABS	<u>0.1MG</u>	<u>N40431</u>	<u>001</u>	Mar 18, 2002

FLUMAZENIL

INJECTABLE; INJECTION

FLUMAZENIL

<u>AP</u>	ABRAXIS PHARM	<u>0.5MG/5ML (0.1MG/ML)</u>	<u>N76955</u>	<u>002</u>	Oct 12, 2004
<u>AP</u>		<u>1MG/10ML (0.1MG/ML)</u>	<u>N76955</u>	<u>001</u>	Oct 12, 2004
<u>AP</u>	APOTEX INC	<u>0.5MG/5ML (0.1MG/ML)</u>	<u>N76755</u>	<u>002</u>	Oct 12, 2004
<u>AP</u>		<u>1MG/10ML (0.1MG/ML)</u>	<u>N76755</u>	<u>001</u>	Oct 12, 2004
<u>AP</u>	BAXTER HLTHCARE	<u>0.5MG/5ML (0.1MG/ML)</u>	<u>N76787</u>	<u>002</u>	Oct 12, 2004
<u>AP</u>		<u>1MG/10ML (0.1MG/ML)</u>	<u>N76787</u>	<u>001</u>	Oct 12, 2004
<u>AP</u>	BEDFORD LABS	<u>0.5MG/5ML (0.1MG/ML)</u>	<u>N76256</u>	<u>002</u>	Oct 12, 2004
<u>AP</u>		<u>1MG/10ML (0.1MG/ML)</u>	<u>N76256</u>	<u>001</u>	Oct 12, 2004
<u>AP</u>	SANDOZ	<u>1MG/10ML (0.1MG/ML)</u>	<u>N77071</u>	<u>002</u>	May 03, 2005
<u>AP</u>		<u>0.5MG/5ML (0.1MG/ML)</u>	<u>N77071</u>	<u>001</u>	May 03, 2005
<u>AP</u>	SICOR PHARMS	<u>0.5MG/5ML (0.1MG/ML)</u>	<u>N76589</u>	<u>002</u>	Oct 12, 2004
<u>AP</u>		<u>1MG/10ML (0.1MG/ML)</u>	<u>N76589</u>	<u>001</u>	Oct 12, 2004
	<u>ROMAZICON</u>				
<u>AP</u>	+ HLR	<u>1MG/10ML (0.1MG/ML)</u>	<u>N20073</u>	<u>001</u>	Dec 20, 1991
<u>AP</u>		<u>0.5MG/5ML (0.1MG/ML)</u>	<u>N20073</u>	<u>002</u>	Dec 20, 1991

FLUNISOLIDE

AEROSOL, METERED; INHALATION

AEROBID

+	ROCHE PALO	0.25MG/INH	N18340	001	Aug 17, 1984
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27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 167 (of 370)

FLUNISOLIDE

AEROSOL, METERED; INHALATION

AEROSPAN HFA

+ FOREST LABS EQ 78UGM BASE/INH N21247 001 Jan 27, 2006

SPRAY, METERED; NASAL

FLUNISOLIDEAB + BAUSCH AND LOMB 0.025MG/SPRAY N74805 001 Feb 20, 2002AB QPHARMA 0.025MG/SPRAY N77704 001 Aug 03, 2006

NASAREL

+ IVAX RES 0.029MG/SPRAY N20409 001 Mar 08, 1995

FLUOCINOLONE ACETONIDE

CREAM; TOPICAL

FLUOCINOLONE ACETONIDEAT FOUGERA 0.01% N88170 001 Dec 16, 1982AT 0.025% N88169 001 Dec 16, 1982AT G AND W LABS 0.01% N89526 001 Jul 26, 1988AT 0.025% N89525 001 Jul 26, 1988AT TARO 0.01% N87102 001 Apr 27, 1982AT 0.025% N87104 001 Apr 27, 1982SYNALARAT + MEDICIS 0.01% N12787 004AT + 0.025% N12787 002AT + 0.025% N12787 005

IMPLANT; INTRAVITREAL

RETISERT

+ BAUSCH AND LOMB 0.59MG N21737 001 Apr 08, 2005

OIL; TOPICAL

DERMA-SMOOTH/FS

+ HILL DERMAC 0.01% N19452 001 Feb 03, 1988

OIL/DROPS; OTIC

+ HILL DERMAC 0.01% N21930 001 Nov 09, 2005

OINTMENT; TOPICAL

FLUOCINOLONE ACETONIDEAT FOUGERA 0.025% N88168 001 Dec 16, 1982AT G AND W LABS 0.025% N89524 001 Jul 26, 1988AT TARO 0.025% N40041 001 Sep 15, 1994SYNALARAT + MEDICIS 0.025% N13960 001

SHAMPOO; TOPICAL

FS SHAMPOO

+ GALDERMA LABS LP 0.01% N20001 001 Aug 27, 1990

SOLUTION; TOPICAL

FLUOCINOLONE ACETONIDEAT FOUGERA 0.01% N88167 001 Dec 16, 1982AT TARO 0.01% N89124 001 Sep 11, 1985SYNALARAT + MEDICIS 0.01% N15296 001FLUOCINOLONE ACETONIDE; HYDROQUINONE; TRETINOIN

CREAM; TOPICAL

TRI-LUMA

+ GALDERMA LABS LP 0.01%;4%;0.05% N21112 001 Jan 18, 2002

PRESCRIPTION DRUG PRODUCT LIST

3 - 168 (of 370)

FLUOCINONIDE

CREAM; TOPICAL

FLUOCINONIDE

<u>AB1</u>	ACTAVIS MID ATLANTIC	<u>0.05%</u>	<u>N73085</u>	<u>001</u>	Feb 14, 1992
<u>AB1</u>	FOUGERA	<u>0.05%</u>	<u>N73030</u>	<u>001</u>	Oct 17, 1994
<u>AB1</u>	TARO	<u>0.05%</u>	<u>N19117</u>	<u>001</u>	Jun 26, 1984
<u>AB1</u>		<u>0.05%</u>	<u>N71500</u>	<u>001</u>	Jun 10, 1987
<u>AB1</u>	TEVA	<u>0.05%</u>	<u>N72488</u>	<u>001</u>	Feb 06, 1989

FLUOCINONIDE EMULSIFIED BASE

<u>AB2</u>	ACTAVIS MID ATLANTIC	<u>0.05%</u>	<u>N74204</u>	<u>001</u>	Jun 13, 1995
<u>AB2</u>	ALTANA	<u>0.05%</u>	<u>N76586</u>	<u>001</u>	Jun 23, 2004
<u>AB2</u>	TARO	<u>0.05%</u>	<u>N72494</u>	<u>001</u>	Jan 19, 1989
<u>AB2</u>	TEVA	<u>0.05%</u>	<u>N72490</u>	<u>001</u>	Feb 07, 1989

LIDEX

<u>AB1</u> +	MEDICIS	<u>0.05%</u>	<u>N16908</u>	<u>002</u>	
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LIDEX-E

<u>AB2</u> +	MEDICIS	<u>0.05%</u>	<u>N16908</u>	<u>003</u>	
	VANOS				
	+ MEDICIS	0.1%	N21758	001	Feb 11, 2005

GEL; TOPICAL

FLUOCINONIDE

<u>AB</u>	FOUGERA	<u>0.05%</u>	<u>N72933</u>	<u>001</u>	Dec 30, 1994
<u>AB</u>	TARO	<u>0.05%</u>	<u>N74935</u>	<u>001</u>	Jul 29, 1997
<u>AB</u>	TEVA	<u>0.05%</u>	<u>N72537</u>	<u>001</u>	Feb 07, 1989

LIDEX

<u>AB</u> +	MEDICIS	<u>0.05%</u>	<u>N17373</u>	<u>001</u>	
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OINTMENT; TOPICAL

FLUOCINONIDE

<u>AB</u>	ALTANA	<u>0.05%</u>	<u>N74905</u>	<u>001</u>	Aug 26, 1997
<u>AB</u>	TARO	<u>0.05%</u>	<u>N75008</u>	<u>001</u>	Jun 30, 1999
<u>AB</u>	TEVA	<u>0.05%</u>	<u>N73481</u>	<u>001</u>	Dec 27, 1991

LIDEX

<u>AB</u> +	MEDICIS	<u>0.05%</u>	<u>N16909</u>	<u>002</u>	
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SOLUTION; TOPICAL

FLUOCINONIDE

<u>AT</u>	ACTAVIS MID ATLANTIC	<u>0.05%</u>	<u>N71535</u>	<u>001</u>	Dec 02, 1988
<u>AT</u>	FOUGERA	<u>0.05%</u>	<u>N72934</u>	<u>001</u>	Feb 27, 1995
<u>AT</u>	TARO	<u>0.05%</u>	<u>N72857</u>	<u>001</u>	Aug 02, 1989
<u>AT</u>		<u>0.05%</u>	<u>N74799</u>	<u>001</u>	Dec 31, 1996
<u>AT</u>	TEVA	<u>0.05%</u>	<u>N72511</u>	<u>001</u>	Feb 07, 1989
<u>AT</u>	TEVA PHARMS	<u>0.05%</u>	<u>N72522</u>	<u>001</u>	Sep 28, 1990

LIDEX

<u>AT</u> +	MEDICIS	<u>0.05%</u>	<u>N18849</u>	<u>001</u>	Apr 06, 1984
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FLUORESCEIN SODIUM

INJECTABLE; INTRAVENOUS

FLUORESCITE

	+ ALCON	EQ 500MG BASE/5ML (EQ 100MG BASE/ML)	N21980	001	Mar 28, 2006
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FLUOROMETHOLONE

OINTMENT; OPHTHALMIC

FML

	+ ALLERGAN	0.1%	N17760	001	Sep 04, 1985
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SUSPENSION/DROPS; OPHTHALMIC

FLUOR-OP

<u>AB</u>	NOVARTIS	<u>0.1%</u>	<u>N70185</u>	<u>001</u>	Feb 27, 1986
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FML

<u>AB</u> +	ALLERGAN	<u>0.1%</u>	<u>N16851</u>	<u>002</u>	Jul 28, 1982
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27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 169 (of 370)

FLUOROMETHOLONE

SUSPENSION/DROPS; OPHTHALMIC

FML FORTE

ALLERGAN 0.25%

N19216 001 Apr 23, 1986

FLUOROMETHOLONE ACETATE

SUSPENSION/DROPS; OPHTHALMIC

FLAREX

+ ALCON 0.1%

N19079 001 Feb 11, 1986

FLUOROMETHOLONE ACETATE; TOBRAMYCIN

SUSPENSION/DROPS; OPHTHALMIC

TOBRASONE

+ ALCON 0.1%;0.3%

N50628 001 Jul 21, 1989

FLUOROMETHOLONE; SULFACETAMIDE SODIUM

SUSPENSION/DROPS; OPHTHALMIC

+ ALLERGAN 0.1%;10%

N19525 001 Sep 29, 1989

FLUOROURACIL

CREAM; TOPICAL

CARAC

+ SANOFI AVENTIS US 0.5%

N20985 001 Oct 27, 2000

EFUDEX

+ VALEANT PHARM INTL 5%

N16831 003

FLUOROPLEX

+ ALLERGAN HERBERT 1%

N16988 001

INJECTABLE; INJECTION

FLUOROURACILAP + ABRAXIS PHARM50MG/MLN40278 001

Sep 30, 1998

AP +50MG/MLN40279 001

Sep 30, 1998

AP +50MG/MLN40291 001

Mar 24, 1999

AP +50MG/MLN40379 001

Nov 15, 2000

AP + SICOR PHARMS50MG/MLN40333 001

Jan 27, 2000

AP +50MG/MLN40334 001

Feb 25, 2000

AP + VALEANT50MG/MLN12209 001

SOLUTION; TOPICAL

EFUDEXAT + VALEANT PHARM INTL2%N16831 001AT +5%N16831 002FLUOROURACILAT TARO2%N76526 001

Nov 05, 2003

AT5%N76526 002

Nov 05, 2003

FLUOXETINE HYDROCHLORIDE

CAPSULE; ORAL

FLUOXETINEAB ALPHAPHARMEQ 10MG BASEN75577 001

Jan 29, 2002

ABEQ 20MG BASEN75577 002

Jan 29, 2002

AB BARREQ 10MG BASEN74803 002

Jan 30, 2002

ABEQ 20MG BASEN74803 001

Aug 02, 2001

ABEQ 40MG BASEN76251 001

May 18, 2005

AB CARLSBADEQ 10MG BASEN76022 001

Jan 30, 2002

ABEQ 20MG BASEN76022 002

Jan 30, 2002

AB DR REDDYS LABS INCEQ 10MG BASEN75465 001

Jan 29, 2002

ABEQ 20MG BASEN75465 002

Jan 29, 2002

ABEQ 40MG BASEN75465 003

Aug 02, 2001

AB IVAX PHARMSEQ 10MG BASEN75245 002

Jan 31, 2002

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 170 (of 370)

FLUOXETINE HYDROCHLORIDE

CAPSULE; ORAL

FLUOXETINE

<u>AB</u>	IVAX PHARMS	<u>EQ 20MG BASE</u>	<u>N75245</u>	<u>001</u>	Jan 31, 2002
<u>AB</u>		<u>EQ 40MG BASE</u>	<u>N75245</u>	<u>003</u>	Sep 28, 2004
<u>AB</u>	MALLINCKRODT	<u>EQ 10MG BASE</u>	<u>N75658</u>	<u>001</u>	Jan 29, 2002
<u>AB</u>		<u>EQ 20MG BASE</u>	<u>N75658</u>	<u>002</u>	Jan 29, 2002
<u>AB</u>	MUTUAL PHARMA	<u>EQ 10MG BASE</u>	<u>N75787</u>	<u>001</u>	Jan 29, 2002
<u>AB</u>		<u>EQ 20MG BASE</u>	<u>N75787</u>	<u>002</u>	Jan 29, 2002
<u>AB</u>	MYLAN	<u>EQ 10MG BASE</u>	<u>N75207</u>	<u>001</u>	Jan 30, 2002
<u>AB</u>		<u>EQ 20MG BASE</u>	<u>N75207</u>	<u>002</u>	Jan 30, 2002
<u>AB</u>	PAR PHARM	<u>EQ 10MG BASE</u>	<u>N76922</u>	<u>001</u>	Dec 16, 2004
<u>AB</u>		<u>EQ 20MG BASE</u>	<u>N76922</u>	<u>002</u>	Dec 16, 2004
<u>AB</u>		<u>EQ 40MG BASE</u>	<u>N76922</u>	<u>003</u>	Dec 16, 2004
<u>AB</u>	PLIVA	<u>EQ 10MG BASE</u>	<u>N76001</u>	<u>001</u>	Jan 29, 2002
<u>AB</u>		<u>EQ 20MG BASE</u>	<u>N76001</u>	<u>002</u>	Jan 29, 2002
<u>AB</u>	RANBAXY	<u>EQ 10MG BASE</u>	<u>N76165</u>	<u>001</u>	Feb 01, 2002
<u>AB</u>		<u>EQ 20MG BASE</u>	<u>N76165</u>	<u>002</u>	Feb 01, 2002
<u>AB</u>		<u>EQ 40MG BASE</u>	<u>N76990</u>	<u>001</u>	Dec 13, 2004
<u>AB</u>	RXELITE	<u>EQ 10MG BASE</u>	<u>N75464</u>	<u>001</u>	Jan 30, 2002
<u>AB</u>		<u>EQ 20MG BASE</u>	<u>N75464</u>	<u>002</u>	Jan 30, 2002
<u>AB</u>	SANDOZ	<u>EQ 10MG BASE</u>	<u>N75049</u>	<u>001</u>	Aug 02, 2001
<u>AB</u>		<u>EQ 10MG BASE</u>	<u>N75807</u>	<u>001</u>	Jan 29, 2002
<u>AB</u>		<u>EQ 20MG BASE</u>	<u>N75807</u>	<u>002</u>	Jan 29, 2002
<u>AB</u>	TEVA	<u>EQ 10MG BASE</u>	<u>N75452</u>	<u>001</u>	Jan 29, 2002
<u>AB</u>		<u>EQ 20MG BASE</u>	<u>N75452</u>	<u>002</u>	Jan 29, 2002
<u>AB</u>		<u>EQ 40MG BASE</u>	<u>N75452</u>	<u>003</u>	Jan 29, 2002
<u>AB</u>	WATSON LABS	<u>EQ 10MG BASE</u>	<u>N75662</u>	<u>001</u>	Jan 29, 2002
<u>AB</u>		<u>EQ 20MG BASE</u>	<u>N75662</u>	<u>002</u>	Jan 29, 2002

FLUOXETINE HYDROCHLORIDE

<u>AB</u>	SANDOZ	<u>EQ 20MG BASE</u>	<u>N75049</u>	<u>002</u>	Jan 29, 2002
<u>AB</u>		<u>EQ 40MG BASE</u>	<u>N75049</u>	<u>003</u>	Jan 29, 2002
	<u>PROZAC</u>				
<u>AB</u>	LILLY	<u>EQ 10MG BASE</u>	<u>N18936</u>	<u>006</u>	Dec 23, 1992
<u>AB</u>		<u>EQ 20MG BASE</u>	<u>N18936</u>	<u>001</u>	Dec 29, 1987
<u>AB</u>	+	<u>EQ 40MG BASE</u>	<u>N18936</u>	<u>003</u>	Jun 15, 1999
	SARAFEM				
	LILLY	EQ 10MG BASE	N18936	007	Jul 06, 2000
	+	EQ 20MG BASE	N18936	008	Jul 06, 2000

CAPSULE, DELAYED REL PELLETS; ORAL

PROZAC WEEKLY

+	LILLY	EQ 90MG BASE	N21235	001	Feb 26, 2001
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SOLUTION; ORAL

FLUOXETINE

<u>AA</u>	ACTAVIS MID ATLANTIC	<u>EQ 20MG BASE/5ML</u>	<u>N75690</u>	<u>001</u>	Jan 31, 2002
<u>AA</u>	MALLINCKRODT	<u>EQ 20MG BASE/5ML</u>	<u>N75920</u>	<u>001</u>	Jan 29, 2002
<u>AA</u>	NOVEX	<u>EQ 20MG BASE/5ML</u>	<u>N75292</u>	<u>001</u>	Feb 07, 2002
<u>AA</u>	PHARM ASSOC	<u>EQ 20MG BASE/5ML</u>	<u>N76015</u>	<u>001</u>	Jan 30, 2002
<u>AA</u>	TEVA	<u>EQ 20MG BASE/5ML</u>	<u>N75506</u>	<u>001</u>	Aug 02, 2001

FLUOXETINE HYDROCHLORIDE

<u>AA</u>	HI TECH PHARMA	<u>EQ 20MG BASE/5ML</u>	<u>N75525</u>	<u>001</u>	Jun 27, 2002
<u>AA</u>	MORTON GROVE	<u>EQ 20MG BASE/5ML</u>	<u>N75514</u>	<u>001</u>	Aug 29, 2002
<u>AA</u>	PAR PHARM	<u>EQ 20MG BASE/5ML</u>	<u>N76458</u>	<u>001</u>	May 14, 2004

PROZAC

<u>AA</u>	+	LILLY	<u>EQ 20MG BASE/5ML</u>	<u>N20101</u>	<u>001</u>	Apr 24, 1991
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TABLET; ORAL

FLUOXETINE HYDROCHLORIDE

<u>AB</u>	ALPHAPHARM	<u>EQ 10MG BASE</u>	<u>N75755</u>	<u>001</u>	Aug 02, 2001
	+	EQ 20MG BASE	N75755	002	Aug 02, 2001
<u>AB</u>	BARR	<u>EQ 10MG BASE</u>	<u>N75810</u>	<u>001</u>	Feb 01, 2002

PRESCRIPTION DRUG PRODUCT LIST

3 - 171 (of 370)

FLUOXETINE HYDROCHLORIDE

TABLET; ORAL

FLUOXETINE HYDROCHLORIDE

AB	DR REDDYS LABS INC	<u>EQ 10MG BASE</u>	<u>N76006</u>	<u>001</u>	Jan 30, 2002
AB	IVAX PHARMS	<u>EQ 10MG BASE</u>	<u>N75865</u>	<u>001</u>	Feb 28, 2002
	+	EQ 40MG BASE	N75865	003	Aug 30, 2004
AB	SANDOZ	<u>EQ 10MG BASE</u>	<u>N76024</u>	<u>001</u>	Jan 29, 2002
AB	TEVA	<u>EQ 10MG BASE</u>	<u>N75872</u>	<u>001</u>	Jan 29, 2002
	SARAFEM				
	WARNER CHILCOTT	EQ 10MG BASE	N21860	001	May 19, 2006
		EQ 15MG BASE	N21860	002	May 19, 2006
	+	EQ 20MG BASE	N21860	003	May 19, 2006

FLUOXETINE HYDROCHLORIDE; OLANZAPINE

CAPSULE; ORAL

SYMBYAX

LILLY

		EQ 25MG BASE;EQ 6MG BASE	N21520	002	Dec 24, 2003
		EQ 25MG BASE;EQ 12MG BASE	N21520	004	Dec 24, 2003
	+	EQ 50MG BASE;EQ 6MG BASE	N21520	003	Dec 24, 2003
		EQ 50MG BASE;EQ 12MG BASE	N21520	005	Dec 24, 2003

FLUOXYMESTERONE

TABLET; ORAL

FLUOXYMESTERONE

BP	USL PHARMA	10MG	N88342	001	Oct 21, 1983
	HALOTESTIN				
BP	+	PHARMACIA AND UPJOHN 10MG	N10611	010	
		2MG	N10611	002	
		5MG	N10611	006	

FLUPHENAZINE DECANOATE

INJECTABLE; IM-SC

FLUPHENAZINE DECANOATE

AO	APOTEX INC	<u>25MG/ML</u>	<u>N75918</u>	<u>001</u>	Aug 17, 2001
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INJECTABLE; INJECTION

FLUPHENAZINE DECANOATE

AO	ABRAXIS PHARM	<u>25MG/ML</u>	<u>N71413</u>	<u>001</u>	Jul 14, 1987
AO	BEDFORD	<u>25MG/ML</u>	<u>N74531</u>	<u>001</u>	Aug 30, 1996
AO	SICOR PHARMS	<u>25MG/ML</u>	<u>N74795</u>	<u>001</u>	Sep 10, 1996

PROLIXIN DECANOATE

AO	+	BRISTOL MYERS SQUIBB	<u>25MG/ML</u>	<u>N16727</u>	<u>001</u>
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FLUPHENAZINE HYDROCHLORIDE

CONCENTRATE; ORAL

FLUPHENAZINE HYDROCHLORIDE

	+	PHARM ASSOC	5MG/ML	N74725	001	Sep 16, 1996
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ELIXIR; ORAL

FLUPHENAZINE HYDROCHLORIDE

	+	PHARM ASSOC	2.5MG/5ML	N40146	001	Aug 21, 1996
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INJECTABLE; INJECTION

FLUPHENAZINE HYDROCHLORIDE

AP	ABRAXIS PHARM	<u>2.5MG/ML</u>	<u>N89556</u>	<u>001</u>	Apr 16, 1987
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PROLIXIN

AP	+	APOTHECON	<u>2.5MG/ML</u>	<u>N11751</u>	<u>005</u>
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TABLET; ORAL

FLUPHENAZINE HYDROCHLORIDE

AB	MYLAN	<u>1MG</u>	<u>N89801</u>	<u>001</u>	Aug 12, 1988
AB		<u>2.5MG</u>	<u>N89802</u>	<u>001</u>	Aug 12, 1988
AB		<u>5MG</u>	<u>N89803</u>	<u>001</u>	Aug 12, 1988

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 172 (of 370)

FLUPHENAZINE HYDROCHLORIDE

TABLET; ORAL

FLUPHENAZINE HYDROCHLORIDE

<u>AB</u>	MYLAN	<u>10MG</u>	<u>N89804</u>	<u>001</u>	Aug 12, 1988
<u>AB</u>	PAR PHARM	<u>1MG</u>	<u>N89740</u>	<u>001</u>	Aug 25, 1988
<u>AB</u>		<u>2.5MG</u>	<u>N89741</u>	<u>001</u>	Aug 25, 1988
<u>AB</u>		<u>5MG</u>	<u>N89742</u>	<u>001</u>	Aug 25, 1988
<u>AB</u>		<u>10MG</u>	<u>N89743</u>	<u>001</u>	Aug 25, 1988
<u>AB</u>	SANDOZ	<u>1MG</u>	<u>N89583</u>	<u>001</u>	Oct 16, 1987
<u>AB</u>		<u>2.5MG</u>	<u>N89584</u>	<u>001</u>	Oct 16, 1987
<u>AB</u>		<u>5MG</u>	<u>N89585</u>	<u>001</u>	Oct 16, 1987
<u>AB</u>		<u>10MG</u>	<u>N89586</u>	<u>001</u>	Oct 16, 1987
	<u>PROLIXIN</u>				
<u>AB</u>	APOTHECON	<u>1MG</u>	<u>N11751</u>	<u>004</u>	
<u>AB</u>		<u>2.5MG</u>	<u>N11751</u>	<u>001</u>	
<u>AB</u>		<u>5MG</u>	<u>N11751</u>	<u>003</u>	
<u>AB</u>	+	<u>10MG</u>	<u>N11751</u>	<u>002</u>	

FLURANDRENOLIDE

CREAM; TOPICAL

CORDRAN SP

+	OCLASSEN	0.025%	N12806	003	
+		0.05%	N12806	002	

LOTION; TOPICAL

CORDRAN

+	WATSON LABS	0.05%	N13790	001	
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OINTMENT; TOPICAL

CORDRAN

+	OCLASSEN	0.025%	N12806	004	
+		0.05%	N12806	001	

TAPE; TOPICAL

CORDRAN

+	OCLASSEN	0.004MG/SQ CM	N16455	001	
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FLURAZEPAM HYDROCHLORIDE

CAPSULE; ORAL

DALMANE

<u>AB</u>	VALEANT PHARM INTL	<u>15MG</u>	<u>N16721</u>	<u>001</u>	
<u>AB</u>	+	<u>30MG</u>	<u>N16721</u>	<u>002</u>	

FLURAZEPAM HYDROCHLORIDE

<u>AB</u>	MYLAN	<u>15MG</u>	<u>N70344</u>	<u>001</u>	Nov 27, 1985
<u>AB</u>		<u>30MG</u>	<u>N70345</u>	<u>001</u>	Nov 27, 1985
<u>AB</u>	PAR PHARM	<u>15MG</u>	<u>N70444</u>	<u>001</u>	Mar 20, 1986
<u>AB</u>		<u>30MG</u>	<u>N70445</u>	<u>001</u>	Mar 20, 1986
<u>AB</u>	SANDOZ	<u>15MG</u>	<u>N71716</u>	<u>001</u>	Jul 31, 1991
<u>AB</u>		<u>30MG</u>	<u>N71717</u>	<u>001</u>	Jul 31, 1991
<u>AB</u>	WATSON LABS	<u>15MG</u>	<u>N71205</u>	<u>001</u>	Nov 25, 1986
<u>AB</u>		<u>30MG</u>	<u>N71068</u>	<u>001</u>	Nov 25, 1986
<u>AB</u>		<u>30MG</u>	<u>N72369</u>	<u>001</u>	Mar 30, 1989
<u>AB</u>	WEST WARD	<u>15MG</u>	<u>N71107</u>	<u>001</u>	Dec 08, 1986
<u>AB</u>		<u>30MG</u>	<u>N71108</u>	<u>001</u>	Dec 08, 1986

FLURBIPROFEN

TABLET; ORAL

ANSAID

<u>AB</u>	PHARMACIA AND UPJOHN	<u>50MG</u>	<u>N18766</u>	<u>002</u>	Oct 31, 1988
<u>AB</u>	+	<u>100MG</u>	<u>N18766</u>	<u>003</u>	Oct 31, 1988
	<u>FLURBIPROFEN</u>				
<u>AB</u>	CARACO	<u>50MG</u>	<u>N75058</u>	<u>001</u>	Apr 27, 2001

PRESCRIPTION DRUG PRODUCT LIST

3 - 173 (of 370)

FLURBIPROFEN

TABLET; ORAL

FLURBIPROFEN

<u>AB</u>	CARACO	<u>100MG</u>	<u>N75058</u>	<u>002</u>	Apr 27, 2001
<u>AB</u>	MYLAN	<u>50MG</u>	<u>N74358</u>	<u>001</u>	Jun 20, 1994
<u>AB</u>		<u>100MG</u>	<u>N74358</u>	<u>002</u>	Jun 20, 1994
<u>AB</u>	PLIVA	<u>50MG</u>	<u>N74647</u>	<u>001</u>	Apr 01, 1997
<u>AB</u>		<u>100MG</u>	<u>N74647</u>	<u>002</u>	Apr 01, 1997
<u>AB</u>	SANDOZ	<u>50MG</u>	<u>N74448</u>	<u>001</u>	Jul 28, 1995
<u>AB</u>		<u>100MG</u>	<u>N74448</u>	<u>002</u>	Jul 28, 1995
<u>AB</u>	TEVA	<u>50MG</u>	<u>N74405</u>	<u>002</u>	May 24, 1995
<u>AB</u>		<u>100MG</u>	<u>N74405</u>	<u>001</u>	May 24, 1995
<u>AB</u>		<u>100MG</u>	<u>N74431</u>	<u>001</u>	May 31, 1995
<u>AB</u>	WARNER CHILCOTT	<u>100MG</u>	<u>N74560</u>	<u>002</u>	May 16, 1997

FLURBIPROFEN SODIUM

SOLUTION/DROPS; OPHTHALMIC

FLURBIPROFEN SODIUM

<u>AT</u>	BAUSCH AND LOMB	<u>0.03%</u>	<u>N74447</u>	<u>001</u>	Jan 04, 1995
	<u>OCUFEN</u>				
<u>AT</u>	+ ALLERGAN	<u>0.03%</u>	<u>N19404</u>	<u>001</u>	Dec 31, 1986

FLUTAMIDE

CAPSULE; ORAL

FLUTAMIDE

<u>AB</u>	BARR	<u>125MG</u>	<u>N75820</u>	<u>001</u>	Sep 18, 2001
<u>AB</u>	GENPHARM	<u>125MG</u>	<u>N76224</u>	<u>001</u>	May 09, 2003
<u>AB</u>	IVAX PHARMS	<u>125MG</u>	<u>N75780</u>	<u>001</u>	Sep 19, 2001
<u>AB</u>	PAR PHARM	<u>125MG</u>	<u>N75298</u>	<u>001</u>	Sep 18, 2001
<u>AB</u>	+ SANDOZ	<u>125MG</u>	<u>N75818</u>	<u>001</u>	Sep 18, 2001

FLUTICASONE PROPIONATE

AEROSOL, METERED; INHALATION

FLOVENT HFA

	+ GLAXO GRP LTD	0.044MG/INH	N21433	003	May 14, 2004
	+	0.11MG/INH	N21433	002	May 14, 2004
	+	0.22MG/INH	N21433	001	May 14, 2004

CREAM; TOPICAL

CUTIVATE

<u>AB</u>	+ ALTANA	<u>0.05%</u>	<u>N19958</u>	<u>001</u>	Dec 18, 1990
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FLUTICASONE PROPIONATE

<u>AB</u>	ALTANA	<u>0.05%</u>	<u>N76451</u>	<u>001</u>	May 14, 2004
<u>AB</u>	G AND W LABS	<u>0.05%</u>	<u>N77055</u>	<u>001</u>	Jun 30, 2006
<u>AB</u>	KV PHARM	<u>0.05%</u>	<u>N76865</u>	<u>001</u>	Sep 10, 2004
<u>AB</u>	PERRIGO NEW YORK	<u>0.05%</u>	<u>N76793</u>	<u>001</u>	May 14, 2004
<u>AB</u>	QLT USA	<u>0.05%</u>	<u>N76633</u>	<u>001</u>	May 14, 2004

LOTION; TOPICAL

CUTIVATE

	+ ALTANA	0.05%	N21152	001	Mar 31, 2005
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OINTMENT; TOPICAL

CUTIVATE

<u>AB</u>	+ ALTANA	<u>0.005%</u>	<u>N19957</u>	<u>001</u>	Dec 14, 1990
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FLUTICASONE PROPIONATE

<u>AB</u>	ALTANA	<u>0.005%</u>	<u>N76300</u>	<u>001</u>	May 14, 2004
<u>AB</u>	G AND W LABS	<u>0.005%</u>	<u>N77168</u>	<u>001</u>	Mar 03, 2006
<u>AB</u>	PERRIGO NEW YORK	<u>0.005%</u>	<u>N76668</u>	<u>001</u>	May 14, 2004
<u>AB</u>	TARO PHARM INDS	<u>0.005%</u>	<u>N77145</u>	<u>001</u>	Jun 14, 2005

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 174 (of 370)

FLUTICASONE PROPIONATE

POWDER; INHALATION

FLOVENT

+	GLAXOSMITHKLINE	0.044MG/INH	N20549	001	Nov 07, 1997
+		0.088MG/INH	N20549	002	Nov 07, 1997
+		0.22MG/INH	N20549	003	Nov 07, 1997

FLOVENT DISKUS 100

+	GLAXOSMITHKLINE	0.1MG/INH	N20833	002	Sep 29, 2000
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FLOVENT DISKUS 250

+	GLAXOSMITHKLINE	0.25MG/INH	N20833	003	Sep 29, 2000
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FLOVENT DISKUS 50

+	GLAXOSMITHKLINE	0.05MG/INH	N20833	001	Sep 29, 2000
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SPRAY, METERED; NASAL

FLONASE

<u>AB</u>	+	GLAXOSMITHKLINE	<u>0.05MG/SPRAY</u>	<u>N20121</u>	<u>001</u>	Oct 19, 1994
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FLUTICASONE PROPIONATE

<u>AB</u>		ROXANE	<u>0.05MG/SPRAY</u>	<u>N76504</u>	<u>001</u>	Feb 22, 2006
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FLUTICASONE PROPIONATE; SALMETEROL XINAFOATE

AEROSOL, METERED; INHALATION

ADVAIR HFA

+	GLAXOSMITHKLINE	0.045MG/INH;EQ 0.021MG BASE/INH	N21254	001	Jun 08, 2006
+		0.115MG/INH;EQ 0.021MG BASE/INH	N21254	002	Jun 08, 2006
+		0.23MG/INH;EQ 0.021MG BASE/INH	N21254	003	Jun 08, 2006

POWDER; INHALATION

ADVAIR DISKUS 100/50

+	GLAXOSMITHKLINE	0.1MG/INH;EQ 0.05MG BASE/INH	N21077	001	Aug 24, 2000
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ADVAIR DISKUS 250/50

+	GLAXOSMITHKLINE	0.25MG/INH;EQ 0.05MG BASE/INH	N21077	002	Aug 24, 2000
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ADVAIR DISKUS 500/50

+	GLAXOSMITHKLINE	0.5MG/INH;EQ 0.05MG BASE/INH	N21077	003	Aug 24, 2000
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FLUVASTATIN SODIUM

CAPSULE; ORAL

LESCOL

	NOVARTIS	EQ 20MG BASE	N20261	001	Dec 31, 1993
+		EQ 40MG BASE	N20261	002	Dec 31, 1993

TABLET, EXTENDED RELEASE; ORAL

LESCOL XL

+	NOVARTIS	80MG	N21192	001	Oct 06, 2000
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FLUVOXAMINE MALEATE

TABLET; ORAL

FLUVOXAMINE MALEATE

<u>AB</u>		<u>ACTAVIS ELIZABETH</u>	<u>25MG</u>	<u>N75901</u>	<u>001</u>	Dec 28, 2000
<u>AB</u>			<u>50MG</u>	<u>N75901</u>	<u>002</u>	Dec 28, 2000
<u>AB</u>			<u>100MG</u>	<u>N75901</u>	<u>003</u>	Dec 28, 2000
<u>AB</u>		<u>BARR</u>	<u>25MG</u>	<u>N75897</u>	<u>001</u>	Jan 25, 2001
<u>AB</u>			<u>50MG</u>	<u>N75897</u>	<u>002</u>	Jan 25, 2001
<u>AB</u>			<u>100MG</u>	<u>N75897</u>	<u>003</u>	Jan 25, 2001
<u>AB</u>		<u>CARACO</u>	<u>25MG</u>	<u>N75900</u>	<u>001</u>	Feb 23, 2006
<u>AB</u>			<u>50MG</u>	<u>N75900</u>	<u>002</u>	Feb 23, 2006
<u>AB</u>			<u>100MG</u>	<u>N75900</u>	<u>003</u>	Feb 23, 2006
<u>AB</u>		<u>GENPHARM</u>	<u>50MG</u>	<u>N75950</u>	<u>001</u>	Oct 15, 2001
<u>AB</u>			<u>100MG</u>	<u>N75950</u>	<u>002</u>	Oct 15, 2001
<u>AB</u>		<u>IVAX PHARMS</u>	<u>25MG</u>	<u>N75898</u>	<u>001</u>	Mar 12, 2001
<u>AB</u>			<u>50MG</u>	<u>N75898</u>	<u>002</u>	Mar 12, 2001
<u>AB</u>			<u>100MG</u>	<u>N75898</u>	<u>003</u>	Mar 12, 2001
<u>AB</u>		<u>MUTUAL PHARM</u>	<u>25MG</u>	<u>N76125</u>	<u>001</u>	Apr 29, 2002

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 175 (of 370)

FLUVOXAMINE MALEATE

TABLET; ORAL

FLUVOXAMINE MALEATE

<u>AB</u>	MUTUAL PHARM	<u>50MG</u>	<u>N76125</u>	<u>002</u>	Apr 29, 2002
<u>AB</u>		<u>100MG</u>	<u>N76125</u>	<u>003</u>	Apr 29, 2002
<u>AB</u>	MYLAN	<u>25MG</u>	<u>N75889</u>	<u>001</u>	Nov 29, 2000
<u>AB</u>		<u>50MG</u>	<u>N75889</u>	<u>002</u>	Nov 29, 2000
<u>AB</u>		<u>100MG</u>	<u>N75889</u>	<u>003</u>	Nov 29, 2000
<u>AB</u>	SANDOZ	<u>25MG</u>	<u>N75887</u>	<u>001</u>	Jan 05, 2001
<u>AB</u>		<u>25MG</u>	<u>N75888</u>	<u>001</u>	Nov 29, 2000
<u>AB</u>		<u>50MG</u>	<u>N75887</u>	<u>002</u>	Jan 05, 2001
<u>AB</u>		<u>50MG</u>	<u>N75888</u>	<u>002</u>	Nov 29, 2000
<u>AB</u>		<u>100MG</u>	<u>N75887</u>	<u>003</u>	Jan 05, 2001
<u>AB</u>	+	<u>100MG</u>	<u>N75888</u>	<u>003</u>	Nov 29, 2000
<u>AB</u>	SYNTHON PHARMS	<u>25MG</u>	<u>N75899</u>	<u>001</u>	Jan 17, 2001
<u>AB</u>		<u>50MG</u>	<u>N75899</u>	<u>002</u>	Jan 17, 2001
<u>AB</u>		<u>100MG</u>	<u>N75899</u>	<u>003</u>	Jan 17, 2001
<u>AB</u>	TEVA	<u>25MG</u>	<u>N75893</u>	<u>001</u>	Sep 10, 2002
<u>AB</u>		<u>50MG</u>	<u>N75893</u>	<u>002</u>	Sep 10, 2002
<u>AB</u>		<u>100MG</u>	<u>N75893</u>	<u>003</u>	Sep 10, 2002
<u>AB</u>	TORPHARM	<u>25MG</u>	<u>N75902</u>	<u>001</u>	May 07, 2001
<u>AB</u>		<u>50MG</u>	<u>N75902</u>	<u>002</u>	May 07, 2001
<u>AB</u>		<u>100MG</u>	<u>N75902</u>	<u>003</u>	May 07, 2001
<u>AB</u>	WATSON LABS	<u>25MG</u>	<u>N75894</u>	<u>001</u>	Apr 18, 2001
<u>AB</u>		<u>50MG</u>	<u>N75894</u>	<u>002</u>	Apr 18, 2001
<u>AB</u>		<u>100MG</u>	<u>N75894</u>	<u>003</u>	Apr 18, 2001

FOLIC ACID

INJECTABLE; INJECTION

FOLIC ACID

<u>AP</u>	+	ABRAXIS PHARM	<u>5MG/ML</u>	<u>N89202</u>	<u>001</u>	Feb 18, 1986
<u>AP</u>		LOCH	<u>5MG/ML</u>	<u>N81066</u>	<u>001</u>	Dec 29, 1993

TABLET; ORAL

FOLIC ACID

<u>AA</u>	JUBILANT PHARMS	<u>1MG</u>	<u>N40514</u>	<u>001</u>	Jun 14, 2005
<u>AA</u>	LEINER	<u>1MG</u>	<u>N85061</u>	<u>001</u>	
<u>AA</u>	MUTUAL PHARM	<u>1MG</u>	<u>N40582</u>	<u>001</u>	Jul 18, 2005
<u>AA</u>	PHARMAX	<u>1MG</u>	<u>N40625</u>	<u>001</u>	Jul 21, 2005
<u>AA</u>	TABLICAPS	<u>1MG</u>	<u>N83133</u>	<u>002</u>	
<u>AA</u>	+	WATSON LABS	<u>N80680</u>	<u>001</u>	
<u>AA</u>	WEST WARD	<u>1MG</u>	<u>N80600</u>	<u>001</u>	
<u>FOLICET</u>					
<u>AA</u>	MISSION PHARMA	<u>1MG</u>	<u>N87438</u>	<u>001</u>	

FOLLITROPIN ALFA/BETA

INJECTABLE; SUBCUTANEOUS

FOLLISTIM AQ

+	ORGANON USA INC	75 IU/0.5ML	N21273	001	Aug 26, 2005
+		150 IU/0.5ML	N21273	002	Aug 26, 2005
+		300 IU/0.36ML	N21211	001	Mar 23, 2004
+		600 IU/0.72ML	N21211	002	Mar 23, 2004
+		900 IU/1.08ML	N21211	004	Feb 11, 2005
GONAL-F					
+	SERONO INC	450 IU/VIAL	N20378	005	Mar 26, 2004
GONAL-F RFF					
+	SERONO INC	75 IU/VIAL	N21765	002	Mar 25, 2004
GONAL-F RFF PEN					
+	SERONO INC	300 IU/0.5ML	N21684	001	May 25, 2004
+		450 IU/0.75ML	N21684	002	May 25, 2004
+		900 IU/1.5ML	N21684	003	May 25, 2004

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 176 (of 370)

FOMEPIZOLEINJECTABLE; INJECTION
ANTIZOL

+ JAZZ 1GM/ML N20696 001 Dec 04, 1997

FONDAPARINUX SODIUMINJECTABLE; SUBCUTANEOUS
ARIXTRA

+	GLAXOSMITHKLINE	2.5MG/0.5ML	N21345	001	Dec 07, 2001
+		5MG/0.4ML	N21345	002	May 28, 2004
+		7.5MG/0.6ML	N21345	003	May 28, 2004
+		10MG/0.8ML	N21345	004	May 28, 2004

FORMOTEROL FUMARATEPOWDER; INHALATION
FORADIL

+ NOVARTIS 0.012MG/INH N20831 001 Feb 16, 2001

FORADIL CERTIHALER
NOVARTIS EQ 0.0085MG BASE/INH N21592 001 Dec 15, 2006FOSAMPRENAVIR CALCIUMTABLET; ORAL
LEXIVA

+ GLAXOSMITHKLINE EQ 700MG BASE N21548 001 Oct 20, 2003

FOSCARNET SODIUMINJECTABLE; INJECTION
FOSCARNET SODIUMAP HOSPIRA 2.4GM/100ML N77174 001 May 31, 2005AP + FOSCAVIR
ASTRAZENECA 2.4GM/100ML N20068 001 Sep 27, 1991FOSFOMYCIN TROMETHAMINEFOR SUSPENSION; ORAL
MONUROL

+ ZAMBON EQ 3GM BASE/PACKET N50717 001 Dec 19, 1996

FOSINOPRIL SODIUMTABLET; ORAL
FOSINOPRIL SODIUM

<u>AB</u>	ANDRX PHARMS	<u>10MG</u>	<u>N76620</u>	<u>001</u>	Oct 15, 2004
<u>AB</u>		<u>20MG</u>	<u>N76620</u>	<u>002</u>	Oct 15, 2004
<u>AB</u>		<u>40MG</u>	<u>N76620</u>	<u>003</u>	Oct 15, 2004
<u>AB</u>	APOTEX INC	<u>10MG</u>	<u>N76906</u>	<u>001</u>	May 17, 2005
<u>AB</u>		<u>20MG</u>	<u>N76906</u>	<u>002</u>	May 17, 2005
<u>AB</u>		<u>40MG</u>	<u>N76906</u>	<u>003</u>	May 17, 2005
<u>AB</u>	COBALT	<u>10MG</u>	<u>N77531</u>	<u>001</u>	Aug 31, 2006
<u>AB</u>		<u>20MG</u>	<u>N77531</u>	<u>002</u>	Aug 31, 2006
<u>AB</u>		<u>40MG</u>	<u>N77531</u>	<u>003</u>	Aug 31, 2006
<u>AB</u>	INVAGEN PHARMS	<u>10MG</u>	<u>N77222</u>	<u>001</u>	Apr 20, 2005
<u>AB</u>		<u>20MG</u>	<u>N77222</u>	<u>002</u>	Apr 20, 2005
<u>AB</u>		<u>40MG</u>	<u>N77222</u>	<u>003</u>	Apr 20, 2005
<u>AB</u>	RANBAXY	<u>10MG</u>	<u>N76580</u>	<u>001</u>	Apr 23, 2004
<u>AB</u>		<u>20MG</u>	<u>N76580</u>	<u>002</u>	Apr 23, 2004
<u>AB</u>		<u>40MG</u>	<u>N76580</u>	<u>003</u>	Apr 23, 2004
<u>AB</u>	SANDOZ	<u>10MG</u>	<u>N76188</u>	<u>001</u>	Oct 08, 2004
<u>AB</u>		<u>10MG</u>	<u>N76483</u>	<u>001</u>	Apr 23, 2004
<u>AB</u>		<u>20MG</u>	<u>N76188</u>	<u>002</u>	Oct 08, 2004
<u>AB</u>		<u>20MG</u>	<u>N76483</u>	<u>002</u>	Apr 23, 2004

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 177 (of 370)

FOSINOPRIL SODIUM

TABLET; ORAL

FOSINOPRIL SODIUM

<u>AB</u>	SANDOZ	<u>40MG</u>	<u>N76188</u>	<u>003</u>	Oct 08, 2004
<u>AB</u>		<u>40MG</u>	<u>N76483</u>	<u>003</u>	Apr 23, 2004
<u>AB</u>	TEVA	<u>10MG</u>	<u>N76139</u>	<u>001</u>	Nov 25, 2003
<u>AB</u>		<u>20MG</u>	<u>N76139</u>	<u>002</u>	Nov 25, 2003
<u>AB</u>		<u>40MG</u>	<u>N76139</u>	<u>003</u>	Nov 25, 2003
<u>AB</u>	WATSON LABS	<u>10MG</u>	<u>N76987</u>	<u>001</u>	Dec 23, 2004
<u>AB</u>		<u>20MG</u>	<u>N76987</u>	<u>002</u>	Dec 23, 2004
<u>AB</u>		<u>40MG</u>	<u>N76987</u>	<u>003</u>	Dec 23, 2004
<u>MONOPRIL</u>					
<u>AB</u>	BRISTOL MYERS SQUIBB	<u>10MG</u>	<u>N19915</u>	<u>002</u>	May 16, 1991
<u>AB</u>		<u>20MG</u>	<u>N19915</u>	<u>003</u>	May 16, 1991
<u>AB</u>	+	<u>40MG</u>	<u>N19915</u>	<u>004</u>	Mar 28, 1995

FOSINOPRIL SODIUM; HYDROCHLOROTHIAZIDE

TABLET; ORAL

FOSINOPRIL SODIUM AND HYDROCHLOROTHIAZIDE

<u>AB</u>	ANDRX PHARMS	<u>10MG;12.5MG</u>	<u>N76608</u>	<u>001</u>	Dec 03, 2004
<u>AB</u>		<u>20MG;12.5MG</u>	<u>N76608</u>	<u>002</u>	Dec 03, 2004
<u>AB</u>	GENPHARM	<u>10MG;12.5MG</u>	<u>N77705</u>	<u>001</u>	Aug 14, 2006
<u>AB</u>		<u>20MG;12.5MG</u>	<u>N77705</u>	<u>002</u>	Aug 14, 2006
<u>AB</u>	RANBAXY	<u>10MG;12.5MG</u>	<u>N76739</u>	<u>001</u>	Dec 17, 2004
<u>AB</u>		<u>20MG;12.5MG</u>	<u>N76739</u>	<u>002</u>	Dec 17, 2004
<u>AB</u>	SANDOZ	<u>10MG;12.5MG</u>	<u>N76961</u>	<u>001</u>	Sep 28, 2005
<u>AB</u>		<u>20MG;12.5MG</u>	<u>N76961</u>	<u>002</u>	Sep 28, 2005
<u>AB</u>	TEVA	<u>10MG;12.5MG</u>	<u>N76945</u>	<u>001</u>	Jul 05, 2006
<u>AB</u>		<u>20MG;12.5MG</u>	<u>N76945</u>	<u>002</u>	Jul 05, 2006
<u>AB</u>	WATSON LABS	<u>10MG;12.5MG</u>	<u>N77144</u>	<u>001</u>	Aug 16, 2005
<u>AB</u>		<u>20MG;12.5MG</u>	<u>N77144</u>	<u>002</u>	Aug 16, 2005
<u>MONOPRIL - HCT</u>					
<u>AB</u>	BRISTOL MYERS SQUIBB	<u>10MG;12.5MG</u>	<u>N20286</u>	<u>002</u>	Nov 30, 1994
<u>AB</u>	+	<u>20MG;12.5MG</u>	<u>N20286</u>	<u>001</u>	Nov 30, 1994

FOSPHENYTOIN SODIUMINJECTABLE; INJECTION
CEREBYX

+ PARKE DAVIS EQ 50MG PHENYTOIN NA/ML N20450 001 Aug 05, 1996

FROVATRIPTAN SUCCINATE

TABLET; ORAL

FROVA

+ ENDO PHARMS EQ 2.5MG BASE N21006 001 Nov 08, 2001

FULVESTRANTINJECTABLE; INTRAMUSCULAR
FASLODEX

+ ASTRAZENECA 50MG/ML N21344 001 Apr 25, 2002

FUROSEMIDE

INJECTABLE; INJECTION

FUROSEMIDE

<u>AP</u>	ABRAXIS PHARM	<u>10MG/ML</u>	<u>N18902</u>	<u>001</u>	May 22, 1984
<u>AP</u>	BAXTER HLTHCARE	<u>10MG/ML</u>	<u>N71439</u>	<u>001</u>	Sep 14, 1990
<u>AP</u>	HOSPIRA	<u>10MG/ML</u>	<u>N18667</u>	<u>001</u>	May 28, 1982
<u>AP</u>		<u>10MG/ML</u>	<u>N70578</u>	<u>001</u>	Jul 08, 1987
<u>AP</u>		<u>10MG/ML</u>	<u>N72080</u>	<u>001</u>	Aug 13, 1991
<u>AP</u>		<u>10MG/ML</u>	<u>N74337</u>	<u>001</u>	Oct 31, 1994

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 178 (of 370)

FUROSEMIDE

INJECTABLE; INJECTION

FUROSEMIDE

<u>AP</u>	HOSPIRA	<u>10MG/ML</u>	<u>N75241</u>	<u>001</u>	May 28, 1999
<u>AP</u>	INTL MEDICATION	<u>10MG/ML</u>	<u>N18025</u>	<u>001</u>	
<u>AP</u>	+ LUITPOLD	<u>10MG/ML</u>	<u>N18579</u>	<u>001</u>	Nov 30, 1983

SOLUTION; ORAL

FUROSEMIDE

<u>AA</u>	MORTON GROVE	<u>10MG/ML</u>	<u>N70655</u>	<u>001</u>	Oct 02, 1987
<u>AA</u>	+ ROXANE	<u>10MG/ML</u>	<u>N70434</u>	<u>001</u>	Apr 22, 1987
		40MG/5ML	N70433	001	Apr 22, 1987

TABLET; ORAL

FUROSEMIDE

<u>AB</u>	EXCELLIUM	<u>20MG</u>	<u>N77293</u>	<u>001</u>	Nov 09, 2005
<u>AB</u>		<u>40MG</u>	<u>N77293</u>	<u>002</u>	Nov 09, 2005
<u>AB</u>		<u>80MG</u>	<u>N77293</u>	<u>003</u>	Nov 09, 2005
<u>AB</u>	INTL MEDICATION	<u>20MG</u>	<u>N18753</u>	<u>001</u>	Feb 28, 1984
<u>AB</u>		<u>40MG</u>	<u>N18753</u>	<u>002</u>	Feb 28, 1984
<u>AB</u>	IVAX PHARMS	<u>20MG</u>	<u>N18413</u>	<u>001</u>	Nov 30, 1983
<u>AB</u>		<u>40MG</u>	<u>N18413</u>	<u>002</u>	Nov 30, 1983
<u>AB</u>	MYLAN	<u>20MG</u>	<u>N18487</u>	<u>001</u>	
<u>AB</u>		<u>40MG</u>	<u>N18487</u>	<u>002</u>	
<u>AB</u>		<u>80MG</u>	<u>N70082</u>	<u>001</u>	Oct 29, 1986
<u>AB</u>	OHM LABS	<u>20MG</u>	<u>N78010</u>	<u>001</u>	Sep 18, 2006
<u>AB</u>		<u>40MG</u>	<u>N78010</u>	<u>002</u>	Sep 18, 2006
<u>AB</u>		<u>80MG</u>	<u>N78010</u>	<u>003</u>	Sep 18, 2006
<u>AB</u>	ROXANE	<u>20MG</u>	<u>N18823</u>	<u>001</u>	Nov 10, 1983
<u>AB</u>		<u>40MG</u>	<u>N18823</u>	<u>002</u>	Nov 10, 1983
<u>AB</u>		<u>80MG</u>	<u>N70086</u>	<u>001</u>	Jan 24, 1986
<u>AB</u>	SANDOZ	<u>20MG</u>	<u>N18569</u>	<u>002</u>	
<u>AB</u>		<u>40MG</u>	<u>N18569</u>	<u>001</u>	
<u>AB</u>		<u>80MG</u>	<u>N18569</u>	<u>005</u>	Aug 14, 1984
<u>AB</u>	VINTAGE PHARMS	<u>20MG</u>	<u>N76796</u>	<u>001</u>	Mar 26, 2004
<u>AB</u>		<u>40MG</u>	<u>N76796</u>	<u>002</u>	Mar 26, 2004
<u>AB</u>		<u>80MG</u>	<u>N76796</u>	<u>003</u>	Mar 26, 2004
<u>AB</u>	WATSON LABS	<u>20MG</u>	<u>N70412</u>	<u>001</u>	Feb 26, 1986
<u>AB</u>		<u>20MG</u>	<u>N70449</u>	<u>001</u>	Nov 22, 1985
<u>AB</u>		<u>20MG</u>	<u>N71379</u>	<u>001</u>	Jan 02, 1987
<u>AB</u>		<u>40MG</u>	<u>N70450</u>	<u>001</u>	Nov 22, 1985
<u>AB</u>		<u>80MG</u>	<u>N70528</u>	<u>001</u>	Jan 07, 1986
<u>AB</u>		<u>80MG</u>	<u>N71594</u>	<u>001</u>	Feb 09, 1988
	<u>LASIX</u>				
<u>AB</u>	SANOFI AVENTIS US	<u>20MG</u>	<u>N16273</u>	<u>002</u>	
<u>AB</u>		<u>40MG</u>	<u>N16273</u>	<u>001</u>	
<u>AB</u>	+	<u>80MG</u>	<u>N16273</u>	<u>003</u>	

GABAPENTIN

CAPSULE; ORAL

GABAPENTIN

<u>AB</u>	ACTAVIS ELIZABETH	<u>100MG</u>	<u>N75350</u>	<u>001</u>	Sep 12, 2003
<u>AB</u>		<u>300MG</u>	<u>N75350</u>	<u>002</u>	Sep 12, 2003
<u>AB</u>		<u>400MG</u>	<u>N75350</u>	<u>003</u>	Sep 12, 2003
<u>AB</u>	APOTEX INC	<u>100MG</u>	<u>N75360</u>	<u>001</u>	Apr 06, 2005
<u>AB</u>		<u>300MG</u>	<u>N75360</u>	<u>002</u>	Apr 06, 2005
<u>AB</u>		<u>400MG</u>	<u>N75360</u>	<u>003</u>	Apr 06, 2005
<u>AB</u>	IVAX PHARMS	<u>100MG</u>	<u>N75477</u>	<u>001</u>	Mar 23, 2005
<u>AB</u>		<u>300MG</u>	<u>N75477</u>	<u>002</u>	Mar 23, 2005
<u>AB</u>		<u>400MG</u>	<u>N75477</u>	<u>003</u>	Mar 23, 2005
<u>AB</u>	MUTUAL PHARM	<u>100MG</u>	<u>N76537</u>	<u>001</u>	Jun 30, 2005
<u>AB</u>		<u>300MG</u>	<u>N76537</u>	<u>002</u>	Jun 30, 2005

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 179 (of 370)

GABAPENTIN

CAPSULE; ORAL

GABAPENTIN

<u>AB</u>	MUTUAL PHARM	<u>400MG</u>	<u>N76537</u>	<u>003</u>	Jun 30, 2005
<u>AB</u>	RANBAXY	<u>100MG</u>	<u>N76606</u>	<u>001</u>	Oct 07, 2005
<u>AB</u>		<u>300MG</u>	<u>N76606</u>	<u>002</u>	Oct 07, 2005
<u>AB</u>		<u>400MG</u>	<u>N76606</u>	<u>003</u>	Oct 07, 2005
<u>AB</u>	SANDOZ	<u>100MG</u>	<u>N75428</u>	<u>001</u>	Jan 24, 2006
<u>AB</u>		<u>100MG</u>	<u>N75539</u>	<u>001</u>	Apr 06, 2005
<u>AB</u>		<u>300MG</u>	<u>N75428</u>	<u>002</u>	Jan 24, 2006
<u>AB</u>		<u>300MG</u>	<u>N75539</u>	<u>002</u>	Apr 06, 2005
<u>AB</u>		<u>400MG</u>	<u>N75428</u>	<u>003</u>	Jan 24, 2006
<u>AB</u>		<u>400MG</u>	<u>N75539</u>	<u>003</u>	Apr 06, 2005
<u>AB</u>	SUN PHARM INDS LTD	<u>100MG</u>	<u>N77242</u>	<u>001</u>	Aug 24, 2006
<u>AB</u>		<u>300MG</u>	<u>N77242</u>	<u>002</u>	Aug 24, 2006
<u>AB</u>		<u>400MG</u>	<u>N77242</u>	<u>003</u>	Aug 24, 2006
<u>AB</u>	TEVA PHARMS	<u>100MG</u>	<u>N75435</u>	<u>001</u>	Oct 08, 2004
<u>AB</u>		<u>300MG</u>	<u>N75435</u>	<u>002</u>	Oct 08, 2004
<u>AB</u>		<u>400MG</u>	<u>N75435</u>	<u>003</u>	Oct 08, 2004
<u>NEURONTIN</u>					
<u>AB</u>	PFIZER PHARMS	<u>100MG</u>	<u>N20235</u>	<u>001</u>	Dec 30, 1993
<u>AB</u>		<u>300MG</u>	<u>N20235</u>	<u>002</u>	Dec 30, 1993
<u>AB</u>	+	<u>400MG</u>	<u>N20235</u>	<u>003</u>	Dec 30, 1993

SOLUTION; ORAL

NEURONTIN

+	PARKE DAVIS	250MG/5ML	N21129	001	Mar 02, 2000
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TABLET; ORAL

GABAPENTIN

<u>AB</u>	ACTAVIS ELIZABETH	<u>600MG</u>	<u>N75694</u>	<u>001</u>	Oct 21, 2004
<u>AB</u>		<u>800MG</u>	<u>N75694</u>	<u>002</u>	Oct 21, 2004
<u>AB</u>	APOTEX INC	<u>100MG</u>	<u>N77894</u>	<u>001</u>	Oct 10, 2006
<u>AB</u>		<u>300MG</u>	<u>N77894</u>	<u>002</u>	Oct 10, 2006
<u>AB</u>		<u>400MG</u>	<u>N77894</u>	<u>003</u>	Oct 10, 2006
<u>AB</u>		<u>600MG</u>	<u>N77661</u>	<u>004</u>	Sep 13, 2006
<u>AB</u>		<u>800MG</u>	<u>N77661</u>	<u>005</u>	Sep 13, 2006
<u>AB</u>	GLENMARK PHARMS	<u>600MG</u>	<u>N77662</u>	<u>001</u>	Aug 18, 2006
<u>AB</u>		<u>800MG</u>	<u>N77662</u>	<u>002</u>	Aug 18, 2006
<u>AB</u>	IVAX PHARMS	<u>100MG</u>	<u>N76017</u>	<u>001</u>	Apr 28, 2004
<u>AB</u>		<u>300MG</u>	<u>N76017</u>	<u>002</u>	Apr 28, 2004
<u>AB</u>		<u>400MG</u>	<u>N76017</u>	<u>003</u>	Apr 28, 2004
<u>AB</u>		<u>600MG</u>	<u>N76017</u>	<u>004</u>	Apr 29, 2005
<u>AB</u>		<u>800MG</u>	<u>N76017</u>	<u>005</u>	Apr 29, 2005
<u>AB</u>	RANBAXY	<u>600MG</u>	<u>N76605</u>	<u>001</u>	Sep 14, 2005
<u>AB</u>		<u>800MG</u>	<u>N76605</u>	<u>002</u>	Sep 14, 2005
<u>AB</u>	SANDOZ	<u>600MG</u>	<u>N76120</u>	<u>001</u>	Jan 27, 2006
<u>AB</u>		<u>600MG</u>	<u>N76877</u>	<u>001</u>	Jul 06, 2006
<u>AB</u>		<u>800MG</u>	<u>N76120</u>	<u>002</u>	Jan 27, 2006
<u>AB</u>		<u>800MG</u>	<u>N76877</u>	<u>002</u>	Jul 06, 2006
<u>AB</u>	SUN PHARM INDS LTD	<u>600MG</u>	<u>N77525</u>	<u>001</u>	Aug 24, 2006
<u>AB</u>		<u>800MG</u>	<u>N77525</u>	<u>002</u>	Aug 24, 2006
<u>AB</u>	TEVA	<u>600MG</u>	<u>N75827</u>	<u>001</u>	Dec 15, 2004
<u>AB</u>		<u>800MG</u>	<u>N75827</u>	<u>002</u>	Dec 15, 2004
<u>NEURONTIN</u>					
<u>AB</u>	PFIZER PHARMS	<u>600MG</u>	<u>N20882</u>	<u>001</u>	Oct 09, 1998
<u>AB</u>	+	<u>800MG</u>	<u>N20882</u>	<u>002</u>	Oct 09, 1998

GADOBENATE DIMEGLUMINE

INJECTABLE; INTRAVENOUS

MULTIHANCE

+	BRACCO	2.645GM/5ML (529MG/ML)	N21357	001	Nov 23, 2004
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27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 180 (of 370)

GADOBENATE DIMEGLUMINEINJECTABLE; INTRAVENOUS
MULTIHANCE

+ BRACCO	5.29GM/10ML (529MG/ML)	N21357 002	Nov 23, 2004
+	7.935GM/15ML (529MG/ML)	N21357 003	Nov 23, 2004
+	10.58GM/20ML (529MG/ML)	N21357 004	Nov 23, 2004
MULTIHANCE MULTIPACK			
+ BRACCO	26.45GM/50ML (529MG/ML)	N21358 001	Nov 23, 2004
+	52.9GM/100ML (529MG/ML)	N21358 002	Nov 23, 2004

GADODIAMIDEINJECTABLE; INJECTION
OMNISCAN

+ GE HEALTHCARE	287MG/ML	N20123 001	Jan 08, 1993
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GADOPENTETATE DIMEGLUMINEINJECTABLE; INJECTION
MAGNEVIST

+ BERLEX	469.01MG/ML	N19596 001	Jun 02, 1988
+	469.01MG/ML	N21037 001	Mar 10, 2000

GADOTERIDOLINJECTABLE; INJECTION
PROHANCE

+ BRACCO	279.3MG/ML	N20131 001	Nov 16, 1992
PROHANCE MULTIPACK			
+ BRACCO	279.3MG/ML	N21489 001	Oct 09, 2003

GADOVERSETAMIDEINJECTABLE; INJECTION
OPTIMARK

+ MALLINCKRODT	1654.5MG/5ML (330.9MG/ML)	N20937 001	Dec 08, 1999
+	3309MG/10ML (330.9MG/ML)	N20937 002	Dec 08, 1999
+	4963.5MG/15ML (330.9MG/ML)	N20937 003	Dec 08, 1999
+	6618MG/20ML (330.9MG/ML)	N20937 004	Dec 08, 1999
+	16.545GM/50ML (330.9MG/ML)	N20975 001	Dec 08, 1999
OPTIMARK IN PLASTIC CONTAINER			
+ MALLINCKRODT	1654.5MG/5ML (330.9MG/ML)	N20976 001	Dec 08, 1999
+	3309MG/10ML (330.9MG/ML)	N20976 002	Dec 08, 1999
+	4963.5MG/15ML (330.9MG/ML)	N20976 003	Dec 08, 1999
+	6618MG/20ML (330.9MG/ML)	N20976 004	Dec 08, 1999

GALANTAMINE HYDROBROMIDECAPSULE, EXTENDED RELEASE; ORAL
RAZADYNE ER

+ JANSSEN	EQ 8MG BASE	N21615 001	Apr 01, 2005
	EQ 16MG BASE	N21615 002	Apr 01, 2005
	EQ 24MG BASE	N21615 003	Apr 01, 2005

SOLUTION; ORAL
RAZADYNE

+ JANSSEN PHARMA	4MG/ML	N21224 001	Jun 22, 2001
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TABLET; ORAL
RAZADYNE

+ JANSSEN PHARMA	EQ 4MG BASE	N21169 001	Feb 28, 2001
	EQ 8MG BASE	N21169 002	Feb 28, 2001
	EQ 12MG BASE	N21169 003	Feb 28, 2001

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 181 (of 370)

GALLIUM CITRATE, GA-67

INJECTABLE; INJECTION				
GALLIUM CITRATE GA 67				
BS	BRISTOL MYERS SQUIBB	2mCi/ML	N17478	001
BS	MALLINCKRODT	2mCi/ML	N18058	001

GALLIUM NITRATE

INJECTABLE; INJECTION				
GANITE				
+	GENTA	25MG/ML	N19961	002 Jan 17, 1991

GANCICLOVIR

CAPSULE; ORAL				
GANCICLOVIR				
	RANBAXY	250MG	N76457	001 Jun 27, 2003
+		500MG	N76457	002 Jun 27, 2003

IMPLANT; IMPLANTATION				
VITRASERT				
+	BAUSCH AND LOMB	4.5MG	N20569	001 Mar 04, 1996

GANCICLOVIR SODIUM

INJECTABLE; INJECTION				
<u>CYTOVENE IV</u>				
<u>AP</u>	+ ROCHE PALO	<u>EQ 500MG BASE/VIAL</u>	<u>N19661</u>	<u>001</u> Jun 23, 1989
<u>GANCICLOVIR SODIUM</u>				
<u>AP</u>	BEDFORD	<u>EQ 500MG BASE/VIAL</u>	<u>N76222</u>	<u>001</u> Jul 16, 2003

GANIRELIX ACETATE

INJECTABLE; INJECTION				
GANIRELIX ACETATE INJECTION				
+	ORGANON USA INC	EQ 250UGM BASE/0.5ML	N21057	001 Jul 29, 1999

GATIFLOXACIN

SOLUTION/DROPS; OPHTHALMIC				
+	ALLERGAN	0.3%	N21493	001 Mar 28, 2003

GEFITINIB

TABLET; ORAL				
IRESSA				
+	ASTRAZENECA	250MG	N21399	001 May 05, 2003

GEMCITABINE HYDROCHLORIDE

INJECTABLE; INJECTION				
GEMZAR				
+	LILLY	EQ 200MG BASE/VIAL	N20509	001 May 15, 1996
+		EQ 1GM BASE/VIAL	N20509	002 May 15, 1996

GEMFIBROZIL

TABLET; ORAL				
<u>GEMFIBROZIL</u>				
<u>AB</u>	INVAGEN PHARMS	<u>600MG</u>	<u>N77836</u>	<u>001</u> Jul 27, 2006
<u>AB</u>	MYLAN	<u>600MG</u>	<u>N74452</u>	<u>001</u> Feb 16, 1995
<u>AB</u>	SANDOZ	<u>600MG</u>	<u>N74615</u>	<u>001</u> Sep 29, 1995
<u>AB</u>	TEVA	<u>600MG</u>	<u>N74256</u>	<u>001</u> Oct 31, 1993
<u>AB</u>	TORPHARM	<u>600MG</u>	<u>N75034</u>	<u>001</u> Jul 20, 1998
<u>AB</u>	WATSON LABS	<u>600MG</u>	<u>N74442</u>	<u>001</u> Apr 28, 1995
<u>LOPID</u>				
<u>AB</u>	+ PFIZER PHARMS	<u>600MG</u>	<u>N18422</u>	<u>003</u> Nov 20, 1986

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 182 (of 370)

GEMIFLOXACIN MESYLATETABLET; ORAL
FACTIVE

+ OSCIENT

EQ 320MG BASE

N21158 001

Apr 04, 2003

GEMTUZUMAB OZOGAMICININJECTABLE; INJECTION
MYLOTARG

+ WYETH PHARMS INC

5MG/VIAL

N21174 001

May 17, 2000

GENTAMICIN SULFATE

CREAM; TOPICAL

GENTAMICINAT PERRIGO NEW YORKEQ 0.1% BASEN62307 001GENTAMICIN SULFATEAT + FOUGERAEQ 0.1% BASEN62531 001

Jul 05, 1984

AT TAROEQ 0.1% BASEN62427 001

May 26, 1983

INJECTABLE; INJECTION

GENTAMICIN SULFATEAP + ABRAXIS PHARMEQ 10MG BASE/MLN62356 001

Mar 04, 1982

AP +EQ 40MG BASE/MLN62356 002

Mar 04, 1982

APEQ 40MG BASE/MLN62366 001

Aug 04, 1983

AP HOSPIRAEQ 10MG BASE/MLN62420 001

Aug 15, 1983

APEQ 10MG BASE/MLN62612 004

Feb 20, 1986

APEQ 40MG BASE/MLN62420 002

Aug 15, 1983

AP PHARM SPECEQ 40MG BASE/MLN62340 001

Mar 28, 1983

GENTAMICIN SULFATE IN SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINERAP + B BRAUNEQ 0.8MG BASE/MLN62814 001

Aug 28, 1987

AP +EQ 1.2MG BASE/MLN62814 002

Aug 28, 1987

AP +EQ 1.4MG BASE/MLN62814 003

Aug 28, 1987

AP +EQ 1.6MG BASE/MLN62814 004

Aug 28, 1987

AP +EQ 1.8MG BASE/MLN62814 005

Aug 28, 1987

AP +EQ 2MG BASE/MLN62814 006

Aug 28, 1987

AP +EQ 2.4MG BASE/MLN62814 007

Aug 28, 1987

AP +EQ 40MG BASE/100MLN62814 008

Aug 28, 1987

AP +EQ 60MG BASE/100MLN62814 009

Aug 28, 1987

AP +EQ 70MG BASE/100MLN62814 010

Aug 28, 1987

AP +EQ 80MG BASE/100MLN62814 011

Aug 28, 1987

AP +EQ 90MG BASE/100MLN62814 012

Aug 28, 1987

AP +EQ 100MG BASE/100MLN62814 013

Aug 28, 1987

AP +EQ 120MG BASE/100MLN62814 014

Aug 28, 1987

AP HOSPIRAEQ 1.2MG BASE/MLN62414 001

Aug 15, 1983

APEQ 1.4MG BASE/MLN62414 002

Aug 15, 1983

APEQ 1.6MG BASE/MLN62414 003

Aug 15, 1983

APEQ 1.8MG BASE/MLN62414 004

Aug 15, 1983

APEQ 2MG BASE/MLN62414 005

Aug 15, 1983

APEQ 60MG BASE/100MLN62414 006

Aug 15, 1983

APEQ 70MG BASE/100MLN62414 007

Aug 15, 1983

APEQ 80MG BASE/100MLN62414 008

Aug 15, 1983

APEQ 90MG BASE/100MLN62414 009

Aug 15, 1983

APEQ 100MG BASE/100MLN62414 010

Aug 15, 1983

ISOTONIC GENTAMICIN SULFATE IN PLASTIC CONTAINERAP BAXTER HLTHCAREEQ 0.8MG BASE/MLN62373 001

Sep 07, 1982

APEQ 1.2MG BASE/MLN62373 007

Sep 07, 1982

APEQ 1.6MG BASE/MLN62373 008

Sep 07, 1982

APEQ 2MG BASE/MLN62373 009

Sep 07, 1982

APEQ 2.4MG BASE/MLN62373 010

Sep 07, 1982

APEQ 40MG BASE/100MLN62373 003

Sep 07, 1982

APEQ 60MG BASE/100MLN62373 004

Sep 07, 1982

APEQ 80MG BASE/100MLN62373 002

Sep 07, 1982

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 183 (of 370)

GENTAMICIN SULFATE

INJECTABLE; INJECTION

ISOTONIC GENTAMICIN SULFATE IN PLASTIC CONTAINER

<u>AP</u>	BAXTER HLTHCARE	<u>EQ 100MG BASE/100ML</u>	<u>N62373</u>	<u>005</u>	Sep 07, 1982
<u>AP</u>		<u>EQ 120MG BASE/100ML</u>	<u>N62373</u>	<u>006</u>	Sep 07, 1982

OINTMENT; OPHTHALMIC

GENTAMICIN SULFATE

<u>AT</u>	+ AKORN	<u>EQ 0.3% BASE</u>	<u>N64093</u>	<u>001</u>	Aug 31, 1995
<u>AT</u>	ALTANA	<u>EQ 0.3% BASE</u>	<u>N65024</u>	<u>001</u>	Jul 30, 2004

OINTMENT; TOPICAL

GENTAMICIN

<u>AT</u>	+ PERRIGO NEW YORK	<u>EQ 0.1% BASE</u>	<u>N62351</u>	<u>001</u>	Feb 18, 1982
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GENTAMICIN SULFATE

<u>AT</u>	FOUGERA	<u>EQ 0.1% BASE</u>	<u>N62533</u>	<u>001</u>	Oct 05, 1984
<u>AT</u>	TARO	<u>EQ 0.1% BASE</u>	<u>N62477</u>	<u>001</u>	Dec 23, 1983

SOLUTION/DROPS; OPHTHALMIC

GENOPTIC

<u>AT</u>	ALLERGAN	<u>EQ 0.3% BASE</u>	<u>N62452</u>	<u>001</u>	Oct 10, 1984
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GENTAK

<u>AT</u>	AKORN	<u>EQ 0.3% BASE</u>	<u>N64163</u>	<u>001</u>	Oct 12, 2001
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GENTAMICIN SULFATE

<u>AT</u>	AKORN	<u>EQ 0.3% BASE</u>	<u>N62635</u>	<u>001</u>	Jan 08, 1987
<u>AT</u>	ALTANA	<u>EQ 0.3% BASE</u>	<u>N65121</u>	<u>001</u>	Jan 30, 2004
<u>AT</u>	+ BAUSCH AND LOMB	<u>EQ 0.3% BASE</u>	<u>N64048</u>	<u>001</u>	May 11, 1994
<u>AT</u>	FALCON PHARMS	<u>EQ 0.3% BASE</u>	<u>N62196</u>	<u>001</u>	

GENTAMICIN SULFATE; PREDNISOLONE ACETATE

OINTMENT; OPHTHALMIC

PRED-G

	+ ALLERGAN	EQ 0.3% BASE; 0.6%	N50612	001	Dec 01, 1989
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SUSPENSION/DROPS; OPHTHALMIC

	+ ALLERGAN	EQ 0.3% BASE; 1%	N50586	001	Jun 10, 1988
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GLATIRAMER ACETATE

FOR SOLUTION; SUBCUTANEOUS

COPAXONE

	+ TEVA	20MG/VIAL	N20622	001	Dec 20, 1996
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INJECTABLE; SUBCUTANEOUS

COPAXONE

	+ TEVA	20MG/ML	N20622	002	Feb 12, 2002
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GLIMEPIRIDE

TABLET; ORAL

AMARYL

<u>AB</u>	+ SANOFI AVENTIS US	<u>1MG</u>	<u>N20496</u>	<u>001</u>	Nov 30, 1995
<u>AB</u>		<u>2MG</u>	<u>N20496</u>	<u>002</u>	Nov 30, 1995
<u>AB</u>		<u>4MG</u>	<u>N20496</u>	<u>003</u>	Nov 30, 1995

GLIMEPIRIDE

<u>AB</u>	COBALT	<u>1MG</u>	<u>N77280</u>	<u>001</u>	Feb 03, 2006
<u>AB</u>		<u>2MG</u>	<u>N77280</u>	<u>002</u>	Feb 03, 2006
<u>AB</u>		<u>4MG</u>	<u>N77280</u>	<u>003</u>	Feb 03, 2006
<u>AB</u>	COREPHARMA	<u>1MG</u>	<u>N77274</u>	<u>001</u>	Oct 06, 2005
<u>AB</u>		<u>2MG</u>	<u>N77274</u>	<u>002</u>	Oct 06, 2005
<u>AB</u>		<u>4MG</u>	<u>N77274</u>	<u>003</u>	Oct 06, 2005
<u>AB</u>	DR REDDYS LABS LTD	<u>1MG</u>	<u>N77091</u>	<u>001</u>	Oct 06, 2005
<u>AB</u>		<u>2MG</u>	<u>N77091</u>	<u>002</u>	Oct 06, 2005
<u>AB</u>		<u>4MG</u>	<u>N77091</u>	<u>003</u>	Oct 06, 2005
<u>AB</u>	GENPHARM	<u>1MG</u>	<u>N77486</u>	<u>001</u>	Feb 10, 2006

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 184 (of 370)

GLIMEPIRIDE

TABLET; ORAL

GLIMEPIRIDE

<u>AB</u>	GENPHARM	<u>2MG</u>	<u>N77486</u>	<u>002</u>	Feb 10, 2006
<u>AB</u>		<u>4MG</u>	<u>N77486</u>	<u>003</u>	Feb 10, 2006
<u>AB</u>	INVAGEN PHARMS	<u>1MG</u>	<u>N77295</u>	<u>001</u>	Oct 06, 2005
<u>AB</u>		<u>2MG</u>	<u>N77295</u>	<u>002</u>	Oct 06, 2005
<u>AB</u>		<u>4MG</u>	<u>N77295</u>	<u>003</u>	Oct 06, 2005
<u>AB</u>	MYLAN	<u>1MG</u>	<u>N77624</u>	<u>001</u>	Nov 28, 2005
<u>AB</u>		<u>2MG</u>	<u>N77624</u>	<u>002</u>	Nov 28, 2005
<u>AB</u>		<u>4MG</u>	<u>N77624</u>	<u>003</u>	Nov 28, 2005
<u>AB</u>	PAR PHARM	<u>1MG</u>	<u>N77370</u>	<u>001</u>	Dec 23, 2005
<u>AB</u>		<u>2MG</u>	<u>N77370</u>	<u>002</u>	Dec 23, 2005
<u>AB</u>		<u>4MG</u>	<u>N77370</u>	<u>003</u>	Dec 23, 2005
<u>AB</u>		<u>8MG</u>	<u>N77370</u>	<u>004</u>	Dec 23, 2005
<u>AB</u>	RANBAXY	<u>1MG</u>	<u>N76875</u>	<u>001</u>	Oct 06, 2005
<u>AB</u>		<u>2MG</u>	<u>N76875</u>	<u>002</u>	Oct 06, 2005
<u>AB</u>		<u>4MG</u>	<u>N76875</u>	<u>003</u>	Oct 06, 2005
<u>AB</u>		<u>8MG</u>	<u>N76875</u>	<u>004</u>	Oct 06, 2005
		3MG	N77366	001	Oct 06, 2005
		6MG	N77366	002	Oct 06, 2005
<u>AB</u>	TEVA	<u>1MG</u>	<u>N76802</u>	<u>001</u>	Oct 06, 2005
<u>AB</u>		<u>2MG</u>	<u>N76802</u>	<u>002</u>	Oct 06, 2005
<u>AB</u>		<u>4MG</u>	<u>N76802</u>	<u>003</u>	Oct 06, 2005

GLIMEPIRIDE; PIOGLITAZONE HYDROCHLORIDE

TABLET; ORAL

DUETACT

TAKEDA GLOBAL	2MG; 30MG	N21925	001	Jul 28, 2006
+	4MG; 30MG	N21925	002	Jul 28, 2006

GLIMEPIRIDE; ROSIGLITAZONE MALEATE

TABLET; ORAL

AVANDARYL

SB PHARMCO	1MG; 4MG	N21700	001	Nov 23, 2005
	2MG; 4MG	N21700	002	Nov 23, 2005
+	4MG; 4MG	N21700	003	Nov 23, 2005

GLIPIZIDE

TABLET; ORAL

GLIPIZIDE

<u>AB</u>	ALPHAPHARM	<u>5MG</u>	<u>N74438</u>	<u>001</u>	Jun 20, 1995
<u>AB</u>		<u>10MG</u>	<u>N74438</u>	<u>002</u>	Jun 20, 1995
<u>AB</u>	CARACO	<u>5MG</u>	<u>N77820</u>	<u>001</u>	Jul 11, 2006
<u>AB</u>		<u>10MG</u>	<u>N77820</u>	<u>002</u>	Jul 11, 2006
<u>AB</u>	IVAX PHARMS	<u>5MG</u>	<u>N74497</u>	<u>001</u>	Aug 31, 1995
<u>AB</u>		<u>10MG</u>	<u>N74497</u>	<u>002</u>	Aug 31, 1995
<u>AB</u>	KALI LABS	<u>5MG</u>	<u>N74550</u>	<u>001</u>	Sep 11, 1997
<u>AB</u>		<u>10MG</u>	<u>N74550</u>	<u>002</u>	Sep 11, 1997
<u>AB</u>	MYLAN	<u>5MG</u>	<u>N74226</u>	<u>001</u>	May 10, 1994
<u>AB</u>		<u>10MG</u>	<u>N74226</u>	<u>002</u>	May 10, 1994
<u>AB</u>	PLIVA	<u>5MG</u>	<u>N74619</u>	<u>001</u>	Apr 04, 1997
<u>AB</u>		<u>10MG</u>	<u>N74619</u>	<u>002</u>	Apr 04, 1997
<u>AB</u>	SANDOZ	<u>5MG</u>	<u>N74305</u>	<u>001</u>	Apr 07, 1995
<u>AB</u>		<u>5MG</u>	<u>N74542</u>	<u>001</u>	Jun 20, 1995
<u>AB</u>		<u>10MG</u>	<u>N74305</u>	<u>002</u>	Apr 07, 1995
<u>AB</u>		<u>10MG</u>	<u>N74542</u>	<u>002</u>	Jun 20, 1995
<u>AB</u>	TEVA	<u>5MG</u>	<u>N74387</u>	<u>001</u>	Mar 04, 1996
<u>AB</u>		<u>10MG</u>	<u>N74387</u>	<u>002</u>	Mar 04, 1996
<u>AB</u>	TORPHARM	<u>5MG</u>	<u>N75795</u>	<u>001</u>	Jun 13, 2001

PRESCRIPTION DRUG PRODUCT LIST

3 - 185 (of 370)

GLIPIZIDE

TABLET; ORAL

GLIPIZIDE

<u>AB</u>	TORPHARM	<u>10MG</u>	<u>N75795</u>	<u>002</u>	Jun 13, 2001
<u>AB</u>	WATSON LABS	<u>5MG</u>	<u>N74223</u>	<u>001</u>	Feb 27, 1995
<u>AB</u>		<u>5MG</u>	<u>N74370</u>	<u>001</u>	Nov 22, 1994
<u>AB</u>		<u>10MG</u>	<u>N74223</u>	<u>002</u>	Feb 27, 1995
<u>AB</u>		<u>10MG</u>	<u>N74370</u>	<u>002</u>	Nov 22, 1994

GLUCOTROL

<u>AB</u>	PFIZER	<u>5MG</u>	<u>N17783</u>	<u>001</u>	May 08, 1984
<u>AB</u>	+	<u>10MG</u>	<u>N17783</u>	<u>002</u>	May 08, 1984

TABLET, EXTENDED RELEASE; ORAL

GLIPIZIDE

<u>AB</u>	WATSON LABS	<u>2.5MG</u>	<u>N76467</u>	<u>003</u>	Mar 27, 2006
<u>AB</u>		<u>5MG</u>	<u>N76467</u>	<u>001</u>	Sep 08, 2003
<u>AB</u>		<u>10MG</u>	<u>N76467</u>	<u>002</u>	Nov 07, 2003

GLUCOTROL XL

<u>AB</u>	PFIZER	<u>2.5MG</u>	<u>N20329</u>	<u>003</u>	Aug 10, 1999
<u>AB</u>		<u>5MG</u>	<u>N20329</u>	<u>001</u>	Apr 26, 1994
<u>AB</u>	+	<u>10MG</u>	<u>N20329</u>	<u>002</u>	Apr 26, 1994

GLIPIZIDE; METFORMIN HYDROCHLORIDE

TABLET; ORAL

GLIPIZIDE AND METFORMIN HYDROCHLORIDE

<u>AB</u>	COREPHARMA	<u>2.5MG;250MG</u>	<u>N77507</u>	<u>001</u>	Oct 27, 2005
<u>AB</u>		<u>2.5MG;500MG</u>	<u>N77507</u>	<u>002</u>	Oct 27, 2005
<u>AB</u>		<u>5MG;500MG</u>	<u>N77507</u>	<u>003</u>	Oct 27, 2005
<u>AB</u>	TEVA PHARMS	<u>2.5MG;250MG</u>	<u>N77270</u>	<u>001</u>	Oct 28, 2005
<u>AB</u>		<u>2.5MG;500MG</u>	<u>N77270</u>	<u>002</u>	Oct 28, 2005
<u>AB</u>		<u>5MG;500MG</u>	<u>N77270</u>	<u>003</u>	Oct 28, 2005
<u>AB</u>	<u>METAGLIP</u>				
<u>AB</u>	BRISTOL MYERS SQUIBB	<u>2.5MG;250MG</u>	<u>N21460</u>	<u>001</u>	Oct 21, 2002
<u>AB</u>		<u>2.5MG;500MG</u>	<u>N21460</u>	<u>002</u>	Oct 21, 2002
<u>AB</u>	+	<u>5MG;500MG</u>	<u>N21460</u>	<u>003</u>	Oct 21, 2002

GLUCAGON HYDROCHLORIDE RECOMBINANT

INJECTABLE; INJECTION

GLUCAGON

+	NOVO NORDISK	EQ 1MG BASE/VIAL	N20918	001	Jun 22, 1998
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GLUCAGON RECOMBINANT

INJECTABLE; INJECTION

GLUCAGON

+	LILLY	1MG/VIAL	N20928	001	Sep 11, 1998
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GLYBURIDE

TABLET; ORAL

DIABETA

BX	SANOFI AVENTIS US	1.25MG	N17532	001	May 01, 1984
BX		2.5MG	N17532	002	May 01, 1984
BX	+	5MG	N17532	003	May 01, 1984

GLYBURIDE

<u>AB</u>	ACTAVIS TOTOWA	<u>1.5MG</u>	<u>N75947</u>	<u>001</u>	Nov 14, 2002
<u>AB</u>		<u>3MG</u>	<u>N75947</u>	<u>002</u>	Nov 14, 2002
<u>AB</u>		<u>6MG</u>	<u>N75947</u>	<u>003</u>	Nov 14, 2002
<u>AB</u>	COREPHARMA	<u>1.25MG</u>	<u>N76257</u>	<u>001</u>	Jun 27, 2002
<u>AB</u>		<u>2.5MG</u>	<u>N76257</u>	<u>002</u>	Jun 27, 2002
<u>AB</u>		<u>5MG</u>	<u>N76257</u>	<u>003</u>	Jun 27, 2002
<u>AB</u>	TEVA	<u>1.25MG</u>	<u>N74388</u>	<u>001</u>	Aug 29, 1995

PRESCRIPTION DRUG PRODUCT LIST

3 - 186 (of 370)

GLYBURIDE

TABLET; ORAL					
<u>GLYBURIDE</u>					
<u>AB</u>	TEVA	<u>2.5MG</u>	<u>N74388</u>	<u>002</u>	Aug 29, 1995
<u>AB</u>		<u>5MG</u>	<u>N74388</u>	<u>003</u>	Aug 29, 1995
<u>GLYBURIDE (MICRONIZED)</u>					
<u>AB</u>	CLONMEL HLTHCARE	<u>1.5MG</u>	<u>N74591</u>	<u>001</u>	Dec 22, 1997
<u>AB</u>		<u>3MG</u>	<u>N74591</u>	<u>002</u>	Dec 22, 1997
<u>AB</u>		<u>4.5MG</u>	<u>N74591</u>	<u>003</u>	Dec 22, 1997
<u>AB</u>		<u>6MG</u>	<u>N74591</u>	<u>004</u>	Dec 22, 1997
<u>AB</u>	HIKMA	<u>1.5MG</u>	<u>N75890</u>	<u>001</u>	Jul 31, 2003
<u>AB</u>		<u>3MG</u>	<u>N75890</u>	<u>002</u>	Jul 31, 2003
<u>AB</u>		<u>6MG</u>	<u>N75890</u>	<u>003</u>	Jul 31, 2003
<u>AB</u>	MYLAN	<u>1.5MG</u>	<u>N74792</u>	<u>001</u>	Jun 26, 1998
<u>AB</u>		<u>3MG</u>	<u>N74792</u>	<u>002</u>	Jun 26, 1998
<u>AB</u>		<u>6MG</u>	<u>N74792</u>	<u>003</u>	Aug 17, 1999
<u>AB</u>	SANDOZ	<u>1.5MG</u>	<u>N75174</u>	<u>001</u>	Jun 22, 1998
<u>AB</u>		<u>3MG</u>	<u>N75174</u>	<u>002</u>	Jun 22, 1998
<u>AB</u>	TEVA	<u>1.5MG</u>	<u>N74686</u>	<u>001</u>	Apr 20, 1999
<u>AB</u>		<u>3MG</u>	<u>N74686</u>	<u>002</u>	Apr 20, 1999
<u>AB</u>		<u>4.5MG</u>	<u>N74686</u>	<u>003</u>	Apr 20, 1999
<u>AB</u>		<u>6MG</u>	<u>N74686</u>	<u>004</u>	Apr 20, 1999
<u>GLYNASE</u>					
<u>AB</u>	PHARMACIA AND UPJOHN	<u>1.5MG</u>	<u>N20051</u>	<u>001</u>	Mar 04, 1992
<u>AB</u>		<u>3MG</u>	<u>N20051</u>	<u>002</u>	Mar 04, 1992
<u>AB</u>	+	<u>6MG</u>	<u>N20051</u>	<u>004</u>	Sep 24, 1993
<u>MICRONASE</u>					
<u>AB</u>	PHARMACIA AND UPJOHN	<u>1.25MG</u>	<u>N17498</u>	<u>001</u>	May 01, 1984
<u>AB</u>		<u>2.5MG</u>	<u>N17498</u>	<u>002</u>	May 01, 1984
<u>AB</u>	+	<u>5MG</u>	<u>N17498</u>	<u>003</u>	May 01, 1984

GLYBURIDE; METFORMIN HYDROCHLORIDE

TABLET; ORAL					
<u>GLUCOVANCE</u>					
<u>AB</u>	BRISTOL MYERS SQUIBB	<u>1.25MG;250MG</u>	<u>N21178</u>	<u>001</u>	Jul 31, 2000
<u>AB</u>		<u>2.5MG;500MG</u>	<u>N21178</u>	<u>002</u>	Jul 31, 2000
<u>AB</u>	+	<u>5MG;500MG</u>	<u>N21178</u>	<u>003</u>	Jul 31, 2000
<u>GLYBURIDE AND METFORMIN HYDROCHLORIDE</u>					
<u>AB</u>	ACTAVIS ELIZABETH	<u>1.25MG;250MG</u>	<u>N76716</u>	<u>001</u>	Jun 28, 2005
<u>AB</u>		<u>2.5MG;500MG</u>	<u>N76716</u>	<u>002</u>	Jun 28, 2005
<u>AB</u>		<u>5MG;500MG</u>	<u>N76716</u>	<u>003</u>	Jun 28, 2005
<u>AB</u>	COREPHARMA	<u>1.25MG;250MG</u>	<u>N76731</u>	<u>001</u>	Nov 19, 2004
<u>AB</u>		<u>2.5MG;500MG</u>	<u>N76731</u>	<u>002</u>	Nov 19, 2004
<u>AB</u>		<u>5MG;500MG</u>	<u>N76731</u>	<u>003</u>	Nov 19, 2004
<u>AB</u>	IVAX PHARMS	<u>1.25MG;250MG</u>	<u>N76345</u>	<u>001</u>	Feb 18, 2004
<u>AB</u>		<u>2.5MG;500MG</u>	<u>N76345</u>	<u>002</u>	Feb 18, 2004
<u>AB</u>		<u>5MG;500MG</u>	<u>N76345</u>	<u>003</u>	Feb 18, 2004
<u>AB</u>	TEVA	<u>1.25MG;250MG</u>	<u>N76821</u>	<u>001</u>	Jan 27, 2005
<u>AB</u>		<u>2.5MG;500MG</u>	<u>N76821</u>	<u>002</u>	Jan 27, 2005
<u>AB</u>		<u>5MG;500MG</u>	<u>N76821</u>	<u>003</u>	Jan 27, 2005

GLYCINE

SOLUTION; IRRIGATION					
<u>AMINOACETIC ACID 1.5% IN PLASTIC CONTAINER</u>					
<u>AT</u>	BAXTER HLTHCARE	<u>1.5GM/100ML</u>	<u>N17865</u>	<u>001</u>	
<u>GLYCINE 1.5% IN PLASTIC CONTAINER</u>					
<u>AT</u>	B BRAUN	<u>1.5GM/100ML</u>	<u>N16784</u>	<u>001</u>	
<u>AT</u>	HOSPIRA	<u>1.5GM/100ML</u>	<u>N17633</u>	<u>001</u>	
<u>AT</u>		<u>1.5GM/100ML</u>	<u>N18315</u>	<u>001</u>	

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 187 (of 370)

GLYCOPYRROLATE

INJECTABLE; INJECTION

GLYCOPYRROLATE

<u>AP</u>	LUITPOLD	<u>0.2MG/ML</u>	<u>N89335</u>	<u>001</u>	Jul 23, 1986
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ROBINUL

<u>AP</u>	+ BAXTER HLTHCARE	<u>0.2MG/ML</u>	<u>N17558</u>	<u>001</u>	
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TABLET; ORAL

GLYCOPYRROLATE

<u>AA</u>	COREPHARMA	<u>1MG</u>	<u>N40568</u>	<u>001</u>	Dec 22, 2004
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<u>AA</u>		<u>2MG</u>	<u>N40568</u>	<u>002</u>	Dec 22, 2004
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<u>AA</u>	KALI LABS	<u>1MG</u>	<u>N40653</u>	<u>001</u>	Aug 31, 2006
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<u>AA</u>		<u>2MG</u>	<u>N40653</u>	<u>002</u>	Aug 31, 2006
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ROBINUL

<u>AA</u>	+ SCIELE PHARMA INC	<u>1MG</u>	<u>N12827</u>	<u>001</u>	
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ROBINUL FORTE

<u>AA</u>	+ SCIELE PHARMA INC	<u>2MG</u>	<u>N12827</u>	<u>002</u>	
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GONADOTROPIN, CHORIONIC

INJECTABLE; INJECTION

CHORIONIC GONADOTROPIN

<u>AP</u>	+ ABRAXIS PHARM	<u>10,000 UNITS/VIAL</u>	<u>N17067</u>	<u>002</u>	
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<u>AP</u>	+ WATSON LABS (UTAH)	<u>10,000 UNITS/VIAL</u>	<u>N17016</u>	<u>007</u>	
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PREGNYL

<u>AP</u>	+ ORGANON USA INC	<u>10,000 UNITS/VIAL</u>	<u>N17692</u>	<u>001</u>	
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GOSERELIN ACETATE

IMPLANT; IMPLANTATION

ZOLADEX

	+ ASTRAZENECA	EQ 3.6MG BASE	N19726	001	Dec 29, 1989
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		EQ 10.8MG BASE	N20578	001	Jan 11, 1996
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GRAMICIDIN; NEOMYCIN SULFATE; POLYMYXIN B SULFATE

SOLUTION/DROPS; OPHTHALMIC

NEOMYCIN AND POLYMYXIN B SULFATES AND GRAMICIDIN

<u>AT</u>	+ BAUSCH AND LOMB	<u>0.025MG/ML;EQ 1.75MG BASE/ML;10,000 UNITS/ML</u>	<u>N64047</u>	<u>001</u>	Jan 31, 1996
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<u>AT</u>	PHARMAFORCE	<u>0.025MG/ML;EQ 1.75MG BASE/ML;10,000 UNITS/ML</u>	<u>N65187</u>	<u>001</u>	Oct 28, 2005
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NEOSPORIN

<u>AT</u>	+ MONARCH PHARMS	<u>0.025MG/ML;EQ 1.75MG BASE/ML;10,000 UNITS/ML</u>	<u>N60582</u>	<u>001</u>	
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GRANISETRON HYDROCHLORIDE

INJECTABLE; INJECTION

KYTRIL

	+ ROCHE	EQ 0.1MG BASE/ML (EQ 0.1MG BASE/ML)	N20239	003	Sep 17, 2004
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		EQ 1MG BASE/ML (EQ 1MG BASE/ML)	N20239	004	Mar 11, 1994
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		EQ 4MG BASE/4ML (EQ 1MG BASE /ML)	N20239	002	Mar 11, 1994
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SOLUTION; ORAL

KYTRIL

	+ ROCHE	EQ 2MG BASE/10ML	N21238	001	Jun 27, 2001
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TABLET; ORAL

KYTRIL

	+ ROCHE	EQ 1MG BASE	N20305	001	Mar 16, 1995
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GRISEOFULVIN, MICROCRYSTALLINE

SUSPENSION; ORAL

GRIFULVIN V

<u>AB</u>	+ J AND J	<u>125MG/5ML</u>	<u>N62483</u>	<u>001</u>	Jan 26, 1984
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27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 188 (of 370)

GRISEOFULVIN, MICROCRYSTALLINE

SUSPENSION; ORAL

GRISEOFULVIN

<u>AB</u>	STIEFEL	<u>125MG/5ML</u>	<u>N65200</u>	<u>001</u>	Mar 02, 2005
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TABLET; ORAL

FULVICIN-U/F

<u>AB</u>	+	SCHERING	<u>250MG</u>	<u>N60569</u>	<u>002</u>
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<u>AB</u>	+		<u>500MG</u>	<u>N60569</u>	<u>001</u>
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GRIFULVIN V

<u>AB</u>	J AND J	<u>250MG</u>	<u>N62279</u>	<u>002</u>	
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<u>AB</u>		<u>500MG</u>	<u>N62279</u>	<u>003</u>	
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	125MG	N62279	001		
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GRISEOFULVIN, ULTRAMICROCRYSTALLINE

TABLET; ORAL

GRIS-PEG

PEDINOL	125MG	N50475	001
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+	250MG	N50475	002
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GUANABENZ ACETATE

TABLET; ORAL

GUANABENZ ACETATE

<u>AB</u>	IVAX PHARMS	<u>EQ 4MG BASE</u>	<u>N74149</u>	<u>001</u>	Apr 07, 1995
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<u>AB</u>		<u>EQ 8MG BASE</u>	<u>N74149</u>	<u>002</u>	Apr 07, 1995
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<u>AB</u>	SANDOZ	<u>EQ 4MG BASE</u>	<u>N74517</u>	<u>001</u>	Sep 30, 1998
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<u>AB</u>		<u>EQ 8MG BASE</u>	<u>N74517</u>	<u>002</u>	Sep 30, 1998
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<u>AB</u>	TEVA PHARMS	<u>EQ 4MG BASE</u>	<u>N74267</u>	<u>001</u>	Jun 01, 1994
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<u>AB</u>		<u>EQ 8MG BASE</u>	<u>N74267</u>	<u>002</u>	Jun 01, 1994
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<u>AB</u>	WATSON LABS	<u>EQ 4MG BASE</u>	<u>N74025</u>	<u>001</u>	Feb 28, 1994
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<u>AB</u>	+	<u>EQ 8MG BASE</u>	<u>N74025</u>	<u>002</u>	Feb 28, 1994
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GUANFACINE HYDROCHLORIDE

TABLET; ORAL

GUANFACINE HYDROCHLORIDE

<u>AB</u>	ACTAVIS TOTOWA	<u>EQ 1MG BASE</u>	<u>N74673</u>	<u>001</u>	Feb 28, 1997
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<u>AB</u>		<u>EQ 2MG BASE</u>	<u>N74673</u>	<u>002</u>	Feb 28, 1997
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<u>AB</u>	GENPHARM	<u>EQ 1MG BASE</u>	<u>N75109</u>	<u>001</u>	Nov 25, 1998
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<u>AB</u>		<u>EQ 2MG BASE</u>	<u>N75109</u>	<u>002</u>	Nov 25, 1998
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<u>AB</u>	MYLAN	<u>EQ 1MG BASE</u>	<u>N74796</u>	<u>001</u>	Jan 27, 1997
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<u>AB</u>		<u>EQ 2MG BASE</u>	<u>N74796</u>	<u>002</u>	Jan 27, 1997
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<u>AB</u>	WATSON LABS	<u>EQ 1MG BASE</u>	<u>N74145</u>	<u>001</u>	Oct 17, 1995
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<u>AB</u>		<u>EQ 1MG BASE</u>	<u>N74762</u>	<u>001</u>	Jun 25, 1997
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<u>AB</u>		<u>EQ 2MG BASE</u>	<u>N74145</u>	<u>002</u>	Oct 17, 1995
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<u>AB</u>		<u>EQ 2MG BASE</u>	<u>N74762</u>	<u>002</u>	Jun 25, 1997
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TENEX

<u>AB</u>	DR REDDYS LABS INC	<u>EQ 1MG BASE</u>	<u>N19032</u>	<u>001</u>	Oct 27, 1986
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<u>AB</u>	+	<u>EQ 2MG BASE</u>	<u>N19032</u>	<u>002</u>	Nov 07, 1988
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GUANIDINE HYDROCHLORIDE

TABLET; ORAL

GUANIDINE HYDROCHLORIDE

SCHERING	125MG	N01546	001
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HALCINONIDE

CREAM; TOPICAL

HALOG

+	WESTWOOD SQUIBB	0.1%	N17556	001
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27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 189 (of 370)

HALCINONIDE

OINTMENT; TOPICAL

HALOG

+ WESTWOOD SQUIBB 0.1% N17824 001

SOLUTION; TOPICAL

HALOG

WESTWOOD SQUIBB 0.1% N17823 001

HALOBETASOL PROPIONATE

CREAM; TOPICAL

HALOBETASOL PROPIONATEAB ALTANA 0.05% N77001 001 Dec 16, 2004AB PERRIGO ISRAEL 0.05% N77123 001 Dec 16, 2004AB TARO 0.05% N77227 001 Aug 04, 2005ULTRAVATEAB + WESTWOOD SQUIBB 0.05% N19967 001 Dec 27, 1990

OINTMENT; TOPICAL

HALOBETASOL PROPIONATEAB ACTAVIS MID ATLANTIC 0.05% N77109 001 Jun 14, 2005AB ALTANA 0.05% N76903 001 Dec 16, 2004AB G AND W LABS 0.05% N77721 001 Sep 07, 2006AB PERRIGO 0.05% N76872 001 Dec 16, 2004AB TARO 0.05% N76994 001 Dec 16, 2004ULTRAVATEAB + WESTWOOD SQUIBB 0.05% N19968 001 Dec 17, 1990HALOPERIDOL

TABLET; ORAL

HALOPERIDOLAB MYLAN 0.5MG N70276 001 Jun 10, 1986AB 1MG N70277 001 Jun 10, 1986AB 2MG N70278 001 Jun 10, 1986AB 5MG N70279 001 Jun 10, 1986AB PAR PHARM 0.5MG N71233 001 Nov 03, 1986AB 1MG N71234 001 Nov 03, 1986AB 2MG N71235 001 Nov 03, 1986AB 5MG N71236 001 Nov 03, 1986AB 10MG N71237 001 Jul 20, 1987AB SANDOZ 0.5MG N71206 001 Nov 17, 1986AB 1MG N71207 001 Nov 17, 1986AB + 2MG N71208 001 Nov 17, 1986AB 5MG N71209 001 Nov 17, 1986AB 10MG N71210 001 Mar 11, 1988AB 20MG N71211 001 Mar 11, 1988AB WATSON LABS 0.5MG N70981 001 Mar 06, 1987AB 2MG N70983 001 Mar 06, 1987AB 5MG N70984 001 Mar 06, 1987HALOPERIDOL DECANOATE

INJECTABLE; INJECTION

HALDOLAO + ORTHO MCNEIL EQ 50MG BASE/ML N18701 001 Jan 14, 1986AO + EQ 100MG BASE/ML N18701 002 Jan 31, 1997HALOPERIDOL DECANOATEAO ABRAXIS PHARM EQ 50MG BASE/ML N74893 001 Dec 19, 1997AO EQ 100MG BASE/ML N74893 002 Dec 19, 1997AO APOTEX INC EQ 50MG BASE/ML N75440 001 Feb 28, 2000AO EQ 100MG BASE/ML N75440 002 Feb 28, 2000AO BEDFORD EQ 50MG BASE/ML N74811 001 Jan 30, 1998

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 190 (of 370)

HALOPERIDOL DECANOATE

INJECTABLE; INJECTION

HALOPERIDOL DECANOATE

<u>AO</u>	BEDFORD	<u>EQ 100MG BASE/ML</u>	<u>N75305</u>	<u>001</u>	Sep 28, 1998
<u>AO</u>	SANDOZ	<u>EQ 50MG BASE/ML</u>	<u>N76463</u>	<u>001</u>	Jun 24, 2005
<u>AO</u>		<u>EQ 100MG BASE/ML</u>	<u>N76463</u>	<u>002</u>	Jun 24, 2005
<u>AO</u>	SICOR PHARMS	<u>EQ 50MG BASE/ML</u>	<u>N75393</u>	<u>001</u>	May 11, 1999
<u>AO</u>		<u>EQ 100MG BASE/ML</u>	<u>N75393</u>	<u>002</u>	May 11, 1999

HALOPERIDOL LACTATE

CONCENTRATE; ORAL

HALOPERIDOL

<u>AA</u>	PHARM ASSOC	<u>EQ 2MG BASE/ML</u>	<u>N73037</u>	<u>001</u>	Feb 26, 1993
<u>AA</u>	SILARX	<u>EQ 2MG BASE/ML</u>	<u>N73364</u>	<u>001</u>	Sep 28, 1993
<u>AA</u>	+ TEVA PHARMS	<u>EQ 2MG BASE/ML</u>	<u>N71617</u>	<u>001</u>	Dec 01, 1988

INJECTABLE; INJECTION

HALDOL

<u>AP</u>	+ ORTHO MCNEIL	<u>EQ 5MG BASE/ML</u>	<u>N15923</u>	<u>001</u>	
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HALOPERIDOL

<u>AP</u>	ABRAXIS PHARM	<u>EQ 5MG BASE/ML</u>	<u>N75689</u>	<u>001</u>	Mar 09, 2001
<u>AP</u>	APOTEX INC	<u>EQ 5MG BASE/ML</u>	<u>N76791</u>	<u>001</u>	Aug 25, 2004
<u>AP</u>		<u>EQ 5MG BASE/ML</u>	<u>N76828</u>	<u>001</u>	Aug 25, 2004
<u>AP</u>	BEDFORD	<u>EQ 5MG BASE/ML</u>	<u>N75858</u>	<u>001</u>	Jun 18, 2001
<u>AP</u>	GLAND PHARMA LTD	<u>EQ 5MG BASE/ML</u>	<u>N76774</u>	<u>001</u>	Aug 25, 2004
<u>AP</u>	SANDOZ	<u>EQ 5MG BASE/ML</u>	<u>N76464</u>	<u>001</u>	Sep 29, 2004
<u>AP</u>	SICOR PHARMS	<u>EQ 5MG BASE/ML</u>	<u>N76035</u>	<u>001</u>	Aug 29, 2001
<u>AP</u>	WATSON LABS	<u>EQ 5MG BASE/ML</u>	<u>N70714</u>	<u>001</u>	May 17, 1988

SOLUTION; ORAL

HALOPERIDOL LACTATE

	ACTAVIS MID ATLANTIC	EQ 1MG BASE/ML	N74536	001	Nov 02, 1995
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HEPARIN SODIUM

INJECTABLE; INJECTION

HEPARIN LOCK FLUSH

<u>AP</u>	HOSPIRA	<u>10 UNITS/ML</u>	<u>N88346</u>	<u>001</u>	May 18, 1983
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HEPARIN LOCK FLUSH IN PLASTIC CONTAINER

<u>AP</u>	HOSPIRA	<u>10 UNITS/ML</u>	<u>N05264</u>	<u>015</u>	May 21, 1985
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HEPARIN SODIUM

<u>AP</u>	ABRAXIS PHARM	<u>1,000 UNITS/ML</u>	<u>N17029</u>	<u>001</u>	
<u>AP</u>		<u>1,000 UNITS/ML</u>	<u>N17979</u>	<u>001</u>	
<u>AP</u>		<u>5,000 UNITS/ML</u>	<u>N17651</u>	<u>006</u>	
<u>AP</u>		<u>10,000 UNITS/ML</u>	<u>N17029</u>	<u>003</u>	
<u>AP</u>		<u>10,000 UNITS/ML</u>	<u>N17979</u>	<u>002</u>	
<u>AP</u>	+	<u>20,000 UNITS/ML</u>	<u>N17029</u>	<u>004</u>	
<u>AP</u>	BAXTER HLTHCARE	<u>1,000 UNITS/ML</u>	<u>N17037</u>	<u>001</u>	
<u>AP</u>		<u>5,000 UNITS/0.5ML</u>	<u>N17037</u>	<u>013</u>	Apr 07, 1986
<u>AP</u>		<u>5,000 UNITS/ML</u>	<u>N17037</u>	<u>002</u>	
<u>AP</u>		<u>10,000 UNITS/ML</u>	<u>N17037</u>	<u>003</u>	
<u>AP</u>	HOSPIRA	<u>5,000 UNITS/ML</u>	<u>N88100</u>	<u>001</u>	Apr 28, 1983
<u>AP</u>	MARSAM PHARMS LLC	<u>1,000 UNITS/ML</u>	<u>N40008</u>	<u>001</u>	Oct 10, 1995
<u>AP</u>	+ PHARMACIA AND UPJOHN	<u>1,000 UNITS/ML</u>	<u>N04570</u>	<u>001</u>	
<u>AP</u>	+	<u>5,000 UNITS/ML</u>	<u>N04570</u>	<u>002</u>	
<u>AP</u>	+	<u>10,000 UNITS/ML</u>	<u>N04570</u>	<u>003</u>	

HEPARIN SODIUM 1,000 UNITS AND SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER

<u>AP</u>	BAXTER HLTHCARE	<u>200 UNITS/100ML</u>	<u>N18609</u>	<u>001</u>	Apr 28, 1982
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HEPARIN SODIUM 1,000 UNITS IN SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER

<u>AP</u>	B BRAUN	<u>200 UNITS/100ML</u>	<u>N19953</u>	<u>001</u>	Jul 20, 1992
<u>AP</u>	HOSPIRA	<u>200 UNITS/100ML</u>	<u>N18916</u>	<u>010</u>	Jun 23, 1989

HEPARIN SODIUM 10,000 UNITS IN DEXTROSE 5% IN PLASTIC CONTAINER

<u>AP</u>	HOSPIRA	<u>10,000 UNITS/100ML</u>	<u>N19339</u>	<u>003</u>	Mar 27, 1985
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27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 191 (of 370)

HEPARIN SODIUM

INJECTABLE; INJECTION

HEPARIN SODIUM 12,500 UNITS IN DEXTROSE 5% IN PLASTIC CONTAINER			
	HOSPIRA	5,000 UNITS/100ML	N19339 001 Mar 27, 1985
HEPARIN SODIUM 12,500 UNITS IN SODIUM CHLORIDE 0.45% IN PLASTIC CONTAINER			
	HOSPIRA	5,000 UNITS/100ML	N18916 006 Jan 31, 1984
HEPARIN SODIUM 2,000 UNITS AND SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER			
<u>AP</u>	BAXTER HLTHCARE	<u>200 UNITS/100ML</u>	<u>N18609</u> <u>002</u> Apr 28, 1982
HEPARIN SODIUM 2,000 UNITS IN SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER			
<u>AP</u>	HOSPIRA	<u>200 UNITS/100ML</u>	<u>N18916</u> <u>011</u> Jun 23, 1989
HEPARIN SODIUM 20,000 UNITS AND DEXTROSE 5% IN PLASTIC CONTAINER			
<u>AP</u>	BAXTER HLTHCARE	<u>4,000 UNITS/100ML</u>	<u>N18814</u> <u>001</u> Oct 31, 1983
HEPARIN SODIUM 20,000 UNITS IN DEXTROSE 5% IN PLASTIC CONTAINER			
<u>AP</u>	B BRAUN	<u>4,000 UNITS/100ML</u>	<u>N19952</u> <u>001</u> Jul 20, 1992
<u>AP</u>	HOSPIRA	<u>4,000 UNITS/100ML</u>	<u>N19805</u> <u>001</u> Jan 25, 1989
HEPARIN SODIUM 25,000 UNITS AND DEXTROSE 5% IN PLASTIC CONTAINER			
<u>AP</u>	BAXTER HLTHCARE	<u>5,000 UNITS/100ML</u>	<u>N18814</u> <u>003</u> Jul 09, 1985
<u>AP</u>		<u>10,000 UNITS/100ML</u>	<u>N18814</u> <u>004</u> Jul 02, 1987
HEPARIN SODIUM 25,000 UNITS IN DEXTROSE 5% IN PLASTIC CONTAINER			
<u>AP</u>	B BRAUN	<u>5,000 UNITS/100ML</u>	<u>N19952</u> <u>004</u> Jul 20, 1992
<u>AP</u>		<u>10,000 UNITS/100ML</u>	<u>N19952</u> <u>005</u> Jul 20, 1992
<u>AP</u>	HOSPIRA	<u>5,000 UNITS/100ML</u>	<u>N19339</u> <u>004</u> Mar 27, 1985
<u>AP</u>		<u>5,000 UNITS/100ML</u>	<u>N19805</u> <u>002</u> Jan 25, 1989
<u>AP</u>		<u>10,000 UNITS/100ML</u>	<u>N19339</u> <u>002</u> Mar 27, 1985
HEPARIN SODIUM 25,000 UNITS IN SODIUM CHLORIDE 0.45% IN PLASTIC CONTAINER			
	HOSPIRA	5,000 UNITS/100ML	N18916 007 Jan 31, 1984
		10,000 UNITS/100ML	N18916 008 Jan 31, 1984
HEPARIN SODIUM IN PLASTIC CONTAINER			
<u>AP</u>	ABRAXIS PHARM	<u>1,000 UNITS/ML</u>	<u>N17029</u> <u>013</u> Dec 05, 1985
<u>AP</u>		<u>5,000 UNITS/ML</u>	<u>N17029</u> <u>014</u> Dec 05, 1985
<u>AP</u>		<u>10,000 UNITS/ML</u>	<u>N17029</u> <u>015</u> Dec 05, 1985
<u>AP</u>		<u>20,000 UNITS/ML</u>	<u>N17029</u> <u>016</u> Dec 05, 1985
HEPARIN SODIUM PRESERVATIVE FREE			
<u>AP</u> +	ABRAXIS PHARM	<u>1,000 UNITS/ML</u>	<u>N17029</u> <u>010</u> Apr 28, 1986
<u>AP</u> +	HOSPIRA	<u>2,500 UNITS/ML</u>	<u>N05264</u> <u>014</u> Apr 07, 1986
<u>AP</u> +		<u>10,000 UNITS/ML</u>	<u>N89522</u> <u>001</u> May 04, 1987
	+	2,000 UNITS/ML	N05264 013 Apr 07, 1986

HEXACHLOROPHENE

EMULSION; TOPICAL

PHISOHEX

+ SANOFI AVENTIS US 3% N06882 001

SPONGE; TOPICAL

PRE-OPAT + DAVIS AND GECK 480MG N17433 001PRE-OP IIAT DAVIS AND GECK 480MG N17433 002HISTRELIN ACETATE

IMPLANT; SUBCUTANEOUS

VANTAS

+ VALERA 50MG N21732 001 Oct 12, 2004

HOMATROPINE METHYLBROMIDE; HYDROCODONE BITARTRATE

SYRUP; ORAL

HYCODANAA + ENDO PHARMS 1.5MG/5ML; 5MG/5ML N05213 002 Jul 26, 1988HYDROCODONE BITARTRATE AND HOMATROPINE METHYLBROMIDEAA IVAX PHARMS 1.5MG/5ML; 5MG/5ML N40285 001 Jul 19, 1999

PRESCRIPTION DRUG PRODUCT LIST

3 - 192 (of 370)

HOMATROPINE METHYLBROMIDE; HYDROCODONE BITARTRATE

SYRUP; ORAL

HYDROCODONE COMPOUND

<u>AA</u>	ACTAVIS MID ATLANTIC	<u>1.5MG/5ML;5MG/5ML</u>	<u>N88017</u>	<u>001</u>	Jul 05, 1983
<u>AA</u>	MORTON GROVE	<u>1.5MG/5ML;5MG/5ML</u>	<u>N88008</u>	<u>001</u>	Mar 03, 1983

TABLET; ORAL

HOMATROPINE METHYLBROMIDE AND HYDROCODONE BITARTRATE

<u>AA</u>	ACTAVIS TOTOWA	<u>1.5MG;5MG</u>	<u>N40295</u>	<u>001</u>	Dec 01, 2000
<u>AA</u>	<u>HYCODAN</u>				
<u>AA</u>	+ ENDO PHARMS	<u>1.5MG;5MG</u>	<u>N05213</u>	<u>001</u>	Jul 26, 1988
<u>AA</u>	<u>TUSSIGON</u>				
<u>AA</u>	KING PHARMS	<u>1.5MG;5MG</u>	<u>N88508</u>	<u>001</u>	Jul 30, 1985

HYALURONIDASE

INJECTABLE; INJECTION

AMPHADASE

	+ AMPHASTAR PHARM	150 UNITS/ML	N21665	001	Oct 26, 2004
	HYDASE				
	+ PRIMAPHARM	150 UNITS/ML	N21716	001	Oct 25, 2005
	VITRASE				
	ISTA PHARMS	200 UNITS/VIAL	N21640	002	Dec 02, 2004
	+	6,200 UNITS/VIAL	N21640	001	May 05, 2004

HYALURONIDASE RECOMBINANT HUMAN

INJECTABLE; INJECTION

HYLENEX RECOMBINANT

	+ HALOZYME THERAP	150 UNITS/ML	N21859	001	Dec 02, 2005
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HYDRALAZINE HYDROCHLORIDE

INJECTABLE; INJECTION

HYDRALAZINE HYDROCHLORIDE

<u>AP</u>	ABRAXIS PHARM	<u>20MG/ML</u>	<u>N40388</u>	<u>001</u>	Mar 13, 2001
<u>AP</u>	+ LUITPOLD	<u>20MG/ML</u>	<u>N40136</u>	<u>001</u>	Jun 30, 1997

TABLET; ORAL

HYDRALAZINE HYDROCHLORIDE

<u>AA</u>	MUTUAL PHARM	<u>10MG</u>	<u>N89359</u>	<u>001</u>	Jul 25, 1986
<u>AA</u>		<u>25MG</u>	<u>N89258</u>	<u>001</u>	May 05, 1986
<u>AA</u>		<u>50MG</u>	<u>N89259</u>	<u>001</u>	May 05, 1986
<u>AA</u>	PAR PHARM	<u>10MG</u>	<u>N87836</u>	<u>001</u>	Oct 05, 1982
<u>AA</u>		<u>25MG</u>	<u>N86961</u>	<u>002</u>	
<u>AA</u>		<u>50MG</u>	<u>N86962</u>	<u>001</u>	
<u>AA</u>		<u>100MG</u>	<u>N88391</u>	<u>001</u>	Sep 27, 1983
<u>AA</u>	+ PLIVA	<u>10MG</u>	<u>N89097</u>	<u>001</u>	Dec 18, 1985
<u>AA</u>	+	<u>25MG</u>	<u>N88467</u>	<u>001</u>	May 01, 1984
<u>AA</u>	+	<u>50MG</u>	<u>N88468</u>	<u>001</u>	May 01, 1984
<u>AA</u>	+	<u>100MG</u>	<u>N89098</u>	<u>001</u>	Dec 18, 1985
<u>AA</u>	SANDOZ	<u>10MG</u>	<u>N83241</u>	<u>001</u>	
<u>AA</u>		<u>25MG</u>	<u>N83560</u>	<u>001</u>	
<u>AA</u>		<u>50MG</u>	<u>N83561</u>	<u>001</u>	
<u>AA</u>	WATSON LABS	<u>25MG</u>	<u>N84504</u>	<u>001</u>	
<u>AA</u>		<u>50MG</u>	<u>N84503</u>	<u>001</u>	

HYDRALAZINE HYDROCHLORIDE; HYDROCHLOROTHIAZIDE

CAPSULE; ORAL

APRESAZIDE

<u>AB</u>	NOVARTIS	<u>25MG;25MG</u>	<u>N84735</u>	<u>001</u>	
<u>AB</u>		<u>50MG;50MG</u>	<u>N84810</u>	<u>001</u>	

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 193 (of 370)

HYDRALAZINE HYDROCHLORIDE; HYDROCHLOROTHIAZIDE

CAPSULE; ORAL

HYDRA-ZIDE

<u>AB</u>	PAR PHARM	<u>25MG;25MG</u>	<u>N88957</u>	<u>001</u>	Oct 21, 1985
<u>AB</u>		<u>50MG;50MG</u>	<u>N88946</u>	<u>001</u>	Oct 21, 1985
	+	100MG;50MG	N88961	001	Oct 21, 1985

HYDRALAZINE HYDROCHLORIDE; ISOSORBIDE DINITRATE

TABLET; ORAL

BIDIL

	+	NITROMED	37.5MG;20MG	N20727	001	Jun 23, 2005
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HYDROCHLOROTHIAZIDE

CAPSULE; ORAL

HYDROCHLOROTHIAZIDE

<u>AB</u>	IVAX PHARMS	<u>12.5MG</u>	<u>N77005</u>	<u>001</u>	Jul 13, 2005
<u>AB</u>	MYLAN	<u>12.5MG</u>	<u>N75640</u>	<u>001</u>	Jan 28, 2000
<u>AB</u>	VINTAGE PHARMS	<u>12.5MG</u>	<u>N75907</u>	<u>001</u>	Sep 17, 2002
	<u>MICROZIDE</u>				
<u>AB</u>	+ WATSON LABS	<u>12.5MG</u>	<u>N20504</u>	<u>001</u>	Dec 27, 1996

SOLUTION; ORAL

HYDROCHLOROTHIAZIDE

	+	ROXANE	50MG/5ML	N88587	001	Jul 02, 1984
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TABLET; ORAL

ESIDRIX

<u>AB</u>	NOVARTIS	<u>25MG</u>	<u>N11793</u>	<u>005</u>	
<u>AB</u>		<u>50MG</u>	<u>N11793</u>	<u>008</u>	
<u>AB</u>	<u>HYDROCHLOROTHIAZIDE</u>				
<u>AB</u>	ACTAVIS ELIZABETH	<u>25MG</u>	<u>N85054</u>	<u>002</u>	
<u>AB</u>		<u>50MG</u>	<u>N85208</u>	<u>001</u>	
<u>AB</u>	CLONMEL HLTHCARE	<u>25MG</u>	<u>N87059</u>	<u>001</u>	
<u>AB</u>		<u>50MG</u>	<u>N87068</u>	<u>001</u>	
<u>AB</u>	IVAX PHARMS	<u>25MG</u>	<u>N83177</u>	<u>001</u>	
<u>AB</u>	+	<u>50MG</u>	<u>N83177</u>	<u>002</u>	
<u>AB</u>	LEINER	<u>25MG</u>	<u>N85181</u>	<u>001</u>	
<u>AB</u>		<u>50MG</u>	<u>N85182</u>	<u>001</u>	
<u>AB</u>	SANDOZ	<u>25MG</u>	<u>N87565</u>	<u>001</u>	Mar 09, 1982
<u>AB</u>		<u>50MG</u>	<u>N84912</u>	<u>001</u>	
<u>AB</u>		<u>50MG</u>	<u>N85219</u>	<u>001</u>	
<u>AB</u>	VINTAGE PHARMS	<u>25MG</u>	<u>N40412</u>	<u>001</u>	Mar 29, 2002
<u>AB</u>		<u>50MG</u>	<u>N40412</u>	<u>002</u>	Mar 29, 2002
<u>AB</u>	WATSON LABS	<u>25MG</u>	<u>N81189</u>	<u>001</u>	Jan 24, 1992
<u>AB</u>	WEST WARD	<u>25MG</u>	<u>N84878</u>	<u>002</u>	Jul 12, 2006
<u>AB</u>	<u>ORETIC</u>				
<u>AB</u>	ABBOTT	<u>50MG</u>	<u>N11971</u>	<u>002</u>	

HYDROCHLOROTHIAZIDE; IRBESARTAN

TABLET; ORAL

AVALIDE

	SANOFI AVENTIS US	12.5MG;150MG	N20758	002	Sep 30, 1997
		12.5MG;300MG	N20758	003	Aug 31, 1998
	+	25MG;300MG	N20758	004	Mar 15, 2005

HYDROCHLOROTHIAZIDE; LISINAPRIL

TABLET; ORAL

LISINAPRIL AND HYDROCHLOROTHIAZIDE

<u>AB</u>	ACTAVIS ELIZABETH	<u>12.5MG;10MG</u>	<u>N76230</u>	<u>001</u>	Jul 01, 2002
<u>AB</u>		<u>12.5MG;20MG</u>	<u>N76230</u>	<u>002</u>	Jul 01, 2002
<u>AB</u>		<u>25MG;20MG</u>	<u>N76230</u>	<u>003</u>	Jul 01, 2002

PRESCRIPTION DRUG PRODUCT LIST

3 - 194 (of 370)

HYDROCHLOROTHIAZIDE; LISINOPRIL

TABLET; ORAL

LISINOPRIL AND HYDROCHLOROTHIAZIDE

<u>AB</u>	APOTEX INC	<u>12.5MG;10MG</u>	<u>N76674</u>	<u>001</u>	Oct 05, 2004
<u>AB</u>		<u>12.5MG;20MG</u>	<u>N76674</u>	<u>002</u>	Oct 05, 2004
<u>AB</u>		<u>25MG;20MG</u>	<u>N76674</u>	<u>003</u>	Oct 05, 2004
<u>AB</u>	AUROBINDO	<u>12.5MG;10MG</u>	<u>N77606</u>	<u>001</u>	Mar 14, 2006
<u>AB</u>		<u>12.5MG;20MG</u>	<u>N77606</u>	<u>002</u>	Mar 14, 2006
<u>AB</u>		<u>25MG;20MG</u>	<u>N77606</u>	<u>003</u>	Mar 14, 2006
<u>AB</u>	IVAX PHARMS	<u>12.5MG;10MG</u>	<u>N75776</u>	<u>001</u>	Jul 01, 2002
<u>AB</u>		<u>12.5MG;20MG</u>	<u>N75776</u>	<u>002</u>	Jul 01, 2002
<u>AB</u>		<u>25MG;20MG</u>	<u>N75776</u>	<u>003</u>	Jul 01, 2002
<u>AB</u>	LUPIN	<u>12.5MG;10MG</u>	<u>N77912</u>	<u>001</u>	Sep 27, 2006
<u>AB</u>		<u>12.5MG;20MG</u>	<u>N77912</u>	<u>002</u>	Sep 27, 2006
<u>AB</u>		<u>25MG;20MG</u>	<u>N77912</u>	<u>003</u>	Sep 27, 2006
<u>AB</u>	MYLAN	<u>12.5MG;10MG</u>	<u>N76113</u>	<u>001</u>	Jul 01, 2002
<u>AB</u>		<u>12.5MG;20MG</u>	<u>N76113</u>	<u>002</u>	Jul 01, 2002
<u>AB</u>		<u>25MG;20MG</u>	<u>N76113</u>	<u>003</u>	Jul 01, 2002
<u>AB</u>	RANBAXY	<u>12.5MG;10MG</u>	<u>N76007</u>	<u>001</u>	Jul 01, 2002
<u>AB</u>		<u>12.5MG;20MG</u>	<u>N76007</u>	<u>002</u>	Jul 01, 2002
<u>AB</u>		<u>25MG;20MG</u>	<u>N76007</u>	<u>003</u>	Jul 01, 2002
<u>AB</u>	SANDOZ	<u>12.5MG;10MG</u>	<u>N75926</u>	<u>001</u>	Jul 01, 2002
<u>AB</u>		<u>12.5MG;10MG</u>	<u>N76262</u>	<u>001</u>	Jul 01, 2002
<u>AB</u>		<u>12.5MG;20MG</u>	<u>N75926</u>	<u>002</u>	Jul 01, 2002
<u>AB</u>		<u>12.5MG;20MG</u>	<u>N76262</u>	<u>002</u>	Jul 01, 2002
<u>AB</u>		<u>25MG;20MG</u>	<u>N75926</u>	<u>003</u>	Jul 01, 2002
<u>AB</u>		<u>25MG;20MG</u>	<u>N76262</u>	<u>003</u>	Jul 01, 2002
<u>AB</u>	TEVA	<u>12.5MG;10MG</u>	<u>N75869</u>	<u>001</u>	Jul 01, 2002
<u>AB</u>		<u>12.5MG;20MG</u>	<u>N75869</u>	<u>002</u>	Jul 01, 2002
<u>AB</u>		<u>25MG;20MG</u>	<u>N75869</u>	<u>003</u>	Jul 01, 2002
<u>AB</u>	WATSON LABS	<u>12.5MG;10MG</u>	<u>N76194</u>	<u>003</u>	Jul 01, 2002
<u>AB</u>		<u>12.5MG;20MG</u>	<u>N76194</u>	<u>001</u>	Jul 01, 2002
<u>AB</u>		<u>25MG;20MG</u>	<u>N76194</u>	<u>002</u>	Jul 01, 2002
<u>AB</u>	WEST WARD	<u>12.5MG;10MG</u>	<u>N76265</u>	<u>001</u>	Jul 08, 2002
<u>AB</u>		<u>12.5MG;20MG</u>	<u>N76265</u>	<u>002</u>	Jul 08, 2002
<u>AB</u>		<u>25MG;20MG</u>	<u>N76265</u>	<u>003</u>	Jul 08, 2002
<u>PRINZIDE</u>					
<u>AB</u>	MERCK	<u>12.5MG;10MG</u>	<u>N19778</u>	<u>003</u>	Nov 18, 1993
<u>AB</u>		<u>12.5MG;20MG</u>	<u>N19778</u>	<u>001</u>	Feb 16, 1989
<u>AB</u>	+	<u>25MG;20MG</u>	<u>N19778</u>	<u>002</u>	Feb 16, 1989
<u>ZESTORETIC</u>					
<u>AB</u>	ASTRAZENECA	<u>12.5MG;10MG</u>	<u>N19888</u>	<u>003</u>	Nov 18, 1993
<u>AB</u>		<u>12.5MG;20MG</u>	<u>N19888</u>	<u>001</u>	Sep 20, 1990
<u>AB</u>		<u>25MG;20MG</u>	<u>N19888</u>	<u>002</u>	Jul 20, 1989

HYDROCHLOROTHIAZIDE; LOSARTAN POTASSIUM

TABLET; ORAL

HYZAAR

MERCK

12.5MG;50MG

N20387 001 Apr 28, 1995

12.5MG;100MG

N20387 003 Oct 20, 2005

+

25MG;100MG

N20387 002 Nov 10, 1998

HYDROCHLOROTHIAZIDE; METHYLDOPA

TABLET; ORAL

METHYLDOPA AND HYDROCHLOROTHIAZIDE

MYLAN

15MG;250MG

N70264 001 Jan 23, 1986

+

25MG;250MG

N70265 001 Jan 23, 1986

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 195 (of 370)

HYDROCHLOROTHIAZIDE; METOPROLOL SUCCINATE

TABLET, EXTENDED RELEASE; ORAL

DUTOPROL

ASTRAZENECA	12.5MG;EQ 25MG TARTRATE	N21956	001	Aug 28, 2006
	12.5MG;EQ 50MG TARTRATE	N21956	002	Aug 28, 2006
+	12.5MG;EQ 100MG TARTRATE	N21956	003	Aug 28, 2006

HYDROCHLOROTHIAZIDE; METOPROLOL TARTRATE

TABLET; ORAL

LOPRESSOR HCT

<u>AB</u>	NOVARTIS	<u>25MG;50MG</u>	<u>N18303</u>	<u>001</u>	Dec 31, 1984
<u>AB</u>		<u>25MG;100MG</u>	<u>N18303</u>	<u>002</u>	Dec 31, 1984
<u>AB</u>	+	<u>50MG;100MG</u>	<u>N18303</u>	<u>003</u>	Dec 31, 1984
	<u>METOPROLOL TARTRATE AND HYDROCHLOROTHIAZIDE</u>				
<u>AB</u>	MYLAN	<u>25MG;50MG</u>	<u>N76792</u>	<u>001</u>	Aug 20, 2004
<u>AB</u>		<u>25MG;100MG</u>	<u>N76792</u>	<u>002</u>	Aug 20, 2004
<u>AB</u>		<u>50MG;100MG</u>	<u>N76792</u>	<u>003</u>	Aug 20, 2004

HYDROCHLOROTHIAZIDE; MOEXIPRIL HYDROCHLORIDE

TABLET; ORAL

UNIRETIC

SCHWARZ PHARMA	12.5MG;7.5MG	N20729	001	Jun 27, 1997
	12.5MG;15MG	N20729	003	Feb 14, 2002
+	25MG;15MG	N20729	002	Jun 27, 1997

HYDROCHLOROTHIAZIDE; OLMESARTAN MEDOXOMIL

TABLET; ORAL

BENICAR HCT

DAIICHI SANKYO	12.5MG;20MG	N21532	002	Jun 05, 2003
	12.5MG;40MG	N21532	003	Jun 05, 2003
+	25MG;40MG	N21532	005	Jun 05, 2003

HYDROCHLOROTHIAZIDE; PROPRANOLOL HYDROCHLORIDE

TABLET; ORAL

INDERIDE-40/25

<u>AB</u>	+	WYETH PHARMS INC	<u>25MG;40MG</u>	<u>N18031</u>	<u>001</u>	
		<u>PROPRANOLOL HYDROCHLORIDE AND HYDROCHLOROTHIAZIDE</u>				
<u>AB</u>		ACTAVIS ELIZABETH	<u>25MG;40MG</u>	<u>N70851</u>	<u>001</u>	May 15, 1986
<u>AB</u>			<u>25MG;80MG</u>	<u>N70852</u>	<u>001</u>	May 15, 1986
<u>AB</u>		BARR	<u>25MG;40MG</u>	<u>N70704</u>	<u>001</u>	Oct 01, 1986
<u>AB</u>			<u>25MG;80MG</u>	<u>N70705</u>	<u>001</u>	Oct 01, 1986
<u>AB</u>		MYLAN	<u>25MG;40MG</u>	<u>N70946</u>	<u>001</u>	Mar 04, 1987
<u>AB</u>			<u>25MG;80MG</u>	<u>N70947</u>	<u>001</u>	Apr 01, 1987
<u>AB</u>		PLIVA	<u>25MG;40MG</u>	<u>N72042</u>	<u>001</u>	Mar 14, 1988
<u>AB</u>			<u>25MG;80MG</u>	<u>N72043</u>	<u>001</u>	Mar 14, 1988
<u>AB</u>		SANDOZ	<u>25MG;40MG</u>	<u>N71060</u>	<u>001</u>	Aug 26, 1987
<u>AB</u>			<u>25MG;80MG</u>	<u>N71061</u>	<u>001</u>	Aug 26, 1987
<u>AB</u>		WATSON LABS	<u>25MG;40MG</u>	<u>N70301</u>	<u>001</u>	Apr 18, 1986
<u>AB</u>			<u>25MG;80MG</u>	<u>N70305</u>	<u>001</u>	Apr 18, 1986

HYDROCHLOROTHIAZIDE; QUINAPRIL HYDROCHLORIDE

TABLET; ORAL

ACCURETIC

<u>AB</u>		PFIZER PHARMS	<u>12.5MG;EQ 10MG BASE</u>	<u>N20125</u>	<u>001</u>	Dec 28, 1999
<u>AB</u>			<u>12.5MG;EQ 20MG BASE</u>	<u>N20125</u>	<u>002</u>	Dec 28, 1999
<u>AB</u>	+		<u>25MG;EQ 20MG BASE</u>	<u>N20125</u>	<u>003</u>	Dec 28, 1999
		<u>QUINAPRIL HYDROCHLORIDE AND HYDROCHLOROTHIAZIDE</u>				
<u>AB</u>		MYLAN	<u>12.5MG;EQ 10MG BASE</u>	<u>N77093</u>	<u>001</u>	Mar 28, 2005
<u>AB</u>			<u>12.5MG;EQ 20MG BASE</u>	<u>N77093</u>	<u>002</u>	Mar 28, 2005

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 196 (of 370)

HYDROCHLOROTHIAZIDE; QUINAPRIL HYDROCHLORIDE

TABLET; ORAL

<u>QUINAPRIL HYDROCHLORIDE AND HYDROCHLOROTHIAZIDE</u>					
<u>AB</u>	MYLAN	<u>25MG;EQ 20MG BASE</u>	<u>N77093</u>	<u>003</u>	Mar 28, 2005
<u>QUINARETIC</u>					
<u>AB</u>	ACTAVIS TOTOWA	<u>12.5MG;EQ 10MG BASE</u>	<u>N76374</u>	<u>001</u>	Mar 31, 2004
<u>AB</u>		<u>12.5MG;EQ 20MG BASE</u>	<u>N76374</u>	<u>002</u>	Mar 31, 2004
<u>AB</u>		<u>25MG;EQ 20MG BASE</u>	<u>N76374</u>	<u>003</u>	Mar 31, 2004

HYDROCHLOROTHIAZIDE; SPIRONOLACTONE

TABLET; ORAL

<u>ALDACTAZIDE</u>					
<u>AB</u>	GD SEARLE LLC	<u>25MG;25MG</u>	<u>N12616</u>	<u>004</u>	Dec 30, 1982
	+	50MG;50MG	N12616	005	Dec 30, 1982
<u>SPIRONOLACTONE AND HYDROCHLOROTHIAZIDE</u>					
<u>AB</u>	MUTUAL PHARM	<u>25MG;25MG</u>	<u>N87267</u>	<u>001</u>	
<u>AB</u>		<u>25MG;25MG</u>	<u>N89534</u>	<u>001</u>	Jul 02, 1987
<u>AB</u>	MYLAN	<u>25MG;25MG</u>	<u>N86513</u>	<u>001</u>	
<u>AB</u>	SANDOZ	<u>25MG;25MG</u>	<u>N86881</u>	<u>001</u>	
<u>AB</u>	WATSON LABS	<u>25MG;25MG</u>	<u>N87398</u>	<u>001</u>	

HYDROCHLOROTHIAZIDE; TELMISARTAN

TABLET; ORAL

MICARDIS HCT					
	BOEHRINGER INGELHEIM	12.5MG;40MG	N21162	001	Nov 17, 2000
		12.5MG;80MG	N21162	002	Nov 17, 2000
	+	25MG;80MG	N21162	003	Apr 19, 2004

HYDROCHLOROTHIAZIDE; TRIAMTERENE

CAPSULE; ORAL

<u>DYAZIDE</u>					
<u>AB</u>	+ GLAXOSMITHKLINE	<u>25MG;37.5MG</u>	<u>N16042</u>	<u>003</u>	Mar 03, 1994
<u>TRIAMTERENE AND HYDROCHLOROTHIAZIDE</u>					
<u>AB</u>	BARR	<u>25MG;37.5MG</u>	<u>N74970</u>	<u>001</u>	Jan 06, 1998
<u>AB</u>	DURAMED PHARMS BARR	<u>25MG;37.5MG</u>	<u>N75052</u>	<u>001</u>	Jun 18, 1999
<u>AB</u>	IVAX PHARMS	<u>25MG;50MG</u>	<u>N74259</u>	<u>001</u>	Mar 30, 1995
<u>AB</u>	MYLAN	<u>25MG;37.5MG</u>	<u>N74701</u>	<u>001</u>	Jun 07, 1996
<u>AB</u>	SANDOZ	<u>25MG;37.5MG</u>	<u>N74821</u>	<u>001</u>	Jun 05, 1997
<u>AB</u>	+	<u>25MG;50MG</u>	<u>N73191</u>	<u>001</u>	Jul 31, 1991

TABLET; ORAL

<u>MAXZIDE</u>					
<u>AB</u>	+ MYLAN BERTEK	<u>50MG;75MG</u>	<u>N19129</u>	<u>001</u>	Oct 22, 1984
<u>MAXZIDE-25</u>					
<u>AB</u>	MYLAN BERTEK	<u>25MG;37.5MG</u>	<u>N19129</u>	<u>003</u>	May 13, 1988
<u>TRIAMTERENE AND HYDROCHLOROTHIAZIDE</u>					
<u>AB</u>	APOTEX INC	<u>25MG;37.5MG</u>	<u>N71251</u>	<u>002</u>	May 05, 1998
<u>AB</u>		<u>50MG;75MG</u>	<u>N71251</u>	<u>001</u>	Apr 17, 1988
<u>AB</u>	PLIVA	<u>25MG;37.5MG</u>	<u>N74026</u>	<u>001</u>	Apr 26, 1996
<u>AB</u>		<u>50MG;75MG</u>	<u>N73467</u>	<u>001</u>	Jan 31, 1996
<u>AB</u>	SANDOZ	<u>25MG;37.5MG</u>	<u>N73281</u>	<u>001</u>	Apr 30, 1992
<u>AB</u>		<u>50MG;75MG</u>	<u>N72011</u>	<u>001</u>	Jun 17, 1988
<u>AB</u>	WATSON LABS	<u>25MG;37.5MG</u>	<u>N73449</u>	<u>001</u>	Sep 23, 1993
<u>AB</u>		<u>50MG;75MG</u>	<u>N71851</u>	<u>001</u>	Nov 30, 1988
<u>AB</u>		<u>50MG;75MG</u>	<u>N71969</u>	<u>001</u>	Apr 17, 1988

HYDROCHLOROTHIAZIDE; VALSARTAN

TABLET; ORAL

DIOVAN HCT					
	NOVARTIS	12.5MG;80MG	N20818	001	Mar 06, 1998
		12.5MG;160MG	N20818	002	Mar 06, 1998

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 197 (of 370)

HYDROCHLOROTHIAZIDE; VALSARTAN

TABLET; ORAL

DIOVAN HCT

NOVARTIS

12.5MG;320MG

N20818 004 Apr 28, 2006

25MG;160MG

N20818 003 Jan 17, 2002

+

25MG;320MG

N20818 005 Apr 28, 2006

HYDROCODONE BITARTRATE; IBUPROFEN

TABLET; ORAL

HYDROCODONE BITARTRATE AND IBUPROFENAB ANDRX PHARMS7.5MG;200MGN76604 001 Dec 31, 2003AB + INTERPHARM5MG;200MGN76642 002 Mar 18, 2004AB7.5MG;200MGN76642 001 Oct 12, 2004AB TEVA7.5MG;200MGN76023 001 Apr 11, 2003AB VINTAGE PHARMS5MG;200MGN77727 001 Nov 06, 2006AB7.5MG;200MGN77723 001 Nov 06, 2006

10MG;200MG

N77723 002 Nov 06, 2006

VICOPROFENAB + ABBOTT7.5MG;200MGN20716 001 Sep 23, 1997HYDROCORTISONE

CREAM; TOPICAL

ALA-CORTAT DEL RAY LABS1%N80706 006ANUSOL HCAT SALIX PHARMS2.5%N88250 001 Jun 06, 1984HI-CORAT C AND M PHARMA2.5%N80483 001HYDROCORTISONEAT ACTAVIS MID ATLANTIC1%N87795 001 May 03, 1983AT2.5%N89682 001 Mar 10, 1988AT EVERYLIFE1%N80452 002AT FOUGERA1%N80693 003AT2.5%N89414 001 Dec 16, 1986AT INGRAM PHARM0.5%N80456 002AT1%N80456 003AT PERRIGO NEW YORK2.5%N85025 001AT TARO1%N86155 001AT2.5%N88799 001 Nov 09, 1984AT VINTAGE PHARMS2.5%N40503 001 Mar 12, 2004HYTONEAT + SANOFI AVENTIS US1%N80472 003AT +2.5%N80472 004SYNACORTAT MEDICIS1%N87458 001AT2.5%N87457 001

ENEMA; RECTAL

COLOCORTAB PADDOCK100MG/60MLN75172 001 Dec 03, 1999CORTENEMAAB + SOLVAY100MG/60MLN16199 001HYDROCORTISONEAB TEVA PHARMS100MG/60MLN74171 001 May 27, 1994

LOTION; TOPICAL

ALA-CORTAT DEL RAY LABS1%N83201 001

ALA-SCALP

DEL RAY LABS

2%

N83231 001

CETACORTAT HEALTHPOINT0.5%N80426 002

PRESCRIPTION DRUG PRODUCT LIST

3 - 198 (of 370)

HYDROCORTISONE

LOTION; TOPICAL				
<u>CETACORT</u>				
<u>AT</u>	HEALTHPOINT	<u>1%</u>	<u>N80426</u>	<u>001</u>
<u>HYDROCORTISONE</u>				
<u>AT</u>	ALTANA	<u>2.5%</u>	<u>N40351</u>	<u>001</u> Jul 25, 2000
<u>AT</u>	TARO	<u>2.5%</u>	<u>N40247</u>	<u>001</u> Jul 23, 1999
<u>AT</u>	VINTAGE PHARMS	<u>2.5%</u>	<u>N40417</u>	<u>001</u> Jul 30, 2003
<u>HYTONE</u>				
<u>AT</u>	+ SANOFI AVENTIS US	<u>1%</u>	<u>N80473</u>	<u>003</u>
<u>AT</u>	+	<u>2.5%</u>	<u>N80473</u>	<u>004</u> Nov 30, 1982
<u>NUTRACORT</u>				
<u>AT</u>	HEALTHPOINT	<u>1%</u>	<u>N80443</u>	<u>003</u>
<u>AT</u>		<u>2.5%</u>	<u>N87644</u>	<u>001</u> Aug 24, 1982
<u>STIE-CORT</u>				
<u>AT</u>	STIEFEL	<u>1%</u>	<u>N89066</u>	<u>001</u> Nov 25, 1985
<u>AT</u>		<u>2.5%</u>	<u>N89074</u>	<u>001</u> Nov 26, 1985
OINTMENT; TOPICAL				
<u>HYDROCORTISONE</u>				
<u>AT</u>	ACTAVIS MID ATLANTIC	<u>1%</u>	<u>N87796</u>	<u>001</u> Oct 13, 1982
<u>AT</u>	ALTANA	<u>1%</u>	<u>N80692</u>	<u>001</u>
<u>AT</u>	FOUGERA	<u>2.5%</u>	<u>N81203</u>	<u>001</u> May 28, 1993
<u>AT</u>	PERRIGO NEW YORK	<u>2.5%</u>	<u>N85027</u>	<u>001</u>
<u>AT</u>	TARO	<u>1%</u>	<u>N86257</u>	<u>001</u>
<u>AT</u>		<u>2.5%</u>	<u>N40310</u>	<u>001</u> Dec 29, 2000
<u>HYDROCORTISONE IN ABSORBASE</u>				
<u>AT</u>	CAROLINA MEDCL	<u>1%</u>	<u>N88138</u>	<u>001</u> Sep 06, 1985
POWDER; FOR RX COMPOUNDING				
HYDRO-RX				
	+ X GEN PHARMS	100%	N85982	001
SOLUTION; TOPICAL				
<u>TEXACORT</u>				
<u>AT</u>	+ SIRIUS LABS	<u>1%</u>	<u>N80425</u>	<u>001</u>
	+	2.5%	N81271	001 Apr 17, 1992
TABLET; ORAL				
CORTEF				
BP	+ PHARMACIA AND UPJOHN	20MG	N08697	002
		5MG	N08697	003
HYDROCORTISONE				
BP	WEST WARD	20MG	N83365	001
<u>HYDROCORTISONE ACETATE</u>				
AEROSOL, METERED; RECTAL				
CORTIFOAM				
	+ SCHWARZ PHARMA	10%	N17351	001 Feb 10, 1982
CREAM; TOPICAL				
HYDROCORTISONE ACETATE				
	+ FERNDAL LABS	2.5%	N40259	001 Jul 29, 1999
MICORT-HC				
	+ FERNDAL LABS	2%	N40398	001 Mar 29, 2002
		2.5%	N40396	001 Feb 27, 2001
LOTION; TOPICAL				
DRICORT				
	+ INGRAM PHARM	0.5%	N86207	001
PASTE; TOPICAL				
ORABASE HCA				
	COLGATE	0.5%	N83205	001

PRESCRIPTION DRUG PRODUCT LIST

3 - 199 (of 370)

HYDROCORTISONE ACETATE

POWDER; FOR RX COMPOUNDING

HYDROCORTISONE ACETATE

X GEN PHARMS 100%

N85981 001

HYDROCORTISONE ACETATE; NEOMYCIN SULFATE; POLYMYXIN B SULFATE

CREAM; TOPICAL

CORTISPORIN

+ MONARCH PHARMS 0.5%;EQ 3.5MG BASE/GM;10,000 UNITS/GM

N50218 001 Aug 09, 1985

HYDROCORTISONE ACETATE; OXYTETRACYCLINE HYDROCHLORIDE

SUSPENSION/DROPS; OPHTHALMIC

+ PFIZER 1.5%;EQ 5MG BASE/ML

N61016 001

HYDROCORTISONE ACETATE; PRAMOXINE HYDROCHLORIDE

AEROSOL, METERED; TOPICAL

EPIFOAM

BX SCHWARZ PHARMA 1%;1%

N86457 001

HYDROCORTISONE ACETATE 1% AND PRAMOXINE HYDROCHLORIDE 1%

BX BOCA PHARMA 1%;1%

N89440 001 May 17, 1988

PROCTOFOAM HC

BX SCHWARZ PHARMA 1%;1%

N86195 001

CREAM; TOPICAL

PRAMOSONE

FERNDAL LABS 0.5%;1%

N83778 001

1%;1%

N85368 001

LOTION; TOPICAL

PRAMOSONE

FERNDAL LABS 1%;1%

N85980 001

2.5%;1%

N85979 001

HYDROCORTISONE ACETATE; UREA

CREAM; TOPICAL

CARMOL HC

AT + KENWOOD LABS 1%;10%

N80505 001

U-CORT

AT TARO 1%;10%

N89472 001 Jun 13, 1988

HYDROCORTISONE BUTYRATE

CREAM; TOPICAL

HYDROCORTISONE BUTYRATE

AB TARO PHARM INDS 0.1%

N76654 001 Aug 03, 2005

LOCOID

AB + FERNDAL LABS 0.1%

N18514 001 Mar 31, 1982

LOCOID LIPOCREAM

+ FERNDAL LABS 0.1%

N20769 001 Sep 08, 1997

OINTMENT; TOPICAL

HYDROCORTISONE BUTYRATE

AB TARO 0.1%

N76842 001 Dec 27, 2004

LOCOID

AB + FERNDAL LABS 0.1%

N18652 001 Oct 29, 1982

SOLUTION; TOPICAL

HYDROCORTISONE BUTYRATE

AT TARO PHARM INDS 0.1%

N76364 001 Jan 14, 2004

LOCOID

AT + FERNDAL LABS 0.1%

N19116 001 Feb 25, 1987

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 200 (of 370)

HYDROCORTISONE PROBUTATE

CREAM; TOPICAL

PANDEL

+ SAVAGE LABS

0.1%

N20453 001

Feb 28, 1997

HYDROCORTISONE SODIUM SUCCINATE

INJECTABLE; INJECTION

A-HYDROCORTAP HOSPIRAEQ 100MG BASE/VIALN40666 001

Apr 06, 2006

APEQ 100MG BASE/VIALN85929 001APEQ 250MG BASE/VIALN85930 001APEQ 500MG BASE/VIALN85931 001APEQ 1GM BASE/VIALN85932 001SOLU-CORTEFAP + PHARMACIA AND UPJOHNEQ 100MG BASE/VIALN09866 001AP +EQ 250MG BASE/VIALN09866 002AP +EQ 500MG BASE/VIALN09866 003AP +EQ 1GM BASE/VIALN09866 004HYDROCORTISONE VALERATE

CREAM; TOPICAL

HYDROCORTISONE VALERATEAB PERRIGO NEW YORK0.2%N75666 001

May 24, 2000

AB TARO0.2%N75042 001

Aug 25, 1998

AB TEVA PHARMS0.2%N74489 001

Aug 12, 1998

WESTCORTAB + WESTWOOD SQUIBB0.2%N17950 001

OINTMENT; TOPICAL

HYDROCORTISONE VALERATEAB ALTANA0.2%N75085 001

Jul 31, 2001

AB TARO0.2%N75043 001

Aug 25, 1998

WESTCORTAB + WESTWOOD SQUIBB0.2%N18726 001

Aug 08, 1983

HYDROCORTISONE; NEOMYCIN SULFATE; POLYMYXIN B SULFATE

SOLUTION/DROPS; OTIC

CORTISPORINAT + MONARCH PHARMS1%;EQ 3.5MG BASE/ML;10,000 UNITS/MLN50479 001NEOMYCIN AND POLYMYXIN B SULFATES AND HYDROCORTISONEAT ALCON1%;EQ 3.5MG BASE/ML;10,000 UNITS/MLN62423 001

Aug 25, 1983

AT BAUSCH AND LOMB1%;EQ 3.5MG BASE/ML;10,000 UNITS/MLN64053 001

Dec 29, 1995

AT PHARMAFORCE1%;EQ 3.5MG BASE/ML;10,000 UNITS/MLN65216 001

Oct 31, 2005

SUSPENSION/DROPS; OPHTHALMIC

CORTISPORIN

+ MONARCH PHARMS

1%;EQ 3.5MG BASE/ML;10,000 UNITS/ML

N50169 001

NEOMYCIN AND POLYMYXIN B SULFATES AND HYDROCORTISONEAT ALCON UNIVERSAL1%;EQ 3.5MG BASE/ML;10,000 UNITS/MLN62874 001

May 11, 1988

SUSPENSION/DROPS; OTIC

CORTISPORINAT + MONARCH PHARMS1%;EQ 3.5MG BASE/ML;10,000 UNITS/MLN60613 001NEOMYCIN AND POLYMYXIN B SULFATES AND HYDROCORTISONEAT ALCON UNIVERSAL1%;EQ 3.5MG BASE/ML;10,000 UNITS/MLN62488 001

Nov 06, 1985

OTICAIRAT BAUSCH AND LOMB1%;EQ 3.5MG BASE/ML;10,000 UNITS/MLN64065 001

Aug 28, 1996

PEDIOTICAT MONARCH PHARMS1%;EQ 3.5MG BASE/ML;10,000 UNITS/MLN62822 001

Sep 29, 1987

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 201 (of 370)

HYDROCORTISONE; NEOMYCIN; POLYMYXIN B SULFATE

SUSPENSION/DROPS; OTIC

NEOMYCIN AND POLYMYXIN B SULFATES AND HYDROCORTISONE

<u>AT</u>	PHARMAFORCE	<u>1%;EQ 3.5MG BASE;10,000 UNITS</u>	<u>N65219</u>	<u>001</u>	May 01, 2006
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HYDROFLUMETHIAZIDE

TABLET; ORAL

HYDROFLUMETHIAZIDE

<u>AB</u>	PAR PHARM	<u>50MG</u>	<u>N88850</u>	<u>001</u>	May 31, 1985
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SALURON

<u>AB</u>	+ SHIRE	<u>50MG</u>	<u>N11949</u>	<u>001</u>	
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HYDROMORPHONE HYDROCHLORIDE

INJECTABLE; INJECTION

DILAUDID-HP

<u>AP</u>	+ ABBOTT	<u>10MG/ML</u>	<u>N19034</u>	<u>001</u>	Jan 11, 1984
	+	250MG/VIAL	N19034	002	Aug 04, 1994

HYDROMORPHONE HYDROCHLORIDE

<u>AP</u>	HOSPIRA	<u>10MG/ML</u>	<u>N74598</u>	<u>001</u>	Jun 19, 1997
<u>AP</u>	MAYNE PHARMA USA	<u>10MG/ML</u>	<u>N76444</u>	<u>001</u>	Apr 25, 2003
<u>AP</u>	WATSON LABS	<u>10MG/ML</u>	<u>N74317</u>	<u>001</u>	Aug 23, 1995

SOLUTION; ORAL

DILAUDID

<u>AA</u>	+ ABBOTT	<u>5MG/5ML</u>	<u>N19891</u>	<u>001</u>	Dec 07, 1992
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HYDROMORPHONE HYDROCHLORIDE

<u>AA</u>	ROXANE	<u>5MG/5ML</u>	<u>N74653</u>	<u>001</u>	Jul 29, 1998
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TABLET; ORAL

DILAUDID

<u>AB</u>	+ ABBOTT	<u>8MG</u>	<u>N19892</u>	<u>001</u>	Dec 07, 1992
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HYDROMORPHONE HYDROCHLORIDE

<u>AB</u>	ACTAVIS TOTOWA	<u>8MG</u>	<u>N76723</u>	<u>001</u>	Oct 18, 2005
<u>AB</u>	KV PHARM	<u>8MG</u>	<u>N77311</u>	<u>003</u>	Nov 09, 2005
		2MG	N77311	001	Nov 09, 2005
		4MG	N77311	002	Nov 09, 2005
<u>AB</u>	MALLINCKRODT	<u>8MG</u>	<u>N76855</u>	<u>001</u>	Dec 23, 2004
<u>AB</u>	ROXANE	<u>8MG</u>	<u>N74597</u>	<u>001</u>	Jul 29, 1998

HYDROXOCOBALAMIN

INJECTABLE; INJECTION

CYANOKIT

	+ EMD PHARMS	2.5GM/VIAL (5GM/KIT)	N22041	002	Dec 15, 2006
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HYDROXOCOBALAMIN

	+ WATSON LABS	1MG/ML	N85998	001	
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HYDROXYAMPHETAMINE HYDROBROMIDE; TROPICAMIDE

SOLUTION/DROPS; OPHTHALMIC

	+ AKORN	1%;0.25%	N19261	001	Jan 30, 1992
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HYDROXYCHLOROQUINE SULFATE

TABLET; ORAL

HYDROXYCHLOROQUINE SULFATE

<u>AB</u>	MYLAN	<u>200MG</u>	<u>N40274</u>	<u>001</u>	May 29, 1998
<u>AB</u>	SANDOZ	<u>200MG</u>	<u>N40104</u>	<u>001</u>	Nov 30, 1995
<u>AB</u>		<u>200MG</u>	<u>N40150</u>	<u>001</u>	Jan 27, 1996
<u>AB</u>	TEVA PHARMS	<u>200MG</u>	<u>N40081</u>	<u>001</u>	Sep 30, 1994
<u>AB</u>	WATSON LABS	<u>200MG</u>	<u>N40133</u>	<u>001</u>	Nov 30, 1995
	<u>PLAQUENIL</u>				
<u>AB</u>	+ SANOFI AVENTIS US	<u>200MG</u>	<u>N09768</u>	<u>001</u>	

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 202 (of 370)

HYDROXYPROPYL CELLULOSEINSERT; OPHTHALMIC
LACRISERT

+ ATON 5MG N18771 001

HYDROXYUREACAPSULE; ORAL
DROXIA

BRISTOL MYERS SQUIBB 200MG N16295 002 Feb 25, 1998

300MG N16295 003 Feb 25, 1998

+ 400MG N16295 004 Feb 25, 1998

HYDREAAB + BRISTOL MYERS SQUIBB 500MG N16295 001HYDROXYUREAAB BARR 250MG N75143 002 Sep 21, 2000AB 500MG N75143 001 Oct 16, 1998AB + DURAMED PHARMS BARR 250MG N75020 002 Jun 26, 2000AB 500MG N75020 001 Jul 30, 1998AB PAR PHARM 500MG N75340 001 Feb 24, 1999AB ROXANE 500MG N74476 001 Aug 18, 1995TABLET; ORAL
HYDROXYUREA

+ BARR 1GM N75734 001 Aug 29, 2000

HYDROXYZINE HYDROCHLORIDE

INJECTABLE; INJECTION

HYDROXYZINE HYDROCHLORIDEAP ABRAXIS PHARM 25MG/ML N87329 001AP 50MG/ML N87329 002AP HOSPIRA 25MG/ML N87416 001AP 50MG/ML N87546 001AP LUITPOLD 25MG/ML N87408 001AP 50MG/ML N87408 002AP WATSON LABS 50MG/ML N85779 001VISTARILAP + PFIZER 25MG/ML N11111 001AP + 50MG/ML N11111 002

SYRUP; ORAL

HYDROXYZINE HYDROCHLORIDEAA + ACTAVIS MID ATLANTIC 10MG/5ML N86880 001AA + HI TECH PHARMA 10MG/5ML N40010 001 Oct 28, 1994AA + MORTON GROVE 10MG/5ML N87294 001 Apr 12, 1982AA + VINTAGE PHARMS 10MG/5ML N40391 001 Apr 10, 2002

TABLET; ORAL

HYDROXYZINE HYDROCHLORIDEAB ACTAVIS TOTOWA 10MG N40600 001 Dec 28, 2004AB 25MG N40602 001 Dec 28, 2004AB 50MG N40604 001 Dec 28, 2004AB MUTUAL PHARM 10MG N89381 001 May 19, 1986AB 25MG N89382 001 May 19, 1986AB 50MG N89383 001 May 19, 1986AB + PLIVA 10MG N88617 001 Jan 10, 1986AB + 25MG N88618 001 Jan 10, 1986AB + 50MG N88619 001 Jan 10, 1986AB SANDOZ 10MG N87869 001 Dec 20, 1982AB 25MG N87870 001 Dec 20, 1982AB 50MG N87871 001 Dec 20, 1982AB VINTAGE PHARMS 10MG N40579 001 May 27, 2005AB 25MG N40574 001 May 27, 2005

PRESCRIPTION DRUG PRODUCT LIST

3 - 203 (of 370)

HYDROXYZINE HYDROCHLORIDE

TABLET; ORAL

HYDROXYZINE HYDROCHLORIDE

<u>AB</u>	VINTAGE PHARMS	<u>50MG</u>	<u>N40580</u>	<u>001</u>	May 27, 2005
<u>AB</u>	WATSON LABS	<u>10MG</u>	<u>N81149</u>	<u>001</u>	Mar 18, 1994
<u>AB</u>		<u>10MG</u>	<u>N88348</u>	<u>001</u>	Sep 15, 1983
<u>AB</u>		<u>25MG</u>	<u>N81150</u>	<u>001</u>	Mar 18, 1994
<u>AB</u>		<u>25MG</u>	<u>N88349</u>	<u>001</u>	Sep 15, 1983
<u>AB</u>		<u>50MG</u>	<u>N81151</u>	<u>001</u>	Mar 18, 1994
<u>AB</u>		<u>50MG</u>	<u>N88350</u>	<u>001</u>	Sep 15, 1983

HYDROXYZINE PAMOATE

CAPSULE; ORAL

HYDROXYZINE PAMOATE

<u>AB</u>	BARR	<u>EQ 25MG HCL</u>	<u>N88496</u>	<u>001</u>	Jun 15, 1984
<u>AB</u>		<u>EQ 50MG HCL</u>	<u>N88487</u>	<u>001</u>	Jun 15, 1984
		EQ 100MG HCL	N88488	001	Jun 15, 1984
<u>AB</u>	IVAX PHARMS	<u>EQ 25MG HCL</u>	<u>N87761</u>	<u>001</u>	Mar 05, 1982
<u>AB</u>		<u>EQ 50MG HCL</u>	<u>N87760</u>	<u>001</u>	Mar 05, 1982
<u>AB</u>	SANDOZ	<u>EQ 25MG HCL</u>	<u>N81127</u>	<u>001</u>	Jun 28, 1991
<u>AB</u>		<u>EQ 25MG HCL</u>	<u>N87479</u>	<u>001</u>	
<u>AB</u>		<u>EQ 50MG HCL</u>	<u>N86183</u>	<u>001</u>	
<u>AB</u>	WATSON LABS	<u>EQ 25MG HCL</u>	<u>N40156</u>	<u>001</u>	Jul 15, 1996
<u>AB</u>		<u>EQ 25MG HCL</u>	<u>N81165</u>	<u>001</u>	Jul 31, 1991
<u>AB</u>		<u>EQ 50MG HCL</u>	<u>N40156</u>	<u>002</u>	Jul 15, 1996
	<u>VISTARIL</u>				
<u>AB</u>	PFIZER	<u>EQ 25MG HCL</u>	<u>N11459</u>	<u>002</u>	
<u>AB</u>	+	<u>EQ 50MG HCL</u>	<u>N11459</u>	<u>004</u>	

SUSPENSION; ORAL

VISTARIL

+ PFIZER

EQ 25MG HCL/5ML

N11795 001

IBANDRONATE SODIUM

INJECTABLE; INTRAVENOUS

BONIVA

+ ROCHE

EQ 3MG BASE/3ML

N21858 001 Jan 06, 2006

TABLET; ORAL

BONIVA

+ ROCHE

EQ 2.5MG BASE

N21455 001 May 16, 2003

+

EQ 150MG BASE

N21455 002 Mar 24, 2005

IBUPROFEN

SUSPENSION; ORAL

CHILDREN'S ADVIL

BX WYETH CONS

100MG/5ML

N19833 002 Sep 19, 1989

IBUPROFEN

AB ACTAVIS MID ATLANTIC

100MG/5ML

N74978 001 Mar 25, 1998

AB PERRIGO R AND D

100MG/5ML

N76925 001 Sep 23, 2004

MOTRIN

AB + MCNEIL PED

100MG/5ML

N19842 001 Sep 19, 1989

TABLET; ORAL

IBUPROFEN

AB BASF

400MG

N75682 001 Nov 14, 2001

AB

600MG

N75682 002 Nov 14, 2001

AB

800MG

N75682 003 Nov 14, 2001

AB DR REDDYS LABS INC

400MG

N76112 001 Oct 31, 2001

AB

600MG

N76112 002 Oct 31, 2001

AB

800MG

N76112 003 Oct 31, 2001

AB INTERPHARM

400MG

N71334 001 Nov 25, 1986

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 204 (of 370)

IBUPROFEN

TABLET; ORAL

IBUPROFEN

<u>AB</u>	INTERPHARM	<u>600MG</u>	<u>N71335</u>	<u>001</u>	Nov 25, 1986
<u>AB</u>		<u>800MG</u>	<u>N71935</u>	<u>001</u>	Oct 13, 1987
<u>AB</u>	LEINER	<u>300MG</u>	<u>N71266</u>	<u>001</u>	Oct 15, 1986
<u>AB</u>		<u>400MG</u>	<u>N71267</u>	<u>001</u>	Oct 15, 1986
<u>AB</u>		<u>600MG</u>	<u>N71268</u>	<u>001</u>	Oct 15, 1986
<u>AB</u>		<u>800MG</u>	<u>N72300</u>	<u>001</u>	Jul 01, 1988
<u>AB</u>	MUTUAL PHARM	<u>300MG</u>	<u>N71230</u>	<u>001</u>	Oct 22, 1986
<u>AB</u>		<u>400MG</u>	<u>N71231</u>	<u>001</u>	Oct 22, 1986
<u>AB</u>		<u>600MG</u>	<u>N71232</u>	<u>001</u>	Oct 22, 1986
<u>AB</u>		<u>800MG</u>	<u>N72004</u>	<u>001</u>	Nov 18, 1987
<u>AB</u>	MYLAN	<u>400MG</u>	<u>N70045</u>	<u>001</u>	Sep 24, 1985
<u>AB</u>		<u>600MG</u>	<u>N70057</u>	<u>001</u>	Sep 24, 1985
<u>AB</u>		<u>800MG</u>	<u>N71999</u>	<u>001</u>	Dec 03, 1987
<u>AB</u>	OHM LABS	<u>400MG</u>	<u>N70818</u>	<u>001</u>	Dec 26, 1985
<u>AB</u>	PAR PHARM	<u>400MG</u>	<u>N70329</u>	<u>001</u>	Aug 06, 1985
<u>AB</u>		<u>600MG</u>	<u>N70330</u>	<u>001</u>	Aug 06, 1985
<u>AB</u>		<u>800MG</u>	<u>N70986</u>	<u>001</u>	Jul 25, 1986
<u>AB</u>	PERRIGO R AND D	<u>400MG</u>	<u>N77114</u>	<u>001</u>	Jul 18, 2005
<u>AB</u>		<u>600MG</u>	<u>N77114</u>	<u>002</u>	Jul 18, 2005
<u>AB</u>		<u>800MG</u>	<u>N77114</u>	<u>003</u>	Jul 18, 2005
<u>AB</u>	PLIVA	<u>400MG</u>	<u>N71666</u>	<u>001</u>	Jun 18, 1987
<u>AB</u>		<u>600MG</u>	<u>N71667</u>	<u>001</u>	Jun 18, 1987
<u>AB</u>		<u>800MG</u>	<u>N71668</u>	<u>001</u>	Jun 18, 1987
<u>AB</u>	SANDOZ	<u>300MG</u>	<u>N70734</u>	<u>001</u>	Jun 12, 1986
<u>AB</u>		<u>400MG</u>	<u>N70735</u>	<u>001</u>	Jun 12, 1986
<u>AB</u>		<u>600MG</u>	<u>N70736</u>	<u>001</u>	Jun 12, 1986
<u>AB</u>		<u>800MG</u>	<u>N72169</u>	<u>001</u>	Dec 11, 1987
<u>AB</u>	VINTAGE PHARMS	<u>400MG</u>	<u>N71644</u>	<u>001</u>	Feb 01, 1988
<u>AB</u>	WATSON LABS	<u>400MG</u>	<u>N70436</u>	<u>001</u>	Aug 21, 1985
<u>AB</u>		<u>600MG</u>	<u>N70437</u>	<u>001</u>	Aug 21, 1985
<u>AB</u>		<u>800MG</u>	<u>N71547</u>	<u>001</u>	Jul 02, 1987
<u>AB</u>	<u>IBUPROHM</u>				
<u>AB</u>	OHM LABS	<u>400MG</u>	<u>N70469</u>	<u>001</u>	Aug 29, 1985
<u>AB</u>	<u>IBU - TAB</u>				
<u>AB</u>	ALRA	<u>400MG</u>	<u>N71058</u>	<u>001</u>	Aug 11, 1988
<u>AB</u>		<u>600MG</u>	<u>N71059</u>	<u>001</u>	Aug 11, 1988
<u>AB</u>	<u>MOTRIN</u>				
<u>AB</u>	MCNEIL PED	<u>300MG</u>	<u>N17463</u>	<u>003</u>	
<u>AB</u>		<u>400MG</u>	<u>N17463</u>	<u>002</u>	
<u>AB</u>		<u>600MG</u>	<u>N17463</u>	<u>004</u>	
<u>AB</u>	+	<u>800MG</u>	<u>N17463</u>	<u>005</u>	May 22, 1985

IBUPROFEN LYSINEINJECTABLE; INTRAVENOUS
NEOPROFEN

+ FARMACON IL EQ 20MG BASE/2ML (EQ 10MG BASE/ML) N21903 001 Apr 13, 2006

IBUPROFEN; OXYCODONE HYDROCHLORIDETABLET; ORAL
COMBUNOX

+ FOREST LABS 400MG;5MG N21378 001 Nov 26, 2004

IBUTILIDE FUMARATEINJECTABLE; INJECTION
CORVERT

+ PHARMACIA AND UPJOHN 0.1MG/ML N20491 001 Dec 28, 1995

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 205 (of 370)

ICODEXTRINSOLUTION; INTRAPERITONEAL
EXTRANEAL

+	BAXTER HLTHCARE	7.5GM/100ML	N21321	001	Dec 20, 2002
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IDARUBICIN HYDROCHLORIDE

INJECTABLE; INJECTION

IDAMYCIN PFS

<u>AP</u>	+	PHARMACIA AND UPJOHN	<u>1MG/ML</u>	<u>N50734</u>	<u>001</u>	Feb 17, 1997
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IDARUBICIN HYDROCHLORIDE

<u>AP</u>		BEDFORD LABS	<u>1MG/ML</u>	<u>N65275</u>	<u>001</u>	Dec 14, 2006
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+	SICOR PHARMS	5MG/VIAL	N65037	003	May 01, 2002
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+		10MG/VIAL	N65037	002	May 01, 2002
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+		20MG/VIAL	N65037	001	May 01, 2002
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IDARUBICIN HYDROCHLORIDE PFS

<u>AP</u>		SICOR PHARMS	<u>1MG/ML</u>	<u>N65036</u>	<u>001</u>	May 01, 2002
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IDOXURIDINE

SOLUTION/DROPS; OPHTHALMIC

DENDRID

<u>AT</u>	+	ALCON	<u>0.1%</u>	<u>N14169</u>	<u>001</u>	
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HERPLEX

<u>AT</u>	+	ALLERGAN	<u>0.1%</u>	<u>N13935</u>	<u>002</u>	
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IFOSFAMIDE

INJECTABLE; INJECTION

IFOSFAMIDE

+	ABRAXIS PHARM	1GM/VIAL	N76078	001	May 28, 2002
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+		3GM/VIAL	N76078	002	May 28, 2002
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IFOSFAMIDE; MESNA

INJECTABLE; INJECTION

IFEX/MESNEX KIT

+	BRISTOL MYERS SQUIBB	1GM/VIAL;100MG/ML	N19763	003	Oct 10, 1992
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+		3GM/VIAL;100MG/ML	N19763	004	Oct 10, 1992
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INJECTABLE; INTRAVENOUS

IFOSFAMIDE/MESNA KIT

+	SICOR PHARMS	EQ 1GM /20ML(50MG/ML);EQ 1GM /10ML(100MG/ML)	N75874	001	Feb 26, 2002
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+		EQ 1GM/10ML (100MG/ML)	N75874	002	Feb 26, 2002
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ILOPROST

SOLUTION; INHALATION

VENTAVIS

+	COTHERIX	10UGM/ML (10UGM/ML)	N21779	002	Dec 08, 2005
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+		20UGM/2ML (10UGM/ML)	N21779	001	Dec 29, 2004
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IMATINIB MESYLATE

TABLET; ORAL

GLEEVEC

NOVARTIS

+		EQ 100MG BASE	N21588	001	Apr 18, 2003
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+		EQ 400MG BASE	N21588	002	Apr 18, 2003
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IMIGLUCERASE

INJECTABLE; INJECTION

CEREZYME

GENZYME

+		200 UNITS/VIAL	N20367	001	May 23, 1994
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+		400 UNITS/VIAL	N20367	002	Sep 22, 1999
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27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 206 (of 370)

IMIPRAMINE HYDROCHLORIDE

TABLET; ORAL

IMIPRAMINE HYDROCHLORIDE

<u>AB</u>	MUTUAL PHARM	<u>10MG</u>	<u>N81048</u>	<u>001</u>	Jun 05, 1990
<u>AB</u>		<u>25MG</u>	<u>N81049</u>	<u>001</u>	Jun 05, 1990
<u>AB</u>		<u>50MG</u>	<u>N81050</u>	<u>001</u>	Jun 05, 1990
<u>AB</u>	PAR PHARM	<u>10MG</u>	<u>N88292</u>	<u>001</u>	Oct 21, 1983
<u>AB</u>		<u>10MG</u>	<u>N89422</u>	<u>001</u>	Jul 14, 1987
<u>AB</u>		<u>25MG</u>	<u>N88262</u>	<u>001</u>	Oct 21, 1983
<u>AB</u>		<u>25MG</u>	<u>N89497</u>	<u>001</u>	Jul 14, 1987
<u>AB</u>		<u>50MG</u>	<u>N88276</u>	<u>001</u>	Oct 21, 1983
<u>AB</u>	SANDOZ	<u>10MG</u>	<u>N84936</u>	<u>002</u>	
<u>AB</u>		<u>25MG</u>	<u>N83745</u>	<u>001</u>	
<u>AB</u>		<u>50MG</u>	<u>N84937</u>	<u>001</u>	
<u>TOFRANIL</u>					
<u>AB</u>	TYCO HLTHCARE	<u>10MG</u>	<u>N87844</u>	<u>001</u>	May 22, 1984
<u>AB</u>		<u>25MG</u>	<u>N87845</u>	<u>001</u>	May 22, 1984
<u>AB</u>	+	<u>50MG</u>	<u>N87846</u>	<u>001</u>	May 22, 1984

IMIPRAMINE PAMOATE

CAPSULE; ORAL

TOFRANIL-PM

	TYCO HLTHCARE	EQ 75MG HCL	N17090	001	
		EQ 100MG HCL	N17090	004	
		EQ 125MG HCL	N17090	003	
	+	EQ 150MG HCL	N17090	002	

IMIQUIMOD

CREAM; TOPICAL

ALDARA

	+ 3M	5%	N20723	001	Feb 27, 1997
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INAMRINONE LACTATE

INJECTABLE; INJECTION

AMRINONE

<u>AP</u>	BEDFORD	<u>EQ 5MG BASE/ML</u>	<u>N75513</u>	<u>001</u>	May 09, 2000
<u>AP</u>	+ HOSPIRA	<u>EQ 5MG BASE/ML</u>	<u>N74616</u>	<u>001</u>	Aug 03, 1998
<u>AMRINONE LACTATE</u>					
<u>AP</u>	BAXTER HLTHCARE CORP	<u>EQ 5MG BASE/ML</u>	<u>N75542</u>	<u>001</u>	May 10, 2000

INDAPAMIDE

TABLET; ORAL

INDAPAMIDE

<u>AB</u>	ACTAVIS ELIZABETH	<u>1.25MG</u>	<u>N74722</u>	<u>001</u>	Jun 17, 1996
<u>AB</u>		<u>2.5MG</u>	<u>N74722</u>	<u>002</u>	Jun 17, 1996
<u>AB</u>	ALPHAPHARM	<u>1.25MG</u>	<u>N75105</u>	<u>001</u>	Jul 23, 1998
<u>AB</u>		<u>2.5MG</u>	<u>N75105</u>	<u>002</u>	Jul 23, 1998
<u>AB</u>	IVAX PHARMS	<u>1.25MG</u>	<u>N74299</u>	<u>002</u>	Apr 29, 1996
<u>AB</u>		<u>2.5MG</u>	<u>N74299</u>	<u>001</u>	Jul 27, 1995
<u>AB</u>	JUBILANT PHARMS	<u>1.25MG</u>	<u>N75201</u>	<u>001</u>	Dec 04, 1998
<u>AB</u>		<u>2.5MG</u>	<u>N75201</u>	<u>002</u>	Dec 04, 1998
<u>AB</u>	MYLAN	<u>1.25MG</u>	<u>N74461</u>	<u>002</u>	Mar 26, 1997
<u>AB</u>		<u>2.5MG</u>	<u>N74461</u>	<u>001</u>	Mar 27, 1996
<u>AB</u>	SANDOZ	<u>1.25MG</u>	<u>N74594</u>	<u>001</u>	May 23, 1996
<u>AB</u>		<u>2.5MG</u>	<u>N74594</u>	<u>002</u>	May 23, 1996
<u>AB</u>	TEVA	<u>1.25MG</u>	<u>N74498</u>	<u>002</u>	Feb 12, 1998
<u>AB</u>		<u>2.5MG</u>	<u>N74498</u>	<u>001</u>	Oct 31, 1996
<u>AB</u>	WATSON LABS	<u>1.25MG</u>	<u>N74585</u>	<u>001</u>	Sep 26, 1996
<u>AB</u>		<u>2.5MG</u>	<u>N74585</u>	<u>002</u>	Sep 26, 1996

PRESCRIPTION DRUG PRODUCT LIST

3 - 207 (of 370)

INDAPAMIDE

TABLET; ORAL					
<u>LOZOL</u>					
<u>AB</u>	SANOFI AVENTIS US	<u>1.25MG</u>	<u>N18538</u>	<u>002</u>	Apr 29, 1993
<u>AB</u>	+	<u>2.5MG</u>	<u>N18538</u>	<u>001</u>	Jul 06, 1983

INDINAVIR SULFATE

CAPSULE; ORAL					
CRIXIVAN					
MERCK					
		EQ 100MG BASE	N20685	006	Apr 19, 2000
		EQ 200MG BASE	N20685	003	Mar 13, 1996
		EQ 333MG BASE	N20685	005	Dec 17, 1998
	+	EQ 400MG BASE	N20685	001	Mar 13, 1996

INDIUM IN-111 CHLORIDE

INJECTABLE; INJECTION					
INDICLOR					
	+	AMERSHAM	2mCi/0.2ML	N19862	001
		INDIUM IN 111 CHLORIDE			
	+	MALLINCKRODT	5mCi/0.5ML	N19841	001
					Sep 27, 1994

INDIUM IN-111 OXYQUINOLINE

INJECTABLE; INJECTION					
INDIUM IN-111 OXYQUINOLINE					
	+	GE HEALTHCARE	1mCi/ML	N19044	001
					Dec 24, 1985

INDIUM IN-111 PENTETATE DISODIUM

INJECTABLE; INTRATHECAL					
MPI INDIUM DTPA IN 111					
	+	GE HEALTHCARE	1mCi/ML	N17707	001
					Feb 18, 1982

INDIUM IN-111 PENTETREOTIDE KIT

INJECTABLE; INJECTION					
OCTREOSCAN					
	+	MALLINCKRODT	3mCi/ML	N20314	001
					Jun 02, 1994

INDOCYANINE GREEN

INJECTABLE; INJECTION					
IC-GREEN					
	+	AKORN	25MG/VIAL	N11525	001

INDOMETHACIN

CAPSULE; ORAL					
<u>INDOMETHACIN</u>					
<u>AB</u>	+	MYLAN	<u>50MG</u>	<u>N18858</u>	<u>002</u>
<u>AB</u>			<u>50MG</u>	<u>N70624</u>	<u>001</u>
			25MG	N18858	001
					Apr 20, 1984
					Sep 04, 1985
					Apr 20, 1984
CAPSULE, EXTENDED RELEASE; ORAL					
INDOCIN SR					
	+	SANDOZ	75MG	N74464	001
					May 28, 1998
SUPPOSITORY; RECTAL					
INDOMETHACIN					
	+	G AND W LABS	50MG	N73314	001
					Aug 31, 1992
SUSPENSION; ORAL					
INDOCIN					
	+	MERCK	25MG/5ML	N18332	001
					Oct 10, 1985

PRESCRIPTION DRUG PRODUCT LIST

3 - 208 (of 370)

INDOMETHACIN SODIUM

INJECTABLE; INJECTION
INDOCIN I.V.

+ OVATION PHARMS	EQ 1MG BASE/VIAL	N18878 001	Jan 30, 1985
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INSULIN ASPART PROTAMINE RECOMBINANT; INSULIN ASPART RECOMBINANT

INJECTABLE; SUBCUTANEOUS
NOVOLOG MIX 70/30

+ NOVO NORDISK INC	70 UNITS/ML;30 UNITS/ML	N21172 001	Nov 01, 2001
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INSULIN ASPART RECOMBINANT

INJECTABLE; SUBCUTANEOUS
NOVOLOG

+ NOVO NORDISK INC	100 UNITS/ML	N20986 001	Jun 07, 2000
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INSULIN DETEMIR RECOMBINANT

INJECTABLE; SUBCUTANEOUS
LEVEMIR

+ NOVO NORDISK INC	100 UNITS/ML	N21536 001	Jun 16, 2005
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INSULIN GLARGINE RECOMBINANT

INJECTABLE; INJECTION
LANTUS

+ SANOFI AVENTIS US	100 UNITS/ML	N21081 001	Apr 20, 2000
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INSULIN GLULISINE RECOMBINANT

INJECTABLE; SUBCUTANEOUS
APIDRA

+ SANOFI AVENTIS US	1000 UNITS/10ML (100 UNITS/ML)	N21629 001	Apr 16, 2004
+	300 UNITS/3ML (100 UNITS/ML)	N21629 002	Dec 20, 2005

INSULIN LISPRO PROTAMINE RECOMBINANT; INSULIN LISPRO RECOMBINANT

INJECTABLE; INJECTION
HUMALOG MIX 50/50

+ LILLY	50 UNITS/ML;50 UNITS/ML	N21018 001	Dec 22, 1999
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+ LILLY	75 UNITS/ML;25 UNITS/ML	N21017 001	Dec 22, 1999
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INSULIN LISPRO RECOMBINANT

INJECTABLE; INJECTION
HUMALOG

+ LILLY	100 UNITS/ML	N20563 001	Jun 14, 1996
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+ LILLY	100 UNITS/ML	N20563 002	Aug 06, 1998
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INSULIN RECOMBINANT HUMAN

INJECTABLE; INJECTION
HUMULIN R

+ LILLY	500 UNITS/ML	N18780 004	Mar 31, 1994
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POWDER; INHALATION
EXUBERA

PFIZER	1MG/INH	N21868 001	Jan 27, 2006
+	3MG/INH	N21868 002	Jan 27, 2006

IOBENGUANE SULFATE I 131

INJECTABLE; INJECTION
IOBENGUANE SULFATE I 131
CIS

2.3mCi/ML	N20084 001	Mar 25, 1994
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27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 209 (of 370)

IODIPAMIDE MEGLUMINEINJECTABLE; INJECTION
CHOLOGRAFIN MEGLUMINE

+ BRACCO 52%

N09321 003

IODIXANOLINJECTABLE; INJECTION
VISIPAQUE 270+ GE HEALTHCARE 55%
55%N20351 001 Mar 22, 1996
N20808 001 Aug 29, 1997VISIPAQUE 320
+ GE HEALTHCARE 65.2%
65.2%N20351 002 Mar 22, 1996
N20808 002 Aug 29, 1997IOHEXOLINJECTABLE; INJECTION
OMNIPAQUE 140

+ GE HEALTHCARE 30.2%

N18956 005 Nov 30, 1988

SOLUTION; INJECTION, ORAL
OMNIPAQUE 350+ GE HEALTHCARE 75.5%
75.5%N18956 004 Dec 26, 1985
N20608 003 Oct 24, 1995SOLUTION; INJECTION, ORAL, RECTAL
OMNIPAQUE 180

+ GE HEALTHCARE 38.8%

N18956 001 Dec 26, 1985

OMNIPAQUE 240
+ GE HEALTHCARE 51.8%
51.8%N18956 002 Dec 26, 1985
N20608 001 Oct 24, 1995OMNIPAQUE 300
+ GE HEALTHCARE 64.7%
64.7%N18956 003 Dec 26, 1985
N20608 002 Oct 24, 1995IOPAMIDOL

INJECTABLE; INJECTION

IOPAMIDOL-200AP COOK IMAGING 41%N74881 001 Jul 28, 2000AP HOSPIRA 41%N74898 001 Dec 30, 1997IOPAMIDOL-200 IN PLASTIC CONTAINERAP HOSPIRA 41%N74636 001 Dec 30, 1997IOPAMIDOL-250AP ABRAXIS PHARM 51%N74679 001 Apr 02, 1997AP COOK IMAGING 51%N74881 002 Jul 28, 2000AP HOSPIRA 51%N74898 002 Dec 30, 1997AP 51%N75005 001 Feb 24, 1998IOPAMIDOL-250 IN PLASTIC CONTAINERAP HOSPIRA 51%N74636 002 Dec 30, 1997IOPAMIDOL-300AP ABRAXIS PHARM 61%N74679 002 Apr 02, 1997AP COOK IMAGING 61%N74881 003 Jul 28, 2000AP HOSPIRA 61%N74898 003 Dec 30, 1997AP 61%N75005 002 Feb 24, 1998IOPAMIDOL-300 IN PLASTIC CONTAINERAP HOSPIRA 61%N74636 003 Dec 30, 1997AP 61%N74637 001 Apr 03, 1997IOPAMIDOL-370AP ABRAXIS PHARM 76%N74679 003 Apr 02, 1997AP COOK IMAGING 76%N74881 004 Jul 28, 2000AP HOSPIRA 76%N74898 004 Dec 30, 1997AP 76%N75005 003 Feb 24, 1998

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 210 (of 370)

IOPAMIDOL

INJECTABLE; INJECTION

IOPAMIDOL-370 IN PLASTIC CONTAINER

<u>AP</u>	HOSPIRA	<u>76%</u>	<u>N74636</u>	<u>004</u>	Dec 30, 1997
	<u>ISOVUE-200</u>				
<u>AP</u>	+ BRACCO	<u>41%</u>	<u>N18735</u>	<u>006</u>	Jul 07, 1987
	<u>ISOVUE-250</u>				
<u>AP</u>	+ BRACCO	<u>51%</u>	<u>N18735</u>	<u>007</u>	Jul 06, 1992
<u>AP</u>	+	<u>51%</u>	<u>N20327</u>	<u>002</u>	Oct 12, 1994
	<u>ISOVUE-300</u>				
<u>AP</u>	+ BRACCO	<u>61%</u>	<u>N18735</u>	<u>002</u>	Dec 31, 1985
<u>AP</u>	+	<u>61%</u>	<u>N20327</u>	<u>003</u>	Oct 12, 1994
	<u>ISOVUE-370</u>				
<u>AP</u>	+ BRACCO	<u>76%</u>	<u>N18735</u>	<u>003</u>	Dec 31, 1985
<u>AP</u>	+	<u>76%</u>	<u>N20327</u>	<u>004</u>	Oct 12, 1994
	ISOVUE-M 200				
	+ BRACCO	41%	N18735	001	Dec 31, 1985
	ISOVUE-M 300				
	+ BRACCO	61%	N18735	004	Dec 31, 1985

IOPROMIDE

INJECTABLE; INJECTION

ULTRAVIST (PHARMACY BULK)

+	BERLEX	49.9%	N21425	003	Mar 12, 2004
+		62.3%	N21425	001	Sep 20, 2002
+		76.9%	N21425	002	Sep 20, 2002
	ULTRAVIST 150				
+	BERLEX	31.2%	N20220	004	May 10, 1995
	ULTRAVIST 240				
+	BERLEX	49.9%	N20220	003	May 10, 1995
	ULTRAVIST 300				
+	BERLEX	62.3%	N20220	002	May 10, 1995
	ULTRAVIST 370				
+	BERLEX	76.9%	N20220	001	May 10, 1995

IOTHALAMATE MEGLUMINE

INJECTABLE; INJECTION

CONRAY

+	MALLINCKRODT	60%	N13295	001	
	CONRAY 30				
+	MALLINCKRODT	30%	N16983	001	
	CONRAY 43				
+	MALLINCKRODT	43%	N13295	002	

SOLUTION; INTRAVESICAL

CYSTO-CONRAY II

MALLINCKRODT	17.2%	N17057	002	
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IOTHALAMATE SODIUM

INJECTABLE; INJECTION

CONRAY 400

+	MALLINCKRODT	66.8%	N14295	001	
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IOTHALAMATE SODIUM, I-125

INJECTABLE; INJECTION

GLOFIL-125

QOL MEDCL	250-300uCi/ML	N17279	001	
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PRESCRIPTION DRUG PRODUCT LIST

3 - 211 (of 370)

IOVERSOL

INJECTABLE; INJECTION

OPTIRAY 160				
+ MALLINCKRODT	34%	N19710	003	Dec 30, 1988
OPTIRAY 240				
+ MALLINCKRODT	51%	N19710	002	Dec 30, 1988
OPTIRAY 300				
+ MALLINCKRODT	64%	N19710	004	Jan 22, 1992
+	64%	N20923	004	May 13, 1999
OPTIRAY 320				
+ MALLINCKRODT	68%	N19710	001	Dec 30, 1988
OPTIRAY 350				
+ MALLINCKRODT	74%	N19710	005	Jan 22, 1992
+	74%	N20923	003	May 28, 1998

IOXAGLATE MEGLUMINE; IOXAGLATE SODIUM

INJECTABLE; INJECTION

HEXABRIX				
+ MALLINCKRODT	39.3%;19.6%	N18905	002	Jul 26, 1985

IOXILAN

INJECTABLE; INJECTION

OXILAN-300				
GUERBET	62%	N20316	001	Dec 21, 1995
OXILAN-350				
GUERBET	73%	N20316	002	Dec 21, 1995

IPRATROPIUM BROMIDE

AEROSOL, METERED; INHALATION

ATROVENT HFA				
+ BOEHRINGER INGELHEIM	0.021MG/INH	N21527	001	Nov 27, 2004

SOLUTION; INHALATION

IPRATROPIUM BROMIDE

<u>AN</u>	ACTAVIS MID ATLANTIC	<u>0.02%</u>	<u>N75111</u>	<u>001</u>	Apr 22, 1999
<u>AN</u>	BAUSCH AND LOMB	<u>0.02%</u>	<u>N75835</u>	<u>001</u>	Oct 15, 2001
<u>AN</u>	+ DEY	<u>0.02%</u>	<u>N74755</u>	<u>001</u>	Jan 10, 1997
<u>AN</u>	HOLOPACK INTL	<u>0.02%</u>	<u>N75693</u>	<u>001</u>	Jan 26, 2001
<u>AN</u>	IVAX PHARMS	<u>0.02%</u>	<u>N75313</u>	<u>001</u>	Feb 07, 2000
<u>AN</u>	NEPHRON	<u>0.02%</u>	<u>N75562</u>	<u>001</u>	Sep 27, 2001
<u>AN</u>	NOVEX	<u>0.02%</u>	<u>N75441</u>	<u>001</u>	Mar 28, 2001
<u>AN</u>	RESPIRARE	<u>0.02%</u>	<u>N76291</u>	<u>001</u>	May 09, 2005
<u>AN</u>	RXELITE	<u>0.02%</u>	<u>N77072</u>	<u>001</u>	Jul 19, 2005
<u>AN</u>	WARRICK PHARMS	<u>0.02%</u>	<u>N75507</u>	<u>001</u>	Jan 19, 2001

SPRAY, METERED; NASAL

ATROVENT

<u>AB</u>	+ BOEHRINGER INGELHEIM	<u>0.021MG/SPRAY</u>	<u>N20393</u>	<u>001</u>	Oct 20, 1995
<u>AB</u>	+	<u>0.042MG/SPRAY</u>	<u>N20394</u>	<u>001</u>	Oct 20, 1995

IPRATROPIUM BROMIDE

<u>AB</u>	BAUSCH AND LOMB	<u>0.021MG/SPRAY</u>	<u>N76025</u>	<u>001</u>	Mar 31, 2003
<u>AB</u>		<u>0.042MG/SPRAY</u>	<u>N76103</u>	<u>001</u>	Mar 31, 2003
<u>AB</u>	DEY	<u>0.021MG/SPRAY</u>	<u>N75552</u>	<u>001</u>	Mar 31, 2003
<u>AB</u>		<u>0.042MG/SPRAY</u>	<u>N75553</u>	<u>001</u>	Mar 31, 2003
<u>AB</u>	NOVEX	<u>0.021MG/SPRAY</u>	<u>N76156</u>	<u>001</u>	Apr 18, 2003
<u>AB</u>		<u>0.042MG/SPRAY</u>	<u>N76155</u>	<u>001</u>	Apr 18, 2003
<u>AB</u>	ROXANE	<u>0.021MG/SPRAY</u>	<u>N76664</u>	<u>001</u>	Nov 05, 2003
<u>AB</u>		<u>0.042MG/SPRAY</u>	<u>N76598</u>	<u>001</u>	Nov 05, 2003

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 212 (of 370)

IRBESARTAN

TABLET; ORAL

AVAPRO

SANOFI AVENTIS US

75MG

N20757 001

Sep 30, 1997

150MG

N20757 002

Sep 30, 1997

+

300MG

N20757 003

Sep 30, 1997

IRINOTECAN HYDROCHLORIDE

INJECTABLE; INJECTION

CAMPTOSAR

+ PHARMACIA AND UPJOHN 20MG/ML

N20571 001

Jun 14, 1996

IRON DEXTRAN

INJECTABLE; INJECTION

DEXFERRUM

BP LUITPOLD

EQ 50MG IRON/ML

N40024 001

Feb 23, 1996

INFED

BP + WATSON LABS (UTAH)

EQ 50MG IRON/ML

N17441 001

PROFERDEX

BP NEW RIVER

EQ 50MG IRON/ML

N17807 001

IRON SUCROSE

INJECTABLE; INTRAVENOUS

VENOFER

+ LUITPOLD

EQ 100MG BASE/5ML (EQ 20MG BASE/ML)

N21135 001

Nov 06, 2000

EQ 50MG BASE/2.5ML (EQ 20MG BASE/ML)

N21135 002

Mar 20, 2005

EQ 75MG BASE/3.75ML (EQ 20MG BASE/ML)

N21135 003

Mar 29, 2005

ISOCARBOXAZID

TABLET; ORAL

MARPLAN

+ OXFORD PHARM

10MG

N11961 001

ISOETHARINE HYDROCHLORIDE

SOLUTION; INHALATION

BETA-2

+ NEPHRON

1%

N86711 001

ISOFLURANE

LIQUID; INHALATION

FORANEAN + BAXTER HLTHCARE CORP99.9%N17624 001ISOFLURANEAN HALOCARBON PRODS99.9%N75225 001

Oct 20, 1999

AN HOSPIRA99.9%N74097 001

Jan 25, 1993

AN MARSAM PHARMS LLC99.9%N74393 001

May 12, 1995

AN MINRAD99.9%N74416 001

Sep 30, 1994

AN RHODIA99.9%N74502 001

Jun 27, 1995

ISONIAZID

INJECTABLE; INJECTION

ISONIAZIDAP SANDOZ100MG/MLN40648 001

Jul 05, 2005

NYDRAZIDAP + SANDOZ100MG/MLN08662 001

SYRUP; ORAL

ISONIAZID

+ CAROLINA MEDCL

50MG/5ML

N88235 001

Nov 10, 1983

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 213 (of 370)

ISONIAZID

TABLET; ORAL

ISONIAZID

<u>AA</u>	BARR	<u>100MG</u>	<u>N80936</u>	<u>001</u>	
<u>AA</u>		<u>300MG</u>	<u>N80937</u>	<u>002</u>	
<u>AA</u>	DURAMED PHARMS BARR	<u>300MG</u>	<u>N88119</u>	<u>001</u>	Mar 17, 1983
<u>AA</u>	MIKART	<u>100MG</u>	<u>N40090</u>	<u>001</u>	Jun 26, 1997
<u>AA</u>		<u>300MG</u>	<u>N40090</u>	<u>002</u>	Jun 26, 1997
<u>AA</u>	MUTUAL PHARM	<u>300MG</u>	<u>N83633</u>	<u>001</u>	
<u>AA</u>	+ SANDOZ	<u>100MG</u>	<u>N08678</u>	<u>002</u>	
<u>AA</u>	+	<u>300MG</u>	<u>N08678</u>	<u>003</u>	
<u>AA</u>	WATSON LABS	<u>300MG</u>	<u>N80521</u>	<u>001</u>	
<u>AA</u>	WEST WARD	<u>100MG</u>	<u>N80212</u>	<u>001</u>	
<u>AA</u>		<u>300MG</u>	<u>N87425</u>	<u>001</u>	
<u>LANIAZID</u>					
<u>AA</u>	LANNETT	<u>100MG</u>	<u>N80140</u>	<u>002</u>	
<u>AA</u>		<u>300MG</u>	<u>N89776</u>	<u>001</u>	Jun 13, 1988
		50MG	N80140	001	

ISONIAZID; PYRAZINAMIDE; RIFAMPIN

TABLET; ORAL

RIFATER

+	SANOFI AVENTIS US	50MG;300MG;120MG	N50705	001	May 31, 1994
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ISONIAZID; RIFAMPIN

CAPSULE; ORAL

RIFAMATE

<u>AB</u>	+ SANOFI AVENTIS US	<u>150MG;300MG</u>	<u>N61884</u>	<u>001</u>	
<u>RIFAMPIN AND ISONIAZID</u>					
<u>AB</u>	WESTWARD	<u>150MG;300MG</u>	<u>N65221</u>	<u>001</u>	Jul 29, 2005

ISOPROTERENOL HYDROCHLORIDE

INJECTABLE; INJECTION

ISOPROTERENOL HYDROCHLORIDE

<u>AP</u>	HOSPIRA	<u>0.2MG/ML</u>	<u>N83346</u>	<u>001</u>	
		0.02MG/ML	N83283	001	
<u>AP</u>	INTL MEDICATION	<u>0.2MG/ML</u>	<u>N83724</u>	<u>001</u>	
<u>ISUPREL</u>					
<u>AP</u>	+ HOSPIRA	<u>0.2MG/ML</u>	<u>N10515</u>	<u>001</u>	

ISOSORBIDE DINITRATE

CAPSULE, EXTENDED RELEASE; ORAL

DILATRATE-SR

+	SCHWARZ PHARMA	40MG	N19790	001	Sep 02, 1988
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TABLET; ORAL

ISORDIL

<u>AB</u>	BIOVAIL	<u>5MG</u>	<u>N12093</u>	<u>007</u>	Jul 29, 1988
<u>AB</u>		<u>10MG</u>	<u>N12093</u>	<u>002</u>	Jul 29, 1988
<u>AB</u>		<u>20MG</u>	<u>N12093</u>	<u>006</u>	Jul 29, 1988
<u>AB</u>	+	<u>30MG</u>	<u>N12093</u>	<u>005</u>	Jul 29, 1988
	+	40MG	N12093	001	Jul 29, 1988

ISOSORBIDE DINITRATE

<u>AB</u>	PAR PHARM	<u>5MG</u>	<u>N86923</u>	<u>001</u>	Mar 12, 1987
<u>AB</u>		<u>10MG</u>	<u>N86925</u>	<u>001</u>	Mar 12, 1987
<u>AB</u>		<u>20MG</u>	<u>N87537</u>	<u>001</u>	Oct 02, 1987
<u>AB</u>		<u>30MG</u>	<u>N87946</u>	<u>001</u>	Jan 12, 1988
<u>AB</u>	SANDOZ	<u>5MG</u>	<u>N86221</u>	<u>001</u>	Jan 07, 1988
<u>AB</u>		<u>10MG</u>	<u>N86223</u>	<u>001</u>	Jan 07, 1988
<u>AB</u>		<u>20MG</u>	<u>N89367</u>	<u>001</u>	Apr 07, 1988

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 214 (of 370)

ISOSORBIDE DINITRATE

TABLET; ORAL

ISOSORBIDE DINITRATE

<u>AB</u>	WATSON LABS	<u>5MG</u>	<u>N86034</u>	<u>001</u>	Jan 06, 1988
<u>AB</u>		<u>10MG</u>	<u>N86032</u>	<u>001</u>	Jan 07, 1988
<u>AB</u>	WEST WARD	<u>5MG</u>	<u>N86067</u>	<u>001</u>	Oct 29, 1987
<u>AB</u>		<u>10MG</u>	<u>N86066</u>	<u>001</u>	Oct 29, 1987
<u>AB</u>		<u>20MG</u>	<u>N88088</u>	<u>001</u>	Nov 02, 1987

TABLET; SUBLINGUAL

ISORDIL

<u>AB</u>	BIOVAIL	<u>2.5MG</u>	<u>N12940</u>	<u>004</u>	Jul 29, 1988
<u>AB</u>		<u>5MG</u>	<u>N12940</u>	<u>003</u>	Jul 29, 1988
	+	10MG	N12940	005	Jul 29, 1988

ISOSORBIDE DINITRATE

<u>AB</u>	SANDOZ	<u>2.5MG</u>	<u>N86225</u>	<u>001</u>	Feb 19, 1988
<u>AB</u>		<u>5MG</u>	<u>N86222</u>	<u>001</u>	Feb 19, 1988
<u>AB</u>	WATSON LABS	<u>2.5MG</u>	<u>N86033</u>	<u>001</u>	Feb 26, 1988
<u>AB</u>		<u>5MG</u>	<u>N86031</u>	<u>001</u>	Sep 29, 1987
<u>AB</u>	WEST WARD	<u>2.5MG</u>	<u>N86054</u>	<u>001</u>	Oct 29, 1987
<u>AB</u>		<u>5MG</u>	<u>N86055</u>	<u>001</u>	Nov 02, 1987

TABLET, EXTENDED RELEASE; ORAL

ISOSORBIDE DINITRATE

	+	INWOOD LABS	40MG	N40009	001	Dec 30, 1998
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ISOSORBIDE MONONITRATE

TABLET; ORAL

ISMO

<u>AB</u>	DR REDDYS LABS INC	<u>20MG</u>	<u>N19091</u>	<u>001</u>	Dec 30, 1991
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ISOSORBIDE MONONITRATE

<u>AB</u>	ACTAVIS ELIZABETH	<u>10MG</u>	<u>N75037</u>	<u>002</u>	Oct 30, 1998
<u>AB</u>		<u>20MG</u>	<u>N75037</u>	<u>001</u>	Oct 30, 1998
<u>AB</u>	TEVA	<u>20MG</u>	<u>N75147</u>	<u>001</u>	Nov 27, 1998
<u>AB</u>	WEST WARD	<u>20MG</u>	<u>N75361</u>	<u>001</u>	Oct 05, 2000

MONOKET

<u>AB</u>	SCHWARZ	<u>10MG</u>	<u>N20215</u>	<u>002</u>	Jun 30, 1993
<u>AB</u>	+	<u>20MG</u>	<u>N20215</u>	<u>001</u>	Jun 30, 1993

TABLET, EXTENDED RELEASE; ORAL

IMDUR

<u>AB</u>	SCHERING PLOUGH	<u>30MG</u>	<u>N20225</u>	<u>001</u>	Aug 12, 1993
<u>AB</u>		<u>60MG</u>	<u>N20225</u>	<u>002</u>	Aug 12, 1993
<u>AB</u>	+	<u>120MG</u>	<u>N20225</u>	<u>003</u>	Mar 30, 1995

ISOSORBIDE MONONITRATE

<u>AB</u>	ACTAVIS ELIZABETH	<u>30MG</u>	<u>N75306</u>	<u>001</u>	Dec 31, 1998
<u>AB</u>		<u>60MG</u>	<u>N75306</u>	<u>002</u>	Dec 31, 1998
<u>AB</u>	BRIGHTSTONE	<u>60MG</u>	<u>N75166</u>	<u>001</u>	Oct 07, 1999
<u>AB</u>	DEXCEL LTD	<u>60MG</u>	<u>N75522</u>	<u>001</u>	Apr 17, 2000
<u>AB</u>	ELAN PHARM	<u>60MG</u>	<u>N75041</u>	<u>001</u>	Sep 22, 1998
<u>AB</u>	KREMERS URBAN	<u>30MG</u>	<u>N75155</u>	<u>002</u>	Jan 13, 2000
<u>AB</u>		<u>60MG</u>	<u>N75155</u>	<u>001</u>	Oct 30, 1998
<u>AB</u>		<u>120MG</u>	<u>N75155</u>	<u>003</u>	Aug 04, 2000
<u>AB</u>	KV PHARM	<u>30MG</u>	<u>N75395</u>	<u>001</u>	Mar 16, 2000
<u>AB</u>		<u>60MG</u>	<u>N75395</u>	<u>002</u>	Mar 16, 2000
<u>AB</u>		<u>120MG</u>	<u>N75395</u>	<u>003</u>	Mar 16, 2000
<u>AB</u>	WEST WARD	<u>30MG</u>	<u>N76813</u>	<u>002</u>	Mar 30, 2006
<u>AB</u>		<u>60MG</u>	<u>N76813</u>	<u>001</u>	Jan 07, 2005

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 215 (of 370)

ISOSULFAN BLUE

INJECTABLE; INJECTION

LYMPHAZURIN

+ US SURGCL 1%

N18310 001

ISOTRETINOIN

CAPSULE; ORAL

ACCUTANEAB HLR 10MGN18662 002 May 07, 1982AB + 20MGN18662 004 Mar 28, 1983AB + 40MGN18662 003 May 07, 1982AMNESTEEMAB GENPHARM 10MGN75945 001 Nov 08, 2002AB 20MGN75945 002 Nov 08, 2002AB 40MGN75945 003 Nov 08, 2002CLARAVISAB BARR 10MGN76356 001 Apr 11, 2003AB 20MGN76135 002 Apr 11, 2003AB 30MGN76135 003 May 11, 2006AB 40MGN76135 001 Apr 11, 2003SOTRETAB RANBAXY 10MGN76041 001 Dec 24, 2002AB 20MGN76041 002 Dec 24, 2002AB 30MGN76503 001 Jun 20, 2003AB 40MGN76041 003 Dec 24, 2002ISRADIPINE

CAPSULE; ORAL

ISRADIPINEAB ABRIKA PHARMS 2.5MGN77317 001 Jan 05, 2006AB + 5MGN77317 002 Jan 05, 2006AB ACTAVIS TOTOWA 2.5MGN77169 001 Apr 24, 2006AB 5MGN77169 002 Apr 24, 2006

TABLET, EXTENDED RELEASE; ORAL

DYNACIRC CR

RELIANT PHARMS 5MG

N20336 001 Jun 01, 1994

+ 10MG

N20336 002 Jun 01, 1994

ITRACONAZOLE

CAPSULE; ORAL

ITRACONAZOLEAB SANDOZ 100MGN76104 001 May 28, 2004SPORANOXAB + JANSSEN PHARMA 100MGN20083 001 Sep 11, 1992

INJECTABLE; INJECTION

SPORANOX

+ JANSSEN PHARMA 10MG/ML

N20966 001 Mar 30, 1999

SOLUTION; ORAL

SPORANOX

+ JANSSEN PHARMA 10MG/ML

N20657 001 Feb 21, 1997

IVERMECTIN

TABLET; ORAL

STROMECTOL

+ MERCK 3MG

N50742 002 Oct 08, 1998

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 216 (of 370)

KANAMYCIN SULFATE

INJECTABLE; INJECTION

KANAMYCIN

<u>AP</u>	ABRAXIS PHARM	<u>EQ 500MG BASE/2ML</u>	<u>N65111</u>	<u>001</u>	Dec 17, 2002
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<u>AP</u>		<u>EQ 1GM BASE/3ML</u>	<u>N65111</u>	<u>002</u>	Dec 17, 2002
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KANTREX

<u>AP</u>	+ APOTHECON	<u>EQ 500MG BASE/2ML</u>	<u>N61901</u>	<u>001</u>	
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<u>AP</u>	+	<u>EQ 1GM BASE/3ML</u>	<u>N61901</u>	<u>002</u>	
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	+	<u>EQ 75MG BASE/2ML</u>	<u>N61901</u>	<u>003</u>	
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KETAMINE HYDROCHLORIDE

INJECTABLE; INJECTION

KETALAR

<u>AP</u>	+ PARKEDALE	<u>EQ 50MG BASE/ML</u>	<u>N16812</u>	<u>002</u>	
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<u>AP</u>	+	<u>EQ 100MG BASE/ML</u>	<u>N16812</u>	<u>003</u>	
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	+	<u>EQ 10MG BASE/ML</u>	<u>N16812</u>	<u>001</u>	
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KETAMINE HYDROCHLORIDE

<u>AP</u>	BEDFORD	<u>EQ 50MG BASE/ML</u>	<u>N74524</u>	<u>001</u>	Mar 22, 1996
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<u>AP</u>		<u>EQ 100MG BASE/ML</u>	<u>N74524</u>	<u>002</u>	Mar 22, 1996
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<u>AP</u>	BIONICHE (CANADA)	<u>EQ 50MG BASE/ML</u>	<u>N76092</u>	<u>002</u>	Dec 28, 2001
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<u>AP</u>		<u>EQ 100MG BASE/ML</u>	<u>N76092</u>	<u>003</u>	Oct 25, 2002
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<u>AP</u>	HOSPIRA	<u>EQ 50MG BASE/ML</u>	<u>N74549</u>	<u>001</u>	Jun 27, 1996
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<u>AP</u>		<u>EQ 100MG BASE/ML</u>	<u>N74549</u>	<u>002</u>	Jun 27, 1996
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KETOCONAZOLE

CREAM; TOPICAL

KETOCONAZOLE

<u>AB</u>	ALTANA	<u>2%</u>	<u>N76294</u>	<u>001</u>	Apr 28, 2004
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<u>AB</u>	+ TEVA	<u>2%</u>	<u>N75581</u>	<u>001</u>	Apr 25, 2000
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KETOZOLE

<u>AB</u>	TARO	<u>2%</u>	<u>N75638</u>	<u>001</u>	Dec 18, 2002
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GEL; TOPICAL

XOLEGEL

	+ BARRIER THERAP	<u>2%</u>	<u>N21946</u>	<u>001</u>	Jul 28, 2006
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SHAMPOO; TOPICAL

KETOCONAZOLE

<u>AB</u>	PERRIGO NEW YORK	<u>2%</u>	<u>N76419</u>	<u>001</u>	Jan 07, 2004
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<u>AB</u>	QLT USA	<u>2%</u>	<u>N76942</u>	<u>001</u>	Apr 11, 2005
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NIZORAL

<u>AB</u>	+ MCNEIL PED	<u>2%</u>	<u>N19927</u>	<u>001</u>	Aug 31, 1990
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TABLET; ORAL

KETOCONAZOLE

<u>AB</u>	AAIPHARMA LLC	<u>200MG</u>	<u>N75341</u>	<u>001</u>	Jul 27, 1999
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<u>AB</u>	MUTUAL PHARMA	<u>200MG</u>	<u>N75314</u>	<u>001</u>	Jun 15, 1999
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<u>AB</u>	MYLAN	<u>200MG</u>	<u>N75597</u>	<u>001</u>	Dec 23, 1999
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<u>AB</u>	PLIVA	<u>200MG</u>	<u>N75362</u>	<u>001</u>	Jun 15, 1999
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<u>AB</u>	TARO	<u>200MG</u>	<u>N75319</u>	<u>001</u>	Jun 15, 1999
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<u>AB</u>	TEVA	<u>200MG</u>	<u>N74971</u>	<u>001</u>	Jun 15, 1999
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<u>AB</u>		<u>200MG</u>	<u>N75273</u>	<u>001</u>	Jun 15, 1999
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<u>AB</u>	TORPHARM	<u>200MG</u>	<u>N75912</u>	<u>001</u>	Jan 10, 2002
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NIZORAL

<u>AB</u>	+ JANSSEN PHARMA	<u>200MG</u>	<u>N18533</u>	<u>001</u>	
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KETOPROFEN

CAPSULE; ORAL

KETOPROFEN

<u>AB</u>	MYLAN	<u>50MG</u>	<u>N74035</u>	<u>002</u>	Dec 31, 1996
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<u>AB</u>		<u>75MG</u>	<u>N74035</u>	<u>003</u>	Dec 31, 1996
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<u>AB</u>	RADIUS PHARMS	<u>25MG</u>	<u>N74014</u>	<u>001</u>	Jan 29, 1993
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27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 217 (of 370)

KETOPROFEN

CAPSULE; ORAL

KETOPROFEN

<u>AB</u>	RADIUS PHARMS	<u>50MG</u>	<u>N74014</u>	<u>002</u>	Jan 29, 1993
<u>AB</u>		<u>75MG</u>	<u>N74014</u>	<u>003</u>	Jan 29, 1993
<u>AB</u>	SANDOZ	<u>50MG</u>	<u>N74024</u>	<u>001</u>	Dec 29, 1995
<u>AB</u>		<u>75MG</u>	<u>N74024</u>	<u>002</u>	Dec 29, 1995
<u>AB</u>	TEVA	<u>25MG</u>	<u>N73515</u>	<u>001</u>	Dec 22, 1992
<u>AB</u>		<u>50MG</u>	<u>N73516</u>	<u>001</u>	Dec 22, 1992
<u>AB</u>	+	<u>75MG</u>	<u>N73517</u>	<u>001</u>	Dec 22, 1992

CAPSULE, EXTENDED RELEASE; ORAL

KETOPROFEN

<u>AB</u>	ANDRX PHARMS	<u>100MG</u>	<u>N75270</u>	<u>002</u>	Mar 24, 1999
<u>AB</u>		<u>150MG</u>	<u>N75270</u>	<u>003</u>	Mar 24, 1999
<u>AB</u>		<u>200MG</u>	<u>N75270</u>	<u>001</u>	Mar 24, 1999
<u>AB</u>	ELAN PHARM	<u>200MG</u>	<u>N74879</u>	<u>001</u>	Dec 10, 1997
<u>AB</u>	MYLAN	<u>100MG</u>	<u>N75679</u>	<u>003</u>	Feb 20, 2002
<u>AB</u>		<u>150MG</u>	<u>N75679</u>	<u>002</u>	Feb 20, 2002
<u>AB</u>		<u>200MG</u>	<u>N75679</u>	<u>001</u>	Feb 20, 2002

ORUVAIL

<u>AB</u>	+	WYETH PHARMS INC	<u>200MG</u>	<u>N19816</u>	<u>001</u>	Sep 24, 1993
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KETOROLAC TROMETHAMINE

INJECTABLE; INJECTION

KETOROLAC TROMETHAMINE

AP	ABRAXIS PHARM	15MG/ML	N75784	001	Jan 11, 2002
AP		30MG/ML	N75784	002	Jan 11, 2002
AP	AMPHASTAR PHARM	15MG/ML	N76209	001	Jul 21, 2004
AP		30MG/ML	N76209	002	Jul 21, 2004
AP	APOTEX INC	15MG/ML	N75631	002	Jun 29, 2001
AP		30MG/ML	N75626	001	Jul 24, 2001
AP		30MG/ML	N75631	001	Jun 29, 2001
AP		30MG/ML	N77201	001	Oct 14, 2005
AP	BAXTER HLTHCARE	15MG/ML	N75772	001	Jul 21, 2004
AP		30MG/ML	N75772	002	Jul 21, 2004
AP	BAXTER HLTHCARE CORP	15MG/ML	N75299	001	Nov 03, 1999
AP		30MG/ML	N75299	002	Nov 03, 1999
AP +	BEDFORD	15MG/ML	N75222	001	Apr 26, 1999
AP +		30MG/ML	N75222	002	Apr 26, 1999
AP +		30MG/ML	N75228	001	Apr 26, 1999
AP	GLAND PHARMA LTD	15MG/ML	N76722	001	Jul 27, 2004
AP		30MG/ML	N76722	002	Jul 27, 2004
AP	HOSPIRA	15MG/ML	N74802	001	Jun 05, 1997
AP		15MG/ML	N74993	001	Jan 27, 1999
AP		30MG/ML	N74802	002	Jun 05, 1997
AP		30MG/ML	N74993	002	Jan 27, 1999
AP	SANDOZ	15MG/ML	N76271	001	Oct 06, 2004
AP		30MG/ML	N76271	002	Oct 06, 2004

SOLUTION/DROPS; OPHTHALMIC

+	ALLERGAN	0.5%	N19700	001	Nov 09, 1992
	ACULAR LS				
+	ALLERGAN	0.4%	N21528	001	May 30, 2003
	ACULAR PRESERVATIVE FREE				
+	ALLERGAN	0.5%	N20811	001	Nov 03, 1997

TABLET; ORAL

KETOROLAC TROMETHAMINE

<u>AB</u>	MYLAN	<u>10MG</u>	<u>N74761</u>	<u>001</u>	May 16, 1997
<u>AB</u>	PLIVA	<u>10MG</u>	<u>N75284</u>	<u>001</u>	Jun 23, 1999

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 218 (of 370)

KETOROLAC TROMETHAMINE

TABLET; ORAL

KETOROLAC TROMETHAMINE

<u>AB</u>	TEVA	<u>10MG</u>	<u>N74754</u>	<u>001</u>	May 16, 1997
<u>AB</u>	WATSON LABS	<u>10MG</u>	<u>N74955</u>	<u>001</u>	Sep 19, 1997
<u>TORADOL</u>					
<u>AB</u>	+ ROCHE PALO	<u>10MG</u>	<u>N19645</u>	<u>001</u>	Dec 20, 1991

KETOTIFEN FUMARATE

SOLUTION/DROPS; OPHTHALMIC

APOTEX INC	EQ 0.025% BASE	N77354	001	May 09, 2006
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KUNECATECHINS

OINTMENT; TOPICAL

VEREGEN

+ MEDIGENE	15%	N21902	001	Oct 31, 2006
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LABETALOL HYDROCHLORIDE

INJECTABLE; INJECTION

LABETALOL HYDROCHLORIDE

<u>AP</u>	APOTEX INC	<u>5MG/ML</u>	<u>N76051</u>	<u>001</u>	Jul 05, 2002
<u>AP</u>	BEDFORD	<u>5MG/ML</u>	<u>N75303</u>	<u>001</u>	May 28, 1999
<u>AP</u>	HOSPIRA	<u>5MG/ML</u>	<u>N75239</u>	<u>001</u>	Nov 29, 1999
<u>AP</u>		<u>5MG/ML</u>	<u>N75240</u>	<u>001</u>	Nov 29, 1999
<u>AP</u>	MAYNE PHARMA USA	<u>5MG/ML</u>	<u>N75242</u>	<u>001</u>	Sep 30, 1999
<u>AP</u>	TAYLOR	<u>5MG/ML</u>	<u>N75431</u>	<u>001</u>	Nov 29, 1999
<u>AP</u>		<u>5MG/ML</u>	<u>N75524</u>	<u>001</u>	Nov 29, 1999
<u>TRANDATE</u>					
<u>AP</u>	+ PROMETHEUS LABS	<u>5MG/ML</u>	<u>N19425</u>	<u>001</u>	Dec 31, 1985

TABLET; ORAL

LABETALOL HYDROCHLORIDE

<u>AB</u>	IVAX PHARMS	<u>100MG</u>	<u>N74787</u>	<u>001</u>	Aug 03, 1998
<u>AB</u>		<u>200MG</u>	<u>N74787</u>	<u>002</u>	Aug 03, 1998
<u>AB</u>		<u>300MG</u>	<u>N74787</u>	<u>003</u>	Aug 03, 1998
<u>AB</u>	MUTUAL PHARM	<u>100MG</u>	<u>N75215</u>	<u>001</u>	Jul 29, 1999
<u>AB</u>		<u>200MG</u>	<u>N75215</u>	<u>002</u>	Jul 29, 1999
<u>AB</u>		<u>300MG</u>	<u>N75215</u>	<u>003</u>	Jul 29, 1999
<u>AB</u>	SANDOZ	<u>100MG</u>	<u>N75113</u>	<u>001</u>	Aug 04, 1998
<u>AB</u>		<u>200MG</u>	<u>N75113</u>	<u>002</u>	Aug 04, 1998
<u>AB</u>		<u>300MG</u>	<u>N75113</u>	<u>003</u>	Aug 04, 1998
<u>AB</u>	TEVA	<u>100MG</u>	<u>N74989</u>	<u>001</u>	Sep 30, 1998
<u>AB</u>		<u>200MG</u>	<u>N74989</u>	<u>002</u>	Sep 30, 1998
<u>AB</u>		<u>300MG</u>	<u>N74989</u>	<u>003</u>	Sep 30, 1998
<u>AB</u>	WATSON LABS	<u>100MG</u>	<u>N75133</u>	<u>001</u>	Aug 03, 1998
<u>AB</u>		<u>200MG</u>	<u>N75133</u>	<u>002</u>	Aug 03, 1998
<u>AB</u>		<u>300MG</u>	<u>N75133</u>	<u>003</u>	Aug 03, 1998
<u>TRANDATE</u>					
<u>AB</u>	PROMETHEUS LABS	<u>100MG</u>	<u>N18716</u>	<u>001</u>	May 24, 1985
<u>AB</u>		<u>200MG</u>	<u>N18716</u>	<u>002</u>	Aug 01, 1984
<u>AB</u>	+	<u>300MG</u>	<u>N18716</u>	<u>003</u>	Aug 01, 1984

LACTULOSE

FOR SOLUTION; ORAL

LACTULOSE

+ INALCO	10GM/PACKET	N74712	001	Dec 10, 1997
+	20GM/PACKET	N74712	002	Dec 10, 1997

PRESCRIPTION DRUG PRODUCT LIST

3 - 219 (of 370)

LACTULOSE

SOLUTION; ORAL					
<u>CONSTILAC</u>					
<u>AA</u>	ALRA	<u>10GM/15ML</u>	<u>N71054</u>	<u>001</u>	Jul 26, 1988
<u>CONSTULOSE</u>					
<u>AA</u>	+ ACTAVIS MID ATLANTIC	<u>10GM/15ML</u>	<u>N70288</u>	<u>001</u>	Aug 15, 1988
<u>EVALOSE</u>					
<u>AA</u>	TEVA PHARMS	<u>10GM/15ML</u>	<u>N73497</u>	<u>001</u>	May 28, 1993
<u>LACTULOSE</u>					
<u>AA</u>	HI TECH PHARMA	<u>10GM/15ML</u>	<u>N74076</u>	<u>001</u>	Jul 03, 1995
<u>AA</u>	MORTON GROVE	<u>10GM/15ML</u>	<u>N74602</u>	<u>001</u>	Nov 14, 1996
<u>AA</u>	NOVEX	<u>10GM/15ML</u>	<u>N75911</u>	<u>001</u>	Feb 21, 2002
<u>AA</u>	PHARM ASSOC	<u>10GM/15ML</u>	<u>N74623</u>	<u>001</u>	Jul 30, 1996
<u>AA</u>	ROXANE	<u>10GM/15ML</u>	<u>N73591</u>	<u>001</u>	May 29, 1992
<u>AA</u>	VINTAGE PHARMS	<u>10GM/15ML</u>	<u>N75993</u>	<u>001</u>	Jul 26, 2001
<u>AA</u>	VISTAPHARM	<u>10GM/15ML</u>	<u>N74138</u>	<u>001</u>	Sep 30, 1992
<u>LAXILOSE</u>					
<u>AA</u>	TECHNILAB	<u>10GM/15ML</u>	<u>N73686</u>	<u>001</u>	May 28, 1993
SOLUTION; ORAL, RECTAL					
<u>ACILAC</u>					
<u>AA</u>	TECHNILAB	<u>10GM/15ML</u>	<u>N73685</u>	<u>001</u>	May 28, 1993
<u>CHOLAC</u>					
<u>AA</u>	ALRA	<u>10GM/15ML</u>	<u>N71331</u>	<u>001</u>	Jul 26, 1988
<u>ENULOSE</u>					
<u>AA</u>	+ ACTAVIS MID ATLANTIC	<u>10GM/15ML</u>	<u>N71548</u>	<u>001</u>	Aug 15, 1988
<u>GENERLAC</u>					
<u>AA</u>	MORTON GROVE	<u>10GM/15ML</u>	<u>N74603</u>	<u>001</u>	Oct 31, 1996
<u>HEPTALAC</u>					
<u>AA</u>	TEVA PHARMS	<u>10GM/15ML</u>	<u>N73504</u>	<u>001</u>	May 28, 1993
<u>LACTULOSE</u>					
<u>AA</u>	HI TECH PHARMA	<u>10GM/15ML</u>	<u>N74077</u>	<u>001</u>	Jul 03, 1995
<u>AA</u>	NOVEX	<u>10GM/15ML</u>	<u>N76645</u>	<u>001</u>	Jul 28, 2003

LAMIVUDINE

SOLUTION; ORAL					
EPIVIR					
	+ GLAXOSMITHKLINE	10MG/ML	N20596	001	Nov 17, 1995
EPIVIR-HBV					
	+ GLAXOSMITHKLINE	5MG/ML	N21004	001	Dec 08, 1998
TABLET; ORAL					
EPIVIR					
	GLAXOSMITHKLINE	150MG	N20564	001	Nov 17, 1995
	+	300MG	N20564	003	Jun 24, 2002
EPIVIR-HBV					
	+ GLAXOSMITHKLINE	100MG	N21003	001	Dec 08, 1998

LAMIVUDINE; ZIDOVUDINE

TABLET; ORAL					
COMBIVIR					
	+ GLAXOSMITHKLINE	150MG; 300MG	N20857	001	Sep 26, 1997

LAMOTRIGINE

TABLET; ORAL					
<u>LAMICTAL</u>					
<u>AB</u>	+ GLAXOSMITHKLINE	<u>25MG</u>	<u>N20241</u>	<u>005</u>	Dec 27, 1994
<u>AB</u>		<u>100MG</u>	<u>N20241</u>	<u>001</u>	Dec 27, 1994
<u>AB</u>		<u>150MG</u>	<u>N20241</u>	<u>002</u>	Dec 27, 1994
<u>AB</u>		<u>200MG</u>	<u>N20241</u>	<u>003</u>	Dec 27, 1994
<u>LAMOTRIGINE</u>					
<u>AB</u>	TEVA	<u>25MG</u>	<u>N76388</u>	<u>001</u>	Aug 30, 2006

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 220 (of 370)

LAMOTRIGINE

TABLET; ORAL

LAMOTRIGINE

<u>AB</u>	TEVA	<u>100MG</u>	<u>N76388</u>	<u>002</u>	Aug 30, 2006
<u>AB</u>		<u>150MG</u>	<u>N76388</u>	<u>003</u>	Aug 30, 2006
<u>AB</u>		<u>200MG</u>	<u>N76388</u>	<u>004</u>	Aug 30, 2006

TABLET, CHEWABLE; ORAL

LAMICTAL CD

<u>AB</u>	GLAXOSMITHKLINE	<u>5MG</u>	<u>N20764</u>	<u>001</u>	Aug 24, 1998
<u>AB</u>	+	<u>25MG</u>	<u>N20764</u>	<u>002</u>	Aug 24, 1998
		2MG	N20764	004	Sep 08, 2000

LAMOTRIGINE

<u>AB</u>	TEVA	<u>5MG</u>	<u>N76420</u>	<u>001</u>	Jun 21, 2006
<u>AB</u>		<u>25MG</u>	<u>N76420</u>	<u>002</u>	Jun 21, 2006

LANSOPRAZOLE

CAPSULE, DELAYED REL PELLETS; ORAL

PREVACID

	TAP PHARM	15MG	N20406	001	May 10, 1995
+		30MG	N20406	002	May 10, 1995

FOR SUSPENSION, DELAYED RELEASE; ORAL

PREVACID

	TAP PHARM	15MG/PACKET	N21281	001	May 03, 2001
+		30MG/PACKET	N21281	002	May 03, 2001

INJECTABLE; INTRAVENOUS

PREVACID IV

+	TAP PHARM	30MG/VIAL	N21566	001	May 27, 2004
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TABLET, DELAYED RELEASE, ORALLY DISINTEGRATING; ORAL

PREVACID

	TAP PHARM	15MG	N21428	001	Aug 30, 2002
+		30MG	N21428	002	Aug 30, 2002

LANSOPRAZOLE; NAPROXEN

CAPSULE, DELAYED REL PELLETS, TABLET; ORAL

PREVACID NAPRAPAC 500 (COPACKAGED)

+	TAP PHARM	15MG, N/A; N/A, 500MG	N21507	004	Nov 14, 2003
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LANTHANUM CARBONATE

TABLET, CHEWABLE; ORAL

FOSRENOL

	SHIRE	EQ 250MG BASE	N21468	001	Oct 26, 2004
		EQ 500MG BASE	N21468	002	Oct 26, 2004
		EQ 750MG BASE	N21468	003	Nov 23, 2005
+		EQ 1GM BASE	N21468	004	Nov 23, 2005

LATANOPROST

SOLUTION/DROPS; OPHTHALMIC

+	PHARMACIA AND UPJOHN	0.005%	N20597	001	Jun 05, 1996
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LEFLUNOMIDE

TABLET; ORAL

ARAVA

<u>AB</u>	SANOFI AVENTIS US	<u>10MG</u>	<u>N20905</u>	<u>001</u>	Sep 10, 1998
<u>AB</u>	+	<u>20MG</u>	<u>N20905</u>	<u>002</u>	Sep 10, 1998
	+	100MG	N20905	003	Sep 10, 1998

LEFLUNOMIDE

<u>AB</u>	APOTEX INC	<u>10MG</u>	<u>N77090</u>	<u>001</u>	Sep 13, 2005
<u>AB</u>		<u>20MG</u>	<u>N77090</u>	<u>002</u>	Sep 13, 2005

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 221 (of 370)

LEFLUNOMIDE

TABLET; ORAL

LEFLUNOMIDE

<u>AB</u>	BARR	<u>10MG</u>	<u>N77083</u>	<u>001</u>	Sep 13, 2005
<u>AB</u>		<u>20MG</u>	<u>N77083</u>	<u>002</u>	Sep 13, 2005
<u>AB</u>	KALI LABS	<u>10MG</u>	<u>N77086</u>	<u>001</u>	Sep 13, 2005
<u>AB</u>		<u>20MG</u>	<u>N77086</u>	<u>002</u>	Sep 13, 2005
<u>AB</u>	SANDOZ	<u>10MG</u>	<u>N77087</u>	<u>001</u>	Sep 13, 2005
<u>AB</u>		<u>20MG</u>	<u>N77087</u>	<u>002</u>	Sep 13, 2005
<u>AB</u>	TEVA PHARMS	<u>10MG</u>	<u>N77084</u>	<u>001</u>	Sep 13, 2005
<u>AB</u>		<u>20MG</u>	<u>N77084</u>	<u>002</u>	Sep 13, 2005

LENALIDOMIDE

CAPSULE; ORAL

REVLIMID

CELGENE

	5MG	N21880	001	Dec 27, 2005
	10MG	N21880	002	Dec 27, 2005
	15MG	N21880	003	Jun 29, 2006
+	25MG	N21880	004	Jun 29, 2006

LEPIRUDIN RECOMBINANT

INJECTABLE; INJECTION

REFLUDAN

+	BERLEX	50MG/VIAL	N20807	001	Mar 06, 1998
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LETROZOLE

TABLET; ORAL

FEMARA

+	NOVARTIS PHARMS	2.5MG	N20726	001	Jul 25, 1997
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LEUCOVORIN CALCIUM

INJECTABLE; INJECTION

LEUCOVORIN CALCIUM

<u>AP</u>	BEDFORD	<u>EQ 50MG BASE/VIAL</u>	<u>N89384</u>	<u>001</u>	Sep 14, 1987
<u>AP</u>		<u>EQ 100MG BASE/VIAL</u>	<u>N89717</u>	<u>001</u>	Mar 28, 1988
<u>AP</u>	+	<u>EQ 350MG BASE/VIAL</u>	<u>N08107</u>	<u>005</u>	Apr 05, 1989
<u>AP</u>	MAYNE PHARMA USA	<u>EQ 350MG BASE/VIAL</u>	<u>N40262</u>	<u>001</u>	Dec 15, 1999
<u>AP</u>	PHARMACHEMIE	<u>EQ 50MG BASE/VIAL</u>	<u>N89628</u>	<u>001</u>	Apr 17, 1997
<u>AP</u>	PHARMACHEMIE USA	<u>EQ 50MG BASE/VIAL</u>	<u>N81278</u>	<u>001</u>	Sep 28, 1993
<u>AP</u>	SICOR PHARMS	<u>EQ 100MG BASE/VIAL</u>	<u>N81277</u>	<u>001</u>	Sep 28, 1993
<u>AP</u>		<u>EQ 350MG BASE/VIAL</u>	<u>N40174</u>	<u>001</u>	Jun 12, 1997
<u>LEUCOVORIN CALCIUM PRESERVATIVE FREE</u>					
<u>AP</u>	ABRAXIS PHARM	<u>EQ 200MG BASE/VIAL</u>	<u>N40258</u>	<u>001</u>	Feb 26, 1999
+		EQ 500MG BASE/VIAL	N40286	001	Feb 26, 1999
<u>AP</u>	+	<u>EQ 10MG BASE/ML</u>	<u>N40347</u>	<u>001</u>	Apr 25, 2000
<u>AP</u>	+	<u>EQ 200MG BASE/VIAL</u>	<u>N40056</u>	<u>001</u>	May 23, 1995
<u>AP</u>		<u>EQ 350MG BASE/VIAL</u>	<u>N40335</u>	<u>001</u>	Apr 20, 2000
<u>AP</u>	LUITPOLD	<u>EQ 50MG BASE/VIAL</u>	<u>N40338</u>	<u>001</u>	Jan 31, 2001
<u>AP</u>	SICOR PHARMS	<u>EQ 10MG BASE/ML</u>	<u>N40332</u>	<u>001</u>	Jun 28, 1999

TABLET; ORAL

LEUCOVORIN CALCIUM

<u>AB</u>	BARR	<u>EQ 5MG BASE</u>	<u>N71198</u>	<u>001</u>	Sep 24, 1987
<u>AB</u>		<u>EQ 25MG BASE</u>	<u>N71199</u>	<u>001</u>	Sep 24, 1987
<u>AB</u>	PAR PHARM	<u>EQ 5MG BASE</u>	<u>N74544</u>	<u>001</u>	Aug 28, 1997
<u>AB</u>		<u>EQ 25MG BASE</u>	<u>N74544</u>	<u>002</u>	Aug 28, 1997
<u>AB</u>	PHARMACHEMIE	<u>EQ 5MG BASE</u>	<u>N73099</u>	<u>001</u>	Mar 28, 1997
<u>AB</u>		<u>EQ 25MG BASE</u>	<u>N73101</u>	<u>001</u>	Mar 28, 1997
<u>AB</u>	ROXANE	<u>EQ 5MG BASE</u>	<u>N72733</u>	<u>001</u>	Feb 22, 1993
<u>AB</u>		<u>EQ 10MG BASE</u>	<u>N72734</u>	<u>001</u>	Feb 22, 1993

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 222 (of 370)

LEUCOVORIN CALCIUM

TABLET; ORAL

LEUCOVORIN CALCIUM

<u>AB</u>	ROXANE	<u>EQ 15MG BASE</u>	<u>N72735</u>	<u>001</u>	Feb 22, 1993
<u>AB</u>	+	<u>EQ 25MG BASE</u>	<u>N72736</u>	<u>001</u>	Feb 22, 1993
<u>AB</u>	SANDOZ	<u>EQ 15MG BASE</u>	<u>N75327</u>	<u>001</u>	Mar 24, 1999
<u>AB</u>	XANODYNE PHARM	<u>EQ 10MG BASE</u>	<u>N71962</u>	<u>001</u>	Nov 19, 1987
<u>AB</u>		<u>EQ 15MG BASE</u>	<u>N71104</u>	<u>001</u>	Mar 04, 1987
<u>BX</u>		EQ 5MG BASE	N18459	001	Jan 30, 1986

LEUPROLIDE ACETATE

IMPLANT; IMPLANTATION

VIADUR

+	ALZA	EQ 65MG BASE	N21088	001	Mar 03, 2000
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INJECTABLE; INJECTION

LEUPROLIDE ACETATE

<u>AP</u>	BEDFORD LABS	<u>1MG/0.2ML</u>	<u>N74728</u>	<u>001</u>	Aug 04, 1998
<u>AP</u>	GENZYME	<u>1MG/0.2ML</u>	<u>N75721</u>	<u>001</u>	Nov 29, 2001
<u>AP</u>	SICOR PHARMS	<u>1MG/0.2ML</u>	<u>N75471</u>	<u>001</u>	Oct 25, 2000
	<u>LUPRON</u>				
<u>AP</u>	+	<u>1MG/0.2ML</u>	<u>N19010</u>	<u>001</u>	Apr 09, 1985
	LUPRON DEPOT				
+	TAP PHARM	3.75MG/VIAL	N20011	001	Oct 22, 1990
+		7.5MG/VIAL	N19732	001	Jan 26, 1989
+		22.5MG/VIAL	N20517	001	Dec 22, 1995
	LUPRON DEPOT-3				
+	TAP PHARM	11.25MG/VIAL	N20708	001	Mar 07, 1997
	LUPRON DEPOT-4				
+	TAP PHARM	30MG/VIAL	N20517	002	May 30, 1997
	LUPRON DEPOT-PED				
+	TAP PHARM	7.5MG/VIAL	N20263	002	Apr 16, 1993
+		11.25MG/VIAL	N20263	005	Jan 21, 1994
+		15MG/VIAL	N20263	006	Jan 21, 1994

INJECTABLE; SUBCUTANEOUS

ELIGARD

+	QLT USA	7.5MG/VIAL	N21343	001	Jan 23, 2002
+		22.5MG/VIAL	N21379	001	Jul 24, 2002
+		30MG/VIAL	N21488	001	Feb 13, 2003
+		45MG/VIAL	N21731	001	Dec 14, 2004

LEVALBUTEROL HYDROCHLORIDE

SOLUTION; INHALATION

XOPENEX

+	SEPRACOR	EQ 0.0103% BASE	N20837	003	Jan 30, 2002
+		EQ 0.021% BASE	N20837	001	Mar 25, 1999
+		EQ 0.042% BASE	N20837	002	Mar 25, 1999
+		EQ 0.25% BASE	N20837	004	Jul 18, 2003

LEVALBUTEROL TARTRATE

AEROSOL, METERED; INHALATION

XOPENEX HFA

+	SEPRACOR	EQ 0.045MG BASE/INH	N21730	001	Mar 11, 2005
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LEVETIRACETAM

INJECTABLE; IV (INFUSION)

+	UCB INC	500MG/5ML (100MG/ML)	N21872	001	Jul 31, 2006
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27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 223 (of 370)

LEVETIRACETAM

SOLUTION; ORAL

KEPPRA

+	UCB INC	100MG/ML	N21505	001	Jul 15, 2003
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TABLET; ORAL

KEPPRA

	UCB INC	250MG	N21035	001	Nov 30, 1999
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		500MG	N21035	002	Nov 30, 1999
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		750MG	N21035	003	Nov 30, 1999
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+		1GM	N21035	004	Jan 06, 2006
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LEVOBUNOLOL HYDROCHLORIDE

SOLUTION/DROPS; OPHTHALMIC

AKBETA

<u>AT</u>	AKORN	<u>0.25%</u>	<u>N74779</u>	<u>001</u>	Oct 29, 1996
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<u>AT</u>		<u>0.5%</u>	<u>N74780</u>	<u>001</u>	Oct 29, 1996
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BETAGAN

<u>AT</u>	+	ALLERGAN	<u>0.25%</u>	<u>N19814</u>	<u>001</u>	Jun 28, 1989
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<u>AT</u>	+		<u>0.5%</u>	<u>N19219</u>	<u>002</u>	Dec 19, 1985
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LEVOBUNOLOL HYDROCHLORIDE

<u>AT</u>	BAUSCH AND LOMB	<u>0.25%</u>	<u>N74307</u>	<u>001</u>	Mar 04, 1994
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<u>AT</u>		<u>0.5%</u>	<u>N74326</u>	<u>001</u>	Mar 04, 1994
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<u>AT</u>	FALCON PHARMS	<u>0.25%</u>	<u>N74851</u>	<u>001</u>	Oct 28, 1996
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<u>AT</u>		<u>0.5%</u>	<u>N74850</u>	<u>001</u>	Oct 28, 1996
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<u>AT</u>	NOVEX	<u>0.25%</u>	<u>N75473</u>	<u>001</u>	Aug 03, 2000
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<u>AT</u>		<u>0.5%</u>	<u>N75475</u>	<u>001</u>	Aug 03, 2000
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LEVOCARNITINE

INJECTABLE; INJECTION

CARNITOR

<u>AP</u>	+	SIGMA TAU	<u>200MG/ML</u>	<u>N20182</u>	<u>001</u>	Dec 16, 1992
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LEVOCARNITINE

<u>AP</u>		BEDFORD	<u>200MG/ML</u>	<u>N75567</u>	<u>001</u>	Mar 29, 2001
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<u>AP</u>		LUITPOLD	<u>200MG/ML</u>	<u>N75861</u>	<u>001</u>	Jun 22, 2001
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<u>AP</u>		SICOR PHARMS	<u>200MG/ML</u>	<u>N75881</u>	<u>001</u>	Mar 29, 2001
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SOLUTION; ORAL

CARNITOR

<u>AA</u>	+	SIGMA TAU	<u>1GM/10ML</u>	<u>N19257</u>	<u>001</u>	Apr 10, 1986
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LEVOCARNITINE

<u>AA</u>		LYNE	<u>1GM/10ML</u>	<u>N76851</u>	<u>001</u>	Aug 10, 2004
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TABLET; ORAL

CARNITOR

<u>AB</u>	+	SIGMA TAU	<u>330MG</u>	<u>N18948</u>	<u>001</u>	Dec 27, 1985
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LEVOCARNITINE

<u>AB</u>		COREPHARMA	<u>330MG</u>	<u>N76858</u>	<u>001</u>	Sep 20, 2004
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LEVOFLOXACIN

INJECTABLE; INJECTION

LEVAQUIN

+	ORTHO MCNEIL PHARM	EQ 500MG /20ML(25MG/ML)	N20635	001	Dec 20, 1996
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+		EQ 750MG /30ML(25MG/ML)	N20635	004	Dec 20, 1996
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LEVAQUIN IN DEXTROSE 5% IN PLASTIC CONTAINER

+	ORTHO MCNEIL PHARM	EQ 250MG /50ML(5MG/ML)	N20635	002	Dec 20, 1996
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+		EQ 500MG /100ML(5MG/ML)	N20635	003	Dec 20, 1996
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+		EQ 750MG /150ML(5MG/ML)	N20635	005	Dec 20, 1996
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SOLUTION; ORAL

LEVAQUIN

+	ORTHO MCNEIL PHARM	250MG/10ML	N21721	001	Oct 21, 2004
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PRESCRIPTION DRUG PRODUCT LIST

3 - 224 (of 370)

LEVOFLOXACIN

SOLUTION/DROPS; OPHTHALMIC

+ SANTEN	1.5%	N21571	001	Mar 01, 2004
QUIXIN				
+ SANTEN	0.5%	N21199	001	Aug 18, 2000

TABLET; ORAL

LEVAQUIN

ORTHO MCNEIL PHARM	250MG	N20634	001	Dec 20, 1996
	500MG	N20634	002	Dec 20, 1996
+	750MG	N20634	003	Sep 08, 2000

LEVONORDEFIN; MEPIVACAINE HYDROCHLORIDE

INJECTABLE; INJECTION

CARBOCAINE W/ NEO-COBEFRIN

<u>AP</u>	+ EASTMAN KODAK	0.05MG/ML;2%	N12125	002	
	<u>ISOCAINE HYDROCHLORIDE W/ LEVONORDEFIN</u>				
<u>AP</u>	NOVOCOL	0.05MG/ML;2%	N84697	001	
	<u>MEPIVACAINE HYDROCHLORIDE W/ LEVONORDEFIN</u>				
<u>AP</u>	GRAHAM CHEM	0.05MG/ML;2%	N84850	002	Oct 21, 1983
	<u>POLOCAINE W/ LEVONORDEFIN</u>				
<u>AP</u>	DENTSPLY PHARM	0.05MG/ML;2%	N89517	001	Apr 14, 1988
	<u>SCANDONEST L</u>				
<u>AP</u>	DEPROCO	0.05MG/ML;2%	N88388	001	Oct 10, 1984

LEVONORGESTREL

IMPLANT; IMPLANTATION

NORPLANT II

BX	+ POPULATION COUNCIL	75MG/IMPLANT	N20544	001	Nov 01, 1996
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INTRAUTERINE DEVICE; INTRAUTERINE

MIRENA

+ BERLEX	52MG	N21225	001	Dec 06, 2000
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TABLET; ORAL

PLAN B

+ DURAMED	0.75MG	N21045	002	Aug 24, 2006
+	0.75MG	N21045	001	Jul 28, 1999

LEVORPHANOL TARTRATE

TABLET; ORAL

LEVO-DROMORAN

<u>AB</u>	+ VALEANT PHARM INTL	2MG	N08720	001	Dec 19, 1991
	<u>LEVORPHANOL TARTRATE</u>				
<u>AB</u>	ROXANE	2MG	N74278	001	Mar 31, 2000

LEVOTHYROXINE SODIUM**

Refer to Preface Section 1.8 Levothyroxine Sodium for amplifying information

CAPSULE; ORAL

TIROSINT

INST BIOCHIMIQUE	0.025MG	N21924	002	Oct 13, 2006
	0.05MG	N21924	003	Oct 13, 2006
	0.075MG	N21924	004	Oct 13, 2006
	0.1MG	N21924	005	Oct 13, 2006
	0.125MG	N21924	006	Oct 13, 2006
+	0.15MG	N21924	007	Oct 13, 2006

TABLET; ORAL

LEVOLET

BX	VINTAGE	0.025MG	N21137	001	Jun 06, 2003
BX		0.05MG	N21137	002	Jun 06, 2003
BX		0.075MG	N21137	003	Jun 06, 2003

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 225 (of 370)

LEVOTHYROXINE SODIUM**

Refer to Preface Section 1.8 Levothyroxine Sodium for amplifying information

TABLET; ORAL

LEVOLET

BX	VINTAGE	0.088MG	N21137	004	Jun 06, 2003
BX		0.1MG	N21137	005	Jun 06, 2003
BX		0.112MG	N21137	006	Jun 06, 2003
BX		0.125MG	N21137	007	Jun 06, 2003
BX		0.137MG	N21137	008	Jun 06, 2003
BX		0.15MG	N21137	009	Jun 06, 2003
BX		0.175MG	N21137	010	Jun 06, 2003
BX		0.2MG	N21137	011	Jun 06, 2003
BX		0.3MG	N21137	012	Jun 06, 2003

LEVO-T

-->	ALARA PHARM	--> AB2, AB3 0.025MG	<u>N21342</u>	<u>001</u>	Mar 01, 2002
-->		--> AB2, AB3 0.05MG	<u>N21342</u>	<u>002</u>	Mar 01, 2002
-->		--> AB2, AB3 0.075MG	<u>N21342</u>	<u>003</u>	Mar 01, 2002
-->		--> AB2, AB3 0.088MG	<u>N21342</u>	<u>004</u>	Mar 01, 2002
-->		--> AB2, AB3 0.1MG	<u>N21342</u>	<u>005</u>	Mar 01, 2002
-->		--> AB2, AB3 0.112MG	<u>N21342</u>	<u>006</u>	Mar 01, 2002
-->		--> AB2, AB3 0.125MG	<u>N21342</u>	<u>007</u>	Mar 01, 2002
-->		--> AB2, AB3 0.137MG	<u>N21342</u>	<u>012</u>	Dec 08, 2003
-->		--> AB2, AB3 0.15MG	<u>N21342</u>	<u>008</u>	Mar 01, 2002
-->		--> AB2, AB3 0.175MG	<u>N21342</u>	<u>009</u>	Mar 01, 2002
-->		--> AB2, AB3 0.2MG	<u>N21342</u>	<u>010</u>	Mar 01, 2002
--> +		--> AB2, AB3 0.3MG	<u>N21342</u>	<u>011</u>	Mar 01, 2002

LEVOTHROID

<u>AB4</u>	LLOYD	<u>0.025MG</u>	<u>N21116</u>	<u>001</u>	Oct 24, 2002
<u>AB4</u>		<u>0.05MG</u>	<u>N21116</u>	<u>002</u>	Oct 24, 2002
<u>AB4</u>		<u>0.075MG</u>	<u>N21116</u>	<u>003</u>	Oct 24, 2002
<u>AB4</u>		<u>0.088MG</u>	<u>N21116</u>	<u>010</u>	Oct 24, 2002
<u>AB4</u>		<u>0.1MG</u>	<u>N21116</u>	<u>004</u>	Oct 24, 2002
<u>AB4</u>		<u>0.112MG</u>	<u>N21116</u>	<u>011</u>	Oct 24, 2002
<u>AB4</u>		<u>0.125MG</u>	<u>N21116</u>	<u>005</u>	Oct 24, 2002
<u>AB4</u>		<u>0.137MG</u>	<u>N21116</u>	<u>012</u>	Dec 07, 2004
<u>AB4</u>		<u>0.15MG</u>	<u>N21116</u>	<u>006</u>	Oct 24, 2002
<u>AB4</u>		<u>0.175MG</u>	<u>N21116</u>	<u>007</u>	Oct 24, 2002
<u>AB4</u>		<u>0.2MG</u>	<u>N21116</u>	<u>008</u>	Oct 24, 2002
<u>AB4</u> +		<u>0.3MG</u>	<u>N21116</u>	<u>009</u>	Oct 24, 2002

LEVOTHYROXINE SODIUM

-->	GENPHARM	--> AB2, AB3 0.025MG	<u>N76752</u>	<u>001</u>	Jun 16, 2005
-->		--> AB2, AB3 0.05MG	<u>N76752</u>	<u>002</u>	Jun 16, 2005
-->		--> AB2, AB3 0.075MG	<u>N76752</u>	<u>003</u>	Jun 16, 2005
-->		--> AB2, AB3 0.088MG	<u>N76752</u>	<u>004</u>	Jun 16, 2005
-->		--> AB2, AB3 0.1MG	<u>N76752</u>	<u>005</u>	Jun 16, 2005
-->		--> AB2, AB3 0.112MG	<u>N76752</u>	<u>006</u>	Jun 16, 2005
-->		--> AB2, AB3 0.125MG	<u>N76752</u>	<u>007</u>	Jun 16, 2005
-->		--> AB2, AB3 0.15MG	<u>N76752</u>	<u>008</u>	Jun 16, 2005
-->		--> AB2, AB3 0.175MG	<u>N76752</u>	<u>009</u>	Jun 16, 2005
-->		--> AB2, AB3 0.2MG	<u>N76752</u>	<u>010</u>	Jun 16, 2005
-->		--> AB2, AB3 0.3MG	<u>N76752</u>	<u>011</u>	Jun 16, 2005
-->	MYLAN	--> AB1, AB2, AB3, AB4 0.025MG	<u>N76187</u>	<u>001</u>	Jun 05, 2002
-->		--> AB1, AB2, AB3, AB4 0.05MG	<u>N76187</u>	<u>002</u>	Jun 05, 2002
-->		--> AB1, AB2, AB3, AB4 0.075MG	<u>N76187</u>	<u>003</u>	Jun 05, 2002
-->		--> AB1, AB2, AB3, AB4 0.088MG	<u>N76187</u>	<u>004</u>	Jun 05, 2002
-->		--> AB1, AB2, AB3, AB4 0.1MG	<u>N76187</u>	<u>005</u>	Jun 05, 2002
-->		--> AB1, AB2, AB3, AB4 0.112MG	<u>N76187</u>	<u>006</u>	Jun 05, 2002
-->		--> AB1, AB2, AB3, AB4 0.125MG	<u>N76187</u>	<u>007</u>	Jun 05, 2002
-->		--> AB1, AB2, AB3, AB4 0.137MG	<u>N76187</u>	<u>012</u>	Dec 13, 2006
-->		--> AB1, AB2, AB3, AB4 0.15MG	<u>N76187</u>	<u>008</u>	Jun 05, 2002

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 226 (of 370)

LEVOTHYROXINE SODIUM**

Refer to Preface Section 1.8 Levothyroxine Sodium for amplifying information

TABLET; ORAL

<u>LEVOTHYROXINE SODIUM</u>					
-->	MYLAN	--> AB1,AB2,AB3,AB4 0.175MG	<u>N76187</u>	<u>009</u>	Jun 05, 2002
-->		--> AB1,AB2,AB3,AB4 0.2MG	<u>N76187</u>	<u>010</u>	Jun 05, 2002
-->		--> AB1,AB2,AB3,AB4 0.3MG	<u>N76187</u>	<u>011</u>	Jun 05, 2002
<u>LEVOXYL</u>					
-->	JONES PHARMA	--> AB1,AB3 0.025MG	<u>N21301</u>	<u>001</u>	May 25, 2001
-->		--> AB1,AB3 0.05MG	<u>N21301</u>	<u>002</u>	May 25, 2001
-->		--> AB1,AB3 0.075MG	<u>N21301</u>	<u>003</u>	May 25, 2001
-->		--> AB1,AB3 0.088MG	<u>N21301</u>	<u>004</u>	May 25, 2001
-->		--> AB1,AB3 0.1MG	<u>N21301</u>	<u>005</u>	May 25, 2001
-->		--> AB1,AB3 0.112MG	<u>N21301</u>	<u>006</u>	May 25, 2001
-->		--> AB1,AB3 0.125MG	<u>N21301</u>	<u>007</u>	May 25, 2001
-->		--> AB1,AB3 0.137MG	<u>N21301</u>	<u>008</u>	May 25, 2001
-->		--> AB1,AB3 0.15MG	<u>N21301</u>	<u>009</u>	May 25, 2001
-->		--> AB1,AB3 0.175MG	<u>N21301</u>	<u>010</u>	May 25, 2001
-->		--> AB1,AB3 0.2MG	<u>N21301</u>	<u>011</u>	May 25, 2001
--> +		--> AB1,AB3 0.3MG	<u>N21301</u>	<u>012</u>	May 25, 2001
NOVOTHYROX					
BX	GENPHARM	0.025MG	N21292	001	May 31, 2002
BX		0.05MG	N21292	002	May 31, 2002
BX		0.075MG	N21292	003	May 31, 2002
BX		0.088MG	N21292	004	May 31, 2002
BX		0.1MG	N21292	005	May 31, 2002
BX		0.112MG	N21292	006	May 31, 2002
BX		0.125MG	N21292	007	May 31, 2002
BX		0.137MG	N21292	008	May 31, 2002
BX		0.15MG	N21292	009	May 31, 2002
BX		0.175MG	N21292	010	May 31, 2002
BX		0.2MG	N21292	011	May 31, 2002
BX +		0.3MG	N21292	012	May 31, 2002
<u>SYNTHROID</u>					
-->	ABBOTT	--> AB1,AB2 0.025MG	<u>N21402</u>	<u>001</u>	Jul 24, 2002
-->		--> AB1,AB2 0.05MG	<u>N21402</u>	<u>002</u>	Jul 24, 2002
-->		--> AB1,AB2 0.075MG	<u>N21402</u>	<u>003</u>	Jul 24, 2002
-->		--> AB1,AB2 0.088MG	<u>N21402</u>	<u>004</u>	Jul 24, 2002
-->		--> AB1,AB2 0.1MG	<u>N21402</u>	<u>005</u>	Jul 24, 2002
-->		--> AB1,AB2 0.112MG	<u>N21402</u>	<u>006</u>	Jul 24, 2002
-->		--> AB1,AB2 0.125MG	<u>N21402</u>	<u>007</u>	Jul 24, 2002
-->		--> AB1,AB2 0.137MG	<u>N21402</u>	<u>008</u>	Jul 24, 2002
-->		--> AB1,AB2 0.15MG	<u>N21402</u>	<u>009</u>	Jul 24, 2002
-->		--> AB1,AB2 0.175MG	<u>N21402</u>	<u>010</u>	Jul 24, 2002
-->		--> AB1,AB2 0.2MG	<u>N21402</u>	<u>012</u>	Jul 24, 2002
--> +		--> AB1,AB2 0.3MG	<u>N21402</u>	<u>011</u>	Jul 24, 2002
<u>UNITHROID</u>					
-->	STEVENS J	--> AB1,AB2,AB3 0.025MG	<u>N21210</u>	<u>001</u>	Aug 21, 2000
-->		--> AB1,AB2,AB3 0.05MG	<u>N21210</u>	<u>002</u>	Aug 21, 2000
-->		--> AB1,AB2,AB3 0.075MG	<u>N21210</u>	<u>003</u>	Aug 21, 2000
-->		--> AB1,AB2,AB3 0.088MG	<u>N21210</u>	<u>004</u>	Aug 21, 2000
-->		--> AB1,AB2,AB3 0.1MG	<u>N21210</u>	<u>005</u>	Aug 21, 2000
-->		--> AB1,AB2,AB3 0.112MG	<u>N21210</u>	<u>006</u>	Aug 21, 2000
-->		--> AB1,AB2,AB3 0.125MG	<u>N21210</u>	<u>007</u>	Aug 21, 2000
-->		--> AB1,AB2,AB3 0.15MG	<u>N21210</u>	<u>008</u>	Aug 21, 2000
-->		--> AB1,AB2,AB3 0.175MG	<u>N21210</u>	<u>009</u>	Aug 21, 2000
-->		--> AB1,AB2,AB3 0.2MG	<u>N21210</u>	<u>010</u>	Aug 21, 2000
--> +		--> AB1,AB2,AB3 0.3MG	<u>N21210</u>	<u>011</u>	Aug 21, 2000

PRESCRIPTION DRUG PRODUCT LIST

3 - 227 (of 370)

LIDOCAINE

FILM, EXTENDED RELEASE; BUCCAL					
LIDOCAINE					
	+ NOVEN	46.1MG/PATCH	N20575	002	May 21, 1996
OINTMENT; TOPICAL					
<u>LIDOCAINE</u>					
<u>AT</u>	+ FOUGERA	5%	<u>N80198</u>	<u>001</u>	
<u>AT</u>	GRAHAM CHEM	5%	<u>N80210</u>	<u>001</u>	
<u>AT</u>	TARO	5%	<u>N86724</u>	<u>001</u>	
PATCH; TOPICAL					
LIDODERM					
	+ TEIKOKU PHARMA USA	5%	N20612	001	Mar 19, 1999

LIDOCAINE HYDROCHLORIDE

INJECTABLE; INJECTION					
<u>LIDOCAINE HYDROCHLORIDE</u>					
<u>AP</u>	GRAHAM CHEM	2%	<u>N80504</u>	<u>001</u>	
<u>AP</u>	HOSPIRA	0.5%	<u>N88328</u>	<u>001</u>	May 17, 1984
<u>AP</u>		1%	<u>N40013</u>	<u>001</u>	Jun 23, 1995
<u>AP</u>		1%	<u>N83158</u>	<u>001</u>	
<u>AP</u>		1%	<u>N88329</u>	<u>001</u>	May 17, 1984
<u>AP</u>		2%	<u>N40078</u>	<u>001</u>	Jun 23, 1995
<u>AP</u>		2%	<u>N83158</u>	<u>002</u>	
<u>AP</u>		2%	<u>N88294</u>	<u>001</u>	May 17, 1984
<u>AP</u>		2%	<u>N88331</u>	<u>001</u>	May 17, 1984
<u>AP</u>		20%	<u>N83158</u>	<u>003</u>	
<u>AP</u>	INTL MEDICATION	1%	<u>N83173</u>	<u>001</u>	
<u>AP</u>		2%	<u>N83173</u>	<u>002</u>	
<u>AP</u>	LUITPOLD	1%	<u>N80850</u>	<u>001</u>	
<u>AP</u>		2%	<u>N83198</u>	<u>001</u>	
<u>LIDOCAINE HYDROCHLORIDE 0.2% AND DEXTROSE 5% IN PLASTIC CONTAINER</u>					
<u>AP</u>	B BRAUN	200MG/100ML	<u>N19830</u>	<u>002</u>	Apr 08, 1992
<u>AP</u>	BAXTER HLTHCARE	200MG/100ML	<u>N18461</u>	<u>002</u>	
<u>LIDOCAINE HYDROCHLORIDE 0.2% IN DEXTROSE 5%</u>					
<u>AP</u>	HOSPIRA	200MG/100ML	<u>N83158</u>	<u>005</u>	
<u>LIDOCAINE HYDROCHLORIDE 0.2% IN DEXTROSE 5% IN PLASTIC CONTAINER</u>					
<u>AP</u>	HOSPIRA	200MG/100ML	<u>N18388</u>	<u>001</u>	
<u>LIDOCAINE HYDROCHLORIDE 0.4% AND DEXTROSE 5% IN PLASTIC CONTAINER</u>					
<u>AP</u>	B BRAUN	400MG/100ML	<u>N19830</u>	<u>003</u>	Apr 08, 1992
<u>AP</u>	BAXTER HLTHCARE	400MG/100ML	<u>N18461</u>	<u>003</u>	
<u>LIDOCAINE HYDROCHLORIDE 0.4% IN DEXTROSE 5%</u>					
<u>AP</u>	HOSPIRA	400MG/100ML	<u>N83158</u>	<u>006</u>	
<u>LIDOCAINE HYDROCHLORIDE 0.4% IN DEXTROSE 5% IN PLASTIC CONTAINER</u>					
<u>AP</u>	HOSPIRA	400MG/100ML	<u>N18388</u>	<u>002</u>	
<u>LIDOCAINE HYDROCHLORIDE 0.8% AND DEXTROSE 5% IN PLASTIC CONTAINER</u>					
<u>AP</u>	B BRAUN	800MG/100ML	<u>N19830</u>	<u>004</u>	Apr 08, 1992
<u>AP</u>	BAXTER HLTHCARE	800MG/100ML	<u>N18461</u>	<u>004</u>	Feb 22, 1982
<u>LIDOCAINE HYDROCHLORIDE 0.8% IN DEXTROSE 5% IN PLASTIC CONTAINER</u>					
<u>AP</u>	HOSPIRA	800MG/100ML	<u>N18388</u>	<u>003</u>	Nov 05, 1982
<u>LIDOCAINE HYDROCHLORIDE IN PLASTIC CONTAINER</u>					
<u>AP</u>	ABRAXIS PHARM	1%	<u>N88586</u>	<u>001</u>	Jul 24, 1985
<u>AP</u>	HOSPIRA	0.5%	<u>N88325</u>	<u>001</u>	Jul 31, 1984
<u>AP</u>		1%	<u>N88299</u>	<u>001</u>	Jul 31, 1984
<u>AP</u>		1.5%	<u>N88326</u>	<u>001</u>	Jul 31, 1984
<u>AP</u>		2%	<u>N88327</u>	<u>001</u>	Jul 31, 1984
<u>AP</u>		10%	<u>N88367</u>	<u>001</u>	Jul 31, 1984
<u>AP</u>		20%	<u>N88368</u>	<u>001</u>	Jul 31, 1984
<u>LIDOCAINE HYDROCHLORIDE PRESERVATIVE FREE</u>					
<u>AP</u>	ABRAXIS PHARM	1%	<u>N80404</u>	<u>002</u>	
<u>AP</u>		2%	<u>N17584</u>	<u>001</u>	

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 228 (of 370)

LIDOCAINE HYDROCHLORIDE

INJECTABLE; INJECTION

LIDOCAINE HYDROCHLORIDE PRESERVATIVE FREE

<u>AP</u>	ABRAXIS PHARM	<u>2%</u>	<u>N80404</u>	<u>003</u>	
<u>AP</u>		<u>4%</u>	<u>N17584</u>	<u>002</u>	
<u>AP</u>	HOSPIRA	<u>1%</u>	<u>N80408</u>	<u>001</u>	
<u>AP</u>		<u>1.5%</u>	<u>N80408</u>	<u>002</u>	
<u>AP</u>		<u>4%</u>	<u>N88295</u>	<u>001</u>	May 17, 1984
<u>AP</u>	INTL MEDICATION	<u>20%</u>	<u>N17702</u>	<u>001</u>	

LIDOCAINE HYDROCHLORIDE PRESERVATIVE FREE IN PLASTIC CONTAINER

<u>AP</u>	HOSPIRA	<u>1%</u>	<u>N40302</u>	<u>001</u>	Sep 28, 1998
<u>AP</u>		<u>2%</u>	<u>N40302</u>	<u>002</u>	Sep 28, 1998

LIDOPEN

<u>AP</u>	MERIDIAN MEDCL TECHN	<u>10%</u>	<u>N17549</u>	<u>001</u>	
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XYLOCAINE

<u>AP</u>	+ ABRAXIS BIOSCIENCE	<u>0.5%</u>	<u>N06488</u>	<u>008</u>	
<u>AP</u>	+	<u>1%</u>	<u>N06488</u>	<u>007</u>	
<u>AP</u>	+	<u>1.5%</u>	<u>N06488</u>	<u>010</u>	
<u>AP</u>	+ DENTSPLY PHARM	<u>2%</u>	<u>N21380</u>	<u>001</u>	

XYLOCAINE 4% PRESERVATIVE FREE

<u>AP</u>	+ ABRAXIS BIOSCIENCE	<u>4%</u>	<u>N10417</u>	<u>001</u>	
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XYLOCAINE PRESERVATIVE FREE

<u>AP</u>	ABRAXIS BIOSCIENCE	<u>1%</u>	<u>N16801</u>	<u>005</u>	Jan 19, 1988
<u>AP</u>	+	<u>2%</u>	<u>N16801</u>	<u>001</u>	
<u>AP</u>		<u>4%</u>	<u>N16801</u>	<u>002</u>	
<u>AP</u>	+	<u>10%</u>	<u>N16801</u>	<u>003</u>	
<u>AP</u>	+	<u>20%</u>	<u>N16801</u>	<u>004</u>	

INJECTABLE; SPINAL

	LIDOCAINE HYDROCHLORIDE	5% AND DEXTROSE 7.5%			
	+ HOSPIRA	5%	N83914	001	

JELLY; TOPICAL

ANESTACON

<u>AT</u>	+ POLYMEDICA	<u>2%</u>	<u>N80429</u>	<u>001</u>	
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LIDOCAINE HYDROCHLORIDE

<u>AT</u>	AKORN	<u>2%</u>	<u>N40433</u>	<u>001</u>	Feb 12, 2003
<u>AT</u>	INTL MEDICATION	<u>2%</u>	<u>N86283</u>	<u>001</u>	
<u>AT</u>	TEVA PHARMS	<u>2%</u>	<u>N81318</u>	<u>001</u>	Apr 29, 1993

XYLOCAINE

<u>AT</u>	+ ABRAXIS BIOSCIENCE	<u>2%</u>	<u>N08816</u>	<u>001</u>	
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SOLUTION; ORAL

LIDOCAINE HYDROCHLORIDE

<u>AT</u>	HI TECH PHARMA	<u>2%</u>	<u>N40014</u>	<u>001</u>	Jul 10, 1995
<u>AT</u>	MORTON GROVE	<u>2%</u>	<u>N87872</u>	<u>001</u>	Nov 18, 1982

LIDOCAINE HYDROCHLORIDE VISCOUS

<u>AT</u>	ACTAVIS MID ATLANTIC	<u>2%</u>	<u>N86578</u>	<u>001</u>	
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LIDOCAINE VISCOUS

<u>AT</u>	ROXANE	<u>2%</u>	<u>N88802</u>	<u>001</u>	Apr 26, 1985
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XYLOCAINE VISCOUS

<u>AT</u>	+ ABRAXIS BIOSCIENCE	<u>2%</u>	<u>N09470</u>	<u>001</u>	
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SOLUTION; TOPICAL

LARYNG-O-JET KIT

<u>AT</u>	INTL MEDICATION	<u>4%</u>	<u>N86364</u>	<u>001</u>	
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LIDOCAINE HYDROCHLORIDE

<u>AT</u>	MORTON GROVE	<u>4%</u>	<u>N87881</u>	<u>001</u>	Nov 18, 1982
<u>AT</u>	ROXANE	<u>4%</u>	<u>N88803</u>	<u>001</u>	Apr 03, 1985

LTA II KIT

<u>AT</u>	HOSPIRA	<u>4%</u>	<u>N80409</u>	<u>001</u>	
<u>AT</u>		<u>4%</u>	<u>N88542</u>	<u>001</u>	Jul 31, 1984

PEDIATRIC LTA KIT

<u>AT</u>	HOSPIRA	<u>2%</u>	<u>N85995</u>	<u>001</u>	
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27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 229 (of 370)

LIDOCAINE HYDROCHLORIDE

SOLUTION; TOPICAL

XYLOCAINE 4% PRESERVATIVE FREE

<u>AT</u>	+	ABRAXIS BIOSCIENCE	<u>4%</u>	<u>N10417</u>	<u>002</u>	
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LIDOCAINE HYDROCHLORIDE; OXYTETRACYCLINE

INJECTABLE; INJECTION

TERRAMYCIN

+	PFIZER	2%;50MG/ML	N60567	001	
+		2%;125MG/ML	N60567	002	

LIDOCAINE; PRILOCAINE

CREAM; TOPICAL

EMLA

<u>AB</u>	+	ABRAXIS BIOSCIENCE	<u>2.5%;2.5%</u>	<u>N19941</u>	<u>001</u>	Dec 30, 1992
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LIDOCAINE AND PRILOCAINE

<u>AB</u>		ALTANA	<u>2.5%;2.5%</u>	<u>N76453</u>	<u>001</u>	Aug 18, 2003
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<u>AB</u>		HI TECH PHARMA	<u>2.5%;2.5%</u>	<u>N76290</u>	<u>001</u>	Sep 25, 2003
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<u>AB</u>		QLT USA	<u>2.5%;2.5%</u>	<u>N76320</u>	<u>001</u>	Aug 27, 2003
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GEL; PERIODONTAL

ORAQIX

+	DENTSPLY PHARM	2.5%;2.5%	N21451	001	Dec 19, 2003
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LIDOCAINE; TETRACAINE

CREAM; TOPICAL

LIDOCAINE AND TETRACAINE

+	ZARS	7%;7%	N21717	001	Jun 29, 2006
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PATCH; TOPICAL

SYNERA

+	ENDO PHARMS	70MG;70MG	N21623	001	Jun 23, 2005
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LINCOMYCIN HYDROCHLORIDE

INJECTABLE; INJECTION

LINCOCIN

+	PHARMACIA AND UPJOHN	EQ 300MG BASE/ML	N50317	001	
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LINDANE

LOTION; TOPICAL

LINDANE

<u>AT</u>	+	ACTAVIS MID ATLANTIC	<u>1%</u>	<u>N87313</u>	<u>001</u>	
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<u>AT</u>		MORTON GROVE	<u>1%</u>	<u>N88190</u>	<u>001</u>	Aug 16, 1984
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SHAMPOO; TOPICAL

LINDANE

<u>AT</u>	+	ACTAVIS MID ATLANTIC	<u>1%</u>	<u>N87266</u>	<u>001</u>	
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<u>AT</u>		MORTON GROVE	<u>1%</u>	<u>N88191</u>	<u>001</u>	Sep 18, 1984
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LINEZOLID

FOR SUSPENSION; ORAL

ZYVOX

+	PHARMACIA AND UPJOHN	100MG/5ML	N21132	001	Apr 18, 2000
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INJECTABLE; INJECTION

ZYVOX

+	PHARMACIA AND UPJOHN	200MG/100ML	N21131	001	Apr 18, 2000
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TABLET; ORAL

ZYVOX

+	PHARMACIA AND UPJOHN	600MG	N21130	002	Apr 18, 2000
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27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 230 (of 370)

LIOTHYRONINE SODIUM

INJECTABLE; INJECTION

LIOTHYRONINE SODIUM

<u>AP</u>	X GEN PHARMS	<u>EQ 0.01MG BASE/ML</u>	<u>N76923</u>	<u>001</u>	Aug 17, 2005
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TRIOSTAT

<u>AP</u>	+ JONES PHARMA	<u>EQ 0.01MG BASE/ML</u>	<u>N20105</u>	<u>001</u>	Dec 31, 1991
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TABLET; ORAL

CYTOMEL

KING PHARMS

EQ 0.005MG BASE

N10379 001

EQ 0.025MG BASE

N10379 002

+

EQ 0.05MG BASE

N10379 003

LIOTRIX (T4;T3)

TABLET; ORAL

THYROLAR-0.25

FOREST LABS

0.0125MG;0.0031MG

N16807 001

THYROLAR-0.5

FOREST LABS

0.025MG;0.0063MG

N16807 005

THYROLAR-1

FOREST LABS

0.05MG;0.0125MG

N16807 004

THYROLAR-2

FOREST LABS

0.1MG;0.025MG

N16807 002

THYROLAR-3

+ FOREST LABS

0.15MG;0.0375MG

N16807 003

LISINOPRIL

TABLET; ORAL

LISINOPRIL

<u>AB</u>	ACTAVIS ELIZABETH	<u>2.5MG</u>	<u>N76180</u>	<u>001</u>	Jul 01, 2002
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<u>AB</u>		<u>5MG</u>	<u>N76180</u>	<u>002</u>	Jul 01, 2002
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<u>AB</u>		<u>10MG</u>	<u>N76180</u>	<u>003</u>	Jul 01, 2002
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<u>AB</u>		<u>20MG</u>	<u>N76164</u>	<u>001</u>	Jul 01, 2002
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<u>AB</u>		<u>30MG</u>	<u>N76164</u>	<u>002</u>	Jul 01, 2002
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<u>AB</u>		<u>40MG</u>	<u>N76164</u>	<u>003</u>	Jul 01, 2002
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<u>AB</u>	APOTEX INC	<u>2.5MG</u>	<u>N76102</u>	<u>001</u>	Sep 30, 2002
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<u>AB</u>		<u>5MG</u>	<u>N76102</u>	<u>002</u>	Sep 30, 2002
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<u>AB</u>		<u>10MG</u>	<u>N76102</u>	<u>003</u>	Sep 30, 2002
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<u>AB</u>		<u>20MG</u>	<u>N76102</u>	<u>004</u>	Sep 30, 2002
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<u>AB</u>		<u>30MG</u>	<u>N76102</u>	<u>005</u>	Sep 30, 2002
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<u>AB</u>		<u>40MG</u>	<u>N76102</u>	<u>006</u>	Sep 30, 2002
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<u>AB</u>	AUROBINDO	<u>2.5MG</u>	<u>N77622</u>	<u>001</u>	Feb 22, 2006
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<u>AB</u>		<u>5MG</u>	<u>N77622</u>	<u>002</u>	Feb 22, 2006
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<u>AB</u>		<u>10MG</u>	<u>N77622</u>	<u>003</u>	Feb 22, 2006
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<u>AB</u>		<u>20MG</u>	<u>N77622</u>	<u>004</u>	Feb 22, 2006
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<u>AB</u>		<u>30MG</u>	<u>N77622</u>	<u>005</u>	Feb 22, 2006
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<u>AB</u>		<u>40MG</u>	<u>N77622</u>	<u>006</u>	Feb 22, 2006
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<u>AB</u>	IVAX PHARMS	<u>2.5MG</u>	<u>N75752</u>	<u>001</u>	Jul 01, 2002
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<u>AB</u>		<u>5MG</u>	<u>N75752</u>	<u>002</u>	Jul 01, 2002
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<u>AB</u>		<u>10MG</u>	<u>N75752</u>	<u>003</u>	Jul 01, 2002
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<u>AB</u>		<u>20MG</u>	<u>N75752</u>	<u>004</u>	Jul 01, 2002
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<u>AB</u>		<u>30MG</u>	<u>N75752</u>	<u>005</u>	Jul 01, 2002
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<u>AB</u>		<u>40MG</u>	<u>N75752</u>	<u>006</u>	Jul 01, 2002
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<u>AB</u>	LEK PHARMS	<u>2.5MG</u>	<u>N75999</u>	<u>001</u>	Jul 01, 2002
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<u>AB</u>		<u>5MG</u>	<u>N75999</u>	<u>002</u>	Jul 01, 2002
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<u>AB</u>		<u>10MG</u>	<u>N75999</u>	<u>003</u>	Jul 01, 2002
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<u>AB</u>		<u>20MG</u>	<u>N75999</u>	<u>004</u>	Jul 01, 2002
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<u>AB</u>		<u>30MG</u>	<u>N75999</u>	<u>005</u>	Jul 01, 2002
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<u>AB</u>		<u>40MG</u>	<u>N75999</u>	<u>006</u>	Jul 01, 2002
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<u>AB</u>	LUPIN	<u>2.5MG</u>	<u>N77321</u>	<u>001</u>	Sep 09, 2005
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<u>AB</u>		<u>5MG</u>	<u>N77321</u>	<u>002</u>	Sep 09, 2005
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PRESCRIPTION DRUG PRODUCT LIST

3 - 231 (of 370)

LISINOPRIL

TABLET; ORAL

LISINOPRIL

<u>AB</u>	<u>LUPIN</u>	<u>10MG</u>	<u>N77321</u>	<u>003</u>	Sep 09, 2005
<u>AB</u>		<u>20MG</u>	<u>N77321</u>	<u>004</u>	Sep 09, 2005
<u>AB</u>		<u>30MG</u>	<u>N77321</u>	<u>005</u>	Sep 09, 2005
<u>AB</u>		<u>40MG</u>	<u>N77321</u>	<u>006</u>	Sep 09, 2005
<u>AB</u>	<u>MYLAN</u>	<u>2.5MG</u>	<u>N76071</u>	<u>001</u>	Jul 01, 2002
<u>AB</u>		<u>5MG</u>	<u>N76071</u>	<u>002</u>	Jul 01, 2002
<u>AB</u>		<u>10MG</u>	<u>N76071</u>	<u>003</u>	Jul 01, 2002
<u>AB</u>		<u>20MG</u>	<u>N76071</u>	<u>004</u>	Jul 01, 2002
<u>AB</u>		<u>30MG</u>	<u>N76071</u>	<u>005</u>	Jul 01, 2002
<u>AB</u>		<u>40MG</u>	<u>N76071</u>	<u>006</u>	Jul 01, 2002
<u>AB</u>	<u>PAR PHARM</u>	<u>2.5MG</u>	<u>N75743</u>	<u>001</u>	Jul 01, 2002
<u>AB</u>		<u>5MG</u>	<u>N75743</u>	<u>002</u>	Jul 01, 2002
<u>AB</u>		<u>10MG</u>	<u>N75743</u>	<u>003</u>	Jul 01, 2002
<u>AB</u>		<u>20MG</u>	<u>N75743</u>	<u>004</u>	Jul 01, 2002
<u>AB</u>		<u>30MG</u>	<u>N75743</u>	<u>005</u>	Jul 01, 2002
<u>AB</u>		<u>40MG</u>	<u>N75743</u>	<u>006</u>	Jul 01, 2002
<u>AB</u>	<u>RANBAXY</u>	<u>2.5MG</u>	<u>N75944</u>	<u>001</u>	Jul 01, 2002
<u>AB</u>		<u>5MG</u>	<u>N75944</u>	<u>002</u>	Jul 01, 2002
<u>AB</u>		<u>10MG</u>	<u>N75944</u>	<u>003</u>	Jul 01, 2002
<u>AB</u>		<u>20MG</u>	<u>N75944</u>	<u>004</u>	Jul 01, 2002
<u>AB</u>		<u>30MG</u>	<u>N75944</u>	<u>006</u>	Feb 11, 2003
<u>AB</u>		<u>40MG</u>	<u>N75944</u>	<u>005</u>	Jul 01, 2002
<u>AB</u>	<u>SANDOZ</u>	<u>2.5MG</u>	<u>N75903</u>	<u>001</u>	Jul 01, 2002
<u>AB</u>		<u>2.5MG</u>	<u>N75994</u>	<u>001</u>	Jul 01, 2002
<u>AB</u>		<u>5MG</u>	<u>N75903</u>	<u>002</u>	Jul 01, 2002
<u>AB</u>		<u>5MG</u>	<u>N75994</u>	<u>002</u>	Jul 01, 2002
<u>AB</u>		<u>10MG</u>	<u>N75903</u>	<u>003</u>	Jul 01, 2002
<u>AB</u>		<u>10MG</u>	<u>N75994</u>	<u>003</u>	Jul 01, 2002
<u>AB</u>		<u>20MG</u>	<u>N75903</u>	<u>004</u>	Jul 01, 2002
<u>AB</u>		<u>20MG</u>	<u>N75994</u>	<u>004</u>	Jul 01, 2002
<u>AB</u>		<u>30MG</u>	<u>N75903</u>	<u>005</u>	Jul 01, 2002
<u>AB</u>		<u>30MG</u>	<u>N75994</u>	<u>005</u>	Jul 01, 2002
<u>AB</u>		<u>40MG</u>	<u>N75903</u>	<u>006</u>	Jul 01, 2002
<u>AB</u>		<u>40MG</u>	<u>N75994</u>	<u>006</u>	Jul 01, 2002
<u>AB</u>	<u>TEVA</u>	<u>2.5MG</u>	<u>N75783</u>	<u>001</u>	Jul 01, 2002
<u>AB</u>		<u>5MG</u>	<u>N75783</u>	<u>002</u>	Jul 01, 2002
<u>AB</u>		<u>10MG</u>	<u>N75783</u>	<u>003</u>	Jul 01, 2002
<u>AB</u>		<u>20MG</u>	<u>N75783</u>	<u>004</u>	Jul 01, 2002
<u>AB</u>		<u>30MG</u>	<u>N75783</u>	<u>005</u>	Jul 01, 2002
<u>AB</u>		<u>40MG</u>	<u>N75783</u>	<u>006</u>	Jul 01, 2002
<u>AB</u>	<u>WATSON LABS</u>	<u>2.5MG</u>	<u>N76059</u>	<u>001</u>	Jul 01, 2002
<u>AB</u>		<u>5MG</u>	<u>N76059</u>	<u>002</u>	Jul 01, 2002
<u>AB</u>		<u>10MG</u>	<u>N76059</u>	<u>003</u>	Jul 01, 2002
<u>AB</u>		<u>20MG</u>	<u>N76059</u>	<u>004</u>	Jul 01, 2002
<u>AB</u>		<u>30MG</u>	<u>N76059</u>	<u>005</u>	Jul 01, 2002
<u>AB</u>		<u>40MG</u>	<u>N76059</u>	<u>006</u>	Jul 01, 2002
<u>AB</u>	<u>WEST WARD</u>	<u>2.5MG</u>	<u>N76063</u>	<u>001</u>	Jul 01, 2002
<u>AB</u>		<u>5MG</u>	<u>N76063</u>	<u>002</u>	Jul 01, 2002
<u>AB</u>		<u>10MG</u>	<u>N76063</u>	<u>003</u>	Jul 01, 2002
<u>AB</u>		<u>20MG</u>	<u>N76063</u>	<u>004</u>	Jul 01, 2002
<u>AB</u>		<u>30MG</u>	<u>N76063</u>	<u>006</u>	Jun 27, 2003
<u>AB</u>		<u>40MG</u>	<u>N76063</u>	<u>005</u>	Jul 01, 2002
<u>PRINIVIL</u>					
<u>AB</u>	<u>MERCK</u>	<u>5MG</u>	<u>N19558</u>	<u>001</u>	Dec 29, 1987
<u>AB</u>		<u>10MG</u>	<u>N19558</u>	<u>002</u>	Dec 29, 1987
<u>AB</u>		<u>20MG</u>	<u>N19558</u>	<u>003</u>	Dec 29, 1987
<u>AB</u>		<u>40MG</u>	<u>N19558</u>	<u>004</u>	Oct 25, 1988

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 232 (of 370)

LISINOPRIL

TABLET; ORAL

ZESTRIL

<u>AB</u>	ASTRAZENECA	<u>2.5MG</u>	<u>N19777</u>	<u>005</u>	Apr 29, 1993
<u>AB</u>		<u>5MG</u>	<u>N19777</u>	<u>001</u>	May 19, 1988
<u>AB</u>		<u>10MG</u>	<u>N19777</u>	<u>002</u>	May 19, 1988
<u>AB</u>		<u>20MG</u>	<u>N19777</u>	<u>003</u>	May 19, 1988
<u>AB</u>		<u>30MG</u>	<u>N19777</u>	<u>006</u>	Jan 20, 1999
<u>AB</u>	+	<u>40MG</u>	<u>N19777</u>	<u>004</u>	May 19, 1988

LITHIUM CARBONATE

CAPSULE; ORAL

ESKALITH

<u>AB</u>	GLAXOSMITHKLINE	<u>300MG</u>	<u>N16860</u>	<u>001</u>	
	<u>LITHIUM CARBONATE</u>				
<u>AB</u>	APOTEX INC	<u>300MG</u>	<u>N76795</u>	<u>001</u>	Nov 22, 2004
<u>AB</u>	ROXANE	<u>150MG</u>	<u>N17812</u>	<u>002</u>	Jan 28, 1987
<u>AB</u>		<u>300MG</u>	<u>N17812</u>	<u>001</u>	
	+	600MG	N17812	003	Jan 28, 1987
<u>AB</u>	WEST WARD	<u>150MG</u>	<u>N76243</u>	<u>002</u>	Feb 24, 2003
<u>AB</u>		<u>300MG</u>	<u>N76243</u>	<u>001</u>	Jun 27, 2002

TABLET; ORAL

LITHIUM CARBONATE

+ ROXANE 300MG N18558 001 Jan 29, 1982

TABLET, EXTENDED RELEASE; ORAL

LITHIUM CARBONATE

<u>AB</u>	BARR	<u>300MG</u>	<u>N76170</u>	<u>001</u>	Jun 10, 2002
<u>AB</u>	ROXANE	<u>300MG</u>	<u>N76832</u>	<u>001</u>	Oct 28, 2004
<u>AB</u>	+	<u>450MG</u>	<u>N76691</u>	<u>001</u>	Jan 05, 2004
<u>AB</u>	WEST WARD	<u>450MG</u>	<u>N76490</u>	<u>001</u>	Jun 17, 2003
	<u>LITHOBID</u>				
<u>AB</u>	+	JDS PHARMS 300MG	<u>N18027</u>	<u>001</u>	

LITHIUM CITRATE

SYRUP; ORAL

LITHIUM CITRATE

<u>AA</u>	MORTON GROVE	<u>EQ 300MG CARBONATE/5ML</u>	<u>N70755</u>	<u>001</u>	May 21, 1986
<u>AA</u>	+	<u>EQ 300MG CARBONATE/5ML</u>	<u>N18421</u>	<u>001</u>	

LODOXAMIDE TROMETHAMINE

SOLUTION/DROPS; OPHTHALMIC

+ ALCON EQ 0.1% BASE N20191 001 Sep 23, 1993

LOMEFLOXACIN HYDROCHLORIDE

TABLET; ORAL

MAXAQUIN

+ PHARMACIA EQ 400MG BASE N20013 001 Feb 21, 1992

LOMUSTINE

CAPSULE; ORAL

CEENU

	BRISTOL MYERS SQUIBB	10MG	N17588	001	
		40MG	N17588	002	
	+	100MG	N17588	003	

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 233 (of 370)

LOPERAMIDE HYDROCHLORIDE

CAPSULE; ORAL

IMODIUMAB + MCNEIL PED 2MGN17694 001LOPERAMIDE HYDROCHLORIDEAB MYLAN 2MGN72741 001

Sep 18, 1991

AB SANDOZ 2MGN72993 001

Aug 28, 1992

AB TEVA 2MGN73122 001

Aug 30, 1991

AB 2MGN73192 001

Apr 30, 1992

LOPINAVIR; RITONAVIR

CAPSULE; ORAL

KALETRA

+ ABBOTT 133.3MG;33.3MG

N21226 001

Sep 15, 2000

SOLUTION; ORAL

KALETRA

+ ABBOTT 80MG/ML;20MG/ML

N21251 001

Sep 15, 2000

TABLET; ORAL

KALETRA

+ ABBOTT 200MG;50MG

N21906 001

Oct 28, 2005

LORAZEPAM

CONCENTRATE; ORAL

LORAZEPAM INTENSOL

+ ROXANE 2MG/ML

N72755 001

Jun 28, 1991

INJECTABLE; INJECTION

ATIVANAP + BAXTER HLTHCARE CORP 2MG/MLN18140 001AP + 4MG/MLN18140 002LORAZEPAMAP BEDFORD 2MG/MLN77076 001

Jul 13, 2005

AP 4MG/MLN77076 002

Jul 13, 2005

AP HOSPIRA 2MG/MLN74243 001

Apr 12, 1994

AP 2MG/MLN74280 001

May 27, 1994

AP 2MG/MLN74282 001

May 27, 1994

AP 2MG/MLN74300 001

Apr 12, 1994

AP 4MG/MLN74243 002

Apr 12, 1994

AP 4MG/MLN74280 002

May 27, 1994

AP 4MG/MLN74282 002

May 27, 1994

AP 4MG/MLN74300 003

Mar 19, 1997

AP INTL MEDICATION SYS 2MG/MLN76150 001

Nov 15, 2004

AP WATSON LABS 2MG/MLN74276 001

Apr 15, 1994

AP 4MG/MLN74276 002

Apr 15, 1994

LORAZEPAM PRESERVATIVE FREEAP BEDFORD LABS 2MG/MLN77074 001

Jul 13, 2005

AP 4MG/MLN77074 002

Jul 13, 2005

SOLUTION; ORAL

LORAZEPAM

ROXANE 0.5MG/5ML

N74648 001

Mar 18, 1997

TABLET; ORAL

ATIVANAB BIOVAIL 0.5MGN17794 001AB 1MGN17794 002AB + 2MGN17794 003LORAZEPAMAB ACTAVIS ELIZABETH 0.5MGN71403 001

Apr 21, 1987

AB 1MGN71404 001

Apr 21, 1987

AB 2MGN71141 001

Apr 21, 1987

AB IVAX PHARMS 0.5MGN77396 001

Dec 13, 2006

PRESCRIPTION DRUG PRODUCT LIST

3 - 234 (of 370)

LORAZEPAM

TABLET; ORAL

LORAZEPAM

<u>AB</u>	IVAX PHARMS	<u>1MG</u>	<u>N77396</u>	<u>002</u>	Dec 13, 2006
<u>AB</u>		<u>2MG</u>	<u>N77396</u>	<u>003</u>	Dec 13, 2006
<u>AB</u>	MUTUAL PHARM	<u>0.5MG</u>	<u>N72553</u>	<u>001</u>	Mar 29, 1991
<u>AB</u>		<u>1MG</u>	<u>N72554</u>	<u>001</u>	Mar 29, 1991
<u>AB</u>		<u>2MG</u>	<u>N72555</u>	<u>001</u>	Mar 29, 1991
<u>AB</u>	MYLAN	<u>0.5MG</u>	<u>N71589</u>	<u>001</u>	Oct 13, 1987
<u>AB</u>		<u>0.5MG</u>	<u>N77657</u>	<u>001</u>	Mar 16, 2006
<u>AB</u>		<u>1MG</u>	<u>N71590</u>	<u>001</u>	Oct 13, 1987
<u>AB</u>		<u>1MG</u>	<u>N77657</u>	<u>002</u>	Mar 16, 2006
<u>AB</u>		<u>2MG</u>	<u>N71591</u>	<u>001</u>	Oct 13, 1987
<u>AB</u>		<u>2MG</u>	<u>N77657</u>	<u>003</u>	Mar 16, 2006
<u>AB</u>	RANBAXY	<u>0.5MG</u>	<u>N76045</u>	<u>001</u>	Aug 29, 2001
<u>AB</u>		<u>1MG</u>	<u>N76045</u>	<u>002</u>	Aug 29, 2001
<u>AB</u>		<u>2MG</u>	<u>N76045</u>	<u>003</u>	Aug 29, 2001
<u>AB</u>	SANDOZ	<u>0.5MG</u>	<u>N71193</u>	<u>001</u>	Apr 15, 1988
<u>AB</u>		<u>1MG</u>	<u>N71194</u>	<u>001</u>	Apr 15, 1988
<u>AB</u>		<u>2MG</u>	<u>N71195</u>	<u>001</u>	Apr 15, 1988
<u>AB</u>	VINTAGE PHARMS	<u>0.5MG</u>	<u>N77754</u>	<u>001</u>	May 10, 2006
<u>AB</u>		<u>1MG</u>	<u>N77754</u>	<u>002</u>	May 10, 2006
<u>AB</u>		<u>2MG</u>	<u>N77754</u>	<u>003</u>	May 10, 2006
<u>AB</u>	WATSON LABS	<u>0.5MG</u>	<u>N71117</u>	<u>001</u>	Jul 24, 1986
<u>AB</u>		<u>0.5MG</u>	<u>N72926</u>	<u>001</u>	Oct 31, 1991
<u>AB</u>		<u>1MG</u>	<u>N71118</u>	<u>001</u>	Jul 24, 1986
<u>AB</u>		<u>1MG</u>	<u>N72927</u>	<u>001</u>	Oct 31, 1991
<u>AB</u>		<u>2MG</u>	<u>N71110</u>	<u>001</u>	Jul 24, 1986
<u>AB</u>		<u>2MG</u>	<u>N72928</u>	<u>001</u>	Oct 31, 1991

LOSARTAN POTASSIUM

TABLET; ORAL

COZAAR

MERCK

25MG

N20386 001 Apr 14, 1995

50MG

N20386 002 Apr 14, 1995

+

100MG

N20386 003 Oct 13, 1998

LOTEPREDNOL ETABONATE

SUSPENSION/DROPS; OPHTHALMIC

ALREX

+ BAUSCH AND LOMB 0.2%

N20803 001 Mar 09, 1998

LOTEMAX

+ BAUSCH AND LOMB 0.5%

N20583 001 Mar 09, 1998

LOTEPREDNOL ETABONATE; TOBRAMYCIN

SUSPENSION/DROPS; OPHTHALMIC

+ BAUSCH AND LOMB 0.5%;0.3%

N50804 001 Dec 14, 2004

LOVASTATIN

TABLET; ORAL

LOVASTATIN

<u>AB</u>	ACTAVIS ELIZABETH	<u>10MG</u>	<u>N75828</u>	<u>001</u>	Dec 17, 2001
<u>AB</u>		<u>20MG</u>	<u>N75828</u>	<u>002</u>	Dec 17, 2001
<u>AB</u>		<u>40MG</u>	<u>N75828</u>	<u>003</u>	Dec 17, 2001
<u>AB</u>	CARLSBAD	<u>10MG</u>	<u>N75991</u>	<u>001</u>	Jun 05, 2002
<u>AB</u>		<u>20MG</u>	<u>N75991</u>	<u>002</u>	Jun 05, 2002
<u>AB</u>		<u>40MG</u>	<u>N75991</u>	<u>003</u>	Jun 05, 2002
<u>AB</u>	GENPHARM	<u>10MG</u>	<u>N75935</u>	<u>001</u>	Dec 17, 2001

PRESCRIPTION DRUG PRODUCT LIST

3 - 235 (of 370)

LOVASTATIN

TABLET; ORAL

LOVASTATIN

<u>AB</u>	GENPHARM	<u>20MG</u>	<u>N75935</u>	<u>002</u>	Dec 17, 2001
<u>AB</u>		<u>40MG</u>	<u>N75935</u>	<u>003</u>	Dec 17, 2001
<u>AB</u>	MUTUAL PHARM	<u>10MG</u>	<u>N77520</u>	<u>001</u>	Apr 14, 2006
<u>AB</u>		<u>20MG</u>	<u>N77520</u>	<u>002</u>	Apr 14, 2006
<u>AB</u>		<u>40MG</u>	<u>N77520</u>	<u>003</u>	Apr 14, 2006
<u>AB</u>	MYLAN	<u>10MG</u>	<u>N75451</u>	<u>001</u>	Dec 17, 2001
<u>AB</u>		<u>20MG</u>	<u>N75451</u>	<u>002</u>	Dec 17, 2001
<u>AB</u>		<u>40MG</u>	<u>N75451</u>	<u>003</u>	Dec 17, 2001
<u>AB</u>	SANDOZ	<u>10MG</u>	<u>N75300</u>	<u>001</u>	Dec 17, 2001
<u>AB</u>		<u>10MG</u>	<u>N75636</u>	<u>001</u>	Dec 17, 2001
<u>AB</u>		<u>20MG</u>	<u>N75300</u>	<u>002</u>	Dec 17, 2001
<u>AB</u>		<u>20MG</u>	<u>N75636</u>	<u>002</u>	Dec 17, 2001
<u>AB</u>		<u>40MG</u>	<u>N75300</u>	<u>003</u>	Dec 17, 2001
<u>AB</u>		<u>40MG</u>	<u>N75636</u>	<u>003</u>	Dec 17, 2001
<u>AB</u>	TEVA	<u>10MG</u>	<u>N75551</u>	<u>003</u>	Dec 17, 2001
<u>AB</u>		<u>20MG</u>	<u>N75551</u>	<u>002</u>	Dec 17, 2001
<u>AB</u>		<u>40MG</u>	<u>N75551</u>	<u>001</u>	Dec 17, 2001
<u>MEVACOR</u>					
<u>AB</u>	MERCK	<u>20MG</u>	<u>N19643</u>	<u>003</u>	Aug 31, 1987
<u>AB</u>	+	<u>40MG</u>	<u>N19643</u>	<u>004</u>	Dec 14, 1988

TABLET, EXTENDED RELEASE; ORAL

ALTOPREV

	ANDRX LABS LLC	10MG	N21316	001	Jun 26, 2002
		20MG	N21316	002	Jun 26, 2002
		40MG	N21316	003	Jun 26, 2002
	+	60MG	N21316	004	Jun 26, 2002

LOVASTATIN; NIACIN

TABLET, EXTENDED RELEASE; ORAL

ADVICOR

+	KOS LIFE	20MG;500MG	N21249	001	Dec 17, 2001
+		20MG;750MG	N21249	002	Dec 17, 2001
+		20MG;1GM	N21249	003	Dec 17, 2001
+		40MG;1GM	N21249	004	Apr 27, 2006

LOXAPINE SUCCINATE

CAPSULE; ORAL

LOXAPINE

<u>AB</u>	MYLAN	<u>EQ 5MG BASE</u>	<u>N76762</u>	<u>001</u>	Nov 01, 2004
<u>AB</u>		<u>EQ 10MG BASE</u>	<u>N76762</u>	<u>002</u>	Nov 01, 2004
<u>AB</u>		<u>EQ 25MG BASE</u>	<u>N76762</u>	<u>003</u>	Nov 01, 2004
<u>AB</u>		<u>EQ 50MG BASE</u>	<u>N76762</u>	<u>004</u>	Nov 01, 2004

LOXAPINE SUCCINATE

<u>AB</u>	ACTAVIS TOTOWA	<u>EQ 5MG BASE</u>	<u>N76868</u>	<u>001</u>	Aug 04, 2005
<u>AB</u>		<u>EQ 10MG BASE</u>	<u>N76868</u>	<u>002</u>	Aug 04, 2005
<u>AB</u>		<u>EQ 25MG BASE</u>	<u>N76868</u>	<u>003</u>	Aug 04, 2005
<u>AB</u>		<u>EQ 50MG BASE</u>	<u>N76868</u>	<u>004</u>	Aug 04, 2005
<u>AB</u>	WATSON LABS	<u>EQ 5MG BASE</u>	<u>N72204</u>	<u>001</u>	Jun 15, 1988
<u>AB</u>		<u>EQ 10MG BASE</u>	<u>N72205</u>	<u>001</u>	Jun 15, 1988
<u>AB</u>		<u>EQ 25MG BASE</u>	<u>N72206</u>	<u>001</u>	Jun 15, 1988
<u>AB</u>		<u>EQ 50MG BASE</u>	<u>N72062</u>	<u>001</u>	Jun 15, 1988

LOXITANE

<u>AB</u>	WATSON LABS	<u>EQ 5MG BASE</u>	<u>N17525</u>	<u>001</u>	
<u>AB</u>		<u>EQ 10MG BASE</u>	<u>N17525</u>	<u>002</u>	
<u>AB</u>	+	<u>EQ 25MG BASE</u>	<u>N17525</u>	<u>003</u>	
<u>AB</u>		<u>EQ 50MG BASE</u>	<u>N17525</u>	<u>004</u>	

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 236 (of 370)

LUBIPROSTONE

CAPSULE; ORAL

AMITIZA

+ SUCAMPO PHARMS 24UGM N21908 001 Jan 31, 2006

LUTROPIN ALFA

INJECTABLE; SUBCUTANEOUS

LUVERIS

+ SERONO INC 75 IU/VIAL N21322 001 Oct 08, 2004

MAFENIDE ACETATE

CREAM; TOPICAL

SULFAMYLON

+ MYLAN BERTEK EQ 85MG BASE/GM N16763 001

FOR SOLUTION; TOPICAL

SULFAMYLON

+ MYLAN BERTEK 5% N19832 003 Jun 05, 1998

MAGNESIUM ACETATE TETRAHYDRATE; POTASSIUM ACETATE; SODIUM CHLORIDE

INJECTABLE; INJECTION

PLASMA-LYTE 56 IN PLASTIC CONTAINER

BAXTER HLTHCARE 32MG/100ML;128MG/100ML;234MG/100ML N19047 001 Jun 15, 1984

MAGNESIUM CHLORIDE; POTASSIUM CHLORIDE; POTASSIUM PHOSPHATE, MONOBASIC; SODIUM ACETATE; SODIUM CHLORIDE; SODIUM GLUCONATE; SODIUM PHOSPHATE, DIBASIC, HEPTAHYDRATE

INJECTABLE; INJECTION

ISOLYTE S PH 7.4 IN PLASTIC CONTAINER

B BRAUN 30MG/100ML;37MG/100ML;0.82MG/100ML;370MG/100ML;530MG/100ML;500MG/100ML;12MG/100ML N19696 001 Sep 29, 1989

MAGNESIUM CHLORIDE; POTASSIUM CHLORIDE; SODIUM ACETATE; SODIUM CHLORIDE; SODIUM GLUCONATE

INJECTABLE; INJECTION

ISOLYTE S IN PLASTIC CONTAINERAP B BRAUN 30MG/100ML;37MG/100ML;370MG/100ML;530MG/100ML;500MG/100ML N19711 001 Sep 29, 1989

NORMOSOL-R IN PLASTIC CONTAINER

HOSPIRA 30MG/100ML;37MG/100ML;222MG/100ML;526MG/100ML;502MG/100ML N17586 001

PLASMA-LYTE 148 IN WATER IN PLASTIC CONTAINERAP BAXTER HLTHCARE 30MG/100ML;37MG/100ML;368MG/100ML;526MG/100ML;502MG/100ML N17378 001PLASMA-LYTE A IN PLASTIC CONTAINERAP BAXTER HLTHCARE 30MG/100ML;37MG/100ML;368MG/100ML;526MG/100ML;502MG/100ML N17378 002 Nov 22, 1982

SOLUTION; IRRIGATION

PHYSIOLYTE IN PLASTIC CONTAINER

B BRAUN 30MG/100ML;37MG/100ML;370MG/100ML;530MG/100ML;500MG/100ML N19024 001 Jun 08, 1984

PHYSIOSOL IN PLASTIC CONTAINER

HOSPIRA 30MG/100ML;37MG/100ML;222MG/100ML;526MG/100ML;502MG/100ML N17637 002 Jul 08, 1982

PHYSIOSOL PH 7.4 IN PLASTIC CONTAINER

HOSPIRA 30MG/100ML;37MG/100ML;222MG/100ML;526MG/100ML;502MG/100ML N18406 002 Jul 08, 1982

MAGNESIUM CHLORIDE; SODIUM BICARBONATE; SODIUM CHLORIDE

SOLUTION; INJECTION

NORMOCARB HF 25

+ DIALYSIS SUPS 0.21GM/100ML;2.8GM/100ML;9.07GM/100ML N21910 001 Jul 26, 2006

PRESCRIPTION DRUG PRODUCT LIST

3 - 237 (of 370)

MAGNESIUM CHLORIDE; SODIUM BICARBONATE; SODIUM CHLORIDE

SOLUTION; INJECTION

NORMOCARB HF 35

+ DIALYSIS SUPS	0.21GM/100ML; 3.97GM/100ML; 8.3GM/100ML	N21910	002	Jul 26, 2006
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MAGNESIUM HYDROXIDE; OMEPRAZOLE; SODIUM BICARBONATE

TABLET, CHEWABLE; ORAL

ZEGERID

SANTARUS

700MG; 20MG; 600MG

N21850 001 Mar 24, 2006

+	700MG; 40MG; 600MG	N21850	002	Mar 24, 2006
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MAGNESIUM SULFATE

INJECTABLE; INJECTION

MAGNESIUM SULFATE

<u>AP</u> + ABRAXIS PHARM	<u>500MG/ML</u>	<u>N19316</u>	<u>001</u>	Sep 08, 1986
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<u>AP</u> HOSPIRA	<u>500MG/ML</u>	<u>N75151</u>	<u>001</u>	Apr 25, 2000
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MAGNESIUM SULFATE IN DEXTROSE 5% IN PLASTIC CONTAINER

+ HOSPIRA	1GM/100ML	N20488	001	Jul 11, 1995
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+	2GM/100ML	N20488	002	Jul 11, 1995
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MAGNESIUM SULFATE IN PLASTIC CONTAINER

+ HOSPIRA	80MG/ML	N20309	002	Jun 24, 1994
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+	4GM/100ML	N20309	001	Jun 24, 1994
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MAGNESIUM SULFATE; POTASSIUM CHLORIDE; POTASSIUM PHOSPHATE, MONOBASIC; SODIUM CHLORIDE; SODIUM PHOSPHATE

SOLUTION; IRRIGATION

TIS-U-SOL

<u>AT</u> BAXTER HLTHCARE	<u>20MG/100ML; 40MG/100ML; 6.25MG/100ML; 800MG/100ML; 8.75MG/100ML</u>	<u>N18508</u>	<u>001</u>	Feb 19, 1982
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TIS-U-SOL IN PLASTIC CONTAINER

<u>AT</u> BAXTER HLTHCARE	<u>20MG/100ML; 40MG/100ML; 6.25MG/100ML; 800MG/100ML; 8.75MG/100ML</u>	<u>N18336</u>	<u>001</u>	
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MALATHION

LOTION; TOPICAL

OVIDE

+ TARO PHARMS NORTH	0.5%	N18613	001	Aug 02, 1982
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MANGANESE CHLORIDE

INJECTABLE; INJECTION

MANGANESE CHLORIDE IN PLASTIC CONTAINER

HOSPIRA

EQ 0.1MG MANGANESE/ML

N18962 001 Jun 26, 1986

MANNITOL

INJECTABLE; INJECTION

MANNITOL 10%

<u>AP</u> B BRAUN	<u>10GM/100ML</u>	<u>N16080</u>	<u>002</u>	
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MANNITOL 10% IN PLASTIC CONTAINER

<u>AP</u> B BRAUN	<u>10GM/100ML</u>	<u>N20006</u>	<u>002</u>	Jul 26, 1993
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<u>AP</u> HOSPIRA	<u>10GM/100ML</u>	<u>N19603</u>	<u>002</u>	Jan 08, 1987
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MANNITOL 10% W/ DEXTROSE 5% IN DISTILLED WATER

<u>AP</u> B BRAUN	<u>10GM/100ML</u>	<u>N16080</u>	<u>006</u>	
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MANNITOL 15%

<u>AP</u> B BRAUN	<u>15GM/100ML</u>	<u>N16080</u>	<u>003</u>	
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MANNITOL 15% IN PLASTIC CONTAINER

<u>AP</u> B BRAUN	<u>15GM/100ML</u>	<u>N20006</u>	<u>003</u>	Jul 26, 1993
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<u>AP</u> HOSPIRA	<u>15GM/100ML</u>	<u>N19603</u>	<u>003</u>	Jan 08, 1990
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MANNITOL 15% W/ DEXTROSE 5% IN SODIUM CHLORIDE 0.45%

<u>AP</u> B BRAUN	<u>15GM/100ML</u>	<u>N16080</u>	<u>005</u>	
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PRESCRIPTION DRUG PRODUCT LIST

3 - 238 (of 370)

MANNITOL

INJECTABLE; INJECTION

MANNITOL 20%

<u>AP</u>	B BRAUN	<u>20GM/100ML</u>	<u>N14738</u>	<u>001</u>	
<u>AP</u>		<u>20GM/100ML</u>	<u>N16080</u>	<u>004</u>	
	<u>MANNITOL 20% IN PLASTIC CONTAINER</u>				
<u>AP</u>	B BRAUN	<u>20GM/100ML</u>	<u>N20006</u>	<u>004</u>	Jul 26, 1993
<u>AP</u>	HOSPIRA	<u>20GM/100ML</u>	<u>N19603</u>	<u>004</u>	Jan 08, 1990

MANNITOL 25%

<u>AP</u>	ABRAXIS PHARM	<u>12.5GM/50ML</u>	<u>N80677</u>	<u>001</u>	
<u>AP</u>	HOSPIRA	<u>12.5GM/50ML</u>	<u>N16269</u>	<u>006</u>	Aug 25, 1994
<u>AP</u>	INTL MEDICATION	<u>12.5GM/50ML</u>	<u>N83051</u>	<u>001</u>	
<u>AP</u>	LUITPOLD	<u>12.5GM/50ML</u>	<u>N87409</u>	<u>001</u>	Jan 21, 1982

MANNITOL 5%

<u>AP</u>	B BRAUN	<u>5GM/100ML</u>	<u>N16080</u>	<u>001</u>	
	<u>MANNITOL 5% IN PLASTIC CONTAINER</u>				
<u>AP</u>	B BRAUN	<u>5GM/100ML</u>	<u>N20006</u>	<u>001</u>	Jul 26, 1993
<u>AP</u>	HOSPIRA	<u>5GM/100ML</u>	<u>N19603</u>	<u>001</u>	Jan 08, 1987

MANNITOL 5% W/ DEXTROSE 5% IN SODIUM CHLORIDE 0.12%

<u>AP</u>	B BRAUN	<u>5GM/100ML</u>	<u>N16080</u>	<u>007</u>	
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OSMITROL 10% IN WATER

<u>AP</u>	BAXTER HLTHCARE	<u>10GM/100ML</u>	<u>N13684</u>	<u>002</u>	
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OSMITROL 10% IN WATER IN PLASTIC CONTAINER

<u>AP</u>	BAXTER HLTHCARE	<u>10GM/100ML</u>	<u>N13684</u>	<u>006</u>	
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OSMITROL 15% IN WATER

<u>AP</u>	BAXTER HLTHCARE	<u>15GM/100ML</u>	<u>N13684</u>	<u>004</u>	
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OSMITROL 15% IN WATER IN PLASTIC CONTAINER

<u>AP</u>	BAXTER HLTHCARE	<u>15GM/100ML</u>	<u>N13684</u>	<u>008</u>	
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OSMITROL 20% IN WATER

<u>AP</u>	BAXTER HLTHCARE	<u>20GM/100ML</u>	<u>N13684</u>	<u>003</u>	
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OSMITROL 20% IN WATER IN PLASTIC CONTAINER

<u>AP</u>	BAXTER HLTHCARE	<u>20GM/100ML</u>	<u>N13684</u>	<u>007</u>	
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OSMITROL 5% IN WATER

<u>AP</u>	BAXTER HLTHCARE	<u>5GM/100ML</u>	<u>N13684</u>	<u>001</u>	
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OSMITROL 5% IN WATER IN PLASTIC CONTAINER

<u>AP</u>	BAXTER HLTHCARE	<u>5GM/100ML</u>	<u>N13684</u>	<u>005</u>	
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SOLUTION; IRRIGATION

RESECTISOL IN PLASTIC CONTAINER

	B BRAUN	5GM/100ML	N16772	002	
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MANNITOL; SORBITOL

SOLUTION; IRRIGATION

SORBITOL-MANNITOL IN PLASTIC CONTAINER

<u>AT</u>	HOSPIRA	<u>540MG/100ML; 2.7GM/100ML</u>	<u>N17636</u>	<u>001</u>	
<u>AT</u>		<u>540MG/100ML; 2.7GM/100ML</u>	<u>N18316</u>	<u>001</u>	

MAPROTILINE HYDROCHLORIDE

TABLET; ORAL

MAPROTILINE HYDROCHLORIDE

<u>AB</u>	MYLAN	<u>25MG</u>	<u>N72284</u>	<u>001</u>	Oct 03, 1988
<u>AB</u>	+	<u>50MG</u>	<u>N72285</u>	<u>001</u>	Oct 03, 1988
<u>AB</u>		<u>75MG</u>	<u>N72286</u>	<u>001</u>	Oct 03, 1988
<u>AB</u>	WATSON LABS	<u>25MG</u>	<u>N72162</u>	<u>001</u>	Jun 01, 1988
<u>AB</u>		<u>50MG</u>	<u>N72163</u>	<u>001</u>	Jun 01, 1988
<u>AB</u>		<u>75MG</u>	<u>N72164</u>	<u>001</u>	Jun 01, 1988

MEBENDAZOLE

TABLET, CHEWABLE; ORAL

MEBENDAZOLE

+	TEVA PHARMS	100MG	N73580	001	Jan 04, 1995
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27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 239 (of 370)

MECAMYLAMINE HYDROCHLORIDE

TABLET; ORAL

INVERSINE

+ TARGACEPT

2.5MG

N10251 001

MECASERMIN RECOMBINANT

INJECTABLE; SUBCUTANEOUS

INCRELEX

+ TERCICA

40MG/4ML(10MG/ML)

N21839 001

Aug 30, 2005

MECASERMIN RINFABATE RECOMBINANT

INJECTABLE; SUBCUTANEOUS

IPLEX

+ INSMED

36MG/0.6ML

N21884 001

Dec 12, 2005

MECHLORETHAMINE HYDROCHLORIDE

INJECTABLE; INJECTION

MUSTARGEN

+ OVATION PHARMS

10MG/VIAL

N06695 001

MECLIZINE HYDROCHLORIDE

TABLET; ORAL

ANTIVERTAA + PFIZER12.5MGN10721 006AA +25MGN10721 004AA +50MGN10721 001

Jan 20, 1982

MECLIZINE HYDROCHLORIDEAA BUNDY12.5MGN84382 001AA25MGN84872 001AA

IVAX PHARMS

12.5MGN84975 001AA25MGN84657 001AA

PAR PHARM

12.5MGN87127 001AA25MGN87128 001AA50MGN89674 001

Mar 31, 1988

AA

PLIVA

12.5MGN88732 001

Dec 11, 1985

AA25MGN88734 001

Dec 11, 1985

AA

SANDOZ

12.5MGN84843 002

May 22, 1989

AA25MGN84092 003

May 22, 1989

AA

VINTAGE PHARMS

12.5MGN40179 001

Jan 30, 1997

AA25MGN40179 002

Jan 30, 1997

AA

WATSON LABS

25MGN85740 001

TABLET, CHEWABLE; ORAL

ANTIVERTAA + PFIZER25MGN10721 005MECLIZINE HYDROCHLORIDEAA

PLIVA

25MGN88733 001

Dec 11, 1985

MECLOFENAMATE SODIUM

CAPSULE; ORAL

MECLOFENAMATE SODIUMAB MYLANEQ 50MG BASEN71080 001

Sep 03, 1986

AB +EQ 100MG BASEN71081 001

Sep 03, 1986

AB

SANDOZ

EQ 50MG BASEN72262 001

Nov 29, 1988

ABEQ 100MG BASEN72263 001

Nov 29, 1988

AB

WATSON LABS

EQ 50MG BASEN71468 001

Apr 15, 1987

ABEQ 100MG BASEN71469 001

Apr 15, 1987

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 240 (of 370)

MEDROXYPROGESTERONE ACETATE

INJECTABLE; INJECTION

DEPO-PROVERA

<u>AB</u>	+	PHARMACIA AND UPJOHN	<u>150MG/ML</u>	<u>N20246</u>	<u>001</u>	Oct 29, 1992
	+		400MG/ML	N12541	003	

MEDROXYPROGESTERONE ACETATE

<u>AB</u>		SICOR PHARMS	<u>150MG/ML</u>	<u>N76552</u>	<u>001</u>	Oct 27, 2004
<u>AB</u>			<u>150MG/ML</u>	<u>N76553</u>	<u>001</u>	Jul 28, 2004

INJECTABLE; SUBCUTANEOUS

DEPO-SUBQ PROVERA 104

+	PHARMACIA AND UPJOHN	104MG/0.65ML	N21583	001	Dec 17, 2004
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TABLET; ORAL

MEDROXYPROGESTERONE ACETATE

<u>AB</u>		BARR	<u>2.5MG</u>	<u>N40159</u>	<u>001</u>	Aug 09, 1996
<u>AB</u>			<u>5MG</u>	<u>N40159</u>	<u>002</u>	Aug 09, 1996
<u>AB</u>			<u>10MG</u>	<u>N40159</u>	<u>003</u>	Aug 09, 1996
<u>AB</u>		DURAMED PHARMS BARR	<u>2.5MG</u>	<u>N40311</u>	<u>001</u>	Dec 01, 1999
<u>AB</u>			<u>5MG</u>	<u>N40311</u>	<u>002</u>	Dec 01, 1999
<u>AB</u>			<u>10MG</u>	<u>N40311</u>	<u>003</u>	Dec 01, 1999
BP		USL PHARMA	10MG	N88484	001	Jul 26, 1984

PROVERA

<u>AB</u>		PHARMACIA AND UPJOHN	<u>2.5MG</u>	<u>N11839</u>	<u>001</u>	
<u>AB</u>			<u>5MG</u>	<u>N11839</u>	<u>003</u>	
<u>AB</u>	+		<u>10MG</u>	<u>N11839</u>	<u>004</u>	

MEDRYSONE

SUSPENSION; OPHTHALMIC

HMS

+	ALLERGAN	1%	N16624	003	
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MEFENAMIC ACID

CAPSULE; ORAL

PONSTEL

+	SCIELE PHARMA INC	250MG	N15034	003	
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MEFLOQUINE HYDROCHLORIDE

TABLET; ORAL

LARIAM

<u>AB</u>	+	ROCHE	<u>250MG</u>	<u>N19591</u>	<u>001</u>	May 02, 1989
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MEFLOQUINE

<u>AB</u>		BARR	<u>250MG</u>	<u>N76392</u>	<u>001</u>	Dec 29, 2003
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MEFLOQUINE HYDROCHLORIDE

<u>AB</u>		ROXANE	<u>250MG</u>	<u>N76523</u>	<u>001</u>	Oct 01, 2004
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<u>AB</u>		SANDOZ	<u>250MG</u>	<u>N76175</u>	<u>001</u>	Feb 20, 2002
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MEGESTROL ACETATE

SUSPENSION; ORAL

MEGACE

<u>AB</u>	+	BRISTOL MYERS SQUIBB	<u>40MG/ML</u>	<u>N20264</u>	<u>001</u>	Sep 10, 1993
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MEGACE ES

+	PAR PHARM	125MG/ML	N21778	001	Jul 05, 2005
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MEGESTROL ACETATE

<u>AB</u>		APOTEX INC	<u>40MG/ML</u>	<u>N77404</u>	<u>001</u>	Feb 16, 2006
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<u>AB</u>		MORTON GROVE	<u>40MG/ML</u>	<u>N76721</u>	<u>001</u>	Nov 01, 2004
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<u>AB</u>		PAR PHARM	<u>40MG/ML</u>	<u>N75671</u>	<u>001</u>	Jul 25, 2001
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<u>AB</u>		ROXANE	<u>40MG/ML</u>	<u>N75997</u>	<u>001</u>	Feb 15, 2002
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<u>AB</u>		TEVA PHARMS	<u>40MG/ML</u>	<u>N75681</u>	<u>001</u>	May 05, 2003
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27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 241 (of 370)

MEGESTROL ACETATE

TABLET; ORAL

MEGACE

AB BRISTOL MYERS SQUIBB 20MG
AB + 40MG

N16979 001
N16979 002

MEGESTROL ACETATE

<u>AB</u>	BARR	<u>20MG</u>	<u>N74621</u>	<u>002</u>	Aug 16, 1996
<u>AB</u>		<u>40MG</u>	<u>N74621</u>	<u>001</u>	Nov 30, 1995
<u>AB</u>	PAR PHARM	<u>20MG</u>	<u>N72422</u>	<u>001</u>	Aug 08, 1988
<u>AB</u>		<u>40MG</u>	<u>N72423</u>	<u>001</u>	Aug 08, 1988
<u>AB</u>	ROXANE	<u>20MG</u>	<u>N74458</u>	<u>001</u>	Sep 29, 1995
<u>AB</u>		<u>40MG</u>	<u>N74458</u>	<u>002</u>	Sep 29, 1995
<u>AB</u>	TEVA	<u>40MG</u>	<u>N74745</u>	<u>001</u>	Feb 27, 1998

MELOXICAM

SUSPENSION; ORAL

MOBIC

+ BOEHRINGER INGELHEIM 7.5MG/5ML N21530 001 Jun 01, 2004

TABLET; ORAL

MELOXICAM

<u>AB</u>	ACTAVIS TOTOWA	<u>7.5MG</u>	<u>N77938</u>	<u>001</u>	Jul 19, 2006
<u>AB</u>		<u>15MG</u>	<u>N77938</u>	<u>002</u>	Jul 19, 2006
<u>AB</u>	APOTEX INC	<u>7.5MG</u>	<u>N77882</u>	<u>001</u>	Jul 20, 2006
<u>AB</u>		<u>15MG</u>	<u>N77882</u>	<u>002</u>	Jul 20, 2006
<u>AB</u>	AUROBINDO PHARMA	<u>7.5MG</u>	<u>N78008</u>	<u>001</u>	Oct 02, 2006
<u>AB</u>		<u>15MG</u>	<u>N78008</u>	<u>002</u>	Oct 02, 2006
<u>AB</u>	BRECKENRIDGE PHARM	<u>7.5MG</u>	<u>N77920</u>	<u>001</u>	Jul 19, 2006
<u>AB</u>		<u>15MG</u>	<u>N77920</u>	<u>002</u>	Jul 19, 2006
<u>AB</u>	CARACO	<u>7.5MG</u>	<u>N77937</u>	<u>001</u>	Jul 19, 2006
<u>AB</u>		<u>15MG</u>	<u>N77937</u>	<u>002</u>	Jul 19, 2006
<u>AB</u>	CARLSBAD	<u>7.5MG</u>	<u>N77918</u>	<u>001</u>	Dec 07, 2006
<u>AB</u>		<u>15MG</u>	<u>N77918</u>	<u>002</u>	Dec 07, 2006
<u>AB</u>	COREPHARMA	<u>7.5MG</u>	<u>N77930</u>	<u>001</u>	Jul 19, 2006
<u>AB</u>		<u>15MG</u>	<u>N77930</u>	<u>002</u>	Jul 19, 2006
<u>AB</u>	DR REDDYS LABS INC	<u>7.5MG</u>	<u>N77931</u>	<u>001</u>	Jul 25, 2006
<u>AB</u>		<u>15MG</u>	<u>N77931</u>	<u>002</u>	Jul 25, 2006
<u>AB</u>	GENPHARM	<u>7.5MG</u>	<u>N77934</u>	<u>001</u>	Jul 20, 2006
<u>AB</u>		<u>15MG</u>	<u>N77934</u>	<u>002</u>	Jul 20, 2006
<u>AB</u>	GLENMARK PHARMS LTD	<u>7.5MG</u>	<u>N77932</u>	<u>001</u>	Jul 19, 2006
<u>AB</u>		<u>15MG</u>	<u>N77932</u>	<u>002</u>	Jul 19, 2006
<u>AB</u>	LUPIN PHARMS	<u>7.5MG</u>	<u>N77944</u>	<u>001</u>	Jul 19, 2006
<u>AB</u>		<u>15MG</u>	<u>N77944</u>	<u>002</u>	Jul 19, 2006
<u>AB</u>	MUTUAL PHARM	<u>7.5MG</u>	<u>N77935</u>	<u>001</u>	Jul 19, 2006
<u>AB</u>		<u>15MG</u>	<u>N77935</u>	<u>002</u>	Jul 19, 2006
<u>AB</u>	MYLAN	<u>7.5MG</u>	<u>N77923</u>	<u>001</u>	Jul 19, 2006
<u>AB</u>		<u>15MG</u>	<u>N77923</u>	<u>002</u>	Jul 19, 2006
<u>AB</u>	PAR PHARM	<u>7.5MG</u>	<u>N77933</u>	<u>001</u>	Jul 19, 2006
<u>AB</u>		<u>15MG</u>	<u>N77933</u>	<u>002</u>	Jul 19, 2006
<u>AB</u>	RANBAXY	<u>7.5MG</u>	<u>N78039</u>	<u>001</u>	Dec 14, 2006
<u>AB</u>		<u>15MG</u>	<u>N78039</u>	<u>002</u>	Dec 14, 2006
<u>AB</u>	ROXANE	<u>7.5MG</u>	<u>N77925</u>	<u>001</u>	Jul 19, 2006
<u>AB</u>		<u>15MG</u>	<u>N77925</u>	<u>002</u>	Jul 19, 2006
<u>AB</u>	TARO	<u>7.5MG</u>	<u>N78102</u>	<u>001</u>	Nov 07, 2006
<u>AB</u>		<u>15MG</u>	<u>N78102</u>	<u>002</u>	Nov 07, 2006
<u>AB</u>	TEVA PHARMS	<u>7.5MG</u>	<u>N77936</u>	<u>001</u>	Jul 19, 2006
<u>AB</u>		<u>15MG</u>	<u>N77936</u>	<u>002</u>	Jul 19, 2006
<u>AB</u>	UNICHEM	<u>7.5MG</u>	<u>N77927</u>	<u>001</u>	Dec 20, 2006
<u>AB</u>		<u>15MG</u>	<u>N77927</u>	<u>002</u>	Dec 20, 2006
<u>AB</u>	WATSON LABS	<u>7.5MG</u>	<u>N77929</u>	<u>001</u>	Jul 19, 2006
<u>AB</u>		<u>15MG</u>	<u>N77929</u>	<u>002</u>	Jul 19, 2006

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 242 (of 370)

MELOXICAM

TABLET; ORAL

MELOXICAM

<u>AB</u>	ZYDUS PHARMS USA	<u>7.5MG</u>	<u>N77921</u>	<u>001</u>	Jul 19, 2006
<u>AB</u>		<u>15MG</u>	<u>N77921</u>	<u>002</u>	Jul 19, 2006

MOBIC

<u>AB</u>	BOEHRINGER INGELHEIM	<u>7.5MG</u>	<u>N20938</u>	<u>001</u>	Apr 13, 2000
<u>AB</u>	+	<u>15MG</u>	<u>N20938</u>	<u>002</u>	Aug 23, 2000

MELPHALAN

TABLET; ORAL

ALKERAN

+	GLAXOSMITHKLINE	2MG	N14691	002	
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MELPHALAN HYDROCHLORIDE

INJECTABLE; INJECTION

ALKERAN

+	GLAXOSMITHKLINE	EQ 50MG BASE/VIAL	N20207	001	Nov 18, 1992
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MEMANTINE HYDROCHLORIDE

SOLUTION; ORAL

NAMENDA

+	FOREST LABS	2MG/ML	N21627	001	Apr 18, 2005
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TABLET; ORAL

NAMENDA

	FOREST LABS	5MG	N21487	001	Oct 16, 2003
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+		10MG	N21487	002	Oct 16, 2003
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MENOTROPINS (FSH;LH)

INJECTABLE; IM-SC

REPRONEX

+	FERRING	75 IU/VIAL;75 IU/VIAL	N21047	001	Aug 27, 1999
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INJECTABLE; SUBCUTANEOUS

MENOPUR

+	FERRING	75 IU/VIAL;75 IU/VIAL	N21663	001	Oct 29, 2004
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MEPENZOLATE BROMIDE

TABLET; ORAL

CANTIL

+	SANOFI AVENTIS US	25MG	N10679	003	
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MEPERIDINE HYDROCHLORIDE

INJECTABLE; INJECTION

DEMEROL

<u>AP</u>	+	HOSPIRA	<u>25MG/ML</u>	<u>N21171</u>	<u>001</u>
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<u>AP</u>	+		<u>50MG/ML</u>	<u>N21171</u>	<u>002</u>
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<u>AP</u>	+		<u>75MG/ML</u>	<u>N21171</u>	<u>003</u>
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<u>AP</u>	+		<u>100MG/ML</u>	<u>N21171</u>	<u>004</u>
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MEPERIDINE HYDROCHLORIDE

<u>AP</u>		BAXTER HLTHCARE CORP	<u>25MG/ML</u>	<u>N80455</u>	<u>007</u>
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<u>AP</u>			<u>50MG/ML</u>	<u>N80455</u>	<u>008</u>
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<u>AP</u>			<u>75MG/ML</u>	<u>N80455</u>	<u>009</u>
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<u>AP</u>			<u>100MG/ML</u>	<u>N80455</u>	<u>010</u>
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<u>AP</u>		ELKINS SINN	<u>25MG/ML</u>	<u>N80445</u>	<u>001</u>
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<u>AP</u>			<u>50MG/ML</u>	<u>N80445</u>	<u>002</u>
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<u>AP</u>			<u>75MG/ML</u>	<u>N80445</u>	<u>003</u>
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<u>AP</u>			<u>100MG/ML</u>	<u>N80445</u>	<u>004</u>
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<u>AP</u>		WATSON LABS	<u>50MG/ML</u>	<u>N73444</u>	<u>001</u>	Mar 17, 1992
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<u>AP</u>			<u>100MG/ML</u>	<u>N73445</u>	<u>001</u>	Mar 17, 1992
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27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 243 (of 370)

MEPERIDINE HYDROCHLORIDE

INJECTABLE; INJECTION

MEPERIDINE HYDROCHLORIDE PRESERVATIVE FREE

<u>AP</u>	ASTRAZENECA	<u>10MG/ML</u>	<u>N81002</u>	<u>001</u>	Jul 30, 1993
<u>AP</u>	+ HOSPIRA	<u>10MG/ML</u>	<u>N88432</u>	<u>001</u>	Aug 16, 1984
<u>AP</u>	INTL MEDICATION	<u>10MG/ML</u>	<u>N81309</u>	<u>001</u>	Aug 30, 1993
<u>AP</u>	MALLINCKRODT	<u>10MG/ML</u>	<u>N40163</u>	<u>001</u>	May 12, 1997
<u>AP</u>	WATSON LABS	<u>10MG/ML</u>	<u>N73443</u>	<u>001</u>	Mar 17, 1992

SYRUP; ORAL

DEMEROL

<u>AA</u>	+ SANOFI AVENTIS US	<u>50MG/5ML</u>	<u>N05010</u>	<u>005</u>	
<u>AA</u>	<u>MEPERIDINE HYDROCHLORIDE</u>				
<u>AA</u>	ROXANE	<u>50MG/5ML</u>	<u>N88744</u>	<u>001</u>	Jan 30, 1985

TABLET; ORAL

DEMEROL

<u>AA</u>	+ SANOFI AVENTIS US	<u>50MG</u>	<u>N05010</u>	<u>001</u>	
<u>AA</u>	+	<u>100MG</u>	<u>N05010</u>	<u>004</u>	
	<u>MEPERIDINE HYDROCHLORIDE</u>				
<u>AA</u>	ACTAVIS TOTOWA	<u>50MG</u>	<u>N40331</u>	<u>001</u>	May 28, 1999
<u>AA</u>		<u>100MG</u>	<u>N40331</u>	<u>002</u>	May 28, 1999
<u>AA</u>	BARR	<u>50MG</u>	<u>N88639</u>	<u>001</u>	Jul 02, 1984
<u>AA</u>		<u>100MG</u>	<u>N88640</u>	<u>001</u>	Sep 19, 1984
<u>AA</u>	CARACO	<u>50MG</u>	<u>N40446</u>	<u>001</u>	Aug 08, 2002
<u>AA</u>		<u>100MG</u>	<u>N40446</u>	<u>002</u>	Aug 08, 2002
<u>AA</u>	DURAMED PHARMS BARR	<u>50MG</u>	<u>N40318</u>	<u>001</u>	Oct 05, 1999
<u>AA</u>		<u>100MG</u>	<u>N40318</u>	<u>002</u>	Oct 05, 1999
<u>AA</u>	MALLINCKRODT	<u>50MG</u>	<u>N40352</u>	<u>001</u>	Jun 13, 2000
<u>AA</u>		<u>100MG</u>	<u>N40352</u>	<u>002</u>	Jun 13, 2000
<u>AA</u>	MUTUAL PHARM	<u>50MG</u>	<u>N80448</u>	<u>001</u>	
<u>AA</u>		<u>100MG</u>	<u>N80448</u>	<u>002</u>	
<u>AA</u>	ROXANE	<u>50MG</u>	<u>N40110</u>	<u>001</u>	Mar 12, 1997
<u>AA</u>		<u>100MG</u>	<u>N40110</u>	<u>002</u>	Mar 12, 1997
<u>AA</u>	VINTAGE PHARMS	<u>50MG</u>	<u>N40191</u>	<u>001</u>	Dec 17, 1998
<u>AA</u>		<u>100MG</u>	<u>N40191</u>	<u>002</u>	Dec 17, 1998
<u>AA</u>	WATSON LABS	<u>50MG</u>	<u>N40186</u>	<u>001</u>	Jun 30, 1997
<u>AA</u>		<u>100MG</u>	<u>N40186</u>	<u>002</u>	Jun 30, 1997

MEPIVACAINE HYDROCHLORIDE

INJECTABLE; INJECTION

CARBOCAINE

<u>AP</u>	+ EASTMAN KODAK	<u>3%</u>	<u>N12125</u>	<u>003</u>	
<u>AP</u>	+ HOSPIRA	<u>1%</u>	<u>N12250</u>	<u>001</u>	
<u>AP</u>	+	<u>1.5%</u>	<u>N12250</u>	<u>005</u>	
<u>AP</u>	+	<u>2%</u>	<u>N12250</u>	<u>002</u>	

ISOCAINE HYDROCHLORIDE

<u>AP</u>	NOVOCOL	<u>3%</u>	<u>N80925</u>	<u>001</u>	
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MEPIVACAINE HYDROCHLORIDE

<u>AP</u>	GRAHAM CHEM	<u>3%</u>	<u>N83559</u>	<u>001</u>	
<u>AP</u>	WATSON LABS	<u>1%</u>	<u>N88769</u>	<u>001</u>	Nov 20, 1984
<u>AP</u>		<u>2%</u>	<u>N88770</u>	<u>001</u>	Nov 20, 1984

POLOCAINE

<u>AP</u>	ASTRAZENECA	<u>1%</u>	<u>N89407</u>	<u>001</u>	Dec 01, 1986
<u>AP</u>		<u>2%</u>	<u>N89410</u>	<u>001</u>	Dec 01, 1986
<u>AP</u>	DENTSPLY PHARM	<u>3%</u>	<u>N88653</u>	<u>001</u>	Aug 21, 1984

POLOCAINE-MPF

<u>AP</u>	ASTRAZENECA	<u>1%</u>	<u>N89406</u>	<u>001</u>	Dec 01, 1986
<u>AP</u>		<u>1.5%</u>	<u>N89408</u>	<u>001</u>	Dec 01, 1986
<u>AP</u>		<u>2%</u>	<u>N89409</u>	<u>001</u>	Dec 01, 1986

SCANDONEST PLAIN

<u>AP</u>	DEPROCO	<u>3%</u>	<u>N88387</u>	<u>001</u>	Oct 10, 1984
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27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 244 (of 370)

MEPROBAMATE

TABLET; ORAL

MEPROBAMATE

<u>AA</u>	+	WATSON LABS	<u>200MG</u>	<u>N83304</u>	<u>001</u>	
	+		400MG	N83308	001	

MEQUINOL; TRETINOINSOLUTION; TOPICAL
SOLAGE

+	BARRIER	2%;0.01%	N20922	001	Dec 10, 1999
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MERCAPTOPURINE

TABLET; ORAL

MERCAPTOPURINE

<u>AB</u>		PROMETHEUS LABS	<u>50MG</u>	<u>N40461</u>	<u>001</u>	Feb 11, 2004
<u>AB</u>		ROXANE	<u>50MG</u>	<u>N40528</u>	<u>001</u>	Feb 13, 2004

PURINETHOL

<u>AB</u>	+	TEVA	<u>50MG</u>	<u>N09053</u>	<u>002</u>	
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MERCAPTOPURINE ANHYDROUS

TABLET; ORAL

MERCAPTOPURINE

<u>AB</u>		MYLAN	<u>50MG</u>	<u>N40594</u>	<u>001</u>	Jul 01, 2005
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MEROPENEM

INJECTABLE; INJECTION

MERREM I.V.

+	ASTRAZENECA	500MG/VIAL	N50706	003	Jun 21, 1996
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+		1GM/VIAL	N50706	001	Jun 21, 1996
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MESALAMINE

CAPSULE, EXTENDED RELEASE; ORAL

PENTASA

SHIRE

250MG

N20049 001 May 10, 1993

+		500MG	N20049	002	Jul 08, 2004
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ENEMA; RECTAL

MESALAMINE

<u>AB</u>		PERRIGO NEW YORK	<u>4GM/60ML</u>	<u>N76751</u>	<u>001</u>	Sep 17, 2004
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<u>AB</u>		TEVA	<u>4GM/60ML</u>	<u>N76841</u>	<u>001</u>	Sep 30, 2004
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ROWASA

<u>AB</u>	+	ALAVEN PHARM	<u>4GM/60ML</u>	<u>N19618</u>	<u>001</u>	Dec 24, 1987
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SUPPOSITORY; RECTAL

CANASA

+	AXCAN SCANDIPHARM	1GM	N21252	002	Nov 05, 2004
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TABLET, DELAYED RELEASE; ORAL

ASACOL

+	PROCTER AND GAMBLE	400MG	N19651	001	Jan 31, 1992
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MESNA

INJECTABLE; INTRAVENOUS

MESNA

<u>AP</u>		ABRAXIS PHARM	<u>100MG/ML</u>	<u>N75811</u>	<u>001</u>	Apr 26, 2001
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<u>AP</u>		BEDFORD	<u>100MG/ML</u>	<u>N75739</u>	<u>001</u>	Jan 09, 2004
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<u>AP</u>		SICOR PHARMS	<u>100MG/ML</u>	<u>N75764</u>	<u>001</u>	Apr 27, 2001
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MESNEX

<u>AP</u>	+	BAXTER HLTHCARE	<u>100MG/ML</u>	<u>N19884</u>	<u>001</u>	Dec 30, 1988
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PRESCRIPTION DRUG PRODUCT LIST

3 - 245 (of 370)

MESNA

TABLET; ORAL

MESNEX

+ BAXTER HLTHCARE	400MG	N20855	001	Mar 21, 2002
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MESTRANOL; NORETHINDRONE

TABLET; ORAL-21

NORETHIN 1/50M-21

AB WATSON LABS	0.05MG;1MG	N71539	001	Apr 12, 1988
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NORETHINDRONE AND MESTRANOL

AB WATSON LABS	0.05MG;1MG	N70758	001	Jul 01, 1988
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NORINYL 1+50 21-DAY

AB + WATSON LABS	0.05MG;1MG	N13625	002	
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TABLET; ORAL-28

NORETHIN 1/50M-28

AB WATSON LABS	0.05MG;1MG	N71540	001	Apr 12, 1988
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NORETHINDRONE AND MESTRANOL

AB WATSON LABS	0.05MG;1MG	N70759	001	Jul 01, 1988
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NORINYL 1+50 28-DAY

AB WATSON LABS	0.05MG;1MG	N16659	001	
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ORTHO-NOVUM 1/50 28

AB ORTHO MCNEIL PHARM	0.05MG;1MG	N16709	001	
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METAPROTERENOL SULFATE

AEROSOL, METERED; INHALATION

ALUPENT

+ BOEHRINGER INGELHEIM	0.65MG/INH	N16402	001	
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SOLUTION; INHALATION

ALUPENT

AN + BOEHRINGER INGELHEIM	0.4%	N18761	002	Oct 10, 1986
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AN +	0.6%	N18761	001	Jun 30, 1983
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METAPROTERENOL SULFATE

AN DEY	0.4%	N71786	001	Aug 05, 1988
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AN	0.6%	N70804	001	Aug 17, 1987
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AN MORTON GROVE	0.4%	N75586	001	May 30, 2002
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AN	0.6%	N75586	002	May 30, 2002
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AN NEPHRON	0.4%	N71855	001	Jul 14, 1988
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AN	0.6%	N71726	001	Jul 14, 1988
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AN NOVEX	0.4%	N75402	001	Feb 28, 2001
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AN	0.6%	N75403	001	Feb 28, 2001
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SYRUP; ORAL

METAPROTERENOL SULFATE

AA + MORTON GROVE	10MG/5ML	N74702	001	Mar 24, 1997
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AA NOVEX	10MG/5ML	N75235	001	Jan 27, 2000
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AA SILARX	10MG/5ML	N73632	001	Jul 22, 1992
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TABLET; ORAL

METAPROTERENOL SULFATE

AB PAR PHARM	10MG	N72024	001	Jun 28, 1988
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AB +	20MG	N72025	001	Jun 28, 1988
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AB TEVA	10MG	N72519	001	Mar 30, 1990
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AB	20MG	N72520	001	Mar 30, 1990
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AB WATSON LABS	10MG	N73013	001	Jan 31, 1991
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AB	20MG	N72795	001	Jan 31, 1991
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METARAMINOL BITARTRATE

INJECTABLE; INJECTION

METARAMINOL BITARTRATE

+ ABRAXIS PHARM	EQ 10MG BASE/ML	N80722	001	
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PRESCRIPTION DRUG PRODUCT LIST

3 - 246 (of 370)

METAXALONE

TABLET; ORAL
SKELAXIN

+ KING PHARMS	800MG	N13217	003	Aug 30, 2002
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METFORMIN HYDROCHLORIDE

SOLUTION; ORAL
RIOMET

+ RANBAXY	500MG/5ML	N21591	001	Sep 11, 2003
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TABLET; ORAL

GLUCOPHAGE

<u>AB</u> BRISTOL MYERS SQUIBB	<u>500MG</u>	<u>N20357</u>	<u>001</u>	Mar 03, 1995
<u>AB</u>	<u>850MG</u>	<u>N20357</u>	<u>002</u>	Mar 03, 1995
<u>AB</u> +	<u>1GM</u>	<u>N20357</u>	<u>005</u>	Nov 05, 1998

METFORMIN HYDROCHLORIDE

<u>AB</u> ACTAVIS ELIZABETH	<u>500MG</u>	<u>N76033</u>	<u>001</u>	Jan 24, 2002
<u>AB</u>	<u>850MG</u>	<u>N76033</u>	<u>002</u>	Jan 24, 2002
<u>AB</u>	<u>1GM</u>	<u>N76033</u>	<u>003</u>	Jan 24, 2002
<u>AB</u> ALPHAPHARM	<u>500MG</u>	<u>N75969</u>	<u>001</u>	Jan 29, 2002
<u>AB</u>	<u>850MG</u>	<u>N75969</u>	<u>002</u>	Jan 29, 2002
<u>AB</u>	<u>1GM</u>	<u>N75969</u>	<u>003</u>	Jan 29, 2002
<u>AB</u> AMNEAL PHARM	<u>500MG</u>	<u>N77853</u>	<u>001</u>	Jul 28, 2006
<u>AB</u>	<u>850MG</u>	<u>N77853</u>	<u>002</u>	Jul 28, 2006
<u>AB</u>	<u>1GM</u>	<u>N77853</u>	<u>003</u>	Jul 28, 2006
<u>AB</u> ANDRX PHARMS	<u>500MG</u>	<u>N75961</u>	<u>001</u>	Jan 25, 2002
<u>AB</u>	<u>850MG</u>	<u>N75961</u>	<u>002</u>	Jan 25, 2002
<u>AB</u>	<u>1GM</u>	<u>N75961</u>	<u>003</u>	Jan 25, 2002
<u>AB</u> AUROBINDO	<u>500MG</u>	<u>N77095</u>	<u>001</u>	Jan 14, 2005
<u>AB</u>	<u>850MG</u>	<u>N77095</u>	<u>002</u>	Jan 14, 2005
<u>AB</u>	<u>1GM</u>	<u>N77095</u>	<u>003</u>	Jan 14, 2005
<u>AB</u> BARR	<u>500MG</u>	<u>N75971</u>	<u>001</u>	Jan 25, 2002
<u>AB</u>	<u>850MG</u>	<u>N75971</u>	<u>002</u>	Jan 25, 2002
<u>AB</u>	<u>1GM</u>	<u>N75971</u>	<u>003</u>	Jan 25, 2002
<u>AB</u> CARACO	<u>500MG</u>	<u>N75967</u>	<u>001</u>	Jan 29, 2002
<u>AB</u>	<u>850MG</u>	<u>N75967</u>	<u>002</u>	Jan 29, 2002
<u>AB</u>	<u>1GM</u>	<u>N75967</u>	<u>003</u>	Jan 29, 2002
<u>AB</u> DR REDDYS LABS INC	<u>500MG</u>	<u>N77787</u>	<u>001</u>	Aug 23, 2006
<u>AB</u>	<u>850MG</u>	<u>N77787</u>	<u>002</u>	Aug 23, 2006
<u>AB</u>	<u>1GM</u>	<u>N77787</u>	<u>003</u>	Aug 23, 2006
<u>AB</u> GENPHARM	<u>500MG</u>	<u>N75973</u>	<u>001</u>	Jan 25, 2002
<u>AB</u>	<u>850MG</u>	<u>N75973</u>	<u>002</u>	Jan 25, 2002
<u>AB</u>	<u>1GM</u>	<u>N75973</u>	<u>003</u>	Jan 25, 2002
<u>AB</u> GOLDLINE	<u>500MG</u>	<u>N75972</u>	<u>001</u>	Jan 24, 2002
<u>AB</u>	<u>625MG</u>	<u>N75972</u>	<u>005</u>	Jan 24, 2002
<u>AB</u>	<u>750MG</u>	<u>N75972</u>	<u>004</u>	Jan 24, 2002
<u>AB</u>	<u>850MG</u>	<u>N75972</u>	<u>002</u>	Jan 24, 2002
<u>AB</u>	<u>1GM</u>	<u>N75972</u>	<u>003</u>	Jan 24, 2002
<u>AB</u> INTERPHARM	<u>500MG</u>	<u>N77880</u>	<u>001</u>	Jun 05, 2006
<u>AB</u>	<u>850MG</u>	<u>N77880</u>	<u>002</u>	Jun 05, 2006
<u>AB</u>	<u>1GM</u>	<u>N77880</u>	<u>003</u>	Jun 05, 2006
<u>AB</u> MUTUAL PHARMA	<u>500MG</u>	<u>N76038</u>	<u>001</u>	Feb 21, 2002
<u>AB</u>	<u>850MG</u>	<u>N76038</u>	<u>002</u>	Feb 21, 2002
<u>AB</u>	<u>1GM</u>	<u>N76038</u>	<u>003</u>	Feb 21, 2002
<u>AB</u> MYLAN	<u>500MG</u>	<u>N75976</u>	<u>001</u>	Jan 24, 2002
<u>AB</u>	<u>850MG</u>	<u>N75976</u>	<u>002</u>	Jan 24, 2002
<u>AB</u>	<u>1GM</u>	<u>N75976</u>	<u>003</u>	Jan 24, 2002
<u>AB</u> SANDOZ	<u>500MG</u>	<u>N75965</u>	<u>001</u>	Jan 25, 2002
<u>AB</u>	<u>500MG</u>	<u>N75985</u>	<u>001</u>	Jan 25, 2002
<u>AB</u>	<u>850MG</u>	<u>N75965</u>	<u>002</u>	Jan 25, 2002
<u>AB</u>	<u>850MG</u>	<u>N75985</u>	<u>002</u>	Jan 25, 2002

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 247 (of 370)

METFORMIN HYDROCHLORIDE

TABLET; ORAL

METFORMIN HYDROCHLORIDE

<u>AB</u>	SANDOZ	<u>1GM</u>	<u>N75965</u>	<u>003</u>	Jan 25, 2002
<u>AB</u>		<u>1GM</u>	<u>N75985</u>	<u>003</u>	Jan 25, 2002
<u>AB</u>	TEVA	<u>500MG</u>	<u>N75978</u>	<u>001</u>	Jan 25, 2002
<u>AB</u>		<u>500MG</u>	<u>N76328</u>	<u>001</u>	Dec 16, 2002
<u>AB</u>		<u>850MG</u>	<u>N75978</u>	<u>002</u>	Jan 25, 2002
<u>AB</u>		<u>850MG</u>	<u>N76328</u>	<u>002</u>	Dec 16, 2002
<u>AB</u>		<u>1GM</u>	<u>N75978</u>	<u>003</u>	Nov 05, 2002
<u>AB</u>		<u>1GM</u>	<u>N76328</u>	<u>003</u>	Dec 16, 2002
<u>AB</u>	TORPHARM	<u>500MG</u>	<u>N75984</u>	<u>001</u>	Apr 23, 2002
<u>AB</u>		<u>850MG</u>	<u>N75984</u>	<u>002</u>	Apr 23, 2002
<u>AB</u>		<u>1GM</u>	<u>N75984</u>	<u>003</u>	Apr 23, 2002
<u>AB</u>	WATSON LABS	<u>500MG</u>	<u>N75979</u>	<u>001</u>	Jan 24, 2002
<u>AB</u>		<u>850MG</u>	<u>N75979</u>	<u>002</u>	Jan 24, 2002
<u>AB</u>		<u>1GM</u>	<u>N75979</u>	<u>003</u>	Jan 24, 2002
<u>AB</u>	ZENITH GOLDLINE	<u>500MG</u>	<u>N75975</u>	<u>001</u>	Jan 24, 2002
<u>AB</u>		<u>625MG</u>	<u>N75975</u>	<u>004</u>	Jan 24, 2002
<u>AB</u>		<u>750MG</u>	<u>N75975</u>	<u>005</u>	Jan 24, 2002
<u>AB</u>		<u>850MG</u>	<u>N75975</u>	<u>002</u>	Jan 24, 2002
<u>AB</u>		<u>1GM</u>	<u>N75975</u>	<u>003</u>	Jan 24, 2002
<u>AB</u>	ZYDUS PHARMS USA	<u>500MG</u>	<u>N77064</u>	<u>001</u>	Apr 18, 2005
<u>AB</u>		<u>850MG</u>	<u>N77064</u>	<u>002</u>	Apr 18, 2005
<u>AB</u>		<u>1GM</u>	<u>N77064</u>	<u>003</u>	Apr 18, 2005

TABLET, EXTENDED RELEASE; ORAL

FORTAMET

BX	ANDRX LABS LLC	500MG	N21574	001	Apr 27, 2004
BX +		1GM	N21574	002	Apr 27, 2004

GLUCOPHAGE XR

<u>AB</u>	BRISTOL MYERS SQUIBB	<u>500MG</u>	<u>N21202</u>	<u>001</u>	Oct 13, 2000
<u>AB</u> +		<u>750MG</u>	<u>N21202</u>	<u>004</u>	Apr 11, 2003

GLUMETZA

BX	DEPOMED INC	500MG	N21748	001	Jun 03, 2005
BX +		1GM	N21748	002	Jun 03, 2005

METFORMIN HYDROCHLORIDE

<u>AB</u>	ACTAVIS ELIZABETH	<u>500MG</u>	<u>N76450</u>	<u>001</u>	Oct 01, 2004
<u>AB</u>		<u>750MG</u>	<u>N76878</u>	<u>001</u>	Apr 13, 2005
<u>AB</u>	ANDRX PHARMS	<u>500MG</u>	<u>N76172</u>	<u>001</u>	Jun 16, 2004
<u>AB</u>		<u>750MG</u>	<u>N76869</u>	<u>001</u>	Apr 12, 2005
<u>AB</u>	APOTEX INC	<u>500MG</u>	<u>N76706</u>	<u>001</u>	Dec 14, 2004
<u>AB</u>		<u>750MG</u>	<u>N76706</u>	<u>002</u>	Dec 29, 2005
<u>AB</u>	BARR	<u>500MG</u>	<u>N76496</u>	<u>001</u>	Nov 25, 2005
<u>AB</u>		<u>750MG</u>	<u>N76863</u>	<u>001</u>	Oct 14, 2004
<u>AB</u>	COBALT	<u>500MG</u>	<u>N76818</u>	<u>001</u>	Dec 14, 2004
<u>AB</u>	IMPAX LABS	<u>500MG</u>	<u>N76249</u>	<u>001</u>	Jul 30, 2004
<u>AB</u>		<u>750MG</u>	<u>N76985</u>	<u>001</u>	Sep 13, 2005
<u>AB</u>	IVAX PHARMS	<u>500MG</u>	<u>N76545</u>	<u>001</u>	Dec 01, 2003
<u>AB</u>	MUTUAL PHARM	<u>500MG</u>	<u>N77124</u>	<u>001</u>	Dec 21, 2005
<u>AB</u>	MYLAN	<u>500MG</u>	<u>N76650</u>	<u>001</u>	Sep 13, 2005
<u>AB</u>		<u>750MG</u>	<u>N77113</u>	<u>001</u>	Sep 08, 2005
<u>AB</u>	NOSTRUM	<u>500MG</u>	<u>N76756</u>	<u>001</u>	Jul 26, 2006
<u>AB</u>	RANBAXY	<u>500MG</u>	<u>N76413</u>	<u>001</u>	Jun 18, 2004
<u>AB</u>		<u>750MG</u>	<u>N77211</u>	<u>001</u>	Jun 29, 2005
<u>AB</u>	SANDOZ	<u>500MG</u>	<u>N76223</u>	<u>001</u>	Dec 14, 2004
<u>AB</u>		<u>500MG</u>	<u>N76873</u>	<u>001</u>	Dec 14, 2004
<u>AB</u>	SUN PHARM INDS (IN)	<u>500MG</u>	<u>N77336</u>	<u>001</u>	Feb 09, 2006
<u>AB</u>		<u>750MG</u>	<u>N77336</u>	<u>002</u>	Feb 09, 2006
<u>AB</u>	TEVA	<u>500MG</u>	<u>N76269</u>	<u>001</u>	Jun 18, 2004
<u>AB</u>		<u>750MG</u>	<u>N76864</u>	<u>001</u>	Apr 12, 2005

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 248 (of 370)

METFORMIN HYDROCHLORIDE

TABLET, EXTENDED RELEASE; ORAL

METFORMIN HYDROCHLORIDE

<u>AB</u>	ZYDUS PHARMS USA	<u>500MG</u>	<u>N77060</u>	<u>001</u>	Apr 20, 2005
<u>AB</u>		<u>750MG</u>	<u>N77078</u>	<u>001</u>	Apr 21, 2005

METFORMIN HYDROCHLORIDE; ROSIGLITAZONE MALEATE

TABLET; ORAL

AVANDAMET

SB PHARMC0

500MG;EQ 1MG BASE

N21410 001 Oct 10, 2002

500MG;EQ 2MG BASE

N21410 002 Oct 10, 2002

500MG;EQ 4MG BASE

N21410 003 Oct 10, 2002

1GM;EQ 2MG BASE

N21410 004 Aug 25, 2003

+

1GM;EQ 4MG BASE

N21410 005 Aug 25, 2003

METFORMIN; PIOGLITAZONE HYDROCHLORIDE

TABLET; ORAL

ACTOPLUS MET

TAKEDA GLOBAL

500MG;EQ 15MG BASE

N21842 001 Aug 29, 2005

+

850MG;EQ 15MG BASE

N21842 002 Aug 29, 2005

METHACHOLINE CHLORIDE

FOR SOLUTION; INHALATION

PROVOCHOLINE

+ METHAPHARM

100MG/VIAL

N19193 001 Oct 31, 1986

METHADONE HYDROCHLORIDE

CONCENTRATE; ORAL

METHADONE HYDROCHLORIDE

<u>AA</u>	ROXANE	<u>10MG/ML</u>	<u>N40180</u>	<u>001</u>	Apr 30, 1998
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<u>AA</u>	UDL	<u>10MG/ML</u>	<u>N40088</u>	<u>001</u>	Nov 30, 1994
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METHADONE HYDROCHLORIDE INTENSOL

<u>AA</u>	ROXANE	<u>10MG/ML</u>	<u>N89897</u>	<u>001</u>	Sep 06, 1988
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METHADOSE

<u>AA</u>	+ MALLINCKRODT	<u>10MG/ML</u>	<u>N17116</u>	<u>002</u>	
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INJECTABLE; INJECTION

DOLOPHINE HYDROCHLORIDE

+ XANODYNE PHARM

10MG/ML

N21624 001

POWDER; FOR RX COMPOUNDING

METHADONE HYDROCHLORIDE

MALLINCKRODT

50GM/BOT

N06383 002

100GM/BOT

N06383 003

500GM/BOT

N06383 004

SOLUTION; ORAL

METHADONE HYDROCHLORIDE

+ ROXANE

5MG/5ML

N87393 001

+

10MG/5ML

N87997 001 Aug 30, 1982

TABLET; ORAL

DOLOPHINE HYDROCHLORIDE

<u>AA</u>	+ ROXANE	<u>5MG</u>	<u>N06134</u>	<u>002</u>	
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<u>AA</u>	+	<u>10MG</u>	<u>N06134</u>	<u>010</u>	
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METHADONE HYDROCHLORIDE

<u>AA</u>	MALLINCKRODT	<u>5MG</u>	<u>N40517</u>	<u>001</u>	Apr 27, 2004
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<u>AA</u>		<u>10MG</u>	<u>N40517</u>	<u>002</u>	Apr 27, 2004
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<u>AA</u>		<u>40MG</u>	<u>N77142</u>	<u>001</u>	Jul 12, 2005
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<u>AA</u>	ROXANE	<u>5MG</u>	<u>N88108</u>	<u>001</u>	Mar 08, 1983
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<u>AA</u>		<u>10MG</u>	<u>N88109</u>	<u>001</u>	Mar 08, 1983
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<u>AA</u>	+	<u>40MG</u>	<u>N17058</u>	<u>001</u>	
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<u>AA</u>		<u>40MG</u>	<u>N74081</u>	<u>001</u>	Apr 28, 1995
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27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 249 (of 370)

METHADONE HYDROCHLORIDE

TABLET; ORAL

METHADONE HYDROCHLORIDE

<u>AA</u>	SANDOZ	<u>10MG</u>	<u>N40241</u>	<u>002</u>	May 29, 1998
<u>AA</u>		<u>40MG</u>	<u>N75082</u>	<u>001</u>	Mar 25, 1998
	<u>METHADOSE</u>				
<u>AA</u>	MALLINCKRODT	<u>5MG</u>	<u>N40050</u>	<u>001</u>	Apr 15, 1993
<u>AA</u>		<u>10MG</u>	<u>N40050</u>	<u>002</u>	Apr 15, 1993
<u>AA</u>		<u>40MG</u>	<u>N74184</u>	<u>001</u>	Apr 29, 1993

METHAMPHETAMINE HYDROCHLORIDE

TABLET; ORAL

DESOXYN

+ OVATION PHARMS 5MG N05378 002

METHAZOLAMIDE

TABLET; ORAL

METHAZOLAMIDE

<u>AB</u>	MIKART	<u>25MG</u>	<u>N40062</u>	<u>001</u>	Jan 27, 1994
<u>AB</u>	+	<u>50MG</u>	<u>N40062</u>	<u>002</u>	Jan 27, 1994
<u>AB</u>	SANDOZ	<u>25MG</u>	<u>N40036</u>	<u>001</u>	Jun 30, 1993
<u>AB</u>		<u>25MG</u>	<u>N40102</u>	<u>001</u>	Aug 28, 1996
<u>AB</u>		<u>50MG</u>	<u>N40036</u>	<u>002</u>	Jun 30, 1993
<u>AB</u>		<u>50MG</u>	<u>N40102</u>	<u>002</u>	Aug 28, 1996
<u>AB</u>	TEVA PHARMS	<u>25MG</u>	<u>N40001</u>	<u>001</u>	Jun 30, 1993
<u>AB</u>		<u>50MG</u>	<u>N40001</u>	<u>002</u>	Jun 30, 1993

METHENAMINE HIPPURATE

TABLET; ORAL

HIPREXAB + SANOFI AVENTIS US 1GM N17681 001METHENAMINE HIPPURATE

<u>AB</u>	COREPHARMA	<u>1GM</u>	<u>N76411</u>	<u>001</u>	Jun 20, 2003
	<u>UREX</u>				
<u>AB</u>	VATRING PHARMS	<u>1GM</u>	<u>N16151</u>	<u>001</u>	

METHIMAZOLE

TABLET; ORAL

METHIMAZOLE

<u>AB</u>	CEDAR PHARMS	<u>5MG</u>	<u>N40547</u>	<u>001</u>	Feb 18, 2005
<u>AB</u>		<u>10MG</u>	<u>N40547</u>	<u>002</u>	Feb 18, 2005
		15MG	N40619	003	Jul 12, 2005
	+	20MG	N40547	004	Feb 18, 2005
<u>AB</u>	GENPHARM	<u>5MG</u>	<u>N40350</u>	<u>001</u>	Mar 29, 2000
<u>AB</u>	+	<u>10MG</u>	<u>N40350</u>	<u>002</u>	Mar 29, 2000
<u>AB</u>	SANDOZ	<u>5MG</u>	<u>N40411</u>	<u>001</u>	Mar 27, 2001
<u>AB</u>		<u>10MG</u>	<u>N40411</u>	<u>002</u>	Mar 27, 2001
	<u>TAPAZOLE</u>				
<u>AB</u>	KING PHARMS	<u>5MG</u>	<u>N40320</u>	<u>001</u>	Mar 31, 2000
<u>AB</u>		<u>10MG</u>	<u>N40320</u>	<u>002</u>	Mar 31, 2000

METHOCARBAMOL

INJECTABLE; INJECTION

METHOCARBAMOLAP WATSON LABS 100MG/ML N86459 001ROBAXINAP + BAXTER HLTHCARE CORP 100MG/ML N11790 001

PRESCRIPTION DRUG PRODUCT LIST

3 - 250 (of 370)

METHOCARBAMOL

TABLET; ORAL

METHOCARBAMOL

<u>AA</u>	IMPAX LABS	<u>500MG</u>	<u>N84927</u>	<u>001</u>	
<u>AA</u>		<u>750MG</u>	<u>N84928</u>	<u>001</u>	
<u>AA</u>	LANNETT	<u>500MG</u>	<u>N84756</u>	<u>002</u>	Mar 31, 2003
<u>AA</u>		<u>750MG</u>	<u>N84756</u>	<u>001</u>	
<u>AA</u>	PAR PHARM	<u>500MG</u>	<u>N86989</u>	<u>001</u>	
<u>AA</u>		<u>750MG</u>	<u>N86988</u>	<u>001</u>	
<u>AA</u>	SANDOZ	<u>500MG</u>	<u>N84616</u>	<u>001</u>	
<u>AA</u>		<u>500MG</u>	<u>N87283</u>	<u>001</u>	
<u>AA</u>		<u>750MG</u>	<u>N84615</u>	<u>001</u>	
<u>AA</u>		<u>750MG</u>	<u>N87282</u>	<u>001</u>	
<u>AA</u>	TABLICAPS	<u>500MG</u>	<u>N84846</u>	<u>001</u>	
<u>AA</u>	VINTAGE PHARMS	<u>500MG</u>	<u>N40489</u>	<u>001</u>	Jan 29, 2003
<u>AA</u>		<u>750MG</u>	<u>N40489</u>	<u>002</u>	Jan 29, 2003
<u>AA</u>	WATSON LABS	<u>500MG</u>	<u>N84277</u>	<u>001</u>	
<u>AA</u>		<u>500MG</u>	<u>N85180</u>	<u>001</u>	
<u>AA</u>		<u>750MG</u>	<u>N84276</u>	<u>002</u>	
<u>AA</u>		<u>750MG</u>	<u>N85192</u>	<u>001</u>	
<u>AA</u>	WEST WARD	<u>500MG</u>	<u>N85159</u>	<u>001</u>	
<u>AA</u>		<u>750MG</u>	<u>N85123</u>	<u>001</u>	
<u>ROBAXIN</u>					
<u>AA</u>	+ SCHWARZ PHARMA	<u>500MG</u>	<u>N11011</u>	<u>004</u>	
<u>ROBAXIN-750</u>					
<u>AA</u>	+ SCHWARZ PHARMA	<u>750MG</u>	<u>N11011</u>	<u>006</u>	

METHOHEXITAL SODIUM

INJECTABLE; INJECTION

BREVITAL SODIUM

+	KING PHARMS	500MG/VIAL	N11559	001	
+		2.5GM/VIAL	N11559	002	
+		5GM/VIAL	N11559	003	

METHOTREXATE SODIUM

INJECTABLE; INJECTION

METHOTREXATE

<u>AP</u>	+ ABRAXIS PHARM	<u>EQ 50MG BASE/2ML (25MG/ML)</u>	<u>N40263</u>	<u>001</u>	Feb 26, 1999
<u>AP</u>	+	<u>EQ 250MG BASE/10ML (25MG/ML)</u>	<u>N40263</u>	<u>002</u>	Feb 26, 1999
<u>AP</u>	+ MAYNE PHARMA USA	<u>EQ 50MG BASE/2ML (25 MG/ML)</u>	<u>N11719</u>	<u>010</u>	Dec 15, 2004
METHOTREXATE PRESERVATIVE FREE					
+	BEDFORD	EQ 1GM BASE/VIAL	N40632	001	Aug 12, 2005
+	MAYNE PHARMA USA	EQ 1GM BASE/40ML (25MG/ML)	N11719	012	Apr 13, 2005
<u>METHOTREXATE SODIUM</u>					
<u>AP</u>	+ BEDFORD	<u>EQ 50MG BASE/2ML (25MG/ML)</u>	<u>N89340</u>	<u>001</u>	Sep 16, 1986
+		EQ 100MG BASE/4ML (25MG/ML)	N89341	001	Sep 16, 1986
+		EQ 200MG BASE/8ML (25MG/ML)	N89342	001	Sep 16, 1986
+		EQ 250MG BASE/10ML (25 MG/ML)	N89343	001	Sep 16, 1986

TABLET; ORAL

METHOTREXATE SODIUM

<u>AB</u>	BARR	<u>EQ 2.5MG BASE</u>	<u>N81099</u>	<u>001</u>	Oct 15, 1990
<u>AB</u>	DURAMED PHARMS BARR	<u>EQ 2.5MG BASE</u>	<u>N40233</u>	<u>001</u>	Jun 17, 1999
<u>AB</u>	MYLAN	<u>EQ 2.5MG BASE</u>	<u>N81235</u>	<u>001</u>	May 15, 1992
<u>AB</u>	ROXANE	<u>EQ 2.5MG BASE</u>	<u>N40054</u>	<u>001</u>	Aug 01, 1994
<u>AB</u>	+ STADA PHARMS	<u>EQ 2.5MG BASE</u>	<u>N08085</u>	<u>002</u>	
TREXALL					
	BARR	EQ 5MG BASE	N40385	001	Mar 21, 2001
		EQ 7.5MG BASE	N40385	002	Mar 21, 2001
		EQ 10MG BASE	N40385	003	Mar 21, 2001
+		EQ 15MG BASE	N40385	004	Mar 21, 2001

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 251 (of 370)

METHOXSALENCAPSULE; ORAL
8-MOP

+ VALEANT PHARM INTL	10MG	N09048	001	
OXSORALEN-ULTRA				
+ VALEANT PHARM INTL	10MG	N19600	001	Oct 30, 1986

INJECTABLE; INJECTION
UVADEX

+ THERAKOS	0.02MG/ML	N20969	001	Feb 25, 1999
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LOTION; TOPICAL
OXSORALEN

+ VALEANT PHARM INTL	1%	N09048	002	
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METHSCOPOLAMINE BROMIDE

TABLET; ORAL

METHSCOPOLAMINE BROMIDE

<u>AA</u>	BOCA PHARMA	<u>2.5MG</u>	<u>N40624</u>	<u>001</u>	Dec 28, 2006
<u>AA</u>		<u>5MG</u>	<u>N40624</u>	<u>002</u>	Dec 28, 2006

PAMINE

<u>AA</u>	+ BRADLEY PHARMS	<u>2.5MG</u>	<u>N08848</u>	<u>001</u>	
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PAMINE FORTE

<u>AA</u>	+ BRADLEY PHARMS	<u>5MG</u>	<u>N08848</u>	<u>002</u>	Mar 25, 2003
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METHSUXIMIDECAPSULE; ORAL
CELONTIN

PARKE DAVIS	150MG	N10596	007	
+	300MG	N10596	008	

METHYCLOTHIAZIDE

TABLET; ORAL

ENDURON

<u>AB</u>	ABBOTT	<u>2.5MG</u>	<u>N12524</u>	<u>001</u>	
<u>AB</u>	+	<u>5MG</u>	<u>N12524</u>	<u>004</u>	

METHYCLOTHIAZIDE

<u>AB</u>	IVAX PHARMS	<u>2.5MG</u>	<u>N87913</u>	<u>001</u>	Jun 03, 1982
<u>AB</u>	MYLAN	<u>5MG</u>	<u>N87672</u>	<u>001</u>	Aug 17, 1982
<u>AB</u>	SANDOZ	<u>2.5MG</u>	<u>N89835</u>	<u>001</u>	Aug 18, 1988
<u>AB</u>		<u>5MG</u>	<u>N89837</u>	<u>001</u>	Aug 18, 1988
<u>AB</u>	WATSON LABS	<u>5MG</u>	<u>N88724</u>	<u>001</u>	Sep 06, 1984

METHYL AMINOLEVULINATE HYDROCHLORIDECREAM; TOPICAL
METVIXIA

+ PHOTOCURE ASA	EQ 16.8% BASE	N21415	001	Jul 27, 2004
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METHYLDOPA

TABLET; ORAL

METHYLDOPA

<u>AB</u>	ACCORD HLTH	<u>250MG</u>	<u>N70084</u>	<u>001</u>	Oct 15, 1985
<u>AB</u>		<u>500MG</u>	<u>N70085</u>	<u>001</u>	Oct 15, 1985
<u>AB</u>	IVAX PHARMS	<u>250MG</u>	<u>N70098</u>	<u>001</u>	Feb 20, 1986
<u>AB</u>		<u>500MG</u>	<u>N70343</u>	<u>001</u>	Feb 20, 1986
<u>AB</u>	MYLAN	<u>250MG</u>	<u>N70075</u>	<u>001</u>	Apr 18, 1985
<u>AB</u>	+	<u>500MG</u>	<u>N70076</u>	<u>001</u>	Apr 18, 1985
<u>AB</u>	PAR PHARM	<u>125MG</u>	<u>N70535</u>	<u>001</u>	Jan 02, 1987
<u>AB</u>		<u>250MG</u>	<u>N70536</u>	<u>001</u>	Jan 02, 1987
<u>AB</u>		<u>500MG</u>	<u>N70537</u>	<u>001</u>	Jan 02, 1987
<u>AB</u>	PLIVA	<u>125MG</u>	<u>N72126</u>	<u>001</u>	Jul 07, 1988

PRESCRIPTION DRUG PRODUCT LIST

3 - 252 (of 370)

METHYLDOPA

TABLET; ORAL

METHYLDOPA

<u>AB</u>	PLIVA	<u>250MG</u>	<u>N72127</u>	<u>001</u>	Jul 07, 1988
<u>AB</u>		<u>500MG</u>	<u>N72128</u>	<u>001</u>	Jul 07, 1988
<u>AB</u>	SANDOZ	<u>125MG</u>	<u>N71700</u>	<u>001</u>	Mar 02, 1988
<u>AB</u>		<u>250MG</u>	<u>N18934</u>	<u>001</u>	Jun 29, 1984
<u>AB</u>		<u>500MG</u>	<u>N18934</u>	<u>002</u>	Jun 29, 1984
<u>AB</u>	WATSON LABS	<u>250MG</u>	<u>N70703</u>	<u>001</u>	Jun 06, 1986
<u>AB</u>		<u>500MG</u>	<u>N70625</u>	<u>001</u>	Jun 06, 1986

METHYLDOPATE HYDROCHLORIDE

INJECTABLE; INJECTION

METHYLDOPATE HYDROCHLORIDE

<u>AP</u>	HOSPIRA	<u>50MG/ML</u>	<u>N70698</u>	<u>001</u>	Jun 15, 1987
<u>AP</u>		<u>50MG/ML</u>	<u>N70699</u>	<u>001</u>	Jun 15, 1987
<u>AP</u>	+ LUITPOLD	<u>50MG/ML</u>	<u>N71279</u>	<u>001</u>	Oct 02, 1987
<u>AP</u>	SICOR PHARMS	<u>50MG/ML</u>	<u>N72974</u>	<u>001</u>	Nov 22, 1991

METHYLERGONOVINE MALEATE

INJECTABLE; INJECTION

METHERGINE

+ NOVARTIS 0.2MG/ML

N06035 004

TABLET; ORAL

METHERGINE

+ NOVARTIS 0.2MG

N06035 003

METHYLPHENIDATE

FILM, EXTENDED RELEASE; TRANSDERMAL

DAYTRANA

+ SHIRE 10MG/9HR (1.1MG/HR)

N21514 001 Apr 06, 2006

+ 15MG/9HR (1.6MG/HR)

N21514 002 Apr 06, 2006

+ 20MG/9HR (2.2MG/HR)

N21514 003 Apr 06, 2006

+ 30MG/9HR (3.3MG/HR)

N21514 004 Apr 06, 2006

METHYLPHENIDATE HYDROCHLORIDE

CAPSULE, EXTENDED RELEASE; ORAL

METADATE CD

BX	UCB INC	10MG	N21259	003	May 27, 2003
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BX		20MG	N21259	001	Apr 03, 2001
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BX	+	30MG	N21259	002	Jun 19, 2003
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BX		40MG	N21259	004	Feb 19, 2006
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		50MG	N21259	005	Feb 19, 2006
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	+	60MG	N21259	006	Feb 19, 2006
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RITALIN LA

BX	NOVARTIS	10MG	N21284	004	Apr 10, 2004
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BX		20MG	N21284	001	Jun 05, 2002
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BX		30MG	N21284	002	Jun 05, 2002
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BX	+	40MG	N21284	003	Jun 05, 2002
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SOLUTION; ORAL

METHYLIN

+ MALLINCKRODT 5MG/5ML

N21419 001 Dec 19, 2002

+ 10MG/5ML

N21419 002 Dec 19, 2002

TABLET; ORAL

METHYLPHENIDATE HYDROCHLORIDE

<u>AB</u>	ACTAVIS ELIZABETH	<u>5MG</u>	<u>N40321</u>	<u>001</u>	Feb 05, 2002
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<u>AB</u>		<u>10MG</u>	<u>N40321</u>	<u>002</u>	Feb 05, 2002
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<u>AB</u>		<u>20MG</u>	<u>N40321</u>	<u>003</u>	Feb 05, 2002
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<u>AB</u>	MALLINCKRODT	<u>5MG</u>	<u>N40300</u>	<u>001</u>	Nov 27, 1998
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27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 253 (of 370)

METHYLPHENIDATE HYDROCHLORIDE

TABLET; ORAL

METHYLPHENIDATE HYDROCHLORIDE

<u>AB</u>	MALLINCKRODT	<u>10MG</u>	<u>N40300</u>	<u>002</u>	Nov 27, 1998
<u>AB</u>		<u>20MG</u>	<u>N40300</u>	<u>003</u>	Nov 27, 1998
<u>AB</u>	UCB INC	<u>5MG</u>	<u>N86429</u>	<u>001</u>	
<u>AB</u>		<u>10MG</u>	<u>N85799</u>	<u>001</u>	
<u>AB</u>		<u>20MG</u>	<u>N86428</u>	<u>001</u>	
<u>AB</u>	WATSON LABS	<u>5MG</u>	<u>N40220</u>	<u>001</u>	Aug 29, 1997
<u>AB</u>		<u>10MG</u>	<u>N40220</u>	<u>002</u>	Aug 29, 1997
<u>AB</u>		<u>20MG</u>	<u>N40220</u>	<u>003</u>	Aug 29, 1997
	<u>RITALIN</u>				
<u>AB</u>	NOVARTIS	<u>5MG</u>	<u>N10187</u>	<u>003</u>	
<u>AB</u>		<u>10MG</u>	<u>N10187</u>	<u>006</u>	
<u>AB</u>	+	<u>20MG</u>	<u>N10187</u>	<u>010</u>	

TABLET, CHEWABLE; ORAL

METHYLIN

	MALLINCKRODT	2.5MG	N21475	001	Apr 15, 2003
		5MG	N21475	002	Apr 15, 2003
	+	10MG	N21475	003	Apr 15, 2003

TABLET, EXTENDED RELEASE; ORAL

CONCERTA

	ALZA	18MG	N21121	001	Aug 01, 2000
		27MG	N21121	004	Apr 01, 2002
		36MG	N21121	002	Aug 01, 2000
	+	54MG	N21121	003	Dec 08, 2000

METADATE ER

<u>AB</u>	UCB INC	<u>10MG</u>	<u>N40306</u>	<u>001</u>	Oct 20, 1999
<u>AB</u>	+	<u>20MG</u>	<u>N89601</u>	<u>001</u>	Jun 01, 1988

METHYLIN ER

<u>AB</u>	MALLINCKRODT	<u>10MG</u>	<u>N75629</u>	<u>001</u>	May 09, 2000
<u>AB</u>		<u>20MG</u>	<u>N75629</u>	<u>002</u>	May 09, 2000

METHYLPHENIDATE HYDROCHLORIDE

<u>AB</u>	ACTAVIS ELIZABETH	<u>20MG</u>	<u>N75450</u>	<u>001</u>	Dec 21, 2001
<u>AB</u>	WATSON LABS	<u>20MG</u>	<u>N40410</u>	<u>001</u>	Feb 09, 2001
	<u>RITALIN-SR</u>				
<u>AB</u>	NOVARTIS	<u>20MG</u>	<u>N18029</u>	<u>001</u>	Mar 30, 1982

METHYLPREDNISOLONE

TABLET; ORAL

MEDROL

<u>AB</u>	PHARMACIA AND UPJOHN	<u>4MG</u>	<u>N11153</u>	<u>001</u>	
<u>AB</u>		<u>8MG</u>	<u>N11153</u>	<u>004</u>	
<u>AB</u>		<u>16MG</u>	<u>N11153</u>	<u>003</u>	
<u>AB</u>		<u>24MG</u>	<u>N11153</u>	<u>005</u>	
<u>AB</u>	+	<u>32MG</u>	<u>N11153</u>	<u>006</u>	

METHYLPREDNISOLONE

<u>AB</u>	DURAMED PHARMS BARR	<u>4MG</u>	<u>N88497</u>	<u>001</u>	Feb 21, 1984
<u>AB</u>	JUBILANT PHARMS	<u>4MG</u>	<u>N40189</u>	<u>001</u>	Oct 31, 1997
<u>AB</u>		<u>8MG</u>	<u>N40189</u>	<u>002</u>	Oct 31, 1997
<u>AB</u>	PAR PHARM	<u>16MG</u>	<u>N89207</u>	<u>001</u>	Apr 25, 1988
<u>AB</u>		<u>24MG</u>	<u>N89208</u>	<u>001</u>	Apr 25, 1988
<u>AB</u>		<u>32MG</u>	<u>N89209</u>	<u>001</u>	Apr 25, 1988
<u>AB</u>	SANDOZ	<u>4MG</u>	<u>N40194</u>	<u>001</u>	Oct 31, 1997
<u>AB</u>	VINTAGE PHARMS	<u>4MG</u>	<u>N40183</u>	<u>001</u>	Dec 22, 1998
<u>AB</u>	WATSON LABS	<u>4MG</u>	<u>N40232</u>	<u>001</u>	Oct 16, 1997

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 254 (of 370)

METHYLPREDNISOLONE ACETATE

INJECTABLE; INJECTION

DEPO-MEDROL

<u>AB</u>	+	PHARMACIA AND UPJOHN	<u>40MG/ML</u>	<u>N11757</u>	<u>001</u>	
<u>AB</u>	+		<u>80MG/ML</u>	<u>N11757</u>	<u>004</u>	
	+		20MG/ML	N11757	002	

METHYLPREDNISOLONE ACETATE

<u>AB</u>		SICOR PHARMS	<u>40MG/ML</u>	<u>N40557</u>	<u>001</u>	Feb 23, 2005
<u>AB</u>			<u>40MG/ML</u>	<u>N40620</u>	<u>001</u>	Oct 27, 2006
<u>AB</u>			<u>80MG/ML</u>	<u>N40557</u>	<u>002</u>	Feb 23, 2005
<u>AB</u>			<u>80MG/ML</u>	<u>N40620</u>	<u>002</u>	Oct 27, 2006

METHYLPREDNISOLONE SODIUM SUCCINATE

INJECTABLE; INJECTION

A-METHAPRED

<u>AP</u>		HOSPIRA	<u>EQ 40MG BASE/VIAL</u>	<u>N40664</u>	<u>001</u>	Dec 20, 2005
<u>AP</u>			<u>EQ 40MG BASE/VIAL</u>	<u>N85853</u>	<u>001</u>	
<u>AP</u>			<u>EQ 125MG BASE/VIAL</u>	<u>N40665</u>	<u>001</u>	Dec 20, 2005
<u>AP</u>			<u>EQ 125MG BASE/VIAL</u>	<u>N85855</u>	<u>001</u>	
<u>AP</u>			<u>EQ 500MG BASE/VIAL</u>	<u>N85854</u>	<u>001</u>	
<u>AP</u>			<u>EQ 500MG BASE/VIAL</u>	<u>N89173</u>	<u>001</u>	Aug 18, 1987
<u>AP</u>			<u>EQ 1GM BASE/VIAL</u>	<u>N85852</u>	<u>001</u>	
<u>AP</u>			<u>EQ 1GM BASE/VIAL</u>	<u>N89174</u>	<u>001</u>	Aug 18, 1987

METHYLPREDNISOLONE SODIUM SUCCINATE

<u>AP</u>		ABRAXIS PHARM	<u>EQ 40MG BASE/VIAL</u>	<u>N40583</u>	<u>001</u>	Jul 30, 2004
<u>AP</u>			<u>EQ 125MG BASE/VIAL</u>	<u>N40583</u>	<u>002</u>	Jul 30, 2004
<u>AP</u>			<u>EQ 1GM BASE/VIAL</u>	<u>N40612</u>	<u>001</u>	Aug 12, 2004
<u>AP</u>		SICOR PHARMS	<u>EQ 125MG BASE/VIAL</u>	<u>N81266</u>	<u>001</u>	Nov 30, 1992

SOLU-MEDROL

<u>AP</u>	+	PHARMACIA AND UPJOHN	<u>EQ 40MG BASE/VIAL</u>	<u>N11856</u>	<u>003</u>	
<u>AP</u>	+		<u>EQ 125MG BASE/VIAL</u>	<u>N11856</u>	<u>004</u>	
<u>AP</u>	+		<u>EQ 500MG BASE/VIAL</u>	<u>N11856</u>	<u>005</u>	
<u>AP</u>	+		<u>EQ 1GM BASE/VIAL</u>	<u>N11856</u>	<u>006</u>	
	+		EQ 2GM BASE/VIAL	N11856	007	Feb 27, 1985

METHYLTESTOSTERONECAPSULE; ORAL
TESTRED

BP	+	VALEANT PHARM INTL	10MG	N83976	001	
		VIRILON				
BP		STAR PHARMS FL	10MG	N87750	001	Nov 24, 1982

TABLET; ORAL
ANDROID 10

BP		VALEANT PHARM INTL	10MG	N86450	001	
		ANDROID 25				

BP	+	VALEANT PHARM INTL	25MG	N87147	001	
		METHYLTESTOSTERONE				

BP		IMPAX LABS	10MG	N80767	002	
BP			25MG	N84310	001	

METIPRANOLOL HYDROCHLORIDE

SOLUTION/DROPS; OPHTHALMIC

METIPRANOLOL

<u>AT</u>		FALCON PHARMS	<u>0.3%</u>	<u>N75720</u>	<u>001</u>	Aug 06, 2001
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OPTIPRANOLOL

<u>AT</u>	+	BAUSCH AND LOMB	<u>0.3%</u>	<u>N19907</u>	<u>001</u>	Dec 29, 1989
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27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 255 (of 370)

METOCLOPRAMIDE HYDROCHLORIDE

INJECTABLE; INJECTION

METOCLOPRAMIDE HYDROCHLORIDE

<u>AP</u>	HOSPIRA	<u>EQ 5MG BASE/ML</u>	<u>N70505</u>	<u>001</u>	Jun 23, 1989
<u>AP</u>		<u>EQ 5MG BASE/ML</u>	<u>N73117</u>	<u>001</u>	Jan 17, 1991
<u>AP</u>		<u>EQ 5MG BASE/ML</u>	<u>N73118</u>	<u>001</u>	Jan 17, 1991
<u>AP</u>		<u>EQ 5MG BASE/ML</u>	<u>N74147</u>	<u>001</u>	Aug 02, 1996
<u>AP</u>	MAYNE PHARMA USA	<u>EQ 5MG BASE/ML</u>	<u>N70847</u>	<u>001</u>	Nov 07, 1988
<u>AP</u>		<u>EQ 5MG BASE/ML</u>	<u>N71291</u>	<u>001</u>	Mar 03, 1989
<u>AP</u>	SICOR PHARMS	<u>EQ 5MG BASE/ML</u>	<u>N73135</u>	<u>001</u>	Nov 27, 1991

REGLAN

<u>AP</u>	+ BAXTER HLTHCARE CORP	<u>EQ 5MG BASE/ML</u>	<u>N17862</u>	<u>001</u>	
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SOLUTION; ORAL

METOCLOPRAMIDE

<u>AA</u>	+ JVL	<u>EQ 5MG BASE/5ML</u>	<u>N74703</u>	<u>001</u>	Oct 31, 1997
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METOCLOPRAMIDE HYDROCHLORIDE

<u>AA</u>	ACTAVIS MID ATLANTIC	<u>EQ 5MG BASE/5ML</u>	<u>N71340</u>	<u>001</u>	Aug 18, 1988
<u>AA</u>	PHARM ASSOC	<u>EQ 5MG BASE/5ML</u>	<u>N72744</u>	<u>001</u>	May 28, 1991
<u>AA</u>	PHARM VENTURES	<u>EQ 5MG BASE/5ML</u>	<u>N71402</u>	<u>001</u>	Jun 25, 1993
<u>AA</u>	SILARX	<u>EQ 5MG BASE/5ML</u>	<u>N73680</u>	<u>001</u>	Oct 27, 1992

TABLET; ORAL

METOCLOPRAMIDE

<u>AB</u>	VINTAGE PHARMS	<u>EQ 5MG BASE</u>	<u>N77878</u>	<u>001</u>	Aug 28, 2006
<u>AB</u>		<u>EQ 10MG BASE</u>	<u>N77878</u>	<u>002</u>	Aug 28, 2006

METOCLOPRAMIDE HYDROCHLORIDE

<u>AB</u>	ACTAVIS ELIZABETH	<u>EQ 10MG BASE</u>	<u>N70581</u>	<u>001</u>	Oct 17, 1985
<u>AB</u>	MUTUAL PHARM	<u>EQ 5MG BASE</u>	<u>N71536</u>	<u>002</u>	Jan 16, 1997
<u>AB</u>		<u>EQ 10MG BASE</u>	<u>N71536</u>	<u>001</u>	Apr 28, 1993
<u>AB</u>	PLIVA	<u>EQ 5MG BASE</u>	<u>N72750</u>	<u>001</u>	Dec 28, 1995
<u>AB</u>		<u>EQ 10MG BASE</u>	<u>N71250</u>	<u>001</u>	Feb 03, 1988
<u>AB</u>	SANDOZ	<u>EQ 5MG BASE</u>	<u>N74478</u>	<u>001</u>	Oct 05, 1995
<u>AB</u>		<u>EQ 10MG BASE</u>	<u>N72215</u>	<u>001</u>	Jan 30, 1990
<u>AB</u>		<u>EQ 10MG BASE</u>	<u>N74478</u>	<u>002</u>	Oct 05, 1995
<u>AB</u>	TEVA	<u>EQ 5MG BASE</u>	<u>N72801</u>	<u>001</u>	Jun 15, 1993
<u>AB</u>		<u>EQ 10MG BASE</u>	<u>N70184</u>	<u>001</u>	Jul 29, 1985
<u>AB</u>	WATSON LABS	<u>EQ 10MG BASE</u>	<u>N70511</u>	<u>001</u>	Jan 22, 1986

REGLAN

<u>AB</u>	SCHWARZ PHARMA	<u>EQ 5MG BASE</u>	<u>N17854</u>	<u>002</u>	May 05, 1987
<u>AB</u>	+	<u>EQ 10MG BASE</u>	<u>N17854</u>	<u>001</u>	

METOLAZONE

TABLET; ORAL

METOLAZONE

<u>AB</u>	MYLAN	<u>2.5MG</u>	<u>N76698</u>	<u>001</u>	Dec 23, 2003
<u>AB</u>		<u>5MG</u>	<u>N76698</u>	<u>002</u>	Oct 19, 2004
<u>AB</u>		<u>10MG</u>	<u>N76698</u>	<u>003</u>	Oct 19, 2004
<u>AB</u>	ROXANE	<u>10MG</u>	<u>N76482</u>	<u>002</u>	Apr 29, 2004
<u>AB</u>	SANDOZ	<u>2.5MG</u>	<u>N76732</u>	<u>001</u>	Dec 19, 2003
<u>AB</u>		<u>5MG</u>	<u>N76466</u>	<u>001</u>	Dec 19, 2003
<u>AB</u>		<u>10MG</u>	<u>N76466</u>	<u>002</u>	Dec 19, 2003
<u>AB</u>	TEVA	<u>2.5MG</u>	<u>N76600</u>	<u>001</u>	Jan 06, 2004
<u>AB</u>		<u>5MG</u>	<u>N76833</u>	<u>001</u>	Mar 01, 2004
<u>AB</u>		<u>10MG</u>	<u>N75543</u>	<u>003</u>	Dec 24, 2003
<u>AB</u>	WATSON LABS	<u>10MG</u>	<u>N76891</u>	<u>001</u>	Jul 21, 2004
<u>ZAROXOLYN</u>					
<u>AB</u>	UCB INC	<u>2.5MG</u>	<u>N17386</u>	<u>001</u>	
<u>AB</u>	+	<u>5MG</u>	<u>N17386</u>	<u>002</u>	
<u>AB</u>	+	<u>10MG</u>	<u>N17386</u>	<u>003</u>	

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 256 (of 370)

METOPROLOL SUCCINATE

TABLET, EXTENDED RELEASE; ORAL

METOPROLOL SUCCINATE

<u>AB</u>	SANDOZ	<u>EQ 25MG TARTRATE</u>	<u>N76969</u>	<u>001</u>	Jul 31, 2006
<u>AB</u>	<u>TOPROL-XL</u>				
<u>AB</u>	ASTRAZENECA	<u>EQ 25MG TARTRATE</u>	<u>N19962</u>	<u>004</u>	Feb 05, 2001
+		EQ 50MG TARTRATE	N19962	001	Jan 10, 1992
		EQ 100MG TARTRATE	N19962	002	Jan 10, 1992
+		EQ 200MG TARTRATE	N19962	003	Jan 10, 1992

METOPROLOL TARTRATE

INJECTABLE; INJECTION

LOPRESSOR

<u>AP</u>	+ NOVARTIS	<u>1MG/ML</u>	<u>N18704</u>	<u>001</u>	Mar 30, 1984
<u>AP</u>	<u>METOPROLOL TARTRATE</u>				
<u>AP</u>	BEDFORD LABS	<u>1MG/ML</u>	<u>N76495</u>	<u>001</u>	Jul 07, 2003
<u>AP</u>	HOSPIRA	<u>1MG/ML</u>	<u>N74133</u>	<u>001</u>	Dec 21, 1993
<u>AP</u>		<u>1MG/ML</u>	<u>N75160</u>	<u>001</u>	Jul 06, 1998
<u>AP</u>	WATSON LABS	<u>1MG/ML</u>	<u>N74032</u>	<u>001</u>	Dec 21, 1993

TABLET; ORAL

LOPRESSOR

<u>AB</u>	NOVARTIS	<u>50MG</u>	<u>N17963</u>	<u>001</u>	
<u>AB</u>		<u>100MG</u>	<u>N17963</u>	<u>002</u>	
<u>AB</u>	<u>METOPROLOL TARTRATE</u>				
<u>AB</u>	CARACO	<u>25MG</u>	<u>N76670</u>	<u>001</u>	Jan 15, 2004
<u>AB</u>		<u>50MG</u>	<u>N74644</u>	<u>001</u>	Dec 10, 1996
<u>AB</u>		<u>100MG</u>	<u>N74644</u>	<u>002</u>	Dec 10, 1996
<u>AB</u>	MUTUAL PHARM	<u>50MG</u>	<u>N73653</u>	<u>001</u>	Dec 21, 1993
<u>AB</u>		<u>100MG</u>	<u>N73654</u>	<u>001</u>	Dec 21, 1993
<u>AB</u>	MYLAN	<u>25MG</u>	<u>N76704</u>	<u>001</u>	Jan 16, 2004
<u>AB</u>		<u>50MG</u>	<u>N73666</u>	<u>001</u>	Dec 21, 1993
<u>AB</u>		<u>50MG</u>	<u>N76704</u>	<u>002</u>	Jan 16, 2004
<u>AB</u>		<u>100MG</u>	<u>N73666</u>	<u>002</u>	Dec 21, 1993
<u>AB</u>	+	<u>100MG</u>	<u>N76704</u>	<u>003</u>	Jan 16, 2004
<u>AB</u>	PAR PHARM	<u>50MG</u>	<u>N74453</u>	<u>001</u>	Apr 27, 1995
<u>AB</u>		<u>100MG</u>	<u>N74453</u>	<u>002</u>	Apr 27, 1995
<u>AB</u>	SANDOZ	<u>50MG</u>	<u>N73288</u>	<u>001</u>	Mar 25, 1994
<u>AB</u>		<u>100MG</u>	<u>N73289</u>	<u>001</u>	Mar 25, 1994
<u>AB</u>	TEVA	<u>50MG</u>	<u>N74141</u>	<u>001</u>	Jan 31, 1995
<u>AB</u>		<u>50MG</u>	<u>N74143</u>	<u>001</u>	Sep 30, 1994
<u>AB</u>		<u>100MG</u>	<u>N74141</u>	<u>002</u>	Jan 31, 1995
<u>AB</u>		<u>100MG</u>	<u>N74143</u>	<u>002</u>	Sep 30, 1994
<u>AB</u>	TEVA PHARMS	<u>50MG</u>	<u>N74333</u>	<u>001</u>	Jan 27, 1994
<u>AB</u>		<u>100MG</u>	<u>N74333</u>	<u>002</u>	Jan 27, 1994
<u>AB</u>	WATSON LABS	<u>50MG</u>	<u>N74217</u>	<u>001</u>	May 27, 1994
<u>AB</u>		<u>100MG</u>	<u>N74217</u>	<u>002</u>	May 27, 1994

METRONIDAZOLE

CAPSULE; ORAL

FLAGYL

<u>AB</u>	+ GD SEARLE LLC	<u>375MG</u>	<u>N20334</u>	<u>001</u>	May 03, 1995
<u>AB</u>	<u>METRONIDAZOLE</u>				
<u>AB</u>	KALI LABS	<u>375MG</u>	<u>N76522</u>	<u>001</u>	Jan 29, 2004

CREAM; TOPICAL

METROCREAM

<u>AB</u>	+ GALDERMA LABS LP	<u>0.75%</u>	<u>N20531</u>	<u>001</u>	Sep 20, 1995
<u>AB</u>	<u>METRONIDAZOLE</u>				
<u>AB</u>	ALTANA	<u>0.75%</u>	<u>N76408</u>	<u>001</u>	May 28, 2004
	NORITATE				
+	SANOFI AVENTIS US	1%	N20743	001	Sep 26, 1997

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 257 (of 370)

METRONIDAZOLE

GEL; TOPICAL

METROGEL

<u>AB</u>	+	GALDERMA LABS LP	<u>0.75%</u>	<u>N19737</u>	<u>001</u>	Nov 22, 1988
	+		1%	N21789	001	Jun 30, 2005

METRONIDAZOLE

<u>AB</u>		ALTANA	<u>0.75%</u>	<u>N77018</u>	<u>001</u>	Jun 06, 2006
<u>AB</u>		QLT USA	<u>0.75%</u>	<u>N77547</u>	<u>001</u>	Jul 13, 2006
<u>AB</u>		TARO	<u>0.75%</u>	<u>N77819</u>	<u>001</u>	Jul 18, 2006

GEL; VAGINAL

METROGEL-VAGINAL

<u>AB</u>	+	3M	<u>0.75%</u>	<u>N20208</u>	<u>001</u>	Aug 17, 1992
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METRONIDAZOLE

<u>AB</u>		QLT USA	<u>0.75%</u>	<u>N77264</u>	<u>001</u>	Oct 31, 2006
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VANDA ZOLE

BX		TEVA PHARMS	0.75%	N21806	001	May 20, 2005
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INJECTABLE; INJECTION

FLAGYL I.V. RTU IN PLASTIC CONTAINER

<u>AP</u>	+	BAXTER HLTHCARE	<u>500MG/100ML</u>	<u>N18657</u>	<u>001</u>	
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<u>AP</u>	+	GD SEARLE LLC	<u>500MG/100ML</u>	<u>N18353</u>	<u>002</u>	
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METRO I.V. IN PLASTIC CONTAINER

<u>AP</u>	+	B BRAUN	<u>500MG/100ML</u>	<u>N18900</u>	<u>001</u>	Sep 29, 1983
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METRONIDAZOLE IN PLASTIC CONTAINER

<u>AP</u>	+	HOSPIRA	<u>500MG/100ML</u>	<u>N18890</u>	<u>002</u>	Nov 18, 1983
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LOTION; TOPICAL

METROLOTION

<u>AB</u>	+	GALDERMA LABS LP	<u>0.75%</u>	<u>N20901</u>	<u>001</u>	Nov 24, 1998
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METRONIDAZOLE

<u>AB</u>		ALTANA	<u>0.75%</u>	<u>N77197</u>	<u>001</u>	May 24, 2006
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TABLET; ORAL

FLAGYL

<u>AB</u>		GD SEARLE LLC	<u>250MG</u>	<u>N12623</u>	<u>001</u>	
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<u>AB</u>	+		<u>500MG</u>	<u>N12623</u>	<u>003</u>	
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METRONIDAZOLE

<u>AB</u>		IVAX PHARMS	<u>250MG</u>	<u>N18517</u>	<u>001</u>	
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<u>AB</u>			<u>500MG</u>	<u>N18517</u>	<u>002</u>	May 05, 1982
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<u>AB</u>		LNK	<u>250MG</u>	<u>N19029</u>	<u>001</u>	Apr 10, 1984
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<u>AB</u>		MUTUAL PHARM	<u>250MG</u>	<u>N70772</u>	<u>001</u>	Jul 16, 1986
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<u>AB</u>			<u>500MG</u>	<u>N70773</u>	<u>001</u>	Jul 16, 1986
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<u>AB</u>		PAR PHARM	<u>250MG</u>	<u>N18845</u>	<u>001</u>	Aug 18, 1983
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<u>AB</u>			<u>250MG</u>	<u>N70040</u>	<u>001</u>	Jan 29, 1985
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<u>AB</u>			<u>500MG</u>	<u>N18930</u>	<u>001</u>	Aug 18, 1983
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<u>AB</u>			<u>500MG</u>	<u>N70039</u>	<u>001</u>	Jan 29, 1985
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<u>AB</u>		PLIVA	<u>250MG</u>	<u>N70027</u>	<u>001</u>	Nov 06, 1984
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<u>AB</u>			<u>500MG</u>	<u>N70033</u>	<u>001</u>	Dec 06, 1984
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<u>AB</u>		SANDOZ	<u>250MG</u>	<u>N18620</u>	<u>001</u>	Mar 04, 1982
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<u>AB</u>			<u>250MG</u>	<u>N18740</u>	<u>001</u>	Oct 22, 1982
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<u>AB</u>			<u>500MG</u>	<u>N18620</u>	<u>002</u>	Jun 02, 1983
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<u>AB</u>			<u>500MG</u>	<u>N18740</u>	<u>002</u>	Oct 22, 1982
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<u>AB</u>		TEVA	<u>250MG</u>	<u>N70035</u>	<u>001</u>	Dec 20, 1984
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<u>AB</u>			<u>500MG</u>	<u>N70044</u>	<u>001</u>	Feb 08, 1985
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<u>AB</u>		WATSON LABS	<u>250MG</u>	<u>N18764</u>	<u>001</u>	Sep 17, 1982
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<u>AB</u>			<u>500MG</u>	<u>N18764</u>	<u>002</u>	Dec 20, 1982
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TABLET, EXTENDED RELEASE; ORAL

FLAGYL ER

	+	GD SEARLE LLC	750MG	N20868	001	Nov 26, 1997
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27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 258 (of 370)

METRONIDAZOLE HYDROCHLORIDE

INJECTABLE; INJECTION

FLAGYL I.V.

+ GD SEARLE LLC EQ 500MG BASE/VIAL N18353 001

METYRAPONE

CAPSULE; ORAL

METOPIRONE

+ NOVARTIS 250MG N12911 002 Aug 09, 1996

METYROSINE

CAPSULE; ORAL

DEMSEER

+ ATON 250MG N17871 001

MEXILETINE HYDROCHLORIDE

CAPSULE; ORAL

MEXILETINE HYDROCHLORIDE

<u>AB</u>	SANDOZ	<u>150MG</u>	<u>N74450</u>	<u>001</u>	May 16, 1996
<u>AB</u>		<u>200MG</u>	<u>N74450</u>	<u>002</u>	May 16, 1996
<u>AB</u>		<u>250MG</u>	<u>N74450</u>	<u>003</u>	May 16, 1996
<u>AB</u>	TEVA	<u>150MG</u>	<u>N74377</u>	<u>001</u>	May 16, 1995
<u>AB</u>		<u>200MG</u>	<u>N74377</u>	<u>002</u>	May 16, 1995
<u>AB</u>		<u>250MG</u>	<u>N74377</u>	<u>003</u>	May 16, 1995
<u>AB</u>	WATSON LABS	<u>150MG</u>	<u>N74711</u>	<u>001</u>	Feb 26, 1997
<u>AB</u>		<u>150MG</u>	<u>N74865</u>	<u>001</u>	Apr 13, 1998
<u>AB</u>		<u>200MG</u>	<u>N74711</u>	<u>002</u>	Feb 26, 1997
<u>AB</u>		<u>200MG</u>	<u>N74865</u>	<u>002</u>	Apr 13, 1998
<u>AB</u>		<u>250MG</u>	<u>N74711</u>	<u>003</u>	Feb 26, 1997
<u>AB</u>		<u>250MG</u>	<u>N74865</u>	<u>003</u>	Apr 13, 1998
<u>MEXITIL</u>					
<u>AB</u>	BOEHRINGER INGELHEIM	<u>150MG</u>	<u>N18873</u>	<u>002</u>	Dec 30, 1985
<u>AB</u>		<u>200MG</u>	<u>N18873</u>	<u>003</u>	Dec 30, 1985
<u>AB</u>	+	<u>250MG</u>	<u>N18873</u>	<u>004</u>	Dec 30, 1985

MICAFUNGIN SODIUM

INJECTABLE; IV (INFUSION)

+ ASTELLAS 50MG/VIAL N21506 002 Mar 16, 2005

100MG/VIAL N21506 003 Jun 27, 2006

MICONAZOLE NITRATE

CREAM; TOPICAL

MONISTAT-DERM

+ JOHNSON AND JOHNSON 2% N17494 001

SUPPOSITORY; VAGINAL

MICONAZOLE NITRATE

<u>AB</u>	ACTAVIS MID ATLANTIC	<u>200MG</u>	<u>N73508</u>	<u>001</u>	Nov 19, 1993
<u>MONISTAT 3</u>					
<u>AB</u>	+ PERSONAL PRODS	<u>200MG</u>	<u>N18888</u>	<u>001</u>	Aug 15, 1984

MICONAZOLE NITRATE; PETROLATUM, WHITE; ZINC OXIDE

OINTMENT; TOPICAL

VUSION

+ BARRIER 0.25%;81.35%;15% N21026 001 Feb 16, 2006

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 259 (of 370)

MIDAZOLAM HYDROCHLORIDE

INJECTABLE; INJECTION

MIDAZOLAM HYDROCHLORIDE

<u>AP</u>	ABRAXIS PHARM	<u>EQ 1MG BASE/ML</u>	<u>N75154</u>	<u>002</u>	Jun 20, 2000
<u>AP</u>		<u>EQ 5MG BASE/ML</u>	<u>N75154</u>	<u>001</u>	Jun 20, 2000
<u>AP</u>	APOTEX INC	<u>EQ 1MG BASE/ML</u>	<u>N75637</u>	<u>001</u>	Oct 31, 2000
<u>AP</u>		<u>EQ 5MG BASE/ML</u>	<u>N75637</u>	<u>002</u>	Oct 31, 2000
<u>AP</u>	BAXTER HLTHCARE	<u>EQ 1MG BASE/ML</u>	<u>N75243</u>	<u>001</u>	Jun 20, 2000
<u>AP</u>		<u>EQ 5MG BASE/ML</u>	<u>N75243</u>	<u>002</u>	Jun 20, 2000
<u>AP</u>	BAXTER HLTHCARE CORP	<u>EQ 1MG BASE/ML</u>	<u>N75324</u>	<u>001</u>	Jun 20, 2000
<u>AP</u>		<u>EQ 5MG BASE/ML</u>	<u>N75324</u>	<u>002</u>	Jun 20, 2000
<u>AP</u>	BEDFORD	<u>EQ 1MG BASE/ML</u>	<u>N75247</u>	<u>002</u>	Jun 23, 2000
<u>AP</u>		<u>EQ 5MG BASE/ML</u>	<u>N75247</u>	<u>001</u>	Jun 23, 2000
<u>AP</u>	BEN VENUE	<u>EQ 1MG BASE/ML</u>	<u>N75421</u>	<u>002</u>	Jun 20, 2000
<u>AP</u>		<u>EQ 5MG BASE/ML</u>	<u>N75421</u>	<u>001</u>	Jun 20, 2000
<u>AP</u>	HOSPIRA	<u>EQ 1MG BASE/ML</u>	<u>N75293</u>	<u>001</u>	Jun 20, 2000
<u>AP</u>		<u>EQ 1MG BASE/ML</u>	<u>N75409</u>	<u>002</u>	Jun 20, 2000
<u>AP</u>		<u>EQ 1MG BASE/ML</u>	<u>N75856</u>	<u>001</u>	Jun 13, 2002
<u>AP</u>	+	<u>EQ 1MG BASE/ML</u>	<u>N75857</u>	<u>001</u>	Jul 22, 2002
<u>AP</u>	+	<u>EQ 5MG BASE/ML</u>	<u>N75293</u>	<u>002</u>	Jun 20, 2000
<u>AP</u>		<u>EQ 5MG BASE/ML</u>	<u>N75409</u>	<u>001</u>	Jun 20, 2000
<u>AP</u>		<u>EQ 5MG BASE/ML</u>	<u>N75856</u>	<u>002</u>	Jun 13, 2002
<u>AP</u>	+	<u>EQ 5MG BASE/ML</u>	<u>N75857</u>	<u>002</u>	Jul 22, 2002
<u>AP</u>	INTL MEDICATED	<u>EQ 1MG BASE/ML</u>	<u>N76144</u>	<u>001</u>	Jan 26, 2005
<u>AP</u>		<u>EQ 5MG BASE/ML</u>	<u>N76144</u>	<u>002</u>	Jan 26, 2005
<u>AP</u>	INTL MEDICATION	<u>EQ 1MG BASE/ML</u>	<u>N76020</u>	<u>001</u>	Jul 16, 2004
<u>AP</u>		<u>EQ 5MG BASE/ML</u>	<u>N76020</u>	<u>002</u>	Jul 16, 2004
<u>AP</u>	MAYNE PHARMA USA	<u>EQ 1MG BASE/ML</u>	<u>N75396</u>	<u>001</u>	Jun 20, 2000
<u>AP</u>		<u>EQ 5MG BASE/ML</u>	<u>N75396</u>	<u>002</u>	Jun 20, 2000
<u>AP</u>		<u>EQ 5MG BASE/ML</u>	<u>N75484</u>	<u>001</u>	Jun 20, 2000
<u>AP</u>	TAYLOR	<u>EQ 5MG BASE/ML</u>	<u>N75481</u>	<u>001</u>	Jun 30, 2000
<u>AP</u>	TAYLOR PHARMA	<u>EQ 1MG BASE/ML</u>	<u>N75494</u>	<u>001</u>	Jun 30, 2000
<u>AP</u>		<u>EQ 5MG BASE/ML</u>	<u>N75494</u>	<u>002</u>	Jun 30, 2000

SYRUP; ORAL

MIDAZOLAM HYDROCHLORIDE

<u>AA</u>	APOTEX INC	<u>EQ 2MG BASE/ML</u>	<u>N77115</u>	<u>001</u>	Sep 09, 2005
<u>AA</u>	HI TECH PHARMA	<u>EQ 2MG BASE/ML</u>	<u>N75958</u>	<u>001</u>	Sep 04, 2003
<u>AA</u>	PADDOCK	<u>EQ 2MG BASE/ML</u>	<u>N76379</u>	<u>001</u>	May 02, 2005
<u>AA</u>	RANBAXY	<u>EQ 2MG BASE/ML</u>	<u>N76058</u>	<u>001</u>	Mar 15, 2002
<u>AA</u>	+	<u>EQ 2MG BASE/ML</u>	<u>N75873</u>	<u>001</u>	Apr 30, 2002

MIDODRINE HYDROCHLORIDE

TABLET; ORAL

MIDODRINE HYDROCHLORIDE

<u>AB</u>	APOTEX INC	<u>2.5MG</u>	<u>N77746</u>	<u>001</u>	Sep 12, 2006
<u>AB</u>		<u>5MG</u>	<u>N77746</u>	<u>002</u>	Sep 12, 2006
<u>AB</u>		<u>10MG</u>	<u>N77746</u>	<u>003</u>	Sep 12, 2006
<u>AB</u>	IMPAX PHARMS	<u>2.5MG</u>	<u>N76449</u>	<u>001</u>	May 27, 2004
<u>AB</u>		<u>5MG</u>	<u>N76449</u>	<u>002</u>	May 27, 2004
<u>AB</u>		<u>10MG</u>	<u>N76449</u>	<u>003</u>	Dec 16, 2005
<u>AB</u>	MYLAN	<u>2.5MG</u>	<u>N76577</u>	<u>001</u>	Sep 10, 2003
<u>AB</u>		<u>5MG</u>	<u>N76577</u>	<u>002</u>	Sep 10, 2003
<u>AB</u>		<u>10MG</u>	<u>N76577</u>	<u>003</u>	Sep 10, 2003
<u>AB</u>	SANDOZ	<u>2.5MG</u>	<u>N76514</u>	<u>001</u>	Sep 11, 2003
<u>AB</u>		<u>5MG</u>	<u>N76514</u>	<u>002</u>	Sep 11, 2003
<u>AB</u>		<u>10MG</u>	<u>N76514</u>	<u>003</u>	Jul 02, 2004
	<u>ORVATEN</u>				
<u>AB</u>	UPSHER SMITH	<u>2.5MG</u>	<u>N76725</u>	<u>001</u>	Nov 03, 2004
<u>AB</u>		<u>5MG</u>	<u>N76725</u>	<u>002</u>	Nov 03, 2004
<u>AB</u>		<u>10MG</u>	<u>N76725</u>	<u>003</u>	Nov 03, 2004

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 260 (of 370)

MIDODRINE HYDROCHLORIDE

TABLET; ORAL

PROAMATINE

<u>AB</u>	SHIRE	<u>2.5MG</u>	<u>N19815</u>	<u>001</u>	Sep 06, 1996
<u>AB</u>	+	<u>5MG</u>	<u>N19815</u>	<u>002</u>	Sep 06, 1996
<u>AB</u>		<u>10MG</u>	<u>N19815</u>	<u>003</u>	Mar 20, 2002

MIFEPRISTONE

TABLET; ORAL

MIFEPREX

+	DANCO LABS LLC	200MG	N20687	001	Sep 28, 2000
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MIGLITOL

TABLET; ORAL

GLYSET

	PHARMACIA AND UPJOHN	25MG	N20682	001	Dec 18, 1996
		50MG	N20682	002	Dec 18, 1996
+		100MG	N20682	003	Dec 18, 1996

MIGLUSTAT

CAPSULE; ORAL

ZAVESCA

+	ACTELION PHARMS	100MG	N21348	001	Jul 31, 2003
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MILRINONE LACTATE

INJECTABLE; INJECTION

MILRINONE LACTATE

<u>AP</u>	ABRAXIS PHARM	<u>EQ 1MG BASE/ML</u>	<u>N75936</u>	<u>001</u>	May 28, 2002
<u>AP</u>	APOTEX INC	<u>EQ 1MG BASE/ML</u>	<u>N76427</u>	<u>001</u>	Sep 21, 2004
<u>AP</u>	BAXTER HLTHCARE	<u>EQ 1MG BASE/ML</u>	<u>N75530</u>	<u>001</u>	May 28, 2002
<u>AP</u>	BAXTER HLTHCARE CORP	<u>EQ 1MG BASE/ML</u>	<u>N75852</u>	<u>001</u>	May 28, 2002
<u>AP</u>	BEDFORD	<u>EQ 1MG BASE/ML</u>	<u>N75660</u>	<u>001</u>	May 28, 2002
<u>AP</u>	BIONICHE (CANADA)	<u>EQ 1MG BASE/ML</u>	<u>N76428</u>	<u>001</u>	Jun 16, 2003
<u>AP</u>	GLAND PHARMA LTD	<u>EQ 1MG BASE/ML</u>	<u>N77190</u>	<u>001</u>	Oct 31, 2006
<u>AP</u>	HOSPIRA	<u>EQ 1MG BASE/ML</u>	<u>N75884</u>	<u>001</u>	May 28, 2002
<u>AP</u>	INTL MEDICATED	<u>EQ 1MG BASE/ML</u>	<u>N76013</u>	<u>001</u>	Aug 02, 2002
<u>AP</u>	MAYNE PHARMA USA	<u>EQ 1MG BASE/ML</u>	<u>N75830</u>	<u>001</u>	May 28, 2002

MILRINONE LACTATE IN 5% DEXTROSE IN PLASTIC CONTAINER

<u>AP</u>	APOTEX INC	<u>EQ 20MG BASE/100ML</u>	<u>N77151</u>	<u>001</u>	Jul 20, 2005
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MILRINONE LACTATE IN DEXTROSE 5% IN PLASTIC CONTAINER

<u>AP</u>	B BRAUN	<u>EQ 20MG BASE/100ML</u>	<u>N76414</u>	<u>001</u>	Aug 18, 2004
<u>AP</u>	BAXTER HLTHCARE	<u>EQ 20MG BASE/100ML</u>	<u>N75510</u>	<u>001</u>	May 28, 2002
<u>AP</u>		<u>EQ 20MG BASE/100ML</u>	<u>N75834</u>	<u>001</u>	May 28, 2002
<u>AP</u>	HOSPIRA	<u>EQ 20MG BASE/100ML</u>	<u>N75885</u>	<u>001</u>	May 28, 2002

MILRINONE LACTOSE IN 5% DEXTROSE

<u>AP</u>	BAXTER HLTHCARE	<u>EQ 20MG BASE/100ML</u>	<u>N76259</u>	<u>001</u>	Aug 08, 2002
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PRIMACOR

<u>AP</u>	+	SANOFI AVENTIS US	<u>EQ 1MG BASE/ML</u>	<u>N19436</u>	<u>001</u>	Dec 31, 1987
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PRIMACOR IN DEXTROSE 5% IN PLASTIC CONTAINER

<u>AP</u>	+	SANOFI AVENTIS US	<u>EQ 20MG BASE/100ML</u>	<u>N20343</u>	<u>003</u>	Aug 09, 1994
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MINOCYCLINE HYDROCHLORIDE

CAPSULE; ORAL

DYNACIN

<u>AB</u>	MEDICIS	<u>EQ 75MG BASE</u>	<u>N63067</u>	<u>002</u>	Sep 15, 1999
<u>AB</u>		<u>EQ 100MG BASE</u>	<u>N63067</u>	<u>001</u>	Jul 31, 1990

MINOCIN

<u>AB</u>	TRIAx PHARMS	<u>EQ 50MG BASE</u>	<u>N50649</u>	<u>001</u>	May 31, 1990
<u>AB</u>	+	<u>EQ 100MG BASE</u>	<u>N50649</u>	<u>002</u>	May 31, 1990

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 261 (of 370)

MINOCYCLINE HYDROCHLORIDE

CAPSULE; ORAL

MINOCYCLINE HYDROCHLORIDE

<u>AB</u>	IMPAX LABS	<u>EQ 50MG BASE</u>	<u>N65005</u>	<u>001</u>	Mar 23, 1999
<u>AB</u>		<u>EQ 75MG BASE</u>	<u>N65005</u>	<u>003</u>	Apr 18, 2001
<u>AB</u>		<u>EQ 100MG BASE</u>	<u>N65005</u>	<u>002</u>	Mar 23, 1999
<u>AB</u>	RANBAXY	<u>EQ 50MG BASE</u>	<u>N65062</u>	<u>001</u>	Nov 30, 2000
<u>AB</u>		<u>EQ 75MG BASE</u>	<u>N65062</u>	<u>002</u>	Nov 30, 2000
<u>AB</u>		<u>EQ 100MG BASE</u>	<u>N65062</u>	<u>003</u>	Nov 30, 2000
<u>AB</u>	TEVA	<u>EQ 50MG BASE</u>	<u>N63011</u>	<u>001</u>	Mar 02, 1992
<u>AB</u>		<u>EQ 75MG BASE</u>	<u>N63009</u>	<u>002</u>	Aug 12, 2003
<u>AB</u>		<u>EQ 100MG BASE</u>	<u>N63009</u>	<u>001</u>	Mar 02, 1992
<u>AB</u>	WATSON LABS	<u>EQ 50MG BASE</u>	<u>N63181</u>	<u>001</u>	Dec 30, 1991
<u>AB</u>		<u>EQ 75MG BASE</u>	<u>N63065</u>	<u>002</u>	Jun 10, 1999
<u>AB</u>		<u>EQ 100MG BASE</u>	<u>N63065</u>	<u>001</u>	Dec 30, 1991

POWDER, EXTENDED RELEASE; DENTAL

ARESTIN

+	ORAPHARMA	EQ 1MG BASE	N50781	001	Feb 16, 2001
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TABLET; ORAL

MINOCYCLINE HYDROCHLORIDE

<u>AB</u>	MEDICIS	<u>EQ 50MG BASE</u>	<u>N65131</u>	<u>001</u>	Apr 16, 2003
<u>AB</u>		<u>EQ 75MG BASE</u>	<u>N65131</u>	<u>002</u>	Apr 16, 2003
<u>AB</u>	+	<u>EQ 100MG BASE</u>	<u>N65131</u>	<u>003</u>	Apr 16, 2003
<u>AB</u>	RANBAXY	<u>EQ 50MG BASE</u>	<u>N65156</u>	<u>001</u>	Jan 06, 2004
<u>AB</u>		<u>EQ 75MG BASE</u>	<u>N65156</u>	<u>002</u>	Jan 06, 2004
<u>AB</u>		<u>EQ 100MG BASE</u>	<u>N65156</u>	<u>003</u>	Jan 06, 2004

TABLET, EXTENDED RELEASE; ORAL

SOLODYN

	MEDICIS	EQ 45MG BASE	N50808	001	May 08, 2006
		EQ 90MG BASE	N50808	002	May 08, 2006
+		EQ 135MG BASE	N50808	003	May 08, 2006

MINOXIDIL

TABLET; ORAL

LONITEN

<u>AB</u>	PHARMACIA AND UPJOHN	<u>2.5MG</u>	<u>N18154</u>	<u>001</u>	
<u>AB</u>	+	<u>10MG</u>	<u>N18154</u>	<u>003</u>	

MINOXIDIL

<u>AB</u>	MUTUAL PHARM	<u>2.5MG</u>	<u>N72708</u>	<u>001</u>	Dec 14, 1995
<u>AB</u>		<u>10MG</u>	<u>N72709</u>	<u>001</u>	Dec 14, 1995
<u>AB</u>	PAR PHARM	<u>2.5MG</u>	<u>N71826</u>	<u>001</u>	Nov 14, 1988
<u>AB</u>		<u>10MG</u>	<u>N71839</u>	<u>001</u>	Nov 14, 1988
<u>AB</u>	WATSON LABS	<u>2.5MG</u>	<u>N71344</u>	<u>001</u>	Mar 03, 1987
<u>AB</u>		<u>10MG</u>	<u>N71345</u>	<u>001</u>	Mar 03, 1987

MIRTAZAPINE

TABLET; ORAL

MIRTAZAPINE

<u>AB</u>	ACTAVIS ELIZABETH	<u>15MG</u>	<u>N76308</u>	<u>001</u>	Jun 20, 2003
<u>AB</u>		<u>30MG</u>	<u>N76308</u>	<u>002</u>	Jun 20, 2003
<u>AB</u>		<u>45MG</u>	<u>N76308</u>	<u>003</u>	Jun 20, 2003
<u>AB</u>	ACTAVIS TOTOWA	<u>15MG</u>	<u>N76241</u>	<u>001</u>	Jun 25, 2003
<u>AB</u>		<u>30MG</u>	<u>N76241</u>	<u>002</u>	Jun 25, 2003
<u>AB</u>		<u>45MG</u>	<u>N76241</u>	<u>003</u>	Jun 25, 2003
<u>AB</u>	ALPHAPHARM	<u>15MG</u>	<u>N76176</u>	<u>001</u>	Jun 19, 2003
<u>AB</u>		<u>30MG</u>	<u>N76176</u>	<u>002</u>	Jun 19, 2003
<u>AB</u>		<u>45MG</u>	<u>N76176</u>	<u>003</u>	Jun 19, 2003
<u>AB</u>	ANDRX PHARMS	<u>15MG</u>	<u>N76336</u>	<u>001</u>	Jun 20, 2003
<u>AB</u>		<u>30MG</u>	<u>N76336</u>	<u>002</u>	Jun 20, 2003

PRESCRIPTION DRUG PRODUCT LIST

3 - 262 (of 370)

MIRTAZAPINE

TABLET; ORAL

MIRTAZAPINE

<u>AB</u>	ANDRX PHARMS	<u>45MG</u>	<u>N76336</u>	<u>003</u>	Jun 20, 2003
<u>AB</u>	AUROBINDO	<u>7.5MG</u>	<u>N76921</u>	<u>001</u>	Oct 22, 2004
<u>AB</u>		<u>15MG</u>	<u>N76921</u>	<u>002</u>	Oct 22, 2004
<u>AB</u>		<u>30MG</u>	<u>N76921</u>	<u>003</u>	Oct 22, 2004
<u>AB</u>		<u>45MG</u>	<u>N76921</u>	<u>004</u>	Oct 22, 2004
<u>AB</u>	CARACO	<u>7.5MG</u>	<u>N76541</u>	<u>004</u>	Apr 22, 2004
<u>AB</u>		<u>15MG</u>	<u>N76541</u>	<u>001</u>	Apr 22, 2004
<u>AB</u>		<u>30MG</u>	<u>N76541</u>	<u>002</u>	Apr 22, 2004
<u>AB</u>		<u>45MG</u>	<u>N76541</u>	<u>003</u>	Apr 22, 2004
<u>AB</u>	IVAX PHARMS	<u>15MG</u>	<u>N76244</u>	<u>001</u>	Dec 22, 2003
<u>AB</u>		<u>30MG</u>	<u>N76244</u>	<u>002</u>	Dec 22, 2003
<u>AB</u>		<u>45MG</u>	<u>N76244</u>	<u>003</u>	Dec 22, 2003
<u>AB</u>	MYLAN	<u>15MG</u>	<u>N76122</u>	<u>001</u>	Jun 19, 2003
<u>AB</u>		<u>30MG</u>	<u>N76122</u>	<u>002</u>	Jun 19, 2003
<u>AB</u>		<u>45MG</u>	<u>N76122</u>	<u>003</u>	Jun 19, 2003
<u>AB</u>	ROXANE	<u>15MG</u>	<u>N76270</u>	<u>001</u>	Jun 19, 2003
<u>AB</u>		<u>30MG</u>	<u>N76270</u>	<u>002</u>	Jun 19, 2003
<u>AB</u>		<u>45MG</u>	<u>N76270</u>	<u>003</u>	Jun 19, 2003
<u>AB</u>	SANDOZ	<u>15MG</u>	<u>N76189</u>	<u>001</u>	Jun 19, 2003
<u>AB</u>		<u>15MG</u>	<u>N76219</u>	<u>001</u>	Jun 19, 2003
<u>AB</u>		<u>30MG</u>	<u>N76189</u>	<u>002</u>	Jun 19, 2003
<u>AB</u>		<u>30MG</u>	<u>N76219</u>	<u>002</u>	Jun 19, 2003
<u>AB</u>		<u>45MG</u>	<u>N76189</u>	<u>003</u>	Jun 19, 2003
<u>AB</u>		<u>45MG</u>	<u>N76219</u>	<u>003</u>	Jun 19, 2003
<u>AB</u>	TEVA	<u>15MG</u>	<u>N76119</u>	<u>001</u>	Jan 24, 2003
<u>AB</u>		<u>30MG</u>	<u>N76119</u>	<u>002</u>	Jan 24, 2003
<u>AB</u>		<u>45MG</u>	<u>N76119</u>	<u>003</u>	Jun 19, 2003
<u>AB</u>	WATSON LABS	<u>15MG</u>	<u>N76312</u>	<u>001</u>	Jun 19, 2003
<u>AB</u>		<u>30MG</u>	<u>N76312</u>	<u>002</u>	Jun 19, 2003
<u>AB</u>		<u>45MG</u>	<u>N76312</u>	<u>003</u>	Jun 19, 2003
<u>REMERON</u>					
<u>AB</u>	+ ORGANON USA INC	<u>15MG</u>	<u>N20415</u>	<u>001</u>	Jun 14, 1996
<u>AB</u>		<u>30MG</u>	<u>N20415</u>	<u>002</u>	Jun 14, 1996
<u>AB</u>		<u>45MG</u>	<u>N20415</u>	<u>003</u>	Mar 17, 1997

TABLET, ORALLY DISINTEGRATING; ORAL

MIRTAZAPINE

<u>AB</u>	ACTAVIS TOTOWA	<u>15MG</u>	<u>N76689</u>	<u>001</u>	Aug 31, 2005
<u>AB</u>		<u>30MG</u>	<u>N76689</u>	<u>002</u>	Aug 31, 2005
<u>AB</u>		<u>45MG</u>	<u>N76689</u>	<u>003</u>	Aug 31, 2005
<u>AB</u>	AUROBINDO PHARMA LTD	<u>15MG</u>	<u>N77376</u>	<u>002</u>	Dec 08, 2005
<u>AB</u>		<u>30MG</u>	<u>N77376</u>	<u>003</u>	Dec 08, 2005
<u>AB</u>		<u>45MG</u>	<u>N77376</u>	<u>004</u>	Feb 28, 2006
<u>AB</u>	BARR	<u>15MG</u>	<u>N76307</u>	<u>001</u>	Dec 17, 2003
<u>AB</u>		<u>30MG</u>	<u>N76307</u>	<u>002</u>	Dec 17, 2003
<u>AB</u>		<u>45MG</u>	<u>N76307</u>	<u>003</u>	Feb 28, 2006
<u>AB</u>	TEVA	<u>15MG</u>	<u>N76901</u>	<u>001</u>	Jun 28, 2005
<u>AB</u>		<u>30MG</u>	<u>N76901</u>	<u>002</u>	Jun 28, 2005
<u>AB</u>		<u>45MG</u>	<u>N76901</u>	<u>003</u>	Jun 28, 2005
<u>REMERON SOLTAB</u>					
<u>AB</u>	+ ORGANON USA INC	<u>15MG</u>	<u>N21208</u>	<u>001</u>	Jan 12, 2001
<u>AB</u>		<u>30MG</u>	<u>N21208</u>	<u>002</u>	Jan 12, 2001
<u>AB</u>		<u>45MG</u>	<u>N21208</u>	<u>003</u>	Jan 12, 2001

MISOPROSTOL

TABLET; ORAL

CYTOTEC

<u>AB</u>	GD SEARLE LLC	<u>0.1MG</u>	<u>N19268</u>	<u>003</u>	Sep 21, 1990
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PRESCRIPTION DRUG PRODUCT LIST

3 - 263 (of 370)

MISOPROSTOL

TABLET; ORAL					
<u>CYTOTEC</u>					
<u>AB</u>	+ GD SEARLE LLC	<u>0.2MG</u>	<u>N19268</u>	<u>001</u>	Dec 27, 1988
<u>MISOPROSTOL</u>					
<u>AB</u>	IVAX PHARMS	<u>0.1MG</u>	<u>N76095</u>	<u>001</u>	Jul 10, 2002
<u>AB</u>		<u>0.2MG</u>	<u>N76095</u>	<u>002</u>	Jul 10, 2002

MITOMYCIN

INJECTABLE; INJECTION					
<u>MITOMYCIN</u>					
<u>AP</u>	BAXTER HLTHCARE	<u>5MG/VIAL</u>	<u>N64180</u>	<u>001</u>	Dec 23, 1999
<u>AP</u>		<u>20MG/VIAL</u>	<u>N64180</u>	<u>002</u>	Dec 23, 1999
<u>AP</u>	BEDFORD	<u>5MG/VIAL</u>	<u>N64117</u>	<u>001</u>	Apr 19, 1995
<u>AP</u>		<u>20MG/VIAL</u>	<u>N64117</u>	<u>002</u>	Apr 19, 1995
<u>AP</u>	SUPERGEN	<u>5MG/VIAL</u>	<u>N64144</u>	<u>001</u>	Apr 30, 1998
<u>AP</u>		<u>20MG/VIAL</u>	<u>N64144</u>	<u>002</u>	Apr 30, 1998
<u>MUTAMYCIN</u>					
<u>AP</u>	+ BRISTOL MYERS	<u>5MG/VIAL</u>	<u>N62336</u>	<u>001</u>	
<u>AP</u>	+	<u>20MG/VIAL</u>	<u>N62336</u>	<u>002</u>	
	+	40MG/VIAL	N62336	003	Mar 10, 1988
	MYTOZYTREX				
	+ SUPERGEN	5MG/VIAL	N50763	001	Nov 14, 2002

MITOTANE

TABLET; ORAL					
LYSODREN					
	+ BRISTOL	500MG	N16885	001	

MITOXANTRONE HYDROCHLORIDE

INJECTABLE; INJECTION					
<u>MITOXANTRONE</u>					
<u>AP</u>	ABRAXIS PHARM	<u>EQ 20MG BASE/10ML (2MG/ML)</u>	<u>N77496</u>	<u>001</u>	Apr 11, 2006
<u>AP</u>		<u>EQ 25MG BASE/12.5ML (2MG/ML)</u>	<u>N77496</u>	<u>002</u>	Apr 11, 2006
<u>AP</u>		<u>EQ 30MG BASE/15ML (2MG/ML)</u>	<u>N77496</u>	<u>003</u>	Apr 11, 2006
<u>AP</u>	BEDFORD	<u>EQ 20MG BASE/10ML (2MG/ML)</u>	<u>N76611</u>	<u>001</u>	Apr 11, 2006
<u>AP</u>		<u>EQ 25MG BASE/12.5ML (2MG/ML)</u>	<u>N76611</u>	<u>002</u>	Apr 11, 2006
<u>AP</u>		<u>EQ 30MG BASE/15ML (2MG/ML)</u>	<u>N76611</u>	<u>003</u>	Apr 11, 2006
<u>AP</u>	MAYNE PHARMA USA	<u>EQ 20MG BASE/10ML (2MG/ML)</u>	<u>N76871</u>	<u>001</u>	Apr 11, 2006
<u>AP</u>		<u>EQ 25MG BASE/12.5ML (2MG/ML)</u>	<u>N76871</u>	<u>002</u>	Apr 11, 2006
<u>AP</u>		<u>EQ 30MG BASE/15ML (2MG/ML)</u>	<u>N76871</u>	<u>003</u>	Apr 11, 2006
<u>AP</u>	SICOR PHARMS	<u>EQ 20MG BASE/10ML (2MG/ML)</u>	<u>N77356</u>	<u>001</u>	Apr 11, 2006
<u>AP</u>		<u>EQ 25MG BASE/12.5ML (2MG/ML)</u>	<u>N77356</u>	<u>002</u>	Apr 11, 2006
<u>AP</u>		<u>EQ 30MG BASE/15ML (2MG/ML)</u>	<u>N77356</u>	<u>003</u>	Apr 11, 2006
<u>NOVANTRONE</u>					
<u>AP</u>	+ SERONO INC	<u>EQ 20MG BASE/10ML (2MG/ML)</u>	<u>N19297</u>	<u>001</u>	Dec 23, 1987
<u>AP</u>	+	<u>EQ 25MG BASE/12.5ML (2MG/ML)</u>	<u>N19297</u>	<u>002</u>	Dec 23, 1987
<u>AP</u>	+	<u>EQ 30MG BASE/15ML (2MG/ML)</u>	<u>N19297</u>	<u>003</u>	Dec 23, 1987

MODAFINIL

TABLET; ORAL					
PROVIGIL					
	CEPHALON	100MG	N20717	001	Dec 24, 1998
	+	200MG	N20717	002	Dec 24, 1998

MOEXIPRIL HYDROCHLORIDE

TABLET; ORAL					
<u>MOEXIPRIL HYDROCHLORIDE</u>					
<u>AB</u>	PADDOCK	<u>7.5MG</u>	<u>N77536</u>	<u>001</u>	Nov 30, 2006

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 264 (of 370)

MOEXIPRIL HYDROCHLORIDE

TABLET; ORAL

MOEXIPRIL HYDROCHLORIDE

<u>AB</u>	PADDOCK	<u>15MG</u>	<u>N77536</u>	<u>002</u>	Nov 30, 2006
<u>AB</u>	TEVA	<u>7.5MG</u>	<u>N76204</u>	<u>001</u>	May 08, 2003
<u>AB</u>		<u>15MG</u>	<u>N76204</u>	<u>002</u>	May 08, 2003
<u>UNIVASC</u>					
<u>AB</u>	SCHWARZ PHARMA	<u>7.5MG</u>	<u>N20312</u>	<u>001</u>	Apr 19, 1995
<u>AB</u>	+	<u>15MG</u>	<u>N20312</u>	<u>002</u>	Apr 19, 1995

MOLINDONE HYDROCHLORIDE

TABLET; ORAL

MOBAN

	ENDO PHARMS	5MG	N17111	004	
		10MG	N17111	005	
+		25MG	N17111	006	
		50MG	N17111	007	

MOMETASONE FUROATE

CREAM; TOPICAL

ELOCON

<u>AB</u>	+	SCHERING	<u>0.1%</u>	<u>N19625</u>	<u>001</u>	May 06, 1987
<u>MOMETASONE FUROATE</u>						
<u>AB</u>		ALTANA	<u>0.1%</u>	<u>N76171</u>	<u>001</u>	Apr 08, 2005
<u>AB</u>		G AND W LABS	<u>0.1%</u>	<u>N77447</u>	<u>001</u>	May 22, 2006
<u>AB</u>		TARO	<u>0.1%</u>	<u>N76679</u>	<u>001</u>	Dec 21, 2004

LOTION; TOPICAL

ELOCON

<u>AB</u>	+	SCHERING	<u>0.1%</u>	<u>N19796</u>	<u>001</u>	Mar 30, 1989
<u>MOMETASONE FUROATE</u>						
<u>AB</u>		PERRIGO	<u>0.1%</u>	<u>N77180</u>	<u>001</u>	Apr 06, 2005
<u>AB</u>		TARO	<u>0.1%</u>	<u>N76788</u>	<u>001</u>	Mar 15, 2006

OINTMENT; TOPICAL

ELOCON

<u>AB</u>	+	SCHERING	<u>0.1%</u>	<u>N19543</u>	<u>001</u>	Apr 30, 1987
<u>MOMETASONE FUROATE</u>						
<u>AB</u>		ALTANA	<u>0.1%</u>	<u>N77061</u>	<u>001</u>	Mar 28, 2005
<u>AB</u>		G AND W LABS	<u>0.1%</u>	<u>N77401</u>	<u>001</u>	Jun 20, 2006
<u>AB</u>		PERRIGO NEW YORK	<u>0.1%</u>	<u>N76067</u>	<u>001</u>	Mar 18, 2002
<u>AB</u>		QLT USA	<u>0.1%</u>	<u>N76481</u>	<u>001</u>	Nov 14, 2003
<u>AB</u>		TARO	<u>0.1%</u>	<u>N76624</u>	<u>001</u>	Dec 03, 2004

POWDER; INHALATION

ASMANEX TWISTHALER

+	SCHERING	0.22MG/INH	N21067	001	Mar 30, 2005
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MOMETASONE FUROATE MONOHYDRATE

SPRAY, METERED; NASAL

NASONEX

+	SCHERING PLOUGH	EQ 0.05MG BASE/SPRAY	N20762	001	Oct 01, 1997
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MONOBENZONE

CREAM; TOPICAL

BENOQUIN

+	VALEANT PHARM INTL	20%	N08173	003	
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MONTELUKAST SODIUM

GRANULE; ORAL

SINGULAIR

+	MERCK	EQ 4MG BASE/PACKET	N21409	001	Jul 26, 2002
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PRESCRIPTION DRUG PRODUCT LIST

3 - 265 (of 370)

MONTELUKAST SODIUM

TABLET; ORAL
SINGULAIR

+ MERCK	EQ 10MG BASE	N20829	002	Feb 20, 1998
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TABLET, CHEWABLE; ORAL
SINGULAIR

MERCK	EQ 4MG BASE	N20830	002	Mar 03, 2000
+	EQ 5MG BASE	N20830	001	Feb 20, 1998

MORICIZINE HYDROCHLORIDE

TABLET; ORAL
ETHMOZINE

SHIRE	200MG	N19753	001	Jun 19, 1990
	250MG	N19753	002	Jun 19, 1990
+	300MG	N19753	003	Jun 19, 1990

MORPHINE SULFATE

CAPSULE, EXTENDED RELEASE; ORAL
AVINZA

BX	LIGAND	30MG	N21260	001	Mar 20, 2002
BX		60MG	N21260	002	Mar 20, 2002
		90MG	N21260	003	Mar 20, 2002
	+	120MG	N21260	004	Mar 20, 2002

	KADIAN				
BX	ALPHARMA US PHARMS	30MG	N20616	004	Mar 09, 2001
BX		60MG	N20616	005	Mar 09, 2001
		20MG	N20616	001	Jul 03, 1996
		50MG	N20616	002	Jul 03, 1996
		80MG	N20616	006	Oct 27, 2006
	+	100MG	N20616	003	Jul 03, 1996

INJECTABLE; INJECTION

ASTRAMORPH PF

<u>AP</u>	ASTRAZENECA	<u>0.5MG/ML</u>	<u>N71050</u>	<u>001</u>	Oct 07, 1986
<u>AP</u>		<u>0.5MG/ML</u>	<u>N71051</u>	<u>001</u>	Oct 07, 1986
<u>AP</u>		<u>1MG/ML</u>	<u>N71052</u>	<u>001</u>	Oct 07, 1986
<u>AP</u>		<u>1MG/ML</u>	<u>N71053</u>	<u>001</u>	Oct 07, 1986

DURAMORPH PF

<u>AP</u>	+ BAXTER HLTHCARE	<u>0.5MG/ML</u>	<u>N18565</u>	<u>001</u>	Sep 18, 1984
<u>AP</u>	+	<u>1MG/ML</u>	<u>N18565</u>	<u>002</u>	Sep 18, 1984
	INFUMORPH				
	+ BAXTER HLTHCARE	10MG/ML	N18565	003	Jul 19, 1991
	+	25MG/ML	N18565	004	Jul 19, 1991

MORPHINE SULFATE

<u>AP</u>	+ HOSPIRA	<u>0.5MG/ML</u>	<u>N19917</u>	<u>001</u>	Oct 30, 1992
<u>AP</u>		<u>0.5MG/ML</u>	<u>N71849</u>	<u>001</u>	May 11, 1988
<u>AP</u>		<u>0.5MG/ML</u>	<u>N73509</u>	<u>001</u>	Sep 30, 1992
<u>AP</u>	+	<u>1MG/ML</u>	<u>N19916</u>	<u>001</u>	Oct 30, 1992
<u>AP</u>		<u>1MG/ML</u>	<u>N71850</u>	<u>001</u>	May 11, 1988
<u>AP</u>		<u>1MG/ML</u>	<u>N73510</u>	<u>001</u>	Sep 30, 1992
	+	5MG/ML	N19916	002	Oct 27, 2006
<u>AP</u>	MALLINCKRODT	<u>1MG/ML</u>	<u>N20631</u>	<u>001</u>	Jul 03, 1996
	+	2MG/ML	N20631	002	Jul 03, 1996
	+ MERIDIAN MEDCL TECHN	15MG/ML	N19999	001	Jul 12, 1990
<u>AP</u>	WATSON LABS	<u>0.5MG/ML</u>	<u>N73373</u>	<u>001</u>	Sep 30, 1991
<u>AP</u>		<u>0.5MG/ML</u>	<u>N73375</u>	<u>001</u>	Sep 30, 1991
<u>AP</u>		<u>1MG/ML</u>	<u>N73374</u>	<u>001</u>	Sep 30, 1991
<u>AP</u>		<u>1MG/ML</u>	<u>N73376</u>	<u>001</u>	Sep 30, 1991

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 266 (of 370)

MORPHINE SULFATEINJECTABLE, LIPOSOMAL; EPIDURAL
DEPODUR

SKYEPHARMA	10MG/ML	N21671	001	May 18, 2004
	15MG/1.5ML (10MG/ML)	N21671	002	May 18, 2004
+	20MG/2ML (10MG/ML)	N21671	003	May 18, 2004

TABLET, EXTENDED RELEASE; ORAL

MORPHINE SULFATE

<u>AB</u>	AB GENERICS	<u>15MG</u>	<u>N74862</u>	<u>001</u>	Jul 07, 1998
<u>AB</u>		<u>30MG</u>	<u>N74862</u>	<u>002</u>	Jul 07, 1998
<u>AB</u>		<u>60MG</u>	<u>N74862</u>	<u>003</u>	Jul 07, 1998
<u>AB</u>		<u>100MG</u>	<u>N74769</u>	<u>001</u>	Jul 02, 1998
<u>AB</u>		<u>200MG</u>	<u>N74769</u>	<u>002</u>	Jul 02, 1998
<u>AB</u>	ENDO PHARMS	<u>15MG</u>	<u>N75295</u>	<u>001</u>	Oct 28, 1998
<u>AB</u>		<u>30MG</u>	<u>N75295</u>	<u>002</u>	Oct 28, 1998
<u>AB</u>		<u>60MG</u>	<u>N75295</u>	<u>003</u>	Oct 28, 1998
<u>AB</u>		<u>100MG</u>	<u>N75295</u>	<u>004</u>	Sep 15, 2000
<u>AB</u>		<u>200MG</u>	<u>N75295</u>	<u>005</u>	Sep 15, 2000
<u>AB</u>	KV PHARM	<u>15MG</u>	<u>N76733</u>	<u>001</u>	May 19, 2004
<u>AB</u>		<u>30MG</u>	<u>N76720</u>	<u>002</u>	Dec 23, 2005
<u>AB</u>		<u>60MG</u>	<u>N76720</u>	<u>001</u>	May 19, 2004
<u>AB</u>	MALLINCKRODT	<u>15MG</u>	<u>N76412</u>	<u>001</u>	Jul 31, 2003
<u>AB</u>		<u>30MG</u>	<u>N76412</u>	<u>002</u>	Jul 31, 2003
<u>AB</u>		<u>60MG</u>	<u>N76412</u>	<u>003</u>	Jul 31, 2003
<u>AB</u>		<u>100MG</u>	<u>N76438</u>	<u>001</u>	Jul 03, 2003
<u>AB</u>		<u>200MG</u>	<u>N76438</u>	<u>002</u>	Jul 03, 2003
<u>AB</u>	WATSON LABS	<u>100MG</u>	<u>N75656</u>	<u>001</u>	Jan 30, 2001
	<u>MS CONTIN</u>				
<u>AB</u>	PURDUE FREDERICK	<u>15MG</u>	<u>N19516</u>	<u>003</u>	Sep 12, 1989
<u>AB</u>		<u>30MG</u>	<u>N19516</u>	<u>001</u>	May 29, 1987
<u>AB</u>	+	<u>60MG</u>	<u>N19516</u>	<u>002</u>	Apr 08, 1988
<u>AB</u>		<u>100MG</u>	<u>N19516</u>	<u>004</u>	Jan 16, 1990
<u>AB</u>		<u>200MG</u>	<u>N19516</u>	<u>005</u>	Nov 08, 1993
	ORAMORPH SR				
BC	XANODYNE PHARM	15MG	N19977	004	Nov 23, 1994
BC		30MG	N19977	001	Aug 15, 1991
BC	+	60MG	N19977	002	Aug 15, 1991
BC		100MG	N19977	003	Aug 15, 1991

MOXIFLOXACIN HYDROCHLORIDE

INJECTABLE; IV (INFUSION)

AVELOX IN SODIUM CHLORIDE 0.8% IN PLASTIC CONTAINER

+	BAYER PHARMS	160MG/100ML	N21277	001	Nov 30, 2001
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SOLUTION/DROPS; OPHTHALMIC

+	ALCON	0.5%	N21598	001	Apr 15, 2003
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TABLET; ORAL

AVELOX

+	BAYER PHARMS	EQ 400MG BASE	N21085	001	Dec 10, 1999
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MUPIROICIN

OINTMENT; TOPICAL

BACTROBAN

<u>AB</u>	+	GLAXOSMITHKLINE	<u>2%</u>	<u>N50591</u>	<u>001</u>	Dec 31, 1987
		CENTANY				
BX		JOHNSON AND JOHNSON	2%	N50788	001	Dec 04, 2002
		<u>MUPIROICIN</u>				
<u>AB</u>		ALTANA	<u>2%</u>	<u>N65192</u>	<u>001</u>	Nov 30, 2005
<u>AB</u>		PERRIGO NEW YORK	<u>2%</u>	<u>N65123</u>	<u>001</u>	Nov 07, 2003

PRESCRIPTION DRUG PRODUCT LIST

3 - 267 (of 370)

MUPIROCIN

OINTMENT; TOPICAL

MUPIROCIN

<u>AB</u>	TARO	<u>2%</u>	<u>N65170</u>	<u>001</u>	Sep 23, 2005
<u>AB</u>	TEVA	<u>2%</u>	<u>N65085</u>	<u>001</u>	Nov 07, 2003

MUPIROCIN CALCIUM

CREAM, AUGMENTED; TOPICAL
BACTROBAN

+	GLAXOSMITHKLINE	2%	N50746	001	Dec 11, 1997
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OINTMENT; NASAL
BACTROBAN

+	GLAXOSMITHKLINE	EQ 2% BASE	N50703	001	Sep 18, 1995
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MYCOPHENOLATE MOFETIL

CAPSULE; ORAL
CELLCEPT

+	ROCHE PALO	250MG	N50722	001	May 03, 1995
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SUSPENSION; ORAL
CELLCEPT

+	ROCHE PALO	200MG/ML	N50759	001	Oct 01, 1998
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TABLET; ORAL
CELLCEPT

+	ROCHE PALO	500MG	N50723	001	Jun 19, 1997
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MYCOPHENOLATE MOFETIL HYDROCHLORIDE

INJECTABLE; INJECTION
CELLCEPT

+	ROCHE PALO	500MG/VIAL	N50758	001	Aug 12, 1998
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MYCOPHENOLIC ACID

TABLET, DELAYED RELEASE; ORAL
MYFORTIC

	NOVARTIS	180MG	N50791	001	Feb 27, 2004
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+		360MG	N50791	002	Feb 27, 2004
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NABILONE

CAPSULE; ORAL
CESAMET

+	VALEANT	1MG	N18677	001	Dec 26, 1985
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NABUMETONE

TABLET; ORAL

NABUMETONE

<u>AB</u>	PAR PHARM	<u>500MG</u>	<u>N76009</u>	<u>001</u>	Jan 24, 2003
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<u>AB</u>		<u>750MG</u>	<u>N76009</u>	<u>002</u>	Jan 24, 2003
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<u>AB</u>	SANDOZ	<u>500MG</u>	<u>N75280</u>	<u>001</u>	Feb 25, 2002
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<u>AB</u>		<u>500MG</u>	<u>N75590</u>	<u>001</u>	Feb 25, 2002
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<u>AB</u>		<u>750MG</u>	<u>N75280</u>	<u>002</u>	Feb 25, 2002
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<u>AB</u>		<u>750MG</u>	<u>N75590</u>	<u>002</u>	Feb 25, 2002
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<u>AB</u>	TEVA	<u>500MG</u>	<u>N75189</u>	<u>001</u>	May 26, 2000
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<u>AB</u>	+	<u>750MG</u>	<u>N75189</u>	<u>002</u>	Sep 24, 2001
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NADOLOL

TABLET; ORAL

CORGARD

<u>AB</u>	KING PHARMS	<u>20MG</u>	<u>N18063</u>	<u>005</u>	Oct 28, 1986
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<u>AB</u>		<u>40MG</u>	<u>N18063</u>	<u>001</u>	
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27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 268 (of 370)

NADOLOL

TABLET; ORAL

CORGARD

<u>AB</u>	KING PHARMS	<u>80MG</u>	<u>N18063</u>	<u>002</u>	
<u>AB</u>		<u>120MG</u>	<u>N18063</u>	<u>003</u>	
<u>AB</u>	+	<u>160MG</u>	<u>N18063</u>	<u>004</u>	

NADOLOL

<u>AB</u>	IVAX PHARMS	<u>20MG</u>	<u>N74229</u>	<u>001</u>	Aug 30, 1996
<u>AB</u>		<u>40MG</u>	<u>N74229</u>	<u>002</u>	Aug 30, 1996
<u>AB</u>		<u>80MG</u>	<u>N74255</u>	<u>001</u>	Jan 24, 1996
<u>AB</u>		<u>120MG</u>	<u>N74255</u>	<u>002</u>	Jan 24, 1996
<u>AB</u>		<u>160MG</u>	<u>N74255</u>	<u>003</u>	Jan 24, 1996
<u>AB</u>	MYLAN	<u>20MG</u>	<u>N74172</u>	<u>001</u>	Oct 31, 1993
<u>AB</u>		<u>40MG</u>	<u>N74172</u>	<u>002</u>	Oct 31, 1993
<u>AB</u>		<u>80MG</u>	<u>N74172</u>	<u>003</u>	Oct 31, 1993
<u>AB</u>	SANDOZ	<u>20MG</u>	<u>N74501</u>	<u>001</u>	Nov 09, 1995
<u>AB</u>		<u>40MG</u>	<u>N74501</u>	<u>002</u>	Nov 09, 1995
<u>AB</u>		<u>80MG</u>	<u>N74501</u>	<u>003</u>	Nov 09, 1995
<u>AB</u>	TEVA PHARMS	<u>80MG</u>	<u>N74368</u>	<u>001</u>	Aug 31, 1994
<u>AB</u>		<u>120MG</u>	<u>N74368</u>	<u>002</u>	Aug 31, 1994
<u>AB</u>		<u>160MG</u>	<u>N74368</u>	<u>003</u>	Aug 31, 1994

NAFARELIN ACETATESPRAY, METERED; NASAL
SYNAREL

+	GD SEARLE LLC	EQ 0.2MG BASE/SPRAY	N19886	001	Feb 13, 1990
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NAFCILLIN SODIUM

INJECTABLE; INJECTION

NAFCILLIN SODIUM

<u>AP</u>	+	SANDOZ	<u>EQ 1GM BASE/VIAL</u>	<u>N62527</u>	<u>002</u>	Aug 02, 1984
<u>AP</u>	+		<u>EQ 1GM BASE/VIAL</u>	<u>N62732</u>	<u>001</u>	Dec 23, 1986
<u>AP</u>	+		<u>EQ 2GM BASE/VIAL</u>	<u>N62527</u>	<u>003</u>	Aug 02, 1984
<u>AP</u>	+		<u>EQ 2GM BASE/VIAL</u>	<u>N62732</u>	<u>002</u>	Dec 23, 1986
<u>AP</u>	+		<u>EQ 10GM BASE/VIAL</u>	<u>N62527</u>	<u>004</u>	Aug 02, 1984
	+		EQ 500MG BASE/VIAL	N62527	001	Aug 02, 1984
		NALLPEN IN PLASTIC CONTAINER				
	+	BAXTER HLTHCARE	EQ 20MG BASE/ML	N50655	001	Oct 31, 1989
	+		EQ 2GM BASE/100ML	N50655	002	Oct 31, 1989

NAFTIFINE HYDROCHLORIDE

CREAM; TOPICAL

NAFTIN

+	MERZ PHARMS	1%	N19599	001	Feb 29, 1988
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GEL; TOPICAL

NAFTIN

+	MERZ PHARMS	1%	N19356	001	Jun 18, 1990
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NALBUPHINE HYDROCHLORIDE

INJECTABLE; INJECTION

NALBUPHINE HYDROCHLORIDE

<u>AP</u>	HOSPIRA	<u>10MG/ML</u>	<u>N70914</u>	<u>001</u>	Feb 03, 1989
<u>AP</u>		<u>10MG/ML</u>	<u>N70915</u>	<u>001</u>	Feb 03, 1989
<u>AP</u>		<u>20MG/ML</u>	<u>N70916</u>	<u>001</u>	Feb 03, 1989
<u>AP</u>		<u>20MG/ML</u>	<u>N70918</u>	<u>001</u>	Feb 03, 1989
<u>AP</u>	MAYNE PHARMA USA	<u>10MG/ML</u>	<u>N74471</u>	<u>001</u>	Mar 19, 1998
<u>AP</u>		<u>20MG/ML</u>	<u>N74471</u>	<u>002</u>	Mar 19, 1998
		<u>NUBAIN</u>			
<u>AP</u>	+	ENDO PHARMS	<u>10MG/ML</u>	<u>N18024</u>	<u>001</u>

PRESCRIPTION DRUG PRODUCT LIST

3 - 269 (of 370)

NALBUPHINE HYDROCHLORIDE

INJECTABLE; INJECTION

NUBAIN

<u>AP</u>	+ ENDO PHARMS	<u>20MG/ML</u>	<u>N18024</u>	<u>002</u>	May 27, 1982
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NALIDIXIC ACID

TABLET; ORAL

NALIDIXIC ACID

<u>AB</u>	WATSON LABS	<u>250MG</u>	<u>N71936</u>	<u>001</u>	Jun 29, 1988
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<u>AB</u>		<u>500MG</u>	<u>N72061</u>	<u>001</u>	Jun 29, 1988
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<u>AB</u>		<u>1GM</u>	<u>N71919</u>	<u>001</u>	Jun 29, 1988
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NEGGRAM

<u>AB</u>	SANOFI AVENTIS US	<u>250MG</u>	<u>N14214</u>	<u>002</u>	
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<u>AB</u>		<u>500MG</u>	<u>N14214</u>	<u>004</u>	
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<u>AB</u>	+	<u>1GM</u>	<u>N14214</u>	<u>005</u>	
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NALMEFENE HYDROCHLORIDE

INJECTABLE; INJECTION

REVEX

	+ BAXTER HLTHCARE CORP	EQ 0.1MG BASE/ML	N20459	001	Apr 17, 1995
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	+	EQ 1MG BASE/ML	N20459	002	Apr 17, 1995
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NALOXONE HYDROCHLORIDE

INJECTABLE; INJECTION

NALOXONE HYDROCHLORIDE

<u>AP</u>	HOSPIRA	<u>0.02MG/ML</u>	<u>N70171</u>	<u>001</u>	Sep 24, 1986
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<u>AP</u>		<u>0.02MG/ML</u>	<u>N70252</u>	<u>001</u>	Jan 16, 1987
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<u>AP</u>		<u>0.02MG/ML</u>	<u>N70253</u>	<u>001</u>	Jan 16, 1987
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<u>AP</u>		<u>0.4MG/ML</u>	<u>N70172</u>	<u>001</u>	Sep 24, 1986
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<u>AP</u>		<u>0.4MG/ML</u>	<u>N70254</u>	<u>001</u>	Jan 07, 1987
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<u>AP</u>		<u>0.4MG/ML</u>	<u>N70255</u>	<u>001</u>	Jan 07, 1987
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<u>AP</u>		<u>0.4MG/ML</u>	<u>N70256</u>	<u>001</u>	Jan 07, 1987
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<u>AP</u>		<u>0.4MG/ML</u>	<u>N70257</u>	<u>001</u>	Jan 07, 1987
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<u>AP</u>	INTL MEDICATION	<u>0.4MG/ML</u>	<u>N70639</u>	<u>001</u>	Sep 24, 1986
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<u>AP</u>		<u>1MG/ML</u>	<u>N72076</u>	<u>001</u>	Mar 24, 1988
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NARCAN

<u>AP</u>	+ ENDO PHARMS	<u>0.02MG/ML</u>	<u>N16636</u>	<u>002</u>	
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<u>AP</u>	+	<u>0.4MG/ML</u>	<u>N16636</u>	<u>001</u>	
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<u>AP</u>	+	<u>1MG/ML</u>	<u>N16636</u>	<u>003</u>	Jun 14, 1982
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NALOXONE HYDROCHLORIDE; PENTAZOCINE HYDROCHLORIDE

TABLET; ORAL

NALOXONE HYDROCHLORIDE AND PENTAZOCAINE

<u>AB</u>	ACTAVIS TOTOWA	<u>EQ 0.5MG BASE;EQ 50MG BASE</u>	<u>N75735</u>	<u>001</u>	Jul 11, 2001
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PENTAZOCINE AND NALOXONE HYDROCHLORIDES

<u>AB</u>	RANBAXY	<u>EQ 0.5MG BASE;EQ 50MG BASE</u>	<u>N75523</u>	<u>001</u>	Mar 17, 2000
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<u>AB</u>	WATSON LABS	<u>EQ 0.5MG BASE;EQ 50MG BASE</u>	<u>N74736</u>	<u>001</u>	Jan 21, 1997
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TALWIN NX

<u>AB</u>	+ SANOFI AVENTIS US	<u>EQ 0.5MG BASE;EQ 50MG BASE</u>	<u>N18733</u>	<u>001</u>	Dec 16, 1982
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NALTREXONE

FOR SUSPENSION, EXTENDED RELEASE; INTRAMUSCULAR

VIVITROL

	+ ALKERMES	380MG/VIAL	N21897	001	Apr 13, 2006
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NALTREXONE HYDROCHLORIDE

TABLET; ORAL

NALTREXONE HYDROCHLORIDE

<u>AB</u>	ACTAVIS TOTOWA	<u>50MG</u>	<u>N75274</u>	<u>001</u>	May 26, 1999
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27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 270 (of 370)

NALTREXONE HYDROCHLORIDE

TABLET; ORAL

NALTREXONE HYDROCHLORIDE

<u>AB</u>	BARR	<u>50MG</u>	<u>N74918</u>	<u>001</u>	May 08, 1998
<u>AB</u>	MALLINCKRODT	<u>50MG</u>	<u>N76264</u>	<u>002</u>	Mar 22, 2002
		25MG	N76264	001	Mar 22, 2002
	+	100MG	N76264	003	Mar 22, 2002
<u>AB</u>	SANDOZ	<u>50MG</u>	<u>N75434</u>	<u>001</u>	Mar 08, 2000
	<u>REVIA</u>				
<u>AB</u>	+ DURAMED	<u>50MG</u>	<u>N18932</u>	<u>001</u>	Nov 20, 1984

NANDROLONE DECANOATE

INJECTABLE; INJECTION

NANDROLONE DECANOATE

+	WATSON LABS	50MG/ML	N86385	001	Jan 13, 1984
+		100MG/ML	N86598	001	Jan 13, 1984
+		200MG/ML	N88128	001	Dec 05, 1983

NAPHAZOLINE HYDROCHLORIDE

SOLUTION/DROPS; OPHTHALMIC

ALBALON

<u>AT</u>	+ ALLERGAN	<u>0.1%</u>	<u>N80248</u>	<u>001</u>	
	<u>NAFAZAIR</u>				
<u>AT</u>	BAUSCH AND LOMB	<u>0.1%</u>	<u>N40073</u>	<u>001</u>	May 25, 1994
	<u>NAPHAZOLINE HYDROCHLORIDE</u>				
<u>AT</u>	TAYLOR	<u>0.1%</u>	<u>N83590</u>	<u>001</u>	
	<u>VASOCON</u>				
<u>AT</u>	NOVARTIS	<u>0.1%</u>	<u>N80235</u>	<u>002</u>	Mar 24, 1983

NAPROXEN

SUSPENSION; ORAL

NAPROSYN

<u>AB</u>	+ ROCHE PALO	<u>25MG/ML</u>	<u>N18965</u>	<u>001</u>	Mar 23, 1987
	<u>NAPROXEN</u>				
<u>AB</u>	ROXANE	<u>25MG/ML</u>	<u>N74190</u>	<u>001</u>	Mar 30, 1994

TABLET; ORAL

NAPROSYN

<u>AB</u>	ROCHE PALO	<u>250MG</u>	<u>N17581</u>	<u>002</u>	
<u>AB</u>		<u>375MG</u>	<u>N17581</u>	<u>003</u>	
<u>AB</u>	+	<u>500MG</u>	<u>N17581</u>	<u>004</u>	Apr 15, 1982
	<u>NAPROXEN</u>				
<u>AB</u>	CLONMEL HLTHCARE	<u>250MG</u>	<u>N74410</u>	<u>001</u>	Apr 28, 1995
<u>AB</u>		<u>375MG</u>	<u>N74410</u>	<u>002</u>	Apr 28, 1995
<u>AB</u>		<u>500MG</u>	<u>N74410</u>	<u>003</u>	Apr 28, 1995
<u>AB</u>	INTERPHARM	<u>250MG</u>	<u>N75927</u>	<u>001</u>	Dec 18, 2001
<u>AB</u>		<u>375MG</u>	<u>N75927</u>	<u>002</u>	Dec 18, 2001
<u>AB</u>		<u>500MG</u>	<u>N75927</u>	<u>003</u>	Dec 18, 2001
<u>AB</u>	MYLAN	<u>250MG</u>	<u>N74121</u>	<u>001</u>	Dec 21, 1993
<u>AB</u>		<u>375MG</u>	<u>N74121</u>	<u>002</u>	Dec 21, 1993
<u>AB</u>		<u>500MG</u>	<u>N74121</u>	<u>003</u>	Dec 21, 1993
<u>AB</u>	PERRIGO R AND D	<u>250MG</u>	<u>N77339</u>	<u>001</u>	Apr 27, 2005
<u>AB</u>		<u>375MG</u>	<u>N77339</u>	<u>002</u>	Apr 27, 2005
<u>AB</u>		<u>500MG</u>	<u>N77339</u>	<u>003</u>	Apr 27, 2005
<u>AB</u>	PLIVA	<u>250MG</u>	<u>N74182</u>	<u>001</u>	Jun 27, 1996
<u>AB</u>		<u>375MG</u>	<u>N74182</u>	<u>002</u>	Jun 27, 1996
<u>AB</u>		<u>500MG</u>	<u>N74182</u>	<u>003</u>	Jun 27, 1996
<u>AB</u>	SANDOZ	<u>250MG</u>	<u>N74140</u>	<u>001</u>	Dec 21, 1993
<u>AB</u>		<u>375MG</u>	<u>N74140</u>	<u>002</u>	Dec 21, 1993
<u>AB</u>		<u>500MG</u>	<u>N74140</u>	<u>003</u>	Dec 21, 1993
<u>AB</u>	TEVA	<u>250MG</u>	<u>N74129</u>	<u>001</u>	Dec 21, 1993

PRESCRIPTION DRUG PRODUCT LIST

3 - 271 (of 370)

NAPROXEN

TABLET; ORAL

NAPROXEN

<u>AB</u>	TEVA	<u>250MG</u>	<u>N74201</u>	<u>001</u>	Dec 21, 1993
<u>AB</u>		<u>250MG</u>	<u>N74216</u>	<u>001</u>	Apr 11, 1996
<u>AB</u>		<u>375MG</u>	<u>N74129</u>	<u>002</u>	Dec 21, 1993
<u>AB</u>		<u>375MG</u>	<u>N74201</u>	<u>002</u>	Dec 21, 1993
<u>AB</u>		<u>375MG</u>	<u>N74216</u>	<u>002</u>	Apr 11, 1996
<u>AB</u>		<u>500MG</u>	<u>N74129</u>	<u>003</u>	Dec 21, 1993
<u>AB</u>		<u>500MG</u>	<u>N74201</u>	<u>003</u>	Dec 21, 1993
<u>AB</u>		<u>500MG</u>	<u>N74216</u>	<u>003</u>	Apr 11, 1996
<u>AB</u>	TEVA PHARMS	<u>250MG</u>	<u>N74207</u>	<u>001</u>	Dec 21, 1993
<u>AB</u>		<u>375MG</u>	<u>N74207</u>	<u>002</u>	Dec 21, 1993
<u>AB</u>		<u>500MG</u>	<u>N74207</u>	<u>003</u>	Dec 21, 1993
<u>AB</u>	WATSON LABS	<u>250MG</u>	<u>N74163</u>	<u>001</u>	Feb 10, 1995
<u>AB</u>		<u>250MG</u>	<u>N74457</u>	<u>001</u>	May 31, 1995
<u>AB</u>		<u>375MG</u>	<u>N74163</u>	<u>002</u>	Feb 10, 1995
<u>AB</u>		<u>375MG</u>	<u>N74457</u>	<u>002</u>	May 31, 1995
<u>AB</u>		<u>500MG</u>	<u>N74163</u>	<u>003</u>	Feb 10, 1995
<u>AB</u>		<u>500MG</u>	<u>N74457</u>	<u>003</u>	May 31, 1995
<u>AB</u>	WESTWARD	<u>250MG</u>	<u>N76494</u>	<u>001</u>	Jan 14, 2004
<u>AB</u>		<u>375MG</u>	<u>N76494</u>	<u>002</u>	Jan 14, 2004
<u>AB</u>		<u>500MG</u>	<u>N76494</u>	<u>003</u>	Jan 14, 2004

TABLET, DELAYED RELEASE; ORAL

EC-NAPROSYN

<u>AB</u>	+ ROCHE PALO	<u>375MG</u>	<u>N20067</u>	<u>002</u>	Oct 14, 1994
<u>AB</u>	+	<u>500MG</u>	<u>N20067</u>	<u>003</u>	Oct 14, 1994

NAPROXEN

<u>AB</u>	ACTAVIS ELIZABETH	<u>375MG</u>	<u>N74936</u>	<u>001</u>	Feb 24, 1998
<u>AB</u>		<u>500MG</u>	<u>N74936</u>	<u>002</u>	Feb 24, 1998
<u>AB</u>	ALPHAPHARM	<u>375MG</u>	<u>N75390</u>	<u>001</u>	Apr 19, 2001
<u>AB</u>		<u>500MG</u>	<u>N75390</u>	<u>002</u>	Apr 19, 2001
<u>AB</u>	PLIVA	<u>375MG</u>	<u>N75337</u>	<u>001</u>	May 26, 1999
<u>AB</u>		<u>500MG</u>	<u>N75337</u>	<u>002</u>	May 26, 1999
<u>AB</u>	SANDOZ	<u>375MG</u>	<u>N75061</u>	<u>001</u>	Feb 18, 1998
<u>AB</u>		<u>500MG</u>	<u>N75061</u>	<u>002</u>	Feb 18, 1998
<u>AB</u>	TEVA	<u>375MG</u>	<u>N75227</u>	<u>001</u>	Jun 30, 1998
<u>AB</u>		<u>500MG</u>	<u>N75227</u>	<u>002</u>	Jun 30, 1998

NAPROXEN SODIUM

TABLET; ORAL

ANAPROX

<u>AB</u>	ROCHE PALO	<u>EQ 250MG BASE</u>	<u>N18164</u>	<u>001</u>	
<u>AB</u>	<u>ANAPROX DS</u>				
<u>AB</u>	+ ROCHE PALO	<u>EQ 500MG BASE</u>	<u>N18164</u>	<u>003</u>	Sep 30, 1987

NAPROXEN SODIUM

<u>AB</u>	HIKMA	<u>EQ 250MG BASE</u>	<u>N74480</u>	<u>002</u>	Feb 18, 1998
<u>AB</u>		<u>EQ 500MG BASE</u>	<u>N74480</u>	<u>001</u>	May 14, 1996
<u>AB</u>	MYLAN	<u>EQ 250MG BASE</u>	<u>N74367</u>	<u>001</u>	Aug 31, 1994
<u>AB</u>		<u>EQ 500MG BASE</u>	<u>N74367</u>	<u>002</u>	Aug 31, 1994
<u>AB</u>	PLIVA	<u>EQ 250MG BASE</u>	<u>N74242</u>	<u>001</u>	Jun 20, 1996
<u>AB</u>		<u>EQ 500MG BASE</u>	<u>N74242</u>	<u>002</u>	Jun 20, 1996
<u>AB</u>	SANDOZ	<u>EQ 250MG BASE</u>	<u>N74162</u>	<u>001</u>	Dec 21, 1993
<u>AB</u>		<u>EQ 250MG BASE</u>	<u>N74495</u>	<u>001</u>	Dec 05, 1994
<u>AB</u>		<u>EQ 500MG BASE</u>	<u>N74162</u>	<u>002</u>	Dec 21, 1993
<u>AB</u>		<u>EQ 500MG BASE</u>	<u>N74495</u>	<u>002</u>	Dec 05, 1994
<u>AB</u>	TEVA	<u>EQ 250MG BASE</u>	<u>N74142</u>	<u>001</u>	Dec 21, 1993
<u>AB</u>		<u>EQ 250MG BASE</u>	<u>N74198</u>	<u>001</u>	Dec 21, 1993
<u>AB</u>		<u>EQ 500MG BASE</u>	<u>N74142</u>	<u>002</u>	Dec 21, 1993
<u>AB</u>		<u>EQ 500MG BASE</u>	<u>N74198</u>	<u>002</u>	Dec 21, 1993

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 272 (of 370)

NAPROXEN SODIUM

TABLET; ORAL

NAPROXEN SODIUM

<u>AB</u>	TEVA PHARMS	<u>EQ 250MG BASE</u>	<u>N74289</u>	<u>001</u>	Jan 27, 1994
<u>AB</u>		<u>EQ 500MG BASE</u>	<u>N74289</u>	<u>002</u>	Jan 27, 1994
<u>AB</u>	WATSON LABS	<u>EQ 250MG BASE</u>	<u>N74195</u>	<u>001</u>	Dec 21, 1993
<u>AB</u>		<u>EQ 250MG BASE</u>	<u>N74455</u>	<u>001</u>	May 31, 1995
<u>AB</u>		<u>EQ 500MG BASE</u>	<u>N74195</u>	<u>002</u>	Dec 21, 1993
<u>AB</u>		<u>EQ 500MG BASE</u>	<u>N74455</u>	<u>002</u>	May 31, 1995

TABLET, EXTENDED RELEASE; ORAL

NAPRELAN

<u>AB</u>	+ STAT TRADE	<u>EQ 375MG BASE</u>	<u>N20353</u>	<u>001</u>	Jan 05, 1996
<u>AB</u>	+	<u>EQ 500MG BASE</u>	<u>N20353</u>	<u>002</u>	Jan 05, 1996
<u>AB</u>	<u>NAPROXEN SODIUM</u>				
<u>AB</u>	ANDRX PHARMS	<u>EQ 375MG BASE</u>	<u>N75416</u>	<u>002</u>	Apr 23, 2003
<u>AB</u>		<u>EQ 500MG BASE</u>	<u>N75416</u>	<u>001</u>	Aug 27, 2002

NARATRIPTAN HYDROCHLORIDE

TABLET; ORAL

AMERGE

	GLAXOSMITHKLINE	EQ 1MG BASE	N20763	002	Feb 10, 1998
+		EQ 2.5MG BASE	N20763	001	Feb 10, 1998

NATAMYCIN

SUSPENSION; OPHTHALMIC

NATACYN

+	ALCON	5%	N50514	001	
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NATEGLINIDE

TABLET; ORAL

STARLIX

	NOVARTIS	60MG	N21204	001	Dec 22, 2000
+		120MG	N21204	002	Dec 22, 2000

NEDOCROMIL SODIUM

AEROSOL, METERED; INHALATION

TILADE

+	KING PHARMS	1.75MG/INH	N19660	001	Dec 30, 1992
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SOLUTION/DROPS; OPHTHALMIC

+	ALLERGAN	2%	N21009	001	Dec 08, 1999
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NEFAZODONE HYDROCHLORIDE

TABLET; ORAL

NEFAZODONE HYDROCHLORIDE

<u>AB</u>	DR REDDYS LABS INC	<u>50MG</u>	<u>N76309</u>	<u>001</u>	Sep 16, 2003
<u>AB</u>		<u>100MG</u>	<u>N76309</u>	<u>002</u>	Sep 16, 2003
<u>AB</u>		<u>150MG</u>	<u>N76309</u>	<u>003</u>	Sep 16, 2003
<u>AB</u>		<u>200MG</u>	<u>N76309</u>	<u>004</u>	Sep 16, 2003
<u>AB</u>		<u>250MG</u>	<u>N76309</u>	<u>005</u>	Sep 16, 2003
<u>AB</u>	IVAX PHARMS	<u>50MG</u>	<u>N75763</u>	<u>001</u>	Sep 16, 2003
<u>AB</u>		<u>100MG</u>	<u>N75763</u>	<u>002</u>	Sep 16, 2003
<u>AB</u>		<u>150MG</u>	<u>N75763</u>	<u>003</u>	Sep 16, 2003
<u>AB</u>		<u>200MG</u>	<u>N75763</u>	<u>004</u>	Sep 16, 2003
<u>AB</u>		<u>250MG</u>	<u>N75763</u>	<u>005</u>	Sep 16, 2003
<u>AB</u>	MYLAN	<u>100MG</u>	<u>N76129</u>	<u>002</u>	Sep 16, 2003
<u>AB</u>		<u>150MG</u>	<u>N76129</u>	<u>003</u>	Sep 16, 2003
<u>AB</u>		<u>200MG</u>	<u>N76129</u>	<u>004</u>	Sep 16, 2003
<u>AB</u>		<u>250MG</u>	<u>N76129</u>	<u>005</u>	Sep 16, 2003

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 273 (of 370)

NEFAZODONE HYDROCHLORIDE

TABLET; ORAL

NEFAZODONE HYDROCHLORIDE

<u>AB</u>	RANBAXY	<u>50MG</u>	<u>N76409</u>	<u>001</u>	Sep 16, 2003
<u>AB</u>		<u>100MG</u>	<u>N76409</u>	<u>002</u>	Sep 16, 2003
<u>AB</u>		<u>150MG</u>	<u>N76409</u>	<u>003</u>	Sep 16, 2003
<u>AB</u>		<u>200MG</u>	<u>N76409</u>	<u>004</u>	Sep 16, 2003
<u>AB</u>		<u>250MG</u>	<u>N76409</u>	<u>005</u>	Sep 16, 2003
<u>AB</u>	SANDOZ	<u>50MG</u>	<u>N76072</u>	<u>001</u>	Sep 16, 2003
<u>AB</u>		<u>50MG</u>	<u>N76302</u>	<u>001</u>	Sep 16, 2003
<u>AB</u>		<u>100MG</u>	<u>N76072</u>	<u>002</u>	Sep 16, 2003
<u>AB</u>		<u>100MG</u>	<u>N76302</u>	<u>002</u>	Sep 16, 2003
<u>AB</u>		<u>150MG</u>	<u>N76072</u>	<u>003</u>	Sep 16, 2003
<u>AB</u>		<u>150MG</u>	<u>N76302</u>	<u>003</u>	Sep 16, 2003
<u>AB</u>		<u>200MG</u>	<u>N76072</u>	<u>004</u>	Sep 16, 2003
<u>AB</u>		<u>200MG</u>	<u>N76302</u>	<u>004</u>	Sep 16, 2003
<u>AB</u>		<u>250MG</u>	<u>N76072</u>	<u>005</u>	Sep 16, 2003
<u>AB</u>		<u>250MG</u>	<u>N76302</u>	<u>005</u>	Sep 16, 2003
<u>AB</u>	TEVA	<u>50MG</u>	<u>N76037</u>	<u>001</u>	Sep 16, 2003
<u>AB</u>		<u>100MG</u>	<u>N76037</u>	<u>002</u>	Sep 16, 2003
<u>AB</u>		<u>150MG</u>	<u>N76037</u>	<u>003</u>	Sep 16, 2003
<u>AB</u>		<u>200MG</u>	<u>N76037</u>	<u>004</u>	Sep 16, 2003
<u>AB</u>	+	<u>250MG</u>	<u>N76037</u>	<u>005</u>	Sep 16, 2003
<u>AB</u>	WATSON LABS	<u>100MG</u>	<u>N76073</u>	<u>002</u>	Sep 16, 2003
<u>AB</u>		<u>150MG</u>	<u>N76073</u>	<u>003</u>	Sep 16, 2003
<u>AB</u>		<u>200MG</u>	<u>N76073</u>	<u>004</u>	Sep 16, 2003
<u>AB</u>		<u>250MG</u>	<u>N76073</u>	<u>005</u>	Sep 16, 2003

NELARABINE

INJECTABLE; IV (INFUSION)

+	SMITHKLINE BEECHAM	250MG/50ML (5MG/ML)	N21877	001	Oct 28, 2005
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NELFINAVIR MESYLATE

FOR SUSPENSION; ORAL

VIRACEPT

+	AGOURON	EQ 50MG BASE/SCOOPFUL	N20778	001	Mar 14, 1997
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TABLET; ORAL

VIRACEPT

+	AGOURON	EQ 250MG BASE	N20779	001	Mar 14, 1997
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+

		EQ 625MG BASE	N21503	001	Apr 30, 2003
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NEOMYCIN SULFATE

POWDER; FOR RX COMPOUNDING

NEO-RX

	X GEN PHARMS	100%	N61579	001	
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SOLUTION; ORAL

NEO-FRADIN

+	X GEN PHARMS	EQ 87.5MG BASE/5ML	N65010	001	May 23, 2002
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TABLET; ORAL

NEOMYCIN SULFATE

<u>AB</u>	+	TEVA	<u>500MG</u>	<u>N60304</u>	<u>001</u>	
<u>AB</u>		X GEN PHARMS	<u>500MG</u>	<u>N65220</u>	<u>001</u>	Jul 28, 2006

NEOMYCIN SULFATE; POLYMYXIN B SULFATE

SOLUTION; IRRIGATION

NEOMYCIN AND POLYMYXIN B SULFATE

<u>AT</u>	WATSON LABS	<u>EQ 40MG BASE/ML;200,000 UNITS/ML</u>	<u>N62664</u>	<u>001</u>	Apr 08, 1986
<u>AT</u>	X GEN PHARMS	<u>EQ 40MG BASE/ML;200,000 UNITS/ML</u>	<u>N65106</u>	<u>001</u>	Jan 31, 2006

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 274 (of 370)

NEOMYCIN SULFATE; POLYMYXIN B SULFATE

SOLUTION; IRRIGATION

NEOMYCIN AND POLYMYXIN B SULFATE

<u>AT</u>	X GEN PHARMS	<u>EQ 800MG BASE/20ML;4,000,000 UNITS/20ML</u>	<u>N65108</u>	<u>001</u>	Jan 31, 2006
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NEOSPORIN G.U. IRRIGANT

<u>AT</u>	+ MONARCH PHARMS	<u>EQ 40MG BASE/ML;200,000 UNITS/ML</u>	<u>N60707</u>	<u>001</u>	
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<u>AT</u>	+	<u>EQ 800MG BASE/20ML;4,000,000 UNITS/20ML (EQ 40MG BASE/ML;200,000 UNITS/ML)</u>	<u>N60707</u>	<u>002</u>	
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NEOMYCIN SULFATE; POLYMYXIN B SULFATE; PREDNISOLONE ACETATE

SUSPENSION/DROPS; OPHTHALMIC

POLY-PRED

+	ALLERGAN	EQ 0.35% BASE;10,000 UNITS/ML;0.5%	N50081	002	
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NEPAFENAC

SUSPENSION/DROPS; OPHTHALMIC

+	ALCON	0.1%	N21862	001	Aug 19, 2005
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NESIRITIDE RECOMBINANT

FOR SOLUTION; INTRAVENOUS

NATRECOR

+	SCIOS	1.5MG/VIAL	N20920	001	Aug 10, 2001
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NEVIRAPINE

SUSPENSION; ORAL

VIRAMUNE

+	BOEHRINGER INGELHEIM	50MG/5ML	N20933	001	Sep 11, 1998
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TABLET; ORAL

VIRAMUNE

+	BOEHRINGER INGELHEIM	200MG	N20636	001	Jun 21, 1996
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NIACIN

TABLET; ORAL

NIACIN

<u>AA</u>	WOCKHARDT	<u>500MG</u>	<u>N81134</u>	<u>001</u>	Apr 28, 1992
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NIACOR

<u>AA</u>	+	UPSHER SMITH	<u>500MG</u>	<u>N40378</u>	<u>001</u>	May 03, 2000
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TABLET, EXTENDED RELEASE; ORAL

NIASPAN

+	KOS LIFE	500MG	N20381	002	Jul 28, 1997
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+		750MG	N20381	003	Jul 28, 1997
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+		1GM	N20381	004	Jul 28, 1997
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NICARDIPINE HYDROCHLORIDE

CAPSULE; ORAL

CARDENE

<u>AB</u>	PDL BIOPHARMA INC	<u>20MG</u>	<u>N19488</u>	<u>001</u>	Dec 21, 1988
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<u>AB</u>	+	<u>30MG</u>	<u>N19488</u>	<u>002</u>	Dec 21, 1988
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NICARDIPINE HYDROCHLORIDE

<u>AB</u>	BARR	<u>20MG</u>	<u>N74439</u>	<u>001</u>	Dec 10, 1996
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<u>AB</u>		<u>30MG</u>	<u>N74439</u>	<u>002</u>	Dec 10, 1996
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<u>AB</u>	GENPHARM	<u>20MG</u>	<u>N74928</u>	<u>001</u>	Mar 19, 1998
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<u>AB</u>		<u>30MG</u>	<u>N74928</u>	<u>002</u>	Mar 19, 1998
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<u>AB</u>	MYLAN	<u>20MG</u>	<u>N74642</u>	<u>001</u>	Jul 18, 1996
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<u>AB</u>		<u>30MG</u>	<u>N74642</u>	<u>002</u>	Jul 18, 1996
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<u>AB</u>	TEVA	<u>20MG</u>	<u>N74540</u>	<u>001</u>	Oct 28, 1996
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<u>AB</u>		<u>30MG</u>	<u>N74540</u>	<u>002</u>	Oct 28, 1996
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27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 275 (of 370)

NICARDIPINE HYDROCHLORIDE

CAPSULE; ORAL

NICARDIPINE HYDROCHLORIDE

<u>AB</u>	WATSON LABS	<u>20MG</u>	<u>N74670</u>	<u>001</u>	Oct 28, 1996
<u>AB</u>		<u>30MG</u>	<u>N74670</u>	<u>002</u>	Oct 28, 1996

CAPSULE, EXTENDED RELEASE; ORAL

CARDENE SR

+	PDL BIOPHARMA INC	30MG	N20005	001	Feb 21, 1992
+		45MG	N20005	002	Feb 21, 1992
+		60MG	N20005	003	Feb 21, 1992

INJECTABLE; INJECTION

CARDENE

+	PDL BIOPHARMA INC	2.5MG/ML	N19734	001	Jan 30, 1992
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NICOTINE

FILM, EXTENDED RELEASE; TRANSDERMAL

NICOTINE

AVEVA	7MG/24HR	N74645	001	Oct 20, 1997
	14MG/24HR	N74611	001	Oct 20, 1997
	21MG/24HR	N74612	001	Oct 20, 1997

INHALANT; ORAL

NICOTROL

+	PHARMACIA AND UPJOHN	4MG/CARTRIDGE	N20714	001	May 02, 1997
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SPRAY, METERED; NASAL

NICOTROL

+	PFIZER INC	0.5MG/SPRAY	N20385	001	Mar 22, 1996
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NIFEDIPINE

CAPSULE; ORAL

NIFEDIPINE

<u>AB</u>	ACTAVIS ELIZABETH	<u>10MG</u>	<u>N72579</u>	<u>001</u>	Jan 08, 1991
<u>AB</u>		<u>20MG</u>	<u>N72556</u>	<u>001</u>	Sep 20, 1990
<u>AB</u>	FLEMINGTON PHARM	<u>10MG</u>	<u>N72781</u>	<u>001</u>	Jul 30, 1993
<u>AB</u>	SCHERER RP	<u>10MG</u>	<u>N73250</u>	<u>001</u>	Oct 08, 1991
<u>AB</u>		<u>20MG</u>	<u>N74045</u>	<u>001</u>	Apr 30, 1992
<u>AB</u>	TEVA	<u>10MG</u>	<u>N72651</u>	<u>001</u>	Feb 19, 1992

PROCARDIA

<u>AB</u>	PFIZER	<u>10MG</u>	<u>N18482</u>	<u>001</u>	
<u>AB</u>	+	<u>20MG</u>	<u>N18482</u>	<u>002</u>	Jul 24, 1986

TABLET, EXTENDED RELEASE; ORAL

ADALAT CC

<u>AB1</u>	BAYER PHARMS	<u>30MG</u>	<u>N20198</u>	<u>001</u>	Apr 21, 1993
<u>AB1</u>	+	<u>60MG</u>	<u>N20198</u>	<u>002</u>	Apr 21, 1993
<u>AB1</u>	+	<u>90MG</u>	<u>N20198</u>	<u>003</u>	Apr 21, 1993

AFEDITAB CR

<u>AB1</u>	WATSON LABS	<u>30MG</u>	<u>N75128</u>	<u>001</u>	Mar 10, 2000
<u>AB1</u>		<u>60MG</u>	<u>N75659</u>	<u>001</u>	Oct 26, 2001

NIFEDIPINE

<u>AB1</u>	ABRIKA PHARMS	<u>30MG</u>	<u>N77899</u>	<u>001</u>	Dec 13, 2006
<u>AB1</u>		<u>60MG</u>	<u>N77899</u>	<u>002</u>	Dec 13, 2006
<u>AB1</u>	BIOVAIL	<u>30MG</u>	<u>N75269</u>	<u>001</u>	Dec 04, 2000
<u>AB1</u>		<u>60MG</u>	<u>N75269</u>	<u>002</u>	Dec 04, 2000
<u>AB1</u>		<u>90MG</u>	<u>N76070</u>	<u>001</u>	Aug 16, 2002
<u>AB2</u>		<u>30MG</u>	<u>N75289</u>	<u>002</u>	Feb 06, 2001
<u>AB2</u>		<u>60MG</u>	<u>N75289</u>	<u>001</u>	Sep 27, 2000
<u>AB2</u>	MARTEC USA LLC	<u>90MG</u>	<u>N75414</u>	<u>003</u>	Mar 23, 2004
<u>AB2</u>	OSMOTICA PHARM	<u>30MG</u>	<u>N77127</u>	<u>001</u>	Nov 21, 2005
<u>AB2</u>		<u>60MG</u>	<u>N77127</u>	<u>002</u>	Nov 21, 2005

PRESCRIPTION DRUG PRODUCT LIST

3 - 276 (of 370)

NIFEDIPINE

TABLET, EXTENDED RELEASE; ORAL					
PROCARDIA XL					
AB2	PFIZER	30MG	N19684	001	Sep 06, 1989
AB2		60MG	N19684	002	Sep 06, 1989
AB2 +		90MG	N19684	003	Sep 06, 1989

NILUTAMIDE

TABLET; ORAL					
NILANDRON					
	+ SANOFI AVENTIS US	150MG	N20169	002	Apr 30, 1999

NIMODIPINE

CAPSULE; ORAL					
NIMOTOP					
	+ BAYER PHARMS	30MG	N18869	001	Dec 28, 1988

NISOLDIPINE

TABLET, EXTENDED RELEASE; ORAL					
SULAR					
	+ SCIELE PHARMA INC	10MG	N20356	001	Feb 02, 1995
		20MG	N20356	002	Feb 02, 1995
		30MG	N20356	003	Feb 02, 1995
		40MG	N20356	004	Feb 02, 1995

NITAZOXANIDE

FOR SUSPENSION; ORAL					
ALINIA					
	+ ROMARK	100MG/5ML	N21498	001	Nov 22, 2002
TABLET; ORAL					
ALINIA					
	+ ROMARK	500MG	N21497	001	Jul 21, 2004

NITISINONE

CAPSULE; ORAL					
ORFADIN					
	SWEDISH ORPHAN	2MG	N21232	001	Jan 18, 2002
		5MG	N21232	002	Jan 18, 2002
		10MG	N21232	003	Jan 18, 2002

NITRIC OXIDE

GAS; INHALATION					
INOMAX					
	INO	100PPM	N20845	002	Dec 23, 1999
		800PPM	N20845	003	Dec 23, 1999

NITROFURANTOIN

SUSPENSION; ORAL					
FURADANTIN					
	+ SCIELE PHARMA INC	25MG/5ML	N09175	001	

NITROFURANTOIN, MACROCRYSTALLINE

CAPSULE; ORAL					
MACRODANTIN					
AB	PROCTER AND GAMBLE	25MG	N16620	003	
AB		50MG	N16620	001	
AB +		100MG	N16620	002	
NITROFURANTOIN					
AB	IVAX PHARMS	50MG	N73671	001	Jan 28, 1993

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 277 (of 370)

NITROFURANTOIN, MACROCRYSTALLINE

CAPSULE; ORAL

NITROFURANTOIN

<u>AB</u>	IVAX PHARMS	<u>100MG</u>	<u>N73652</u>	<u>001</u>	Jan 28, 1993
<u>AB</u>	MYLAN	<u>50MG</u>	<u>N74967</u>	<u>001</u>	Jul 09, 1997
<u>AB</u>		<u>100MG</u>	<u>N77025</u>	<u>001</u>	Aug 18, 2004
<u>AB</u>	SANDOZ	<u>25MG</u>	<u>N74336</u>	<u>001</u>	Jan 25, 1995
<u>AB</u>		<u>50MG</u>	<u>N74336</u>	<u>002</u>	Jan 25, 1995
<u>AB</u>		<u>100MG</u>	<u>N74336</u>	<u>003</u>	Jan 25, 1995
<u>AB</u>	WATSON LABS	<u>25MG</u>	<u>N73696</u>	<u>001</u>	Dec 31, 1992
<u>AB</u>		<u>50MG</u>	<u>N73696</u>	<u>002</u>	Dec 31, 1992
<u>AB</u>		<u>100MG</u>	<u>N73696</u>	<u>003</u>	Dec 31, 1992

NITROFURANTOIN; NITROFURANTOIN, MACROCRYSTALLINE

CAPSULE; ORAL

MACROBID

<u>AB</u>	+ PROCTER AND GAMBLE	<u>75MG;25MG</u>	<u>N20064</u>	<u>001</u>	Dec 24, 1991
	<u>NITROFURANTOIN (MONOHYDRATE/MACROCRYSTALS)</u>				
<u>AB</u>	MYLAN	<u>75MG;25MG</u>	<u>N76648</u>	<u>001</u>	Mar 22, 2004
<u>AB</u>	RANBAXY	<u>75MG;25MG</u>	<u>N76951</u>	<u>001</u>	Mar 30, 2005
<u>AB</u>	SANDOZ	<u>75MG;25MG</u>	<u>N77066</u>	<u>001</u>	Apr 05, 2005

NITROFURAZONE

OINTMENT; TOPICAL

NITROFURAZONE

+ TARO	0.2%	N86156	001	
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NITROGLYCERIN

AEROSOL, METERED; SUBLINGUAL

NITROMIST

+ NOVADEL	0.4MG/SPRAY	N21780	001	Nov 02, 2006
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FILM, EXTENDED RELEASE; TRANSDERMAL

MINITRAN

<u>AB1</u>	3M	<u>0.1MG/HR</u>	<u>N89771</u>	<u>001</u>	Aug 30, 1996
<u>AB1</u>		<u>0.2MG/HR</u>	<u>N89772</u>	<u>001</u>	Aug 30, 1996
<u>AB1</u>		<u>0.4MG/HR</u>	<u>N89773</u>	<u>001</u>	Aug 30, 1996
<u>AB1</u>		<u>0.6MG/HR</u>	<u>N89774</u>	<u>001</u>	Aug 30, 1996

NITRO-DUR

<u>AB1</u>	+ KEY PHARMS	<u>0.1MG/HR</u>	<u>N20145</u>	<u>001</u>	Apr 04, 1995
<u>AB1</u>	+	<u>0.2MG/HR</u>	<u>N20145</u>	<u>002</u>	Apr 04, 1995
<u>AB1</u>	+	<u>0.4MG/HR</u>	<u>N20145</u>	<u>004</u>	Apr 04, 1995
<u>AB1</u>	+	<u>0.6MG/HR</u>	<u>N20145</u>	<u>005</u>	Apr 04, 1995
BX	+	0.8MG/HR	N20145	006	Apr 04, 1995
	+	0.3MG/HR	N20145	003	Apr 04, 1995

NITROGLYCERIN

<u>AB2</u>	HERCON LABS	<u>0.2MG/HR</u>	<u>N89884</u>	<u>001</u>	Oct 30, 1998
<u>AB2</u>		<u>0.4MG/HR</u>	<u>N89885</u>	<u>001</u>	Oct 30, 1998
<u>AB2</u>		<u>0.6MG/HR</u>	<u>N89886</u>	<u>001</u>	Oct 30, 1998
<u>AB1</u>	KREMERS URBAN	<u>0.2MG/HR</u>	<u>N75115</u>	<u>001</u>	Aug 10, 2004
<u>AB1</u>		<u>0.4MG/HR</u>	<u>N75115</u>	<u>002</u>	Aug 10, 2004
<u>AB1</u>	MYLAN TECHNOLOGIES	<u>0.1MG/HR</u>	<u>N74992</u>	<u>004</u>	Nov 12, 1999
<u>AB1</u>		<u>0.2MG/HR</u>	<u>N74992</u>	<u>003</u>	Nov 12, 1999
<u>AB1</u>		<u>0.4MG/HR</u>	<u>N74992</u>	<u>002</u>	Nov 12, 1999
<u>AB1</u>		<u>0.6MG/HR</u>	<u>N74992</u>	<u>001</u>	Nov 12, 1999
<u>AB2</u>	+	<u>0.1MG/HR</u>	<u>N74559</u>	<u>004</u>	Feb 06, 1998
<u>AB2</u>	+	<u>0.2MG/HR</u>	<u>N74559</u>	<u>003</u>	Aug 30, 1996
<u>AB2</u>	+	<u>0.4MG/HR</u>	<u>N74559</u>	<u>002</u>	Aug 30, 1996
<u>AB2</u>	+	<u>0.6MG/HR</u>	<u>N74559</u>	<u>001</u>	Aug 30, 1996

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 278 (of 370)

NITROGLYCERIN

INJECTABLE; INJECTION

NITROGLYCERIN

<u>AP</u>	+	HOSPIRA	<u>5MG/ML</u>	<u>N18531</u>	<u>001</u>	
		+ LUITPOLD	5MG/ML	N72034	001	May 24, 1988

NITROGLYCERIN IN DEXTROSE 5%

<u>AP</u>	+	BAXTER HLTHCARE	<u>10MG/100ML</u>	<u>N19970</u>	<u>001</u>	Dec 29, 1989
<u>AP</u>	+		<u>20MG/100ML</u>	<u>N19970</u>	<u>002</u>	Dec 29, 1989
<u>AP</u>	+		<u>40MG/100ML</u>	<u>N19970</u>	<u>003</u>	Dec 29, 1989
<u>AP</u>		HOSPIRA	<u>10MG/100ML</u>	<u>N71846</u>	<u>001</u>	Aug 31, 1990
<u>AP</u>			<u>20MG/100ML</u>	<u>N71847</u>	<u>001</u>	Aug 31, 1990
<u>AP</u>			<u>40MG/100ML</u>	<u>N71848</u>	<u>001</u>	Aug 31, 1990

OINTMENT; TRANSDERMAL

NITROGLYCERIN

	+	FOUGERA	2%	N87355	001	Jul 08, 1988
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SPRAY, METERED; SUBLINGUAL

NITROLINGUAL PUMPSPRAY

	+	POHL BOSKAMP	0.4MG/SPRAY	N18705	002	Jan 10, 1997
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TABLET; SUBLINGUAL

NITROSTAT

		PFIZER PHARMS	0.3MG	N21134	001	May 01, 2000
			0.4MG	N21134	002	May 01, 2000
	+		0.6MG	N21134	003	May 01, 2000

NIZATIDINE

CAPSULE; ORAL

AXID

<u>AB</u>		RELIANT PHARMS	<u>150MG</u>	<u>N19508</u>	<u>001</u>	Apr 12, 1988
<u>AB</u>	+		<u>300MG</u>	<u>N19508</u>	<u>002</u>	Apr 12, 1988

NIZATIDINE

<u>AB</u>		DR REDDYS LABS LTD	<u>150MG</u>	<u>N77314</u>	<u>001</u>	Sep 15, 2005
<u>AB</u>			<u>300MG</u>	<u>N77314</u>	<u>002</u>	Sep 15, 2005
<u>AB</u>		GENPHARM	<u>150MG</u>	<u>N75934</u>	<u>001</u>	Jul 09, 2002
<u>AB</u>			<u>300MG</u>	<u>N75934</u>	<u>002</u>	Jul 09, 2002
<u>AB</u>		MYLAN	<u>150MG</u>	<u>N75806</u>	<u>001</u>	Jul 05, 2002
<u>AB</u>			<u>300MG</u>	<u>N75806</u>	<u>002</u>	Jul 05, 2002
<u>AB</u>		SANDOZ	<u>150MG</u>	<u>N76178</u>	<u>001</u>	Jul 05, 2002
<u>AB</u>			<u>300MG</u>	<u>N76178</u>	<u>002</u>	Jul 05, 2002
<u>AB</u>		TEVA	<u>150MG</u>	<u>N75668</u>	<u>001</u>	Sep 12, 2002
<u>AB</u>			<u>300MG</u>	<u>N75668</u>	<u>002</u>	Sep 12, 2002
<u>AB</u>		TORPHARM	<u>150MG</u>	<u>N76383</u>	<u>001</u>	Jan 23, 2003
<u>AB</u>			<u>300MG</u>	<u>N76383</u>	<u>002</u>	Jan 23, 2003
<u>AB</u>		WATSON LABS	<u>150MG</u>	<u>N75616</u>	<u>001</u>	Jul 09, 2002
<u>AB</u>			<u>300MG</u>	<u>N75616</u>	<u>002</u>	Jul 09, 2002
<u>AB</u>		ZENITH GOLDLINE	<u>150MG</u>	<u>N75461</u>	<u>001</u>	Jul 08, 2002
<u>AB</u>			<u>300MG</u>	<u>N75461</u>	<u>002</u>	Jul 08, 2002

SOLUTION; ORAL

AXID

	+	BRAINTREE	15MG/ML	N21494	001	May 25, 2004
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NOREPINEPHRINE BITARTRATE

INJECTABLE; INJECTION

LEVOPHED

<u>AP</u>	+	HOSPIRA	<u>EQ 1MG BASE/ML</u>	<u>N07513</u>	<u>001</u>	
<u>AP</u>		BEDFORD	<u>EQ 1MG BASE/ML</u>	<u>N40462</u>	<u>001</u>	Oct 31, 2003
<u>AP</u>		METRICS PHARM	<u>EQ 1MG BASE/ML</u>	<u>N40522</u>	<u>001</u>	Sep 30, 2004
<u>AP</u>		SICOR PHARMS	<u>EQ 1MG BASE/ML</u>	<u>N40455</u>	<u>001</u>	Mar 03, 2003

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 279 (of 370)

NORETHINDRONE

TABLET; ORAL-28

<u>AB1</u>	<u>CAMILA</u> BARR	<u>0.35MG</u>	<u>N76177</u>	<u>001</u>	Oct 21, 2002
<u>AB2</u>	<u>ERRIN</u> BARR	<u>0.35MG</u>	<u>N76225</u>	<u>001</u>	Oct 21, 2002
<u>AB2</u>	<u>MICRONOR</u> + ORTHO MCNEIL PHARM	<u>0.35MG</u>	<u>N16954</u>	<u>001</u>	
<u>AB1</u>	<u>NOR-QD</u> + WATSON LABS (UTAH)	<u>0.35MG</u>	<u>N17060</u>	<u>001</u>	

NORETHINDRONE ACETATE

TABLET; ORAL

<u>AB</u>	<u>AYGESTIN</u> + DURAMED PHARMS BARR	<u>5MG</u>	<u>N18405</u>	<u>001</u>	Apr 21, 1982
<u>AB</u>	<u>NORETHINDRONE ACETATE</u> BARR	<u>5MG</u>	<u>N75951</u>	<u>001</u>	May 25, 2001

NORFLOXACIN

TABLET; ORAL

NOROXIN

+	MERCK	400MG	N19384	002	Oct 31, 1986
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NORTRIPTYLINE HYDROCHLORIDE

CAPSULE; ORAL

	<u>NORTRIPTYLINE HYDROCHLORIDE</u>				
<u>AB</u>	MYLAN	<u>EQ 10MG BASE</u>	<u>N74234</u>	<u>001</u>	Jul 26, 1993
<u>AB</u>		<u>EQ 25MG BASE</u>	<u>N74234</u>	<u>002</u>	Jul 26, 1993
<u>AB</u>		<u>EQ 50MG BASE</u>	<u>N74234</u>	<u>003</u>	Jul 26, 1993
<u>AB</u>		<u>EQ 75MG BASE</u>	<u>N74234</u>	<u>004</u>	Jul 26, 1993
<u>AB</u>	SANDOZ	<u>EQ 10MG BASE</u>	<u>N74054</u>	<u>001</u>	Dec 31, 1992
<u>AB</u>		<u>EQ 10MG BASE</u>	<u>N74835</u>	<u>001</u>	Jun 30, 1997
<u>AB</u>		<u>EQ 25MG BASE</u>	<u>N74054</u>	<u>002</u>	Dec 31, 1992
<u>AB</u>		<u>EQ 25MG BASE</u>	<u>N74835</u>	<u>002</u>	Jun 30, 1997
<u>AB</u>		<u>EQ 50MG BASE</u>	<u>N74054</u>	<u>003</u>	Dec 31, 1992
<u>AB</u>		<u>EQ 50MG BASE</u>	<u>N74835</u>	<u>003</u>	Jun 30, 1997
<u>AB</u>		<u>EQ 75MG BASE</u>	<u>N74054</u>	<u>004</u>	Dec 31, 1992
<u>AB</u>		<u>EQ 75MG BASE</u>	<u>N74835</u>	<u>004</u>	Jun 30, 1997
<u>AB</u>	TARO	<u>EQ 10MG BASE</u>	<u>N75520</u>	<u>004</u>	May 08, 2000
<u>AB</u>		<u>EQ 25MG BASE</u>	<u>N75520</u>	<u>003</u>	May 08, 2000
<u>AB</u>		<u>EQ 50MG BASE</u>	<u>N75520</u>	<u>001</u>	May 08, 2000
<u>AB</u>		<u>EQ 75MG BASE</u>	<u>N75520</u>	<u>002</u>	May 08, 2000
<u>AB</u>	TEVA	<u>EQ 10MG BASE</u>	<u>N73667</u>	<u>001</u>	Apr 11, 1996
<u>AB</u>		<u>EQ 10MG BASE</u>	<u>N74132</u>	<u>001</u>	Mar 27, 1995
<u>AB</u>		<u>EQ 25MG BASE</u>	<u>N73667</u>	<u>002</u>	Apr 11, 1996
<u>AB</u>		<u>EQ 25MG BASE</u>	<u>N74132</u>	<u>002</u>	Mar 27, 1995
<u>AB</u>		<u>EQ 50MG BASE</u>	<u>N73667</u>	<u>003</u>	Apr 11, 1996
<u>AB</u>		<u>EQ 50MG BASE</u>	<u>N74132</u>	<u>003</u>	Mar 27, 1995
<u>AB</u>		<u>EQ 75MG BASE</u>	<u>N73667</u>	<u>004</u>	Apr 11, 1996
<u>AB</u>		<u>EQ 75MG BASE</u>	<u>N74132</u>	<u>004</u>	Mar 27, 1995
<u>AB</u>	WATSON LABS	<u>EQ 10MG BASE</u>	<u>N73553</u>	<u>001</u>	Mar 30, 1992
<u>AB</u>		<u>EQ 25MG BASE</u>	<u>N73554</u>	<u>001</u>	Mar 30, 1992
<u>AB</u>		<u>EQ 50MG BASE</u>	<u>N73555</u>	<u>001</u>	Mar 30, 1992
<u>AB</u>		<u>EQ 75MG BASE</u>	<u>N73556</u>	<u>001</u>	Mar 30, 1992
	<u>PAMELOR</u>				
<u>AB</u>	TYCO HLTHCARE	<u>EQ 10MG BASE</u>	<u>N18013</u>	<u>001</u>	
<u>AB</u>		<u>EQ 25MG BASE</u>	<u>N18013</u>	<u>002</u>	
<u>AB</u>		<u>EQ 50MG BASE</u>	<u>N18013</u>	<u>004</u>	
<u>AB</u>	+	<u>EQ 75MG BASE</u>	<u>N18013</u>	<u>003</u>	

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 280 (of 370)

NORTRIPTYLINE HYDROCHLORIDE

SOLUTION; ORAL

AVENTYL HYDROCHLORIDE

<u>AA</u>	+	RANBAXY	<u>EQ 10MG BASE/5ML</u>	<u>N14685</u>	<u>001</u>	
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NORTRIPTYLINE HYDROCHLORIDE

<u>AA</u>		PHARM ASSOC	<u>EQ 10MG BASE/5ML</u>	<u>N75606</u>	<u>001</u>	Aug 28, 2000
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<u>AA</u>		TARO	<u>EQ 10MG BASE/5ML</u>	<u>N77965</u>	<u>001</u>	Jun 20, 2006
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PAMELOR

<u>AA</u>		TYCO HLTHCARE	<u>EQ 10MG BASE/5ML</u>	<u>N18012</u>	<u>001</u>	
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NYSTATIN

CREAM; TOPICAL

MYCOSTATIN

<u>AT</u>	+	WESTWOOD SQUIBB	<u>100,000 UNITS/GM</u>	<u>N60575</u>	<u>001</u>	
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NYSTATIN

<u>AT</u>		ACTAVIS MID ATLANTIC	<u>100,000 UNITS/GM</u>	<u>N62949</u>	<u>001</u>	Jun 13, 1988
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<u>AT</u>		ALTANA	<u>100,000 UNITS/GM</u>	<u>N62129</u>	<u>001</u>	
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<u>AT</u>		PERRIGO NEW YORK	<u>100,000 UNITS/GM</u>	<u>N62225</u>	<u>001</u>	
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<u>AT</u>		TARO	<u>100,000 UNITS/GM</u>	<u>N64022</u>	<u>001</u>	Jan 29, 1993
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<u>AT</u>		VINTAGE	<u>100,000 UNITS/GM</u>	<u>N65315</u>	<u>001</u>	May 31, 2006
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OINTMENT; TOPICAL

NYSTATIN

<u>AT</u>		ACTAVIS MID ATLANTIC	<u>100,000 UNITS/GM</u>	<u>N62840</u>	<u>001</u>	Nov 13, 1987
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<u>AT</u>	+	ALTANA	<u>100,000 UNITS/GM</u>	<u>N62124</u>	<u>002</u>	Sep 23, 1982
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<u>AT</u>		PERRIGO NEW YORK	<u>100,000 UNITS/GM</u>	<u>N62472</u>	<u>001</u>	Feb 13, 1984
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POWDER; TOPICAL

MYCOSTATIN

<u>AT</u>	+	WESTWOOD SQUIBB	<u>100,000 UNITS/GM</u>	<u>N60578</u>	<u>001</u>	
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NYSTATIN

<u>AT</u>		KALI LABS	<u>100,000 UNITS/GM</u>	<u>N65138</u>	<u>001</u>	Jul 23, 2004
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<u>AT</u>		KV PHARM	<u>100,000 UNITS/GM</u>	<u>N65321</u>	<u>001</u>	Aug 18, 2006
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<u>AT</u>		PHARMAFORCE	<u>100,000 UNITS/GM</u>	<u>N65203</u>	<u>001</u>	Jul 15, 2004
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<u>AT</u>		UPSHER SMITH	<u>100,000 UNITS/GM</u>	<u>N65183</u>	<u>001</u>	May 03, 2005
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<u>AT</u>		X GEN PHARMS	<u>100,000 UNITS/GM</u>	<u>N65175</u>	<u>001</u>	Dec 17, 2004
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NYSTOP

<u>AT</u>		PADDOCK	<u>100,000 UNITS/GM</u>	<u>N64118</u>	<u>001</u>	Aug 16, 1996
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SUSPENSION; ORAL

NILSTAT

<u>AA</u>	+	GLENMARK PHARMA	<u>100,000 UNITS/ML</u>	<u>N50299</u>	<u>001</u>	
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NYSTATIN

<u>AA</u>		ACTAVIS MID ATLANTIC	<u>100,000 UNITS/ML</u>	<u>N62349</u>	<u>001</u>	Jul 14, 1982
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<u>AA</u>		BAUSCH AND LOMB	<u>100,000 UNITS/ML</u>	<u>N64042</u>	<u>001</u>	Feb 28, 1994
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<u>AA</u>		FOUGERA	<u>100,000 UNITS/ML</u>	<u>N62517</u>	<u>001</u>	Jun 07, 1984
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<u>AA</u>	+	MORTON GROVE	<u>100,000 UNITS/ML</u>	<u>N62512</u>	<u>001</u>	Oct 29, 1984
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<u>AA</u>		TARO	<u>100,000 UNITS/ML</u>	<u>N62876</u>	<u>001</u>	Feb 29, 1988
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<u>AA</u>		VINTAGE PHARMS	<u>100,000 UNITS/ML</u>	<u>N65148</u>	<u>001</u>	Jun 28, 2005
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<u>AA</u>		VISTAPHARM	<u>100,000 UNITS/ML</u>	<u>N64142</u>	<u>001</u>	Jun 25, 1998
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TABLET; ORAL

MYCOSTATIN

<u>AA</u>	+	APOTHECON	<u>500,000 UNITS</u>	<u>N60574</u>	<u>001</u>	
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NYSTATIN

<u>AA</u>		MUTUAL PHARM	<u>500,000 UNITS</u>	<u>N62838</u>	<u>001</u>	Dec 22, 1988
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<u>AA</u>		PAR PHARM	<u>500,000 UNITS</u>	<u>N62474</u>	<u>001</u>	Dec 22, 1983
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<u>AA</u>		TEVA	<u>500,000 UNITS</u>	<u>N62506</u>	<u>001</u>	Jan 16, 1984
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TABLET; VAGINAL

NYSTATIN

	+	ODYSSEY PHARMS	100,000 UNITS	N62615	001	Oct 17, 1985
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27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 281 (of 370)

NYSTATIN; TRIAMCINOLONE ACETONIDE

CREAM; TOPICAL

MYKACET

<u>AT</u>	ACTAVIS MID ATLANTIC	<u>100,000 UNITS/GM;0.1%</u>	<u>N62367</u>	<u>001</u>	May 28, 1985
	<u>NYSTATIN AND TRIAMCINOLONE ACETONIDE</u>				
<u>AT</u>	+ TARO	<u>100,000 UNITS/GM;0.1%</u>	<u>N62364</u>	<u>001</u>	Dec 22, 1987
	<u>NYSTATIN-TRIAMCINOLONE ACETONIDE</u>				
<u>AT</u>	FOUGERA	<u>100,000 UNITS/GM;0.1%</u>	<u>N62599</u>	<u>001</u>	Oct 08, 1985

OINTMENT; TOPICAL

MYKACET

<u>AT</u>	ACTAVIS MID ATLANTIC	<u>100,000 UNITS/GM;0.1%</u>	<u>N62733</u>	<u>001</u>	Mar 09, 1987
	<u>NYSTATIN AND TRIAMCINOLONE ACETONIDE</u>				
<u>AT</u>	+ TARO	<u>100,000 UNITS/GM;0.1%</u>	<u>N63305</u>	<u>001</u>	Mar 29, 1993
	<u>NYSTATIN-TRIAMCINOLONE ACETONIDE</u>				
<u>AT</u>	FOUGERA	<u>100,000 UNITS/GM;0.1%</u>	<u>N62602</u>	<u>001</u>	Oct 09, 1985

OCTREOTIDE ACETATE

INJECTABLE; INJECTION

OCTREOTIDE ACETATE

<u>AP</u>	ABRAXIS PHARM	<u>EQ 0.2MG BASE/ML</u>	<u>N77450</u>	<u>001</u>	Feb 10, 2006
<u>AP</u>		<u>EQ 1MG BASE/ML</u>	<u>N77450</u>	<u>002</u>	Feb 10, 2006
<u>AP</u>	BEDFORD	<u>EQ 0.2MG BASE/ML</u>	<u>N76330</u>	<u>001</u>	Apr 08, 2005
<u>AP</u>		<u>EQ 1MG BASE/ML</u>	<u>N76330</u>	<u>002</u>	Apr 08, 2005
<u>AP</u>	SICOR PHARMS	<u>EQ 0.05MG BASE/ML</u>	<u>N75957</u>	<u>001</u>	Oct 03, 2005
<u>AP</u>		<u>EQ 0.1MG BASE/ML</u>	<u>N75957</u>	<u>002</u>	Oct 03, 2005
<u>AP</u>		<u>EQ 0.2MG BASE/ML</u>	<u>N75959</u>	<u>001</u>	Nov 21, 2005
<u>AP</u>		<u>EQ 0.5MG BASE/ML</u>	<u>N75957</u>	<u>003</u>	Oct 03, 2005
<u>AP</u>		<u>EQ 1MG BASE/ML</u>	<u>N75959</u>	<u>002</u>	Nov 21, 2005

OCTREOTIDE ACETATE (PRESERVATIVE FREE)

<u>AP</u>	ABRAXIS PHARM	<u>EQ 0.05MG BASE/ML</u>	<u>N77457</u>	<u>001</u>	Feb 10, 2006
<u>AP</u>		<u>EQ 0.1MG BASE/ML</u>	<u>N77457</u>	<u>002</u>	Feb 10, 2006
<u>AP</u>		<u>EQ 0.5MG BASE/ML</u>	<u>N77457</u>	<u>003</u>	Feb 10, 2006
<u>AP</u>	BEDFORD	<u>EQ 0.05MG BASE/ML</u>	<u>N76313</u>	<u>001</u>	Mar 28, 2005
<u>AP</u>		<u>EQ 0.1MG BASE/ML</u>	<u>N76313</u>	<u>003</u>	Mar 28, 2005
<u>AP</u>		<u>EQ 0.5MG BASE/ML</u>	<u>N76313</u>	<u>002</u>	Mar 28, 2005

SANDOSTATIN

<u>AP</u>	+ NOVARTIS	<u>EQ 0.05MG BASE/ML</u>	<u>N19667</u>	<u>001</u>	Oct 21, 1988
<u>AP</u>		<u>EQ 0.1MG BASE/ML</u>	<u>N19667</u>	<u>002</u>	Oct 21, 1988
<u>AP</u>		<u>EQ 0.2MG BASE/ML</u>	<u>N19667</u>	<u>004</u>	Jun 12, 1991
<u>AP</u>		<u>EQ 0.5MG BASE/ML</u>	<u>N19667</u>	<u>003</u>	Oct 21, 1988
<u>AP</u>		<u>EQ 1MG BASE/ML</u>	<u>N19667</u>	<u>005</u>	Jun 12, 1991
	SANDOSTATIN LAR				
	NOVARTIS	EQ 10MG BASE/VIAL	N21008	001	Nov 25, 1998
		EQ 20MG BASE/VIAL	N21008	002	Nov 25, 1998
	+	EQ 30MG BASE/VIAL	N21008	003	Nov 25, 1998

OFLOXACIN

INJECTABLE; INJECTION

OFLOXACIN

	+ BEDFORD	40MG/ML	N75762	001	Jan 16, 2002
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SOLUTION/DROPS; OPHTHALMIC

OCUFLOX

<u>AT</u>	+ ALLERGAN	<u>0.3%</u>	<u>N19921</u>	<u>001</u>	Jul 30, 1993
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OFLOXACIN

<u>AT</u>	ALCON	<u>0.3%</u>	<u>N76231</u>	<u>001</u>	May 14, 2004
<u>AT</u>	BAUSCH AND LOMB	<u>0.3%</u>	<u>N76622</u>	<u>001</u>	May 14, 2004
<u>AT</u>	HI TECH PHARMA	<u>0.3%</u>	<u>N76615</u>	<u>001</u>	May 14, 2004
<u>AT</u>	NOVEX	<u>0.3%</u>	<u>N76513</u>	<u>001</u>	May 14, 2004
<u>AT</u>	PHARMAFORCE	<u>0.3%</u>	<u>N76830</u>	<u>001</u>	Aug 31, 2004

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 282 (of 370)

OFLOXACIN

SOLUTION/DROPS; OTIC

FLOXIN OTIC

+	DAIICHI	0.3%	N20799	001	Dec 16, 1997
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TABLET; ORAL

FLOXIN

<u>AB</u>	ORTHO MCNEIL PHARM	<u>200MG</u>	<u>N19735</u>	<u>001</u>	Dec 28, 1990
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<u>AB</u>		<u>300MG</u>	<u>N19735</u>	<u>002</u>	Dec 28, 1990
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<u>AB</u>	+	<u>400MG</u>	<u>N19735</u>	<u>003</u>	Dec 28, 1990
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OFLOXACIN

<u>AB</u>	DR REDDYS LABS LTD	<u>200MG</u>	<u>N77098</u>	<u>001</u>	Feb 10, 2006
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<u>AB</u>		<u>300MG</u>	<u>N77098</u>	<u>002</u>	Feb 10, 2006
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<u>AB</u>		<u>400MG</u>	<u>N77098</u>	<u>003</u>	Feb 10, 2006
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<u>AB</u>	PAR PHARM	<u>200MG</u>	<u>N76093</u>	<u>001</u>	Sep 02, 2003
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<u>AB</u>		<u>300MG</u>	<u>N76093</u>	<u>002</u>	Sep 02, 2003
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<u>AB</u>		<u>400MG</u>	<u>N76093</u>	<u>003</u>	Sep 02, 2003
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<u>AB</u>	RANBAXY	<u>200MG</u>	<u>N76220</u>	<u>001</u>	Sep 02, 2003
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<u>AB</u>		<u>300MG</u>	<u>N76220</u>	<u>002</u>	Sep 02, 2003
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<u>AB</u>		<u>400MG</u>	<u>N76220</u>	<u>003</u>	Sep 02, 2003
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<u>AB</u>	TEVA	<u>200MG</u>	<u>N76182</u>	<u>001</u>	Sep 02, 2003
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<u>AB</u>		<u>300MG</u>	<u>N76182</u>	<u>002</u>	Sep 02, 2003
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<u>AB</u>		<u>400MG</u>	<u>N76182</u>	<u>003</u>	Sep 02, 2003
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OLANZAPINE

INJECTABLE; INTRAMUSCULAR

ZYPREXA

+	LILLY	10MG/VIAL	N21253	001	Mar 29, 2004
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TABLET; ORAL

ZYPREXA

	LILLY	2.5MG	N20592	001	Sep 30, 1996
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+		5MG	N20592	002	Sep 30, 1996
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		7.5MG	N20592	003	Sep 30, 1996
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		10MG	N20592	004	Sep 30, 1996
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		15MG	N20592	005	Sep 09, 1997
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		20MG	N20592	006	Sep 09, 1997
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TABLET, ORALLY DISINTEGRATING; ORAL

ZYPREXA ZYDIS

+	LILLY	5MG	N21086	001	Apr 06, 2000
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		10MG	N21086	002	Apr 06, 2000
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		15MG	N21086	003	Apr 06, 2000
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		20MG	N21086	004	Apr 06, 2000
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OLMESARTAN MEDOXOMIL

TABLET; ORAL

BENICAR

	DAIICHI SANKYO	5MG	N21286	001	Apr 25, 2002
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		20MG	N21286	003	Apr 25, 2002
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+		40MG	N21286	004	Apr 25, 2002
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OLOPATADINE HYDROCHLORIDE

SOLUTION/DROPS; OPHTHALMIC

+	ALCON	0.2%	N21545	001	Dec 22, 2004
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PATANOL

+	ALCON	EQ 0.1% BASE	N20688	001	Dec 18, 1996
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27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 283 (of 370)

OLSALAZINE SODIUMCAPSULE; ORAL
DIPENTUM

+	UCB INC	250MG	N19715	001	Jul 31, 1990
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OMEGA-3-ACID ETHYL ESTERSCAPSULE; ORAL
OMACOR

+	RELIANT PHARMS INC	1GM	N21654	001	Nov 10, 2004
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OMEPRAZOLE

CAPSULE, DELAYED REL PELLETS; ORAL

OMEPRAZOLE

<u>AB</u>	IMPAX LABS	<u>10MG</u>	<u>N75785</u>	<u>001</u>	Nov 08, 2002
<u>AB</u>		<u>20MG</u>	<u>N75785</u>	<u>002</u>	Nov 08, 2002
<u>AB</u>	KREMERS URBAN DEV	<u>10MG</u>	<u>N75410</u>	<u>001</u>	Nov 01, 2002
<u>AB</u>		<u>20MG</u>	<u>N75410</u>	<u>002</u>	Nov 01, 2002
<u>AB</u>	LEK PHARMS	<u>10MG</u>	<u>N75757</u>	<u>001</u>	Jan 28, 2003
<u>AB</u>		<u>20MG</u>	<u>N75757</u>	<u>002</u>	Jan 28, 2003
<u>AB</u>	MYLAN	<u>10MG</u>	<u>N75876</u>	<u>001</u>	May 29, 2003
<u>AB</u>		<u>20MG</u>	<u>N75876</u>	<u>002</u>	May 29, 2003
<u>AB</u>	TORPHARM	<u>10MG</u>	<u>N76048</u>	<u>001</u>	Oct 06, 2003
<u>AB</u>		<u>20MG</u>	<u>N76048</u>	<u>002</u>	Oct 06, 2003
	<u>PRILOSEC</u>				
<u>AB</u>	ASTRAZENECA	<u>10MG</u>	<u>N19810</u>	<u>003</u>	Oct 05, 1995
<u>AB</u>	+	<u>20MG</u>	<u>N19810</u>	<u>001</u>	Sep 14, 1989
	+	40MG	N19810	002	Jan 15, 1998

OMEPRAZOLE; SODIUM BICARBONATECAPSULE; ORAL
ZEGERID

	SANTARUS	20MG;1.1GM	N21849	001	Feb 27, 2006
+		40MG;1.1GM	N21849	002	Feb 27, 2006

FOR SUSPENSION; ORAL
ZEGERID

	SANTARUS	20MG/PACKET;1.68GM/PACKET	N21636	001	Jun 15, 2004
+		40MG/PACKET;1.68GM/PACKET	N21706	001	Dec 21, 2004

ONDANSETRON

TABLET, ORALLY DISINTEGRATING; ORAL

ONDANSETRON

<u>AB</u>	KALI LABS	<u>4MG</u>	<u>N76506</u>	<u>001</u>	Dec 26, 2006
<u>AB</u>		<u>8MG</u>	<u>N76506</u>	<u>002</u>	Dec 26, 2006
		16MG	N77406	001	Dec 26, 2006
		24MG	N77406	002	Dec 26, 2006
	<u>ZOFRAN ODT</u>				
<u>AB</u>	GLAXOSMITHKLINE	<u>4MG</u>	<u>N20781</u>	<u>001</u>	Jan 27, 1999
<u>AB</u>	+	<u>8MG</u>	<u>N20781</u>	<u>002</u>	Jan 27, 1999

ONDANSETRON HYDROCHLORIDE

INJECTABLE; INJECTION

ONDANSETRON HYDROCHLORIDE

<u>AP</u>	ABRAXIS PHARM	<u>EQ 2MG BASE/ML</u>	<u>N76974</u>	<u>001</u>	Dec 26, 2006
<u>AP</u>	APOTEX	<u>EQ 2MG BASE/ML</u>	<u>N77368</u>	<u>001</u>	Dec 26, 2006
<u>AP</u>	BAXTER HLTHCARE	<u>EQ 2MG BASE/ML</u>	<u>N77365</u>	<u>001</u>	Dec 26, 2006
<u>AP</u>	BEDFORD	<u>EQ 2MG BASE/ML</u>	<u>N76967</u>	<u>001</u>	Dec 26, 2006
<u>AP</u>	HIKMA FARMACEUTICA	<u>EQ 2MG BASE/ML</u>	<u>N76781</u>	<u>001</u>	Dec 26, 2006
<u>AP</u>	HOSPIRA	<u>EQ 2MG BASE/ML</u>	<u>N77473</u>	<u>001</u>	Dec 26, 2006
<u>AP</u>	MAYNE PHARMA USA	<u>EQ 2MG BASE/ML</u>	<u>N76695</u>	<u>001</u>	Dec 26, 2006

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 284 (of 370)

ONDANSETRON HYDROCHLORIDE

INJECTABLE; INJECTION

ONDANSETRON HYDROCHLORIDE

<u>AP</u>	PHARMAFORCE	<u>EQ 2MG BASE/ML</u>	<u>N77582</u>	<u>001</u>	Dec 26, 2006
<u>AP</u>	PLIVA HRVATSKA DOO	<u>EQ 2MG BASE/ML</u>	<u>N77544</u>	<u>001</u>	Dec 26, 2006
<u>AP</u>	SUN PHARM INDS (IN)	<u>EQ 2MG BASE/ML</u>	<u>N77172</u>	<u>001</u>	Dec 26, 2006
<u>AP</u>	TEVA	<u>EQ 2MG BASE/ML</u>	<u>N76876</u>	<u>001</u>	Nov 22, 2006
<u>AP</u>	WOCKHARDT	<u>EQ 2MG BASE/ML</u>	<u>N77577</u>	<u>001</u>	Dec 26, 2006

ONDANSETRON HYDROCHLORIDE AND DEXTROSE IN PLASTIC CONTAINER

<u>AP</u>	SICOR PHARMS	<u>EQ 0.64MG BASE/ML</u>	<u>N77480</u>	<u>001</u>	Nov 22, 2006
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ONDANSETRON HYDROCHLORIDE AND SODIUM CHLORIDE IN PLASTIC CONTAINER

<u>AP</u> +	BAXTER HLTHCARE	<u>EQ 0.64MG BASE/ML</u>	<u>N21915</u>	<u>002</u>	Dec 27, 2006
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ONDANSETRON HYDROCHLORIDE PRESERVATIVE FREE

<u>AP</u>	ABRAXIS PHARM	<u>EQ 2MG BASE/ML</u>	<u>N76972</u>	<u>001</u>	Dec 26, 2006
<u>AP</u>	APOTEX INC	<u>EQ 2MG BASE/ML</u>	<u>N77343</u>	<u>001</u>	Dec 26, 2006
<u>AP</u>	BAXTER HLTHCARE	<u>EQ 2MG BASE/ML</u>	<u>N77541</u>	<u>001</u>	Dec 26, 2006
<u>AP</u>	BEDFORD LABS	<u>EQ 2MG BASE/ML</u>	<u>N77011</u>	<u>001</u>	Dec 26, 2006
<u>AP</u>	HIKMA FARMACEUTICA	<u>EQ 2MG BASE/ML</u>	<u>N76780</u>	<u>001</u>	Dec 26, 2006
<u>AP</u>	HOSPIRA	<u>EQ 2MG BASE/ML</u>	<u>N77548</u>	<u>001</u>	Dec 26, 2006
<u>AP</u>	MAYNE PHARMA USA	<u>EQ 2MG BASE/ML</u>	<u>N76696</u>	<u>001</u>	Dec 26, 2006
<u>AP</u>	PHARMAFORCE	<u>EQ 2MG BASE/ML</u>	<u>N77387</u>	<u>001</u>	Dec 26, 2006
<u>AP</u>	SUN PHARM INDS LTD	<u>EQ 2MG BASE/ML</u>	<u>N77173</u>	<u>001</u>	Dec 26, 2006
<u>AP</u>	TEVA	<u>EQ 2MG BASE/ML</u>	<u>N76759</u>	<u>001</u>	Nov 22, 2006
<u>AP</u>	WOCKHARDT	<u>EQ 2MG BASE/ML</u>	<u>N77716</u>	<u>001</u>	Dec 26, 2006

ZOFRAN

<u>AP</u> +	GLAXOSMITHKLINE	<u>EQ 2MG BASE/ML</u>	<u>N20007</u>	<u>001</u>	Jan 04, 1991
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ZOFRAN IN PLASTIC CONTAINER

<u>AP</u> +	GLAXOSMITHKLINE	<u>EQ 0.64MG BASE/ML</u>	<u>N20403</u>	<u>001</u>	Jan 31, 1995
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ZOFRAN PRESERVATIVE FREE

<u>AP</u> +	GLAXOSMITHKLINE	<u>EQ 2MG BASE/ML</u>	<u>N20007</u>	<u>003</u>	Dec 10, 1993
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SOLUTION; ORAL

ONDANSETRON HYDROCHLORIDE

<u>AA</u>	ROXANE	<u>EQ 4MG BASE/5ML</u>	<u>N76960</u>	<u>001</u>	Dec 26, 2006
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ZOFRAN

<u>AA</u> +	GLAXOSMITHKLINE	<u>EQ 4MG BASE/5ML</u>	<u>N20605</u>	<u>001</u>	Jan 24, 1997
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TABLET; ORAL

ONDANSETRON HYDROCHLORIDE

<u>AB</u>	DR REDDYS LABS LTD	<u>EQ 4MG BASE</u>	<u>N76183</u>	<u>003</u>	Dec 26, 2006
<u>AB</u>		<u>EQ 8MG BASE</u>	<u>N76183</u>	<u>002</u>	Dec 26, 2006
<u>AB</u>		<u>EQ 24MG BASE</u>	<u>N76183</u>	<u>001</u>	Dec 26, 2006
		EQ 16MG BASE	N76559	001	Dec 26, 2006

ZOFRAN

<u>AB</u>	GLAXOSMITHKLINE	<u>EQ 4MG BASE</u>	<u>N20103</u>	<u>001</u>	Dec 31, 1992
<u>AB</u>		<u>EQ 8MG BASE</u>	<u>N20103</u>	<u>002</u>	Dec 31, 1992
<u>AB</u> +		<u>EQ 24MG BASE</u>	<u>N20103</u>	<u>003</u>	Aug 27, 1999

ORLISTAT

CAPSULE; ORAL

XENICAL

+	HLR	120MG	N20766	001	Apr 23, 1999
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ORPHENADRINE CITRATE

INJECTABLE; INJECTION

NORFLEX

<u>AP</u> +	3M	<u>30MG/ML</u>	<u>N13055</u>	<u>001</u>	
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ORPHENADRINE CITRATE

<u>AP</u>	AKORN	<u>30MG/ML</u>	<u>N40484</u>	<u>001</u>	May 24, 2006
<u>AP</u>	BEDFORD LABS	<u>30MG/ML</u>	<u>N40463</u>	<u>001</u>	Mar 04, 2003
<u>AP</u>	WATSON LABS	<u>30MG/ML</u>	<u>N84779</u>	<u>001</u>	Mar 15, 1982
<u>AP</u>		<u>30MG/ML</u>	<u>N87062</u>	<u>001</u>	

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 285 (of 370)

ORPHENADRINE CITRATE

TABLET, EXTENDED RELEASE; ORAL

ORPHENADRINE CITRATE

<u>AB</u>	IMPAX PHARMS	<u>100MG</u>	<u>N40368</u>	<u>001</u>	Jun 23, 2000
<u>AB</u>	KIEL	<u>100MG</u>	<u>N40249</u>	<u>001</u>	Jan 29, 1999
<u>AB</u>	SANDOZ	<u>100MG</u>	<u>N40284</u>	<u>001</u>	Jun 19, 1998
<u>AB</u>	+	<u>100MG</u>	<u>N40327</u>	<u>001</u>	Feb 15, 2000

OSELTAMIVIR PHOSPHATE

CAPSULE; ORAL

TAMIFLU

+	ROCHE	EQ 75MG BASE	N21087	001	Oct 27, 1999
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FOR SUSPENSION; ORAL

TAMIFLU

+	ROCHE	EQ 12MG BASE/ML	N21246	001	Dec 14, 2000
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OXACILLIN SODIUM

CAPSULE; ORAL

BACTOCILL

<u>AB</u>	GLAXOSMITHKLINE	<u>EQ 250MG BASE</u>	<u>N61336</u>	<u>001</u>	
<u>AB</u>	+	<u>EQ 500MG BASE</u>	<u>N61336</u>	<u>002</u>	
<u>AB</u>	<u>OXACILLIN SODIUM</u>				
<u>AB</u>	TEVA	<u>EQ 250MG BASE</u>	<u>N62222</u>	<u>001</u>	
<u>AB</u>		<u>EQ 500MG BASE</u>	<u>N62222</u>	<u>002</u>	

FOR SOLUTION; ORAL

BACTOCILL

<u>AA</u>	GLAXOSMITHKLINE	<u>EQ 250MG BASE/5ML</u>	<u>N62321</u>	<u>001</u>	
<u>AA</u>	<u>OXACILLIN SODIUM</u>				
<u>AA</u>	TEVA	<u>EQ 250MG BASE/5ML</u>	<u>N62252</u>	<u>001</u>	

INJECTABLE; INJECTION

BACTOCILL IN PLASTIC CONTAINER

+	BAXTER HLTHCARE	EQ 20MG BASE/ML	N50640	001	Oct 26, 1989
+		EQ 40MG BASE/ML	N50640	002	Oct 26, 1989

OXACILLIN SODIUM

<u>AP</u>	MARSAM PHARMS LLC	<u>EQ 250MG BASE/VIAL</u>	<u>N62856</u>	<u>001</u>	Oct 26, 1988
<u>AP</u>		<u>EQ 500MG BASE/VIAL</u>	<u>N62856</u>	<u>002</u>	Oct 26, 1988
<u>AP</u>		<u>EQ 1GM BASE/VIAL</u>	<u>N62856</u>	<u>003</u>	Oct 26, 1988
<u>AP</u>		<u>EQ 2GM BASE/VIAL</u>	<u>N62856</u>	<u>004</u>	Oct 26, 1988
<u>AP</u>		<u>EQ 10GM BASE/VIAL</u>	<u>N62984</u>	<u>001</u>	Sep 29, 1988
		EQ 4GM BASE/VIAL	N62856	005	Oct 26, 1988
<u>AP</u>	+	<u>EQ 250MG BASE/VIAL</u>	<u>N61490</u>	<u>001</u>	
<u>AP</u>	+	<u>EQ 500MG BASE/VIAL</u>	<u>N61490</u>	<u>002</u>	
<u>AP</u>	+	<u>EQ 1GM BASE/VIAL</u>	<u>N61490</u>	<u>003</u>	
<u>AP</u>		<u>EQ 1GM BASE/VIAL</u>	<u>N62737</u>	<u>001</u>	Dec 23, 1986
<u>AP</u>	+	<u>EQ 2GM BASE/VIAL</u>	<u>N61490</u>	<u>004</u>	
<u>AP</u>		<u>EQ 2GM BASE/VIAL</u>	<u>N62737</u>	<u>002</u>	Dec 23, 1986
<u>AP</u>	+	<u>EQ 10GM BASE/VIAL</u>	<u>N61490</u>	<u>006</u>	May 09, 1991

OXALIPLATIN

INJECTABLE; IV (INFUSION)

+	SANOFI AVENTIS US	50MG/10ML (5MG/ML)	N21759	001	Jan 31, 2005
+		100MG/20ML (5MG/ML)	N21759	002	Jan 31, 2005
+		200MG/40ML (5MG/ML)	N21759	003	Nov 17, 2006

OXANDROLONE

TABLET; ORAL

OXANDRIN

<u>AB</u>	SAVIENT PHARMS	<u>2.5MG</u>	<u>N13718</u>	<u>001</u>	
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27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 286 (of 370)

OXANDROLONE

TABLET; ORAL

OXANDRIN

<u>AB</u>	+	SAVIENT PHARMS	<u>10MG</u>	<u>N13718</u>	<u>002</u>	Nov 05, 2001
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OXANDROLONE

<u>AB</u>		SANDOZ	<u>2.5MG</u>	<u>N76897</u>	<u>001</u>	Dec 01, 2006
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<u>AB</u>			<u>10MG</u>	<u>N76897</u>	<u>002</u>	Dec 01, 2006
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<u>AB</u>		UPSHER SMITH	<u>2.5MG</u>	<u>N76761</u>	<u>001</u>	Dec 01, 2006
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OXAPROZIN

TABLET; ORAL

DAYPRO

<u>AB</u>	+	GD SEARLE	<u>600MG</u>	<u>N18841</u>	<u>004</u>	Oct 29, 1992
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OXAPROZIN

<u>AB</u>		ACTAVIS ELIZABETH	<u>600MG</u>	<u>N75843</u>	<u>001</u>	Oct 03, 2001
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<u>AB</u>		APOTEX INC	<u>600MG</u>	<u>N75987</u>	<u>001</u>	Sep 02, 2004
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<u>AB</u>		CARACO	<u>600MG</u>	<u>N75844</u>	<u>001</u>	Jan 03, 2002
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<u>AB</u>		DR REDDYS LABS LTD	<u>600MG</u>	<u>N75855</u>	<u>001</u>	Jan 31, 2001
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<u>AB</u>		GENPHARM	<u>600MG</u>	<u>N75847</u>	<u>001</u>	Feb 28, 2001
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<u>AB</u>		IVAX PHARMS	<u>600MG</u>	<u>N75846</u>	<u>001</u>	May 13, 2002
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<u>AB</u>		MYLAN	<u>600MG</u>	<u>N75851</u>	<u>001</u>	Aug 17, 2001
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<u>AB</u>		SANDOZ	<u>600MG</u>	<u>N75842</u>	<u>001</u>	Apr 12, 2001
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<u>AB</u>			<u>600MG</u>	<u>N75845</u>	<u>001</u>	Jan 31, 2001
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<u>AB</u>			<u>600MG</u>	<u>N75850</u>	<u>001</u>	Apr 27, 2001
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<u>AB</u>		TEVA	<u>600MG</u>	<u>N75849</u>	<u>001</u>	Jul 03, 2002
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<u>AB</u>		WATSON LABS	<u>600MG</u>	<u>N75848</u>	<u>001</u>	Feb 09, 2001
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OXAZEPAM

CAPSULE; ORAL

OXAZEPAM

<u>AB</u>		ACTAVIS ELIZABETH	<u>10MG</u>	<u>N72251</u>	<u>001</u>	Apr 14, 1988
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<u>AB</u>			<u>15MG</u>	<u>N72252</u>	<u>001</u>	Apr 14, 1988
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<u>AB</u>			<u>30MG</u>	<u>N72253</u>	<u>001</u>	Apr 14, 1988
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<u>AB</u>		IVAX PHARMS	<u>10MG</u>	<u>N70943</u>	<u>001</u>	Aug 03, 1987
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<u>AB</u>			<u>15MG</u>	<u>N70944</u>	<u>001</u>	Aug 03, 1987
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<u>AB</u>	+		<u>30MG</u>	<u>N70945</u>	<u>001</u>	Aug 03, 1987
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<u>AB</u>		SANDOZ	<u>10MG</u>	<u>N71813</u>	<u>001</u>	Apr 19, 1988
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<u>AB</u>			<u>15MG</u>	<u>N71756</u>	<u>001</u>	Apr 19, 1988
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<u>AB</u>			<u>30MG</u>	<u>N71814</u>	<u>001</u>	Apr 19, 1988
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<u>AB</u>		WATSON LABS	<u>10MG</u>	<u>N72952</u>	<u>001</u>	Sep 28, 1990
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<u>AB</u>			<u>15MG</u>	<u>N72953</u>	<u>001</u>	Sep 28, 1990
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<u>AB</u>			<u>30MG</u>	<u>N72954</u>	<u>001</u>	Sep 28, 1990
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OXCARBAZEPINE

SUSPENSION; ORAL

TRILEPTAL

+	NOVARTIS	300MG/5ML	N21285	001	May 25, 2001
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TABLET; ORAL

TRILEPTAL

	NOVARTIS	150MG	N21014	001	Jan 14, 2000
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		300MG	N21014	002	Jan 14, 2000
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+		600MG	N21014	003	Jan 14, 2000
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OXICONAZOLE NITRATE

CREAM; TOPICAL

OXISTAT

+	ALTANA	EQ 1% BASE	N19828	001	Dec 30, 1988
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27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 287 (of 370)

OXICONAZOLE NITRATELOTION; TOPICAL
OXISTAT

+ ALTANA EQ 1% BASE N20209 001 Sep 30, 1992

OXTRIPHYLLINETABLET, EXTENDED RELEASE; ORAL
CHOLEDYL SA+ PARKE DAVIS 400MG N87863 001 May 24, 1983
+ WARNER CHILCOTT 600MG N86742 001OXYBUTYNINFILM, EXTENDED RELEASE; TRANSDERMAL
OXYTROL

+ WATSON LABS (UTAH) 3.9MG/24HR N21351 002 Feb 26, 2003

OXYBUTYNIN CHLORIDE

SYRUP; ORAL

DITROPANAA + ALZA 5MG/5ML N18211 001OXYBUTYNIN CHLORIDEAA MIKART 5MG/5ML N75039 001 Jan 29, 1999
AA MORTON GROVE 5MG/5ML N74868 001 Feb 12, 1997
AA NOVEX 5MG/5ML N74997 001 Oct 15, 1997
AA PHARM ASSOC 5MG/5ML N75137 001 Dec 18, 1998
AA SILARX 5MG/5ML N74520 001 Mar 29, 1996
AA VINTAGE PHARMS 5MG/5ML N76682 001 Dec 28, 2004

TABLET; ORAL

DITROPANAB + ALZA 5MG N17577 001OXYBUTYNIN CHLORIDEAB PLIVA 5MG N71655 001 Nov 14, 1988
AB USL PHARMA 5MG N74625 001 Jul 31, 1996
AB VINTAGE PHARMS 5MG N75079 001 Oct 31, 1997

TABLET, EXTENDED RELEASE; ORAL

DITROPAN XLAB ALZA 5MG N20897 001 Dec 16, 1998
AB 10MG N20897 002 Dec 16, 1998
AB + 15MG N20897 003 Jun 22, 1999OXYBUTYNIN CHLORIDEAB IMPAX PHARMS 15MG N76745 001 Nov 09, 2006
AB MYLAN 5MG N76702 001 Nov 09, 2006
AB 10MG N76644 001 Nov 09, 2006OXYCODONE HYDROCHLORIDE

TABLET; ORAL

OXYCODONE HYDROCHLORIDEAB ACTAVIS TOTOWA 15MG N76636 001 Feb 06, 2004
AB 30MG N76636 002 Feb 06, 2004
AB KV PHARM 15MG N77290 003 Dec 08, 2005
AB 30MG N77290 005 Dec 08, 2005
5MG N77290 001 Dec 08, 2005
10MG N77290 002 Dec 08, 2005
20MG N77290 004 Dec 08, 2005
AB MALLINCKRODT 15MG N76758 001 Jun 30, 2004
AB 30MG N76758 002 Jun 30, 2004ROXICODONEAB + XANODYNE PHARM 15MG N21011 001 Aug 31, 2000
AB 30MG N21011 002 Aug 31, 2000

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 288 (of 370)

OXYCODONE HYDROCHLORIDE

TABLET, EXTENDED RELEASE; ORAL					
<u>OXYCODONE HYDROCHLORIDE</u>					
<u>AB</u>	ENDO PHARMS	<u>10MG</u>	<u>N75923</u>	<u>001</u>	Mar 23, 2004
<u>AB</u>		<u>20MG</u>	<u>N75923</u>	<u>002</u>	Mar 23, 2004
<u>AB</u>		<u>40MG</u>	<u>N75923</u>	<u>003</u>	Mar 23, 2004
<u>AB</u>		<u>80MG</u>	<u>N75923</u>	<u>004</u>	Sep 29, 2004
<u>AB</u>	IMPAX LABS	<u>80MG</u>	<u>N76318</u>	<u>001</u>	Sep 27, 2004
<u>AB</u>	IMPAX PHARMS	<u>10MG</u>	<u>N76446</u>	<u>001</u>	Dec 06, 2005
<u>AB</u>		<u>20MG</u>	<u>N76446</u>	<u>002</u>	Dec 06, 2005
<u>AB</u>		<u>40MG</u>	<u>N76446</u>	<u>003</u>	Dec 06, 2005
<u>AB</u>	TEVA	<u>10MG</u>	<u>N76610</u>	<u>001</u>	Dec 06, 2005
<u>AB</u>		<u>20MG</u>	<u>N76610</u>	<u>002</u>	Dec 06, 2005
<u>AB</u>		<u>40MG</u>	<u>N76610</u>	<u>003</u>	Dec 06, 2005
<u>AB</u>		<u>80MG</u>	<u>N76168</u>	<u>001</u>	Mar 23, 2004
<u>OXYCONTIN</u>					
<u>AB</u>	PURDUE PHARMA LP	<u>10MG</u>	<u>N20553</u>	<u>001</u>	Dec 12, 1995
<u>AB</u>		<u>20MG</u>	<u>N20553</u>	<u>002</u>	Dec 12, 1995
<u>AB</u>	+	<u>40MG</u>	<u>N20553</u>	<u>003</u>	Dec 12, 1995
<u>AB</u>		<u>80MG</u>	<u>N20553</u>	<u>004</u>	Jan 06, 1997
		15MG	N20553	006	Sep 18, 2006
		30MG	N20553	007	Sep 18, 2006
		60MG	N20553	008	Sep 18, 2006

OXYMETHOLONE

TABLET; ORAL					
ANADROL-50					
	+ ALAVEN PHARM	50MG	N16848	001	

OXYMORPHONE HYDROCHLORIDE

INJECTABLE; INJECTION					
NUMORPHAN					
	+ ENDO PHARMS	1MG/ML	N11707	002	
TABLET; ORAL					
OPANA					
	ENDO PHARMS	5MG	N21611	001	Jun 22, 2006
	+	10MG	N21611	002	Jun 22, 2006
TABLET, EXTENDED RELEASE; ORAL					
OPANA ER					
	ENDO PHARMS	5MG	N21610	001	Jun 22, 2006
		10MG	N21610	002	Jun 22, 2006
		20MG	N21610	003	Jun 22, 2006
	+	40MG	N21610	004	Jun 22, 2006

OXYTETRACYCLINE HYDROCHLORIDE

CAPSULE; ORAL					
TERRAMYCIN					
	+ PFIZER	EQ 250MG BASE	N50286	002	

OXYTETRACYCLINE HYDROCHLORIDE; POLYMYXIN B SULFATE

OINTMENT; OPHTHALMIC					
TERRAMYCIN W/ POLYMYXIN B SULFATE					
	+ PFIZER	EQ 5MG BASE/GM;10,000 UNITS/GM	N61015	001	

OXYTOCIN

INJECTABLE; INJECTION					
<u>OXYTOCIN</u>					
<u>AP</u>	+ ABRAXIS PHARM	<u>10USP UNITS/ML</u>	<u>N18248</u>	<u>001</u>	

PRESCRIPTION DRUG PRODUCT LIST

3 - 289 (of 370)

OXYTOCIN

INJECTABLE; INJECTION

PITOCIN

<u>AP</u>	+ PARKEDALE	<u>10USP UNITS/ML</u>	<u>N18261</u>	<u>001</u>	
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PACLITAXEL

FOR SUSPENSION; IV (INFUSION)

	+ ABRAXIS BIOSCIENCE	100MG/VIAL	N21660	001	Jan 07, 2005
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INJECTABLE; INJECTION

PACLITAXEL

<u>AP</u>	BEDFORD	<u>6MG/ML</u>	<u>N75190</u>	<u>001</u>	Jan 28, 2002
<u>AP</u>	DABUR ONCOLOGY PLC	<u>6MG/ML</u>	<u>N77574</u>	<u>001</u>	Nov 27, 2006
<u>AP</u>	MAYNE PHARMA USA	<u>6MG/ML</u>	<u>N76131</u>	<u>001</u>	May 08, 2002
<u>AP</u>		<u>6MG/ML</u>	<u>N76233</u>	<u>001</u>	Aug 01, 2002
<u>AP</u>	MYLAN	<u>6MG/ML</u>	<u>N75278</u>	<u>001</u>	Jan 25, 2002
<u>AP</u>	SICOR PHARMS	<u>6MG/ML</u>	<u>N75184</u>	<u>001</u>	Jan 25, 2002
<u>AP</u>		<u>6MG/ML</u>	<u>N75297</u>	<u>001</u>	Jan 25, 2002
<u>AP</u>	SUPERGEN	<u>6MG/ML</u>	<u>N75436</u>	<u>001</u>	Nov 12, 2004

TAXOL

<u>AP</u>	+ BRISTOL MYERS SQUIBB	<u>6MG/ML</u>	<u>N20262</u>	<u>001</u>	Dec 29, 1992
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PALIPERIDONE

TABLET, EXTENDED RELEASE; ORAL

INVEGA

	JANSSEN LP	3MG	N21999	001	Dec 19, 2006
		6MG	N21999	002	Dec 19, 2006
+		9MG	N21999	003	Dec 19, 2006

PALONOSETRON HYDROCHLORIDE

INJECTABLE; INTRAVENOUS

ALOXI

	+ HELSINN HLTHCARE	EQ 0.25MG BASE/5ML	N21372	001	Jul 25, 2003
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PAMIDRONATE DISODIUM

INJECTABLE; INJECTION

AREDIA

<u>AP</u>	+ NOVARTIS	<u>30MG/VIAL</u>	<u>N20036</u>	<u>001</u>	Oct 31, 1991
<u>AP</u>	+	<u>90MG/VIAL</u>	<u>N20036</u>	<u>004</u>	May 06, 1993

PAMIDRONATE DISODIUM

<u>AP</u>	ABRAXIS PHARM	<u>30MG/VIAL</u>	<u>N75773</u>	<u>001</u>	May 06, 2002
<u>AP</u>		<u>30MG /10ML (3MG/ML)</u>	<u>N76207</u>	<u>001</u>	May 17, 2002
<u>AP</u>		<u>90MG/VIAL</u>	<u>N75773</u>	<u>002</u>	May 06, 2002
<u>AP</u>		<u>90MG /10ML (9MG/ML)</u>	<u>N76207</u>	<u>002</u>	May 17, 2002
<u>AP</u>	AESGEN	<u>30MG/VIAL</u>	<u>N75594</u>	<u>001</u>	May 06, 2002
<u>AP</u>		<u>90MG/VIAL</u>	<u>N75594</u>	<u>002</u>	May 06, 2002
<u>AP</u>	+ BEDFORD	<u>30MG /10ML (3MG/ML)</u>	<u>N21113</u>	<u>001</u>	Mar 04, 2002
<u>AP</u>		<u>30MG/VIAL</u>	<u>N75290</u>	<u>001</u>	Apr 30, 2001
<u>AP</u>	+	<u>90MG /10ML (9MG/ML)</u>	<u>N21113</u>	<u>002</u>	Mar 04, 2002
<u>AP</u>		<u>90MG/VIAL</u>	<u>N75290</u>	<u>003</u>	Apr 30, 2001
<u>AP</u>	+ MAYNE PHARMA USA	<u>30MG/10ML (3MG/ML)</u>	<u>N75841</u>	<u>001</u>	Jun 27, 2002
<u>AP</u>	+	<u>60MG/10ML (6MG/ML)</u>	<u>N75841</u>	<u>002</u>	Jun 27, 2002
<u>AP</u>	+	<u>90MG/10ML (9MG/ML)</u>	<u>N75841</u>	<u>003</u>	Jun 27, 2002
<u>AP</u>	SICOR PHARMS	<u>30MG /10ML (3MG/ML)</u>	<u>N76153</u>	<u>001</u>	Mar 27, 2002
<u>AP</u>		<u>90MG /10ML (9MG/ML)</u>	<u>N76153</u>	<u>002</u>	Mar 27, 2002

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 290 (of 370)

PANCURONIUM BROMIDE

INJECTABLE; INJECTION

PANCURONIUM

<u>AP</u>	ELKINS SINN	<u>1MG/ML</u>	<u>N72058</u>	<u>001</u>	Mar 23, 1988
<u>AP</u>		<u>2MG/ML</u>	<u>N72059</u>	<u>001</u>	Mar 23, 1988
<u>AP</u>		<u>2MG/ML</u>	<u>N72060</u>	<u>001</u>	Mar 23, 1988
<u>PANCURONIUM BROMIDE</u>					
<u>AP</u>	HOSPIRA	<u>1MG/ML</u>	<u>N72320</u>	<u>001</u>	Jan 19, 1989
<u>AP</u>		<u>2MG/ML</u>	<u>N72321</u>	<u>001</u>	Jan 19, 1989
<u>AP</u>	+ SICOR PHARMS	<u>1MG/ML</u>	<u>N72759</u>	<u>001</u>	Jul 31, 1990
<u>AP</u>	+	<u>2MG/ML</u>	<u>N72760</u>	<u>001</u>	Jul 31, 1990

PANTOPRAZOLE SODIUM

INJECTABLE; IV (INFUSION)

	+ WYETH PHARMS INC	EQ 40MG BASE/VIAL	N20988	001	Mar 22, 2001
TABLET, DELAYED RELEASE; ORAL					
PROTONIX					
	WYETH PHARMS INC	EQ 20MG BASE	N20987	002	Jun 12, 2001
	+	EQ 40MG BASE	N20987	001	Feb 02, 2000

PARICALCITOL

CAPSULE; ORAL

ZEMPLAR

	ABBOTT	1UGM	N21606	001	May 26, 2005
		2UGM	N21606	002	May 26, 2005
	+	4UGM	N21606	003	May 26, 2005

INJECTABLE; INJECTION

ZEMPLAR

	+ ABBOTT	0.002MG/ML	N20819	002	Feb 01, 2000
	+	0.005MG/ML	N20819	001	Apr 17, 1998

PAROMOMYCIN SULFATE

CAPSULE; ORAL

HUMATIN

<u>AA</u>	+ KING PHARMS	<u>EQ 250MG BASE</u>	<u>N62310</u>	<u>001</u>	
<u>PAROMOMYCIN SULFATE</u>					
<u>AA</u>	CARACO	<u>EQ 250MG BASE</u>	<u>N64171</u>	<u>001</u>	Jun 30, 1997

PAROXETINE HYDROCHLORIDE

SUSPENSION; ORAL

PAROXETINE HYDROCHLORIDE

<u>AB</u>	APOTEX INC	<u>EQ 10MG BASE/5ML</u>	<u>N77395</u>	<u>001</u>	Dec 05, 2006
<u>PAXIL</u>					
<u>AB</u>	+ GLAXOSMITHKLINE	<u>EQ 10MG BASE/5ML</u>	<u>N20710</u>	<u>001</u>	Jun 25, 1997

TABLET; ORAL

PAROXETINE HYDROCHLORIDE

<u>AB</u>	ALPHAPHARM	<u>EQ 10MG BASE</u>	<u>N75716</u>	<u>001</u>	Mar 08, 2004
<u>AB</u>		<u>EQ 20MG BASE</u>	<u>N75716</u>	<u>002</u>	Mar 08, 2004
<u>AB</u>		<u>EQ 30MG BASE</u>	<u>N75716</u>	<u>003</u>	Mar 08, 2004
<u>AB</u>		<u>EQ 40MG BASE</u>	<u>N75716</u>	<u>004</u>	Mar 08, 2004
<u>AB</u>	SANDOZ	<u>EQ 10MG BASE</u>	<u>N75566</u>	<u>001</u>	Mar 08, 2004
<u>AB</u>		<u>EQ 20MG BASE</u>	<u>N75566</u>	<u>002</u>	Mar 08, 2004
<u>AB</u>		<u>EQ 30MG BASE</u>	<u>N75566</u>	<u>003</u>	Mar 08, 2004
<u>AB</u>		<u>EQ 40MG BASE</u>	<u>N75566</u>	<u>004</u>	Mar 08, 2004
<u>AB</u>	TEVA	<u>EQ 10MG BASE</u>	<u>N76618</u>	<u>001</u>	Aug 15, 2005
<u>AB</u>		<u>EQ 20MG BASE</u>	<u>N76618</u>	<u>002</u>	Aug 15, 2005
<u>AB</u>		<u>EQ 30MG BASE</u>	<u>N76618</u>	<u>003</u>	Aug 15, 2005
<u>AB</u>		<u>EQ 40MG BASE</u>	<u>N76618</u>	<u>004</u>	Aug 15, 2005

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 291 (of 370)

PAROXETINE HYDROCHLORIDE

TABLET; ORAL

PAROXETINE HYDROCHLORIDE

<u>AB</u>	TORPHARM	<u>EQ 10MG BASE</u>	<u>N75356</u>	<u>001</u>	Jul 30, 2003
<u>AB</u>		<u>EQ 20MG BASE</u>	<u>N75356</u>	<u>002</u>	Jul 30, 2003
<u>AB</u>		<u>EQ 30MG BASE</u>	<u>N75356</u>	<u>003</u>	Jul 30, 2003
<u>AB</u>		<u>EQ 40MG BASE</u>	<u>N75356</u>	<u>004</u>	Jul 30, 2003

PAXIL

<u>AB</u>	GLAXOSMITHKLINE	<u>EQ 10MG BASE</u>	<u>N20031</u>	<u>001</u>	Dec 29, 1992
<u>AB</u>		<u>EQ 20MG BASE</u>	<u>N20031</u>	<u>002</u>	Dec 29, 1992
<u>AB</u>		<u>EQ 30MG BASE</u>	<u>N20031</u>	<u>003</u>	Dec 29, 1992
<u>AB</u>	+	<u>EQ 40MG BASE</u>	<u>N20031</u>	<u>005</u>	Dec 29, 1992

TABLET, EXTENDED RELEASE; ORAL

PAXIL CR

	GLAXOSMITHKLINE	EQ 12.5MG BASE	N20936	001	Feb 16, 1999
		EQ 25MG BASE	N20936	002	Feb 16, 1999
+		EQ 37.5MG BASE	N20936	003	Dec 06, 2000

PAROXETINE MESYLATE

TABLET; ORAL

PEXEVA

	JDS PHARMS	EQ 10MG BASE	N21299	001	Jul 03, 2003
		EQ 20MG BASE	N21299	002	Jul 03, 2003
		EQ 30MG BASE	N21299	003	Jul 03, 2003
+		EQ 40MG BASE	N21299	004	Jul 03, 2003

PEGADEMASE BOVINE

INJECTABLE; INJECTION

ADAGEN

+	ENZON	250 UNITS/ML	N19818	001	Mar 21, 1990
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PEGAPTANIB SODIUM

INJECTABLE; INTRAVITREAL

MACUGEN

+	OSI EYETECH	EQ 0.3MG ACID/0.09ML	N21756	001	Dec 17, 2004
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PEGVISOMANT

INJECTABLE; SUBCUTANEOUS

SOMAVERT

+	PHARMACIA AND UPJOHN	10MG/VIAL	N21106	001	Mar 25, 2003
+		15MG/VIAL	N21106	002	Mar 25, 2003
+		20MG/VIAL	N21106	003	Mar 25, 2003

PEMETREXED DISODIUM

INJECTABLE; IV (INFUSION)

ALIMTA

+	LILLY	EQ 500MG BASE/VIAL	N21462	001	Feb 04, 2004
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PEMIROLAST POTASSIUM

SOLUTION/DROPS; OPHTHALMIC

+	SANTEN	0.1%	N21079	001	Sep 24, 1999
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PENBUTOLOL SULFATE

TABLET; ORAL

LEVATOL

+	SCHWARZ PHARMA	20MG	N18976	004	Jan 05, 1989
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PRESCRIPTION DRUG PRODUCT LIST

3 - 292 (of 370)

PENCICLOVIR SODIUM

CREAM; TOPICAL
DENA VIR

+ NOVARTIS	1%	N20629 001	Sep 24, 1996
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PENICILLAMINE

CAPSULE; ORAL
CUPRIMINE

+ ATON	250MG	N19853 001	
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TABLET; ORAL
DEPEN

+ MEDPOINTE PHARM HLC	250MG	N19854 001	
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PENICILLIN G BENZATHINE

INJECTABLE; INJECTION
BICILLIN L-A

BC + KING PHARMS	600,000 UNITS/ML	N50141 001	
+	300,000 UNITS/ML	N50141 003	

PERMAPEN

BC PFIZER	600,000 UNITS/ML	N60014 001	
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PENICILLIN G BENZATHINE; PENICILLIN G PROCAINE

INJECTABLE; INJECTION
BICILLIN C-R

+ KING PHARMS	150,000 UNITS/ML;150,000 UNITS/ML	N50138 002	
+	300,000 UNITS/ML;300,000 UNITS/ML	N50138 001	

BICILLIN C-R 900/300

+ KING PHARMS	900,000 UNITS/2ML;300,000 UNITS/2ML	N50138 003	
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PENICILLIN G POTASSIUM

INJECTABLE; INJECTION

PENICILLIN G POTASSIUM

<u>AP</u> PFIZER	<u>20,000,000 UNITS/VIAL</u>	<u>N60074 003</u>	
<u>AP</u> SANDOZ	<u>1,000,000 UNITS/VIAL</u>	<u>N65079 001</u>	Aug 30, 2002
<u>AP</u>	<u>5,000,000 UNITS/VIAL</u>	<u>N65079 002</u>	Aug 30, 2002
<u>AP</u>	<u>20,000,000 UNITS/VIAL</u>	<u>N65079 003</u>	Aug 30, 2002

PENICILLIN G POTASSIUM IN PLASTIC CONTAINER

+ BAXTER HLTHCARE	20,000 UNITS/ML	N50638 001	Jun 25, 1990
+	40,000 UNITS/ML	N50638 002	Jun 25, 1990
+	60,000 UNITS/ML	N50638 003	Jun 25, 1990

PFIZERPEN

<u>AP</u> + PFIZER	<u>1,000,000 UNITS/VIAL</u>	<u>N60657 001</u>	
<u>AP</u> +	<u>5,000,000 UNITS/VIAL</u>	<u>N60657 002</u>	
<u>AP</u> +	<u>20,000,000 UNITS/VIAL</u>	<u>N60657 003</u>	

PENICILLIN G PROCAINE

INJECTABLE; INJECTION
PENICILLIN G PROCAINE

+ KING PHARMS	300,000 UNITS/ML	N60101 002	
+	600,000 UNITS/ML	N60101 001	

PENICILLIN G SODIUM

INJECTABLE; IM-IV
PENICILLIN G SODIUM

+ BIOCHEMIE	5,000,000 UNITS/VIAL	N65068 001	Feb 26, 2001
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27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 293 (of 370)

PENICILLIN V POTASSIUM

FOR SOLUTION; ORAL

PENICILLIN V POTASSIUM

<u>AA</u>	AM ANTIBIOTICS	<u>EQ 125MG BASE/5ML</u>	<u>N61529</u>	<u>001</u>	
<u>AA</u>		<u>EQ 250MG BASE/5ML</u>	<u>N61529</u>	<u>002</u>	
<u>AA</u>	CLONMEL	<u>EQ 125MG BASE/5ML</u>	<u>N62981</u>	<u>001</u>	Feb 10, 1989
<u>AA</u>		<u>EQ 250MG BASE/5ML</u>	<u>N62981</u>	<u>002</u>	Feb 10, 1989
	<u>PENICILLIN-VK</u>				
<u>AA</u>	TEVA	<u>EQ 125MG BASE/5ML</u>	<u>N60456</u>	<u>001</u>	
<u>AA</u>		<u>EQ 250MG BASE/5ML</u>	<u>N60456</u>	<u>002</u>	
	<u>VEETIDS</u>				
<u>AA</u>	APOTHECON	<u>EQ 125MG BASE/5ML</u>	<u>N61410</u>	<u>001</u>	
<u>AA</u>		<u>EQ 250MG BASE/5ML</u>	<u>N61410</u>	<u>002</u>	

TABLET; ORAL

PENICILLIN V POTASSIUM

<u>AB</u>	BIOCHEMIE	<u>EQ 250MG BASE</u>	<u>N64071</u>	<u>001</u>	Nov 30, 1995
<u>AB</u>	+	<u>EQ 500MG BASE</u>	<u>N64071</u>	<u>002</u>	Nov 30, 1995
<u>AB</u>	CLONMEL	<u>EQ 250MG BASE</u>	<u>N62936</u>	<u>001</u>	Nov 25, 1988
<u>AB</u>		<u>EQ 500MG BASE</u>	<u>N62935</u>	<u>001</u>	Nov 23, 1988
	<u>PENICILLIN-VK</u>				
<u>AB</u>	TEVA	<u>EQ 250MG BASE</u>	<u>N60711</u>	<u>002</u>	
<u>AB</u>		<u>EQ 500MG BASE</u>	<u>N60711</u>	<u>003</u>	

PENTAMIDINE ISETHIONATE

FOR SOLUTION; INHALATION

NEBUPENT

+	ABRAXIS PHARM	300MG/VIAL	N19887	001	Jun 15, 1989
		600MG/VIAL	N19887	002	Mar 22, 1996

INJECTABLE; INJECTION

PENTAM

<u>AP</u>	+	ABRAXIS PHARM	<u>300MG/VIAL</u>	<u>N19264</u>	<u>001</u>	Oct 16, 1984
		<u>PENTAMIDINE ISETHIONATE</u>				
<u>AP</u>		HOSPIRA	<u>300MG/VIAL</u>	<u>N73479</u>	<u>001</u>	Jun 30, 1992
<u>AP</u>		WATSON LABS	<u>300MG/VIAL</u>	<u>N74303</u>	<u>001</u>	Aug 17, 1995

PENTAZOCINE LACTATE

INJECTABLE; INJECTION

TALWIN

+	HOSPIRA	EQ 30MG BASE/ML	N16194	001	
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PENTETATE CALCIUM TRISODIUM

SOLUTION; INHALATION, INTRAVENOUS

PENTETATE CALCIUM TRISODIUM

+	HAMELN PHARMS	EQ 1GM/5ML(EQ 200MG/ML)	N21749	001	Aug 11, 2004
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PENTETATE ZINC TRISODIUM

SOLUTION; INHALATION, INTRAVENOUS

PENTETATE ZINC TRISODIUM

+	HAMELN PHARMS	EQ 1GM/5ML(EQ 200MG/ML)	N21751	001	Aug 11, 2004
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PENTOBARBITAL SODIUM

INJECTABLE; INJECTION

NEMBUTAL SODIUM

+	OVATION PHARMS	50MG/ML	N83246	001	
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PENTOSAN POLYSULFATE SODIUM

CAPSULE; ORAL

ELMIRON

+	ORTHO MCNEIL PHARM	100MG	N20193	001	Sep 26, 1996
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PRESCRIPTION DRUG PRODUCT LIST

3 - 294 (of 370)

PENTOSTATIN

INJECTABLE; INJECTION
NIPENT

+ MAYNE PHARMA USA	10MG/VIAL	N20122	001	Oct 11, 1991
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PENTOXIFYLLINE

TABLET, EXTENDED RELEASE; ORAL

PENTOXIFYLLINE

<u>AB</u>	ACTAVIS ELIZABETH	<u>400MG</u>	<u>N74878</u>	<u>001</u>	Jul 09, 1997
<u>AB</u>	BIOVAIL	<u>400MG</u>	<u>N75028</u>	<u>001</u>	Jul 20, 1998
<u>AB</u>	IMPAX LABS	<u>400MG</u>	<u>N75093</u>	<u>001</u>	Aug 10, 1999
<u>AB</u>	MYLAN	<u>400MG</u>	<u>N74425</u>	<u>001</u>	Jul 08, 1997
<u>AB</u>	PLIVA	<u>400MG</u>	<u>N74874</u>	<u>001</u>	May 25, 1999
<u>AB</u>	RADIUS PHARMS	<u>400MG</u>	<u>N74877</u>	<u>001</u>	Jul 08, 1997
<u>AB</u>	TEVA	<u>400MG</u>	<u>N75199</u>	<u>001</u>	Sep 03, 1999
<u>AB</u>	TORPHARM	<u>400MG</u>	<u>N75191</u>	<u>001</u>	Jun 09, 1999
<u>AB</u>	WATSON LABS	<u>400MG</u>	<u>N75107</u>	<u>001</u>	Sep 04, 1998

PENTOXIL

<u>AB</u>	UPSHER SMITH	<u>400MG</u>	<u>N74962</u>	<u>001</u>	Mar 31, 1999
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TRENTAL

<u>AB</u>	+ SANOFI AVENTIS US	<u>400MG</u>	<u>N18631</u>	<u>001</u>	Aug 30, 1984
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PERFLUTREN

INJECTABLE; INTRAVENOUS
DEFINITY

+ BRISTOL MYERS SQUIBB	6.52MG/ML	N21064	001	Jul 31, 2001
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PERGOLIDE MESYLATE

TABLET; ORAL

PERGOLIDE MESYLATE

<u>AB</u>	IVAX PHARMS	<u>EQ 0.05MG BASE</u>	<u>N76094</u>	<u>001</u>	Sep 04, 2003
<u>AB</u>		<u>EQ 0.25MG BASE</u>	<u>N76094</u>	<u>002</u>	Sep 04, 2003
<u>AB</u>		<u>EQ 1MG BASE</u>	<u>N76094</u>	<u>003</u>	Sep 04, 2003
<u>AB</u>	PAR PHARM	<u>EQ 0.05MG BASE</u>	<u>N76061</u>	<u>001</u>	Nov 27, 2002
<u>AB</u>		<u>EQ 0.25MG BASE</u>	<u>N76061</u>	<u>002</u>	Nov 27, 2002
<u>AB</u>		<u>EQ 1MG BASE</u>	<u>N76061</u>	<u>003</u>	Nov 27, 2002

PERMAX

<u>AB</u>	VALEANT	<u>EQ 0.05MG BASE</u>	<u>N19385</u>	<u>001</u>	Dec 30, 1988
<u>AB</u>	+	<u>EQ 0.25MG BASE</u>	<u>N19385</u>	<u>002</u>	Dec 30, 1988
<u>AB</u>		<u>EQ 1MG BASE</u>	<u>N19385</u>	<u>003</u>	Dec 30, 1988

PERINDOPRIL ERBUMINE

TABLET; ORAL

ACEON

SOLVAY PHARMS	2MG	N20184	001	Dec 30, 1993
	4MG	N20184	002	Dec 30, 1993
+	8MG	N20184	003	Dec 30, 1993

PERMETHRIN

CREAM; TOPICAL

ELIMITE

<u>AB</u>	+ ALLERGAN	<u>5%</u>	<u>N19855</u>	<u>001</u>	Aug 25, 1989
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PERMETHRIN

<u>AB</u>	ACTAVIS MID ATLANTIC	<u>5%</u>	<u>N74806</u>	<u>001</u>	Jan 23, 1998
<u>AB</u>	PERRIGO NEW YORK	<u>5%</u>	<u>N76369</u>	<u>001</u>	Apr 21, 2003

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 295 (of 370)

PERPHENAZINE

CONCENTRATE; ORAL

PERPHENAZINE

+ PHARM ASSOC

16MG/5ML

N40360 001

May 25, 2001

TABLET; ORAL

PERPHENAZINEAB IVAX PHARMS2MGN89707 001

Sep 10, 1987

AB4MGN89708 001

Sep 10, 1987

AB8MGN89456 001

Sep 10, 1987

AB +16MGN89457 001

Sep 10, 1987

AB SANDOZ2MGN89683 001

Dec 08, 1988

AB4MGN89684 001

Dec 08, 1988

AB8MGN89685 001

Dec 08, 1988

AB16MGN89686 001

Dec 08, 1988

AB VINTAGE PHARMS2MGN40226 001

Dec 31, 1998

AB4MGN40226 002

Dec 31, 1998

AB8MGN40226 003

Dec 31, 1998

AB16MGN40226 004

Dec 31, 1998

PHENDIMETRAZINE TARTRATE

CAPSULE; ORAL

PHENDIMETRAZINE TARTRATEAA + SANDOZ35MGN85695 001X-TROZINEAA SHIRE RICHWOOD35MGN87394 001

Sep 22, 1982

CAPSULE, EXTENDED RELEASE; ORAL

BONTRIL

BC MALLINCKRODT

105MG

N88021 001

Sep 21, 1982

PHENDIMETRAZINE TARTRATE

BC + SANDOZ

105MG

N18074 001

X-TROZINE L.A.

BC SHIRE RICHWOOD

105MG

N87371 001

Aug 24, 1982

TABLET; ORAL

BONTRIL PDMAA + VALEANT35MGN85272 001PHENDIMETRAZINE TARTRATEAA MIKART35MGN89452 001

Oct 30, 1991

AA SANDOZ35MGN85588 001PHENELZINE SULFATE

TABLET; ORAL

NARDIL

+ PARKE DAVIS

EQ 15MG BASE

N11909 002

PHENOXYBENZAMINE HYDROCHLORIDE

CAPSULE; ORAL

DIBENZYLINE

+ WELLSRING PHARM

10MG

N08708 001

PHENTERMINE HYDROCHLORIDE

CAPSULE; ORAL

ADIPEX-PAA + TEVA37.5MGN88023 001

Aug 02, 1983

PHENTERMINE HYDROCHLORIDEAA ACTAVIS TOTOWA15MGN40460 001

Jan 14, 2003

AA30MGN40227 001

Jun 18, 1997

AA30MGN40448 001

Jan 22, 2003

AA37.5MGN40228 001

Jun 19, 1997

AA AMBI PHARMS30MGN40083 001

Mar 07, 1997

PRESCRIPTION DRUG PRODUCT LIST

3 - 296 (of 370)

PHENTERMINE HYDROCHLORIDE

CAPSULE; ORAL

PHENTERMINE HYDROCHLORIDE

<u>AA</u>	MUTUAL PHARM	<u>30MG</u>	<u>N40525</u>	<u>001</u>	Oct 23, 2003
<u>AA</u>		<u>37.5MG</u>	<u>N40527</u>	<u>001</u>	Oct 23, 2003
<u>AA</u>	+ SANDOZ	<u>15MG</u>	<u>N87190</u>	<u>002</u>	
<u>AA</u>		<u>30MG</u>	<u>N86945</u>	<u>001</u>	Jul 20, 1983
<u>AA</u>		<u>30MG</u>	<u>N87190</u>	<u>001</u>	
<u>AA</u>	VINTAGE PHARMS	<u>37.5MG</u>	<u>N40377</u>	<u>001</u>	Jan 04, 2002

TABLET; ORAL

ADIPEX-P

<u>AA</u>	+ TEVA	<u>37.5MG</u>	<u>N85128</u>	<u>001</u>	
<u>PHENTERMINE HYDROCHLORIDE</u>					
<u>AA</u>	ACTAVIS ELIZABETH	<u>37.5MG</u>	<u>N40276</u>	<u>001</u>	Nov 25, 1998
<u>AA</u>	ACTAVIS TOTOWA	<u>37.5MG</u>	<u>N40190</u>	<u>001</u>	May 30, 1997
<u>AA</u>	LANNETT	<u>37.5MG</u>	<u>N40555</u>	<u>001</u>	Apr 15, 2005
<u>AA</u>	MUTUAL PHARM	<u>37.5MG</u>	<u>N40526</u>	<u>001</u>	Oct 23, 2003
	+ SANDOZ	30MG	N88605	001	Sep 28, 1987

PHENTERMINE RESIN COMPLEX

CAPSULE, EXTENDED RELEASE; ORAL

IONAMIN

UCB INC

EQ 15MG BASE

N11613 004

+

EQ 30MG BASE

N11613 002

PHENTOLAMINE MESYLATE

INJECTABLE; INJECTION

PHENTOLAMINE MESYLATE

<u>AP</u>	BEDFORD	<u>5MG/VIAL</u>	<u>N40235</u>	<u>001</u>	Mar 11, 1998
<u>REGITINE</u>					
<u>AP</u>	+ NOVARTIS	<u>5MG/VIAL</u>	<u>N08278</u>	<u>003</u>	

PHENYLEPHRINE HYDROCHLORIDE; PROMETHAZINE HYDROCHLORIDE

SYRUP; ORAL

PHENYLEPHRINE HYDROCHLORIDE AND PROMETHAZINE HYDROCHLORIDE

<u>AA</u>	VINTAGE	<u>5MG/5ML; 6.25MG/5ML</u>	<u>N40654</u>	<u>001</u>	Dec 07, 2006
<u>PROMETH VC PLAIN</u>					
<u>AA</u>	+ ACTAVIS MID ATLANTIC	<u>5MG/5ML; 6.25MG/5ML</u>	<u>N88761</u>	<u>001</u>	Nov 08, 1984

PHENYTOIN

SUSPENSION; ORAL

DILANTIN-125

<u>AB</u>	+ PARKE DAVIS	<u>125MG/5ML</u>	<u>N08762</u>	<u>001</u>	
<u>PHENTYTOIN</u>					
<u>AB</u>	MORTON GROVE	<u>125MG/5ML</u>	<u>N40420</u>	<u>001</u>	Apr 19, 2002
<u>PHENYTOIN</u>					
<u>AB</u>	ACTAVIS MID ATLANTIC	<u>125MG/5ML</u>	<u>N89892</u>	<u>001</u>	Sep 25, 1992
<u>AB</u>	TARO	<u>125MG/5ML</u>	<u>N40521</u>	<u>001</u>	Mar 08, 2004
<u>AB</u>	VISTAPHARM	<u>125MG/5ML</u>	<u>N40342</u>	<u>001</u>	Jan 31, 2001
<u>AB</u>		<u>125MG/5ML</u>	<u>N40610</u>	<u>001</u>	Aug 18, 2005

TABLET, CHEWABLE; ORAL

DILANTIN

+ PFIZER PHARMS

50MG

N84427 001

PHENYTOIN SODIUM

CAPSULE; ORAL

DILANTIN

<u>AB</u>	+ PARKE DAVIS	<u>100MG EXTENDED</u>	<u>N84349</u>	<u>002</u>	
	+	30MG EXTENDED	N84349	001	

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 297 (of 370)

PHENYTOIN SODIUM

CAPSULE; ORAL

EXTENDED PHENYTOIN SODIUM

<u>AB</u>	BARR	<u>100MG EXTENDED</u>	<u>N40435</u>	<u>001</u>	Jun 20, 2003
<u>AB</u>	MYLAN	<u>100MG EXTENDED</u>	<u>N40298</u>	<u>001</u>	Dec 28, 1998
<u>AB</u>	SUN PHARM INDS (IN)	<u>100MG EXTENDED</u>	<u>N40621</u>	<u>001</u>	Dec 11, 2006
<u>AB</u>	TARO	<u>100MG EXTENDED</u>	<u>N40684</u>	<u>001</u>	Sep 05, 2006
	PHENYTEK				
	+ MYLAN	200MG EXTENDED	N40298	002	Dec 06, 2001
	+	300MG EXTENDED	N40298	003	Dec 06, 2001
	PROMPT PHENYTOIN SODIUM				
BX	+ IVAX PHARMS	100MG PROMPT	N80259	001	

INJECTABLE; INJECTION

PHENYTOIN

<u>AP</u>	+ ELKINS SINN	<u>50MG/ML</u>	<u>N84307</u>	<u>001</u>	
	<u>PHENYTOIN SODIUM</u>				
<u>AP</u>	HIKMA FARMACEUTICA	<u>50MG/ML</u>	<u>N40573</u>	<u>001</u>	Sep 13, 2006
<u>AP</u>	HOSPIRA	<u>50MG/ML</u>	<u>N89521</u>	<u>001</u>	Mar 17, 1987
<u>AP</u>		<u>50MG/ML</u>	<u>N89744</u>	<u>001</u>	Dec 18, 1987

PHYTONADIONE

INJECTABLE; INJECTION

PHYTONADIONE

BP	INTL MEDICATION	1MG/0.5ML	N83722	001	
	VITAMIN K1				
BP	+ HOSPIRA	1MG/0.5ML	N87954	001	Jul 25, 1983
	+	10MG/ML	N87955	001	Jul 25, 1983

TABLET; ORAL

MEPHYTON

	+ ATON	5MG	N10104	003	
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PILOCARPINE HYDROCHLORIDE

GEL; OPHTHALMIC

PILOPINE HS

	+ ALCON	4%	N18796	001	Oct 01, 1984
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TABLET; ORAL

PILOCARPINE HYDROCHLORIDE

<u>AB</u>	COREPHARMA	<u>5MG</u>	<u>N76746</u>	<u>001</u>	Nov 16, 2004
<u>AB</u>	IMPAX LABS	<u>5MG</u>	<u>N77248</u>	<u>001</u>	Mar 31, 2006
<u>AB</u>		<u>7.5MG</u>	<u>N77248</u>	<u>002</u>	Mar 31, 2006
<u>AB</u>	LANNETT	<u>5MG</u>	<u>N77220</u>	<u>001</u>	Oct 14, 2005
<u>AB</u>	ROXANE	<u>5MG</u>	<u>N76963</u>	<u>001</u>	Dec 22, 2004
	<u>SALAGEN</u>				
<u>AB</u>	MGI PHARMA INC	<u>5MG</u>	<u>N20237</u>	<u>001</u>	Mar 22, 1994
<u>AB</u>	+	<u>7.5MG</u>	<u>N20237</u>	<u>002</u>	Apr 18, 2003

PIMECROLIMUS

CREAM; TOPICAL

ELIDEL

	+ NOVARTIS	1%	N21302	001	Dec 13, 2001
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PIMOZIDE

TABLET; ORAL

ORAP

	TEVA	1MG	N17473	003	Aug 27, 1997
	+	2MG	N17473	001	Jul 31, 1984

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 298 (of 370)

PINDOLOL

TABLET; ORAL

PINDOLOL

<u>AB</u>	GENPHARM	<u>5MG</u>	<u>N74013</u>	<u>001</u>	Sep 24, 1992
<u>AB</u>		<u>10MG</u>	<u>N74018</u>	<u>001</u>	Sep 24, 1992
<u>AB</u>	IVAX PHARMS	<u>5MG</u>	<u>N73687</u>	<u>001</u>	Feb 26, 1993
<u>AB</u>		<u>10MG</u>	<u>N73687</u>	<u>002</u>	Feb 26, 1993
<u>AB</u>	MARTEC USA LLC	<u>5MG</u>	<u>N74474</u>	<u>001</u>	Oct 28, 1996
<u>AB</u>		<u>10MG</u>	<u>N74474</u>	<u>002</u>	Oct 28, 1996
<u>AB</u>	MUTUAL PHARM	<u>5MG</u>	<u>N74063</u>	<u>001</u>	Jan 27, 1994
<u>AB</u>		<u>10MG</u>	<u>N74063</u>	<u>002</u>	Jan 27, 1994
<u>AB</u>	MYLAN	<u>5MG</u>	<u>N74019</u>	<u>001</u>	Sep 03, 1992
<u>AB</u>		<u>10MG</u>	<u>N74019</u>	<u>002</u>	Sep 03, 1992
<u>AB</u>	SANDOZ	<u>5MG</u>	<u>N73608</u>	<u>001</u>	Mar 29, 1993
<u>AB</u>		<u>10MG</u>	<u>N73609</u>	<u>001</u>	Mar 29, 1993
<u>AB</u>	TEVA	<u>5MG</u>	<u>N73661</u>	<u>001</u>	Oct 31, 1993
<u>AB</u>		<u>5MG</u>	<u>N74123</u>	<u>001</u>	Apr 17, 1997
<u>AB</u>		<u>10MG</u>	<u>N73661</u>	<u>002</u>	Oct 31, 1993
<u>AB</u>		<u>10MG</u>	<u>N74123</u>	<u>002</u>	Apr 17, 1997
<u>AB</u>	WATSON LABS	<u>5MG</u>	<u>N74437</u>	<u>001</u>	Feb 27, 1995
<u>AB</u>		<u>10MG</u>	<u>N74437</u>	<u>002</u>	Feb 27, 1995
<u>VISKEN</u>					
<u>AB</u>	NOVARTIS	<u>5MG</u>	<u>N18285</u>	<u>001</u>	Sep 03, 1982
<u>AB</u>	+	<u>10MG</u>	<u>N18285</u>	<u>002</u>	Sep 03, 1982

PIOGLITAZONE HYDROCHLORIDE

TABLET; ORAL

ACTOS

TAKEDA GLOBAL	EQ 15MG BASE	N21073	001	Jul 15, 1999
	EQ 30MG BASE	N21073	002	Jul 15, 1999
+	EQ 45MG BASE	N21073	003	Jul 15, 1999

PIPERACILLIN SODIUM

INJECTABLE; INJECTION

PIPERACILLIN

+	ABRAXIS PHARM	EQ 2GM BASE/VIAL	N65114	001	Nov 14, 2003
+		EQ 3GM BASE/VIAL	N65114	002	Nov 14, 2003
+		EQ 4GM BASE/VIAL	N65114	003	Nov 14, 2003
+		EQ 40GM BASE/VIAL	N65157	001	Jul 12, 2004

PIPERACILLIN SODIUM; TAZOBACTAM SODIUM

INJECTABLE; INJECTION

ZOSYN

+	WYETH PHARMS INC	EQ 2GM BASE/VIAL;EQ 250MG BASE/VIAL	N50684	001	Oct 22, 1993
+		EQ 3GM BASE/VIAL;EQ 375MG BASE/VIAL	N50684	002	Oct 22, 1993
+		EQ 4GM BASE/VIAL;EQ 500MG BASE/VIAL	N50684	003	Oct 22, 1993
+		EQ 36GM BASE/VIAL;EQ 4.5GM BASE/VIAL	N50684	004	Oct 22, 1993
<u>ZOSYN IN PLASTIC CONTAINER</u>					
+	WYETH PHARMS INC	EQ 40MG BASE/ML;EQ 5MG BASE/ML	N50750	001	Feb 24, 1998
+		EQ 60MG BASE/ML;EQ 7.5MG BASE/ML	N50750	002	Feb 24, 1998
+		EQ 4GM BASE/100ML;EQ 500MG BASE/100ML	N50750	003	Feb 24, 1998

PIRBUTEROL ACETATE

AEROSOL, METERED; INHALATION

MAXAIR

+	3M	EQ 0.2MG BASE/INH	N20014	001	Nov 30, 1992
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PRESCRIPTION DRUG PRODUCT LIST

3 - 299 (of 370)

PIROXICAM

CAPSULE; ORAL

FELDENE

<u>AB</u>	PFIZER	<u>10MG</u>	<u>N18147</u>	<u>002</u>	Apr 06, 1982
<u>AB</u>	+	<u>20MG</u>	<u>N18147</u>	<u>003</u>	Apr 06, 1982
	<u>PIROXICAM</u>				
<u>AB</u>	EGIS	<u>10MG</u>	<u>N74808</u>	<u>001</u>	Jul 08, 1997
<u>AB</u>		<u>20MG</u>	<u>N74808</u>	<u>002</u>	Jul 08, 1997
<u>AB</u>	GENPHARM	<u>10MG</u>	<u>N74043</u>	<u>001</u>	Sep 22, 1992
<u>AB</u>		<u>20MG</u>	<u>N74043</u>	<u>002</u>	Sep 22, 1992
<u>AB</u>	MEPHA	<u>10MG</u>	<u>N74116</u>	<u>001</u>	Jun 15, 1993
<u>AB</u>		<u>20MG</u>	<u>N74118</u>	<u>001</u>	Jun 15, 1993
<u>AB</u>	MUTUAL PHARM	<u>10MG</u>	<u>N73535</u>	<u>001</u>	Mar 12, 1993
<u>AB</u>		<u>20MG</u>	<u>N73536</u>	<u>001</u>	Mar 12, 1993
<u>AB</u>	MYLAN	<u>10MG</u>	<u>N74102</u>	<u>001</u>	Jul 31, 1992
<u>AB</u>		<u>20MG</u>	<u>N74102</u>	<u>002</u>	Jul 31, 1992
<u>AB</u>	TEVA	<u>10MG</u>	<u>N73637</u>	<u>001</u>	Jan 28, 1994
<u>AB</u>		<u>10MG</u>	<u>N74131</u>	<u>001</u>	Dec 11, 1992
<u>AB</u>		<u>20MG</u>	<u>N73638</u>	<u>001</u>	Jan 28, 1994
<u>AB</u>		<u>20MG</u>	<u>N74131</u>	<u>002</u>	Dec 11, 1992
<u>AB</u>	TEVA PHARMS	<u>10MG</u>	<u>N74103</u>	<u>001</u>	Aug 28, 1992
<u>AB</u>		<u>20MG</u>	<u>N74103</u>	<u>002</u>	Aug 28, 1992
<u>AB</u>	WATSON LABS	<u>10MG</u>	<u>N74287</u>	<u>001</u>	May 16, 1996
<u>AB</u>		<u>10MG</u>	<u>N74460</u>	<u>001</u>	Sep 29, 1995
<u>AB</u>		<u>20MG</u>	<u>N74287</u>	<u>002</u>	May 16, 1996
<u>AB</u>		<u>20MG</u>	<u>N74460</u>	<u>002</u>	Sep 29, 1995

PLICAMYCIN

INJECTABLE; INJECTION

MITHRACIN

+ PFIZER 2.5MG/VIAL N50109 001

PODOFILOX

GEL; TOPICAL

CONDYLOX

+ OCLASSEN 0.5% N20529 001 Mar 13, 1997

SOLUTION; TOPICAL

CONDYLOX

AT + OCLASSEN 0.5% N19795 001 Dec 13, 1990

PODOFILOX

AT PADDOCK 0.5% N75600 001 Jan 29, 2002

POLYETHYLENE GLYCOL 3350

FOR SOLUTION; ORAL

GLYCOLAX

AA SCHWARZ PHARMA 17GM/SC00PFUL N76652 001 Jul 02, 2004

POLYETHYLENE GLYCOL 3350

AA COASTAL PHARMS 17GM/SC00PFUL N77893 001 May 26, 2006

AA KALI LABS 17GM/SC00PFUL N77736 001 May 26, 2006

AA TEVA PHARMS 17GM/SC00PFUL N77445 001 May 04, 2006

AA YVR THERAP 17GM/SC00PFUL N77706 001 Sep 27, 2006

POLYETHYLENE GLYCOL 3350; POTASSIUM CHLORIDE; SODIUM BICARBONATE; SODIUM CHLORIDE

FOR SOLUTION; ORAL

NULYTELY

AA + BRAINTREE 420GM/BOT;1.48GM/BOT;5.72GM/BOT;11.2GM/BOT N19797 001 Apr 22, 1991

NULYTELY-FLAVORED

AA + BRAINTREE 420GM/BOT;1.48GM/BOT;5.72GM/BOT;11.2GM/BOT N19797 002 Nov 18, 1994

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 300 (of 370)

POLYETHYLENE GLYCOL 3350; POTASSIUM CHLORIDE; SODIUM BICARBONATE; SODIUM CHLORIDE

FOR SOLUTION; ORAL

TRILYTE

<u>AA</u>	SCHWARZ PHARMA	<u>420GM/BOT;1.48GM/BOT;5.72GM/BOT;11.2GM/BOT</u>	<u>N76491</u>	<u>001</u>	Feb 05, 2004
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POLYETHYLENE GLYCOL 3350; POTASSIUM CHLORIDE; SODIUM BICARBONATE; SODIUM CHLORIDE; SODIUM SULFATE, ANHYDROUS

FOR SOLUTION; ORAL

COLYTE

<u>AA</u>	+ SCHWARZ PHARMA	<u>227.1GM/BOT;2.82GM/BOT;6.36GM/BOT;5.53GM/BOT;21.5GM/BOT</u>	<u>N18983</u>	<u>010</u>	Jan 31, 1989
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<u>AA</u>	+	<u>240GM/BOT;2.98GM/BOT;6.72GM/BOT;5.84GM/BOT;22.72GM/BOT</u>	<u>N18983</u>	<u>007</u>	Jun 12, 1987
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COLYTE WITH FLAVOR PACKS

<u>AA</u>	+ SCHWARZ PHARMA	<u>240GM/BOT;2.98GM/BOT;6.72GM/BOT;5.84GM/BOT;22.72GM/BOT</u>	<u>N18983</u>	<u>012</u>	Oct 08, 1998
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COLYTE-FLAVORED

<u>AA</u>	+ SCHWARZ PHARMA	<u>227.1GM/BOT;2.82GM/BOT;6.36GM/BOT;5.53GM/BOT;21.5GM/BOT</u>	<u>N18983</u>	<u>008</u>	Nov 14, 1991
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<u>AA</u>	+	<u>240GM/BOT;2.98GM/BOT;6.72GM/BOT;5.84GM/BOT;22.72GM/BOT</u>	<u>N18983</u>	<u>009</u>	Nov 14, 1991
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GOLYTELY

<u>AA</u>	+ BRAINTREE	<u>227.1GM/PACKET;2.82GM/PACKET;6.36GM/PACKET;5.53GM/PACKET;21.5GM/PACKET</u>	<u>N19011</u>	<u>002</u>	Jun 02, 1992
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<u>AA</u>	+	<u>236GM/BOT;2.97GM/BOT;6.74GM/BOT;5.86GM/BOT;22.74GM/BOT</u>	<u>N19011</u>	<u>001</u>	Jul 13, 1984
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FOR SUSPENSION; ORAL

CO-LAV

<u>AA</u>	COPLEY PHARM	<u>240GM/BOT;2.98GM/BOT;6.72GM/BOT;5.84GM/BOT;22.72GM/BOT</u>	<u>N73428</u>	<u>001</u>	Jan 28, 1992
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GO-EVAC

<u>AA</u>	COPLEY PHARM	<u>236GM/BOT;2.97GM/BOT;6.74GM/BOT;5.86GM/BOT;22.74GM/BOT</u>	<u>N73433</u>	<u>001</u>	Apr 28, 1992
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POLYMYXIN B SULFATE

INJECTABLE; INJECTION

POLYMYXIN B SULFATE

<u>AP</u>	+ BEDFORD	<u>EQ 500,000 U BASE/VIAL</u>	<u>N60716</u>	<u>001</u>	
<u>AP</u>	X GEN PHARMS	<u>EQ 500,000 U BASE/VIAL</u>	<u>N63000</u>	<u>001</u>	Sep 30, 1994

POWDER; FOR RX COMPOUNDING

POLY-RX

+	X GEN PHARMS	100,000,000 UNITS/BOT	N61578	001	
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POLYMYXIN B SULFATE; TRIMETHOPRIM

SOLUTION/DROPS; OPHTHALMIC

TRIMETHOPRIM SULFATE AND POLYMYXIN B SULFATE

<u>AT</u>	TAYLOR PHARMA	<u>10,000 UNITS/ML;EQ 1MG BASE/ML</u>	<u>N65006</u>	<u>001</u>	Dec 17, 1998
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POLYMYXIN B SULFATE; TRIMETHOPRIM SULFATE

SOLUTION/DROPS; OPHTHALMIC

POLYTRIM

<u>AT</u>	+ ALLERGAN	<u>10,000 UNITS/ML;EQ 1MG BASE/ML</u>	<u>N50567</u>	<u>001</u>	Oct 20, 1988
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TRIMETHOPRIM SULFATE AND POLYMYXIN B SULFATE

<u>AT</u>	BAUSCH AND LOMB	<u>10,000 UNITS/ML;EQ 1MG BASE/ML</u>	<u>N64120</u>	<u>001</u>	Feb 14, 1997
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<u>AT</u>	FALCON PHARMS	<u>10,000 UNITS/ML;EQ 1MG BASE/ML</u>	<u>N64211</u>	<u>001</u>	Apr 13, 1998
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POLYTHIAZIDE

TABLET; ORAL

RENESE

PFIZER

1MG

N12845 001

PRESCRIPTION DRUG PRODUCT LIST

3 - 301 (of 370)

POLYTHIAZIDE

TABLET; ORAL			
RENESE			
PFIZER	2MG	N12845	002
+	4MG	N12845	003

POLYTHIAZIDE; PRAZOSIN HYDROCHLORIDE

CAPSULE; ORAL			
MINIZIDE			
PFIZER	0.5MG;EQ 1MG BASE	N17986	001
	0.5MG;EQ 2MG BASE	N17986	002
+	0.5MG;EQ 5MG BASE	N17986	003

POLYTHIAZIDE; RESERPINE

TABLET; ORAL			
RENESE-R			
+	PFIZER	2MG;0.25MG	N13636 001

PORACTANT ALFA

SUSPENSION; INTRATRACHEAL			
CUROSURF			
+	DEY	80MG/ML	N20744 001 Nov 18, 1999

PORFIMER SODIUM

INJECTABLE; INJECTION			
PHOTOFRIN			
+	AXCAN SCANDIPHARM	75MG/VIAL	N20451 001 Dec 27, 1995

POSACONAZOLE

SUSPENSION; ORAL			
NOXAFIL			
+	SCHERING	40MG/ML	N22003 001 Sep 15, 2006

POTASSIUM ACETATE

INJECTABLE; INJECTION			
POTASSIUM ACETATE IN PLASTIC CONTAINER			
+	HOSPIRA	2MEQ/ML	N18896 001 Jul 20, 1984

POTASSIUM AMINOSALICYLATE

CAPSULE; ORAL			
PASKALIUM			
+	GLENWOOD	500MG	N09395 004

TABLET; ORAL			
PASKALIUM			
+	GLENWOOD	1GM	N09395 003

POTASSIUM CHLORIDE

CAPSULE, EXTENDED RELEASE; ORAL			
<u>MICRO-K</u>			
<u>AB</u> KV PHARM	<u>8MEQ</u>	<u>N18238</u>	<u>001</u>
<u>MICRO-K 10</u>			
<u>AB</u> + KV PHARM	<u>10MEQ</u>	<u>N18238</u>	<u>002</u> May 14, 1984
<u>POTASSIUM CHLORIDE</u>			
<u>AB</u> KV PHARM	<u>10MEQ</u>	<u>N70980</u>	<u>001</u> Feb 17, 1987
<u>AB</u> TEVA	<u>8MEQ</u>	<u>N73531</u>	<u>001</u> Apr 26, 1996
<u>AB</u>	<u>10MEQ</u>	<u>N73532</u>	<u>001</u> Apr 26, 1996

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 302 (of 370)

POTASSIUM CHLORIDE

INJECTABLE; INJECTION

POTASSIUM CHLORIDE

<u>AP</u>	ABRAXIS PHARM	<u>2MEQ/ML</u>	<u>N80225</u>	<u>001</u>	
	+	3MEQ/ML	N80225	003	
<u>AP</u>	B BRAUN	<u>2MEQ/ML</u>	<u>N85870</u>	<u>001</u>	
<u>AP</u>	BAXTER HLTHCARE	<u>2MEQ/ML</u>	<u>N85499</u>	<u>001</u>	
<u>AP</u>	+	<u>2MEQ/ML</u>	<u>N80205</u>	<u>001</u>	
<u>AP</u>	HOSPIRA	<u>2MEQ/ML</u>	<u>N83345</u>	<u>002</u>	
	+	1.5MEQ/ML	N83345	001	
<u>AP</u>	INTL MEDICATION	<u>2MEQ/ML</u>	<u>N83163</u>	<u>001</u>	
<u>AP</u>	LUITPOLD	<u>2MEQ/ML</u>	<u>N87584</u>	<u>001</u>	
<u>AP</u>		<u>2MEQ/ML</u>	<u>N87585</u>	<u>001</u>	
<u>POTASSIUM CHLORIDE 10MEQ IN PLASTIC CONTAINER</u>					
<u>AP</u>	+	<u>14.9MG/ML</u>	<u>N19904</u>	<u>001</u>	Dec 26, 1989
<u>AP</u>	+	<u>746MG/100ML</u>	<u>N19904</u>	<u>005</u>	Dec 17, 1990
<u>AP</u>	HOSPIRA	<u>14.9MG/ML</u>	<u>N20161</u>	<u>005</u>	Nov 30, 1992
<u>AP</u>		<u>745MG/100ML</u>	<u>N20161</u>	<u>001</u>	Nov 30, 1992
<u>POTASSIUM CHLORIDE 20MEQ IN PLASTIC CONTAINER</u>					
<u>AP</u>	+	<u>29.8MG/ML</u>	<u>N19904</u>	<u>002</u>	Dec 26, 1989
<u>AP</u>	+	<u>1.49GM/100ML</u>	<u>N19904</u>	<u>006</u>	Dec 17, 1990
<u>AP</u>	+	<u>29.8MG/ML</u>	<u>N20161</u>	<u>006</u>	Aug 11, 1998
<u>AP</u>		<u>1.49GM/100ML</u>	<u>N20161</u>	<u>002</u>	Nov 30, 1992
<u>POTASSIUM CHLORIDE 30MEQ IN PLASTIC CONTAINER</u>					
<u>AP</u>	+	<u>2.24GM/100ML</u>	<u>N19904</u>	<u>003</u>	Dec 26, 1989
<u>AP</u>	+	<u>2.24GM/100ML</u>	<u>N20161</u>	<u>003</u>	Aug 11, 1998
<u>POTASSIUM CHLORIDE 40MEQ IN PLASTIC CONTAINER</u>					
<u>AP</u>	+	<u>2.98GM/100ML</u>	<u>N19904</u>	<u>004</u>	Dec 26, 1989
<u>AP</u>	+	<u>2.98GM/100ML</u>	<u>N20161</u>	<u>004</u>	Aug 11, 1998
<u>POTASSIUM CHLORIDE IN PLASTIC CONTAINER</u>					
<u>AP</u>	ABRAXIS PHARM	<u>2MEQ/ML</u>	<u>N88901</u>	<u>001</u>	Jan 25, 1985
<u>AP</u>		<u>2MEQ/ML</u>	<u>N88908</u>	<u>001</u>	Jan 25, 1985
TABLET, EXTENDED RELEASE; ORAL					
K+10					
BC	ALRA	10MEQ	N70999	001	Oct 22, 1987
<u>K+8</u>					
<u>AB</u>	ALRA	<u>8MEQ</u>	<u>N70998</u>	<u>001</u>	Jan 25, 1993
KAON CL-10					
BC	SAVAGE LABS	10MEQ	N17046	002	
<u>K-DUR 10</u>					
<u>AB</u>	KEY PHARMS	<u>10MEQ</u>	<u>N19439</u>	<u>002</u>	Jun 13, 1986
<u>K-DUR 20</u>					
<u>AB</u>	+	<u>20MEQ</u>	<u>N19439</u>	<u>001</u>	Jun 13, 1986
<u>KLOR-CON</u>					
<u>AB</u>	UPSHER SMITH	<u>8MEQ</u>	<u>N19123</u>	<u>001</u>	Apr 17, 1986
BC		10MEQ	N19123	002	Apr 17, 1986
<u>KLOR-CON M10</u>					
<u>AB</u>	UPSHER SMITH	<u>10MEQ</u>	<u>N74726</u>	<u>002</u>	Aug 09, 2000
KLOR-CON M15					
	UPSHER SMITH	15MEQ	N74726	003	Jun 06, 2003
<u>KLOR-CON M20</u>					
<u>AB</u>	UPSHER SMITH	<u>20MEQ</u>	<u>N74726</u>	<u>001</u>	Nov 20, 1998
KLOTRIX					
BC	APOTHECON	10MEQ	N17850	001	
K-TAB					
BC	ABBOTT	10MEQ	N18279	001	
POTASSIUM CHLORIDE					
BC	ABBOTT	8MEQ	N18279	002	Aug 01, 1988
<u>AB</u>	ANDRX PHARMS	<u>10MEQ</u>	<u>N75604</u>	<u>001</u>	Apr 10, 2002
<u>AB</u>		<u>20MEQ</u>	<u>N75604</u>	<u>002</u>	Apr 10, 2002
<u>AB</u>	+	<u>8MEQ</u>	<u>N70618</u>	<u>001</u>	Sep 09, 1987

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 303 (of 370)

POTASSIUM CHLORIDETABLET, EXTENDED RELEASE; ORAL
POTASSIUM CHLORIDE

<u>AB</u>	EURAND	<u>20MEQ</u>	<u>N76368</u>	<u>001</u>	Aug 18, 2004
<u>AB</u>	KV PHARM	<u>20MEQ</u>	<u>N76044</u>	<u>001</u>	Apr 05, 2002

POTASSIUM CHLORIDE; SODIUM CHLORIDE

INJECTABLE; INJECTION

<u>POTASSIUM CHLORIDE 0.15% IN SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER</u>					
<u>AP</u>	B BRAUN	<u>150MG/100ML;900MG/100ML</u>	<u>N19708</u>	<u>004</u>	Sep 29, 1989
<u>POTASSIUM CHLORIDE 20MEQ IN SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER</u>					
<u>AP</u>	HOSPIRA	<u>149MG/100ML;900MG/100ML</u>	<u>N19686</u>	<u>001</u>	Oct 17, 1988
<u>POTASSIUM CHLORIDE 40MEQ IN SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER</u>					
<u>AP</u>	HOSPIRA	<u>298MG/100ML;900MG/100ML</u>	<u>N19686</u>	<u>002</u>	Oct 17, 1988
SODIUM CHLORIDE 0.45% AND POTASSIUM CHLORIDE 0.15% IN PLASTIC CONTAINER					
+	BAXTER HLTHCARE	450MG/100ML;150MG/100ML	N17648	005	Nov 26, 2002
<u>SODIUM CHLORIDE 0.9% AND POTASSIUM CHLORIDE 0.15% IN PLASTIC CONTAINER</u>					
<u>AP</u>	BAXTER HLTHCARE	<u>150MG/100ML;900MG/100ML</u>	<u>N17648</u>	<u>001</u>	
SODIUM CHLORIDE 0.9% AND POTASSIUM CHLORIDE 0.224%					
	BAXTER HLTHCARE	224MG/100ML;900MG/100ML	N17648	003	
<u>SODIUM CHLORIDE 0.9% AND POTASSIUM CHLORIDE 0.3% IN PLASTIC CONTAINER</u>					
<u>AP</u>	BAXTER HLTHCARE	<u>300MG/100ML;900MG/100ML</u>	<u>N17648</u>	<u>002</u>	

POTASSIUM CITRATE

TABLET, EXTENDED RELEASE; ORAL

<u>POTASSIUM CITRATE</u>					
<u>AB</u>	COREPHARMA	<u>5MEQ</u>	<u>N77440</u>	<u>001</u>	Jun 09, 2006
<u>AB</u>		<u>10MEQ</u>	<u>N77440</u>	<u>002</u>	Jun 09, 2006
<u>UROCIT-K</u>					
<u>AB</u>	MISSION PHARMA	<u>5MEQ</u>	<u>N19071</u>	<u>001</u>	Aug 30, 1985
<u>AB</u>	+	<u>10MEQ</u>	<u>N19071</u>	<u>002</u>	Aug 31, 1992

POVIDONE-IODINE

SOLUTION/DROPS; OPHTHALMIC

+	ALCON	5%	N18634	001	Dec 17, 1986
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PRALIDOXIME CHLORIDE

INJECTABLE; INJECTION

PRALIDOXIME CHLORIDE

+	MERIDIAN MEDCL TECHN	300MG/ML	N18986	001	Apr 26, 1983
PROTOPAM CHLORIDE					
+	BAXTER HLTHCARE CORP	1GM/VIAL	N14134	001	

PRAMIPEXOLE DIHYDROCHLORIDE

TABLET; ORAL

MIRAPEX

	BOEHRINGER INGELHEIM	0.125MG	N20667	001	Jul 01, 1997
+		0.25MG	N20667	002	Jul 01, 1997
		0.5MG	N20667	006	Feb 12, 1998
		1MG	N20667	003	Jul 01, 1997
		1.5MG	N20667	005	Jul 01, 1997

PRAMLINTIDE ACETATE

INJECTABLE; SUBCUTANEOUS

SYMLIN

+	AMYLIN	EQ 3MG BASE/5ML (EQ 0.6MG BASE/ML)	N21332	001	Mar 16, 2005
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27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 304 (of 370)

PRAVASTATIN SODIUM

TABLET; ORAL

PRAVACHOL

<u>AB</u>	BRISTOL MYERS SQUIBB	<u>10MG</u>	<u>N19898</u>	<u>002</u>	Oct 31, 1991
<u>AB</u>		<u>20MG</u>	<u>N19898</u>	<u>003</u>	Oct 31, 1991
<u>AB</u>		<u>40MG</u>	<u>N19898</u>	<u>004</u>	Mar 22, 1993
	+	80MG	N19898	008	Dec 18, 2001

PRAVASTATIN SODIUM

<u>AB</u>	APOTEX	<u>10MG</u>	<u>N76341</u>	<u>001</u>	Oct 23, 2006
<u>AB</u>		<u>20MG</u>	<u>N76341</u>	<u>002</u>	Oct 23, 2006
<u>AB</u>		<u>40MG</u>	<u>N76341</u>	<u>003</u>	Oct 23, 2006
<u>AB</u>	COBALT	<u>10MG</u>	<u>N76939</u>	<u>004</u>	Oct 23, 2006
<u>AB</u>		<u>20MG</u>	<u>N76939</u>	<u>003</u>	Oct 23, 2006
<u>AB</u>		<u>40MG</u>	<u>N76939</u>	<u>002</u>	Oct 23, 2006
<u>AB</u>	DR REDDYS LABS INC	<u>10MG</u>	<u>N76714</u>	<u>001</u>	Oct 23, 2006
<u>AB</u>		<u>20MG</u>	<u>N76714</u>	<u>002</u>	Oct 23, 2006
<u>AB</u>		<u>40MG</u>	<u>N76714</u>	<u>003</u>	Oct 23, 2006
<u>AB</u>	GENPHARM	<u>10MG</u>	<u>N77013</u>	<u>001</u>	Oct 23, 2006
<u>AB</u>		<u>20MG</u>	<u>N77013</u>	<u>002</u>	Oct 23, 2006
<u>AB</u>		<u>40MG</u>	<u>N77013</u>	<u>003</u>	Oct 23, 2006
<u>AB</u>	LEK PHARMS DD	<u>10MG</u>	<u>N76397</u>	<u>003</u>	Oct 23, 2006
<u>AB</u>		<u>20MG</u>	<u>N76397</u>	<u>002</u>	Oct 23, 2006
<u>AB</u>		<u>40MG</u>	<u>N76397</u>	<u>001</u>	Oct 23, 2006
<u>AB</u>	PLIVA HRVATSKA DOO	<u>10MG</u>	<u>N77730</u>	<u>001</u>	Nov 21, 2006
<u>AB</u>		<u>20MG</u>	<u>N77730</u>	<u>002</u>	Nov 21, 2006
<u>AB</u>		<u>40MG</u>	<u>N77730</u>	<u>005</u>	Nov 21, 2006
		30MG	N77730	003	Nov 21, 2006
<u>AB</u>	TEVA	<u>10MG</u>	<u>N76056</u>	<u>001</u>	Apr 24, 2006
<u>AB</u>		<u>20MG</u>	<u>N76056</u>	<u>002</u>	Apr 24, 2006
<u>AB</u>		<u>40MG</u>	<u>N76056</u>	<u>003</u>	Apr 24, 2006
<u>AB</u>	WATSON LABS	<u>10MG</u>	<u>N77904</u>	<u>001</u>	Oct 23, 2006
<u>AB</u>		<u>20MG</u>	<u>N77904</u>	<u>002</u>	Oct 23, 2006
<u>AB</u>		<u>40MG</u>	<u>N77904</u>	<u>003</u>	Oct 23, 2006

PRAZQUANTEL

TABLET; ORAL

BILTRICIDE

	+	BAYER PHARMS	600MG	N18714	001	Dec 29, 1982
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PRAZOSIN HYDROCHLORIDE

CAPSULE; ORAL

MINIPRESS

<u>AB</u>	PFIZER	<u>EQ 1MG BASE</u>	<u>N17442</u>	<u>002</u>	
<u>AB</u>	+	<u>EQ 2MG BASE</u>	<u>N17442</u>	<u>003</u>	
<u>AB</u>		<u>EQ 5MG BASE</u>	<u>N17442</u>	<u>001</u>	

PRAZOSIN HYDROCHLORIDE

<u>AB</u>	IVAX PHARMS	<u>EQ 1MG BASE</u>	<u>N71994</u>	<u>001</u>	Sep 12, 1988
<u>AB</u>		<u>EQ 2MG BASE</u>	<u>N71995</u>	<u>001</u>	Sep 12, 1988
<u>AB</u>		<u>EQ 5MG BASE</u>	<u>N71745</u>	<u>001</u>	Sep 12, 1988
<u>AB</u>	MYLAN	<u>EQ 1MG BASE</u>	<u>N72573</u>	<u>001</u>	May 16, 1989
<u>AB</u>		<u>EQ 2MG BASE</u>	<u>N72574</u>	<u>001</u>	May 16, 1989
<u>AB</u>		<u>EQ 5MG BASE</u>	<u>N72575</u>	<u>001</u>	May 16, 1989
<u>AB</u>	SANDOZ	<u>EQ 1MG BASE</u>	<u>N72576</u>	<u>001</u>	May 16, 1989
<u>AB</u>		<u>EQ 2MG BASE</u>	<u>N72577</u>	<u>001</u>	May 16, 1989
<u>AB</u>		<u>EQ 5MG BASE</u>	<u>N72578</u>	<u>001</u>	May 16, 1989
<u>AB</u>	WATSON LABS	<u>EQ 1MG BASE</u>	<u>N72352</u>	<u>001</u>	May 16, 1989
<u>AB</u>		<u>EQ 2MG BASE</u>	<u>N72333</u>	<u>001</u>	May 16, 1989
<u>AB</u>		<u>EQ 5MG BASE</u>	<u>N72609</u>	<u>001</u>	May 16, 1989

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 305 (of 370)

PRAZOSIN HYDROCHLORIDE

TABLET, EXTENDED RELEASE; ORAL

MINIPRESS XL

+	PFIZER	2.5MG	N19775	001	Jan 29, 1992
+		5MG	N19775	002	Jan 29, 1992

PREDNICARBATE

CREAM; TOPICAL

DERMATOP E EMOLLIENT

<u>AB</u>	+	SANOFI AVENTIS US	<u>0.1%</u>	<u>N20279</u>	<u>001</u>	Oct 29, 1993
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PREDNICARBATE

<u>AB</u>		ALTANA	<u>0.1%</u>	<u>N77287</u>	<u>001</u>	Sep 19, 2006
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OINTMENT; TOPICAL

DERMATOP

+	SANOFI AVENTIS US	0.1%	N19568	001	Sep 23, 1991
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PREDNISOLONE

SYRUP; ORAL

PREDNISOLONE

<u>AA</u>		APOTEX INC	<u>5MG/5ML</u>	<u>N40570</u>	<u>001</u>	Aug 25, 2005
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<u>AA</u>			<u>15MG/5ML</u>	<u>N40571</u>	<u>001</u>	Aug 25, 2005
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<u>AA</u>		HI TECH PHARMA	<u>15MG/5ML</u>	<u>N40401</u>	<u>001</u>	Feb 27, 2003
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<u>AA</u>		IVAX PHARMS	<u>15MG/5ML</u>	<u>N40287</u>	<u>001</u>	May 28, 1999
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<u>AA</u>	+	KV PHARM	<u>5MG/5ML</u>	<u>N40423</u>	<u>001</u>	Oct 22, 2001
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<u>AA</u>	+		<u>15MG/5ML</u>	<u>N40364</u>	<u>001</u>	Apr 10, 2002
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<u>AA</u>		MORTON GROVE	<u>15MG/5ML</u>	<u>N40313</u>	<u>001</u>	Sep 10, 2003
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<u>AA</u>		PHARM ASSOC	<u>15MG/5ML</u>	<u>N40399</u>	<u>001</u>	Mar 05, 2003
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<u>AA</u>		TEVA PHARMS	<u>15MG/5ML</u>	<u>N40322</u>	<u>001</u>	Jan 19, 2000
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<u>AA</u>		UDL	<u>15MG/5ML</u>	<u>N40323</u>	<u>001</u>	May 13, 1999
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<u>AA</u>		WE PHARMS	<u>15MG/5ML</u>	<u>N40192</u>	<u>001</u>	May 28, 1998
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PRELONE

<u>AA</u>		TEVA	<u>15MG/5ML</u>	<u>N89081</u>	<u>001</u>	Feb 04, 1986
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TABLET; ORAL

PREDNISOLONE

BX		LANNETT	5MG	N80531	002	
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BX	+	WATSON LABS	5MG	N80354	001	
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PREDNISOLONE ACETATE

SUSPENSION/DROPS; OPHTHALMIC

OMNIPRED

<u>AB</u>		ALCON	<u>1%</u>	<u>N17469</u>	<u>001</u>	
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PRED FORTE

<u>AB</u>	+	ALLERGAN	<u>1%</u>	<u>N17011</u>	<u>001</u>	
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PRED MILD

+	ALLERGAN	0.12%	N17100	001	
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PREDNISOLONE ACETATE; SULFACETAMIDE SODIUM

OINTMENT; OPHTHALMIC

BLEPHAMIDE S.O.P.

+	ALLERGAN	0.2%;10%	N87748	001	Dec 03, 1986
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SUSPENSION; OPHTHALMIC

BLEPHAMIDE

+	ALLERGAN	0.2%;10%	N12813	002	
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PREDNISOLONE SODIUM PHOSPHATE

SOLUTION; ORAL

ORAPRED

<u>AA</u>	+	ASCENT PEDS	<u>EQ 15MG BASE/5ML</u>	<u>N75117</u>	<u>001</u>	Dec 14, 2000
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27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 306 (of 370)

PREDNISOLONE SODIUM PHOSPHATE

SOLUTION; ORAL

PEDIAPRED

<u>AA</u>	+	UCB INC	<u>EQ 5MG BASE/5ML</u>	<u>N19157</u>	<u>001</u>	May 28, 1986
<u>PREDNISOLONE SODIUM PHOSPHATE</u>						
<u>AA</u>		HI TECH PHARMA	<u>EQ 5MG BASE/5ML</u>	<u>N75183</u>	<u>001</u>	Mar 26, 2003
<u>AA</u>		KV PHARM	<u>EQ 5MG BASE/5ML</u>	<u>N76982</u>	<u>001</u>	May 24, 2005
<u>AA</u>			<u>EQ 15MG BASE/5ML</u>	<u>N76988</u>	<u>001</u>	May 24, 2005
<u>AA</u>		MORTON GROVE	<u>EQ 5MG BASE/5ML</u>	<u>N75099</u>	<u>001</u>	Jun 28, 2002
<u>AA</u>			<u>EQ 15MG BASE/5ML</u>	<u>N76895</u>	<u>001</u>	Oct 04, 2004
<u>AA</u>		PADDOCK	<u>EQ 5MG BASE/5ML</u>	<u>N75988</u>	<u>001</u>	May 25, 2004
<u>AA</u>		PHARM ASSOC	<u>EQ 5MG BASE/5ML</u>	<u>N76123</u>	<u>001</u>	Dec 23, 2002
<u>AA</u>			<u>EQ 15MG BASE/5ML</u>	<u>N76913</u>	<u>001</u>	Apr 25, 2005
<u>AA</u>		WE PHARMS	<u>EQ 5MG BASE/5ML</u>	<u>N75181</u>	<u>001</u>	Dec 23, 2002
<u>AA</u>			<u>EQ 15MG BASE/5ML</u>	<u>N75250</u>	<u>001</u>	Jul 12, 2002

SOLUTION/DROPS; OPHTHALMIC

INFLAMASE FORTE

<u>AT</u>	+	NOVARTIS	<u>EQ 0.9% PHOSPHATE</u>	<u>N80751</u>	<u>002</u>	
<u>INFLAMASE MILD</u>						
<u>AT</u>	+	NOVARTIS	<u>EQ 0.11% PHOSPHATE</u>	<u>N80751</u>	<u>001</u>	
<u>PREDNISOLONE SODIUM PHOSPHATE</u>						
<u>AT</u>		BAUSCH AND LOMB	<u>EQ 0.11% PHOSPHATE</u>	<u>N40065</u>	<u>001</u>	Jul 29, 1994
<u>AT</u>			<u>EQ 0.9% PHOSPHATE</u>	<u>N40070</u>	<u>001</u>	Jul 29, 1994

TABLET, ORALLY DISINTEGRATING; ORAL

ORAPRED ODT

MEDICIS

			EQ 10MG BASE	N21959	001	Jun 01, 2006
			EQ 15MG BASE	N21959	002	Jun 01, 2006
+			EQ 30MG BASE	N21959	003	Jun 01, 2006

PREDNISOLONE SODIUM PHOSPHATE; SULFACETAMIDE SODIUM

SOLUTION/DROPS; OPHTHALMIC

SULFACETAMIDE SODIUM AND PREDNISOLONE SODIUM PHOSPHATE

<u>AT</u>		ALCON	<u>EQ 0.23% PHOSPHATE;10%</u>	<u>N73630</u>	<u>001</u>	May 27, 1993
<u>AT</u>		BAUSCH AND LOMB	<u>EQ 0.23% PHOSPHATE;10%</u>	<u>N74449</u>	<u>001</u>	Dec 29, 1995
<u>VASOCIDIN</u>						
<u>AT</u>	+	NOVARTIS	<u>EQ 0.23% PHOSPHATE;10%</u>	<u>N18988</u>	<u>001</u>	Aug 26, 1988

PREDNISONE

SOLUTION; ORAL

PREDNISONE

+	ROXANE	5MG/5ML	N88703	001	Nov 08, 1984
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PREDNISONE INTENSOL

+	ROXANE	5MG/ML	N88810	001	Feb 20, 1985
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TABLET; ORAL

PREDNISONE

<u>AB</u>		JUBILANT PHARMS	<u>1MG</u>	<u>N40611</u>	<u>001</u>	Jun 06, 2005
<u>AB</u>			<u>5MG</u>	<u>N40362</u>	<u>002</u>	Aug 29, 2001
<u>AB</u>			<u>10MG</u>	<u>N40362</u>	<u>001</u>	Aug 29, 2001
<u>AB</u>			<u>20MG</u>	<u>N40362</u>	<u>003</u>	Jun 29, 2005
<u>AB</u>		LEINER	<u>5MG</u>	<u>N80209</u>	<u>001</u>	
<u>AB</u>		MUTUAL PHARM	<u>5MG</u>	<u>N89245</u>	<u>001</u>	Dec 04, 1985
<u>AB</u>			<u>10MG</u>	<u>N89246</u>	<u>001</u>	Dec 04, 1985
<u>AB</u>			<u>20MG</u>	<u>N89247</u>	<u>001</u>	Dec 04, 1985
<u>AB</u>	+	ROXANE	<u>1MG</u>	<u>N87800</u>	<u>001</u>	Apr 22, 1982
<u>AB</u>	+		<u>2.5MG</u>	<u>N87801</u>	<u>001</u>	Apr 22, 1982
<u>AB</u>	+		<u>5MG</u>	<u>N80352</u>	<u>001</u>	
<u>AB</u>	+		<u>10MG</u>	<u>N84122</u>	<u>001</u>	
<u>AB</u>	+		<u>20MG</u>	<u>N87342</u>	<u>001</u>	
<u>AB</u>	+		<u>50MG</u>	<u>N84283</u>	<u>001</u>	

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 307 (of 370)

PREDNISONE

TABLET; ORAL

PREDNISONE

<u>AB</u>	SANDOZ	<u>10MG</u>	<u>N89983</u>	<u>001</u>	Jan 12, 1989
<u>AB</u>		<u>50MG</u>	<u>N89984</u>	<u>001</u>	Jan 12, 1989
<u>AB</u>	VINTAGE PHARMS	<u>1MG</u>	<u>N40584</u>	<u>001</u>	Dec 21, 2004
<u>AB</u>		<u>2.5MG</u>	<u>N40581</u>	<u>001</u>	Dec 21, 2004
<u>AB</u>		<u>5MG</u>	<u>N40256</u>	<u>001</u>	Jul 12, 2002
<u>AB</u>		<u>10MG</u>	<u>N40256</u>	<u>002</u>	Jul 12, 2002
<u>AB</u>		<u>20MG</u>	<u>N40392</u>	<u>001</u>	Feb 12, 2003
<u>AB</u>	WATSON LABS	<u>5MG</u>	<u>N80356</u>	<u>001</u>	
<u>AB</u>		<u>10MG</u>	<u>N85162</u>	<u>001</u>	
<u>AB</u>		<u>10MG</u>	<u>N87773</u>	<u>001</u>	Jul 13, 1982
<u>AB</u>		<u>20MG</u>	<u>N85161</u>	<u>001</u>	
<u>AB</u>		<u>20MG</u>	<u>N86813</u>	<u>001</u>	
<u>AB</u>		<u>50MG</u>	<u>N87772</u>	<u>001</u>	Jul 13, 1982
<u>AB</u>	WEST WARD	<u>2.5MG</u>	<u>N40538</u>	<u>001</u>	Jan 08, 2004
<u>AB</u>		<u>5MG</u>	<u>N80292</u>	<u>001</u>	
<u>AB</u>		<u>10MG</u>	<u>N88832</u>	<u>001</u>	Dec 04, 1985
<u>AB</u>		<u>20MG</u>	<u>N83677</u>	<u>001</u>	

PREGABALIN

CAPSULE; ORAL

LYRICA

CP PHARMS

25MG	N21446	001	Dec 30, 2004
50MG	N21446	002	Dec 30, 2004
75MG	N21446	003	Dec 30, 2004
100MG	N21446	004	Dec 30, 2004
150MG	N21446	005	Dec 30, 2004
200MG	N21446	006	Dec 30, 2004
225MG	N21446	007	Dec 30, 2004
+	N21446	008	Dec 30, 2004
300MG			

PRILOCAINE HYDROCHLORIDE

INJECTABLE; INJECTION

CITANEST PLAIN

+ DENTSPLY PHARM 4%

N21382 001

PRIMAQUINE PHOSPHATE

TABLET; ORAL

PRIMAQUINE

+ SANOFI AVENTIS US EQ 15MG BASE

N08316 001

PRIMIDONE

TABLET; ORAL

MYSOLINE

<u>AB</u>	+ VALEANT	<u>50MG</u>	<u>N09170</u>	<u>003</u>	
<u>AB</u>		<u>250MG</u>	<u>N09170</u>	<u>002</u>	

PRIMIDONE

<u>AB</u>	LANNETT	<u>50MG</u>	<u>N84903</u>	<u>002</u>	May 24, 2001
<u>AB</u>		<u>250MG</u>	<u>N84903</u>	<u>001</u>	
<u>AB</u>	MUTUAL PHARM	<u>50MG</u>	<u>N40626</u>	<u>001</u>	Sep 29, 2005
<u>AB</u>		<u>250MG</u>	<u>N40626</u>	<u>002</u>	Sep 29, 2005
<u>AB</u>	VINTAGE PHARMS	<u>50MG</u>	<u>N40586</u>	<u>001</u>	Feb 24, 2005
<u>AB</u>		<u>250MG</u>	<u>N40586</u>	<u>002</u>	Feb 24, 2005
<u>AB</u>	WATSON LABS	<u>250MG</u>	<u>N83551</u>	<u>001</u>	
<u>AB</u>	WEST WARD	<u>50MG</u>	<u>N40667</u>	<u>001</u>	Jul 27, 2006
<u>AB</u>		<u>250MG</u>	<u>N40667</u>	<u>002</u>	Jul 27, 2006

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 308 (of 370)

PROBENECID

TABLET; ORAL

PROBALAN

<u>AB</u>	LANNETT	<u>500MG</u>	<u>N80966</u>	<u>001</u>	
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PROBENECID

<u>AB</u>	IVAX PHARMS	<u>500MG</u>	<u>N83740</u>	<u>001</u>	May 09, 1984
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<u>AB</u>	+	MYLAN	<u>500MG</u>	<u>N84211</u>	<u>002</u>
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<u>AB</u>		WATSON LABS	<u>500MG</u>	<u>N84442</u>	<u>004</u>
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Mar 29, 1983

PROCAINAMIDE HYDROCHLORIDE

CAPSULE; ORAL

PROCAINAMIDE HYDROCHLORIDE

<u>AB</u>	IVAX PHARMS	<u>250MG</u>	<u>N84604</u>	<u>001</u>	
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<u>AB</u>		<u>375MG</u>	<u>N84595</u>	<u>001</u>	
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<u>AB</u>		<u>500MG</u>	<u>N84606</u>	<u>001</u>	
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<u>AB</u>	WATSON LABS	<u>250MG</u>	<u>N83287</u>	<u>001</u>	
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<u>AB</u>		<u>375MG</u>	<u>N84403</u>	<u>001</u>	
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<u>AB</u>		<u>500MG</u>	<u>N84280</u>	<u>001</u>	
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PRONESTYL

<u>AB</u>	APOTHECON	<u>250MG</u>	<u>N07335</u>	<u>001</u>	
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<u>AB</u>		<u>375MG</u>	<u>N07335</u>	<u>004</u>	
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<u>AB</u>	+	<u>500MG</u>	<u>N07335</u>	<u>003</u>	
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INJECTABLE; INJECTION

PROCAINAMIDE HYDROCHLORIDE

<u>AP</u>	HOSPIRA	<u>100MG/ML</u>	<u>N89069</u>	<u>001</u>	Feb 12, 1986
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<u>AP</u>		<u>500MG/ML</u>	<u>N89070</u>	<u>001</u>	Feb 12, 1986
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<u>AP</u>		<u>500MG/ML</u>	<u>N89537</u>	<u>001</u>	Aug 25, 1987
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<u>AP</u>	INTL MEDICATION	<u>100MG/ML</u>	<u>N88636</u>	<u>001</u>	Jul 31, 1984
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<u>AP</u>		<u>500MG/ML</u>	<u>N88637</u>	<u>001</u>	Jul 31, 1984
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PRONESTYL

<u>AP</u>	+	APOTHECON	<u>100MG/ML</u>	<u>N07335</u>	<u>002</u>
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<u>AP</u>	+	<u>500MG/ML</u>	<u>N07335</u>	<u>005</u>	
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TABLET; ORAL

PRONESTYL

	APOTHECON	250MG	N17371	001	
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		375MG	N17371	002	
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+		500MG	N17371	003	
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TABLET, EXTENDED RELEASE; ORAL

PROCANBID

+	KING PHARMS	500MG	N20545	001	Jan 31, 1996
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+		1GM	N20545	002	Jan 31, 1996
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PRONESTYL-SR

BC	APOTHECON	500MG	N87361	001	
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PROCAINE HYDROCHLORIDE

INJECTABLE; INJECTION

NOVOCAIN

<u>AP</u>	+	HOSPIRA	<u>1%</u>	<u>N85362</u>	<u>003</u>
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<u>AP</u>	+		<u>2%</u>	<u>N85362</u>	<u>004</u>
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+		10%	N86797	001	
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PROCAINE HYDROCHLORIDE

<u>AP</u>	HOSPIRA	<u>1%</u>	<u>N80416</u>	<u>001</u>	
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<u>AP</u>		<u>2%</u>	<u>N80416</u>	<u>002</u>	
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<u>AP</u>	WATSON LABS	<u>1%</u>	<u>N80658</u>	<u>001</u>	
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PROCARBAZINE HYDROCHLORIDE

CAPSULE; ORAL

MATULANE

+	SIGMA TAU	EQ 50MG BASE	N16785	001	
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27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 309 (of 370)

PROCHLORPERAZINE

SUPPOSITORY; RECTAL

COMPAZINE

<u>AB</u>	+	GLAXOSMITHKLINE	<u>25MG</u>	<u>N11127</u>	<u>002</u>	
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COMPRO

<u>AB</u>		PADDOCK	<u>25MG</u>	<u>N40246</u>	<u>001</u>	Jun 28, 2000
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PROCHLORPERAZINE

<u>AB</u>		G AND W LABS	<u>25MG</u>	<u>N40058</u>	<u>001</u>	Nov 24, 1993
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PROCHLORPERAZINE EDISYLATE

INJECTABLE; INJECTION

COMPAZINE

<u>AP</u>	+	GLAXOSMITHKLINE	<u>EQ 5MG BASE/ML</u>	<u>N10742</u>	<u>002</u>	
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PROCHLORPERAZINE

<u>AP</u>		BAXTER HLTHCARE	<u>EQ 5MG BASE/ML</u>	<u>N87759</u>	<u>001</u>	Oct 01, 1982
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PROCHLORPERAZINE EDISYLATE

<u>AP</u>		BAXTER HLTHCARE	<u>EQ 5MG BASE/ML</u>	<u>N89903</u>	<u>001</u>	Aug 29, 1989
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<u>AP</u>		BEDFORD	<u>EQ 5MG BASE/ML</u>	<u>N40540</u>	<u>001</u>	May 28, 2004
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<u>AP</u>		HOSPIRA	<u>EQ 5MG BASE/ML</u>	<u>N89703</u>	<u>001</u>	Apr 07, 1988
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<u>AP</u>		SICOR PHARMS	<u>EQ 5MG BASE/ML</u>	<u>N40505</u>	<u>001</u>	May 30, 2003
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<u>AP</u>		WATSON LABS	<u>EQ 5MG BASE/ML</u>	<u>N89530</u>	<u>001</u>	Jul 08, 1987
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PROCHLORPERAZINE MALEATE

CAPSULE, EXTENDED RELEASE; ORAL

COMPAZINE

	+	GLAXOSMITHKLINE	EQ 15MG BASE	N21019	002	Oct 06, 1999
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TABLET; ORAL

COMPAZINE

<u>AB</u>		GLAXOSMITHKLINE	<u>EQ 5MG BASE</u>	<u>N10571</u>	<u>001</u>	
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<u>AB</u>			<u>EQ 10MG BASE</u>	<u>N10571</u>	<u>002</u>	
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<u>AB</u>	+		<u>EQ 25MG BASE</u>	<u>N10571</u>	<u>003</u>	
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PROCHLORPERAZINE MALEATE

<u>AB</u>		DURAMED PHARMS BARR	<u>EQ 5MG BASE</u>	<u>N40207</u>	<u>001</u>	May 01, 1997
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<u>AB</u>			<u>EQ 10MG BASE</u>	<u>N40207</u>	<u>002</u>	May 01, 1997
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<u>AB</u>		IVAX PHARMS	<u>EQ 5MG BASE</u>	<u>N40162</u>	<u>001</u>	Jan 20, 1998
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<u>AB</u>			<u>EQ 10MG BASE</u>	<u>N40162</u>	<u>002</u>	Jan 20, 1998
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<u>AB</u>		JUBILANT PHARMS	<u>EQ 5MG BASE</u>	<u>N40268</u>	<u>001</u>	Feb 27, 1998
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<u>AB</u>			<u>EQ 10MG BASE</u>	<u>N40268</u>	<u>002</u>	Feb 27, 1998
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<u>AB</u>		MYLAN	<u>EQ 5MG BASE</u>	<u>N40185</u>	<u>002</u>	Oct 28, 1996
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<u>AB</u>			<u>EQ 10MG BASE</u>	<u>N40185</u>	<u>001</u>	Oct 28, 1996
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<u>AB</u>		SANDOZ	<u>EQ 5MG BASE</u>	<u>N40101</u>	<u>001</u>	Jul 19, 1996
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<u>AB</u>			<u>EQ 10MG BASE</u>	<u>N40101</u>	<u>002</u>	Jul 19, 1996
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<u>AB</u>			<u>EQ 25MG BASE</u>	<u>N40101</u>	<u>003</u>	Jul 19, 1996
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<u>AB</u>		TEVA PHARMS	<u>EQ 5MG BASE</u>	<u>N40120</u>	<u>001</u>	Jul 11, 1996
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<u>AB</u>			<u>EQ 10MG BASE</u>	<u>N40120</u>	<u>002</u>	Jul 11, 1996
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PROCYCLIDINE HYDROCHLORIDE

TABLET; ORAL

KEMADRIN

	+	MONARCH PHARMS	5MG	N09818	003	
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PROGESTERONE

CAPSULE; ORAL

PROMETRIUM

		UNIMED PHARMS	100MG	N19781	001	May 14, 1998
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	+		200MG	N19781	002	Oct 15, 1999
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GEL; VAGINAL

CRINONE

		COLUMBIA LABS	4%	N20701	001	Jul 31, 1997
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PRESCRIPTION DRUG PRODUCT LIST

3 - 310 (of 370)

PROGESTERONE

GEL; VAGINAL
CRINONE

+ COLUMBIA LABS	8%	N20701	002	Jul 31, 1997
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INJECTABLE; INJECTION

PROGESTERONE

<u>AO</u> ABRAXIS PHARM	<u>50MG/ML</u>	<u>N75906</u>	<u>001</u>	Apr 25, 2001
<u>AO</u> + WATSON LABS (UTAH)	<u>50MG/ML</u>	<u>N17362</u>	<u>002</u>	

PROMETHAZINE HYDROCHLORIDE

INJECTABLE; INJECTION

PROMETHAZINE HYDROCHLORIDE

<u>AP</u> BAXTER HLTHCARE	<u>25MG/ML</u>	<u>N83312</u>	<u>001</u>	
<u>AP</u>	<u>50MG/ML</u>	<u>N83312</u>	<u>002</u>	
<u>AP</u> BEDFORD LABS	<u>25MG/ML</u>	<u>N40524</u>	<u>001</u>	Mar 17, 2004
<u>AP</u>	<u>50MG/ML</u>	<u>N40524</u>	<u>002</u>	Mar 17, 2004
<u>AP</u> BIONICHE PHARMA	<u>25MG/ML</u>	<u>N40471</u>	<u>001</u>	Nov 21, 2002
<u>AP</u> HOSPIRA	<u>25MG/ML</u>	<u>N40372</u>	<u>001</u>	Jun 08, 2000
<u>AP</u>	<u>50MG/ML</u>	<u>N40372</u>	<u>002</u>	Jun 08, 2000
<u>AP</u>	<u>50MG/ML</u>	<u>N83838</u>	<u>002</u>	
<u>AP</u> PHARMAFORCE	<u>25MG/ML</u>	<u>N40515</u>	<u>001</u>	Mar 19, 2003
<u>AP</u> SANDOZ	<u>25MG/ML</u>	<u>N40593</u>	<u>001</u>	Nov 08, 2006
<u>AP</u>	<u>50MG/ML</u>	<u>N40593</u>	<u>002</u>	Nov 08, 2006
<u>AP</u> + SICOR PHARMS	<u>25MG/ML</u>	<u>N40454</u>	<u>001</u>	Aug 22, 2002
<u>AP</u> +	<u>50MG/ML</u>	<u>N40454</u>	<u>002</u>	Aug 22, 2002
<u>AP</u> WATSON LABS	<u>25MG/ML</u>	<u>N84591</u>	<u>001</u>	
<u>AP</u>	<u>50MG/ML</u>	<u>N80629</u>	<u>002</u>	

SUPPOSITORY; RECTAL

PROMETHACON

BR POLYMEDICA	25MG	N84901	001	
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PROMETHAZINE HYDROCHLORIDE

<u>AB</u> G AND W LABS	<u>12.5MG</u>	<u>N40428</u>	<u>002</u>	Mar 31, 2003
<u>AB</u> +	<u>25MG</u>	<u>N40428</u>	<u>001</u>	Feb 05, 2002
<u>AB</u> PADDOCK	<u>12.5MG</u>	<u>N40479</u>	<u>001</u>	Jun 24, 2003
<u>AB</u>	<u>25MG</u>	<u>N40479</u>	<u>002</u>	Jun 24, 2003
<u>AB</u> PERRIGO NEW YORK	<u>12.5MG</u>	<u>N40500</u>	<u>001</u>	Jun 30, 2003
<u>AB</u>	<u>25MG</u>	<u>N40500</u>	<u>002</u>	Jun 30, 2003
<u>AB</u> TARO	<u>12.5MG</u>	<u>N40603</u>	<u>001</u>	Oct 26, 2006
<u>AB</u>	<u>25MG</u>	<u>N40603</u>	<u>002</u>	Oct 26, 2006

PROMETHEGAN

+ G AND W LABS	50MG	N87165	001	Aug 14, 1987
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SYRUP; ORAL

PROMETHAZINE HYDROCHLORIDE

<u>AA</u> HI TECH PHARMA	<u>6.25MG/5ML</u>	<u>N40026</u>	<u>001</u>	Sep 25, 1998
<u>AA</u> VINTAGE	<u>6.25MG/5ML</u>	<u>N40643</u>	<u>001</u>	Apr 26, 2006

PROMETHAZINE PLAIN

<u>AA</u> + MORTON GROVE	<u>6.25MG/5ML</u>	<u>N87953</u>	<u>001</u>	Nov 15, 1982
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TABLET; ORAL

PHENERGAN

<u>AB</u> WYETH PHARMS INC	<u>25MG</u>	<u>N07935</u>	<u>003</u>	
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PROMETHAZINE HYDROCHLORIDE

<u>AB</u> KVK-TECH INC	<u>25MG</u>	<u>N40712</u>	<u>001</u>	Jul 31, 2006
<u>AB</u>	<u>50MG</u>	<u>N40713</u>	<u>001</u>	Jul 31, 2006
<u>AB</u> SANDOZ	<u>25MG</u>	<u>N84234</u>	<u>001</u>	
<u>AB</u> +	<u>50MG</u>	<u>N84176</u>	<u>001</u>	
<u>AB</u> VINTAGE PHARMS	<u>12.5MG</u>	<u>N40622</u>	<u>001</u>	Jul 18, 2006
<u>AB</u>	<u>25MG</u>	<u>N40622</u>	<u>002</u>	Jul 18, 2006
<u>AB</u>	<u>50MG</u>	<u>N40622</u>	<u>003</u>	Jul 18, 2006
BP WATSON LABS	25MG	N83426	001	

PRESCRIPTION DRUG PRODUCT LIST

3 - 311 (of 370)

PROMETHAZINE HYDROCHLORIDE

TABLET; ORAL

PROMETHAZINE HYDROCHLORIDE

BP	WATSON LABS	50MG	N83711	001	
<u>AB</u>	ZYDUS PHARMS USA	<u>12.5MG</u>	<u>N40596</u>	<u>001</u>	Nov 18, 2005
<u>AB</u>		<u>25MG</u>	<u>N40596</u>	<u>002</u>	Nov 18, 2005
<u>AB</u>		<u>50MG</u>	<u>N40596</u>	<u>003</u>	Nov 18, 2005

PROPAFENONE HYDROCHLORIDE

CAPSULE, EXTENDED RELEASE; ORAL
RYTHMOL SR

	RELIANT PHARMS	225MG	N21416	001	Sep 04, 2003
		325MG	N21416	002	Sep 04, 2003
+		425MG	N21416	003	Sep 04, 2003

TABLET; ORAL

PROPAFENONE HYDROCHLORIDE

<u>AB</u>	KV PHARM	<u>150MG</u>	<u>N76193</u>	<u>001</u>	Feb 07, 2002
<u>AB</u>		<u>225MG</u>	<u>N76193</u>	<u>002</u>	Feb 07, 2002
<u>AB</u>		<u>300MG</u>	<u>N76193</u>	<u>003</u>	Feb 07, 2002
<u>AB</u>	MUTUAL PHARM	<u>150MG</u>	<u>N75998</u>	<u>001</u>	Nov 29, 2001
<u>AB</u>		<u>225MG</u>	<u>N75998</u>	<u>002</u>	Nov 29, 2001
<u>AB</u>		<u>300MG</u>	<u>N75998</u>	<u>003</u>	Nov 29, 2001
<u>AB</u>	PLIVA	<u>150MG</u>	<u>N76550</u>	<u>001</u>	Apr 23, 2004
<u>AB</u>		<u>225MG</u>	<u>N76550</u>	<u>002</u>	Apr 23, 2004
<u>AB</u>		<u>300MG</u>	<u>N76550</u>	<u>003</u>	Apr 23, 2004
<u>AB</u>	VINTAGE PHARMS	<u>150MG</u>	<u>N75938</u>	<u>001</u>	Oct 17, 2002
<u>AB</u>		<u>225MG</u>	<u>N75938</u>	<u>002</u>	Oct 17, 2002
<u>AB</u>		<u>300MG</u>	<u>N75938</u>	<u>003</u>	Oct 17, 2002
<u>AB</u>	WATSON LABS	<u>150MG</u>	<u>N75203</u>	<u>001</u>	Oct 24, 2000
<u>AB</u>		<u>225MG</u>	<u>N75203</u>	<u>002</u>	Oct 24, 2000
	<u>RYTHMOL</u>				
<u>AB</u>	RELIANT PHARMS	<u>150MG</u>	<u>N19151</u>	<u>001</u>	Nov 27, 1989
<u>AB</u>		<u>225MG</u>	<u>N19151</u>	<u>003</u>	Nov 20, 1992
<u>AB</u>	+	<u>300MG</u>	<u>N19151</u>	<u>002</u>	Nov 27, 1989

PROPANTHELINE BROMIDE

TABLET; ORAL
PRO-BANTHINE

BP	+	SHIRE	7.5MG	N08732	003
BP	+		15MG	N08732	002
		PROPANTHELINE BROMIDE			
BP		ROXANE	7.5MG	N80927	001
BP			15MG	N80927	002

PROPARACAINE HYDROCHLORIDE

SOLUTION; OPHTHALMIC

PROPARACAINE HYDROCHLORIDE

<u>AT</u>	TAYLOR PHARMA	<u>0.5%</u>	<u>N40277</u>	<u>001</u>	Mar 16, 2000
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SOLUTION/DROPS; OPHTHALMIC

ALCAINE

<u>AT</u>	ALCON	<u>0.5%</u>	<u>N80027</u>	<u>001</u>	
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OPHTHAINE

<u>AT</u>	+	APOTHECON	<u>0.5%</u>	<u>N08883</u>	<u>001</u>
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OPHTHETIC

<u>AT</u>	+	ALLERGAN	<u>0.5%</u>	<u>N12583</u>	<u>001</u>
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PARACAINE

<u>AT</u>	OPTOPICS	<u>0.5%</u>	<u>N87681</u>	<u>001</u>	Aug 05, 1982
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PROPARACAINE HYDROCHLORIDE

<u>AT</u>	BAUSCH AND LOMB	<u>0.5%</u>	<u>N40074</u>	<u>001</u>	Sep 29, 1995
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27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 312 (of 370)

PROPOFOL

INJECTABLE; INJECTION

DIPRIVAN

<u>AB</u>	+	ABRAXIS BIOSCIENCE	<u>10MG/ML</u>	<u>N19627</u>	<u>002</u>	Jun 11, 1996
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PROPOFOL

<u>AB</u>		BEDFORD	<u>10MG/ML</u>	<u>N74848</u>	<u>001</u>	Apr 19, 2005
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<u>AB</u>		HOSPIRA	<u>10MG/ML</u>	<u>N77908</u>	<u>001</u>	Mar 17, 2006
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<u>AB</u>		SICOR PHARMS	<u>10MG/ML</u>	<u>N75102</u>	<u>001</u>	Jan 04, 1999
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<u>AB</u>			<u>10MG/ML</u>	<u>N75392</u>	<u>001</u>	Sep 19, 2000
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PROPOXYPHENE HYDROCHLORIDE

CAPSULE; ORAL

DARVON

<u>AA</u>	+	XANODYNE PHARM	<u>65MG</u>	<u>N10997</u>	<u>003</u>	
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PROPOXYPHENE HYDROCHLORIDE

<u>AA</u>		MYLAN	<u>65MG</u>	<u>N40569</u>	<u>001</u>	Dec 16, 2004
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<u>AA</u>		PAR PHARM	<u>65MG</u>	<u>N80269</u>	<u>001</u>	
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<u>AA</u>		TEVA	<u>65MG</u>	<u>N88615</u>	<u>001</u>	Oct 22, 1984
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<u>AA</u>		WEST WARD	<u>65MG</u>	<u>N83501</u>	<u>001</u>	
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PROPOXYPHENE NAPSYLATE

TABLET; ORAL

DARVON-N

	+	XANODYNE PHARM	100MG	N16862	002	
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PROPRANOLOL HYDROCHLORIDE

CAPSULE, EXTENDED RELEASE; ORAL

INDERAL LA

BX		WYETH PHARMS INC	80MG	N18553	002	Apr 19, 1983
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BX			120MG	N18553	003	Apr 19, 1983
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			60MG	N18553	004	Mar 18, 1987
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	+		160MG	N18553	001	Apr 19, 1983
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INNOPRAN XL

BX		RELIANT PHARMS	80MG	N21438	001	Mar 12, 2003
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BX			120MG	N21438	002	Mar 12, 2003
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CONCENTRATE; ORAL

PROPRANOLOL HYDROCHLORIDE INTENSOL

	+	ROXANE	80MG/ML	N71388	001	May 15, 1987
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INJECTABLE; INJECTION

PROPRANOLOL

<u>AP</u>		ABRAXIS PHARM	<u>1MG/ML</u>	<u>N75826</u>	<u>001</u>	Aug 31, 2001
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PROPRANOLOL HYDROCHLORIDE

<u>AP</u>	+	BAXTER HLTHCARE CORP	<u>1MG/ML</u>	<u>N16419</u>	<u>001</u>	
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<u>AP</u>		BEDFORD	<u>1MG/ML</u>	<u>N75792</u>	<u>001</u>	Aug 29, 2000
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<u>AP</u>		SANDOZ	<u>1MG/ML</u>	<u>N76400</u>	<u>001</u>	Feb 26, 2003
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SOLUTION; ORAL

PROPRANOLOL HYDROCHLORIDE

	+	ROXANE	20MG/5ML	N70979	001	May 15, 1987
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	+		40MG/5ML	N70690	001	May 15, 1987
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TABLET; ORAL

INDERAL

<u>AB</u>		WYETH PHARMS INC	<u>10MG</u>	<u>N16418</u>	<u>001</u>	
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<u>AB</u>			<u>20MG</u>	<u>N16418</u>	<u>003</u>	
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<u>AB</u>			<u>40MG</u>	<u>N16418</u>	<u>002</u>	
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<u>AB</u>			<u>60MG</u>	<u>N16418</u>	<u>009</u>	Oct 18, 1982
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<u>AB</u>	+		<u>80MG</u>	<u>N16418</u>	<u>004</u>	
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PROPRANOLOL HYDROCHLORIDE

<u>AB</u>		MYLAN	<u>10MG</u>	<u>N70211</u>	<u>001</u>	Nov 19, 1985
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<u>AB</u>			<u>20MG</u>	<u>N70212</u>	<u>001</u>	Nov 19, 1985
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27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 313 (of 370)

PROPRANOLOL HYDROCHLORIDE

TABLET; ORAL

PROPRANOLOL HYDROCHLORIDE

<u>AB</u>	MYLAN	<u>40MG</u>	<u>N70213</u>	<u>001</u>	Nov 19, 1985
<u>AB</u>		<u>80MG</u>	<u>N70214</u>	<u>001</u>	Nov 19, 1985
<u>AB</u>	PAR PHARM	<u>10MG</u>	<u>N70217</u>	<u>001</u>	Aug 01, 1986
<u>AB</u>		<u>20MG</u>	<u>N70218</u>	<u>001</u>	Aug 01, 1986
<u>AB</u>		<u>40MG</u>	<u>N70219</u>	<u>001</u>	Aug 01, 1986
<u>AB</u>		<u>60MG</u>	<u>N70220</u>	<u>001</u>	Sep 24, 1986
<u>AB</u>		<u>80MG</u>	<u>N70221</u>	<u>001</u>	Apr 14, 1986
<u>AB</u>		<u>90MG</u>	<u>N71288</u>	<u>001</u>	Oct 22, 1986
<u>AB</u>	PLIVA	<u>10MG</u>	<u>N71972</u>	<u>001</u>	Apr 06, 1988
<u>AB</u>		<u>20MG</u>	<u>N71973</u>	<u>001</u>	Apr 06, 1988
<u>AB</u>		<u>40MG</u>	<u>N71974</u>	<u>001</u>	Apr 06, 1988
<u>AB</u>		<u>60MG</u>	<u>N71975</u>	<u>001</u>	Apr 06, 1988
<u>AB</u>		<u>80MG</u>	<u>N71976</u>	<u>001</u>	Apr 06, 1988
<u>AB</u>		<u>90MG</u>	<u>N71977</u>	<u>001</u>	Apr 06, 1988
<u>AB</u>	SANDOZ	<u>10MG</u>	<u>N70663</u>	<u>001</u>	Jun 13, 1986
<u>AB</u>		<u>20MG</u>	<u>N70664</u>	<u>001</u>	Jun 13, 1986
<u>AB</u>		<u>40MG</u>	<u>N70665</u>	<u>001</u>	Jun 13, 1986
<u>AB</u>		<u>60MG</u>	<u>N70666</u>	<u>001</u>	Oct 10, 1986
<u>AB</u>		<u>80MG</u>	<u>N70667</u>	<u>001</u>	Jun 13, 1986
<u>AB</u>	TEVA	<u>40MG</u>	<u>N70234</u>	<u>001</u>	Jun 23, 1986
<u>AB</u>	WATSON LABS	<u>10MG</u>	<u>N70175</u>	<u>001</u>	May 13, 1986
<u>AB</u>		<u>10MG</u>	<u>N70548</u>	<u>001</u>	Jul 10, 1986
<u>AB</u>		<u>20MG</u>	<u>N70176</u>	<u>001</u>	May 13, 1986
<u>AB</u>		<u>40MG</u>	<u>N70177</u>	<u>001</u>	May 13, 1986
<u>AB</u>		<u>40MG</u>	<u>N70550</u>	<u>001</u>	Apr 11, 1986
<u>AB</u>		<u>60MG</u>	<u>N71791</u>	<u>001</u>	Jul 15, 1987
<u>AB</u>		<u>80MG</u>	<u>N70178</u>	<u>001</u>	May 13, 1986
<u>AB</u>		<u>80MG</u>	<u>N70551</u>	<u>001</u>	Jul 10, 1986
<u>AB</u>		<u>90MG</u>	<u>N71792</u>	<u>001</u>	Jul 15, 1987

PROPYLTHIOURACIL

TABLET; ORAL

PROPYLTHIOURACIL

BD	ACTAVIS ELIZABETH	50MG	N80172	001	
BD +	STADA PHARMS	50MG	N06188	001	
BD	WEST WARD	50MG	N80154	001	

PROTAMINE SULFATE

INJECTABLE; INJECTION

PROTAMINE SULFATE

+	ABRAXIS PHARM	10MG/ML	N89454	001	Apr 07, 1987
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PROTRIPTYLINE HYDROCHLORIDE

TABLET; ORAL

VIVACTIL

<u>AB</u>	ODYSSEY PHARMS	<u>5MG</u>	<u>N73644</u>	<u>001</u>	Aug 24, 1995
<u>AB</u> +		<u>10MG</u>	<u>N73645</u>	<u>001</u>	Aug 24, 1995

PSEUDOEPHEDRINE HYDROCHLORIDE; TRIPROLIDINE HYDROCHLORIDE

TABLET; ORAL

CORPHED

<u>AA</u>	SANDOZ	<u>60MG;2.5MG</u>	<u>N88602</u>	<u>001</u>	Apr 11, 1985
<u>AA</u> +	SANDOZ	<u>60MG;2.5MG</u>	<u>N88193</u>	<u>001</u>	May 17, 1983

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 314 (of 370)

PYRAZINAMIDE

TABLET; ORAL

PYRAZINAMIDE

<u>AB</u>	+	CLONMEL HLTHCARE	<u>500MG</u>	<u>N80157</u>	<u>001</u>	
<u>AB</u>		MIKART	<u>500MG</u>	<u>N81319</u>	<u>001</u>	Jun 30, 1992

PYRIDOSTIGMINE BROMIDE

INJECTABLE; INJECTION

MESTINON

<u>AP</u>	+	VALEANT PHARM INTL	<u>5MG/ML</u>	<u>N09830</u>	<u>001</u>	
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REGONOL

<u>AP</u>		SANDOZ	<u>5MG/ML</u>	<u>N17398</u>	<u>001</u>	
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SYRUP; ORAL

MESTINON

	+	VALEANT PHARM INTL	60MG/5ML	N15193	001	
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TABLET; ORAL

MESTINON

<u>AB</u>	+	VALEANT PHARM INTL	<u>60MG</u>	<u>N09829</u>	<u>002</u>	
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PYRIDOSTIGMINE BROMIDE

<u>AB</u>		BARR	<u>60MG</u>	<u>N40512</u>	<u>001</u>	Oct 08, 2003
			30MG	N40512	002	Jul 20, 2005

<u>AB</u>		COREPHARMA	<u>60MG</u>	<u>N40457</u>	<u>001</u>	Dec 26, 2002
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<u>AB</u>		IMPAX LABS	<u>60MG</u>	<u>N40502</u>	<u>001</u>	Apr 24, 2003
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TABLET, EXTENDED RELEASE; ORAL

MESTINON

	+	VALEANT PHARM INTL	180MG	N11665	001	
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PYRIDOXINE HYDROCHLORIDE

INJECTABLE; INJECTION

PYRIDOXINE HYDROCHLORIDE

<u>AP</u>		ABRAXIS PHARM	<u>100MG/ML</u>	<u>N80618</u>	<u>001</u>	
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<u>AP</u>	+	WATSON LABS	<u>100MG/ML</u>	<u>N80572</u>	<u>001</u>	
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PRIMETHAMINE

TABLET; ORAL

DARAPRIM

	+	GLAXOSMITHKLINE	25MG	N08578	001	
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PRIMETHAMINE; SULFADOXINE

TABLET; ORAL

FANSIDAR

	+	ROCHE	25MG;500MG	N18557	001	
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QUAZEPAM

TABLET; ORAL

DORAL

	+	QUESTCOR PHARMS	15MG	N18708	001	Dec 27, 1985
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QUETIAPINE FUMARATE

TABLET; ORAL

SEROQUEL

	+	ASTRAZENECA	EQ 25MG BASE	N20639	001	Sep 26, 1997
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			EQ 50MG BASE	N20639	007	Oct 04, 2005
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			EQ 100MG BASE	N20639	002	Sep 26, 1997
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			EQ 200MG BASE	N20639	003	Sep 26, 1997
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+			EQ 300MG BASE	N20639	005	Jul 26, 2000
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			EQ 400MG BASE	N20639	006	Oct 04, 2005
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27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 315 (of 370)

QUINAPRIL HYDROCHLORIDE

TABLET; ORAL

ACCUPRIL

<u>AB</u>	PFIZER PHARMS	<u>EQ 5MG BASE</u>	<u>N19885</u>	<u>001</u>	Nov 19, 1991
<u>AB</u>		<u>EQ 10MG BASE</u>	<u>N19885</u>	<u>002</u>	Nov 19, 1991
<u>AB</u>		<u>EQ 20MG BASE</u>	<u>N19885</u>	<u>003</u>	Nov 19, 1991
<u>AB</u>	+	<u>EQ 40MG BASE</u>	<u>N19885</u>	<u>004</u>	Nov 19, 1991

QUINAPRIL

<u>AB</u>	LUPIN	<u>EQ 5MG BASE</u>	<u>N77690</u>	<u>001</u>	Jun 20, 2006
<u>AB</u>		<u>EQ 10MG BASE</u>	<u>N77690</u>	<u>002</u>	Jun 20, 2006
<u>AB</u>		<u>EQ 20MG BASE</u>	<u>N77690</u>	<u>003</u>	Jun 20, 2006
<u>AB</u>		<u>EQ 40MG BASE</u>	<u>N77690</u>	<u>004</u>	Jun 20, 2006
<u>AB</u>	RANBAXY	<u>EQ 5MG BASE</u>	<u>N76607</u>	<u>001</u>	Dec 15, 2004
<u>AB</u>		<u>EQ 10MG BASE</u>	<u>N76607</u>	<u>002</u>	Dec 15, 2004
<u>AB</u>		<u>EQ 20MG BASE</u>	<u>N76607</u>	<u>003</u>	Dec 15, 2004
<u>AB</u>		<u>EQ 40MG BASE</u>	<u>N76607</u>	<u>004</u>	Dec 15, 2004

QUINAPRIL HYDROCHLORIDE

<u>AB</u>	ACTAVIS TOTOWA	<u>EQ 5MG BASE</u>	<u>N76459</u>	<u>001</u>	Dec 22, 2004
<u>AB</u>		<u>EQ 10MG BASE</u>	<u>N76459</u>	<u>002</u>	Dec 22, 2004
<u>AB</u>		<u>EQ 20MG BASE</u>	<u>N76459</u>	<u>003</u>	Dec 22, 2004
<u>AB</u>		<u>EQ 40MG BASE</u>	<u>N76459</u>	<u>004</u>	Dec 22, 2004
<u>AB</u>	ANDRX PHARMS	<u>EQ 5MG BASE</u>	<u>N76049</u>	<u>001</u>	Jan 14, 2005
<u>AB</u>		<u>EQ 10MG BASE</u>	<u>N76049</u>	<u>002</u>	Jan 14, 2005
<u>AB</u>		<u>EQ 20MG BASE</u>	<u>N76049</u>	<u>003</u>	Jan 14, 2005
<u>AB</u>		<u>EQ 40MG BASE</u>	<u>N76049</u>	<u>004</u>	Jan 14, 2005
<u>AB</u>	GENPHARM	<u>EQ 5MG BASE</u>	<u>N76036</u>	<u>001</u>	Jan 28, 2005
<u>AB</u>		<u>EQ 10MG BASE</u>	<u>N76036</u>	<u>002</u>	Jan 28, 2005
<u>AB</u>		<u>EQ 20MG BASE</u>	<u>N76036</u>	<u>003</u>	Jan 28, 2005
<u>AB</u>		<u>EQ 40MG BASE</u>	<u>N76036</u>	<u>004</u>	Jan 28, 2005
<u>AB</u>	MYLAN	<u>EQ 5MG BASE</u>	<u>N76694</u>	<u>001</u>	Dec 23, 2004
<u>AB</u>		<u>EQ 10MG BASE</u>	<u>N76694</u>	<u>002</u>	Dec 23, 2004
<u>AB</u>		<u>EQ 20MG BASE</u>	<u>N76694</u>	<u>003</u>	Dec 23, 2004
<u>AB</u>		<u>EQ 40MG BASE</u>	<u>N76694</u>	<u>004</u>	Dec 23, 2004
<u>AB</u>	SANDOZ	<u>EQ 5MG BASE</u>	<u>N76803</u>	<u>001</u>	Mar 02, 2005
<u>AB</u>		<u>EQ 10MG BASE</u>	<u>N76803</u>	<u>002</u>	Mar 02, 2005
<u>AB</u>		<u>EQ 20MG BASE</u>	<u>N76803</u>	<u>003</u>	Mar 02, 2005
<u>AB</u>		<u>EQ 40MG BASE</u>	<u>N76803</u>	<u>004</u>	Mar 02, 2005
<u>AB</u>	TORPHARM	<u>EQ 5MG BASE</u>	<u>N76240</u>	<u>001</u>	Jan 26, 2006
<u>AB</u>		<u>EQ 10MG BASE</u>	<u>N76240</u>	<u>002</u>	Jan 26, 2006
<u>AB</u>		<u>EQ 20MG BASE</u>	<u>N76240</u>	<u>003</u>	Jan 26, 2006
<u>AB</u>		<u>EQ 40MG BASE</u>	<u>N76240</u>	<u>004</u>	Jan 26, 2006

QUINIDINE GLUCONATE

INJECTABLE; INJECTION

QUINIDINE GLUCONATE

+	LILLY	80MG/ML	N07529	002	Feb 10, 1989
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TABLET, EXTENDED RELEASE; ORAL

QUINIDINE GLUCONATE

BX	+	MUTUAL PHARM	324MG	N89338	001	Feb 11, 1987
BX		WATSON LABS	324MG	N87810	001	Sep 29, 1982

QUINIDINE SULFATE

TABLET; ORAL

QUINIDINE SULFATE

<u>AB</u>	MUTUAL PHARM	<u>200MG</u>	<u>N81030</u>	<u>001</u>	Apr 14, 1989
<u>AB</u>		<u>300MG</u>	<u>N81031</u>	<u>001</u>	Apr 14, 1989
<u>AB</u>	SANDOZ	<u>200MG</u>	<u>N88072</u>	<u>002</u>	
<u>AB</u>		<u>300MG</u>	<u>N88072</u>	<u>001</u>	Sep 26, 1983
<u>AB</u>	WATSON LABS	<u>200MG</u>	<u>N83288</u>	<u>001</u>	
<u>AB</u>		<u>300MG</u>	<u>N85583</u>	<u>001</u>	

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 316 (of 370)

QUINIDINE SULFATE

TABLET, EXTENDED RELEASE; ORAL

QUINIDINE SULFATE

+	TEVA PHARMS	300MG	N40045	001	Jun 30, 1994
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QUININE SULFATE

CAPSULE; ORAL

QUININE SULFATE

+	AR HOLDING CO INC	324MG	N21799	001	Aug 12, 2005
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RABEPRAZOLE SODIUM

TABLET, DELAYED RELEASE; ORAL

ACIPHEX

+	EISAI MEDCL RES	20MG	N20973	002	Aug 19, 1999
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RALOXIFENE HYDROCHLORIDE

TABLET; ORAL

EVISTA

+	LILLY	60MG	N20815	001	Dec 09, 1997
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RAMELTEON

TABLET; ORAL

ROZEREM

+	TAKEDA GLOBAL	8MG	N21782	001	Jul 22, 2005
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RAMIPRIL

CAPSULE; ORAL

ALTACE

<u>AB</u>	KING PHARMS	<u>1.25MG</u>	<u>N19901</u>	<u>001</u>	Jan 28, 1991
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<u>AB</u>		<u>2.5MG</u>	<u>N19901</u>	<u>002</u>	Jan 28, 1991
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<u>AB</u>		<u>5MG</u>	<u>N19901</u>	<u>003</u>	Jan 28, 1991
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<u>AB</u>	+	<u>10MG</u>	<u>N19901</u>	<u>004</u>	Jan 28, 1991
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RAMIPRIL

<u>AB</u>	COBALT	<u>1.25MG</u>	<u>N76549</u>	<u>001</u>	Oct 24, 2005
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<u>AB</u>		<u>2.5MG</u>	<u>N76549</u>	<u>002</u>	Oct 24, 2005
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<u>AB</u>		<u>5MG</u>	<u>N76549</u>	<u>003</u>	Oct 24, 2005
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<u>AB</u>		<u>10MG</u>	<u>N76549</u>	<u>004</u>	Oct 24, 2005
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RANITIDINE HYDROCHLORIDE

CAPSULE; ORAL

RANITIDINE HYDROCHLORIDE

<u>AB</u>	DR REDDYS LABS LTD	<u>EQ 150MG BASE</u>	<u>N75742</u>	<u>001</u>	Nov 29, 2000
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<u>AB</u>		<u>EQ 300MG BASE</u>	<u>N75742</u>	<u>002</u>	Nov 29, 2000
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<u>AB</u>	GENPHARM	<u>EQ 150MG BASE</u>	<u>N75564</u>	<u>001</u>	Oct 27, 2000
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<u>AB</u>		<u>EQ 300MG BASE</u>	<u>N75564</u>	<u>002</u>	Oct 27, 2000
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<u>AB</u>	SANDOZ	<u>EQ 150MG BASE</u>	<u>N74655</u>	<u>001</u>	Oct 22, 1997
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<u>AB</u>	+	<u>EQ 300MG BASE</u>	<u>N74655</u>	<u>002</u>	Oct 22, 1997
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<u>AB</u>	TEVA	<u>EQ 150MG BASE</u>	<u>N75557</u>	<u>001</u>	Oct 31, 2003
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<u>AB</u>		<u>EQ 300MG BASE</u>	<u>N75557</u>	<u>002</u>	Oct 31, 2003
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GRANULE, EFFERVESCENT; ORAL

ZANTAC 150

+	GLAXOSMITHKLINE	EQ 150MG BASE/PACKET	N20251	002	Mar 31, 1994
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INJECTABLE; INJECTION

RANITIDINE

<u>AP</u>	BEDFORD	<u>EQ 25MG BASE/ML</u>	<u>N74764</u>	<u>001</u>	Nov 19, 2004
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RANITIDINE HYDROCHLORIDE

<u>AP</u>	BEDFORD	<u>EQ 25MG BASE/ML</u>	<u>N77458</u>	<u>001</u>	Feb 16, 2006
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<u>AP</u>	BEN VENUE	<u>EQ 25MG BASE/ML</u>	<u>N74777</u>	<u>001</u>	Mar 02, 2005
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27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 317 (of 370)

RANITIDINE HYDROCHLORIDE

INJECTABLE; INJECTION

ZANTAC

<u>AP</u>	+	GLAXOSMITHKLINE	<u>EQ 25MG BASE/ML</u>	<u>N19090</u>	<u>001</u>	Oct 19, 1984
		ZANTAC IN PLASTIC CONTAINER				
	+	GLAXOSMITHKLINE	EQ 1MG BASE/ML	N19593	002	Sep 27, 1991

SYRUP; ORAL

ZANTAC

	+	GLAXOSMITHKLINE	EQ 15MG BASE/ML	N19675	001	Dec 30, 1988
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TABLET; ORAL

RANITIDINE

<u>AB</u>		RANBAXY	<u>EQ 150MG BASE</u>	<u>N75439</u>	<u>001</u>	Apr 19, 2000
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<u>AB</u>			<u>EQ 300MG BASE</u>	<u>N75439</u>	<u>002</u>	Apr 19, 2000
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RANITIDINE HYDROCHLORIDE

<u>AB</u>		COBALT	<u>EQ 150MG BASE</u>	<u>N77426</u>	<u>001</u>	Dec 19, 2005
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<u>AB</u>			<u>EQ 300MG BASE</u>	<u>N77426</u>	<u>002</u>	Dec 19, 2005
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<u>AB</u>		DR REDDYS LABS INC	<u>EQ 150MG BASE</u>	<u>N76705</u>	<u>001</u>	Jul 27, 2005
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<u>AB</u>			<u>EQ 300MG BASE</u>	<u>N76705</u>	<u>002</u>	Jul 27, 2005
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<u>AB</u>		GENPHARM	<u>EQ 150MG BASE</u>	<u>N74023</u>	<u>001</u>	Aug 22, 1997
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<u>AB</u>			<u>EQ 300MG BASE</u>	<u>N74023</u>	<u>002</u>	Aug 22, 1997
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<u>AB</u>		INTERPHARM	<u>EQ 150MG BASE</u>	<u>N77824</u>	<u>001</u>	Oct 13, 2006
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<u>AB</u>			<u>EQ 300MG BASE</u>	<u>N77824</u>	<u>002</u>	Oct 13, 2006
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<u>AB</u>		IVAX PHARMS	<u>EQ 150MG BASE</u>	<u>N75165</u>	<u>001</u>	Sep 30, 1998
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<u>AB</u>			<u>EQ 300MG BASE</u>	<u>N75165</u>	<u>002</u>	Sep 30, 1998
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<u>AB</u>		MYLAN	<u>EQ 150MG BASE</u>	<u>N74552</u>	<u>001</u>	Jul 30, 1998
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<u>AB</u>			<u>EQ 300MG BASE</u>	<u>N74552</u>	<u>002</u>	Jul 30, 1998
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<u>AB</u>		PAR PHARM	<u>EQ 150MG BASE</u>	<u>N75180</u>	<u>001</u>	Jan 28, 1999
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<u>AB</u>			<u>EQ 300MG BASE</u>	<u>N75180</u>	<u>002</u>	Jan 28, 1999
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<u>AB</u>		SANDOZ	<u>EQ 150MG BASE</u>	<u>N74467</u>	<u>001</u>	Aug 29, 1997
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<u>AB</u>			<u>EQ 300MG BASE</u>	<u>N74467</u>	<u>002</u>	Aug 29, 1997
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<u>AB</u>		TEVA	<u>EQ 150MG BASE</u>	<u>N74488</u>	<u>001</u>	Jul 31, 1997
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<u>AB</u>			<u>EQ 300MG BASE</u>	<u>N74488</u>	<u>002</u>	Jul 31, 1997
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<u>AB</u>		TORPHARM	<u>EQ 150MG BASE</u>	<u>N74680</u>	<u>001</u>	Sep 12, 1997
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<u>AB</u>			<u>EQ 300MG BASE</u>	<u>N74680</u>	<u>002</u>	Sep 12, 1997
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<u>AB</u>		WATSON LABS	<u>EQ 150MG BASE</u>	<u>N74864</u>	<u>001</u>	Oct 20, 1997
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<u>AB</u>			<u>EQ 300MG BASE</u>	<u>N74864</u>	<u>002</u>	Oct 20, 1997
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<u>AB</u>		WOCKHARDT	<u>EQ 150MG BASE</u>	<u>N75208</u>	<u>001</u>	Dec 17, 1998
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<u>AB</u>			<u>EQ 300MG BASE</u>	<u>N75208</u>	<u>002</u>	Dec 17, 1998
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ZANTAC 150

<u>AB</u>		GLAXOSMITHKLINE	<u>EQ 150MG BASE</u>	<u>N18703</u>	<u>001</u>	Jun 09, 1983
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ZANTAC 300

<u>AB</u>	+	GLAXOSMITHKLINE	<u>EQ 300MG BASE</u>	<u>N18703</u>	<u>002</u>	Dec 09, 1985
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TABLET, EFFERVESCENT; ORAL

ZANTAC 150

	+	GLAXOSMITHKLINE	EQ 150MG BASE	N20251	001	Mar 31, 1994
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ZANTAC 25

		GLAXOSMITHKLINE	EQ 25MG BASE	N20251	003	Apr 01, 2004
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RANOLAZINE

TABLET, EXTENDED RELEASE; ORAL

RANEXA

	+	CV THERAP	500MG	N21526	002	Jan 27, 2006
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RASAGILINE MESYLATE

TABLET; ORAL

AZILECT

TEVA

	+		EQ 0.5MG BASE	N21641	001	May 16, 2006
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			EQ 1MG BASE	N21641	002	May 16, 2006
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27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 318 (of 370)

REMIFENTANIL HYDROCHLORIDE

INJECTABLE; INJECTION

ULTIVA

ABBOTT

EQ 1MG BASE/VIAL

N20630 001

Jul 12, 1996

EQ 2MG BASE/VIAL

N20630 002

Jul 12, 1996

+

EQ 5MG BASE/VIAL

N20630 003

Jul 12, 1996

REPAGLINIDE

TABLET; ORAL

PRANDIN

NOVO NORDISK INC

0.5MG

N20741 001

Dec 22, 1997

1MG

N20741 002

Dec 22, 1997

+

2MG

N20741 003

Dec 22, 1997

RESCINNAMINE

TABLET; ORAL

MODERIL

PFIZER

0.25MG

N10686 003

+

0.5MG

N10686 006

RESERPINE

TABLET; ORAL

RESERPINE

BP SANDOZ

0.1MG

N09838 001

BP +

0.25MG

N09838 002

SERPALAN

BP LANNETT

0.1MG

N10124 001

BP

0.25MG

N10124 002

RIBAVIRIN

CAPSULE; ORAL

REBETOLAB + SCHERING PLOUGH RES200MGN20903 002

Jul 25, 2001

+

200MG ***Indicated for use and
comarketed with interferon alfa-2b,
recombinant (Intron A), as Rebetron
Combination Therapy***

N20903 001

Jun 03, 1998

RIBASPHEREAB THREE RIVERS PHARMS200MGN76203 001

Apr 06, 2004

RIBAVIRINAB SANDOZ200MGN76192 001

Apr 06, 2004

AB TEVA200MGN76277 001

Oct 04, 2004

AB ZYDUS PHARMS USA200MGN77224 001

Oct 28, 2005

FOR SOLUTION; INHALATION

VIRAZOLE

+ VALEANT PHARM INTL

6GM/VIAL

N18859 001

Dec 31, 1985

SOLUTION; ORAL

REBETOL

+ SCHERING

40MG/ML

N21546 001

Jul 29, 2003

TABLET; ORAL

COPEGUSAB ROCHE200MGN21511 001

Dec 03, 2002

RIBAVIRINAB SANDOZ200MGN77743 001

Oct 03, 2006

AB TEVA200MGN77053 001

Dec 05, 2005

AB THREE RIVERS PHARMS200MGN77456 001

Dec 05, 2005

400MG

N77456 002

Dec 05, 2005

+

600MG

N77456 003

Dec 05, 2005

AB ZYDUS PHARMS USA200MGN77094 001

Dec 05, 2005

PRESCRIPTION DRUG PRODUCT LIST

3 - 319 (of 370)

RIFABUTIN

CAPSULE; ORAL
MYCOBUTIN

+ PHARMACIA AND UPJOHN	150MG	N50689	001	Dec 23, 1992
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RIFAMPIN

CAPSULE; ORAL

RIFADIN

AB SANOFI AVENTIS US	150MG	N62303	001	
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AB +	300MG	N50420	001	
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RIFAMPIN

AB SANDOZ	150MG	N64150	002	Jan 02, 1998
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AB	300MG	N64150	001	May 28, 1997
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AB VERSAPHARM	150MG	N65028	001	Mar 14, 2001
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AB	300MG	N65028	002	Mar 14, 2001
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RIMACTANE

AB ACTAVIS TOTOWA	300MG	N50429	001	
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INJECTABLE; INJECTION

RIFADIN

AP + SANOFI AVENTIS US	600MG/VIAL	N50627	001	May 25, 1989
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RIFAMPIN

AP BEDFORD	600MG/VIAL	N64217	001	Oct 29, 1999
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RIFAPENTINE

TABLET; ORAL
PRIFTIN

+ SANOFI AVENTIS US	150MG	N21024	001	Jun 22, 1998
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RIFAXIMIN

TABLET; ORAL
XIFAXAN

+ SALIX PHARMS	200MG	N21361	001	May 25, 2004
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RILUZOLE

TABLET; ORAL

RILUTEK

AB + SANOFI AVENTIS US	50MG	N20599	001	Dec 12, 1995
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RILUZOLE

AB IMPAX LABS	50MG	N76173	001	Jan 29, 2003
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RIMANTADINE HYDROCHLORIDE

SYRUP; ORAL
FLUMADINE

+ FOREST LABS	50MG/5ML	N19650	001	Sep 17, 1993
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TABLET; ORAL

FLUMADINE

AB + FOREST LABS	100MG	N19649	001	Sep 17, 1993
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RIMANTADINE HYDROCHLORIDE

AB ACTAVIS TOTOWA	100MG	N76375	001	Jan 14, 2003
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AB COREPHARMA	100MG	N75916	001	Nov 02, 2001
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AB IMPAX LABS	100MG	N76132	001	Aug 30, 2002
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RIMEXOLONE

SUSPENSION/DROPS; OPHTHALMIC

+ ALCON	1%	N20474	001	Dec 30, 1994
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27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 320 (of 370)

RISEDRONATE SODIUM

TABLET; ORAL

ACTONEL

PROCTER AND GAMBLE

5MG

N20835 002 Apr 14, 2000

30MG

N20835 001 Mar 27, 1998

+

35MG

N20835 003 May 25, 2002

RISPERIDONE

INJECTABLE; INTRAMUSCULAR

RISPERDAL CONSTA

JANSSEN PHARMA

25MG/VIAL

N21346 001 Oct 29, 2003

37.5MG/VIAL

N21346 002 Oct 29, 2003

+

50MG/VIAL

N21346 003 Oct 29, 2003

SOLUTION; ORAL

RISPERDAL

+ JANSSEN PHARMA

1MG/ML

N20588 001 Jun 10, 1996

TABLET; ORAL

RISPERDAL

JANSSEN PHARMA

0.25MG

N20272 008 May 10, 1999

0.5MG

N20272 007 Jan 27, 1999

+

1MG

N20272 001 Dec 29, 1993

2MG

N20272 002 Dec 29, 1993

3MG

N20272 003 Dec 29, 1993

4MG

N20272 004 Dec 29, 1993

TABLET, ORALLY DISINTEGRATING; ORAL

RISPERDAL

JANSSEN PHARMA

0.5MG

N21444 001 Apr 02, 2003

+

1MG

N21444 002 Apr 02, 2003

2MG

N21444 003 Apr 02, 2003

3MG

N21444 004 Dec 23, 2004

4MG

N21444 005 Dec 23, 2004

RITODRINE HYDROCHLORIDE

INJECTABLE; INJECTION

RITODRINE HYDROCHLORIDE

+ HOSPIRA

10MG/ML

N71618 001 Feb 28, 1991

+

15MG/ML

N71619 001 Feb 28, 1991

RITODRINE HYDROCHLORIDE IN DEXTROSE 5% IN PLASTIC CONTAINER

+ HOSPIRA

30MG/100ML

N71438 001 Jan 22, 1991

RITONAVIR

CAPSULE; ORAL

NORVIR

+ ABBOTT

100MG

N20945 001 Jun 29, 1999

SOLUTION; ORAL

NORVIR

+ ABBOTT

80MG/ML

N20659 001 Mar 01, 1996

RIVASTIGMINE TARTRATE

CAPSULE; ORAL

EXELON

+ NOVARTIS

EQ 1.5MG BASE

N20823 003 Apr 21, 2000

EQ 3MG BASE

N20823 004 Apr 21, 2000

EQ 4.5MG BASE

N20823 005 Apr 21, 2000

+

EQ 6MG BASE

N20823 006 Apr 21, 2000

SOLUTION; ORAL

EXELON

+ NOVARTIS

EQ 2MG BASE/ML

N21025 001 Apr 21, 2000

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 321 (of 370)

RIZATRIPTAN BENZOATE

TABLET; ORAL

MAXALT

MERCK

EQ 5MG BASE

N20864 001

Jun 29, 1998

+

EQ 10MG BASE

N20864 002

Jun 29, 1998

TABLET, ORALLY DISINTEGRATING; ORAL

MAXALT-MLT

MERCK

EQ 5MG BASE

N20865 001

Jun 29, 1998

+

EQ 10MG BASE

N20865 002

Jun 29, 1998

ROCURONIUM BROMIDE

INJECTABLE; INJECTION

ZEMURON

+ ORGANON USA INC

50MG/5ML (10MG/ML)

N20214 001

Mar 17, 1994

+

100MG/10ML (10MG/ML)

N20214 003

Mar 17, 1994

ROPINIROLE HYDROCHLORIDE

TABLET; ORAL

REQUIP

+ GLAXOSMITHKLINE

EQ 0.25MG BASE

N20658 001

Sep 19, 1997

EQ 0.5MG BASE

N20658 002

Sep 19, 1997

EQ 1MG BASE

N20658 003

Sep 19, 1997

EQ 2MG BASE

N20658 004

Sep 19, 1997

EQ 3MG BASE

N20658 006

Jan 27, 1999

EQ 4MG BASE

N20658 007

Jan 27, 1999

EQ 5MG BASE

N20658 005

Sep 19, 1997

ROIIVACAINE HYDROCHLORIDE MONOHYDRATE

INJECTABLE; INJECTION

NAROPIN

ABRAXIS BIOSCIENCE

2MG/ML

N20533 001

Sep 24, 1996

5MG/ML

N20533 003

Sep 24, 1996

7.5MG/ML

N20533 004

Sep 24, 1996

+

10MG/ML

N20533 005

Sep 24, 1996

ROSIGLITAZONE MALEATE

TABLET; ORAL

AVANDIA

SB PHARMCO

EQ 2MG BASE

N21071 002

May 25, 1999

EQ 4MG BASE

N21071 003

May 25, 1999

+

EQ 8MG BASE

N21071 004

May 25, 1999

ROSUVASTATIN CALCIUM

TABLET; ORAL

CRESTOR

ASTRAZENECA

5MG

N21366 002

Aug 12, 2003

10MG

N21366 003

Aug 12, 2003

20MG

N21366 004

Aug 12, 2003

+

40MG

N21366 005

Aug 12, 2003

RUBIDIUM CHLORIDE RB-82

INJECTABLE; INJECTION

CARDIOGEN-82

BRACCO

N/A

N19414 001

Dec 29, 1989

SACROSIDASE

SOLUTION; ORAL

SUCRAID

+ QOL MEDCL

8,500 IU/ML

N20772 001

Apr 09, 1998

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 322 (of 370)

SAFFLOWER OIL; SOYBEAN OIL

INJECTABLE; INJECTION

LIPOSYN II 10%

+ HOSPIRA 5%;5% (5GM/100ML) N18997 001 Aug 27, 1984

LIPOSYN II 20%

+ HOSPIRA 10%;10% (10GM/100ML) N18991 001 Aug 27, 1984

SALMETEROL XINAFOATE

POWDER; INHALATION

SEREVENT

+ GLAXO GRP LTD EQ 0.046MG BASE/INH N20692 001 Sep 19, 1997

SAMARIUM SM 153 LEXIDRONAM PENTASODIUM

INJECTABLE; INJECTION

QUADRAMET

+ CYTOGEN 50mCi/ML N20570 001 Mar 28, 1997

SAQUINAVIR MESYLATE

CAPSULE; ORAL

INVIRASE

+ HLR EQ 200MG BASE N20628 001 Dec 06, 1995

TABLET; ORAL

INVIRASE

+ ROCHE EQ 500MG BASE N21785 001 Dec 17, 2004

SCOPOLAMINE

FILM, EXTENDED RELEASE; TRANSDERMAL

TRANSDERM SCOP

+ NOVARTIS 1MG/72HR N17874 001

SECOBARBITAL SODIUM

CAPSULE; ORAL

SECOBARBITAL SODIUMAA EVERYLIFE 100MG N85895 001SECONAL SODIUMAA + RANBAXY 100MG N86101 002 Oct 03, 1983

+ 50MG N86101 001 Oct 03, 1983

SECRETIN SYNTHETIC HUMAN

FOR SOLUTION; INTRAVENOUS

CHIRHOSTIM

+ CHIRHOCLIN 16UGM/VIAL N21256 001 Apr 09, 2004

SECRETIN SYNTHETIC PORCINE

FOR SOLUTION; INTRAVENOUS

SECREFLO

+ CHIRHOCLIN 16UGM/VIAL N21136 001 Apr 04, 2002

SELEGILINE

FILM, EXTENDED RELEASE; TRANSDERMAL

EMSAM

SOMERSET

6MG/24HR N21336 001 Feb 27, 2006

9MG/24HR N21336 002 Feb 27, 2006

+ 12MG/24HR N21336 003 Feb 27, 2006

SELEGILINE HYDROCHLORIDE

CAPSULE; ORAL

ELDEPRYLAB + SOMERSET 5MG N20647 001 May 15, 1996

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 323 (of 370)

SELEGILINE HYDROCHLORIDE

CAPSULE; ORAL

SELEGILINE HYDROCHLORIDE

<u>AB</u>	CLONMEL HLTHCARE	<u>5MG</u>	<u>N75352</u>	<u>001</u>	Nov 30, 1998
<u>AB</u>	TORPHARM	<u>5MG</u>	<u>N75321</u>	<u>001</u>	Dec 04, 1998

TABLET; ORAL

SELEGILINE HYDROCHLORIDE

<u>AB</u>	ALPHAPHARM	<u>5MG</u>	<u>N74866</u>	<u>001</u>	Nov 26, 1997
<u>AB</u>	+ APOTEX INC	<u>5MG</u>	<u>N74871</u>	<u>001</u>	Jun 06, 1997
<u>AB</u>	CLONMEL HLTHCARE	<u>5MG</u>	<u>N74641</u>	<u>001</u>	Aug 02, 1996
<u>AB</u>	ENDO PHARMS	<u>5MG</u>	<u>N74565</u>	<u>001</u>	Aug 02, 1996
<u>AB</u>	IVAX PHARMS	<u>5MG</u>	<u>N74756</u>	<u>001</u>	Nov 25, 1998
<u>AB</u>	SIEGFRIED	<u>5MG</u>	<u>N74672</u>	<u>001</u>	Apr 01, 1997
<u>AB</u>	STASON	<u>5MG</u>	<u>N74912</u>	<u>001</u>	Apr 30, 1998
<u>AB</u>	TEVA	<u>5MG</u>	<u>N74537</u>	<u>001</u>	Aug 02, 1996
<u>AB</u>		<u>5MG</u>	<u>N74744</u>	<u>001</u>	Jan 27, 1997

TABLET, ORALLY DISINTEGRATING; ORAL

ZELAPAR

	+ VALEANT PHARM INTL	1.25MG	N21479	001	Jun 14, 2006
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SELENIUM SULFIDE

LOTION/SHAMPOO; TOPICAL

SELENIUM SULFIDE

<u>AT</u>	ACTAVIS MID ATLANTIC	<u>2.5%</u>	<u>N84394</u>	<u>001</u>	
<u>AT</u>	MORTON GROVE	<u>2.5%</u>	<u>N88228</u>	<u>001</u>	Sep 01, 1983
<u>AT</u>	PERRIGO NEW YORK	<u>2.5%</u>	<u>N89996</u>	<u>001</u>	Jan 10, 1991

SELSUN

<u>AT</u>	+ CHATTEM	<u>2.5%</u>	<u>N07936</u>	<u>001</u>	
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SERMORELIN ACETATE

INJECTABLE; INJECTION

GEREF

	+ SERONO	EQ 0.05MG BASE/AMP	N19863	001	Dec 28, 1990
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SERTACONAZOLE NITRATE

CREAM; TOPICAL

ERTACZO

	+ JOHNSON AND JOHNSON	2%	N21385	001	Dec 10, 2003
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SERTRALINE HYDROCHLORIDE

CONCENTRATE; ORAL

SERTRALINE HYDROCHLORIDE

<u>AB</u>	ROXANE	<u>EQ 20MG BASE/ML</u>	<u>N76934</u>	<u>001</u>	Jun 30, 2006
	<u>ZOLOFT</u>				
<u>AB</u>	+ PFIZER	<u>EQ 20MG BASE/ML</u>	<u>N20990</u>	<u>001</u>	Dec 07, 1999

TABLET; ORAL

SERTRALINE HYDROCHLORIDE

<u>AB</u>	IVAX PHARMS	<u>EQ 25MG BASE</u>	<u>N75719</u>	<u>003</u>	Jun 30, 2006
<u>AB</u>		<u>EQ 50MG BASE</u>	<u>N75719</u>	<u>001</u>	Jun 30, 2006
<u>AB</u>		<u>EQ 100MG BASE</u>	<u>N75719</u>	<u>002</u>	Jun 30, 2006
<u>AB</u>	TEVA	<u>EQ 25MG BASE</u>	<u>N76465</u>	<u>001</u>	Aug 11, 2006
<u>AB</u>		<u>EQ 50MG BASE</u>	<u>N76465</u>	<u>002</u>	Aug 11, 2006
<u>AB</u>		<u>EQ 100MG BASE</u>	<u>N76465</u>	<u>003</u>	Aug 11, 2006
	<u>ZOLOFT</u>				
<u>AB</u>	PFIZER	<u>EQ 25MG BASE</u>	<u>N19839</u>	<u>005</u>	Mar 06, 1996
<u>AB</u>		<u>EQ 50MG BASE</u>	<u>N19839</u>	<u>001</u>	Dec 30, 1991
<u>AB</u>	+	<u>EQ 100MG BASE</u>	<u>N19839</u>	<u>002</u>	Dec 30, 1991

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 324 (of 370)

SEVELAMER HYDROCHLORIDE

TABLET; ORAL

RENAGEL

GENZYME

400MG

N21179 001

Jul 12, 2000

+

800MG

N21179 002

Jul 12, 2000

SEVOFLURANE

LIQUID; INHALATION

SEVOFLURANEAN BAXTER HLTHCARE100%N75895 001

Jul 02, 2002

ULTANEAN + ABBOTT100%N20478 001

Jun 07, 1995

SIBUTRAMINE HYDROCHLORIDE

CAPSULE; ORAL

MERIDIA

ABBOTT

5MG

N20632 001

Nov 22, 1997

10MG

N20632 002

Nov 22, 1997

+

15MG

N20632 003

Nov 22, 1997

SILDENAFIL CITRATE

TABLET; ORAL

REVATIO

+ PFIZER

EQ 20MG BASE

N21845 001

Jun 03, 2005

VIAGRA

PFIZER IRELAND

EQ 25MG BASE

N20895 001

Mar 27, 1998

EQ 50MG BASE

N20895 002

Mar 27, 1998

+

EQ 100MG BASE

N20895 003

Mar 27, 1998

SILVER SULFADIAZINE

CREAM; TOPICAL

SILVADENEAB + KING PHARMS1%N17381 001SSDAB BASF1%N18578 001

Feb 25, 1982

SSD AF

BX BASF

1%

N18578 003

Jul 11, 1990

THERMAZENEAB KENDALL LP1%N18810 001

Dec 23, 1985

SIMVASTATIN

TABLET; ORAL

SIMVASTATINAB AUROBINDO PHARMA5MGN77691 001

Dec 20, 2006

AB 10MGN77691 002

Dec 20, 2006

AB 20MGN77691 003

Dec 20, 2006

AB 40MGN77691 004

Dec 20, 2006

AB 80MGN77691 005

Dec 20, 2006

AB COBALT 5MGN76685 001

Dec 20, 2006

AB 10MGN76685 002

Dec 20, 2006

AB 20MGN76685 003

Dec 20, 2006

AB 40MGN76685 004

Dec 20, 2006

AB 80MGN76685 005

Dec 20, 2006

AB DR REDDYS LABS INC 10MGN77752 001

Dec 20, 2006

AB 20MGN77752 002

Dec 20, 2006

AB 40MGN77752 003

Dec 20, 2006

AB 80MGN77752 004

Dec 20, 2006

AB IVAX PHARMS 5MGN76052 001

Jun 23, 2006

AB 10MGN76052 002

Jun 23, 2006

AB 20MGN76052 003

Jun 23, 2006

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 325 (of 370)

SIMVASTATIN

TABLET; ORAL

SIMVASTATIN

<u>AB</u>	IVAX PHARMS	<u>40MG</u>	<u>N76052</u>	<u>004</u>	Jun 23, 2006
<u>AB</u>		<u>80MG</u>	<u>N76052</u>	<u>005</u>	Dec 20, 2006
<u>AB</u>	PERRIGO R AND D	<u>5MG</u>	<u>N78034</u>	<u>001</u>	Dec 20, 2006
<u>AB</u>		<u>10MG</u>	<u>N78034</u>	<u>002</u>	Dec 20, 2006
<u>AB</u>		<u>20MG</u>	<u>N78034</u>	<u>003</u>	Dec 20, 2006
<u>AB</u>		<u>40MG</u>	<u>N78034</u>	<u>004</u>	Dec 20, 2006
<u>AB</u>		<u>80MG</u>	<u>N78034</u>	<u>005</u>	Dec 20, 2006
<u>AB</u>	RANBAXY	<u>5MG</u>	<u>N76285</u>	<u>001</u>	Dec 20, 2006
<u>AB</u>		<u>10MG</u>	<u>N76285</u>	<u>002</u>	Dec 20, 2006
<u>AB</u>		<u>20MG</u>	<u>N76285</u>	<u>003</u>	Dec 20, 2006
<u>AB</u>		<u>40MG</u>	<u>N76285</u>	<u>004</u>	Dec 20, 2006
<u>AB</u>		<u>80MG</u>	<u>N76285</u>	<u>005</u>	Jun 23, 2006
<u>AB</u>	SANDOZ	<u>5MG</u>	<u>N77766</u>	<u>001</u>	Dec 20, 2006
<u>AB</u>		<u>10MG</u>	<u>N77766</u>	<u>002</u>	Dec 20, 2006
<u>AB</u>		<u>20MG</u>	<u>N77766</u>	<u>003</u>	Dec 20, 2006
<u>AB</u>		<u>40MG</u>	<u>N77766</u>	<u>004</u>	Dec 20, 2006
<u>AB</u>		<u>80MG</u>	<u>N77766</u>	<u>005</u>	Dec 20, 2006
<u>AB</u>	ZYDUS PHARMS USA	<u>5MG</u>	<u>N77837</u>	<u>001</u>	Dec 20, 2006
<u>AB</u>		<u>10MG</u>	<u>N77837</u>	<u>002</u>	Dec 20, 2006
<u>AB</u>		<u>20MG</u>	<u>N77837</u>	<u>003</u>	Dec 20, 2006
<u>AB</u>		<u>40MG</u>	<u>N77837</u>	<u>004</u>	Dec 20, 2006
<u>AB</u>		<u>80MG</u>	<u>N77837</u>	<u>005</u>	Dec 20, 2006
<u>ZOCOR</u>					
<u>AB</u>	MERCK	<u>5MG</u>	<u>N19766</u>	<u>001</u>	Dec 23, 1991
<u>AB</u>		<u>10MG</u>	<u>N19766</u>	<u>002</u>	Dec 23, 1991
<u>AB</u>		<u>20MG</u>	<u>N19766</u>	<u>003</u>	Dec 23, 1991
<u>AB</u>		<u>40MG</u>	<u>N19766</u>	<u>004</u>	Dec 23, 1991
<u>AB</u>	+	<u>80MG</u>	<u>N19766</u>	<u>005</u>	Jul 10, 1998

SINCALIDE

INJECTABLE; INJECTION

KINEVAC

+ BRACCO 0.005MG/VIAL

N17697 001

SIROLIMUS

SOLUTION; ORAL

RAPAMUNE

+ WYETH PHARMS INC 1MG/ML

N21083 001 Sep 15, 1999

TABLET; ORAL

RAPAMUNE

WYETH PHARMS INC 1MG

N21110 001 Aug 25, 2000

+ 2MG

N21110 002 Aug 22, 2002

SITAGLIPTIN PHOSPHATE

TABLET; ORAL

JANUVIA

MERCK CO INC EQ 25MG BASE

N21995 001 Oct 16, 2006

EQ 50MG BASE

N21995 002 Oct 16, 2006

+ EQ 100MG BASE

N21995 003 Oct 16, 2006

SODIUM ACETATE, ANHYDROUS

INJECTABLE; INJECTION

SODIUM ACETATE IN PLASTIC CONTAINER

+ HOSPIRA 2MEQ/ML

N18893 001 May 04, 1983

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 326 (of 370)

SODIUM BENZOATE; SODIUM PHENYLACETATE

SOLUTION; IV (INFUSION)

+	UCYCLYD	10%;10% (5GM/50ML;5GM/50ML)	N20645	001	Feb 17, 2005
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SODIUM BICARBONATE

INJECTABLE; INJECTION

SODIUM BICARBONATE

	HOSPIRA	0.9MEQ/ML	N77394	001	Nov 09, 2005
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+		1MEQ/ML	N77394	002	Nov 09, 2005
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SODIUM BICARBONATE; TARTARIC ACID

GRANULE, EFFERVESCENT; ORAL

BAROS

	LAFAYETTE PHARMS	460MG/GM;420MG/GM	N18509	001	Aug 07, 1985
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SODIUM CHLORIDE

INJECTABLE; INJECTION

BACTERIOSTATIC SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER

<u>AP</u>	ABRAXIS PHARM	<u>9MG/ML</u>	<u>N88911</u>	<u>001</u>	Feb 07, 1985
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<u>AP</u>	+ HOSPIRA	<u>9MG/ML</u>	<u>N18800</u>	<u>001</u>	Oct 29, 1982
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SODIUM CHLORIDE 0.45% IN PLASTIC CONTAINER

<u>AP</u>	B BRAUN	<u>450MG/100ML</u>	<u>N19635</u>	<u>001</u>	Mar 09, 1988
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<u>AP</u>	BAXTER HLTHCARE	<u>450MG/100ML</u>	<u>N18016</u>	<u>001</u>	
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<u>AP</u>	HOSPIRA	<u>450MG/100ML</u>	<u>N18090</u>	<u>001</u>	
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<u>AP</u>		<u>450MG/100ML</u>	<u>N19759</u>	<u>001</u>	Jun 08, 1988
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SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER

<u>AP</u>	+ ABRAXIS PHARM	<u>9MG/ML</u>	<u>N88912</u>	<u>001</u>	Jan 10, 1985
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<u>AP</u>	+ B BRAUN	<u>900MG/100ML</u>	<u>N17464</u>	<u>001</u>	
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<u>AP</u>	+	<u>900MG/100ML</u>	<u>N19635</u>	<u>002</u>	Mar 09, 1988
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<u>AP</u>	+ BAXTER HLTHCARE	<u>9MG/ML</u>	<u>N16677</u>	<u>004</u>	Oct 30, 1985
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<u>AP</u>		<u>9MG/ML</u>	<u>N20178</u>	<u>002</u>	Dec 07, 1992
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<u>AP</u>	+	<u>900MG/100ML</u>	<u>N16677</u>	<u>001</u>	
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<u>AP</u>	+	<u>900MG/100ML</u>	<u>N20178</u>	<u>001</u>	Dec 07, 1992
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<u>AP</u>	HAEMONETICS	<u>900MG/100ML</u>	<u>N76316</u>	<u>001</u>	Oct 27, 2004
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<u>AP</u>	+ HOSPIRA	<u>9MG/ML</u>	<u>N18803</u>	<u>001</u>	Oct 29, 1982
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<u>AP</u>	+	<u>9MG/ML</u>	<u>N19217</u>	<u>001</u>	Jul 13, 1984
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<u>AP</u>		<u>9MG/ML</u>	<u>N19465</u>	<u>002</u>	Jul 15, 1985
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<u>AP</u>	+	<u>900MG/100ML</u>	<u>N16366</u>	<u>001</u>	
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<u>AP</u>	+	<u>900MG/100ML</u>	<u>N19465</u>	<u>001</u>	Jul 15, 1985
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<u>AP</u>	+	<u>900MG/100ML</u>	<u>N19480</u>	<u>001</u>	Sep 17, 1985
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	+ MALLINCKRODT	45MG/50ML (9MG/ML)	N21569	001	Jul 27, 2006
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		112.5MG/125ML (9MG/ML)	N21569	002	Jul 27, 2006
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<u>AP</u>	+ TARO PHARMS IRELAND	<u>9MG/ML</u>	<u>N77407</u>	<u>001</u>	Aug 11, 2006
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SODIUM CHLORIDE 3% IN PLASTIC CONTAINER

	BAXTER HLTHCARE	3GM/100ML	N19022	001	Nov 01, 1983
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SODIUM CHLORIDE 5% IN PLASTIC CONTAINER

	BAXTER HLTHCARE	5GM/100ML	N19022	002	Nov 01, 1983
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SODIUM CHLORIDE IN PLASTIC CONTAINER

	HOSPIRA	2.5MEQ/ML	N18897	001	Jul 20, 1984
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SOLUTION FOR SLUSH; IRRIGATION

SODIUM CHLORIDE 0.9% IN STERILE PLASTIC CONTAINER

	BAXTER HLTHCARE	900MG/100ML	N19319	002	May 17, 1985
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SOLUTION; IRRIGATION

SODIUM CHLORIDE 0.45% IN PLASTIC CONTAINER

<u>AT</u>	BAXTER HLTHCARE	<u>450MG/100ML</u>	<u>N17864</u>	<u>001</u>	
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<u>AT</u>	HOSPIRA	<u>450MG/100ML</u>	<u>N17670</u>	<u>001</u>	
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<u>AT</u>		<u>450MG/100ML</u>	<u>N18380</u>	<u>001</u>	
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SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER

<u>AT</u>	B BRAUN	<u>900MG/100ML</u>	<u>N16733</u>	<u>001</u>	
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27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 327 (of 370)

SODIUM CHLORIDE

SOLUTION; IRRIGATION

SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER

<u>AT</u>	BAXTER HLTHCARE	<u>900MG/100ML</u>	<u>N17427</u>	<u>001</u>	
<u>AT</u>		<u>900MG/100ML</u>	<u>N17867</u>	<u>001</u>	
<u>AT</u>	HOSPIRA	<u>900MG/100ML</u>	<u>N17514</u>	<u>001</u>	
<u>AT</u>		<u>900MG/100ML</u>	<u>N18314</u>	<u>001</u>	

SODIUM CHROMATE, CR-51INJECTABLE; INJECTION
CHROMITOPES SODIUM

BRACCO 200uCi/ML N13993 001

SODIUM CHROMATE CR 51
MALLINCKRODT 100uCi/ML N16708 001SODIUM FERRIC GLUCONATE COMPLEXINJECTABLE; INJECTION
FERRLECIT

+ WATSON PHARMS 62.5MG/5ML N20955 001 Feb 18, 1999

SODIUM IODIDE, I-123

CAPSULE; ORAL

SODIUM IODIDE I 123

<u>AA</u>	+	GE HEALTHCARE	<u>100uCi</u>	<u>N17630</u>	<u>001</u>	
<u>AA</u>		MALLINCKRODT	<u>100uCi</u>	<u>N71909</u>	<u>001</u>	Feb 28, 1989
<u>AA</u>			<u>200uCi</u>	<u>N71910</u>	<u>001</u>	Feb 28, 1989
<u>AA</u>	+	SYNCOR PHARMS	<u>100uCi</u>	<u>N18671</u>	<u>001</u>	May 27, 1982
<u>AA</u>	+		<u>200uCi</u>	<u>N18671</u>	<u>002</u>	May 27, 1982

SOLUTION; ORAL

SODIUM IODIDE I 123

+ GE HEALTHCARE 2mCi/ML N17630 002

SODIUM IODIDE, I-131

CAPSULE; ORAL

IODOTOPE

+ BRACCO 1-130mCi N10929 001

SODIUM IODIDE I 131

DRAXIMAGE 100uCi N21305 004 Nov 18, 2004

+ MALLINCKRODT 15-100uCi N16517 002

+ 0.8-100mCi N16517 001

SOLUTION; ORAL

SODIUM IODIDE I 131

+ MALLINCKRODT 3.5-150mCi/VIAL N16515 001

SODIUM IODIDE I 131, KIT

+ DRAXIMAGE 1-500mCi/0.5ML N21305 003 Jan 24, 2003

+ 1-250mCi/0.25ML N21305 002 Jan 24, 2003

SODIUM LACTATE

INJECTABLE; INJECTION

SODIUM LACTATE 0.167 MOLAR IN PLASTIC CONTAINER

<u>AP</u>	BAXTER HLTHCARE	<u>1.87GM/100ML</u>	<u>N16692</u>	<u>001</u>	
<u>AP</u>	HOSPIRA	<u>1.87GM/100ML</u>	<u>N18249</u>	<u>001</u>	

SODIUM LACTATE 1/6 MOLAR IN PLASTIC CONTAINER

<u>AP</u>	B BRAUN	<u>1.87GM/100ML</u>	<u>N20004</u>	<u>001</u>	Apr 21, 1992
	SODIUM LACTATE IN PLASTIC CONTAINER				
	HOSPIRA	5MEQ/ML	N18947	001	Sep 05, 1984

PRESCRIPTION DRUG PRODUCT LIST

3 - 328 (of 370)

SODIUM NITROPRUSSIDE

INJECTABLE; INJECTION

NITROPRESS

<u>AP</u>	+	HOSPIRA	<u>25MG/ML</u>	<u>N71961</u>	<u>001</u>	Aug 01, 1988
	+		50MG/VIAL	N70566	001	Jun 09, 1986

SODIUM NITROPRUSSIDE

<u>AP</u>		SICOR PHARMS	<u>25MG/ML</u>	<u>N73465</u>	<u>001</u>	Mar 30, 1992
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SODIUM OXYBATE

SOLUTION; ORAL

XYREM

	+	JAZZ	500MG/ML	N21196	001	Jul 17, 2002
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SODIUM PHENYLBUTYRATE

POWDER; ORAL

BUPHENYL

	+	MEDICIS	3GM/TEASPOONFUL	N20573	001	Apr 30, 1996
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TABLET; ORAL

BUPHENYL

	+	MEDICIS	500MG	N20572	001	May 13, 1996
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SODIUM PHOSPHATE, DIBASIC, ANHYDROUS; SODIUM PHOSPHATE, MONOBASIC, MONOHYDRATE

TABLET; ORAL

OSMOPREP

	+	SALIX PHARMS	0.398GM;1.102GM	N21892	001	Mar 16, 2006
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VISICOL

	+	SALIX PHARMS	0.398GM;1.102GM	N21097	001	Sep 21, 2000
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SODIUM PHOSPHATE, DIBASIC, HEPTAHYDRATE; SODIUM PHOSPHATE, MONOBASIC, ANHYDROUS

INJECTABLE; INJECTION

SODIUM PHOSPHATES IN PLASTIC CONTAINER

		HOSPIRA	142MG/ML;276MG/ML	N18892	001	May 10, 1983
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SODIUM PHOSPHATE, P-32

SOLUTION; INJECTION, ORAL

SODIUM PHOSPHATE P 32

		MALLINCKRODT	0.67mCi/ML	N11777	001	
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SODIUM POLYSTYRENE SULFONATE

POWDER; ORAL, RECTAL

KAYEXALATE

<u>AA</u>	+	SANOFI AVENTIS US	<u>453.6GM/BOT</u>	<u>N11287</u>	<u>001</u>	
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KIONEX

<u>AA</u>		PADDOCK	<u>454GM/BOT</u>	<u>N40029</u>	<u>001</u>	Feb 06, 1998
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SODIUM POLYSTYRENE SULFONATE

<u>AA</u>		CAROLINA MEDCL	<u>454GM/BOT</u>	<u>N89910</u>	<u>001</u>	Jan 19, 1989
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SUSPENSION; ORAL, RECTAL

SODIUM POLYSTYRENE SULFONATE

<u>AA</u>		ROXANE	<u>15GM/60ML</u>	<u>N89049</u>	<u>001</u>	Nov 17, 1986
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SPS

<u>AA</u>	+	CAROLINA MEDCL	<u>15GM/60ML</u>	<u>N87859</u>	<u>001</u>	Dec 08, 1982
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SODIUM TETRADECYL SULFATE

INJECTABLE; INJECTION

SOTRADECOL

		BIONICHE PHARM USA	20MG/2ML (10MG/ML)	N40541	001	Nov 12, 2004
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	+		60MG/2ML (30MG/ML)	N40541	002	Nov 12, 2004
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PRESCRIPTION DRUG PRODUCT LIST

3 - 329 (of 370)

SOLIFENACIN SUCCINATE

TABLET; ORAL

VESICARE

ASTELLAS

5MG

N21518 001

Nov 19, 2004

+

10MG

N21518 002

Nov 19, 2004

SOMATROPIN RECOMBINANT

INJECTABLE; INJECTION

GENOTROPIN

BX + PHARMACIA AND UPJOHN 5.8MG/VIAL
+ 13.8MG/VIAL

N20280 006

Aug 24, 1995

N20280 007

Oct 23, 1996

GENOTROPIN PRESERVATIVE FREE

BX PHARMACIA AND UPJOHN 1.5MG/VIAL
0.2MG/VIAL
0.4MG/VIAL
0.6MG/VIAL
0.8MG/VIAL
1MG/VIAL
1.2MG/VIAL
1.4MG/VIAL
1.6MG/VIAL
1.8MG/VIAL
+ 2MG/VIAL

N20280 004

Aug 24, 1995

N20280 001

Jan 27, 1998

N20280 002

Jan 27, 1998

N20280 003

Jan 27, 1998

N20280 005

Jan 27, 1998

N20280 008

Jan 27, 1998

N20280 009

Jan 27, 1998

N20280 010

Jan 27, 1998

N20280 011

Jan 27, 1998

N20280 012

Jan 27, 1998

N20280 013

Jan 27, 1998

HUMATROPE

BX + LILLY 5MG/VIAL
BX 6MG/VIAL
+ 12MG/VIAL
+ 24MG/VIAL

N19640 004

Mar 08, 1987

N19640 005

Feb 04, 1999

N19640 006

Feb 04, 1999

N19640 007

Feb 04, 1999

NORDITROPIN

BX NOVO NORDISK INC 4MG/VIAL
5MG/1.5ML
8MG/VIAL
10MG/1.5ML
+ 15MG/1.5ML

N19721 001

May 08, 1995

N21148 001

Jun 20, 2000

N19721 002

May 08, 1995

N21148 002

Jun 20, 2000

N21148 003

Jun 20, 2000

NORDITROPIN NORDIFLEX

NOVO NORDISK INC

5MG/1.5ML
10MG/1.5ML
15MG/1.5ML

N21148 004

Oct 01, 2004

N21148 005

Oct 01, 2004

N21148 006

Oct 01, 2004

NUTROPIN

BX GENENTECH 5MG/VIAL
+ 10MG/VIAL

N20168 001

Nov 17, 1993

N20168 002

Nov 17, 1993

NUTROPIN AQ

+ GENENTECH 5MG/ML

N20522 001

Dec 29, 1995

NUTROPIN AQ PEN

BX + GENENTECH 5MG/ML

N20522 002

Apr 22, 2002

OMNITROPE

BX SANDOZ 1.5MG/VIAL
BX 5.8MG/VIAL

N21426 002

May 30, 2006

N21426 001

May 30, 2006

SAIZEN

BX SERONO 5MG/VIAL
+ 8.8MG/VIAL

N19764 002

Oct 08, 1996

N19764 003

Aug 29, 2000

SEROSTIM

BX SERONO 4MG/VIAL
BX 5MG/VIAL
BX 6MG/VIAL

N20604 003

Jul 25, 1997

N20604 002

Aug 23, 1996

N20604 001

Aug 23, 1996

TEV-TROPIN

BX + FERRING 5MG/ML

N19774 002

Jan 04, 2002

ZORBTIVE

+ SERONO INC 8.8MG/VIAL

N21597 004

Dec 01, 2003

PRESCRIPTION DRUG PRODUCT LIST

3 - 330 (of 370)

SORAFENIB TOSYLATE

TABLET; ORAL

NEXAVAR

+ BAYER PHARMS

EQ 200MG BASE

N21923 001

Dec 20, 2005

SORBITOL

SOLUTION; IRRIGATION

SORBITOL 3% IN PLASTIC CONTAINER

BAXTER HLTHCARE 3GM/100ML

N17863 001

SORBITOL 3.3% IN PLASTIC CONTAINER

B BRAUN 3.3GM/100ML

N16741 001

SOTALOL HYDROCHLORIDE

TABLET; ORAL

BETAPACE

AB1 BERLEX

80MG

N19865 001

Oct 30, 1992

AB1

120MG

N19865 005

Apr 20, 1994

AB1 +

160MG

N19865 002

Oct 30, 1992

AB1

240MG

N19865 003

Oct 30, 1992

BETAPACE AF

AB2 BERLEX

80MG

N21151 001

Feb 22, 2000

AB2

120MG

N21151 002

Feb 22, 2000

AB2 +

160MG

N21151 003

Feb 22, 2000

SORINE

AB1 UPSHER SMITH

80MG

N75500 001

Apr 27, 2001

AB1

120MG

N75500 004

Apr 27, 2001

AB1

160MG

N75500 002

Apr 27, 2001

AB1

240MG

N75500 003

Apr 27, 2001

SOTALOL HYDROCHLORIDE

AB1 APOTEX INC

80MG

N76140 001

Sep 26, 2002

AB1

120MG

N76140 002

Sep 26, 2002

AB1

160MG

N76140 003

Sep 26, 2002

AB1

240MG

N76140 004

Sep 26, 2002

AB1 GENPHARM

80MG

N75237 001

May 01, 2000

AB1

120MG

N75237 002

May 01, 2000

AB1

160MG

N75237 003

May 01, 2000

AB1

240MG

N75237 004

May 01, 2000

AB2

80MG

N77070 001

Nov 04, 2005

AB2

120MG

N77070 002

Nov 04, 2005

AB2

160MG

N77070 003

Nov 04, 2005

AB1 IMPAX PHARMS

80MG

N75663 001

Nov 07, 2000

AB1

120MG

N75663 002

Nov 07, 2000

AB1

160MG

N75663 003

Nov 07, 2000

AB1

240MG

N75663 004

Nov 07, 2000

AB1 MUTUAL PHARM

80MG

N75515 001

Oct 15, 2001

AB1

120MG

N75515 004

Oct 15, 2001

AB1

160MG

N75515 002

Oct 15, 2001

AB1

240MG

N75515 003

Oct 15, 2001

AB2

80MG

N76576 001

Apr 08, 2004

AB2

120MG

N76576 002

Apr 08, 2004

AB2

160MG

N76576 003

Apr 08, 2004

AB1 MYLAN

80MG

N75725 001

Dec 19, 2000

AB1

120MG

N75725 002

Dec 19, 2000

AB1

160MG

N75725 003

Dec 19, 2000

AB1

240MG

N75725 004

Dec 19, 2000

AB1 SANDOZ

80MG

N75366 001

May 01, 2000

AB1

120MG

N75366 002

May 01, 2000

AB1

160MG

N75366 003

May 01, 2000

AB1

240MG

N75366 004

May 01, 2000

AB1 TEVA

80MG

N75429 001

May 01, 2000

PRESCRIPTION DRUG PRODUCT LIST

3 - 331 (of 370)

SOTALOL HYDROCHLORIDE

TABLET; ORAL

SOTALOL HYDROCHLORIDE

<u>AB1</u>	TEVA	<u>120MG</u>	<u>N75429</u>	<u>002</u>	May 01, 2000
<u>AB1</u>		<u>160MG</u>	<u>N75429</u>	<u>003</u>	May 01, 2000
<u>AB1</u>		<u>240MG</u>	<u>N75429</u>	<u>004</u>	May 01, 2000
<u>AB2</u>		<u>80MG</u>	<u>N76883</u>	<u>001</u>	Jul 26, 2004
<u>AB2</u>		<u>120MG</u>	<u>N76883</u>	<u>002</u>	Jul 26, 2004
<u>AB2</u>		<u>160MG</u>	<u>N76883</u>	<u>003</u>	Jul 26, 2004
<u>AB2</u>	TORPHARM	<u>80MG</u>	<u>N76214</u>	<u>001</u>	Aug 27, 2003
<u>AB2</u>		<u>120MG</u>	<u>N76214</u>	<u>002</u>	Aug 27, 2003
<u>AB2</u>		<u>160MG</u>	<u>N76214</u>	<u>003</u>	Aug 27, 2003
<u>AB1</u>	VINTAGE PHARMS	<u>80MG</u>	<u>N75563</u>	<u>001</u>	Nov 07, 2003
<u>AB1</u>		<u>120MG</u>	<u>N75563</u>	<u>002</u>	Nov 07, 2003
<u>AB1</u>		<u>160MG</u>	<u>N75563</u>	<u>003</u>	Nov 07, 2003
<u>AB1</u>		<u>240MG</u>	<u>N75563</u>	<u>004</u>	Nov 07, 2003
<u>AB1</u>	WATSON LABS	<u>80MG</u>	<u>N75238</u>	<u>001</u>	Jul 13, 2000
<u>AB1</u>		<u>120MG</u>	<u>N75238</u>	<u>002</u>	Jul 13, 2000
<u>AB1</u>		<u>160MG</u>	<u>N75238</u>	<u>003</u>	Jul 13, 2000
<u>AB1</u>		<u>240MG</u>	<u>N75238</u>	<u>004</u>	Jul 13, 2000

SOYBEAN OIL

INJECTABLE; INJECTION

INTRALIPID 10%

<u>AP</u>	+ FRESINIUS	<u>10%</u>	<u>N17643</u>	<u>001</u>	
	<u>INTRALIPID 20%</u>				
<u>AP</u>	+ FRESINIUS	<u>20%</u>	<u>N18449</u>	<u>001</u>	
<u>AP</u>	+	<u>20%</u>	<u>N20248</u>	<u>001</u>	Aug 07, 1996
	<u>INTRALIPID 30%</u>				
<u>AP</u>	+ FRESINIUS	<u>30%</u>	<u>N19942</u>	<u>001</u>	Dec 30, 1993
	<u>LIPOSYN III 10%</u>				
<u>AP</u>	+ HOSPIRA	<u>10%</u>	<u>N18969</u>	<u>001</u>	Sep 24, 1984
	<u>LIPOSYN III 20%</u>				
<u>AP</u>	+ HOSPIRA	<u>20%</u>	<u>N18970</u>	<u>001</u>	Sep 25, 1984
	<u>LIPOSYN III 30%</u>				
<u>AP</u>	+ HOSPIRA	<u>30%</u>	<u>N20181</u>	<u>001</u>	Jan 13, 1998
	<u>NUTRILIPID 10%</u>				
<u>AP</u>	+ B BRAUN	<u>10%</u>	<u>N19531</u>	<u>001</u>	May 28, 1993
	<u>NUTRILIPID 20%</u>				
<u>AP</u>	+ B BRAUN	<u>20%</u>	<u>N19531</u>	<u>002</u>	May 28, 1993

SPECTINOMYCIN HYDROCHLORIDE

INJECTABLE; INJECTION

TROBICIN

+ PHARMACIA AND UPJOHN EQ 2GM BASE/VIAL

N50347 001

SPIRONOLACTONE

TABLET; ORAL

ALDACTONE

<u>AB</u>	GD SEARLE LLC	<u>25MG</u>	<u>N12151</u>	<u>009</u>	Dec 30, 1983
<u>AB</u>		<u>50MG</u>	<u>N12151</u>	<u>008</u>	Dec 30, 1982
<u>AB</u>	+	<u>100MG</u>	<u>N12151</u>	<u>010</u>	Dec 30, 1983
	<u>SPIRONOLACTONE</u>				
<u>AB</u>	ACTAVIS ELIZABETH	<u>25MG</u>	<u>N40353</u>	<u>003</u>	Mar 15, 2006
<u>AB</u>		<u>50MG</u>	<u>N40353</u>	<u>001</u>	Jul 29, 1999
<u>AB</u>		<u>100MG</u>	<u>N40353</u>	<u>002</u>	Jul 29, 1999
<u>AB</u>	MUTUAL PHARM	<u>25MG</u>	<u>N87265</u>	<u>001</u>	
<u>AB</u>		<u>25MG</u>	<u>N89424</u>	<u>001</u>	Jul 23, 1986
<u>AB</u>		<u>50MG</u>	<u>N89424</u>	<u>002</u>	Aug 11, 1999
<u>AB</u>		<u>100MG</u>	<u>N89424</u>	<u>003</u>	Aug 11, 1999

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 332 (of 370)

SPIRONOLACTONE

TABLET; ORAL

SPIRONOLACTONE

<u>AB</u>	MYLAN	<u>25MG</u>	<u>N40424</u>	<u>001</u>	Aug 20, 2001
<u>AB</u>		<u>25MG</u>	<u>N87086</u>	<u>001</u>	
<u>AB</u>		<u>50MG</u>	<u>N40424</u>	<u>002</u>	Aug 20, 2001
<u>AB</u>		<u>100MG</u>	<u>N40424</u>	<u>003</u>	Aug 20, 2001
<u>AB</u>	SANDOZ	<u>25MG</u>	<u>N86809</u>	<u>001</u>	
<u>AB</u>	VINTAGE	<u>25MG</u>	<u>N40750</u>	<u>001</u>	Aug 29, 2006
<u>AB</u>		<u>50MG</u>	<u>N40750</u>	<u>002</u>	Aug 29, 2006
<u>AB</u>		<u>100MG</u>	<u>N40750</u>	<u>003</u>	Aug 29, 2006

STAVUDINE

CAPSULE; ORAL

ZERIT

	BRISTOL MYERS SQUIBB	15MG	N20412	002	Jun 24, 1994
		20MG	N20412	003	Jun 24, 1994
		30MG	N20412	004	Jun 24, 1994
+		40MG	N20412	005	Jun 24, 1994

FOR SOLUTION; ORAL

ZERIT

+	BRISTOL MYERS SQUIBB	1MG/ML	N20413	001	Sep 06, 1996
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STREPTOMYCIN SULFATE

INJECTABLE; INJECTION

STREPTOMYCIN SULFATE

+	PFIZER	EQ 1GM BASE/2.5ML	N60111	001	
+	X GEN PHARMS	EQ 1GM BASE/VIAL	N64210	001	Jun 30, 1998

STREPTOZOCIN

INJECTABLE; INJECTION

ZANOSAR

+	SICOR PHARMS	1GM/VIAL	N50577	001	May 07, 1982
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STRONTIUM CHLORIDE, SR-89

INJECTABLE; INJECTION

METASTRON

<u>AP</u>	+	GE HEALTHCARE	<u>1mCi/ML</u>	<u>N20134</u>	<u>001</u>	Jun 18, 1993
		<u>STRONTIUM CHLORIDE SR-89</u>				
<u>AP</u>		BIO NUCLEONICS	<u>1mCi/ML</u>	<u>N75941</u>	<u>001</u>	Jan 06, 2003

SUCCIMER

CAPSULE; ORAL

CHEMET

+	OVATION PHARMS	100MG	N19998	002	Jan 30, 1991
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SUCCINYLCHOLINE CHLORIDE

INJECTABLE; INJECTION

ANECTINE

<u>AP</u>	+	SANDOZ	<u>20MG/ML</u>	<u>N08453</u>	<u>002</u>	
		<u>QUELICIN</u>				
<u>AP</u>	+	HOSPIRA	<u>20MG/ML</u>	<u>N08845</u>	<u>006</u>	
		<u>QUELICIN PRESERVATIVE FREE</u>				
<u>AP</u>	+	HOSPIRA	<u>20MG/ML</u>	<u>N08845</u>	<u>001</u>	
	+		100MG/ML	N08845	004	

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 333 (of 370)

SUCRALFATE

SUSPENSION; ORAL

CARAFATE

+ AXCAN SCANDIPHARM 1GM/10ML N19183 001 Dec 16, 1993

TABLET; ORAL

CARAFATEAB + AXCAN SCANDIPHARM 1GM N18333 001SUCRALFATEAB RATIOPHARM 1GM N74415 001 Jun 08, 1998AB TEVA 1GM N70848 001 Mar 29, 1996SUFENTANIL CITRATE

INJECTABLE; INJECTION

SUFENTAAP + AKORN EQ 0.05MG BASE/ML N19050 001 May 04, 1984SUFENTANIL CITRATEAP BAXTER HLTHCARE EQ 0.05MG BASE/ML N74413 001 Dec 15, 1995SUFENTANIL CITRATEAP HOSPIRA EQ 0.05MG BASE/ML N74534 001 Dec 11, 1996SULCONAZOLE NITRATE

CREAM; TOPICAL

EXELDERM

+ WESTWOOD SQUIBB 1% N18737 001 Feb 28, 1989

SOLUTION; TOPICAL

EXELDERM

+ WESTWOOD SQUIBB 1% N18738 001 Aug 30, 1985

SULFACETAMIDE SODIUM

LOTION; TOPICAL

KLARONAB + SANOFI AVENTIS US 10% N19931 001 Dec 23, 1996SULFACETAMIDE SODIUMAB ALTANA 10% N77015 001 Nov 17, 2006

OINTMENT; OPHTHALMIC

CETAMIDEAT + ALCON 10% N80021 001SULFACETAMIDE SODIUMAT ALTANA 10% N80029 001

SOLUTION/DROPS; OPHTHALMIC

BLEPH-10AT + ALLERGAN 10% N80028 001BLEPH-30AT + ALLERGAN 30% N80028 002ISOPTO CETAMIDEAT + ALCON 15% N80020 002OCUSULF-10AT MIZA PHARMS USA 10% N80660 001SULF-10AT NOVARTIS 10% N80025 001SULFACEL-15AT OPTOPICS 15% N80024 001SULFACETAMIDE SODIUMAT AKORN 10% N40215 001 May 25, 1999AT 30% N40216 001 May 25, 1999AT ALCON 30% N89068 001 May 05, 1987AT ALCON UNIVERSAL 10% N89560 001 Oct 18, 1988AT BAUSCH AND LOMB 10% N40066 001 Dec 28, 1994

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 334 (of 370)

SULFADIAZINETABLET; ORAL
SULFADIAZINE

+ SANDOZ 500MG N40091 001 Jul 29, 1994

SULFAMETHOXAZOLE; TRIMETHOPRIM

INJECTABLE; INJECTION

BACTRIM

AP + MUTUAL PHARM 80MG/ML;16MG/ML N18374 001

SULFAMETHOXAZOLE AND TRIMETHOPRIM

AP BAXTER HLTHCARE 80MG/ML;16MG/ML N70628 001 Dec 29, 1987

AP HOSPIRA 80MG/ML;16MG/ML N73199 001 Sep 11, 1992

AP SICOR PHARMS 80MG/ML;16MG/ML N73303 001 Oct 31, 1991

SUSPENSION; ORAL

BACTRIM PEDIATRIC

AB MUTUAL PHARM 200MG/5ML;40MG/5ML N17560 002

SEPTRA

AB + MONARCH PHARMS 200MG/5ML;40MG/5ML N17598 001

SEPTRA GRAPE

AB MONARCH PHARMS 200MG/5ML;40MG/5ML N17598 002 Feb 12, 1986

SULFAMETHOXAZOLE AND TRIMETHOPRIM

AB HI TECH PHARMA 200MG/5ML;40MG/5ML N74650 001 Dec 29, 1997

AB TEVA 200MG/5ML;40MG/5ML N18812 001 Jan 28, 1983

AB 200MG/5ML;40MG/5ML N18812 002 Jun 10, 1983

AB TEVA PHARMS 200MG/5ML;40MG/5ML N77612 001 Nov 13, 2006

SULFATRIM

AB ACTAVIS MID ATLANTIC 200MG/5ML;40MG/5ML N18615 002 Jan 07, 1983

SULFATRIM PEDIATRIC

AB ACTAVIS MID ATLANTIC 200MG/5ML;40MG/5ML N18615 001 Jan 07, 1983

TABLET; ORAL

BACTRIM

AB MUTUAL PHARM 400MG;80MG N17377 001

BACTRIM DS

AB + MUTUAL PHARM 800MG;160MG N17377 002

COTRIM

AB TEVA 400MG;80MG N70034 001 May 16, 1985

COTRIM D.S.

AB TEVA 800MG;160MG N70048 001 Mar 18, 1985

SEPTRA

AB MONARCH PHARMS 400MG;80MG N17376 001

SEPTRA DS

AB MONARCH PHARMS 800MG;160MG N17376 002

SULFAMETHOPRIM

AB PAR PHARM 400MG;80MG N70022 001 Feb 15, 1985

SULFAMETHOXAZOLE AND TRIMETHOPRIM

AB INTERPHARM 400MG;80MG N76899 001 Jan 27, 2005

AB 800MG;160MG N76899 002 Jan 27, 2005

AB MUTUAL PHARM 400MG;80MG N71016 001 Aug 25, 1986

AB 800MG;160MG N71017 001 Aug 25, 1986

AB PLIVA 400MG;80MG N70215 001 Sep 10, 1985

AB 800MG;160MG N70216 001 Sep 10, 1985

AB SANDOZ 800MG;160MG N70890 001 Nov 13, 1986

AB VISTA PHARMS 400MG;80MG N76817 001 Oct 07, 2005

AB 800MG;160MG N76817 002 Oct 07, 2005

AB WATSON LABS 400MG;80MG N18852 001 May 09, 1983

SULFAMETHOXAZOLE AND TRIMETHOPRIM DOUBLE STRENGTH

AB SANDOZ 800MG;160MG N18598 004 May 19, 1982

AB TEVA 800MG;160MG N70037 001 Jun 02, 1987

AB WATSON LABS 800MG;160MG N18854 001 May 09, 1983

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 335 (of 370)

SULFAMETHOXAZOLE; TRIMETHOPRIM

TABLET; ORAL

SULFAMETHOXAZOLE AND TRIMETHOPRIM SINGLE STRENGTH

<u>AB</u>	PLANTEX	<u>400MG;80MG</u>	<u>N70030</u>	<u>001</u>	Jun 02, 1987
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SULFANILAMIDE

CREAM; VAGINAL

AVC

+	PHARMELLE	15%	N06530	003	Jan 27, 1987
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SULFASALAZINE

SUSPENSION; ORAL

AZULFIDINE

+	PHARMACIA AND UPJOHN	250MG/5ML	N86983	001	
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TABLET; ORAL

AZULFIDINE

<u>AB</u>	+	PHARMACIA AND UPJOHN	<u>500MG</u>	<u>N07073</u>	<u>001</u>	
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SULFASALAZINE

<u>AB</u>	MUTUAL PHARM	<u>500MG</u>	<u>N89590</u>	<u>001</u>	Oct 19, 1987
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<u>AB</u>	VINTAGE PHARMS	<u>500MG</u>	<u>N40349</u>	<u>001</u>	Jan 11, 2002
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<u>AB</u>	WATSON LABS	<u>500MG</u>	<u>N85828</u>	<u>001</u>	
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<u>AB</u>		<u>500MG</u>	<u>N87197</u>	<u>001</u>	
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TABLET, DELAYED RELEASE; ORAL

AZULFIDINE EN-TABS

<u>AB</u>	+	PHARMACIA AND UPJOHN	<u>500MG</u>	<u>N07073</u>	<u>002</u>	Apr 06, 1983
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SULFASALAZINE

<u>AB</u>	VINTAGE PHARMS	<u>500MG</u>	<u>N75339</u>	<u>001</u>	Jan 11, 2002
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SULFINPYRAZONE

CAPSULE; ORAL

ANTURANE

<u>AB</u>	+	NOVARTIS	<u>200MG</u>	<u>N11556</u>	<u>004</u>	
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SULFINPYRAZONE

<u>AB</u>	BARR	<u>200MG</u>	<u>N87666</u>	<u>001</u>	Sep 17, 1982
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<u>AB</u>	IVAX PHARMS	<u>200MG</u>	<u>N87770</u>	<u>001</u>	Nov 19, 1982
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<u>AB</u>	PAR PHARM	<u>200MG</u>	<u>N88934</u>	<u>001</u>	Sep 06, 1985
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TABLET; ORAL

ANTURANE

<u>AB</u>	+	NOVARTIS	<u>100MG</u>	<u>N11556</u>	<u>003</u>	
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SULFINPYRAZONE

<u>AB</u>	BARR	<u>100MG</u>	<u>N87665</u>	<u>001</u>	Sep 17, 1982
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<u>AB</u>	PAR PHARM	<u>100MG</u>	<u>N88933</u>	<u>001</u>	Sep 06, 1985
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SULFISOXAZOLE

TABLET; ORAL

SULFISOXAZOLE

+	IVAX PHARMS	500MG	N80142	001	
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SULFISOXAZOLE ACETYL

SUSPENSION; ORAL

GANTRISIN PEDIATRIC

+	ROCHE	EQ 500MG BASE/5ML	N09182	004	
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SULINDAC

TABLET; ORAL

CLINORIL

<u>AB</u>	+	MERCK	<u>200MG</u>	<u>N17911</u>	<u>002</u>	
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SULINDAC

<u>AB</u>	MUTUAL PHARM	<u>150MG</u>	<u>N72050</u>	<u>001</u>	Apr 17, 1991
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PRESCRIPTION DRUG PRODUCT LIST

3 - 336 (of 370)

SULINDAC

TABLET; ORAL

SULINDAC

<u>AB</u>	MUTUAL PHARM	<u>200MG</u>	<u>N72051</u>	<u>001</u>	Apr 17, 1991
<u>AB</u>	MYLAN	<u>150MG</u>	<u>N73039</u>	<u>002</u>	Jun 22, 1993
<u>AB</u>		<u>200MG</u>	<u>N73039</u>	<u>001</u>	Jun 22, 1993
<u>AB</u>	SANDOZ	<u>150MG</u>	<u>N72712</u>	<u>001</u>	Aug 30, 1991
<u>AB</u>		<u>200MG</u>	<u>N72713</u>	<u>001</u>	Aug 30, 1991
<u>AB</u>	TEVA	<u>150MG</u>	<u>N72972</u>	<u>001</u>	Feb 28, 1992
<u>AB</u>		<u>200MG</u>	<u>N72973</u>	<u>001</u>	Feb 28, 1992
<u>AB</u>	WATSON LABS	<u>150MG</u>	<u>N71891</u>	<u>001</u>	Apr 03, 1990
<u>AB</u>		<u>200MG</u>	<u>N71795</u>	<u>001</u>	Apr 03, 1990

SUMATRIPTAN

SPRAY; NASAL

IMITREX

+	GLAXOSMITHKLINE	5MG/SPRAY	N20626	001	Aug 26, 1997
+		10MG/SPRAY	N20626	002	Aug 26, 1997
+		20MG/SPRAY	N20626	003	Aug 26, 1997

SUMATRIPTAN SUCCINATE

INJECTABLE; SUBCUTANEOUS

IMITREX

+	GLAXOSMITHKLINE	EQ 6MG BASE/0.5ML (12MG/ML)	N20080	001	Dec 28, 1992
	IMITREX STATDOSE				
+	GLAXOSMITHKLINE	EQ 4MG BASE/0.5ML (8MG/ML)	N20080	002	Feb 01, 2006
+		EQ 6MG BASE/0.5ML (12MG/ML)	N20080	003	Dec 23, 1996

TABLET; ORAL

IMITREX

	GLAXOSMITHKLINE	EQ 25MG BASE	N20132	002	Jun 01, 1995
		EQ 50MG BASE	N20132	003	Jun 01, 1995
+		EQ 100MG BASE	N20132	001	Jun 01, 1995

SUNITINIB MALATE

CAPSULE; ORAL

SUTENT

	CPPI CV	EQ 12.5MG BASE	N21938	001	Jan 26, 2006
		EQ 25MG BASE	N21938	002	Jan 26, 2006
+		EQ 50MG BASE	N21938	003	Jan 26, 2006

TACRINE HYDROCHLORIDE

CAPSULE; ORAL

COGNEX

	SCIELE PHARMA INC	EQ 10MG BASE	N20070	001	Sep 09, 1993
		EQ 20MG BASE	N20070	002	Sep 09, 1993
		EQ 30MG BASE	N20070	003	Sep 09, 1993
+		EQ 40MG BASE	N20070	004	Sep 09, 1993

TACROLIMUS

CAPSULE; ORAL

PROGRAF

	ASTELLAS	EQ 0.5MG BASE	N50708	003	Aug 24, 1998
		EQ 1MG BASE	N50708	001	Apr 08, 1994
+		EQ 5MG BASE	N50708	002	Apr 08, 1994

INJECTABLE; INJECTION

PROGRAF

+	ASTELLAS	EQ 5MG BASE/ML	N50709	001	Apr 08, 1994
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PRESCRIPTION DRUG PRODUCT LIST

3 - 337 (of 370)

TACROLIMUS

OINTMENT; TOPICAL
PROTOPIC

ASTELLAS	0.03%	N50777 001	Dec 08, 2000
+	0.1%	N50777 002	Dec 08, 2000

TADALAFIL

TABLET; ORAL
CIALIS

LILLY ICOS	5MG	N21368 001	Nov 21, 2003
	10MG	N21368 002	Nov 21, 2003
+	20MG	N21368 003	Nov 21, 2003

TALC

AEROSOL, METERED; INTRAPLEURAL
SCLEROSOL

+ BRYAN	400MG/SPRAY	N20587 001	Dec 24, 1997
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POWDER; INTRAPLEURAL
TALC

+ BRYAN	5GM/BOT	N21388 001	Dec 15, 2003
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TAMOXIFEN CITRATE

SOLUTION; ORAL
SOLTAMOX

+ ROSEMONT	EQ 10MG BASE/5ML	N21807 001	Oct 29, 2005
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TABLET; ORAL

NOLVADEX

<u>AB</u> ASTRAZENECA	<u>EQ 10MG BASE</u>	<u>N17970 001</u>	
<u>AB</u> +	<u>EQ 20MG BASE</u>	<u>N17970 002</u>	Mar 21, 1994

TAMOXIFEN CITRATE

<u>AB</u> AEGIS PHARMS	<u>EQ 10MG BASE</u>	<u>N76398 001</u>	Mar 31, 2003
<u>AB</u>	<u>EQ 20MG BASE</u>	<u>N76398 002</u>	Mar 31, 2003
<u>AB</u> ANDRX PHARMS	<u>EQ 10MG BASE</u>	<u>N76179 001</u>	Feb 20, 2003
<u>AB</u>	<u>EQ 20MG BASE</u>	<u>N76179 002</u>	Feb 20, 2003
<u>AB</u> BARR	<u>EQ 10MG BASE</u>	<u>N70929 001</u>	Feb 20, 2003
<u>AB</u>	<u>EQ 20MG BASE</u>	<u>N70929 002</u>	Feb 20, 2003
<u>AB</u> IVAX PHARMS	<u>EQ 10MG BASE</u>	<u>N75740 001</u>	Feb 20, 2003
<u>AB</u>	<u>EQ 20MG BASE</u>	<u>N75740 002</u>	Feb 20, 2003
<u>AB</u> MYLAN	<u>EQ 10MG BASE</u>	<u>N74732 002</u>	Feb 20, 2003
<u>AB</u>	<u>EQ 20MG BASE</u>	<u>N74732 001</u>	Feb 20, 2003
<u>AB</u> PHARMACHEMIE	<u>EQ 20MG BASE</u>	<u>N74858 001</u>	Feb 20, 2003
<u>AB</u> ROXANE	<u>EQ 10MG BASE</u>	<u>N76027 001</u>	Feb 20, 2003
<u>AB</u>	<u>EQ 20MG BASE</u>	<u>N76027 002</u>	Feb 20, 2003
<u>AB</u> TEVA	<u>EQ 10MG BASE</u>	<u>N74504 001</u>	Apr 28, 2003
<u>AB</u>	<u>EQ 10MG BASE</u>	<u>N75797 001</u>	Feb 20, 2003
<u>AB</u>	<u>EQ 20MG BASE</u>	<u>N74504 002</u>	Apr 28, 2003

TAMSULOSIN HYDROCHLORIDE

CAPSULE; ORAL
FLOMAX

+ BOEHRINGER INGELHEIM	0.4MG	N20579 001	Apr 15, 1997
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TAZAROTENE

CREAM; TOPICAL
AVAGE

+ ALLERGAN	0.1%	N21184 003	Sep 30, 2002
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TAZORAC

+ ALLERGAN	0.05%	N21184 001	Sep 29, 2000
+	0.1%	N21184 002	Sep 29, 2000

PRESCRIPTION DRUG PRODUCT LIST

3 - 338 (of 370)

TAZAROTENE

GEL; TOPICAL
TAZORAC

+ ALLERGAN	0.05%	N20600	001	Jun 13, 1997
+	0.1%	N20600	002	Jun 13, 1997

TECHNETIUM TC-99M ALBUMIN AGGREGATED KIT

INJECTABLE; INJECTION
MACROTEC

BS BRACCO	N/A	N17833	001	
BS PULMOLITE				
BS CIS	N/A	N17776	001	
BS TECHNESCAN MAA				
BS MALLINCKRODT	N/A	N17842	001	
BS TECHNITIUM TC 99M ALBUMIN AGGREGATED KIT				
BS DRAXIMAGE	N/A	N17881	001	Dec 30, 1987

TECHNETIUM TC-99M BICISATE KIT

INJECTABLE; INJECTION
NEUROLITE

BRISTOL MYERS SQUIBB	N/A/VIAL	N20256	001	Nov 23, 1994
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TECHNETIUM TC-99M DISOFENIN KIT

INJECTABLE; INJECTION
HEPATOLITE

CIS	N/A	N18467	001	Mar 16, 1982
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TECHNETIUM TC-99M EXAMETAZIME KIT

INJECTABLE; INJECTION
CERETEC

GE HEALTHCARE	N/A	N19829	001	Dec 30, 1988
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TECHNETIUM TC-99M GLUCEPTATE KIT

INJECTABLE; INJECTION
TECHNESCAN GLUCEPTATE

DRAXIMAGE	N/A	N18272	001	Jan 27, 1982
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TECHNETIUM TC-99M MEBROFENIN KIT

INJECTABLE; INJECTION
CHOLETEC

+ BRACCO	N/A	N18963	001	Jan 21, 1987
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TECHNETIUM TC-99M MEDRONATE KIT

INJECTABLE; INJECTION
CIS-MDP

<u>AP</u> CIS	N/A	<u>N18124</u>	<u>001</u>	
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<u>AP</u> + <u>DRAXIMAGE MDP</u>	N/A	<u>N18035</u>	<u>001</u>	
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<u>AP</u> + <u>MDP-BRACCO</u>	N/A	<u>N18107</u>	<u>001</u>	
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<u>AP</u> BRACCO	N/A	<u>N18107</u>	<u>001</u>	
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<u>AP</u> <u>TECHNETIUM TC 99M MPI MDP</u>	N/A	<u>N18141</u>	<u>001</u>	
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<u>AP</u> GE HEALTHCARE	N/A	<u>N18141</u>	<u>001</u>	
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TECHNETIUM TC-99M MERTIATIDE KIT

INJECTABLE; INJECTION
TECHNESCAN MAG3

MALLINCKRODT	N/A	N19882	001	Jun 15, 1990
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PRESCRIPTION DRUG PRODUCT LIST

3 - 339 (of 370)

TECHNETIUM TC-99M OXIDRONATE KIT

INJECTABLE; INJECTION
TECHNESCAN

+ MALLINCKRODT	N/A	N18321	001
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TECHNETIUM TC-99M PENTETATE KIT

INJECTABLE; INJECTION

<u>AN-DTPA</u>		<u>N17714</u>	<u>001</u>	
<u>AP</u> CIS	N/A			
<u>DTPA</u>				
<u>AP</u> DRAXIMAGE	N/A	<u>N18511</u>	<u>001</u>	Dec 29, 1989
<u>MPI DTPA KIT - CHELATE</u>				
<u>AP</u> GE HEALTHCARE	N/A	<u>N17255</u>	<u>001</u>	
<u>TECHNETIUM TC-99M PENTETATE KIT</u>				
<u>AP</u> GE HEALTHCARE	N/A	<u>N17264</u>	<u>002</u>	

TECHNETIUM TC-99M PYROPHOSPHATE KIT

INJECTABLE; INJECTION

<u>CIS-PYRO</u>		<u>N19039</u>	<u>001</u>	
<u>AP</u> CIS	N/A			Jun 30, 1987
<u>PHOSPHOTEC</u>				
<u>AP</u> BRACCO	N/A	<u>N17680</u>	<u>001</u>	
<u>TECHNESCAN PYP KIT</u>				
<u>AP</u> MALLINCKRODT	N/A	<u>N17538</u>	<u>001</u>	

TECHNETIUM TC-99M RED BLOOD CELL KIT

INJECTABLE; INJECTION
ULTRATAG

MALLINCKRODT	N/A	N19981	001	Jun 10, 1991
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TECHNETIUM TC-99M SESTAMIBI KIT

INJECTABLE; INJECTION
CARDIOLITE

+ BRISTOL MYERS SQUIBB	N/A	N19785	001	Dec 21, 1990
MIRALUMA				
+ BRISTOL MYERS SQUIBB	N/A	N19785	003	May 23, 1997

TECHNETIUM TC-99M SODIUM PERTECHNETATE GENERATOR

SOLUTION; INJECTION, ORAL
TECHNELITE

BRISTOL MYERS SQUIBB	0.0083-2.7 CI/GENERATOR	N17771	001
ULTRA-TECHNEKOW FM			
MALLINCKRODT	0.25-3 CI/GENERATOR	N17243	002

TECHNETIUM TC-99M SUCCIMER KIT

INJECTABLE; INJECTION
MPI DMSA KIDNEY REAGENT

GE HEALTHCARE	N/A	N17944	001	May 18, 1982
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TECHNETIUM TC-99M SULFUR COLLOID KIT

SOLUTION; INJECTION, ORAL

<u>AN-SULFUR COLLOID</u>		<u>N17858</u>	<u>001</u>
<u>AP</u> CIS	N/A		
<u>TECHNECOLL</u>			
<u>AP</u> MALLINCKRODT	N/A	<u>N17059</u>	<u>001</u>

PRESCRIPTION DRUG PRODUCT LIST

3 - 340 (of 370)

TECHNETIUM TC-99M TETROFOSMIN KIT

INJECTABLE; INJECTION
MYOVUE

+ GE HEALTHCARE	N/A	N20372 001	Feb 09, 1996
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TEGASEROD MALEATE

TABLET; ORAL
ZELNORM

NOVARTIS	EQ 2MG BASE	N21200 001	Jul 24, 2002
+	EQ 6MG BASE	N21200 002	Jul 24, 2002

TELBIVUDINE

TABLET; ORAL
TYZEKA

+ IDENIX	600MG	N22011 001	Oct 25, 2006
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TELITHROMYCIN

TABLET; ORAL
KETEK

SANOFI AVENTIS US	300MG	N21144 002	Feb 09, 2005
+	400MG	N21144 001	Apr 01, 2004

TELMISARTAN

TABLET; ORAL
MICARDIS

BOEHRINGER INGELHEIM	20MG	N20850 003	Apr 04, 2000
	40MG	N20850 001	Nov 10, 1998
+	80MG	N20850 002	Nov 10, 1998

TEMAZEPAM

CAPSULE; ORAL

RESTORIL

<u>AB</u> TYCO HLTHCARE	<u>15MG</u>	<u>N18163</u> <u>001</u>	
<u>AB</u> +	<u>30MG</u>	<u>N18163</u> <u>002</u>	
	7.5MG	N18163 003	Oct 25, 1991
	22.5MG	N18163 004	Nov 02, 2004

TEMAZEPAM

<u>AB</u> ACTAVIS ELIZABETH	<u>15MG</u>	<u>N71638</u> <u>001</u>	Aug 07, 1987
<u>AB</u>	<u>30MG</u>	<u>N71620</u> <u>001</u>	Aug 07, 1987
<u>AB</u> MYLAN	<u>15MG</u>	<u>N70919</u> <u>001</u>	Jul 07, 1986
<u>AB</u>	<u>30MG</u>	<u>N70920</u> <u>001</u>	Jul 10, 1986
<u>AB</u> PAR PHARM	<u>15MG</u>	<u>N71456</u> <u>001</u>	Apr 21, 1987
<u>AB</u>	<u>30MG</u>	<u>N71457</u> <u>001</u>	Apr 21, 1987
<u>AB</u> SANDOZ	<u>15MG</u>	<u>N71427</u> <u>001</u>	Jan 12, 1988
<u>AB</u>	<u>30MG</u>	<u>N71428</u> <u>001</u>	Jan 12, 1988
<u>AB</u> WATSON LABS	<u>15MG</u>	<u>N71446</u> <u>001</u>	May 21, 1993
<u>AB</u>	<u>30MG</u>	<u>N71447</u> <u>001</u>	May 21, 1993

TEMOZOLOMIDE

CAPSULE; ORAL
TEMODAR

SCHERING	5MG	N21029 001	Aug 11, 1999
	20MG	N21029 002	Aug 11, 1999
	100MG	N21029 003	Aug 11, 1999
	140MG	N21029 005	Oct 19, 2006
	180MG	N21029 006	Oct 19, 2006
+	250MG	N21029 004	Aug 11, 1999

PRESCRIPTION DRUG PRODUCT LIST

3 - 341 (of 370)

TENIPOSIDE

INJECTABLE; INJECTION
VUMON

+ BRISTOL MYERS SQUIBB	10MG/ML	N20119	001	Jul 14, 1992
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TENOFOVIR DISOPROXIL FUMARATE

TABLET; ORAL
VIREAD

+ GILEAD	300MG	N21356	001	Oct 26, 2001
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TERAZOSIN HYDROCHLORIDE

CAPSULE; ORAL

HYTRIN

<u>AB</u> ABBOTT	<u>EQ 1MG BASE</u>	<u>N20347</u>	<u>001</u>	Dec 14, 1994
<u>AB</u> +	<u>EQ 2MG BASE</u>	<u>N20347</u>	<u>002</u>	Dec 14, 1994
<u>AB</u>	<u>EQ 5MG BASE</u>	<u>N20347</u>	<u>003</u>	Dec 14, 1994
<u>AB</u>	<u>EQ 10MG BASE</u>	<u>N20347</u>	<u>004</u>	Dec 14, 1994

TERAZOSIN HYDROCHLORIDE

<u>AB</u> JUBILANT PHARMS	<u>EQ 1MG BASE</u>	<u>N75317</u>	<u>001</u>	Dec 20, 2004
<u>AB</u>	<u>EQ 2MG BASE</u>	<u>N75317</u>	<u>002</u>	Dec 20, 2004
<u>AB</u>	<u>EQ 5MG BASE</u>	<u>N75317</u>	<u>003</u>	Dec 20, 2004
<u>AB</u>	<u>EQ 10MG BASE</u>	<u>N75317</u>	<u>004</u>	Dec 20, 2004
<u>AB</u> MYLAN	<u>EQ 1MG BASE</u>	<u>N75140</u>	<u>002</u>	Feb 11, 2000
<u>AB</u>	<u>EQ 2MG BASE</u>	<u>N75140</u>	<u>003</u>	Feb 11, 2000
<u>AB</u>	<u>EQ 5MG BASE</u>	<u>N75140</u>	<u>001</u>	Feb 11, 2000
<u>AB</u>	<u>EQ 10MG BASE</u>	<u>N75140</u>	<u>004</u>	Feb 11, 2000
<u>AB</u> MYLAN TECHNOLOGIES	<u>EQ 1MG BASE</u>	<u>N75384</u>	<u>001</u>	Dec 01, 2000
<u>AB</u>	<u>EQ 2MG BASE</u>	<u>N75384</u>	<u>002</u>	Dec 01, 2000
<u>AB</u>	<u>EQ 5MG BASE</u>	<u>N75384</u>	<u>003</u>	Dec 01, 2000
<u>AB</u>	<u>EQ 10MG BASE</u>	<u>N75384</u>	<u>004</u>	Dec 01, 2000
<u>AB</u> RANBAXY	<u>EQ 1MG BASE</u>	<u>N76021</u>	<u>001</u>	Aug 22, 2002
<u>AB</u>	<u>EQ 2MG BASE</u>	<u>N76021</u>	<u>002</u>	Aug 22, 2002
<u>AB</u>	<u>EQ 5MG BASE</u>	<u>N76021</u>	<u>003</u>	Aug 22, 2002
<u>AB</u>	<u>EQ 10MG BASE</u>	<u>N76021</u>	<u>004</u>	Aug 22, 2002
<u>AB</u> SANDOZ	<u>EQ 1MG BASE</u>	<u>N74823</u>	<u>001</u>	Mar 30, 1998
<u>AB</u>	<u>EQ 1MG BASE</u>	<u>N75667</u>	<u>001</u>	Jul 28, 2000
<u>AB</u>	<u>EQ 2MG BASE</u>	<u>N74823</u>	<u>002</u>	Mar 30, 1998
<u>AB</u>	<u>EQ 2MG BASE</u>	<u>N75667</u>	<u>002</u>	Jul 28, 2000
<u>AB</u>	<u>EQ 5MG BASE</u>	<u>N74823</u>	<u>003</u>	Mar 30, 1998
<u>AB</u>	<u>EQ 5MG BASE</u>	<u>N75667</u>	<u>003</u>	Jul 28, 2000
<u>AB</u>	<u>EQ 10MG BASE</u>	<u>N74823</u>	<u>004</u>	Mar 30, 1998
<u>AB</u>	<u>EQ 10MG BASE</u>	<u>N75667</u>	<u>004</u>	Jul 28, 2000
<u>AB</u> TORPHARM	<u>EQ 1MG BASE</u>	<u>N75498</u>	<u>001</u>	Apr 12, 2001
<u>AB</u>	<u>EQ 2MG BASE</u>	<u>N75498</u>	<u>002</u>	Apr 12, 2001
<u>AB</u>	<u>EQ 5MG BASE</u>	<u>N75498</u>	<u>003</u>	Apr 12, 2001
<u>AB</u>	<u>EQ 10MG BASE</u>	<u>N75498</u>	<u>004</u>	Apr 12, 2001

TABLET; ORAL

HYTRIN

<u>AB</u> ABBOTT	<u>EQ 1MG BASE</u>	<u>N19057</u>	<u>001</u>	Aug 07, 1987
<u>AB</u> +	<u>EQ 2MG BASE</u>	<u>N19057</u>	<u>002</u>	Aug 07, 1987
<u>AB</u>	<u>EQ 5MG BASE</u>	<u>N19057</u>	<u>003</u>	Aug 07, 1987
<u>AB</u>	<u>EQ 10MG BASE</u>	<u>N19057</u>	<u>004</u>	Aug 07, 1987

TERAZOSIN HYDROCHLORIDE

<u>AB</u> SANDOZ	<u>EQ 1MG BASE</u>	<u>N74315</u>	<u>001</u>	Dec 31, 1998
<u>AB</u>	<u>EQ 1MG BASE</u>	<u>N74657</u>	<u>001</u>	Apr 28, 2000
<u>AB</u>	<u>EQ 2MG BASE</u>	<u>N74315</u>	<u>002</u>	Dec 31, 1998
<u>AB</u>	<u>EQ 2MG BASE</u>	<u>N74657</u>	<u>002</u>	Apr 28, 2000
<u>AB</u>	<u>EQ 5MG BASE</u>	<u>N74315</u>	<u>003</u>	Dec 31, 1998
<u>AB</u>	<u>EQ 5MG BASE</u>	<u>N74657</u>	<u>003</u>	Apr 28, 2000
<u>AB</u>	<u>EQ 10MG BASE</u>	<u>N74315</u>	<u>004</u>	Dec 31, 1998

PRESCRIPTION DRUG PRODUCT LIST

3 - 342 (of 370)

TERAZOSIN HYDROCHLORIDE

TABLET; ORAL

TERAZOSIN HYDROCHLORIDE

<u>AB</u>	SANDOZ	<u>EQ 10MG BASE</u>	<u>N74657</u>	<u>004</u>	Apr 28, 2000
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TERBINAFINE

GEL; TOPICAL

LAMISIL

	NOVARTIS	1%	N20846	001	Apr 29, 1998
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TERBINAFINE HYDROCHLORIDE

SOLUTION; TOPICAL

LAMISIL

	+ NOVARTIS	1%	N20749	001	Oct 17, 1997
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TABLET; ORAL

LAMISIL

	+ NOVARTIS	EQ 250MG BASE	N20539	001	May 10, 1996
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TERBUTALINE SULFATE

INJECTABLE; INJECTION

TERBUTALINE SULFATE

<u>AP</u>	ABRAXIS PHARM	<u>1MG/ML</u>	<u>N76887</u>	<u>001</u>	May 26, 2004
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<u>AP</u>	+ BEDFORD	<u>1MG/ML</u>	<u>N76770</u>	<u>001</u>	Apr 23, 2004
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<u>AP</u>	SICOR PHARMS	<u>1MG/ML</u>	<u>N76853</u>	<u>001</u>	Jul 20, 2004
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TABLET; ORAL

BRETHINE

<u>AB</u>	AAIPHARMA LLC	<u>2.5MG</u>	<u>N17849</u>	<u>001</u>	
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<u>AB</u>	+	<u>5MG</u>	<u>N17849</u>	<u>002</u>	
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TERBUTALINE SULFATE

<u>AB</u>	IMPAX LABS	<u>2.5MG</u>	<u>N75877</u>	<u>001</u>	Jun 26, 2001
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<u>AB</u>		<u>5MG</u>	<u>N75877</u>	<u>002</u>	Jun 26, 2001
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<u>AB</u>	LANNETT	<u>2.5MG</u>	<u>N77152</u>	<u>001</u>	Mar 25, 2005
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<u>AB</u>		<u>5MG</u>	<u>N77152</u>	<u>002</u>	Mar 25, 2005
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TERCONAZOLE

CREAM; VAGINAL

TERAZOL 3

<u>AB</u>	+ ORTHO MCNEIL PHARM	<u>0.8%</u>	<u>N19964</u>	<u>001</u>	Feb 21, 1991
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TERAZOL 7

<u>AB</u>	+ ORTHO MCNEIL PHARM	<u>0.4%</u>	<u>N19579</u>	<u>001</u>	Dec 31, 1987
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TERCONAZOLE

<u>AB</u>	ALTANA	<u>0.4%</u>	<u>N76712</u>	<u>001</u>	Feb 18, 2005
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BX	+	0.8%	N21735	001	Oct 01, 2004
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<u>AB</u>	TARO	<u>0.4%</u>	<u>N76043</u>	<u>001</u>	Jan 19, 2005
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<u>AB</u>		<u>0.8%</u>	<u>N75953</u>	<u>001</u>	Apr 06, 2004
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SUPPOSITORY; VAGINAL

TERAZOL 3

<u>AB</u>	+ ORTHO MCNEIL PHARM	<u>80MG</u>	<u>N19641</u>	<u>001</u>	May 24, 1988
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TERCONAZOLE

<u>AB</u>	ALTANA	<u>80MG</u>	<u>N76850</u>	<u>001</u>	Jul 12, 2006
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<u>AB</u>	PERRIGO NEW YORK	<u>80MG</u>	<u>N77149</u>	<u>001</u>	Mar 17, 2006
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TERIPARATIDE RECOMBINANT HUMAN

INJECTABLE; SUBCUTANEOUS

FORTEO

	+ LILLY	EQ 0.75MG/3ML (0.25MG/ML)	N21318	001	Nov 26, 2002
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27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 343 (of 370)

TESTOLACTONE

TABLET; ORAL

TESLAC

+ BRISTOL MYERS SQUIBB 50MG

N16118 001

TESTOSTERONE

FILM, EXTENDED RELEASE; TRANSDERMAL

ANDRODERM

+ WATSON LABS 2.5MG/24HR

N20489 001 Sep 29, 1995

+ 5MG/24HR

N20489 002 May 02, 1997

GEL; TRANSDERMAL

ANDROGELAB + UNIMED PHARMS 1%N21015 001 Feb 28, 2000

TESTIM

BX + AUXILIUM PHARMS 1%

N21454 001 Oct 31, 2002

TESTOSTERONEAB WATSON LABS 1%N76737 001 Jan 27, 2006

INJECTABLE; INJECTION

TESTOSTERONE

+ WATSON LABS 100MG/ML

N86417 001 Jul 07, 1983

PELLET; IMPLANTATION

TESTOSTERONE

BARTOR 75MG

N80911 001

TABLET, EXTENDED RELEASE; BUCCAL

STRIANT

+ COLUMBIA LABS 30MG

N21543 001 Jun 19, 2003

TESTOSTERONE CYPIONATE

INJECTABLE; INJECTION

DEPO-TESTOSTERONEAO + PHARMACIA AND UPJOHN 100MG/MLN85635 002AO + 200MG/MLN85635 003TESTOSTERONE CYPIONATEAO PADDOCK 200MG/MLN40530 001 Jan 31, 2005AO SANDOZ 100MG/MLN40615 001 Aug 10, 2006AO 200MG/MLN40615 002 Aug 10, 2006AO SYNERX PHARMA 200MG/MLN40652 001 Dec 11, 2006AO WATSON LABS 100MG/MLN86029 001AO 200MG/MLN86030 001TESTOSTERONE ENANTHATE

INJECTABLE; INJECTION

DELATESTRYLAO + INDEVUS PHARMS 200MG/MLN09165 003TESTOSTERONE ENANTHATEAO WATSON LABS 200MG/MLN85598 001TESTOSTERONE ETHANATEAO PADDOCK 200MG/MLN40575 001 Jun 14, 2006TESTOSTERONE PROPIONATE

INJECTABLE; INJECTION

TESTOSTERONE PROPIONATE

+ WATSON LABS 25MG/ML

N80188 001

+ 50MG/ML

N80188 002

+ 100MG/ML

N80188 003

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 344 (of 370)

TETRACYCLINE HYDROCHLORIDE

CAPSULE; ORAL

BRISTACYCLINE

<u>AB</u>	BRISTOL	<u>250MG</u>	<u>N61658</u>	<u>001</u>
<u>AB</u>		<u>250MG</u>	<u>N61888</u>	<u>001</u>
<u>AB</u>		<u>500MG</u>	<u>N61658</u>	<u>002</u>
<u>AB</u>		<u>500MG</u>	<u>N61888</u>	<u>002</u>

TETRACYCLINE HYDROCHLORIDE

<u>AB</u>	BARR	<u>250MG</u>	<u>N61837</u>	<u>001</u>
<u>AB</u>		<u>500MG</u>	<u>N61837</u>	<u>002</u>
<u>AB</u>	IMPAX LABS	<u>250MG</u>	<u>N60469</u>	<u>001</u>
<u>AB</u>		<u>500MG</u>	<u>N60469</u>	<u>003</u>
		100MG	N60469	002
<u>AB</u>	IVAX PHARMS	<u>250MG</u>	<u>N60704</u>	<u>001</u>
<u>AB</u>	+	<u>500MG</u>	<u>N60704</u>	<u>002</u>
<u>AB</u>	LABS ATRAL	<u>250MG</u>	<u>N62752</u>	<u>001</u>
<u>AB</u>		<u>500MG</u>	<u>N62752</u>	<u>002</u>
<u>AB</u>	MUTUAL PHARM	<u>250MG</u>	<u>N60736</u>	<u>001</u>
<u>AB</u>		<u>500MG</u>	<u>N60736</u>	<u>002</u>
<u>AB</u>	MYLAN	<u>250MG</u>	<u>N60783</u>	<u>001</u>
<u>AB</u>		<u>500MG</u>	<u>N60783</u>	<u>002</u>

Aug 12, 1988

Aug 12, 1988

SUSPENSION; ORAL

SUMYCIN

+	PAR PHARM	125MG/5ML	N60400	001
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TABLET; ORAL

SUMYCIN

	PAR PHARM	250MG	N61147	001
+		500MG	N61147	004

TETRACYCLINE PHOSPHATE COMPLEX

CAPSULE; ORAL

TETREX

	BRISTOL	EQ 100MG HCL	N61653	001
		EQ 250MG HCL	N61653	002
		EQ 250MG HCL	N61889	002
+		EQ 500MG HCL	N61653	003
		EQ 500MG HCL	N61889	001

TETRAHYDROZOLINE HYDROCHLORIDE

SOLUTION; NASAL

TYZINE

+	KENWOOD LABS	0.05%	N86576	002
		0.1%	N86576	001

SPRAY; NASAL

TYZINE

+	KENWOOD LABS	0.1%	N86576	003
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THALIDOMIDE

CAPSULE; ORAL

THALOMID

	CELGENE	50MG	N20785	001	Jul 16, 1998
		100MG	N20785	002	Jan 17, 2003
+		200MG	N20785	003	Jan 17, 2003

THALLOUS CHLORIDE, TL-201

INJECTABLE; INJECTION

THALLOUS CHLORIDE TL 201

<u>AP</u>	+	BRISTOL MYERS SQUIBB	<u>1mCi/ML</u>	<u>N17806</u>	<u>001</u>	
<u>AP</u>	+	GE HEALTHCARE	<u>1mCi/ML</u>	<u>N18110</u>	<u>002</u>	Feb 27, 1996

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 345 (of 370)

THALLOUS CHLORIDE, TL-201

INJECTABLE; INJECTION

THALLOUS CHLORIDE TL 201

<u>AP</u>	+	MALLINCKRODT	<u>1mCi/ML</u>	<u>N18150</u>	<u>001</u>	
<u>AP</u>		TRACE RADIOCHEMICALS	<u>1mCi/ML</u>	<u>N75569</u>	<u>001</u>	Nov 21, 2001

INJECTABLE; INTRAVENOUS

THALLOUS CHLORIDE TL 201

<u>AP</u>	+	BRISTOL MYERS SQUIBB	<u>2mCi/ML</u>	<u>N17806</u>	<u>002</u>	Oct 09, 1998
<u>AP</u>		TCI MEDCL	<u>2mCi/ML</u>	<u>N77698</u>	<u>001</u>	Nov 09, 2006

THEOPHYLLINECAPSULE, EXTENDED RELEASE; ORAL
THEO-24

BC		UCB INC	100MG	N87942	001	Aug 22, 1983
BC			200MG	N87943	001	Aug 22, 1983
BC			300MG	N87944	001	Aug 22, 1983
			400MG	N81034	001	Feb 28, 1992

THEOPHYLLINE

BC		INWOOD LABS	100MG	N40052	001	Feb 14, 1994
BC			200MG	N40052	003	Feb 14, 1994
BC	+		300MG	N40052	004	Feb 14, 1994
			125MG	N40052	002	Feb 14, 1994

ELIXIR; ORAL

ELIXOPHYLLIN

<u>AA</u>		FOREST LABS	<u>80MG/15ML</u>	<u>N85186</u>	<u>001</u>	
		<u>THEOPHYLLINE</u>				
<u>AA</u>		ACTAVIS MID ATLANTIC	<u>80MG/15ML</u>	<u>N85863</u>	<u>001</u>	
<u>AA</u>		MORTON GROVE	<u>80MG/15ML</u>	<u>N86748</u>	<u>001</u>	
<u>AA</u>		PHARM ASSOC	<u>80MG/15ML</u>	<u>N86720</u>	<u>001</u>	
<u>AA</u>		TARO	<u>80MG/15ML</u>	<u>N89626</u>	<u>001</u>	Oct 28, 1988

INJECTABLE; INJECTION

THEOPHYLLINE 0.04% AND DEXTROSE 5% IN PLASTIC CONTAINER

<u>AP</u>	+	B BRAUN	<u>40MG/100ML</u>	<u>N19826</u>	<u>001</u>	Aug 14, 1992
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THEOPHYLLINE 0.08% AND DEXTROSE 5% IN PLASTIC CONTAINER

<u>AP</u>	+	B BRAUN	<u>80MG/100ML</u>	<u>N19826</u>	<u>002</u>	Aug 14, 1992
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THEOPHYLLINE 0.16% AND DEXTROSE 5% IN PLASTIC CONTAINER

<u>AP</u>	+	B BRAUN	<u>160MG/100ML</u>	<u>N19826</u>	<u>003</u>	Aug 14, 1992
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THEOPHYLLINE 0.32% AND DEXTROSE 5% IN PLASTIC CONTAINER

<u>AP</u>	+	B BRAUN	<u>320MG/100ML</u>	<u>N19826</u>	<u>006</u>	Aug 14, 1992
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THEOPHYLLINE AND DEXTROSE 5% IN PLASTIC CONTAINER

<u>AP</u>	+	BAXTER HLTHCARE	<u>4MG/ML</u>	<u>N18649</u>	<u>007</u>	Jul 26, 1982
<u>AP</u>	+		<u>40MG/100ML</u>	<u>N18649</u>	<u>001</u>	Jul 26, 1982
<u>AP</u>	+		<u>80MG/100ML</u>	<u>N18649</u>	<u>002</u>	Jul 26, 1982
<u>AP</u>	+		<u>160MG/100ML</u>	<u>N18649</u>	<u>003</u>	Jul 26, 1982
<u>AP</u>	+		<u>200MG/100ML</u>	<u>N18649</u>	<u>004</u>	Jul 26, 1982
<u>AP</u>	+		<u>320MG/100ML</u>	<u>N18649</u>	<u>006</u>	Nov 13, 1985
<u>AP</u>	+		<u>400MG/100ML</u>	<u>N18649</u>	<u>005</u>	Jul 26, 1982

THEOPHYLLINE IN DEXTROSE 5% IN PLASTIC CONTAINER

<u>AP</u>	+	HOSPIRA	<u>4MG/ML</u>	<u>N19211</u>	<u>007</u>	Dec 14, 1984
<u>AP</u>	+		<u>40MG/100ML</u>	<u>N19211</u>	<u>001</u>	Dec 14, 1984
<u>AP</u>	+		<u>80MG/100ML</u>	<u>N19211</u>	<u>002</u>	Dec 14, 1984
<u>AP</u>	+		<u>160MG/100ML</u>	<u>N19211</u>	<u>003</u>	Dec 14, 1984
<u>AP</u>	+		<u>200MG/100ML</u>	<u>N19211</u>	<u>004</u>	Dec 14, 1984
<u>AP</u>	+		<u>320MG/100ML</u>	<u>N19211</u>	<u>006</u>	Jan 20, 1988
<u>AP</u>	+		<u>400MG/100ML</u>	<u>N19211</u>	<u>005</u>	Dec 14, 1984

SOLUTION; ORAL

THEOPHYLLINE

	+	ROXANE	80MG/15ML	N87449	001	Sep 15, 1983
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PRESCRIPTION DRUG PRODUCT LIST

3 - 346 (of 370)

THEOPHYLLINE

TABLET; ORAL				
QUIBRON-T				
	+ MONARCH PHARMS	300MG	N88656 001	Aug 22, 1985
THEOLAIR				
	+ 3M	125MG	N86399 001	
	+	250MG	N86399 002	
TABLET, EXTENDED RELEASE; ORAL				
QUIBRON-T/SR				
BC	MONARCH PHARMS	300MG	N87563 001	Jun 21, 1983
<u>THEOCHRON</u>				
<u>AB</u>	INWOOD LABS	<u>100MG</u>	<u>N88320</u> <u>001</u>	Feb 21, 1985
<u>AB</u>		<u>200MG</u>	<u>N88321</u> <u>001</u>	Feb 21, 1985
<u>AB</u>		<u>300MG</u>	<u>N87400</u> <u>002</u>	Jan 11, 1983
<u>THEOPHYLLINE</u>				
<u>AB</u>	INWOOD LABS	<u>450MG</u>	<u>N40034</u> <u>001</u>	Apr 28, 1995
<u>AB</u>	NOSTRUM	<u>400MG</u>	<u>N40595</u> <u>001</u>	Apr 21, 2006
<u>AB</u>		<u>600MG</u>	<u>N40560</u> <u>002</u>	Apr 21, 2006
<u>AB</u>	+ PLIVA	<u>100MG</u>	<u>N89807</u> <u>001</u>	Apr 30, 1990
<u>AB</u>	+	<u>200MG</u>	<u>N89808</u> <u>001</u>	Apr 30, 1990
<u>AB</u>		<u>300MG</u>	<u>N89763</u> <u>001</u>	Apr 30, 1990
<u>AB</u>	+	<u>450MG</u>	<u>N81236</u> <u>001</u>	Nov 09, 1992
<u>UNIPHYL</u>				
<u>AB</u>	PURDUE FREDERICK	<u>400MG</u>	<u>N87571</u> <u>001</u>	Sep 01, 1982
<u>AB</u>	+	<u>600MG</u>	<u>N40086</u> <u>001</u>	Apr 15, 1996

THIABENDAZOLE

TABLET, CHEWABLE; ORAL				
MINTEZOL				
	+ MERCK	500MG	N16096 001	

THIAMINE HYDROCHLORIDE

INJECTABLE; INJECTION				
<u>THIAMINE HYDROCHLORIDE</u>				
<u>AP</u>	+ ABRAXIS PHARM	<u>100MG/ML</u>	<u>N80556</u> <u>001</u>	
<u>AP</u>	BAXTER HLTHCARE	<u>100MG/ML</u>	<u>N80575</u> <u>001</u>	
<u>AP</u>	HOSPIRA	<u>100MG/ML</u>	<u>N40079</u> <u>001</u>	May 03, 1996
<u>AP</u>	WATSON LABS	<u>100MG/ML</u>	<u>N80571</u> <u>001</u>	
	+	200MG/ML	N80571 002	

THIETHYLPERAZINE MALEATE

TABLET; ORAL				
TORECAN				
	+ NOVARTIS	10MG	N12753 001	

THIOGUANINE

TABLET; ORAL				
THIOGUANINE				
	+ GLAXOSMITHKLINE	40MG	N12429 001	

THIORIDAZINE HYDROCHLORIDE

CONCENTRATE; ORAL				
<u>THIORIDAZINE HYDROCHLORIDE</u>				
<u>AA</u>	ACTAVIS MID ATLANTIC	<u>100MG/ML</u>	<u>N88229</u> <u>001</u>	Aug 23, 1983
<u>AA</u>	PHARM ASSOC	<u>30MG/ML</u>	<u>N40187</u> <u>001</u>	Aug 28, 1997
<u>AA</u>		<u>100MG/ML</u>	<u>N40213</u> <u>001</u>	May 29, 1998
<u>AA</u>	+ TEVA PHARMS	<u>30MG/ML</u>	<u>N89602</u> <u>001</u>	Nov 09, 1987
<u>AA</u>	+	<u>100MG/ML</u>	<u>N89603</u> <u>001</u>	Nov 09, 1987

PRESCRIPTION DRUG PRODUCT LIST

3 - 347 (of 370)

THIORIDAZINE HYDROCHLORIDE

CONCENTRATE; ORAL

THIORIDAZINE HYDROCHLORIDE INTENSOL

<u>AA</u>	ROXANE	<u>30MG/ML</u>	<u>N88941</u>	<u>001</u>	Dec 16, 1985
<u>AA</u>		<u>100MG/ML</u>	<u>N88942</u>	<u>001</u>	Dec 16, 1985

TABLET; ORAL

THIORIDAZINE HYDROCHLORIDE

<u>AB</u>	IVAX PHARMS	<u>10MG</u>	<u>N88270</u>	<u>001</u>	Apr 14, 1983
<u>AB</u>		<u>15MG</u>	<u>N88271</u>	<u>001</u>	Apr 14, 1983
<u>AB</u>		<u>25MG</u>	<u>N88272</u>	<u>001</u>	Apr 14, 1983
<u>AB</u>	MUTUAL PHARM	<u>10MG</u>	<u>N89431</u>	<u>001</u>	Aug 01, 1986
<u>AB</u>		<u>25MG</u>	<u>N89432</u>	<u>001</u>	Aug 01, 1986
<u>AB</u>		<u>50MG</u>	<u>N88370</u>	<u>001</u>	Nov 18, 1983
<u>AB</u>		<u>50MG</u>	<u>N89433</u>	<u>001</u>	Aug 01, 1986
<u>AB</u>		<u>100MG</u>	<u>N89953</u>	<u>001</u>	Oct 07, 1988
<u>AB</u>	MYLAN	<u>10MG</u>	<u>N88001</u>	<u>001</u>	Mar 15, 1983
<u>AB</u>		<u>25MG</u>	<u>N88002</u>	<u>001</u>	Mar 15, 1983
<u>AB</u>		<u>50MG</u>	<u>N88003</u>	<u>001</u>	Mar 15, 1983
<u>AB</u>		<u>100MG</u>	<u>N88004</u>	<u>001</u>	Nov 18, 1983
<u>AB</u>	SANDOZ	<u>10MG</u>	<u>N88131</u>	<u>001</u>	Aug 30, 1983
<u>AB</u>		<u>15MG</u>	<u>N88132</u>	<u>001</u>	Aug 30, 1983
<u>AB</u>	+	<u>25MG</u>	<u>N88133</u>	<u>001</u>	Aug 30, 1983
<u>AB</u>		<u>50MG</u>	<u>N88134</u>	<u>001</u>	Aug 30, 1983
<u>AB</u>	+	<u>100MG</u>	<u>N88135</u>	<u>001</u>	Nov 20, 1984
<u>AB</u>		<u>150MG</u>	<u>N88136</u>	<u>001</u>	Sep 17, 1986
<u>AB</u>	+	<u>200MG</u>	<u>N88137</u>	<u>001</u>	Sep 17, 1986
<u>AB</u>	TEVA	<u>100MG</u>	<u>N88456</u>	<u>001</u>	May 17, 1985
<u>AB</u>	WATSON LABS	<u>10MG</u>	<u>N88476</u>	<u>001</u>	Nov 08, 1983
<u>AB</u>		<u>15MG</u>	<u>N88477</u>	<u>001</u>	Nov 08, 1983
<u>AB</u>		<u>25MG</u>	<u>N88478</u>	<u>001</u>	Nov 08, 1983
<u>AB</u>		<u>25MG</u>	<u>N88567</u>	<u>001</u>	May 11, 1984
<u>AB</u>		<u>50MG</u>	<u>N88479</u>	<u>001</u>	Nov 08, 1983
<u>AB</u>		<u>50MG</u>	<u>N88563</u>	<u>001</u>	May 11, 1984
<u>AB</u>		<u>100MG</u>	<u>N88564</u>	<u>001</u>	May 11, 1984
<u>AB</u>		<u>100MG</u>	<u>N88736</u>	<u>001</u>	Jul 24, 1984
<u>AB</u>		<u>150MG</u>	<u>N88869</u>	<u>001</u>	Jun 28, 1985
<u>AB</u>		<u>200MG</u>	<u>N88872</u>	<u>001</u>	Apr 26, 1985

THIOTEPA

INJECTABLE; INJECTION

THIOTEPA

<u>AP</u>	ABRAXIS PHARM	<u>15MG/VIAL</u>	<u>N75698</u>	<u>001</u>	Sep 20, 2001
<u>AP</u>	BEDFORD	<u>15MG/VIAL</u>	<u>N75547</u>	<u>001</u>	Apr 02, 2001
<u>AP</u>	+	<u>15MG/VIAL</u>	<u>N75730</u>	<u>001</u>	Apr 20, 2001
	+	<u>30MG/VIAL</u>	<u>N75730</u>	<u>002</u>	Apr 20, 2001

THIOTHIXENE

CAPSULE; ORAL

NAVANE

<u>AB</u>	PFIZER	<u>1MG</u>	<u>N16584</u>	<u>001</u>	
<u>AB</u>		<u>2MG</u>	<u>N16584</u>	<u>002</u>	
<u>AB</u>	+	<u>5MG</u>	<u>N16584</u>	<u>003</u>	
<u>AB</u>		<u>10MG</u>	<u>N16584</u>	<u>004</u>	
		<u>20MG</u>	<u>N16584</u>	<u>005</u>	

THIOTHIXENE

<u>AB</u>	MYLAN	<u>1MG</u>	<u>N71090</u>	<u>001</u>	Jun 23, 1987
<u>AB</u>		<u>2MG</u>	<u>N71091</u>	<u>001</u>	Jun 23, 1987
<u>AB</u>		<u>5MG</u>	<u>N71092</u>	<u>001</u>	Jun 23, 1987
<u>AB</u>		<u>10MG</u>	<u>N71093</u>	<u>001</u>	Jun 23, 1987
<u>AB</u>	SANDOZ	<u>1MG</u>	<u>N71610</u>	<u>001</u>	Jun 24, 1987

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 348 (of 370)

THIOTHIXENE

CAPSULE; ORAL

THIOTHIXENE

<u>AB</u>	SANDOZ	<u>2MG</u>	<u>N71570</u>	<u>001</u>	Jun 24, 1987
<u>AB</u>		<u>5MG</u>	<u>N71529</u>	<u>001</u>	Jun 24, 1987
<u>AB</u>		<u>10MG</u>	<u>N71530</u>	<u>001</u>	Jun 24, 1987
<u>AB</u>	WATSON LABS	<u>1MG</u>	<u>N70600</u>	<u>001</u>	Jun 05, 1987
<u>AB</u>		<u>2MG</u>	<u>N70601</u>	<u>001</u>	Jun 05, 1987
<u>AB</u>		<u>5MG</u>	<u>N70602</u>	<u>001</u>	Jun 05, 1987
<u>AB</u>		<u>10MG</u>	<u>N70603</u>	<u>001</u>	Jun 05, 1987

THIOTHIXENE HYDROCHLORIDE

CONCENTRATE; ORAL

NAVANE

<u>AA</u>	+ PFIZER	<u>EQ 5MG BASE/ML</u>	<u>N16758</u>	<u>001</u>	
		<u>THIOTHIXENE HYDROCHLORIDE</u>			
<u>AA</u>	TEVA PHARMS	<u>EQ 5MG BASE/ML</u>	<u>N71554</u>	<u>001</u>	Oct 16, 1987

INJECTABLE; INJECTION

NAVANE

+ PFIZER	EQ 10MG BASE/VIAL	N16904	002	
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THYROTROPIN ALFA

INJECTABLE; INJECTION

THYROGEN

+ GENZYME	1.1MG/VIAL	N20898	001	Nov 30, 1998
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TIAGABINE HYDROCHLORIDE

TABLET; ORAL

GABITRIL

+ CEPHALON	2MG	N20646	005	Apr 16, 1999
	4MG	N20646	001	Sep 30, 1997
	6MG	N20646	006	Nov 29, 2005
	8MG	N20646	007	Nov 29, 2005
	10MG	N20646	008	Nov 29, 2005
	12MG	N20646	002	Sep 30, 1997
	16MG	N20646	003	Sep 30, 1997

TICLOPIDINE HYDROCHLORIDE

TABLET; ORAL

TICLID

<u>AB</u>	+ ROCHE PALO	<u>250MG</u>	<u>N19979</u>	<u>002</u>	Oct 31, 1991
		<u>TICLOPIDINE HYDROCHLORIDE</u>			
<u>AB</u>	ACTAVIS ELIZABETH	<u>250MG</u>	<u>N75253</u>	<u>001</u>	Aug 20, 1999
<u>AB</u>	CARACO	<u>250MG</u>	<u>N75526</u>	<u>001</u>	Sep 26, 2002
<u>AB</u>	GENPHARM	<u>250MG</u>	<u>N75161</u>	<u>001</u>	Sep 13, 1999
<u>AB</u>	MYLAN	<u>250MG</u>	<u>N75316</u>	<u>001</u>	Nov 02, 1999
<u>AB</u>	SANDOZ	<u>250MG</u>	<u>N75318</u>	<u>001</u>	Aug 20, 1999
<u>AB</u>		<u>250MG</u>	<u>N75326</u>	<u>001</u>	Aug 20, 1999
<u>AB</u>	TEVA	<u>250MG</u>	<u>N75149</u>	<u>001</u>	Aug 20, 1999
<u>AB</u>	TORPHARM	<u>250MG</u>	<u>N75089</u>	<u>001</u>	Jul 01, 1999

TIGECYCLINE

INJECTABLE; IV (INFUSION)

+ WYETH PHARMS INC	50MG/VIAL	N21821	001	Jun 15, 2005
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27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 349 (of 370)

TILUDRONATE DISODIUM

TABLET; ORAL

SKELID

+	SANOFI AVENTIS US	EQ 200MG BASE	N20707	001	Mar 07, 1997
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TIMOLOL

SOLUTION/DROPS; OPHTHALMIC

BETIMOL

+	SANTEN OY	EQ 0.25% BASE	N20439	001	Mar 31, 1995
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+		EQ 0.5% BASE	N20439	002	Mar 31, 1995
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TIMOLOL MALEATE

SOLUTION, GEL FORMING/DROPS; OPHTHALMIC

TIMOLOL MALEATE

<u>AB</u>	FALCON PHARMS	<u>EQ 0.25% BASE</u>	<u>N20963</u>	<u>001</u>	Oct 21, 1998
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<u>AB</u>		<u>EQ 0.5% BASE</u>	<u>N20963</u>	<u>002</u>	Oct 21, 1998
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TIMOPTIC-XE

<u>AB</u>	+	MERCK	<u>EQ 0.25% BASE</u>	<u>N20330</u>	<u>001</u>	Nov 04, 1993
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<u>AB</u>	+		<u>EQ 0.5% BASE</u>	<u>N20330</u>	<u>002</u>	Nov 04, 1993
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SOLUTION/DROPS; OPHTHALMIC

BT	+	ISTA PHARMS	EQ 0.5% BASE	N21516	001	Jun 04, 2004
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TIMOLOL MALEATE

<u>AT</u>		AKORN	<u>EQ 0.25% BASE</u>	<u>N74515</u>	<u>001</u>	Mar 25, 1997
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<u>AT</u>			<u>EQ 0.5% BASE</u>	<u>N74466</u>	<u>001</u>	Mar 25, 1997
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<u>AT</u>			<u>EQ 0.5% BASE</u>	<u>N74516</u>	<u>001</u>	Mar 25, 1997
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<u>AT</u>		BAUSCH AND LOMB	<u>EQ 0.25% BASE</u>	<u>N74778</u>	<u>001</u>	Mar 25, 1997
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<u>AT</u>			<u>EQ 0.5% BASE</u>	<u>N74776</u>	<u>001</u>	Mar 25, 1997
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<u>AT</u>		FALCON PHARMS	<u>EQ 0.25% BASE</u>	<u>N74261</u>	<u>001</u>	Apr 28, 1995
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<u>AT</u>			<u>EQ 0.5% BASE</u>	<u>N74262</u>	<u>001</u>	Apr 28, 1995
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<u>AT</u>		HI TECH PHARMA	<u>EQ 0.5% BASE</u>	<u>N75163</u>	<u>001</u>	Sep 10, 2002
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<u>AT</u>		NOVEX	<u>EQ 0.25% BASE</u>	<u>N75411</u>	<u>001</u>	Sep 08, 2000
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<u>AT</u>			<u>EQ 0.5% BASE</u>	<u>N75412</u>	<u>001</u>	Sep 08, 2000
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<u>AT</u>		PACIFIC PHARMA	<u>EQ 0.25% BASE</u>	<u>N74746</u>	<u>001</u>	Mar 25, 1997
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<u>AT</u>			<u>EQ 0.5% BASE</u>	<u>N74747</u>	<u>001</u>	Mar 25, 1997
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TIMOPTIC

<u>AT</u>	+	MERCK	<u>EQ 0.25% BASE</u>	<u>N18086</u>	<u>001</u>	
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<u>AT</u>	+		<u>EQ 0.5% BASE</u>	<u>N18086</u>	<u>002</u>	
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TIMOPTIC IN OCUDOSE

+	MERCK	EQ 0.25% BASE	N19463	001	Nov 05, 1986
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+		EQ 0.5% BASE	N19463	002	Nov 05, 1986
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TABLET; ORAL

TIMOLOL MALEATE

<u>AB</u>		MYLAN	<u>5MG</u>	<u>N72666</u>	<u>001</u>	Jun 08, 1990
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<u>AB</u>			<u>10MG</u>	<u>N72667</u>	<u>001</u>	Jun 08, 1990
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<u>AB</u>	+		<u>20MG</u>	<u>N72668</u>	<u>001</u>	Jun 08, 1990
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<u>AB</u>		SANDOZ	<u>5MG</u>	<u>N72550</u>	<u>001</u>	Apr 13, 1989
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<u>AB</u>			<u>10MG</u>	<u>N72551</u>	<u>001</u>	Apr 13, 1989
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<u>AB</u>			<u>20MG</u>	<u>N72552</u>	<u>001</u>	Apr 13, 1989
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<u>AB</u>		WATSON LABS	<u>5MG</u>	<u>N72917</u>	<u>001</u>	Jul 31, 1991
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<u>AB</u>			<u>10MG</u>	<u>N72918</u>	<u>001</u>	Jul 31, 1991
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<u>AB</u>			<u>20MG</u>	<u>N72919</u>	<u>001</u>	Jul 31, 1991
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TINIDAZOLE

TABLET; ORAL

TINDAMAX

	MISSION PHARMA	250MG	N21618	001	May 17, 2004
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+		500MG	N21618	002	May 17, 2004
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27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 350 (of 370)

TINZAPARIN SODIUM

INJECTABLE; INJECTION

INNOHEP

+ PHARMION

20,000 IU/ML

N20484 001

Jul 14, 2000

TIOPRONIN

TABLET; ORAL

TIOPRONIN

+ MISSION PHARMA

100MG

N19569 001

Aug 11, 1988

TIOTROPIUM BROMIDE MONOHYDRATE

POWDER; INHALATION

SPIRIVA

+ BOEHRINGER INGELHEIM EQ 0.018 BASE/INH

N21395 001

Jan 30, 2004

TIPRANAVIR

CAPSULE; ORAL

APTIVUS

+ BOEHRINGER INGELHEIM 250MG

N21814 001

Jun 22, 2005

TIROFIBAN HYDROCHLORIDE

INJECTABLE; INJECTION

AGGRASTAT

+ MEDICURE

EQ 0.05MG BASE/ML

N20913 001

May 14, 1998

+

EQ 0.25MG BASE/ML

N20912 001

May 14, 1998

TIZANIDINE HYDROCHLORIDE

CAPSULE; ORAL

ZANAFLEX

ACORDA

EQ 2MG BASE

N21447 001

Aug 29, 2002

EQ 4MG BASE

N21447 002

Aug 29, 2002

+

EQ 6MG BASE

N21447 003

Aug 29, 2002

TABLET; ORAL

TIZANIDINE HYDROCHLORIDE

<u>AB</u>	ACTAVIS ELIZABETH	<u>EQ 2MG BASE</u>	<u>N76283</u>	<u>001</u>	Jul 12, 2002
<u>AB</u>		<u>EQ 4MG BASE</u>	<u>N76283</u>	<u>002</u>	Jul 12, 2002
<u>AB</u>	ACTAVIS TOTOWA	<u>EQ 2MG BASE</u>	<u>N76281</u>	<u>001</u>	Oct 20, 2003
<u>AB</u>		<u>EQ 4MG BASE</u>	<u>N76281</u>	<u>002</u>	Oct 20, 2003
<u>AB</u>	ALPHAPHARM	<u>EQ 2MG BASE</u>	<u>N76282</u>	<u>001</u>	Dec 16, 2003
<u>AB</u>		<u>EQ 4MG BASE</u>	<u>N76282</u>	<u>002</u>	Dec 16, 2003
<u>AB</u>	BARR	<u>EQ 2MG BASE</u>	<u>N76371</u>	<u>001</u>	Apr 09, 2003
<u>AB</u>		<u>EQ 4MG BASE</u>	<u>N76371</u>	<u>002</u>	Apr 09, 2003
<u>AB</u>	CARACO	<u>EQ 2MG BASE</u>	<u>N76416</u>	<u>001</u>	Sep 29, 2003
<u>AB</u>		<u>EQ 4MG BASE</u>	<u>N76416</u>	<u>002</u>	Sep 29, 2003
<u>AB</u>	COREPHARMA	<u>EQ 2MG BASE</u>	<u>N76347</u>	<u>001</u>	Oct 11, 2002
<u>AB</u>		<u>EQ 4MG BASE</u>	<u>N76347</u>	<u>002</u>	Oct 11, 2002
<u>AB</u>	DR REDDYS LABS INC	<u>EQ 2MG BASE</u>	<u>N76286</u>	<u>001</u>	Jul 03, 2002
<u>AB</u>		<u>EQ 4MG BASE</u>	<u>N76286</u>	<u>002</u>	Jul 03, 2002
<u>AB</u>	IVAX PHARMS	<u>EQ 2MG BASE</u>	<u>N76321</u>	<u>001</u>	Sep 30, 2004
<u>AB</u>		<u>EQ 4MG BASE</u>	<u>N76321</u>	<u>002</u>	Sep 30, 2004
<u>AB</u>	MYLAN	<u>EQ 2MG BASE</u>	<u>N76354</u>	<u>001</u>	Mar 28, 2003
<u>AB</u>		<u>EQ 4MG BASE</u>	<u>N76354</u>	<u>002</u>	Mar 28, 2003
<u>AB</u>	SANDOZ	<u>EQ 2MG BASE</u>	<u>N76399</u>	<u>001</u>	Nov 26, 2002
<u>AB</u>		<u>EQ 4MG BASE</u>	<u>N76280</u>	<u>002</u>	Jun 27, 2002
<u>AB</u>	TEVA	<u>EQ 2MG BASE</u>	<u>N76284</u>	<u>001</u>	Jul 03, 2002
<u>AB</u>		<u>EQ 4MG BASE</u>	<u>N76284</u>	<u>002</u>	Jul 03, 2002
<u>AB</u>	TORPHARM	<u>EQ 2MG BASE</u>	<u>N76533</u>	<u>001</u>	Jan 16, 2004
<u>AB</u>		<u>EQ 4MG BASE</u>	<u>N76533</u>	<u>002</u>	Jan 16, 2004

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 351 (of 370)

TIZANIDINE HYDROCHLORIDE

TABLET; ORAL

ZANAFLEX

<u>AB</u>	ACORDA	<u>EQ 2MG BASE</u>	<u>N20397</u>	<u>002</u>	Feb 04, 2000
<u>AB</u>	+	<u>EQ 4MG BASE</u>	<u>N20397</u>	<u>001</u>	Nov 27, 1996

TOBRAMYCIN

OINTMENT; OPHTHALMIC

TOBREX

+	ALCON	0.3%	N50555	001	
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SOLUTION; INHALATION

TOBI

+	NOVARTIS PHARMS	300MG/5ML	N50753	001	Dec 22, 1997
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SOLUTION/DROPS; OPHTHALMIC

AKTOB

<u>AT</u>	AKORN	<u>0.3%</u>	<u>N64096</u>	<u>001</u>	Jan 31, 1996
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TOBRAMYCIN

<u>AT</u>	ALTANA	<u>0.3%</u>	<u>N65026</u>	<u>001</u>	Sep 11, 2001
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<u>AT</u>	BAUSCH AND LOMB	<u>0.3%</u>	<u>N64052</u>	<u>001</u>	Nov 29, 1993
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<u>AT</u>	NOVEX	<u>0.3%</u>	<u>N65087</u>	<u>001</u>	Feb 25, 2002
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TOBREX

<u>AT</u>	ALCON	<u>0.3%</u>	<u>N62535</u>	<u>001</u>	Dec 13, 1984
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<u>AT</u>	+	FALCON PHARMS	<u>0.3%</u>	<u>N50541</u>	<u>001</u>
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TOBRAMYCIN SULFATE

INJECTABLE; INJECTION

TOBRAMYCIN

<u>AP</u>	ABRAXIS PHARM	<u>EQ 10MG BASE/ML</u>	<u>N65122</u>	<u>001</u>	Nov 29, 2002
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<u>AP</u>	+	<u>EQ 40MG BASE/ML</u>	<u>N65122</u>	<u>002</u>	Nov 29, 2002
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		EQ 1.2GM BASE/VIAL	N50789	001	Jul 13, 2004
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+	X GEN PHARMS	EQ 1.2GM BASE/VIAL	N65013	001	Aug 17, 2001
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TOBRAMYCIN (PHARMACY BULK)

<u>AP</u>	ABRAXIS PHARM	<u>EQ 40MG BASE/ML</u>	<u>N65120</u>	<u>001</u>	Nov 29, 2002
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TOBRAMYCIN SULFATE

<u>AP</u>	APOTHECON	<u>EQ 10MG BASE/ML</u>	<u>N64021</u>	<u>001</u>	May 31, 1994
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<u>AP</u>		<u>EQ 40MG BASE/ML</u>	<u>N64021</u>	<u>002</u>	May 31, 1994
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<u>AP</u>	BAXTER HLTHCARE	<u>EQ 40MG BASE/ML</u>	<u>N63117</u>	<u>001</u>	Apr 26, 1991
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<u>AP</u>		<u>EQ 40MG BASE/ML</u>	<u>N63118</u>	<u>001</u>	Jul 29, 1991
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<u>AP</u>	+	HOSPIRA	<u>EQ 10MG BASE/ML</u>	<u>N63080</u>	<u>001</u>
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<u>AP</u>		<u>EQ 10MG BASE/ML</u>	<u>N63112</u>	<u>001</u>	Apr 30, 1991
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<u>AP</u>		<u>EQ 40MG BASE/ML</u>	<u>N63111</u>	<u>001</u>	Apr 30, 1991
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<u>AP</u>	+	<u>EQ 40MG BASE/ML</u>	<u>N63116</u>	<u>001</u>	May 18, 1992
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<u>AP</u>		<u>EQ 40MG BASE/ML</u>	<u>N63161</u>	<u>001</u>	May 29, 1991
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<u>AP</u>	MARSAM PHARMS LLC	<u>EQ 10MG BASE/ML</u>	<u>N62945</u>	<u>001</u>	Aug 09, 1989
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<u>AP</u>		<u>EQ 40MG BASE/ML</u>	<u>N62945</u>	<u>002</u>	Aug 09, 1989
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<u>AP</u>	SICOR PHARMS	<u>EQ 40MG BASE/ML</u>	<u>N63100</u>	<u>001</u>	Jan 30, 1992
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TOBRAMYCIN SULFATE IN SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER

+	HOSPIRA	EQ 1.2MG BASE/ML	N63081	003	Jul 31, 1990
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+		EQ 1.6MG BASE/ML	N63081	006	Jun 02, 1993
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+		EQ 80MG BASE/100ML	N63081	001	Jul 31, 1990
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TOLAZAMIDE

TABLET; ORAL

TOLAZAMIDE

<u>AB</u>	IVAX PHARMS	<u>100MG</u>	<u>N18894</u>	<u>001</u>	Nov 02, 1984
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<u>AB</u>		<u>250MG</u>	<u>N18894</u>	<u>002</u>	Nov 02, 1984
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<u>AB</u>		<u>500MG</u>	<u>N18894</u>	<u>003</u>	Nov 02, 1984
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<u>AB</u>	MUTUAL PHARM	<u>100MG</u>	<u>N71357</u>	<u>001</u>	Jul 16, 1987
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<u>AB</u>		<u>250MG</u>	<u>N71358</u>	<u>001</u>	Jul 16, 1987
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27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 352 (of 370)

TOLAZAMIDE

TABLET; ORAL

TOLAZAMIDE

<u>AB</u>	MUTUAL PHARM	<u>500MG</u>	<u>N71359</u>	<u>001</u>	Jul 16, 1987
<u>AB</u>	MYLAN	<u>250MG</u>	<u>N70259</u>	<u>001</u>	Jan 02, 1986
<u>AB</u>		<u>500MG</u>	<u>N70913</u>	<u>001</u>	Mar 17, 1986
<u>AB</u>	PAR PHARM	<u>100MG</u>	<u>N70159</u>	<u>001</u>	Jan 06, 1986
<u>AB</u>		<u>250MG</u>	<u>N70160</u>	<u>001</u>	Jan 06, 1986
<u>AB</u>		<u>500MG</u>	<u>N70161</u>	<u>001</u>	Jan 06, 1986
<u>AB</u>	SANDOZ	<u>100MG</u>	<u>N71633</u>	<u>001</u>	Dec 09, 1987
<u>AB</u>		<u>250MG</u>	<u>N70289</u>	<u>001</u>	Mar 13, 1986
<u>AB</u>		<u>500MG</u>	<u>N70290</u>	<u>001</u>	Mar 13, 1986
<u>AB</u>	WATSON LABS	<u>250MG</u>	<u>N70514</u>	<u>001</u>	Jan 09, 1986
<u>AB</u>		<u>500MG</u>	<u>N70515</u>	<u>001</u>	Jan 09, 1986
<u>TOLINASE</u>					
<u>AB</u>	PHARMACIA AND UPJOHN	<u>100MG</u>	<u>N15500</u>	<u>002</u>	
<u>AB</u>		<u>250MG</u>	<u>N15500</u>	<u>004</u>	
<u>AB</u>	+	<u>500MG</u>	<u>N15500</u>	<u>005</u>	

TOLBUTAMIDE

TABLET; ORAL

TOLBUTAMIDE

<u>AB</u>	IVAX PHARMS	<u>500MG</u>	<u>N87093</u>	<u>001</u>	
<u>AB</u>	MYLAN	<u>500MG</u>	<u>N86445</u>	<u>001</u>	
<u>AB</u>	SANDOZ	<u>500MG</u>	<u>N86574</u>	<u>001</u>	
<u>AB</u>	+	<u>500MG</u>	<u>N86109</u>	<u>001</u>	
<u>AB</u>	WATSON LABS	<u>500MG</u>	<u>N87318</u>	<u>001</u>	

TOLCAPONE

TABLET; ORAL

TASMAR

	VALEANT	100MG	N20697	001	Jan 29, 1998
	+	200MG	N20697	002	Jan 29, 1998

TOLMETIN SODIUM

CAPSULE; ORAL

TOLECTIN DS

<u>AB</u>	+	ORTHO MCNEIL PHARM	<u>EQ 400MG BASE</u>	<u>N18084</u>	<u>001</u>
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TOLMETIN SODIUM

<u>AB</u>	ACTAVIS ELIZABETH	<u>EQ 400MG BASE</u>	<u>N73308</u>	<u>001</u>	Jan 24, 1992
<u>AB</u>	IVAX PHARMS	<u>EQ 400MG BASE</u>	<u>N73392</u>	<u>001</u>	Jan 24, 1992
<u>AB</u>	MUTUAL PHARM	<u>EQ 400MG BASE</u>	<u>N73311</u>	<u>001</u>	Nov 27, 1991
<u>AB</u>	MYLAN	<u>EQ 400MG BASE</u>	<u>N73393</u>	<u>001</u>	May 27, 1993
<u>AB</u>	TEVA	<u>EQ 400MG BASE</u>	<u>N73290</u>	<u>001</u>	Nov 27, 1991

TABLET; ORAL

TOLECTIN

<u>AB</u>	ORTHO MCNEIL PHARM	<u>EQ 200MG BASE</u>	<u>N17628</u>	<u>001</u>	
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TOLECTIN 600

<u>AB</u>	+	ORTHO MCNEIL PHARM	<u>EQ 600MG BASE</u>	<u>N17628</u>	<u>002</u> Mar 08, 1989
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TOLMETIN SODIUM

<u>AB</u>	ACTAVIS ELIZABETH	<u>EQ 600MG BASE</u>	<u>N73527</u>	<u>001</u>	Jun 30, 1992
<u>AB</u>	IVAX PHARMS	<u>EQ 600MG BASE</u>	<u>N74399</u>	<u>001</u>	Mar 28, 1996
<u>AB</u>	MUTUAL PHARM	<u>EQ 200MG BASE</u>	<u>N73310</u>	<u>001</u>	Nov 27, 1991
<u>AB</u>	MYLAN	<u>EQ 600MG BASE</u>	<u>N74473</u>	<u>001</u>	Aug 30, 1994
<u>AB</u>	SANDOZ	<u>EQ 200MG BASE</u>	<u>N73588</u>	<u>001</u>	Jul 31, 1992
<u>AB</u>		<u>EQ 600MG BASE</u>	<u>N74002</u>	<u>001</u>	Sep 27, 1993

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 353 (of 370)

TOLTERODINE TARTRATE

CAPSULE, EXTENDED RELEASE; ORAL

DETROL LA

PHARMACIA AND UPJOHN 2MG

N21228 001 Dec 22, 2000

+ 4MG

N21228 002 Dec 22, 2000

TABLET; ORAL

DETROL

PHARMACIA AND UPJOHN 1MG

N20771 001 Mar 25, 1998

+ 2MG

N20771 002 Mar 25, 1998

TOPIRAMATE

CAPSULE; ORAL

TOPAMAX SPRINKLE

ORTHO MCNEIL PHARM 15MG

N20844 001 Oct 26, 1998

+ 25MG

N20844 002 Oct 26, 1998

TABLET; ORAL

TOPAMAXAB + ORTHO MCNEIL PHARM25MGN20505 004 Dec 24, 1996AB 100MGN20505 001 Dec 24, 1996AB 200MGN20505 002 Dec 24, 1996

50MG

N20505 005 Dec 24, 1996

TOPIRAMATEAB MYLAN25MGN76314 001 Sep 11, 2006AB 100MGN76314 002 Sep 11, 2006AB 200MGN76314 003 Sep 11, 2006TOPOTECAN HYDROCHLORIDE

INJECTABLE; INJECTION

HYCAMTIN

+ GLAXOSMITHKLINE EQ 4MG BASE/VIAL

N20671 001 May 28, 1996

TOREMIFENE CITRATE

TABLET; ORAL

FARESTON

+ GTX INC EQ 60MG BASE

N20497 001 May 29, 1997

TORSEMIDE

INJECTABLE; INJECTION

DEMADEX

+ ROCHE 10MG/ML

N20137 002 Aug 23, 1993

TABLET; ORAL

DEMADEXAB ROCHE5MGN20136 001 Aug 23, 1993AB 10MGN20136 002 Aug 23, 1993AB + 20MGN20136 003 Aug 23, 1993AB 100MGN20136 004 Aug 23, 1993TORSEMIDEAB APOTEX INC5MGN76894 001 May 31, 2005AB 10MGN76894 002 May 31, 2005AB 20MGN76894 003 May 31, 2005AB 100MGN76894 004 May 31, 2005AB PAR PHARM5MGN76226 001 May 27, 2003AB 10MGN76226 002 May 27, 2003AB 20MGN76226 003 May 27, 2003AB 100MGN76226 004 May 27, 2003AB PLIVA PHARM IND5MGN76346 001 May 30, 2003AB 10MGN76346 002 May 30, 2003AB 20MGN76346 003 May 30, 2003AB 100MGN76346 004 Oct 19, 2004

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 354 (of 370)

TORSEMIDE

TABLET; ORAL

TORSEMIDE

<u>AB</u>	ROXANE	<u>5MG</u>	<u>N76943</u>	<u>001</u>	Mar 01, 2005
<u>AB</u>		<u>10MG</u>	<u>N76943</u>	<u>002</u>	Mar 01, 2005
<u>AB</u>		<u>20MG</u>	<u>N76943</u>	<u>003</u>	Mar 01, 2005
<u>AB</u>	TEVA	<u>5MG</u>	<u>N76110</u>	<u>001</u>	May 14, 2002
<u>AB</u>		<u>10MG</u>	<u>N76110</u>	<u>002</u>	May 14, 2002
<u>AB</u>		<u>20MG</u>	<u>N76110</u>	<u>003</u>	May 14, 2002
<u>AB</u>		<u>100MG</u>	<u>N76110</u>	<u>004</u>	May 14, 2002

TRAMADOL HYDROCHLORIDE

TABLET; ORAL

TRAMADOL HYDROCHLORIDE

<u>AB</u>	ACTAVIS ELIZABETH	<u>50MG</u>	<u>N75960</u>	<u>001</u>	Jun 19, 2002
<u>AB</u>	ALPHAPHARM	<u>50MG</u>	<u>N75980</u>	<u>001</u>	Nov 21, 2002
<u>AB</u>	CARACO	<u>50MG</u>	<u>N75964</u>	<u>001</u>	Jun 19, 2002
<u>AB</u>	COREPHARMA	<u>50MG</u>	<u>N76003</u>	<u>001</u>	Jun 20, 2002
<u>AB</u>	MALLINCKRODT	<u>50MG</u>	<u>N75983</u>	<u>001</u>	Jun 25, 2002
<u>AB</u>	MUTUAL PHARM	<u>50MG</u>	<u>N76100</u>	<u>001</u>	Jun 20, 2002
<u>AB</u>	MYLAN	<u>50MG</u>	<u>N75986</u>	<u>001</u>	Jun 21, 2002
<u>AB</u>	PLIVA	<u>50MG</u>	<u>N75982</u>	<u>001</u>	Jul 01, 2002
<u>AB</u>	SANDOZ	<u>50MG</u>	<u>N75968</u>	<u>001</u>	Jun 25, 2002
<u>AB</u>	TEVA	<u>50MG</u>	<u>N75977</u>	<u>001</u>	Jun 19, 2002
<u>AB</u>	TORPHARM	<u>50MG</u>	<u>N75981</u>	<u>001</u>	Jul 10, 2002
<u>AB</u>	WATSON LABS	<u>50MG</u>	<u>N75962</u>	<u>001</u>	Jun 24, 2002

ULTRAM

<u>AB</u>	+ ORTHO MCNEIL PHARM	<u>50MG</u>	<u>N20281</u>	<u>002</u>	Mar 03, 1995
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TABLET, EXTENDED RELEASE; ORAL

TRAMADOL HYDROCHLORIDE

	BIOVAIL LABS INTL	100MG	N21692	001	Sep 08, 2005
		200MG	N21692	002	Sep 08, 2005
+		300MG	N21692	003	Sep 08, 2005

TABLET, ORALLY DISINTEGRATING; ORAL

TRAMADOL HYDROCHLORIDE

+	BIOVAIL	50MG	N21693	001	May 05, 2005
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TRANDOLAPRIL

TABLET; ORAL

MAVIK

<u>AB</u>	ABBOTT	<u>1MG</u>	<u>N20528</u>	<u>001</u>	Apr 26, 1996
<u>AB</u>		<u>2MG</u>	<u>N20528</u>	<u>002</u>	Apr 26, 1996
<u>AB</u>	+	<u>4MG</u>	<u>N20528</u>	<u>003</u>	Apr 26, 1996

TRANDOLAPRIL

<u>AB</u>	TEVA PHARMS	<u>1MG</u>	<u>N77489</u>	<u>001</u>	Dec 12, 2006
<u>AB</u>		<u>2MG</u>	<u>N77489</u>	<u>002</u>	Dec 12, 2006
<u>AB</u>		<u>4MG</u>	<u>N77489</u>	<u>003</u>	Dec 12, 2006

TRANDOLAPRIL; VERAPAMIL HYDROCHLORIDE

TABLET, EXTENDED RELEASE; ORAL

TARKA

+	ABBOTT	1MG; 240MG	N20591	003	Oct 22, 1996
+		2MG; 180MG	N20591	001	Oct 22, 1996
+		2MG; 240MG	N20591	004	Oct 22, 1996
+		4MG; 240MG	N20591	002	Oct 22, 1996

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 355 (of 370)

TRANEXAMIC ACID

INJECTABLE; INJECTION

CYKLOKAPRON

+ PHARMACIA AND UPJOHN 100MG/ML

N19281 001 Dec 30, 1986

TRANLYCYPROMINE SULFATE

TABLET; ORAL

PARNATEAB + GLAXOSMITHKLINE EQ 10MG BASEN12342 003 Aug 16, 1985TRANLYCYPROMINE SULFATEAB KALI LABS EQ 10MG BASEN40640 001 Jun 29, 2006TRAVOPROST

SOLUTION/DROPS; OPHTHALMIC

TRAVATAN

+ ALCON 0.004%

N21257 001 Mar 16, 2001

TRAVATAN Z

+ ALCON 0.004%

N21994 001 Sep 21, 2006

TRAZODONE HYDROCHLORIDE

TABLET; ORAL

TRAZODONE HYDROCHLORIDEAB ACTAVIS ELIZABETH 50MG N71636 001 Apr 18, 1988AB 100MG N71514 001 Apr 18, 1988AB APOTEX INC 100MG N71196 001 Mar 25, 1987AB 150MG N71196 002 Apr 26, 1999AB + 300MG N71196 003 Apr 26, 1999AB BARR 50MG N71258 001 Mar 25, 1987AB MUTUAL PHARM 50MG N73136 001 Mar 24, 1993AB 100MG N73137 001 Mar 24, 1993AB 150MG N73138 001 Dec 22, 1995AB MYLAN 50MG N71405 001 Feb 27, 1991AB PLIVA 50MG N71523 001 Dec 11, 1987AB 100MG N71524 001 Dec 11, 1987AB 150MG N71525 001 Mar 09, 1988AB SANDOZ 50MG N72484 001 Apr 30, 1990AB 100MG N72483 001 Apr 30, 1990AB TEVA 50MG N72192 001 Feb 02, 1989AB 100MG N72193 001 Feb 02, 1989AB 150MG N74357 001 Apr 30, 1997AB WATSON LABS 50MG N70857 001 Oct 10, 1986AB 100MG N70858 001 Oct 10, 1986TREPROSTINIL SODIUM

INJECTABLE; IV (INFUSION)-SC

UNITED THERAP 1MG/ML

N21272 001 May 21, 2002

2.5MG/ML

N21272 002 May 21, 2002

5MG/ML

N21272 003 May 21, 2002

+ 10MG/ML

N21272 004 May 21, 2002

TRETINOIN

CAPSULE; ORAL

VESANOID

+ ROCHE 10MG

N20438 001 Nov 22, 1995

CREAM; TOPICAL

AVITAAB MYLAN BERTEK 0.025%N20404 003 Jan 14, 1997

PRESCRIPTION DRUG PRODUCT LIST

3 - 356 (of 370)

TRETINOIN

CREAM; TOPICAL				
<u>RENOVA</u>				
<u>AB2</u> +	JOHNSON AND JOHNSON	<u>0.05%</u>	<u>N19963</u>	<u>001</u> Dec 29, 1995
+		<u>0.02%</u>	<u>N21108</u>	<u>001</u> Aug 31, 2000
<u>RETIN-A</u>				
<u>AB</u> +	JOHNSON AND JOHNSON	<u>0.025%</u>	<u>N19049</u>	<u>001</u> Sep 16, 1988
<u>AB</u> +		<u>0.1%</u>	<u>N17340</u>	<u>001</u>
<u>AB1</u> +		<u>0.05%</u>	<u>N17522</u>	<u>001</u>
<u>TRETINOIN</u>				
<u>AB2</u>	SPEAR PHARMS	<u>0.05%</u>	<u>N76498</u>	<u>001</u> Sep 15, 2005
<u>AB</u>	TRIAx PHARMS LLC	<u>0.025%</u>	<u>N75264</u>	<u>001</u> Dec 24, 1998
<u>AB</u>		<u>0.1%</u>	<u>N75213</u>	<u>001</u> Dec 24, 1998
<u>AB1</u>		<u>0.05%</u>	<u>N75265</u>	<u>001</u> Dec 24, 1998
GEL; TOPICAL				
AVITA				
BT	MYLAN BERTEK	0.025%	N20400	001 Jan 29, 1998
<u>RETIN-A</u>				
<u>AB</u> +	JOHNSON AND JOHNSON	<u>0.01%</u>	<u>N17955</u>	<u>001</u>
<u>AB</u> +		<u>0.025%</u>	<u>N17579</u>	<u>002</u>
	RETIN-A MICRO			
+	JOHNSON AND JOHNSON	0.04%	N20475	002 May 10, 2002
+		0.1%	N20475	001 Feb 07, 1997
<u>TRETINOIN</u>				
<u>AB</u>	SPEAR PHARMS	<u>0.01%</u>	<u>N75589</u>	<u>001</u> Jun 11, 2002
<u>AB</u>		<u>0.025%</u>	<u>N75529</u>	<u>001</u> Feb 22, 2000
SOLUTION; TOPICAL				
<u>RETIN-A</u>				
<u>AT</u> +	JOHNSON AND JOHNSON	<u>0.05%</u>	<u>N16921</u>	<u>001</u>
<u>TRETINOIN</u>				
<u>AT</u>	MORTON GROVE	<u>0.05%</u>	<u>N75260</u>	<u>001</u> Jan 25, 1999
<u>AT</u>	TEVA PHARMS	<u>0.05%</u>	<u>N74873</u>	<u>001</u> Jun 19, 1998

TRIAMCINOLONE ACETONIDE

AEROSOL, METERED; INHALATION				
AZMACORT				
+	KOS LIFE	0.1MG/INH	N18117	001 Apr 23, 1982
CREAM; TOPICAL				
<u>KENALOG</u>				
<u>AT</u> +	APOTHECON	<u>0.025%</u>	<u>N11601</u>	<u>003</u>
<u>AT</u> +		<u>0.1%</u>	<u>N11601</u>	<u>006</u>
<u>AT</u> +		<u>0.5%</u>	<u>N83943</u>	<u>001</u>
<u>TRIA CET</u>				
<u>AT</u>	TEVA	<u>0.025%</u>	<u>N84908</u>	<u>001</u>
<u>AT</u>		<u>0.1%</u>	<u>N84908</u>	<u>002</u>
<u>AT</u>		<u>0.5%</u>	<u>N84908</u>	<u>003</u>
<u>TRIAMCINOLONE ACETONIDE</u>				
<u>AT</u>	ACTAVIS MID ATLANTIC	<u>0.1%</u>	<u>N87798</u>	<u>001</u> Jun 04, 1982
<u>AT</u>	ALTANA	<u>0.025%</u>	<u>N85692</u>	<u>001</u>
<u>AT</u>		<u>0.1%</u>	<u>N85692</u>	<u>003</u>
<u>AT</u>		<u>0.5%</u>	<u>N85692</u>	<u>002</u>
<u>AT</u>	G AND W LABS	<u>0.025%</u>	<u>N89797</u>	<u>001</u> May 31, 1991
<u>AT</u>		<u>0.1%</u>	<u>N89798</u>	<u>001</u> May 31, 1991
<u>AT</u>	PERRIGO NEW YORK	<u>0.025%</u>	<u>N86415</u>	<u>001</u>
<u>AT</u>		<u>0.1%</u>	<u>N86414</u>	<u>001</u>
<u>AT</u>		<u>0.5%</u>	<u>N86413</u>	<u>001</u>
<u>AT</u>	TARO	<u>0.025%</u>	<u>N86277</u>	<u>001</u>
<u>AT</u>		<u>0.1%</u>	<u>N40039</u>	<u>001</u> Nov 26, 1997
<u>AT</u>		<u>0.1%</u>	<u>N86276</u>	<u>001</u>
<u>AT</u>		<u>0.5%</u>	<u>N86275</u>	<u>001</u>

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 357 (of 370)

TRIAMCINOLONE ACETONIDE

CREAM; TOPICAL

TRIAMCINOLONE ACETONIDE

<u>AT</u>	VINTAGE	<u>0.025%</u>	<u>N40671</u>	<u>001</u>	Jun 09, 2006
<u>AT</u>		<u>0.1%</u>	<u>N40671</u>	<u>002</u>	Jun 09, 2006

TRIDERM

<u>AT</u>	DEL RAY LABS	<u>0.1%</u>	<u>N88042</u>	<u>001</u>	Mar 19, 1984
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INJECTABLE; INJECTION

KENALOG-10

APOTHECON

10MG/ML

N12041 001

KENALOG-40

+ APOTHECON

40MG/ML

N14901 001

LOTION; TOPICAL

KENALOG

<u>AT</u>	APOTHECON	<u>0.025%</u>	<u>N84343</u>	<u>001</u>	
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<u>AT</u>	+	<u>0.1%</u>	<u>N84343</u>	<u>002</u>	
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TRIAMCINOLONE ACETONIDE

<u>AT</u>	ALTANA	<u>0.025%</u>	<u>N40467</u>	<u>001</u>	Apr 21, 2003
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<u>AT</u>		<u>0.1%</u>	<u>N40467</u>	<u>002</u>	Apr 21, 2003
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<u>AT</u>	MORTON GROVE	<u>0.025%</u>	<u>N88450</u>	<u>001</u>	Apr 01, 1985
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<u>AT</u>		<u>0.1%</u>	<u>N88451</u>	<u>001</u>	Apr 03, 1985
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<u>AT</u>	TARO	<u>0.1%</u>	<u>N89129</u>	<u>001</u>	Aug 14, 1986
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<u>AT</u>	VINTAGE	<u>0.1%</u>	<u>N40672</u>	<u>002</u>	Dec 13, 2006
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OINTMENT; TOPICAL

KENALOG

<u>AT</u>	APOTHECON	<u>0.025%</u>	<u>N11600</u>	<u>003</u>	
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<u>AT</u>		<u>0.1%</u>	<u>N11600</u>	<u>001</u>	
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TRIAMCINOLONE ACETONIDE

<u>AT</u>	ACTAVIS MID ATLANTIC	<u>0.1%</u>	<u>N87799</u>	<u>001</u>	Jun 07, 1982
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<u>AT</u>	ALTANA	<u>0.025%</u>	<u>N85691</u>	<u>001</u>	
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<u>AT</u>		<u>0.1%</u>	<u>N85691</u>	<u>003</u>	
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<u>AT</u>		<u>0.5%</u>	<u>N85691</u>	<u>002</u>	
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<u>AT</u>	+	<u>0.025%</u>	<u>N87356</u>	<u>001</u>	
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<u>AT</u>	+	<u>0.1%</u>	<u>N87357</u>	<u>001</u>	
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<u>AT</u>	+	<u>0.5%</u>	<u>N87385</u>	<u>001</u>	
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<u>AT</u>	TARO	<u>0.025%</u>	<u>N40040</u>	<u>001</u>	Sep 30, 1994
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<u>AT</u>		<u>0.025%</u>	<u>N40374</u>	<u>001</u>	Jun 05, 2001
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<u>AT</u>		<u>0.1%</u>	<u>N40037</u>	<u>001</u>	Sep 30, 1994
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<u>AT</u>		<u>0.1%</u>	<u>N87902</u>	<u>001</u>	Dec 27, 1982
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<u>AT</u>		<u>0.5%</u>	<u>N40386</u>	<u>001</u>	Jun 05, 2001
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TRIAMCINOLONE ACETONIDE IN ABSORBASE

+ CAROLINA MEDCL

0.05%

N89595 001 Mar 23, 1995

PASTE; DENTAL

KENALOG IN ORABASE

<u>AT</u>	+	APOTHECON	<u>0.1%</u>	<u>N12097</u>	<u>001</u>
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ORACORT

<u>AT</u>	TARO	<u>0.1%</u>	<u>N70730</u>	<u>001</u>	Oct 01, 1986
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ORALONE

<u>AT</u>	TARO	<u>0.1%</u>	<u>N71383</u>	<u>001</u>	Jul 06, 1987
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SPRAY; TOPICAL

KENALOG

+ APOTHECON

0.147MG/GM

N12104 001

SPRAY, METERED; NASAL

NASACORT AQ

+ SANOFI AVENTIS US

0.055MG/SPRAY

N20468 001 May 20, 1996

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 358 (of 370)

TRIAMCINOLONE HEXACETONIDE

INJECTABLE; INJECTION

ARISTOSPAN

+	SANDOZ	5MG/ML	N16466	001
+		20MG/ML	N16466	002

TRIAMTERENE

CAPSULE; ORAL

DYRENIUM

	WELLSPRING PHARM	50MG	N13174	001
+		100MG	N13174	002

TRIAZOLAM

TABLET; ORAL

HALCION

<u>AB</u>	PHARMACIA AND UPJOHN	<u>0.125MG</u>	<u>N17892</u>	<u>003</u>	Apr 26, 1985
<u>AB</u>	+	<u>0.25MG</u>	<u>N17892</u>	<u>001</u>	Nov 15, 1982

TRIAZOLAM

<u>AB</u>	ALPHAPHARM	<u>0.125MG</u>	<u>N74031</u>	<u>001</u>	Mar 25, 1994
<u>AB</u>		<u>0.25MG</u>	<u>N74031</u>	<u>002</u>	Mar 25, 1994
<u>AB</u>	ROXANE	<u>0.125MG</u>	<u>N74224</u>	<u>001</u>	Jun 01, 1994
<u>AB</u>		<u>0.25MG</u>	<u>N74224</u>	<u>002</u>	Jun 01, 1994
<u>AB</u>	WATSON LABS	<u>0.125MG</u>	<u>N74445</u>	<u>001</u>	Oct 20, 1995
<u>AB</u>		<u>0.25MG</u>	<u>N74445</u>	<u>002</u>	Oct 20, 1995

TRIENTINE HYDROCHLORIDE

CAPSULE; ORAL

SYPRINE

+	ATON	250MG	N19194	001	Nov 08, 1985
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TRIFLUOPERAZINE HYDROCHLORIDE

CONCENTRATE; ORAL

STELAZINE

<u>AA</u>	+	GLAXOSMITHKLINE	<u>EQ 10MG BASE/ML</u>	<u>N11552</u>	<u>006</u>
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TABLET; ORAL

STELAZINE

<u>AB</u>	GLAXOSMITHKLINE	<u>EQ 1MG BASE</u>	<u>N11552</u>	<u>001</u>
<u>AB</u>		<u>EQ 2MG BASE</u>	<u>N11552</u>	<u>002</u>
<u>AB</u>		<u>EQ 5MG BASE</u>	<u>N11552</u>	<u>003</u>
<u>AB</u>	+	<u>EQ 10MG BASE</u>	<u>N11552</u>	<u>004</u>

TRIFLUOPERAZINE HYDROCHLORIDE

<u>AB</u>	MYLAN	<u>EQ 1MG BASE</u>	<u>N40209</u>	<u>001</u>	Jul 07, 1997
<u>AB</u>		<u>EQ 2MG BASE</u>	<u>N40209</u>	<u>002</u>	Jul 07, 1997
<u>AB</u>		<u>EQ 5MG BASE</u>	<u>N40209</u>	<u>003</u>	Jul 07, 1997
<u>AB</u>		<u>EQ 10MG BASE</u>	<u>N40209</u>	<u>004</u>	Jul 07, 1997
<u>AB</u>	SANDOZ	<u>EQ 1MG BASE</u>	<u>N40153</u>	<u>001</u>	Oct 25, 1996
<u>AB</u>		<u>EQ 1MG BASE</u>	<u>N85785</u>	<u>001</u>	
<u>AB</u>		<u>EQ 2MG BASE</u>	<u>N40153</u>	<u>002</u>	Oct 25, 1996
<u>AB</u>		<u>EQ 2MG BASE</u>	<u>N85786</u>	<u>001</u>	
<u>AB</u>		<u>EQ 5MG BASE</u>	<u>N40153</u>	<u>003</u>	Oct 25, 1996
<u>AB</u>		<u>EQ 5MG BASE</u>	<u>N85789</u>	<u>001</u>	
<u>AB</u>		<u>EQ 10MG BASE</u>	<u>N40153</u>	<u>004</u>	Oct 25, 1996
<u>AB</u>		<u>EQ 10MG BASE</u>	<u>N85788</u>	<u>001</u>	

TRIFLURIDINE

SOLUTION/DROPS; OPHTHALMIC

TRIFLURIDINE

<u>AT</u>	ALCON	<u>1%</u>	<u>N74311</u>	<u>001</u>	Oct 06, 1995
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PRESCRIPTION DRUG PRODUCT LIST

3 - 359 (of 370)

TRIFLURIDINE

SOLUTION/DROPS; OPHTHALMIC

VIROPTIC

<u>AT</u>	+	MONARCH PHARMS	<u>1%</u>	<u>N18299</u>	<u>001</u>
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TRIHEXYPHENIDYL HYDROCHLORIDE

ELIXIR; ORAL

TRIHEXYPHENIDYL HYDROCHLORIDE

<u>AA</u>	MIKART	<u>2MG/5ML</u>	<u>N40251</u>	<u>001</u>	Sep 27, 1999	
<u>AA</u>	PHARM ASSOC	<u>2MG/5ML</u>	<u>N40177</u>	<u>001</u>	Apr 17, 1997	
<u>AA</u>	+	PHARM VENTURES	<u>2MG/5ML</u>	<u>N89514</u>	<u>001</u>	Apr 07, 1989

TABLET; ORAL

TRIHEXYPHENIDYL HYDROCHLORIDE

<u>AA</u>	VINTAGE PHARMS	<u>2MG</u>	<u>N40254</u>	<u>001</u>	Dec 24, 1998
<u>AA</u>		<u>5MG</u>	<u>N40254</u>	<u>002</u>	Dec 24, 1998
<u>AA</u>	WATSON LABS	<u>2MG</u>	<u>N40184</u>	<u>001</u>	Feb 06, 1998
<u>AA</u>	+	<u>2MG</u>	<u>N84363</u>	<u>001</u>	
<u>AA</u>		<u>5MG</u>	<u>N40184</u>	<u>002</u>	Feb 06, 1998
<u>AA</u>	+	<u>5MG</u>	<u>N84364</u>	<u>001</u>	
<u>AA</u>	WEST WARD	<u>2MG</u>	<u>N40337</u>	<u>002</u>	Feb 16, 2000
<u>AA</u>		<u>5MG</u>	<u>N40337</u>	<u>001</u>	Feb 16, 2000

TRIMETHADIONE

TABLET; ORAL

TRIDIONE

+	ABBOTT	150MG	N05856	009
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TRIMETHOBENZAMIDE HYDROCHLORIDE

CAPSULE; ORAL

TIGAN

<u>AB</u>	+	KING PHARMS	<u>300MG</u>	<u>N17531</u>	<u>006</u>	Dec 13, 2001
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TRIMETHOBENZAMIDE HYDROCHLORIDE

<u>AB</u>	ACTAVIS TOTOWA	<u>300MG</u>	<u>N76546</u>	<u>001</u>	Aug 20, 2003
<u>AB</u>	MUTUAL PHARMA	<u>300MG</u>	<u>N76570</u>	<u>001</u>	Aug 28, 2003

INJECTABLE; INJECTION

TIGAN

<u>AP</u>	+	KING PHARMS	<u>100MG/ML</u>	<u>N17530</u>	<u>001</u>
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TRIMETHOBENZAMIDE HYDROCHLORIDE

<u>AP</u>	HOSPIRA	<u>100MG/ML</u>	<u>N88804</u>	<u>001</u>	Apr 03, 1987
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TRIMETHOPRIM

TABLET; ORAL

PROLOPRIM

<u>AB</u>	MONARCH PHARMS	<u>100MG</u>	<u>N17943</u>	<u>001</u>
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TRIMETHOPRIM

<u>AB</u>	TEVA	<u>100MG</u>	<u>N18679</u>	<u>001</u>	Jul 30, 1982
	+	200MG	N71259	001	Jun 18, 1987
<u>AB</u>	WATSON LABS	<u>100MG</u>	<u>N70049</u>	<u>001</u>	Jun 06, 1985

TRIMETHOPRIM HYDROCHLORIDE

SOLUTION; ORAL

PRIMSOL

+	FSC	EQ 50MG BASE/5ML	N74973	001	Jan 24, 2000
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TRIMETREXATE GLUCURONATE

INJECTABLE; INJECTION

NEUTREXIN

+	MEDIMMUNE ONCOLOGY	EQ 25MG BASE/VIAL	N20326	001	Dec 17, 1993
+		EQ 200MG BASE/VIAL	N20326	002	Jul 31, 1998

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 360 (of 370)

TRIMIPRAMINE MALEATE

CAPSULE; ORAL

SURMONTIL

<u>AB</u>	ODYSSEY PHARMS	<u>EQ 25MG BASE</u>	<u>N16792</u>	<u>001</u>	
<u>AB</u>		<u>EQ 50MG BASE</u>	<u>N16792</u>	<u>002</u>	
<u>AB</u>	+	<u>EQ 100MG BASE</u>	<u>N16792</u>	<u>003</u>	Sep 15, 1982
	<u>TRIMIPRAMINE MALEATE</u>				
<u>AB</u>	ACTAVIS TOTOWA	<u>EQ 25MG BASE</u>	<u>N77361</u>	<u>001</u>	Aug 02, 2006
<u>AB</u>		<u>EQ 50MG BASE</u>	<u>N77361</u>	<u>002</u>	Aug 02, 2006
<u>AB</u>		<u>EQ 100MG BASE</u>	<u>N77361</u>	<u>003</u>	Aug 02, 2006

TRIPTORELIN PAMOATE

INJECTABLE; INTRAMUSCULAR

TRELSTAR

+	WATSON LABS	11.25MG/VIAL	N21288	001	Jun 29, 2001
	TRELSTAR DEPOT				
+	WATSON LABS	EQ 3.75MG BASE/VIAL	N20715	001	Jun 15, 2000

TROMETHAMINE

INJECTABLE; INJECTION

THAM

+	HOSPIRA	3.6GM/100ML	N13025	002	
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TROPICAMIDE

SOLUTION/DROPS; OPHTHALMIC

MYDRIACYL

<u>AT</u>	+	ALCON	<u>0.5%</u>	<u>N84305</u>	<u>001</u>	
<u>AT</u>	+		<u>1%</u>	<u>N84306</u>	<u>001</u>	
	<u>TROPICACYL</u>					
<u>AT</u>		AKORN	<u>0.5%</u>	<u>N40314</u>	<u>001</u>	Sep 29, 2000
<u>AT</u>			<u>1%</u>	<u>N40315</u>	<u>001</u>	Sep 29, 2000
	<u>TROPICAMIDE</u>					
<u>AT</u>		ALCON UNIVERSAL	<u>1%</u>	<u>N89172</u>	<u>001</u>	Dec 28, 1990
<u>AT</u>		BAUSCH AND LOMB	<u>0.5%</u>	<u>N40067</u>	<u>001</u>	Jul 27, 1994
<u>AT</u>			<u>1%</u>	<u>N40064</u>	<u>001</u>	Jul 27, 1994
<u>AT</u>		MIZA PHARMS USA	<u>0.5%</u>	<u>N87636</u>	<u>001</u>	Jul 30, 1982
<u>AT</u>			<u>1%</u>	<u>N87637</u>	<u>001</u>	Aug 09, 1982

TROSPIUM CHLORIDE

TABLET; ORAL

SANCTURA

+	INDEVUS	20MG	N21595	001	May 28, 2004
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TRYPAN BLUE

SOLUTION; OPHTHALMIC

VISIONBLUE

+	DORC	0.06%	N21670	001	Dec 16, 2004
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UREA, C-13

FOR SOLUTION; ORAL

BREATHTEK UBT FOR H-PYLORI

+	MERETEK	EQ 75MG /POUCHE	N20586	002	May 10, 2001
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UREA, C-14

CAPSULE; ORAL

PYTEST

+	BALLARD MEDCL	1uCi	N20617	001	May 09, 1997
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PYTEST KIT

+	BALLARD MEDCL	1uCi	N20617	002	May 09, 1997
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27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 361 (of 370)

UROFOLLITROPIN

INJECTABLE; INTRAMUSCULAR, SUBCUTANEOUS

BRAVELLE

+ FERRING

75 IU/VIAL

N21289 001

May 06, 2002

UROKINASE

INJECTABLE; INJECTION

ABBOKINASE

+ IMARX THERAP

250,000 IU/VIAL

N21846 001

URSODIOL

CAPSULE; ORAL

ACTIGALLAB + WATSON PHARMS300MGN19594 002

Dec 31, 1987

URSODIOLAB ACTAVIS TOTOWA300MGN75517 001

Mar 14, 2000

AB COREPHARMA300MGN77895 001

Jul 27, 2006

AB TEVA PHARMS300MGN75592 001

May 25, 2000

TABLET; ORAL

URSO 250

AXCAN SCANDIPHARM

250MG

N20675 001

Dec 10, 1997

URSO FORTE

+ AXCAN SCANDIPHARM

500MG

N20675 002

Jul 21, 2004

VALACYCLOVIR HYDROCHLORIDE

TABLET; ORAL

VALTREX

GLAXOSMITHKLINE

EQ 500MG BASE

N20487 001

Jun 23, 1995

+

EQ 1GM BASE

N20487 002

Jun 23, 1995

VALDECOXIB

TABLET; ORAL

BEXTRA

GD SEARLE

10MG

N21341 002

Nov 16, 2001

+

20MG

N21341 003

Nov 16, 2001

VALGANCICLOVIR HYDROCHLORIDE

TABLET; ORAL

VALCYTE

+ ROCHE PALO

EQ 450MG BASE

N21304 001

Mar 29, 2001

VALPROATE SODIUM

INJECTABLE; INJECTION

DEPACONAP + ABBOTTEQ 100MG BASE/MLN20593 001

Dec 30, 1996

VALPROATE SODIUMAP ABRAXIS PHARMEQ 100MG BASE/MLN76539 001

Jun 26, 2003

AP BEDFORDEQ 100MG BASE/MLN76295 001

Nov 14, 2002

VALPROIC ACID

CAPSULE; ORAL

DEPAKENEAB + ABBOTT250MGN18081 001VALPROIC ACIDAB BANNER PHARMACAPS250MGN73484 001

Jun 29, 1993

AB SCHERER RP250MGN73229 001

Oct 29, 1991

AB USL PHARMA250MGN70631 001

Jun 11, 1987

PRESCRIPTION DRUG PRODUCT LIST

3 - 362 (of 370)

VALPROIC ACID

SOLUTION; ORAL					
<u>VALPROIC ACID</u>					
<u>AA</u>	VINTAGE	<u>250MG/5ML</u>	<u>N77960</u>	<u>001</u>	Oct 13, 2006
SYRUP; ORAL					
<u>DEPAKENE</u>					
<u>AA</u>	+ ABBOTT	<u>250MG/5ML</u>	<u>N18082</u>	<u>001</u>	
<u>MYPROIC ACID</u>					
<u>AA</u>	MORTON GROVE	<u>250MG/5ML</u>	<u>N70868</u>	<u>001</u>	Jul 01, 1986
<u>VALPROIC ACID</u>					
<u>AA</u>	APOTEX INC	<u>250MG/5ML</u>	<u>N77105</u>	<u>001</u>	Jul 29, 2005
<u>AA</u>	HIGH TECH PHARMA	<u>250MG/5ML</u>	<u>N74060</u>	<u>001</u>	Jan 13, 1995
<u>AA</u>	PHARM ASSOC	<u>250MG/5ML</u>	<u>N75379</u>	<u>001</u>	Dec 15, 2000
<u>AA</u>	TEVA PHARMS	<u>250MG/5ML</u>	<u>N73178</u>	<u>001</u>	Aug 25, 1992
<u>AA</u>	UDL	<u>250MG/5ML</u>	<u>N75782</u>	<u>001</u>	Dec 22, 2000

VALRUBICIN

SOLUTION; INTRAVESICAL					
VALSTAR PRESERVATIVE FREE					
	+ VALERA	40MG/ML	N20892	001	Sep 25, 1998

VALSARTAN

TABLET; ORAL					
DIOVAN					
	NOVARTIS	40MG	N21283	004	Aug 14, 2002
		80MG	N21283	001	Jul 18, 2001
		160MG	N21283	002	Jul 18, 2001
	+	320MG	N21283	003	Jul 18, 2001

VANCOMYCIN HYDROCHLORIDE

CAPSULE; ORAL					
VANCOCIN HYDROCHLORIDE					
	VIROPHARMA	EQ 125MG BASE	N50606	001	Apr 15, 1986
	+	EQ 250MG BASE	N50606	002	Apr 15, 1986
INJECTABLE; INJECTION					
VANCOCIN HYDROCHLORIDE IN PLASTIC CONTAINER					
	+ BAXTER HLTHCARE	EQ 500MG BASE/100ML	N50671	001	Apr 29, 1993
<u>VANCOMYCIN HYDROCHLORIDE</u>					
<u>AP</u>	+ ABRAXIS PHARM	<u>EQ 500MG BASE/VIAL</u>	<u>N62663</u>	<u>001</u>	Mar 17, 1987
<u>AP</u>	+	<u>EQ 1GM BASE/VIAL</u>	<u>N62663</u>	<u>002</u>	Jul 31, 1987
<u>AP</u>	+	<u>EQ 5GM BASE/VIAL</u>	<u>N62663</u>	<u>003</u>	Jun 03, 1988
	+	EQ 10GM BASE/VIAL	N62663	004	Nov 28, 1997
<u>AP</u>	+ HOSPIRA	<u>EQ 500MG BASE/VIAL</u>	<u>N62911</u>	<u>001</u>	Aug 04, 1988
<u>AP</u>	+	<u>EQ 500MG BASE/VIAL</u>	<u>N62931</u>	<u>001</u>	Oct 29, 1992
<u>AP</u>	+	<u>EQ 1GM BASE/VIAL</u>	<u>N62912</u>	<u>001</u>	Aug 04, 1988
<u>AP</u>	+	<u>EQ 1GM BASE/VIAL</u>	<u>N62933</u>	<u>001</u>	Oct 29, 1992
<u>AP</u>	+	<u>EQ 5GM BASE/VIAL</u>	<u>N63076</u>	<u>001</u>	Dec 21, 1990

VARDENAFIL HYDROCHLORIDE

TABLET; ORAL					
LEVITRA					
	BAYER PHARMS	2.5MG	N21400	003	Aug 19, 2003
		5MG	N21400	001	Aug 19, 2003
		10MG	N21400	002	Aug 19, 2003
	+	20MG	N21400	004	Aug 19, 2003

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 363 (of 370)

VARENICLINE TARTRATE

TABLET; ORAL

CHANTIX

PFIZER INC

EQ 0.5MG BASE

N21928 001

May 10, 2006

+

EQ 1MG BASE

N21928 002

May 10, 2006

VECURONIUM BROMIDE

INJECTABLE; INJECTION

VECURONIUM BROMIDE

<u>AP</u>	BAXTER HLTHCARE	<u>10MG/VIAL</u>	<u>N75218</u>	<u>001</u>	Aug 23, 1999
<u>AP</u>		<u>20MG/VIAL</u>	<u>N75218</u>	<u>002</u>	Aug 23, 1999
<u>AP</u>	+ BEDFORD	<u>10MG/VIAL</u>	<u>N75549</u>	<u>001</u>	Jun 13, 2000
<u>AP</u>	+	<u>20MG/VIAL</u>	<u>N75549</u>	<u>002</u>	Jun 13, 2000
<u>AP</u>	HOSPIRA	<u>10MG/VIAL</u>	<u>N75164</u>	<u>001</u>	Oct 21, 1999
<u>AP</u>		<u>20MG/VIAL</u>	<u>N75164</u>	<u>002</u>	Oct 21, 1999
	+	4MG/VIAL	N75558	001	Sep 11, 2001
<u>AP</u>	SICOR PHARMS	<u>10MG/VIAL</u>	<u>N74688</u>	<u>001</u>	Aug 25, 1999
<u>AP</u>		<u>20MG/VIAL</u>	<u>N74688</u>	<u>002</u>	Aug 25, 1999
<u>AP</u>	WATSON LABS	<u>10MG/VIAL</u>	<u>N74334</u>	<u>001</u>	Aug 31, 1995
<u>AP</u>		<u>20MG/VIAL</u>	<u>N74334</u>	<u>002</u>	Aug 31, 1995

VENLAFAXINE HYDROCHLORIDE

CAPSULE, EXTENDED RELEASE; ORAL

EFFEXOR XR

WYETH PHARMS INC

EQ 37.5MG BASE

N20699 001

Oct 20, 1997

EQ 75MG BASE

N20699 002

Oct 20, 1997

+

EQ 150MG BASE

N20699 004

Oct 20, 1997

TABLET; ORAL

EFFEXOR

<u>AB</u>	WYETH PHARMS INC	<u>EQ 25MG BASE</u>	<u>N20151</u>	<u>002</u>	Dec 28, 1993
<u>AB</u>		<u>EQ 37.5MG BASE</u>	<u>N20151</u>	<u>006</u>	Dec 28, 1993
<u>AB</u>	+	<u>EQ 50MG BASE</u>	<u>N20151</u>	<u>003</u>	Dec 28, 1993
<u>AB</u>		<u>EQ 75MG BASE</u>	<u>N20151</u>	<u>004</u>	Dec 28, 1993
<u>AB</u>		<u>EQ 100MG BASE</u>	<u>N20151</u>	<u>005</u>	Dec 28, 1993
	<u>VENLAFAXINE HYDROCHLORIDE</u>				
<u>AB</u>	TEVA	<u>EQ 25MG BASE</u>	<u>N76690</u>	<u>001</u>	Aug 03, 2006
<u>AB</u>		<u>EQ 37.5MG BASE</u>	<u>N76690</u>	<u>002</u>	Aug 03, 2006
<u>AB</u>		<u>EQ 50MG BASE</u>	<u>N76690</u>	<u>003</u>	Aug 03, 2006
<u>AB</u>		<u>EQ 75MG BASE</u>	<u>N76690</u>	<u>004</u>	Aug 03, 2006
<u>AB</u>		<u>EQ 100MG BASE</u>	<u>N76690</u>	<u>005</u>	Aug 03, 2006

VERAPAMIL HYDROCHLORIDE

CAPSULE, EXTENDED RELEASE; ORAL

VERAPAMIL HYDROCHLORIDE

<u>AB</u>	MYLAN	<u>120MG</u>	<u>N75138</u>	<u>001</u>	Apr 20, 1999
<u>AB</u>		<u>180MG</u>	<u>N75138</u>	<u>002</u>	Apr 20, 1999
<u>AB</u>		<u>240MG</u>	<u>N75138</u>	<u>003</u>	Apr 20, 1999
	<u>VERELAN</u>				
<u>AB</u>	+ ELAN DRUG	<u>120MG</u>	<u>N19614</u>	<u>001</u>	May 29, 1990
<u>AB</u>	+	<u>180MG</u>	<u>N19614</u>	<u>003</u>	Jan 09, 1992
<u>AB</u>	+	<u>240MG</u>	<u>N19614</u>	<u>002</u>	May 29, 1990
	+	360MG	N19614	004	May 10, 1996
	VERELAN PM				
	ELAN DRUG	100MG	N20943	001	Nov 25, 1998
		200MG	N20943	002	Nov 25, 1998
	+	300MG	N20943	003	Nov 25, 1998

INJECTABLE; INJECTION

ISOPTIN

<u>AP</u>	+ FSC	<u>2.5MG/ML</u>	<u>N18485</u>	<u>001</u>	
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PRESCRIPTION DRUG PRODUCT LIST

3 - 364 (of 370)

VERAPAMIL HYDROCHLORIDE

INJECTABLE; INJECTION

VERAPAMIL HYDROCHLORIDE

<u>AP</u>	BEDFORD	<u>2.5MG/ML</u>	<u>N72888</u>	<u>001</u>	Jul 28, 1995
<u>AP</u>	HOSPIRA	<u>2.5MG/ML</u>	<u>N70577</u>	<u>001</u>	Feb 02, 1987
<u>AP</u>		<u>2.5MG/ML</u>	<u>N70737</u>	<u>001</u>	May 06, 1987
<u>AP</u>		<u>2.5MG/ML</u>	<u>N70738</u>	<u>001</u>	May 06, 1987
<u>AP</u>		<u>2.5MG/ML</u>	<u>N75136</u>	<u>001</u>	Oct 20, 1998
<u>AP</u>	INTL MEDICATION	<u>2.5MG/ML</u>	<u>N70451</u>	<u>001</u>	Dec 16, 1985
<u>AP</u>	LUITPOLD	<u>2.5MG/ML</u>	<u>N70225</u>	<u>001</u>	Nov 12, 1985
<u>AP</u>		<u>2.5MG/ML</u>	<u>N70617</u>	<u>001</u>	Nov 12, 1985

TABLET; ORAL

CALAN

<u>AB</u>	GD SEARLE LLC	<u>40MG</u>	<u>N18817</u>	<u>003</u>	Feb 23, 1988
<u>AB</u>		<u>80MG</u>	<u>N18817</u>	<u>001</u>	Sep 10, 1984
<u>AB</u>		<u>120MG</u>	<u>N18817</u>	<u>002</u>	Sep 10, 1984

ISOPTIN

<u>AB</u>	FSC	<u>40MG</u>	<u>N18593</u>	<u>003</u>	Nov 23, 1987
<u>AB</u>		<u>80MG</u>	<u>N18593</u>	<u>001</u>	Mar 08, 1982
<u>AB</u>	+	<u>120MG</u>	<u>N18593</u>	<u>002</u>	Mar 08, 1982

VERAPAMIL HYDROCHLORIDE

<u>AB</u>	ACTAVIS ELIZABETH	<u>80MG</u>	<u>N71019</u>	<u>001</u>	Sep 24, 1986
<u>AB</u>		<u>120MG</u>	<u>N70468</u>	<u>001</u>	Sep 24, 1986
<u>AB</u>	MUTUAL PHARM	<u>80MG</u>	<u>N71488</u>	<u>001</u>	Jan 13, 1988
<u>AB</u>		<u>120MG</u>	<u>N71489</u>	<u>001</u>	Jan 13, 1988
<u>AB</u>	MYLAN	<u>80MG</u>	<u>N71482</u>	<u>001</u>	Feb 15, 1989
<u>AB</u>		<u>120MG</u>	<u>N71483</u>	<u>001</u>	Feb 15, 1989
<u>AB</u>	PLIVA	<u>80MG</u>	<u>N72124</u>	<u>001</u>	Jan 26, 1989
<u>AB</u>		<u>120MG</u>	<u>N72125</u>	<u>001</u>	Jan 26, 1989
<u>AB</u>	SANDOZ	<u>40MG</u>	<u>N73168</u>	<u>001</u>	Jul 31, 1992
<u>AB</u>		<u>80MG</u>	<u>N71423</u>	<u>001</u>	May 24, 1988
<u>AB</u>		<u>120MG</u>	<u>N71424</u>	<u>001</u>	May 25, 1988
<u>AB</u>	WATSON LABS	<u>40MG</u>	<u>N72923</u>	<u>001</u>	Jun 29, 1993
<u>AB</u>		<u>40MG</u>	<u>N72924</u>	<u>001</u>	Jun 29, 1993
<u>AB</u>		<u>80MG</u>	<u>N70855</u>	<u>001</u>	Sep 24, 1986
<u>AB</u>		<u>80MG</u>	<u>N70995</u>	<u>001</u>	Oct 01, 1986
<u>AB</u>		<u>80MG</u>	<u>N71366</u>	<u>001</u>	Oct 01, 1986
<u>AB</u>		<u>120MG</u>	<u>N70856</u>	<u>001</u>	Sep 24, 1986
<u>AB</u>		<u>120MG</u>	<u>N70994</u>	<u>001</u>	Oct 01, 1986
<u>AB</u>		<u>120MG</u>	<u>N71367</u>	<u>001</u>	Oct 01, 1986

TABLET, EXTENDED RELEASE; ORAL

COVERA-HS

BC	+	GD SEARLE LLC	180MG	N20552	001	Feb 26, 1996
BC	+		240MG	N20552	002	Feb 26, 1996

ISOPTIN SR

<u>AB</u>	+	FSC	<u>120MG</u>	<u>N19152</u>	<u>003</u>	Mar 06, 1991
<u>AB</u>	+		<u>180MG</u>	<u>N19152</u>	<u>002</u>	Dec 15, 1989
<u>AB</u>	+		<u>240MG</u>	<u>N19152</u>	<u>001</u>	Dec 16, 1986

VERAPAMIL HYDROCHLORIDE

<u>AB</u>	IVAX PHARMS	<u>120MG</u>	<u>N73568</u>	<u>002</u>	Oct 10, 1997
<u>AB</u>		<u>180MG</u>	<u>N74330</u>	<u>001</u>	Jan 31, 1994
<u>AB</u>		<u>240MG</u>	<u>N73568</u>	<u>001</u>	Jul 31, 1992
<u>AB</u>	KALI LABS	<u>120MG</u>	<u>N75072</u>	<u>001</u>	May 25, 1999
<u>AB</u>		<u>240MG</u>	<u>N75072</u>	<u>003</u>	May 25, 1999
<u>AB</u>	MYLAN	<u>120MG</u>	<u>N74587</u>	<u>002</u>	Feb 21, 1997
<u>AB</u>		<u>180MG</u>	<u>N74587</u>	<u>003</u>	Sep 09, 1997
<u>AB</u>		<u>240MG</u>	<u>N74587</u>	<u>001</u>	Mar 23, 1996
<u>AB</u>	PLIVA	<u>240MG</u>	<u>N72922</u>	<u>001</u>	Mar 01, 1996

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 365 (of 370)

VERTEPORFININJECTABLE; INJECTION
VISUDYNE

+ QLT 15MG/VIAL N21119 001 Apr 12, 2000

VINBLASTINE SULFATEINJECTABLE; INJECTION
VINBLASTINE SULFATE

+ ABRAXIS PHARM 1MG/ML N89515 001 Apr 29, 1987

+ BEDFORD 10MG/VIAL N89395 001 Apr 09, 1987

VINCRISTINE SULFATE

INJECTABLE; INJECTION

VINCRISTINE SULFATEAP + ABRAXIS PHARM 1MG/ML N76296 001 Dec 20, 2002AP 1MG/ML N76401 001 Oct 28, 2003VINCRISTINE SULFATE PFSAP MAYNE PHARMA USA 1MG/ML N71484 001 Apr 19, 1988AP SICOR PHARMS 1MG/ML N75493 001 Sep 01, 1999VINORELBINE TARTRATE

INJECTABLE; INJECTION

NAVELBINEAP + PIERRE FABRE EQ 10MG BASE/ML N20388 001 Dec 23, 1994VINORELBINE TARTRATEAP ABRAXIS PHARM EQ 10MG BASE/ML N76849 001 Apr 18, 2005AP BAXTER HLTHCARE EQ 10MG BASE/ML N75992 001 Jun 10, 2003AP BEDFORD EQ 10MG BASE/ML N76461 001 Dec 11, 2003AP MAYNE PHARMA USA EQ 10MG BASE/ML N76827 001 Jun 02, 2005AP SICOR PHARMS EQ 10MG BASE/ML N76028 001 Feb 03, 2003VITAMIN A PALMITATE

CAPSULE; ORAL

VITAMIN A PALMITATEAA ARCUM EQ 50,000 UNITS BASE N83311 001AA EQ 50,000 UNITS BASE N83321 001

INJECTABLE; INJECTION

AQUASOL A

+ MAYNE PHARMA USA EQ 50,000 UNITS BASE/ML N06823 001

VORICONAZOLE

FOR SUSPENSION; ORAL

VFEND

+ PFIZER 200MG/5ML N21630 001 Dec 19, 2003

INJECTABLE; IV (INFUSION)

+ PFIZER 200MG/VIAL N21267 001 May 24, 2002

TABLET; ORAL

VFEND

PFIZER 50MG N21266 001 May 24, 2002

+ 200MG N21266 002 May 24, 2002

VORINOSTAT

CAPSULE; ORAL

ZOLINZA

+ MERCK 100MG N21991 001 Oct 06, 2006

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 366 (of 370)

WARFARIN SODIUM

INJECTABLE; INJECTION

COUMADIN

+ BRISTOL MYERS SQUIBB 5MG/VIAL

N09218 024 Feb 07, 1995

TABLET; ORAL

COUMADIN

<u>AB</u>	BRISTOL MYERS SQUIBB	<u>1MG</u>	<u>N09218</u>	<u>022</u>	Mar 01, 1990
<u>AB</u>		<u>2MG</u>	<u>N09218</u>	<u>013</u>	
<u>AB</u>		<u>2.5MG</u>	<u>N09218</u>	<u>018</u>	
<u>AB</u>		<u>3MG</u>	<u>N09218</u>	<u>025</u>	Nov 18, 1996
<u>AB</u>		<u>4MG</u>	<u>N09218</u>	<u>023</u>	Aug 24, 1993
<u>AB</u>		<u>5MG</u>	<u>N09218</u>	<u>007</u>	
<u>AB</u>		<u>6MG</u>	<u>N09218</u>	<u>026</u>	Nov 18, 1996
<u>AB</u>		<u>7.5MG</u>	<u>N09218</u>	<u>016</u>	
<u>AB</u>	+	<u>10MG</u>	<u>N09218</u>	<u>005</u>	

JANTOVEN

<u>AB</u>	USL PHARMA	<u>1MG</u>	<u>N40416</u>	<u>001</u>	Oct 02, 2003
<u>AB</u>		<u>2MG</u>	<u>N40416</u>	<u>002</u>	Oct 02, 2003
<u>AB</u>		<u>2.5MG</u>	<u>N40416</u>	<u>003</u>	Oct 02, 2003
<u>AB</u>		<u>3MG</u>	<u>N40416</u>	<u>004</u>	Oct 02, 2003
<u>AB</u>		<u>4MG</u>	<u>N40416</u>	<u>005</u>	Oct 02, 2003
<u>AB</u>		<u>5MG</u>	<u>N40416</u>	<u>006</u>	Oct 02, 2003
<u>AB</u>		<u>6MG</u>	<u>N40416</u>	<u>007</u>	Oct 02, 2003
<u>AB</u>		<u>7.5MG</u>	<u>N40416</u>	<u>008</u>	Oct 02, 2003
<u>AB</u>		<u>10MG</u>	<u>N40416</u>	<u>009</u>	Oct 02, 2003

WARFARIN SODIUM

<u>AB</u>	BARR	<u>1MG</u>	<u>N40145</u>	<u>001</u>	Mar 26, 1997
<u>AB</u>		<u>2MG</u>	<u>N40145</u>	<u>002</u>	Mar 26, 1997
<u>AB</u>		<u>2.5MG</u>	<u>N40145</u>	<u>003</u>	Mar 26, 1997
<u>AB</u>		<u>3MG</u>	<u>N40145</u>	<u>008</u>	Nov 05, 1998
<u>AB</u>		<u>4MG</u>	<u>N40145</u>	<u>004</u>	Mar 26, 1997
<u>AB</u>		<u>5MG</u>	<u>N40145</u>	<u>005</u>	Mar 26, 1997
<u>AB</u>		<u>6MG</u>	<u>N40145</u>	<u>009</u>	Nov 05, 1998
<u>AB</u>		<u>7.5MG</u>	<u>N40145</u>	<u>006</u>	Mar 26, 1997
<u>AB</u>		<u>10MG</u>	<u>N40145</u>	<u>007</u>	Mar 26, 1997
<u>AB</u>	GENPHARM	<u>1MG</u>	<u>N40415</u>	<u>001</u>	Sep 27, 2004
<u>AB</u>		<u>2MG</u>	<u>N40415</u>	<u>002</u>	Sep 27, 2004
<u>AB</u>		<u>2.5MG</u>	<u>N40415</u>	<u>003</u>	Sep 29, 2004
<u>AB</u>		<u>3MG</u>	<u>N40415</u>	<u>004</u>	Sep 27, 2004
<u>AB</u>		<u>4MG</u>	<u>N40415</u>	<u>005</u>	Sep 27, 2004
<u>AB</u>		<u>5MG</u>	<u>N40415</u>	<u>006</u>	Sep 27, 2004
<u>AB</u>		<u>6MG</u>	<u>N40415</u>	<u>007</u>	Sep 27, 2004
<u>AB</u>		<u>7.5MG</u>	<u>N40415</u>	<u>008</u>	Sep 27, 2004
<u>AB</u>		<u>10MG</u>	<u>N40415</u>	<u>009</u>	Sep 27, 2004
<u>AB</u>	PLIVA	<u>1MG</u>	<u>N40616</u>	<u>009</u>	Jul 05, 2006
<u>AB</u>		<u>2MG</u>	<u>N40616</u>	<u>001</u>	Jul 05, 2006
<u>AB</u>		<u>2.5MG</u>	<u>N40616</u>	<u>002</u>	Jul 05, 2006
<u>AB</u>		<u>3MG</u>	<u>N40616</u>	<u>003</u>	Jul 05, 2006
<u>AB</u>		<u>4MG</u>	<u>N40616</u>	<u>004</u>	Jul 05, 2006
<u>AB</u>		<u>5MG</u>	<u>N40616</u>	<u>005</u>	Jul 05, 2006
<u>AB</u>		<u>6MG</u>	<u>N40616</u>	<u>006</u>	Jul 05, 2006
<u>AB</u>		<u>7.5MG</u>	<u>N40616</u>	<u>007</u>	Jul 05, 2006
<u>AB</u>		<u>10MG</u>	<u>N40616</u>	<u>008</u>	Jul 05, 2006
<u>AB</u>	SANDOZ	<u>1MG</u>	<u>N40196</u>	<u>001</u>	Sep 30, 1997
<u>AB</u>		<u>2MG</u>	<u>N40196</u>	<u>002</u>	Sep 30, 1997
<u>AB</u>		<u>2.5MG</u>	<u>N40196</u>	<u>003</u>	Sep 30, 1997
<u>AB</u>		<u>3MG</u>	<u>N40196</u>	<u>008</u>	Jul 26, 2000
<u>AB</u>		<u>4MG</u>	<u>N40196</u>	<u>004</u>	Sep 30, 1997
<u>AB</u>		<u>5MG</u>	<u>N40196</u>	<u>005</u>	Sep 30, 1997
<u>AB</u>		<u>6MG</u>	<u>N40196</u>	<u>009</u>	Jul 26, 2000

PRESCRIPTION DRUG PRODUCT LIST

3 - 367 (of 370)

WARFARIN SODIUM

TABLET; ORAL

WARFARIN SODIUM

<u>AB</u>	SANDOZ	<u>7.5MG</u>	<u>N40196</u>	<u>006</u>	Sep 30, 1997
<u>AB</u>		<u>10MG</u>	<u>N40196</u>	<u>007</u>	Sep 30, 1997
<u>AB</u>	TARO	<u>1MG</u>	<u>N40301</u>	<u>002</u>	Jul 15, 1999
<u>AB</u>		<u>2MG</u>	<u>N40301</u>	<u>003</u>	Jul 15, 1999
<u>AB</u>		<u>2.5MG</u>	<u>N40301</u>	<u>004</u>	Jul 15, 1999
<u>AB</u>		<u>3MG</u>	<u>N40301</u>	<u>005</u>	Jul 15, 1999
<u>AB</u>		<u>4MG</u>	<u>N40301</u>	<u>006</u>	Jul 15, 1999
<u>AB</u>		<u>5MG</u>	<u>N40301</u>	<u>007</u>	Jul 15, 1999
<u>AB</u>		<u>6MG</u>	<u>N40301</u>	<u>008</u>	Jul 15, 1999
<u>AB</u>		<u>7.5MG</u>	<u>N40301</u>	<u>009</u>	Jul 15, 1999
<u>AB</u>		<u>10MG</u>	<u>N40301</u>	<u>001</u>	Jul 15, 1999
<u>AB</u>	ZYDUS PHARMS USA	<u>1MG</u>	<u>N40663</u>	<u>001</u>	May 30, 2006
<u>AB</u>		<u>2MG</u>	<u>N40663</u>	<u>002</u>	May 30, 2006
<u>AB</u>		<u>2.5MG</u>	<u>N40663</u>	<u>003</u>	May 30, 2006
<u>AB</u>		<u>3MG</u>	<u>N40663</u>	<u>004</u>	May 30, 2006
<u>AB</u>		<u>4MG</u>	<u>N40663</u>	<u>005</u>	May 30, 2006
<u>AB</u>		<u>5MG</u>	<u>N40663</u>	<u>006</u>	May 30, 2006
<u>AB</u>		<u>6MG</u>	<u>N40663</u>	<u>007</u>	May 30, 2006
<u>AB</u>		<u>7.5MG</u>	<u>N40663</u>	<u>008</u>	May 30, 2006
<u>AB</u>		<u>10MG</u>	<u>N40663</u>	<u>009</u>	May 30, 2006

WATER FOR INJECTION, STERILE

LIQUID; N/A

BACTERIOSTATIC WATER FOR INJECTION IN PLASTIC CONTAINER

<u>AP</u>	+ HOSPIRA	<u>100%</u>	<u>N18802</u>	<u>001</u>	Oct 27, 1982
<u>STERILE WATER FOR INJECTION IN PLASTIC CONTAINER</u>					
<u>AP</u>	ABRAXIS PHARM	<u>100%</u>	<u>N88400</u>	<u>001</u>	Jan 16, 1984
<u>AP</u>	+ B BRAUN	<u>100%</u>	<u>N19633</u>	<u>001</u>	Feb 29, 1988
<u>AP</u>	+ BAXTER HLTHCARE	<u>100%</u>	<u>N18632</u>	<u>002</u>	Apr 19, 1988
<u>AP</u>	+	<u>100%</u>	<u>N18632</u>	<u>001</u>	Jun 30, 1982
<u>AP</u>	+ HOSPIRA	<u>100%</u>	<u>N18233</u>	<u>001</u>	
<u>AP</u>	+	<u>100%</u>	<u>N18801</u>	<u>001</u>	Oct 27, 1982
<u>AP</u>	+	<u>100%</u>	<u>N19869</u>	<u>001</u>	Dec 26, 1989
<u>AP</u>	TARO PHARMS IRELAND	<u>100%</u>	<u>N77393</u>	<u>001</u>	Aug 11, 2006

WATER FOR IRRIGATION, STERILE

LIQUID; IRRIGATION

STERILE WATER

<u>AT</u>	BAXTER HLTHCARE	<u>100%</u>	<u>N17428</u>	<u>001</u>	
<u>STERILE WATER IN PLASTIC CONTAINER</u>					
<u>AT</u>	B BRAUN	<u>100%</u>	<u>N16734</u>	<u>001</u>	
<u>AT</u>	BAXTER HLTHCARE	<u>100%</u>	<u>N17866</u>	<u>001</u>	
<u>AT</u>	HOSPIRA	<u>100%</u>	<u>N17513</u>	<u>001</u>	
<u>AT</u>		<u>100%</u>	<u>N18313</u>	<u>001</u>	

XENON, XE-133

GAS; INHALATION

XENON XE 133

<u>AA</u>	BRISTOL MYERS SQUIBB	<u>10mCi/VIAL</u>	<u>N17284</u>	<u>001</u>	
<u>AA</u>		<u>20mCi/VIAL</u>	<u>N17284</u>	<u>002</u>	
<u>AA</u>	MALLINCKRODT	<u>10mCi/VIAL</u>	<u>N18327</u>	<u>001</u>	Mar 09, 1982
<u>AA</u>		<u>20mCi/VIAL</u>	<u>N18327</u>	<u>002</u>	Mar 09, 1982

ZAFIRLUKAST

TABLET; ORAL

ACCOLATE

	ASTRAZENECA	10MG	N20547	003	Sep 17, 1999
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27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 368 (of 370)

ZAFIRLUKASTTABLET; ORAL
ACCOLATE

+ ASTRAZENECA 20MG N20547 001 Sep 26, 1996

ZALCITABINETABLET; ORAL
HIVID

ROCHE 0.375MG N20199 001 Jun 19, 1992

+ 0.75MG N20199 002 Jun 19, 1992

ZALEPLONCAPSULE; ORAL
SONATA

KING PHARMS 5MG N20859 001 Aug 13, 1999

+ 10MG N20859 002 Aug 13, 1999

ZANAMIVIRPOWDER; INHALATION
RELENZA

+ GLAXOSMITHKLINE 5MG N21036 001 Jul 26, 1999

ZICONOTIDEINJECTABLE; INTRATHECAL
PRIALT

+ ELAN PHARMS 100UGM/1ML (100MG/ML) N21060 002 Dec 28, 2004

+ 500UGM/20ML (25MG/ML) N21060 001 Dec 28, 2004

+ 500UGM/5ML (100MG/ML) N21060 004 Dec 28, 2004

ZIDOVUDINE

CAPSULE; ORAL

RETROVIRAB + GLAXOSMITHKLINE 100MG N19655 001 Mar 19, 1987ZIDOVUDINEAB AUROBINDO PHARMA LTD 100MG N78128 001 Mar 27, 2006INJECTABLE; INJECTION
RETROVIR

+ GLAXOSMITHKLINE 10MG/ML N19951 001 Feb 02, 1990

SYRUP; ORAL

RETROVIRAA + GLAXOSMITHKLINE 50MG/5ML N19910 001 Sep 28, 1989ZIDOVUDINEAA AUROBINDO 50MG/5ML N77268 001 Sep 19, 2005

TABLET; ORAL

RETROVIRAB + GLAXOSMITHKLINE 300MG N20518 002 Oct 04, 1996ZIDOVUDINEAB AUROBINDO 300MG N77267 001 Sep 19, 2005AB RANBAXY 300MG N77327 001 Sep 19, 2005AB ROXANE 300MG N76844 001 Sep 19, 2005ZILEUTONTABLET; ORAL
ZYFLO

+ CRITICAL 600MG N20471 003 Dec 09, 1996

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 369 (of 370)

ZINC ACETATE

CAPSULE; ORAL

GALZIN

TEVA

EQ 25MG ZINC

N20458 001

Jan 28, 1997

+

EQ 50MG ZINC

N20458 002

Jan 28, 1997

ZINC CHLORIDE

INJECTABLE; INJECTION

ZINC CHLORIDE IN PLASTIC CONTAINER

+ HOSPIRA

EQ 1MG ZINC/ML

N18959 001

Jun 26, 1986

ZIPRASIDONE HYDROCHLORIDE

CAPSULE; ORAL

GEODON

+ PFIZER

EQ 20MG BASE

N20825 001

Feb 05, 2001

EQ 40MG BASE

N20825 002

Feb 05, 2001

EQ 60MG BASE

N20825 003

Feb 05, 2001

EQ 80MG BASE

N20825 004

Feb 05, 2001

SUSPENSION; ORAL

GEODON

+ PFIZER INC

EQ 10MG BASE/ML

N21483 001

Mar 29, 2006

ZIPRASIDONE MESYLATE

INJECTABLE; INTRAMUSCULAR

GEODON

+ PFIZER

EQ 20MG BASE/ML

N20919 001

Jun 21, 2002

ZOLEDRONIC ACID

INJECTABLE; IV (INFUSION)

+ NOVARTIS

EQ 4MG BASE/5ML

N21223 002

Mar 07, 2003

+

EQ 4MG BASE/VIAL

N21223 001

Aug 20, 2001

ZOLMITRIPTAN

SPRAY; NASAL

ZOMIG

+ ASTRAZENECA

5MG/SPRAY

N21450 004

Sep 30, 2003

TABLET; ORAL

ZOMIG

IPR

2.5MG

N20768 001

Nov 25, 1997

+

5MG

N20768 002

Nov 25, 1997

TABLET, ORALLY DISINTEGRATING; ORAL

ZOMIG-ZMT

ASTRAZENECA

2.5MG

N21231 001

Feb 13, 2001

+

5MG

N21231 002

Sep 17, 2001

ZOLPIDEM TARTRATE

TABLET; ORAL

AMBIEN

SANOFI AVENTIS US

5MG

N19908 001

Dec 16, 1992

+

10MG

N19908 002

Dec 16, 1992

TABLET, EXTENDED RELEASE; ORAL

AMBIEN CR

SANOFI AVENTIS US

6.25MG

N21774 002

Sep 02, 2005

+

12.5MG

N21774 001

Sep 02, 2005

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

PRESCRIPTION DRUG PRODUCT LIST

3 - 370 (of 370)

ZONISAMIDE

CAPSULE; ORAL

ZONEGRAN

<u>AB</u>	DAINIPPON	<u>25MG</u>	<u>N20789</u>	<u>003</u>	Aug 22, 2003
<u>AB</u>		<u>50MG</u>	<u>N20789</u>	<u>002</u>	Aug 22, 2003
<u>AB</u>	+	<u>100MG</u>	<u>N20789</u>	<u>001</u>	Mar 27, 2000
	<u>ZONISAMIDE</u>				
<u>AB</u>	ALPHAPHARM	<u>25MG</u>	<u>N77647</u>	<u>001</u>	Dec 22, 2005
<u>AB</u>		<u>50MG</u>	<u>N77647</u>	<u>002</u>	Dec 22, 2005
<u>AB</u>		<u>100MG</u>	<u>N77647</u>	<u>003</u>	Dec 22, 2005
<u>AB</u>	APOTEX INC	<u>25MG</u>	<u>N77642</u>	<u>001</u>	Dec 22, 2005
<u>AB</u>		<u>50MG</u>	<u>N77642</u>	<u>002</u>	Dec 22, 2005
<u>AB</u>		<u>100MG</u>	<u>N77642</u>	<u>003</u>	Dec 22, 2005
<u>AB</u>	BANNER PHARMACAPS	<u>25MG</u>	<u>N77813</u>	<u>001</u>	Aug 16, 2006
<u>AB</u>		<u>50MG</u>	<u>N77813</u>	<u>002</u>	Aug 16, 2006
<u>AB</u>		<u>100MG</u>	<u>N77813</u>	<u>003</u>	Aug 16, 2006
<u>AB</u>	BARR	<u>25MG</u>	<u>N77639</u>	<u>001</u>	Dec 22, 2005
<u>AB</u>		<u>50MG</u>	<u>N77639</u>	<u>002</u>	Dec 22, 2005
<u>AB</u>		<u>100MG</u>	<u>N77639</u>	<u>003</u>	Dec 22, 2005
<u>AB</u>	DR REDDYS LABS LTD	<u>25MG</u>	<u>N77891</u>	<u>001</u>	Sep 29, 2006
<u>AB</u>		<u>50MG</u>	<u>N77891</u>	<u>002</u>	Sep 29, 2006
<u>AB</u>		<u>100MG</u>	<u>N77645</u>	<u>001</u>	Dec 22, 2005
<u>AB</u>	GLENMARK PHARMS	<u>25MG</u>	<u>N77651</u>	<u>001</u>	Jan 30, 2006
<u>AB</u>		<u>50MG</u>	<u>N77651</u>	<u>002</u>	Jan 30, 2006
<u>AB</u>		<u>100MG</u>	<u>N77651</u>	<u>003</u>	Jan 30, 2006
<u>AB</u>	INVAGEN PHARMS	<u>25MG</u>	<u>N77869</u>	<u>001</u>	May 31, 2006
<u>AB</u>		<u>50MG</u>	<u>N77869</u>	<u>002</u>	May 31, 2006
<u>AB</u>		<u>100MG</u>	<u>N77869</u>	<u>003</u>	May 31, 2006
<u>AB</u>	MUTUAL PHARM	<u>25MG</u>	<u>N77635</u>	<u>001</u>	Dec 22, 2005
<u>AB</u>		<u>50MG</u>	<u>N77635</u>	<u>002</u>	Dec 22, 2005
<u>AB</u>		<u>100MG</u>	<u>N77635</u>	<u>003</u>	Dec 22, 2005
<u>AB</u>	MYLAN	<u>25MG</u>	<u>N77637</u>	<u>001</u>	Dec 22, 2005
<u>AB</u>		<u>50MG</u>	<u>N77637</u>	<u>002</u>	Dec 22, 2005
<u>AB</u>		<u>100MG</u>	<u>N77637</u>	<u>003</u>	Dec 22, 2005
<u>AB</u>	ROXANE	<u>25MG</u>	<u>N77648</u>	<u>001</u>	Dec 22, 2005
<u>AB</u>		<u>50MG</u>	<u>N77648</u>	<u>002</u>	Dec 22, 2005
<u>AB</u>		<u>100MG</u>	<u>N77648</u>	<u>003</u>	Dec 22, 2005
<u>AB</u>	SANDOZ	<u>25MG</u>	<u>N77644</u>	<u>001</u>	Dec 22, 2005
<u>AB</u>		<u>50MG</u>	<u>N77644</u>	<u>002</u>	Dec 22, 2005
<u>AB</u>		<u>100MG</u>	<u>N77644</u>	<u>003</u>	Dec 22, 2005
<u>AB</u>	SUN PHARM INDS (IN)	<u>25MG</u>	<u>N77634</u>	<u>001</u>	Mar 17, 2006
<u>AB</u>		<u>50MG</u>	<u>N77634</u>	<u>002</u>	Mar 17, 2006
<u>AB</u>		<u>100MG</u>	<u>N77634</u>	<u>003</u>	Mar 17, 2006
<u>AB</u>	TEVA PHARMS	<u>25MG</u>	<u>N77641</u>	<u>003</u>	Dec 22, 2005
<u>AB</u>		<u>50MG</u>	<u>N77641</u>	<u>002</u>	Dec 22, 2005
<u>AB</u>		<u>100MG</u>	<u>N77641</u>	<u>001</u>	Dec 22, 2005
<u>AB</u>	WATSON LABS	<u>25MG</u>	<u>N77650</u>	<u>001</u>	Apr 20, 2006
<u>AB</u>		<u>50MG</u>	<u>N77650</u>	<u>002</u>	Apr 20, 2006
<u>AB</u>		<u>100MG</u>	<u>N77650</u>	<u>003</u>	Apr 20, 2006
<u>AB</u>	WOCKHARDT	<u>25MG</u>	<u>N77636</u>	<u>003</u>	Jul 27, 2006
<u>AB</u>		<u>50MG</u>	<u>N77636</u>	<u>002</u>	Jul 27, 2006
<u>AB</u>		<u>100MG</u>	<u>N77636</u>	<u>001</u>	Dec 22, 2005
<u>AB</u>	ZYDUS PHARMS USA	<u>25MG</u>	<u>N77625</u>	<u>001</u>	Oct 16, 2006
<u>AB</u>		<u>50MG</u>	<u>N77625</u>	<u>002</u>	Oct 16, 2006
<u>AB</u>		<u>100MG</u>	<u>N77625</u>	<u>003</u>	Oct 16, 2006

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

OTC DRUG PRODUCT LIST

4 - 1 (of 15)

ACETAMINOPHEN

SUPPOSITORY; RECTAL

ACEPHEN

G AND W LABS

120MG

N18060 001

120MG

N72218 001

Mar 27, 1992

325MG

N18060 003

Dec 18, 1986

325MG

N72344 001

Mar 27, 1992

650MG

N18060 002

650MG

N72237 001

Mar 27, 1992

ACETAMINOPHEN

ACTAVIS MID ATLANTIC

120MG

N18337 003

Sep 12, 1983

325MG

N18337 002

+

650MG

N18337 001

SUPPOSITORIA

120MG

N70607 001

Apr 06, 1987

650MG

N70608 001

Dec 01, 1986

INFANTS' FEVERALL

ACTAVIS MID ATLANTIC

80MG

N18337 004

Aug 26, 1992

NEOPAP

POLYMEDICA

120MG

N16401 001

TABLET, EXTENDED RELEASE; ORAL

ACETAMINOPHEN

COREPHARMA

650MG

N76200 001

Mar 19, 2002

PERRIGO

650MG

N75077 001

Feb 25, 2000

TYLENOL (CAPLET)

+

MCNEIL CONS

650MG

N19872 001

Jun 08, 1994

TYLENOL (GELTAB)

+

MCNEIL CONS

650MG

N19872 002

Jan 11, 2001

ACETAMINOPHEN; ASPIRIN; CAFFEINE

TABLET; ORAL

ACETAMINOPHEN, ASPIRIN AND CAFFEINE

PERRIGO

250MG;250MG;65MG

N75794 001

Nov 26, 2001

EXCEDRIN (MIGRAINE)

+

NOVARTIS

250MG;250MG;65MG

N20802 001

Jan 14, 1998

ACETAMINOPHEN; CLEMASTINE FUMARATE; PSEUDOEPHEDRINE HYDROCHLORIDE

TABLET; ORAL

TAVIST ALLERGY/SINUS/HEADACHE

+

NOVARTIS

500MG;EQ 0.25MG BASE;30MG

N21082 001

Mar 01, 2001

ACETAMINOPHEN; DEXBROMPHENIRAMINE MALEATE; PSEUDOEPHEDRINE SULFATE

TABLET, EXTENDED RELEASE; ORAL

DRIXORAL PLUS

+

SCHERING PLOUGH

500MG;3MG;60MG

N19453 001

May 22, 1987

ALCOHOL; CHLORHEXIDINE GLUCONATE

SOLUTION; TOPICAL

AVAGARD

+

3M

61%;1%

N21074 001

Jun 07, 2001

ALUMINUM HYDROXIDE; MAGNESIUM TRISILICATE

TABLET, CHEWABLE; ORAL

FOAMCOAT

GUARDIAN DRUG

80MG;20MG

N71793 001

Sep 04, 1987

GAVISCON

SANOFI AVENTIS US

80MG;20MG

N18685 001

Dec 09, 1983

+

160MG;40MG

N18685 002

Dec 09, 1983

ANTAZOLINE PHOSPHATE; NAPHAZOLINE HYDROCHLORIDE

SOLUTION/DROPS; OPHTHALMIC

VASOCON-A

+

NOVARTIS

0.5%;0.05%

N18746 002

Jul 11, 1994

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

OTC DRUG PRODUCT LIST

4 - 2 (of 15)

ASPIRIN

TABLET; ORAL

BAYER EXTRA STRENGTH ASPIRIN FOR MIGRAINE PAIN

BAYER

500MG

N21317 001

Oct 18, 2001

AVOBENZONE; ECAMSULE; OCTOCRYLENE

CREAM; TOPICAL

ANTHELIOS SX

+ LOREAL USA

2%;2%;10%

N21502 001

Jul 21, 2006

CAPITAL SOLEIL 15

+ LOREAL USA

2%;3%;10%

N21501 001

Oct 02, 2006

AVOBENZONE; ECAMSULE; OCTOCRYLENE; TITANIUM DIOXIDE

CREAM; TOPICAL

ANTHELIOS 20

+ LOREAL USA

2%;2%;10%;2%

N21471 001

Oct 05, 2006

AVOBENZONE; OCTINOXATE; OXYBENZONE

LOTION; TOPICAL

SHADE UVAGUARD

+ SCHERING PLOUGH

3%;7.5%;3%

N20045 001

Dec 07, 1992

BENTOQUATAM

LOTION; TOPICAL

IVY BLOCK

+ STAND HOMEOPATH

5%

N20532 001

Aug 26, 1996

BROMPHENIRAMINE MALEATE; PSEUDOEPHEDRINE HYDROCHLORIDE

TABLET, EXTENDED RELEASE; ORAL

EFIDAC 24 PSEUDOEPHEDRINE HYDROCHLORIDE/BROMPHENIRAMINE MALEATE

+ ALZA

16MG;240MG

N19672 001

Mar 29, 1996

BUTENAFINE HYDROCHLORIDE

CREAM; TOPICAL

LOTRIMIN ULTRA

+ SCHERING PLOUGH

1%

N21307 001

Dec 07, 2001

BUTOCONAZOLE NITRATE

CREAM; VAGINAL

FEMSTAT 3

+ BAYER

2%

N20421 001

Dec 21, 1995

CALCIUM CARBONATE; FAMOTIDINE; MAGNESIUM HYDROXIDE

TABLET, CHEWABLE; ORAL

PEPCID COMPLETE

+ MERCK

800MG;10MG;165MG

N20958 001

Oct 16, 2000

CHLORHEXIDINE GLUCONATE

AEROSOL, METERED; TOPICAL

EXIDINE

+ XTTRIUM

4%

N19127 001

Dec 24, 1984

CLOTH; TOPICAL

CHLORHEXIDINE GLUCONATE

+ SAGE PRODS

2%

N21669 001

Apr 25, 2005

SOLUTION; TOPICAL

BRIAN CARE

SOAPCO

4%

N71419 001

Dec 17, 1987

CHG SCRUB

ECOLAB

4%

N19258 002

Jul 22, 1986

CIDA-STAT

ECOLAB

2%

N19258 001

Jul 22, 1986

DYNA-HEX

XTTRIUM

0.75%

N20111 001

Sep 11, 1997

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

OTC DRUG PRODUCT LIST

4 - 3 (of 15)

CHLORHEXIDINE GLUCONATE

SOLUTION; TOPICAL

EXIDINE

+ XTTRIUM	2%	N19422	001	Dec 17, 1985
	4%	N19125	001	Dec 24, 1984
HIBICLENS				
+ REGENT	4%	N17768	001	
HIBISTAT				
+ REGENT	0.5%	N18300	001	
MICRODERM				
J AND J	4%	N72255	001	Apr 15, 1991
PREVACARE R				
J AND J	0.5%	N72292	001	Jan 28, 1992
STERI-STAT				
MATRIX MEDCL	4%	N70104	001	Jul 24, 1986
SPONGE; TOPICAL				
BIOSCRUB				
GRIFFEN	4%	N19822	001	Mar 31, 1989
CHLORHEXIDINE GLUCONATE				
BECTON DICKINSON	4%	N72525	001	Oct 24, 1989
HIBICLENS				
+ REGENT	4%	N18423	001	
MICRODERM				
J AND J	4%	N72295	001	Feb 28, 1991
PHARMASEAL SCRUB CARE				
PHARMASEAL	4%	N19793	001	Dec 02, 1988

CHLORHEXIDINE GLUCONATE; ISOPROPYL ALCOHOL

SPONGE; TOPICAL

CHLORAPREP ONE-STEP

+ MEDI FLEX INC	2%;70% (3ML)	N20832	001	Jul 14, 2000
+	2%;70% (10.5ML)	N20832	004	Aug 20, 2003
+	2%;70% (26ML)	N20832	006	Nov 21, 2006
CHLORAPREP ONE-STEP FREPP				
+ MEDI FLEX INC	2%;70% (1.5ML)	N20832	003	Apr 26, 2002
CHLORAPREP WITH TINT				
+ MEDI FLEX INC	2%;70% (26ML)	N20832	002	May 03, 2005
+	2%;70% (10.5ML)	N20832	005	Apr 03, 2006

SWAB; TOPICAL

CHLORAPREP ONE-STEP SEPP

+ MEDI FLEX INC	2%;70% (0.67ML)	N21555	001	Oct 07, 2002
CHLORAPREP SINGLE SWABSTICK				
+ MEDI FLEX INC	2%;70% (1.75ML)	N21555	002	May 10, 2005
CHLORASCRUB MAXI SWABSTICK				
+ NICE PAK	3.15%;70% (5.1ML)	N21524	003	Jun 03, 2005
CHLORASCRUB SWAB				
+ NICE PAK	3.15%;70% (1ML)	N21524	001	Jun 03, 2005
CHLORASCRUB SWABSTICK				
+ NICE PAK	3.15%;70% (1.6ML)	N21524	002	Jun 03, 2005

CHLORPHENIRAMINE MALEATE

TABLET, EXTENDED RELEASE; ORAL

CHLOR-TRIMETON

+ SCHERING PLOUGH	12MG	N07638	002	
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CHLORPHENIRAMINE MALEATE; IBUPROFEN; PSEUDOEPHEDRINE HYDROCHLORIDE

SUSPENSION; ORAL

CHILDREN'S ADVIL ALLERGY SINUS

+ WYETH CONS	1MG/5ML;100MG/5ML;15MG/5ML	N21587	001	Feb 24, 2004
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TABLET; ORAL

ADVIL ALLERGY SINUS

+ WYETH CONS	2MG;200MG;30MG	N21441	001	Dec 19, 2002
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27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

OTC DRUG PRODUCT LIST

4 - 4 (of 15)

CHLORPHENIRAMINE MALEATE; PSEUDOEPHEDRINE SULFATE

TABLET, EXTENDED RELEASE; ORAL

CHLOR-TRIMETON

+ SCHERING PLOUGH 8MG;120MG N18397 001

CIMETIDINE

TABLET; ORAL

CIMETIDINE

IVAX PHARMS 200MG N75345 001 Jun 16, 1999

LEINER 200MG N74961 001 Jun 19, 1998

200MG N74963 001 Jun 19, 1998

LEK PHARMS 200MG N75122 002 Jun 19, 1998

PERRIGO 200MG N75285 001 Oct 29, 1998

TORPHARM 100MG N74948 001 Jun 19, 1998

200MG N74948 002 Jul 26, 2002

WATSON LABS 200MG N75425 001 Jul 29, 1999

TAGAMET HB

+ GLAXOSMITHKLINE 200MG N20238 002 Aug 21, 1996

CLEMASTINE FUMARATE

TABLET; ORAL

CLEMASTINE FUMARATE

PERRIGO 1.34MG N74512 001 Nov 22, 1995

SANDOZ 1.34MG N73458 001 Oct 31, 1993

TEVA 1.34MG N73282 002 Dec 03, 1992

TAVIST-1

+ NOVARTIS 1.34MG N20925 001 Aug 21, 1992

CLOTRIMAZOLE

CREAM, TABLET; TOPICAL, VAGINAL

GYNE-LOTRIMIN 3 COMBINATION PACK

+ SCHERING PLOUGH 1%,200MG N20526 002 Jul 29, 1996

GYNE-LOTRIMIN COMBINATION PACK

+ SCHERING PLOUGH 1%,100MG N20289 002 Apr 26, 1993

MYCELEX-7 COMBINATION PACK

BAYER PHARMS 1%,100MG N20389 002 Jun 23, 1994

CREAM; VAGINAL

CLOTRIMAZOLE

ACTAVIS MID ATLANTIC 1% N74165 001 Jul 16, 1993

TARO 1% N72641 001 Dec 04, 1995

GYNE-LOTRIMIN

+ SCHERING PLOUGH 1% N18052 002 Nov 30, 1990

GYNE-LOTRIMIN 3

+ SCHERING PLOUGH 2% N20574 001 Nov 24, 1998

MYCELEX-7

BAYER PHARMS 1% N18230 002 Dec 26, 1991

TRIVAGIZOLE 3

TARO 2% N21143 001 Apr 12, 2000

TABLET; VAGINAL

GYNE-LOTRIMIN

+ SCHERING PLOUGH 100MG N17717 002 Nov 30, 1990

GYNE-LOTRIMIN 3

+ SCHERING PLOUGH 200MG N20525 001 Jul 29, 1996

GYNIX

TEVA PHARMS 100MG N73249 001 Feb 13, 1998

MYCELEX-7

BAYER PHARMS 100MG N18182 002 Dec 26, 1991

CROMOLYN SODIUM

SPRAY, METERED; NASAL

CROMOLYN SODIUM

ACTAVIS MID ATLANTIC 5.2MG/SPRAY N74800 001 Jul 26, 2001

+ BAUSCH AND LOMB 5.2MG/SPRAY N75702 001 Jul 03, 2001

PERRIGO 5.2MG/SPRAY N75427 001 Dec 12, 2001

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

OTC DRUG PRODUCT LIST

4 - 5 (of 15)

DEXBROMPHENIRAMINE MALEATE; PSEUDOEPHEDRINE SULFATE

TABLET, EXTENDED RELEASE; ORAL

DISOPHROL

SCHERING PLOUGH 6MG;120MG

N13483 004 Sep 13, 1982

DRIXORAL

+ SCHERING PLOUGH 6MG;120MG

N13483 003 Sep 13, 1982

DEXTROMETHORPHAN HYDROBROMIDE; GUAIFENESIN

TABLET, EXTENDED RELEASE; ORAL

MUCINEX DM

ADAMS RESP THERAP 30MG;600MG

N21620 002 Apr 29, 2004

+ 60MG;1.2GM

N21620 001 Apr 29, 2004

DEXTROMETHORPHAN POLISTIREX

SUSPENSION, EXTENDED RELEASE; ORAL

DELSYM

+ ADAMS RESP THERAP EQ 30MG HBR/5ML

N18658 001 Oct 08, 1982

DIPHENHYDRAMINE CITRATE; IBUPROFEN

TABLET; ORAL

ADVIL PM

+ WYETH CONS 38MG;200MG

N21394 001 Dec 21, 2005

DIPHENHYDRAMINE HYDROCHLORIDE; IBUPROFEN POTASSIUM

CAPSULE; ORAL

ADVIL PM

+ WYETH CONS 25MG;200MG

N21393 001 Dec 21, 2005

DOCOSANOL

CREAM; TOPICAL

ABREVA

+ GLAXOSMITHKLINE 10%

N20941 001 Jul 25, 2000

DOXYLAMINE SUCCINATE

TABLET; ORAL

DOXYLAMINE SUCCINATE

COPLY PHARM 25MG

N88900 002 Feb 12, 1988

LNK 25MG

N40564 001 Aug 27, 2004

PERRIGO 25MG

N40167 001 Sep 18, 1996

UNISOM

+ PFIZER 25MG

N18066 001

EPINEPHRINE

AEROSOL, METERED; INHALATION

EPINEPHRINE

ARMSTRONG PHARMS 0.2MG/INH

N87907 001 May 23, 1984

PRIMATENE MIST

+ WYETH CONS 0.2MG/INH

N16126 001

EPINEPHRINE BITARTRATE

AEROSOL, METERED; INHALATION

BRONITIN MIST

WYETH CONS 0.3MG/INH

N16126 002

FAMOTIDINE

TABLET, CHEWABLE; ORAL

FAMOTIDINE

PERRIGO 10MG

N75715 001 Aug 22, 2003

PEPCID AC

+ MERCK 10MG

N20801 001 Sep 24, 1998

TABLET; ORAL

FAMOTIDINE

DR REDDYS LABS LTD 10MG

N75758 001 Aug 17, 2001

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

OTC DRUG PRODUCT LIST

4 - 6 (of 15)

FAMOTIDINE

TABLET; ORAL

FAMOTIDINE

DR REDDYS LABS LTD	20MG	N77367	001	Sep 25, 2006
GENPHARM	10MG	N75674	001	Dec 21, 2001
IVAX PHARMS	10MG	N75512	001	Jul 26, 2001
PERRIGO	10MG	N75400	001	Mar 18, 2005
	20MG	N77351	001	Sep 25, 2006
SANDOZ	10MG	N76101	001	Oct 21, 2002
TEVA	10MG	N75312	001	May 31, 2001
TORPHARM	10MG	N75610	001	Mar 12, 2002
WATSON LABS	10MG	N75404	001	Nov 28, 2001
WOCKHARDT	10MG	N77146	001	Mar 07, 2005
PEPCID AC				
MERCK	10MG	N20325	001	Apr 28, 1995
+	20MG	N20325	002	Sep 23, 2003
PEPCID AC (GELTAB)				
MERCK	10MG	N20902	001	Aug 05, 1999

GUAIFENESIN

TABLET, EXTENDED RELEASE; ORAL

HUMABID

+ ADAMS RESP THERAP	1.2GM	N21282	002	Dec 18, 2002
MUCINEX				
ADAMS RESP THERAP	600MG	N21282	001	Jul 12, 2002

GUAIFENESIN; PSEUDOEPHEDRINE HYDROCHLORIDE

TABLET, EXTENDED RELEASE; ORAL

MUCINEX D

ADAMS RESP THERAP	600MG;60MG	N21585	001	Jun 22, 2004
+	1.2GM;120MG	N21585	002	Jun 22, 2004

IBUPROFEN

CAPSULE; ORAL

IBUPROFEN

+ BANNER PHARMACAPS	200MG	N21472	001	Oct 18, 2002
+ LEINER	200MG	N74782	001	Jul 06, 1998

SUSPENSION/DROPS; ORAL

CHILDREN'S MOTRIN

+ MCNEIL CONS	40MG/ML	N20603	001	Jun 10, 1996
IBUPROFEN				
PERRIGO	40MG/ML	N75217	001	Dec 16, 1998
PEDIATRIC ADVIL				
+ WYETH CONS	100MG/2.5ML	N20812	001	Jan 30, 1998

SUSPENSION; ORAL

CHILDREN'S ADVIL

WYETH CONS	100MG/5ML	N20589	001	Jun 27, 1996
CHILDREN'S ADVIL-FLAVORED				
WYETH CONS	100MG/5ML	N20589	002	Nov 07, 1997
CHILDREN'S ELIXSURE				
ALTERNA-TCHP LLC	100MG/5ML	N21604	001	Jan 07, 2004
CHILDREN'S IBUPROFEN				
PERRIGO	100MG/5ML	N74937	001	Dec 22, 1998
CHILDREN'S MOTRIN				
+ MCNEIL	100MG/5ML	N20516	001	Jun 16, 1995
IBUPROFEN				
ACTAVIS MID ATLANTIC	100MG/5ML	N74916	001	Apr 30, 1999
TABLET, CHEWABLE; ORAL				
CHILDREN'S ADVIL				
WYETH CONS	50MG	N20944	001	Dec 18, 1998
CHILDREN'S MOTRIN				
MCNEIL CONS	50MG	N20601	001	Nov 15, 1996
IBUPROFEN				
PERRIGO	50MG	N76359	001	Jan 16, 2004

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

OTC DRUG PRODUCT LIST

4 - 7 (of 15)

IBUPROFEN

TABLET, CHEWABLE; ORAL

IBUPROFEN

PERRIGO 100MG N76359 002 Jan 16, 2004

JUNIOR STRENGTH ADVIL

WYETH CONS 100MG N20944 002 Dec 18, 1998

JUNIOR STRENGTH MOTRIN

+ MCNEIL CONS 100MG N20601 003 Nov 15, 1996

TABLET; ORAL

ADVIL

WYETH CONS 200MG N18989 001 May 18, 1984

CAP-PROFEN

PERRIGO 200MG N72097 001 Dec 08, 1987

IBUPROFEN

BASF 200MG N75661 001 Dec 12, 2001

DR REDDYS LABS INC 100MG N76117 001 Nov 20, 2001

INTERPHARM 200MG N71333 001 Feb 17, 1987

200MG N72199 001 May 23, 1988

IVAX PHARMS 200MG N71144 001 Jan 20, 1987

LEINER 200MG N71732 001 Sep 10, 1987

200MG N71735 001 Sep 10, 1987

200MG N72299 001 Jul 01, 1988

200MG N73691 001 Feb 25, 1994

200MG N74931 001 Jul 20, 1998

LNK 100MG N76741 001 Jun 17, 2004

200MG N75010 001 Mar 01, 1999

200MG N75139 001 Mar 01, 1999

MCNEIL 200MG N73019 001 Mar 30, 1994

MUTUAL PHARM 200MG N71229 001 Apr 01, 1987

200MG N72249 001 Jan 10, 1989

MYLAN 200MG N71870 001 May 05, 1988

NEIL 200MG N76460 001 Nov 26, 2003

OHM 200MG N71163 001 Jul 15, 1986

PAR PHARM 200MG N70481 001 Sep 24, 1986

200MG N70985 001 Oct 02, 1987

200MG N71575 001 May 08, 1987

PERRIGO 200MG N72096 001 Dec 08, 1987

200MG N72098 001 Dec 08, 1987

200MG N75995 001 Mar 14, 2002

PERRIGO R AND D 200MG N77349 001 Jun 21, 2005

SANDOZ 200MG N70733 001 Sep 19, 1986

200MG N71807 001 Feb 25, 1988

200MG N74525 001 Dec 15, 1995

200MG N74533 001 Dec 15, 1995

VINTAGE PHARMS 200MG N71639 001 Feb 02, 1988

WATSON LABS 200MG N70435 001 Mar 05, 1986

IBUPROHM

OHM LABS 200MG N71214 001 Dec 01, 1986

IBU-TAB 200

ALRA 200MG N71057 001 Aug 11, 1988

JUNIOR STRENGTH ADVIL

WYETH CONS 100MG N20267 002 Dec 13, 1996

JUNIOR STRENGTH IBUPROFEN

PERRIGO 100MG N75367 001 Apr 22, 1999

JUNIOR STRENGTH MOTRIN

MCNEIL CONS 100MG N20602 001 Jun 10, 1996

MEDIPREN

MCNEIL 200MG N70475 001 Feb 06, 1986

200MG N71215 001 Jun 26, 1986

MOTRIN IB

+ MCNEIL 200MG N19012 003 Dec 17, 1990

MOTRIN MIGRAINE PAIN

+ MCNEIL 200MG N19012 004 Feb 25, 2000

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

OTC DRUG PRODUCT LIST

4 - 8 (of 15)

IBUPROFEN

TABLET; ORAL

PROFEN

LEINER

200MG

N71265 001

Oct 15, 1986

TAB-PROFEN

PERRIGO

200MG

N72095 001

Dec 08, 1987

IBUPROFEN POTASSIUM

CAPSULE; ORAL

ADVIL LIQUI-GELS

+ WYETH CONS

200MG

N20402 001

Apr 20, 1995

ADVIL MIGRAINE LIQUI-GELS

+ WYETH CONS

200MG

N20402 002

Mar 16, 2000

IBUPROFEN POTASSIUM; PSEUDOEPHEDRINE HYDROCHLORIDE

CAPSULE; ORAL

ADVIL COLD AND SINUS

+ WYETH CONS

200MG;30MG

N21374 001

May 30, 2002

IBUPROFEN; PSEUDOEPHEDRINE HYDROCHLORIDE

SUSPENSION; ORAL

CHILDREN'S ADVIL COLD

WYETH CONS

100MG/5ML;15MG/5ML

N21373 001

Apr 18, 2002

CHILDREN'S MOTRIN COLD

+ MCNEIL CONS

100MG/5ML;15MG/5ML

N21128 001

Aug 01, 2000

IBUPROFEN AND PSEUDOEPHEDRINE HYDROCHLORIDE

PERRIGO

100MG/5ML;15MG/5ML

N76478 001

Nov 05, 2003

TABLET; ORAL

ADVIL COLD AND SINUS

+ WYETH CONS

200MG;30MG

N19771 001

Sep 19, 1989

IBUPROFEN AND PSEUDOEPHEDRINE HYDROCHLORIDE

DR REDDYS LABS LTD

200MG;30MG

N77628 001

Aug 14, 2006

LEINER

200MG;30MG

N75588 001

Apr 08, 2002

IBUPROHM COLD AND SINUS

OHM LABS

200MG;30MG

N74567 001

Apr 17, 2001

SINE-AID IB

MCNEIL CONS

200MG;30MG

N19899 001

Dec 31, 1992

INSULIN RECOMBINANT HUMAN

INJECTABLE; INJECTION

HUMULIN R

+ LILLY

100 UNITS/ML

N18780 001

Oct 28, 1982

HUMULIN R PEN

+ LILLY

100 UNITS/ML

N18780 005

Aug 06, 1998

NOVOLIN R

+ NOVO NORDISK INC

100 UNITS/ML

N19938 001

Jun 25, 1991

INSULIN RECOMBINANT HUMAN; INSULIN SUSP ISOPHANE RECOMBINANT HUMAN

INJECTABLE; INJECTION

HUMULIN 50/50

+ LILLY

50 UNITS/ML;50 UNITS/ML

N20100 001

Apr 29, 1992

HUMULIN 70/30

+ LILLY

30 UNITS/ML;70 UNITS/ML

N19717 001

Apr 25, 1989

HUMULIN 70/30 PEN

+ LILLY

30 UNITS/ML;70 UNITS/ML

N19717 002

Aug 06, 1998

NOVOLIN 70/30

+ NOVO NORDISK INC

30 UNITS/ML;70 UNITS/ML

N19991 001

Jun 25, 1991

INSULIN SUSP ISOPHANE RECOMBINANT HUMAN

INJECTABLE; INJECTION

HUMULIN N

+ LILLY

100 UNITS/ML

N18781 001

Oct 28, 1982

NOVOLIN N

NOVO NORDISK INC

100 UNITS/ML

N19959 001

Jul 01, 1991

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

OTC DRUG PRODUCT LIST

4 - 9 (of 15)

INSULIN ZINC SUSP RECOMBINANT HUMAN

INJECTABLE; INJECTION

HUMULIN L

+ LILLY 100 UNITS/ML

N19377 002 Sep 30, 1985

IODINE POVACRYLEX; ISOPROPYL ALCOHOL

SPONGE; TOPICAL

DURAPREP

+ 3M EQ 0.7% IODINE; 74% (6ML)

N21586 001 Sep 29, 2006

+ EQ 0.7% IODINE; 74% (26ML)

N21586 002 Sep 29, 2006

KETOCONAZOLE

SHAMPOO; TOPICAL

NIZORAL A-D

+ MCNEIL CONS 1%

N20310 001 Oct 10, 1997

KETOTIFEN FUMARATE

SOLUTION/DROPS; OPHTHALMIC

ALAWAY

+ ALIMERA SCIENCES INC EQ 0.025% BASE

N21996 001 Dec 01, 2006

ZADITOR

+ NOVARTIS EQ 0.025% BASE

N21066 002 Oct 19, 2006

LEVONORGESTREL

TABLET; ORAL

PLAN B

+ DURAMED 0.75MG

N21045 002 Aug 24, 2006

LOPERAMIDE HYDROCHLORIDE

CAPSULE; ORAL

LOPERAMIDE HYDROCHLORIDE

BANNER PHARMACAPS 1MG

N21855 001 Aug 04, 2005

+ 2MG

N21855 002 Aug 04, 2005

SOLUTION; ORAL

IMODIUM A-D

+ MCNEIL CONS 1MG/5ML

N19487 001 Mar 01, 1988

LOPERAMIDE HYDROCHLORIDE

DURAMED PHARMS BARR 1MG/5ML

N74991 001 Dec 29, 1997

HI TECH PHARMA 1MG/5ML

N74352 001 Nov 17, 1995

MORTON GROVE 1MG/5ML

N74730 001 Aug 28, 1997

PERRIGO 1MG/5ML

N73243 001 Jan 21, 1992

ROXANE 1MG/5ML

N73079 001 Apr 30, 1992

TEVA 1MG/5ML

N73478 001 Jun 23, 1995

SUSPENSION; ORAL

IMODIUM A-D

+ MCNEIL CONS 1MG/7.5ML

N19487 002 Jul 08, 2004

TABLET, CHEWABLE; ORAL

IMODIUM A-D

+ MCNEIL 2MG

N20448 001 Jul 24, 1997

TABLET; ORAL

IMODIUM A-D

+ MCNEIL CONS 2MG

N19860 001 Nov 22, 1989

LOPERAMIDE HYDROCHLORIDE

LEINER 2MG

N73254 001 Jul 30, 1993

LNK 2MG

N76497 001 Jun 10, 2003

OHM LABS 2MG

N74091 001 Dec 10, 1992

PERRIGO 2MG

N75232 001 Jan 06, 2000

LOPERAMIDE HYDROCHLORIDE; SIMETHICONE

TABLET, CHEWABLE; ORAL

IMODIUM ADVANCED

+ MCNEIL 2MG;125MG

N20606 001 Jun 26, 1996

LOPERAMIDE HYDROCHLORIDE AND SIMETHICONE

PERRIGO 2MG;125MG

N76029 001 Aug 30, 2002

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

OTC DRUG PRODUCT LIST

4 - 10 (of 15)

LOPERAMIDE HYDROCHLORIDE; SIMETHICONE

TABLET; ORAL

IMODIUM ADVANCED

+ MCNEIL CONS 2MG;125MG N21140 001 Nov 30, 2000

LOPERAMIDE HYDROCHLORIDE AND SIMETHICONE

RANBAXY 2MG;125MG N77500 001 Sep 06, 2006

LORATADINE

SUSPENSION; ORAL

LORATADINE

+ TARO 1MG/ML N21734 001 Oct 04, 2005

SYRUP; ORAL

CLARITIN

+ SCHERING PLOUGH 1MG/ML N20641 002 Nov 27, 2002

LORATADINE

APOTEX INC 1MG/ML N75565 001 Oct 05, 2004

MORTON GROVE 1MG/ML N75815 001 Aug 20, 2004

PERRIGO 1MG/ML N75728 001 Aug 20, 2004

RANBAXY 1MG/ML N76529 001 Aug 20, 2004

SILARX 1MG/ML N77421 001 Jun 29, 2006

TARO 1MG/ML N76805 001 Aug 20, 2004

TEVA 1MG/ML N75505 001 Nov 07, 2003

TABLET, CHEWABLE; ORAL

CHILDREN'S CLARITIN

+ SCHERING PLOUGH 5MG N21891 001 Aug 23, 2006

TABLET, ORALLY DISINTEGRATING; ORAL

ALAVERT

WYETH CONS 10MG N21375 001 Dec 19, 2002

CLARITIN HIVES RELIEF REDITAB

+ SCHERING PLOUGH 10MG N20704 003 Nov 19, 2003

CLARITIN REDITABS

+ SCHERING PLOUGH 5MG N21993 001 Dec 12, 2006

+ 10MG N20704 002 Nov 27, 2002

LORATADINE

ANDRX PHARMS 10MG N75990 001 Nov 03, 2003

IMPAX LABS 10MG N76011 001 Sep 29, 2003

WYETH CONS 10MG N75822 001 Feb 10, 2003

TABLET; ORAL

CLARITIN

+ SCHERING PLOUGH 10MG N19658 002 Nov 27, 2002

CLARITIN HIVES RELIEF

+ SCHERING PLOUGH 10MG N19658 003 Nov 19, 2003

LORATADINE

APOTEX INC 10MG N76471 001 Feb 14, 2006

GENPHARM 10MG N76154 001 Aug 20, 2003

PERRIGO 10MG N76301 001 Jun 25, 2004

RANBAXY 10MG N76134 001 Aug 18, 2003

SANDOZ 10MG N75209 001 Jan 21, 2003

LORATADINE; PSEUDOEPHEDRINE SULFATE

TABLET, EXTENDED RELEASE; ORAL

CLARITIN-D 24 HOUR

+ SCHERING PLOUGH 10MG;240MG N20470 002 Nov 27, 2002

LORATADINE AND PSEUDOEPHEDRINE SULFATE

ANDRX PHARMS 5MG;120MG N76208 001 Jan 28, 2004

10MG;240MG N75706 001 Feb 21, 2003

+ IMPAX LABS 5MG;120MG N76050 001 Jan 30, 2003

10MG;240MG N75989 001 Mar 04, 2004

RANBAXY 10MG;240MG N76557 001 Sep 22, 2004

MICONAZOLE NITRATE

CREAM, SUPPOSITORY; TOPICAL, VAGINAL

MICONAZOLE 7 COMBINATION PACK

G AND W LABS 2%,100MG N76585 001 Mar 26, 2004

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

OTC DRUG PRODUCT LIST

4 - 11 (of 15)

MICONAZOLE NITRATE

CREAM, SUPPOSITORY; TOPICAL, VAGINAL			
MICONAZOLE NITRATE COMBINATION PACK			
PERRIGO	2%, 200MG	N75329 001	Apr 20, 1999
MONISTAT 1 COMBINATION PACK			
+ PERSONAL PRODS	1.2GM, 2%	N21308 001	Jun 29, 2001
MONISTAT 7 COMBINATION PACK			
+ ADV CARE	2%, 100MG	N20288 002	Apr 26, 1993
MONISTAT-3 COMBINATION PACK			
+ ADV CARE	2%, 200MG	N20670 002	Apr 16, 1996
M-ZOLE 3 COMBINATION PACK			
ACTAVIS MID ATLANTIC	2%, 200MG	N74926 001	Apr 16, 1999
M-ZOLE 7 DUAL PACK			
ACTAVIS MID ATLANTIC	2%, 100MG	N74586 001	Jul 17, 1997
CREAM; TOPICAL, VAGINAL			
MICONAZOLE 3 COMBINATION PACK			
PERRIGO	2%, 4%	N76357 001	Mar 30, 2004
MONISTAT 3 COMBINATION PACK			
PERSONAL PRODS	2%, 4%	N21261 003	Jun 17, 2003
MONISTAT 3 COMBINATION PACK (PREFILLED)			
+ PERSONAL PRODS	2%, 4%	N21261 001	Feb 02, 2001
CREAM; VAGINAL			
MICONAZOLE 3			
TARO	4%	N76773 001	Mar 02, 2005
MICONAZOLE 7			
ACTAVIS MID ATLANTIC	2%	N74164 001	Mar 29, 1996
MICONAZOLE NITRATE			
G AND W LABS	2%	N74366 001	Feb 22, 1996
PERRIGO	2%	N74760 001	May 15, 1997
TARO	2%	N74444 001	Jan 13, 1997
TEVA	2%	N74136 001	Jan 04, 1995
TEVA PHARMS	2%	N74030 001	Oct 30, 1992
MONISTAT 3			
+ ADV CARE PRODS	4%	N20827 001	Mar 30, 1998
MONISTAT 7			
+ ADV CARE	2%	N17450 002	Feb 15, 1991
SUPPOSITORY; VAGINAL			
MICONAZOLE NITRATE			
ACTAVIS MID ATLANTIC	100MG	N73507 001	Nov 19, 1993
G AND W LABS	100MG	N74414 001	Apr 30, 1997
+ PERRIGO	100MG	N74395 001	Mar 20, 1997
MONISTAT 7			
+ ADV CARE PRODS	100MG	N18520 002	Feb 15, 1991

MINOXIDIL

AEROSOL, FOAM; TOPICAL			
MEN'S ROGAINE			
+ JOHNSON AND JOHNSON	5%	N21812 001	Jan 20, 2006
SOLUTION; TOPICAL			
MINOXIDIL (FOR MEN)			
ACTAVIS MID ATLANTIC	2%	N74588 001	Apr 05, 1996
BAUSCH AND LOMB	2%	N74643 001	Apr 09, 1996
COPLEY PHARM	2%	N74500 001	May 23, 1996
HI TECH PHARMA	2%	N74731 001	Dec 24, 1996
MORTON GROVE	2%	N74767 001	Feb 28, 1997
NOVEX	2%	N74924 001	Apr 29, 1998
PERRIGO	2%	N75357 001	Jul 30, 1999
SIGHT PHARMS	2%	N74743 002	Oct 18, 1996
TEVA	2%	N74589 001	Apr 05, 1996
MINOXIDIL (FOR WOMEN)			
NOVEX	2%	N74924 002	Apr 29, 1998
PERRIGO	2%	N75357 002	Jul 30, 1999
SIGHT PHARMS	2%	N74743 001	Oct 18, 1996
MINOXIDIL EXTRA STRENGTH (FOR MEN)			
ACTAVIS MID ATLANTIC	5%	N75518 001	Nov 17, 2000

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

OTC DRUG PRODUCT LIST

4 - 12 (of 15)

MINOXIDIL

SOLUTION; TOPICAL

MINOXIDIL EXTRA STRENGTH (FOR MEN)

MORTON GROVE	5%	N75438	001	Feb 27, 2003
NOVEX	5%	N75839	001	Oct 01, 2001
PERRIGO	5%	N75598	001	Jun 13, 2001
PERRIGO NEW YORK	5%	N75737	001	Mar 15, 2002
TEVA	5%	N75619	001	Nov 17, 2000
ROGAINE (FOR MEN)				
+ JOHNSON AND JOHNSON	2%	N19501	002	Feb 09, 1996
ROGAINE (FOR WOMEN)				
+ JOHNSON AND JOHNSON	2%	N19501	003	Feb 09, 1996
ROGAINE EXTRA STRENGTH (FOR MEN)				
+ JOHNSON AND JOHNSON	5%	N20834	001	Nov 14, 1997
THEROXIDIL				
HARMONY LABS	5%	N76239	001	Aug 24, 2004

NAPHAZOLINE HYDROCHLORIDE; PHENIRAMINE MALEATE

SOLUTION/DROPS; OPHTHALMIC

NAPHCN-A

+ ALCON	0.025%;0.3%	N20226	001	Jun 08, 1994
OPCN-A				
+ BAUSCH AND LOMB	0.02675%;0.315%	N20065	001	Jun 08, 1994
VISINE-A				
JOHNSON AND JOHNSON	0.025%;0.3%	N20485	001	Jan 31, 1996

NAPROXEN SODIUM

CAPSULE; ORAL

NAPROXEN SODIUM

+ BANNER PHARMACAPS	EQ 200MG BASE	N21920	001	Feb 17, 2006
TABLET; ORAL				
ALEVE				
+ BAYER	EQ 200MG BASE	N20204	002	Jan 11, 1994
NAPROXEN SODIUM				
DR REDDYS LABS INC	EQ 200MG BASE	N75168	001	Jul 28, 1998
LEINER	EQ 200MG BASE	N74635	001	Jan 13, 1997
	EQ 200MG BASE	N74789	001	Feb 27, 1997
PERRIGO	EQ 200MG BASE	N74661	001	Jan 13, 1997
SANDOZ	EQ 200MG BASE	N74646	001	Jan 13, 1997

NAPROXEN SODIUM; PSEUDOEPHEDRINE HYDROCHLORIDE

TABLET, EXTENDED RELEASE; ORAL

ALEVE COLD AND SINUS

+ BAYER	EQ 200MG BASE;120MG	N21076	001	Nov 29, 1999
NAPROXEN SODIUM AND PSEUDOEPHEDRINE HYDROCHLORIDE				
DR REDDYS LABS INC	EQ 220MG BASE;120MG	N77381	001	Sep 27, 2006
PERRIGO	EQ 200MG BASE;120MG	N76518	001	Mar 17, 2004

NICOTINE

FILM, EXTENDED RELEASE; TRANSDERMAL

HABITROL

+ NOVARTIS	7MG/24HR	N20076	004	Nov 12, 1999
+	14MG/24HR	N20076	005	Nov 12, 1999
+	21MG/24HR	N20076	006	Nov 12, 1999
NICODERM CQ				
+ SANOFI AVENTIS US	7MG/24HR	N20165	006	Aug 02, 1996
+	14MG/24HR	N20165	005	Aug 02, 1996
+	21MG/24HR	N20165	004	Aug 02, 1996
PROSTEP				
+ AVEVA	11MG/24HR	N19983	003	Dec 23, 1998
+	22MG/24HR	N19983	004	Dec 23, 1998

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

OTC DRUG PRODUCT LIST

4 - 13 (of 15)

NICOTINE POLACRILEXGUM, CHEWING; BUCCAL
NICORETTE

+	GLAXOSMITHKLINE	EQ 2MG BASE	N18612	002	Feb 09, 1996
		EQ 2MG BASE	N18612	004	Sep 25, 2000
+		EQ 4MG BASE	N20066	002	Feb 09, 1996
		EQ 4MG BASE	N20066	004	Sep 25, 2000

NICORETTE (MINT)

GLAXOSMITHKLINE	EQ 2MG BASE	N18612	003	Dec 23, 1998
	EQ 4MG BASE	N20066	003	Dec 23, 1998

NICOTINE POLACRILEX
PERRIGO

EQ 2MG BASE	N76775	001	Sep 16, 2004	
EQ 2MG BASE	N76776	001	Sep 16, 2004	
EQ 2MG BASE	N76777	001	Sep 16, 2004	
EQ 4MG BASE	N76778	001	Sep 16, 2004	
EQ 4MG BASE	N76779	001	Sep 16, 2004	
EQ 4MG BASE	N76789	001	Sep 16, 2004	
PERRIGO R AND D	EQ 2MG BASE	N78325	001	Oct 30, 2006
	EQ 4MG BASE	N78326	001	Oct 30, 2006
WATSON LABS	EQ 2MG BASE	N74507	001	Mar 15, 1999
	EQ 2MG BASE	N76569	001	Jul 29, 2004
	EQ 4MG BASE	N74707	001	Mar 19, 1999
	EQ 4MG BASE	N76568	002	Jul 29, 2004

TROCHE/LOZENGE; ORAL
COMMIT

GLAXOSMITHKLINE CONS	EQ 2MG BASE	N21330	001	Oct 31, 2002
+	EQ 4MG BASE	N21330	002	Oct 31, 2002
NICOTINE POLACRILEX				
PERRIGO R AND D	EQ 2MG BASE	N77007	001	Jan 31, 2006
	EQ 4MG BASE	N77007	002	Jan 31, 2006

NIZATIDINETABLET; ORAL
AXID AR

+	WYETH CONS	75MG	N20555	001	May 09, 1996
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OMEPRAZOLE MAGNESIUMTABLET, DELAYED RELEASE; ORAL
PRILOSEC OTC

+	ASTRAZENECA	EQ 20MG BASE	N21229	001	Jun 20, 2003
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OXYMETAZOLINE HYDROCHLORIDESOLUTION/DROPS; OPHTHALMIC
OCUCLEAR

SCHERING PLOUGH	0.025%	N18471	001	May 30, 1986	
VISINE L.R.					
+	JOHNSON AND JOHNSON	0.025%	N19407	001	Mar 31, 1989

PERMETHRINLOTION; TOPICAL
NIX

+	INSIGHT PHARMS	1%	N19918	001	May 02, 1990
PERMETHRIN					
ACTAVIS MID ATLANTIC	1%	N75014	001	Mar 28, 2000	
PERRIGO NEW YORK	1%	N76090	001	Dec 20, 2001	

PIPERONYL BUTOXIDE; PYRETHRINSAEROSOL; TOPICAL
RID MOUSSE

+	PFIZER	4%;EQ 0.33% BASE	N21043	001	Mar 07, 2000
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27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

OTC DRUG PRODUCT LIST

4 - 14 (of 15)

POLYETHYLENE GLYCOL 3350FOR SOLUTION; ORAL
MIRALAX

+ SCHERING PLOUGH 17GM/SCOOPFUL N22015 001 Oct 06, 2006

POTASSIUM IODIDESOLUTION; ORAL
THYROSHIELD

FLEMING 65MG/ML N77218 001 Jan 12, 2005

TABLET; ORAL
IOSAT

+ ANBEX 130MG N18664 001 Oct 14, 1982

THYROSAFE

+ R R REGISTRATIONS 65MG N76350 001 Sep 10, 2002

POVIDONE-IODINESOLUTION; TOPICAL
POVIDONE IODINE

+ ALLEGIANCE HLTHCARE 1% N19522 001 Mar 31, 1989

SPONGE; TOPICAL

E-Z SCRUB 201

+ BECTON DICKINSON 20% N19240 001 Nov 29, 1985

E-Z SCRUB 241

+ BECTON DICKINSON 10% N19476 001 Jan 07, 1987

PSEUDOEPHEDRINE HYDROCHLORIDE

TABLET, EXTENDED RELEASE; ORAL

EFIDAC 24 PSEUDOEPHEDRINE HYDROCHLORIDE

+ ALZA 240MG N20021 002 Dec 15, 1992

PSEUDOEPHEDRINE HYDROCHLORIDE

PERRIGO 120MG N75153 001 Feb 26, 1999

RANBAXY 120MG N77442 001 Sep 28, 2005

SUDAFED 12 HOUR

+ WARNER LAMBERT 120MG N73585 001 Oct 31, 1991

PSEUDOEPHEDRINE SULFATE

TABLET, EXTENDED RELEASE; ORAL

AFRINOL

+ SCHERING PLOUGH 120MG N18191 001

RANITIDINE HYDROCHLORIDE

TABLET; ORAL

RANITIDINE

DR REDDYS LABS LTD EQ 75MG BASE N75294 001 Mar 28, 2000

GENPHARM EQ 75MG BASE N75497 001 Jan 14, 2000

IVAX PHARMS EQ 75MG BASE N75296 001 Jan 14, 2000

LEINER EQ 75MG BASE N75094 001 Jun 21, 1999

PERRIGO EQ 75MG BASE N76195 001 Aug 30, 2002

RANBAXY EQ 75MG BASE N75132 001 Jan 14, 2000

SANDOZ EQ 75MG BASE N75519 001 Sep 26, 2002

TORPHARM EQ 75MG BASE N75167 001 May 04, 2000

WATSON LABS EQ 75MG BASE N75212 001 Jan 14, 2000

WOCKHARDT EQ 75MG BASE N76760 001 Feb 24, 2006

ZANTAC 150

+ MCNEIL CONS EQ 150MG BASE N21698 001 Aug 31, 2004

ZANTAC 75

MCNEIL CONS EQ 75MG BASE N20520 001 Dec 19, 1995

SODIUM CHLORIDE

AEROSOL, METERED; INHALATION

BRONCHO SALINE

+ BLAIREX 0.9% N19912 001 Sep 03, 1992

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

OTC DRUG PRODUCT LIST

4 - 15 (of 15)

SODIUM FLUORIDE; TRICLOSAN

PASTE; DENTAL				
COLGATE TOTAL				
+ COLGATE PALMOLIVE	0.24%;0.3%	N20231	001	Jul 11, 1997

SODIUM MONOFLUOROPHOSPHATE

GEL; DENTAL				
EXTRA-STRENGTH AIM				
+ CHESEBROUGH PONDS	1.2%	N19518	002	Aug 06, 1986
PASTE; DENTAL				
EXTRA-STRENGTH AIM				
+ CHESEBROUGH PONDS	1.2%	N19518	001	Jun 03, 1987

TERBINAFINE

GEL; TOPICAL				
LAMISIL AT				
+ NOVARTIS	1%	N21958	001	Jul 24, 2006

TERBINAFINE HYDROCHLORIDE

CREAM; TOPICAL				
LAMISIL				
+ NOVARTIS	1%	N20980	001	Mar 09, 1999
SOLUTION; TOPICAL				
LAMISIL AT				
+ NOVARTIS	1%	N21124	001	Mar 17, 2000
SPRAY; TOPICAL				
LAMISIL AT				
+ NOVARTIS	1%	N21124	002	Mar 17, 2000

TIOCONAZOLE

OINTMENT; VAGINAL				
TIOCONAZOLE				
PERRIGO	6.5%	N75915	001	Nov 21, 2001
VAGISTAT-1				
+ NOVARTIS	6.5%	N20676	001	Feb 11, 1997

27TH EDITION - 2007 - APPROVED DRUG PRODUCT LIST

5 - 1 of 5

**DRUG PRODUCTS WITH APPROVAL UNDER SECTION 505 OF THE ACT
ADMINISTERED BY THE CENTER FOR BIOLOGICS EVALUATION AND RESEARCH LIST**

ANTICOAGULANT 4% SODIUM CITRATE SOLUTION USP

INJECTABLE; INJECTION

NONE

MEDSHEP CORPORATION

N760305

Jun 30, 1978

ANTICOAGULANT CITRATE DEXTROSE SOLUTION (ACD)

INJECTABLE; INJECTION

CYTOSOL

LABORATORIES INC

N020037

Aug 26, 2003

ACD-A SOLUTION

GAMBRO BCT INC

N010228

ANDA

Feb 25, 2002

ADSOL WITH ACD-A

BAXTER HEALTHCARE

CORP FENWAL DIVISION

N000922

Aug 29, 2002

ANTICOAGULANT CITRATE DEXTROSE SOLUTION FORMULA A

HAEMONETICS CORP

N980728

ANDA

Feb 06, 2002

AS3 SOLUTION/ACD-A

GAMBRO BCT INC

N001214

May 29, 2002

ANTICOAGULANT CITRATE DEXTROSE SOLUTION USP

INJECTABLE; INJECTION

NONE

BAXTER HEALTHCARE

CORP - FENWAL

DIVISION

N100855

Jun 03, 1959

FRESENIUS USA INC

N160918

Mar 17, 1978

MEDSEP CORPORATION

N110912

Sep 02, 1959

MILES INC

N710497

ANDA

N100102

Dec 14, 1961

ANTICOAGULANT CITRATE PHOSPHATE 2X DEXTROSE SOLUTION (CP2D)

INJECTABLE; INJECTION

CITRATE PHOSPHATE DOUBLE DEXTROSE/ADDITIVE SOLUTION 3

HAEMONETICS CORP

N000127

Jan 18, 2002

ANTICOAGULANT CITRATE PHOSPHATE DEXTROSE ADENINE SOLUTION

INJECTABLE; INJECTION

NONE

FRESENIUS USA INC

N780519

Apr 23, 1980

TERUMO MEDICAL CORP

N820528

Nov 03, 1982

ANTICOAGULANT CITRATE PHOSPHATE DEXTROSE ADENINE SOLUTION (CPDA)

INJECTABLE; INJECTION

CPDA-1 BLOOD-PACK UNIT (PL 146 PLASTIC) 250, 450, 500 ML BLOOD PACK UNITS

BAXTER HEALTHCARE

N770420

May 12, 1978

CORP

ANTICOAGULANT CITRATE PHOSPHATE DEXTROSE ADENINE SOLUTION USP

INJECTABLE; INJECTION

BLOOD PACK UNIT CPDA-1 IN PLASTIC CONTAINER

BAXTER HEALTHCARE

N940404

Jul 28, 1994

CORP - FENWAL

DIVISION

ANTICOAGULANT CITRATE PHOSPHATE DEXTROSE ADENINE-1 SOLUTION

INJECTABLE; INJECTION

NONE

MEDSEP CORPORATION

N800077

Nov 06, 1980

27TH EDITION - 2007 - APPROVED DRUG PRODUCT LIST

5 - 2 of 5

**DRUG PRODUCTS WITH APPROVAL UNDER SECTION 505 OF THE ACT
ADMINISTERED BY THE CENTER FOR BIOLOGICS EVALUATION AND RESEARCH LIST**

ANTICOAGULANT CITRATE PHOSPHATE DEXTROSE SOLUTION

INJECTABLE; INJECTION		
ADSOL IN PLASTIC CONTAINER		
BAXTER HEALTHCARE	N900223	Dec 27, 1991
CORP - FENWAL		
DIVISION		
CDP BLOOD BAG UNIT IN PLASTIC CONTAINER		
BAXTER HLTHCARE -	N900224	Dec 27, 1991
FENWAL DIVISION		

ANTICOAGULANT CITRATE PHOSPHATE DEXTROSE SOLUTION (CPD)

INJECTABLE; INJECTION		
MACOPRODUCTIONS SAS CDP/AS-1: MACOPHARMA LEUCOFLEX MTL1 LEUKOREDUCTION SYSTEM FOR BLOOD		
COMPONENTS KNOWN AS MTL1-WB		
MACOPRODUCTIONS SAS	N040083	Nov 21, 2005

ANTICOAGULANT CITRATE PHOSPHATE DEXTROSE SOLUTION USP

INJECTABLE; INJECTION		
NONE		
BAXTER HEALTHCARE	N170401	Dec 06, 1977
CORP - FENWAL		
DIVISION		
FRESENIUS USA INC	N811012	Jun 28, 1983
MEDSEP CORPORATION	N160907	May 16, 1973
MILES INC	N800222	Aug 23, 1982
TERUMO MEDICAL CORP	N160527	Jun 22, 1970
	N781211	Jun 10, 1981

ANTICOAGULANT CITRATE PHOSPHATE DEXTROSE SOLUTION USP WITH: AS-1:
DEXTROSE USP; SODIUM CHLORIDE USP; MANNITOL USP; ADENINE

INJECTABLE; INJECTION		
ADSOL RED BLOOD CELL PRESERVATIVE SOLUTION		
BAXTER HEALTHCARE	N811104	May 16, 1983
CORP - FENWAL		
DIVISION		

ANTICOAGULANT CITRATE PHOSPHATE DEXTROSE SOLUTION USP WITH: AS-5:
DEXTROSE USP; SODIUM CHLORIDE USP; MANNITOL USP; ADENINE

INJECTABLE; INJECTION		
OPTISOL RED BLOOD CELL PRESERVATIVE SOLUTION		
TERUMO MEDICAL CORP	N880217	Oct 07, 1988

ANTICOAGULANT CITRATE PHOSPHATE DOUBLE DEXTROSE SOLUTION WITH:
AS-3: CITRIC ACID USP; MONOBASIC SODIUM PHOSPHATE USP; SODIUM CHLORIDE USP; ADENINE;
DEXTROSE USP; SODIUM CITRATE USP

INJECTABLE; INJECTION		
AS-3 NUTRICEL ADDITIVE SYSTEM		
MEDSEP CORP	0.042GM/100ML;0.276GM/100ML; 0.410GM/100ML;0.30GM/100ML; 1.10GM/100ML;0.588GM/100ML	N820915 Oct 19, 1984

ANTICOAGULANT CITRATE PHOSPHATE DOUBLE DEXTROSE SOLUTION WITH:
AS-2: CITRIC ACID USP; DIBASIC SODIUM PHOSPHATE USP; SODIUM CHLORIDE USP; ADENINE;
DEXTROSE USP; SODIUM CITRATE USP

INJECTABLE; INJECTION		
AS-2 NUTRICEL ADDITIVE SYSTEM		
MEDSEP CORP	0.042GM/100ML;0.285GM/100ML; 0.718GM/100ML;0.017GM/100ML; 0.396GM/100ML;0.588GM/100ML	N820915 Sep 22, 1983

27TH EDITION - 2007 - APPROVED DRUG PRODUCT LIST

5 - 3 of 5

**DRUG PRODUCTS WITH APPROVAL UNDER SECTION 505 OF THE ACT
ADMINISTERED BY THE CENTER FOR BIOLOGICS EVALUATION AND RESEARCH LIST**

ANTICOAGULANT ETHYLENEDIAMINE TETRAACETIC ACID (EDTA)

INJECTABLE; INJECTION

NONE

BAXTER HEALTHCARE
CORP

N090480

Mar 07, 1955

ANTICOAGULANT HEPARIN SOLUTION USP

INJECTABLE; INJECTION

NONE

FRESENIUS USA INC

N770822

May 17, 1978

ANTICOAGULANT SODIUM CITRATE 4% SOLUTION

INJECTABLE; INJECTION

NONE

HAEMONETICS
CORPORATION

N980123

Mar 03, 2000

ANTICOAGULANT SODIUM CITRATE SOLUTION

INJECTABLE; INJECTION

TRICITRASOL

CYTOSOL
LABORATORIES INC

N010409

Jul 10, 2003

ANTICOAGULANT SODIUM CITRATE SOLUTION USP

INJECTABLE; INJECTION

NONE

BAXTER HEALTHCARE
CORP - FENWAL
DIVISION

N770923

Jan 20, 1978

FRESENIUS USA INC

N160702

Dec 28, 1970

TERUMO MEDICAL CORP

N781214

Feb 08, 1980

DEXTRAN 1 IN SODIUM CHLORIDE 0.6%

INJECTABLE; INJECTION

PROMIT

PHARMALINK
BASLAKEMEDEL AB

N830715

Oct 30, 1984

DEXTRAN 40, 10% IN DEXTROSE 5%

INJECTABLE; INJECTION

GENTRAN 40

BAXTER HLTHCARE IV 10GM/100ML;5GM/100ML
SYSTEMS

N840619

Feb 22, 1985

LMD IN PLASTIC CONTAINER

ABBOTT LABS 10GM/100ML;5GM/100ML

N720563

ANDA

Oct 30, 1992

NONE

ABBOTT LABS 10GM/100ML;5GM/100ML

N160375

Jul 25, 1967

B BRAUN MEDICAL INC 10GM/100ML;5GM/100ML

N160767

Apr 06, 1970

MILES INC 10GM/100ML;5GM/100ML

N160653

Sep 23, 1969

PHARMACHEM 10GM/100ML;5GM/100ML

N160836

Nov 14, 1972

RHEOMACRODEX

MEDA AB 10GM/100ML;5GM/100ML

N140716

Jan 18, 1967

27TH EDITION - 2007 - APPROVED DRUG PRODUCT LIST

5 - 4 of 5

**DRUG PRODUCTS WITH APPROVAL UNDER SECTION 505 OF THE ACT
ADMINISTERED BY THE CENTER FOR BIOLOGICS EVALUATION AND RESEARCH LIST**

DEXTRAN 40, 10% IN SODIUM CHLORIDE 0.9%

INJECTABLE; INJECTION

GENTRAN 40

BAXTER HLTHCARE IV SYSTEMS	10GM/100ML;0.9GM/100ML	N840620	Feb 22, 1985
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LMD IN PLASTIC CONTAINER

ABBOTT LABS	10GM/100ML;0.9GM/100ML	N720562	ANDA Oct 30, 1992
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NONE

ABBOTT LABS	10GM/100ML;0.9GM/100ML	N160375	Jul 25, 1967
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MCGAW INC	10GM/100ML;0.9GM/100ML	N160767	Apr 06, 1970
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MILES INC	10GM/100ML;0.9GM/100ML	N160653	Sep 23, 1969
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PHARMACHEM	10GM/100ML;0.9GM/100ML	N160836	Nov 14, 1972
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RHEOMACRODEX

PHARMALINK	10GM/100ML;0.9GM/100ML	N140716	Jan 18, 1967
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BASLAKEMEDEL AB

DEXTRAN 40, 10% W/V DEXTRAN 40 IN 0.8% NACL 500 ML PVC BAGS

INJECTABLE; INJECTION

RHEOMACRODEX

PHARMALINK	N830527	Mar 27, 1985
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BASLAKEMEDEL AB

DEXTRAN 40, 10% W/V DEXTRAN 40 IN 5% DEXTROSE 500 ML PVC BAGS

INJECTABLE; INJECTION

RHEOMACRODEX

PHARMALINK	N830627	Mar 27, 1985
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BASLAKEMEDEL AB

DEXTRAN 70, 6% IN DEXTROSE 5%

INJECTABLE; INJECTION

MACRODEX

MEDA AB	6GM/100ML;5GM/100ML	N060826	Jun 08, 1954
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NONE

ABBOTT LABS	6GM/100ML;5GM/100ML	N080819	Mar 31, 1953
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DEXTRAN 70, 6% IN SODIUM CHLORIDE 0.9%

INJECTABLE; INJECTION

GENTRAIN 70

BAXTER HLTHCARE IV SYSTEMS	N160607	Jan 26, 1970
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MACRODEX

PHARMALINK	6GM/100ML;5GM/100ML	N060826	Jun 08, 1954
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BASLAKEMEDEL AB

NONE

MCGAW INC	6GM/100ML;0.9 GM/100ML	N090024	Aug 18, 1969
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MILES INC	6GM/100ML;0.9 GM/100ML	N080716	Mar 13, 1953
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DEXTRAN 70, 6% W/V DEXTRAN 70 IN 0.9% NACL IN 500 ML PVC BAGS

INJECTABLE; INJECTION

MACRODEX

PHARMALINK	N830613	Mar 27, 1985
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BASLAKEMEDEL AB

DEXTRAN 70, 6% W/V DEXTRAN 70 IN 5% DEXTROSE

INJECTABLE; INJECTION

MACRODEX

PHARMALINK	N830629	Mar 27, 1985
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BASLAKEMEDEL AB

27TH EDITION - 2007 - APPROVED DRUG PRODUCT LIST

5 - 5 of 5

**DRUG PRODUCTS WITH APPROVAL UNDER SECTION 505 OF THE ACT
ADMINISTERED BY THE CENTER FOR BIOLOGICS EVALUATION AND RESEARCH LIST**

DEXTRAN 75, 6% IN SODIUM CHLORIDE 0.9%

INJECTABLE; INJECTION

NONE

PHARMACHEM

6GM/100ML;0.9GM/100ML

N080564

Sep 19, 1952

6GM/100ML;0.9GM/100ML

N160759

Aug 19, 1970

HETASTARCH 6% IN 0.9% NACL

INJECTABLE; INJECTION

NONE

GENSIA LABS INC

6GM/100ML;0.9GM/100ML

N740592

ANDA

Nov 12, 1998

MCGAW INC

6GM/100ML;0.9GM/100ML

N740283

ANDA

Oct 21, 1998

HETASTARCH 6% IN LACTATED ELECTROLYTE INJECTION

INJECTABLE; INJECTION

HEXTEND

BIOTIME INC

6GM/100ML

N200952

Mar 31, 1999

HETASTARCH 6% IN SODIUM CHLORIDE 0.9%

INJECTABLE; INJECTION

6% HETASTARCH IN SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER

ABBOTT

6GM/100ML;0.9GM/100ML

N740193

ANDA

Jan 30, 1995

HESPAN

MCGAW INC

6GM/100ML;0.9GM/100ML

N160889

Jul 17, 1972

HESPAN IN PLASTIC CONTAINER

MCGAW INC

6GM/100ML;0.9GM/100ML

N890105

Apr 04, 1991

HETASTARCH 6% IN SODIUM CHLORIDE 0.9% NACL 500 ML GLASS BOTTLES

INJECTABLE; INJECTION

NONE

ABBOTT

6GM/100ML;0.9 GM/100ML

N720746

ANDA

Feb 07, 1996

LABORATORIES -

HOSPITAL PRODUCTS

DIV

PENTASTARCH 10% IN SODIUM CHLORIDE 0.9%

INJECTABLE; INJECTION

PENTASPAN

MCGAW INC

10GM/100ML;0.9GM/100ML

N841207

May 19, 1987

PENTASPAN IN PLASTIC CONTAINER

MCGAW INC

10GM/100ML;0.9GM/100ML

N890104

Apr 04, 1991

PERFLUORODECALIN; PERFLUOROTRI-N-PROPYLAMINE

INJECTABLE; INJECTION

FLUOSOL

ALPHA THERPTC

17.5GM/100ML;7.5GM/100ML

N860909

Dec 26, 1989

RED BLOOD CELL PROCESSING SOLUTION

INJECTABLE; INJECTION

REJUVESOL

CYTOSOL LABS INC

N950522

Feb 26, 1997

DISCONTINUED DRUG PRODUCT LIST

6 - 1 (of 274)

ABARELIX

INJECTABLE; INTRAMUSCULAR PLENAXIS PRAECIS	100MG/VIAL	N21320 001	Nov 25, 2003
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ACETAMINOPHEN

INJECTABLE; INJECTION INJECTAPAP ORTHO MCNEIL PHARM	100MG/ML	N17785 001	Mar 07, 1986
SUPPOSITORY; RECTAL ACETAMINOPHEN ABLE	120MG	N73106 001	Feb 27, 1995
	325MG	N73107 001	Feb 27, 1995
	650MG	N73108 001	Feb 27, 1995
ROXANE	120MG	N71010 001	May 12, 1987
	650MG	N71011 001	May 12, 1987
TYLENOL MCNEIL CONS	120MG	N17756 002	
	650MG	N17756 001	

ACETAMINOPHEN; ASPIRIN; CODEINE PHOSPHATE

CAPSULE; ORAL ACETAMINOPHEN, ASPIRIN, AND CODEINE PHOSPHATE MIKART	150MG;180MG;15MG 150MG;180MG;60MG	N81095 001 N81097 001	Oct 26, 1990 Oct 26, 1990
CODEINE, ASPIRIN, APAP FORMULA NO. 2 SCHERER LABS	150MG;180MG;15MG	N85640 001	
CODEINE, ASPIRIN, APAP FORMULA NO. 3 SCHERER LABS	150MG;180MG;30MG	N85639 001	
CODEINE, ASPIRIN, APAP FORMULA NO. 4 SCHERER LABS	150MG;180MG;60MG	N85638 001	

ACETAMINOPHEN; BUTALBITAL

CAPSULE; ORAL BANCAP FOREST PHARMS	325MG;50MG	N88889 001	Jan 16, 1986
TRIAPRIN DUNHALL	325MG;50MG	N89268 001	Jul 02, 1987
TABLET; ORAL BUTALBITAL AND ACETAMINOPHEN HALSEY	325MG;50MG	N89568 001	Oct 05, 1988
WATSON LABS	325MG;50MG	N87550 001	Oct 19, 1984
SEDAPAP MAYRAND	650MG;50MG	N88944 001	Oct 17, 1985

ACETAMINOPHEN; BUTALBITAL; CAFFEINE

CAPSULE; ORAL ACETAMINOPHEN, BUTALBITAL AND CAFFEINE GILBERT LABS	325MG;50MG;40MG	N88825 001	Dec 05, 1984
ANOQUAN SHIRE	325MG;50MG;40MG	N87628 001	Oct 01, 1986
BUTALBITAL, ACETAMINOPHEN AND CAFFEINE GRAHAM DM	325MG;50MG;40MG 325MG;50MG;40MG 325MG;50MG;40MG	N88743 001 N88765 001 N89067 001	Jul 18, 1985 Mar 27, 1985 Apr 19, 1985
MALLINCKRODT FEMCET	325MG;50MG;40MG	N88758 001	Mar 27, 1985
MALLINCKRODT	325MG;50MG;40MG	N89102 001	Jun 19, 1985
MEDIGESIC PLUS US CHEM	325MG;50MG;40MG	N89115 001	Jan 14, 1986
TRIAD MALLINCKRODT	325MG;50MG;40MG	N89023 001	Jun 19, 1985

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 2 (of 274)

ACETAMINOPHEN; BUTALBITAL; CAFFEINE

TABLET; ORAL

ACETAMINOPHEN, BUTALBITAL AND CAFFEINE

GILBERT LABS

325MG;50MG;40MG

N87629 001

Nov 13, 1984

BUTALBITAL, ACETAMINOPHEN AND CAFFEINE

ABLE

325MG;50MG;40MG

N40390 001

Jul 23, 2001

500MG;50MG;40MG

N40394 001

Jul 23, 2001

ESGIC

FOREST PHARMS

325MG;50MG;40MG

N89660 001

Dec 23, 1988

ACETAMINOPHEN; BUTALBITAL; CAFFEINE; CODEINE PHOSPHATE

CAPSULE; ORAL

BUTALBITAL, ACETAMINOPHEN, CAFFEINE AND CODEINE PHOSPHATE

ABLE

325MG;50MG;40MG;30MG

N76528 001

Aug 21, 2003

ACETAMINOPHEN; CAFFEINE; DIHYDROCODEINE BITARTRATE

CAPSULE; ORAL

DHC PLUS

PURDUE FREDERICK

356.4MG;30MG;16MG

N88584 001

Mar 04, 1986

SYNALGOS-DC-A

LEITNER PHARMS

356.4MG;30MG;16MG

N89166 001

May 14, 1986

ACETAMINOPHEN; CODEINE PHOSPHATE

CAPSULE; ORAL

ACETAMINOPHEN AND CODEINE PHOSPHATE

TEVA

300MG;30MG

N88324 001

Dec 29, 1983

ACETAMINOPHEN W/ CODEINE #2

TEVA

300MG;15MG

N88537 001

Jun 04, 1984

ACETAMINOPHEN W/ CODEINE #4

TEVA

300MG;60MG

N88599 001

Jun 01, 1984

PHENAPHEN W/ CODEINE NO. 2

ROBINS AH

325MG;15MG

N84444 001

PHENAPHEN W/ CODEINE NO. 3

ROBINS AH

325MG;30MG

N84445 001

PHENAPHEN W/ CODEINE NO. 4

ROBINS AH

325MG;60MG

N84446 001

PROVAL #3

SOLVAY

325MG;30MG

N85685 001

TYLENOL W/ CODEINE NO. 3

ORTHO MCNEIL PHARM

300MG;30MG

N87422 001

TYLENOL W/ CODEINE NO. 4

ORTHO MCNEIL PHARM

300MG;60MG

N87421 001

SOLUTION; ORAL

ACETAMINOPHEN AND CODEINE PHOSPHATE

CLONMEL

120MG/5ML;12MG/5ML

N40098 001

Sep 20, 1996

TYLENOL W/ CODEINE

ORTHO MCNEIL PHARM

120MG/5ML;12MG/5ML

N85057 001

TABLET; ORAL

ACETAMINOPHEN AND CODEINE PHOSPHATE

ABLE

300MG;30MG

N40452 001

Aug 01, 2002

300MG;60MG

N40459 001

Aug 01, 2002

DURAMED PHARMS BARR

300MG;15MG

N88353 001

Feb 06, 1984

300MG;30MG

N88354 001

Feb 06, 1984

300MG;60MG

N88355 001

Feb 06, 1984

HALSEY

300MG;60MG

N86549 001

KV PHARM

300MG;30MG

N85288 001

300MG;60MG

N85365 001

325MG;15MG

N85364 001

325MG;45MG **Federal Register
determination that product was not
discontinued or withdrawn for safety
or efficacy reasons**

N85363 001

MIKART

300MG;60MG

N89244 001

Feb 25, 1986

MUTUAL PHARM

300MG;15MG

N85795 001

300MG;30MG

N85794 001

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 3 (of 274)

ACETAMINOPHEN; CODEINE PHOSPHATE

TABLET; ORAL

ACETAMINOPHEN AND CODEINE PHOSPHATE

MUTUAL PHARM	300MG;60MG	N89673	001	Feb 10, 1988
PUREPAC PHARM	300MG;30MG	N86681	001	
	300MG;60MG	N86683	001	
ROXANE	500MG;15MG	N89511	001	Apr 25, 1989
	500MG;30MG	N89512	001	Apr 25, 1989
	500MG;60MG	N89513	001	Apr 25, 1989
SANDOZ	300MG;15MG	N87433	001	
	300MG;30MG	N85291	002	
	300MG;30MG	N85917	001	
	300MG;60MG	N85964	001	
	300MG;60MG	N87423	001	
WARNER CHILCOTT	300MG;15MG	N85992	001	
	300MG;30MG	N85218	002	

ACETAMINOPHEN AND CODEINE PHOSPHATE #2

SUPERPHARM	300MG;15MG	N89183	001	Oct 18, 1985
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ACETAMINOPHEN AND CODEINE PHOSPHATE #3

PUREPAC PHARM	300MG;30MG	N89080	001	Jul 17, 1986
SUPERPHARM	300MG;30MG	N89184	001	Oct 18, 1985
	300MG;30MG	N89253	001	May 19, 1986

ACETAMINOPHEN AND CODEINE PHOSPHATE #4

SUPERPHARM	300MG;60MG	N89185	001	Oct 18, 1985
	300MG;60MG	N89254	001	May 19, 1986

ACETAMINOPHEN AND CODEINE PHOSPHATE NO. 2

AM THERAP	300MG;15MG	N89478	001	Mar 03, 1987
	300MG;15MG	N89481	001	Mar 03, 1987

ACETAMINOPHEN AND CODEINE PHOSPHATE NO. 3

AM THERAP	300MG;30MG	N89479	001	Mar 03, 1987
	300MG;30MG	N89482	001	Mar 03, 1987

ACETAMINOPHEN AND CODEINE PHOSPHATE NO. 4

AM THERAP	300MG;60MG	N89480	001	Mar 03, 1987
	300MG;60MG	N89483	001	Mar 03, 1987
ROXANE	300MG;60MG	N84667	001	

ACETAMINOPHEN W/ CODEINE

LEDERLE	300MG;30MG	N87141	001	
MUTUAL PHARM	300MG;60MG	N87653	001	Apr 13, 1982

ACETAMINOPHEN W/ CODEINE NO. 2

ROXANE	300MG;15MG	N84659	001	
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ACETAMINOPHEN W/ CODEINE NO. 3

ROXANE	300MG;30MG	N84656	001	
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ACETAMINOPHEN W/ CODEINE PHOSPHATE

HALSEY	300MG;15MG	N83871	001	
	300MG;30MG	N83872	001	
USL PHARMA	300MG;30MG	N87919	001	Jun 22, 1982
	300MG;60MG	N87920	001	Jun 22, 1982
VITARINE	300MG;30MG	N85676	001	
WARNER CHILCOTT	300MG;60MG	N87306	001	
WATSON LABS	300MG;15MG	N87277	001	May 26, 1982
	300MG;30MG	N87276	001	May 26, 1982
	300MG;60MG	N87275	001	May 26, 1982
WHITEWORTH TOWN PLSN	300MG;30MG	N84360	001	
	300MG;60MG	N85607	001	

APAP W/ CODEINE PHOSPHATE

EVERYLIFE	325MG;30MG	N85217	001	
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CAPITAL WITH CODEINE

CARNRICK	325MG;30MG	N83643	001	
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CODEINE PHOSPHATE AND ACETAMINOPHEN

VALEANT PHARM INTL	300MG;30MG	N85896	001	
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EMPRACET W/ CODEINE PHOSPHATE #3

GLAXOSMITHKLINE	300MG;30MG	N83951	001	
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EMPRACET W/ CODEINE PHOSPHATE #4

GLAXOSMITHKLINE	300MG;60MG	N83951	002	
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27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 4 (of 274)

ACETAMINOPHEN; CODEINE PHOSPHATE

TABLET; ORAL

PAPA-DEINE #3

VANGARD

300MG;30MG

N88037 001

Mar 20, 1984

PAPA-DEINE #4

VANGARD

300MG;60MG

N88715 001

Mar 20, 1984

PHENAPHEN-650 W/ CODEINE

ROBINS AH

650MG;30MG

N85856 001

TYLENOL W/ CODEINE

ORTHO MCNEIL PHARM

325MG;7.5MG

N85056 001

325MG;15MG

N85056 002

325MG;30MG

N85056 003

325MG;60MG

N85056 004

TYLENOL W/ CODEINE NO. 1

ORTHO MCNEIL PHARM

300MG;7.5MG

N85055 001

TYLENOL W/ CODEINE NO. 2

ORTHO MCNEIL PHARM

300MG;15MG

N85055 002

TYLENOL W/ CODEINE NO. 4

ORTHO MCNEIL PHARM

300MG;60MG

N85055 004

ACETAMINOPHEN; HYDROCODONE BITARTRATE

CAPSULE; ORAL

ACETAMINOPHEN AND HYDROCODONE BITARTRATE

CENT PHARMS

500MG;5MG

N88898 001

Mar 27, 1985

BANCAP HC

FOREST PHARMS

500MG;5MG

N87961 001

Mar 17, 1983

CO-GESIC

CENT PHARMS

500MG;5MG

N89360 001

Mar 02, 1988

TABLET; ORAL

DURADYNE DHC

FOREST PHARMS

500MG;5MG

N87809 001

Mar 17, 1983

HYDROCODONE BITARTRATE AND ACETAMINOPHEN

ABLE

325MG;5MG

N40478 001

Nov 08, 2002

325MG;7.5MG

N40464 001

Oct 23, 2002

325MG;10MG

N40464 002

Oct 23, 2002

500MG;5MG

N40477 001

Nov 06, 2002

500MG;7.5MG

N40490 001

May 21, 2003

500MG;10MG

N40473 001

Nov 06, 2002

650MG;7.5MG

N40474 001

Jan 02, 2003

650MG;10MG

N40476 001

Oct 23, 2002

750MG;7.5MG

N40469 001

Oct 25, 2002

BARR

500MG;5MG

N88577 001

Dec 21, 1984

ENDO PHARMS

500MG;5MG

N40281 001

Sep 30, 1998

500MG;7.5MG

N40280 001

Sep 30, 1998

650MG;7.5MG

N40280 002

Sep 30, 1998

650MG;10MG

N40280 003

Sep 30, 1998

750MG;7.5MG

N40281 002

Sep 30, 1998

HALSEY

500MG;5MG

N89554 001

Jun 12, 1987

USL PHARMA

500MG;5MG

N89290 001

May 29, 1987

500MG;5MG

N89291 001

May 29, 1987

WATSON LABS

325MG;7.5MG

N40248 001

Apr 28, 2000

HY-PHEN

ASCHER

500MG;5MG

N87677 001

May 03, 1982

NORCET

ABANA

500MG;5MG

N88871 001

May 15, 1986

TYCOLET

ORTHO MCNEIL PHARM

500MG;5MG

N89385 001

Aug 27, 1986

VICODIN

ABBOTT

500MG;5MG

N85667 001

ACETAMINOPHEN; OXYCODONE HYDROCHLORIDE

CAPSULE; ORAL

OXYCODONE AND ACETAMINOPHEN

HALSEY

500MG;5MG

N89994 001

May 04, 1989

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 5 (of 274)

ACETAMINOPHEN; OXYCODONE HYDROCHLORIDE

CAPSULE; ORAL

TYLOX-325

ORTHO MCNEIL PHARM 325MG;5MG

N88246 001 Nov 08, 1984

TABLET; ORAL

OXYCODONE 2.5/APAP 500

BRISTOL MYERS SQUIBB 500MG;2.5MG

N85910 001

OXYCODONE 5/APAP 500

BRISTOL MYERS SQUIBB 500MG;5MG

N85911 001

ACETAMINOPHEN; OXYCODONE HYDROCHLORIDE; OXYCODONE TEREPHTHALATE

CAPSULE; ORAL

TYLOX

ORTHO MCNEIL PHARM 500MG;4.5MG;0.38MG

N85375 001

ACETAMINOPHEN; PROPOXYPHENE HYDROCHLORIDE

TABLET; ORAL

DARVOCET

AAIPHARMA LLC 325MG;32.5MG

N16844 001

DOLENE AP-65

LEDERLE 650MG;65MG

N85100 001

PROPOXYPHENE HYDROCHLORIDE AND ACETAMINOPHEN

MYLAN 325MG;32MG

N83689 001

ACETAMINOPHEN; PROPOXYPHENE NAPSYLATE

TABLET; ORAL

PROPACET 100

TEVA 650MG;100MG

N70107 001 Jun 12, 1985

PROPOXYPHENE NAPSYLATE AND ACETAMINOPHEN

ABLE 650MG;100MG

N75838 001 Jul 11, 2001

HALSEY 325MG;50MG

N72105 001 May 13, 1988

650MG;100MG

N72106 001 May 13, 1988

MUTUAL PHARM 325MG;50MG

N70115 001 Jun 12, 1985

650MG;100MG

N70116 001 Jun 12, 1985

SUPERPHARM 650MG;100MG

N71319 001 Jan 06, 1987

TEVA 650MG;100MG

N70732 001 Jan 03, 1986

WATSON LABS 325MG;50MG

N70398 001 Dec 18, 1986

650MG;100MG

N70399 001 Dec 18, 1986

ACETAZOLAMIDE

CAPSULE, EXTENDED RELEASE; ORAL

DIAMOX

DURAMED PHARMS BARR 500MG

N12945 001

TABLET; ORAL

ACETAZOLAMIDE

ALRA 250MG

N83320 001

ASCOT 250MG

N87686 001 Oct 20, 1982

VANGARD 250MG

N87654 001 Feb 05, 1982

WATSON LABS 250MG

N84498 002

DIAMOX

DURAMED PHARMS BARR 125MG

N08943 001

250MG

N08943 002

ACETIC ACID, GLACIAL

SOLUTION/DROPS; OTIC

ACETIC ACID

KV PHARM 2%

N85493 001

ORLEX

PROCTER AND GAMBLE 2%

N86845 001

VOSOL

MEDPOINTE PHARM HLC 2%

N12179 001

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 6 (of 274)

ACETIC ACID, GLACIAL; ALUMINUM ACETATE

SOLUTION/DROPS; OTIC

BOROFAIR

PHARMAFAIR

2%;0.79%

N88606 001

Aug 21, 1985

DOMEBORO

BAYER PHARMS

2%;0.79%

N84476 001

ACETIC ACID, GLACIAL; DESONIDE

SOLUTION/DROPS; OTIC

TRIDESILON

BAYER PHARMS

2%;0.05%

N17914 001

ACETIC ACID, GLACIAL; HYDROCORTISONE

SOLUTION/DROPS; OTIC

ACETIC ACID W/ HYDROCORTISONE

KV PHARM

2%;1%

N85492 001

HYDROCORTISONE AND ACETIC ACID

BAUSCH AND LOMB

2%;1%

N40097 001

Oct 31, 1994

ORLEX HC

PROCTER AND GAMBLE

2%;1%

N86844 001

ACETIC ACID, GLACIAL; HYDROCORTISONE; NEOMYCIN SULFATE

SUSPENSION/DROPS; OTIC

NEO-CORT-DOME

BAYER PHARMS

2%;1%;EQ 0.35% BASE

N50238 001

ACETOHEXAMIDE

TABLET; ORAL

ACETOHEXAMIDE

USL PHARMA

250MG

N70753 001

Nov 03, 1986

500MG

N70754 001

Nov 03, 1986

DYMELOR

LILLY

250MG

N13378 002

500MG

N13378 001

ACETOPHENAZINE MALEATE

TABLET; ORAL

TINDAL

SCHERING

20MG

N12254 002

ACETRIZOATE SODIUM

SOLUTION; INTRAUTERINE

SALPIX

ORTHO MCNEIL PHARM

53%

N09008 001

ACETYLCHOLINE CHLORIDE

FOR SOLUTION; OPHTHALMIC

MIOCHOL

NOVARTIS

20MG/VIAL

N16211 001

ACETYLCYSTEINE

SOLUTION; INHALATION, ORAL

ACETYLCYSTEINE

ROXANE

10%

N72621 001

Sep 30, 1992

20%

N72622 001

Sep 30, 1992

MUCOSIL-10

DEY

10%

N70575 001

Oct 14, 1986

MUCOSIL-20

DEY

20%

N70576 001

Oct 14, 1986

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 7 (of 274)

ACETYLCYSTEINE; ISOPROTERENOL HYDROCHLORIDE

SOLUTION; INHALATION				
MUCOMYST W/ ISOPROTERENOL				
MEAD JOHNSON	10%;0.05%	N17366	001	

ACETYLDIGITOXIN

TABLET; ORAL				
ACYLANID				
NOVARTIS	0.1MG	N09436	001	

ACRISORCIN

CREAM; TOPICAL				
AKRINOL				
SCHERING	2MG/GM	N12470	001	

ACYCLOVIR

CAPSULE; ORAL				
ACYCLOVIR				
LEK PHARM	200MG	N74750	001	Apr 22, 1997
ROXANE	200MG	N74570	002	Apr 22, 1997
TABLET; ORAL				
ACYCLOVIR				
CLONMEL HLTHCARE	400MG	N74834	001	Apr 24, 1997
	800MG	N74834	002	Apr 24, 1997
LEK PHARM	400MG	N74658	001	Apr 22, 1997
	800MG	N74658	002	Apr 22, 1997
TEVA	200MG **Federal Register determination that product was not discontinued or withdrawn for safety or efficacy reasons**	N74556	001	Apr 22, 1997

ACYCLOVIR SODIUM

INJECTABLE; INJECTION				
ACYCLOVIR				
ABBOTT	EQ 50MG BASE/ML	N75114	001	Jul 26, 1999
ACYCLOVIR SODIUM				
APOTHECON	EQ 500MG BASE/VIAL	N74897	001	Feb 27, 1998
	EQ 1GM BASE/VIAL	N74897	002	Feb 27, 1998
HOSPIRA	EQ 500MG BASE/VIAL	N74663	001	Apr 22, 1997
	EQ 1GM BASE/VIAL	N74663	002	Apr 22, 1997
ZOVIRAX				
GLAXOSMITHKLINE	EQ 250MG BASE/VIAL	N18603	003	Aug 30, 1983

ADAPALENE

SOLUTION; TOPICAL				
DIFFERIN				
GALDERMA LABS LP	0.1%	N20338	001	May 31, 1996

ALATROFLOXACIN MESYLATE

INJECTABLE; INJECTION				
TROVAN PRESERVATIVE FREE				
PFIZER	EQ 200MG BASE/VIAL	N20760	001	Dec 18, 1997
	EQ 300MG BASE/VIAL	N20760	002	Dec 18, 1997

ALBUMIN CHROMATED CR-51 SERUM

INJECTABLE; INJECTION				
CHROMALBIN				
ISO TEX	100uCi/VIAL	N17835	001	
	250uCi/VIAL	N17835	002	
	500uCi/VIAL	N17835	003	

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 8 (of 274)

ALBUMIN IODINATED I-125 SERUM

INJECTABLE; INJECTION		
RADIO-IODINATED (I 125) SERUM ALBUMIN (HUMAN)		
BAYER PHARMS	2.5uCi/AMP	N17846 001
RADIOIODINATED SERUM ALBUMIN (HUMAN) IHSA I 125		
MALLINCKRODT	6.67uCi/ML	N17844 003
	10uCi/ML	N17844 001
	100uCi/ML	N17844 002

ALBUMIN IODINATED I-131 SERUM

INJECTABLE; INJECTION		
MEGATOPE		
ISO TEX	2mCi/VIAL	N17837 003
	5uCi/AMP	N17837 004
	20uCi/AMP	N17837 005

ALBUTEROL

AEROSOL, METERED; INHALATION			
ALBUTEROL			
GENPHARM	0.09MG/INH	N73045 001	Aug 19, 1997
PLIVA	0.09MG/INH	N74072 001	Aug 01, 1996

ALBUTEROL SULFATE

CAPSULE; INHALATION			
VENTOLIN ROTACAPS			
GLAXOSMITHKLINE	EQ 0.2MG BASE	N19489 001	May 04, 1988
SOLUTION; INHALATION			
ALBUTEROL SULFATE			
COPLEY PHARM	EQ 0.083% BASE	N73495 001	May 28, 1993
	EQ 0.5% BASE	N73307 001	Nov 27, 1991
PROVENTIL			
SCHERING	EQ 0.083% BASE	N19243 002	Jan 14, 1987
	EQ 0.5% BASE	N19243 001	Jan 14, 1987
VENTOLIN			
GLAXOSMITHKLINE	EQ 0.083% BASE	N19773 001	Apr 23, 1992
	EQ 0.5% BASE	N19269 002	Jan 16, 1987
SYRUP; ORAL			
ALBUTEROL SULFATE			
MOVA	EQ 2MG BASE/5ML	N74302 001	Sep 30, 1994
WATSON LABS	EQ 2MG BASE/5ML	N73165 001	Apr 29, 1993
PROVENTIL			
SCHERING	EQ 2MG BASE/5ML	N18062 001	Jan 19, 1983
VENTOLIN			
GLAXOSMITHKLINE	EQ 2MG BASE/5ML	N19621 001	Jun 10, 1987
TABLET; ORAL			
ALBUTEROL SULFATE			
AM THERAP	EQ 2MG BASE	N72449 001	Dec 05, 1989
	EQ 4MG BASE	N72450 001	Dec 05, 1989
CLONMEL HLTHCARE	EQ 2MG BASE	N72859 001	Dec 20, 1989
	EQ 4MG BASE	N72860 001	Dec 20, 1989
COPLEY PHARM	EQ 2MG BASE	N72966 001	Nov 22, 1991
	EQ 4MG BASE	N72967 001	Nov 22, 1991
UCB INC	EQ 2MG BASE	N73120 001	Sep 29, 1992
	EQ 4MG BASE	N73121 001	Sep 29, 1992
WARNER CHILCOTT	EQ 2MG BASE	N72817 001	Jan 09, 1990
	EQ 4MG BASE	N72818 001	Jan 09, 1990
PROVENTIL			
SCHERING	EQ 2MG BASE	N17853 001	May 07, 1982
	EQ 4MG BASE	N17853 002	May 07, 1982
VENTOLIN			
GLAXOSMITHKLINE	EQ 2MG BASE	N19112 001	Jul 10, 1986
	EQ 4MG BASE	N19112 002	Jul 10, 1986

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 9 (of 274)

ALBUTEROL SULFATE

TABLET, EXTENDED RELEASE; ORAL

PROVENTIL

SCHERING

EQ 4MG BASE

N19383 001

Jul 13, 1987

VOLMAX

MURO

EQ 4MG BASE

N19604 002

Dec 23, 1992

EQ 8MG BASE

N19604 001

Dec 23, 1992

ALCOHOL

INJECTABLE; INJECTION

ALCOHOL 5% IN DEXTROSE 5%

MILES

5ML/100ML

N83483 001

ALCOHOL; DEXTROSE

INJECTABLE; INJECTION

ALCOHOL 5% IN DEXTROSE 5% IN WATER

BAXTER HLTHCARE

5ML/100ML;5GM/100ML

N83256 001

ALGLUCERASE

INJECTABLE; INJECTION

CEREDASE

GENZYME

10 UNITS/ML

N20057 004

May 08, 1992

ALKAVERVIR

TABLET; ORAL

VERILOID

3M

2MG

N07336 002

3MG

N07336 003

ALLOPURINOL

TABLET; ORAL

ALLOPURINOL

MUTUAL PHARM

100MG

N70466 001

Dec 24, 1985

300MG

N70467 001

Dec 24, 1985

PUREPAC PHARM

100MG

N70579 001

Apr 14, 1986

300MG

N70580 001

Apr 14, 1986

SUPERPHARM

100MG

N70950 001

Nov 30, 1988

300MG

N70951 001

Nov 30, 1988

WATSON LABS

100MG

N18241 001

Nov 16, 1984

100MG

N18785 001

Sep 28, 1984

300MG

N18241 002

Nov 16, 1984

300MG

N18785 002

Sep 28, 1984

LOPURIN

ABBOTT

100MG

N18297 001

300MG

N18297 002

ALPHA-TOCOPHEROL; ASCORBIC ACID; BIOTIN; CHOLECALCIFEROL; CYANOCOBALAMIN; FOLIC ACID; NIACINAMIDE; PANTOTHENIC ACID; PYRIDOXINE; RIBOFLAVIN; THIAMINE; VITAMIN A

INJECTABLE; INJECTION

CERNEVIT-12

BAXTER HLTHCARE

11.2 IU/VIAL;125MG/VIAL;60UGM/VIAL;200
IU/VIAL;5.5MG/VIAL;414UGM/VIAL;46MG/VIA
L;17.25MG/VIAL;4.53MG/VIAL;4.14MG/VIAL;
3.51MG/VIAL;3,500 IU/VIAL

N20924 001

Apr 06, 1999

ALPRAZOLAM

SOLUTION; ORAL

ALPRAZOLAM

ROXANE

0.5MG/5ML

N74314 001

Oct 31, 1993

TABLET; ORAL

ALPRAZOLAM

CLONMEL HLTHCARE

0.25MG

N74174 001

Oct 19, 1993

0.5MG

N74174 002

Oct 19, 1993

1MG

N74174 003

Oct 19, 1993

DISCONTINUED DRUG PRODUCT LIST

6 - 10 (of 274)

ALPRAZOLAM

TABLET; ORAL
ALPRAZOLAM

CLONMEL HLTHCARE	2MG	N74174	004	Oct 19, 1993
ROXANE	0.25MG	N74199	001	Oct 19, 1993
	0.5MG	N74199	002	Oct 19, 1993
	1MG	N74199	003	Oct 19, 1993

ALPROSTADIL

INJECTABLE; INJECTION
CAVERJECT

PFIZER	0.005MG/ML	N20755	001	Oct 31, 1997
	0.01MG/ML	N20755	002	Oct 01, 1997
	0.02MG/ML	N20755	003	Oct 01, 1997

EDEX				
SCHWARZ PHARMA	0.005MG/VIAL	N20649	001	Jun 12, 1997

ALSEROXYLON

TABLET; ORAL
RAUTENSIN

NOVARTIS	2MG	N09215	001	
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RAUWILOID				
3M	2MG	N08867	001	

ALUMINUM HYDROXIDE; MAGNESIUM TRISILICATE

TABLET, CHEWABLE; ORAL
ALUMINUM HYDROXIDE AND MAGNESIUM TRISILICATE

PENNEX	80MG;20MG	N89449	001	Nov 27, 1987
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FOAMICON				
NOVARTIS	80MG;20MG	N72687	001	Jun 28, 1989

AMANTADINE HYDROCHLORIDE

CAPSULE; ORAL
AMANTADINE HYDROCHLORIDE

WATSON LABS	100MG	N71382	001	Jan 21, 1987
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SYMADINE				
SOLVAY	100MG	N71000	001	Sep 04, 1986

SYMMETREL				
ENDO PHARMS	100MG	N16020	001	

SYRUP; ORAL				
SYMMETREL				
ENDO PHARMS	50MG/5ML	N16023	002	

AMCINONIDE

CREAM; TOPICAL
CYCLOCORT

ASTELLAS	0.025%	N18116	001	
	0.1%	N18116	002	

LOTION; TOPICAL				
CYCLOCORT				
ASTELLAS	0.1%	N19729	001	Jun 13, 1988

OINTMENT; TOPICAL				
CYCLOCORT				
ASTELLAS	0.1%	N18498	001	

AMDINOCILLIN

INJECTABLE; INJECTION
COACTIN

ROCHE	250MG/VIAL	N50565	001	Dec 21, 1984
	500MG/VIAL	N50565	002	Dec 21, 1984
	1GM/VIAL	N50565	003	Dec 21, 1984

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 11 (of 274)

AMIFOSTINEINJECTABLE; INJECTION
ETHYOL

MEDIMMUNE ONCOLOGY 375MG/VIAL N20221 002 Sep 10, 1999

AMIKACIN SULFATEINJECTABLE; INJECTION
AMIKACIN SULFATE

ABBOTT	EQ 250MG BASE/ML	N63265	001	Nov 30, 1994
	EQ 250MG BASE/ML	N63266	001	Oct 31, 1994
ASTRAZENECA	EQ 50MG BASE/ML	N63167	001	Dec 14, 1995
	EQ 250MG BASE/ML	N63169	001	Dec 14, 1995
BAXTER HLTHCARE	EQ 50MG BASE/ML	N63274	001	May 18, 1992
	EQ 250MG BASE/ML	N63275	001	May 18, 1992
HOSPIRA	EQ 250MG BASE/ML	N64099	001	Jun 20, 1995
MAYNE PHARMA USA	EQ 50MG BASE/ML	N63350	001	Jul 30, 1993
	EQ 250MG BASE/ML	N63350	002	Jul 30, 1993

AMIKIN

APOTHECON	EQ 50MG BASE/ML	N50495	001	
	EQ 50MG BASE/ML	N62562	001	Sep 20, 1984
	EQ 250MG BASE/ML	N50495	002	
	EQ 250MG BASE/ML	N62562	002	Sep 20, 1984
AMIKIN IN SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER				
APOTHECON	EQ 5MG BASE/ML	N50618	002	Nov 30, 1987
	EQ 10MG BASE/ML	N50618	001	Nov 30, 1987

AMILORIDE HYDROCHLORIDETABLET; ORAL
MIDAMOR

MERCK 5MG N18200 001

AMILORIDE HYDROCHLORIDE; HYDROCHLOROTHIAZIDETABLET; ORAL
MODURETIC 5-50
MERCK

EQ 5MG ANHYDROUS;50MG N18201 001

AMINO ACIDS

INJECTABLE; INJECTION

AMINESS 5.2% ESSENTIAL AMINO ACIDS W/ HISTADINE

HOSPIRA 5.2% (5.2GM/100ML) N18901 001 Apr 06, 1984

AMINOSYN 3.5% IN PLASTIC CONTAINER

ABBOTT 3.5% (3.5GM/100ML) N18804 001 May 15, 1984

3.5% (3.5GM/100ML) N18875 001 Aug 08, 1984

AMINOSYN II 3.5%

HOSPIRA 3.5% (3.5GM/100ML) N19438 001 Apr 03, 1986

AMINOSYN II 3.5% IN PLASTIC CONTAINER

ABBOTT 3.5% (3.5GM/100ML) N19491 001 Oct 10, 1986

AMINOSYN II 5%

HOSPIRA 5% (5GM/100ML) N19438 002 Apr 03, 1986

AMINOSYN-HBC 7% IN PLASTIC CONTAINER

ABBOTT 7% (7GM/100ML) N19400 001 Jul 23, 1986

BRANCHAMIN 4%

BAXTER HLTHCARE 4% (4GM/100ML) N18678 001 Sep 28, 1984

FREAMINE 8.5%

B BRAUN 8.5% (8.5GM/100ML) N16822 001

FREAMINE II 8.5%

B BRAUN 8.5% (8.5GM/100ML) N16822 002

NEOPHAM 6.4%

HOSPIRA 6.4% (6.4GM/100ML) N18792 001 Jan 17, 1984

NOVAMINE 15% SULFITE FREE IN PLASTIC CONTAINER

BAXTER HLTHCARE 15% (15GM/100ML) N20107 001 Feb 05, 1993

NOVAMINE 8.5%

HOSPIRA 8.5% (8.5GM/100ML) N17957 002 Aug 09, 1982

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 12 (of 274)

AMINO ACIDS; CALCIUM CHLORIDE; DEXTROSE; MAGNESIUM CHLORIDE; POTASSIUM CHLORIDE;
POTASSIUM PHOSPHATE, DIBASIC; SODIUM CHLORIDEINJECTABLE; INJECTION

AMINOSYN II 3.5% W/ ELECTROLYTES IN DEXTROSE 25% W/ CALCIUM IN PLASTIC CONTAINER			
ABBOTT	3.5%;36.8MG/100ML;25GM/100ML;51MG/100ML ;22.4MG/100ML;261MG/100ML;205MG/100ML	N19714 001	Sep 12, 1988
AMINOSYN II 4.25% W/ ELECTROLYTES IN DEXTROSE 20% W/ CALCIUM IN PLASTIC CONTAINER			
ABBOTT	4.25%;36.8MG/100ML;20GM/100ML;51MG/100M L;22.4MG/100ML;261MG/100ML;205MG/100ML	N19714 002	Sep 12, 1988
AMINOSYN II 4.25% W/ ELECTROLYTES IN DEXTROSE 25% W/ CALCIUM IN PLASTIC CONTAINER			
ABBOTT	4.25%;36.8MG/100ML;25GM/100ML;51MG/100M L;22.4MG/100ML;261MG/100ML;205MG/100ML	N19714 004	Sep 12, 1988
AMINOSYN II 5% W/ ELECTROLYTES IN DEXTROSE 25% W/ CALCIUM IN PLASTIC CONTAINER			
ABBOTT	5%;36.8MG/100ML;25GM/100ML;51MG/100ML;2 2.4MG/100ML;261MG/100ML;205MG/100ML	N19714 003	Sep 12, 1988
HOSPIRA	5%;36.8MG/100ML;25GM/100ML;51MG/100ML;2 2.4MG/100ML;261MG/100ML;205MG/100ML	N19683 004	Nov 07, 1988

AMINO ACIDS; DEXTROSEINJECTABLE; INJECTION

AMINOSYN 3.5% W/ DEXTROSE 25% IN PLASTIC CONTAINER			
ABBOTT	3.5%;25GM/100ML	N19118 001	Oct 11, 1984
AMINOSYN 3.5% W/ DEXTROSE 5% IN PLASTIC CONTAINER			
ABBOTT	3.5%;5GM/100ML	N19120 001	Oct 11, 1984
AMINOSYN 4.25% W/ DEXTROSE 25% IN PLASTIC CONTAINER			
ABBOTT	4.25%;25GM/100ML	N19119 001	Oct 11, 1984
AMINOSYN II 3.5% IN DEXTROSE 25% IN PLASTIC CONTAINER			
ABBOTT	3.5%;25GM/100ML	N19505 002	Nov 07, 1986
	3.5%;25GM/100ML	N19713 006	Sep 09, 1988
AMINOSYN II 3.5% IN DEXTROSE 5% IN PLASTIC CONTAINER			
ABBOTT	3.5%;5GM/100ML	N19506 001	Nov 07, 1986
	3.5%;5GM/100ML	N19713 002	Sep 09, 1988
AMINOSYN II 4.25% IN DEXTROSE 10% IN PLASTIC CONTAINER			
ABBOTT	4.25%;10GM/100ML	N19713 001	Sep 09, 1988
AMINOSYN II 4.25% IN DEXTROSE 20% IN PLASTIC CONTAINER			
ABBOTT	4.25%;20GM/100ML	N19713 004	Sep 09, 1988
AMINOSYN II 4.25% IN DEXTROSE 25% IN PLASTIC CONTAINER			
ABBOTT	4.25%;25GM/100ML	N19504 002	Nov 07, 1986
	4.25%;25GM/100ML	N19713 005	Sep 09, 1988
AMINOSYN II 5% IN DEXTROSE 25% IN PLASTIC CONTAINER			
ABBOTT	5%;25GM/100ML	N19565 001	Dec 17, 1986
	5%;25GM/100ML	N19713 003	Sep 09, 1988
TRAVASOL 2.75% IN DEXTROSE 10% IN PLASTIC CONTAINER			
BAXTER HLTHCARE	2.75%;10GM/100ML	N19520 002	Sep 23, 1988
TRAVASOL 2.75% IN DEXTROSE 15% IN PLASTIC CONTAINER			
BAXTER HLTHCARE	2.75%;15GM/100ML	N19520 003	Sep 23, 1988
TRAVASOL 2.75% IN DEXTROSE 20% IN PLASTIC CONTAINER			
BAXTER HLTHCARE	2.75%;20GM/100ML	N19520 004	Sep 23, 1988
TRAVASOL 2.75% IN DEXTROSE 25% IN PLASTIC CONTAINER			
BAXTER HLTHCARE	2.75%;25GM/100ML	N19520 005	Sep 23, 1988
TRAVASOL 2.75% IN DEXTROSE 5% IN PLASTIC CONTAINER			
BAXTER HLTHCARE	2.75%;5GM/100ML	N19520 001	Sep 23, 1988
TRAVASOL 4.25% IN DEXTROSE 10% IN PLASTIC CONTAINER			
BAXTER HLTHCARE	4.25%;10GM/100ML	N19520 007	Sep 23, 1988
TRAVASOL 4.25% IN DEXTROSE 15% IN PLASTIC CONTAINER			
BAXTER HLTHCARE	4.25%;15GM/100ML	N19520 008	Sep 23, 1988
TRAVASOL 4.25% IN DEXTROSE 20% IN PLASTIC CONTAINER			
BAXTER HLTHCARE	4.25%;20GM/100ML	N19520 009	Sep 23, 1988
TRAVASOL 4.25% IN DEXTROSE 25% IN PLASTIC CONTAINER			
BAXTER HLTHCARE	4.25%;25GM/100ML	N19520 010	Sep 23, 1988
TRAVASOL 4.25% IN DEXTROSE 5% IN PLASTIC CONTAINER			
BAXTER HLTHCARE	4.25%;5GM/100ML	N19520 006	Sep 23, 1988

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 13 (of 274)

AMINO ACIDS; DEXTROSE; MAGNESIUM CHLORIDE; POTASSIUM ACETATE; POTASSIUM CHLORIDE; POTASSIUM PHOSPHATE, DIBASIC; SODIUM CHLORIDEINJECTABLE; INJECTION

AMINOSYN II 4.25% W/ ELECT AND ADJUSTED PHOSPHATE IN DEXTROSE 10% IN PLASTIC CONTAINER			
ABBOTT	4.25%;10GM/100ML;51MG/100ML;176.5MG/100ML;22.4MG/100ML;104.5MG/100ML;205MG/100ML	N19712 002	Sep 08, 1988
HOSPIRA			
	4.25%;10GM/100ML;51MG/100ML;176.5MG/100ML;22.4MG/100ML;104.5MG/100ML;205MG/100ML	N19682 003	Nov 01, 1988

AMINO ACIDS; DEXTROSE; MAGNESIUM CHLORIDE; POTASSIUM CHLORIDE; POTASSIUM PHOSPHATE, DIBASIC; SODIUM CHLORIDEINJECTABLE; INJECTION

AMINOSYN II 3.5% W/ ELECTROLYTES IN DEXTROSE 25% IN PLASTIC CONTAINER			
ABBOTT	3.5%;25GM/100ML;51MG/100ML;22.4MG/100ML;261MG/100ML;205MG/100ML	N19564 002	Dec 16, 1986
AMINOSYN II 4.25% W/ ELECTROLYTES IN DEXTROSE 25% IN PLASTIC CONTAINER			
ABBOTT	4.25%;25GM/100ML;51MG/100ML;22.4MG/100ML;261MG/100ML;205MG/100ML	N19564 004	Dec 16, 1986

AMINO ACIDS; DEXTROSE; MAGNESIUM CHLORIDE; POTASSIUM CHLORIDE; SODIUM CHLORIDE; SODIUM PHOSPHATE, DIBASIC, HEPTAHYDRATEINJECTABLE; INJECTION

AMINOSYN II 3.5% M IN DEXTROSE 5% IN PLASTIC CONTAINER			
ABBOTT	3.5%;5GM/100ML;30MG/100ML;97MG/100ML;120MG/100ML;49.3MG/100ML	N19564 001	Dec 16, 1986
	3.5%;5GM/100ML;30MG/100ML;97MG/100ML;120MG/100ML;49.3MG/100ML	N19712 001	Sep 08, 1988
AMINOSYN II 4.25% M IN DEXTROSE 10% IN PLASTIC CONTAINER			
ABBOTT	4.25%;10GM/100ML;30MG/100ML;97MG/100ML;120MG/100ML;49.3MG/100ML	N19564 003	Dec 16, 1986

AMINO ACIDS; DEXTROSE; MAGNESIUM CHLORIDE; POTASSIUM PHOSPHATE, DIBASIC; SODIUM ACETATE; SODIUM CHLORIDEINJECTABLE; INJECTION

TRAVASOL 2.75% SULFITE FREE W/ ELECTROLYTES IN DEXTROSE 10% IN PLASTIC CONTAINER			
BAXTER HLTHCARE	2.75%;10GM/100ML;51MG/100ML;261MG/100ML;216MG/100ML;112MG/100ML	N20147 002	Oct 23, 1995
TRAVASOL 2.75% SULFITE FREE W/ ELECTROLYTES IN DEXTROSE 15% IN PLASTIC CONTAINER			
BAXTER HLTHCARE	2.75%;15GM/100ML;51MG/100ML;261MG/100ML;216MG/100ML;112MG/100ML	N20147 003	Oct 23, 1995
TRAVASOL 2.75% SULFITE FREE W/ ELECTROLYTES IN DEXTROSE 20% IN PLASTIC CONTAINER			
BAXTER HLTHCARE	2.75%;20GM/100ML;51MG/100ML;261MG/100ML;216MG/100ML;112MG/100ML	N20147 004	Oct 23, 1995
TRAVASOL 2.75% SULFITE FREE W/ ELECTROLYTES IN DEXTROSE 25% IN PLASTIC CONTAINER			
BAXTER HLTHCARE	2.75%;25GM/100ML;51MG/100ML;261MG/100ML;216MG/100ML;112MG/100ML	N20147 005	Oct 23, 1995
TRAVASOL 2.75% SULFITE FREE W/ ELECTROLYTES IN DEXTROSE 5% IN PLASTIC CONTAINER			
BAXTER HLTHCARE	2.75%;5GM/100ML;51MG/100ML;261MG/100ML;216MG/100ML;112MG/100ML	N20147 001	Oct 23, 1995
TRAVASOL 4.25% SULFITE FREE W/ ELECTROLYTES IN DEXTROSE 10% IN PLASTIC CONTAINER			
BAXTER HLTHCARE	4.25%;10GM/100ML;51MG/100ML;261MG/100ML;297MG/100ML;77MG/100ML	N20147 007	Oct 23, 1995
TRAVASOL 4.25% SULFITE FREE W/ ELECTROLYTES IN DEXTROSE 15% IN PLASTIC CONTAINER			
BAXTER HLTHCARE	4.25%;15GM/100ML;51MG/100ML;261MG/100ML;297MG/100ML;77MG/100ML	N20147 008	Oct 23, 1995
TRAVASOL 4.25% SULFITE FREE W/ ELECTROLYTES IN DEXTROSE 20% IN PLASTIC CONTAINER			
BAXTER HLTHCARE	4.25%;20GM/100ML;51MG/100ML;261MG/100ML;297MG/100ML;77MG/100ML	N20147 009	Oct 23, 1995
TRAVASOL 4.25% SULFITE FREE W/ ELECTROLYTES IN DEXTROSE 25% IN PLASTIC CONTAINER			
BAXTER HLTHCARE	4.25%;25GM/100ML;51MG/100ML;261MG/100ML;297MG/100ML;77MG/100ML	N20147 010	Oct 23, 1995
TRAVASOL 4.25% SULFITE FREE W/ ELECTROLYTES IN DEXTROSE 5% IN PLASTIC CONTAINER			
BAXTER HLTHCARE	4.25%;5GM/100ML;51MG/100ML;261MG/100ML;297MG/100ML;77MG/100ML	N20147 006	Oct 23, 1995

DISCONTINUED DRUG PRODUCT LIST

6 - 14 (of 274)

AMINO ACIDS; MAGNESIUM ACETATE; PHOSPHORIC ACID; POTASSIUM ACETATE; SODIUM CHLORIDE

INJECTABLE; INJECTION

AMINOSYN 3.5% M IN PLASTIC CONTAINER

ABBOTT	3.5%;21MG/100ML;40MG/100ML;128MG/100ML; 234MG/100ML	N18804 002	May 15, 1984
	3.5%;21MG/100ML;40MG/100ML;128MG/100ML; 234MG/100ML	N18875 002	Aug 08, 1984

AMINO ACIDS; MAGNESIUM ACETATE; POTASSIUM ACETATE; SODIUM CHLORIDE

INJECTABLE; INJECTION

AMINOSYN 3.5% M

HOSPIRA	3.5%;21MG/100ML;128MG/100ML;234MG/100ML	N17789 005	
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AMINO ACIDS; MAGNESIUM ACETATE; POTASSIUM ACETATE; SODIUM CHLORIDE; SODIUM PHOSPHATE, DIBASIC, HEPTAHYDRATE

INJECTABLE; INJECTION

AMINOSYN II 3.5% M IN PLASTIC CONTAINER

ABBOTT	3.5%;32MG/100ML;128MG/100ML;222MG/100ML ;49MG/100ML	N19493 001	Oct 16, 1986
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AMINO ACIDS; MAGNESIUM CHLORIDE; POTASSIUM ACETATE; POTASSIUM CHLORIDE; SODIUM ACETATE

INJECTABLE; INJECTION

VEINAMINE 8%

HOSPIRA	8%;61MG/100ML;211MG/100ML;56MG/100ML;38 8MG/100ML	N17957 001	
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AMINO ACIDS; MAGNESIUM CHLORIDE; POTASSIUM CHLORIDE; POTASSIUM PHOSPHATE, DIBASIC; SODIUM CHLORIDE

INJECTABLE; INJECTION

AMINOSYN II 7% W/ ELECTROLYTES

HOSPIRA	7%;102MG/100ML;45MG/100ML;522MG/100ML;4 10MG/100ML	N19437 006	Apr 03, 1986
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AMINO ACIDS; MAGNESIUM CHLORIDE; POTASSIUM CHLORIDE; SODIUM CHLORIDE; SODIUM PHOSPHATE, DIBASIC, HEPTAHYDRATE

INJECTABLE; INJECTION

AMINOSYN II 3.5% M

HOSPIRA	3.5%;30MG/100ML;97MG/100ML;120MG/100ML; 49MG/100ML	N19437 007	Apr 03, 1986
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AMINOCAPROIC ACID

INJECTABLE; INJECTION

AMINOCAPROIC ACID

ABRAXIS PHARM	250MG/ML	N70522 001	Jun 17, 1986
BAXTER HLTHCARE	250MG/ML	N18590 001	Oct 29, 1982

AMINOPHYLLINE

ENEMA; RECTAL

SOMOPHYLLIN

FISONS	300MG/5ML	N18232 001	Apr 02, 1982
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INJECTABLE; INJECTION

AMINOPHYLLIN

GD SEARLE LLC	25MG/ML	N87243 001	May 24, 1982
	25MG/ML	N87621 001	May 24, 1982

AMINOPHYLLINE

ABRAXIS PHARM	25MG/ML	N84568 001	
	25MG/ML	N87200 001	
	25MG/ML	N87250 001	Jan 06, 1982
	25MG/ML	N87431 001	
	25MG/ML	N87886 001	Aug 30, 1983
	25MG/ML	N88407 001	Jan 25, 1984
ELKINS SINN	25MG/ML	N87239 001	
INTL MEDICATION	25MG/ML	N87867 001	Nov 10, 1983
	25MG/ML	N87868 001	Nov 10, 1983
KING PHARMS	25MG/ML	N86606 001	

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 15 (of 274)

AMINOPHYLLINE

INJECTABLE; INJECTION

AMINOPHYLLINE

PHARMA SERVE NY	25MG/ML	N87387	001	Jun 03, 1983
SMITH AND NEPHEW	25MG/ML	N88429	001	May 30, 1985
	25MG/ML	N88749	001	May 30, 1985

AMINOPHYLLINE IN SODIUM CHLORIDE 0.45% IN PLASTIC CONTAINER

HOSPIRA	100MG/100ML	N18924	001	Dec 12, 1984
	200MG/100ML	N18924	002	Dec 12, 1984
	400MG/100ML	N18924	003	Dec 12, 1984
	500MG/100ML	N18924	004	Dec 12, 1984

SOLUTION; ORAL

AMINOPHYLLINE

MORTON GROVE	105MG/5ML	N88156	001	Dec 05, 1983
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SOMOPHYLLIN

FISONS	105MG/5ML	N86466	001	
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SOMOPHYLLIN-DF

FISONS	105MG/5ML	N87045	001	
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TABLET; ORAL

AMINOPHYLLIN

GD SEARLE LLC	100MG	N02386	002	
	200MG	N02386	003	

AMINOPHYLLINE

ASCOT	100MG	N87522	001	Feb 12, 1982
	200MG	N87523	001	Feb 12, 1982

BARR	100MG	N88297	001	Aug 19, 1983
	200MG	N88298	001	Aug 19, 1983

DURAMED PHARMS BARR	100MG	N88182	001	Mar 31, 1983
	200MG	N88183	001	Mar 31, 1983

HALSEY	100MG	N84674	001	
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IMPAX LABS	100MG	N84574	001	
	200MG	N84576	001	

KV PHARM	100MG	N85284	001	
	200MG	N85289	001	

LANNETT	100MG	N84588	001	
	200MG	N84588	002	

PAL PAK	100MG	N84533	001	
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PANRAY	100MG	N84552	001	
	200MG	N84552	002	

PUREPAC PHARM	100MG	N84699	001	
	200MG	N85333	001	

ROXANE	100MG	N87500	001	Feb 09, 1982
	200MG	N87501	001	Feb 09, 1982

SANDOZ	100MG	N85261	003	
	100MG	N85262	002	

	200MG	N85261	002	
VALEANT PHARM INTL	200MG	N84563	001	

VANGARD	100MG	N88314	001	Oct 03, 1983
	200MG	N88319	001	Oct 03, 1983

VINTAGE PHARMS	100MG	N85409	001	
	200MG	N85410	001	

WATSON LABS	100MG	N85567	001	
	200MG	N85564	001	

TABLET, DELAYED RELEASE; ORAL

AMINOPHYLLINE

IMPAX LABS	100MG	N84577	001	
	200MG	N84575	001	

TABLICAPS	100MG	N84632	002	
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VALE	100MG	N84531	001	
	200MG	N84530	001	

TABLET, EXTENDED RELEASE; ORAL

PHYLLLOCONTIN

PURDUE FREDERICK	225MG	N86760	001	
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27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 16 (of 274)

AMINOSALICYLATE SODIUM

POWDER; ORAL		
P.A.S. SODIUM		
CENTURY PHARMS	4GM/PACKET	N80947 001
SODIUM AMINOSALICYLATE		
HEXCEL	100%	N80097 001
TABLET; ORAL		
PARASAL SODIUM		
PANRAY	500MG	N06811 006
	1GM	N06811 011
TEEBACIN		
CONSOLIDATED MIDLAND	500MG	N07320 002

AMINOSALICYLATE SODIUM; AMINOSALICYLIC ACID

TABLET; ORAL		
NEOPASALATE		
MEDPOINTE PHARM HLC	846MG;112MG	N80059 002

AMINOSALICYLIC ACID

TABLET; ORAL		
PARASAL		
PANRAY	500MG	N06811 001
	1GM	N06811 002

AMINOSALICYLIC ACID RESIN COMPLEX

POWDER; ORAL		
REZIPAS		
BRISTOL MYERS SQUIBB	EQ 500MG BASE/GM	N09052 001

AMITRIPTYLINE HYDROCHLORIDE

CONCENTRATE; ORAL			
ENDEP			
ROCHE	40MG/ML	N85749 001	
INJECTABLE; INJECTION			
AMITRIPTYLINE HYDROCHLORIDE			
WATSON LABS	10MG/ML	N85594 001	
ELAVIL			
ASTRAZENECA	10MG/ML	N12704 001	
TABLET; ORAL			
AMITID			
BRISTOL MYERS SQUIBB	10MG	N86454 001	
	25MG	N86454 002	
	50MG	N86454 003	
	75MG	N86454 004	
	100MG	N86454 005	
AMITRIL			
WARNER CHILCOTT	10MG	N83939 001	
	25MG	N83937 001	
	50MG	N83938 002	
	75MG	N84957 001	
	100MG	N85093 001	
	150MG	N86295 001	
AMITRIPTYLINE HYDROCHLORIDE			
AM THERAP	25MG	N88672 001	Nov 20, 1984
	50MG	N88673 001	Nov 20, 1984
	75MG	N88674 001	Nov 20, 1984
	100MG	N88675 001	Nov 20, 1984
COPLEY PHARM	10MG	N88421 001	Apr 30, 1984
	25MG	N88422 001	Apr 30, 1984
	50MG	N88423 001	Apr 30, 1984
	75MG	N88424 001	Apr 30, 1984
	100MG	N88425 001	Apr 30, 1984
	150MG	N88426 001	Apr 30, 1984
HALSEY	10MG	N85923 001	

DISCONTINUED DRUG PRODUCT LIST

6 - 17 (of 274)

AMITRIPTYLINE HYDROCHLORIDE

TABLET; ORAL

AMITRIPTYLINE HYDROCHLORIDE

HALSEY	25MG	N85922	001	
	50MG	N85925	001	
	50MG	N87557	001	Mar 05, 1982
	75MG	N85926	001	May 20, 1983
	100MG	N85927	001	May 20, 1983
LEDERLE	10MG	N86744	001	
	10MG	N87366	001	Jan 04, 1982
	25MG	N86746	001	
	25MG	N87367	001	May 03, 1982
	50MG	N86743	001	
	50MG	N87181	001	Jan 04, 1982
	75MG	N86745	001	
	75MG	N87369	001	Jan 04, 1982
	100MG	N86747	001	
	100MG	N87368	001	May 03, 1982
	150MG	N87370	001	Jan 04, 1982
MUTUAL PHARM	10MG	N85744	001	
	25MG	N85627	001	
	50MG	N85745	001	
	75MG	N85743	001	
	100MG	N85742	002	May 11, 1982
	150MG	N89423	001	Feb 17, 1987
PAR PHARM	10MG	N88697	001	Sep 25, 1984
	25MG	N88698	001	Sep 25, 1984
	50MG	N88699	001	Sep 25, 1984
	75MG	N88700	001	Sep 25, 1984
	100MG	N88701	001	Sep 25, 1984
	150MG	N88702	001	Sep 25, 1984
PUREPAC PHARM	10MG	N88075	001	Sep 16, 1983
	10MG	N88084	001	Jul 18, 1983
	25MG	N88076	001	May 20, 1983
	25MG	N88085	001	Jul 18, 1983
	50MG	N88077	001	Sep 16, 1983
	50MG	N88105	001	Jul 18, 1983
	75MG	N88078	001	Sep 16, 1983
	75MG	N88106	001	Jul 18, 1983
	100MG	N88079	001	Sep 16, 1983
	100MG	N88107	001	Jul 18, 1983
ROXANE	10MG	N86002	001	
	10MG	N86144	001	
	25MG	N85944	001	
	25MG	N86145	001	
	50MG	N85945	001	
	50MG	N86143	001	
	75MG	N86004	001	
	75MG	N86147	001	
	100MG	N86003	001	
	100MG	N86146	001	
	150MG	N86090	001	
	150MG	N86148	001	
SUPERPHARM	10MG	N88853	001	Nov 13, 1984
	25MG	N88854	001	Nov 13, 1984
	50MG	N88855	001	Nov 13, 1984
	75MG	N88856	001	Nov 13, 1984
	100MG	N88857	001	Nov 13, 1984
TEVA	10MG	N86610	001	
	25MG	N86859	001	
	50MG	N85032	001	
	50MG	N86857	001	
	75MG	N85030	001	
	75MG	N86860	001	

DISCONTINUED DRUG PRODUCT LIST

6 - 18 (of 274)

AMITRIPTYLINE HYDROCHLORIDE

TABLET; ORAL

AMITRIPTYLINE HYDROCHLORIDE

TEVA	100MG	N86854	001	
	150MG	N86853	001	
UCB INC	10MG	N85864	001	
	25MG	N85935	001	
	50MG	N85936	001	
	75MG	N86337	001	
	100MG	N86336	001	
	150MG	N86335	001	
USL PHARMA	25MG	N87775	001	Feb 10, 1982
VANGARD	10MG	N87632	001	Feb 01, 1982
	50MG	N87616	001	Feb 08, 1982
	75MG	N87617	001	Feb 05, 1982
	100MG	N87639	001	Feb 08, 1982
WATSON LABS	10MG	N85816	001	
	10MG	N88620	001	Mar 02, 1984
	25MG	N85817	001	
	25MG	N88621	001	Mar 02, 1984
	50MG	N85815	001	
	50MG	N88622	001	Mar 02, 1984
	75MG	N85819	001	
	75MG	N88633	001	Mar 02, 1984
	100MG	N85820	001	
	100MG	N88634	001	Mar 02, 1984
	150MG	N85821	001	
	150MG	N88635	001	Mar 02, 1984
	10MG	N87647	001	Mar 05, 1982
	25MG	N87278	001	
ELAVIL				
ASTRAZENECA	10MG	N12703	001	
	25MG	N12703	003	
	50MG	N12703	004	
	75MG	N12703	005	
	100MG	N12703	006	
	150MG	N12703	007	
ENDEP				
ROCHE	10MG	N83639	001	
	25MG	N83639	002	
	50MG	N83639	003	
	75MG	N83639	004	
	100MG	N83639	005	
	150MG	N85303	001	

AMITRIPTYLINE HYDROCHLORIDE; CHLORDIAZEPOXIDE

TABLET; ORAL

CHLORDIAZEPOXIDE AND AMITRIPTYLINE HYDROCHLORIDE

MUTUAL PHARM	EQ 12.5MG BASE;5MG	N70765	001	Dec 10, 1986
	EQ 25MG BASE;10MG	N70766	001	Dec 10, 1986
USL PHARMA	EQ 12.5MG BASE;5MG	N70477	001	Jan 12, 1988
	EQ 25MG BASE;10MG	N70478	001	Jan 12, 1988

AMITRIPTYLINE HYDROCHLORIDE; PERPHENAZINE

TABLET; ORAL

ETRAFON 2-10

SCHERING 10MG;2MG N14713 007

ETRAFON 2-25

SCHERING 25MG;2MG N14713 004

ETRAFON -A

SCHERING 10MG;4MG N14713 002

ETRAFON-FORTE

SCHERING 25MG;4MG N14713 006

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 19 (of 274)

AMITRIPTYLINE HYDROCHLORIDE; PERPHENAZINE

TABLET; ORAL

PERPHENAZINE AND AMITRIPTYLINE HYDROCHLORIDE

IVAX PHARMS	10MG;2MG	N70935	001	Sep 11, 1986
	10MG;4MG	N70937	001	Sep 11, 1986
	25MG;2MG	N70936	001	Sep 11, 1986
	25MG;4MG	N70938	001	Sep 11, 1986
	50MG;4MG	N70939	001	Sep 12, 1986
MUTUAL PHARM	10MG;2MG	N71077	001	Nov 12, 1986
	10MG;4MG	N71078	001	Nov 12, 1986
	25MG;2MG	N70297	001	Nov 12, 1986
	25MG;4MG	N71079	001	Nov 12, 1986
PAR PHARM	10MG;2MG	N70565	001	Sep 11, 1986
	10MG;4MG	N70620	001	Sep 11, 1986
	25MG;2MG	N70621	001	Sep 11, 1986
	25MG;4MG	N70595	001	Sep 11, 1986
	50MG;4MG	N70574	001	Sep 11, 1986
SANDOZ	10MG;2MG	N71062	001	Nov 27, 1987
	10MG;4MG	N71862	001	Dec 21, 1987
	25MG;2MG	N71063	001	Nov 27, 1987
	25MG;4MG	N71064	001	Nov 27, 1987
	50MG;4MG	N71863	001	Dec 21, 1987
WATSON LABS	10MG;2MG	N70373	001	Aug 25, 1986
	10MG;2MG	N72539	001	Feb 15, 1989
	10MG;2MG	N73007	001	Oct 17, 1991
	10MG;4MG	N70375	001	Aug 25, 1986
	10MG;4MG	N72540	001	Feb 15, 1989
	10MG;4MG	N73009	001	Oct 17, 1991
	25MG;2MG	N70374	001	Aug 25, 1986
	25MG;2MG	N72541	001	Feb 15, 1989
	25MG;2MG	N73008	001	Oct 17, 1991
	25MG;4MG	N70376	001	Aug 25, 1986
	25MG;4MG	N72134	001	Feb 15, 1989
	25MG;4MG	N73010	001	Oct 17, 1991
	50MG;4MG	N70377	001	Nov 04, 1986
	50MG;4MG	N71558	001	Mar 02, 1987
	50MG;4MG	N72135	001	Feb 15, 1989
TRIAVIL 2-10				
NEW RIVER	10MG;2MG	N14715	004	
TRIAVIL 2-25				
NEW RIVER	25MG;2MG	N14715	002	
TRIAVIL 4-10				
NEW RIVER	10MG;4MG	N14715	003	
TRIAVIL 4-25				
NEW RIVER	25MG;4MG	N14715	005	
TRIAVIL 4-50				
NEW RIVER	50MG;4MG	N14715	006	

AMLODIPINE MALEATE

TABLET; ORAL

AMVAZ

DR REDDYS LABS INC	2.5MG	N21435	001	Oct 31, 2003
	5MG	N21435	002	Oct 31, 2003
	10MG	N21435	003	Oct 31, 2003

AMMONIUM CHLORIDE

INJECTABLE; INJECTION

AMMONIUM CHLORIDE

ABBOTT	5MEQ/ML	N83130	001	
GD SEARLE LLC	3MEQ/ML	N86205	001	
AMMONIUM CHLORIDE 0.9% IN NORMAL SALINE				
MCGAW	900MG/100ML	N06580	001	
AMMONIUM CHLORIDE 2.14%				
B BRAUN	40MEQ/100ML	N85734	001	

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 20 (of 274)

AMODIAQUINE HYDROCHLORIDE

TABLET; ORAL

CAMOQUIN HYDROCHLORIDE

PARKE DAVIS

EQ 200MG BASE

N06441 001

AMOXAPINE

TABLET; ORAL

ASENDIN

LEDERLE

25MG

N18021 001

50MG

N18021 002

100MG

N18021 003

150MG

N18021 004

AMOXICILLIN

CAPSULE; ORAL

AMOXICILLIN

LABS ATRAL

250MG

N62528 001

Aug 07, 1985

500MG

N62528 002

Aug 07, 1985

MYLAN

250MG

N62067 001

500MG

N62067 002

TEVA

250MG

N63030 001

Feb 28, 1989

500MG

N63031 001

Feb 28, 1989

AMOXIL

GLAXOSMITHKLINE

250MG

N50459 001

250MG

N62216 003

500MG

N50459 002

TRIMOX

APOTHECON

250MG

N62098 001

250MG

N62152 001

250MG

N63099 001

Mar 20, 1992

500MG

N62098 002

500MG

N62152 002

500MG

N63099 002

Mar 20, 1992

UTIMOX

PARKE DAVIS

250MG

N62107 001

500MG

N62107 002

WYMOX

WYETH AYERST

250MG

N62120 001

500MG

N62120 002

FOR SUSPENSION; ORAL

AMOXICILLIN

MYLAN

125MG/5ML

N62090 001

250MG/5ML

N62090 002

AMOXIL

GLAXOSMITHKLINE

50MG/ML

N50460 005

125MG/5ML

N50460 001

250MG/5ML

N50460 002

LAROTID

GLAXOSMITHKLINE

50MG/ML

N50460 006

POLYMOX

APOTHECON

125MG/5ML

N61851 001

125MG/5ML

N62323 001

250MG/5ML

N61851 002

250MG/5ML

N62323 002

TRIMOX

APOTHECON

125MG/5ML

N62099 001

125MG/5ML

N62154 001

250MG/5ML

N62099 002

250MG/5ML

N62154 002

UTIMOX

PARKE DAVIS

125MG/5ML

N62127 001

250MG/5ML

N62127 002

WYMOX

WYETH AYERST

125MG/5ML

N62131 001

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 21 (of 274)

AMOXICILLIN

FOR SUSPENSION; ORAL

WYMOX

WYETH AYERST

250MG/5ML

N62131 002

TABLET, CHEWABLE; ORAL

AMOXICILLIN

APOTHECON

125MG

N64131 001

May 06, 1996

250MG

N64131 002

May 06, 1996

AMOXIL

GLAXOSMITHKLINE

125MG

N50542 002

250MG

N50542 001

AMPHETAMINE ADIPATE; AMPHETAMINE SULFATE; DEXTROAMPHETAMINE ADIPATE;
DEXTROAMPHETAMINE SULFATE

CAPSULE; ORAL

DELCOBESE

TEVA

1.25MG;1.25MG;1.25MG;1.25MG **Federal
Register determination that product
was not discontinued or withdrawn for
safety or efficacy reasons**

N83564 001

2.5MG;2.5MG;2.5MG;2.5MG **Federal
Register determination that product
was not discontinued or withdrawn for
safety or efficacy reasons**

N83564 002

3.75MG;3.75MG;3.75MG;3.75MG **Federal
Register determination that product
was not discontinued or withdrawn for
safety or efficacy reasons**

N83564 003

5MG;5MG;5MG;5MG **Federal Register
determination that product was not
discontinued or withdrawn for safety
or efficacy reasons**

N83564 004

TABLET; ORAL

DELCOBESE

TEVA

1.25MG;1.25MG;1.25MG;1.25MG **Federal
Register determination that product
was not discontinued or withdrawn for
safety or efficacy reasons**

N83563 004

2.5MG;2.5MG;2.5MG;2.5MG **Federal
Register determination that product
was not discontinued or withdrawn for
safety or efficacy reasons**

N83563 003

3.75MG;3.75MG;3.75MG;3.75MG **Federal
Register determination that product
was not discontinued or withdrawn for
safety or efficacy reasons**

N83563 002

5MG;5MG;5MG;5MG **Federal Register
determination that product was not
discontinued or withdrawn for safety
or efficacy reasons**

N83563 001

AMPHETAMINE RESIN COMPLEX; DEXTROAMPHETAMINE RESIN COMPLEX

CAPSULE, EXTENDED RELEASE; ORAL

BIPHETAMINE 12.5

UCB INC

EQ 6.25MG BASE;EQ 6.25MG BASE

N10093 007

BIPHETAMINE 20

UCB INC

EQ 10MG BASE;EQ 10MG BASE

N10093 003

BIPHETAMINE 7.5

UCB INC

EQ 3.75MG BASE;EQ 3.75MG BASE

N10093 009

AMPHETAMINE SULFATE

TABLET; ORAL

AMPHETAMINE SULFATE

LANNETT

5MG

N83901 001

Aug 31, 1984

10MG

N83901 002

Aug 31, 1984

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 22 (of 274)

AMPHOTERICIN B

INJECTABLE; INJECTION

AMPHOTERICIN B

ABBOTT

50MG/VIAL

N64141 001

Dec 23, 1996

ABRAXIS PHARM

50MG/VIAL

N62728 001

Apr 13, 1987

LOTION; TOPICAL

FUNGIZONE

APOTHECON

3%

N60570 001

OINTMENT; TOPICAL

FUNGIZONE

APOTHECON

3%

N50313 001

SUSPENSION; ORAL

FUNGIZONE

BRISTOL MYERS SQUIBB

100MG/ML

N50341 003

AMPICILLIN SODIUM

INJECTABLE; INJECTION

AMPICILLIN SODIUM

APOTHECON

EQ 125MG BASE/VIAL

N62860 001

Feb 05, 1988

EQ 250MG BASE/VIAL

N62860 002

Feb 05, 1988

EQ 500MG BASE/VIAL

N62860 003

Feb 05, 1988

EQ 1GM BASE/VIAL

N62860 004

Feb 05, 1988

EQ 2GM BASE/VIAL

N62860 005

Feb 05, 1988

BAXTER HLTHCARE

EQ 125MG BASE/VIAL

N62692 001

Jun 24, 1986

EQ 250MG BASE/VIAL

N62692 002

Jun 24, 1986

EQ 500MG BASE/VIAL

N62692 003

Jun 24, 1986

EQ 1GM BASE/VIAL

N62692 004

Jun 24, 1986

EQ 2GM BASE/VIAL

N62692 005

Jun 24, 1986

EQ 10GM BASE/VIAL

N62692 006

Jun 24, 1986

CONSOLIDATED PHARM

EQ 125MG BASE/VIAL

N61936 005

EQ 250MG BASE/VIAL

N61936 001

EQ 500MG BASE/VIAL

N61936 002

EQ 1GM BASE/VIAL

N61936 003

EQ 2GM BASE/VIAL

N61936 004

HANFORD GC

EQ 125MG BASE/VIAL

N63143 001

Apr 15, 1993

EQ 250MG BASE/VIAL

N63145 001

Apr 15, 1993

EQ 500MG BASE/VIAL

N63146 001

Apr 15, 1993

EQ 500MG BASE/VIAL

N63147 001

Apr 15, 1993

EQ 1GM BASE/VIAL

N62772 001

Apr 15, 1993

EQ 1GM BASE/VIAL

N63139 001

Apr 15, 1993

EQ 2GM BASE/VIAL

N63140 001

Apr 15, 1993

EQ 2GM BASE/VIAL

N63141 001

Apr 15, 1993

EQ 10GM BASE/VIAL

N63142 001

Apr 15, 1993

INTL MEDICATION

EQ 1GM BASE/VIAL

N62634 002

Jan 09, 1987

EQ 2GM BASE/VIAL

N62634 003

Jan 09, 1987

LILLY

EQ 500MG BASE/VIAL

N62565 001

Apr 04, 1985

EQ 1GM BASE/VIAL

N62565 002

Apr 04, 1985

EQ 2GM BASE/VIAL

N62565 003

Jun 24, 1986

MARSAM PHARMS LLC

EQ 125MG BASE/VIAL

N62816 001

Oct 24, 1988

EQ 250MG BASE/VIAL

N62816 002

Oct 24, 1988

EQ 500MG BASE/VIAL

N62816 003

Oct 24, 1988

EQ 1GM BASE/VIAL

N62816 004

Oct 24, 1988

EQ 2GM BASE/VIAL

N62816 005

Oct 24, 1988

EQ 10GM BASE/VIAL

N62994 001

Sep 15, 1988

OMNIPEN-N

WYETH AYERST

EQ 125MG BASE/VIAL

N60626 001

EQ 125MG BASE/VIAL

N62718 001

Dec 16, 1986

EQ 250MG BASE/VIAL

N60626 002

EQ 250MG BASE/VIAL

N62718 002

Dec 16, 1986

EQ 500MG BASE/VIAL

N60626 003

EQ 500MG BASE/VIAL

N62718 003

Dec 16, 1986

EQ 1GM BASE/VIAL

N60626 004

EQ 1GM BASE/VIAL

N62718 004

Dec 16, 1986

EQ 2GM BASE/VIAL

N60626 005

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 23 (of 274)

AMPICILLIN SODIUM

INJECTABLE; INJECTION

OMNIPEN-N

WYETH AYERST

EQ 2GM BASE/VIAL

N62718 005

Dec 16, 1986

PENBRITIN-S

WYETH AYERST

EQ 125MG BASE/VIAL

N50072 001

EQ 250MG BASE/VIAL

N50072 002

EQ 500MG BASE/VIAL

N50072 003

EQ 1GM BASE/VIAL

N50072 004

EQ 2GM BASE/VIAL

N50072 005

EQ 4GM BASE/VIAL

N50072 006

POLYCILLIN-N

BRISTOL

EQ 125MG BASE/VIAL

N50309 001

EQ 250MG BASE/VIAL

N50309 002

EQ 500MG BASE/VIAL

N50309 003

EQ 1GM BASE/VIAL

N50309 004

EQ 2GM BASE/VIAL

N50309 005

TOTACILLIN-N

GLAXOSMITHKLINE

EQ 125MG BASE/VIAL

N60677 001

EQ 250MG BASE/VIAL

N60677 002

EQ 500MG BASE/VIAL

N60677 003

EQ 1GM BASE/VIAL

N60677 004

EQ 1GM BASE/VIAL

N62727 001

Dec 19, 1986

EQ 2GM BASE/VIAL

N60677 005

EQ 2GM BASE/VIAL

N62727 002

Dec 19, 1986

EQ 10GM BASE/VIAL

N60677 006

AMPICILLIN SODIUM; SULBACTAM SODIUM

INJECTABLE; INJECTION

UNASYN

PFIZER

EQ 500MG BASE/VIAL;EQ 250MG BASE/VIAL

N50608 003

Dec 31, 1986

AMPICILLIN/AMPICILLIN TRIHYDRATE

CAPSULE; ORAL

AMCILL

PARKE DAVIS

EQ 250MG BASE

N62041 001

EQ 500MG BASE

N62041 002

AMPICILLIN

LEDERLE

EQ 250MG BASE

N62208 001

EQ 500MG BASE

N62208 002

VITARINE

EQ 250MG BASE

N61387 001

EQ 500MG BASE

N61387 003

AMPICILLIN TRIHYDRATE

AM ANTIBIOTICS

EQ 250MG BASE

N61602 001

EQ 500MG BASE

N61602 002

BIOCHEMIE

EQ 250MG BASE

N64082 001

Aug 29, 1995

EQ 500MG BASE

N64082 002

Aug 29, 1995

IVAX PHARMS

EQ 250MG BASE

N60765 001

EQ 500MG BASE

N60765 002

MYLAN

EQ 250MG BASE

N61755 001

EQ 500MG BASE

N61755 002

PUREPAC PHARM

EQ 250MG BASE

N61853 001

EQ 500MG BASE

N61853 002

TEVA

EQ 250MG BASE

N61502 001

EQ 500MG BASE

N61502 002

OMNIPEN (AMPICILLIN)

WYETH AYERST

250MG

N60624 001

500MG

N60624 002

PENBRITIN

WYETH AYERST

EQ 250MG BASE

N60908 001

EQ 500MG BASE

N60908 002

PFIZERPEN-A

PFIZER

EQ 250MG BASE

N62050 001

EQ 500MG BASE

N62050 002

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 24 (of 274)

AMPICILLIN/AMPICILLIN TRIHYDRATE

CAPSULE; ORAL

POLYCILLIN

BRISTOL

EQ 250MG BASE

N50310 001

EQ 500MG BASE

N50310 002

PRINCIPEN

APOTHECON

EQ 250MG BASE

N61392 001

EQ 250MG BASE

N62888 001

Mar 04, 1988

EQ 500MG BASE

N61392 002

EQ 500MG BASE

N62888 002

Mar 04, 1988

PRINCIPEN '250'

APOTHECON

EQ 250MG BASE

N50056 001

EQ 250MG BASE

N62157 002

PRINCIPEN '500'

APOTHECON

EQ 500MG BASE

N50056 002

EQ 500MG BASE

N62157 001

TOTACILLIN

GLAXOSMITHKLINE

EQ 250MG BASE

N60060 001

EQ 250MG BASE

N62212 001

EQ 500MG BASE

N60060 002

EQ 500MG BASE

N62212 002

FOR SUSPENSION; ORAL

AMCILL

PARKE DAVIS

EQ 125MG BASE/5ML

N62030 001

EQ 250MG BASE/5ML

N62030 002

AMPICILLIN TRIHYDRATE

AM ANTIBIOTICS

EQ 125MG BASE/5ML

N61601 001

EQ 250MG BASE/5ML

N61601 002

MYLAN

EQ 125MG BASE/5ML

N61829 002

EQ 250MG BASE/5ML

N61829 001

PUREPAC PHARM

EQ 125MG BASE/5ML

N61980 001

TEVA

EQ 125MG BASE/5ML

N61370 001

EQ 250MG BASE/5ML

N61370 002

OMNIPEN (AMPICILLIN)

WYETH AYERST

100MG/ML

N60625 001

125MG/5ML

N60625 002

250MG/5ML

N60625 003

500MG/5ML

N60625 004

PENBRITIN

WYETH AYERST

EQ 100MG BASE/ML

N50019 001

EQ 125MG BASE/5ML

N50019 002

EQ 250MG BASE/5ML

N50019 003

PFIZERPEN-A

PFIZER

EQ 125MG BASE/5ML

N62049 001

EQ 250MG BASE/5ML

N62049 002

POLYCILLIN

APOTHECON

EQ 125MG BASE/5ML

N62297 001

EQ 250MG BASE/5ML

N62297 002

BRISTOL

EQ 100MG BASE/ML

N50308 004

EQ 125MG BASE/5ML

N50308 001

EQ 250MG BASE/5ML

N50308 002

EQ 500MG BASE/5ML

N50308 003

PRINCIPEN

APOTHECON

EQ 100MG BASE/ML

N61394 001

PRINCIPEN '125'

APOTHECON

EQ 125MG BASE/5ML

N60127 002

EQ 125MG BASE/5ML

N62151 001

PRINCIPEN '250'

APOTHECON

EQ 250MG BASE/5ML

N60127 001

EQ 250MG BASE/5ML

N62151 002

TOTACILLIN

GLAXOSMITHKLINE

EQ 125MG BASE/5ML

N60666 001

EQ 125MG BASE/5ML

N62223 001

EQ 250MG BASE/5ML

N60666 002

DISCONTINUED DRUG PRODUCT LIST

6 - 25 (of 274)

AMPICILLIN/AMPICILLIN TRIHYDRATE

FOR SUSPENSION; ORAL		
TOTACILLIN		
GLAXOSMITHKLINE	EQ 250MG BASE/5ML	N62223 002
TABLET, CHEWABLE; ORAL		
POLYCILLIN		
BRISTOL	EQ 125MG BASE	N50093 001

AMPICILLIN/AMPICILLIN TRIHYDRATE; PROBENECID

CAPSULE; ORAL		
PRINCIPEN W/ PROBENECID		
APOTHECON	EQ 389MG BASE;111MG	N50488 001
	EQ 389MG BASE;111MG	N62150 001
FOR SUSPENSION; ORAL		
POLYCILLIN-PRB		
APOTHECON	EQ 3.5GM BASE/BOT;1GM/BOT	N61898 001
BRISTOL	EQ 3.5GM BASE/BOT;1GM/BOT	N50457 001
PROBAMPACIN		
TEVA	EQ 3.5GM BASE/BOT;1GM/BOT	N61741 001

AMPRENAVIR

CAPSULE; ORAL		
AGENERASE		
GLAXOSMITHKLINE	150MG	N21007 002 Apr 15, 1999

ANILERIDINE HYDROCHLORIDE

TABLET; ORAL		
LERITINE		
MERCK	EQ 25MG BASE	N10585 002

ANILERIDINE PHOSPHATE

INJECTABLE; INJECTION		
LERITINE		
MERCK	25MG/ML	N10520 003

ANISINDIONE

TABLET; ORAL		
MIRADON		
SCHERING	50MG	N10909 003

ANISOTROPINE METHYLBROMIDE

TABLET; ORAL		
ANISOTROPINE METHYLBROMIDE		
WATSON LABS	50MG	N86046 001
VALPIN 50		
ENDO PHARMS	50MG	N13428 001

APOMORPHINE HYDROCHLORIDE

INJECTABLE; SUBCUTANEOUS		
APOKYN		
VERNALIS	20MG/2ML (10MG/ML)	N21264 001 Apr 20, 2004

ARBUTAMINE HYDROCHLORIDE

INJECTABLE; INJECTION		
GENESA		
GENSIA AUTOMEDICS	0.05MG/ML	N20420 001 Sep 12, 1997

ARDEPARIN SODIUM

INJECTABLE; INJECTION		
NORMIFLO		
PHARMACIA AND UPJOHN	5,000 UNITS/0.5ML **Federal Register determination that product was not discontinued or withdrawn for safety or efficacy reasons**	N20227 002 May 23, 1997

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 26 (of 274)

ARDEPARIN SODIUMINJECTABLE; INJECTION
NORMIFLO

PHARMACIA AND UPJOHN	10,000 UNITS/0.5ML **Federal Register determination that product was not discontinued or withdrawn for safety or efficacy reasons**	N20227 001	May 23, 1997
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ASCORBIC ACID; BIOTIN; CYANOCOBALAMIN; DEXPANTHENOL; ERGOCALCIFEROL; FOLIC ACID; NIACINAMIDE; PYRIDOXINE HYDROCHLORIDE; RIBOFLAVIN PHOSPHATE SODIUM; THIAMINE HYDROCHLORIDE; VITAMIN A PALMITATE; VITAMIN EINJECTABLE; INJECTION
BEROCCA PN

ROCHE	50MG/ML;0.03MG/ML;0.0025MG/ML;7.5MG/ML;100 IU/ML;0.2MG/ML;20MG/ML;2MG/ML;1.8MG/ML;1.5MG/ML;1,650 IU/ML;5 IU/ML	N06071 003	Oct 10, 1985
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ASCORBIC ACID; BIOTIN; CYANOCOBALAMIN; DEXPANTHENOL; ERGOCALCIFEROL; FOLIC ACID; NIACINAMIDE; PYRIDOXINE HYDROCHLORIDE; RIBOFLAVIN PHOSPHATE SODIUM; THIAMINE HYDROCHLORIDE; VITAMIN A; VITAMIN EINJECTABLE; INJECTION
M.V.C. 9+3

ABRAXIS PHARM	10MG/ML;0.006MG/ML;0.5UGM/ML;1.5MG/ML;20 IU/ML;0.04MG/ML;4MG/ML;0.4MG/ML;0.36MG/ML;0.3MG/ML;330 UNITS/ML;1 IU/ML	N18440 002	Aug 08, 1985
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M.V.I. -12

MAYNE PHARMA USA	10MG/ML;0.006MG/ML;0.5UGM/ML;1.5MG/ML;20 IU/ML;0.04MG/ML;4MG/ML;0.4MG/ML;0.36MG/ML;0.3MG/ML;330 UNITS/ML;1 IU/ML	N08809 004	Aug 08, 1985
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MVC PLUS

WATSON LABS	10MG/ML;0.006MG/ML;0.5UGM/ML;1.5MG/ML;20 IU/ML;0.04MG/ML;4MG/ML;0.4MG/ML;0.36MG/ML;0.3MG/ML;330 UNITS/ML;1 IU/ML	N18439 002	Aug 08, 1985
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ASCORBIC ACID; BIOTIN; CYANOCOBALAMIN; DEXPANTHENOL; ERGOCALCIFEROL; FOLIC ACID; NIACINAMIDE; PYRIDOXINE HYDROCHLORIDE; RIBOFLAVIN; THIAMINE HYDROCHLORIDE; VITAMIN A; VITAMIN EINJECTABLE; INJECTION
M.V.I. -12

MAYNE PHARMA USA	20MG/ML;0.006MG/ML;0.5UGM/ML;1.5MG/ML;20 IU/ML;0.6MG/ML;4MG/ML;0.4MG/ML;0.36MG/ML;0.6MG/ML;330 UNITS/ML;1 IU/ML	N08809 005	Apr 22, 2004
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ASCORBIC ACID; BIOTIN; CYANOCOBALAMIN; DEXPANTHENOL; ERGOCALCIFEROL; FOLIC ACID; NIACINAMIDE; PYRIDOXINE; RIBOFLAVIN PHOSPHATE SODIUM; THIAMINE; VITAMIN A; VITAMIN EINJECTABLE; INJECTION
M.V.I. -12 LYOPHILIZED

ASTRAZENECA	100MG/VIAL;0.06MG/VIAL;0.005MG/VIAL;15MG/VIAL;5UGM/VIAL;0.4MG/VIAL;40MG/VIAL;4MG/VIAL;3.6MG/VIAL;3MG/VIAL;1MG/VIAL;10MG/VIAL	N18933 002	Aug 08, 1985
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ASCORBIC ACID; BIOTIN; CYANOCOBALAMIN; ERGOCALCIFEROL; FOLIC ACID; NIACINAMIDE; PANTOTHENIC ACID; PHYTONADIONE; PYRIDOXINE; RIBOFLAVIN; THIAMINE; VITAMIN A PALMITATE; VITAMIN EINJECTABLE; INJECTION
VITAPED

HOSPIRA	N/A, 80MG/VIAL;N/A, 0.02MG/VIAL;N/A, 0.001MG/VIAL;400 IU/10ML,N/A;N/A, 0.14MG/VIAL;N/A, 17MG/VIAL;N/A, 5MG/VIAL;0.2MG/10ML,N/A;N/A, 1MG/VIAL;N/A, 1.4MG/VIAL;N/A, 1.2MG/VIAL;EQ 2,300 UNITS BASE/10ML,N/A;7 IU/10ML,N/A	N20176 001	Dec 29, 1993
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27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 27 (of 274)

ASPIRIN

TABLET, EXTENDED RELEASE; ORAL

8-HOUR BAYER

BAYER

650MG

N16030 001

MEASURIN

BAYER

650MG

N16030 002

ASPIRIN; BUTALBITAL

TABLET; ORAL

AXOTAL

SAVAGE LABS

650MG;50MG

N88305 001 Oct 13, 1983

ASPIRIN; BUTALBITAL; CAFFEINE

CAPSULE; ORAL

BUTALBITAL, ASPIRIN AND CAFFEINE

WATSON LABS

325MG;50MG;40MG

N86231 002 Feb 12, 1985

TABLET; ORAL

BUTALBITAL ASPIRIN AND CAFFEINE

QUANTUM PHARMICS

325MG;50MG;40MG

N88972 001 Jun 18, 1985

BUTALBITAL COMPOUND

IVAX PHARMS

325MG;50MG;40MG

N85441 002 Oct 31, 1984

BUTALBITAL W/ ASPIRIN & CAFFEINE

PHARMERAL

325MG;50MG;40MG

N87048 002 Dec 09, 1983

BUTALBITAL, ASPIRIN AND CAFFEINE

HALSEY

325MG;50MG;40MG

N89448 001 Dec 01, 1986

WATSON LABS

325MG;50MG;40MG

N86237 002 Mar 23, 1984

FIORINAL

WATSON PHARMS

325MG;50MG;40MG

N17534 003 Apr 16, 1986

LANORINAL

LANNETT

325MG;50MG;40MG

N86986 002 Oct 18, 1985

ASPIRIN; BUTALBITAL; CAFFEINE; CODEINE PHOSPHATE

CAPSULE; ORAL

BUTALBITAL, ASPIRIN, CAFFEINE, AND CODEINE PHOSPHATE

ENDO PHARMS

325MG;50MG;40MG;30MG

N75351 001 Mar 05, 1999

ASPIRIN; CAFFEINE; PROPOXYPHENE HYDROCHLORIDE

CAPSULE; ORAL

COMPOUND 65

ALRA

389MG;32.4MG;65MG

N84553 002 Aug 17, 1983

DARVON COMPOUND

XANODYNE PHARM

389MG;32.4MG;32MG

N10996 006 Mar 08, 1983

PROPOXYPHENE COMPOUND 65

IVAX PHARMS

389MG;32.4MG;65MG

N83077 002 Dec 07, 1984

SANDOZ

389MG;32.4MG;65MG

N80044 002 Sep 16, 1983

PROPOXYPHENE COMPOUND-65

SANDOZ

389MG;32.4MG;65MG

N83101 002 Jun 24, 1985

PROPOXYPHENE HYDROCHLORIDE W/ ASPIRIN AND CAFFEINE

WATSON LABS

389MG;32.4MG;65MG

N85732 002 Sep 03, 1984

ASPIRIN; CARISOPRODOL

TABLET; ORAL

CARISOPRODOL COMPOUND

WATSON LABS

325MG;200MG

N88809 001 Oct 03, 1985

ASPIRIN; HYDROCODONE BITARTRATE

TABLET; ORAL

VICOPRIN

ABBOTT

500MG;5MG

N86333 001 Sep 14, 1983

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 28 (of 274)

ASPIRIN; MEPROBAMATE

TABLET; ORAL				
MEPRO-ASPIRIN				
SANDOZ	325MG;200MG	N89127	001	Mar 02, 1987
MEPROBAMATE AND ASPIRIN				
PAR PHARM	325MG;200MG	N89126	001	Aug 19, 1986
MICRAININ				
MEDPOINTE PHARM HLC	325MG;200MG	N84978	001	
Q-GESIC				
QUANTUM PHARMICS	325MG;200MG	N88740	001	Jun 01, 1984

ASPIRIN; METHOCARBAMOL

TABLET; ORAL				
METHOCARBAMOL AND ASPIRIN				
MCNEIL	325MG;400MG	N89193	001	Feb 12, 1986
ROBAXISAL				
ROBINS AH	325MG;400MG	N12281	001	

ASPIRIN; OXYCODONE HYDROCHLORIDE; OXYCODONE TEREPHTHALATE

TABLET; ORAL				
CODOXY				
HALSEY	325MG;4.5MG;0.38MG	N87464	001	Jul 01, 1982
OXYCODONE AND ASPIRIN (HALF-STRENGTH)				
ROXANE	325MG;2.25MG;0.19MG	N87742	001	Jun 04, 1982
PERCODAN-DEMI				
ENDO PHARMS	325MG;2.25MG;0.19MG	N07337	005	
ROXIPRIN				
ROXANE	325MG;4.5MG;0.38MG	N87743	001	Jun 04, 1982

ASPIRIN; PENTAZOCINE HYDROCHLORIDE

TABLET; ORAL				
TALWIN COMPOUND				
SANOFI AVENTIS US	325MG;EQ 12.5MG BASE	N16891	001	

ASPIRIN; PROPOXYPHENE HYDROCHLORIDE

CAPSULE; ORAL				
DARVON W/ ASA				
XANODYNE PHARM	325MG;65MG	N10996	005	

ASPIRIN; PROPOXYPHENE NAPSYLATE

CAPSULE; ORAL				
DARVON-N W/ ASA				
AAIPHARMA LLC	325MG;100MG	N16829	001	
TABLET; ORAL				
DARVON-N W/ ASA				
AAIPHARMA LLC	325MG;100MG	N16863	001	

ATENOLOL

TABLET; ORAL				
ATENOLOL				
ABLE	25MG	N76907	001	Jul 30, 2004
	50MG	N76907	002	Jul 30, 2004
	100MG	N76907	003	Jul 30, 2004
		N73317	001	Mar 20, 1992
APOTHECON	50MG	N73317	001	Mar 20, 1992
	100MG	N73318	001	Mar 20, 1992

ATOMOXETINE HYDROCHLORIDE

CAPSULE; ORAL				
STRATTERA				
LILLY	5MG	N21411	001	Nov 26, 2002

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 29 (of 274)

ATOVAQUONETABLET; ORAL
MEPRON

GLAXOSMITHKLINE 250MG

N20259 001 Nov 25, 1992

ATracurium BESYLATE

INJECTABLE; INJECTION

ATracurium BESYLATE PRESERVATIVE FREE

BAXTER HLTHCARE 10MG/ML

N74825 001 Sep 30, 1997

ATROPINE

INJECTABLE; INJECTION

ATROPINE

SOLVAY EQ 2MG SULFATE/0.7ML

N71295 001 Jan 30, 1987

ATROPINE SULFATE

AEROSOL, METERED; INHALATION

ATROPINE SULFATE

US ARMY EQ 0.36MG BASE/INH

N20056 001 Sep 19, 1990

ATROPINE SULFATE; DIFENOXIN HYDROCHLORIDE

TABLET; ORAL

MOTOFEN HALF-STRENGTH

VALEANT 0.025MG;0.5MG

N17744 001

ATROPINE SULFATE; DIPHENOXYLATE HYDROCHLORIDE

SOLUTION; ORAL

COLONAIID

MEDPOINTE PHARM HLC 0.025MG/5ML;2.5MG/5ML

N85735 001

LOMANATE

ALPHARMA US PHARMS 0.025MG/5ML;2.5MG/5ML

N85746 001

TABLET; ORAL

COLONAIID

MEDPOINTE PHARM HLC 0.025MG;2.5MG

N85737 001

DI-ATRO

MD PHARM 0.025MG;2.5MG

N85266 001

DIPHENOXYLATE HYDROCHLORIDE AND ATROPINE SULFATE

ABLE 0.025MG;2.5MG

N40395 001 Nov 27, 2000

ASCOT 0.025MG;2.5MG

N87934 001 Jul 19, 1983

HEATHER 0.025MG;2.5MG

N86798 001

INWOOD LABS 0.025MG;2.5MG

N85509 001

LEDERLE 0.025MG;2.5MG

N86950 001

MUTUAL PHARM 0.025MG;2.5MG

N85506 001

PARKE DAVIS 0.025MG;2.5MG

N87131 001

R AND S PHARMA 0.025MG;2.5MG

N85035 001

ROXANE 0.025MG;2.5MG

N86057 001

WATSON LABS 0.025MG;2.5MG

N85876 001

WEST WARD 0.025MG;2.5MG

N87765 001 Mar 15, 1982

DIPHENOXYLATE HYDROCHLORIDE W/ ATROPINE SULFATE

KV PHARM 0.025MG;2.5MG

N85659 001

LEINER 0.025MG;2.5MG

N85766 001

USL PHARMA 0.025MG;2.5MG

N87842 001 Mar 29, 1982

VALEANT PHARM INTL 0.025MG;2.5MG

N87195 001 Feb 16, 1982

LOGEN

SUPERPHARM 0.025MG;2.5MG

N88962 001 May 10, 1985

LO-TROL

VANGARD 0.025MG;2.5MG

N88009 001 Mar 25, 1983

LOW-QUEL

HALSEY 0.025MG;2.5MG

N85211 001

ATROPINE SULFATE; MEPERIDINE HYDROCHLORIDE

INJECTABLE; INJECTION

ATROPINE AND DEMEROL

ABBOTT 0.4MG/ML;50MG/ML

N87853 001 Nov 26, 1982

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 30 (of 274)

ATROPINE SULFATE; MEPERIDINE HYDROCHLORIDEINJECTABLE; INJECTION
ATROPINE AND DEMEROL

ABBOTT

0.4MG/ML;75MG/ML
0.4MG/ML;100MG/ML

N87847 001 Nov 26, 1982

N87848 001 Nov 26, 1982

MEPERIDINE AND ATROPINE SULFATE

WYETH AYERST

0.4MG/ML;50MG/ML
0.4MG/ML;75MG/ML
0.4MG/ML;100MG/ML

N85121 001

N85121 002

N85121 003

ATROPINE; PRALIDOXIME CHLORIDEINJECTABLE; INTRAMUSCULAR
ATNAA

US ARMY

2.1MG/0.7ML,N/A;N/A,600MG/2ML

N21175 001 Jan 17, 2002

AZATADINE MALEATETABLET; ORAL
OPTIMINE

SCHERING

1MG

N17601 001

AZATADINE MALEATE; PSEUDOEPHEDRINE SULFATETABLET, EXTENDED RELEASE; ORAL
TRINALIN

SCHERING

1MG;120MG

N18506 001 Mar 23, 1982

AZATHIOPRINETABLET; ORAL
IMURAN

PROMETHEUS LABS

25MG

N16324 002

AZATHIOPRINE SODIUMINJECTABLE; INJECTION
IMURAN

PROMETHEUS LABS

EQ 100MG BASE/VIAL

N17391 001

AZITHROMYCINCAPSULE; ORAL
ZITHROMAX

PFIZER

EQ 250MG BASE **Federal Register
determination that product was not
discontinued or withdrawn for safety
or efficacy reasons**

N50670 001 Nov 01, 1991

AZITHROMYCIN DIHYDRATE; TROVAFLOXACIN MESYLATEFOR SUSPENSION, TABLET; ORAL
TROVAN/ZITHROMAX COMPLIANCE PAK

PFIZER

EQ 1GM BASE,N/A;N/A,EQ 100MG BASE

N50762 001 Dec 18, 1998

AZLOCILLIN SODIUMINJECTABLE; INJECTION
AZLIN

BAYER PHARMS

EQ 2GM BASE/VIAL
EQ 2GM BASE/VIAL
EQ 2GM BASE/VIAL
EQ 3GM BASE/VIAL
EQ 3GM BASE/VIAL
EQ 3GM BASE/VIAL
EQ 4GM BASE/VIAL
EQ 4GM BASE/VIAL
EQ 4GM BASE/VIAL

N50562 001 Sep 03, 1982

N62388 001 Sep 08, 1982

N62417 001 Oct 12, 1982

N50562 002 Sep 03, 1982

N62388 002 Sep 08, 1982

N62417 002 Oct 12, 1982

N50562 003 Sep 03, 1982

N62388 003 Sep 08, 1982

N62417 003 Oct 12, 1982

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 31 (of 274)

AZTREONAM

INJECTABLE; INJECTION				
AZACTAM IN PLASTIC CONTAINER				
BRISTOL MYERS SQUIBB	10MG/ML	N50632	003	May 24, 1989

BACAMPICILLIN HYDROCHLORIDE

FOR SUSPENSION; ORAL				
SPECTROBID				
PFIZER	125MG/5ML	N50556	001	Mar 23, 1982
TABLET; ORAL				
SPECTROBID				
PFIZER	400MG	N50520	001	
	800MG	N50520	002	Sep 12, 1983

BACITRACIN

INJECTABLE; INJECTION				
BACITRACIN				
PFIZER	50,000 UNITS/VIAL	N60282	001	
OINTMENT; OPHTHALMIC				
BACIGUENT				
PHARMACIA AND UPJOHN	500 UNITS/GM	N60734	001	
BACITRACIN				
LILLY	500 UNITS/GM	N60687	001	
PHARMADERM	500 UNITS/GM	N62158	001	
PHARMAFAIR	500 UNITS/GM	N62453	001	Mar 28, 1984
OINTMENT; TOPICAL				
BACITRACIN				
COMBE	500 UNITS/GM	N62799	001	May 14, 1987
NASKA	500 UNITS/GM	N62857	001	Nov 13, 1987
POWDER; FOR RX COMPOUNDING				
BACITRACIN				
PADDOCK	5,000,000 UNITS/BOT	N62456	001	Jul 27, 1983

BACITRACIN ZINC

POWDER; FOR RX COMPOUNDING				
ZIBA-RX				
X GEN PHARMS	500,000 UNITS/BOT	N61737	001	

BACITRACIN ZINC; HYDROCORTISONE; NEOMYCIN SULFATE; POLYMYXIN B SULFATE

OINTMENT; OPHTHALMIC				
ZINC BACITRACIN, NEOMYCIN SULFATE, POLYMYXIN B SULFATE & HYDROCORTISONE				
PHARMAFAIR	400 UNITS/GM; 1%; EQ 3.5MG BASE/GM; 10,000 UNITS/GM	N62389	001	Jul 02, 1982
OINTMENT; TOPICAL				
NEOMYCIN & POLYMYXIN B SULFATES & BACITRACIN ZINC & HYDROCORTISONE				
PHARMAFAIR	400 UNITS/GM; 1%; EQ 3.5MG BASE/GM; 5,000 UNITS/GM	N62381	001	Sep 06, 1985

BACITRACIN ZINC; LIDOCAINE; NEOMYCIN SULFATE; POLYMYXIN B SULFATE

OINTMENT; TOPICAL				
LANABIOTIC				
COMBE	400 UNITS/GM; 40MG/GM; EQ 5MG BASE/GM; 5,000 UNITS/GM	N62499	001	Jun 03, 1985

BACITRACIN ZINC; NEOMYCIN SULFATE; POLYMYXIN B SULFATE

OINTMENT; OPHTHALMIC				
BACITRACIN ZINC-NEOMYCIN SULFATE-POLYMYXIN B SULFATE				
PHARMAFAIR	400 UNITS/GM; EQ 3.5MG BASE/GM; 10,000 UNITS/GM	N62386	001	Sep 09, 1982
BACITRACIN-NEOMYCIN-POLYMYXIN				
PHARMADERM	400 UNITS/GM; EQ 3.5MG BASE/GM; 5,000 UNITS/GM	N62167	001	
NEO-POLYCIN				
DOW PHARM	500 UNITS/GM; EQ 3.5MG BASE/GM; 10,000 UNITS/GM	N60647	001	

DISCONTINUED DRUG PRODUCT LIST

6 - 32 (of 274)

BACITRACIN ZINC; NEOMYCIN SULFATE; POLYMYXIN B SULFATE

OINTMENT; TOPICAL			
BACITRACIN ZINC-NEOMYCIN SULFATE-POLYMYXIN B SULFATE			
NASKA	400 UNITS/GM;EQ 3.5MG BASE/GM;5,000 UNITS/GM	N62833 001	Nov 09, 1987

BACITRACIN ZINC; POLYMYXIN B SULFATE

AEROSOL; TOPICAL			
POLYSPORIN			
GLAXOSMITHKLINE	10,000 UNITS/GM;2,000,000 UNITS/GM	N50167 002	Mar 01, 1985
OINTMENT; OPHTHALMIC			
OCUMYCIN			
PHARMAFAIR	500 UNITS/GM;10,000 UNITS/GM	N62430 001	Apr 08, 1983
OINTMENT; TOPICAL			
BACITRACIN ZINC-POLYMYXIN B SULFATE			
NASKA	500 UNITS/GM;10,000 UNITS/GM	N62849 001	Nov 13, 1987

BACITRACIN; HYDROCORTISONE ACETATE; NEOMYCIN SULFATE; POLYMYXIN B SULFATE

OINTMENT; OPHTHALMIC			
BACITRACIN-NEOMYCIN-POLYMYXIN W/ HYDROCORTISONE ACETATE			
ALTANA	400 UNITS/GM;1%;EQ 3.5MG BASE/GM;10,000 UNITS/GM	N60731 002	

BACITRACIN; NEOMYCIN SULFATE; POLYMYXIN B SULFATE

OINTMENT; OPHTHALMIC			
MYCITRACIN			
PHARMACIA AND UPJOHN	500 UNITS/GM;EQ 3.5MG BASE/GM;10,000 UNITS/GM	N61048 001	

BACITRACIN; POLYMYXIN B SULFATE

DISC; TOPICAL			
LANABIOTIC			
COMBE	500 UNITS/GM;5,000 UNITS/GM	N50598 001	Sep 22, 1986

BACLOFEN

TABLET; ORAL			
BACLOFEN			
TEVA	10MG	N73043 001	Feb 27, 1992
	20MG	N73044 001	Feb 27, 1992
USL PHARMA	10MG	N71260 001	May 06, 1988
	20MG	N71261 001	May 06, 1988
WATSON LABS	10MG	N74698 001	Aug 20, 1996
	20MG	N73093 001	Jan 28, 1994
	20MG	N74698 002	Aug 20, 1996
LIORESAL			
NOVARTIS	10MG	N17851 001	
	20MG	N17851 003	Jan 20, 1982

BECLOMETHASONE DIPROPIONATE

AEROSOL, METERED; INHALATION			
BECLOVENT			
GLAXOSMITHKLINE	0.042MG/INH	N18153 001	
VANCERIL			
SCHERING	0.042MG/INH	N17573 001	
VANCERIL DOUBLE STRENGTH			
SCHERING	0.084MG/INH	N20486 001	Dec 24, 1996
AEROSOL, METERED; NASAL			
BECONASE			
GLAXOSMITHKLINE	0.042MG/INH	N18584 001	
VANCENASE			
SCHERING	0.042MG/INH	N18521 001	

DISCONTINUED DRUG PRODUCT LIST

6 - 33 (of 274)

BECLOMETHASONE DIPROPIONATE MONOHYDRATE

SPRAY, METERED; NASAL
VANCENASE AQ
SCHERING

EQ 0.042MG DIPROP/SPRAY
EQ 0.084MG DIPROP/SPRAY

N19589 001 Dec 23, 1987
N20469 001 Jun 26, 1996

BENDROFLUMETHIAZIDE

TABLET; ORAL
NATURETIN-10
APOTHECON
NATURETIN-2.5
APOTHECON
NATURETIN-5
APOTHECON

10MG
2.5MG
5MG

N12164 003
N12164 001
N12164 002

BENOXINATE HYDROCHLORIDE

SOLUTION/DROPS; OPHTHALMIC
BENOXINATE HYDROCHLORIDE
SOLA BARNES HIND

0.4%

N84149 001

BENTIROMIDE

SOLUTION; ORAL
CHYMEX
SAVAGE LABS

500MG/7.5ML

N18366 001 Dec 29, 1983

BENZPHETAMINE HYDROCHLORIDE

TABLET; ORAL
DIDREX

PHARMACIA AND UPJOHN 25MG

N12427 003

BENZQUINAMIDE HYDROCHLORIDE

INJECTABLE; INJECTION
EMETE-CON

PFIZER

EQ 50MG BASE/VIAL

N16820 001

SUPPOSITORY; RECTAL
EMETE-CON

ROERIG

EQ 100MG BASE

N16818 006

BENZTHIAZIDE

TABLET; ORAL
AQUATAG

SOLVAY

25MG
50MG

N16001 001
N16001 002

BENZTHIAZIDE

LEINER

50MG

N83206 001

EXNA

AH ROBINS INC

50MG

N12489 001

FOVANE

PFIZER

50MG

N12128 002

URESE

PFIZER

25MG

N12128 003

BENZTROPINE MESYLATE

TABLET; ORAL
BENZTROPINE MESYLATE

QUANTUM PHARMICS

0.5MG
1MG
2MG
0.5MG
1MG
2MG

N88514 001 Jan 31, 1984
N88510 001 Jan 31, 1984
N88511 001 Jan 31, 1984
N89211 001 Jun 14, 1988
N89212 001 Jun 14, 1988
N89213 001 Jun 14, 1988

USL PHARMA

COGENTIN

MERCK

0.5MG

N09193 004

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 34 (of 274)

BENZTROPINE MESYLATE

TABLET; ORAL

COGENTIN

MERCK

1MG

N09193 003

2MG

N09193 002

BENZYL BENZOATE

EMULSION; TOPICAL

BENZYL BENZOATE

LANNETT

50%

N84535 001

BENZYL PENICILLOYL-POLYLYSINE

INJECTABLE; INJECTION

PRE-PEN

ALLERQUEST

60UMOLAR

N50114 001

BEPRIDIL HYDROCHLORIDE

TABLET; ORAL

BEPADIN

MEDPOINTE PHARM HLC

200MG

N19001 001

Dec 28, 1990

300MG

N19001 002

Dec 28, 1990

400MG

N19001 003

Dec 28, 1990

VASCOR

JOHNSON AND JOHNSON

200MG

N19002 001

Dec 28, 1990

300MG

N19002 002

Dec 28, 1990

400MG

N19002 003

Dec 28, 1990

BETA-CAROTENE

CAPSULE; ORAL

SOLATENE

ROCHE

30MG

N17589 001

BETAMETHASONE

CREAM; TOPICAL

CELESTONE

SCHERING

0.2%

N14762 001

TABLET; ORAL

CELESTONE

SCHERING

0.6MG

N12657 003

BETAMETHASONE BENZOATE

CREAM; TOPICAL

UTICORT

PARKE DAVIS

0.025%

N16998 002

GEL; TOPICAL

UTICORT

PARKE DAVIS

0.025%

N17244 001

LOTION; TOPICAL

UTICORT

PARKE DAVIS

0.025%

N17528 001

OINTMENT; TOPICAL

UTICORT

PARKE DAVIS

0.025%

N18089 001

BETAMETHASONE DIPROPIONATE

CREAM; TOPICAL

ALPHATREX

SAVAGE LABS

EQ 0.05% BASE

N19138 001

Jun 26, 1984

BETAMETHASONE DIPROPIONATE

PERRIGO NEW YORK

EQ 0.05% BASE

N72536 001

Jan 31, 1990

EQ 0.05% BASE

N74579 001

Nov 26, 1997

PHARMADERM

EQ 0.05% BASE

N19136 001

Jun 26, 1984

DIPROSONE

SCHERING

EQ 0.05% BASE

N17536 001

DISCONTINUED DRUG PRODUCT LIST

6 - 35 (of 274)

BETAMETHASONE DIPROPIONATE

CREAM, AUGMENTED; TOPICAL				
DIPROLENE				
SCHERING	EQ 0.05% BASE	N19408	001	Jan 31, 1986
DISC; TOPICAL				
DIPROSONE				
SCHERING	EQ 0.1% BASE	N17829	001	
GEL, AUGMENTED; TOPICAL				
DIPROLENE				
SCHERING	EQ 0.05% BASE	N19408	002	Nov 22, 1991
LOTION; TOPICAL				
ALPHATREX				
SAVAGE LABS	EQ 0.05% BASE	N70273	001	Aug 12, 1985
BETAMETHASONE DIPROPIONATE				
ALPHARMA US PHARMS	EQ 0.05% BASE	N71085	001	Feb 03, 1987
PHARMADERM	EQ 0.05% BASE	N70274	001	Aug 12, 1985
DIPROSONE				
SCHERING	EQ 0.05% BASE	N17781	001	
OINTMENT; TOPICAL				
ALPHATREX				
SAVAGE LABS	EQ 0.05% BASE	N19143	001	Sep 04, 1984
BETAMETHASONE DIPROPIONATE				
PERRIGO NEW YORK	EQ 0.05% BASE	N72526	001	Jan 31, 1990
PHARMADERM	EQ 0.05% BASE	N19140	001	Sep 04, 1984
DIPROSONE				
SCHERING	EQ 0.05% BASE	N17691	001	

BETAMETHASONE SODIUM PHOSPHATE

INJECTABLE; INJECTION				
BETAMETHASONE SODIUM PHOSPHATE				
WATSON LABS	EQ 3MG BASE/ML	N85738	001	
CELESTONE				
SCHERING	EQ 3MG BASE/ML **Federal Register determination that product was not discontinued or withdrawn for safety or efficacy reasons**	N17561	001	

BETAMETHASONE VALERATE

CREAM; TOPICAL				
BETADERM				
ROACO	EQ 0.1% BASE	N18839	001	Jun 30, 1983
BETAMETHASONE VALERATE				
PERRIGO NEW YORK	EQ 0.1% BASE	N70053	001	Jun 10, 1986
PHARMADERM	EQ 0.1% BASE	N18860	002	Aug 31, 1983
PHARMAFAIR	EQ 0.1% BASE	N70485	001	May 29, 1987
BETATREX				
SAVAGE LABS	EQ 0.1% BASE	N18862	001	Aug 31, 1983
VALISONE				
SCHERING	EQ 0.01% BASE	N16322	002	
	EQ 0.1% BASE	N16322	001	
LOTION; TOPICAL				
BETAMETHASONE VALERATE				
PHARMADERM	EQ 0.1% BASE	N18870	001	Aug 31, 1983
PHARMAFAIR	EQ 0.1% BASE	N70484	001	May 29, 1987
BETATREX				
SAVAGE LABS	EQ 0.1% BASE	N18867	001	Aug 31, 1983
VALISONE				
SCHERING	EQ 0.1% BASE	N16932	001	
OINTMENT; TOPICAL				
BETAMETHASONE VALERATE				
PERRIGO NEW YORK	EQ 0.1% BASE	N71478	001	Dec 23, 1987
PHARMADERM	EQ 0.1% BASE	N18864	001	Aug 31, 1983
PHARMAFAIR	EQ 0.1% BASE	N70486	001	May 29, 1987
BETATREX				
SAVAGE LABS	EQ 0.1% BASE	N18863	001	Aug 31, 1983

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 36 (of 274)

BETAMETHASONE VALERATEOINTMENT; TOPICAL
VALISONE

SCHERING

EQ 0.1% BASE

N16740 001

BETAXOLOL HYDROCHLORIDE; CHLORTHALIDONETABLET; ORAL
KERLEDEX

SANOFI AVENTIS US

5MG;12.5MG

N19807 001

Oct 30, 1992

10MG;12.5MG

N19807 002

Oct 30, 1992

BETAXOLOL HYDROCHLORIDE; PILOCARPINE HYDROCHLORIDE

SUSPENSION/DROPS; OPHTHALMIC

BETOPTIC PILO

ALCON

EQ 0.25% BASE;1.75%

N20619 001

Apr 17, 1997

BETAZOLE HYDROCHLORIDEINJECTABLE; INJECTION
HISTALOG

LILLY

50MG/ML

N09344 001

BETHANECHOL CHLORIDEINJECTABLE; INJECTION
URECHOLINE

ODYSSEY PHARMS

5MG/ML **Federal Register
determination that product was not
discontinued or withdrawn for safety
or efficacy reasons**

N06536 001

TABLET; ORAL

BETHANECHOL CHLORIDE

ABLE

5MG

N40492 001

Jul 27, 2004

10MG

N40483 001

Jul 27, 2004

25MG

N40485 001

Jul 27, 2004

50MG

N40509 001

Jul 27, 2004

ASCOT

10MG

N88288 001

Jun 08, 1983

25MG

N88289 001

Jun 08, 1983

IVAX PHARMS

25MG

N84689 001

LANNETT

5MG

N84702 001

10MG

N84712 001

25MG

N84074 001

SANDOZ

5MG

N84353 001

10MG

N84378 001

10MG

N84379 001

25MG

N84383 001

25MG

N84384 001

WATSON LABS

5MG

N84402 001

5MG

N85230 002

5MG

N85841 001

10MG

N84408 001

10MG

N85228 001

10MG

N85842 001

25MG

N84441 001

25MG

N85229 001

25MG

N85839 001

50MG

N87397 001

50MG

N87444 001

MYOTONACHOL

GLENWOOD

5MG

N84188 001

10MG

N84188 003

25MG

N84188 004

URECHOLINE

ODYSSEY PHARMS

5MG

N06536 003

10MG

N06536 002

25MG

N06536 004

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 37 (of 274)

BETHANECHOL CHLORIDETABLET; ORAL
URECHOLINE

ODYSSEY PHARMS

50MG

N06536 005

BETHANIDINE SULFATETABLET; ORAL
TENATHAN

ROBINS AH

10MG

N17675 001

25MG

N17675 002

BIPERIDEN LACTATEINJECTABLE; INJECTION
AKINETON

ABBOTT

5MG/ML

N12418 002

BISOPROLOL FUMARATE; HYDROCHLOROTHIAZIDE

TABLET; ORAL

BISOPROLOL FUMARATE AND HYDROCHLOROTHIAZIDE

APOTHECON

2.5MG;6.25MG

N75642 002

Dec 27, 2000

5MG;6.25MG

N75642 001

Dec 27, 2000

10MG;6.25MG

N75642 003

Dec 27, 2000

IVAX PHARMS

2.5MG;6.25MG

N75632 001

Sep 27, 2000

5MG;6.25MG

N75632 002

Sep 27, 2000

10MG;6.25MG

N75632 003

Sep 27, 2000

BITOLTEROL MESYLATE

AEROSOL, METERED; INHALATION

TORNALATE

SANOFI AVENTIS US

0.37MG/INH

N18770 001

Dec 28, 1984

SOLUTION; INHALATION

TORNALATE

SANOFI AVENTIS US

0.2%

N19548 001

Feb 19, 1992

BLEOMYCIN SULFATE

INJECTABLE; INJECTION

BLEOMYCIN SULFATE

SICOR PHARMS

EQ 15 UNITS BASE/VIAL

N64084 001

Jun 01, 1996

EQ 30 UNITS BASE/VIAL

N64084 002

Jun 01, 1996

BRETYLIUM TOSYLATE

INJECTABLE; INJECTION

BRETYLIUM TOSYLATE

ABRAXIS PHARM

50MG/ML

N70134 001

Apr 29, 1986

100MG/ML

N71298 001

Feb 13, 1987

ASTRAZENECA

50MG/ML

N71151 001

Aug 10, 1987

50MG/ML

N71152 001

Aug 10, 1987

50MG/ML

N71153 001

Aug 10, 1987

BAXTER HLTHCARE

50MG/ML

N70545 001

May 14, 1986

50MG/ML

N70546 001

May 14, 1986

HOSPIRA

50MG/ML

N19033 001

Apr 29, 1986

BRETYLIUM TOSYLATE IN DEXTROSE 5%

ABBOTT

200MG/100ML

N19005 002

Apr 29, 1986

400MG/100ML

N19005 003

Apr 29, 1986

800MG/100ML

N19005 001

Apr 29, 1986

BRETYLIUM TOSYLATE IN DEXTROSE 5% IN PLASTIC CONTAINER

BAXTER HLTHCARE

200MG/100ML

N19837 002

Apr 12, 1989

400MG/100ML

N19837 001

Apr 12, 1989

HOSPIRA

800MG/100ML

N19008 001

Apr 29, 1986

BRETYLOL

MAYNE PHARMA USA

50MG/ML

N17954 001

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 38 (of 274)

BRIMONIDINE TARTRATE

SOLUTION/DROPS; OPHTHALMIC

ALPHAGAN

ALLERGAN

0.2%

N20613 001

Sep 06, 1996

0.5%

N20490 001

Mar 13, 1997

BROMOCRIPTINE MESYLATE

CAPSULE; ORAL

BROMOCRIPTINE MESYLATE

LEK PHARM

EQ 5MG BASE

N75100 001

Dec 10, 1998

BROMODIPHENHYDRAMINE HYDROCHLORIDE

CAPSULE; ORAL

AMBODRYL

PARKE DAVIS

25MG

N07984 001

BROMODIPHENHYDRAMINE HYDROCHLORIDE; CODEINE PHOSPHATE

SYRUP; ORAL

AMBENYL

FOREST LABS

12.5MG/5ML;10MG/5ML

N09319 006

Jan 10, 1984

BROMANYL

ALPHARMA US PHARMS

12.5MG/5ML;10MG/5ML

N88343 001

Aug 15, 1984

BROMPHENIRAMINE MALEATE

ELIXIR; ORAL

BROMPHENIRAMINE MALEATE

ALPHARMA US PHARMS

2MG/5ML

N86936 001

PHARM ASSOC

2MG/5ML

N87517 001

USL PHARMA

2MG/5ML

N87964 001

Jan 25, 1983

INJECTABLE; INJECTION

BROMPHENIRAMINE MALEATE

WATSON LABS

100MG/ML

N83820 001

DIMETANE-TEN

WYETH AYERST

10MG/ML

N11418 002

TABLET; ORAL

BROMPHENIRAMINE MALEATE

ANABOLIC

4MG

N86187 001

BARR

4MG

N84468 001

IVAX PHARMS

4MG

N84351 001

NEWTRON PHARMS

4MG

N86987 001

PAR PHARM

4MG

N87009 001

PIONEER PHARMS

4MG

N88604 001

Jul 13, 1984

SANDOZ

4MG

N83215 001

VITARINE

4MG

N85850 001

WATSON LABS

4MG

N83123 001

N85769 001

DIMETANE

WYETH CONS

4MG

N10799 003

TABLET, EXTENDED RELEASE; ORAL

DIMETANE

WYETH CONS

8MG

N10799 010

Jun 10, 1983

12MG

N10799 011

Jun 10, 1983

BROMPHENIRAMINE MALEATE; CODEINE PHOSPHATE; PHENYLPROPANOLAMINE HYDROCHLORIDE

SYRUP; ORAL

BROMANATE DC

ALPHARMA US PHARMS

2MG/5ML;10MG/5ML;12.5MG/5ML

N88723 001

Feb 25, 1985

DIMETANE-DC

ROBINS AH

2MG/5ML;10MG/5ML;12.5MG/5ML

N11694 006

Mar 29, 1984

MYPHETANE DC

MORTON GROVE

2MG/5ML;10MG/5ML;12.5MG/5ML

N88904 001

Feb 21, 1985

DISCONTINUED DRUG PRODUCT LIST

6 - 39 (of 274)

BROMPHENIRAMINE MALEATE; DEXTROMETHORPHAN HYDROBROMIDE; PSEUDOEPHEDRINE HYDROCHLORIDE

SYRUP; ORAL		
BROMANATE DM		
ALPHARMA US PHARMS	2MG/5ML;10MG/5ML;30MG/5ML	N88722 001 Mar 07, 1985
DIMETANE-DX		
ROBINS AH	2MG/5ML;10MG/5ML;30MG/5ML	N19279 001 Aug 24, 1984

BROMPHENIRAMINE MALEATE; PHENYLPROPANOLAMINE HYDROCHLORIDE

ELIXIR; ORAL		
BIPHETAP		
MORTON GROVE	4MG/5ML;25MG/5ML	N88687 001 Sep 26, 1984
BROMANATE		
ALPHARMA US PHARMS	4MG/5ML;25MG/5ML	N88688 001 Feb 06, 1985
DIMETAPP		
WYETH CONS	2MG/5ML;12.5MG/5ML	N13087 003 Mar 29, 1984
TABLET, EXTENDED RELEASE; ORAL		
BROMATAPP		
COPLEY PHARM	12MG;75MG	N71099 001 Jul 02, 1987
DIMETAPP		
WYETH CONS	12MG;75MG	N12436 003 May 14, 1985

BUCLIZINE HYDROCHLORIDE

TABLET; ORAL		
BUCLADIN-S		
STUART PHARMS	50MG	N10911 006

BUDESONIDE

AEROSOL, METERED; NASAL		
RHINOCORT		
ASTRAZENECA	0.032MG/INH	N20233 001 Feb 14, 1994
POWDER, METERED; INHALATION		
PULMICORT		
ASTRAZENECA	0.32MG/INH	N20441 003 Jun 24, 1997
SPRAY, METERED; NASAL		
RHINOCORT		
ASTRAZENECA	0.064MG/INH	N20746 002 Oct 01, 1999
SUSPENSION; INHALATION		
PULMICORT RESPULES		
ASTRAZENECA	1MG/2ML	N20929 003 Aug 08, 2000

BUMETANIDE

INJECTABLE; INJECTION		
BUMEX		
ROCHE	0.25MG/ML	N18226 001 Feb 28, 1983

BUPIVACAINE HYDROCHLORIDE

INJECTABLE; INJECTION		
BUPIVACAINE HYDROCHLORIDE KIT		
HOSPIRA	0.075%	N19978 001 Sep 03, 1992
	0.114%	N19978 002 Sep 03, 1992
	0.23%	N19978 003 Sep 03, 1992

BUPROPION HYDROCHLORIDE

TABLET; ORAL		
WELLBUTRIN		
GLAXOSMITHKLINE	50MG	N18644 001 Dec 30, 1985
TABLET, EXTENDED RELEASE; ORAL		
ZYBAN		
GLAXOSMITHKLINE	100MG	N20711 002 May 14, 1997

BUSPIRONE HYDROCHLORIDE

CAPSULE; ORAL		
BUSPAR		
BRISTOL MYERS SQUIBB	5MG	N21190 001 Dec 20, 2000

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 40 (of 274)

BUSPIRONE HYDROCHLORIDE

CAPSULE; ORAL

BUSPAR

BRISTOL MYERS SQUIBB 7.5MG

N21190 002 Dec 20, 2000

10MG

N21190 003 Dec 20, 2000

15MG

N21190 004 Dec 20, 2000

BUTABARBITAL SODIUM

CAPSULE; ORAL

BUTICAPS

MEDPOINTE PHARM HLC 15MG

N85381 001

30MG

N85381 002

50MG

N85381 003

100MG

N85381 004

ELIXIR; ORAL

BUTABARB

ALPHARMA US PHARMS 30MG/5ML

N85873 001

BUTABARBITAL SODIUM

MORTON GROVE 30MG/5ML

N85383 001

BUTALAN

LANNETT 33.3MG/5ML

N85880 001

SARISOL

HALSEY 30MG/5ML

N84723 001

TABLET; ORAL

BUTABARBITAL SODIUM

SANDOZ 15MG

N84292 003 Feb 09, 1982

15MG

N85938 001

30MG

N84272 002

30MG

N85934 001

SOLVAY 16.2MG

N83606 001

32.4MG

N83898 001

48.6MG

N83897 001

97.2MG

N83896 001

TEVA

15MG

N88632 001

May 18, 1985

30MG

N88631 001

May 01, 1985

WATSON LABS

15MG

N85764 001

30MG

N85772 001

WHITEWORTH TOWN PLSN

15MG

N83325 002

30MG

N83337 001

BUTISOL SODIUM

MEDPOINTE PHARM HLC 15MG

N00793 002

100MG

N00793 005

SARISOL NO. 1

HALSEY 15MG

N84719 001

SARISOL NO. 2

HALSEY 30MG

N84719 002

SODIUM BUTABARBITAL

IVAX PHARMS 15MG

N83484 001

30MG

N84040 001

LANNETT

15MG

N85849 001

30MG

N85866 001

100MG

N85881 001

WEST WARD

15MG

N85418 001

30MG

N85432 001

BUTOCONAZOLE NITRATE

CREAM; VAGINAL

FEMSTAT

ROCHE PALO 2%

N19215 001 Nov 25, 1985

SUPPOSITORY; VAGINAL

FEMSTAT

ROCHE PALO 100MG

N19359 001 Nov 25, 1985

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 41 (of 274)

BUTORPHANOL TARTRATESPRAY, METERED; NASAL
STADOL

BRISTOL MYERS SQUIBB 1MG/SPRAY

N19890 001 Dec 12, 1991

CAFFEINE; ERGOTAMINE TARTRATE

SUPPOSITORY; RECTAL

CAFERGOT

NOVARTIS 100MG;2MG

N09000 002

TABLET; ORAL

CAFERGOT

NOVARTIS 100MG;1MG

N06620 001

WIGRAINE

ORGANON USA INC 100MG;1MG

N86562 001

CALCIFEDIOL

CAPSULE; ORAL

CALDEROL

ORGANON USA INC 0.02MG

N18312 001

0.05MG

N18312 002

CALCITONIN HUMAN

INJECTABLE; INJECTION

CIBACALCIN

NOVARTIS 0.5MG/VIAL

N18470 001 Oct 31, 1986

CALCITONIN, SALMON

INJECTABLE; INJECTION

CALCIMAR

SANOFI AVENTIS US 200 IU/ML
400 IU/VIAL

N17769 001

N17497 001

CALCITONIN-SALMON

ASTRAZENECA 200 IU/ML

N73690 001 Apr 14, 1995

MIACALCIN

NOVARTIS 100 IU/ML

N17808 001 Jul 03, 1986

CALCITRIOL

INJECTABLE; INJECTION

CALCITRIOL

MAYNE PHARMA USA 0.001MG/ML

N75816 001 Jan 16, 2004

0.002MG/ML

N75816 002 Jan 16, 2004

CALCIUM ACETATE

CAPSULE; ORAL

PHOSLO

FRESENIUS MEDCL EQ 84.5MG CALCIUM
EQ 169MG CALCIUM

N21160 001 Apr 02, 2001

N21160 002 Apr 02, 2001

TABLET; ORAL

PHOSLO

FRESENIUS MEDCL EQ 169MG CALCIUM

N19976 001 Dec 10, 1990

CALCIUM CHLORIDE; DEXTROSE; MAGNESIUM CHLORIDE; POTASSIUM CHLORIDE; SODIUM ACETATE; SODIUM CHLORIDE

INJECTABLE; INJECTION

ISOLYTE R W/ DEXTROSE 5% IN PLASTIC CONTAINER

B BRAUN 37MG/100ML;5GM/100ML;31MG/100ML;120MG/100ML;330MG/100ML;88MG/100ML N18271 001

CALCIUM CHLORIDE; DEXTROSE; MAGNESIUM CHLORIDE; POTASSIUM CHLORIDE; SODIUM ACETATE; SODIUM CHLORIDE; SODIUM CITRATE

INJECTABLE; INJECTION

ISOLYTE E W/ DEXTROSE 5% IN PLASTIC CONTAINER

B BRAUN 35MG/100ML;5GM/100ML;30MG/100ML;74MG/100ML;640MG/100ML;500MG/100ML;74MG/100ML N18269 002 Jan 17, 1983

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 42 (of 274)

CALCIUM CHLORIDE; DEXTROSE; MAGNESIUM CHLORIDE; SODIUM ACETATE; SODIUM CHLORIDE

SOLUTION; INTRAPERITONEAL

DIALYTE LM/ DEXTROSE 2.5% IN PLASTIC CONTAINER

B BRAUN	29MG/100ML;2.5GM/100ML;15MG/100ML;610MG /100ML;560MG/100ML	N18460	006	Jan 29, 1986
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DIALYTE W/ DEXTROSE 1.5% IN PLASTIC CONTAINER

B BRAUN	29MG/100ML;1.5GM/100ML;15MG/100ML;610MG /100ML;560MG/100ML	N18460	001	
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DIALYTE W/ DEXTROSE 4.25% IN PLASTIC CONTAINER

B BRAUN	29MG/100ML;4.25GM/100ML;15MG/100ML;610M G/100ML;560MG/100ML	N18460	003	
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CALCIUM CHLORIDE; DEXTROSE; MAGNESIUM CHLORIDE; SODIUM CHLORIDE; SODIUM LACTATE

SOLUTION; INTRAPERITONEAL

DIALYTE LM/ DEXTROSE 1.5% IN PLASTIC CONTAINER

B BRAUN	26MG/100ML;1.5GM/100ML;15MG/100ML;560MG /100ML;390MG/100ML	N18460	002	
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DIALYTE LM/ DEXTROSE 2.5% IN PLASTIC CONTAINER

B BRAUN	26MG/100ML;5GM/100ML;5MG/100ML;530MG/10 0ML;450MG/100ML	N18460	008	Jan 29, 1986
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DIALYTE LM/ DEXTROSE 4.25% IN PLASTIC CONTAINER

B BRAUN	26MG/100ML;4.25GM/100ML;15MG/100ML;560M G/100ML;390MG/100ML	N18460	004	
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CALCIUM CHLORIDE; DEXTROSE; POTASSIUM CHLORIDE; SODIUM CHLORIDE

INJECTABLE; INJECTION

DEXTROSE 5% IN RINGER'S IN PLASTIC CONTAINER

B BRAUN	33MG/100ML;5GM/100ML;30MG/100ML;860MG/1 00ML	N18256	001	
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CALCIUM CHLORIDE; DEXTROSE; POTASSIUM CHLORIDE; SODIUM CHLORIDE; SODIUM LACTATE

INJECTABLE; INJECTION

DEXTROSE 4% IN MODIFIED LACTATED RINGER'S IN PLASTIC CONTAINER

B BRAUN	4MG/100ML;4GM/100ML;6MG/100ML;120MG/100 ML;62MG/100ML	N19634	002	Feb 24, 1988
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DEXTROSE 5% IN LACTATED RINGER'S IN PLASTIC CONTAINER

B BRAUN	20MG/100ML;5GM/100ML;30MG/100ML;600MG/1 00ML;310MG/100ML	N17510	001	
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MILES	20MG/100ML;5GM/100ML;30MG/100ML;600MG/1 00ML;310MG/100ML	N18499	001	
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POTASSIUM CHLORIDE 10MEQ IN DEXTROSE 5% AND LACTATED RINGER'S IN PLASTIC CONTAINER

HOSPIRA	20MG/100ML;5GM/100ML;104MG/100ML;600MG/ 100ML;310MG/100ML	N19685	005	Oct 17, 1988
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	20MG/100ML;5GM/100ML;179MG/100ML;600MG/ 100ML;310MG/100ML	N19685	006	Oct 17, 1988
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POTASSIUM CHLORIDE 15MEQ IN DEXTROSE 5% AND LACTATED RINGER'S IN PLASTIC CONTAINER

HOSPIRA	20MG/100ML;5GM/100ML;254MG/100ML;600MG/ 100ML;310MG/100ML	N19685	007	Oct 17, 1988
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POTASSIUM CHLORIDE 30MEQ IN DEXTROSE 5% AND LACTATED RINGER'S IN PLASTIC CONTAINER

HOSPIRA	20MG/100ML;5GM/100ML;254MG/100ML;600MG/ 100ML;310MG/100ML	N19685	003	Oct 17, 1988
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POTASSIUM CHLORIDE 5MEQ IN DEXTROSE 5% AND LACTATED RINGER'S IN PLASTIC CONTAINER

HOSPIRA	20MG/100ML;5GM/100ML;104MG/100ML;600MG/ 100ML;310MG/100ML	N19685	001	Oct 17, 1988
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CALCIUM CHLORIDE; DEXTROSE; SODIUM CHLORIDE; SODIUM LACTATE

SOLUTION; INTRAPERITONEAL

INPERSOL-ZM W/ DEXTROSE 1.5% IN PLASTIC CONTAINER

FRESENIUS MEDCL	25.7MG/100ML;1.5GM/100ML;538MG/100ML;44 8MG/100ML	N19395	001	Mar 26, 1986
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INPERSOL-ZM W/ DEXTROSE 2.5% IN PLASTIC CONTAINER

FRESENIUS MEDCL	25.7MG/100ML;2.5GM/100ML;538MG/100ML;44 8MG/100ML	N19395	002	Mar 26, 1986
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INPERSOL-ZM W/ DEXTROSE 4.25% IN PLASTIC CONTAINER

FRESENIUS MEDCL	25.7MG/100ML;4.25GM/100ML;538MG/100ML;4 48MG/100ML	N19395	003	Mar 26, 1986
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27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 43 (of 274)

CALCIUM CHLORIDE; MAGNESIUM CHLORIDE; POTASSIUM CHLORIDE; SODIUM ACETATE; SODIUM CHLORIDE

INJECTABLE; INJECTION

TPN ELECTROLYTES IN PLASTIC CONTAINER

ABBOTT	16.5MG/ML;25.4MG/ML;74.6MG/ML;121MG/ML; 16.1MG/ML	N19399	001	Jun 16, 1986
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CALCIUM CHLORIDE; MAGNESIUM CHLORIDE; POTASSIUM CHLORIDE; SODIUM ACETATE; SODIUM CHLORIDE; SODIUM CITRATE

INJECTABLE; INJECTION

ISOLYTE E IN PLASTIC CONTAINER

B BRAUN	35MG/100ML;30MG/100ML;74MG/100ML;640MG/ 100ML;500MG/100ML;74MG/100ML	N18899	001	Oct 31, 1983
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CALCIUM CHLORIDE; POTASSIUM CHLORIDE; SODIUM ACETATE; SODIUM CHLORIDE

INJECTABLE; INJECTION

ACETATED RINGER'S IN PLASTIC CONTAINER

B BRAUN	20MG/100ML;30MG/100ML;380MG/100ML;600MG/ /100ML	N18725	001	Nov 29, 1982
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CALCIUM CHLORIDE; POTASSIUM CHLORIDE; SODIUM CHLORIDE

INJECTABLE; INJECTION

RINGER'S IN PLASTIC CONTAINER

B BRAUN	33MG/100ML;30MG/100ML;860MG/100ML	N18721	001	Nov 09, 1982
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SOLUTION; IRRIGATION

RINGER'S IN PLASTIC CONTAINER

ABBOTT	33MG/100ML;30MG/100ML;860MG/100ML	N18462	001	
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CALCIUM CHLORIDE; POTASSIUM CHLORIDE; SODIUM CHLORIDE; SODIUM LACTATE

INJECTABLE; INJECTION

LACTATED RINGER'S IN PLASTIC CONTAINER

ABBOTT	20MG/100ML;30MG/100ML;600MG/100ML;310MG/ /100ML	N19485	001	Oct 24, 1985
B BRAUN	20MG/100ML;30MG/100ML;600MG/100ML;310MG/ /100ML	N18023	001	
MILES	20MG/100ML;30MG/100ML;600MG/100ML;310MG/ /100ML	N18417	001	

CALCIUM GLUCEPTATE

INJECTABLE; INJECTION

CALCIUM GLUCEPTATE

ABBOTT	EQ 90MG CALCIUM/5ML	N80001	001	
	EQ 90MG CALCIUM/5ML	N83159	001	
ABRAXIS PHARM	EQ 90MG CALCIUM/5ML	N89373	001	Apr 30, 1987
LILLY	EQ 90MG CALCIUM/5ML	N06470	001	

CALCIUM METRIZOATE; MEGLUMINE METRIZOATE; METRIZOATE MAGNESIUM; METRIZOATE SODIUM

INJECTABLE; INJECTION

ISOPAQUE 440

GE HEALTHCARE	0.78MG/ML;75.9MG/ML;0.15MG/ML;16.6MG/ML	N16847	001	
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CALCIUM; MEGLUMINE; METRIZOIC ACID

INJECTABLE; INJECTION

ISOPAQUE 280

GE HEALTHCARE	0.35MG/ML;140.1MG/ML;461.8MG/ML	N17506	001	
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CANDICIDIN

OINTMENT; VAGINAL

VANOBIID

SANOFI AVENTIS US	0.6MG/GM	N61596	001	
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TABLET; VAGINAL

VANOBIID

SANOFI AVENTIS US	3MG	N61613	001	
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DISCONTINUED DRUG PRODUCT LIST

6 - 44 (of 274)

CAPTOPRIL

TABLET; ORAL
CAPOTEN

PAR PHARM	37.5MG	N18343	006	Sep 17, 1986
	75MG	N18343	007	Jun 13, 1995
	150MG	N18343	004	Jun 13, 1995

CAPTOPRIL

APOTHECON	12.5MG	N74472	001	Mar 31, 1995
	25MG	N74472	002	Mar 31, 1995
	50MG	N74472	003	Mar 31, 1995
	100MG	N74472	004	Mar 31, 1995

CLONMEL HLTHCARE	12.5MG	N74423	001	Feb 13, 1996
	25MG	N74423	002	Feb 13, 1996
	50MG	N74423	003	Feb 13, 1996
	100MG	N74423	004	Feb 13, 1996

ENDO LABS	12.5MG	N74418	001	Feb 13, 1996
	25MG	N74418	002	Feb 13, 1996
	50MG	N74418	003	Feb 13, 1996
	100MG	N74418	004	Feb 13, 1996

PAR PHARM	12.5MG	N74493	001	Feb 13, 1996
	25MG	N74493	002	Feb 13, 1996
	50MG	N74493	003	Feb 13, 1996
	100MG	N74493	004	Feb 13, 1996

PUREPAC PHARM	12.5MG	N74640	001	Mar 31, 1997
	25MG	N74640	002	Mar 31, 1997
	50MG	N74640	003	Mar 31, 1997
	100MG	N74640	004	Mar 31, 1997

CAPTOPRIL; HYDROCHLOROTHIAZIDE

TABLET; ORAL

CAPTOPRIL AND HYDROCHLOROTHIAZIDE

ENDO LABS	25MG;15MG	N74788	001	Dec 29, 1997
	25MG;25MG	N74788	002	Dec 29, 1997
	50MG;15MG	N74788	004	Dec 29, 1997
	50MG;25MG	N74788	003	Dec 29, 1997
WATSON LABS	50MG;25MG	N74832	001	Dec 29, 1997

CARBACHOL

SOLUTION; INTRAOCULAR

CARBACHOL

PHARMAFAIR	0.01%	N70292	001	May 21, 1986
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CARBAMAZEPINE

SUSPENSION; ORAL

CARBAMAZEPINE

TARO	100MG/5ML	N75875	001	Dec 21, 2000
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TABLET; ORAL

CARBAMAZEPINE

USL PHARMA	200MG	N70300	001	May 15, 1986
WARNER CHILCOTT	200MG	N70429	001	Jan 02, 1987

CARBENICILLIN DISODIUM

INJECTABLE; INJECTION

GEOPEN

ROERIG	EQ 1GM BASE/VIAL	N50306	001	
	EQ 2GM BASE/VIAL	N50306	004	
	EQ 5GM BASE/VIAL	N50306	002	
	EQ 10GM BASE/VIAL	N50306	006	
	EQ 30GM BASE/VIAL	N50306	007	

PYOPEN

GLAXOSMITHKLINE	EQ 1GM BASE/VIAL	N50298	001	
	EQ 2GM BASE/VIAL	N50298	002	
	EQ 5GM BASE/VIAL	N50298	003	
	EQ 10GM BASE/VIAL	N50298	006	

DISCONTINUED DRUG PRODUCT LIST

6 - 45 (of 274)

CARBENICILLIN DISODIUM

INJECTABLE; INJECTION
PYOPEN

GLAXOSMITHKLINE	EQ 20GM BASE/VIAL	N50298	007
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CARBIDOPA; LEVODOPA

TABLET; ORAL
CARBIDOPA AND LEVODOPA
SCS

	10MG;100MG	N74080	001	Mar 25, 1994
	25MG;100MG	N74080	002	Mar 25, 1994
	25MG;250MG	N74080	003	Mar 25, 1994
WATSON LABS	10MG;100MG	N73381	001	Sep 28, 1993
	25MG;100MG	N73382	001	Sep 28, 1993
	25MG;250MG	N73383	001	Sep 28, 1993

CARBINOXAMINE MALEATE

ELIXIR; ORAL
CLISTIN
MCNEIL

	4MG/5ML **Federal Register determination that product was not discontinued or withdrawn for safety or efficacy reasons**	N08955	001
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TABLET; ORAL
CLISTIN

ORTHO MCNEIL PHARM	4MG	N08915	001
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CARBOPLATIN

INJECTABLE; INJECTION
CARBOPLATIN

MAYNE PHARMA USA	50MG/VIAL	N76473	001	Oct 27, 2004
	150MG/VIAL	N76473	002	Oct 27, 2004
	450MG/VIAL	N76473	003	Oct 27, 2004

INJECTABLE; IV (INFUSION)
CARBOPLATIN

ABRAXIS PHARM	EQ 50MG/5ML (10MG/ML)	N77247	001	Oct 21, 2004
	EQ 150MG/15ML (10MG/ML)	N77247	002	Oct 21, 2004

CARISOPRODOL

CAPSULE; ORAL
SOMA

MEDPOINTE PHARM HLC	250MG	N11792	003
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TABLET; ORAL
CARISOPRODOL

ABLE	350MG	N40421	001	Jun 21, 2001
PIIONEER PHARMS	350MG	N89390	001	Oct 13, 1988
SANDOZ	350MG	N89566	001	Aug 30, 1988
WATSON LABS	350MG	N85433	001	
RELA				
SCHERING	350MG	N12155	001	

CARPHENAZINE MALEATE

CONCENTRATE; ORAL
PROKETAZINE

WYETH AYERST	50MG/ML	N14173	001
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TABLET; ORAL
PROKETAZINE

WYETH AYERST	12.5MG	N12768	001
	25MG	N12768	002
	50MG	N12768	004

CARPROFEN

TABLET; ORAL
RIMADYL

ROCHE	100MG	N18550	002	Dec 31, 1987
	150MG	N18550	003	Dec 31, 1987

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 46 (of 274)

CARTEOLOL HYDROCHLORIDE

TABLET; ORAL

CARTROL

ABBOTT

2.5MG

N19204 001

Dec 28, 1988

5MG

N19204 002

Dec 28, 1988

10MG

N19204 003

Dec 28, 1988

CEFACLOR

CAPSULE; ORAL

CECLOR

LILLY

EQ 250MG BASE

N50521 001

EQ 500MG BASE

N50521 002

CEFACLOR

CLONMEL HLTHCARE

EQ 250MG BASE

N64107 001

Apr 27, 1995

EQ 500MG BASE

N64107 002

Apr 27, 1995

TEVA

EQ 250MG BASE

N64081 001

Sep 16, 1996

EQ 500MG BASE

N64081 002

Sep 16, 1996

FOR SUSPENSION; ORAL

CECLOR

LILLY

EQ 125MG BASE/5ML

N50522 001

EQ 250MG BASE/5ML

N50522 002

CEFACLOR

CLONMEL HLTHCARE

EQ 125MG BASE/5ML

N64114 001

Apr 28, 1995

EQ 187MG BASE/5ML

N64115 001

Apr 28, 1995

EQ 250MG BASE/5ML

N64116 001

Apr 28, 1995

EQ 375MG BASE/5ML

N64110 001

Apr 28, 1995

IVAX PHARMS

EQ 125MG BASE/5ML

N64087 001

Apr 28, 1995

EQ 187MG BASE/5ML

N64086 001

Apr 28, 1995

EQ 250MG BASE/5ML

N64085 001

Apr 28, 1995

TABLET, EXTENDED RELEASE; ORAL

CECLOR CD

LILLY

EQ 375MG BASE

N50673 001

Jun 28, 1996

EQ 500MG BASE

N50673 002

Jun 28, 1996

CEFADROXIL/CEFADROXIL HEMIHYDRATE

CAPSULE; ORAL

CEFADROXIL

PUREPAC PHARM

EQ 500MG BASE

N63017 001

Jan 05, 1989

TEVA

EQ 500MG BASE

N62695 001

Feb 10, 1989

DURICEF

WARNER CHILCOTT

EQ 250MG BASE

N50512 002

EQ 500MG BASE

N50512 001

ULTRACEF

BRISTOL

EQ 500MG BASE

N62378 001

Mar 16, 1982

FOR SUSPENSION; ORAL

CEFADROXIL

APOTHECON

EQ 125MG BASE/5ML

N62334 001

EQ 250MG BASE/5ML

N62334 002

EQ 500MG BASE/5ML

N62334 003

TEVA

EQ 125MG BASE/5ML

N62698 001

Mar 01, 1989

EQ 250MG BASE/5ML

N62698 002

Mar 01, 1989

EQ 500MG BASE/5ML

N62698 003

Mar 01, 1989

DURICEF

WARNER CHILCOTT

EQ 125MG BASE/5ML

N50527 002

ULTRACEF

BRISTOL

EQ 125MG BASE/5ML

N62376 001

Mar 16, 1982

EQ 250MG BASE/5ML

N62376 002

Mar 16, 1982

EQ 500MG BASE/5ML

N62376 003

Mar 16, 1982

TABLET; ORAL

DURICEF

WARNER CHILCOTT

EQ 1GM BASE

N50528 001

ULTRACEF

APOTHECON

EQ 1GM BASE

N62390 001

Jun 10, 1982

BRISTOL

EQ 1GM BASE

N62408 001

Aug 31, 1982

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 47 (of 274)

CEFAMANDOLE NAFATEINJECTABLE; INJECTION
MANDOL

LILLY

EQ 500MG BASE/VIAL	N50504	001	
EQ 1GM BASE/VIAL	N50504	002	
EQ 1GM BASE/VIAL	N62560	001	Sep 10, 1985
EQ 2GM BASE/VIAL	N50504	003	
EQ 2GM BASE/VIAL	N62560	002	Sep 10, 1985
EQ 10GM BASE/VIAL	N50504	004	

CEFAZOLIN SODIUMINJECTABLE; INJECTION
ANCEF

GLAXOSMITHKLINE

EQ 250MG BASE/VIAL	N50461	001	
EQ 500MG BASE/VIAL	N50461	002	
EQ 5GM BASE/VIAL	N50461	004	

ANCEF IN DEXTROSE 5% IN PLASTIC CONTAINER

BAXTER HLTHCARE

EQ 10MG BASE/ML	N50566	003	Jun 08, 1983
EQ 20MG BASE/ML	N50566	004	Jun 08, 1983

ANCEF IN SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER

BAXTER HLTHCARE

EQ 10MG BASE/ML	N50566	001	Jun 08, 1983
EQ 20MG BASE/ML	N50566	002	Jun 08, 1983

CEFAZOLIN AND DEXTROSE

B BRAUN

EQ 500MG BASE/VIAL	N50779	001	Jul 27, 2000
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CEFAZOLIN SODIUM

ABRAXIS PHARM

EQ 500MG BASE/VIAL	N62688	002	Nov 17, 1986
EQ 1GM BASE/VIAL	N62688	003	Nov 17, 1986
EQ 10GM BASE/VIAL	N62688	004	Nov 17, 1986
EQ 20GM BASE/VIAL	N62688	005	Aug 03, 1987

BEDFORD

EQ 250MG BASE/VIAL	N62894	001	Jul 21, 1988
EQ 500MG BASE/VIAL	N62894	002	Jul 21, 1988
EQ 1GM BASE/VIAL	N62894	003	Jul 21, 1988
EQ 5GM BASE/VIAL	N62894	004	Jul 21, 1988
EQ 10GM BASE/VIAL	N62894	005	Jul 21, 1988

ELKINS SINN

EQ 250MG BASE/VIAL	N62807	001	Jan 12, 1988
EQ 500MG BASE/VIAL	N62807	002	Jan 12, 1988
EQ 1GM BASE/VIAL	N62807	003	Jan 12, 1988
EQ 5GM BASE/VIAL	N62807	004	Jan 12, 1988
EQ 10GM BASE/VIAL	N62807	005	Jan 12, 1988
EQ 20GM BASE/VIAL	N62807	006	Jan 12, 1988

MARSAM PHARMS LLC

EQ 250MG BASE/VIAL	N62988	001	Dec 29, 1989
EQ 500MG BASE/VIAL	N62988	002	Dec 29, 1989
EQ 1GM BASE/VIAL	N62988	003	Dec 29, 1989
EQ 5GM BASE/VIAL	N62989	001	Dec 29, 1989
EQ 10GM BASE/VIAL	N62989	002	Dec 29, 1989
EQ 20GM BASE/VIAL	N62989	003	Dec 29, 1989

TEVA

EQ 250MG BASE/VIAL	N63016	001	Mar 14, 1989
EQ 500MG BASE/VIAL	N63016	002	Mar 14, 1989
EQ 1GM BASE/VIAL	N63016	003	Mar 14, 1989
EQ 5GM BASE/VIAL	N63018	001	Mar 05, 1990
EQ 10GM BASE/VIAL	N63018	002	Mar 05, 1990

KEFZOL

LILLY

EQ 250MG BASE/VIAL	N61773	001	
EQ 500MG BASE/VIAL	N61773	002	
EQ 500MG BASE/VIAL	N62557	001	Sep 10, 1985
EQ 1GM BASE/VIAL	N61773	003	
EQ 1GM BASE/VIAL	N62557	002	Sep 10, 1985
EQ 10GM BASE/VIAL	N61773	004	
EQ 20GM BASE/VIAL	N61773	005	Sep 08, 1987

CEFIXIMEFOR SUSPENSION; ORAL
SUPRAX

LEDERLE

100MG/5ML	N50622	001	Apr 28, 1989
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DISCONTINUED DRUG PRODUCT LIST

6 - 48 (of 274)

CEFIXIME

TABLET; ORAL
SUPRAX

LEDERLE	200MG **Federal Register determination that product was not discontinued or withdrawn for safety or efficacy reasons**	N50621	001	Apr 28, 1989
	400MG	N50621	002	Apr 28, 1989
LUPIN	400MG	N65130	001	Feb 12, 2004

CEFMENOXIME HYDROCHLORIDE

INJECTABLE; INJECTION
CEFMAX

TAP PHARM	EQ 500MG BASE/VIAL	N50571	001	Dec 30, 1987
	EQ 1GM BASE/VIAL	N50571	002	Dec 30, 1987
	EQ 2GM BASE/VIAL	N50571	003	Dec 30, 1987

CEFMETAZOLE SODIUM

INJECTABLE; INJECTION
ZEFAZONE

PHARMACIA AND UPJOHN	EQ 1GM BASE/VIAL	N50637	001	Dec 11, 1989
	EQ 2GM BASE/VIAL	N50637	002	Dec 11, 1989
ZEFAZONE IN PLASTIC CONTAINER				
PHARMACIA AND UPJOHN	EQ 20MG BASE/ML	N50683	001	Dec 29, 1992
	EQ 40MG BASE/ML	N50683	002	Dec 29, 1992

CEFONICID SODIUM

INJECTABLE; INJECTION
MONOCID

GLAXOSMITHKLINE	EQ 500MG BASE/VIAL	N50579	001	May 23, 1984
	EQ 1GM BASE/VIAL	N50579	002	May 23, 1984
	EQ 1GM BASE/VIAL	N63295	001	Jul 26, 1993
	EQ 2GM BASE/VIAL	N50579	003	May 23, 1984
	EQ 10GM BASE/VIAL	N50579	004	May 23, 1984

CEFOPERAZONE SODIUM

INJECTABLE; INJECTION
CEFOBID

PFIZER	EQ 1GM BASE/VIAL	N63333	001	Mar 31, 1995
	EQ 2GM BASE/VIAL	N63333	002	Mar 31, 1995

CEFORANIDE

INJECTABLE; INJECTION
PRECEF

APOTHECON	500MG/VIAL	N62579	001	Nov 26, 1984
	1GM/VIAL	N62579	002	Nov 26, 1984
	2GM/VIAL	N62579	003	Nov 26, 1984
	10GM/VIAL	N62579	004	Nov 26, 1984
	20GM/VIAL	N62579	005	Nov 26, 1984
BRISTOL	500MG/VIAL	N50554	001	May 24, 1984
	1GM/VIAL	N50554	002	May 24, 1984
	2GM/VIAL	N50554	003	May 24, 1984
	10GM/VIAL	N50554	004	May 24, 1984
	20GM/VIAL	N50554	005	May 24, 1984

CEFOTAXIME SODIUM

INJECTABLE; INJECTION

CEFOTAXIME AND DEXTROSE 2.4% IN PLASTIC CONTAINER				
B BRAUN	EQ 2GM BASE	N50792	001	Jul 29, 2004
CEFOTAXIME AND DEXTROSE 3.9% IN PLASTIC CONTAINER				
B BRAUN	EQ 1GM BASE	N50792	002	Jul 29, 2004
CLAFORAN IN SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER				
SANOFI AVENTIS US	EQ 20MG BASE/ML	N50596	001	May 20, 1985
	EQ 40MG BASE/ML	N50596	003	May 20, 1985

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 49 (of 274)

CEFOTETAN DISODIUMINJECTABLE; INJECTION
CEFOTAN

ASTRAZENECA

EQ 1GM BASE/VIAL
EQ 1GM BASE/VIAL
EQ 2GM BASE/VIAL
EQ 2GM BASE/VIAL
EQ 10GM BASE/VIALN50588 001 Dec 27, 1985
N63293 001 Apr 29, 1993
N50588 002 Dec 27, 1985
N63293 002 Apr 29, 1993
N50588 003 Apr 25, 1988

CEFOTAN IN PLASTIC CONTAINER

ASTRAZENECA

EQ 20MG BASE/ML
EQ 40MG BASE/MLN50694 002 Jul 30, 1993
N50694 001 Jul 30, 1993CEFOTIAM HYDROCHLORIDEINJECTABLE; INJECTION
CERADON

TAKEDA

EQ 1GM BASE/VIAL

N50601 001 Dec 30, 1988

CEFOXITIN SODIUMINJECTABLE; INJECTION
MEFOXIN

MERCK

EQ 1GM BASE/VIAL
EQ 2GM BASE/VIAL
EQ 10GM BASE/VIALN50517 001
N50517 002
N50517 003

MEFOXIN IN DEXTROSE 5% IN PLASTIC CONTAINER

MERCK

EQ 20MG BASE/ML
EQ 40MG BASE/MLN50581 003 Sep 20, 1984
N50581 004 Sep 20, 1984

MEFOXIN IN SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER

MERCK

EQ 20MG BASE/ML
EQ 40MG BASE/MLN50581 002 Sep 20, 1984
N50581 001 Sep 20, 1984CEFPIRAMIDE SODIUMINJECTABLE; INJECTION
CEFPIRAMIDE SODIUM

WYETH AYERST

EQ 1GM BASE/VIAL
EQ 2GM BASE/VIAL
EQ 10GM BASE/VIALN50633 002 Jan 31, 1989
N50633 003 Jan 31, 1989
N50633 005 Jan 31, 1989CEFPODOXIME PROXETILFOR SUSPENSION; ORAL
BANAN

SANKYO

EQ 50MG BASE/5ML
EQ 100MG BASE/5MLN50688 002 Aug 07, 1992
N50688 001 Aug 07, 1992TABLET; ORAL
BANAN

SANKYO

EQ 100MG BASE
EQ 200MG BASEN50687 001 Aug 07, 1992
N50687 002 Aug 07, 1992CEFTAZIDIMEINJECTABLE; INJECTION
TAZIDIME

LILLY

1GM/VIAL
2GM/VIALN62655 001 Nov 20, 1985
N62655 002 Nov 20, 1985

TAZIDIME IN PLASTIC CONTAINER

LILLY

1GM/VIAL
2GM/VIALN62739 001 Jul 10, 1986
N62739 002 Jul 10, 1986CEFTAZIDIME (ARGININE FORMULATION)INJECTABLE; INJECTION
CEPTAZ

GLAXOSMITHKLINE

500MG/VIAL
1GM/VIAL
2GM/VIAL
10GM/VIALN50646 001 Sep 27, 1990
N50646 002 Sep 27, 1990
N50646 003 Sep 27, 1990
N50646 004 Sep 27, 1990

DISCONTINUED DRUG PRODUCT LIST

6 - 50 (of 274)

CEFTAZIDIME (ARGININE FORMULATION)

INJECTABLE; INJECTION
PENTACEF

GLAXOSMITHKLINE	1GM/VIAL	N63322	001	Nov 07, 1995
	1GM/VIAL	N64006	001	Mar 31, 1992
	2GM/VIAL	N63322	002	Nov 07, 1995
	2GM/VIAL	N64006	002	Mar 31, 1992
	6GM/VIAL	N64008	001	Mar 31, 1992
	10GM/VIAL	N64008	002	Mar 31, 1992

CEFTAZIDIME SODIUM

INJECTABLE; INJECTION
CEFTAZIDIME SODIUM IN PLASTIC CONTAINER

BAXTER HLTHCARE	EQ 10MG BASE/ML	N63221	001	Apr 29, 1993
	EQ 20MG BASE/ML	N63221	002	Apr 29, 1993
	EQ 40MG BASE/ML	N63221	003	Apr 29, 1993

FORTAZ IN PLASTIC CONTAINER

GLAXOSMITHKLINE	EQ 10MG BASE/ML	N50634	001	Apr 28, 1989
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CEFTIBUTEN DIHYDRATE

FOR SUSPENSION; ORAL
CEDAX

SHIONOGI	EQ 180MG BASE/5ML	N50686	002	Dec 20, 1995
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CEFTIZOXIME SODIUM

INJECTABLE; INJECTION
CEFIZOX

ASTELLAS	EQ 500MG BASE/VIAL	N50560	001	Sep 15, 1983
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CEFIZOX IN DEXTROSE 5% IN PLASTIC CONTAINER

ASTELLAS	EQ 20MG BASE/ML	N50589	001	Oct 03, 1984
	EQ 40MG BASE/ML	N50589	002	Oct 03, 1984

CEFTRIAXONE SODIUM

INJECTABLE; INJECTION
ROCEPHIN

HLR	EQ 250MG BASE/VIAL	N63239	001	Aug 13, 1993
	EQ 500MG BASE/VIAL	N62654	001	Apr 30, 1987
	EQ 500MG BASE/VIAL	N63239	002	Aug 13, 1993
	EQ 1GM BASE/VIAL	N63239	003	Aug 13, 1993
ROCHE	EQ 250MG BASE/VIAL	N62510	001	Mar 12, 1985
	EQ 500MG BASE/VIAL	N62510	002	Mar 12, 1985
	EQ 1GM BASE/VIAL	N62510	003	Mar 12, 1985

ROCEPHIN W/ DEXTROSE IN PLASTIC CONTAINER

HLR	EQ 10MG BASE/ML	N50624	001	Feb 11, 1987
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CEFTRIAXONE SODIUM; LIDOCAINE

INJECTABLE; INJECTION
ROCEPHIN KIT

HLR	EQ 500MG BASE/VIAL, N/A; N/A, 1%	N50585	007	May 08, 1996
	EQ 1GM BASE/VIAL, N/A; N/A, 1%	N50585	006	May 08, 1996

CEFUROXIME SODIUM

INJECTABLE; IM-IV
KEFUROX

LILLY	EQ 750MG BASE/VIAL	N62591	001	Jan 10, 1986
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INJECTABLE; INJECTION

KEFUROX	EQ 1.5GM BASE/VIAL	N62591	002	Jan 10, 1986
LILLY	EQ 1.5GM BASE/VIAL	N62592	002	Jan 10, 1986
	EQ 7.5GM BASE/VIAL	N62591	003	Dec 17, 1987

KEFUROX IN PLASTIC CONTAINER

LILLY	EQ 1.5GM BASE/VIAL	N62590	002	Jan 10, 1986
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DISCONTINUED DRUG PRODUCT LIST

6 - 51 (of 274)

CEFUROXIME SODIUM

INJECTABLE; INTRAVENOUS

KEFUROX

LILLY

EQ 750MG BASE/VIAL

N62592 001

Jan 10, 1986

KEFUROX IN PLASTIC CONTAINER

LILLY

EQ 750MG BASE/VIAL

N62590 001

Jan 10, 1986

CELLULOSE SODIUM PHOSPHATE

POWDER; ORAL

CALCIBIND

MISSION PHARMA

2.5GM/PACKET

N18757 002

Dec 28, 1982

300GM/BOT

N18757 003

Oct 16, 1984

CEPHALEXIN

CAPSULE; ORAL

CEPHALEXIN

APOTHECON

EQ 250MG BASE

N62973 001

Nov 08, 1988

EQ 250MG BASE

N63186 001

Dec 30, 1994

EQ 500MG BASE

N62974 001

Nov 23, 1988

EQ 500MG BASE

N63186 002

Dec 30, 1994

PUREPAC PHARM

EQ 250MG BASE

N62809 001

Apr 22, 1987

EQ 500MG BASE

N62809 002

Apr 22, 1987

STEVENS J

EQ 500MG BASE

N62869 001

Mar 17, 1988

TEVA

EQ 250MG BASE

N62821 001

Feb 05, 1988

EQ 500MG BASE

N62823 001

Feb 05, 1988

YOSHITOMI

EQ 250MG BASE

N62872 001

Jun 20, 1988

EQ 500MG BASE

N62871 001

Jul 05, 1988

FOR SUSPENSION; ORAL

CEPHALEXIN

APOTHECON

EQ 125MG BASE/5ML

N62986 001

Apr 18, 1991

EQ 250MG BASE/5ML

N62987 001

Jul 25, 1989

BARR

EQ 125MG BASE/5ML

N62778 001

Aug 06, 1987

TEVA

EQ 125MG BASE/5ML

N62873 001

May 23, 1988

EQ 250MG BASE/5ML

N62867 001

Apr 15, 1988

VITARINE

EQ 125MG BASE/5ML

N62779 001

Dec 22, 1987

EQ 250MG BASE/5ML

N62781 001

Dec 22, 1987

KEFLEX

ADVANCIS PHARM

EQ 100MG BASE/ML

N50406 003

EQ 125MG BASE/5ML

N50406 001

EQ 250MG BASE/5ML

N50406 002

TABLET; ORAL

CEPHALEXIN

BARR

EQ 250MG BASE

N62826 001

Aug 17, 1987

EQ 500MG BASE

N62827 001

Aug 17, 1987

VITARINE

EQ 250MG BASE

N62863 001

Aug 11, 1988

EQ 500MG BASE

N62863 002

Aug 11, 1988

EQ 1GM BASE

N62863 003

Aug 11, 1988

KEFLET

LILLY

EQ 250MG BASE

N50440 003

Feb 26, 1987

EQ 250MG BASE

N62745 001

Dec 01, 1986

EQ 500MG BASE

N50440 001

EQ 500MG BASE

N62745 002

Dec 01, 1986

EQ 1GM BASE

N50440 002

CEPHALEXIN HYDROCHLORIDE

TABLET; ORAL

KEFTAB

LILLY

EQ 250MG BASE

N50614 001

Oct 29, 1987

EQ 333MG BASE

N50614 003

May 16, 1988

EQ 500MG BASE

N50614 002

Oct 29, 1987

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 52 (of 274)

CEPHALOGLYCINCAPSULE; ORAL
KAFOCIN
LILLY

250MG

N50219 001

CEPHALOTHIN SODIUMINJECTABLE; INJECTION
CEPHALOTHIN

INTL MEDICATION

EQ 500MG BASE/VIAL
EQ 1GM BASE/VIAL
EQ 2GM BASE/VIAL
EQ 4GM BASE/VIALN62426 001 May 03, 1985
N62426 002 May 03, 1985
N62426 003 May 03, 1985
N62426 004 May 03, 1985CEPHALOTHIN SODIUM
ABBOTTEQ 1GM BASE/VIAL
EQ 1GM BASE/VIAL
EQ 2GM BASE/VIAL
EQ 2GM BASE/VIALN62547 001 Sep 11, 1985
N62548 001 Sep 11, 1985
N62547 002 Sep 11, 1985
N62548 002 Sep 11, 1985

ABRAXIS PHARM

EQ 1GM BASE/VIAL
EQ 2GM BASE/VIALN62666 002 Jun 10, 1987
N62666 001 Jun 10, 1987

BRISTOL

EQ 1GM BASE/VIAL
EQ 2GM BASE/VIAL
EQ 4GM BASE/VIALN62464 001 May 07, 1984
N62464 002 May 07, 1984
N62464 003 May 07, 1984

CEPHALOTHIN SODIUM W/ DEXTROSE IN PLASTIC CONTAINER

BAXTER HLTHCARE

EQ 20MG BASE/ML
EQ 20MG BASE/ML
EQ 20MG BASE/ML
EQ 40MG BASE/ML
EQ 40MG BASE/ML
EQ 40MG BASE/MLN62422 003 Jan 31, 1984
N62422 005 Jul 16, 1991
N62730 001 Mar 05, 1987
N62422 004 Jan 31, 1984
N62422 006 Jul 16, 1991
N62730 002 Mar 05, 1987

CEPHALOTHIN SODIUM W/ SODIUM CHLORIDE IN PLASTIC CONTAINER

BAXTER HLTHCARE

EQ 20MG BASE/ML
EQ 40MG BASE/MLN62422 001 Jan 31, 1984
N62422 002 Jan 31, 1984

KEFLIN

LILLY

EQ 1GM BASE/VIAL
EQ 2GM BASE/VIAL
EQ 4GM BASE/VIAL
EQ 20GM BASE/VIALN50482 001
N50482 002
N50482 003
N50482 007

KEFLIN IN PLASTIC CONTAINER

LILLY

EQ 1GM BASE/VIAL
EQ 2GM BASE/VIALN62549 001 Sep 10, 1985
N62549 002 Sep 10, 1985

SEFFIN

GLAXOSMITHKLINE

EQ 1GM BASE/VIAL
EQ 2GM BASE/VIAL
EQ 10GM BASE/VIALN62435 001 Nov 15, 1983
N62435 002 Nov 15, 1983
N62435 003 Nov 15, 1983CEPHAPIRIN SODIUMINJECTABLE; INJECTION
CEFADYL

APOTHECON

EQ 500MG BASE/VIAL
EQ 500MG BASE/VIAL
EQ 1GM BASE/VIAL
EQ 1GM BASE/VIAL
EQ 1GM BASE/VIAL
EQ 1GM BASE/VIAL
EQ 2GM BASE/VIAL
EQ 2GM BASE/VIAL
EQ 2GM BASE/VIAL
EQ 2GM BASE/VIAL
EQ 2GM BASE/VIAL
EQ 2GM BASE/VIAL
EQ 4GM BASE/VIAL
EQ 4GM BASE/VIAL
EQ 4GM BASE/VIAL
EQ 20GM BASE/VIALN50446 005
N62961 001 Sep 20, 1988
N50446 001
N61769 001
N62724 001 Dec 23, 1986
N62961 002 Sep 20, 1988
N50446 002
N61769 002
N62724 002 Dec 23, 1986
N62961 003 Sep 20, 1988
N50446 003
N61769 003
N62961 004 Sep 20, 1988
N50446 004

CEPHAPIRIN SODIUM

ABRAXIS PHARM

EQ 500MG BASE/VIAL

N62723 001 Nov 17, 1986

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 53 (of 274)

CEPHAPIRIN SODIUMINJECTABLE; INJECTION
CEPHAPIRIN SODIUM

ABRAXIS PHARM	EQ 1GM BASE/VIAL	N62723	002	Nov 17, 1986
	EQ 2GM BASE/VIAL	N62723	003	Nov 17, 1986
	EQ 4GM BASE/VIAL	N62723	004	Nov 17, 1986
	EQ 20GM BASE/VIAL	N62723	005	Nov 17, 1986
BAXTER HLTHCARE	EQ 500MG BASE/VIAL	N62720	001	Jul 02, 1987
	EQ 1GM BASE/VIAL	N62720	002	Jul 02, 1987
	EQ 2GM BASE/VIAL	N62720	003	Jul 02, 1987
	EQ 20GM BASE/VIAL	N62720	004	Jul 02, 1987

CEPHRADINECAPSULE; ORAL
ANSPOR

GLAXOSMITHKLINE	250MG	N61859	001	
	500MG	N61859	002	
CEPHRADINE				
BARR	250MG	N62850	001	Apr 22, 1988
	500MG	N62851	001	Apr 22, 1988
IVAX PHARMS	250MG	N62762	001	Mar 06, 1987
	500MG	N62762	002	Mar 06, 1987
VITARINE	250MG	N62813	001	Feb 25, 1988
	500MG	N62813	002	Feb 25, 1988

VELOSEF

APOTHECON	250MG	N61764	001	
	500MG	N61764	002	
VELOSEF '250'				
ERSANA	250MG	N50548	001	
VELOSEF '500'				
ERSANA	500MG	N50548	002	

FOR SUSPENSION; ORAL
CEPHRADINE

BARR	125MG/5ML	N62858	001	May 19, 1988
	250MG/5ML	N62859	001	May 19, 1988

INJECTABLE; INJECTION

VELOSEF

APOTHECON	250MG/VIAL	N61976	001	
	500MG/VIAL	N61976	002	
	1GM/VIAL	N61976	004	
	2GM/VIAL	N61976	003	
	4GM/VIAL	N61976	005	

TABLET; ORAL

VELOSEF

BRISTOL MYERS SQUIBB	1GM	N50530	001	
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CERIVASTATIN SODIUM

TABLET; ORAL

BAYCOL

BAYER PHARMS	0.05MG	N20740	001	Jun 26, 1997
	0.1MG	N20740	002	Jun 26, 1997
	0.2MG	N20740	003	Jun 26, 1997
	0.3MG	N20740	004	Jun 26, 1997
	0.4MG	N20740	005	May 24, 1999
	0.8MG	N20740	006	Jul 24, 2000

CERULETIDE DIETHYLAMINE

INJECTABLE; INJECTION

TYMTRAN

PHARMACIA AND UPJOHN	0.02MG/ML	N18296	001	
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27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 54 (of 274)

CETYL ALCOHOL; COLFOSCERIL PALMITATE; TYLOXAPOL

FOR SUSPENSION; INTRATRACHEAL

EXOSURF NEONATAL

GLAXOSMITHKLINE 12MG/VIAL;108MG/VIAL;8MG/VIAL

N20044 001 Aug 02, 1990

CHENODIOL

TABLET; ORAL

CHENIX

AXCAN SCANDIPHARM 250MG

N18513 002 Jul 28, 1983

CHLOPHEDIANOL HYDROCHLORIDE

SYRUP; ORAL

ULO

3M 25MG/5ML

N12126 001

CHLORAMPHENICOL

CAPSULE; ORAL

AMPHICOL

FERRANTE 100MG

N60058 001

250MG

N60058 002

CHLORAMPHENICOL

IVAX PHARMS 250MG

N62247 001

CHLOROMYCETIN

PARKEDALE 50MG

N60591 001

100MG

N60591 003

250MG

N60591 002

MYCHEL

ARMENPHARM 250MG

N60851 001

CREAM; TOPICAL

CHLOROMYCETIN

PARKE DAVIS 1%

N50183 001

FOR SOLUTION; OPHTHALMIC

CHLOROMYCETIN

PARKEDALE 25MG/VIAL

N50143 001

INJECTABLE; INJECTION

CHLOROMYCETIN

PARKE DAVIS 250MG/ML

N50153 001

OINTMENT; OPHTHALMIC

CHLOROFAIR

PHARMAFAIR 1%

N62439 001 Apr 21, 1983

CHLOROMYCETIN

PARKEDALE 1%

N50156 001

CHLOROPTIC S.O.P.

ALLERGAN 1%

N61187 001

ECONOCHLOR

ALCON 1%

N61648 001

SOLUTION/DROPS; OPHTHALMIC

CHLORAMPHENICOL

AKORN 0.5%

N62042 001

ALCON 0.5%

N62628 001 Sep 25, 1985

CHLOROFAIR

PHARMAFAIR 0.5%

N62437 001 Apr 14, 1983

CHLOROPTIC

ALLERGAN 0.5%

N50091 001

ECONOCHLOR

ALCON 0.5%

N61645 001

OPHTHOCHLOR

PARKEDALE 0.5%

N61220 001

OPTOMYCIN

OPTOPICS 0.5%

N62171 001 Mar 31, 1982

SOLUTION/DROPS; OTIC

CHLOROMYCETIN

PARKEDALE 0.5%

N50205 001

DISCONTINUED DRUG PRODUCT LIST

6 - 55 (of 274)

CHLORAMPHENICOL PALMITATE

SUSPENSION; ORAL		
CHLOROMYCETIN PALMITATE		
PARKE DAVIS	EQ 150MG BASE/5ML	N50152 001
	EQ 150MG BASE/5ML	N62301 001

CHLORAMPHENICOL SODIUM SUCCINATE

INJECTABLE; INJECTION		
CHLORAMPHENICOL		
ELKINS SINN	EQ 1GM BASE/VIAL	N62406 001 Nov 09, 1982
CHLORAMPHENICOL SODIUM SUCCINATE		
GRUPPO LEPETIT	EQ 1GM BASE/VIAL	N62278 001
MYCHEL-S		
ANGUS	EQ 1GM BASE/VIAL	N60132 001

CHLORAMPHENICOL; DESOXYRIBONUCLEASE; FIBRINOLYSIN

OINTMENT; TOPICAL		
ELASE-CHLOROMYCETIN		
PARKE DAVIS	10MG/GM;666 UNITS/GM;1 UNITS/GM	N50294 001

CHLORAMPHENICOL; HYDROCORTISONE ACETATE

FOR SUSPENSION; OPHTHALMIC		
CHLOROMYCETIN HYDROCORTISONE		
PARKEDALE	12.5MG/VIAL;25MG/VIAL	N50202 001

CHLORAMPHENICOL; HYDROCORTISONE ACETATE; POLYMYXIN B SULFATE

OINTMENT; OPHTHALMIC		
OPHTHOCORT		
PARKEDALE	10MG/GM;5MG/GM;10,000 UNITS/GM	N50201 002

CHLORAMPHENICOL; POLYMYXIN B SULFATE

OINTMENT; OPHTHALMIC		
CHLOROMYXIN		
PARKE DAVIS	1%;10,000 UNITS/GM	N50203 002

CHLORAMPHENICOL; PREDNISOLONE

OINTMENT; OPHTHALMIC		
CHLOROPTIC-P S.O.P.		
ALLERGAN	1%;0.5%	N61188 001

CHLORDIAZEPOXIDE

CAPSULE, EXTENDED RELEASE; ORAL		
LIBRELEASE		
VALEANT PHARM INTL	30MG	N17813 001 Sep 12, 1983
TABLET; ORAL		
LIBRITABS		
VALEANT PHARM INTL	5MG	N85482 001
	10MG	N85481 001
	25MG	N85488 001

CHLORDIAZEPOXIDE HYDROCHLORIDE

CAPSULE; ORAL		
A-POXIDE		
ABBOTT	5MG	N85447 001
	5MG	N85517 001
	10MG	N85447 002
	10MG	N85518 001
	25MG	N85447 003
	25MG	N85513 001
CHLORDIAZACHEL		
RACHELLE	5MG	N85086 001
	10MG	N84639 001
	25MG	N85087 001

DISCONTINUED DRUG PRODUCT LIST

6 - 56 (of 274)

CHLORDIAZEPOXIDE HYDROCHLORIDE

CAPSULE; ORAL

CHLORDIAZEPOXIDE HYDROCHLORIDE

ASCOT	5MG	N87525	001	Jan 07, 1982
	10MG	N87524	001	Jan 07, 1982
	25MG	N87512	001	Jan 07, 1982
FERRANTE	5MG	N85118	001	
	10MG	N85119	001	
	25MG	N85120	001	
HALSEY	5MG	N85340	001	
	10MG	N85339	001	
	25MG	N84685	001	
IMPAX LABS	5MG	N86213	001	
	10MG	N85113	001	
	25MG	N86212	001	
IVAX PHARMS	5MG	N83741	001	
	10MG	N83742	001	
	25MG	N83570	001	
LEDERLE	5MG	N86892	001	
	5MG	N87234	001	
	10MG	N86876	001	
	10MG	N87037	001	
	25MG	N86893	001	
	25MG	N87231	001	
MAST MM	10MG	N86217	001	
MYLAN	5MG	N84886	001	
	10MG	N84601	001	
	25MG	N84887	001	
PARKE DAVIS	5MG	N85163	001	
	10MG	N84598	001	
	25MG	N85164	001	
PIONEER PHARMS	10MG	N89533	001	Jul 15, 1988
	25MG	N89558	001	Jul 15, 1988
PUREPAC PHARM	5MG	N85155	001	
	10MG	N84939	002	
	25MG	N85144	001	
ROXANE	5MG	N84706	001	
	10MG	N84700	001	
	25MG	N84705	001	
SANDOZ	5MG	N84678	001	
	5MG	N84919	001	
	10MG	N84041	001	
	10MG	N84920	001	
	25MG	N84679	002	
	25MG	N84823	001	
SUPERPHARM	5MG	N88987	001	Apr 25, 1985
	10MG	N88986	001	Apr 25, 1985
	25MG	N88988	001	Apr 25, 1985
TEVA	5MG	N88705	001	Jan 18, 1985
	10MG	N88706	001	Jan 18, 1985
	25MG	N86494	001	
	25MG	N88707	001	Jan 18, 1985
USL PHARMA	5MG	N84644	001	
	25MG	N84645	001	
VANGARD	5MG	N88129	001	Mar 28, 1983
	10MG	N88010	001	Mar 28, 1983
	25MG	N88130	001	Mar 28, 1983
WEST WARD	5MG	N85014	001	
	10MG	N85000	001	
	25MG	N85294	001	
LIBRIUM				
VALEANT PHARM INTL	5MG	N12249	002	
	10MG	N12249	001	
	25MG	N12249	003	

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 57 (of 274)

CHLORDIAZEPOXIDE HYDROCHLORIDECAPSULE; ORAL
LYGEN

ALRA	5MG	N85107	001
	10MG	N85009	001
	25MG	N85108	001

INJECTABLE; INJECTION
LIBRIUM

VALEANT PHARM INTL	100MG/AMP	N12301	001
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CHLORDIAZEPOXIDE; ESTROGENS, ESTERIFIED

TABLET; ORAL

MENRIUM 10-4

ROCHE	10MG;0.4MG	N14740	006
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MENRIUM 5-2

ROCHE	5MG;0.2MG	N14740	002
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MENRIUM 5-4

ROCHE	5MG;0.4MG	N14740	004
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CHLORHEXIDINE GLUCONATESOLUTION; TOPICAL
EXIDINE

XTTRIUM	2.5%	N19421	001	Dec 17, 1985
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SPONGE; TOPICAL

CHLORHEXIDINE GLUCONATE

KENDALL IL	4%	N19490	001	Mar 27, 1987
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E-Z SCRUB

BECTON DICKINSON	4%	N73416	001	Mar 14, 2000
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CHLORMERODRIN, HG-197

INJECTABLE; INJECTION

CHLORMERODRIN HG 197

BRACCO	0.6-1.4mCi/ML	N17269	001
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CHLORMEZANONE

TABLET; ORAL

TRANCOPAL

SANOFI AVENTIS US	100MG	N11467	003
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	200MG	N11467	005
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CHLOROPROCAINE HYDROCHLORIDE

INJECTABLE; INJECTION

NESACAINE-MPF

ABRAXIS BIOSCIENCE	2%	N09435	003
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	3%	N09435	004
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CHLOROQUINE HYDROCHLORIDE

INJECTABLE; INJECTION

ARALEN HYDROCHLORIDE

SANOFI AVENTIS US	EQ 40MG BASE/ML	N06002	002
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CHLOROQUINE PHOSPHATE

TABLET; ORAL

CHLOROQUINE PHOSPHATE

MD PHARM	EQ 150MG BASE	N87228	001
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PUREPAC PHARM	EQ 150MG BASE	N80886	001
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TEVA	EQ 150MG BASE	N87504	001	Jan 13, 1982
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WATSON LABS	EQ 150MG BASE	N87979	001	Dec 21, 1982
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	EQ 300MG BASE	N88030	001	Dec 21, 1982
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27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 58 (of 274)

CHLOROQUINE PHOSPHATE; PRIMAQUINE PHOSPHATE

TABLET; ORAL

ARALEN PHOSPHATE W/ PRIMAQUINE PHOSPHATE

SANOFI AVENTIS US	EQ 300MG BASE;EQ 45MG BASE	N14860	002
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CHLOROTHIAZIDE

TABLET; ORAL

CHLOROTHIAZIDE

ABC HOLDING	250MG	N85569	001	
LEDERLE	250MG	N86940	001	
	500MG	N86938	001	
SANDOZ	250MG	N85485	001	
WATSON LABS	250MG	N85165	001	
	250MG	N85173	001	
	250MG	N86795	001	Aug 15, 1983
	500MG	N84026	001	Sep 01, 1982
	500MG	N86796	001	Aug 15, 1983
DIURIL				
OVATION PHARMS	250MG	N11145	004	
	500MG	N11145	002	

CHLOROTHIAZIDE; METHYLDOPA

TABLET; ORAL

ALDOCLOR-150

MERCK	150MG;250MG	N16016	001
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ALDOCLOR-250

MERCK	250MG;250MG	N16016	002
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METHYLDOPA AND CHLOROTHIAZIDE

PAR PHARM	150MG;250MG	N70783	001	Nov 06, 1987
	250MG;250MG	N70654	001	Nov 06, 1987

CHLOROTHIAZIDE; RESERPINE

TABLET; ORAL

CHLOROTHIAZIDE AND RESERPINE

WEST WARD	250MG;0.125MG	N88557	001	Dec 22, 1983
	500MG;0.125MG	N88365	001	Dec 22, 1983

CHLOROTHIAZIDE W/ RESERPINE

WATSON LABS	250MG;0.125MG	N84853	001	
	500MG;0.125MG	N88151	001	Jun 09, 1983

CHLOROTHIAZIDE-RESERPINE

MYLAN	250MG;0.125MG	N87744	001	May 06, 1982
	500MG;0.125MG	N87745	001	May 06, 1982

DIUPRES-250

MERCK	250MG;0.125MG	N11635	003	Aug 26, 1987
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DIUPRES-500

MERCK	500MG;0.125MG	N11635	006	Aug 26, 1987
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CHLOROTRIANISENE

CAPSULE; ORAL

CHLOROTRIANISENE

BANNER PHARMACAPS	12MG	N84652	001
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TACE

SANOFI AVENTIS US	12MG	N08102	004
	25MG	N11444	001
	72MG	N16235	001

CHLORPHENESIN CARBAMATE

TABLET; ORAL

MAOLATE

PHARMACIA AND UPJOHN	400MG	N14217	002
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27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 59 (of 274)

CHLORPHENIRAMINE MALEATE

CAPSULE, EXTENDED RELEASE; ORAL

CHLORPHENIRAMINE MALEATE

SANDOZ 12MG

N70797 001 Aug 12, 1988

TELDRIN

GLAXOSMITHKLINE 8MG

N17369 001

12MG

N17369 002

INJECTABLE; INJECTION

CHLORPHENIRAMINE MALEATE

BEL MAR 10MG/ML

N80821 001

ELKINS SINN 10MG/ML

N80797 001

WATSON LABS 10MG/ML

N83593 001

10MG/ML

N86096 001

100MG/ML

N86095 001

CHLOR-TRIMETON

SCHERING PLOUGH 10MG/ML

N08826 001

100MG/ML

N08794 001

PYRIDAMAL 100

BEL MAR 100MG/ML

N83733 001

SYRUP; ORAL

CHLORPHENIRAMINE MALEATE

PHARM ASSOC 2MG/5ML

N87520 001 Feb 10, 1982

CHLOR-TRIMETON

SCHERING 2MG/5ML

N06921 006

TABLET; ORAL

ANTAGONATE

BAYER PHARMS 4MG

N83381 001

CHLORPHENIRAMINE MALEATE

ANABOLIC 4MG

N83078 001

BELL PHARMA 4MG

N83062 001

ELKINS SINN 4MG

N80938 001

IMPAX LABS 4MG

N80809 001

IVAX PHARMS 4MG

N80779 001

KV PHARM 4MG

N87164 001

LEDERLE 4MG

N86941 001

LEINER 4MG

N80786 001

MUTUAL PHARM 4MG

N80700 001

NEWTRON PHARMS 4MG

N86519 001

PANRAY 4MG

N83243 001

PHARMAVITE 4MG

N85104 001

PHARMERAL 4MG

N83753 001

PIONEER PHARMS 4MG

N88556 001 Jul 13, 1984

PUREPAC PHARM 4MG

N86306 001

ROXANE 4MG

N80626 001

SANDOZ 4MG

N80961 001

VITARINE 4MG

N85837 001

WATSON LABS 4MG

N80696 001

4MG

N80791 001

4MG

N85139 001

4MG

N83787 001

WEST WARD

CHLOR-TRIMETON

SCHERING 4MG

N06921 002

KLOROMIN

HALSEY 4MG

N83629 001

PHENETRON

LANNETT 4MG

N80846 001

TABLET, EXTENDED RELEASE; ORAL

CHLOR-TRIMETON

SCHERING PLOUGH 8MG

N07638 001

EFIDAC 24 CHLORPHENIRAMINE MALEATE

ALZA 16MG

N19746 002 Nov 18, 1994

DISCONTINUED DRUG PRODUCT LIST

6 - 60 (of 274)

CHLORPHENIRAMINE MALEATE; PHENYLPROPANOLAMINE HYDROCHLORIDE

CAPSULE, EXTENDED RELEASE; ORAL			
CHLORPHENIRAMINE MALEATE AND PHENYLPROPANOLAMINE HYDROCHLORIDE			
WATSON LABS	12MG;75MG	N88681 001	Sep 29, 1987
CHLORPHENIRAMINE MALEATE AND PHENYLPROPANOLAMINE HYDROCHLORIDE			
SANDOZ	12MG;75MG	N88940 001	Jan 26, 1989
COLD CAPSULE IV			
GRAHAM DM	12MG;75MG	N18793 001	Apr 25, 1985
COLD CAPSULE V			
GRAHAM DM	8MG;75MG	N18794 001	Apr 23, 1985
CONTAC 12 HOUR			
GLAXOSMITHKLINE	8MG;75MG	N18099 001	
DRIZE			
ASCHER	12MG;75MG	N88359 001	Feb 13, 1986
ORNADE			
GLAXOSMITHKLINE	12MG;75MG	N12152 004	
PHENYLPROPANOLAMINE HYDROCHLORIDE W/ CHLORPHENIRAMINE MALEATE			
CENT PHARMS	8MG;75MG	N18809 001	May 07, 1984
TABLET, EXTENDED RELEASE; ORAL			
CONTAC			
NOVARTIS	12MG;75MG	N19613 001	Jun 13, 1986
DEMAZIN			
SCHERING PLOUGH	4MG;25MG	N18556 001	May 14, 1984
TRIAMINIC-12			
NOVARTIS	12MG;75MG	N18115 001	

CHLORPHENIRAMINE MALEATE; PSEUDOEPHEDRINE HYDROCHLORIDE

CAPSULE, EXTENDED RELEASE; ORAL			
CODIMAL-L.A. 12			
SCHWARZ PHARMA	12MG;120MG	N18935 001	Apr 15, 1985
ISOCLOR			
FISONS	8MG;120MG	N18747 001	Mar 06, 1986
PSEUDOEPHEDRINE HYDROCHLORIDE AND CHLORPHENIRAMINE MALEATE			
CENT PHARMS	8MG;120MG	N19428 001	Aug 02, 1988
GRAHAM DM	8MG;120MG	N18844 001	Mar 20, 1985
	12MG;120MG	N18843 001	Mar 18, 1985
KV PHARM	12MG;120MG	N71455 001	Mar 01, 1989

CHLORPHENIRAMINE POLISTIREX; CODEINE POLISTIREX

SUSPENSION, EXTENDED RELEASE; ORAL			
PENNTUSS			
FISONS	EQ 4MG MALEATE/5ML;EQ 10MG BASE/5ML	N18928 001	Aug 14, 1985

CHLORPHENIRAMINE POLISTIREX; PHENYLPROPANOLAMINE POLISTIREX

SUSPENSION, EXTENDED RELEASE; ORAL			
CORSYM			
UCB INC	EQ 4MG MALEATE/5ML;EQ 37.5MG HCL/5ML	N18050 001	Jan 04, 1984

CHLORPHENTERMINE HYDROCHLORIDE

TABLET; ORAL			
PRE-SATE			
PARKE DAVIS	EQ 65MG BASE	N14696 001	

CHLORPROMAZINE

SUPPOSITORY; RECTAL			
THORAZINE			
GLAXOSMITHKLINE	25MG	N09149 024	
	100MG	N09149 033	

CHLORPROMAZINE HYDROCHLORIDE

CAPSULE, EXTENDED RELEASE; ORAL			
THORAZINE			
GLAXOSMITHKLINE	30MG	N11120 016	
	75MG	N11120 017	

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 61 (of 274)

CHLORPROMAZINE HYDROCHLORIDECAPSULE, EXTENDED RELEASE; ORAL
THORAZINE

GLAXOSMITHKLINE	150MG	N11120	018
	200MG	N11120	019
	300MG	N11120	020

CONCENTRATE; ORAL

CHLORPROMAZINE HYDROCHLORIDE

MORTON GROVE	30MG/ML	N87032	001	Jul 08, 1982
	100MG/ML	N87053	001	

INJECTABLE; INJECTION

CHLORPROMAZINE HYDROCHLORIDE

ABRAXIS PHARM	25MG/ML	N84911	001	
MARSAM PHARMS LLC	25MG/ML	N89563	001	Apr 15, 1988
WATSON LABS	25MG/ML	N80365	001	
	25MG/ML	N85591	001	
WYETH AYERST	25MG/ML	N80370	001	

SYRUP; ORAL

CHLORPROMAZINE HYDROCHLORIDE

ALPHARMA US PHARMS	10MG/5ML	N86712	001
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TABLET; ORAL

CHLORPROMAZINE HYDROCHLORIDE

ABBOTT	10MG	N84414	001	
	25MG	N84415	001	
	50MG	N84411	001	
	100MG	N84412	001	
	200MG	N84413	001	
IVAX PHARMS	10MG	N83549	001	
	25MG	N83549	002	
	50MG	N83549	003	
	100MG	N83574	001	
	200MG	N83575	001	
KV PHARM	10MG	N85750	002	Jan 04, 1982
	25MG	N85751	001	
	50MG	N85484	001	
	100MG	N85752	001	
	200MG	N85748	002	Jan 04, 1982
LEDERLE	10MG	N84803	001	
	25MG	N84801	001	
	50MG	N84800	001	
	100MG	N84789	001	
	200MG	N84802	001	
LEINER	25MG	N80340	001	
	50MG	N80340	002	
	200MG	N80340	003	
PUREPAC PHARM	10MG	N80403	004	
	25MG	N80403	001	
	50MG	N80403	002	
	100MG	N80403	003	
	200MG	N80403	005	
ROXANE	10MG	N85331	001	
	25MG	N85331	002	
	50MG	N85331	003	
	100MG	N85331	004	
	200MG	N85331	005	
VANGARD	10MG	N88038	001	Aug 16, 1982
	25MG	N87645	001	
	50MG	N87646	001	
WATSON LABS	10MG	N85959	001	
	25MG	N85956	001	
	50MG	N85960	001	
	100MG	N85957	001	
	200MG	N85958	001	
WEST WARD	10MG	N87783	001	Sep 16, 1982
	25MG	N87865	001	Sep 16, 1982

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 62 (of 274)

CHLORPROMAZINE HYDROCHLORIDE

TABLET; ORAL

CHLORPROMAZINE HYDROCHLORIDE

WEST WARD	50MG	N87878	001	Sep 15, 1982
	100MG	N87884	001	Sep 15, 1982
	200MG	N87880	001	Sep 16, 1982
PROMAPAR PARKE DAVIS	10MG	N86886	001	
	25MG	N84423	001	
	50MG	N86887	001	
	100MG	N86888	001	
	200MG	N86885	001	
THORAZINE GLAXOSMITHKLINE	10MG	N09149	002	
	25MG	N09149	007	
	50MG	N09149	013	
	100MG	N09149	018	
	200MG	N09149	020	

CHLORPROPAMIDE

TABLET; ORAL

CHLORPROPAMIDE

BARR	100MG	N88812	001	Oct 19, 1984
	100MG	N89446	001	Nov 17, 1986
	250MG	N88813	001	Oct 19, 1984
	250MG	N89447	001	Nov 17, 1986
CLONMEL HLTHCARE	100MG	N89561	001	Sep 04, 1987
	250MG	N89562	001	Sep 04, 1987
DURAMED PHARMS BARR	100MG	N88918	001	Oct 16, 1984
	250MG	N88919	001	Oct 16, 1984
HALSEY	100MG	N89321	001	Jan 16, 1986
	250MG	N88662	001	Jan 09, 1986
IVAX PHARMS	100MG	N88840	001	Oct 25, 1984
	250MG	N87353	001	
SANDOZ	250MG	N84669	001	
SUPERPHARM	100MG	N88694	001	Sep 17, 1984
	250MG	N88695	001	Sep 17, 1984
TEVA	100MG	N88768	001	Oct 11, 1984
USL PHARMA	100MG	N88708	001	Aug 30, 1984
	250MG	N88709	001	Aug 30, 1984
WATSON LABS	100MG	N86865	001	Sep 24, 1984
	100MG	N88608	001	Apr 12, 1984
	250MG	N86866	001	
	250MG	N88568	001	Apr 12, 1984
GLUCAMIDE				
TEVA	250MG	N88641	001	Oct 11, 1984

CHLORPROTHIXENE

CONCENTRATE; ORAL

TARACTAN

ROCHE 100MG/5ML

N16149 002

INJECTABLE; INJECTION

TARACTAN

ROCHE 12.5MG/ML

N12487 001

TABLET; ORAL

TARACTAN

ROCHE 10MG

N12486 005

25MG

N12486 004

50MG

N12486 003

100MG

N12486 001

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 63 (of 274)

CHLORTETRACYCLINE HYDROCHLORIDE

OINTMENT; OPHTHALMIC

AUREOMYCIN

LEDERLE

1%

N50404 001

CHLORTHALIDONE

TABLET; ORAL

CHLORTHALIDONE

ABBOTT

25MG

N87364 001

50MG

N87384 001

ASCOT

25MG

N87698 001

Oct 20, 1982

50MG

N87699 001

Oct 20, 1982

CLONMEL HLTHCARE

25MG

N87451 001

50MG

N87450 001

IVAX PHARMS

25MG

N87555 001

25MG

N88164 001

Jan 09, 1984

50MG

N87176 001

50MG

N87947 001

Feb 27, 1984

KV PHARM

25MG

N87311 001

50MG

N87312 001

MUTUAL PHARM

25MG

N87292 001

25MG

N89285 001

Jul 21, 1986

25MG

N89738 001

Sep 19, 1988

50MG

N87293 001

50MG

N89286 001

Jul 21, 1986

50MG

N89739 001

Sep 19, 1988

PIONEER PHARMS

50MG

N89591 001

Jul 21, 1988

PUREPAC PHARM

25MG

N88139 001

Jul 16, 1986

50MG

N88140 001

Aug 11, 1983

SANDOZ

25MG

N87380 001

50MG

N87118 001

50MG

N87381 001

SUPERPHARM

25MG

N87473 001

Feb 09, 1983

50MG

N87247 001

Feb 09, 1983

TEVA

50MG

N88651 001

May 30, 1985

USL PHARMA

25MG

N89051 001

Jun 01, 1987

50MG

N89052 001

Jun 01, 1987

VANGARD

25MG

N88012 001

Jul 14, 1982

50MG

N88073 001

Mar 25, 1983

WARNER CHILCOTT

25MG

N87515 001

Jan 24, 1983

50MG

N87516 001

Feb 09, 1983

WATSON LABS

25MG

N87050 001

25MG

N87100 001

25MG

N87296 001

25MG

N87706 001

50MG

N87029 001

50MG

N87082 001

50MG

N87521 001

50MG

N87689 001

HYGROTON

SANOFI AVENTIS US

25MG

N12283 004

50MG

N12283 003

THALITONE

MONARCH PHARMS

25MG

N19574 002

Feb 12, 1992

25MG

N88051 001

Nov 12, 1982

CHLORTHALIDONE; CLONIDINE HYDROCHLORIDE

TABLET; ORAL

CLONIDINE HYDROCHLORIDE AND CHLORTHALIDONE

PAR PHARM

15MG;0.1MG

N71179 001

Dec 16, 1987

15MG;0.2MG

N71178 001

Dec 16, 1987

15MG;0.3MG

N71142 001

Dec 16, 1987

COMBIPRES

BOEHRINGER INGELHEIM

15MG;0.1MG

N17503 001

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 64 (of 274)

CHLORTHALIDONE; CLONIDINE HYDROCHLORIDETABLET; ORAL
COMBIPRESBOEHRINGER INGELHEIM 15MG;0.2MG
15MG;0.3MG

N17503 002

N17503 003 Apr 10, 1984

CHLORTHALIDONE; METOPROLOL TARTRATECAPSULE; ORAL
LOPRESSIDONENOVARTIS 25MG;100MG
25MG;200MG

N19451 001

Dec 31, 1987

N19451 002

Dec 31, 1987

CHLORTHALIDONE; RESERPINETABLET; ORAL
DEMI-REGROTON

SANOFI AVENTIS US 25MG;0.125MG

N15103 002

REGROTON

SANOFI AVENTIS US 50MG;0.25MG

N15103 001

CHLORZOXAZONETABLET; ORAL
CHLORZOXAZONE

PIONEER PHARMS 250MG

N89592 001

Jan 06, 1989

500MG

N89948 001

Jan 06, 1989

WATSON LABS 250MG

N86901 001

250MG

N86948 001

Aug 09, 1982

500MG

N81019 001

Jul 29, 1991

PARAFLEX

ORTHO MCNEIL PHARM 250MG

N11300 003

CHOLESTYRAMINEBAR, CHEWABLE; ORAL
CHOLYBAR

PARKE DAVIS EQ 4GM RESIN/BAR

N71621 001

May 26, 1988

EQ 4GM RESIN/BAR

N71739 001

May 26, 1988

POWDER; ORAL
CHOLESTYRAMINE

IVAX PHARMS EQ 4GM RESIN/PACKET

N74771 001

Jul 09, 1997

EQ 4GM RESIN/SCOOPFUL

N74771 002

Jul 09, 1997

TABLET; ORAL
QUESTRAN

APOTHECON EQ 800MG RESIN

N73403 002

Dec 27, 1999

EQ 1GM RESIN

N73403 001

Apr 28, 1994

CHORIOGONADOTROPIN ALFAINJECTABLE; INJECTION
OVIDREL

SERONO INC 0.25MG/VIAL

N21149 001

Sep 20, 2000

CHROMIC CHLORIDEINJECTABLE; INJECTION
CHROMIC CHLORIDE

ABRAXIS PHARM EQ 0.004MG CHROMIUM/ML

N19271 001

May 05, 1987

CHYMOPAPAININJECTABLE; INJECTION
CHYMODIACTIN

ABBOTT 4,000 UNITS/VIAL

N18663 002

Aug 21, 1984

10,000 UNITS/VIAL **Federal Register
determination that product was not
discontinued or withdrawn for safety
or efficacy reasons**

N18663 001

Nov 10, 1982

DISCASE

ABBOTT 12,500 UNITS/VIAL

N18625 001

Jan 18, 1984

DISCONTINUED DRUG PRODUCT LIST

6 - 65 (of 274)

CHYMOTRYPSIN

FOR SOLUTION; OPHTHALMIC

ALPHA CHYMAR

SOLA BARNES HIND 750 UNITS/VIAL

N11837 001

CATARASE

CIBA 300 UNITS/VIAL

N16938 001

NOVARTIS 150 UNITS/VIAL

N18121 001

ZOLYSE

ALCON 750 UNITS/VIAL

N11903 001

CILASTATIN SODIUM; IMIPENEM

INJECTABLE; INJECTION

PRIMAXIN

MERCK EQ 750MG BASE/VIAL;750MG/VIAL

N50630 002 Dec 14, 1990

CIMETIDINE

SUSPENSION; ORAL

TAGAMET HB 200

GLAXOSMITHKLINE 200MG/20ML

N20951 001 Jul 09, 1999

TABLET; ORAL

CIMETIDINE

ENDO PHARMS 200MG

N74281 001 May 17, 1994

300MG

N74281 002 May 17, 1994

400MG

N74281 003 May 17, 1994

800MG

N74329 001 May 17, 1994

IVAX PHARMS 200MG

N74401 001 May 30, 1995

300MG

N74401 002 May 30, 1995

400MG

N74401 003 May 30, 1995

LEK PHARMS 100MG

N75122 001 Jun 19, 1998

200MG

N74250 001 Jun 29, 1995

PERRIGO 100MG

N74972 001 Jun 19, 1998

ROXANE 300MG

N74361 001 Dec 23, 1994

400MG

N74361 002 Dec 23, 1994

800MG

N74371 001 Dec 23, 1994

TAGAMET HB 100MG

N20238 001 Jun 19, 1995

CIMETIDINE HYDROCHLORIDE

INJECTABLE; INJECTION

CIMETIDINE HYDROCHLORIDE

ENDO PHARMS EQ 300MG BASE/2ML

N74005 001 Aug 31, 1994

HOSPIRA EQ 300MG BASE/2ML

N74296 001 Mar 28, 1997

EQ 300MG BASE/2ML

N74412 001 Mar 28, 1997

EQ 300MG BASE/2ML

N74422 001 Jan 31, 1995

LUITPOLD EQ 300MG BASE/2ML

N74353 001 Dec 20, 1994

CIMETIDINE HYDROCHLORIDE IN SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER

HOSPIRA EQ 90MG BASE/100ML

N74468 005 Dec 29, 1994

EQ 120MG BASE/100ML

N74468 006 Dec 29, 1994

EQ 180MG BASE/100ML

N74468 003 Dec 29, 1994

EQ 240MG BASE/100ML

N74468 004 Dec 29, 1994

EQ 360MG BASE/100ML

N74468 001 Dec 29, 1994

EQ 480MG BASE/100ML

N74468 002 Dec 29, 1994

TAGAMET

GLAXOSMITHKLINE EQ 300MG BASE/2ML

N17939 002

TAGAMET HYDROCHLORIDE IN SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER

GLAXOSMITHKLINE EQ 6MG BASE/ML

N19434 001 Oct 31, 1985

SOLUTION; ORAL

CIMETIDINE HYDROCHLORIDE

ENDO PHARMS EQ 300MG BASE/5ML

N74251 001 Dec 22, 1994

ROXANE EQ 300MG BASE/5ML

N74541 001 Aug 05, 1997

TAGAMET

GLAXOSMITHKLINE EQ 300MG BASE/5ML

N17924 001

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 66 (of 274)

CINOXACINCAPSULE; ORAL
CINOBAC
LILLY250MG
500MGN18067 001
N18067 002CINOXACIN
TEVA250MG
500MGN73005 001 Feb 28, 1992
N73006 001 Feb 28, 1992CIPROFLOXACININJECTABLE; INJECTION
CIPROBAYER PHARMS 1200MG/120ML (10MG/ML)
CIPRO IN SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER
BAYER PHARMS 200MG/100MLN19847 003 Dec 26, 1990
N19858 001 Dec 26, 1990CISAPRIDE MONOHYDRATESUSPENSION; ORAL
PROPULSID

JANSSEN PHARMA EQ 1MG BASE/ML

N20398 001 Sep 15, 1995

TABLET; ORAL
PROPULSIDJANSSEN PHARMA EQ 10MG BASE
EQ 20MG BASEN20210 001 Jul 29, 1993
N20210 002 Dec 23, 1993TABLET, ORALLY DISINTEGRATING; ORAL
PROPULSID QUICKSOLV

JANSSEN PHARMA EQ 20MG BASE

N20767 001 Nov 07, 1997

CISPLATININJECTABLE; INJECTION
PLATINOL-AQ

BRISTOL MYERS 0.5MG/ML

N18057 003 Jul 18, 1984

CITALOPRAM HYDROBROMIDETABLET; ORAL
CELEXA

FOREST LABS EQ 60MG BASE

N20822 004 Jul 17, 1998

TABLET, ORALLY DISINTEGRATING; ORAL
CITALOPRAM HYDROBROMIDEBIOVAIL LABS INTL EQ 10MG BASE
EQ 20MG BASE
EQ 40MG BASEN21763 001 Dec 20, 2005
N21763 002 Dec 20, 2005
N21763 003 Dec 20, 2005CITRIC ACID; MAGNESIUM OXIDE; SODIUM CARBONATESOLUTION; IRRIGATION
IRRIGATING SOLUTION G IN PLASTIC CONTAINER

BAXTER HLTHCARE 3.24GM/100ML; 380MG/100ML; 430MG/100ML

N18519 001 Jun 22, 1982

CITRIC ACID; UREA, C-13FOR SOLUTION, TABLET, FOR SOLUTION; ORAL
IDKIT:HP

BREATHID 2006 N/A, 4GM; 75MG, N/A

N21314 001 Dec 17, 2002

CLARITHROMYCINFOR SUSPENSION; ORAL
BIAXIN
ABBOTT

187MG/5ML

N50698 003 Sep 30, 1998

CLEMASTINE FUMARATESYRUP; ORAL
TAVIST

NOVARTIS EQ 0.5MG BASE/5ML

N18675 001 Jun 28, 1985

DISCONTINUED DRUG PRODUCT LIST

6 - 67 (of 274)

CLEMASTINE FUMARATE

TABLET; ORAL				
CLEMASTINE FUMARATE				
TEVA	1.34MG	N73282	001	Jan 31, 1992
TAVIST				
NOVARTIS	2.68MG	N17661	001	
TAVIST-1				
NOVARTIS	1.34MG	N17661	002	
	1.34MG	N17661	003	Aug 21, 1992

CLEMASTINE FUMARATE; PHENYLPROPANOLAMINE HYDROCHLORIDE

TABLET, EXTENDED RELEASE; ORAL				
TAVIST-D				
NOVARTIS	EQ 1MG BASE;75MG	N18298	001	Dec 15, 1982
	1.34MG;75MG	N18298	002	Aug 21, 1992
	1.34MG;75MG	N20640	001	Aug 09, 1996

CLIDINIUM BROMIDE

CAPSULE; ORAL				
QUARZAN				
ROCHE	2.5MG	N10355	001	
	5MG	N10355	002	

CLINDAMYCIN HYDROCHLORIDE

CAPSULE; ORAL				
CLEOCIN				
PHARMACIA AND UPJOHN	EQ 75MG BASE	N61809	001	
	EQ 150MG BASE	N61809	002	
CLINDAMYCIN HYDROCHLORIDE				
WATSON LABS	EQ 75MG BASE	N63082	001	Jul 31, 1991

CLINDAMYCIN PALMITATE HYDROCHLORIDE

FOR SOLUTION; ORAL				
CLEOCIN				
PHARMACIA AND UPJOHN	EQ 75MG BASE/5ML	N61827	001	

CLINDAMYCIN PHOSPHATE

CREAM; VAGINAL				
CLEOCIN				
PHARMACIA AND UPJOHN	EQ 2% BASE	N50680	001	Aug 11, 1992
INJECTABLE; INJECTION				
CLEOCIN PHOSPHATE				
PHARMACIA AND UPJOHN	EQ 150MG BASE/ML	N61839	001	
CLINDAMYCIN PHOSPHATE				
ABRAXIS PHARM	EQ 150MG BASE/ML	N62747	001	Jun 03, 1988
ASTRAZENECA	EQ 150MG BASE/ML	N62928	001	Feb 13, 1989
BAXTER HLTHCARE	EQ 150MG BASE/ML	N62806	001	Oct 15, 1987
	EQ 150MG BASE/ML	N62953	001	Apr 21, 1988
	EQ 150MG BASE/ML	N63068	001	Aug 28, 1989
BEDFORD	EQ 150MG BASE/ML	N63163	001	Jun 30, 1994
BRISTOL MYERS SQUIBB	EQ 150MG BASE/ML	N62908	001	Feb 01, 1989
LOCH	EQ 150MG BASE/ML	N62905	001	May 09, 1988
MARSAM PHARMS LLC	EQ 150MG BASE/ML	N62913	001	Oct 20, 1988
SICOR PHARMS	EQ 150MG BASE/ML	N63041	001	Dec 29, 1989
	EQ 150MG BASE/ML	N63282	001	May 29, 1992
SOLOPAK	EQ 150MG BASE/ML	N62819	001	Mar 15, 1988
	EQ 150MG BASE/ML	N62852	001	Mar 17, 1988
WATSON LABS	EQ 150MG BASE/ML	N62900	001	Jun 08, 1988
	EQ 150MG BASE/ML	N63079	001	Mar 05, 1990
CLINDAMYCIN PHOSPHATE IN DEXTROSE 5%				
ABRAXIS PHARM	EQ 12MG BASE/ML	N50636	001	Dec 22, 1989
CLINDAMYCIN PHOSPHATE IN DEXTROSE 5% IN PLASTIC CONTAINER				
ABBOTT	EQ 6MG BASE/ML	N65027	001	Jun 29, 2001
	EQ 12MG BASE/ML	N65027	002	Jun 29, 2001

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 68 (of 274)

CLINDAMYCIN PHOSPHATE

INJECTABLE; INJECTION

CLINDAMYCIN PHOSPHATE IN DEXTROSE 5% IN PLASTIC CONTAINER

ABBOTT	EQ 18MG BASE/ML	N65027	003	Jun 29, 2001
BAXTER HLTHCARE	EQ 6MG BASE/ML	N50648	001	Dec 29, 1989
	EQ 12MG BASE/ML	N50648	002	Dec 29, 1989
	EQ 900MG BASE/100ML	N50648	003	Dec 29, 1989

SOLUTION; TOPICAL

CLEOCIN T

PHARMACIA AND UPJOHN	EQ 1% BASE	N62363	001	Feb 08, 1982
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CLINDAMYCIN PHOSPHATE

COPLEY PHARM	EQ 1% BASE	N62944	001	Jan 11, 1989
TEVA	EQ 1% BASE	N62930	001	Jun 28, 1989

CLIOQUINOL; NYSTATIN

OINTMENT; TOPICAL

NYSTAFORM

BAYER PHARMS	10MG/GM;100,000 UNITS/GM	N50235	001	
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CLOBETASOL PROPIONATE

SOLUTION; TOPICAL

CLOBETASOL PROPIONATE

ALTANA	0.05%	N75391	001	Feb 08, 1999
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CLOFAZIMINE

CAPSULE; ORAL

LAMPRENE

NOVARTIS	100MG	N19500	001	Dec 15, 1986
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CLOFIBRATE

CAPSULE; ORAL

ATROMID-S

WYETH AYERST	500MG	N16099	002	
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CLOFIBRATE

SANDOZ	500MG	N72191	001	May 02, 1988
WATSON LABS	500MG	N71603	001	Sep 18, 1987

CLONAZEPAM

TABLET; ORAL

KLONOPIN

ROCHE	0.125MG	N17533	005	Apr 09, 1997
	0.25MG	N17533	006	Apr 09, 1997

CLONIDINE HYDROCHLORIDE

TABLET; ORAL

CLONIDINE HYDROCHLORIDE

AM THERAP	0.1MG	N70881	001	Jul 08, 1986
	0.2MG	N70882	001	Jul 08, 1986
	0.3MG	N70883	001	Jul 08, 1986
DURAMED PHARMS BARR	0.1MG	N71103	001	Aug 14, 1986
	0.2MG	N71102	001	Aug 14, 1986
	0.3MG	N71101	001	Aug 14, 1986
INTERPHARM	0.1MG	N71252	001	Oct 01, 1986
	0.2MG	N71253	001	Oct 01, 1986
	0.3MG	N71254	001	Oct 01, 1986
PAR PHARM	0.1MG	N70461	001	Jul 08, 1986
	0.2MG	N70460	001	Jul 08, 1986
	0.3MG	N70459	001	Jul 08, 1986
TEVA	0.1MG	N70747	001	Jul 08, 1986
	0.2MG	N70702	001	Jul 08, 1986
	0.3MG	N70659	001	Jul 08, 1986
WARNER CHILCOTT	0.1MG	N72138	001	Jun 13, 1988
	0.2MG	N72139	001	Jun 13, 1988
	0.3MG	N72140	001	Jun 13, 1988

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 69 (of 274)

CLONIDINE HYDROCHLORIDE

TABLET; ORAL

CLONIDINE HYDROCHLORIDE

WATSON LABS

0.1MG

N70395 001

Mar 23, 1987

0.2MG

N70396 001

Mar 23, 1987

0.3MG

N70397 001

Mar 23, 1987

CLORAZEPATE DIPOTASSIUM

CAPSULE; ORAL

CLORAZEPATE DIPOTASSIUM

ABLE

3.75MG

N71777 001

Jul 14, 1987

7.5MG

N71778 001

Jul 14, 1987

15MG

N71779 001

Jul 14, 1987

AM THERAP

3.75MG

N71429 001

Jun 23, 1987

7.5MG

N71430 001

Jun 23, 1987

15MG

N71431 001

Jun 23, 1987

CLONMEL HLTHCARE

3.75MG

N71742 001

Dec 14, 1987

7.5MG

N71743 001

Dec 14, 1987

15MG

N71744 001

Dec 14, 1987

GD SEARLE LLC

3.75MG

N71727 001

Dec 18, 1987

7.5MG

N71728 001

Dec 18, 1987

15MG

N71729 001

Dec 18, 1987

PUREPAC PHARM

3.75MG

N71924 001

Apr 25, 1988

7.5MG

N71925 001

Apr 25, 1988

15MG

N71926 001

Apr 25, 1988

QUANTUM PHARMICS

3.75MG

N71549 001

Sep 12, 1988

7.5MG

N71550 001

Sep 12, 1988

15MG

N71522 001

Sep 12, 1988

SANDOZ

7.5MG

N72220 001

Aug 26, 1988

USL PHARMA

3.75MG

N71242 001

Jun 23, 1987

7.5MG

N71243 001

Jun 23, 1987

15MG

N71244 001

Jun 23, 1987

WARNER CHILCOTT

3.75MG

N71774 001

Mar 01, 1988

7.5MG

N71775 001

Mar 01, 1988

15MG

N71776 001

Mar 01, 1988

WATSON LABS

3.75MG

N71878 001

Mar 15, 1988

7.5MG

N71879 001

Mar 15, 1988

15MG

N71860 001

Mar 15, 1988

TRANXENE

OVATION PHARMS

3.75MG

N17105 001

7.5MG

N17105 002

15MG

N17105 003

TABLET; ORAL

CLORAZEPATE DIPOTASSIUM

ABLE

3.75MG

N71780 001

Jun 26, 1987

7.5MG

N71781 001

Jun 26, 1987

15MG

N71782 001

Jun 26, 1987

AM THERAP

3.75MG

N71747 001

Jun 23, 1987

7.5MG

N71748 001

Jun 23, 1987

15MG

N71749 001

Jun 23, 1987

LEDERLE

3.75MG

N72013 001

Dec 15, 1987

7.5MG

N72014 001

Dec 15, 1987

15MG

N72015 001

Dec 15, 1987

PUREPAC PHARM

3.75MG

N72330 001

Aug 08, 1988

7.5MG

N72331 001

Aug 08, 1988

15MG

N72332 001

Aug 08, 1988

QUANTUM PHARMICS

3.75MG

N71730 001

Oct 26, 1987

7.5MG

N71731 001

Oct 26, 1987

15MG

N71702 001

Oct 26, 1987

SANDOZ

3.75MG

N72512 001

May 11, 1990

WARNER CHILCOTT

3.75MG

N71828 001

Mar 03, 1988

7.5MG

N71829 001

Mar 03, 1988

15MG

N71830 001

Mar 03, 1988

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 70 (of 274)

CLOTRIMAZOLECREAM; TOPICAL
LOTRIMIN

SCHERING PLOUGH 1%

N17619 001

MYCELEX

BAYER PHARMS 1%

N18183 001

LOTION; TOPICAL
LOTRIMIN

SCHERING 1%

N18813 001 Feb 17, 1984

SOLUTION; TOPICAL
LOTRIMIN

SCHERING PLOUGH 1%

N17613 001

TABLET; VAGINAL
MYCELEX-G

BAYER PHARMS 500MG

N19069 001 Apr 19, 1985

CLOXACILLIN SODIUM

CAPSULE; ORAL

CLOXACILLIN SODIUM

APOTHECON EQ 250MG BASE
EQ 500MG BASE

N61452 001

N61452 002

CLOXAPEN

GLAXOSMITHKLINE EQ 250MG BASE
EQ 500MG BASE

N62233 001

N62233 002

FOR SOLUTION; ORAL
CLOXACILLIN SODIUM

TEVA EQ 125MG BASE/5ML

N62978 001 Apr 06, 1989

TEGOPEN

APOTHECON EQ 125MG BASE/5ML

N50192 001

EQ 125MG BASE/5ML

N61453 001

CLOZAPINE

TABLET; ORAL

CLOZAPINE

SANDOZ 25MG

N74546 001 Aug 30, 1996

100MG

N74546 002 Aug 30, 1996

COBALT CHLORIDE, CO-57; CYANOCOBALAMIN; CYANOCOBALAMIN, CO-57; INTRINSIC FACTOR

N/A; N/A

RUBRATOPE-57 KIT

BRACCO N/A;N/A;N/A;N/A

N16089 001

COBALT CHLORIDE, CO-60; CYANOCOBALAMIN; CYANOCOBALAMIN, CO-60; INTRINSIC FACTOR

N/A; N/A

RUBRATOPE-60 KIT

BRACCO N/A;N/A;N/A;N/A

N16090 001

CODEINE PHOSPHATE; PHENYLEPHRINE HYDROCHLORIDE; PROMETHAZINE HYDROCHLORIDE

SYRUP; ORAL

PHENERGAN VC W/ CODEINE

WYETH AYERST 10MG/5ML;5MG/5ML;6.25MG/5ML

N08306 005 Apr 02, 1984

PHERAZINE VC W/ CODEINE

HALSEY 10MG/5ML;5MG/5ML;6.25MG/5ML

N88870 001 Mar 02, 1987

PROMETHAZINE VC W/ CODEINE

CENCI 10MG/5ML;5MG/5ML;6.25MG/5ML

N88816 001 Nov 22, 1985

MORTON GROVE 10MG/5ML;5MG/5ML;6.25MG/5ML

N88896 001 Jan 04, 1985

CODEINE PHOSPHATE; PROMETHAZINE HYDROCHLORIDE

SYRUP; ORAL

PHENERGAN W/ CODEINE

WYETH AYERST 10MG/5ML;6.25MG/5ML

N08306 004 Apr 02, 1984

PHERAZINE W/ CODEINE

HALSEY 10MG/5ML;6.25MG/5ML

N88739 001 Dec 23, 1988

DISCONTINUED DRUG PRODUCT LIST

6 - 71 (of 274)

CODEINE PHOSPHATE; PROMETHAZINE HYDROCHLORIDE

SYRUP; ORAL		
PROMETHAZINE W/ CODEINE		
CENCI	10MG/5ML;6.25MG/5ML	N88814 001 Nov 22, 1985

CODEINE PHOSPHATE; PSEUDOEPHEDRINE HYDROCHLORIDE; TRIPROLIDINE HYDROCHLORIDE

SYRUP; ORAL		
ACTIFED W/ CODEINE		
GLAXOSMITHKLINE	10MG/5ML;30MG/5ML;1.25MG/5ML	N12575 003 Apr 04, 1984
TRIPROLIDINE AND PSEUDOEPHEDRINE HYDROCHLORIDES W/ CODEINE		
CENCI	10MG/5ML;30MG/5ML;1.25MG/5ML	N89018 001 Jul 23, 1986

COLCHICINE; PROBENECID

TABLET; ORAL		
COLBENEMID		
MERCK	0.5MG;500MG	N12383 001
PROBEN-C		
WATSON LABS	0.5MG;500MG	N85552 001
PROBENECID AND COLCHICINE		
BEECHAM	0.5MG;500MG	N84321 001
IMPAX LABS	0.5MG;500MG	N83720 002
SANDOZ	0.5MG;500MG	N86130 001
PROBENECID W/ COLCHICINE		
LEDERLE	0.5MG;500MG	N86954 001
WATSON LABS	0.5MG;500MG	N83221 001

COLESEVELAM HYDROCHLORIDE

CAPSULE; ORAL		
WELCHOL		
DAIICHI SANKYO	375MG	N21141 001 May 26, 2000

COLISTIN SULFATE

SUSPENSION; ORAL		
COLY-MYCIN S		
PARKE DAVIS	EQ 25MG BASE/5ML	N50355 001

COPPER

INTRAUTERINE DEVICE; INTRAUTERINE		
CU-7		
GD SEARLE LLC	89MG	N17408 001
TATUM-T		
GD SEARLE LLC	120MG	N18205 001

CORTICOTROPIN

INJECTABLE; INJECTION		
ACTH		
PARKEDALE	25 UNITS/VIAL	N08317 002
	40 UNITS/VIAL	N08317 004
ACTHAR		
SANOFI AVENTIS US	25 UNITS/VIAL	N07504 002
	40 UNITS/VIAL	N07504 003
CORTICOTROPIN		
ORGANICS LAGRANGE	40 UNITS/ML	N10831 001
	80 UNITS/ML	N10831 002
WATSON LABS	40 UNITS/VIAL	N88772 001 Nov 21, 1984
H.P. ACTHAR GEL		
QUESTCOR PHARMS	40 UNITS/ML	N08372 006
PURIFIED CORTROPHIN GEL		
ORGANON USA INC	40 UNITS/ML	N08975 001
	80 UNITS/ML	N08975 002

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 72 (of 274)

CORTICOTROPIN-ZINC HYDROXIDEINJECTABLE; INJECTION
CORTROPHIN-ZINC

ORGANON USA INC 40 UNITS/ML N09854 001

CORTISONE ACETATEINJECTABLE; INJECTION
CORTISONE ACETATE

PHARMACIA AND UPJOHN 25MG/ML N08126 002

WATSON LABS 25MG/ML N83147 003

25MG/ML N85677 001

50MG/ML N83147 004

50MG/ML N85677 002

CORTONE

MERCK 25MG/ML N07110 002

50MG/ML N07110 003

TABLET; ORAL

CORTISONE ACETATE

BARR 25MG N83471 001

ELKINS SINN 25MG N80836 001

EVERYLIFE 25MG N84246 001

HEATHER 25MG N85736 001

IMPAX LABS 25MG N09458 001

INWOOD LABS 25MG N80731 001

IVAX PHARMS 25MG N80630 001

25MG N83536 001

LANNETT 25MG N80694 001

PANRAY 5MG N08284 002

25MG N08284 001

PUREPAC PHARM 25MG N80493 001

VITARINE 25MG N80333 001

WATSON LABS 25MG N85884 001

WHITEWORTH TOWN PLSN 25MG N80341 001

CORTONE

MERCK 25MG N07750 003

CROMOLYN SODIUM

CAPSULE; INHALATION

INTAL

SANOFI AVENTIS US 20MG N16990 001

CAPSULE; ORAL

GASTROCROM

UCB INC 100MG N19188 001 Dec 22, 1989

SOLUTION/DROPS; OPHTHALMIC

CROMOPTIC

KING PHARMS 4% N75088 001 Apr 27, 1999

SPRAY, METERED; NASAL

NASALCROM

MCNEIL CONS 5.2MG/SPRAY N20463 001 Jan 03, 1997

CRYPTENAMINE ACETATES

INJECTABLE; INJECTION

UNITENSEN

MEDPOINTE PHARM HLC 260CSR UNIT/ML N08814 001

CRYPTENAMINE TANNATES

TABLET; ORAL

UNITENSEN

MEDPOINTE PHARM HLC 260CSR UNIT N09217 001

CUPRIC SULFATE

INJECTABLE; INJECTION

CUPRIC SULFATE

ABRAXIS PHARM EQ 0.4MG COPPER/ML N19350 001 May 05, 1987

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 73 (of 274)

CYANOCOBALAMIN

GEL, METERED; NASAL				
NASCOBAL				
QOL MEDCL	0.5MG/INH	N19722	001	Nov 05, 1996
INJECTABLE; INJECTION				
BERUBIGEN				
PHARMACIA AND UPJOHN	1MG/ML	N06798	001	
BETALIN 12				
LILLY	0.1MG/ML	N80855	001	
	1MG/ML	N80855	002	
COBAVITE				
WATSON LABS	0.1MG/ML	N83013	001	
	1MG/ML	N83064	001	
CYANOCOBALAMIN				
ABRAXIS PHARM	0.03MG/ML	N80510	003	
	0.1MG/ML	N80510	001	
	0.1MG/ML	N80557	002	
	1MG/ML	N80510	002	
	1MG/ML	N83075	001	
AKORN	1MG/ML	N87969	001	Nov 10, 1983
BAXTER HLTHCARE	1MG/ML	N80515	002	
DELL LABS	0.03MG/ML	N80689	001	
	0.1MG/ML	N80689	002	
	1MG/ML	N80689	003	
LUITPOLD	0.03MG/ML	N80668	001	
SANOFI AVENTIS US	1MG/ML	N80564	001	
SOLOPAK	1MG/ML	N87551	001	Feb 29, 1984
WARNER CHILCOTT	1MG/ML	N07085	002	
WATSON LABS	0.1MG/ML	N80573	002	
	0.1MG/ML	N83120	001	
	1MG/ML	N80573	001	
	1MG/ML	N83120	002	
WYETH AYERST	0.1MG/ML	N80554	001	
	1MG/ML	N80554	002	
REDISOL				
MERCK	1MG/ML	N06668	010	
RUBIVITE				
BEL MAR	0.03MG/ML	N10791	004	
	0.05MG/ML	N10791	001	
	0.1MG/ML	N10791	002	
	0.12MG/ML	N10791	005	
	1MG/ML	N10791	003	
RUBRAMIN PC				
BRISTOL MYERS SQUIBB	0.1MG/ML	N06799	002	
	1MG/ML	N06799	004	
	1MG/ML	N06799	010	Apr 28, 1988
RUVITE				
SAVAGE LABS	1MG/ML	N80570	002	
VI-TWEL				
BERLEX	1MG/ML	N07012	002	
TABLET; ORAL				
CYANOCOBALAMIN				
WEST WARD	1MG	N84264	001	

CYANOCOBALAMIN, CO-60

CAPSULE; ORAL				
RUBRATOPE-60				
BRACCO	0.5-1uCi	N16090	002	

CYANOCOBALAMIN; CYANOCOBALAMIN, CO-57; CYANOCOBALAMIN, CO-58

N/A; N/A				
DICOPAC KIT				
GE HEALTHCARE	N/A;N/A;N/A	N17406	001	

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 74 (of 274)

CYANOCOBALAMIN; CYANOCOBALAMIN, CO-57; INTRINSIC FACTOR

N/A; N/A

CYANOCOBALAMIN CO 57 SCHILLING TEST KIT

MALLINCKRODT

0.1MG;0.5uCi;60MG

N16635 001

CYANOCOBALAMIN; TANNIC ACID; ZINC ACETATE

INJECTABLE; INJECTION

DEPINAR

ARMOUR PHARM

0.5MG/ML;2.3MG/ML;1MG/ML

N11208 001

CYCLACILLIN

FOR SUSPENSION; ORAL

CYCLAPEN-W

WYETH AYERST

125MG/5ML

N50508 001

250MG/5ML

N50508 002

500MG/5ML

N50508 003

TABLET; ORAL

CYCLACILLIN

TEVA

250MG

N62895 001

Aug 04, 1988

500MG

N62895 002

Aug 04, 1988

CYCLAPEN-W

WYETH AYERST

250MG

N50509 001

500MG

N50509 002

CYCLIZINE LACTATE

INJECTABLE; INJECTION

MAREZINE

GLAXOSMITHKLINE

50MG/ML

N09495 001

CYCLOPENTOLATE HYDROCHLORIDE

SOLUTION/DROPS; OPHTHALMIC

AK-PENTOLATE

AKORN

1%

N85555 001

CYCLOPENTOLATE HYDROCHLORIDE

SOLA BARNES HIND

1%

N84150 001

1%

N84863 001

PENTOLAIR

PHARMAFAIR

0.5%

N88643 001

Feb 09, 1987

1%

N88150 001

Feb 25, 1983

CYCLOPHOSPHAMIDE

INJECTABLE; INJECTION

CYTOXAN

BRISTOL MYERS SQUIBB

100MG/VIAL

N12142 001

200MG/VIAL

N12142 002

500MG/VIAL

N12142 003

1GM/VIAL

N12142 004

Aug 30, 1982

2GM/VIAL

N12142 005

Aug 30, 1982

NEOSAR

SICOR PHARMS

100MG/VIAL

N40015 001

Apr 29, 1993

200MG/VIAL

N40015 002

Apr 29, 1993

500MG/VIAL

N40015 003

Apr 29, 1993

1GM/VIAL

N40015 004

Apr 29, 1993

2GM/VIAL

N40015 005

Apr 29, 1993

CYCLOSPORINE

CAPSULE; ORAL

NEORAL

NOVARTIS

50MG

N50715 003

Jul 14, 1995

DISCONTINUED DRUG PRODUCT LIST

6 - 75 (of 274)

CYCLOTHIAZIDE

TABLET; ORAL
ANHYDRON

LILLY 2MG

N13157 002

FLUIDIL

PHARMACIA AND UPJOHN 2MG

N18173 001

CYCRIMINE HYDROCHLORIDE

TABLET; ORAL
PAGITANE

LILLY 1.25MG
2.5MG

N08951 001

N08951 002

CYPROHEPTADINE HYDROCHLORIDE

SYRUP; ORAL

CYPROHEPTADINE HYDROCHLORIDE

HALSEY 2MG/5ML

N89199 001 Jul 03, 1986

MORTON GROVE 2MG/5ML

N87001 001 Nov 04, 1982

NASKA 2MG/5ML

N89021 001 Dec 21, 1987

PERIACTIN

MERCK 2MG/5ML

N13220 002

TABLET; ORAL

CYPROHEPTADINE HYDROCHLORIDE

AM THERAP 4MG

N88798 001 Feb 15, 1985

ASCOT 4MG

N87685 001 Oct 25, 1982

DURAMED PHARMS BARR 4MG

N88232 001 Oct 25, 1983

HALSEY 4MG

N89057 001 Jul 03, 1986

KV PHARM 4MG

N86737 001

MD PHARM 4MG

N87566 001 Nov 10, 1982

MYLAN 4MG

N86678 001

PIONEER PHARMS 4MG

N87839 001 Feb 08, 1984

SANDOZ 4MG

N86808 001

SUPERPHARM 4MG

N87405 001

TG UNITED LABS 4MG

N88212 001 May 26, 1983

VITARINE 4MG

N87284 001

WATSON LABS 4MG

N85245 001

4MG

N86165 001

4MG

N86580 001

PERIACTIN

MERCK 4MG

N12649 001

CYSTEINE HYDROCHLORIDE

INJECTABLE; INJECTION

CYSTEINE HYDROCHLORIDE

HOSPIRA 7.25%

N19523 001 Oct 22, 1986

CYTARABINE

INJECTABLE; INJECTION

CYTARABINE

SICOR PHARMS 100MG/VIAL

N16793 001

500MG/VIAL

N16793 002

1GM/VIAL

N16793 003 Dec 21, 1987

2GM/VIAL

N16793 004 Dec 21, 1987

DACARBAZINE

INJECTABLE; INJECTION

DACARBAZINE

ABRAXIS PHARM 100MG/VIAL

N70962 001 Aug 28, 1986

200MG/VIAL

N70990 001 Aug 28, 1986

DALFOPRISTIN; QUINUPRISTIN

INJECTABLE; IV (INFUSION)

SYNERCID

KING PHARMS 420MG/VIAL;180MG/VIAL

N50748 002 Aug 24, 2000

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 76 (of 274)

DALTEPARIN SODIUMINJECTABLE; INJECTION
FRAGMIN

PHARMACIA AND UPJOHN 7,500 IU/0.75ML

N20287 008 Apr 04, 2002

DANAPAROID SODIUMINJECTABLE; INJECTION
ORGARAN

ORGANON USA INC 750 UNITS/0.6ML

N20430 001 Dec 24, 1996

DANAZOLCAPSULE; ORAL
DANAZOL

AM THERAP 200MG

N71569 001 Dec 30, 1987

DANOCRINE

SANOFI AVENTIS US 50MG

N17557 003

100MG

N17557 004

200MG

N17557 002

DAPSONEGEL; TOPICAL
ACZONE

QLT USA 5%

N21794 001 Jul 07, 2005

DAPTOMYCININJECTABLE; IV (INFUSION)
CUBICIN

CUBIST 250MG/VIAL

N21572 001 Sep 12, 2003

DAUNORUBICIN HYDROCHLORIDEINJECTABLE; INJECTION
CERUBIDINE

SANOFI AVENTIS US EQ 20MG BASE/VIAL

N61876 001

WYETH AYERST EQ 20MG BASE/VIAL

N50484 001

DECAMETHONIUM BROMIDEINJECTABLE; INJECTION
SYNCURINE

GLAXOSMITHKLINE 1MG/ML

N06931 002

DEMECARIUM BROMIDESOLUTION/DROPS; OPHTHALMIC
HUMORSOL

MERCK 0.125%

N11860 002

0.25%

N11860 001

DEMECLOCYCLINE HYDROCHLORIDECAPSULE; ORAL
DECLOMYCIN

LEDERLE 150MG

N50262 001

SYRUP; ORAL
DECLOMYCIN

LEDERLE 75MG/5ML

N50257 001

TABLET; ORAL
DECLOMYCIN

GLADES PHARMS LLC 75MG

N50261 001

DESERPIDINETABLET; ORAL
HARMONYL

ABBOTT 0.1MG

N10796 001

0.25MG

N10796 002

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 77 (of 274)

DESERPIDINE; HYDROCHLOROTHIAZIDE

TABLET; ORAL			
ORETICYL 25			
ABBOTT	0.125MG;25MG	N12148	001
ORETICYL 50			
ABBOTT	0.125MG;50MG	N12148	003
ORETICYL FORTE			
ABBOTT	0.25MG;25MG	N12148	002

DESERPIDINE; METHYCLOTHIAZIDE

TABLET; ORAL			
ENDURONYL			
ABBOTT	0.25MG;5MG	N12775	001
ENDURONYL FORTE			
ABBOTT	0.5MG;5MG	N12775	002
METHYCLOTHIAZIDE AND DESERPIDINE			
WATSON LABS	0.25MG;5MG	N88486	001 Aug 10, 1984
	0.5MG;5MG	N88452	001 Aug 10, 1984

DESIPRAMINE HYDROCHLORIDE

CAPSULE; ORAL			
PERTOFRANE			
SANOFI AVENTIS US	25MG	N13621	001
	50MG	N13621	002
TABLET; ORAL			
DESIPRAMINE HYDROCHLORIDE			
USL PHARMA	25MG	N71864	001 Sep 09, 1987
	50MG	N71865	001 Sep 09, 1987
	75MG	N71866	001 Sep 09, 1987
	100MG	N71867	001 Sep 09, 1987

DESLANOSIDE

INJECTABLE; INJECTION			
CEDILANID-D			
NOVARTIS	0.2MG/ML	N09282	002

DESMOPRESSIN ACETATE

INJECTABLE; INJECTION			
DESMOPRESSIN ACETATE			
BEDFORD	0.004MG/ML	N74575	001 Feb 18, 2000
DESMOPRESSIN ACETATE PRESERVATIVE FREE			
BEDFORD	0.004MG/ML	N74574	001 Feb 18, 2000
SOLUTION; NASAL			
CONCENTRAID			
FERRING	0.01%	N19776	001 Dec 26, 1990
SPRAY, METERED; NASAL			
DDAVP			
SANOFI AVENTIS US	0.01MG/SPRAY	N17922	002 Feb 06, 1989

DESOGESTREL; ETHINYL ESTRADIOL

TABLET; ORAL-21			
DESOGEN			
ORGANON USA INC	0.15MG;0.03MG	N20071	001 Dec 10, 1992
ORTHO-CEPT			
ORTHO MCNEIL PHARM	0.15MG;0.03MG	N20301	001 Dec 14, 1992

DESOXIMETASONE

OINTMENT; TOPICAL			
DESOXIMETASONE			
ALTANA	0.25%	N73440	001 Apr 01, 1998
TOPICORT			
TARO PHARMS NORTH	0.05%	N18594	001 Jan 17, 1985

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 78 (of 274)

DESOXYCORTICOSTERONE ACETATEINJECTABLE; INJECTION
DOCA

ORGANON USA INC 5MG/ML N01104 001

PELLET; IMPLANTATION
PERCORTEN

NOVARTIS 125MG N05151 001

DESOXYCORTICOSTERONE PIVALATEINJECTABLE; INJECTION
PERCORTEN

NOVARTIS 25MG/ML N08822 001

DEXAMETHASONE

AEROSOL; TOPICAL

AEROSEB-DEX

ALLERGAN HERBERT 0.01% N83296 002

DECASPRAY

MERCK 0.04% N12731 002

ELIXIR; ORAL

DECADRON

MERCK 0.5MG/5ML N12376 002

DEXAMETHASONE

ALPHARMA US PHARMS 0.5MG/5ML N88997 001 Oct 10, 1986

HEXADRÖL

ORGANON USA INC 0.5MG/5ML N12674 001

GEL; TOPICAL

DECADERM

MERCK 0.1% N13538 001

SUSPENSION/DROPS; OPHTHALMIC

DEXAMETHASONE

WATSON LABS 0.1% N89170 001 May 09, 1989

TABLET; ORAL

DECADRON

MERCK 0.25MG N11664 004

0.5MG N11664 001

0.75MG N11664 002

1.5MG N11664 003

4MG N11664 005

6MG N11664 006 Jul 30, 1982

DEXAMETHASONE

IMPAX LABS 0.75MG N85376 001

LEINER 0.75MG N83420 001

MUTUAL PHARM 0.25MG N84013 001

0.25MG N84764 001

0.5MG N84084 001

0.5MG N84766 001

0.75MG N84081 001

0.75MG N84765 001

1.5MG N84086 001

1.5MG N84763 001

PHOENIX LABS NY 0.75MG N83806 001

ROXANE 0.25MG N84614 001

SANDOZ 0.75MG N80399 001

UPSHER SMITH 0.75MG N87534 001

1.5MG N87533 001

WATSON LABS 0.25MG N85455 001

0.5MG N85458 001

0.75MG N80968 001

0.75MG N84457 001

0.75MG N85818 001

1.5MG N85456 001

1.5MG N85840 001

WHITEWORTH TOWN PLSN 0.75MG N84327 001

DISCONTINUED DRUG PRODUCT LIST

6 - 79 (of 274)

DEXAMETHASONE

TABLET; ORAL

DEXONE 0.5

SOLVAY

0.5MG

N84991 001

DEXONE 0.75

SOLVAY

0.75MG

N84993 001

DEXONE 1.5

SOLVAY

1.5MG

N84990 001

DEXONE 4

SOLVAY

4MG

N84992 001

HEXADROL

ORGANON USA INC

0.5MG

N12675 004

0.75MG

N12675 007

1.5MG

N12675 009

4MG

N12675 010

DEXAMETHASONE ACETATE

INJECTABLE; INJECTION

DECADRON-LA

MERCK

EQ 8MG BASE/ML **Federal Register
determination that product was not
discontinued or withdrawn for safety
or efficacy reasons**

N16675 001

DEXAMETHASONE ACETATE

WATSON LABS

EQ 8MG BASE/ML

N84315 001

EQ 16MG BASE/ML

N87711 001

May 24, 1982

DEXAMETHASONE SODIUM PHOSPHATE

AEROSOL; NASAL

DEXACORT

UCB INC

EQ 0.1MG PHOSPHATE/INH

N14242 001

AEROSOL, METERED; INHALATION

DEXACORT

UCB INC

EQ 0.1MG PHOSPHATE/INH

N13413 001

CREAM; TOPICAL

DECADRON

MERCK

EQ 0.1% PHOSPHATE

N11983 002

INJECTABLE; INJECTION

DECADRON

MERCK

EQ 4MG PHOSPHATE/ML

N12071 002

EQ 24MG PHOSPHATE/ML **Federal
Register determination that product
was not discontinued or withdrawn for
safety or efficacy reasons**

N12071 004

DEXACEN-4

CENT PHARMS

EQ 4MG PHOSPHATE/ML

N84342 001

DEXAMETHASONE

ABRAXIS PHARM

EQ 4MG PHOSPHATE/ML

N88448 001

EQ 10MG PHOSPHATE/ML

N88469 001

Jan 25, 1984

Jan 25, 1984

DEXAMETHASONE SODIUM PHOSPHATE

ABRAXIS PHARM

EQ 4MG PHOSPHATE/ML

N87065 001

AKORN

EQ 4MG PHOSPHATE/ML

N84493 001

BEL MAR

EQ 4MG PHOSPHATE/ML

N84752 001

DELL LABS

EQ 4MG PHOSPHATE/ML

N83161 001

INTL MEDICATION

EQ 20MG PHOSPHATE/ML

N88522 001

Feb 17, 1984

SICOR PHARMS

EQ 4MG PHOSPHATE/ML

N81125 001

Aug 31, 1990

WATSON LABS

EQ 4MG PHOSPHATE/ML

N83702 001

EQ 4MG PHOSPHATE/ML

N84355 001

EQ 4MG PHOSPHATE/ML

N89169 001

Apr 09, 1986

EQ 10MG PHOSPHATE/ML

N87668 001

Jul 01, 1982

EQ 24MG PHOSPHATE/ML

N85606 001

WYETH AYERST

EQ 4MG PHOSPHATE/ML

N85641 001

HEXADROL

ORGANON USA INC

EQ 4MG PHOSPHATE/ML

N14694 002

EQ 10MG PHOSPHATE/ML

N14694 003

DISCONTINUED DRUG PRODUCT LIST

6 - 80 (of 274)

DEXAMETHASONE SODIUM PHOSPHATE

INJECTABLE; INJECTION				
HEXADROL				
ORGANON USA INC	EQ 20MG PHOSPHATE/ML	N14694	004	
OINTMENT; OPHTHALMIC				
DECADRON				
MERCK	EQ 0.05% PHOSPHATE	N11977	001	
DEXAIR				
PHARMAFAIR	EQ 0.05% PHOSPHATE	N88071	001	Dec 28, 1982
MAXIDEX				
ALCON	EQ 0.05% PHOSPHATE	N83342	001	
SOLUTION/DROPS; OPHTHALMIC				
DEXAIR				
PHARMAFAIR	EQ 0.1% PHOSPHATE	N88433	001	Dec 15, 1983
DEXAMETHASONE SODIUM PHOSPHATE				
SOLA BARNES HIND	EQ 0.1% PHOSPHATE	N84170	001	
	EQ 0.1% PHOSPHATE	N84173	001	
SOLUTION/DROPS; OPHTHALMIC, OTIC				
DECADRON				
MERCK	EQ 0.1% PHOSPHATE	N11984	001	
SOLUTION/DROPS; OTIC				
DEXAMETHASONE SODIUM PHOSPHATE				
AKORN	EQ 0.1% PHOSPHATE	N84855	001	

DEXAMETHASONE SODIUM PHOSPHATE; LIDOCAINE HYDROCHLORIDE

INJECTABLE; INJECTION				
DECADRON W/ XYLOCAINE				
MERCK	EQ 4MG PHOSPHATE/ML;10MG/ML	N13334	002	

DEXAMETHASONE SODIUM PHOSPHATE; NEOMYCIN SULFATE

OINTMENT; OPHTHALMIC				
NEODECADRON				
MERCK	EQ 0.05% PHOSPHATE;EQ 3.5MG BASE/GM	N50324	001	
SOLUTION/DROPS; OPHTHALMIC				
NEODECADRON				
MERCK	EQ 0.1% PHOSPHATE;EQ 3.5MG BASE/ML	N50322	001	
NEOMYCIN SULFATE AND DEXAMETHASONE SODIUM PHOSPHATE				
BAUSCH AND LOMB	EQ 0.1% PHOSPHATE;EQ 3.5MG BASE/ML	N64055	001	Oct 30, 1995
NEOMYCIN SULFATE-DEXAMETHASONE SODIUM PHOSPHATE				
ALCON UNIVERSAL	EQ 0.1% PHOSPHATE;EQ 3.5MG BASE/ML	N62714	001	Jul 21, 1986
PHARMAFAIR	EQ 0.1% PHOSPHATE;EQ 3.5MG BASE/ML	N62539	001	Jan 10, 1985

DEXAMETHASONE; NEOMYCIN SULFATE; POLYMYXIN B SULFATE

OINTMENT; OPHTHALMIC				
DEXACIDIN				
NOVARTIS	0.1%;EQ 3.5MG BASE/GM;10,000 UNITS/GM	N62566	001	Feb 22, 1985
DEXASPORIN				
PHARMAFAIR	0.1%;EQ 3.5MG BASE/GM;10,000 UNITS/GM	N62411	001	May 16, 1983
SUSPENSION/DROPS; OPHTHALMIC				
DEXACIDIN				
NOVARTIS	0.1%;EQ 3.5MG BASE/ML;10,000 UNITS/ML	N62544	001	Oct 29, 1984
DEXASPORIN				
PHARMAFAIR	0.1%;EQ 3.5MG BASE/ML;10,000 UNITS/ML	N62428	001	May 18, 1983

DEXBROMPHENIRAMINE MALEATE

SYRUP; ORAL				
DISOMER				
SCHERING	2MG/5ML	N11814	002	
TABLET; ORAL				
DISOMER				
SCHERING	2MG	N11814	001	

DISCONTINUED DRUG PRODUCT LIST

6 - 81 (of 274)

DEXBROMPHENIRAMINE MALEATE; PSEUDOEPHEDRINE SULFATE

TABLET; ORAL			
DISOPHROL			
SCHERING	2MG;60MG	N12394	002
TABLET, EXTENDED RELEASE; ORAL			
BROMPHERIL			
COPLEY PHARM	6MG;120MG	N89116	001 Jan 22, 1987
DISOBROM			
SANDOZ	6MG;120MG	N70770	001 Sep 30, 1991
RESPORAL			
PIONEER PHARMS	6MG;120MG	N89139	001 Jun 16, 1988

DEXCHLORPHENIRAMINE MALEATE

SYRUP; ORAL			
POLARAMINE			
SCHERING	2MG/5ML	N86837	001 Jul 19, 1982
TABLET; ORAL			
POLARAMINE			
SCHERING	2MG	N86835	001

DEXTROAMPHETAMINE SULFATE

CAPSULE; ORAL			
DEXAMPEX			
TEVA	15MG	N85355	001
CAPSULE, EXTENDED RELEASE; ORAL			
DEXTROAMPHETAMINE SULFATE			
ABLE	5MG	N76814	001 Aug 25, 2004
	10MG	N76814	002 Aug 25, 2004
	15MG	N76814	003 Aug 25, 2004
ELIXIR; ORAL			
DEXEDRINE			
GLAXOSMITHKLINE	5MG/5ML	N83902	001
TABLET; ORAL			
DEXAMPEX			
TEVA	5MG	N83735	001
	10MG	N83735	002
DEXTROAMPHETAMINE SULFATE			
ENDO PHARMS	5MG	N40299	001 May 13, 1999
HALSEY	10MG	N83930	001
LANNETT	5MG	N83903	001
	10MG	N83903	003
	15MG **Federal Register determination that product was not discontinued or withdrawn for safety or efficacy reasons**	N85652	001
MAST MM	5MG	N86521	001
PUREPAC PHARM	5MG	N84125	001
SANDOZ	5MG	N85370	001
	10MG	N85371	001
VITARINE	5MG	N84986	001
	10MG	N85892	001
FERNDEX			
FERNDALE LABS	5MG	N84001	001

DEXTROMETHORPHAN HYDROBROMIDE; PROMETHAZINE HYDROCHLORIDE

SYRUP; ORAL			
PHENERGAN W/ DEXTROMETHORPHAN			
WYETH AYERST	15MG/5ML;6.25MG/5ML	N11265	002 Apr 02, 1984
PHERAZINE DM			
HALSEY	15MG/5ML;6.25MG/5ML	N88913	001 Mar 02, 1987

DEXTROSE

INJECTABLE; INJECTION			
DEXTROSE 10% IN PLASTIC CONTAINER			
B BRAUN	10GM/100ML	N18046	001

DISCONTINUED DRUG PRODUCT LIST

6 - 82 (of 274)

DEXTROSE

INJECTABLE; INJECTION

DEXTROSE 10% IN PLASTIC CONTAINER			
MILES	10GM/100ML	N18504	001
DEXTROSE 2.5% IN PLASTIC CONTAINER			
B BRAUN	2.5GM/100ML	N18358	001
	2.5GM/100ML	N19626	001
			Feb 02, 1988
DEXTROSE 38.5% IN PLASTIC CONTAINER			
ABBOTT	38.5GM/100ML	N18923	001
			Sep 19, 1984
DEXTROSE 60%			
B BRAUN	60GM/100ML	N17995	002
			Sep 22, 1982
DEXTROSE 60% IN PLASTIC CONTAINER			
B BRAUN	60GM/100ML	N17995	001
BAXTER HLTHCARE	60GM/100ML	N20047	002
			Jul 02, 1991
DEXTROSE 7.7% IN PLASTIC CONTAINER			
B BRAUN	7.7GM/100ML	N19626	003
			Feb 02, 1988

DEXTROSE; MAGNESIUM CHLORIDE; POTASSIUM CHLORIDE; POTASSIUM PHOSPHATE, DIBASIC; SODIUM ACETATE

INJECTABLE; INJECTION

ISOLYTE P W/ DEXTROSE 5% IN PLASTIC CONTAINER			
B BRAUN	5GM/100ML;31MG/100ML;130MG/100ML;26MG/100ML;320MG/100ML	N19025	001
			Dec 27, 1984

DEXTROSE; MAGNESIUM CHLORIDE; POTASSIUM CHLORIDE; SODIUM ACETATE; SODIUM CHLORIDE

INJECTABLE; INJECTION

ISOLYTE H W/ DEXTROSE 5% IN PLASTIC CONTAINER			
B BRAUN	5GM/100ML;30MG/100ML;97MG/100ML;220MG/100ML;140MG/100ML	N18273	001

DEXTROSE; MAGNESIUM CHLORIDE; POTASSIUM CHLORIDE; SODIUM ACETATE; SODIUM CHLORIDE; SODIUM GLUCONATE

INJECTABLE; INJECTION

ISOLYTE S W/ DEXTROSE 5% IN PLASTIC CONTAINER			
B BRAUN	5GM/100ML;30MG/100ML;37MG/100ML;370MG/100ML;530MG/100ML;500MG/100ML	N18274	001

DEXTROSE; POTASSIUM CHLORIDE

INJECTABLE; INJECTION

POTASSIUM CHLORIDE 0.037% IN DEXTROSE 5% IN PLASTIC CONTAINER			
B BRAUN	5GM/100ML;37MG/100ML	N19699	001
			Sep 29, 1989
POTASSIUM CHLORIDE 0.075% IN DEXTROSE 5% IN PLASTIC CONTAINER			
B BRAUN	5GM/100ML;75MG/100ML	N19699	002
			Sep 29, 1989
POTASSIUM CHLORIDE 0.11% IN DEXTROSE 5% IN PLASTIC CONTAINER			
B BRAUN	5GM/100ML;110MG/100ML	N19699	003
			Sep 29, 1989
POTASSIUM CHLORIDE 0.22% IN DEXTROSE 5% IN PLASTIC CONTAINER			
B BRAUN	5GM/100ML;220MG/100ML	N19699	005
			Sep 29, 1989

DEXTROSE; POTASSIUM CHLORIDE; POTASSIUM PHOSPHATE, DIBASIC; SODIUM ACETATE; SODIUM CHLORIDE

INJECTABLE; INJECTION

ISOLYTE M W/ DEXTROSE 5% IN PLASTIC CONTAINER			
B BRAUN	5GM/100ML;150MG/100ML;130MG/100ML;280MG/100ML;91MG/100ML	N18270	001

DEXTROSE; POTASSIUM CHLORIDE; SODIUM CHLORIDE

INJECTABLE; INJECTION

DEXTROSE 5%, SODIUM CHLORIDE 0.2% AND POTASSIUM CHLORIDE 0.075%			
B BRAUN	5GM/100ML;75MG/100ML;200MG/100ML	N18268	009
DEXTROSE 5%, SODIUM CHLORIDE 0.2% AND POTASSIUM CHLORIDE 0.15% IN PLASTIC CONTAINER			
B BRAUN	5GM/100ML;150MG/100ML;200MG/100ML	N18268	004
DEXTROSE 5%, SODIUM CHLORIDE 0.2% AND POTASSIUM CHLORIDE 0.224% IN PLASTIC CONTAINER			
B BRAUN	5GM/100ML;220MG/100ML;200MG/100ML	N18268	005
DEXTROSE 5%, SODIUM CHLORIDE 0.2% AND POTASSIUM CHLORIDE 0.3% IN PLASTIC CONTAINER			
B BRAUN	5GM/100ML;300MG/100ML;200MG/100ML	N18268	006

DISCONTINUED DRUG PRODUCT LIST

6 - 83 (of 274)

DEXTROSE; POTASSIUM CHLORIDE; SODIUM CHLORIDE

INJECTABLE; INJECTION

DEXTROSE 5%, SODIUM CHLORIDE 0.33% AND POTASSIUM CHLORIDE 0.075% IN PLASTIC CONTAINER			
B BRAUN	5GM/100ML;75MG/100ML;330MG/100ML	N18268 011	Jan 18, 1986
DEXTROSE 5%, SODIUM CHLORIDE 0.33% AND POTASSIUM CHLORIDE 0.15% IN PLASTIC CONTAINER			
B BRAUN	5GM/100ML;150MG/100ML;330MG/100ML	N18268 012	Jan 18, 1986
DEXTROSE 5%, SODIUM CHLORIDE 0.33% AND POTASSIUM CHLORIDE 0.22% IN PLASTIC CONTAINER			
B BRAUN	5GM/100ML;220MG/100ML;330MG/100ML	N18268 013	Jan 18, 1986
DEXTROSE 5%, SODIUM CHLORIDE 0.33% AND POTASSIUM CHLORIDE 0.30% IN PLASTIC CONTAINER			
B BRAUN	5GM/100ML;300MG/100ML;330MG/100ML	N18268 014	Jan 18, 1986
DEXTROSE 5%, SODIUM CHLORIDE 0.45% AND POTASSIUM CHLORIDE 0.075%			
B BRAUN	5GM/100ML;75MG/100ML;450MG/100ML	N18268 010	
DEXTROSE 5%, SODIUM CHLORIDE 0.45% AND POTASSIUM CHLORIDE 0.15% IN PLASTIC CONTAINER			
B BRAUN	5GM/100ML;150MG/100ML;450MG/100ML	N18268 001	
DEXTROSE 5%, SODIUM CHLORIDE 0.45% AND POTASSIUM CHLORIDE 0.22% IN PLASTIC CONTAINER			
B BRAUN	5GM/100ML;220MG/100ML;450MG/100ML	N18268 002	
DEXTROSE 5%, SODIUM CHLORIDE 0.45% AND POTASSIUM CHLORIDE 0.3% IN PLASTIC CONTAINER			
B BRAUN	5GM/100ML;300MG/100ML;450MG/100ML	N18268 003	
DEXTROSE 5%, SODIUM CHLORIDE 0.45% AND POTASSIUM CHLORIDE 15MEQ IN PLASTIC CONTAINER			
BAXTER HLTHCARE	5GM/100ML;224MG/100ML;450MG/100ML	N18008 003	
DEXTROSE 5%, SODIUM CHLORIDE 0.45% AND POTASSIUM CHLORIDE 20MEQ (K) IN PLASTIC CONTAINER			
BAXTER HLTHCARE	5GM/100ML;300MG/100ML;450MG/100ML	N18008 001	
DEXTROSE 5%, SODIUM CHLORIDE 0.45% AND POTASSIUM CHLORIDE 5MEQ IN PLASTIC CONTAINER			
BAXTER HLTHCARE	5GM/100ML;75MG/100ML;450MG/100ML	N18008 002	

DEXTROSE; SODIUM CHLORIDE

INJECTABLE; INJECTION

DEXTROSE 10% AND SODIUM CHLORIDE 0.2% IN PLASTIC CONTAINER			
B BRAUN	10GM/100ML;200MG/100ML	N18386 001	
DEXTROSE 10% AND SODIUM CHLORIDE 0.45% IN PLASTIC CONTAINER			
B BRAUN	10GM/100ML;450MG/100ML	N18229 001	
DEXTROSE 10% AND SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER			
B BRAUN	10GM/100ML;900MG/100ML	N18047 001	
DEXTROSE 2.5% AND SODIUM CHLORIDE 0.45% IN PLASTIC CONTAINER			
B BRAUN	2.5GM/100ML;450MG/100ML	N18030 001	
DEXTROSE 2.5% AND SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER			
B BRAUN	2.5GM/100ML;900MG/100ML	N18376 001	
DEXTROSE 3.3% AND SODIUM CHLORIDE 0.3% IN PLASTIC CONTAINER			
ABBOTT	3.3GM/100ML;300MG/100ML	N18055 001	
DEXTROSE 5% AND SODIUM CHLORIDE 0.11% IN PLASTIC CONTAINER			
B BRAUN	5GM/100ML;110MG/100ML	N18030 005	
DEXTROSE 5% AND SODIUM CHLORIDE 0.2% IN PLASTIC CONTAINER			
B BRAUN	5GM/100ML;200MG/100ML	N18030 004	
MILES	5GM/100ML;200MG/100ML	N18399 001	
DEXTROSE 5% AND SODIUM CHLORIDE 0.225% IN PLASTIC CONTAINER			
ABBOTT	5GM/100ML;225MG/100ML	N19482 001	Oct 04, 1985
DEXTROSE 5% AND SODIUM CHLORIDE 0.3% IN PLASTIC CONTAINER			
ABBOTT	5GM/100ML;300MG/100ML	N19486 001	Oct 04, 1985
MILES	5GM/100ML;300MG/100ML	N18501 001	
DEXTROSE 5% AND SODIUM CHLORIDE 0.33% IN PLASTIC CONTAINER			
B BRAUN	5GM/100ML;330MG/100ML	N18030 003	
DEXTROSE 5% AND SODIUM CHLORIDE 0.45% IN PLASTIC CONTAINER			
ABBOTT	5GM/100ML;450MG/100ML	N19484 001	Oct 04, 1985
B BRAUN	5GM/100ML;450MG/100ML	N18030 002	
MILES	5GM/100ML;450MG/100ML	N18400 001	
DEXTROSE 5% AND SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER			
ABBOTT	5GM/100ML;900MG/100ML	N19483 001	Oct 04, 1985
B BRAUN	5GM/100ML;900MG/100ML	N18026 001	
MILES	5GM/100ML;900MG/100ML	N18500 001	

DEXTROTHYROXINE SODIUM

TABLET; ORAL

CHOLOXIN			
ABBOTT	1MG	N12302 005	

DISCONTINUED DRUG PRODUCT LIST

6 - 84 (of 274)

DEXTROTHYROXINE SODIUM

TABLET; ORAL			
CHOLOXIN			
ABBOTT	2MG	N12302	002
	4MG	N12302	004
	6MG	N12302	006

DEZOCINE

INJECTABLE; INJECTION			
DALGAN			
ASTRAZENECA	5MG/ML	N19082	001 Dec 29, 1989
	10MG/ML	N19082	002 Dec 29, 1989
	15MG/ML	N19082	003 Dec 29, 1989

DIATRIZOATE MEGLUMINE

INJECTABLE; INJECTION			
ANGIOVIST 282			
BERLEX	60%	N87726	001 Sep 23, 1982
CARDIOGRAFIN			
BRACCO	85%	N11620	002
DIATRIZOATE MEGLUMINE			
BRACCO	76%	N10040	017
HYPaque			
GE HEALTHCARE	30%	N16403	002
UROVIST MEGLUMINE DIU/CT			
BERLEX	30%	N87739	001 Sep 23, 1982
SOLUTION; URETERAL			
UROVIST CYSTO			
BERLEX	30%	N87729	001 Sep 23, 1982
UROVIST CYSTO PEDIATRIC			
BERLEX	30%	N87731	001 Sep 23, 1982

DIATRIZOATE MEGLUMINE; DIATRIZOATE SODIUM

INJECTABLE; INJECTION			
ANGIOVIST 292			
BERLEX	52%;8%	N87724	001 Sep 23, 1982
ANGIOVIST 370			
BERLEX	66%;10%	N87723	001 Sep 23, 1982
DIATRIZOATE-60			
INTL MEDICATION	52%;8%	N88166	001 Jun 17, 1983
HYPaque-M, 75%			
GE HEALTHCARE	50%;25%	N10220	003
HYPaque-M, 90%			
GE HEALTHCARE	60%;30%	N10220	002
MD-60			
MALLINCKRODT	52%;8%	N87074	001
MD-76			
MALLINCKRODT	66%;10%	N87073	001
RENOVIST			
BRACCO	34.3%;35%	N10040	020
RENOVIST II			
BRACCO	28.5%;29.1%	N10040	019
SOLUTION; ORAL, RECTAL			
GASTROVIST			
BERLEX	66%;10%	N87728	001 Sep 23, 1982

DIATRIZOATE SODIUM

INJECTABLE; INJECTION			
HYPaque			
GE HEALTHCARE	25%	N09561	003
MD-50			
MALLINCKRODT	50%	N87075	001
UROVIST SODIUM 300			
BERLEX	50%	N87725	001 Sep 23, 1982

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 85 (of 274)

DIATRIZOATE SODIUMSOLUTION; ORAL, RECTAL
HYPAQUE

GE HEALTHCARE 40%

N11386 003

SOLUTION; URETERAL
HYPAQUE SODIUM 20%

GE HEALTHCARE 20%

N09561 002

DIAZEPAMCAPSULE, EXTENDED RELEASE; ORAL
VALRELEASE

ROCHE 15MG

N18179 001

GEL; RECTAL
DIASTATVALEANT 5MG/ML (5MG/ML)
10MG/2ML (5MG/ML)
15MG/3ML (5MG/ML)
20MG/4ML (5MG/ML)N20648 002 Jul 29, 1997
N20648 003 Jul 29, 1997
N20648 004 Jul 29, 1997
N20648 005 Jul 29, 1997INJECTABLE; INJECTION
DIAZEPAM

ABRAXIS PHARM 5MG/ML

N70662 001 Jun 25, 1986

BAXTER HLTHCARE 5MG/ML

N70311 001 Dec 16, 1985

5MG/ML

N70312 001 Dec 16, 1985

5MG/ML

N70313 001 Dec 16, 1985

5MG/ML

N71308 001 Jul 17, 1987

MARSAM PHARMS LLC 5MG/ML

N72371 001 Jan 29, 1993

US ARMY 5MG/ML

N20124 001 Dec 05, 1990

WARNER CHILCOTT 5MG/ML

N71613 001 Oct 22, 1987

5MG/ML

N71614 001 Oct 22, 1987

WATSON LABS 5MG/ML

N70911 001 Aug 28, 1986

5MG/ML

N70912 001 Aug 28, 1986

5MG/ML

N70930 001 Dec 01, 1986

DIZAC

PHARMACIA AND UPJOHN 5MG/ML

N19287 001 Jun 18, 1993

VALIUM

ROCHE 5MG/ML

N16087 001

TABLET; ORAL
DIAZEPAM

CLONMEL HLTHCARE 5MG

N70227 001 Sep 26, 1985

10MG

N70228 001 Sep 26, 1985

DURAMED PHARMS BARR 2MG

N70894 001 Aug 27, 1986

5MG

N70895 001 Aug 27, 1986

10MG

N70896 001 Aug 27, 1986

FERNDAL LABS 2MG

N70903 001 Apr 01, 1987

5MG

N70904 001 Apr 01, 1987

10MG

N70905 001 Apr 01, 1987

HALSEY 2MG

N70987 001 Aug 15, 1986

5MG

N70996 001 Aug 15, 1986

10MG

N70956 001 Aug 15, 1986

IVAX PHARMS 2MG

N70360 001 Sep 04, 1985

5MG

N70361 001 Sep 04, 1985

10MG

N70362 001 Sep 04, 1985

MARTEC USA LLC 10MG

N72402 001 Apr 25, 1989

PAR PHARM 10MG

N70464 001 Feb 25, 1986

PIONEER PHARMS 2MG

N70787 001 Aug 02, 1988

5MG

N70788 001 Aug 02, 1988

10MG

N70776 001 Aug 02, 1988

ROXANE 2MG

N70356 001 Jun 17, 1986

5MG

N70357 001 Jun 17, 1986

10MG

N70358 001 Jun 17, 1986

WARNER CHILCOTT 2MG

N70209 001 Sep 04, 1985

5MG

N70210 001 Sep 04, 1985

10MG

N70222 001 Sep 04, 1985

WATSON LABS 2MG

N70456 001 Nov 01, 1985

5MG

N70457 001 Nov 01, 1985

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 86 (of 274)

DIAZEPAMTABLET; ORAL
DIAZEPAM

WATSON LABS

10MG

N70458 001

Nov 01, 1985

Q-PAM

QUANTUM PHARMICS

2MG

N70423 001

Dec 12, 1985

2MG

N72431 001

Apr 29, 1988

5MG

N70424 001

Dec 12, 1985

5MG

N72432 001

Apr 29, 1988

10MG

N70425 001

Dec 12, 1985

10MG

N72433 001

Apr 29, 1988

DIAZOXIDECAPSULE; ORAL
PROGLYCEM

BAKER NORTON

50MG

N17425 001

100MG

N17425 002

INJECTABLE; INJECTION
DIAZOXIDE

ABRAXIS PHARM

15MG/ML

N71519 001

Aug 26, 1987

HYPERSTAT

SCHERING

15MG/ML

N16996 001

DIBUCAINE HYDROCHLORIDEINJECTABLE; INJECTION
HEAVY SOLUTION NUPERCALINE

NOVARTIS

2.5MG/ML

N06203 001

DICHLORPHENAMIDETABLET; ORAL
DARANIDE

MERCK

50MG

N11366 001

DICLOFENAC POTASSIUMTABLET; ORAL
CATAFLAM

NOVARTIS

25MG **Federal Register determination
that product was not discontinued or
withdrawn for safety or efficacy
reasons**

N20142 001

Nov 24, 1993

DICLOFENAC SODIUMSOLUTION/DROPS; OPHTHALMIC
DICLOFENAC SODIUM

FALCON PHARMS

0.1%

N20809 001

May 04, 1998

TABLET, DELAYED RELEASE; ORAL
VOLTAREN

NOVARTIS

25MG

N19201 001

Jul 28, 1988

50MG

N19201 002

Jul 28, 1988

DICLOXACILLIN SODIUMCAPSULE; ORAL
DYCILL

GLAXOSMITHKLINE

EQ 250MG BASE

N60254 002

EQ 250MG BASE

N62238 001

EQ 500MG BASE

N60254 003

EQ 500MG BASE

N62238 002

PATHOCIL

WYETH AYERST

EQ 250MG BASE

N50011 002

EQ 500MG BASE

N50011 003

Mar 28, 1983

FOR SUSPENSION; ORAL
DYNAPEN

APOTHECON

EQ 62.5MG BASE/5ML

N50337 002

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 87 (of 274)

DICLOXACILLIN SODIUMFOR SUSPENSION; ORAL
PATHOCIL

WYETH AYERST EQ 62.5MG BASE/5ML N50092 001

DICUMAROLCAPSULE; ORAL
DICUMAROLLILLY 25MG N05509 003
50MG N05509 001TABLET; ORAL
DICUMAROLABBOTT 25MG N05545 003
50MG N05545 004
100MG N05545 005DICYCLOMINE HYDROCHLORIDECAPSULE; ORAL
DICYCLOMINE HYDROCHLORIDEMUTUAL PHARM 10MG N84505 001 Oct 21, 1986
PIONEER PHARMS 10MG N89361 001 Jan 10, 1989
WATSON LABS 10MG N83179 001 Feb 12, 1986

INJECTABLE; INJECTION

DICYCLOMINE HYDROCHLORIDE

WATSON LABS 10MG/ML N80614 001 Feb 11, 1986

SYRUP; ORAL

DICYCLOMINE HYDROCHLORIDE

ALPHARMA US PHARMS 10MG/5ML N84479 001

TABLET; ORAL

DICYCLOMINE HYDROCHLORIDE

MUTUAL PHARM 20MG N84600 001 Jul 29, 1985
PIONEER PHARMS 20MG N88585 001 Aug 20, 1986
WATSON LABS 20MG N84361 001 Feb 06, 1986DIDANOSINEFOR SOLUTION; ORAL
VIDEXBRISTOL MYERS SQUIBB 100MG/PACKET N20155 003 Oct 09, 1991
167MG/PACKET N20155 004 Oct 09, 1991
250MG/PACKET N20155 005 Oct 09, 1991
375MG/PACKET N20155 006 Oct 09, 1991DIENESTROL

CREAM; VAGINAL

DIENESTROL

ORTHO MCNEIL PHARM 0.01% N06110 005

DV

SANOFI AVENTIS US 0.01% N83518 001

ESTRAGUARD

SOLVAY 0.01% N84436 001

SUPPOSITORY; VAGINAL

DV

SANOFI AVENTIS US 0.7MG N83517 001

DIETHYLCARBAMAZINE CITRATE

TABLET; ORAL

HETRAZAN

LEDERLE 50MG N06459 001

DIETHYLPROPION HYDROCHLORIDE

TABLET; ORAL

DIETHYLPROPION HYDROCHLORIDE

SANDOZ 25MG N85916 001

TEVA 25MG N88642 001 Sep 20, 1984

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 88 (of 274)

DIETHYLPROPION HYDROCHLORIDE

TABLET; ORAL

DIETHYLPROPION HYDROCHLORIDE

TG UNITED LABS	25MG	N88267	001	Aug 25, 1983
	25MG	N88268	001	Aug 25, 1983
UCB INC	25MG	N85544	001	
WATSON LABS	25MG	N85741	001	

TENUATE

SANOFI AVENTIS US	25MG	N17668	001	
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TEPANIL

3M	25MG	N11673	001	
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TABLET, EXTENDED RELEASE; ORAL

TENUATE

SANOFI AVENTIS US	75MG	N17669	001	
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TEPANIL TEN-TAB

3M	75MG	N17956	001	
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DIETHYLSTILBESTROL

INJECTABLE; INJECTION

STILBESTROL

BRISTOL MYERS SQUIBB	0.2MG/ML	N04056	003	
	0.5MG/ML	N04056	004	
	1MG/ML	N04056	005	
	5MG/ML	N04056	006	

SUPPOSITORY; VAGINAL

DIETHYLSTILBESTROL

LILLY	0.1MG	N04040	001	
	0.5MG	N04040	002	

STILBESTROL

BRISTOL MYERS SQUIBB	0.1MG	N04056	001	
	0.5MG	N04056	002	

TABLET; ORAL

DIETHYLSTILBESTROL

LILLY	0.1MG	N04041	002	
	0.5MG	N04041	003	
	1MG	N04041	004	
	5MG	N04041	005	

STILBESTROL

TABLICAPS	0.5MG	N83004	001	
	1MG	N83002	001	
	5MG	N83006	001	

STILBETIN

BRISTOL MYERS SQUIBB	0.1MG	N04056	007	
	0.25MG	N04056	017	
	0.5MG	N04056	008	
	1MG	N04056	009	
	5MG	N04056	010	

TABLET, DELAYED RELEASE; ORAL

DIETHYLSTILBESTROL

LILLY	0.1MG	N04039	002	
	0.25MG	N04039	005	
	0.5MG	N04039	003	
	1MG	N04039	004	
	5MG	N04039	006	

STILBESTROL

TABLICAPS	0.5MG	N83003	001	
	1MG	N83005	001	
	5MG	N83007	001	

STILBETIN

BRISTOL MYERS SQUIBB	0.1MG	N04056	011	
	0.5MG	N04056	012	
	1MG	N04056	013	
	5MG	N04056	014	

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 89 (of 274)

DIETHYLSTILBESTROL DIPHOSPHATEINJECTABLE; INJECTION
STILPHOSTROL

BAYER PHARMS 250MG/5ML

N10010 001

TABLET; ORAL
STILPHOSTROL

BAYER PHARMS 50MG

N10010 002

DIFLORASONE DIACETATE

CREAM; TOPICAL

FLORONE

PHARMACIA AND UPJOHN 0.05%

N17741 001

FLORONE E

PHARMACIA AND UPJOHN 0.05%

N19259 001 Aug 28, 1985

OINTMENT; TOPICAL
FLORONE

PHARMACIA AND UPJOHN 0.05%

N17994 001

PSORCON

PHARMACIA AND UPJOHN 0.05%

N19260 001 Aug 28, 1985

DIFLUNISAL

TABLET; ORAL

DIFLUNISAL

PUREPAC PHARM 250MG

N74285 001 May 07, 1996

500MG

N74285 002 May 07, 1996

ROXANE

250MG

N73562 001 Nov 27, 1992

500MG

N73563 001 Nov 27, 1992

DOLOBID

MERCK 250MG

N18445 001 Apr 19, 1982

500MG

N18445 002 Apr 19, 1982

DIGITOXININJECTABLE; INJECTION
CRYSTODIGIN

LILLY 0.2MG/ML

N84100 005

DIGOXIN

CAPSULE; ORAL

LANOXICAPS

SMITHKLINE BEECHAM 0.15MG

N18118 004 Sep 24, 1984

INJECTABLE; INJECTION
DIGOXIN

ABRAXIS PHARM 0.25MG/ML

N83217 001

WYETH AYERST 0.25MG/ML

N84386 001

TABLET; ORAL

LANOXIN

SMITHKLINE BEECHAM 0.0625MG

N20405 001 Sep 30, 1997

0.1875MG

N20405 003 Sep 30, 1997

0.375MG

N20405 005 Sep 30, 1997

0.5MG

N20405 006 Sep 30, 1997

DIHYDROERGOTAMINE MESYLATE; HEPARIN SODIUM; LIDOCAINE HYDROCHLORIDE

INJECTABLE; INJECTION

EMBOLEX

NOVARTIS 0.5MG/0.5ML; 2,500
UNITS/0.5ML; 5.33MG/0.5ML

N18885 001 Nov 30, 1984

0.5MG/0.7ML; 5,000

N18885 002 Nov 30, 1984

UNITS/0.7ML; 7.46MG/0.7ML

DILTIAZEM HYDROCHLORIDE

CAPSULE, EXTENDED RELEASE; ORAL

CARDIZEM SR

BIOVAIL 60MG

N19471 001 Jan 23, 1989

90MG

N19471 002 Jan 23, 1989

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 90 (of 274)

DILTIAZEM HYDROCHLORIDECAPSULE, EXTENDED RELEASE; ORAL
CARDIZEM SR

BIOVAIL	120MG	N19471	003	Jan 23, 1989
	180MG	N19471	004	Jan 23, 1989

DILTIAZEM HYDROCHLORIDE

BIOVAIL	60MG	N74845	001	Sep 15, 1999
	90MG	N74845	002	Sep 15, 1999
	120MG	N74845	003	Sep 15, 1999
TEVA	60MG	N74079	001	Nov 30, 1993
	90MG	N74079	002	Nov 30, 1993
	120MG	N74079	003	Nov 30, 1993

TABLET; ORAL

DILTIAZEM HYDROCHLORIDE

APOTHECON	30MG	N74051	001	Mar 31, 1993
	60MG	N74051	002	Mar 31, 1993
	90MG	N74051	003	Mar 31, 1993
	120MG	N74051	004	Mar 31, 1993
TEVA	30MG	N74084	001	Feb 25, 1994
	60MG	N74084	002	Feb 25, 1994
TEVA PHARMS	30MG	N74067	001	Nov 05, 1992
	60MG	N74067	002	Nov 05, 1992
	90MG	N74067	003	Nov 05, 1992
	120MG	N74067	004	Nov 05, 1992

DILTIAZEM MALATE

TABLET, EXTENDED RELEASE; ORAL

TIAMATE

MERCK	EQ 120MG HCL	N20506	001	Oct 04, 1996
	EQ 180MG HCL	N20506	002	Oct 04, 1996
	EQ 240MG HCL	N20506	003	Oct 04, 1996

DILTIAZEM MALATE; ENALAPRIL MALEATE

TABLET, EXTENDED RELEASE; ORAL

TECZEM

BIOVAIL	EQ 180MG HCL;5MG	N20507	001	Oct 04, 1996
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DIMENHYDRINATE

INJECTABLE; INJECTION

DIMENHYDRINATE

BAXTER HLTHCARE	50MG/ML	N84767	001	
WATSON LABS	50MG/ML	N83531	001	
WYETH AYERST	50MG/ML	N84316	001	

LIQUID; ORAL

DIMENHYDRINATE

ALRA	12.5MG/4ML	N80715	001	
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TABLET; ORAL

DIMENHYDRINATE

ANABOLIC	50MG	N85985	001	
HEATHER	50MG	N80841	001	
WATSON LABS	50MG	N85166	001	

DIMYRISTOYL LECITHIN; PERFLEXANE

INJECTABLE; INTRAVENOUS

IMAGENT

IMCOR PH	0.92MG/VIAL;0.092MG/VIAL	N21191	001	May 31, 2002
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DINOPROST TROMETHAMINE

INJECTABLE; INJECTION

PROSTIN F2 ALPHA

PHARMACIA AND UPJOHN	EQ 5MG BASE/ML	N17434	001	
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DISCONTINUED DRUG PRODUCT LIST

6 - 91 (of 274)

DIPHEMANIL METHYLSULFATE

TABLET; ORAL
PRANTAL

SCHERING	100MG		N08114	004
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DIPHENHYDRAMINE HYDROCHLORIDE

CAPSULE; ORAL
BENADRYL

MCNEIL CONS	25MG		N05845	007
	50MG		N05845	001

DIPHENHYDRAMINE HYDROCHLORIDE

ALRA	25MG		N80519	004	
	50MG		N80519	003	
ANABOLIC	25MG		N83634	001	
	50MG		N83275	001	
ELKINS SINN	25MG		N85701	001	
	50MG		N85701	002	
HALSEY	50MG		N87914	001	Jun 04, 1984
HEATHER	25MG		N84524	001	
	50MG		N83953	001	
IMPAX LABS	25MG		N80807	001	
	50MG		N80807	002	
IVAX PHARMS	25MG		N80762	001	
	50MG		N80762	002	
LANNETT	25MG		N80868	001	
	50MG		N80868	002	
LEDERLE	25MG		N86874	001	
	50MG		N86875	001	
LEINER	25MG		N83027	001	
	50MG		N83027	002	
MK LABS	25MG		N83087	001	
	50MG		N83087	002	
NEWTRON PHARMS	25MG		N86543	001	
	50MG		N86544	001	
PERRIGO	25MG		N83061	001	
	50MG		N83061	002	
PIIONEER PHARMS	25MG		N89101	001	Dec 20, 1985
	50MG		N88880	001	Dec 20, 1985
PUREPAC PHARM	25MG		N85156	001	
	50MG		N85150	001	
ROXANE	50MG		N80635	001	
SANDOZ	25MG		N80845	002	
	50MG		N80845	001	
SUPERPHARM	25MG		N89040	001	May 15, 1985
	50MG		N89041	001	May 15, 1985
TEVA	25MG		N85874	001	
	50MG		N85874	002	
VALEANT PHARM INTL	25MG		N80596	001	
VANGARD	25MG		N88034	001	Oct 27, 1982
	50MG		N87630	001	
WATSON LABS	25MG		N83797	001	
	25MG		N85138	001	
	50MG		N83797	002	
	50MG		N85083	001	
WEST WARD	50MG		N83567	001	
WHITEWORTH TOWN PLSN	25MG		N83441	001	
	50MG		N80800	001	

ELIXIR; ORAL
BELIX

HALSEY	12.5MG/5ML		N86586	001	Oct 03, 1983
BENADRYL					
MCNEIL CONS	12.5MG/5ML		N05845	004	
DIBENIL					
CENCI	12.5MG/5ML		N88304	001	Dec 16, 1983

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 92 (of 274)

DIPHENHYDRAMINE HYDROCHLORIDE

ELIXIR; ORAL

DIPHEN

USL PHARMA 12.5MG/5ML

N84640 001

DIPHENHYDRAMINE HYDROCHLORIDE

BUNDY 12.5MG/5ML

N83674 001

CENCI 12.5MG/5ML

N87941 001

Dec 17, 1982

KV PHARM 12.5MG/5ML

N85621 001

LANNETT 12.5MG/5ML

N80939 002

LEDERLE 12.5MG/5ML

N86937 001

LEINER 12.5MG/5ML

N85287 001

MK LABS 12.5MG/5ML

N83088 002

NASKA 12.5MG/5ML

N88680 001

May 31, 1985

PERRIGO 12.5MG/5ML

N83063 001

PUREPAC PHARM 12.5MG/5ML

N83237 001

Jan 25, 1982

ROXANE 12.5MG/5ML

N80643 001

HYDRAMINE

ALPHARMA US PHARMS 12.5MG/5ML

N80763 002

INJECTABLE; INJECTION

BENADRYL

PARKE DAVIS 10MG/ML

N06146 001

DIPHENHYDRAMINE HYDROCHLORIDE

ABRAXIS PHARM 10MG/ML

N87066 001

BAXTER HLTHCARE 50MG/ML

N83183 001

BEL MAR 10MG/ML

N80822 001

WATSON LABS 10MG/ML

N83533 001

WYETH AYERST 50MG/ML

N80577 001

DIPHENHYDRAMINE HYDROCHLORIDE PRESERVATIVE FREE

ABRAXIS PHARM 50MG/ML

N80586 002

SYRUP; ORAL

ANTITUSSIVE

PERRIGO 12.5MG/5ML

N71292 001

Apr 10, 1987

BELDIN

HALSEY 12.5MG/5ML

N89179 001

Jun 05, 1986

BENYLIN

PARKE DAVIS 12.5MG/5ML

N06514 004

DIPHEN

MORTON GROVE 12.5MG/5ML

N70118 001

Oct 01, 1985

DIPHENHYDRAMINE HYDROCHLORIDE

ALPHARMA US PHARMS 12.5MG/5ML

N70497 001

Apr 25, 1989

CUMBERLAND SWAN 12.5MG/5ML

N73611 001

Aug 20, 1992

HI TECH PHARMA 12.5MG/5ML

N72416 001

Sep 28, 1990

HYDRAMINE

ALPHARMA US PHARMS 12.5MG/5ML

N70205 001

Jan 28, 1986

SILPHEN

SILARX 12.5MG/5ML

N72646 001

Feb 27, 1992

VICKS FORMULA 44

PROCTER AND GAMBLE 12.5MG/5ML

N70524 001

Jan 14, 1987

DIPHENHYDRAMINE HYDROCHLORIDE; PSEUDOEPHEDRINE HYDROCHLORIDE

SOLUTION; ORAL

BENYLIN

PARKE DAVIS 12.5MG/5ML; 30MG/5ML

N19014 001

Jun 11, 1985

DIPHENIDOL HYDROCHLORIDE

TABLET; ORAL

VONTROL

GLAXOSMITHKLINE EQ 25MG BASE

N16033 001

DIPHENYLPYRALINE HYDROCHLORIDE

CAPSULE, EXTENDED RELEASE; ORAL

HISPRIL

GLAXOSMITHKLINE 5MG

N11945 001

DISCONTINUED DRUG PRODUCT LIST

6 - 93 (of 274)

DIPYRIDAMOLE

INJECTABLE; INJECTION					
IV PERSANTINE					
BOEHRINGER INGELHEIM	5MG/ML		N19817	001	Dec 13, 1990
TABLET; ORAL					
DIPYRIDAMOLE					
PUREPAC PHARM	50MG		N89426	001	Jul 12, 1990
	75MG		N89427	001	Jul 12, 1990

DIRITHROMYCIN

TABLET, DELAYED RELEASE; ORAL					
DYNABAC					
LILLY RES LABS	250MG		N50678	001	Jun 19, 1995

DISOPYRAMIDE PHOSPHATE

CAPSULE; ORAL					
DISOPYRAMIDE PHOSPHATE					
INTERPHARM	EQ 100MG BASE		N71190	001	Jan 15, 1987
	EQ 150MG BASE		N71191	001	Jan 15, 1987
IVAX PHARMS	EQ 100MG BASE		N70186	001	Nov 18, 1985
	EQ 150MG BASE		N70187	001	Nov 18, 1985
MUTUAL PHARM	EQ 100MG BASE		N70351	001	Dec 17, 1985
	EQ 150MG BASE		N70352	001	Dec 17, 1985
MYLAN	EQ 100MG BASE		N70138	001	Jun 14, 1985
	EQ 150MG BASE		N70139	001	Jun 14, 1985
SANDOZ	EQ 100MG BASE		N70470	001	Dec 10, 1985
	EQ 150MG BASE		N70471	001	Dec 10, 1985
SUPERPHARM	EQ 100MG BASE		N70940	001	Feb 09, 1987
	EQ 150MG BASE		N70941	001	Feb 09, 1987
WATSON LABS	EQ 100MG BASE		N70240	001	Feb 02, 1986
	EQ 150MG BASE		N70241	001	Feb 02, 1986

DISULFIRAM

TABLET; ORAL					
ANTABUSE					
ODYSSEY PHARMS	250MG		N07883	003	
	500MG		N07883	002	
DISULFIRAM					
PAR PHARM	250MG		N88792	001	Aug 14, 1984
	500MG		N88793	001	Aug 14, 1984
WATSON LABS	250MG		N86889	001	
	250MG		N87973	001	Aug 05, 1983
	500MG		N86890	001	
	500MG		N87974	001	Aug 05, 1983

DIVALPROEX SODIUM

TABLET, DELAYED RELEASE; ORAL					
DEPAKOTE CP					
ABBOTT	EQ 250MG BASE		N19794	001	Jul 11, 1990
	EQ 500MG BASE		N19794	002	Jul 11, 1990

DOBUTAMINE HYDROCHLORIDE

INJECTABLE; INJECTION					
DOBUTAMINE HYDROCHLORIDE					
ASTRAZENECA	EQ 12.5MG BASE/ML		N74098	001	Feb 21, 1995
BAXTER HLTHCARE	EQ 12.5MG BASE/ML		N74381	001	Sep 26, 1996
DOBUTREX					
LILLY	EQ 12.5MG BASE/ML		N17820	002	

DONEPEZIL HYDROCHLORIDE

SOLUTION; ORAL					
ARICEPT					
EISAI MEDCL RES	5MG/5ML		N21719	001	Oct 18, 2004

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 94 (of 274)

DOPAMINE HYDROCHLORIDEINJECTABLE; INJECTION
DOPAMINE HYDROCHLORIDE

ABBOTT	40MG/ML	N70656	001	Jan 24, 1989
	80MG/ML	N70657	001	Jan 24, 1989
ABRAXIS PHARM	40MG/ML	N18549	001	Mar 11, 1983
	40MG/ML	N70012	001	Jun 12, 1985
	40MG/ML	N70058	001	Mar 20, 1985
	80MG/ML	N70013	001	Jun 12, 1985
	80MG/ML	N70059	001	Mar 20, 1985
	160MG/ML	N70364	001	Dec 04, 1985
ASTRAZENECA	40MG/ML	N18656	001	Jun 28, 1983
	40MG/ML	N70087	001	Oct 23, 1985
	80MG/ML	N70089	001	Oct 23, 1985
	80MG/ML	N70090	001	Oct 23, 1985
	80MG/ML	N70091	001	Oct 23, 1985
	160MG/ML	N70092	001	Oct 23, 1985
	160MG/ML	N70093	001	Oct 23, 1985
	160MG/ML	N70094	001	Oct 23, 1985
BAXTER HLTHCARE	40MG/ML	N18398	001	
	80MG/ML	N18398	002	Mar 22, 1982
HOSPIRA	40MG/ML	N74403	001	May 23, 1996
INTL MEDICATION	40MG/ML	N18014	001	
SICOR PHARMS	40MG/ML	N72999	001	Oct 23, 1991
	80MG/ML	N73000	001	Oct 23, 1991
SMITH AND NEPHEW	40MG/ML	N70011	001	Aug 29, 1985
	40MG/ML	N70046	001	Aug 29, 1985
	80MG/ML	N70047	001	Aug 29, 1985
WARNER CHILCOTT	40MG/ML	N18138	001	
	40MG/ML	N70558	001	Sep 20, 1985
	80MG/ML	N70559	001	Sep 20, 1985
DOPAMINE HYDROCHLORIDE IN DEXTROSE 5%				
HOSPIRA	1.6MG/ML	N20542	001	Aug 30, 1995
INTROPIN				
MAYNE PHARMA USA	40MG/ML	N17395	001	
	80MG/ML	N17395	002	
	160MG/ML	N17395	003	

DOXACURIUM CHLORIDEINJECTABLE; INJECTION
NUROMAX

ABBOTT	EQ 1MG BASE/ML	N19946	001	Mar 07, 1991
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DOXEPIN HYDROCHLORIDECAPSULE; ORAL
DOXEPIN HYDROCHLORIDE

CLONMEL HLTHCARE	EQ 10MG BASE	N71685	001	Jan 05, 1988
	EQ 25MG BASE	N71686	001	Jan 05, 1988
	EQ 50MG BASE	N71673	001	Jan 05, 1988
	EQ 75MG BASE	N71674	001	Jan 05, 1988
	EQ 100MG BASE	N71675	001	Jan 05, 1988
	EQ 150MG BASE	N71676	001	Jan 05, 1988
MUTUAL PHARM	EQ 25MG BASE	N71502	001	Feb 18, 1988
	EQ 50MG BASE	N71653	001	Feb 18, 1988
	EQ 75MG BASE	N71654	001	Feb 18, 1988
	EQ 100MG BASE	N71521	001	Feb 18, 1988
NEW RIVER	EQ 10MG BASE	N16987	001	
	EQ 25MG BASE	N16987	002	
	EQ 50MG BASE	N16987	003	
	EQ 75MG BASE	N16987	006	
	EQ 100MG BASE	N16987	004	
	EQ 150MG BASE	N16987	007	Apr 13, 1987
PUREPAC PHARM	EQ 10MG BASE	N73054	001	Dec 28, 1990
	EQ 25MG BASE	N72109	001	Dec 28, 1990

DISCONTINUED DRUG PRODUCT LIST

6 - 95 (of 274)

DOXEPIN HYDROCHLORIDE

CAPSULE; ORAL

DOXEPIN HYDROCHLORIDE

PUREPAC PHARM	EQ 50MG BASE	N73055	001	Dec 28, 1990
	EQ 75MG BASE	N72386	001	Sep 08, 1988
	EQ 100MG BASE	N72110	001	Sep 08, 1988
	EQ 150MG BASE	N72387	001	Sep 08, 1988
QUANTUM PHARMICS	EQ 10MG BASE	N70972	001	Sep 29, 1987
	EQ 25MG BASE	N70973	001	Sep 29, 1987
	EQ 50MG BASE	N70931	001	Sep 29, 1987
	EQ 75MG BASE	N70932	001	Sep 29, 1987
	EQ 100MG BASE	N72375	001	Mar 15, 1989
	EQ 150MG BASE	N72376	001	Mar 15, 1989
SANDOZ	EQ 25MG BASE	N70827	001	May 15, 1986
	EQ 50MG BASE	N70828	001	May 15, 1986
	EQ 75MG BASE	N70825	001	May 15, 1986
	EQ 100MG BASE	N71562	001	Mar 02, 1987
WATSON LABS	EQ 10MG BASE	N70952	001	Mar 04, 1987
	EQ 25MG BASE	N70953	001	May 15, 1986
	EQ 50MG BASE	N70954	001	May 15, 1986
	EQ 75MG BASE	N71763	001	Feb 09, 1988
	EQ 100MG BASE	N70955	001	May 15, 1986
	EQ 150MG BASE	N71764	001	Feb 09, 1988

DOXORUBICIN HYDROCHLORIDE

INJECTABLE; INJECTION

ADRIAMYCIN RDF

PHARMACIA AND UPJOHN	10MG/VIAL	N50467	001	
	20MG/VIAL	N50467	003	May 20, 1985
	50MG/VIAL	N50467	002	
	150MG/VIAL	N50467	004	Jul 22, 1987

DOXYCYCLINE

CAPSULE; ORAL

DOXYCYCLINE

PAR PHARM	EQ 75MG BASE	N65055	004	Apr 18, 2005
	EQ 150MG BASE	N65055	003	Jul 15, 2005

FOR SUSPENSION; ORAL

DOXYCHEL

RACHELLE	EQ 25MG BASE/5ML	N61720	001	
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DOXYCYCLINE HYCLATE

CAPSULE; ORAL

DOXYCYCLINE HYCLATE

HALSEY	EQ 50MG BASE	N62119	002	May 24, 1985
	EQ 100MG BASE	N62119	001	May 24, 1985
HEATHER	EQ 50MG BASE	N62463	001	Dec 07, 1983
	EQ 100MG BASE	N62463	002	Dec 07, 1983
INTERPHARM	EQ 50MG BASE	N62763	001	Sep 02, 1988
	EQ 100MG BASE	N62763	002	Sep 02, 1988
LEINER	EQ 50MG BASE	N62631	001	Jul 24, 1986
	EQ 100MG BASE	N62631	002	Jul 24, 1986
MUTUAL PHARM	EQ 50MG BASE	N62418	001	Jan 28, 1983
	EQ 100MG BASE	N62418	002	Jan 28, 1983
PAR PHARM	EQ 50MG BASE	N62434	001	Oct 19, 1984
	EQ 100MG BASE	N62442	001	Dec 22, 1983
RANBAXY	EQ 50MG BASE	N62479	001	Dec 23, 1983
	EQ 100MG BASE	N62479	002	Dec 23, 1983
SUPERPHARM	EQ 50MG BASE	N62469	001	Oct 31, 1984
	EQ 100MG BASE	N62469	002	Oct 31, 1984
WARNER CHILCOTT	EQ 50MG BASE	N62594	001	Dec 05, 1985
	EQ 100MG BASE	N62594	002	Dec 05, 1985
WATSON LABS	EQ 50MG BASE	N62142	001	
	EQ 100MG BASE	N62142	002	

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 96 (of 274)

DOXYCYCLINE HYCLATE

CAPSULE; ORAL

DOXY-LEMMON

TEVA

EQ 50MG BASE

N62497 001 Aug 23, 1984

EQ 100MG BASE

N62497 002 Jun 15, 1984

PERIOSTAT

COLLAGENEX

EQ 20MG BASE

N50744 001 Sep 30, 1998

CAPSULE, COATED PELLETS; ORAL

DOXYCYCLINE HYCLATE

PLIVA

EQ 100MG BASE

N63187 001 Jun 30, 1992

CAPSULE, DELAYED RELEASE; ORAL

DORYX

MAYNE PHARMA INTL

EQ 75MG BASE

N50582 002 Aug 13, 2001

EQ 100MG BASE

N50582 001 Jul 22, 1985

EQ 100MG BASE

N62653 001 Oct 30, 1985

INJECTABLE; INJECTION

DOXYCHEL HYCLATE

RACHELLE

EQ 100MG BASE/VIAL

N61953 001

DOXYCYCLINE

BAXTER HLTHCARE

EQ 100MG BASE/VIAL

N62450 001 Oct 27, 1983

EQ 200MG BASE/VIAL

N62450 002 Oct 27, 1983

EQ 200MG BASE/VIAL

N62569 002 Mar 09, 1988

DOXYCYCLINE HYCLATE

BAXTER HLTHCARE

EQ 100MG BASE/VIAL

N62992 001 Feb 16, 1989

EQ 200MG BASE/VIAL

N62992 002 Feb 16, 1989

VIBRAMYCIN

PFIZER

EQ 100MG BASE/VIAL

N50442 002

EQ 200MG BASE/VIAL

N50442 001

TABLET; ORAL

DOXYCYCLINE HYCLATE

HEATHER

EQ 100MG BASE

N62462 001 May 11, 1983

INTERPHARM

EQ 100MG BASE

N62764 001 Sep 02, 1988

MUTUAL PHARM

EQ 100MG BASE

N62391 001 Sep 30, 1982

SUPERPHARM

EQ 100MG BASE

N62494 001 Feb 20, 1985

TRUXTON INC

EQ 50MG BASE

N62269 003

WARNER CHILCOTT

EQ 100MG BASE

N62593 001 Aug 28, 1985

WATSON LABS

EQ 50MG BASE

N62392 001 Mar 31, 1983

EQ 100MG BASE

N62392 002 Mar 31, 1983

DOXY-LEMMON

TEVA

EQ 100MG BASE

N62581 001 Mar 15, 1985

DOXYLAMINE SUCCINATE

CAPSULE; ORAL

UNISOM

PFIZER

25MG

N19440 001 Feb 05, 1986

TABLET; ORAL

DECAPRYN

SANOFI AVENTIS US

12.5MG

N06412 015

25MG

N06412 014

DOXYLAMINE SUCCINATE

QUANTUM PHARMICS

25MG

N88603 001 Aug 07, 1984

DOXY-SLEEP-AID

PAR PHARM

25MG

N70156 001 Jul 02, 1987

DOXYLAMINE SUCCINATE; PYRIDOXINE HYDROCHLORIDE

TABLET, EXTENDED RELEASE; ORAL

BENDECTIN

SANOFI AVENTIS US

10MG;10MG **Federal Register
determination that product was not
discontinued or withdrawn for safety
or efficacy reasons**

N10598 002

DISCONTINUED DRUG PRODUCT LIST

6 - 97 (of 274)

DROMOSTANOLONE PROPIONATE

INJECTABLE; INJECTION

DROLBAN

LILLY

50MG/ML

N12936 001

DROPERIDOL

INJECTABLE; INJECTION

DROPERIDOL

ABRAXIS PHARM

2.5MG/ML

N70992 001

Nov 17, 1986

ASTRAZENECA

2.5MG/ML

N70993 001

Nov 17, 1986

ASTRAZENECA

2.5MG/ML

N72018 001

Oct 20, 1988

2.5MG/ML

N72019 001

Oct 19, 1988

2.5MG/ML

N72020 001

Oct 19, 1988

2.5MG/ML

N72021 001

Oct 19, 1988

MAYNE PHARMA USA

2.5MG/ML

N71645 001

Apr 07, 1988

SMITH AND NEPHEW

2.5MG/ML

N71750 001

Sep 06, 1988

SOLOPAK

2.5MG/ML

N71754 001

Sep 06, 1988

2.5MG/ML

N71755 001

Sep 06, 1988

WATSON LABS

2.5MG/ML

N73520 001

Nov 27, 1991

2.5MG/ML

N73521 001

Nov 27, 1991

2.5MG/ML

N73523 001

Nov 27, 1991

DROPERIDOL; FENTANYL CITRATE

INJECTABLE; INJECTION

FENTANYL CITRATE AND DROPERIDOL

ASTRAZENECA

2.5MG/ML;EQ 0.05MG BASE/ML

N72026 001

Apr 13, 1989

2.5MG/ML;EQ 0.05MG BASE/ML

N72027 001

Apr 13, 1989

2.5MG/ML;EQ 0.05MG BASE/ML

N72028 001

Apr 13, 1989

HOSPIRA

2.5MG/ML;EQ 0.05MG BASE/ML

N71982 001

May 04, 1988

INNOVAR

AKORN MFG

2.5MG/ML;EQ 0.05MG BASE/ML

N16049 001

DYCLONINE HYDROCHLORIDE

SOLUTION; TOPICAL

DYCLONE

ASTRAZENECA

0.5% **Federal Register determination
that product was not discontinued or
withdrawn for safety or efficacy
reasons**

N09925 002

1% **Federal Register determination
that product was not discontinued or
withdrawn for safety or efficacy
reasons**

N09925 001

DYDROGESTERONE

TABLET; ORAL

GYNOREST

SOLVAY

5MG

N17388 001

10MG

N17388 002

DYPHYLLINE

ELIXIR; ORAL

NEOTHYLLINE

TEVA

160MG/15ML

N07794 003

INJECTABLE; INJECTION

NEOTHYLLINE

TEVA

250MG/ML

N09088 001

TABLET; ORAL

DILOR

SAVAGE LABS

200MG

N84514 001

DILOR-400

SAVAGE LABS

400MG

N84751 001

NEOTHYLLINE

TEVA

200MG

N07794 001

400MG

N07794 002

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 98 (of 274)

ECHOTHIOPHATE IODIDEFOR SOLUTION; OPHTHALMIC
PHOSPHOLINE IODIDE

AYERST	0.03%	N11963	002
	0.06%	N11963	004
	0.25%	N11963	003

EDETATE CALCIUM DISODIUM

TABLET; ORAL

CALCIUM DISODIUM VERSENATE

3M	500MG	N08922	002
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EDETATE DISODIUM

INJECTABLE; INJECTION

DISODIUM EDETATE

WATSON LABS	150MG/ML	N84356	001
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EDETATE DISODIUM

WATSON LABS	150MG/ML	N80391	001
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SODIUM VERSENATE

3M	200MG/ML	N10573	001
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EDROPHONIUM CHLORIDE

INJECTABLE; INJECTION

EDROPHONIUM CHLORIDE

WATSON LABS	10MG/ML	N40044	001	Mar 20, 1996
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EDROPHONIUM CHLORIDE PRESERVATIVE FREE

WATSON LABS	10MG/ML	N40043	001	Mar 20, 1996
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REVERSOL

ORGANON USA INC	10MG/ML	N89624	001	May 13, 1988
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EFAVIRENZ

TABLET; ORAL

SUSTIVA

BRISTOL MYERS SQUIBB	300MG	N21360	001	Feb 01, 2002
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EFLORNITHINE HYDROCHLORIDE

INJECTABLE; INJECTION

ORNIDYL

SANOFI AVENTIS US	200MG/ML	N19879	002	Nov 28, 1990
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ENALAPRIL MALEATE

TABLET; ORAL

ENALAPRIL MALEATE

APOTHECON	2.5MG	N75583	001	Aug 22, 2000
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5MG	N75583	002	Aug 22, 2000
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10MG	N75583	003	Aug 22, 2000
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20MG	N75583	004	Aug 22, 2000
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IVAX PHARMS	2.5MG	N75482	001	Aug 22, 2000
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5MG	N75482	002	Aug 22, 2000
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10MG	N75482	003	Aug 22, 2000
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20MG	N75482	004	Aug 22, 2000
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ENALAPRIL MALEATE; HYDROCHLOROTHIAZIDE

TABLET; ORAL

ENALAPRIL MALEATE AND HYDROCHLOROTHIAZIDE

IVAX PHARMS	5MG;12.5MG	N75736	001	Mar 25, 2003
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10MG;25MG	N75736	002	Mar 25, 2003
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ENFLURANE

LIQUID; INHALATION

ENFLURANE

ABBOTT	99.9%	N70803	001	Sep 08, 1987
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27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 99 (of 274)

ENOXACINTABLET; ORAL
PENETREXSANOFI AVENTIS US 200MG
400MGN19616 004 Dec 31, 1991
N19616 005 Dec 31, 1991ENOXAPARIN SODIUMINJECTABLE; SUBCUTANEOUS
LOVENOX (PRESERVATIVE FREE)SANOFI AVENTIS US 90MG/0.6ML (150MG/ML) **Federal
Register determination that product
was not discontinued or withdrawn for
safety or efficacy reasons**

N20164 006 Jun 02, 2000

EPINEPHRINEAEROSOL, METERED; INHALATION
BRONKAID MIST

STERLING 0.25MG/INH

N16803 001

INJECTABLE; INJECTION
SUS-PHRINE SULFITE-FREEFOREST LABS 1.5MG/AMP
5MG/MLN07942 003 Feb 05, 1999
N07942 001INJECTABLE; INTRAMUSCULAR
EPI E Z PEN JR

MERIDIAN MEDCL TECHN 0.15MG/DELIVERY

N19430 004 Aug 03, 1995

EPIPEN E Z PEN

MERIDIAN MEDCL TECHN 0.3MG/DELIVERY

N19430 003 Aug 03, 1995

EPINEPHRINE BITARTRATEAEROSOL, METERED; INHALATION
MEDIHALER-EPI

3M 0.3MG/INH

N10374 003

EPINEPHRINE BITARTRATE; ETIDOCAINE HYDROCHLORIDEINJECTABLE; INJECTION
DURANESTASTRAZENECA 0.005MG/ML;1%
0.005MG/ML;1.5%
DENTSPLY PHARM 0.005MG/ML;1.5%N17751 006
N17751 007
N21384 001EPINEPHRINE BITARTRATE; PRILOCAINE HYDROCHLORIDEINJECTABLE; INJECTION
CITANEST FORTE

ASTRAZENECA 0.005MG/ML;4%

N14763 008

EPINEPHRINE; ETIDOCAINE HYDROCHLORIDEINJECTABLE; INJECTION
DURANEST

ASTRAZENECA 0.005MG/ML;0.5%

N17751 004

EPINEPHRINE; LIDOCAINE HYDROCHLORIDE

INJECTABLE; INJECTION

ALPHACAINE HYDROCHLORIDE W/ EPINEPHRINE

CARLISLE 0.01MG/ML;2%
0.02MG/ML;2%N84720 001
N84732 001

LIDOCAINE HYDROCHLORIDE AND EPINEPHRINE

ELKINS SINN 0.01MG/ML;1%
0.01MG/ML;2%N80406 001
N80406 002

LIDOCAINE HYDROCHLORIDE W/ EPINEPHRINE

ABBOTT 0.01MG/ML;1%
BEL MAR 0.01MG/ML;1%
0.01MG/ML;2%N83154 001
N80820 001
N80757 001

INTL MEDICATION 0.01MG/ML;1%

N86402 001

WATSON LABS 0.01MG/ML;1%

N80377 003

DISCONTINUED DRUG PRODUCT LIST

6 - 100 (of 274)

EPINEPHRINE; LIDOCAINE HYDROCHLORIDE

INJECTABLE; INJECTION

LIDOCAINE HYDROCHLORIDE W/ EPINEPHRINE

WATSON LABS	0.01MG/ML;1%	N85463	001
	0.01MG/ML;2%	N80377	004

LIDOCATON

PHARMATON	0.02MG/ML;2%	N84728	001	Aug 17, 1983
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XYLOCAINE W/ EPINEPHRINE

ABRAXIS BIOSCIENCE	0.01MG/ML;2%	N06488	003
	0.02MG/ML;2%	N06488	005
ASTRAZENECA	0.005MG/ML;1%	N10418	006
	0.005MG/ML;1.5%	N10418	010
	0.005MG/ML;2%	N10418	008

SOLUTION; IONTOPHORESIS

IONTOCAINE

IOMED	0.01MG/ML;2%	N20530	001	Dec 21, 1995
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EPINEPHRINE; PROCAINE HYDROCHLORIDE

INJECTABLE; INJECTION

PROCAINE HYDROCHLORIDE W/ EPINEPHRINE

BEL MAR	0.02MG/ML;1%	N80758	001
	0.02MG/ML;2%	N80759	001

EPLERENONE

TABLET; ORAL

INSPRA

GD SEARLE LLC	100MG	N21437	003	Sep 27, 2002
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EPROSARTAN MESYLATE

TABLET; ORAL

TEVETEN

KOS LIFE	EQ 300MG BASE	N20738	004	Dec 22, 1997
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ERGOCALCIFEROL

CAPSULE; ORAL

DELTALIN

LILLY	50,000 IU	N80884	001
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VITAMIN D

CHASE CHEM	50,000 IU	N80747	001
EVERYLIFE	50,000 IU	N80956	001
IMPAX LABS	50,000 IU	N80951	001
LANNETT	50,000 IU	N80825	001
VITARINE	50,000 IU	N84053	001
WEST WARD	50,000 IU	N83102	001

ERGOLOID MESYLATES

CAPSULE; ORAL

HYDERGINE LC

NOVARTIS	1MG	N18706	001	Jan 18, 1983
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SOLUTION; ORAL

HYDERGINE

NOVARTIS	1MG/ML	N18418	001
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TABLET; ORAL

ERGOLOID MESYLATES

MUTUAL PHARM	1MG	N88891	001	Nov 01, 1985
WATSON LABS	1MG	N86433	001	May 27, 1982
	1MG	N87244	001	Aug 16, 1982

GERIMAL

WATSON LABS	1MG	N88207	001	Mar 22, 1984
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HYDERGINE

NOVARTIS	0.5MG	N17993	003
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TABLET; SUBLINGUAL

ALKERGOT

SANDOZ	0.5MG	N85153	001
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DISCONTINUED DRUG PRODUCT LIST

6 - 101 (of 274)

ERGOLOID MESYLATES

TABLET; SUBLINGUAL

ALKERGOT

SANDOZ

1MG

N87417 001

CIRCANOL

3M

0.5MG

N84868 001

1MG

N85809 001

DEAPRIL-ST

BRISTOL MYERS SQUIBB

1MG

N85020 002

ERGOLOID MESYLATES

KV PHARM

0.5MG

N85899 001

0.5MG

N86265 001

1MG

N85900 001

1MG

N86264 001

LEDERLE

0.5MG

N86984 001

1MG

N86985 001

MUTUAL PHARM

0.5MG

N87407 001

1MG

N87552 001

SUPERPHARM

0.5MG

N89233 001

Sep 23, 1986

1MG

N89234 001

Sep 23, 1986

VANGARD

0.5MG

N88013 001

Sep 20, 1982

1MG

N88014 001

Sep 20, 1982

WATSON LABS

0.5MG

N84930 001

1MG

N85177 001

1MG

N87183 001

HYDROGENATED ERGOT ALKALOIDS

IVAX PHARMS

0.5MG

N87186 001

ERGOTAMINE TARTRATE

AEROSOL, METERED; INHALATION

MEDIHALER ERGOTAMINE

3M

0.36MG/INH

N12102 001

TABLET; SUBLINGUAL

ERGOSTAT

PARKE DAVIS

2MG

N88337 001

Jun 08, 1984

WIGRETTES

ORGANON USA INC

2MG

N86750 001

Jul 29, 1982

ERYTHROMYCIN

CAPSULE, DELAYED REL PELLETS; ORAL

ERYC

PARKE DAVIS

250MG

N62546 001

Jul 25, 1985

250MG

N62618 001

Sep 25, 1985

ERYC 125

PARKE DAVIS

125MG

N62648 001

Oct 24, 1985

ERYC SPRINKLES

MAYNE PHARMA USA

125MG

N50593 001

Jul 22, 1985

GEL; TOPICAL

EMGEL

ALTANA

2%

N63107 001

Aug 23, 1991

OINTMENT; OPHTHALMIC

ERYTHROMYCIN

PHARMADERM

5MG/GM

N62446 001

Sep 26, 1983

PHARMAFAIR

5MG/GM

N62481 001

Apr 05, 1984

ILOTYCIN

DISTA

0.5%

N50368 001

POWDER; FOR RX COMPOUNDING

ERYTHROMYCIN

PADDOCK

100%

N50610 001

Nov 07, 1986

SOLUTION; TOPICAL

C-SOLVE-2

BIOGLAN PHARMA

2%

N62468 001

Jul 03, 1985

ERYMAX

MERZ PHARMS

2%

N62508 002

Jul 11, 1985

DISCONTINUED DRUG PRODUCT LIST

6 - 102 (of 274)

ERYTHROMYCIN

SOLUTION; TOPICAL ERYTHROMYCIN

ALPHARMA US PHARMS	1.5%	N62328	001	Apr 19, 1982
	2%	N62326	001	Apr 19, 1982
	2%	N62327	001	Apr 19, 1982
	2%	N62342	001	Feb 25, 1982
	2%	N62957	001	Jul 21, 1988
BAUSCH AND LOMB	2%	N64039	001	Jan 27, 1994
LILLY	2%	N50532	001	
PHARMAFAIR	1.5%	N62485	001	Jul 11, 1984
	2%	N62616	001	Jul 25, 1985
STIEFEL	2%	N64127	001	Feb 14, 1997
ERYTHRO-STATIN				
HI TECH PHARMA	2%	N64101	001	Oct 22, 1996
SANSAC				
HEALTHPOINT	2%	N62522	001	Jan 24, 1985
STATICIN				
WESTWOOD SQUIBB	1.5%	N50526	001	
T-STAT				
WESTWOOD SQUIBB	2%	N62436	001	Mar 09, 1983
SWAB; TOPICAL				
T-STAT				
WESTWOOD SQUIBB	2%	N62748	001	Jul 23, 1987
TABLET, DELAYED RELEASE; ORAL				
E-BASE				
BARR	333MG	N63028	001	May 15, 1990
ILOTYCIN				
DISTA	250MG	N61910	001	
ROBIMYCIN				
ROBINS AH	250MG	N61633	001	
R-P MYCIN				
SOLVAY	250MG	N61659	001	

ERYTHROMYCIN ESTOLATE

CAPSULE; ORAL

ERYTHROMYCIN ESTOLATE				
BARR	EQ 125MG BASE	N62162	001	
	EQ 250MG BASE	N62162	002	
IVAX PHARMS	EQ 250MG BASE	N62237	001	
WATSON LABS	EQ 250MG BASE	N62087	001	
ILOSONE				
LILLY	EQ 125MG BASE	N61897	001	
	EQ 250MG BASE	N61897	002	
FOR SUSPENSION; ORAL				
ILOSONE				
DISTA	EQ 125MG BASE/5ML	N61893	001	
SUSPENSION; ORAL				
ERYTHROMYCIN ESTOLATE				
BARR	EQ 125MG BASE/5ML	N62169	001	Oct 17, 1990
	EQ 250MG BASE/5ML	N62169	002	Oct 17, 1990
LIFE LABS	EQ 250MG BASE/5ML	N62362	001	Dec 17, 1982
ILOSONE				
LILLY	EQ 125MG BASE/5ML	N50010	001	
	EQ 125MG BASE/5ML	N61894	001	
	EQ 250MG BASE/5ML	N50010	002	
	EQ 250MG BASE/5ML	N61894	002	
SUSPENSION/DROPS; ORAL				
ILOSONE				
LILLY	EQ 100MG BASE/ML	N61894	003	
TABLET; ORAL				
ILOSONE				
LILLY	EQ 500MG BASE	N61896	001	

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 103 (of 274)

ERYTHROMYCIN ESTOLATETABLET, CHEWABLE; ORAL
ILOSONE

DISTA

EQ 125MG BASE

N61895 001

EQ 250MG BASE

N61895 002

ERYTHROMYCIN ESTOLATE; SULFISOXAZOLE ACETYL

SUSPENSION; ORAL

ILOSONE SULFA

LILLY

EQ 125MG BASE/5ML;EQ 600MG BASE/5ML

N50599 001

Sep 29, 1989

ERYTHROMYCIN ETHYLSUCCINATE

GRANULE; ORAL

PEDIAMYCIN

ROSS LABS

EQ 200MG BASE/5ML

N62305 001

SUSPENSION; ORAL

E-MYCIN E

PHARMACIA AND UPJOHN

EQ 200MG BASE/5ML

N62198 001

EQ 400MG BASE/5ML

N62198 002

ERYTHROMYCIN ETHYLSUCCINATE

DISTA

EQ 200MG BASE/5ML

N62177 001

EQ 400MG BASE/5ML

N62177 002

NASKA

EQ 400MG BASE/5ML

N62674 001

Mar 10, 1987

PARKE DAVIS

EQ 200MG BASE/5ML

N62231 001

EQ 400MG BASE/5ML

N62231 002

PHARMAFAIR

EQ 200MG BASE/5ML

N62559 001

Mar 15, 1985

EQ 400MG BASE/5ML

N62558 001

Mar 15, 1985

WYAMYCIN E

WYETH AYERST

EQ 200MG BASE/5ML

N62123 002

EQ 400MG BASE/5ML

N62123 001

SUSPENSION/DROPS; ORAL

PEDIAMYCIN

ROSS LABS

EQ 100MG BASE/2.5ML

N62305 002

TABLET; ORAL

E.E.S. 400

ABBOTT

EQ 400MG BASE

N61905 001

ERYTHROMYCIN ETHYLSUCCINATE

BARR

EQ 400MG BASE

N62256 001

TABLET, CHEWABLE; ORAL

E.E.S.

ABBOTT

EQ 200MG BASE

N50297 002

ERYPED

ABBOTT

EQ 200MG BASE

N50297 003

Jul 05, 1988

PEDIAMYCIN

ROSS LABS

EQ 200MG BASE

N62306 001

ERYTHROMYCIN GLUCEPTATE

INJECTABLE; INJECTION

ILOTYCIN GLUCEPTATE

DISTA

EQ 250MG BASE/VIAL

N50370 001

EQ 500MG BASE/VIAL

N50370 002

EQ 1GM BASE/VIAL

N50370 003

ERYTHROMYCIN LACTOBIONATE

INJECTABLE; INJECTION

ERYTHROCIN

ABBOTT

EQ 500MG BASE/VIAL

N62586 001

Jan 04, 1988

EQ 1GM BASE/VIAL

N62586 002

Jan 04, 1988

ERYTHROMYCIN

ELKINS SINN

EQ 500MG BASE/VIAL

N62563 001

Mar 28, 1985

EQ 1GM BASE/VIAL

N62563 002

Mar 28, 1985

ERYTHROMYCIN LACTOBIONATE

ABRAXIS PHARM

EQ 500MG BASE/VIAL

N62604 001

Nov 24, 1986

EQ 1GM BASE/VIAL

N62604 002

Nov 24, 1986

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 104 (of 274)

ERYTHROMYCIN STEARATETABLET; ORAL
BRISTAMYCIN

BRISTOL	EQ 250MG BASE	N61304	001	
	EQ 250MG BASE	N61887	001	

ERYPAR

PARKE DAVIS	EQ 250MG BASE	N62032	001	
	EQ 500MG BASE	N62032	002	
WARNER CHILCOTT	EQ 250MG BASE	N62322	001	

ERYTHROCIN STEARATE

ABBOTT	EQ 125MG BASE	N60359	002	
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ERYTHROMYCIN STEARATE

BARR	EQ 500MG BASE	N63179	001	May 15, 1990
IVAX PHARMS	EQ 250MG BASE	N61461	001	
	EQ 500MG BASE	N61461	002	
LEDERLE	EQ 250MG BASE	N62089	001	
	EQ 500MG BASE	N62089	002	
PUREPAC PHARM	EQ 250MG BASE	N61743	001	
WATSON LABS	EQ 250MG BASE	N62121	002	
	EQ 500MG BASE	N62121	001	

ETHRIL 250

BRISTOL MYERS SQUIBB	EQ 250MG BASE	N61605	001	
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ETHRIL 500

BRISTOL MYERS SQUIBB	EQ 500MG BASE	N61605	002	
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PFIZER-E

PFIZER	EQ 250MG BASE	N61791	001	
	EQ 500MG BASE	N61791	002	

WYAMYCIN S

WYETH AYERST	EQ 250MG BASE	N61675	001	
	EQ 500MG BASE	N61675	002	

ESMOLOL HYDROCHLORIDEINJECTABLE; INJECTION
BREVIBLOC

BAXTER HLTHCARE CORP	100MG/ML	N19386	003	Dec 31, 1986
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ESTRADIOLFILM, EXTENDED RELEASE; TRANSDERMAL
ESCLIM

WOMEN FIRST HLTHCARE	0.025MG/24HR	N20847	001	Aug 04, 1998
	0.0375MG/24HR	N20847	002	Aug 04, 1998
	0.05MG/24HR	N20847	003	Aug 04, 1998
	0.075MG/24HR	N20847	004	Aug 04, 1998
	0.1MG/24HR	N20847	005	Aug 04, 1998

ESTRADIOL

ORTHO MCNEIL PHARM	0.05MG/24HR	N21048	001	Sep 20, 1999
	0.075MG/24HR	N21048	002	Sep 20, 1999
	0.1MG/24HR	N21048	003	Sep 20, 1999

FEMPATCH

PARKE DAVIS	0.025MG/24HR	N20417	001	Dec 03, 1996
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VIVELLE

NOVARTIS	0.025MG/24HR	N20323	005	Aug 16, 2000
	0.0375MG/24HR	N20323	001	Oct 28, 1994
	0.075MG/24HR	N20323	003	Oct 28, 1994

GEL; TOPICAL

ESTROGEL

ASCEND	0.06%	N21166	001	Feb 09, 2004
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TABLET; ORAL

ESTRADIOL

RADIUS PHARMS	0.5MG	N40275	001	Dec 29, 1998
	1MG	N40275	002	Dec 29, 1998
	2MG	N40275	003	Dec 29, 1998

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 105 (of 274)

ESTRADIOL CYPIONATEINJECTABLE; INJECTION
DEPO-ESTRADIOL

PHARMACIA AND UPJOHN	1MG/ML	N85470	001
	3MG/ML	N85470	002

ESTRADIOL CYPIONATE; MEDROXYPROGESTERONE ACETATEINJECTABLE; INTRAMUSCULAR
LUNELLE

PHARMACIA AND UPJOHN	5MG/0.5ML;25MG/0.5ML	N20874	001	Oct 05, 2000
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ESTRADIOL CYPIONATE; TESTOSTERONE CYPIONATEINJECTABLE; INJECTION
DEPO-TESTADIOL

PHARMACIA AND UPJOHN	2MG/ML;50MG/ML	N17968	001
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TESTOSTERONE CYPIONATE-ESTRADIOL CYPIONATE

WATSON LABS	2MG/ML;50MG/ML	N85603	001	Mar 13, 1986
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ESTRADIOL VALERATEINJECTABLE; INJECTION
ESTRADIOL VALERATE

WATSON LABS	10MG/ML	N83546	001
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ESTRADIOL VALERATE; TESTOSTERONE ENANTHATEINJECTABLE; INJECTION
DITATE-DS

SAVAGE LABS	8MG/ML;180MG/ML	N86423	001
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TESTOSTERONE ENANTHATE AND ESTRADIOL VALERATE

WATSON LABS	4MG/ML;90MG/ML	N85865	001
	8MG/ML;180MG/ML	N85860	001

ESTROGENS, CONJUGATEDTABLET; ORAL
PREMARIN

WYETH PHARMS INC	2.5MG	N04782	002
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ESTROGENS, CONJUGATED; MEDROXYPROGESTERONE ACETATE

TABLET; ORAL-28

PREMPHASE (PREMARIN;CYCRIN 14/14)

WYETH PHARMS INC	0.625MG, 0.625MG;N/A, 5MG	N20303	002	Dec 30, 1994
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PREMPRO (PREMARIN;CYCRIN)

WYETH PHARMS INC	0.625MG, 0.625MG;2.5MG, 2.5MG	N20303	001	Dec 30, 1994
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ESTROGENS, CONJUGATED; MEPROBAMATE

TABLET; ORAL

MILPREM-200

MEDPOINTE PHARM HLC	0.45MG;200MG	N11045	002
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MILPREM-400

MEDPOINTE PHARM HLC	0.45MG;400MG	N11045	001
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PMB 200

WYETH AYERST	0.45MG;200MG	N10971	005
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PMB 400

WYETH AYERST	0.45MG;400MG	N10971	003
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ESTROGENS, ESTERIFIED

TABLET; ORAL

AMNESTROGEN

BRISTOL MYERS SQUIBB	0.3MG	N83266	001
	0.625MG	N83266	002
	1.25MG	N83266	003
	2.5MG	N83266	004

ESTERIFIED ESTROGENS

LEINER	0.625MG	N83414	001
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27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 106 (of 274)

ESTROGENS, ESTERIFIED

TABLET; ORAL

ESTERIFIED ESTROGENS

LEINER	1.25MG	N83765	001
	2.5MG	N85907	001
SANDOZ	1.25MG	N85302	001
ESTRATAB			
SOLVAY	0.3MG	N86715	001
	0.625MG	N83209	001
	1.25MG	N83856	001
	2.5MG	N83857	001
EVEX			
ROCHE PALO	0.625MG	N84215	001
	1.25MG	N83376	002
FEMOGEN			
LEINER	0.625MG	N85076	001
	1.25MG	N85008	001
	2.5MG	N85007	001

ESTRONE

INJECTABLE; INJECTION

ESTROGENIC SUBSTANCE

WYETH AYERST	2MG/ML	N83488	001
ESTRONE			
WATSON LABS	2MG/ML	N83397	001
NATURAL ESTROGENIC SUBSTANCE-ESTRONE			
WATSON LABS	2MG/ML	N85237	001 Nov 23, 1982
THEELIN			
PARKEDALE	1MG/ML	N03977	001
	2MG/ML	N03977	002
	5MG/ML	N03977	003

ETHACRYNIC ACID

TABLET; ORAL

EDECIN

ATON	50MG	N16092	002
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ETHAMBUTOL HYDROCHLORIDE

TABLET; ORAL

MYAMBUTOL

STAT TRADE	200MG	N16320	002
	500MG	N16320	004

ETHCHLORVYNOL

CAPSULE; ORAL

ETHCHLORVYNOL

BANNER PHARMACAPS	100MG	N84463	001
	200MG	N84463	002
	500MG	N84463	003
	750MG	N84463	004
PLACIDYL			
ABBOTT	100MG	N10021	004
	200MG	N10021	007
	500MG	N10021	002
	750MG	N10021	010

ETHINAMATE

CAPSULE; ORAL

VALMID

DISTA	500MG	N09750	001
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DISCONTINUED DRUG PRODUCT LIST

6 - 107 (of 274)

ETHINYL ESTRADIOL

TABLET; ORAL
ESTINYL

SCHERING	0.02MG	N05292	001
	0.05MG	N05292	002
	0.5MG	N05292	003

FEMINONE

PHARMACIA AND UPJOHN	0.05MG	N16649	001
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LYNORAL

ORGANON USA INC	0.01MG	N05490	003
	0.05MG	N05490	002

ETHINYL ESTRADIOL; FERROUS FUMARATE; NORETHINDRONE

TABLET; ORAL-28

NORQUEST FE

GD SEARLE LLC	0.035MG;75MG;1MG	N18926	001	Jul 18, 1986
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ETHINYL ESTRADIOL; FERROUS FUMARATE; NORETHINDRONE ACETATE

TABLET; ORAL-28

NORLESTRIN FE 1/50

PARKE DAVIS	0.05MG;75MG;1MG	N16766	001
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NORLESTRIN FE 2.5/50

PARKE DAVIS	0.05MG;75MG;2.5MG	N16854	001
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ETHINYL ESTRADIOL; LEVONORGESTREL

TABLET; ORAL

PREVEN EMERGENCY CONTRACEPTIVE KIT

DURAMED	0.05MG;0.25MG	N20946	001	Sep 01, 1998
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TABLET; ORAL-21

ALESSE

WYETH PHARMS INC	0.02MG;0.1MG	N20683	001	Mar 27, 1997
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AVIANE-21

DURAMED PHARMS BARR	0.02MG;0.1MG	N75796	002	Apr 30, 2001
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ENPRESSE-21

DURAMED PHARMS BARR	0.03MG,0.04MG,0.03MG;0.05MG,0.075MG,0.125MG	N75809	001	Jul 16, 2001
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LESSINA-21

BARR	0.02MG;0.1MG	N75803	001	Mar 20, 2002
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LEVLITE

BERLEX	0.02MG;0.1MG	N20860	001	Jul 13, 1998
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LEVONORGESTREL AND ETHINYL ESTRADIOL

BARR	0.02MG;0.1MG	N75862	001	Apr 29, 2003
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LEVORA 0.15/30-21

WATSON LABS	0.03MG;0.15MG	N73592	001	Dec 13, 1993
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NORDETTE-21

DURAMED	0.03MG;0.15MG	N18668	001	May 10, 1982
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PORTIA-21

BARR	0.03MG;0.15MG	N75866	001	May 23, 2002
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TRIPHASIL-21

WYETH PHARMS INC	0.03MG,0.04MG,0.03MG;0.05MG,0.075MG,0.125MG	N19192	001	Nov 01, 1984
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TRIVORA-21

WATSON LABS	0.03MG,0.04MG,0.03MG;0.05MG,0.075MG,0.125MG	N74538	001	Dec 18, 1997
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ETHINYL ESTRADIOL; NORETHINDRONE

TABLET; ORAL-21

MODICON 21

ORTHO MCNEIL PHARM	0.035MG;0.5MG	N17488	001
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N.E.E. 1/35 21

LPI	0.035MG;1MG	N71541	001	Dec 14, 1987
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NORCEPT-E 1/35 21

ORTHO MCNEIL PHARM	0.035MG;1MG	N71545	001	Feb 09, 1989
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DISCONTINUED DRUG PRODUCT LIST

6 - 108 (of 274)

ETHINYL ESTRADIOL; NORETHINDRONE

TABLET; ORAL-21				
ORTHO-NOVUM 1/35-21				
ORTHO MCNEIL PHARM	0.035MG;1MG	N17489	002	
ORTHO-NOVUM 10/11-21				
ORTHO MCNEIL PHARM	0.035MG, 0.035MG;0.5MG, 1MG	N18354	001	Jan 11, 1982
ORTHO-NOVUM 7/14-21				
ORTHO MCNEIL PHARM	0.035MG, 0.035MG;0.5MG, 1MG	N19004	001	Apr 04, 1984
ORTHO-NOVUM 7/7/7-21				
ORTHO MCNEIL PHARM	0.035MG, 0.035MG, 0.035MG;0.5MG, 0.75MG, 1MG	N18985	001	Apr 04, 1984
OVCON-35				
WARNER CHILCOTT	0.035MG;0.4MG	N18127	001	
OVCON-50				
WARNER CHILCOTT	0.05MG;1MG	N18128	001	
TRI-NORINYL 21-DAY				
WATSON LABS	0.035MG, 0.035MG, 0.035MG;0.5MG, 1MG, 0.5MG	N18977	001	Apr 13, 1984
TABLET; ORAL-28				
N.E.E. 1/35 28				
LPI	0.035MG;1MG	N71542	001	Dec 14, 1987
NORCEPT-E 1/35 28				
ORTHO MCNEIL PHARM	0.035MG;1MG	N71546	001	Feb 09, 1989
ORTHO-NOVUM 7/14-28				
ORTHO MCNEIL PHARM	0.035MG, 0.035MG;0.5MG, 1MG	N19004	002	Apr 04, 1984

ETHINYL ESTRADIOL; NORETHINDRONE ACETATE

TABLET; ORAL-21				
ESTROSTEP 21				
WARNER CHILCOTT	0.02MG, 0.03MG, 0.035MG;1MG, 1MG, 1MG	N20130	001	Oct 09, 1996
NORLESTRIN 21 1/50				
PARKE DAVIS	0.05MG;1MG	N16749	001	
NORLESTRIN 21 2.5/50				
PARKE DAVIS	0.05MG;2.5MG	N16852	001	
TABLET; ORAL-28				
NORLESTRIN 28 1/50				
PARKE DAVIS	0.05MG;1MG	N16723	001	

ETHINYL ESTRADIOL; NORGESTIMATE

TABLET; ORAL-21				
ORTHO CYCLEN-21				
JOHNSON RW	0.035MG;0.25MG	N19653	001	Dec 29, 1989
ORTHO TRI-CYCLEN				
JOHNSON RW	0.035MG, 0.035MG, 0.035MG;0.18MG, 0.215MG, 0.25MG	N19697	002	Jul 03, 1992

ETHINYL ESTRADIOL; NORGESTREL

TABLET; ORAL-21				
LO/OVRAL				
WYETH PHARMS INC	0.03MG;0.3MG	N17612	001	
OVRAL				
WYETH PHARMS INC	0.05MG;0.5MG	N16672	001	
TABLET; ORAL-28				
OVRAL-28				
WYETH PHARMS INC	0.05MG;0.5MG	N16806	001	

ETHOPROPAZINE HYDROCHLORIDE

TABLET; ORAL				
PARSIDOL				
PARKE DAVIS	10MG	N09078	003	
	50MG	N09078	006	
	100MG	N09078	008	

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 109 (of 274)

ETHOTOIN

TABLET; ORAL PEGANONE OVATION PHARMS	500MG	N10841	003
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ETHOXZOLAMIDE

TABLET; ORAL CARDRASE PHARMACIA AND UPJOHN	62.5MG 125MG	N11047	002 001
ETHAMIDE ALLERGAN	125MG	N16144	001

ETHYLESTRENOL

ELIXIR; ORAL MAXIBOLIN ORGANON USA INC	2MG/5ML	N14006	002
TABLET; ORAL MAXIBOLIN ORGANON USA INC	2MG	N14005	002

ETHYNODIOL DIACETATE; MESTRANOL

TABLET; ORAL-20 OVULEN GD SEARLE LLC	1MG;0.1MG	N16029	002
TABLET; ORAL-21 OVULEN-21 GD SEARLE LLC	1MG;0.1MG	N16029	003
TABLET; ORAL-28 OVULEN-28 GD SEARLE LLC	1MG;0.1MG	N16705	001

ETIDOCAINE HYDROCHLORIDE

INJECTABLE; INJECTION DURANEST ASTRAZENECA	0.5% 1%	N17751	003 005
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ETIDRONATE DISODIUM

INJECTABLE; INJECTION DIDRONEL MGI PHARMA INC	50MG/ML	N19545	001	Apr 20, 1987
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ETODOLAC

CAPSULE; ORAL ETODOLAC ENDO PHARMS	200MG 300MG	N74842	001	Jul 17, 1997
		N74842	002	Jul 17, 1997
IVAX PHARMS	200MG 300MG	N74899	001	Jul 08, 1997
		N74899	002	Jul 08, 1997
LODINE WYETH PHARMS INC	200MG 300MG	N18922	002	Jan 31, 1991
		N18922	003	Jan 31, 1991
TABLET; ORAL ETODOLAC ENDO PHARMS	400MG	N74841	001	Jun 27, 1997
LODINE WYETH PHARMS INC	400MG 500MG	N18922	004	Jul 29, 1993
		N18922	005	Jun 28, 1996
TABLET, EXTENDED RELEASE; ORAL ETODOLAC POINT HOLDINGS	400MG	N75696	001	Jul 31, 2000
LODINE XL WYETH PHARMS INC	400MG	N20584	001	Oct 25, 1996

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 110 (of 274)

ETODOLAC

TABLET, EXTENDED RELEASE; ORAL
 LODINE XL

WYETH PHARMS INC	500MG	N20584	003	Jan 20, 1998
	600MG	N20584	002	Oct 25, 1996

ETOPOSIDE

CAPSULE; ORAL
 VEPESID

BRISTOL MYERS SQUIBB	100MG	N19557	002	Dec 30, 1986
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INJECTABLE; INJECTION
 ETOPOSIDE

PIERRE FABRE	20MG/ML	N74813	001	Jul 09, 1997
SICOR PHARMS	20MG/ML	N74510	001	Jun 29, 1995
WATSON LABS	20MG/ML	N74228	001	Oct 15, 1996
TOPOSAR				
SICOR PHARMS	20MG/ML	N74166	001	Feb 27, 1995

ETOPOSIDE PHOSPHATE

INJECTABLE; INJECTION
 ETOPOPHOS PRESERVATIVE FREE

BRISTOL MYERS SQUIBB	EQ 500MG BASE/VIAL	N20906	001	Feb 27, 1998
	EQ 1GM BASE/VIAL	N20906	002	Feb 27, 1998

ETRETINATE

CAPSULE; ORAL
 TEGISON

ROCHE	10MG	N19369	001	Sep 30, 1986
	25MG	N19369	002	Sep 30, 1986

EVANS BLUE

INJECTABLE; INJECTION
 EVANS BLUE

PARKE DAVIS	0.5% **Federal Register determination that product was not discontinued or withdrawn for safety or efficacy reasons**	N08041	001	
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FAMOTIDINE

INJECTABLE; INJECTION
 FAMOTIDINE

APOTHECON	10MG/ML	N75707	001	Apr 16, 2001
MAYNE PHARMA USA	10MG/ML	N75705	001	Apr 16, 2001
FAMOTIDINE PRESERVATIVE FREE				
APOTHECON	10MG/ML	N75708	001	Apr 16, 2001
MAYNE PHARMA USA	10MG/ML	N75669	001	Apr 16, 2001
FAMOTIDINE PRESERVATIVE FREE IN PLASTIC CONTAINER				
ABBOTT	0.4MG/ML	N75729	001	Dec 17, 2001

TABLET, ORALLY DISINTEGRATING; ORAL
 PEPCID RPD

MERCK	20MG	N20752	001	May 28, 1998
	40MG	N20752	002	May 28, 1998

FENOFIBRATE

CAPSULE; ORAL
 ANTARA (MICRONIZED)

OSCIENT	87MG	N21695	002	Nov 30, 2004
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LIPIDIL

ABBOTT	100MG	N19304	001	Dec 31, 1993
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TRICOR (MICRONIZED)

ABBOTT	67MG	N19304	002	Feb 09, 1998
	134MG	N19304	003	Jun 30, 1999
	200MG	N19304	004	Jun 30, 1999

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 111 (of 274)

FENOFIBRATETABLET; ORAL
TRICOR

ABBOTT

54MG
160MGN21203 001 Sep 04, 2001
N21203 003 Sep 04, 2001FENOPROFEN CALCIUM

CAPSULE; ORAL

FENOPROFEN CALCIUM

AM THERAP

EQ 200MG BASE
EQ 300MG BASEN72307 001 Aug 22, 1988
N72308 001 Aug 22, 1988

HALSEY

EQ 200MG BASE
EQ 300MG BASEN72355 001 Aug 17, 1988
N72356 001 Aug 17, 1988

QUANTUM PHARMICS

EQ 200MG BASE
EQ 300MG BASEN72214 001 Aug 17, 1988
N71738 001 Aug 17, 1988

WARNER CHILCOTT

EQ 200MG BASE
EQ 300MG BASEN72946 001 Apr 30, 1991
N72472 001 Apr 30, 1991

WATSON LABS

EQ 200MG BASE
EQ 200MG BASE
EQ 300MG BASE
EQ 300MG BASEN72294 001 Aug 17, 1988
N72981 001 Aug 19, 1991
N72293 001 Aug 17, 1988
N72982 001 Aug 19, 1991

TABLET; ORAL

FENOPROFEN CALCIUM

AM THERAP

EQ 600MG BASE

N72309 001 Aug 17, 1988

CLONMEL HLTHCARE

EQ 600MG BASE

N72326 001 Aug 17, 1988

HALSEY

EQ 600MG BASE

N72357 001 Aug 17, 1988

QUANTUM PHARMICS

EQ 600MG BASE

N72194 001 Aug 17, 1988

USL PHARMA

EQ 600MG BASE

N72362 001 Aug 17, 1988

WATSON LABS

EQ 600MG BASE

N72165 001 Aug 17, 1988

NALFON

DISTA

EQ 600MG BASE

N17710 001

FENTANYL CITRATE

INJECTABLE; INJECTION

FENTANYL CITRATE

ABBOTT

EQ 0.05MG BASE/ML
EQ 0.05MG BASE/MLN70636 001 Apr 30, 1990
N70637 001 Apr 30, 1990

WATSON LABS

EQ 0.05MG BASE/ML

N73488 001 Jun 30, 1992

FENTANYL CITRATE PRESERVATIVE FREE

MARSAM PHARMS LLC

EQ 0.05MG BASE/ML

N74917 001 Feb 03, 1998

TROCHE/LOZENGE; ORAL

FENTANYL

CEPHALON

EQ 0.1MG BASE
EQ 0.2MG BASE
EQ 0.3MG BASE
EQ 0.4MG BASEN20195 007 Oct 30, 1995
N20195 001 Oct 04, 1993
N20195 002 Oct 04, 1993
N20195 003 Oct 04, 1993FERRIC AMMONIUM CITRATE

FOR SOLUTION; ORAL

FERRISELTZ

OTSUKA

600MG/PACKET

N20292 001 Oct 14, 1997

FERROUS CITRATE, FE-59

INJECTABLE; INJECTION

FERROUS CITRATE FE 59

MALLINCKRODT

25uCi/ML

N16729 001

FERROUS SULFATE; FOLIC ACID

CAPSULE; ORAL

FOLVRON

LEDERLE

182MG;0.33MG

N06012 003

DISCONTINUED DRUG PRODUCT LIST

6 - 112 (of 274)

FIBRINOGEN, I-125

INJECTABLE; INJECTION
IBRIN

GE HEALTHCARE	154uCi/VIAL	N17879	001	
RADIONUCLIDE-LABELED (125 I) FIBRINOGEN (HUMAN) SENSOR				
ABBOTT	140uCi/ML	N17787	001	

FLECAINIDE ACETATE

TABLET; ORAL
TAMBOCOR
3M

	200MG	N18830	002	Oct 31, 1985
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FLUCONAZOLE

INJECTABLE; INJECTION

DIFLUCAN IN DEXTROSE 5% IN PLASTIC CONTAINER

PFIZER	2MG/ML	N19950	005	Jul 08, 1994
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DIFLUCAN IN SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER

PFIZER	2MG/ML	N19950	004	Jul 08, 1994
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TABLET; ORAL

FLUCONAZOLE

GEDEON RICHTER USA	50MG	N76432	001	Jul 29, 2004
	100MG	N76432	002	Jul 29, 2004
	150MG	N76432	003	Jul 29, 2004
	200MG	N76432	004	Jul 29, 2004

FLUDEOXYGLUCOSE F-18

INJECTABLE; INJECTION

FLUDEOXYGLUCOSE F 18

DOWNSSTATE CLINCL	4-40mCi/ML	N20306	001	Aug 19, 1994
	4-90mCi/ML	N20306	002	Sep 25, 2001

FLUMETHASONE PIVALATE

CREAM; TOPICAL
LOCORTEN

NOVARTIS	0.03%	N16379	001	
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FLUNISOLIDE

SPRAY, METERED; NASAL
NASALIDE

IVAX RES	0.025MG/SPRAY	N18148	001	
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FLUOCINOLONE ACETONIDE

CREAM; TOPICAL
FLUOCET

ALPHARMA US PHARMS	0.025%	N88360	001	Jan 16, 1984
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FLUOCINOLONE ACETONIDE

ALPHARMA US PHARMS	0.01%	N88361	001	Jan 16, 1984
PERRIGO NEW YORK	0.01%	N86810	001	Mar 04, 1982
	0.025%	N86811	001	Mar 04, 1982
PHARMADERM	0.01%	N88047	001	Dec 16, 1982
	0.025%	N88045	001	Dec 16, 1982
PHARMAFAIR	0.01%	N88499	001	Aug 02, 1984
	0.025%	N88506	001	Aug 02, 1984
TARO	0.01%	N40035	001	Oct 31, 1994
	0.025%	N40042	001	Oct 31, 1994
USL PHARMA	0.01%	N88757	001	Feb 11, 1985
	0.025%	N88756	001	Mar 28, 1985

FLUONID

ALLERGAN HERBERT	0.025%	N87156	002	Sep 06, 1984
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FLUOTREX

SAVAGE LABS	0.01%	N88174	001	May 06, 1983
	0.025%	N88173	001	Mar 09, 1983

SYNALAR-HP

MEDICIS	0.2%	N16161	002	
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27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 113 (of 274)

FLUOCINOLONE ACETONIDE

GEL; TOPICAL

FLUONID

ALLERGAN HERBERT

0.025%

N87300 001

May 27, 1982

OINTMENT; TOPICAL

FLUOCINOLONE ACETONIDE

PHARMADERM

0.025%

N88046 001

Dec 16, 1982

PHARMAFAIR

0.025%

N88507 001

Feb 27, 1984

USL PHARMA

0.025%

N88742 001

Feb 08, 1985

FLUONID

ALLERGAN HERBERT

0.025%

N87157 001

Sep 06, 1984

FLUOTREX

SAVAGE LABS

0.025%

N88172 001

Mar 09, 1983

SOLUTION; TOPICAL

FLUOCINOLONE ACETONIDE

ALPHARMA US PHARMS

0.01%

N87159 001

Jun 16, 1982

BAUSCH AND LOMB

0.01%

N40059 001

Dec 20, 1993

MORTON GROVE

0.01%

N88312 001

Jan 27, 1984

PHARMADERM

0.01%

N88048 001

Dec 16, 1982

PHARMAFAIR

0.01%

N88449 001

Feb 08, 1984

FLUONID

ALLERGAN HERBERT

0.01%

N87158 001

Mar 17, 1983

FLUOTREX

SAVAGE LABS

0.01%

N88171 001

Mar 09, 1983

FLUOCINOLONE ACETONIDE; NEOMYCIN SULFATE

CREAM; TOPICAL

NEO-SYNALAR

MEDICIS

0.025%;EQ 3.5MG BASE/GM

N60700 001

FLUOCINONIDE

CREAM; TOPICAL

FLUOCINONIDE

PERRIGO NEW YORK

0.05%

N71790 001

Jul 13, 1988

FLUORESCEIN SODIUM

INJECTABLE; INJECTION

FUNDUSCEIN-25

NOVARTIS

25%

N17869 001

FLUOROMETHOLONE

CREAM; TOPICAL

OXYLONE

PHARMACIA AND UPJOHN

0.025%

N11748 001

FLUOROURACIL

INJECTABLE; INJECTION

ADRUCIL

PHARMACIA AND UPJOHN

50MG/ML

N17959 001

50MG/ML

N81222 001

Jun 28, 1991

SICOR PHARMS

50MG/ML

N40023 001

Oct 18, 1991

50MG/ML

N81225 001

Aug 28, 1991

FLUOROURACIL

ABIC

50MG/ML

N88929 001

Mar 04, 1986

ABRAXIS PHARM

50MG/ML

N89152 001

Mar 21, 1986

50MG/ML

N89428 001

Jan 12, 1987

50MG/ML

N89519 001

Mar 12, 1987

BEDFORD

50MG/ML

N89508 001

Jan 26, 1988

MARCHAR

50MG/ML

N87791 001

Jan 18, 1983

SMITH AND NEPHEW

50MG/ML

N88766 001

Dec 28, 1984

50MG/ML

N88767 001

Dec 28, 1984

50MG/ML

N89434 001

Mar 26, 1987

WATSON LABS

50MG/ML

N87792 001

Oct 13, 1982

DISCONTINUED DRUG PRODUCT LIST

6 - 114 (of 274)

FLUOROURACIL

SOLUTION; TOPICAL FLUOROPLEX				
ALLERGAN HERBERT	1%	N16765	001	

FLUOXETINE HYDROCHLORIDE

CAPSULE; ORAL PROZAC				
LILLY	EQ 60MG BASE	N18936	004	Jun 15, 1999
TABLET; ORAL PROZAC				
LILLY	EQ 10MG BASE	N20974	001	Mar 09, 1999
	EQ 20MG BASE **Federal Register determination that product was not discontinued or withdraw for safety or efficacy reasons**	N20974	002	Mar 09, 1999

FLUOXYMESTERONE

TABLET; ORAL ANDROID-F				
VALEANT PHARM INTL	10MG	N87196	001	
FLUOXYMESTERONE				
VALEANT PHARM INTL	10MG	N88221	001	May 05, 1983
WATSON LABS	2MG	N88260	001	Dec 06, 1983
	5MG	N88265	001	Dec 06, 1983
	10MG	N88309	001	Dec 06, 1983
ORA-TESTRYL				
BRISTOL MYERS SQUIBB	2MG	N11359	001	
	5MG	N11359	002	

FLUPHENAZINE DECANOATE

INJECTABLE; INJECTION FLUPHENAZINE DECANOATE				
MAYNE PHARMA USA	25MG/ML	N74966	001	Apr 16, 1998

FLUPHENAZINE ENANTHATE

INJECTABLE; INJECTION PROLIXIN ENANTHATE				
APOTHECON	25MG/ML	N16110	001	

FLUPHENAZINE HYDROCHLORIDE

CONCENTRATE; ORAL FLUPHENAZINE HYDROCHLORIDE				
TEVA PHARMS	5MG/ML	N73058	001	Aug 30, 1991
PERMITIL				
SCHERING	5MG/ML	N16008	001	
PROLIXIN				
APOTHECON	5MG/ML	N70533	001	Nov 07, 1985
ELIXIR; ORAL FLUPHENAZINE HYDROCHLORIDE				
TEVA PHARMS	2.5MG/5ML	N81310	001	Apr 29, 1993
PROLIXIN				
APOTHECON	2.5MG/5ML	N12145	003	
TABLET; ORAL FLUPHENAZINE HYDROCHLORIDE				
WATSON LABS	1MG	N88555	001	Dec 18, 1987
	2.5MG	N88544	001	Dec 18, 1987
	5MG	N88527	001	Dec 18, 1987
	10MG	N88550	001	Dec 18, 1987
PERMITIL				
SCHERING	0.25MG	N12034	001	
	2.5MG	N12034	004	
	5MG	N12034	005	
	10MG	N12034	006	

DISCONTINUED DRUG PRODUCT LIST

6 - 115 (of 274)

FLUPHENAZINE HYDROCHLORIDE

TABLET, EXTENDED RELEASE; ORAL
PERMITIL
SCHERING 1MG

N12419 004

FLUPREDNISOLONE

TABLET; ORAL
ALPHADROL
PHARMACIA AND UPJOHN 1.5MG

N12259 002

FLURANDRENOLIDE

LOTION; TOPICAL
FLURANDRENOLIDE
ALPHARMA US PHARMS 0.05%

N87203 001 Apr 29, 1982

FLURANDRENOLIDE; NEOMYCIN SULFATE

CREAM; TOPICAL
CORDRAN-N
LILLY 0.05%;EQ 3.5MG BASE/GM
OINTMENT; TOPICAL
CORDRAN-N
LILLY 0.05%;EQ 3.5MG BASE/GM

N50346 001

N50345 001

FLURAZEPAM HYDROCHLORIDE

CAPSULE; ORAL
FLURAZEPAM HYDROCHLORIDE
HALSEY 15MG
30MG
MUTUAL PHARM 15MG
30MG
PUREPAC PHARM 15MG
30MG
SUPERPHARM 15MG
30MG
USL PHARMA 15MG
30MG
WARNER CHILCOTT 15MG
30MG
WATSON LABS 15MG

N71808 001 Jan 07, 1988
N71809 001 Jan 07, 1988
N70454 001 Aug 04, 1986
N70455 001 Aug 04, 1986
N71927 001 Sep 09, 1987
N71551 001 Sep 09, 1987
N71659 001 Aug 04, 1988
N71660 001 Aug 04, 1988
N70562 001 Jul 09, 1987
N70563 001 Jul 09, 1987
N71767 001 Dec 04, 1987
N71768 001 Dec 04, 1987
N72368 001 Mar 30, 1989

FLURBIPROFEN

TABLET; ORAL
FLURBIPROFEN
IVAX PHARMS 50MG
100MG

N74411 001 May 31, 1995
N74411 002 May 31, 1995

FLUTAMIDE

CAPSULE; ORAL
EULEXIN
SCHERING 125MG

N18554 001 Jan 27, 1989

FLUTICASONE PROPIONATE

AEROSOL, METERED; INHALATION
FLOVENT
GLAXOSMITHKLINE 0.044MG/INH
0.11MG/INH
0.22MG/INH

N20548 001 Mar 27, 1996
N20548 002 Mar 27, 1996
N20548 003 Mar 27, 1996

FLUVOXAMINE MALEATE

TABLET; ORAL
LUVOX
SOLVAY 25MG
50MG

N20243 001 Dec 05, 1994
N20243 002 Dec 05, 1994

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 116 (of 274)

FLUVOXAMINE MALEATETABLET; ORAL
LUVOX

SOLVAY	100MG	N20243	003	Dec 05, 1994
	150MG	N20243	004	Dec 05, 1994

FOLIC ACIDINJECTABLE; INJECTION
FOLVITE

WYETH PHARMS INC	5MG/ML	N05897	008	
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TABLET; ORAL
FOLIC ACID

ANABOLIC	1MG	N84915	001	
BARR	1MG	N89177	001	Jan 08, 1986
EVERYLIFE	1MG	N80755	001	
HALSEY	1MG	N83598	001	
IMPAX LABS	1MG	N80686	001	
IVAX PHARMS	1MG	N83000	001	
LANNETT	1MG	N80816	001	
LILLY	1MG	N06135	003	
MK LABS	1MG	N83526	001	
PHARMERAL	1MG	N84158	001	
PIONEER PHARMS	1MG	N88949	001	Sep 13, 1985
PUREPAC PHARM	1MG	N80784	001	
SANDOZ	1MG	N84472	001	
UDL	1MG	N88199	001	Mar 29, 1983
USL PHARMA	1MG	N87828	001	May 13, 1982
VALEANT PHARM INTL	1MG	N80903	001	
VANGARD	1MG	N88730	001	Mar 23, 1984
VINTAGE PHARMS	1MG	N86296	001	
WATSON LABS	1MG	N83141	001	
	1MG	N85141	002	
WHITEWORTH TOWN PLSN	1MG	N80691	002	
FOLVITE				
WYETH PHARMS INC	1MG	N05897	004	

FOLLITROPIN ALFA/BETAINJECTABLE; IM-SC
FOLLISTIM

ORGANON USA INC	75 IU/VIAL	N20582	001	Sep 29, 1997
	150 IU/VIAL	N20582	002	Sep 29, 1997

INJECTABLE; SUBCUTANEOUS
FOLLISTIM AQ

ORGANON USA INC	150 IU/0.18ML	N21211	003	Feb 11, 2004
GONAL-F				
SERONO INC	37.5 IU/VIAL	N20378	003	May 25, 2000
	37.5 IU/VIAL	N21765	001	Mar 25, 2004
	75 IU/VIAL	N20378	001	Sep 29, 1997
	150 IU/VIAL	N20378	002	Sep 29, 1997
	150 IU/VIAL	N21765	003	Mar 25, 2004
	1,050 IU/VIAL	N20378	004	Feb 28, 2001

FOMIVIRSEN SODIUMINJECTABLE; INJECTION
VITRAVENE PRESERVATIVE FREE
NOVARTIS

6.6MG/ML	N20961	001	Aug 26, 1998
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FURAZOLIDONESUSPENSION; ORAL
FUROXONE

SHIRE	50MG/15ML	N11323	002	
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TABLET; ORAL
FUROXONE

SHIRE	100MG	N11270	002	
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DISCONTINUED DRUG PRODUCT LIST

6 - 117 (of 274)

FUROSEMIDE

INJECTABLE; INJECTION FUROSEMIDE

ABRAXIS PHARM	10MG/ML	N18507	001	Jul 30, 1982
	10MG/ML	N19036	001	Aug 13, 1984
ASTRAZENECA	10MG/ML	N70014	001	Sep 09, 1985
	10MG/ML	N70095	001	Sep 09, 1985
	10MG/ML	N70096	001	Sep 09, 1985
BAXTER HLTHCARE	10MG/ML	N18267	001	
MARSAM PHARMS LLC	10MG/ML	N74017	001	Jun 30, 1994
ORGANON USA INC	10MG/ML	N70017	001	Dec 15, 1986
SMITH AND NEPHEW	10MG/ML	N70023	001	Feb 05, 1986
	10MG/ML	N70078	001	Feb 05, 1986
WARNER CHILCOTT	10MG/ML	N18420	001	Feb 26, 1982
WATSON LABS	10MG/ML	N70019	001	Sep 22, 1986
	10MG/ML	N70604	001	Jan 02, 1987
WYETH AYERST	10MG/ML	N18670	001	Jul 20, 1982

LASIX

SANOFI AVENTIS US	10MG/ML	N16363	001	
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SOLUTION; ORAL

LASIX

SANOFI AVENTIS US	10MG/ML	N17688	001	
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TABLET; ORAL

FUROSEMIDE

CLONMEL HLTHCARE	20MG	N18415	001	Jul 27, 1982
	40MG	N18415	002	Jul 27, 1982
	80MG	N18415	003	Nov 26, 1984
KALAPHARM	20MG	N18868	001	Jun 28, 1983
	40MG	N18868	002	Jun 28, 1983
MUTUAL PHARM	20MG	N70043	001	Sep 26, 1985
	40MG	N18790	001	Nov 29, 1983
	80MG	N70100	001	Jan 26, 1988
SANDOZ	40MG	N18750	002	Jul 30, 1984
SUPERPHARM	20MG	N18370	002	Jun 26, 1984
	40MG	N18370	001	Feb 10, 1983
WARNER CHILCOTT	20MG	N18419	001	Jan 31, 1983
	40MG	N18419	002	Jan 31, 1983
	80MG	N18419	003	Nov 13, 1984
WATSON LABS	20MG	N18369	001	May 14, 1982
	40MG	N18369	002	May 14, 1982
	40MG	N70413	001	Feb 26, 1986

GALLAMINE TRIETHIODIDE

INJECTABLE; INJECTION FLAXEDIL

DAVIS AND GECK	20MG/ML	N07842	001	
	100MG/ML	N07842	002	

GALLIUM CITRATE, GA-67

INJECTABLE; INJECTION GALLIUM CITRATE GA 67

GE HEALTHCARE	1mCi/ML	N17700	001	
NEOSCAN				
GE HEALTHCARE	2mCi/ML	N17655	001	

GANCICLOVIR

CAPSULE; ORAL CYTOVENE

ROCHE PALO	250MG	N20460	001	Dec 22, 1994
	500MG	N20460	002	Dec 12, 1997

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 118 (of 274)

GATIFLOXACIN

INJECTABLE; INJECTION

TEQUIN

BRISTOL MYERS SQUIBB	EQ 10MG /ML(200MG)	N21062	003	Dec 17, 1999
	400MG/40ML(10MG/ML)	N21062	004	Dec 17, 1999

TEQUIN IN DEXTROSE 5% IN PLASTIC CONTAINER

BRISTOL MYERS SQUIBB	200MG/100ML(2MG/ML)	N21062	001	Dec 17, 1999
	400MG/200ML(2MG/ML)	N21062	002	Dec 17, 1999

SUSPENSION; ORAL

TEQUIN

BRISTOL MYERS SQUIBB	200MG/5ML	N21678	001	Aug 27, 2004
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TABLET; ORAL

TEQUIN

BRISTOL MYERS SQUIBB	200MG	N21061	001	Dec 17, 1999
	400MG	N21061	002	Dec 17, 1999

GEMFIBROZIL

CAPSULE; ORAL

GEMFIBROZIL

MYLAN	300MG	N73466	001	Jan 25, 1993
PUREPAC PHARM	300MG	N72929	001	Jan 29, 1993

LOPID

PFIZER PHARMS	200MG	N18422	001	
	300MG	N18422	002	

TABLET; ORAL

GEMFIBROZIL

CLONMEL HLTHCARE	600MG	N74270	001	Sep 27, 1993
PUREPAC PHARM	600MG	N74360	001	Aug 31, 1994
WATSON LABS	600MG	N74156	001	Oct 24, 1994

GENTAMICIN SULFATE

CREAM; TOPICAL

GARAMYCIN

SCHERING	EQ 0.1% BASE	N60462	001	
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GENTAFAIR

PHARMAFAIR	EQ 0.1% BASE	N62458	001	Sep 01, 1983
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GENTAMICIN SULFATE

ALPHARMA US PHARMS	EQ 0.1% BASE	N62471	001	Sep 27, 1983
BAUSCH AND LOMB	EQ 0.1% BASE	N64056	001	Apr 29, 1994
PHARMADERM	EQ 1MG BASE/GM	N62530	001	Jul 05, 1984

INJECTABLE; INJECTION

APOGEN

KING PHARMS	EQ 10MG BASE/ML	N62289	001	
	EQ 40MG BASE/ML	N62289	002	

BRISTAGEN

BRISTOL	EQ 40MG BASE/ML	N62288	001	
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GARAMYCIN

SCHERING	EQ 1MG BASE/ML	N61716	002	
	EQ 10MG BASE/ML	N61739	001	
	EQ 40MG BASE/ML	N61716	001	

GENTAFAIR

PHARMAFAIR	EQ 40MG BASE/ML	N62493	001	Aug 28, 1985
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GENTAMICIN

INTL MEDICATION	EQ 1MG BASE/ML	N62325	003	Jun 23, 1982
	EQ 40MG BASE/ML	N62325	001	
	EQ 100MG BASE/100ML	N62325	004	Jun 23, 1982

GENTAMICIN SULFATE

ABBOTT	EQ 1.2MG BASE/ML	N62413	001	Aug 11, 1983
	EQ 1.4MG BASE/ML	N62413	002	Aug 11, 1983
	EQ 1.6MG BASE/ML	N62413	003	Aug 11, 1983
	EQ 1.8MG BASE/ML	N62413	004	Aug 11, 1983
	EQ 2MG BASE/ML	N62413	005	Aug 11, 1983
	EQ 60MG BASE/100ML	N62413	006	Aug 11, 1983
	EQ 70MG BASE/100ML	N62413	007	Aug 11, 1983

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 119 (of 274)

GENTAMICIN SULFATE

INJECTABLE; INJECTION

GENTAMICIN SULFATE

ABBOTT

EQ 80MG BASE/100ML

N62413 008 Aug 11, 1983

EQ 90MG BASE/100ML

N62413 009 Aug 11, 1983

EQ 100MG BASE/100ML

N62413 010 Aug 11, 1983

BAXTER HLTHCARE

EQ 10MG BASE/ML

N62251 002

EQ 40MG BASE/ML

N62251 001

KALAPHARM

EQ 40MG BASE/ML

N62354 001 Apr 05, 1982

SICOR PHARMS

EQ 10MG BASE/ML

N63149 001 Nov 21, 1991

EQ 40MG BASE/ML

N63106 002 Nov 21, 1991

SOLOPAK

EQ 10MG BASE/ML

N62507 001 Jun 06, 1985

EQ 40MG BASE/ML

N62507 002 Jun 06, 1985

WATSON LABS

EQ 10MG BASE/ML

N62318 002

EQ 40MG BASE/ML

N62318 001

WYETH AYERST

EQ 10MG BASE/ML

N62264 001

EQ 40MG BASE/ML

N62264 002

GENTAMICIN SULFATE IN SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER

HOSPIRA

EQ 1.2MG BASE/ML

N62588 001 Jan 06, 1986

EQ 1.4MG BASE/ML

N62588 002 Jan 06, 1986

EQ 1.6MG BASE/ML

N62588 003 Jan 06, 1986

EQ 1.8MG BASE/ML

N62588 004 Jan 06, 1986

EQ 2MG BASE/ML

N62588 005 Jan 06, 1986

EQ 60MG BASE/100ML

N62588 006 Jan 06, 1986

EQ 70MG BASE/100ML

N62588 007 Jan 06, 1986

EQ 80MG BASE/100ML

N62588 008 Jan 06, 1986

EQ 90MG BASE/100ML

N62588 009 Jan 06, 1986

EQ 100MG BASE/100ML

N62588 010 Jan 06, 1986

U-GENCIN

PHARMACIA AND UPJOHN

EQ 10MG BASE/ML

N62248 001

EQ 40MG BASE/ML

N62248 002

INJECTABLE; INTRATHECAL

GARAMYCIN

SCHERING

EQ 2MG BASE/ML

N50505 001

OINTMENT; OPHTHALMIC

GARAMYCIN

SCHERING

EQ 0.3% BASE

N50425 001

GENTACIDIN

NOVARTIS

EQ 0.3% BASE

N62501 001 Jul 26, 1984

GENTAFAIR

PHARMAFAIR

EQ 3MG BASE/GM

N62443 001 May 26, 1983

OINTMENT; TOPICAL

GARAMYCIN

SCHERING

EQ 0.1% BASE

N60463 001

GENTAFAIR

PHARMAFAIR

EQ 0.1% BASE

N62444 001 May 26, 1983

GENTAMICIN SULFATE

ALPHARMA US PHARMS

EQ 0.1% BASE

N62496 001 Mar 14, 1984

BAUSCH AND LOMB

EQ 0.1% BASE

N64054 001 Apr 29, 1994

PHARMADERM

EQ 0.1% BASE

N62534 001 Oct 10, 1984

SOLUTION/DROPS; OPHTHALMIC

GARAMYCIN

SCHERING

EQ 0.3% BASE

N50039 002

GENTACIDIN

NOVARTIS

EQ 0.3% BASE

N62480 001 Mar 30, 1984

GENTAFAIR

PHARMAFAIR

EQ 0.3% BASE

N62440 001 May 03, 1983

GENTAMICIN SULFATE

ALCON UNIVERSAL

EQ 0.3% BASE

N62523 001 Nov 25, 1985

PACO

EQ 3MG BASE/ML

N62932 001 Nov 07, 1988

GENTIAN VIOLET

SUPPOSITORY; VAGINAL

GVS

SAVAGE LABS

0.4%

N83513 001

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 120 (of 274)

GENTIAN VIOLET

TAMPON; VAGINAL
GENAPAX
KEY PHARMS

5MG

N85017 001

GLIPIZIDE

TABLET; ORAL
GLIPIZIDE
ENDO PHARMS

5MG

N74378 001

Nov 28, 1994

10MG

N74378 002

Nov 28, 1994

GLUCOTROL
PFIZER

2.5MG

N17783 003

May 11, 1993

GLUCAGON HYDROCHLORIDE

INJECTABLE; INJECTION
GLUCAGON
LILLY

EQ 1MG BASE/VIAL

N12122 001

EQ 10MG BASE/VIAL

N12122 002

GLUTAMINE

FOR SOLUTION; ORAL
NUTRESTORE

NUTRITIONAL RESTART

5GM/PACKET

N21667 001

Jun 10, 2004

GLUTETHIMIDE

CAPSULE; ORAL
DORIDEN

SANOFI AVENTIS US

500MG

N09519 008

TABLET; ORAL
DORIDEN

SANOFI AVENTIS US

250MG

N09519 002

500MG

N09519 005

GLUTETHIMIDE
HALSEY

250MG

N89458 001

Oct 10, 1986

500MG

N89459 001

Oct 10, 1986

LANNETT

250MG

N83475 001

500MG

N85571 001

SANDOZ

500MG

N83234 002

UCB INC

500MG

N85171 001

VITARINE

500MG

N87297 001

WATSON LABS

500MG

N84362 001

500MG

N85763 001

GLYBURIDE

TABLET; ORAL
GLYBURIDE (MICRONIZED)
SANOFI AVENTIS US

1.5MG

N20055 001

Apr 17, 1992

3MG

N20055 002

Apr 17, 1992

6MG

N20055 003

Mar 08, 2000

GLYNASE

PHARMACIA AND UPJOHN

4.5MG

N20051 003

Sep 24, 1993

GLYCINE

SOLUTION; IRRIGATION
GLYCINE 1.5% IN PLASTIC CONTAINER
BAXTER HLTHCARE

1.5GM/100ML

N18522 001

Feb 19, 1982

GLYCOPYRROLATE

INJECTABLE; INJECTION
GLYCOPYRROLATE

ABRAXIS PHARM

0.2MG/ML

N88475 001

Jun 12, 1984

HOSPIRA

0.2MG/ML

N89393 001

Jun 15, 1988

SICOR PHARMS

0.2MG/ML

N81169 001

Sep 10, 1991

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 121 (of 274)

GLYCOPYRROLATEINJECTABLE; INJECTION
GLYCOPYRROLATE

WATSON LABS 0.2MG/ML

N86947 001 Jun 24, 1983

ROBINUL

ROBINS AH 0.2MG/ML

N14764 001

TABLET; ORAL

GLYCOPYRROLATE

WATSON LABS 1MG

N85562 001

1MG

N86902 001

2MG

N85563 001

2MG

N86178 001

2MG

N86900 001

GONADORELIN ACETATEINJECTABLE; INJECTION
LUTREPULSE KIT

FERRING 0.8MG/VIAL

N19687 001 Oct 10, 1989

3.2MG/VIAL

N19687 002 Oct 10, 1989

GONADORELIN HYDROCHLORIDEINJECTABLE; INJECTION
FACTREL

BAXTER HLTHCARE CORP EQ 0.1MG BASE/VIAL

N18123 001 Sep 30, 1982

EQ 0.2MG BASE/VIAL

N18123 002 Sep 30, 1982

EQ 0.5MG BASE/VIAL

N18123 003 Sep 30, 1982

GONADOTROPIN, CHORIONICINJECTABLE; INJECTION
A.P.L.

FERRING 5,000 UNITS/VIAL

N17055 001

10,000 UNITS/VIAL

N17055 002

20,000 UNITS/VIAL

N17055 003

CHORIONIC GONADOTROPIN

ABRAXIS PHARM 5,000 UNITS/VIAL

N17067 001

15,000 UNITS/VIAL

N17067 004

20,000 UNITS/VIAL

N17067 003

BEL MAR 5,000 UNITS/VIAL

N17054 001

10,000 UNITS/VIAL

N17054 002

WATSON LABS (UTAH) 2,000 UNITS/VIAL

N17016 009 Dec 27, 1984

2,000 UNITS/VIAL

N17016 011 Feb 16, 1990

5,000 UNITS/VIAL

N17016 006

15,000 UNITS/VIAL

N17016 010 Feb 15, 1985

20,000 UNITS/VIAL

N17016 004

FOLLUTEIN

BRISTOL MYERS SQUIBB 10,000 UNITS/VIAL

N17056 001

GRAMICIDIN; NEOMYCIN SULFATE; POLYMYXIN B SULFATE

SOLUTION/DROPS; OPHTHALMIC

NEOMYCIN AND POLYMYXIN B SULFATES AND GRAMICIDIN

IPHARM 0.025MG/ML;EQ 1.75MG BASE/ML;10,000 UNITS/ML

N62818 001 Oct 11, 1988

WATSON LABS 0.025MG/ML;EQ 1.75MG BASE/ML;10,000 UNITS/ML

N62788 001 Jun 11, 1987

NEOMYCIN SULFATE AND POLYMYXIN B SULFATE GRAMICIDIN

PHARMAFAIR 0.025MG/ML;EQ 1.75MG BASE/ML;10,000 UNITS/ML

N62383 001 Aug 31, 1982

NEO-POLYCIN

DOW PHARM 0.025MG/ML;EQ 1.75MG BASE/ML;10,000 UNITS/ML

N60427 001

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 122 (of 274)

GRANISETRON HYDROCHLORIDE

INJECTABLE; INJECTION

KYTRIL

ROCHE

EQ 3MG BASE/ML

N20239 001

Dec 29, 1993

TABLET; ORAL

KYTRIL

ROCHE

EQ 2MG BASE

N20305 002

Jun 15, 1998

GREPAFLOXACIN HYDROCHLORIDE

TABLET; ORAL

RAXAR

OTSUKA

EQ 200MG BASE

N20695 001

Nov 06, 1997

EQ 400MG BASE

N20695 002

May 14, 1998

EQ 600MG BASE

N20695 003

May 14, 1998

GRISEOFULVIN, MICROCRYSTALLINE

CAPSULE; ORAL

GRISACTIN

WYETH AYERST

125MG

N50051 002

250MG

N50051 001

SUSPENSION; ORAL

GRIFULVIN V

JOHNSON AND JOHNSON

125MG/5ML

N50448 001

TABLET; ORAL

GRIFULVIN V

J AND J

125MG

N60618 001

250MG

N60618 002

500MG

N60618 003

GRISACTIN

WYETH AYERST

500MG

N60212 001

GRISEOFULVIN, ULTRAMICROCRYSTALLINE

TABLET; ORAL

FULVICIN P/G

SCHERING

125MG

N61996 001

250MG

N61996 002

FULVICIN P/G 165

SCHERING

165MG

N61996 003

Apr 06, 1982

FULVICIN P/G 330

SCHERING

330MG

N61996 004

Apr 06, 1982

GRISACTIN ULTRA

WYETH AYERST

125MG

N62178 001

165MG

N62438 001

Nov 17, 1983

250MG

N62178 002

330MG

N62438 002

Nov 17, 1983

ULTRAGRIS-165

PLIVA

165MG

N62645 001

Jun 30, 1992

ULTRAGRIS-330

PLIVA

330MG

N62646 001

Jun 30, 1992

GUANABENZ ACETATE

TABLET; ORAL

WYTENSIN

WYETH AYERST

EQ 4MG BASE

N18587 001

Sep 07, 1982

EQ 8MG BASE

N18587 002

Sep 07, 1982

EQ 16MG BASE

N18587 003

Sep 07, 1982

GUANADREL SULFATE

TABLET; ORAL

HYLOREL

PHARMACIA AND UPJOHN

10MG

N18104 001

Dec 29, 1982

25MG

N18104 002

Dec 29, 1982

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 123 (of 274)

GUANETHIDINE MONOSULFATE

TABLET; ORAL

GUANETHIDINE MONOSULFATE

WATSON LABS

EQ 10MG SULFATE

N86113 001

Mar 26, 1985

EQ 25MG SULFATE

N86114 001

Mar 26, 1985

ISMELIN

NOVARTIS

EQ 10MG SULFATE

N12329 001

EQ 25MG SULFATE

N12329 002

GUANETHIDINE MONOSULFATE; HYDROCHLOROTHIAZIDE

TABLET; ORAL

ESIMIL

NOVARTIS

10MG;25MG

N13553 001

GUANFACINE HYDROCHLORIDE

TABLET; ORAL

TENEX

DR REDDYS LABS INC

EQ 3MG BASE

N19032 003

Nov 07, 1988

HALAZEPAM

TABLET; ORAL

PAXIPAM

SCHERING

20MG

N17736 003

40MG

N17736 004

HALCINONIDE

CREAM; TOPICAL

HALOG

WESTWOOD SQUIBB

0.025%

N17818 001

HALOG-E

WESTWOOD SQUIBB

0.1%

N18234 001

OINTMENT; TOPICAL

HALOG

BRISTOL MYERS SQUIBB

0.025%

N18125 001

HALOFANTRINE HYDROCHLORIDE

TABLET; ORAL

HALFAN

GLAXOSMITHKLINE

250MG

N20250 001

Jul 24, 1992

HALOPERIDOL

TABLET; ORAL

HALDOL

ORTHO MCNEIL

0.5MG

N15921 001

1MG

N15921 002

2MG

N15921 003

5MG

N15921 004

10MG

N15921 005

20MG

N15921 006

Feb 02, 1982

HALDOL SOLUTAB

ORTHO MCNEIL PHARM

1MG

N17079 001

HALOPERIDOL

DURAMED PHARMS BARR

0.5MG

N71216 001

Dec 04, 1986

1MG

N71217 001

Dec 04, 1986

2MG

N71218 001

Dec 04, 1986

5MG

N71219 001

Dec 04, 1986

10MG

N71220 001

Jul 07, 1987

20MG

N71221 001

Jul 07, 1987

LEDERLE

0.5MG

N72727 001

Sep 19, 1989

1MG

N72728 001

Sep 19, 1989

2MG

N72729 001

Sep 19, 1989

5MG

N72730 001

Sep 19, 1989

10MG

N72731 001

Sep 19, 1989

20MG

N72732 001

Sep 19, 1989

DISCONTINUED DRUG PRODUCT LIST

6 - 124 (of 274)

HALOPERIDOL

TABLET; ORAL

HALOPERIDOL

MUTUAL PHARM	0.5MG	N71156	001	Jan 02, 1987
	1MG	N71157	001	Jan 02, 1987
	2MG	N71172	001	Jan 02, 1987
	5MG	N71212	001	Jan 07, 1988
	10MG	N71173	001	Jan 07, 1988
	20MG	N71177	001	Jan 07, 1988
PAR PHARM	20MG	N71328	001	Jul 20, 1987
PUREPAC PHARM	0.5MG	N71071	001	Nov 03, 1986
	1MG	N71072	001	Nov 03, 1986
	2MG	N71073	001	Nov 03, 1986
	5MG	N71074	001	Nov 03, 1986
	10MG	N71075	001	Aug 04, 1987
	20MG	N71076	001	Aug 04, 1987
QUANTUM PHARMICS	0.5MG	N71255	001	Feb 17, 1987
	1MG	N71269	001	Feb 17, 1987
	2MG	N71256	001	Feb 17, 1987
	5MG	N71257	001	Feb 17, 1987
ROXANE	0.5MG	N71128	001	Feb 17, 1987
	1MG	N71129	001	Feb 17, 1987
	2MG	N71130	001	Feb 17, 1987
	5MG	N71131	001	Feb 17, 1987
	10MG	N71132	001	May 12, 1987
	20MG	N71133	001	May 12, 1987
ROYCE LABS	0.5MG	N71722	001	Dec 24, 1987
	1MG	N71723	001	Dec 24, 1987
	2MG	N71724	001	Dec 24, 1987
	5MG	N71725	001	Dec 24, 1987
	10MG	N72121	001	Dec 24, 1987
	20MG	N72122	001	Dec 24, 1987
SCS	0.5MG	N70720	001	Jun 10, 1986
	1MG	N70721	001	Jun 10, 1986
	2MG	N70722	001	Jun 10, 1986
	5MG	N70723	001	Jun 10, 1986
	10MG	N70724	001	Jun 10, 1986
	20MG	N70725	001	Sep 24, 1986
WATSON LABS	0.5MG	N71571	001	Jun 03, 1988
	1MG	N70982	001	Mar 06, 1987
	1MG	N71572	001	Jun 03, 1988
	2MG	N71573	001	Jun 03, 1988
	5MG	N71374	001	Jun 03, 1988
	10MG	N71375	001	Jun 03, 1988
	10MG	N72113	001	Aug 27, 1991
	20MG	N71376	001	Jun 03, 1988
	20MG	N72353	001	Aug 27, 1991

HALOPERIDOL DECANOATE

INJECTABLE; INJECTION

HALOPERIDOL DECANOATE

MAYNE PHARMA USA	EQ 50MG BASE/ML	N75176	001	Feb 09, 2000
	EQ 100MG BASE/ML	N75176	002	Feb 09, 2000

HALOPERIDOL LACTATE

CONCENTRATE; ORAL

HALDOL

ORTHO MCNEIL	EQ 2MG BASE/ML	N15922	001	
HALOPERIDOL				
ALPHARMA	EQ 2MG BASE/ML	N70318	001	Apr 11, 1986
MORTON GROVE	EQ 2MG BASE/ML	N70710	001	Mar 07, 1986
SCS	EQ 2MG BASE/ML	N70726	001	Jun 10, 1986
TEVA	EQ 2MG BASE/ML	N71015	001	Aug 25, 1987

DISCONTINUED DRUG PRODUCT LIST

6 - 125 (of 274)

HALOPERIDOL LACTATE

CONCENTRATE; ORAL					
HALOPERIDOL INTENSOL					
ROXANE	EQ 2MG BASE/ML		N72045	001	Apr 12, 1988
INJECTABLE; INJECTION					
HALOPERIDOL					
ABRAXIS PHARM	EQ 5MG BASE/ML		N71187	001	Jan 20, 1987
MARSAM PHARMS LLC	EQ 5MG BASE/ML		N72516	001	Feb 25, 1993
	EQ 5MG BASE/ML		N72517	001	Feb 25, 1993
SMITH AND NEPHEW	EQ 5MG BASE/ML		N70802	001	Dec 14, 1987
SOLOPAK	EQ 5MG BASE/ML		N70800	001	Dec 14, 1987
	EQ 5MG BASE/ML		N70801	001	Dec 14, 1987
	EQ 5MG BASE/ML		N70864	001	Dec 14, 1987
WATSON LABS	EQ 5MG BASE/ML		N70713	001	May 17, 1988
	EQ 5MG BASE/ML		N70744	001	May 17, 1988

HALOPROGIN

CREAM; TOPICAL					
HALOTEX					
WESTWOOD SQUIBB	1%		N16942	001	
SOLUTION; TOPICAL					
HALOTEX					
WESTWOOD SQUIBB	1%		N16943	001	

HALOTHANE

LIQUID; INHALATION					
FLUOTHANE					
WYETH AYERST	99.99%		N11338	001	
HALOTHANE					
BH	99.99%		N84977	001	
HALOCARBON	99.99%		N80810	001	
HOSPIRA	99.99%		N83254	001	

HEPARIN CALCIUM

INJECTABLE; INJECTION					
CALCIPARINE					
SANOFI AVENTIS US	25,000 UNITS/ML		N18237	001	

HEPARIN SODIUM

INJECTABLE; INJECTION					
HEPARIN LOCK FLUSH					
HOSPIRA	100 UNITS/ML		N05264	010	
INTL MEDICATION	10 UNITS/ML		N86357	001	
	500 UNITS/ML		N86357	002	
	10 UNITS/ML		N89063	001	Oct 09, 1985
	100 UNITS/ML		N89064	001	Oct 09, 1985
PARKE DAVIS	10 UNITS/ML		N17346	006	
SMITH AND NEPHEW	10 UNITS/ML		N87904	001	Apr 20, 1983
	10 UNITS/ML		N87958	001	Apr 20, 1983
	10 UNITS/ML		N88458	001	Jul 26, 1984
	10 UNITS/ML		N88580	001	Oct 25, 1984
	100 UNITS/ML		N87906	001	Apr 20, 1983
	100 UNITS/ML		N87959	001	Apr 20, 1983
	100 UNITS/ML		N88460	001	Jul 26, 1984
	100 UNITS/ML		N88581	001	Oct 25, 1984
	10 UNITS/ML		N87903	001	Apr 20, 1983
	10 UNITS/ML		N88457	001	Oct 25, 1984
SOLOPAK	100 UNITS/ML		N87905	001	Apr 20, 1983
	100 UNITS/ML		N88459	001	Jul 26, 1984
HEPARIN SODIUM					
ABRAXIS PHARM	1,000 UNITS/ML		N17033	001	
	1,000 UNITS/ML		N17651	005	
	5,000 UNITS/ML		N17029	002	
	5,000 UNITS/ML		N17979	003	

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 126 (of 274)

HEPARIN SODIUM

INJECTABLE; INJECTION

HEPARIN SODIUM

ABRAXIS PHARM	10,000 UNITS/ML	N17651	003	
	20,000 UNITS/ML	N17651	008	
AKORN	1,000 UNITS/ML	N17486	001	
	5,000 UNITS/ML	N17486	002	
	10,000 UNITS/ML	N17486	003	
	20,000 UNITS/ML	N17486	004	
	40,000 UNITS/ML	N17486	005	
BAXTER HLTHCARE CORP	1,000 UNITS/ML	N17007	001	
	2,500 UNITS/ML	N17007	007	
	5,000 UNITS/ML	N17007	002	
	5,000 UNITS/0.5ML	N17007	010	
	7,500 UNITS/ML	N17007	003	
	10,000 UNITS/ML	N17007	004	
	15,000 UNITS/ML	N17007	005	
	20,000 UNITS/ML	N17007	006	
CHAMBERLIN PARENTERL	1,000 UNITS/ML	N17130	001	
	5,000 UNITS/ML	N17130	002	
	10,000 UNITS/ML	N17130	003	
	20,000 UNITS/ML	N17130	004	
DELL LABS	1,000 UNITS/ML	N17540	001	
	5,000 UNITS/ML	N17540	002	
	10,000 UNITS/ML	N17540	003	
	20,000 UNITS/ML	N17540	004	
	40,000 UNITS/ML	N17540	005	
HOSPIRA	2,500 UNITS/ML	N88099	001	Apr 28, 1983
	10,000 UNITS/ML	N40095	001	Jul 26, 1996
LILLY	1,000 UNITS/ML	N05521	001	
	10,000 UNITS/ML	N05521	002	
	20,000 UNITS/ML	N05521	004	
	1,000 UNITS/ML	N87452	001	Oct 31, 1983
MARSAM PHARMS LLC	1,000 UNITS/ML	N40007	001	Jun 07, 1996
ORGANON USA INC	1,000 UNITS/ML	N00552	008	
	5,000 UNITS/ML	N00552	009	
	10,000 UNITS/ML	N00552	010	
	1,000 UNITS/ML	N17346	001	
PARKE DAVIS	5,000 UNITS/ML	N17346	002	
	7,500 UNITS/ML	N17346	003	
	10,000 UNITS/ML	N17346	004	
	20,000 UNITS/ML	N17346	005	
	1,000 UNITS/ML	N17780	001	
PHARM SPEC	5,000 UNITS/ML	N17780	002	
	10,000 UNITS/ML	N17780	003	
	20,000 UNITS/ML	N17780	004	
	40,000 UNITS/ML	N17780	005	
	1,000 UNITS/ML	N88239	001	Jul 26, 1984
SMITH AND NEPHEW SOLOPAK	1,000 UNITS/ML	N87043	001	
	5,000 UNITS/ML	N87077	001	
	5,000 UNITS/0.5ML	N87395	001	
	10,000 UNITS/ML	N87107	001	
	10,000 UNITS/0.5ML	N87363	001	
	1,000 UNITS/ML	N17064	002	
WATSON PHARMS	2,500 UNITS/ML	N17064	015	
	3,000 UNITS/ML	N17064	016	
	4,000 UNITS/ML	N17064	017	
	5,000 UNITS/ML	N17064	003	
	6,000 UNITS/ML	N17064	018	
	7,500 UNITS/ML	N17064	019	
	10,000 UNITS/ML	N17064	004	
	20,000 UNITS/ML	N17064	005	
	40,000 UNITS/ML	N17064	006	

DISCONTINUED DRUG PRODUCT LIST

6 - 127 (of 274)

HEPARIN SODIUM

INJECTABLE; INJECTION

HEPARIN SODIUM 1,000 UNITS IN DEXTROSE 5% IN PLASTIC CONTAINER				
MCGAW	200 UNITS/100ML	N19130	001	Dec 31, 1984
HEPARIN SODIUM 1,000 UNITS IN SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER				
B BRAUN	200 UNITS/100ML	N19042	001	Mar 29, 1985
HEPARIN SODIUM 10,000 UNITS IN DEXTROSE 5%				
HOSPIRA	10,000 UNITS/100ML	N18911	006	Jan 30, 1985
HEPARIN SODIUM 10,000 UNITS IN DEXTROSE 5% IN PLASTIC CONTAINER				
BAXTER HLTHCARE	2,000 UNITS/100ML	N18814	002	Jul 09, 1985
HEPARIN SODIUM 10,000 UNITS IN SODIUM CHLORIDE 0.45%				
HOSPIRA	10,000 UNITS/100ML	N18911	001	Jan 30, 1985
	10,000 UNITS/100ML	N18916	005	Jan 31, 1984
HEPARIN SODIUM 10,000 UNITS IN SODIUM CHLORIDE 0.9%				
HOSPIRA	10,000 UNITS/100ML	N18911	003	Jan 30, 1985
	10,000 UNITS/100ML	N18916	002	Jan 31, 1984
HEPARIN SODIUM 12,500 UNITS IN DEXTROSE 5%				
HOSPIRA	5,000 UNITS/100ML	N18911	007	Jan 30, 1985
HEPARIN SODIUM 12,500 UNITS IN SODIUM CHLORIDE 0.45% IN PLASTIC CONTAINER				
B BRAUN	5,000 UNITS/100ML	N19802	001	Jul 20, 1992
HEPARIN SODIUM 12,500 UNITS IN SODIUM CHLORIDE 0.9%				
HOSPIRA	5,000 UNITS/100ML	N18911	005	Jan 30, 1985
	5,000 UNITS/100ML	N18916	003	Jan 31, 1984
HEPARIN SODIUM 2,000 UNITS IN DEXTROSE 5% IN PLASTIC CONTAINER				
MCGAW	200 UNITS/100ML	N19130	003	Dec 31, 1984
HEPARIN SODIUM 2,000 UNITS IN SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER				
B BRAUN	200 UNITS/100ML	N19042	002	Mar 29, 1985
HEPARIN SODIUM 25,000 UNITS IN DEXTROSE 5%				
HOSPIRA	5,000 UNITS/100ML	N18911	009	Jan 30, 1985
	10,000 UNITS/100ML	N18911	008	Jan 30, 1985
HEPARIN SODIUM 25,000 UNITS IN DEXTROSE 5% IN PLASTIC CONTAINER				
B BRAUN	5,000 UNITS/100ML	N19134	001	Mar 29, 1985
HEPARIN SODIUM 25,000 UNITS IN SODIUM CHLORIDE 0.45% IN PLASTIC CONTAINER				
B BRAUN	5,000 UNITS/100ML	N19802	005	Jul 20, 1992
	10,000 UNITS/100ML	N19802	002	Jul 20, 1992
HEPARIN SODIUM 25,000 UNITS IN SODIUM CHLORIDE 0.9%				
HOSPIRA	5,000 UNITS/100ML	N18911	004	Jan 30, 1985
HEPARIN SODIUM 25,000 UNITS IN SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER				
B BRAUN	5,000 UNITS/100ML	N19135	001	Mar 29, 1985
	5,000 UNITS/100ML	N19802	003	Jul 20, 1992
HOSPIRA	5,000 UNITS/100ML	N18916	009	Jan 31, 1984
HEPARIN SODIUM 5,000 UNITS AND SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER				
BAXTER HLTHCARE	500 UNITS/100ML	N18609	003	Apr 28, 1982
HEPARIN SODIUM 5,000 UNITS IN DEXTROSE 5% IN PLASTIC CONTAINER				
MCGAW	1,000 UNITS/100ML	N19130	002	Dec 31, 1984
HEPARIN SODIUM 5,000 UNITS IN SODIUM CHLORIDE 0.45%				
HOSPIRA	100 UNITS/ML	N18911	002	Jan 30, 1985
	100 UNITS/ML	N18916	004	Jan 31, 1984
HEPARIN SODIUM 5,000 UNITS IN SODIUM CHLORIDE 0.9%				
HOSPIRA	1,000 UNITS/100ML	N18916	001	Jan 31, 1984
HEPARIN SODIUM 5,000 UNITS IN SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER				
B BRAUN	1,000 UNITS/100ML	N19042	004	Mar 29, 1985
HEPARIN SODIUM PRESERVATIVE FREE				
MARSAM PHARMS LLC	1,000 UNITS/ML	N89464	001	Jun 03, 1986
PHARMA SERVE NY	1,000 UNITS/ML	N86129	001	
LIPO-HEPIN				
3M	1,000 UNITS/0.5ML	N17027	001	
	1,000 UNITS/ML	N17027	006	
	5,000 UNITS/0.5ML	N17027	002	
	5,000 UNITS/ML	N17027	008	
	7,500 UNITS/0.5ML	N17027	010	
	10,000 UNITS/0.5ML	N17027	003	
	10,000 UNITS/ML	N17027	009	
	15,000 UNITS/0.5ML	N17027	011	

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 128 (of 274)

HEPARIN SODIUMINJECTABLE; INJECTION
LIPO-HEPIN

3M	20,000 UNITS/0.5ML	N17027	004
	20,000 UNITS/ML	N17027	007
	40,000 UNITS/ML	N17027	005

LIQUAEMIN LOCK FLUSH

ORGANON USA INC	100 UNITS/ML	N00552	007
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LIQUAEMIN SODIUM

ORGANON USA INC	1,000 UNITS/ML	N00552	004
	5,000 UNITS/ML	N00552	003
	10,000 UNITS/ML	N00552	005
	20,000 UNITS/ML	N00552	001
	40,000 UNITS/ML	N00552	002

LIQUAEMIN SODIUM PRESERVATIVE FREE

ORGANON USA INC	1,000 UNITS/ML	N00552	011	Apr 11, 1986
	5,000 UNITS/ML	N00552	012	Apr 11, 1986
	10,000 UNITS/ML	N00552	013	Apr 11, 1986

PANHEPRIN

HOSPIRA	1,000 UNITS/ML	N05264	004
	5,000 UNITS/ML	N05264	006
	10,000 UNITS/ML	N05264	007
	20,000 UNITS/ML	N05264	008
	40,000 UNITS/ML	N05264	009

SODIUM HEPARIN

ABRAXIS PHARM	5,000 UNITS/ML	N17033	002
	10,000 UNITS/ML	N17033	003
	20,000 UNITS/ML	N17033	004

BAXTER HLTHCARE	1,000 UNITS/ML	N17036	001	Mar 04, 1988
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HETACILLINFOR SUSPENSION; ORAL
VERSAPEN

BRISTOL	EQ 112.5MG AMPICIL/5ML	N50060	001
	EQ 112.5MG AMPICIL/ML	N50060	003
	EQ 112.5MG AMPICIL/ML	N61398	001
	EQ 225MG AMPICIL/5ML	N61398	002

HETACILLIN POTASSIUMCAPSULE; ORAL
VERSAPEN-K

BRISTOL	EQ 225MG AMPICIL	N61396	001
	EQ 450MG AMPICIL	N61396	002

HEXACHLOROPHENEAEROSOL; TOPICAL
SEPTISOL

VESTAL LABS	0.23%	N17424	001
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TURGEX

XTTRIUM	3%	N18375	001
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EMULSION; TOPICAL

HEXA-GERM

HUNTINGTON LABS	3%	N17411	001
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PHISOHEX

SANOFI AVENTIS US	3%	N08402	001
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SOY-DOME

BAYER PHARMS	3%	N17405	001
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TURGEX

XTTRIUM	3%	N19055	001	Nov 30, 1984
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SOAP; TOPICAL

GAMOPHEN

ARBROOK	2%	N06270	003
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27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 129 (of 274)

HEXACHLOROPHENESOLUTION; TOPICAL
DIAL

DIAL 0.25% N17421 002

GERMA-MEDICA

HUNTINGTON LABS 1% N17412 001

GERMA-MEDICA "MG"

HUNTINGTON LABS 0.25% N17412 002

SEPTI-SOFT

CALGON 0.25% N17460 001

SEPTISOL

VESTAL LABS 0.25% N17423 001

SPONGE; TOPICAL

E-Z SCRUB

BECTON DICKINSON 450MG N17452 001

HEXASCRUB

PROF DSPLS 3% N18363 001

PHISO-SCRUB

SANOFI AVENTIS US 3% N17446 001

SCRUBTEAM SURGICAL SPONGEBRUSH

3M 330MG N17413 001

HEXAFLUORENIUM BROMIDEINJECTABLE; INJECTION
MYLAXEN

MEDPOINTE PHARM HLC 20MG/ML N09789 003

HEXOCYCLIUM METHYLSULFATETABLET; ORAL
TRAL

ABBOTT 25MG N10599 001

HEXYLCAINE HYDROCHLORIDESOLUTION; TOPICAL
CYCLAINE

MERCK 5% N08472 001

HISTAMINE PHOSPHATEINJECTABLE; INJECTION
HISTAMINE PHOSPHATE

LILLY EQ 0.1MG BASE/ML N00734 003

EQ 0.2MG BASE/ML N00734 002

EQ 1MG BASE/ML N00734 001

HISTRELIN ACETATEINJECTABLE; INJECTION
SUPPRELIN

SHIRE EQ 0.2MG BASE/ML N19836 001 Dec 24, 1991

EQ 0.5MG BASE/ML N19836 002 Dec 24, 1991

EQ 1MG BASE/ML N19836 003 Dec 24, 1991

HOMATROPINE METHYLBROMIDE

TABLET; ORAL

HOMAPIN-10

MISSION PHARMA 10MG N86308 001

HOMAPIN-5

MISSION PHARMA 5MG N86309 001

TABLET, CHEWABLE; ORAL
EQUIPIN

MISSION PHARMA 3MG N86310 001

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 130 (of 274)

HOMATROPINE METHYLBROMIDE; HYDROCODONE BITARTRATE

SYRUP; ORAL				
HYDRODANE				
HALSEY	1.5MG/5ML;5MG/5ML	N88066	001	Jun 28, 1985

HYALURONIDASE

INJECTABLE; INJECTION				
WYDASE				
BAXTER HLTHCARE	150 UNITS/ML	N06343	002	
	150 UNITS/VIAL **Federal Register determination that product was not discontinued or withdrawn for safety or efficacy reasons**	N06343	006	
	1,500 UNITS/VIAL **Federal Register determination that product was not discontinued or withdrawn for safety or efficacy reasons**	N06343	005	

HYDRALAZINE HYDROCHLORIDE

INJECTABLE; INJECTION				
APRESOLINE				
NOVARTIS	20MG/ML	N08303	003	
HYDRALAZINE HYDROCHLORIDE				
ABRAXIS PHARM	20MG/ML	N89532	001	Aug 11, 1987
SICOR PHARMS	20MG/ML	N40373	001	Feb 23, 2000
SMITH AND NEPHEW	20MG/ML	N88518	001	Apr 20, 1984
SOLOPAK	20MG/ML	N88517	001	Aug 22, 1985
TABLET; ORAL				
APRESOLINE				
NOVARTIS	10MG	N08303	004	
	25MG	N08303	001	
	50MG	N08303	002	
	100MG	N08303	005	
DRALZINE				
TEVA	25MG	N84301	001	
HYDRALAZINE HYDROCHLORIDE				
ACTAVIS TOTOWA	25MG	N88560	001	Oct 04, 1984
	50MG	N88649	001	Oct 18, 1984
ASCOT	25MG	N88310	001	Dec 19, 1984
	50MG	N88311	001	Dec 19, 1984
HALSEY	10MG	N89218	001	Jan 22, 1986
	25MG	N89130	001	Jan 15, 1986
	50MG	N89222	001	Jan 22, 1986
	100MG	N89178	001	Jan 15, 1986
IMPAX LABS	25MG	N84922	001	
	50MG	N84923	001	
IVAX PHARMS	10MG	N84443	001	
	25MG	N84437	001	
	50MG	N84469	002	
	100MG	N84581	001	
MUTUAL PHARM	10MG	N88728	001	Apr 11, 1985
	25MG	N84106	002	
	50MG	N84107	002	
	100MG	N88729	001	Apr 11, 1985
PUREPAC PHARM	25MG	N88177	001	Jul 29, 1983
	50MG	N88178	001	Aug 15, 1983
QUANTUM PHARMICS	10MG	N88671	001	May 01, 1984
	25MG	N88657	001	Jun 15, 1984
	50MG	N88652	001	May 08, 1984
	100MG	N88686	001	May 01, 1984
RADIUS PHARMS	25MG	N86243	001	
	50MG	N86242	002	
SANDOZ	50MG	N85088	001	
SUPERPHARM	10MG	N88787	001	Aug 28, 1984
	25MG	N88788	001	Aug 28, 1984

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 131 (of 274)

HYDRALAZINE HYDROCHLORIDE

TABLET; ORAL

HYDRALAZINE HYDROCHLORIDE

SUPERPHARM	50MG	N88789	001	Aug 28, 1984
TG UNITED LABS	10MG	N88846	001	Feb 26, 1985
	25MG	N88847	001	Feb 26, 1985
	50MG	N88848	001	Feb 26, 1985
	100MG	N88849	001	Feb 26, 1985
USL PHARMA	25MG	N87780	001	Mar 29, 1982
	50MG	N87751	001	Mar 29, 1982
VANGARD	25MG	N87712	001	
	50MG	N87908	001	May 07, 1982
VITARINE	25MG	N86088	001	
WATSON LABS	25MG	N85532	002	May 24, 1982
	50MG	N85533	002	May 25, 1982
WEST WARD	25MG	N88240	001	May 27, 1983
	50MG	N88241	001	May 27, 1983

HYDRALAZINE HYDROCHLORIDE; HYDROCHLOROTHIAZIDE

CAPSULE; ORAL

APRESAZIDE

NOVARTIS 100MG;50MG

N84811 001

HYDRALAZINE HYDROCHLORIDE AND HYDROCHLOROTHIAZIDE

SOLVAY 25MG;25MG

N87608 001 Feb 08, 1982

50MG;50MG

N87213 001 Feb 08, 1982

100MG;50MG

N87609 001 Feb 08, 1982

SUPERPHARM 25MG;25MG

N89200 001 Feb 09, 1987

50MG;50MG

N89201 001 Feb 09, 1987

WATSON LABS 25MG;25MG

N85457 001 Mar 04, 1982

50MG;50MG

N85446 001 Mar 04, 1982

100MG;50MG

N85440 001 Mar 04, 1982

HYDRALAZINE HYDROCHLORIDE W/ HYDROCHLOROTHIAZIDE 100/50

IVAX PHARMS 100MG;50MG

N88358 001 Apr 10, 1984

HYDRALAZINE HYDROCHLORIDE W/ HYDROCHLOROTHIAZIDE 25/25

IVAX PHARMS 25MG;25MG

N88356 001 Apr 10, 1984

HYDRALAZINE HYDROCHLORIDE W/ HYDROCHLOROTHIAZIDE 50/50

IVAX PHARMS 50MG;50MG

N88357 001 Apr 10, 1984

TABLET; ORAL

APRESOLINE-ESIDRIX

NOVARTIS 25MG;15MG

N12026 002

HYDRALAZINE AND HYDROCHLOROTHIAZIDE

WATSON LABS 25MG;15MG

N85827 001

HYDROCHLOROTHIAZIDE W/ HYDRALAZINE

WATSON LABS 25MG;15MG

N85373 001

HYDRALAZINE HYDROCHLORIDE; HYDROCHLOROTHIAZIDE; RESERPINE

TABLET; ORAL

CAM-AP-ES

TG UNITED LABS 25MG;15MG;0.1MG

N84897 001

HYDRALAZINE HYDROCHLORIDE, HYDROCHLOROTHIAZIDE AND RESERPINE

IVAX PHARMS 25MG;15MG;0.1MG

N84291 001

HYDRALAZINE HYDROCHLORIDE-HYDROCHLOROTHIAZIDE-RESERPINE

MYLAN 25MG;15MG;0.1MG

N87085 001

HYDRALAZINE, HYDROCHLOROTHIAZIDE W/ RESERPINE

WATSON LABS 25MG;15MG;0.1MG

N85771 001

HYDRAP-ES

SANDOZ 25MG;15MG;0.1MG

N84876 001

HYDROCHLOROTHIAZIDE W/ RESERPINE AND HYDRALAZINE

WATSON LABS 25MG;15MG;0.1MG

N83770 001

HYDROSERPINE PLUS (R-H-H)

IVAX PHARMS 25MG;15MG;0.1MG

N83877 001

RESERPINE, HYDRALAZINE HYDROCHLORIDE AND HYDROCHLOROTHIAZIDE

MUTUAL PHARM 25MG;15MG;0.1MG

N88570 001 Apr 10, 1984

SOLVAY 25MG;15MG;0.1MG

N88376 001 Oct 28, 1983

DISCONTINUED DRUG PRODUCT LIST

6 - 132 (of 274)

HYDRALAZINE HYDROCHLORIDE; HYDROCHLOROTHIAZIDE; RESERPINE

TABLET; ORAL

RESERPINE, HYDRALAZINE HYDROCHLORIDE AND HYDROCHLOROTHIAZIDE

WATSON LABS 25MG;15MG;0.1MG N85549 001

25MG;15MG;0.1MG N87556 001

RESERPINE, HYDROCHLOROTHIAZIDE, AND HYDRALAZINE HYDROCHLORIDE

LEDERLE 25MG;15MG;0.1MG N87709 001

May 13, 1982

SER-A-GEN

SOLVAY 25MG;15MG;0.1MG N87210 001

SER-AP-ES

NOVARTIS 25MG;15MG;0.1MG N12193 005

UNIPRES

SOLVAY 25MG;15MG;0.1MG N85893 001

25MG;15MG;0.1MG N86298 001

HYDRALAZINE HYDROCHLORIDE; RESERPINE

TABLET; ORAL

DRALSERP

SANDOZ 25MG;0.1MG N84617 001

SERPASIL-APRESOLINE

NOVARTIS 25MG;0.1MG N09296 004

50MG;0.2MG N09296 002

HYDROCHLOROTHIAZIDE

SOLUTION; ORAL

HYDROCHLOROTHIAZIDE

MORTON GROVE 50MG/5ML N89661 001

Jun 20, 1988

HYDROCHLOROTHIAZIDE INTENSOL

ROXANE 100MG/ML N88588 001

Jul 02, 1984

TABLET; ORAL

ESIDRIX

NOVARTIS 100MG N11793 009

HYDROCHLOROTHIAZIDE

ABC HOLDING 50MG N85672 001

ALRA 25MG N86369 001

50MG N83554 001

ASCOT 25MG N87539 001

Feb 03, 1982

50MG N87540 001

Feb 03, 1982

50MG N84771 001

BARR 50MG N87060 001

CLONMEL HLTHCARE 100MG N85152 002

ELKINS SINN 50MG N84135 001

HEATHER 50MG N84029 001

IMPAX LABS 25MG N83607 002

50MG N85098 001

100MG N84776 001

INWOOD LABS 25MG N85067 001

25MG N84776 002

50MG N84658 001

IVAX PHARMS 50MG N85022 001

100MG N86597 001

LEINER 50MG N86192 001

MAST MM 25MG N86192 002

50MG N83972 001

MUTUAL PHARM 25MG N83972 002

50MG N83972 003

100MG N84880 001

MYLAN 25MG N85112 001

50MG N84325 001

PHARMERAL 25MG N84324 001

50MG N85004 001

ROXANE 25MG N84536 002

50MG N85005 001

50MG N83899 001

SANDOZ 25MG

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 133 (of 274)

HYDROCHLOROTHIAZIDE

TABLET; ORAL

HYDROCHLOROTHIAZIDE

SOLVAY	25MG	N85323	001	
SUPERPHARM	25MG	N88827	001	Dec 28, 1984
	50MG	N88828	001	Dec 28, 1984
	100MG	N88829	001	Dec 28, 1984
	25MG	N88924	001	Feb 07, 1985
TEVA	50MG	N88923	001	Feb 07, 1985
	25MG	N85683	001	
TG UNITED LABS	50MG	N83965	001	
	25MG	N87827	001	Apr 19, 1982
USL PHARMA	50MG	N87752	001	Apr 19, 1982
	25MG	N87638	001	
VANGARD	50MG	N87610	001	
	25MG	N87586	001	May 03, 1982
WARNER CHILCOTT	50MG	N87587	001	May 03, 1982
	25MG	N83458	001	
WATSON LABS	25MG	N85232	002	
	50MG	N83232	001	
	50MG	N83456	001	
	50MG	N85233	001	
	50MG	N86087	001	
	50MG	N86594	001	
	100MG	N81190	001	Jan 24, 1992
	100MG	N85099	001	
	100MG	N87002	001	
	25MG	N84899	001	
WEST WARD	50MG	N84878	001	
	25MG	N83809	002	
WHITEWORTH TOWN PLSN	50MG	N83809	001	
	100MG	N85347	001	
HYDRO-D HALSEY	25MG	N86504	001	
	50MG	N83891	002	
HYDRODIURIL MERCK	25MG	N11835	003	
	50MG	N11835	006	
	100MG	N11835	007	
ORETIC ABBOTT	25MG	N11971	001	
ZIDE SOLVAY	50MG	N83925	001	

HYDROCHLOROTHIAZIDE; IRBESARTAN

TABLET; ORAL

AVALIDE

SANOFI AVENTIS US	12.5MG;75MG	N20758	001	Sep 30, 1997
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HYDROCHLOROTHIAZIDE; LABETALOL HYDROCHLORIDE

TABLET; ORAL

NORMOZIDE

SCHERING	25MG;100MG	N19046	001	Apr 06, 1987
	25MG;200MG	N19046	002	Apr 06, 1987
	25MG;300MG	N19046	003	Apr 06, 1987
	25MG;400MG	N19046	004	Apr 06, 1987
TRANDATE HCT GLAXOSMITHKLINE	25MG;100MG	N19174	001	Apr 10, 1987
	25MG;200MG	N19174	002	Apr 10, 1987
	25MG;300MG	N19174	003	Apr 10, 1987
	25MG;400MG	N19174	004	Apr 10, 1987

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 134 (of 274)

HYDROCHLOROTHIAZIDE; METHYLDOPA

TABLET; ORAL

ALDORIL 15

MERCK

15MG;250MG

N13402 001

ALDORIL 25

MERCK

25MG;250MG

N13402 002

ALDORIL D30

MERCK

30MG;500MG

N13402 003

ALDORIL D50

MERCK

50MG;500MG

N13402 004

METHYLDOPA AND HYDROCHLOROTHIAZIDE

CLONMEL HLTHCARE

15MG;250MG

N72507 001

Jun 02, 1989

25MG;250MG

N72508 001

Jun 02, 1989

30MG;500MG

N72509 001

Jun 02, 1989

50MG;500MG

N72510 001

Jun 02, 1989

IVAX PHARMS

15MG;250MG

N71458 001

Mar 08, 1988

25MG;250MG

N71459 001

Mar 08, 1988

30MG;500MG

N71460 001

Mar 08, 1988

50MG;500MG

N71461 001

Mar 08, 1988

PAR PHARM

15MG;250MG

N70616 001

Feb 02, 1987

25MG;250MG

N70612 001

Feb 02, 1987

30MG;500MG

N70613 001

Feb 02, 1987

50MG;500MG

N70614 001

Feb 02, 1987

PARKE DAVIS

15MG;250MG

N71897 001

Nov 23, 1987

25MG;250MG

N71898 001

Nov 23, 1987

30MG;500MG

N71899 001

Nov 23, 1987

50MG;500MG

N71900 001

Nov 23, 1987

PUREPAC PHARM

15MG;250MG

N70853 001

Oct 08, 1986

25MG;250MG

N70688 001

Apr 24, 1986

30MG;500MG

N70854 001

Oct 08, 1986

50MG;500MG

N70689 001

Apr 24, 1986

SANDOZ

15MG;250MG

N70182 001

Jan 15, 1986

15MG;250MG

N70829 001

Mar 09, 1987

25MG;250MG

N70183 001

Jan 15, 1986

25MG;250MG

N70830 001

Mar 09, 1987

30MG;500MG

N70543 001

Jan 15, 1986

50MG;500MG

N70544 001

Jan 15, 1986

TEVA

15MG;250MG

N71819 001

Apr 08, 1988

25MG;250MG

N71820 001

Apr 08, 1988

30MG;500MG

N71821 001

Apr 08, 1988

50MG;500MG

N71822 001

Apr 08, 1988

WATSON LABS

15MG;250MG

N70365 001

Mar 19, 1986

15MG;250MG

N70958 001

Feb 06, 1989

15MG;250MG

N71920 001

Aug 29, 1988

25MG;250MG

N70366 001

Apr 16, 1986

25MG;250MG

N70959 001

Jan 19, 1989

25MG;250MG

N71921 001

Aug 29, 1988

30MG;500MG

N70367 001

Mar 19, 1986

30MG;500MG

N71069 001

Jan 19, 1989

30MG;500MG

N71922 001

Aug 29, 1988

50MG;500MG

N70368 001

Apr 16, 1986

50MG;500MG

N70960 001

Feb 06, 1989

50MG;500MG

N71923 001

Aug 29, 1988

HYDROCHLOROTHIAZIDE; PINDOLOL

TABLET; ORAL

VISKAZIDE

NOVARTIS

25MG;5MG

N18872 001

Jul 22, 1987

25MG;10MG

N18872 002

Jul 22, 1987

HYDROCHLOROTHIAZIDE; PROPRANOLOL HYDROCHLORIDE

CAPSULE, EXTENDED RELEASE; ORAL

INDERIDE LA 120/50

WYETH AYERST

50MG;120MG

N19059 002

Jul 03, 1985

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 135 (of 274)

HYDROCHLOROTHIAZIDE; PROPRANOLOL HYDROCHLORIDE

CAPSULE, EXTENDED RELEASE; ORAL

INDERIDE LA 160/50

WYETH AYERST 50MG;160MG

N19059 003 Jul 03, 1985

INDERIDE LA 80/50

WYETH AYERST 50MG;80MG

N19059 001 Jul 03, 1985

TABLET; ORAL

INDERIDE-80/25

WYETH PHARMS INC 25MG;80MG

N18031 002

PROPRANOLOL HYDROCHLORIDE & HYDROCHLOROTHIAZIDE

DURAMED PHARMS BARR 25MG;40MG

N71126 001 Mar 02, 1987

25MG;80MG

N71127 001 Mar 02, 1987

PROPRANOLOL HYDROCHLORIDE AND HYDROCHLOROTHIAZIDE

IVAX PHARMS 25MG;40MG

N71552 001 Dec 01, 1988

25MG;80MG

N71553 001 Dec 01, 1988

WARNER CHILCOTT 25MG;40MG

N71771 001 Jan 26, 1988

25MG;80MG

N71772 001 Jan 26, 1988

WATSON LABS 25MG;40MG

N71498 001 Dec 18, 1991

25MG;80MG

N71501 001 Dec 18, 1991

HYDROCHLOROTHIAZIDE; RESERPINE

TABLET; ORAL

H.R.-50

WHITEWORTH TOWN PLSN 50MG;0.125MG

N85338 001

HYDROCHLOROTHIAZIDE W/ RESERPINE

IVAX PHARMS 25MG;0.1MG

N83572 001

25MG;0.125MG

N83571 001

50MG;0.1MG

N83568 001

50MG;0.125MG

N83573 001

PHARMERAL 25MG;0.125MG

N85421 001

50MG;0.125MG

N85420 001

ROXANE 50MG;0.125MG

N84603 001

WATSON LABS 25MG;0.125MG

N84466 001

25MG;0.125MG

N85317 001

25MG;0.125MG

N86330 002

50MG;0.125MG

N83666 001

50MG;0.125MG

N84467 001

50MG;0.125MG

N86331 001

HYDROPRES 25

MERCK 25MG;0.125MG

N11958 002

HYDROPRES 50

MERCK 50MG;0.125MG

N11958 003

HYDRO-RESERP

ABC HOLDING 50MG;0.125MG

N84714 002 Jun 29, 1982

HYDRO-SERP "25"

SANDOZ 25MG;0.125MG

N84827 001

HYDRO-SERP "50"

SANDOZ 50MG;0.125MG

N85213 001

RESERPINE AND HYDROCHLOROTHIAZIDE

BARR 25MG;0.125MG

N84580 001

50MG;0.125MG

N84579 001

SANDOZ 50MG;0.125MG

N88200 001 Jan 31, 1984

RESERPINE AND HYDROCHLOROTHIAZIDE-50

WEST WARD 50MG;0.125MG

N88189 001 May 10, 1984

SERPASIL-ESIDRIX #1

NOVARTIS 25MG;0.1MG

N11878 003

SERPASIL-ESIDRIX #2

NOVARTIS 50MG;0.1MG

N11878 005

HYDROCHLOROTHIAZIDE; SPIRONOLACTONE

TABLET; ORAL

SPIRONOLACTONE AND HYDROCHLOROTHIAZIDE

ASCOT 25MG;25MG

N88025 001 Nov 23, 1984

PUREPAC PHARM 25MG;25MG

N87999 001 Nov 06, 1985

DISCONTINUED DRUG PRODUCT LIST

6 - 136 (of 274)

HYDROCHLOROTHIAZIDE; SPIRONOLACTONE

TABLET; ORAL

SPIRONOLACTONE AND HYDROCHLOROTHIAZIDE

SUPERPHARM 25MG;25MG

N89137 001 Aug 26, 1985

SPIRONOLACTONE W/ HYDROCHLOROTHIAZIDE

IVAX PHARMS 25MG;25MG

N87004 002 May 24, 1982

LEDERLE 25MG;25MG

N87511 001

PARKE DAVIS 25MG;25MG

N87948 001 Feb 22, 1983

PUREPAC PHARM 25MG;25MG

N88054 001 Aug 18, 1983

UPSHER SMITH 25MG;25MG

N87553 001

USL PHARMA 25MG;25MG

N87651 001

VANGARD 25MG;25MG

N87655 001

WATSON LABS 25MG;25MG

N85974 001

25MG;25MG

N86026 001

HYDROCHLOROTHIAZIDE; TIMOLOL MALEATE

TABLET; ORAL

TIMOLIDE 10-25

MERCK 25MG;10MG

N18061 001

HYDROCHLOROTHIAZIDE; TRIAMTERENE

CAPSULE; ORAL

DYAZIDE

GLAXOSMITHKLINE 25MG;50MG

N16042 002

TRIAMTERENE AND HYDROCHLOROTHIAZIDE

NOVARTIS 25MG;37.5MG

N74857 001 Sep 09, 1997

VITARINE 25MG;50MG

N71737 001 Feb 12, 1988

TABLET; ORAL

TRIAMTERENE AND HYDROCHLOROTHIAZIDE

AM THERAP 50MG;75MG

N72022 001 Apr 17, 1988

QUANTUM PHARMICS 50MG;75MG

N71980 001 Apr 17, 1988

HYDROCODONE BITARTRATE; PHENYLPROPANOLAMINE HYDROCHLORIDE

SYRUP; ORAL

CODAMINE

ALPHARMA US PHARMS 5MG/5ML;25MG/5ML

N75103 001 Sep 29, 2000

HYCOMINE

ENDO PHARMS 5MG/5ML;25MG/5ML

N19410 001 Aug 17, 1990

HYCOMINE PEDIATRIC

ENDO PHARMS 2.5MG/5ML;12.5MG/5ML

N19411 001 Aug 17, 1990

HYDROCORTAMATE HYDROCHLORIDE

OINTMENT; TOPICAL

MAGNACORT

PFIZER 0.5%

N10554 001

HYDROCORTISONE

AEROSOL; TOPICAL

AEROSEB-HC

ALLERGAN HERBERT 0.5%

N85805 001

CREAM; TOPICAL

CORT-DOME

BAYER PHARMS 0.5%

N09585 003

1%

N09585 001

DERMACORT

MONARCH PHARMS 1%

N83011 002

ELDECORT

VALEANT PHARM INTL 1%

N80459 001

2.5%

N84055 001

FLEXICORT

WESTWOOD SQUIBB 0.5%

N87136 003 Apr 08, 1982

1%

N87136 002 Apr 08, 1982

2.5%

N87136 001 Apr 08, 1982

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 137 (of 274)

HYDROCORTISONE

CREAM; TOPICAL

HC #1

BAYER PHARMS	0.5%	N80438	001
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HC #4

BAYER PHARMS	1%	N80438	002
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HC (HYDROCORTISONE)

C AND M PHARMA	0.5%	N80482	003
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	1%	N80482	004
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H-CORT

PHARM ASSOC	0.5%	N86823	001
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HYDROCORTISONE

ALPHARMA US PHARMS	2.5%	N89754	001	Feb 01, 1989
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ALTANA	0.5%	N80848	002	
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	1%	N80848	003	
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AMBIX	1%	N86080	001	
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	2.5%	N86271	001	
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EVERYLIFE	0.5%	N80452	001	
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G AND W LABS	1%	N84059	001	
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IVAX PHARMS	1%	N85733	001	
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NASKA	1%	N89706	001	Mar 10, 1988
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PERRIGO NEW YORK	0.5%	N84970	002	
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	1%	N85026	001	
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PHARMADERM	1%	N88845	001	Feb 27, 1986
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	2.5%	N89413	001	Dec 16, 1986
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PHARMAFAIR	1%	N87838	001	Jul 28, 1982
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STIEFEL	1%	N86170	001	
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SYOSSET	0.5%	N85527	001	
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TARO	0.5%	N86154	001	
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TEVA	0.5%	N80400	002	
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	1%	N80400	003	
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	1%	N85191	001	
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	2.5%	N80400	004	
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TOPIDERM	1%	N89273	001	Feb 17, 1989
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USL PHARMA	1%	N88027	001	Sep 27, 1983
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	2.5%	N88029	001	Sep 27, 1983
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WHITEWORTH TOWN PLSN	1%	N80496	002	
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NOGENIC HC

IVAX PHARMS	1%	N87427	001	Apr 04, 1988
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NUTRACORT

HEALTHPOINT	0.5%	N80442	002	
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	1%	N80442	003	
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PENECORT

ALLERGAN HERBERT	1%	N88216	001	Jun 06, 1984
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PROCTOCORT

MONARCH PHARMS	1%	N83011	001	
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SYNACORT

MEDICIS	0.5%	N87459	001	
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GEL; TOPICAL

NUTRACORT

HEALTHPOINT	1%	N84698	001	
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PENECORT

ALLERGAN HERBERT	1%	N88215	001	Jun 06, 1984
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INJECTABLE; INJECTION

CORTEF

PHARMACIA AND UPJOHN	50MG/ML	N09864	001	
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LOTION; TOPICAL

ACTICORT

BAKER NORTON	1%	N86535	001	
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BALNEOL-HC

SOLVAY	1%	N88041	001	Dec 03, 1982
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BETA-HC

BETA DERMAC	1%	N89495	001	Jan 25, 1988
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CORT-DOME

BAYER PHARMS	0.5%	N09895	003	
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27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 138 (of 274)

HYDROCORTISONE

LOTION; TOPICAL

CORT-DOME

BAYER PHARMS

1%

N09895 001

DERMACORT

SOLVAY

0.5%

N84573 002

1%

N86462 001

EPICORT

BLULINE

0.5%

N83219 002

GLYCORT

HERAN

1%

N87489 001

Oct 03, 1983

H-CORT

PHARM ASSOC

0.5%

N86824 001

HYDROCORTISONE

ALPHARMA US PHARMS

0.5%

N87317 001

Jun 07, 1982

1%

N87315 001

Jun 07, 1982

MERICON

0.5%

N85282 001

1%

N85282 002

Feb 26, 1987

NASKA

1%

N89705 001

Apr 25, 1988

PERRIGO NEW YORK

0.5%

N85662 001

1%

N85663 001

TARO

1%

N89024 001

Feb 12, 1986

NUTRACORT

HEALTHPOINT

0.5%

N80443 002

OINTMENT; TOPICAL

CORTRIL

PFIZER GLOBAL

1%

N09176 001

2.5%

N09176 002

HC (HYDROCORTISONE)

C AND M PHARMA

0.5%

N80481 001

1%

N80481 002

HYDROCORTISONE

ALTANA

0.5%

N80489 002

1%

N80489 003

AMBIX

1%

N86079 001

2.5%

N86272 001

NASKA

1%

N89704 001

Mar 10, 1988

PERRIGO NEW YORK

0.5%

N84969 003

1%

N85028 001

PHARMADERM

1%

N88842 001

Feb 09, 1987

TARO

0.5%

N86256 001

USL PHARMA

1%

N88061 001

Sep 27, 1983

2.5%

N88039 001

Sep 27, 1983

HYTONE

DERMIK LABS

1%

N80474 003

2.5%

N80474 004

PENECORT

ALLERGAN HERBERT

2.5%

N88217 001

Jun 06, 1984

POWDER; FOR RX COMPOUNDING

H-CORT

TORCH

100%

N87834 001

Mar 29, 1982

HYDROCORTISONE

PADDOCK

100%

N88082 001

Apr 08, 1983

SOLUTION; TOPICAL

PENECORT

ALLERGAN HERBERT

1%

N88214 001

Jun 06, 1984

TABLET; ORAL

CORTEF

PHARMACIA AND UPJOHN

10MG

N08697 001

CORTRIL

PFIZER

10MG

N09127 005

20MG

N09127 003

HYDROCORTISONE

ANABOLIC

20MG

N83140 001

BARR

20MG

N83999 001

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 139 (of 274)

HYDROCORTISONE

TABLET; ORAL

HYDROCORTISONE

ELKINS SINN

20MG

N80624 001

FERRANTE

10MG

N80568 001

20MG

N80568 002

IMPAX LABS

20MG

N80781 001

INWOOD LABS

20MG

N80732 001

LANNETT

20MG

N85070 001

PANRAY

10MG

N09659 001

20MG

N09659 002

PARKE DAVIS

20MG

N84243 001

PUREPAC PHARM

10MG

N84247 003

Aug 31, 1982

20MG

N80395 001

20MG

N84247 002

ROXANE

10MG

N88539 001

Mar 21, 1984

SANDOZ

20MG

N80642 002

WATSON LABS

20MG

N80355 001

WHITEWORTH TOWN PLSN

10MG

N80344 001

20MG

N80344 002

HYDROCORTONE

MERCK

10MG

N08506 007

20MG

N08506 011

TABLET; VAGINAL

CORTRIL

PFIPHARMECS

10MG

N09796 001

HYDROCORTISONE ACETATE

CREAM; TOPICAL

HEMSOL-HC

ABLE

1%

N81274 001

Jun 19, 1992

HYDROCORTISONE ACETATE

CENCI

1%

N80419 001

Jan 25, 1982

PARKE DAVIS

1%

N89914 001

Jan 03, 1989

PUREPAC PHARM

0.5%

N86050 001

1%

N86052 001

INJECTABLE; INJECTION

CORTEF ACETATE

PHARMACIA AND UPJOHN

50MG/ML

N09378 002

CORTRIL

PFIZER

25MG/ML

N09164 001

HYDROCORTISONE ACETATE

AKORN

25MG/ML

N09637 001

50MG/ML

N09637 002

BEL MAR

25MG/ML

N83739 001

50MG/ML

N83739 002

WATSON LABS

25MG/ML

N83128 001

25MG/ML

N83759 001

50MG/ML

N83759 002

50MG/ML

N85214 001

HYDROCORTONE

MERCK

25MG/ML

N08228 001

50MG/ML

N08228 004

OINTMENT; OPHTHALMIC

HYDROCORTISONE ACETATE

ALTANA

0.5%

N80828 001

OINTMENT; OPHTHALMIC, OTIC

HYDROCORTONE

MERCK

1.5%

N09018 003

OINTMENT; TOPICAL

CORTEF ACETATE

PHARMACIA AND UPJOHN

1%

N08917 002

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 140 (of 274)

HYDROCORTISONE ACETATEOINTMENT; TOPICAL
CORTEF ACETATE

PHARMACIA AND UPJOHN	2.5% **Federal Register determination that product was not discontinued or withdrawn for safety or efficacy reasons**	N08917	001
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HYDROCORTISONE ACETATE; NEOMYCIN SULFATECREAM; TOPICAL
NEO-CORTEF

PHARMACIA AND UPJOHN	1%;EQ 3.5MG BASE/GM	N61049	001
	2.5%;EQ 3.5MG BASE/GM	N61049	002

OINTMENT; OPHTHALMIC
NEO-CORTEF

PHARMACIA AND UPJOHN	0.5%;EQ 3.5MG BASE/GM	N60610	001
	1.5%;EQ 3.5MG BASE/GM	N60610	002

OINTMENT; TOPICAL
NEO-CORTEF

PHARMACIA AND UPJOHN	0.5%;EQ 3.5MG BASE/GM	N60751	001
	1%;EQ 3.5MG BASE/GM	N60751	002
	2.5%;EQ 3.5MG BASE/GM	N60751	003

SUSPENSION/DROPS; OPHTHALMIC
COR-OTICIN

AKORN	1.5%;EQ 3.5MG BASE/ML	N60188	001
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NEO-CORTEF

PHARMACIA AND UPJOHN	0.5%;EQ 3.5MG BASE/ML	N60612	002
	1.5%;EQ 3.5MG BASE/ML	N60612	001

HYDROCORTISONE ACETATE; PRAMOXINE HYDROCHLORIDELOTION; TOPICAL
PRAMOSONE

FERNDAL LABS	0.5%;1%	N83213	002
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HYDROCORTISONE BUTYRATECREAM; TOPICAL
LOCOID

YAMANOUCHI	0.1%	N18795	001	Jan 07, 1983
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OINTMENT; TOPICAL
LOCOID

YAMANOUCHI	0.1%	N19106	001	Jul 03, 1984
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SOLUTION; TOPICAL
LOCOID

YAMANOUCHI	0.1%	N19819	001	Sep 15, 1988
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HYDROCORTISONE CYPIONATESUSPENSION; ORAL
CORTEF

PHARMACIA AND UPJOHN	EQ 10MG BASE/5ML	N09900	001
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HYDROCORTISONE SODIUM PHOSPHATEINJECTABLE; INJECTION
HYDROCORTONE

MERCK	EQ 50MG BASE/ML	N12052	001
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HYDROCORTISONE SODIUM SUCCINATEINJECTABLE; INJECTION
A-HYDROCORT

ABBOTT	EQ 100MG BASE/VIAL	N85928	001	
	EQ 100MG BASE/VIAL	N89577	001	Apr 11, 1989
	EQ 250MG BASE/VIAL	N89578	001	Apr 11, 1989
	EQ 500MG BASE/VIAL	N89579	001	Apr 11, 1989
	EQ 1GM BASE/VIAL	N89580	001	Apr 11, 1989

HYDROCORTISONE SODIUM SUCCINATE

ABRAXIS PHARM	EQ 100MG BASE/VIAL	N88667	001	Jun 08, 1984
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27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 141 (of 274)

HYDROCORTISONE SODIUM SUCCINATE

INJECTABLE; INJECTION

HYDROCORTISONE SODIUM SUCCINATE

ABRAXIS PHARM	EQ 100MG BASE/VIAL	N88712	001	Jun 08, 1984
	EQ 250MG BASE/VIAL	N88668	001	Jun 08, 1984
	EQ 500MG BASE/VIAL	N88669	001	Jun 08, 1984
	EQ 1GM BASE/VIAL	N88670	001	Jun 08, 1984
BAXTER HLTHCARE	EQ 100MG BASE/VIAL	N86619	001	
	EQ 250MG BASE/VIAL	N87567	001	
	EQ 500MG BASE/VIAL	N87568	001	
	EQ 1GM BASE/VIAL	N87569	001	
INTL MEDICATION	EQ 100MG BASE/VIAL	N87532	001	Mar 19, 1982
WATSON LABS	EQ 100MG BASE/VIAL	N84737	002	
	EQ 100MG BASE/VIAL	N84738	001	
	EQ 250MG BASE/VIAL	N84737	001	
	EQ 500MG BASE/VIAL	N84747	001	
	EQ 1GM BASE/VIAL	N84748	001	

HYDROCORTISONE; NEOMYCIN SULFATE

CREAM; TOPICAL

NEO-CORT-DOME

BAYER PHARMS	0.5%;EQ 3.5MG BASE/GM	N50237	006	Jun 05, 1984
	1%;EQ 3.5MG BASE/GM	N50237	005	Jun 05, 1984

HYDROCORTISONE; NEOMYCIN SULFATE; POLYMYXIN B SULFATE

SOLUTION/DROPS; OTIC

NEOMYCIN SULFATE-POLYMYXIN B SULFATE-HYDROCORTISONE

PHARMAFAIR	1%;EQ 3.5MG BASE/ML;10,000 UNITS/ML	N62394	001	Sep 29, 1982
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OTOCORT

WATSON LABS	1%;EQ 3.5MG BASE/ML;10,000 UNITS/ML	N60730	002	
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SUSPENSION/DROPS; OPHTHALMIC

NEOMYCIN SULFATE-POLYMYXIN B SULFATE-HYDROCORTISONE

PHARMAFAIR	1%;EQ 3.5MG BASE/ML;10,000 UNITS/ML	N62623	001	Sep 24, 1985
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SUSPENSION/DROPS; OTIC

NEOMYCIN SULFATE, POLYMYXIN B SULFATE & HYDROCORTISONE

PHARMAFAIR	1%;EQ 3.5MG BASE/ML;10,000 UNITS/ML	N62617	001	Sep 18, 1985
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OTICAIR

PHARMAFAIR	1%;EQ 3.5MG BASE/ML;10,000 UNITS/ML	N62399	001	Nov 18, 1982
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OTOBIONE

SCHERING	1%;EQ 3.5MG BASE/ML;10,000 UNITS/ML	N61816	001	
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OTOCORT

WATSON LABS	1%;EQ 3.5MG BASE/ML;10,000 UNITS/ML	N62521	001	Jul 11, 1985
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HYDROCORTISONE; POLYMYXIN B SULFATE

SOLUTION/DROPS; OTIC

OTOBiotic

SCHERING	5MG/ML;EQ 10,000 UNITS BASE/ML	N62302	001	
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PYOCIDIN

FOREST LABS	5MG/ML;EQ 10,000 UNITS BASE/ML	N61606	001	
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HYDROCORTISONE; TETRACYCLINE HYDROCHLORIDE

OINTMENT; OPHTHALMIC

ACHROMYCIN

LEDERLE	1.5%;1%	N50272	001	
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HYDROCORTISONE; UREA

CREAM; TOPICAL

ALPHADERM

BIOGLAN	1%;10%	N86008	001	
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CALMURID HC

PHARMACIA AND UPJOHN	1%;10%	N83947	001	
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27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 142 (of 274)

HYDROFLUMETHIAZIDE

TABLET; ORAL

DIUCARDIN

WYETH AYERST

50MG

N83383 001

HYDROFLUMETHIAZIDE

WATSON LABS

50MG

N88031 001

Apr 06, 1983

50MG

N88528 001

Aug 15, 1984

HYDROFLUMETHIAZIDE; RESERPINE

TABLET; ORAL

HYDROFLUMETHIAZIDE AND RESERPINE

USL PHARMA

50MG;0.125MG

N88195 001

Oct 26, 1983

WATSON LABS

25MG;0.125MG

N88127 001

Mar 22, 1983

50MG;0.125MG

N88110 001

Mar 22, 1983

RESERPINE AND HYDROFLUMETHIAZIDE

IVAX PHARMS

50MG;0.125MG

N88932 001

Jan 11, 1985

PAR PHARM

50MG;0.125MG

N88907 001

Sep 20, 1985

SALUTENSIN

SHIRE

50MG;0.125MG

N12359 003

SALUTENSIN-DEMI

SHIRE

25MG;0.125MG

N12359 004

HYDROMORPHONE HYDROCHLORIDE

CAPSULE, EXTENDED RELEASE; ORAL

PALLADONE

PURDUE PHARMA LP

12MG

N21044 001

Sep 24, 2004

16MG

N21044 002

Sep 24, 2004

24MG

N21044 003

Sep 24, 2004

32MG

N21044 004

Sep 24, 2004

HYDROXOCOBALAMIN

INJECTABLE; INJECTION

ALPHAREDISOL

MERCK

1MG/ML

N80778 001

HYDROXOCOBALAMIN

ABRAXIS PHARM

1MG/ML

N84921 001

WATSON LABS

1MG/ML

N85528 001

HYDROXOMIN

BEL MAR

1MG/ML

N84629 001

HYDROXYAMPHETAMINE HYDROBROMIDE

SOLUTION/DROPS; OPHTHALMIC

PAREDRIINE

AKORN

1%

N00004 004

HYDROXYPROGESTERONE CAPROATE

INJECTABLE; INJECTION

DELALUTIN

BRISTOL MYERS SQUIBB

125MG/ML

N10347 004

125MG/ML

N16911 001

250MG/ML

N10347 002

250MG/ML

N16911 002

HYDROXYPROGESTERONE CAPROATE

AKORN

125MG/ML

N18004 001

WATSON LABS

125MG/ML

N17439 001

250MG/ML

N17439 002

HYDROXYSTILBAMIDINE ISETHIONATE

INJECTABLE; INJECTION

HYDROXYSTILBAMIDINE ISETHIONATE

SANOFI AVENTIS US

225MG/AMP

N09166 001

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 143 (of 274)

HYDROXYZINE HYDROCHLORIDE

INJECTABLE; INJECTION

HYDROXYZINE

BAXTER HLTHCARE 50MG/ML

N85551 002

HYDROXYZINE HYDROCHLORIDE

ABRAXIS PHARM 25MG/ML

N88184 001 Mar 31, 1983

50MG/ML

N88185 001 Mar 31, 1983

ALTANA 25MG/ML

N87273 001 Apr 20, 1982

50MG/ML

N87273 002 Apr 20, 1982

BAXTER HLTHCARE 25MG/ML

N85551 001

HOSPIRA 50MG/ML

N86821 001

PHARMAFAIR 25MG/ML

N88862 001 Feb 14, 1986

25MG/ML

N89106 001 Feb 14, 1986

50MG/ML

N88881 001 Feb 14, 1986

50MG/ML

N89107 001 Feb 14, 1986

SMITH AND NEPHEW 25MG/ML

N87592 001

SOLOPAK 25MG/ML

N86822 001

25MG/ML

N87591 001

50MG/ML

N87310 001

50MG/ML

N87593 001

50MG/ML

N87595 001

50MG/ML

N87596 001

WATSON LABS 25MG/ML

N85778 001

25MG/ML

N87274 001

50MG/ML

N87274 002

WYETH AYERST 25MG/ML

N86258 001

50MG/ML

N86258 002

ORGATRAX

ORGANON USA INC 25MG/ML

N87014 001

50MG/ML

N87014 002

SYRUP; ORAL

ATARAX

ROERIG 10MG/5ML

N10485 001

HYDROXYZINE HYDROCHLORIDE

ALPHARMA US PHARMS 10MG/5ML

N88785 001 Feb 03, 1988

KV PHARM 10MG/5ML

N87730 001 Jul 01, 1982

TABLET; ORAL

ATARAX

PFIZER 10MG

N10392 001

25MG

N10392 004

50MG

N10392 006

100MG

N10392 005

HYDROXYZINE HYDROCHLORIDE

ABLE 10MG

N40559 001 Jul 22, 2004

25MG

N40562 001 Jul 22, 2004

50MG

N40563 001 Jul 22, 2004

ACTAVIS TOTOWA 10MG

N89071 001 Jul 22, 1986

25MG

N89072 001 Jul 22, 1986

50MG

N89073 001 Jul 22, 1986

HALSEY 10MG

N89366 001 May 02, 1988

25MG

N89117 001 May 02, 1988

50MG

N89396 001 May 02, 1988

IVAX PHARMS 10MG

N87216 001

25MG

N87410 001

50MG

N87411 001

KV PHARM 10MG

N87819 001 Jun 23, 1982

25MG

N87820 001 Jun 23, 1982

50MG

N87821 001 Jun 23, 1982

100MG

N87822 001 Jun 23, 1982

MUTUAL PHARM 10MG

N88409 001 Nov 15, 1983

25MG

N87857 001 Apr 18, 1983

50MG

N87860 001 Apr 18, 1983

100MG

N87862 001 Apr 18, 1983

PAR PHARM 10MG

N87602 001 Jan 22, 1982

DISCONTINUED DRUG PRODUCT LIST

6 - 144 (of 274)

HYDROXYZINE HYDROCHLORIDE

TABLET; ORAL

HYDROXYZINE HYDROCHLORIDE

PAR PHARM	25MG	N87603	001	Jan 22, 1982
	50MG	N87604	001	Jan 22, 1982
PLIVA	100MG	N81054	001	Sep 25, 1995
PUREPAC PHARM	10MG	N88120	001	Sep 25, 1984
	25MG	N88121	001	Sep 25, 1984
	50MG	N88122	001	Sep 25, 1984
QUANTUM PHARMICS	10MG	N88540	001	Oct 22, 1985
	25MG	N88551	001	Oct 22, 1985
	50MG	N88529	001	Oct 22, 1985
SANDOZ	10MG	N87246	002	
	25MG	N85247	001	
	50MG	N87245	001	
SUPERPHARM	10MG	N88794	001	Dec 05, 1984
	25MG	N88795	001	Dec 05, 1984
	50MG	N88796	001	Dec 05, 1984
USL PHARMA	10MG	N89121	001	Mar 20, 1986
	25MG	N89122	001	Mar 20, 1986
	50MG	N89123	001	Mar 20, 1986
WATSON LABS	10MG	N86827	001	
	25MG	N86829	001	
	50MG	N86836	001	

HYDROXYZINE PAMOATE

CAPSULE; ORAL

HYDROXYZINE PAMOATE

DURAMED PHARMS BARR	EQ 25MG HCL	N88593	001	Feb 29, 1984
	EQ 50MG HCL	N88594	001	Feb 29, 1984
	EQ 100MG HCL	N88595	001	Feb 29, 1984
PAR PHARM	EQ 25MG HCL	N87656	001	Jun 11, 1982
	EQ 25MG HCL	N89145	001	Mar 17, 1986
	EQ 50MG HCL	N87657	001	Jun 11, 1982
	EQ 50MG HCL	N89146	001	Mar 17, 1986
	EQ 100MG HCL	N87658	001	Jun 11, 1982
SANDOZ	EQ 50MG HCL	N81128	001	Jun 28, 1991
	EQ 100MG HCL	N81129	001	Jun 28, 1991
SUPERPHARM	EQ 25MG HCL	N89031	001	Jan 02, 1987
	EQ 50MG HCL	N89032	001	Jan 02, 1987
	EQ 100MG HCL	N89033	001	Jan 02, 1987
VANGARD	EQ 25MG HCL	N88392	001	Sep 19, 1983
	EQ 50MG HCL	N88393	001	Sep 19, 1983
WATSON LABS	EQ 25MG HCL	N86698	001	
	EQ 25MG HCL	N86840	001	Jul 01, 1982
	EQ 50MG HCL	N86695	001	
	EQ 50MG HCL	N86705	001	Jul 01, 1982
	EQ 50MG HCL	N87767	001	Aug 16, 1982
	EQ 100MG HCL	N86697	001	
	EQ 100MG HCL	N86728	001	Oct 05, 1982
	EQ 100MG HCL	N87790	001	Aug 16, 1982
HY-PAM "25"				
TEVA	EQ 25MG HCL	N88713	001	Mar 04, 1985
VISTARIL				
PFIZER	EQ 100MG HCL	N11459	006	

IBUPROFEN

CAPSULE; ORAL

MIDOL

BAYER	200MG **Federal Register determination that product was not discontinued or withdrawn for safety or efficacy reasons**	N70626	001	Sep 02, 1987
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27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 145 (of 274)

IBUPROFENCAPSULE; ORAL
MIDOL

BAYER

200MG **Federal Register determination
that product was not discontinued or
withdrawn for safety or efficacy
reasons**

N71002 001

Sep 02, 1987

SUSPENSION; ORAL

IBU

ABBOTT

100MG/5ML

N19784 001

Dec 18, 1989

SUSPENSION/DROPS; ORAL

MOTRIN

MCNEIL

40MG/ML

N20476 001

May 25, 1995

TABLET; ORAL

ACHES-N-PAIN

LEDERLE

200MG

N71065 001

May 28, 1987

IBU

BASF

400MG

N18197 001

400MG

N70083 001

Feb 22, 1985

600MG

N70088 001

Feb 08, 1985

600MG

N70099 001

Mar 29, 1985

800MG

N70745 001

Jul 23, 1986

IBUPRIN

PLIVA

200MG

N71773 001

Jul 16, 1987

IBUPROFEN

ABBOTT

600MG

N70556 001

Jun 14, 1985

800MG

N71264 001

Jul 25, 1986

HALSEY

200MG

N71027 001

Sep 29, 1987

300MG

N71028 001

Mar 23, 1987

400MG

N71029 001

Mar 23, 1987

600MG

N71030 001

Mar 23, 1987

800MG

N72137 001

Feb 05, 1988

IVAX PHARMS

200MG

N71154 001

Oct 27, 1987

200MG

N72040 001

Apr 29, 1988

200MG

N72901 001

Dec 19, 1991

200MG

N72903 001

Dec 19, 1991

400MG

N71145 001

Sep 23, 1986

600MG

N71146 001

Sep 23, 1986

800MG

N71769 001

May 08, 1987

LEDERLE

400MG

N70629 001

Sep 19, 1986

600MG

N70630 001

Sep 19, 1986

MCNEIL

400MG

N70081 001

Jun 16, 1986

600MG

N70476 001

Jun 16, 1986

MUTUAL PHARM

200MG

N70493 001

Dec 24, 1985

200MG

N70908 001

Sep 26, 1986

200MG

N71462 001

Oct 02, 1986

400MG

N70079 001

Jul 24, 1985

600MG

N70080 001

Jul 24, 1985

800MG

N71448 001

Feb 18, 1987

PAR PHARM

300MG

N70328 001

Aug 06, 1985

PUREPAC PHARM

200MG

N71122 001

Oct 03, 1986

200MG

N71664 001

Feb 03, 1987

300MG

N71123 001

Sep 19, 1986

400MG

N71124 001

Sep 19, 1986

600MG

N71125 001

Sep 19, 1986

800MG

N71964 001

Feb 01, 1988

SANDOZ

400MG

N72064 001

Jan 14, 1988

600MG

N72065 001

Jan 14, 1988

800MG

N71938 001

Jan 14, 1988

SUPERPHARM

600MG

N70709 001

Apr 25, 1986

TEVA

200MG

N73141 001

May 29, 1992

400MG

N73343 001

Jun 30, 1992

600MG

N73344 001

Jun 30, 1992

800MG

N73345 001

Jun 30, 1992

WATSON LABS

200MG

N71765 001

Sep 04, 1987

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 146 (of 274)

IBUPROFEN

TABLET; ORAL

IBUPROFEN

WATSON LABS

200MG

N71905 001

Mar 08, 1988

300MG

N71338 001

Dec 01, 1986

400MG

N70038 001

Sep 06, 1985

600MG

N70041 001

Sep 06, 1985

800MG

N71911 001

Oct 13, 1987

IBU-TAB

ALRA

800MG

N71965 001

Aug 11, 1988

MIDOL

BAYER

200MG

N70591 001

Sep 02, 1987

200MG

N71001 001

Sep 02, 1987

MOTRIN

MCNEIL PED

100MG

N20418 001

Nov 16, 1994

NUPRIN

BRISTOL MYERS

200MG

N72035 001

Feb 16, 1988

200MG

N72036 001

Feb 16, 1988

MCNEIL

200MG

N19012 001

May 18, 1984

200MG

N19012 002

Jul 29, 1987

RUFEN

BASF

600MG

N18197 002

Mar 05, 1984

TABLET, CHEWABLE; ORAL

MOTRIN

MCNEIL PED

50MG

N20135 001

Nov 16, 1994

100MG

N20135 002

Nov 16, 1994

IDARUBICIN HYDROCHLORIDE

INJECTABLE; INJECTION

IDAMYCIN

PHARMACIA AND UPJOHN

5MG/VIAL

N50661 002

Sep 27, 1990

10MG/VIAL

N50661 001

Sep 27, 1990

20MG/VIAL

N50661 003

Apr 25, 1995

IDOXURIDINE

OINTMENT; OPHTHALMIC

STOXIL

GLAXOSMITHKLINE

0.5%

N15868 001

SOLUTION/DROPS; OPHTHALMIC

STOXIL

GLAXOSMITHKLINE

0.1%

N13934 001

IFOSFAMIDE

INJECTABLE; INJECTION

IFEX

BRISTOL MYERS SQUIBB

1GM/VIAL

N19763 001

Dec 30, 1988

3GM/VIAL

N19763 002

Dec 30, 1988

IMATINIB MESYLATE

CAPSULE; ORAL

GLEEVEC

NOVARTIS

EQ 50MG BASE

N21335 001

May 10, 2001

EQ 100MG BASE

N21335 002

May 10, 2001

IMIPRAMINE HYDROCHLORIDE

CONCENTRATE; ORAL

IMIPRAMINE HYDROCHLORIDE

NOVARTIS

25MG/ML

N86765 001

INJECTABLE; INJECTION

TOFRANIL

NOVARTIS

12.5MG/ML

N11838 002

TABLET; ORAL

IMIPRAMINE HYDROCHLORIDE

LEDERLE

10MG

N86269 001

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 147 (of 274)

IMIPRAMINE HYDROCHLORIDE

TABLET; ORAL

IMIPRAMINE HYDROCHLORIDE

LEDERLE	25MG	N86267	001	
	50MG	N86268	001	
ROXANE	10MG	N83799	001	
	25MG	N83799	002	
	50MG	N83799	003	
SANDOZ	10MG	N85200	001	
	25MG	N84869	002	
	50MG	N85133	001	
TEVA	10MG	N83729	001	
	25MG	N83729	004	
	50MG	N83729	003	
USL PHARMA	25MG	N87776	001	Feb 10, 1982
VANGARD	10MG	N88036	001	Nov 03, 1982
	25MG	N87619	001	Feb 09, 1982
	50MG	N87631	001	Jan 04, 1982
WATSON LABS	10MG	N85220	001	
	10MG	N85875	001	
	25MG	N84252	002	
	25MG	N85878	001	
	50MG	N85221	001	
	50MG	N85877	001	
WEST WARD	25MG	N88222	001	May 26, 1983
	50MG	N88223	001	May 26, 1983
JANIMINE				
ABBOTT	10MG	N17895	001	
	25MG	N17895	002	
	50MG	N17895	003	
PRAMINE				
ALRA	10MG	N83827	001	
	25MG	N83827	002	
	50MG	N83827	003	
PRESAMINE				
SANOFI AVENTIS US	10MG	N11836	006	
	25MG	N11836	003	
	50MG	N11836	007	

INAMRINONE LACTATE

INJECTABLE; INJECTION

INOCOR

SANOFI AVENTIS US	EQ 5MG BASE/ML	N18700	001	Jul 31, 1984
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INDAPAMIDE

TABLET; ORAL

INDAPAMIDE

TEVA	1.25MG	N74665	001	Apr 04, 1997
	2.5MG	N74665	002	Apr 04, 1997

INDECAINIDE HYDROCHLORIDE

TABLET, EXTENDED RELEASE; ORAL

DECABID

LILLY	EQ 50MG BASE	N19693	001	Dec 29, 1989
	EQ 75MG BASE	N19693	002	Dec 29, 1989
	EQ 100MG BASE	N19693	003	Dec 29, 1989

INDOCYANINE GREEN

INJECTABLE; INJECTION

IC-GREEN

AKORN	10MG/VIAL	N11525	003	
	40MG/VIAL	N11525	004	
	50MG/VIAL	N11525	002	

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 148 (of 274)

INDOMETHACIN

CAPSULE; ORAL

INDOCIN

MERCK

25MG

N16059 001

50MG

N16059 002

INDO-LEMMON

TEVA

25MG

N70266 001

Nov 07, 1985

50MG

N70267 001

Nov 07, 1985

INDOMETHACIN

ABLE

25MG

N76666 001

Dec 17, 2003

50MG

N76666 002

Dec 17, 2003

DURAMED PHARMS BARR

25MG

N70326 001

Oct 18, 1985

50MG

N70327 001

Oct 18, 1985

HALSEY

25MG

N70782 001

Jun 03, 1987

50MG

N70635 001

Jun 03, 1987

IVAX PHARMS

25MG

N18730 001

May 04, 1984

25MG

N70719 001

Feb 12, 1986

50MG

N18730 002

May 04, 1984

50MG

N70756 001

Feb 12, 1986

MUTUAL PHARM

25MG

N70067 001

Oct 03, 1986

25MG

N70899 001

Feb 09, 1987

50MG

N70068 001

Oct 03, 1986

50MG

N70900 001

Feb 09, 1987

PAR PHARM

25MG

N18829 002

Aug 06, 1984

50MG

N18829 001

Aug 06, 1984

50MG

N70651 001

Mar 05, 1986

PARKE DAVIS

25MG

N18806 001

Nov 23, 1984

50MG

N18806 002

Nov 23, 1984

PIONEER PHARMS

25MG

N70813 001

Aug 11, 1986

50MG

N70592 001

Aug 11, 1986

PLIVA

25MG

N71148 001

Mar 18, 1987

50MG

N71149 001

Mar 18, 1987

RADIUS PHARMS

25MG

N18851 001

May 18, 1984

50MG

N18851 002

May 18, 1984

ROXANE

25MG

N70353 001

Jun 18, 1985

50MG

N70354 001

Jun 18, 1985

SANDOZ

25MG

N70673 001

Apr 29, 1987

50MG

N70674 001

Apr 29, 1987

SUPERPHARM

25MG

N70487 001

Oct 10, 1986

50MG

N70488 001

Oct 10, 1986

TEVA

25MG

N71342 001

Apr 18, 1988

50MG

N71343 001

Apr 18, 1988

WATSON LABS

25MG

N18690 001

Jul 31, 1984

25MG

N70529 001

Oct 18, 1985

25MG

N70784 001

Aug 20, 1986

25MG

N72996 001

Jul 31, 1991

50MG

N18690 002

Jul 31, 1984

50MG

N70530 001

Oct 18, 1985

50MG

N70785 001

Aug 20, 1986

50MG

N71635 001

May 18, 1987

50MG

N72997 001

Jul 31, 1991

CAPSULE, EXTENDED RELEASE; ORAL

INDOCIN SR

MERCK

75MG

N18185 001

Feb 23, 1982

INDOMETHACIN

ABLE

75MG

N76114 001

Feb 06, 2002

INWOOD LABS

75MG

N72410 001

Mar 15, 1989

SUPPOSITORY; RECTAL

INDOCIN

MERCK

50MG

N17814 001

Aug 13, 1984

SUSPENSION; ORAL

INDOMETHACIN

ROXANE

25MG/5ML

N71412 001

Mar 18, 1987

DISCONTINUED DRUG PRODUCT LIST

6 - 149 (of 274)

INSULIN PORK

INJECTABLE; INJECTION			
Iletin I			
Lilly	500 UNITS/ML	N17931	001
INSULIN			
NOVO NORDISK INC	40 UNITS/ML	N17926	001
REGULAR INSULIN			
NOVO NORDISK INC	100 UNITS/ML	N17926	003

INSULIN PURIFIED BEEF

INJECTABLE; INJECTION			
REGULAR Iletin II			
Lilly	100 UNITS/ML	N18478	001

INSULIN PURIFIED PORK

INJECTABLE; INJECTION			
Iletin II			
Lilly	500 UNITS/ML	N18344	002
REGULAR Iletin II (PORK)			
Lilly	100 UNITS/ML	N18344	001
REGULAR PURIFIED PORK INSULIN			
NOVO NORDISK INC	100 UNITS/ML	N18381	001
VELOSULIN			
NOVO NORDISK INC	100 UNITS/ML	N18193	001

INSULIN PURIFIED PORK; INSULIN SUSP ISOPHANE PURIFIED PORK

INJECTABLE; INJECTION			
INSULIN NORDISK MIXTARD (PORK)			
NOVO NORDISK INC	30 UNITS/ML;70 UNITS/ML	N18195	001

INSULIN RECOMBINANT HUMAN

INJECTABLE; INJECTION			
HUMULIN BR			
Lilly	100 UNITS/ML	N19529	001 Apr 28, 1986
VELOSULIN BR			
NOVO NORDISK INC	100 UNITS/ML	N21028	001 Jul 19, 1999

INSULIN RECOMBINANT PURIFIED HUMAN

INJECTABLE; INJECTION			
NOVOLIN R			
NOVO NORDISK INC	100 UNITS/ML	N18778	001 Aug 30, 1983
VELOSULIN BR HUMAN			
NOVO NORDISK INC	100 UNITS/ML	N19450	001 May 30, 1986

INSULIN RECOMBINANT PURIFIED HUMAN; INSULIN SUSP ISOPHANE SEMISYNTHETIC PURIFIED HUMAN

INJECTABLE; INJECTION			
MIXTARD HUMAN 70/30			
BAYER PHARMS	30 UNITS/ML;70 UNITS/ML	N19585	001 Mar 11, 1988
NOVOLIN 70/30			
NOVO NORDISK INC	30 UNITS/ML;70 UNITS/ML	N19441	001 Jul 11, 1986

INSULIN SUSP ISOPHANE BEEF

INJECTABLE; INJECTION			
NPH INSULIN			
NOVO NORDISK INC	40 UNITS/ML	N17929	001
	100 UNITS/ML	N17929	003

INSULIN SUSP ISOPHANE BEEF/PORK

INJECTABLE; INJECTION			
NPH Iletin I (BEEF-PORK)			
Lilly	40 UNITS/ML	N17936	001
	100 UNITS/ML	N17936	002

DISCONTINUED DRUG PRODUCT LIST

6 - 150 (of 274)

INSULIN SUSP ISOPHANE PURIFIED BEEF

INJECTABLE; INJECTION		
NPH ILETIN II		
LILLY	100 UNITS/ML	N18479 001

INSULIN SUSP ISOPHANE PURIFIED PORK

INJECTABLE; INJECTION		
INSULIN INSULATARD NPH NORDISK		
NOVO NORDISK INC	100 UNITS/ML	N18194 001
NPH ILETIN II (PORK)		
LILLY	100 UNITS/ML	N18345 001
NPH PURIFIED PORK ISOPHANE INSULIN		
NOVO NORDISK INC	100 UNITS/ML	N18623 001

INSULIN SUSP ISOPHANE SEMISYNTHETIC PURIFIED HUMAN

INJECTABLE; INJECTION			
INSULATARD NPH HUMAN			
NOVO NORDISK INC	100 UNITS/ML	N19449 001	May 30, 1986
NOVOLIN N			
NOVO NORDISK INC	100 UNITS/ML	N19065 001	Jan 23, 1985

INSULIN SUSP PROTAMINE ZINC BEEF/PORK

INJECTABLE; INJECTION		
PROTAMINE ZINC & ILETIN I (BEEF-PORK)		
LILLY	40 UNITS/ML	N17932 001
	100 UNITS/ML	N17932 002

INSULIN SUSP PROTAMINE ZINC PURIFIED BEEF

INJECTABLE; INJECTION		
PROTAMINE ZINC AND ILETIN II		
LILLY	100 UNITS/ML	N18476 001
PROTAMINE ZINC INSULIN		
BRISTOL MYERS SQUIBB	40 UNITS/ML	N17928 001
	100 UNITS/ML	N17928 003

INSULIN SUSP PROTAMINE ZINC PURIFIED PORK

INJECTABLE; INJECTION		
PROTAMINE ZINC AND ILETIN II (PORK)		
LILLY	100 UNITS/ML	N18346 001

INSULIN ZINC SUSP BEEF

INJECTABLE; INJECTION		
LENTE INSULIN		
NOVO NORDISK INC	40 UNITS/ML	N17998 001
	100 UNITS/ML	N17998 003

INSULIN ZINC SUSP EXTENDED BEEF

INJECTABLE; INJECTION		
ULTRALENTE INSULIN		
NOVO NORDISK INC	100 UNITS/ML	N17997 003

INSULIN ZINC SUSP EXTENDED PURIFIED BEEF

INJECTABLE; INJECTION		
ULTRALENTE		
NOVO NORDISK INC	100 UNITS/ML	N18385 001

INSULIN ZINC SUSP EXTENDED RECOMBINANT HUMAN

INJECTABLE; INJECTION			
HUMULIN U			
LILLY	40 UNITS/ML	N19571 001	Jun 10, 1987
	100 UNITS/ML	N19571 002	Jun 10, 1987

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 151 (of 274)

INSULIN ZINC SUSP PROMPT BEEF

INJECTABLE; INJECTION			
SEMILENTE INSULIN			
NOVO NORDISK INC	100 UNITS/ML	N17996	003

INSULIN ZINC SUSP PROMPT PURIFIED PORK

INJECTABLE; INJECTION			
SEMILENTE			
NOVO NORDISK INC	100 UNITS/ML	N18382	001

INSULIN ZINC SUSP PURIFIED BEEF

INJECTABLE; INJECTION			
LENTE ILETIN II			
LILLY	100 UNITS/ML	N18477	001

INSULIN ZINC SUSP PURIFIED BEEF/PORK

INJECTABLE; INJECTION			
LENTARD			
NOVO NORDISK INC	100 UNITS/ML	N18384	001

INSULIN ZINC SUSP PURIFIED PORK

INJECTABLE; INJECTION			
LENTE			
NOVO NORDISK INC	100 UNITS/ML	N18383	001
LENTE ILETIN II (PORK)			
LILLY	100 UNITS/ML	N18347	001

INSULIN ZINC SUSP RECOMBINANT HUMAN

INJECTABLE; INJECTION			
NOVOLIN L			
NOVO NORDISK INC	100 UNITS/ML	N19965	001 Jun 25, 1991

INSULIN ZINC SUSP SEMISYNTHETIC PURIFIED HUMAN

INJECTABLE; INJECTION			
NOVOLIN L			
NOVO NORDISK INC	100 UNITS/ML	N18777	001 Aug 30, 1983

INULIN

INJECTABLE; INJECTION			
INULIN AND SODIUM CHLORIDE			
ISO TEX	100MG/ML	N02282	001

INVERT SUGAR

INJECTABLE; INJECTION			
TRAVERS 10% IN PLASTIC CONTAINER			
BAXTER HLTHCARE	10GM/100ML	N16717	001

IOCEAMIC ACID

TABLET; ORAL			
CHOLEBRINE			
MALLINCKRODT	750MG	N17129	001

IODAMIDE MEGLUMINE

INJECTABLE; INJECTION			
RENOVUE-65			
BRACCO	65%	N17902	001
RENOVUE-DIP			
BRACCO	24%	N17903	001

IODIPAMIDE MEGLUMINE

INJECTABLE; INJECTION			
CHOLOGRAFIN MEGLUMINE			
BRACCO	10.3%	N09321	007

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 152 (of 274)

IODIPAMIDE SODIUM

INJECTABLE; INJECTION
CHOLOGRAFIN SODIUM
BRACCO

20%

N09321 001

IODOHIPPURATE SODIUM, I-123

INJECTABLE; INJECTION
NEPHROFLOW
GE HEALTHCARE

1mCi/ML

N18289 001 Dec 28, 1984

IODOHIPPURATE SODIUM, I-131

INJECTABLE; INJECTION
HIPURAN I 131
MALLINCKRODT
HIPPUTOPE

0.25mCi/ML

N16666 001

BRACCO

1-2mCi/VIAL

N15419 002

IODOHIPPURATE SODIUM I 131

CIS

0.2mCi/ML

N17313 001

IODOXAMATE MEGLUMINE

INJECTABLE; INJECTION
CHOLOVUE
BRACCO

9.9%

N18077 001

40.3%

N18076 001

IOFETAMINE HYDROCHLORIDE I-123

INJECTABLE; INJECTION
SPECTAMINE
IMP

1mCi/ML

N19432 001 Dec 24, 1987

IOHEXOL

INJECTABLE; INJECTION
OMNIPAQUE 210
GE HEALTHCARE

45.3%

N18956 006 Jun 30, 1989

SOLUTION; URETHRAL
OMNIPAQUE 70

GE HEALTHCARE

15.1%

N18956 007 Jun 01, 1994

IOPAMIDOL

INJECTABLE; INJECTION
IOPAMIDOL

BAXTER HLTHCARE

41%

N74629 001 Nov 06, 1996

51%

N74629 004 Mar 31, 1998

61%

N74629 002 Nov 06, 1996

76%

N74629 003 Nov 06, 1996

MAYNE PHARMA USA

61%

N74734 001 Dec 10, 1996

76%

N74734 002 Dec 10, 1996

IOPAMIDOL-300

ABBOTT

61%

N74638 001 Apr 30, 1997

ISOVUE-128

BRACCO

26%

N18735 005 Oct 21, 1986

ISOVUE-200

BRACCO

41%

N20327 001 Oct 12, 1994

IOPANOIC ACID

TABLET; ORAL
TELEPAQUE

GE HEALTHCARE

500MG

N08032 001

IOPHENDYLATE

INJECTABLE; INJECTION
PANTOPAQUE
ALCON

100%

N05319 001

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 153 (of 274)

IOTHALAMATE MEGLUMINE; IOTHALAMATE SODIUMINJECTABLE; INJECTION
VASCORAY

MALLINCKRODT 52%;26% N16783 001

IOTHALAMATE SODIUMINJECTABLE; INJECTION
ANGIO-CONRAY

MALLINCKRODT 80% N13319 001

CONRAY 325

MALLINCKRODT 54.3% N17685 001

IOTROLANINJECTABLE; INTRATHECAL
OSMOVIST 190

BERLEX 40.6% N19580 001 Dec 07, 1989

OSMOVIST 240

BERLEX 51.3% N19580 002 Dec 07, 1989

IOVERSOLINJECTABLE; INJECTION
OPTIRAY 240

MALLINCKRODT 51% N20923 001 May 28, 1998

OPTIRAY 320

MALLINCKRODT 68% N20923 002 May 29, 1998

IPODATE CALCIUMGRANULE; ORAL
ORAGRAFIN CALCIUM

BRACCO 3GM/PACKET N12968 001

IPODATE SODIUMCAPSULE; ORAL
BILIVIST

BERLEX 500MG N87768 001 Aug 11, 1982

ORAGRAFIN SODIUM

BRACCO 500MG N12967 001

IPRATROPIUM BROMIDEAEROSOL, METERED; INHALATION
ATROVENT

BOEHRINGER INGELHEIM 0.018MG/INH N19085 001 Dec 29, 1986

SOLUTION; INHALATION
ATROVENT

BOEHRINGER INGELHEIM 0.02% N20228 001 Sep 29, 1993

IPRATROPIUM BROMIDE

ROXANE 0.02% N75867 001 Jul 22, 2002

IRON DEXTRANINJECTABLE; INJECTION
IRON DEXTRAN

SANOFI AVENTIS US EQ 50MG IRON/ML N10787 002

ISOETHARINE HYDROCHLORIDESOLUTION; INHALATION
BRONKOSOL

SANOFI AVENTIS US 0.25% N12339 009

1% N12339 008

ISOETHARINE HYDROCHLORIDE

ALPHARMA US PHARMS 1% N87101 001

ASTRAZENECA 0.062% N87937 001 Nov 15, 1982

0.062% N89614 001 Jun 13, 1991

0.125% N87938 001 Nov 15, 1982

DISCONTINUED DRUG PRODUCT LIST

6 - 154 (of 274)

ISOETHARINE HYDROCHLORIDE

SOLUTION; INHALATION

ISOETHARINE HYDROCHLORIDE

ASTRAZENECA	0.125%	N89615	001	Jun 13, 1991
	0.167%	N88470	001	Mar 14, 1984
	0.167%	N89616	001	Jun 13, 1991
	0.2%	N88471	001	Mar 14, 1984
	0.2%	N89617	001	Jun 13, 1991
	0.25%	N88472	001	Mar 14, 1984
	0.25%	N89618	001	Jun 13, 1991
BAXTER HLTHCARE	0.08%	N88144	001	Jul 29, 1983
	0.14%	N88145	001	Mar 26, 1984
	0.25%	N88146	001	Aug 01, 1983
DEY	0.08%	N88187	001	Dec 03, 1982
	0.1%	N87389	001	
	0.17%	N87390	001	
	0.25%	N88188	001	Dec 03, 1982
	1%	N86763	001	
INTL MEDICATION	0.077%	N86651	001	
	0.08%	N86651	002	
	0.1%	N86651	003	
	0.143%	N86651	004	
	0.167%	N86651	005	
	0.2%	N86651	006	
	0.25%	N86651	007	
	1%	N86651	008	
PARKE DAVIS	0.5%	N85997	001	
	1%	N85889	001	
ROXANE	0.1%	N87396	001	
	0.125%	N87025	001	
	0.167%	N88226	001	Sep 16, 1983
	0.2%	N87324	001	
	0.25%	N88275	001	Jun 03, 1983
	1%	N86899	001	
ISOETHARINE HYDROCHLORIDE S/F				
DEY	0.08%	N89817	001	Nov 22, 1988
	0.1%	N89818	001	Nov 22, 1988
	0.17%	N89819	001	Nov 22, 1988
	0.25%	N89820	001	Nov 22, 1988
	1%	N89252	001	Sep 15, 1986

ISOETHARINE MESYLATE

AEROSOL, METERED; INHALATION

BRONKOMETER

SANOFI AVENTIS US	0.34MG/INH	N12339	007	
ISOETHARINE MESYLATE				
ALPHARMA US PHARMS	0.34MG/INH	N87858	001	Aug 21, 1984

ISOFLUROPHATE

OINTMENT; OPHTHALMIC

FLOROPRYL

MERCK	0.025%	N10656	001	
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ISONIAZID

INJECTABLE; INJECTION

RIMIFON

ROCHE	25MG/ML	N08420	002	
	100MG/ML	N08420	003	

SYRUP; ORAL

ISONIAZID

MIKART	50MG/5ML	N81118	001	Jul 21, 1997
LANIAZID				
LANNETT	50MG/5ML	N89243	001	Feb 03, 1986

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 155 (of 274)

ISONIAZID

SYRUP; ORAL				
RIMIFON				
ROCHE	50MG/5ML	N08420	001	
TABLET; ORAL				
DOW-ISONIAZID				
DOW PHARM	300MG	N80330	002	
HYZYD				
MEDPOINTE PHARM HLC	100MG	N80134	003	
	300MG	N80134	004	
INH				
NOVARTIS	300MG	N80935	001	
ISONIAZID				
ANABOLIC	100MG	N84050	001	
DURAMED PHARMS BARR	100MG	N88231	001	Mar 17, 1983
HALSEY	50MG	N83632	001	
IMPAX LABS	100MG	N80153	001	
IVAX PHARMS	100MG	N80270	001	
	300MG	N83610	001	
LILLY	100MG	N08499	002	
	300MG	N08499	003	
MK LABS	100MG	N80941	001	
MUTUAL PHARM	100MG	N80136	001	
PANRAY	50MG	N08428	001	
	100MG	N08428	002	
	300MG	N08428	003	
PERRIGO	100MG	N83060	001	
PHARMAVITE	100MG	N85091	001	
PHOENIX LABS NY	50MG	N80368	001	
	100MG	N80368	002	
PUREPAC PHARM	50MG	N80132	003	Jul 14, 1982
	100MG	N80132	004	Jul 14, 1982
WATSON LABS	50MG	N80522	001	
	100MG	N80401	001	
	100MG	N80523	001	
	100MG	N85790	001	
	300MG	N83178	001	
	300MG	N85784	001	
WHITEWORTH TOWN PLSN	100MG	N80120	002	
NYDRAZID				
BRISTOL MYERS SQUIBB	100MG	N08392	003	
STANOZIDE				
EVERYLIFE	100MG	N80126	001	
	300MG	N80126	002	

ISOPROPAMIDE IODIDE

TABLET; ORAL				
DARBID				
GLAXOSMITHKLINE	EQ 5MG BASE	N10744	001	

ISOPROTERENOL HYDROCHLORIDE

AEROSOL, METERED; INHALATION				
ISOPROTERENOL HYDROCHLORIDE				
3M	0.12MG/INH	N10375	004	
ALPHARMA US PHARMS	0.12MG/INH	N85904	001	
ISUPREL				
SANOFI AVENTIS US	0.103MG/INH	N11178	001	
DISC; INHALATION				
NORISODRINE AEROTROL				
ABBOTT	0.25%	N16814	001	
INJECTABLE; INJECTION				
ISOPROTERENOL HYDROCHLORIDE				
ABRAXIS PHARM	0.2MG/ML	N83431	001	
BAXTER HLTHCARE	0.2MG/ML	N83486	001	

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 156 (of 274)

ISOPROTERENOL HYDROCHLORIDE

SOLUTION; INHALATION

AEROLONE

LILLY 0.25%

N07245 001

ISOPROTERENOL HYDROCHLORIDE

ARMOUR PHARM 0.031%

N87935 001

Nov 18, 1982

0.062%

N87936 001

Nov 18, 1982

DEY 0.5%

N86764 001

Jan 04, 1982

PARKE DAVIS 0.25%

N85994 001

0.5%

N85540 001

ISUPREL

SANOFI AVENTIS US 0.5%

N06327 002

1%

N06327 003

VAPO-ISO

FISONS 0.5%

N16813 001

TABLET; RECTAL, SUBLINGUAL

ISUPREL

SANOFI AVENTIS US 10MG

N06328 001

15MG

N06328 002

ISOPROTERENOL HYDROCHLORIDE; PHENYLEPHRINE BITARTRATE

AEROSOL, METERED; INHALATION

DUO-MEDIAHALER

3M 0.16MG/INH;0.24MG/INH

N13296 001

ISOPROTERENOL SULFATE

AEROSOL, METERED; INHALATION

MEDIAHALER-ISO

3M 0.08MG/INH

N10375 003

POWDER; INHALATION

NORISODRINE

ABBOTT 10%

N06905 003

25%

N06905 002

ISOSORBIDE

SOLUTION; ORAL

ISMOTIC

ALCON 100GM/220ML

N17063 001

ISOSORBIDE DINITRATE

CAPSULE, EXTENDED RELEASE; ORAL

ISORDIL

WYETH AYERST 40MG

N12882 002

Jul 29, 1988

TABLET; ORAL

ISOSORBIDE DINITRATE

MUTUAL PHARM 5MG

N86166 002

Sep 19, 1986

10MG

N86169 001

Sep 19, 1986

20MG

N86167 001

Sep 19, 1986

30MG

N87564 001

Sep 18, 1986

5MG

N89190 001

Feb 17, 1987

10MG

N89191 001

Feb 17, 1987

20MG

N89192 001

Feb 17, 1987

SORBITRATE

ASTRAZENECA 5MG

N16192 001

Apr 01, 1996

10MG

N16192 002

Apr 01, 1996

20MG

N86405 002

Aug 21, 1990

30MG

N88124 001

Aug 21, 1990

40MG

N88125 001

Aug 21, 1990

TABLET; SUBLINGUAL

ISOSORBIDE DINITRATE

MUTUAL PHARM 2.5MG

N84204 001

Sep 18, 1986

5MG

N86168 001

Sep 18, 1986

10MG

N87545 001

Sep 18, 1986

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 157 (of 274)

ISOSORBIDE DINITRATETABLET; SUBLINGUAL
SORBITRATE

ASTRAZENECA	2.5MG	N16191	002	Apr 01, 1996
	5MG	N16191	001	Apr 01, 1996

TABLET, CHEWABLE; ORAL
SORBITRATE

ASTRAZENECA	5MG	N16776	002	Apr 01, 1996
	10MG	N16776	003	Apr 01, 1996

TABLET, EXTENDED RELEASE; ORAL
ISORDIL

WYETH AYERST	40MG	N12882	001	Jul 29, 1988
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ISOSORBIDE MONONITRATETABLET, EXTENDED RELEASE; ORAL
ISOSORBIDE MONONITRATE

IVAX PHARMS	30MG	N75448	002	Aug 07, 2001
	60MG	N75448	001	Jun 19, 2000
	120MG	N75448	003	Aug 07, 2001

ISRADIPINECAPSULE; ORAL
DYNACIRC

RELIANT PHARMS	2.5MG	N19546	001	Dec 20, 1990
	5MG	N19546	002	Dec 20, 1990

IVERMECTINTABLET; ORAL
STROMEKTOL

MERCK	6MG	N50742	001	Nov 22, 1996
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KANAMYCIN SULFATECAPSULE; ORAL
KANTREX

APOTHECON	EQ 500MG BASE	N60516	001	
	EQ 500MG BASE	N61911	001	
	EQ 500MG BASE	N62726	001	Mar 06, 1987

INJECTABLE; INJECTION
KANAMYCIN

BAXTER HLTHCARE	EQ 75MG BASE/2ML	N62324	001	
	EQ 500MG BASE/2ML	N62324	002	
	EQ 1GM BASE/3ML	N62324	003	

KANAMYCIN SULFATE

ABRAXIS PHARM	EQ 75MG BASE/2ML	N62504	001	Apr 05, 1984
	EQ 500MG BASE/2ML	N62504	002	Apr 05, 1984
	EQ 1GM BASE/3ML	N62504	003	Apr 05, 1984

INTL MEDICATION	EQ 500MG BASE/2ML	N62466	001	Sep 30, 1983
	EQ 1GM BASE/3ML	N62466	002	Sep 30, 1983

LOCH	EQ 75MG BASE/2ML	N63021	001	Jul 31, 1992
	EQ 500MG BASE/2ML	N63022	001	Jul 31, 1992
	EQ 1GM BASE/3ML	N63025	001	Jul 31, 1992

PHARMAFAIR	EQ 75MG BASE/2ML	N62668	001	May 07, 1987
	EQ 500MG BASE/2ML	N62672	001	May 07, 1987
	EQ 1GM BASE/3ML	N62669	001	May 07, 1987

SOLOPAK	EQ 75MG BASE/2ML	N62605	003	Feb 26, 1986
	EQ 500MG BASE/2ML	N62605	001	Feb 26, 1986
	EQ 1GM BASE/3ML	N62605	002	Feb 26, 1986

WARNER CHILCOTT	EQ 1GM BASE/3ML	N63092	001	Oct 11, 1989
WATSON LABS	EQ 1GM BASE/3ML	N62520	003	May 09, 1985

KANTREX

APOTHECON	EQ 75MG BASE/2ML	N61655	003	
	EQ 75MG BASE/2ML	N62564	001	Sep 21, 1984
	EQ 500MG BASE/2ML	N61655	001	
	EQ 500MG BASE/2ML	N62564	002	Sep 21, 1984

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 158 (of 274)

KANAMYCIN SULFATE

INJECTABLE; INJECTION

KANTREX

APOTHECON

EQ 1GM BASE/3ML

N61655 002

EQ 1GM BASE/3ML

N62564 003

Sep 21, 1984

KLEBCIL

KING PHARMS

EQ 75MG BASE/2ML

N62170 001

EQ 500MG BASE/2ML

N62170 002

EQ 1GM BASE/3ML

N62170 003

KETOCONAZOLE

CREAM; TOPICAL

NIZORAL

JANSSEN PHARMA

2%

N19084 001

Dec 31, 1985

SUSPENSION; ORAL

NIZORAL

JANSSEN PHARMA

100MG/5ML

N70767 001

Nov 07, 1986

KETOPROFEN

CAPSULE; ORAL

ORUDIS

WYETH AYERST

25MG

N18754 001

Jul 31, 1987

50MG

N18754 002

Jan 09, 1986

75MG

N18754 003

Jan 09, 1986

CAPSULE, EXTENDED RELEASE; ORAL

ORUVAIL

WYETH PHARMS INC

100MG

N19816 003

Feb 08, 1995

150MG

N19816 002

Feb 08, 1995

TABLET; ORAL

ACTRON

BAYER

12.5MG

N20499 001

Oct 06, 1995

KETOPROFEN

PERRIGO

12.5MG

N75364 001

Feb 07, 2002

ORUDIS KT

WYETH CONS

12.5MG

N20429 001

Oct 06, 1995

KETOROLAC TROMETHAMINE

INJECTABLE; INJECTION

KETOROLAC TROMETHAMINE

APOTHECON

15MG/ML

N75348 001

Nov 28, 2000

30MG/ML

N75348 002

Nov 28, 2000

BEDFORD

15MG/ML

N75230 002

Oct 25, 1999

30MG/ML

N75230 001

Oct 25, 1999

HOSPIRA

15MG/ML

N74801 001

Jun 05, 1997

30MG/ML

N74801 002

Jun 05, 1997

TORADOL

ROCHE PALO

15MG/ML

N19698 001

Nov 30, 1989

30MG/ML

N19698 002

Nov 30, 1989

TABLET; ORAL

KETOROLAC TROMETHAMINE

ROXANE

10MG

N74790 001

Jun 26, 1997

KRYPTON, KR-81M

GAS; INHALATION

MPI KRYPTON 81M GAS GENERATOR

GE HEALTHCARE

N/A

N18088 001

LABETALOL HYDROCHLORIDE

INJECTABLE; INJECTION

LABETALOL HYDROCHLORIDE

APOTHECON

5MG/ML

N75355 001

Nov 29, 1999

NORMODYNE

SCHERING

5MG/ML

N18686 001

Aug 01, 1984

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 159 (of 274)

LABETALOL HYDROCHLORIDE

TABLET; ORAL

LABETALOL HYDROCHLORIDE

APOTHECON

100MG

N75223 001

Nov 20, 1998

200MG

N75223 002

Nov 20, 1998

300MG

N75223 003

Nov 20, 1998

NORMODYNE

SCHERING

100MG

N18687 001

Aug 31, 1987

200MG

N18687 002

Aug 01, 1984

300MG

N18687 003

Aug 01, 1984

400MG

N18687 004

Aug 01, 1984

TRANDATE

PROMETHEUS LABS

400MG

N18716 004

Aug 01, 1984

LACTULOSE

SOLUTION; ORAL

CHRONULAC

SANOFI AVENTIS US

10GM/15ML

N17884 001

DUPHALAC

SOLVAY

10GM/15ML

N72372 001

Mar 22, 1989

LACTULOSE

MORTON GROVE

10GM/15ML

N71841 001

Sep 22, 1988

PACO

10GM/15ML

N73160 001

Aug 25, 1992

SOLUTION; ORAL, RECTAL

CEPHULAC

SANOFI AVENTIS US

10GM/15ML

N17657 001

GENERLAC

MORTON GROVE

10GM/15ML

N71842 001

Sep 27, 1988

LACTULOSE

PACO

10GM/15ML

N72029 001

Aug 25, 1992

ROXANE

10GM/15ML

N73590 001

May 29, 1992

SOLVAY

10GM/15ML

N17906 001

PORTALAC

SOLVAY

10GM/15ML

N72374 001

Mar 22, 1989

LAMOTRIGINE

TABLET; ORAL

LAMICTAL

GLAXOSMITHKLINE

50MG

N20241 006

Dec 27, 1994

250MG

N20241 004

Dec 27, 1994

TABLET, CHEWABLE; ORAL

LAMICTAL CD

GLAXOSMITHKLINE

100MG

N20764 003

Aug 24, 1998

LANSOPRAZOLE; NAPROXEN

CAPSULE, DELAYED REL PELLETS, TABLET; ORAL

PREVACID NAPRAPAC 250 (COPACKAGED)

TAP PHARM

15MG, N/A; N/A, 250MG

N21507 002

Nov 14, 2003

PREVACID NAPRAPAC 375 (COPACKAGED)

TAP PHARM

15MG, N/A; N/A, 375MG

N21507 003

Nov 14, 2003

LEFLUNOMIDE

TABLET; ORAL

LEFLUNOMIDE

SANDOZ

10MG

N77085 001

Sep 13, 2005

20MG

N77085 002

Sep 13, 2005

LEUCOVORIN CALCIUM

FOR SOLUTION; ORAL

LEUCOVORIN CALCIUM

MAYNE PHARMA USA

EQ 60MG BASE/VIAL

N08107 003

Jan 30, 1987

INJECTABLE; INJECTION

LEUCOVORIN CALCIUM

ABIC

EQ 3MG BASE/ML

N89352 001

Jun 01, 1988

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 160 (of 274)

LEUCOVORIN CALCIUM

INJECTABLE; INJECTION

LEUCOVORIN CALCIUM

ABIC	EQ 50MG BASE/VIAL	N89353	001	Jun 01, 1988
ABRAXIS PHARM	EQ 50MG BASE/VIAL	N88939	001	Dec 01, 1986
ELKINS SINN	EQ 50MG BASE/VIAL	N70480	001	Jan 02, 1987
	EQ 100MG BASE/VIAL	N81224	001	Jun 03, 1994
MAYNE PHARMA USA	EQ 3MG BASE/ML	N08107	001	
	EQ 50MG BASE/VIAL	N08107	002	
	EQ 100MG BASE/VIAL	N08107	004	May 23, 1988
PHARMACHEMIE USA	EQ 100MG BASE/VIAL	N89915	001	Apr 17, 1997
LEUCOVORIN CALCIUM PRESERVATIVE FREE				
HOSPIRA	EQ 10MG BASE/ML	N40147	001	Jun 25, 1997
WELLCOVORIN				
GLAXOSMITHKLINE	EQ 5MG BASE/ML	N87439	001	Oct 19, 1982
	EQ 25MG BASE/VIAL	N89833	001	Jan 23, 1989
	EQ 50MG BASE/VIAL	N89465	001	Jan 23, 1989
	EQ 100MG BASE/VIAL	N89834	001	Jan 23, 1989

TABLET; ORAL

LEUCOVORIN CALCIUM

PAR PHARM	EQ 5MG BASE	N71600	001	Oct 14, 1987
	EQ 25MG BASE	N71598	001	Oct 14, 1987
WELLCOVORIN				
GLAXOSMITHKLINE	EQ 5MG BASE	N18342	001	Jul 08, 1983
	EQ 25MG BASE	N18342	002	Jul 08, 1983

LEUPROLIDE ACETATE

INJECTABLE; INJECTION

LUPRON DEPOT-PED

TAP PHARM	3.75MG/VIAL, 7.5MG/VIAL	N20263	003	Apr 16, 1993
	7.5MG/VIAL, 7.5MG/VIAL	N20263	004	Apr 16, 1993

LEVALLORPHAN TARTRATE

INJECTABLE; INJECTION

LORFAN

ROCHE	1MG/ML	N10423	001	
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LEVAMISOLE HYDROCHLORIDE

TABLET; ORAL

ERGAMISOL

JANSSEN PHARMA	EQ 50MG BASE	N20035	001	Jun 18, 1990
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LEVOBETAXOLOL HYDROCHLORIDE

SUSPENSION/DROPS; OPHTHALMIC

BETAXON

ALCON	EQ 0.5% BASE	N21114	001	Feb 23, 2000
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LEVOPUPIVACAINE HYDROCHLORIDE

INJECTABLE; INJECTION

CHIROCAINE

PURDUE PHARMA LP	EQ 2.5MG BASE/ML	N20997	001	Aug 05, 1999
	EQ 5MG BASE/ML	N20997	002	Aug 05, 1999
	EQ 7.5MG BASE/ML	N20997	003	Aug 05, 1999

LEVOCABASTINE HYDROCHLORIDE

SUSPENSION/DROPS; OPHTHALMIC

LIVOSTIN

NOVARTIS	EQ 0.05% BASE	N20219	001	Nov 10, 1993
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LEVOCARNITINE

SOLUTION; ORAL

CARNITOR

SIGMA TAU	1GM/10ML	N18948	002	Apr 27, 1988
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DISCONTINUED DRUG PRODUCT LIST

6 - 161 (of 274)

LEVODOPA

CAPSULE; ORAL

BENDOPA

VALEANT PHARM INTL

100MG

N16948 003

250MG

N16948 001

500MG

N16948 002

DOPAR

SHIRE

100MG

N16913 003

250MG

N16913 001

500MG

N16913 002

LARODOPA

ROCHE

100MG

N16912 002

250MG

N16912 001

500MG

N16912 006

TABLET; ORAL

DOPAR

SHIRE

250MG

N16913 004

500MG

N16913 005

LARODOPA

ROCHE

100MG

N16912 005

250MG

N16912 003

500MG

N16912 004

LEVOMEPRMAZINE

INJECTABLE; INJECTION

LEVOPROME

IMMUNEX

20MG/ML

N15865 001

LEVOMETHADYL ACETATE HYDROCHLORIDE

CONCENTRATE; ORAL

ORLAAM

ROXANE

10MG/ML

N20315 001

Jul 09, 1993

LEVONORDEFIN; MEPIVACAINE HYDROCHLORIDE

INJECTABLE; INJECTION

ARESTOCAINE HYDROCHLORIDE W/ LEVONORDEFIN

SOLVAY

0.05MG/ML; 2%

N85010 001

LEVONORDEFIN; PROCAINE HYDROCHLORIDE; PROPOXYCAINE HYDROCHLORIDE

INJECTABLE; INJECTION

RAVOCAINE AND NOVOCAIN W/ NEO-COBEFRIN

EASTMAN KODAK

0.05MG/ML; 2%; 0.4%

N08592 007

LEVONORGESTREL

IMPLANT; IMPLANTATION

LEVONORGESTREL

WYETH PHARMS INC

75MG/IMPLANT

N20627 001

Aug 15, 1996

NORPLANT

POPULATION COUNCIL

36MG/IMPLANT

N19897 001

Dec 10, 1990

NORPLANT SYSTEM IN PLASTIC CONTAINER

WYETH PHARMS INC

36MG/IMPLANT

N20088 001

Dec 10, 1990

LEVOPROPOXYPHENE NAPSYLATE, ANHYDROUS

CAPSULE; ORAL

NOVRAD

LILLY

EQ 50MG BASE

N12928 006

EQ 100MG BASE

N12928 004

SUSPENSION; ORAL

NOVRAD

LILLY

EQ 50MG BASE/5ML

N12928 002

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 162 (of 274)

LEVORPHANOL TARTRATEINJECTABLE; INJECTION
LEVO-DROMORAN

VALEANT PHARM INTL 2MG/ML

N08719 001 Dec 19, 1991

LIDOCAINE

AEROSOL; ORAL

XYLOCAINE

ASTRAZENECA 10%

N14394 001

FILM, EXTENDED RELEASE; BUCCAL

LIDOCAINE

NOVEN 23MG/PATCH

N20575 001 May 21, 1996

OINTMENT; TOPICAL

ALPHACAINE

CARLISLE

5%

N84944 001

5%

N84946 001

5%

N84947 001

XYLOCAINE

ASTRAZENECA 5%

N08048 001

SOLUTION; TOPICAL

XYLOCAINE

ASTRAZENECA 5%

N14127 001

SUPPOSITORY; RECTAL

XYLOCAINE

ASTRAZENECA 100MG

N13077 001

LIDOCAINE HYDROCHLORIDE

INJECTABLE; INJECTION

ALPHACAINE HYDROCHLORIDE

CARLISLE 2%

N84721 001

LIDOCAINE HYDROCHLORIDE

ABBOTT 10%

N87980 001 Feb 02, 1983

20%

N89362 001 May 25, 1988

ABRAXIS PHARM

1%

N80390 001

1%

N80420 001

1%

N86761 001

1.5%

N80420 005

2%

N17508 001

2%

N80390 002

2%

N80420 002

2%

N80420 004

2%

N86761 002

4%

N17508 002

20%

N17508 004

AKORN

1%

N85037 001

2%

N85037 002

BAXTER HLTHCARE

1%

N80407 001

2%

N80407 002

BEL MAR

1%

N80710 001

2%

N80760 001

DELL LABS

1%

N83387 001

2%

N83388 001

ELKINS SINN

0.5%

N85131 001

4%

N84626 001

GD SEARLE LLC

1%

N83135 001

2%

N83135 002

HOSPIRA

1.5%

N88330 001 May 17, 1984

INTL MEDICATION

1%

N17701 002

2%

N17701 001

1GM/VIAL

N18543 001

2GM/VIAL

N18543 002

MILES

1%

N80414 001

2%

N80414 002

WATSON LABS

1%

N80377 001

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 163 (of 274)

LIDOCAINE HYDROCHLORIDE

INJECTABLE; INJECTION

LIDOCAINE HYDROCHLORIDE

WATSON LABS

1%

N83627 001

2%

N80377 002

2%

N83627 002

WYETH AYERST

1%

N83083 001

2%

N83083 002

LIDOCAINE HYDROCHLORIDE 0.1% AND DEXTROSE 5% IN PLASTIC CONTAINER

BAXTER HLTHCARE

100MG/100ML

N18461 001

LIDOCAINE HYDROCHLORIDE 0.2% AND DEXTROSE 5% IN PLASTIC CONTAINER

B BRAUN

200MG/100ML

N18967 001

Mar 30, 1984

LIDOCAINE HYDROCHLORIDE 0.2% IN DEXTROSE 5% IN PLASTIC CONTAINER

ABBOTT

200MG/100ML

N18954 001

Jul 09, 1985

LIDOCAINE HYDROCHLORIDE 0.4% AND DEXTROSE 5% IN PLASTIC CONTAINER

B BRAUN

400MG/100ML

N18967 002

Mar 30, 1984

LIDOCAINE HYDROCHLORIDE 0.8% AND DEXTROSE 5% IN PLASTIC CONTAINER

B BRAUN

800MG/100ML

N18967 003

Mar 30, 1984

LIDOCAINE HYDROCHLORIDE PRESERVATIVE FREE

BAXTER HLTHCARE

1%

N84625 001

2%

N84625 002

INTL MEDICATION

4%

N17702 002

LIDOCATON

PHARMATON

2%

N84727 001

Aug 17, 1983

XYLOCAINE

ABRAXIS BIOSCIENCE

2%

N06488 002

ASTRAZENECA

1%

N10418 005

1.5%

N10418 009

2%

N10418 007

INJECTABLE; SPINAL

XYLOCAINE 1.5% W/ DEXTROSE 7.5%

ABRAXIS BIOSCIENCE

1.5%

N16297 001

XYLOCAINE 5% W/ GLUCOSE 7.5%

ASTRAZENECA

5%

N10496 002

Jul 07, 1982

SOLUTION; ORAL

LIDOCAINE HYDROCHLORIDE VISCOUS

INTL MEDICATION

2%

N86389 001

Feb 02, 1982

SOLUTION; TOPICAL

LARYNGOTRACHEAL ANESTHESIA KIT

KENDALL IL

4%

N87931 001

Jun 10, 1983

LIDOCAINE HYDROCHLORIDE

PACO

4%

N89688 001

Jun 30, 1989

PEDIATRIC LTA KIT

ABBOTT

2%

N88572 001

Jul 31, 1984

LIDOCAINE; PRILOCAINE

DISC; TOPICAL

EMLA

ASTRAZENECA

2.5%;2.5%

N20962 001

Feb 04, 1998

LINCOMYCIN HYDROCHLORIDE

CAPSULE; ORAL

LINCOCIN

PHARMACIA AND UPJOHN EQ 250MG BASE

N50316 001

EQ 500MG BASE

N50316 002

INJECTABLE; INJECTION

LINCOMYCIN HYDROCHLORIDE

WATSON LABS

EQ 300MG BASE/ML

N63180 001

Apr 16, 1991

LINDANE

CREAM; TOPICAL

KWELL

REED AND CARNRICK

1%

N06309 001

1%

N84218 001

DISCONTINUED DRUG PRODUCT LIST

6 - 164 (of 274)

LINDANE

LOTION; TOPICAL
GAMENE

SOLA BARNES HIND 1%

N84989 001

KWELL

REED AND CARNRICK 1%

N06309 003

1%

N84218 002

SCABENE

STIEFEL 1%

N86769 001

SHAMPOO; TOPICAL
GAMENE

SOLA BARNES HIND 1%

N84988 001

KWELL

REED AND CARNRICK 1%

N10718 001

1%

N84219 001

SCABENE

STIEFEL 1%

N87940 001

Apr 08, 1983

LINEZOLID

TABLET; ORAL
ZYVOX

PHARMACIA AND UPJOHN 400MG **Federal Register determination
that product was not discontinued or
withdrawn for safety or efficacy
reasons**

N21130 001

Apr 18, 2000

LIOTHYRONINE SODIUM

TABLET; ORAL
LIOTHYRONINE SODIUM

WATSON LABS EQ 0.025MG BASE
EQ 0.05MG BASE

N85755 001

Jan 25, 1982

N85753 001

Feb 03, 1982

LIOTRIX (T4;T3)

TABLET; ORAL

EUTHROID-0.5

PARKE DAVIS 0.03MG;0.0075MG

N16680 001

EUTHROID-1

PARKE DAVIS 0.06MG;0.015MG

N16680 002

EUTHROID-2

PARKE DAVIS 0.12MG;0.03MG

N16680 003

EUTHROID-3

PARKE DAVIS 0.18MG;0.045MG

N16680 004

THYROLAR-5

FOREST LABS 0.25MG;0.0625MG

N16807 006

LISINAPRIL

TABLET; ORAL
PRINIVIL

MERCK 2.5MG

N19558 006

Jan 28, 1994

LITHIUM CARBONATE

CAPSULE; ORAL
LITHIUM CARBONATE
ABLE

150MG

N76823 001

Jun 29, 2004

300MG

N76121 001

Sep 27, 2001

300MG

N76823 002

Jun 29, 2004

600MG

N76823 003

Jun 29, 2004

USL PHARMA

300MG

N72542 001

Feb 01, 1989

WATSON LABS

300MG

N70407 001

Mar 19, 1987

LITHONATE

SOLVAY 300MG

N16782 001

TABLET; ORAL

ESKALITH

GLAXOSMITHKLINE 300MG

N17971 001

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 165 (of 274)

LITHIUM CARBONATE

TABLET; ORAL

LITHANE

BAYER PHARMS	300MG	N18833	001	Jul 18, 1985
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LITHIUM CARBONATE

PFIZER	300MG	N16834	001	
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LITHOTABS

SOLVAY	300MG	N16980	001	
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TABLET, EXTENDED RELEASE; ORAL

ESKALITH CR

GLAXOSMITHKLINE	450MG	N18152	001	Mar 29, 1982
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LITHIUM CARBONATE

ABLE	300MG	N76382	001	Apr 21, 2003
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BARR	450MG	N76366	001	Aug 21, 2003
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LITHIUM CITRATE

SYRUP; ORAL

LITHONATE

SOLVAY	EQ 300MG CARBONATE/5ML	N17672	001	
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LOPERAMIDE HYDROCHLORIDE

CAPSULE; ORAL

IMODIUM

MCNEIL PED	2MG	N17690	001	
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LOPERAMIDE HYDROCHLORIDE

ROXANE	2MG	N73080	001	Nov 27, 1991
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SOLUTION; ORAL

IMODIUM

JANSSEN PHARMA	1MG/5ML	N19037	001	Jul 31, 1984
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LOPERAMIDE HYDROCHLORIDE

ALPHARMA US PHARMS	1MG/5ML	N73187	001	Sep 15, 1992
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WATSON LABS	1MG/5ML	N73062	001	May 28, 1993
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TABLET; ORAL

LOPERAMIDE HYDROCHLORIDE

ABLE	2MG	N73528	001	Nov 30, 1993
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PERRIGO	2MG	N74194	001	Oct 30, 1992
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LORACARBEF

CAPSULE; ORAL

LORABID

KING PHARMS	200MG	N50668	001	Dec 31, 1991
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	400MG	N50668	002	Apr 05, 1996
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FOR SUSPENSION; ORAL

LORABID

KING PHARMS	100MG/5ML	N50667	001	Dec 31, 1991
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	200MG/5ML	N50667	002	Dec 31, 1991
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LORATADINE

SYRUP; ORAL

CLARITIN HIVES RELIEF

SCHERING PLOUGH	1MG/ML **Federal Register	N20641	003	Nov 19, 2003
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determination that product was not discontinued or withdrawn for safety or efficacy reasons**

TABLET; ORAL

LORATADINE

PERRIGO	10MG	N21512	001	Jun 24, 2004
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LORATADINE; PSEUDOEPHEDRINE SULFATE

TABLET, EXTENDED RELEASE; ORAL

CLARITIN-D

SCHERING PLOUGH	5MG;120MG	N19670	002	Nov 27, 2002
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DISCONTINUED DRUG PRODUCT LIST

6 - 166 (of 274)

LORAZEPAM

INJECTABLE; INJECTION

LORAZEPAM

AKORN	2MG/ML	N74974	001	Jul 23, 1998
BAXTER HLTHCARE	2MG/ML	N74496	001	Sep 28, 1998
	4MG/ML	N74496	002	Sep 28, 1998
CLONMEL HLTHCARE	2MG/ML	N74793	001	Mar 16, 2000
	4MG/ML	N74793	002	Mar 16, 2000
MARSAM PHARMS LLC	1MG/0.5ML	N74551	003	Sep 12, 1996
	2MG/ML	N74535	001	Sep 12, 1996
	2MG/ML	N74551	001	Sep 12, 1996
	4MG/ML	N74535	002	Sep 12, 1996
	4MG/ML	N74551	002	Sep 12, 1996
TAYLOR	2MG/ML	N75025	001	Jul 23, 1998

TABLET; ORAL

LORAZ

QUANTUM PHARMICS	0.5MG	N70200	001	Aug 09, 1985
	1MG	N70201	001	Aug 09, 1985
	2MG	N70202	001	Aug 09, 1985

LORAZEPAM

AM THERAP	0.5MG	N70727	001	Mar 07, 1986
	1MG	N70728	001	Mar 07, 1986
	2MG	N70729	001	Mar 07, 1986
HALSEY	0.5MG	N71434	001	Sep 01, 1987
	1MG	N71435	001	Sep 01, 1987
	2MG	N71436	001	Sep 01, 1987
MUTUAL PHARM	0.5MG	N70472	001	Dec 10, 1985
	1MG	N70473	001	Dec 10, 1985
	2MG	N70474	001	Dec 10, 1985
PAR PHARM	0.5MG	N70675	001	Dec 01, 1986
	1MG	N70676	001	Dec 01, 1986
	2MG	N70677	001	Dec 01, 1986
SUPERPHARM	0.5MG	N71245	001	Feb 09, 1987
	1MG	N71246	001	Feb 09, 1987
	2MG	N71247	001	Feb 09, 1987
USL PHARMA	1MG	N70539	001	Dec 22, 1986
	2MG	N70540	001	Dec 22, 1986
WARNER CHILCOTT	1MG	N71038	001	Jan 12, 1988
	2MG	N71039	001	Jan 12, 1988
WATSON LABS	0.5MG	N71086	001	Mar 23, 1987
	1MG	N71087	001	Mar 23, 1987
	2MG	N71088	001	Mar 23, 1987

LOTEPREDNOL ETABONATE

SUSPENSION/DROPS; OPHTHALMIC

LOTEMAX

PHARMOS	0.5%	N20841	001	Mar 09, 1998
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LOVASTATIN

TABLET; ORAL

MEVACOR

MERCK	10MG	N19643	002	Mar 28, 1991
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LOXAPINE HYDROCHLORIDE

CONCENTRATE; ORAL

LOXITANE C

WATSON LABS	EQ 25MG BASE/ML	N17658	001	
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INJECTABLE; INJECTION

LOXITANE IM

WATSON LABS	EQ 50MG BASE/ML	N18039	001	
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DISCONTINUED DRUG PRODUCT LIST

6 - 167 (of 274)

LOXAPINE SUCCINATE

TABLET; ORAL
LOXITANE

WATSON LABS

EQ 10MG BASE

N17525 006

EQ 25MG BASE

N17525 007

EQ 50MG BASE

N17525 008

LYPRESSIN

SOLUTION; NASAL
DIAPID

NOVARTIS

0.185MG/ML

N16755 001

MAGNESIUM CHLORIDE; POTASSIUM CHLORIDE; POTASSIUM PHOSPHATE, MONOBASIC; SODIUM ACETATE; SODIUM CHLORIDE; SODIUM GLUCONATE; SODIUM PHOSPHATE, DIBASIC, HEPTAHYDRATE

INJECTABLE; INJECTION

ISOLYTE S PH 7.4 IN PLASTIC CONTAINER

B BRAUN

30MG/100ML;37MG/100ML;0.82MG/100ML;370MG/100ML;530MG/100ML;500MG/100ML;12MG/100ML

N19006 001

Apr 04, 1984

MAGNESIUM CHLORIDE; POTASSIUM CHLORIDE; SODIUM ACETATE; SODIUM CHLORIDE; SODIUM GLUCONATE

INJECTABLE; INJECTION

ISOLYTE S IN PLASTIC CONTAINER

B BRAUN

30MG/100ML;37MG/100ML;370MG/100ML;530MG/100ML;500MG/100ML

N18252 001

SOLUTION; IRRIGATION

PHYSIOSOL IN PLASTIC CONTAINER

HOSPIRA

14MG/100ML;37MG/100ML;222MG/100ML;526MG/100ML;502MG/100ML

N18406 001

SYNOVALYTE IN PLASTIC CONTAINER

BAXTER HLTHCARE

30MG/100ML;37MG/100ML;368MG/100ML;526MG/100ML;502MG/100ML

N19326 001

Jan 25, 1985

MANGAFODIPIR TRISODIUM

INJECTABLE; INJECTION

TESLASCAN

GE HEALTHCARE

37.9MG/ML

N20652 001

Nov 26, 1997

MANGANESE CHLORIDE TETRAHYDRATE

FOR SOLUTION; ORAL

LUMENHANCE

BRACCO

3.49MG/GM

N20686 001

Dec 19, 1997

MANGANESE SULFATE

INJECTABLE; INJECTION

MANGANESE SULFATE

ABRAXIS PHARM

EQ 0.1MG MANGANESE/ML

N19228 001

May 05, 1987

MANNITOL

INJECTABLE; INJECTION

MANNITOL 10%

HOSPIRA

10GM/100ML

N16269 002

MILES

10GM/100ML

N16472 002

MANNITOL 15%

HOSPIRA

15GM/100ML

N16269 003

MILES

15GM/100ML

N16472 005

MANNITOL 20%

HOSPIRA

20GM/100ML

N16269 004

MILES

20GM/100ML

N16472 004

MANNITOL 25%

ABRAXIS PHARM

12.5GM/50ML

N86754 001

ASTRAZENECA

12.5GM/50ML

N89239 001

May 06, 1987

12.5GM/50ML

N89240 001

May 06, 1987

HOSPIRA

12.5GM/50ML

N16269 005

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 168 (of 274)

MANNITOL

INJECTABLE; INJECTION

MANNITOL 25%

MERCK

12.5GM/50ML

N05620 001

WATSON LABS

12.5GM/50ML

N87460 001

Jun 27, 1983

MANNITOL 5%

HOSPIRA

5GM/100ML

N16269 001

SOLUTION; IRRIGATION

RESECTISOL

B BRAUN

5GM/100ML

N16704 002

MANNITOL; SORBITOL

SOLUTION; IRRIGATION

SORBITOL-MANNITOL

HOSPIRA

540MG/100ML; 2.7GM/100ML

N80224 001

MAPROTILINE HYDROCHLORIDE

TABLET; ORAL

LUDIOMIL

NOVARTIS

25MG

N17543 001

50MG

N17543 002

75MG

N17543 003

Sep 30, 1982

MAPROTILINE HYDROCHLORIDE

AM THERAP

25MG

N72129 001

Jan 14, 1988

50MG

N72130 001

Jan 14, 1988

75MG

N72131 001

Jan 14, 1988

WATSON LABS

25MG

N71943 001

Dec 30, 1987

50MG

N71944 001

Dec 30, 1987

75MG

N71945 001

Dec 30, 1987

MASOPROCOL

CREAM; TOPICAL

ACTINEX

UNIV AZ CANCER CTR

10%

N19940 001

Sep 04, 1992

MAZINDOL

TABLET; ORAL

MAZANOR

WYETH AYERST

1MG

N17980 002

2MG

N17980 001

SANOEX

NOVARTIS

1MG

N17247 001

2MG

N17247 002

MEBENDAZOLE

TABLET, CHEWABLE; ORAL

VERMOX

MCNEIL PED

100MG

N17481 001

MEBUTAMATE

TABLET; ORAL

DORMATE

MEDPOINTE PHARM HLC

600MG

N17374 001

MECLIZINE HYDROCHLORIDE

TABLET; ORAL

MECLIZINE HYDROCHLORIDE

ABC HOLDING

12.5MG

N85253 001

25MG

N85252 001

ANABOLIC

25MG

N85891 001

IVAX PHARMS

12.5MG

N83784 001

KV PHARM

12.5MG

N85524 001

25MG

N85523 001

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 169 (of 274)

MECLIZINE HYDROCHLORIDE

TABLET; ORAL

MECLIZINE HYDROCHLORIDE

SUPERPHARM	12.5MG	N89113	001	Aug 20, 1985
	25MG	N89114	001	Aug 20, 1985
UDL	12.5MG	N88256	001	Jun 13, 1983
	25MG	N88257	001	Jun 13, 1983
VANGARD	12.5MG	N87877	001	Apr 20, 1982
	25MG	N87620	001	Jan 04, 1982
WATSON LABS	12.5MG	N85195	001	
	12.5MG	N85269	001	

TABLET, CHEWABLE; ORAL

MECLIZINE HYDROCHLORIDE

ANABOLIC	25MG	N86392	001	
IVAX PHARMS	25MG	N84976	001	

MECLOCYCLINE SULFOSALICYLATE

CREAM; TOPICAL

MECLAN

JOHNSON AND JOHNSON	1%	N50518	001	
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MECLOFENAMATE SODIUM

CAPSULE; ORAL

MECLODIUM

QUANTUM PHARMICS	EQ 50MG BASE	N71380	001	Jul 14, 1987
	EQ 100MG BASE	N71381	001	Jul 14, 1987

MECLOFENAMATE SODIUM

AM THERAP	EQ 50MG BASE	N71362	001	Feb 10, 1987
	EQ 100MG BASE	N71363	001	Feb 10, 1987
BARR	EQ 50MG BASE	N72848	001	Mar 20, 1989
	EQ 100MG BASE	N72809	001	Mar 20, 1989
PAR PHARM	EQ 50MG BASE	N72077	001	Mar 10, 1988
	EQ 100MG BASE	N72078	001	Mar 10, 1988
USL PHARMA	EQ 50MG BASE	N71007	001	Mar 25, 1988
	EQ 100MG BASE	N71008	001	Mar 25, 1988
VITARINE	EQ 50MG BASE	N71710	001	Jun 15, 1988
	EQ 100MG BASE	N71684	001	Jun 15, 1988
WATSON LABS	EQ 50MG BASE	N70400	001	Nov 25, 1986
	EQ 50MG BASE	N71640	001	Aug 11, 1987
	EQ 100MG BASE	N70401	001	Nov 25, 1986
	EQ 100MG BASE	N71641	001	Aug 11, 1987

MECLOMEN

PARKE DAVIS	EQ 50MG BASE	N18006	001	
	EQ 100MG BASE	N18006	002	

MEDROXYPROGESTERONE ACETATE

INJECTABLE; INJECTION

DEPO-PROVERA

PHARMACIA AND UPJOHN	100MG/ML **Federal Register determination that product was not discontinued or withdrawn for safety or efficacy reasons**	N12541	002	
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TABLET; ORAL

AMEN

AMARIN PHARMS	10MG	N83242	001	
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CURRETAB

SOLVAY	10MG	N85686	001	
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CYCRIN

ESI	2.5MG	N81239	001	Oct 30, 1992
	5MG	N81240	001	Oct 30, 1992
	10MG	N89386	001	Sep 09, 1987

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 170 (of 274)

MEFLOQUINE HYDROCHLORIDE

TABLET; ORAL

MEFLOQUINE HYDROCHLORIDE

US ARMY WALTER REED 250MG

N19578 001 May 02, 1989

MEGESTROL ACETATE

TABLET; ORAL

MEGESTROL ACETATE

USL PHARMA 20MG
40MGN70646 001 Oct 02, 1987
N70647 001 Oct 02, 1987MEMANTINE HYDROCHLORIDE

TABLET; ORAL

NAMENDA

FOREST LABS 15MG
20MGN21487 003 Oct 16, 2003
N21487 004 Oct 16, 2003MENADIOL SODIUM DIPHOSPHATE

INJECTABLE; INJECTION

KAPPADIONE

LILLY 10MG/ML

N05725 001

SYNKAYVITE

ROCHE 5MG/ML
10MG/ML
37.5MG/MLN03718 004
N03718 006
N03718 008

TABLET; ORAL

SYNKAYVITE

ROCHE 5MG

N03718 010

MENADIONE

TABLET; ORAL

MENADIONE

LILLY 5MG

N02139 003

MENOTROPINS (FSH; LH)

INJECTABLE; INJECTION

HUMEGON

ORGANON USA INC 75 IU/VIAL; 75 IU/VIAL
150 IU/VIAL; 150 IU/VIALN20328 001 Sep 01, 1994
N20328 002 Sep 01, 1994

MENOTROPINS

FERRING 75 IU/VIAL; 75 IU/VIAL
150 IU/VIAL; 150 IU/VIALN73598 001 Jan 30, 1997
N73599 001 Jan 30, 1997

PERGONAL

SERONO 75 IU/AMP; 75 IU/AMP
150 IU/AMP; 150 IU/AMPN17646 001
N17646 002 May 20, 1985

REPRONEX

FERRING 150 IU/VIAL; 150 IU/VIAL

N21047 002 Aug 27, 1999

MEPENZOLATE BROMIDE

SOLUTION; ORAL

CANTIL

SANOFI AVENTIS US 25MG/5ML

N10679 004

MEPERIDINE HYDROCHLORIDE

INJECTABLE; INJECTION

DEMEROL

SANOFI AVENTIS US 25MG/ML
50MG/ML
75MG/ML
100MG/MLN05010 007
N05010 002
N05010 009
N05010 003

MEPERIDINE HYDROCHLORIDE

ABBOTT 25MG/ML
50MG/MLN80388 001
N80385 001

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 171 (of 274)

MEPERIDINE HYDROCHLORIDE

INJECTABLE; INJECTION

MEPERIDINE HYDROCHLORIDE

ABBOTT

50MG/ML

N80387 001

75MG/ML

N80389 001

100MG/ML

N80386 001

ASTRAZENECA

25MG/ML

N89781 001 Mar 31, 1989

50MG/ML

N89782 001 Mar 31, 1989

50MG/ML

N89783 001 Mar 31, 1989

50MG/ML

N89784 001 Mar 31, 1989

75MG/ML

N89785 001 Mar 31, 1989

100MG/ML

N89786 001 Mar 31, 1989

100MG/ML

N89787 001 Mar 31, 1989

100MG/ML

N89788 001 Mar 31, 1989

BAXTER HLTHCARE

25MG/ML

N88279 001 Jun 15, 1984

50MG/ML

N88280 001 Jun 15, 1984

75MG/ML

N88281 001 Jun 15, 1984

100MG/ML

N88282 001 Jun 15, 1984

INTL MEDICATION

10MG/ML

N86332 001

PARKE DAVIS

50MG/ML

N80364 002

75MG/ML

N80364 003

100MG/ML

N80364 001

MEPERIDINE HYDROCHLORIDE PRESERVATIVE FREE

MAYNE PHARMA USA

10MG/ML

N40305 001 Mar 10, 1999

TABLET; ORAL

MEPERIDINE HYDROCHLORIDE

WYETH AYERST

50MG

N80454 001

MEPERIDINE HYDROCHLORIDE; PROMETHAZINE HYDROCHLORIDE

INJECTABLE; INJECTION

MEPERGAN

BAXTER HLTHCARE CORP 25MG/ML;25MG/ML

N11730 001

MEPHENTERMINE SULFATE

INJECTABLE; INJECTION

WYAMINE SULFATE

BAXTER HLTHCARE CORP EQ 15MG BASE/ML

N08248 002

EQ 30MG BASE/ML

N08248 001

MEPHENYTOIN

TABLET; ORAL

MESANTOIN

NOVARTIS

100MG

N06008 001

MEPIVACAINE HYDROCHLORIDE

INJECTABLE; INJECTION

ARESTOCAINE HYDROCHLORIDE

SOLVAY

3%

N84777 002 Apr 18, 1982

MEPIVACAINE HYDROCHLORIDE

INTL MEDICATION

1%

N87509 001 Oct 05, 1982

MEPREDNISONE

TABLET; ORAL

BETAPAR

SCHERING

4MG

N16053 002

MEPROBAMATE

CAPSULE; ORAL

EQUANIL

WYETH AYERST

400MG

N12455 002

CAPSULE, EXTENDED RELEASE; ORAL

MEPROSPAN

MEDPOINTE PHARM HLC

200MG

N11284 001

400MG

N11284 002

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 172 (of 274)

MEPROBAMATE

TABLET; ORAL

AMOSENE

FERNDAL LABS

400MG

N84030 001

BAMATE

ALRA

200MG

N80380 001

400MG

N80380 002

EQUANIL

WYETH AYERST

200MG

N10028 005

400MG

N10028 004

MEPRIAM

TEVA

400MG

N16069 001

MEPROBAMATE

ANABOLIC

200MG

N84220 001

400MG

N84589 001

BARR

600MG

N84230 001

ELKINS SINN

200MG

N15426 002

400MG

N15426 001

HEATHER

400MG

N16928 003

600MG

N84329 001

IMPAX LABS

200MG

N14322 002

400MG

N14322 001

IVAX PHARMS

200MG

N15438 001

400MG

N15438 002

600MG

N84181 001

LANNETT

200MG

N14882 002

400MG

N14882 001

LEDERLE

400MG

N86299 001

LEE KM

400MG

N89538 001

Nov 25, 1987

LEINER

400MG

N14601 001

MALLARD

400MG

N15072 002

MK LABS

200MG

N14368 004

400MG

N14368 002

MUTUAL PHARM

200MG

N80699 001

400MG

N80699 002

MYLAN

400MG

N83618 001

PARKE DAVIS

200MG

N84744 001

400MG

N84744 002

PERRIGO

200MG

N84546 001

400MG

N84547 001

PHARMAVITE

400MG

N84438 001

PHARMERAL

400MG

N84153 001

PUREPAC PHARM

200MG

N84804 001

400MG

N84804 002

ROXANE

600MG

N84332 001

SANDOZ

200MG

N14547 002

400MG

N14547 001

400MG

N80655 001

SCHERER LABS

400MG

N83343 001

SOLVAY

200MG

N84435 001

STANLABS PHARM

200MG

N14474 002

400MG

N14474 004

TABLICAPS

400MG

N83494 001

USL PHARMA

200MG

N87825 001

Mar 18, 1982

400MG

N87826 001

Mar 18, 1982

VALEANT PHARM INTL

200MG

N15139 006

400MG

N15139 005

VANGARD

400MG

N88011 001

Jul 14, 1982

WATSON LABS

200MG

N85720 001

400MG

N85721 001

600MG

N84274 001

600MG

N85719 001

WEST WARD

200MG

N15417 003

400MG

N15417 002

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 173 (of 274)

MEPROBAMATETABLET; ORAL
MEPROBAMATE

WHITEWORTH TOWN PLSN	200MG	N83830	001
	400MG	N83442	001

MILTOWN

MEDPOINTE PHARM HLC	200MG	N09698	004
	400MG	N09698	002
	600MG	N83919	001

NEURAMATE

HALSEY	200MG	N14359	002
	400MG	N14359	001

TRANMEP

SOLVAY	400MG	N16249	001
	400MG	N84369	001

MERSALYL SODIUM; THEOPHYLLINEINJECTABLE; INJECTION
MERSALYL-THEOPHYLLINE

WATSON LABS	100MG/ML ; 50MG/ML	N84875	001
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MESALAMINE

SUPPOSITORY; RECTAL

CANASA

AXCAN SCANDIPHARM	500MG	N21252	001	Jan 05, 2001
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ROWASA

ALAVEN PHARM	500MG	N19919	001	Dec 18, 1990
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MESORIDAZINE BESYLATE

CONCENTRATE; ORAL

SERENTIL

NOVARTIS	EQ 25MG BASE/ML	N16997	001
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INJECTABLE; INJECTION

SERENTIL

NOVARTIS	EQ 25MG BASE/ML	N16775	001
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TABLET; ORAL

SERENTIL

NOVARTIS	EQ 10MG BASE	N16774	001
	EQ 25MG BASE	N16774	002
	EQ 50MG BASE	N16774	003
	EQ 100MG BASE	N16774	004

MESTRANOL; NORETHINDRONE

TABLET; ORAL-20

NORINYL

WATSON LABS	0.1MG; 2MG	N13625	004
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TABLET; ORAL-21

NORINYL 1+80 21-DAY

GD SEARLE LLC	0.08MG; 1MG	N16724	001
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ORTHO-NOVUM 1/50 21

ORTHO MCNEIL PHARM	0.05MG; 1MG	N12728	004
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ORTHO-NOVUM 1/80 21

ORTHO MCNEIL PHARM	0.08MG; 1MG	N16715	001
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ORTHO-NOVUM 10-21

ORTHO MCNEIL PHARM	0.06MG; 10MG	N12728	001
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ORTHO-NOVUM 2-21

ORTHO MCNEIL PHARM	0.1MG; 2MG	N12728	005
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TABLET; ORAL-28

NORINYL 1+80 28-DAY

GD SEARLE LLC	0.08MG; 1MG	N16725	001
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ORTHO-NOVUM 1/80 28

ORTHO MCNEIL PHARM	0.08MG; 1MG	N16715	002
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27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 174 (of 274)

MESTRANOL; NORETHYNODREL

TABLET; ORAL

ENOVID

GD SEARLE LLC

0.075MG;5MG
0.15MG;9.85MGN10976 008
N10976 005

TABLET; ORAL-20

ENOVID

GD SEARLE LLC

0.075MG;5MG

N10976 004

ENOVID-E

GD SEARLE LLC

0.1MG;2.5MG

N10976 006

TABLET; ORAL-21

ENOVID-E 21

GD SEARLE LLC

0.1MG;2.5MG

N10976 007

METAPROTERENOL SULFATE

SOLUTION; INHALATION

ALUPENT

BOEHRINGER INGELHEIM

5%

N17659 001

METAPROTERENOL SULFATE

ASTRAZENECA

0.4%

N71275 001 Jul 27, 1988

0.6%

N71018 001 Jul 27, 1988

DEY

0.33%

N71806 001 Aug 05, 1988

0.5%

N71805 001 Aug 05, 1988

5%

N70805 001 Aug 17, 1987

MORTON GROVE

5%

N72190 001 Jun 07, 1988

PROMETA

MURO

5%

N73340 001 Mar 30, 1992

SYRUP; ORAL

ALUPENT

BOEHRINGER INGELHEIM

10MG/5ML

N17571 001

METAPROTERENOL SULFATE

MORTON GROVE

10MG/5ML

N71656 001 Oct 13, 1987

TEVA

10MG/5ML

N72761 001 Feb 27, 1992

TEVA PHARMS

10MG/5ML

N73034 001 Aug 30, 1991

PROMETA

MURO

10MG/5ML

N72023 001 Sep 15, 1988

TABLET; ORAL

ALUPENT

BOEHRINGER INGELHEIM

10MG

N15874 002

20MG

N15874 001

METAPROTERENOL SULFATE

AM THERAP

10MG

N72054 001 Jun 23, 1988

20MG

N72055 001 Jun 23, 1988

USL PHARMA

10MG

N71013 001 Jan 25, 1988

20MG

N71014 001 Jan 25, 1988

METARAMINOL BITARTRATE

INJECTABLE; INJECTION

ARAMINE

MERCK

EQ 10MG BASE/ML

N09509 002 Dec 22, 1987

METARAMINOL BITARTRATE

ABRAXIS PHARM

EQ 10MG BASE/ML

N80431 001

ELKINS SINN

EQ 10MG BASE/ML

N83363 001

GD SEARLE LLC

EQ 10MG BASE/ML

N86418 001

EQ 20MG BASE/ML

N86418 002

METAXALONE

TABLET; ORAL

SKELAXIN

KING PHARMS

400MG

N13217 001

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 175 (of 274)

METFORMIN HYDROCHLORIDETABLET; ORAL
GLUCOPHAGEBRISTOL MYERS SQUIBB 625MG
750MGN20357 003 Nov 05, 1998
N20357 004 Nov 05, 1998METHACYCLINE HYDROCHLORIDECAPSULE; ORAL
RONDONMYCINMEDPOINTE PHARM HLC EQ 140MG BASE
EQ 280MG BASEN60641 001
N60641 002SYRUP; ORAL
RONDONMYCIN

MEDPOINTE PHARM HLC EQ 70MG BASE/5ML

N60641 003

METHADONE HYDROCHLORIDESYRUP; ORAL
DOLOPHINE HYDROCHLORIDE
ROXANE

10MG/30ML

N06134 004

TABLET; ORAL
METHADONE HYDROCHLORIDE

SANDOZ 5MG

N40241 001 May 29, 1998

TABLET, DISPERSIBLE; ORAL
WESTADONE

SANDOZ 2.5MG

N17108 001

TABLET, EFFERVESCENT; ORAL
WESTADONESANDOZ 5MG
10MG
40MGN17108 002
N17108 003
N17108 004METHAMPHETAMINE HYDROCHLORIDETABLET; ORAL
METHAMPEX

TEVA 10MG

N83889 001

METHAMPHETAMINE HYDROCHLORIDE

ABLE 5MG

N40529 001 Feb 25, 2004

REXAR 5MG

N84931 001

10MG

N84931 002

TEVA 5MG

N86359 001

TABLET, EXTENDED RELEASE; ORAL
DESOXYN

OVATION PHARMS 5MG

N05378 004

10MG

N05378 003

15MG

N05378 005

METHANTHELINE BROMIDETABLET; ORAL
BANTHINE

SHIRE 50MG

N07390 001

METHARBITALTABLET; ORAL
GEMONIL

ABBOTT 100MG

N08322 001

METHAZOLAMIDETABLET; ORAL
METHAZOLAMIDEAPPLIED ANAL 25MG
50MGN40011 001 Jul 17, 1997
N40011 002 Jul 17, 1997

NEPTAZANE

LEDERLE 25MG
50MGN11721 002 Nov 25, 1991
N11721 001

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 176 (of 274)

METHDILAZINE

TABLET, CHEWABLE; ORAL

TACARYL

WESTWOOD SQUIBB 3.6MG

N11950 009

METHDILAZINE HYDROCHLORIDE

SYRUP; ORAL

METHDILAZINE HYDROCHLORIDE

ALPHARMA US PHARMS 4MG/5ML

N87122 001

TACARYL

WESTWOOD SQUIBB 4MG/5ML

N11950 007

TABLET; ORAL

TACARYL

WESTWOOD SQUIBB 8MG

N11950 006

METHICILLIN SODIUM

INJECTABLE; INJECTION

STAPHICILLIN

APOTHECON

EQ 900MG BASE/VIAL

N50117 001

EQ 900MG BASE/VIAL

N61449 001

EQ 3.6GM BASE/VIAL

N50117 002

EQ 3.6GM BASE/VIAL

N61449 002

EQ 5.4GM BASE/VIAL

N50117 003

EQ 5.4GM BASE/VIAL

N61449 003

METHIMAZOLE

TABLET; ORAL

METHIMAZOLE

GENPHARM 20MG

N40350 003 Jun 07, 2001

TAPAZOLE

KING PHARMS 5MG

N07517 002

10MG

N07517 004

METHIXENE HYDROCHLORIDE

TABLET; ORAL

TREST

NOVARTIS 1MG

N13420 001

METHOCARBAMOL

INJECTABLE; INJECTION

METHOCARBAMOL

MARSAM PHARMS LLC 100MG/ML

N89849 001 Dec 27, 1991

TABLET; ORAL

DELAXIN

FERNDAL LABS 500MG

N85454 001

FORBAXIN

FOREST LABS 750MG

N85136 001

METHOCARBAMOL

ABLE 500MG

N40413 001 Mar 17, 2003

750MG

N40413 002 Mar 17, 2003

AM THERAP 500MG

N89417 001 Feb 11, 1987

750MG

N89418 001 Feb 11, 1987

ASCOT 500MG

N87660 001 Oct 27, 1982

750MG

N87661 001 Oct 27, 1982

CLONMEL HLTHCARE 500MG

N85961 001

750MG

N85963 001

HEATHER 500MG

N84675 001

750MG

N84924 001

INWOOD LABS 500MG

N85137 001

IVAX PHARMS 500MG

N84648 001

750MG

N84649 001

KV PHARM 500MG

N85660 001

750MG

N85658 001

MUTUAL PHARM 500MG

N84488 001

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 177 (of 274)

METHOCARBAMOL

TABLET; ORAL

METHOCARBAMOL

MUTUAL PHARM	750MG	N84486	001	
MYLAN	500MG	N84259	001	
	750MG	N84323	001	
NYLOS	750MG	N85033	001	
PHARMERAL	500MG	N84231	002	
	750MG	N84471	001	
PIONEER PHARMS	500MG	N88731	001	Dec 13, 1985
	750MG	N89082	001	Dec 13, 1985
PUREPAC PHARM	500MG	N85718	001	
	750MG	N85718	002	
ROXANE	500MG	N88646	001	Feb 29, 1984
	750MG	N88647	001	Feb 29, 1984
SOLVAY	500MG	N84448	001	
	750MG	N84449	001	
SUPERPHARM	500MG	N87589	001	Jan 22, 1982
	750MG	N87590	001	Jan 22, 1982
UPSHER SMITH	500MG	N87453	001	
	750MG	N87454	001	
WATSON LABS	500MG	N83605	001	
	750MG	N83605	002	

METHOTREXATE SODIUM

INJECTABLE; INJECTION

ABITREXATE

ABIC	EQ 25MG BASE/ML	N89161	001	Mar 10, 1987
	EQ 50MG BASE/VIAL	N89354	001	Jul 17, 1987
	EQ 100MG BASE/VIAL	N89355	001	Jul 17, 1987
	EQ 250MG BASE/VIAL	N89356	001	Jul 17, 1987

FOLEX

PHARMACIA AND UPJOHN	EQ 25MG BASE/VIAL	N87695	001	Apr 08, 1983
	EQ 50MG BASE/VIAL	N87695	002	Apr 08, 1983
	EQ 100MG BASE/VIAL	N87695	003	Apr 08, 1983
	EQ 250MG BASE/VIAL	N88954	001	Oct 24, 1985

FOLEX PFS

PHARMACIA AND UPJOHN	EQ 25MG BASE/ML	N81242	001	Aug 23, 1991
	EQ 25MG BASE/ML	N89180	001	Jan 03, 1986

METHOTREXATE LPF

MAYNE PHARMA USA	EQ 25MG BASE/ML	N11719	007	Mar 31, 1982
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METHOTREXATE PRESERVATIVE FREE

ABRAXIS PHARM	EQ 25MG BASE/ML	N40265	001	Feb 26, 1999
	EQ 1GM BASE/VIAL	N40266	001	Feb 26, 1999
MAYNE PHARMA USA	EQ 20MG BASE/2ML (10 MG/ML)	N11719	014	Apr 13, 2005
	EQ 500MG BASE/20ML (25 MG/ML)	N11719	013	Apr 13, 2005
	EQ 2.5GM BASE/100ML (25 MG/ML)	N11719	011	Apr 13, 2005

METHOTREXATE SODIUM

ABRAXIS PHARM	EQ 2.5MG BASE/ML	N89323	001	Jun 13, 1986
	EQ 20MG BASE/VIAL	N88935	001	Oct 11, 1985
	EQ 25MG BASE/ML	N89263	001	Jun 13, 1986
	EQ 25MG BASE/ML	N89322	001	Jun 13, 1986
	EQ 50MG BASE/VIAL	N88936	001	Oct 11, 1985
	EQ 100MG BASE/VIAL	N88937	001	Oct 11, 1985
MAYNE PHARMA USA	EQ 2.5MG BASE/ML	N11719	004	
	EQ 20MG BASE/VIAL	N11719	001	
	EQ 25MG BASE/ML	N11719	005	
	EQ 50MG BASE/VIAL	N11719	003	
	EQ 100MG BASE/VIAL	N11719	006	
NORBROOK	EQ 25MG BASE/ML	N88648	001	May 09, 1986
PHARMACHEMIE USA	EQ 25MG BASE/ML	N89158	001	Jul 08, 1988
METHOTREXATE SODIUM PRESERVATIVE FREE				
MAYNE PHARMA USA	EQ 1GM BASE/VIAL	N11719	009	Apr 07, 1988

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 178 (of 274)

METHOTREXATE SODIUM

INJECTABLE; INJECTION

MEXATE

BRISTOL

EQ 20MG BASE/VIAL

N86358 001

EQ 50MG BASE/VIAL

N86358 002

EQ 100MG BASE/VIAL

N86358 003

EQ 250MG BASE/VIAL

N86358 004

MEXATE-AQ

BRISTOL MYERS

EQ 25MG BASE/ML

N88760 001

Feb 14, 1985

MEXATE-AQ PRESERVED

BRISTOL MYERS SQUIBB

EQ 25MG BASE/ML

N89887 001

Apr 14, 1989

METHOXAMINE HYDROCHLORIDE

INJECTABLE; INJECTION

VASOXYL

GLAXOSMITHKLINE

10MG/ML

N06772 002

20MG/ML

N06772 001

METHOXSALEN

CAPSULE; ORAL

METHOXSALEN

SANDOZ

10MG

N87781 001

Jun 08, 1982

METHSCOPOLAMINE BROMIDE

TABLET; ORAL

METHSCOPOLAMINE BROMIDE

LEINER

2.5MG

N80970 001

METHYCLOTHIAZIDE

TABLET; ORAL

AQUATENSEN

MEDPOINTE PHARM HLC

5MG

N17364 001

METHYCLOTHIAZIDE

IVAX PHARMS

5MG

N87786 001

May 18, 1982

MYLAN

2.5MG

N87671 001

Aug 17, 1982

PAR PHARM

2.5MG

N89135 001

Feb 12, 1986

5MG

N89136 001

Feb 12, 1986

USL PHARMA

5MG

N88745 001

Mar 21, 1985

WATSON LABS

2.5MG

N85487 001

Mar 11, 1982

2.5MG

N88750 001

Sep 06, 1984

5MG

N85476 001

Mar 11, 1982

METHYCLOTHIAZIDE; PARGYLINE HYDROCHLORIDE

TABLET; ORAL

EUTRON

ABBOTT

5MG;25MG

N16047 001

METHYCLOTHIAZIDE; RESERPINE

TABLET; ORAL

DIUTENSEN-R

MEDPOINTE PHARM HLC

2.5MG;0.1MG

N12708 005

METHYLDOPA

SUSPENSION; ORAL

ALDOMET

MERCK

250MG/5ML

N18389 001

TABLET; ORAL

ALDOMET

MERCK

125MG

N13400 003

250MG

N13400 001

500MG

N13400 002

METHYLDOPA

ACCORD HLTH

125MG

N70070 003

Oct 15, 1985

DISCONTINUED DRUG PRODUCT LIST

6 - 179 (of 274)

METHYLDOPA

TABLET; ORAL
METHYLDOPA

DURAMED PHARMS BARR	250MG	N71006	001	Dec 16, 1986
	500MG	N71009	001	Dec 16, 1986
HALSEY	125MG	N71751	001	Mar 28, 1988
	250MG	N71752	001	Mar 28, 1988
	500MG	N71753	001	Mar 28, 1988
MUTUAL PHARM	125MG	N70073	001	Oct 09, 1986
	250MG	N70060	001	Oct 09, 1986
	500MG	N70074	001	Oct 09, 1986
PARKE DAVIS	125MG	N70331	001	Apr 15, 1986
	250MG	N70332	001	Apr 15, 1986
	500MG	N70333	001	Apr 15, 1986
PUREPAC PHARM	125MG	N70749	001	Feb 07, 1986
	250MG	N70750	001	Feb 07, 1986
	500MG	N70452	001	Feb 07, 1986
ROXANE	125MG	N70192	001	Apr 25, 1986
	250MG	N70193	001	Apr 25, 1986
	500MG	N70194	001	Apr 25, 1986
SUPERPHARM	250MG	N70669	001	Jun 23, 1989
	500MG	N70670	001	Jun 23, 1989
TEVA	125MG	N71105	001	Dec 05, 1986
	250MG	N71106	001	Dec 05, 1986
	500MG	N71067	001	Dec 05, 1986
WATSON LABS	125MG	N70245	001	Feb 25, 1986
	125MG	N70260	001	Jun 24, 1985
	250MG	N70246	001	Feb 25, 1986
	250MG	N70261	001	Jun 24, 1985
	500MG	N70247	001	Feb 25, 1986
	500MG	N70262	001	Jun 24, 1985

METHYLDOPATE HYDROCHLORIDE

INJECTABLE; INJECTION
ALDOMET

MERCK	50MG/ML	N13401	001	
METHYLDOPATE HYDROCHLORIDE				
ABRAXIS PHARM	50MG/ML	N70652	001	Jun 03, 1986
BAXTER HLTHCARE	50MG/ML	N70291	001	Jul 01, 1986
MARSAM PHARMS LLC	50MG/ML	N71812	001	Dec 22, 1987
MAYNE PHARMA USA	50MG/ML	N70691	001	Jun 19, 1987
	50MG/ML	N70849	001	Jun 19, 1987
SMITH AND NEPHEW	50MG/ML	N70841	001	Jan 02, 1987

METHYLPHENIDATE HYDROCHLORIDE

TABLET; ORAL

METHYLPHENIDATE HYDROCHLORIDE

ABLE	5MG	N40404	001	Mar 29, 2001
	10MG	N40404	002	Mar 29, 2001
	20MG	N40404	003	Mar 29, 2001

TABLET, EXTENDED RELEASE; ORAL

METHYLPHENIDATE HYDROCHLORIDE

ABLE	20MG	N76032	001	May 09, 2001
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METHYLPREDNISOLONE

TABLET; ORAL

MEDROL

PHARMACIA AND UPJOHN	2MG	N11153	002	
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METHYLPREDNISOLONE

HEATHER	4MG	N85650	001	
SANDOZ	4MG	N87341	001	
WATSON LABS	4MG	N86161	001	Feb 09, 1982
	16MG	N86159	001	Feb 09, 1982

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 180 (of 274)

METHYLPREDNISOLONE ACETATE

ENEMA; RECTAL

MEDROL

PHARMACIA AND UPJOHN 40MG/BOT

N18102 001

INJECTABLE; INJECTION

METHYLPREDNISOLONE ACETATE

AKORN

40MG/ML

N86903 001

Oct 20, 1982

80MG/ML

N86903 002

Oct 20, 1982

WATSON LABS

20MG/ML

N85597 001

20MG/ML

N87248 001

40MG/ML

N85374 001

40MG/ML

N85600 001

80MG/ML

N85595 001

80MG/ML

N86507 001

M-PREDROL

BEL MAR

40MG/ML

N86666 001

80MG/ML

N87135 001

OINTMENT; TOPICAL

MEDROL ACETATE

PHARMACIA AND UPJOHN 0.25%

N12421 001

1%

N12421 002

METHYLPREDNISOLONE ACETATE; NEOMYCIN SULFATE

CREAM; TOPICAL

NEO-MEDROL ACETATE

PHARMACIA AND UPJOHN 0.25%;EQ 3.5MG BASE/GM

N60611 002

1%;EQ 3.5MG BASE/GM

N60611 001

METHYLPREDNISOLONE SODIUM SUCCINATE

INJECTABLE; INJECTION

A-METHAPRED

ABBOTT

EQ 40MG BASE/VIAL

N89573 001

Feb 22, 1991

EQ 125MG BASE/VIAL

N89574 001

Feb 22, 1991

EQ 500MG BASE/VIAL

N89575 001

Feb 22, 1991

EQ 1GM BASE/VIAL

N89576 001

Feb 22, 1991

METHYLPREDNISOLONE

ELKINS SINN

EQ 125MG BASE/VIAL

N86906 002

EQ 500MG BASE/VIAL

N86906 003

EQ 1GM BASE/VIAL

N86906 004

ORGANON USA INC

EQ 500MG BASE/VIAL

N87535 001

Jun 25, 1982

EQ 1GM BASE/VIAL

N87535 002

Jun 25, 1982

METHYLPREDNISOLONE SODIUM SUCCINATE

ABRAXIS PHARM

EQ 40MG BASE/VIAL

N88676 001

Jun 08, 1984

EQ 40MG BASE/VIAL

N89143 001

Mar 28, 1986

EQ 125MG BASE/VIAL

N88677 001

Jun 08, 1984

EQ 125MG BASE/VIAL

N89144 001

Mar 28, 1986

EQ 500MG BASE/VIAL

N88678 001

Jun 08, 1984

EQ 500MG BASE/VIAL

N89186 001

Mar 28, 1986

EQ 500MG BASE/VIAL

N89187 001

Mar 28, 1986

EQ 1GM BASE/VIAL

N88679 001

Jun 08, 1984

EQ 1GM BASE/VIAL

N89188 001

Mar 28, 1986

EQ 1GM BASE/VIAL

N89189 001

Mar 28, 1986

ELKINS SINN

EQ 40MG BASE/VIAL

N86906 001

INTL MEDICATION

EQ 40MG BASE/VIAL

N87812 001

Feb 09, 1983

EQ 125MG BASE/VIAL

N87813 001

Feb 09, 1983

EQ 500MG BASE/VIAL

N87851 001

Feb 09, 1983

EQ 1GM BASE/VIAL

N87852 001

Feb 09, 1983

SICOR PHARMS

EQ 500MG BASE/VIAL

N81267 001

Nov 30, 1992

EQ 1GM BASE/VIAL

N81268 001

Nov 30, 1992

WATSON LABS

EQ 40MG BASE/VIAL

N86953 001

Jul 22, 1982

EQ 125MG BASE/VIAL

N87030 001

Jul 22, 1982

EQ 500MG BASE/VIAL

N88523 001

Jul 24, 1984

EQ 1GM BASE/VIAL

N88524 001

Jul 24, 1984

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 181 (of 274)

METHYLPREDNISOLONE; NEOMYCIN SULFATEOINTMENT; OPHTHALMIC
NEO-MEDROL

PHARMACIA AND UPJOHN 0.1%;EQ 3.5MG BASE/GM

N60645 001

METHYLTESTOSTERONECAPSULE; ORAL
METHYLTESTOSTERONE

HEATHER 10MG

N84967 001

TABLET; BUCCAL
ANDROID 5

VALEANT PHARM INTL 5MG

N87222 001

ORETON

SCHERING 10MG

N80281 001

TABLET; BUCCAL/SUBLINGUAL
METANDREN

NOVARTIS 5MG

N03240 004

10MG

N03240 005

METHYLTESTOSTERONE

IMPAX LABS 10MG

N84287 001

LEINER 5MG

N83836 001

LILLY 10MG

N80256 001

PUREPAC PHARM 10MG

N80308 001

10MG

N80475 001

TABLICAPS 10MG

N85125 001

USL PHARMA 10MG

N80271 001

TABLET; ORAL

METANDREN

NOVARTIS 10MG

N03240 001

25MG

N03240 003

METHYLTESTOSTERONE

INWOOD LABS 10MG

N80839 001

25MG

N80973 001

KV PHARM 10MG

N84312 001

LANNETT 10MG

N87092 001

25MG

N87111 001

Nov 05, 1982

Jan 27, 1983

LEINER 5MG

N80214 001

10MG

N80214 002

25MG

N80214 003

LILLY 25MG

N80256 002

PARKE DAVIS 10MG

N84244 001

25MG

N84241 001

PUREPAC PHARM 10MG

N80309 001

10MG

N80475 002

25MG

N80310 001

25MG

N80475 003

TABLICAPS 10MG

N80313 001

25MG

N85270 001

WATSON LABS 10MG

N80933 001

25MG

N80931 001

WEST WARD 10MG

N84331 001

25MG

N84331 002

25MG

N84642 001

ORETON METHYL

SCHERING 10MG

N03158 001

25MG

N03158 002

METHYPRYLONCAPSULE; ORAL
NOLUDAR

ROCHE 300MG

N09660 008

ELIXIR; ORAL

NOLUDAR

ROCHE 50MG/5ML

N09660 007

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 182 (of 274)

METHYPRYLON

TABLET; ORAL
NOLUDAR
ROCHE

50MG
200MG

N09660 002
N09660 004

METHYSERGIDE MALEATE

TABLET; ORAL
SANSERT
NOVARTIS

2MG

N12516 001

METOCLOPRAMIDE HYDROCHLORIDE

CONCENTRATE; ORAL

METOCLOPRAMIDE INTENSOL

ROXANE

EQ 10MG BASE/ML

N72995 001 Jan 30, 1992

INJECTABLE; INJECTION

METOCLOPRAMIDE HYDROCHLORIDE

ABRAXIS PHARM

EQ 10MG BASE/2ML

N70293 001 Jan 24, 1986

BEDFORD

EQ 5MG BASE/ML

N72155 001 Mar 30, 1992

EQ 5MG BASE/ML

N72244 001 Mar 30, 1992

EQ 5MG BASE/ML

N72247 001 May 18, 1992

HOSPIRA

EQ 5MG BASE/ML

N70506 001 Jun 22, 1989

MAYNE PHARMA USA

EQ 5MG BASE/ML

N71990 001 Jan 18, 1989

NORBROOK

EQ 10MG BASE/2ML

N70892 001 Aug 26, 1988

SMITH AND NEPHEW

EQ 5MG BASE/ML

N70623 001 Mar 02, 1987

EQ 10MG BASE/2ML

N70622 001 Mar 02, 1987

REGLAN

BAXTER HLTHCARE CORP

EQ 10MG BASE/ML

N17862 004 May 28, 1987

SOLUTION; ORAL

METOCLOPRAMIDE

VISTAPHARM

EQ 5MG BASE/5ML

N75051 001 Jan 26, 2001

METOCLOPRAMIDE HYDROCHLORIDE

MORTON GROVE

EQ 5MG BASE/5ML

N70949 001 Mar 06, 1987

PACO

EQ 5MG BASE/5ML

N71665 001 Dec 05, 1988

ROXANE

EQ 5MG BASE/5ML

N72038 001 Dec 05, 1988

TEVA

EQ 5MG BASE/5ML

N70819 001 Jul 10, 1987

EQ 5MG BASE/5ML

N71315 001 Jun 30, 1993

REGLAN

ROBINS AH

EQ 5MG BASE/5ML

N18821 001 Mar 25, 1983

TABLET; ORAL

CLOPRA

QUANTUM PHARMICS

EQ 5MG BASE

N72384 001 Jun 02, 1988

EQ 10MG BASE

N70294 001 Jul 29, 1985

CLOPRA-"YELLOW"

QUANTUM PHARMICS

EQ 10MG BASE

N70632 001 Oct 28, 1985

MAXOLON

KING PHARMS

EQ 10MG BASE

N70106 001 Mar 04, 1986

METOCLOPRAMIDE HYDROCHLORIDE

CLONMEL HLTHCARE

EQ 10MG BASE

N72639 001 May 09, 1991

HALSEY

EQ 10MG BASE

N70906 001 Oct 28, 1986

INTERPHARM

EQ 10MG BASE

N71213 001 Sep 24, 1986

MUTUAL PHARM

EQ 10MG BASE

N70660 001 Feb 10, 1987

PAR PHARM

EQ 10MG BASE

N70342 001 Mar 25, 1986

SANDOZ

EQ 5MG BASE

N72436 001 Jun 22, 1989

EQ 10MG BASE

N70850 001 Feb 03, 1987

SCHERING

EQ 10MG BASE

N70598 001 Feb 02, 1987

SUPERPHARM

EQ 10MG BASE

N70926 001 Jun 26, 1987

USL PHARMA

EQ 10MG BASE

N70339 001 Jul 29, 1985

WATSON LABS

EQ 10MG BASE

N70363 001 Mar 02, 1987

EQ 10MG BASE

N70453 001 Jun 06, 1986

EQ 10MG BASE

N70645 001 May 11, 1987

TABLET, ORALLY DISINTEGRATING; ORAL

REGLAN ODT

SCHWARZ PHARMA

EQ 5MG BASE

N21793 001 Jun 10, 2005

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 183 (of 274)

METOCLOPRAMIDE HYDROCHLORIDETABLET, ORALLY DISINTEGRATING; ORAL
REGLAN ODT

SCHWARZ PHARMA	EQ 10MG BASE	N21793	002	Jun 10, 2005
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METOCURINE IODIDEINJECTABLE; INJECTION
METUBINE IODIDE

LILLY	2MG/ML	N06632	003	
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METOLAZONETABLET; ORAL
DIULO

GD SEARLE LLC	2.5MG	N18535	001	
	5MG	N18535	002	
	10MG	N18535	003	

MYKROX

UCB INC	0.5MG	N19532	001	Oct 30, 1987
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METOPROLOL FUMARATETABLET, EXTENDED RELEASE; ORAL
LOPRESSOR

NOVARTIS	EQ 100MG TARTRATE	N19786	001	Dec 27, 1989
	EQ 200MG TARTRATE	N19786	002	Dec 27, 1989
	EQ 300MG TARTRATE	N19786	003	Dec 27, 1989
	EQ 400MG TARTRATE	N19786	004	Dec 27, 1989

METOPROLOL TARTRATETABLET; ORAL
METOPROLOL TARTRATE

APOTHECON	50MG	N74258	001	Jan 27, 1994
	100MG	N74258	002	Jan 27, 1994
PUREPAC PHARM	50MG	N74380	001	Jul 29, 1994
	100MG	N74380	002	Jul 29, 1994

METRIZAMIDEINJECTABLE; INJECTION
AMIPAQUE

GE HEALTHCARE	2.5GM/VIAL	N17982	003	Sep 12, 1983
	3.75GM/VIAL	N17982	001	
	6.75GM/VIAL	N17982	002	
	13.5GM/VIAL	N17982	004	Sep 12, 1983

METRONIDAZOLECAPSULE; ORAL
METRONIDAZOLE

ABLE	375MG	N76505	001	Nov 13, 2003
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INJECTABLE; INJECTION
METRO I.V.

B BRAUN	500MG/100ML	N18674	001	Aug 31, 1982
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METRONIDAZOLE

ABBOTT	500MG/100ML	N18889	001	Nov 18, 1983
ABRAXIS PHARM	500MG/100ML	N70071	001	Dec 03, 1984
ELKINS SINN	500MG/100ML	N18907	001	Mar 30, 1984
INTL MEDICATION	500MG/100ML	N70004	001	May 08, 1985
WATSON LABS	500MG/100ML	N70042	001	Dec 20, 1984
	500MG/100ML	N70170	001	Apr 01, 1986

TABLET; ORAL
METROMIDOL

LABS AF	250MG	N74523	001	Oct 24, 1996
	500MG	N74523	002	Oct 24, 1996

METRONIDAZOLE

ABLE	250MG	N76519	001	Jun 27, 2003
	500MG	N76519	002	Jun 27, 2003

DISCONTINUED DRUG PRODUCT LIST

6 - 184 (of 274)

METRONIDAZOLE

TABLET; ORAL
METRONIDAZOLE

HALSEY	250MG	N70021	001	Apr 02, 1985
	500MG	N70593	001	Feb 27, 1986
MUTUAL PHARM	250MG	N18818	001	Feb 16, 1983
	500MG	N18818	002	Feb 16, 1983
SUPERPHARM	250MG	N70008	001	Dec 11, 1984
	500MG	N70009	001	Dec 11, 1984
WATSON LABS	250MG	N18599	001	Sep 17, 1982
	500MG	N18599	002	Feb 13, 1984
PROTOSTAT				
ORTHO MCNEIL PHARM	250MG	N18871	001	Mar 02, 1983
	500MG	N18871	002	Mar 02, 1983
SATRIC				
SAVAGE LABS	250MG	N70029	001	Mar 19, 1985
	500MG	N70731	001	Jun 08, 1987
TABLET, EXTENDED RELEASE; ORAL METRONIDAZOLE				
ABLE	750MG	N76462	001	Jun 25, 2003

METRONIDAZOLE HYDROCHLORIDE

INJECTABLE; INJECTION
METRONIDAZOLE HYDROCHLORIDE

ABRAXIS PHARM	EQ 500MG BASE/VIAL	N70295	001	Oct 15, 1985
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METYRAPONE

TABLET; ORAL
METOPIRONE

NOVARTIS	250MG	N12911	001	
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MEZLOCILLIN SODIUM MONOHYDRATE

INJECTABLE; INJECTION
MEZLIN

BAYER PHARMS	EQ 1GM BASE/VIAL	N50549	001	
	EQ 1GM BASE/VIAL	N62333	001	
	EQ 1GM BASE/VIAL	N62372	005	Jan 13, 1983
	EQ 2GM BASE/VIAL	N50549	002	
	EQ 2GM BASE/VIAL	N62333	002	
	EQ 2GM BASE/VIAL	N62372	001	May 13, 1982
	EQ 3GM BASE/VIAL	N50549	003	
	EQ 3GM BASE/VIAL	N62333	003	
	EQ 3GM BASE/VIAL	N62372	002	May 13, 1982
	EQ 3GM BASE/VIAL	N62697	001	Jan 22, 1987
	EQ 4GM BASE/VIAL	N50549	004	
	EQ 4GM BASE/VIAL	N62333	004	
	EQ 4GM BASE/VIAL	N62372	003	May 13, 1982
	EQ 4GM BASE/VIAL	N62697	002	Jan 22, 1987
	EQ 20GM BASE/VIAL	N50549	005	Mar 02, 1988
	EQ 20GM BASE/VIAL	N62372	004	Mar 02, 1988

MICONAZOLE

INJECTABLE; INJECTION
MONISTAT

JANSSEN PHARMA	10MG/ML	N18040	001	
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MICONAZOLE NITRATE

LOTION; TOPICAL
MONISTAT-DERM

JOHNSON AND JOHNSON	2%	N17739	001	
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TAMPON; VAGINAL
MONISTAT 5

PERSONAL PRODS	100MG	N18592	001	Oct 27, 1989
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27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 185 (of 274)

MIDAZOLAM HYDROCHLORIDEINJECTABLE; INJECTION
MIDAZOLAM HYDROCHLORIDE

APOTHECON

EQ 1MG BASE/ML

N75620 001 Nov 01, 2000

EQ 5MG BASE/ML

N75620 002 Nov 01, 2000

EQ 5MG BASE/ML

N75641 001 Oct 19, 2000

ASTRAZENECA

EQ 5MG BASE/ML

N75263 001 Jun 26, 2000

BEDFORD

EQ 5MG BASE/ML

N75249 001 Jun 23, 2000

BEN VENUE

EQ 5MG BASE/ML

N75455 001 Jun 20, 2000

VERSED

HLR

EQ 1MG BASE/ML

N18654 002 May 26, 1987

EQ 5MG BASE/ML

N18654 001 Dec 20, 1985

SYRUP; ORAL

VERSED

ROCHE

EQ 2MG BASE/ML

N20942 001 Oct 15, 1998

MILRINONE LACTATE

INJECTABLE; INJECTION

PRIMACOR IN DEXTROSE 5% IN PLASTIC CONTAINER

SANOFI AVENTIS US

EQ 10MG BASE/100ML

N20343 001 Aug 09, 1994

EQ 15MG BASE/100ML

N20343 002 Aug 09, 1994

MINOCYCLINE HYDROCHLORIDE

CAPSULE; ORAL

DYNACIN

MEDICIS

EQ 50MG BASE

N63066 001 Aug 14, 1990

MINOCIN

TRIAx PHARMS

EQ 50MG BASE

N50315 002

EQ 75MG BASE

N50649 003 Feb 12, 2001

EQ 100MG BASE

N50315 001

INJECTABLE; INJECTION

MINOCIN

LEDERLE

EQ 100MG BASE/VIAL

N62139 001

TRIAx PHARMS

EQ 100MG BASE/VIAL

N50444 001

SUSPENSION; ORAL

MINOCIN

TRIAx PHARMS

EQ 50MG BASE/5ML

N50445 001

TABLET; ORAL

MINOCYCLINE HYDROCHLORIDE

TRIAx PHARMS

EQ 50MG BASE

N50451 003 Aug 10, 1982

EQ 100MG BASE

N50451 002 Aug 10, 1982

MINOXIDIL

TABLET; ORAL

MINODYL

QUANTUM PHARMICS

2.5MG

N72153 001 Jul 13, 1988

10MG

N71534 001 Mar 19, 1987

MINOXIDIL

ROYCE LABS

2.5MG

N71799 001 Nov 10, 1987

10MG

N71796 001 Nov 10, 1987

USL PHARMA

2.5MG

N71537 001 Dec 16, 1988

MITOMYCIN

INJECTABLE; INJECTION

MITOMYCIN

MAYNE PHARMA USA

20MG/VIAL

N64106 001 Nov 29, 1995

MUTAMYCIN

BRISTOL

5MG/VIAL

N50450 001

20MG/VIAL

N50450 002

MIVACURIUM CHLORIDE

INJECTABLE; INJECTION

MIVACRON

ABBOTT

EQ 2MG BASE/ML

N20098 001 Jan 22, 1992

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 186 (of 274)

MIVACURIUM CHLORIDE

INJECTABLE; INJECTION

MIVACRON IN DEXTROSE 5% IN PLASTIC CONTAINER

ABBOTT

EQ 0.5MG BASE/ML

N20098 002 Jan 22, 1992

EQ 50MG BASE/100ML

N20098 003 Jan 22, 1992

MOLINDONE HYDROCHLORIDE

CAPSULE; ORAL

MOBAN

ENDO PHARMS

5MG

N17111 001

10MG

N17111 002

25MG

N17111 003

CONCENTRATE; ORAL

MOBAN

ENDO PHARMS

20MG/ML

N17938 001

TABLET; ORAL

MOBAN

ENDO PHARMS

100MG

N17111 008

MONOCTANOIN

LIQUID; PERFUSION, BILIARY

MOCTANIN

ETHITEK

100%

N19368 001 Oct 29, 1985

MORPHINE SULFATE

TABLET, EXTENDED RELEASE; ORAL

MORPHINE SULFATE

CLONMEL HLTHCARE

15MG

N75407 001 Jan 28, 2000

MOXALACTAM DISODIUM

INJECTABLE; INJECTION

MOXAM

LILLY

EQ 250MG BASE/VIAL

N50550 001

EQ 500MG BASE/VIAL

N50550 002

EQ 1GM BASE/VIAL

N50550 003

EQ 2GM BASE/VIAL

N50550 004

EQ 10GM BASE/VIAL

N50550 008

NABUMETONE

TABLET; ORAL

NABUMETONE

COPLEY PHARM

750MG

N75179 001 Jun 06, 2000

RELAFEN

SMITHKLINE BEECHAM

500MG

N19583 001 Dec 24, 1991

750MG

N19583 002 Dec 24, 1991

NAFCILLIN SODIUM

CAPSULE; ORAL

UNIPEN

WYETH AYERST

EQ 250MG BASE

N50111 001

FOR SOLUTION; ORAL

UNIPEN

WYETH AYERST

EQ 250MG BASE/5ML

N50199 001

INJECTABLE; INJECTION

NAFCILLIN SODIUM

APOTHECON

EQ 500MG BASE/VIAL

N61984 001

EQ 1GM BASE/VIAL

N61984 002

EQ 2GM BASE/VIAL

N61984 003

EQ 4GM BASE/VIAL

N61984 005

MARSAM PHARMS LLC

EQ 500MG BASE/VIAL

N62844 001 Oct 26, 1988

EQ 1GM BASE/VIAL

N62844 002 Oct 26, 1988

EQ 1.5GM BASE/VIAL

N62844 003 Oct 26, 1988

EQ 2GM BASE/VIAL

N62844 004 Oct 26, 1988

EQ 4GM BASE/VIAL

N62844 005 Oct 26, 1988

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 187 (of 274)

NAFCILLIN SODIUM

INJECTABLE; INJECTION

NAFCILLIN SODIUM

MARSAM PHARMS LLC

EQ 10GM BASE/VIAL

N63008 001

Sep 29, 1988

NALLPEN

GLAXOSMITHKLINE

EQ 500MG BASE/VIAL

N61999 001

EQ 1GM BASE/VIAL

N61999 002

EQ 1GM BASE/VIAL

N62755 001

Dec 19, 1986

EQ 2GM BASE/VIAL

N61999 003

EQ 2GM BASE/VIAL

N62755 002

Dec 19, 1986

EQ 10GM BASE/VIAL

N61999 004

UNIPEN

WYETH AYERST

EQ 500MG BASE/VIAL

N50320 001

EQ 500MG BASE/VIAL

N62717 001

Dec 16, 1986

EQ 1GM BASE/VIAL

N62717 002

Dec 16, 1986

EQ 2GM BASE/VIAL

N50320 003

EQ 2GM BASE/VIAL

N62717 004

Dec 16, 1986

EQ 4GM BASE/VIAL

N50320 004

EQ 10GM BASE/VIAL

N50320 005

EQ 20GM BASE/VIAL

N50320 006

UNIPEN IN PLASTIC CONTAINER

WYETH AYERST

EQ 1GM BASE/VIAL

N50320 002

TABLET; ORAL

UNIPEN

WYETH AYERST

EQ 500MG BASE

N50462 001

NALBUPHINE HYDROCHLORIDE

INJECTABLE; INJECTION

NALBUPHINE

ABRAXIS PHARM

10MG/ML

N70751 001

Jul 02, 1986

20MG/ML

N70752 001

Sep 24, 1986

NALBUPHINE HYDROCHLORIDE

ABBOTT

1.5MG/ML

N20200 001

Mar 12, 1993

20MG/ML

N70917 001

Feb 03, 1989

ASTRAZENECA

10MG/ML

N72070 001

Apr 10, 1989

10MG/ML

N72071 001

Apr 10, 1989

10MG/ML

N72072 001

Apr 10, 1989

20MG/ML

N72073 001

Apr 10, 1989

20MG/ML

N72074 001

Apr 10, 1989

20MG/ML

N72075 001

Apr 10, 1989

NALIDIXIC ACID

SUSPENSION; ORAL

NEGGRAM

SANOFI AVENTIS US

250MG/5ML

N17430 001

TABLET; ORAL

NALIDIXIC ACID

MUTUAL PHARM

250MG

N70270 001

Jun 29, 1988

500MG

N70271 001

Jun 29, 1988

1GM

N70272 001

Jun 29, 1988

NALOXONE HYDROCHLORIDE

INJECTABLE; INJECTION

NALOXONE

BAXTER HLTHCARE

0.4MG/ML

N70298 001

Sep 24, 1986

0.4MG/ML

N70299 001

Sep 24, 1986

0.4MG/ML

N70496 001

Sep 24, 1986

WYETH AYERST

0.02MG/ML

N70188 001

Sep 24, 1986

0.02MG/ML

N70189 001

Sep 24, 1986

0.4MG/ML

N70190 001

Sep 24, 1986

0.4MG/ML

N70191 001

Sep 24, 1986

NALOXONE HYDROCHLORIDE

ABRAXIS PHARM

0.02MG/ML

N70648 001

Nov 17, 1986

0.02MG/ML

N70661 001

Nov 17, 1986

DISCONTINUED DRUG PRODUCT LIST

6 - 188 (of 274)

NALOXONE HYDROCHLORIDE

INJECTABLE; INJECTION
NALOXONE HYDROCHLORIDE

ABRAXIS PHARM	0.4MG/ML	N70649	001	Nov 17, 1986
	1MG/ML	N71604	001	Dec 16, 1988
ASTRAZENECA	0.02MG/ML	N72081	001	Apr 11, 1989
	0.02MG/ML	N72082	001	Apr 11, 1989
	0.02MG/ML	N72083	001	Apr 11, 1989
	0.02MG/ML	N72084	001	Apr 11, 1989
	0.02MG/ML	N72085	001	Apr 11, 1989
	0.4MG/ML	N72086	001	Apr 11, 1989
	0.4MG/ML	N72087	001	Apr 11, 1989
	0.4MG/ML	N72088	001	Apr 11, 1989
	0.4MG/ML	N72089	001	Apr 11, 1989
	0.4MG/ML	N72090	001	Apr 11, 1989
	1MG/ML	N72091	001	Apr 11, 1989
	1MG/ML	N72092	001	Apr 11, 1989
	1MG/ML	N72093	001	Apr 11, 1989
BAXTER HLTHCARE	0.02MG/ML	N71272	001	May 24, 1988
	1MG/ML	N71273	001	May 24, 1988
	1MG/ML	N71274	001	May 24, 1988
	1MG/ML	N71287	001	May 24, 1988
INTL MEDICATION	0.4MG/ML	N70417	001	Sep 24, 1986
	1MG/ML	N72115	001	Apr 27, 1988
MARSAM PHARMS LLC	0.4MG/ML	N71811	001	Jul 19, 1988
SMITH AND NEPHEW	0.02MG/ML	N71671	001	Nov 17, 1987
	0.4MG/ML	N71681	001	Nov 17, 1987
	0.4MG/ML	N71682	001	Nov 17, 1987
SOLOPAK	0.02MG/ML	N71672	001	Nov 17, 1987
	0.4MG/ML	N71683	001	Nov 17, 1987
WATSON LABS	0.4MG/ML	N71339	001	Nov 18, 1987
NARCAN				
BRISTOL MYERS SQUIBB	0.4MG/ML	N71083	001	Jul 28, 1988
	1MG/ML	N71084	001	Jul 28, 1988
	1MG/ML	N71311	001	Jul 28, 1988

NANDROLONE DECANOATE

INJECTABLE; INJECTION
DECA-DURABOLIN

ORGANON USA INC	50MG/ML	N13132	001	Jun 12, 1986
	100MG/ML	N13132	002	Jun 12, 1986
	200MG/ML	N13132	003	Jun 12, 1986
NANDROLONE DECANOATE				
ABRAXIS PHARM	100MG/ML	N88290	001	Oct 03, 1983
	200MG/ML	N88317	001	Oct 14, 1983
AKORN	100MG/ML	N87519	001	Sep 28, 1983
WATSON LABS	50MG/ML	N87598	001	Oct 06, 1983
	50MG/ML	N88554	001	Feb 10, 1986
	100MG/ML	N87599	001	Oct 06, 1983

NANDROLONE PHENPROPIONATE

INJECTABLE; INJECTION
DURABOLIN

ORGANON USA INC	25MG/ML	N11891	001	
	50MG/ML	N11891	002	
NANDROLONE PHENPROPIONATE				
WATSON LABS	25MG/ML	N86386	001	Jun 17, 1983
	50MG/ML	N87488	001	Jun 17, 1983

NAPHAZOLINE HYDROCHLORIDE

SOLUTION/DROPS; OPHTHALMIC
NAFAZAIR

PHARMAFAIR	0.1%	N88101	001	Apr 15, 1983
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DISCONTINUED DRUG PRODUCT LIST

6 - 189 (of 274)

NAPHAZOLINE HYDROCHLORIDE

SOLUTION/DROPS; OPHTHALMIC
NAPHCN FORTE

ALCON	0.1%	N80229	001	
OPCON				
BAUSCH AND LOMB	0.1%	N87506	001	

NAPROXEN

TABLET; ORAL
NAPROXEN

BAXTER HLTHCARE	250MG	N74105	001	Dec 21, 1993
	375MG	N74105	002	Dec 21, 1993
	500MG	N74105	003	Dec 21, 1993
HAMILTON PHARMS	250MG	N74110	001	Oct 30, 1992
	375MG	N74110	002	Oct 30, 1992
	500MG	N74110	003	Oct 30, 1992
IVAX PHARMS	250MG	N74111	001	Feb 28, 1995
	375MG	N74111	002	Feb 28, 1995
	500MG	N74111	003	Feb 28, 1995
PUREPAC PHARM	250MG	N74263	001	Dec 21, 1993
	375MG	N74263	002	Dec 21, 1993
	500MG	N74263	003	Dec 21, 1993
ROXANE	250MG	N74211	001	Feb 28, 1994
	375MG	N74211	002	Feb 28, 1994
	500MG	N74211	003	Feb 28, 1994

NAPROXEN SODIUM

TABLET; ORAL
NAPROXEN SODIUM

ABLE	EQ 250MG BASE	N76544	001	Aug 22, 2003
	EQ 500MG BASE	N76544	002	Aug 22, 2003
HAMILTON PHARMS	EQ 250MG BASE	N74106	001	Aug 31, 1993
	EQ 500MG BASE	N74106	002	Aug 31, 1993
IVAX PHARMS	EQ 250MG BASE	N74230	001	Mar 14, 1995
	EQ 500MG BASE	N74230	002	Mar 14, 1995
PUREPAC PHARM	EQ 250MG BASE	N74319	001	Mar 20, 1995
	EQ 500MG BASE	N74319	002	Mar 20, 1995
ROXANE	EQ 250MG BASE	N74257	001	Dec 21, 1993
	EQ 500MG BASE	N74257	002	Dec 21, 1993

TABLET, EXTENDED RELEASE; ORAL
NAPRELAN

STAT TRADE	EQ 750MG BASE	N20353	003	Jan 05, 1996
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NEDOCROMIL SODIUM

SOLUTION; INHALATION
TILADE

SANOFI AVENTIS US	0.5%	N20750	001	Oct 01, 1997
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NEFAZODONE HYDROCHLORIDE

TABLET; ORAL
NEFAZODONE HYDROCHLORIDE

ROXANE	50MG	N76196	001	Sep 16, 2003
	100MG	N76196	002	Sep 16, 2003
	150MG	N76196	003	Sep 16, 2003
	200MG	N76196	004	Sep 16, 2003
	250MG	N76196	005	Sep 16, 2003

SERZONE

BRISTOL MYERS SQUIBB	50MG	N20152	001	Dec 22, 1994
	100MG	N20152	002	Dec 22, 1994
	150MG	N20152	003	Dec 22, 1994
	200MG	N20152	004	Dec 22, 1994
	250MG	N20152	005	Dec 22, 1994
	300MG	N20152	006	Dec 22, 1994

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 190 (of 274)

NEOMYCIN SULFATE

SOLUTION; ORAL			
MYCIFRADIN			
PHARMACIA AND UPJOHN	EQ 87.5MG BASE/5ML	N50285	001
TABLET; ORAL			
MYCIFRADIN			
PHARMACIA AND UPJOHN	EQ 350MG BASE	N60520	001
NEOBIOTIC			
PFIZER	EQ 350MG BASE	N60475	001
NEOMYCIN SULFATE			
BRISTOL MYERS SQUIBB	500MG	N60365	001
LANNETT	500MG	N60607	001
LILLY	500MG	N60385	001
ROXANE	500MG	N62173	001
SANDOZ	500MG	N61586	001

NEOMYCIN SULFATE; POLYMYXIN B SULFATE

CREAM; TOPICAL			
NEOSPORIN			
GLAXOSMITHKLINE	EQ 3.5MG BASE/GM;10,000 UNITS/GM	N50176	002 Jan 14, 1985
OINTMENT; OPHTHALMIC			
STATROL			
ALCON	EQ 3.5MG BASE/GM;10,000 UNITS/GM	N50344	002
SOLUTION/DROPS; OPHTHALMIC			
STATROL			
ALCON	EQ 3.5MG BASE/ML;16,250 UNITS/ML	N50456	001
	EQ 3.5MG BASE/ML;16,250 UNITS/ML	N62339	001 Nov 30, 1984

NEOMYCIN SULFATE; PREDNISOLONE ACETATE

OINTMENT; OPHTHALMIC			
NEO-DELTA-CORTEF			
PHARMACIA AND UPJOHN	EQ 3.5MG BASE/GM;0.25%	N61039	002
	EQ 3.5MG BASE/GM;0.5%	N61039	001
SUSPENSION/DROPS; OPHTHALMIC			
NEO-DELTA-CORTEF			
PHARMACIA AND UPJOHN	EQ 3.5MG BASE/ML;0.25%	N61037	001

NEOMYCIN SULFATE; PREDNISOLONE SODIUM PHOSPHATE

OINTMENT; OPHTHALMIC			
NEO-HYDELTRASOL			
MERCK	EQ 3.5MG BASE/GM;EQ 0.25% PHOSPHATE	N50378	001

NEOMYCIN SULFATE; TRIAMCINOLONE ACETONIDE

CREAM; TOPICAL			
MYTREX A			
SAVAGE LABS	EQ 3.5MG BASE/GM;0.1%	N62598	001 Jul 21, 1986
NEOMYCIN SULFATE-TRIAMCINOLONE ACETONIDE			
FOUGERA	EQ 3.5MG BASE/GM;0.1%	N62600	001 Jul 21, 1986
PHARMADERM	EQ 3.5MG BASE/GM;0.1%	N62595	001 Jul 21, 1986
OINTMENT; TOPICAL			
MYTREX A			
SAVAGE LABS	EQ 3.5MG BASE/GM;0.1%	N62609	001 May 23, 1986
NEOMYCIN SULFATE-TRIAMCINOLONE ACETONIDE			
FOUGERA	EQ 3.5MG BASE/GM;0.1%	N62608	001 May 23, 1986
PHARMADERM	EQ 3.5MG BASE/GM;0.1%	N62607	001 May 23, 1986

NETILMICIN SULFATE

INJECTABLE; INJECTION			
NETROMYCIN			
SCHERING	EQ 10MG BASE/ML	N50544	001 Feb 28, 1983
	EQ 25MG BASE/ML	N50544	002 Feb 28, 1983
	EQ 100MG BASE/ML	N50544	003 Feb 28, 1983

DISCONTINUED DRUG PRODUCT LIST

6 - 191 (of 274)

NIACIN

CAPSULE; ORAL			
WAMPOCAP			
MEDPOINTE PHARM HLC	500MG	N11073	003
TABLET; ORAL			
NIACIN			
EVERYLIFE	500MG	N83203	001
HALSEY	500MG	N83453	001
IMPAX LABS	500MG	N83115	001
IVAX PHARMS	500MG	N83180	001
MK LABS	500MG	N83525	001
PUREPAC PHARM	500MG	N83271	001
SANDOZ	500MG	N83306	001
TABLICAPS	500MG	N84237	001
WATSON LABS	500MG	N83136	001
	500MG	N83305	001
	500MG	N85172	001
WEST WARD	500MG	N83718	001
NICOLAR			
SANOFI AVENTIS US	500MG	N83823	001
TABLET, EXTENDED RELEASE; ORAL			
NIACIN			
BARR	500MG	N76378	001 Apr 26, 2005
	750MG	N76378	002 Apr 26, 2005
	1GM	N76250	001 Apr 14, 2005
NIASPAN			
KOS LIFE	375MG	N20381	001 Jul 28, 1997
NIASPAN TITRATION STARTER PACK			
KOS LIFE	375MG;500MG;750MG	N20381	005 Jul 28, 1997

NIACINAMIDE; PYRIDOXINE HYDROCHLORIDE; TYROSINE

SUSPENSION; ORAL			
TPN SUSPENSION			
INTL MINERALS	15MG/5ML;3.75MG/5ML;600MG/5ML	N08378	003

NICLOSAMIDE

TABLET, CHEWABLE; ORAL			
NICLOCIDE			
BAYER PHARMS	500MG	N18669	001 May 14, 1982

NICOTINE

FILM, EXTENDED RELEASE; TRANSDERMAL			
NICOTROL			
MCNEIL CONS	15MG/16HR	N20536	001 Jul 03, 1996

NICOTINE POLACRILEX

GUM, CHEWING; BUCCAL			
NICOTINE POLACRILEX			
WATSON LABS	EQ 2MG BASE	N76568	001 Jul 29, 2004
	EQ 4MG BASE	N76569	002 Jul 29, 2004

NIFEDIPINE

CAPSULE; ORAL			
ADALAT			
BAYER PHARMS	10MG	N19478	001 Nov 27, 1985
	20MG	N19478	002 Sep 17, 1986
NIFEDIPINE			
CHASE LABS NJ	10MG	N72409	001 Jul 04, 1990
	20MG	N73421	001 Jun 19, 1991
TABLET, EXTENDED RELEASE; ORAL			
NIFEDIPINE			
MYLAN	30MG	N75108	001 Dec 17, 1999

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 192 (of 274)

NILUTAMIDETABLET; ORAL
NILANDRON

SANOFI AVENTIS US 50MG

N20169 001 Sep 19, 1996

NITROFURANTOINCAPSULE; ORAL
NITROFURANTOINWATSON LABS 50MG
100MGN84326 001
N84326 002TABLET; ORAL
FURADANTINPROCTER AND GAMBLE 50MG
100MGN08693 001
N08693 002

FURALAN

LANNETT 50MG
100MGN80017 001
N80017 002

NITROFURANTOIN

ELKINS SINN 50MG
100MGN80003 001
N80003 002IVAX PHARMS 50MG
100MGN80078 002
N80078 001SANDOZ 50MG
100MGN80043 001
N80043 002WATSON LABS 50MG
50MGN80447 001
N85797 001

100MG

N80447 002

100MG

N85796 001

WHITEWORTH TOWN PLSN 100MG

N84085 002

NITROFURANTOIN SODIUMINJECTABLE; INJECTION
IVADANTIN

PROCTER AND GAMBLE EQ 180MG BASE/VIAL

N12402 001

NITROFURANTOIN, MACROCRYSTALLINECAPSULE; ORAL
NITROFURANTOIN

MYLAN 100MG

N74967 002 Jul 09, 1997

NITROFURANTOIN MACROCRYSTALLINE

WATSON LABS 50MG
100MGN70248 001 Jun 24, 1988
N70249 001 Jun 24, 1988NITROFURAZONECREAM; TOPICAL
FURACIN

SHIRE 0.2%

N83789 001

DRESSING; TOPICAL
ACTIN-N

SHERWOOD MEDCL 0.2%

N17343 001

OINTMENT; TOPICAL
FURACIN

SHIRE 0.2%

N05795 001

NITROFURAZONE

AMBIX 0.2%

N86077 001

LANNETT 0.2%

N84393 001

PERRIGO NEW YORK 0.2%

N84968 001

WENDT 0.2%

N86766 001

POWDER; TOPICAL
FURACIN

SHIRE 0.2%

N83791 001

SOLUTION; TOPICAL
NITROFURAZONE

PERRIGO NEW YORK 0.2%

N85130 001

DISCONTINUED DRUG PRODUCT LIST

6 - 193 (of 274)

NITROFURAZONE

SOLUTION; TOPICAL NITROFURAZONE				
WENDT	0.2%		N87081	001

NITROGLYCERIN

AEROSOL; SUBLINGUAL NITROLINGUAL				
POHL BOSKAMP	0.4MG/SPRAY		N18705	001
FILM, EXTENDED RELEASE; TRANSDERMAL				Oct 31, 1985
TRANSDERM-NITRO				
NOVARTIS	0.1MG/HR		N20144	001
	0.2MG/HR		N20144	002
	0.4MG/HR		N20144	003
	0.6MG/HR		N20144	004
	0.8MG/HR		N20144	005
				Feb 27, 1996
				Feb 27, 1996
				Feb 27, 1996
				Feb 27, 1996
				Feb 27, 1996
INJECTABLE; INJECTION				
NITRO IV				
POHL BOSKAMP	5MG/ML		N18672	002
				Aug 30, 1983
NITRO-BID				
SANOFI AVENTIS US	5MG/ML		N18621	001
	10MG/ML		N71159	001
				Feb 28, 1990
NITROGLYCERIN				
ABRAXIS PHARM	5MG/ML		N70077	001
	5MG/ML		N71203	001
				May 08, 1987
INTL MEDICATION	5MG/ML		N70026	001
				Sep 10, 1985
LUITPOLD	5MG/ML		N71492	001
				May 24, 1988
SMITH AND NEPHEW	5MG/ML		N70633	001
	5MG/ML		N70634	001
				Jun 19, 1986
				Jun 19, 1986
NITROGLYCERIN IN DEXTROSE 5%				
HOSPIRA	0.1MG/ML		N74083	001
				Oct 26, 1994
NITROL				
RORER	0.8MG/ML		N18774	001
				Jan 19, 1983
NITRONAL				
POHL BOSKAMP	1MG/ML		N18672	001
				Aug 30, 1983
NITROSTAT				
PARKE DAVIS	0.8MG/ML		N18588	001
	5MG/ML		N18588	002
	5MG/ML		N70863	001
	10MG/ML		N70871	001
	10MG/ML		N70872	001
				Jan 08, 1987
TRIDIL				
MAYNE PHARMA USA	0.5MG/ML		N18537	002
	5MG/ML		N18537	001
				Jun 16, 1983

NONOXYNOL-9

AEROSOL; VAGINAL DOLFEN				
PERSONAL PRODS	12.5%		N14349	002
SPONGE; VAGINAL				
TODAY				
ALLENDALDE PHARMS	1GM		N18683	001
				Apr 01, 1983

NOREPINEPHRINE BITARTRATE; PROCAINE HYDROCHLORIDE; PROPOXYCAINE HYDROCHLORIDE

INJECTABLE; INJECTION				
RAVOCAINE AND NOVOCAIN W/ LEVOPHED				
EASTMAN KODAK	EQ 0.033MG BASE/ML; 2%; 0.4%		N08592	003

NORETHINDRONE

TABLET; ORAL NORLUTIN				
PARKE DAVIS	5MG		N10895	002

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 194 (of 274)

NORETHINDRONE ACETATE

TABLET; ORAL			
NORLUTATE			
PARKE DAVIS	5MG	N12184	002

NORFLOXACIN

SOLUTION/DROPS; OPHTHALMIC			
CHIBROXIN			
MERCK	0.3%	N19757	001 Jun 17, 1991

NORGESTREL

TABLET; ORAL			
OVRETTE			
WYETH PHARMS INC	0.075MG	N17031	001

NORTRIPTYLINE HYDROCHLORIDE

CAPSULE; ORAL			
AVENTYL HYDROCHLORIDE			
LILLY	EQ 10MG BASE	N14684	001
	EQ 25MG BASE	N14684	002

NOVOBIOCIN SODIUM

CAPSULE; ORAL			
ALBAMYCIN			
PHARMACIA AND UPJOHN	EQ 250MG BASE	N50339	001

NYSTATIN

CREAM; TOPICAL			
CANDEX			
BAYER PHARMS	100,000 UNITS/GM	N61810	001
MYKINAC			
ALPHARMA US PHARMS	100,000 UNITS/GM	N62387	001 Jul 29, 1982
NILSTAT			
LEDERLE	100,000 UNITS/GM	N61445	001
NYSTATIN			
TARO	100,000 UNITS/GM	N62457	001 Jul 28, 1983
TEVA	100,000 UNITS/GM	N61966	001
LOTION; TOPICAL			
CANDEX			
BAYER PHARMS	100,000 UNITS/ML	N50233	001
OINTMENT; TOPICAL			
MYCOSTATIN			
WESTWOOD SQUIBB	100,000 UNITS/GM	N60571	001
MYKINAC			
ALPHARMA US PHARMS	100,000 UNITS/GM	N62731	001 Sep 22, 1986
NILSTAT			
LEDERLE	100,000 UNITS/GM	N61444	001
PASTILLE; ORAL			
MYCOSTATIN			
BRISTOL MYERS SQUIBB	200,000 UNITS	N50619	001 Apr 09, 1987
POWDER; ORAL			
BARSTATIN 100			
BARLAN	100%	N62489	001 Apr 27, 1988
NILSTAT			
STADA PHARMS	100%	N50576	001 Dec 22, 1983
NYSTATIN			
PADDOCK	100%	N62613	001 Nov 26, 1985
SUPPOSITORY; VAGINAL			
NYSERT			
PROCTER AND GAMBLE	100,000 UNITS	N50478	001
SUSPENSION; ORAL			
MYCOSTATIN			
APOTHECON	100,000 UNITS/ML	N61533	001

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 195 (of 274)

NYSTATIN

SUSPENSION; ORAL

NYSTATIN

ALPHARMA US PHARMS	100,000 UNITS/ML	N62571	001	Oct 29, 1985
MORTON GROVE	100,000 UNITS/ML	N62835	001	Nov 19, 1987
PHARMADERM	100,000 UNITS/ML	N62518	001	Jul 06, 1984
PHARMAFAIR	100,000 UNITS/ML	N62541	001	Jan 16, 1985
ROXANE	100,000 UNITS/ML	N62832	001	Dec 27, 1991
TEVA	100,000 UNITS/ML	N62670	001	Jun 18, 1987
	100,000 UNITS/ML	N62776	001	Dec 17, 1987

NYSTEX

SAVAGE LABS	100,000 UNITS/ML	N62519	001	Jul 06, 1984
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TABLET; ORAL

NILSTAT

LEDERLE	500,000 UNITS	N61151	001	
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NYSTATIN

QUANTUM PHARMICS	500,000 UNITS	N62525	001	Oct 29, 1984
SANDOZ	500,000 UNITS	N62065	001	
USL PHARMA	500,000 UNITS	N62524	001	Nov 26, 1985
WATSON LABS	500,000 UNITS	N62402	001	Dec 16, 1982

TABLET; VAGINAL

KOROSTATIN

HOLLAND RANTOS	100,000 UNITS	N61718	001	
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MYCOSTATIN

BRISTOL MYERS SQUIBB	100,000 UNITS	N60577	001	
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NILSTAT

LEDERLE	100,000 UNITS	N61325	001	
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NYSTATIN

FOUGERA	100,000 UNITS	N62459	001	Nov 09, 1983
PHARMADERM	100,000 UNITS	N62460	001	Nov 09, 1983
QUANTUM PHARMICS	100,000 UNITS	N62509	001	Apr 03, 1984
SANDOZ	100,000 UNITS	N61965	001	
TEVA	100,000 UNITS	N62502	001	Dec 23, 1983
WATSON LABS	100,000 UNITS	N62176	001	

NYSTATIN; TRIAMCINOLONE ACETONIDE

CREAM; TOPICAL

MYCOLOG-II

APOTHECON	100,000 UNITS/GM;0.1%	N60576	002	May 01, 1985
	100,000 UNITS/GM;0.1%	N62606	001	May 15, 1985

MYCO-TRIACET II

TEVA	100,000 UNITS/GM;0.1%	N61954	002	Sep 20, 1985
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MYTREX F

SAVAGE LABS	100,000 UNITS/GM;0.1%	N62597	001	Oct 08, 1985
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NYSTATIN AND TRIAMCINOLONE ACETONIDE

ALPHARMA US PHARMS	100,000 UNITS/GM;0.1%	N63010	001	Dec 20, 1988
PERRIGO NEW YORK	100,000 UNITS/GM;0.1%	N62186	002	Jun 06, 1985
PHARMAFAIR	100,000 UNITS/GM;0.1%	N62657	001	Jul 30, 1986
TARO	100,000 UNITS/GM;0.1%	N62347	001	Mar 30, 1987

NYSTATIN-TRIAMCINOLONE ACETONIDE

PHARMADERM	100,000 UNITS/GM;0.1%	N62596	001	Oct 08, 1985
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OINTMENT; TOPICAL

MYCOLOG-II

APOTHECON	100,000 UNITS/GM;0.1%	N60572	001	Jun 28, 1985
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MYCO-TRIACET II

TEVA	100,000 UNITS/GM;0.1%	N62045	002	Nov 26, 1985
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MYTREX F

SAVAGE LABS	100,000 UNITS/GM;0.1%	N62601	001	Oct 09, 1985
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NYSTATIN AND TRIAMCINOLONE ACETONIDE

PERRIGO NEW YORK	100,000 UNITS/GM;0.1%	N62280	002	Oct 10, 1985
PHARMAFAIR	100,000 UNITS/GM;0.1%	N62656	001	Jul 30, 1986

NYSTATIN-TRIAMCINOLONE ACETONIDE

PHARMADERM	100,000 UNITS/GM;0.1%	N62603	001	Oct 09, 1985
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27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 196 (of 274)

OFLOXACININJECTABLE; INJECTION
FLOXIN

ORTHO MCNEIL PHARM	20MG/ML	N20087	002	Mar 31, 1992
	40MG/ML	N20087	003	Mar 31, 1992
FLOXIN IN DEXTROSE 5%				
ORTHO MCNEIL PHARM	400MG/100ML	N20087	001	Mar 31, 1992
FLOXIN IN DEXTROSE 5% IN PLASTIC CONTAINER				
ORTHO MCNEIL PHARM	4MG/ML	N20087	004	Mar 31, 1992
	400MG/100ML	N20087	005	Mar 31, 1992

ORPHENADRINE CITRATETABLET, EXTENDED RELEASE; ORAL
NORFLEX

3M	100MG	N12157	001	
ORPHENADRINE CITRATE				
ASCOT	100MG	N88067	001	Apr 06, 1983
SANDOZ	100MG	N85046	001	
WATSON LABS	100MG	N84303	001	

ORPHENADRINE HYDROCHLORIDETABLET; ORAL
DISIPAL

3M	50MG	N10653	001	
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OXACILLIN SODIUM

CAPSULE; ORAL

BACTOCILL

GLAXOSMITHKLINE	EQ 250MG BASE	N62241	001	
	EQ 500MG BASE	N62241	002	

OXACILLIN SODIUM

APOTHECON	EQ 250MG BASE	N61450	002	
	EQ 500MG BASE	N61450	001	

PROSTAPHLIN

APOTHECON	EQ 500MG BASE	N50118	002	
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FOR SOLUTION; ORAL

OXACILLIN SODIUM				
APOTHECON	EQ 250MG BASE/5ML	N61457	001	

PROSTAPHLIN

APOTHECON	EQ 250MG BASE/5ML	N50194	001	
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INJECTABLE; INJECTION

BACTOCILL

GLAXOSMITHKLINE	EQ 500MG BASE/VIAL	N61334	009	Mar 26, 1982
	EQ 1GM BASE/VIAL	N61334	006	Mar 26, 1982
	EQ 1GM BASE/VIAL	N62736	001	Dec 19, 1986
	EQ 2GM BASE/VIAL	N61334	007	Mar 26, 1982
	EQ 2GM BASE/VIAL	N62736	002	Dec 19, 1986
	EQ 4GM BASE/VIAL	N61334	008	Mar 26, 1982
	EQ 10GM BASE/VIAL	N61334	010	

OXACILLIN SODIUM

APOTHECON	EQ 250MG BASE/VIAL	N50195	001	
	EQ 500MG BASE/VIAL	N50195	002	
	EQ 1GM BASE/VIAL	N50195	003	
	EQ 2GM BASE/VIAL	N50195	004	
	EQ 4GM BASE/VIAL	N50195	005	

ELKINS SINN

	EQ 250MG BASE/VIAL	N62711	001	Feb 03, 1989
	EQ 500MG BASE/VIAL	N62711	002	Feb 03, 1989
	EQ 1GM BASE/VIAL	N62711	003	Feb 03, 1989
	EQ 2GM BASE/VIAL	N62711	004	Feb 03, 1989
	EQ 4GM BASE/VIAL	N62711	005	Feb 03, 1989
	EQ 10GM BASE/VIAL	N62711	006	Feb 03, 1989

IBI

	EQ 125MG BASE/VIAL	N62798	003	Dec 11, 1995
	EQ 250MG BASE/VIAL	N62798	004	Dec 11, 1995
	EQ 500MG BASE/VIAL	N62798	005	Dec 11, 1995

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 197 (of 274)

OXACILLIN SODIUMINJECTABLE; INJECTION
OXACILLIN SODIUM

IBI	EQ 1GM BASE/VIAL	N62798	001	Dec 11, 1995
	EQ 2GM BASE/VIAL	N62798	002	Dec 11, 1995

OXALIPLATININJECTABLE; IV (INFUSION)
ELOXATIN

SANOFI AVENTIS US	50MG/VIAL	N21492	001	Aug 09, 2002
	100MG/VIAL	N21492	002	Aug 09, 2002

OXAMNIQUINECAPSULE; ORAL
VANSIL

PFIZER	250MG	N18069	001	
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OXAPROZIN POTASSIUMTABLET; ORAL
DAYPRO ALTA
GD SEARLE

600MG	N20776	001	Oct 17, 2002
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OXAZEPAMCAPSULE; ORAL
OXAZEPAM

AM THERAP	10MG	N71955	001	Mar 03, 1988
	15MG	N71956	001	Mar 03, 1988
	30MG	N71957	001	Mar 03, 1988
MUTUAL PHARM	10MG	N70957	001	Aug 10, 1987
	15MG	N71025	001	Aug 10, 1987
	30MG	N71026	001	Aug 10, 1987
MYLAN	10MG	N71713	001	Oct 20, 1987
	15MG	N71714	001	Oct 20, 1987
	30MG	N71715	001	Oct 20, 1987
SERAX				
ALPHARMA US PHARMS	10MG	N15539	002	
	15MG	N15539	004	
	30MG	N15539	006	

ZAXOPAM

QUANTUM PHARMICS	10MG	N70650	001	Mar 01, 1988
	15MG	N70640	001	Mar 01, 1988
	30MG	N70641	001	Mar 01, 1988

TABLET; ORAL
OXAZEPAM

MUTUAL PHARM	15MG	N70683	001	Jan 16, 1987
PARKE DAVIS	15MG	N71508	001	Feb 02, 1987
WATSON LABS	15MG	N71494	001	Apr 21, 1987
SERAX				
ALPHARMA US PHARMS	15MG	N15539	008	

OPRENOLOL HYDROCHLORIDECAPSULE; ORAL
TRASICOR
NOVARTIS

20MG	N18166	001	Dec 28, 1983
40MG	N18166	002	Dec 28, 1983
80MG	N18166	003	Dec 28, 1983
160MG	N18166	004	Dec 28, 1983

OXTRIPHYLLINESOLUTION; ORAL
CHOLEDYL

PARKE DAVIS	100MG/5ML	N09268	012	Nov 27, 1984
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OXTRIPHYLLINE

MORTON GROVE	100MG/5ML	N88243	001	Dec 05, 1983
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27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 198 (of 274)

OXTRIPHYLLINESYRUP; ORAL
CHOLEDYL

PARKE DAVIS 50MG/5ML

N09268 011

OXTRIPHYLLINE PEDIATRIC

MORTON GROVE 50MG/5ML

N88242 001 Dec 05, 1983

TABLET, DELAYED RELEASE; ORAL
CHOLEDYL

PARKE DAVIS 100MG

N09268 003

200MG

N09268 007

OXTRIPHYLLINE

WATSON LABS 100MG

N87866 001 Aug 25, 1983

200MG

N87835 001 Aug 25, 1983

OXYBUTYNIN CHLORIDE

TABLET; ORAL

OXYBUTYNIN CHLORIDE

QUANTUM PHARMICS 5MG

N72296 001 Dec 08, 1988

USL PHARMA 5MG

N70746 001 Mar 10, 1988

WATSON LABS 5MG

N72485 001 Apr 19, 1989

OXYCODONE HYDROCHLORIDE

TABLET, EXTENDED RELEASE; ORAL

OXYCONTIN

PURDUE PHARMA LP 160MG

N20553 005 Mar 15, 2000

ROXICODONE

ROXANE 10MG

N20932 001 Oct 26, 1998

30MG

N20932 002 Oct 26, 1998

OXYMORPHONE HYDROCHLORIDE

INJECTABLE; INJECTION

NUMORPHAN

ENDO PHARMS 1.5MG/ML

N11707 001

SUPPOSITORY; RECTAL

NUMORPHAN

ENDO PHARMS 5MG

N11738 004

OXYPHENBUTAZONE

TABLET; ORAL

OXYPHENBUTAZONE

WATSON LABS 100MG

N88399 001 Sep 17, 1984

TANDEARIL

NOVARTIS 100MG

N12542 004 Sep 03, 1982

OXYPHENCYCLIMINE HYDROCHLORIDE

TABLET; ORAL

DARICON

PFIZER 10MG

N11612 001

OXYPHENONIUM BROMIDE

TABLET; ORAL

ANTRENYL

NOVARTIS 5MG

N08492 002

OXYTETRACYCLINE

TABLET; ORAL

TERRAMYCIN

PFIZER 250MG

N50287 001

OXYTETRACYCLINE CALCIUM

SYRUP; ORAL

TERRAMYCIN

PFIZER EQ 125MG BASE/5ML

N60595 001

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 199 (of 274)

OXYTETRACYCLINE HYDROCHLORIDE

CAPSULE; ORAL			
OXY-KESSO-TETRA			
FERRANTE	EQ 250MG BASE	N60179	001
OXYTETRACYCLINE HYDROCHLORIDE			
IMPAX LABS	EQ 250MG BASE	N60760	001
PROTER	EQ 250MG BASE	N60869	001
PUREPAC PHARM	EQ 250MG BASE	N60634	001
WEST WARD	EQ 250MG BASE	N60770	001
TERRAMYCIN			
PFIZER	EQ 125MG BASE	N50286	001
INJECTABLE; INJECTION			
TERRAMYCIN			
PFIZER	EQ 250MG BASE/VIAL	N60586	001
	EQ 500MG BASE/VIAL	N60586	002

OXYTETRACYCLINE HYDROCHLORIDE; POLYMYXIN B SULFATE

OINTMENT; OTIC			
TERRAMYCIN W/ POLYMYXIN			
PFIZER	EQ 5MG BASE/GM;10,000 UNITS/GM	N61841	001
TABLET; VAGINAL			
TERRAMYCIN-POLYMYXIN			
PFIZER	EQ 100MG BASE;100,000 UNITS	N61009	001

OXYTOCIN

INJECTABLE; INJECTION			
OXYTOCIN			
BAXTER HLTHCARE CORP	10USP UNITS/ML	N18243	001
OXYTOCIN 10 USP UNITS IN DEXTROSE 5%			
ABBOTT	1USP UNITS/100ML	N19185	004 Mar 29, 1985
	2USP UNITS/100ML	N19185	003 Mar 29, 1985
OXYTOCIN 20 USP UNITS IN DEXTROSE 5%			
ABBOTT	2USP UNITS/100ML	N19185	002 Mar 29, 1985
OXYTOCIN 5 USP UNITS IN DEXTROSE 5%			
ABBOTT	1USP UNITS/100ML	N19185	001 Mar 29, 1985
SYNTOCINON			
NOVARTIS	10USP UNITS/ML	N18245	001
SOLUTION; NASAL			
SYNTOCINON			
NOVARTIS	40USP UNITS/ML	N12285	001

PALIPERIDONE

TABLET, EXTENDED RELEASE; ORAL			
INVEGA			
JANSSEN LP	12MG	N21999	004 Dec 19, 2006

PAMIDRONATE DISODIUM

INJECTABLE; INJECTION			
AREDIA			
NOVARTIS	60MG/VIAL	N20036	003 May 06, 1993

PANCRELIPASE (AMYLASE;LIPASE;PROTEASE)

CAPSULE; ORAL			
COTAZYM			
ORGANON USA INC	30,000USP UNITS;8,000USP UNITS;30,000USP UNITS	N20580	001 Dec 09, 1996

PANCURONIUM BROMIDE

INJECTABLE; INJECTION			
PANCURONIUM BROMIDE			
ASTRAZENECA	1MG/ML	N72210	001 Mar 31, 1988
	2MG/ML	N72211	001 Mar 31, 1988
	2MG/ML	N72212	001 Mar 31, 1988
	2MG/ML	N72213	001 Mar 31, 1988

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 200 (of 274)

PANCURONIUM BROMIDE

INJECTABLE; INJECTION

PAVULON

ORGANON USA INC

1MG/ML

N17015 002

2MG/ML

N17015 001

PARAMETHADIONE

CAPSULE; ORAL

PARADIONE

ABBOTT

150MG

N06800 003

300MG

N06800 001

SOLUTION; ORAL

PARADIONE

ABBOTT

300MG/ML

N06800 002

PARAMETHASONE ACETATE

TABLET; ORAL

HALDRONE

LILLY

1MG

N12772 005

2MG

N12772 006

PARGYLINE HYDROCHLORIDE

TABLET; ORAL

EUTONYL

ABBOTT

10MG

N13448 002

25MG

N13448 003

50MG

N13448 004

PAROMOMYCIN SULFATE

CAPSULE; ORAL

HUMATIN

PARKEDALE

EQ 250MG BASE

N60521 001

SYRUP; ORAL

HUMATIN

PARKE DAVIS

EQ 125MG BASE/5ML

N60522 001

PAROXETINE HYDROCHLORIDE

CAPSULE; ORAL

PAXIL

GLAXOSMITHKLINE

EQ 10MG BASE **Federal Register
determination that product was not
discontinued or withdrawn for safety
or efficacy reasons**

N20885 001

Oct 09, 1998

EQ 20MG BASE **Federal Register
determination that product was not
discontinued or withdrawn for safety
or efficacy reasons**

N20885 002

Oct 09, 1998

EQ 30MG BASE **Federal Register
determination that product was not
discontinued or withdrawn for safety
or efficacy reasons**

N20885 003

Oct 09, 1998

EQ 40MG BASE **Federal Register
determination that product was not
discontinued or withdrawn for safety
or efficacy reasons**

N20885 004

Oct 09, 1998

TABLET; ORAL

PAXIL

GLAXOSMITHKLINE

EQ 50MG BASE

N20031 004

Dec 29, 1992

PEMOLINE

TABLET; ORAL

CYLERT

ABBOTT

18.75MG

N16832 001

37.5MG

N16832 002

75MG

N16832 003

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 201 (of 274)

PEMOLINETABLET; ORAL
PEMOLINE

ACTAVIS TOTOWA	18.75MG	N75595 001	Feb 28, 2000
	37.5MG	N75595 002	Feb 28, 2000
	75MG	N75595 003	Feb 28, 2000
MALLINCKRODT	18.75MG	N75726 003	Mar 30, 2001
	37.5MG	N75726 002	Mar 30, 2001
	75MG	N75726 001	Mar 30, 2001
SANDOZ	18.75MG	N75286 001	Dec 27, 1999
	37.5MG	N75286 002	Jun 30, 1999
	75MG	N75286 003	Jun 30, 1999
TEVA PHARMS	18.75MG	N75030 003	Feb 22, 2000
	37.5MG	N75030 001	Jan 29, 1999
	75MG	N75030 002	Jan 29, 1999
VINTAGE PHARMS	18.75MG	N75328 001	Apr 19, 2000
	37.5MG	N75328 002	Apr 19, 2000
	75MG	N75328 003	Apr 19, 2000
WATSON LABS	18.75MG	N75287 001	Jun 13, 2001
	37.5MG	N75287 002	Sep 18, 2000
	75MG	N75287 003	Sep 18, 2000

TABLET, CHEWABLE; ORAL
CYLERT

ABBOTT	37.5MG	N17703 001	
PEMOLINE			
ACTAVIS TOTOWA	37.5MG	N75678 001	Jul 26, 2000
TEVA PHARMS	37.5MG	N75555 001	Feb 18, 2000

PENBUTOLOL SULFATETABLET; ORAL
LEVATOL

SCHWARZ PHARMA	10MG	N18976 001	Dec 30, 1987
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PENICILLAMINECAPSULE; ORAL
CUPRIMINE

ATON	125MG	N19853 002	
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PENICILLIN G BENZATHINEINJECTABLE; INJECTION
BICILLIN L-A

WYETH AYERST	300,000 UNITS/ML	N50131 001	
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SUSPENSION; ORAL
BICILLIN

WYETH AYERST	300,000 UNITS/5ML	N50126 002	
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TABLET; ORAL
BICILLIN

WYETH AYERST	200,000 UNITS	N50128 001	
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PENICILLIN G POTASSIUMFOR SOLUTION; ORAL
PENICILLIN

TEVA	200,000 UNITS/5ML	N60307 002	
	400,000 UNITS/5ML	N60307 004	

PENICILLIN G POTASSIUM
MYLAN

200,000 UNITS/5ML	N60752 003	
250,000 UNITS/5ML	N60752 002	
400,000 UNITS/5ML	N60752 001	
PUREPAC PHARM	250,000 UNITS/5ML	N61740 001
	400,000 UNITS/5ML	N61740 002

PENICILLIN-2

TEVA	250,000 UNITS/5ML	N60307 003	
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PENTIDS '200'

APOTHECON	200,000 UNITS/5ML	N62149 001	
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27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 202 (of 274)

PENICILLIN G POTASSIUM

FOR SOLUTION; ORAL

PENTIDS '400'

APOTHECON

400,000 UNITS/5ML

N62149 002

PFIZERPEN G

PFIZER

400,000 UNITS/5ML

N60587 001

INJECTABLE; INJECTION

PENICILLIN G POTASSIUM

APOTHECON

1,000,000 UNITS/VIAL

N60362 001

5,000,000 UNITS/VIAL

N60362 003

10,000,000 UNITS/VIAL

N60362 004

20,000,000 UNITS/VIAL

N60362 002

CONSOLIDATED PHARM

500,000 UNITS/VIAL

N60806 001

1,000,000 UNITS/VIAL

N60806 002

5,000,000 UNITS/VIAL

N60806 003

10,000,000 UNITS/VIAL

N60806 004

LILLY

200,000 UNITS/VIAL

N60384 004

500,000 UNITS/VIAL

N60384 003

1,000,000 UNITS/VIAL

N60384 002

5,000,000 UNITS/VIAL

N60384 001

20,000,000 UNITS/VIAL

N60384 005

20,000,000 UNITS/VIAL

N60601 001

MARSAM PHARMS LLC

1,000,000 UNITS/VIAL

N62991 001

Sep 13, 1988

5,000,000 UNITS/VIAL

N62991 002

Sep 13, 1988

10,000,000 UNITS/VIAL

N62991 003

Sep 13, 1988

20,000,000 UNITS/VIAL

N62991 004

Sep 13, 1988

PARKE DAVIS

1,000,000 UNITS/VIAL

N62003 001

5,000,000 UNITS/VIAL

N62003 002

TABLET; ORAL

PENICILLIN G POTASSIUM

APOTHECON

250,000 UNITS

N60392 003

IVAX PHARMS

400,000 UNITS

N60073 004

LILLY

250,000 UNITS

N60403 001

MYLAN

200,000 UNITS

N60781 001

250,000 UNITS

N60781 002

400,000 UNITS

N60781 003

500,000 UNITS

N60781 005

800,000 UNITS

N60781 004

PUREPAC PHARM

200,000 UNITS

N61588 001

250,000 UNITS

N61588 002

400,000 UNITS

N61588 003

TEVA

200,000 UNITS

N60306 001

250,000 UNITS

N60306 002

400,000 UNITS

N60306 003

500,000 UNITS

N60306 004

WYETH AYERST

200,000 UNITS

N60413 001

250,000 UNITS

N60413 002

400,000 UNITS

N60413 003

PENTIDS '200'

APOTHECON

200,000 UNITS

N62155 001

PENTIDS '250'

APOTHECON

250,000 UNITS

N62155 002

PENTIDS '400'

APOTHECON

400,000 UNITS

N60392 004

400,000 UNITS

N62155 003

PENTIDS '800'

APOTHECON

800,000 UNITS

N60392 005

800,000 UNITS

N62155 004

PFIZERPEN G

PFIZER

50,000 UNITS

N60075 001

100,000 UNITS

N60075 002

200,000 UNITS

N60075 003

250,000 UNITS

N60075 004

400,000 UNITS

N60075 005

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 203 (of 274)

PENICILLIN G POTASSIUM

TABLET; ORAL			
PFIZERPEN G			
PFIZER	800,000 UNITS	N60075	006

PENICILLIN G PROCAINE

INJECTABLE; INJECTION			
DURACILLIN A.S.			
LILLY	300,000 UNITS/ML	N60093	001
PENICILLIN G PROCAINE			
CONSOLIDATED PHARM	300,000 UNITS/ML	N60800	001
	600,000 UNITS/1.2ML	N60800	002
PARKE DAVIS	300,000 UNITS/ML	N62029	001
PFIZER	300,000 UNITS/VIAL	N60099	001
	1,500,000 UNITS/VIAL	N60099	002
PFIZERPEN-AS			
PFIZER	300,000 UNITS/ML	N60286	001
	600,000 UNITS/ML	N60286	002

PENICILLIN G SODIUM

INJECTABLE; IM-IV			
PENICILLIN G SODIUM			
MARSAM PHARMS LLC	5,000,000 UNITS/VIAL	N63014	001 Sep 13, 1988
INJECTABLE; INJECTION			
PENICILLIN G SODIUM			
BRISTOL MYERS SQUIBB	5,000,000 UNITS/VIAL	N61935	001
COPANOS	5,000,000 UNITS/VIAL	N61051	001
PHARMACIA AND UPJOHN	1,000,000 UNITS/VIAL	N61046	001

PENICILLIN V

FOR SUSPENSION; ORAL			
V-CILLIN			
LILLY	125MG/0.6ML	N60002	001

PENICILLIN V POTASSIUM

FOR SOLUTION; ORAL			
BEEPEN-VK			
GLAXOSMITHKLINE	EQ 125MG BASE/5ML	N62270	001
	EQ 250MG BASE/5ML	N62270	002
BETAPEN-VK			
APOTHECON	EQ 125MG BASE/5ML	N61149	001
	EQ 250MG BASE/5ML	N61149	002
LEDERCILLIN VK			
LEDERLE	EQ 125MG BASE/5ML	N60136	001
	EQ 250MG BASE/5ML	N60136	002
PENAPAR-VK			
PARKE DAVIS	EQ 125MG BASE/5ML	N62002	001
	EQ 250MG BASE/5ML	N62002	002
PENICILLIN V POTASSIUM			
MYLAN	EQ 125MG BASE/5ML	N61624	002
	EQ 250MG BASE/5ML	N61624	001
PUREPAC PHARM	EQ 125MG BASE/5ML	N61758	001
	EQ 250MG BASE/5ML	N61758	002
PEN-VEE K			
WYETH AYERST	EQ 125MG BASE/5ML	N60007	001
	EQ 250MG BASE/5ML	N60007	002
PFIZERPEN VK			
PFIZER	EQ 125MG BASE/5ML	N61815	001
	EQ 250MG BASE/5ML	N61815	002
V-CILLIN K			
LILLY	EQ 125MG BASE/5ML	N60004	001
	EQ 250MG BASE/5ML	N60004	002
VEETIDS '125'			
APOTHECON	EQ 125MG BASE/5ML	N61206	001

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 204 (of 274)

PENICILLIN V POTASSIUM

FOR SOLUTION; ORAL

VEETIDS '125'

APOTHECON

EQ 125MG BASE/5ML

N62153 001

VEETIDS '250'

APOTHECON

EQ 250MG BASE/5ML

N61206 002

EQ 250MG BASE/5ML

N62153 002

TABLET; ORAL

BEEPEN-VK

GLAXOSMITHKLINE

EQ 250MG BASE

N62273 001

EQ 500MG BASE

N62273 002

BETAPEN-VK

BRISTOL

EQ 250MG BASE

N61150 001

EQ 500MG BASE

N61150 002

LEDERCILLIN VK

LEDERLE

EQ 250MG BASE

N60134 001

EQ 500MG BASE

N60134 002

PENAPAR-VK

PARKE DAVIS

EQ 250MG BASE

N62001 001

EQ 500MG BASE

N62001 002

PENICILLIN V POTASSIUM

AM ANTIBIOTICS

EQ 250MG BASE

N61528 001

EQ 500MG BASE

N61528 002

IVAX PHARMS

EQ 125MG BASE

N60518 001

EQ 250MG BASE

N60518 002

EQ 500MG BASE

N60518 003

MYLAN

EQ 250MG BASE

N61530 001

EQ 500MG BASE

N61530 002

PUREPAC PHARM

EQ 125MG BASE

N61571 001

EQ 250MG BASE

N61571 002

EQ 500MG BASE

N61571 003

PEN-VEE K

WYETH AYERST

EQ 125MG BASE

N60006 001

EQ 250MG BASE

N60006 002

EQ 500MG BASE

N60006 003

PFIZERPEN VK

PFIZER

EQ 250MG BASE

N61836 001

EQ 500MG BASE

N61836 002

UTICILLIN VK

PHARMACIA AND UPJOHN

EQ 250MG BASE

N61651 001

EQ 500MG BASE

N61651 002

V-CILLIN K

LILLY

EQ 125MG BASE

N60003 001

EQ 250MG BASE

N60003 002

EQ 500MG BASE

N60003 003

VEETIDS

APOTHECON

EQ 250MG BASE

N61411 001

EQ 500MG BASE

N61411 002

VEETIDS '250'

APOTHECON

EQ 250MG BASE

N61164 001

EQ 250MG BASE

N62156 002

VEETIDS '500'

APOTHECON

EQ 500MG BASE

N61164 002

EQ 500MG BASE

N62156 001

PENTAGASTRIN

INJECTABLE; INJECTION

PEPTAVLON

WYETH AYERST

0.25MG/ML **Federal Register
determination that product was not
discontinued or withdrawn for safety
or efficacy reasons**

N17048 001

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 205 (of 274)

PENTAMIDINE ISETHIONATEINJECTABLE; INJECTION
PENTACARINAT

ARMOUR PHARM 300MG/VIAL

N73447 001 Apr 28, 1994

PENTAMIDINE ISETHIONATE

BAXTER HLTHCARE 300MG/VIAL

N73617 001 Dec 18, 1995

PENTAZOCINE HYDROCHLORIDE

TABLET; ORAL

TALWIN 50

SANOFI AVENTIS US EQ 50MG BASE

N16732 001

PENTETATE CALCIUM TRISODIUM YB-169

INJECTABLE; INJECTION

YTTERBIUM YB 169 DTPA

3M 2mCi/ML

N17518 001

PENTOBARBITAL

ELIXIR; ORAL

NEMBUTAL

OVATION PHARMS 18.2MG/5ML

N83244 001

PENTOBARBITAL SODIUM

CAPSULE; ORAL

NEMBUTAL SODIUM

OVATION PHARMS 30MG
50MG
100MG

N84095 001

N84093 001

N83245 001

PENTOBARBITAL SODIUM

LANNETT 50MG
100MG

N85937 001

N85915 001

VITARINE 100MG

N83284 001

WHITEWORTH TOWN PLSN 100MG

N83338 001

SODIUM PENTOBARBITAL

ANABOLIC 100MG

N84590 001

ELKINS SINN 100MG

N83368 001

EVERYLIFE 100MG

N83259 001

HALSEY 100MG

N84677 001

IVAX PHARMS 50MG

N83461 001

100MG

N83461 002

PARKE DAVIS 100MG

N84156 001

PERRIGO 100MG

N84560 001

PUREPAC PHARM 100MG

N83301 001

VALEANT PHARM INTL 100MG

N83264 001

WATSON LABS 100MG

N85791 001

WYETH AYERST 100MG

N83239 001

INJECTABLE; INJECTION

PENTOBARBITAL SODIUM

ELKINS SINN 50MG/ML

N83270 001

SODIUM PENTOBARBITAL

WYETH AYERST 50MG/ML

N83261 001

SUPPOSITORY; RECTAL

NEMBUTAL

OVATION PHARMS 30MG

N83247 001 Jan 25, 1982

60MG

N83247 002 Jan 25, 1982

120MG

N83247 003 Jan 25, 1982

200MG

N83247 004 Jan 25, 1982

TABLET; ORAL

PENTOBARBITAL SODIUM

VITARINE 100MG

N83285 001

SODIUM PENTOBARBITAL

ANABOLIC 100MG

N84238 001

DISCONTINUED DRUG PRODUCT LIST

6 - 206 (of 274)

PENTOLINIUM TARTRATE

INJECTABLE; INJECTION
ANSOLYSEN

WYETH AYERST 10MG/ML N09372 001

PERFLUBRON

LIQUID; ORAL
IMAGENT

ALLIANCE PHARM 100% N20091 001 Aug 13, 1993

PERFLUOROPOLYMETHYLISOPROPYL ETHER; POLYTETRAFLUOROETHYLENE

PASTE; TOPICAL
SKIN EXPOSURE REDUCTION PASTE AGAINST CHEMICAL WARFARE AGENTS
US ARMY 50%;50%

N21084 001 Feb 17, 2000

PERMETHRIN

LOTION; TOPICAL
NIX

GLAXOSMITHKLINE 1% N19435 001 Mar 31, 1986

PERPHENAZINE

CONCENTRATE; ORAL
TRILAFON

SCHERING 16MG/5ML N11557 001

INJECTABLE; INJECTION
TRILAFON

SCHERING 5MG/ML N11213 002

SYRUP; ORAL
TRILAFON

SCHERING 2MG/5ML N11294 002

TABLET; ORAL
TRILAFON

SCHERING 2MG N10775 001

4MG N10775 002

8MG N10775 003

16MG N10775 004

TABLET, EXTENDED RELEASE; ORAL
TRILAFON

SCHERING 8MG N11361 002

PHENACEMIDE

TABLET; ORAL
PHENURONE

ABBOTT 500MG N07707 001

PHENAZOPYRIDINE HYDROCHLORIDE; SULFAMETHOXAZOLE

TABLET; ORAL
AZO GANTANOL

ROCHE 100MG;500MG N13294 001 Sep 10, 1987

PHENAZOPYRIDINE HYDROCHLORIDE; SULFAMETHOXAZOLE; TRIMETHOPRIM

TABLET; ORAL

SULFAMETHOXAZOLE AND TRIMETHOPRIM AND PENAZOPYRIDINE HYDROCHLORIDE
ABLE

200MG, N/A, N/A; N/A, 800MG, 160MG N21105 001 Jun 26, 2001

PHENAZOPYRIDINE HYDROCHLORIDE; SULFISOXAZOLE

TABLET; ORAL
AZO GANTRISIN

ROCHE 50MG;500MG N19358 001 Aug 31, 1990

PHENDIMETRAZINE TARTRATE

CAPSULE; ORAL
PHENAZINE

MAST MM 35MG N86523 001

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 207 (of 274)

PHENDIMETRAZINE TARTRATE

CAPSULE; ORAL

PHENAZINE

MAST MM

35MG

N86524 001

35MG

N86525 001

PHENDIMETRAZINE TARTRATE

SANDOZ

35MG

N85633 001

35MG

N85694 001

35MG

N85702 001

VITARINE

35MG

N85634 001

35MG

N85645 001

35MG

N85670 001

35MG

N86403 001

35MG

N86408 001

35MG

N86410 001

35MG

N87424 001

SPRX-3

SOLVAY

35MG

N85897 001

STATOBEX

TEVA

35MG

N85507 001

CAPSULE, EXTENDED RELEASE; ORAL

MELFIAT-105

NUMARK

105MG

N87487 001

Oct 13, 1982

PHENDIMETRAZINE TARTRATE

GRAHAM DM

105MG

N87214 001

May 26, 1982

105MG

N88020 001

Aug 16, 1982

105MG

N88028 001

Aug 16, 1982

105MG

N88062 001

Sep 13, 1982

105MG

N88063 001

Sep 10, 1982

105MG

N88111 001

Oct 18, 1982

SANDOZ

105MG

N87378 001

SPRX-105

NUMARK

105MG

N88024 001

Dec 22, 1982

TABLET; ORAL

ADPHEN

FERNDAL LABS

35MG

N83655 001

ALPHAZINE

SANDOZ

35MG

N85034 001

CAM-METRAZINE

ABC HOLDING

35MG

N85511 001

CAMALL

35MG

N85756 001

TG UNITED LABS

35MG

N83922 001

35MG

N85318 001

35MG

N85320 001

35MG

N85321 001

DI-METREX

LEINER

35MG

N85698 001

MELFIAT

NUMARK

35MG

N83790 002

METRA

FOREST PHARMS

35MG

N83754 001

PHENAZINE

MAST MM

35MG

N87305 001

PHENAZINE-35

ABC HOLDING

35MG

N85512 001

PHENDIMETRAZINE TARTRATE

ANABOLIC

35MG

N86020 001

BARR

35MG

N83644 001

35MG

N83684 001

35MG

N83686 001

35MG

N83687 001

35MG

N84831 001

35MG

N84834 001

35MG

N84835 001

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 208 (of 274)

PHENDIMETRAZINE TARTRATE

TABLET; ORAL

PHENDIMETRAZINE TARTRATE

FERNDAL LABS	35MG	N86834	001	Sep 15, 1983
INWOOD LABS	35MG	N84740	001	
	35MG	N84741	001	
	35MG	N84742	001	
	35MG	N84743	001	
IVAX PHARMS	35MG	N83682	001	
	35MG	N85611	001	
	35MG	N85612	001	
KV PHARM	35MG	N84138	001	
	35MG	N84141	001	
	35MG	N85525	001	
LEINER	35MG	N85199	001	
	35MG	N85697	001	
MFG CHEMISTS	35MG	N85914	001	
NUMARK	35MG	N83790	001	
SANDOZ	35MG	N85402	001	
	35MG	N85497	001	
	35MG	N85830	001	
	35MG	N86365	001	
	35MG	N86370	001	
SOLVAY	35MG	N83993	001	
TG UNITED LABS	35MG	N85761	001	
	35MG	N85941	001	Jun 27, 1983
USL PHARMA	35MG	N83805	001	
	35MG	N84398	001	
	35MG	N84399	001	
VITARINE	35MG	N85519	001	
	35MG	N86005	001	
	35MG	N86106	001	
WATSON LABS	35MG	N85767	001	
	35MG	N85768	001	
	35MG	N85770	001	
	35MG	N85773	001	
PLEGINE				
WYETH AYERST	35MG	N12248	001	
STATOBEX				
TEVA	35MG	N86013	001	
STATOBEX-G				
TEVA	35MG	N85095	001	
X-TROZINE				
SHIRE RICHWOOD	35MG	N86550	001	
	35MG	N86551	001	
	35MG	N86552	001	
	35MG	N86553	001	
	35MG	N86554	001	

PHENINDIONE

TABLET; ORAL

HEDULIN

SANOFI AVENTIS US	50MG	N08767	002
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PHENMETRAZINE HYDROCHLORIDE

TABLET; ORAL

PRELUDIN

BOEHRINGER INGELHEIM	25MG	N10460	005
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TABLET, EXTENDED RELEASE; ORAL

PRELUDIN

BOEHRINGER INGELHEIM	50MG	N11752	004
	75MG	N11752	003

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 209 (of 274)

PHENPROCOUONTABLET; ORAL
LIQUAMAR

ORGANON USA INC 3MG

N11228 001

PHENSUXIMIDECAPSULE; ORAL
MILONTIN

PARKE DAVIS 500MG

N08855 004

PHENTERMINE HYDROCHLORIDECAPSULE; ORAL
FASTIN

GLAXOSMITHKLINE 30MG

N17352 001

OBESTIN-30

FERNDAL LABS 30MG

N87144 001

OBY-TRIM

SHIRE RICHWOOD 30MG

N87764 001 Mar 18, 1982

ONA-MAST

MAST MM 30MG

N86511 001

30MG

N86516 001

PHENTERMINE HYDROCHLORIDE

ABC HOLDING 30MG

N85411 001

ABLE 15MG

N40497 001 Mar 13, 2003

30MG

N40403 001 Aug 30, 2001

30MG

N40427 001 Aug 30, 2001

CAMALL 15MG

N86735 001

30MG

N87226 001

DURAMED PHARMS BARR 30MG

N88948 001 Apr 25, 1986

IVAX PHARMS 30MG

N86329 001

LANNETT 30MG

N87022 001 Feb 03, 1983

SANDOZ 15MG

N87301 001

30MG

N87208 001

30MG

N87223 001

37.5MG

N88414 001 Oct 19, 1983

TEVA 30MG

N86911 001

30MG

N87126 001

30MG

N87777 001 Nov 01, 1985

30MG

N88612 001 Apr 04, 1984

30MG

N88613 001 Apr 09, 1984

30MG

N88614 001 Apr 09, 1984

TG UNITED LABS 18.75MG

N88576 001 May 23, 1984

30MG

N85417 001

30MG

N86732 002

30MG

N87215 001

37.5MG

N87915 001 Dec 22, 1983

37.5MG

N87918 001 Dec 22, 1983

37.5MG

N87930 001 Oct 14, 1983

37.5MG

N88610 001 Jun 04, 1984

37.5MG

N88611 001 Jun 04, 1984

37.5MG

N88625 001 Aug 23, 1984

USL PHARMA 30MG

N84487 001 Apr 09, 1982

30MG

N88430 001 Mar 27, 1984

30MG

N88797 001 Dec 10, 1984

VITARINE 30MG

N87202 001

30MG

N87235 001

WATSON LABS 30MG

N86740 001 Mar 21, 1985

TABLET; ORAL

ONA-MAST

MAST MM 8MG

N86260 001

PHENTERMINE HYDROCHLORIDE

ABLE 37.5MG

N40402 001 Aug 30, 2001

IVAX PHARMS 8MG

N85553 001

SANDOZ 8MG

N85671 001

DISCONTINUED DRUG PRODUCT LIST

6 - 210 (of 274)

PHENTERMINE HYDROCHLORIDE

TABLET; ORAL

PHENTERMINE HYDROCHLORIDE

SANDOZ	8MG	N85689	001	
TG UNITED LABS	8MG	N83923	001	
	8MG	N85319	001	
	37.5MG	N87805	001	Dec 06, 1982
	37.5MG	N88596	001	Apr 04, 1984
USL PHARMA	8MG	N83804	001	
	37.5MG	N88910	001	Jul 17, 1985
	37.5MG	N88917	001	Jul 17, 1985
VITARINE	8MG	N86453	001	
	8MG	N86456	001	
WATSON LABS	8MG	N85739	001	
TORA				
SOLVAY	8MG	N84035	001	

PHENTERMINE RESIN COMPLEX

CAPSULE, EXTENDED RELEASE; ORAL

PHENTERMINE RESIN 30

QUANTUM PHARMICS	EQ 30MG BASE	N89120	001	Feb 04, 1988
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PHENYL AMINOSALICYLATE

POWDER; ORAL

PHENY-PAS-TEBAMIN

PURDUE FREDERICK	50%	N11695	002	
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TABLET; ORAL

PHENY-PAS-TEBAMIN

PURDUE FREDERICK	500MG	N11695	003	
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PHENYLBUTAZONE

CAPSULE; ORAL

AZOLID

SANOFI AVENTIS US	100MG	N87260	001	
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BUTAZOLIDIN

NOVARTIS	100MG	N08319	009	
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PHENYLBUTAZONE

IVAX PHARMS	100MG	N88218	001	Jun 24, 1983
MUTUAL PHARM	100MG	N88994	001	Dec 04, 1985
SANDOZ	100MG	N87774	001	Jun 16, 1982
WATSON LABS	100MG	N87756	001	Dec 17, 1982

TABLET; ORAL

AZOLID

SANOFI AVENTIS US	100MG	N87091	001	
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BUTAZOLIDIN

NOVARTIS	100MG	N08319	008	
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PHENYLBUTAZONE

MUTUAL PHARM	100MG	N88863	001	Dec 04, 1985
SANDOZ	100MG	N84339	001	
WATSON LABS	100MG	N86151	001	
	100MG	N87674	001	Apr 21, 1982

PHENYLEPHRINE HYDROCHLORIDE; PROMETHAZINE HYDROCHLORIDE

SYRUP; ORAL

PHENERGAN VC

WYETH AYERST	5MG/5ML; 6.25MG/5ML	N08604	003	Apr 02, 1984
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PHERAZINE VC

HALSEY	5MG/5ML; 6.25MG/5ML	N88868	001	Mar 02, 1987
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PROMETHAZINE VC PLAIN

CENCI	5MG/5ML; 6.25MG/5ML	N88815	001	Nov 22, 1985
MORTON GROVE	5MG/5ML; 6.25MG/5ML	N88897	001	Jan 04, 1985

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 211 (of 274)

PHENYLEPHRINE HYDROCHLORIDE; PYRILAMINE MALEATE

SOLUTION/DROPS; OPHTHALMIC

PREFRIN-A

ALLERGAN

0.12%;0.1%

N07953 001

PHENYTOIN

SUSPENSION; ORAL

DILANTIN-30

PARKE DAVIS

30MG/5ML

N08762 002

PHENYTOIN SODIUM

CAPSULE; ORAL

DIPHENYLAN SODIUM

LANNETT

30MG PROMPT

N80857 001

100MG PROMPT

N80857 002

EXTENDED PHENYTOIN SODIUM

PLIVA

100MG EXTENDED

N89441 001

Dec 18, 1986

PHENYTEX

WATSON LABS

100MG EXTENDED

N88711 001

Dec 21, 1984

PHENYTOIN SODIUM

PHARMERAL

100MG PROMPT

N85435 001

WATSON LABS

100MG PROMPT

N85894 001

PROMPT PHENYTOIN SODIUM

WATSON LABS

100MG PROMPT

N80905 001

INJECTABLE; INJECTION

DILANTIN

PARKE DAVIS

50MG/ML

N10151 001

PHENYTOIN SODIUM

ABRAXIS PHARM

50MG/ML

N89003 001

May 31, 1985

MARSAM PHARMS LLC

50MG/ML

N89501 001

Oct 13, 1987

50MG/ML

N89779 001

Nov 27, 1992

SMITH AND NEPHEW

50MG/ML

N88519 001

Dec 19, 1984

50MG/ML

N88521 001

Dec 18, 1984

SOLOPAK

50MG/ML

N88520 001

Dec 17, 1984

WARNER CHILCOTT

50MG/ML

N89900 001

Mar 30, 1990

WATSON LABS

50MG/ML

N85434 001

PHYTONADIONE

INJECTABLE; INJECTION

AQUAMEPHYTON

ATON

1MG/0.5ML

N12223 002

10MG/ML

N12223 001

KONAKION

ROCHE

1MG/0.5ML

N11745 001

10MG/ML

N11745 003

PHYTONADIONE

GLAXOSMITHKLINE

1MG/0.5ML

N84060 001

10MG/ML

N84060 002

VITAMIN K1

HOSPIRA

10MG/ML

N87956 001

Jul 25, 1983

PILOCARPINE

INSERT, EXTENDED RELEASE; OPHTHALMIC

OCUSERT PILO-20

AKORN

5MG

N17431 001

OCUSERT PILO-40

AKORN

11MG

N17548 001

PINACIDIL

CAPSULE, EXTENDED RELEASE; ORAL

PINDAC

LEO PHARM

12.5MG

N19456 001

Dec 28, 1989

25MG

N19456 002

Dec 28, 1989

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 212 (of 274)

PINDOLOLTABLET; ORAL
PINDOLOL

PUREPAC PHARM

5MG
10MGN74125 001 Apr 28, 1993
N74125 002 Apr 28, 1993PIPECURONIUM BROMIDEINJECTABLE; INJECTION
ARDUAN

ORGANON USA INC

10MG/VIAL

N19638 001 Jun 26, 1990

PIPERACETAZINETABLET; ORAL
QUIDE

DOW PHARM

10MG
25MGN13615 001
N13615 002PIPERACILLIN SODIUMINJECTABLE; INJECTION
PIPRACIL

WYETH PHARMS INC

EQ 2GM BASE/VIAL
EQ 2GM BASE/VIAL
EQ 3GM BASE/VIAL
EQ 3GM BASE/VIAL
EQ 4GM BASE/VIAL
EQ 4GM BASE/VIAL
EQ 40GM BASE/VIALN50545 002
N62750 001 Oct 13, 1987
N50545 003
N62750 002 Oct 13, 1987
N50545 004
N62750 003 Oct 13, 1987
N50545 006 Sep 30, 1985PIPERAZINE CITRATESYRUP; ORAL
ANTEPAR

GLAXOSMITHKLINE

EQ 500MG BASE/5ML

N09102 001

BRYREL

SANOFI AVENTIS US

EQ 500MG BASE/5ML

N17796 001

MULTIFUGE

BLULINE

EQ 500MG BASE/5ML

N09452 001

PIPERAZINE CITRATE

ALPHARMA US PHARMS

EQ 500MG BASE/5ML

N80774 001

LANNETT

EQ 500MG BASE/5ML

N80963 001

LUITPOLD

EQ 500MG BASE/5ML

N80671 001

VERMIDOL

SOLVAY

EQ 500MG BASE/5ML

N80992 001

TABLET; ORAL
ANTEPAR

GLAXOSMITHKLINE

EQ 500MG BASE

N09102 003

PIPERAZINE CITRATE

IMPAX LABS

EQ 250MG BASE

N80874 001

PIPOBROMANTABLET; ORAL
VERCYTE

ABBOTT

10MG
25MGN16245 001
N16245 002PIRBUTEROL ACETATEAEROSOL, METERED; INHALATION
MAXAIR

3M

EQ 0.2MG BASE/INH

N19009 001 Dec 30, 1986

PIROXICAMCAPSULE; ORAL
PIROXICAM

IVAX PHARMS

10MG

N74148 001 Jun 03, 1996

DISCONTINUED DRUG PRODUCT LIST

6 - 213 (of 274)

PIROXICAM

CAPSULE; ORAL
PIROXICAM

IVAX PHARMS	20MG	N74148	002	Jun 03, 1996
ROXANE	10MG	N73651	001	Feb 26, 1993
	20MG	N73651	002	Feb 26, 1993
SCS	10MG	N74036	001	May 29, 1992
	20MG	N74036	002	May 29, 1992

POLYESTRADIOL PHOSPHATE

INJECTABLE; INJECTION
ESTRADURIN

WYETH AYERST	40MG/AMP	N10753	001	
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POLYETHYLENE GLYCOL 3350; POTASSIUM CHLORIDE; SODIUM BICARBONATE; SODIUM CHLORIDE; SODIUM SULFATE

SOLUTION; ORAL
OCL

HOSPIRA	6GM/100ML; 75MG/100ML; 168MG/100ML; 146MG/100ML; 1.29GM/100ML	N19284	001	Apr 30, 1986
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POLYETHYLENE GLYCOL 3350; POTASSIUM CHLORIDE; SODIUM BICARBONATE; SODIUM CHLORIDE; SODIUM SULFATE, ANHYDROUS

FOR SOLUTION; ORAL
COLYTE

SCHWARZ PHARMA	120GM/PACKET; 1.49GM/PACKET; 3.36GM/PACKET; 2.92GM/PACKET; 11.36GM/PACKET	N18983	005	Oct 26, 1984
	227.1GM/PACKET; 2.82GM/PACKET; 6.36GM/PACKET; 5.53GM/PACKET; 21.5GM/PACKET	N18983	004	Oct 26, 1984
	360GM/PACKET; 4.47GM/PACKET; 10.08GM/PACKET; 8.76GM/PACKET; 34.08GM/PACKET	N18983	006	Oct 26, 1984

FOR SUSPENSION; ORAL
COLOVAGE

DYNAPHARM	227.1GM/PACKET; 2.82GM/PACKET; 6.36GM/PACKET; 5.53GM/PACKET; 21.5GM/PACKET	N71320	001	Apr 20, 1988
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E-Z-EM PREP LYTE

E Z EM	236GM/BOT; 2.97GM/BOT; 6.74GM/BOT; 5.86GM/BOT; 22.74GM/BOT	N71278	001	Nov 21, 1988
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GLYCOPREP

GOLDLINE	236GM/BOT; 2.97GM/BOT; 6.74GM/BOT; 5.86GM/BOT; 22.74GM/BOT	N72319	001	Dec 23, 1988
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PEG-LYTE

SANDOZ	236GM/BOT; 2.97GM/BOT; 6.74GM/BOT; 5.86GM/BOT; 22.74GM/BOT	N73098	001	Aug 31, 1993
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POLYMYXIN B SULFATE

INJECTABLE; INJECTION
AEROSPORIN

GLAXOSMITHKLINE	EQ 500,000 U BASE/VIAL	N62036	001	
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POWDER; FOR RX COMPOUNDING

POLYMYXIN B SULFATE

PADDOCK	100,000,000 UNITS/BOT	N62455	001	Jul 27, 1983
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POTASSIUM AMINOSALICYLATE

POWDER; ORAL

POTASSIUM AMINOSALICYLATE

HEXCEL	100%	N80098	001	
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POTASSIUM CHLORIDE

CAPSULE, EXTENDED RELEASE; ORAL
K-LEASE

SAVAGE LABS	8MEQ	N73398	001	Jan 28, 1992
	10MEQ	N72427	001	Mar 28, 1990

FOR SUSPENSION, EXTENDED RELEASE; ORAL

MICRO-K LS

KV PHARM	20MEQ/PACKET	N19561	003	Aug 26, 1988
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27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 214 (of 274)

POTASSIUM CHLORIDE

INJECTABLE; INJECTION

POTASSIUM CHLORIDE

ABRAXIS PHARM

2MEQ/ML

N80204 001

2MEQ/ML

N84290 001

2MEQ/ML

N86713 001

2MEQ/ML

N86714 001

2MEQ/ML

N87787 001

Apr 20, 1982

2MEQ/ML

N87817 001

Oct 20, 1982

2MEQ/ML

N87885 001

Feb 03, 1983

AKORN

2MEQ/ML

N88286 001

Sep 05, 1985

BAXTER HLTHCARE

2MEQ/ML

N80203 001

GD SEARLE LLC

1MEQ/ML

N86219 001

2MEQ/ML

N86219 002

2MEQ/ML

N86220 002

3MEQ/ML

N86219 003

3MEQ/ML

N86220 001

4MEQ/ML

N86219 004

HOSPIRA

1MEQ/ML

N80205 003

1MEQ/ML

N83345 003

2.4MEQ/ML

N80205 004

3.2MEQ/ML

N80205 005

LILLY

2MEQ/ML

N07865 002

LUITPOLD

2MEQ/ML

N80221 001

2MEQ/ML

N80736 001

MILES

1MEQ/ML

N80195 002

2MEQ/ML

N80195 001

3MEQ/ML

N80195 003

4MEQ/ML

N80195 004

PHARMA SERVE NY

2MEQ/ML

N86297 001

2MEQ/ML

N87362 001

Mar 08, 1983

WATSON LABS

2MEQ/ML

N86208 001

2MEQ/ML

N89163 001

Mar 10, 1988

2MEQ/ML

N89421 001

Jan 02, 1987

3MEQ/ML

N86210 001

TABLET, EXTENDED RELEASE; ORAL

KAON CL

SAVAGE LABS

6.7MEQ

N17046 001

SLOW-K

NOVARTIS

8MEQ

N17476 002

TEN-K

NOVARTIS

10MEQ

N19381 001

Apr 16, 1986

POTASSIUM CHLORIDE; SODIUM CHLORIDE

INJECTABLE; INJECTION

POTASSIUM CHLORIDE 0.037% IN SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER

B BRAUN

37MG/100ML;900MG/100ML

N19708 001

Sep 29, 1989

POTASSIUM CHLORIDE 0.075% IN SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER

B BRAUN

75MG/100ML;900MG/100ML

N19708 002

Sep 29, 1989

POTASSIUM CHLORIDE 0.11% IN SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER

B BRAUN

110MG/100ML;900MG/100ML

N19708 003

Sep 29, 1989

POTASSIUM CHLORIDE 0.22% IN SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER

B BRAUN

220MG/100ML;900MG/100ML

N19708 005

Sep 29, 1989

POTASSIUM CHLORIDE 0.3% IN SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER

B BRAUN

300MG/100ML;900MG/100ML

N19708 006

Sep 29, 1989

SODIUM CHLORIDE 0.9% AND POTASSIUM CHLORIDE 0.075% IN PLASTIC CONTAINER

B BRAUN

75MG/100ML;900MG/100ML

N18722 001

Nov 09, 1982

BAXTER HLTHCARE

75MG/100ML;900MG/100ML

N17648 004

SODIUM CHLORIDE 0.9% AND POTASSIUM CHLORIDE 0.15% IN PLASTIC CONTAINER

B BRAUN

150MG/100ML;900MG/100ML

N18722 002

Nov 09, 1982

SODIUM CHLORIDE 0.9% AND POTASSIUM CHLORIDE 0.22% IN PLASTIC CONTAINER

B BRAUN

220MG/100ML;900MG/100ML

N18722 003

Nov 09, 1982

SODIUM CHLORIDE 0.9% AND POTASSIUM CHLORIDE 0.3% IN PLASTIC CONTAINER

B BRAUN

300MG/100ML;900MG/100ML

N18722 004

Nov 09, 1982

DISCONTINUED DRUG PRODUCT LIST

6 - 215 (of 274)

POTASSIUM CHLORIDE; SODIUM CHLORIDE; TROMETHAMINE

INJECTABLE; INJECTION

THAM-E

HOSPIRA

370MG/VIAL;1.75GM/VIAL;36GM/VIAL

N13025 001

POTASSIUM CITRATE

FOR SOLUTION; ORAL

POTASSIUM CITRATE

MISSION PHARMA

10MEQ/PACKET

N19647 002

Oct 13, 1988

20MEQ/PACKET

N19647 001

Oct 13, 1988

POTASSIUM IODIDE

SOLUTION; ORAL

POTASSIUM IODIDE

ROXANE

1GM/ML

N18551 001

Feb 19, 1982

TABLET; ORAL

THYRO-BLOCK

MEDPOINTE PHARM HLC

130MG

N18307 001

POTASSIUM PERCHLORATE

CAPSULE; ORAL

PERCHLORACAP

MALLINCKRODT

200MG

N17551 001

POVIDONE- IODINE

SOLUTION; TOPICAL

E-Z PREP

CLINIPAD

10%

N19382 001

Jul 25, 1989

SPONGE; TOPICAL

E-Z PREP

CLINIPAD

5%

N19382 002

Jul 25, 1989

E-Z PREP 220

CLINIPAD

5%

N19382 003

Jul 25, 1989

PRALIDOXIME CHLORIDE

INJECTABLE; INJECTION

PRALIDOXIME CHLORIDE

BAXTER HLTHCARE CORP

300MG/ML

N18799 001

Dec 13, 1982

TABLET; ORAL

PROTOPAM CHLORIDE

WYETH AYERST

500MG

N14122 002

PRAMIPEXOLE DIHYDROCHLORIDE

TABLET; ORAL

MIRAPEX

BOEHRINGER INGELHEIM

1.25MG

N20667 004

Jul 01, 1997

PRAZEPAM

CAPSULE; ORAL

CENTRAX

PARKE DAVIS

5MG

N18144 001

10MG

N18144 002

20MG

N18144 003

May 10, 1982

PRAZEPAM

USL PHARMA

5MG

N70427 001

Nov 06, 1987

10MG

N70428 001

Nov 06, 1987

TABLET; ORAL

CENTRAX

PARKE DAVIS

10MG

N17415 001

PRAZOSIN HYDROCHLORIDE

CAPSULE; ORAL

PRAZOSIN HYDROCHLORIDE

AM THERAP

EQ 1MG BASE

N72782 001

May 16, 1989

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 216 (of 274)

PRAZOSIN HYDROCHLORIDE

CAPSULE; ORAL

PRAZOSIN HYDROCHLORIDE

AM THERAP

EQ 2MG BASE

N72783 001 May 16, 1989

EQ 5MG BASE

N72784 001 May 16, 1989

CLONMEL HLTHCARE

EQ 1MG BASE

N72705 001 May 16, 1989

EQ 2MG BASE

N72706 001 May 16, 1989

EQ 5MG BASE

N72707 001 May 16, 1989

PUREPAC PHARM

EQ 1MG BASE

N72991 001 May 16, 1989

EQ 2MG BASE

N72921 001 May 16, 1989

EQ 5MG BASE

N72992 001 May 16, 1989

PREDNISOLONE

CREAM; TOPICAL

METI-DERM

SCHERING

0.5%

N10209 002

SYRUP; ORAL

PRELONE

MURO

5MG/5ML

N89654 001 Jan 17, 1989

TABLET; ORAL

CORTALONE

HALSEY

1MG

N80304 003

2.5MG

N80304 002

5MG

N80304 001

DELTA-CORTEF

PHARMACIA AND UPJOHN

5MG

N09987 004

FERNISOLONE-P

FERNDAL LABS

5MG

N83941 001

PREDNISOLONE

BARR

5MG

N84426 002

BUNDY

5MG

N83675 001

ELKINS SINN

5MG

N80625 001

EVERYLIFE

1MG

N84439 001

2.5MG

N84439 002

5MG

N84439 003

FERRANTE

2.5MG

N80562 001

5MG

N80562 002

HEATHER

5MG

N80326 001

IMPAX LABS

5MG

N80780 001

INWOOD LABS

5MG

N80748 001

IVAX PHARMS

5MG

N80378 001

LEINER

5MG

N80211 001

MARSHALL PHARMA

5MG

N80307 001

PANRAY

1MG

N80351 001

5MG

N80351 002

PERRIGO

5MG

N84542 001

PHOENIX LABS NY

5MG

N80322 001

PUREPAC PHARM

5MG

N80325 001

ROXANE

5MG

N80327 002

SANDOZ

5MG

N80339 001

5MG

N84773 001

SPERTI

1MG

N80358 001

2.5MG

N80358 002

5MG

N80358 003

SUPERPHARM

5MG

N88892 001 Feb 26, 1985

TABLICAPS

5MG

N85170 001

TEVA

5MG

N80398 001

UDL

5MG

N87987 001 Jan 18, 1983

VALEANT PHARM INTL

5MG

N80236 001

VITARINE

5MG

N80534 001

WATSON LABS

5MG

N85085 002

5MG

N85415 001

5MG

N85416 001

WEST WARD

5MG

N80324 001

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 217 (of 274)

PREDNISOLONETABLET; ORAL
PREDNISOLONE

WHITEWORTH TOWN PLSN 5MG

N80342 001

STERANE

PFIZER 5MG

N09996 001

PREDNISOLONE ACETATE

INJECTABLE; INJECTION

METICORTELONE

SCHERING 25MG/ML

N10255 002

PREDNISOLONE ACETATE

AKORN 25MG/ML

N83032 001

50MG/ML

N84492 001

BEL MAR 25MG/ML

N83738 001

50MG/ML

N83738 002

CENT PHARMS 25MG/ML

N84717 001

50MG/ML

N84717 002

WATSON LABS 25MG/ML

N83398 001

25MG/ML

N83654 001

40MG/ML

N83767 001

50MG/ML

N83764 001

50MG/ML

N85781 001

STERANE

PFIZER 25MG/ML

N11446 001

SUSPENSION/DROPS; OPHTHALMIC

ECONOPRED

ALCON 0.125%

N17468 001

PREDNISOLONE ACETATE; SULFACETAMIDE SODIUM

OINTMENT; OPHTHALMIC

CETAPRED

ALCON 0.25%;10%

N87771 001 Aug 06, 1993

METIMYD

SCHERING 0.5%;10%

N10210 002 Sep 09, 1984

PREDSULFAIR

PHARMAFAIR 0.5%;10%

N88032 001 Apr 15, 1983

VASOCIDIN

NOVARTIS 0.5%;10%

N88791 001 Oct 05, 1984

SUSPENSION; OPHTHALMIC

ISOPTO CETAPRED

ALCON 0.25%;10%

N87547 001

SUSPENSION/DROPS; OPHTHALMIC

METIMYD

SCHERING 0.5%;10%

N10210 001

PREDAMIDE

AKORN 0.5%;10%

N88059 001 Jul 29, 1983

PREDSULFAIR

PHARMAFAIR 0.5%;10%

N88007 001 Apr 19, 1983

PREDSULFAIR II

PHARMAFAIR 0.2%;10%

N88837 001 Dec 24, 1985

SULPHRIN

BAUSCH AND LOMB 0.5%;10%

N88089 001 Dec 28, 1982

PREDNISOLONE SODIUM PHOSPHATE

INJECTABLE; INJECTION

HYDELTRASOL

MERCK EQ 20MG PHOSPHATE/ML

N11583 002

PREDNISOLONE SODIUM PHOSPHATE

WATSON LABS EQ 20MG PHOSPHATE/ML

N80517 001

OINTMENT; OPHTHALMIC, OTIC

HYDELTRASOL

MERCK EQ 0.25% PHOSPHATE

N11028 001

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 218 (of 274)

PREDNISOLONE SODIUM PHOSPHATE

SOLUTION/DROPS; OPHTHALMIC

METRETON

SCHERING

EQ 0.5% PHOSPHATE

N83834 001

PREDAIR

PHARMAFAIR

EQ 0.11% PHOSPHATE

N88415 001

Feb 29, 1984

PREDAIR FORTE

PHARMAFAIR

EQ 0.9% PHOSPHATE

N88165 001

Mar 28, 1983

PREDNISOLONE SODIUM PHOSPHATE

AKORN

EQ 0.11% PHOSPHATE

N83358 001

EQ 0.9% PHOSPHATE

N83358 002

ALCON UNIVERSAL

EQ 0.11% PHOSPHATE

N81043 001

Oct 24, 1991

EQ 0.9% PHOSPHATE

N81044 001

Oct 24, 1991

SOLA BARNES HIND

EQ 0.11% PHOSPHATE

N84171 001

EQ 0.9% PHOSPHATE

N84168 001

EQ 0.9% PHOSPHATE

N84169 001

EQ 0.9% PHOSPHATE

N84172 001

PREDNISOLONE SODIUM PHOSPHATE; SULFACETAMIDE SODIUM

SOLUTION/DROPS; OPHTHALMIC

SULSTER

AKORN

EQ 0.23% PHOSPHATE;10%

N74511 001

Jul 30, 1996

PREDNISOLONE TEBUTATE

INJECTABLE; INJECTION

HYDELTRA-TBA

MERCK

20MG/ML

N10562 001

PREDNISOLONE TEBUTATE

WATSON LABS

20MG/ML

N83362 001

Feb 17, 1984

PREDNISONE

SOLUTION; ORAL

PREDNISONE

MORTON GROVE

5MG/5ML

N89726 001

Aug 02, 1988

SYRUP; ORAL

LIQUID PRED

MURO

5MG/5ML

N87611 002

Sep 07, 1982

TABLET; ORAL

CORTAN

HALSEY

20MG

N87480 001

DELTA-DOME

BAYER PHARMS

5MG

N80293 001

DELTASONE

PHARMACIA AND UPJOHN

2.5MG

N09986 005

5MG

N09986 002

10MG

N09986 006

20MG

N09986 007

50MG

N09986 008

FERNISONE

FERNDAL LABS

5MG

N83364 001

METICORTEN

SCHERING

1MG

N09766 002

5MG

N09766 001

ORASONE

SOLVAY

1MG

N83009 001

5MG

N83009 002

10MG

N83009 003

20MG

N83009 004

50MG

N85999 001

PARACORT

PARKE DAVIS

5MG

N10962 002

PREDNICEN-M

SCHWARZ PHARMA

5MG

N84655 001

DISCONTINUED DRUG PRODUCT LIST

6 - 219 (of 274)

PREDNISONE

TABLET; ORAL
PREDNISONE

AM THERAP	5MG	N89387	001	Nov 06, 1986
	10MG	N89388	001	Nov 06, 1986
	20MG	N89389	001	Nov 06, 1986
BUNDY	5MG	N83676	001	
DURAMED PHARMS BARR	5MG	N88394	001	Oct 04, 1983
	10MG	N88395	001	Oct 04, 1983
	20MG	N88396	001	Oct 04, 1983
ELKINS SINN	5MG	N80491	001	
	20MG	N85811	001	
EVERYLIFE	1MG	N84440	001	
	2.5MG	N84440	002	
	5MG	N84440	003	
FERRANTE	2.5MG	N80563	001	
	5MG	N80563	002	
HALSEY	5MG	N80300	001	
HEATHER	5MG	N80320	001	
	10MG	N84341	001	
	20MG	N84417	001	
	20MG	N85543	001	
	50MG	N86946	001	
IMPAX LABS	5MG	N80782	001	
INTERPHARM	5MG	N89597	001	Oct 05, 1987
	10MG	N89598	001	Oct 05, 1987
	20MG	N89599	001	Oct 05, 1987
INWOOD LABS	1MG	N80328	001	
	2.5MG	N80306	001	
	5MG	N80279	001	
IVAX PHARMS	5MG	N80283	001	
	10MG	N84133	001	
	20MG	N84134	001	
KV PHARM	5MG	N84236	001	
LANNETT	5MG	N80514	001	
	20MG	N84275	001	
LEDERLE	5MG	N86968	001	
LEINER	20MG	N85151	001	
MARSHALL PHARMA	5MG	N80301	001	
MUTUAL PHARM	5MG	N80701	001	
	10MG	N86595	001	
	20MG	N84634	001	
	50MG	N86596	001	
NYLOS	5MG	N85115	001	
PANRAY	1MG	N80350	001	
	2.5MG	N80350	002	
	5MG	N80350	003	
PERRIGO	5MG	N83059	001	
PHARMAVITE	5MG	N84662	002	
PHOENIX LABS NY	5MG	N80321	001	
	20MG	N83807	001	
PUREPAC PHARM	5MG	N80353	001	
	10MG	N86062	001	
	20MG	N86061	001	
REXALL	5MG	N80232	001	
ROXANE	20MG	N17109	001	
	25MG	N87833	001	May 04, 1982
SANDOZ	5MG	N80336	002	
	5MG	N84774	001	
	20MG	N85813	001	
SCHERER LABS	5MG	N80371	001	
SPERTI	1MG	N80359	001	
	2.5MG	N80359	002	
	5MG	N80359	003	

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 220 (of 274)

PREDNISONETABLET; ORAL
PREDNISONE

SUPERPHARM	5MG	N88865	001	Oct 25, 1984
	10MG	N88866	001	Oct 25, 1984
	20MG	N88867	001	Oct 25, 1984
TEVA	5MG	N80397	001	
UDL	5MG	N87984	001	Jan 18, 1983
	10MG	N87985	001	Jan 18, 1983
	20MG	N87986	001	Jan 18, 1983
UPSHER SMITH	5MG	N87471	001	
	20MG	N87470	001	
VALEANT PHARM INTL	5MG	N80237	001	
VANGARD	5MG	N87682	001	Jan 15, 1982
	20MG	N87701	001	Jan 15, 1982
VITARINE	5MG	N80334	001	
	5MG	N80506	001	
WATSON LABS	5MG	N85084	002	
	50MG	N86867	001	
WEST WARD	50MG	N88465	001	Jun 01, 1984
WHITEWORTH TOWN PLSN	2.5MG	N84913	001	
	5MG	N80343	001	
	10MG	N89028	001	Jul 24, 1986
	20MG	N84913	002	
SERVISONE				
LEDERLE	5MG	N80223	001	

PRILOCAINE HYDROCHLORIDEINJECTABLE; INJECTION
CITANEST

ASTRAZENECA	1%	N14763	004	
	2%	N14763	005	
	3%	N14763	003	
CITANEST PLAIN				
ASTRAZENECA	4%	N14763	007	

PRIMIDONESUSPENSION; ORAL
MYSOLINE

XCEL PHARMS	250MG/5ML	N10401	001	
TABLET; ORAL PRIMIDONE				
WATSON LABS	250MG	N85052	001	

PROBENECIDTABLET; ORAL
BENEMID

MERCK	500MG	N07898	004	
PROBENECID				
LEDERLE	500MG	N86917	001	
WATSON LABS	500MG	N86150	002	Apr 23, 1982

PROBUCOLTABLET; ORAL
LORELCO

SANOFI AVENTIS US	250MG	N17535	001	
	500MG	N17535	002	Jul 06, 1988

PROCAINAMIDE HYDROCHLORIDE

CAPSULE; ORAL

PROCAINAMIDE HYDROCHLORIDE

ASCOT	250MG	N87542	001	Jan 08, 1982
	375MG	N87697	001	Mar 01, 1983
	500MG	N87543	001	Jan 08, 1982

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 221 (of 274)

PROCAINAMIDE HYDROCHLORIDE

CAPSULE; ORAL

PROCAINAMIDE HYDROCHLORIDE

LANNETT	250MG	N83693	001	
	500MG	N84696	001	
LEDERLE	250MG	N86942	001	
	375MG	N86952	001	
	500MG	N86943	001	
ROXANE	250MG	N88989	001	Apr 26, 1985
	500MG	N88990	001	Apr 26, 1985
SANDOZ	250MG	N89219	001	Jul 01, 1986
	375MG	N89220	001	Jul 01, 1986
	500MG	N89221	001	Jul 01, 1986
VANGARD	250MG	N87643	001	Jun 01, 1982
	500MG	N87875	001	Jun 01, 1982
WATSON LABS	250MG	N83795	001	
	250MG	N85167	001	
	375MG	N87020	001	
	500MG	N84357	001	
	500MG	N87021	001	
PROCAN				
PARKE DAVIS	250MG	N85804	001	
	375MG	N87502	001	
	500MG	N85079	001	
PROCAPAN				
PANRAY	250MG	N83553	002	

INJECTABLE; INJECTION

PROCAINAMIDE HYDROCHLORIDE

ABRAXIS PHARM	100MG/ML	N89415	001	Nov 17, 1986
	500MG/ML	N89416	001	Nov 17, 1986
BAXTER HLTHCARE	100MG/ML	N89029	001	Apr 17, 1986
	500MG/ML	N89030	001	Apr 17, 1986
PHARMAFAIR	100MG/ML	N88824	001	Nov 20, 1985
	500MG/ML	N88830	001	Nov 20, 1985
SMITH AND NEPHEW	100MG/ML	N88530	001	Mar 04, 1985
	500MG/ML	N88531	001	Mar 04, 1985
SOLOPAK	500MG/ML	N88532	001	Mar 04, 1985
WARNER CHILCOTT	100MG/ML	N89528	001	May 03, 1988
	500MG/ML	N89529	001	May 03, 1988
WATSON LABS	100MG/ML	N87079	001	
	500MG/ML	N87080	001	

TABLET, EXTENDED RELEASE; ORAL

PROCAINAMIDE HYDROCHLORIDE

COPLEY PHARM	500MG	N88974	001	Jul 22, 1985
	750MG	N89438	001	Mar 23, 1987
	1GM	N40111	001	Dec 13, 1996
INWOOD LABS	500MG	N89840	001	Mar 06, 1989
PLIVA	250MG	N88958	001	Dec 02, 1985
	500MG	N88959	001	Dec 02, 1985
SANDOZ	250MG	N89369	001	Aug 14, 1987
	500MG	N89284	001	Jun 23, 1986
	500MG	N89370	001	Jan 09, 1987
WATSON LABS	750MG	N89371	001	Aug 14, 1987
	250MG	N88533	001	Dec 03, 1984
	250MG	N89026	001	Oct 22, 1985
	500MG	N88534	001	Dec 03, 1984
	500MG	N89027	001	Oct 22, 1985
	750MG	N88535	001	Nov 03, 1984
	750MG	N89042	001	Oct 22, 1985
	1GM	N89520	001	Jan 15, 1987
PROCAN SR				
PARKE DAVIS	250MG	N86468	001	
PARKEDALE	500MG	N86065	001	
	750MG	N87510	001	Apr 01, 1982

DISCONTINUED DRUG PRODUCT LIST

6 - 222 (of 274)

PROCAINAMIDE HYDROCHLORIDE

TABLET, EXTENDED RELEASE; ORAL
 PROCAN SR
 PARKEDALE 1GM

N88489 001 Jan 16, 1985

PROCAINE HYDROCHLORIDE

INJECTABLE; INJECTION
 PROCAINE HYDROCHLORIDE

ABRAXIS PHARM 1%
 1%
 2%
 2%
 BEL MAR 1%
 2%
 ELKINS SINN 1%
 2%
 GD SEARLE LLC 1%
 2%
 MILES 1%
 2%
 WATSON LABS 1%
 2%
 2%

N80384 002
 N80421 001
 N80384 003
 N80421 002
 N80711 001
 N80756 001
 N83315 001
 N83315 002
 N86202 001
 N86202 002
 N80415 001
 N80415 002
 N83535 001
 N80658 002
 N83535 002

PROCAINE HYDROCHLORIDE; TETRACYCLINE HYDROCHLORIDE

INJECTABLE; INJECTION
 ACHROMYCIN

LEDERLE 40MG/VIAL;100MG/VIAL
 40MG/VIAL;250MG/VIAL

N50276 001
 N50276 003

TETRACYN

PFIZER 40MG/VIAL;100MG/VIAL
 40MG/VIAL;250MG/VIAL

N60285 002
 N60285 003

PROCAINE MERETHOXYLLINE; THEOPHYLLINE

INJECTABLE; INJECTION
 DICURIN PROCAINE

LILLY 100MG/ML;50MG/ML

N08869 001

PROCHLORPERAZINE

SUPPOSITORY; RECTAL
 COMPAZINE

GLAXOSMITHKLINE 2.5MG
 5MG

N11127 003
 N11127 001

PROCHLORPERAZINE

ABLE 2.5MG
 5MG
 25MG

N40407 001 Jul 11, 2001
 N40407 002 Jul 11, 2001
 N40407 003 Jul 11, 2001

PROCHLORPERAZINE EDISYLATE

CONCENTRATE; ORAL
 COMPAZINE

GLAXOSMITHKLINE EQ 10MG BASE/ML

N11276 001

PROCHLORPERAZINE

ALPHARMA US PHARMS EQ 10MG BASE/ML

N87153 001 Jun 08, 1982

PROCHLORPERAZINE EDISYLATE

MORTON GROVE EQ 10MG BASE/ML

N88598 001 Oct 25, 1984

INJECTABLE; INJECTION

PROCHLORPERAZINE EDISYLATE

BAXTER HLTHCARE EQ 5MG BASE/ML
 MARSAM PHARMS LLC EQ 5MG BASE/ML
 SMITH AND NEPHEW EQ 5MG BASE/ML
 WATSON LABS EQ 5MG BASE/ML
 EQ 5MG BASE/ML
 WYETH AYERST EQ 5MG BASE/ML

N89523 001 May 03, 1988
 N89675 001 Dec 05, 1988
 N89251 001 Dec 04, 1986
 N89605 001 Jul 08, 1987
 N89606 001 Jul 08, 1987
 N86348 001

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 223 (of 274)

PROCHLORPERAZINE EDISYLATE

SYRUP; ORAL

COMPAZINE

GLAXOSMITHKLINE EQ 5MG BASE/5ML

N11188 001

PROCHLORPERAZINE EDISYLATE

ALPHARMA US PHARMS EQ 5MG BASE/5ML

N87154 001 Sep 01, 1982

MORTON GROVE EQ 5MG BASE/5ML

N88597 001 Oct 25, 1984

PROCHLORPERAZINE MALEATE

CAPSULE, EXTENDED RELEASE; ORAL

COMPAZINE

GLAXOSMITHKLINE EQ 10MG BASE

N11000 001

EQ 10MG BASE

N21019 001 Oct 06, 1999

EQ 15MG BASE

N11000 002

EQ 30MG BASE

N11000 003

EQ 75MG BASE

N11000 004

TABLET; ORAL

PROCHLORPERAZINE

WATSON LABS EQ 5MG BASE

N85580 001

EQ 10MG BASE

N85178 001

EQ 25MG BASE

N85579 001

PROCHLORPERAZINE MALEATE

DURAMED PHARMS BARR EQ 5MG BASE

N89484 001 Jan 20, 1987

EQ 10MG BASE

N89485 001 Jan 20, 1987

EQ 25MG BASE

N89486 001 Jan 20, 1987

PROCYCLIDINE HYDROCHLORIDE

TABLET; ORAL

KEMADRIN

MONARCH PHARMS 2MG

N09818 005

PROGESTERONE

CAPSULE; ORAL

PROMETRIUM

UNIMED PHARMS 300MG

N19781 003 Oct 15, 1999

INJECTABLE; INJECTION

PROGESTERONE

LILLY 25MG/ML

N09238 002

50MG/ML

N09238 001

INSERT, EXTENDED RELEASE; INTRAUTERINE

PROGESTASERT

ALZA 38MG

N17553 001

PROMAZINE HYDROCHLORIDE

CONCENTRATE; ORAL

SPARINE

WYETH AYERST 30MG/ML

N10942 001

100MG/ML

N10942 004

INJECTABLE; INJECTION

PROMAZINE HYDROCHLORIDE

WATSON LABS 25MG/ML

N84510 001

50MG/ML

N84517 001

SPARINE

BAXTER HLTHCARE CORP 25MG/ML

N10349 008

50MG/ML

N10349 006

SYRUP; ORAL

SPARINE

WYETH AYERST 10MG/5ML

N10942 003

TABLET; ORAL

SPARINE

WYETH AYERST 10MG

N10348 006

25MG

N10348 001

50MG

N10348 002

100MG

N10348 003

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 224 (of 274)

PROMAZINE HYDROCHLORIDETABLET; ORAL
SPARINE

WYETH AYERST	200MG	N10348	004
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PROMETHAZINE HYDROCHLORIDEINJECTABLE; INJECTION
PHENERGAN

WYETH AYERST	25MG/ML	N08857	002
	50MG/ML	N08857	003

PROMETHAZINE HYDROCHLORIDE

ABBOTT	25MG/ML	N84223	001
	50MG/ML	N84222	001
AKORN	25MG/ML	N83955	002
	50MG/ML	N83955	001
MARSAM PHARMS LLC	25MG/ML	N89463	001
	50MG/ML	N89477	001
WATSON LABS	25MG/ML	N83532	001
	50MG/ML	N83532	002

May 02, 1988

May 02, 1988

ZIPAN-25

ALTANA	25MG/ML	N83997	001
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ZIPAN-50

ALTANA	50MG/ML	N83997	002
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SUPPOSITORY; RECTALPHENERGAN

WYETH PHARMS INC	12.5MG	N10926	002
	25MG	N10926	001
	50MG	N11689	001

PROMETHACON

POLYMEDICA	50MG	N84902	001
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PROMETHAZINE HYDROCHLORIDE

ABLE	12.5MG	N40504	001	Apr 11, 2003
	25MG	N40504	002	Apr 11, 2003
	50MG	N40449	001	Feb 27, 2003

SYRUP; ORALMYMETHAZINE FORTIS

USL PHARMA	25MG/5ML	N87996	001	Jan 18, 1983
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PHENERGAN FORTIS

WYETH AYERST	25MG/5ML	N08381	003
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PHENERGAN PLAIN

WYETH AYERST	6.25MG/5ML	N08381	004	Apr 18, 1984
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PROMETH FORTIS

ALPHARMA US PHARMS	25MG/5ML	N84772	001
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PROMETH PLAIN

ACTAVIS MID ATLANTIC	6.25MG/5ML	N85953	001
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PROMETHAZINE

CENCI	6.25MG/5ML	N89013	001	Sep 20, 1985
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PROMETHAZINE HYDROCHLORIDE

KV PHARM	6.25MG/5ML	N85388	001
	25MG/5ML	N85385	001
PHARM ASSOC	6.25MG/5ML	N87518	001
WHITEWORTH TOWN PLSN	6.25MG/5ML	N86395	001

TABLET; ORALPHENERGAN

WYETH PHARMS INC	12.5MG	N07935	002
	50MG	N07935	004

PROMETHAZINE HYDROCHLORIDE

ABBOTT	12.5MG	N84160	001
	25MG	N84166	001
	50MG	N84539	001

ABLE	12.5MG	N40558	001	Jul 01, 2004
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	25MG	N40558	002	Jul 01, 2004
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	50MG	N40558	003	Jul 01, 2004
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IMPAX LABS	25MG	N84214	002	Jul 07, 1982
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27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 225 (of 274)

PROMETHAZINE HYDROCHLORIDE

TABLET; ORAL

PROMETHAZINE HYDROCHLORIDE

IVAX PHARMS	12.5MG	N83604	001	Sep 10, 1985
	25MG	N83603	001	
	50MG	N83613	001	
LANNETT	12.5MG	N80949	001	
	25MG	N80949	002	
	50MG	N80949	003	
LEINER	12.5MG	N83214	001	
	25MG	N83658	001	
MUTUAL PHARM	12.5MG	N84555	001	
	25MG	N84554	001	
	50MG	N84557	001	
SANDOZ	12.5MG	N84233	001	
	25MG	N85146	001	
	50MG	N85146	002	
TABLICAPS	12.5MG	N84080	001	
	25MG	N84027	001	
TEVA	25MG	N89109	001	
WATSON LABS	12.5MG	N83401	001	
	12.5MG	N83712	001	
	12.5MG	N85986	001	
	25MG	N83204	001	
	25MG	N85684	001	
	50MG	N83403	001	
	50MG	N85664	001	
REMSED				
BRISTOL MYERS SQUIBB	25MG	N83176	002	
	50MG	N83176	001	

PROPANTHELINE BROMIDE

INJECTABLE; INJECTION

PRO-BANTHINE

GD SEARLE LLC	30MG/VIAL	N08843	001	
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TABLET; ORAL

PROPANTHELINE BROMIDE

ASCOT	15MG	N87663	001	Oct 25, 1982
HEATHER	15MG	N85780	001	
IMPAX LABS	15MG	N84541	002	Dec 08, 1983
LEINER	15MG	N80977	001	
MYLAN	15MG	N83706	001	
PAR PHARM	15MG	N88377	001	
SANDOZ	15MG	N80928	001	
TABLICAPS	15MG	N84428	001	
WATSON LABS	15MG	N83029	002	
	15MG	N83151	001	

PROPARACAINE HYDROCHLORIDE

SOLUTION/DROPS; OPHTHALMIC

KAINAIR

PHARMAFAIR	0.5%	N88087	001	Jun 07, 1983
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PROPARACAINE HYDROCHLORIDE

SOLA BARNES HIND	0.5%	N84144	001	
	0.5%	N84151	001	

PROPIOLACTONE

SOLUTION; IRRIGATION

BETAPRONE

FOREST LABS	N/A	N11657	001	
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27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 226 (of 274)

PROPIOMAZINE HYDROCHLORIDE

INJECTABLE; INJECTION

LARGON

BAXTER HLTHCARE CORP 20MG/ML

N12382 002

PROPOFOL

INJECTABLE; INJECTION

DIPRIVAN

ABRAXIS BIOSCIENCE 10MG/ML

N19627 001 Oct 02, 1989

PROPOXYPHENE HYDROCHLORIDE

CAPSULE; ORAL

DARVON

XANODYNE PHARM 32MG

N10997 001

DOLENE

RADIUS PHARMS 65MG

N80530 001

KESSO-GESIC

MK LABS 65MG

N83544 001

PROPHENE 65

HALSEY 65MG

N83538 002

PROPOXYPHENE HYDROCHLORIDE

ALRA 65MG

N83184 001

ANABOLIC 65MG

N83185 001

IMPAX LABS 65MG

N83317 001

IVAX PHARMS 32MG

N83597 001

LEINER 32MG

N83464 001

65MG

N83113 001

MUTUAL PHARM 65MG

N83186 001

MYLAN 32MG

N83528 001

65MG

N83299 001

PUREPAC PHARM 65MG

N83278 001

ROXANE 32MG

N83089 001

65MG

N83089 002

SANDOZ 32MG

N84014 001

65MG

N83125 002

65MG

N83688 001

65MG

N83870 002

65MG

N86495 001

VALEANT PHARM INTL 65MG

N80783 001

WATSON LABS 65MG

N80908 002

65MG

N85190 001

WHITEWORTH TOWN PLSN 65MG

N84551 001

PROPOXYPHENE HYDROCHLORIDE 65

WARNER CHILCOTT 65MG

N83786 001

PROPOXYPHENE NAPSYLATE

SUSPENSION; ORAL

DARVON-N

AAIPHARMA LLC 50MG/5ML

N16861 001

PROPRANOLOL HYDROCHLORIDE

CAPSULE, EXTENDED RELEASE; ORAL

PROPRANOLOL HYDROCHLORIDE

INWOOD LABS 60MG

N72499 001 Apr 11, 1989

80MG

N72500 001 Apr 11, 1989

120MG

N72501 001 Apr 11, 1989

160MG

N72502 001 Apr 11, 1989

INJECTABLE; INJECTION

PROPRANOLOL HYDROCHLORIDE

SMITH AND NEPHEW 1MG/ML

N70135 001 Apr 15, 1986

1MG/ML

N70137 001 Apr 15, 1986

SOLOPAK 1MG/ML

N70136 001 Apr 15, 1986

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 227 (of 274)

PROPRANOLOL HYDROCHLORIDE

SOLUTION; ORAL

PROPRANOLOL HYDROCHLORIDE

MORTON GROVE

20MG/5ML

N71984 001

Mar 03, 1989

40MG/5ML

N71985 001

Mar 03, 1989

SUSPENSION; ORAL

INDERAL

WYETH AYERST

10MG/ML

N19536 001

Dec 12, 1986

TABLET; ORAL

INDERAL

WYETH PHARMS INC

90MG

N16418 010

Oct 18, 1982

PROPRANOLOL HYDROCHLORIDE

CLONMEL HLTHCARE

10MG

N70125 001

Jul 30, 1985

20MG

N70126 001

Jul 30, 1985

40MG

N70127 001

Jul 30, 1985

60MG

N71495 001

Dec 31, 1987

80MG

N70128 001

Jul 30, 1985

90MG

N71496 001

Dec 31, 1987

DURAMED PHARMS BARR

10MG

N70306 001

Sep 09, 1985

20MG

N70307 001

Sep 09, 1985

40MG

N70308 001

Sep 09, 1985

60MG

N70309 001

Oct 01, 1986

80MG

N70310 001

Sep 09, 1985

90MG

N71327 001

Oct 01, 1986

INTERPHARM

10MG

N71368 001

May 05, 1987

20MG

N71369 001

May 05, 1987

40MG

N71370 001

May 05, 1987

80MG

N71371 001

May 05, 1987

IVAX PHARMS

10MG

N72063 001

Jul 29, 1988

20MG

N72066 001

Jul 29, 1988

40MG

N72067 001

Jul 29, 1988

60MG

N72068 001

Jul 29, 1988

80MG

N72069 001

Jul 29, 1988

LEDERLE

10MG

N72117 001

Jun 23, 1988

20MG

N72118 001

Jun 23, 1988

40MG

N72119 001

Jun 23, 1988

80MG

N72120 001

Jun 23, 1988

MUTUAL PHARM

10MG

N70319 001

Oct 22, 1985

20MG

N70320 001

Oct 22, 1985

40MG

N70103 001

Oct 22, 1985

60MG

N70321 001

Sep 24, 1986

80MG

N70322 001

Aug 04, 1986

MYLAN

60MG

N72275 001

Jun 09, 1989

PUREPAC PHARM

10MG

N70814 001

Nov 03, 1986

20MG

N70815 001

Nov 03, 1986

40MG

N70816 001

Nov 03, 1986

60MG

N70817 001

Nov 03, 1986

80MG

N70757 001

Nov 03, 1986

ROXANE

10MG

N70516 001

Jul 07, 1986

20MG

N70517 001

Jul 07, 1986

40MG

N70518 001

Jul 07, 1986

60MG

N70519 001

Sep 24, 1986

80MG

N70520 001

Jul 07, 1986

90MG

N70521 001

Sep 24, 1986

SANDOZ

10MG

N71658 001

Jul 05, 1988

20MG

N71687 001

Jul 05, 1988

40MG

N71688 001

Jul 05, 1988

60MG

N72197 001

Jul 05, 1988

80MG

N71689 001

Jul 05, 1988

90MG

N72198 001

Jul 05, 1988

SCHERING

10MG

N70120 001

Aug 06, 1985

20MG

N70121 001

Aug 06, 1985

40MG

N70122 001

Aug 06, 1985

60MG

N70123 001

Oct 29, 1986

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 228 (of 274)

PROPRANOLOL HYDROCHLORIDE

TABLET; ORAL

PROPRANOLOL HYDROCHLORIDE

SCHERING	80MG	N70124	001	Aug 06, 1985
SUPERPHARM	10MG	N71515	001	Jun 08, 1988
	20MG	N71516	001	Jun 08, 1988
	40MG	N71517	001	Jun 08, 1988
	80MG	N71518	001	Jun 08, 1988
TEVA	10MG	N70232	001	Oct 07, 1987
	20MG	N70233	001	Jun 23, 1986
WARNER CHILCOTT	10MG	N70438	001	Sep 15, 1986
	20MG	N70439	001	Sep 15, 1986
	40MG	N70440	001	Sep 15, 1986
	60MG	N70441	001	Sep 24, 1986
	80MG	N70442	001	Sep 15, 1986
WATSON LABS	10MG	N70140	001	Jul 30, 1985
	10MG	N70378	001	Mar 19, 1987
	20MG	N70141	001	Jul 30, 1985
	20MG	N70379	001	Mar 19, 1987
	20MG	N70549	001	Apr 11, 1986
	40MG	N70142	001	Jul 30, 1985
	40MG	N70380	001	Mar 19, 1987
	60MG	N70143	001	Jan 15, 1987
	60MG	N70381	001	Mar 19, 1987
	60MG	N71098	001	Oct 06, 1986
	80MG	N70144	001	Jul 30, 1985
	80MG	N70382	001	Mar 19, 1987
	90MG	N71183	001	Oct 06, 1986

PROPYLIODONE

SUSPENSION; INTRATRACHEAL

DIONOSIL AQUEOUS

GLAXOSMITHKLINE	50%	N09309	001
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DIONOSIL OILY

GLAXOSMITHKLINE	60%	N09309	002
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PROPYLTHIOURACIL

TABLET; ORAL

PROPYLTHIOURACIL

ABBOTT	50MG	N84075	001
ANABOLIC	50MG	N80285	001
HALSEY	50MG	N80015	001
IMPAX LABS	50MG	N80159	001
IVAX PHARMS	50MG	N80215	001
LANNETT	50MG	N80016	001
LILLY	50MG	N06213	001
MUTUAL PHARM	50MG	N83982	001
PERRIGO	50MG	N84543	001
TABLICAPS	50MG	N80840	001
WATSON LABS	50MG	N80932	001
	50MG	N85201	001

PROTAMINE SULFATE

INJECTABLE; INJECTION

PROTAMINE SULFATE

BAXTER HLTHCARE	10MG/ML	N89474	001	Nov 05, 1986
	10MG/ML	N89475	001	Nov 05, 1986
LILLY	10MG/ML	N06460	002	
PHARMACIA AND UPJOHN	50MG/VIAL	N07413	001	
	250MG/VIAL	N07413	002	Aug 02, 1984

DISCONTINUED DRUG PRODUCT LIST

6 - 229 (of 274)

PROTEIN HYDROLYSATE

INJECTABLE; INJECTION

AMINOSOL 5%

ABBOTT

5%

N05932 012

Jan 31, 1985

HYPROTIGEN 5%

B BRAUN

5%

N06170 003

Jan 10, 1984

PROTIRELIN

INJECTABLE; INJECTION

THYPINONE

ABBOTT

0.5MG/ML

N17638 001

THYREL TRH

FERRING

0.5MG/ML

N18087 001

PROTOKYLOL HYDROCHLORIDE

TABLET; ORAL

VENTAIRE

SANOFI AVENTIS US

2MG

N83459 001

PROTRIPTYLINE HYDROCHLORIDE

TABLET; ORAL

VIVACTIL

ODYSSEY PHARMS

5MG

N16012 001

10MG

N16012 002

PSEUDOEPHEDRINE HYDROCHLORIDE

CAPSULE, EXTENDED RELEASE; ORAL

NOVAFED

SANOFI AVENTIS US

120MG

N17603 001

SUDAFED 12 HOUR

GLAXOSMITHKLINE

120MG **Federal Register determination
that product was not discontinued or
withdrawn for safety or efficacy
reasons**

N17941 002

PSEUDOEPHEDRINE HYDROCHLORIDE; TRIPROLIDINE HYDROCHLORIDE

CAPSULE, EXTENDED RELEASE; ORAL

ACTIFED

GLAXOSMITHKLINE

120MG;5MG

N18996 001

Jun 17, 1985

TRIPROLIDINE AND PSEUDOEPHEDRINE HYDROCHLORIDES

KV PHARM

120MG;5MG

N71798 001

Mar 16, 1989

SYRUP; ORAL

ACTAHIST

CENCI

30MG/5ML;1.25MG/5ML

N88344 001

Feb 09, 1984

HISTAFED

CENCI

30MG/5ML;1.25MG/5ML

N88283 001

Apr 20, 1984

MYFED

USL PHARMA

30MG/5ML;1.25MG/5ML

N88116 001

Mar 04, 1983

TRILITRON

NEWTRON PHARMS

30MG/5ML;1.25MG/5ML

N88474 001

Feb 12, 1985

TABLET; ORAL

ALLERFED

LEINER

60MG;2.5MG

N88860 001

Jan 31, 1985

TRILITRON

NEWTRON PHARMS

60MG;2.5MG

N88515 001

Jan 09, 1985

TRIPHED

TEVA

60MG;2.5MG

N88630 001

May 17, 1984

TRIPROLIDINE AND PSEUDOEPHEDRINE

WATSON LABS

60MG;2.5MG

N88318 002

Jan 13, 1984

WEST WARD

60MG;2.5MG

N88117 001

Apr 19, 1983

TRIPROLIDINE HYDROCHLORIDE AND PSEUDOEPHEDRINE HYDROCHLORIDE

IVAX PHARMS

60MG;2.5MG

N85273 001

Dec 12, 1984

SUPERPHARM

60MG;2.5MG

N88578 001

Feb 21, 1985

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 230 (of 274)

PSEUDOEPHEDRINE HYDROCHLORIDE; TRIPROLIDINE HYDROCHLORIDE

TABLET, EXTENDED RELEASE; ORAL				
TRIPROLIDINE AND PSEUDOEPHEDRINE HYDROCHLORIDES				
KV PHARM	120MG;5MG	N72758	001	Nov 25, 1991

PSEUDOEPHEDRINE POLISTIREX

SUSPENSION, EXTENDED RELEASE; ORAL				
PSEUDO-12				
UCB INC	EQ 60MG HCL/5ML	N19401	001	Jun 19, 1987

PYRIDOSTIGMINE BROMIDE

TABLET; ORAL				
PYRIDOSTIGMINE BROMIDE				
SOLVAY	30MG	N89572	001	Nov 27, 1990
US ARMY	30MG	N20414	001	Feb 05, 2003

PYRIDOXINE HYDROCHLORIDE

INJECTABLE; INJECTION				
HEXA-BETALIN				
LILLY	100MG/ML	N80854	001	
PYRIDOXINE HYDROCHLORIDE				
AKORN	100MG/ML	N87967	001	Oct 01, 1982
BEL MAR	100MG/ML	N80761	001	
DELL LABS	50MG/ML	N83771	001	
	100MG/ML	N83772	001	
ELKINS SINN	100MG/ML	N80581	001	
LUITPOLD	100MG/ML	N80669	001	
WATSON LABS	100MG/ML	N83760	001	

PYRILAMINE MALEATE

TABLET; ORAL				
PYRILAMINE MALEATE				
IMPAX LABS	25MG	N80808	001	
WATSON LABS	25MG	N85231	001	

PYRITHIONE ZINC

LOTION; TOPICAL				
HEAD & SHOULDERS CONDITIONER				
PROCTER AND GAMBLE	0.3%	N19412	002	Mar 10, 1986

PYRVINIUM PAMOATE

SUSPENSION; ORAL				
POVAN				
PARKE DAVIS	EQ 50MG BASE/5ML	N11964	001	
TABLET; ORAL				
POVAN				
PARKE DAVIS	EQ 50MG BASE	N12485	002	

QUAZEPAM

TABLET; ORAL				
DORAL				
QUESTCOR PHARMS	7.5MG	N18708	003	Feb 26, 1987

QUETIAPINE FUMARATE

TABLET; ORAL				
SEROQUEL				
ASTRAZENECA	EQ 150MG BASE	N20639	004	Dec 20, 1998

QUINESTROL

TABLET; ORAL				
ESTROVIS				
PARKE DAVIS	0.1MG	N16768	002	
	0.2MG	N16768	003	

DISCONTINUED DRUG PRODUCT LIST

6 - 231 (of 274)

QUINETHAZONE

TABLET; ORAL HYDROMOX LEDERLE	50MG	N13264	001
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QUINETHAZONE; RESERPINE

TABLET; ORAL HYDROMOX R LEDERLE	50MG;0.125MG	N13927	001
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QUINIDINE GLUCONATE

TABLET; ORAL QUINACT BERLEX	266MG 400MG	N85978 N86099	001 001
TABLET, EXTENDED RELEASE; ORAL			
DURAQUIN			
WARNER CHILCOTT	330MG	N17917	001
QUINAGLUTE			
BERLEX	324MG	N16647	001
QUINALAN			
LANNETT	324MG	N88081	001 Feb 10, 1986
QUINATIME			
WATSON LABS	324MG	N87448	001
QUINIDINE GLUCONATE			
ASCOT	324MG	N88582	001 Jun 17, 1985
HALSEY	324MG	N89476	001 Apr 10, 1987
ROXANE	324MG	N88431	001 Jan 06, 1984
SANDOZ	324MG	N89894	001 Dec 15, 1988
SUPERPHARM	324MG	N89164	001 Nov 21, 1985
WATSON LABS	324MG	N87785	001 Jan 24, 1983

QUINIDINE POLYGALACTURONATE

TABLET; ORAL CARDIOQUIN PURDUE FREDERICK	275MG	N11642	002
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QUINIDINE SULFATE

CAPSULE; ORAL CIN-QUIN SOLVAY	200MG 300MG	N85296 N85297	001 001
QUINIDINE SULFATE			
LILLY	200MG	N85103	001
TABLET; ORAL			
CIN-QUIN			
SOLVAY	100MG	N85299	001
	200MG	N84932	001
	300MG	N85298	001
QUINIDINE SULFATE			
BARR	200MG	N84177	001
CLONMEL HLTHCARE	200MG	N87011	001
ELKINS SINN	200MG	N83622	001
EVERYLIFE	200MG	N83439	001
HALSEY	200MG	N83583	001
IMPAX LABS	200MG	N83347	001
IVAX PHARMS	200MG	N84549	001
KING PHARMS	200MG	N85175	001
KV PHARM	200MG	N85276	001
LANNETT	200MG	N83743	001
LEDERLE	200MG	N86176	001
LEINER	200MG	N83808	001
LILLY	200MG	N85038	001
MUTUAL PHARM	100MG	N81029	001 Apr 14, 1989

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 232 (of 274)

QUINIDINE SULFATE

TABLET; ORAL

QUINIDINE SULFATE

PERRIGO	200MG	N85322	001	
PHARMAVITE	200MG	N84627	001	
PUREPAC PHARM	200MG	N84003	001	
ROXANE	200MG	N83640	001	
	300MG	N85632	001	
SANDOZ	200MG	N84631	001	
	200MG	N84914	001	
	300MG	N89839	001	Sep 29, 1988
SCHERER LABS	200MG	N85068	001	
SUPERPHARM	200MG	N88973	001	Apr 10, 1985
USL PHARMA	200MG	N87837	001	Apr 14, 1982
VALEANT PHARM INTL	200MG	N83393	001	
VANGARD	200MG	N87909	001	Jul 13, 1982
VINTAGE PHARMS	200MG	N83963	001	
WARNER CHILCOTT	200MG	N83879	001	
WATSON LABS	100MG	N85584	001	
	200MG	N85140	002	
WEST WARD	200MG	N83862	001	
WHITEWORTH TOWN PLSN	200MG	N85444	001	
QUINORA				
KEY PHARMS	200MG	N83576	001	
SCHERING	300MG	N85222	001	
TABLET, EXTENDED RELEASE; ORAL				
QUINIDEX				
WYETH PHARMS INC	300MG	N12796	002	

RABEPRAZOLE SODIUM

TABLET, DELAYED RELEASE; ORAL

ACIPHEX

EISAI MEDCL RES	10MG **Federal Register determination that product was not discontinued or withdrawn for safety or efficacy reasons**	N20973	001	May 29, 2002
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RANITIDINE BISMUTH CITRATE

TABLET; ORAL

TRITEC

GLAXOSMITHKLINE	400MG	N20559	001	Aug 08, 1996
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RANITIDINE HYDROCHLORIDE

CAPSULE; ORAL

ZANTAC 150

GLAXOSMITHKLINE	EQ 150MG BASE	N20095	001	Mar 08, 1994
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ZANTAC 300

GLAXOSMITHKLINE	EQ 300MG BASE	N20095	002	Mar 08, 1994
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INJECTABLE; INJECTION

ZANTAC IN PLASTIC CONTAINER

GLAXOSMITHKLINE	EQ 50MG BASE/100ML	N19593	001	Dec 17, 1986
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TABLET; ORAL

RANITIDINE

RANBAXY	EQ 75MG BASE	N75254	001	Jan 14, 2000
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RANITIDINE HYDROCHLORIDE

BOEHRINGER INGELHEIM	EQ 150MG BASE	N74662	001	Aug 29, 1997
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	EQ 300MG BASE	N74662	002	Aug 29, 1997
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RANBAXY	EQ 150MG BASE	N75000	001	Jan 30, 1998
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	EQ 300MG BASE	N75000	002	Jan 30, 1998
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TABLET, EFFERVESCENT; ORAL

ZANTAC 75

MCNEIL CONS	EQ 75MG BASE **Federal Register determination that product was not discontinued or withdrawn for safety or efficacy reasons**	N20745	001	Feb 26, 1998
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27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 233 (of 274)

RAPACURONIUM BROMIDE

INJECTABLE; INJECTION

RAPLON

ORGANON USA INC

100MG/VIAL

N20984 001

Aug 18, 1999

200MG/VIAL

N20984 002

Aug 18, 1999

RAUWOLFIA SERPENTINA

TABLET; ORAL

HIWOLFIA

BOWMAN PHARMS

50MG

N09276 003

50MG

N09276 005

100MG

N09276 004

HYSERPIN

PHYS PRODS VA

50MG

N10581 001

KONGLUCOID

PANRAY

50MG

N09278 001

100MG

N09278 002

RAUDIXIN

APOTHECON

50MG

N08842 001

100MG

N08842 002

RAUSERPIN

FERNDAL LABS

50MG

N09926 002

100MG

N09926 004

RAUVAL

PAL PAK

50MG

N09108 002

100MG

N09108 004

RAUWOLFIA SERPENTINA

BUNDY

50MG

N09477 001

100MG

N09477 002

HALSEY

50MG

N80498 001

100MG

N80498 002

IMPAX LABS

50MG

N09273 001

100MG

N09273 002

IVAX PHARMS

50MG

N11521 001

100MG

N11521 002

LEINER

50MG

N80583 001

100MG

N80583 002

PUREPAC PHARM

50MG

N80842 001

100MG

N80842 002

SOLVAY

50MG

N80500 001

100MG

N80500 002

TABLICAPS

50MG

N83867 001

100MG

N83444 001

VALEANT PHARM INTL

50MG

N09668 001

100MG

N09668 002

WATSON LABS

50MG

N80907 001

100MG

N80914 001

WOLFINA

FOREST PHARMS

50MG

N09255 008

100MG

N09255 006

RESCINNAMINE

CAPSULE; ORAL

CINNASIL

PANRAY

0.5MG

N84736 001

RESERPINE

ELIXIR; ORAL

SERPASIL

NOVARTIS

0.2MG/4ML

N09115 005

INJECTABLE; INJECTION

SANDRIL

LILLY

2.5MG/ML

N10012 001

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 234 (of 274)

RESERPINE

INJECTABLE; INJECTION

SERPASIL

NOVARTIS

2.5MG/ML

N09434 002

TABLET; ORAL

HISERPIA

BOWMAN PHARMS

0.1MG

N09631 002

0.25MG

N09631 004

RAU-SED

BRISTOL MYERS SQUIBB

0.1MG

N09357 001

0.25MG

N09357 004

0.5MG

N09357 006

1MG

N09357 008

RESERPINE

BARR

0.25MG

N80721 002

BELL PHARMA

0.1MG

N83058 001

0.25MG

N83058 002

BUNDY

0.1MG

N09663 001

0.25MG

N09663 003

ELKINS SINN

0.1MG

N83145 001

0.25MG

N83145 002

EVERYLIFE

0.1MG

N10441 001

0.25MG

N10441 002

0.5MG

N10441 003

1MG

N10441 004

HALSEY

0.1MG

N80457 002

0.25MG

N80457 001

1MG

N80457 003

IMPAX LABS

0.1MG

N09627 001

0.25MG

N09627 002

IVAX PHARMS

0.1MG

N11185 001

0.25MG

N11185 002

LEINER

0.1MG

N86117 001

0.25MG

N80582 001

0.25MG

N85775 001

1MG

N80582 002

MARSHALL PHARMA

0.1MG

N80492 001

0.25MG

N80492 002

MK LABS

0.1MG

N80525 002

0.25MG

N80525 001

MYLAN

1MG

N84974 001

PHARMAVITE

0.25MG

N84663 001

PUREPAC PHARM

0.1MG

N80753 002

0.25MG

N80753 001

REXALL

0.25MG

N80637 001

ROXANE

0.1MG

N09859 001

0.25MG

N09859 002

SOLVAY

0.25MG

N80446 001

TABLICAPS

0.25MG

N85207 001

TEVA

0.1MG

N89020 001

Mar 07, 1985

0.25MG

N89019 001

Mar 07, 1985

VALEANT PHARM INTL

0.1MG

N09667 001

0.25MG

N09667 002

WATSON LABS

0.1MG

N80679 001

0.25MG

N80393 001

0.25MG

N85401 001

1MG

N80749 001

WEST WARD

0.1MG

N80975 001

0.25MG

N80975 002

1MG

N80975 003

WHITEWORTH TOWN PLSN

0.1MG

N80723 001

0.25MG

N80723 002

1MG

N80723 003

DISCONTINUED DRUG PRODUCT LIST

6 - 235 (of 274)

RESERPINE

TABLET; ORAL

SANDRIL

LILLY

0.1MG

N09376 004

0.25MG

N09376 001

SERPANRAY

PANRAY

0.1MG

N09391 001

0.25MG

N09391 002

1MG

N09391 004

SERPASIL

NOVARTIS

0.1MG

N09115 001

0.25MG

N09115 003

1MG

N09115 004

SERPATE

VALE

0.1MG

N09453 001

0.25MG

N09453 002

SERPIVITE

VITARINE

0.25MG

N09645 002

RESERPINE; TRICHLORMETHIAZIDE

TABLET; ORAL

METATENSIN #2

SANOFI AVENTIS US

0.1MG;2MG

N12972 001

METATENSIN #4

SANOFI AVENTIS US

0.1MG;4MG

N12972 002

NAQUIVAL

SCHERING

0.1MG;4MG

N12265 003

TRICHLORMETHIAZIDE W/ RESERPINE

WATSON LABS

0.1MG;4MG

N85248 001

RIBAVIRIN

TABLET; ORAL

COPEGUS

ROCHE

400MG

N21511 002

Jun 21, 2005

RISPERIDONE

TABLET; ORAL

RISPERDAL

JANSSEN PHARMA

5MG

N20272 005

Dec 29, 1993

RITODRINE HYDROCHLORIDE

INJECTABLE; INJECTION

RITODRINE HYDROCHLORIDE

ABRAXIS PHARM

10MG/ML

N71188 001

Jul 23, 1987

15MG/ML

N71189 001

Jul 23, 1987

YUTOPAR

ASTRAZENECA

10MG/ML

N18580 001

15MG/ML

N18580 002

TABLET; ORAL

YUTOPAR

ASTRAZENECA

10MG

N18555 001

RITONAVIR

CAPSULE; ORAL

NORVIR

ABBOTT

100MG

N20680 001

Mar 01, 1996

ROCURONIUM BROMIDE

INJECTABLE; INJECTION

ZEMURON

ORGANON USA INC

10MG/ML

N20214 002

Mar 17, 1994

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 236 (of 274)

ROFECOXIBSUSPENSION; ORAL
VIOXX

MERCK	12.5MG/5ML	N21052	001	May 20, 1999
	25MG/5ML	N21052	002	May 20, 1999

TABLET; ORAL
VIOXX

MERCK	12.5MG	N21042	001	May 20, 1999
	25MG	N21042	002	May 20, 1999
	50MG	N21042	003	Feb 25, 2000

ROSE BENGAL SODIUM, I-131INJECTABLE; INJECTION
ROBENGATOPE

BRACCO	0.5mCi/VIAL	N16224	001	
	1mCi/VIAL	N16224	002	
	2mCi/VIAL	N16224	003	

SODIUM ROSE BENGAL I 131
SORIN

0.5mCi/ML	N17318	001	
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SAFFLOWER OILINJECTABLE; INJECTION
LIPOSYN 10%

ABBOTT	10% (10GM/100ML)	N18203	001	
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LIPOSYN 20%
ABBOTT

20% (20GM/100ML)	N18614	001	
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SALMETEROL XINAFOATEAEROSOL, METERED; INHALATION
SEREVENT

GLAXOSMITHKLINE	EQ 0.021MG BASE/INH	N20236	001	Feb 04, 1994
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SAQUINAVIRCAPSULE; ORAL
FORTOVASE

HLR	200MG	N20828	001	Nov 07, 1997
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SARALASIN ACETATEINJECTABLE; INJECTION
SARENIN

PROCTER AND GAMBLE	EQ 0.6MG BASE/ML	N18009	001	
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SECOBARBITAL SODIUM

CAPSULE; ORAL

SECOBARBITAL SODIUM

IVAX PHARMS	100MG	N85869	001	
LANNETT	50MG	N85909	001	

	100MG	N85903	001	
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PARKE DAVIS	100MG	N84762	001	
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PUREPAC PHARM	100MG	N85867	001	
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VALEANT PHARM INTL	100MG	N85477	001	
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VITARINE	100MG	N85898	001	
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	100MG	N86273	001	
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WHITEWORTH TOWN PLSN	100MG	N85798	001	
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WYETH AYERST	100MG	N86390	001	
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SODIUM SECOBARBITAL

ANABOLIC	100MG	N84422	001	
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BARR	100MG	N84225	001	
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HALSEY	100MG	N84676	001	
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KV PHARM	100MG	N85285	001	
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PERRIGO	100MG	N84561	001	
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WATSON LABS	100MG	N85792	001	
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WEST WARD	100MG	N84926	001	
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DISCONTINUED DRUG PRODUCT LIST

6 - 237 (of 274)

SECOBARBITAL SODIUM

INJECTABLE; INJECTION		
SECOBARBITAL SODIUM		
ELKINS SINN	100MG/VIAL	N83281 001
SECONAL SODIUM		
LILLY	50MG/ML	N07392 002
SODIUM SECOBARBITAL		
WYETH AYERST	50MG/ML	N83262 001
SUPPOSITORY; RECTAL		
SECONAL SODIUM		
LILLY	30MG	N86530 001
	60MG	N86530 002
	120MG	N86530 003
	200MG	N86530 004

SECRETIN

INJECTABLE; INJECTION		
SECRETIN-FERRING		
FERRING	75CU/VIAL	N18290 001

SELEGILINE HYDROCHLORIDE

CAPSULE; ORAL			
SELEGILINE HYDROCHLORIDE			
AAIPHARMA LLC	5MG	N75145 001	Sep 15, 2003
TABLET; ORAL			
SELEGILINE HYDROCHLORIDE			
SOMERSET	5MG	N19334 001	Jun 05, 1989

SELENIUM SULFIDE

LOTION/SHAMPOO; TOPICAL		
EXSEL		
ALLERGAN HERBERT	2.5%	N83892 001
SELENIUM SULFIDE		
IVAX PHARMS	2.5%	N85777 001
TARO	2.5%	N86209 001

SELENOMETHIONINE, SE-75

INJECTABLE; INJECTION		
SELENOMETHIONINE SE 75		
CIS	500uCi/ML	N17322 001
GE HEALTHCARE	250uCi/ML	N17257 001
MALLINCKRODT	100uCi/ML	N17098 001
SETHOTOPE		
BRACCO	85-550uCi/ML	N17047 001

SERACTIDE ACETATE

INJECTABLE; INJECTION		
ACTHAR GEL-SYNTHETIC		
ARMOUR PHARM	40 UNITS/ML	N17861 001
	80 UNITS/ML	N17861 002

SERMORELIN ACETATE

INJECTABLE; INJECTION			
GEREF			
SERONO	EQ 0.5MG BASE/VIAL	N20443 001	Sep 26, 1997
	EQ 1MG BASE/VIAL	N20443 002	Sep 26, 1997

SERTRALINE HYDROCHLORIDE

TABLET; ORAL			
ZOLOFT			
PFIZER	EQ 150MG BASE **Federal Register determination that product was not discontinued or withdrawn for safety or efficacy reasons**	N19839 003	Dec 30, 1991

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 238 (of 274)

SERTRALINE HYDROCHLORIDETABLET; ORAL
ZOLOFT

PFIZER

EQ 200MG BASE **Federal Register
determination that product was not
discontinued or withdrawn for safety
or efficacy reasons**

N19839 004 Dec 30, 1991

SEVELAMER HYDROCHLORIDECAPSULE; ORAL
RENAGEL

GENZYME

403MG

N20926 001 Oct 30, 1998

SILVER SULFADIAZINEDRESSING; TOPICAL
SILDAFLO

FRANKLIN PHARMS

1%

N19608 001 Nov 30, 1989

SIMETHICONE-CELLULOSESUSPENSION; ORAL
SONORX

BRACCO

7.5MG/ML

N20773 001 Oct 29, 1998

SIROLIMUSTABLET; ORAL
RAPAMUNE

WYETH PHARMS INC

5MG

N21110 003 Feb 23, 2004

SODIUM BENZOATE; SODIUM PHENYLACETATESOLUTION; ORAL
UCEPHAN

B BRAUN

100MG/ML;100MG/ML

N19530 001 Dec 23, 1987

SODIUM BICARBONATE

INJECTABLE; INJECTION

SODIUM BICARBONATE IN PLASTIC CONTAINER

ABBOTT

0.9MEQ/ML

N19443 001 Jun 03, 1986

1MEQ/ML

N19443 002 Jun 03, 1986

SODIUM CHLORIDE

INJECTABLE; INJECTION

BACTERIOSTATIC SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER

ABRAXIS PHARM

9MG/ML

N88909 001 Feb 07, 1985

SODIUM CHLORIDE

ABBOTT

20GM/100ML

N17013 001

B BRAUN

20GM/100ML

N17038 001

SODIUM CHLORIDE 0.45% IN PLASTIC CONTAINER

B BRAUN

450MG/100ML

N18184 001

MILES

450MG/100ML

N18503 001

SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER

ABBOTT

9MG/ML

N19218 001

MILES

900MG/100ML

N18502 001

SODIUM CHLORIDE 23.4% IN PLASTIC CONTAINER

ABRAXIS PHARM

234MG/ML

N19329 001 Apr 22, 1987

SODIUM CHLORIDE 3% IN PLASTIC CONTAINER

B BRAUN

3GM/100ML

N19635 003 Mar 09, 1988

SODIUM CHLORIDE 5% IN PLASTIC CONTAINER

B BRAUN

5GM/100ML

N19635 004 Mar 09, 1988

SOLUTION; IRRIGATION

SODIUM CHLORIDE 0.45% IN PLASTIC CONTAINER

BAXTER HLTHCARE

450MG/100ML

N18497 001 Feb 19, 1982

SODIUM CHLORIDE IN PLASTIC CONTAINER

MILES

900MG/100ML

N18247 001

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 239 (of 274)

SODIUM CHROMATE, CR-51

INJECTABLE; INJECTION
CHROMITOPES SODIUM
BRACCO

2mCi/VIAL

N13993 002

SODIUM FLUORIDE, F-18

INJECTABLE; INTRAVENOUS
FLUORINE F-18
GE HEALTHCARE

2mCi/ML **Federal Register
determination that product was not
discontinued or withdrawn for safety
or efficacy reasons**

N17042 001

SODIUM IODIDE, I-123

CAPSULE; ORAL
SODIUM IODIDE I 123
SYNCOR PHARMS

400uCi

N18671 003 May 27, 1982

SODIUM IODIDE, I-131

CAPSULE; ORAL
IODOTOPE

BRACCO

1-150mCi

N10929 003

SODIUM IODIDE I 131

CIS

50uCi

N17316 001

100uCi

N17316 002

MALLINCKRODT

0.8-100mCi

N16515 002

SOLUTION; ORAL

IODOTOPE

BRACCO

7-106mCi/BOT

N10929 002

SODIUM IODIDE I 131

CIS

50mCi/ML

N17315 001

SODIUM LACTATE

INJECTABLE; INJECTION

SODIUM LACTATE 0.167 MOLAR IN PLASTIC CONTAINER

B BRAUN

1.87GM/100ML

N18186 001

SODIUM NITROPRUSSIDE

INJECTABLE; INJECTION

NIPRIDE

ROCHE

50MG/VIAL

N17546 001

NITROPRESS

ABBOTT

50MG/VIAL

N18450 001

50MG/VIAL

N71555 001 Nov 16, 1987

SODIUM NITROPRUSSIDE

ABRAXIS PHARM

50MG/VIAL

N70031 001

BAXTER HLTHCARE

50MG/VIAL

N18581 001

Jan 17, 1985

Jul 28, 1982

SODIUM PHOSPHATE, P-32

SOLUTION; INJECTION, ORAL
PHOSPHOTOPE

BRACCO

1-8mCi/VIAL

N10927 001

SODIUM PHOSPHATE P 32

MALLINCKRODT

1.5mCi/VIAL

N11777 002

SODIUM POLYSTYRENE SULFONATE

POWDER; ORAL, RECTAL

SODIUM POLYSTYRENE SULFONATE

MORTON GROVE

453.6GM/BOT

N88786 001

Sep 11, 1984

SUSPENSION; ORAL, RECTAL

SODIUM POLYSTYRENE SULFONATE

MORTON GROVE

15GM/60ML

N88717 001

Sep 11, 1984

ROXANE

15GM/60ML

N88453 001

Nov 17, 1983

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 240 (of 274)

SODIUM SUCCINATE

INJECTABLE; INJECTION			
SODIUM SUCCINATE			
ELKINS SINN	30%	N80516	001

SODIUM TETRADECYL SULFATE

INJECTABLE; INJECTION			
SOTRADECOL			
ELKINS SINN	1% **Federal Register determination that product was not discontinued or withdrawn for safety or efficacy reasons**	N05970	004
	3% **Federal Register determination that product was not discontinued or withdrawn for safety or efficacy reasons**	N05970	005

SODIUM THIOSULFATE

INJECTABLE; INJECTION			
SODIUM THIOSULFATE			
US ARMY	250MG/ML	N20166	001 Feb 14, 1992

SOMATREM

INJECTABLE; INJECTION			
PROTROPIN			
GENENTECH	5MG/VIAL	N19107	001 Oct 17, 1985
	10MG/VIAL	N19107	002 Oct 24, 1989

SOMATROPIN

INJECTABLE; INJECTION			
ASELLACRIN 10			
SERONO	10 IU/VIAL	N17726	001
ASELLACRIN 2			
SERONO	2 IU/VIAL	N17726	002 Jul 21, 1983
CRESCORMON			
GENENTECH	4 IU/VIAL	N17992	001

SOMATROPIN RECOMBINANT

INJECTABLE; INJECTION			
BIO-TROPIN			
FERRING	4.8MG/VIAL	N19774	001 May 25, 1995
HUMATROPE			
LILLY	2MG/VIAL	N19640	001 Jun 23, 1987
NUTROPIN DEPOT			
GENENTECH	13.5MG/VIAL	N21075	001 Dec 22, 1999
	18MG/VIAL	N21075	002 Dec 22, 1999
	22.5MG/VIAL	N21075	003 Dec 22, 1999
SAIZEN			
SERONO	6MG/VIAL	N19764	001 Oct 08, 1996
SEROSTIM			
SERONO	8.8MG/VIAL	N20604	004 Sep 06, 2001
ZORBTIVE			
SERONO INC	4MG/VIAL	N21597	001 Dec 01, 2003
	5MG/VIAL	N21597	002 Dec 01, 2003
	6MG/VIAL	N21597	003 Dec 01, 2003
INJECTABLE; SUBCUTANEOUS			
SEROSTIM LQ			
SERONO	6MG/0.5ML	N20604	005 Feb 11, 2005

SORBITOL

SOLUTION; IRRIGATION			
SORBITOL 3% IN PLASTIC CONTAINER			
BAXTER HLTHCARE	3GM/100ML	N18512	001 May 27, 1982

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 241 (of 274)

SOTALOL HYDROCHLORIDE

TABLET; ORAL

BETAPACE

BERLEX

320MG

N19865 004

Oct 30, 1992

BETAPACE AF

BERLEX

40MG

N21151 006

Apr 02, 2003

60MG

N21151 007

Apr 02, 2003

100MG

N21151 005

Mar 14, 2003

SOYBEAN OIL

INJECTABLE; INJECTION

SOYACAL 10%

ALPHA THERA

10%

N18465 001

Jun 29, 1983

SOYACAL 20%

ALPHA THERA

20%

N18786 001

Jun 29, 1983

TRAVAMULSION 10%

BAXTER HLTHCARE

10%

N18660 001

Feb 26, 1982

TRAVAMULSION 20%

BAXTER HLTHCARE

20%

N18758 001

Feb 15, 1983

SPARFLOXACIN

TABLET; ORAL

ZAGAM

MYLAN

200MG

N20677 001

Dec 19, 1996

SPECTINOMYCIN HYDROCHLORIDE

INJECTABLE; INJECTION

TROBICIN

PHARMACIA AND UPJOHN EQ 4GM BASE/VIAL

N50347 002

SPIRAPRIL HYDROCHLORIDE

TABLET; ORAL

RENORMAX

SCHERING

3MG

N20240 001

Dec 29, 1994

6MG

N20240 002

Dec 29, 1994

12MG

N20240 003

Dec 29, 1994

24MG

N20240 004

Dec 29, 1994

SPIRONOLACTONE

TABLET; ORAL

SPIRONOLACTONE

ASCOT

25MG

N87687 001

Oct 20, 1982

IVAX PHARMS

25MG

N87108 001

LEDERLE

25MG

N87634 001

PUREPAC PHARM

25MG

N87998 001

Oct 14, 1983

25MG

N88053 001

Aug 25, 1983

SUPERPHARM

25MG

N89364 001

Nov 07, 1986

UPSHER SMITH

25MG

N87554 001

VANGARD

25MG

N87648 001

Feb 01, 1982

WARNER CHILCOTT

25MG

N87952 001

Nov 18, 1982

WATSON LABS

25MG

N86898 002

Mar 02, 1982

25MG

N87078 001

STANOZOLOL

TABLET; ORAL

WINSTROL

OVATION PHARMS

2MG

N12885 001

May 14, 1984

STAVUDINE

CAPSULE; ORAL

ZERIT

BRISTOL MYERS SQUIBB 5MG

N20412 001

Jun 24, 1994

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 242 (of 274)

STAVUDINECAPSULE, EXTENDED RELEASE; ORAL
ZERIT XR

BRISTOL MYERS SQUIBB	37.5MG	N21453	001	Dec 31, 2002
	50MG	N21453	002	Dec 31, 2002
	75MG	N21453	003	Dec 31, 2002
	100MG	N21453	004	Dec 31, 2002

STREPTOMYCIN SULFATEINJECTABLE; INJECTION
STREPTOMYCIN SULFATE

COPANOS	EQ 500MG BASE/ML	N60684	001	
LILLY	EQ 1GM BASE/VIAL	N60107	001	
	EQ 1GM BASE/2ML	N60404	001	
	EQ 5GM BASE/VIAL	N60107	002	
PFIZER	EQ 1GM BASE/VIAL	N60076	001	
	EQ 5GM BASE/VIAL	N60076	002	

SUCCINYLCHOLINE CHLORIDEINJECTABLE; INJECTION
ANECTINE

SANDOZ	50MG/ML	N08453	003	
	500MG/VIAL	N08453	001	
	1GM/VIAL	N08453	004	

QUELICIN PRESERVATIVE FREE

HOSPIRA	50MG/ML	N08845	002	
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SUCCINYLCHOLINE CHLORIDE

INTL MEDICATION	100MG/VIAL	N85400	001	Feb 04, 1982
ORGANON USA INC	20MG/ML	N80997	001	

SUCOSTRIN

APOTHECON	20MG/ML	N08847	001	
	100MG/ML	N08847	003	

SUFENTANIL CITRATEINJECTABLE; INJECTION
SUFENTANIL CITRATE

WATSON LABS	EQ 0.05MG BASE/ML	N74406	001	Dec 15, 1995
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SULFACETAMIDE SODIUM

OINTMENT; OPHTHALMIC

BLEPH-10

ALLERGAN	10%	N84015	001	
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SODIUM SULAMYD

SCHERING	10%	N05963	002	
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SULFAIR 10

PHARMAFAIR	10%	N88000	001	Dec 22, 1982
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SOLUTION/DROPS; OPHTHALMIC

OCUSULF-30

MIZA PHARMS USA	30%	N80660	002	
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SODIUM SULAMYD

SCHERING	10%	N05963	001	
	30%	N05963	003	

SODIUM SULFACETAMIDE

AKORN	10%	N83021	001	
	15%	N83021	002	
	30%	N83021	003	
SOLA BARNES HIND	10%	N84143	001	
	10%	N84145	001	
	30%	N84146	001	
	30%	N84147	001	

SULF-15

NOVARTIS	15%	N89047	001	Oct 31, 1995
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SULFACETAMIDE SODIUM

PHARMAFAIR	10%	N88947	001	May 17, 1985
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27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 243 (of 274)

SULFACETAMIDE SODIUM

SOLUTION/DROPS; OPHTHALMIC

SULFAIR 10

PHARMAFAIR 10%

N87949 001 Dec 13, 1982

SULFAIR FORTE

PHARMAFAIR 30%

N88385 001 Oct 13, 1983

SULFAIR-15

PHARMAFAIR 15%

N88186 001 May 25, 1983

SULTEN-10

BAUSCH AND LOMB 10%

N87818 001 Feb 03, 1983

SULFACYTINE

TABLET; ORAL

RENOQUID

GLENWOOD 250MG

N17569 001

SULFADIAZINE

TABLET; ORAL

SULFADIAZINE

ABBOTT 300MG

N04125 005

EVERYLIFE 500MG

N80088 001

IMPAX LABS 500MG

N80081 001

LANNETT 500MG

N80084 001

LEDERLE 500MG

N04054 001

LILLY 500MG

N04122 002

SULFADIAZINE SODIUM

INJECTABLE; INJECTION

SULFADIAZINE SODIUM

LEDERLE 250MG/ML

N04054 002

SULFADIAZINE; SULFAMERAZINE

SUSPENSION; ORAL

SULFONAMIDES DUPLEX

LILLY 250MG/5ML; 250MG/5ML

N06317 007

SULFAMETER

TABLET; ORAL

SULLA

BERLEX 500MG

N16000 002

SULFAMETHIZOLE

TABLET; ORAL

MICROSUL

FOREST PHARMS 1GM

N86012 001

PROKLAR

FOREST PHARMS 500MG

N80273 001

THIOSULFIL

WYETH AYERST 250MG

N08565 001

500MG

N08565 004

SULFAMETHOXAZOLE

SUSPENSION; ORAL

GANTANOL

ROCHE 500MG/5ML

N13664 002

TABLET; ORAL

GANTANOL

ROCHE 500MG

N12715 002

GANTANOL-DS

ROCHE 1GM

N12715 003

SULFAMETHOXAZOLE

ASCOT 500MG

N87662 001 Oct 20, 1982

BARR 500MG

N87189 001 Jul 25, 1983

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 244 (of 274)

SULFAMETHOXAZOLE

TABLET; ORAL

SULFAMETHOXAZOLE

HEATHER	500MG	N86163	001
SANDOZ	500MG	N85844	001
WATSON LABS	500MG	N85053	001
	1GM	N86000	001
UROBAK			
SHIONOGI	500MG	N87307	001

SULFAMETHOXAZOLE; TRIMETHOPRIM

INJECTABLE; INJECTION

SEPTRA

MONARCH PHARMS	80MG/ML;16MG/ML	N18452	001
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SULFAMETHOXAZOLE AND TRIMETHOPRIM

ABRAXIS PHARM	80MG/ML;16MG/ML	N70223	001	Dec 29, 1987
BAXTER HLTHCARE	80MG/ML;16MG/ML	N70627	001	Dec 29, 1987
BEDFORD	80MG/ML;16MG/ML	N72383	001	Apr 29, 1992
WATSON LABS	80MG/ML;16MG/ML	N71556	001	Dec 29, 1987

SUSPENSION; ORAL

BACTRIM

MUTUAL PHARM	200MG/5ML;40MG/5ML	N17560	001
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SULFAMETHOXAZOLE AND TRIMETHOPRIM

TEVA	200MG/5ML;40MG/5ML	N70028	001	Jun 02, 1987
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SULMEPRIM

USL PHARMA	200MG/5ML;40MG/5ML	N70063	001	Aug 01, 1986
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SULMEPRIM PEDIATRIC

USL PHARMA	200MG/5ML;40MG/5ML	N70064	001	Aug 01, 1986
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TRIMETH/SULFA

ALPHARMA US PHARMS	200MG/5ML;40MG/5ML	N72289	001	May 23, 1988
	200MG/5ML;40MG/5ML	N72398	001	May 23, 1988
NASKA	200MG/5ML;40MG/5ML	N72399	001	May 23, 1988

TABLET; ORAL

SULFAMETHOPRIM-DS

PAR PHARM	800MG;160MG	N70032	001	Feb 15, 1985
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SULFAMETHOXAZOLE AND TRIMETHOPRIM

HEATHER	400MG;80MG	N18946	001	Aug 10, 1984
	800MG;160MG	N18946	002	Aug 10, 1984
INTERPHARM	400MG;80MG	N71299	001	Oct 27, 1987
	800MG;160MG	N71300	001	Oct 27, 1987
MARTEC USA LLC	400MG;80MG	N72408	001	Dec 07, 1988
MUTUAL PHARM	400MG;80MG	N70006	001	Nov 14, 1984
ROXANE	400MG;80MG	N72768	001	Aug 30, 1991
SANDOZ	400MG;80MG	N18598	003	May 19, 1982
	400MG;80MG	N70889	001	Nov 13, 1986
TEVA	400MG;80MG	N18242	001	
	800MG;160MG	N18242	002	
USL PHARMA	400MG;80MG	N70203	001	Nov 08, 1985
	800MG;160MG	N70204	001	Nov 08, 1985
WATSON LABS	400MG;80MG	N70002	001	Nov 07, 1984
	800MG;160MG	N70000	001	Nov 07, 1984

SULFAMETHOXAZOLE AND TRIMETHOPRIM DOUBLE STRENGTH

MARTEC USA LLC	800MG;160MG	N72417	001	Dec 07, 1988
MUTUAL PHARM	800MG;160MG	N70007	001	Nov 14, 1984
ROXANE	800MG;160MG	N72769	001	Aug 30, 1991

SULFATRIM-DS

SUPERPHARM	800MG;160MG	N70066	001	Jun 24, 1985
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SULFATRIM-SS

SUPERPHARM	400MG;80MG	N70065	002	Jun 24, 1985
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UROPLUS DS

SHIONOGI	800MG;160MG	N71816	001	Sep 28, 1987
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UROPLUS SS

SHIONOGI	400MG;80MG	N71815	001	Sep 28, 1987
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27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 245 (of 274)

SULFANILAMIDE

CREAM; VAGINAL SULFANILAMIDE TEVA	15%	N88718	001	Sep 19, 1985
SUPPOSITORY; VAGINAL AVC PHARMELLE	1.05GM	N06530	004	Jan 27, 1987

SULFAPHENAZOLE

SUSPENSION; ORAL SULFABID PURDUE FREDERICK	500MG/5ML	N13093	001	
TABLET; ORAL SULFABID PURDUE FREDERICK	500MG	N13092	002	

SULFAPYRIDINE

TABLET; ORAL SULFAPYRIDINE LILLY	500MG	N00159	001	
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SULFASALAZINE

SUSPENSION; ORAL AZULFIDINE PHARMACIA AND UPJOHN	250MG/5ML	N18605	001	
TABLET; ORAL S.A.S. - 500 SOLVAY	500MG	N83450	001	
SULFASALAZINE RADIUS PHARMS	500MG	N80197	001	
SANDOZ	500MG	N86184	001	
SUPERPHARM	500MG	N89339	001	Oct 26, 1987
WATSON LABS	500MG	N84964	001	
TABLET, DELAYED RELEASE; ORAL SULFASALAZINE WATSON LABS	500MG	N88052	001	May 24, 1983

SULFINPYRAZONE

CAPSULE; ORAL SULFINPYRAZONE VANGARD	200MG	N88666	001	Feb 17, 1984
TABLET; ORAL SULFINPYRAZONE IVAX PHARMS	100MG	N87769	001	Jun 01, 1982
WATSON LABS	100MG	N87667	001	May 26, 1982

SULFISOXAZOLE

TABLET; ORAL GANTRISIN ROCHE	500MG	N06525	001	
SOSOL MK LABS	500MG	N80036	001	
SOXAZOLE ALRA	500MG	N80366	001	
SULFALAR PARKE DAVIS	500MG	N84955	001	
SULFISOXAZOLE BARR	500MG	N84031	001	
HEATHER	500MG	N80189	001	
IMPAX LABS	500MG	N80109	001	
LANNETT	500MG	N80085	001	
LEDERLE	500MG	N87649	001	
PHARMERAL	500MG	N84385	001	
PUREPAC PHARM	500MG	N80087	001	

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 246 (of 274)

SULFISOXAZOLETABLET; ORAL
SULFISOXAZOLE

ROXANE	500MG	N80082	001
SANDOZ	500MG	N85628	001
VALEANT PHARM INTL	500MG	N80268	002
VITARINE	500MG	N87332	001
WATSON LABS	500MG	N85534	001
WEST WARD	500MG	N80379	001
SULSOXIN			
SOLVAY	500MG	N80040	001

SULFISOXAZOLE ACETYLEMULSION; ORAL
LIPO GANTRISIN

ROCHE	EQ 1GM BASE/5ML	N09182	009
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SYRUP; ORAL
GANTRISIN

ROCHE	EQ 500MG BASE/5ML	N09182	002
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SULFISOXAZOLE DIOLAMINEINJECTABLE; INJECTION
GANTRISIN

ROCHE	EQ 400MG BASE/ML	N06917	001
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OINTMENT; OPHTHALMIC
GANTRISIN

ROCHE	EQ 4% BASE	N08414	002
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SOLUTION/DROPS; OPHTHALMIC
GANTRISIN

ROCHE	EQ 4% BASE	N07757	002
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SULFISOXAZOLE DIOLAMINE

SOLA BARNES HIND	EQ 4% BASE	N84148	001
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SULFOXONE SODIUMTABLET, DELAYED RELEASE; ORAL
DIASONE SODIUM

ABBOTT	165MG	N06044	003
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SULFURPOWDER; TOPICAL
BENSULFOID

POYTHRESS	33.32%	N02918	001
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SULINDACTABLET; ORAL
CLINORIL

MERCK	150MG	N17911	001
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SULINDAC

RADIUS PHARMS	150MG	N73262	002	Sep 06, 1991
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	200MG	N73262	001	Sep 06, 1991
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WARNER CHILCOTT	150MG	N72710	001	Mar 25, 1991
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	200MG	N72711	001	Mar 25, 1991
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SUPROFENSOLUTION/DROPS; OPHTHALMIC
PROFENAL

ALCON	1%	N19387	001	Dec 23, 1988
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27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 247 (of 274)

SUTILAINS

OINTMENT; TOPICAL

TRAVASE

ABBOTT

82,000 UNITS/GM **Federal Register
determination that product was not
discontinued or withdrawn for safety
or efficacy reasons**

N12828 001

TALBUTAL

TABLET; ORAL

LOTUSATE

SANOFI AVENTIS US

120MG

N09410 005

TAMOXIFEN CITRATE

TABLET; ORAL

TAMOXIFEN CITRATE

PHARMACHEMIE

EQ 10MG BASE

N74539 001 Mar 31, 2003

TECHNETIUM TC-99M ALBUMIN AGGREGATED

INJECTABLE; INJECTION

TC 99M-LUNGAGGREGATE

GE HEALTHCARE

5mCi/ML

N17848 001

TECHNETIUM TC-99M ALBUMIN AGGREGATED KIT

INJECTABLE; INJECTION

A-N STANNOUS AGGREGATED ALBUMIN

SYNCOR PHARMS

N/A

N17916 001

AN-MAA

CIS

N/A

N17792 001

LUNGAGGREGATE REAGENT

GE HEALTHCARE

N/A

N17838 001

TECHNETIUM TC 99M MAA

GE HEALTHCARE

N/A

N17773 001

TECHNETIUM TC-99M ALBUMIN COLLOID KIT

INJECTABLE; INJECTION

MICROLITE

CIS

N/A

N18263 001 Mar 25, 1983

TECHNETIUM TC-99M ALBUMIN KIT

INJECTABLE; INJECTION

TECHNETIUM TC 99M HSA

GE HEALTHCARE

N/A

N17775 001

TECHNETIUM TC-99M ALBUMIN MICROSPHERES KIT

INJECTABLE; INJECTION

INSTANT MICROSPHERES

3M

N/A

N17832 001

TECHNETIUM TC-99M APCITIDE

INJECTABLE; INJECTION

ACUTECT

CIS BIO INTL SA

N/A

N20887 001 Sep 14, 1998

TECHNETIUM TC-99M DEPREOTIDE

INJECTABLE; INJECTION

NEO TECT KIT

CIS BIO INTL SA

N/A

N21012 001 Aug 03, 1999

TECHNETIUM TC-99M ETIDRONATE KIT

INJECTABLE; INJECTION

CINTICHEM TECHNETIUM 99M HEDSPA

GE HEALTHCARE

N/A

N17653 001

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 248 (of 274)

TECHNETIUM TC-99M ETIDRONATE KIT

INJECTABLE; INJECTION		
MPI STANNOUS DIPHOSPHONATE		
GE HEALTHCARE	N/A	N17667 001
OSTEOSCAN		
MALLINCKRODT	N/A	N17454 001
TECHNETIUM TC 99M DIPHOSPHONATE-TIN KIT		
GE HEALTHCARE	N/A	N17562 001

TECHNETIUM TC-99M FERMENTETATE KIT

INJECTABLE; INJECTION		
RENOTEC		
BRACCO	N/A	N17045 001

TECHNETIUM TC-99M GLUCEPTATE KIT

INJECTABLE; INJECTION		
GLUCOSCAN		
BRISTOL MYERS SQUIBB	N/A	N17907 001

TECHNETIUM TC-99M LIDOFENIN KIT

INJECTABLE; INJECTION		
TECHNISCAN HIDA		
DRAXIMAGE	N/A	N18489 001 Oct 31, 1986

TECHNETIUM TC-99M MEDRONATE KIT

INJECTABLE; INJECTION		
AMERSCAN MDP KIT		
GE HEALTHCARE	N/A	N18335 001 Aug 05, 1982
OSTEOLITE		
CIS	N/A	N17972 001

TECHNETIUM TC-99M POLYPHOSPHATE KIT

INJECTABLE; INJECTION		
SODIUM POLYPHOSPHATE-TIN KIT		
GE HEALTHCARE	N/A	N17664 001

TECHNETIUM TC-99M PYRO/TRIMETA PHOSPHATES KIT

INJECTABLE; INJECTION		
PYROLITE		
CIS	N/A	N17684 001

TECHNETIUM TC-99M RED BLOOD CELL KIT

INJECTABLE; INJECTION		
RBC-SCAN		
CADEMA	N/A	N20063 001 Jun 11, 1992

TECHNETIUM TC-99M SODIUM PERTECHNETATE

SOLUTION; INJECTION, ORAL		
SODIUM PERTECHNETATE TC 99M		
CIS	12mCi/ML	N17321 001
	24mCi/ML	N17321 002
	48mCi/ML	N17321 003
GE HEALTHCARE	2-100mCi/ML	N17471 001
MALLINCKRODT	10-60mCi/ML	N17725 001

TECHNETIUM TC-99M SODIUM PERTECHNETATE GENERATOR

SOLUTION; INJECTION, ORAL		
MINITEC		
BRACCO	0.22-2.22 CI/GENERATOR	N17339 001
TECHNETIUM TC 99M GENERATOR		
GE HEALTHCARE	830-16600mCi/GENERATOR	N17693 001

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 249 (of 274)

TECHNETIUM TC-99M SULFUR COLLOID

SOLUTION; INJECTION, ORAL		
TECHNETIUM TC 99M SULFUR COLLOID		
GE HEALTHCARE	4mCi/ML	N17456 001
SOLUTION; ORAL		
TECHNETIUM TC 99M SULFUR COLLOID		
MALLINCKRODT	3mCi/ML	N17724 001

TECHNETIUM TC-99M SULFUR COLLOID KIT

SOLUTION; INJECTION, ORAL		
TECHNETIUM TC 99M TSC		
GE HEALTHCARE	N/A	N17784 001
TESULOID		
BRACCO	N/A	N16923 001

TECHNETIUM TC-99M TEBOROXIME KIT

INJECTABLE; INJECTION			
CARDIOTEC			
BRACCO	N/A	N19928 001	Dec 19, 1990

TECHNETIUM TC-99M TETROFOSMIN KIT

INJECTABLE; INJECTION			
MYOVUE 30ML			
GE HEALTHCARE	N/A	N20372 002	Jul 07, 2005

TEMAZEPAM

CAPSULE; ORAL			
TEMAZ			
QUANTUM PHARMICS	15MG	N70564 001	Oct 15, 1985
	30MG	N70547 001	Oct 15, 1985
TEMAZEPAM			
DURAMED PHARMS BARR	15MG	N71708 001	Sep 29, 1988
	30MG	N71709 001	Sep 29, 1988
MUTUAL PHARM	15MG	N71174 001	Jul 10, 1986
	30MG	N71175 001	Jul 10, 1986
USL PHARMA	15MG	N70489 001	Jul 07, 1986
	30MG	N70490 001	Jul 07, 1986
WATSON LABS	15MG	N70383 001	Mar 23, 1987
	30MG	N70384 001	Mar 23, 1987

TERAZOSIN HYDROCHLORIDE

CAPSULE; ORAL			
TERAZOSIN HYDROCHLORIDE			
IVAX PHARMS	EQ 1MG BASE	N75614 002	Jan 30, 2001
	EQ 2MG BASE	N75614 001	Jan 30, 2001
	EQ 5MG BASE	N75614 003	Jan 30, 2001
	EQ 10MG BASE	N75614 004	Jan 30, 2001
TABLET; ORAL			
TERAZOSIN HYDROCHLORIDE			
IVAX PHARMS	EQ 1MG BASE	N74530 001	Apr 21, 2000
	EQ 2MG BASE	N74530 002	Apr 21, 2000
	EQ 5MG BASE	N74530 003	Apr 21, 2000
	EQ 10MG BASE	N74530 004	Apr 21, 2000
TEVA	EQ 1MG BASE	N74446 001	May 18, 2000
	EQ 2MG BASE	N74446 002	May 18, 2000
	EQ 5MG BASE	N74446 003	May 18, 2000
	EQ 10MG BASE	N74446 004	May 18, 2000

TERBINAFINE HYDROCHLORIDE

CREAM; TOPICAL			
LAMISIL			
NOVARTIS	1%	N20192 001	Dec 30, 1992

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 250 (of 274)

TERBUTALINE SULFATE

AEROSOL, METERED; INHALATION				
BRETHAIRE				
NOVARTIS	0.2MG/INH	N18762	001	Aug 17, 1984
BRICANYL				
SANOFI AVENTIS US	0.2MG/INH	N18000	001	Mar 19, 1985
INJECTABLE; INJECTION				
BRETHINE				
AAIPHARMA LLC	1MG/ML	N18571	001	
BRICANYL				
SANOFI AVENTIS US	1MG/ML	N17466	001	
TABLET; ORAL				
BRICANYL				
SANOFI AVENTIS US	2.5MG	N17618	001	
	5MG	N17618	002	

TERIPARATIDE ACETATE

INJECTABLE; INJECTION				
PARATHAR				
SANOFI AVENTIS US	200 UNITS/VIAL	N19498	001	Dec 23, 1987

TESTOLACTONE

INJECTABLE; INJECTION				
TESLAC				
BRISTOL MYERS SQUIBB	100MG/ML	N16119	001	
TABLET; ORAL				
TESLAC				
BRISTOL MYERS SQUIBB	250MG	N16118	002	

TESTOSTERONE

FILM, EXTENDED RELEASE; TRANSDERMAL				
TESTODERM				
ALZA	4MG/24HR	N19762	001	Oct 12, 1993
	6MG/24HR	N19762	002	Oct 12, 1993
TESTODERM TTS				
ALZA	5MG/24HR	N20791	001	Dec 18, 1997
INJECTABLE; INJECTION				
TESTOSTERONE				
WATSON LABS	25MG/ML	N86420	001	May 10, 1983
	50MG/ML	N86419	001	Aug 23, 1983

TESTOSTERONE CYPIONATE

INJECTABLE; INJECTION				
DEPO-TESTOSTERONE				
PHARMACIA AND UPJOHN	50MG/ML	N85635	001	
TESTOSTERONE CYPIONATE				
WATSON LABS	100MG/ML	N84401	001	
	200MG/ML	N84401	002	

TESTOSTERONE ENANTHATE

INJECTABLE; INJECTION				
DELATESTRYL				
INDEVUS PHARMS	200MG/ML	N09165	001	
TESTOSTERONE ENANTHATE				
WATSON LABS	100MG/ML	N83667	001	
	100MG/ML	N85599	001	
	200MG/ML	N83667	002	

TESTOSTERONE PROPIONATE

INJECTABLE; INJECTION				
TESTOSTERONE PROPIONATE				
BEL MAR	25MG/ML	N80741	001	
	50MG/ML	N80742	001	
	100MG/ML	N80743	001	

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 251 (of 274)

TESTOSTERONE PROPIONATEINJECTABLE; INJECTION
TESTOSTERONE PROPIONATE

ELKINS SINN	25MG/ML	N80276	001
LILLY	50MG/ML	N80254	002
WATSON LABS	25MG/ML	N85490	001
	50MG/ML	N85490	002
	100MG/ML	N83595	003

TETRACYCLINE HYDROCHLORIDECAPSULE; ORAL
ACHROMYCIN V

RADIUS PHARMS	250MG	N50278	003
	500MG	N50278	001

CYCLOPAR

WARNER CHILCOTT	250MG	N61725	001
	250MG	N62175	001
	250MG	N62332	001
	500MG	N61725	002
	500MG	N62332	002

PANMYCIN

PHARMACIA AND UPJOHN	250MG	N60347	001
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RETET

SOLVAY	250MG	N61443	001
	500MG	N61443	002

ROBITET

WYETH AYERST	250MG	N61734	001
	500MG	N61734	002

SUMYCIN

APOTHECON	100MG	N60429	002
	125MG	N60429	004
	250MG	N60429	001
	500MG	N60429	003

TETRACHEL

ANGUS	250MG	N60343	001
	500MG	N60343	003

TETRACYCLINE HYDROCHLORIDE

ABBOTT	250MG	N61802	001
	500MG	N61802	002

ELKINS SINN	250MG	N60059	001
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FERRANTE	125MG	N60173	001
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	250MG	N60173	002
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HEATHER	250MG	N61148	001
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	500MG	N61148	002
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LEINER	250MG	N62686	001
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	500MG	N62686	002
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MAST MM	250MG	N62085	001
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PUREPAC PHARM	250MG	N60290	001
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	500MG	N60290	002
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ROXANE	500MG	N61214	002
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SANDOZ	250MG	N61471	001
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SUPERPHARM	250MG	N62540	001
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	500MG	N62540	002
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VALEANT PHARM INTL	250MG	N60471	001
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	500MG	N60471	002
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WARNER CHILCOTT	250MG	N62300	001
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	500MG	N62300	002
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WATSON LABS	250MG	N62103	001
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	250MG	N62343	001
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	500MG	N62103	002
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	500MG	N62343	002
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WEST WARD	250MG	N60768	001
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	500MG	N60768	002
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WYETH AYERST	250MG	N61685	001
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Jul 24, 1986

Jul 24, 1986

Mar 21, 1985

Mar 21, 1985

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 252 (of 274)

TETRACYCLINE HYDROCHLORIDE

CAPSULE; ORAL

TETRACYCLINE HYDROCHLORIDE

WYETH AYERST 500MG

N61685 002

TETRACYN

PFIPHARMECS 250MG

N60082 003

500MG

N60082 004

FIBER, EXTENDED RELEASE; PERIODONTAL

ACTISITE

ALZA 12.7MG/FIBER

N50653 001

Mar 25, 1994

FOR SOLUTION; TOPICAL

TOPICYCLINE

SHIRE 2.2MG/ML

N50493 001

INJECTABLE; INJECTION

ACHROMYCIN

LEDERLE 250MG/VIAL

N50273 002

500MG/VIAL

N50273 003

TETRACYN

PFIZER 250MG/VIAL

N60096 001

500MG/VIAL

N60096 002

OINTMENT; OPHTHALMIC

ACHROMYCIN

STORZ 10MG/GM

N50266 001

SUSPENSION; ORAL

ACHROMYCIN V

LEDERLE 125MG/5ML

N50263 002

TETRACYCLINE HYDROCHLORIDE

ALPHARMA US PHARMS 125MG/5ML

N60633 001

FERRANTE 125MG/5ML

N60174 001

PROTER 125MG/5ML

N60446 001

PUREPAC PHARM 125MG/5ML

N60291 001

TETRACYN

PFIPHARMECS 125MG/5ML

N60095 001

TETRAMED

IVAX PHARMS 125MG/5ML

N61468 001

SUSPENSION/DROPS; OPHTHALMIC

ACHROMYCIN

STORZ 1%

N50268 001

TABLET; ORAL

PANMYCIN

PHARMACIA AND UPJOHN 250MG

N61705 001

500MG

N61705 002

SUMYCIN

PAR PHARM 50MG

N61147 003

100MG

N61147 002

TETRACYCLINE PHOSPHATE COMPLEX

CAPSULE; ORAL

TETREX

BRISTOL EQ 250MG HCL

N50212 002

EQ 500MG HCL

N50212 003

THALLOUS CHLORIDE, TL-201

INJECTABLE; INJECTION

THALLOUS CHLORIDE TL 201

BRACCO 1mCi/ML

N18548 001

Dec 30, 1982

THEOPHYLLINE

CAPSULE; ORAL

BRONKODYL

SANOFI AVENTIS US 100MG

N85264 001

200MG

N85264 002

ELIXOPHYLLIN

FOREST LABS 100MG

N85545 001

Jul 31, 1984

200MG

N83921 001

Jul 31, 1984

DISCONTINUED DRUG PRODUCT LIST

6 - 253 (of 274)

THEOPHYLLINE

CAPSULE; ORAL SOMOPHYLLIN-T

FISONS	100MG	N87155	001	Feb 25, 1985
	200MG	N87155	002	Feb 25, 1985
	250MG	N87155	003	Feb 25, 1985

THEOPHYLLINE

KV PHARM	100MG	N85263	001	
	200MG	N85263	002	
SCHERER RP	100MG	N84731	002	Nov 07, 1986
	200MG	N84731	001	Nov 07, 1986
	250MG	N84731	003	Nov 07, 1986

CAPSULE, EXTENDED RELEASE; ORAL

AEROLATE III

FLEMING PHARMS	65MG	N85075	003	Nov 24, 1986
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AEROLATE JR

FLEMING PHARMS	130MG	N85075	002	Nov 24, 1986
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AEROLATE SR

FLEMING PHARMS	260MG	N85075	001	Nov 24, 1986
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ELIXOPHYLLIN SR

FOREST LABS	125MG	N86826	001	Jan 29, 1985
	250MG	N86826	002	Jan 29, 1985

SLO-BID

SANOFI AVENTIS US	50MG	N88269	001	Jan 31, 1985
	75MG	N89539	001	May 10, 1989
	100MG	N87892	001	Jan 31, 1985
	125MG	N89540	001	May 10, 1989
	200MG	N87893	001	Jan 31, 1985
	300MG	N87894	001	Jan 31, 1985

SLO-PHYLLIN

SANOFI AVENTIS US	60MG	N85206	001	May 24, 1982
	125MG	N85203	001	May 24, 1982
	250MG	N85205	001	May 24, 1982

SOMOPHYLLIN-CRT

GRAHAM DM	50MG	N87763	001	Feb 27, 1985
	100MG	N87194	001	
	200MG	N88382	001	Feb 27, 1985
	250MG	N87193	001	
	300MG	N88383	001	Feb 27, 1985

THEOBID

WHITBY	260MG	N85983	001	Mar 20, 1985
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THEOBID JR.

WHITBY	130MG	N87854	001	Mar 20, 1985
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THEOCLEAR L.A.-130

SCHWARZ PHARMA	130MG	N86569	001	May 27, 1982
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THEOCLEAR L.A.-260

SCHWARZ PHARMA	260MG	N86569	002	May 27, 1982
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THEO-DUR

SCHERING	50MG	N88022	001	Sep 10, 1985
	75MG	N88015	001	Sep 10, 1985
	125MG	N88016	001	Sep 10, 1985
	200MG	N87995	001	Sep 10, 1985

THEOPHYLLINE

CENT PHARMS	125MG	N88654	001	Feb 12, 1985
	250MG	N88689	001	Feb 12, 1985
MAYNE PHARMA USA	100MG	N89976	001	Jan 04, 1995
	200MG	N89977	001	Jan 04, 1995
	300MG	N89932	001	Jan 04, 1995
SANDOZ	260MG	N87462	001	May 11, 1982

THEOPHYLLINE-SR

SCHERER RP	300MG	N88255	001	Jun 12, 1986
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THEOPHYL-SR

ORTHO MCNEIL PHARM	125MG	N86480	001	Feb 08, 1985
	250MG	N86471	001	Feb 08, 1985

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 254 (of 274)

THEOPHYLLINE

CAPSULE, EXTENDED RELEASE; ORAL

THEOVENT

SCHERING

125MG

N87010 001

Jan 31, 1985

250MG

N87910 001

Jan 31, 1985

ELIXIR; ORAL

ELIXOMIN

CENCI

80MG/15ML

N88303 001

Jan 25, 1984

LANOPHYLLIN

LANNETT

80MG/15ML

N84578 001

THEOLIXIR

PANRAY

80MG/15ML

N84559 001

THEOPHYL-225

ORTHO MCNEIL PHARM

112.5MG/15ML

N86485 001

THEOPHYLLINE

ALPHARMA US PHARMS

80MG/15ML

N89223 001

May 27, 1988

CENCI

80MG/15ML

N87679 001

Apr 15, 1982

HALSEY

80MG/15ML

N85169 001

PERRIGO

80MG/15ML

N85952 001

ROXANE

80MG/15ML

N84739 001

INJECTABLE; INJECTION

THEOPHYLLINE 0.04% AND DEXTROSE 5% IN PLASTIC CONTAINER

B BRAUN

40MG/100ML

N19083 001

Nov 07, 1984

THEOPHYLLINE 0.08% AND DEXTROSE 5% IN PLASTIC CONTAINER

B BRAUN

80MG/100ML

N19083 002

Nov 07, 1984

THEOPHYLLINE 0.16% AND DEXTROSE 5% IN PLASTIC CONTAINER

B BRAUN

160MG/100ML

N19083 003

Nov 07, 1984

THEOPHYLLINE 0.2% AND DEXTROSE 5% IN PLASTIC CONTAINER

B BRAUN

200MG/100ML

N19212 001

Nov 07, 1984

200MG/100ML

N19826 004

Aug 14, 1992

THEOPHYLLINE 0.4% AND DEXTROSE 5% IN PLASTIC CONTAINER

B BRAUN

4MG/ML

N19212 003

Nov 07, 1984

400MG/100ML

N19212 002

Nov 07, 1984

400MG/100ML

N19826 005

Aug 14, 1992

SOLUTION; ORAL

AEROLATE

FLEMING PHARMS

150MG/15ML

N89141 001

Dec 03, 1986

THEOLAIR

3M

80MG/15ML

N86107 001

SUSPENSION; ORAL

ELIXICON

FOREST LABS

100MG/5ML

N85502 001

SYRUP; ORAL

ACCURBRON

SANOFI AVENTIS US

150MG/15ML

N88746 001

Nov 22, 1985

AQUAPHYLLIN

FERNDAL LABS

80MG/15ML

N87917 001

Jan 18, 1983

SLO-PHYLLIN

SANOFI AVENTIS US

80MG/15ML

N85187 001

THEOCLEAR-80

CENT PHARMS

80MG/15ML

N87095 001

Mar 01, 1982

THEOPHYLLINE

ALPHARMA US PHARMS

80MG/15ML

N86001 001

150MG/15ML

N86545 001

TABLET; ORAL

SLO-PHYLLIN

SANOFI AVENTIS US

100MG

N85202 001

200MG

N85204 001

THEOCLEAR-100

CENT PHARMS

100MG

N85353 002

THEOCLEAR-200

CENT PHARMS

200MG

N85353 001

THEOPHYL-225

ORTHO MCNEIL PHARM

225MG

N84726 001

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 255 (of 274)

THEOPHYLLINETABLET, CHEWABLE; ORAL
THEOPHYL

ORTHO MCNEIL PHARM 100MG

N86506 001 Sep 12, 1985

TABLET, EXTENDED RELEASE; ORAL
DURAPHYL

FOREST LABS 100MG

N88503 001 Apr 03, 1985

200MG

N88504 001 Apr 03, 1985

300MG

N88505 001 Apr 03, 1985

LABID

PROCTER AND GAMBLE 250MG

N87225 001

SUSTAIRE

ROERIG 100MG

N85665 001

300MG

N85665 002

THEO-DUR

SCHERING 100MG

N85328 001

200MG

N86998 001

300MG

N85328 002

450MG

N89131 001 Jun 25, 1986

THEOLAIR-SR

3M 200MG

N88369 001 Jul 16, 1987

250MG

N86363 002 Jul 16, 1987

300MG

N88364 001 Jul 16, 1987

500MG

N89132 001 Jul 16, 1987

THEOPHYLLINE

ABLE 300MG

N40548 001 Apr 30, 2004

400MG

N40543 001 Apr 27, 2004

450MG

N40546 001 Apr 30, 2004

600MG

N40539 001 Apr 27, 2004

T-PHYL

PURDUE FREDERICK 200MG

N88253 001 Aug 17, 1983

UNI-DUR

SCHERING 400MG

N89822 001 Jan 04, 1995

600MG

N89823 001 Jan 04, 1995

THEOPHYLLINE SODIUM GLYCINATE

ELIXIR; ORAL

SYNOPHYLATE

CENT PHARMS EQ 165MG BASE/15ML

N06333 008

TABLET; ORAL

ASBRON

NOVARTIS EQ 150MG BASE

N85148 001

THIABENDAZOLE

SUSPENSION; ORAL

MINTEZOL

MERCK 500MG/5ML

N16097 001

THIAMINE HYDROCHLORIDE

INJECTABLE; INJECTION

BETALIN S

LILLY 100MG/ML

N80853 001

THIAMINE HYDROCHLORIDE

ABRAXIS PHARM 100MG/ML

N80509 001

AKORN 100MG/ML

N87968 001 Oct 01, 1982

BEL MAR 100MG/ML

N80718 001

200MG/ML

N80712 001

DELL LABS 100MG/ML

N83775 001

LUITPOLD 100MG/ML

N80667 001

PARKE DAVIS 100MG/ML

N80770 001

WATSON LABS 100MG/ML

N83534 001

200MG/ML

N83534 002

WYETH AYERST 100MG/ML

N80553 001

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 256 (of 274)

THIAMYLAL SODIUMINJECTABLE; INJECTION
SURITAL

PARKEDALE	1GM/VIAL	N07600	003
	5GM/VIAL	N07600	005
	10GM/VIAL	N07600	009

THIETHYLPERAZINE MALATEINJECTABLE; INJECTION
TORECAN

NOVARTIS	5MG/ML	N12754	002
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THIETHYLPERAZINE MALEATESUPPOSITORY; RECTAL
TORECAN

NOVARTIS	10MG	N13247	001
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THIOPENTAL SODIUMSUSPENSION; RECTAL
PENTOTHAL

ABBOTT	400MG/GM	N11679	001
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THIORIDAZINESUSPENSION; ORAL
MELLARIL-S

NOVARTIS	EQ 25MG HCL/5ML	N17923	001
	EQ 100MG HCL/5ML	N17923	002

THIORIDAZINE HYDROCHLORIDECONCENTRATE; ORAL
MELLARIL

NOVARTIS	30MG/ML	N11808	012
	100MG/ML	N11808	018

THIORIDAZINE HYDROCHLORIDE

ALPHARMA US PHARMS	30MG/ML	N87766	001	Apr 26, 1983
HI TECH PHARMA	30MG/ML	N40125	001	Aug 16, 1996
	100MG/ML	N40126	001	Aug 16, 1996
MORTON GROVE	30MG/ML	N88258	001	Jul 25, 1983
	100MG/ML	N88227	001	Jul 05, 1983
SANDOZ	30MG/ML	N88307	001	Nov 23, 1983
	100MG/ML	N88308	001	Nov 23, 1983

TABLET; ORAL
MELLARIL

NOVARTIS	10MG	N11808	003
	15MG	N11808	016
	25MG	N11808	006
	50MG	N11808	011
	100MG	N11808	009
	150MG	N11808	017
	200MG	N11808	015

THIORIDAZINE HYDROCHLORIDE

IVAX PHARMS	50MG	N88194	001	Apr 14, 1983
	100MG	N88273	001	Oct 03, 1983
MUTUAL PHARM	10MG	N88375	001	Nov 18, 1983
	15MG	N88461	001	Nov 18, 1983
	25MG	N87264	001	Nov 18, 1983
	100MG	N88379	001	Nov 16, 1983
	150MG	N88737	001	Sep 26, 1984
	200MG	N88738	001	Oct 16, 1984
MYLAN	10MG	N88332	001	Jun 27, 1983
	25MG	N88333	001	Jun 27, 1983
	50MG	N88334	001	Jun 27, 1983
	100MG	N88335	001	Nov 18, 1983
PAR PHARM	10MG	N88351	001	Dec 05, 1983

DISCONTINUED DRUG PRODUCT LIST

6 - 257 (of 274)

THIORIDAZINE HYDROCHLORIDE

TABLET; ORAL

THIORIDAZINE HYDROCHLORIDE

PAR PHARM	15MG	N88352	001	Dec 05, 1983
	25MG	N88336	001	Dec 05, 1983
	50MG	N88322	001	Dec 05, 1983
	100MG	N88480	001	Dec 29, 1983
	150MG	N89764	001	Feb 09, 1988
ROXANE	200MG	N89765	001	Feb 09, 1988
	10MG	N88663	001	Mar 15, 1984
	25MG	N88664	001	Mar 15, 1984
	50MG	N88665	001	Mar 15, 1984
	100MG	N89048	001	Feb 26, 1985
SUPERPHARM	10MG	N89103	001	Jul 02, 1985
	25MG	N89104	001	Jul 02, 1985
	50MG	N89105	001	Jul 02, 1985
TEVA	10MG	N88493	001	May 17, 1985
WATSON LABS	10MG	N88412	001	Sep 12, 1983
	10MG	N88561	001	May 11, 1984
	15MG	N88345	001	Jul 28, 1983
	15MG	N88562	001	May 11, 1984
	25MG	N88296	001	Jul 28, 1983
	25MG	N88755	001	Jul 24, 1984
	50MG	N88323	001	Jul 28, 1983
	100MG	N88284	001	Aug 25, 1983
	150MG	N88410	001	Mar 05, 1984
	200MG	N88381	001	Mar 14, 1984
	10MG	N88658	001	Mar 26, 1984
	15MG	N88659	001	Mar 26, 1984
	25MG	N88660	001	Mar 26, 1984
	50MG	N88661	001	Mar 26, 1984

THIOTEPA

INJECTABLE; INJECTION

THIOPLEX

IMMUNEX	15MG/VIAL	N20058	001	Dec 22, 1994
THIOTEPA				
IMMUNEX	15MG/VIAL	N11683	001	

THIOTHIXENE

CAPSULE; ORAL

THIOTHIXENE

AM THERAP	1MG	N71884	001	Aug 12, 1987
	2MG	N71885	001	Aug 12, 1987
	5MG	N71886	001	Aug 12, 1987
	10MG	N71887	001	Aug 12, 1987
	20MG	N72200	001	Dec 17, 1987
	2MG	N71626	001	Jun 25, 1987
WATSON LABS	5MG	N71627	001	Jun 25, 1987
	10MG	N71628	001	Jun 25, 1987

THIOTHIXENE HYDROCHLORIDE

CONCENTRATE; ORAL

THIOTHIXENE HYDROCHLORIDE

ALPHARMA US PHARMS	EQ 5MG BASE/ML	N70969	001	Oct 16, 1987
PACO	EQ 1MG BASE/ML	N71917	001	Sep 20, 1989
	EQ 5MG BASE/ML	N71939	001	Dec 16, 1988
TEVA	EQ 5MG BASE/ML	N71184	001	Jun 22, 1987
THIOTHIXENE HYDROCHLORIDE INTENSOL				
ROXANE	EQ 5MG BASE/ML	N73494	001	Jun 30, 1992
INJECTABLE; INJECTION				
NAVANE				
PFIZER	EQ 2MG BASE/ML	N16904	001	

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 258 (of 274)

THYROGLOBULINTABLET; ORAL
PROLOID

PARKE DAVIS

16MG

N02245 009

32MG

N02245 005

65MG

N02245 002

100MG

N02245 008

130MG

N02245 010

200MG

N02245 007

325MG

N02245 004

THYROGLOBULIN

IMPAX LABS

64.8MG

N80151 001

THYROTROPININJECTABLE; INJECTION
THYTROPAR

SANOFI AVENTIS US

10 IU/VIAL

N08682 001

TIAGABINE HYDROCHLORIDETABLET; ORAL
GABITRIL

CEPHALON

20MG

N20646 004

Sep 30, 1997

TICARCILLIN DISODIUMINJECTABLE; INJECTION
TICAR

GLAXOSMITHKLINE

EQ 1GM BASE/VIAL

N50497 001

EQ 3GM BASE/VIAL

N50497 002

EQ 3GM BASE/VIAL

N62690 001

Dec 19, 1986

EQ 6GM BASE/VIAL

N50497 003

EQ 20GM BASE/VIAL

N50497 004

EQ 30GM BASE/VIAL

N50497 005

Apr 04, 1984

TICLOPIDINE HYDROCHLORIDETABLET; ORAL
TICLID

ROCHE PALO

125MG

N19979 001

Mar 24, 1993

TICLOPIDINE HYDROCHLORIDE

WATSON LABS

250MG

N75309 001

Apr 26, 2000

TIMOLOL MALEATESOLUTION/DROPS; OPHTHALMIC
TIMOLOL MALEATE

AKORN

EQ 0.25% BASE

N74465 001

Mar 25, 1997

FOUGERA

EQ 0.25% BASE

N74667 001

Mar 25, 1997

EQ 0.5% BASE

N74668 001

Mar 25, 1997

TABLET; ORAL
BLOCADREN

MERCK

5MG

N18017 001

10MG

N18017 002

20MG

N18017 004

TIMOLOL MALEATE

QUANTUM PHARMICS

5MG

N72466 001

May 19, 1989

10MG

N72467 001

May 19, 1989

20MG

N72468 001

May 19, 1989

TEVA

5MG

N72648 001

Jun 16, 1993

10MG

N72649 001

Jun 16, 1993

20MG

N72650 001

Jun 16, 1993

USL PHARMA

5MG

N72001 001

Apr 11, 1989

10MG

N72002 001

Apr 11, 1989

20MG

N72003 001

Apr 11, 1989

WATSON LABS

5MG

N72269 001

Apr 11, 1989

10MG

N72270 001

Apr 11, 1989

20MG

N72271 001

Apr 11, 1989

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 259 (of 274)

TIOCONAZOLECREAM; TOPICAL
TZ-3

PFIZER

1%

N18682 001

Feb 18, 1983

TOBRAMYCINSOLUTION/DROPS; OPHTHALMIC
TOBRAMYCIN

ALCON UNIVERSAL

0.3%

N63176 001

May 25, 1994

TOBRAMYCIN SULFATEINJECTABLE; INJECTION
NEBCIN

LILLY

EQ 10MG BASE/ML

N50477 005

EQ 10MG BASE/ML

N62008 004

EQ 10MG BASE/ML

N62707 001

Apr 29, 1987

EQ 40MG BASE/ML

N62008 001

EQ 1.2GM BASE/VIAL

N50519 001

TOBRAMYCIN SULFATE

APOTHECON

EQ 40MG BASE/ML

N64026 001

May 31, 1994

ASTRAZENECA

EQ 10MG BASE/ML

N63119 001

Oct 31, 1994

EQ 40MG BASE/ML

N63120 001

Oct 31, 1994

EQ 40MG BASE/ML

N63121 001

Oct 31, 1994

EQ 40MG BASE/ML

N63122 001

Oct 31, 1994

BAXTER HLTHCARE

EQ 10MG BASE/ML

N63113 001

Apr 26, 1991

EQ 10MG BASE/ML

N63128 001

Nov 27, 1991

EQ 40MG BASE/ML

N63127 001

Nov 27, 1991

TOCAINIDE HYDROCHLORIDETABLET; ORAL
TONOCARD

ASTRAZENECA

400MG

N18257 001

Nov 09, 1984

600MG

N18257 002

Nov 09, 1984

TOLAZAMIDETABLET; ORAL
TOLAZAMIDE

BARR

100MG

N70162 001

Jan 14, 1986

250MG

N70163 001

Jan 14, 1986

500MG

N70164 001

Jan 14, 1986

DURAMED PHARMS BARR

100MG

N70165 001

Jan 10, 1986

250MG

N70166 001

Jan 10, 1986

500MG

N70167 001

Jan 10, 1986

INTERPHARM

250MG

N71270 001

Sep 23, 1986

500MG

N71271 001

Sep 23, 1986

SUPERPHARM

250MG

N70763 001

Jun 16, 1986

500MG

N70764 001

Jun 16, 1986

USL PHARMA

100MG

N71355 001

Jan 11, 1988

250MG

N70168 001

Apr 02, 1986

500MG

N70169 001

Apr 02, 1986

WATSON LABS

100MG

N70242 001

Aug 01, 1986

100MG

N70513 001

Jan 09, 1986

250MG

N70243 001

Aug 01, 1986

500MG

N70244 001

Aug 01, 1986

TOLAZOLINE HYDROCHLORIDEINJECTABLE; INJECTION
PRISCOLINE

NOVARTIS

25MG/ML

N06403 005

Feb 22, 1985

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 260 (of 274)

TOLBUTAMIDETABLET; ORAL
ORINASEPHARMACIA AND UPJOHN 250MG
500MGN10670 002
N10670 001

TOLBUTAMIDE

ALRA 500MG
ASCOT 500MG
BARR 500MG
CLONMEL HLTHCARE 500MG
PARKE DAVIS 500MG
PUREPAC PHARM 500MG
SANDOZ 500MG
SUPERPHARM 500MG
VANGARD 500MG
WATSON LABS 250MG
500MGN86141 001
N87541 001 Mar 01, 1983
N87121 001
N86926 001
N86047 001
N88950 001 Jun 17, 1985
N12678 001
N88893 001 Nov 19, 1984
N87876 001 Apr 20, 1982
N89110 001 May 29, 1987
N89111 001 May 29, 1987TOLBUTAMIDE SODIUMINJECTABLE; INJECTION
ORINASE DIAGNOSTIC

PHARMACIA AND UPJOHN EQ 1GM BASE/VIAL

N12095 001

TOLMETIN SODIUMCAPSULE; ORAL
TOLMETIN SODIUMSANDOZ EQ 400MG BASE
TEVA EQ 400MG BASEN73462 001 Apr 30, 1992
N73519 001 May 29, 1992TABLET; ORAL
TOLMETIN SODIUM

TEVA EQ 600MG BASE

N74729 001 Feb 27, 1997

TOPIRAMATECAPSULE; ORAL
TOPAMAX SPRINKLE

ORTHO MCNEIL PHARM 50MG

N20844 003 Oct 26, 1998

TABLET; ORAL

TOPAMAX

ORTHO MCNEIL PHARM 300MG
400MGN20505 003 Dec 24, 1996
N20505 006 Dec 24, 1996TRAMADOL HYDROCHLORIDETABLET; ORAL
TRAMADOL HYDROCHLORIDE

ASTA 50MG

N75974 001 Jul 12, 2002

IVAX PHARMS 50MG

N75963 001 Jul 03, 2002

ULTRAM

ORTHO MCNEIL PHARM 100MG

N20281 001 Mar 03, 1995

TRANEXAMIC ACIDTABLET; ORAL
CYKLOKAPRON

PHARMACIA AND UPJOHN 500MG

N19280 001 Dec 30, 1986

TRAZODONE HYDROCHLORIDETABLET; ORAL
DESYRELAPOTHECON 50MG
100MG
150MG
300MGN18207 001
N18207 002
N18207 003 Mar 25, 1985
N18207 004 Nov 07, 1988

TRAZODONE HYDROCHLORIDE

AM THERAP 50MG
100MGN71139 001 Oct 29, 1986
N71140 001 Oct 29, 1986

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 261 (of 274)

TRAZODONE HYDROCHLORIDE

TABLET; ORAL

TRAZODONE HYDROCHLORIDE

MYLAN	100MG	N71406	001	Feb 27, 1991
QUANTUM PHARMICS	100MG	N70921	001	Dec 01, 1986
USL PHARMA	50MG	N70491	001	Apr 29, 1987
	100MG	N70492	001	Apr 29, 1987
WATSON LABS	50MG	N71112	001	Nov 17, 1986
	100MG	N71113	001	Nov 17, 1986
TRIALODINE				
QUANTUM PHARMICS	50MG	N70942	001	Dec 01, 1986

TRETINOIN

SWAB; TOPICAL

RETIN-A

JOHNSON AND JOHNSON	0.05%	N16921	002	
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TRIAMCINOLONE

TABLET; ORAL

ARISTOCORT

ASTELLAS	1MG	N11161	009	
	2MG	N11161	004	
	4MG	N11161	007	
	8MG	N11161	011	
	16MG	N11161	010	

KENACORT

BRISTOL MYERS SQUIBB	1MG	N11283	003	
	2MG	N11283	008	
	4MG	N11283	006	
	8MG	N11283	010	

TRIAMCINOLONE

BARR	2MG	N84286	001	
	2MG	N84318	001	
	4MG	N84267	001	
	4MG	N84319	001	
	8MG	N84268	001	
	8MG	N84320	001	
IMPAX LABS	4MG	N84340	001	
IVAX PHARMS	4MG	N83750	001	
MYLAN	2MG	N84406	001	
PUREPAC PHARM	2MG	N84020	002	
	4MG	N84020	003	
ROXANE	2MG	N84708	001	
	4MG	N84709	001	
	8MG	N84707	001	
SANDOZ	4MG	N85601	001	
TEVA	4MG	N84775	001	
WATSON LABS	4MG	N84270	001	
	4MG	N85834	001	

TRIAMCINOLONE ACETONIDE

AEROSOL, METERED; NASAL

NASACORT

SANOFI AVENTIS US	0.055MG/INH	N19798	001	Jul 11, 1991
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CREAM; TOPICAL

ARISTOCORT

ASTELLAS	0.025%	N83017	003	
	0.1%	N83016	004	
	0.5%	N83015	002	

ARISTOCORT A

ASTELLAS	0.025%	N83017	004	
	0.025%	N88818	001	Oct 16, 1984
	0.1%	N83016	005	
	0.1%	N88819	001	Oct 16, 1984

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 262 (of 274)

TRIAMCINOLONE ACETONIDE

CREAM; TOPICAL			
ARISTOCORT A			
ASTELLAS	0.5%	N83015 003	
	0.5%	N88820 001	Oct 16, 1984
FLUTEX			
IVAX PHARMS	0.025%	N85539 001	
	0.1%	N85539 002	
	0.5%	N85539 003	
KENALOG -H			
APOTHECON	0.1%	N86240 001	
TRIAECORT			
SOLVAY	0.1%	N87113 001	
TRIAMCINOLONE ACETONIDE			
ALPHARMA US PHARMS	0.025%	N87797 001	Jun 07, 1982
AMBIX	0.025%	N87932 001	May 09, 1983
MORTON GROVE	0.025%	N88094 001	Sep 01, 1983
	0.1%	N88095 001	Sep 01, 1983
	0.5%	N88096 001	Sep 01, 1983
PHARMADERM	0.025%	N87990 001	Jul 07, 1983
	0.1%	N87991 001	Jul 07, 1983
	0.5%	N87992 001	Jul 07, 1983
PHARMAFAIR	0.025%	N87921 001	Aug 10, 1982
	0.1%	N87912 001	Aug 10, 1982
	0.5%	N87922 001	Aug 10, 1982
TARO	0.025%	N40038 001	Oct 26, 1994
TOPIDERM	0.025%	N89274 001	Feb 21, 1989
	0.1%	N89275 001	Feb 21, 1989
	0.5%	N89276 001	Feb 21, 1989
TRIAEX			
IVAX PHARMS	0.025%	N87430 001	Nov 01, 1988
	0.1%	N87429 001	Nov 01, 1988
	0.5%	N87428 001	Nov 01, 1988
TRYMEX			
SAVAGE LABS	0.025%	N88196 001	Mar 25, 1983
	0.1%	N88197 001	Mar 25, 1983
	0.5%	N88198 001	Mar 25, 1983
GEL; TOPICAL			
ARISTOGEL			
ASTELLAS	0.1%	N83380 001	
INJECTABLE; INJECTION			
TRIAMCINOLONE ACETONIDE			
PARNELL	3MG/ML	N19503 001	Oct 16, 1987
WATSON LABS	40MG/ML	N85825 001	
LOTION; TOPICAL			
KENALOG			
BRISTOL MYERS SQUIBB	0.025%	N11602 003	
	0.1%	N11602 001	
TRIAMCINOLONE ACETONIDE			
ALPHARMA US PHARMS	0.025%	N87191 001	Sep 08, 1982
	0.1%	N87192 001	Sep 08, 1982
OINTMENT; TOPICAL			
ARISTOCORT			
ASTELLAS	0.1%	N80750 004	
	0.5%	N80745 002	
ARISTOCORT A			
ASTELLAS	0.1%	N80750 003	
	0.1%	N88780 001	Oct 01, 1984
	0.5%	N80745 003	
	0.5%	N88781 001	Oct 05, 1984
FLUTEX			
IVAX PHARMS	0.025%	N87375 001	Nov 01, 1988
	0.1%	N87377 001	Nov 01, 1988
	0.5%	N87376 001	Nov 01, 1988

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 263 (of 274)

TRIAMCINOLONE ACETONIDE

OINTMENT; TOPICAL

KENALOG

APOTHECON 0.5%

N83944 001

TRIAMCINOLONE ACETONIDE

ALPHARMA US PHARMS 0.5%

N89913 001 Dec 23, 1988

G AND W LABS 0.025%

N89795 001 Dec 23, 1988

0.1%

N89796 001 Dec 23, 1988

MORTON GROVE 0.025%

N88090 001 Sep 01, 1983

0.1%

N88091 001 Sep 01, 1983

0.5%

N88092 001 Sep 01, 1983

PHARMADERM 0.025%

N88692 001 Aug 02, 1984

0.1%

N88690 001 Aug 02, 1984

TRYMEX

SAVAGE LABS 0.025%

N88693 001 Aug 02, 1984

0.1%

N88691 001 Aug 02, 1984

SPRAY, METERED; NASAL

NASACORT HFA

SANOFI AVENTIS US 0.055MG/SPRAY

N20784 001 Apr 07, 2004

TRI-NASAL

COLLEGIUM 0.05MG/SPRAY

N20120 001 Feb 04, 2000

TRIAMCINOLONE DIACETATE

INJECTABLE; INJECTION

ARISTOCORT

SANDOZ 25MG/ML

N11685 003

40MG/ML

N12802 001

TRIAMCINOLONE DIACETATE

AKORN 25MG/ML

N85122 001

40MG/ML

N86394 001

WATSON LABS 40MG/ML

N84072 001

40MG/ML

N85529 001

SYRUP; ORAL

ARISTOCORT

ASTELLAS 2MG/5ML

N11960 004

KENACORT

BRISTOL MYERS SQUIBB EQ 4MG BASE/5ML

N12515 001

TRIAZOLAM

TABLET; ORAL

HALCION

PHARMACIA AND UPJOHN 0.5MG

N17892 002 Nov 15, 1982

TRICHLORMETHIAZIDE

TABLET; ORAL

METAHYDRIN

SANOFI AVENTIS US 2MG

N12594 001 Jun 16, 1988

4MG

N12594 002 Jun 16, 1988

NAQUA

SCHERING 2MG

N12265 001

4MG

N12265 002

TRICHLOREX

LANNETT 4MG

N83436 001

4MG

N85630 001

TRICHLORMAS

MAST MM 4MG

N86259 001

TRICHLORMETHIAZIDE

IMPAX LABS 4MG

N83967 001

PAR PHARM 2MG

N87007 001

4MG

N87005 001

SANDOZ 4MG

N86171 001

TG UNITED LABS 4MG

N85568 001

WATSON LABS 2MG

N83847 001

2MG

N86458 001

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 264 (of 274)

TRICHLORMETHIAZIDE

TABLET; ORAL				
TRICHLORMETHIAZIDE				
WATSON LABS	4MG	N83462	001	
	4MG	N83855	001	
	4MG	N85962	001	

TRICLOFOS SODIUM

SOLUTION; ORAL				
TRICLOS				
SANOFI AVENTIS US	1.5GM/15ML	N16830	001	
TABLET; ORAL				
TRICLOS				
SANOFI AVENTIS US	750MG	N16809	002	

TRIDIHEXETHYL CHLORIDE

INJECTABLE; INJECTION				
PATHILON				
LEDERLE	10MG/ML	N09729	001	
TABLET; ORAL				
PATHILON				
LEDERLE	25MG	N09489	005	

TRIFLUOPERAZINE HYDROCHLORIDE

CONCENTRATE; ORAL				
TRIFLUOPERAZINE HYDROCHLORIDE				
MORTON GROVE	EQ 10MG BASE/ML	N88143	001	Jul 26, 1983
SANDOZ	EQ 10MG BASE/ML	N85787	001	Apr 15, 1982
INJECTABLE; INJECTION				
STELAZINE				
GLAXOSMITHKLINE	EQ 2MG BASE/ML	N11552	005	
TABLET; ORAL				
TRIFLUOPERAZINE HYDROCHLORIDE				
DURAMED PHARMS BARR	EQ 1MG BASE	N88967	001	Apr 23, 1985
	EQ 2MG BASE	N88968	001	Apr 23, 1985
	EQ 5MG BASE	N88969	001	Apr 23, 1985
	EQ 10MG BASE	N88970	001	Apr 23, 1985
IVAX PHARMS	EQ 1MG BASE	N87612	001	Nov 19, 1982
	EQ 2MG BASE	N87613	001	Nov 19, 1982
	EQ 5MG BASE	N87328	001	Nov 19, 1982
	EQ 10MG BASE	N87614	001	Nov 19, 1982
WATSON LABS	EQ 1MG BASE	N85975	001	Jun 23, 1988
	EQ 2MG BASE	N85976	001	Jun 23, 1988
	EQ 5MG BASE	N85973	001	Jun 23, 1988
	EQ 10MG BASE	N88710	001	Jun 23, 1988

TRIFLUPROMAZINE

SUSPENSION; ORAL				
VESPRIN				
APOTHECON	EQ 50MG HCL/5ML	N11491	004	

TRIFLUPROMAZINE HYDROCHLORIDE

INJECTABLE; INJECTION				
VESPRIN				
APOTHECON	3MG/ML	N11325	005	
	10MG/ML	N11325	004	
	20MG/ML	N11325	001	
TABLET; ORAL				
VESPRIN				
BRISTOL MYERS SQUIBB	10MG	N11123	001	
	25MG	N11123	002	
	50MG	N11123	003	

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 265 (of 274)

TRIHEXYPHENIDYL HYDROCHLORIDE

CAPSULE, EXTENDED RELEASE; ORAL

ARTANE

LEDERLE

5MG

N06773 010

5MG

N12947 001

ELIXIR; ORAL

ARTANE

LEDERLE

2MG/5ML

N06773 009

TABLET; ORAL

ARTANE

LEDERLE

2MG

N06773 005

5MG

N06773 003

TREMIN

SCHERING

2MG

N80381 001

5MG

N80381 003

TRIHEXYPHENIDYL HYDROCHLORIDE

NYLOS

5MG

N85622 001

VANGARD

2MG

N88035 001

WATSON LABS

2MG

N85117 001

5MG

N85105 001

Jul 30, 1982

TRILOSTANE

CAPSULE; ORAL

MODRASTANE

BIOENVISION

30MG

N18719 002

Dec 31, 1984

60MG

N18719 001

Dec 31, 1984

TRIMEPRAZINE TARTRATE

CAPSULE, EXTENDED RELEASE; ORAL

TEMARIL

ALLERGAN HERBERT

EQ 5MG BASE

N11316 004

SYRUP; ORAL

TEMARIL

ALLERGAN HERBERT

EQ 2.5MG BASE/5ML

N11316 003

TRIMEPRAZINE TARTRATE

ALPHARMA US PHARMS

EQ 2.5MG BASE/5ML

N85015 001

Feb 18, 1982

MORTON GROVE

EQ 2.5MG BASE/5ML

N88285 001

Apr 11, 1985

TABLET; ORAL

TEMARIL

ALLERGAN HERBERT

EQ 2.5MG BASE

N11316 001

TRIMETHADIONE

CAPSULE; ORAL

TRIDIONE

ABBOTT

300MG

N05856 005

SOLUTION; ORAL

TRIDIONE

ABBOTT

200MG/5ML

N05856 002

TRIMETHAPHAN CAMSYLATE

INJECTABLE; INJECTION

ARFONAD

ROCHE

50MG/ML

N08983 001

TRIMETHOBENZAMIDE HYDROCHLORIDE

INJECTABLE; INJECTION

TRIMETHOBENZAMIDE HYDROCHLORIDE

SMITH AND NEPHEW

100MG/ML

N88960 001

Apr 04, 1986

100MG/ML

N89043 001

Apr 04, 1986

SOLOPAK

100MG/ML

N89094 001

Apr 04, 1986

WATSON LABS

100MG/ML

N86577 001

Oct 19, 1982

100MG/ML

N87939 001

Dec 28, 1982

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 266 (of 274)

TRIMETHOPRIMTABLET; ORAL
PROLOPRIM

MONARCH PHARMS 200MG N17943 003 Jul 14, 1982

TRIMETHOPRIM

MUTUAL PHARM 100MG N70494 001 Jan 22, 1986

200MG N70495 001 Sep 24, 1986

TRIMPEX

ROCHE 100MG N17952 001

TRIMPEX 200

ROCHE 200MG N17952 002 Nov 09, 1982

TRIMETHOPRIM HYDROCHLORIDESOLUTION; ORAL
PRIMSOL

FSC EQ 25MG BASE/5ML N74374 001 Jun 23, 1995

TRIMIPRAMINE MALEATE

CAPSULE; ORAL

TRIMIPRAMINE MALEATE

USL PHARMA EQ 25MG BASE N71283 001 Dec 08, 1987

EQ 50MG BASE N71284 001 Dec 08, 1987

EQ 100MG BASE N71285 001 Dec 08, 1987

TRIOXSALENTABLET; ORAL
TRISORALEN

VALEANT PHARM INTL 5MG N12697 001

TRIPLENNAMINE CITRATEELIXIR; ORAL
PBZ

NOVARTIS EQ 25MG HCL/5ML N05914 004

TRIPLENNAMINE HYDROCHLORIDETABLET; ORAL
PBZ

NOVARTIS 25MG N83149 001

50MG N05914 002

TRIPLENNAMINE HYDROCHLORIDE

ANABOLIC 50MG N83037 001

BARR 50MG N80744 001

HEATHER 50MG N83989 001

IMPAX LABS 50MG N80785 001

LANNETT 50MG N83557 001

NYLOS 50MG N85412 001

PARKE DAVIS 25MG N83625 001

50MG N83626 001

WATSON LABS 50MG N80713 001

50MG N80790 001

50MG N85188 001

TABLET, EXTENDED RELEASE; ORAL

PBZ-SR

NOVARTIS 50MG N10533 002

100MG N10533 001

TRIPLE SULFA (SULFABENZAMIDE;SULFACETAMIDE;SULFATHIAZOLE)CREAM; VAGINAL
GYNE-SULF

G AND W LABS 3.7%;2.86%;3.42% N88607 001 Jun 09, 1986

SULTRIN

ORTHO MCNEIL PHARM 3.7%;2.86%;3.42% N05794 001

TRIPLE SULFA

ALPHARMA US PHARMS 3.7%;2.86%;3.42% N87864 001 Sep 01, 1982

DISCONTINUED DRUG PRODUCT LIST

6 - 267 (of 274)

TRIPLE SULFA (SULFABENZAMIDE;SULFACETAMIDE;SULFATHIAZOLE)

CREAM; VAGINAL				
TRIPLE SULFA				
FOUGERA	3.7%;2.86%;3.42%	N86424	001	
PERRIGO NEW YORK	3.7%;2.86%;3.42%	N87285	001	Nov 15, 1982
TRYSQL				
SAVAGE LABS	3.7%;2.86%;3.42%	N87887	001	Jul 23, 1982
VAGILIA				
TEVA	3.7%;2.86%;3.42%	N88821	001	Nov 09, 1987
TABLET; VAGINAL				
SULTRIN				
ORTHO MCNEIL PHARM	184MG;143.75MG;172.5MG	N05794	002	
TRIPLE SULFA				
FOUGERA	184MG;143.75MG;172.5MG	N88463	001	Jan 03, 1985
PHARMADERM	184MG;143.75MG;172.5MG	N88462	001	Jan 03, 1985

TRIPROLIDINE HYDROCHLORIDE

SYRUP; ORAL				
ACTIDIL				
GLAXOSMITHKLINE	1.25MG/5ML	N11496	002	Jul 01, 1983
MYIDYL				
USL PHARMA	1.25MG/5ML	N87963	001	Jan 18, 1983
TRIPROLIDINE HYDROCHLORIDE				
ALPHARMA US PHARMS	1.25MG/5ML	N85940	001	
HALSEY	1.25MG/5ML	N88735	001	Jan 17, 1985
PHARM ASSOC	1.25MG/5ML	N87514	001	Feb 10, 1982
TABLET; ORAL				
ACTIDIL				
GLAXOSMITHKLINE	2.5MG	N11110	002	Jul 01, 1983
TRIPROLIDINE HYDROCHLORIDE				
VITARINE	2.5MG	N85610	001	
WATSON LABS	2.5MG	N85094	001	

TRISULFAPYRIMIDINES (SULFADIAZINE;SULFAMERAZINE;SULFAMETHAZINE)

SUSPENSION; ORAL				
LANTRISUL				
LANNETT	167MG/5ML;167MG/5ML;167MG/5ML	N80123	002	
NEOTRIZINE				
LILLY	167MG/5ML;167MG/5ML;167MG/5ML	N06317	012	
SULFALOID				
FOREST PHARMS	167MG/5ML;167MG/5ML;167MG/5ML	N80100	001	
SULFOSE				
WYETH AYERST	167MG/5ML;167MG/5ML;167MG/5ML	N80013	002	
TERFONYL				
BRISTOL MYERS SQUIBB	167MG/5ML;167MG/5ML;167MG/5ML	N06904	002	
TRIPLE SULFA				
ALPHARMA US PHARMS	167MG/5ML;167MG/5ML;167MG/5ML	N80280	001	
TRIPLE SULFAS				
LEDERLE	167MG/5ML;167MG/5ML;167MG/5ML	N06920	003	
TABLET; ORAL				
NEOTRIZINE				
LILLY	167MG;167MG;167MG	N06317	011	
SULFALOID				
FOREST PHARMS	167MG;167MG;167MG	N80099	001	
SULFA-TRIPLE #2				
IMPAX LABS	167MG;167MG;167MG	N80079	001	
SULFOSE				
WYETH AYERST	167MG;167MG;167MG	N80013	001	
TERFONYL				
BRISTOL MYERS SQUIBB	167MG;167MG;167MG	N06904	001	
TRIPLE SULFA				
PUREPAC PHARM	167MG;167MG;167MG	N80086	001	
TRIPLE SULFAS				
LEDERLE	167MG;167MG;167MG	N06920	002	

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 268 (of 274)

TRISULFAPYRIMIDINES (SULFADIAZINE;SULFAMERAZINE;SULFAMETHAZINE)

TABLET; ORAL				
TRIPLE SULFOID				
PAL PAK	167MG;167MG;167MG	N80094	001	

TROGLITAZONE

TABLET; ORAL				
PRELAY				
SANKYO	200MG	N20719	001	Jan 29, 1997
	300MG	N20719	003	Aug 04, 1997
	400MG	N20719	002	Jan 29, 1997
REZULIN				
PFIZER PHARMS	200MG	N20720	001	Jan 29, 1997
	300MG	N20720	003	Aug 04, 1997
	400MG	N20720	002	Jan 29, 1997

TROLAMINE POLYPEPTIDE OLEATE CONDENSATE

SOLUTION/DROPS; OTIC				
CERUMENEX				
PURDUE FREDERICK	10%	N11340	002	

TROLEANDOMYCIN

CAPSULE; ORAL				
TAO				
PFIZER	EQ 250MG BASE	N50336	002	
SUSPENSION; ORAL				
TAO				
PFIZER	EQ 125MG BASE/5ML	N50332	001	

TROPICAMIDE

SOLUTION/DROPS; OPHTHALMIC				
MYDRIACYL				
ALCON	0.5%	N12111	002	
	1%	N12111	004	
MYDRIAFAIR				
PHARMAFAIR	0.5%	N88274	001	Sep 16, 1983
	1%	N88230	001	Sep 16, 1983
TROPICAMIDE				
AKORN	1%	N88447	001	Aug 28, 1985
WATSON LABS	0.5%	N89171	001	Dec 28, 1990

TROVAFLOXACIN MESYLATE

TABLET; ORAL				
TROVAN				
PFIZER	EQ 100MG BASE	N20759	001	Dec 18, 1997
	EQ 200MG BASE	N20759	002	Dec 18, 1997

TUBOCURARINE CHLORIDE

INJECTABLE; INJECTION				
TUBOCURARINE CHLORIDE				
BRISTOL MYERS SQUIBB	3MG/ML	N05657	001	
HOSPIRA	3MG/ML	N06095	001	
LILLY	3MG/ML	N06325	001	

TYROPANOATE SODIUM

CAPSULE; ORAL				
BILOPAQUE				
GE HEALTHCARE	750MG	N13731	001	

UNDECOYLUM CHLORIDE; UNDECOYLUM CHLORIDE IODINE COMPLEX

SOLUTION; TOPICAL				
VIRAC REX				
CHESEBROUGH PONDS	0.5%;1.8%	N11914	001	

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 269 (of 274)

UNOPROSTONE ISOPROPYLSOLUTION/DROPS; OPHTHALMIC
RESCULA

R TECH UENO LTD 0.15%

N21214 001 Aug 03, 2000

URACIL MUSTARDCAPSULE; ORAL
URACIL MUSTARD
SHIRE

1MG

N12892 001

UREAINJECTABLE; INJECTION
STERILE UREA

HOSPIRA 40GM/VIAL

N17698 001

UREAPHIL

HOSPIRA 40GM/VIAL

N12154 001

UREA, C-13FOR SOLUTION; ORAL
HELICOSOL

METABOLIC SOLUTIONS 125MG/VIAL

N21092 001 Dec 17, 1999

MERETEK UBT KIT (W/ PRANACTIN)

MERETEK 125MG/VIAL

N20586 001 Sep 17, 1996

PYLORI-CHEK BREATH TEST

DXS DEVICES 100MG/VIAL

N20900 001 Feb 04, 1999

UROFOLLITROPININJECTABLE; INTRAMUSCULAR
METRODIN

SERONO 75 IU/AMP

N19415 002 Sep 18, 1986

150 IU/AMP

N19415 003 Sep 18, 1986

INJECTABLE; SUBCUTANEOUS

FERTINEX

SERONO 75 IU/AMP

N19415 005 Aug 23, 1996

150 IU/AMP

N19415 004 Aug 23, 1996

UROKINASEINJECTABLE; INJECTION
ABBOKINASE

IMARX THERAP 5,000 IU/VIAL

N21846 003

9,000 IU/VIAL

N21846 002

URSODIOLCAPSULE; ORAL
ACTIGALL

WATSON PHARMS 150MG

N19594 001 Dec 31, 1987

VALPROIC ACIDCAPSULE; ORAL
VALPROIC ACID

PAR PHARM 250MG

N70431 001 Feb 28, 1986

SCHERER RP 250MG

N70195 001 Jul 02, 1987

VALSARTANCAPSULE; ORAL
DIOVAN

NOVARTIS 80MG

N20665 001 Dec 23, 1996

160MG

N20665 002 Dec 23, 1996

VANCOMYCIN HYDROCHLORIDEFOR SOLUTION; ORAL
VANCOCIN HYDROCHLORIDE

VIROPHARMA EQ 250MG BASE/5ML

N61667 002 Jul 13, 1983

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 270 (of 274)

VANCOMYCIN HYDROCHLORIDE

FOR SOLUTION; ORAL

VANOCIN HYDROCHLORIDE

VIOPHARMA

EQ 500MG BASE/6ML

N61667 001

VANCOLED

LEDERLE

EQ 250MG BASE/5ML

N63321 002

Oct 15, 1993

EQ 500MG BASE/6ML

N63321 003

Oct 15, 1993

INJECTABLE; INJECTION

VANOCIN HYDROCHLORIDE

VIOPHARMA

EQ 500MG BASE/VIAL

N60180 001

EQ 500MG BASE/VIAL

N62476 001

Mar 15, 1984

EQ 500MG BASE/VIAL

N62716 001

Mar 13, 1987

EQ 500MG BASE/VIAL

N62812 001

Nov 17, 1987

EQ 1GM BASE/VIAL

N60180 002

Mar 21, 1986

EQ 1GM BASE/VIAL

N62476 002

Mar 21, 1986

EQ 1GM BASE/VIAL

N62716 002

Mar 13, 1987

EQ 1GM BASE/VIAL

N62812 002

Nov 17, 1987

EQ 10GM BASE/VIAL

N62812 003

Nov 17, 1987

VANCOLED

BAXTER HLTHCARE

EQ 500MG BASE/VIAL

N62682 001

Jul 22, 1986

EQ 1GM BASE/VIAL

N62682 002

Mar 30, 1988

EQ 2GM BASE/VIAL

N62682 003

May 11, 1988

EQ 5GM BASE/VIAL

N62682 004

May 11, 1988

EQ 10GM BASE/VIAL

N62682 005

May 11, 1988

VANCOMYCIN HYDROCHLORIDE

BAXTER HLTHCARE

EQ 500MG BASE/VIAL

N62879 001

Aug 02, 1988

EQ 1GM BASE/VIAL

N62879 002

Aug 02, 1988

VANCOR

PHARMACIA AND UPJOHN

EQ 500MG BASE/VIAL

N62956 001

Aug 01, 1988

EQ 1GM BASE/VIAL

N62956 002

Aug 01, 1988

VASOPRESSIN TANNATE

INJECTABLE; INJECTION

PITRESSIN TANNATE

PARKE DAVIS

5PRESSOR UNITS/ML

N03402 001

VECURONIUM BROMIDE

INJECTABLE; INJECTION

NORCURON

ORGANON USA INC

10MG/VIAL

N18776 002

Apr 30, 1984

20MG/VIAL

N18776 003

Jan 03, 1992

VENLAFAXINE HYDROCHLORIDE

CAPSULE, EXTENDED RELEASE; ORAL

EFFEXOR XR

WYETH PHARMS INC

EQ 100MG BASE

N20699 003

Oct 20, 1997

TABLET; ORAL

EFFEXOR

WYETH PHARMS INC

EQ 12.5MG BASE

N20151 001

Dec 28, 1993

VERAPAMIL HYDROCHLORIDE

INJECTABLE; INJECTION

CALAN

GD SEARLE LLC

2.5MG/ML

N18925 001

Mar 30, 1984

2.5MG/ML

N19038 001

Mar 30, 1984

VERAPAMIL HYDROCHLORIDE

ABRAXIS PHARM

2.5MG/ML

N70348 001

May 01, 1986

HOSPIRA

2.5MG/ML

N70739 001

May 06, 1987

2.5MG/ML

N70740 001

May 06, 1987

MARSAM PHARMS LLC

2.5MG/ML

N72233 001

Feb 26, 1993

2.5MG/ML

N73485 001

Sep 27, 1993

SMITH AND NEPHEW

2.5MG/ML

N70696 001

Jul 31, 1987

2.5MG/ML

N70697 001

Jul 31, 1987

SOLOPAK

2.5MG/ML

N70695 001

Jul 31, 1987

DISCONTINUED DRUG PRODUCT LIST

6 - 271 (of 274)

VERAPAMIL HYDROCHLORIDE

TABLET; ORAL
CALAN

GD SEARLE LLC	160MG	N18817	004	Feb 23, 1988
VERAPAMIL HYDROCHLORIDE				
MUTUAL PHARM	80MG	N70482	001	Sep 24, 1986
	120MG	N70483	001	Sep 24, 1986
PLIVA	40MG	N72751	001	Feb 23, 1996
RADIUS PHARMS	80MG	N71880	001	Apr 05, 1988
	120MG	N71881	001	Apr 05, 1988
WARNER CHILCOTT	80MG	N70340	001	Sep 24, 1986
	120MG	N70341	001	Sep 24, 1986
WATSON LABS	40MG	N72799	001	Apr 28, 1989

VERATRUM VIRIDE

TABLET; ORAL
VERTAVIS

MEDPOINTE PHARM HLC	130CSR UNIT	N05691	002	
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VIDARABINE

INJECTABLE; INJECTION
VIRA-A

PARKEDALE	EQ 187.4MG BASE/ML	N50523	001	
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OINTMENT; OPHTHALMIC
VIRA-A

PARKEDALE	3%	N50486	001	
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VINBLASTINE SULFATE

INJECTABLE; INJECTION
VELBAN

LILLY	10MG/VIAL	N12665	001	
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VINBLASTINE SULFATE

ABRAXIS PHARM	10MG/VIAL	N89011	001	Nov 18, 1985
MAYNE PHARMA USA	10MG/VIAL	N89565	001	Aug 18, 1987

VINCRISTINE SULFATE

INJECTABLE; INJECTION
ONCOVIN

LILLY	1MG/VIAL	N14103	001	
	1MG/ML	N14103	003	Mar 07, 1984
	5MG/VIAL	N14103	002	

VINCASAR PFS

SICOR PHARMS	1MG/ML	N71426	001	Jul 17, 1987
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VINCAREX

BRISTOL MYERS SQUIBB	5MG/VIAL	N70867	001	Jul 12, 1988
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VINCRISTINE SULFATE

ABIC	1MG/ML	N70873	001	Feb 19, 1987
ABRAXIS PHARM	1MG/ML	N70411	001	Sep 10, 1986
MAYNE PHARMA USA	1MG/VIAL	N71559	001	Apr 11, 1988
	2MG/VIAL	N71560	001	Apr 11, 1988
	5MG/VIAL	N71561	001	Apr 11, 1988

VIOMYCIN SULFATE

INJECTABLE; INJECTION
VIOCIN SULFATE

PFIZER	EQ 1GM BASE/VIAL	N61086	001	
	EQ 5GM BASE/VIAL	N61086	002	

VITAMIN A

CAPSULE; ORAL
AQUASOL A

ASTRAZENECA	25,000USP UNITS	N83080	002	
	50,000USP UNITS	N83080	001	

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 272 (of 274)

VITAMIN A

CAPSULE; ORAL

VITAMIN A

BANNER PHARMACAPS	50,000USP UNITS	N83973	001
CHASE CHEM	50,000 IU	N83351	001
EVERYLIFE	50,000 IU	N83134	001
IMPAX LABS	50,000USP UNITS	N80952	001
WEST WARD	50,000USP UNITS	N80985	001

VITAMIN A PALMITATE

CAPSULE; ORAL

AFAXIN

STERLING WINTHROP	EQ 50,000 UNITS BASE	N83187	001
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ALPHALIN

LILLY	EQ 50,000 UNITS BASE	N80883	001
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DEL-VI-A

DEL RAY LABS	EQ 50,000 UNITS BASE	N80830	001
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VI-DOM-A

BAYER PHARMS	EQ 50,000 UNITS BASE	N80972	001
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VITAMIN A

BANNER PHARMACAPS	EQ 50,000 UNITS BASE	N80702	001
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BRISTOL MYERS SQUIBB	EQ 50,000 UNITS BASE	N80860	001
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CHASE CHEM	EQ 50,000 UNITS BASE	N80746	001
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	EQ 50,000 UNITS BASE	N83207	001
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ELKINS SINN	EQ 50,000 UNITS BASE	N85479	001
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EVERYLIFE	EQ 50,000 UNITS BASE	N80943	001
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	EQ 50,000 UNITS BASE	N83114	001
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IMPAX LABS	EQ 50,000 UNITS BASE	N80953	001
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	EQ 50,000 UNITS BASE	N80955	001
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IVAX PHARMS	EQ 50,000 UNITS BASE	N83035	001
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	EQ 50,000 UNITS BASE	N83190	001
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MK LABS	EQ 25,000 UNITS BASE	N83457	002
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	EQ 50,000 UNITS BASE	N83457	001
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WEST WARD	EQ 50,000 UNITS BASE	N80967	001
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WHARTON LABS	EQ 50,000 UNITS BASE	N83665	001
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VITAMIN A PALMITATE

BANNER PHARMACAPS	EQ 50,000 UNITS BASE	N83948	001
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	EQ 50,000 UNITS BASE	N83981	001
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VITAMIN A SOLUBILIZED

TEVA	EQ 50,000 UNITS BASE	N80921	001
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INJECTABLE; INJECTION

VITAMIN A PALMITATE

BEL MAR	EQ 50,000 UNITS BASE/ML	N80819	001
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WARFARIN POTASSIUM

TABLET; ORAL

ATHROMBIN-K

PURDUE FREDERICK	2MG	N11771	007
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	5MG	N11771	004
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	10MG	N11771	005
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	25MG	N11771	006
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WARFARIN SODIUM

INJECTABLE; INJECTION

COUMADIN

BRISTOL MYERS SQUIBB	50MG/VIAL	N09218	020
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	75MG/VIAL	N09218	012
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TABLET; ORAL

ATHROMBIN

PURDUE FREDERICK	5MG	N11771	003
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	10MG	N11771	002
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	25MG	N11771	001
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PANWARFIN

ABBOTT	2MG	N17020	001
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DISCONTINUED DRUG PRODUCT LIST

6 - 273 (of 274)

WARFARIN SODIUM

TABLET; ORAL				
PANWARFIN				
ABBOTT	2.5MG	N17020	002	
	5MG	N17020	003	
	7.5MG	N17020	004	
	10MG	N17020	005	
WARFARIN SODIUM				
USL PHARMA	2MG	N88719	001	Jun 27, 1985
	2.5MG	N88720	001	Aug 06, 1985
	5MG	N88721	001	Jul 02, 1985
WATSON LABS	2MG	N86123	001	Aug 17, 1982
	2.5MG	N86120	001	Aug 17, 1982
	5MG	N86119	001	Aug 17, 1982
	7.5MG	N86118	001	Aug 17, 1982
	10MG	N86122	001	Aug 17, 1982

WATER FOR INJECTION, STERILE

LIQUID; N/A				
BACTERIOSTATIC WATER FOR INJECTION IN PLASTIC CONTAINER				
ABRAXIS PHARM	100%	N89099	001	Dec 29, 1987
	100%	N89100	001	Dec 29, 1987
STERILE WATER FOR INJECTION IN PLASTIC CONTAINER				
B BRAUN	100%	N19077	001	Mar 02, 1984

WATER FOR IRRIGATION, STERILE

LIQUID; IRRIGATION				
STERILE WATER IN PLASTIC CONTAINER				
MILES	100%	N18246	001	

XENON, XE-127

GAS; INHALATION				
XENON XE 127				
MALLINCKRODT	5mCi/VIAL	N18536	001	Oct 01, 1982
	10mCi/VIAL	N18536	002	Oct 01, 1982

XENON, XE-133

GAS; INHALATION				
XENON XE 133				
GE HEALTHCARE	1 CI/AMP	N17256	002	
	10mCi/VIAL	N17687	002	
	20mCi/VIAL	N17687	003	
GEN ELECTRIC	5-100 CI/CYLINDER	N17550	001	
	0.25-5 CI/AMP	N17550	003	
XENON XE 133-V.S.S.				
GE HEALTHCARE	10mCi/VIAL	N17687	001	
INJECTABLE; INJECTION				
XENON XE 133				
BRISTOL MYERS SQUIBB	6.3mCi/ML	N17283	001	
GE HEALTHCARE	1.3-1.7 CI/AMP	N17256	001	
SOLUTION; INHALATION, INJECTION				
XENEISOL				
MALLINCKRODT	18-25mCi/AMP	N17262	002	

XYLOSE

POWDER; ORAL				
XYLO-PFAN				
SAVAGE LABS	25GM/BOT	N17605	001	
XYLOSE				
LYNE	25GM/BOT	N18856	001	Mar 26, 1987

27TH EDITION - 2007 - APPROVED DRUG PRODUCTS LIST

DISCONTINUED DRUG PRODUCT LIST

6 - 274 (of 274)

ZICONOTIDEINJECTABLE; INTRATHECAL
PRIALT

ELAN PHARMS

200UGM/2ML (100MG/ML)

N21060 003

Dec 28, 2004

ZIDOVUDINETABLET; ORAL
RETROVIR

GLAXOSMITHKLINE

200MG

N20518 001

Dec 19, 1995

ZILEUTONTABLET; ORAL
ZYFLO

CRITICAL

300MG

N20471 001

Dec 09, 1996

ZINC SULFATEINJECTABLE; INJECTION
ZINC SULFATE

ABRAXIS PHARM

EQ 1MG ZINC/ML

N19229 002

May 05, 1987

ORPHAN PRODUCT DESIGNATIONS AND APPROVALS LIST

The list of List of Orphan Designations and Approvals is available at:
<http://www.fda.gov/orphan/designat/list.htm>

**DRUG PRODUCTS WHICH MUST DEMONSTRATE *IN VIVO* BIOAVAILABILITY
ONLY IF PRODUCT FAILS TO ACHIEVE ADEQUATE DISSOLUTION**

ACETAMINOPHEN; ASPIRIN;
BUTALBITAL
CAPSULE OR TABLET; ORAL
160-165MG; 160-165MG; 50MG

ASPIRIN; CAFFEINE;
CARISOPRODOL
TABLET; ORAL
160MG; 32MG; 200MG

ACETAMINOPHEN; ASPIRIN;
BUTALBITAL
CAPSULE OR TABLET; ORAL
325MG; 325MG; 50MG

ASPIRIN; CAFFEINE;
CARISOPRODOL;
CODEINE PHOSPHATE
TABLET; ORAL
160MG; 32MG;
200MG; 16MG

ACETAMINOPHEN; ASPIRIN;
BUTALBITAL; CAFFEINE
CAPSULE OR TABLET; ORAL
160-165MG; 160-165MG; 50MG; 40MG

ASPIRIN;
CARISOPRODOL
TABLET; ORAL
325MG; 200MG

ACETAMINOPHEN; ASPIRIN;
BUTALBITAL; CAFFEINE
CAPSULE OR TABLET; ORAL
325MG; 325MG; 50MG; 40MG

ASPIRIN;
CARISOPRODOL;
CODEINE PHOSPHATE
TABLET; ORAL
325MG; 200MG; 16MG

ACETAMINOPHEN; BUTALBITAL
CAPSULE OR TABLET; ORAL
325MG; 50MG
650MG; 50MG

ASPIRIN; MEPROBAMATE
TABLET; ORAL
325MG; 200MG

ACETAMINOPHEN; BUTALBITAL;
CAFFEINE
CAPSULE OR TABLET; ORAL
325MG; 50MG; 40MG
650MG; 50MG; 40MG
500MG; 50MG; 40MG

ASPIRIN;
METHOCARBAMOL
TABLET; ORAL
325MG; 400MG

AMINOPHYLLINE
TABLET; ORAL
100MG; 200MG

CHLOROTHIAZIDE
TABLET; ORAL
250MG

ASPIRIN; BUTALBITAL
CAPSULE OR TABLET; ORAL
325MG; 50MG
650MG; 50MG

HYDROXYZINE
HYDROCHLORIDE
TABLET; ORAL
10MG; 25MG;
50MG; 100MG

ASPIRIN; BUTALBITAL; CAFFEINE
CAPSULE OR TABLET; ORAL
325MG; 50MG; 40MG
650MG; 50MG; 40MG

PREDNISONE
TABLET; ORAL
1MG; 2.5MG;
5MG; 10MG;
20MG; 25MG; 50MG

APPENDIX A - PRODUCT NAME INDEX

A - 1

**** 8 ****

8-MOP, METHOXSALEN

**** A ****

A/T/S, ERYTHROMYCIN
ABBOKINASE, UROKINASE
ABELCET, AMPHOTERICIN B
ABILIFY, ARIPIPRAZOLE
ABRAXANE, PACLITAXEL
ABREVA, DOCOSANOL (OTC)
ACCOLATE, ZAFIRLUKAST
ACCUNEB, ALBUTEROL SULFATE
ACCUPRIL, QUINAPRIL HYDROCHLORIDE
ACCURETIC, HYDROCHLOROTHIAZIDE
AC CUTANE, ISOTRETINOIN
ACEBUTOLOL HYDROCHLORIDE, ACEBUTOLOL HYDROCHLORIDE
ACEON, PERINDOPRIL ERBUMINE
ACEPHEN, ACETAMINOPHEN (OTC)
ACETADOTE, ACETYLCYSTEINE
ACETAMINOPHEN AND CODEINE PHOSPHATE, ACETAMINOPHEN
ACETAMINOPHEN AND PENTAZOCINE HYDROCHLORIDE, ACETAMINOPHEN
ACETAMINOPHEN W/ CODEINE PHOSPHATE #3, ACETAMINOPHEN
ACETAMINOPHEN W/ CODEINE, ACETAMINOPHEN
ACETAMINOPHEN, ACETAMINOPHEN (OTC)
ACETAMINOPHEN, ASPIRIN AND CAFFEINE, ACETAMINOPHEN (OTC)
ACETAMINOPHEN, ASPIRIN, AND CODEINE PHOSPHATE, ACETAMINOPHEN
ACETAMINOPHEN, BUTALBITAL, AND CAFFEINE, ACETAMINOPHEN
ACETAMINOPHEN, BUTALBITAL, CAFFEINE, AND CODEINE PHOSPHATE, ACETAMINOPHEN
ACETAMINOPHEN, CAFFEINE, AND DIHYDROCODEINE BITARTRATE, ACETAMINOPHEN
ACETAMINOPHEN, CAFFEINE, AND DIHYDROCODEINE BITARTRATE, ACETAMINOPHEN
ACETASOL HC, ACETIC ACID, GLACIAL
ACETASOL, ACETIC ACID, GLACIAL
ACETAZOLAMIDE SODIUM, ACETAZOLAMIDE SODIUM
ACETAZOLAMIDE, ACETAZOLAMIDE
ACETIC ACID 0.25% IN PLASTIC CONTAINER, ACETIC ACID, GLACIAL
ACETIC ACID 2% IN AQUEOUS ALUMINUM ACETATE, ACETIC ACID, GLACIAL
ACETIC ACID, ACETIC ACID, GLACIAL
ACETOHEXAMIDE, ACETOHEXAMIDE
ACETYLCYSTEINE, ACETYLCYSTEINE
ACILAC, LACTULOSE
ACIPHEX, RABEPRAZOLE SODIUM
ACLOVATE, ALCLOMETASONE DIPROPIONATE
ACTHREL, CORTICORELIN OVINE TRIFLUTATE
ACTIGALL, URSODIOL
ACTIQ, FENTANYL CITRATE
ACTIVELLA, ESTRADIOL
ACTONEL WITH CALCIUM (COPACKAGED), CALCIUM CARBONATE
ACTONEL, RISEDRONATE SODIUM
ACTOPLUS MET, METFORMIN
ACTOS, PIOGLITAZONE HYDROCHLORIDE
ACULAR LS, KETOROLAC TROMETHAMINE
ACULAR PRESERVATIVE FREE, KETOROLAC TROMETHAMINE
ACULAR, KETOROLAC TROMETHAMINE
ACYCLOVIR IN SODIUM CHLORIDE 0.9% PRESERVATIVE FREE, ACYCLOVIR SODIUM
ACYCLOVIR SODIUM, ACYCLOVIR SODIUM
ACYCLOVIR, ACYCLOVIR
ACYCLOVIR, ACYCLOVIR SODIUM
ADAGEN, PEGADEMASE BOVINE
ADALAT CC, NIFEDIPINE
ADDERALL 10, AMPHETAMINE ASPARTATE
ADDERALL 12.5, AMPHETAMINE ASPARTATE
ADDERALL 15, AMPHETAMINE ASPARTATE
ADDERALL 20, AMPHETAMINE ASPARTATE
ADDERALL 30, AMPHETAMINE ASPARTATE

APPENDIX A - PRODUCT NAME INDEX

A - 2

**** A ****

ADDERALL 5, AMPHETAMINE ASPARTATE
ADDERALL 7.5, AMPHETAMINE ASPARTATE
ADDERALL XR 10, AMPHETAMINE ASPARTATE
ADDERALL XR 15, AMPHETAMINE ASPARTATE
ADDERALL XR 20, AMPHETAMINE ASPARTATE
ADDERALL XR 25, AMPHETAMINE ASPARTATE
ADDERALL XR 30, AMPHETAMINE ASPARTATE
ADDERALL XR 5, AMPHETAMINE ASPARTATE
ADENOCARD, ADENOSINE
ADENOSCAN, ADENOSINE
ADENOSINE, ADENOSINE
ADIPEX-P, PHENTERMINE HYDROCHLORIDE
ADRIAMYCIN PFS, DOXORUBICIN HYDROCHLORIDE
ADVAIR DISKUS 100/50, FLUTICASONE PROPIONATE
ADVAIR DISKUS 250/50, FLUTICASONE PROPIONATE
ADVAIR DISKUS 500/50, FLUTICASONE PROPIONATE
ADVAIR HFA, FLUTICASONE PROPIONATE
ADVICOR, LOVASTATIN
ADVIL ALLERGY SINUS, CHLORPHENIRAMINE MALEATE (OTC)
ADVIL COLD AND SINUS, IBUPROFEN (OTC)
ADVIL COLD AND SINUS, IBUPROFEN POTASSIUM (OTC)
ADVIL LIQUI-GELS, IBUPROFEN POTASSIUM (OTC)
ADVIL MIGRAINE LIQUI-GELS, IBUPROFEN POTASSIUM (OTC)
ADVIL PM, DIPHENHYDRAMINE CITRATE (OTC)
ADVIL PM, DIPHENHYDRAMINE HYDROCHLORIDE (OTC)
ADVIL, IBUPROFEN (OTC)
AEROBID, FLUNISOLIDE
AEROSPAN HFA, FLUNISOLIDE
AFEDITAB CR, NIFEDIPINE
AFRINOL, PSEUDOEPHEDRINE SULFATE (OTC)
AGENERASE, AMPRENAVIR
AGGRASTAT, TIROFIBAN HYDROCHLORIDE
AGGRENOLX, ASPIRIN
AGRYLIN, ANAGRELIDE HYDROCHLORIDE
A-HYDROCORT, HYDROCORTISONE SODIUM SUCCINATE
AKBETA, LEVOBUNOLOL HYDROCHLORIDE
AKINETON, BIPERIDEN HYDROCHLORIDE
AKNE-MYCIN, ERYTHROMYCIN
AKPENTOLATE, CYCLOPENTOLATE HYDROCHLORIDE
AKPRO, DIPIVEFRIN HYDROCHLORIDE
AKTOB, TOBRAMYCIN
ALA-CORT, HYDROCORTISONE
ALAMAST, PEMIROLAST POTASSIUM
ALA-SCALP, HYDROCORTISONE
ALAVERT, LORATADINE (OTC)
ALAWAY, KETOTIFEN FUMARATE (OTC)
ALBALON, NAPHAZOLINE HYDROCHLORIDE
ALBENZA, ALBENDAZOLE
ALBUTEROL SULFATE AND IPRATROPIUM BROMIDE, ALBUTEROL SULFATE
ALBUTEROL SULFATE, ALBUTEROL SULFATE
ALBUTEROL, ALBUTEROL
ALCAINE, PROPARACAIN HYDROCHLORIDE
ALCLOMETASONE DIPROPIONATE, ALCLOMETASONE DIPROPIONATE
ALCOHOL 10% AND DEXTROSE 5%, ALCOHOL
ALCOHOL 5% AND DEXTROSE 5%, ALCOHOL
ALCOHOL 5% IN D5-W, ALCOHOL
ALDACTAZIDE, HYDROCHLOROTHIAZIDE
ALDACTONE, SPIRONOLACTONE
ALDARA, IMIQUIMOD
ALESSE, ETHINYL ESTRADIOL
ALEVE COLD AND SINUS, NAPROXEN SODIUM (OTC)
ALEVE, NAPROXEN SODIUM (OTC)
ALFENTA, ALFENTANIL HYDROCHLORIDE

APPENDIX A - PRODUCT NAME INDEX

A - 3

**** A ****

ALFENTANIL, ALFENTANIL HYDROCHLORIDE
ALIMTA, PEMETREXED DISODIUM
ALINIA, NITAZOXANIDE
ALKERAN, MELPHALAN
ALKERAN, MELPHALAN HYDROCHLORIDE
ALLAY, ACETAMINOPHEN
ALLEGRA D 24 HOUR, FEXOFENADINE HYDROCHLORIDE
ALLEGRA, FEXOFENADINE HYDROCHLORIDE
ALLEGRA-D 12 HOUR, FEXOFENADINE HYDROCHLORIDE
ALLOPURINOL SODIUM, ALLOPURINOL SODIUM
ALLOPURINOL, ALLOPURINOL
ALOCRIL, NEDOCROMIL SODIUM
ALOMIDE, LODOXAMIDE TROMETHAMINE
ALOPRIM, ALLOPURINOL SODIUM
ALORA, ESTRADIOL
ALOXI, PALONOSETRON HYDROCHLORIDE
ALPHAGAN P, BRIMONIDINE TARTRATE
ALPRAZOLAM, ALPRAZOLAM
ALPROSTADIL, ALPROSTADIL
ALREX, LOTEPREDNOL ETABONATE
ALTACE, RAMIPRIL
ALTOPREV, LOVASTATIN
ALUPENT, METAPROTERENOL SULFATE
AMANTADINE HYDROCHLORIDE, AMANTADINE HYDROCHLORIDE
AMARYL, GLIMEPIRIDE
AMBIEN CR, ZOLPIDEM TARTRATE
AMBIEN, ZOLPIDEM TARTRATE
AMBISOME, AMPHOTERICIN B
AMCINONIDE, AMCINONIDE
AMERGE, NARATRIPTAN HYDROCHLORIDE
A-METHAPRED, METHYLPREDNISOLONE SODIUM SUCCINATE
AMICAR, AMINOCAPROIC ACID
AMIDATE, ETOMIDATE
AMIKACIN SULFATE IN SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER, AMIKACIN SULFATE
AMIKACIN SULFATE, AMIKACIN SULFATE
AMIKIN, AMIKACIN SULFATE
AMILORIDE HYDROCHLORIDE AND HYDROCHLOROTHIAZIDE, AMILORIDE HYDROCHLORIDE
AMILORIDE HYDROCHLORIDE, AMILORIDE HYDROCHLORIDE
AMINOACETIC ACID 1.5% IN PLASTIC CONTAINER, GLYCINE
AMINOCAPROIC ACID IN PLASTIC CONTAINER, AMINOCAPROIC ACID
AMINOCAPROIC ACID, AMINOCAPROIC ACID
AMINOCAPROIC, AMINOCAPROIC ACID
AMINOHIPPUATE SODIUM, AMINOHIPPUATE SODIUM
AMINOPHYLLINE DYE FREE, AMINOPHYLLINE
AMINOPHYLLINE IN SODIUM CHLORIDE 0.45%, AMINOPHYLLINE
AMINOPHYLLINE, AMINOPHYLLINE
AMINOSYN 10% (PH6), AMINO ACIDS
AMINOSYN 10%, AMINO ACIDS
AMINOSYN 3.5% M, AMINO ACIDS
AMINOSYN 3.5%, AMINO ACIDS
AMINOSYN 5%, AMINO ACIDS
AMINOSYN 7% (PH6), AMINO ACIDS
AMINOSYN 7% W/ ELECTROLYTES, AMINO ACIDS
AMINOSYN 7%, AMINO ACIDS
AMINOSYN 8.5% (PH6), AMINO ACIDS
AMINOSYN 8.5% W/ ELECTROLYTES, AMINO ACIDS
AMINOSYN 8.5%, AMINO ACIDS
AMINOSYN II 10% IN PLASTIC CONTAINER, AMINO ACIDS
AMINOSYN II 10% W/ ELECTROLYTES, AMINO ACIDS
AMINOSYN II 10%, AMINO ACIDS
AMINOSYN II 15% IN PLASTIC CONTAINER, AMINO ACIDS
AMINOSYN II 3.5% IN DEXTROSE 25% IN PLASTIC CONTAINER, AMINO ACIDS
AMINOSYN II 3.5% IN DEXTROSE 5% IN PLASTIC CONTAINER, AMINO ACIDS

APPENDIX A - PRODUCT NAME INDEX

A - 4

**** A ****

AMINOSYN II 3.5% M IN DEXTROSE 5% IN PLASTIC CONTAINER, AMINO ACIDS
AMINOSYN II 3.5% W/ ELECTROLYTES IN DEXTROSE 25% W/ CALCIUM IN PLASTIC CONTAINER, AMINO ACIDS
AMINOSYN II 4.25% IN DEXTROSE 10% IN PLASTIC CONTAINER, AMINO ACIDS
AMINOSYN II 4.25% IN DEXTROSE 20% IN PLASTIC CONTAINER, AMINO ACIDS
AMINOSYN II 4.25% IN DEXTROSE 25% IN PLASTIC CONTAINER, AMINO ACIDS
AMINOSYN II 4.25% M IN DEXTROSE 10% IN PLASTIC CONTAINER, AMINO ACIDS
AMINOSYN II 4.25% W/ ELECTROLYTES IN DEXTROSE 20% W/ CALCIUM IN PLASTIC CONTAINER, AMINO ACIDS
AMINOSYN II 4.25% W/ ELECTROLYTES IN DEXTROSE 25% W/ CALCIUM IN PLASTIC CONTAINER, AMINO ACIDS
AMINOSYN II 5% IN DEXTROSE 25% IN PLASTIC CONTAINER, AMINO ACIDS
AMINOSYN II 7%, AMINO ACIDS
AMINOSYN II 8.5% W/ ELECTROLYTES, AMINO ACIDS
AMINOSYN II 8.5%, AMINO ACIDS
AMINOSYN-HBC 7%, AMINO ACIDS
AMINOSYN-HF 8%, AMINO ACIDS
AMINOSYN-PF 10%, AMINO ACIDS
AMINOSYN-PF 7%, AMINO ACIDS
AMINOSYN-RF 5.2%, AMINO ACIDS
AMIODARONE HYDROCHLORIDE, AMIODARONE HYDROCHLORIDE
AMITIZA, LUBIPROSTONE
AMITRIPTYLINE HYDROCHLORIDE, AMITRIPTYLINE HYDROCHLORIDE
AMLEXANOX, AMLEXANOX
AMLODIPINE BESYLATE, AMLODIPINE BESYLATE
AMMONIUM CHLORIDE IN PLASTIC CONTAINER, AMMONIUM CHLORIDE
AMMONIUM LACTATE, AMMONIUM LACTATE
AMMONUL, SODIUM BENZOATE
AMNESTEEM, ISOTRETINOIN
AMOXAPINE, AMOXAPINE
AMOXICILLIN AND CLAVULANATE POTASSIUM, AMOXICILLIN
AMOXICILLIN PEDIATRIC, AMOXICILLIN
AMOXICILLIN, AMOXICILLIN
AMOXIL, AMOXICILLIN
AMPHADASE, HYALURONIDASE
AMPHOTEC, AMPHOTERICIN B
AMPHOTERICIN B, AMPHOTERICIN B
AMPICILLIN AND SULBACTAM, AMPICILLIN SODIUM
AMPICILLIN SODIUM, AMPICILLIN SODIUM
AMPICILLIN TRIHYDRATE, AMPICILLIN/AMPICILLIN TRIHYDRATE
AMRINONE LACTATE, INAMRINONE LACTATE
AMRINONE, INAMRINONE LACTATE
ANADROL-50, OXYMETHOLONE
ANAFRANIL, CLOMIPRAMINE HYDROCHLORIDE
ANAGRELIDE HYDROCHLORIDE, ANAGRELIDE HYDROCHLORIDE
ANAPROX DS, NAPROXEN SODIUM
ANAPROX, NAPROXEN SODIUM
ANCEF IN PLASTIC CONTAINER, CEFAZOLIN SODIUM
ANCEF, CEFAZOLIN SODIUM
ANCOBON, FLUCYTOSINE
ANDRODERM, TESTOSTERONE
ANDROGEL, TESTOSTERONE
ANDROID 10, METHYLTESTOSTERONE
ANDROID 25, METHYLTESTOSTERONE
AN-DTPA, TECHNETIUM TC-99M PENTETATE KIT
ANECTINE, SUCCINYLCHOLINE CHLORIDE
ANESTACON, LIDOCAINE HYDROCHLORIDE
ANEXSIA 5/325, ACETAMINOPHEN
ANEXSIA 7.5/325, ACETAMINOPHEN
ANEXSIA 7.5/650, ACETAMINOPHEN
ANEXSIA, ACETAMINOPHEN
ANGELIQ, DROSPIRENONE
ANGIOMAX, BIVALIRUDIN
ANSAID, FLURBIPROFEN
ANSPOR, CEPHRADINE
AN-SULFUR COLLOID, TECHNETIUM TC-99M SULFUR COLLOID KIT

APPENDIX A - PRODUCT NAME INDEX

A - 5

**** A ****

ANTABUSE, DISULFIRAM
ANTARA (MICRONIZED), FENOFIBRATE
ANTHELIOS 20, AVOBENZONE (OTC)
ANTHELIOS SX, AVOBENZONE (OTC)
ANTIVERT, MECLIZINE HYDROCHLORIDE
ANTIZOL, FOMEPIZOLE
ANTURANE, SULFINPYRAZONE
ANUSOL HC, HYDROCORTISONE
ANZEMET, DOLASETRON MESYLATE
APHTHASOL, AMLEXANOX
APIDRA, INSULIN GLULISINE RECOMBINANT
APOKYN, APOMORPHINE HYDROCHLORIDE
APRESAZIDE, HYDRALAZINE HYDROCHLORIDE
APTIVUS, TIPRANAVIR
AQUASOL A, VITAMIN A PALMITATE
ARALEN, CHLOROQUINE PHOSPHATE
ARANELLE, ETHINYL ESTRADIOL
ARAVA, LEFLUNOMIDE
AREDIA, PAMIDRONATE DISODIUM
ARESTIN, MINOCYCLINE HYDROCHLORIDE
ARGATROBAN, ARGATROBAN
ARICEPT ODT, DONEPEZIL HYDROCHLORIDE
ARICEPT, DONEPEZIL HYDROCHLORIDE
ARIMIDEX, ANASTROZOLE
ARISTOSPAN, TRIAMCINOLONE HEXACETONIDE
ARIXTRA, FONDAPARINUX SODIUM
AROMASIN, EXEMESTANE
ARRANON, NELARABINE
ARTHROTEC, DICLOFENAC SODIUM
ASACOL, MESALAMINE
ASMANEX TWISTHALER, MOMETASONE FUROATE
ASPIRIN AND CAFFEINE W/ BUTALBITAL, ASPIRIN
ASPIRIN AND OXYCODONE, ASPIRIN
ASTELIN, AZELASTINE HYDROCHLORIDE
ASTRAMORPH PF, MORPHINE SULFATE
ATACAND HCT, CANDESARTAN CILEXETIL
ATACAND, CANDESARTAN CILEXETIL
ATENOLOL AND CHLORTHALIDONE, ATENOLOL
ATENOLOL, ATENOLOL
ATIVAN, LORAZEPAM
ATRACURIUM BESYLATE PRESERVATIVE FREE, ATRACURIUM BESYLATE
ATRACURIUM BESYLATE, ATRACURIUM BESYLATE
ATRIDOX, DOXYCYCLINE HYCLATE
ATRIPLA, EFAVIRENZ
ATROPEN, ATROPINE
ATROPINE SULFATE ANSYR PLASTIC SYRINGE, ATROPINE SULFATE
ATROVENT HFA, IPRATROPIUM BROMIDE
ATROVENT, IPRATROPIUM BROMIDE
AUGMENTIN '125', AMOXICILLIN
AUGMENTIN '200', AMOXICILLIN
AUGMENTIN '250', AMOXICILLIN
AUGMENTIN '400', AMOXICILLIN
AUGMENTIN '500', AMOXICILLIN
AUGMENTIN '875', AMOXICILLIN
AUGMENTIN ES-600, AMOXICILLIN
AUGMENTIN XR, AMOXICILLIN
AVAGARD, ALCOHOL (OTC)
AVAGE, TAZAROTENE
AVALIDE, HYDROCHLOROTHIAZIDE
AVANDAMET, METFORMIN HYDROCHLORIDE
AVANDARYL, GLIMEPIRIDE
AVANDIA, ROSIGLITAZONE MALEATE
AVAPRO, IRBESARTAN

APPENDIX A - PRODUCT NAME INDEX

A - 6

**** A ****

AVC, SULFANILAMIDE
AVELOX IN SODIUM CHLORIDE 0.8% IN PLASTIC CONTAINER, MOXIFLOXACIN HYDROCHLORIDE
AVELOX, MOXIFLOXACIN HYDROCHLORIDE
AVENTYL HYDROCHLORIDE, NORTRIPTYLINE HYDROCHLORIDE
AVIANE-28, ETHINYL ESTRADIOL
AVINZA, MORPHINE SULFATE
AVITA, TRETINOIN
AVODART, DUTASTERIDE
AXERT, ALMOTRIPTAN MALATE
AXID AR, NIZATIDINE (OTC)
AXID, NIZATIDINE
AYGESTIN, NORETHINDRONE ACETATE
AZACTAM IN PLASTIC CONTAINER, AZTREONAM
AZACTAM, AZTREONAM
AZASAN, AZATHIOPRINE
AZATHIOPRINE SODIUM, AZATHIOPRINE SODIUM
AZATHIOPRINE, AZATHIOPRINE
AZDONE, ASPIRIN
AZELEX, AZELAIC ACID
AZILECT, RASAGILINE MESYLATE
AZITHROMYCIN, AZITHROMYCIN
AZITHROMYCIN, AZITHROMYCIN HYDROGENCITRATE
AZMACORT, TRIAMCINOLONE ACETONIDE
AZOPT, BRINZOLAMIDE
AZULFIDINE EN-TABS, SULFASALAZINE
AZULFIDINE, SULFASALAZINE

**** B ****

BACIIM, BACITRACIN
BACI-RX, BACITRACIN
BACITRACIN ZINC AND POLYMYXIN B SULFATE, BACITRACIN ZINC
BACITRACIN, BACITRACIN
BACITRACIN-NEOMYCIN-POLYMYXIN W/ HYDROCORTISONE ACETATE, BACITRACIN
BACITRACIN-NEOMYCIN-POLYMYXIN, BACITRACIN ZINC
BACLOFEN, BACLOFEN
BACTERIOSTATIC SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER, SODIUM CHLORIDE
BACTERIOSTATIC WATER FOR INJECTION IN PLASTIC CONTAINER, WATER FOR INJECTION, STERILE
BACTOCILL IN PLASTIC CONTAINER, OXACILLIN SODIUM
BACTOCILL, OXACILLIN SODIUM
BACTRIM DS, SULFAMETHOXAZOLE
BACTRIM PEDIATRIC, SULFAMETHOXAZOLE
BACTRIM, SULFAMETHOXAZOLE
BACTROBAN, MUPIROCIN
BACTROBAN, MUPIROCIN CALCIUM
BAL, DIMERCAPROL
BALANCED SALT SOLUTION, CALCIUM CHLORIDE
BALZIVA-21, ETHINYL ESTRADIOL
BALZIVA-28, ETHINYL ESTRADIOL
BARACLUDE, ENTECAVIR
BAROS, SODIUM BICARBONATE
BAYER EXTRA STRENGTH ASPIRIN FOR MIGRAINE PAIN, ASPIRIN (OTC)
BECONASE AQ, BECLOMETHASONE DIPROPIONATE MONOHYDRATE
BENADRYL PRESERVATIVE FREE, DIPHENHYDRAMINE HYDROCHLORIDE
BENADRYL, DIPHENHYDRAMINE HYDROCHLORIDE
BENZAEPRIIL HYDROCHLORIDE AND HYDROCHLOROTHIAZIDE, BENAZEPRIL HYDROCHLORIDE
BENZAEPRIIL HYDROCHLORIDE, BENAZEPRIL HYDROCHLORIDE
BENICAR HCT, HYDROCHLOROTHIAZIDE
BENICAR, OLMESARTAN MEDOXOMIL
BENOQUIN, MONOBENZONE
BENTYL PRESERVATIVE FREE, DICYCLOMINE HYDROCHLORIDE
BENTYL, DICYCLOMINE HYDROCHLORIDE
BENZACLIN, BENZOYL PEROXIDE
BENZAMYCIN PAK, BENZOYL PEROXIDE

APPENDIX A - PRODUCT NAME INDEX

A - 7

**** B ****

BENZAMYCIN, BENZOYL PEROXIDE
BENZONATATE, BENZONATATE
BENZPHETAMINE HYDROCHLORIDE, BENZPHETAMINE HYDROCHLORIDE
BENZTROPINE MESYLATE, BENZTROPINE MESYLATE
BETA-2, ISOETHARINE HYDROCHLORIDE
BETADINE, POVIDONE-IODINE
BETAGAN, LEVOBUNOLOL HYDROCHLORIDE
BETAMETHASONE DIPROPIONATE, BETAMETHASONE DIPROPIONATE
BETAMETHASONE VALERATE, BETAMETHASONE VALERATE
BETAPACE AF, SOTALOL HYDROCHLORIDE
BETAPACE, SOTALOL HYDROCHLORIDE
BETA-VAL, BETAMETHASONE VALERATE
BETAXOLOL HYDROCHLORIDE, BETAXOLOL HYDROCHLORIDE
BETAXOLOL, BETAXOLOL HYDROCHLORIDE
BETHANECHOL CHLORIDE, BETHANECHOL CHLORIDE
BETIMOL, TIMOLOL
BETOPTIC S, BETAXOLOL HYDROCHLORIDE
BETOPTIC, BETAXOLOL HYDROCHLORIDE
BEXTRA, VALDECOXIB
BIAXIN XL, CLARITHROMYCIN
BIAXIN, CLARITHROMYCIN
BICILLIN C-R 900/300, PENICILLIN G BENZATHINE
BICILLIN C-R, PENICILLIN G BENZATHINE
BICILLIN L-A, PENICILLIN G BENZATHINE
BICNU, CARMUSTINE
BIDIL, HYDRAZINE HYDROCHLORIDE
BILTRICIDE, PRAZIQUANTEL
BIOSCRUB, CHLORHEXIDINE GLUCONATE (OTC)
BISOPROLOL FUMARATE AND HYDROCHLOROTHIAZIDE, BISOPROLOL FUMARATE
BISOPROLOL FUMARATE, BISOPROLOL FUMARATE
BLENOXANE, BLEOMYCIN SULFATE
BLEOMYCIN, BLEOMYCIN SULFATE
BLEPH-10, SULFACETAMIDE SODIUM
BLEPH-30, SULFACETAMIDE SODIUM
BLEPHAMIDE S.O.P., PREDNISOLONE ACETATE
BLEPHAMIDE, PREDNISOLONE ACETATE
BONIVA, IBANDRONATE SODIUM
BONTRIL PDM, PHENDIMETRAZINE TARTRATE
BONTRIL, PHENDIMETRAZINE TARTRATE
BRANCHAMIN 4% IN PLASTIC CONTAINER, AMINO ACIDS
BRAVELLE, UROFOLLITROPIN
BREATHTEK UBT FOR H-PYLORI, UREA, C-13
BRETHINE, TERBUTALINE SULFATE
BRETILIUM TOSYLATE IN DEXTROSE 5% IN PLASTIC CONTAINER, BRETILIUM TOSYLATE
BRETILIUM TOSYLATE IN PLASTIC CONTAINER, BRETILIUM TOSYLATE
BRETILIUM TOSYLATE, BRETILIUM TOSYLATE
BREVIBLOC DOUBLE STRENGTH IN PLASTIC CONTAINER, ESMOLOL HYDROCHLORIDE
BREVIBLOC IN PLASTIC CONTAINER, ESMOLOL HYDROCHLORIDE
BREVIBLOC, ESMOLOL HYDROCHLORIDE
BREVICON 21-DAY, ETHINYL ESTRADIOL
BREVICON 28-DAY, ETHINYL ESTRADIOL
BREVITAL SODIUM, METHOHEXITAL SODIUM
BRIAN CARE, CHLORHEXIDINE GLUCONATE (OTC)
BRIMONIDINE TARTRATE, BRIMONIDINE TARTRATE
BRISTACYCLINE, TETRACYCLINE HYDROCHLORIDE
BROMFED-DM, BROMPHENIRAMINE MALEATE
BROMOCRIPTINE MESYLATE, BROMOCRIPTINE MESYLATE
BROMPHENIRAMINE MALEATE, BROMPHENIRAMINE MALEATE
BRONCHO SALINE, SODIUM CHLORIDE (OTC)
BRONITIN MIST, EPINEPHRINE BITARTRATE (OTC)
BROVANA, ARFORMOTEROL TARTRATE
BSS PLUS, CALCIUM CHLORIDE
BSS, CALCIUM CHLORIDE

APPENDIX A - PRODUCT NAME INDEX

A - 8

**** B ****

BUCET, ACETAMINOPHEN
BUDESONIDE, BUDESONIDE
BUMETANIDE, BUMETANIDE
BUMEX, BUMETANIDE
BUPHENYL, SODIUM PHENYLBUTYRATE
BUPIVACAINE HYDROCHLORIDE AND EPINEPHRINE, BUPIVACAINE HYDROCHLORIDE
BUPIVACAINE HYDROCHLORIDE PRESERVATIVE FREE, BUPIVACAINE HYDROCHLORIDE
BUPIVACAINE HYDROCHLORIDE W/EPINEPHRINE, BUPIVACAINE HYDROCHLORIDE
BUPIVACAINE HYDROCHLORIDE, BUPIVACAINE HYDROCHLORIDE
BUPIVACAINE, BUPIVACAINE HYDROCHLORIDE
BUPRENEX, BUPRENORPHINE HYDROCHLORIDE
BUPRENORPHINE HYDROCHLORIDE, BUPRENORPHINE HYDROCHLORIDE
BUPROPION HYDROCHLORIDE, BUPROPION HYDROCHLORIDE
BUSPAR, BUSPIRONE HYDROCHLORIDE
BUSPIRONE HYDROCHLORIDE, BUSPIRONE HYDROCHLORIDE
BUSULFEX, BUSULFAN
BUTABARBITAL, BUTABARBITAL SODIUM
BUTAL COMPOUND, ASPIRIN
BUTALBITAL, ACETAMINOPHEN AND CAFFEINE, ACETAMINOPHEN
BUTALBITAL, ACETAMINOPHEN, AND CAFFEINE WITH CODEINE PHOSPHATE, ACETAMINOPHEN
BUTALBITAL, ACETAMINOPHEN, CAFFEINE AND CODEINE PHOSPHATE, ACETAMINOPHEN
BUTALBITAL, APAP, AND CAFFEINE, ACETAMINOPHEN
BUTALBITAL, ASPIRIN AND CAFFEINE, ASPIRIN
BUTALBITAL, ASPIRIN, CAFFEINE, AND CODEINE PHOSPHATE, ASPIRIN
BUTAPAP, ACETAMINOPHEN
BUTISOL SODIUM, BUTABARBITAL SODIUM
BUTORPHANOL TARTRATE PRESERVATIVE FREE, BUTORPHANOL TARTRATE
BUTORPHANOL TARTRATE, BUTORPHANOL TARTRATE
BYETTA, EXENATIDE SYNTHETIC

**** C ****

CABERGOLINE, CABERGOLINE
CADUET, AMLODIPINE BESYLATE
CAFCIT, CAFFEINE CITRATE
CAFERGOT, CAFFEINE
CAFFEINE CITRATE, CAFFEINE CITRATE
CALAN, VERAPAMIL HYDROCHLORIDE
CALCIJEX, CALCITRIOL
CALCITRIOL, CALCITRIOL
CALCIUM CHLORIDE 10% IN PLASTIC CONTAINER, CALCIUM CHLORIDE
CALCIUM DISODIUM VERSENATE, EDETATE CALCIUM DISODIUM
CAMILA, NORETHINDRONE
CAMPRAL, ACAMPROSATE CALCIUM
CAMPTOSAR, IRINOTECAN HYDROCHLORIDE
CANASA, MESALAMINE
CANCIDAS, CASPOFUNGIN ACETATE
CANTIL, MEPENZOLATE BROMIDE
CAPASTAT SULFATE, CAPREOMYCIN SULFATE
CAPITAL AND CODEINE, ACETAMINOPHEN
CAPITAL SOLEIL 15, AVOBENZONE (OTC)
CAPITROL, CHLOROXINE
CAPOTEN, CAPTOPRIL
CAPOZIDE 25/15, CAPTOPRIL
CAPOZIDE 25/25, CAPTOPRIL
CAPOZIDE 50/15, CAPTOPRIL
CAPOZIDE 50/25, CAPTOPRIL
CAP-PROFEN, IBUPROFEN (OTC)
CAPTOPRIL AND HYDROCHLOROTHIAZIDE, CAPTOPRIL
CAPTOPRIL, CAPTOPRIL
CARAC, FLUOROURACIL
CARAFATE, SUCRALFATE
CARBAMAZEPINE, CARBAMAZEPINE
CARBASTAT, CARBACHOL

APPENDIX A - PRODUCT NAME INDEX

A - 9

**** C ****

CARBATROL, CARBAMAZEPINE
CARBIDOPA AND LEVODOPA, CARBIDOPA
CARBILEV, CARBIDOPA
CARBINOXAMINE MALEATE, CARBINOXAMINE MALEATE
CARBOCAINE W/ NEO-COBEFRIN, LEVONORDEFIN
CARBOCAINE, MEPIVACAINE HYDROCHLORIDE
CARBOPLATIN, CARBOPLATIN
CARDENE SR, NICARDIPINE HYDROCHLORIDE
CARDENE, NICARDIPINE HYDROCHLORIDE
CARDIOGEN-82, RUBIDIUM CHLORIDE RB-82
CARDIOLITE, TECHNETIUM TC-99M SESTAMIBI KIT
CARDIOPLEGIC IN PLASTIC CONTAINER, CALCIUM CHLORIDE
CARDIZEM CD, DILTIAZEM HYDROCHLORIDE
CARDIZEM LA, DILTIAZEM HYDROCHLORIDE
CARDIZEM, DILTIAZEM HYDROCHLORIDE
CARDURA XL, DOXAZOSIN MESYLATE
CARDURA, DOXAZOSIN MESYLATE
CARISOPRODOL AND ASPIRIN, ASPIRIN
CARISOPRODOL, ASPIRIN AND CODEINE PHOSPHATE, ASPIRIN
CARISOPRODOL, CARISOPRODOL
CARMOL HC, HYDROCORTISONE ACETATE
CARNITOR, LEVOCARNITINE
CARTEOLOL HYDROCHLORIDE, CARTEOLOL HYDROCHLORIDE
CARTIA XT, DILTIAZEM HYDROCHLORIDE
CASODEX, BICALUTAMIDE
CATAFLAM, DICLOFENAC POTASSIUM
CATAPRES, CLONIDINE HYDROCHLORIDE
CATAPRES-TTS-1, CLONIDINE
CATAPRES-TTS-2, CLONIDINE
CATAPRES-TTS-3, CLONIDINE
CAVERJECT, ALPROSTADIL
CEDAX, CEFTIBUTEN DIHYDRATE
CEENU, LOMUSTINE
CEFACLOR, CEFACLOR
CEFADROXIL, CEFADROXIL/CEFADROXIL HEMIHYDRATE
CEFAZOLIN AND DEXTROSE, CEFAZOLIN SODIUM
CEFAZOLIN SODIUM, CEFAZOLIN SODIUM
CEFAZOLIN, CEFAZOLIN SODIUM
CEFDINIR, CEFDINIR
CEFIZOX IN PLASTIC CONTAINER, CEFTIZOXIME SODIUM
CEFIZOX, CEFTIZOXIME SODIUM
CEFOBID IN PLASTIC CONTAINER, CEFOPERAZONE SODIUM
CEFOBID, CEFOPERAZONE SODIUM
CEFOTAXIME SODIUM, CEFOTAXIME SODIUM
CEFOTAXIME, CEFOTAXIME SODIUM
CEFOXITIN AND DEXTROSE IN DUPLEX CONTAINER, CEFOXITIN SODIUM
CEFOXITIN, CEFOXITIN SODIUM
CEFPODOXIME PROXETIL, CEFPODOXIME PROXETIL
CEFPROZIL, CEFPROZIL
CEFTAZIDIME, CEFTAZIDIME
CEFTIN, CEFUROXIME AXETIL
CEFTRIAZONE AND DEXTROSE IN DUPLEX CONTAINER, CEFTRIAZONE SODIUM
CEFTRIAZONE IN PLASTIC CONTAINER, CEFTRIAZONE SODIUM
CEFTRIAZONE SODIUM, CEFTRIAZONE SODIUM
CEFTRIAZONE, CEFTRIAZONE SODIUM
CEFUROXIME AND DEXTROSE IN DUPLEX CONTAINER, CEFUROXIME SODIUM
CEFUROXIME AXETIL, CEFUROXIME AXETIL
CEFUROXIME SODIUM, CEFUROXIME SODIUM
CEFUROXIME, CEFUROXIME SODIUM
CEFZIL, CEFPROZIL
CELEBREX, CELECOXIB
CELESTONE SOLUSPAN, BETAMETHASONE ACETATE
CELESTONE, BETAMETHASONE

APPENDIX A - PRODUCT NAME INDEX

A - 10

**** C ****

CELEXA, CITALOPRAM HYDROBROMIDE
CELLCEPT, MYCOPHENOLATE MOFETIL
CELLCEPT, MYCOPHENOLATE MOFETIL HYDROCHLORIDE
CELONTIN, METHSUXIMIDE
CENESTIN, ESTROGENS, CONJUGATED SYNTHETIC A
CENTANY, MUPIROCIN
CEPHALEXIN, CEPHALEXIN
CEPHRADINE, CEPHRADINE
CEREBYX, FOSPHENYTOIN SODIUM
CEREDASE, ALGLUCERASE
CERETEC, TECHNETIUM TC-99M EXAMETAZIME KIT
CEREZYME, IMIGLUCERASE
CERUBIDINE, DAUNORUBICIN HYDROCHLORIDE
CERVIDIL, DINOPROSTONE
CESAMET, NABILONE
CETACORT, HYDROCORTISONE
CETAMIDE, SULFACETAMIDE SODIUM
CETROTIDE, CETRORELIX
CHANTIX, VARENICLINE TARTRATE
CHEMET, SUCCIMER
CHG SCRUB, CHLORHEXIDINE GLUCONATE (OTC)
CHILDREN'S ADVIL ALLERGY SINUS, CHLORPHENIRAMINE MALEATE (OTC)
CHILDREN'S ADVIL COLD, IBUPROFEN (OTC)
CHILDREN'S ADVIL, IBUPROFEN
CHILDREN'S ADVIL, IBUPROFEN (OTC)
CHILDREN'S ADVIL-FLAVORED, IBUPROFEN (OTC)
CHILDREN'S CLARITIN, LORATADINE (OTC)
CHILDREN'S ELIXSURE, IBUPROFEN (OTC)
CHILDREN'S IBUPROFEN, IBUPROFEN (OTC)
CHILDREN'S MOTRIN COLD, IBUPROFEN (OTC)
CHILDREN'S MOTRIN, IBUPROFEN (OTC)
CHIRHOSTIM, SECRETIN SYNTHETIC HUMAN
CHLORAMPHENICOL SODIUM SUCCINATE, CHLORAMPHENICOL SODIUM SUCCINATE
CHLORAMPHENICOL, CHLORAMPHENICOL
CHLORAPREP ONE-STEP FREPP, CHLORHEXIDINE GLUCONATE (OTC)
CHLORAPREP ONE-STEP SEPP, CHLORHEXIDINE GLUCONATE (OTC)
CHLORAPREP ONE-STEP, CHLORHEXIDINE GLUCONATE (OTC)
CHLORAPREP SINGLE SWABSTICK, CHLORHEXIDINE GLUCONATE (OTC)
CHLORAPREP WITH TINT, CHLORHEXIDINE GLUCONATE (OTC)
CHLORASCRUB MAXI SWABSTICK, CHLORHEXIDINE GLUCONATE (OTC)
CHLORASCRUB SWAB, CHLORHEXIDINE GLUCONATE (OTC)
CHLORASCRUB SWABSTICK, CHLORHEXIDINE GLUCONATE (OTC)
CHLORDIAZEPOXIDE AND AMITRIPTYLINE HYDROCHLORIDE, AMITRIPTYLINE HYDROCHLORIDE
CHLORDIAZEPOXIDE HYDROCHLORIDE, CHLORDIAZEPOXIDE HYDROCHLORIDE
CHLORHEXIDINE GLUCONATE, CHLORHEXIDINE GLUCONATE
CHLORHEXIDINE GLUCONATE, CHLORHEXIDINE GLUCONATE (OTC)
CHLOROMYCETIN, CHLORAMPHENICOL SODIUM SUCCINATE
CHLOROPROCAINE HYDROCHLORIDE, CHLOROPROCAINE HYDROCHLORIDE
CHLOROQUINE PHOSPHATE, CHLOROQUINE PHOSPHATE
CHLOROTHIAZIDE, CHLOROTHIAZIDE
CHLORPROMAZINE HYDROCHLORIDE INTENSOL, CHLORPROMAZINE HYDROCHLORIDE
CHLORPROMAZINE HYDROCHLORIDE, CHLORPROMAZINE HYDROCHLORIDE
CHLORPROPAMIDE, CHLORPROPAMIDE
CHLORTHALIDONE, CHLORTHALIDONE
CHLOR-TRIMETON, CHLORPHENIRAMINE MALEATE (OTC)
CHLORZOXAZONE, CHLORZOXAZONE
CHOLAC, LACTULOSE
CHOLEDYL SA, OXTRIPHYLLINE
CHOLESTYRAMINE LIGHT, CHOLESTYRAMINE
CHOLESTYRAMINE, CHOLESTYRAMINE
CHOLETEC, TECHNETIUM TC-99M MEBROFENIN KIT
CHOLOGRAFIN MEGLUMINE, IODIPAMIDE MEGLUMINE
CHORIONIC GONADOTROPIN, GONADOTROPIN, CHORIONIC

APPENDIX A - PRODUCT NAME INDEX

A - 11

**** C ****

CHROMIC CHLORIDE IN PLASTIC CONTAINER, CHROMIC CHLORIDE
CHROMITOPES SODIUM, SODIUM CHROMATE, CR-51
CIALIS, TADALAFIL
CICLOPIROX, CICLOPIROX
CIDA-STAT, CHLORHEXIDINE GLUCONATE (OTC)
CILOSTAZOL, CILOSTAZOL
CILOXAN, CIPROFLOXACIN HYDROCHLORIDE
CIMETIDINE HYDROCHLORIDE IN SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER, CIMETIDINE HYDROCHLORIDE
CIMETIDINE HYDROCHLORIDE, CIMETIDINE HYDROCHLORIDE
CIMETIDINE, CIMETIDINE
CIMETIDINE, CIMETIDINE (OTC)
CIPRO HC, CIPROFLOXACIN HYDROCHLORIDE
CIPRO IN DEXTROSE 5% IN PLASTIC CONTAINER, CIPROFLOXACIN
CIPRO XR, CIPROFLOXACIN
CIPRO, CIPROFLOXACIN
CIPRO, CIPROFLOXACIN HYDROCHLORIDE
CIPRODEX, CIPROFLOXACIN
CIPROFLOXACIN, CIPROFLOXACIN
CIPROFLOXACIN, CIPROFLOXACIN HYDROCHLORIDE
CIS-MDP, TECHNETIUM TC-99M MEDRONATE KIT
CISPLATIN, CISPLATIN
CIS-PYRO, TECHNETIUM TC-99M PYROPHOSPHATE KIT
CITALOPRAM HYDROBROMIDE, CITALOPRAM HYDROBROMIDE
CITANEST FORTE, EPINEPHRINE BITARTRATE
CITANEST PLAIN, PRILOCAINE HYDROCHLORIDE
CLADRIBINE, CLADRIBINE
CLAFORAN IN DEXTROSE 5% IN PLASTIC CONTAINER, CEFOTAXIME SODIUM
CLAFORAN, CEFOTAXIME SODIUM
CLARAVIS, ISOTRETINOIN
CLARINEX D 24 HOUR, DESLORATADINE
CLARINEX, DESLORATADINE
CLARINEX-D 12 HOUR, DESLORATADINE
CLARITHROMYCIN EXTENDED RELEASE, CLARITHROMYCIN
CLARITHROMYCIN, CLARITHROMYCIN
CLARITIN HIVES RELIEF REDITAB, LORATADINE (OTC)
CLARITIN HIVES RELIEF, LORATADINE (OTC)
CLARITIN REDITABS, LORATADINE (OTC)
CLARITIN, LORATADINE (OTC)
CLARITIN-D 24 HOUR, LORATADINE (OTC)
CLEMASTINE FUMARATE, CLEMASTINE FUMARATE
CLEMASTINE FUMARATE, CLEMASTINE FUMARATE (OTC)
CLEOCIN HYDROCHLORIDE, CLINDAMYCIN HYDROCHLORIDE
CLEOCIN PHOSPHATE IN DEXTROSE 5% IN PLASTIC CONTAINER, CLINDAMYCIN PHOSPHATE
CLEOCIN PHOSPHATE, CLINDAMYCIN PHOSPHATE
CLEOCIN T, CLINDAMYCIN PHOSPHATE
CLEOCIN, CLINDAMYCIN PALMITATE HYDROCHLORIDE
CLEOCIN, CLINDAMYCIN PHOSPHATE
CLIMARA PRO, ESTRADIOL
CLIMARA, ESTRADIOL
CLINDA-DERM, CLINDAMYCIN PHOSPHATE
CLINDAGEL, CLINDAMYCIN PHOSPHATE
CLINDAMYCIN HYDROCHLORIDE, CLINDAMYCIN HYDROCHLORIDE
CLINDAMYCIN PHOSPHATE IN DEXTROSE 5%, CLINDAMYCIN PHOSPHATE
CLINDAMYCIN PHOSPHATE, CLINDAMYCIN PHOSPHATE
CLINDESSE, CLINDAMYCIN PHOSPHATE
CLINDETS, CLINDAMYCIN PHOSPHATE
CLINIMIX 2.75/10 SULFITE FREE IN DEXTROSE 10% IN PLASTIC CONTAINER, AMINO ACIDS
CLINIMIX 2.75/25 SULFITE FREE IN DEXTROSE 25% IN PLASTIC CONTAINER, AMINO ACIDS
CLINIMIX 2.75/5 SULFITE FREE IN DEXTROSE 5% IN PLASTIC CONTAINER, AMINO ACIDS
CLINIMIX 4.25/10 SULFITE FREE IN DEXTROSE 10% IN PLASTIC CONTAINER, AMINO ACIDS
CLINIMIX 4.25/20 SULFITE FREE IN DEXTROSE 20% IN PLASTIC CONTAINER, AMINO ACIDS
CLINIMIX 4.25/25 SULFITE FREE IN DEXTROSE 25% IN PLASTIC CONTAINER, AMINO ACIDS
CLINIMIX 4.25/5 SULFITE FREE IN DEXTROSE 5% IN PLASTIC CONTAINER, AMINO ACIDS

APPENDIX A - PRODUCT NAME INDEX

A - 12

**** C ****

CLINIMIX 5/10 SULFITE FREE IN DEXTROSE 10% IN PLASTIC CONTAINER, AMINO ACIDS
CLINIMIX 5/15 SULFITE FREE IN DEXTROSE 15% IN PLASTIC CONTAINER, AMINO ACIDS
CLINIMIX 5/20 SULFITE FREE IN DEXTROSE 20% IN PLASTIC CONTAINER, AMINO ACIDS
CLINIMIX 5/25 SULFITE FREE IN DEXTROSE 25% IN PLASTIC CONTAINER, AMINO ACIDS
CLINIMIX 5/35 SULFITE FREE IN DEXTROSE 35% IN PLASTIC CONTAINER, AMINO ACIDS
CLINIMIX E 2.75/10 SULFITE-FREE W/ ELECT IN DEXTROSE 10% W/ CALCIUM IN PLASTIC CONTAINER, AMINO ACIDS
CLINIMIX E 2.75/25 SULFITE-FREE W/ ELECT IN DEXTROSE 25% W/ CALCIUM IN PLASTIC CONTAINER, AMINO ACIDS
CLINIMIX E 2.75/5 SULFITE-FREE W/ ELECT IN DEXTROSE 5% W/ CALCIUM IN PLASTIC CONTAINER, AMINO ACIDS
CLINIMIX E 4.25/10 SULFITE-FREE W/ ELECT IN DEXTROSE 10% W/ CALCIUM IN PLASTIC CONTAINER, AMINO ACIDS
CLINIMIX E 4.25/20 SULFITE-FREE W/ ELECT IN DEXTROSE 20% W/ CALCIUM IN PLASTIC CONTAINER, AMINO ACIDS
CLINIMIX E 4.25/25 SULFITE-FREE W/ ELECT IN DEXTROSE 25% W/ CALCIUM IN PLASTIC CONTAINER, AMINO ACIDS
CLINIMIX E 4.25/5 SULFITE-FREE W/ ELECT IN DEXTROSE 5% W/ CALCIUM IN PLASTIC CONTAINER, AMINO ACIDS
CLINIMIX E 5/10 SULFITE-FREE W/ ELECT IN DEXTROSE 10% W/ CALCIUM IN PLASTIC CONTAINER, AMINO ACIDS
CLINIMIX E 5/15 SULFITE-FREE W/ ELECT IN DEXTROSE 15% W/ CALCIUM IN PLASTIC CONTAINER, AMINO ACIDS
CLINIMIX E 5/20 SULFITE-FREE W/ ELECT IN 20% DEXTROSE W/ CALCIUM IN PLASTIC CONTAINER, AMINO ACIDS
CLINIMIX E 5/25 SULFITE-FREE W/ ELECT IN DEXTROSE 25% W/ CALCIUM IN PLASTIC CONTAINER, AMINO ACIDS
CLINIMIX E 5/35 SULFITE-FREE W/ ELECT IN DEXTROSE 35% W/ CALCIUM IN PLASTIC CONTAINER, AMINO ACIDS
CLINISOL 15% SULFITE FREE IN PLASTIC CONTAINER, AMINO ACIDS
CLINORIL, SULINDAC
CLOBETASOL PROPIONATE (EMOLLIENT), CLOBETASOL PROPIONATE
CLOBETASOL PROPIONATE, CLOBETASOL PROPIONATE
CLOBEX, CLOBETASOL PROPIONATE
CLODERM, CLOCORTOLONE PIVALATE
CLOFIBRATE, CLOFIBRATE
CLOLAR, CLOFARABINE
CLOMID, CLOMIPHENE CITRATE
CLOMIPHENE CITRATE, CLOMIPHENE CITRATE
CLOMIPRAMINE HYDROCHLORIDE, CLOMIPRAMINE HYDROCHLORIDE
CLONAZEPAM, CLONAZEPAM
CLONIDINE HYDROCHLORIDE, CLONIDINE HYDROCHLORIDE
CLOPIDOGREL BISULFATE, CLOPIDOGREL BISULFATE
CLORAZEPATE DIPOTASSIUM, CLORAZEPATE DIPOTASSIUM
CLORPRES, CHLORTHALIDONE
CLOTRIMAZOLE AND BETAMETHASONE DIPROPIONATE, BETAMETHASONE DIPROPIONATE
CLOTRIMAZOLE, CLOTRIMAZOLE
CLOTRIMAZOLE, CLOTRIMAZOLE (OTC)
CLOXACILLIN SODIUM, CLOXACILLIN SODIUM
CLOXAPEN, CLOXACILLIN SODIUM
CLOZAPINE, CLOZAPINE
CLOZARIL, CLOZAPINE
CODEPREX, CHLORPHENIRAMINE POLISTIREX
CODRIX, ACETAMINOPHEN
COGENTIN, BENZTROPINE MESYLATE
CO-GESIC, ACETAMINOPHEN
COGNEX, TACRINE HYDROCHLORIDE
CO-LAV, POLYETHYLENE GLYCOL 3350
COLAZAL, BALSALAZIDE DISODIUM
COLESTID, COLESTIPOL HYDROCHLORIDE
COLESTIPOL HYDROCHLORIDE, COLESTIPOL HYDROCHLORIDE
COLGATE TOTAL, SODIUM FLUORIDE (OTC)
COLISTIMETHATE, COLISTIMETHATE SODIUM
COLOCORT, HYDROCORTISONE
COL-PROBENECID, COLCHICINE

APPENDIX A - PRODUCT NAME INDEX

A - 13

**** C ****

COLY-MYCIN M, COLISTIMETHATE SODIUM
COLY-MYCIN S, COLISTIN SULFATE
COLYTE WITH FLAVOR PACKS, POLYETHYLENE GLYCOL 3350
COLYTE, POLYETHYLENE GLYCOL 3350
COLYTE-FLAVORED, POLYETHYLENE GLYCOL 3350
COMBIPATCH, ESTRADIOL
COMBIVENT, ALBUTEROL SULFATE
COMBIVIR, LAMIVUDINE
COMBUNOX, IBUPROFEN
COMMIT, NICOTINE POLACRILEX (OTC)
COMPAZINE, PROCHLORPERAZINE
COMPAZINE, PROCHLORPERAZINE EDISYLATE
COMPAZINE, PROCHLORPERAZINE MALEATE
COMPRO, PROCHLORPERAZINE
COMTAN, ENTACAPONE
CONCERTA, METHYLPHENIDATE HYDROCHLORIDE
CONDYLOX, PODOFILOX
CONRAY 30, IOTHALAMATE MEGLUMINE
CONRAY 400, IOTHALAMATE SODIUM
CONRAY 43, IOTHALAMATE MEGLUMINE
CONRAY, IOTHALAMATE MEGLUMINE
CONSTILAC, LACTULOSE
CONSTULOSE, LACTULOSE
COPAXONE, GLATIRAMER ACETATE
COPEGUS, RIBAVIRIN
CORDARONE, AMIODARONE HYDROCHLORIDE
CORDRAN SP, FLURANDRENOLIDE
CORDRAN, FLURANDRENOLIDE
COREG CR, CARVEDILOL PHOSPHATE
COREG, CARVEDILOL
CORGARD, NADOLOL
CORLOPAM, FENOLDOPAM MESYLATE
CORMAX, CLOBETASOL PROPIONATE
CORPHED, PSEUDOEPHEDRINE HYDROCHLORIDE
CORTEF, HYDROCORTISONE
CORTENEMA, HYDROCORTISONE
CORTIFOAM, HYDROCORTISONE ACETATE
CORTISONE ACETATE, CORTISONE ACETATE
CORTISPORIN, BACITRACIN ZINC
CORTISPORIN, HYDROCORTISONE
CORTISPORIN, HYDROCORTISONE ACETATE
CORTROSYN, COSYNTROPIN
CORVERT, IBUTILIDE FUMARATE
CORZIDE, BENDROFLUMETHIAZIDE
COSMEGEN, DACTINOMYCIN
COSOPT, DORZOLAMIDE HYDROCHLORIDE
COTRIM D.S., SULFAMETHOXAZOLE
COTRIM, SULFAMETHOXAZOLE
COUMADIN, WARFARIN SODIUM
COVERA-HS, VERAPAMIL HYDROCHLORIDE
COZAAR, LOSARTAN POTASSIUM
CRESTOR, ROSUVASTATIN CALCIUM
CRINONE, PROGESTERONE
CRIXIVAN, INDINAVIR SULFATE
CROLOM, CROMOLYN SODIUM
CROMOLYN SODIUM, CROMOLYN SODIUM
CROMOLYN SODIUM, CROMOLYN SODIUM (OTC)
CROTAN, CROTAMITON
CRYSELLE, ETHINYL ESTRADIOL
C-SOLVE-2, ERYTHROMYCIN
CUBICIN, DAPTOMYCIN
CUPRIC CHLORIDE IN PLASTIC CONTAINER, CUPRIC CHLORIDE
CUPRIMINE, PENICILLAMINE

APPENDIX A - PRODUCT NAME INDEX

A - 14

**** C ****

CUROSURF, PORACTANT ALFA
CUTIVATE, FLUTICASONE PROPIONATE
CYANOCOBALAMIN, CYANOCOBALAMIN
CYANOKIT, HYDROXOCOBALAMIN
CYCLESSA, DESOGESTREL
CYCLOBENZAPRINE HYDROCHLORIDE, CYCLOBENZAPRINE HYDROCHLORIDE
CYCLOGYL, CYCLOPENTOLATE HYDROCHLORIDE
CYCLOMYDRIL, CYCLOPENTOLATE HYDROCHLORIDE
CYCLOPENTOLATE HYDROCHLORIDE, CYCLOPENTOLATE HYDROCHLORIDE
CYCLOPHOSPHAMIDE, CYCLOPHOSPHAMIDE
CYCLOSPORINE, CYCLOSPORINE
CYKLOKAPRON, TRANEXAMIC ACID
CYMBALTA, DULOXETINE HYDROCHLORIDE
CYPROHEPTADINE HYDROCHLORIDE, CYPROHEPTADINE HYDROCHLORIDE
CYPROHEPTADINE HYDROCHLORIDE, CYPROHEPTADINE HYDROCHLORIDE
CYSTADANE, BETAINE HYDROCHLORIDE
CYSTAGON, CYSTEAMINE BITARTRATE
CYSTO-CONRAY II, IOTHALAMATE MEGLUMINE
CYSTOGRAFIN DILUTE, DIATRIZOATE MEGLUMINE
CYSTOGRAFIN, DIATRIZOATE MEGLUMINE
CYTADREN, AMINOGLUTETHIMIDE
CYTARABINE, CYTARABINE
CYTOMEL, LIOTHYRONINE SODIUM
CYTOSAR-U, CYTARABINE
CYTOTEC, MISOPROSTOL
CYTOVENE IV, GANCICLOVIR SODIUM
CYTOXAN, CYCLOPHOSPHAMIDE

**** D ****

D.H.E. 45, DIHYDROERGOTAMINE MESYLATE
DACARBAZINE, DACARBAZINE
DACOGEN, DECITABINE
DALMANE, FLURAZEPAM HYDROCHLORIDE
DANAZOL, DANAZOL
DANTRIUM, DANTROLENE SODIUM
DANTROLENE SODIUM, DANTROLENE SODIUM
DAPSONE, DAPSONE
DARAPRIM, PYRIMETHAMINE
DARVOCET A500, ACETAMINOPHEN
DARVOCET-N 100, ACETAMINOPHEN
DARVOCET-N 50, ACETAMINOPHEN
DARVON COMPOUND-65, ASPIRIN
DARVON, PROPOXYPHENE HYDROCHLORIDE
DARVON-N, PROPOXYPHENE NAPSYLATE
DAUNORUBICIN HYDROCHLORIDE, DAUNORUBICIN HYDROCHLORIDE
DAUNOXOME, DAUNORUBICIN CITRATE
DAYPRO, OXAPROZIN
DAYTRANA, METHYLPHENIDATE
DDAVP (NEEDS NO REFRIGERATION), DESMOPRESSIN ACETATE
DDAVP, DESMOPRESSIN ACETATE
DECLOMYCIN, DEMECLOCYCLINE HYDROCHLORIDE
DEFEROXAMINE MESYLATE, DEFEROXAMINE MESYLATE
DEFINITY, PERFLUTREN
DELATESTRYL, TESTOSTERONE ENANTHATE
DELESTROGEN, ESTRADIOL VALERATE
DELFLEX W/ DEXTROSE 1.5% IN PLASTIC CONTAINER, CALCIUM CHLORIDE
DELFLEX W/ DEXTROSE 1.5% LOW MAGNESIUM IN PLASTIC CONTAINER, CALCIUM CHLORIDE
DELFLEX W/ DEXTROSE 1.5% LOW MAGNESIUM LOW CALCIUM IN PLASTIC CONTAINER, CALCIUM CHLORIDE
DELFLEX W/ DEXTROSE 2.5% IN PLASTIC CONTAINER, CALCIUM CHLORIDE
DELFLEX W/ DEXTROSE 2.5% LOW MAGNESIUM IN PLASTIC CONTAINER, CALCIUM CHLORIDE
DELFLEX W/ DEXTROSE 2.5% LOW MAGNESIUM LOW CALCIUM IN PLASTIC CONTAINER, CALCIUM CHLORIDE
DELFLEX W/ DEXTROSE 3.5% IN PLASTIC CONTAINER, CALCIUM CHLORIDE
DELFLEX W/ DEXTROSE 4.25% IN PLASTIC CONTAINER, CALCIUM CHLORIDE

APPENDIX A - PRODUCT NAME INDEX

A - 15

**** D ****

DELFLX W/ DEXTROSE 4.25% LOW MAGNESIUM IN PLASTIC CONTAINER, CALCIUM CHLORIDE
DELFLX W/ DEXTROSE 4.25% LOW MAGNESIUM LOW CALCIUM IN PLASTIC CONTAINER, CALCIUM CHLORIDE
DELFLX-LM W/ DEXTROSE 1.5% IN PLASTIC CONTAINER, CALCIUM CHLORIDE
DELFLX-LM W/ DEXTROSE 2.5% IN PLASTIC CONTAINER, CALCIUM CHLORIDE
DELFLX-LM W/ DEXTROSE 3.5% IN PLASTIC CONTAINER, CALCIUM CHLORIDE
DELFLX-LM W/ DEXTROSE 4.25% IN PLASTIC CONTAINER, CALCIUM CHLORIDE
DELSYM, DEXTROMETHORPHAN POLISTIREX (OTC)
DEMADEX, TORSEMIDE
DEMECLOCYCLINE HYDROCHLORIDE, DEMECLOCYCLINE HYDROCHLORIDE
DEMEROL, MEPERIDINE HYDROCHLORIDE
DEMSEER, METYROSINE
DEMULEN 1/35-21, ETHINYL ESTRADIOL
DEMULEN 1/35-28, ETHINYL ESTRADIOL
DEMULEN 1/50-21, ETHINYL ESTRADIOL
DEMULEN 1/50-28, ETHINYL ESTRADIOL
DENAVER, PENCICLOVIR SODIUM
DENDRID, IDOXURIDINE
DEPAON, VALPROATE SODIUM
DEPAKENE, VALPROIC ACID
DEPAKOTE ER, DIVALPROEX SODIUM
DEPAKOTE, DIVALPROEX SODIUM
DEPEN, PENICILLAMINE
DEPOCYT, CYTARABINE
DEPODUR, MORPHINE SULFATE
DEPO-ESTRADIOL, ESTRADIOL CYPIONATE
DEPO-MEDROL, METHYLPREDNISOLONE ACETATE
DEPO-PROVERA, MEDROXYPROGESTERONE ACETATE
DEPO-SUBQ PROVERA 104, MEDROXYPROGESTERONE ACETATE
DEPO-TESTOSTERONE, TESTOSTERONE CYPIONATE
DERMABET, BETAMETHASONE VALERATE
DERMA-SMOOTH/FS, FLUOCINOLONE ACETONIDE
DERMATOP E EMOLLIENT, PREDNICARBATE
DERMATOP, PREDNICARBATE
DESFERAL, DEFEROXAMINE MESYLATE
DESIPRAMINE HYDROCHLORIDE, DESIPRAMINE HYDROCHLORIDE
DESMOPRESSIN ACETATE (NEEDS NO REFRIGERATION), DESMOPRESSIN ACETATE
DESMOPRESSIN ACETATE, DESMOPRESSIN ACETATE
DESOGEN, DESOGESTREL
DESOGESTREL AND ETHINYL ESTRADIOL, DESOGESTREL
DESONATE, DESONIDE
DESONIDE, DESONIDE
DESOWEN, DESONIDE
DESOXIMETASONE, DESOXIMETASONE
DESOXYN, METHAMPHETAMINE HYDROCHLORIDE
DETROL LA, TOLTERODINE TARTRATE
DETROL, TOLTERODINE TARTRATE
DEXAMETHASONE INTENSOL, DEXAMETHASONE
DEXAMETHASONE SODIUM PHOSPHATE, DEXAMETHASONE SODIUM PHOSPHATE
DEXAMETHASONE, DEXAMETHASONE
DEXAMETHASONE, DEXAMETHASONE SODIUM PHOSPHATE
DEXASPORIN, DEXAMETHASONE
DEXCHLORPHENIRAMINE MALEATE, DEXCHLORPHENIRAMINE MALEATE
DEXEDRINE, DEXTROAMPHETAMINE SULFATE
DEXFERRUM, IRON DEXTRAN
DEXRAZOXANE, DEXRAZOXANE HYDROCHLORIDE
DEXTROAMP SACCHARATE, AMP ASPARTATE, DEXTROAMP SULFATE AND AMP SULFATE, AMPHETAMINE ASPARTATE
DEXTROAMPHETAMINE SULFATE, DEXTROAMPHETAMINE SULFATE
DEXTROSE 10% AND SODIUM CHLORIDE 0.11% IN PLASTIC CONTAINER, DEXTROSE
DEXTROSE 10% AND SODIUM CHLORIDE 0.2% IN PLASTIC CONTAINER, DEXTROSE
DEXTROSE 10% AND SODIUM CHLORIDE 0.33% IN PLASTIC CONTAINER, DEXTROSE
DEXTROSE 10% AND SODIUM CHLORIDE 0.45% IN PLASTIC CONTAINER, DEXTROSE
DEXTROSE 10% AND SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER, DEXTROSE
DEXTROSE 10% IN PLASTIC CONTAINER, DEXTROSE

APPENDIX A - PRODUCT NAME INDEX

A - 16

**** D ****

DEXTROSE 2.5% AND SODIUM CHLORIDE 0.11% IN PLASTIC CONTAINER, DEXTROSE
DEXTROSE 2.5% AND SODIUM CHLORIDE 0.2% IN PLASTIC CONTAINER, DEXTROSE
DEXTROSE 2.5% AND SODIUM CHLORIDE 0.33% IN PLASTIC CONTAINER, DEXTROSE
DEXTROSE 2.5% AND SODIUM CHLORIDE 0.45% IN PLASTIC CONTAINER, DEXTROSE
DEXTROSE 2.5% AND SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER, DEXTROSE
DEXTROSE 2.5% IN HALF-STRENGTH LACTATED RINGER'S IN PLASTIC CONTAINER, CALCIUM CHLORIDE
DEXTROSE 20% IN PLASTIC CONTAINER, DEXTROSE
DEXTROSE 25%, DEXTROSE
DEXTROSE 3.3% AND SODIUM CHLORIDE 0.3% IN PLASTIC CONTAINER, DEXTROSE
DEXTROSE 30% IN PLASTIC CONTAINER, DEXTROSE
DEXTROSE 40% IN PLASTIC CONTAINER, DEXTROSE
DEXTROSE 5% AND ELECTROLYTE NO 75 IN PLASTIC CONTAINER, DEXTROSE
DEXTROSE 5% AND ELECTROLYTE NO.48 IN PLASTIC CONTAINER, DEXTROSE
DEXTROSE 5% AND LACTATED RINGER'S IN PLASTIC CONTAINER, CALCIUM CHLORIDE
DEXTROSE 5% AND POTASSIUM CHLORIDE 0.075% IN PLASTIC CONTAINER, DEXTROSE
DEXTROSE 5% AND POTASSIUM CHLORIDE 0.15% IN PLASTIC CONTAINER, DEXTROSE
DEXTROSE 5% AND POTASSIUM CHLORIDE 0.224% IN PLASTIC CONTAINER, DEXTROSE
DEXTROSE 5% AND POTASSIUM CHLORIDE 0.3% IN PLASTIC CONTAINER, DEXTROSE
DEXTROSE 5% AND RINGER'S IN PLASTIC CONTAINER, CALCIUM CHLORIDE
DEXTROSE 5% AND SODIUM CHLORIDE 0.11% IN PLASTIC CONTAINER, DEXTROSE
DEXTROSE 5% AND SODIUM CHLORIDE 0.2% IN PLASTIC CONTAINER, DEXTROSE
DEXTROSE 5% AND SODIUM CHLORIDE 0.225% IN PLASTIC CONTAINER, DEXTROSE
DEXTROSE 5% AND SODIUM CHLORIDE 0.3% IN PLASTIC CONTAINER, DEXTROSE
DEXTROSE 5% AND SODIUM CHLORIDE 0.33% IN PLASTIC CONTAINER, DEXTROSE
DEXTROSE 5% AND SODIUM CHLORIDE 0.45% IN PLASTIC CONTAINER, DEXTROSE
DEXTROSE 5% AND SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER, DEXTROSE
DEXTROSE 5% IN ACETATED RINGER'S IN PLASTIC CONTAINER, CALCIUM CHLORIDE
DEXTROSE 5% IN LACTATED RINGER'S IN PLASTIC CONTAINER, CALCIUM CHLORIDE
DEXTROSE 5% IN PLASTIC CONTAINER, DEXTROSE
DEXTROSE 5% IN RINGER'S IN PLASTIC CONTAINER, CALCIUM CHLORIDE
DEXTROSE 5% IN SODIUM CHLORIDE 0.2% IN PLASTIC CONTAINER, DEXTROSE
DEXTROSE 5% IN SODIUM CHLORIDE 0.33% IN PLASTIC CONTAINER, DEXTROSE
DEXTROSE 5% IN SODIUM CHLORIDE 0.45% IN PLASTIC CONTAINER, DEXTROSE
DEXTROSE 5% IN SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER, DEXTROSE
DEXTROSE 5%, SODIUM CHLORIDE 0.2% AND POTASSIUM CHLORIDE 10MEQ, DEXTROSE
DEXTROSE 5%, SODIUM CHLORIDE 0.2% AND POTASSIUM CHLORIDE 15MEQ (K), DEXTROSE
DEXTROSE 5%, SODIUM CHLORIDE 0.2% AND POTASSIUM CHLORIDE 20MEQ (K), DEXTROSE
DEXTROSE 5%, SODIUM CHLORIDE 0.2% AND POTASSIUM CHLORIDE 20MEQ, DEXTROSE
DEXTROSE 5%, SODIUM CHLORIDE 0.2% AND POTASSIUM CHLORIDE 30MEQ, DEXTROSE
DEXTROSE 5%, SODIUM CHLORIDE 0.2% AND POTASSIUM CHLORIDE 40MEQ, DEXTROSE
DEXTROSE 5%, SODIUM CHLORIDE 0.2% AND POTASSIUM CHLORIDE 5MEQ (K), DEXTROSE
DEXTROSE 5%, SODIUM CHLORIDE 0.2% AND POTASSIUM CHLORIDE 5MEQ, DEXTROSE
DEXTROSE 5%, SODIUM CHLORIDE 0.33% AND POTASSIUM CHLORIDE 10MEQ IN PLASTIC CONTAINER, DEXTROSE
DEXTROSE 5%, SODIUM CHLORIDE 0.33% AND POTASSIUM CHLORIDE 15MEQ IN PLASTIC CONTAINER, DEXTROSE
DEXTROSE 5%, SODIUM CHLORIDE 0.33% AND POTASSIUM CHLORIDE 20MEQ IN PLASTIC CONTAINER, DEXTROSE
DEXTROSE 5%, SODIUM CHLORIDE 0.33% AND POTASSIUM CHLORIDE 30MEQ IN PLASTIC CONTAINER, DEXTROSE
DEXTROSE 5%, SODIUM CHLORIDE 0.33% AND POTASSIUM CHLORIDE 40MEQ IN PLASTIC CONTAINER, DEXTROSE
DEXTROSE 5%, SODIUM CHLORIDE 0.33% AND POTASSIUM CHLORIDE 5MEQ IN PLASTIC CONTAINER, DEXTROSE
DEXTROSE 5%, SODIUM CHLORIDE 0.45% AND POTASSIUM CHLORIDE 20MEQ (K) IN PLASTIC CONTAINER, DEXTROSE
DEXTROSE 50% IN PLASTIC CONTAINER, DEXTROSE
DEXTROSE 60% IN PLASTIC CONTAINER, DEXTROSE
DEXTROSE 70% IN PLASTIC CONTAINER, DEXTROSE
DEXTROSTAT, DEXTROAMPHETAMINE SULFATE
DIABETA, GLYBURIDE
DIABINESE, CHLORPROPAMIDE
DIALYTE CONCENTRATE W/ DEXTROSE 30% IN PLASTIC CONTAINER, CALCIUM CHLORIDE
DIALYTE CONCENTRATE W/ DEXTROSE 50% IN PLASTIC CONTAINER, CALCIUM CHLORIDE
DIALYTE LM/ DEXTROSE 1.5% IN PLASTIC CONTAINER, CALCIUM CHLORIDE
DIALYTE LM/ DEXTROSE 2.5% IN PLASTIC CONTAINER, CALCIUM CHLORIDE
DIALYTE LM/ DEXTROSE 4.25% IN PLASTIC CONTAINER, CALCIUM CHLORIDE
DIAMOX, ACETAZOLAMIDE
DIAMOX, ACETAZOLAMIDE SODIUM

APPENDIX A - PRODUCT NAME INDEX

A - 17

**** D ****

DIANEAL 137 W/ DEXTROSE 1.5% IN PLASTIC CONTAINER, CALCIUM CHLORIDE
DIANEAL 137 W/ DEXTROSE 2.5% IN PLASTIC CONTAINER, CALCIUM CHLORIDE
DIANEAL 137 W/ DEXTROSE 4.25% IN PLASTIC CONTAINER, CALCIUM CHLORIDE
DIANEAL LOW CALCIUM W/ DEXTROSE 1.5% IN PLASTIC CONTAINER, CALCIUM CHLORIDE
DIANEAL LOW CALCIUM W/ DEXTROSE 2.5% IN PLASTIC CONTAINER, CALCIUM CHLORIDE
DIANEAL LOW CALCIUM W/ DEXTROSE 3.5% IN PLASTIC CONTAINER, CALCIUM CHLORIDE
DIANEAL LOW CALCIUM W/ DEXTROSE 4.25% IN PLASTIC CONTAINER, CALCIUM CHLORIDE
DIANEAL PD-1 W/ DEXTROSE 1.5% IN PLASTIC CONTAINER, CALCIUM CHLORIDE
DIANEAL PD-1 W/ DEXTROSE 2.5% IN PLASTIC CONTAINER, CALCIUM CHLORIDE
DIANEAL PD-1 W/ DEXTROSE 3.5% IN PLASTIC CONTAINER, CALCIUM CHLORIDE
DIANEAL PD-1 W/ DEXTROSE 4.25% IN PLASTIC CONTAINER, CALCIUM CHLORIDE
DIANEAL PD-2 W/ DEXTROSE 1.5% IN PLASTIC CONTAINER, CALCIUM CHLORIDE
DIANEAL PD-2 W/ DEXTROSE 2.5% IN PLASTIC CONTAINER, CALCIUM CHLORIDE
DIANEAL PD-2 W/ DEXTROSE 3.5% IN PLASTIC CONTAINER, CALCIUM CHLORIDE
DIANEAL PD-2 W/ DEXTROSE 4.25% IN PLASTIC CONTAINER, CALCIUM CHLORIDE
DIASTAT ACUDIAL, DIAZEPAM
DIASTAT, DIAZEPAM
DIAZEPAM INTENSOL, DIAZEPAM
DIAZEPAM, DIAZEPAM
DIBENZYLINE, PHENOXYBENZAMINE HYDROCHLORIDE
DICLOFENAC POTASSIUM, DICLOFENAC POTASSIUM
DICLOFENAC SODIUM, DICLOFENAC SODIUM
DICLOXACILLIN SODIUM, DICLOXACILLIN SODIUM
DICYCLOMINE HYDROCHLORIDE (PRESERVATIVE-FREE), DICYCLOMINE HYDROCHLORIDE
DICYCLOMINE HYDROCHLORIDE, DICYCLOMINE HYDROCHLORIDE
DIDANOSINE, DIDANOSINE
DIDREX, BENZPHETAMINE HYDROCHLORIDE
DIDRONEL, ETIDRONATE DISODIUM
DIFFERIN, ADAPALENE
DIFLORASONE DIACETATE, DIFLORASONE DIACETATE
DIFLUCAN IN DEXTROSE 5% IN PLASTIC CONTAINER, FLUCONAZOLE
DIFLUCAN IN SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER, FLUCONAZOLE
DIFLUCAN IN SODIUM CHLORIDE 0.9%, FLUCONAZOLE
DIFLUCAN, FLUCONAZOLE
DIFLUNISAL, DIFLUNISAL
DIGOXIN PEDIATRIC, DIGOXIN
DIGOXIN, DIGOXIN
DIHYDROERGOTAMINE MESYLATE, DIHYDROERGOTAMINE MESYLATE
DILACOR XR, DILTIAZEM HYDROCHLORIDE
DILANTIN, PHENYTOIN
DILANTIN, PHENYTOIN SODIUM
DILANTIN-125, PHENYTOIN
DILATRATE-SR, ISOSORBIDE DINITRATE
DILAUDID, HYDROMORPHONE HYDROCHLORIDE
DILAUDID-HP, HYDROMORPHONE HYDROCHLORIDE
DILT-CD, DILTIAZEM HYDROCHLORIDE
DILTIAZEM HYDROCHLORIDE, DILTIAZEM HYDROCHLORIDE
DILTZAC, DILTIAZEM HYDROCHLORIDE
DIMENHYDRINATE, DIMENHYDRINATE
DIMETHYL SULFOXIDE, DIMETHYL SULFOXIDE
DIOVAN HCT, HYDROCHLOROTHIAZIDE
DIOVAN, VALSARTAN
DIPENTUM, OLSALAZINE SODIUM
DIPHENHYDRAMINE HYDROCHLORIDE PRESERVATIVE FREE, DIPHENHYDRAMINE HYDROCHLORIDE
DIPHENHYDRAMINE HYDROCHLORIDE, DIPHENHYDRAMINE HYDROCHLORIDE
DIPHENOXYLATE HYDROCHLORIDE AND ATROPINE SULFATE, ATROPINE SULFATE
DIPHENOXYLATE HYDROCHLORIDE W/ ATROPINE SULFATE, ATROPINE SULFATE
DIPIVEFRIN HYDROCHLORIDE, DIPIVEFRIN HYDROCHLORIDE
DIPRIVAN, PROPOFOL
DIPROLENE AF, BETAMETHASONE DIPROPIONATE
DIPROLENE, BETAMETHASONE DIPROPIONATE
DIPYRIDAMOLE, DIPYRIDAMOLE
DISOPHROL, DEXBROMPHENIRAMINE MALEATE (OTC)

APPENDIX A - PRODUCT NAME INDEX

A - 18

**** D ****

DISOPYRAMIDE PHOSPHATE, DISOPYRAMIDE PHOSPHATE
DISPERMOX, AMOXICILLIN
DITROPAN XL, OXYBUTYNIN CHLORIDE
DITROPAN, OXYBUTYNIN CHLORIDE
DIURIL, CHLOROTHIAZIDE
DIURIL, CHLOROTHIAZIDE SODIUM
DOBUTAMINE HYDROCHLORIDE IN DEXTROSE 5% IN PLASTIC CONTAINER, DOBUTAMINE HYDROCHLORIDE
DOBUTAMINE HYDROCHLORIDE IN DEXTROSE 5%, DOBUTAMINE HYDROCHLORIDE
DOBUTAMINE HYDROCHLORIDE, DOBUTAMINE HYDROCHLORIDE
DOLOPHINE HYDROCHLORIDE, METHADONE HYDROCHLORIDE
DOPAMINE HYDROCHLORIDE AND DEXTROSE 5% IN PLASTIC CONTAINER, DOPAMINE HYDROCHLORIDE
DOPAMINE HYDROCHLORIDE AND DEXTROSE 5%, DOPAMINE HYDROCHLORIDE
DOPAMINE HYDROCHLORIDE IN DEXTROSE 5% IN PLASTIC CONTAINER, DOPAMINE HYDROCHLORIDE
DOPAMINE HYDROCHLORIDE, DOPAMINE HYDROCHLORIDE
DOPRAM, DOXAPRAM HYDROCHLORIDE
DORAL, QUAZEPAM
DORYX, DOXYCYCLINE HYCLATE
DOSTINEX, CABERGOLINE
DOVONEX, CALCIPOTRIENE
DOXAPRAM HYDROCHLORIDE, DOXAPRAM HYDROCHLORIDE
DOXAZOSIN MESYLATE, DOXAZOSIN MESYLATE
DOXEPIN HYDROCHLORIDE, DOXEPIN HYDROCHLORIDE
DOXIL, DOXORUBICIN HYDROCHLORIDE
DOXORUBICIN HYDROCHLORIDE, DOXORUBICIN HYDROCHLORIDE
DOXY 100, DOXYCYCLINE HYCLATE
DOXY 200, DOXYCYCLINE HYCLATE
DOXYCYCLINE HYCLATE, DOXYCYCLINE HYCLATE
DOXYCYCLINE, DOXYCYCLINE
DOXYCYCLINE, DOXYCYCLINE HYCLATE
DOXYLAMINE SUCCINATE, DOXYLAMINE SUCCINATE (OTC)
DRAXIMAGE MDP, TECHNETIUM TC-99M MEDRONATE KIT
DRICORT, HYDROCORTISONE ACETATE
DRISDOL, ERGOCALCIFEROL
DRIXORAL PLUS, ACETAMINOPHEN (OTC)
DRIXORAL, DEXBROMPHENIRAMINE MALEATE (OTC)
DROPERIDOL, DROPERIDOL
DROXIA, HYDROXYUREA
DTIC-DOME, DACARBAZINE
DTPA, TECHNETIUM TC-99M PENTETATE KIT
DUAC, BENZOYL PEROXIDE
DUETACT, GLIMEPIRIDE
DUOCAINE, BUPIVACAINE HYDROCHLORIDE
DUODOTE, ATROPINE
DUONEB, ALBUTEROL SULFATE
DURACLON, CLONIDINE HYDROCHLORIDE
DURAGESIC-100, FENTANYL
DURAGESIC-12, FENTANYL
DURAGESIC-25, FENTANYL
DURAGESIC-50, FENTANYL
DURAGESIC-75, FENTANYL
DURAMORPH PF, MORPHINE SULFATE
DURAPREP, IODINE POVACRYLEX (OTC)
DURICEF, CEFADROXIL/CEFADROXIL HEMIHYDRATE
DUTOPROL, HYDROCHLOROTHIAZIDE
DUVOID, BETHANECHOL CHLORIDE
DYAZIDE, HYDROCHLOROTHIAZIDE
DYNACIN, MINOCYCLINE HYDROCHLORIDE
DYNACIRC CR, ISRADIPINE
DYNA-HEX, CHLORHEXIDINE GLUCONATE (OTC)
DYRENIUM, TRIAMTERENE

**** E ****

E.E.S. 200, ERYTHROMYCIN ETHYLSUCCINATE

APPENDIX A - PRODUCT NAME INDEX

A - 19

**** E ****

E.E.S. 400, ERYTHROMYCIN ETHYLSUCCINATE
E.E.S., ERYTHROMYCIN ETHYLSUCCINATE
E-BASE, ERYTHROMYCIN
EC-NAPROSYN, NAPROXEN
ECONAZOLE NITRATE, ECONAZOLE NITRATE
EDECIN, ETHACRYNATE SODIUM
EDECIN, ETHACRYNIC ACID
EDETATE DISODIUM, EDETATE DISODIUM
EDEX, ALPROSTADIL
EDROPHONIUM CHLORIDE, EDROPHONIUM CHLORIDE
EFFEXOR XR, VENLAFAXINE HYDROCHLORIDE
EFFEXOR, VENLAFAXINE HYDROCHLORIDE
EFIDAC 24 PSEUDOEPHEDRINE HYDROCHLORIDE, PSEUDOEPHEDRINE HYDROCHLORIDE (OTC)
EFIDAC 24 PSEUDOEPHEDRINE HYDROCHLORIDE/BROMPHENIRAMINE MALEATE, BROMPHENIRAMINE MALEATE (OTC)
EFUDEX, FLUOROURACIL
E-GLADES, ERYTHROMYCIN
ELDEPRYL, SELEGILINE HYDROCHLORIDE
ELESTAT, EPINASTINE HYDROCHLORIDE
ELESTRIN, ESTRADIOL
ELIDEL, PIMECROLIMUS
ELIGARD, LEUPROLIDE ACETATE
ELIMITE, PERMETHRIN
ELIXOPHYLLIN, THEOPHYLLINE
ELLENCE, EPIRUBICIN HYDROCHLORIDE
ELLIOTTS B SOLUTION, CALCIUM CHLORIDE
ELMIRON, PENTOSAN POLYSULFATE SODIUM
ELOCON, MOMETASONE FUROATE
ELOXATIN, OXALIPLATIN
EMADINE, EMEDASTINE DIFUMARATE
EMBELINE E, CLOBETASOL PROPIONATE
EMBELINE, CLOBETASOL PROPIONATE
EMCYT, ESTRAMUSTINE PHOSPHATE SODIUM
EMEND, APREPITANT
EMLA, LIDOCAINE
EMSAM, SELEGILINE
EMTRIVA, EMTRICITABINE
E-MYCIN, ERYTHROMYCIN
ENABLEX, DARIFENACIN HYDROBROMIDE
ENALAPRIL MALEATE AND HYDROCHLOROTHIAZIDE, ENALAPRIL MALEATE
ENALAPRIL MALEATE, ENALAPRIL MALEATE
ENALAPRILAT, ENALAPRILAT
ENDOSOL EXTRA, CALCIUM CHLORIDE
ENDRATE, EDETATE DISODIUM
ENDURON, METHYLCLOTHIAZIDE
ENFLURANE, ENFLURANE
ENJUVIA, ESTROGENS, CONJUGATED SYNTHETIC B
ENLON, EDROPHONIUM CHLORIDE
ENLON-PLUS, ATROPINE SULFATE
ENPRESSE-28, ETHINYL ESTRADIOL
ENTOCORT EC, BUDESONIDE
ENULOSE, LACTULOSE
EPIFOAM, HYDROCORTISONE ACETATE
EPINEPHRINE, EPINEPHRINE (OTC)
EPIPEN JR., EPINEPHRINE
EPIPEN, EPINEPHRINE
EPIRUBICIN HYDROCHLORIDE, EPIRUBICIN HYDROCHLORIDE
EPITOL, CARBAMAZEPINE
EPIVIR, LAMIVUDINE
EPIVIR-HBV, LAMIVUDINE
EPZICOM, ABACAVIR SULFATE
EQUAGESIC, ASPIRIN
EQUETRO, CARBAMAZEPINE
ERAXIS, ANIDULAFUNGIN

APPENDIX A - PRODUCT NAME INDEX

A - 20

**** E ****

ERGOLOID MESYLATES, ERGOLOID MESYLATES
ERGOMAR, ERGOTAMINE TARTRATE
ERGOTAMINE TARTRATE AND CAFFEINE, CAFFEINE
ERRIN, NORETHINDRONE
ERTACZO, SERTACONAZOLE NITRATE
ERYC, ERYTHROMYCIN
ERYCETTE, ERYTHROMYCIN
ERYDERM, ERYTHROMYCIN
ERYGEL, ERYTHROMYCIN
ERYPED, ERYTHROMYCIN ETHYLSUCCINATE
ERY-TAB, ERYTHROMYCIN
ERYTHRA-DERM, ERYTHROMYCIN
ERYTHROCIN STEARATE, ERYTHROMYCIN STEARATE
ERYTHROCIN, ERYTHROMYCIN LACTOBIONATE
ERYTHROMYCIN AND BENZOYL PEROXIDE, BENZOYL PEROXIDE
ERYTHROMYCIN ESTOLATE, ERYTHROMYCIN ESTOLATE
ERYTHROMYCIN ETHYLSUCCINATE AND SULFISOXAZOLE ACETYL, ERYTHROMYCIN ETHYLSUCCINATE
ERYTHROMYCIN ETHYLSUCCINATE, ERYTHROMYCIN ETHYLSUCCINATE
ERYTHROMYCIN LACTOBIONATE, ERYTHROMYCIN LACTOBIONATE
ERYTHROMYCIN STEARATE, ERYTHROMYCIN STEARATE
ERYTHROMYCIN, ERYTHROMYCIN
ERYZOLE, ERYTHROMYCIN ETHYLSUCCINATE
ESCITALOPRAM OXALATE, ESCITALOPRAM OXALATE
ESGIC-PLUS, ACETAMINOPHEN
ESIDRIX, HYDROCHLOROTHIAZIDE
ESKALITH, LITHIUM CARBONATE
ESMOLOL HYDROCHLORIDE, ESMOLOL HYDROCHLORIDE
E-SOLVE 2, ERYTHROMYCIN
ESTAZOLAM, ESTAZOLAM
ESTRACE, ESTRADIOL
ESTRADERM, ESTRADIOL
ESTRADIOL AND NORGESTIMATE, ESTRADIOL
ESTRADIOL CYPIONATE, ESTRADIOL CYPIONATE
ESTRADIOL VALERATE, ESTRADIOL VALERATE
ESTRADIOL, ESTRADIOL
ESTRASORB, ESTRADIOL HEMIHYDRATE
ESTRING, ESTRADIOL
ESTROGEL, ESTRADIOL
ESTRONE, ESTRONE
ESTROPIPATE, ESTROPIPATE
ESTROSTEP FE, ETHINYL ESTRADIOL
ETHAMBUTOL HYDROCHLORIDE, ETHAMBUTOL HYDROCHLORIDE
ETHAMOLIN, ETHANOLAMINE OLEATE
ETHIODOL, ETHIODIZED OIL
ETHMOZINE, MORICIZINE HYDROCHLORIDE
ETHOSUXIMIDE, ETHOSUXIMIDE
ETHRANE, ENFLURANE
ETHYOL, AMIFOSTINE
ETIDRONATE DISODIUM, ETIDRONATE DISODIUM
ETODOLAC, ETODOLAC
ETOMIDATE, ETOMIDATE
ETOPOPHOS PRESERVATIVE FREE, ETOPOSIDE PHOSPHATE
ETOPOSIDE, ETOPOSIDE
EURAX, CROTAMITON
EVALOSE, LACTULOSE
EVISTA, RALOXIFENE HYDROCHLORIDE
EVOCLIN, CLINDAMYCIN PHOSPHATE
EVOXAC, CEVIMELINE HYDROCHLORIDE
EXCEDRIN (MIGRAINE), ACETAMINOPHEN (OTC)
EXELDERM, SULCONAZOLE NITRATE
EXELON, RIVASTIGMINE TARTRATE
EXIDINE, CHLORHEXIDINE GLUCONATE (OTC)
EXJADE, DEFERASIROX

APPENDIX A - PRODUCT NAME INDEX

A - 21

**** E ****

EXTENDED PHENYTOIN SODIUM, PHENYTOIN SODIUM
EXTRANEAL, ICODextrin
EXTRA-STRENGTH AIM, SODIUM MONOFLUOROPHOSPHATE (OTC)
EXUBERA, INSULIN RECOMBINANT HUMAN
E-Z SCRUB 201, POVIDONE-IODINE (OTC)
E-Z SCRUB 241, POVIDONE-IODINE (OTC)

**** F ****

FACTIVE, GEMIFLOXACIN MESYLATE
FAMOTIDINE PRESERVATIVE FREE (PHARMACY BULK), FAMOTIDINE
FAMOTIDINE PRESERVATIVE FREE IN PLASTIC CONTAINER, FAMOTIDINE
FAMOTIDINE PRESERVATIVE FREE, FAMOTIDINE
FAMOTIDINE, FAMOTIDINE
FAMOTIDINE, FAMOTIDINE (OTC)
FAMVIR, FAMCICLOVIR
FANSIDAR, PYRIMETHAMINE
FARESTON, TOREMIFENE CITRATE
FASLODEX, FULVESTRANT
FAZACLO ODT, CLOZAPINE
FELBATOL, FELBAMATE
FELDENE, PIROXICAM
FELODIPINE, FELODIPINE
FEMARA, LETROZOLE
FEMHRT, ETHINYL ESTRADIOL
FEMRING, ESTRADIOL ACETATE
FEMSTAT 3, BUTOCONAZOLE NITRATE (OTC)
FEMTRACE, ESTRADIOL ACETATE
FENOFIBRATE (MICRONIZED), FENOFIBRATE
FENOFIBRATE, FENOFIBRATE
FENOLDOPAM MESYLATE, FENOLDOPAM MESYLATE
FENOPROFEN CALCIUM, FENOPROFEN CALCIUM
FENTANYL CITRATE PRESERVATIVE FREE, FENTANYL CITRATE
FENTANYL CITRATE, FENTANYL CITRATE
FENTANYL, FENTANYL
FENTORA, FENTANYL CITRATE
FERIDEX I.V., FERUMOXIDES
FERRLECIT, SODIUM FERRIC GLUCONATE COMPLEX
FEXOFENADINE HYDROCHLORIDE AND PSEUDOEPHEDRINE HYDROCHLORIDE, FEXOFENADINE HYDROCHLORIDE
FEXOFENADINE HYDROCHLORIDE, FEXOFENADINE HYDROCHLORIDE
FINACEA, AZELAIC ACID
FINASTERIDE, FINASTERIDE
FIORICET W/ CODEINE, ACETAMINOPHEN
FIORICET, ACETAMINOPHEN
FIORINAL W/CODEINE, ASPIRIN
FIORINAL, ASPIRIN
FLAGYL ER, METRONIDAZOLE
FLAGYL I.V. RTU IN PLASTIC CONTAINER, METRONIDAZOLE
FLAGYL I.V., METRONIDAZOLE HYDROCHLORIDE
FLAGYL, METRONIDAZOLE
FLAREX, FLUOROMETHOLONE ACETATE
FLAVORED COLESTID, COLESTIPOL HYDROCHLORIDE
FLAVOXATE HYDROCHLORIDE, FLAVOXATE HYDROCHLORIDE
FLECAINIDE ACETATE, FLECAINIDE ACETATE
FLEXERIL, CYCLOBENZAPRINE HYDROCHLORIDE
FLOLAN, EPOPROSTENOL SODIUM
FLOMAX, TAMSULOSIN HYDROCHLORIDE
FLONASE, FLUTICASONE PROPIONATE
FLORINEF, FLUDROCORTISONE ACETATE
FLOVENT DISKUS 100, FLUTICASONE PROPIONATE
FLOVENT DISKUS 250, FLUTICASONE PROPIONATE
FLOVENT DISKUS 50, FLUTICASONE PROPIONATE
FLOVENT HFA, FLUTICASONE PROPIONATE
FLOVENT, FLUTICASONE PROPIONATE

APPENDIX A - PRODUCT NAME INDEX

A - 22

**** F ****

FLOXIN OTIC, OFLOXACIN
FLOXIN, OFLOXACIN
FLOXURIDINE, FLOXURIDINE
FLUCONAZOLE IN DEXTROSE 5% IN PLASTIC CONTAINER, FLUCONAZOLE
FLUCONAZOLE IN SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER, FLUCONAZOLE
FLUCONAZOLE IN SODIUM CHLORIDE 0.9%, FLUCONAZOLE
FLUCONAZOLE, FLUCONAZOLE
FLUDARA, FLUDARABINE PHOSPHATE
FLUDARABINE PHOSPHATE, FLUDARABINE PHOSPHATE
FLUDEOXYGLUCOSE F 18, FLUDEOXYGLUCOSE F-18
FLUDEOXYGLUCOSE F18, FLUDEOXYGLUCOSE F-18
FLUDROCORTISONE ACETATE, FLUDROCORTISONE ACETATE
FLUMADINE, RIMANTADINE HYDROCHLORIDE
FLUMAZENIL, FLUMAZENIL
FLUNISOLIDE, FLUNISOLIDE
FLUOCINOLONE ACETONIDE, FLUOCINOLONE ACETONIDE
FLUOCINONIDE EMULSIFIED BASE, FLUOCINONIDE
FLUOCINONIDE, FLUOCINONIDE
FLUORESCITE, FLUORESCIN SODIUM
FLUOR-OP, FLUOROMETHOLONE
FLUOROPLEX, FLUOROURACIL
FLUOROURACIL, FLUOROURACIL
FLUOXETINE HYDROCHLORIDE, FLUOXETINE HYDROCHLORIDE
FLUOXETINE, FLUOXETINE HYDROCHLORIDE
FLUOXYMESTERONE, FLUOXYMESTERONE
FLUPHENAZINE DECANOATE, FLUPHENAZINE DECANOATE
FLUPHENAZINE HYDROCHLORIDE, FLUPHENAZINE HYDROCHLORIDE
FLURAZEPAM HYDROCHLORIDE, FLURAZEPAM HYDROCHLORIDE
FLURBIPROFEN SODIUM, FLURBIPROFEN SODIUM
FLURBIPROFEN, FLURBIPROFEN
FLUTAMIDE, FLUTAMIDE
FLUTICASONE PROPIONATE, FLUTICASONE PROPIONATE
FLUVOXAMINE MALEATE, FLUVOXAMINE MALEATE
FLUXID, FAMOTIDINE
FML FORTE, FLUOROMETHOLONE
FML, FLUOROMETHOLONE
FML-S, FLUOROMETHOLONE
FOAMCOAT, ALUMINUM HYDROXIDE (OTC)
FOCALIN XR, DEXMETHYLPHENIDATE HYDROCHLORIDE
FOCALIN, DEXMETHYLPHENIDATE HYDROCHLORIDE
FOLIC ACID, FOLIC ACID
FOLICET, FOLIC ACID
FOLLISTIM AQ, FOLLITROPIN ALFA/BETA
FORADIL CERTIHALER, FORMOTEROL FUMARATE
FORADIL, FORMOTEROL FUMARATE
FORANE, ISOFLURANE
FORTAMET, METFORMIN HYDROCHLORIDE
FORTAZ IN PLASTIC CONTAINER, CEFTAZIDIME SODIUM
FORTAZ, CEFTAZIDIME
FORTEO, TERIPARATIDE RECOMBINANT HUMAN
FORTICAL, CALCITONIN SALMON RECOMBINANT
FOSAMAX PLUS D, ALENDRONATE SODIUM
FOSAMAX, ALENDRONATE SODIUM
FOSCARNET SODIUM, FOSCARNET SODIUM
FOSCAVIR, FOSCARNET SODIUM
FOSINOPRIL SODIUM AND HYDROCHLOROTHIAZIDE, FOSINOPRIL SODIUM
FOSINOPRIL SODIUM, FOSINOPRIL SODIUM
FOSRENOL, LANTHANUM CARBONATE
FRAGMIN, DALTEPARIN SODIUM
FREAMINE HBC 6.9%, AMINO ACIDS
FREAMINE III 10%, AMINO ACIDS
FREAMINE III 3% W/ ELECTROLYTES, AMINO ACIDS
FREAMINE III 8.5% W/ ELECTROLYTES, AMINO ACIDS

APPENDIX A - PRODUCT NAME INDEX

A - 23

**** F ****

FREAMINE III 8.5%, AMINO ACIDS
FROVA, FROVATRIPTAN SUCCINATE
FS SHAMPOO, FLUOCINOLONE ACETONIDE
FUDR, FLOXURIDINE
FULVICIN-U/F, GRISEOFULVIN, MICROCRYSTALLINE
FUNGIZONE, AMPHOTERICIN B
FURADANTIN, NITROFURANTOIN
FUROSEMIDE, FUROSEMIDE
FUZEON, ENFUVIRTIDE

**** G ****

GABAPENTIN, GABAPENTIN
GABITRIL, TIAGABINE HYDROCHLORIDE
GALLIUM CITRATE GA 67, GALLIUM CITRATE, GA-67
GALZIN, ZINC ACETATE
GANCICLOVIR SODIUM, GANCICLOVIR SODIUM
GANCICLOVIR, GANCICLOVIR
GANIRELIX ACETATE INJECTION, GANIRELIX ACETATE
GANITE, GALLIUM NITRATE
GANTRISIN PEDIATRIC, SULFISOXAZOLE ACETYL
GASTROCROM, CROMOLYN SODIUM
GASTROGRAFIN, DIATRIZOATE MEGLUMINE
GASTROMARK, FERUMOXASIL
GAVISCON, ALUMINUM HYDROXIDE (OTC)
GEMFIBROZIL, GEMFIBROZIL
GEMZAR, GEMCITABINE HYDROCHLORIDE
GENCEPT 10/11-21, ETHINYL ESTRADIOL
GENCEPT 10/11-28, ETHINYL ESTRADIOL
GENERLAC, LACTULOSE
GENGRAF, CYCLOSPORINE
GENOPTIC, GENTAMICIN SULFATE
GENOTROPIN PRESERVATIVE FREE, SOMATROPIN RECOMBINANT
GENOTROPIN, SOMATROPIN RECOMBINANT
GENTAK, GENTAMICIN SULFATE
GENTAMICIN SULFATE IN SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER, GENTAMICIN SULFATE
GENTAMICIN SULFATE, GENTAMICIN SULFATE
GENTAMICIN, GENTAMICIN SULFATE
GEN-XENE, CLORAZEPATE DIPOTASSIUM
GEOCILLIN, CARBENICILLIN INDANYL SODIUM
GEODON, ZIPRASIDONE HYDROCHLORIDE
GEODON, ZIPRASIDONE MESYLATE
GEREF, SERMORELIN ACETATE
GERIMAL, ERGOLOID MESYLATES
GLEEVEC, IMATINIB MESYLATE
GLIADEL, CARMUSTINE
GLIMEPIRIDE, GLIMEPIRIDE
GLIPIZIDE AND METFORMIN HYDROCHLORIDE, GLIPIZIDE
GLIPIZIDE, GLIPIZIDE
GLOFIL-125, IOTHALAMATE SODIUM, I-125
GLUCAGEN, GLUCAGON HYDROCHLORIDE RECOMBINANT
GLUCAGON, GLUCAGON RECOMBINANT
GLUCOPHAGE XR, METFORMIN HYDROCHLORIDE
GLUCOPHAGE, METFORMIN HYDROCHLORIDE
GLUCOTROL XL, GLIPIZIDE
GLUCOTROL, GLIPIZIDE
GLUCOVANCE, GLYBURIDE
GLUMETZA, METFORMIN HYDROCHLORIDE
GLYBURIDE (MICRONIZED), GLYBURIDE
GLYBURIDE AND METFORMIN HYDROCHLORIDE, GLYBURIDE
GLYBURIDE, GLYBURIDE
GLYCINE 1.5% IN PLASTIC CONTAINER, GLYCINE
GLYCOLAX, POLYETHYLENE GLYCOL 3350
GLYCOPYRROLATE, GLYCOPYRROLATE

APPENDIX A - PRODUCT NAME INDEX

A - 24

**** G ****

GLYNASE, GLYBURIDE
GLYSET, MIGLITOL
GO-EVAC, POLYETHYLENE GLYCOL 3350
GOLYTELY, POLYETHYLENE GLYCOL 3350
GONAL-F RFF PEN, FOLLITROPIN ALFA/BETA
GONAL-F RFF, FOLLITROPIN ALFA/BETA
GONAL-F, FOLLITROPIN ALFA/BETA
GRIFULVIN V, GRISEOFULVIN, MICROCRYSTALLINE
GRISEOFULVIN, GRISEOFULVIN, MICROCRYSTALLINE
GRIS-PEG, GRISEOFULVIN, ULTRAMICROCRYSTALLINE
GUANABENZ ACETATE, GUANABENZ ACETATE
GUANFACINE HYDROCHLORIDE, GUANFACINE HYDROCHLORIDE
GUANIDINE HYDROCHLORIDE, GUANIDINE HYDROCHLORIDE
GYNAZOLE-1, BUTOCONAZOLE NITRATE
GYNE-LOTRIMIN 3 COMBINATION PACK, CLOTRIMAZOLE (OTC)
GYNE-LOTRIMIN 3, CLOTRIMAZOLE (OTC)
GYNE-LOTRIMIN COMBINATION PACK, CLOTRIMAZOLE (OTC)
GYNE-LOTRIMIN, CLOTRIMAZOLE (OTC)
GYNIX, CLOTRIMAZOLE (OTC)
GYNODIOL, ESTRADIOL

**** H ****

H.P. ACTHAR GEL, CORTICOTROPIN
HABITROL, NICOTINE (OTC)
HALCION, TRIAZOLAM
HALDOL, HALOPERIDOL DECANOATE
HALDOL, HALOPERIDOL LACTATE
HALFLYTELY, BISACODYL
HALOBETASOL PROPIONATE, HALOBETASOL PROPIONATE
HALOG, HALCINONIDE
HALOPERIDOL DECANOATE, HALOPERIDOL DECANOATE
HALOPERIDOL LACTATE, HALOPERIDOL LACTATE
HALOPERIDOL, HALOPERIDOL
HALOPERIDOL, HALOPERIDOL LACTATE
HALOTESTIN, FLUOXYMESTERONE
HECTOROL, DOXERCALCIFEROL
HELIDAC, BISMUTH SUBSALICYLATE
HEMABATE, CARBOPROST TROMETHAMINE
HEPARIN LOCK FLUSH IN PLASTIC CONTAINER, HEPARIN SODIUM
HEPARIN LOCK FLUSH, HEPARIN SODIUM
HEPARIN SODIUM 1,000 UNITS AND SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER, HEPARIN SODIUM
HEPARIN SODIUM 1,000 UNITS IN SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER, HEPARIN SODIUM
HEPARIN SODIUM 10,000 UNITS IN DEXTROSE 5% IN PLASTIC CONTAINER, HEPARIN SODIUM
HEPARIN SODIUM 12,500 UNITS IN DEXTROSE 5% IN PLASTIC CONTAINER, HEPARIN SODIUM
HEPARIN SODIUM 12,500 UNITS IN SODIUM CHLORIDE 0.45% IN PLASTIC CONTAINER, HEPARIN SODIUM
HEPARIN SODIUM 2,000 UNITS AND SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER, HEPARIN SODIUM
HEPARIN SODIUM 2,000 UNITS IN SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER, HEPARIN SODIUM
HEPARIN SODIUM 20,000 UNITS AND DEXTROSE 5% IN PLASTIC CONTAINER, HEPARIN SODIUM
HEPARIN SODIUM 20,000 UNITS IN DEXTROSE 5% IN PLASTIC CONTAINER, HEPARIN SODIUM
HEPARIN SODIUM 25,000 UNITS AND DEXTROSE 5% IN PLASTIC CONTAINER, HEPARIN SODIUM
HEPARIN SODIUM 25,000 UNITS IN DEXTROSE 5% IN PLASTIC CONTAINER, HEPARIN SODIUM
HEPARIN SODIUM 25,000 UNITS IN SODIUM CHLORIDE 0.45% IN PLASTIC CONTAINER, HEPARIN SODIUM
HEPARIN SODIUM IN PLASTIC CONTAINER, HEPARIN SODIUM
HEPARIN SODIUM PRESERVATIVE FREE, HEPARIN SODIUM
HEPARIN SODIUM, HEPARIN SODIUM
HEPATAMINE 8%, AMINO ACIDS
HEPATASOL 8%, AMINO ACIDS
HEPATOLITE, TECHNETIUM TC-99M DISOFENIN KIT
HEPSERA, ADEFOVIR DIPIVOXIL
HEPTALAC, LACTULOSE
HERPLEX, IDOXURIDINE
HEXABRIX, IOXAGLATE MEGLUMINE
HEXALEN, ALTRETAMINE

APPENDIX A - PRODUCT NAME INDEX

A - 25

**** H ****

HIBICLENS, CHLORHEXIDINE GLUCONATE (OTC)
HIBISTAT, CHLORHEXIDINE GLUCONATE (OTC)
HI-COR, HYDROCORTISONE
HIPREX, METHENAMINE HIPPURATE
HIVID, ZALCITABINE
HMS, MEDRYSONE
HOMATROPRINE METHYLBROMIDE AND HYDROCODONE BITARTRATE, HOMATROPINE METHYLBROMIDE
HUMABID, GUAIFENESIN (OTC)
HUMALOG MIX 50/50, INSULIN LISPRO PROTAMINE RECOMBINANT
HUMALOG MIX 75/25, INSULIN LISPRO PROTAMINE RECOMBINANT
HUMALOG PEN, INSULIN LISPRO RECOMBINANT
HUMALOG, INSULIN LISPRO RECOMBINANT
HUMATIN, PAROMOMYCIN SULFATE
HUMATROPE, SOMATROPIN RECOMBINANT
HUMULIN 50/50, INSULIN RECOMBINANT HUMAN (OTC)
HUMULIN 70/30 PEN, INSULIN RECOMBINANT HUMAN (OTC)
HUMULIN 70/30, INSULIN RECOMBINANT HUMAN (OTC)
HUMULIN L, INSULIN ZINC SUSP RECOMBINANT HUMAN (OTC)
HUMULIN N, INSULIN SUSP ISOPHANE RECOMBINANT HUMAN (OTC)
HUMULIN R PEN, INSULIN RECOMBINANT HUMAN (OTC)
HUMULIN R, INSULIN RECOMBINANT HUMAN
HUMULIN R, INSULIN RECOMBINANT HUMAN (OTC)
HYCAMTIN, TOPOTECAN HYDROCHLORIDE
HYCODAN, HOMATROPINE METHYLBROMIDE
HYDASE, HYALURONIDASE
HYDERGINE, ERGOLOID MESYLATES
HYDRALAZINE HYDROCHLORIDE, HYDRALAZINE HYDROCHLORIDE
HYDRA-ZIDE, HYDRALAZINE HYDROCHLORIDE
HYDREA, HYDROXYUREA
HYDROCET, ACETAMINOPHEN
HYDROCHLOROTHIAZIDE, HYDROCHLOROTHIAZIDE
HYDROCODONE BITARTRATE AND ACETAMINOPHEN, ACETAMINOPHEN
HYDROCODONE BITARTRATE AND HOMATROPINE METHYLBROMIDE, HOMATROPINE METHYLBROMIDE
HYDROCODONE BITARTRATE AND IBUPROFEN, HYDROCODONE BITARTRATE
HYDROCODONE COMPOUND, HOMATROPINE METHYLBROMIDE
HYDROCORTISONE ACETATE 1% AND PRAMOXINE HYDROCHLORIDE 1%, HYDROCORTISONE ACETATE
HYDROCORTISONE ACETATE, HYDROCORTISONE ACETATE
HYDROCORTISONE AND ACETIC ACID, ACETIC ACID, GLACIAL
HYDROCORTISONE BUTYRATE, HYDROCORTISONE BUTYRATE
HYDROCORTISONE IN ABSORBASE, HYDROCORTISONE
HYDROCORTISONE VALERATE, HYDROCORTISONE VALERATE
HYDROCORTISONE, HYDROCORTISONE
HYDROFLUMETHIAZIDE, HYDROFLUMETHIAZIDE
HYDROGENATED ERGOT ALKALOIDS, ERGOLOID MESYLATES
HYDROMORPHONE HYDROCHLORIDE, HYDROMORPHONE HYDROCHLORIDE
HYDRO-RIDE, AMILORIDE HYDROCHLORIDE
HYDRO-RX, HYDROCORTISONE
HYDROXOCOBALAMIN, HYDROXOCOBALAMIN
HYDROXYCHLOROQUINE SULFATE, HYDROXYCHLOROQUINE SULFATE
HYDROXYUREA, HYDROXYUREA
HYDROXYZINE HYDROCHLORIDE, HYDROXYZINE HYDROCHLORIDE
HYDROXYZINE PAMOATE, HYDROXYZINE PAMOATE
HYLENEX RECOMBINANT, HYALURONIDASE RECOMBINANT HUMAN
HYPAQUE, DIATRIZOATE MEGLUMINE
HYPAQUE, DIATRIZOATE SODIUM
HYPAQUE-76, DIATRIZOATE MEGLUMINE
HYPAQUE-CYSTO, DIATRIZOATE MEGLUMINE
HYTONE, HYDROCORTISONE
HYTRIN, TERAZOSIN HYDROCHLORIDE
HYZAAR, HYDROCHLOROTHIAZIDE

**** I ****

IBUPROFEN AND PSEUDOEPHEDRINE HYDROCHLORIDE, IBUPROFEN (OTC)

APPENDIX A - PRODUCT NAME INDEX

A - 26

**** I ****

IBUPROFEN, IBUPROFEN
IBUPROFEN, IBUPROFEN (OTC)
IBUPROHM COLD AND SINUS, IBUPROFEN (OTC)
IBUPROHM, IBUPROFEN
IBUPROHM, IBUPROFEN (OTC)
IBU-TAB 200, IBUPROFEN (OTC)
IBU-TAB, IBUPROFEN
IC-GREEN, INDOCYANINE GREEN
IDAMYCIN PFS, IDARUBICIN HYDROCHLORIDE
IDARUBICIN HYDROCHLORIDE PFS, IDARUBICIN HYDROCHLORIDE
IDARUBICIN HYDROCHLORIDE, IDARUBICIN HYDROCHLORIDE
IFEX/MESNEX KIT, IFOSFAMIDE
IFOSFAMIDE, IFOSFAMIDE
IFOSFAMIDE/MESNA KIT, IFOSFAMIDE
IMDUR, ISOSORBIDE MONONITRATE
IMIPRAMINE HYDROCHLORIDE, IMIPRAMINE HYDROCHLORIDE
IMITREX STATDOSE, SUMATRIPTAN SUCCINATE
IMITREX, SUMATRIPTAN
IMITREX, SUMATRIPTAN SUCCINATE
IMODIUM A-D, LOPERAMIDE HYDROCHLORIDE (OTC)
IMODIUM ADVANCED, LOPERAMIDE HYDROCHLORIDE (OTC)
IMODIUM, LOPERAMIDE HYDROCHLORIDE
IMPLANON, ETONOGESTREL
IMURAN, AZATHIOPRINE
INAPSINE, DROPERIDOL
INCRELEX, MECASERMIN RECOMBINANT
INDAPAMIDE, INDAPAMIDE
INDERAL LA, PROPRANOLOL HYDROCHLORIDE
INDERAL, PROPRANOLOL HYDROCHLORIDE
INDERIDE-40/25, HYDROCHLOROTHIAZIDE
INDICLOR, INDIUM IN-111 CHLORIDE
INDIUM IN 111 CHLORIDE, INDIUM IN-111 CHLORIDE
INDIUM IN-111 OXYQUINOLINE, INDIUM IN-111 OXYQUINOLINE
INDOCIN I.V., INDOMETHACIN SODIUM
INDOCIN SR, INDOMETHACIN
INDOCIN, INDOMETHACIN
INDOMETHACIN, INDOMETHACIN
INFANTS' FEVERALL, ACETAMINOPHEN (OTC)
INFASURF PRESERVATIVE FREE, CALFACTANT
INFED, IRON DEXTRAN
INFLAMASE FORTE, PREDNISOLONE SODIUM PHOSPHATE
INFLAMASE MILD, PREDNISOLONE SODIUM PHOSPHATE
INFUMORPH, MORPHINE SULFATE
INFUVITE ADULT, ALPHA-TOCOPHEROL ACETATE
INFUVITE PEDIATRIC (PHARMACY BULK PACKAGE), ASCORBIC ACID
INFUVITE PEDIATRIC, ASCORBIC ACID
INNOFEM, ESTRADIOL
INNOHEP, TINZAPARIN SODIUM
INNOPRAN XL, PROPRANOLOL HYDROCHLORIDE
INOMAX, NITRIC OXIDE
INPERSOL-LC/LM W/ DEXTROSE 1.5% IN PLASTIC CONTAINER, CALCIUM CHLORIDE
INPERSOL-LC/LM W/ DEXTROSE 2.5% IN PLASTIC CONTAINER, CALCIUM CHLORIDE
INPERSOL-LC/LM W/ DEXTROSE 3.5% IN PLASTIC CONTAINER, CALCIUM CHLORIDE
INPERSOL-LC/LM W/ DEXTROSE 4.25% IN PLASTIC CONTAINER, CALCIUM CHLORIDE
INSPIRA, EPLERENONE
INTAL, CROMOLYN SODIUM
INTEGRILIN, EPTIFIBATIDE
INTRALIPID 10%, SOYBEAN OIL
INTRALIPID 20%, SOYBEAN OIL
INTRALIPID 30%, SOYBEAN OIL
INVAGESIC FORTE, ASPIRIN
INVAGESIC, ASPIRIN
INVANZ, ERTAPENEM SODIUM

APPENDIX A - PRODUCT NAME INDEX

A - 27

**** I ****

INVEGA, PALIPERIDONE
INVERSINE, MECAMYLAMINE HYDROCHLORIDE
INVIRASE, SAQUINAVIR MESYLATE
IOBENGUANE SULFATE I 131, IOBENGUANE SULFATE I 131
IODOTOPE, SODIUM IODIDE, I-131
IONAMIN, PHENTERMINE RESIN COMPLEX
IONOSOL B AND DEXTROSE 5% IN PLASTIC CONTAINER, DEXTROSE
IONOSOL MB AND DEXTROSE 5% IN PLASTIC CONTAINER, DEXTROSE
IONOSOL T AND DEXTROSE 5% IN PLASTIC CONTAINER, DEXTROSE
IONSYS, FENTANYL HYDROCHLORIDE
IOPAMIDOL-200 IN PLASTIC CONTAINER, IOPAMIDOL
IOPAMIDOL-200, IOPAMIDOL
IOPAMIDOL-250 IN PLASTIC CONTAINER, IOPAMIDOL
IOPAMIDOL-250, IOPAMIDOL
IOPAMIDOL-300 IN PLASTIC CONTAINER, IOPAMIDOL
IOPAMIDOL-300, IOPAMIDOL
IOPAMIDOL-370 IN PLASTIC CONTAINER, IOPAMIDOL
IOPAMIDOL-370, IOPAMIDOL
IOPIDINE, APRACLONIDINE HYDROCHLORIDE
IOSAT, POTASSIUM IODIDE (OTC)
IPLEX, MECASERMIN RINFABATE RECOMBINANT
IPRATROPIUM BROMIDE, IPRATROPIUM BROMIDE
IPRIVASK, DESIRUDIN RECOMBINANT
IQUIX, LEVOFLOXACIN
IRESSA, GEFITINIB
ISMO, ISOSORBIDE MONONITRATE
ISOCAINE HYDROCHLORIDE W/ LEVONORDEFIN, LEVONORDEFIN
ISOCAINE HYDROCHLORIDE, MEPIVACAINE HYDROCHLORIDE
ISOFLURANE, ISOFLURANE
ISOLYTE E IN DEXTROSE 5% IN PLASTIC CONTAINER, CALCIUM CHLORIDE
ISOLYTE E IN PLASTIC CONTAINER, CALCIUM CHLORIDE
ISOLYTE H IN DEXTROSE 5% IN PLASTIC CONTAINER, DEXTROSE
ISOLYTE M IN DEXTROSE 5% IN PLASTIC CONTAINER, DEXTROSE
ISOLYTE P IN DEXTROSE 5% IN PLASTIC CONTAINER, DEXTROSE
ISOLYTE R IN DEXTROSE 5% IN PLASTIC CONTAINER, CALCIUM CHLORIDE
ISOLYTE S IN DEXTROSE 5% IN PLASTIC CONTAINER, DEXTROSE
ISOLYTE S IN PLASTIC CONTAINER, MAGNESIUM CHLORIDE
ISOLYTE S PH 7.4 IN PLASTIC CONTAINER, MAGNESIUM CHLORIDE
ISONIAZID, ISONIAZID
ISOPROTERENOL HYDROCHLORIDE, ISOPROTERENOL HYDROCHLORIDE
ISOPTIN SR, VERAPAMIL HYDROCHLORIDE
ISOPTIN, VERAPAMIL HYDROCHLORIDE
ISOPTO CETAMIDE, SULFACETAMIDE SODIUM
ISORDIL, ISOSORBIDE DINITRATE
ISOSORBIDE DINITRATE, ISOSORBIDE DINITRATE
ISOSORBIDE MONONITRATE, ISOSORBIDE MONONITRATE
ISOTONIC GENTAMICIN SULFATE IN PLASTIC CONTAINER, GENTAMICIN SULFATE
ISOVUE-200, IOPAMIDOL
ISOVUE-250, IOPAMIDOL
ISOVUE-300, IOPAMIDOL
ISOVUE-370, IOPAMIDOL
ISOVUE-M 200, IOPAMIDOL
ISOVUE-M 300, IOPAMIDOL
ISRADIPINE, ISRADIPINE
ISTALOL, TIMOLOL MALEATE
ISUPREL, ISOPROTERENOL HYDROCHLORIDE
ITRACONAZOLE, ITRACONAZOLE
IVY BLOCK, BENTOQUATAM (OTC)

**** J ****

JANTOVEN, WARFARIN SODIUM
JANUVIA, SITAGLIPTIN PHOSPHATE
JEANATOPE, ALBUMIN IODINATED I-125 SERUM

APPENDIX A - PRODUCT NAME INDEX

A - 28

**** J ****

JUNEL 1.5/30, ETHINYL ESTRADIOL
JUNEL 1/20, ETHINYL ESTRADIOL
JUNEL FE 1.5/30, ETHINYL ESTRADIOL
JUNEL FE 1/20, ETHINYL ESTRADIOL
JUNIOR STRENGTH ADVIL, IBUPROFEN (OTC)
JUNIOR STRENGTH IBUPROFEN, IBUPROFEN (OTC)
JUNIOR STRENGTH MOTRIN, IBUPROFEN (OTC)

**** K ****

K+10, POTASSIUM CHLORIDE
K+8, POTASSIUM CHLORIDE
KADIAN, MORPHINE SULFATE
KALETRA, LOPINAVIR
KANAMYCIN, KANAMYCIN SULFATE
KANTREX, KANAMYCIN SULFATE
KAON CL-10, POTASSIUM CHLORIDE
KARIVA, DESOGESTREL
KAYEXALATE, SODIUM POLYSTYRENE SULFONATE
K-DUR 10, POTASSIUM CHLORIDE
K-DUR 20, POTASSIUM CHLORIDE
KEFLEX, CEPHALEXIN
KELNOR, ETHINYL ESTRADIOL
KEMADRIN, PROCYCLIDINE HYDROCHLORIDE
KEMSTRO, BACLOFEN
KENALOG IN ORABASE, TRIAMCINOLONE ACETONIDE
KENALOG, TRIAMCINOLONE ACETONIDE
KENALOG-10, TRIAMCINOLONE ACETONIDE
KENALOG-40, TRIAMCINOLONE ACETONIDE
KEPPRA, LEVETIRACETAM
KERLONE, BETAXOLOL HYDROCHLORIDE
KETALAR, KETAMINE HYDROCHLORIDE
KETAMINE HYDROCHLORIDE, KETAMINE HYDROCHLORIDE
KETEK, TELITHROMYCIN
KETOCONAZOLE, KETOCONAZOLE
KETOPROFEN, KETOPROFEN
KETOROLAC TROMETHAMINE, KETOROLAC TROMETHAMINE
KETOTIFEN FUMARATE, KETOTIFEN FUMARATE
KETOZOLE, KETOCONAZOLE
KINEVAC, SINCALIDE
KIONEX, SODIUM POLYSTYRENE SULFONATE
KLARON, SULFACETAMIDE SODIUM
KLONOPIN RAPIDLY DISINTEGRATING, CLONAZEPAM
KLONOPIN, CLONAZEPAM
KLOR-CON M10, POTASSIUM CHLORIDE
KLOR-CON M15, POTASSIUM CHLORIDE
KLOR-CON M20, POTASSIUM CHLORIDE
KLOR-CON, POTASSIUM CHLORIDE
KLOTRIX, POTASSIUM CHLORIDE
K-TAB, POTASSIUM CHLORIDE
KYTRIL, GRANISETRON HYDROCHLORIDE

**** L ****

LABETALOL HYDROCHLORIDE, LABETALOL HYDROCHLORIDE
LAC-HYDRIN, AMMONIUM LACTATE
LACRISERT, HYDROXYPROPYL CELLULOSE
LACTATED RINGER'S AND DEXTROSE 5% IN PLASTIC CONTAINER, CALCIUM CHLORIDE
LACTATED RINGER'S IN PLASTIC CONTAINER, CALCIUM CHLORIDE
LACTULOSE, LACTULOSE
LAMICTAL CD, LAMOTRIGINE
LAMICTAL, LAMOTRIGINE
LAMISIL AT, TERBINAFINE (OTC)
LAMISIL AT, TERBINAFINE HYDROCHLORIDE (OTC)

APPENDIX A - PRODUCT NAME INDEX

A - 29

**** L ****

LAMISIL, TERBINAFINE
LAMISIL, TERBINAFINE HYDROCHLORIDE
LAMISIL, TERBINAFINE HYDROCHLORIDE (OTC)
LAMOTRIGINE, LAMOTRIGINE
LAMPRENE, CLOFAZIMINE
LANIAZID, ISONIAZID
LANORINAL, ASPIRIN
LANOXICAPS, DIGOXIN
LANOXIN PEDIATRIC, DIGOXIN
LANOXIN, DIGOXIN
LANTUS, INSULIN GLARGINE RECOMBINANT
LARIAM, MEFLOROQUINE HYDROCHLORIDE
LAROTID, AMOXICILLIN
LARYNG-O-JET KIT, LIDOCAINE HYDROCHLORIDE
LASIX, FUROSEMIDE
LAXILOSE, LACTULOSE
LEFLUNOMIDE, LEFLUNOMIDE
LESCOL XL, FLUVASTATIN SODIUM
LESCOL, FLUVASTATIN SODIUM
LESSINA-28, ETHINYL ESTRADIOL
LEUCOVORIN CALCIUM PRESERVATIVE FREE, LEUCOVORIN CALCIUM
LEUCOVORIN CALCIUM, LEUCOVORIN CALCIUM
LEUKERAN, CHLORAMBUCIL
LEUPROLIDE ACETATE, LEUPROLIDE ACETATE
LEUSTATIN, CLADRIBINE
LEVAQUIN IN DEXTROSE 5% IN PLASTIC CONTAINER, LEVOFLOXACIN
LEVAQUIN, LEVOFLOXACIN
LEVATOL, PENBUTOLOL SULFATE
LEVEMIR, INSULIN DETEMIR RECOMBINANT
LEVITRA, VARDENAFIL HYDROCHLORIDE
LEVLITE, ETHINYL ESTRADIOL
LEVOBUNOLOL HYDROCHLORIDE, LEVOBUNOLOL HYDROCHLORIDE
LEVOCARNITINE, LEVOCARNITINE
LEVO-DROMORAN, LEVORPHANOL TARTRATE
LEVOLET, LEVOTHYROXINE SODIUM
LEVONORGESTREL AND ETHINYL ESTRADIOL, ETHINYL ESTRADIOL
LEVOPHED, NOREPINEPHRINE BITARTRATE
LEVORA 0.15/30-28, ETHINYL ESTRADIOL
LEVORPHANOL TARTRATE, LEVORPHANOL TARTRATE
LEVO-T, LEVOTHYROXINE SODIUM
LEVOTHROID, LEVOTHYROXINE SODIUM
LEVOTHYROXINE SODIUM, LEVOTHYROXINE SODIUM
LEVOXYL, LEVOTHYROXINE SODIUM
LEVULAN, AMINOLEVULINIC ACID HYDROCHLORIDE
LEXAPRO, ESCITALOPRAM OXALATE
LEXIVA, FOSAMPRENAVIR CALCIUM
LEXXEL, ENALAPRIL MALEATE
LIBRIUM, CHLORDIAZEPOXIDE HYDROCHLORIDE
LIDEX, FLUOCINONIDE
LIDEX-E, FLUOCINONIDE
LIDOCAINE AND PRILOCAINE, LIDOCAINE
LIDOCAINE AND TETRACAINE, LIDOCAINE
LIDOCAINE HYDROCHLORIDE 0.2% AND DEXTROSE 5% IN PLASTIC CONTAINER, LIDOCAINE HYDROCHLORIDE
LIDOCAINE HYDROCHLORIDE 0.2% IN DEXTROSE 5% IN PLASTIC CONTAINER, LIDOCAINE HYDROCHLORIDE
LIDOCAINE HYDROCHLORIDE 0.2% IN DEXTROSE 5%, LIDOCAINE HYDROCHLORIDE
LIDOCAINE HYDROCHLORIDE 0.4% AND DEXTROSE 5% IN PLASTIC CONTAINER, LIDOCAINE HYDROCHLORIDE
LIDOCAINE HYDROCHLORIDE 0.4% IN DEXTROSE 5% IN PLASTIC CONTAINER, LIDOCAINE HYDROCHLORIDE
LIDOCAINE HYDROCHLORIDE 0.4% IN DEXTROSE 5%, LIDOCAINE HYDROCHLORIDE
LIDOCAINE HYDROCHLORIDE 0.8% AND DEXTROSE 5% IN PLASTIC CONTAINER, LIDOCAINE HYDROCHLORIDE
LIDOCAINE HYDROCHLORIDE 0.8% IN DEXTROSE 5% IN PLASTIC CONTAINER, LIDOCAINE HYDROCHLORIDE
LIDOCAINE HYDROCHLORIDE 5% AND DEXTROSE 7.5%, LIDOCAINE HYDROCHLORIDE
LIDOCAINE HYDROCHLORIDE AND EPINEPHRINE, EPINEPHRINE
LIDOCAINE HYDROCHLORIDE IN PLASTIC CONTAINER, LIDOCAINE HYDROCHLORIDE

APPENDIX A - PRODUCT NAME INDEX

A - 30

**** L ****

LIDOCAINE HYDROCHLORIDE PRESERVATIVE FREE IN PLASTIC CONTAINER, LIDOCAINE HYDROCHLORIDE
LIDOCAINE HYDROCHLORIDE PRESERVATIVE FREE, LIDOCAINE HYDROCHLORIDE
LIDOCAINE HYDROCHLORIDE VISCOUS, LIDOCAINE HYDROCHLORIDE
LIDOCAINE HYDROCHLORIDE W/ EPINEPHRINE, EPINEPHRINE
LIDOCAINE HYDROCHLORIDE, LIDOCAINE HYDROCHLORIDE
LIDOCAINE VISCOUS, LIDOCAINE HYDROCHLORIDE
LIDOCAINE, LIDOCAINE
LIDOCATON, EPINEPHRINE
LIDODERM, LIDOCAINE
LIDOPEN, LIDOCAINE HYDROCHLORIDE
LIDOSITE TOPICAL SYSTEM KIT, EPINEPHRINE
LIGNOSPAN FORTE, EPINEPHRINE BITARTRATE
LIGNOSPAN STANDARD, EPINEPHRINE BITARTRATE
LIMBITROL DS, AMITRIPTYLINE HYDROCHLORIDE
LIMBITROL, AMITRIPTYLINE HYDROCHLORIDE
LINCOCIN, LINCOMYCIN HYDROCHLORIDE
LINDANE, LINDANE
LIORESAL, BACLOFEN
LIOTHYRONINE SODIUM, LIOTHYRONINE SODIUM
LIPITOR, ATORVASTATIN CALCIUM
LIPOFEN, FENOFIBRATE
LIPOSYN II 10%, SAFFLOWER OIL
LIPOSYN II 20%, SAFFLOWER OIL
LIPOSYN III 10%, SOYBEAN OIL
LIPOSYN III 20%, SOYBEAN OIL
LIPOSYN III 30%, SOYBEAN OIL
LISINOPRIL AND HYDROCHLOROTHIAZIDE, HYDROCHLOROTHIAZIDE
LISINOPRIL, LISINOPRIL
LITHIUM CARBONATE, LITHIUM CARBONATE
LITHIUM CITRATE, LITHIUM CITRATE
LITHOBID, LITHIUM CARBONATE
LITHOSTAT, ACETOHYDROXAMIC ACID
LO/OVRAL-28, ETHINYL ESTRADIOL
LOCHOLEST LIGHT, CHOLESTYRAMINE
LOCHOLEST, CHOLESTYRAMINE
LOCOID LIPOCREAM, HYDROCORTISONE BUTYRATE
LOCOID, HYDROCORTISONE BUTYRATE
LODOSYN, CARBIDOPA
LOESTRIN 21 1.5/30, ETHINYL ESTRADIOL
LOESTRIN 21 1/20, ETHINYL ESTRADIOL
LOESTRIN 24 FE, ETHINYL ESTRADIOL
LOESTRIN FE 1.5/30, ETHINYL ESTRADIOL
LOESTRIN FE 1/20, ETHINYL ESTRADIOL
LOMOTIL, ATROPINE SULFATE
LONITEN, MINOXIDIL
LONOX, ATROPINE SULFATE
LOPERAMIDE HYDROCHLORIDE AND SIMETHICONE, LOPERAMIDE HYDROCHLORIDE (OTC)
LOPERAMIDE HYDROCHLORIDE, LOPERAMIDE HYDROCHLORIDE
LOPERAMIDE HYDROCHLORIDE, LOPERAMIDE HYDROCHLORIDE (OTC)
LOPID, GEMFIBROZIL
LOPRESSOR HCT, HYDROCHLOROTHIAZIDE
LOPRESSOR, METOPROLOL TARTRATE
LOPROX, CICLOPIROX
LOPURIN, ALLOPURINOL
LORATADINE AND PSEUDOEPHEDRINE SULFATE, LORATADINE (OTC)
LORATADINE, LORATADINE (OTC)
LORAZEPAM INTENSOL, LORAZEPAM
LORAZEPAM PRESERVATIVE FREE, LORAZEPAM
LORAZEPAM, LORAZEPAM
LORCET-HD, ACETAMINOPHEN
LORTAB, ACETAMINOPHEN
LOTEMAX, LOTEPREDNOL ETABONATE
LOTENSIN HCT, BENAZEPRIL HYDROCHLORIDE

APPENDIX A - PRODUCT NAME INDEX

A - 31

**** L ****

LOTENSIN, BENAZEPRIL HYDROCHLORIDE
LOTREL, AMLODIPINE BESYLATE
LOTRIMIN ULTRA, BUTENAFINE HYDROCHLORIDE (OTC)
LOTRISONE, BETAMETHASONE DIPROPIONATE
LOTRONEX, ALOSETRON HYDROCHLORIDE
LOVASTATIN, LOVASTATIN
LOVENOX (PRESERVATIVE FREE), ENOXAPARIN SODIUM
LOVENOX, ENOXAPARIN SODIUM
LOW-OGESTREL-21, ETHINYL ESTRADIOL
LOW-OGESTREL-28, ETHINYL ESTRADIOL
LOXAPINE SUCCINATE, LOXAPINE SUCCINATE
LOXAPINE, LOXAPINE SUCCINATE
LOXITANE, LOXAPINE SUCCINATE
LOZOL, INDAPAMIDE
LTA II KIT, LIDOCAINE HYDROCHLORIDE
LUFYLLIN, DYPHYLLINE
LUMIGAN, BIMATOPROST
LUNESTA, ESZOPICLONE
LUPRON DEPOT, LEUPROLIDE ACETATE
LUPRON DEPOT-3, LEUPROLIDE ACETATE
LUPRON DEPOT-4, LEUPROLIDE ACETATE
LUPRON DEPOT-PED, LEUPROLIDE ACETATE
LUPRON, LEUPROLIDE ACETATE
LUVERIS, LUTROPIN ALFA
LUXIQ, BETAMETHASONE VALERATE
LYMPHAZURIN, ISOSULFAN BLUE
LYOPHILIZED CYTOXAN, CYCLOPHOSPHAMIDE
LYRICA, PREGABALIN
LYSODREN, MITOTANE

**** M ****

M.V.I. ADULT (PHARMACY BULK PACKAGE), ASCORBIC ACID
M.V.I. ADULT, ASCORBIC ACID
M.V.I. PEDIATRIC, ASCORBIC ACID
M.V.I.-12 (WITHOUT VITAMIN K), ASCORBIC ACID
MACROBID, NITROFURANTOIN
MACRODANTIN, NITROFURANTOIN, MACROCRYSTALLINE
MACROTEC, TECHNETIUM TC-99M ALBUMIN AGGREGATED KIT
MACUGEN, PEGAPTANIB SODIUM
MAGNESIUM SULFATE IN DEXTROSE 5% IN PLASTIC CONTAINER, MAGNESIUM SULFATE
MAGNESIUM SULFATE IN PLASTIC CONTAINER, MAGNESIUM SULFATE
MAGNESIUM SULFATE, MAGNESIUM SULFATE
MAGNEVIST, GADOPENTETATE DIMEGLUMINE
MALARONE PEDIATRIC, ATOVAQUONE
MALARONE, ATOVAQUONE
MANGANESE CHLORIDE IN PLASTIC CONTAINER, MANGANESE CHLORIDE
MANNITOL 10% IN PLASTIC CONTAINER, MANNITOL
MANNITOL 10% W/ DEXTROSE 5% IN DISTILLED WATER, MANNITOL
MANNITOL 10%, MANNITOL
MANNITOL 15% IN PLASTIC CONTAINER, MANNITOL
MANNITOL 15% W/ DEXTROSE 5% IN SODIUM CHLORIDE 0.45%, MANNITOL
MANNITOL 15%, MANNITOL
MANNITOL 20% IN PLASTIC CONTAINER, MANNITOL
MANNITOL 20%, MANNITOL
MANNITOL 25%, MANNITOL
MANNITOL 5% IN PLASTIC CONTAINER, MANNITOL
MANNITOL 5% W/ DEXTROSE 5% IN SODIUM CHLORIDE 0.12%, MANNITOL
MANNITOL 5%, MANNITOL
MAPROTILINE HYDROCHLORIDE, MAPROTILINE HYDROCHLORIDE
MARCAINE HYDROCHLORIDE PRESERVATIVE FREE, BUPIVACAINE HYDROCHLORIDE
MARCAINE HYDROCHLORIDE W/ EPINEPHRINE PRESERVATIVE FREE, BUPIVACAINE HYDROCHLORIDE
MARCAINE HYDROCHLORIDE W/ EPINEPHRINE, BUPIVACAINE HYDROCHLORIDE
MARCAINE HYDROCHLORIDE, BUPIVACAINE HYDROCHLORIDE

APPENDIX A - PRODUCT NAME INDEX

A - 32

**** M ****

MARCAINE, BUPIVACAINE HYDROCHLORIDE
MARINOL, DRONABINOL
MARPLAN, ISOCARBOXAZID
MATULANE, PROCARBAZINE HYDROCHLORIDE
MAVIK, TRANDOLAPRIL
MAXAIR, PIRBUTEROL ACETATE
MAXALT, RIZATRIPTAN BENZOATE
MAXALT-MLT, RIZATRIPTAN BENZOATE
MAXAQUIN, LOMEFLOXACIN HYDROCHLORIDE
MAXIDEX, DEXAMETHASONE
MAXIPIME, CEFEPIME HYDROCHLORIDE (ARGININE FORMULATION)
MAXITROL, DEXAMETHASONE
MAXZIDE, HYDROCHLOROTHIAZIDE
MAXZIDE-25, HYDROCHLOROTHIAZIDE
MD-76R, DIATRIZOATE MEGLUMINE
MD-GASTROVIEW, DIATRIZOATE MEGLUMINE
MDP-BRACCO, TECHNETIUM TC-99M MEDRONATE KIT
MEBENDAZOLE, MEBENDAZOLE
MECLIZINE HYDROCHLORIDE, MECLIZINE HYDROCHLORIDE
MECLOFENAMATE SODIUM, MECLOFENAMATE SODIUM
MEDIPREN, IBUPROFEN (OTC)
MEDROL, METHYLPREDNISOLONE
MEDROXYPROGESTERONE ACETATE, MEDROXYPROGESTERONE ACETATE
MEFLOQUINE HYDROCHLORIDE, MEFLOQUINE HYDROCHLORIDE
MEFLOQUINE, MEFLOQUINE HYDROCHLORIDE
MEFOXIN IN PLASTIC CONTAINER, CEFOXITIN SODIUM
MEFOXIN, CEFOXITIN SODIUM
MEGACE ES, MEGESTROL ACETATE
MEGACE, MEGESTROL ACETATE
MEGATOPE, ALBUMIN IODINATED I-131 SERUM
MEGESTROL ACETATE, MEGESTROL ACETATE
MELOXICAM, MELOXICAM
MENEST, ESTROGENS, ESTERIFIED
MENOPUR, LUTEINIZING HORMONE
MENOSTAR, ESTRADIOL
MEN'S ROGAINE, MINOXIDIL (OTC)
MENTAX, BUTENAFINE HYDROCHLORIDE
MENTAX-TC, BUTENAFINE HYDROCHLORIDE
MEPERIDINE HYDROCHLORIDE PRESERVATIVE FREE, MEPERIDINE HYDROCHLORIDE
MEPERIDINE HYDROCHLORIDE, MEPERIDINE HYDROCHLORIDE
MEPHYTON, PHYTONADIONE
MEPIVACAINE HYDROCHLORIDE W/ LEVONORDEFIN, LEVONORDEFIN
MEPIVACAINE HYDROCHLORIDE, MEPIVACAINE HYDROCHLORIDE
MEPROBAMATE, MEPROBAMATE
MEPRON, ATOVAQUONE
MERCAPTOPYRINE, MERCAPTOPYRINE
MERCAPTOPYRINE, MERCAPTOPYRINE ANHYDROUS
MERIDIA, SIBUTRAMINE HYDROCHLORIDE
MERREM I.V., MEROPENEM
MESALAMINE, MESALAMINE
MESNA, MESNA
MESNEX, MESNA
MESTINON, PYRIDOSTIGMINE BROMIDE
METADATE CD, METHYLPHENIDATE HYDROCHLORIDE
METADATE ER, METHYLPHENIDATE HYDROCHLORIDE
METAGLIP, GLIPIZIDE
METAPROTERENOL SULFATE, METAPROTERENOL SULFATE
METARAMINOL BITARTRATE, METARAMINOL BITARTRATE
METASTRON, STRONTIUM CHLORIDE, SR-89
METFORMIN HYDROCHLORIDE, METFORMIN HYDROCHLORIDE
METHADONE HYDROCHLORIDE INTENSOL, METHADONE HYDROCHLORIDE
METHADONE HYDROCHLORIDE, METHADONE HYDROCHLORIDE
METHADOSE, METHADONE HYDROCHLORIDE

APPENDIX A - PRODUCT NAME INDEX

A - 33

**** M ****

METHAZOLAMIDE, METHAZOLAMIDE
METHENAMINE HIPPURATE, METHENAMINE HIPPURATE
METHERGINE, METHYLERGONOVINE MALEATE
METHIMAZOLE, METHIMAZOLE
METHOCARBAMOL AND ASPIRIN, ASPIRIN
METHOCARBAMOL, METHOCARBAMOL
METHOTREXATE PRESERVATIVE FREE, METHOTREXATE SODIUM
METHOTREXATE SODIUM, METHOTREXATE SODIUM
METHOTREXATE, METHOTREXATE SODIUM
METHSCOPOLAMINE BROMIDE, METHSCOPOLAMINE BROMIDE
METHYCLOTHIAZIDE, METHYCLOTHIAZIDE
METHYLDOPA AND HYDROCHLOROTHIAZIDE, HYDROCHLOROTHIAZIDE
METHYLDOPA, METHYLDOPA
METHYLDOPATE HYDROCHLORIDE, METHYLDOPATE HYDROCHLORIDE
METHYLIN ER, METHYLPHENIDATE HYDROCHLORIDE
METHYLIN, METHYLPHENIDATE HYDROCHLORIDE
METHYLPHENIDATE HYDROCHLORIDE, METHYLPHENIDATE HYDROCHLORIDE
METHYLPREDNISOLONE ACETATE, METHYLPREDNISOLONE ACETATE
METHYLPREDNISOLONE SODIUM SUCCINATE, METHYLPREDNISOLONE SODIUM SUCCINATE
METHYLPREDNISOLONE, METHYLPREDNISOLONE
METHYLTESTOSTERONE, METHYLTESTOSTERONE
METIPRANOLOL, METIPRANOLOL HYDROCHLORIDE
METOCLOPRAMIDE HYDROCHLORIDE, METOCLOPRAMIDE HYDROCHLORIDE
METOCLOPRAMIDE, METOCLOPRAMIDE HYDROCHLORIDE
METOLAZONE, METOLAZONE
METOPIRONE, METYRAPONE
METOPROLOL SUCCINATE, METOPROLOL SUCCINATE
METOPROLOL TARTRATE AND HYDROCHLOROTHIAZIDE, HYDROCHLOROTHIAZIDE
METOPROLOL TARTRATE, METOPROLOL TARTRATE
METRO I.V. IN PLASTIC CONTAINER, METRONIDAZOLE
METROCREAM, METRONIDAZOLE
METROGEL, METRONIDAZOLE
METROGEL-VAGINAL, METRONIDAZOLE
METROLOTION, METRONIDAZOLE
METRONIDAZOLE IN PLASTIC CONTAINER, METRONIDAZOLE
METRONIDAZOLE, METRONIDAZOLE
METVIXIA, METHYL AMINOLEVULINATE HYDROCHLORIDE
MEVACOR, LOVASTATIN
MEXILETINE HYDROCHLORIDE, MEXILETINE HYDROCHLORIDE
MEXITIL, MEXILETINE HYDROCHLORIDE
MIACALCIN, CALCITONIN, SALMON
MICARDIS HCT, HYDROCHLOROTHIAZIDE
MICARDIS, TELMISARTAN
MICONAZOLE 3 COMBINATION PACK, MICONAZOLE NITRATE (OTC)
MICONAZOLE 3, MICONAZOLE NITRATE (OTC)
MICONAZOLE 7 COMBINATION PACK, MICONAZOLE NITRATE (OTC)
MICONAZOLE 7, MICONAZOLE NITRATE (OTC)
MICONAZOLE NITRATE COMBINATION PACK, MICONAZOLE NITRATE (OTC)
MICONAZOLE NITRATE, MICONAZOLE NITRATE
MICONAZOLE NITRATE, MICONAZOLE NITRATE (OTC)
MICORT-HC, HYDROCORTISONE ACETATE
MICRODERM, CHLORHEXIDINE GLUCONATE (OTC)
MICROGESTIN FE 1.5/30, ETHINYL ESTRADIOL
MICROGESTIN FE 1/20, ETHINYL ESTRADIOL
MICRO-K 10, POTASSIUM CHLORIDE
MICRO-K, POTASSIUM CHLORIDE
MICRONASE, GLYBURIDE
MICRONOR, NORETHINDRONE
MICROZIDE, HYDROCHLOROTHIAZIDE
MIDAZOLAM HYDROCHLORIDE, MIDAZOLAM HYDROCHLORIDE
MIDODRINE HYDROCHLORIDE, MIDODRINE HYDROCHLORIDE
MIFEPREX, MIFEPRISTONE
MIGERGOT, CAFFEINE

APPENDIX A - PRODUCT NAME INDEX

A - 34

**** M ****

MIGRANAL, DIHYDROERGOTAMINE MESYLATE
MILOPHENE, CLOMIPHENE CITRATE
MILRINONE LACTATE IN 5% DEXTROSE IN PLASTIC CONTAINER, MILRINONE LACTATE
MILRINONE LACTATE IN DEXTROSE 5% IN PLASTIC CONTAINER, MILRINONE LACTATE
MILRINONE LACTATE, MILRINONE LACTATE
MILRINONE LACTOSE IN 5% DEXTROSE, MILRINONE LACTATE
MINIPRESS XL, PRAZOSIN HYDROCHLORIDE
MINIPRESS, PRAZOSIN HYDROCHLORIDE
MINIRIN, DESMOPRESSIN ACETATE
MINITRAN, NITROGLYCERIN
MINIZIDE, POLYTHIAZIDE
MINOCIN, MINOCYCLINE HYDROCHLORIDE
MINOCYCLINE HYDROCHLORIDE, MINOCYCLINE HYDROCHLORIDE
MINOXIDIL (FOR MEN), MINOXIDIL (OTC)
MINOXIDIL (FOR WOMEN), MINOXIDIL (OTC)
MINOXIDIL EXTRA STRENGTH (FOR MEN), MINOXIDIL (OTC)
MINOXIDIL, MINOXIDIL
MINTEZOL, THIABENDAZOLE
MIOCHOL-E, ACETYLCHOLINE CHLORIDE
MIOSSTAT, CARBACHOL
MIRALAX, POLYETHYLENE GLYCOL 3350 (OTC)
MIRALUMA, TECHNETIUM TC-99M SESTAMIBI KIT
MIRAPEX, PRAMIPEXOLE DIHYDROCHLORIDE
MIRCETTE, DESOGESTREL
MIRENA, LEVONORGESTREL
MIRTAZAPINE, MIRTAZAPINE
MISOPROSTOL, MISOPROSTOL
MITHRACIN, PLICAMYCIN
MITOMYCIN, MITOMYCIN
MITOXANTRONE, MITOXANTRONE HYDROCHLORIDE
MOBAN, MOLINDONE HYDROCHLORIDE
MOBIC, MELOXICAM
MODERIL, RESCINNAMINE
MODICON 28, ETHINYL ESTRADIOL
MOEXIPRIL HYDROCHLORIDE, MOEXIPRIL HYDROCHLORIDE
MOMETASONE FUROATE, MOMETASONE FUROATE
MONISTAT 1 COMBINATION PACK, MICONAZOLE NITRATE (OTC)
MONISTAT 3 COMBINATION PACK (PREFILLED), MICONAZOLE NITRATE (OTC)
MONISTAT 3 COMBINATION PACK, MICONAZOLE NITRATE (OTC)
MONISTAT 3, MICONAZOLE NITRATE
MONISTAT 3, MICONAZOLE NITRATE (OTC)
MONISTAT 7 COMBINATION PACK, MICONAZOLE NITRATE (OTC)
MONISTAT 7, MICONAZOLE NITRATE (OTC)
MONISTAT-3 COMBINATION PACK, MICONAZOLE NITRATE (OTC)
MONISTAT-DERM, MICONAZOLE NITRATE
MONODOX, DOXYCYCLINE
MONOKET, ISOSORBIDE MONONITRATE
MONOPRIL, FOSINOPRIL SODIUM
MONOPRIL-HCT, FOSINOPRIL SODIUM
MONUROL, FOSFOMYCIN TROMETHAMINE
MORPHINE SULFATE, MORPHINE SULFATE
MOTOFEN, ATROPINE SULFATE
MOTRIN IB, IBUPROFEN (OTC)
MOTRIN MIGRAINE PAIN, IBUPROFEN (OTC)
MOTRIN, IBUPROFEN
MOVIPREP, ASCORBIC ACID
MPI DMSA KIDNEY REAGENT, TECHNETIUM TC-99M SUCCIMER KIT
MPI DTPA KIT - CHELATE, TECHNETIUM TC-99M PENTETATE KIT
MPI INDIUM DTPA IN 111, INDIUM IN-111 PENTETATE DISODIUM
MS CONTIN, MORPHINE SULFATE
MUCINEX D, GUAIFENESIN (OTC)
MUCINEX DM, DEXTROMETHORPHAN HYDROBROMIDE (OTC)
MUCINEX, GUAIFENESIN (OTC)

APPENDIX A - PRODUCT NAME INDEX

A - 35

**** M ****

MUCOMYST, ACETYLCYSTEINE
MULTIHANCE MULTIPACK, GADOBENATE DIMEGLUMINE
MULTIHANCE, GADOBENATE DIMEGLUMINE
MUPIROCIN, MUPIROCIN
MUSE, ALPROSTADIL
MUSTARGEN, MECHLORETHAMINE HYDROCHLORIDE
MUTAMYCIN, MITOMYCIN
MYAMBUTOL, ETHAMBUTOL HYDROCHLORIDE
MYBANIL, BROMODIPHENHYDRAMINE HYDROCHLORIDE
MYCAMINE, MICA FUNGIN SODIUM
MYCELEX, CLOTRIMAZOLE
MYCELEX-7 COMBINATION PACK, CLOTRIMAZOLE (OTC)
MYCELEX-7, CLOTRIMAZOLE (OTC)
MYCOBUTIN, RIFABUTIN
MYCODONE, HOMATROPINE METHYLBROMIDE
MYCOSTATIN, NYSTATIN
MYDRIACYL, TROPICAMIDE
MYFORTIC, MYCOPHENOLIC ACID
MYKACET, NYSTATIN
MYLARAMINE, DEXCHLORPHENIRAMINE MALEATE
MYLERAN, BUSULFAN
MYLOTARG, GEMTUZUMAB OZOGAMICIN
MYMETHASONE, DEXAMETHASONE
MYOVIEW, TECHNETIUM TC-99M TETROFOSMIN KIT
MYPHETANE DX, BROMPHENIRAMINE MALEATE
MYPROIC ACID, VALPROIC ACID
MYSOLINE, PRIMIDONE
MYTELASE, AMBENONIUM CHLORIDE
MYTOZYTREX, MITOMYCIN
M-ZOLE 3 COMBINATION PACK, MICONAZOLE NITRATE (OTC)
M-ZOLE 7 DUAL PACK, MICONAZOLE NITRATE (OTC)

**** N ****

NABUMETONE, NABUMETONE
NADOLOL, NADOLOL
NAFAZAIR, NAPHAZOLINE HYDROCHLORIDE
NAFCILLIN SODIUM, NAFCILLIN SODIUM
NAFTIN, NAFTIFINE HYDROCHLORIDE
NALBUPHINE HYDROCHLORIDE, NALBUPHINE HYDROCHLORIDE
NALFON 200, FENOPROFEN CALCIUM
NALFON, FENOPROFEN CALCIUM
NALIDIXIC ACID, NALIDIXIC ACID
NALLPEN IN PLASTIC CONTAINER, NAFCILLIN SODIUM
NALOXONE HYDROCHLORIDE AND PENTAZOCAINE, NALOXONE HYDROCHLORIDE
NALOXONE HYDROCHLORIDE, NALOXONE HYDROCHLORIDE
NALTREXONE HYDROCHLORIDE, NALTREXONE HYDROCHLORIDE
NAMENDA, MEMANTINE HYDROCHLORIDE
NANDROLONE DECANOATE, NANDROLONE DECANOATE
NAPHAZOLINE HYDROCHLORIDE, NAPHAZOLINE HYDROCHLORIDE
NAPHCON-A, NAPHAZOLINE HYDROCHLORIDE (OTC)
NAPRELAN, NAPROXEN SODIUM
NAPROSYN, NAPROXEN
NAPROXEN SODIUM AND PSEUDOEPHEDRINE HYDROCHLORIDE, NAPROXEN SODIUM (OTC)
NAPROXEN SODIUM, NAPROXEN SODIUM
NAPROXEN SODIUM, NAPROXEN SODIUM (OTC)
NAPROXEN, NAPROXEN
NARCAN, NALOXONE HYDROCHLORIDE
NARDIL, PHENELZINE SULFATE
NAROPIN, ROPIVACAINE HYDROCHLORIDE MONOHYDRATE
NASACORT AQ, TRIAMCINOLONE ACETONIDE
NASAREL, FLUNISOLIDE
NASCOBAL, CYANOCOBALAMIN
NASONEX, MOMETASONE FUROATE MONOHYDRATE

APPENDIX A - PRODUCT NAME INDEX

A - 36

**** N ****

NATACYN, NATAMYCIN
NATRECOR, NESIRITIDE RECOMBINANT
NAVANE, THIOTHIXENE
NAVANE, THIOTHIXENE HYDROCHLORIDE
NAVELBINE, VINORELBINE TARTRATE
NEBUPENT, PENTAMIDINE ISETHIONATE
NEFAZODONE HYDROCHLORIDE, NEFAZODONE HYDROCHLORIDE
NEGGAM, NALIDIXIC ACID
NEMBUTAL SODIUM, PENTOBARBITAL SODIUM
NEO-FRADIN, NEOMYCIN SULFATE
NEOMYCIN AND POLYMYXIN B SULFATE AND BACITRACIN ZINC, BACITRACIN ZINC
NEOMYCIN AND POLYMYXIN B SULFATE, NEOMYCIN SULFATE
NEOMYCIN AND POLYMYXIN B SULFATES AND BACITRACIN ZINC, BACITRACIN ZINC
NEOMYCIN AND POLYMYXIN B SULFATES AND DEXAMETHASONE, DEXAMETHASONE
NEOMYCIN AND POLYMYXIN B SULFATES AND GRAMICIDIN, GRAMICIDIN
NEOMYCIN AND POLYMYXIN B SULFATES AND HYDROCORTISONE, HYDROCORTISONE
NEOMYCIN AND POLYMYXIN B SULFATES, BACITRACIN ZINC AND HYDROCORTISONE, BACITRACIN ZINC
NEOMYCIN SULFATE, NEOMYCIN SULFATE
NEOPAP, ACETAMINOPHEN (OTC)
NEOPROFEN, IBUPROFEN LYSINE
NEORAL, CYCLOSPORINE
NEO-RX, NEOMYCIN SULFATE
NEOSAR, CYCLOPHOSPHAMIDE
NEOSPORIN G.U. IRRIGANT, NEOMYCIN SULFATE
NEOSPORIN, BACITRACIN ZINC
NEOSPORIN, GRAMICIDIN
NEPHRAMINE 5.4%, AMINO ACIDS
NESACAINE, CHLOROPROCAINE HYDROCHLORIDE
NESACAINE-MPF, CHLOROPROCAINE HYDROCHLORIDE
NEUROLITE, TECHNETIUM TC-99M BICISATE KIT
NEURONTIN, GABAPENTIN
NEUTREXIN, TRIMETREXATE GLUCURONATE
NEVANAC, NEPAFENAC
NEXAVAR, SORAFENIB TOSYLATE
NEXIUM IV, ESOMEPRAZOLE SODIUM
NEXIUM, ESOMEPRAZOLE MAGNESIUM
NIACIN, NIACIN
NIACOR, NIACIN
NIASPAN, NIACIN
NICARDIPINE HYDROCHLORIDE, NICARDIPINE HYDROCHLORIDE
NICODERM CQ, NICOTINE (OTC)
NICORETTE (MINT), NICOTINE POLACRILEX (OTC)
NICORETTE, NICOTINE POLACRILEX (OTC)
NICOTINE POLACRILEX, NICOTINE POLACRILEX (OTC)
NICOTINE, NICOTINE
NICOTROL, NICOTINE
NIFEDIPINE, NIFEDIPINE
NILANDRON, NILUTAMIDE
NILSTAT, NYSTATIN
NIMBEX PRESERVATIVE FREE, CISATRACURIUM BESYLATE
NIMBEX, CISATRACURIUM BESYLATE
NIMOTOP, NIMODIPINE
NIPENT, PENTOSTATIN
NIRAVAM, ALPRAZOLAM
NITRO-DUR, NITROGLYCERIN
NITROFURANTOIN (MONOHYDRATE/MACROCRYSTALS), NITROFURANTOIN
NITROFURANTOIN, NITROFURANTOIN, MACROCRYSTALLINE
NITROFURAZONE, NITROFURAZONE
NITROGLYCERIN IN DEXTROSE 5%, NITROGLYCERIN
NITROGLYCERIN, NITROGLYCERIN
NITROLINGUAL PUMPSPRAY, NITROGLYCERIN
NITROMIST, NITROGLYCERIN
NITROPRESS, SODIUM NITROPRUSSIDE

APPENDIX A - PRODUCT NAME INDEX

A - 37

**** N ****

NITROSTAT, NITROGLYCERIN
NIX, PERMETHRIN (OTC)
NIZATIDINE, NIZATIDINE
NIZORAL A-D, KETOCONAZOLE (OTC)
NIZORAL, KETOCONAZOLE
NOLVADEX, TAMOXIFEN CITRATE
NORCO, ACETAMINOPHEN
NORDETTE-28, ETHINYL ESTRADIOL
NORDITROPIN NORDIFLEX, SOMATROPIN RECOMBINANT
NORDITROPIN, SOMATROPIN RECOMBINANT
NOREPINEPHRINE BITARTRATE, NOREPINEPHRINE BITARTRATE
NORETHIN 1/35E-21, ETHINYL ESTRADIOL
NORETHIN 1/35E-28, ETHINYL ESTRADIOL
NORETHIN 1/50M-21, MESTRANOL
NORETHIN 1/50M-28, MESTRANOL
NORETHINDRONE ACETATE AND ETHINYL ESTRADIOL AND FERROUS FUMARATE, ETHINYL ESTRADIOL
NORETHINDRONE ACETATE, NORETHINDRONE ACETATE
NORETHINDRONE AND ETHINYL ESTRADIOL (10/11), ETHINYL ESTRADIOL
NORETHINDRONE AND ETHINYL ESTRADIOL (7/14), ETHINYL ESTRADIOL
NORETHINDRONE AND ETHINYL ESTRADIOL, ETHINYL ESTRADIOL
NORETHINDRONE AND MESTRANOL, MESTRANOL
NORFLEX, ORPHENADRINE CITRATE
NORGESIC FORTE, ASPIRIN
NORGESIC, ASPIRIN
NORGESTIMATE AND ETHINYL ESTRADIOL, ETHINYL ESTRADIOL
NORINYL 1+35 21-DAY, ETHINYL ESTRADIOL
NORINYL 1+35 28-DAY, ETHINYL ESTRADIOL
NORINYL 1+50 21-DAY, MESTRANOL
NORINYL 1+50 28-DAY, MESTRANOL
NORITATE, METRONIDAZOLE
NORMOCARB HF 25, MAGNESIUM CHLORIDE
NORMOCARB HF 35, MAGNESIUM CHLORIDE
NORMOSOL-M AND DEXTROSE 5% IN PLASTIC CONTAINER, DEXTROSE
NORMOSOL-R AND DEXTROSE 5% IN PLASTIC CONTAINER, DEXTROSE
NORMOSOL-R IN PLASTIC CONTAINER, MAGNESIUM CHLORIDE
NOROXIN, NORFLOXACIN
NORPACE CR, DISOPYRAMIDE PHOSPHATE
NORPACE, DISOPYRAMIDE PHOSPHATE
NORPLANT II, LEVONORGESTREL
NORPRAMIN, DESIPRAMINE HYDROCHLORIDE
NOR-QD, NORETHINDRONE
NORTREL 0.5/35-21, ETHINYL ESTRADIOL
NORTREL 0.5/35-28, ETHINYL ESTRADIOL
NORTREL 1/35-21, ETHINYL ESTRADIOL
NORTREL 1/35-28, ETHINYL ESTRADIOL
NORTREL 7/7/7, ETHINYL ESTRADIOL
NORTRIPTYLINE HYDROCHLORIDE, NORTRIPTYLINE HYDROCHLORIDE
NORVASC, AMLODIPINE BESYLATE
NORVIR, RITONAVIR
NOVAMINE 11.4%, AMINO ACIDS
NOVAMINE 15%, AMINO ACIDS
NOVANTRONE, MITOXANTRONE HYDROCHLORIDE
NOVOCAIN, PROCAINE HYDROCHLORIDE
NOVOLIN 70/30, INSULIN RECOMBINANT HUMAN (OTC)
NOVOLIN N, INSULIN SUSP ISOPHANE RECOMBINANT HUMAN (OTC)
NOVOLIN R, INSULIN RECOMBINANT HUMAN (OTC)
NOVOLOG MIX 70/30, INSULIN ASPART PROTAMINE RECOMBINANT
NOVOLOG, INSULIN ASPART RECOMBINANT
NOVOTHYROX, LEVOTHYROXINE SODIUM
NOXAFIL, POSACONAZOLE
NUBAIN, NALBUPHINE HYDROCHLORIDE
NULYTELY, POLYETHYLENE GLYCOL 3350
NULYTELY-FLAVORED, POLYETHYLENE GLYCOL 3350

APPENDIX A - PRODUCT NAME INDEX

A - 38

**** N ****

NUMORPHAN, OXYMORPHONE HYDROCHLORIDE
NUTRACORT, HYDROCORTISONE
NUTRILIPID 10%, SOYBEAN OIL
NUTRILIPID 20%, SOYBEAN OIL
NUTROPIN AQ PEN, SOMATROPIN RECOMBINANT
NUTROPIN AQ, SOMATROPIN RECOMBINANT
NUTROPIN, SOMATROPIN RECOMBINANT
NUVARING, ETHINYL ESTRADIOL
NYDRAZID, ISONIAZID
NYSTATIN AND TRIAMCINOLONE ACETONIDE, NYSTATIN
NYSTATIN, NYSTATIN
NYSTATIN-TRIAMCINOLONE ACETONIDE, NYSTATIN
NYSTOP, NYSTATIN

**** O ****

OCTOCAINE, EPINEPHRINE
OCTREOSCAN, INDIUM IN-111 PENTETREOTIDE KIT
OCTREOTIDE ACETATE (PRESERVATIVE FREE), OCTREOTIDE ACETATE
OCTREOTIDE ACETATE, OCTREOTIDE ACETATE
OCUCLEAR, OXYMETAZOLINE HYDROCHLORIDE (OTC)
OCUFEN, FLURBIPROFEN SODIUM
OCUFLOX, OFLOXACIN
OCUPRESS, CARTEOLOL HYDROCHLORIDE
OCUSULF-10, SULFACETAMIDE SODIUM
OFLOXACIN, OFLOXACIN
OGEN .625, ESTROPIPATE
OGEN 1.25, ESTROPIPATE
OGEN 2.5, ESTROPIPATE
OGEN 5, ESTROPIPATE
OGEN, ESTROPIPATE
OGESTREL 0.5/50-21, ETHINYL ESTRADIOL
OGESTREL 0.5/50-28, ETHINYL ESTRADIOL
OLUX, CLOBETASOL PROPIONATE
OMACOR, OMEGA-3-ACID ETHYL ESTERS
OMEPRazole, OMEPRazole
OMNARIS, CICLESonIDE
OMNICEF, CEFDINIR
OMNIPaque 140, IOHEXOL
OMNIPaque 180, IOHEXOL
OMNIPaque 240, IOHEXOL
OMNIPaque 300, IOHEXOL
OMNIPaque 350, IOHEXOL
OMNIPRED, PREDNISOLONE ACETATE
OMNISCAN, GADODIAMIDE
OMNITROPE, SOMATROPIN RECOMBINANT
ONDANSETRON HYDROCHLORIDE AND DEXTROSE IN PLASTIC CONTAINER, ONDANSETRON HYDROCHLORIDE
ONDANSETRON HYDROCHLORIDE AND SODIUM CHLORIDE IN PLASTIC CONTAINER, ONDANSETRON HYDROCHLORIDE
ONDANSETRON HYDROCHLORIDE PRESERVATIVE FREE, ONDANSETRON HYDROCHLORIDE
ONDANSETRON HYDROCHLORIDE, ONDANSETRON HYDROCHLORIDE
ONDANSETRON, ONDANSETRON
OPANA ER, OXYMORPHONE HYDROCHLORIDE
OPANA, OXYMORPHONE HYDROCHLORIDE
OPCON-A, NAPHAZOLINE HYDROCHLORIDE (OTC)
OPHTHAINE, PROPARACAINE HYDROCHLORIDE
OPHTHETIC, PROPARACAINE HYDROCHLORIDE
OPTICROM, CROMOLYN SODIUM
OPTIMARK IN PLASTIC CONTAINER, GADOVERSETAMIDE
OPTIMARK, GADOVERSETAMIDE
OPTIPRANOLOL, METIPRANOLOL HYDROCHLORIDE
OPTIRAY 160, IOVERSOL
OPTIRAY 240, IOVERSOL
OPTIRAY 300, IOVERSOL
OPTIRAY 320, IOVERSOL

APPENDIX A - PRODUCT NAME INDEX

A - 39

**** 0 ****

OPTIRAY 350, IOVERSOL
OPTISON, ALBUMIN HUMAN
OPTIVAR, AZELASTINE HYDROCHLORIDE
ORABASE HCA, HYDROCORTISONE ACETATE
ORACEA, DOXYCYCLINE
ORACORT, TRIAMCINOLONE ACETONIDE
ORALONE, TRIAMCINOLONE ACETONIDE
ORAMORPH SR, MORPHINE SULFATE
ORAP, PIMOZIDE
ORAPRED ODT, PREDNISOLONE SODIUM PHOSPHATE
ORAPRED, PREDNISOLONE SODIUM PHOSPHATE
ORAQIX, LIDOCAINE
ORETIC, HYDROCHLOROTHIAZIDE
ORFADIN, NITISINONE
ORPHENADRINE CITRATE, ASPIRIN, AND CAFFEINE, ASPIRIN
ORPHENADRINE CITRATE, ORPHENADRINE CITRATE
ORPHENGESIC FORTE, ASPIRIN
ORPHENGESIC, ASPIRIN
ORTHO CYCLEN-28, ETHINYL ESTRADIOL
ORTHO EVRA, ETHINYL ESTRADIOL
ORTHO TRI-CYCLEN LO, ETHINYL ESTRADIOL
ORTHO TRI-CYCLEN, ETHINYL ESTRADIOL
ORTHO-CEPT, DESOGESTREL
ORTHO-EST, ESTROPIPATE
ORTHO-NOVUM 1/35-28, ETHINYL ESTRADIOL
ORTHO-NOVUM 1/50 28, MESTRANOL
ORTHO-NOVUM 10/11-28, ETHINYL ESTRADIOL
ORTHO-NOVUM 7/7/7-28, ETHINYL ESTRADIOL
ORUVAIL, KETOPROFEN
ORVATEN, MIDODRINE HYDROCHLORIDE
OSMITROL 10% IN WATER IN PLASTIC CONTAINER, MANNITOL
OSMITROL 10% IN WATER, MANNITOL
OSMITROL 15% IN WATER IN PLASTIC CONTAINER, MANNITOL
OSMITROL 15% IN WATER, MANNITOL
OSMITROL 20% IN WATER IN PLASTIC CONTAINER, MANNITOL
OSMITROL 20% IN WATER, MANNITOL
OSMITROL 5% IN WATER IN PLASTIC CONTAINER, MANNITOL
OSMITROL 5% IN WATER, MANNITOL
OSMOPREP, SODIUM PHOSPHATE, DIBASIC, ANHYDROUS
OTICAIR, HYDROCORTISONE
OVCON-35, ETHINYL ESTRADIOL
OVCON-50, ETHINYL ESTRADIOL
OVIDE, MALATHION
OVIDREL, CHORIOGONADOTROPIN ALFA
OXACILLIN SODIUM, OXACILLIN SODIUM
OXANDRIN, OXANDROLONE
OXANDROLONE, OXANDROLONE
OXAPROZIN, OXAPROZIN
OXAZEPAM, OXAZEPAM
OXILAN-300, IOXILAN
OXILAN-350, IOXILAN
OXISTAT, OXICONAZOLE NITRATE
OXSORALEN, METHOXSALEN
OXSORALEN-ULTRA, METHOXSALEN
OXYBUTYNIN CHLORIDE, OXYBUTYNIN CHLORIDE
OXYCET, ACETAMINOPHEN
OXYCODONE AND ACETAMINOPHEN, ACETAMINOPHEN
OXYCODONE AND ASPIRIN, ASPIRIN
OXYCODONE HYDROCHLORIDE, OXYCODONE HYDROCHLORIDE
OXYCONTIN, OXYCODONE HYDROCHLORIDE
OXYTOCIN, OXYTOCIN
OXYTROL, OXYBUTYNIN

APPENDIX A - PRODUCT NAME INDEX

A - 40

**** P ****

PACERONE, AMIODARONE HYDROCHLORIDE
PACLITAXEL, PACLITAXEL
PAMELOR, NORTRIPTYLINE HYDROCHLORIDE
PAMIDRONATE DISODIUM, PAMIDRONATE DISODIUM
PAMINE FORTE, METHSCOPOLAMINE BROMIDE
PAMINE, METHSCOPOLAMINE BROMIDE
PANCURONIUM BROMIDE, PANCURONIUM BROMIDE
PANCURONIUM, PANCURONIUM BROMIDE
PANDEL, HYDROCORTISONE PROBUTATE
PANIXINE DISPERDOSE, CEPHALEXIN
PANRETIN, ALITRETINOIN
PARACAINE, PROPACAINE HYDROCHLORIDE
PARAFON FORTE DSC, CHLORZOXAZONE
PARAGARD T 380A, COPPER
PARAPLATIN, CARBOPLATIN
PARCOPA, CARBIDOPA
PAREMYD, HYDROXYAMPHETAMINE HYDROBROMIDE
PARLODEL, BROMOCRIPTINE MESYLATE
PARNATE, TRANLYCYPROMINE SULFATE
PAROMOMYCIN SULFATE, PAROMOMYCIN SULFATE
PAROXETINE HYDROCHLORIDE, PAROXETINE HYDROCHLORIDE
PASER, AMINOSALICYLIC ACID
PASKALIUM, POTASSIUM AMINOSALICYLATE
PATADAY, OLOPATADINE HYDROCHLORIDE
PATANOL, OLOPATADINE HYDROCHLORIDE
PAXIL CR, PAROXETINE HYDROCHLORIDE
PAXIL, PAROXETINE HYDROCHLORIDE
PCE, ERYTHROMYCIN
PEDIAMYCIN 400, ERYTHROMYCIN ETHYLSUCCINATE
PEDIAMYCIN, ERYTHROMYCIN ETHYLSUCCINATE
PEDIAPRED, PREDNISOLONE SODIUM PHOSPHATE
PEDIATRIC ADVIL, IBUPROFEN (OTC)
PEDIATRIC LTA KIT, LIDOCAINE HYDROCHLORIDE
PEDIAZOLE, ERYTHROMYCIN ETHYLSUCCINATE
PEDIOTIC, HYDROCORTISONE
PEGANONE, ETHOTOIN
PENICILLIN G POTASSIUM IN PLASTIC CONTAINER, PENICILLIN G POTASSIUM
PENICILLIN G POTASSIUM, PENICILLIN G POTASSIUM
PENICILLIN G PROCAINE, PENICILLIN G PROCAINE
PENICILLIN G SODIUM, PENICILLIN G SODIUM
PENICILLIN V POTASSIUM, PENICILLIN V POTASSIUM
PENICILLIN-VK, PENICILLIN V POTASSIUM
PENLAC, CICLOPIROX
PENTAM, PENTAMIDINE ISETHIONATE
PENTAMIDINE ISETHIONATE, PENTAMIDINE ISETHIONATE
PENTASA, MESALAMINE
PENTAZOCINE AND NALOXONE HYDROCHLORIDES, NALOXONE HYDROCHLORIDE
PENTAZOCINE HYDROCHLORIDE AND ACETAMINOPHEN, ACETAMINOPHEN
PENTETATE CALCIUM TRISODIUM, PENTETATE CALCIUM TRISODIUM
PENTETATE ZINC TRISODIUM, PENTETATE ZINC TRISODIUM
PENTOLAIR, CYCLOPENTOLATE HYDROCHLORIDE
PENTOXIFYLLINE, PENTOXIFYLLINE
PENTOXIL, PENTOXIFYLLINE
PEPCID AC (GELTAB), FAMOTIDINE (OTC)
PEPCID AC, FAMOTIDINE (OTC)
PEPCID COMPLETE, CALCIUM CARBONATE (OTC)
PEPCID PRESERVATIVE FREE IN PLASTIC CONTAINER, FAMOTIDINE
PEPCID PRESERVATIVE FREE, FAMOTIDINE
PEPCID, FAMOTIDINE
PERCOCET, ACETAMINOPHEN
PERCODAN, ASPIRIN
PERGOLIDE MESYLATE, PERGOLIDE MESYLATE
PERIDEX, CHLORHEXIDINE GLUCONATE

APPENDIX A - PRODUCT NAME INDEX

A - 41

**** P ****

PERIOCHIP, CHLORHEXIDINE GLUCONATE
PERIOGARD, CHLORHEXIDINE GLUCONATE
PERIOSTAT, DOXYCYCLINE HYCLATE
PERMAPEN, PENICILLIN G BENZATHINE
PERMAX, PERGOLIDE MESYLATE
PERMETHRIN, PERMETHRIN
PERMETHRIN, PERMETHRIN (OTC)
PERPHENAZINE AND AMITRIPTYLINE HYDROCHLORIDE, AMITRIPTYLINE HYDROCHLORIDE
PERPHENAZINE, PERPHENAZINE
PERSANTINE, DIPYRIDAMOLE
PEXEVA, PAROXETINE MESYLATE
PFIZERPEN, PENICILLIN G POTASSIUM
PHARMASEAL SCRUB CARE, CHLORHEXIDINE GLUCONATE (OTC)
PHENDIMETRAZINE TARTRATE, PHENDIMETRAZINE TARTRATE
PHENERGAN, PROMETHAZINE HYDROCHLORIDE
PHENTERMINE HYDROCHLORIDE, PHENTERMINE HYDROCHLORIDE
PHENTOLAMINE MESYLATE, PHENTOLAMINE MESYLATE
PHENTYTOIN, PHENYTOIN
PHENYLEPHRINE HYDROCHLORIDE AND PROMETHAZINE HYDROCHLORIDE, PHENYLEPHRINE HYDROCHLORIDE
PHENYTEK, PHENYTOIN SODIUM
PHENYTOIN SODIUM, PHENYTOIN SODIUM
PHENYTOIN, PHENYTOIN
PHENYTOIN, PHENYTOIN SODIUM
PHISOHEX, HEXACHLOROPHENE
PHOSLO GELCAPS, CALCIUM ACETATE
PHOSPHOCOL P32, CHROMIC PHOSPHATE, P-32
PHOSPHOLINE IODIDE, ECHOTHIOPHATE IODIDE
PHOSPHOTEC, TECHNETIUM TC-99M PYROPHOSPHATE KIT
PHOTOFRIN, PORFIMER SODIUM
PHRENILIN FORTE, ACETAMINOPHEN
PHRENILIN WITH CAFFEINE AND CODEINE, ACETAMINOPHEN
PHRENILIN, ACETAMINOPHEN
PHYSIOLYTE IN PLASTIC CONTAINER, MAGNESIUM CHLORIDE
PHYSIOSOL IN PLASTIC CONTAINER, MAGNESIUM CHLORIDE
PHYSIOSOL PH 7.4 IN PLASTIC CONTAINER, MAGNESIUM CHLORIDE
PHYTONADIONE, PHYTONADIONE
PILOCARPINE HYDROCHLORIDE, PILOCARPINE HYDROCHLORIDE
PILOPINE HS, PILOCARPINE HYDROCHLORIDE
PINDOLOL, PINDOLOL
PIPERACILLIN, PIPERACILLIN SODIUM
PIROXICAM, PIROXICAM
PITOCIN, OXYTOCIN
PLAN B, LEVONORGESTREL
PLAN B, LEVONORGESTREL (OTC)
PLAQUENIL, HYDROXYCHLOROQUINE SULFATE
PLASMA-LYTE 148 AND DEXTROSE 5% IN PLASTIC CONTAINER, DEXTROSE
PLASMA-LYTE 148 IN WATER IN PLASTIC CONTAINER, MAGNESIUM CHLORIDE
PLASMA-LYTE 56 AND DEXTROSE 5% IN PLASTIC CONTAINER, DEXTROSE
PLASMA-LYTE 56 IN PLASTIC CONTAINER, MAGNESIUM ACETATE TETRAHYDRATE
PLASMA-LYTE A IN PLASTIC CONTAINER, MAGNESIUM CHLORIDE
PLASMA-LYTE M AND DEXTROSE 5% IN PLASTIC CONTAINER, CALCIUM CHLORIDE
PLASMA-LYTE R IN PLASTIC CONTAINER, CALCIUM CHLORIDE
PLATINOL, CISPLATIN
PLATINOL-AQ, CISPLATIN
PLAVIX, CLOPIDOGREL BISULFATE
PLEGISOL IN PLASTIC CONTAINER, CALCIUM CHLORIDE
PLENDIL, FELODIPINE
PLETAL, CILOSTAZOL
PODOFILOX, PODOFILOX
POLOCAINE W/ LEVONORDEFIN, LEVONORDEFIN
POLOCAINE, MEPIVACAINE HYDROCHLORIDE
POLOCAINE-MPF, MEPIVACAINE HYDROCHLORIDE
POLYETHYLENE GLYCOL 3350, POLYETHYLENE GLYCOL 3350

APPENDIX A - PRODUCT NAME INDEX

A - 42

**** P ****

POLYMYXIN B SULFATE, POLYMYXIN B SULFATE
POLY-PRED, NEOMYCIN SULFATE
POLY-RX, POLYMYXIN B SULFATE
POLYSPORIN, BACITRACIN ZINC
POLYTRIM, POLYMYXIN B SULFATE
PONSTEL, MEFENAMIC ACID
PORTIA-28, ETHINYL ESTRADIOL
POTASSIUM ACETATE IN PLASTIC CONTAINER, POTASSIUM ACETATE
POTASSIUM CHLORIDE 0.037% IN DEXTROSE 10% AND SODIUM CHLORIDE 0.2% IN PLASTIC CONTAINER, DEXTROSE
POTASSIUM CHLORIDE 0.037% IN DEXTROSE 10% AND SODIUM CHLORIDE 0.45% IN PLASTIC CONTAINER, DEXTROSE
POTASSIUM CHLORIDE 0.037% IN DEXTROSE 10% AND SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER, DEXTROSE
POTASSIUM CHLORIDE 0.037% IN DEXTROSE 5% AND SODIUM CHLORIDE 0.11% IN PLASTIC CONTAINER, DEXTROSE
POTASSIUM CHLORIDE 0.037% IN DEXTROSE 5% AND SODIUM CHLORIDE 0.2% IN PLASTIC CONTAINER, DEXTROSE
POTASSIUM CHLORIDE 0.037% IN DEXTROSE 5% AND SODIUM CHLORIDE 0.33% IN PLASTIC CONTAINER, DEXTROSE
POTASSIUM CHLORIDE 0.037% IN DEXTROSE 5% AND SODIUM CHLORIDE 0.45% IN PLASTIC CONTAINER, DEXTROSE
POTASSIUM CHLORIDE 0.037% IN DEXTROSE 5% AND SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER, DEXTROSE
POTASSIUM CHLORIDE 0.075% IN DEXTROSE 10% AND SODIUM CHLORIDE 0.2% IN PLASTIC CONTAINER, DEXTROSE
POTASSIUM CHLORIDE 0.075% IN DEXTROSE 10% AND SODIUM CHLORIDE 0.45% IN PLASTIC CONTAINER, DEXTROSE
POTASSIUM CHLORIDE 0.075% IN DEXTROSE 10% AND SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER, DEXTROSE
POTASSIUM CHLORIDE 0.075% IN DEXTROSE 3.3% AND SODIUM CHLORIDE 0.3% IN PLASTIC CONTAINER, DEXTROSE
POTASSIUM CHLORIDE 0.075% IN DEXTROSE 5% AND SODIUM CHLORIDE 0.11% IN PLASTIC CONTAINER, DEXTROSE
POTASSIUM CHLORIDE 0.075% IN DEXTROSE 5% AND SODIUM CHLORIDE 0.2% IN PLASTIC CONTAINER, DEXTROSE
POTASSIUM CHLORIDE 0.075% IN DEXTROSE 5% AND SODIUM CHLORIDE 0.33% IN PLASTIC CONTAINER, DEXTROSE
POTASSIUM CHLORIDE 0.075% IN DEXTROSE 5% AND SODIUM CHLORIDE 0.45% IN PLASTIC CONTAINER, DEXTROSE
POTASSIUM CHLORIDE 0.075% IN DEXTROSE 5% AND SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER, DEXTROSE
POTASSIUM CHLORIDE 0.075% IN DEXTROSE 5% IN PLASTIC CONTAINER, DEXTROSE
POTASSIUM CHLORIDE 0.11% IN DEXTROSE 10% AND SODIUM CHLORIDE 0.2% IN PLASTIC CONTAINER, DEXTROSE
POTASSIUM CHLORIDE 0.11% IN DEXTROSE 10% AND SODIUM CHLORIDE 0.45% IN PLASTIC CONTAINER, DEXTROSE
POTASSIUM CHLORIDE 0.11% IN DEXTROSE 10% AND SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER, DEXTROSE
POTASSIUM CHLORIDE 0.11% IN DEXTROSE 3.3% AND SODIUM CHLORIDE 0.3% IN PLASTIC CONTAINER, DEXTROSE
POTASSIUM CHLORIDE 0.11% IN DEXTROSE 5% AND SODIUM CHLORIDE 0.11% IN PLASTIC CONTAINER, DEXTROSE
POTASSIUM CHLORIDE 0.11% IN DEXTROSE 5% AND SODIUM CHLORIDE 0.2% IN PLASTIC CONTAINER, DEXTROSE
POTASSIUM CHLORIDE 0.11% IN DEXTROSE 5% AND SODIUM CHLORIDE 0.33% IN PLASTIC CONTAINER, DEXTROSE
POTASSIUM CHLORIDE 0.11% IN DEXTROSE 5% AND SODIUM CHLORIDE 0.45% IN PLASTIC CONTAINER, DEXTROSE
POTASSIUM CHLORIDE 0.11% IN DEXTROSE 5% AND SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER, DEXTROSE
POTASSIUM CHLORIDE 0.15% IN DEXTROSE 10% AND SODIUM CHLORIDE 0.2% IN PLASTIC CONTAINER, DEXTROSE
POTASSIUM CHLORIDE 0.15% IN DEXTROSE 10% AND SODIUM CHLORIDE 0.45% IN PLASTIC CONTAINER, DEXTROSE
POTASSIUM CHLORIDE 0.15% IN DEXTROSE 10% AND SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER, DEXTROSE
POTASSIUM CHLORIDE 0.15% IN DEXTROSE 3.3% AND SODIUM CHLORIDE 0.3% IN PLASTIC CONTAINER, DEXTROSE
POTASSIUM CHLORIDE 0.15% IN DEXTROSE 5% AND SODIUM CHLORIDE 0.11% IN PLASTIC CONTAINER, DEXTROSE
POTASSIUM CHLORIDE 0.15% IN DEXTROSE 5% AND SODIUM CHLORIDE 0.2% IN PLASTIC CONTAINER, DEXTROSE
POTASSIUM CHLORIDE 0.15% IN DEXTROSE 5% AND SODIUM CHLORIDE 0.33% IN PLASTIC CONTAINER, DEXTROSE
POTASSIUM CHLORIDE 0.15% IN DEXTROSE 5% AND SODIUM CHLORIDE 0.45% IN PLASTIC CONTAINER, DEXTROSE
POTASSIUM CHLORIDE 0.15% IN DEXTROSE 5% AND SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER, DEXTROSE
POTASSIUM CHLORIDE 0.15% IN DEXTROSE 5% IN PLASTIC CONTAINER, DEXTROSE
POTASSIUM CHLORIDE 0.15% IN SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER, POTASSIUM CHLORIDE
POTASSIUM CHLORIDE 0.22% IN DEXTROSE 10% AND SODIUM CHLORIDE 0.2% IN PLASTIC CONTAINER, DEXTROSE
POTASSIUM CHLORIDE 0.22% IN DEXTROSE 10% AND SODIUM CHLORIDE 0.45% IN PLASTIC CONTAINER, DEXTROSE
POTASSIUM CHLORIDE 0.22% IN DEXTROSE 10% AND SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER, DEXTROSE
POTASSIUM CHLORIDE 0.22% IN DEXTROSE 3.3% AND SODIUM CHLORIDE 0.3% IN PLASTIC CONTAINER, DEXTROSE
POTASSIUM CHLORIDE 0.22% IN DEXTROSE 5% AND SODIUM CHLORIDE 0.11% IN PLASTIC CONTAINER, DEXTROSE
POTASSIUM CHLORIDE 0.22% IN DEXTROSE 5% AND SODIUM CHLORIDE 0.2% IN PLASTIC CONTAINER, DEXTROSE
POTASSIUM CHLORIDE 0.22% IN DEXTROSE 5% AND SODIUM CHLORIDE 0.33% IN PLASTIC CONTAINER, DEXTROSE
POTASSIUM CHLORIDE 0.22% IN DEXTROSE 5% AND SODIUM CHLORIDE 0.45% IN PLASTIC CONTAINER, DEXTROSE
POTASSIUM CHLORIDE 0.22% IN DEXTROSE 5% AND SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER, DEXTROSE
POTASSIUM CHLORIDE 0.22% IN DEXTROSE 5% IN PLASTIC CONTAINER, DEXTROSE
POTASSIUM CHLORIDE 0.3% IN DEXTROSE 10% AND SODIUM CHLORIDE 0.2% IN PLASTIC CONTAINER, DEXTROSE
POTASSIUM CHLORIDE 0.3% IN DEXTROSE 10% AND SODIUM CHLORIDE 0.45% IN PLASTIC CONTAINER, DEXTROSE
POTASSIUM CHLORIDE 0.3% IN DEXTROSE 10% AND SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER, DEXTROSE
POTASSIUM CHLORIDE 0.3% IN DEXTROSE 3.3% AND SODIUM CHLORIDE 0.3% IN PLASTIC CONTAINER, DEXTROSE
POTASSIUM CHLORIDE 0.3% IN DEXTROSE 5% AND SODIUM CHLORIDE 0.11% IN PLASTIC CONTAINER, DEXTROSE

APPENDIX A - PRODUCT NAME INDEX

A - 43

**** P ****

POTASSIUM CHLORIDE 0.3% IN DEXTROSE 5% AND SODIUM CHLORIDE 0.2% IN PLASTIC CONTAINER, DEXTROSE
POTASSIUM CHLORIDE 0.3% IN DEXTROSE 5% AND SODIUM CHLORIDE 0.33% IN PLASTIC CONTAINER, DEXTROSE
POTASSIUM CHLORIDE 0.3% IN DEXTROSE 5% AND SODIUM CHLORIDE 0.45% IN PLASTIC CONTAINER, DEXTROSE
POTASSIUM CHLORIDE 0.3% IN DEXTROSE 5% AND SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER, DEXTROSE
POTASSIUM CHLORIDE 0.3% IN DEXTROSE 5% IN PLASTIC CONTAINER, DEXTROSE
POTASSIUM CHLORIDE 10MEQ IN DEXTROSE 5% AND LACTATED RINGER'S IN PLASTIC CONTAINER, CALCIUM
CHLORIDE
POTASSIUM CHLORIDE 10MEQ IN DEXTROSE 5% AND SODIUM CHLORIDE 0.225% IN PLASTIC CONTAINER, DEXTROSE
POTASSIUM CHLORIDE 10MEQ IN DEXTROSE 5% AND SODIUM CHLORIDE 0.3% IN PLASTIC CONTAINER, DEXTROSE
POTASSIUM CHLORIDE 10MEQ IN DEXTROSE 5% AND SODIUM CHLORIDE 0.45% IN PLASTIC CONTAINER, DEXTROSE
POTASSIUM CHLORIDE 10MEQ IN DEXTROSE 5% AND SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER, DEXTROSE
POTASSIUM CHLORIDE 10MEQ IN PLASTIC CONTAINER, POTASSIUM CHLORIDE
POTASSIUM CHLORIDE 15MEQ IN DEXTROSE 5% AND LACTATED RINGER'S IN PLASTIC CONTAINER, CALCIUM
CHLORIDE
POTASSIUM CHLORIDE 15MEQ IN DEXTROSE 5% AND SODIUM CHLORIDE 0.225% IN PLASTIC CONTAINER, DEXTROSE
POTASSIUM CHLORIDE 15MEQ IN DEXTROSE 5% AND SODIUM CHLORIDE 0.3% IN PLASTIC CONTAINER, DEXTROSE
POTASSIUM CHLORIDE 15MEQ IN DEXTROSE 5% AND SODIUM CHLORIDE 0.45% IN PLASTIC CONTAINER, DEXTROSE
POTASSIUM CHLORIDE 15MEQ IN DEXTROSE 5% AND SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER, DEXTROSE
POTASSIUM CHLORIDE 20MEQ IN DEXTROSE 5% AND LACTATED RINGER'S IN PLASTIC CONTAINER, CALCIUM
CHLORIDE
POTASSIUM CHLORIDE 20MEQ IN DEXTROSE 5% AND SODIUM CHLORIDE 0.225% IN PLASTIC CONTAINER, DEXTROSE
POTASSIUM CHLORIDE 20MEQ IN DEXTROSE 5% AND SODIUM CHLORIDE 0.3% IN PLASTIC CONTAINER, DEXTROSE
POTASSIUM CHLORIDE 20MEQ IN DEXTROSE 5% AND SODIUM CHLORIDE 0.45% IN PLASTIC CONTAINER, DEXTROSE
POTASSIUM CHLORIDE 20MEQ IN DEXTROSE 5% AND SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER, DEXTROSE
POTASSIUM CHLORIDE 20MEQ IN DEXTROSE 5% IN PLASTIC CONTAINER, DEXTROSE
POTASSIUM CHLORIDE 20MEQ IN DEXTROSE 5% IN SODIUM CHLORIDE 0.3% IN PLASTIC CONTAINER, DEXTROSE
POTASSIUM CHLORIDE 20MEQ IN PLASTIC CONTAINER, POTASSIUM CHLORIDE
POTASSIUM CHLORIDE 20MEQ IN SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER, POTASSIUM CHLORIDE
POTASSIUM CHLORIDE 30MEQ IN DEXTROSE 5% AND LACTATED RINGER'S IN PLASTIC CONTAINER, CALCIUM
CHLORIDE
POTASSIUM CHLORIDE 30MEQ IN DEXTROSE 5% AND SODIUM CHLORIDE 0.225% IN PLASTIC CONTAINER, DEXTROSE
POTASSIUM CHLORIDE 30MEQ IN DEXTROSE 5% AND SODIUM CHLORIDE 0.3% IN PLASTIC CONTAINER, DEXTROSE
POTASSIUM CHLORIDE 30MEQ IN DEXTROSE 5% AND SODIUM CHLORIDE 0.45% IN PLASTIC CONTAINER, DEXTROSE
POTASSIUM CHLORIDE 30MEQ IN DEXTROSE 5% AND SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER, DEXTROSE
POTASSIUM CHLORIDE 30MEQ IN DEXTROSE 5% IN PLASTIC CONTAINER, DEXTROSE
POTASSIUM CHLORIDE 30MEQ IN PLASTIC CONTAINER, POTASSIUM CHLORIDE
POTASSIUM CHLORIDE 40MEQ IN DEXTROSE 5% AND LACTATED RINGER'S IN PLASTIC CONTAINER, CALCIUM
CHLORIDE
POTASSIUM CHLORIDE 40MEQ IN DEXTROSE 5% AND SODIUM CHLORIDE 0.225% IN PLASTIC CONTAINER, DEXTROSE
POTASSIUM CHLORIDE 40MEQ IN DEXTROSE 5% AND SODIUM CHLORIDE 0.3% IN PLASTIC CONTAINER, DEXTROSE
POTASSIUM CHLORIDE 40MEQ IN DEXTROSE 5% AND SODIUM CHLORIDE 0.45% IN PLASTIC CONTAINER, DEXTROSE
POTASSIUM CHLORIDE 40MEQ IN DEXTROSE 5% AND SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER, DEXTROSE
POTASSIUM CHLORIDE 40MEQ IN DEXTROSE 5% IN PLASTIC CONTAINER, DEXTROSE
POTASSIUM CHLORIDE 40MEQ IN PLASTIC CONTAINER, POTASSIUM CHLORIDE
POTASSIUM CHLORIDE 40MEQ IN SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER, POTASSIUM CHLORIDE
POTASSIUM CHLORIDE 5MEQ IN DEXTROSE 5% AND LACTATED RINGER'S IN PLASTIC CONTAINER, CALCIUM
CHLORIDE
POTASSIUM CHLORIDE 5MEQ IN DEXTROSE 5% AND SODIUM CHLORIDE 0.225% IN PLASTIC CONTAINER, DEXTROSE
POTASSIUM CHLORIDE 5MEQ IN DEXTROSE 5% AND SODIUM CHLORIDE 0.3% IN PLASTIC CONTAINER, DEXTROSE
POTASSIUM CHLORIDE 5MEQ IN DEXTROSE 5% AND SODIUM CHLORIDE 0.45% IN PLASTIC CONTAINER, DEXTROSE
POTASSIUM CHLORIDE 5MEQ IN DEXTROSE 5% AND SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER, DEXTROSE
POTASSIUM CHLORIDE IN PLASTIC CONTAINER, POTASSIUM CHLORIDE
POTASSIUM CHLORIDE, POTASSIUM CHLORIDE
POTASSIUM CITRATE, POTASSIUM CITRATE
POVIDONE IODINE, POVIDONE-IODINE (OTC)
PRALIDOXIME CHLORIDE, PRALIDOXIME CHLORIDE
PRAMOSONE, HYDROCORTISONE ACETATE
PRANDIN, REPAGLINIDE
PRAVACHOL, PRAVASTATIN SODIUM
PRAVASTATIN SODIUM, PRAVASTATIN SODIUM
PRAVIGARD PAC (COPACKAGED), ASPIRIN
PRAZOSIN HYDROCHLORIDE, PRAZOSIN HYDROCHLORIDE
PRECEDEX, DEXMEDETOMIDINE
PRECOSE, ACARBOSE

APPENDIX A - PRODUCT NAME INDEX

A - 44

**** P ****

PRED FORTE, PREDNISOLONE ACETATE
PRED MILD, PREDNISOLONE ACETATE
PRED-G, GENTAMICIN SULFATE
PREDNICARBATE, PREDNICARBATE
PREDNISOLONE SODIUM PHOSPHATE, PREDNISOLONE SODIUM PHOSPHATE
PREDNISOLONE, PREDNISOLONE
PREDNISONE INTENSOL, PREDNISONE
PREDNISONE, PREDNISONE
PREFEST, ESTRADIOL
PREGNYL, GONADOTROPIN, CHORIONIC
PRELONE, PREDNISOLONE
PREMARIN, ESTROGENS, CONJUGATED
PREMASOL 10% IN PLASTIC CONTAINER, AMINO ACIDS
PREMASOL 6% IN PLASTIC CONTAINER, AMINO ACIDS
PREMPHASE 14/14, ESTROGENS, CONJUGATED
PREMPRO, ESTROGENS, CONJUGATED
PRE-OP II, HEXACHLOROPHENE
PRE-OP, HEXACHLOROPHENE
PREPIDIL, DINOPROSTONE
PREVACARE R, CHLORHEXIDINE GLUCONATE (OTC)
PREVACID IV, LANSOPRAZOLE
PREVACID NAPRAPAC 500 (COPACKAGED), LANSOPRAZOLE
PREVACID, LANSOPRAZOLE
PREVALITE, CHOLESTYRAMINE
PREVIFEM, ETHINYL ESTRADIOL
PREVPAC, AMOXICILLIN
PREZISTA, DARUNAVIR ETHANOLATE
PRIALT, ZICONOTIDE
PRIFTIN, RIFAPENTINE
PRILOSEC OTC, OMEPRAZOLE MAGNESIUM (OTC)
PRILOSEC, OMEPRAZOLE
PRIMACOR IN DEXTROSE 5% IN PLASTIC CONTAINER, MILRINONE LACTATE
PRIMACOR, MILRINONE LACTATE
PRIMAQUINE, PRIMAQUINE PHOSPHATE
PRIMATENE MIST, EPINEPHRINE (OTC)
PRIMAXIN, CILASTATIN SODIUM
PRIMIDONE, PRIMIDONE
PRIMSOL, TRIMETHOPRIM HYDROCHLORIDE
PRINCIPEN, AMPICILLIN/AMPICILLIN TRIHYDRATE
PRINIVIL, LISINAPRIL
PRINZIDE, HYDROCHLOROTHIAZIDE
PRISMASOL BGK 0/2.5 IN PLASTIC CONTAINER, CALCIUM CHLORIDE
PRISMASOL BGK 2/0 IN PLASTIC CONTAINER, CALCIUM CHLORIDE
PRISMASOL BGK 2/3.5 IN PLASTIC CONTAINER, CALCIUM CHLORIDE
PRISMASOL BGK 4/0 IN PLASTIC CONTAINER, CALCIUM CHLORIDE
PRISMASOL BGK 4/2.5 IN PLASTIC CONTAINER, CALCIUM CHLORIDE
PRISMASOL BK 0/0 IN PLASTIC CONTAINER, CALCIUM CHLORIDE
PRISMASOL BK 0/3.5 IN PLASTIC CONTAINER, CALCIUM CHLORIDE
PROAIR HFA, ALBUTEROL SULFATE
PROAMATINE, MIDODRINE HYDROCHLORIDE
PROBALAN, PROBENECID
PRO-BANTHINE, PROPANTHELINE BROMIDE
PROBENECID AND COLCHICINE, COLCHICINE
PROBENECID, PROBENECID
PROCAINAMIDE HYDROCHLORIDE, PROCAINAMIDE HYDROCHLORIDE
PROCAINE HYDROCHLORIDE, PROCAINE HYDROCHLORIDE
PROCALAMINE, AMINO ACIDS
PROCANBID, PROCAINAMIDE HYDROCHLORIDE
PROCARDIA XL, NIFEDIPINE
PROCARDIA, NIFEDIPINE
PROCHLORPERAZINE EDISYLATE, PROCHLORPERAZINE EDISYLATE
PROCHLORPERAZINE MALEATE, PROCHLORPERAZINE MALEATE
PROCHLORPERAZINE, PROCHLORPERAZINE

APPENDIX A - PRODUCT NAME INDEX

A - 45

**** P ****

PROCHLORPERAZINE, PROCHLORPERAZINE EDISYLATE
PROCTOFOAM HC, HYDROCORTISONE ACETATE
PROFEN, IBUPROFEN (OTC)
PROFERDEX, IRON DEXTRAN
PROGESTERONE, PROGESTERONE
PROGLYCEM, DIAZOXIDE
PROGRAF, TACROLIMUS
PROHANCE MULTIPACK, GADOTERIDOL
PROHANCE, GADOTERIDOL
PROLIXIN DECANOATE, FLUPHENAZINE DECANOATE
PROLIXIN, FLUPHENAZINE HYDROCHLORIDE
PROLOPRIM, TRIMETHOPRIM
PROMETH HYDROCHLORIDE, PHENYLEPHRINE HYDROCHLORIDE W/CODEINE PHOSPHATE, CODEINE PHOSPHATE
PROMETH VC PLAIN, PHENYLEPHRINE HYDROCHLORIDE
PROMETH VC W/ CODEINE, CODEINE PHOSPHATE
PROMETH W/ CODEINE, CODEINE PHOSPHATE
PROMETH W/ DEXTROMETHORPHAN, DEXTROMETHORPHAN HYDROBROMIDE
PROMETHACON, PROMETHAZINE HYDROCHLORIDE
PROMETHAZINE DM, DEXTROMETHORPHAN HYDROBROMIDE
PROMETHAZINE HYDROCHLORIDE AND CODEINE PHOSPHATE, CODEINE PHOSPHATE
PROMETHAZINE HYDROCHLORIDE AND DEXTROMETHORPHAN HYDROBROMIDE, DEXTROMETHORPHAN HYDROBROMIDE
PROMETHAZINE HYDROCHLORIDE, PROMETHAZINE HYDROCHLORIDE
PROMETHAZINE PLAIN, PROMETHAZINE HYDROCHLORIDE
PROMETHAZINE W/ CODEINE, CODEINE PHOSPHATE
PROMETHAZINE W/ DEXTROMETHORPHAN, DEXTROMETHORPHAN HYDROBROMIDE
PROMETHAZINE WITH CODEINE SYRUP, CODEINE PHOSPHATE
PROMETHEGAN, PROMETHAZINE HYDROCHLORIDE
PROMETRIUM, PROGESTERONE
PROMPT PHENYTOIN SODIUM, PHENYTOIN SODIUM
PRONESTYL, PROCAINAMIDE HYDROCHLORIDE
PRONESTYL-SR, PROCAINAMIDE HYDROCHLORIDE
PROPAPENONE HYDROCHLORIDE, PROPAPENONE HYDROCHLORIDE
PROPANTHELINE BROMIDE, PROPANTHELINE BROMIDE
PROPARACAINE HYDROCHLORIDE, PROPARACAINE HYDROCHLORIDE
PROPECIA, FINASTERIDE
PROPINE, DIPIVEFRIN HYDROCHLORIDE
PROPOFOL, PROPOFOL
PROPOXYPHENE COMPOUND 65, ASPIRIN
PROPOXYPHENE HYDROCHLORIDE AND ACETAMINOPHEN, ACETAMINOPHEN
PROPOXYPHENE HYDROCHLORIDE, PROPOXYPHENE HYDROCHLORIDE
PROPOXYPHENE NAPSYLATE AND ACETAMINOPHEN, ACETAMINOPHEN
PROPRANOLOL HYDROCHLORIDE AND HYDROCHLOROTHIAZIDE, HYDROCHLOROTHIAZIDE
PROPRANOLOL HYDROCHLORIDE INTENSOL, PROPRANOLOL HYDROCHLORIDE
PROPRANOLOL HYDROCHLORIDE, PROPRANOLOL HYDROCHLORIDE
PROPRANOLOL, PROPRANOLOL HYDROCHLORIDE
PROPYLTHIOURACIL, PROPYLTHIOURACIL
PROQUIN XR, CIPROFLOXACIN HYDROCHLORIDE
PROSCAR, FINASTERIDE
PROSOL 20% SULFITE FREE IN PLASTIC CONTAINER, AMINO ACIDS
PROSOM, ESTAZOLAM
PROSTEP, NICOTINE (OTC)
PROSTIN E2, DINOPROSTONE
PROSTIN VR PEDIATRIC, ALPROSTADIL
PROTAMINE SULFATE, PROTAMINE SULFATE
PROTONIX IV, PANTOPRAZOLE SODIUM
PROTONIX, PANTOPRAZOLE SODIUM
PROTOPAM CHLORIDE, PRALIDOXIME CHLORIDE
PROTOPIC, TACROLIMUS
PROVENTIL, ALBUTEROL
PROVENTIL-HFA, ALBUTEROL SULFATE
PROVERA, MEDROXYPROGESTERONE ACETATE
PROVIGIL, MODAFINIL
PROVOCHOLINE, METHACHOLINE CHLORIDE

APPENDIX A - PRODUCT NAME INDEX

A - 46

**** P ****

PROZAC WEEKLY, FLUOXETINE HYDROCHLORIDE
PROZAC, FLUOXETINE HYDROCHLORIDE
PSEUDOEPHEDRINE HYDROCHLORIDE AND TRIPROLIDINE HYDROCHLORIDE, PSEUDOEPHEDRINE HYDROCHLORIDE
PSEUDOEPHEDRINE HYDROCHLORIDE, PSEUDOEPHEDRINE HYDROCHLORIDE (OTC)
PSORCON, DIFLORASONE DIACETATE
PULMICORT RESPULES, BUDESONIDE
PULMICORT, BUDESONIDE
PULMOLITE, TECHNETIUM TC-99M ALBUMIN AGGREGATED KIT
PURINETHOL, MERCAPTOPURINE
PYLERA, BISKALCITRATE
PYRAZINAMIDE, PYRAZINAMIDE
PYRIDOSTIGMINE BROMIDE, PYRIDOSTIGMINE BROMIDE
PYRIDOXINE HYDROCHLORIDE, PYRIDOXINE HYDROCHLORIDE
PYTEST KIT, UREA, C-14
PYTEST, UREA, C-14

**** Q ****

QUADRAMET, SAMARIUM SM 153 LEXIDRONAM PENTASODIUM
QUASENSE, ETHINYL ESTRADIOL
QUELICIN PRESERVATIVE FREE, SUCCINYLCHOLINE CHLORIDE
QUELICIN, SUCCINYLCHOLINE CHLORIDE
QUESTRAN LIGHT, CHOLESTYRAMINE
QUESTRAN, CHOLESTYRAMINE
QUIBRON-T, THEOPHYLLINE
QUIBRON-T/SR, THEOPHYLLINE
QUINAPRIL HYDROCHLORIDE AND HYDROCHLOROTHIAZIDE, HYDROCHLOROTHIAZIDE
QUINAPRIL HYDROCHLORIDE, QUINAPRIL HYDROCHLORIDE
QUINAPRIL, QUINAPRIL HYDROCHLORIDE
QUINARETIC, HYDROCHLOROTHIAZIDE
QUINIDINE GLUCONATE, QUINIDINE GLUCONATE
QUINIDINE SULFATE, QUINIDINE SULFATE
QUININE SULFATE, QUININE SULFATE
QUIXIN, LEVOFLOXACIN
QVAR 40, BECLOMETHASONE DIPROPIONATE
QVAR 80, BECLOMETHASONE DIPROPIONATE

**** R ****

RADIOGARDASE (PRUSSIAN BLUE), FERRIC HEXACYANOFERRATE(II)
RAMIPRIL, RAMIPRIL
RANEXA, RANOLAZINE
RANICLOR, CEFACLOX
RANITIDINE HYDROCHLORIDE, RANITIDINE HYDROCHLORIDE
RANITIDINE, RANITIDINE HYDROCHLORIDE
RANITIDINE, RANITIDINE HYDROCHLORIDE (OTC)
RAPAMUNE, SIROLIMUS
RAZADYNE ER, GALANTAMINE HYDROBROMIDE
RAZADYNE, GALANTAMINE HYDROBROMIDE
REBETOL, RIBAVIRIN
REFLUDAN, LEPIRUDIN RECOMBINANT
REGITINE, PHENTOLAMINE MESYLATE
REGLAN, METOCLOPRAMIDE HYDROCHLORIDE
REGONOL, PYRIDOSTIGMINE BROMIDE
RELENZA, ZANAMIVIR
RELPAZ, ELETRIPTAN HYDROBROMIDE
REMERON SOLTAB, MIRTAZAPINE
REMERON, MIRTAZAPINE
REMODULIN, TREPROSTINIL SODIUM
RENACIDIN, CITRIC ACID
RENAGEL, SEVELAMER HYDROCHLORIDE
RENAMIN W/O ELECTROLYTES, AMINO ACIDS
RENESE, POLYTHIAZIDE
RENESE-R, POLYTHIAZIDE

APPENDIX A - PRODUCT NAME INDEX

A - 47

**** R ****

RENO-30, DIATRIZOATE MEGLUMINE
RENO-60, DIATRIZOATE MEGLUMINE
RENOCAL-76, DIATRIZOATE MEGLUMINE
RENO-DIP, DIATRIZOATE MEGLUMINE
RENOGRAFIN-60, DIATRIZOATE MEGLUMINE
RENOGRAFIN-76, DIATRIZOATE MEGLUMINE
RENOVA, TRETINOIN
REPRONEX, LUTEINIZING HORMONE
REQUIP, ROPINIROLE HYDROCHLORIDE
RESCRIPTOR, DELAVIRDINE MESYLATE
RESECTISOL IN PLASTIC CONTAINER, MANNITOL
RESERPINE, RESERPINE
RESTASIS, CYCLOSPORINE
RESTORIL, TEMAZEPAM
RETIN-A MICRO, TRETINOIN
RETIN-A, TRETINOIN
RETISERT, FLUOCINOLONE ACETONIDE
RETROVIR, ZIDOVUDINE
REVATIO, SILDENAFIL CITRATE
REVEX, NALMEFENE HYDROCHLORIDE
REV-EYES, DAPIPRAZOLE HYDROCHLORIDE
REVIA, NALTREXONE HYDROCHLORIDE
REVLIMID, LENALIDOMIDE
REYATAZ, ATAZANAVIR SULFATE
R-GENE 10, ARGININE HYDROCHLORIDE
RHINOCORT, BUDESONIDE
RIBASPHERE, RIBAVIRIN
RIBAVIRIN, RIBAVIRIN
RID MOUSSE, PIPERONYL BUTOXIDE (OTC)
RIDAURA, AURANOFIN
RIFADIN, RIFAMPIN
RIFAMATE, ISONIAZID
RIFAMPIN AND ISONIAZID, ISONIAZID
RIFAMPIN, RIFAMPIN
RIFATER, ISONIAZID
RILUTEK, RILUZOLE
RILUZOLE, RILUZOLE
RIMACTANE, RIFAMPIN
RIMANTADINE HYDROCHLORIDE, RIMANTADINE HYDROCHLORIDE
RIMSO-50, DIMETHYL SULFOXIDE
RINGER'S IN PLASTIC CONTAINER, CALCIUM CHLORIDE
RIOMET, METFORMIN HYDROCHLORIDE
RISPERDAL CONSTA, RISPERIDONE
RISPERDAL, RISPERIDONE
RITALIN LA, METHYLPHENIDATE HYDROCHLORIDE
RITALIN, METHYLPHENIDATE HYDROCHLORIDE
RITALIN-SR, METHYLPHENIDATE HYDROCHLORIDE
RITODRINE HYDROCHLORIDE IN DEXTROSE 5% IN PLASTIC CONTAINER, RITODRINE HYDROCHLORIDE
RITODRINE HYDROCHLORIDE, RITODRINE HYDROCHLORIDE
ROBAXIN, METHOCARBAMOL
ROBAXIN-750, METHOCARBAMOL
ROBINUL FORTE, GLYCOPYRROLATE
ROBINUL, GLYCOPYRROLATE
ROCALTROL, CALCITRIOL
ROCEPHIN W/ DEXTROSE IN PLASTIC CONTAINER, CEFTRIAXONE SODIUM
ROCEPHIN, CEFTRIAXONE SODIUM
ROGAINE (FOR MEN), MINOXIDIL (OTC)
ROGAINE (FOR WOMEN), MINOXIDIL (OTC)
ROGAINE EXTRA STRENGTH (FOR MEN), MINOXIDIL (OTC)
ROMAZICON, FLUMAZENIL
ROWASA, MESALAMINE
ROXICET 5/500, ACETAMINOPHEN
ROXICET, ACETAMINOPHEN

APPENDIX A - PRODUCT NAME INDEX

A - 48

**** R ****

ROXICODONE, OXYCODONE HYDROCHLORIDE
ROXILOX, ACETAMINOPHEN
ROZEREM, RAMELTEON
RUBEX, DOXORUBICIN HYDROCHLORIDE
RUBRATOPE-57, CYANOCOBALAMIN, CO-57
RYTHMOL SR, PROPAFENONE HYDROCHLORIDE
RYTHMOL, PROPAFENONE HYDROCHLORIDE

**** S ****

SAIZEN, SOMATROPIN RECOMBINANT
SALAGEN, PILOCARPINE HYDROCHLORIDE
SALURON, HYDROFLUMETHIAZIDE
SANCTURA, TROSPIMUM CHLORIDE
SANDIMMUNE, CYCLOSPORINE
SANDOSTATIN LAR, OCTREOTIDE ACETATE
SANDOSTATIN, OCTREOTIDE ACETATE
SARAFEM, FLUOXETINE HYDROCHLORIDE
SCANDONEST L, LEVONORDEFIN
SCANDONEST PLAIN, MEPIVACAINE HYDROCHLORIDE
SCLEROSOL, TALC
SEASONALE, ETHINYL ESTRADIOL
SEASONIQUE, ETHINYL ESTRADIOL
SECOBARBITAL SODIUM, SECOBARBITAL SODIUM
SECONAL SODIUM, SECOBARBITAL SODIUM
SECREFO, SECRETIN SYNTHETIC PORCINE
SECTRAL, ACEBUTOLOL HYDROCHLORIDE
SELEGILINE HYDROCHLORIDE, SELEGILINE HYDROCHLORIDE
SELENIUM SULFIDE, SELENIUM SULFIDE
SELSUN, SELENIUM SULFIDE
SEMPREX-D, ACRIVASTINE
SENSIPAR, CINACALCET HYDROCHLORIDE
SENSORCAINE, BUPIVACAINE HYDROCHLORIDE
SEPTOCAINE, ARTICAIN HYDROCHLORIDE
SEPTRA DS, SULFAMETHOXAZOLE
SEPTRA GRAPE, SULFAMETHOXAZOLE
SEPTRA, SULFAMETHOXAZOLE
SEREVENT, SALMETEROL XINAFOATE
SEROMYCIN, CYCLOSERINE
SEROPHENE, CLOMIPHENE CITRATE
SEROQUEL, QUETIAPINE FUMARATE
SEROSTIM, SOMATROPIN RECOMBINANT
SERPALAN, RESERPINE
SERTRALINE HYDROCHLORIDE, SERTRALINE HYDROCHLORIDE
SEVOFLURANE, SEVOFLURANE
SHADE UVAGUARD, AVOBENZONE (OTC)
SILVADENE, SILVER SULFADIAZINE
SIMVASTATIN, SIMVASTATIN
SINE-AID IB, IBUPROFEN (OTC)
SINEMET CR, CARBIDOPA
SINEMET, CARBIDOPA
SINEQUAN, DOXEPIN HYDROCHLORIDE
SINGULAIR, MONTELUKAST SODIUM
SINOGRAPHIN, DIATRIZOATE MEGLUMINE
SKELAXIN, METAXALONE
SKELID, TILUDRONATE DISODIUM
SODIUM ACETATE IN PLASTIC CONTAINER, SODIUM ACETATE, ANHYDROUS
SODIUM BICARBONATE, SODIUM BICARBONATE
SODIUM BUTABARBITAL, BUTABARBITAL SODIUM
SODIUM CHLORIDE 0.45% AND POTASSIUM CHLORIDE 0.15% IN PLASTIC CONTAINER, POTASSIUM CHLORIDE
SODIUM CHLORIDE 0.45% IN PLASTIC CONTAINER, SODIUM CHLORIDE
SODIUM CHLORIDE 0.9% AND POTASSIUM CHLORIDE 0.15% IN PLASTIC CONTAINER, POTASSIUM CHLORIDE
SODIUM CHLORIDE 0.9% AND POTASSIUM CHLORIDE 0.224%, POTASSIUM CHLORIDE
SODIUM CHLORIDE 0.9% AND POTASSIUM CHLORIDE 0.3% IN PLASTIC CONTAINER, POTASSIUM CHLORIDE

APPENDIX A - PRODUCT NAME INDEX

A - 49

**** S ****

SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER, SODIUM CHLORIDE
SODIUM CHLORIDE 0.9% IN STERILE PLASTIC CONTAINER, SODIUM CHLORIDE
SODIUM CHLORIDE 3% IN PLASTIC CONTAINER, SODIUM CHLORIDE
SODIUM CHLORIDE 5% IN PLASTIC CONTAINER, SODIUM CHLORIDE
SODIUM CHLORIDE IN PLASTIC CONTAINER, SODIUM CHLORIDE
SODIUM CHROMATE CR 51, SODIUM CHROMATE, CR-51
SODIUM IODIDE I 123, SODIUM IODIDE, I-123
SODIUM IODIDE I 131, KIT, SODIUM IODIDE, I-131
SODIUM IODIDE I 131, SODIUM IODIDE, I-131
SODIUM LACTATE 0.167 MOLAR IN PLASTIC CONTAINER, SODIUM LACTATE
SODIUM LACTATE 1/6 MOLAR IN PLASTIC CONTAINER, SODIUM LACTATE
SODIUM LACTATE IN PLASTIC CONTAINER, SODIUM LACTATE
SODIUM NITROPRUSSIDE, SODIUM NITROPRUSSIDE
SODIUM P.A.S., AMINOSALICYLATE SODIUM
SODIUM PHOSPHATE P 32, SODIUM PHOSPHATE, P-32
SODIUM PHOSPHATES IN PLASTIC CONTAINER, SODIUM PHOSPHATE, DIBASIC, HEPTAHYDRATE
SODIUM POLYSTYRENE SULFONATE, SODIUM POLYSTYRENE SULFONATE
SOLAGE, MEQUINOL
SOLARAZE, DICLOFENAC SODIUM
SOLODYN, MINOCYCLINE HYDROCHLORIDE
SOLTAMOX, TAMOXIFEN CITRATE
SOLU-CORTEF, HYDROCORTISONE SODIUM SUCCINATE
SOLU-MEDROL, METHYLPREDNISOLONE SODIUM SUCCINATE
SOMA COMPOUND W/ CODEINE, ASPIRIN
SOMA COMPOUND, ASPIRIN
SOMA, CARISOPRODOL
SOMAVERT, PEGVISOMANT
SONATA, ZALEPLON
SONAZINE, CHLORPROMAZINE HYDROCHLORIDE
SORBITOL 3% IN PLASTIC CONTAINER, SORBITOL
SORBITOL 3.3% IN PLASTIC CONTAINER, SORBITOL
SORBITOL-MANNITOL IN PLASTIC CONTAINER, MANNITOL
SORIATANE, ACITRETIN
SORINE, SOTALOL HYDROCHLORIDE
SOTALOL HYDROCHLORIDE, SOTALOL HYDROCHLORIDE
SOTRADECOL, SODIUM TETRADECYL SULFATE
SOTRET, ISOTRETINOIN
SPECTAZOLE, ECONAZOLE NITRATE
SPECTRACEF, CEFditoren PIVOXIL
SPIRIVA, TIOTROPIUM BROMIDE MONOHYDRATE
SPIRONOLACTONE AND HYDROCHLOROTHIAZIDE, HYDROCHLOROTHIAZIDE
SPIRONOLACTONE, SPIRONOLACTONE
SPORANOX, ITRACONAZOLE
SPRINTEC, ETHINYL ESTRADIOL
SPRYCEL, DASATINIB
SPS, SODIUM POLYSTYRENE SULFONATE
SSD AF, SILVER SULFADIAZINE
SSD, SILVER SULFADIAZINE
STADOL PRESERVATIVE FREE, BUTORPHANOL TARTRATE
STADOL, BUTORPHANOL TARTRATE
STALEVO 100, CARBIDOPA
STALEVO 150, CARBIDOPA
STALEVO 50, CARBIDOPA
STARLIX, NATEGLINIDE
STELAZINE, TRIFLUOPERAZINE HYDROCHLORIDE
STERILE WATER FOR INJECTION IN PLASTIC CONTAINER, WATER FOR INJECTION, STERILE
STERILE WATER IN PLASTIC CONTAINER, WATER FOR IRRIGATION, STERILE
STERILE WATER, WATER FOR IRRIGATION, STERILE
STERI-STAT, CHLORHEXIDINE GLUCONATE (OTC)
STIE-CORT, HYDROCORTISONE
STIMATE, DESMOPRESSIN ACETATE
STRATTERA, ATOMOXETINE HYDROCHLORIDE
STREPTOMYCIN SULFATE, STREPTOMYCIN SULFATE

APPENDIX A - PRODUCT NAME INDEX

A - 50

**** S ****

STRIANT, TESTOSTERONE
STRIFON FORTE DSC, CHLORZOXAZONE
STROMEKTOL, IVERMECTIN
STRONTIUM CHLORIDE SR-89, STRONTIUM CHLORIDE, SR-89
SUBLIMAZE PRESERVATIVE FREE, FENTANYL CITRATE
SUBOXONE, BUPRENORPHINE HYDROCHLORIDE
SUBUTEX, BUPRENORPHINE HYDROCHLORIDE
SUCRAID, SACROSIDASE
SUCRALFATE, SUCRALFATE
SUDAFED 12 HOUR, PSEUDOEPHEDRINE HYDROCHLORIDE (OTC)
SUFENTA, SUFENTANIL CITRATE
SUFENTANIL CITRATE, SUFENTANIL CITRATE
SULAR, NISOLDIPINE
SULF-10, SULFACETAMIDE SODIUM
SULFACEL-15, SULFACETAMIDE SODIUM
SULFACETAMIDE SODIUM AND PREDNISOLONE SODIUM PHOSPHATE, PREDNISOLONE SODIUM PHOSPHATE
SULFACETAMIDE SODIUM, SULFACETAMIDE SODIUM
SULFADIAZINE, SULFADIAZINE
SULFAMETHOPRIM, SULFAMETHOXAZOLE
SULFAMETHOXAZOLE AND TRIMETHOPRIM DOUBLE STRENGTH, SULFAMETHOXAZOLE
SULFAMETHOXAZOLE AND TRIMETHOPRIM SINGLE STRENGTH, SULFAMETHOXAZOLE
SULFAMETHOXAZOLE AND TRIMETHOPRIM, SULFAMETHOXAZOLE
SULFAMYLON, MAFENIDE ACETATE
SULFASALAZINE, SULFASALAZINE
SULFATRIM PEDIATRIC, SULFAMETHOXAZOLE
SULFATRIM, SULFAMETHOXAZOLE
SUFENTANIL CITRATE, SUFENTANIL CITRATE
SULFINPYRAZONE, SULFINPYRAZONE
SULFISOXAZOLE, SULFISOXAZOLE
SULINDAC, SULINDAC
SUMYCIN, TETRACYCLINE HYDROCHLORIDE
SUPRANE, DESFLURANE
SUPRAX, CEFIXIME
SURMONTIL, TRIMIPRAMINE MALEATE
SURVANTA, BERACTANT
SUSTIVA, EFAVIRENZ
SUTENT, SUNITINIB MALATE
SYMBICORT, BUDESONIDE
SYMBYAX, FLUOXETINE HYDROCHLORIDE
SYMLIN, PRAMLINTIDE ACETATE
SYMMETREL, AMANTADINE HYDROCHLORIDE
SYNACORT, HYDROCORTISONE
SYNALAR, FLUOCINOLONE ACETONIDE
SYNALGOS-DC, ASPIRIN
SYNAREL, NAFARELIN ACETATE
SYNERA, LIDOCAINE
SYNERCID, DALFOPRISTIN
SYNTHROID, LEVOTHYROXINE SODIUM
SYPRINE, TRIENTINE HYDROCHLORIDE

**** T ****

TAB-PROFEN, IBUPROFEN (OTC)
TACLONEX, BETAMETHASONE DIPROPIONATE
TAGAMET HB, CIMETIDINE (OTC)
TAGAMET, CIMETIDINE
TALACEN, ACETAMINOPHEN
TALC, TALC
TALWIN NX, NALOXONE HYDROCHLORIDE
TALWIN, PENTAZOCINE LACTATE
TAMBOCOR, FLECAINIDE ACETATE
TAMIFLU, OSELTAMIVIR PHOSPHATE
TAMOXIFEN CITRATE, TAMOXIFEN CITRATE
TAPAZOLE, METHIMAZOLE

APPENDIX A - PRODUCT NAME INDEX

A - 51

**** T ****

TARCEVA, ERLOTINIB HYDROCHLORIDE
TARGRETIN, BEXAROTENE
TARKA, TRANDOLAPRIL
TASMAR, TOLCAPONE
TAVIST ALLERGY/SINUS/HEADACHE, ACETAMINOPHEN (OTC)
TAVIST-1, CLEMASTINE FUMARATE (OTC)
TAXOL, PACLITAXEL
TAXOTERE, DOCETAXEL
TAZICEF, CEFTAZIDIME
TAZORAC, TAZAROTENE
TAZTIA XT, DILTIAZEM HYDROCHLORIDE
TECHNECOLL, TECHNETIUM TC-99M SULFUR COLLOID KIT
TECHNELITE, TECHNETIUM TC-99M SODIUM PERTECHNETATE GENERATOR
TECHNISCAN GLUCEPTATE, TECHNETIUM TC-99M GLUCEPTATE KIT
TECHNISCAN MAA, TECHNETIUM TC-99M ALBUMIN AGGREGATED KIT
TECHNISCAN MAG3, TECHNETIUM TC-99M MERTIATIDE KIT
TECHNISCAN PYP KIT, TECHNETIUM TC-99M PYROPHOSPHATE KIT
TECHNISCAN, TECHNETIUM TC-99M OXIDRONATE KIT
TECHNETIUM TC 99M ALBUMIN AGGREGATED KIT, TECHNETIUM TC-99M ALBUMIN AGGREGATED KIT
TECHNETIUM TC 99M MPI MDP, TECHNETIUM TC-99M MEDRONATE KIT
TECHNETIUM TC-99M PENTETATE KIT, TECHNETIUM TC-99M PENTETATE KIT
TEGRETOL, CARBAMAZEPINE
TEGRETOL-XR, CARBAMAZEPINE
TEMAZEPAM, TEMAZEPAM
TEMODAR, TEMOZOLOMIDE
TEMOVATE E, CLOBETASOL PROPIONATE
TEMOVATE, CLOBETASOL PROPIONATE
TENCON, ACETAMINOPHEN
TENEX, GUANFACINE HYDROCHLORIDE
TENORETIC 100, ATENOLOL
TENORETIC 50, ATENOLOL
TENORMIN, ATENOLOL
TENSILON PRESERVATIVE FREE, EDROPHONIUM CHLORIDE
TENSILON, EDROPHONIUM CHLORIDE
TENUATE DOSPAN, DIETHYLPROPION HYDROCHLORIDE
TENUATE, DIETHYLPROPION HYDROCHLORIDE
TERAZOL 3, TERCONAZOLE
TERAZOL 7, TERCONAZOLE
TERAZOSIN HYDROCHLORIDE, TERAZOSIN HYDROCHLORIDE
TERBUTALINE SULFATE, TERBUTALINE SULFATE
TERCONAZOLE, TERCONAZOLE
TERIL, CARBAMAZEPINE
TERRA-CORTRIL, HYDROCORTISONE ACETATE
TERRAMYCIN W/ POLYMYXIN B SULFATE, OXYTETRACYCLINE HYDROCHLORIDE
TERRAMYCIN, LIDOCAINE HYDROCHLORIDE
TERRAMYCIN, OXYTETRACYCLINE HYDROCHLORIDE
TESLAC, TESTOLACTONE
TESSALON, BENZONATATE
TESTIM, TESTOSTERONE
TESTOSTERONE CYPIONATE, TESTOSTERONE CYPIONATE
TESTOSTERONE ENANTHATE, TESTOSTERONE ENANTHATE
TESTOSTERONE ETHANATE, TESTOSTERONE ENANTHATE
TESTOSTERONE PROPIONATE, TESTOSTERONE PROPIONATE
TESTOSTERONE, TESTOSTERONE
TESTRED, METHYLTESTOSTERONE
TETRACYCLINE HYDROCHLORIDE, TETRACYCLINE HYDROCHLORIDE
TETREX, TETRACYCLINE PHOSPHATE COMPLEX
TEVETEN HCT, EPROSARTAN MESYLATE
TEVETEN, EPROSARTAN MESYLATE
TEV-TROPIN, SOMATROPIN RECOMBINANT
TEXACORT, HYDROCORTISONE
THALITONE, CHLORTHALIDONE
THALLOUS CHLORIDE TL 201, THALLOUS CHLORIDE, TL-201

APPENDIX A - PRODUCT NAME INDEX

A - 52

**** T ****

THALOMID, THALIDOMIDE
THAM, TROMETHAMINE
THEO-24, THEOPHYLLINE
THEOCHRON, THEOPHYLLINE
THEOLAIR, THEOPHYLLINE
THEOPHYLLINE 0.04% AND DEXTROSE 5% IN PLASTIC CONTAINER, THEOPHYLLINE
THEOPHYLLINE 0.08% AND DEXTROSE 5% IN PLASTIC CONTAINER, THEOPHYLLINE
THEOPHYLLINE 0.16% AND DEXTROSE 5% IN PLASTIC CONTAINER, THEOPHYLLINE
THEOPHYLLINE 0.32% AND DEXTROSE 5% IN PLASTIC CONTAINER, THEOPHYLLINE
THEOPHYLLINE AND DEXTROSE 5% IN PLASTIC CONTAINER, THEOPHYLLINE
THEOPHYLLINE IN DEXTROSE 5% IN PLASTIC CONTAINER, THEOPHYLLINE
THEOPHYLLINE, THEOPHYLLINE
THERMAZENE, SILVER SULFADIAZINE
THEROXIDIL, MINOXIDIL (OTC)
THIAMINE HYDROCHLORIDE, THIAMINE HYDROCHLORIDE
THIOGUANINE, THIOGUANINE
THIORIDAZINE HYDROCHLORIDE INTENSOL, THIORIDAZINE HYDROCHLORIDE
THIORIDAZINE HYDROCHLORIDE, THIORIDAZINE HYDROCHLORIDE
THIOTEPA, THIOTEPA
THIOTHIXENE HYDROCHLORIDE, THIOTHIXENE HYDROCHLORIDE
THIOTHIXENE, THIOTHIXENE
THORAZINE, CHLORPROMAZINE HYDROCHLORIDE
THYROGEN, THYROTROPIN ALFA
THYROLAR-0.25, LIOETHYRONINE SODIUM
THYROLAR-0.5, LIOETHYRONINE SODIUM
THYROLAR-1, LIOETHYRONINE SODIUM
THYROLAR-2, LIOETHYRONINE SODIUM
THYROLAR-3, LIOETHYRONINE SODIUM
THYROSAFE, POTASSIUM IODIDE (OTC)
THYROSHIELD, POTASSIUM IODIDE (OTC)
TIAZAC, DILTIAZEM HYDROCHLORIDE
TICLID, TICLOPIDINE HYDROCHLORIDE
TICLOPIDINE HYDROCHLORIDE, TICLOPIDINE HYDROCHLORIDE
TIGAN, TRIMETHOBENZAMIDE HYDROCHLORIDE
TIKOSYN, DOFETILIDE
TILADE, NEDOCROMIL SODIUM
TIMENTIN IN PLASTIC CONTAINER, CLAVULANATE POTASSIUM
TIMENTIN, CLAVULANATE POTASSIUM
TIMOLOL MALEATE, TIMOLOL MALEATE
TIMOPTIC IN OCUDOSE, TIMOLOL MALEATE
TIMOPTIC, TIMOLOL MALEATE
TIMOPTIC-XE, TIMOLOL MALEATE
TINDAMAX, TINIDAZOLE
TIOCONAZOLE, TIOCONAZOLE (OTC)
TIOPRONIN, TIOPRONIN
TIROSINT, LEVOTHYROXINE SODIUM
TIS-U-SOL IN PLASTIC CONTAINER, MAGNESIUM SULFATE
TIS-U-SOL, MAGNESIUM SULFATE
TIZANIDINE HYDROCHLORIDE, TIZANIDINE HYDROCHLORIDE
TOBI, TOBRAMYCIN
TOBRADEX, DEXAMETHASONE
TOBRAMYCIN (PHARMACY BULK), TOBRAMYCIN SULFATE
TOBRAMYCIN AND DEXAMETHASONE, DEXAMETHASONE
TOBRAMYCIN SULFATE IN SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER, TOBRAMYCIN SULFATE
TOBRAMYCIN SULFATE, TOBRAMYCIN SULFATE
TOBRAMYCIN, TOBRAMYCIN
TOBRAMYCIN, TOBRAMYCIN SULFATE
TOBRASONE, FLUOROMETHOLONE ACETATE
TOBREX, TOBRAMYCIN
TOFRANIL, IMIPRAMINE HYDROCHLORIDE
TOFRANIL-PM, IMIPRAMINE PAMOATE
TOLAZAMIDE, TOLAZAMIDE
TOLBUTAMIDE, TOLBUTAMIDE

APPENDIX A - PRODUCT NAME INDEX

A - 53

**** T ****

TOLECTIN 600, TOLMETIN SODIUM
TOLECTIN DS, TOLMETIN SODIUM
TOLECTIN, TOLMETIN SODIUM
TOLINASE, TOLAZAMIDE
TOLMETIN SODIUM, TOLMETIN SODIUM
TOPAMAX SPRINKLE, TOPIRAMATE
TOPAMAX, TOPIRAMATE
TOPICORT LP, DESOXIMETASONE
TOPICORT, DESOXIMETASONE
TOPIRAMATE, TOPIRAMATE
TOPROL-XL, METOPROLOL SUCCINATE
TORADOL, KETOROLAC TROMETHAMINE
TORECAN, THIETHYLPERAZINE MALEATE
TORSEMIDE, TORSEMIDE
TPN ELECTROLYTES IN PLASTIC CONTAINER, CALCIUM CHLORIDE
TRACLEER, BOSENTAN
TRACRIUM PRESERVATIVE FREE, ATRACURIUM BESYLATE
TRACRIUM, ATRACURIUM BESYLATE
TRAMADOL HYDROCHLORIDE AND ACETAMINOPHEN, ACETAMINOPHEN
TRAMADOL HYDROCHLORIDE, TRAMADOL HYDROCHLORIDE
TRANDATE, LABETALOL HYDROCHLORIDE
TRANDOLAPRIL, TRANDOLAPRIL
TRANSDERM SCOP, SCOPOLAMINE
TRANXENE SD, CLORAZEPATE DIPOTASSIUM
TRANXENE, CLORAZEPATE DIPOTASSIUM
TRANLYCYPROMINE SULFATE, TRANLYCYPROMINE SULFATE
TRASYLOL, APROTININ BOVINE
TRAVASOL 10% IN PLASTIC CONTAINER, AMINO ACIDS
TRAVASOL 10% W/O ELECTROLYTES, AMINO ACIDS
TRAVASOL 3.5% SULFITE FREE W/ ELECTROLYTES IN PLASTIC CONTAINER, AMINO ACIDS
TRAVASOL 3.5% W/ ELECTROLYTES, AMINO ACIDS
TRAVASOL 5.5% IN PLASTIC CONTAINER, AMINO ACIDS
TRAVASOL 5.5% SULFITE FREE W/ ELECTROLYTES IN PLASTIC CONTAINER, AMINO ACIDS
TRAVASOL 5.5% W/ ELECTROLYTES, AMINO ACIDS
TRAVASOL 5.5% W/O ELECTROLYTES, AMINO ACIDS
TRAVASOL 8.5% IN PLASTIC CONTAINER, AMINO ACIDS
TRAVASOL 8.5% SULFITE FREE W/ ELECTROLYTES IN PLASTIC CONTAINER, AMINO ACIDS
TRAVASOL 8.5% W/ ELECTROLYTES, AMINO ACIDS
TRAVASOL 8.5% W/O ELECTROLYTES, AMINO ACIDS
TRAVATAN Z, TRAVOPROST
TRAVATAN, TRAVOPROST
TRAZODONE HYDROCHLORIDE, TRAZODONE HYDROCHLORIDE
TRECATOR-SC, ETHIONAMIDE
TRELSTAR DEPOT, TRIPTORELIN PAMOATE
TRELSTAR, TRIPTORELIN PAMOATE
TRENTAL, PENTOXIFYLLINE
TRETINOIN, TRETINOIN
TREXALL, METHOTREXATE SODIUM
TRIAcet, TRIAMCINOLONE ACETONIDE
TRIAcin-C, CODEINE PHOSPHATE
TRIAMCINOLONE ACETONIDE IN ABSORBASE, TRIAMCINOLONE ACETONIDE
TRIAMCINOLONE ACETONIDE, TRIAMCINOLONE ACETONIDE
TRIAMTERENE AND HYDROCHLOROTHIAZIDE, HYDROCHLOROTHIAZIDE
TRIAZOLAM, TRIAZOLAM
TRICOR, FENOFIBRATE
TRIDERM, TRIAMCINOLONE ACETONIDE
TRIDESILON, DESONIDE
TRIDIONE, TRIMETHADIONE
TRIFLUOPERAZINE HYDROCHLORIDE, TRIFLUOPERAZINE HYDROCHLORIDE
TRIFLURIDINE, TRIFLURIDINE
TRIGLIDE, FENOFIBRATE
TRIHENXYPHENIDYL HYDROCHLORIDE, TRIHENXYPHENIDYL HYDROCHLORIDE
TRILEPTAL, OXCARBAZEPINE

APPENDIX A - PRODUCT NAME INDEX

A - 54

**** T ****

TRI-LUMA, FLUOCINOLONE ACETONIDE
TRILYTE, POLYETHYLENE GLYCOL 3350
TRIMETHOBENZAMIDE HYDROCHLORIDE, TRIMETHOBENZAMIDE HYDROCHLORIDE
TRIMETHOPRIM SULFATE AND POLYMYXIN B SULFATE, POLYMYXIN B SULFATE
TRIMETHOPRIM, TRIMETHOPRIM
TRIMIPRAMINE MALEATE, TRIMIPRAMINE MALEATE
TRIMOX, AMOXICILLIN
TRI-NORINYL 28-DAY, ETHINYL ESTRADIOL
TRIOSTAT, LIOTHYRONINE SODIUM
TRIPHASIL-28, ETHINYL ESTRADIOL
TRI-PREVIFEM, ETHINYL ESTRADIOL
TRIPROLIDINE HCL, PSEUDOEPHEDRINE HCL AND CODEINE PHOSPHATE, CODEINE PHOSPHATE
TRISENOX, ARSENIC TRIOXIDE
TRI-SPRINTEC, ETHINYL ESTRADIOL
TRIVAGIZOLE 3, CLOTRIMAZOLE (OTC)
TRIVORA-28, ETHINYL ESTRADIOL
TRIZIVIR, ABACAVIR SULFATE
TROBICIN, SPECTINOMYCIN HYDROCHLORIDE
TROPHAMINE 10%, AMINO ACIDS
TROPHAMINE, AMINO ACIDS
TROPICACYL, TROPICAMIDE
TROPICAMIDE, TROPICAMIDE
TRUPHYLLINE, AMINOPHYLLINE
TRUSOPT, DORZOLAMIDE HYDROCHLORIDE
TRUVADA, EMTRICITABINE
TUSSIGON, HOMATROPINE METHYLBROMIDE
TUSSIONEX PENNKINETIC, CHLORPHENIRAMINE POLISTIREX
TWINJECT 0.3, EPINEPHRINE
TWINJECT, EPINEPHRINE
TYGACIL, TIGECYCLINE
TYLENOL (CAPLET), ACETAMINOPHEN (OTC)
TYLENOL (GELTAB), ACETAMINOPHEN (OTC)
TYLENOL W/ CODEINE NO. 3, ACETAMINOPHEN
TYLOX, ACETAMINOPHEN
TYZEKA, TELBIVUDINE
TYZINE, TETRAHYDROZOLINE HYDROCHLORIDE

**** U ****

U-CORT, HYDROCORTISONE ACETATE
ULTANE, SEVOFLURANE
ULTIVA, REMIFENTANIL HYDROCHLORIDE
ULTRACET, ACETAMINOPHEN
ULTRAM, TRAMADOL HYDROCHLORIDE
ULTRATAG, TECHNETIUM TC-99M RED BLOOD CELL KIT
ULTRA-TECHNEKOW FM, TECHNETIUM TC-99M SODIUM PERTECHNETATE GENERATOR
ULTRAVATE, HALOBETASOL PROPIONATE
ULTRAVIST (PHARMACY BULK), IOPROMIDE
ULTRAVIST 150, IOPROMIDE
ULTRAVIST 240, IOPROMIDE
ULTRAVIST 300, IOPROMIDE
ULTRAVIST 370, IOPROMIDE
UNASYN, AMPICILLIN SODIUM
UNIPHYL, THEOPHYLLINE
UNIRETIC, HYDROCHLOROTHIAZIDE
UNISOM, DOXYLAMINE SUCCINATE (OTC)
UNITHROID, LEVOTHYROXINE SODIUM
UNIVASC, MOEXIPRIL HYDROCHLORIDE
URECHOLINE, BETHANECHOL CHLORIDE
UREX, METHENAMINE HIPPURATE
URISPAS, FLAVOXATE HYDROCHLORIDE
UROCIT-K, POTASSIUM CITRATE
UROLOGIC G IN PLASTIC CONTAINER, CITRIC ACID
UROXATRAL, ALFUZOSIN HYDROCHLORIDE

APPENDIX A - PRODUCT NAME INDEX

A - 55

**** U ****

URSO 250, URSODIOL
URSO FORTE, URSODIOL
URSODIOL, URSODIOL
UVADEX, METHOXSALEN

**** V ****

VAGIFEM, ESTRADIOL
VAGISTAT-1, TIOCONAZOLE (OTC)
VALCYTE, VALGANCICLOVIR HYDROCHLORIDE
VALIUM, DIAZEPAM
VALNAC, BETAMETHASONE VALERATE
VALPROATE SODIUM, VALPROATE SODIUM
VALPROIC ACID, VALPROIC ACID
VALSTAR PRESERVATIVE FREE, VALRUBICIN
VALTREX, VALACYCLOVIR HYDROCHLORIDE
VANCOCIN HYDROCHLORIDE IN PLASTIC CONTAINER, VANCOMYCIN HYDROCHLORIDE
VANCOCIN HYDROCHLORIDE, VANCOMYCIN HYDROCHLORIDE
VANCOMYCIN HYDROCHLORIDE, VANCOMYCIN HYDROCHLORIDE
VANDAZOLE, METRONIDAZOLE
VANIQA, EFLORNITHINE HYDROCHLORIDE
VANOS, FLUOCINONIDE
VANTAS, HISTRELIN ACETATE
VANTIN, CEFPODOXIME PROXETIL
VAPRISOL, CONIVAPTAN HYDROCHLORIDE
VASERETIC, ENALAPRIL MALEATE
VASOCIDIN, PREDNISOLONE SODIUM PHOSPHATE
VASOCON, NAPHAZOLINE HYDROCHLORIDE
VASOCON-A, ANTAZOLINE PHOSPHATE (OTC)
VASOTEC, ENALAPRIL MALEATE
VASOTEC, ENALAPRILAT
VECURONIUM BROMIDE, VECURONIUM BROMIDE
VEETIDS, PENICILLIN V POTASSIUM
VELCADE, BORTEZOMIB
VELIVET, DESOGESTREL
VELOSEF '125', CEPHRADINE
VELOSEF '250', CEPHRADINE
VENLAFAXINE HYDROCHLORIDE, VENLAFAXINE HYDROCHLORIDE
VENOFER, IRON SUCROSE
VENTAVIS, ILOPROST
VENTOLIN HFA, ALBUTEROL SULFATE
VENTOLIN, ALBUTEROL
VEPESID, ETOPOSIDE
VERAPAMIL HYDROCHLORIDE, VERAPAMIL HYDROCHLORIDE
VERDESO, DESONIDE
VEREGEN, KUNECATECHINS
VERELAN PM, VERAPAMIL HYDROCHLORIDE
VERELAN, VERAPAMIL HYDROCHLORIDE
VESANOID, TRETINOIN
VESICARE, SOLIFENACIN SUCCINATE
VEXOL, RIMEXOLONE
VFEND, VORICONAZOLE
VIADUR, LEUPROLIDE ACETATE
VIAGRA, SILDENAFIL CITRATE
VIBISONE, CYANOCOBALAMIN
VIBRAMYCIN, DOXYCYCLINE
VIBRAMYCIN, DOXYCYCLINE CALCIUM
VIBRAMYCIN, DOXYCYCLINE HYCLATE
VIBRA-TABS, DOXYCYCLINE HYCLATE
VICODIN ES, ACETAMINOPHEN
VICODIN HP, ACETAMINOPHEN
VICODIN, ACETAMINOPHEN
VICOPROFEN, HYDROCODONE BITARTRATE
VIDAZA, AZACITIDINE

APPENDIX A - PRODUCT NAME INDEX

A - 56

**** V ****

VIDEX EC, DIDANOSINE
 VIDEX, DIDANOSINE
 VIGAMOX, MOXIFLOXACIN HYDROCHLORIDE
 VINBLASTINE SULFATE, VINBLASTINE SULFATE
 VINCRISTINE SULFATE PFS, VINCRISTINE SULFATE
 VINCRISTINE SULFATE, VINCRISTINE SULFATE
 VINORELBINE TARTRATE, VINORELBINE TARTRATE
 VIRACEPT, NELFINAVIR MESYLATE
 VIRAMUNE, NEVIRAPINE
 VIRAZOLE, RIBAVIRIN
 VIREAD, TENOFOVIR DISOPROXIL FUMARATE
 VIRILON, METHYLTESTOSTERONE
 VIROPTIC, TRIFLURIDINE
 VISICOL, SODIUM PHOSPHATE, DIBASIC, ANHYDROUS
 VISINE L.R., OXYMETAZOLINE HYDROCHLORIDE (OTC)
 VISINE-A, NAPHAZOLINE HYDROCHLORIDE (OTC)
 VISIONBLUE, TRYPAN BLUE
 VISIPAQUE 270, IODIXANOL
 VISIPAQUE 320, IODIXANOL
 VISKEN, PINDOLOL
 VISTARIL, HYDROXYZINE HYDROCHLORIDE
 VISTARIL, HYDROXYZINE PAMOATE
 VISTIDE, CIDOFOVIR
 VISUDYNE, VERTEPORFIN
 VITAMIN A PALMITATE, VITAMIN A PALMITATE
 VITAMIN D, ERGOCALCIFEROL
 VITAMIN K1, PHYTONADIONE
 VITRASE, HYALURONIDASE
 VITRASERT, GANCICLOVIR
 VIVACTIL, PROTRIPTYLINE HYDROCHLORIDE
 VIVELLE, ESTRADIOL
 VIVELLE-DOT, ESTRADIOL
 VIVITROL, NALTREXONE
 VOLTAREN, DICLOFENAC SODIUM
 VOLTAREN-XR, DICLOFENAC SODIUM
 VOSOL HC, ACETIC ACID, GLACIAL
 VOSPIRE ER, ALBUTEROL SULFATE
 VUMON, TENIPOSIDE
 VUSION, MICONAZOLE NITRATE
 VYTORIN, EZETIMIBE

**** W ****

WARFARIN SODIUM, WARFARIN SODIUM
 WELCHOL, COLESEVELAM HYDROCHLORIDE
 WELLBUTRIN SR, BUPROPION HYDROCHLORIDE
 WELLBUTRIN XL, BUPROPION HYDROCHLORIDE
 WELLBUTRIN, BUPROPION HYDROCHLORIDE
 WESTCORT, HYDROCORTISONE VALERATE
 WYGESIC, ACETAMINOPHEN

**** X ****

XALATAN, LATANOPROST
 XANAX XR, ALPRAZOLAM
 XANAX, ALPRAZOLAM
 XELODA, CAPECITABINE
 XENICAL, ORLISTAT
 XENON XE 133, XENON, XE-133
 XIBROM, BROMFENAC SODIUM
 XIFAXAN, RIFAXIMIN
 XOLEGEL, KETOCONAZOLE
 XOPENEX HFA, LEVALBUTEROL TARTRATE
 XOPENEX, LEVALBUTEROL HYDROCHLORIDE

APPENDIX A - PRODUCT NAME INDEX

A - 57

**** X ****

X-TROZINE L.A., PHENDIMETRAZINE TARTRATE
X-TROZINE, PHENDIMETRAZINE TARTRATE
XYLOCAINE 4% PRESERVATIVE FREE, LIDOCAINE HYDROCHLORIDE
XYLOCAINE PRESERVATIVE FREE, LIDOCAINE HYDROCHLORIDE
XYLOCAINE VISCOUS, LIDOCAINE HYDROCHLORIDE
XYLOCAINE W/ EPINEPHRINE, EPINEPHRINE
XYLOCAINE, LIDOCAINE HYDROCHLORIDE
XYREM, SODIUM OXYBATE

**** Y ****

YASMIN, DROSPIRENONE
YAZ, DROSPIRENONE

**** Z ****

ZADITOR, KETOTIFEN FUMARATE (OTC)
ZANAFLEX, TIZANIDINE HYDROCHLORIDE
ZANOSAR, STREPTOZOCIN
ZANTAC 150, RANITIDINE HYDROCHLORIDE
ZANTAC 150, RANITIDINE HYDROCHLORIDE (OTC)
ZANTAC 25, RANITIDINE HYDROCHLORIDE
ZANTAC 300, RANITIDINE HYDROCHLORIDE
ZANTAC 75, RANITIDINE HYDROCHLORIDE (OTC)
ZANTAC IN PLASTIC CONTAINER, RANITIDINE HYDROCHLORIDE
ZANTAC, RANITIDINE HYDROCHLORIDE
ZARONTIN, ETHOSUXIMIDE
ZAROXOLYN, METOLAZONE
ZAVESCA, MIGLUSTAT
ZEBETA, BISOPROLOL FUMARATE
ZEGERID, MAGNESIUM HYDROXIDE
ZEGERID, OMEPRAZOLE
ZELAPAR, SELEGILINE HYDROCHLORIDE
ZELNORM, TEGASEROD MALEATE
ZEMPLAR, PARICALCITOL
ZEMURON, ROCURONIUM BROMIDE
ZERIT, STAVUDINE
ZESTORETIC, HYDROCHLOROTHIAZIDE
ZESTRIL, LISINAPRIL
ZETIA, EZETIMIBE
ZIAC, BISOPROLOL FUMARATE
ZIAGEN, ABACAVIR SULFATE
ZIANA, CLINDAMYCIN PHOSPHATE
ZIDOVUDINE, ZIDOVUDINE
ZINACEF IN PLASTIC CONTAINER, CEFUROXIME SODIUM
ZINACEF, CEFUROXIME SODIUM
ZINC CHLORIDE IN PLASTIC CONTAINER, ZINC CHLORIDE
ZINECARD, DEXRAZOXANE HYDROCHLORIDE
ZITHROMAX, AZITHROMYCIN
ZMAX, AZITHROMYCIN
ZOCOR, SIMVASTATIN
ZOFRAN IN PLASTIC CONTAINER, ONDANSETRON HYDROCHLORIDE
ZOFRAN ODT, ONDANSETRON
ZOFRAN PRESERVATIVE FREE, ONDANSETRON HYDROCHLORIDE
ZOFRAN, ONDANSETRON HYDROCHLORIDE
ZOLADEX, GOSERELIN ACETATE
ZOLINZA, VORINOSTAT
ZOLOFT, SERTRALINE HYDROCHLORIDE
ZOMETA, ZOLEDRONIC ACID
ZOMIG, ZOLMITRIPTAN
ZOMIG-ZMT, ZOLMITRIPTAN
ZONALON, DOXEPIN HYDROCHLORIDE
ZONEGRAN, ZONISAMIDE
ZONISAMIDE, ZONISAMIDE

APPENDIX A - PRODUCT NAME INDEX

A - 58

**** Z ****

ZORBTIVE, SOMATROPIN RECOMBINANT
ZOSYN IN PLASTIC CONTAINER, PIPERACILLIN SODIUM
ZOSYN, PIPERACILLIN SODIUM
ZOVIA 1/35E-21, ETHINYL ESTRADIOL
ZOVIA 1/35E-28, ETHINYL ESTRADIOL
ZOVIA 1/50E-21, ETHINYL ESTRADIOL
ZOVIA 1/50E-28, ETHINYL ESTRADIOL
ZOVIRAX, ACYCLOVIR
ZOVIRAX, ACYCLOVIR SODIUM
ZYBAN, BUPROPION HYDROCHLORIDE
ZYDONE, ACETAMINOPHEN
ZYFLO, ZILEUTON
ZYLET, LOTEPREDNOL ETABONATE
ZYLOPRIM, ALLOPURINOL
ZYMAR, GATIFLOXACIN
ZYPREXA ZYDIS, OLANZAPINE
ZYPREXA, OLANZAPINE
ZYRTEC, CETIRIZINE HYDROCHLORIDE
ZYRTEC-D 12 HOUR, CETIRIZINE HYDROCHLORIDE
ZYVOX, LINEZOLID

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

**** 3 ****

3M

- * 3M HEALTH CARE INC
AVAGARD, ALCOHOL (OTC)
DURAPREP, IODINE POVACRYLEX (OTC)
- * 3M PHARMACEUTICALS INC
ALDARA, IMIQUIMOD
CALCIUM DISODIUM VERSENATE, EDETATE CALCIUM DISODIUM
MAXAIR, PIRBUTEROL ACETATE
METROGEL-VAGINAL, METRONIDAZOLE
MINITRAN, NITROGLYCERIN
NORFLEX, ORPHENADRINE CITRATE
NORGESIC FORTE, ASPIRIN
NORGESIC, ASPIRIN
PROVENTIL-HFA, ALBUTEROL SULFATE
QVAR 40, BECLOMETHASONE DIPROPIONATE
QVAR 80, BECLOMETHASONE DIPROPIONATE
TAMBOCOR, FLECAINIDE ACETATE
THEOLAIR, THEOPHYLLINE

AAIPHARMA LLC

- * AAIPHARMA LLC
AZASAN, AZATHIOPRINE
BRETHINE, TERBUTALINE SULFATE
ETODOLAC, ETODOLAC
KETOCONAZOLE, KETOCONAZOLE

AB GENERICS

- * AB GENERICS LP
MORPHINE SULFATE, MORPHINE SULFATE

ABBOTT

- * ABBOTT LABORATORIES
AKINETON, BIPERIDEN HYDROCHLORIDE
BIAXIN XL, CLARITHROMYCIN
BIAXIN, CLARITHROMYCIN
CYCLOSPORINE, CYCLOSPORINE
DEPAKOTE ER, DIVALPROEX SODIUM
DEPAKOTE, DIVALPROEX SODIUM
GENGRAF, CYCLOSPORINE
KALETRA, LOPINAVIR
NIMBEX PRESERVATIVE FREE, CISATRACURIUM BESYLATE
NIMBEX, CISATRACURIUM BESYLATE
NORVIR, RITONAVIR
OMNICEF, CEFDINIR
SYNTHROID, LEVOTHYROXINE SODIUM
ULTANE, SEVOFLURANE
ULTIVA, REMIFENTANIL HYDROCHLORIDE
ZEMPLAR, PARICALCITOL
- * ABBOTT LABORATORIES HOSP PRODUCTS DIV
CALCIJEX, CALCITRIOL
- * ABBOTT LABORATORIES PHARMACEUTICAL PRODUCTS DIV
BIAXIN, CLARITHROMYCIN
DEPACON, VALPROATE SODIUM
DEPAKENE, VALPROIC ACID
DEPAKOTE, DIVALPROEX SODIUM
DILAUDID, HYDROMORPHONE HYDROCHLORIDE
DILAUDID-HP, HYDROMORPHONE HYDROCHLORIDE
E.E.S. 200, ERYTHROMYCIN ETHYLSUCCINATE
E.E.S. 400, ERYTHROMYCIN ETHYLSUCCINATE
E.E.S., ERYTHROMYCIN ETHYLSUCCINATE
E-MYCIN, ERYTHROMYCIN
ENDURON, METHYCLOTHIAZIDE
ERYDERM, ERYTHROMYCIN
ERYPED, ERYTHROMYCIN ETHYLSUCCINATE
ERY-TAB, ERYTHROMYCIN
ERYTHROCIN STEARATE, ERYTHROMYCIN STEARATE

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

** A **

* ABBOTT LABORATORIES PHARMACEUTICAL PRODUCTS DIV

ERYTHROMYCIN, ERYTHROMYCIN
HYTRIN, TERAZOSIN HYDROCHLORIDE
K-TAB, POTASSIUM CHLORIDE
MAVIK, TRANDOLAPRIL
MERIDIA, SIBUTRAMINE HYDROCHLORIDE
NORVIR, RITONAVIR
ORETIC, HYDROCHLOROTHIAZIDE
PCE, ERYTHROMYCIN
POTASSIUM CHLORIDE, POTASSIUM CHLORIDE
PROSOM, ESTAZOLAM
TARKA, TRANDOLAPRIL
TRICOR, FENOFIBRATE
TRIDIONE, TRIMETHADIONE
VICODIN ES, ACETAMINOPHEN
VICODIN HP, ACETAMINOPHEN
VICODIN, ACETAMINOPHEN
VICOPROFEN, HYDROCODONE BITARTRATE
ZEMPLAR, PARICALCITOL

ABRAXIS BIOSCIENCE

* ABRAXIS BIOSCIENCE INC

ABRAXANE, PACLITAXEL
DIPRIVAN, PROPOFOL
EMLA, LIDOCAINE
NAROPIN, ROPIVACAINE HYDROCHLORIDE MONOHYDRATE
NESACAINE, CHLOROPROCAINE HYDROCHLORIDE
NESACAINE-MPF, CHLOROPROCAINE HYDROCHLORIDE
XYLOCAINE 4% PRESERVATIVE FREE, LIDOCAINE HYDROCHLORIDE
XYLOCAINE PRESERVATIVE FREE, LIDOCAINE HYDROCHLORIDE
XYLOCAINE VISCOUS, LIDOCAINE HYDROCHLORIDE
XYLOCAINE W/ EPINEPHRINE, EPINEPHRINE
XYLOCAINE, LIDOCAINE HYDROCHLORIDE

ABRAXIS PHARM

* ABRAXIS PHARMACEUTICAL PRODUCTS

ACYCLOVIR SODIUM, ACYCLOVIR SODIUM
ADENOSINE, ADENOSINE
AMIODARONE HYDROCHLORIDE, AMIODARONE HYDROCHLORIDE
AZITHROMYCIN, AZITHROMYCIN
BACITRACIN, BACITRACIN
BACTERIOSTATIC SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER, SODIUM CHLORIDE
CALCITRIOL, CALCITRIOL
CARBOPLATIN, CARBOPLATIN
CEFAZOLIN SODIUM, CEFAZOLIN SODIUM
CEFOTAXIME, CEFOTAXIME SODIUM
CEFOXITIN, CEFOTAXIME SODIUM
CEFTRIAXONE, CEFTRIAXONE SODIUM
CEFUROXIME SODIUM, CEFUROXIME SODIUM
CEFUROXIME, CEFUROXIME SODIUM
CHLORAMPHENICOL SODIUM SUCCINATE, CHLORAMPHENICOL SODIUM SUCCINATE
CHORIONIC GONADOTROPIN, GONADOTROPIN, CHORIONIC
CIPROFLOXACIN, CIPROFLOXACIN
CISPLATIN, CISPLATIN
CLADRIBINE, CLADRIBINE
CLINDAMYCIN PHOSPHATE IN DEXTROSE 5%, CLINDAMYCIN PHOSPHATE
CYTARABINE, CYTARABINE
DACARBAZINE, DACARBAZINE
DAUNORUBICIN HYDROCHLORIDE, DAUNORUBICIN HYDROCHLORIDE
DEXAMETHASONE SODIUM PHOSPHATE, DEXAMETHASONE SODIUM PHOSPHATE
DIMENHYDRINATE, DIMENHYDRINATE
DIPHENHYDRAMINE HYDROCHLORIDE, DIPHENHYDRAMINE HYDROCHLORIDE
DIPYRIDAMOLE, DIPYRIDAMOLE
DOXORUBICIN HYDROCHLORIDE, DOXORUBICIN HYDROCHLORIDE
DOXY 100, DOXYCYCLINE HYCLATE

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

** A **

* ABRAXIS PHARMACEUTICAL PRODUCTS
 DOXY 200, DOXYCYCLINE HYCLATE
 ESMOLOL HYDROCHLORIDE, ESMOLOL HYDROCHLORIDE
 ETOPOSIDE, ETOPOSIDE
 FAMOTIDINE PRESERVATIVE FREE, FAMOTIDINE
 FAMOTIDINE, FAMOTIDINE
 FLOXURIDINE, FLOXURIDINE
 FLUCONAZOLE IN SODIUM CHLORIDE 0.9%, FLUCONAZOLE
 FLUMAZENIL, FLUMAZENIL
 FLUOROURACIL, FLUOROURACIL
 FLUPHENAZINE DECANOATE, FLUPHENAZINE DECANOATE
 FLUPHENAZINE HYDROCHLORIDE, FLUPHENAZINE HYDROCHLORIDE
 FOLIC ACID, FOLIC ACID
 FUROSEMIDE, FUROSEMIDE
 GENTAMICIN SULFATE, GENTAMICIN SULFATE
 HALOPERIDOL DECANOATE, HALOPERIDOL DECANOATE
 HALOPERIDOL, HALOPERIDOL LACTATE
 HEPARIN SODIUM IN PLASTIC CONTAINER, HEPARIN SODIUM
 HEPARIN SODIUM PRESERVATIVE FREE, HEPARIN SODIUM
 HEPARIN SODIUM, HEPARIN SODIUM
 HYDRALAZINE HYDROCHLORIDE, HYDRALAZINE HYDROCHLORIDE
 HYDROXYZINE HYDROCHLORIDE, HYDROXYZINE HYDROCHLORIDE
 IFOSFAMIDE, IFOSFAMIDE
 IOPAMIDOL-250, IOPAMIDOL
 IOPAMIDOL-300, IOPAMIDOL
 IOPAMIDOL-370, IOPAMIDOL
 KANAMYCIN, KANAMYCIN SULFATE
 KETOROLAC TROMETHAMINE, KETOROLAC TROMETHAMINE
 LEUCOVORIN CALCIUM PRESERVATIVE FREE, LEUCOVORIN CALCIUM
 LIDOCAINE HYDROCHLORIDE IN PLASTIC CONTAINER, LIDOCAINE HYDROCHLORIDE
 LIDOCAINE HYDROCHLORIDE PRESERVATIVE FREE, LIDOCAINE HYDROCHLORIDE
 MAGNESIUM SULFATE, MAGNESIUM SULFATE
 MANNITOL 25%, MANNITOL
 MESNA, MESNA
 METARAMINOL BITARTRATE, METARAMINOL BITARTRATE
 METHOTREXATE, METHOTREXATE SODIUM
 METHYLPREDNISOLONE SODIUM SUCCINATE, METHYLPREDNISOLONE SODIUM SUCCINATE
 MIDAZOLAM HYDROCHLORIDE, MIDAZOLAM HYDROCHLORIDE
 MILRINONE LACTATE, MILRINONE LACTATE
 MITOXANTRONE, MITOXANTRONE HYDROCHLORIDE
 NEBUPENT, PENTAMIDINE ISETHIONATE
 OCTREOTIDE ACETATE (PRESERVATIVE FREE), OCTREOTIDE ACETATE
 OCTREOTIDE ACETATE, OCTREOTIDE ACETATE
 ONDANSETRON HYDROCHLORIDE PRESERVATIVE FREE, ONDANSETRON HYDROCHLORIDE
 ONDANSETRON HYDROCHLORIDE, ONDANSETRON HYDROCHLORIDE
 OXYTOCIN, OXYTOCIN
 PAMIDRONATE DISODIUM, PAMIDRONATE DISODIUM
 PENTAM, PENTAMIDINE ISETHIONATE
 PIPERACILLIN, PIPERACILLIN SODIUM
 POTASSIUM CHLORIDE IN PLASTIC CONTAINER, POTASSIUM CHLORIDE
 POTASSIUM CHLORIDE, POTASSIUM CHLORIDE
 PROGESTERONE, PROGESTERONE
 PROPRANOLOL, PROPRANOLOL HYDROCHLORIDE
 PROTAMINE SULFATE, PROTAMINE SULFATE
 PYRIDOXINE HYDROCHLORIDE, PYRIDOXINE HYDROCHLORIDE
 SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER, SODIUM CHLORIDE
 STERILE WATER FOR INJECTION IN PLASTIC CONTAINER, WATER FOR INJECTION, STERILE
 TERBUTALINE SULFATE, TERBUTALINE SULFATE
 THIAMINE HYDROCHLORIDE, THIAMINE HYDROCHLORIDE
 THIOTEPA, THIOTEPA
 TOBRAMYCIN (PHARMACY BULK), TOBRAMYCIN SULFATE
 TOBRAMYCIN, TOBRAMYCIN SULFATE
 VALPROATE SODIUM, VALPROATE SODIUM
 VANCOMYCIN HYDROCHLORIDE, VANCOMYCIN HYDROCHLORIDE

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

**** A ****

* ABRAXIS PHARMACEUTICAL PRODUCTS
VIBISONE, CYANOCOBALAMIN
VINBLASTINE SULFATE, VINBLASTINE SULFATE
VINCRISTINE SULFATE, VINCRISTINE SULFATE
VINORELBINE TARTRATE, VINORELBINE TARTRATE

ABRIKA PHARMS

* ABRIKA PHARMACEUTICALS LLLP
ISRADIPINE, ISRADIPINE
NIFEDIPINE, NIFEDIPINE

ACCORD HLTH

* ACCORD HEALTH CARE INC
METHYLDOPA, METHYLDOPA

ACORDA

* ACORDA THERAPEUTICS INC
ZANAFLEX, TIZANIDINE HYDROCHLORIDE

ACS DOBFAR

* ACS DOBFAR SPA
CEFTAZIDIME, CEFTAZIDIME

ACTAVIS ELIZABETH

* ACTAVIS ELIZABETH LLC
ACYCLOVIR, ACYCLOVIR
ALPRAZOLAM, ALPRAZOLAM
ASPIRIN AND CAFFEINE W/ BUTALBITAL, ASPIRIN
BISOPROLOL FUMARATE AND HYDROCHLOROTHIAZIDE, BISOPROLOL FUMARATE
CARBAMAZEPINE, CARBAMAZEPINE
CARBIDOPA AND LEVODOPA, CARBIDOPA
CITALOPRAM HYDROBROMIDE, CITALOPRAM HYDROBROMIDE
CLONAZEPAM, CLONAZEPAM
CLONIDINE HYDROCHLORIDE, CLONIDINE HYDROCHLORIDE
DIAZEPAM, DIAZEPAM
DICLOFENAC SODIUM, DICLOFENAC SODIUM
DILTIAZEM HYDROCHLORIDE, DILTIAZEM HYDROCHLORIDE
DOXAZOSIN MESYLATE, DOXAZOSIN MESYLATE
ETODOLAC, ETODOLAC
FAMOTIDINE, FAMOTIDINE
FENOPROFEN CALCIUM, FENOPROFEN CALCIUM
FLUVOXAMINE MALEATE, FLUVOXAMINE MALEATE
GABAPENTIN, GABAPENTIN
GLYBURIDE AND METFORMIN HYDROCHLORIDE, GLYBURIDE
HYDROCHLOROTHIAZIDE, HYDROCHLOROTHIAZIDE
INDAPAMIDE, INDAPAMIDE
ISOSORBIDE MONONITRATE, ISOSORBIDE MONONITRATE
LISINOPRIL AND HYDROCHLOROTHIAZIDE, HYDROCHLOROTHIAZIDE
LISINOPRIL, LISINOPRIL
LORAZEPAM, LORAZEPAM
LOVASTATIN, LOVASTATIN
METFORMIN HYDROCHLORIDE, METFORMIN HYDROCHLORIDE
METHYLPHENIDATE HYDROCHLORIDE, METHYLPHENIDATE HYDROCHLORIDE
METOCLOPRAMIDE HYDROCHLORIDE, METOCLOPRAMIDE HYDROCHLORIDE
MIRTAZAPINE, MIRTAZAPINE
NAPROXEN, NAPROXEN
NIFEDIPINE, NIFEDIPINE
OXAPROZIN, OXAPROZIN
OXAZEPAM, OXAZEPAM
PENTOXIFYLLINE, PENTOXIFYLLINE
PHENTERMINE HYDROCHLORIDE, PHENTERMINE HYDROCHLORIDE
PROPOXYPHENE NAPSYLATE AND ACETAMINOPHEN, ACETAMINOPHEN
PROPRANOLOL HYDROCHLORIDE AND HYDROCHLOROTHIAZIDE, HYDROCHLOROTHIAZIDE
PROPYLTHIOURACIL, PROPYLTHIOURACIL
SPIRONOLACTONE, SPIRONOLACTONE
TEMAZEPAM, TEMAZEPAM
TICLOPIDINE HYDROCHLORIDE, TICLOPIDINE HYDROCHLORIDE

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

** A **

* ACTAVIS ELIZABETH LLC
TIZANIDINE HYDROCHLORIDE, TIZANIDINE HYDROCHLORIDE
TOLMETIN SODIUM, TOLMETIN SODIUM
TRAMADOL HYDROCHLORIDE, TRAMADOL HYDROCHLORIDE
TRAZODONE HYDROCHLORIDE, TRAZODONE HYDROCHLORIDE
VERAPAMIL HYDROCHLORIDE, VERAPAMIL HYDROCHLORIDE

ACTAVIS MID ATLANTIC

* ACTAVIS MID ATLANTIC LLC
ACETAMINOPHEN AND CODEINE PHOSPHATE, ACETAMINOPHEN
ACETAMINOPHEN, ACETAMINOPHEN (OTC)
ACETASOL HC, ACETIC ACID, GLACIAL
ACETASOL, ACETIC ACID, GLACIAL
ACYCLOVIR, ACYCLOVIR
ALBUTEROL SULFATE, ALBUTEROL SULFATE
AMANTADINE HYDROCHLORIDE, AMANTADINE HYDROCHLORIDE
AMINOPHYLLINE DYE FREE, AMINOPHYLLINE
BETAMETHASONE DIPROPIONATE, BETAMETHASONE DIPROPIONATE
BETAMETHASONE VALERATE, BETAMETHASONE VALERATE
CAPITAL AND CODEINE, ACETAMINOPHEN
CHLORHEXIDINE GLUCONATE, CHLORHEXIDINE GLUCONATE
CHLORPROMAZINE HYDROCHLORIDE, CHLORPROMAZINE HYDROCHLORIDE
CIMETIDINE HYDROCHLORIDE, CIMETIDINE HYDROCHLORIDE
CLEMASTINE FUMARATE, CLEMASTINE FUMARATE
CLINDAMYCIN PHOSPHATE, CLINDAMYCIN PHOSPHATE
CLOBETASOL PROPIONATE, CLOBETASOL PROPIONATE
CLOTRIMAZOLE AND BETAMETHASONE DIPROPIONATE, BETAMETHASONE DIPROPIONATE
CLOTRIMAZOLE, CLOTRIMAZOLE (OTC)
CONSTULOSE, LACTULOSE
CROMOLYN SODIUM, CROMOLYN SODIUM
CROMOLYN SODIUM, CROMOLYN SODIUM (OTC)
CYPROHEPTADINE HYDROCHLORIDE, CYPROHEPTADINE HYDROCHLORIDE
DEXAMETHASONE, DEXAMETHASONE
ENULOSE, LACTULOSE
FLUOCINONIDE EMULSIFIED BASE, FLUOCINONIDE
FLUOCINONIDE, FLUOCINONIDE
FLUOXETINE, FLUOXETINE HYDROCHLORIDE
HALOBETASOL PROPIONATE, HALOBETASOL PROPIONATE
HALOPERIDOL LACTATE, HALOPERIDOL LACTATE
HYDROCODONE COMPOUND, HOMATROPINE METHYLBROMIDE
HYDROCORTISONE, HYDROCORTISONE
HYDROXYZINE HYDROCHLORIDE, HYDROXYZINE HYDROCHLORIDE
IBUPROFEN, IBUPROFEN
IBUPROFEN, IBUPROFEN (OTC)
INFANTS' FEVERALL, ACETAMINOPHEN (OTC)
IPRATROPIUM BROMIDE, IPRATROPIUM BROMIDE
LIDOCAINE HYDROCHLORIDE VISCOUS, LIDOCAINE HYDROCHLORIDE
LINDANE, LINDANE
METOCLOPRAMIDE HYDROCHLORIDE, METOCLOPRAMIDE HYDROCHLORIDE
MICONAZOLE 7, MICONAZOLE NITRATE (OTC)
MICONAZOLE NITRATE, MICONAZOLE NITRATE
MICONAZOLE NITRATE, MICONAZOLE NITRATE (OTC)
MINOXIDIL (FOR MEN), MINOXIDIL (OTC)
MINOXIDIL EXTRA STRENGTH (FOR MEN), MINOXIDIL (OTC)
MYKACET, NYSTATIN
M-ZOLE 3 COMBINATION PACK, MICONAZOLE NITRATE (OTC)
M-ZOLE 7 DUAL PACK, MICONAZOLE NITRATE (OTC)
NYSTATIN, NYSTATIN
PERMETHRIN, PERMETHRIN
PERMETHRIN, PERMETHRIN (OTC)
PHENYTOIN, PHENYTOIN
PROMETH VC PLAIN, PHENYLEPHRINE HYDROCHLORIDE
PROMETH VC W/ CODEINE, CODEINE PHOSPHATE
PROMETH W/ CODEINE, CODEINE PHOSPHATE

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

** A **

* ACTAVIS MID ATLANTIC LLC
PROMETH W/ DEXTROMETHORPHAN, DEXTROMETHORPHAN HYDROBROMIDE
SELENIUM SULFIDE, SELENIUM SULFIDE
SULFATRIM PEDIATRIC, SULFAMETHOXAZOLE
SULFATRIM, SULFAMETHOXAZOLE
THEOPHYLLINE, THEOPHYLLINE
THIORIDAZINE HYDROCHLORIDE, THIORIDAZINE HYDROCHLORIDE
TRIAFIN-C, CODEINE PHOSPHATE
TRIAMCINOLONE ACETONIDE, TRIAMCINOLONE ACETONIDE
VALNAC, BETAMETHASONE VALERATE

ACTAVIS TOTOWA

* ACTAVIS TOTOWA LLC
ACETAMINOPHEN AND PENTAZOCINE HYDROCHLORIDE, ACETAMINOPHEN
AMANTADINE HYDROCHLORIDE, AMANTADINE HYDROCHLORIDE
BETAXOLOL HYDROCHLORIDE, BETAXOLOL HYDROCHLORIDE
BETHANECHOL CHLORIDE, BETHANECHOL CHLORIDE
BUSPIRONE HYDROCHLORIDE, BUSPIRONE HYDROCHLORIDE
CARISOPRODOL AND ASPIRIN, ASPIRIN
CARISOPRODOL, CARISOPRODOL
CARISOPRODOL, ASPIRIN AND CODEINE PHOSPHATE, ASPIRIN
CHLORZOXAZONE, CHLORZOXAZONE
CILOSTAZOL, CILOSTAZOL
CYCLOBENZAPRINE HYDROCHLORIDE, CYCLOBENZAPRINE HYDROCHLORIDE
DANTROLENE SODIUM, DANTROLENE SODIUM
DIGOXIN, DIGOXIN
DIPYRIDAMOLE, DIPYRIDAMOLE
GLYBURIDE, GLYBURIDE
GUANFACINE HYDROCHLORIDE, GUANFACINE HYDROCHLORIDE
HOMATROPINE METHYLBROMIDE AND HYDROCODONE BITARTRATE, HOMATROPINE METHYLBROMIDE
HYDROMORPHONE HYDROCHLORIDE, HYDROMORPHONE HYDROCHLORIDE
HYDROXYZINE HYDROCHLORIDE, HYDROXYZINE HYDROCHLORIDE
ISRADIPINE, ISRADIPINE
LOXAPINE SUCCINATE, LOXAPINE SUCCINATE
MELOXICAM, MELOXICAM
MEPERIDINE HYDROCHLORIDE, MEPERIDINE HYDROCHLORIDE
MIRTAZAPINE, MIRTAZAPINE
NALOXONE HYDROCHLORIDE AND PENTAZOCAINE, NALOXONE HYDROCHLORIDE
NALTREXONE HYDROCHLORIDE, NALTREXONE HYDROCHLORIDE
OXYCODONE AND ACETAMINOPHEN, ACETAMINOPHEN
OXYCODONE HYDROCHLORIDE, OXYCODONE HYDROCHLORIDE
PHENTERMINE HYDROCHLORIDE, PHENTERMINE HYDROCHLORIDE
QUINAPRIL HYDROCHLORIDE, QUINAPRIL HYDROCHLORIDE
QUINARETIC, HYDROCHLOROTHIAZIDE
RIMACTANE, RIFAMPIN
RIMANTADINE HYDROCHLORIDE, RIMANTADINE HYDROCHLORIDE
TIZANIDINE HYDROCHLORIDE, TIZANIDINE HYDROCHLORIDE
TRIMETHOBENZAMIDE HYDROCHLORIDE, TRIMETHOBENZAMIDE HYDROCHLORIDE
TRIMIPRAMINE MALEATE, TRIMIPRAMINE MALEATE
URSODIOL, URSODIOL

ACTELION

* ACTELION LTD
TRACLEER, BOSENTAN

ACTELION PHARMS

* ACTELION PHARMACEUTICALS US INC
ZAVESCA, MIGLUSTAT

ADAMS RESP THERAP

* ADAMS RESPIRATORY THERAPEUTICS
DELSYM, DEXTROMETHORPHAN POLISTIREX (OTC)
HUMABID, GUAIFENESIN (OTC)
MUCINEX D, GUAIFENESIN (OTC)
MUCINEX DM, DEXTROMETHORPHAN HYDROBROMIDE (OTC)
MUCINEX, GUAIFENESIN (OTC)

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

** A **

ADV CARE

- * ADVANCED CARE PRODUCTS DIV ORTHO PHARMACEUTICAL CORP
MONISTAT 7 COMBINATION PACK, MICONAZOLE NITRATE (OTC)
MONISTAT 7, MICONAZOLE NITRATE (OTC)
MONISTAT-3 COMBINATION PACK, MICONAZOLE NITRATE (OTC)

ADV CARE PRODS

- * ADVANCED CARE PRODUCTS
MONISTAT 3, MICONAZOLE NITRATE (OTC)
MONISTAT 7, MICONAZOLE NITRATE (OTC)

ADV MAGNETICS

- * ADVANCED MAGNETICS INC
FERIDEX I.V., FERUMOXIDES
GASTROMARK, FERUMOXIL

ADVANCIS PHARM

- * ADVANCIS PHARMACEUTICAL CORP
KEFLEX, CEPHALEXIN

AEGIS PHARMS

- * AEGIS PHARMACEUTICALS INC
TAMOXIFEN CITRATE, TAMOXIFEN CITRATE

AESGEN

- * AESGEN INC
PAMIDRONATE DISODIUM, PAMIDRONATE DISODIUM

AGOURON

- * AGOURON PHARMACEUTICALS INC
RESCRIPTOR, DELAVIRDINE MESYLATE
VIRACEPT, NELFINAVIR MESYLATE

AKORN

- * AKORN INC
AKBETA, LEVOBUNOLOL HYDROCHLORIDE
AKPENTOLATE, CYCLOPENTOLATE HYDROCHLORIDE
AKPRO, DIPIVEFRIN HYDROCHLORIDE
AKTOB, TOBRAMYCIN
ALFENTA, ALFENTANIL HYDROCHLORIDE
AMIODARONE HYDROCHLORIDE, AMIODARONE HYDROCHLORIDE
BACITRACIN ZINC AND POLYMYXIN B SULFATE, BACITRACIN ZINC
BAL, DIMERCAPROL
BALANCED SALT SOLUTION, CALCIUM CHLORIDE
BETAXOLOL, BETAXOLOL HYDROCHLORIDE
BRIMONIDINE TARTRATE, BRIMONIDINE TARTRATE
CROMOLYN SODIUM, CROMOLYN SODIUM
ENDOSOL EXTRA, CALCIUM CHLORIDE
ERYTHROMYCIN, ERYTHROMYCIN
GENTAK, GENTAMICIN SULFATE
GENTAMICIN SULFATE, GENTAMICIN SULFATE
IC-GREEN, INDOCYANINE GREEN
INAPSINE, DROPERIDOL
LIDOCAINE HYDROCHLORIDE, LIDOCAINE HYDROCHLORIDE
NEOMYCIN AND POLYMYXIN B SULFATE AND BACITRACIN ZINC, BACITRACIN ZINC
ORPHENADRINE CITRATE, ORPHENADRINE CITRATE
PAREMYD, HYDROXYAMPHETAMINE HYDROBROMIDE
SUFENTA, SUFENTANIL CITRATE
SULFACETAMIDE SODIUM, SULFACETAMIDE SODIUM
TIMOLOL MALEATE, TIMOLOL MALEATE
TROPICACYL, TROPICAMIDE

AKORN MFG

- * AKORN MANUFACTURING INC
SUBLIMAZE PRESERVATIVE FREE, FENTANYL CITRATE

AKYMA PHARMS

- * AKYMA PHARMACEUTICALS LLC
CITALOPRAM HYDROBROMIDE, CITALOPRAM HYDROBROMIDE

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

**** A ****

ALARA PHARM

* ALARA PHARMACEUTICAL CORPORATION
LEVO-T, LEVOTHYROXINE SODIUM

ALAVEN PHARM

* ALAVEN PHARMACEUTICAL LLC
ANADROL-50, OXYMETHOLONE
ROWASA, MESALAMINE

ALCON

* ALCON INC
AZOPT, BRINZOLAMIDE
BRIMONIDINE TARTRATE, BRIMONIDINE TARTRATE
CILOXAN, CIPROFLOXACIN HYDROCHLORIDE
CIPRO HC, CIPROFLOXACIN HYDROCHLORIDE
CIPRODEX, CIPROFLOXACIN
FLUORESCITE, FLUORESCIN SODIUM
NEVANAC, NEPAFENAC
OFLOXACIN, OFLOXACIN
PATADAY, OLOPATADINE HYDROCHLORIDE
SULFACETAMIDE SODIUM AND PREDNISOLONE SODIUM PHOSPHATE, PREDNISOLONE SODIUM PHOSPHATE
SULFACETAMIDE SODIUM, SULFACETAMIDE SODIUM
TRAVATAN Z, TRAVOPROST
TRAVATAN, TRAVOPROST
VIGAMOX, MOXIFLOXACIN HYDROCHLORIDE
* ALCON LABORATORIES INC
ALCAINE, PROPARACAINE HYDROCHLORIDE
ALOMIDE, LODOXAMIDE TROMETHAMINE
BETADINE, POVIDONE-IODINE
BETOPTIC S, BETAXOLOL HYDROCHLORIDE
BETOPTIC, BETAXOLOL HYDROCHLORIDE
BSS PLUS, CALCIUM CHLORIDE
BSS, CALCIUM CHLORIDE
CARTEOLOL HYDROCHLORIDE, CARTEOLOL HYDROCHLORIDE
CETAMIDE, SULFACETAMIDE SODIUM
CROMOLYN SODIUM, CROMOLYN SODIUM
CYCLOGYL, CYCLOPENTOLATE HYDROCHLORIDE
CYCLOMYDRIL, CYCLOPENTOLATE HYDROCHLORIDE
DENDRID, IDOXURIDINE
EMADINE, EMEDASTINE DIFUMARATE
FLAREX, FLUOROMETHOLONE ACETATE
IOPIDINE, APRACLOXIDINE HYDROCHLORIDE
ISOPTO CETAMIDE, SULFACETAMIDE SODIUM
MAXIDEX, DEXAMETHASONE
MAXITROL, DEXAMETHASONE
MIOSTAT, CARBACHOL
MYDRIACYL, TROPICAMIDE
NAPHCON-A, NAPHAZOLINE HYDROCHLORIDE (OTC)
NATACYN, NATAMYCIN
NEOMYCIN AND POLYMYXIN B SULFATES AND HYDROCORTISONE, HYDROCORTISONE
OMNIPRED, PREDNISOLONE ACETATE
PATANOL, OLOPATADINE HYDROCHLORIDE
PILOPINE HS, PILOCARPINE HYDROCHLORIDE
TOBRADEX, DEXAMETHASONE
TOBRASONE, FLUOROMETHOLONE ACETATE
TOBREX, TOBRAMYCIN
TRIFLURIDINE, TRIFLURIDINE
VEXOL, RIMEXOLONE

ALCON RES

* ALCON RESEARCH LTD
BRIMONIDINE TARTRATE, BRIMONIDINE TARTRATE

ALCON UNIVERSAL

* ALCON UNIVERSAL LTD
CYCLOPENTOLATE HYDROCHLORIDE, CYCLOPENTOLATE HYDROCHLORIDE

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

** A **

- * ALCON UNIVERSAL LTD
DEXAMETHASONE SODIUM PHOSPHATE, DEXAMETHASONE SODIUM PHOSPHATE
NEOMYCIN AND POLYMYXIN B SULFATES AND DEXAMETHASONE, DEXAMETHASONE
NEOMYCIN AND POLYMYXIN B SULFATES AND HYDROCORTISONE, HYDROCORTISONE
SULFACETAMIDE SODIUM, SULFACETAMIDE SODIUM
TROPICAMIDE, TROPICAMIDE

ALIMERA SCIENCES INC

- * ALIMERA SCIENCES INC
ALAWAY, KETOTIFEN FUMARATE (OTC)

ALKERMES

- * ALKERMES INC
VIVITROL, NALTREXONE

ALLEGIANCE HLTHCARE

- * ALLEGIANCE HEALTHCARE CORP
POVIDONE IODINE, POVIDONE-IODINE (OTC)

ALLERGAN

- * ALLERGAN
ACULAR LS, KETOROLAC TROMETHAMINE
ALPHAGAN P, BRIMONIDINE TARTRATE
BLEPH-10, SULFACETAMIDE SODIUM
BLEPH-30, SULFACETAMIDE SODIUM
GENOPTIC, GENTAMICIN SULFATE
- * ALLERGAN INC
ACULAR PRESERVATIVE FREE, KETOROLAC TROMETHAMINE
ACULAR, KETOROLAC TROMETHAMINE
ALBALON, NAPHAZOLINE HYDROCHLORIDE
ALOCRIL, NEDOCROMIL SODIUM
ALPHAGAN P, BRIMONIDINE TARTRATE
AVAGE, TAZAROTENE
AZELEX, AZELAIC ACID
ELESTAT, EPINASTINE HYDROCHLORIDE
ELIMITE, PERMETHRIN
LUMIGAN, BIMATOPROST
OCUFLOX, OFLOXACIN
OPTICROM, CROMOLYN SODIUM
POLYTRIM, POLYMYXIN B SULFATE
RESTASIS, CYCLOSPORINE
TAZORAC, TAZAROTENE
ZYMAR, GATIFLOXACIN
- * ALLERGAN PHARMACEUTICAL
BETAGAN, LEVOBUNOLOL HYDROCHLORIDE
BLEPHAMIDE S.O.P., PREDNISOLONE ACETATE
BLEPHAMIDE, PREDNISOLONE ACETATE
FML FORTE, FLUOROMETHOLONE
FML, FLUOROMETHOLONE
FML-S, FLUOROMETHOLONE
HERPLEX, IDOXURIDINE
HMS, MEDRYSONE
OCUFEN, FLURBIPROFEN SODIUM
OPHTHETIC, PROPARACAINE HYDROCHLORIDE
POLY-PRED, NEOMYCIN SULFATE
PRED FORTE, PREDNISOLONE ACETATE
PRED MILD, PREDNISOLONE ACETATE
PRED-G, GENTAMICIN SULFATE
PROPINE, DIPIVEFRIN HYDROCHLORIDE

ALLERGAN HERBERT

- * ALLERGAN HERBERT SKIN CARE DIV ALLERGAN INC
FLUOROPLEX, FLUOROURACIL

ALPHAPHARM

- * ALPHAPHARM PARTY LTD
ALPRAZOLAM, ALPRAZOLAM
AMIODARONE HYDROCHLORIDE, AMIODARONE HYDROCHLORIDE

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

**** A ****

* ALPHAPHARM PARTY LTD
BACLOFEN, BACLOFEN
CILOSTAZOL, CILOSTAZOL
CITALOPRAM HYDROBROMIDE, CITALOPRAM HYDROBROMIDE
CLONAZEPAM, CLONAZEPAM
DICLOFENAC SODIUM, DICLOFENAC SODIUM
FLECAINIDE ACETATE, FLECAINIDE ACETATE
FLUOXETINE HYDROCHLORIDE, FLUOXETINE HYDROCHLORIDE
FLUOXETINE, FLUOXETINE HYDROCHLORIDE
GLIPIZIDE, GLIPIZIDE
INDAPAMIDE, INDAPAMIDE
METFORMIN HYDROCHLORIDE, METFORMIN HYDROCHLORIDE
MIRTAZAPINE, MIRTAZAPINE
NAPROXEN, NAPROXEN
SELEGILINE HYDROCHLORIDE, SELEGILINE HYDROCHLORIDE
TIZANIDINE HYDROCHLORIDE, TIZANIDINE HYDROCHLORIDE
TRAMADOL HYDROCHLORIDE, TRAMADOL HYDROCHLORIDE
TRIAZOLAM, TRIAZOLAM
ZONISAMIDE, ZONISAMIDE
* ALPHAPHARM PTY LTD
ACEBUTOLOL HYDROCHLORIDE, ACEBUTOLOL HYDROCHLORIDE
ANAGRELIDE HYDROCHLORIDE, ANAGRELIDE HYDROCHLORIDE
PAROXETINE HYDROCHLORIDE, PAROXETINE HYDROCHLORIDE

ALPHARMA US PHARMS

* ALPHARMA US PHARMACEUTICALS DIVISION
ERYTHROMYCIN ESTOLATE, ERYTHROMYCIN ESTOLATE
ERYTHROMYCIN ETHYLSUCCINATE, ERYTHROMYCIN ETHYLSUCCINATE
KADIAN, MORPHINE SULFATE

ALRA

* ALRA LABORATORIES INC
CHOLAC, LACTULOSE
CONSTILAC, LACTULOSE
ERYZOLE, ERYTHROMYCIN ETHYLSUCCINATE
GEN-XENE, CLORAZEPATE DIPOTASSIUM
IBU-TAB 200, IBUPROFEN (OTC)
IBU-TAB, IBUPROFEN
K+10, POTASSIUM CHLORIDE
K+8, POTASSIUM CHLORIDE

ALTANA

* ALTANA INC
ALCLOMETASONE DIPROPIONATE, ALCLOMETASONE DIPROPIONATE
AMCINONIDE, AMCINONIDE
BACITRACIN ZINC AND POLYMYXIN B SULFATE, BACITRACIN ZINC
BACITRACIN, BACITRACIN
BACITRACIN-NEOMYCIN-POLYMYXIN, BACITRACIN ZINC
BETAMETHASONE DIPROPIONATE, BETAMETHASONE DIPROPIONATE
CHLORAMPHENICOL, CHLORAMPHENICOL
CICLOPIROX, CICLOPIROX
CLINDAMYCIN PHOSPHATE, CLINDAMYCIN PHOSPHATE
CLOBETASOL PROPIONATE (EMOLLIENT), CLOBETASOL PROPIONATE
CLOBETASOL PROPIONATE, CLOBETASOL PROPIONATE
CLOTRIMAZOLE AND BETAMETHASONE DIPROPIONATE, BETAMETHASONE DIPROPIONATE
CUTIVATE, FLUTICASONE PROPIONATE
DESONIDE, DESONIDE
DIFLORASONE DIACETATE, DIFLORASONE DIACETATE
ECONAZOLE NITRATE, ECONAZOLE NITRATE
ERYTHROMYCIN, ERYTHROMYCIN
FLUOCINONIDE EMULSIFIED BASE, FLUOCINONIDE
FLUOCINONIDE, FLUOCINONIDE
FLUTICASONE PROPIONATE, FLUTICASONE PROPIONATE
GENTAMICIN SULFATE, GENTAMICIN SULFATE
HALOBETASOL PROPIONATE, HALOBETASOL PROPIONATE
HYDROCORTISONE VALERATE, HYDROCORTISONE VALERATE

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

** A **

* ALTANA INC
HYDROCORTISONE, HYDROCORTISONE
KETOCONAZOLE, KETOCONAZOLE
LIDOCAINE AND PRILOCAINE, LIDOCAINE
METRONIDAZOLE, METRONIDAZOLE
MOMETASONE FUROATE, MOMETASONE FUROATE
MUPIROCIN, MUPIROCIN
NYSTATIN, NYSTATIN
OXISTAT, OXICONAZOLE NITRATE
PREDNICARBATE, PREDNICARBATE
SULFACETAMIDE SODIUM, SULFACETAMIDE SODIUM
TEMOVATE E, CLOBETASOL PROPIONATE
TEMOVATE, CLOBETASOL PROPIONATE
TERCONAZOLE, TERCONAZOLE
TOBRAMYCIN, TOBRAMYCIN
TRIAMCINOLONE ACETONIDE, TRIAMCINOLONE ACETONIDE

ALTANA PHARMA

* ALTANA PHARMA
OMNARIS, CICLESONIDE
* ALTANA PHARMA AG
CLINDAMYCIN PHOSPHATE, CLINDAMYCIN PHOSPHATE
CLOTRIMAZOLE AND BETAMETHASONE DIPROPIONATE, BETAMETHASONE DIPROPIONATE

ALTERNA-TCHP LLC

* ALTERNA-TCHP LLC
CHILDREN'S ELIXSURE, IBUPROFEN (OTC)

ALZA

* ALZA CORP
CONCERTA, METHYLPHENIDATE HYDROCHLORIDE
DITROPAN XL, OXYBUTYNIN CHLORIDE
DITROPAN, OXYBUTYNIN CHLORIDE
DOXIL, DOXORUBICIN HYDROCHLORIDE
DURAGESIC-100, FENTANYL
DURAGESIC-12, FENTANYL
DURAGESIC-25, FENTANYL
DURAGESIC-50, FENTANYL
DURAGESIC-75, FENTANYL
EFIDAC 24 PSEUDOEPHEDRINE HYDROCHLORIDE, PSEUDOEPHEDRINE HYDROCHLORIDE (OTC)
EFIDAC 24 PSEUDOEPHEDRINE HYDROCHLORIDE/BROMPHENIRAMINE MALEATE, BROMPHENIRAMINE MALEATE (OTC)
IONSYS, FENTANYL HYDROCHLORIDE
VIADUR, LEUPROLIDE ACETATE

AM ANTIBIOTICS

* AMERICAN ANTIBIOTICS LLC
AMOXICILLIN, AMOXICILLIN
PENICILLIN V POTASSIUM, PENICILLIN V POTASSIUM

AMBI PHARMS

* AMBI PHARMACEUTICALS INC
PHENTERMINE HYDROCHLORIDE, PHENTERMINE HYDROCHLORIDE

AMERSHAM

* AMERSHAM CORP SUB RADIOCHEMICAL CENTER
INDICLOR, INDIUM IN-111 CHLORIDE

AMGEN

* AMGEN INC
SENSIPAR, CINACALCET HYDROCHLORIDE

AMNEAL PHARM

* AMNEAL PHARMACEUTICAL
METFORMIN HYDROCHLORIDE, METFORMIN HYDROCHLORIDE

AMPHASTAR

* AMPHASTAR PHARMACEUTICALS INC
CORTROSYN, COSYNTROPIN
DUOCAINE, BUPIVACAINE HYDROCHLORIDE

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

**** A ****

AMPHASTAR PHARM

- * AMPHASTAR PHARMACEUTICAL INC
AMPHADASE, HYALURONIDASE
KETOROLAC TROMETHAMINE, KETOROLAC TROMETHAMINE

AMYLIN

- * AMYLIN PHARMACEUTICALS INC
BYETTA, EXENATIDE SYNTHETIC
SYMLIN, PRAMLINTIDE ACETATE

ANABOLIC

- * ANABOLIC INC
BUTALBITAL, ASPIRIN, CAFFEINE, AND CODEINE PHOSPHATE, ASPIRIN

ANABOLIC LABS

- * ANABOLIC LABORATORIES INC
BUTALBITAL, ACETAMINOPHEN, CAFFEINE AND CODEINE PHOSPHATE, ACETAMINOPHEN

ANBEX

- * ANBEX INC
IOSAT, POTASSIUM IODIDE (OTC)

ANCHEN PHARMS

- * ANCHEN PHARMACEUTICALS INC
BUPROPION HYDROCHLORIDE, BUPROPION HYDROCHLORIDE

ANDRX

- * ANDRX CORP
DILTIAZEM HYDROCHLORIDE, DILTIAZEM HYDROCHLORIDE

ANDRX LABS LLC

- * ANDRX LABS LLC
ALTOPREV, LOVASTATIN
FORTAMET, METFORMIN HYDROCHLORIDE

ANDRX PHARMS

- * ANDRX PHARMACEUTICALS INC
ACETAMINOPHEN AND CODEINE PHOSPHATE, ACETAMINOPHEN
BENAZEPRIL HYDROCHLORIDE AND HYDROCHLOROTHIAZIDE, BENAZEPRIL HYDROCHLORIDE
BENAZEPRIL HYDROCHLORIDE, BENAZEPRIL HYDROCHLORIDE
CARTIA XT, DILTIAZEM HYDROCHLORIDE
CLARITHROMYCIN, CLARITHROMYCIN
CODRIX, ACETAMINOPHEN
FOSINOPRIL SODIUM AND HYDROCHLOROTHIAZIDE, FOSINOPRIL SODIUM
FOSINOPRIL SODIUM, FOSINOPRIL SODIUM
HYDROCODONE BITARTRATE AND ACETAMINOPHEN, ACETAMINOPHEN
HYDROCODONE BITARTRATE AND IBUPROFEN, HYDROCODONE BITARTRATE
KETOPROFEN, KETOPROFEN
LORATADINE AND PSEUDOEPHEDRINE SULFATE, LORATADINE (OTC)
LORATADINE, LORATADINE (OTC)
METFORMIN HYDROCHLORIDE, METFORMIN HYDROCHLORIDE
MIRTAZAPINE, MIRTAZAPINE
NAPROXEN SODIUM, NAPROXEN SODIUM
NORETHINDRONE ACETATE AND ETHINYL ESTRADIOL AND FERROUS FUMARATE, ETHINYL ESTRADIOL
POTASSIUM CHLORIDE, POTASSIUM CHLORIDE
PREVIFEM, ETHINYL ESTRADIOL
PROPOXYPHENE NAPSYLATE AND ACETAMINOPHEN, ACETAMINOPHEN
QUINAPRIL HYDROCHLORIDE, QUINAPRIL HYDROCHLORIDE
TAMOXIFEN CITRATE, TAMOXIFEN CITRATE
TAZTIA XT, DILTIAZEM HYDROCHLORIDE
TRI-PREVIFEM, ETHINYL ESTRADIOL
- * ANDRX PHARMACEUTICALS LLC
CODRIX, ACETAMINOPHEN
ETODOLAC, ETODOLAC
LORATADINE AND PSEUDOEPHEDRINE SULFATE, LORATADINE (OTC)
PROPOXYPHENE NAPSYLATE AND ACETAMINOPHEN, ACETAMINOPHEN

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

**** A ****

ANGELINI

* ANGELINI PHARMACEUTICALS INC
REV-EYES, DAPIPRAZOLE HYDROCHLORIDE

APOTEX

* APOTEX CORP
PRAVASTATIN SODIUM, PRAVASTATIN SODIUM
* APOTEX INC
ONDANSETRON HYDROCHLORIDE, ONDANSETRON HYDROCHLORIDE

APOTEX INC

* APOTEX INC
PAROXETINE HYDROCHLORIDE, PAROXETINE HYDROCHLORIDE
TRAZODONE HYDROCHLORIDE, TRAZODONE HYDROCHLORIDE
TRIAMTERENE AND HYDROCHLOROTHIAZIDE, HYDROCHLOROTHIAZIDE
* APOTEX INC ETOBICOKE SITE
ACYCLOVIR, ACYCLOVIR
ALLOPURINOL, ALLOPURINOL
BENAZEPRIL HYDROCHLORIDE, BENAZEPRIL HYDROCHLORIDE
BUPROPION HYDROCHLORIDE, BUPROPION HYDROCHLORIDE
CARBAMAZEPINE, CARBAMAZEPINE
CEFUROXIME AXETIL, CEFUROXIME AXETIL
CILOSTAZOL, CILOSTAZOL
CITALOPRAM HYDROBROMIDE, CITALOPRAM HYDROBROMIDE
CLOPIDOGREL BISULFATE, CLOPIDOGREL BISULFATE
DESMOPRESSIN ACETATE, DESMOPRESSIN ACETATE
DILTIZAC, DILTIAZEM HYDROCHLORIDE
ENALAPRIL MALEATE AND HYDROCHLOROTHIAZIDE, ENALAPRIL MALEATE
ETODOLAC, ETODOLAC
FOSINOPRIL SODIUM, FOSINOPRIL SODIUM
GABAPENTIN, GABAPENTIN
KETOTIFEN FUMARATE, KETOTIFEN FUMARATE
LEFLUNOMIDE, LEFLUNOMIDE
LISINOPRIL AND HYDROCHLOROTHIAZIDE, HYDROCHLOROTHIAZIDE
LISINOPRIL, LISINOPRIL
LITHIUM CARBONATE, LITHIUM CARBONATE
LORATADINE, LORATADINE (OTC)
MELOXICAM, MELOXICAM
METFORMIN HYDROCHLORIDE, METFORMIN HYDROCHLORIDE
MIDODRINE HYDROCHLORIDE, MIDODRINE HYDROCHLORIDE
OXAPROZIN, OXAPROZIN
SELEGILINE HYDROCHLORIDE, SELEGILINE HYDROCHLORIDE
SOTALOL HYDROCHLORIDE, SOTALOL HYDROCHLORIDE
TORSEMIDE, TORSEMIDE
ZONISAMIDE, ZONISAMIDE
* APOTEX INC RICHMOND HILL
AMIODARONE HYDROCHLORIDE, AMIODARONE HYDROCHLORIDE
BUTORPHANOL TARTRATE PRESERVATIVE FREE, BUTORPHANOL TARTRATE
BUTORPHANOL TARTRATE, BUTORPHANOL TARTRATE
CITALOPRAM HYDROBROMIDE, CITALOPRAM HYDROBROMIDE
DESMOPRESSIN ACETATE (NEEDS NO REFRIGERATION), DESMOPRESSIN ACETATE
DILTIAZEM HYDROCHLORIDE, DILTIAZEM HYDROCHLORIDE
DIPYRIDAMOLE, DIPYRIDAMOLE
EDETATE DISODIUM, EDETATE DISODIUM
FAMOTIDINE PRESERVATIVE FREE (PHARMACY BULK), FAMOTIDINE
FAMOTIDINE PRESERVATIVE FREE, FAMOTIDINE
FAMOTIDINE, FAMOTIDINE
FLUCONAZOLE IN DEXTROSE 5% IN PLASTIC CONTAINER, FLUCONAZOLE
FLUCONAZOLE IN SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER, FLUCONAZOLE
FLUMAZENIL, FLUMAZENIL
FLUPHENAZINE DECANOATE, FLUPHENAZINE DECANOATE
HALOPERIDOL DECANOATE, HALOPERIDOL DECANOATE
HALOPERIDOL, HALOPERIDOL LACTATE
KETOROLAC TROMETHAMINE, KETOROLAC TROMETHAMINE
LABETALOL HYDROCHLORIDE, LABETALOL HYDROCHLORIDE

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

**** A ****

* APOTEX INC RICHMOND HILL
LORATADINE, LORATADINE (OTC)
MEGESTROL ACETATE, MEGESTROL ACETATE
MIDAZOLAM HYDROCHLORIDE, MIDAZOLAM HYDROCHLORIDE
MILRINONE LACTATE IN 5% DEXTROSE IN PLASTIC CONTAINER, MILRINONE LACTATE
MILRINONE LACTATE, MILRINONE LACTATE
ONDANSETRON HYDROCHLORIDE PRESERVATIVE FREE, ONDANSETRON HYDROCHLORIDE
PREDNISOLONE, PREDNISOLONE
VALPROIC ACID, VALPROIC ACID

APOTHECON

* APOTHECON INC DIV BRISTOL MYERS SQUIBB
ACYCLOVIR, ACYCLOVIR
AMIKIN, AMIKACIN SULFATE
CAPOZIDE 25/15, CAPTOPRIL
CAPOZIDE 25/25, CAPTOPRIL
CAPOZIDE 50/15, CAPTOPRIL
CAPOZIDE 50/25, CAPTOPRIL
CEFAZOLIN SODIUM, CEFZOLIN SODIUM
CEPHALEXIN, CEPHALEXIN
DICLOXACILLIN SODIUM, DICLOXACILLIN SODIUM
FUNGIZONE, AMPHOTERICIN B
KANTREX, KANAMYCIN SULFATE
KENALOG IN ORABASE, TRIAMCINOLONE ACETONIDE
KENALOG, TRIAMCINOLONE ACETONIDE
KENALOG-10, TRIAMCINOLONE ACETONIDE
KENALOG-40, TRIAMCINOLONE ACETONIDE
KLOTRIX, POTASSIUM CHLORIDE
MUCOMYST, ACETYLCYSTEINE
MYCOSTATIN, NYSTATIN
OPHTHAINE, PROPARACAINE HYDROCHLORIDE
PRINCIPEN, AMPICILLIN/AMPICILLIN TRIHYDRATE
PROLIXIN, FLUPHENAZINE HYDROCHLORIDE
PRONESTYL, PROCAINAMIDE HYDROCHLORIDE
PRONESTYL-SR, PROCAINAMIDE HYDROCHLORIDE
STADOL PRESERVATIVE FREE, BUTORPHANOL TARTRATE
STADOL, BUTORPHANOL TARTRATE
TOBRAMYCIN SULFATE, TOBRAMYCIN SULFATE
TRIMOX, AMOXICILLIN
VEETIDS, PENICILLIN V POTASSIUM
VELOSEF '125', CEPHRADINE
VELOSEF '250', CEPHRADINE

APOTHEKERNES

* APOTHEKERNES LABORATORIUM A/S
BACITRACIN, BACITRACIN

APPLIED ANAL

* APPLIED ANALYTICAL INDUSTRIES INC
ESTRADIOL, ESTRADIOL

AR HOLDING CO INC

* AR HOLDING CO INC
QUININE SULFATE, QUININE SULFATE

ARCUM

* ARCUM PHARMACEUTICAL CORP
VITAMIN A PALMITATE, VITAMIN A PALMITATE

ARMSTRONG PHARMS

* ARMSTRONG PHARMACEUTICALS INC
ALBUTEROL, ALBUTEROL
EPINEPHRINE, EPINEPHRINE (OTC)

ASCEND

* ASCEND THERAPEUTICS INC
ESTROGEL, ESTRADIOL

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

** A **

ASCENT PEDS

- * ASCENT PEDIATRICS INC
ORAPRED, PREDNISOLONE SODIUM PHOSPHATE

ASTELLAS

- * ASTELLAS PHARMA US INC
ADENOCARD, ADENOSINE
ADENOSCAN, ADENOSINE
AMBISOME, AMPHOTERICIN B
CEFIZOX IN PLASTIC CONTAINER, CEFTIZOXIME SODIUM
CEFIZOX, CEFTIZOXIME SODIUM
MYCAMINE, MICAFUNGIN SODIUM
PROGRAF, TACROLIMUS
PROTOPIC, TACROLIMUS
VAPRISOL, CONIVAPTAN HYDROCHLORIDE
VESICARE, SOLIFENACIN SUCCINATE

ASTRAZENECA

- * ASTRAZENECA LP
ASTRAMORPH PF, MORPHINE SULFATE
BUDESONIDE, BUDESONIDE
DUTOPROL, HYDROCHLOROTHIAZIDE
ENTOCORT EC, BUDESONIDE
FOSCAVIR, FOSCARNET SODIUM
MEPERIDINE HYDROCHLORIDE PRESERVATIVE FREE, MEPERIDINE HYDROCHLORIDE
NEXIUM IV, ESOMEPRAZOLE SODIUM
NEXIUM, ESOMEPRAZOLE MAGNESIUM
POLOCAINE, MEPIVACAINE HYDROCHLORIDE
POLOCAINE-MPF, MEPIVACAINE HYDROCHLORIDE
PRILOSEC OTC, OMEPRAZOLE MAGNESIUM (OTC)
PRILOSEC, OMEPRAZOLE
PULMICORT RESPULES, BUDESONIDE
RHINOCORT, BUDESONIDE
SENSORCAINE, BUPIVACAINE HYDROCHLORIDE
SEROQUEL, QUETIAPINE FUMARATE
SYMBICORT, BUDESONIDE
TENORMIN, ATENOLOL
TOPROL-XL, METOPROLOL SUCCINATE
- * ASTRAZENECA PHARMACEUTICALS LP
ATACAND HCT, CANDESARTAN CILEXETIL
ATACAND, CANDESARTAN CILEXETIL
CRESTOR, ROSUVASTATIN CALCIUM
FASLODEX, FULVESTRANT
LEXXEL, ENALAPRIL MALEATE
NOLVADEX, TAMOXIFEN CITRATE
PLENDIL, FELODIPINE
PULMICORT, BUDESONIDE
TENORETIC 100, ATENOLOL
TENORETIC 50, ATENOLOL
TENORMIN, ATENOLOL
ZOMIG, ZOLMITRIPTAN
ZOMIG-ZMT, ZOLMITRIPTAN
- * ASTRAZENECA UK LTD
ACCOLATE, ZAFIRLUKAST
ARIMIDEX, ANASTROZOLE
CASODEX, BICALUTAMIDE
IRESSA, GEFITINIB
MERREM I.V., MEROPENEM
ZESTORETIC, HYDROCHLOROTHIAZIDE
ZESTRIL, LISINAPRIL
ZOLADEX, GOSERELIN ACETATE

ATON

- * ATON PHARMA INC
CUPRIMINE, PENICILLAMINE

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

**** A ****

* ATON PHARMA INC
DEMSE, METYROSINE
EDECRIN, ETHACRYNATE SODIUM
EDECRIN, ETHACRYNIC ACID
LACRISERT, HYDROXYPROPYL CELLULOSE
MEPHYTON, PHYTONADIONE
SYPRINE, TRIENTINE HYDROCHLORIDE

AUROBINDO

* AUROBINDO PHARMA LTD
AMOXICILLIN, AMOXICILLIN
CITALOPRAM HYDROBROMIDE, CITALOPRAM HYDROBROMIDE
LISINAPRIL AND HYDROCHLOROTHIAZIDE, HYDROCHLOROTHIAZIDE
LISINAPRIL, LISINAPRIL
METFORMIN HYDROCHLORIDE, METFORMIN HYDROCHLORIDE
MIRTAZAPINE, MIRTAZAPINE
ZIDOVUDINE, ZIDOVUDINE

AUROBINDO PHARMA

* AUROBINDO PHARMA LTD
BISOPROLOL FUMARATE, BISOPROLOL FUMARATE
MELOXICAM, MELOXICAM
SIMVASTATIN, SIMVASTATIN

AUROBINDO PHARMA LTD

* AUROBINDO PHARMA LTD INC
CEFUROXIME AXETIL, CEFUROXIME AXETIL
CEPHALEXIN, CEPHALEXIN
CITALOPRAM HYDROBROMIDE, CITALOPRAM HYDROBROMIDE
MIRTAZAPINE, MIRTAZAPINE
ZIDOVUDINE, ZIDOVUDINE

AUROSAL PHARMS

* AUROSAL PHARMACEUTICALS LLC
AMIODARONE HYDROCHLORIDE, AMIODARONE HYDROCHLORIDE

AUXILIUM PHARMS

* AUXILIUM PHARMACEUTICALS
TESTIM, TESTOSTERONE

AVANIR PHARMS

* AVANIR PHARMACEUTICALS
FAZACLO ODT, CLOZAPINE

AVEVA

* AVEVA DRUG DELIVERY SYSTEMS INC
NICOTINE, NICOTINE
PROSTEP, NICOTINE (OTC)

AXCAN SCANDIPHARM

* AXCAN SCANDIPHARM INC
BENTYL PRESERVATIVE FREE, DICYCLOMINE HYDROCHLORIDE
BENTYL, DICYCLOMINE HYDROCHLORIDE
CANASA, MESALAMINE
CARAFATE, SUCRALFATE
PHOTOFRIN, PORFIMER SODIUM
PYLERA, BISKALCITRATE
URSO 250, URSODIOL
URSO FORTE, URSODIOL

AYERST

* AYERST LABORATORIES INC
PHOSPHOLINE IODIDE, ECHOTHIOPHATE IODIDE

AZUR PHARMA

* AZUR PHARMA INTERNATIONAL LTD
GASTROCROM, CROMOLYN SODIUM

B BRAUN

* B BRAUN MEDICAL INC
ACETIC ACID 0.25% IN PLASTIC CONTAINER, ACETIC ACID, GLACIAL

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

** B **

* B BRAUN MEDICAL INC
 ALCOHOL 10% AND DEXTROSE 5%, ALCOHOL
 ALCOHOL 5% AND DEXTROSE 5%, ALCOHOL
 BRETYLIUM TOSYLATE IN DEXTROSE 5% IN PLASTIC CONTAINER, BRETYLIUM TOSYLATE
 CEFAZOLIN AND DEXTROSE, CEFAZOLIN SODIUM
 CEFOXITIN AND DEXTROSE IN DUPLEX CONTAINER, CEFOXITIN SODIUM
 CEFTRIAXONE AND DEXTROSE IN DUPLEX CONTAINER, CEFTRIAXONE SODIUM
 CEFUROXIME AND DEXTROSE IN DUPLEX CONTAINER, CEFUROXIME SODIUM
 DEXTROSE 10% AND SODIUM CHLORIDE 0.11% IN PLASTIC CONTAINER, DEXTROSE
 DEXTROSE 10% AND SODIUM CHLORIDE 0.2% IN PLASTIC CONTAINER, DEXTROSE
 DEXTROSE 10% AND SODIUM CHLORIDE 0.33% IN PLASTIC CONTAINER, DEXTROSE
 DEXTROSE 10% AND SODIUM CHLORIDE 0.45% IN PLASTIC CONTAINER, DEXTROSE
 DEXTROSE 10% AND SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER, DEXTROSE
 DEXTROSE 10% IN PLASTIC CONTAINER, DEXTROSE
 DEXTROSE 2.5% AND SODIUM CHLORIDE 0.11% IN PLASTIC CONTAINER, DEXTROSE
 DEXTROSE 2.5% AND SODIUM CHLORIDE 0.2% IN PLASTIC CONTAINER, DEXTROSE
 DEXTROSE 2.5% AND SODIUM CHLORIDE 0.33% IN PLASTIC CONTAINER, DEXTROSE
 DEXTROSE 2.5% AND SODIUM CHLORIDE 0.45% IN PLASTIC CONTAINER, DEXTROSE
 DEXTROSE 2.5% AND SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER, DEXTROSE
 DEXTROSE 2.5% IN HALF-STRENGTH LACTATED RINGER'S IN PLASTIC CONTAINER, CALCIUM CHLORIDE
 DEXTROSE 3.3% AND SODIUM CHLORIDE 0.3% IN PLASTIC CONTAINER, DEXTROSE
 DEXTROSE 5% AND SODIUM CHLORIDE 0.11% IN PLASTIC CONTAINER, DEXTROSE
 DEXTROSE 5% AND SODIUM CHLORIDE 0.2% IN PLASTIC CONTAINER, DEXTROSE
 DEXTROSE 5% AND SODIUM CHLORIDE 0.33% IN PLASTIC CONTAINER, DEXTROSE
 DEXTROSE 5% AND SODIUM CHLORIDE 0.45% IN PLASTIC CONTAINER, DEXTROSE
 DEXTROSE 5% AND SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER, DEXTROSE
 DEXTROSE 5% IN ACETATED RINGER'S IN PLASTIC CONTAINER, CALCIUM CHLORIDE
 DEXTROSE 5% IN LACTATED RINGER'S IN PLASTIC CONTAINER, CALCIUM CHLORIDE
 DEXTROSE 5% IN PLASTIC CONTAINER, DEXTROSE
 DEXTROSE 5% IN RINGER'S IN PLASTIC CONTAINER, CALCIUM CHLORIDE
 DIALYTE CONCENTRATE W/ DEXTROSE 30% IN PLASTIC CONTAINER, CALCIUM CHLORIDE
 DIALYTE CONCENTRATE W/ DEXTROSE 50% IN PLASTIC CONTAINER, CALCIUM CHLORIDE
 DIALYTE LM/ DEXTROSE 1.5% IN PLASTIC CONTAINER, CALCIUM CHLORIDE
 DIALYTE LM/ DEXTROSE 2.5% IN PLASTIC CONTAINER, CALCIUM CHLORIDE
 DIALYTE LM/ DEXTROSE 4.25% IN PLASTIC CONTAINER, CALCIUM CHLORIDE
 DOPAMINE HYDROCHLORIDE AND DEXTROSE 5% IN PLASTIC CONTAINER, DOPAMINE HYDROCHLORIDE
 DOPAMINE HYDROCHLORIDE AND DEXTROSE 5%, DOPAMINE HYDROCHLORIDE
 FREAMINE HBC 6.9%, AMINO ACIDS
 FREAMINE III 10%, AMINO ACIDS
 FREAMINE III 3% W/ ELECTROLYTES, AMINO ACIDS
 FREAMINE III 8.5% W/ ELECTROLYTES, AMINO ACIDS
 FREAMINE III 8.5%, AMINO ACIDS
 GENTAMICIN SULFATE IN SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER, GENTAMICIN SULFATE
 GLYCINE 1.5% IN PLASTIC CONTAINER, GLYCINE
 HEPARIN SODIUM 1,000 UNITS IN SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER, HEPARIN SODIUM
 HEPARIN SODIUM 20,000 UNITS IN DEXTROSE 5% IN PLASTIC CONTAINER, HEPARIN SODIUM
 HEPARIN SODIUM 25,000 UNITS IN DEXTROSE 5% IN PLASTIC CONTAINER, HEPARIN SODIUM
 HEPATAMINE 8%, AMINO ACIDS
 ISOLYTE E IN DEXTROSE 5% IN PLASTIC CONTAINER, CALCIUM CHLORIDE
 ISOLYTE E IN PLASTIC CONTAINER, CALCIUM CHLORIDE
 ISOLYTE H IN DEXTROSE 5% IN PLASTIC CONTAINER, DEXTROSE
 ISOLYTE M IN DEXTROSE 5% IN PLASTIC CONTAINER, DEXTROSE
 ISOLYTE P IN DEXTROSE 5% IN PLASTIC CONTAINER, DEXTROSE
 ISOLYTE R IN DEXTROSE 5% IN PLASTIC CONTAINER, CALCIUM CHLORIDE
 ISOLYTE S IN DEXTROSE 5% IN PLASTIC CONTAINER, DEXTROSE
 ISOLYTE S IN PLASTIC CONTAINER, MAGNESIUM CHLORIDE
 ISOLYTE S PH 7.4 IN PLASTIC CONTAINER, MAGNESIUM CHLORIDE
 LACTATED RINGER'S IN PLASTIC CONTAINER, CALCIUM CHLORIDE
 LIDOCAINE HYDROCHLORIDE 0.2% AND DEXTROSE 5% IN PLASTIC CONTAINER, LIDOCAINE
 HYDROCHLORIDE
 LIDOCAINE HYDROCHLORIDE 0.4% AND DEXTROSE 5% IN PLASTIC CONTAINER, LIDOCAINE
 HYDROCHLORIDE
 LIDOCAINE HYDROCHLORIDE 0.8% AND DEXTROSE 5% IN PLASTIC CONTAINER, LIDOCAINE
 HYDROCHLORIDE

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

** B **

* B BRAUN MEDICAL INC
 MANNITOL 10% IN PLASTIC CONTAINER, MANNITOL
 MANNITOL 10% W/ DEXTROSE 5% IN DISTILLED WATER, MANNITOL
 MANNITOL 10%, MANNITOL
 MANNITOL 15% IN PLASTIC CONTAINER, MANNITOL
 MANNITOL 15% W/ DEXTROSE 5% IN SODIUM CHLORIDE 0.45%, MANNITOL
 MANNITOL 15%, MANNITOL
 MANNITOL 20% IN PLASTIC CONTAINER, MANNITOL
 MANNITOL 20%, MANNITOL
 MANNITOL 5% IN PLASTIC CONTAINER, MANNITOL
 MANNITOL 5% W/ DEXTROSE 5% IN SODIUM CHLORIDE 0.12%, MANNITOL
 MANNITOL 5%, MANNITOL
 METRO I.V. IN PLASTIC CONTAINER, METRONIDAZOLE
 MILRINONE LACTATE IN DEXTROSE 5% IN PLASTIC CONTAINER, MILRINONE LACTATE
 NEPHRAMINE 5.4%, AMINO ACIDS
 NUTRILIPID 10%, SOYBEAN OIL
 NUTRILIPID 20%, SOYBEAN OIL
 PHYSIOLYTE IN PLASTIC CONTAINER, MAGNESIUM CHLORIDE
 POTASSIUM CHLORIDE 0.037% IN DEXTROSE 10% AND SODIUM CHLORIDE 0.2% IN PLASTIC CONTAINER, DEXTROSE
 POTASSIUM CHLORIDE 0.037% IN DEXTROSE 10% AND SODIUM CHLORIDE 0.45% IN PLASTIC CONTAINER, DEXTROSE
 POTASSIUM CHLORIDE 0.037% IN DEXTROSE 10% AND SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER, DEXTROSE
 POTASSIUM CHLORIDE 0.037% IN DEXTROSE 5% AND SODIUM CHLORIDE 0.11% IN PLASTIC CONTAINER, DEXTROSE
 POTASSIUM CHLORIDE 0.037% IN DEXTROSE 5% AND SODIUM CHLORIDE 0.2% IN PLASTIC CONTAINER, DEXTROSE
 POTASSIUM CHLORIDE 0.037% IN DEXTROSE 5% AND SODIUM CHLORIDE 0.33% IN PLASTIC CONTAINER, DEXTROSE
 POTASSIUM CHLORIDE 0.037% IN DEXTROSE 5% AND SODIUM CHLORIDE 0.45% IN PLASTIC CONTAINER, DEXTROSE
 POTASSIUM CHLORIDE 0.037% IN DEXTROSE 5% AND SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER, DEXTROSE
 POTASSIUM CHLORIDE 0.075% IN DEXTROSE 10% AND SODIUM CHLORIDE 0.2% IN PLASTIC CONTAINER, DEXTROSE
 POTASSIUM CHLORIDE 0.075% IN DEXTROSE 10% AND SODIUM CHLORIDE 0.45% IN PLASTIC CONTAINER, DEXTROSE
 POTASSIUM CHLORIDE 0.075% IN DEXTROSE 10% AND SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER, DEXTROSE
 POTASSIUM CHLORIDE 0.075% IN DEXTROSE 3.3% AND SODIUM CHLORIDE 0.3% IN PLASTIC CONTAINER, DEXTROSE
 POTASSIUM CHLORIDE 0.075% IN DEXTROSE 5% AND SODIUM CHLORIDE 0.11% IN PLASTIC CONTAINER, DEXTROSE
 POTASSIUM CHLORIDE 0.075% IN DEXTROSE 5% AND SODIUM CHLORIDE 0.2% IN PLASTIC CONTAINER, DEXTROSE
 POTASSIUM CHLORIDE 0.075% IN DEXTROSE 5% AND SODIUM CHLORIDE 0.33% IN PLASTIC CONTAINER, DEXTROSE
 POTASSIUM CHLORIDE 0.075% IN DEXTROSE 5% AND SODIUM CHLORIDE 0.45% IN PLASTIC CONTAINER, DEXTROSE
 POTASSIUM CHLORIDE 0.075% IN DEXTROSE 5% AND SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER, DEXTROSE
 POTASSIUM CHLORIDE 0.075% IN DEXTROSE 5% IN PLASTIC CONTAINER, DEXTROSE
 POTASSIUM CHLORIDE 0.11% IN DEXTROSE 10% AND SODIUM CHLORIDE 0.2% IN PLASTIC CONTAINER, DEXTROSE
 POTASSIUM CHLORIDE 0.11% IN DEXTROSE 10% AND SODIUM CHLORIDE 0.45% IN PLASTIC CONTAINER, DEXTROSE
 POTASSIUM CHLORIDE 0.11% IN DEXTROSE 10% AND SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER, DEXTROSE
 POTASSIUM CHLORIDE 0.11% IN DEXTROSE 3.3% AND SODIUM CHLORIDE 0.3% IN PLASTIC CONTAINER, DEXTROSE
 POTASSIUM CHLORIDE 0.11% IN DEXTROSE 5% AND SODIUM CHLORIDE 0.11% IN PLASTIC CONTAINER, DEXTROSE
 POTASSIUM CHLORIDE 0.11% IN DEXTROSE 5% AND SODIUM CHLORIDE 0.2% IN PLASTIC CONTAINER, DEXTROSE
 POTASSIUM CHLORIDE 0.11% IN DEXTROSE 5% AND SODIUM CHLORIDE 0.33% IN PLASTIC CONTAINER, DEXTROSE

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

** B **

* B BRAUN MEDICAL INC
 POTASSIUM CHLORIDE 0.11% IN DEXTROSE 5% AND SODIUM CHLORIDE 0.45% IN PLASTIC CONTAINER, DEXTROSE
 POTASSIUM CHLORIDE 0.11% IN DEXTROSE 5% AND SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER, DEXTROSE
 POTASSIUM CHLORIDE 0.15% IN DEXTROSE 10% AND SODIUM CHLORIDE 0.2% IN PLASTIC CONTAINER, DEXTROSE
 POTASSIUM CHLORIDE 0.15% IN DEXTROSE 10% AND SODIUM CHLORIDE 0.45% IN PLASTIC CONTAINER, DEXTROSE
 POTASSIUM CHLORIDE 0.15% IN DEXTROSE 10% AND SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER, DEXTROSE
 POTASSIUM CHLORIDE 0.15% IN DEXTROSE 3.3% AND SODIUM CHLORIDE 0.3% IN PLASTIC CONTAINER, DEXTROSE
 POTASSIUM CHLORIDE 0.15% IN DEXTROSE 5% AND SODIUM CHLORIDE 0.11% IN PLASTIC CONTAINER, DEXTROSE
 POTASSIUM CHLORIDE 0.15% IN DEXTROSE 5% AND SODIUM CHLORIDE 0.2% IN PLASTIC CONTAINER, DEXTROSE
 POTASSIUM CHLORIDE 0.15% IN DEXTROSE 5% AND SODIUM CHLORIDE 0.33% IN PLASTIC CONTAINER, DEXTROSE
 POTASSIUM CHLORIDE 0.15% IN DEXTROSE 5% AND SODIUM CHLORIDE 0.45% IN PLASTIC CONTAINER, DEXTROSE
 POTASSIUM CHLORIDE 0.15% IN DEXTROSE 5% AND SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER, DEXTROSE
 POTASSIUM CHLORIDE 0.15% IN DEXTROSE 5% IN PLASTIC CONTAINER, DEXTROSE
 POTASSIUM CHLORIDE 0.15% IN SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER, POTASSIUM CHLORIDE
 POTASSIUM CHLORIDE 0.22% IN DEXTROSE 10% AND SODIUM CHLORIDE 0.2% IN PLASTIC CONTAINER, DEXTROSE
 POTASSIUM CHLORIDE 0.22% IN DEXTROSE 10% AND SODIUM CHLORIDE 0.45% IN PLASTIC CONTAINER, DEXTROSE
 POTASSIUM CHLORIDE 0.22% IN DEXTROSE 10% AND SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER, DEXTROSE
 POTASSIUM CHLORIDE 0.22% IN DEXTROSE 3.3% AND SODIUM CHLORIDE 0.3% IN PLASTIC CONTAINER, DEXTROSE
 POTASSIUM CHLORIDE 0.22% IN DEXTROSE 5% AND SODIUM CHLORIDE 0.11% IN PLASTIC CONTAINER, DEXTROSE
 POTASSIUM CHLORIDE 0.22% IN DEXTROSE 5% AND SODIUM CHLORIDE 0.2% IN PLASTIC CONTAINER, DEXTROSE
 POTASSIUM CHLORIDE 0.22% IN DEXTROSE 5% AND SODIUM CHLORIDE 0.33% IN PLASTIC CONTAINER, DEXTROSE
 POTASSIUM CHLORIDE 0.22% IN DEXTROSE 5% AND SODIUM CHLORIDE 0.45% IN PLASTIC CONTAINER, DEXTROSE
 POTASSIUM CHLORIDE 0.22% IN DEXTROSE 5% AND SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER, DEXTROSE
 POTASSIUM CHLORIDE 0.22% IN DEXTROSE 5% IN PLASTIC CONTAINER, DEXTROSE
 POTASSIUM CHLORIDE 0.3% IN DEXTROSE 10% AND SODIUM CHLORIDE 0.2% IN PLASTIC CONTAINER, DEXTROSE
 POTASSIUM CHLORIDE 0.3% IN DEXTROSE 10% AND SODIUM CHLORIDE 0.45% IN PLASTIC CONTAINER, DEXTROSE
 POTASSIUM CHLORIDE 0.3% IN DEXTROSE 10% AND SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER, DEXTROSE
 POTASSIUM CHLORIDE 0.3% IN DEXTROSE 3.3% AND SODIUM CHLORIDE 0.3% IN PLASTIC CONTAINER, DEXTROSE
 POTASSIUM CHLORIDE 0.3% IN DEXTROSE 5% AND SODIUM CHLORIDE 0.11% IN PLASTIC CONTAINER, DEXTROSE
 POTASSIUM CHLORIDE 0.3% IN DEXTROSE 5% AND SODIUM CHLORIDE 0.2% IN PLASTIC CONTAINER, DEXTROSE
 POTASSIUM CHLORIDE 0.3% IN DEXTROSE 5% AND SODIUM CHLORIDE 0.33% IN PLASTIC CONTAINER, DEXTROSE
 POTASSIUM CHLORIDE 0.3% IN DEXTROSE 5% AND SODIUM CHLORIDE 0.45% IN PLASTIC CONTAINER, DEXTROSE
 POTASSIUM CHLORIDE 0.3% IN DEXTROSE 5% AND SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER, DEXTROSE
 POTASSIUM CHLORIDE 0.3% IN DEXTROSE 5% IN PLASTIC CONTAINER, DEXTROSE
 POTASSIUM CHLORIDE, POTASSIUM CHLORIDE
 PROCALAMINE, AMINO ACIDS
 RESECTISOL IN PLASTIC CONTAINER, MANNITOL
 RINGER'S IN PLASTIC CONTAINER, CALCIUM CHLORIDE
 SODIUM CHLORIDE 0.45% IN PLASTIC CONTAINER, SODIUM CHLORIDE
 SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER, SODIUM CHLORIDE

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

**** B ****

* B BRAUN MEDICAL INC
SODIUM LACTATE 1/6 MOLAR IN PLASTIC CONTAINER, SODIUM LACTATE
SORBITOL 3.3% IN PLASTIC CONTAINER, SORBITOL
STERILE WATER FOR INJECTION IN PLASTIC CONTAINER, WATER FOR INJECTION, STERILE
STERILE WATER IN PLASTIC CONTAINER, WATER FOR IRRIGATION, STERILE
THEOPHYLLINE 0.04% AND DEXTROSE 5% IN PLASTIC CONTAINER, THEOPHYLLINE
THEOPHYLLINE 0.08% AND DEXTROSE 5% IN PLASTIC CONTAINER, THEOPHYLLINE
THEOPHYLLINE 0.16% AND DEXTROSE 5% IN PLASTIC CONTAINER, THEOPHYLLINE
THEOPHYLLINE 0.32% AND DEXTROSE 5% IN PLASTIC CONTAINER, THEOPHYLLINE
TROPHAMINE 10%, AMINO ACIDS
TROPHAMINE, AMINO ACIDS

BAKER NORTON

* BAKER NORTON PHARMACEUTICALS INC
PROGLYCEM, DIAZOXIDE

BALLARD MEDCL

* BALLARD MEDICAL PRODUCTS INC
PYTEST KIT, UREA, C-14
PYTEST, UREA, C-14

BANNER PHARMACAPS

* BANNER PHARMACAPS INC
BENZONATATE, BENZONATATE
CLOFIBRATE, CLOFIBRATE
ETHOSUXIMIDE, ETHOSUXIMIDE
IBUPROFEN, IBUPROFEN (OTC)
LOPERAMIDE HYDROCHLORIDE, LOPERAMIDE HYDROCHLORIDE (OTC)
NAPROXEN SODIUM, NAPROXEN SODIUM (OTC)
VALPROIC ACID, VALPROIC ACID
VITAMIN D, ERGOCALCIFEROL
ZONISAMIDE, ZONISAMIDE

BARR

* BARR LABORATORIES INC
ACETOHEXAMIDE, ACETOHEXAMIDE
ALPRAZOLAM, ALPRAZOLAM
AMILORIDE HYDROCHLORIDE AND HYDROCHLOROTHIAZIDE, AMILORIDE HYDROCHLORIDE
AMIODARONE HYDROCHLORIDE, AMIODARONE HYDROCHLORIDE
ANAGRELIDE HYDROCHLORIDE, ANAGRELIDE HYDROCHLORIDE
ARANELLE, ETHINYL ESTRADIOL
BALZIVA-21, ETHINYL ESTRADIOL
BALZIVA-28, ETHINYL ESTRADIOL
CAMILA, NORETHINDRONE
CEPHALEXIN, CEPHALEXIN
CHLORDIAZEPOXIDE HYDROCHLORIDE, CHLORDIAZEPOXIDE HYDROCHLORIDE
CHLORZOXAZONE, CHLORZOXAZONE
CIPROFLOXACIN, CIPROFLOXACIN HYDROCHLORIDE
CLARAVIS, ISOTRETINOIN
CLONAZEPAM, CLONAZEPAM
DANAZOL, DANAZOL
DEMECLOCYCLINE HYDROCHLORIDE, DEMECLOCYCLINE HYDROCHLORIDE
DESMOPRESSIN ACETATE, DESMOPRESSIN ACETATE
DEXTROAMP SACCHARATE, AMP ASPARTATE, DEXTROAMP SULFATE AND AMP SULFATE, AMPHETAMINE
ASPARTATE
DEXTROAMPHETAMINE SULFATE, DEXTROAMPHETAMINE SULFATE
DIAZEPAM, DIAZEPAM
DIDANOSINE, DIDANOSINE
DIPHENHYDRAMINE HYDROCHLORIDE, DIPHENHYDRAMINE HYDROCHLORIDE
DIPYRIDAMOLE, DIPYRIDAMOLE
E-BASE, ERYTHROMYCIN
ERRIN, NORETHINDRONE
ERYTHROMYCIN ETHYLSUCCINATE AND SULFISOXAZOLE ACETYL, ERYTHROMYCIN ETHYLSUCCINATE
ERYTHROMYCIN ETHYLSUCCINATE, ERYTHROMYCIN ETHYLSUCCINATE
ERYTHROMYCIN STEARATE, ERYTHROMYCIN STEARATE
ERYTHROMYCIN, ERYTHROMYCIN

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

**** B ****

* BARR LABORATORIES INC
ESTRADIOL AND NORGESTIMATE, ESTRADIOL
ESTRADIOL, ESTRADIOL
ESTROPIPATE, ESTROPIPATE
ETHAMBUTOL HYDROCHLORIDE, ETHAMBUTOL HYDROCHLORIDE
EXTENDED PHENYTOIN SODIUM, PHENYTOIN SODIUM
FEXOFENADINE HYDROCHLORIDE AND PSEUDOEPHEDRINE HYDROCHLORIDE, FEXOFENADINE HYDROCHLORIDE
FEXOFENADINE HYDROCHLORIDE, FEXOFENADINE HYDROCHLORIDE
FLECAINIDE ACETATE, FLECAINIDE ACETATE
FLUDROCORTISONE ACETATE, FLUDROCORTISONE ACETATE
FLUOXETINE HYDROCHLORIDE, FLUOXETINE HYDROCHLORIDE
FLUOXETINE, FLUOXETINE HYDROCHLORIDE
FLUTAMIDE, FLUTAMIDE
FLUVOXAMINE MALEATE, FLUVOXAMINE MALEATE
GENCEPT 10/11-21, ETHINYL ESTRADIOL
GENCEPT 10/11-28, ETHINYL ESTRADIOL
HYDROCODONE BITARTRATE AND ACETAMINOPHEN, ACETAMINOPHEN
HYDROXYUREA, HYDROXYUREA
HYDROXYZINE PAMOATE, HYDROXYZINE PAMOATE
ISONIAZID, ISONIAZID
JUNEL 1.5/30, ETHINYL ESTRADIOL
JUNEL 1/20, ETHINYL ESTRADIOL
JUNEL FE 1.5/30, ETHINYL ESTRADIOL
JUNEL FE 1/20, ETHINYL ESTRADIOL
KARIVA, DESOGESTREL
KELNOR, ETHINYL ESTRADIOL
LEFLUNOMIDE, LEFLUNOMIDE
LESSINA-28, ETHINYL ESTRADIOL
LEVONORGESTREL AND ETHINYL ESTRADIOL, ETHINYL ESTRADIOL
LITHIUM CARBONATE, LITHIUM CARBONATE
MEDROXYPROGESTERONE ACETATE, MEDROXYPROGESTERONE ACETATE
MEFLOQUINE, MEFLOQUINE HYDROCHLORIDE
MEGESTROL ACETATE, MEGESTROL ACETATE
MEPERIDINE HYDROCHLORIDE, MEPERIDINE HYDROCHLORIDE
METFORMIN HYDROCHLORIDE, METFORMIN HYDROCHLORIDE
METHOTREXATE SODIUM, METHOTREXATE SODIUM
MIRTAZAPINE, MIRTAZAPINE
NALTREXONE HYDROCHLORIDE, NALTREXONE HYDROCHLORIDE
NICARDIPINE HYDROCHLORIDE, NICARDIPINE HYDROCHLORIDE
NORETHINDRONE ACETATE, NORETHINDRONE ACETATE
NORTREL 0.5/35-21, ETHINYL ESTRADIOL
NORTREL 0.5/35-28, ETHINYL ESTRADIOL
NORTREL 1/35-21, ETHINYL ESTRADIOL
NORTREL 1/35-28, ETHINYL ESTRADIOL
NORTREL 7/7/7, ETHINYL ESTRADIOL
OXYCODONE AND ACETAMINOPHEN, ACETAMINOPHEN
PORTIA-28, ETHINYL ESTRADIOL
PROPRANOLOL HYDROCHLORIDE AND HYDROCHLOROTHIAZIDE, HYDROCHLOROTHIAZIDE
PYRIDOSTIGMINE BROMIDE, PYRIDOSTIGMINE BROMIDE
SPRINTEC, ETHINYL ESTRADIOL
SULFINPYRAZONE, SULFINPYRAZONE
TAMOXIFEN CITRATE, TAMOXIFEN CITRATE
TETRACYCLINE HYDROCHLORIDE, TETRACYCLINE HYDROCHLORIDE
TIZANIDINE HYDROCHLORIDE, TIZANIDINE HYDROCHLORIDE
TRAMADOL HYDROCHLORIDE AND ACETAMINOPHEN, ACETAMINOPHEN
TRAZODONE HYDROCHLORIDE, TRAZODONE HYDROCHLORIDE
TREXALL, METHOTREXATE SODIUM
TRIAMTERENE AND HYDROCHLOROTHIAZIDE, HYDROCHLOROTHIAZIDE
TRI-SPRINTEC, ETHINYL ESTRADIOL
WARFARIN SODIUM, WARFARIN SODIUM
ZONISAMIDE, ZONISAMIDE

* BARR PHARMACEUTICALS
LEUCOVORIN CALCIUM, LEUCOVORIN CALCIUM

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

** B **

BARRIER

- * BARRIER THERAPEUTICS
SOLAGE, MEQUINOL
VUSION, MICONAZOLE NITRATE

BARRIER THERAP

- * BARRIER THERAPEUTICS INC
XOLEGEL, KETOCONAZOLE

BARTOR

- * BARTOR PHARMACAL CO
TESTOSTERONE, TESTOSTERONE

BASF

- * BASF CORP
IBUPROFEN, IBUPROFEN
IBUPROFEN, IBUPROFEN (OTC)
LOPURIN, ALLOPURINOL
SSD AF, SILVER SULFADIAZINE
SSD, SILVER SULFADIAZINE

BAUSCH AND LOMB

- * BAUSCH AND LOMB INC
ALBUTEROL SULFATE, ALBUTEROL SULFATE
ALREX, LOTEPIREDNOL ETABONATE
CARTEOLOL HYDROCHLORIDE, CARTEOLOL HYDROCHLORIDE
FLURBIPROFEN SODIUM, FLURBIPROFEN SODIUM
IPRATROPIUM BROMIDE, IPRATROPIUM BROMIDE
LOTEMAX, LOTEPIREDNOL ETABONATE
OFLOXACIN, OFLOXACIN
OPCON-A, NAPHAZOLINE HYDROCHLORIDE (OTC)
RETISERT, FLUOCINOLONE ACETONIDE
TIMOLOL MALEATE, TIMOLOL MALEATE
VITRASERT, GANCICLOVIR
ZYLET, LOTEPIREDNOL ETABONATE
- * BAUSCH AND LOMB PHARMACEUTICALS INC
ACETIC ACID 2% IN AQUEOUS ALUMINUM ACETATE, ACETIC ACID, GLACIAL
BACITRACIN ZINC AND POLYMYXIN B SULFATE, BACITRACIN ZINC
BETAXOLOL HYDROCHLORIDE, BETAXOLOL HYDROCHLORIDE
BRIMONIDINE TARTRATE, BRIMONIDINE TARTRATE
CIPROFLOXACIN, CIPROFLOXACIN HYDROCHLORIDE
CROLOM, CROMOLYN SODIUM
CROMOLYN SODIUM, CROMOLYN SODIUM
CROMOLYN SODIUM, CROMOLYN SODIUM (OTC)
DESMOPRESSIN ACETATE, DESMOPRESSIN ACETATE
DEXAMETHASONE SODIUM PHOSPHATE, DEXAMETHASONE SODIUM PHOSPHATE
DEXASPORIN, DEXAMETHASONE
DIPIVEFRIN HYDROCHLORIDE, DIPIVEFRIN HYDROCHLORIDE
ERYTHROMYCIN, ERYTHROMYCIN
FLUNISOLIDE, FLUNISOLIDE
GENTAMICIN SULFATE, GENTAMICIN SULFATE
IPRATROPIUM BROMIDE, IPRATROPIUM BROMIDE
LEVOBUNOLOL HYDROCHLORIDE, LEVOBUNOLOL HYDROCHLORIDE
MINOXIDIL (FOR MEN), MINOXIDIL (OTC)
NAFAZAIR, NAPHAZOLINE HYDROCHLORIDE
NEOMYCIN AND POLYMYXIN B SULFATES AND BACITRACIN ZINC, BACITRACIN ZINC
NEOMYCIN AND POLYMYXIN B SULFATES AND DEXAMETHASONE, DEXAMETHASONE
NEOMYCIN AND POLYMYXIN B SULFATES AND GRAMICIDIN, GRAMICIDIN
NEOMYCIN AND POLYMYXIN B SULFATES AND HYDROCORTISONE, HYDROCORTISONE
NEOMYCIN AND POLYMYXIN B SULFATES, BACITRACIN ZINC AND HYDROCORTISONE, BACITRACIN ZINC
NYSTATIN, NYSTATIN
OPTIPRANOLOL, METIPRANOLOL HYDROCHLORIDE
OTICAIR, HYDROCORTISONE
PENTOLAIR, CYCLOPENTOLATE HYDROCHLORIDE
PREDNISOLONE SODIUM PHOSPHATE, PREDNISOLONE SODIUM PHOSPHATE
PROPACARINE HYDROCHLORIDE, PROPACARINE HYDROCHLORIDE

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

** B **

* BAUSCH AND LOMB PHARMACEUTICALS INC
 SULFACETAMIDE SODIUM AND PREDNISOLONE SODIUM PHOSPHATE, PREDNISOLONE SODIUM PHOSPHATE
 SULFACETAMIDE SODIUM, SULFACETAMIDE SODIUM
 TIMOLOL MALEATE, TIMOLOL MALEATE
 TOBRAMYCIN AND DEXAMETHASONE, DEXAMETHASONE
 TOBRAMYCIN, TOBRAMYCIN
 TRIMETHOPRIM SULFATE AND POLYMYXIN B SULFATE, POLYMYXIN B SULFATE
 TROPICAMIDE, TROPICAMIDE

BAXTER HLTHCARE

* BAXTER HEALTHCARE CORP
 ACETIC ACID 0.25% IN PLASTIC CONTAINER, ACETIC ACID, GLACIAL
 ADENOSINE, ADENOSINE
 AMINOACETIC ACID 1.5% IN PLASTIC CONTAINER, GLYCINE
 ANCEF IN PLASTIC CONTAINER, CEFAZOLIN SODIUM
 BACTOCILL IN PLASTIC CONTAINER, OXACILLIN SODIUM
 BRANCHAMIN 4% IN PLASTIC CONTAINER, AMINO ACIDS
 CARDIOPLEGIC IN PLASTIC CONTAINER, CALCIUM CHLORIDE
 CEFTRIAXONE IN PLASTIC CONTAINER, CEFTRIAXONE SODIUM
 CLINIMIX 2.75/10 SULFITE FREE IN DEXTROSE 10% IN PLASTIC CONTAINER, AMINO ACIDS
 CLINIMIX 2.75/25 SULFITE FREE IN DEXTROSE 25% IN PLASTIC CONTAINER, AMINO ACIDS
 CLINIMIX 2.75/5 SULFITE FREE IN DEXTROSE 5% IN PLASTIC CONTAINER, AMINO ACIDS
 CLINIMIX 4.25/10 SULFITE FREE IN DEXTROSE 10% IN PLASTIC CONTAINER, AMINO ACIDS
 CLINIMIX 4.25/20 SULFITE FREE IN DEXTROSE 20% IN PLASTIC CONTAINER, AMINO ACIDS
 CLINIMIX 4.25/25 SULFITE FREE IN DEXTROSE 25% IN PLASTIC CONTAINER, AMINO ACIDS
 CLINIMIX 4.25/5 SULFITE FREE IN DEXTROSE 5% IN PLASTIC CONTAINER, AMINO ACIDS
 CLINIMIX 5/10 SULFITE FREE IN DEXTROSE 10% IN PLASTIC CONTAINER, AMINO ACIDS
 CLINIMIX 5/15 SULFITE FREE IN DEXTROSE 15% IN PLASTIC CONTAINER, AMINO ACIDS
 CLINIMIX 5/20 SULFITE FREE IN DEXTROSE 20% IN PLASTIC CONTAINER, AMINO ACIDS
 CLINIMIX 5/25 SULFITE FREE IN DEXTROSE 25% IN PLASTIC CONTAINER, AMINO ACIDS
 CLINIMIX 5/35 SULFITE FREE IN DEXTROSE 35% IN PLASTIC CONTAINER, AMINO ACIDS
 CLINIMIX E 2.75/10 SULFITE-FREE W/ ELECT IN DEXTROSE 10% W/ CALCIUM IN PLASTIC CONTAINER, AMINO ACIDS
 CLINIMIX E 2.75/25 SULFITE-FREE W/ ELECT IN DEXTROSE 25% W/ CALCIUM IN PLASTIC CONTAINER, AMINO ACIDS
 CLINIMIX E 2.75/5 SULFITE-FREE W/ ELECT IN DEXTROSE 5% W/ CALCIUM IN PLASTIC CONTAINER, AMINO ACIDS
 CLINIMIX E 4.25/10 SULFITE-FREE W/ ELECT IN DEXTROSE 10% W/ CALCIUM IN PLASTIC CONTAINER, AMINO ACIDS
 CLINIMIX E 4.25/20 SULFITE-FREE W/ ELECT IN DEXTROSE 20% W/ CALCIUM IN PLASTIC CONTAINER, AMINO ACIDS
 CLINIMIX E 4.25/25 SULFITE-FREE W/ ELECT IN DEXTROSE 25% W/ CALCIUM IN PLASTIC CONTAINER, AMINO ACIDS
 CLINIMIX E 4.25/5 SULFITE-FREE W/ ELECT IN DEXTROSE 5% W/ CALCIUM IN PLASTIC CONTAINER, AMINO ACIDS
 CLINIMIX E 5/10 SULFITE-FREE W/ ELECT IN DEXTROSE 10% W/ CALCIUM IN PLASTIC CONTAINER, AMINO ACIDS
 CLINIMIX E 5/15 SULFITE-FREE W/ ELECT IN DEXTROSE 15% W/ CALCIUM IN PLASTIC CONTAINER, AMINO ACIDS
 CLINIMIX E 5/20 SULFITE-FREE W/ ELECT IN 20% DEXTROSE W/ CALCIUM IN PLASTIC CONTAINER, AMINO ACIDS
 CLINIMIX E 5/25 SULFITE-FREE W/ ELECT IN DEXTROSE 25% W/ CALCIUM IN PLASTIC CONTAINER, AMINO ACIDS
 CLINIMIX E 5/35 SULFITE-FREE W/ ELECT IN DEXTROSE 35% W/ CALCIUM IN PLASTIC CONTAINER, AMINO ACIDS
 CYCLOPHOSPHAMIDE, CYCLOPHOSPHAMIDE
 DEXTROSE 10% AND SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER, DEXTROSE
 DEXTROSE 10% IN PLASTIC CONTAINER, DEXTROSE
 DEXTROSE 2.5% AND SODIUM CHLORIDE 0.45% IN PLASTIC CONTAINER, DEXTROSE
 DEXTROSE 20% IN PLASTIC CONTAINER, DEXTROSE
 DEXTROSE 30% IN PLASTIC CONTAINER, DEXTROSE
 DEXTROSE 40% IN PLASTIC CONTAINER, DEXTROSE
 DEXTROSE 5% AND ELECTROLYTE NO 75 IN PLASTIC CONTAINER, DEXTROSE
 DEXTROSE 5% AND ELECTROLYTE NO.48 IN PLASTIC CONTAINER, DEXTROSE
 DEXTROSE 5% IN PLASTIC CONTAINER, DEXTROSE
 DEXTROSE 5% IN RINGER'S IN PLASTIC CONTAINER, CALCIUM CHLORIDE
 DEXTROSE 5% IN SODIUM CHLORIDE 0.2% IN PLASTIC CONTAINER, DEXTROSE

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

**** B ****

* BAXTER HEALTHCARE CORP
DEXTROSE 5% IN SODIUM CHLORIDE 0.33% IN PLASTIC CONTAINER, DEXTROSE
DEXTROSE 5% IN SODIUM CHLORIDE 0.45% IN PLASTIC CONTAINER, DEXTROSE
DEXTROSE 5% IN SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER, DEXTROSE
DEXTROSE 5%, SODIUM CHLORIDE 0.2% AND POTASSIUM CHLORIDE 10MEQ, DEXTROSE
DEXTROSE 5%, SODIUM CHLORIDE 0.2% AND POTASSIUM CHLORIDE 15MEQ (K), DEXTROSE
DEXTROSE 5%, SODIUM CHLORIDE 0.2% AND POTASSIUM CHLORIDE 20MEQ (K), DEXTROSE
DEXTROSE 5%, SODIUM CHLORIDE 0.2% AND POTASSIUM CHLORIDE 20MEQ, DEXTROSE
DEXTROSE 5%, SODIUM CHLORIDE 0.2% AND POTASSIUM CHLORIDE 30MEQ, DEXTROSE
DEXTROSE 5%, SODIUM CHLORIDE 0.2% AND POTASSIUM CHLORIDE 40MEQ, DEXTROSE
DEXTROSE 5%, SODIUM CHLORIDE 0.2% AND POTASSIUM CHLORIDE 5MEQ (K), DEXTROSE
DEXTROSE 5%, SODIUM CHLORIDE 0.2% AND POTASSIUM CHLORIDE 5MEQ, DEXTROSE
DEXTROSE 5%, SODIUM CHLORIDE 0.33% AND POTASSIUM CHLORIDE 10MEQ IN PLASTIC CONTAINER, DEXTROSE
DEXTROSE 5%, SODIUM CHLORIDE 0.33% AND POTASSIUM CHLORIDE 15MEQ IN PLASTIC CONTAINER, DEXTROSE
DEXTROSE 5%, SODIUM CHLORIDE 0.33% AND POTASSIUM CHLORIDE 20MEQ IN PLASTIC CONTAINER, DEXTROSE
DEXTROSE 5%, SODIUM CHLORIDE 0.33% AND POTASSIUM CHLORIDE 30MEQ IN PLASTIC CONTAINER, DEXTROSE
DEXTROSE 5%, SODIUM CHLORIDE 0.33% AND POTASSIUM CHLORIDE 40MEQ IN PLASTIC CONTAINER, DEXTROSE
DEXTROSE 5%, SODIUM CHLORIDE 0.33% AND POTASSIUM CHLORIDE 5MEQ IN PLASTIC CONTAINER, DEXTROSE
DEXTROSE 5%, SODIUM CHLORIDE 0.45% AND POTASSIUM CHLORIDE 20MEQ (K) IN PLASTIC CONTAINER, DEXTROSE
DEXTROSE 50% IN PLASTIC CONTAINER, DEXTROSE
DEXTROSE 60% IN PLASTIC CONTAINER, DEXTROSE
DEXTROSE 70% IN PLASTIC CONTAINER, DEXTROSE
DIANEAL 137 W/ DEXTROSE 1.5% IN PLASTIC CONTAINER, CALCIUM CHLORIDE
DIANEAL 137 W/ DEXTROSE 2.5% IN PLASTIC CONTAINER, CALCIUM CHLORIDE
DIANEAL 137 W/ DEXTROSE 4.25% IN PLASTIC CONTAINER, CALCIUM CHLORIDE
DIANEAL LOW CALCIUM W/ DEXTROSE 1.5% IN PLASTIC CONTAINER, CALCIUM CHLORIDE
DIANEAL LOW CALCIUM W/ DEXTROSE 2.5% IN PLASTIC CONTAINER, CALCIUM CHLORIDE
DIANEAL LOW CALCIUM W/ DEXTROSE 3.5% IN PLASTIC CONTAINER, CALCIUM CHLORIDE
DIANEAL LOW CALCIUM W/ DEXTROSE 4.25% IN PLASTIC CONTAINER, CALCIUM CHLORIDE
DIANEAL PD-1 W/ DEXTROSE 1.5% IN PLASTIC CONTAINER, CALCIUM CHLORIDE
DIANEAL PD-1 W/ DEXTROSE 2.5% IN PLASTIC CONTAINER, CALCIUM CHLORIDE
DIANEAL PD-1 W/ DEXTROSE 3.5% IN PLASTIC CONTAINER, CALCIUM CHLORIDE
DIANEAL PD-1 W/ DEXTROSE 4.25% IN PLASTIC CONTAINER, CALCIUM CHLORIDE
DIANEAL PD-2 W/ DEXTROSE 1.5% IN PLASTIC CONTAINER, CALCIUM CHLORIDE
DIANEAL PD-2 W/ DEXTROSE 2.5% IN PLASTIC CONTAINER, CALCIUM CHLORIDE
DIANEAL PD-2 W/ DEXTROSE 3.5% IN PLASTIC CONTAINER, CALCIUM CHLORIDE
DIANEAL PD-2 W/ DEXTROSE 4.25% IN PLASTIC CONTAINER, CALCIUM CHLORIDE
DOBUTAMINE HYDROCHLORIDE IN DEXTROSE 5% IN PLASTIC CONTAINER, DOBUTAMINE HYDROCHLORIDE
DOPAMINE HYDROCHLORIDE IN DEXTROSE 5% IN PLASTIC CONTAINER, DOPAMINE HYDROCHLORIDE
EXTRANEAL, ICODEXTRIN
FAMOTIDINE PRESERVATIVE FREE, FAMOTIDINE
FAMOTIDINE, FAMOTIDINE
FLAGYL I.V. RTU IN PLASTIC CONTAINER, METRONIDAZOLE
FLUCONAZOLE IN SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER, FLUCONAZOLE
HEPARIN SODIUM 1,000 UNITS AND SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER, HEPARIN SODIUM
HEPARIN SODIUM 2,000 UNITS AND SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER, HEPARIN SODIUM
HEPARIN SODIUM 20,000 UNITS AND DEXTROSE 5% IN PLASTIC CONTAINER, HEPARIN SODIUM
HEPARIN SODIUM 25,000 UNITS AND DEXTROSE 5% IN PLASTIC CONTAINER, HEPARIN SODIUM
HEPATASOL 8%, AMINO ACIDS
ISOTONIC GENTAMICIN SULFATE IN PLASTIC CONTAINER, GENTAMICIN SULFATE
LACTATED RINGER'S AND DEXTROSE 5% IN PLASTIC CONTAINER, CALCIUM CHLORIDE
LACTATED RINGER'S IN PLASTIC CONTAINER, CALCIUM CHLORIDE
LIDOCAINE HYDROCHLORIDE 0.2% AND DEXTROSE 5% IN PLASTIC CONTAINER, LIDOCAINE HYDROCHLORIDE
LIDOCAINE HYDROCHLORIDE 0.4% AND DEXTROSE 5% IN PLASTIC CONTAINER, LIDOCAINE HYDROCHLORIDE
LIDOCAINE HYDROCHLORIDE 0.8% AND DEXTROSE 5% IN PLASTIC CONTAINER, LIDOCAINE HYDROCHLORIDE
MESNEX, MESNA

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

** B **

* BAXTER HEALTHCARE CORP
 MILRINONE LACTATE IN DEXTROSE 5% IN PLASTIC CONTAINER, MILRINONE LACTATE
 NALLPEN IN PLASTIC CONTAINER, NAFCILLIN SODIUM
 NITROGLYCERIN IN DEXTROSE 5%, NITROGLYCERIN
 ONDANSETRON HYDROCHLORIDE AND SODIUM CHLORIDE IN PLASTIC CONTAINER, ONDANSETRON
 HYDROCHLORIDE
 OSMITROL 10% IN WATER IN PLASTIC CONTAINER, MANNITOL
 OSMITROL 10% IN WATER, MANNITOL
 OSMITROL 15% IN WATER IN PLASTIC CONTAINER, MANNITOL
 OSMITROL 15% IN WATER, MANNITOL
 OSMITROL 20% IN WATER IN PLASTIC CONTAINER, MANNITOL
 OSMITROL 20% IN WATER, MANNITOL
 OSMITROL 5% IN WATER IN PLASTIC CONTAINER, MANNITOL
 OSMITROL 5% IN WATER, MANNITOL
 PENICILLIN G POTASSIUM IN PLASTIC CONTAINER, PENICILLIN G POTASSIUM
 PLASMA-LYTE 148 AND DEXTROSE 5% IN PLASTIC CONTAINER, DEXTROSE
 PLASMA-LYTE 148 IN WATER IN PLASTIC CONTAINER, MAGNESIUM CHLORIDE
 PLASMA-LYTE 56 AND DEXTROSE 5% IN PLASTIC CONTAINER, DEXTROSE
 PLASMA-LYTE 56 IN PLASTIC CONTAINER, MAGNESIUM ACETATE TETRAHYDRATE
 PLASMA-LYTE A IN PLASTIC CONTAINER, MAGNESIUM CHLORIDE
 PLASMA-LYTE M AND DEXTROSE 5% IN PLASTIC CONTAINER, CALCIUM CHLORIDE
 PLASMA-LYTE R IN PLASTIC CONTAINER, CALCIUM CHLORIDE
 POTASSIUM CHLORIDE 10MEQ IN DEXTROSE 5% AND LACTATED RINGER'S IN PLASTIC CONTAINER,
 CALCIUM CHLORIDE
 POTASSIUM CHLORIDE 10MEQ IN DEXTROSE 5% AND SODIUM CHLORIDE 0.45% IN PLASTIC CONTAINER,
 DEXTROSE
 POTASSIUM CHLORIDE 10MEQ IN DEXTROSE 5% AND SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER,
 DEXTROSE
 POTASSIUM CHLORIDE 10MEQ IN PLASTIC CONTAINER, POTASSIUM CHLORIDE
 POTASSIUM CHLORIDE 15MEQ IN DEXTROSE 5% AND LACTATED RINGER'S IN PLASTIC CONTAINER,
 CALCIUM CHLORIDE
 POTASSIUM CHLORIDE 20MEQ IN DEXTROSE 5% AND LACTATED RINGER'S IN PLASTIC CONTAINER,
 CALCIUM CHLORIDE
 POTASSIUM CHLORIDE 20MEQ IN DEXTROSE 5% AND SODIUM CHLORIDE 0.45% IN PLASTIC CONTAINER,
 DEXTROSE
 POTASSIUM CHLORIDE 20MEQ IN DEXTROSE 5% AND SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER,
 DEXTROSE
 POTASSIUM CHLORIDE 20MEQ IN PLASTIC CONTAINER, POTASSIUM CHLORIDE
 POTASSIUM CHLORIDE 30MEQ IN DEXTROSE 5% AND LACTATED RINGER'S IN PLASTIC CONTAINER,
 CALCIUM CHLORIDE
 POTASSIUM CHLORIDE 30MEQ IN DEXTROSE 5% AND SODIUM CHLORIDE 0.45% IN PLASTIC CONTAINER,
 DEXTROSE
 POTASSIUM CHLORIDE 30MEQ IN DEXTROSE 5% AND SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER,
 DEXTROSE
 POTASSIUM CHLORIDE 30MEQ IN PLASTIC CONTAINER, POTASSIUM CHLORIDE
 POTASSIUM CHLORIDE 40MEQ IN DEXTROSE 5% AND LACTATED RINGER'S IN PLASTIC CONTAINER,
 CALCIUM CHLORIDE
 POTASSIUM CHLORIDE 40MEQ IN DEXTROSE 5% AND SODIUM CHLORIDE 0.45% IN PLASTIC CONTAINER,
 DEXTROSE
 POTASSIUM CHLORIDE 40MEQ IN DEXTROSE 5% AND SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER,
 DEXTROSE
 POTASSIUM CHLORIDE 40MEQ IN PLASTIC CONTAINER, POTASSIUM CHLORIDE
 POTASSIUM CHLORIDE 5MEQ IN DEXTROSE 5% AND LACTATED RINGER'S IN PLASTIC CONTAINER,
 CALCIUM CHLORIDE
 POTASSIUM CHLORIDE 5MEQ IN DEXTROSE 5% AND SODIUM CHLORIDE 0.45% IN PLASTIC CONTAINER,
 DEXTROSE
 POTASSIUM CHLORIDE 5MEQ IN DEXTROSE 5% AND SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER,
 DEXTROSE
 POTASSIUM CHLORIDE, POTASSIUM CHLORIDE
 PREMASOL 10% IN PLASTIC CONTAINER, AMINO ACIDS
 PREMASOL 6% IN PLASTIC CONTAINER, AMINO ACIDS
 RENAMIN W/O ELECTROLYTES, AMINO ACIDS
 RINGER'S IN PLASTIC CONTAINER, CALCIUM CHLORIDE
 SEVOFLURANE, SEVOFLURANE
 SODIUM CHLORIDE 0.45% AND POTASSIUM CHLORIDE 0.15% IN PLASTIC CONTAINER, POTASSIUM
 CHLORIDE
 SODIUM CHLORIDE 0.45% IN PLASTIC CONTAINER, SODIUM CHLORIDE

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

** B **

* BAXTER HEALTHCARE CORP
 SODIUM CHLORIDE 0.9% AND POTASSIUM CHLORIDE 0.15% IN PLASTIC CONTAINER, POTASSIUM CHLORIDE
 SODIUM CHLORIDE 0.9% AND POTASSIUM CHLORIDE 0.224%, POTASSIUM CHLORIDE
 SODIUM CHLORIDE 0.9% AND POTASSIUM CHLORIDE 0.3% IN PLASTIC CONTAINER, POTASSIUM CHLORIDE
 SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER, SODIUM CHLORIDE
 SODIUM CHLORIDE 0.9% IN STERILE PLASTIC CONTAINER, SODIUM CHLORIDE
 SODIUM CHLORIDE 3% IN PLASTIC CONTAINER, SODIUM CHLORIDE
 SODIUM CHLORIDE 5% IN PLASTIC CONTAINER, SODIUM CHLORIDE
 SODIUM LACTATE 0.167 MOLAR IN PLASTIC CONTAINER, SODIUM LACTATE
 SORBITOL 3% IN PLASTIC CONTAINER, SORBITOL
 STERILE WATER FOR INJECTION IN PLASTIC CONTAINER, WATER FOR INJECTION, STERILE
 STERILE WATER IN PLASTIC CONTAINER, WATER FOR IRRIGATION, STERILE
 STERILE WATER, WATER FOR IRRIGATION, STERILE
 THEOPHYLLINE AND DEXTROSE 5% IN PLASTIC CONTAINER, THEOPHYLLINE
 TIS-U-SOL IN PLASTIC CONTAINER, MAGNESIUM SULFATE
 TIS-U-SOL, MAGNESIUM SULFATE
 TRAVASOL 10% IN PLASTIC CONTAINER, AMINO ACIDS
 TRAVASOL 10% W/O ELECTROLYTES, AMINO ACIDS
 TRAVASOL 3.5% SULFITE FREE W/ ELECTROLYTES IN PLASTIC CONTAINER, AMINO ACIDS
 TRAVASOL 3.5% W/ ELECTROLYTES, AMINO ACIDS
 TRAVASOL 5.5% IN PLASTIC CONTAINER, AMINO ACIDS
 TRAVASOL 5.5% SULFITE FREE W/ ELECTROLYTES IN PLASTIC CONTAINER, AMINO ACIDS
 TRAVASOL 5.5% W/ ELECTROLYTES, AMINO ACIDS
 TRAVASOL 5.5% W/O ELECTROLYTES, AMINO ACIDS
 TRAVASOL 8.5% IN PLASTIC CONTAINER, AMINO ACIDS
 TRAVASOL 8.5% SULFITE FREE W/ ELECTROLYTES IN PLASTIC CONTAINER, AMINO ACIDS
 TRAVASOL 8.5% W/ ELECTROLYTES, AMINO ACIDS
 TRAVASOL 8.5% W/O ELECTROLYTES, AMINO ACIDS
 VANCOCIN HYDROCHLORIDE IN PLASTIC CONTAINER, VANCOMYCIN HYDROCHLORIDE

* BAXTER HEALTHCARE CORP ANESTHESIA AND CRITICAL CARE
 ACYCLOVIR IN SODIUM CHLORIDE 0.9% PRESERVATIVE FREE, ACYCLOVIR SODIUM
 ACYCLOVIR SODIUM, ACYCLOVIR SODIUM
 AMPICILLIN AND SULBACTAM, AMPICILLIN SODIUM
 ATRACURIUM BESYLATE, ATRACURIUM BESYLATE
 CEFOXITIN, CEFOXITIN SODIUM
 CHLORPROMAZINE HYDROCHLORIDE, CHLORPROMAZINE HYDROCHLORIDE
 CLINDAMYCIN PHOSPHATE, CLINDAMYCIN PHOSPHATE
 DEXAMETHASONE, DEXAMETHASONE SODIUM PHOSPHATE
 DIAZEPAM, DIAZEPAM
 DIGOXIN, DIGOXIN
 DIPHENHYDRAMINE HYDROCHLORIDE, DIPHENHYDRAMINE HYDROCHLORIDE
 DIPYRIDAMOLE, DIPYRIDAMOLE
 DOPRAM, DOXAPRAM HYDROCHLORIDE
 DURAMORPH PF, MORPHINE SULFATE
 ERYTHROMYCIN LACTOBIONATE, ERYTHROMYCIN LACTOBIONATE
 FAMOTIDINE PRESERVATIVE FREE, FAMOTIDINE
 FAMOTIDINE, FAMOTIDINE
 FENTANYL CITRATE PRESERVATIVE FREE, FENTANYL CITRATE
 FLUMAZENIL, FLUMAZENIL
 FUROSEMIDE, FUROSEMIDE
 HEPARIN SODIUM, HEPARIN SODIUM
 INFUMORPH, MORPHINE SULFATE
 KETOROLAC TROMETHAMINE, KETOROLAC TROMETHAMINE
 MIDAZOLAM HYDROCHLORIDE, MIDAZOLAM HYDROCHLORIDE
 MILRINONE LACTATE IN DEXTROSE 5% IN PLASTIC CONTAINER, MILRINONE LACTATE
 MILRINONE LACTATE, MILRINONE LACTATE
 MILRINONE LACTOSE IN 5% DEXTROSE, MILRINONE LACTATE
 MITOMYCIN, MITOMYCIN
 ONDANSETRON HYDROCHLORIDE PRESERVATIVE FREE, ONDANSETRON HYDROCHLORIDE
 ONDANSETRON HYDROCHLORIDE, ONDANSETRON HYDROCHLORIDE
 PROCHLORPERAZINE EDISYLATE, PROCHLORPERAZINE EDISYLATE
 PROCHLORPERAZINE, PROCHLORPERAZINE EDISYLATE
 PROMETHAZINE HYDROCHLORIDE, PROMETHAZINE HYDROCHLORIDE

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

** B **

- * BAXTER HEALTHCARE CORP ANESTHESIA AND CRITICAL CARE
ROBINUL, GLYCOPYRROLATE
SUFENTANIL CITRATE, SUFENTANIL CITRATE
SULFAMETHOXAZOLE AND TRIMETHOPRIM, SULFAMETHOXAZOLE
THIAMINE HYDROCHLORIDE, THIAMINE HYDROCHLORIDE
TOBRAMYCIN SULFATE, TOBRAMYCIN SULFATE
VECURONIUM BROMIDE, VECURONIUM BROMIDE
VINORELBINE TARTRATE, VINORELBINE TARTRATE
- * BAXTER HEALTHCARE INTERNATIONAL SPECIALTY THERAPIES DIV
FAMOTIDINE PRESERVATIVE FREE IN PLASTIC CONTAINER, FAMOTIDINE
PROSOL 20% SULFITE FREE IN PLASTIC CONTAINER, AMINO ACIDS

BAXTER HLTHCARE CORP

- * BAXTER HEALTHCARE CORP ANESTHESIA CRITICAL CARE
AMRINONE LACTATE, INAMRINONE LACTATE
ATIVAN, LORAZEPAM
ATracurium BESYLATE PRESERVATIVE FREE, ATracurium BESYLATE
ATracurium BESYLATE, ATracurium BESYLATE
BREVIBLOC DOUBLE STRENGTH IN PLASTIC CONTAINER, ESMOLOL HYDROCHLORIDE
BREVIBLOC IN PLASTIC CONTAINER, ESMOLOL HYDROCHLORIDE
BREVIBLOC, ESMOLOL HYDROCHLORIDE
ENLON, EDROPHONIUM CHLORIDE
ENLON-PLUS, ATROPINE SULFATE
ETHRANE, ENFLURANE
FORANE, ISOFLURANE
KETOROLAC TROMETHAMINE, KETOROLAC TROMETHAMINE
MEPERIDINE HYDROCHLORIDE, MEPERIDINE HYDROCHLORIDE
MIDAZOLAM HYDROCHLORIDE, MIDAZOLAM HYDROCHLORIDE
MILRINONE LACTATE, MILRINONE LACTATE
PROPRANOLOL HYDROCHLORIDE, PROPRANOLOL HYDROCHLORIDE
PROTOPAM CHLORIDE, PRALIDOXIME CHLORIDE
REGLAN, METOCLOPRAMIDE HYDROCHLORIDE
REVEX, NALMEFENE HYDROCHLORIDE
ROBAXIN, METHOCARBAMOL
SUPRANE, DESFLURANE

BAYER

- * BAYER HEALTHCARE LLC
ALEVE COLD AND SINUS, NAPROXEN SODIUM (OTC)
ALEVE, NAPROXEN SODIUM (OTC)
BAYER EXTRA STRENGTH ASPIRIN FOR MIGRAINE PAIN, ASPIRIN (OTC)
FEMSTAT 3, BUTOCONAZOLE NITRATE (OTC)

BAYER PHARMS

- * BAYER PHARMACEUTICALS CORP
ADALAT CC, NIFEDIPINE
AVELOX IN SODIUM CHLORIDE 0.8% IN PLASTIC CONTAINER, MOXIFLOXACIN HYDROCHLORIDE
AVELOX, MOXIFLOXACIN HYDROCHLORIDE
BILTRICIDE, PRAZIQUANTEL
CIPRO IN DEXTROSE 5% IN PLASTIC CONTAINER, CIPROFLOXACIN
CIPRO XR, CIPROFLOXACIN
CIPRO, CIPROFLOXACIN
CIPRO, CIPROFLOXACIN HYDROCHLORIDE
DTIC-DOME, DACARBAZINE
LEVITRA, VARDENAFIL HYDROCHLORIDE
MYCELEX, CLOTRIMAZOLE
MYCELEX-7 COMBINATION PACK, CLOTRIMAZOLE (OTC)
MYCELEX-7, CLOTRIMAZOLE (OTC)
NEXAVAR, SORAFENIB TOSYLATE
NIMOTOP, NIMODIPINE
PRECOSE, ACARBOSE
TRASYLOL, APROTININ BOVINE

BECTON DICKINSON

- * BECTON DICKINSON AND CO
CHLORHEXIDINE GLUCONATE, CHLORHEXIDINE GLUCONATE (OTC)

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

**** B ****

* BECTON DICKINSON AND CO
E-Z SCRUB 201, POVIDONE-IODINE (OTC)
E-Z SCRUB 241, POVIDONE-IODINE (OTC)

BEDFORD

* BEDFORD LABORATORIES DIV BEN VENUE LABORATORIES INC
ACETAZOLAMIDE SODIUM, ACETAZOLAMIDE SODIUM
ACETYLCYSTEINE, ACETYLCYSTEINE
ACYCLOVIR SODIUM, ACYCLOVIR SODIUM
ADENOSINE, ADENOSINE
ALPROSTADIL, ALPROSTADIL
AMIKACIN SULFATE, AMIKACIN SULFATE
AMIODARONE HYDROCHLORIDE, AMIODARONE HYDROCHLORIDE
AMRINONE, INAMRINONE LACTATE
ATRACURIUM BESYLATE PRESERVATIVE FREE, ATRACURIUM BESYLATE
ATRACURIUM BESYLATE, ATRACURIUM BESYLATE
AZATHIOPRINE SODIUM, AZATHIOPRINE SODIUM
BLEOMYCIN, BLEOMYCIN SULFATE
BUMETANIDE, BUMETANIDE
BUPRENORPHINE HYDROCHLORIDE, BUPRENORPHINE HYDROCHLORIDE
BUTORPHANOL TARTRATE PRESERVATIVE FREE, BUTORPHANOL TARTRATE
BUTORPHANOL TARTRATE, BUTORPHANOL TARTRATE
CARBOPLATIN, CARBOPLATIN
CERUBIDINE, DAUNORUBICIN HYDROCHLORIDE
CHLOROPROCAINE HYDROCHLORIDE, CHLOROPROCAINE HYDROCHLORIDE
CISPLATIN, CISPLATIN
CLADRIBINE, CLADRIBINE
CLINDAMYCIN PHOSPHATE, CLINDAMYCIN PHOSPHATE
CYCLOSPORINE, CYCLOSPORINE
CYTARABINE, CYTARABINE
DACARBAZINE, DACARBAZINE
DAUNORUBICIN HYDROCHLORIDE, DAUNORUBICIN HYDROCHLORIDE
DEXRAZOXANE, DEXRAZOXANE HYDROCHLORIDE
DICYCLOMINE HYDROCHLORIDE (PRESERVATIVE-FREE), DICYCLOMINE HYDROCHLORIDE
DILTIAZEM HYDROCHLORIDE, DILTIAZEM HYDROCHLORIDE
DIPYRIDAMOLE, DIPYRIDAMOLE
DOBUTAMINE HYDROCHLORIDE, DOBUTAMINE HYDROCHLORIDE
DOXAPRAM HYDROCHLORIDE, DOXAPRAM HYDROCHLORIDE
DOXORUBICIN HYDROCHLORIDE, DOXORUBICIN HYDROCHLORIDE
DOXYCYCLINE, DOXYCYCLINE HYCLATE
ENALAPRILAT, ENALAPRILAT
ETOMIDATE, ETOMIDATE
ETOPOSIDE, ETOPOSIDE
FAMOTIDINE PRESERVATIVE FREE, FAMOTIDINE
FAMOTIDINE, FAMOTIDINE
FLOXURIDINE, FLOXURIDINE
FLUCONAZOLE IN SODIUM CHLORIDE 0.9%, FLUCONAZOLE
FLUPHENAZINE DECANOATE, FLUPHENAZINE DECANOATE
GANCICLOVIR SODIUM, GANCICLOVIR SODIUM
HALOPERIDOL DECANOATE, HALOPERIDOL DECANOATE
HALOPERIDOL, HALOPERIDOL LACTATE
KETAMINE HYDROCHLORIDE, KETAMINE HYDROCHLORIDE
KETOROLAC TROMETHAMINE, KETOROLAC TROMETHAMINE
LABETALOL HYDROCHLORIDE, LABETALOL HYDROCHLORIDE
LEUCOVORIN CALCIUM PRESERVATIVE FREE, LEUCOVORIN CALCIUM
LEUCOVORIN CALCIUM, LEUCOVORIN CALCIUM
LEVOCARNITINE, LEVOCARNITINE
LORAZEPAM, LORAZEPAM
MESNA, MESNA
METHOTREXATE PRESERVATIVE FREE, METHOTREXATE SODIUM
METHOTREXATE SODIUM, METHOTREXATE SODIUM
MIDAZOLAM HYDROCHLORIDE, MIDAZOLAM HYDROCHLORIDE
MILRINONE LACTATE, MILRINONE LACTATE
MITOMYCIN, MITOMYCIN

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

** B **

* BEDFORD LABORATORIES DIV BEN VENUE LABORATORIES INC
MITOXANTRONE, MITOXANTRONE HYDROCHLORIDE
NOREPINEPHRINE BITARTRATE, NOREPINEPHRINE BITARTRATE
OCTREOTIDE ACETATE (PRESERVATIVE FREE), OCTREOTIDE ACETATE
OCTREOTIDE ACETATE, OCTREOTIDE ACETATE
OFLOXACIN, OFLOXACIN
ONDANSETRON HYDROCHLORIDE, ONDANSETRON HYDROCHLORIDE
PACLITAXEL, PACLITAXEL
PAMIDRONATE DISODIUM, PAMIDRONATE DISODIUM
PHENTOLAMINE MESYLATE, PHENTOLAMINE MESYLATE
POLYMYXIN B SULFATE, POLYMYXIN B SULFATE
PROCHLORPERAZINE EDISYLATE, PROCHLORPERAZINE EDISYLATE
PROPOFOL, PROPOFOL
PROPRANOLOL HYDROCHLORIDE, PROPRANOLOL HYDROCHLORIDE
RANITIDINE HYDROCHLORIDE, RANITIDINE HYDROCHLORIDE
RANITIDINE, RANITIDINE HYDROCHLORIDE
RIFAMPIN, RIFAMPIN
TERBUTALINE SULFATE, TERBUTALINE SULFATE
THIOTEPA, THIOTEPA
VALPROATE SODIUM, VALPROATE SODIUM
VECURONIUM BROMIDE, VECURONIUM BROMIDE
VERAPAMIL HYDROCHLORIDE, VERAPAMIL HYDROCHLORIDE
VINBLASTINE SULFATE, VINBLASTINE SULFATE
VINORELBINE TARTRATE, VINORELBINE TARTRATE

BEDFORD LABS

* BEDFORD LABORATORIES
ALLOPURINOL SODIUM, ALLOPURINOL SODIUM
AMIODARONE HYDROCHLORIDE, AMIODARONE HYDROCHLORIDE
CARBOPLATIN, CARBOPLATIN
CIPROFLOXACIN, CIPROFLOXACIN
DIHYDROERGOTAMINE MESYLATE, DIHYDROERGOTAMINE MESYLATE
ESMOLOL HYDROCHLORIDE, ESMOLOL HYDROCHLORIDE
FENOLDOPAM MESYLATE, FENOLDOPAM MESYLATE
FLUMAZENIL, FLUMAZENIL
IDARUBICIN HYDROCHLORIDE, IDARUBICIN HYDROCHLORIDE
LORAZEPAM PRESERVATIVE FREE, LORAZEPAM
METOPROLOL TARTRATE, METOPROLOL TARTRATE
ONDANSETRON HYDROCHLORIDE PRESERVATIVE FREE, ONDANSETRON HYDROCHLORIDE
ORPHENADRINE CITRATE, ORPHENADRINE CITRATE
PROMETHAZINE HYDROCHLORIDE, PROMETHAZINE HYDROCHLORIDE
* BEDFORD LABORATORIES INC
LEUPROLIDE ACETATE, LEUPROLIDE ACETATE

BELCHER

* BELCHER PHARMACEUTICALS INC
CEPHALEXIN, CEPHALEXIN

BEN VENUE

* BEN VENUE LABORATORIES INC
AMIODARONE HYDROCHLORIDE, AMIODARONE HYDROCHLORIDE
FAMOTIDINE PRESERVATIVE FREE, FAMOTIDINE
MIDAZOLAM HYDROCHLORIDE, MIDAZOLAM HYDROCHLORIDE
RANITIDINE HYDROCHLORIDE, RANITIDINE HYDROCHLORIDE

BERLEX

* BERLEX INC
ANGELIQ, DROSPIRENONE
BETAPACE AF, SOTALOL HYDROCHLORIDE
BETAPACE, SOTALOL HYDROCHLORIDE
CLIMARA PRO, ESTRADIOL
CLIMARA, ESTRADIOL
FLUDARA, FLUDARABINE PHOSPHATE
LEVITE, ETHINYL ESTRADIOL
MAGNEVIST, GADOPENTETATE DIMEGLUMINE
MENOSTAR, ESTRADIOL

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

**** B ****

* BERLEX INC
MIRENA, LEVONORGESTREL
REFLUDAN, LEPYRIDIN RECOMBINANT
ULTRAVIST (PHARMACY BULK), IOPROMIDE
ULTRAVIST 150, IOPROMIDE
ULTRAVIST 240, IOPROMIDE
ULTRAVIST 300, IOPROMIDE
ULTRAVIST 370, IOPROMIDE
YASMIN, DROSPIRENONE
YAZ, DROSPIRENONE

BIO NUCLEONICS

* BIO NUCLEONICS INC
STRONTIUM CHLORIDE SR-89, STRONTIUM CHLORIDE, SR-89

BIOCHEMIE

* BIOCHEMIE GMBH
AMOXICILLIN, AMOXICILLIN
PENICILLIN G SODIUM, PENICILLIN G SODIUM
PENICILLIN V POTASSIUM, PENICILLIN V POTASSIUM

BIOGLAN PHARMS CORP

* BIOGLAN PHARMS CORP DBA DOAK DERMATOLOGICS
SOLARAZE, DICLOFENAC SODIUM

BIOKEY

* BIOKEY INC
BENZAEPRIIL HYDROCHLORIDE, BENAZEPRIL HYDROCHLORIDE

BIONICHE (CANADA)

* BIONICHE PHARMA (CANADA) LTD
AMIODARONE HYDROCHLORIDE, AMIODARONE HYDROCHLORIDE
DIMETHYL SULFOXIDE, DIMETHYL SULFOXIDE
EDETATE DISODIUM, EDETATE DISODIUM
KETAMINE HYDROCHLORIDE, KETAMINE HYDROCHLORIDE
MILRINONE LACTATE, MILRINONE LACTATE

BIONICHE PHARM USA

* BIONICHE PHARMA USA INC
SOTRADECOL, SODIUM TETRADECYL SULFATE

BIONICHE PHARMA

* BIONICHE PHARMA
CYANOCOBALAMIN, CYANOCOBALAMIN
DIPHENHYDRAMINE HYDROCHLORIDE, DIPHENHYDRAMINE HYDROCHLORIDE
PROMETHAZINE HYDROCHLORIDE, PROMETHAZINE HYDROCHLORIDE
RIMS0-50, DIMETHYL SULFOXIDE

BIOSANTE

* BIOSANTE PHARMACEUTICALS INC
ELESTRIN, ESTRADIOL

BIOVAIL

* BIOVAIL CORP INTERNATIONAL
TIAZAC, DILTIAZEM HYDROCHLORIDE
* BIOVAIL LABORATORIES INC
ATIVAN, LORAZEPAM
CARDIZEM CD, DILTIAZEM HYDROCHLORIDE
CARDIZEM, DILTIAZEM HYDROCHLORIDE
DICLOFENAC SODIUM, DICLOFENAC SODIUM
DILTIAZEM HYDROCHLORIDE, DILTIAZEM HYDROCHLORIDE
ISORDIL, ISOSORBIDE DINITRATE
NIFEDIPINE, NIFEDIPINE
PENTOXIFYLLINE, PENTOXIFYLLINE
TRAMADOL HYDROCHLORIDE, TRAMADOL HYDROCHLORIDE

BIOVAIL LABS INTL

* BIOVAIL LABORATORIES INTERNATIONAL SRL
CARDIZEM LA, DILTIAZEM HYDROCHLORIDE
CARDIZEM, DILTIAZEM HYDROCHLORIDE

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

** B **

* BIOVAIL LABORATORIES INTERNATIONAL SRL
TRAMADOL HYDROCHLORIDE, TRAMADOL HYDROCHLORIDE
VASERETIC, ENALAPRIL MALEATE
VASOTEC, ENALAPRIL MALEATE
VASOTEC, ENALAPRILAT

BLAIREX

* BLAIREX LABORATORIES INC
BRONCHO SALINE, SODIUM CHLORIDE (OTC)

BOCA PHARMA

* BOCA PHARMACAL INC
HYDROCORTISONE ACETATE 1% AND PRAMOXINE HYDROCHLORIDE 1%, HYDROCORTISONE ACETATE
METHSCOPOLAMINE BROMIDE, METHSCOPOLAMINE BROMIDE

BOEHRINGER INGELHEIM

* BOEHRINGER INGELHEIM
CATAPRES, CLONIDINE HYDROCHLORIDE
CATAPRES-TTS-1, CLONIDINE
CATAPRES-TTS-2, CLONIDINE
CATAPRES-TTS-3, CLONIDINE
MEXITIL, MEXILETINE HYDROCHLORIDE
MICARDIS HCT, HYDROCHLOROTHIAZIDE
MICARDIS, TELMISARTAN
MIRAPEX, PRAMIPEXOLE DIHYDROCHLORIDE
* BOEHRINGER INGELHEIM PHARMACEUTICALS INC
AGGRENOX, ASPIRIN
ALUPENT, METAPROTERENOL SULFATE
APTIVUS, TIPRANAVIR
ATROVENT HFA, IPRATROPIUM BROMIDE
ATROVENT, IPRATROPIUM BROMIDE
COMBIVENT, ALBUTEROL SULFATE
FLOMAX, TAMSULOSIN HYDROCHLORIDE
MOBIC, MELOXICAM
PERSANTINE, DIPYRIDAMOLE
SPIRIVA, TIOTROPIUM BROMIDE MONOHYDRATE
VIRAMUNE, NEVIRAPINE

BRACCO

* BRACCO DIAGNOSTICS INC
CARDIOGEN-82, RUBIDIUM CHLORIDE RB-82
CHOLETEC, TECHNETIUM TC-99M MEBROFENIN KIT
CHOLOGRAFIN MEGLUMINE, IODIPAMIDE MEGLUMINE
CHROMITOPES SODIUM, SODIUM CHROMATE, CR-51
CYSTOGRAFIN DILUTE, DIATRIZOATE MEGLUMINE
CYSTOGRAFIN, DIATRIZOATE MEGLUMINE
GASTROGRAFIN, DIATRIZOATE MEGLUMINE
IODOTOPE, SODIUM IODIDE, I-131
ISOVUE-200, IOPAMIDOL
ISOVUE-250, IOPAMIDOL
ISOVUE-300, IOPAMIDOL
ISOVUE-370, IOPAMIDOL
ISOVUE-M 200, IOPAMIDOL
ISOVUE-M 300, IOPAMIDOL
KINEVAC, SINICALIDE
MACROTEC, TECHNETIUM TC-99M ALBUMIN AGGREGATED KIT
MDP-BRACCO, TECHNETIUM TC-99M MEDRONATE KIT
MULTIHANCE MULTIPACK, GADOBENATE DIMEGLUMINE
MULTIHANCE, GADOBENATE DIMEGLUMINE
PHOSPHOTEC, TECHNETIUM TC-99M PYROPHOSPHATE KIT
PROHANCE MULTIPACK, GADOTERIDOL
PROHANCE, GADOTERIDOL
RENO-30, DIATRIZOATE MEGLUMINE
RENO-60, DIATRIZOATE MEGLUMINE
RENOCAL-76, DIATRIZOATE MEGLUMINE
RENO-DIP, DIATRIZOATE MEGLUMINE

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

**** B ****

* BRACCO DIAGNOSTICS INC
RENOGRAFIN-60, DIATRIZOATE MEGLUMINE
RENOGRAFIN-76, DIATRIZOATE MEGLUMINE
RUBRATOPE-57, CYANOCOBALAMIN, CO-57
SINOGRAFIN, DIATRIZOATE MEGLUMINE

BRADLEY PHARMS

* BRADLEY PHARMACEUTICALS INC
PAMINE FORTE, METHSCOPOLAMINE BROMIDE
PAMINE, METHSCOPOLAMINE BROMIDE
ZONALON, DOXEPIN HYDROCHLORIDE

BRAINTREE

* BRAINTREE LABORATORIES INC
AXID, NIZATIDINE
GOLYTELY, POLYETHYLENE GLYCOL 3350
HALFLYTELY, BISACODYL
NULYTELY, POLYETHYLENE GLYCOL 3350
NULYTELY-FLAVORED, POLYETHYLENE GLYCOL 3350

BRECKENRIDGE PHARM

* BRECKENRIDGE PHARMACEUTICAL INC
MELOXICAM, MELOXICAM

BRIGHTON PHARMS INC

* BRIGHTON PHARMACEUTICALS INC
BROMFED-DM, BROMPHENIRAMINE MALEATE

BRIGHTSTONE

* BRIGHTSTONE PHARMA INC
ISOSORBIDE MONONITRATE, ISOSORBIDE MONONITRATE

BRISTOL

* BRISTOL LABORATORIES INC DIV BRISTOL MYERS CO
BICNU, CARMUSTINE
BRISTACYCLINE, TETRACYCLINE HYDROCHLORIDE
LYSODREN, MITOTANE
TETREX, TETRACYCLINE PHOSPHATE COMPLEX

BRISTOL MYERS

* BRISTOL MYERS CO
MUTAMYCIN, MITOMYCIN
PLATINOL, CISPLATIN
PLATINOL-AQ, CISPLATIN
QUESTRAN LIGHT, CHOLESTYRAMINE
QUESTRAN, CHOLESTYRAMINE

BRISTOL MYERS SQUIBB

* BRISTOL MYERS SQUIBB
AZACTAM, AZTREONAM
BARACLUDE, ENTECAVIR
DEXTROSE 5% AND POTASSIUM CHLORIDE 0.075% IN PLASTIC CONTAINER, DEXTROSE
DEXTROSE 5% AND POTASSIUM CHLORIDE 0.15% IN PLASTIC CONTAINER, DEXTROSE
DEXTROSE 5% AND POTASSIUM CHLORIDE 0.224% IN PLASTIC CONTAINER, DEXTROSE
DEXTROSE 5% AND POTASSIUM CHLORIDE 0.3% IN PLASTIC CONTAINER, DEXTROSE
GLUCOVANCE, GLYBURIDE
MEGACE, MEGESTROL ACETATE
MONOPRIL-HCT, FOSINOPRIL SODIUM
PRAVACHOL, PRAVASTATIN SODIUM
* BRISTOL MYERS SQUIBB CO
AZACTAM IN PLASTIC CONTAINER, AZTREONAM
CEENU, LOMUSTINE
DROXIA, HYDROXYUREA
ESTRACE, ESTRADIOL
GLUCOPHAGE XR, METFORMIN HYDROCHLORIDE
HYDREA, HYDROXYUREA
METAGLIP, GLIPIZIDE
PARAPLATIN, CARBOPLATIN
PRAVIGARD PAC (COPACKAGED), ASPIRIN

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

**** B ****

- * BRISTOL MYERS SQUIBB CO
PROLIXIN DECANOATE, FLUPHENAZINE DECANOATE
REYATAZ, ATAZANAVIR SULFATE
RUBEX, DOXORUBICIN HYDROCHLORIDE
SPRYCEL, DASATINIB
SUSTIVA, EFAVIRENZ
TESLAC, TESTOLACTONE
VEPESID, ETOPOSIDE
VIDEX EC, DIDANOSINE
- * BRISTOL MYERS SQUIBB CO PHARMACEUTICAL RESEARCH INSTITUTE
BLENOXANE, BLEOMYCIN SULFATE
BUSPAR, BUSPIRONE HYDROCHLORIDE
CEFZIL, CEFPROZIL
CYTOXAN, CYCLOPHOSPHAMIDE
ESTRACE, ESTRADIOL
ETOPOPHOS PRESERVATIVE FREE, ETOPOSIDE PHOSPHATE
GLUCOPHAGE, METFORMIN HYDROCHLORIDE
IFEX/MESNEX KIT, IFOSFAMIDE
LYOPHILIZED CYTOXAN, CYCLOPHOSPHAMIDE
MAXIPIME, CEFEPIME HYDROCHLORIDE (ARGININE FORMULATION)
MONOPRIL, FOSINOPRIL SODIUM
PARAPLATIN, CARBOPLATIN
TAXOL, PACLITAXEL
VIDEX, DIDANOSINE
VUMON, TENIPOSIDE
ZERIT, STAVUDINE
- * BRISTOL MYERS SQUIBB MEDICAL IMAGING
DEFINITY, PERFLUTREN
GALLIUM CITRATE GA 67, GALLIUM CITRATE, GA-67
NEUROLITE, TECHNETIUM TC-99M BICISATE KIT
TECHNELITE, TECHNETIUM TC-99M SODIUM PERTECHNETATE GENERATOR
THALLOUS CHLORIDE TL 201, THALLOUS CHLORIDE, TL-201
XENON XE 133, XENON, XE-133
- * BRISTOL MYERS SQUIBB PHARMA CO
CARDIOLITE, TECHNETIUM TC-99M SESTAMIBI KIT
COUMADIN, WARFARIN SODIUM
LODOSYN, CARBIDOPA
MIRALUMA, TECHNETIUM TC-99M SESTAMIBI KIT
SINEMET CR, CARBIDOPA
SINEMET, CARBIDOPA
SUSTIVA, EFAVIRENZ

BRYAN

- * BRYAN CORP
SCLEROSOL, TALC
TALC, TALC

BUNDY

- * CM BUNDY CO
BUTABARBITAL, BUTABARBITAL SODIUM
MECLIZINE HYDROCHLORIDE, MECLIZINE HYDROCHLORIDE

C AND M PHARMA

- * C AND M PHARMACAL INC
HI-COR, HYDROCORTISONE

CANYON

- * CANYON PHARMACEUTICALS INC
IPRIVASK, DESIRUDIN RECOMBINANT

CARACO

- * CARACO PHARMACEUTICAL LABORATORIES LTD
BACLOFEN, BACLOFEN
CARBAMAZEPINE, CARBAMAZEPINE
CITALOPRAM HYDROBROMIDE, CITALOPRAM HYDROBROMIDE
CLONAZEPAM, CLONAZEPAM
CLOZAPINE, CLOZAPINE

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

**** C ****

* CARACO PHARMACEUTICAL LABORATORIES LTD
DIGOXIN, DIGOXIN
FLURBIPROFEN, FLURBIPROFEN
FLUVOXAMINE MALEATE, FLUVOXAMINE MALEATE
GLIPIZIDE, GLIPIZIDE
MELOXICAM, MELOXICAM
MEPERIDINE HYDROCHLORIDE, MEPERIDINE HYDROCHLORIDE
METFORMIN HYDROCHLORIDE, METFORMIN HYDROCHLORIDE
METOPROLOL TARTRATE, METOPROLOL TARTRATE
MIRTAZAPINE, MIRTAZAPINE
OXAPROZIN, OXAPROZIN
PAROMOMYCIN SULFATE, PAROMOMYCIN SULFATE
TICLOPIDINE HYDROCHLORIDE, TICLOPIDINE HYDROCHLORIDE
TIZANIDINE HYDROCHLORIDE, TIZANIDINE HYDROCHLORIDE
TRAMADOL HYDROCHLORIDE AND ACETAMINOPHEN, ACETAMINOPHEN
TRAMADOL HYDROCHLORIDE, TRAMADOL HYDROCHLORIDE

CARLSBAD

* CARLSBAD TECHNOLOGY INC
ACYCLOVIR, ACYCLOVIR
CEFACLOL, CEFACLOL
CIPROFLOXACIN, CIPROFLOXACIN HYDROCHLORIDE
DICLOFENAC SODIUM, DICLOFENAC SODIUM
FAMOTIDINE, FAMOTIDINE
FLUOXETINE, FLUOXETINE HYDROCHLORIDE
LOVASTATIN, LOVASTATIN
MELOXICAM, MELOXICAM

CAROLINA MEDCL

* CAROLINA MEDICAL PRODUCTS CO
AMANTADINE HYDROCHLORIDE, AMANTADINE HYDROCHLORIDE
HYDROCORTISONE IN ABSORBASE, HYDROCORTISONE
ISONIAZID, ISONIAZID
SODIUM POLYSTYRENE SULFONATE, SODIUM POLYSTYRENE SULFONATE
SPS, SODIUM POLYSTYRENE SULFONATE
TRIAMCINOLONE ACETONIDE IN ABSORBASE, TRIAMCINOLONE ACETONIDE

CEDAR PHARMS

* CEDAR PHARMACEUTICALS LLC
METHIMAZOLE, METHIMAZOLE

CELGENE

* CELGENE CORP
REVLIMID, LENALIDOMIDE
THALOMID, THALIDOMIDE

CEPH INTL

* CEPH INTERNATIONAL CORP
CEFACLOL, CEFACLOL
CEPHALEXIN, CEPHALEXIN

CEPHALON

* CEPHALON INC
ACTIQ, FENTANYL CITRATE
FENTORA, FENTANYL CITRATE
GABITRIL, TIAGABINE HYDROCHLORIDE
PROVIGIL, MODAFINIL
TRISENOX, ARSENIC TRIOXIDE

CHATTEM

* CHATTEM INC
SELSUN, SELENIUM SULFIDE

CHESEBROUGH PONDS

* CHESEBROUGH PONDS INC
EXTRA-STRENGTH AIM, SODIUM MONOFLUOROPHOSPHATE (OTC)

CHIRHOCLIN

* CHIRHOCLIN INC
CHIRHOSTIM, SECRETIN SYNTHETIC HUMAN

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

**** C ****

* CHIRHOCLIN INC
SECREFLO, SECRETIN SYNTHETIC PORCINE

CIPHER

* CIPHER PHARMACEUTICALS LTD
LIPOFEN, FENOFIBRATE

CIS

* CIS US INC
AN-DTPA, TECHNETIUM TC-99M PENTETATE KIT
AN-SULFUR COLLOID, TECHNETIUM TC-99M SULFUR COLLOID KIT
CIS-MDP, TECHNETIUM TC-99M MEDRONATE KIT
CIS-PYRO, TECHNETIUM TC-99M PYROPHOSPHATE KIT
HEPATOLITE, TECHNETIUM TC-99M DISOFENIN KIT
IOBENGUANE SULFATE I 131, IOBENGUANE SULFATE I 131
PULMOLITE, TECHNETIUM TC-99M ALBUMIN AGGREGATED KIT

CLINTEC NUTR

* CLINTEC NUTRITION CO SUB CLINIQUE
CLINISOL 15% SULFITE FREE IN PLASTIC CONTAINER, AMINO ACIDS

CLONMEL

* CLONMEL HEALTHCARE
AMOXICILLIN, AMOXICILLIN
AMPICILLIN TRIHYDRATE, AMPICILLIN/AMPICILLIN TRIHYDRATE
PENICILLIN V POTASSIUM, PENICILLIN V POTASSIUM

CLONMEL HLTHCARE

* CLONMEL HEALTHCARE LTD
ACYCLOVIR, ACYCLOVIR
ATENOLOL, ATENOLOL
CIMETIDINE HYDROCHLORIDE, CIMETIDINE HYDROCHLORIDE
CIMETIDINE, CIMETIDINE
CLONIDINE HYDROCHLORIDE, CLONIDINE HYDROCHLORIDE
DIAZEPAM, DIAZEPAM
DILTIAZEM HYDROCHLORIDE, DILTIAZEM HYDROCHLORIDE
DOXAZOSIN MESYLATE, DOXAZOSIN MESYLATE
GLYBURIDE (MICRONIZED), GLYBURIDE
HYDROCHLOROTHIAZIDE, HYDROCHLOROTHIAZIDE
NAPROXEN, NAPROXEN
PYRAZINAMIDE, PYRAZINAMIDE
SELEGILINE HYDROCHLORIDE, SELEGILINE HYDROCHLORIDE

COASTAL PHARMS

* COASTAL PHARMACEUTICALS INC
POLYETHYLENE GLYCOL 3350, POLYETHYLENE GLYCOL 3350

COBALT

* COBALT PHARMACEUTICALS INC
CIPROFLOXACIN, CIPROFLOXACIN HYDROCHLORIDE
CITALOPRAM HYDROBROMIDE, CITALOPRAM HYDROBROMIDE
FOSINOPRIL SODIUM, FOSINOPRIL SODIUM
GLIMEPIRIDE, GLIMEPIRIDE
METFORMIN HYDROCHLORIDE, METFORMIN HYDROCHLORIDE
PRAVASTATIN SODIUM, PRAVASTATIN SODIUM
RAMIPRIL, RAMIPRIL
RANITIDINE HYDROCHLORIDE, RANITIDINE HYDROCHLORIDE
SIMVASTATIN, SIMVASTATIN

COLGATE

* COLGATE ORAL PHARMACEUTICALS INC
ORABASE HCA, HYDROCORTISONE ACETATE
PERIOGARD, CHLORHEXIDINE GLUCONATE

COLGATE PALMOLIVE

* COLGATE PALMOLIVE
COLGATE TOTAL, SODIUM FLUORIDE (OTC)

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

** C **

COLLAGENEX PHARMS

- * COLLAGENEX PHARMACEUTICALS INC
ORACEA, DOXYCYCLINE
PERIOSTAT, DOXYCYCLINE HYCLATE

COLUMBIA LABS

- * COLUMBIA LABORATORIES INC
CRINONE, PROGESTERONE
STRIANT, TESTOSTERONE

CONNETICS

- * CONNETICS CORP
EVOCLIN, CLINDAMYCIN PHOSPHATE
LUXIQ, BETAMETHASONE VALERATE
OLUX, CLOBETASOL PROPIONATE
SORIATANE, ACITRETIN
VERDESO, DESONIDE

CONTROLLED THERAP

- * CONTROLLED THERAPEUTICS (SCOTLAND) LTD
CERVIDIL, DINOPROSTONE

COOK IMAGING

- * COOK IMAGING CORP
IOPAMIDOL-200, IOPAMIDOL
IOPAMIDOL-250, IOPAMIDOL
IOPAMIDOL-300, IOPAMIDOL
IOPAMIDOL-370, IOPAMIDOL

COPLEY PHARM

- * COPLEY PHARMACEUTICAL INC
CO-LAV, POLYETHYLENE GLYCOL 3350
DOXYLAMINE SUCCINATE, DOXYLAMINE SUCCINATE (OTC)
GO-EVAC, POLYETHYLENE GLYCOL 3350
MINOXIDIL (FOR MEN), MINOXIDIL (OTC)
POTASSIUM CHLORIDE, POTASSIUM CHLORIDE

COREPHARMA

- * COREPHARMA LLC
ACETAMINOPHEN, ACETAMINOPHEN (OTC)
BENZTROPINE MESYLATE, BENZTROPINE MESYLATE
CARISOPRODOL, CARISOPRODOL
CILOSTAZOL, CILOSTAZOL
CITALOPRAM HYDROBROMIDE, CITALOPRAM HYDROBROMIDE
CLINDAMYCIN HYDROCHLORIDE, CLINDAMYCIN HYDROCHLORIDE
CYPROHEPTADINE HYDROCHLORIDE, CYPROHEPTADINE HYDROCHLORIDE
DEXTROAMP SACCHARATE, AMP ASPARTATE, DEXTROAMP SULFATE AND AMP SULFATE, AMPHETAMINE
ASPARTATE
DOXYCYCLINE HYCLATE, DOXYCYCLINE HYCLATE
GLIMEPIRIDE, GLIMEPIRIDE
GLIPIZIDE AND METFORMIN HYDROCHLORIDE, GLIPIZIDE
GLYBURIDE AND METFORMIN HYDROCHLORIDE, GLYBURIDE
GLYBURIDE, GLYBURIDE
GLYCOPYRROLATE, GLYCOPYRROLATE
LEVOCARNITINE, LEVOCARNITINE
MELOXICAM, MELOXICAM
METHENAMINE HIPPURATE, METHENAMINE HIPPURATE
PILOCARPINE HYDROCHLORIDE, PILOCARPINE HYDROCHLORIDE
POTASSIUM CITRATE, POTASSIUM CITRATE
PYRIDOSTIGMINE BROMIDE, PYRIDOSTIGMINE BROMIDE
RIMANTADINE HYDROCHLORIDE, RIMANTADINE HYDROCHLORIDE
TIZANIDINE HYDROCHLORIDE, TIZANIDINE HYDROCHLORIDE
TRAMADOL HYDROCHLORIDE, TRAMADOL HYDROCHLORIDE
URSODIOL, URSODIOL

CORNERSTONE

- * CORNERSTONE BIOPHARMA INC
PROPOXYPHENE NAPSYLATE AND ACETAMINOPHEN, ACETAMINOPHEN

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

**** C ****

* CORNERSTONE BIOPHARMA INC
SPECTRACEF, CEFEDITOREN PIVOXIL

COTHERIX

* COTHERIX INC
VENTAVIS, ILOPROST

CP PHARMS

* CP PHARMACEUTICALS CV
LYRICA, PREGABALIN

CPPI CV

* CP PHARMACEUTICALS INTERNATIONAL CV
SUTENT, SUNITINIB MALATE

CRITICAL

* CRITICAL THERAPEUTICS INC
ZYFLO, ZILEUTON

CUBIST

* CUBIST PHARMACEUTICALS INC
CUBICIN, DAPTOMYCIN

CUMBERLAND PHARMS

* CUMBERLAND PHARMACEUTICALS INC
ACETADOTE, ACETYLCYSTEINE

CV THERAP

* CV THERAPEUTICS INC
RANEXA, RANOLAZINE

CYTOGEN

* CYTOGEN CORP
QUADRAMET, SAMARIUM SM 153 LEXIDRONAM PENTASODIUM

DABUR ONCOLOGY PLC

* DABUR ONCOLOGY PLC
CARBOPLATIN, CARBOPLATIN
PACLITAXEL, PACLITAXEL

DAIICHI

* DAIICHI MEDICAL RESEARCH INC
EVOXAC, CEVIMELINE HYDROCHLORIDE
* DAIICHI PHARMACEUTICAL CORP
FLOXIN OTIC, OFLOXACIN

DAIICHI SANKYO

* DAIICHI SANKYO INC
BENICAR HCT, HYDROCHLOROTHIAZIDE
BENICAR, OLMESARTAN MEDOXOMIL
WELCHOL, COLESEVELAM HYDROCHLORIDE

DAINIPPON

* DAINIPPON PHARMACEUTICAL USA CORP
ZONEGRAN, ZONISAMIDE

DANCO LABS LLC

* DANCO LABORATORIES LLC
MIFEPREX, MIFEPRISTONE

DAVA PHARMS INC

* DAVA PHARMACEUTICALS INC
VOSPIRE ER, ALBUTEROL SULFATE

DAVIS AND GECK

* DAVIS AND GECK DIV AMERICAN CYANAMID CO
PRE-OP II, HEXACHLOROPHENE
PRE-OP, HEXACHLOROPHENE

DEL RAY LABS

* DEL RAY LABORATORIES INC
ALA-CORT, HYDROCORTISONE
ALA-SCALP, HYDROCORTISONE
TRIDERM, TRIAMCINOLONE ACETONIDE

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

** D **

DELL LABS

- * DELL LABORATORIES INC
LIDOCAINE HYDROCHLORIDE W/ EPINEPHRINE, EPINEPHRINE

DENTSPLY PHARM

- * DENTSPLY PHARMACEUTICAL
CITANEST FORTE, EPINEPHRINE BITARTRATE
CITANEST PLAIN, PRILOCAINE HYDROCHLORIDE
ORAQIX, LIDOCAINE
POLOCAINE W/ LEVONORDEFIN, LEVONORDEFIN
POLOCAINE, MEPIVACAINE HYDROCHLORIDE
XYLOCAINE W/ EPINEPHRINE, EPINEPHRINE
XYLOCAINE, LIDOCAINE HYDROCHLORIDE

DEPOMED INC

- * DEPOMED INC
GLUMETZA, METFORMIN HYDROCHLORIDE

DEPROCO

- * DEPROCO INC
LIGNOSPAN FORTE, EPINEPHRINE BITARTRATE
LIGNOSPAN STANDARD, EPINEPHRINE BITARTRATE
SCANDONEST L, LEVONORDEFIN
SCANDONEST PLAIN, MEPIVACAINE HYDROCHLORIDE
SEPTOCAINE, ARTICAINE HYDROCHLORIDE

DEXCEL LTD

- * DEXCEL LTD
DICLOFENAC SODIUM, DICLOFENAC SODIUM
ISOSORBIDE MONONITRATE, ISOSORBIDE MONONITRATE

DEXCEL PHARMA

- * DEXCEL PHARMA TECHNOLOGIES LTD
PERIOCHIP, CHLORHEXIDINE GLUCONATE

DEY

- * DEY LP
ACCUNEB, ALBUTEROL SULFATE
ALBUTEROL SULFATE, ALBUTEROL SULFATE
CROMOLYN SODIUM, CROMOLYN SODIUM
CUROSURF, PORACTANT ALFA
DUONEB, ALBUTEROL SULFATE
IPRATROPIUM BROMIDE, IPRATROPIUM BROMIDE
METAPROTERENOL SULFATE, METAPROTERENOL SULFATE

DHL

- * DHL LABORATORIES INC
DEXTROSE 5% IN PLASTIC CONTAINER, DEXTROSE

DIALYSIS SUPS

- * DIALYSIS SUPPLIES INC
NORMOCARB HF 25, MAGNESIUM CHLORIDE
NORMOCARB HF 35, MAGNESIUM CHLORIDE

DORC

- * DORC INTERNATIONAL BV
VISIONBLUE, TRYPAN BLUE

DPT

- * DPT LABORATORIES INC
EMBELINE, CLOBETASOL PROPIONATE

DR REDDYS LABS INC

- * DR REDDYS LABORATORIES INC
FINASTERIDE, FINASTERIDE
FLUCONAZOLE, FLUCONAZOLE
FLUOXETINE HYDROCHLORIDE, FLUOXETINE HYDROCHLORIDE
FLUOXETINE, FLUOXETINE HYDROCHLORIDE
IBUPROFEN, IBUPROFEN
IBUPROFEN, IBUPROFEN (OTC)

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

**** D ****

* DR REDDYS LABORATORIES INC
ISMO, ISOSORBIDE MONONITRATE
MELOXICAM, MELOXICAM
METFORMIN HYDROCHLORIDE, METFORMIN HYDROCHLORIDE
NAPROXEN SODIUM AND PSEUDOEPHEDRINE HYDROCHLORIDE, NAPROXEN SODIUM (OTC)
NAPROXEN SODIUM, NAPROXEN SODIUM (OTC)
NEFAZODONE HYDROCHLORIDE, NEFAZODONE HYDROCHLORIDE
PRAVASTATIN SODIUM, PRAVASTATIN SODIUM
RANITIDINE HYDROCHLORIDE, RANITIDINE HYDROCHLORIDE
SECTRAL, ACEBUTOLOL HYDROCHLORIDE
SIMVASTATIN, SIMVASTATIN
TENEX, GUANFACINE HYDROCHLORIDE
TIZANIDINE HYDROCHLORIDE, TIZANIDINE HYDROCHLORIDE

DR REDDYS LABS LTD

* DR REDDYS LABORATORIES LTD
CIPROFLOXACIN, CIPROFLOXACIN HYDROCHLORIDE
CITALOPRAM HYDROBROMIDE, CITALOPRAM HYDROBROMIDE
ENALAPRIL MALEATE AND HYDROCHLOROTHIAZIDE, ENALAPRIL MALEATE
FAMOTIDINE, FAMOTIDINE
FAMOTIDINE, FAMOTIDINE (OTC)
FEXOFENADINE HYDROCHLORIDE, FEXOFENADINE HYDROCHLORIDE
GLIMEPIRIDE, GLIMEPIRIDE
IBUPROFEN AND PSEUDOEPHEDRINE HYDROCHLORIDE, IBUPROFEN (OTC)
NIZATIDINE, NIZATIDINE
OFLOXACIN, OFLOXACIN
ONDANSETRON HYDROCHLORIDE, ONDANSETRON HYDROCHLORIDE
OXAPROZIN, OXAPROZIN
RANITIDINE HYDROCHLORIDE, RANITIDINE HYDROCHLORIDE
RANITIDINE, RANITIDINE HYDROCHLORIDE (OTC)
ZONISAMIDE, ZONISAMIDE

DRAXIMAGE

* DRAXIMAGE INC
DRAXIMAGE MDP, TECHNETIUM TC-99M MEDRONATE KIT
DTPA, TECHNETIUM TC-99M PENTETATE KIT
SODIUM IODIDE I 131, SODIUM IODIDE, I-131
SODIUM IODIDE I 131, KIT, SODIUM IODIDE, I-131
TECHNISCAN GLUCEPTATE, TECHNETIUM TC-99M GLUCEPTATE KIT
TECHNETIUM TC 99M ALBUMIN AGGREGATED KIT, TECHNETIUM TC-99M ALBUMIN AGGREGATED KIT

DURAMED

* DURAMED PHARMACEUTICALS INC
CENESTIN, ESTROGENS, CONJUGATED SYNTHETIC A
ENJUVIA, ESTROGENS, CONJUGATED SYNTHETIC B
MIRCETTE, DESOGESTREL
NORDETTE-28, ETHINYL ESTRADIOL
PLAN B, LEVONORGESTREL
PLAN B, LEVONORGESTREL (OTC)
PREFEST, ESTRADIOL
REVIA, NALTREXONE HYDROCHLORIDE
SEASONALE, ETHINYL ESTRADIOL

DURAMED PHARMS BARR

* DURAMED PHARMACEUTICALS INC SUB BARR LABORATORIES INC
ACETAMINOPHEN AND CODEINE PHOSPHATE, ACETAMINOPHEN
AVIANE-28, ETHINYL ESTRADIOL
AYGESTIN, NORETHINDRONE ACETATE
CIMETIDINE HYDROCHLORIDE, CIMETIDINE HYDROCHLORIDE
CRYSSELLE, ETHINYL ESTRADIOL
DESOGESTREL AND ETHINYL ESTRADIOL, DESOGESTREL
DIAMOX, ACETAZOLAMIDE
DIAMOX, ACETAZOLAMIDE SODIUM
ENPRESSE-28, ETHINYL ESTRADIOL
ESTROPIPATE, ESTROPIPATE
GYNODIOL, ESTRADIOL

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

** D **

* DURAMED PHARMACEUTICALS INC SUB BARR LABORATORIES INC
HYDROXYUREA, HYDROXYUREA
ISONIAZID, ISONIAZID
LOPERAMIDE HYDROCHLORIDE, LOPERAMIDE HYDROCHLORIDE (OTC)
MEDROXYPROGESTERONE ACETATE, MEDROXYPROGESTERONE ACETATE
MEPERIDINE HYDROCHLORIDE, MEPERIDINE HYDROCHLORIDE
METHOTREXATE SODIUM, METHOTREXATE SODIUM
METHYLPREDNISOLONE, METHYLPREDNISOLONE
OXYCODONE AND ACETAMINOPHEN, ACETAMINOPHEN
PARAGARD T 380A, COPPER
PROCHLORPERAZINE MALEATE, PROCHLORPERAZINE MALEATE
TRIAMTERENE AND HYDROCHLOROTHIAZIDE, HYDROCHLOROTHIAZIDE
VELIVET, DESOGESTREL
ZEBETA, BISOPROLOL FUMARATE
ZIAC, BISOPROLOL FUMARATE

DURAMED RES

* DURAMED RESEARCH INC
SEASONIQUE, ETHINYL ESTRADIOL

DUSA

* DUSA PHARMACEUTICALS INC
LEVULAN, AMINOLEVULINIC ACID HYDROCHLORIDE

EASTMAN KODAK

* EASTMAN KODAK CO
CARBOCAINE W/ NEO-COBEFRIN, LEVONORDEFIN
CARBOCAINE, MEPIVACAINE HYDROCHLORIDE
LIDOCAINE HYDROCHLORIDE AND EPINEPHRINE, EPINEPHRINE

ECOLAB

* ECOLAB INC
CHG SCRUB, CHLORHEXIDINE GLUCONATE (OTC)
CIDA-STAT, CHLORHEXIDINE GLUCONATE (OTC)

EGIS

* EGIS PHARMACEUTICALS
BUSPIRONE HYDROCHLORIDE, BUSPIRONE HYDROCHLORIDE
PIROXICAM, PIROXICAM

EGIS PHARMS

* EGIS PHARMACEUTICALS LTD
CAPTOPRIL, CAPTOPRIL

EISAI MEDCL RES

* EISAI MEDICAL RESEARCH INC
ACIPHEX, RABEPRAZOLE SODIUM
ARICEPT ODT, DONEPEZIL HYDROCHLORIDE
ARICEPT, DONEPEZIL HYDROCHLORIDE
PANRETIN, ALITRETINOIN
TARGETIN, BEXAROTENE

ELAN DRUG

* ELAN DRUG DELIVERY INC
VERELAN PM, VERAPAMIL HYDROCHLORIDE
VERELAN, VERAPAMIL HYDROCHLORIDE

ELAN PHARM

* ELAN PHARMACEUTICAL RESEARCH CORP
ISOSORBIDE MONONITRATE, ISOSORBIDE MONONITRATE
KETOPROFEN, KETOPROFEN

ELAN PHARMS

* ELAN PHARMACEUTICALS INC
PRIALT, ZICONOTIDE

ELKINS SINN

* ELKINS SINN DIV AH ROBINS CO INC
MEPERIDINE HYDROCHLORIDE, MEPERIDINE HYDROCHLORIDE
PANCURONIUM, PANCURONIUM BROMIDE
PHENYTOIN, PHENYTOIN SODIUM

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

** E **

EMD PHARMS

* EMD PHARMACEUTICALS INC
CYANOKIT, HYDROXOCOBALAMIN

EMPI

* EMPI INC
LIDOCAINE HYDROCHLORIDE AND EPINEPHRINE, EPINEPHRINE

ENCYSIVE

* ENCYSIVE PHARMACEUTICALS INC
ARGATROBAN, ARGATROBAN

ENDO PHARMS

* ENDO PHARMACEUTICALS INC
ASPIRIN AND OXYCODONE, ASPIRIN
FROVA, FROVATRIPTAN SUCCINATE
HYCODAN, HOMATROPINE METHYLBROMIDE
MOBAN, MOLINDONE HYDROCHLORIDE
MORPHINE SULFATE, MORPHINE SULFATE
NARCAN, NALOXONE HYDROCHLORIDE
NUBAIN, NALBUPHINE HYDROCHLORIDE
NUMORPHAN, OXYMORPHONE HYDROCHLORIDE
OPANA ER, OXYMORPHONE HYDROCHLORIDE
OPANA, OXYMORPHONE HYDROCHLORIDE
OXYCODONE AND ACETAMINOPHEN, ACETAMINOPHEN
OXYCODONE HYDROCHLORIDE, OXYCODONE HYDROCHLORIDE
PERCOCET, ACETAMINOPHEN
PERCODAN, ASPIRIN
SELEGILINE HYDROCHLORIDE, SELEGILINE HYDROCHLORIDE
SYMMETREL, AMANTADINE HYDROCHLORIDE
SYNERA, LIDOCAINE
ZYDONE, ACETAMINOPHEN

ENZON

* ENZON INC
ABELCET, AMPHOTERICIN B
ADAGEN, PEGADEMASE BOVINE

ESPRIT PHARMA

* ESPRIT PHARMA INC
ESTRASORB, ESTRADIOL HEMIHYDRATE
PROQUIN XR, CIPROFLOXACIN HYDROCHLORIDE

EURAND

* EURAND AMERICA INC
POTASSIUM CHLORIDE, POTASSIUM CHLORIDE

EVERYLIFE

* EVERYLIFE
HYDROCORTISONE, HYDROCORTISONE
SECOBARBITAL SODIUM, SECOBARBITAL SODIUM

EXCELLIUM

* EXCELLIUM PHARMACEUTICAL INC
FUROSEMIDE, FUROSEMIDE

FALCON PHARMS

* FALCON PHARMACEUTICALS INC
TIMOLOL MALEATE, TIMOLOL MALEATE
* FALCON PHARMACEUTICALS LTD
DIPIVEFRIN HYDROCHLORIDE, DIPIVEFRIN HYDROCHLORIDE
GENTAMICIN SULFATE, GENTAMICIN SULFATE
LEVOBUNOLOL HYDROCHLORIDE, LEVOBUNOLOL HYDROCHLORIDE
MAXITROL, DEXAMETHASONE
METIPRANOLOL, METIPRANOLOL HYDROCHLORIDE
TIMOLOL MALEATE, TIMOLOL MALEATE
TOBREX, TOBRAMYCIN
TRIMETHOPRIM SULFATE AND POLYMYXIN B SULFATE, POLYMYXIN B SULFATE

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

** F **

FARMACON IL

- * FARMACON IL LLC
NEOPROFEN, IBUPROFEN LYSINE

FERNDALE LABS

- * FERNDALE LABORATORIES INC
HYDROCORTISONE ACETATE, HYDROCORTISONE ACETATE
LOCOID LIPOCREAM, HYDROCORTISONE BUTYRATE
LOCOID, HYDROCORTISONE BUTYRATE
MICORT-HC, HYDROCORTISONE ACETATE
PRAMOSONE, HYDROCORTISONE ACETATE
STRIFON FORTE DSC, CHLORZOXAZONE

FERRING

- * FERRING PHARMACEUTICALS INC
ACTHREL, CORTICORELIN OVINE TRIFLUTATE
BRAVELLE, UROFOLLITROPIN
MENOPUR, LUTEINIZING HORMONE
MINIRIN, DESMOPRESSIN ACETATE
REPRONEX, LUTEINIZING HORMONE
TEV-TROPIN, SOMATROPIN RECOMBINANT

FLEMING

- * FLEMING AND CO PHARMACEUTICALS
THYROSHIELD, POTASSIUM IODIDE (OTC)

FLEMINGTON PHARM

- * FLEMINGTON PHARMACEUTICAL CORP
NIFEDIPINE, NIFEDIPINE

FOREST LABS

- * FOREST LABORATORIES INC
AEROSPAN HFA, FLUNISOLIDE
CAMPRAL, ACAMPROSATE CALCIUM
CELEXA, CITALOPRAM HYDROBROMIDE
COMBUNOX, IBUPROFEN
ELIXOPHYLLIN, THEOPHYLLINE
FLUMADINE, RIMANTADINE HYDROCHLORIDE
LEXAPRO, ESCITALOPRAM OXALATE
NAMENDA, MEMANTINE HYDROCHLORIDE
TESSALON, BENZONATATE
THYROLAR-0.25, LIOETHYRONINE SODIUM
THYROLAR-0.5, LIOETHYRONINE SODIUM
THYROLAR-1, LIOETHYRONINE SODIUM
THYROLAR-2, LIOETHYRONINE SODIUM
THYROLAR-3, LIOETHYRONINE SODIUM

FOUGERA

- * E FOUGERA DIV ALTANA INC
BETAMETHASONE DIPROPIONATE, BETAMETHASONE DIPROPIONATE
BETAMETHASONE VALERATE, BETAMETHASONE VALERATE
CLINDAMYCIN PHOSPHATE, CLINDAMYCIN PHOSPHATE
CLOBETASOL PROPIONATE, CLOBETASOL PROPIONATE
ERYTHROMYCIN, ERYTHROMYCIN
FLUOCINOLONE ACETONIDE, FLUOCINOLONE ACETONIDE
FLUOCINONIDE, FLUOCINONIDE
GENTAMICIN SULFATE, GENTAMICIN SULFATE
HYDROCORTISONE, HYDROCORTISONE
LIDOCAINE, LIDOCAINE
NEOMYCIN AND POLYMYXIN B SULFATES AND DEXAMETHASONE, DEXAMETHASONE
NITROGLYCERIN, NITROGLYCERIN
NYSTATIN, NYSTATIN
NYSTATIN-TRIAMCINOLONE ACETONIDE, NYSTATIN

FRESENIUS

- * FRESENIUS KABI DEUTSCHLAND GMBH
INTRALIPID 10%, SOYBEAN OIL
INTRALIPID 20%, SOYBEAN OIL

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

**** F ****

- * FRESENIUS KABI DEUTSCHLAND GMBH
INTRALIPID 30%, SOYBEAN OIL
- * FRESENIUS USA INC
INPERSOL-LC/LM W/ DEXTROSE 1.5% IN PLASTIC CONTAINER, CALCIUM CHLORIDE
INPERSOL-LC/LM W/ DEXTROSE 2.5% IN PLASTIC CONTAINER, CALCIUM CHLORIDE
INPERSOL-LC/LM W/ DEXTROSE 3.5% IN PLASTIC CONTAINER, CALCIUM CHLORIDE
INPERSOL-LC/LM W/ DEXTROSE 4.25% IN PLASTIC CONTAINER, CALCIUM CHLORIDE

FRESENIUS MEDCL

- * FRESENIUS MEDICAL CARE NORTH AMERICA
CALCITRIOL, CALCITRIOL
DELFLEX W/ DEXTROSE 1.5% IN PLASTIC CONTAINER, CALCIUM CHLORIDE
DELFLEX W/ DEXTROSE 1.5% LOW MAGNESIUM IN PLASTIC CONTAINER, CALCIUM CHLORIDE
DELFLEX W/ DEXTROSE 1.5% LOW MAGNESIUM LOW CALCIUM IN PLASTIC CONTAINER, CALCIUM CHLORIDE
DELFLEX W/ DEXTROSE 2.5% IN PLASTIC CONTAINER, CALCIUM CHLORIDE
DELFLEX W/ DEXTROSE 2.5% LOW MAGNESIUM IN PLASTIC CONTAINER, CALCIUM CHLORIDE
DELFLEX W/ DEXTROSE 2.5% LOW MAGNESIUM LOW CALCIUM IN PLASTIC CONTAINER, CALCIUM CHLORIDE
DELFLEX W/ DEXTROSE 3.5% IN PLASTIC CONTAINER, CALCIUM CHLORIDE
DELFLEX W/ DEXTROSE 4.25% IN PLASTIC CONTAINER, CALCIUM CHLORIDE
DELFLEX W/ DEXTROSE 4.25% LOW MAGNESIUM IN PLASTIC CONTAINER, CALCIUM CHLORIDE
DELFLEX W/ DEXTROSE 4.25% LOW MAGNESIUM LOW CALCIUM IN PLASTIC CONTAINER, CALCIUM CHLORIDE
DELFLEX-LM W/ DEXTROSE 1.5% IN PLASTIC CONTAINER, CALCIUM CHLORIDE
DELFLEX-LM W/ DEXTROSE 2.5% IN PLASTIC CONTAINER, CALCIUM CHLORIDE
DELFLEX-LM W/ DEXTROSE 3.5% IN PLASTIC CONTAINER, CALCIUM CHLORIDE
DELFLEX-LM W/ DEXTROSE 4.25% IN PLASTIC CONTAINER, CALCIUM CHLORIDE
PHOSLO GELCAPS, CALCIUM ACETATE

FSC

- * FSC LABORATORIES INC
ISOPTIN SR, VERAPAMIL HYDROCHLORIDE
ISOPTIN, VERAPAMIL HYDROCHLORIDE
PRIMSOL, TRIMETHOPRIM HYDROCHLORIDE

G AND W LABS

- * G AND W LABORATORIES INC
ACEPHEN, ACETAMINOPHEN (OTC)
FLUOCINOLONE ACETONIDE, FLUOCINOLONE ACETONIDE
FLUTICASONE PROPIONATE, FLUTICASONE PROPIONATE
HALOBETASOL PROPIONATE, HALOBETASOL PROPIONATE
INDOMETHACIN, INDOMETHACIN
MICONAZOLE 7 COMBINATION PACK, MICONAZOLE NITRATE (OTC)
MICONAZOLE NITRATE, MICONAZOLE NITRATE (OTC)
MIGERGOT, CAFFEINE
MOMETASONE FUROATE, MOMETASONE FUROATE
PROCHLORPERAZINE, PROCHLORPERAZINE
PROMETHAZINE HYDROCHLORIDE, PROMETHAZINE HYDROCHLORIDE
PROMETHEGAN, PROMETHAZINE HYDROCHLORIDE
TRIAMCINOLONE ACETONIDE, TRIAMCINOLONE ACETONIDE
TRUPHYLLINE, AMINOPHYLLINE

GALDERMA LABS

- * GALDERMA LABORATORIES INC
CLOBEX, CLOBETASOL PROPIONATE

GALDERMA LABS LP

- * GALDERMA LABORATORIES L P
CLOBEX, CLOBETASOL PROPIONATE
- * GALDERMA LABORATORIES LP
CLINDAGEL, CLINDAMYCIN PHOSPHATE
CLOBEX, CLOBETASOL PROPIONATE
DESOWEN, DESONIDE
DIFFERIN, ADAPALENE
FS SHAMPOO, FLUOCINOLONE ACETONIDE
METROCREAM, METRONIDAZOLE
METROGEL, METRONIDAZOLE
METROLOTION, METRONIDAZOLE

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

**** G ****

* GALDERMA LABORATORIES LP
TRI-LUMA, FLUOCINOLONE ACETONIDE

GALEN LTD

* GALEN LTD
FEMRING, ESTRADIOL ACETATE

GAMBRO RENAL PRODS

* GAMBRO RENAL PRODUCTS
PRISMASOL BGK 0/2.5 IN PLASTIC CONTAINER, CALCIUM CHLORIDE
PRISMASOL BGK 2/0 IN PLASTIC CONTAINER, CALCIUM CHLORIDE
PRISMASOL BGK 2/3.5 IN PLASTIC CONTAINER, CALCIUM CHLORIDE
PRISMASOL BGK 4/0 IN PLASTIC CONTAINER, CALCIUM CHLORIDE
PRISMASOL BGK 4/2.5 IN PLASTIC CONTAINER, CALCIUM CHLORIDE
PRISMASOL BK 0/0 IN PLASTIC CONTAINER, CALCIUM CHLORIDE
PRISMASOL BK 0/3.5 IN PLASTIC CONTAINER, CALCIUM CHLORIDE

GD SEARLE

* GD SEARLE LLC
BEXTRA, VALDECOXIB
CELEBREX, CELECOXIB
DAYPRO, OXAPROZIN

GD SEARLE LLC

* GD SEARLE LLC
ALDACTAZIDE, HYDROCHLOROTHIAZIDE
ALDACTONE, SPIRONOLACTONE
ARTHROTEC, DICLOFENAC SODIUM
CALAN, VERAPAMIL HYDROCHLORIDE
COVERA-HS, VERAPAMIL HYDROCHLORIDE
CYTOTEC, MISOPROSTOL
DEMULEN 1/35-21, ETHINYL ESTRADIOL
DEMULEN 1/35-28, ETHINYL ESTRADIOL
DEMULEN 1/50-21, ETHINYL ESTRADIOL
DEMULEN 1/50-28, ETHINYL ESTRADIOL
FLAGYL ER, METRONIDAZOLE
FLAGYL I.V. RTU IN PLASTIC CONTAINER, METRONIDAZOLE
FLAGYL I.V., METRONIDAZOLE HYDROCHLORIDE
FLAGYL, METRONIDAZOLE
INSPRA, EPLERENONE
LOMOTIL, ATROPINE SULFATE
NORPACE CR, DISOPYRAMIDE PHOSPHATE
NORPACE, DISOPYRAMIDE PHOSPHATE
SYNAREL, NAFARELIN ACETATE

GE HEALTHCARE

* GE HEALTHCARE
CERETEC, TECHNETIUM TC-99M EXAMETAZIME KIT
HYPAQUE, DIATRIZOATE MEGLUMINE
HYPAQUE, DIATRIZOATE SODIUM
HYPAQUE-76, DIATRIZOATE MEGLUMINE
HYPAQUE-CYSTO, DIATRIZOATE MEGLUMINE
INDIUM IN-111 OXYQUINOLINE, INDIUM IN-111 OXYQUINOLINE
METASTRON, STRONTIUM CHLORIDE, SR-89
MPI DMSA KIDNEY REAGENT, TECHNETIUM TC-99M SUCCIMER KIT
MPI DTPA KIT - CHELATE, TECHNETIUM TC-99M PENTETATE KIT
MPI INDIUM DTPA IN 111, INDIUM IN-111 PENTETATE DISODIUM
MYOVUE, TECHNETIUM TC-99M TETROFOSMIN KIT
OMNIPAQUE 140, IOHEXOL
OMNIPAQUE 180, IOHEXOL
OMNIPAQUE 240, IOHEXOL
OMNIPAQUE 300, IOHEXOL
OMNIPAQUE 350, IOHEXOL
OMNISCAN, GADODIAMIDE
OPTISON, ALBUMIN HUMAN
SODIUM IODIDE I 123, SODIUM IODIDE, I-123
TECHNETIUM TC 99M MPI MDP, TECHNETIUM TC-99M MEDRONATE KIT

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

** G **

* GE HEALTHCARE
TECHNETIUM TC-99M PENTETATE KIT, TECHNETIUM TC-99M PENTETATE KIT
THALLOUS CHLORIDE TL 201, THALLOUS CHLORIDE, TL-201
VISIPAQUE 270, IODIXANOL
VISIPAQUE 320, IODIXANOL

GEDEON RICHTER USA

* GEDEON RICHTER USA INC
FINASTERIDE, FINASTERIDE

GENENTECH

* GENENTECH INC
NUTROPIN AQ PEN, SOMATROPIN RECOMBINANT
NUTROPIN AQ, SOMATROPIN RECOMBINANT
NUTROPIN, SOMATROPIN RECOMBINANT

GENIX THERAP

* GENIX THERAPEUTICS INC
CALCITRIOL, CALCITRIOL

GENPHARM

* GENPHARM INC
ACYCLOVIR, ACYCLOVIR
AMNESTEEM, ISOTRETINOIN
AZATHIOPRINE, AZATHIOPRINE
BENAZEPRIL HYDROCHLORIDE AND HYDROCHLOROTHIAZIDE, BENAZEPRIL HYDROCHLORIDE
BENAZEPRIL HYDROCHLORIDE, BENAZEPRIL HYDROCHLORIDE
CIPROFLOXACIN, CIPROFLOXACIN HYDROCHLORIDE
CLARITHROMYCIN, CLARITHROMYCIN
DOXAZOSIN MESYLATE, DOXAZOSIN MESYLATE
ENALAPRIL MALEATE, ENALAPRIL MALEATE
ETIDRONATE DISODIUM, ETIDRONATE DISODIUM
ETODOLAC, ETODOLAC
ETOPOSIDE, ETOPOSIDE
FAMOTIDINE, FAMOTIDINE
FAMOTIDINE, FAMOTIDINE (OTC)
FLUCONAZOLE, FLUCONAZOLE
FLUTAMIDE, FLUTAMIDE
FLUVOXAMINE MALEATE, FLUVOXAMINE MALEATE
FOSINOPRIL SODIUM AND HYDROCHLOROTHIAZIDE, FOSINOPRIL SODIUM
GLIMEPIRIDE, GLIMEPIRIDE
GUANFACINE HYDROCHLORIDE, GUANFACINE HYDROCHLORIDE
LEVOTHYROXINE SODIUM, LEVOTHYROXINE SODIUM
LORATADINE, LORATADINE (OTC)
LOVASTATIN, LOVASTATIN
MELOXICAM, MELOXICAM
METFORMIN HYDROCHLORIDE, METFORMIN HYDROCHLORIDE
METHIMAZOLE, METHIMAZOLE
NICARDIPINE HYDROCHLORIDE, NICARDIPINE HYDROCHLORIDE
NIZATIDINE, NIZATIDINE
NOVOTHYROX, LEVOTHYROXINE SODIUM
OXAPROZIN, OXAPROZIN
PRAVASTATIN SODIUM, PRAVASTATIN SODIUM
QUINAPRIL HYDROCHLORIDE, QUINAPRIL HYDROCHLORIDE
RANITIDINE HYDROCHLORIDE, RANITIDINE HYDROCHLORIDE
RANITIDINE, RANITIDINE HYDROCHLORIDE (OTC)
SOTALOL HYDROCHLORIDE, SOTALOL HYDROCHLORIDE
TICLOPIDINE HYDROCHLORIDE, TICLOPIDINE HYDROCHLORIDE
WARFARIN SODIUM, WARFARIN SODIUM

* GENPHARM PHARMACEUTICALS INC
ATENOLOL, ATENOLOL
PINDOLOL, PINDOLOL
PIROXICAM, PIROXICAM

GENTA

* GENTA INC
GANITE, GALLIUM NITRATE

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

**** G ****

GENZYME

* GENZYME CORP
CEREDASE, ALGLUCERASE
CEREZYME, IMIGLUCERASE
CLOLAR, CLOFARABINE
HECTOROL, DOXERCALCIFEROL
LEUPROLIDE ACETATE, LEUPROLIDE ACETATE
RENAGEL, SEVELAMER HYDROCHLORIDE
THYROGEN, THYROTROPIN ALFA

GILEAD

* GILEAD SCIENCES INC
ATRIPLA, EFAVIRENZ
DAUNOXOME, DAUNORUBICIN CITRATE
EMTRIVA, EMTRICITABINE
HEPSERA, ADEFOVIR DIPIVOXIL
TRUVADA, EMTRICITABINE
VIREAD, TENOFOVIR DISOPROXIL FUMARATE
VISTIDE, CIDOFOVIR

GLADES PHARMS

* GLADES PHARMACEUTICALS INC
E-GLADES, ERYTHROMYCIN

GLADES PHARMS LLC

* GLADES PHARMACEUTICALS LLC
DECLOMYCIN, DEMECLOCYCLINE HYDROCHLORIDE

GLAND PHARMA LTD

* GLAND PHARMA LTD
AMIODARONE HYDROCHLORIDE, AMIODARONE HYDROCHLORIDE
HALOPERIDOL, HALOPERIDOL LACTATE
KETOROLAC TROMETHAMINE, KETOROLAC TROMETHAMINE
MILRINONE LACTATE, MILRINONE LACTATE

GLAXO GRP LTD

* GLAXO GROUP LTD DBA GLAXOSMITHKLINE
FLOVENT HFA, FLUTICASONE PROPIONATE
SEREVENT, SALMETEROL XINAFOATE

GLAXOSMITHKLINE

* GLAXOSMITHKLINE
ABREVA, DOCOSANOL (OTC)
ACLOVATE, ALCLOMETASONE DIPROPIONATE
ADVAIR DISKUS 100/50, FLUTICASONE PROPIONATE
ADVAIR DISKUS 250/50, FLUTICASONE PROPIONATE
ADVAIR DISKUS 500/50, FLUTICASONE PROPIONATE
ADVAIR HFA, FLUTICASONE PROPIONATE
AGENERASE, AMPRENAVIR
ALBENZA, ALBENDAZOLE
ALKERAN, MELPHALAN
ALKERAN, MELPHALAN HYDROCHLORIDE
AMERGE, NARATRIPTAN HYDROCHLORIDE
AMOXIL, AMOXICILLIN
ANCEF, CEFAZOLIN SODIUM
ANSPOR, CEPHRADINE
ARIXTRA, FONDAPARINUX SODIUM
AUGMENTIN '125', AMOXICILLIN
AUGMENTIN '200', AMOXICILLIN
AUGMENTIN '250', AMOXICILLIN
AUGMENTIN '400', AMOXICILLIN
AUGMENTIN '500', AMOXICILLIN
AUGMENTIN '875', AMOXICILLIN
AUGMENTIN ES-600, AMOXICILLIN
AUGMENTIN XR, AMOXICILLIN
AVODART, DUTASTERIDE
BACTOCILL, OXACILLIN SODIUM

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

**** G ****

* GLAXOSMITHKLINE
BACTROBAN, MUPIROCI
BACTROBAN, MUPIROCI
BECONASE AQ, BECLOMETHASONE DIPROPIONATE MONOHYDRATE
CEFAZOLIN SODIUM, CEFZOLIN SODIUM
CEFTIN, CEFUROXIME AXETIL
CLOXAPEN, CLOXACILLIN SODIUM
COMBIVIR, LAMIVUDINE
COMPAZINE, PROCHLORPERAZINE
COMPAZINE, PROCHLORPERAZINE EDISYLATE
COMPAZINE, PROCHLORPERAZINE MALEATE
DARAPRIM, PYRIMETHAMINE
DEXEDRINE, DEXTROAMPHETAMINE SULFATE
DYAZIDE, HYDROCHLOROTHIAZIDE
EPIVIR, LAMIVUDINE
EPIVIR-HBV, LAMIVUDINE
ESKALITH, LITHIUM CARBONATE
FLOLAN, EPOPROSTENOL SODIUM
FLONASE, FLUTICASONE PROPIONATE
FLOVENT DISKUS 100, FLUTICASONE PROPIONATE
FLOVENT DISKUS 250, FLUTICASONE PROPIONATE
FLOVENT DISKUS 50, FLUTICASONE PROPIONATE
FLOVENT, FLUTICASONE PROPIONATE
FORTAZ IN PLASTIC CONTAINER, CEFTAZIDIME SODIUM
FORTAZ, CEFTAZIDIME
HYCAMTIN, TOPOTECAN HYDROCHLORIDE
IMITREX STATDOSE, SUMATRIPTAN SUCCINATE
IMITREX, SUMATRIPTAN
IMITREX, SUMATRIPTAN SUCCINATE
LAMICTAL CD, LAMOTRIGINE
LAMICTAL, LAMOTRIGINE
LANOXIN PEDIATRIC, DIGOXIN
LANOXIN, DIGOXIN
LAROTID, AMOXICILLIN
LEXIVA, FOSAMPRENAVIR CALCIUM
LOTRONEX, ALOSETRON HYDROCHLORIDE
MALARONE PEDIATRIC, ATOVAQUONE
MALARONE, ATOVAQUONE
MEPRON, ATOVAQUONE
MYLERAN, BUSULFAN
NICORETTE (MINT), NICOTINE POLACRILEX (OTC)
NICORETTE, NICOTINE POLACRILEX (OTC)
PARNATE, TRANLYCYPROMINE SULFATE
PAXIL CR, PAROXETINE HYDROCHLORIDE
PAXIL, PAROXETINE HYDROCHLORIDE
RELENZA, ZANAMIVIR
REQUIP, ROPINIROLE HYDROCHLORIDE
RETROVIR, ZIDOVUDINE
STELAZINE, TRIFLUOPERAZINE HYDROCHLORIDE
TAGAMET HB, CIMETIDINE (OTC)
TAGAMET, CIMETIDINE
THIOGUANINE, THIOGUANINE
THORAZINE, CHLORPROMAZINE HYDROCHLORIDE
TIMENTIN IN PLASTIC CONTAINER, CLAVULANATE POTASSIUM
TIMENTIN, CLAVULANATE POTASSIUM
TRIZIVIR, ABACAVIR SULFATE
VALTREX, VALACYCLOVIR HYDROCHLORIDE
VENTOLIN HFA, ALBUTEROL SULFATE
VENTOLIN, ALBUTEROL
WELLBUTRIN SR, BUPROPION HYDROCHLORIDE
WELLBUTRIN, BUPROPION HYDROCHLORIDE
ZANTAC 150, RANITIDINE HYDROCHLORIDE
ZANTAC 25, RANITIDINE HYDROCHLORIDE
ZANTAC 300, RANITIDINE HYDROCHLORIDE

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

** G **

- * GLAXOSMITHKLINE
 - ZANTAC IN PLASTIC CONTAINER, RANITIDINE HYDROCHLORIDE
 - ZANTAC, RANITIDINE HYDROCHLORIDE
 - ZIAGEN, ABACAVIR SULFATE
 - ZINACEF IN PLASTIC CONTAINER, CEFUROXIME SODIUM
 - ZINACEF, CEFUROXIME SODIUM
 - ZOFRAN IN PLASTIC CONTAINER, ONDANSETRON HYDROCHLORIDE
 - ZOFRAN ODT, ONDANSETRON
 - ZOFRAN PRESERVATIVE FREE, ONDANSETRON HYDROCHLORIDE
 - ZOFRAN, ONDANSETRON HYDROCHLORIDE
 - ZOVIRAX, ACYCLOVIR
 - ZOVIRAX, ACYCLOVIR SODIUM
 - ZYBAN, BUPROPION HYDROCHLORIDE

GLAXOSMITHKLINE CONS

- * GLAXOSMITHKLINE CONSUMER HEALTHCARE
 - COMMIT, NICOTINE POLACRILEX (OTC)

GLENMARK PHARMA

- * GLENMARK PHARMACEUTICALS INC USA
 - DIPYRIDAMOLE, DIPYRIDAMOLE
 - FLUCONAZOLE, FLUCONAZOLE
 - NILSTAT, NYSTATIN

GLENMARK PHARMS

- * GLENMARK PHARMACEUTICALS LTD
 - GABAPENTIN, GABAPENTIN
 - ZONISAMIDE, ZONISAMIDE

GLENMARK PHARMS LTD

- * GLENMARK PHARMACEUTICALS LTD INDIA
 - MELOXICAM, MELOXICAM

GLENWOOD

- * GLENWOOD INC
 - PASKALIUM, POTASSIUM AMINOSALICYLATE

GOLDLINE

- * GOLDLINE LABORATORIES INC
 - CLOZAPINE, CLOZAPINE
 - METFORMIN HYDROCHLORIDE, METFORMIN HYDROCHLORIDE

GRAHAM CHEM

- * GRAHAM CHEMICAL CO
 - LIDOCAINE HYDROCHLORIDE AND EPINEPHRINE, EPINEPHRINE
 - LIDOCAINE HYDROCHLORIDE, LIDOCAINE HYDROCHLORIDE
 - LIDOCAINE, LIDOCAINE
 - MEPIVACAINE HYDROCHLORIDE W/ LEVONORDEFIN, LEVONORDEFIN
 - MEPIVACAINE HYDROCHLORIDE, MEPIVACAINE HYDROCHLORIDE

GRIFFEN

- * KW GRIFFEN CO
 - BIOSCRUB, CHLORHEXIDINE GLUCONATE (OTC)

GTX INC

- * GTX INC
 - FARESTON, TOREMIFENE CITRATE

GUARDIAN DRUG

- * GUARDIAN DRUG CO INC
 - FOAMCOAT, ALUMINUM HYDROXIDE (OTC)

GUERBET

- * GUERBET LLC
 - OXILAN-300, IOXILAN
 - OXILAN-350, IOXILAN

HAEMONETICS

- * HAEMONETICS CORP
 - SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER, SODIUM CHLORIDE

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

** H **

HALOCARBON PRODS

* HALOCARBON PRODUCTS CORP
ISOFLURANE, ISOFLURANE

HALOZYME THERAP

* HALOZYME THERAPEUTICS INC
HYLENEX RECOMBINANT, HYALURONIDASE RECOMBINANT HUMAN

HAMELN PHARMS

* HAMELN PHARMACEUTICALS GMBH
PENTETATE CALCIUM TRISODIUM, PENTETATE CALCIUM TRISODIUM
PENTETATE ZINC TRISODIUM, PENTETATE ZINC TRISODIUM

HAMPTON LAINE LLC

* HAMPTON LAINE LLC
FLAVOXATE HYDROCHLORIDE, FLAVOXATE HYDROCHLORIDE

HANFORD GC

* GC HANFORD MANUFACTURING CO
AMPICILLIN AND SULBACTAM, AMPICILLIN SODIUM
CEFAZOLIN SODIUM, CEFZOLIN SODIUM
CEFUROXIME SODIUM, CEFUROXIME SODIUM

HARMONY LABS

* HARMONY LABORATORIES
THEROXIDIL, MINOXIDIL (OTC)

HARVEST PHARMS

* HARVEST PHARMACEUTICALS INC
ERGOMAR, ERGOTAMINE TARTRATE

HEALTHPOINT

* HEALTHPOINT LTD
AKNE-MYCIN, ERYTHROMYCIN
CETACORT, HYDROCORTISONE
CLODERM, CLOCORTOLONE PIVALATE
CORMAX, CLOBETASOL PROPIONATE
ECONAZOLE NITRATE, ECONAZOLE NITRATE
EMBELINE E, CLOBETASOL PROPIONATE
EMBELINE, CLOBETASOL PROPIONATE
NUTRACORT, HYDROCORTISONE

HELSINN HLTHCARE

* HELSINN HEALTHCARE SA
ALOXI, PALONOSETRON HYDROCHLORIDE

HERCON LABS

* HERCON LABORATORIES CORP
NITROGLYCERIN, NITROGLYCERIN

HEYL CHEMISCH

* HEYL CHEMISCH PHARMAZEUTISCHE FABRIK
RADIOGARDASE (PRUSSIAN BLUE), FERRIC HEXACYANOFERRATE(II)

HI TECH PHARMA

* HI TECH PHARMACAL CO INC
ACETAMINOPHEN AND CODEINE PHOSPHATE, ACETAMINOPHEN
ACYCLOVIR, ACYCLOVIR
ALBUTEROL SULFATE, ALBUTEROL SULFATE
AMANTADINE HYDROCHLORIDE, AMANTADINE HYDROCHLORIDE
CHLORHEXIDINE GLUCONATE, CHLORHEXIDINE GLUCONATE
CIMETIDINE HYDROCHLORIDE, CIMETIDINE HYDROCHLORIDE
FLUOXETINE HYDROCHLORIDE, FLUOXETINE HYDROCHLORIDE
HYDROXYZINE HYDROCHLORIDE, HYDROXYZINE HYDROCHLORIDE
LACTULOSE, LACTULOSE
LIDOCAINE AND PRILOCAINE, LIDOCAINE
LIDOCAINE HYDROCHLORIDE, LIDOCAINE HYDROCHLORIDE
LOPERAMIDE HYDROCHLORIDE, LOPERAMIDE HYDROCHLORIDE (OTC)
MIDAZOLAM HYDROCHLORIDE, MIDAZOLAM HYDROCHLORIDE
MINOXIDIL (FOR MEN), MINOXIDIL (OTC)

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

** H **

* HI TECH PHARMACAL CO INC
OFLOXACIN, OFLOXACIN
PREDNISOLONE SODIUM PHOSPHATE, PREDNISOLONE SODIUM PHOSPHATE
PREDNISOLONE, PREDNISOLONE
PROMETHAZINE HYDROCHLORIDE AND CODEINE PHOSPHATE, CODEINE PHOSPHATE
PROMETHAZINE HYDROCHLORIDE AND DEXTROMETHORPHAN HYDROBROMIDE, DEXTROMETHORPHAN
HYDROBROMIDE
PROMETHAZINE HYDROCHLORIDE, PROMETHAZINE HYDROCHLORIDE
SULFAMETHOXAZOLE AND TRIMETHOPRIM, SULFAMETHOXAZOLE
TIMOLOL MALEATE, TIMOLOL MALEATE

HIGH TECH PHARMA

* HIGH TECHNOLOGY PHARMACAL CO INC
VALPROIC ACID, VALPROIC ACID

HIKMA

* HIKMA FARMACEUTICA LDA
CEFOTAXIME, CEFOTAXIME SODIUM
* HIKMA PHARMACEUTICALS
AMOXICILLIN, AMOXICILLIN
CEFADROXIL, CEFADROXIL/CEFADROXIL HEMIHYDRATE
CEPHALEXIN, CEPHALEXIN
CIPROFLOXACIN, CIPROFLOXACIN HYDROCHLORIDE
GLYBURIDE (MICRONIZED), GLYBURIDE
NAPROXEN SODIUM, NAPROXEN SODIUM

HIKMA FARMACEUTICA

* HIKMA FARMACEUTICA (PORTUGAL) SA
ONDANSETRON HYDROCHLORIDE PRESERVATIVE FREE, ONDANSETRON HYDROCHLORIDE
ONDANSETRON HYDROCHLORIDE, ONDANSETRON HYDROCHLORIDE
PHENYTOIN SODIUM, PHENYTOIN SODIUM
* HIKMA FARMACEUTICA PORTUGAL LDA
CEFAZOLIN SODIUM, CEFAZOLIN SODIUM
CEFUROXIME, CEFUROXIME SODIUM
FLUCONAZOLE IN SODIUM CHLORIDE 0.9%, FLUCONAZOLE

HIKMA PHARMS

* HIKMA PHARMACEUTICALS
AMOXICILLIN AND CLAVULANATE POTASSIUM, AMOXICILLIN

HILL DERMAC

* HILL DERMACEUTICALS INC
DERMA-SMOOTH/FS, FLUOCINOLONE ACETONIDE
FLUOCINOLONE ACETONIDE, FLUOCINOLONE ACETONIDE

HITECH PHARMA

* HITECH PHARMACAL CORP
CIPROFLOXACIN, CIPROFLOXACIN HYDROCHLORIDE

HLR

* HLR TECHNOLOGY
ACCUTANE, ISOTRETINOIN
INVIRASE, SAQUINAVIR MESYLATE
ROCEPHIN W/ DEXTROSE IN PLASTIC CONTAINER, CEFTRIAXONE SODIUM
ROCEPHIN, CEFTRIAXONE SODIUM
ROMAZICON, FLUMAZENIL
XELODA, CAPECITABINE
XENICAL, ORLISTAT

HOLOPACK INTL

* HOLOPACK INTERNATIONAL CORP
IPRATROPIUM BROMIDE, IPRATROPIUM BROMIDE

HOSPIRA

* HOSPIRA INC
ACETAZOLAMIDE SODIUM, ACETAZOLAMIDE SODIUM
ACETIC ACID 0.25% IN PLASTIC CONTAINER, ACETIC ACID, GLACIAL
ACETYLCYSTEINE, ACETYLCYSTEINE
ACYCLOVIR SODIUM, ACYCLOVIR SODIUM
A-HYDROCORT, HYDROCORTISONE SODIUM SUCCINATE

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

**** H ****

* HOSPIRA INC
ALCOHOL 5% IN D5-W, ALCOHOL
ALFENTANIL, ALFENTANIL HYDROCHLORIDE
A-METHAPRED, METHYLPREDNISOLONE SODIUM SUCCINATE
AMIDATE, ETOMIDATE
AMIKACIN SULFATE IN SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER, AMIKACIN SULFATE
AMIKACIN SULFATE, AMIKACIN SULFATE
AMINOCAPROIC ACID IN PLASTIC CONTAINER, AMINOCAPROIC ACID
AMINOCAPROIC ACID, AMINOCAPROIC ACID
AMINOPHYLLINE IN SODIUM CHLORIDE 0.45%, AMINOPHYLLINE
AMINOPHYLLINE, AMINOPHYLLINE
AMINOSYN 10% (PH6), AMINO ACIDS
AMINOSYN 10%, AMINO ACIDS
AMINOSYN 3.5% M, AMINO ACIDS
AMINOSYN 3.5%, AMINO ACIDS
AMINOSYN 5%, AMINO ACIDS
AMINOSYN 7% (PH6), AMINO ACIDS
AMINOSYN 7% W/ ELECTROLYTES, AMINO ACIDS
AMINOSYN 7%, AMINO ACIDS
AMINOSYN 8.5% (PH6), AMINO ACIDS
AMINOSYN 8.5% W/ ELECTROLYTES, AMINO ACIDS
AMINOSYN 8.5%, AMINO ACIDS
AMINOSYN II 10% IN PLASTIC CONTAINER, AMINO ACIDS
AMINOSYN II 10% W/ ELECTROLYTES, AMINO ACIDS
AMINOSYN II 10%, AMINO ACIDS
AMINOSYN II 15% IN PLASTIC CONTAINER, AMINO ACIDS
AMINOSYN II 3.5% IN DEXTROSE 25% IN PLASTIC CONTAINER, AMINO ACIDS
AMINOSYN II 3.5% IN DEXTROSE 5% IN PLASTIC CONTAINER, AMINO ACIDS
AMINOSYN II 3.5% M IN DEXTROSE 5% IN PLASTIC CONTAINER, AMINO ACIDS
AMINOSYN II 3.5% W/ ELECTROLYTES IN DEXTROSE 25% W/ CALCIUM IN PLASTIC CONTAINER, AMINO ACIDS
AMINOSYN II 4.25% IN DEXTROSE 10% IN PLASTIC CONTAINER, AMINO ACIDS
AMINOSYN II 4.25% IN DEXTROSE 20% IN PLASTIC CONTAINER, AMINO ACIDS
AMINOSYN II 4.25% IN DEXTROSE 25% IN PLASTIC CONTAINER, AMINO ACIDS
AMINOSYN II 4.25% M IN DEXTROSE 10% IN PLASTIC CONTAINER, AMINO ACIDS
AMINOSYN II 4.25% W/ ELECTROLYTES IN DEXTROSE 20% W/ CALCIUM IN PLASTIC CONTAINER, AMINO ACIDS
AMINOSYN II 4.25% W/ ELECTROLYTES IN DEXTROSE 25% W/ CALCIUM IN PLASTIC CONTAINER, AMINO ACIDS
AMINOSYN II 5% IN DEXTROSE 25% IN PLASTIC CONTAINER, AMINO ACIDS
AMINOSYN II 7%, AMINO ACIDS
AMINOSYN II 8.5% W/ ELECTROLYTES, AMINO ACIDS
AMINOSYN II 8.5%, AMINO ACIDS
AMINOSYN-HBC 7%, AMINO ACIDS
AMINOSYN-HF 8%, AMINO ACIDS
AMINOSYN-PF 10%, AMINO ACIDS
AMINOSYN-PF 7%, AMINO ACIDS
AMINOSYN-RF 5.2%, AMINO ACIDS
AMIODARONE HYDROCHLORIDE, AMIODARONE HYDROCHLORIDE
AMMONIUM CHLORIDE IN PLASTIC CONTAINER, AMMONIUM CHLORIDE
AMRINONE, INAMRINONE LACTATE
ATRACURIUM BESYLATE PRESERVATIVE FREE, ATRACURIUM BESYLATE
ATRACURIUM BESYLATE, ATRACURIUM BESYLATE
ATROPINE SULFATE ANSYR PLASTIC SYRINGE, ATROPINE SULFATE
BACTERIOSTATIC SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER, SODIUM CHLORIDE
BACTERIOSTATIC WATER FOR INJECTION IN PLASTIC CONTAINER, WATER FOR INJECTION, STERILE
BRETYLIUM TOSYLATE IN DEXTROSE 5% IN PLASTIC CONTAINER, BRETYLIUM TOSYLATE
BRETYLIUM TOSYLATE IN PLASTIC CONTAINER, BRETYLIUM TOSYLATE
BUMETANIDE, BUMETANIDE
BUPIVACAINE HYDROCHLORIDE AND EPINEPHRINE, BUPIVACAINE HYDROCHLORIDE
BUPIVACAINE HYDROCHLORIDE W/EPINEPHRINE, BUPIVACAINE HYDROCHLORIDE
BUPIVACAINE HYDROCHLORIDE, BUPIVACAINE HYDROCHLORIDE
BUPIVACAINE, BUPIVACAINE HYDROCHLORIDE
BUPRENORPHINE HYDROCHLORIDE, BUPRENORPHINE HYDROCHLORIDE

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

** H **

* HOSPIRA INC

BUTORPHANOL TARTRATE PRESERVATIVE FREE, BUTORPHANOL TARTRATE

BUTORPHANOL TARTRATE, BUTORPHANOL TARTRATE

CALCIUM CHLORIDE 10% IN PLASTIC CONTAINER, CALCIUM CHLORIDE

CARBOCAINE, MEPIVACAINE HYDROCHLORIDE

CHLOROPROCAINE HYDROCHLORIDE, CHLOROPROCAINE HYDROCHLORIDE

CHROMIC CHLORIDE IN PLASTIC CONTAINER, CHROMIC CHLORIDE

CIMETIDINE HYDROCHLORIDE IN SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER, CIMETIDINE HYDROCHLORIDE

CIMETIDINE HYDROCHLORIDE, CIMETIDINE HYDROCHLORIDE

CIPROFLOXACIN, CIPROFLOXACIN

CLINDAMYCIN PHOSPHATE, CLINDAMYCIN PHOSPHATE

CORLOPAM, FENOLDOPAM MESYLATE

CUPRIC CHLORIDE IN PLASTIC CONTAINER, CUPRIC CHLORIDE

DEFEROXAMINE MESYLATE, DEFEROXAMINE MESYLATE

DEMOROL, MEPERIDINE HYDROCHLORIDE

DESMOPRESSIN ACETATE, DESMOPRESSIN ACETATE

DEXTROSE 10% IN PLASTIC CONTAINER, DEXTROSE

DEXTROSE 2.5% AND SODIUM CHLORIDE 0.45% IN PLASTIC CONTAINER, DEXTROSE

DEXTROSE 20% IN PLASTIC CONTAINER, DEXTROSE

DEXTROSE 25%, DEXTROSE

DEXTROSE 30% IN PLASTIC CONTAINER, DEXTROSE

DEXTROSE 40% IN PLASTIC CONTAINER, DEXTROSE

DEXTROSE 5% AND LACTATED RINGER'S IN PLASTIC CONTAINER, CALCIUM CHLORIDE

DEXTROSE 5% AND RINGER'S IN PLASTIC CONTAINER, CALCIUM CHLORIDE

DEXTROSE 5% AND SODIUM CHLORIDE 0.225% IN PLASTIC CONTAINER, DEXTROSE

DEXTROSE 5% AND SODIUM CHLORIDE 0.3% IN PLASTIC CONTAINER, DEXTROSE

DEXTROSE 5% AND SODIUM CHLORIDE 0.45% IN PLASTIC CONTAINER, DEXTROSE

DEXTROSE 5% AND SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER, DEXTROSE

DEXTROSE 5% IN PLASTIC CONTAINER, DEXTROSE

DEXTROSE 50% IN PLASTIC CONTAINER, DEXTROSE

DEXTROSE 60% IN PLASTIC CONTAINER, DEXTROSE

DEXTROSE 70% IN PLASTIC CONTAINER, DEXTROSE

DIAZEPAM, DIAZEPAM

DIGOXIN PEDIATRIC, DIGOXIN

DIGOXIN, DIGOXIN

DILTIAZEM HYDROCHLORIDE, DILTIAZEM HYDROCHLORIDE

DIPHENHYDRAMINE HYDROCHLORIDE, DIPHENHYDRAMINE HYDROCHLORIDE

DIPYRIDAMOLE, DIPYRIDAMOLE

DOBUTAMINE HYDROCHLORIDE IN DEXTROSE 5% IN PLASTIC CONTAINER, DOBUTAMINE HYDROCHLORIDE

DOBUTAMINE HYDROCHLORIDE IN DEXTROSE 5%, DOBUTAMINE HYDROCHLORIDE

DOBUTAMINE HYDROCHLORIDE, DOBUTAMINE HYDROCHLORIDE

DOPAMINE HYDROCHLORIDE IN DEXTROSE 5% IN PLASTIC CONTAINER, DOPAMINE HYDROCHLORIDE

DOPAMINE HYDROCHLORIDE, DOPAMINE HYDROCHLORIDE

DROPERIDOL, DROPERIDOL

EDROPHONIUM CHLORIDE, EDROPHONIUM CHLORIDE

ENALAPRILAT, ENALAPRILAT

ENDRATE, EDETATE DISODIUM

ERYTHROCIN, ERYTHROMYCIN LACTOBIONATE

ETOPOSIDE, ETOPOSIDE

FAMOTIDINE, FAMOTIDINE

FENTANYL CITRATE PRESERVATIVE FREE, FENTANYL CITRATE

FENTANYL CITRATE, FENTANYL CITRATE

FLUCONAZOLE IN DEXTROSE 5% IN PLASTIC CONTAINER, FLUCONAZOLE

FLUCONAZOLE IN SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER, FLUCONAZOLE

FOSCARNET SODIUM, FOSCARNET SODIUM

FUROSEMIDE, FUROSEMIDE

GENTAMICIN SULFATE IN SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER, GENTAMICIN SULFATE

GENTAMICIN SULFATE, GENTAMICIN SULFATE

GLYCINE 1.5% IN PLASTIC CONTAINER, GLYCINE

HEPARIN LOCK FLUSH IN PLASTIC CONTAINER, HEPARIN SODIUM

HEPARIN LOCK FLUSH, HEPARIN SODIUM

HEPARIN SODIUM 1,000 UNITS IN SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER, HEPARIN SODIUM

HEPARIN SODIUM 10,000 UNITS IN DEXTROSE 5% IN PLASTIC CONTAINER, HEPARIN SODIUM

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

**** H ****

* HOSPIRA INC

HEPARIN SODIUM 12,500 UNITS IN DEXTROSE 5% IN PLASTIC CONTAINER, HEPARIN SODIUM
HEPARIN SODIUM 12,500 UNITS IN SODIUM CHLORIDE 0.45% IN PLASTIC CONTAINER, HEPARIN SODIUM
HEPARIN SODIUM 2,000 UNITS IN SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER, HEPARIN SODIUM
HEPARIN SODIUM 20,000 UNITS IN DEXTROSE 5% IN PLASTIC CONTAINER, HEPARIN SODIUM
HEPARIN SODIUM 25,000 UNITS IN DEXTROSE 5% IN PLASTIC CONTAINER, HEPARIN SODIUM
HEPARIN SODIUM 25,000 UNITS IN SODIUM CHLORIDE 0.45% IN PLASTIC CONTAINER, HEPARIN SODIUM
HEPARIN SODIUM PRESERVATIVE FREE, HEPARIN SODIUM
HEPARIN SODIUM, HEPARIN SODIUM
HYDROMORPHONE HYDROCHLORIDE, HYDROMORPHONE HYDROCHLORIDE
HYDROXYZINE HYDROCHLORIDE, HYDROXYZINE HYDROCHLORIDE
IONOSOL B AND DEXTROSE 5% IN PLASTIC CONTAINER, DEXTROSE
IONOSOL MB AND DEXTROSE 5% IN PLASTIC CONTAINER, DEXTROSE
IONOSOL T AND DEXTROSE 5% IN PLASTIC CONTAINER, DEXTROSE
IOPAMIDOL-200 IN PLASTIC CONTAINER, IOPAMIDOL
IOPAMIDOL-200, IOPAMIDOL
IOPAMIDOL-250 IN PLASTIC CONTAINER, IOPAMIDOL
IOPAMIDOL-250, IOPAMIDOL
IOPAMIDOL-300 IN PLASTIC CONTAINER, IOPAMIDOL
IOPAMIDOL-300, IOPAMIDOL
IOPAMIDOL-370 IN PLASTIC CONTAINER, IOPAMIDOL
IOPAMIDOL-370, IOPAMIDOL
ISOFLURANE, ISOFLURANE
ISOPROTERENOL HYDROCHLORIDE, ISOPROTERENOL HYDROCHLORIDE
ISUPREL, ISOPROTERENOL HYDROCHLORIDE
KETAMINE HYDROCHLORIDE, KETAMINE HYDROCHLORIDE
KETOROLAC TROMETHAMINE, KETOROLAC TROMETHAMINE
LABETALOL HYDROCHLORIDE, LABETALOL HYDROCHLORIDE
LACTATED RINGER'S IN PLASTIC CONTAINER, CALCIUM CHLORIDE
LEVOPHED, NOREPINEPHRINE BITARTRATE
LIDOCAINE HYDROCHLORIDE 0.2% IN DEXTROSE 5% IN PLASTIC CONTAINER, LIDOCAINE HYDROCHLORIDE
LIDOCAINE HYDROCHLORIDE 0.2% IN DEXTROSE 5%, LIDOCAINE HYDROCHLORIDE
LIDOCAINE HYDROCHLORIDE 0.4% IN DEXTROSE 5% IN PLASTIC CONTAINER, LIDOCAINE HYDROCHLORIDE
LIDOCAINE HYDROCHLORIDE 0.4% IN DEXTROSE 5%, LIDOCAINE HYDROCHLORIDE
LIDOCAINE HYDROCHLORIDE 0.8% IN DEXTROSE 5% IN PLASTIC CONTAINER, LIDOCAINE HYDROCHLORIDE
LIDOCAINE HYDROCHLORIDE 5% AND DEXTROSE 7.5%, LIDOCAINE HYDROCHLORIDE
LIDOCAINE HYDROCHLORIDE AND EPINEPHRINE, EPINEPHRINE
LIDOCAINE HYDROCHLORIDE IN PLASTIC CONTAINER, LIDOCAINE HYDROCHLORIDE
LIDOCAINE HYDROCHLORIDE PRESERVATIVE FREE IN PLASTIC CONTAINER, LIDOCAINE HYDROCHLORIDE
LIDOCAINE HYDROCHLORIDE PRESERVATIVE FREE, LIDOCAINE HYDROCHLORIDE
LIDOCAINE HYDROCHLORIDE, LIDOCAINE HYDROCHLORIDE
LIPOSYN II 10%, SAFFLOWER OIL
LIPOSYN II 20%, SAFFLOWER OIL
LIPOSYN III 10%, SOYBEAN OIL
LIPOSYN III 20%, SOYBEAN OIL
LIPOSYN III 30%, SOYBEAN OIL
LORAZEPAM, LORAZEPAM
LTA II KIT, LIDOCAINE HYDROCHLORIDE
MAGNESIUM SULFATE IN DEXTROSE 5% IN PLASTIC CONTAINER, MAGNESIUM SULFATE
MAGNESIUM SULFATE IN PLASTIC CONTAINER, MAGNESIUM SULFATE
MAGNESIUM SULFATE, MAGNESIUM SULFATE
MANGANESE CHLORIDE IN PLASTIC CONTAINER, MANGANESE CHLORIDE
MANNITOL 10% IN PLASTIC CONTAINER, MANNITOL
MANNITOL 15% IN PLASTIC CONTAINER, MANNITOL
MANNITOL 20% IN PLASTIC CONTAINER, MANNITOL
MANNITOL 25%, MANNITOL
MANNITOL 5% IN PLASTIC CONTAINER, MANNITOL
MARCAINE HYDROCHLORIDE PRESERVATIVE FREE, BUPIVACAINE HYDROCHLORIDE
MARCAINE HYDROCHLORIDE W/ EPINEPHRINE PRESERVATIVE FREE, BUPIVACAINE HYDROCHLORIDE
MARCAINE HYDROCHLORIDE W/ EPINEPHRINE, BUPIVACAINE HYDROCHLORIDE
MARCAINE HYDROCHLORIDE, BUPIVACAINE HYDROCHLORIDE
MARCAINE, BUPIVACAINE HYDROCHLORIDE
MEPERIDINE HYDROCHLORIDE PRESERVATIVE FREE, MEPERIDINE HYDROCHLORIDE
METHYLDOPATE HYDROCHLORIDE, METHYLDOPATE HYDROCHLORIDE

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

** H **

* HOSPIRA INC

METOCLOPRAMIDE HYDROCHLORIDE, METOCLOPRAMIDE HYDROCHLORIDE

METOPROLOL TARTRATE, METOPROLOL TARTRATE

METRONIDAZOLE IN PLASTIC CONTAINER, METRONIDAZOLE

MIDAZOLAM HYDROCHLORIDE, MIDAZOLAM HYDROCHLORIDE

MILRINONE LACTATE IN DEXTROSE 5% IN PLASTIC CONTAINER, MILRINONE LACTATE

MILRINONE LACTATE, MILRINONE LACTATE

MORPHINE SULFATE, MORPHINE SULFATE

NALBUPHINE HYDROCHLORIDE, NALBUPHINE HYDROCHLORIDE

NALOXONE HYDROCHLORIDE, NALOXONE HYDROCHLORIDE

NITROGLYCERIN IN DEXTROSE 5%, NITROGLYCERIN

NITROGLYCERIN, NITROGLYCERIN

NITROPRESS, SODIUM NITROPRUSSIDE

NORMOSOL-M AND DEXTROSE 5% IN PLASTIC CONTAINER, DEXTROSE

NORMOSOL-R AND DEXTROSE 5% IN PLASTIC CONTAINER, DEXTROSE

NORMOSOL-R IN PLASTIC CONTAINER, MAGNESIUM CHLORIDE

NOVAMINE 11.4%, AMINO ACIDS

NOVAMINE 15%, AMINO ACIDS

NOVOCAIN, PROCAINE HYDROCHLORIDE

ONDANSETRON HYDROCHLORIDE PRESERVATIVE FREE, ONDANSETRON HYDROCHLORIDE

ONDANSETRON HYDROCHLORIDE, ONDANSETRON HYDROCHLORIDE

PANCURONIUM BROMIDE, PANCURONIUM BROMIDE

PEDIATRIC LTA KIT, LIDOCAINE HYDROCHLORIDE

PENTAMIDINE ISETHIONATE, PENTAMIDINE ISETHIONATE

PHENYTOIN SODIUM, PHENYTOIN SODIUM

PHYSIOSOL IN PLASTIC CONTAINER, MAGNESIUM CHLORIDE

PHYSIOSOL PH 7.4 IN PLASTIC CONTAINER, MAGNESIUM CHLORIDE

PLEGISOL IN PLASTIC CONTAINER, CALCIUM CHLORIDE

POTASSIUM ACETATE IN PLASTIC CONTAINER, POTASSIUM ACETATE

POTASSIUM CHLORIDE 10MEQ IN DEXTROSE 5% AND SODIUM CHLORIDE 0.225% IN PLASTIC CONTAINER, DEXTROSE

POTASSIUM CHLORIDE 10MEQ IN DEXTROSE 5% AND SODIUM CHLORIDE 0.3% IN PLASTIC CONTAINER, DEXTROSE

POTASSIUM CHLORIDE 10MEQ IN DEXTROSE 5% AND SODIUM CHLORIDE 0.45% IN PLASTIC CONTAINER, DEXTROSE

POTASSIUM CHLORIDE 10MEQ IN DEXTROSE 5% AND SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER, DEXTROSE

POTASSIUM CHLORIDE 10MEQ IN PLASTIC CONTAINER, POTASSIUM CHLORIDE

POTASSIUM CHLORIDE 15MEQ IN DEXTROSE 5% AND SODIUM CHLORIDE 0.225% IN PLASTIC CONTAINER, DEXTROSE

POTASSIUM CHLORIDE 15MEQ IN DEXTROSE 5% AND SODIUM CHLORIDE 0.3% IN PLASTIC CONTAINER, DEXTROSE

POTASSIUM CHLORIDE 15MEQ IN DEXTROSE 5% AND SODIUM CHLORIDE 0.45% IN PLASTIC CONTAINER, DEXTROSE

POTASSIUM CHLORIDE 15MEQ IN DEXTROSE 5% AND SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER, DEXTROSE

POTASSIUM CHLORIDE 20MEQ IN DEXTROSE 5% AND LACTATED RINGER'S IN PLASTIC CONTAINER, CALCIUM CHLORIDE

POTASSIUM CHLORIDE 20MEQ IN DEXTROSE 5% AND SODIUM CHLORIDE 0.225% IN PLASTIC CONTAINER, DEXTROSE

POTASSIUM CHLORIDE 20MEQ IN DEXTROSE 5% AND SODIUM CHLORIDE 0.3% IN PLASTIC CONTAINER, DEXTROSE

POTASSIUM CHLORIDE 20MEQ IN DEXTROSE 5% AND SODIUM CHLORIDE 0.45% IN PLASTIC CONTAINER, DEXTROSE

POTASSIUM CHLORIDE 20MEQ IN DEXTROSE 5% AND SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER, DEXTROSE

POTASSIUM CHLORIDE 20MEQ IN DEXTROSE 5% IN PLASTIC CONTAINER, DEXTROSE

POTASSIUM CHLORIDE 20MEQ IN DEXTROSE 5% IN SODIUM CHLORIDE 0.3% IN PLASTIC CONTAINER, DEXTROSE

POTASSIUM CHLORIDE 20MEQ IN PLASTIC CONTAINER, POTASSIUM CHLORIDE

POTASSIUM CHLORIDE 20MEQ IN SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER, POTASSIUM CHLORIDE

POTASSIUM CHLORIDE 30MEQ IN DEXTROSE 5% AND SODIUM CHLORIDE 0.225% IN PLASTIC CONTAINER, DEXTROSE

POTASSIUM CHLORIDE 30MEQ IN DEXTROSE 5% AND SODIUM CHLORIDE 0.3% IN PLASTIC CONTAINER, DEXTROSE

POTASSIUM CHLORIDE 30MEQ IN DEXTROSE 5% AND SODIUM CHLORIDE 0.45% IN PLASTIC CONTAINER, DEXTROSE

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

** H **

* HOSPIRA INC
 POTASSIUM CHLORIDE 30MEQ IN DEXTROSE 5% AND SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER, DEXTROSE
 POTASSIUM CHLORIDE 30MEQ IN DEXTROSE 5% IN PLASTIC CONTAINER, DEXTROSE
 POTASSIUM CHLORIDE 30MEQ IN PLASTIC CONTAINER, POTASSIUM CHLORIDE
 POTASSIUM CHLORIDE 40MEQ IN DEXTROSE 5% AND LACTATED RINGER'S IN PLASTIC CONTAINER, CALCIUM CHLORIDE
 POTASSIUM CHLORIDE 40MEQ IN DEXTROSE 5% AND SODIUM CHLORIDE 0.225% IN PLASTIC CONTAINER, DEXTROSE
 POTASSIUM CHLORIDE 40MEQ IN DEXTROSE 5% AND SODIUM CHLORIDE 0.3% IN PLASTIC CONTAINER, DEXTROSE
 POTASSIUM CHLORIDE 40MEQ IN DEXTROSE 5% AND SODIUM CHLORIDE 0.45% IN PLASTIC CONTAINER, DEXTROSE
 POTASSIUM CHLORIDE 40MEQ IN DEXTROSE 5% AND SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER, DEXTROSE
 POTASSIUM CHLORIDE 40MEQ IN DEXTROSE 5% IN PLASTIC CONTAINER, DEXTROSE
 POTASSIUM CHLORIDE 40MEQ IN PLASTIC CONTAINER, POTASSIUM CHLORIDE
 POTASSIUM CHLORIDE 40MEQ IN SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER, POTASSIUM CHLORIDE
 POTASSIUM CHLORIDE 5MEQ IN DEXTROSE 5% AND SODIUM CHLORIDE 0.225% IN PLASTIC CONTAINER, DEXTROSE
 POTASSIUM CHLORIDE 5MEQ IN DEXTROSE 5% AND SODIUM CHLORIDE 0.3% IN PLASTIC CONTAINER, DEXTROSE
 POTASSIUM CHLORIDE 5MEQ IN DEXTROSE 5% AND SODIUM CHLORIDE 0.45% IN PLASTIC CONTAINER, DEXTROSE
 POTASSIUM CHLORIDE 5MEQ IN DEXTROSE 5% AND SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER, DEXTROSE
 POTASSIUM CHLORIDE, POTASSIUM CHLORIDE
 PRECEDEX, DEXMEDETOMIDINE
 PROCAINAMIDE HYDROCHLORIDE, PROCAINAMIDE HYDROCHLORIDE
 PROCAINE HYDROCHLORIDE, PROCAINE HYDROCHLORIDE
 PROCHLORPERAZINE EDISYLATE, PROCHLORPERAZINE EDISYLATE
 PROMETHAZINE HYDROCHLORIDE, PROMETHAZINE HYDROCHLORIDE
 PROPOFOL, PROPOFOL
 QUELICIN PRESERVATIVE FREE, SUCCINYLCHOLINE CHLORIDE
 QUELICIN, SUCCINYLCHOLINE CHLORIDE
 RINGER'S IN PLASTIC CONTAINER, CALCIUM CHLORIDE
 RITODRINE HYDROCHLORIDE IN DEXTROSE 5% IN PLASTIC CONTAINER, RITODRINE HYDROCHLORIDE
 RITODRINE HYDROCHLORIDE, RITODRINE HYDROCHLORIDE
 SODIUM ACETATE IN PLASTIC CONTAINER, SODIUM ACETATE, ANHYDROUS
 SODIUM BICARBONATE, SODIUM BICARBONATE
 SODIUM CHLORIDE 0.45% IN PLASTIC CONTAINER, SODIUM CHLORIDE
 SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER, SODIUM CHLORIDE
 SODIUM CHLORIDE IN PLASTIC CONTAINER, SODIUM CHLORIDE
 SODIUM LACTATE 0.167 MOLAR IN PLASTIC CONTAINER, SODIUM LACTATE
 SODIUM LACTATE IN PLASTIC CONTAINER, SODIUM LACTATE
 SODIUM PHOSPHATES IN PLASTIC CONTAINER, SODIUM PHOSPHATE, DIBASIC, HEPTAHYDRATE
 SORBITOL-MANNITOL IN PLASTIC CONTAINER, MANNITOL
 STERILE WATER FOR INJECTION IN PLASTIC CONTAINER, WATER FOR INJECTION, STERILE
 STERILE WATER IN PLASTIC CONTAINER, WATER FOR IRRIGATION, STERILE
 SULFAMETHOXAZOLE AND TRIMETHOPRIM, SULFAMETHOXAZOLE
 SULFENTANIL CITRATE, SUFENTANIL CITRATE
 TALWIN, PENTAZOCINE LACTATE
 TAZICEF, CEFTAZIDIME
 THAM, TROMETHAMINE
 THEOPHYLLINE IN DEXTROSE 5% IN PLASTIC CONTAINER, THEOPHYLLINE
 THIAMINE HYDROCHLORIDE, THIAMINE HYDROCHLORIDE
 TOBRAMYCIN SULFATE IN SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER, TOBRAMYCIN SULFATE
 TOBRAMYCIN SULFATE, TOBRAMYCIN SULFATE
 TPN ELECTROLYTES IN PLASTIC CONTAINER, CALCIUM CHLORIDE
 TRACRIUM PRESERVATIVE FREE, ATRACURIUM BESYLATE
 TRACRIUM, ATRACURIUM BESYLATE
 TRIMETHOBENZAMIDE HYDROCHLORIDE, TRIMETHOBENZAMIDE HYDROCHLORIDE
 UROLOGIC G IN PLASTIC CONTAINER, CITRIC ACID
 VANCOMYCIN HYDROCHLORIDE, VANCOMYCIN HYDROCHLORIDE
 VECURONIUM BROMIDE, VECURONIUM BROMIDE
 VERAPAMIL HYDROCHLORIDE, VERAPAMIL HYDROCHLORIDE

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

**** H ****

* HOSPIRA INC
VITAMIN K1, PHYTONADIONE
ZINC CHLORIDE IN PLASTIC CONTAINER, ZINC CHLORIDE

IBI

* ISTITUTO BIOCHIMICO ITALIANO SPA
AMPICILLIN SODIUM, AMPICILLIN SODIUM

IDENIX

* IDENIX PHARMACEUTICALS INC
TYZEKA, TELBIVUDINE

IMARX THERAP

* IMARX THERAPEUTICS
ABBOKINASE, UROKINASE

IMPAX LABS

* IMPAX LABORATORIES INC
ANAGRELIDE HYDROCHLORIDE, ANAGRELIDE HYDROCHLORIDE
BETHANECHOL CHLORIDE, BETHANECHOL CHLORIDE
BUPROPION HYDROCHLORIDE, BUPROPION HYDROCHLORIDE
CARBIDOPA AND LEVODOPA, CARBIDOPA
CHLOROQUINE PHOSPHATE, CHLOROQUINE PHOSPHATE
COLESTIPOL HYDROCHLORIDE, COLESTIPOL HYDROCHLORIDE
DANTROLENE SODIUM, DANTROLENE SODIUM
DEMECLOCYCLINE HYDROCHLORIDE, DEMECLOCYCLINE HYDROCHLORIDE
FENOFIBRATE (MICRONIZED), FENOFIBRATE
FLUDROCORTISONE ACETATE, FLUDROCORTISONE ACETATE
LORATADINE AND PSEUDOEPHEDRINE SULFATE, LORATADINE (OTC)
LORATADINE, LORATADINE (OTC)
METFORMIN HYDROCHLORIDE, METFORMIN HYDROCHLORIDE
METHOCARBAMOL, METHOCARBAMOL
METHYLTESTOSTERONE, METHYLTESTOSTERONE
MINOCYCLINE HYDROCHLORIDE, MINOCYCLINE HYDROCHLORIDE
OMEPRazole, OMEPRazole
OXYCODONE HYDROCHLORIDE, OXYCODONE HYDROCHLORIDE
PENTOXIFYLLINE, PENTOXIFYLLINE
PILOCARPINE HYDROCHLORIDE, PILOCARPINE HYDROCHLORIDE
PYRIDOSTIGMINE BROMIDE, PYRIDOSTIGMINE BROMIDE
RILUZOLE, RILUZOLE
RIMANTADINE HYDROCHLORIDE, RIMANTADINE HYDROCHLORIDE
TERBUTALINE SULFATE, TERBUTALINE SULFATE
TETRACYCLINE HYDROCHLORIDE, TETRACYCLINE HYDROCHLORIDE

IMPAX PHARMS

* IMPAX PHARMACEUTICALS
BUPROPION HYDROCHLORIDE, BUPROPION HYDROCHLORIDE
FLAVOXATE HYDROCHLORIDE, FLAVOXATE HYDROCHLORIDE
MIDODRINE HYDROCHLORIDE, MIDODRINE HYDROCHLORIDE
ORPHENADRINE CITRATE, ORPHENADRINE CITRATE
OXYBUTYNIN CHLORIDE, OXYBUTYNIN CHLORIDE
OXYCODONE HYDROCHLORIDE, OXYCODONE HYDROCHLORIDE
SOTALOL HYDROCHLORIDE, SOTALOL HYDROCHLORIDE

INALCO

* INALCO SPA
LACTULOSE, LACTULOSE

INDEVUS

* INDEVUS PHARMACEUTICALS INC
SANCTURA, TROSPIMUM CHLORIDE

INDEVUS PHARMS

* INDEVUS PHARMACEUTICALS
DELATESTRYL, TESTOSTERONE ENANTHATE

INGRAM PHARM

* INGRAM PHARMACEUTICAL CO
DRICORT, HYDROCORTISONE ACETATE
HYDROCORTISONE, HYDROCORTISONE

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

**** I ****

INO

* INO THERAPEUTICS INC
INOMAX, NITRIC OXIDE

INSIGHT PHARMS

* INSIGHT PHARMACEUTICALS CORP
NIX, PERMETHRIN (OTC)

INSMED

* INSMED INC
IPLEX, MECASERMIN RINFABATE RECOMBINANT

INST BIOCHIMIQUE

* INSTITUTE BIOCHIMIQUE SA (IBSA)
TIROSINT, LEVOTHYROXINE SODIUM

INSTITUTO BIOCHEMICO

* INSTITUTO BIOCHEMICO ITALIANO SPA
AMPICILLIN SODIUM, AMPICILLIN SODIUM

INSTITUTO BIOCHIMICO

* INSTITUTO BIOCHIMICO ITALIANO SPA
AMPICILLIN AND SULBACTAM, AMPICILLIN SODIUM

INTENDIS

* INTENDIS INC
FINACEA, AZELAIC ACID

INTERPHARM

* INTERPHARM INC
CITALOPRAM HYDROBROMIDE, CITALOPRAM HYDROBROMIDE
HYDROCODONE BITARTRATE AND ACETAMINOPHEN, ACETAMINOPHEN
HYDROCODONE BITARTRATE AND IBUPROFEN, HYDROCODONE BITARTRATE
IBUPROFEN, IBUPROFEN
IBUPROFEN, IBUPROFEN (OTC)
METFORMIN HYDROCHLORIDE, METFORMIN HYDROCHLORIDE
NAPROXEN, NAPROXEN
RANITIDINE HYDROCHLORIDE, RANITIDINE HYDROCHLORIDE
SULFAMETHOXAZOLE AND TRIMETHOPRIM, SULFAMETHOXAZOLE

INTL MEDICATED

* INTERNATIONAL MEDICATED SYSTEMS LTD
BUPIVACAINE HYDROCHLORIDE PRESERVATIVE FREE, BUPIVACAINE HYDROCHLORIDE
MIDAZOLAM HYDROCHLORIDE, MIDAZOLAM HYDROCHLORIDE
MILRINONE LACTATE, MILRINONE LACTATE

INTL MEDICATION

* INTERNATIONAL MEDICATION SYSTEM
AMINOPHYLLINE, AMINOPHYLLINE
BRETYLIUM TOSYLATE, BRETYLIUM TOSYLATE
DIPHENHYDRAMINE HYDROCHLORIDE PRESERVATIVE FREE, DIPHENHYDRAMINE HYDROCHLORIDE
FUROSEMIDE, FUROSEMIDE
ISOPROTERENOL HYDROCHLORIDE, ISOPROTERENOL HYDROCHLORIDE
LARYNG-O-JET KIT, LIDOCAINE HYDROCHLORIDE
LIDOCAINE HYDROCHLORIDE PRESERVATIVE FREE, LIDOCAINE HYDROCHLORIDE
LIDOCAINE HYDROCHLORIDE, LIDOCAINE HYDROCHLORIDE
MANNITOL 25%, MANNITOL
MEPERIDINE HYDROCHLORIDE PRESERVATIVE FREE, MEPERIDINE HYDROCHLORIDE
NALOXONE HYDROCHLORIDE, NALOXONE HYDROCHLORIDE
PHYTONADIONE, PHYTONADIONE
POTASSIUM CHLORIDE, POTASSIUM CHLORIDE
PROCAINAMIDE HYDROCHLORIDE, PROCAINAMIDE HYDROCHLORIDE
VERAPAMIL HYDROCHLORIDE, VERAPAMIL HYDROCHLORIDE
* INTERNATIONAL MEDICATION SYSTEMS LTD
DILTIAZEM HYDROCHLORIDE, DILTIAZEM HYDROCHLORIDE
FUROSEMIDE, FUROSEMIDE
MIDAZOLAM HYDROCHLORIDE, MIDAZOLAM HYDROCHLORIDE

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

** I **

INTL MEDICATION SYS

* INTERNATIONAL MEDICATION SYSTEMS LTD
AMIODARONE HYDROCHLORIDE, AMIODARONE HYDROCHLORIDE
LORAZEPAM, LORAZEPAM

INVAGEN PHARMS

* INVAGEN PHARMACEUTICALS INC
CITALOPRAM HYDROBROMIDE, CITALOPRAM HYDROBROMIDE
FOSINOPRIL SODIUM, FOSINOPRIL SODIUM
GEMFIBROZIL, GEMFIBROZIL
GLIMEPIRIDE, GLIMEPIRIDE
ZONISAMIDE, ZONISAMIDE

INWOOD LABS

* INWOOD LABORATORIES INC SUB FOREST LABORATORIES INC
CARBAMAZEPINE, CARBAMAZEPINE
ISOSORBIDE DINITRATE, ISOSORBIDE DINITRATE
THEOCHRON, THEOPHYLLINE
THEOPHYLLINE, THEOPHYLLINE

IPR

* IPR PHARMACEUTICALS INC
ATENOLOL AND CHLORTHALIDONE, ATENOLOL
ATENOLOL, ATENOLOL
ZOMIG, ZOLMITRIPTAN

ISO TEX

* ISO TEX DIAGNOSTICS INC
JEANATOPE, ALBUMIN IODINATED I-125 SERUM
MEGATOPE, ALBUMIN IODINATED I-131 SERUM

ISTA PHARMS

* ISTA PHARMACEUTICALS
ISTALOL, TIMOLOL MALEATE
VITRASE, HYALURONIDASE
XIBROM, BROMFENAC SODIUM

IVAX PHARMS

* IVAX PHARMACEUTICALS INC
ACYCLOVIR, ACYCLOVIR
ALBUTEROL SULFATE, ALBUTEROL SULFATE
ALBUTEROL, ALBUTEROL
ALLAY, ACETAMINOPHEN
ALPRAZOLAM, ALPRAZOLAM
ANAGRELIDE HYDROCHLORIDE, ANAGRELIDE HYDROCHLORIDE
BACLOFEN, BACLOFEN
BENAZEPRIL HYDROCHLORIDE AND HYDROCHLOROTHIAZIDE, BENAZEPRIL HYDROCHLORIDE
BENAZEPRIL HYDROCHLORIDE, BENAZEPRIL HYDROCHLORIDE
BRIMONIDINE TARTRATE, BRIMONIDINE TARTRATE
BUMETANIDE, BUMETANIDE
CAPTOPRIL AND HYDROCHLOROTHIAZIDE, CAPTOPRIL
CAPTOPRIL, CAPTOPRIL
CEFACLOR, CEFACLOL
CEFADROXIL, CEFADROXIL/CEFADROXIL HEMIHYDRATE
CEPHALEXIN, CEPHALEXIN
CILOSTAZOL, CILOSTAZOL
CIMETIDINE, CIMETIDINE
CIMETIDINE, CIMETIDINE (OTC)
CIPROFLOXACIN, CIPROFLOXACIN HYDROCHLORIDE
CITALOPRAM HYDROBROMIDE, CITALOPRAM HYDROBROMIDE
CLARITHROMYCIN, CLARITHROMYCIN
CLOZAPINE, CLOZAPINE
CROMOLYN SODIUM, CROMOLYN SODIUM
C-SOLVE-2, ERYTHROMYCIN
CYCLOSPORINE, CYCLOSPORINE
CYPROHEPTADINE HYDROCHLORIDE, CYPROHEPTADINE HYDROCHLORIDE
DIAZEPAM, DIAZEPAM

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

**** I ****

* IVAX PHARMACEUTICALS INC
DILTIAZEM HYDROCHLORIDE, DILTIAZEM HYDROCHLORIDE
DIPHENOXYLATE HYDROCHLORIDE W/ ATROPINE SULFATE, ATROPINE SULFATE
DOXAZOSIN MESYLATE, DOXAZOSIN MESYLATE
DOXYCYCLINE HYCLATE, DOXYCYCLINE HYCLATE
ESCITALOPRAM OXALATE, ESCITALOPRAM OXALATE
ETODOLAC, ETODOLAC
FAMOTIDINE, FAMOTIDINE
FAMOTIDINE, FAMOTIDINE (OTC)
FENOPROFEN CALCIUM, FENOPROFEN CALCIUM
FINASTERIDE, FINASTERIDE
FLUCONAZOLE, FLUCONAZOLE
FLUOXETINE HYDROCHLORIDE, FLUOXETINE HYDROCHLORIDE
FLUOXETINE, FLUOXETINE HYDROCHLORIDE
FLUTAMIDE, FLUTAMIDE
FLUVOXAMINE MALEATE, FLUVOXAMINE MALEATE
FUROSEMIDE, FUROSEMIDE
GABAPENTIN, GABAPENTIN
GLIPIZIDE, GLIPIZIDE
GLYBURIDE AND METFORMIN HYDROCHLORIDE, GLYBURIDE
GUANABENZ ACETATE, GUANABENZ ACETATE
HYDROCHLOROTHIAZIDE, HYDROCHLOROTHIAZIDE
HYDROCODONE BITARTRATE AND ACETAMINOPHEN, ACETAMINOPHEN
HYDROCODONE BITARTRATE AND HOMATROPINE METHYLBROMIDE, HOMATROPINE METHYLBROMIDE
HYDROGENATED ERGOT ALKALOIDS, ERGOLOID MESYLATES
HYDROXYZINE PAMOATE, HYDROXYZINE PAMOATE
IBUPROFEN, IBUPROFEN (OTC)
INDAPAMIDE, INDAPAMIDE
IPRATROPIUM BROMIDE, IPRATROPIUM BROMIDE
LABETALOL HYDROCHLORIDE, LABETALOL HYDROCHLORIDE
LISINOPRIL AND HYDROCHLOROTHIAZIDE, HYDROCHLOROTHIAZIDE
LISINOPRIL, LISINOPRIL
LORAZEPAM, LORAZEPAM
MECLIZINE HYDROCHLORIDE, MECLIZINE HYDROCHLORIDE
METFORMIN HYDROCHLORIDE, METFORMIN HYDROCHLORIDE
METHOCARBAMOL AND ASPIRIN, ASPIRIN
METHYCLOTHIAZIDE, METHYCLOTHIAZIDE
METHYLDOPA, METHYLDOPA
METRONIDAZOLE, METRONIDAZOLE
MIRTAZAPINE, MIRTAZAPINE
MISOPROSTOL, MISOPROSTOL
NADOLOL, NADOLOL
NEFAZODONE HYDROCHLORIDE, NEFAZODONE HYDROCHLORIDE
NITROFURANTOIN, NITROFURANTOIN, MACROCRYSTALLINE
OXAPROZIN, OXAPROZIN
OXAZEPAM, OXAZEPAM
PERGOLIDE MESYLATE, PERGOLIDE MESYLATE
PERPHENAZINE, PERPHENAZINE
PINDOLOL, PINDOLOL
PRAZOSIN HYDROCHLORIDE, PRAZOSIN HYDROCHLORIDE
PREDNISOLONE, PREDNISOLONE
PROBENECID AND COLCHICINE, COLCHICINE
PROBENECID, PROBENECID
PROCAINAMIDE HYDROCHLORIDE, PROCAINAMIDE HYDROCHLORIDE
PROCHLORPERAZINE MALEATE, PROCHLORPERAZINE MALEATE
PROMPT PHENYTOIN SODIUM, PHENYTOIN SODIUM
PROPOXYPHENE NAPSYLATE AND ACETAMINOPHEN, ACETAMINOPHEN
RANITIDINE HYDROCHLORIDE, RANITIDINE HYDROCHLORIDE
RANITIDINE, RANITIDINE HYDROCHLORIDE (OTC)
SELEGILINE HYDROCHLORIDE, SELEGILINE HYDROCHLORIDE
SERTRALINE HYDROCHLORIDE, SERTRALINE HYDROCHLORIDE
SIMVASTATIN, SIMVASTATIN
SULFINPYRAZONE, SULFINPYRAZONE
SULFISOXAZOLE, SULFISOXAZOLE

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

**** I ****

* IVAX PHARMACEUTICALS INC
TAMOXIFEN CITRATE, TAMOXIFEN CITRATE
TETRACYCLINE HYDROCHLORIDE, TETRACYCLINE HYDROCHLORIDE
THIORIDAZINE HYDROCHLORIDE, THIORIDAZINE HYDROCHLORIDE
TIZANIDINE HYDROCHLORIDE, TIZANIDINE HYDROCHLORIDE
TOLAZAMIDE, TOLAZAMIDE
TOLBUTAMIDE, TOLBUTAMIDE
TOLMETIN SODIUM, TOLMETIN SODIUM
TRIAMTERENE AND HYDROCHLOROTHIAZIDE, HYDROCHLOROTHIAZIDE
VERAPAMIL HYDROCHLORIDE, VERAPAMIL HYDROCHLORIDE

IVAX RES

* IVAX RESEARCH INC
NASAREL, FLUNISOLIDE
PROAIR HFA, ALBUTEROL SULFATE

J AND J

* JOHNSON AND JOHNSON CONSUMER PRODUCTS INC
ERYCETTE, ERYTHROMYCIN
GRIFULVIN V, GRISEOFULVIN, MICROCRYSTALLINE
* JOHNSON AND JOHNSON MEDICAL INC
MICRODERM, CHLORHEXIDINE GLUCONATE (OTC)
PREVACARE R, CHLORHEXIDINE GLUCONATE (OTC)

JACOBUS

* JACOBUS PHARMACEUTICAL CO
DAPSONE, DAPSONE
PASER, AMINOSALICYLIC ACID

JANSSEN

* JANSSEN PHARMACEUTICA INC
RAZADYNE ER, GALANTAMINE HYDROBROMIDE

JANSSEN LP

* JANSSEN LP
INVEGA, PALIPERIDONE

JANSSEN PHARMA

* JANSSEN PHARMACEUTICA PRODUCTS LP
NIZORAL, KETOCONAZOLE
RAZADYNE, GALANTAMINE HYDROBROMIDE
RISPERDAL CONSTA, RISPERIDONE
RISPERDAL, RISPERIDONE
SPORANOX, ITRACONAZOLE

JAZZ

* JAZZ PHARMACEUTICALS
ANTIZOL, FOMEPIZOLE
CYSTADANE, BETAINE HYDROCHLORIDE
XYREM, SODIUM OXYBATE

JDS PHARMS

* JDS PHARMACEUTICALS LLC
LITHOBID, LITHIUM CARBONATE
PEXEVA, PAROXETINE MESYLATE

JOHN O BUTLER CO

* JOHN O BUTLER CO
CHLORHEXIDINE GLUCONATE, CHLORHEXIDINE GLUCONATE

JOHNSON AND JOHNSON

* JOHNSON AND JOHNSON CONSUMER COMPANIES INC
CENTANY, MUPIROCIN
MONISTAT-DERM, MICONAZOLE NITRATE
RENOVA, TRETINOIN
RETIN-A MICRO, TRETINOIN
RETIN-A, TRETINOIN
SPECTAZOLE, ECONAZOLE NITRATE
* JOHNSON AND JOHNSON CONSUMER PRODUCTS WORLDWIDE
ERTACZO, SERTACONAZOLE NITRATE

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

** J **

- * JOHNSON AND JOHNSON GROUP CONSUMER COMPANIES
 - MEN'S ROGAINE, MINOXIDIL (OTC)
 - ROGAINE (FOR MEN), MINOXIDIL (OTC)
 - ROGAINE (FOR WOMEN), MINOXIDIL (OTC)
 - ROGAINE EXTRA STRENGTH (FOR MEN), MINOXIDIL (OTC)
 - VISINE L.R., OXYMETAZOLINE HYDROCHLORIDE (OTC)
 - VISINE-A, NAPHAZOLINE HYDROCHLORIDE (OTC)

JOHNSON RW

- * RW JOHNSON PHARMACEUTICAL RESEARCH INSTITUTE
 - ORTHO CYCLEN-28, ETHINYL ESTRADIOL
 - ORTHO TRI-CYCLEN, ETHINYL ESTRADIOL

JONES PHARMA

- * JONES PHARMA INC SUB KING PHARMACEUTICALS INC
 - LEVOXYL, LEVOTHYROXINE SODIUM
 - TRIOSTAT, LIOTHYRONINE SODIUM

JUBILANT PHARMS

- * JUBILANT PHARMACEUTICALS INC
 - CARBAMAZEPINE, CARBAMAZEPINE
 - CYCLOBENZAPRINE HYDROCHLORIDE, CYCLOBENZAPRINE HYDROCHLORIDE
 - FOLIC ACID, FOLIC ACID
 - INDAPAMIDE, INDAPAMIDE
 - METHYLPREDNISOLONE, METHYLPREDNISOLONE
 - PREDNISONE, PREDNISONE
 - PROCHLORPERAZINE MALEATE, PROCHLORPERAZINE MALEATE
 - TERAZOSIN HYDROCHLORIDE, TERAZOSIN HYDROCHLORIDE

JVL

- * JVL CORP
 - METOCLOPRAMIDE, METOCLOPRAMIDE HYDROCHLORIDE

KALI LABS

- * KALI LABORATORIES INC
 - CAPTOPRIL, CAPTOPRIL
 - CITALOPRAM HYDROBROMIDE, CITALOPRAM HYDROBROMIDE
 - CLONAZEPAM, CLONAZEPAM
 - GLIPIZIDE, GLIPIZIDE
 - GLYCOPYRROLATE, GLYCOPYRROLATE
 - LEFLUNOMIDE, LEFLUNOMIDE
 - METRONIDAZOLE, METRONIDAZOLE
 - NYSTATIN, NYSTATIN
 - ONDANSETRON, ONDANSETRON
 - POLYETHYLENE GLYCOL 3350, POLYETHYLENE GLYCOL 3350
 - TRAMADOL HYDROCHLORIDE AND ACETAMINOPHEN, ACETAMINOPHEN
 - TRANLYCYPROMINE SULFATE, TRANLYCYPROMINE SULFATE
 - VERAPAMIL HYDROCHLORIDE, VERAPAMIL HYDROCHLORIDE

KENDALL LP

- * KENDALL CO LP
 - THERMAZENE, SILVER SULFADIAZINE

KENWOOD LABS

- * KENWOOD LABORATORIES DIV BRADLEY PHARMACEUTICALS
 - CARMOL HC, HYDROCORTISONE ACETATE
 - TYZINE, TETRAHYDROZOLINE HYDROCHLORIDE

KEY PHARMS

- * KEY PHARMACEUTICALS INC SUB SCHERING PLOUGH CORP
 - K-DUR 10, POTASSIUM CHLORIDE
 - K-DUR 20, POTASSIUM CHLORIDE
 - NITRO-DUR, NITROGLYCERIN

KIEL

- * KIEL LABORATORIES INC
 - ORPHENADRINE CITRATE, ORPHENADRINE CITRATE

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

**** K ****

KING PHARMS

- * KING PHARMACEUTICALS INC
ALTACE, RAMIPRIL
BICILLIN C-R 900/300, PENICILLIN G BENZATHINE
BICILLIN C-R, PENICILLIN G BENZATHINE
BICILLIN L-A, PENICILLIN G BENZATHINE
BREVITAL SODIUM, METHOHEXITAL SODIUM
CORGARD, NADOLOL
CORZIDE, BENDROFLUMETHIAZIDE
CYTOMEL, LIOTHYRONINE SODIUM
DELESTROGEN, ESTRADIOL VALERATE
FLORINEF, FLUDROCORTISONE ACETATE
HUMATIN, PAROMOMYCIN SULFATE
INTAL, CROMOLYN SODIUM
PENICILLIN G PROCAINE, PENICILLIN G PROCAINE
PROCANBID, PROCAINAMIDE HYDROCHLORIDE
SILVADENE, SILVER SULFADIAZINE
SKELAXIN, METAXALONE
SYNERCID, DALFOPRISTIN
TIGAN, TRIMETHOBENZAMIDE HYDROCHLORIDE
TILADE, NEDOCROMIL SODIUM
- * KING PHARMACEUTICALS RESEARCH AND DEVELOPMENT INC SUB KING PHARMACEUTICALS INC
SONATA, ZALEPLON
TAPAZOLE, METHIMAZOLE
TUSSIGON, HOMATROPINE METHYLBROMIDE

KOS LIFE

- * KOS LIFE SCIENCES INC
ADVICOR, LOVASTATIN
AZMACORT, TRIAMCINOLONE ACETONIDE
NIASPAN, NIACIN
TEVETEN HCT, EPROSARTAN MESYLATE
TEVETEN, EPROSARTAN MESYLATE

KREMERS URBAN

- * KREMERS URBAN CO
ISOSORBIDE MONONITRATE, ISOSORBIDE MONONITRATE
NITROGLYCERIN, NITROGLYCERIN

KREMERS URBAN DEV

- * KREMERS URBAN DEVELOPMENT CO
OMEPRAZOLE, OMEPRAZOLE

KRKA DD NOVO MESTO

- * KRKA DD NOVO MESTO
ENALAPRIL MALEATE, ENALAPRIL MALEATE

KV PHARM

- * KV PHARMACEUTICAL CO
BENAZEPRIL HYDROCHLORIDE, BENAZEPRIL HYDROCHLORIDE
BROMPHENIRAMINE MALEATE, BROMPHENIRAMINE MALEATE
BUSPIRONE HYDROCHLORIDE, BUSPIRONE HYDROCHLORIDE
CARBIDOPA AND LEVODOPA, CARBIDOPA
CLINDESSE, CLINDAMYCIN PHOSPHATE
DEXTROAMPHETAMINE SULFATE, DEXTROAMPHETAMINE SULFATE
DILTIAZEM HYDROCHLORIDE, DILTIAZEM HYDROCHLORIDE
DISOPYRAMIDE PHOSPHATE, DISOPYRAMIDE PHOSPHATE
DOXAZOSIN MESYLATE, DOXAZOSIN MESYLATE
FLUTICASONE PROPIONATE, FLUTICASONE PROPIONATE
GYNAZOLE-1, BUTOCONAZOLE NITRATE
HYDROCODONE BITARTRATE AND ACETAMINOPHEN, ACETAMINOPHEN
HYDROMORPHONE HYDROCHLORIDE, HYDROMORPHONE HYDROCHLORIDE
ISOSORBIDE MONONITRATE, ISOSORBIDE MONONITRATE
MICRO-K 10, POTASSIUM CHLORIDE
MICRO-K, POTASSIUM CHLORIDE
MORPHINE SULFATE, MORPHINE SULFATE
NYSTATIN, NYSTATIN

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

**** K ****

* KV PHARMACEUTICAL CO
OXYCODONE HYDROCHLORIDE, OXYCODONE HYDROCHLORIDE
POTASSIUM CHLORIDE, POTASSIUM CHLORIDE
PREDNISOLONE SODIUM PHOSPHATE, PREDNISOLONE SODIUM PHOSPHATE
PREDNISOLONE, PREDNISOLONE
PROPAFENONE HYDROCHLORIDE, PROPAFENONE HYDROCHLORIDE

KVK-TECH INC

* KVK-TECH, INC.
PROMETHAZINE HYDROCHLORIDE, PROMETHAZINE HYDROCHLORIDE

LABS ATRAL

* LABORATORIOS ATRAL SARL
TETRACYCLINE HYDROCHLORIDE, TETRACYCLINE HYDROCHLORIDE

LAFAYETTE PHARMS

* LAFAYETTE PHARMACEUTICALS INC
BAROS, SODIUM BICARBONATE

LANNETT

* LANNETT CO INC
ACETAZOLAMIDE, ACETAZOLAMIDE
DICYCLOMINE HYDROCHLORIDE, DICYCLOMINE HYDROCHLORIDE
DIPHENOXYLATE HYDROCHLORIDE AND ATROPINE SULFATE, ATROPINE SULFATE
LANIAZID, ISONIAZID
LANORINAL, ASPIRIN
METHOCARBAMOL, METHOCARBAMOL
PHENTERMINE HYDROCHLORIDE, PHENTERMINE HYDROCHLORIDE
PREDNISOLONE, PREDNISOLONE
PRIMIDONE, PRIMIDONE
PROBALAN, PROBENECID
SERPALAN, RESERPINE
SODIUM P.A.S., AMINOSALICYLATE SODIUM
* LANNETT HOLDINGS INC
BACLOFEN, BACLOFEN
CLINDAMYCIN HYDROCHLORIDE, CLINDAMYCIN HYDROCHLORIDE
DANAZOL, DANAZOL
DOXYCYCLINE HYCLATE, DOXYCYCLINE HYCLATE
DOXYCYCLINE, DOXYCYCLINE
PILOCARPINE HYDROCHLORIDE, PILOCARPINE HYDROCHLORIDE
TERBUTALINE SULFATE, TERBUTALINE SULFATE

LAVIPHARM LABS

* LAVIPHARM LABORATORIES INC
FENTANYL, FENTANYL

LEINER

* LEINER HEALTH PRODUCTS INC
CIMETIDINE, CIMETIDINE (OTC)
FOLIC ACID, FOLIC ACID
HYDROCHLOROTHIAZIDE, HYDROCHLOROTHIAZIDE
IBUPROFEN AND PSEUDOEPHEDRINE HYDROCHLORIDE, IBUPROFEN (OTC)
IBUPROFEN, IBUPROFEN
IBUPROFEN, IBUPROFEN (OTC)
LOPERAMIDE HYDROCHLORIDE, LOPERAMIDE HYDROCHLORIDE (OTC)
NAPROXEN SODIUM, NAPROXEN SODIUM (OTC)
PREDNISONE, PREDNISONE
PROFEN, IBUPROFEN (OTC)
RANITIDINE, RANITIDINE HYDROCHLORIDE (OTC)

LEITNER PHARMS

* LEITNER PHARMACEUTICALS LLC
EQUAGESIC, ASPIRIN
SYNALGOS-DC, ASPIRIN
WYGESIC, ACETAMINOPHEN

LEK PHARMS

* LEK PHARMACEUTICALS D D
AMOXICILLIN AND CLAVULANATE POTASSIUM, AMOXICILLIN

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

** L **

* LEK PHARMACEUTICALS D D
BROMOCRIPTINE MESYLATE, BROMOCRIPTINE MESYLATE
CIMETIDINE, CIMETIDINE
CIMETIDINE, CIMETIDINE (OTC)
ENALAPRIL MALEATE, ENALAPRIL MALEATE
LISINOPRIL, LISINOPRIL
OMEPRazole, OMEPRazole

LEK PHARMS DD

* LEK PHARMACEUTICALS DD
PRAVASTATIN SODIUM, PRAVASTATIN SODIUM

LEO PHARM

* LEO PHARMACEUTICAL PRODUCTS LTD
DOVONEX, CALCIPOTRIENE

LEO PHARM PRODS

* LEO PHARMACEUTICAL PRODUCTS LTD
TACLONEX, BETAMETHASONE DIPROPIONATE

LIGAND

* LIGAND PHARMACEUTICALS INC
AVINZA, MORPHINE SULFATE

LILLY

* ELI LILLY AND CO
ALIMTA, PEMETREXED DISODIUM
CAPASTAT SULFATE, CAPREOMYCIN SULFATE
CYMBALTA, DULOXETINE HYDROCHLORIDE
EVISTA, RALOXIFENE HYDROCHLORIDE
FORTEO, TERIPARATIDE RECOMBINANT HUMAN
GEMZAR, GEMCITABINE HYDROCHLORIDE
GLUCAGON, GLUCAGON RECOMBINANT
HUMALOG MIX 50/50, INSULIN LISPRO PROTAMINE RECOMBINANT
HUMALOG MIX 75/25, INSULIN LISPRO PROTAMINE RECOMBINANT
HUMALOG PEN, INSULIN LISPRO RECOMBINANT
HUMALOG, INSULIN LISPRO RECOMBINANT
HUMATROPE, SOMATROPIN RECOMBINANT
HUMULIN 50/50, INSULIN RECOMBINANT HUMAN (OTC)
HUMULIN 70/30 PEN, INSULIN RECOMBINANT HUMAN (OTC)
HUMULIN 70/30, INSULIN RECOMBINANT HUMAN (OTC)
HUMULIN L, INSULIN ZINC SUSP RECOMBINANT HUMAN (OTC)
HUMULIN N, INSULIN SUSP ISOPHANE RECOMBINANT HUMAN (OTC)
HUMULIN R PEN, INSULIN RECOMBINANT HUMAN (OTC)
HUMULIN R, INSULIN RECOMBINANT HUMAN
HUMULIN R, INSULIN RECOMBINANT HUMAN (OTC)
PROZAC WEEKLY, FLUOXETINE HYDROCHLORIDE
PROZAC, FLUOXETINE HYDROCHLORIDE
QUINIDINE GLUCONATE, QUINIDINE GLUCONATE
SEROMYCIN, CYCLOSERINE
STRATTERA, ATOMOXETINE HYDROCHLORIDE
SYMBYAX, FLUOXETINE HYDROCHLORIDE
ZYPREXA ZYDIS, OLANZAPINE
ZYPREXA, OLANZAPINE
* LILLY RESEARCH LABORATORIES DIV ELI LILLY AND CO
PROZAC, FLUOXETINE HYDROCHLORIDE
SARAFEM, FLUOXETINE HYDROCHLORIDE

LILLY ICOS

* LILLY ICOS LLC
CIALIS, TADALAFIL

LLOYD

* LLOYD INC
LEVOTHROID, LEVOTHYROXINE SODIUM

LNK

* LNK INTERNATIONAL INC
DIPHENHYDRAMINE HYDROCHLORIDE, DIPHENHYDRAMINE HYDROCHLORIDE

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

**** L ****

* LNK INTERNATIONAL INC
DOXYLAMINE SUCCINATE, DOXYLAMINE SUCCINATE (OTC)
IBUPROFEN, IBUPROFEN (OTC)
LOPERAMIDE HYDROCHLORIDE, LOPERAMIDE HYDROCHLORIDE (OTC)
METRONIDAZOLE, METRONIDAZOLE

LOCH

* LOCH PHARMACEUTICALS INC
FOLIC ACID, FOLIC ACID

LOREAL USA

* LOREAL USA PRODUCTS INC
ANTHELIOS 20, AVOBENZONE (OTC)
ANTHELIOS SX, AVOBENZONE (OTC)
CAPITAL SOLEIL 15, AVOBENZONE (OTC)

LUITPOLD

* LUITPOLD PHARMACEUTICALS INC
ACETYLCYSTEINE, ACETYLCYSTEINE
AMINOCAPROIC ACID, AMINOCAPROIC ACID
AMINOPHYLLINE, AMINOPHYLLINE
BRETILUM TOSYLATE, BRETILUM TOSYLATE
CALCITRIOL, CALCITRIOL
CYANOCOBALAMIN, CYANOCOBALAMIN
DEXAMETHASONE SODIUM PHOSPHATE, DEXAMETHASONE SODIUM PHOSPHATE
DEXFERRUM, IRON DEXTRAN
DOBUTAMINE HYDROCHLORIDE, DOBUTAMINE HYDROCHLORIDE
DOPAMINE HYDROCHLORIDE, DOPAMINE HYDROCHLORIDE
DROPERIDOL, DROPERIDOL
FUROSEMIDE, FUROSEMIDE
GLYCOPYRROLATE, GLYCOPYRROLATE
HYDRALAZINE HYDROCHLORIDE, HYDRALAZINE HYDROCHLORIDE
HYDROXYZINE HYDROCHLORIDE, HYDROXYZINE HYDROCHLORIDE
LEUCOVORIN CALCIUM PRESERVATIVE FREE, LEUCOVORIN CALCIUM
LEVOCARNITINE, LEVOCARNITINE
LIDOCAINE HYDROCHLORIDE, LIDOCAINE HYDROCHLORIDE
MANNITOL 25%, MANNITOL
METHYLDOPATE HYDROCHLORIDE, METHYLDOPATE HYDROCHLORIDE
NITROGLYCERIN, NITROGLYCERIN
POTASSIUM CHLORIDE, POTASSIUM CHLORIDE
VENOFER, IRON SUCROSE
VERAPAMIL HYDROCHLORIDE, VERAPAMIL HYDROCHLORIDE

LUPIN

* LUPIN LTD
CEFDINIR, CEFDINIR
CEFOTAXIME SODIUM, CEFOTAXIME SODIUM
CEFPROZIL, CEFPROZIL
CEFTRIAZONE, CEFTRIAZONE SODIUM
CEFUROXIME AXETIL, CEFUROXIME AXETIL
CEPHALEXIN, CEPHALEXIN
LISINAPRIL AND HYDROCHLOROTHIAZIDE, HYDROCHLOROTHIAZIDE
LISINAPRIL, LISINAPRIL
QUINAPRIL, QUINAPRIL HYDROCHLORIDE
SUPRAX, CEFIXIME

LUPIN PHARMS

* LUPIN PHARMACEUTICALS INC
MELOXICAM, MELOXICAM

LYNE

* LYNE LABORATORIES INC
CALCITRIOL, CALCITRIOL
CYPROHEPTADINE HYDROCHLORIDE, CYPROHEPTADINE HYDROCHLORIDE
LEVOCARNITINE, LEVOCARNITINE

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

**** M ****

MALLINCKRODT

- * MALLINCKRODT CHEMICAL INC
 - ANEXSIA 7.5/650, ACETAMINOPHEN
 - ANEXSIA, ACETAMINOPHEN
 - BONTRIL, PHENDIMETRAZINE TARTRATE
 - BUCET, ACETAMINOPHEN
 - BUTALBITAL, ACETAMINOPHEN AND CAFFEINE, ACETAMINOPHEN
 - HYDROCET, ACETAMINOPHEN
 - HYDROCODONE BITARTRATE AND ACETAMINOPHEN, ACETAMINOPHEN
 - LORCET-HD, ACETAMINOPHEN
 - LORTAB, ACETAMINOPHEN
 - MEPERIDINE HYDROCHLORIDE PRESERVATIVE FREE, MEPERIDINE HYDROCHLORIDE
 - METHADONE HYDROCHLORIDE, METHADONE HYDROCHLORIDE
 - METHADOSE, METHADONE HYDROCHLORIDE
 - MORPHINE SULFATE, MORPHINE SULFATE
 - OXYCODONE AND ACETAMINOPHEN, ACETAMINOPHEN
 - TENCON, ACETAMINOPHEN
- * MALLINCKRODT INC
 - ACETAMINOPHEN AND CODEINE PHOSPHATE, ACETAMINOPHEN
 - ANEXSIA 5/325, ACETAMINOPHEN
 - ANEXSIA 7.5/325, ACETAMINOPHEN
 - ANEXSIA, ACETAMINOPHEN
 - DEXTROAMP SACCHARATE, AMP ASPARTATE, DEXTROAMP SULFATE AND AMP SULFATE, AMPHETAMINE ASPARTATE
 - DEXTROAMPHETAMINE SULFATE, DEXTROAMPHETAMINE SULFATE
 - FLUOXETINE, FLUOXETINE HYDROCHLORIDE
 - HYDROCODONE BITARTRATE AND ACETAMINOPHEN, ACETAMINOPHEN
 - HYDROMORPHONE HYDROCHLORIDE, HYDROMORPHONE HYDROCHLORIDE
 - MEPERIDINE HYDROCHLORIDE, MEPERIDINE HYDROCHLORIDE
 - METHADONE HYDROCHLORIDE, METHADONE HYDROCHLORIDE
 - METHYLIN ER, METHYLPHENIDATE HYDROCHLORIDE
 - METHYLIN, METHYLPHENIDATE HYDROCHLORIDE
 - METHYLPHENIDATE HYDROCHLORIDE, METHYLPHENIDATE HYDROCHLORIDE
 - MORPHINE SULFATE, MORPHINE SULFATE
 - NALTREXONE HYDROCHLORIDE, NALTREXONE HYDROCHLORIDE
 - OPTIMARK IN PLASTIC CONTAINER, GADOVERSETAMIDE
 - OPTIMARK, GADOVERSETAMIDE
 - OXYCET, ACETAMINOPHEN
 - OXYCODONE AND ACETAMINOPHEN, ACETAMINOPHEN
 - OXYCODONE HYDROCHLORIDE, OXYCODONE HYDROCHLORIDE
 - PROPOXYPHENE NAPSYLATE AND ACETAMINOPHEN, ACETAMINOPHEN
 - SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER, SODIUM CHLORIDE
 - TRAMADOL HYDROCHLORIDE, TRAMADOL HYDROCHLORIDE
- * MALLINCKRODT MEDICAL INC
 - CONRAY 30, IOTHALAMATE MEGLUMINE
 - CONRAY 400, IOTHALAMATE SODIUM
 - CONRAY 43, IOTHALAMATE MEGLUMINE
 - CONRAY, IOTHALAMATE MEGLUMINE
 - CYSTO-CONRAY II, IOTHALAMATE MEGLUMINE
 - GALLIUM CITRATE GA 67, GALLIUM CITRATE, GA-67
 - HEXABRIX, IOXAGLATE MEGLUMINE
 - INDIUM IN 111 CHLORIDE, INDIUM IN-111 CHLORIDE
 - MD-76R, DIATRIZOATE MEGLUMINE
 - MD-GASTROVIEW, DIATRIZOATE MEGLUMINE
 - OCTREOSCAN, INDIUM IN-111 PENTETREOTIDE KIT
 - OPTIRAY 160, IOVERSOL
 - OPTIRAY 240, IOVERSOL
 - OPTIRAY 300, IOVERSOL
 - OPTIRAY 320, IOVERSOL
 - OPTIRAY 350, IOVERSOL
 - PHOSPHOCOL P32, CHROMIC PHOSPHATE, P-32
 - SODIUM CHROMATE CR 51, SODIUM CHROMATE, CR-51
 - SODIUM IODIDE I 123, SODIUM IODIDE, I-123
 - SODIUM IODIDE I 131, SODIUM IODIDE, I-131

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

** M **

* MALLINCKRODT MEDICAL INC
SODIUM PHOSPHATE P 32, SODIUM PHOSPHATE, P-32
TECHNECOLL, TECHNETIUM TC-99M SULFUR COLLOID KIT
TECHNESCAN MAA, TECHNETIUM TC-99M ALBUMIN AGGREGATED KIT
TECHNESCAN MAG3, TECHNETIUM TC-99M MERTIATIDE KIT
TECHNESCAN PYP KIT, TECHNETIUM TC-99M PYROPHOSPHATE KIT
TECHNESCAN, TECHNETIUM TC-99M OXIDRONATE KIT
THALLOUS CHLORIDE TL 201, THALLOUS CHLORIDE, TL-201
ULTRATAG, TECHNETIUM TC-99M RED BLOOD CELL KIT
ULTRA-TECHNEKOW FM, TECHNETIUM TC-99M SODIUM PERTECHNETATE GENERATOR
XENON XE 133, XENON, XE-133

MARSAM PHARMS LLC

* MARSAM PHARMACEUTICALS LLC
ATRACURIUM BESYLATE PRESERVATIVE FREE, ATRACURIUM BESYLATE
ATRACURIUM BESYLATE, ATRACURIUM BESYLATE
CEFACLOL, CEFACLOL
CEFUROXIME SODIUM, CEFUROXIME SODIUM
DIAZEPAM, DIAZEPAM
DOBUTAMINE HYDROCHLORIDE, DOBUTAMINE HYDROCHLORIDE
ETOPOSIDE, ETOPOSIDE
HEPARIN SODIUM, HEPARIN SODIUM
ISOFLURANE, ISOFLURANE
OXACILLIN SODIUM, OXACILLIN SODIUM
TOBRAMYCIN SULFATE, TOBRAMYCIN SULFATE

MARSHALL PHARMA

* MARSHALL PHARMACAL CORP
SODIUM BUTABARBITAL, BUTABARBITAL SODIUM

MARTEC USA LLC

* MARTEC USA LLC
ATENOLOL AND CHLORTHALIDONE, ATENOLOL
ATENOLOL, ATENOLOL
CIPROFLOXACIN, CIPROFLOXACIN HYDROCHLORIDE
DICLOFENAC SODIUM, DICLOFENAC SODIUM
NIFEDIPINE, NIFEDIPINE
PINDOLOL, PINDOLOL

MATRIX MEDCL

* MATRIX MEDICAL CORP
STERI-STAT, CHLORHEXIDINE GLUCONATE (OTC)

MAYNE PHARMA INTL

* MAYNE PHARMA INTERNATIONAL FAULDING PHARM
DORYX, DOXYCYCLINE HYCLATE

MAYNE PHARMA USA

* MAYNE PHARMA USA INC
ACETYLCYSTEINE, ACETYLCYSTEINE
ACYCLOVIR SODIUM, ACYCLOVIR SODIUM
AMIODARONE HYDROCHLORIDE, AMIODARONE HYDROCHLORIDE
AQUASOL A, VITAMIN A PALMITATE
ATRACURIUM BESYLATE PRESERVATIVE FREE, ATRACURIUM BESYLATE
ATRACURIUM BESYLATE, ATRACURIUM BESYLATE
BLEOMYCIN, BLEOMYCIN SULFATE
BUTORPHANOL TARTRATE PRESERVATIVE FREE, BUTORPHANOL TARTRATE
BUTORPHANOL TARTRATE, BUTORPHANOL TARTRATE
CARBOPLATIN, CARBOPLATIN
CYTARABINE, CYTARABINE
DACARBAZINE, DACARBAZINE
DILTIAZEM HYDROCHLORIDE, DILTIAZEM HYDROCHLORIDE
ENALAPRILAT, ENALAPRILAT
EPIRUBICIN HYDROCHLORIDE, EPIRUBICIN HYDROCHLORIDE
ERYC, ERYTHROMYCIN
FLUCONAZOLE IN SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER, FLUCONAZOLE
FUDR, FLOXURIDINE
HYDROMORPHONE HYDROCHLORIDE, HYDROMORPHONE HYDROCHLORIDE

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

** M **

* MAYNE PHARMA USA INC
LABETALOL HYDROCHLORIDE, LABETALOL HYDROCHLORIDE
LEUCOVORIN CALCIUM, LEUCOVORIN CALCIUM
M.V.I. ADULT (PHARMACY BULK PACKAGE), ASCORBIC ACID
M.V.I. ADULT, ASCORBIC ACID
M.V.I. PEDIATRIC, ASCORBIC ACID
M.V.I.-12 (WITHOUT VITAMIN K), ASCORBIC ACID
METHOTREXATE PRESERVATIVE FREE, METHOTREXATE SODIUM
METHOTREXATE, METHOTREXATE SODIUM
METOCLOPRAMIDE HYDROCHLORIDE, METOCLOPRAMIDE HYDROCHLORIDE
MIDAZOLAM HYDROCHLORIDE, MIDAZOLAM HYDROCHLORIDE
MILRINONE LACTATE, MILRINONE LACTATE
MITOXANTRONE, MITOXANTRONE HYDROCHLORIDE
NALBUPHINE HYDROCHLORIDE, NALBUPHINE HYDROCHLORIDE
NIPENT, PENTOSTATIN
ONDANSETRON HYDROCHLORIDE PRESERVATIVE FREE, ONDANSETRON HYDROCHLORIDE
ONDANSETRON HYDROCHLORIDE, ONDANSETRON HYDROCHLORIDE
PACLITAXEL, PACLITAXEL
PAMIDRONATE DISODIUM, PAMIDRONATE DISODIUM
VINCRISTINE SULFATE PFS, VINCRISTINE SULFATE
VINORELBINE TARTRATE, VINORELBINE TARTRATE

MCNEIL

* MCNEIL CONSUMER PRODUCTS CO DIV MCNEILAB INC
CHILDREN'S MOTRIN, IBUPROFEN (OTC)
IBUPROFEN, IBUPROFEN (OTC)
IMODIUM A-D, LOPERAMIDE HYDROCHLORIDE (OTC)
IMODIUM ADVANCED, LOPERAMIDE HYDROCHLORIDE (OTC)
MEDIPREN, IBUPROFEN (OTC)
MOTRIN IB, IBUPROFEN (OTC)
MOTRIN MIGRAINE PAIN, IBUPROFEN (OTC)

MCNEIL CONS

* MCNEIL CONSUMER HEALTHCARE
CHILDREN'S MOTRIN COLD, IBUPROFEN (OTC)
CHILDREN'S MOTRIN, IBUPROFEN (OTC)
IMODIUM A-D, LOPERAMIDE HYDROCHLORIDE (OTC)
IMODIUM ADVANCED, LOPERAMIDE HYDROCHLORIDE (OTC)
JUNIOR STRENGTH MOTRIN, IBUPROFEN (OTC)
NIZORAL A-D, KETOCONAZOLE (OTC)
SINE-AID IB, IBUPROFEN (OTC)
TYLENOL (CAPLET), ACETAMINOPHEN (OTC)
TYLENOL (GELTAB), ACETAMINOPHEN (OTC)
ZANTAC 150, RANITIDINE HYDROCHLORIDE (OTC)
ZANTAC 75, RANITIDINE HYDROCHLORIDE (OTC)

MCNEIL PED

* MCNEIL PEDIATRICS
FLEXERIL, CYCLOBENZAPRINE HYDROCHLORIDE
IMODIUM, LOPERAMIDE HYDROCHLORIDE
MOTRIN, IBUPROFEN
NIZORAL, KETOCONAZOLE

MEAD JOHNSON

* MEAD JOHNSON AND CO
CAFCIT, CAFFEINE CITRATE

MEDI FLEX INC

* MEDI FLEX HOSP PRODUCTS INC
CHLORAPREP ONE-STEP FREPP, CHLORHEXIDINE GLUCONATE (OTC)
CHLORAPREP ONE-STEP SEPP, CHLORHEXIDINE GLUCONATE (OTC)
CHLORAPREP ONE-STEP, CHLORHEXIDINE GLUCONATE (OTC)
CHLORAPREP SINGLE SWABSTICK, CHLORHEXIDINE GLUCONATE (OTC)
CHLORAPREP WITH TINT, CHLORHEXIDINE GLUCONATE (OTC)

MEDICINES CO

* THE MEDICINES CO
ANGIOMAX, BIVALIRUDIN

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

** M **

MEDICIS

* MEDICIS PHARMACEUTICAL CORP
BUPHENYL, SODIUM PHENYLBUTYRATE
DYNACIN, MINOCYCLINE HYDROCHLORIDE
LIDEX, FLUOCINONIDE
LIDEX-E, FLUOCINONIDE
LOPROX, CICLOPIROX
MINOCYCLINE HYDROCHLORIDE, MINOCYCLINE HYDROCHLORIDE
ORAPRED ODT, PREDNISOLONE SODIUM PHOSPHATE
SOLODYN, MINOCYCLINE HYDROCHLORIDE
SYNACORT, HYDROCORTISONE
SYNALAR, FLUOCINOLONE ACETONIDE
VANOS, FLUOCINONIDE
ZIANA, CLINDAMYCIN PHOSPHATE

MEDICURE

* MEDICURE INTERNATIONAL INC
AGGRASTAT, TIROFIBAN HYDROCHLORIDE

MEDIGENE

* MEDIGENE INC
VEREGEN, KUNECATECHINS

MEDIMMUNE ONCOLOGY

* MEDIMMUNE ONCOLOGY INC
ETHYOL, AMIFOSTINE
NEUTREXIN, TRIMETREXATE GLUCURONATE

MEDPOINTE

* MEDPOINTE HEALTHCARE INC
FELBATOL, FELBAMATE

MEDPOINTE PHARM HLC

* MEDPOINTE PHARMACEUTICALS MEDPOINTE HEALTHCARE INC
ASTELIN, AZELASTINE HYDROCHLORIDE
BUTISOL SODIUM, BUTABARBITAL SODIUM
DEPEN, PENICILLAMINE
LUFYLLIN, DYPHYLLINE
OPTIVAR, AZELASTINE HYDROCHLORIDE
SOMA COMPOUND W/ CODEINE, ASPIRIN
SOMA COMPOUND, ASPIRIN
SOMA, CARISOPRODOL
VOSOL HC, ACETIC ACID, GLACIAL

MEDTRONIC

* MEDTRONIC INC
LIORESAL, BACLOFEN

MEPHA

* MEPHA AG
PIROXICAM, PIROXICAM

MERCK

* MERCK AND CO INC
AMINOHIPPURATE SODIUM, AMINOHIPPURATE SODIUM
CANCIDAS, CASPOFUNGIN ACETATE
DIURIL, CHLOROTHIAZIDE
EMEND, APREPITANT
FOSAMAX PLUS D, ALENDRONATE SODIUM
FOSAMAX, ALENDRONATE SODIUM
HYZAAR, HYDROCHLOROTHIAZIDE
INVANZ, ERTAPENEM SODIUM
MAXALT, RIZATRIPTAN BENZOATE
MAXALT-MLT, RIZATRIPTAN BENZOATE
MEFOXIN, CEFOXITIN SODIUM
MINTEZOL, THIABENDAZOLE
PRIMAXIN, CILASTATIN SODIUM
PROSCAR, FINASTERIDE

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

** M **

- * MERCK AND CO INC
 - SINGULAIR, MONTELUKAST SODIUM
 - STROMEKTOL, IVERMECTIN
 - ZOLINZA, VORINOSTAT
- * MERCK RESEARCH LABORATORIES DIV MERCK CO INC
 - CLINORIL, SULINDAC
 - COSOPT, DORZOLAMIDE HYDROCHLORIDE
 - COZAAR, LOSARTAN POTASSIUM
 - CRIXIVAN, INDINAVIR SULFATE
 - INDOCIN, INDOMETHACIN
 - MEFOXIN IN PLASTIC CONTAINER, CEFOXITIN SODIUM
 - MEVACOR, LOVASTATIN
 - NOROXIN, NORFLOXACIN
 - PEPCID AC (GELTAB), FAMOTIDINE (OTC)
 - PEPCID AC, FAMOTIDINE (OTC)
 - PEPCID COMPLETE, CALCIUM CARBONATE (OTC)
 - PEPCID PRESERVATIVE FREE IN PLASTIC CONTAINER, FAMOTIDINE
 - PEPCID PRESERVATIVE FREE, FAMOTIDINE
 - PEPCID, FAMOTIDINE
 - PRINIVIL, LISINOPRIL
 - PRINZIDE, HYDROCHLOROTHIAZIDE
 - PROPECIA, FINASTERIDE
 - SINGULAIR, MONTELUKAST SODIUM
 - TIMOPTIC IN OCUDOSE, TIMOLOL MALEATE
 - TIMOPTIC, TIMOLOL MALEATE
 - TIMOPTIC-XE, TIMOLOL MALEATE
 - TRUSOPT, DORZOLAMIDE HYDROCHLORIDE
 - ZOCOR, SIMVASTATIN

MERCK CO INC

- * MERCK CO INC
 - JANUVIA, SITAGLIPTIN PHOSPHATE

MERETEK

- * MERETEK DIAGNOSTICS INC
 - BREATHTEK UBT FOR H-PYLORI, UREA, C-13

MERIDIAN MEDCL

- * MERIDIAN MEDICAL TECHNOLOGIES INC
 - DUODOTE, ATROPINE

MERIDIAN MEDCL TECHN

- * MERIDIAN MEDICAL TECHNOLOGIES INC
 - ATROPEN, ATROPINE
 - EPIPEN JR., EPINEPHRINE
 - EPIPEN, EPINEPHRINE
 - LIDOPEN, LIDOCAINE HYDROCHLORIDE
 - MORPHINE SULFATE, MORPHINE SULFATE
 - PRALIDOXIME CHLORIDE, PRALIDOXIME CHLORIDE

MERZ PHARMS

- * MERZ PHARMACEUTICALS LLC
 - ERYGEL, ERYTHROMYCIN
 - NAFTIN, NAFTIFINE HYDROCHLORIDE

METHAPHARM

- * METHAPHARM INC
 - PROVOCHOLINE, METHACHOLINE CHLORIDE

METRICS PHARM

- * METRICS PHARMACEUTICAL VENTURES LLC
 - NOREPINEPHRINE BITARTRATE, NOREPINEPHRINE BITARTRATE

MGI PHARMA INC

- * MGI PHARMA INC
 - DACOGEN, DECITABINE
 - GLIADEL, CARMUSTINE
 - HEXALEN, ALTRETAMINE
 - SALAGEN, PILOCARPINE HYDROCHLORIDE

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

**** M ****

MIKART

* MIKART INC
ACETAMINOPHEN AND CODEINE PHOSPHATE, ACETAMINOPHEN
ACETAMINOPHEN, ASPIRIN, AND CODEINE PHOSPHATE, ACETAMINOPHEN
ACETAMINOPHEN, BUTALBITAL, AND CAFFEINE, ACETAMINOPHEN
ACETAMINOPHEN, CAFFEINE, AND DIHYDROCODEINE BITARTRATE, ACETAMINOPHEN
AMANTADINE HYDROCHLORIDE, AMANTADINE HYDROCHLORIDE
AMINOCAPROIC ACID, AMINOCAPROIC ACID
AMINOCAPROIC, AMINOCAPROIC ACID
BUTALBITAL, ACETAMINOPHEN AND CAFFEINE, ACETAMINOPHEN
BUTAPAP, ACETAMINOPHEN
CARBINOXAMINE MALEATE, CARBINOXAMINE MALEATE
DICYCLOMINE HYDROCHLORIDE, DICYCLOMINE HYDROCHLORIDE
ERGOTAMINE TARTRATE AND CAFFEINE, CAFFEINE
ESGIC-PLUS, ACETAMINOPHEN
ETHOSUXIMIDE, ETHOSUXIMIDE
HYDROCODONE BITARTRATE AND ACETAMINOPHEN, ACETAMINOPHEN
ISONIAZID, ISONIAZID
METHAZOLAMIDE, METHAZOLAMIDE
OXYBUTYNIN CHLORIDE, OXYBUTYNIN CHLORIDE
OXYCODONE AND ACETAMINOPHEN, ACETAMINOPHEN
PHENDIMETRAZINE TARTRATE, PHENDIMETRAZINE TARTRATE
PYRAZINAMIDE, PYRAZINAMIDE
TRIHXYPHENIDYL HYDROCHLORIDE, TRIHXYPHENIDYL HYDROCHLORIDE

MILEX

* MILEX PRODUCTS INC
MILOPHENE, CLOMIPHENE CITRATE

MILLENNIUM PHARMS

* MILLENNIUM PHARMACEUTICALS INC
VELCADE, BORTEZOMIB

MINRAD

* MINRAD INC
ENFLURANE, ENFLURANE
ISOFLURANE, ISOFLURANE

MISSION PHARMA

* MISSION PHARMACAL CO
FOLICET, FOLIC ACID
LITHOSTAT, ACETOHYDROXAMIC ACID
TINDAMAX, TINIDAZOLE
TIOPRONIN, TIOPRONIN
UROKIT-K, POTASSIUM CITRATE

MIZA PHARMS USA

* MIZA PHARMACEUTICALS USA INC
OCUSULF-10, SULFACETAMIDE SODIUM
TROPICAMIDE, TROPICAMIDE

MONARCH PHARMS

* MONARCH PHARMACEUTICALS INC
CORTISPORIN, BACITRACIN ZINC
CORTISPORIN, HYDROCORTISONE
CORTISPORIN, HYDROCORTISONE ACETATE
KEMADRIN, PROCYCLIDINE HYDROCHLORIDE
MENEST, ESTROGENS, ESTERIFIED
NEOSPORIN G.U. IRRIGANT, NEOMYCIN SULFATE
NEOSPORIN, BACITRACIN ZINC
NEOSPORIN, GRAMICIDIN
PEDIOTIC, HYDROCORTISONE
POLYSPORIN, BACITRACIN ZINC
PROLOPRIM, TRIMETHOPRIM
QUIBRON-T, THEOPHYLLINE
QUIBRON-T/SR, THEOPHYLLINE
SEPTRA DS, SULFAMETHOXAZOLE

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

** M **

* MONARCH PHARMACEUTICALS INC
 SEPTRA GRAPE, SULFAMETHOXAZOLE
 SEPTRA, SULFAMETHOXAZOLE
 THALITONE, CHLORTHALIDONE
 VIROPTIC, TRIFLURIDINE

MORTON GROVE

* MORTON GROVE ACQUISITION CORP
 CLEMASTINE FUMARATE, CLEMASTINE FUMARATE
 FLUOXETINE HYDROCHLORIDE, FLUOXETINE HYDROCHLORIDE
 MINOXIDIL EXTRA STRENGTH (FOR MEN), MINOXIDIL (OTC)
 OXYBUTYNIN CHLORIDE, OXYBUTYNIN CHLORIDE
 PREDNISOLONE SODIUM PHOSPHATE, PREDNISOLONE SODIUM PHOSPHATE

* MORTON GROVE PHARMACEUTICALS INC
 ACETAMINOPHEN AND CODEINE PHOSPHATE, ACETAMINOPHEN
 ACETIC ACID, ACETIC ACID, GLACIAL
 ALBUTEROL SULFATE, ALBUTEROL SULFATE
 AMANTADINE HYDROCHLORIDE, AMANTADINE HYDROCHLORIDE
 CARBAMAZEPINE, CARBAMAZEPINE
 CHLORHEXIDINE GLUCONATE, CHLORHEXIDINE GLUCONATE
 CIMETIDINE HYDROCHLORIDE, CIMETIDINE HYDROCHLORIDE
 CLINDAMYCIN PHOSPHATE, CLINDAMYCIN PHOSPHATE
 CLOBETASOL PROPIONATE, CLOBETASOL PROPIONATE
 CROMOLYN SODIUM, CROMOLYN SODIUM
 CYCLOSPORINE, CYCLOSPORINE
 DOXEPIN HYDROCHLORIDE, DOXEPIN HYDROCHLORIDE
 ERYTHROMYCIN, ERYTHROMYCIN
 FUROSEMIDE, FUROSEMIDE
 GENERLAC, LACTULOSE
 HYDROCORTISONE AND ACETIC ACID, ACETIC ACID, GLACIAL
 HYDROXYZINE HYDROCHLORIDE, HYDROXYZINE HYDROCHLORIDE
 LACTULOSE, LACTULOSE
 LIDOCAINE HYDROCHLORIDE, LIDOCAINE HYDROCHLORIDE
 LINDANE, LINDANE
 LITHIUM CITRATE, LITHIUM CITRATE
 LOPERAMIDE HYDROCHLORIDE, LOPERAMIDE HYDROCHLORIDE (OTC)
 LORATADINE, LORATADINE (OTC)
 MEGESTROL ACETATE, MEGESTROL ACETATE
 METAPROTERENOL SULFATE, METAPROTERENOL SULFATE
 MINOXIDIL (FOR MEN), MINOXIDIL (OTC)
 MYBANIL, BROMODIPHENHYDRAMINE HYDROCHLORIDE
 MYCODONE, HOMATROPINE METHYLBROMIDE
 MYLAMINE, DEXCHLORPHENIRAMINE MALEATE
 MYMETHASONE, DEXAMETHASONE
 MYPHETANE DX, BROMPHENIRAMINE MALEATE
 MYPROIC ACID, VALPROIC ACID
 NYSTATIN, NYSTATIN
 PHENTYTOIN, PHENTYTOIN
 PREDNISOLONE SODIUM PHOSPHATE, PREDNISOLONE SODIUM PHOSPHATE
 PREDNISOLONE, PREDNISOLONE
 PROMETHAZINE PLAIN, PROMETHAZINE HYDROCHLORIDE
 PROMETHAZINE W/ CODEINE, CODEINE PHOSPHATE
 PROMETHAZINE W/ DEXTROMETHORPHAN, DEXTROMETHORPHAN HYDROBROMIDE
 SELENIUM SULFIDE, SELENIUM SULFIDE
 THEOPHYLLINE, THEOPHYLLINE
 TRETINOIN, TRETINOIN
 TRIAMCINOLONE ACETONIDE, TRIAMCINOLONE ACETONIDE
 TRIPROLIDINE HCL, PSEUDOEPHEDRINE HCL AND CODEINE PHOSPHATE, CODEINE PHOSPHATE

MSP SINGAPORE

* MSP SINGAPORE CO LLC
 VYTORIN, EZETIMIBE
 ZETIA, EZETIMIBE

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

**** M ****

MUTUAL PHARM

* MUTUAL PHARMACEUTICAL CO INC
ACETAMINOPHEN AND CODEINE PHOSPHATE, ACETAMINOPHEN
ACETAZOLAMIDE, ACETAZOLAMIDE
ALBUTEROL SULFATE, ALBUTEROL SULFATE
ALLOPURINOL, ALLOPURINOL
AMITRIPTYLINE HYDROCHLORIDE, AMITRIPTYLINE HYDROCHLORIDE
ATENOLOL AND CHLORTHALIDONE, ATENOLOL
ATENOLOL, ATENOLOL
BACTRIM DS, SULFAMETHOXAZOLE
BACTRIM PEDIATRIC, SULFAMETHOXAZOLE
BACTRIM, SULFAMETHOXAZOLE
BENZTROPINE MESYLATE, BENZTROPINE MESYLATE
BISOPROLOL FUMARATE, BISOPROLOL FUMARATE
BUTALBITAL, ACETAMINOPHEN AND CAFFEINE, ACETAMINOPHEN
CARISOPRODOL, CARISOPRODOL
CHLORZOXAZONE, CHLORZOXAZONE
CILOSTAZOL, CILOSTAZOL
CITALOPRAM HYDROBROMIDE, CITALOPRAM HYDROBROMIDE
CLONIDINE HYDROCHLORIDE, CLONIDINE HYDROCHLORIDE
CYCLOBENZAPRINE HYDROCHLORIDE, CYCLOBENZAPRINE HYDROCHLORIDE
DEXTROAMP SACCHARATE, AMP ASPARTATE, DEXTROAMP SULFATE AND AMP SULFATE, AMPHETAMINE
ASPARTATE
DICLOFENAC POTASSIUM, DICLOFENAC POTASSIUM
DIPHENHYDRAMINE HYDROCHLORIDE, DIPHENHYDRAMINE HYDROCHLORIDE
DOXYCYCLINE HYCLATE, DOXYCYCLINE HYCLATE
ERGOLOID MESYLATES, ERGOLOID MESYLATES
FAMOTIDINE, FAMOTIDINE
FELODIPINE, FELODIPINE
FENOPROFEN CALCIUM, FENOPROFEN CALCIUM
FLUVOXAMINE MALEATE, FLUVOXAMINE MALEATE
FOLIC ACID, FOLIC ACID
GABAPENTIN, GABAPENTIN
HYDRALAZINE HYDROCHLORIDE, HYDRALAZINE HYDROCHLORIDE
HYDROCODONE BITARTRATE AND ACETAMINOPHEN, ACETAMINOPHEN
HYDROXYZINE HYDROCHLORIDE, HYDROXYZINE HYDROCHLORIDE
IBUPROFEN, IBUPROFEN
IBUPROFEN, IBUPROFEN (OTC)
IMIPRAMINE HYDROCHLORIDE, IMIPRAMINE HYDROCHLORIDE
ISONIAZID, ISONIAZID
LABETALOL HYDROCHLORIDE, LABETALOL HYDROCHLORIDE
LORAZEPAM, LORAZEPAM
LOVASTATIN, LOVASTATIN
MELOXICAM, MELOXICAM
MEPERIDINE HYDROCHLORIDE, MEPERIDINE HYDROCHLORIDE
METFORMIN HYDROCHLORIDE, METFORMIN HYDROCHLORIDE
METOCLOPRAMIDE HYDROCHLORIDE, METOCLOPRAMIDE HYDROCHLORIDE
METOPROLOL TARTRATE, METOPROLOL TARTRATE
METRONIDAZOLE, METRONIDAZOLE
MINOXIDIL, MINOXIDIL
NYSTATIN, NYSTATIN
OXYCODONE AND ACETAMINOPHEN, ACETAMINOPHEN
OXYCODONE AND ASPIRIN, ASPIRIN
PHENTERMINE HYDROCHLORIDE, PHENTERMINE HYDROCHLORIDE
PINDOLOL, PINDOLOL
PIROXICAM, PIROXICAM
PREDNISONE, PREDNISONE
PRIMIDONE, PRIMIDONE
PROPAFENONE HYDROCHLORIDE, PROPAFENONE HYDROCHLORIDE
PROPOXYPHENE NAPSYLATE AND ACETAMINOPHEN, ACETAMINOPHEN
QUINIDINE GLUCONATE, QUINIDINE GLUCONATE
QUINIDINE SULFATE, QUINIDINE SULFATE
SOTALOL HYDROCHLORIDE, SOTALOL HYDROCHLORIDE

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

** M **

* MUTUAL PHARMACEUTICAL CO INC
SPIRONOLACTONE AND HYDROCHLOROTHIAZIDE, HYDROCHLOROTHIAZIDE
SPIRONOLACTONE, SPIRONOLACTONE
SULFAMETHOXAZOLE AND TRIMETHOPRIM, SULFAMETHOXAZOLE
SULFASALAZINE, SULFASALAZINE
SULINDAC, SULINDAC
TETRACYCLINE HYDROCHLORIDE, TETRACYCLINE HYDROCHLORIDE
THIORIDAZINE HYDROCHLORIDE, THIORIDAZINE HYDROCHLORIDE
TOLAZAMIDE, TOLAZAMIDE
TOLMETIN SODIUM, TOLMETIN SODIUM
TRAMADOL HYDROCHLORIDE, TRAMADOL HYDROCHLORIDE
TRAZODONE HYDROCHLORIDE, TRAZODONE HYDROCHLORIDE
VERAPAMIL HYDROCHLORIDE, VERAPAMIL HYDROCHLORIDE
ZONISAMIDE, ZONISAMIDE

MUTUAL PHARMA

* MUTUAL PHARMACAL CO
DOXYCYCLINE HYCLATE, DOXYCYCLINE HYCLATE
FLUOXETINE, FLUOXETINE HYDROCHLORIDE
KETOCONAZOLE, KETOCONAZOLE
METFORMIN HYDROCHLORIDE, METFORMIN HYDROCHLORIDE
TRIMETHOBENZAMIDE HYDROCHLORIDE, TRIMETHOBENZAMIDE HYDROCHLORIDE

MYLAN

* MYLAN LABORATORIES INC
ACYCLOVIR, ACYCLOVIR
AMLODIPINE BESYLATE, AMLDIPINE BESYLATE
ANAGRELIDE HYDROCHLORIDE, ANAGRELIDE HYDROCHLORIDE
CAPTOPRIL, CAPTOPRIL
ETODOLAC, ETODOLAC
IBUPROFEN, IBUPROFEN
TERAZOSIN HYDROCHLORIDE, TERAZOSIN HYDROCHLORIDE

* MYLAN PHARMACEUTICALS INC
ACEBUTOLOL HYDROCHLORIDE, ACEBUTOLOL HYDROCHLORIDE
ACYCLOVIR, ACYCLOVIR
ALBUTEROL SULFATE, ALBUTEROL SULFATE
ALLOPURINOL, ALLOPURINOL
ALPRAZOLAM, ALPRAZOLAM
AMILORIDE HYDROCHLORIDE AND HYDROCHLOROTHIAZIDE, AMILORIDE HYDROCHLORIDE
AMITRIPTYLINE HYDROCHLORIDE, AMITRIPTYLINE HYDROCHLORIDE
ATENOLOL AND CHLORTHALIDONE, ATENOLOL
ATENOLOL, ATENOLOL
AZITHROMYCIN, AZITHROMYCIN
BENAZEPRIL HYDROCHLORIDE AND HYDROCHLOROTHIAZIDE, BENAZEPRIL HYDROCHLORIDE
BENAZEPRIL HYDROCHLORIDE, BENAZEPRIL HYDROCHLORIDE
BISOPROLOL FUMARATE AND HYDROCHLOROTHIAZIDE, BISOPROLOL FUMARATE
BISOPROLOL FUMARATE, BISOPROLOL FUMARATE
BROMOCRIPTINE MESYLATE, BROMOCRIPTINE MESYLATE
BUPROPION HYDROCHLORIDE, BUPROPION HYDROCHLORIDE
BUSPIRONE HYDROCHLORIDE, BUSPIRONE HYDROCHLORIDE
BUTORPHANOL TARTRATE, BUTORPHANOL TARTRATE
CAPTOPRIL AND HYDROCHLOROTHIAZIDE, CAPTOPRIL
CARBIDOPA AND LEVODOPA, CARBIDOPA
CHLORDIAZEPOXIDE AND AMITRIPTYLINE HYDROCHLORIDE, AMITRIPTYLINE HYDROCHLORIDE
CHLOROTHIAZIDE, CHLOROTHIAZIDE
CHLORPROPAMIDE, CHLORPROPAMIDE
CHLORTHALIDONE, CHLORTHALIDONE
CILOSTAZOL, CILOSTAZOL
CIMETIDINE, CIMETIDINE
CIPROFLOXACIN, CIPROFLOXACIN HYDROCHLORIDE
CITALOPRAM HYDROBROMIDE, CITALOPRAM HYDROBROMIDE
CLOMIPRAMINE HYDROCHLORIDE, CLOMIPRAMINE HYDROCHLORIDE
CLONAZEPAM, CLONAZEPAM
CLONIDINE HYDROCHLORIDE, CLONIDINE HYDROCHLORIDE
CLORAZEPATE DIPOTASSIUM, CLORAZEPATE DIPOTASSIUM

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

**** M ****

* MYLAN PHARMACEUTICALS INC
CLORPRES, CHLORTHALIDONE
CLOZAPINE, CLOZAPINE
CYCLOBENZAPRINE HYDROCHLORIDE, CYCLOBENZAPRINE HYDROCHLORIDE
CYSTAGON, CYSTEAMINE BITARTRATE
DIAZEPAM, DIAZEPAM
DICLOFENAC POTASSIUM, DICLOFENAC POTASSIUM
DICLOFENAC SODIUM, DICLOFENAC SODIUM
DICYCLOMINE HYDROCHLORIDE, DICYCLOMINE HYDROCHLORIDE
DILTIAZEM HYDROCHLORIDE, DILTIAZEM HYDROCHLORIDE
DIPHENOXYLATE HYDROCHLORIDE AND ATROPINE SULFATE, ATROPINE SULFATE
DOXAZOSIN MESYLATE, DOXAZOSIN MESYLATE
DOXEPIN HYDROCHLORIDE, DOXEPIN HYDROCHLORIDE
DOXYCYCLINE HYCLATE, DOXYCYCLINE HYCLATE
DOXYCYCLINE, DOXYCYCLINE
ENALAPRIL MALEATE AND HYDROCHLOROTHIAZIDE, ENALAPRIL MALEATE
ENALAPRIL MALEATE, ENALAPRIL MALEATE
ERYTHROMYCIN ETHYLSUCCINATE, ERYTHROMYCIN ETHYLSUCCINATE
ERYTHROMYCIN STEARATE, ERYTHROMYCIN STEARATE
ESTRADIOL, ESTRADIOL
ESTROPIPATE, ESTROPIPATE
ETODOLAC, ETODOLAC
EXTENDED PHENYTOIN SODIUM, PHENYTOIN SODIUM
FAMOTIDINE, FAMOTIDINE
FENOPROFEN CALCIUM, FENOPROFEN CALCIUM
FINASTERIDE, FINASTERIDE
FLUCONAZOLE, FLUCONAZOLE
FLUOXETINE, FLUOXETINE HYDROCHLORIDE
FLUPHENAZINE HYDROCHLORIDE, FLUPHENAZINE HYDROCHLORIDE
FLURAZEPAM HYDROCHLORIDE, FLURAZEPAM HYDROCHLORIDE
FLURBIPROFEN, FLURBIPROFEN
FLUVOXAMINE MALEATE, FLUVOXAMINE MALEATE
FUROSEMIDE, FUROSEMIDE
GEMFIBROZIL, GEMFIBROZIL
GLIMEPIRIDE, GLIMEPIRIDE
GLIPIZIDE, GLIPIZIDE
GLYBURIDE (MICRONIZED), GLYBURIDE
GUANFACINE HYDROCHLORIDE, GUANFACINE HYDROCHLORIDE
HALOPERIDOL, HALOPERIDOL
HYDROCHLOROTHIAZIDE, HYDROCHLOROTHIAZIDE
HYDROXYCHLOROQUINE SULFATE, HYDROXYCHLOROQUINE SULFATE
IBUPROFEN, IBUPROFEN
IBUPROFEN, IBUPROFEN (OTC)
INDAPAMIDE, INDAPAMIDE
INDOMETHACIN, INDOMETHACIN
KETOCONAZOLE, KETOCONAZOLE
KETOPROFEN, KETOPROFEN
KETOROLAC TROMETHAMINE, KETOROLAC TROMETHAMINE
LEVOTHYROXINE SODIUM, LEVOTHYROXINE SODIUM
LISINOPRIL AND HYDROCHLOROTHIAZIDE, HYDROCHLOROTHIAZIDE
LISINOPRIL, LISINOPRIL
LOPERAMIDE HYDROCHLORIDE, LOPERAMIDE HYDROCHLORIDE
LORAZEPAM, LORAZEPAM
LOVASTATIN, LOVASTATIN
LOXAPINE, LOXAPINE SUCCINATE
MAPROTILINE HYDROCHLORIDE, MAPROTILINE HYDROCHLORIDE
MECLOFENAMATE SODIUM, MECLOFENAMATE SODIUM
MELOXICAM, MELOXICAM
MERCAPTOPYRINE, MERCAPTOPYRINE ANHYDROUS
METFORMIN HYDROCHLORIDE, METFORMIN HYDROCHLORIDE
METHOTREXATE SODIUM, METHOTREXATE SODIUM
METHYLCLOTHIAZIDE, METHYLCLOTHIAZIDE
METHYLDOPA AND HYDROCHLOROTHIAZIDE, HYDROCHLOROTHIAZIDE
METHYLDOPA, METHYLDOPA

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

** M **

* MYLAN PHARMACEUTICALS INC
METOLAZONE, METOLAZONE
METOPROLOL TARTRATE AND HYDROCHLOROTHIAZIDE, HYDROCHLOROTHIAZIDE
METOPROLOL TARTRATE, METOPROLOL TARTRATE
MIDODRINE HYDROCHLORIDE, MIDODRINE HYDROCHLORIDE
MIRTAZAPINE, MIRTAZAPINE
NADOLOL, NADOLOL
NAPROXEN SODIUM, NAPROXEN SODIUM
NAPROXEN, NAPROXEN
NEFAZODONE HYDROCHLORIDE, NEFAZODONE HYDROCHLORIDE
NICARDIPINE HYDROCHLORIDE, NICARDIPINE HYDROCHLORIDE
NITROFURANTOIN (MONOHYDRATE/MACROCRYSTALS), NITROFURANTOIN
NITROFURANTOIN, NITROFURANTOIN, MACROCRYSTALLINE
NIZATIDINE, NIZATIDINE
NORTRIPTYLINE HYDROCHLORIDE, NORTRIPTYLINE HYDROCHLORIDE
OMEPRazole, OMEPRazole
OXAPROZIN, OXAPROZIN
OXYBUTYNIN CHLORIDE, OXYBUTYNIN CHLORIDE
PACLITAXEL, PACLITAXEL
PENTOXIFYLLINE, PENTOXIFYLLINE
PERPHENAZINE AND AMITRIPTYLINE HYDROCHLORIDE, AMITRIPTYLINE HYDROCHLORIDE
PHENYTEK, PHENYTOIN SODIUM
PINDOLOL, PINDOLOL
PIROXICAM, PIROXICAM
PRAZOSIN HYDROCHLORIDE, PRAZOSIN HYDROCHLORIDE
PROBENECID, PROBENECID
PROCHLORPERAZINE MALEATE, PROCHLORPERAZINE MALEATE
PROPOXYPHENE HYDROCHLORIDE AND ACETAMINOPHEN, ACETAMINOPHEN
PROPOXYPHENE HYDROCHLORIDE, PROPOXYPHENE HYDROCHLORIDE
PROPOXYPHENE NAPSYLATE AND ACETAMINOPHEN, ACETAMINOPHEN
PROPRANOLOL HYDROCHLORIDE AND HYDROCHLOROTHIAZIDE, HYDROCHLOROTHIAZIDE
PROPRANOLOL HYDROCHLORIDE, PROPRANOLOL HYDROCHLORIDE
QUINAPRIL HYDROCHLORIDE AND HYDROCHLOROTHIAZIDE, HYDROCHLOROTHIAZIDE
QUINAPRIL HYDROCHLORIDE, QUINAPRIL HYDROCHLORIDE
RANITIDINE HYDROCHLORIDE, RANITIDINE HYDROCHLORIDE
SOTALOL HYDROCHLORIDE, SOTALOL HYDROCHLORIDE
SPIRONOLACTONE AND HYDROCHLOROTHIAZIDE, HYDROCHLOROTHIAZIDE
SPIRONOLACTONE, SPIRONOLACTONE
SULINDAC, SULINDAC
TAMOXIFEN CITRATE, TAMOXIFEN CITRATE
TEMAZEPAM, TEMAZEPAM
TETRACYCLINE HYDROCHLORIDE, TETRACYCLINE HYDROCHLORIDE
THIORIDAZINE HYDROCHLORIDE, THIORIDAZINE HYDROCHLORIDE
THIOTHIXENE, THIOTHIXENE
TICLOPIDINE HYDROCHLORIDE, TICLOPIDINE HYDROCHLORIDE
TIMOLOL MALEATE, TIMOLOL MALEATE
TIZANIDINE HYDROCHLORIDE, TIZANIDINE HYDROCHLORIDE
TOLAZAMIDE, TOLAZAMIDE
TOLBUTAMIDE, TOLBUTAMIDE
TOLMETIN SODIUM, TOLMETIN SODIUM
TOPIRAMATE, TOPIRAMATE
TRAMADOL HYDROCHLORIDE, TRAMADOL HYDROCHLORIDE
TRAZODONE HYDROCHLORIDE, TRAZODONE HYDROCHLORIDE
TRIAMTERENE AND HYDROCHLOROTHIAZIDE, HYDROCHLOROTHIAZIDE
TRIFLUOPERAZINE HYDROCHLORIDE, TRIFLUOPERAZINE HYDROCHLORIDE
VERAPAMIL HYDROCHLORIDE, VERAPAMIL HYDROCHLORIDE
ZONISAMIDE, ZONISAMIDE

MYLAN BERTEK

* MYLAN BERTEK PHARMACEUTICALS INC
AVITA, TRETINOIN
MAXZIDE, HYDROCHLOROTHIAZIDE
MAXZIDE-25, HYDROCHLOROTHIAZIDE
MENTAX, BUTENAFINE HYDROCHLORIDE

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

**** M ****

* MYLAN BERTEK PHARMACEUTICALS INC
MENTAX-TC, BUTENAFINE HYDROCHLORIDE
SULFAMYLON, MAFENIDE ACETATE

MYLAN TECHNOLOGIES

* MYLAN TECHNOLOGIES INC
ESTRADIOL, ESTRADIOL
FENTANYL, FENTANYL
NITROGLYCERIN, NITROGLYCERIN
TERAZOSIN HYDROCHLORIDE, TERAZOSIN HYDROCHLORIDE

NABI

* NABI BIOPHARMACEUTICALS
ALOPRIM, ALLOPURINOL SODIUM

NEIL

* NEIL LABORATORIES INC
CARISOPRODOL, CARISOPRODOL
IBUPROFEN, IBUPROFEN (OTC)

NEPHRON

* NEPHRON CORP
ALBUTEROL SULFATE, ALBUTEROL SULFATE
BETA-2, ISOETHARINE HYDROCHLORIDE
IPRATROPIUM BROMIDE, IPRATROPIUM BROMIDE
* NEPHRON PHARMACEUTICALS CORP
ALBUTEROL SULFATE, ALBUTEROL SULFATE
METAPROTERENOL SULFATE, METAPROTERENOL SULFATE

NEW RIVER

* NEW RIVER PHARMACEUTICALS INC
PROFERDEX, IRON DEXTRAN

NEXUS PHARMS

* NEXUS PHARMACEUTICALS INC
CIPROFLOXACIN, CIPROFLOXACIN

NICE PAK

* NICE PAK PRODUCTS INC
CHLORASCRUB MAXI SWABSTICK, CHLORHEXIDINE GLUCONATE (OTC)
CHLORASCRUB SWAB, CHLORHEXIDINE GLUCONATE (OTC)
CHLORASCRUB SWABSTICK, CHLORHEXIDINE GLUCONATE (OTC)

NITROMED

* NITROMED INC
BIDIL, HYDRALAZINE HYDROCHLORIDE

NORTH SHORE LIJ

* NORTH SHORE LIJ RESEARCH INSTITUTE
FLUDEOXYGLUCOSE F18, FLUDEOXYGLUCOSE F-18

NOSTRUM

* NOSTRUM PHARMACEUTICALS INC
METFORMIN HYDROCHLORIDE, METFORMIN HYDROCHLORIDE
THEOPHYLLINE, THEOPHYLLINE

NOVADEL

* NOVADEL PHARMA INC
NITROMIST, NITROGLYCERIN

NOVARTIS

* NOVARTIS CONSUMER HEALTH INC
DENAVIR, PENCICLOVIR SODIUM
EXCEDRIN (MIGRAINE), ACETAMINOPHEN (OTC)
HABITROL, NICOTINE (OTC)
LAMISIL AT, TERBINAFINE (OTC)
LAMISIL AT, TERBINAFINE HYDROCHLORIDE (OTC)
LAMISIL, TERBINAFINE
LAMISIL, TERBINAFINE HYDROCHLORIDE
LAMISIL, TERBINAFINE HYDROCHLORIDE (OTC)
TAVIST ALLERGY/SINUS/HEADACHE, ACETAMINOPHEN (OTC)
TRANSDERM SCOP, SCOPOLAMINE

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

** N **

* NOVARTIS CONSUMER HEALTH INC
VAGISTAT-1, TIOCONAZOLE (OTC)

* NOVARTIS PHARMACEUTICALS CORP
ANTURANE, SULFINPYRAZONE
APRESAZIDE, HYDRALAZINE HYDROCHLORIDE
AREDIA, PAMIDRONATE DISODIUM
CARBASTAT, CARBACHOL
CATAFLAM, DICLOFENAC POTASSIUM
CLOZARIL, CLOZAPINE
COMBIPATCH, ESTRADIOL
CYTADREN, AMINOGLUTETHIMIDE
DEFERFAL, DEFEROXAMINE MESYLATE
DIOVAN HCT, HYDROCHLOROTHIAZIDE
DIOVAN, VALSARTAN
ELIDEL, PIMECROLIMUS
ENABLEX, DARIFENACIN HYDROBROMIDE
ESIDRIX, HYDROCHLOROTHIAZIDE
ESTRADERM, ESTRADIOL
EXELON, RIVASTIGMINE TARTRATE
EXJADE, DEFERASIROX
FAMVIR, FAMCICLOVIR
FLUOR-OP, FLUOROMETHOLONE
FOCALIN XR, DEXMETHYLPHENIDATE HYDROCHLORIDE
FOCALIN, DEXMETHYLPHENIDATE HYDROCHLORIDE
FORADIL CERTIHALER, FORMOTEROL FUMARATE
FORADIL, FORMOTEROL FUMARATE
GLEEVEC, IMATINIB MESYLATE
HYDERGINE, ERGOLOID MESYLATES
INFLAMASE FORTE, PREDNISOLONE SODIUM PHOSPHATE
INFLAMASE MILD, PREDNISOLONE SODIUM PHOSPHATE
LAMISIL, TERBINAFINE HYDROCHLORIDE
LAMPRENE, CLOFAZIMINE
LESCOL XL, FLUVASTATIN SODIUM
LESCOL, FLUVASTATIN SODIUM
LOPRESSOR HCT, HYDROCHLOROTHIAZIDE
LOPRESSOR, METOPROLOL TARTRATE
LOTENSIN HCT, BENAZEPRIL HYDROCHLORIDE
LOTENSIN, BENAZEPRIL HYDROCHLORIDE
LOTREL, AMLODIPINE BESYLATE
METHERGINE, METHYLERGONOVINE MALEATE
METOPIRONE, METYRAPONE
MIACALCIN, CALCITONIN, SALMON
MIOCHOL-E, ACETYLCHOLINE CHLORIDE
MYFORTIC, MYCOPHENOLIC ACID
NEORAL, CYCLOSPORINE
OCUPRESS, CARTEOLOL HYDROCHLORIDE
PARLODEL, BROMOCRIPTINE MESYLATE
REGITINE, PHENTOLAMINE MESYLATE
RITALIN LA, METHYLPHENIDATE HYDROCHLORIDE
RITALIN, METHYLPHENIDATE HYDROCHLORIDE
RITALIN-SR, METHYLPHENIDATE HYDROCHLORIDE
SANDIMMUNE, CYCLOSPORINE
SANDOSTATIN LAR, OCTREOTIDE ACETATE
SANDOSTATIN, OCTREOTIDE ACETATE
STARLIX, NATEGLINIDE
SULF-10, SULFACETAMIDE SODIUM
TAVIST-1, CLEMISTINE FUMARATE (OTC)
TEGRETOL, CARBAMAZEPINE
TEGRETOL-XR, CARBAMAZEPINE
TORECAN, THIETHYLPERAZINE MALEATE
TRILEPTAL, OXCARBAZEPINE
VASOCIDIN, PREDNISOLONE SODIUM PHOSPHATE
VASOCON, NAPHAZOLINE HYDROCHLORIDE
VASOCON-A, ANTAZOLINE PHOSPHATE (OTC)

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

**** N ****

* NOVARTIS PHARMACEUTICALS CORP
VISKEN, PINDOLOL
VIVELLE, ESTRADIOL
VIVELLE-DOT, ESTRADIOL
VOLTAREN, DICLOFENAC SODIUM
VOLTAREN-XR, DICLOFENAC SODIUM
ZADITOR, KETOTIFEN FUMARATE (OTC)
ZELNORM, TEGASEROD MALEATE
ZOMETA, ZOLEDRONIC ACID

NOVARTIS PHARMS

* NOVARTIS PHARMACEUTICALS CORP
FEMARA, LETROZOLE
TOBI, TOBRAMYCIN

NOVEN

* NOVEN PHARMACEUTICALS INC
LIDOCAINE, LIDOCAINE

NOVEX

* NOVEX PHARMA
ALBUTEROL SULFATE, ALBUTEROL SULFATE
BETAXOLOL, BETAXOLOL HYDROCHLORIDE
BUTORPHANOL TARTRATE, BUTORPHANOL TARTRATE
CARTEOLOL HYDROCHLORIDE, CARTEOLOL HYDROCHLORIDE
CHLORHEXIDINE GLUCONATE, CHLORHEXIDINE GLUCONATE
CIMETIDINE HYDROCHLORIDE, CIMETIDINE HYDROCHLORIDE
CIPROFLOXACIN, CIPROFLOXACIN HYDROCHLORIDE
CLEMASTINE FUMARATE, CLEMASTINE FUMARATE
CROMOLYN SODIUM, CROMOLYN SODIUM
CYCLOSPORINE, CYCLOSPORINE
FLUOXETINE, FLUOXETINE HYDROCHLORIDE
IPRATROPIUM BROMIDE, IPRATROPIUM BROMIDE
LACTULOSE, LACTULOSE
LEVOBUNOLOL HYDROCHLORIDE, LEVOBUNOLOL HYDROCHLORIDE
METAPROTERENOL SULFATE, METAPROTERENOL SULFATE
MINOXIDIL (FOR MEN), MINOXIDIL (OTC)
MINOXIDIL (FOR WOMEN), MINOXIDIL (OTC)
MINOXIDIL EXTRA STRENGTH (FOR MEN), MINOXIDIL (OTC)
OFLOXACIN, OFLOXACIN
OXYBUTYNIN CHLORIDE, OXYBUTYNIN CHLORIDE
TIMOLOL MALEATE, TIMOLOL MALEATE
TOBRAMYCIN, TOBRAMYCIN

NOVO NORDISK

* NOVO NORDISK PHARMACEUTICALS INC
GLUCAGEN, GLUCAGON HYDROCHLORIDE RECOMBINANT

NOVO NORDISK INC

* NOVO NORDISK INC
ACTIVELLA, ESTRADIOL
INNOFEM, ESTRADIOL
LEVEMIR, INSULIN DETEMIR RECOMBINANT
NORDITROPIN NORDIFLEX, SOMATROPIN RECOMBINANT
NORDITROPIN, SOMATROPIN RECOMBINANT
NOVOLIN 70/30, INSULIN RECOMBINANT HUMAN (OTC)
NOVOLIN N, INSULIN SUSP ISOPHANE RECOMBINANT HUMAN (OTC)
NOVOLIN R, INSULIN RECOMBINANT HUMAN (OTC)
NOVOLOG MIX 70/30, INSULIN ASPART PROTAMINE RECOMBINANT
NOVOLOG, INSULIN ASPART RECOMBINANT
PRANDIN, REPAGLINIDE
VAGIFEM, ESTRADIOL

NOVOCOL

* NOVOCOL PHARMACEUTICAL INC
ISOCAINE HYDROCHLORIDE W/ LEVONORDEFIN, LEVONORDEFIN
ISOCAINE HYDROCHLORIDE, MEPIVACAINE HYDROCHLORIDE

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

**** 0 ****

OCLASSEN

* OCLASSEN PHARMACEUTICALS INC
CONDYLOX, PODOFILOX
CORDRAN SP, FLURANDRENOLIDE
CORDRAN, FLURANDRENOLIDE
MONODOX, DOXYCYCLINE

ODYSSEY PHARMS

* ODYSSEY PHARMACEUTICALS INC
ANTABUSE, DISULFIRAM
NYSTATIN, NYSTATIN
SURMONTIL, TRIMIPRAMINE MALEATE
URECHOLINE, BETHANECHOL CHLORIDE
VIVACTIL, PROTRIPTYLINE HYDROCHLORIDE

OHM

* OHM CORP
IBUPROFEN, IBUPROFEN (OTC)

OHM LABS

* OHM LABORATORIES INC
ATENOLOL, ATENOLOL
CHLORZOXAZONE, CHLORZOXAZONE
FUROSEMIDE, FUROSEMIDE
IBUPROFEN, IBUPROFEN
IBUPROHM COLD AND SINUS, IBUPROFEN (OTC)
IBUPROHM, IBUPROFEN
IBUPROHM, IBUPROFEN (OTC)
LOPERAMIDE HYDROCHLORIDE, LOPERAMIDE HYDROCHLORIDE (OTC)

ONY

* ONY INC
INFASURF PRESERVATIVE FREE, CALFACTANT

OPTOPICS

* OPTOPICS LABORATORIES CORP
PARACAINE, PROPARACAINE HYDROCHLORIDE
SULFACEL-15, SULFACETAMIDE SODIUM

ORAPHARMA

* ORAPHARMA INC
ARESTIN, MINOCYCLINE HYDROCHLORIDE

ORCHID HLTHCARE

* ORCHID HEALTHCARE
CEFADROXIL, CEFADROXIL/CEFADROXIL HEMIHYDRATE
CEFAZOLIN SODIUM, CEFAZOLIN SODIUM
CEFAZOLIN, CEFAZOLIN SODIUM
CEFOTAXIME SODIUM, CEFOTAXIME SODIUM
CEFOXITIN, CEFOXITIN SODIUM
CEFPROZIL, CEFPROZIL
CEFTRIAXONE, CEFTRIAXONE SODIUM
CEPHALEXIN, CEPHALEXIN

ORGANON USA INC

* ORGANON USA INC
CYCLESSA, DESOGESTREL
DESOGEN, DESOGESTREL
FOLLISTIM AQ, FOLLITROPIN ALFA/BETA
GANIRELIX ACETATE INJECTION, GANIRELIX ACETATE
IMPLANON, ETONOGESTREL
NUVARING, ETHINYL ESTRADIOL
PREGNYL, GONADOTROPIN, CHORIONIC
REMERON SOLTAB, MIRTAZAPINE
REMERON, MIRTAZAPINE
ZEMURON, ROCURONIUM BROMIDE

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

**** 0 ****

ORION

* ORION CORP
COMTAN, ENTACAPONE

ORION PHARMA INC

* ORION PHARMACEUTICALS INC
STALEVO 100, CARBIDOPA
STALEVO 150, CARBIDOPA
STALEVO 50, CARBIDOPA

ORTHO BIOTECH

* ORTHO BIOTECH PRODUCTS LP
LEUSTATIN, CLADRIBINE

ORTHO MCNEIL

* ORTHO MCNEIL PHARMACEUTICAL
HALDOL, HALOPERIDOL DECANOATE
HALDOL, HALOPERIDOL LACTATE

ORTHO MCNEIL PHARM

* ALZA CORP
URISPAS, FLAVOXATE HYDROCHLORIDE

* ORTHO MCNEIL PHARMACEUTICAL INC
AXERT, ALMOTRIPTAN MALATE
ELMIRON, PENTOSAN POLYSULFATE SODIUM
FLOXIN, OFLOXACIN
LEVAQUIN IN DEXTROSE 5% IN PLASTIC CONTAINER, LEVOFLOXACIN
LEVAQUIN, LEVOFLOXACIN
MICRONOR, NORETHINDRONE
MODICON 28, ETHINYL ESTRADIOL
ORTHO EVRA, ETHINYL ESTRADIOL
ORTHO TRI-CYCLEN LO, ETHINYL ESTRADIOL
ORTHO-CEPT, DESOGESTREL
ORTHO-NOVUM 1/35-28, ETHINYL ESTRADIOL
ORTHO-NOVUM 1/50 28, MESTRANOL
ORTHO-NOVUM 10/11-28, ETHINYL ESTRADIOL
ORTHO-NOVUM 7/7/7-28, ETHINYL ESTRADIOL
PARAFON FORTE DSC, CHLORZOXAZONE
TERAZOL 3, TERCONAZOLE
TERAZOL 7, TERCONAZOLE
TOLECTIN 600, TOLMETIN SODIUM
TOLECTIN DS, TOLMETIN SODIUM
TOLECTIN, TOLMETIN SODIUM
TOPAMAX SPRINKLE, TOPIRAMATE
TOPAMAX, TOPIRAMATE
TYLENOL W/ CODEINE NO. 3, ACETAMINOPHEN
TYLOX, ACETAMINOPHEN
ULTRACET, ACETAMINOPHEN
ULTRAM, TRAMADOL HYDROCHLORIDE

OSCIENT

* OSCIENT PHARMACEUTICALS
ANTARA (MICRONIZED), FENOFIBRATE
FACTIVE, GEMIFLOXACIN MESYLATE

OSI EYETECH

* OSI EYETECH INC
MACUGEN, PEGAPTANIB SODIUM

OSI PHARMS

* OSI PHARMACEUTICALS INC
TARCEVA, ERLOTINIB HYDROCHLORIDE

OSMOTICA PHARM

* OSMOTICA PHARMACEUTICAL CORP
NIFEDIPINE, NIFEDIPINE

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

** 0 **

OTSUKA

- * OTSUKA AMERICA PHARMACEUTICALS INC
ABILIFY, ARIPIPRAZOLE
- * OTSUKA PHARMACEUTICAL CO LTD
ABILIFY, ARIPIPRAZOLE
PLETAL, CILOSTAZOL

OVATION PHARMS

- * OVATION PHARMACEUTICALS INC
CHEMET, SUCCIMER
COGENTIN, BENZTROPINE MESYLATE
COSMEGEN, DACTINOMYCIN
DESOXYN, METHAMPHETAMINE HYDROCHLORIDE
DIURIL, CHLOROTHIAZIDE SODIUM
INDOCIN I.V., INDOMETHACIN SODIUM
MUSTARGEN, MECHLORETHAMINE HYDROCHLORIDE
NEMBUTAL SODIUM, PENTOBARBITAL SODIUM
PEGANONE, ETHOTOIN
TRANXENE SD, CLORAZEPATE DIPOTASSIUM
TRANXENE, CLORAZEPATE DIPOTASSIUM

OXFORD PHARM

- * OXFORD PHARMACEUTICAL SERVICES INC
MARPLAN, ISOCARBOXAZID

PACIFIC PHARMA

- * PACIFIC PHARMA
TIMOLOL MALEATE, TIMOLOL MALEATE
- * PACIFIC PHARMA INC
TIMOLOL MALEATE, TIMOLOL MALEATE

PADDOCK

- * PADDOCK LABORATORIES INC
AMMONIUM LACTATE, AMMONIUM LACTATE
BENZPHETAMINE HYDROCHLORIDE, BENZPHETAMINE HYDROCHLORIDE
CLINDA-DERM, CLINDAMYCIN PHOSPHATE
CLOTRIMAZOLE, CLOTRIMAZOLE
COLISTIMETHATE, COLISTIMETHATE SODIUM
COLOCORT, HYDROCORTISONE
COMPRO, PROCHLORPERAZINE
DIHYDROERGOTAMINE MESYLATE, DIHYDROERGOTAMINE MESYLATE
ERYTHRA-DERM, ERYTHROMYCIN
FLAVOXATE HYDROCHLORIDE, FLAVOXATE HYDROCHLORIDE
KIONEX, SODIUM POLYSTYRENE SULFONATE
MIDAZOLAM HYDROCHLORIDE, MIDAZOLAM HYDROCHLORIDE
MOEXIPRIL HYDROCHLORIDE, MOEXIPRIL HYDROCHLORIDE
NYSTOP, NYSTATIN
PODOFILOX, PODOFILOX
PREDNISOLONE SODIUM PHOSPHATE, PREDNISOLONE SODIUM PHOSPHATE
PROMETHAZINE HYDROCHLORIDE, PROMETHAZINE HYDROCHLORIDE
TESTOSTERONE CYPIONATE, TESTOSTERONE CYPIONATE
TESTOSTERONE ETHANATE, TESTOSTERONE ENANTHATE

PAR PHARM

- * PAR PHARMACEUTICAL
MELOXICAM, MELOXICAM
NABUMETONE, NABUMETONE
PERGOLIDE MESYLATE, PERGOLIDE MESYLATE
SUMYCIN, TETRACYCLINE HYDROCHLORIDE
- * PAR PHARMACEUTICAL INC
ALLOPURINOL, ALLOPURINOL
AMILORIDE HYDROCHLORIDE, AMILORIDE HYDROCHLORIDE
BENZTROPINE MESYLATE, BENZTROPINE MESYLATE
BUSPIRONE HYDROCHLORIDE, BUSPIRONE HYDROCHLORIDE
CABERGOLINE, CABERGOLINE
CAPOTEN, CAPTOPRIL
CARISOPRODOL AND ASPIRIN, ASPIRIN

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

**** P ****

* PAR PHARMACEUTICAL INC
CEFACLOX, CEFACLOX
CHLORDIAZEPOXIDE AND AMITRIPTYLINE HYDROCHLORIDE, AMITRIPTYLINE HYDROCHLORIDE
CHLORPROPAMIDE, CHLORPROPAMIDE
CHLORZOXAZONE, CHLORZOXAZONE
CHOLESTYRAMINE LIGHT, CHOLESTYRAMINE
CHOLESTYRAMINE, CHOLESTYRAMINE
CLOMIPHENE CITRATE, CLOMIPHENE CITRATE
CYPROHEPTADINE HYDROCHLORIDE, CYPROHEPTADINE HYDROCHLORIDE
DEXAMETHASONE, DEXAMETHASONE
DIAZEPAM, DIAZEPAM
DIPHENOXYLATE HYDROCHLORIDE AND ATROPINE SULFATE, ATROPINE SULFATE
DOXEPIN HYDROCHLORIDE, DOXEPIN HYDROCHLORIDE
DOXYCYCLINE HYCLATE, DOXYCYCLINE HYCLATE
DOXYCYCLINE, DOXYCYCLINE
ESTAZOLAM, ESTAZOLAM
FENOFIBRATE, FENOFIBRATE
FENOPROFEN CALCIUM, FENOPROFEN CALCIUM
FLUOXETINE HYDROCHLORIDE, FLUOXETINE HYDROCHLORIDE
FLUOXETINE, FLUOXETINE HYDROCHLORIDE
FLUPHENAZINE HYDROCHLORIDE, FLUPHENAZINE HYDROCHLORIDE
FLURAZEPAM HYDROCHLORIDE, FLURAZEPAM HYDROCHLORIDE
FLUTAMIDE, FLUTAMIDE
GLIMEPIRIDE, GLIMEPIRIDE
HALOPERIDOL, HALOPERIDOL
HYDRALAZINE HYDROCHLORIDE, HYDRALAZINE HYDROCHLORIDE
HYDRA-ZIDE, HYDRALAZINE HYDROCHLORIDE
HYDROFLUMETHIAZIDE, HYDROFLUMETHIAZIDE
HYDRO-RIDE, AMILORIDE HYDROCHLORIDE
HYDROXYUREA, HYDROXYUREA
IBUPROFEN, IBUPROFEN
IBUPROFEN, IBUPROFEN (OTC)
IMIPRAMINE HYDROCHLORIDE, IMIPRAMINE HYDROCHLORIDE
ISOSORBIDE DINITRATE, ISOSORBIDE DINITRATE
LEUCOVORIN CALCIUM, LEUCOVORIN CALCIUM
LISINOPRIL, LISINOPRIL
MECLIZINE HYDROCHLORIDE, MECLIZINE HYDROCHLORIDE
MEGACE ES, MEGESTROL ACETATE
MEGESTROL ACETATE, MEGESTROL ACETATE
METAPROTERENOL SULFATE, METAPROTERENOL SULFATE
METHOCARBAMOL AND ASPIRIN, ASPIRIN
METHOCARBAMOL, METHOCARBAMOL
METHYLDOPA, METHYLDOPA
METHYLPREDNISOLONE, METHYLPREDNISOLONE
METOPROLOL TARTRATE, METOPROLOL TARTRATE
METRONIDAZOLE, METRONIDAZOLE
MINOXIDIL, MINOXIDIL
NYSTATIN, NYSTATIN
OFLOXACIN, OFLOXACIN
ORPHENGESIC FORTE, ASPIRIN
ORPHENGESIC, ASPIRIN
PROPOXYPHENE HYDROCHLORIDE, PROPOXYPHENE HYDROCHLORIDE
PROPRANOLOL HYDROCHLORIDE, PROPRANOLOL HYDROCHLORIDE
RANITIDINE HYDROCHLORIDE, RANITIDINE HYDROCHLORIDE
SULFAMETHOPRIM, SULFAMETHOXAZOLE
SULFINPYRAZONE, SULFINPYRAZONE
SUMYCIN, TETRACYCLINE HYDROCHLORIDE
TEMAZEPAM, TEMAZEPAM
TOLAZAMIDE, TOLAZAMIDE
TORSEMIDE, TORSEMIDE

PARENTA PHARMS

* PARENTA PHARMACEUTICALS INC
DIAZEPAM, DIAZEPAM

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

** P **

PARKE DAVIS

- * PARKE DAVIS DIV WARNER LAMBERT CO
BENADRYL PRESERVATIVE FREE, DIPHENHYDRAMINE HYDROCHLORIDE
BENADRYL, DIPHENHYDRAMINE HYDROCHLORIDE
CELONTIN, METHSUXIMIDE
CEREBYX, FOSPHENYTOIN SODIUM
CHOLEDYL SA, OXTRIPHYLLINE
DILANTIN, PHENYTOIN SODIUM
DILANTIN-125, PHENYTOIN
NARDIL, PHENELZINE SULFATE
NEURONTIN, GABAPENTIN
ZARONTIN, ETHOSUXIMIDE
- * PARKE DAVIS PHARMACEUTICAL RESEARCH DIV WARNER LAMBERT CO
ZARONTIN, ETHOSUXIMIDE

PARKEDALE

- * PARKEDALE PHARMACEUTICALS INC
CHLOROMYCETIN, CHLORAMPHENICOL SODIUM SUCCINATE
COLY-MYCIN M, COLISTIMETHATE SODIUM
COLY-MYCIN S, COLISTIN SULFATE
KETALAR, KETAMINE HYDROCHLORIDE
PITOCIN, OXYTOCIN

PDL BIOPHARMA INC

- * PDL BIOPHARMA INC
BUSULFEX, BUSULFAN
CARDENE SR, NICARDIPINE HYDROCHLORIDE
CARDENE, NICARDIPINE HYDROCHLORIDE

PEDINOL

- * PEDINOL PHARMACAL INC
GRIS-PEG, GRISEOFULVIN, ULTRAMICROCRYSTALLINE
NALFON 200, FENOPROFEN CALCIUM
NALFON, FENOPROFEN CALCIUM

PERRIGO

- * L PERRIGO CO
ACETAMINOPHEN, ACETAMINOPHEN (OTC)
ACETAMINOPHEN, ASPIRIN AND CAFFEINE, ACETAMINOPHEN (OTC)
CAP-PROFEN, IBUPROFEN (OTC)
CHILDREN'S IBUPROFEN, IBUPROFEN (OTC)
CROMOLYN SODIUM, CROMOLYN SODIUM (OTC)
DOXYLAMINE SUCCINATE, DOXYLAMINE SUCCINATE (OTC)
FAMOTIDINE, FAMOTIDINE (OTC)
IBUPROFEN AND PSEUDOEPHEDRINE HYDROCHLORIDE, IBUPROFEN (OTC)
IBUPROFEN, IBUPROFEN (OTC)
LOPERAMIDE HYDROCHLORIDE AND SIMETHICONE, LOPERAMIDE HYDROCHLORIDE (OTC)
LOPERAMIDE HYDROCHLORIDE, LOPERAMIDE HYDROCHLORIDE (OTC)
LORATADINE, LORATADINE (OTC)
MICONAZOLE 3 COMBINATION PACK, MICONAZOLE NITRATE (OTC)
MICONAZOLE NITRATE, MICONAZOLE NITRATE (OTC)
MINOXIDIL EXTRA STRENGTH (FOR MEN), MINOXIDIL (OTC)
NAPROXEN SODIUM AND PSEUDOEPHEDRINE HYDROCHLORIDE, NAPROXEN SODIUM (OTC)
NAPROXEN SODIUM, NAPROXEN SODIUM (OTC)
RANITIDINE, RANITIDINE HYDROCHLORIDE (OTC)
TAB-PROFEN, IBUPROFEN (OTC)
TIOCONAZOLE, TIOCONAZOLE (OTC)
- * PERRIGO CO
CICLOPIROX, CICLOPIROX
CIMETIDINE, CIMETIDINE (OTC)
CLEMASTINE FUMARATE, CLEMASTINE FUMARATE (OTC)
FAMOTIDINE, FAMOTIDINE
FAMOTIDINE, FAMOTIDINE (OTC)
HALOBETASOL PROPIONATE, HALOBETASOL PROPIONATE
IBUPROFEN, IBUPROFEN (OTC)
JUNIOR STRENGTH IBUPROFEN, IBUPROFEN (OTC)

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

** P **

* PERRIGO CO
LOPERAMIDE HYDROCHLORIDE, LOPERAMIDE HYDROCHLORIDE (OTC)
MICONAZOLE NITRATE COMBINATION PACK, MICONAZOLE NITRATE (OTC)
MINOXIDIL (FOR MEN), MINOXIDIL (OTC)
MINOXIDIL (FOR WOMEN), MINOXIDIL (OTC)
MOMETASONE FUROATE, MOMETASONE FUROATE
NICOTINE POLACRILEX, NICOTINE POLACRILEX (OTC)
PSEUDOEPHEDRINE HYDROCHLORIDE, PSEUDOEPHEDRINE HYDROCHLORIDE (OTC)

PERRIGO ISRAEL

* PERRIGO ISRAEL PHARMACEUTICALS LTD
HALOBETASOL PROPIONATE, HALOBETASOL PROPIONATE

PERRIGO NEW YORK

* PERRIGO NEW YORK INC
AMMONIUM LACTATE, AMMONIUM LACTATE
BETAMETHASONE DIPROPIONATE, BETAMETHASONE DIPROPIONATE
CICLOPIROX, CICLOPIROX
CLINDAMYCIN PHOSPHATE, CLINDAMYCIN PHOSPHATE
DESOXIMETASONE, DESOXIMETASONE
ECONAZOLE NITRATE, ECONAZOLE NITRATE
ERYTHROMYCIN, ERYTHROMYCIN
FLUTICASONE PROPIONATE, FLUTICASONE PROPIONATE
GENTAMICIN, GENTAMICIN SULFATE
HYDROCORTISONE VALERATE, HYDROCORTISONE VALERATE
HYDROCORTISONE, HYDROCORTISONE
KETOCONAZOLE, KETOCONAZOLE
MESALAMINE, MESALAMINE
MINOXIDIL EXTRA STRENGTH (FOR MEN), MINOXIDIL (OTC)
MOMETASONE FUROATE, MOMETASONE FUROATE
MUPIROCIN, MUPIROCIN
NYSTATIN, NYSTATIN
PERMETHRIN, PERMETHRIN
PERMETHRIN, PERMETHRIN (OTC)
PROMETHAZINE HYDROCHLORIDE, PROMETHAZINE HYDROCHLORIDE
SELENIUM SULFIDE, SELENIUM SULFIDE
TERCONAZOLE, TERCONAZOLE
TRIAMCINOLONE ACETONIDE, TRIAMCINOLONE ACETONIDE
TRIDESILON, DESONIDE

PERRIGO R AND D

* PERRIGO R AND D CO
IBUPROFEN, IBUPROFEN
IBUPROFEN, IBUPROFEN (OTC)
NAPROXEN, NAPROXEN
NICOTINE POLACRILEX, NICOTINE POLACRILEX (OTC)
SIMVASTATIN, SIMVASTATIN

PERSONAL PRODS

* PERSONAL PRODUCTS CO
MONISTAT 1 COMBINATION PACK, MICONAZOLE NITRATE (OTC)
MONISTAT 3 COMBINATION PACK (PREFILLED), MICONAZOLE NITRATE (OTC)
MONISTAT 3 COMBINATION PACK, MICONAZOLE NITRATE (OTC)
MONISTAT 3, MICONAZOLE NITRATE

PFIZER

* PFIZER AGRICULTURAL DIV
ANTIVERT, MECLIZINE HYDROCHLORIDE
* PFIZER CENTRAL RESEARCH
DIFLUCAN, FLUCONAZOLE
ZITHROMAX, AZITHROMYCIN
* PFIZER CHEMICALS DIV PFIZER INC
DIFLUCAN, FLUCONAZOLE
ZITHROMAX, AZITHROMYCIN
* PFIZER INC
CADUET, AMLODIPINE BESYLATE
CARDURA XL, DOXAZOSIN MESYLATE

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

**** P ****

- * PFIZER INC
 - DIFLUCAN IN DEXTROSE 5% IN PLASTIC CONTAINER, FLUCONAZOLE
 - DIFLUCAN IN SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER, FLUCONAZOLE
 - DIFLUCAN IN SODIUM CHLORIDE 0.9%, FLUCONAZOLE
 - EXUBERA, INSULIN RECOMBINANT HUMAN
 - GEODON, ZIPRASIDONE HYDROCHLORIDE
 - GEODON, ZIPRASIDONE MESYLATE
 - GLUCOTROL XL, GLIPIZIDE
 - GLUCOTROL, GLIPIZIDE
 - LIPITOR, ATORVASTATIN CALCIUM
 - MINIPRESS XL, PRAZOSIN HYDROCHLORIDE
 - MINIZIDE, POLYTHIAZIDE
 - NAVANE, THIOTHIXENE
 - NAVANE, THIOTHIXENE HYDROCHLORIDE
 - NORVASC, AMLODIPINE BESYLATE
 - PROCARDIA, NIFEDIPINE
 - RENESE, POLYTHIAZIDE
 - REVATIO, SILDENAFIL CITRATE
 - RID MOUSSE, PIPERONYL BUTOXIDE (OTC)
 - UNASYN, AMPICILLIN SODIUM
 - VFEND, VORICONAZOLE
 - ZITHROMAX, AZITHROMYCIN
 - ZYRTEC, CETIRIZINE HYDROCHLORIDE
 - ZYRTEC-D 12 HOUR, CETIRIZINE HYDROCHLORIDE
- * PFIZER LABORATORIES DIV PFIZER INC
 - CARDURA, DOXAZOSIN MESYLATE
 - CEFOBID IN PLASTIC CONTAINER, CEFOPERAZONE SODIUM
 - CEFOBID, CEFOPERAZONE SODIUM
 - DIABINESE, CHLORPROPAMIDE
 - FELDENE, PIROXICAM
 - GEOCILLIN, CARBENICILLIN INDANYL SODIUM
 - MINIPRESS, PRAZOSIN HYDROCHLORIDE
 - MITHRACIN, PLICAMYCIN
 - MODERIL, RESCINNAMINE
 - PENICILLIN G POTASSIUM, PENICILLIN G POTASSIUM
 - PERMAPEN, PENICILLIN G BENZATHINE
 - PFIZERPEN, PENICILLIN G POTASSIUM
 - PROCARDIA XL, NIFEDIPINE
 - RENESE-R, POLYTHIAZIDE
 - SINEQUAN, DOXEPIN HYDROCHLORIDE
 - STREPTOMYCIN SULFATE, STREPTOMYCIN SULFATE
 - TERRA-CORTRIL, HYDROCORTISONE ACETATE
 - TERRAMYCIN W/ POLYMYXIN B SULFATE, OXYTETRACYCLINE HYDROCHLORIDE
 - TERRAMYCIN, LIDOCAINE HYDROCHLORIDE
 - TERRAMYCIN, OXYTETRACYCLINE HYDROCHLORIDE
 - UNASYN, AMPICILLIN SODIUM
 - UNISOM, DOXYLAMINE SUCCINATE (OTC)
 - VIBRAMYCIN, DOXYCYCLINE
 - VIBRAMYCIN, DOXYCYCLINE CALCIUM
 - VIBRAMYCIN, DOXYCYCLINE HYCLATE
 - VIBRA-TABS, DOXYCYCLINE HYCLATE
 - VISTARIL, HYDROXYZINE HYDROCHLORIDE
 - VISTARIL, HYDROXYZINE PAMOATE
- * PFIZER PHARMACEUTICALS INC
 - ZOLOFT, SERTRALINE HYDROCHLORIDE
- * PFIZER PHARMACEUTICALS PRODUCTION CORP LTD
 - TIKOSYN, DOFETILIDE

PFIZER GLOBAL

- * PFIZER GLOBAL RESEARCH DEVELOPMENT
 - ZMAX, AZITHROMYCIN

PFIZER INC

- * PFIZER INC
 - CHANTIX, VARENICLINE TARTRATE

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

**** P ****

* PFIZER INC
ELLENC, EPIRUBICIN HYDROCHLORIDE
GEODON, ZIPRASIDONE HYDROCHLORIDE
NICOTROL, NICOTINE

PFIZER IRELAND

* PFIZER IRELAND PHARMACEUTICALS
RELPA, ELETRIPTAN HYDROBROMIDE
VIAGRA, SILDENAFIL CITRATE

PFIZER PHARMS

* PFIZER PHARMACEUTICALS LTD
ACCUPRIL, QUINAPRIL HYDROCHLORIDE
ACCURETIC, HYDROCHLOROTHIAZIDE
DILANTIN, PHENYTOIN
LOPID, GEMFIBROZIL
NEURONTIN, GABAPENTIN
NITROSTAT, NITROGLYCERIN

PHARM ASSOC

* PHARMACEUTICAL ASSOC INC
DOXEPIN HYDROCHLORIDE, DOXEPIN HYDROCHLORIDE
NORTRIPTYLINE HYDROCHLORIDE, NORTRIPTYLINE HYDROCHLORIDE
PERPHENAZINE, PERPHENAZINE
PREDNISOLONE SODIUM PHOSPHATE, PREDNISOLONE SODIUM PHOSPHATE

* PHARMACEUTICAL ASSOC INC DIV BEACH PRODUCTS
ACETAMINOPHEN AND CODEINE PHOSPHATE, ACETAMINOPHEN
AMANTADINE HYDROCHLORIDE, AMANTADINE HYDROCHLORIDE
CHLORPROMAZINE HYDROCHLORIDE, CHLORPROMAZINE HYDROCHLORIDE
CIMETIDINE HYDROCHLORIDE, CIMETIDINE HYDROCHLORIDE
DIPHENHYDRAMINE HYDROCHLORIDE, DIPHENHYDRAMINE HYDROCHLORIDE
ETHOSUXIMIDE, ETHOSUXIMIDE
FLUOXETINE, FLUOXETINE HYDROCHLORIDE
FLUPHENAZINE HYDROCHLORIDE, FLUPHENAZINE HYDROCHLORIDE
HALOPERIDOL, HALOPERIDOL LACTATE
HYDROCODONE BITARTRATE AND ACETAMINOPHEN, ACETAMINOPHEN
LACTULOSE, LACTULOSE
METOCLOPRAMIDE HYDROCHLORIDE, METOCLOPRAMIDE HYDROCHLORIDE
OXYBUTYNIN CHLORIDE, OXYBUTYNIN CHLORIDE
PREDNISOLONE, PREDNISOLONE
PROMETHAZINE HYDROCHLORIDE AND CODEINE PHOSPHATE, CODEINE PHOSPHATE
THEOPHYLLINE, THEOPHYLLINE
THIORIDAZINE HYDROCHLORIDE, THIORIDAZINE HYDROCHLORIDE
TRIHENXYPHENIDYL HYDROCHLORIDE, TRIHENXYPHENIDYL HYDROCHLORIDE
VALPROIC ACID, VALPROIC ACID

PHARM SPEC

* PHARMACEUTICAL SPECIALIST ASSOC
GENTAMICIN SULFATE, GENTAMICIN SULFATE

PHARM VENTURES

* PHARMACEUTICAL VENTURES LTD
METOCLOPRAMIDE HYDROCHLORIDE, METOCLOPRAMIDE HYDROCHLORIDE
TRIHENXYPHENIDYL HYDROCHLORIDE, TRIHENXYPHENIDYL HYDROCHLORIDE

PHARMA SERVE NY

* PHARMA SERVE INC SUB TORIGIAN LABORATORIES
AMINOPHYLLINE, AMINOPHYLLINE

PHARMACHEMIE

* PHARMACHEMIE BV
CARBOPLATIN, CARBOPLATIN
CISPLATIN, CISPLATIN
DOXORUBICIN HYDROCHLORIDE, DOXORUBICIN HYDROCHLORIDE
ETOPOSIDE, ETOPOSIDE
LEUCOVORIN CALCIUM, LEUCOVORIN CALCIUM
TAMOXIFEN CITRATE, TAMOXIFEN CITRATE

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

**** P ****

PHARMACHEMIE USA

* PHARMACHEMIE USA INC
LEUCOVORIN CALCIUM, LEUCOVORIN CALCIUM

PHARMACIA

* PHARMACIA CORP
MAXAQUIN, LOMEFLOXACIN HYDROCHLORIDE

PHARMACIA AND UPJOHN

* PHARMACIA AND UPJOHN
XANAX XR, ALPRAZOLAM

* PHARMACIA AND UPJOHN CO
ADRIAMYCIN PFS, DOXORUBICIN HYDROCHLORIDE
ANSAID, FLURBIPROFEN
AROMASIN, EXEMESTANE
AZULFIDINE EN-TABS, SULFASALAZINE
AZULFIDINE, SULFASALAZINE
BACITRACIN, BACITRACIN
CAMPTOSAR, IRINOTECAN HYDROCHLORIDE
CAVERJECT, ALPROSTADIL
CLEOCIN HYDROCHLORIDE, CLINDAMYCIN HYDROCHLORIDE
CLEOCIN PHOSPHATE IN DEXTROSE 5% IN PLASTIC CONTAINER, CLINDAMYCIN PHOSPHATE
CLEOCIN PHOSPHATE, CLINDAMYCIN PHOSPHATE
CLEOCIN T, CLINDAMYCIN PHOSPHATE
CLEOCIN, CLINDAMYCIN PALMITATE HYDROCHLORIDE
CLEOCIN, CLINDAMYCIN PHOSPHATE
COLESTID, COLESTIPOL HYDROCHLORIDE
CORTEF, HYDROCORTISONE
CORTISONE ACETATE, CORTISONE ACETATE
CORVERT, IBUTILIDE FUMARATE
CYKLOKAPRON, TRANEXAMIC ACID
DEPO-ESTRADIOL, ESTRADIOL CYPIONATE
DEPO-MEDROL, METHYLPREDNISOLONE ACETATE
DEPO-PROVERA, MEDROXYPROGESTERONE ACETATE
DEPO-TESTOSTERONE, TESTOSTERONE CYPIONATE
DETROL LA, TOLTERODINE TARTRATE
DETROL, TOLTERODINE TARTRATE
DIDREX, BENZPHETAMINE HYDROCHLORIDE
DOSTINEX, CABERGOLINE
EMCYT, ESTRAMUSTINE PHOSPHATE SODIUM
ESTRING, ESTRADIOL
FLAVORED COLESTID, COLESTIPOL HYDROCHLORIDE
FRAGMIN, DALTEPARIN SODIUM
GENOTROPIN PRESERVATIVE FREE, SOMATROPIN RECOMBINANT
GENOTROPIN, SOMATROPIN RECOMBINANT
GLYNASE, GLYBURIDE
GLYSET, MIGLITOL
HALCION, TRIAZOLAM
HALOTESTIN, FLUOXYMESTERONE
HEMABATE, CARBOPROST TROMETHAMINE
HEPARIN SODIUM, HEPARIN SODIUM
IDAMYCIN PFS, IDARUBICIN HYDROCHLORIDE
LINCOCIN, LINCOMYCIN HYDROCHLORIDE
LONITEN, MINOXIDIL
MEDROL, METHYLPREDNISOLONE
MICRONASE, GLYBURIDE
MYCOBUTIN, RIFABUTIN
NICOTROL, NICOTINE
OGEN .625, ESTROPIPATE
OGEN 1.25, ESTROPIPATE
OGEN 2.5, ESTROPIPATE
OGEN 5, ESTROPIPATE
OGEN, ESTROPIPATE
PREPIDIL, DINOPROSTONE
PROSTIN E2, DINOPROSTONE

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

**** P ****

- * PHARMACIA AND UPJOHN CO
PROSTIN VR PEDIATRIC, ALPROSTADIL
PROVERA, MEDROXYPROGESTERONE ACETATE
R-GENE 10, ARGININE HYDROCHLORIDE
SOLU-CORTEF, HYDROCORTISONE SODIUM SUCCINATE
SOLU-MEDROL, METHYLPREDNISOLONE SODIUM SUCCINATE
SOMAVERT, PEGVISOMANT
TOLINASE, TOLAZAMIDE
TROBICIN, SPECTINOMYCIN HYDROCHLORIDE
VANTIN, CEFPODOXIME PROXETIL
XALATAN, LATANOPROST
XANAX, ALPRAZOLAM
ZINECARD, DEXRAZOXANE HYDROCHLORIDE
ZYVOX, LINEZOLID
- * PHARMACIA AND UPJOHN SUB PFIZER INC
DEPO-SUBQ PROVERA 104, MEDROXYPROGESTERONE ACETATE

PHARMADERM

- * PHARMADERM DIV ALTANA INC
BACITRACIN-NEOMYCIN-POLYMYXIN W/ HYDROCORTISONE ACETATE, BACITRACIN

PHARMAFORCE

- * PHARMAFORCE INC
CAFFEINE CITRATE, CAFFEINE CITRATE
CYCLOSPORINE, CYCLOSPORINE
ESMOLOL HYDROCHLORIDE, ESMOLOL HYDROCHLORIDE
FENOLDOPAM MESYLATE, FENOLDOPAM MESYLATE
NEOMYCIN AND POLYMYXIN B SULFATES AND GRAMICIDIN, GRAMICIDIN
NEOMYCIN AND POLYMYXIN B SULFATES AND HYDROCORTISONE, HYDROCORTISONE
NYSTATIN, NYSTATIN
OFLOXACIN, OFLOXACIN
ONDANSETRON HYDROCHLORIDE PRESERVATIVE FREE, ONDANSETRON HYDROCHLORIDE
ONDANSETRON HYDROCHLORIDE, ONDANSETRON HYDROCHLORIDE
PROMETHAZINE HYDROCHLORIDE, PROMETHAZINE HYDROCHLORIDE

PHARMASEAL

- * PHARMASEAL DIV BAXTER HEALTHCARE CORP
PHARMASEAL SCRUB CARE, CHLORHEXIDINE GLUCONATE (OTC)

PHARMATON

- * PHARMATON LTD
LIDOCATON, EPINEPHRINE

PHARMAX

- * PHARMAX GROUP INC
FOLIC ACID, FOLIC ACID

PHARMELLE

- * PHARMELLE LLC
AVC, SULFANILAMIDE

PHARMERAL

- * PHARMERAL INC
ACETAMINOPHEN AND CODEINE PHOSPHATE, ACETAMINOPHEN

PHARMION

- * PHARMION CORP
INNOHEP, TINZAPARIN SODIUM
VIDAZA, AZACITIDINE

PHOTOCURE ASA

- * PHOTOCURE ASA
METVIXIA, METHYL AMINOLEVULINATE HYDROCHLORIDE

PIERRE FABRE

- * PIERRE FABRE MEDICAMENT
NAVELBINE, VINORELBINE TARTRATE

PLANTEX

- * PLANTEX USA INC DIV IKAPHARM INC
SULFAMETHOXAZOLE AND TRIMETHOPRIM SINGLE STRENGTH, SULFAMETHOXAZOLE

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

**** P ****

PLIVA

* PLIVA INC
ALBUTEROL SULFATE, ALBUTEROL SULFATE
AMITRIPTYLINE HYDROCHLORIDE, AMITRIPTYLINE HYDROCHLORIDE
ATENOLOL AND CHLORTHALIDONE, ATENOLOL
ATENOLOL, ATENOLOL
AZITHROMYCIN, AZITHROMYCIN
BENZTROPINE MESYLATE, BENZTROPINE MESYLATE
CARBAMAZEPINE, CARBAMAZEPINE
CARBOPLATIN, CARBOPLATIN
CHLORPROPAMIDE, CHLORPROPAMIDE
CHLORTHALIDONE, CHLORTHALIDONE
CIMETIDINE, CIMETIDINE
CIPROFLOXACIN, CIPROFLOXACIN HYDROCHLORIDE
CITALOPRAM HYDROBROMIDE, CITALOPRAM HYDROBROMIDE
CYCLOBENZAPRINE HYDROCHLORIDE, CYCLOBENZAPRINE HYDROCHLORIDE
CYCLOSPORINE, CYCLOSPORINE
CYPROHEPTADINE HYDROCHLORIDE, CYPROHEPTADINE HYDROCHLORIDE
DESIPRAMINE HYDROCHLORIDE, DESIPRAMINE HYDROCHLORIDE
DEXCHLORPHENIRAMINE MALEATE, DEXCHLORPHENIRAMINE MALEATE
DICLOFENAC SODIUM, DICLOFENAC SODIUM
DOXAZOSIN MESYLATE, DOXAZOSIN MESYLATE
FLUCONAZOLE, FLUCONAZOLE
FLUOXETINE, FLUOXETINE HYDROCHLORIDE
FLURBIPROFEN, FLURBIPROFEN
GLIPIZIDE, GLIPIZIDE
HYDRALAZINE HYDROCHLORIDE, HYDRALAZINE HYDROCHLORIDE
HYDROXYZINE HYDROCHLORIDE, HYDROXYZINE HYDROCHLORIDE
IBUPROFEN, IBUPROFEN
KETOCONAZOLE, KETOCONAZOLE
KETOROLAC TROMETHAMINE, KETOROLAC TROMETHAMINE
MECLIZINE HYDROCHLORIDE, MECLIZINE HYDROCHLORIDE
METHYLDOPA, METHYLDOPA
METOCLOPRAMIDE HYDROCHLORIDE, METOCLOPRAMIDE HYDROCHLORIDE
METRONIDAZOLE, METRONIDAZOLE
NAPROXEN SODIUM, NAPROXEN SODIUM
NAPROXEN, NAPROXEN
OXYBUTYNIN CHLORIDE, OXYBUTYNIN CHLORIDE
PENTOXIFYLLINE, PENTOXIFYLLINE
PROPAFENONE HYDROCHLORIDE, PROPAFENONE HYDROCHLORIDE
PROPRANOLOL HYDROCHLORIDE AND HYDROCHLOROTHIAZIDE, HYDROCHLOROTHIAZIDE
PROPRANOLOL HYDROCHLORIDE, PROPRANOLOL HYDROCHLORIDE
SULFAMETHOXAZOLE AND TRIMETHOPRIM, SULFAMETHOXAZOLE
THEOPHYLLINE, THEOPHYLLINE
TRAMADOL HYDROCHLORIDE, TRAMADOL HYDROCHLORIDE
TRAZODONE HYDROCHLORIDE, TRAZODONE HYDROCHLORIDE
TRIAMTERENE AND HYDROCHLOROTHIAZIDE, HYDROCHLOROTHIAZIDE
VERAPAMIL HYDROCHLORIDE, VERAPAMIL HYDROCHLORIDE
WARFARIN SODIUM, WARFARIN SODIUM

PLIVA HRVATSKA DOO

* PLIVA HRVATSKA DOO
ONDANSETRON HYDROCHLORIDE, ONDANSETRON HYDROCHLORIDE
PRAVASTATIN SODIUM, PRAVASTATIN SODIUM

PLIVA PHARM IND

* PLIVA PHARMACEUTICAL INDUSTRY INC
TORSEMIDE, TORSEMIDE

POHL BOSKAMP

* POHL BOSKAMP
NITROLINGUAL PUMPSPRAY, NITROGLYCERIN

POLYMEDICA

* POLYMEDICA INDUSTRIES INC
ANESTACON, LIDOCAINE HYDROCHLORIDE

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

**** P ****

- * POLYMEDICA INDUSTRIES INC
NEOPAP, ACETAMINOPHEN (OTC)
PROMETHACON, PROMETHAZINE HYDROCHLORIDE

POPULATION COUNCIL

- * POPULATION COUNCIL CENTER FOR BIOMEDICAL RESEARCH
NORPLANT II, LEVONORGESTREL

PRIMAPHARM

- * PRIMAPHARM INC
HYDASE, HYALURONIDASE

PROCTER AND GAMBLE

- * PROCTER AND GAMBLE CO
DANTRIUM, DANTROLENE SODIUM
MACRODANTIN, NITROFURANTOIN, MACROCRYSTALLINE
- * PROCTER AND GAMBLE PHARMACEUTICALS INC
ACTONEL WITH CALCIUM (COPACKAGED), CALCIUM CARBONATE
- * PROCTER AND GAMBLE PHARMACEUTICALS INC SUB PROCTER AND GAMBLE CO
ACTONEL, RISEDRONATE SODIUM
ASACOL, MESALAMINE
DANTRIUM, DANTROLENE SODIUM
DIDRONEL, ETIDRONATE DISODIUM
MACROBID, NITROFURANTOIN

PROMETHEUS LABS

- * PROMETHEUS LABORATORIES INC
HELIDAC, BISMUTH SUBSALICYLATE
IMURAN, AZATHIOPRINE
MERCAPTOPYRINE, MERCAPTOPYRINE
RIDAURA, AURANOFIN
TRANDATE, LABETALOL HYDROCHLORIDE
ZYLORIM, ALLOPURINOL

PURDUE FREDERICK

- * PURDUE FREDERICK CO
MS CONTIN, MORPHINE SULFATE
UNIPHYL, THEOPHYLLINE

PURDUE PHARMA LP

- * PURDUE PHARMA LP
OXYCONTIN, OXYCODONE HYDROCHLORIDE

PUREPAC PHARM

- * PUREPAC PHARMACEUTICAL CO
AMPICILLIN TRIHYDRATE, AMPICILLIN/AMPICILLIN TRIHYDRATE
DIPYRIDAMOLE, DIPYRIDAMOLE

QLT

- * QLT INC
VISUDYNE, VERTEPORFIN

QLT USA

- * QLT USA INC
ATRIDOX, DOXYCYCLINE HYCLATE
BETAMETHASONE DIPROPIONATE, BETAMETHASONE DIPROPIONATE
CLOBETASOL PROPIONATE, CLOBETASOL PROPIONATE
ELIGARD, LEUPROLIDE ACETATE
ERYTHROMYCIN AND BENZOYL PEROXIDE, BENZOYL PEROXIDE
FLUTICASONE PROPIONATE, FLUTICASONE PROPIONATE
KETOCONAZOLE, KETOCONAZOLE
LIDOCAINE AND PRILOCAINE, LIDOCAINE
METRONIDAZOLE, METRONIDAZOLE
MOMETASONE FUROATE, MOMETASONE FUROATE

QOL MEDCL

- * QOL MEDICAL LLC
ELLIOTTS B SOLUTION, CALCIUM CHLORIDE
ETHAMOLIN, ETHANOLAMINE OLEATE
GLOFIL-125, IOTHALAMATE SODIUM, I-125

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

**** Q ****

* QOL MEDICAL LLC
NASCOBAL, CYANOCOBALAMIN
SUCRAID, SACROSIDASE

QPHARMA

* QPHARMA, LLC
FLUNISOLIDE, FLUNISOLIDE

QUESTCOR PHARMS

* QUESTCOR PHARMACEUTICALS INC
DORAL, QUAZEPAM
H.P. ACTHAR GEL, CORTICOTROPIN

R R REGISTRATIONS

* R AND R REGISTRATIONS
THYROSAFE, POTASSIUM IODIDE (OTC)

RADIUS PHARMS

* RADIUS PHARMACEUTICALS INC
KETOPROFEN, KETOPROFEN
PENTOXIFYLLINE, PENTOXIFYLLINE

RANBAXY

* RANBAXY LABORATORIES LTD
ACYCLOVIR, ACYCLOVIR
AMOXICILLIN AND CLAVULANATE POTASSIUM, AMOXICILLIN
AMOXICILLIN, AMOXICILLIN
BENAZEPRIL HYDROCHLORIDE AND HYDROCHLOROTHIAZIDE, BENAZEPRIL HYDROCHLORIDE
BENAZEPRIL HYDROCHLORIDE, BENAZEPRIL HYDROCHLORIDE
CARBILEV, CARBIDOPA
CEFADROXIL, CEFADROXIL/CEFADROXIL HEMIHYDRATE
CEFPODOXIME PROXETIL, CEFPODOXIME PROXETIL
CEFPROZIL, CEFPROZIL
CEFUROXIME AXETIL, CEFUROXIME AXETIL
CEPHALEXIN, CEPHALEXIN
CLARITHROMYCIN, CLARITHROMYCIN
CLINDAMYCIN HYDROCHLORIDE, CLINDAMYCIN HYDROCHLORIDE
CLORAZEPATE DIPOTASSIUM, CLORAZEPATE DIPOTASSIUM
DISPERMOX, AMOXICILLIN
DOXYCYCLINE, DOXYCYCLINE
ENALAPRIL MALEATE, ENALAPRIL MALEATE
ETODOLAC, ETODOLAC
FENOFIBRATE, FENOFIBRATE
FLECAINIDE ACETATE, FLECAINIDE ACETATE
FLUCONAZOLE, FLUCONAZOLE
FLUOXETINE, FLUOXETINE HYDROCHLORIDE
FOSINOPRIL SODIUM AND HYDROCHLOROTHIAZIDE, FOSINOPRIL SODIUM
FOSINOPRIL SODIUM, FOSINOPRIL SODIUM
GABAPENTIN, GABAPENTIN
GANCICLOVIR, GANCICLOVIR
GLIMEPIRIDE, GLIMEPIRIDE
LORATADINE AND PSEUDOEPHEDRINE SULFATE, LORATADINE (OTC)
LORATADINE, LORATADINE (OTC)
LORAZEPAM, LORAZEPAM
MELOXICAM, MELOXICAM
METFORMIN HYDROCHLORIDE, METFORMIN HYDROCHLORIDE
MIDAZOLAM HYDROCHLORIDE, MIDAZOLAM HYDROCHLORIDE
MINOCYCLINE HYDROCHLORIDE, MINOCYCLINE HYDROCHLORIDE
NEFAZODONE HYDROCHLORIDE, NEFAZODONE HYDROCHLORIDE
NITROFURANTOIN (MONOHYDRATE/MACROCRYSTALS), NITROFURANTOIN
OFLOXACIN, OFLOXACIN
PANIXINE DISPERDOSE, CEPHALEXIN
PSEUDOEPHEDRINE HYDROCHLORIDE, PSEUDOEPHEDRINE HYDROCHLORIDE (OTC)
QUINAPRIL, QUINAPRIL HYDROCHLORIDE
RANICLOR, CEFACLOR
RANITIDINE, RANITIDINE HYDROCHLORIDE
RANITIDINE, RANITIDINE HYDROCHLORIDE (OTC)

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

**** R ****

- * RANBAXY LABORATORIES LTD
SIMVASTATIN, SIMVASTATIN
SOTRET, ISOTRETINOIN
TERAZOSIN HYDROCHLORIDE, TERAZOSIN HYDROCHLORIDE
ZIDOVUDINE, ZIDOVUDINE
- * RANBAXY PHARMACEUTICALS INC
ACETAMINOPHEN AND CODEINE PHOSPHATE, ACETAMINOPHEN
ACETAMINOPHEN W/ CODEINE PHOSPHATE #3, ACETAMINOPHEN
AMOXICILLIN AND CLAVULANATE POTASSIUM, AMOXICILLIN
AMOXICILLIN, AMOXICILLIN
AVENTYL HYDROCHLORIDE, NORTRIPTYLINE HYDROCHLORIDE
CEFACLOX, CEFACLOX
CEFADROXIL, CEFADROXIL/CEFADROXIL HEMIHYDRATE
CIPROFLOXACIN, CIPROFLOXACIN HYDROCHLORIDE
LISINOPRIL AND HYDROCHLOROTHIAZIDE, HYDROCHLOROTHIAZIDE
LISINOPRIL, LISINOPRIL
LOPERAMIDE HYDROCHLORIDE AND SIMETHICONE, LOPERAMIDE HYDROCHLORIDE (OTC)
PENTAZOCINE AND NALOXONE HYDROCHLORIDES, NALOXONE HYDROCHLORIDE
RIOMET, METFORMIN HYDROCHLORIDE
SECONAL SODIUM, SECOBARBITAL SODIUM
SOTRET, ISOTRETINOIN

RATIOPHARM

- * RATIOPHARM GMBH AND CO ARZNEIMITTEL
SUCRALFATE, SUCRALFATE

RECKITT BENCKISER

- * RECKITT BENCKISER PHARMACEUTICALS INC
BUPRENEX, BUPRENORPHINE HYDROCHLORIDE
SUBOXONE, BUPRENORPHINE HYDROCHLORIDE
SUBUTEX, BUPRENORPHINE HYDROCHLORIDE

REAGENT

- * REAGENT MEDICAL AMERICAS LLC
HIBICLENS, CHLORHEXIDINE GLUCONATE (OTC)
HIBISTAT, CHLORHEXIDINE GLUCONATE (OTC)

RELIANT PHARMS

- * RELIANT PHARMACEUTICALS LLC
AXID, NIZATIDINE
DYNACIRC CR, ISRADIPINE
INNOPRAN XL, PROPRANOLOL HYDROCHLORIDE
RYTHMOL SR, PROPAFENONE HYDROCHLORIDE
RYTHMOL, PROPAFENONE HYDROCHLORIDE

RELIANT PHARMS INC

- * RELIANT PHARMACEUTICALS INC
OMACOR, OMEGA-3-ACID ETHYL ESTERS

RESPIRARE

- * RESPIRARE LLC
ALBUTEROL SULFATE, ALBUTEROL SULFATE
CROMOLYN SODIUM, CROMOLYN SODIUM
IPRATROPIUM BROMIDE, IPRATROPIUM BROMIDE

RHODIA

- * RHODIA LTD
ISOFLURANE, ISOFLURANE

ROCHE

- * HOFFMANN LA ROCHE INC
BONIVA, IBANDRONATE SODIUM
BUMEX, BUMETANIDE
COPEGUS, RIBAVIRIN
DEMADEX, TORSEMIDE
FANSIDAR, PYRIMETHAMINE
FUZEON, ENFUVIRTIDE
GANTRISIN PEDIATRIC, SULFISOXAZOLE ACETYL
HIVID, ZALCITABINE

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

**** R ****

* HOFFMANN LA ROCHE INC
INVIRASE, SAQUINAVIR MESYLATE
KLONOPIN RAPIDLY DISINTEGRATING, CLONAZEPAM
KLONOPIN, CLONAZEPAM
KYTRIL, GRANISETRON HYDROCHLORIDE
LARIAM, MEFLUQUINE HYDROCHLORIDE
ROCALTRON, CALCITRIOL
TAMIFLU, OSELTAMIVIR PHOSPHATE
VALIUM, DIAZEPAM
VESANOID, TRETINOIN

ROCHE PALO

* ROCHE PALO ALTO LLC
AEROBID, FLUNISOLIDE
ANAPROX DS, NAPROXEN SODIUM
ANAPROX, NAPROXEN SODIUM
CELLCEPT, MYCOPHENOLATE MOFETIL
CELLCEPT, MYCOPHENOLATE MOFETIL HYDROCHLORIDE
CYTOVENE IV, GANCICLOVIR SODIUM
EC-NAPROSYN, NAPROXEN
NAPROSYN, NAPROXEN
TICLID, TICLOPIDINE HYDROCHLORIDE
TORADOL, KETOROLAC TROMETHAMINE
VALCYTE, VALGANCICLOVIR HYDROCHLORIDE

ROMARK

* ROMARK LABORATORIES
ALINIA, NITAZOXANIDE

ROSEMONT

* ROSEMONT GROUP LTD
SOLTAMOX, TAMOXIFEN CITRATE

ROSS LABS

* ROSS LABORATORIES DIV ABBOTT LABORATORIES INC
PEDIAMYCIN 400, ERYTHROMYCIN ETHYLSUCCINATE
PEDIAMYCIN, ERYTHROMYCIN ETHYLSUCCINATE
PEDIAZOLE, ERYTHROMYCIN ETHYLSUCCINATE
SURVANTA, BERACTANT

ROXANE

* ROXANE LABORATORIES INC
ACETAMINOPHEN W/ CODEINE, ACETAMINOPHEN
ALBUTEROL SULFATE, ALBUTEROL SULFATE
ALPRAZOLAM, ALPRAZOLAM
AMINOPHYLLINE, AMINOPHYLLINE
ANAGRELIDE HYDROCHLORIDE, ANAGRELIDE HYDROCHLORIDE
AZATHIOPRINE, AZATHIOPRINE
BUTORPHANOL TARTRATE, BUTORPHANOL TARTRATE
CALCITRIOL, CALCITRIOL
CHLORPROMAZINE HYDROCHLORIDE INTENSOL, CHLORPROMAZINE HYDROCHLORIDE
CILOSTAZOL, CILOSTAZOL
CITALOPRAM HYDROBROMIDE, CITALOPRAM HYDROBROMIDE
CLARITHROMYCIN, CLARITHROMYCIN
CLOTRIMAZOLE, CLOTRIMAZOLE
CROMOLYN SODIUM, CROMOLYN SODIUM
CYCLOPHOSPHAMIDE, CYCLOPHOSPHAMIDE
DEXAMETHASONE INTENSOL, DEXAMETHASONE
DEXAMETHASONE, DEXAMETHASONE
DIAZEPAM INTENSOL, DIAZEPAM
DIAZEPAM, DIAZEPAM
DICLOFENAC SODIUM, DICLOFENAC SODIUM
DIGOXIN, DIGOXIN
DIPHENOXYLATE HYDROCHLORIDE AND ATROPINE SULFATE, ATROPINE SULFATE
DOLOPHINE HYDROCHLORIDE, METHADONE HYDROCHLORIDE
FLECAINIDE ACETATE, FLECAINIDE ACETATE
FLUCONAZOLE, FLUCONAZOLE

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

** R **

* ROXANE LABORATORIES INC
FLUTICASONE PROPIONATE, FLUTICASONE PROPIONATE
FUROSEMIDE, FUROSEMIDE
HYDROCHLOROTHIAZIDE, HYDROCHLOROTHIAZIDE
HYDROMORPHONE HYDROCHLORIDE, HYDROMORPHONE HYDROCHLORIDE
HYDROXYUREA, HYDROXYUREA
IPRATROPIUM BROMIDE, IPRATROPIUM BROMIDE
LACTULOSE, LACTULOSE
LEUCOVORIN CALCIUM, LEUCOVORIN CALCIUM
LEVORPHANOL TARTRATE, LEVORPHANOL TARTRATE
LIDOCAINE HYDROCHLORIDE, LIDOCAINE HYDROCHLORIDE
LIDOCAINE VISCOUS, LIDOCAINE HYDROCHLORIDE
LITHIUM CARBONATE, LITHIUM CARBONATE
LITHIUM CITRATE, LITHIUM CITRATE
LOPERAMIDE HYDROCHLORIDE, LOPERAMIDE HYDROCHLORIDE (OTC)
LORAZEPAM INTENSOL, LORAZEPAM
LORAZEPAM, LORAZEPAM
MEFLOQUINE HYDROCHLORIDE, MEFLOQUINE HYDROCHLORIDE
MEGESTROL ACETATE, MEGESTROL ACETATE
MELOXICAM, MELOXICAM
MEPERIDINE HYDROCHLORIDE, MEPERIDINE HYDROCHLORIDE
MERCAPTOPYRINE, MERCAPTOPYRINE
METHADONE HYDROCHLORIDE INTENSOL, METHADONE HYDROCHLORIDE
METHADONE HYDROCHLORIDE, METHADONE HYDROCHLORIDE
METHOTREXATE SODIUM, METHOTREXATE SODIUM
METOLAZONE, METOLAZONE
MIDAZOLAM HYDROCHLORIDE, MIDAZOLAM HYDROCHLORIDE
MIRTAZAPINE, MIRTAZAPINE
NAPROXEN, NAPROXEN
ONDANSETRON HYDROCHLORIDE, ONDANSETRON HYDROCHLORIDE
PILOCARPINE HYDROCHLORIDE, PILOCARPINE HYDROCHLORIDE
PREDNISONE INTENSOL, PREDNISONE
PREDNISONE, PREDNISONE
PROPANTHELINE BROMIDE, PROPANTHELINE BROMIDE
PROPRANOLOL HYDROCHLORIDE INTENSOL, PROPRANOLOL HYDROCHLORIDE
PROPRANOLOL HYDROCHLORIDE, PROPRANOLOL HYDROCHLORIDE
ROXICET 5/500, ACETAMINOPHEN
ROXICET, ACETAMINOPHEN
ROXILOX, ACETAMINOPHEN
SERTRALINE HYDROCHLORIDE, SERTRALINE HYDROCHLORIDE
SODIUM POLYSTYRENE SULFONATE, SODIUM POLYSTYRENE SULFONATE
TAMOXIFEN CITRATE, TAMOXIFEN CITRATE
THEOPHYLLINE, THEOPHYLLINE
THIORIDAZINE HYDROCHLORIDE INTENSOL, THIORIDAZINE HYDROCHLORIDE
TORSEMIDE, TORSEMIDE
TRIAZOLAM, TRIAZOLAM
ZIDOVUDINE, ZIDOVUDINE
ZONISAMIDE, ZONISAMIDE

RXELITE

* RXELITE
ALBUTEROL SULFATE, ALBUTEROL SULFATE
FLUOXETINE, FLUOXETINE HYDROCHLORIDE
IPRATROPIUM BROMIDE, IPRATROPIUM BROMIDE

SAGE PRODS

* SAGE PRODUCTS INC
CHLORHEXIDINE GLUCONATE, CHLORHEXIDINE GLUCONATE (OTC)

SALIX PHARMS

* SALIX PHARMACEUTICALS INC
ANUSOL HC, HYDROCORTISONE
COLAZAL, BALSALAZIDE DISODIUM
MOVIPREP, ASCORBIC ACID
OSMOPREP, SODIUM PHOSPHATE, DIBASIC, ANHYDROUS
VISICOL, SODIUM PHOSPHATE, DIBASIC, ANHYDROUS

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

** S **

* SALIX PHARMACEUTICALS INC
XIFAXAN, RIFAXIMIN

SAMSON MEDCL

* SAMSON MEDICAL TECHNOLOGIES LLC
CEFAZOLIN SODIUM, CEFAZOLIN SODIUM

SANDOZ

* SANDOZ CANADA INC
ANECTINE, SUCCINYLCHOLINE CHLORIDE
ARISTOSPAN, TRIAMCINOLONE HEXACETONIDE
DIGOXIN, DIGOXIN
FENOLDOPAM MESYLATE, FENOLDOPAM MESYLATE
FLUMAZENIL, FLUMAZENIL
HALOPERIDOL DECANOATE, HALOPERIDOL DECANOATE
HALOPERIDOL, HALOPERIDOL LACTATE
INFUVITE ADULT, ALPHA-TOCOPHEROL ACETATE
INFUVITE PEDIATRIC (PHARMACY BULK PACKAGE), ASCORBIC ACID
INFUVITE PEDIATRIC, ASCORBIC ACID
ISONIAZID, ISONIAZID
KETOROLAC TROMETHAMINE, KETOROLAC TROMETHAMINE
PROPRANOLOL HYDROCHLORIDE, PROPRANOLOL HYDROCHLORIDE
REGONOL, PYRIDOSTIGMINE BROMIDE
TESTOSTERONE CYPIONATE, TESTOSTERONE CYPIONATE

* SANDOZ INC
ACETAMINOPHEN AND CODEINE PHOSPHATE, ACETAMINOPHEN
ALBUTEROL SULFATE AND IPRATROPIUM BROMIDE, ALBUTEROL SULFATE
ALBUTEROL SULFATE, ALBUTEROL SULFATE
ALLOPURINOL, ALLOPURINOL
ALPRAZOLAM, ALPRAZOLAM
AMANTADINE HYDROCHLORIDE, AMANTADINE HYDROCHLORIDE
AMILORIDE HYDROCHLORIDE AND HYDROCHLOROTHIAZIDE, AMILORIDE HYDROCHLORIDE
AMIODARONE HYDROCHLORIDE, AMIODARONE HYDROCHLORIDE
AMITRIPTYLINE HYDROCHLORIDE, AMITRIPTYLINE HYDROCHLORIDE
AMOXAPINE, AMOXAPINE
AMOXICILLIN AND CLAVULANATE POTASSIUM, AMOXICILLIN
AMOXICILLIN, AMOXICILLIN
AMPICILLIN AND SULBACTAM, AMPICILLIN SODIUM
AMPICILLIN SODIUM, AMPICILLIN SODIUM
ANAGRELIDE HYDROCHLORIDE, ANAGRELIDE HYDROCHLORIDE
ATENOLOL, ATENOLOL
AZITHROMYCIN, AZITHROMYCIN
BENAZEPRIL HYDROCHLORIDE AND HYDROCHLOROTHIAZIDE, BENAZEPRIL HYDROCHLORIDE
BENAZEPRIL HYDROCHLORIDE, BENAZEPRIL HYDROCHLORIDE
BISOPROLOL FUMARATE AND HYDROCHLOROTHIAZIDE, BISOPROLOL FUMARATE
BISOPROLOL FUMARATE, BISOPROLOL FUMARATE
BUMETANIDE, BUMETANIDE
BUPROPION HYDROCHLORIDE, BUPROPION HYDROCHLORIDE
BUSPIRONE HYDROCHLORIDE, BUSPIRONE HYDROCHLORIDE
BUTAL COMPOUND, ASPIRIN
CAFERGOT, CAFFEINE
CAPTOPRIL, CAPTOPRIL
CARBIDOPA AND LEVODOPA, CARBIDOPA
CARBOPLATIN, CARBOPLATIN
CARISOPRODOL AND ASPIRIN, ASPIRIN
CARISOPRODOL, CARISOPRODOL
CARISOPRODOL, ASPIRIN AND CODEINE PHOSPHATE, ASPIRIN
CEFADROXIL, CEFADROXIL/CEFADROXIL HEMIHYDRATE
CEFPROZIL, CEFPROZIL
CEFTRIAZONE, CEFTRIAZONE SODIUM
CEFUROXIME AXETIL, CEFUROXIME AXETIL
CHLORPROMAZINE HYDROCHLORIDE, CHLORPROMAZINE HYDROCHLORIDE
CHLORPROPAMIDE, CHLORPROPAMIDE
CHLORZOXAZONE, CHLORZOXAZONE
CHOLESTYRAMINE LIGHT, CHOLESTYRAMINE

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

**** S ****

* SANDOZ INC
CHOLESTYRAMINE, CHOLESTYRAMINE
CILOSTAZOL, CILOSTAZOL
CIMETIDINE, CIMETIDINE
CIPROFLOXACIN, CIPROFLOXACIN HYDROCHLORIDE
CITALOPRAM HYDROBROMIDE, CITALOPRAM HYDROBROMIDE
CLARITHROMYCIN EXTENDED RELEASE, CLARITHROMYCIN
CLARITHROMYCIN, CLARITHROMYCIN
CLEMASTINE FUMARATE, CLEMASTINE FUMARATE
CLEMASTINE FUMARATE, CLEMASTINE FUMARATE (OTC)
CLOMIPRAMINE HYDROCHLORIDE, CLOMIPRAMINE HYDROCHLORIDE
CLONAZEPAM, CLONAZEPAM
CLONIDINE HYDROCHLORIDE, CLONIDINE HYDROCHLORIDE
CLORAZEPATE DIPOTASSIUM, CLORAZEPATE DIPOTASSIUM
CORPHED, PSEUDOEPHEDRINE HYDROCHLORIDE
CYCLOBENZAPRINE HYDROCHLORIDE, CYCLOBENZAPRINE HYDROCHLORIDE
CYCLOSPORINE, CYCLOSPORINE
DESIPRAMINE HYDROCHLORIDE, DESIPRAMINE HYDROCHLORIDE
DEXTROAMP SACCHARATE, AMP ASPARTATE, DEXTROAMP SULFATE AND AMP SULFATE, AMPHETAMINE
ASPARTATE
DIAZEPAM, DIAZEPAM
DICLOFENAC POTASSIUM, DICLOFENAC POTASSIUM
DICLOFENAC SODIUM, DICLOFENAC SODIUM
DICLOXACILLIN SODIUM, DICLOXACILLIN SODIUM
DIFLUNISAL, DIFLUNISAL
DIPHENHYDRAMINE HYDROCHLORIDE, DIPHENHYDRAMINE HYDROCHLORIDE
DIPHENOXYLATE HYDROCHLORIDE W/ ATROPINE SULFATE, ATROPINE SULFATE
DIPYRIDAMOLE, DIPYRIDAMOLE
DOXAZOSIN MESYLATE, DOXAZOSIN MESYLATE
DOXEPIN HYDROCHLORIDE, DOXEPIN HYDROCHLORIDE
DOXYCYCLINE HYCLATE, DOXYCYCLINE HYCLATE
DOXYCYCLINE, DOXYCYCLINE
ENALAPRIL MALEATE AND HYDROCHLOROTHIAZIDE, ENALAPRIL MALEATE
ENALAPRIL MALEATE, ENALAPRIL MALEATE
ETODOLAC, ETODOLAC
FAMOTIDINE, FAMOTIDINE
FAMOTIDINE, FAMOTIDINE (OTC)
FENOPROFEN CALCIUM, FENOPROFEN CALCIUM
FLECAINIDE ACETATE, FLECAINIDE ACETATE
FLUCONAZOLE, FLUCONAZOLE
FLUOXETINE HYDROCHLORIDE, FLUOXETINE HYDROCHLORIDE
FLUOXETINE, FLUOXETINE HYDROCHLORIDE
FLUPHENAZINE HYDROCHLORIDE, FLUPHENAZINE HYDROCHLORIDE
FLURAZEPAM HYDROCHLORIDE, FLURAZEPAM HYDROCHLORIDE
FLURBIPROFEN, FLURBIPROFEN
FLUTAMIDE, FLUTAMIDE
FLUVOXAMINE MALEATE, FLUVOXAMINE MALEATE
FOSINOPRIL SODIUM AND HYDROCHLOROTHIAZIDE, FOSINOPRIL SODIUM
FOSINOPRIL SODIUM, FOSINOPRIL SODIUM
FUROSEMIDE, FUROSEMIDE
GABAPENTIN, GABAPENTIN
GEMFIBROZIL, GEMFIBROZIL
GLIPIZIDE, GLIPIZIDE
GLYBURIDE (MICRONIZED), GLYBURIDE
GUANABENZ ACETATE, GUANABENZ ACETATE
HALOPERIDOL, HALOPERIDOL
HYDRALAZINE HYDROCHLORIDE, HYDRALAZINE HYDROCHLORIDE
HYDROCHLOROTHIAZIDE, HYDROCHLOROTHIAZIDE
HYDROCODONE BITARTRATE AND ACETAMINOPHEN, ACETAMINOPHEN
HYDROXYCHLOROQUINE SULFATE, HYDROXYCHLOROQUINE SULFATE
HYDROXYZINE HYDROCHLORIDE, HYDROXYZINE HYDROCHLORIDE
HYDROXYZINE PAMOATE, HYDROXYZINE PAMOATE
IBUPROFEN, IBUPROFEN
IBUPROFEN, IBUPROFEN (OTC)

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

**** S ****

* SANDOZ INC
IMIPRAMINE HYDROCHLORIDE, IMIPRAMINE HYDROCHLORIDE
INDAPAMIDE, INDAPAMIDE
INDOCIN SR, INDOMETHACIN
INVAGESIC FORTE, ASPIRIN
INVAGESIC, ASPIRIN
ISONIAZID, ISONIAZID
ISOSORBIDE DINITRATE, ISOSORBIDE DINITRATE
ITRACONAZOLE, ITRACONAZOLE
KETOPROFEN, KETOPROFEN
LABETALOL HYDROCHLORIDE, LABETALOL HYDROCHLORIDE
LEFLUNOMIDE, LEFLUNOMIDE
LEUCOVORIN CALCIUM, LEUCOVORIN CALCIUM
LISINOPRIL AND HYDROCHLOROTHIAZIDE, HYDROCHLOROTHIAZIDE
LISINOPRIL, LISINOPRIL
LOCHOLEST LIGHT, CHOLESTYRAMINE
LOCHOLEST, CHOLESTYRAMINE
LONOX, ATROPINE SULFATE
LOPERAMIDE HYDROCHLORIDE, LOPERAMIDE HYDROCHLORIDE
LORATADINE, LORATADINE (OTC)
LORAZEPAM, LORAZEPAM
LOVASTATIN, LOVASTATIN
MECLIZINE HYDROCHLORIDE, MECLIZINE HYDROCHLORIDE
MECLOFENAMATE SODIUM, MECLOFENAMATE SODIUM
MEFLOQUINE HYDROCHLORIDE, MEFLOQUINE HYDROCHLORIDE
METFORMIN HYDROCHLORIDE, METFORMIN HYDROCHLORIDE
METHADONE HYDROCHLORIDE, METHADONE HYDROCHLORIDE
METHAZOLAMIDE, METHAZOLAMIDE
METHIMAZOLE, METHIMAZOLE
METHOCARBAMOL, METHOCARBAMOL
METHYLCLOTHIAZIDE, METHYLCLOTHIAZIDE
METHYLDOPA, METHYLDOPA
METHYLPREDNISOLONE, METHYLPREDNISOLONE
METOCLOPRAMIDE HYDROCHLORIDE, METOCLOPRAMIDE HYDROCHLORIDE
METOLAZONE, METOLAZONE
METOPROLOL SUCCINATE, METOPROLOL SUCCINATE
METOPROLOL TARTRATE, METOPROLOL TARTRATE
METRONIDAZOLE, METRONIDAZOLE
MEXILETINE HYDROCHLORIDE, MEXILETINE HYDROCHLORIDE
MIDODRINE HYDROCHLORIDE, MIDODRINE HYDROCHLORIDE
MIRTAZAPINE, MIRTAZAPINE
NABUMETONE, NABUMETONE
NADOLOL, NADOLOL
NAFCILLIN SODIUM, NAFCILLIN SODIUM
NALTREXONE HYDROCHLORIDE, NALTREXONE HYDROCHLORIDE
NAPROXEN SODIUM, NAPROXEN SODIUM
NAPROXEN SODIUM, NAPROXEN SODIUM (OTC)
NAPROXEN, NAPROXEN
NEFAZODONE HYDROCHLORIDE, NEFAZODONE HYDROCHLORIDE
NITROFURANTOIN (MONOHYDRATE/MACROCRYSTALS), NITROFURANTOIN
NITROFURANTOIN, NITROFURANTOIN, MACROCRYSTALLINE
NIZATIDINE, NIZATIDINE
NORTRIPTYLINE HYDROCHLORIDE, NORTRIPTYLINE HYDROCHLORIDE
NYDRAZID, ISONIAZID
OMNITROPE, SOMATROPIN RECOMBINANT
ORPHENADRINE CITRATE, ORPHENADRINE CITRATE
ORPHENADRINE CITRATE, ASPIRIN, AND CAFFEINE, ASPIRIN
OXACILLIN SODIUM, OXACILLIN SODIUM
OXANDROLONE, OXANDROLONE
OXAPROZIN, OXAPROZIN
OXAZEPAM, OXAZEPAM
PAROXETINE HYDROCHLORIDE, PAROXETINE HYDROCHLORIDE
PENICILLIN G POTASSIUM, PENICILLIN G POTASSIUM
PERPHENAZINE, PERPHENAZINE

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

** S **

* SANDOZ INC
 PHENDIMETRAZINE TARTRATE, PHENDIMETRAZINE TARTRATE
 PHENTERMINE HYDROCHLORIDE, PHENTERMINE HYDROCHLORIDE
 PINDOLOL, PINDOLOL
 PRAZOSIN HYDROCHLORIDE, PRAZOSIN HYDROCHLORIDE
 PREDNISONE, PREDNISONE
 PROCHLORPERAZINE MALEATE, PROCHLORPERAZINE MALEATE
 PROMETHAZINE HYDROCHLORIDE, PROMETHAZINE HYDROCHLORIDE
 PROPOXYPHENE HYDROCHLORIDE AND ACETAMINOPHEN, ACETAMINOPHEN
 PROPOXYPHENE NAPSYLATE AND ACETAMINOPHEN, ACETAMINOPHEN
 PROPRANOLOL HYDROCHLORIDE AND HYDROCHLOROTHIAZIDE, HYDROCHLOROTHIAZIDE
 PROPRANOLOL HYDROCHLORIDE, PROPRANOLOL HYDROCHLORIDE
 PSEUDOEPHEDRINE HYDROCHLORIDE AND TRIPROLIDINE HYDROCHLORIDE, PSEUDOEPHEDRINE
 HYDROCHLORIDE
 QUINAPRIL HYDROCHLORIDE, QUINAPRIL HYDROCHLORIDE
 QUINIDINE SULFATE, QUINIDINE SULFATE
 RANITIDINE HYDROCHLORIDE, RANITIDINE HYDROCHLORIDE
 RANITIDINE, RANITIDINE HYDROCHLORIDE (OTC)
 RESERPINE, RESERPINE
 RIBAVIRIN, RIBAVIRIN
 RIFAMPIN, RIFAMPIN
 SIMVASTATIN, SIMVASTATIN
 SONAZINE, CHLORPROMAZINE HYDROCHLORIDE
 SOTALOL HYDROCHLORIDE, SOTALOL HYDROCHLORIDE
 SPIRONOLACTONE AND HYDROCHLOROTHIAZIDE, HYDROCHLOROTHIAZIDE
 SPIRONOLACTONE, SPIRONOLACTONE
 SULFADIAZINE, SULFADIAZINE
 SULFAMETHOXAZOLE AND TRIMETHOPRIM DOUBLE STRENGTH, SULFAMETHOXAZOLE
 SULFAMETHOXAZOLE AND TRIMETHOPRIM, SULFAMETHOXAZOLE
 SULINDAC, SULINDAC
 TEMAZEPAM, TEMAZEPAM
 TERAZOSIN HYDROCHLORIDE, TERAZOSIN HYDROCHLORIDE
 THIORIDAZINE HYDROCHLORIDE, THIORIDAZINE HYDROCHLORIDE
 THIOTHIXENE, THIOTHIXENE
 TICLOPIDINE HYDROCHLORIDE, TICLOPIDINE HYDROCHLORIDE
 TIMOLOL MALEATE, TIMOLOL MALEATE
 TIZANIDINE HYDROCHLORIDE, TIZANIDINE HYDROCHLORIDE
 TOLAZAMIDE, TOLAZAMIDE
 TOLBUTAMIDE, TOLBUTAMIDE
 TOLMETIN SODIUM, TOLMETIN SODIUM
 TRAMADOL HYDROCHLORIDE, TRAMADOL HYDROCHLORIDE
 TRAZODONE HYDROCHLORIDE, TRAZODONE HYDROCHLORIDE
 TRIAMTERENE AND HYDROCHLOROTHIAZIDE, HYDROCHLOROTHIAZIDE
 TRIFLUOPERAZINE HYDROCHLORIDE, TRIFLUOPERAZINE HYDROCHLORIDE
 VERAPAMIL HYDROCHLORIDE, VERAPAMIL HYDROCHLORIDE
 WARFARIN SODIUM, WARFARIN SODIUM
 ZONISAMIDE, ZONISAMIDE

SANOFI AVENTIS US

* SANOFI AVENTIS US LLC
 ALLEGRA D 24 HOUR, FEXOFENADINE HYDROCHLORIDE
 ALLEGRA, FEXOFENADINE HYDROCHLORIDE
 ALLEGRA-D 12 HOUR, FEXOFENADINE HYDROCHLORIDE
 AMARYL, GLIMEPIRIDE
 AMBIEN CR, ZOLPIDEM TARTRATE
 AMBIEN, ZOLPIDEM TARTRATE
 ANZEMET, DOLASETRON MESYLATE
 APIDRA, INSULIN GLULISINE RECOMBINANT
 ARALEN, CHLOROQUINE PHOSPHATE
 ARAVA, LEFLUNOMIDE
 AVALIDE, HYDROCHLOROTHIAZIDE
 AVAPRO, IRBESARTAN
 BENZACLIN, BENZOYL PEROXIDE
 BENZAMYCIN PAK, BENZOYL PEROXIDE
 BENZAMYCIN, BENZOYL PEROXIDE

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

**** S ****

* SANOFI AVENTIS US LLC
CANTIL, MEPENZOLATE BROMIDE
CARAC, FLUOROURACIL
CLAFORAN IN DEXTROSE 5% IN PLASTIC CONTAINER, CEFOTAXIME SODIUM
CLAFORAN, CEFOTAXIME SODIUM
CLOMID, CLOMIPHENE CITRATE
DDAVP (NEEDS NO REFRIGERATION), DESMOPRESSIN ACETATE
DDAVP, DESMOPRESSIN ACETATE
DEMEROL, MEPERIDINE HYDROCHLORIDE
DERMATOP E EMOLLIENT, PREDNICARBATE
DERMATOP, PREDNICARBATE
DIABETA, GLYBURIDE
DRISDOL, ERGOCALCIFEROL
ELOXATIN, OXALIPLATIN
GAVISCON, ALUMINUM HYDROXIDE (OTC)
HIPREX, METHENAMINE HIPPURATE
HYTONE, HYDROCORTISONE
KAYEXALATE, SODIUM POLYSTYRENE SULFONATE
KERLONE, BETAXOLOL HYDROCHLORIDE
KETEK, TELITHROMYCIN
KLARON, SULFACETAMIDE SODIUM
LANTUS, INSULIN GLARGINE RECOMBINANT
LASIX, FUROSEMIDE
LOVENOX (PRESERVATIVE FREE), ENOXAPARIN SODIUM
LOVENOX, ENOXAPARIN SODIUM
LOZOL, INDAPAMIDE
MYTELASE, AMBENONIUM CHLORIDE
NASACORT AQ, TRIAMCINOLONE ACETONIDE
NEGGRAM, NALIDIXIC ACID
NICODERM CQ, NICOTINE (OTC)
NILANDRON, NILUTAMIDE
NORITATE, METRONIDAZOLE
NORPRAMIN, DESIPRAMINE HYDROCHLORIDE
PENLAC, CICLOPIROX
PHISOHEX, HEXACHLOROPHENE
PLAQUENIL, HYDROXYCHLOROQUINE SULFATE
PLAVIX, CLOPIDOGREL BISULFATE
PRIFTIN, RIFAPENTINE
PRIMACOR IN DEXTROSE 5% IN PLASTIC CONTAINER, MILRINONE LACTATE
PRIMACOR, MILRINONE LACTATE
PRIMAQUINE, PRIMAQUINE PHOSPHATE
PSORCON, DIFLORASONE DIACETATE
RIFADIN, RIFAMPIN
RIFAMATE, ISONIAZID
RIFATER, ISONIAZID
RILUTEK, RILUZOLE
SKELID, TILUDRONATE DISODIUM
TALACEN, ACETAMINOPHEN
TALWIN NX, NALOXONE HYDROCHLORIDE
TAXOTERE, DOCETAXEL
TRENTAL, PENTOXIFYLLINE
UROXATRAL, ALFUZOSIN HYDROCHLORIDE

SANTARUS

* SANTARUS INC
ZEGERID, MAGNESIUM HYDROXIDE
ZEGERID, OMEPRAZOLE

SANTEN

* SANTEN INC
ALAMAST, PEMIROLAST POTASSIUM
IQUIX, LEVOFLOXACIN
QUIXIN, LEVOFLOXACIN

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

** S **

SANTEN OY

- * SANTEN OY
BETIMOL, TIMOLOL

SAVAGE LABS

- * SAVAGE LABORATORIES INC DIV ALTANA INC
ETHIODOL, ETHIODIZED OIL
KAON CL-10, POTASSIUM CHLORIDE
PANDEL, HYDROCORTISONE PROBUTATE

SAVIENT PHARMS

- * SAVIENT PHARMACEUTICALS INC
OXANDRIN, OXANDROLONE

SB PHARMCO

- * SB PHARMCO PUERTO RICO INC
AVANDAMET, METFORMIN HYDROCHLORIDE
AVANDARYL, GLIMEPIRIDE
AVANDIA, ROSIGLITAZONE MALEATE
COREG CR, CARVEDILOL PHOSPHATE

SCHERER RP

- * RP SCHERER CORP
NIFEDIPINE, NIFEDIPINE
- * RP SCHERER NORTH AMERICA
NIFEDIPINE, NIFEDIPINE
VALPROIC ACID, VALPROIC ACID
- * RP SCHERER NORTH AMERICA DIV RP SCHERER CORP
DIPHENOXYLATE HYDROCHLORIDE W/ ATROPINE SULFATE, ATROPINE SULFATE

SCHERING

- * SCHERING
ASMANEX TWISTHALER, MOMETASONE FUROATE
CLARINEX, DESLORATADINE
CLARINEX-D 12 HOUR, DESLORATADINE
- * SCHERING CORP
CLARINEX D 24 HOUR, DESLORATADINE
CLARINEX, DESLORATADINE
DIPROLENE AF, BETAMETHASONE DIPROPIONATE
GUANIDINE HYDROCHLORIDE, GUANIDINE HYDROCHLORIDE
INTEGRILIN, EPTIFIBATIDE
LOTRISONE, BETAMETHASONE DIPROPIONATE
NOXAFIL, POSACONAZOLE
REBETOL, RIBAVIRIN
TEMODAR, TEMOZOLOMIDE
- * SCHERING CORP SUB SCHERING PLOUGH CORP
CELESTONE SOLUSPAN, BETAMETHASONE ACETATE
CELESTONE, BETAMETHASONE
DIPROLENE, BETAMETHASONE DIPROPIONATE
ELOCON, MOMETASONE FUROATE
FULVICIN-U/F, GRISEOFULVIN, MICROCRYSTALLINE
PROVENTIL, ALBUTEROL

SCHERING PLOUGH

- * SCHERING PLOUGH CORP
CLARINEX, DESLORATADINE
IMDUR, ISOSORBIDE MONONITRATE
- * SCHERING PLOUGH HEALTHCARE PRODUCTS INC
AFRINOL, PSEUDOEPHEDRINE SULFATE (OTC)
CHILDREN'S CLARITIN, LORATADINE (OTC)
CHLOR-TRIMETON, CHLORPHENIRAMINE MALEATE (OTC)
CLARITIN HIVES RELIEF REDITAB, LORATADINE (OTC)
CLARITIN HIVES RELIEF, LORATADINE (OTC)
CLARITIN REDITABS, LORATADINE (OTC)
CLARITIN, LORATADINE (OTC)
CLARITIN-D 24 HOUR, LORATADINE (OTC)
DISOPHROL, DEXBROMPHENIRAMINE MALEATE (OTC)
DRIXORAL PLUS, ACETAMINOPHEN (OTC)

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

** S **

- * SCHERING PLOUGH HEALTHCARE PRODUCTS INC
DRIXORAL, DEXBROMPHENIRAMINE MALEATE (OTC)
GYNE-LOTTRIMIN 3 COMBINATION PACK, CLOTRIMAZOLE (OTC)
GYNE-LOTTRIMIN 3, CLOTRIMAZOLE (OTC)
GYNE-LOTTRIMIN COMBINATION PACK, CLOTRIMAZOLE (OTC)
GYNE-LOTTRIMIN, CLOTRIMAZOLE (OTC)
LOTTRIMIN ULTRA, BUTENAFINE HYDROCHLORIDE (OTC)
MIRALAX, POLYETHYLENE GLYCOL 3350 (OTC)
NASONEX, MOMETASONE FUROATE MONOHYDRATE
OCUCLEAR, OXYMETAZOLINE HYDROCHLORIDE (OTC)
SHADE UVAGUARD, AVOBENZONE (OTC)

SCHERING PLOUGH RES

- * SCHERING PLOUGH RESEARCH INSTITUTE
LOTTRISONE, BETAMETHASONE DIPROPIONATE
REBETOL, RIBAVIRIN

SCHWARZ

- * SCHWARZ GMBH
MONOKET, ISOSORBIDE MONONITRATE

SCHWARZ PHARMA

- * SCHWARZ PHARMA AG
EDEX, ALPROSTADIL
GLYCOLAX, POLYETHYLENE GLYCOL 3350
- * SCHWARZ PHARMA INC
AZDONE, ASPIRIN
CO-GESIC, ACETAMINOPHEN
COLYTE WITH FLAVOR PACKS, POLYETHYLENE GLYCOL 3350
COLYTE, POLYETHYLENE GLYCOL 3350
COLYTE-FLAVORED, POLYETHYLENE GLYCOL 3350
CORTIFOAM, HYDROCORTISONE ACETATE
DILATRATE-SR, ISOSORBIDE DINITRATE
EPIFOAM, HYDROCORTISONE ACETATE
FLUXID, FAMOTIDINE
KEMSTRO, BACLOFEN
LEVATOL, PENBUTOLOL SULFATE
NIRAVAM, ALPRAZOLAM
PARCOPA, CARBIDOPA
PROCTOFOAM HC, HYDROCORTISONE ACETATE
REGLAN, METOCLOPRAMIDE HYDROCHLORIDE
ROBAXIN, METHOCARBAMOL
ROBAXIN-750, METHOCARBAMOL
TRILYTE, POLYETHYLENE GLYCOL 3350
UNIRETIC, HYDROCHLOROTHIAZIDE
UNIVASC, MOEXIPRIL HYDROCHLORIDE

SCIELE PHARMA INC

- * SCIELE PHARMA INC
COGNEX, TACRINE HYDROCHLORIDE
FURADANTIN, NITROFURANTOIN
PONSTEL, MEFENAMIC ACID
ROBINUL FORTE, GLYCOPYRROLATE
ROBINUL, GLYCOPYRROLATE
SULAR, NISOLDIPINE

SCIOS

- * SCIOS INC
NATRECOR, NESIRITIDE RECOMBINANT

SCS

- * SCS PHARMACEUTICALS
ATENOLOL, ATENOLOL

SEPRACOR

- * SEPRACOR INC
BROVANA, ARFORMOTEROL TARTRATE
XOPENEX, LEVALBUTEROL HYDROCHLORIDE

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

**** S ****

* SEPRACOR PHARMACEUTICALS
LUNESTA, ESZOPICLONE
XOPENEX HFA, LEVALBUTEROL TARTRATE

SEPTODONT

* SEPTODONT INC
BUPIVACAINE HYDROCHLORIDE AND EPINEPHRINE, BUPIVACAINE HYDROCHLORIDE
OCTOCAINE, EPINEPHRINE

SERONO

* SERONO LABORATORIES INC
GEREF, SERMORELIN ACETATE
SAIZEN, SOMATROPIN RECOMBINANT
SEROPHENE, CLOMIPHENE CITRATE
SEROSTIM, SOMATROPIN RECOMBINANT

SERONO INC

* SERONO INC
CETROTIDE, CETRORELIX
GONAL-F RFF PEN, FOLLITROPIN ALFA/BETA
GONAL-F RFF, FOLLITROPIN ALFA/BETA
GONAL-F, FOLLITROPIN ALFA/BETA
LUVERIS, LUTROPIN ALFA
NOVANTRONE, MITOXANTRONE HYDROCHLORIDE
OVIDREL, CHORIOGONADOTROPIN ALFA
ZORBTIVE, SOMATROPIN RECOMBINANT

SHIONOGI

* SHIONOGI USA INC
CEDAX, CEFTIBUTEN DIHYDRATE

SHIRE

* SHIRE DEVELOPMENT INC
ADDERALL 10, AMPHETAMINE ASPARTATE
ADDERALL 12.5, AMPHETAMINE ASPARTATE
ADDERALL 15, AMPHETAMINE ASPARTATE
ADDERALL 20, AMPHETAMINE ASPARTATE
ADDERALL 30, AMPHETAMINE ASPARTATE
ADDERALL 5, AMPHETAMINE ASPARTATE
ADDERALL 7.5, AMPHETAMINE ASPARTATE
ADDERALL XR 10, AMPHETAMINE ASPARTATE
ADDERALL XR 15, AMPHETAMINE ASPARTATE
ADDERALL XR 20, AMPHETAMINE ASPARTATE
ADDERALL XR 25, AMPHETAMINE ASPARTATE
ADDERALL XR 30, AMPHETAMINE ASPARTATE
ADDERALL XR 5, AMPHETAMINE ASPARTATE
AGRYLIN, ANAGRELIDE HYDROCHLORIDE
CARBATROL, CARBAMAZEPINE
DAYTRANA, METHYLPHENIDATE
EQUETRO, CARBAMAZEPINE
ETHMOZINE, MORICIZINE HYDROCHLORIDE
FOSRENOL, LANTHANUM CARBONATE
PENTASA, MESALAMINE
PROAMATINE, MIDODRINE HYDROCHLORIDE
PRO-BANTHINE, PROPANTHELINE BROMIDE
SALURON, HYDROFLUMETHIAZIDE

SHIRE RICHWOOD

* SHIRE RICHWOOD INC
DEXTROSTAT, DEXTROAMPHETAMINE SULFATE
X-TROZINE L.A., PHENDIMETRAZINE TARTRATE
X-TROZINE, PHENDIMETRAZINE TARTRATE

SICOR PHARMS

* SICOR PHARMACEUTICALS INC
ACYCLOVIR SODIUM, ACYCLOVIR SODIUM
ACYCLOVIR, ACYCLOVIR SODIUM
ADENOSINE, ADENOSINE

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

** S **

* SICOR PHARMACEUTICALS INC
 ALPROSTADIL, ALPROSTADIL
 AMIKACIN SULFATE, AMIKACIN SULFATE
 AMINOPHYLLINE, AMINOPHYLLINE
 AMIODARONE HYDROCHLORIDE, AMIODARONE HYDROCHLORIDE
 AMPHOTERICIN B, AMPHOTERICIN B
 ATRACURIUM BESYLATE, ATRACURIUM BESYLATE
 AZITHROMYCIN, AZITHROMYCIN HYDROGENCITRATE
 BLEOMYCIN, BLEOMYCIN SULFATE
 BUMETANIDE, BUMETANIDE
 CALCITRIOL, CALCITRIOL
 CARBOPLATIN, CARBOPLATIN
 CIMETIDINE HYDROCHLORIDE, CIMETIDINE HYDROCHLORIDE
 CIPROFLOXACIN, CIPROFLOXACIN
 CISPLATIN, CISPLATIN
 CYTOSAR-U, CYTARABINE
 DACARBAZINE, DACARBAZINE
 DAUNORUBICIN HYDROCHLORIDE, DAUNORUBICIN HYDROCHLORIDE
 DEFEROXAMINE MESYLATE, DEFEROXAMINE MESYLATE
 DESMOPRESSIN ACETATE, DESMOPRESSIN ACETATE
 DEXAMETHASONE SODIUM PHOSPHATE, DEXAMETHASONE SODIUM PHOSPHATE
 DILTIAZEM HYDROCHLORIDE, DILTIAZEM HYDROCHLORIDE
 DIPYRIDAMOLE, DIPYRIDAMOLE
 DOBUTAMINE HYDROCHLORIDE, DOBUTAMINE HYDROCHLORIDE
 DOXORUBICIN HYDROCHLORIDE, DOXORUBICIN HYDROCHLORIDE
 ENALAPRILAT, ENALAPRILAT
 ERYTHROMYCIN LACTOBIONATE, ERYTHROMYCIN LACTOBIONATE
 ETOPOSIDE, ETOPOSIDE
 FLUCONAZOLE IN SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER, FLUCONAZOLE
 FLUCONAZOLE IN SODIUM CHLORIDE 0.9%, FLUCONAZOLE
 FLUDARABINE PHOSPHATE, FLUDARABINE PHOSPHATE
 FLUMAZENIL, FLUMAZENIL
 FLUOROURACIL, FLUOROURACIL
 FLUPHENAZINE DECANOATE, FLUPHENAZINE DECANOATE
 HALOPERIDOL DECANOATE, HALOPERIDOL DECANOATE
 HALOPERIDOL, HALOPERIDOL LACTATE
 IDARUBICIN HYDROCHLORIDE PFS, IDARUBICIN HYDROCHLORIDE
 IDARUBICIN HYDROCHLORIDE, IDARUBICIN HYDROCHLORIDE
 IFOSFAMIDE/MESNA KIT, IFOSFAMIDE
 LEUCOVORIN CALCIUM PRESERVATIVE FREE, LEUCOVORIN CALCIUM
 LEUCOVORIN CALCIUM, LEUCOVORIN CALCIUM
 LEUPROLIDE ACETATE, LEUPROLIDE ACETATE
 LEVOCARNITINE, LEVOCARNITINE
 MEDROXYPROGESTERONE ACETATE, MEDROXYPROGESTERONE ACETATE
 MESNA, MESNA
 METHYLDOPATE HYDROCHLORIDE, METHYLDOPATE HYDROCHLORIDE
 METHYLPREDNISOLONE ACETATE, METHYLPREDNISOLONE ACETATE
 METHYLPREDNISOLONE SODIUM SUCCINATE, METHYLPREDNISOLONE SODIUM SUCCINATE
 METOCLOPRAMIDE HYDROCHLORIDE, METOCLOPRAMIDE HYDROCHLORIDE
 MITOXANTRONE, MITOXANTRONE HYDROCHLORIDE
 NEOSAR, CYCLOPHOSPHAMIDE
 NOREPINEPHRINE BITARTRATE, NOREPINEPHRINE BITARTRATE
 OCTREOTIDE ACETATE, OCTREOTIDE ACETATE
 ONDANSETRON HYDROCHLORIDE AND DEXTROSE IN PLASTIC CONTAINER, ONDANSETRON HYDROCHLORIDE
 PACLITAXEL, PACLITAXEL
 PAMIDRONATE DISODIUM, PAMIDRONATE DISODIUM
 PANCURONIUM BROMIDE, PANCURONIUM BROMIDE
 PROCHLORPERAZINE EDISYLATE, PROCHLORPERAZINE EDISYLATE
 PROMETHAZINE HYDROCHLORIDE, PROMETHAZINE HYDROCHLORIDE
 PROPOFOL, PROPOFOL
 SODIUM NITROPRUSSIDE, SODIUM NITROPRUSSIDE
 SULFAMETHOXAZOLE AND TRIMETHOPRIM, SULFAMETHOXAZOLE
 TERBUTALINE SULFATE, TERBUTALINE SULFATE
 THIOTEPA, THIOTEPA

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

**** S ****

* SICOR PHARMACEUTICALS INC
TOBRAMYCIN SULFATE, TOBRAMYCIN SULFATE
VECURONIUM BROMIDE, VECURONIUM BROMIDE
VINCRIStINE SULFATE PFS, VINCRIStINE SULFATE
VINORELBINE TARTRATE, VINORELBINE TARTRATE
ZANOSAR, STREPTOZOcIN

SIEGFRIED

* SIEGFRIED LTD
SELEGILINE HYDROCHLORIDE, SELEGILINE HYDROCHLORIDE

SIGHT PHARMS

* SIGHT PHARMACEUTICALS INC
MINOXIDIL (FOR MEN), MINOXIDIL (OTC)
MINOXIDIL (FOR WOMEN), MINOXIDIL (OTC)

SIGMA TAU

* SIGMA TAU PHARMACEUTICALS INC
CARNITOR, LEVOCARNITINE
MATULANE, PROCARBAZINE HYDROCHLORIDE

SILARX

* SILARX PHARMACEUTICALS INC
AMANTADINE HYDROCHLORIDE, AMANTADINE HYDROCHLORIDE
CITALOPRAM HYDROBROMIDE, CITALOPRAM HYDROBROMIDE
CLEMAStINE FUMARATE, CLEMAStINE FUMARATE
DOXEPIN HYDROCHLORIDE, DOXEPIN HYDROCHLORIDE
HALOPERIDOL, HALOPERIDOL LACTATE
LORATADINE, LORATADINE (OTC)
METAPROTERENOL SULFATE, METAPROTERENOL SULFATE
METOCLOPRAMIDE HYDROCHLORIDE, METOCLOPRAMIDE HYDROCHLORIDE
OXYBUTYNIN CHLORIDE, OXYBUTYNIN CHLORIDE

SIRIUS LABS

* SIRIUS LABORATORIES INC
TEXACORT, HYDROCORTISONE

SKINMEDICA

* SKINMEDICA INC
DESONATE, DESONIDE
VANIQA, EFLORNITHINE HYDROCHLORIDE

SKYEPHARMA

* SKYEPHARMA INC
DEPOCYT, CYTARABINE
DEPODUR, MORPHINE SULFATE
TRIGLIDE, FENOFIBRATE

SMITHKLINE BEECHAM

* SMITHKLINE BEECHAM CORP DBA GLAXOSMITHKLINE
ARRANON, NELARABINE
COREG, CARVEDILOL
DEXEDRINE, DEXTROAMPHETAMINE SULFATE
EPZICOM, ABACAVIR SULFATE
LANOXICAPS, DIGOXIN
LANOXIN, DIGOXIN
LEUKERAN, CHLORAMBUCIL
WELLBUTRIN XL, BUPROPION HYDROCHLORIDE

SOAPCO

* SOAPCO INC
BRIAN CARE, CHLORHEXIDINE GLUCONATE (OTC)

SOLVAY

* SOLVAY PHARMACEUTICALS
CORTENEMA, HYDROCORTISONE

SOLVAY PHARMS

* SOLVAY PHARMACEUTICALS INC
ACEON, PERINDOPRIL ERBUMINE

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

** S **

SOMERSET

- * SOMERSET PHARMACEUTICALS INC
ELDEPRYL, SELEGILINE HYDROCHLORIDE
EMSAM, SELEGILINE

SPEAR PHARMS

- * SPEAR PHARMACEUTICALS INC
TRETINOIN, TRETINOIN

SPECTRUM PHARMS

- * SPECTRUM PHARMACEUTICALS INC
CARBOPLATIN, CARBOPLATIN

STADA PHARMS

- * STADA PHARMACEUTICALS INC
METHOTREXATE SODIUM, METHOTREXATE SODIUM
PROPYLTHIOURACIL, PROPYLTHIOURACIL

STAND HOMEOPATH

- * STANDARD HOMEOPATHIC CO
IVY BLOCK, BENTOQUATAM (OTC)

STAR PHARMS FL

- * STAR PHARMACEUTICALS INC
VIRILON, METHYLTESTOSTERONE

STASON

- * STASON INDUSTRIAL CORP
ACYCLOVIR, ACYCLOVIR
CAPTOPRIL, CAPTOPRIL
SELEGILINE HYDROCHLORIDE, SELEGILINE HYDROCHLORIDE

STASON PHARMS

- * STASON PHARMACEUTICALS INC
CYPROHEPTADINE HYDROCHLORIDE, CYPROHEPTADINE HYDROCHLORIDE

STAT TRADE

- * STAT TRADE INC
MYAMBUTOL, ETHAMBUTOL HYDROCHLORIDE
NAPRELAN, NAPROXEN SODIUM

STEVENS J

- * JEROME STEVENS PHARMACEUTICALS INC
BUTALBITAL, ASPIRIN, CAFFEINE, AND CODEINE PHOSPHATE, ASPIRIN
CEPHALEXIN, CEPHALEXIN
DIGOXIN, DIGOXIN
METHOCARBAMOL AND ASPIRIN, ASPIRIN
ORPHENADRINE CITRATE, ASPIRIN, AND CAFFEINE, ASPIRIN
UNITHROID, LEVOTHYROXINE SODIUM

STIEFEL

- * STIEFEL LABORATORIES INC
CLINDAMYCIN PHOSPHATE, CLINDAMYCIN PHOSPHATE
CLINDETS, CLINDAMYCIN PHOSPHATE
CLOBETASOL PROPIONATE (EMOLLIENT), CLOBETASOL PROPIONATE
CLOBETASOL PROPIONATE, CLOBETASOL PROPIONATE
DUAC, BENZOYL PEROXIDE
ERYTHROMYCIN, ERYTHROMYCIN
GRISEOFULVIN, GRISEOFULVIN, MICROCRYSTALLINE
STIE-CORT, HYDROCORTISONE

SUCAMPO PHARMS

- * SUCAMPO PHARMACEUTICALS INC
AMITIZA, LUBIPROSTONE

SUMMERS

- * SUMMERS LABORATORIES INC
CROTAN, CROTAMITON

SUN PHARM INDS (IN)

- * SUN PHARMACEUTICAL INDUSTRIES LTD
CEPHALEXIN, CEPHALEXIN

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

** S **

- * SUN PHARMACEUTICAL INDUSTRIES LTD
EXTENDED PHENYTOIN SODIUM, PHENYTOIN SODIUM
METFORMIN HYDROCHLORIDE, METFORMIN HYDROCHLORIDE
ONDANSETRON HYDROCHLORIDE, ONDANSETRON HYDROCHLORIDE
ORTHO-EST, ESTROPIRATE
ZONISAMIDE, ZONISAMIDE

SUN PHARM INDS LTD

- * SUN PHARMACEUTICAL INDUSTRIES LTD
GABAPENTIN, GABAPENTIN
ONDANSETRON HYDROCHLORIDE PRESERVATIVE FREE, ONDANSETRON HYDROCHLORIDE

SUPERGEN

- * SUPERGEN INC
ETOPOSIDE, ETOPOSIDE
MITOMYCIN, MITOMYCIN
MYTOZYTREX, MITOMYCIN
PACLITAXEL, PACLITAXEL

SUPPOSITORIA

- * SUPPOSITORIA LABORATORIES INC
ACETAMINOPHEN, ACETAMINOPHEN (OTC)

SWEDISH ORPHAN

- * SWEDISH ORPHAN AB
ORFADIN, NITISINONE

SYNCOR PHARMS

- * SYNCOR PHARMACEUTICALS INC
SODIUM IODIDE I 123, SODIUM IODIDE, I-123

SYNERX PHARMA

- * SYNERX PHARMA LLC
TESTOSTERONE CYPIONATE, TESTOSTERONE CYPIONATE

SYNTHON PHARMS

- * SYNTHON PHARMACEUTICALS LTD
FLUVOXAMINE MALEATE, FLUVOXAMINE MALEATE

SYOSSET

- * SYOSSET LABORATORIES INC
E-SOLVE 2, ERYTHROMYCIN

TABLICAPS

- * TABLICAPS INC
FOLIC ACID, FOLIC ACID
METHOCARBAMOL, METHOCARBAMOL

TAKEDA GLOBAL

- * TAKEDA GLOBAL RESEARCH DEVELOPMENT CENTER INC
ACTOPLUS MET, METFORMIN
ACTOS, PIOGLITAZONE HYDROCHLORIDE
DUETACT, GLIMEPIRIDE
ROZEREM, RAMELTEON

TAP PHARM

- * TAP PHARMACEUTICAL PRODUCTS INC
LUPRON DEPOT, LEUPROLIDE ACETATE
LUPRON DEPOT-3, LEUPROLIDE ACETATE
LUPRON DEPOT-4, LEUPROLIDE ACETATE
LUPRON DEPOT-PED, LEUPROLIDE ACETATE
LUPRON, LEUPROLIDE ACETATE
PREVACID IV, LANSOPRAZOLE
PREVACID NAPRAPAC 500 (COPACKAGED), LANSOPRAZOLE
PREVACID, LANSOPRAZOLE
PREVPAC, AMOXICILLIN

TARGACEPT

- * TARGACEPT INC
INVERSINE, MECAMYLAMINE HYDROCHLORIDE

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

** T **

TARO

- * TARO PHARMACEUTICAL INDUSTRIES LTD
 - ACETAZOLAMIDE, ACETAZOLAMIDE
 - CARBAMAZEPINE, CARBAMAZEPINE
 - CLOMIPRAMINE HYDROCHLORIDE, CLOMIPRAMINE HYDROCHLORIDE
 - ENALAPRIL MALEATE, ENALAPRIL MALEATE
 - ETODOLAC, ETODOLAC
 - EXTENDED PHENYTOIN SODIUM, PHENYTOIN SODIUM
 - FLUCONAZOLE, FLUCONAZOLE
 - HYDROCORTISONE BUTYRATE, HYDROCORTISONE BUTYRATE
 - KETOCONAZOLE, KETOCONAZOLE
 - LORATADINE, LORATADINE (OTC)
 - MELOXICAM, MELOXICAM
 - METRONIDAZOLE, METRONIDAZOLE
 - NORTRIPTYLINE HYDROCHLORIDE, NORTRIPTYLINE HYDROCHLORIDE
 - PHENYTOIN, PHENYTOIN
 - TERIL, CARBAMAZEPINE
- * TARO PHARMACEUTICALS INC
 - BETAMETHASONE DIPROPIONATE, BETAMETHASONE DIPROPIONATE
 - CLOBETASOL PROPIONATE, CLOBETASOL PROPIONATE
 - CLOTRIMAZOLE, CLOTRIMAZOLE
 - CLOTRIMAZOLE, CLOTRIMAZOLE (OTC)
 - DESOXIMETASONE, DESOXIMETASONE
 - DIFLORASONE DIACETATE, DIFLORASONE DIACETATE
 - FLUOCINOLONE ACETONIDE, FLUOCINOLONE ACETONIDE
 - FLUOCINONIDE, FLUOCINONIDE
 - HYDROCORTISONE VALERATE, HYDROCORTISONE VALERATE
 - MICONAZOLE NITRATE, MICONAZOLE NITRATE (OTC)
 - NORTRIPTYLINE HYDROCHLORIDE, NORTRIPTYLINE HYDROCHLORIDE
 - TRIAMCINOLONE ACETONIDE, TRIAMCINOLONE ACETONIDE
 - WARFARIN SODIUM, WARFARIN SODIUM
- * TARO PHARMACEUTICALS USA INC
 - ACETIC ACID, ACETIC ACID, GLACIAL
 - ALCLOMETASONE DIPROPIONATE, ALCLOMETASONE DIPROPIONATE
 - AMIODARONE HYDROCHLORIDE, AMIODARONE HYDROCHLORIDE
 - AMMONIUM LACTATE, AMMONIUM LACTATE
 - BETAMETHASONE DIPROPIONATE, BETAMETHASONE DIPROPIONATE
 - BETAMETHASONE VALERATE, BETAMETHASONE VALERATE
 - CICLOPIROX, CICLOPIROX
 - CIPROFLOXACIN, CIPROFLOXACIN HYDROCHLORIDE
 - CITALOPRAM HYDROBROMIDE, CITALOPRAM HYDROBROMIDE
 - CLOBETASOL PROPIONATE (EMOLLIENT), CLOBETASOL PROPIONATE
 - CLOBETASOL PROPIONATE, CLOBETASOL PROPIONATE
 - CLORAZEPATE DIPOTASSIUM, CLORAZEPATE DIPOTASSIUM
 - CLOTRIMAZOLE AND BETAMETHASONE DIPROPIONATE, BETAMETHASONE DIPROPIONATE
 - CLOTRIMAZOLE, CLOTRIMAZOLE
 - DERMABET, BETAMETHASONE VALERATE
 - DESONIDE, DESONIDE
 - DIFLORASONE DIACETATE, DIFLORASONE DIACETATE
 - ECONAZOLE NITRATE, ECONAZOLE NITRATE
 - FLUOCINOLONE ACETONIDE, FLUOCINOLONE ACETONIDE
 - FLUOCINONIDE EMULSIFIED BASE, FLUOCINONIDE
 - FLUOCINONIDE, FLUOCINONIDE
 - FLUOROURACIL, FLUOROURACIL
 - GENTAMICIN SULFATE, GENTAMICIN SULFATE
 - HALOBETASOL PROPIONATE, HALOBETASOL PROPIONATE
 - HYDROCORTISONE AND ACETIC ACID, ACETIC ACID, GLACIAL
 - HYDROCORTISONE, HYDROCORTISONE
 - KETOZOLE, KETOCONAZOLE
 - LIDOCAINE, LIDOCAINE
 - LORATADINE, LORATADINE (OTC)
 - MICONAZOLE 3, MICONAZOLE NITRATE (OTC)
 - MOMETASONE FUROATE, MOMETASONE FUROATE
 - MUPIROICIN, MUPIROICIN

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

**** T ****

* TARO PHARMACEUTICALS USA INC
NITROFURAZONE, NITROFURAZONE
NYSTATIN AND TRIAMCINOLONE ACETONIDE, NYSTATIN
NYSTATIN, NYSTATIN
ORACORT, TRIAMCINOLONE ACETONIDE
ORALONE, TRIAMCINOLONE ACETONIDE
PROMETHAZINE HYDROCHLORIDE, PROMETHAZINE HYDROCHLORIDE
TERCONAZOLE, TERCONAZOLE
TERIL, CARBAMAZEPINE
THEOPHYLLINE, THEOPHYLLINE
TRIAMCINOLONE ACETONIDE, TRIAMCINOLONE ACETONIDE
TRIVAGIZOLE 3, CLOTRIMAZOLE (OTC)
U-CORT, HYDROCORTISONE ACETATE

TARO PHARM INDS

* TARO PHARMACEUTICAL INDUSTRIES LTD
AMCINONIDE, AMCINONIDE
CARBAMAZEPINE, CARBAMAZEPINE
CLINDAMYCIN PHOSPHATE, CLINDAMYCIN PHOSPHATE
ENALAPRIL MALEATE AND HYDROCHLOROTHIAZIDE, ENALAPRIL MALEATE
ETODOLAC, ETODOLAC
FLUCONAZOLE, FLUCONAZOLE
FLUTICASONE PROPIONATE, FLUTICASONE PROPIONATE
HYDROCORTISONE BUTYRATE, HYDROCORTISONE BUTYRATE

TARO PHARMS IRELAND

* TARO PHARMACEUTICALS IRELAND LTD
SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER, SODIUM CHLORIDE
STERILE WATER FOR INJECTION IN PLASTIC CONTAINER, WATER FOR INJECTION, STERILE

TARO PHARMS NORTH

* TARO PHARMACEUTICALS NORTH AMERICA INC
A/T/S, ERYTHROMYCIN
OVIDE, MALATHION
TOPICORT LP, DESOXIMETASONE
TOPICORT, DESOXIMETASONE

TAYLOR

* TAYLOR PHARMACEUTICALS
LABETALOL HYDROCHLORIDE, LABETALOL HYDROCHLORIDE
MIDAZOLAM HYDROCHLORIDE, MIDAZOLAM HYDROCHLORIDE
NAPHAZOLINE HYDROCHLORIDE, NAPHAZOLINE HYDROCHLORIDE

TAYLOR PHARMA

* TAYLOR PHARMACAL CO
DILTIAZEM HYDROCHLORIDE, DILTIAZEM HYDROCHLORIDE
MIDAZOLAM HYDROCHLORIDE, MIDAZOLAM HYDROCHLORIDE
PROPARACAINE HYDROCHLORIDE, PROPARACAINE HYDROCHLORIDE
TRIMETHOPRIM SULFATE AND POLYMYXIN B SULFATE, POLYMYXIN B SULFATE

TCI MEDCL

* TCI MEDICAL INC
THALLOUS CHLORIDE TL 201, THALLOUS CHLORIDE, TL-201

TECHNILAB

* TECHNILAB INC
ACILAC, LACTULOSE
LAXILOSE, LACTULOSE

TEIKOKU PHARMA USA

* TEIKOKU PHARMA USA INC
LIDODERM, LIDOCAINE

TERCICA

* TERCICA INC
INCRELEX, MECASERMIN RECOMBINANT

TEVA

* TEVA NEUROSCIENCE INC
AZILECT, RASAGILINE MESYLATE

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

** T **

* TEVA NEUROSCIENCE INC
COPAXONE, GLATIRAMER ACETATE

* TEVA PHARMACEUTICALS USA INC
ACETAMINOPHEN AND CODEINE PHOSPHATE, ACETAMINOPHEN
ACYCLOVIR, ACYCLOVIR
ADIPEX-P, PHENTERMINE HYDROCHLORIDE
ALBUTEROL SULFATE, ALBUTEROL SULFATE
ALPRAZOLAM, ALPRAZOLAM
AMILORIDE HYDROCHLORIDE AND HYDROCHLOROTHIAZIDE, AMILORIDE HYDROCHLORIDE
AMIODARONE HYDROCHLORIDE, AMIODARONE HYDROCHLORIDE
AMITRIPTYLINE HYDROCHLORIDE, AMITRIPTYLINE HYDROCHLORIDE
AMOXICILLIN AND CLAVULANATE POTASSIUM, AMOXICILLIN
AMOXICILLIN PEDIATRIC, AMOXICILLIN
AMOXICILLIN, AMOXICILLIN
ATENOLOL, ATENOLOL
AZITHROMYCIN, AZITHROMYCIN
BENAZEPRIL HYDROCHLORIDE, BENAZEPRIL HYDROCHLORIDE
BETAMETHASONE DIPROPIONATE, BETAMETHASONE DIPROPIONATE
BETA-VAL, BETAMETHASONE VALERATE
BISOPROLOL FUMARATE AND HYDROCHLOROTHIAZIDE, BISOPROLOL FUMARATE
BUPROPION HYDROCHLORIDE, BUPROPION HYDROCHLORIDE
BUSPIRONE HYDROCHLORIDE, BUSPIRONE HYDROCHLORIDE
CALCITRIOL, CALCITRIOL
CAPTOPRIL AND HYDROCHLOROTHIAZIDE, CAPTOPRIL
CAPTOPRIL, CAPTOPRIL
CARBIDOPA AND LEVODOPA, CARBIDOPA
CEFACLOR, CEFACLOL
CEFPROZIL, CEFPROZIL
CEFTRIAZONE SODIUM, CEFTRIAZONE SODIUM
CEFUROXIME AXETIL, CEFUROXIME AXETIL
CEFUROXIME, CEFUROXIME SODIUM
CEPHALEXIN, CEPHALEXIN
CEPHRADINE, CEPHRADINE
CHLORHEXIDINE GLUCONATE, CHLORHEXIDINE GLUCONATE
CHLORZOXAZONE, CHLORZOXAZONE
CHOLESTYRAMINE LIGHT, CHOLESTYRAMINE
CHOLESTYRAMINE, CHOLESTYRAMINE
CILOSTAZOL, CILOSTAZOL
CIMETIDINE HYDROCHLORIDE, CIMETIDINE HYDROCHLORIDE
CIMETIDINE, CIMETIDINE
CIPROFLOXACIN, CIPROFLOXACIN HYDROCHLORIDE
CLARITHROMYCIN, CLARITHROMYCIN
CLEMASTINE FUMARATE, CLEMASTINE FUMARATE
CLEMASTINE FUMARATE, CLEMASTINE FUMARATE (OTC)
CLINDAMYCIN HYDROCHLORIDE, CLINDAMYCIN HYDROCHLORIDE
CLOFIBRATE, CLOFIBRATE
CLOMIPRAMINE HYDROCHLORIDE, CLOMIPRAMINE HYDROCHLORIDE
CLONAZEPAM, CLONAZEPAM
CLOTRIMAZOLE, CLOTRIMAZOLE
CLOXACILLIN SODIUM, CLOXACILLIN SODIUM
CLOZAPINE, CLOZAPINE
COTRIM D.S., SULFAMETHOXAZOLE
COTRIM, SULFAMETHOXAZOLE
DICLOFENAC POTASSIUM, DICLOFENAC POTASSIUM
DICLOFENAC SODIUM, DICLOFENAC SODIUM
DICLOXACILLIN SODIUM, DICLOXACILLIN SODIUM
DIFLUNISAL, DIFLUNISAL
DILTIAZEM HYDROCHLORIDE, DILTIAZEM HYDROCHLORIDE
DISOPYRAMIDE PHOSPHATE, DISOPYRAMIDE PHOSPHATE
DOXAZOSIN MESYLATE, DOXAZOSIN MESYLATE
ENALAPRIL MALEATE AND HYDROCHLOROTHIAZIDE, ENALAPRIL MALEATE
ENALAPRIL MALEATE, ENALAPRIL MALEATE
EPITOL, CARBAMAZEPINE
ESTAZOLAM, ESTAZOLAM

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

** T **

* TEVA PHARMACEUTICALS USA INC
 ETODOLAC, ETODOLAC
 FAMOTIDINE, FAMOTIDINE
 FAMOTIDINE, FAMOTIDINE (OTC)
 FENOFIBRATE (MICRONIZED), FENOFIBRATE
 FENOFIBRATE, FENOFIBRATE
 FEXOFENADINE HYDROCHLORIDE, FEXOFENADINE HYDROCHLORIDE
 FINASTERIDE, FINASTERIDE
 FLUCONAZOLE, FLUCONAZOLE
 FLUOCINONIDE EMULSIFIED BASE, FLUOCINONIDE
 FLUOCINONIDE, FLUOCINONIDE
 FLUOXETINE HYDROCHLORIDE, FLUOXETINE HYDROCHLORIDE
 FLUOXETINE, FLUOXETINE HYDROCHLORIDE
 FLURBIPROFEN, FLURBIPROFEN
 FLUVOXAMINE MALEATE, FLUVOXAMINE MALEATE
 FOSINOPRIL SODIUM AND HYDROCHLOROTHIAZIDE, FOSINOPRIL SODIUM
 FOSINOPRIL SODIUM, FOSINOPRIL SODIUM
 GABAPENTIN, GABAPENTIN
 GALZIN, ZINC ACETATE
 GEMFIBROZIL, GEMFIBROZIL
 GLIMEPIRIDE, GLIMEPIRIDE
 GLIPIZIDE, GLIPIZIDE
 GLYBURIDE (MICRONIZED), GLYBURIDE
 GLYBURIDE AND METFORMIN HYDROCHLORIDE, GLYBURIDE
 GLYBURIDE, GLYBURIDE
 HYDROCODONE BITARTRATE AND IBUPROFEN, HYDROCODONE BITARTRATE
 INDAPAMIDE, INDAPAMIDE
 ISOSORBIDE MONONITRATE, ISOSORBIDE MONONITRATE
 KETOCONAZOLE, KETOCONAZOLE
 KETOPROFEN, KETOPROFEN
 KETOROLAC TROMETHAMINE, KETOROLAC TROMETHAMINE
 LABETALOL HYDROCHLORIDE, LABETALOL HYDROCHLORIDE
 LAMOTRIGINE, LAMOTRIGINE
 LISINOPRIL AND HYDROCHLOROTHIAZIDE, HYDROCHLOROTHIAZIDE
 LISINOPRIL, LISINOPRIL
 LOPERAMIDE HYDROCHLORIDE, LOPERAMIDE HYDROCHLORIDE
 LOPERAMIDE HYDROCHLORIDE, LOPERAMIDE HYDROCHLORIDE (OTC)
 LORATADINE, LORATADINE (OTC)
 LOVASTATIN, LOVASTATIN
 MEGESTROL ACETATE, MEGESTROL ACETATE
 MESALAMINE, MESALAMINE
 METAPROTERENOL SULFATE, METAPROTERENOL SULFATE
 METFORMIN HYDROCHLORIDE, METFORMIN HYDROCHLORIDE
 METOCLOPRAMIDE HYDROCHLORIDE, METOCLOPRAMIDE HYDROCHLORIDE
 METOLAZONE, METOLAZONE
 METOPROLOL TARTRATE, METOPROLOL TARTRATE
 METRONIDAZOLE, METRONIDAZOLE
 MEXILETINE HYDROCHLORIDE, MEXILETINE HYDROCHLORIDE
 MICONAZOLE NITRATE, MICONAZOLE NITRATE (OTC)
 MINOCYCLINE HYDROCHLORIDE, MINOCYCLINE HYDROCHLORIDE
 MINOXIDIL (FOR MEN), MINOXIDIL (OTC)
 MINOXIDIL EXTRA STRENGTH (FOR MEN), MINOXIDIL (OTC)
 MIRTAZAPINE, MIRTAZAPINE
 MOEXIPRIL HYDROCHLORIDE, MOEXIPRIL HYDROCHLORIDE
 MUPIROCIN, MUPIROCIN
 NABUMETONE, NABUMETONE
 NAPROXEN SODIUM, NAPROXEN SODIUM
 NAPROXEN, NAPROXEN
 NEFAZODONE HYDROCHLORIDE, NEFAZODONE HYDROCHLORIDE
 NEOMYCIN SULFATE, NEOMYCIN SULFATE
 NICARDIPINE HYDROCHLORIDE, NICARDIPINE HYDROCHLORIDE
 NIFEDIPINE, NIFEDIPINE
 NIZATIDINE, NIZATIDINE
 NORTRIPTYLINE HYDROCHLORIDE, NORTRIPTYLINE HYDROCHLORIDE

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

** T **

* TEVA PHARMACEUTICALS USA INC
 NYSTATIN, NYSTATIN
 OFLOXACIN, OFLOXACIN
 ONDANSETRON HYDROCHLORIDE PRESERVATIVE FREE, ONDANSETRON HYDROCHLORIDE
 ONDANSETRON HYDROCHLORIDE, ONDANSETRON HYDROCHLORIDE
 ORAP, PIMOZIDE
 OXACILLIN SODIUM, OXACILLIN SODIUM
 OXAPROZIN, OXAPROZIN
 OXYCODONE HYDROCHLORIDE, OXYCODONE HYDROCHLORIDE
 PAROXETINE HYDROCHLORIDE, PAROXETINE HYDROCHLORIDE
 PENICILLIN-VK, PENICILLIN V POTASSIUM
 PENTOXIFYLLINE, PENTOXIFYLLINE
 PINDOLOL, PINDOLOL
 PIROXICAM, PIROXICAM
 POTASSIUM CHLORIDE, POTASSIUM CHLORIDE
 PRAVASTATIN SODIUM, PRAVASTATIN SODIUM
 PRELONE, PREDNISOLONE
 PROPOXYPHENE COMPOUND 65, ASPIRIN
 PROPOXYPHENE HYDROCHLORIDE, PROPOXYPHENE HYDROCHLORIDE
 PROPOXYPHENE NAPSYLATE AND ACETAMINOPHEN, ACETAMINOPHEN
 PROPRANOLOL HYDROCHLORIDE, PROPRANOLOL HYDROCHLORIDE
 PURINETHOL, MERCAPTOPURINE
 RANITIDINE HYDROCHLORIDE, RANITIDINE HYDROCHLORIDE
 RIBAVIRIN, RIBAVIRIN
 SELEGILINE HYDROCHLORIDE, SELEGILINE HYDROCHLORIDE
 SERTRALINE HYDROCHLORIDE, SERTRALINE HYDROCHLORIDE
 SOTALOL HYDROCHLORIDE, SOTALOL HYDROCHLORIDE
 SUCRALFATE, SUCRALFATE
 SULFAMETHOXAZOLE AND TRIMETHOPRIM DOUBLE STRENGTH, SULFAMETHOXAZOLE
 SULFAMETHOXAZOLE AND TRIMETHOPRIM, SULFAMETHOXAZOLE
 SULINDAC, SULINDAC
 TAMOXIFEN CITRATE, TAMOXIFEN CITRATE
 THIORIDAZINE HYDROCHLORIDE, THIORIDAZINE HYDROCHLORIDE
 TICLOPIDINE HYDROCHLORIDE, TICLOPIDINE HYDROCHLORIDE
 TIZANIDINE HYDROCHLORIDE, TIZANIDINE HYDROCHLORIDE
 TOLMETIN SODIUM, TOLMETIN SODIUM
 TORSEMIDE, TORSEMIDE
 TRAMADOL HYDROCHLORIDE, TRAMADOL HYDROCHLORIDE
 TRAZODONE HYDROCHLORIDE, TRAZODONE HYDROCHLORIDE
 TRIACET, TRIAMCINOLONE ACETONIDE
 TRIMETHOPRIM, TRIMETHOPRIM
 VENLAFAXINE HYDROCHLORIDE, VENLAFAXINE HYDROCHLORIDE

TEVA PHARMS

* TEVA PHARMACEUTICALS USA
 ACYCLOVIR, ACYCLOVIR
 AMANTADINE HYDROCHLORIDE, AMANTADINE HYDROCHLORIDE
 AMIODARONE HYDROCHLORIDE, AMIODARONE HYDROCHLORIDE
 ATENOLOL, ATENOLOL
 BETAMETHASONE DIPROPIONATE, BETAMETHASONE DIPROPIONATE
 BETAMETHASONE VALERATE, BETAMETHASONE VALERATE
 BISOPROLOL FUMARATE, BISOPROLOL FUMARATE
 CAPTOPRIL, CAPTOPRIL
 CEFADROXIL, CEFADROXIL/CEFADROXIL HEMIHYDRATE
 CEFPROZIL, CEFPROZIL
 CHOLESTYRAMINE LIGHT, CHOLESTYRAMINE
 CHOLESTYRAMINE, CHOLESTYRAMINE
 CIMETIDINE HYDROCHLORIDE, CIMETIDINE HYDROCHLORIDE
 CITALOPRAM HYDROBROMIDE, CITALOPRAM HYDROBROMIDE
 CLEMASTINE FUMARATE, CLEMASTINE FUMARATE
 CLOBETASOL PROPIONATE, CLOBETASOL PROPIONATE
 DESMOPRESSIN ACETATE, DESMOPRESSIN ACETATE
 DESONIDE, DESONIDE
 DEXTROAMP SACCHARATE, AMP ASPARTATE, DEXTROAMP SULFATE AND AMP SULFATE, AMPHETAMINE
 ASPARTATE

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

**** T ****

* TEVA PHARMACEUTICALS USA
DICLOFENAC SODIUM, DICLOFENAC SODIUM
DOXEPIN HYDROCHLORIDE, DOXEPIN HYDROCHLORIDE
ETHOSUXIMIDE, ETHOSUXIMIDE
EVALOSE, LACTULOSE
FLUOCINONIDE, FLUOCINONIDE
GABAPENTIN, GABAPENTIN
GLIPIZIDE AND METFORMIN HYDROCHLORIDE, GLIPIZIDE
GUANABENZ ACETATE, GUANABENZ ACETATE
GYNIX, CLOTRIMAZOLE (OTC)
HALOPERIDOL, HALOPERIDOL LACTATE
HEPTALAC, LACTULOSE
HYDROCORTISONE VALERATE, HYDROCORTISONE VALERATE
HYDROCORTISONE, HYDROCORTISONE
HYDROXYCHLOROQUINE SULFATE, HYDROXYCHLOROQUINE SULFATE
LEFLUNOMIDE, LEFLUNOMIDE
LIDOCAINE HYDROCHLORIDE, LIDOCAINE HYDROCHLORIDE
MEBENDAZOLE, MEBENDAZOLE
MEGESTROL ACETATE, MEGESTROL ACETATE
MELOXICAM, MELOXICAM
METHAZOLAMIDE, METHAZOLAMIDE
METOPROLOL TARTRATE, METOPROLOL TARTRATE
MICONAZOLE NITRATE, MICONAZOLE NITRATE (OTC)
NADOLOL, NADOLOL
NAPROXEN SODIUM, NAPROXEN SODIUM
NAPROXEN, NAPROXEN
PIROXICAM, PIROXICAM
POLYETHYLENE GLYCOL 3350, POLYETHYLENE GLYCOL 3350
PREDNISOLONE, PREDNISOLONE
PROCHLORPERAZINE MALEATE, PROCHLORPERAZINE MALEATE
QUINIDINE SULFATE, QUINIDINE SULFATE
SULFAMETHOXAZOLE AND TRIMETHOPRIM, SULFAMETHOXAZOLE
THIORIDAZINE HYDROCHLORIDE, THIORIDAZINE HYDROCHLORIDE
THIOTHIXENE HYDROCHLORIDE, THIOTHIXENE HYDROCHLORIDE
TRANDOLAPRIL, TRANDOLAPRIL
TRETINOIN, TRETINOIN
URSODIOL, URSODIOL
VALPROIC ACID, VALPROIC ACID
VANDAZOLE, METRONIDAZOLE
ZONISAMIDE, ZONISAMIDE

THERAKOS

* THERAKOS INC
UVADEX, METHOXSALEN

THREE RIVERS PHARMS

* THREE RIVERS PHARMACEUTICALS LLC
AMPHOTEC, AMPHOTERICIN B
RIBASPHERE, RIBAVIRIN
RIBAVIRIN, RIBAVIRIN

TIBOTEC

* TIBOTEC INC
PREZISTA, DARUNAVIR ETHANOLATE

TORPHARM

* TORPHARM INC
BUSPIRONE HYDROCHLORIDE, BUSPIRONE HYDROCHLORIDE
CAPTOPRIL, CAPTOPRIL
CARBIDOPA AND LEVODOPA, CARBIDOPA
CIMETIDINE, CIMETIDINE
CIMETIDINE, CIMETIDINE (OTC)
CIPROFLOXACIN, CIPROFLOXACIN HYDROCHLORIDE
CLONAZEPAM, CLONAZEPAM
CYCLOSPORINE, CYCLOSPORINE
DICLOFENAC POTASSIUM, DICLOFENAC POTASSIUM

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

** T **

* TORPHARM INC
DILT-CD, DILTIAZEM HYDROCHLORIDE
DILTIAZEM HYDROCHLORIDE, DILTIAZEM HYDROCHLORIDE
DOXAZOSIN MESYLATE, DOXAZOSIN MESYLATE
ENALAPRIL MALEATE, ENALAPRIL MALEATE
ETODOLAC, ETODOLAC
FAMOTIDINE, FAMOTIDINE
FAMOTIDINE, FAMOTIDINE (OTC)
FLUCONAZOLE, FLUCONAZOLE
FLUVOXAMINE MALEATE, FLUVOXAMINE MALEATE
GEMFIBROZIL, GEMFIBROZIL
GLIPIZIDE, GLIPIZIDE
KETOCONAZOLE, KETOCONAZOLE
METFORMIN HYDROCHLORIDE, METFORMIN HYDROCHLORIDE
NIZATIDINE, NIZATIDINE
OMEPRazole, OMEPRazole
PAROXETINE HYDROCHLORIDE, PAROXETINE HYDROCHLORIDE
PENTOXIFYLLINE, PENTOXIFYLLINE
QUINAPRIL HYDROCHLORIDE, QUINAPRIL HYDROCHLORIDE
RANITIDINE HYDROCHLORIDE, RANITIDINE HYDROCHLORIDE
RANITIDINE, RANITIDINE HYDROCHLORIDE (OTC)
SELEGILINE HYDROCHLORIDE, SELEGILINE HYDROCHLORIDE
SOTALOL HYDROCHLORIDE, SOTALOL HYDROCHLORIDE
TERAZOSIN HYDROCHLORIDE, TERAZOSIN HYDROCHLORIDE
TICLOPIDINE HYDROCHLORIDE, TICLOPIDINE HYDROCHLORIDE
TIZANIDINE HYDROCHLORIDE, TIZANIDINE HYDROCHLORIDE
TRAMADOL HYDROCHLORIDE, TRAMADOL HYDROCHLORIDE

TRACE RADIOCHEMICALS

* TRACE RADIOCHEMICALS INC
THALLOUS CHLORIDE TL 201, THALLOUS CHLORIDE, TL-201

TRIAx PHARMS

* TRIAX PHARMACEUTICALS LLC
MINOCIN, MINOCYCLINE HYDROCHLORIDE

TRIAx PHARMS LLC

* TRIAX PHARMACEUTICALS LLC
TRETINOIN, TRETINOIN

TRUXTON INC

* TRUXTON INC
DOXYCYCLINE HYCLATE, DOXYCYCLINE HYCLATE

TYCO HLTHCARE

* TYCO HEALTHCARE GROUP LP
ANAFRANIL, CLOMIPRAMINE HYDROCHLORIDE
PAMELOR, NORTRIPTYLINE HYDROCHLORIDE
RESTORIL, TEMAZEPAM
TOFRANIL, IMIPRAMINE HYDROCHLORIDE
TOFRANIL-PM, IMIPRAMINE PAMOATE

UCB INC

* UCB INC
CODEPREX, CHLORPHENIRAMINE POLISTIREX
DIPENTUM, OLSALAZINE SODIUM
HYDROCODONE BITARTRATE AND ACETAMINOPHEN, ACETAMINOPHEN
IONAMIN, PHENTERMINE RESIN COMPLEX
KEPPRA, LEVETIRACETAM
LORTAB, ACETAMINOPHEN
METADATE CD, METHYLPHENIDATE HYDROCHLORIDE
METADATE ER, METHYLPHENIDATE HYDROCHLORIDE
METHYLPHENIDATE HYDROCHLORIDE, METHYLPHENIDATE HYDROCHLORIDE
PEDIAPRED, PREDNISOLONE SODIUM PHOSPHATE
SEMPREX-D, ACRIVASTINE
THEO-24, THEOPHYLLINE
TUSSIONEX PENNKINETIC, CHLORPHENIRAMINE POLISTIREX
ZAROXOLYN, METOLAZONE

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

**** U ****

UCYCLYD

* UCYCLYD PHARMA INC
AMMONUL, SODIUM BENZOATE

UDL

* UDL LABORATORIES INC
METHADONE HYDROCHLORIDE, METHADONE HYDROCHLORIDE
PREDNISOLONE, PREDNISOLONE
VALPROIC ACID, VALPROIC ACID

ULURU

* ULURU INC
AMLEXANOX, AMLEXANOX
APHTHASOL, AMLEXANOX

UNICHEM

* UNICHEM LABORATORIES LTD
MELOXICAM, MELOXICAM

UNIMED

* UNIMED INC
MARINOL, DRONABINOL

UNIMED PHARMS

* UNIMED PHARMACEUTICALS INC
ANDROGEL, TESTOSTERONE
PROMETRIUM, PROGESTERONE

UNIQUE PHARM LABS

* UNIQUE PHARMACEUTICAL LABORATORIES
ATENOLOL, ATENOLOL
CIPROFLOXACIN, CIPROFLOXACIN HYDROCHLORIDE
FLUCONAZOLE, FLUCONAZOLE

UNITED GUARDIAN

* UNITED GUARDIAN INC
RENACIDIN, CITRIC ACID

UNITED THERAP

* UNITED THERAPEUTICS CORP
REMODULIN, TREPROSTINIL SODIUM

UPSHER SMITH

* UPSHER SMITH LABORATORIES INC
BETHANECHOL CHLORIDE, BETHANECHOL CHLORIDE
FORTICAL, CALCITONIN SALMON RECOMBINANT
KLOR-CON M10, POTASSIUM CHLORIDE
KLOR-CON M15, POTASSIUM CHLORIDE
KLOR-CON M20, POTASSIUM CHLORIDE
KLOR-CON, POTASSIUM CHLORIDE
NIACOR, NIACIN
NYSTATIN, NYSTATIN
ORVATEN, MIDODRINE HYDROCHLORIDE
OXANDROLONE, OXANDROLONE
PACERONE, AMIODARONE HYDROCHLORIDE
PENTOXIL, PENTOXIFYLLINE
PREVALITE, CHOLESTYRAMINE
SORINE, SOTALOL HYDROCHLORIDE

US SURGCL

* UNITED STATES SURGICAL DIV TYCO HEALTHCARE GROUP LP
LYMPHAZURIN, ISOSULFAN BLUE

USL PHARMA

* USL PHARMA INC
AMANTADINE HYDROCHLORIDE, AMANTADINE HYDROCHLORIDE
BACLOFEN, BACLOFEN
BENZTROPINE MESYLATE, BENZTROPINE MESYLATE
CHLORDIAZEPOXIDE HYDROCHLORIDE, CHLORDIAZEPOXIDE HYDROCHLORIDE
CHLORPROMAZINE HYDROCHLORIDE, CHLORPROMAZINE HYDROCHLORIDE

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

** U **

* USL PHARMA INC
CLOFIBRATE, CLOFIBRATE
ESTRADIOL, ESTRADIOL
FLUOXYMESTERONE, FLUOXYMESTERONE
JANTOVEN, WARFARIN SODIUM
MEDROXYPROGESTERONE ACETATE, MEDROXYPROGESTERONE ACETATE
OXYBUTYNIN CHLORIDE, OXYBUTYNIN CHLORIDE
VALPROIC ACID, VALPROIC ACID

VALEANT

* VALEANT PHARMACEUTICALS INTERNATIONAL
ACETAMINOPHEN AND CODEINE PHOSPHATE, ACETAMINOPHEN
ANCOBON, FLUCYTOSINE
BONTRIL PDM, PHENDIMETRAZINE TARTRATE
CESAMET, NABILONE
D.H.E. 45, DIHYDROERGOTAMINE MESYLATE
DIASTAT ACUDIAL, DIAZEPAM
DIASTAT, DIAZEPAM
FLUOROURACIL, FLUOROURACIL
MIGRANAL, DIHYDROERGOTAMINE MESYLATE
MOTOFEN, ATROPINE SULFATE
MYSOLINE, PRIMIDONE
PERMAX, PERGOLIDE MESYLATE
PHRENILIN FORTE, ACETAMINOPHEN
PHRENILIN WITH CAFFEINE AND CODEINE, ACETAMINOPHEN
PHRENILIN, ACETAMINOPHEN
TASMAR, TOLCAPONE

VALEANT PHARM INTL

* VALEANT PHARMACEUTICALS INTERNATIONAL
8-MOP, METHOXSALEN
ANDROID 10, METHYLTESTOSTERONE
ANDROID 25, METHYLTESTOSTERONE
BENOQUIN, MONOBENZONE
DALMANE, FLURAZEPAM HYDROCHLORIDE
DIPHENHYDRAMINE HYDROCHLORIDE, DIPHENHYDRAMINE HYDROCHLORIDE
EFUDEX, FLUOROURACIL
LEVO-DROMORAN, LEVORPHANOL TARTRATE
LIBRIUM, CHLORDIAZEPOXIDE HYDROCHLORIDE
LIMBITROL DS, AMITRIPTYLINE HYDROCHLORIDE
LIMBITROL, AMITRIPTYLINE HYDROCHLORIDE
MESTINON, PYRIDOSTIGMINE BROMIDE
OXSORALEN, METHOXSALEN
OXSORALEN-ULTRA, METHOXSALEN
TENSILON PRESERVATIVE FREE, EDROPHONIUM CHLORIDE
TENSILON, EDROPHONIUM CHLORIDE
TESTRED, METHYLTESTOSTERONE
VIRAZOLE, RIBAVIRIN
ZELAPAR, SELEGILINE HYDROCHLORIDE

VALERA

* VALERA PHARMACEUTICALS INC
VALSTAR PRESERVATIVE FREE, VALRUBICIN
VANTAS, HISTRELIN ACETATE

VATRING PHARMS

* VATRING PHARMACEUTICALS LP
UREX, METHENAMINE HIPPURATE

VERNALIS

* VERNALIS RD LTD
APOKYN, APOMORPHINE HYDROCHLORIDE

VERSAPHARM

* VERSAPHARM INC
RIFAMPIN, RIFAMPIN

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

**** V ****

VERUS PHARMS

* VERUS PHARMACEUTICALS INC
TWINJECT 0.3, EPINEPHRINE
TWINJECT, EPINEPHRINE

VICURON

* VICURON PHARMACEUTICALS INC
ERAXIS, ANIDULAFUNGIN

VINTAGE

* VINTAGE PHARMACEUTICALS LLC
ACETAMINOPHEN AND CODEINE PHOSPHATE, ACETAMINOPHEN
ACETIC ACID, ACETIC ACID, GLACIAL
ALBUTEROL SULFATE, ALBUTEROL SULFATE
AMANTADINE HYDROCHLORIDE, AMANTADINE HYDROCHLORIDE
HYDROCORTISONE AND ACETIC ACID, ACETIC ACID, GLACIAL
LEVOLET, LEVOTHYROXINE SODIUM
NYSTATIN, NYSTATIN
PHENYLEPHRINE HYDROCHLORIDE AND PROMETHAZINE HYDROCHLORIDE, PHENYLEPHRINE HYDROCHLORIDE
PROMETH HYDROCHLORIDE, PHENYLEPHRINE HYDROCHLORIDE W/CODEINE PHOSPHATE, CODEINE PHOSPHATE
PROMETHAZINE DM, DEXTROMETHORPHAN HYDROBROMIDE
PROMETHAZINE HYDROCHLORIDE, PROMETHAZINE HYDROCHLORIDE
PROMETHAZINE WITH CODEINE SYRUP, CODEINE PHOSPHATE
SPIRONOLACTONE, SPIRONOLACTONE
TRIAMCINOLONE ACETONIDE, TRIAMCINOLONE ACETONIDE
VALPROIC ACID, VALPROIC ACID

VINTAGE PHARMS

* VINTAGE PHARMACEUTICALS INC
ACETAMINOPHEN AND CODEINE PHOSPHATE, ACETAMINOPHEN
ACETAMINOPHEN, BUTALBITAL, CAFFEINE, AND CODEINE PHOSPHATE, ACETAMINOPHEN
ALLOPURINOL, ALLOPURINOL
AMITRIPTYLINE HYDROCHLORIDE, AMITRIPTYLINE HYDROCHLORIDE
BACLOFEN, BACLOFEN
BUTALBITAL, ACETAMINOPHEN AND CAFFEINE, ACETAMINOPHEN
CARISOPRODOL, CARISOPRODOL
CLONAZEPAM, CLONAZEPAM
DIAZEPAM, DIAZEPAM
DOXYCYCLINE HYCLATE, DOXYCYCLINE HYCLATE
FUROSEMIDE, FUROSEMIDE
HYDROCHLOROTHIAZIDE, HYDROCHLOROTHIAZIDE
HYDROCODONE BITARTRATE AND ACETAMINOPHEN, ACETAMINOPHEN
HYDROCODONE BITARTRATE AND IBUPROFEN, HYDROCODONE BITARTRATE
HYDROCORTISONE, HYDROCORTISONE
HYDROXYZINE HYDROCHLORIDE, HYDROXYZINE HYDROCHLORIDE
IBUPROFEN, IBUPROFEN
IBUPROFEN, IBUPROFEN (OTC)
LACTULOSE, LACTULOSE
LORAZEPAM, LORAZEPAM
MECLIZINE HYDROCHLORIDE, MECLIZINE HYDROCHLORIDE
MEPERIDINE HYDROCHLORIDE, MEPERIDINE HYDROCHLORIDE
METHOCARBAMOL, METHOCARBAMOL
METHYLPREDNISOLONE, METHYLPREDNISOLONE
METOCLOPRAMIDE, METOCLOPRAMIDE HYDROCHLORIDE
NYSTATIN, NYSTATIN
OXYBUTYNIN CHLORIDE, OXYBUTYNIN CHLORIDE
OXYCODONE AND ACETAMINOPHEN, ACETAMINOPHEN
PERPHENAZINE, PERPHENAZINE
PHENTERMINE HYDROCHLORIDE, PHENTERMINE HYDROCHLORIDE
PREDNISONE, PREDNISONE
PRIMIDONE, PRIMIDONE
PROMETHAZINE HYDROCHLORIDE, PROMETHAZINE HYDROCHLORIDE
PROPAFENONE HYDROCHLORIDE, PROPAFENONE HYDROCHLORIDE
PROPOXYPHENE HYDROCHLORIDE AND ACETAMINOPHEN, ACETAMINOPHEN
PROPOXYPHENE NAPSYLATE AND ACETAMINOPHEN, ACETAMINOPHEN

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

**** V ****

* VINTAGE PHARMACEUTICALS INC
SOTALOL HYDROCHLORIDE, SOTALOL HYDROCHLORIDE
SULFASALAZINE, SULFASALAZINE
TRIHEXYPHENIDYL HYDROCHLORIDE, TRIHEXYPHENIDYL HYDROCHLORIDE

VIROPHARMA

* VIROPHARMA INC
VANCOCIN HYDROCHLORIDE, VANCOMYCIN HYDROCHLORIDE

VISTA PHARMS

* VISTA PHARMACEUTICALS INC
SULFAMETHOXAZOLE AND TRIMETHOPRIM, SULFAMETHOXAZOLE

VISTAPHARM

* VISTAPHARM INC
LACTULOSE, LACTULOSE
NYSTATIN, NYSTATIN
PHENYTOIN, PHENYTOIN

VIVUS

* VIVUS INC
MUSE, ALPROSTADIL

VYTERIS

* VYTERIS INC
LIDOSITE TOPICAL SYSTEM KIT, EPINEPHRINE

WARNER CHILCOTT

* WARNER CHILCOTT BERMUDA LTD
CHOLEDYL SA, OXTRIPHYLLINE
* WARNER CHILCOTT INC
DURICEF, CEFADROXIL/CEFADROXIL HEMIHYDRATE
ERYC, ERYTHROMYCIN
ESTRACE, ESTRADIOL
ESTROSTEP FE, ETHINYL ESTRADIOL
FEMHRT, ETHINYL ESTRADIOL
FEMTRACE, ESTRADIOL ACETATE
FLURBIPROFEN, FLURBIPROFEN
LOESTRIN 21 1.5/30, ETHINYL ESTRADIOL
LOESTRIN 21 1/20, ETHINYL ESTRADIOL
LOESTRIN 24 FE, ETHINYL ESTRADIOL
LOESTRIN FE 1.5/30, ETHINYL ESTRADIOL
LOESTRIN FE 1/20, ETHINYL ESTRADIOL
OVCON-35, ETHINYL ESTRADIOL
OVCON-50, ETHINYL ESTRADIOL
SARAFEM, FLUOXETINE HYDROCHLORIDE

WARNER LAMBERT

* WARNER LAMBERT CO
SUDAFED 12 HOUR, PSEUDOEPHEDRINE HYDROCHLORIDE (OTC)

WARRICK PHARMS

* WARRICK PHARMACEUTICALS
CROMOLYN SODIUM, CROMOLYN SODIUM
IPRATROPIUM BROMIDE, IPRATROPIUM BROMIDE

WATSON LABS

* WATSON LABORATORIES
ANAGRELIDE HYDROCHLORIDE, ANAGRELIDE HYDROCHLORIDE
CARBOPLATIN, CARBOPLATIN
FOLIC ACID, FOLIC ACID
LEVONORGESTREL AND ETHINYL ESTRADIOL, ETHINYL ESTRADIOL
METFORMIN HYDROCHLORIDE, METFORMIN HYDROCHLORIDE
MICROGESTIN FE 1.5/30, ETHINYL ESTRADIOL
MICROGESTIN FE 1/20, ETHINYL ESTRADIOL
NORCO, ACETAMINOPHEN
OXAPROZIN, OXAPROZIN
OXYCODONE AND ACETAMINOPHEN, ACETAMINOPHEN
PENTAZOCINE HYDROCHLORIDE AND ACETAMINOPHEN, ACETAMINOPHEN
PROPAFENONE HYDROCHLORIDE, PROPAFENONE HYDROCHLORIDE

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

**** W ****

* WATSON LABORATORIES
TESTOSTERONE, TESTOSTERONE
TRAMADOL HYDROCHLORIDE, TRAMADOL HYDROCHLORIDE

* WATSON LABORATORIES INC
ACEBUTOLOL HYDROCHLORIDE, ACEBUTOLOL HYDROCHLORIDE
ACETAMINOPHEN AND CODEINE PHOSPHATE, ACETAMINOPHEN
ACETAZOLAMIDE, ACETAZOLAMIDE
ACETOHEXAMIDE, ACETOHEXAMIDE
ACYCLOVIR, ACYCLOVIR
AFEDITAB CR, NIFEDIPINE
ALBUTEROL SULFATE, ALBUTEROL SULFATE
ALLOPURINOL, ALLOPURINOL
ALORA, ESTRADIOL
ALPRAZOLAM, ALPRAZOLAM
AMILORIDE HYDROCHLORIDE AND HYDROCHLOROTHIAZIDE, AMILORIDE HYDROCHLORIDE
AMOXAPINE, AMOXAPINE
ANDRODERM, TESTOSTERONE
ATENOLOL AND CHLORTHALIDONE, ATENOLOL
ATENOLOL, ATENOLOL
BACLOFEN, BACLOFEN
BISOPROLOL FUMARATE AND HYDROCHLOROTHIAZIDE, BISOPROLOL FUMARATE
BREVICON 21-DAY, ETHINYL ESTRADIOL
BREVICON 28-DAY, ETHINYL ESTRADIOL
BROMPHENIRAMINE MALEATE, BROMPHENIRAMINE MALEATE
BUSPIRONE HYDROCHLORIDE, BUSPIRONE HYDROCHLORIDE
BUTALBITAL, ACETAMINOPHEN AND CAFFEINE, ACETAMINOPHEN
BUTALBITAL, APAP, AND CAFFEINE, ACETAMINOPHEN
BUTALBITAL, ASPIRIN, CAFFEINE, AND CODEINE PHOSPHATE, ASPIRIN
CAPTOPRIL, CAPTOPRIL
CARISOPRODOL, CARISOPRODOL
CHLORDIAZEPOXIDE AND AMITRIPTYLINE HYDROCHLORIDE, AMITRIPTYLINE HYDROCHLORIDE
CHLORDIAZEPOXIDE HYDROCHLORIDE, CHLORDIAZEPOXIDE HYDROCHLORIDE
CHLORPROPAMIDE, CHLORPROPAMIDE
CHLORZOXAZONE, CHLORZOXAZONE
CIMETIDINE, CIMETIDINE
CIMETIDINE, CIMETIDINE (OTC)
CITALOPRAM HYDROBROMIDE, CITALOPRAM HYDROBROMIDE
CLINDAMYCIN HYDROCHLORIDE, CLINDAMYCIN HYDROCHLORIDE
CLOMIPRAMINE HYDROCHLORIDE, CLOMIPRAMINE HYDROCHLORIDE
CLONAZEPAM, CLONAZEPAM
CLONIDINE HYDROCHLORIDE, CLONIDINE HYDROCHLORIDE
CLORAZEPATE DIPOTASSIUM, CLORAZEPATE DIPOTASSIUM
COL-PROBENECID, COLCHICINE
CORDRAN, FLURANDRENOLIDE
CYCLOBENZAPRINE HYDROCHLORIDE, CYCLOBENZAPRINE HYDROCHLORIDE
DESOGESTREL AND ETHINYL ESTRADIOL, DESOGESTREL
DEXTROAMP SACCHARATE, AMP ASPARTATE, DEXTROAMP SULFATE AND AMP SULFATE, AMPHETAMINE
ASPARTATE
DIAZEPAM, DIAZEPAM
DICLOFENAC POTASSIUM, DICLOFENAC POTASSIUM
DICYCLOMINE HYDROCHLORIDE, DICYCLOMINE HYDROCHLORIDE
DIFLUNISAL, DIFLUNISAL
DILACOR XR, DILTIAZEM HYDROCHLORIDE
DIMENHYDRINATE, DIMENHYDRINATE
DIPHENHYDRAMINE HYDROCHLORIDE PRESERVATIVE FREE, DIPHENHYDRAMINE HYDROCHLORIDE
DIPHENHYDRAMINE HYDROCHLORIDE, DIPHENHYDRAMINE HYDROCHLORIDE
DIPYRIDAMOLE, DIPYRIDAMOLE
DISOPYRAMIDE PHOSPHATE, DISOPYRAMIDE PHOSPHATE
DOBUTAMINE HYDROCHLORIDE, DOBUTAMINE HYDROCHLORIDE
DOXAPRAM HYDROCHLORIDE, DOXAPRAM HYDROCHLORIDE
DOXAZOSIN MESYLATE, DOXAZOSIN MESYLATE
DOXEPIN HYDROCHLORIDE, DOXEPIN HYDROCHLORIDE
DOXYCYCLINE HYCLATE, DOXYCYCLINE HYCLATE
DOXYCYCLINE, DOXYCYCLINE

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

** W **

* WATSON LABORATORIES INC
 ENALAPRIL MALEATE, ENALAPRIL MALEATE
 ERGOLOID MESYLATES, ERGOLOID MESYLATES
 ESTAZOLAM, ESTAZOLAM
 ESTRADIOL CYPIONATE, ESTRADIOL CYPIONATE
 ESTRADIOL VALERATE, ESTRADIOL VALERATE
 ESTRADIOL, ESTRADIOL
 ESTRONE, ESTRONE
 ESTROPIRATE, ESTROPIRATE
 ETODOLAC, ETODOLAC
 FAMOTIDINE, FAMOTIDINE
 FAMOTIDINE, FAMOTIDINE (OTC)
 FENOPROFEN CALCIUM, FENOPROFEN CALCIUM
 FLUOXETINE, FLUOXETINE HYDROCHLORIDE
 FLURAZEPAM HYDROCHLORIDE, FLURAZEPAM HYDROCHLORIDE
 FLUVOXAMINE MALEATE, FLUVOXAMINE MALEATE
 FOSINOPRIL SODIUM AND HYDROCHLOROTHIAZIDE, FOSINOPRIL SODIUM
 FOSINOPRIL SODIUM, FOSINOPRIL SODIUM
 FUROSEMIDE, FUROSEMIDE
 GEMFIBROZIL, GEMFIBROZIL
 GERIMAL, ERGOLOID MESYLATES
 GLIPIZIDE, GLIPIZIDE
 GUANABENZ ACETATE, GUANABENZ ACETATE
 GUANFACINE HYDROCHLORIDE, GUANFACINE HYDROCHLORIDE
 HALOPERIDOL, HALOPERIDOL
 HALOPERIDOL, HALOPERIDOL LACTATE
 HYDRALAZINE HYDROCHLORIDE, HYDRALAZINE HYDROCHLORIDE
 HYDROCHLOROTHIAZIDE, HYDROCHLOROTHIAZIDE
 HYDROCODONE BITARTRATE AND ACETAMINOPHEN, ACETAMINOPHEN
 HYDROMORPHONE HYDROCHLORIDE, HYDROMORPHONE HYDROCHLORIDE
 HYDROXOCOBALAMIN, HYDROXOCOBALAMIN
 HYDROXYCHLOROQUINE SULFATE, HYDROXYCHLOROQUINE SULFATE
 HYDROXYZINE HYDROCHLORIDE, HYDROXYZINE HYDROCHLORIDE
 HYDROXYZINE PAMOATE, HYDROXYZINE PAMOATE
 IBUPROFEN, IBUPROFEN
 IBUPROFEN, IBUPROFEN (OTC)
 INDAPAMIDE, INDAPAMIDE
 ISONIAZID, ISONIAZID
 ISOSORBIDE DINITRATE, ISOSORBIDE DINITRATE
 KETOROLAC TROMETHAMINE, KETOROLAC TROMETHAMINE
 LABETALOL HYDROCHLORIDE, LABETALOL HYDROCHLORIDE
 LEVONORGESTREL AND ETHINYL ESTRADIOL, ETHINYL ESTRADIOL
 LEVORA 0.15/30-28, ETHINYL ESTRADIOL
 LISINAPRIL, LISINAPRIL
 LORAZEPAM, LORAZEPAM
 LOW-OGESTREL-21, ETHINYL ESTRADIOL
 LOW-OGESTREL-28, ETHINYL ESTRADIOL
 LOXAPINE SUCCINATE, LOXAPINE SUCCINATE
 LOXITANE, LOXAPINE SUCCINATE
 MAPROTILINE HYDROCHLORIDE, MAPROTILINE HYDROCHLORIDE
 MECLIZINE HYDROCHLORIDE, MECLIZINE HYDROCHLORIDE
 MECLOFENAMATE SODIUM, MECLOFENAMATE SODIUM
 MELOXICAM, MELOXICAM
 MEPERIDINE HYDROCHLORIDE PRESERVATIVE FREE, MEPERIDINE HYDROCHLORIDE
 MEPERIDINE HYDROCHLORIDE, MEPERIDINE HYDROCHLORIDE
 MEPIVACAINE HYDROCHLORIDE, MEPIVACAINE HYDROCHLORIDE
 MEPROBAMATE, MEPROBAMATE
 METAPROTERENOL SULFATE, METAPROTERENOL SULFATE
 METHOCARBAMOL, METHOCARBAMOL
 METHYCLOTHIAZIDE, METHYCLOTHIAZIDE
 METHYLDOPA, METHYLDOPA
 METHYLPHENIDATE HYDROCHLORIDE, METHYLPHENIDATE HYDROCHLORIDE
 METHYLPREDNISOLONE, METHYLPREDNISOLONE
 METOCLOPRAMIDE HYDROCHLORIDE, METOCLOPRAMIDE HYDROCHLORIDE

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

**** W ****

* WATSON LABORATORIES INC
METOLAZONE, METOLAZONE
METOPROLOL TARTRATE, METOPROLOL TARTRATE
METRONIDAZOLE, METRONIDAZOLE
MEXILETINE HYDROCHLORIDE, MEXILETINE HYDROCHLORIDE
MICROZIDE, HYDROCHLOROTHIAZIDE
MINOCYCLINE HYDROCHLORIDE, MINOCYCLINE HYDROCHLORIDE
MINOXIDIL, MINOXIDIL
MIRTAZAPINE, MIRTAZAPINE
MORPHINE SULFATE, MORPHINE SULFATE
NALIDIXIC ACID, NALIDIXIC ACID
NANDROLONE DECANOATE, NANDROLONE DECANOATE
NAPROXEN SODIUM, NAPROXEN SODIUM
NAPROXEN, NAPROXEN
NEFAZODONE HYDROCHLORIDE, NEFAZODONE HYDROCHLORIDE
NEOMYCIN AND POLYMYXIN B SULFATE, NEOMYCIN SULFATE
NICARDIPINE HYDROCHLORIDE, NICARDIPINE HYDROCHLORIDE
NICOTINE POLACRILEX, NICOTINE POLACRILEX (OTC)
NITROFURANTOIN, NITROFURANTOIN, MACROCRYSTALLINE
NIZATIDINE, NIZATIDINE
NORCO, ACETAMINOPHEN
NORETHIN 1/35E-21, ETHINYL ESTRADIOL
NORETHIN 1/35E-28, ETHINYL ESTRADIOL
NORETHIN 1/50M-21, MESTRANOL
NORETHIN 1/50M-28, MESTRANOL
NORETHINDRONE AND ETHINYL ESTRADIOL (10/11), ETHINYL ESTRADIOL
NORETHINDRONE AND ETHINYL ESTRADIOL (7/14), ETHINYL ESTRADIOL
NORETHINDRONE AND ETHINYL ESTRADIOL, ETHINYL ESTRADIOL
NORETHINDRONE AND MESTRANOL, MESTRANOL
NORGESTIMATE AND ETHINYL ESTRADIOL, ETHINYL ESTRADIOL
NORINYL 1+35 21-DAY, ETHINYL ESTRADIOL
NORINYL 1+35 28-DAY, ETHINYL ESTRADIOL
NORINYL 1+50 21-DAY, MESTRANOL
NORINYL 1+50 28-DAY, MESTRANOL
NORTRIPTYLINE HYDROCHLORIDE, NORTRIPTYLINE HYDROCHLORIDE
OGESTREL 0.5/50-21, ETHINYL ESTRADIOL
OGESTREL 0.5/50-28, ETHINYL ESTRADIOL
ORPHENADRINE CITRATE, ORPHENADRINE CITRATE
OXAZEPAM, OXAZEPAM
OXYCODONE AND ACETAMINOPHEN, ACETAMINOPHEN
OXYCODONE AND ASPIRIN, ASPIRIN
PENTAMIDINE ISETHIONATE, PENTAMIDINE ISETHIONATE
PENTAZOCINE AND NALOXONE HYDROCHLORIDES, NALOXONE HYDROCHLORIDE
PENTOXIFYLLINE, PENTOXIFYLLINE
PINDOLOL, PINDOLOL
PIROXICAM, PIROXICAM
PRAVASTATIN SODIUM, PRAVASTATIN SODIUM
PRAZOSIN HYDROCHLORIDE, PRAZOSIN HYDROCHLORIDE
PREDNISOLONE, PREDNISOLONE
PREDNISONE, PREDNISONE
PRIMIDONE, PRIMIDONE
PROBENECID, PROBENECID
PROCAINAMIDE HYDROCHLORIDE, PROCAINAMIDE HYDROCHLORIDE
PROCAINE HYDROCHLORIDE, PROCAINE HYDROCHLORIDE
PROCHLORPERAZINE EDISYLATE, PROCHLORPERAZINE EDISYLATE
PROMETHAZINE HYDROCHLORIDE, PROMETHAZINE HYDROCHLORIDE
PROPOXYPHENE HYDROCHLORIDE AND ACETAMINOPHEN, ACETAMINOPHEN
PROPRANOLOL HYDROCHLORIDE AND HYDROCHLOROTHIAZIDE, HYDROCHLOROTHIAZIDE
PROPRANOLOL HYDROCHLORIDE, PROPRANOLOL HYDROCHLORIDE
PYRIDOXINE HYDROCHLORIDE, PYRIDOXINE HYDROCHLORIDE
QUASENSE, ETHINYL ESTRADIOL
QUINIDINE GLUCONATE, QUINIDINE GLUCONATE
QUINIDINE SULFATE, QUINIDINE SULFATE
RANITIDINE HYDROCHLORIDE, RANITIDINE HYDROCHLORIDE

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

** W **

- * WATSON LABORATORIES INC
 - RANITIDINE, RANITIDINE HYDROCHLORIDE (OTC)
 - SOTALOL HYDROCHLORIDE, SOTALOL HYDROCHLORIDE
 - SPIRONOLACTONE AND HYDROCHLOROTHIAZIDE, HYDROCHLOROTHIAZIDE
 - SULFAMETHOXAZOLE AND TRIMETHOPRIM DOUBLE STRENGTH, SULFAMETHOXAZOLE
 - SULFAMETHOXAZOLE AND TRIMETHOPRIM, SULFAMETHOXAZOLE
 - SULFASALAZINE, SULFASALAZINE
 - SULINDAC, SULINDAC
 - TEMAZEPAM, TEMAZEPAM
 - TESTOSTERONE CYPIONATE, TESTOSTERONE CYPIONATE
 - TESTOSTERONE ENANTHATE, TESTOSTERONE ENANTHATE
 - TESTOSTERONE PROPIONATE, TESTOSTERONE PROPIONATE
 - TESTOSTERONE, TESTOSTERONE
 - THIAMINE HYDROCHLORIDE, THIAMINE HYDROCHLORIDE
 - THIORIDAZINE HYDROCHLORIDE, THIORIDAZINE HYDROCHLORIDE
 - THIOTHIXENE, THIOTHIXENE
 - TIMOLOL MALEATE, TIMOLOL MALEATE
 - TOLAZAMIDE, TOLAZAMIDE
 - TOLBUTAMIDE, TOLBUTAMIDE
 - TRAZODONE HYDROCHLORIDE, TRAZODONE HYDROCHLORIDE
 - TRELSTAR DEPOT, TRIPTORELIN PAMOATE
 - TRELSTAR, TRIPTORELIN PAMOATE
 - TRIAMTERENE AND HYDROCHLOROTHIAZIDE, HYDROCHLOROTHIAZIDE
 - TRIAZOLAM, TRIAZOLAM
 - TRIHENXYPHENIDYL HYDROCHLORIDE, TRIHENXYPHENIDYL HYDROCHLORIDE
 - TRIMETHOPRIM, TRIMETHOPRIM
 - TRI-NORINYL 28-DAY, ETHINYL ESTRADIOL
 - TRIVORA-28, ETHINYL ESTRADIOL
 - VECURONIUM BROMIDE, VECURONIUM BROMIDE
 - VERAPAMIL HYDROCHLORIDE, VERAPAMIL HYDROCHLORIDE
 - ZONISAMIDE, ZONISAMIDE
 - ZOVIA 1/35E-21, ETHINYL ESTRADIOL
 - ZOVIA 1/35E-28, ETHINYL ESTRADIOL
 - ZOVIA 1/50E-21, ETHINYL ESTRADIOL
 - ZOVIA 1/50E-28, ETHINYL ESTRADIOL
- * WATSON LABS INC
 - LISINAPRIL AND HYDROCHLOROTHIAZIDE, HYDROCHLOROTHIAZIDE

WATSON LABS (UTAH)

- * WATSON LABORATORIES INC
 - CHORIONIC GONADOTROPIN, GONADOTROPIN, CHORIONIC
 - INFED, IRON DEXTRAN
 - NOR-QD, NORETHINDRONE
 - OXYTROL, OXYBUTYNIN
 - PROGESTERONE, PROGESTERONE

WATSON PHARMS

- * WATSON PHARMACEUTICALS
 - TENUATE DOSPAN, DIETHYLPROPION HYDROCHLORIDE
 - TENUATE, DIETHYLPROPION HYDROCHLORIDE
- * WATSON PHARMACEUTICALS INC
 - ACTIGALL, URSODIOL
 - FERRLECIT, SODIUM FERRIC GLUCONATE COMPLEX
 - FIORICET W/ CODEINE, ACETAMINOPHEN
 - FIORICET, ACETAMINOPHEN
 - FIORINAL W/CODEINE, ASPIRIN
 - FIORINAL, ASPIRIN

WE PHARMS

- * WE PHARMACEUTICALS INC
 - PREDNISOLONE SODIUM PHOSPHATE, PREDNISOLONE SODIUM PHOSPHATE
 - PREDNISOLONE, PREDNISOLONE

WEILL MEDCL COLL

- * WEILL MEDICAL COLLEGE CORNELL UNIV
 - FLUDEOXYGLUCOSE F 18, FLUDEOXYGLUCOSE F-18

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

**** W ****

WELLSPRING PHARM

- * WELLSPRING PHARMACEUTICAL CORP
DIBENZYLIN, PHENOXYBENZAMINE HYDROCHLORIDE
DUVOID, BETHANECHOL CHLORIDE
DYRENIUM, TRIAMTERENE

WEST WARD

- * WEST WARD INC
ERGOTAMINE TARTRATE AND CAFFEINE, CAFFEINE
- * WEST WARD PHARMACEUTICAL CORP
ACETAMINOPHEN, CAFFEINE, AND DIHYDROCODEINE BITRATE, ACETAMINOPHEN
AMINOPHYLLINE, AMINOPHYLLINE
BUTALBITAL, ACETAMINOPHEN AND CAFFEINE, ACETAMINOPHEN
BUTALBITAL, ACETAMINOPHEN, AND CAFFEINE WITH CODEINE PHOSPHATE, ACETAMINOPHEN
BUTALBITAL, ASPIRIN AND CAFFEINE, ASPIRIN
CAPTOPRIL, CAPTOPRIL
CARISOPRODOL, CARISOPRODOL
CHLOROQUINE PHOSPHATE, CHLOROQUINE PHOSPHATE
CHLOROTHIAZIDE, CHLOROTHIAZIDE
CORTISONE ACETATE, CORTISONE ACETATE
DICYCLOMINE HYDROCHLORIDE, DICYCLOMINE HYDROCHLORIDE
DOXYCYCLINE HYCLATE, DOXYCYCLINE HYCLATE
ETHAMBUTOL HYDROCHLORIDE, ETHAMBUTOL HYDROCHLORIDE
FLURAZEPAM HYDROCHLORIDE, FLURAZEPAM HYDROCHLORIDE
FOLIC ACID, FOLIC ACID
HYDROCHLOROTHIAZIDE, HYDROCHLOROTHIAZIDE
HYDROCORTISONE, HYDROCORTISONE
ISONIAZID, ISONIAZID
ISOSORBIDE DINITRATE, ISOSORBIDE DINITRATE
ISOSORBIDE MONONITRATE, ISOSORBIDE MONONITRATE
LISINOPRIL AND HYDROCHLOROTHIAZIDE, HYDROCHLOROTHIAZIDE
LISINOPRIL, LISINOPRIL
LITHIUM CARBONATE, LITHIUM CARBONATE
METHOCARBAMOL, METHOCARBAMOL
PREDNISONE, PREDNISONE
PRIMIDONE, PRIMIDONE
PROPOXYPHENE HYDROCHLORIDE, PROPOXYPHENE HYDROCHLORIDE
PROPYLTHIOURACIL, PROPYLTHIOURACIL
TRIHENYPHENIDYL HYDROCHLORIDE, TRIHENYPHENIDYL HYDROCHLORIDE

WESTWARD

- * WESTWARD PHARMACEUTICAL CORP
CEFADROXIL, CEFADROXIL/CEFADROXIL HEMIHYDRATE
NAPROXEN, NAPROXEN
RIFAMPIN AND ISONIAZID, ISONIAZID

WESTWOOD SQUIBB

- * WESTWOOD SQUIBB PHARMACEUTICALS INC
CAPITROL, CHLOROXINE
EURAX, CROTAMITON
EXELDERM, SULCONAZOLE NITRATE
HALOG, HALCINONIDE
LAC-HYDRIN, AMMONIUM LACTATE
MYCOSTATIN, NYSTATIN
ULTRAVATE, HALOBETASOL PROPIONATE
WESTCORT, HYDROCORTISONE VALERATE

WOCKHARDT

- * WOCKHARDT AMERICAS INC
CAPTOPRIL, CAPTOPRIL
ENALAPRIL MALEATE, ENALAPRIL MALEATE
FAMOTIDINE, FAMOTIDINE
RANITIDINE HYDROCHLORIDE, RANITIDINE HYDROCHLORIDE
- * WOCKHARDT LTD
BETHANECHOL CHLORIDE, BETHANECHOL CHLORIDE
CEFOTAXIME, CEFOTAXIME SODIUM

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

** W **

* WOCKHARDT LTD
CEFTRIAZONE, CEFTRIAZONE SODIUM
CEFUROXIME AXETIL, CEFUROXIME AXETIL
CLARITHROMYCIN, CLARITHROMYCIN
FAMOTIDINE, FAMOTIDINE (OTC)
NIACIN, NIACIN
ONDANSETRON HYDROCHLORIDE PRESERVATIVE FREE, ONDANSETRON HYDROCHLORIDE
ONDANSETRON HYDROCHLORIDE, ONDANSETRON HYDROCHLORIDE
RANITIDINE, RANITIDINE HYDROCHLORIDE (OTC)
ZONISAMIDE, ZONISAMIDE

WYETH CONS

* WYETH CONSUMER HEALTHCARE
ADVIL ALLERGY SINUS, CHLORPHENIRAMINE MALEATE (OTC)
ADVIL COLD AND SINUS, IBUPROFEN (OTC)
ADVIL COLD AND SINUS, IBUPROFEN POTASSIUM (OTC)
ADVIL LIQUI-GELS, IBUPROFEN POTASSIUM (OTC)
ADVIL MIGRAINE LIQUI-GELS, IBUPROFEN POTASSIUM (OTC)
ADVIL PM, DIPHENHYDRAMINE CITRATE (OTC)
ADVIL PM, DIPHENHYDRAMINE HYDROCHLORIDE (OTC)
ADVIL, IBUPROFEN (OTC)
ALAVERT, LORATADINE (OTC)
AXID AR, NIZATIDINE (OTC)
BRONITIN MIST, EPINEPHRINE BITARTRATE (OTC)
CHILDREN'S ADVIL ALLERGY SINUS, CHLORPHENIRAMINE MALEATE (OTC)
CHILDREN'S ADVIL COLD, IBUPROFEN (OTC)
CHILDREN'S ADVIL, IBUPROFEN
CHILDREN'S ADVIL, IBUPROFEN (OTC)
CHILDREN'S ADVIL-FLAVORED, IBUPROFEN (OTC)
JUNIOR STRENGTH ADVIL, IBUPROFEN (OTC)
LORATADINE, LORATADINE (OTC)
PEDIATRIC ADVIL, IBUPROFEN (OTC)
PRIMATENE MIST, EPINEPHRINE (OTC)

WYETH PHARMS INC

* WYETH PHARMACEUTICALS INC
ALESSE, ETHINYL ESTRADIOL
CORDARONE, AMIODARONE HYDROCHLORIDE
EFFEXOR XR, VENLAFAXINE HYDROCHLORIDE
EFFEXOR, VENLAFAXINE HYDROCHLORIDE
INDERAL LA, PROPRANOLOL HYDROCHLORIDE
INDERAL, PROPRANOLOL HYDROCHLORIDE
INDERIDE-40/25, HYDROCHLOROTHIAZIDE
LO/OVRAL-28, ETHINYL ESTRADIOL
MYLOTARG, GEMTUZUMAB OZOGAMICIN
ORUVAIL, KETOPROFEN
PHENERGAN, PROMETHAZINE HYDROCHLORIDE
PREMARIN, ESTROGENS, CONJUGATED
PREMPHASE 14/14, ESTROGENS, CONJUGATED
PREMPRO, ESTROGENS, CONJUGATED
PROTONIX IV, PANTOPRAZOLE SODIUM
PROTONIX, PANTOPRAZOLE SODIUM
RAPAMUNE, SIROLIMUS
TRECATOR-SC, ETHIONAMIDE
TRIPHASIL-28, ETHINYL ESTRADIOL
TYGACIL, TIGECYCLINE
ZOSYN IN PLASTIC CONTAINER, PIPERACILLIN SODIUM
ZOSYN, PIPERACILLIN SODIUM

X GEN PHARMS

* X GEN PHARMACEUTICALS INC
AMPHOTERICIN B, AMPHOTERICIN B
BACIIM, BACITRACIN
BACI-RX, BACITRACIN
COLISTIMETHATE, COLISTIMETHATE SODIUM
HYDROCORTISONE ACETATE, HYDROCORTISONE ACETATE

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT

**** X ****

* X GEN PHARMACEUTICALS INC
HYDRO-RX, HYDROCORTISONE
LIOETHYRONINE SODIUM, LIOETHYRONINE SODIUM
NEO-FRADIN, NEOMYCIN SULFATE
NEOMYCIN AND POLYMYXIN B SULFATE, NEOMYCIN SULFATE
NEOMYCIN SULFATE, NEOMYCIN SULFATE
NEO-RX, NEOMYCIN SULFATE
NYSTATIN, NYSTATIN
POLYMYXIN B SULFATE, POLYMYXIN B SULFATE
POLY-RX, POLYMYXIN B SULFATE
STREPTOMYCIN SULFATE, STREPTOMYCIN SULFATE
TOBRAMYCIN, TOBRAMYCIN SULFATE

XANODYNE PHARM

* XANODYNE PHARMACAL INC
AMICAR, AMINOCAPROIC ACID
DARVOCET A500, ACETAMINOPHEN
DARVOCET-N 100, ACETAMINOPHEN
DARVOCET-N 50, ACETAMINOPHEN
DARVON COMPOUND-65, ASPIRIN
DARVON, PROPOXYPHENE HYDROCHLORIDE
DARVON-N, PROPOXYPHENE NAPSYLATE
DOLOPHINE HYDROCHLORIDE, METHADONE HYDROCHLORIDE
DURACLON, CLONIDINE HYDROCHLORIDE
LEUCOVORIN CALCIUM, LEUCOVORIN CALCIUM
ORAMORPH SR, MORPHINE SULFATE
ROXICODONE, OXYCODONE HYDROCHLORIDE

XTTRIUM

* XTTRIUM LABORATORIES INC
DYNA-HEX, CHLORHEXIDINE GLUCONATE (OTC)
EXIDINE, CHLORHEXIDINE GLUCONATE (OTC)

YUNG SHIN PHARM

* YUNG SHIN PHARMACEUTICAL INDUSTRIAL CO LTD
CEPHALEXIN, CEPHALEXIN

YVR THERAP

* YVR THERAPEUTICS LLC
POLYETHYLENE GLYCOL 3350, POLYETHYLENE GLYCOL 3350

ZAMBON

* ZAMBON CORP
MONUROL, FOSFOMYCIN TROMETHAMINE

ZARS

* ZARS INC
LIDOCAINE AND TETRACAINE, LIDOCAINE

ZENITH GOLDLINE

* ZENITH GOLDLINE LABORATORIES INC
METFORMIN HYDROCHLORIDE, METFORMIN HYDROCHLORIDE
* ZENITH GOLDLINE PHARMACEUTICALS
BUSPIRONE HYDROCHLORIDE, BUSPIRONE HYDROCHLORIDE
* ZENITH GOLDLINE PHARMACEUTICALS INC
NIZATIDINE, NIZATIDINE

ZILA

* ZILA PHARMACEUTICALS INC
PERIDEX, CHLORHEXIDINE GLUCONATE

ZLB BEHRING

* ZLB BEHRING LLC
STIMATE, DESMOPRESSIN ACETATE

ZYDUS PHARMS USA

* ZYDUS PHARMACEUTICALS USA INC
ATENOLOL, ATENOLOL
CLINDAMYCIN HYDROCHLORIDE, CLINDAMYCIN HYDROCHLORIDE
MELOXICAM, MELOXICAM

APPENDIX B - PRODUCT NAME SORTED BY APPLICANT**** Z ****

* ZYDUS PHARMACEUTICALS USA INC
METFORMIN HYDROCHLORIDE, METFORMIN HYDROCHLORIDE
PROMETHAZINE HYDROCHLORIDE, PROMETHAZINE HYDROCHLORIDE
RIBAVIRIN, RIBAVIRIN
SIMVASTATIN, SIMVASTATIN
WARFARIN SODIUM, WARFARIN SODIUM
ZONISAMIDE, ZONISAMIDE

APPENDIX C**UNIFORM TERMS*****DOSAGE FORMS***

AEROSOL	LOTION/SHAMPOO
AEROSOL, FOAM	OIL
AEROSOL, METERED	OIL/DROPS
CAPSULE	OINTMENT
CAPSULE, DELAYED REL PELLETS	OINTMENT, AUGMENTED
CAPSULE, DELAYED RELEASE	PASTE
CAPSULE, EXTENDED RELEASE	PATCH
CLOTH	PELLET
CONCENTRATE	POWDER
CREAM	POWDER, EXTENDED RELEASE
CREAM, AUGMENTED	POWDER, METERED
ELIXIR	RING
EMULSION	SHAMPOO
ENEMA	SOLUTION
FILM, EXTENDED RELEASE	SOLUTION FOR SLUSH
FOR SOLUTION	SOLUTION, GEL FORMING/DROPS
FOR SUSPENSION	SOLUTION/DROPS
FOR SUSPENSION, DELAYED RELEASE	SPONGE
FOR SUSPENSION, EXTENDED RELEASE	SPRAY
GAS	SPRAY, METERED
GEL	SUPPOSITORY
GEL, AUGMENTED	SUSPENSION
GEL, METERED	SUSPENSION, EXTENDED RELEASE
GRANULE	SUSPENSION/DROPS
GRANULE, DELAYED RELEASE	SWAB
GRANULE, EFFERVESCENT	SYRUP
GUM, CHEWING	SYSTEM
IMPLANT	TABLET
INHALANT	TABLET, CHEWABLE
INJECTABLE	TABLET, COATED PARTICLES
INJECTABLE, LIPID COMPLEX	TABLET, DELAYED RELEASE
INJECTABLE, LIPOSOMAL	TABLET, DELAYED RELEASE, ORALLY
INSERT	DISINTEGRATING
INSERT, EXTENDED RELEASE	TABLET, EFFERVESCENT
INTRAUTERINE DEVICE	TABLET, EXTENDED RELEASE
JELLY	TABLET, FOR SUSPENSION
LIQUID	TABLET, ORALLY DISINTEGRATING
LIQUID, EXTENDED RELEASE	TAPE
LOTION	TROCHE/LOZENGE
LOTION, AUGMENTED	

Note: Terms comprise currently marketed products

APPENDIX C

UNIFORM TERMS

ROUTES OF ADMINISTRATION

BUCCAL
 DENTAL
 ENDOCERVICAL
 EPIDURAL
 FOR RX COMPOUNDING
 IM-IV
 IM-IV-SC
 IMPLANTATION
 IM-SC
 INHALATION
 INJECTION
 INTRACRANIAL
 INTRALYMPHATIC
 INTRAMUSCULAR
 INTRAOCULAR
 INTRAPERITONEAL
 INTRAPLEURAL
 INTRATHECAL
 INTRATRACHEAL
 INTRAUTERINE
 INTRAVENOUS
 INTRAVESICAL
 INTRAVITREAL
 IONTOPHORESIS, TOPICAL
 IONTOPHORESIS, TRANSDERMAL
 IRRIGATION
 IV (INFUSION)
 IV (INFUSION)-SC
 N/A
 NASAL
 OPHTHALMIC
 ORAL
 ORAL-21
 ORAL-28
 OTIC
 PERFUSION, CARDIAC
 PERIODONTAL
 RECTAL
 SPINAL
 SUBCUTANEOUS
 SUBLINGUAL
 TOPICAL
 TRANSDERMAL
 TRANSMUCOSAL
 URETERAL
 URETHRAL
 VAGINAL

Note: Terms comprise currently marketed products

APPENDIX C**UNIFORM TERMS*****ABBREVIATIONS***

AMP	AMPULE
AMPICIL	AMPICILLIN
APPROX	APPROXIMATELY
BOT	BOTTLE
CI	CURIE
CSR	CAROTID SINUS REFLEX
CU	CLINICAL UNITS
DIPROP	DIPROPIONATE
ELECT	ELECTROLYTE
EQ	EQUIVALENT TO
ER	EXTENDED RELEASE
GM	GRAM
HBR	HYDROBROMIDE
HCL	HYDROCHLORIDE
HR	HOURLY
INH	INHALATION
IU	INTERNATIONAL UNITS
KIU	KALLIKREIN INHIBITOR UNITS
MCI	MILLICURIE
MEQ	MILLIEQUIVALENT
MG	MILLIGRAM
ML	MILLILITER
N/A	NOT APPLICABLE
PPM	PARTS PER MILLION
REL	RELEASE
SQ CM	SQUARE CENTIMETER
U	UNITS
UCI	MICROCURIE
UGM	MICROGRAM
UMOLAR	MICROMOLAR
USP	UNITED STATES PHARMACOPEIA

PATENT AND EXCLUSIVITY INFORMATION ADDENDUM

This *Addendum* identifies drugs that qualify under the Drug Price Competition and Patent Term Restoration Act (1984 Amendments) for periods of exclusivity, during which abbreviated new drug applications (ANDAs) and applications described in Section 505(b)(2) of the Federal Food, Drug, and Cosmetic Act (the Act) for those drug products may, in some instances, not be submitted or made effective as described below, and provides patent information concerning the listed drug products. Those drugs that have qualified for Orphan Drug Exclusivity pursuant to Section 527 of the Act and those drugs that have qualified for Pediatric Exclusivity pursuant to Section 505A are also included in this *Addendum*. This section is arranged in alphabetical order by active ingredient name followed the trade name. Active ingredient headings for multiple ingredient (combination) drug products are arranged alphabetically. For an explanation of the codes used in the *Addendum*, see the *Patent and Exclusivity Terms* Section. Exclusivity prevents the submission or effective approval of ANDAs or applications described in Section 505(b)(2) of the Act. It does not prevent the submission or approval of a second 505(b)(1) application except in the case of Orphan Drug exclusivity. Applications qualifying for periods of exclusivity are:

- (1) A new drug application approved after September 24, 1984, for a drug product all active ingredients (including any ester or salt of the active ingredient) of which had never been approved in any other new drug application under Section 505 (b) of the Act. No subsequent ANDA or application described in Section 505(b)(2) of the Act for the same drug may be *submitted* for a period of *five years* from the date of approval of the original application, except that such an application may be *submitted* after *four years* if it contains a certification that a patent claiming the drug is invalid or will not be infringed by the product for which approval is sought.
- (2) A new drug application approved after September 24, 1984, for a drug product containing an active ingredient (including any ester or salt of that active ingredient) that has been approved in an earlier new drug application and that includes reports of new clinical investigations (other than bioavailability studies). Such investigations must have been conducted or sponsored by the applicant and must have been essential to approval of the application. If these requirements are met, the approval of a subsequent ANDA or an application described in Section 505(b)(2) of the Act may not be *made effective* for the same drug or use, if for a new indication, before the expiration of *three years* from the date of approval of the original application. If an applicant has exclusivity for a new application or 505(b)(2) application for the drug product with indications or use, this does not preclude the approval of an ANDA or 505(b)(2) application not covered by the exclusivity.
- (3) A supplement to a new drug application for a drug containing a previously approved active ingredient including (any ester or salt of the active ingredient) approved after September 24, 1984, that contains reports of new clinical investigations (other than bioavailability studies) essential to the approval of the supplement and conducted or sponsored by the applicant. The approval of a subsequent ANDA or 505(b)(2) application for a change approved in the supplement may not be

made effective for three years from the date of approval of the original supplement.

The Act requires that patent information be filed with all newly submitted Section 505(b) drug applications. No NDA may be approved after September 24, 1984, without the submission of patent information to the Agency. Effective August 18, 2003, this information must be filed using FDA Form 3524a "Patent Information Submitted with the Filing of an NDA, Amendment or Supplement".

Effective August 18, 2003, this information must be submitted to the agency upon approval on FDA Form 3542 "Patent Information Submitted Upon and After Approval of an NDA or Supplement". Patent information on unapproved applications or on patents beyond the scope of the Act (i.e., process or manufacturing patents) will not be published. FDA form 3542 will be the only form used for the purposes of this publication.

The patents that FDA regards as covered by the statutory provisions for submission of patent information are: patents that claim the active ingredient(s); drug product patents which include formulation/composition patents; use patents for a particular approved indication or method of using the product; and certain other patents as detailed on FDA Form 3542. This information, as provided by the sponsor on FDA form 3542, will be published as described above.

Upon approval, patent numbers and expiration dates, in addition to certain other information on appropriate patents claiming drug products that are the subject of approved applications, will be published on a daily basis in the Electronic Orange Book, <http://www.fda.gov/cder/ob/default.htm>. The Addendum lists patent and exclusivity information up to January of the Edition year. The monthly Cumulative Supplements to the annual edition list patent and exclusivity information changes since the Annual Edition Addendum. Since all parts of this publication are subject to changes, additions, or deletions, the Electronic Orange Book, updated daily, should be consulted for the most recent patent and exclusivity information.

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
ABACAVIR SULFATE; ZIAGEN					
020977 001	5034394	Dec 18, 2011		D-40	Aug 02, 2007
	5034394*PED	Jun 18, 2012			
	5089500	Jun 26, 2009	U-248		
	5089500*PED	Dec 26, 2009			
	6294540	May 14, 2018	U-65		
6294540*PED	Nov 14, 2018	U-65			
ABACAVIR SULFATE; ZIAGEN					
020978 001	5034394	Dec 18, 2011		D-40	Aug 02, 2007
	5034394*PED	Jun 18, 2012			
	5089500	Jun 26, 2009	U-248		
	5089500*PED	Dec 26, 2009			
	6294540	May 14, 2018	U-65		
6294540*PED	Nov 14, 2018	U-65			
ABACAVIR SULFATE; LAMIVUDINE; EPZICOM					
021652 001	5034394	Dec 18, 2011	DS DP	D-40	Aug 02, 2007
	5034394*PED	Jun 18, 2012			
	5047407	Nov 17, 2009	DS DP	U-257	
	5047407*PED	May 17, 2010			
	5089500	Jun 26, 2009		U-257	
	5089500*PED	Dec 26, 2009			
	5905082	May 18, 2016	DS DP		
	5905082*PED	Nov 18, 2016			
	6294540	May 14, 2018	DS DP	U-257	
	6294540*PED	Nov 14, 2018			
	6417191	Mar 28, 2016	DP	U-257	
	ABACAVIR SULFATE; LAMIVUDINE; ZIDOVUDINE; TRIZIVIR				
021205 001	5034394	Dec 18, 2011	DS DP		
	5034394*PED	Jun 18, 2012			
	5047407	Nov 17, 2009	DS DP	U-248	
	5047407*PED	May 17, 2010			
	5089500	Jun 26, 2009		U-248	
	5089500*PED	Dec 26, 2009		U-248	
	5905082	May 18, 2016	DS DP	U-248	
	5905082*PED	Nov 18, 2016			
	6294540	May 14, 2018	DS DP	U-65	
	6294540*PED	Nov 14, 2018		U-65	
	6417191	Mar 28, 2016	DP	U-248	
	ABARELIX; PLENAXIS				
021320 001	5843901	Dec 01, 2015	DS DP	NCE	Nov 25, 2008
	5968895	Dec 11, 2016	DP		
	6180608	Dec 11, 2016	DP	U-549	
	6423686	Jun 07, 2015	DS		
	6455499	Jun 07, 2015		U-549	
	6699833	Dec 11, 2016	DP		
ACAMPROSATE CALCIUM; CAMPRAL					
021431 001				NCE	Jul 29, 2009
ACARBOSE; PRECOSE					
020482 001	4904769	Feb 27, 2007			
ACARBOSE; PRECOSE					
020482 002	4904769	Feb 27, 2007			
ACARBOSE; PRECOSE					
020482 004	4904769	Feb 27, 2007			
ACETAMINOPHEN; TYLENOL (CAPLET)					
019872 001	4820522	Jul 27, 2007			
	4968509	Nov 06, 2007			
	5004613	Jul 27, 2007			
ACETAMINOPHEN; ASPIRIN; CAFFEINE; EXCEDRIN (MIGRAINE)					
020802 001	4943565	Jul 24, 2007			
	5972916	Jul 14, 2017	U-296		
ACETAMINOPHEN; TRAMADOL HYDROCHLORIDE; ULTRACET					
021123 001	RE39221	Aug 09, 2011	DS DP	U-55	
ACETYLCHOLINE CHLORIDE; MIOCHOL-E					
020213 001	6261546	Apr 29, 2019		U-506	

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>ACETYLCYSTEINE; ACETADOTE</u>					
021539 001				ODE	Jan 23, 2011
<u>ACRIVASTINE; PSEUDOEPHEDRINE HYDROCHLORIDE; SEMPREX-D</u>					
019806 001	4650807	Mar 26, 2008	U-93		
<u>ACYCLOVIR; ZOVIRAX</u>					
021478 001	4963555	Oct 16, 2007			
<u>ADAPALENE; DIFFERIN</u>					
020338 001	4717720	May 31, 2010			
	RE34440	May 31, 2010	U-275		
<u>ADAPALENE; DIFFERIN</u>					
020380 001	4717720	May 31, 2010			
	RE34440	May 31, 2010	U-275		
<u>ADAPALENE; DIFFERIN</u>					
020748 001	4717720	May 31, 2010			
	RE34440	May 31, 2010	U-275		
<u>ADEFOVIR DIPIVOXIL; HEPSERA</u>					
021449 001	5663159	Sep 02, 2014	DS DP	NCE	Sep 20, 2007
	6451340	Jul 23, 2018	DS DP	U-470	
<u>ADENOSINE; ADENOSCAN</u>					
020059 001	5070877	May 18, 2009		U-116	
	5731296	Mar 24, 2015		U-221	
<u>ALATROFLOXACIN MESYLATE; TROVAN PRESERVATIVE FREE</u>					
020760 001	5164402	Nov 17, 2009		U-282	
	5763454	Jun 15, 2015		U-282	
	6080756	Jul 05, 2016			
	6194429	Jul 23, 2018			
<u>ALATROFLOXACIN MESYLATE; TROVAN PRESERVATIVE FREE</u>					
020760 002	5164402	Nov 17, 2009		U-282	
	5763454	Jun 15, 2015		U-282	
	6080756	Jul 05, 2016			
	6194429	Jul 23, 2018			
<u>ALBUMIN HUMAN; OPTISON</u>					
020899 001	5529766	Jun 25, 2013		U-505	
	5558094	Feb 28, 2012		U-505	
	5573751	Apr 25, 2012			
	6723303	Apr 20, 2021	DP		
<u>ALBUTEROL SULFATE; ACCUNEB</u>					
020949 001	6702997	Dec 28, 2021		U-558	
<u>ALBUTEROL SULFATE; ACCUNEB</u>					
020949 002	6702997	Dec 28, 2021		U-558	
<u>ALBUTEROL SULFATE; PROAIR HFA</u>					
021457 001	5605674	Feb 25, 2014	DP	I-235	Feb 03, 2009
	5695743	Jul 06, 2010	DP	U-491	
	5766573	Nov 28, 2009		U-356	Oct 29, 2007
	6352684	Nov 28, 2009	DP		
<u>ALBUTEROL SULFATE; PROVENTIL-HFA</u>					
020503 001	5225183	Jul 06, 2010			
	5439670	Jul 06, 2010			
	5605674	Feb 25, 2014			
	5695743	Jul 06, 2010		U-491	
	5766573	Jun 16, 2015			
	6352684	Nov 28, 2009			
<u>ALBUTEROL SULFATE; VENTOLIN HFA</u>					
020983 001	6131566	Apr 14, 2015	DP	U-716	
	6131566	Apr 14, 2015	DP	U-589	
	6315173	Dec 23, 2017	DP		
	6510969	Dec 23, 2017	DP		
	6532955	Apr 14, 2015	DP	U-590	
	6532955	Apr 14, 2015	DP	U-716	
<u>ALBUTEROL SULFATE; IPRATROPIUM BROMIDE; COMBIVENT</u>					
020291 001	5603918	Jun 09, 2015			

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>ALBUTEROL SULFATE; IPRATROPIUM BROMIDE; DUONEB</u>					
020950 001	6632842	Dec 28, 2021	U-532		
<u>ALCOHOL; CHLORHEXIDINE GLUCONATE; AVAGARD</u>					
021074 001	5897031	Jun 21, 2016			
<u>ALENDRONATE SODIUM; FOSAMAX</u>					
020560 001	4621077	Aug 06, 2007	U-114	M-51	Dec 21, 2008
	4621077*PED	Feb 06, 2008	U-114		
	5358941	Dec 02, 2012			
	5358941*PED	Jun 02, 2013			
	5681590	Dec 02, 2012			
	5681590*PED	Jun 02, 2013			
	5849726	Jun 06, 2015			
	5849726*PED	Dec 06, 2015			
	6008207	Jun 06, 2015	U-303		
	6008207*PED	Dec 06, 2015	U-303		
	6090410	Dec 02, 2012			
	6090410*PED	Jun 02, 2013			
	6194004	Dec 02, 2012			
	6194004*PED	Jun 02, 2013			
<u>ALENDRONATE SODIUM; FOSAMAX</u>					
020560 002	4621077	Aug 06, 2007	U-114	M-51	Dec 21, 2008
	4621077*PED	Feb 06, 2008	U-114		
	5358941	Dec 02, 2012			
	5358941*PED	Jun 02, 2013			
	5681590	Dec 02, 2012			
	5681590*PED	Jun 02, 2013			
	5849726	Jun 06, 2015			
	5849726*PED	Dec 06, 2015			
	6008207	Jun 06, 2015	U-303		
	6008207*PED	Jun 06, 2015	U-303		
	6090410	Dec 02, 2012			
	6090410*PED	Jun 02, 2013			
<u>ALENDRONATE SODIUM; FOSAMAX</u>					
020560 003	4621077	Aug 06, 2007	U-114	M-51	Dec 21, 2008
	4621077*PED	Feb 06, 2008	U-114		
	5358941	Dec 02, 2012			
	5358941*PED	Jun 02, 2013			
	5681590	Dec 02, 2012			
	5681590*PED	Jun 02, 2013			
	5849726	Jun 06, 2015			
	5849726*PED	Dec 06, 2015			
	6008207	Jun 06, 2015	U-303		
	6008207*PED	Dec 06, 2015	U-303		
	6090410	Dec 02, 2012			
	6090410*PED	Jun 02, 2013			
<u>ALENDRONATE SODIUM; FOSAMAX</u>					
020560 004	4621077	Aug 06, 2007	U-114	M-51	Dec 21, 2008
	4621077*PED	Feb 06, 2008	U-114		
	5358941	Dec 02, 2012			
	5358941*PED	Jun 02, 2013			
	5681590	Dec 02, 2012			
	5681590*PED	Jun 02, 2013			
	5849726	Jun 06, 2015			
	5849726*PED	Dec 16, 2015			
	5994329	Jul 17, 2018			
	5994329*PED	Jan 17, 2019			
	6008207	Jun 06, 2015			
	6008207*PED	Dec 06, 2015			
	6015801	Jul 17, 2018	U-353		
	6015801*PED	Jan 17, 2019	U-353		
	6090410	Dec 02, 2012			
	6090410*PED	Jun 02, 2013			
	6225294	Jul 17, 2018			
	6225294*PED	Jan 17, 2019			

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>ALENDRONATE SODIUM; FOSAMAX</u>					
020560 005	4621077	Aug 06, 2007	U-114	D-87	Apr 16, 2007
	4621077*PED	Feb 06, 2008	U-114	M-51	Dec 21, 2008
	5358941	Dec 02, 2012			
	5358941*PED	Jun 02, 2013			
	5681590	Dec 02, 2012			
	5681590*PED	Jun 02, 2013			
	5849726	Jun 06, 2015			
	5849726*PED	Dec 06, 2015			
	5994329	Jul 17, 2018			
	5994329*PED	Jan 17, 2019			
	6008207	Jun 06, 2015			
	6008207*PED	Dec 06, 2015			
	6015801	Jul 17, 2018	U-353		
	6015801*PED	Jan 17, 2019	U-353		
	6090410	Dec 02, 2012			
	6090410*PED	Jun 02, 2013			
	6225294	Jul 17, 2018			
	6225294*PED	Jan 17, 2019			
<u>ALENDRONATE SODIUM; FOSAMAX</u>					
021575 001	4621077	Aug 06, 2007		D-87	Apr 16, 2007
	4621077*PED	Feb 06, 2008		M-51	Dec 21, 2008
	5462932	May 17, 2014			
	5994329	Jul 17, 2018			
	5994329*PED	Jan 17, 2019			
	6015801	Jul 17, 2018			
	6015801*PED	Jan 17, 2019			
	6225294	Jul 17, 2018			
	6225294*PED	Jan 17, 2019			
<u>ALENDRONATE SODIUM; CHOLECALCIFEROL; FOSAMAX PLUS D</u>					
021762 001	4621077	Aug 06, 2007	U-648	NC	Apr 07, 2008
	4621077*PED	Feb 06, 2008			
	5358941	Dec 02, 2012	DP		
	5358941*PED	Jun 02, 2013			
	5681590	Dec 02, 2012	DP		
	5681590*PED	Jun 02, 2013			
	5994329	Jul 17, 2018	U-647		
	5994329*PED	Jan 17, 2019			
	6090410	Dec 02, 2012	DP		
	6090410*PED	Jun 02, 2013	DP		
<u>ALFUZOSIN HYDROCHLORIDE; UROXATRAL</u>					
021287 001	4661491	May 27, 2007	U-706	NCE	Jun 12, 2008
	6149940	Aug 22, 2017			
<u>ALITRETINOIN; PANRETIN</u>					
020886 001	5932622	Aug 03, 2016	U-562		
	7056954	Aug 02, 2012	DP		
<u>ALMOTRIPTAN MALATE; AXERT</u>					
021001 001	5565447	May 07, 2015			
<u>ALMOTRIPTAN MALATE; AXERT</u>					
021001 002	5565447	May 07, 2015			
<u>ALOSETRON HYDROCHLORIDE; LOTRONEX</u>					
021107 001	5360800	Jan 13, 2013	DS DP	U-405	
<u>ALOSETRON HYDROCHLORIDE; LOTRONEX</u>					
021107 002	5360800	Jan 13, 2013	DS DP	U-405	
<u>ALPRAZOLAM; NIRAVAM</u>					
021726 001	6024981	Apr 09, 2018	DP		
	6221392	Apr 09, 2018	DP		
<u>ALPRAZOLAM; NIRAVAM</u>					
021726 002	6024981	Apr 09, 2018	DP		
	6221392	Apr 09, 2018	DP		
<u>ALPRAZOLAM; NIRAVAM</u>					
021726 003	6024981	Apr 09, 2018	DP		
	6221392	Apr 09, 2018	DP		

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>ALPRAZOLAM; NIRAVAM</u>					
021726 004	6024981	Apr 09, 2018	DP		
	6221392	Apr 09, 2018	DP		
<u>ALPRAZOLAM; XANAX</u>					
018276 004	5061494	Dec 12, 2008			
<u>ALPROSTADIL; CAVERJECT</u>					
020379 001	5741523	Apr 21, 2015			
<u>ALPROSTADIL; CAVERJECT</u>					
020379 002	5741523	Apr 21, 2015			
<u>ALPROSTADIL; CAVERJECT</u>					
020379 003	5741523	Apr 21, 2015			
<u>ALPROSTADIL; CAVERJECT</u>					
020379 004	5741523	Apr 21, 2015			
<u>ALPROSTADIL; MUSE</u>					
020700 001	4801587	Jan 31, 2007		U-155	
	5242391	Sep 07, 2010		U-155	
	5474535	Dec 12, 2012		U-155	
<u>ALPROSTADIL; MUSE</u>					
020700 002	4801587	Jan 31, 2007		U-155	
	5242391	Sep 07, 2010		U-155	
	5474535	Dec 12, 2012		U-155	
<u>ALPROSTADIL; MUSE</u>					
020700 003	4801587	Jan 31, 2007		U-155	
	5242391	Sep 07, 2010		U-155	
	5474535	Dec 12, 2012		U-155	
<u>ALPROSTADIL; MUSE</u>					
020700 004	4801587	Jan 31, 2007		U-155	
	5242391	Sep 07, 2010		U-155	
	5474535	Dec 12, 2012		U-155	
<u>AMIFOSTINE; ETHYOL</u>					
020221 001	5424471	Jul 31, 2012			
	5591731	Jul 31, 2012			
	5994409	Dec 08, 2017		U-305	
<u>AMIFOSTINE; ETHYOL</u>					
020221 002	5424471	Jul 31, 2012			
	5591731	Jul 31, 2012			
	5994409	Dec 08, 2017		U-305	
<u>AMINOLEVULINIC ACID HYDROCHLORIDE; LEVULAN</u>					
020965 001	5079262	Jul 28, 2009		U-289	
	5211938	May 18, 2010		U-289	
	5422093	Jul 28, 2009		U-289	
	5954703	Oct 31, 2017		U-289	
	6710066	Jul 28, 2009		U-289	
<u>AMLEXANOX; AMLEXANOX</u>					
021727 001				NDF	Sep 29, 2007
<u>AMLEXANOX; APHTHASOL</u>					
020511 001	5362737	Nov 08, 2011		U-243	
<u>AMLODIPINE BESYLATE; NORVASC</u>					
019787 001	4572909	Jul 31, 2006	DS DP	U-683	I-472 Sep 28, 2008
	4572909*PED	Jan 31, 2007			NPP Jan 08, 2007
	4879303	Mar 25, 2007	DS DP		PED Jul 08, 2007
	4879303*PED	Sep 25, 2007			
<u>AMLODIPINE BESYLATE; NORVASC</u>					
019787 002	4572909	Jul 31, 2006	DS DP	U-683	I-472 Sep 28, 2008
	4572909*PED	Jan 31, 2007			NPP Jan 08, 2007
	4879303	Mar 25, 2007	DS DP		PED Jul 08, 2007
	4879303*PED	Sep 25, 2007			

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>AMLODIPINE BESYLATE; NORVASC</u>					
019787 003	4572909	Jul 31, 2006	DS DP U-683	I-472	Sep 28, 2008
	4572909*PED	Jan 31, 2007		NPP	Jan 08, 2007
	4879303	Mar 25, 2007	DS DP	PED	Jul 08, 2007
	4879303*PED	Sep 25, 2007			
<u>AMLODIPINE BESYLATE; ATORVASTATIN CALCIUM; CADUET</u>					
021540 001	4572909	Jul 31, 2006	DS DP U-3	NC	Jan 30, 2007
	4572909*PED	Jan 31, 2007			
	4681893	Sep 24, 2009	DS DP U-161		
	4681893*PED	Mar 24, 2010			
	4879303	Mar 25, 2007	DS DP		
	4879303*PED	Sep 25, 2007			
	5273995	Dec 28, 2010	DS DP U-162		
	5273995*PED	Jun 28, 2011			
	5686104	Nov 11, 2014	DP U-213		
	5686104*PED	May 11, 2015			
	5969156	Jul 08, 2016	DS		
	5969156*PED	Jan 08, 2017			
	6126971	Jan 19, 2013	DP		
	6126971*PED	Jul 19, 2013			
	6455574	Aug 11, 2018	U-552		
<u>AMLODIPINE BESYLATE; ATORVASTATIN CALCIUM; CADUET</u>					
021540 002	4572909	Jul 31, 2006	DS DP U-3	NC	Jan 30, 2007
	4572909*PED	Jan 31, 2007			
	4681893	Sep 24, 2009	DS DP U-161		
	4681893*PED	Mar 24, 2010			
	4879303	Mar 25, 2007	DS DP		
	4879303*PED	Sep 25, 2007			
	5273995	Dec 28, 2010	DS DP U-162		
	5273995*PED	Jun 28, 2011			
	5686104	Nov 11, 2014	DP U-213		
	5686104*PED	May 11, 2015			
	5969156	Jul 08, 2016	DS		
	5969156*PED	Jan 08, 2017			
	6126971	Jan 19, 2013	DP		
	6126971*PED	Jul 19, 2013			
	6455574	Aug 11, 2018	U-552		
<u>AMLODIPINE BESYLATE; ATORVASTATIN CALCIUM; CADUET</u>					
021540 003	4572909	Jul 31, 2006	DS DP U-3	NC	Jan 30, 2007
	4572909*PED	Jan 31, 2007			
	4681893	Sep 24, 2009	DS DP U-161		
	4681893*PED	Mar 24, 2010			
	4879303	Mar 25, 2007	DS DP		
	4879303*PED	Sep 25, 2007			
	5273995	Dec 28, 2010	DS DP U-162		
	5273995*PED	Jun 28, 2011			
	5686104	Nov 11, 2014	DP U-213		
	5686104*PED	May 11, 2015			
	5969156	Jul 08, 2016	DS		
	5969156*PED	Jan 08, 2017			
	6126971	Jan 19, 2013	DP		
	6126971*PED	Jul 19, 2013			
	6455574	Aug 11, 2018	U-552		
<u>AMLODIPINE BESYLATE; ATORVASTATIN CALCIUM; CADUET</u>					
021540 004	4572909	Jul 31, 2006	DS DP U-3	NC	Jan 30, 2007
	4572909*PED	Jan 31, 2007			
	4681893	Sep 24, 2009	DS DP U-161		
	4681893*PED	Mar 24, 2010			
	4879303	Mar 25, 2007	DS DP		
	4879303*PED	Sep 25, 2007			
	5273995	Dec 28, 2010	DS DP U-162		
	5273995*PED	Jun 28, 2011			
	5686104	Nov 11, 2014	DP U-213		
	5686104*PED	May 11, 2015			
	5969156	Jul 08, 2016	DS		
	5969156*PED	Jan 08, 2017			
	6126971	Jan 19, 2013	DP		
	6126971*PED	Jul 19, 2013			
	6455574	Aug 11, 2018	U-552		

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
AMLODIPINE BESYLATE; ATORVASTATIN CALCIUM; CADUET					
021540 005	4572909	Jul 31, 2006	DS DP U-3	NC	Jan 30, 2007
	4572909*PED	Jan 31, 2007			
	4681893	Sep 24, 2009	DS DP U-161		
	4681893*PED	Mar 24, 2010			
	4879303	Mar 25, 2007	DS DP		
	4879303*PED	Sep 25, 2007			
	5273995	Dec 28, 2010	DS DP U-162		
	5273995*PED	Jun 28, 2011			
	5686104	Nov 11, 2014	DP U-213		
	5686104*PED	May 11, 2015			
	5969156	Jul 08, 2016	DS		
	5969156*PED	Jan 08, 2017			
	6126971	Jan 19, 2013	DP		
	6126971*PED	Jul 19, 2013			
	6455574	Aug 11, 2018	U-552		
AMLODIPINE BESYLATE; ATORVASTATIN CALCIUM; CADUET					
021540 006	4572909	Jul 31, 2006	DS DP U-3	NC	Jan 30, 2007
	4572909*PED	Jan 31, 2007			
	4681893	Sep 24, 2009	DS DP U-161		
	4681893*PED	Mar 24, 2010			
	4879303	Mar 25, 2007	DS DP		
	4879303*PED	Sep 25, 2007			
	5273995	Dec 28, 2010	DS DP U-162		
	5273995*PED	Jun 28, 2011			
	5686104	Nov 11, 2014	DP U-213		
	5686104*PED	May 11, 2015			
	5969156	Jul 08, 2016	DS		
	5969156*PED	Jan 08, 2017			
	6126971	Jan 19, 2013	DP		
	6126971*PED	Jul 19, 2013			
	6455574	Aug 11, 2018	U-552		
AMLODIPINE BESYLATE; ATORVASTATIN CALCIUM; CADUET					
021540 007	4572909	Jul 31, 2006	DS DP U-3	NC	Jan 30, 2007
	4572909*PED	Jan 31, 2007			
	4681893	Sep 24, 2009	DS DP U-161		
	4681893*PED	Mar 24, 2010			
	4879303	Mar 25, 2007	DS DP		
	4879303*PED	Sep 25, 2007			
	5273995	Dec 28, 2010	DS DP U-162		
	5273995*PED	Jun 28, 2011			
	5686104	Nov 11, 2014	DP U-213		
	5686104*PED	May 11, 2015			
	5969156	Jul 08, 2016	DS		
	5969156*PED	Jan 08, 2017			
	6126971	Jan 19, 2013	DP		
	6126971*PED	Jul 19, 2013			
	6455574	Aug 11, 2018	U-552		
AMLODIPINE BESYLATE; ATORVASTATIN CALCIUM; CADUET					
021540 008	4572909	Jul 31, 2006	DS DP U-3	NC	Jan 30, 2007
	4572909*PED	Jan 31, 2007			
	4681893	Sep 24, 2009	DS DP U-161		
	4681893*PED	Mar 24, 2010			
	4879303	Mar 25, 2007	DS DP		
	4879303*PED	Sep 25, 2007			
	5273995	Dec 28, 2010	DS DP U-162		
	5273995*PED	Jun 28, 2011			
	5686104	Nov 11, 2014	DP U-213		
	5686104*PED	May 11, 2015			
	5969156	Jul 08, 2016	DS		
	5969156*PED	Jan 08, 2017			
	6126971	Jan 19, 2013	DP		
	6126971*PED	Jul 19, 2013			
	6455574	Aug 11, 2018	U-552		

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE		PATENT CODES			EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE	
AMLODIPINE BESYLATE; ATORVASTATIN CALCIUM; CADUET									
021540 009	4572909	Jul 31, 2006	DS	DP	U-3	NC	Jan 30, 2007		
	4572909*PED	Jan 31, 2007							
	4681893	Sep 24, 2009	DS	DP	U-161				
	4681893*PED	Mar 24, 2010							
	4879303	Mar 25, 2007	DS	DP					
	4879303*PED	Sep 25, 2007							
	5273995	Dec 28, 2010	DS	DP	U-162				
	5273995*PED	Jun 28, 2011							
	5686104	Nov 11, 2014	DP	U-213					
	5686104*PED	May 11, 2015							
	5969156	Jul 08, 2016	DS						
	5969156*PED	Jan 08, 2017							
	6126971	Jan 19, 2013	DP						
	6126971*PED	Jul 19, 2013							
	6455574	Aug 11, 2018		U-552					
AMLODIPINE BESYLATE; ATORVASTATIN CALCIUM; CADUET									
021540 010	4572909	Jul 31, 2006	DS	DP	U-3	NC	Jan 30, 2007		
	4572909*PED	Jan 31, 2007							
	4681893	Sep 24, 2009	DS	DP	U-161				
	4681893*PED	Mar 24, 2010							
	4879303	Mar 25, 2007	DS	DP					
	4879303*PED	Sep 25, 2007							
	5273995	Dec 28, 2010	DS	DP	U-162				
	5273995*PED	Jun 28, 2011							
	5686104	Nov 11, 2014	DP	U-213					
	5686104*PED	May 11, 2015							
	5969156	Jul 08, 2016	DS						
	5969156*PED	Jan 08, 2017							
	6126971	Jan 19, 2013	DP						
	6126971*PED	Jul 19, 2013							
	6455574	Aug 11, 2018		U-552					
AMLODIPINE BESYLATE; ATORVASTATIN CALCIUM; CADUET									
021540 011	4572909	Jul 31, 2006	DS	DP	U-3	NC	Jan 30, 2007		
	4572909*PED	Jan 31, 2007							
	4681893	Sep 24, 2009	DS	DP	U-161				
	4681893*PED	Mar 24, 2010							
	4879303	Mar 25, 2007	DS	DP					
	4879303*PED	Sep 25, 2007							
	5273995	Dec 28, 2010	DS	DP	U-162				
	5273995*PED	Jun 28, 2011							
	5686104	Nov 11, 2014	DP	U-213					
	5686104*PED	May 11, 2015							
	5969156	Jul 08, 2016	DS						
	5969156*PED	Jan 08, 2017							
	6126971	Jan 19, 2013	DP						
	6126971*PED	Jul 19, 2013							
	6455574	Aug 11, 2018		U-552					
AMLODIPINE BESYLATE; BENAZEPRIL HYDROCHLORIDE; LOTREL									
020364 002	4572909	Jul 31, 2006							
	4879303	Mar 25, 2007							
	6162802	Dec 19, 2017							
AMLODIPINE BESYLATE; BENAZEPRIL HYDROCHLORIDE; LOTREL									
020364 003	4572909	Jul 31, 2006							
	4879303	Mar 25, 2007							
	6162802	Dec 19, 2017							
AMLODIPINE BESYLATE; BENAZEPRIL HYDROCHLORIDE; LOTREL									
020364 004	4572909	Jul 31, 2006							
	4879303	Mar 25, 2007							
	6162802	Dec 19, 2017							
AMLODIPINE BESYLATE; BENAZEPRIL HYDROCHLORIDE; LOTREL									
020364 005	4572909	Jul 31, 2006							
	4879303	Mar 25, 2007							
	6162802	Dec 19, 2017							
AMLODIPINE BESYLATE; BENAZEPRIL HYDROCHLORIDE; LOTREL									
020364 006	4879303	Mar 25, 2007	DS	DP	U-185	NS	Apr 11, 2009		
	6162802	Dec 19, 2017	DS	DP					

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>AMLODIPINE BESYLATE; BENAZEPRIL HYDROCHLORIDE; LOTREL</u>					
020364 007	4879303	Mar 25, 2007	DS DP	NS	Apr 11, 2009
	6162802	Dec 19, 2017	DS DP U-185		
<u>AMPHETAMINE ASPARTATE; AMPHETAMINE SULFATE; DEXTROAMPHETAMINE SACCHARATE; DEXTROAMPHETAMINE SULFATE; ADDERALL 10</u>					
011522 007	6384020	Jul 06, 2020			
	6384020*PED	Jan 06, 2021			
<u>AMPHETAMINE ASPARTATE; AMPHETAMINE SULFATE; DEXTROAMPHETAMINE SACCHARATE; DEXTROAMPHETAMINE SULFATE; ADDERALL 12.5</u>					
011522 012	6384020	Jul 06, 2020			
	6384020*PED	Jan 06, 2021			
<u>AMPHETAMINE ASPARTATE; AMPHETAMINE SULFATE; DEXTROAMPHETAMINE SACCHARATE; DEXTROAMPHETAMINE SULFATE; ADDERALL 15</u>					
011522 013	6384020	Jul 06, 2020			
	6384020*PED	Jan 06, 2021			
<u>AMPHETAMINE ASPARTATE; AMPHETAMINE SULFATE; DEXTROAMPHETAMINE SACCHARATE; DEXTROAMPHETAMINE SULFATE; ADDERALL 20</u>					
011522 008	6384020	Jul 06, 2020			
	6384020*PED	Jan 06, 2021			
<u>AMPHETAMINE ASPARTATE; AMPHETAMINE SULFATE; DEXTROAMPHETAMINE SACCHARATE; DEXTROAMPHETAMINE SULFATE; ADDERALL 30</u>					
011522 010	6384020	Jul 06, 2020			
	6384020*PED	Jan 06, 2021			
<u>AMPHETAMINE ASPARTATE; AMPHETAMINE SULFATE; DEXTROAMPHETAMINE SACCHARATE; DEXTROAMPHETAMINE SULFATE; ADDERALL 5</u>					
011522 009	6384020	Jul 06, 2020			
	6384020*PED	Jan 06, 2021			
<u>AMPHETAMINE ASPARTATE; AMPHETAMINE SULFATE; DEXTROAMPHETAMINE SACCHARATE; DEXTROAMPHETAMINE SULFATE; ADDERALL 7.5</u>					
011522 011	6384020	Jul 06, 2020			
	6384020*PED	Jan 06, 2021			
<u>AMPHETAMINE ASPARTATE; AMPHETAMINE SULFATE; DEXTROAMPHETAMINE SACCHARATE; DEXTROAMPHETAMINE SULFATE; ADDERALL XR 10</u>					
021303 001	6322819	Oct 21, 2018		NPP	Jul 21, 2008
	6322819*PED	Apr 21, 2019		NPP	Aug 11, 2007
	6605300	Oct 21, 2018		PED	Jan 21, 2009
	6605300*PED	Apr 21, 2019			
<u>AMPHETAMINE ASPARTATE; AMPHETAMINE SULFATE; DEXTROAMPHETAMINE SACCHARATE; DEXTROAMPHETAMINE SULFATE; ADDERALL XR 15</u>					
021303 006	6322819	Oct 21, 2018		NPP	Jul 21, 2008
	6322819*PED	Apr 21, 2019		NPP	Aug 11, 2007
	6605300	Oct 21, 2018		PED	Jan 21, 2009
	6605300*PED	Apr 21, 2019			
<u>AMPHETAMINE ASPARTATE; AMPHETAMINE SULFATE; DEXTROAMPHETAMINE SACCHARATE; DEXTROAMPHETAMINE SULFATE; ADDERALL XR 20</u>					
021303 002	6322819	Oct 21, 2018		NPP	Jul 21, 2008
	6322819*PED	Apr 21, 2019		NPP	Aug 11, 2007
	6605300	Oct 21, 2018		PED	Jan 21, 2009
	6605300*PED	Apr 21, 2019			
<u>AMPHETAMINE ASPARTATE; AMPHETAMINE SULFATE; DEXTROAMPHETAMINE SACCHARATE; DEXTROAMPHETAMINE SULFATE; ADDERALL XR 25</u>					
021303 004	6322819	Oct 21, 2018		NPP	Jul 21, 2008
	6322819*PED	Apr 21, 2019		NPP	Aug 11, 2007
	6605300	Oct 21, 2018		PED	Jan 21, 2009
	6605300*PED	Apr 21, 2019			
<u>AMPHETAMINE ASPARTATE; AMPHETAMINE SULFATE; DEXTROAMPHETAMINE SACCHARATE; DEXTROAMPHETAMINE SULFATE; ADDERALL XR 30</u>					
021303 003	6322819	Oct 21, 2018		NPP	Jul 21, 2008
	6322819*PED	Apr 21, 2019		NPP	Aug 11, 2007
	6605300	Oct 21, 2018		PED	Jan 21, 2009
	6605300*PED	Apr 21, 2019			

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>AMPHETAMINE ASPARTATE; AMPHETAMINE SULFATE; DEXTROAMPHETAMINE SACCHARATE; DEXTROAMPHETAMINE SULFATE; ADDERALL XR 5</u>					
021303 005	6322819	Oct 21, 2018		NPP	Jul 21, 2008
	6322819*PED	Apr 21, 2019		NPP	Aug 11, 2007
	6605300	Oct 21, 2018		PED	Jan 21, 2009
	6605300*PED	Apr 21, 2019			
<u>AMPRENAVIR; AGENERASE</u>					
021007 001	5585397	Dec 17, 2013			
	5646180	Jul 08, 2014		U-257	
	5723490	Mar 03, 2015		U-257	
	6730679	Nov 11, 2017	DP		
<u>AMPRENAVIR; AGENERASE</u>					
021007 002	5585397	Dec 17, 2013			
	5646180	Jul 08, 2014		U-257	
	5723490	Mar 03, 2015		U-257	
	6730679	Nov 11, 2017	DP		
<u>AMPRENAVIR; AGENERASE</u>					
021039 001	5585397	Dec 17, 2013			
	5646180	Jul 08, 2014		U-257	
	5723490	Mar 03, 2015		U-257	
<u>ANAGRELIDE HYDROCHLORIDE; AGRYLIN</u>					
020333 001				M-39	Dec 10, 2007
				PED	Jun 10, 2008
<u>ANAGRELIDE HYDROCHLORIDE; AGRYLIN</u>					
020333 002				M-39	Dec 10, 2007
				PED	Jun 10, 2008
<u>ANASTROZOLE; ARIMIDEX</u>					
020541 001	RE36617	Dec 27, 2009			
<u>ANIDULAFUNGIN; ERAXIS</u>					
021632 001	5965525	Oct 12, 2016	DS DP	U-540	NCE Feb 17, 2011
	6384013	Mar 19, 2012	DS		
	6743777	Mar 19, 2012		DP U-540	
	6960564	Apr 12, 2021		DP U-540	
<u>ANIDULAFUNGIN; ERAXIS</u>					
021632 002				NCE	Feb 17, 2011
<u>APOMORPHINE HYDROCHLORIDE; APOKYN</u>					
021264 001				NCE	Apr 20, 2009
				ODE	Apr 20, 2011
<u>APOMORPHINE HYDROCHLORIDE; APOKYN</u>					
021264 002				NCE	Apr 20, 2009
				ODE	Apr 20, 2011
<u>APRACLONIDINE HYDROCHLORIDE; IOPIDINE</u>					
019779 001	5212196	May 18, 2010		U-120	
<u>APREPITANT; EMEND</u>					
021549 001	5145684	Jan 25, 2011		DP	I-475 Oct 28, 2008
	5538982	Jul 23, 2013		U-745	NCE Mar 26, 2008
	5719147	Jun 29, 2012	DS DP		
	6048859	Jun 29, 2012		U-745	
	6096742	Jul 01, 2018	DS DP	U-745	
	6235735	Jun 29, 2012		U-747	
	6235735	Jun 29, 2012		U-746	
<u>APREPITANT; EMEND</u>					
021549 002	5145684	Jan 25, 2011		DP	I-475 Oct 28, 2008
	5538982	Jul 23, 2013		U-745	NCE Mar 26, 2008
	5719147	Jun 29, 2012	DS DP		
	6048859	Jun 29, 2012		U-745	
	6096742	Jul 01, 2018	DS DP	U-745	
	6235735	Jun 29, 2012		U-746	
	6235735	Jun 29, 2012		U-747	

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>APREPITANT; EMEND</u>					
021549 003	5145684	Jan 25, 2011	DP	I-498	Jun 30, 2009
	5538982	Jul 23, 2013	U-745	NS	Jun 30, 2009
	5719147	Jun 29, 2012	DS DP	NCE	Mar 26, 2008
	6048859	Jun 29, 2012	U-745		
	6096742	Jul 01, 2018	DS DP	U-745	
	6235735	Jun 29, 2012	U-747		
	6235735	Jun 29, 2012	U-746		
<u>ARBUTAMINE HYDROCHLORIDE; GENESA</u>					
020420 001	5108363	Apr 28, 2009	U-220		
	5234404	Aug 10, 2010	U-220		
	5395970	Mar 07, 2012			
<u>ARFORMOTEROL TARTRATE; BROVANA</u>					
021912 001				NP	Oct 06, 2009
<u>ARGATROBAN; ARGATROBAN</u>					
020883 001	5214052	Jun 30, 2014			
<u>ARIPIPRAZOLE; ABILIFY</u>					
021436 001	5006528	Oct 20, 2014	DS DP	U-761	I-488
					Mar 01, 2008
					I-437
					Sep 29, 2007
					I-401
					Aug 28, 2006
					NCE
					Nov 15, 2007
<u>ARIPIPRAZOLE; ABILIFY</u>					
021436 002	5006528	Oct 20, 2014	DS DP	U-761	I-488
					Mar 01, 2008
					I-437
					Sep 29, 2007
					I-401
					Aug 28, 2006
					NCE
					Nov 15, 2007
<u>ARIPIPRAZOLE; ABILIFY</u>					
021436 003	5006528	Oct 20, 2014	DS DP	U-761	I-488
					Mar 01, 2008
					I-437
					Sep 29, 2007
					I-401
					Aug 28, 2006
					NCE
					Nov 15, 2007
<u>ARIPIPRAZOLE; ABILIFY</u>					
021436 004	5006528	Oct 20, 2014	DS DP	U-761	I-488
					Mar 01, 2008
					I-437
					Sep 29, 2007
					I-401
					Aug 28, 2006
					NCE
					Nov 15, 2007
<u>ARIPIPRAZOLE; ABILIFY</u>					
021436 005	5006528	Oct 20, 2014	DS DP	U-761	I-488
					Mar 01, 2008
					I-437
					Sep 29, 2007
					I-401
					Aug 28, 2006
					NCE
					Nov 15, 2007
<u>ARIPIPRAZOLE; ABILIFY</u>					
021436 006	5006528	Oct 20, 2014	DS DP	U-761	I-488
					Mar 01, 2008
					I-437
					Sep 29, 2007
					I-401
					Aug 28, 2006
					NCE
					Nov 15, 2007
<u>ARIPIPRAZOLE; ABILIFY</u>					
021713 001	5006528	Oct 20, 2014	DS DP	U-761	I-488
					Mar 01, 2008
					I-401
					Aug 28, 2006
	6977257	Apr 24, 2022	DS	DP	I-437
					Sep 29, 2007
					NCE
					Nov 15, 2007
<u>ARIPIPRAZOLE; ABILIFY</u>					
021729 002	5006528	Oct 20, 2014	DS DP	U-761	I-488
					Mar 01, 2008
					I-437
					Sep 29, 2007
					NCE
					Nov 15, 2007

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>ARIPIRAZOLE; ABILIFY</u>					
021729 003	5006528	Oct 20, 2014	DS DP U-761	I-488 I-437 NCE	Mar 01, 2008 Sep 29, 2007 Nov 15, 2007
<u>ARIPIRAZOLE; ABILIFY</u>					
021729 004	5006528	Oct 20, 2014	DS DP U-761	I-488 I-437 NCE	Mar 01, 2008 Sep 29, 2007 Nov 15, 2007
<u>ARIPIRAZOLE; ABILIFY</u>					
021729 005	5006528	Oct 20, 2014	DS DP U-761	I-488 I-437 NCE	Mar 01, 2008 Sep 29, 2007 Nov 15, 2007
<u>ARIPIRAZOLE; ABILIFY</u>					
021866 001	5006528 7115587	Oct 20, 2014 Jul 21, 2024	DS DP U-763 DS DP U-764	NCE NDF	Nov 15, 2007 Sep 20, 2009
<u>ARSENIC TRIOXIDE; TRISENOX</u>					
021248 001	6723351 6855339 6861076 6884439	Nov 10, 2018 Nov 10, 2018 Nov 10, 2018 Nov 10, 2018	U-573 U-617 U-617 U-651	ODE	Sep 25, 2007
<u>ARTICAINE HYDROCHLORIDE; EPINEPHRINE BITARTRATE; SEPTOCAINE</u>					
022010 001				NP	Mar 30, 2009
<u>ASCORBIC ACID; BIOTIN; CYANOCOBALAMIN; DEXPANTHENOL; ERGOCALCIFEROL; FOLIC ACID; NIACINAMIDE; PYRIDOXINE HYDROCHLORIDE; RIBOFLAVIN PHOSPHATE SODIUM; THIAMINE HYDROCHLORIDE; VITAMIN A; VITAMIN E; M.V.I.-12 (WITHOUT VITAMIN K)</u>					
008809 006				ODE	Sep 09, 2011
<u>ASCORBIC ACID; POLYETHYLENE GLYCOL 3350; POTASSIUM CHLORIDE; SODIUM ASCORBATE; SODIUM CHLORIDE; SODIUM SULFATE; MOVIPREP</u>					
021881 001				NP	Aug 02, 2009
<u>ASPIRIN; DIPYRIDAMOLE; AGGRENOX</u>					
020884 001	6015577	Jan 18, 2017	U-302		
<u>ASPIRIN; PRAVASTATIN SODIUM; PRAVIGARD PAC (COPACKAGED)</u>					
021387 001	5030447 5030447*PED 5180589 5180589*PED 5622985 5622985*PED	Jul 09, 2008 Jan 09, 2009 Jul 09, 2008 Jan 09, 2009 Apr 22, 2014 Oct 22, 2014	U-335 U-335		
<u>ASPIRIN; PRAVASTATIN SODIUM; PRAVIGARD PAC (COPACKAGED)</u>					
021387 002	5030447 5030447*PED 5180589 5180589*PED 5622985 5622985*PED	Jul 09, 2008 Jan 09, 2009 Jul 09, 2008 Jan 09, 2009 Apr 22, 2014 Oct 22, 2014	U-335 U-335		
<u>ASPIRIN; PRAVASTATIN SODIUM; PRAVIGARD PAC (COPACKAGED)</u>					
021387 003	5030447 5030447*PED 5180589 5180589*PED 5622985 5622985*PED	Jul 09, 2008 Jan 09, 2009 Jul 09, 2008 Jan 09, 2009 Apr 22, 2014 Oct 22, 2014	U-335 U-335		
<u>ASPIRIN; PRAVASTATIN SODIUM; PRAVIGARD PAC (COPACKAGED)</u>					
021387 004	5030447 5030447*PED 5180589 5180589*PED 5622985 5622985*PED	Jul 09, 2008 Jan 09, 2009 Jul 09, 2008 Jan 09, 2009 Apr 22, 2014 Oct 22, 2014	U-335 U-335		

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>ASPIRIN; PRAVASTATIN SODIUM; PRAVIGARD PAC (COPACKAGED)</u>					
021387 005	5030447	Jul 09, 2008			
	5030447*PED	Jan 09, 2009			
	5180589	Jul 09, 2008			
	5180589*PED	Jan 09, 2009			
	5622985	Apr 22, 2014	U-335		
	5622985*PED	Oct 22, 2014	U-335		
<u>ASPIRIN; PRAVASTATIN SODIUM; PRAVIGARD PAC (COPACKAGED)</u>					
021387 006	5030447	Jul 09, 2008			
	5030447*PED	Jan 09, 2009			
	5180589	Jul 09, 2008			
	5180589*PED	Jan 09, 2009			
	5622985	Apr 22, 2014	U-335		
	5622985*PED	Oct 22, 2014	U-335		
<u>ATAZANAVIR SULFATE; REYATAZ</u>					
021567 001	5849911	Apr 09, 2017	DS DP	U-167	D-89
	6087383	Dec 21, 2018	DS DP		NCE
<u>ATAZANAVIR SULFATE; REYATAZ</u>					
021567 002	5849911	Apr 09, 2017	DS DP	U-167	D-89
	6087383	Dec 21, 2018	DS DP		NCE
<u>ATAZANAVIR SULFATE; REYATAZ</u>					
021567 003	5849911	Apr 09, 2017	DS DP	U-167	D-89
	6087383	Dec 21, 2018	DS DP		NCE
<u>ATAZANAVIR SULFATE; REYATAZ</u>					
021567 004	5849911	Apr 09, 2017	DS DP	U-167	D-89
	6087383	Dec 21, 2018	DS DP		NCE
<u>ATOMOXETINE HYDROCHLORIDE; STRATTERA</u>					
021411 001	5658590	Nov 26, 2016		U-494	NCE
	5658590*PED	May 26, 2017		U-494	PED
<u>ATOMOXETINE HYDROCHLORIDE; STRATTERA</u>					
021411 002	5658590	Nov 26, 2016		U-494	NCE
	5658590*PED	May 26, 2017		U-494	PED
<u>ATOMOXETINE HYDROCHLORIDE; STRATTERA</u>					
021411 003	5658590	Nov 26, 2016		U-494	NCE
	5658590*PED	May 26, 2017		U-494	PED
<u>ATOMOXETINE HYDROCHLORIDE; STRATTERA</u>					
021411 004	5658590	Nov 26, 2016		U-494	NCE
	5658590*PED	May 26, 2017		U-494	PED
<u>ATOMOXETINE HYDROCHLORIDE; STRATTERA</u>					
021411 005	5658590	Nov 26, 2016		U-494	NCE
	5658590*PED	May 26, 2017		U-494	PED
<u>ATOMOXETINE HYDROCHLORIDE; STRATTERA</u>					
021411 006	5658590	Nov 26, 2016		U-494	NCE
	5658590*PED	May 26, 2017		U-494	PED
<u>ATOMOXETINE HYDROCHLORIDE; STRATTERA</u>					
021411 007	5658590	Nov 26, 2016		U-494	NCE
	5658590*PED	May 26, 2017		U-494	PED
<u>ATOMOXETINE HYDROCHLORIDE; STRATTERA</u>					
021411 008	5658590	Nov 26, 2016		U-494	NCE
	5658590*PED	May 26, 2017		U-494	PED

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>ATORVASTATIN CALCIUM; LIPITOR</u>					
020702 001	4681893	Sep 24, 2009	DS DP	U-161	I-471 Sep 21, 2008
	4681893*PED	Mar 24, 2010		U-161	I-434 Jul 30, 2007
	5273995	Dec 28, 2010	DS DP	U-162	M-36 Jul 30, 2007
	5273995*PED	Jun 28, 2011		U-162	
	5686104	Nov 11, 2014	DP	U-213	
	5686104*PED	May 11, 2015		U-213	
	5969156	Jul 08, 2016	DS		
	5969156*PED	Jan 08, 2017			
	6126971	Jan 19, 2013	DP		
	6126971*PED	Jul 19, 2013			
<u>ATORVASTATIN CALCIUM; LIPITOR</u>					
020702 002	4681893	Sep 24, 2009	DS DP	U-161	I-471 Sep 21, 2008
	4681893*PED	Mar 24, 2010		U-161	I-434 Jul 30, 2007
	5273995	Dec 28, 2010	DS DP	U-162	M-36 Jul 30, 2007
	5273995*PED	Jun 28, 2011		U-162	
	5686104	Nov 11, 2014	DP	U-213	
	5686104*PED	May 11, 2015		U-213	
	5969156	Jul 08, 2016	DS		
	5969156*PED	Jan 08, 2017			
	6126971	Jan 19, 2013	DP		
	6126971*PED	Jul 19, 2013			
<u>ATORVASTATIN CALCIUM; LIPITOR</u>					
020702 003	4681893	Sep 24, 2009	DS DP	U-161	I-471 Sep 21, 2008
	4681893*PED	Mar 24, 2010		U-161	I-434 Jul 30, 2007
	5273995	Dec 28, 2010	DS DP	U-162	M-36 Jul 30, 2007
	5273995*PED	Jun 28, 2011		U-162	
	5686104	Nov 11, 2014	DP	U-213	
	5686104*PED	May 11, 2015		U-213	
	5969156	Jul 08, 2016	DS		
	5969156*PED	Jan 08, 2017			
	6126971	Jan 19, 2013	DP		
	6126971*PED	Jul 19, 2013			
<u>ATORVASTATIN CALCIUM; LIPITOR</u>					
020702 004	4681893	Sep 24, 2009	DS DP	U-161	I-471 Sep 21, 2008
	4681893*PED	Mar 24, 2010		U-161	I-434 Jul 30, 2007
	5273995	Dec 28, 2010	DS DP	U-162	M-36 Jul 30, 2007
	5273995*PED	Jun 28, 2011		U-162	
	5686104	Nov 11, 2014	DP	U-213	
	5686104*PED	May 11, 2015		U-213	
	5969156	Jul 08, 2016	DS		
	5969156*PED	Jan 08, 2017			
	6126971	Jan 19, 2013	DP		
	6126971*PED	Jul 19, 2013			
<u>ATOVAQUONE; MEPRON</u>					
020259 001	4981874	Aug 15, 2009		U-69	
	4981874*PED	Feb 15, 2010		U-69	
	5053432	Oct 01, 2008			
	5053432*PED	Apr 01, 2009			
<u>ATOVAQUONE; MEPRON</u>					
020500 001	4981874	Aug 15, 2009		U-69	PED Jul 05, 2006
	4981874*PED	Feb 15, 2010		U-69	
	5053432	Oct 01, 2008			
	5053432*PED	Apr 01, 2009			
	6649659	Jul 10, 2016	DS DP	U-69	
	6649659*PED	Jan 10, 2017			
<u>ATOVAQUONE; PROGUANIL HYDROCHLORIDE; MALARONE</u>					
021078 001	5053432	Oct 01, 2008			
	5053432*PED	Apr 01, 2009			
	6166046	Nov 25, 2013		U-406	
	6166046*PED	May 25, 2014			
	6291488	Nov 25, 2013		U-406	
	6291488*PED	May 25, 2014			

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>ATOVAQUONE; PROGUANIL HYDROCHLORIDE; MALARONE PEDIATRIC</u>					
021078 002	5053432	Oct 01, 2008		NPP	Dec 02, 2006
	5053432*PED	Apr 01, 2009		PED	Jun 02, 2007
	6166046	Nov 25, 2013			
	6166046*PED	May 25, 2014			
	6291488	Nov 25, 2013	U-406		
	6291488*PED	May 25, 2014			
<u>ATROPINE SULFATE; EDROPHONIUM CHLORIDE; ENLON-PLUS</u>					
019677 001	4952586	Aug 28, 2007	U-54		
<u>ATROPINE SULFATE; EDROPHONIUM CHLORIDE; ENLON-PLUS</u>					
019678 001	4952586	Aug 28, 2007	U-54		
<u>ATROPINE; PRALIDOXIME CHLORIDE; DUODOTE</u>					
021983 001	5092843	Apr 12, 2010	DP		
<u>AVOBENZONE; ECAMSULE; OCTOCRYLENE; ANTHELIOS SX</u>					
021502 001	4585597	Jun 16, 2007	DS DP U-752	NC	Jul 21, 2009
	5587150	Dec 24, 2013	DP U-752		
<u>AZACITIDINE; VIDAZA</u>					
050794 001				ODE	May 19, 2011
<u>AZELAIC ACID; FINACEA</u>					
021470 001	6534070	Nov 18, 2018			
<u>AZELASTINE HYDROCHLORIDE; ASTELIN</u>					
020114 001	5164194	Nov 01, 2010	U-207	D-102	Feb 17, 2009
	5164194	Nov 01, 2010	U-730		
	5164194*PED	May 01, 2011			
<u>AZELASTINE HYDROCHLORIDE; OPTIVAR</u>					
021127 001	5164194	Nov 01, 2010			
	5164194*PED	May 01, 2011			
<u>BACLOFEN; KEMSTRO</u>					
021589 001	6024981	Apr 09, 2018	DP		
	6221392	Apr 09, 2018	DP		
<u>BACLOFEN; KEMSTRO</u>					
021589 002	6024981	Apr 09, 2018	DP		
	6221392	Apr 09, 2018	DP		
<u>BALSALAZIDE DISODIUM; COLAZAL</u>					
020610 001	4412992	Jul 08, 2006		NPP	Dec 20, 2009
	4412992*PED	Jan 08, 2007		PED	Jun 20, 2010
<u>BECLOMETHASONE DIPROPIONATE; QVAR 40</u>					
020911 002	5605674	Feb 25, 2014			
	5683677	Nov 04, 2014			
	5695743	Jul 06, 2010			
	5766573	Nov 28, 2009	U-356		
	5776432	Jul 07, 2015			
	6352684	Nov 28, 2009			
<u>BECLOMETHASONE DIPROPIONATE; QVAR 80</u>					
020911 001	5605674	Feb 25, 2014			
	5683677	Nov 04, 2014			
	5695743	Jul 06, 2010			
	5766573	Nov 28, 2009	U-356		
	5776432	Jul 07, 2015			
	6352684	Nov 28, 2009			
<u>BENAZEPRIL HYDROCHLORIDE; LOTENSIN</u>					
019851 001				M-30	Mar 02, 2007
				PED	Sep 02, 2007
<u>BENAZEPRIL HYDROCHLORIDE; LOTENSIN</u>					
019851 002				M-30	Mar 02, 2007
				PED	Sep 02, 2007
<u>BENAZEPRIL HYDROCHLORIDE; LOTENSIN</u>					
019851 003				M-30	Mar 02, 2007
				PED	Sep 02, 2007

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>BENAZEPRIL HYDROCHLORIDE; LOTENSIN</u>					
019851 004				M-30 PED	Mar 02, 2007 Sep 02, 2007
<u>BETAMETHASONE DIPROPIONATE; DIPROLENE</u>					
019716 001	4775529	May 21, 2007			
<u>BETAMETHASONE DIPROPIONATE; CALCIPOTRIENE HYDRATE; TACLONEX</u>					
021852 001	4866048	Dec 29, 2007	DS DP	U-88	NC Jan 09, 2009
	4866048	Dec 29, 2007	DS DP	U-193	
	5763426	Jun 09, 2015	DS DP		
	6753013	Jan 27, 2020	DP	U-88	
	6753013	Jan 27, 2020	DP	U-193	
<u>BETAMETHASONE VALERATE; LUXIQ</u>					
020934 001	6126920	Mar 01, 2016		U-484	
	7078058	Mar 01, 2016	DS DP		
<u>BETAXOLOL HYDROCHLORIDE; BETOPTIC S</u>					
019845 001	4911920	Mar 27, 2007			
	4911920*PED	Sep 27, 2007			
<u>BETAXOLOL HYDROCHLORIDE; PILOCARPINE HYDROCHLORIDE; BETOPTIC PILO</u>					
020619 001	5635172	Jun 03, 2014		U-191	
<u>BEXAROTENE; TARGRETIN</u>					
021055 001	5780676	Jul 14, 2015		U-509	ODE Dec 29, 2006
	5962731	Oct 05, 2016		U-475	
	6043279	Apr 22, 2012		U-509	
	6320074	Apr 22, 2012	DS	U-509	
<u>BEXAROTENE; TARGRETIN</u>					
021056 001	5780676	Jul 14, 2015		U-510	ODE Dec 29, 2006
	5962731	Oct 05, 2016			
	6043279	Apr 22, 2012		U-510	
	6320074	Apr 22, 2012	DS	U-510	
<u>BICALUTAMIDE; CASODEX</u>					
020498 001	4636505	Oct 01, 2008			
<u>BIMATOPROST; LUMIGAN</u>					
021275 001	5688819	Aug 19, 2014		U-446	
	6403649	Sep 21, 2012	DS DP	U-446	
<u>BISACODYL; POLYETHYLENE GLYCOL 3350; POTASSIUM CHLORIDE; SODIUM BICARBONATE; SODIUM CHLORIDE; HALFLYTELY</u>					
021551 001				NP	May 10, 2007
<u>BIVALIRUDIN; ANGIOMAX</u>					
020873 001	5196404	Mar 23, 2010		I-486 I-458	Nov 30, 2008 Jun 13, 2008
<u>BORTEZOMIB; VELCADE</u>					
021602 001	5780454	Oct 28, 2014		I-521	Dec 08, 2009
	6083903	Oct 28, 2014		U-515	I-452 NCE ODE ODE
	6297217	Oct 28, 2014		U-515	
	6617317	Oct 28, 2014	DS DP		
	6713446	Jan 25, 2022	DP		
	6747150	Oct 28, 2014	DP		
	6958319	Jan 25, 2022	DP		
<u>BOSENTAN; TRACLEER</u>					
021290 001	5292740	Nov 20, 2015		NCE ODE	Nov 20, 2006 Nov 20, 2008
<u>BOSENTAN; TRACLEER</u>					
021290 002	5292740	Nov 20, 2015		NCE ODE	Nov 20, 2006 Nov 20, 2008

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>BRIMONIDINE TARTRATE; ALPHAGAN P</u>					
021262 001	5424078	Jun 13, 2012			
	5424078*PED	Dec 13, 2012			
	6562873	Jul 10, 2021			
	6562873*PED	Jan 10, 2022			
	6627210	Jul 18, 2021	DP		
	6627210*PED	Jan 18, 2022			
	6641834	Jul 28, 2021	DP		
	6641834*PED	Jan 28, 2022			
	6673337	Jul 26, 2021	DP		
	6673337*PED	Jan 26, 2022			
<u>BRIMONIDINE TARTRATE; ALPHAGAN P</u>					
021770 001	5424078	Jun 13, 2012	DP	NP	Aug 19, 2008
	5424078*PED	Dec 13, 2012			
	6562873	Jul 10, 2021	DP		
	6562873*PED	Jan 10, 2022			
	6627210	Jul 18, 2021	DP		
	6627210*PED	Jan 18, 2022			
	6641834	Jul 28, 2021	DP		
	6641834*PED	Jan 28, 2022			
	6673337	Jul 26, 2021	DP		
	6673337*PED	Jan 26, 2022			
<u>BRINZOLAMIDE; AZOPT</u>					
020816 001	5240923	Aug 31, 2010	DS DP U-224	M-54	Sep 28, 2009
	5240923*PED	Mar 01, 2011		PED	Mar 28, 2010
	5378703	Apr 01, 2012	DS DP U-224		
	5378703*PED	Oct 01, 2012			
	5461081	Oct 24, 2012	DP U-225		
	5461081*PED	Apr 24, 2013			
<u>BROMFENAC SODIUM; XIBROM</u>					
021664 001				I-485	Jan 27, 2009
				NP	Mar 24, 2008
<u>BUDESONIDE; BUDESONIDE</u>					
021949 001	4907583	Mar 13, 2007	DP	NP	Jul 12, 2009
	6027714	Jan 09, 2018	DP U-787		
	6142145	May 08, 2018	DP		
	6287540	Jan 09, 2018	DP		
	7143764	Mar 13, 2018	DP		
<u>BUDESONIDE; BUDESONIDE</u>					
021949 002	4907583	Mar 13, 2007	DP	NP	Jul 12, 2009
	6027714	Jan 09, 2018	DP U-787		
	6142145	May 08, 2018	DP		
	6287540	Jan 09, 2018	DP		
	7143764	Mar 13, 2018	DP		
<u>BUDESONIDE; ENTOCORT EC</u>					
021324 001	5643602	Jul 01, 2014		U-655	
	5643602*PED	Jan 01, 2015	DP	I-454	Apr 29, 2008
	6423340	Nov 15, 2010			
	6423340*PED	May 11, 2011			
<u>BUDESONIDE; PULMICORT</u>					
020441 002	4668218*PED	Oct 11, 2006			
	4907583	Mar 13, 2007			
	4907583*PED	Sep 13, 2007			
<u>BUDESONIDE; PULMICORT</u>					
020441 003	4668218*PED	Oct 11, 2006			
	4907583	Mar 13, 2007			
	4907583*PED	Sep 13, 2007			
<u>BUDESONIDE; PULMICORT RESPULES</u>					
020929 001	4787536*PED	Aug 27, 2006			
	6598603	Dec 23, 2018		U-529	
	6598603*PED	Jun 23, 2019			
	6899099	Dec 23, 2018		U-751	
	6899099*PED	Jun 23, 2019			

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE	
BUDESONIDE; PULMICORT RESPULES						
020929 002	4787536*PED	Aug 27, 2006				
	6598603	Dec 23, 2018		U-529		
	6598603*PED	Jun 23, 2019				
	6899099	Dec 23, 2018		U-751		
	6899099*PED	Jun 23, 2019				
BUDESONIDE; PULMICORT RESPULES						
020929 003	4787536*PED	Aug 27, 2006				
	6598603	Dec 23, 2018		U-529		
	6598603*PED	Jun 23, 2019				
BUDESONIDE; RHINOCORT						
020746 001	6291445	Apr 29, 2017				
	6291445*PED	Oct 29, 2017				
	6686346	Apr 29, 2017	DP	U-557		
	6986904	Apr 29, 2017	DP	U-699		
BUDESONIDE; RHINOCORT						
020746 002	6291445	Apr 29, 2017				
	6291445*PED	Oct 29, 2017				
	6686346	Apr 29, 2017	DP	U-557		
	6986904	Apr 29, 2017	DP	U-699		
BUDESONIDE; FORMOTEROL FUMARATE; SYMBICORT						
021929 001	5674860	Oct 07, 2014	DP	U-754	NC	Jul 21, 2009
	5972919	Dec 17, 2012	DP	U-754		
	6123924	Sep 26, 2017	DP			
	6641800	Sep 26, 2017	DP			
BUDESONIDE; FORMOTEROL FUMARATE; SYMBICORT						
021929 002	5674860	Oct 07, 2014	DP	U-754	NC	Jul 21, 2009
	5972919	Dec 17, 2012	DP	U-754		
	6123924	Sep 26, 2017	DP			
	6641800	Sep 26, 2017	DP			
BUPRENORPHINE HYDROCHLORIDE; SUBUTEX						
020732 002				ODE	Oct 08, 2009	
BUPRENORPHINE HYDROCHLORIDE; SUBUTEX						
020732 003				ODE	Oct 08, 2009	
BUPRENORPHINE HYDROCHLORIDE; NALOXONE HYDROCHLORIDE; SUBOXONE						
020733 001				ODE	Oct 08, 2009	
BUPRENORPHINE HYDROCHLORIDE; NALOXONE HYDROCHLORIDE; SUBOXONE						
020733 002				ODE	Oct 08, 2009	
BUPROPION HYDROCHLORIDE; WELLBUTRIN SR						
020358 001	5358970	Aug 12, 2013				
	5427798	Aug 12, 2013				
	5731000	Aug 12, 2013				
	5763493	Aug 12, 2013				
BUPROPION HYDROCHLORIDE; WELLBUTRIN SR						
020358 002	5358970	Aug 12, 2013				
	5427798	Aug 12, 2013				
	5731000	Aug 12, 2013				
	5763493	Aug 12, 2013				
BUPROPION HYDROCHLORIDE; WELLBUTRIN SR						
020358 003	5358970	Aug 12, 2013				
	5427798	Aug 12, 2013				
	5731000	Aug 12, 2013				
	5763493	Aug 12, 2013				
BUPROPION HYDROCHLORIDE; WELLBUTRIN SR						
020358 004	5358970	Aug 12, 2013				
	5427798	Aug 12, 2013				
	5731000	Aug 12, 2013				
	5763493	Aug 12, 2013				
BUPROPION HYDROCHLORIDE; WELLBUTRIN XL						
021515 001	6096341	Oct 30, 2018		I-497	Jun 12, 2009	
	6143327	Oct 30, 2018				

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
BUPROPION HYDROCHLORIDE; WELLBUTRIN XL					
021515 002	6096341	Oct 30, 2018		I-497	Jun 12, 2009
	6143327	Oct 30, 2018			
BUPROPION HYDROCHLORIDE; ZYBAN					
020711 002	5358970	Aug 12, 2013			
	5427798	Aug 12, 2013			
	5731000	Aug 12, 2013			
	5763493	Aug 12, 2013			
BUPROPION HYDROCHLORIDE; ZYBAN					
020711 003	5358970	Aug 12, 2013			
	5427798	Aug 12, 2013			
	5731000	Aug 12, 2013			
	5763493	Aug 12, 2013			
BUSPIRONE HYDROCHLORIDE; BUSPAR					
021190 001	5015646	May 14, 2008			
	5015646*PED	Nov 14, 2008			
BUSPIRONE HYDROCHLORIDE; BUSPAR					
021190 002	5015646	May 14, 2008			
	5015646*PED	Nov 14, 2008			
BUSPIRONE HYDROCHLORIDE; BUSPAR					
021190 003	5015646	May 14, 2008			
	5015646*PED	Nov 14, 2008			
BUSPIRONE HYDROCHLORIDE; BUSPAR					
021190 004	5015646	May 14, 2008			
	5015646*PED	Nov 14, 2008			
BUSULFAN; BUSULFEX					
020954 001	5430057	Sep 30, 2013	U-263	PED	Aug 04, 2006
	5430057*PED	Mar 30, 2014	U-263		
	5559148	Sep 30, 2013	U-264		
	5559148*PED	Mar 30, 2014	U-264		
BUTENAFINE HYDROCHLORIDE; MENTAX					
020524 001	5021458	Oct 18, 2010	U-177		
BUTOCONAZOLE NITRATE; GYNAZOLE-1					
019881 001	5266329	Nov 30, 2010	U-457		
CAFFEINE CITRATE; CAFCIT					
020793 001				ODE	Sep 21, 2006
CALCIPOTRIENE; DOVONEX					
020273 001	4866048	Dec 29, 2007			
CALCIPOTRIENE; DOVONEX					
020554 001	4866048	Dec 29, 2007			
	5763426	Jun 09, 2015	DS DP		
CALCIPOTRIENE; DOVONEX					
020611 001	4866048	Dec 29, 2007			
	5763426	Jun 09, 2015	DS DP		
CALCITONIN SALMON RECOMBINANT; FORTICAL					
021406 001	6440392	Feb 02, 2021	DP U-227		
CALCITONIN, SALMON; MIACALCIN					
020313 002	5733569	Mar 31, 2015	U-227		
	5759565	Mar 31, 2015			
CALCITRIOL; CALCIJEX					
018874 001	6051567	Aug 02, 2019			
	6051567*PED	Feb 02, 2020			
	6265392	Aug 02, 2019			
	6265392*PED	Feb 02, 2020			
	6274169	Aug 02, 2019			
	6274169*PED	Feb 02, 2020			

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE	
CALCITRIOL; CALCIJEX						
018874 002	6051567	Aug 02, 2019				
	6051567*PED	Feb 02, 2020				
	6265392	Aug 02, 2019				
	6265392*PED	Feb 02, 2020				
	6274169	Aug 02, 2019				
	6274169*PED	Feb 02, 2020				
CALCIUM ACETATE; PHOSLO						
019976 001	4870105	Apr 07, 2007		U-381		
CALCIUM ACETATE; PHOSLO						
021160 001	4870105	Apr 07, 2007		U-381		
CALCIUM ACETATE; PHOSLO						
021160 002	4870105	Apr 07, 2007		U-381		
	6576665	Apr 03, 2021				
CALCIUM ACETATE; PHOSLO GELCAPS						
021160 003	4870105	Apr 07, 2007		U-381		
	6576665	Apr 03, 2021				
CALCIUM CARBONATE; FAMOTIDINE; MAGNESIUM HYDROXIDE; PEPCID COMPLETE						
020958 001	5229137	May 16, 2012		U-349		
	5229137*PED	Nov 16, 2012				
	5817340	Dec 01, 2012				
	5817340*PED	Jun 01, 2013				
	5989588	Sep 30, 2015		U-349		
	5989588*PED	Mar 30, 2016		U-349		
CALCIUM CARBONATE; RISEDRONATE SODIUM; ACTONEL WITH CALCIUM (COPACKAGED)						
021823 001	5583122	Dec 10, 2013	DS DP	U-353	M-52	
	5994329	Jul 17, 2018		U-353		
	6015801	Jul 17, 2018		U-353		
	6096342	Nov 21, 2011		DP		
	6165513	Jun 10, 2018		DP		
	6432932	Jul 17, 2018		U-595		
	6465443	Aug 14, 2018		DP		
CANDESARTAN CILEXETIL; ATACAND						
020838 001	5196444	Jun 04, 2012	DS DP	U-660	I-456	May 18, 2008
	5196444	Jun 04, 2012	DS DP	U-3	I-455	May 18, 2008
	5534534	Jul 09, 2013		DP	I-448	May 18, 2008
	5705517	Apr 18, 2011	DS DP	U-660	I-448	Feb 22, 2008
CANDESARTAN CILEXETIL; ATACAND						
020838 002	5196444	Jun 04, 2012	DS DP	U-3	I-456	May 18, 2008
	5196444	Jun 04, 2012	DS DP	U-660	I-455	May 18, 2008
	5534534	Jul 09, 2013		DP	I-448	May 18, 2008
	5705517	Apr 18, 2011	DS DP	U-660	I-448	Feb 22, 2008
CANDESARTAN CILEXETIL; ATACAND						
020838 003	5196444	Jun 04, 2012	DS DP	U-3	I-456	May 18, 2008
	5196444	Jun 04, 2012	DS DP	U-660	I-455	May 18, 2008
	5534534	Jul 09, 2013		DP	I-448	May 18, 2008
	5705517	Apr 18, 2011	DS DP	U-660	I-448	Feb 22, 2008
CANDESARTAN CILEXETIL; ATACAND						
020838 004	5196444	Jun 04, 2012	DS DP	U-660	I-456	May 18, 2008
	5196444	Jun 04, 2012	DS DP	U-3	I-455	May 18, 2008
	5534534	Jul 09, 2013		DP	I-448	May 18, 2008
	5705517	Apr 18, 2011	DS DP	U-660	I-448	Feb 22, 2008
CANDESARTAN CILEXETIL; HYDROCHLOROTHIAZIDE; ATACAND HCT						
021093 001	5196444	Jun 04, 2012		U-3		
	5534534	Jul 09, 2013				
	5705517	Apr 18, 2011		U-3		
	5721263	Feb 24, 2015		U-3		
	5958961	Jun 06, 2014		U-3		
CANDESARTAN CILEXETIL; HYDROCHLOROTHIAZIDE; ATACAND HCT						
021093 002	5196444	Jun 04, 2012		U-3		
	5534534	Jul 09, 2013				
	5705517	Apr 18, 2011		U-3		
	5721263	Feb 24, 2015		U-3		
	5958961	Jun 06, 2014		U-3		

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>CAPECITABINE; XELODA</u>					
020896 001	4966891 5472949	Jan 13, 2011 Dec 14, 2013	U-272 U-271	I-461	Jun 15, 2008
<u>CAPECITABINE; XELODA</u>					
020896 002	4966891 5472949	Jan 13, 2011 Dec 14, 2013	U-272 U-271	I-461	Jun 15, 2008
<u>CAPTOPRIL; CAPOTEN</u>					
018343 001	5238924	Aug 24, 2010	U-92		
<u>CAPTOPRIL; CAPOTEN</u>					
018343 002	5238924	Aug 24, 2010	U-92		
<u>CAPTOPRIL; CAPOTEN</u>					
018343 003	5238924	Aug 24, 2010	U-92		
<u>CAPTOPRIL; CAPOTEN</u>					
018343 004	5238924	Aug 24, 2010	U-92		
<u>CAPTOPRIL; CAPOTEN</u>					
018343 005	5238924	Aug 24, 2010	U-92		
<u>CAPTOPRIL; CAPOTEN</u>					
018343 006	5238924	Aug 24, 2010	U-92		
<u>CAPTOPRIL; CAPOTEN</u>					
018343 007	5238924	Aug 24, 2010	U-92		
<u>CARBAMAZEPINE; CARBATROL</u>					
020712 001	5326570 5912013	Jul 05, 2011 Jun 15, 2016	U-215 U-277		
<u>CARBAMAZEPINE; CARBATROL</u>					
020712 002	5326570 5912013	Jul 05, 2011 Jun 15, 2016	U-215 U-277		
<u>CARBAMAZEPINE; CARBATROL</u>					
020712 003	5326570 5912013	Jul 05, 2011 Jun 15, 2016	U-215 U-277		
<u>CARBAMAZEPINE; EQUETRO</u>					
021710 001	5326570 5912013 6977253	Jul 23, 2011 Jun 15, 2016 May 19, 2024	DP U-627 DP U-693	NP	Dec 10, 2007
<u>CARBAMAZEPINE; EQUETRO</u>					
021710 002	5326570 5912013 6977253	Jul 23, 2011 Jun 15, 2016 May 19, 2024	DP U-627 DP U-693	NP	Dec 10, 2007
<u>CARBAMAZEPINE; EQUETRO</u>					
021710 003	5326570 5912013 6977253	Jul 23, 2011 Jun 15, 2016 May 19, 2024	DP U-627 DP U-693	NP	Dec 10, 2007
<u>CARBAMAZEPINE; TEGRETOL-XR</u>					
020234 001	5284662 RE34990	Feb 08, 2011 Jul 29, 2007			
<u>CARBAMAZEPINE; TEGRETOL-XR</u>					
020234 002	5284662 RE34990	Feb 08, 2011 Jul 29, 2007			
<u>CARBAMAZEPINE; TEGRETOL-XR</u>					
020234 003	5284662 RE34990	Feb 08, 2011 Jul 29, 2007			
<u>CARBIDOPA; ENTACAPONE; LEVODOPA; STALEVO 100</u>					
021485 002	4963590 5112861 5135950 5446194 6500867 6797732	Nov 27, 2007 May 12, 2009 Oct 31, 2010 Oct 19, 2013 Jun 29, 2020 Jun 29, 2020	DP U-219 DP U-219 DS DP U-219 DS DP U-219 DP		

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>CARBIDOPA; ENTACAPONE; LEVODOPA; STALEVO 150</u>					
021485 003	4963590	Nov 27, 2007	DP	U-219	
	5112861	May 12, 2009		U-219	
	5135950	Oct 31, 2010	DS	DP	U-219
	5446194	Oct 19, 2013	DS		
	6500867	Jun 29, 2020	DP	U-219	
	6797732	Jun 29, 2020	DP		
<u>CARBIDOPA; ENTACAPONE; LEVODOPA; STALEVO 50</u>					
021485 001	4963590	Nov 27, 2007	DP	U-219	
	5112861	May 12, 2009		U-219	
	5135950	Oct 31, 2010	DS	DP	U-219
	5446194	Oct 19, 2013	DS		
	6500867	Jun 29, 2020	DP	U-219	
	6797732	Jun 29, 2020	DP		
<u>CARMUSTINE; GLIADEL</u>					
020637 001	4757128	Aug 01, 2006		ODE	Feb 25, 2010
	4789724	Aug 01, 2006			
<u>CARVEDILOL; COREG</u>					
020297 001	4503067	Mar 05, 2007		U-3	M-56
	4503067*PED	Sep 05, 2007			PED
	5760069	Jun 07, 2015		U-233	
	5760069*PED	Dec 07, 2015			
	5902821	Feb 07, 2016		U-313	
	5902821*PED	Aug 07, 2016			
<u>CARVEDILOL; COREG</u>					
020297 002	4503067	Mar 05, 2007		U-3	M-56
	4503067*PED	Sep 05, 2007			PED
	5760069	Jun 07, 2015		U-233	
	5760069*PED	Dec 07, 2015			
	5902821	Feb 07, 2016		U-313	
	5902821*PED	Aug 07, 2016			
<u>CARVEDILOL; COREG</u>					
020297 003	4503067	Mar 05, 2007		U-3	M-56
	4503067*PED	Sep 05, 2007			PED
	5760069	Jun 07, 2015		U-233	
	5760069*PED	Dec 07, 2015			
	5902821	Feb 07, 2016		U-313	
	5902821*PED	Aug 07, 2016			
<u>CARVEDILOL; COREG</u>					
020297 004	4503067	Mar 05, 2007		U-3	M-56
	4503067*PED	Sep 05, 2007			PED
	5760069	Jun 07, 2015		U-233	
	5760069*PED	Dec 07, 2015			
	5902821	Feb 07, 2016		U-313	
	5902821*PED	Aug 07, 2016			
<u>CARVEDILOL PHOSPHATE; COREG CR</u>					
022012 001	4503067	Mar 05, 2007	DS	DP	U-3
	4503067*PED	Sep 05, 2007			
	5760069	Jun 07, 2015			U-777
	5760069*PED	Dec 07, 2015			
	5902821	Feb 07, 2016			U-313
	5902821	Feb 07, 2016			U-777
	5902821*PED	Aug 07, 2016			
	6022562	Oct 17, 2015		DP	
	6022562*PED	Apr 17, 2016			
<u>CARVEDILOL PHOSPHATE; COREG CR</u>					
022012 002	4503067	Mar 05, 2007	DS	DP	U-3
	4503067*PED	Sep 05, 2007			
	5760069	Jun 07, 2015			U-777
	5760069*PED	Dec 07, 2015			
	5902821	Feb 07, 2016			U-777
	5902821	Feb 07, 2016			U-313
	5902821*PED	Aug 07, 2016			
	6022562	Oct 17, 2015		DP	
	6022562*PED	Apr 17, 2016			

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE		PATENT CODES			EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE		
<u>CARVEDILOL PHOSPHATE; COREG CR</u>										
022012 003	4503067	Mar	05, 2007	DS	DP	U-3	M-56	Aug	28, 2009	
	4503067*PED	Sep	05, 2007			NDF	Oct	20, 2009		
	5760069	Jun	07, 2015			U-777	PED	Feb	28, 2010	
	5760069*PED	Dec	07, 2015			PED	Apr	20, 2010		
	5902821	Feb	07, 2016			U-313				
	5902821	Feb	07, 2016			U-777				
	5902821*PED	Aug	07, 2016							
	6022562	Oct	17, 2015			DP				
6022562*PED	Apr	17, 2016								
<u>CARVEDILOL PHOSPHATE; COREG CR</u>										
022012 004	4503067	Mar	05, 2007	DS	DP	U-3	M-56	Aug	28, 2009	
	4503067*PED	Sep	05, 2007			NDF	Oct	20, 2009		
	5760069	Jun	07, 2015			U-777	PED	Feb	28, 2010	
	5760069*PED	Dec	07, 2015			PED	Apr	20, 2010		
	5902821	Feb	07, 2016			U-313				
	5902821	Feb	07, 2016			U-777				
	5902821*PED	Aug	07, 2016							
	6022562	Oct	17, 2015			DP				
6022562*PED	Apr	17, 2016								
<u>CASPOFUNGIN ACETATE; CANCIDAS</u>										
021227 001	5378804	Mar	16, 2013	DS	DP	U-607	I-438	Sep	29, 2007	
	5514650	Jan	26, 2015							
	5792746	Mar	16, 2013			DS	DP	U-607		
	5952300	Mar	28, 2017			DP				
	6136783	Mar	28, 2017			U-607				
<u>CASPOFUNGIN ACETATE; CANCIDAS</u>										
021227 002	5378804	Mar	16, 2013	DS	DP	U-607	I-438	Sep	29, 2007	
	5514650	Jan	26, 2015							
	5792746	Mar	16, 2013			DS	DP	U-607		
	5952300	Mar	28, 2017			DP				
	6136783	Mar	28, 2017			U-607				
<u>CEFEDITOREN PIVOXIL; SPECTRACEF</u>										
021222 001	4839350	Apr	14, 2009	DS	DP		NCE	Aug	29, 2006	
	5958915	Oct	14, 2016							
<u>CELECOXIB; CELEBREX</u>										
020998 001	5466823	Nov	30, 2013	DS	DP		I-466	Jul	29, 2008	
	5466823*PED	May	30, 2014				NPP	Dec	15, 2009	
	5563165	Nov	30, 2013			DP	PED	Jun	15, 2010	
	5563165*PED	May	30, 2014				PED	Jan	29, 2009	
	5760068	Jun	02, 2015			U-672				
	5760068	Jun	02, 2015			U-299				
	5760068*PED	Dec	02, 2015							
	5972986	Oct	14, 2017			U-299				
	5972986*PED	Apr	14, 2018							
<u>CELECOXIB; CELEBREX</u>										
020998 002	5466823	Nov	30, 2013	DS	DP		I-466	Jul	29, 2008	
	5466823*PED	May	30, 2014				NPP	Dec	15, 2009	
	5563165	Nov	30, 2013			DP	PED	Jun	15, 2010	
	5563165*PED	May	30, 2014				PED	Jan	29, 2009	
	5760068	Jun	02, 2015			U-299				
	5760068	Jun	02, 2015			U-672				
	5760068*PED	Dec	02, 2015							
	5972986	Oct	14, 2017			U-299				
	5972986*PED	Apr	14, 2018							
<u>CELECOXIB; CELEBREX</u>										
020998 003	5466823	Nov	30, 2013	DS	DP		I-466	Jul	29, 2008	
	5466823*PED	May	30, 2014				NPP	Dec	15, 2009	
	5563165	Nov	30, 2013				PED	Jun	15, 2010	
	5563165*PED	May	30, 2014				PED	Jan	29, 2009	
	5760068	Jun	02, 2015			U-299				
	5760068	Jun	02, 2015			U-672				
	5760068*PED	Dec	02, 2015							
	5972986	Oct	14, 2017			U-299				
	5972986*PED	Apr	14, 2018							
<u>CERIVASTATIN SODIUM; BAYCOL</u>										
020740 001	5006530	Jun	26, 2011							
	5177080	Nov	26, 2011							

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>CERIVASTATIN SODIUM; BAYCOL</u>					
020740 002	5006530	Jun 26, 2011			
	5177080	Nov 26, 2011			
<u>CERIVASTATIN SODIUM; BAYCOL</u>					
020740 003	5006530	Jun 26, 2011			
	5177080	Nov 26, 2011			
<u>CERIVASTATIN SODIUM; BAYCOL</u>					
020740 004	5006530	Jun 26, 2011			
	5177080	Nov 26, 2011			
<u>CERIVASTATIN SODIUM; BAYCOL</u>					
020740 005	5006530	Jun 26, 2011			
	5177080	Nov 26, 2011			
<u>CERIVASTATIN SODIUM; BAYCOL</u>					
020740 006	5006530	Jun 26, 2011			
	5177080	Jan 26, 2011			
<u>CETIRIZINE HYDROCHLORIDE; ZYRTEC</u>					
019835 001	4525358	Jun 25, 2007	DS DP	U-565	
	4525358*PED	Dec 25, 2007			
<u>CETIRIZINE HYDROCHLORIDE; ZYRTEC</u>					
019835 002	4525358	Jun 25, 2007	DS DP	U-565	
	4525358*PED	Dec 25, 2007			
<u>CETIRIZINE HYDROCHLORIDE; ZYRTEC</u>					
020346 001	4525358	Jun 25, 2007			
	4525358*PED	Dec 25, 2007			
<u>CETIRIZINE HYDROCHLORIDE; ZYRTEC</u>					
021621 001	4525358	Jun 25, 2007	DS DP	U-565	
	4525358*PED	Dec 25, 2007			
	6455533	Jul 02, 2018	DP		
<u>CETIRIZINE HYDROCHLORIDE; ZYRTEC</u>					
021621 002	4525358	Jun 25, 2007	DS DP	U-565	
	4525358*PED	Dec 25, 2007			
	6455533	Jul 02, 2018	DP		
<u>CETIRIZINE HYDROCHLORIDE; PSEUDOEPHEDRINE HYDROCHLORIDE; ZYRTEC-D 12 HOUR</u>					
021150 001	4525358	Jun 25, 2007		U-295	
	4525358*PED	Dec 25, 2007		U-295	
	6469009	Jul 13, 2019		U-295	
	6489329	Apr 08, 2016			
	7014867	Jun 10, 2022	DP		
<u>CETRORELIX; CETROTIDE</u>					
021197 001	4800191	Jul 17, 2007			
	5198533	Jul 17, 2007			
	6319192	Apr 23, 2018		U-426	
	6863891	Feb 19, 2013		U-426	
<u>CETRORELIX; CETROTIDE</u>					
021197 002	4800191	Jul 17, 2007			
	5198533	Jul 17, 2007			
	6319192	Apr 23, 2018		U-426	
	6863891	Feb 19, 2013		U-426	
<u>CEVIMELINE HYDROCHLORIDE; EVOXAC</u>					
020989 002	4855290	Aug 30, 2009			
	5340821	Jul 07, 2013		U-309	
<u>CHLORHEXIDINE GLUCONATE; CHLORHEXIDINE GLUCONATE</u>					
021669 001	7066916	Feb 17, 2024	U-737	NDF	Apr 25, 2008
<u>CHLORHEXIDINE GLUCONATE; ISOPROPYL ALCOHOL; CHLORAPREP ONE-STEP</u>					
020832 001	5538353	Aug 25, 2015	DP		
	5690958	Sep 30, 2016	DP		
	5752363	Apr 22, 2017	DP		
	5772346	Apr 22, 2017	DP		
	6536975	Nov 10, 2020	DP		
	D386849	Nov 25, 2011	DP		
	D396911	Aug 11, 2012	DP		

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>CHLORHEXIDINE GLUCONATE; ISOPROPYL ALCOHOL; CHLORAPREP ONE-STEP</u>					
020832 004	5690958	Sep 30, 2016	DP		
	6536975	Nov 10, 2020	DP		
<u>CHLORHEXIDINE GLUCONATE; ISOPROPYL ALCOHOL; CHLORAPREP ONE-STEP FREPP</u>					
020832 003	5538353	Aug 25, 2015	DP		
	5690958	Sep 30, 2016	DP		
	5752363	Apr 22, 2017	DP		
	5772346	Apr 22, 2017	DP		
	D386849	Nov 25, 2011	DP		
	D396911	Aug 11, 2012	DP		
<u>CHLORHEXIDINE GLUCONATE; ISOPROPYL ALCOHOL; CHLORAPREP ONE-STEP SEPP</u>					
021555 001	5690958	Sep 30, 2016	DP		
<u>CHLORHEXIDINE GLUCONATE; ISOPROPYL ALCOHOL; CHLORAPREP SINGLE SWABSTICK</u>					
021555 002	5690958	Sep 30, 2016	DP		
<u>CHLORHEXIDINE GLUCONATE; ISOPROPYL ALCOHOL; CHLORAPREP WITH TINT</u>					
020832 002	5538353	Aug 25, 2015	DP	NP	May 03, 2008
	5690958	Sep 30, 2016	DP		
	6536975	Nov 10, 2020	DP		
	6729786	Mar 14, 2023	DP		
	6991393	Mar 14, 2023	DP		
<u>CHLORHEXIDINE GLUCONATE; ISOPROPYL ALCOHOL; CHLORAPREP WITH TINT</u>					
020832 005	5690958	Sep 30, 2016	DP		
	6536975	Nov 10, 2020	DP		
	6729786	Mar 14, 2023	DP		
	6991393	Jan 31, 2024	DP		
<u>CHLORHEXIDINE GLUCONATE; ISOPROPYL ALCOHOL; CHLORASCRUB MAXI SWABSTICK</u>					
021524 003	D468424	Jan 07, 2017		NP	Jun 03, 2008
<u>CHLORHEXIDINE GLUCONATE; ISOPROPYL ALCOHOL; CHLORASCRUB SWAB</u>					
021524 001				NP	Jun 03, 2008
<u>CHLORHEXIDINE GLUCONATE; ISOPROPYL ALCOHOL; CHLORASCRUB SWABSTICK</u>					
021524 002				NP	Jun 03, 2008
<u>CHLORPHENIRAMINE MALEATE; EFIDAC 24 CHLORPHENIRAMINE MALEATE</u>					
019746 002	4857330	Aug 15, 2006			
<u>CHLORPHENIRAMINE MALEATE; IBUPROFEN; PSEUDOEPHEDRINE HYDROCHLORIDE; CHILDREN'S ADVIL ALLERGY SINUS</u>					
021587 001				NP	Feb 24, 2007
<u>CHLORTHALIDONE; THALITONE</u>					
019574 001	4933360	Jun 12, 2007			
<u>CHORIOGONADOTROPIN ALFA; OVIDREL</u>					
021149 001	5767251	Jun 16, 2015			
<u>CHORIOGONADOTROPIN ALFA; OVIDREL</u>					
021149 002	5767251	Jun 16, 2015	DS		
	6706681	Mar 16, 2021	DP		
<u>CICLESONIDE; OMNARIS</u>					
022004 001				NCE	Oct 20, 2011
<u>CICLOPIROX; LOPROX</u>					
020519 001	7018656	Sep 05, 2018	DP		
	7026337	Apr 02, 2018		U-714	
<u>CICLOPIROX; PENLAC</u>					
021022 001	4957730	Sep 18, 2007		U-379	
<u>CIDOFOVIR; VISTIDE</u>					
020638 001	5142051	Jun 26, 2010			
<u>CINACALCET HYDROCHLORIDE; SENSIPAR</u>					
021688 001	6011068	Dec 14, 2016	DS DP	NCE	Mar 08, 2009
	6031003	Dec 14, 2016		U-559	Mar 08, 2011
	6211244	Oct 23, 2015	DS DP	U-560	
	6313146	Dec 14, 2016	DS DP		

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>CINACALCET HYDROCHLORIDE; SENSIPAR</u>					
021688 002	6011068	Dec 14, 2016	DS DP	NCE	Mar 08, 2009
	6031003	Dec 14, 2016		ODE	Mar 08, 2011
	6211244	Oct 23, 2015	DS DP U-560		
	6313146	Dec 14, 2016	DS DP		
<u>CINACALCET HYDROCHLORIDE; SENSIPAR</u>					
021688 003	6011068	Dec 14, 2016	DS DP	NCE	Mar 08, 2009
	6031003	Dec 14, 2016		ODE	Mar 08, 2011
	6211244	Oct 23, 2015	DS DP U-559		
	6313146	Dec 14, 2016	DS DP U-560		
<u>CIPROFLOXACIN; CIPRO</u>					
019847 001	4808583*PED	Aug 28, 2006		I-421 PED	Mar 25, 2007 Sep 25, 2007
<u>CIPROFLOXACIN; CIPRO</u>					
019847 002				I-421 PED	Mar 25, 2007 Sep 25, 2007
<u>CIPROFLOXACIN; CIPRO</u>					
019847 003				I-421 PED	Mar 25, 2007 Sep 25, 2007
<u>CIPROFLOXACIN; CIPRO</u>					
020780 001	5695784	Dec 09, 2014		I-421	Mar 25, 2007
	5695784*PED	Jun 09, 2015		PED	Sep 25, 2007
	6136347	Jan 06, 2013	U-362		
	6136347*PED	Jul 06, 2013			
<u>CIPROFLOXACIN; CIPRO</u>					
020780 002	5695784	Dec 09, 2014		I-421	Mar 25, 2007
	5695784*PED	Jun 09, 2015		PED	Sep 25, 2007
	6136347	Jan 06, 2013	U-362		
	6136347*PED	Jul 06, 2013			
<u>CIPROFLOXACIN; CIPRO IN DEXTROSE 5% IN PLASTIC CONTAINER</u>					
019857 001	4808583*PED	Aug 28, 2006		I-421	Mar 25, 2007
	4957922	Sep 18, 2007		PED	Sep 25, 2007
	4957922*PED	Mar 18, 2008			
<u>CIPROFLOXACIN; CIPRO IN SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER</u>					
019858 001	4808583*PED	Aug 28, 2006			
	4957922	Sep 18, 2007			
	4957922*PED	Mar 18, 2008			
<u>CIPROFLOXACIN HYDROCHLORIDE; CIPRO</u>					
019537 001	5286754	Feb 15, 2011		I-421	Mar 25, 2007
	5286754*PED	Aug 15, 2011		PED	Sep 25, 2007
<u>CIPROFLOXACIN HYDROCHLORIDE; CIPRO</u>					
019537 002	5286754	Feb 15, 2011		I-421	Mar 25, 2007
	5286754*PED	Aug 15, 2011		PED	Sep 25, 2007
<u>CIPROFLOXACIN HYDROCHLORIDE; CIPRO</u>					
019537 003	5286754	Feb 15, 2011		I-421	Mar 25, 2007
	5286754*PED	Aug 15, 2011		PED	Sep 25, 2007
<u>CIPROFLOXACIN HYDROCHLORIDE; CIPRO</u>					
019537 004	5286754	Feb 15, 2011		I-421	Mar 25, 2007
	5286754*PED	Aug 15, 2011		PED	Sep 25, 2007
<u>CIPROFLOXACIN HYDROCHLORIDE; PROQUIN XR</u>					
021744 001	5972389	Sep 19, 2016	DP U-663	NDF	May 19, 2008
	6340475	Sep 19, 2016	DP U-663		
	6488962	Jun 20, 2020	DP		
	6635280	Sep 19, 2016	DP U-663		
<u>CIPROFLOXACIN HYDROCHLORIDE; HYDROCORTISONE; CIPRO HC</u>					
020805 001	4844902	Feb 11, 2008	DP		
	5843930	Jul 06, 2015		U-646	
	5965549	Jun 06, 2015	DP		

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>CIPROFLOXACIN; CIPROFLOXACIN HYDROCHLORIDE; CIPRO XR</u>					
021473 002				I-416 NS PED PED	Aug 28, 2006 Aug 28, 2006 Feb 28, 2007 Feb 28, 2007
<u>CIPROFLOXACIN; DEXAMETHASONE; CIPRODEX</u>					
021537 001	4844902 6284804 6359016	Feb 11, 2008 Aug 10, 2020 Aug 10, 2020		NC	Jul 18, 2006
<u>CISAPRIDE MONOHYDRATE; PROPULSID</u>					
020210 001	4962115	Oct 09, 2007	U-79		
<u>CISAPRIDE MONOHYDRATE; PROPULSID</u>					
020210 002	4962115	Oct 09, 2007	U-79		
<u>CISAPRIDE MONOHYDRATE; PROPULSID</u>					
020398 001	4962115	Oct 09, 2007	U-79		
<u>CISAPRIDE MONOHYDRATE; PROPULSID QUICKSOLV</u>					
020767 001	4962115 5648093	Oct 09, 2007 Jul 15, 2014	U-79		
<u>CISATRACURIUM BESYLATE; NIMBEX</u>					
020551 001	5453510	Sep 26, 2012	U-127		
<u>CISATRACURIUM BESYLATE; NIMBEX PRESERVATIVE FREE</u>					
020551 002	5453510	Sep 26, 2012	U-127		
<u>CISATRACURIUM BESYLATE; NIMBEX PRESERVATIVE FREE</u>					
020551 003	5453510	Sep 26, 2012	U-127		
<u>CLOBETASOL PROPIONATE; CLOBEX</u>					
021535 001	6106848	Sep 22, 2017		NDF	Jul 24, 2006
<u>CLOBETASOL PROPIONATE; CLOBEX</u>					
021644 001				NDF	Feb 05, 2007
<u>CLOBETASOL PROPIONATE; CLOBEX</u>					
021835 001	5972920 5990100	Feb 12, 2018 Mar 24, 2018	DP DP U-742	NDF	Oct 27, 2008
<u>CLOBETASOL PROPIONATE; OLUX</u>					
021142 001	6126920	Oct 03, 2017	U-484		
<u>CLOFARABINE; CLOLAR</u>					
021673 001	5384310 5661136	May 23, 2009 Aug 26, 2014	DS DP U-626	NCE ODE PED PED	Dec 28, 2009 Dec 28, 2011 Jun 28, 2012 Jun 28, 2010
<u>CLOPIDOGREL BISULFATE; CLOPIDOGREL BISULFATE</u>					
076274 001				PC	Feb 04, 2007
<u>CLOPIDOGREL BISULFATE; PLAVIX</u>					
020839 001	4847265 5576328 6429210 6504030	Nov 17, 2011 Jan 31, 2014 Jun 10, 2019 Jun 10, 2019	U-432	I-502	Aug 17, 2009
<u>COLESEVELAM HYDROCHLORIDE; WELCHOL</u>					
021141 001	5607669 5679717 5693675 5917007 5919832 6066678 6433026	Jun 10, 2014 Apr 29, 2014 Dec 02, 2014 Apr 29, 2014 Jun 10, 2014 Jun 10, 2014 Jun 10, 2014	U-323 U-323 U-323 U-323		

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>COLESEVELAM HYDROCHLORIDE; WELCHOL</u>					
021176 001	5607669	Jun 10, 2014		U-323	
	5679717	Apr 29, 2014		U-323	
	5693675	Dec 02, 2014	DS		
	5917007	Apr 29, 2014	DS	U-323	
	5919832	Jun 10, 2014	DS		
	6066678	Jun 10, 2014	DS	U-323	
	6433026	Jun 10, 2014	DS		
	6784254	Jun 10, 2014	DS		
	7101960	Apr 29, 2014	DS DP	U-757	
<u>COLESTIPOL HYDROCHLORIDE; COLESTID</u>					
020222 001	5490987	Feb 13, 2013	DP		
<u>CONIVAPTAN HYDROCHLORIDE; VAPRISOL</u>					
021697 001	5723606	Mar 03, 2015	DS DP	U-698 NCE	Dec 29, 2010
<u>DALTEPARIN SODIUM; FRAGMIN</u>					
020287 001				I-414	Dec 10, 2006
<u>DALTEPARIN SODIUM; FRAGMIN</u>					
020287 003				I-414	Dec 10, 2006
<u>DALTEPARIN SODIUM; FRAGMIN</u>					
020287 004				I-414	Dec 10, 2006
<u>DANAPAROID SODIUM; ORGARAN</u>					
020430 001	5164377	Oct 03, 2010			
<u>DAPSONE; ACZONE</u>					
021794 001	5863560	Sep 11, 2016	DP	NDF	Jul 07, 2008
	6620435	Sep 11, 2016	DP		
<u>DAPTOMYCIN; CUBICIN</u>					
021572 001	6468967	Sep 24, 2019		U-282	NCE Sep 12, 2008
	6852689	Sep 24, 2019		U-282	
	RE39071	Jun 15, 2016	DS DP	U-728	
<u>DAPTOMYCIN; CUBICIN</u>					
021572 002	6468967	Sep 24, 2019		U-282	I-505 May 25, 2009
	6852689	Sep 24, 2019		U-282	NCE Sep 12, 2008
	RE39071	Jun 15, 2016	DS DP	U-728	
<u>DARIFENACIN HYDROBROMIDE; ENABLEX</u>					
021513 001	5096890	Mar 13, 2010	DS	DP U-631	NCE Dec 22, 2009
	6106864	Aug 21, 2016	DP	U-630	
<u>DARIFENACIN HYDROBROMIDE; ENABLEX</u>					
021513 002	5096890	Mar 13, 2010	DS	DP U-631	NCE Dec 22, 2009
	6106864	Aug 21, 2016	DP	U-630	
<u>DARUNAVIR ETHANOLATE; PREZISTA</u>					
021976 001	5843946	Dec 01, 2015		DP U-744	NCE Jun 23, 2011
	6248775	Aug 25, 2012	DS		
	6335460	Aug 25, 2012	DS DP	U-744	
<u>DASATINIB; SPRYCEL</u>					
021986 001	6596746	Apr 13, 2020	DS DP	U-748	NCE Jun 28, 2011
	7125875	Apr 13, 2020		U-779	ODE Jun 28, 2013
	7125875	Apr 13, 2020		U-780	ODE Jun 28, 2013
<u>DASATINIB; SPRYCEL</u>					
021986 002	6596746	Apr 13, 2020	DS DP	U-748	NCE Jun 28, 2011
	7125875	Apr 13, 2020		U-779	ODE Jun 28, 2013
	7125875	Apr 13, 2020		U-780	ODE Jun 28, 2013
<u>DASATINIB; SPRYCEL</u>					
021986 003	6596746	Apr 13, 2020	DS DP	U-748	NCE Jun 28, 2011
	7125875	Apr 13, 2020		U-780	ODE Jun 28, 2013
	7125875	Apr 13, 2020		U-779	ODE Jun 28, 2013
<u>DECITABINE; DACOGEN</u>					
021790 001				NCE	May 02, 2011

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>DEFERASIROX; EXJADE</u>					
021882 001	6465504	Jun 24, 2017	DS DP	NCE	Nov 02, 2010
	6596750	Jun 24, 2017	DS U-735	ODE	Nov 02, 2012
<u>DEFERASIROX; EXJADE</u>					
021882 002	6465504	Jun 24, 2017	DS DP		
	6596750	Jun 24, 2017	DS U-735		
<u>DEFERASIROX; EXJADE</u>					
021882 003	6465504	Jun 24, 2017	DS DP		
	6596750	Jun 24, 2017	DS U-735		
<u>DELAVIRDINE MESYLATE; RESCRIPTOR</u>					
020705 001	5563142	Oct 08, 2013			
<u>DELAVIRDINE MESYLATE; RESCRIPTOR</u>					
020705 002	5563142	Oct 08, 2013			
	6177101	Jun 07, 2019			
<u>DESFLURANE; SUPRANE</u>					
020118 001	4762856	Feb 02, 2007		U-67	
	4762856*PED	Aug 02, 2007			
	5617906	Apr 08, 2014	DP		
	5617906*PED	Oct 08, 2014			
<u>DESIRUDIN RECOMBINANT; IPRIVASK</u>					
021271 001	4745177	May 17, 2010	DS DP	NCE	Apr 04, 2008
	5733874	Mar 31, 2015			
<u>DESLOMATADINE; CLARINEX</u>					
021165 001	4659716	Mar 31, 2007			
	4659716	Mar 31, 2007	DP U-427	NCE	Dec 21, 2006
	4659716*PED	Oct 01, 2007	DP U-725	PED	Jun 21, 2007
	4863931	Sep 15, 2008	U-427		
	4863931*PED	Mar 15, 2009			
	6100274	Jul 07, 2019			
	6100274*PED	Jan 07, 2020			
<u>DESLOMATADINE; CLARINEX</u>					
021300 001	4659716	Mar 31, 2007	DP U-611	NCE	Dec 21, 2006
	4659716	Mar 31, 2007	DP U-725	PED	Jun 21, 2007
	4659716*PED	Oct 01, 2007			
	6514520	Jun 01, 2018	DP		
<u>DESLOMATADINE; CLARINEX</u>					
021312 001	4659716	Mar 31, 2007	DP U-427	NCE	Dec 21, 2006
	4659716	Mar 31, 2007	DP U-725	PED	Jun 21, 2007
	4659716*PED	Oct 01, 2007	U-427		
	4863931	Sep 15, 2008			
	4863931*PED	Mar 15, 2009			
	5178878	Jan 12, 2010	DP		
	5607697	Jun 07, 2015	DP		
	6100274	Jul 07, 2019	DP		
	6100274*PED	Jan 07, 2020			
<u>DESLOMATADINE; CLARINEX</u>					
021312 002	4659716	Mar 31, 2007	DP U-427	NCE	Dec 21, 2006
	4659716	Mar 31, 2007	DP U-725	PED	Jun 21, 2007
	4659716*PED	Oct 01, 2007	DP		
	5178878	Jan 12, 2010	DP		
	5607697	Jun 07, 2015	DP		
	6100274	Jul 07, 2019	DP		
	6100274*PED	Jan 07, 2020			
<u>DESLOMATADINE; PSEUDOEPHEDRINE SULFATE; CLARINEX D 24 HOUR</u>					
021605 001	4659716	Mar 31, 2007	DP U-726	NC	Mar 03, 2008
	4659716	Mar 31, 2007	DP U-644	NCE	Dec 21, 2006
	4659716*PED	Oct 01, 2007		PED	Jun 21, 2007
	6100274	Jul 07, 2019	DP		
	6100274*PED	Jan 07, 2020			
	6979463	Mar 28, 2022	DP		

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE		PATENT CODES			EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE	
DESCLORATADINE; PSEUDOEPHEDRINE SULFATE; CLARINEX-D 12 HOUR									
021313 001	4659716	Mar	31, 2007	DP	U-707	NP		Feb	01, 2009
	4659716*PED	Oct	01, 2007					NC	Mar
	6100274	Jul	07, 2019	DP		NCE		Dec	21, 2006
	6100274*PED	Jan	07, 2020					PED	Jun
	6709676	Feb	18, 2021	DP	U-707				
DESMOPRESSIN ACETATE; DDAVP									
017922 001	5500413	Jun	29, 2013						
	5674850	Dec	23, 2013						
	5763407	Jun	29, 2013						
DESMOPRESSIN ACETATE; DDAVP									
017922 002	5500413	Jun	29, 2013						
	5674850	Dec	23, 2013						
	5763407	Jun	29, 2013						
DESMOPRESSIN ACETATE; DDAVP									
018938 001	5500413	Jun	29, 2013						
	5763407	Jun	29, 2013						
DESMOPRESSIN ACETATE; DDAVP									
018938 002	5500413	Jun	29, 2013						
	5763407	Jun	29, 2013						
DESMOPRESSIN ACETATE; DDAVP									
019955 001	5047398	Sep	10, 2008	DP					
	5500413	Jun	29, 2013						
	5674850	Dec	23, 2013						
	5763407	Jun	29, 2013						
	7022340	Apr	30, 2023						
DESMOPRESSIN ACETATE; DDAVP									
019955 002	5047398	Sep	10, 2008	DP					
	5500413	Jun	29, 2013						
	5674850	Dec	23, 2013						
	5763407	Jun	29, 2013						
	7022340	Apr	30, 2023						
DESMOPRESSIN ACETATE; DDAVP (NEEDS NO REFRIGERATION)									
017922 003	5482931	Jun	29, 2013						
	5500413	Jun	29, 2013						
	5674850	Dec	23, 2013						
	5763407	Jun	29, 2013						
DESOGESTREL; ETHINYL ESTRADIOL; MIRCETTE									
020713 001	RE35724	Oct	20, 2008						
DESONIDE; DESONATE									
021844 001	6387383	Aug	03, 2020	DS	DP	U-783			
DESONIDE; VERDESO									
021978 001	6730288	Sep	08, 2019	DP	DP		NDF	Sep	19, 2009
	7029659	Sep	08, 2019						
DEXMEDETOMIDINE; PRECEDEX									
021038 001	4910214	Jul	15, 2013	DS	DP	U-421			
	6716867	Mar	31, 2019						
DEXMETHYLPHENIDATE HYDROCHLORIDE; FOCALIN									
021278 001	5908850	Dec	04, 2015	DS	DP	U-422			
	6355656	Dec	04, 2015						
	6528530	Dec	04, 2015						
DEXMETHYLPHENIDATE HYDROCHLORIDE; FOCALIN									
021278 002	5908850	Dec	04, 2015	DS	DP	U-422			
	6355656	Dec	04, 2015						
	6528530	Dec	04, 2015						
DEXMETHYLPHENIDATE HYDROCHLORIDE; FOCALIN									
021278 003	5908850	Dec	04, 2015	DS	DP	U-422			
	6355656	Dec	04, 2015						
	6528530	Dec	04, 2015						

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>DEXMETHYLPHENIDATE HYDROCHLORIDE; FOCALIN XR</u>					
021802 001	5837284	Dec 04, 2015	DP	NDF	May 26, 2008
	5908850	Dec 04, 2015	U-678		
	6228398	Nov 01, 2019	DP	U-676	
	6528530	Dec 04, 2015	DP		
	6635284	Dec 04, 2015	DP	U-677	
	6730325	Nov 01, 2019	DP	U-676	
<u>DEXMETHYLPHENIDATE HYDROCHLORIDE; FOCALIN XR</u>					
021802 002	5837284	Dec 04, 2015	DP	NDF	May 26, 2008
	5908850	Dec 04, 2015	U-678		
	6228398	Nov 01, 2019	DP	U-676	
	6528530	Dec 04, 2015	DP		
	6635284	Dec 04, 2015	DP	U-677	
	6730325	Nov 01, 2019	DP	U-676	
<u>DEXMETHYLPHENIDATE HYDROCHLORIDE; FOCALIN XR</u>					
021802 003	5837284	Dec 04, 2015	DP	NDF	May 26, 2008
	5908850	Dec 04, 2015	U-678		
	6228398	Nov 01, 2019	DP	U-676	
	6528530	Dec 04, 2015	DP		
	6635284	Dec 04, 2015	DP	U-677	
	6730325	Nov 01, 2019	DP	U-676	
<u>DEXMETHYLPHENIDATE HYDROCHLORIDE; FOCALIN XR</u>					
021802 004				NDF	May 26, 2008
<u>DEXRAZOXANE HYDROCHLORIDE; ZINECARD</u>					
020212 001	4963551	Dec 21, 2007			
	5242901	Sep 07, 2010	U-339		
<u>DEXRAZOXANE HYDROCHLORIDE; ZINECARD</u>					
020212 002	4963551	Dec 21, 2007			
	5242901	Sep 07, 2010	U-339		
<u>DEXTROMETHORPHAN HYDROBROMIDE; GUAIFENESIN; MUCINEX DM</u>					
021620 001	6372252	Apr 28, 2020	DP		
	6955821	Apr 28, 2020	DP	U-685	
<u>DEXTROMETHORPHAN HYDROBROMIDE; GUAIFENESIN; MUCINEX DM</u>					
021620 002	6372252	Apr 28, 2020	DP		
	6955821	Apr 28, 2020	DP	U-685	
<u>DEXTROMETHORPHAN POLISTIREX; DELSYM</u>					
018658 001	5980882	Apr 16, 2017	DP		
<u>DIAZEPAM; DIASTAT</u>					
020648 001	5462740	Sep 17, 2013	DP		
<u>DIAZEPAM; DIASTAT</u>					
020648 002	5462740	Sep 17, 2013	DP		
<u>DIAZEPAM; DIASTAT</u>					
020648 003	5462740	Sep 17, 2013	DP		
<u>DIAZEPAM; DIASTAT</u>					
020648 004	5462740	Sep 17, 2013	DP		
<u>DIAZEPAM; DIASTAT</u>					
020648 005	5462740	Sep 17, 2013	DP		
<u>DIAZEPAM; DIASTAT ACUDIAL</u>					
020648 006	5462740	Sep 17, 2013	DP		
<u>DIAZEPAM; DIASTAT ACUDIAL</u>					
020648 007	5462740	Sep 17, 2013	DP		
<u>DICLOFENAC SODIUM; DICLOFENAC SODIUM</u>					
020809 001	5603929	Nov 16, 2014		U-239	
	5653972	Nov 16, 2014		U-239	
<u>DICLOFENAC SODIUM; SOLARAZE</u>					
021005 001	5639738	Jun 17, 2014		U-402	
	5792753	Aug 11, 2015			
	5852002	Jun 17, 2014		U-402	
	5914322	Aug 11, 2015			
	5929048	Jul 27, 2016		U-402	
	5985850	Nov 16, 2016			

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>DICLOFENAC SODIUM; VOLTAREN</u>					
020037 001	4829088	Apr 14, 2007			
	4960799	Oct 03, 2007			
<u>DICLOFENAC SODIUM; MISOPROSTOL; ARTHROTEC</u>					
020607 001	5601843	Feb 11, 2014			
	5698225	May 03, 2010		U-392	
<u>DICLOFENAC SODIUM; MISOPROSTOL; ARTHROTEC</u>					
020607 002	5601843	Feb 11, 2014			
	5698225	May 03, 2010		U-392	
<u>DIDANOSINE; VIDEX</u>					
020154 002	4861759	Aug 29, 2006		U-248	
	4861759*PED	Mar 01, 2007		U-248	
	5254539	Aug 29, 2006		U-248	
	5254539*PED	Mar 01, 2007		U-248	
	5616566	Aug 29, 2006		U-180	
	5616566*PED	Mar 01, 2007		U-180	
	5880106	Jul 22, 2011			
	5880106*PED	Jan 22, 2012			
<u>DIDANOSINE; VIDEX</u>					
020154 003	4861759	Aug 29, 2006		U-248	
	4861759*PED	Mar 01, 2007		U-248	
	5254539	Aug 29, 2006		U-248	
	5254539*PED	Mar 01, 2007		U-248	
	5616566	Aug 29, 2006		U-180	
	5616566*PED	Mar 01, 2007		U-180	
	5880106	Jul 22, 2011			
	5880106*PED	Jan 22, 2012			
<u>DIDANOSINE; VIDEX</u>					
020154 004	4861759	Aug 29, 2006		U-248	
	4861759*PED	Mar 01, 2007		U-248	
	5254539	Aug 29, 2006		U-248	
	5254539*PED	Mar 01, 2007		U-248	
	5616566	Aug 29, 2006		U-180	
	5616566*PED	Mar 01, 2007		U-180	
	5880106	Jul 22, 2011			
	5880106*PED	Jan 22, 2012			
<u>DIDANOSINE; VIDEX</u>					
020154 005	4861759	Aug 29, 2006		U-248	
	4861759*PED	Mar 01, 2007		U-248	
	5254539	Aug 29, 2006		U-248	
	5254539*PED	Mar 01, 2007		U-248	
	5616566	Aug 29, 2006		U-180	
	5616566*PED	Mar 01, 2007		U-180	
	5880106	Jul 22, 2011			
	5880106*PED	Jan 22, 2012			
<u>DIDANOSINE; VIDEX</u>					
020154 006	4861759	Aug 29, 2006		U-248	
	4861759*PED	Mar 01, 2007		U-248	
	5254539	Aug 29, 2006		U-248	
	5254539*PED	Mar 01, 2007		U-248	
	5616566	Aug 29, 2006		U-180	
	5616566*PED	Mar 01, 2007		U-180	
	5880106	Jul 22, 2011			
	5880106*PED	Jan 22, 2012			
<u>DIDANOSINE; VIDEX</u>					
020155 003	4861759	Aug 29, 2006		U-248	
	4861759*PED	Mar 01, 2007		U-248	
	5254539	Aug 29, 2006		U-248	
	5254539*PED	Mar 01, 2007		U-248	
	5616566	Aug 29, 2006		U-180	
	5616566*PED	Mar 01, 2007		U-180	

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>DIDANOSINE; VIDEX</u>					
020155 004	4861759	Aug 29, 2006		U-248	
	4861759*PED	Mar 01, 2007		U-248	
	5254539	Aug 29, 2006		U-248	
	5254539*PED	Mar 01, 2007		U-248	
	5616566	Aug 29, 2006		U-180	
	5616566*PED	Mar 01, 2007		U-180	
<u>DIDANOSINE; VIDEX</u>					
020155 005	4861759	Aug 29, 2006		U-248	
	4861759*PED	Mar 01, 2007		U-248	
	5254539	Aug 29, 2006		U-248	
	5254539*PED	Mar 01, 2007		U-248	
	5616566	Aug 29, 2006		U-180	
	5616566*PED	Mar 01, 2007		U-180	
<u>DIDANOSINE; VIDEX</u>					
020155 006	4861759	Aug 29, 2006		U-52	
	4861759*PED	Mar 01, 2007		U-248	
	5254539	Aug 29, 2006		U-248	
	5254539*PED	Mar 01, 2007		U-248	
	5616566	Aug 29, 2006		U-180	
	5616566*PED	Mar 01, 2007		U-180	
<u>DIDANOSINE; VIDEX</u>					
020156 001	4861759	Aug 29, 2006		U-248	
	4861759*PED	Mar 01, 2007		U-248	
	5254539	Aug 29, 2006		U-248	
	5254539*PED	Mar 01, 2007		U-248	
	5616566	Aug 29, 2006		U-180	
	5616566*PED	Mar 01, 2007		U-180	
<u>DIDANOSINE; VIDEX EC</u>					
021183 001	4861759	Aug 29, 2006		U-248	
	4861759*PED	Mar 01, 2007		U-248	
	5254539	Aug 29, 2006		U-248	
	5254539*PED	Mar 01, 2007		U-248	
<u>DIDANOSINE; VIDEX EC</u>					
021183 002	4861759	Aug 29, 2006		U-248	
	4861759*PED	Mar 01, 2007		U-248	
	5254539	Aug 29, 2006		U-248	
	5254539*PED	Mar 01, 2007		U-248	
<u>DIDANOSINE; VIDEX EC</u>					
021183 003	4861759	Aug 29, 2006		U-248	
	4861759*PED	Mar 01, 2007		U-248	
	5254539	Aug 29, 2006		U-248	
	5254539*PED	Mar 01, 2007		U-248	
<u>DIDANOSINE; VIDEX EC</u>					
021183 004	4861759	Aug 29, 2006		U-248	
	4861759*PED	Mar 01, 2007		U-248	
	5254539	Aug 29, 2006		U-248	
	5254539*PED	Mar 01, 2007		U-248	
<u>DIHYDROERGOTAMINE MESYLATE; MIGRANAL</u>					
020148 001	5169849	Dec 08, 2009		U-227	
<u>DILTIAZEM HYDROCHLORIDE; CARDIZEM CD</u>					
020062 001	4894240	Jan 16, 2007			
	5002776	Mar 26, 2008			
	5286497	May 20, 2011			
	5364620	Nov 14, 2011		U-3	
	5439689	Aug 08, 2012		U-107	
	5470584	May 20, 2011			
<u>DILTIAZEM HYDROCHLORIDE; CARDIZEM CD</u>					
020062 002	4894240	Jan 16, 2007			
	5002776	Mar 26, 2008			
	5286497	May 20, 2011			
	5364620	Nov 14, 2011		U-3	
	5439689	Aug 08, 2012		U-107	
	5470584	May 20, 2011			

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>DILTIAZEM HYDROCHLORIDE; CARDIZEM CD</u>					
020062 003	4894240	Jan 16, 2007	U-3 U-107		
	5002776	Mar 26, 2008			
	5286497	May 20, 2011			
	5364620	Nov 14, 2011			
	5439689	Aug 08, 2012			
	5470584	May 20, 2011			
<u>DILTIAZEM HYDROCHLORIDE; CARDIZEM CD</u>					
020062 004	4894240	Jan 16, 2007	U-3 U-107		
	5002776	Mar 26, 2008			
	5286497	May 20, 2011			
	5364620	Nov 14, 2011			
	5439689	Aug 08, 2012			
	5470584	May 20, 2011			
<u>DILTIAZEM HYDROCHLORIDE; CARDIZEM LA</u>					
021392 001	5288505	Jun 26, 2011	DP DP	I-133	Apr 09, 2007
	5529791	Jun 25, 2013			
	6923984	Feb 25, 2021			
	7108866	Dec 17, 2019			
<u>DILTIAZEM HYDROCHLORIDE; CARDIZEM LA</u>					
021392 002	5288505	Jun 26, 2011	DP DP	I-133	Apr 09, 2007
	5529791	Jun 25, 2013			
	6923984	Feb 25, 2021			
	7108866	Dec 17, 2019			
<u>DILTIAZEM HYDROCHLORIDE; CARDIZEM LA</u>					
021392 003	5288505	Jun 26, 2011	DP DP	I-133	Apr 09, 2007
	5529791	Jun 25, 2013			
	6923984	Feb 25, 2021			
	7108866	Dec 17, 2019			
<u>DILTIAZEM HYDROCHLORIDE; CARDIZEM LA</u>					
021392 004	5288505	Jun 26, 2011	DP DP	I-133	Apr 09, 2007
	5529791	Jun 25, 2013			
	6923984	Feb 25, 2021			
	7108866	Dec 17, 2019			
<u>DILTIAZEM HYDROCHLORIDE; CARDIZEM LA</u>					
021392 005	5288505	Jun 26, 2011	DP DP	I-133	Apr 09, 2007
	5529791	Jun 25, 2013			
	6923984	Feb 25, 2021			
	7108866	Dec 17, 2019			
<u>DILTIAZEM HYDROCHLORIDE; CARDIZEM LA</u>					
021392 006	5288505	Jun 26, 2011	DP DP	I-133	Apr 09, 2007
	5529791	Jun 25, 2013			
	6923984	Feb 25, 2021			
	7108866	Dec 17, 2019			
<u>DILTIAZEM HYDROCHLORIDE; DILACOR XR</u>					
020092 001	4839177	Dec 09, 2006			
	5422123	Jun 06, 2012			
<u>DILTIAZEM HYDROCHLORIDE; DILACOR XR</u>					
020092 002	4839177	Dec 09, 2006			
	5422123	Jun 06, 2012			
<u>DILTIAZEM HYDROCHLORIDE; DILACOR XR</u>					
020092 003	4839177	Dec 09, 2006			
	5422123	Jun 06, 2012			
<u>DILTIAZEM HYDROCHLORIDE; DILTIAZEM HYDROCHLORIDE</u>					
020939 001	5288505	Jun 26, 2011			
	5529791	Jun 25, 2013			
<u>DILTIAZEM HYDROCHLORIDE; DILTIAZEM HYDROCHLORIDE</u>					
020939 002	5288505	Jun 26, 2011			
	5529791	Jun 25, 2013			
<u>DILTIAZEM HYDROCHLORIDE; DILTIAZEM HYDROCHLORIDE</u>					
020939 003	5288505	Jun 26, 2011			
	5529791	Jun 25, 2013			

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>DILTIAZEM HYDROCHLORIDE; DILTIAZEM HYDROCHLORIDE</u>					
020939 004	5288505	Jun 26, 2011			
	5529791	Jun 25, 2013			
<u>DILTIAZEM HYDROCHLORIDE; TIAZAC</u>					
020401 001	5529791	Jun 25, 2013			
<u>DILTIAZEM HYDROCHLORIDE; TIAZAC</u>					
020401 002	5529791	Jun 25, 2013			
<u>DILTIAZEM HYDROCHLORIDE; TIAZAC</u>					
020401 003	5529791	Jun 25, 2013			
<u>DILTIAZEM HYDROCHLORIDE; TIAZAC</u>					
020401 004	5529791	Jun 25, 2013			
<u>DILTIAZEM HYDROCHLORIDE; TIAZAC</u>					
020401 005	5529791	Jun 25, 2013			
<u>DILTIAZEM HYDROCHLORIDE; TIAZAC</u>					
020401 006	5529791	Jun 25, 2013			
<u>DILTIAZEM MALATE; TIAMATE</u>					
020506 001	4880631	Sep 24, 2007			
	4968507	Jul 25, 2006			
<u>DILTIAZEM MALATE; TIAMATE</u>					
020506 002	4880631	Sep 24, 2007			
	4968507	Jul 25, 2006			
<u>DILTIAZEM MALATE; TIAMATE</u>					
020506 003	4880631	Sep 24, 2007			
	4968507	Jul 25, 2006			
<u>DILTIAZEM MALATE; ENALAPRIL MALEATE; TECZEM</u>					
020507 001	4880631	Sep 24, 2007			
	4968507	Jul 25, 2006			
	4983598	Jan 08, 2008			
<u>DIMYRISTOYL LECITHIN; PERFLEXANE; IMAGENT</u>					
021191 001	5605673	Feb 25, 2014		NCE	May 31, 2007
	5626833	May 16, 2014	U-458		
	5639443	Jun 17, 2014			
	5695741	Dec 09, 2014	U-458		
	5720938	Feb 24, 2015			
	5798091	Aug 25, 2015	U-458		
	6280704	Jul 30, 2013			
	6280705	Jul 30, 2013			
	6287539	Jul 30, 2013			
<u>DINOPROSTONE; CERVIDIL</u>					
020411 001	4931288	Jan 16, 2007			
	5269321	Dec 14, 2010	U-110		
<u>DIPHENHYDRAMINE CITRATE; IBUPROFEN; ADVIL PM</u>					
021394 001				NC	Dec 21, 2008
<u>DIPHENHYDRAMINE HYDROCHLORIDE; IBUPROFEN POTASSIUM; ADVIL PM</u>					
021393 001				NC	Dec 21, 2008
<u>DIVALPROEX SODIUM; DEPAKOTE</u>					
018723 001	4988731	Jan 29, 2008			
	5212326	Jan 29, 2008			
<u>DIVALPROEX SODIUM; DEPAKOTE</u>					
018723 002	4988731	Jan 29, 2008			
	5212326	Jan 29, 2008			
<u>DIVALPROEX SODIUM; DEPAKOTE</u>					
018723 003	4988731	Jan 29, 2008			
	5212326	Jan 29, 2008			
<u>DIVALPROEX SODIUM; DEPAKOTE</u>					
019680 001	4988731	Jan 29, 2008			
	5212326	Jan 29, 2008			

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>DIVALPROEX SODIUM; DEPAKOTE CP</u>					
019794 001	4988731	Jan 29, 2008			
	5212326	Jan 29, 2008			
<u>DIVALPROEX SODIUM; DEPAKOTE CP</u>					
019794 002	4988731	Jan 29, 2008			
	5212326	Jan 29, 2008			
<u>DIVALPROEX SODIUM; DEPAKOTE ER</u>					
021168 001	4913906	Apr 03, 2007		I-482	Dec 06, 2008
	4988731	Jan 29, 2008			
	5212326	Jan 29, 2008	DS DP		
	6419953	Dec 18, 2018			
	6511678	Dec 18, 2018			
	6528090	Dec 18, 2018	DP		
	6528091	Dec 18, 2018		U-106	
	6713086	Dec 18, 2018	DP	U-579	
	6720004	Dec 18, 2018	DP		
<u>DIVALPROEX SODIUM; DEPAKOTE ER</u>					
021168 002	4913906	Apr 03, 2007		I-482	Dec 06, 2008
	4988731	Jan 29, 2008			
	5212326	Jan 29, 2008	DS DP		
	6511678	Dec 18, 2018			
	6528090	Dec 18, 2018	DP		
	6713086	Dec 18, 2018	DP	U-579	
	6720004	Dec 18, 2018	DP		
<u>DOCETAXEL; TAXOTERE</u>					
020449 001	4814470	May 14, 2010	DS DP	I-519	Oct 17, 2009
	5438072	Nov 22, 2013	DP	I-490	Mar 22, 2009
	5698582	Jul 03, 2012	DP	I-436	Aug 18, 2007
	5714512	Jul 03, 2012	DP		
	5750561	Jul 03, 2012	DP	I-429	May 19, 2007
<u>DOCOSANOL; ABREVA</u>					
020941 001	4874794	Apr 28, 2014			
<u>DOFETILIDE; TIKOSYN</u>					
020931 001	4959366	Sep 25, 2012	DS DP	U-652	
	6124363	Oct 09, 2018			
<u>DOFETILIDE; TIKOSYN</u>					
020931 002	4959366	Sep 25, 2012	DS DP	U-652	
	6124363	Oct 09, 2018			
<u>DOFETILIDE; TIKOSYN</u>					
020931 003	4959366	Sep 25, 2012	DS DP	U-652	
	6124363	Oct 09, 2018			
<u>DOLASETRON MESYLATE; ANZEMET</u>					
020623 001	4906755	Jul 02, 2011			
<u>DOLASETRON MESYLATE; ANZEMET</u>					
020623 002	4906755	Jul 02, 2011			
<u>DOLASETRON MESYLATE; ANZEMET</u>					
020624 001	4906755	Jul 02, 2011			
<u>DOLASETRON MESYLATE; ANZEMET</u>					
020624 002	4906755	Jul 02, 2011			
<u>DOLASETRON MESYLATE; ANZEMET</u>					
020624 003	4906755	Jul 02, 2011			
<u>DONEPEZIL HYDROCHLORIDE; ARICEPT</u>					
020690 001	4895841	Nov 25, 2010			
	5985864	Dec 30, 2016			
	6140321	Dec 30, 2016			
	6245911	Dec 01, 2018			
	6372760	Mar 31, 2019			
<u>DONEPEZIL HYDROCHLORIDE; ARICEPT</u>					
020690 002	4895841	Nov 25, 2010			
	5985864	Dec 30, 2016			
	6140321	Dec 30, 2016			
	6245911	Dec 01, 2018			
	6372760	Mar 31, 2019			

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>DONEPEZIL HYDROCHLORIDE; ARICEPT ODT</u>					
021720 001	4895841	Nov 25, 2010	DS DP U-713		
<u>DONEPEZIL HYDROCHLORIDE; ARICEPT ODT</u>					
021720 002	4895841	Nov 25, 2010	DS DP U-713		
<u>DORZOLAMIDE HYDROCHLORIDE; TRUSOPT</u>					
020408 001	4797413	Apr 28, 2008	DS DP U-103	M-38	Apr 15, 2007
	4797413*PED	Oct 28, 2008		PED	Oct 15, 2007
<u>DORZOLAMIDE HYDROCHLORIDE; TIMOLOL MALEATE; COSOPT</u>					
020869 001	4797413	Apr 28, 2008		U-103	
	4797413*PED	Oct 28, 2008			
	6248735	Apr 17, 2011			
	6316443	Apr 17, 2011	DP	U-561	
<u>DOXAZOSIN MESYLATE; CARDURA XL</u>					
021269 001				NDF	Feb 22, 2008
<u>DOXAZOSIN MESYLATE; CARDURA XL</u>					
021269 002	4837111	Mar 21, 2008	DP	NDF	Feb 22, 2008
<u>DOXERCALCIFEROL; HECTOROL</u>					
020862 001	5602116	Feb 11, 2014		U-278	Apr 23, 2007
	5707980	Aug 17, 2010		U-278	
	5861386	Aug 02, 2008		U-278	
	5869473	Aug 02, 2008		U-278	
	6903083	Jul 18, 2021	DS DP		
<u>DOXERCALCIFEROL; HECTOROL</u>					
020862 002	5869473	Aug 02, 2008		U-278	Apr 23, 2007
	6903083	Jul 18, 2021	DS DP	I-424	
<u>DOXERCALCIFEROL; HECTOROL</u>					
021027 001	5602116	Feb 11, 2014		U-321	
	5707980	Aug 17, 2010		U-321	
	6903083	Jul 18, 2021	DS DP		
<u>DRONABINOL; MARINOL</u>					
018651 001	6703418	Feb 26, 2011		U-563	
<u>DRONABINOL; MARINOL</u>					
018651 002	6703418	Feb 26, 2011		U-563	
<u>DRONABINOL; MARINOL</u>					
018651 003	6703418	Feb 26, 2011		U-563	
<u>DROSPIRENONE; ESTRADIOL; ANGELIQ</u>					
021355 002	6933395	Aug 11, 2017	DS	NP	Sep 28, 2008
<u>DROSPIRENONE; ETHINYL ESTRADIOL; YASMIN</u>					
021098 001	5569652	Oct 29, 2013		U-1	
	6787531	Aug 31, 2020		DP	
	6933395	Aug 11, 2017	DS		
<u>DROSPIRENONE; ETHINYL ESTRADIOL; YAZ</u>					
021676 001	5569652	Oct 29, 2013		U-758	Oct 04, 2009
	5569652	Oct 29, 2013		U-1	Mar 16, 2009
	5798338	Jul 10, 2015			
	6787531	Aug 31, 2020		DP	
	6933395	Aug 11, 2017		DP	
	6958326	Dec 20, 2021		DP	
	6987101	Dec 22, 2017		U-758	
	6987101	Dec 22, 2017		U-1	
	RE37564	Jun 30, 2014		DP	
	RE37838	Jun 30, 2014		DP	
	RE38253	Jun 30, 2014		DP	
<u>DULOXETINE HYDROCHLORIDE; CYMBALTA</u>					
021427 001	5023269	Jun 11, 2008	DS DP U-605	I-470	Sep 03, 2007
	5508276	Jul 18, 2014		NCE	Aug 03, 2009
<u>DULOXETINE HYDROCHLORIDE; CYMBALTA</u>					
021427 002	5023269	Jun 11, 2008	DS DP U-605	I-470	Sep 03, 2007
	5508276	Jul 18, 2014		NCE	Aug 03, 2009

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE		PATENT CODES			EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE	
DULOXETINE HYDROCHLORIDE; CYMBALTA									
021427 004	5023269	Jun 11, 2008	DS	DP	U-605	I-470 NCE	Sep 03, 2007 Aug 03, 2009		
	5508276	Jul 18, 2014							
DUTASTERIDE; AVODART									
021319 001	5565467	Oct 15, 2013	DS		U-476 U-477	NCE	Nov 20, 2006		
	5846976	Sep 17, 2013							
	5998427	Sep 17, 2013							
EFAVIRENZ; SUSTIVA									
020972 001	5519021	May 21, 2013	DS	DP					
	5663169	Sep 02, 2014							
	5811423	Aug 07, 2012	DS	DP	U-257 U-256				
	6238695	Apr 06, 2019							
	6555133	Apr 06, 2019	DS	U-248					
	6639071	Feb 14, 2018							
	6939964	Jan 20, 2018							
EFAVIRENZ; SUSTIVA									
020972 002	5519021	May 21, 2013	DS	DP					
	5663169	Sep 02, 2014							
	5811423	Aug 07, 2012	DS	DP	U-257 U-256				
	6238695	Apr 06, 2019							
	6555133	Apr 06, 2019	DS	U-248					
	6639071	Feb 14, 2018							
	6939964	Jan 20, 2018							
EFAVIRENZ; SUSTIVA									
020972 003	5519021	May 21, 2013	DS	DP					
	5663169	Sep 02, 2014							
	5811423	Aug 07, 2012	DS	DP	U-257 U-256				
	6238695	Apr 06, 2019							
	6555133	Apr 06, 2019	DS	U-248					
	6639071	Feb 14, 2018							
	6939964	Jan 20, 2018							
EFAVIRENZ; SUSTIVA									
021360 001	5519021	May 21, 2013	DS		U-256				
	5663169	Sep 02, 2014							
	5811423	Aug 07, 2012							
	6639071	Feb 14, 2018							
	6939964	Jan 20, 2018							
EFAVIRENZ; SUSTIVA									
021360 002	5519021	May 21, 2013	DS	DP	U-248 U-256				
	5663169	Sep 02, 2014							
	5811423	Aug 07, 2012							
	6639071	Feb 14, 2018							
	6939964	Jan 20, 2018							
EFAVIRENZ; EMTRICITABINE; TENOFOVIR DISOPROXIL FUMARATE; ATRIPLA									
021937 001	5210085	May 11, 2010	DS	DP	U-750 U-750 U-750 U-750 U-750 U-750 U-750 U-750 U-750 U-750 U-750 U-750 U-750 U-750 U-750				
	5210085*PED	Nov 11, 2010							
	5519021	May 21, 2013							
	5663169	Sep 02, 2014							
	5811423	Aug 07, 2012							
	5814639	Sep 29, 2015							
	5814639*PED	Mar 29, 2016							
	5914331	Sep 29, 2015							
	5914331*PED	Mar 29, 2016							
	5922695	Jul 25, 2017							
	5935946	Jul 25, 2017							
	5977089	Jul 25, 2017							
	6043230	Jul 25, 2017							
	6238695	Apr 06, 2019							
	6555133	Apr 06, 2019							
	6639071	Nov 13, 2021							
	6642245	Nov 04, 2020							
	6642245*PED	May 04, 2021							
	6703396	Mar 09, 2021							
	6703396*PED	Sep 09, 2021							
	6939964	Jan 20, 2018							
EFLORNITHINE HYDROCHLORIDE; VANIQA									
021145 001	5648394	Jul 15, 2014			U-334				

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>ELETRIPTAN HYDROBROMIDE; RELPAX</u>					
021016 001	5545644	Aug 13, 2013		NCE	Dec 26, 2007
<u>ELETRIPTAN HYDROBROMIDE; RELPAX</u>					
021016 002	5545644	Aug 13, 2013		NCE	Dec 26, 2007
<u>EMEDASTINE DIFUMARATE; EMADINE</u>					
020706 001	5441958	Dec 08, 2013		U-404	
<u>EMTRICITABINE; EMTRIVA</u>					
021500 001	5210085	May 11, 2010		NCE	Jul 02, 2008
	5210085*PED	Nov 11, 2010		PED	Jan 02, 2009
	5814639	Sep 29, 2015			
	5814639*PED	Mar 29, 2016			
	5914331	Sep 29, 2015			
	5914331*PED	Mar 29, 2016			
	6642245	Nov 04, 2020		U-541	
	6642245*PED	May 04, 2021			
	6703396	Mar 09, 2021	DS DP		
	6703396*PED	Sep 09, 2021			
<u>EMTRICITABINE; EMTRIVA</u>					
021896 001	5210085	May 11, 2010		U-257	NPP Dec 22, 2009
	5210085*PED	Nov 11, 2010		NDF	Sep 27, 2008
	5814639	Sep 29, 2015	DS DP	NCE	Jul 02, 2008
	5814639*PED	Mar 29, 2016		PED	Jun 22, 2010
	5914331	Sep 29, 2015	DS	PED	Mar 27, 2009
	5914331*PED	Mar 29, 2016		PED	Jan 02, 2009
	6642245	Nov 04, 2020		U-257	
	6642245*PED	May 04, 2021			
	6703396	Mar 09, 2021	DS DP		
	6703396*PED	Sep 09, 2021			
<u>EMTRICITABINE; TENOFOVIR DISOPROXIL FUMARATE; TRUVADA</u>					
021752 001	5210085	May 11, 2010		U-541	NCE Oct 26, 2006
	5210085	May 11, 2010		U-248	NCE Jul 02, 2008
	5210085*PED	Nov 11, 2010		PED	Jan 02, 2009
	5814639	Sep 29, 2015	DS DP		
	5814639*PED	Mar 29, 2016			
	5914331	Sep 29, 2015	DS DP	U-248	
	5914331*PED	Mar 29, 2016			
	5922695	Jul 25, 2017	DS	U-541	
	5922695	Jul 25, 2017	DS	U-248	
	5935946	Jul 25, 2017	DS DP	U-541	
	5977089	Jul 25, 2017	DS DP	U-541	
	5977089	Jul 25, 2017	DS DP	U-248	
	6043230	Jul 25, 2017		DP U-248	
	6043230	Jul 25, 2017		DP U-541	
	6642245	Nov 04, 2020		U-541	
	6642245	Nov 04, 2020		U-248	
	6642245*PED	May 04, 2021			
	6703396	Mar 09, 2021	DS DP		
	6703396*PED	Sep 09, 2021			
<u>ENALAPRIL MALEATE; FELODIPINE; LEXXEL</u>					
020668 001	4803081	Apr 03, 2007			
	4803081*PED	Oct 03, 2007			
<u>ENALAPRIL MALEATE; FELODIPINE; LEXXEL</u>					
020668 002	4803081	Apr 03, 2007			
	4803081*PED	Oct 03, 2007			
<u>ENFUVIRTIDE; FUZEON</u>					
021481 001	5464933	Jun 07, 2013		M-59	Sep 29, 2009
	6133418	Jun 07, 2013		NCE	Mar 13, 2008
	6475491	Jun 07, 2015		U-248	
<u>ENOXAPARIN SODIUM; LOVENOX</u>					
020164 009	5389618	Feb 14, 2012	DS DP	U-545	
	RE38743	Feb 14, 2012	DS DP	U-545	
<u>ENOXAPARIN SODIUM; LOVENOX (PRESERVATIVE FREE)</u>					
020164 001	5389618	Feb 14, 2012	DS DP	U-545	
	RE38743	Feb 14, 2012	DS DP	U-545	

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>ENOXAPARIN SODIUM; LOVENOX (PRESERVATIVE FREE)</u>					
020164 002	5389618	Feb 14, 2012	DS DP U-545		
	RE38743	Feb 14, 2012	DS DP U-545		
<u>ENOXAPARIN SODIUM; LOVENOX (PRESERVATIVE FREE)</u>					
020164 003	5389618	Feb 14, 2012	DS DP U-545		
	RE38743	Feb 14, 2012	DS DP U-545		
<u>ENOXAPARIN SODIUM; LOVENOX (PRESERVATIVE FREE)</u>					
020164 004	5389618	Feb 14, 2012	DS DP U-545		
	RE38743	Feb 14, 2012	DS DP U-545		
<u>ENOXAPARIN SODIUM; LOVENOX (PRESERVATIVE FREE)</u>					
020164 005	5389618	Feb 14, 2012	DS DP U-545		
	RE38743	Feb 14, 2012	DS DP U-545		
<u>ENOXAPARIN SODIUM; LOVENOX (PRESERVATIVE FREE)</u>					
020164 006	5389618	Feb 14, 2012	DS DP U-545		
	RE38743	Feb 14, 2012	DS DP U-545		
<u>ENOXAPARIN SODIUM; LOVENOX (PRESERVATIVE FREE)</u>					
020164 007	5389618	Feb 14, 2012	DS DP U-545		
	RE38743	Feb 14, 2012	DS DP U-545		
<u>ENOXAPARIN SODIUM; LOVENOX (PRESERVATIVE FREE)</u>					
020164 008	5389618	Feb 14, 2012	DS DP U-545		
	RE38743	Feb 14, 2012	DS DP U-545		
<u>ENTACAPONE; COMTAN</u>					
020796 001	4963590	Nov 27, 2007		DP U-219	
	5112861	May 12, 2009			U-219
	5135950	Oct 31, 2010	DS DP U-219		
	5446194	Oct 19, 2013	DS		
	6599530	Sep 14, 2018		DP U-219	
<u>ENTECAVIR; BARACLUDE</u>					
021797 001	5206244	Oct 18, 2010	DS	NCE	Mar 29, 2010
<u>ENTECAVIR; BARACLUDE</u>					
021797 002	5206244	Oct 18, 2010	DS	NCE	Mar 29, 2010
<u>ENTECAVIR; BARACLUDE</u>					
021798 001	5206244	Oct 18, 2010	DS	NCE	Mar 29, 2010
<u>EPINASTINE HYDROCHLORIDE; ELESTAT</u>					
021565 001				NCE	Oct 15, 2008
<u>EPINEPHRINE; LIDOCAINE HYDROCHLORIDE; LIDOCAINE HYDROCHLORIDE AND EPINEPHRINE</u>					
021486 001				NP	Oct 26, 2007
<u>EPINEPHRINE; LIDOCAINE HYDROCHLORIDE; LIDOSITE TOPICAL SYSTEM KIT</u>					
021504 001	5246418	Sep 30, 2013	DS DP	NP	May 04, 2007
	5873850	Sep 30, 2013	DS DP		
	6377847	Sep 30, 2013	DS DP		
	6385488	Sep 30, 2013	DS DP		
	6629968	Jun 30, 2020	DS DP		
	6635045	Jun 29, 2021	DS DP		
	6862473	Sep 30, 2013	DP		
<u>EPIRUBICIN HYDROCHLORIDE; ELLENCE</u>					
050778 001				ODE	Sep 15, 2006
<u>EPLERENONE; INSPRA</u>					
021437 001	4559332	Apr 09, 2007	DS DP U-537	I-407	Oct 07, 2006
	6410054	Dec 08, 2019		U-537	
	6410054	Dec 08, 2019		U-3	Sep 27, 2007
	6410524	Nov 05, 2019		U-467	
	6495165	Dec 08, 2019		U-3	
	6495165	Dec 08, 2019		U-537	
	6534093	Dec 08, 2019		U-537	
	6534093	Dec 08, 2019		U-3	
	6558707	Dec 08, 2019	DP	U-537	
	6747020	Nov 05, 2019		U-587	
	6863902	Apr 10, 2020	DP	U-664	

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE		PATENT CODES			EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE	
<u>EPLERENONE; INSPRA</u>									
021437 002	4559332	Apr 09, 2007	DS	DP	U-537	I-407 NCE	Oct 07, 2006 Sep 27, 2007		
	6410054	Dec 08, 2019			U-3				
	6410054	Dec 08, 2019			U-537				
	6410524	Nov 05, 2019			U-467				
	6495165	Dec 08, 2019			U-537				
	6495165	Dec 08, 2019			U-3				
	6534093	Dec 08, 2019			U-3				
	6534093	Dec 08, 2019			U-537				
	6558707	Dec 08, 2019			DP U-537				
	6747020	Nov 05, 2019			U-587				
	6863902	Apr 10, 2020			DP U-664				
<u>EPLERENONE; INSPRA</u>									
021437 003	4559332	Apr 09, 2007	DS	DP	U-537	I-407 NCE	Oct 07, 2006 Sep 27, 2007		
	6410054	Dec 08, 2019			U-3				
	6410054	Dec 08, 2019			U-537				
	6410524	Nov 05, 2019			U-467				
	6495165	Dec 08, 2019			U-537				
	6495165	Dec 08, 2019			U-3				
	6534093	Dec 08, 2019			U-3				
	6534093	Dec 08, 2019			U-537				
	6558707	Dec 08, 2019			DP U-537				
	6747020	Nov 05, 2019			U-587				
	6863902	Apr 10, 2020			DP U-664				
<u>EPOPROSTENOL SODIUM; FLOLAN</u>									
020444 001					ODE	Apr 14, 2007			
<u>EPOPROSTENOL SODIUM; FLOLAN</u>									
020444 002					ODE	Apr 14, 2007			
<u>EPROSARTAN MESYLATE; TEVETEN</u>									
020738 004	5185351	Feb 09, 2010			U-3				
	5656650	Aug 12, 2014			U-3				
<u>EPROSARTAN MESYLATE; TEVETEN</u>									
020738 005	5185351	Feb 09, 2010			U-3				
	5656650	Aug 12, 2014			U-3				
<u>EPROSARTAN MESYLATE; TEVETEN</u>									
020738 006	5185351	Feb 09, 2010			U-3				
	5656650	Aug 12, 2014			U-3				
<u>EPROSARTAN MESYLATE; HYDROCHLOROTHIAZIDE; TEVETEN HCT</u>									
021268 001	5185351	Feb 09, 2010			U-3				
	5656650	Aug 12, 2014			U-3				
<u>EPROSARTAN MESYLATE; HYDROCHLOROTHIAZIDE; TEVETEN HCT</u>									
021268 002	5185351	Feb 09, 2010			U-3				
	5656650	Aug 12, 2014			U-3				
<u>EPTIFIBATIDE; INTEGRILIN</u>									
020718 001	5686570	Nov 11, 2014							
	5747447	May 05, 2015							
	5756451	Nov 11, 2014							
	5807825	Sep 15, 2015			U-244				
	5968902	Jun 02, 2015			U-453				
<u>EPTIFIBATIDE; INTEGRILIN</u>									
020718 002	5686570	Nov 11, 2014							
	5747447	May 05, 2015							
	5756451	Nov 11, 2014							
	5807825	Sep 15, 2015			U-244				
	5968902	Jun 02, 2015			U-453				
<u>ERLOTINIB HYDROCHLORIDE; TARCEVA</u>									
021743 001	5747498	Mar 30, 2015	DS	DP	U-659	I-473 NCE	Nov 02, 2008 Nov 18, 2009		
	6900221	Nov 09, 2020			U-659				
<u>ERLOTINIB HYDROCHLORIDE; TARCEVA</u>									
021743 002	5747498	Mar 30, 2015	DS	DP	U-659	I-473 NCE	Nov 02, 2008 Nov 18, 2009		
	6900221	Nov 09, 2020			U-659				

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>ERLOTINIB HYDROCHLORIDE; TARCEVA</u>					
021743 003	5747498	Mar 30, 2015	DS DP U-659	I-473	Nov 02, 2008
	6900221	Nov 09, 2020	DS DP U-659	NCE	Nov 18, 2009
<u>ERTAPENEM SODIUM; INVANZ</u>					
021337 001	5478820	Feb 02, 2013	DS DP U-160	I-515	Aug 10, 2009
	5478820*PED	Aug 02, 2013		I-476	Oct 14, 2008
	5652233	Feb 02, 2013	DS DP U-160	NPP	May 18, 2008
	5652233*PED	Aug 02, 2013		NCE	Nov 21, 2006
	5952323	May 15, 2017	DP	PED	Nov 18, 2008
	5952323*PED	Nov 15, 2017		PED	May 21, 2007
<u>ESCITALOPRAM OXALATE; LEXAPRO</u>					
021323 001	6916941	Jul 25, 2022	DS DP	I-251	Dec 18, 2006
	6916941*PED	Jan 25, 2023	DS DP		
	RE34712	Sep 14, 2011	DS DP		
	RE34712*PED	Mar 14, 2012			
<u>ESCITALOPRAM OXALATE; LEXAPRO</u>					
021323 002	6916941	Jul 25, 2022	DS DP	I-251	Dec 18, 2006
	6916941*PED	Jan 25, 2023	DS DP		
	RE34712	Sep 14, 2011	DS DP		
	RE34712*PED	Mar 14, 2012			
<u>ESCITALOPRAM OXALATE; LEXAPRO</u>					
021323 003	6916941	Jul 25, 2022	DS DP	I-251	Dec 18, 2006
	6916941*PED	Jan 25, 2023	DS DP		
	RE34712	Sep 14, 2011	DS DP		
	RE34712*PED	Mar 14, 2012			
<u>ESCITALOPRAM OXALATE; LEXAPRO</u>					
021365 001	RE34712	Sep 14, 2011	DS DP	I-251	Dec 18, 2006
	RE34712*PED	Mar 14, 2012			
<u>ESMOLOL HYDROCHLORIDE; BREVIBLOC</u>					
019386 002	5017609	May 21, 2008			
	5017609*PED	Nov 21, 2008			
<u>ESMOLOL HYDROCHLORIDE; BREVIBLOC</u>					
019386 003	5017609	May 21, 2008			
	5017609*PED	Nov 21, 2008			
<u>ESMOLOL HYDROCHLORIDE; BREVIBLOC</u>					
019386 006	6310094	Jan 12, 2021			
	6310094*PED	Jul 12, 2021			
	6528540	Jan 12, 2021			
	6528540*PED	Jul 12, 2021			
<u>ESMOLOL HYDROCHLORIDE; BREVIBLOC</u>					
019386 007	6310094	Jan 12, 2021			
	6310094*PED	Jul 12, 2021			
	6528540	Jan 12, 2021			
	6528540*PED	Jul 12, 2021			
<u>ESMOLOL HYDROCHLORIDE; BREVIBLOC DOUBLE STRENGTH IN PLASTIC CONTAINER</u>					
019386 005	6310094	Jan 12, 2021			
	6310094*PED	Jul 12, 2021			
	6528540	Jan 12, 2021			
	6528540*PED	Jul 12, 2021			
<u>ESMOLOL HYDROCHLORIDE; BREVIBLOC IN PLASTIC CONTAINER</u>					
019386 004	6310094	Jan 12, 2021			
	6310094*PED	Jul 12, 2021			
	6528540	Jan 12, 2021			
	6528540*PED	Jul 12, 2021			

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>ESOMEPRAZOLE MAGNESIUM; NEXIUM</u>					
021153 001	4738974	Apr 19, 2007	DS DP	U-770	I-504 Oct 11, 2009
	4738974	Apr 19, 2007	DS DP	U-635	I-484 Nov 24, 2007
	4738974	Apr 19, 2007	DS DP	U-373	NPP Apr 28, 2009
	4738974*PED	Oct 19, 2007		U-373	
	4786505	Apr 20, 2007	DP	U-770	
	4786505	Apr 20, 2007	DP	U-729	
	4786505	Apr 20, 2007	DP	U-373	
	4786505*PED	Oct 20, 2007		U-373	
	4853230	Apr 20, 2007	DP	U-373	
	4853230	Apr 20, 2007	DP	U-729	
	4853230	Apr 20, 2007	DP	U-770	
	4853230*PED	Oct 20, 2007		U-373	
	5690960	Nov 25, 2014	DP	U-373	
	5690960	Nov 25, 2014	DP	U-770	
	5690960	Nov 25, 2014	DP	U-729	
	5690960*PED	May 25, 2015		U-373	
	5714504	Feb 03, 2015	DP	U-373	
	5714504	Feb 03, 2015	DP	U-729	
	5714504	Feb 03, 2015	DP	U-770	
	5714504*PED	Aug 03, 2015		U-373	
	5877192	May 27, 2014	DP	U-373	
	5877192	May 27, 2014	DP	U-770	
	5877192	May 27, 2014	DP	U-729	
	5877192*PED	Nov 27, 2014		U-373	
	5900424	May 04, 2016	DS	U-729	
	5900424	May 04, 2016	DS	U-373	
	5900424	May 04, 2016	DS	U-770	
	5900424*PED	Nov 04, 2016		U-373	
	6147103	Oct 09, 2018			
	6147103*PED	Apr 09, 2019			
	6166213	Oct 09, 2018			
	6166213*PED	Apr 09, 2019			
	6191148	Oct 09, 2018			
	6191148*PED	Apr 09, 2019			
	6369085	May 25, 2018	DS DP	U-770	
	6369085	May 25, 2018	DS DP	U-729	
	6369085*PED	Nov 25, 2018			
	6428810	Nov 03, 2019	DP	U-729	
	6428810	Nov 03, 2019	DP	U-770	
	6428810	Nov 03, 2019	DP	U-469	
	6428810*PED	May 03, 2020		U-469	
	6875872	May 27, 2014	DS		
	6875872*PED	Nov 27, 2014			

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>ESOMEPRAZOLE MAGNESIUM; NEXIUM</u>					
021153 002	4738974	Apr 19, 2007	DS DP	U-373	I-504 Oct 11, 2009
	4738974	Apr 19, 2007	DS DP	U-635	I-484 Nov 24, 2007
	4738974	Apr 19, 2007	DS DP	U-770	NPP Apr 28, 2009
	4738974*PED	Oct 19, 2007		U-373	
	4786505	Apr 20, 2007	DP	U-770	
	4786505	Apr 20, 2007	DP	U-729	
	4786505	Apr 20, 2007	DP	U-373	
	4786505*PED	Oct 20, 2007		U-373	
	4853230	Apr 20, 2007	DP	U-373	
	4853230	Apr 20, 2007	DP	U-770	
	4853230	Apr 20, 2007	DP	U-729	
	4853230*PED	Oct 20, 2007		U-373	
	5690960	Nov 25, 2014	DP	U-770	
	5690960	Nov 25, 2014	DP	U-729	
	5690960	Nov 25, 2014	DP	U-373	
	5690960*PED	May 25, 2015		U-373	
	5714504	Feb 03, 2015	DP	U-729	
	5714504	Feb 03, 2015	DP	U-770	
	5714504	Feb 03, 2015	DP	U-373	
	5714504*PED	Aug 03, 2015		U-373	
	5877192	May 27, 2014	DP	U-770	
	5877192	May 27, 2014	DP	U-373	
	5877192	May 27, 2014	DP	U-729	
	5877192*PED	Nov 27, 2014		U-373	
	5900424	May 04, 2016	DS	U-770	
	5900424	May 04, 2016	DS	U-729	
	5900424	May 04, 2016	DS	U-373	
	5900424*PED	Nov 04, 2016		U-373	
	6147103	Oct 09, 2018			
	6147103*PED	Apr 09, 2019			
	6166213	Oct 09, 2018			
	6166213*PED	Apr 09, 2019			
	6191148	Oct 09, 2018			
	6191148*PED	Apr 09, 2019			
	6369085	May 25, 2018	DS DP	U-729	
	6369085	May 25, 2018	DS DP	U-770	
	6369085*PED	Nov 25, 2018			
	6428810	Nov 03, 2019	DP	U-469	
	6428810	Nov 03, 2019	DP	U-729	
	6428810	Nov 03, 2019	DP	U-770	
	6428810*PED	May 03, 2020		U-469	
	6875872	May 27, 2014	DS		
	6875872*PED	Nov 27, 2014			

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>ESOMEPRAZOLE MAGNESIUM; NEXIUM</u>					
021957 001	4738974	Apr 19, 2007	DS DP	U-729	I-504 Oct 11, 2009
	4738974	Apr 19, 2007	DS DP	U-773	I-484 Nov 24, 2007
	4783974*PED	Oct 19, 2007			NPP Apr 28, 2009
	4786505	Apr 20, 2007	DP	U-729	
	4786505	Apr 20, 2007	DP	U-773	
	4786505*PED	Oct 20, 2007			
	4853230	Apr 20, 2007	DP	U-773	
	4853230	Apr 20, 2007	DP	U-729	
	4853230*PED	Oct 20, 2007			
	5690960	Nov 25, 2014	DP	U-729	
	5690960	Nov 25, 2014	DP	U-773	
	5690960*PED	May 25, 2015			
	5714504	Feb 03, 2015	DP	U-729	
	5714504	Feb 03, 2015	DP	U-773	
	5714504*PED	Aug 03, 2015			
	5877192	May 27, 2014		U-773	
	5877192	May 27, 2014		U-729	
	5877192*PED	Nov 27, 2014			
	5900424	May 04, 2016	DS	U-773	
	5900424	May 04, 2016	DS	U-729	
	5900424*PED	Nov 04, 2016			
	6369085	May 25, 2018	DS DP	U-773	
	6369085	May 25, 2018	DS DP	U-729	
	6369085*PED	Nov 25, 2018			
	6428810	Nov 03, 2019	DP	U-773	
	6428810	Nov 03, 2019	DP	U-729	
	6428810*PED	May 03, 2020			
	6875872	May 27, 2014	DS		
	6875872*PED	Nov 27, 2014			
<u>ESOMEPRAZOLE MAGNESIUM; NEXIUM</u>					
021957 002	4738974	Apr 19, 2007	DS DP	U-729	I-504 Oct 11, 2009
	4738974	Apr 19, 2007	DS DP	U-773	I-484 Nov 24, 2007
	4783974*PED	Oct 19, 2007			NPP Apr 28, 2009
	4786505	Apr 20, 2007	DP	U-773	
	4786505	Apr 20, 2007	DP	U-729	
	4786505*PED	Oct 20, 2007			
	4853230	Apr 20, 2007	DP	U-773	
	4853230	Apr 20, 2007	DP	U-729	
	4853230*PED	Oct 20, 2007			
	5690960	Nov 25, 2014	DP	U-729	
	5690960	Nov 25, 2014	DP	U-773	
	5690960*PED	May 25, 2015			
	5714504	Feb 03, 2015	DP	U-729	
	5714504	Feb 03, 2015	DP	U-773	
	5714504*PED	Aug 03, 2015			
	5877192	May 27, 2014		U-729	
	5877192	May 27, 2014		U-773	
	5877192*PED	Nov 27, 2014			
	5900424	May 04, 2016	DS	U-773	
	5900424	May 04, 2016	DS	U-729	
	5900424*PED	Nov 04, 2016			
	6369085	May 25, 2018	DS DP	U-729	
	6369085	May 25, 2018	DS DP	U-773	
	6369085*PED	Nov 25, 2018			
	6428810	Nov 03, 2019	DP	U-773	
	6428810	Nov 03, 2019	DP	U-729	
	6428810*PED	May 03, 2020			
	6875872	May 27, 2014	DS		
	6875872*PED	Nov 27, 2014			
<u>ESOMEPRAZOLE SODIUM; NEXIUM IV</u>					
021689 001	5877192	May 27, 2014		U-643	NE Mar 31, 2008
	5877192*PED	Nov 27, 2014			
	6143771	May 27, 2014	DP	U-643	
<u>ESOMEPRAZOLE SODIUM; NEXIUM IV</u>					
021689 002	5877192	May 27, 2014		U-643	NE Mar 31, 2008
	5877192*PED	Nov 27, 2014			
	6143771	May 27, 2014	DP	U-643	

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>ESTRADIOL; ALORA</u>					
020655 001	5122383	May 17, 2011			
	5164190	Dec 11, 2010			
	5212199	May 17, 2011			
	5227169	May 17, 2011			
<u>ESTRADIOL; ALORA</u>					
020655 002	5122383	May 17, 2011			
	5164190	Dec 11, 2010			
	5212199	May 17, 2011			
	5227169	May 17, 2011			
<u>ESTRADIOL; ALORA</u>					
020655 003	5122383	May 17, 2011			
	5164190	Dec 11, 2010			
	5212199	May 17, 2011			
	5227169	May 17, 2011			
<u>ESTRADIOL; ALORA</u>					
020655 004	5122383	May 17, 2011			
	5164190	Dec 11, 2010			
	5212199	May 17, 2011			
	5227169	May 17, 2011			
<u>ESTRADIOL; CLIMARA</u>					
020375 001	5223261	Jun 29, 2010			
<u>ESTRADIOL; CLIMARA</u>					
020375 002	5223261	Jun 29, 2010			
<u>ESTRADIOL; CLIMARA</u>					
020375 003	5223261	Jun 29, 2010			
<u>ESTRADIOL; CLIMARA</u>					
020375 004	5223261	Jun 29, 2010			
<u>ESTRADIOL; CLIMARA</u>					
020375 005	5223261	Jun 29, 2010			
<u>ESTRADIOL; CLIMARA</u>					
020375 006	5223261	Jun 29, 2010			
<u>ESTRADIOL; ELESTRIN</u>					
021813 001				NP	Dec 15, 2009
<u>ESTRADIOL; ESCLIM</u>					
020847 001	4842864	Mar 25, 2008			
<u>ESTRADIOL; ESCLIM</u>					
020847 002	4842864	Mar 25, 2008			
<u>ESTRADIOL; ESCLIM</u>					
020847 003	4842864	Mar 25, 2008			
<u>ESTRADIOL; ESCLIM</u>					
020847 004	4842864	Mar 25, 2008			
<u>ESTRADIOL; ESCLIM</u>					
020847 005	4842864	Mar 25, 2008			
<u>ESTRADIOL; ESTRADIOL</u>					
075182 004				PC	Feb 06, 2007
<u>ESTRADIOL; ESTRADIOL</u>					
075182 005				PC	Feb 06, 2007
<u>ESTRADIOL; ESTRING</u>					
020472 001	4871543	Jun 12, 2007	U-306		
	5188835	Jun 12, 2007	U-306		
<u>ESTRADIOL; ESTROGEL</u>					
021166 001				NDF	Feb 09, 2007
<u>ESTRADIOL; FEMPATCH</u>					
020417 001	4906463	Mar 06, 2007			
	5006342	Apr 09, 2008			

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE		PATENT CODES		EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE	
<u>ESTRADIOL; MENOSTAR</u>								
021674 001	5223261	Jun	29, 2010	DP	U-594	NP	Jun	08, 2007
	5891868	Nov	21, 2017	DP	U-594			
	6692763	Nov	21, 2017	DP	U-594			
<u>ESTRADIOL; VIVELLE</u>								
020323 001	4814168	Mar	04, 2008					
	4994267	Mar	04, 2008					
	4994278	Mar	04, 2008					
	5300291	Apr	05, 2011					
<u>ESTRADIOL; VIVELLE</u>								
020323 002	4814168	Mar	04, 2008					
	4994267	Mar	04, 2008					
	4994278	Mar	04, 2008					
	5300291	Apr	05, 2011					
<u>ESTRADIOL; VIVELLE</u>								
020323 003	4814168	Mar	04, 2008					
	4994267	Mar	04, 2008					
	4994278	Mar	04, 2008					
	5300291	Apr	05, 2011					
<u>ESTRADIOL; VIVELLE</u>								
020323 004	4814168	Mar	04, 2008					
	4994267	Mar	04, 2008					
	4994278	Mar	04, 2008					
	5300291	Apr	05, 2011					
<u>ESTRADIOL; VIVELLE</u>								
020323 005	4814168	Mar	04, 2008					
	4994267	Mar	04, 2008					
	4994278	Mar	04, 2008					
	5300291	Apr	05, 2011					
<u>ESTRADIOL; VIVELLE-DOT</u>								
020538 005	5474783	Dec	12, 2012					
	5656286	Aug	12, 2014					
	5958446	Dec	12, 2012					
	6024976	Jan	07, 2014					
<u>ESTRADIOL; VIVELLE-DOT</u>								
020538 006	5474783	Dec	12, 2012					
	5656286	Aug	12, 2014					
	5958446	Dec	12, 2012					
	6024976	Jan	07, 2014					
<u>ESTRADIOL; VIVELLE-DOT</u>								
020538 007	5474783	Dec	12, 2012					
	5656286	Aug	12, 2014					
	5958446	Dec	12, 2012					
	6024976	Jan	07, 2014					
<u>ESTRADIOL; VIVELLE-DOT</u>								
020538 008	5474783	Dec	12, 2012					
	5656286	Aug	12, 2014					
	5958446	Dec	12, 2012					
	6024976	Jan	07, 2014					
<u>ESTRADIOL ACETATE; FEMRING</u>								
021367 001	5855906	Dec	19, 2015		U-508			
<u>ESTRADIOL ACETATE; FEMRING</u>								
021367 002	5855906	Dec	19, 2015		U-508			
<u>ESTRADIOL ACETATE; FEMTRACE</u>								
021633 001	6962908	Dec	21, 2021	DP		NDF	Aug	20, 2007
<u>ESTRADIOL ACETATE; FEMTRACE</u>								
021633 002	6962908	Dec	21, 2021	DP		NDF	Aug	20, 2007
<u>ESTRADIOL ACETATE; FEMTRACE</u>								
021633 003	6962908	Dec	21, 2021	DP		NDF	Aug	20, 2007
<u>ESTRADIOL HEMIHYDRATE; ESTRASORB</u>								
021371 001	5629021	Jan	31, 2015	DP		NDF	Oct	09, 2006

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
ESTRADIOL; LEVONORGESTREL; CLIMARA PRO					
021258 001	5252334	Oct 12, 2010	DP	I-483	Dec 27, 2008
	5393529	Feb 28, 2012	DP	NC	Nov 21, 2006
	5676968	Oct 14, 2014	DP		
	5770219	Oct 12, 2010	DP		
ESTRADIOL; NORETHINDRONE ACETATE; ACTIVELLA					
020907 002				D-104	Dec 29, 2009
				NS	Dec 28, 2009
ESTRADIOL; NORETHINDRONE ACETATE; COMBIPATCH					
020870 001	5474783	Dec 12, 2012			
	5656286	Aug 12, 2014			
	5958446	Dec 12, 2012			
	6024976	Jan 07, 2014			
ESTRADIOL; NORETHINDRONE ACETATE; COMBIPATCH					
020870 002	5474783	Dec 12, 2012			
	5656286	Aug 12, 2014			
	5958446	Dec 12, 2012			
	6024976	Jan 07, 2014			
ESTRADIOL; NORGESTIMATE; PREFEST					
021040 001	5108995	Apr 28, 2009		U-311	
	5382573	Jan 17, 2012			
	6747019	Mar 20, 2020		U-311	
ESTROGENS, CONJUGATED; PREMARIN					
004782 001	5210081	Feb 26, 2012			
ESTROGENS, CONJUGATED; PREMARIN					
004782 002	5210081	Feb 26, 2012			
ESTROGENS, CONJUGATED; PREMARIN					
004782 003	5210081	Feb 26, 2012		D-82	Jul 16, 2006
ESTROGENS, CONJUGATED; PREMARIN					
004782 004	5210081	Feb 26, 2012			
ESTROGENS, CONJUGATED; PREMARIN					
004782 005	5210081	Feb 26, 2012			
ESTROGENS, CONJUGATED; PREMARIN					
004782 006				D-82	Jul 16, 2006
ESTROGENS, CONJUGATED SYNTHETIC A; CENESTIN					
020992 001	5908638	Jul 26, 2015			
ESTROGENS, CONJUGATED SYNTHETIC A; CENESTIN					
020992 002	5908638	Jul 26, 2015		D-85	Feb 05, 2007
ESTROGENS, CONJUGATED SYNTHETIC A; CENESTIN					
020992 003	5908638	Jul 26, 2015		D-85	Feb 05, 2007
ESTROGENS, CONJUGATED SYNTHETIC A; CENESTIN					
020992 004	5908638	Jul 26, 2015		D-85	Feb 05, 2007
ESTROGENS, CONJUGATED SYNTHETIC B; ENJUVIA					
021443 001	6660726	Mar 08, 2021	DS DP	U-284	NP
	6660726	Mar 08, 2021	DS DP	U-196	
	6855703	Feb 12, 2021	DS DP	U-284	
ESTROGENS, CONJUGATED SYNTHETIC B; ENJUVIA					
021443 002	6660726	Mar 08, 2021	DS DP	U-284	NP
	6660726	Mar 08, 2021	DS DP	U-196	
	6855703	Feb 12, 2021	DS DP	U-284	
ESTROGENS, CONJUGATED SYNTHETIC B; ENJUVIA					
021443 003	6660726	Mar 08, 2021	DS DP	U-196	NP
	6660726	Mar 08, 2021	DS DP	U-284	
	6855703	Feb 12, 2021	DS DP	U-284	
ESTROGENS, CONJUGATED SYNTHETIC B; ENJUVIA					
021443 004	6660726	Mar 08, 2021	DS DP	U-284	NP
	6660726	Mar 08, 2021	DS DP	U-196	
	6855703	Feb 12, 2021	DS DP	U-284	

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>ESTROGENS, CONJUGATED; MEDROXYPROGESTERONE ACETATE; PREMPHASE (PREMARIN;CYCRIN 14/14)</u>					
020303 002	5210081	Feb 26, 2012			
<u>ESTROGENS, CONJUGATED; MEDROXYPROGESTERONE ACETATE; PREMPHASE 14/14</u>					
020527 002	5547948	Jan 17, 2015			
<u>ESTROGENS, CONJUGATED; MEDROXYPROGESTERONE ACETATE; PREMPRO</u>					
020527 001	5547948	Jan 17, 2015			
<u>ESTROGENS, CONJUGATED; MEDROXYPROGESTERONE ACETATE; PREMPRO</u>					
020527 003	5547948	Jan 17, 2015			
<u>ESTROGENS, CONJUGATED; MEDROXYPROGESTERONE ACETATE; PREMPRO</u>					
020527 004	5547948	Jan 17, 2015			
<u>ESTROGENS, CONJUGATED; MEDROXYPROGESTERONE ACETATE; PREMPRO</u>					
020527 005	5547948	Jan 17, 2015			
<u>ESTROGENS, CONJUGATED; MEDROXYPROGESTERONE ACETATE; PREMPRO (PREMARIN;CYCRIN)</u>					
020303 001	5210081	Feb 26, 2012			
<u>ESTROGENS, CONJUGATED; MEPROBAMATE; PMB 200</u>					
010971 005	5210081	Feb 26, 2012			
<u>ESTROGENS, CONJUGATED; MEPROBAMATE; PMB 400</u>					
010971 003	5210081	Feb 26, 2012			
<u>ESZOPICLONE; LUNESTA</u>					
021476 001	6319926 6444673 6864257	Jan 16, 2012 Jan 16, 2012 Aug 30, 2012	DS DP	U-620 U-629	NCE Dec 15, 2009
<u>ESZOPICLONE; LUNESTA</u>					
021476 002	6319926 6444673 6864257	Jan 16, 2012 Jan 16, 2012 Aug 30, 2012	DS DP	U-620 U-629	NCE Dec 15, 2009
<u>ESZOPICLONE; LUNESTA</u>					
021476 003	6319926 6444673 6864257	Jan 16, 2012 Jan 16, 2012 Aug 30, 2012	DS DP	U-620 U-629	NCE Dec 15, 2009
<u>ETHINYL ESTRADIOL; ETONOGESTREL; NUVARING</u>					
021187 001	5989581	Apr 08, 2018			
<u>ETHINYL ESTRADIOL; LEVONORGESTREL; PREVEN EMERGENCY CONTRACEPTIVE KIT</u>					
020946 001	6156742	Dec 05, 2020		U-374	
<u>ETHINYL ESTRADIOL; LEVONORGESTREL; SEASONALE</u>					
021544 001	5898032	Jun 23, 2017		U-1	NP Sep 05, 2006
<u>ETHINYL ESTRADIOL; LEVONORGESTREL; SEASONIQUE</u>					
021840 001				NP	May 25, 2009
<u>ETHINYL ESTRADIOL; NORELGESTROMIN; ORTHO EVRA</u>					
021180 001	5876746 5972377	Nov 20, 2015 Jun 07, 2015	DP	U-514 U-514	
<u>ETHINYL ESTRADIOL; NORETHINDRONE; OVCON-35</u>					
021490 001	6667050	Apr 06, 2019	DP	U-1	NDF Nov 14, 2006
<u>ETHINYL ESTRADIOL; NORETHINDRONE ACETATE; ESTROSTEP 21</u>					
020130 001	4962098 5010070	Oct 09, 2007 Apr 23, 2008		U-112	
<u>ETHINYL ESTRADIOL; NORETHINDRONE ACETATE; ESTROSTEP FE</u>					
020130 002	4962098 5010070	Oct 09, 2007 Apr 23, 2008		U-112	
<u>ETHINYL ESTRADIOL; NORETHINDRONE ACETATE; FEMHRT</u>					
021065 001	5208225	May 04, 2010		U-283	
<u>ETHINYL ESTRADIOL; NORETHINDRONE ACETATE; FEMHRT</u>					
021065 002	5208225	May 04, 2010		U-283	
<u>ETHINYL ESTRADIOL; NORETHINDRONE ACETATE; LOESTRIN 24 FE</u>					
021871 001	5552394	Jul 22, 2014	U-1	NP	Feb 17, 2009

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>ETHINYL ESTRADIOL; NORGESTIMATE; ORTHO CYCLEN-21</u>					
019653 001				M-41 PED	May 13, 2008 Nov 13, 2008
<u>ETHINYL ESTRADIOL; NORGESTIMATE; ORTHO CYCLEN-28</u>					
019653 002				M-41 PED	May 13, 2008 Nov 13, 2008
<u>ETHINYL ESTRADIOL; NORGESTIMATE; ORTHO TRI-CYCLEN</u>					
019697 001				M-41 PED	May 13, 2008 Nov 13, 2008
<u>ETHINYL ESTRADIOL; NORGESTIMATE; ORTHO TRI-CYCLEN</u>					
019697 002				M-41 PED	May 13, 2008 Nov 13, 2008
<u>ETHINYL ESTRADIOL; NORGESTIMATE; ORTHO TRI-CYCLEN LO</u>					
021241 001	6214815 6214815*PED	Jun 09, 2019 Dec 09, 2019	U-112		
<u>ETODOLAC; LODINE XL</u>					
020584 001	4966768 4966768*PED	Oct 30, 2007 Apr 30, 2008			
<u>ETODOLAC; LODINE XL</u>					
020584 002	4966768 4966768*PED	Oct 30, 2007 Apr 30, 2008			
<u>ETODOLAC; LODINE XL</u>					
020584 003	4966768 4966768*PED	Oct 30, 2007 Apr 30, 2008			
<u>ETONOGESTREL; IMPLANON</u>					
021529 001	4957119 5150718	Aug 05, 2008 Sep 29, 2009	DP U-749	NDF	Jul 17, 2009
<u>ETOPOSIDE PHOSPHATE; ETOPOPHOS PRESERVATIVE FREE</u>					
020457 001	5041424 RE35524	Aug 20, 2008 May 17, 2010	U-135		
<u>ETOPOSIDE PHOSPHATE; ETOPOPHOS PRESERVATIVE FREE</u>					
020906 001	5041424 RE35524	Aug 20, 2008 May 17, 2010	U-135		
<u>ETOPOSIDE PHOSPHATE; ETOPOPHOS PRESERVATIVE FREE</u>					
020906 002	5041424 RE35524	Aug 20, 2008 May 17, 2010	U-135		
<u>EXEMESTANE; AROMASIN</u>					
020753 001	4808616 4904650	Apr 01, 2011 Jul 07, 2006	DS DP DS DP	U-658 I-495 ODE	Oct 05, 2008 Oct 21, 2006
<u>EXENATIDE SYNTHETIC; BYETTA</u>					
021773 001	5424286 6858576 6872700 6902744 6956026	May 24, 2013 Jan 06, 2017 Jan 14, 2020 Jan 14, 2020 Jan 07, 2018	U-653 U-656 U-654 DP U-687	I-520 NCE	Dec 22, 2009 Apr 28, 2010
<u>EXENATIDE SYNTHETIC; BYETTA</u>					
021773 002	5424286 6858576 6872700 6902744 6956026	May 24, 2013 Jan 06, 2017 Jan 14, 2020 Jan 14, 2020 Jan 07, 2018	U-653 U-656 U-654 DP U-687	I-520 NCE	Dec 22, 2009 Apr 28, 2010
<u>EZETIMIBE; ZETIA</u>					
021445 001	5846966 7030106 RE37721	Sep 21, 2013 Jan 25, 2022 Oct 25, 2016	U-474 DP DS DP U-473	I-493 M-57 NCE	May 23, 2009 Mar 16, 2009 Oct 25, 2007
<u>EZETIMIBE; SIMVASTATIN; VYTORIN</u>					
021687 001	5846966 RE37721	Sep 21, 2013 Oct 25, 2016	DS DP DS DP U-593 U-473	M-60 M-58 NCE	Oct 03, 2009 Mar 16, 2009 Oct 25, 2007

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE		PATENT CODES			EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE	
<u>EZETIMIBE; SIMVASTATIN; VYTORIN</u>									
021687 002	5846966	Sep 21, 2013		DP	U-593	M-60	Oct 03, 2009		
	RE37721	Oct 25, 2016		DS DP	U-473	M-58	Mar 16, 2009		
						NCE	Oct 25, 2007		
<u>EZETIMIBE; SIMVASTATIN; VYTORIN</u>									
021687 003	5846966	Sep 21, 2013		DP	U-593	M-60	Oct 03, 2009		
	RE37721	Oct 25, 2016		DS DP	U-473	M-58	Mar 16, 2009		
						NCE	Oct 25, 2007		
<u>EZETIMIBE; SIMVASTATIN; VYTORIN</u>									
021687 004	5846966	Sep 21, 2013		DP	U-593	M-60	Oct 03, 2009		
	RE37721	Oct 25, 2016		DS DP	U-473	M-58	Mar 16, 2009		
						NCE	Oct 25, 2007		
<u>FAMCICLOVIR; FAMVIR</u>									
020363 001	5246937	Sep 21, 2010			U-96	D-103	Jul 28, 2009		
	5840763	Sep 01, 2015			U-96	I-501	Jul 28, 2009		
	5866581	Oct 04, 2014			U-96				
	5916893	Sep 01, 2015			U-96				
	6124304	Oct 04, 2014			U-96				
<u>FAMCICLOVIR; FAMVIR</u>									
020363 002	5246937	Sep 21, 2010			U-96	D-103	Jul 28, 2009		
	5840763	Sep 01, 2015			U-96	I-501	Jul 28, 2009		
	5866581	Oct 04, 2014			U-96				
	5916893	Sep 01, 2015			U-96				
	6124304	Oct 04, 2014			U-96				
<u>FAMCICLOVIR; FAMVIR</u>									
020363 003	5246937	Sep 21, 2010			U-96	D-103	Jul 28, 2009		
	5840763	Sep 01, 2015			U-96	I-501	Jul 28, 2009		
	5866581	Oct 04, 2014			U-96				
	5916893	Sep 01, 2015			U-96				
	6124304	Oct 04, 2014			U-96				
<u>FAMOTIDINE; FLUXID</u>									
021712 001	6024981	Apr 09, 2018		DP					
	6221392	Apr 09, 2018		DP					
<u>FAMOTIDINE; FLUXID</u>									
021712 002	6024981	Apr 09, 2018		DP					
	6221392	Apr 09, 2018		DP					
<u>FAMOTIDINE; PEPCID AC</u>									
020325 001	5854267	Dec 29, 2015			U-267				
	5854267*PED	Jun 29, 2016			U-267				
<u>FAMOTIDINE; PEPCID AC</u>									
020325 002						NP	Sep 23, 2006		
<u>FAMOTIDINE; PEPCID AC</u>									
020801 001	5075114	Dec 24, 2008							
	5075114*PED	Jun 24, 2009							
	5667794	May 02, 2015							
	5667794*PED	Nov 02, 2015							
	5854267	Dec 29, 2015			U-267				
	5854267*PED	Jun 29, 2016							
<u>FAMOTIDINE; PEPCID AC (GELTAB)</u>									
020902 001	4820524	Feb 20, 2007							
	4820524*PED	Aug 20, 2007							
	5854267	Dec 29, 2015			U-368				
	5854267*PED	Jun 29, 2016			U-368				
<u>FELBAMATE; FELBATOL</u>									
020189 001	4978680	Sep 26, 2009			U-83				
	5082861	Sep 26, 2009			U-83				
<u>FELBAMATE; FELBATOL</u>									
020189 002	4978680	Sep 26, 2009			U-83				
	5082861	Sep 26, 2009			U-83				

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>FELBAMATE; FELBATOL</u>					
020189 003	4978680	Sep 26, 2009		U-83	
	5082861	Sep 26, 2009		U-83	
<u>FELODIPINE; PLENDIL</u>					
019834 001	4803081	Apr 03, 2007			
	4803081*PED	Oct 03, 2007			
<u>FELODIPINE; PLENDIL</u>					
019834 002	4803081	Apr 03, 2007			
	4803081*PED	Oct 03, 2007			
<u>FELODIPINE; PLENDIL</u>					
019834 004	4803081	Apr 03, 2007			
	4803081*PED	Oct 03, 2007			
<u>FENOFIBRATE; ANTARA (MICRONIZED)</u>					
021695 001	4800079	Aug 10, 2007		M-47	Oct 21, 2008
	7101574	Aug 20, 2020	DS DP		
<u>FENOFIBRATE; ANTARA (MICRONIZED)</u>					
021695 002	4800079	Aug 10, 2007		DP	
<u>FENOFIBRATE; ANTARA (MICRONIZED)</u>					
021695 003	4800079	Aug 10, 2007		M-47	Oct 21, 2008
	7101574	Aug 20, 2020	DS DP		
<u>FENOFIBRATE; LIPOFEN</u>					
021612 001	5545628	Jan 10, 2015		U-701	
<u>FENOFIBRATE; LIPOFEN</u>					
021612 002	5545628	Jan 10, 2015		U-701	
<u>FENOFIBRATE; LIPOFEN</u>					
021612 003	5545628	Jan 10, 2015		U-701	
<u>FENOFIBRATE; TRICOR</u>					
021203 001	4895726	Jan 19, 2009			
	6074670	Jan 09, 2018			
	6277405	Jan 09, 2018			
	6589552	Jan 09, 2018			
	6652881	Jan 09, 2018		DP	
	7037529	Jan 09, 2018		DP	
	7041319	Jan 09, 2018		DP	
<u>FENOFIBRATE; TRICOR</u>					
021203 003	4895726	Jan 19, 2009			
	6074670	Jan 09, 2018			
	6277405	Jan 09, 2018			
	6589552	Jan 09, 2018			
	6652881	Jan 09, 2018		DP	
	7037529	Jan 09, 2018		DP	
	7041319	Jan 09, 2018		DP	
<u>FENOFIBRATE; TRICOR</u>					
021656 001	5145684	Jan 25, 2011		DP U-615	
	6277405	Jan 09, 2018	DS		
	6375986	Sep 21, 2020		DP U-615	
	6652881	Jan 09, 2018	DS		
	7037529	Jan 09, 2018		DP	
	7041319	Jan 09, 2018		DP	
<u>FENOFIBRATE; TRICOR</u>					
021656 002	5145684	Jan 25, 2011		DP U-615	
	6277405	Jan 09, 2018	DS		
	6375986	Sep 21, 2020		DP U-615	
	6652881	Jan 09, 2018	DS		
	7037529	Jan 09, 2018		DP	
	7041319	Jan 09, 2018		DP	
<u>FENOFIBRATE; TRICOR (MICRONIZED)</u>					
019304 002	4895726	Jan 19, 2009			
<u>FENOFIBRATE; TRICOR (MICRONIZED)</u>					
019304 003	4895726	Jan 19, 2009			
<u>FENOFIBRATE; TRICOR (MICRONIZED)</u>					
019304 004	4895726	Jan 19, 2009			

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>FENOFIBRATE; TRIGLIDE</u>					
021350 001	6696084	Sep 11, 2021	DS DP U-680		
<u>FENOFIBRATE; TRIGLIDE</u>					
021350 002	6696084	Sep 11, 2021	DS DP U-680		
<u>FENOLDOPAM MESYLATE; CORLOPAM</u>					
019922 001				I-422	Apr 01, 2007
<u>FENTANYL; DURAGESIC-100</u>					
019813 001				PED	Nov 20, 2006
<u>FENTANYL; DURAGESIC-12</u>					
019813 005				PED	Nov 20, 2006
<u>FENTANYL; DURAGESIC-25</u>					
019813 004				PED	Nov 20, 2006
<u>FENTANYL; DURAGESIC-50</u>					
019813 003				PED	Nov 20, 2006
<u>FENTANYL; DURAGESIC-75</u>					
019813 002				PED	Nov 20, 2006
<u>FENTANYL CITRATE; ACTIQ</u>					
020747 001	4863737	Sep 05, 2006			
<u>FENTANYL CITRATE; ACTIQ</u>					
020747 002	4863737	Sep 05, 2006			
<u>FENTANYL CITRATE; ACTIQ</u>					
020747 003	4863737	Sep 05, 2006			
<u>FENTANYL CITRATE; ACTIQ</u>					
020747 004	4863737	Sep 05, 2006			
<u>FENTANYL CITRATE; ACTIQ</u>					
020747 005	4863737	Sep 05, 2006			
<u>FENTANYL CITRATE; ACTIQ</u>					
020747 006	4863737	Sep 05, 2006			
<u>FENTANYL CITRATE; FENTORA</u>					
021947 001	6200604 6974590	Mar 26, 2019 Mar 26, 2019	U-767 U-767	NDF	Sep 25, 2009
<u>FENTANYL CITRATE; FENTORA</u>					
021947 002	6200604 6974590	Mar 26, 2019 Mar 26, 2019	U-767 U-767	NDF	Sep 25, 2009
<u>FENTANYL CITRATE; FENTORA</u>					
021947 003	6200604 6974590	Mar 26, 2019 Mar 26, 2019	U-767 U-767	NDF	Sep 25, 2009
<u>FENTANYL CITRATE; FENTORA</u>					
021947 004	6200604 6974590	Mar 26, 2019 Mar 26, 2019	U-767 U-767	NDF	Sep 25, 2009
<u>FENTANYL CITRATE; FENTORA</u>					
021947 005	6200604 6974590	Mar 26, 2019 Mar 26, 2019	U-767 U-767	NDF	Sep 25, 2009

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE		PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>FENTANYL HYDROCHLORIDE; IONSYS</u>						
021338 001	5232438	Oct 03, 2008	DP	U-736	NDF	May 22, 2009
	5445606	Dec 11, 2011	DP			
	5697896	Dec 16, 2014	DP			
	5843014	Dec 01, 2015	DP			
	6169920	Jan 02, 2018	DP			
	6171294	Jun 05, 2015		U-736		
	6181963	Nov 02, 2019	DP			
	6195582	Jan 28, 2019	DP	U-736		
	6216033	Jun 05, 2015	DP			
	6317629	Jun 02, 2012	DP			
	6425892	Jun 05, 2015		U-736		
	6842640	Jun 02, 2015	DP			
	6881208	Apr 19, 2022		U-736		
	6975902	Apr 01, 2024	DP			
	7018370	Jun 05, 2015		U-736		
	7027859	Sep 26, 2014	DP			
<u>FERRIC HEXACYANOFERRATE(II); RADIOGARDASE (PRUSSIAN BLUE)</u>						
021626 001					NCE	Oct 02, 2008
					ODE	Oct 02, 2010
<u>FERUMOXIDES; FERIDEX I.V.</u>						
020416 001	4951675	Aug 28, 2007		U-143		
	5055288	Oct 08, 2008				
	5102652	Feb 06, 2009				
	5219554	Jun 15, 2010				
	5248492	Sep 28, 2010				
<u>FERUMOXASIL; GASTROMARK</u>						
020410 001	5055288	Oct 08, 2008				
	5219554	Jun 15, 2010				
<u>FEXOFENADINE HYDROCHLORIDE; ALLEGRA</u>						
020625 001	5578610	Nov 26, 2013		U-192	PED	Nov 12, 2006
	5578610*PED	May 26, 2014		U-192		
	5738872	Feb 28, 2015				
	5738872*PED	Aug 28, 2015				
	5855912	Feb 28, 2015				
	5855912*PED	Aug 28, 2015				
	5932247	Feb 28, 2015				
	5932247*PED	Aug 28, 2015				
	6037353	Mar 14, 2017		U-138		
	6037353*PED	Sep 14, 2017		U-138		
	6113942	Feb 28, 2015				
	6113942*PED	Aug 28, 2015				
	6187791	May 11, 2012		U-138		
	6187791*PED	Nov 11, 2012		U-138		
	6399632	May 11, 2012		U-468		
	6399632*PED	Nov 11, 2012		U-468		
	7135571	May 18, 2014	DS			
	7135571*PED	Nov 18, 2014				
	7138524	May 18, 2014	DS			
	7138524*PED	Nov 18, 2014				

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE		PATENT CODES			EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE	
FEXOFENADINE HYDROCHLORIDE; ALLEGRA									
020872 001	5578610	Nov 26, 2013	DS	DP	U-139	PED	Nov 12, 2006		
	5578610	Nov 26, 2013	DS	DP	U-684				
	5578610*PED	May 26, 2014			U-139				
	5855912	Feb 28, 2015		DP					
	5855912*PED	Aug 28, 2015							
	5932247	Feb 28, 2015		DP					
	5932247*PED	Aug 28, 2015							
	6037353	Mar 14, 2017			U-684				
	6037353	Mar 14, 2017			U-138				
	6037353*PED	Sep 14, 2017			U-138				
	6113942	Feb 28, 2015		DP					
	6113942*PED	Aug 28, 2015							
	6187791	May 11, 2012			U-684				
	6187791	May 11, 2012			U-138				
	6187791*PED	Nov 11, 2012			U-138				
	6399632	May 11, 2012			U-468				
	6399632	May 11, 2012			U-684				
	6399632*PED	Nov 11, 2012			U-468				
	7135571	May 18, 2014	DS						
	7135571*PED	Nov 18, 2014							
	7138524	May 18, 2014	DS						
	7138524*PED	Nov 18, 2014							
FEXOFENADINE HYDROCHLORIDE; ALLEGRA									
020872 002	5578610	Nov 26, 2013	DS	DP	U-684	PED	Nov 12, 2006		
	5578610	Nov 26, 2013	DS	DP	U-139				
	5578610*PED	May 26, 2014			U-139				
	5855912	Feb 28, 2015		DP					
	5855912*PED	Aug 28, 2015							
	5932247	Feb 28, 2015		DP					
	5932247*PED	Aug 28, 2015							
	6037353	Mar 14, 2017			U-684				
	6037353	Mar 14, 2017			U-138				
	6037353*PED	Sep 14, 2017			U-138				
	6113942	Feb 28, 2015		DP					
	6113942*PED	Aug 28, 2015							
	6187791	May 11, 2012			U-684				
	6187791	May 11, 2012			U-138				
	6187791*PED	Nov 11, 2012			U-138				
	6399632	May 11, 2012			U-468				
	6399632	May 11, 2012			U-684				
	6399632*PED	Nov 11, 2012			U-468				
	7135571	May 18, 2014	DS						
	7135571*PED	Nov 18, 2014							
	7138524	May 18, 2014	DS						
	7138524*PED	Nov 18, 2014							
FEXOFENADINE HYDROCHLORIDE; ALLEGRA									
020872 004	5578610	Nov 26, 2013	DS	DP	U-139	D-101 PED	Oct 13, 2008		
	5578610	Nov 26, 2013	DS	DP	U-684		Nov 12, 2006		
	5578610*PED	May 26, 2014			U-139				
	5855912	Feb 28, 2015		DP					
	5855912*PED	Aug 28, 2015							
	5932247	Feb 28, 2015		DP					
	5932247*PED	Aug 28, 2015							
	6037353	Mar 14, 2017			U-684				
	6037353	Mar 14, 2017			U-138				
	6037353*PED	Sep 14, 2017			U-138				
	6113942	Feb 28, 2015		DP					
	6113942*PED	Aug 28, 2015							
	6187791	May 11, 2012			U-684				
	6187791	May 11, 2012			U-138				
	6187791*PED	Nov 11, 2012			U-138				
	6399632	May 11, 2012			U-684				
	6399632	May 11, 2012			U-468				
	6399632*PED	Nov 11, 2012			U-468				
	7135571	May 18, 2014	DS						
	7135571*PED	Nov 18, 2014							
	7138524	May 18, 2014	DS						
	7138524*PED	Nov 18, 2014							

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE		PATENT CODES			EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE	
FEXOFENADINE HYDROCHLORIDE; ALLEGRA									
021963 001	5578610	Nov	26,	2013	DS	DP	U-772		
	6037353	Mar	14,	2017			U-772		
	6187791	May	11,	2012			U-772		
	6399632	May	11,	2012	U-772				
	7138524	May	18,	2014	DS				
	7138524*PED	Nov	18,	2014					
FEXOFENADINE HYDROCHLORIDE; PSEUDOEPHEDRINE HYDROCHLORIDE; ALLEGRA D 24 HOUR									
021704 001	5578610	Nov	26,	2013	DS	DP	U-612		
	5578610*PED	May	26,	2014					
	6004582	May	29,	2018		DP			
	6037353	Mar	17,	2017			U-612		
	6037353*PED	Sep	14,	2017					
	6187791	May	11,	2012			U-612		
	6187791*PED	Nov	11,	2012					
	6399632	May	11,	2012			U-612		
	6399632*PED	Nov	11,	2012					
	6613357	Dec	25,	2020		DP	U-612		
	7138524	May	18,	2014	DS				
	7138524*PED	Nov	18,	2014					
	RE39069	May	29,	2018		DP			
FEXOFENADINE HYDROCHLORIDE; PSEUDOEPHEDRINE HYDROCHLORIDE; ALLEGRA-D 12 HOUR									
020786 001	5578610	Nov	26,	2013	DS				
	5578610*PED	May	26,	2014					
	5855912	Feb	28,	2015					
	5855912*PED	Aug	28,	2015					
	6037353	Mar	14,	2017				U-138	
	6037353*PED	Sep	14,	2017				U-138	
	6039974	Jul	31,	2018					
	6113942	Feb	28,	2015					
	6113942*PED	Aug	28,	2015					
	6187791	May	11,	2012				U-138	
	6187791*PED	Nov	11,	2012				U-138	
	6399632	May	11,	2012				U-468	
	6399632*PED	Nov	11,	2012				U-468	
	7135571	May	18,	2014		DS			
	7135571*PED	Nov	18,	2014					
	7138524	May	18,	2014		DS			
	7138524*PED	Nov	18,	2014					
FINASTERIDE; FINASTERIDE									
076340 001							PC	Dec	16, 2006
FINASTERIDE; PROPECIA									
020788 001	5547957	Oct	15,	2013			U-236		
	5571817	Nov	05,	2013			U-259		
	5886184	Nov	19,	2012					
FINASTERIDE; PROSCAR									
020180 001	5886184	Nov	19,	2012	DS				
	5942519	Oct	23,	2018				U-280	
	6046183	Mar	20,	2011		DP	U-577		
FLUNISOLIDE; AEROBID									
018340 001	4933168	Jun	12,	2007					
FLUNISOLIDE; AEROSPAN HFA									
021247 001							NP	Jan	27, 2009
FLUNISOLIDE; NASALIDE									
018148 001	4933168	Jun	12,	2007					
FLUNISOLIDE; NASAREL									
020409 001	4933168	Jun	12,	2007					
FLUOCINOLONE ACETONIDE; FLUOCINOLONE ACETONIDE									
021930 001							NP	Nov	09, 2008
FLUOCINOLONE ACETONIDE; RETISERT									
021737 001	6217895	Mar	22,	2019		DP	U-708	NDF	Apr 08, 2008
	6548078	Mar	22,	2019		DP	U-708	ODE	Apr 08, 2012

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>FLUOCINONIDE; VANOS</u>					
021758 001	6765001	Dec 21, 2021	DP	I-487 NP	Mar 02, 2009 Feb 11, 2008
<u>FLUOROURACIL; CARAC</u>					
020985 001	6670335	Jun 02, 2021	DP U-68		
<u>FLUOXETINE HYDROCHLORIDE; PROZAC</u>					
018936 001				PED	Jul 03, 2006
<u>FLUOXETINE HYDROCHLORIDE; PROZAC</u>					
018936 003				PED	Jul 03, 2006
<u>FLUOXETINE HYDROCHLORIDE; PROZAC</u>					
018936 004				PED	Jul 03, 2006
<u>FLUOXETINE HYDROCHLORIDE; PROZAC</u>					
018936 006				PED	Jul 03, 2006
<u>FLUOXETINE HYDROCHLORIDE; PROZAC</u>					
020101 001				PED	Jul 03, 2006
<u>FLUOXETINE HYDROCHLORIDE; PROZAC</u>					
020974 001				PED	Jul 03, 2006
<u>FLUOXETINE HYDROCHLORIDE; PROZAC</u>					
020974 002				PED	Jul 03, 2006
<u>FLUOXETINE HYDROCHLORIDE; PROZAC WEEKLY</u>					
021235 001	5910319	May 29, 2017		U-396	
	5985322	May 29, 2017		U-397	
	RE39030	May 29, 2017	DP	U-397	
	RE39030	May 29, 2017	DP	U-396	
<u>FLUOXETINE HYDROCHLORIDE; SARAFEM</u>					
018936 007	4971998	Nov 20, 2007		U-338	
	4971998*PED	May 20, 2008		U-338	
<u>FLUOXETINE HYDROCHLORIDE; SARAFEM</u>					
018936 008	4971998	Nov 20, 2007		U-338	
	4971998*PED	May 20, 2008		U-338	
<u>FLUOXETINE HYDROCHLORIDE; OLANZAPINE; SYMBYAX</u>					
021520 002	5229382	Apr 23, 2011	DS DP	NC	Dec 24, 2006
	5945416	Mar 24, 2017	DS DP		
<u>FLUOXETINE HYDROCHLORIDE; OLANZAPINE; SYMBYAX</u>					
021520 003	5229382	Apr 23, 2011	DS DP	NC	Dec 24, 2006
	5945416	Mar 24, 2017	DS DP		
<u>FLUOXETINE HYDROCHLORIDE; OLANZAPINE; SYMBYAX</u>					
021520 004	5229382	Apr 23, 2011	DS DP	NC	Dec 24, 2006
	5945416	Mar 24, 2017	DS DP		
<u>FLUOXETINE HYDROCHLORIDE; OLANZAPINE; SYMBYAX</u>					
021520 005	5229382	Apr 23, 2011	DS DP	NC	Dec 24, 2006
	5945416	Mar 24, 2017	DS DP		
<u>FLUTICASONE PROPIONATE; CUTIVATE</u>					
021152 001				NDF PED	Mar 31, 2008 Sep 30, 2008
<u>FLUTICASONE PROPIONATE; FLONASE</u>					
020121 001				PED	Nov 01, 2006
<u>FLUTICASONE PROPIONATE; FLOVENT DISKUS 100</u>					
020833 002	6536427	Mar 01, 2011	DP		
	6536427*PED	Sep 01, 2011	DP		
<u>FLUTICASONE PROPIONATE; FLOVENT DISKUS 250</u>					
020833 003	6536427	Mar 01, 2011	DP		
	6536427*PED	Sep 01, 2011	DP		
<u>FLUTICASONE PROPIONATE; FLOVENT DISKUS 50</u>					
020833 001	6536427	Mar 01, 2011	DP		
	6536427*PED	Sep 01, 2011	DP		

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>FLUTICASONE PROPIONATE; FLOVENT HFA</u>					
021433 001	5658549	Sep 19, 2014	DP U-710	NPP	Feb 28, 2009
	5674472	Oct 07, 2014	DP	NP	May 14, 2007
	6170717	Dec 23, 2017	DP		
	6251368	Dec 04, 2012	DP		
	6253762	Apr 14, 2015	DP U-582		
	6315173	Dec 23, 2017	DP		
	6510969	Dec 23, 2017	DP		
	6546928	Apr 14, 2015	DP U-583		
	6743413	Jun 01, 2021	DP U-581		
<u>FLUTICASONE PROPIONATE; FLOVENT HFA</u>					
021433 002	5658549	Sep 19, 2014	DP U-710	NPP	Feb 28, 2009
	5674472	Oct 07, 2014	DP	NP	May 14, 2007
	6170717	Dec 23, 2017	DP		
	6251368	Dec 04, 2012	DP		
	6253762	Apr 14, 2015	DP U-582		
	6315173	Dec 23, 2017	DP		
	6510969	Dec 23, 2017	DP		
	6546928	Apr 14, 2015	DP U-583		
	6743413	Jun 01, 2021	DP U-581		
<u>FLUTICASONE PROPIONATE; FLOVENT HFA</u>					
021433 003	5658549	Sep 19, 2014	DP U-710	NPP	Feb 28, 2009
	5674472	Oct 07, 2014	DP	NP	May 14, 2007
	6170717	Dec 23, 2017	DP		
	6251368	Dec 04, 2012	DP		
	6253762	Apr 14, 2015	DP U-582		
	6315173	Dec 23, 2017	DP		
	6510969	Dec 23, 2017	DP		
	6546928	Apr 14, 2015	DP U-583		
	6743413	Jun 01, 2021	DP U-581		
<u>FLUTICASONE PROPIONATE; SALMETEROL XINAFOATE; ADVAIR DISKUS 100/50</u>					
021077 001	4992474	Feb 12, 2008	U-211	M-33	Apr 21, 2007
	4992474*PED	Aug 12, 2008	U-211		
	5126375	Feb 12, 2008			
	5126375*PED	Aug 12, 2008			
	5225445	Feb 12, 2008	U-211		
	5225445*PED	Aug 12, 2008	U-211		
	5270305	Sep 07, 2010	U-387		
	6536427	Mar 01, 2011	DP		
	6536427*PED	Sep 01, 2011			
<u>FLUTICASONE PROPIONATE; SALMETEROL XINAFOATE; ADVAIR DISKUS 250/50</u>					
021077 002	4992474	Feb 12, 2008	U-211	I-410	Nov 17, 2006
	4992474*PED	Aug 12, 2008	U-211		
	5126375	Feb 12, 2008			
	5126375*PED	Aug 12, 2008			
	5225445	Feb 12, 2008	U-211		
	5225445*PED	Aug 12, 2008	U-211		
	5270305	Sep 07, 2010	U-387		
	6536427	Mar 01, 2011	DP		
	6536427*PED	Sep 01, 2011			
<u>FLUTICASONE PROPIONATE; SALMETEROL XINAFOATE; ADVAIR DISKUS 500/50</u>					
021077 003	4992474	Feb 12, 2008	U-211		
	4992474*PED	Aug 12, 2008	U-211		
	5126375	Feb 12, 2008			
	5126375*PED	Aug 12, 2008			
	5225445	Feb 12, 2008	U-211		
	5225445*PED	Aug 12, 2008	U-211		
	5270305	Sep 07, 2010	U-387		
	6536427	Mar 01, 2011	DP		
	6536427*PED	Sep 01, 2011			

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
FLUTICASONE PROPIONATE; SALMETEROL XINAFOATE; ADVAIR HFA					
021254 001	4992474	Feb 12, 2008	DS DP	NP	Jun 08, 2009
	4992474*PED	Aug 12, 2008			
	5126375	Feb 12, 2008	DS DP		
	5126375*PED	Aug 12, 2008			
	5225445	Feb 12, 2008			
	5225445*PED	Aug 12, 2008			
	5270305	Sep 07, 2010			
	5658549	Aug 19, 2014	DP		
	5674472	Oct 07, 2014	DP		
	6143277	Apr 14, 2015	DP		
	6251368	Dec 04, 2012	DP		
	6253762	Apr 14, 2015	DP		
	6315173	Dec 23, 2017	DP		
	6510969	Dec 23, 2017	DP		
	6524555	Apr 14, 2015	DP		
	6546928	Apr 14, 2015	DP		
FLUTICASONE PROPIONATE; SALMETEROL XINAFOATE; ADVAIR HFA					
021254 002	4992474	Feb 12, 2008	DS DP	NP	Jun 08, 2009
	4992474*PED	Aug 12, 2008			
	5126375	Feb 12, 2008	DS DP		
	5126375*PED	Aug 12, 2008			
	5225445	Feb 12, 2008			
	5225445*PED	Aug 12, 2008			
	5270305	Sep 07, 2010			
	5658549	Aug 19, 2014	DP		
	5674472	Oct 07, 2014	DP		
	6143277	Apr 14, 2015	DP		
	6251368	Dec 04, 2012	DP		
	6253762	Apr 14, 2015	DP		
	6315173	Dec 23, 2017	DP		
	6510969	Dec 23, 2017	DP		
	6524555	Apr 14, 2015	DP		
	6546928	Apr 14, 2015	DP		
FLUTICASONE PROPIONATE; SALMETEROL XINAFOATE; ADVAIR HFA					
021254 003	4992474	Feb 12, 2008	DS DP	NP	Jun 08, 2009
	4992474*PED	Aug 12, 2008			
	5126375	Feb 12, 2008	DS DP		
	5126375*PED	Aug 12, 2008			
	5225445	Feb 12, 2008			
	5225445*PED	Aug 12, 2008			
	5270305	Sep 07, 2010			
	5658549	Aug 19, 2014	DP		
	5674472	Oct 07, 2014	DP		
	6143277	Apr 14, 2015	DP		
	6251368	Dec 04, 2012	DP		
	6253762	Apr 14, 2015	DP		
	6315173	Dec 23, 2017	DP		
	6510969	Dec 23, 2017	DP		
	6524555	Apr 14, 2015	DP		
	6546928	Apr 14, 2015	DP		
FLUVASTATIN SODIUM; LESCOL					
020261 001	5354772	Oct 11, 2011		I-507 U-109 PED PED	Apr 10, 2009 Oct 10, 2009 Nov 27, 2006
	5354772	Oct 11, 2011			
	5354772*PED	Apr 11, 2012			
	5356896	Dec 12, 2011			
	5356896*PED	Jun 12, 2012			
FLUVASTATIN SODIUM; LESCOL					
020261 002	5354772	Oct 11, 2011		I-507 U-413 PED PED	Apr 10, 2009 Oct 10, 2009 Nov 27, 2006
	5354772	Oct 11, 2011			
	5354772*PED	Apr 11, 2012			
	5356896	Dec 12, 2011			
	5356896*PED	Jun 12, 2012			

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>FLUVASTATIN SODIUM; LESCOL XL</u>					
021192 001	5354772	Oct 11, 2011	U-109	I-507	Apr 10, 2009
	5354772	Oct 11, 2011	U-413	PED	Oct 10, 2009
	5354772*PED	Apr 11, 2012		PED	Nov 27, 2006
	5356896	Dec 12, 2011			
	5356896*PED	Jun 12, 2012			
	6242003	Apr 13, 2020			
	6242003*PED	Oct 13, 2020			
<u>FOLLITROPIN ALFA/BETA; FOLLISTIM</u>					
020582 001	5270057	Mar 20, 2011			
	5767251	Jun 16, 2015			
<u>FOLLITROPIN ALFA/BETA; FOLLISTIM</u>					
020582 002	5270057	Mar 20, 2011			
	5767251	Jun 16, 2015			
<u>FOLLITROPIN ALFA/BETA; FOLLISTIM AQ</u>					
021211 001	5767251	Jun 16, 2015	DS		
	5929028	Jan 14, 2018	DP U-567		
<u>FOLLITROPIN ALFA/BETA; FOLLISTIM AQ</u>					
021211 002	5767251	Jun 16, 2015	DS		
	5929028	Jan 14, 2018	DP U-567		
<u>FOLLITROPIN ALFA/BETA; GONAL-F</u>					
020378 001	5767251	Jun 16, 2015	DS	ODE	May 24, 2007
<u>FOLLITROPIN ALFA/BETA; GONAL-F</u>					
020378 002	5767251	Jun 16, 2015	DS	ODE	May 24, 2007
<u>FOLLITROPIN ALFA/BETA; GONAL-F</u>					
020378 003	5767251	Jun 16, 2015	DS	ODE	May 24, 2007
<u>FOLLITROPIN ALFA/BETA; GONAL-F</u>					
020378 004	5767067	Jun 16, 2015	DS	ODE	May 24, 2007
	5767251	Jun 16, 2015	DS		
<u>FOLLITROPIN ALFA/BETA; GONAL-F</u>					
020378 005	5767067	Jun 16, 2015	DS	ODE	May 24, 2007
	5767251	Jun 16, 2015	DS		
<u>FOLLITROPIN ALFA/BETA; GONAL-F</u>					
021765 001	5767251	Jun 16, 2015	DS	NP	Mar 25, 2007
<u>FOLLITROPIN ALFA/BETA; GONAL-F</u>					
021765 003	5767251	Jun 16, 2015	DS	NP	Mar 25, 2007
<u>FOLLITROPIN ALFA/BETA; GONAL-F RFF</u>					
021765 002	5767067	Jun 16, 2015	DS	NP	Mar 25, 2007
	5767251	Jun 16, 2015	DS		
<u>FOLLITROPIN ALFA/BETA; GONAL-F RFF PEN</u>					
021684 001	5767067	Jun 16, 2015	DS		
	5767251	Jun 16, 2015	DS		
<u>FOLLITROPIN ALFA/BETA; GONAL-F RFF PEN</u>					
021684 002	5767067	Jun 16, 2015	DS		
	5767251	Jun 16, 2015	DS		
<u>FOLLITROPIN ALFA/BETA; GONAL-F RFF PEN</u>					
021684 003	5767067	Jun 16, 2015	DS		
	5767251	Jun 16, 2015	DS		
<u>FOMEPIZOLE; ANTIZOL</u>					
020696 001				ODE	Dec 08, 2007
<u>FOMIVIRSEN SODIUM; VITRAVENE PRESERVATIVE FREE</u>					
020961 001	5264423	Nov 23, 2010	U-522		
	5276019	Jan 04, 2011	U-522		
	5442049	Aug 15, 2012			
	5595978	Aug 15, 2012	U-522		

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>FONDAPARINUX SODIUM; ARIXTRA</u>					
021345 001				I-457	May 26, 2008
				I-427	May 31, 2007
				I-426	May 31, 2007
				NCE	Dec 07, 2006
<u>FONDAPARINUX SODIUM; ARIXTRA</u>					
021345 002				I-457	May 26, 2008
				I-427	May 31, 2007
				I-426	May 31, 2007
				NS	May 31, 2007
				NCE	Dec 07, 2006
<u>FONDAPARINUX SODIUM; ARIXTRA</u>					
021345 003				I-457	May 26, 2008
				I-427	May 31, 2007
				I-426	May 31, 2007
				NS	May 31, 2007
				NCE	Dec 07, 2006
<u>FONDAPARINUX SODIUM; ARIXTRA</u>					
021345 004				I-457	May 26, 2008
				I-427	May 31, 2007
				I-426	May 31, 2007
				NS	May 31, 2007
				NCE	Dec 07, 2006
<u>FORMOTEROL FUMARATE; FORADIL</u>					
020831 001	6488027	Mar 08, 2019			
	6887459	Nov 28, 2020		U-762	
<u>FOSAMPRENAVIR CALCIUM; LEXIVA</u>					
021548 001	6436989	Dec 24, 2017		U-257	
	6514953	Jul 15, 2019		U-257	
<u>FOSINOPRIL SODIUM; MONOPRIL</u>					
019915 002	5006344	Jul 10, 2009		PED	
	5006344*PED	Jan 10, 2010			Nov 27, 2006
<u>FOSINOPRIL SODIUM; MONOPRIL</u>					
019915 003	5006344	Jul 10, 2009		PED	
	5006344*PED	Jan 10, 2010			Nov 27, 2006
<u>FOSINOPRIL SODIUM; MONOPRIL</u>					
019915 004	5006344	Jul 10, 2009		PED	
	5006344*PED	Jan 10, 2010			Nov 27, 2006
<u>FOSINOPRIL SODIUM; HYDROCHLOROTHIAZIDE; MONOPRIL-HCT</u>					
020286 001	5006344	Jul 10, 2009			
	5006344*PED	Jan 10, 2010			
<u>FOSINOPRIL SODIUM; HYDROCHLOROTHIAZIDE; MONOPRIL-HCT</u>					
020286 002	5006344	Jul 10, 2009			
	5006344*PED	Jan 10, 2010			
<u>FOSPHENYTOIN SODIUM; CEREBYX</u>					
020450 001	4925860	Aug 05, 2007			
<u>FROVATRIPTAN SUCCINATE; FROVA</u>					
021006 001	5464864	Nov 07, 2015		U-436	
	5616603	Apr 01, 2014		U-436	
	5637611	Jun 10, 2014		U-436	
	5827871	Oct 27, 2015		U-436	
	5962501	Dec 16, 2013		U-436	
<u>FULVESTRANT; FASLODEX</u>					
021344 001	4659516	Dec 11, 2007	DS DP	U-596	
	6774122	Jan 09, 2021		U-596	
				NCE	Apr 25, 2007

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>GABAPENTIN; NEURONTIN</u>					
020235 001	4894476	May 02, 2008			
	4894476*PED	Nov 02, 2008			
	6054482	Apr 25, 2017			
	6054482*PED	Oct 25, 2017			
<u>GABAPENTIN; NEURONTIN</u>					
020235 002	4894476	May 02, 2008			
	4894476*PED	Nov 02, 2008			
	6054482	Apr 25, 2017			
	6054482*PED	Oct 25, 2017			
<u>GABAPENTIN; NEURONTIN</u>					
020235 003	4894476	May 02, 2008			
	4894476*PED	Nov 02, 2008			
	6054482	Apr 25, 2017			
	6054482*PED	Oct 25, 2017			
<u>GABAPENTIN; NEURONTIN</u>					
020882 001	4894476	May 02, 2008			
	4894476*PED	Nov 02, 2008			
	6054482	Apr 25, 2017			
	6054482*PED	Oct 25, 2017			
<u>GABAPENTIN; NEURONTIN</u>					
020882 002	4894476	May 02, 2008			
	4894476*PED	Nov 02, 2008			
	6054482	Apr 25, 2017			
	6054482*PED	Oct 25, 2017			
<u>GABAPENTIN; NEURONTIN</u>					
021129 001	4894476	May 02, 2008			
	4894476*PED	Nov 02, 2008			
	6054482	Apr 25, 2017			
	6054482*PED	Oct 25, 2017			
<u>GADOBENATE DIMEGLUMINE; MULTIANCE</u>					
021357 001				NCE	Nov 23, 2009
<u>GADOBENATE DIMEGLUMINE; MULTIANCE</u>					
021357 002				NCE	Nov 23, 2009
<u>GADOBENATE DIMEGLUMINE; MULTIANCE</u>					
021357 003				NCE	Nov 23, 2009
<u>GADOBENATE DIMEGLUMINE; MULTIANCE</u>					
021357 004				NCE	Nov 23, 2009
<u>GADOBENATE DIMEGLUMINE; MULTIANCE MULTIPACK</u>					
021358 001				NCE	Nov 23, 2009
<u>GADOBENATE DIMEGLUMINE; MULTIANCE MULTIPACK</u>					
021358 002				NCE	Nov 23, 2009
<u>GADODIAMIDE; OMNISCAN</u>					
020123 001	4687659	May 04, 2007		U-76	
	5362475	Nov 08, 2011	DS		
<u>GADOPENTETATE DIMEGLUMINE; MAGNEVIST</u>					
019596 001	5362475	Nov 08, 2011			
	5560903	Oct 01, 2013			
<u>GADOPENTETATE DIMEGLUMINE; MAGNEVIST</u>					
021037 001	5362475	Nov 08, 2011			
	5560903	Oct 01, 2013			
<u>GADOTERIDOL; PROHANCE</u>					
020131 001	4885363	Dec 05, 2006			
	5474756	Dec 12, 2012		U-480	
	5846519	Dec 08, 2015			
	6143274	Dec 12, 2012		U-480	
<u>GADOTERIDOL; PROHANCE MULTIPACK</u>					
021489 001	4885363	Dec 05, 2006			
	5474756	Dec 12, 2012		U-536	
	5846519	Dec 08, 2015			
	6143274	Dec 12, 2012		U-536	

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>GADOVERSETAMIDE; OPTIMARK</u>					
020937 001	5130120	Jul 14, 2009			
	5137711	Jul 14, 2009			
<u>GADOVERSETAMIDE; OPTIMARK</u>					
020937 002	5130120	Jul 14, 2009			
	5137711	Jul 14, 2009			
<u>GADOVERSETAMIDE; OPTIMARK</u>					
020937 003	5130120	Jul 14, 2009			
	5137711	Jul 14, 2009			
<u>GADOVERSETAMIDE; OPTIMARK</u>					
020937 004	5130120	Jul 14, 2009			
	5137711	Jul 14, 2009			
<u>GADOVERSETAMIDE; OPTIMARK</u>					
020975 001	5130120	Jul 14, 2009			
	5137711	Jul 14, 2009			
<u>GADOVERSETAMIDE; OPTIMARK IN PLASTIC CONTAINER</u>					
020976 001	5130120	Jul 14, 2009			
	5137711	Jul 14, 2009			
<u>GALANTAMINE HYDROBROMIDE; RAZADYNE</u>					
021169 001	4663318	Dec 14, 2008		U-322	
	6099863	Jun 06, 2017			
	6358527	Jun 06, 2017	DP	U-322	
<u>GALANTAMINE HYDROBROMIDE; RAZADYNE</u>					
021169 002	4663318	Dec 14, 2008		U-322	
	6099863	Jun 06, 2017			
	6358527	Jun 06, 2017	DP	U-322	
<u>GALANTAMINE HYDROBROMIDE; RAZADYNE</u>					
021169 003	4663318	Dec 14, 2008		U-322	
	6099863	Jun 06, 2017			
	6358527	Jun 06, 2017	DP	U-322	
<u>GALANTAMINE HYDROBROMIDE; RAZADYNE</u>					
021224 001	4663318	Dec 14, 2008		U-322	
<u>GALANTAMINE HYDROBROMIDE; RAZADYNE ER</u>					
021615 001	4663318	Dec 14, 2008		U-322	NDF Apr 01, 2008
<u>GALANTAMINE HYDROBROMIDE; RAZADYNE ER</u>					
021615 002	4663318	Dec 14, 2008		U-322	NDF Apr 01, 2008
<u>GALANTAMINE HYDROBROMIDE; RAZADYNE ER</u>					
021615 003	4663318	Dec 14, 2008		U-322	NDF Apr 01, 2008
<u>GANCICLOVIR; VITRASERT</u>					
020569 001	5378475	Jan 03, 2012			
<u>GANIRELIX ACETATE; GANIRELIX ACETATE INJECTION</u>					
021057 001	4801577	Feb 05, 2012	DS	DP	
	5767082	Jun 16, 2015			
<u>GATIFLOXACIN; TEQUIN</u>					
021061 001	4980470	Dec 15, 2009	DS	I-432	Jun 30, 2007
	5880283	Dec 05, 2015			
<u>GATIFLOXACIN; TEQUIN</u>					
021061 002	4980470	Dec 15, 2009	DS	I-432	Jun 30, 2007
	5880283	Dec 05, 2015			
<u>GATIFLOXACIN; TEQUIN</u>					
021062 003	4980470	Dec 15, 2009	DS	I-432	Jun 30, 2007
	5880283	Dec 05, 2015			
<u>GATIFLOXACIN; TEQUIN</u>					
021062 004	4980470	Dec 15, 2009	DS	I-432	Jun 30, 2007
	5880283	Dec 05, 2015			
<u>GATIFLOXACIN; TEQUIN</u>					
021678 001	4980470	Dec 15, 2009	DS		
	5880283	Dec 05, 2015	DS		
	6589955	May 09, 2022	DS	U-603	

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>GATIFLOXACIN; TEQUIN IN DEXTROSE 5% IN PLASTIC CONTAINER</u>					
021062 001	4980470 5880283	Dec 15, 2009 Dec 05, 2015	DS	I-432	Jun 30, 2007
<u>GATIFLOXACIN; TEQUIN IN DEXTROSE 5% IN PLASTIC CONTAINER</u>					
021062 002	4980470 5880283	Dec 15, 2009 Dec 05, 2015	DS	I-432	Jun 30, 2007
<u>GATIFLOXACIN; ZYMAR</u>					
021493 001	4980470 5880283 6333045	Dec 15, 2009 Dec 05, 2015 Aug 20, 2019	DS DP DP		
<u>GEFITINIB; IRESSA</u>					
021399 001	5457105 5616582 5770599	Jan 19, 2013 Jan 19, 2013 Apr 26, 2016		NCE	May 05, 2008
<u>GEMCITABINE HYDROCHLORIDE; GEMZAR</u>					
020509 001	4808614 4808614*PED 5464826 5464826*PED	May 15, 2010 Nov 15, 2010 Nov 07, 2012 May 07, 2013	DS U-146	I-499 I-428 M-40 PED PED	Jul 14, 2009 May 19, 2007 Apr 26, 2008 Oct 26, 2008 Nov 19, 2007
<u>GEMCITABINE HYDROCHLORIDE; GEMZAR</u>					
020509 002	4808614 4808614*PED 5464826 5464826*PED	May 15, 2010 Nov 15, 2010 Nov 07, 2012 May 07, 2013	DS U-146	I-499 I-428 M-40 PED PED	Jul 14, 2009 May 19, 2007 Apr 26, 2008 Oct 26, 2008 Nov 19, 2007
<u>GEMIFLOXACIN MESYLATE; FACTIVE</u>					
021158 001	5633262 5776944 5962468 6262071 6331550 6340689 6455540 6723734 6803376 6803376	Jun 15, 2015 Apr 04, 2017 Jun 15, 2015 Sep 21, 2019 Sep 21, 2019 Sep 14, 2019 Sep 21, 2019 Mar 20, 2018 Sep 21, 2019 Sep 21, 2019	DS DP U-282 U-513 U-511 U-512 U-511 DS DP DS DP	NCE	Apr 04, 2008
<u>GEMTUZUMAB OZOGAMICIN; MYLOTARG</u>					
021174 001	4970198 5079233 5585089 5606040 5693762 5739116 5767285 5773001	Nov 30, 2007 Jan 07, 2009 Dec 17, 2013 Feb 25, 2014 Dec 02, 2014 Apr 14, 2015 Jun 16, 2015 Jun 30, 2015		ODE U-320	May 17, 2007
<u>GLATIRAMER ACETATE; COPAXONE</u>					
020622 001	5981589 6054430 6342476 6362161 6620847 6939539	May 24, 2014 May 24, 2014 May 24, 2014 May 24, 2014 May 24, 2014 May 24, 2014	DS DS	U-441 U-441	
<u>GLATIRAMER ACETATE; COPAXONE</u>					
020622 002	5981589 6054430 6342476 6362161 6620847 6939539	May 24, 2014 May 24, 2014 May 24, 2014 May 24, 2014 May 24, 2014 May 24, 2014	DS DS	U-441 U-441	
<u>GLIMEPIRIDE; AMARYL</u>					
020496 001				M-54 PED	Nov 28, 2008 May 28, 2009

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>GLIMEPIRIDE; AMARYL</u>					
020496 002				M-54 PED	Nov 28, 2008 May 28, 2009
<u>GLIMEPIRIDE; AMARYL</u>					
020496 003				M-54 PED	Nov 28, 2008 May 28, 2009
<u>GLIMEPIRIDE; PIOGLITAZONE HYDROCHLORIDE; DUETACT</u>					
021925 001	4687777	Jan 17, 2011	DS		
	6150383	Jun 19, 2016		U-753	
	6211205	Jun 19, 2016		U-753	
	6303640	Aug 09, 2016		U-753	
	6329404	Jun 19, 2016	DP	U-753	
<u>GLIMEPIRIDE; PIOGLITAZONE HYDROCHLORIDE; DUETACT</u>					
021925 002	4687777	Jan 17, 2011	DS		
	6150383	Jun 19, 2016		U-753	
	6211205	Jun 19, 2016		U-753	
	6303640	Aug 09, 2016		U-753	
	6329404	Jun 19, 2016	DP	U-753	
<u>GLIMEPIRIDE; ROSIGLITAZONE MALEATE; AVANDARYL</u>					
021700 001	5002953	Sep 17, 2011	DS	DP	U-690
	5002953	Sep 17, 2011	DS	DP	U-781
	5002953*PED	Mar 17, 2012			U-781
	5741803	Apr 21, 2015	DS	DP	U-781
	5741803	Apr 21, 2015	DS	DP	U-690
	5741803*PED	Oct 21, 2015			
<u>GLIMEPIRIDE; ROSIGLITAZONE MALEATE; AVANDARYL</u>					
021700 002	5002953	Sep 17, 2011	DS	DP	U-781
	5002953	Sep 17, 2011	DS	DP	U-690
	5002953*PED	Mar 17, 2012			
	5741803	Apr 21, 2015	DS	DP	U-781
	5741803	Apr 21, 2015	DS	DP	U-690
	5741803*PED	Oct 21, 2015			
<u>GLIMEPIRIDE; ROSIGLITAZONE MALEATE; AVANDARYL</u>					
021700 003	5002953	Sep 17, 2011	DS	DP	U-781
	5002953	Sep 17, 2011	DS	DP	U-690
	5002953*PED	Mar 17, 2012			
	5741803	Apr 21, 2015	DS	DP	U-781
	5741803	Apr 21, 2015	DS	DP	U-690
	5741803*PED	Oct 21, 2015			
<u>GLIPIZIDE; GLUCOTROL XL</u>					
020329 001	5024843	Sep 05, 2009			
	5091190	Sep 05, 2009			U-111
	5545413	Jul 02, 2008			U-111
	5591454	Jan 07, 2014			U-150
<u>GLIPIZIDE; GLUCOTROL XL</u>					
020329 002	5024843	Sep 05, 2009			
	5091190	Sep 05, 2009			U-111
	5545413	Jul 02, 2008			U-111
	5591454	Jan 07, 2014			U-150
<u>GLIPIZIDE; GLUCOTROL XL</u>					
020329 003	5024843	Sep 05, 2009			
	5091190	Sep 05, 2009			U-111
	5545413	Jul 02, 2008			U-111
	5591454	Jan 07, 2014			U-111
<u>GLUTAMINE; NUTRESTORE</u>					
021667 001				NCE ODE	Jun 10, 2009 Jun 10, 2011
<u>GLYBURIDE; GLYNASE</u>					
020051 001	4735805	Mar 11, 2007			
	4916163	Apr 10, 2007			
<u>GLYBURIDE; GLYNASE</u>					
020051 002	4735805	Mar 11, 2007			
	4916163	Apr 10, 2007			

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>GLYBURIDE; GLYNASE</u>					
020051 003	4735805	Mar 11, 2007			
	4916163	Apr 10, 2007			
<u>GLYBURIDE; GLYNASE</u>					
020051 004	4735805	Mar 11, 2007			
	4916163	Apr 10, 2007			
<u>GLYBURIDE; METFORMIN HYDROCHLORIDE; GLUCOVANCE</u>					
021178 001	6303146	Jul 14, 2019	U-412	M-37	Mar 15, 2007
	6303146*PED	Jan 14, 2020	U-412	PED	Sep 15, 2007
<u>GLYBURIDE; METFORMIN HYDROCHLORIDE; GLUCOVANCE</u>					
021178 002	6303146	Jul 14, 2019	U-412	M-37	Mar 15, 2007
	6303146*PED	Jan 14, 2020	U-412	PED	Sep 15, 2007
<u>GLYBURIDE; METFORMIN HYDROCHLORIDE; GLUCOVANCE</u>					
021178 003	6303146	Jul 14, 2019	U-412	M-37	Mar 15, 2007
	6303146*PED	Jan 14, 2020	U-412	PED	Sep 15, 2007
<u>GOSERELIN ACETATE; ZOLADEX</u>					
019726 001	5366734	Nov 22, 2011			
<u>GOSERELIN ACETATE; ZOLADEX</u>					
020578 001	5366734	Nov 22, 2011			
<u>GRANISETRON HYDROCHLORIDE; KYTRIL</u>					
020239 001	4886808	Dec 29, 2007	U-89		
	5952340	Sep 14, 2016	U-519		
	6294548	May 04, 2019			
<u>GRANISETRON HYDROCHLORIDE; KYTRIL</u>					
020239 002	4886808	Dec 29, 2007	U-89		
	5952340	Sep 14, 2016	U-519		
	6294548	May 04, 2019			
<u>GRANISETRON HYDROCHLORIDE; KYTRIL</u>					
020239 003	4886808	Dec 29, 2007	DS DP U-89		
<u>GRANISETRON HYDROCHLORIDE; KYTRIL</u>					
020239 004	4886808	Dec 29, 2007	U-89		
	5952340	Sep 14, 2016	U-519		
	6294548	May 04, 2019			
<u>GRANISETRON HYDROCHLORIDE; KYTRIL</u>					
020305 001	4886808	Dec 29, 2007	U-105		
<u>GRANISETRON HYDROCHLORIDE; KYTRIL</u>					
020305 002	4886808	Dec 20, 2007	U-105		
<u>GRANISETRON HYDROCHLORIDE; KYTRIL</u>					
021238 001	4886808	Dec 29, 2007	U-105		
<u>GREPAFLOXACIN HYDROCHLORIDE; RAXAR</u>					
020695 001	5563138	Oct 08, 2013			
<u>GUAIFENESIN; HUMABID</u>					
021282 002	6372252	Apr 28, 2020	U-489		
	6955821	Apr 28, 2020	DP U-489		
<u>GUAIFENESIN; MUCINEX</u>					
021282 001	6372252	Apr 28, 2020	U-489		
	6955821	Apr 28, 2020	DP U-489		
<u>GUAIFENESIN; PSEUDOEPHEDRINE HYDROCHLORIDE; MUCINEX D</u>					
021585 001	6372252	Apr 28, 2020	DP		
	6955821	Apr 28, 2020	DP U-686		
<u>GUAIFENESIN; PSEUDOEPHEDRINE HYDROCHLORIDE; MUCINEX D</u>					
021585 002	6372252	Apr 28, 2020	DP		
	6955821	Apr 28, 2020	DP U-686		
<u>HISTRELIN ACETATE; VANTAS</u>					
021732 001	5266325	Nov 30, 2010	DP	NDF	Oct 12, 2007
	5292515	Mar 08, 2011	DP		
<u>HYALURONIDASE; AMPHADASE</u>					
021665 001				NCE	Oct 26, 2009

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
HYALURONIDASE; HYDASE					
021716 001				NCE	Oct 25, 2010
HYALURONIDASE; VITRASE					
021640 001				NCE	May 05, 2009
HYALURONIDASE; VITRASE					
021640 002				NCE	May 05, 2009
HYALURONIDASE RECOMBINANT HUMAN; HYLENEX RECOMBINANT					
021859 001				NCE	Dec 02, 2010
HYDRALAZINE HYDROCHLORIDE; ISOSORBIDE DINITRATE; BIDIL					
020727 001	4868179	Apr 22, 2007	U-71	NC	Jun 23, 2008
	6465463	Sep 08, 2020	U-71		
	6784177	Sep 08, 2020	U-71		
HYDROCHLOROTHIAZIDE; IRBESARTAN; AVALIDE					
020758 001	5270317	Sep 30, 2011			
	5270317*PED	Mar 30, 2012			
	5994348	Jun 07, 2015			
	5994348*PED	Dec 07, 2015			
HYDROCHLOROTHIAZIDE; IRBESARTAN; AVALIDE					
020758 002	5270317	Sep 30, 2011			
	5270317*PED	Mar 30, 2012			
	5994348	Jun 07, 2015			
	5994348*PED	Dec 07, 2015			
HYDROCHLOROTHIAZIDE; IRBESARTAN; AVALIDE					
020758 003	5270317	Sep 30, 2011			
	5270317*PED	Mar 30, 2012			
	5994348	Jun 07, 2015			
	5994348*PED	Dec 07, 2015			
HYDROCHLOROTHIAZIDE; IRBESARTAN; AVALIDE					
020758 004	5270317	Sep 30, 2011	DS DP		
	5270317*PED	Mar 30, 2012			
	5994348	Jun 07, 2015		DP	
	5994348*PED	Dec 07, 2015			
HYDROCHLOROTHIAZIDE; LOSARTAN POTASSIUM; HYZAAR					
020387 001	5138069	Aug 11, 2009	DS	I-415	Sep 30, 2006
	5138069*PED	Feb 11, 2010			
	5153197	Oct 06, 2009	DP U-3		
	5153197	Oct 06, 2009	DP U-538		
	5153197*PED	Apr 06, 2010	U-538		
	5153197*PED	Apr 06, 2010	U-3		
	5608075	Mar 04, 2014			
	5608075*PED	Sep 04, 2014			
HYDROCHLOROTHIAZIDE; LOSARTAN POTASSIUM; HYZAAR					
020387 002	5138069	Aug 11, 2009	DS	I-415	Sep 30, 2006
	5138069*PED	Feb 11, 2010			
	5153197	Oct 06, 2009	DP U-538		
	5153197	Oct 06, 2009	DP U-3		
	5153197*PED	Apr 06, 2010	U-538		
	5153197*PED	Apr 06, 2010	U-3		
	5608075	Mar 04, 2014			
	5608075*PED	Sep 04, 2014			
HYDROCHLOROTHIAZIDE; LOSARTAN POTASSIUM; HYZAAR					
020387 003	5138069	Aug 11, 2009	DS	I-415	Sep 30, 2006
	5138069*PED	Feb 11, 2010			
	5153197	Oct 08, 2009	DP U-538		
	5153197	Oct 08, 2009	DP U-3		
	5153197*PED	Apr 06, 2010			
HYDROCHLOROTHIAZIDE; METOPROLOL SUCCINATE; DUTOPROL					
021956 001				NC	Aug 28, 2009
HYDROCHLOROTHIAZIDE; METOPROLOL SUCCINATE; DUTOPROL					
021956 002				NC	Aug 28, 2009
HYDROCHLOROTHIAZIDE; METOPROLOL SUCCINATE; DUTOPROL					
021956 003				NC	Aug 28, 2009

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>HYDROCHLOROTHIAZIDE; MOEXIPRIL HYDROCHLORIDE; UNIRETIC</u>					
020729 001	4743450	Feb 24, 2007			
<u>HYDROCHLOROTHIAZIDE; MOEXIPRIL HYDROCHLORIDE; UNIRETIC</u>					
020729 002	4743450	Feb 24, 2007			
<u>HYDROCHLOROTHIAZIDE; MOEXIPRIL HYDROCHLORIDE; UNIRETIC</u>					
020729 003	4743450	Feb 24, 2007			
<u>HYDROCHLOROTHIAZIDE; OLMESARTAN MEDOXOMIL; BENICAR HCT</u>					
021532 002	5616599 6878703	Apr 25, 2016 Nov 19, 2021	DS DP	U-500 U-3	NCE Apr 25, 2007
<u>HYDROCHLOROTHIAZIDE; OLMESARTAN MEDOXOMIL; BENICAR HCT</u>					
021532 003	5616599 6878703	Apr 25, 2016 Nov 19, 2021	DS DP	U-500 U-3	NCE Apr 25, 2007
<u>HYDROCHLOROTHIAZIDE; OLMESARTAN MEDOXOMIL; BENICAR HCT</u>					
021532 005	5616599 6878703	Apr 25, 2016 Nov 19, 2021	DS DP	U-500 U-3	NCE Apr 25, 2007
<u>HYDROCHLOROTHIAZIDE; QUINAPRIL HYDROCHLORIDE; ACCURETIC</u>					
020125 001	4743450 4743450*PED	Feb 24, 2007 Aug 24, 2007			
<u>HYDROCHLOROTHIAZIDE; QUINAPRIL HYDROCHLORIDE; ACCURETIC</u>					
020125 002	4743450 4743450*PED	Feb 24, 2007 Aug 24, 2007			
<u>HYDROCHLOROTHIAZIDE; QUINAPRIL HYDROCHLORIDE; ACCURETIC</u>					
020125 003	4743450 4743450*PED	Feb 24, 2007 Aug 24, 2007			
<u>HYDROCHLOROTHIAZIDE; TELMISARTAN; MICARDIS HCT</u>					
021162 001	5591762 6358986	Jan 07, 2014 Jan 10, 2020		U-3	
<u>HYDROCHLOROTHIAZIDE; TELMISARTAN; MICARDIS HCT</u>					
021162 002	5591762 6358986	Jan 07, 2014 Jan 10, 2020		U-3	
<u>HYDROCHLOROTHIAZIDE; VALSARTAN; DIOVAN HCT</u>					
020818 001	5399578 6294197	Mar 21, 2012 Jun 18, 2017		U-3 U-3	
<u>HYDROCHLOROTHIAZIDE; VALSARTAN; DIOVAN HCT</u>					
020818 002	5399578 6294197	Mar 21, 2012 Jun 18, 2017		U-3 U-3	
<u>HYDROCHLOROTHIAZIDE; VALSARTAN; DIOVAN HCT</u>					
020818 003	5399578 6294197	Mar 21, 2012 Jun 18, 2017		U-3 U-3	
<u>HYDROCHLOROTHIAZIDE; VALSARTAN; DIOVAN HCT</u>					
020818 004				NS	Apr 28, 2009
<u>HYDROCHLOROTHIAZIDE; VALSARTAN; DIOVAN HCT</u>					
020818 005				NS	Apr 28, 2009
<u>HYDROCODONE BITARTRATE; IBUPROFEN; VICOPROFEN</u>					
020716 001	6348216 6599531	Jun 10, 2017 Jun 10, 2017			
<u>HYDROCORTISONE BUTYRATE; LOCROID LIPOCREAM</u>					
020769 001	5635497	Jun 03, 2014			
<u>HYDROMORPHONE HYDROCHLORIDE; PALLADONE</u>					
021044 001	5958452 5965161 5968551 6294195 6335033 6706281 6743442	Nov 04, 2014 Nov 04, 2014 Dec 24, 2011 Dec 24, 2011 Nov 04, 2014 Nov 04, 2014 Nov 04, 2014	DP DP DP DP DP DP DP	U-616 U-616 U-616	NDF Sep 24, 2007

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
HYDROMORPHONE HYDROCHLORIDE; PALLADONE					
021044 002	5958452	Nov 04, 2014	DP	NDF	Sep 24, 2007
	5965161	Nov 04, 2014	DP U-616		
	5968551	Dec 24, 2011	DP		
	6294195	Dec 24, 2011	DP		
	6335033	Nov 04, 2014	DP U-616		
	6706281	Nov 04, 2014	DP U-616		
	6743442	Nov 04, 2014	DP		
HYDROMORPHONE HYDROCHLORIDE; PALLADONE					
021044 003	5958452	Nov 04, 2014	DP	NDF	Sep 24, 2007
	5965161	Nov 04, 2014	DP U-616		
	5968551	Dec 24, 2011	DP		
	6294195	Dec 24, 2011	DP		
	6335033	Nov 04, 2014	DP U-616		
	6706281	Nov 04, 2014	DP U-616		
	6743442	Nov 04, 2014	DP		
HYDROMORPHONE HYDROCHLORIDE; PALLADONE					
021044 004	5958452	Nov 04, 2014	DP	NDF	Sep 24, 2007
	5965161	Nov 04, 2014	DP U-616		
	5968551	Dec 24, 2011	DP		
	6294195	Dec 24, 2011	DP		
	6335033	Nov 04, 2014	DP U-616		
	6706281	Nov 04, 2014	DP U-616		
	6743442	Nov 04, 2014	DP		
HYDROXOCOBALAMIN; CYANOKIT					
022041 002				NP	Dec 15, 2009
IBANDRONATE SODIUM; BONIVA					
021455 001	4927814	Jul 09, 2007	DS DP	NCE	May 16, 2008
	6143326	Apr 21, 2017	U-642		
	6294196	Oct 07, 2019	DP		
IBANDRONATE SODIUM; BONIVA					
021455 002	4927814	Jul 09, 2007	DS DP	D-96 NCE NS	Mar 24, 2008 May 16, 2008 Mar 24, 2008
	6294196	Oct 07, 2019	DP		
IBANDRONATE SODIUM; BONIVA					
021858 001	4927814	Jul 09, 2007	DS DP	NDF	Jan 06, 2009
	5662918	Sep 02, 2014	DP		
IBUPROFEN; CHILDREN'S ADVIL					
020589 001	4788220	Jul 08, 2007			
	4788220*PED	Jan 08, 2008			
IBUPROFEN; CHILDREN'S ELIXSURE					
021604 001				NP	Jan 07, 2007
IBUPROFEN; CHILDREN'S MOTRIN					
020516 001	5374659	Dec 20, 2011			
	5374659*PED	Jun 20, 2012			
IBUPROFEN; CHILDREN'S MOTRIN					
020601 001	5215755	Jun 01, 2010			
	5215755*PED	Dec 01, 2010			
IBUPROFEN; CHILDREN'S MOTRIN					
020603 001	5374659	Dec 20, 2011			
	5374659*PED	Jun 20, 2012			
IBUPROFEN; IBUPROFEN					
021472 001	6251426	Jun 25, 2018			
IBUPROFEN; JUNIOR STRENGTH MOTRIN					
020601 003	5215755	Jun 01, 2010			
	5215755*PED	Dec 01, 2010			
IBUPROFEN; MOTRIN					
019842 001	5374659	Dec 20, 2011			
	5374659*PED	Jun 20, 2012			

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>IBUPROFEN; MOTRIN</u>					
020135 001	5215755	Jun 01, 2010			
	5215755*PED	Dec 01, 2010			
	5320855	Jun 14, 2011			
	5320855*PED	Dec 14, 2011			
<u>IBUPROFEN; MOTRIN</u>					
020135 002	5215755	Jun 01, 2010			
	5215755*PED	Dec 01, 2010			
	5320855	Jun 14, 2011			
	5320855*PED	Dec 14, 2011			
<u>IBUPROFEN LYSINE; NEOPROFEN</u>					
021903 001				NE	Apr 13, 2009
				ODE	Apr 13, 2013
<u>IBUPROFEN POTASSIUM; PSEUDOEPHEDRINE HYDROCHLORIDE; ADVIL COLD AND SINUS</u>					
021374 001	5071643	Dec 10, 2008			
	5071643*PED	Jun 10, 2009			
	5360615	Dec 10, 2008			
	5360615*PED	Jun 10, 2009			
<u>IBUPROFEN; OXYCODONE HYDROCHLORIDE; COMBUNOX</u>					
021378 001				NC	Nov 26, 2007
<u>IBUPROFEN; PSEUDOEPHEDRINE HYDROCHLORIDE; CHILDREN'S MOTRIN COLD</u>					
021128 001	6211246	Jun 10, 2019			
<u>IBUTILIDE FUMARATE; CORVERT</u>					
020491 001	5155268	Dec 28, 2009			
<u>ICODEXTRIN; EXTRANEAL</u>					
021321 001	4761237	Aug 09, 2009	U-495	I-445	Dec 17, 2007
	4886789	Dec 12, 2006	U-495	NCE	Dec 20, 2007
	6077836	Jun 20, 2017	U-495	ODE	Dec 20, 2009
	6248726	Jun 19, 2018	U-495		
<u>IFOSFAMIDE; IFEX</u>					
019763 001	4882452	Mar 03, 2008			
<u>IFOSFAMIDE; IFEX</u>					
019763 002	4882452	Mar 03, 2008			
<u>IFOSFAMIDE; MESNA; IFEX/MESNEX KIT</u>					
019763 003	4882452	Mar 03, 2008			
<u>IFOSFAMIDE; MESNA; IFEX/MESNEX KIT</u>					
019763 004	4882452	Mar 03, 2008			
<u>ILOPROST; VENTAVIS</u>					
021779 001				M-43	Aug 24, 2008
				NCE	Dec 29, 2009
				ODE	Dec 29, 2011
<u>IMATINIB MESYLATE; GLEEVEC</u>					
021335 001	5521184	Jan 04, 2015	DS DP	ODE	Feb 01, 2009
	5521184*PED	Jul 04, 2015		ODE	May 10, 2008
	6894051	May 23, 2019	DS DP U-649	PED	Nov 10, 2008
	6894051*PED	Nov 23, 2019		PED	Nov 20, 2006
	6958335	Dec 19, 2021	DS DP	PED	Aug 01, 2009
	6958335*PED	Jun 19, 2022		PED	Nov 10, 2006
<u>IMATINIB MESYLATE; GLEEVEC</u>					
021335 002	5521184	Jan 04, 2015		ODE	Feb 01, 2009
	5521184*PED	Jul 04, 2015		ODE	May 10, 2008
	6894051	May 23, 2019	DS DP U-649	PED	Nov 10, 2008
	6894051*PED	Nov 23, 2019		PED	Nov 20, 2006
	6958335	Dec 19, 2021	DS DP	PED	Aug 01, 2009
	6958335*PED	Jun 19, 2022		PED	Nov 10, 2006

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>IMATINIB MESYLATE; GLEEVEC</u>					
021588 001	5521184	Jan 04, 2015	DS DP	I-513	Oct 19, 2009
	5521184*PED	Jul 04, 2015		I-512	Oct 19, 2009
	6894051	May 23, 2019	DS DP U-649	I-511	Oct 19, 2009
	6894051*PED	Nov 23, 2019		I-510	Oct 19, 2009
	6958335	Dec 19, 2021	DS DP	I-514	Oct 19, 2009
	6958335*PED	Jun 19, 2022		ODE	Oct 19, 2013
				ODE	Oct 19, 2013
				ODE	Oct 19, 2013
				ODE	Oct 19, 2013
				ODE	Oct 19, 2013
				ODE	Feb 01, 2009
				ODE	May 10, 2008
				PED	Nov 10, 2006
				PED	Nov 20, 2006
				PED	Nov 10, 2008
				PED	Aug 01, 2009
<u>IMATINIB MESYLATE; GLEEVEC</u>					
021588 002	5521184	Jan 04, 2015		I-511	Oct 19, 2009
	5521184*PED	Jul 04, 2015		I-510	Oct 19, 2009
	6894051	May 23, 2019	DS DP U-649	I-513	Oct 19, 2009
	6894051*PED	Nov 23, 2019		I-512	Oct 19, 2009
	6958335	Dec 19, 2021	DS DP	I-514	Oct 19, 2009
	6958335*PED	Jun 19, 2022		ODE	Oct 19, 2013
				ODE	Oct 19, 2013
				ODE	Oct 19, 2013
				ODE	Oct 19, 2013
				ODE	Oct 19, 2013
				ODE	Feb 01, 2009
				ODE	May 10, 2008
				PED	Nov 10, 2008
				PED	Nov 20, 2006
				PED	Aug 01, 2009
				PED	Nov 10, 2006
<u>IMIGLUCERASE; CEREZYME</u>					
020367 001	5549892	Aug 27, 2013		U-252	
<u>IMIGLUCERASE; CEREZYME</u>					
020367 002	5549892	Aug 27, 2013		U-252	
<u>IMIQUIMOD; ALDARA</u>					
020723 001	4689338	Aug 25, 2009		U-172	I-433 Jul 14, 2007
	4689338*PED	Feb 25, 2010			I-420 Mar 02, 2007
	5238944	Aug 24, 2010			PED Sep 02, 2007
	5238944*PED	Feb 24, 2011			PED Jan 14, 2008
<u>INDINAVIR SULFATE; CRIXIVAN</u>					
020685 001	5413999	May 09, 2012		U-132	
	6645961	Mar 04, 2018	DP		
	6689761	Feb 10, 2021		U-554	
<u>INDINAVIR SULFATE; CRIXIVAN</u>					
020685 003	5413999	May 09, 2012		U-132	
	6645961	Mar 04, 2018	DP		
	6689761	Feb 10, 2021		U-554	
<u>INDINAVIR SULFATE; CRIXIVAN</u>					
020685 005	5413999	May 09, 2012		U-132	
	6645961	Mar 04, 2018	DP		
	6689761	Feb 10, 2021		U-554	
<u>INDINAVIR SULFATE; CRIXIVAN</u>					
020685 006	5413999	May 09, 2012		U-132	
	6645961	Mar 04, 2018	DP		
	6689761	Feb 10, 2021		U-554	

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
INSULIN ASPART PROTAMINE RECOMBINANT; INSULIN ASPART RECOMBINANT; NOVOLOG MIX 70/30					
021172 001	5547930	Sep 28, 2013		M-48	Nov 21, 2008
	5618913	Apr 08, 2014			
	5618913*PED	Oct 08, 2014			
	5834422	Sep 28, 2013	U-471		
	5840680	Sep 28, 2013	U-471		
	5866538	Jun 19, 2017	DP		
	5948751	Jun 19, 2017			
INSULIN ASPART RECOMBINANT; NOVOLOG					
020986 001	5618913	Apr 08, 2014		M-44	Sep 13, 2008
	5618913*PED	Oct 08, 2014		PED	Mar 13, 2009
	5866538	Jun 20, 2017			
	5866538*PED	Dec 20, 2017			
INSULIN DETEMIR RECOMBINANT; LEVEMIR					
021536 001	5750497	May 12, 2015	DS DP	U-668	Oct 19, 2008
	5866538	Jun 20, 2017	DP		
	6011007	Feb 02, 2014	DS DP	U-668	Jun 16, 2010
	6869930	Feb 02, 2014	DS DP	U-668	
INSULIN GLARGINE RECOMBINANT; LANTUS					
021081 001	5656722	Sep 12, 2014			
	5656722*PED	Mar 12, 2015			
	6100376	Nov 06, 2009	DS DP	U-771	
	6100376*PED	May 06, 2010			
INSULIN GLULISINE RECOMBINANT; APIDRA					
021629 001	6221633	Jun 18, 2018	DS DP	U-471	NCE Apr 16, 2009
INSULIN LISPRO PROTAMINE RECOMBINANT; INSULIN LISPRO RECOMBINANT; HUMALOG MIX 50/50					
021018 001	5461031	Jun 26, 2014			
	5474978	Jun 16, 2014			
	5514646	May 07, 2013			
	5747642	Jun 16, 2014			
INSULIN LISPRO PROTAMINE RECOMBINANT; INSULIN LISPRO RECOMBINANT; HUMALOG MIX 75/25					
021017 001	5461031	Jun 16, 2014			
	5474978	Jun 16, 2014			
	5514646	May 07, 2013			
	5747642	Jun 16, 2014			
INSULIN LISPRO RECOMBINANT; HUMALOG					
020563 001	5474978	Jun 16, 2014		U-534	Jun 02, 2007
	5514646	May 07, 2013		U-534	
INSULIN LISPRO RECOMBINANT; HUMALOG PEN					
020563 002	5474978	Jun 16, 2014		U-534	Jun 02, 2007
	5514646	May 07, 2013		U-534	
INSULIN RECOMBINANT HUMAN; EXUBERA					
021868 001	5740794	Apr 21, 2015	DP		Jan 27, 2009
	5997848	Mar 07, 2014		U-704	
	6051256	Mar 07, 2014	DP		
	6257233	May 14, 2019		U-704	
	6423344	Mar 07, 2014	DP		
	6543448	Sep 21, 2014	DP		
	6546929	May 14, 2019		U-704	
	6582728	Jun 24, 2020	DP		
	6592904	Mar 07, 2014	DP		
	6685967	Sep 11, 2018	DP		
	6737045	Mar 07, 2014		U-704	
	RE37872	Feb 12, 2010	DP		
	RE38385	Feb 12, 2010	DP		

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE	
INSULIN RECOMBINANT HUMAN; EXUBERA						
021868 002	5740794	Apr 21, 2015	DP	NP	Jan 27, 2009	
	5997848	Mar 07, 2014	U-704			
	6051256	Mar 07, 2014	DP	U-704		
	6257233	May 14, 2019				
	6423344	Mar 07, 2014	DP	U-704		
	6543448	Sep 21, 2014	DP			
	6546929	May 14, 2019		U-704		
	6582728	Jun 24, 2020	DP			
	6592904	Mar 07, 2014	DP	U-704		
	6685967	Sep 11, 2018	DP			
	6737045	Mar 07, 2014				
	RE37872	Feb 12, 2010	DP			
RE38385	Feb 12, 2010	DP				
IODINE POVACRYLEX; ISOPROPYL ALCOHOL; DURAPREP						
021586 001				NC	Sep 29, 2009	
IODINE POVACRYLEX; ISOPROPYL ALCOHOL; DURAPREP						
021586 002				NC	Sep 29, 2009	
IODIXANOL; VISIPAQUE 270						
020351 001	5349085	Sep 20, 2011				
IODIXANOL; VISIPAQUE 270						
020808 001	5349085	Sep 20, 2011				
IODIXANOL; VISIPAQUE 320						
020351 002	5349085	Sep 20, 2011				
IODIXANOL; VISIPAQUE 320						
020808 002	5349085	Sep 20, 2011				
IOXILAN; OXILAN-300						
020316 001	4954348	Dec 21, 2009				
IOXILAN; OXILAN-350						
020316 002	4954348	Dec 21, 2009				
IPRATROPIUM BROMIDE; ATROVENT HFA						
021527 001	5676930	Oct 14, 2014	DP	NP	Nov 17, 2007	
	5683677	Nov 04, 2014	DP			
	5695743	Jul 06, 2010	DP			U-610
	5766573	Nov 28, 2009				U-610
	6739333	May 26, 2020	DP			
	6983743	May 26, 2020	DP			
IRBESARTAN; AVAPRO						
020757 001	5270317	Sep 30, 2011				
	5270317*PED	Mar 30, 2012				
	6342247	Jun 07, 2015				
	6342247*PED	Dec 07, 2015				
IRBESARTAN; AVAPRO						
020757 002	5270317	Sep 30, 2011				
	5270317*PED	Mar 30, 2012				
	6342247	Jun 07, 2015				
	6342247*PED	Dec 07, 2015				
IRBESARTAN; AVAPRO						
020757 003	5270317	Sep 30, 2011				
	5270317*PED	Mar 30, 2012				
	6342247	Jun 07, 2015				
	6342247*PED	Dec 07, 2015				
IRINOTECAN HYDROCHLORIDE; CAMPTOSAR						
020571 001	4604463	Aug 20, 2007		M-34 PED	Jun 24, 2007	
	4604463*PED	Feb 20, 2008				
	6403569	Apr 28, 2020	U-449		Dec 24, 2007	
	6403569*PED	Oct 28, 2020				
	6794370	May 01, 2020	U-606			
	6794370*PED	Nov 01, 2020				
IRON DEXTRAN; DEXFERRUM						
040024 001	5624668	Sep 29, 2015				

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>IRON SUCROSE; VENOFER</u>					
021135	001			I-474	Oct 17, 2008
				I-459	Jun 17, 2008
<u>IRON SUCROSE; VENOFER</u>					
021135	002			I-474	Oct 17, 2008
				I-459	Jun 17, 2008
<u>IRON SUCROSE; VENOFER</u>					
021135	003			I-474	Oct 17, 2008
				I-459	Jun 17, 2008
<u>ISRADIPINE; DYNACIRC CR</u>					
020336	001	4816263	U-3		
		4946687			
		4950486			
		5030456			
<u>ISRADIPINE; DYNACIRC CR</u>					
020336	002	4816263	U-3		
		4946687			
		4950486			
		5030456			
<u>ITRACONAZOLE; SPORANOX</u>					
020083	001	5633015			
<u>ITRACONAZOLE; SPORANOX</u>					
020657	001	5707975			
		6407079			
<u>ITRACONAZOLE; SPORANOX</u>					
020966	001	6407079			
<u>KETOCONAZOLE; NIZORAL A-D</u>					
020310	001	5456851			
<u>KETOCONAZOLE; XOLEGEL</u>					
021946	001			NDF	Jul 28, 2009
<u>KETOROLAC TROMETHAMINE; ACULAR</u>					
019700	001	5110493	U-75 U-75		
		5110493*PED			
<u>KETOROLAC TROMETHAMINE; ACULAR LS</u>					
021528	001	5110493			
		5110493*PED			
<u>KETOTIFEN FUMARATE; KETOTIFEN FUMARATE</u>					
077354	001			PC	Dec 30, 2006
<u>KUNECATECHINS; VEREGEN</u>					
021902	001	5795911	U-172 U-172	NP	Oct 31, 2009
		5968973			
<u>LAMIVUDINE; EPIVIR</u>					
020564	001	5047407			
		5047407*PED			
		5905082			
		5905082*PED			
<u>LAMIVUDINE; EPIVIR</u>					
020564	003	5047407			
		5047407*PED			
		5905082			
		5905082*PED			
<u>LAMIVUDINE; EPIVIR</u>					
020596	001	5047407			
		5047407*PED			
		6004968			
		6004968*PED			

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>LAMIVUDINE; EPIVIR-HBV</u>					
021003 001	5047407	Nov 17, 2009			
	5047407*PED	May 17, 2010			
	5532246	Jul 02, 2013	U-250		
	5532246*PED	Jan 02, 2014	U-250		
	5905082	May 18, 2016			
	5905082*PED	Nov 18, 2016			
<u>LAMIVUDINE; EPIVIR-HBV</u>					
021004 001	5047407	Nov 17, 2009			
	5047407*PED	May 17, 2010			
	5532246	Jul 02, 2013	U-250		
	5532246*PED	Jan 02, 2014	U-250		
	6004968	Mar 20, 2018			
	6004968*PED	Sep 20, 2018			
<u>LAMIVUDINE; ZIDOVUDINE; COMBIVIR</u>					
020857 001	5047407	Nov 17, 2009			
	5047407*PED	May 17, 2010			
	5859021	May 15, 2012	U-248		
	5905082	May 18, 2016	U-248		
	5905082*PED	Nov 18, 2016	U-248		
<u>LAMOTRIGINE; LAMICTAL</u>					
020241 001	4602017	Jul 22, 2008	U-106	I-516	Sep 22, 2009
<u>LAMOTRIGINE; LAMICTAL</u>					
020241 002	4602017	Jul 22, 2008	U-106	I-516	Sep 22, 2009
<u>LAMOTRIGINE; LAMICTAL</u>					
020241 003	4602017	Jul 22, 2008	U-106	I-516	Sep 22, 2009
<u>LAMOTRIGINE; LAMICTAL</u>					
020241 004	4602017	Jul 22, 2008	U-106	I-516	Sep 22, 2009
<u>LAMOTRIGINE; LAMICTAL</u>					
020241 005	4602017	Jul 22, 2008	U-106	I-516	Sep 22, 2009
<u>LAMOTRIGINE; LAMICTAL</u>					
020241 006	4602017	Jul 22, 2008	U-106	I-516	Sep 22, 2009
<u>LAMOTRIGINE; LAMICTAL CD</u>					
020764 001	4602017	Jul 22, 2008	U-106	I-516	Sep 22, 2009
	5698226	Jan 29, 2012			
<u>LAMOTRIGINE; LAMICTAL CD</u>					
020764 002	4602017	Jul 22, 2008	U-106	I-516	Sep 22, 2009
	5698226	Jan 29, 2012			
<u>LAMOTRIGINE; LAMICTAL CD</u>					
020764 003	4602017	Jul 22, 2008	U-106	I-516	Sep 22, 2009
	5698226	Jan 29, 2012			
<u>LAMOTRIGINE; LAMICTAL CD</u>					
020764 004	4602017	Jul 22, 2008	U-106	I-516	Sep 22, 2009
	5698226	Jan 29, 2012			
<u>LAMOTRIGINE; LAMOTRIGINE</u>					
076420 001				PC	Feb 25, 2007
<u>LAMOTRIGINE; LAMOTRIGINE</u>					
076420 002				PC	Dec 25, 2006
<u>LANSOPRAZOLE; PREVACID</u>					
020406 001	4628098	May 10, 2009		NPP	Jun 17, 2007
	5013743	Feb 12, 2010	U-452		
	5026560	Jun 25, 2008			
	5045321	Sep 03, 2008			
	5093132	Sep 03, 2008			
	5433959	Sep 03, 2008			
	6749864	Feb 13, 2007	DP		

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE		PATENT CODES		EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE	
<u>LANSOPRAZOLE; PREVACID</u>								
020406 002	4628098	May 10, 2009	U-452	NPP	Jun 17, 2007			
	5013743	Feb 12, 2010						
	5026560	Jun 25, 2008						
	5045321	Sep 03, 2008						
	5093132	Sep 03, 2008						
	5433959	Sep 03, 2008						
	6749864	Feb 13, 2007				DP		
<u>LANSOPRAZOLE; PREVACID</u>								
021281 001	4628098	May 10, 2009	U-452	NPP	Jun 17, 2007			
	5013743	Feb 12, 2010						
	5026560	Jun 25, 2008						
	5045321	Sep 03, 2008						
	5093132	Sep 03, 2008						
	5433959	Sep 03, 2008						
	6749864	Feb 13, 2007				DP		
<u>LANSOPRAZOLE; PREVACID</u>								
021281 002	4628098	May 10, 2009	U-452	NPP	Jun 17, 2007			
	5013743	Feb 12, 2010						
	5026560	Jun 25, 2008						
	5045321	Sep 03, 2008						
	5093132	Sep 03, 2008						
	5433959	Sep 03, 2008						
	6749864	Feb 13, 2007				DP		
<u>LANSOPRAZOLE; PREVACID</u>								
021428 001	4628098	May 10, 2009	U-452	NPP	Jun 17, 2007			
	5013743	Feb 12, 2010						
	5026560	Jun 25, 2008						
	5045321	Sep 03, 2008						
	5093132	Sep 03, 2008						
	5433959	Sep 03, 2008						
	5464632	Nov 07, 2012						
	6123962	Feb 13, 2007						
	6328994	May 17, 2019						
6749864	Feb 13, 2007	DP						
<u>LANSOPRAZOLE; PREVACID</u>								
021428 002	4628098	May 10, 2009	U-452	NPP	Jun 17, 2007			
	5013743	Feb 12, 2010						
	5026560	Jun 25, 2008						
	5045321	Sep 03, 2008						
	5093132	Sep 03, 2008						
	5433959	Sep 03, 2008						
	5464632	Nov 07, 2012						
	6123962	Feb 13, 2007						
	6328994	May 17, 2019						
6749864	Feb 13, 2007	DP						
<u>LANSOPRAZOLE; PREVACID IV</u>								
021566 001				NDF	May 27, 2007			
<u>LANTHANUM CARBONATE; FOSRENOL</u>								
021468 001	5968976	Mar 19, 2016	DP	U-613	NCE	Oct 26, 2009		
<u>LANTHANUM CARBONATE; FOSRENOL</u>								
021468 002	5968976	Mar 19, 2016	DP	U-613	NCE	Oct 26, 2009		
<u>LANTHANUM CARBONATE; FOSRENOL</u>								
021468 003	5968976	Mar 19, 2016	DP	U-613	NCE	Oct 26, 2009		
<u>LANTHANUM CARBONATE; FOSRENOL</u>								
021468 004	5968976	Mar 19, 2016	DP	U-613	NCE	Oct 26, 2009		
<u>LATANOPROST; XALATAN</u>								
020597 001	4599353	Jul 28, 2006		U-260				
	5296504	Mar 22, 2011	DP	U-778				
	5422368	Mar 22, 2011	DP	U-778				
	6429226	Sep 06, 2009	DP	U-778				

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE		PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE	
<u>LEFLUNOMIDE; ARAVA</u>							
020905 001					M-32	Mar 05, 2007	
					PED	Sep 05, 2007	
					PED	Dec 13, 2006	
<u>LEFLUNOMIDE; ARAVA</u>							
020905 002					M-32	Mar 05, 2007	
					PED	Sep 05, 2007	
					PED	Dec 13, 2006	
<u>LEFLUNOMIDE; ARAVA</u>							
020905 003					M-32	Mar 05, 2007	
					PED	Sep 05, 2007	
					PED	Dec 13, 2006	
<u>LENALIDOMIDE; REVLIMID</u>							
021880 001	5635517	Jul 24, 2016	DS		NCE	Dec 27, 2010	
	6045501	Aug 28, 2018			ODE	Jun 29, 2013	
	6281230	Jul 24, 2016			ODE	Dec 27, 2012	
	6315720	Oct 23, 2020	DP				
	6555554	Jul 24, 2016					
	6561976	Aug 28, 2018					
	6561977	Oct 23, 2020					
	6755784	Oct 23, 2020					
	6908432	Aug 28, 2018					
	7119106	Jul 24, 2016	DP				
<u>LENALIDOMIDE; REVLIMID</u>							
021880 002	5635517	Jul 24, 2016	DS		NCE	Dec 27, 2010	
	6045501	Aug 28, 2018			ODE	Jun 29, 2013	
	6281230	Jul 24, 2016			ODE	Dec 27, 2012	
	6315720	Oct 23, 2020	DP				
	6555554	Jul 24, 2016					
	6561976	Aug 28, 2018					
	6561977	Oct 23, 2020					
	6755784	Oct 23, 2020					
	6908432	Aug 28, 2018					
	7119106	Jul 24, 2016	DP				
<u>LENALIDOMIDE; REVLIMID</u>							
021880 003	5635517	Jul 24, 2016	DS		I-500	Jun 29, 2009	
	6045501	Aug 28, 2018			NCE	Dec 27, 2010	
	6281230	Jul 24, 2016			ODE	Jun 29, 2013	
	6315720	Oct 23, 2020	DP				
	6555554	Jul 24, 2016					
	6561976	Aug 28, 2018					
	6561977	Oct 23, 2020					
	6755784	Oct 23, 2020					
	6908432	Aug 28, 2018					
	7119106	Jul 24, 2016	DP				
<u>LENALIDOMIDE; REVLIMID</u>							
021880 004	5635517	Jul 24, 2016	DS		I-500	Jun 29, 2009	
	6045501	Aug 28, 2018			NCE	Dec 27, 2010	
	6281230	Jul 24, 2016			ODE	Jun 29, 2013	
	6315720	Oct 23, 2020	DP				
	6555554	Jul 24, 2016					
	6561976	Aug 28, 2018					
	6561977	Oct 23, 2020					
	6755784	Oct 23, 2020					
	6908432	Aug 28, 2018					
	7119106	Jul 24, 2016	DP				
<u>LEPIRUDIN RECOMBINANT; REFLUDAN</u>							
020807 001	5180668	Jan 19, 2010					
<u>LETROZOLE; FEMARA</u>							
020726 001	4978672	Jun 03, 2011		U-203	I-481	Dec 28, 2008	
					I-446	Oct 29, 2007	

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
LEUPROLIDE ACETATE; ELIGARD					
021343 001	4938763	Oct 03, 2008			
	5278201	Jan 11, 2011			
	5324519	Oct 20, 2011			
	5599552	Feb 04, 2014			
	5733950	Oct 03, 2008			
	5739176	Oct 03, 2008			
LEUPROLIDE ACETATE; ELIGARD					
021379 001	4938763	Oct 03, 2008			
	5278201	Jan 11, 2011			
	5324519	Oct 20, 2011			
	5599552	Feb 04, 2014			
	5733950	Oct 03, 2008			
	5739176	Oct 03, 2008			
LEUPROLIDE ACETATE; ELIGARD					
021488 001	4938763	Oct 03, 2008			
	5278201	Jan 11, 2011			
	5324519	Oct 20, 2011			
	5599552	Feb 04, 2014			
	5733950	Oct 03, 2008			
	5739176	Oct 03, 2008			
LEUPROLIDE ACETATE; ELIGARD					
021731 001	4938763	Oct 03, 2008	DP	U-621	NP Dec 14, 2007
	5278201	Jan 11, 2011	DP		
	5324519	Jun 28, 2011	DP		
	5599552	Feb 04, 2014	DP	U-621	
	5739176	Oct 03, 2008	DP	U-621	
	6395293	Sep 28, 2013	DP		
	6565874	Oct 28, 2018	DP	U-621	
	6626870	Mar 27, 2020	DP		
	6773714	Oct 28, 2018		U-621	
	RE37950	Oct 03, 2008	DP	U-621	
LEUPROLIDE ACETATE; LUPRON DEPOT					
019732 001	4849228	Jul 18, 2006			
	5575987	Sep 02, 2013			
	5631020	May 20, 2014			
	5716640	Sep 02, 2013			
	6036976	Dec 13, 2016			
LEUPROLIDE ACETATE; LUPRON DEPOT					
020011 001	4849228	Jul 18, 2006			
	5575987	Sep 02, 2013			
	5631021	May 20, 2014			
	5716640	Sep 02, 2013			
LEUPROLIDE ACETATE; LUPRON DEPOT					
020517 001	4849228	Jul 18, 2006			
	5480656	Jan 02, 2013			
	5575987	Nov 19, 2013			
	5631020	May 20, 2014			
	5643607	Jan 02, 2013			
	5716640	Sep 02, 2013			
	5814342	Feb 01, 2011			
	6036976	Dec 13, 2016			
LEUPROLIDE ACETATE; LUPRON DEPOT-3					
020708 001	4849228	Jul 18, 2006			
	5480656	Jan 02, 2013			
	5575987	Nov 19, 2013			
	5631020	May 20, 2014			
	5643607	Jan 02, 2013			
	5716640	Sep 02, 2013			
	5814342	Feb 01, 2011			
	6036976	Dec 13, 2016			

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
LEUPROLIDE ACETATE; LUPRON DEPOT-4					
020517 002	4849228	Jul 18, 2006			
	5480656	Jan 02, 2013			
	5575987	Nov 19, 2013			
	5631020	May 20, 2014			
	5643607	Jan 02, 2013			
	5716640	Sep 02, 2013			
	5814342	Feb 01, 2011			
	6036976	Dec 13, 2016			
LEUPROLIDE ACETATE; LUPRON DEPOT-PED					
020263 002	4849228	Jul 18, 2006			
	5575987	Sep 02, 2013			
	5631020	May 20, 2014			
	5716640	Sep 02, 2013			
	6036976	Dec 13, 2016			
LEUPROLIDE ACETATE; LUPRON DEPOT-PED					
020263 003	4849228	Jul 18, 2006			
	5575987	Sep 02, 2013			
	5631020	May 20, 2014			
	5716640	Sep 02, 2013			
	6036976	Dec 13, 2013			
LEUPROLIDE ACETATE; LUPRON DEPOT-PED					
020263 004	4849228	Jul 18, 2006			
	5575987	Sep 02, 2013			
	5631020	May 20, 2014			
	5716640	Sep 02, 2013			
	6036976	Dec 13, 2013			
LEUPROLIDE ACETATE; LUPRON DEPOT-PED					
020263 005	4849228	Jul 18, 2006			
	5575987	Sep 02, 2013			
	5631020	May 20, 2014			
	5716640	Sep 02, 2013			
	6036976	Dec 13, 2016			
LEUPROLIDE ACETATE; LUPRON DEPOT-PED					
020263 006	4849228	Jul 18, 2006			
	5575987	Sep 02, 2013			
	5631020	May 20, 2014			
	5716640	Sep 02, 2013			
	6036976	Dec 13, 2016			
LEUPROLIDE ACETATE; VIADUR					
021088 001	5728396	Jan 30, 2017		U-316	
	5932547	Jun 13, 2017			
	5985305	Jan 30, 2017			
	6113938	Jul 24, 2018			
	6124261	Jun 13, 2017			
	6132420	Jan 30, 2017			
	6156331	Jan 30, 2017			
	6235712	Jun 13, 2017			
	6375978	Dec 17, 2018			
	6395292	Jan 30, 2017			
LEVALBUTEROL HYDROCHLORIDE; XOPENEX					
020837 001	5362755	Nov 08, 2011		U-332	
	5547994	Aug 20, 2013		U-332	
	5760090	Jan 05, 2010		U-332	
	5844002	Jan 05, 2010		U-332	
	6083993	Jan 05, 2010		U-332	
	6451289	Mar 21, 2021			
LEVALBUTEROL HYDROCHLORIDE; XOPENEX					
020837 002	5362755	Nov 08, 2011		U-332	
	5547994	Aug 20, 2013		U-332	
	5760090	Jan 05, 2010		U-332	
	5844002	Jan 05, 2010		U-332	
	6083993	Jan 05, 2010		U-332	
	6451289	Mar 21, 2021			

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE		PATENT CODES			EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE	
LEVALBUTEROL HYDROCHLORIDE; XOPENEX									
020837 003	5362755	Nov 08, 2011				U-332			
	5547994	Aug 20, 2013				U-332			
	5760090	Jan 05, 2010				U-332			
	5844002	Jan 05, 2010				U-332			
	6083993	Jan 05, 2010				U-332			
	6451289	Mar 21, 2021							
LEVALBUTEROL HYDROCHLORIDE; XOPENEX									
020837 004	5362755	Nov 08, 2011				U-332			
	5547994	Aug 20, 2013				U-332			
	5760090	Jan 05, 2010				U-332			
	5844002	Jan 05, 2010				U-5			
	6083993	Jan 05, 2010				U-332			
	6451289	Mar 21, 2021			DP				
LEVALBUTEROL TARTRATE; XOPENEX HFA									
021730 001	5225183	Jul 06, 2010			DP		NP	Mar 11, 2008	
	5362755	Mar 25, 2013				U-636			
	5439670	Jul 06, 2010			DP				
	5547994	Aug 20, 2013				U-636			
	5605674	Feb 25, 2014			DP				
	5695743	Jul 06, 2010			DP	U-636			
	5760090	Jan 05, 2010				U-636			
	5836299	Nov 17, 2017			DP				
	5844002	Jan 05, 2010				U-636			
	6083993	Jan 05, 2010				U-636			
	6352684	Nov 28, 2009			DP				
LEVETIRACETAM; KEPPRA									
021035 001	4943639	Jul 14, 2008		DS			I-506 NPP	Aug 15, 2009 Jun 21, 2008	
LEVETIRACETAM; KEPPRA									
021035 002	4943639	Jul 14, 2008		DS			I-506 NPP	Aug 15, 2009 Jun 21, 2008	
LEVETIRACETAM; KEPPRA									
021035 003	4943639	Jul 14, 2008		DS			I-506 NPP	Aug 15, 2009 Jun 21, 2008	
LEVETIRACETAM; KEPPRA									
021035 004	4943639	Jul 14, 2008		DS			I-506 NPP	Aug 15, 2009 Jun 21, 2008	
LEVETIRACETAM; KEPPRA									
021505 001	4943639	Jul 14, 2008		DS			I-506 NPP	Aug 15, 2009 Jun 21, 2008	
LEVETIRACETAM; KEPPRA									
021872 001	4943639	Jul 14, 2008		DS			NDF	Jul 31, 2009	
LEVOBETAXOLOL HYDROCHLORIDE; BETAXON									
021114 001	4911920	Mar 27, 2007		DS	DP	U-369	M-54	Sep 28, 2009	
	4911920	Mar 27, 2007		DS	DP	U-191	PED	Mar 28, 2010	
	4911920*PED	Sep 27, 2007							
	5540918	Jul 30, 2013			DP				
	5540918*PED	Jan 30, 2014							
LEVOBUPIVACAINE HYDROCHLORIDE; CHIROCAINE									
020997 001	5708011	Oct 13, 2014				U-276			
LEVOBUPIVACAINE HYDROCHLORIDE; CHIROCAINE									
020997 002	5708011	Oct 13, 2014				U-276			
LEVOBUPIVACAINE HYDROCHLORIDE; CHIROCAINE									
020997 003	5708011	Oct 13, 2014				U-276			
LEVOCARNITINE; CARNITOR									
020182 001	6335369	Jan 18, 2021				U-433	ODE	Dec 15, 2006	
	6429230	Jan 18, 2021				U-433			
	6696493	Jan 18, 2021				U-433			
LEVOFLOXACIN; IQUIX									
021571 001	5053407	Dec 20, 2010		DS	DP	U-600	NP	Mar 01, 2007	

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>LEVOFLOXACIN; LEVAQUIN</u>					
020634 001	5053407	Dec 20, 2010		D-100 D-83	Aug 04, 2008 Oct 23, 2006
<u>LEVOFLOXACIN; LEVAQUIN</u>					
020634 002	5053407	Dec 20, 2010		D-100 D-83	Aug 04, 2008 Oct 23, 2006
<u>LEVOFLOXACIN; LEVAQUIN</u>					
020634 003	5053407	Dec 20, 2010		D-100 D-83	Aug 04, 2008 Oct 23, 2006
<u>LEVOFLOXACIN; LEVAQUIN</u>					
020635 001	5053407	Dec 20, 2010		D-100 D-83	Aug 04, 2008 Oct 23, 2006
<u>LEVOFLOXACIN; LEVAQUIN</u>					
020635 004	5053407	Dec 20, 2010		D-100 D-83	Aug 04, 2008 Oct 23, 2006
<u>LEVOFLOXACIN; LEVAQUIN</u>					
021721 001				D-100	Aug 04, 2008
<u>LEVOFLOXACIN; LEVAQUIN IN DEXTROSE 5% IN PLASTIC CONTAINER</u>					
020635 002	5053407	Dec 20, 2010		D-100 D-83	Aug 04, 2008 Oct 23, 2006
<u>LEVOFLOXACIN; LEVAQUIN IN DEXTROSE 5% IN PLASTIC CONTAINER</u>					
020635 003	5053407	Dec 20, 2010		D-100 D-83	Aug 04, 2008 Oct 23, 2006
<u>LEVOFLOXACIN; LEVAQUIN IN DEXTROSE 5% IN PLASTIC CONTAINER</u>					
020635 005	5053407	Dec 20, 2010		D-100 D-83	Aug 04, 2008 Oct 23, 2006
<u>LEVOFLOXACIN; QUIXIN</u>					
021199 001	5053407	Dec 20, 2010			
<u>LEVONORGESTREL; MIRENA</u>					
021225 001	5785053	Dec 05, 2015	DP		
<u>LEVONORGESTREL; PLAN B</u>					
021045 002				NP	Aug 24, 2009
<u>LEVOTHYROXINE SODIUM; LEVO-T</u>					
021342 001	6399101	Mar 30, 2020			
<u>LEVOTHYROXINE SODIUM; LEVO-T</u>					
021342 002	6399101	Mar 30, 2020			
<u>LEVOTHYROXINE SODIUM; LEVO-T</u>					
021342 003	6399101	Mar 30, 2020			
<u>LEVOTHYROXINE SODIUM; LEVO-T</u>					
021342 004	6399101	Mar 30, 2020			
<u>LEVOTHYROXINE SODIUM; LEVO-T</u>					
021342 005	6399101	Mar 30, 2020			
<u>LEVOTHYROXINE SODIUM; LEVO-T</u>					
021342 006	6399101	Mar 30, 2020			
<u>LEVOTHYROXINE SODIUM; LEVO-T</u>					
021342 007	6399101	Mar 30, 2020			
<u>LEVOTHYROXINE SODIUM; LEVO-T</u>					
021342 008	6399101	Mar 30, 2020			
<u>LEVOTHYROXINE SODIUM; LEVO-T</u>					
021342 009	6399101	Mar 30, 2020			
<u>LEVOTHYROXINE SODIUM; LEVO-T</u>					
021342 010	6399101	Mar 30, 2020			
<u>LEVOTHYROXINE SODIUM; LEVO-T</u>					
021342 011	6399101	Mar 30, 2020			

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE		PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>LEVOTHYROXINE SODIUM; LEVOXYL</u>						
021301 001	6555581	Feb	15, 2022	DP	U-759	
	7067148	Feb	15, 2022			
	7101569	Aug	14, 2022			
<u>LEVOTHYROXINE SODIUM; LEVOXYL</u>						
021301 002	6555581	Feb	15, 2022	DP	U-759	
	7067148	Feb	15, 2022			
	7101569	Aug	14, 2022			
<u>LEVOTHYROXINE SODIUM; LEVOXYL</u>						
021301 003	6555581	Feb	15, 2022	DP	U-759	
	7067148	Feb	15, 2022			
	7101569	Aug	14, 2022			
<u>LEVOTHYROXINE SODIUM; LEVOXYL</u>						
021301 004	6555581	Feb	15, 2022	DP	U-759	
	7067148	Feb	15, 2022			
	7101569	Aug	14, 2022			
<u>LEVOTHYROXINE SODIUM; LEVOXYL</u>						
021301 005	6555581	Feb	15, 2022	DP	U-759	
	7067148	Feb	15, 2022			
	7101569	Aug	14, 2022			
<u>LEVOTHYROXINE SODIUM; LEVOXYL</u>						
021301 006	6555581	Feb	15, 2022	DP	U-759	
	7067148	Feb	15, 2022			
	7101569	Aug	14, 2022			
<u>LEVOTHYROXINE SODIUM; LEVOXYL</u>						
021301 007	6555581	Feb	15, 2022	DP	U-759	
	7067148	Feb	15, 2022			
	7101569	Aug	14, 2022			
<u>LEVOTHYROXINE SODIUM; LEVOXYL</u>						
021301 008	6555581	Feb	15, 2022	DP	U-759	
	7067148	Feb	15, 2022			
	7101569	Aug	14, 2022			
<u>LEVOTHYROXINE SODIUM; LEVOXYL</u>						
021301 009	6555581	Feb	15, 2022	DP	U-759	
	7067148	Feb	15, 2022			
	7101569	Aug	14, 2022			
<u>LEVOTHYROXINE SODIUM; LEVOXYL</u>						
021301 010	6555581	Feb	15, 2022	DP	U-759	
	7067148	Feb	15, 2022			
	7101569	Aug	14, 2022			
<u>LEVOTHYROXINE SODIUM; LEVOXYL</u>						
021301 011	6555581	Feb	15, 2022	DP	U-759	
	7067148	Feb	15, 2022			
	7101569	Aug	14, 2022			
<u>LEVOTHYROXINE SODIUM; LEVOXYL</u>						
021301 012	6555581	Feb	15, 2022	DP	U-759	
	7067148	Feb	15, 2022			
	7101569	Aug	14, 2022			
<u>LIDOCAINE; LIDOCAINE</u>						
020575 001	5234957	Feb	27, 2011			
	5332576	Jul	26, 2011			
	5446070	Feb	27, 2011			
<u>LIDOCAINE; LIDOCAINE</u>						
020575 002	5234957	Feb	27, 2011			
	5332576	Jul	26, 2011			
	5446070	Feb	27, 2011			
<u>LIDOCAINE; LIDODERM</u>						
020612 001	5411738	May	02, 2012		U-485	
	5589180	Mar	17, 2009			
	5601838	May	02, 2012			
	5709869	Mar	17, 2009			
	5827529	Oct	27, 2015			

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>LIDOCAINE; PRILOCAINE; ORAQIX</u>					
021451 001	6031007	Apr 01, 2017	DP U-553	NDF	Dec 19, 2006
<u>LIDOCAINE; TETRACAINE; LIDOCAINE AND TETRACAINE</u>					
021717 001	5919479 6528086	Jul 28, 2015 Sep 28, 2019	DP DP	NP	Jun 29, 2009
<u>LIDOCAINE; TETRACAINE; SYNERA</u>					
021623 001	5658583 5919479 6306431 6465006 6546281 6780426	Jul 28, 2015 Jul 28, 2015 Jul 28, 2015 Jul 28, 2015 Jul 28, 2015 Jul 28, 2015	DP DP DP DP DP DP	NC	Jun 23, 2008
<u>LINEZOLID; ZYVOX</u>					
021130 001	5688792 5688792*PED 6514529 6514529*PED 6559305 6559305*PED	Nov 18, 2014 May 18, 2015 Mar 15, 2021 Sep 15, 2021 Jan 29, 2021 Jul 29, 2021	DS DP DS	U-319 I-431 I-402 PED PED	Jun 23, 2007 Jul 22, 2006 Dec 23, 2007 Jan 22, 2007
<u>LINEZOLID; ZYVOX</u>					
021130 002	5688792 5688792*PED 6514529 6514529*PED 6559305 6559305*PED	Nov 18, 2014 May 18, 2015 Mar 15, 2021 Sep 15, 2021 Jan 29, 2021 Jul 29, 2021	DS DP DS	U-319 I-431 I-402 PED PED	Jun 23, 2007 Jul 22, 2006 Dec 23, 2007 Jan 22, 2007
<u>LINEZOLID; ZYVOX</u>					
021131 001	5688792 5688792*PED 6559305 6559305*PED	Nov 18, 2014 May 18, 2015 Jan 29, 2021 Jul 29, 2021	 DS	U-319 I-431 I-402 PED PED	Jun 23, 2007 Jul 22, 2006 Dec 23, 2007 Jan 22, 2007
<u>LINEZOLID; ZYVOX</u>					
021132 001	5688792 5688792*PED 6559305 6559305*PED	Nov 18, 2014 May 18, 2015 Jan 29, 2021 Jul 29, 2021	DS DS	U-319 I-431 I-402 PED PED	Jun 23, 2007 Jul 22, 2006 Dec 23, 2007 Jan 22, 2007
<u>LISINAPRIL; PRINIVIL</u>					
019558 001				PED	Nov 29, 2006
<u>LISINAPRIL; PRINIVIL</u>					
019558 002				PED	Nov 29, 2006
<u>LISINAPRIL; PRINIVIL</u>					
019558 003				PED	Nov 29, 2006
<u>LISINAPRIL; PRINIVIL</u>					
019558 004				PED	Nov 29, 2006
<u>LISINAPRIL; PRINIVIL</u>					
019558 006				PED	Nov 29, 2006
<u>LISINAPRIL; ZESTRIL</u>					
019777 001				PED	Nov 29, 2006
<u>LISINAPRIL; ZESTRIL</u>					
019777 002				PED	Nov 29, 2006
<u>LISINAPRIL; ZESTRIL</u>					
019777 003				PED	Nov 29, 2006
<u>LISINAPRIL; ZESTRIL</u>					
019777 004				PED	Nov 29, 2006
<u>LISINAPRIL; ZESTRIL</u>					
019777 005				PED	Nov 29, 2006

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>LODOXAMIDE TROMETHAMINE; ALOMIDE</u>					
020191 001	5457126	Oct 10, 2012	U-117		
<u>LOPERAMIDE HYDROCHLORIDE; IMODIUM A-D EZ CHEWS</u>					
020448 001	5489436	Feb 06, 2013			
<u>LOPERAMIDE HYDROCHLORIDE; SIMETHICONE; IMODIUM ADVANCED</u>					
020606 001	5248505	Jul 28, 2010			
	5489436	Feb 06, 2013			
	5612054	Sep 28, 2010			
	5679376	Oct 21, 2014			
	5716641	May 21, 2012	U-226		
<u>LOPERAMIDE HYDROCHLORIDE; SIMETHICONE; IMODIUM ADVANCED</u>					
021140 001	5248505	Sep 28, 2010	U-434		
	5612054	Sep 28, 2010			
	6103260	Jul 17, 2017			
<u>LOPERAMIDE HYDROCHLORIDE; SIMETHICONE; LOPERAMIDE HYDROCHLORIDE AND SIMETHICONE</u>					
077500 001				PC	May 12, 2007
<u>LOPINAVIR; RITONAVIR; KALETRA</u>					
021226 001	5541206	Jul 30, 2013	U-348	D-99	Apr 29, 2008
	5541206*PED	Jan 30, 2014		PED	Oct 29, 2008
	5635523	Jun 30, 2014	U-352		
	5635523*PED	Dec 30, 2014			
	5648497	Jul 15, 2014			
	5648497*PED	Jan 15, 2015			
	5674882	Oct 07, 2014	U-344		
	5674882*PED	Apr 07, 2015			
	5846987	Dec 29, 2012	U-350		
	5846987*PED	Jun 29, 2013			
	5886036	Nov 19, 2013	DS DP		
	5886036*PED	May 19, 2014			
	5914332	Dec 13, 2015	U-351		
	5948436	Sep 13, 2013	DP		
	5948436*PED	Mar 13, 2014			
	6037157	Jun 26, 2016	U-346		
	6037157*PED	Dec 26, 2016			
	6232333	Nov 07, 2017			
	6232333*PED	May 07, 2018			
	6284767	Feb 14, 2016	U-401		
	6458818	Nov 07, 2017			
	6458818*PED	May 07, 2018			
	6521651	Nov 01, 2017			
	6521651*PED	May 01, 2018			
	6703403	Jun 26, 2016	U-257		
	6703403*PED	Dec 26, 2016			
	7141593	May 22, 2020	DP		
	7141593*PED	Nov 22, 2020			
<u>LOPINAVIR; RITONAVIR; KALETRA</u>					
021251 001	5541206	Jul 30, 2013	U-348	D-99	Apr 29, 2008
	5541206*PED	Jan 30, 2014		PED	Oct 29, 2008
	5635523	Jun 03, 2014	U-352		
	5635523*PED	Dec 03, 2014			
	5648497	Jul 15, 2014			
	5648497*PED	Jan 15, 2015			
	5674882	Oct 07, 2014	U-344		
	5674882*PED	Apr 07, 2015			
	5846987	Dec 29, 2012	U-350		
	5846987*PED	Jun 29, 2013			
	5886036	Nov 19, 2013	DS DP		
	5886036*PED	May 19, 2014			
	5948436	Sep 13, 2013	DP		
	5948436*PED	Mar 13, 2014			
	6037157	Jun 26, 2016	U-346		
	6037157*PED	Dec 26, 2016			
	6284767	Feb 14, 2016	U-401		
	6703403	Jun 26, 2016	U-257		
	6703403*PED	Dec 26, 2016			
	6911214	Nov 28, 2021	DP		
	6911214*PED	May 28, 2022			

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE	
LOPINAVIR; RITONAVIR; KALETRA						
021906 001	5541206	Jul 30, 2013	DS DP	U-688		
	5541206*PED	Jan 30, 2014				
	5635523	Jun 03, 2014				
	5635523*PED	Dec 03, 2014	DS DP	U-688		
	5648497	Jul 15, 2014				
	5648497*PED	Jan 15, 2015				
	5674882	Oct 07, 2014				
	5674882*PED	Apr 07, 2015				
	5846987	Dec 29, 2012				
	5846987*PED	Jun 29, 2013				
	5886036	Nov 19, 2013	DS DP			
	5886036*PED	May 19, 2014				
	5914332	Dec 13, 2015	DS DP	U-688		
	6037157	Jun 26, 2016				
	6037157*PED	Dec 26, 2016	DP	U-688		
	6284767	Feb 14, 2016				
	6703403	Jun 26, 2016				
	6703403*PED	Dec 26, 2016				
LORATADINE; CLARITIN						
019658 002	4863931	Sep 15, 2008				
	4863931*PED	Mar 15, 2009				
LORATADINE; CLARITIN						
020641 002	4863931	Sep 15, 2008				
	4863931*PED	Mar 15, 2009				
	6132758	Jun 01, 2018				
LORATADINE; CLARITIN REDITABS						
020704 002	4863931	Sep 15, 2008				
	4863931*PED	Mar 15, 2009				
LORATADINE; PSEUDOEPHEDRINE SULFATE; CLARITIN-D						
019670 002	4863931	Sep 15, 2008				
	4863931*PED	Mar 15, 2009				
LORATADINE; PSEUDOEPHEDRINE SULFATE; CLARITIN-D 24 HOUR						
020470 002	4863931	Sep 15, 2008				
	4863931*PED	Mar 15, 2009				
	5314697	Oct 23, 2012				
LOSARTAN POTASSIUM; COZAAR						
020386 001	5138069	Aug 11, 2009		NPP PED	Mar 11, 2007 Sep 11, 2007	
	5138069*PED	Feb 11, 2010				
	5153197	Oct 06, 2009		U-3 U-3 U-496 U-496		
	5153197*PED	Apr 06, 2010				
	5210079	May 11, 2010				
	5210079*PED	Nov 11, 2010				
	5608075	Mar 04, 2014				
	5608075*PED	Sep 04, 2014				
	LOSARTAN POTASSIUM; COZAAR					
020386 002	5138069	Aug 11, 2009			NPP PED	Mar 11, 2007 Sep 11, 2007
	5138069*PED	Feb 11, 2010				
	5153197	Oct 06, 2009			U-3 U-3 U-496 U-496	
	5153197*PED	Apr 06, 2010				
	5210079	May 11, 2010				
	5210079*PED	Nov 11, 2010				
	5608075	Mar 04, 2014				
	5608075*PED	Sep 04, 2014				
	LOSARTAN POTASSIUM; COZAAR					
020386 003	5138069	Aug 11, 2009			NPP PED	Mar 11, 2007 Sep 11, 2007
	5138069*PED	Feb 11, 2010				
	5153197	Oct 06, 2009			U-3 U-3 U-496 U-496	
	5153197*PED	Apr 06, 2010				
	5210079	May 11, 2010				
	5210079*PED	Nov 11, 2010				
	5608075	Mar 04, 2014				
	5608075*PED	Sep 04, 2014				
	LOTEPREDNOL ETABONATE; ALREX					
020803 001	4996335	Mar 09, 2012		DP	U-576	
	5540930	Oct 25, 2013				
	5747061	Oct 25, 2013				

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE		PATENT CODES		EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE		
<u>LOTEPREDNOL ETABONATE; LOTEMAX</u>									
020583 001	4996335	Mar	09, 2012	DP	U-575				
	5540930	Oct	25, 2013						
	5747061	Oct	25, 2013						
<u>LOTEPREDNOL ETABONATE; LOTEMAX</u>									
020841 001	4996335	Feb	26, 2008						
	5540930	Oct	25, 2013						
<u>LOVASTATIN; ALTOPREV</u>									
021316 001	5916595	Dec	12, 2017	DP	U-456				
	6080778	Mar	23, 2018						
	6485748	Dec	12, 2017						
<u>LOVASTATIN; ALTOPREV</u>									
021316 002	5916595	Dec	12, 2017	DP	U-456				
	6080778	Mar	23, 2018						
	6485748	Dec	12, 2017						
<u>LOVASTATIN; ALTOPREV</u>									
021316 003	5916595	Dec	12, 2017	DP	U-456				
	6080778	Mar	23, 2018						
	6485748	Dec	12, 2017						
<u>LOVASTATIN; ALTOPREV</u>									
021316 004	5916595	Dec	12, 2017	DP	U-456				
	6080778	Mar	23, 2018						
	6485748	Dec	12, 2017						
<u>LOVASTATIN; NIACIN; ADVICOR</u>									
021249 001	6080428	May	27, 2017	DP	U-447				
	6129930	Sep	20, 2013						
	6406715	Oct	31, 2017						
	6469035	Mar	15, 2018						
	6676967	Sep	20, 2013						
	6746691	Sep	20, 2013						
	6818229	Feb	15, 2014						
	7011848	Sep	20, 2013						
<u>LOVASTATIN; NIACIN; ADVICOR</u>									
021249 002	6080428	May	27, 2017	DP	U-447				
	6129930	Sep	20, 2013						
	6406715	Oct	31, 2017						
	6469035	Mar	15, 2018						
	6676967	Sep	20, 2013						
	6746691	Sep	20, 2013						
	6818229	Feb	15, 2014						
	7011848	Sep	20, 2013						
<u>LOVASTATIN; NIACIN; ADVICOR</u>									
021249 003	6080428	May	27, 2017		U-447				
	6129930	Sep	20, 2013						
	6406715	Oct	31, 2017						
	6469035	Mar	15, 2018						
	6676967	Sep	20, 2013						
	6746691	Sep	20, 2013						
	7011848	Sep	20, 2013						
<u>LOVASTATIN; NIACIN; ADVICOR</u>									
021249 004	6469035	Mar	15, 2018		U-768				
<u>LUBIPROSTONE; AMITIZA</u>									
021908 001	5284858	Feb	08, 2011	DS	DP	NCE	Jan	31, 2011	
	5317032	May	31, 2011	DS	DP				U-717
	6414016	Sep	05, 2020	DS	DP				U-717
	6583174	Oct	16, 2020	DS	DP				
	7064148	Aug	30, 2022	DS	DP				U-739
<u>LUTROPIN ALFA; LUVERIS</u>									
021322 001	5767251	Jun	16, 2015	DS		ODE	Oct	08, 2011	
<u>MAGNESIUM CHLORIDE; SODIUM BICARBONATE; SODIUM CHLORIDE; NORMOCARB HF 25</u>									
021910 001	5945449	Oct	31, 2017	DP	U-785	ODE	Jul	26, 2013	
<u>MAGNESIUM CHLORIDE; SODIUM BICARBONATE; SODIUM CHLORIDE; NORMOCARB HF 35</u>									
021910 002	5945449	Oct	31, 2017	DP	U-785	ODE	Jul	26, 2013	

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>MAGNESIUM HYDROXIDE; OMEPRAZOLE; SODIUM BICARBONATE; ZEGERID</u>					
021850 001	6489346	Jul 16, 2016	DS DP	U-623	
	6489346	Jul 16, 2016	DS DP	U-588	
	6645988	Jul 16, 2016	DS DP	U-623	
	6645988	Jul 16, 2016	DS DP	U-588	
	6699885	Jul 16, 2016		U-588	
	6699885	Jul 16, 2016		U-623	
<u>MAGNESIUM HYDROXIDE; OMEPRAZOLE; SODIUM BICARBONATE; ZEGERID</u>					
021850 002	6489346	Jul 16, 2016	DS DP	U-588	
	6489346	Jul 16, 2016	DS DP	U-623	
	6645988	Jul 16, 2016	DS DP	U-588	
	6645988	Jul 16, 2016	DS DP	U-623	
	6699885	Jul 16, 2016		U-623	
	6699885	Jul 16, 2016		U-588	
<u>MANGAFODIPIR TRISODIUM; TESLASCAN</u>					
020652 001	4933456	Nov 27, 2011			
	4992554	Feb 12, 2008			
	5091169	Feb 25, 2009			
	5223243	Jun 29, 2010		U-237	
<u>MASOPROCOL; ACTINEX</u>					
019940 001	4695590	Apr 17, 2008			
	5008294	Apr 15, 2008		U-68	
<u>MECASERMIN RECOMBINANT; INCRELEX</u>					
021839 001	5681814	Oct 28, 2014	DP	NCE	Aug 30, 2010
	5824642	Oct 20, 2015		ODE	Aug 30, 2012
	6207640	Apr 07, 2014		U-681	
<u>MECASERMIN RINFABATE RECOMBINANT; IPLEX</u>					
021884 001	5200509	Apr 06, 2010	DS	NP	Dec 12, 2008
	5681818	Oct 28, 2014		ODE	Dec 12, 2012
<u>MEDROXYPROGESTERONE ACETATE; DEPO-SUBQ PROVERA 104</u>					
021583 001	6495534	May 15, 2020	DP	I-451 NP	Mar 25, 2008 Dec 17, 2007
<u>MEGESTROL ACETATE; MEGACE</u>					
020264 001	5338732	Aug 16, 2011			
<u>MEGESTROL ACETATE; MEGACE ES</u>					
021778 001	5145684	Jan 25, 2011	DP	U-673	
	6592903	Sep 21, 2020	DP		
	7101576	Apr 22, 2024		U-755	
<u>MELOXICAM; MOBIC</u>					
020938 001				I-469	Aug 11, 2008
				I-430	Jul 16, 2007
				ODE	Aug 11, 2012
				PED	Feb 11, 2013
				PED	Feb 11, 2009
				PED	Jan 16, 2008
<u>MELOXICAM; MOBIC</u>					
020938 002				I-469	Aug 11, 2008
				I-430	Jul 16, 2007
				ODE	Aug 11, 2012
				PED	Feb 11, 2013
				PED	Feb 11, 2009
				PED	Jan 16, 2008
<u>MELOXICAM; MOBIC</u>					
021530 001	6184220	Mar 25, 2019	DP	I-469	Aug 11, 2008
	6184220*PED	Sep 25, 2019		I-430	Jul 16, 2007
				ODE	Aug 11, 2012
				PED	Feb 11, 2013
				PED	Dec 01, 2007
				PED	Feb 11, 2009

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>MELPHALAN HYDROCHLORIDE; ALKERAN</u>					
020207 001	4997651	Nov 18, 2008			
<u>MEMANTINE HYDROCHLORIDE; NAMENDA</u>					
021487 001	5061703	Apr 11, 2010	U-539	NCE	Oct 16, 2008
	5614560	Mar 25, 2014	U-539		
<u>MEMANTINE HYDROCHLORIDE; NAMENDA</u>					
021487 002	5061703	Apr 11, 2010	U-539	NCE	Oct 16, 2008
	5614560	Mar 25, 2014	U-539		
<u>MEMANTINE HYDROCHLORIDE; NAMENDA</u>					
021487 003	5061703	Apr 11, 2010	U-539	NCE	Oct 16, 2008
	5614560	Mar 25, 2014	U-539		
<u>MEMANTINE HYDROCHLORIDE; NAMENDA</u>					
021487 004	5061703	Apr 11, 2010	U-539	NCE	Oct 16, 2008
	5614560	Mar 25, 2014	U-539		
<u>MEMANTINE HYDROCHLORIDE; NAMENDA</u>					
021627 001				NCE	Oct 16, 2008
<u>MENOTROPINS (FSH;LH); MENOPUR</u>					
021663 001				NP	Oct 29, 2007
<u>MEQUINOL; TRETINOIN; SOLAGE</u>					
020922 001	5194247	Dec 10, 2013	DP U-294		
	5470567	Mar 19, 2010	U-294		
	6353029	Aug 24, 2020			
<u>MESALAMINE; ASACOL</u>					
019651 001	5541170	Jul 30, 2013	U-141		
	5541171	Jul 30, 2013	U-141		
<u>MESALAMINE; CANASA</u>					
021252 002				D-92 NS	Nov 05, 2007 Nov 05, 2007
<u>MESNA; MESNEX</u>					
019884 001	5696172	Oct 06, 2013			
<u>MESNA; MESNEX</u>					
020855 001	5252341	Jul 16, 2011			
	5262169	Jul 16, 2011			
<u>METAXALONE; SKELAXIN</u>					
013217 001	6407128	Dec 03, 2021	U-189		
	6683102	Dec 03, 2021	U-189		
<u>METAXALONE; SKELAXIN</u>					
013217 003	6407128	Dec 03, 2021	U-189		
	6683102	Dec 03, 2021	U-189		
<u>METFORMIN HYDROCHLORIDE; FORTAMET</u>					
021574 001	6099859	Mar 20, 2018	DP	NP	Apr 27, 2007
	6495162	Mar 20, 2018	DP		
	6790459	Mar 17, 2021	U-604		
	6866866	Mar 17, 2021	DP		
<u>METFORMIN HYDROCHLORIDE; FORTAMET</u>					
021574 002	6099859	Mar 20, 2018	DP	NP	Apr 27, 2007
	6495162	Mar 20, 2018	DP		
	6790459	Mar 17, 2021	U-604		
	6866866	Mar 17, 2021	DP		
<u>METFORMIN HYDROCHLORIDE; GLUCOPHAGE XR</u>					
021202 001	6475521	Mar 19, 2018			
	6660300	Mar 19, 2018	U-542		
<u>METFORMIN HYDROCHLORIDE; GLUCOPHAGE XR</u>					
021202 004	6475521	Mar 19, 2018			
	6660300	Mar 19, 2018	U-542		
<u>METFORMIN HYDROCHLORIDE; GLUMETZA</u>					
021748 001	6340475	Sep 19, 2016	DS DP U-669	NP	Jun 03, 2008
	6488962	Jun 20, 2020	DS DP		
	6723340	Oct 25, 2021	DS DP		

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
METFORMIN HYDROCHLORIDE; GLUMETZA					
021748 002	6340475	Sep 19, 2016	DS DP U-669	NP	Jun 03, 2008
	6488962	Jun 20, 2020	DS DP		
	6723340	Oct 25, 2021	DS DP		
METFORMIN HYDROCHLORIDE; RIOMET					
021591 001	6890957	Sep 14, 2023	DP		
METFORMIN HYDROCHLORIDE; ROSIGLITAZONE MALEATE; AVANDAMET					
021410 001	5002953	Sep 17, 2011	DS DP U-691	I-494	May 19, 2009
	5002953	Sep 17, 2011	DS DP U-690		
	5002953	Sep 17, 2011	DS DP U-734		
	5002953	Sep 17, 2011	DS DP U-493		
	5002953*PED	Mar 17, 2012			
	5741803	Apr 21, 2015	DS DP U-493		
	5741803	Apr 21, 2015	DS DP U-734		
	5741803*PED	Oct 21, 2015			
	5965584	Jun 19, 2016	U-493		
	6166042	Jun 19, 2016	U-493		
	6288095	Feb 11, 2017	U-493		
	6288095*PED	Aug 11, 2017			
	METFORMIN HYDROCHLORIDE; ROSIGLITAZONE MALEATE; AVANDAMET				
021410 002	5002953	Sep 17, 2011	DS DP U-690	I-494	May 19, 2009
	5002953	Sep 17, 2011	DS DP U-734		
	5002953	Sep 17, 2011	DS DP U-493		
	5002953	Sep 17, 2011	DS DP U-691		
	5002953*PED	Mar 17, 2012			
	5741803	Apr 21, 2015	DS DP U-493		
	5741803	Apr 21, 2015	DS DP U-734		
	5741803*PED	Oct 21, 2015			
	5965584	Jun 19, 2016	U-493		
	6166042	Jun 19, 2016	U-493		
	6288095	Feb 11, 2017	U-493		
	6288095*PED	Aug 11, 2017			
	METFORMIN HYDROCHLORIDE; ROSIGLITAZONE MALEATE; AVANDAMET				
021410 003	5002953	Sep 17, 2011	DS DP U-690	I-494	May 19, 2009
	5002953	Sep 17, 2011	DS DP U-734		
	5002953	Sep 17, 2011	DS DP U-691		
	5002953	Sep 17, 2011	DS DP U-493		
	5002953*PED	Mar 17, 2012			
	5741803	Apr 21, 2015	DS DP U-493		
	5741803	Apr 21, 2015	DS DP U-734		
	5741803*PED	Oct 21, 2015			
	5965584	Jun 19, 2016	U-493		
	6166042	Jun 19, 2016	U-493		
	6288095	Feb 11, 2017	U-493		
	6288095*PED	Aug 11, 2017			
	METFORMIN HYDROCHLORIDE; ROSIGLITAZONE MALEATE; AVANDAMET				
021410 004	5002953	Sep 17, 2011	DS DP U-493	I-494	May 19, 2009
	5002953	Sep 17, 2011	DS DP U-690		
	5002953	Sep 17, 2011	DS DP U-691		
	5002953	Sep 17, 2011	DS DP U-734		
	5002953*PED	Mar 17, 2012			
	5741803	Apr 21, 2015	DS DP U-493		
	5741803	Apr 21, 2015	DS DP U-734		
	5741803*PED	Oct 21, 2015			
METFORMIN HYDROCHLORIDE; ROSIGLITAZONE MALEATE; AVANDAMET					
021410 005	5002953	Sep 17, 2011	DS DP U-493	I-494	May 19, 2009
	5002953	Sep 17, 2011	DS DP U-691		
	5002953	Sep 17, 2011	DS DP U-690		
	5002953	Sep 17, 2011	DS DP U-734		
	5002953*PED	Mar 17, 2012			
	5741803	Apr 21, 2015	DS DP U-493		
	5741803	Apr 21, 2015	DS DP U-734		
	5741803*PED	Oct 21, 2015			

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE		PATENT CODES		EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE	
METFORMIN; PIOGLITAZONE HYDROCHLORIDE; ACTOPLUS MET								
021842 001	4687777	Jan	17,	2011	DS	DP	U-679	
	5965584	Jun	19,	2016				
	6166042	Jun	19,	2016				
	6166043	Jun	19,	2016				
	6172090	Jun	19,	2016				
METFORMIN; PIOGLITAZONE HYDROCHLORIDE; ACTOPLUS MET								
021842 002	4687777	Jan	17,	2011	DS	DP	U-679	
	5965584	Jun	19,	2016				
	6166042	Jun	19,	2016				
	6166043	Jun	19,	2016				
	6172090	Jun	19,	2016				
METHOXSALEN; UVADEX								
020969 001	4845075	Apr	11,	2009			U-273	
	4999375	Mar	12,	2008				
	5036102	Jul	30,	2008				
METHYL AMINOLEVULINATE HYDROCHLORIDE; METVIXIA								
021415 001							NE	Jul 27, 2007
METHYLPHENIDATE; DAYTRANA								
021514 001	5958446	Dec	12,	2012		DP	NDF	Apr 06, 2009
	6210705	Sep	30,	2018				
	6348211	Sep	30,	2018				
METHYLPHENIDATE; DAYTRANA								
021514 002	5958446	Dec	12,	2012		DP	NDF	Apr 06, 2009
	6210705	Sep	30,	2018				
	6348211	Sep	30,	2018				
METHYLPHENIDATE; DAYTRANA								
021514 003	5958446	Dec	12,	2012		DP	NDF	Apr 06, 2009
	6210705	Sep	30,	2018				
	6348211	Sep	30,	2018				
METHYLPHENIDATE; DAYTRANA								
021514 004	5958446	Dec	12,	2012		DP	NDF	Apr 06, 2009
	6210705	Sep	30,	2018				
	6348211	Sep	30,	2018				
METHYLPHENIDATE HYDROCHLORIDE; CONCERTA								
021121 001	6919373	Jul	31,	2017		U-666	D-94	Oct 21, 2007
	6919373*PED	Jan	31,	2018			NPP	Oct 21, 2007
	6930129	Jul	31,	2017			PED	Apr 21, 2008
	6930129*PED	Jan	31,	2018			PED	Apr 21, 2008
METHYLPHENIDATE HYDROCHLORIDE; CONCERTA								
021121 002	6919373	Jul	31,	2017		U-666	D-94	Oct 21, 2007
	6919373*PED	Jan	31,	2018			NPP	Oct 21, 2007
	6930129	Jul	31,	2017			PED	Apr 21, 2008
	6930129*PED	Jan	31,	2018			PED	Apr 21, 2008
METHYLPHENIDATE HYDROCHLORIDE; CONCERTA								
021121 003	6919373	Jul	31,	2017		U-666	D-94	Oct 21, 2007
	6919373*PED	Jan	31,	2018			NPP	Oct 21, 2007
	6930129	Jul	31,	2017			PED	Apr 21, 2008
	6930129*PED	Jan	31,	2018			PED	Apr 21, 2008
METHYLPHENIDATE HYDROCHLORIDE; CONCERTA								
021121 004	6919373	Jul	31,	2017		U-666	D-94	Oct 21, 2007
	6919373*PED	Jan	31,	2018			NPP	Oct 21, 2007
	6930129	Jul	31,	2017			PED	Apr 21, 2008
	6930129*PED	Jan	31,	2018			PED	Apr 21, 2008
METHYLPHENIDATE HYDROCHLORIDE; METADATE CD								
021259 001	6344215	Oct	27,	2020	DP			
METHYLPHENIDATE HYDROCHLORIDE; METADATE CD								
021259 002	6344215	Oct	27,	2020	DP			
METHYLPHENIDATE HYDROCHLORIDE; METADATE CD								
021259 003	6344215	Oct	27,	2020	DP			

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>METHYLPHENIDATE HYDROCHLORIDE; METADATE CD</u>					
021259 004	6344215	Oct 27, 2020	DP		
<u>METHYLPHENIDATE HYDROCHLORIDE; RITALIN LA</u>					
021284 001	5837284	Dec 04, 2015	DP		
	6228398	Nov 01, 2019	DP	U-472	
	6635284	Dec 04, 2015	DP	U-591	
<u>METHYLPHENIDATE HYDROCHLORIDE; RITALIN LA</u>					
021284 002	5837284	Dec 04, 2015	DP		
	6228398	Nov 01, 2019	DP	U-472	
	6635284	Dec 04, 2015	DP	U-591	
<u>METHYLPHENIDATE HYDROCHLORIDE; RITALIN LA</u>					
021284 003	5837284	Dec 04, 2015	DP		
	6228398	Nov 01, 2019	DP	U-472	
	6635284	Dec 04, 2015	DP	U-591	
<u>METHYLPHENIDATE HYDROCHLORIDE; RITALIN LA</u>					
021284 004	5837284	Dec 04, 2015	DP		
	6228398	Nov 01, 2019	DP		
	6635284	Dec 04, 2015	DP	U-591	
<u>METOCLOPRAMIDE HYDROCHLORIDE; REGLAN ODT</u>					
021793 001	6024981	Apr 09, 2018	DP		
	6221392	Apr 09, 2018	DP		
<u>METOCLOPRAMIDE HYDROCHLORIDE; REGLAN ODT</u>					
021793 002	6024981	Apr 09, 2018	DP		
	6221392	Apr 09, 2018	DP		
<u>METOPROLOL FUMARATE; LOPRESSOR</u>					
019786 003	4892739	Jan 09, 2007			
<u>METOPROLOL FUMARATE; LOPRESSOR</u>					
019786 004	4892739	Jan 09, 2007			
<u>METOPROLOL SUCCINATE; TOPROL-XL</u>					
019962 001	4927640	May 22, 2007	DP	D-95	Feb 15, 2008
	4927640*PED	Nov 22, 2007		PED	Aug 15, 2008
	4957745	Sep 18, 2007	DP	U-107	
	4957745*PED	Mar 18, 2008			
	5001161	Sep 18, 2007	DP		
	5001161*PED	Mar 18, 2008			
	5081154	Sep 18, 2007	DS		
	5081154*PED	Mar 18, 2008			
	5246714	Sep 21, 2010			
	5246714*PED	Mar 21, 2011			
<u>METOPROLOL SUCCINATE; TOPROL-XL</u>					
019962 002	4927640	May 22, 2007	DP	D-95	Feb 15, 2008
	4927640*PED	Nov 22, 2007		PED	Aug 15, 2008
	4957745	Sep 18, 2007	DP	U-107	
	4957745*PED	Mar 18, 2008			
	5001161	Sep 18, 2007	DP		
	5001161*PED	Mar 18, 2008			
	5081154	Sep 18, 2007	DS		
	5081154*PED	Mar 18, 2008			
	5246714	Sep 21, 2010			
	5246714*PED	Mar 21, 2011			
<u>METOPROLOL SUCCINATE; TOPROL-XL</u>					
019962 003	4927640	May 22, 2007	DP	D-95	Feb 15, 2008
	4927640*PED	Nov 22, 2007		PED	Aug 15, 2008
	4957745	Sep 18, 2007	DP	U-107	
	4957745*PED	Mar 18, 2008			
	5001161	Sep 18, 2007	DP		
	5001161*PED	Mar 18, 2008			
	5081154	Sep 18, 2007	DS		
	5081154*PED	Mar 18, 2008			
	5246714	Sep 21, 2010			
	5246714*PED	Mar 21, 2011			

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>METOPROLOL SUCCINATE; TOPROL-XL</u>					
019962 004	4927640	May 22, 2007	DP	D-95	Feb 15, 2008
	4927640*PED	Nov 22, 2007		PED	Aug 15, 2008
	4957745	Sep 18, 2007	DP U-107		
	4957745*PED	Mar 18, 2008			
	5001161	Sep 18, 2007	DP U-107		
	5001161*PED	Mar 18, 2008			
	5081154	Sep 18, 2007	DS U-107		
	5081154*PED	Mar 18, 2008			
	5246714	Sep 21, 2010			
	5246714*PED	Mar 21, 2011			
<u>METRONIDAZOLE; METROGEL</u>					
021789 001	6881726	Feb 21, 2022	DP U-743	NP	Jun 30, 2008
<u>METRONIDAZOLE; METROGEL-VAGINAL</u>					
020208 001	5536743	Jul 16, 2013	U-137		
	5840744	Jan 15, 2008	U-370		
<u>MICAFUNGIN SODIUM; MYCAMINE</u>					
021506 002	5376634	Dec 27, 2011	DS DP	NCE	Mar 16, 2010
	6107458	Sep 29, 2015	DS DP U-650		
	6265536	Sep 29, 2015	DS DP U-650		
	6774104	Jan 08, 2021	DP U-650		
<u>MICONAZOLE NITRATE; MONISTAT 1 COMBINATION PACK</u>					
021308 001	5514698	Mar 21, 2014		D-90	Oct 01, 2007
	6153635	Nov 28, 2020			
<u>MICONAZOLE NITRATE; PETROLATUM, WHITE; ZINC OXIDE; VUSION</u>					
021026 001	4911932	Mar 27, 2007	DP U-718	NP	Feb 16, 2009
<u>MIGLITOL; GLYSET</u>					
020682 001	4639436	Jan 27, 2009	U-111		
<u>MIGLITOL; GLYSET</u>					
020682 002	4639436	Jan 27, 2009	U-111		
<u>MIGLITOL; GLYSET</u>					
020682 003	4639436	Jan 27, 2009	U-111		
<u>MIGLUSTAT; ZAVESCA</u>					
021348 001	5472969	May 13, 2013		NCE	Jul 31, 2008
	5525616	Jun 11, 2013		ODE	Jul 31, 2010
<u>MINOXIDIL; MEN'S ROGAINE</u>					
021812 001	6946120	Apr 20, 2019	DP U-702	NDF	Jan 20, 2009
<u>MIRTAZAPINE; REMERON SOLTAB</u>					
021208 001	5178878	Jan 12, 2010			
<u>MIRTAZAPINE; REMERON SOLTAB</u>					
021208 002	5178878	Jan 12, 2010			
<u>MIRTAZAPINE; REMERON SOLTAB</u>					
021208 003	5178878	Jan 12, 2010			
<u>MITOXANTRONE HYDROCHLORIDE; NOVANTRONE</u>					
019297 001				ODE	Oct 13, 2007
<u>MITOXANTRONE HYDROCHLORIDE; NOVANTRONE</u>					
019297 002				ODE	Oct 13, 2007
<u>MITOXANTRONE HYDROCHLORIDE; NOVANTRONE</u>					
019297 003				ODE	Oct 13, 2007
<u>MODAFINIL; PROVIGIL</u>					
020717 001	4927855	May 22, 2007	U-255	I-449	Jan 23, 2007
	4927855*PED	Nov 22, 2007		PED	Jul 23, 2007
	RE37516	Oct 06, 2014	U-255		
	RE37516*PED	Apr 06, 2015			
<u>MODAFINIL; PROVIGIL</u>					
020717 002	4927855	May 22, 2007	U-255	I-449	Jan 23, 2007
	4927855*PED	Nov 22, 2007		PED	Jul 23, 2007
	RE37516	Oct 06, 2014	U-255		
	RE37516*PED	Apr 06, 2015			

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>MOEXIPRIL HYDROCHLORIDE; UNIVASC</u>					
020312 001	4743450	Feb 24, 2007			
<u>MOEXIPRIL HYDROCHLORIDE; UNIVASC</u>					
020312 002	4743450	Feb 24, 2007			
<u>MOMETASONE FUROATE; ASMANEX TWISTHALER</u>					
021067 001	5394868	Jun 25, 2012	DP	NP	Mar 30, 2008
	5687710	Nov 18, 2014	DP		
	5829434	Nov 03, 2015	DP		
	5889015	Jan 27, 2014		U-645	
	6057307	Jan 27, 2014	DP	U-645	
	6240918	Feb 20, 2017	DP		
	6365581	Jan 27, 2014		U-645	
	6503537	Mar 17, 2018	DP		
	6677322	Jan 27, 2014		U-645	
	6949532	Jan 27, 2014		U-645	
<u>MOMETASONE FUROATE; ELOCON</u>					
019625 001	4808610	Oct 02, 2006			
	4808610*PED	Apr 02, 2007			
<u>MOMETASONE FUROATE; ELOCON</u>					
019796 001	4775529	May 21, 2007			
	4775529*PED	Nov 21, 2007			
<u>MOMETASONE FUROATE MONOHYDRATE; NASONEX</u>					
020762 001	5837699	Jan 27, 2014	DP	U-625	I-443
	5837699*PED	Jul 27, 2014			
	6127353	Oct 03, 2017	DS	DP	
	6127353*PED	Apr 03, 2018			
	6723713	Jan 27, 2014		U-625	
	6723713*PED	Jul 27, 2014			
<u>MONTELUKAST SODIUM; SINGULAIR</u>					
020829 002	5565473	Feb 03, 2012	DS	DP	U-675
	5565473	Feb 03, 2012	DS	DP	U-228
	5565473*PED	Aug 03, 2012			U-228
<u>MONTELUKAST SODIUM; SINGULAIR</u>					
020830 001	5565473	Feb 03, 2012	DS	DP	U-675
	5565473	Feb 03, 2012	DS	DP	U-228
	5565473*PED	Aug 03, 2012			U-228
<u>MONTELUKAST SODIUM; SINGULAIR</u>					
020830 002	5565473	Feb 03, 2012	DS	DP	U-228
	5565473	Feb 03, 2012	DS	DP	U-675
	5565473*PED	Aug 03, 2012			U-228
<u>MONTELUKAST SODIUM; SINGULAIR</u>					
021409 001	5565473	Feb 03, 2012	DS	DP	U-675
	5565473*PED	Aug 03, 2012			I-465
				M-49	Aug 27, 2008
					Nov 30, 2008
<u>MORPHINE SULFATE; AVINZA</u>					
021260 001	6066339	Nov 25, 2017			
<u>MORPHINE SULFATE; AVINZA</u>					
021260 002	6066339	Nov 25, 2017			
<u>MORPHINE SULFATE; AVINZA</u>					
021260 003	6066339	Nov 25, 2017			
<u>MORPHINE SULFATE; AVINZA</u>					
021260 004	6066339	Nov 25, 2017			
<u>MORPHINE SULFATE; DEPODUR</u>					
021671 001	5723147	Mar 03, 2015	DP	U-584	
	5807572	Sep 15, 2015	DP		
	5891467	Jan 31, 2017	DP		
	5931089	Jul 14, 2015		U-584	
	5962016	Jan 31, 2017		U-584	
	5997899	Sep 01, 2016	DP		
	6071534	Feb 18, 2008			
	6171613	Oct 01, 2016	DP		
	6193998	Sep 01, 2016	DP		
	6241999	Sep 01, 2016	DP		

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>MORPHINE SULFATE; DEPODUR</u>					
021671 002	5723147	Mar 03, 2015	DP	U-584	
	5807572	Sep 15, 2015	DP		
	5891467	Jan 31, 2017	DP		
	5931089	Jul 14, 2015		U-584	
	5962016	Jan 31, 2017	DP	U-584	
	5997899	Sep 01, 2016	DP		
	6071534	Feb 18, 2008			
	6171613	Oct 01, 2016	DP		
	6193998	Sep 01, 2016	DP		
	6241999	Sep 01, 2016	DP		
<u>MORPHINE SULFATE; DEPODUR</u>					
021671 003	5723147	Mar 03, 2015	DP	U-584	
	5807572	Sep 15, 2015	DP		
	5891467	Jan 31, 2017	DP		
	5931089	Jul 14, 2015		U-584	
	5962016	Jan 31, 2017	DP	U-584	
	5997899	Sep 01, 2016	DP		
	6071534	Feb 18, 2008			
	6171613	Oct 01, 2016	DP		
	6193998	Sep 01, 2016	DP		
	6241999	Sep 01, 2016	DP		
<u>MORPHINE SULFATE; KADIAN</u>					
020616 001	5202128	Apr 13, 2010			
	5378474	Mar 23, 2010			
<u>MORPHINE SULFATE; KADIAN</u>					
020616 002	5202128	Apr 13, 2010			
	5378474	Mar 23, 2010			
<u>MORPHINE SULFATE; KADIAN</u>					
020616 003	5202128	Apr 13, 2010			
	5378474	Mar 23, 2010			
<u>MORPHINE SULFATE; KADIAN</u>					
020616 004	5202128	Apr 13, 2010			
	5378474	Mar 23, 2010			
<u>MORPHINE SULFATE; KADIAN</u>					
020616 005	5202128	Apr 13, 2010			
	5378474	Mar 23, 2010			
<u>MOXIFLOXACIN HYDROCHLORIDE; AVELOX</u>					
021085 001	4990517	Dec 08, 2011	DS DP	U-298	Jun 13, 2008
	5607942	Mar 04, 2014		U-298	Nov 18, 2008
	5849752	Dec 05, 2016		U-298	
	6610327	Oct 29, 2019	DP	U-298	
<u>MOXIFLOXACIN HYDROCHLORIDE; AVELOX IN SODIUM CHLORIDE 0.8% IN PLASTIC CONTAINER</u>					
021277 001	4990517	Dec 08, 2011	DS DP	U-298	Jun 13, 2008
	5607942	Mar 04, 2014		U-298	Nov 18, 2008
	5849752	Dec 05, 2016		U-298	
	6548079	Jul 25, 2020	DP	U-298	
<u>MOXIFLOXACIN HYDROCHLORIDE; VIGAMOX</u>					
021598 001	4990517	Dec 08, 2011	DS DP	U-709	Oct 15, 2006
	4990517*PED	Jun 08, 2012			
	5607942	Mar 04, 2014			
	5607942*PED	Sep 04, 2014			
	6716830	Sep 20, 2019	DP		
	6716830*PED	Mar 29, 2020			

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>NALTREXONE; VIVITROL</u>					
021897 001	5792477	May 02, 2017	DP	NDF	Apr 13, 2009
	5916598	May 02, 2017	DP		
	6110503	May 02, 2017	DP		
	6194006	Dec 30, 2018	DP		
	6264987	May 19, 2020	DP		
	6331317	Nov 12, 2019	DP		
	6379703	Dec 30, 2018	DP		
	6379704	May 19, 2020	DP		
	6395304	Nov 12, 2019	DP		
	6403114	May 02, 2017	DP		
	6495164	May 25, 2020	DP		
	6495166	Nov 12, 2019	DP		
	6534092	May 19, 2020	DP		
	6537586	Nov 12, 2019	DP		
	6596316	Dec 30, 2018	DP		
	6667061	May 25, 2020	DP		
	6713090	Nov 12, 2019	DP		
	6939033	Nov 12, 2019	DP		
<u>NAPROXEN SODIUM; NAPRELAN</u>					
020353 001	5637320	Jun 10, 2014			
<u>NAPROXEN SODIUM; NAPRELAN</u>					
020353 002	5637320	Jun 10, 2014			
<u>NAPROXEN SODIUM; NAPRELAN</u>					
020353 003	5637320	Jun 10, 2014			
<u>NARATRIPTAN HYDROCHLORIDE; AMERGE</u>					
020763 001	4997841	Jul 07, 2010		U-232	
<u>NARATRIPTAN HYDROCHLORIDE; AMERGE</u>					
020763 002	4997841	Jul 07, 2010		U-232	
<u>NATEGLINIDE; STARLIX</u>					
021204 001	5463116	Oct 21, 2012		I-411	Oct 20, 2006
	5488150	Jan 30, 2013			
	6559188	Sep 17, 2019			
	6641841	Nov 14, 2017	DP	U-214	
	6844008	Nov 14, 2017	DP	U-214	
	6878749	Sep 15, 2020	DP		
	RE34878	Sep 08, 2009			
<u>NATEGLINIDE; STARLIX</u>					
021204 002	5463116	Oct 21, 2012		I-411	Oct 20, 2006
	5488150	Jan 30, 2013			
	6559188	Sep 17, 2019			
	6641841	Nov 14, 2017	DP	U-214	
	6844008	Nov 14, 2017	DP	U-214	
	6878749	Sep 15, 2020	DP		
	RE34878	Sep 08, 2009			
<u>NEDOCROMIL SODIUM; ALOCRIL</u>					
021009 001	4474787	Oct 02, 2006		U-304	
	RE38628	Aug 22, 2012		U-304	
<u>NEDOCROMIL SODIUM; TILADE</u>					
019660 001	4474787	Oct 02, 2006			
<u>NELARABINE; ARRANON</u>					
021877 001	5424295	Jun 13, 2012	DS DP	NCE	Oct 28, 2010
	5492897	Feb 20, 2013		ODE	Oct 28, 2012
	5747472	Feb 20, 2013		U-689	
	5747472	Feb 20, 2013		U-689	
	5747472	Feb 20, 2013		U-696	
	5747472	Feb 20, 2013		U-695	
	5821236	Feb 20, 2013		U-695	
<u>NELFINAVIR MESYLATE; VIRACEPT</u>					
020778 001	5484926	Oct 07, 2013			
	5484926*PED	Apr 07, 2014			
	5952343	Oct 07, 2013		U-257	
	5952343*PED	Apr 07, 2014		U-257	
	6162812	Oct 07, 2013		U-248	
	6162812*PED	Apr 07, 2014		U-248	

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>NELFINAVIR MESYLATE; VIRACEPT</u>					
020779 001	5484926	Oct 07, 2013			
	5484926*PED	Apr 07, 2014			
	5952343	Oct 07, 2013		U-257	
	5952343*PED	Apr 07, 2014		U-257	
	6162812	Oct 07, 2013		U-248	
	6162812*PED	Apr 07, 2014		U-248	
<u>NELFINAVIR MESYLATE; VIRACEPT</u>					
021503 001	5484926	Oct 07, 2013			
	5484926*PED	Apr 07, 2014			
	5952343	Oct 07, 2013		U-257	
	5952343*PED	Apr 07, 2014		U-257	
	6162812	Oct 07, 2013		U-248	
	6162812*PED	Apr 07, 2014		U-248	
<u>NEPAFENAC; NEVANAC</u>					
021862 001	5475034	Jun 06, 2014	U-100	NCE	Aug 19, 2010
<u>NESIRITIDE RECOMBINANT; NATRECOR</u>					
020920 001	5114923	May 19, 2009		NCE	Aug 10, 2006
<u>NEVIRAPINE; VIRAMUNE</u>					
020636 001	5366972	Nov 22, 2011		U-167	
	5366972*PED	May 22, 2012			
<u>NEVIRAPINE; VIRAMUNE</u>					
020933 001	5366972	Nov 22, 2011			
	5366972*PED	May 22, 2012			
<u>NIACIN; NIASPAN</u>					
020381 001	6080428	May 27, 2017		U-331	
	6129930	Sep 20, 2013		U-354	
	6406715	Oct 31, 2017		U-450	
	6746691	Sep 20, 2013		U-586	
	6818229	Feb 15, 2014	DP		
	7011848	Sep 20, 2013		U-712	
<u>NIACIN; NIASPAN</u>					
020381 002	6080428	May 27, 2017		U-331	
	6129930	Sep 20, 2013		U-354	
	6406715	Oct 31, 2017		U-450	
	6469035	Mar 15, 2018		U-768	
	6676967	Sep 20, 2013		U-548	
	6746691	Sep 20, 2013		U-586	
	6818229	Feb 15, 2014	DP		
	7011848	Sep 20, 2013		U-712	
<u>NIACIN; NIASPAN</u>					
020381 003	6080428	May 27, 2017		U-331	
	6129930	Sep 20, 2013		U-354	
	6406715	Oct 31, 2017		U-450	
	6469035	Mar 15, 2018		U-768	
	6676967	Sep 20, 2013		U-548	
	6746691	Sep 20, 2013		U-586	
	6818229	Feb 15, 2014	DP		
	7011848	Sep 20, 2013		U-712	
<u>NIACIN; NIASPAN</u>					
020381 004	6080428	May 27, 2017		U-331	
	6129930	Sep 20, 2013		U-354	
	6406715	Oct 31, 2017		U-450	
	6469035	Mar 15, 2018		U-768	
	6676967	Sep 20, 2013		U-548	
	6746691	Sep 20, 2013		U-586	
	6818229	Feb 15, 2014	DP		
	7011848	Sep 20, 2013		U-712	
<u>NIACIN; NIASPAN TITRATION STARTER PACK</u>					
020381 005	6080428	May 27, 2017		U-331	
	6129930	Sep 20, 2013		U-354	
	6406715	Oct 31, 2017		U-450	
	6746691	Sep 20, 2013		U-586	
	7011848	Sep 20, 2013		U-712	

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>NICARDIPINE HYDROCHLORIDE; CARDENE</u>					
019734 001	4880823	Nov 14, 2006			
	5164405	Nov 17, 2009			
<u>NICARDIPINE HYDROCHLORIDE; CARDENE SR</u>					
020005 001	5198226	Mar 30, 2010			
<u>NICARDIPINE HYDROCHLORIDE; CARDENE SR</u>					
020005 002	5198226	Mar 30, 2010			
<u>NICARDIPINE HYDROCHLORIDE; CARDENE SR</u>					
020005 003	5198226	Mar 30, 2010			
<u>NICOTINE; HABITROL</u>					
020076 004	5016652	May 21, 2008			
	5834011	May 01, 2007		U-355	
<u>NICOTINE; HABITROL</u>					
020076 005	5016652	May 21, 2008			
	5834011	May 01, 2007		U-355	
<u>NICOTINE; HABITROL</u>					
020076 006	5016652	May 21, 2008			
	5834011	May 01, 2007		U-355	
<u>NICOTINE; NICODERM CQ</u>					
020165 004	5004610	Jun 14, 2008			
	5342623	Jun 14, 2008			
	5344656	Jun 14, 2008			
	5364630	Jun 14, 2008			
	5462745	Jun 14, 2008			
	5508038	Apr 16, 2013			
	5633008	Jun 14, 2008		U-389	
	6165497	Jun 14, 2008		U-388	
<u>NICOTINE; NICODERM CQ</u>					
020165 005	5004610	Jun 14, 2008			
	5342623	Jun 14, 2008			
	5344656	Jun 14, 2008			
	5364630	Jun 14, 2008			
	5462745	Jun 14, 2008			
	5508038	Apr 16, 2013			
	5633008	Jun 14, 2008		U-389	
	6165497	Jun 14, 2008		U-388	
<u>NICOTINE; NICODERM CQ</u>					
020165 006	5004610	Jun 14, 2008			
	5342623	Jun 14, 2008			
	5344656	Jun 14, 2008			
	5364630	Jun 14, 2008			
	5462745	Jun 14, 2008			
	5508038	Apr 16, 2013			
	5633008	Jun 14, 2008		U-389	
	6165497	Jun 14, 2008		U-388	
<u>NICOTINE; NICOTROL</u>					
020385 001	5656255	Aug 12, 2014			
<u>NICOTINE; NICOTROL</u>					
020536 001	5501236	Jun 08, 2010			
	6098632	Jun 08, 2010			
<u>NICOTINE; NICOTROL</u>					
020714 001	5167242	Jun 08, 2010			
	5400808	Jun 08, 2010			
	5501236	Jun 08, 2010			
	6098632	Jun 08, 2010			
<u>NICOTINE; PROSTEP</u>					
019983 003	4946853	Aug 07, 2007		U-56	
<u>NICOTINE; PROSTEP</u>					
019983 004	4946853	Aug 07, 2007		U-56	
<u>NICOTINE POLACRILEX; COMMIT</u>					
021330 001	5110605	Aug 21, 2010			

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>NICOTINE POLACRILEX; COMMIT</u>					
021330 002	5110605	Aug 21, 2010			
<u>NICOTINE POLACRILEX; NICOTINE POLACRILEX</u>					
077007 001				PC	Aug 21, 2006
<u>NICOTINE POLACRILEX; NICOTINE POLACRILEX</u>					
077007 002				PC	Aug 21, 2006
<u>NIFEDIPINE; ADALAT CC</u>					
020198 001	4892741	Jun 08, 2008			
	5264446	Nov 23, 2010			
<u>NIFEDIPINE; ADALAT CC</u>					
020198 002	4892741	Jun 08, 2008			
	5264446	Nov 23, 2010			
<u>NIFEDIPINE; ADALAT CC</u>					
020198 003	4892741	Jun 08, 2008			
	5264446	Nov 23, 2010			
<u>NIFEDIPINE; PROCARDIA XL</u>					
019684 001	5264446	Nov 23, 2010			
<u>NIFEDIPINE; PROCARDIA XL</u>					
019684 002	5264446	Nov 23, 2010			
<u>NIFEDIPINE; PROCARDIA XL</u>					
019684 003	5264446	Nov 23, 2010			
<u>NISOLDIPINE; SULAR</u>					
020356 001	4892741	Jun 08, 2008			
<u>NISOLDIPINE; SULAR</u>					
020356 002	4892741	Jun 08, 2008			
<u>NISOLDIPINE; SULAR</u>					
020356 003	4892741	Jun 08, 2008			
<u>NISOLDIPINE; SULAR</u>					
020356 004	4892741	Jun 08, 2008			
<u>NITAZOXANIDE; ALINIA</u>					
021497 001	5387598	Feb 07, 2012	DP U-524	I-460	Jun 16, 2008
	5578621	Nov 26, 2013	DP U-525	NCE	Nov 22, 2007
	5968961	May 07, 2017	DP	NDF	Jul 21, 2007
	6020353	Sep 18, 2014	DS DP	ODE	Nov 22, 2009
<u>NITAZOXANIDE; ALINIA</u>					
021498 001	5387598	Feb 07, 2012	U-524	I-460	Jun 16, 2008
	5578621	Sep 08, 2014	U-525	NPP	Jul 21, 2007
	5965590	Jul 03, 2017	U-523	NCE	Nov 22, 2007
	5968961	May 07, 2017		ODE	Nov 22, 2009
	6020353	Sep 08, 2014		ODE	Nov 22, 2009
	6117894	May 07, 2017			
<u>NITISINONE; ORFADIN</u>					
021232 001	5006158	Apr 09, 2008		NCE	Jan 18, 2007
	5550165	Aug 27, 2013		ODE	Jan 18, 2009
<u>NITISINONE; ORFADIN</u>					
021232 002	5006158	Apr 09, 2008		NCE	Jan 18, 2007
	5550165	Aug 27, 2013		ODE	Jan 18, 2009
<u>NITISINONE; ORFADIN</u>					
021232 003	5006158	Apr 09, 2008		NCE	Jan 18, 2007
	5550165	Aug 27, 2013		ODE	Jan 18, 2009
<u>NITRIC OXIDE; INOMAX</u>					
020845 002	5485827	Jan 23, 2013	U-297	ODE	Dec 23, 2006
	5873359	Jan 23, 2013	U-297		
<u>NITRIC OXIDE; INOMAX</u>					
020845 003	5485827	Jan 23, 2013	U-297	ODE	Dec 23, 2006
	5873359	Jan 23, 2013	U-297		

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>NITROGLYCERIN; NITRO-DUR</u>					
020145 001	5186938	Feb 16, 2010			
<u>NITROGLYCERIN; NITRO-DUR</u>					
020145 002	5186938	Feb 16, 2010			
<u>NITROGLYCERIN; NITRO-DUR</u>					
020145 003	5186938	Feb 16, 2010			
<u>NITROGLYCERIN; NITRO-DUR</u>					
020145 004	5186938	Feb 16, 2010			
<u>NITROGLYCERIN; NITRO-DUR</u>					
020145 005	5186938	Feb 16, 2010			
<u>NITROGLYCERIN; NITRO-DUR</u>					
020145 006	5186938	Feb 16, 2010			
<u>NITROGLYCERIN; NITROLINGUAL PUMPSPRAY</u>					
018705 002	5186925	Feb 16, 2010			
<u>NITROGLYCERIN; NITROMIST</u>					
021780 001				NP	Nov 02, 2009
<u>NITROGLYCERIN; NITROSTAT</u>					
021134 001	6500456	Sep 16, 2018			
<u>NITROGLYCERIN; NITROSTAT</u>					
021134 002	6500456	Sep 16, 2018			
<u>NITROGLYCERIN; NITROSTAT</u>					
021134 003	6500456	Sep 16, 2018			
<u>NIZATIDINE; AXID</u>					
021494 001	6930119	Jul 17, 2022	DP	NDF	May 25, 2007
<u>OCTREOTIDE ACETATE; SANDOSTATIN</u>					
019667 001	5753618	May 19, 2015			
	5753618*PED	Nov 19, 2015			
<u>OCTREOTIDE ACETATE; SANDOSTATIN</u>					
019667 002	5753618	May 19, 2015			
	5753618*PED	Nov 19, 2015			
<u>OCTREOTIDE ACETATE; SANDOSTATIN</u>					
019667 003	5753618	May 19, 2015			
	5753618*PED	Nov 19, 2015			
<u>OCTREOTIDE ACETATE; SANDOSTATIN</u>					
019667 004	5753618	May 19, 2015			
	5753618*PED	Nov 19, 2015			
<u>OCTREOTIDE ACETATE; SANDOSTATIN</u>					
019667 005	5753618	May 19, 2015			
	5753618*PED	Nov 19, 2015			
<u>OCTREOTIDE ACETATE; SANDOSTATIN LAR</u>					
021008 001	5538739	Jul 23, 2013	DP	M-55	May 10, 2009
	5538739*PED	Jan 23, 2014		PED	Nov 10, 2009
	5639480	Jun 17, 2014	DP		
	5639480*PED	Dec 17, 2014			
	5688530	Nov 18, 2014		U-268	
	5688530*PED	May 18, 2015			
	5922338	Jul 13, 2016	DP		
	5922338*PED	Jan 13, 2017			
	5922682	Jul 13, 2016	DP		
	5922682*PED	Jan 13, 2017			

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>OCTREOTIDE ACETATE; SANDOSTATIN LAR</u>					
021008 002	5538739	Jul 23, 2013		M-55	May 10, 2009
	5538739*PED	Jan 23, 2014		PED	Nov 10, 2009
	5639480	Jun 17, 2014	DP		
	5639480*PED	Dec 17, 2014			
	5688530	Nov 18, 2014	U-268		
	5688530*PED	May 18, 2015			
	5922338	Jul 13, 2016	DP		
	5922338*PED	Jan 13, 2017			
	5922682	Jul 13, 2016	DP		
	5922682*PED	Jan 13, 2017			
<u>OCTREOTIDE ACETATE; SANDOSTATIN LAR</u>					
021008 003	5538739	Jul 23, 2013		M-55	May 25, 2009
	5538739*PED	Jan 23, 2014		PED	Nov 25, 2009
	5639480	Jun 17, 2014	DP		
	5639480*PED	Dec 17, 2014			
	5688530	Nov 18, 2014	U-268		
	5688530*PED	May 18, 2015			
	5922338	Jul 13, 2016	DP		
	5922338*PED	Jan 13, 2017			
	5922682	Jul 13, 2016	DP		
	5922682*PED	Jan 13, 2017			
<u>OFLOXACIN; FLOXIN OTIC</u>					
020799 001	5401741	Mar 27, 2012	U-407	D-84	Sep 30, 2006
<u>OLANZAPINE; ZYPREXA</u>					
020592 001	5229382	Apr 23, 2011	DS DP	U-547	I-417
	5229382	Apr 23, 2011	DS DP	U-149	I-400
<u>OLANZAPINE; ZYPREXA</u>					
020592 002	5229382	Apr 23, 2011	DS DP	U-149	I-417
	5229382	Apr 23, 2011	DS DP	U-547	I-400
<u>OLANZAPINE; ZYPREXA</u>					
020592 003	5229382	Apr 23, 2011	DS DP	U-547	I-417
	5229382	Apr 23, 2011	DS DP	U-149	I-400
<u>OLANZAPINE; ZYPREXA</u>					
020592 004	5229382	Apr 23, 2011	DS DP	U-547	I-417
	5229382	Apr 23, 2011	DS DP	U-149	I-400
<u>OLANZAPINE; ZYPREXA</u>					
020592 005	5229382	Apr 23, 2011	DS DP	U-149	I-417
	5229382	Apr 23, 2011	DS DP	U-547	I-400
<u>OLANZAPINE; ZYPREXA</u>					
020592 006	5229382	Apr 23, 2011	DS DP	U-149	I-417
	5229382	Apr 23, 2011	DS DP	U-547	I-400
<u>OLANZAPINE; ZYPREXA</u>					
021253 001	5229382	Apr 23, 2011	DS DP	U-571	NP
				NDF	Mar 29, 2007
<u>OLANZAPINE; ZYPREXA ZYDIS</u>					
021086 001	5229382	Apr 23, 2011	U-324	I-400	Jul 10, 2006
				I-417	Jan 14, 2007
<u>OLANZAPINE; ZYPREXA ZYDIS</u>					
021086 002	5229382	Apr 23, 2011	U-324	I-400	Jul 10, 2006
				I-417	Jan 14, 2007
<u>OLANZAPINE; ZYPREXA ZYDIS</u>					
021086 003	5229382	Apr 23, 2011	U-324	I-400	Jul 10, 2006
				I-417	Jan 14, 2007
<u>OLANZAPINE; ZYPREXA ZYDIS</u>					
021086 004	5229382	Apr 23, 2011	U-324	I-400	Jul 10, 2006
				I-417	Jan 14, 2007

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>OLMESARTAN MEDOXOMIL; BENICAR</u>					
021286 001	5616599 6878703	Apr 25, 2016 Nov 19, 2021	DS DP U-500 U-3	NCE	Apr 25, 2007
<u>OLMESARTAN MEDOXOMIL; BENICAR</u>					
021286 003	5616599 6878703	Apr 25, 2016 Nov 19, 2021	DS DP U-500 U-3	NCE	Apr 25, 2007
<u>OLMESARTAN MEDOXOMIL; BENICAR</u>					
021286 004	5616599 6878703	Apr 25, 2016 Nov 19, 2021	DS DP U-500 U-3	NCE	Apr 25, 2007
<u>OLOPATADINE HYDROCHLORIDE; PATADAY</u>					
021545 001	5116863 5641805 6995186	Dec 18, 2010 Jun 06, 2015 Nov 12, 2023	DS DP U-765 DP U-765	NP	Dec 22, 2007
<u>OLOPATADINE HYDROCHLORIDE; PATANOL</u>					
020688 001	4871865 4923892 5116863 5641805	Oct 03, 2006 May 08, 2007 Dec 18, 2010 Jun 06, 2015	U-174 U-184		
<u>OMEGA-3-ACID ETHYL ESTERS; OMACOR</u>					
021654 001	5502077 5656667 5698594	Mar 26, 2013 Aug 12, 2014 Dec 16, 2009	DS DS DS	NCE	Nov 10, 2009
<u>OMEPRAZOLE; PRILOSEC</u>					
019810 001	4786505 4786505*PED 4853230 4853230*PED 6147103 6147103*PED 6150380 6150380*PED 6166213 6166213*PED 6191148 6191148*PED	Apr 20, 2007 Oct 20, 2007 Apr 20, 2007 Oct 20, 2007 Oct 09, 2018 Apr 09, 2019 Nov 10, 2018 May 10, 2019 Oct 09, 2018 Apr 09, 2019 Oct 09, 2018 Apr 09, 2019	U-108 U-108 U-108 U-108		
<u>OMEPRAZOLE; PRILOSEC</u>					
019810 002	4786505 4786505*PED 4853230 4853230*PED 6147103 6147103*PED 6150380 6150380*PED 6166213 6166213*PED 6191148 6191148*PED	Apr 20, 2007 Oct 20, 2007 Apr 20, 2007 Oct 20, 2007 Oct 09, 2018 Apr 09, 2019 Nov 10, 2018 May 10, 2019 Oct 09, 2018 Apr 09, 2019 Oct 09, 2018 Apr 09, 2019	U-108 U-108 U-108 U-108		
<u>OMEPRAZOLE; PRILOSEC</u>					
019810 003	4786505 4786505*PED 4853230 4853230*PED 6147103 6147103*PED 6150380 6150380*PED 6166213 6166213*PED 6191148 6191148*PED	Apr 20, 2007 Oct 20, 2007 Apr 20, 2007 Oct 20, 2007 Oct 09, 2018 Apr 09, 2019 Nov 10, 2018 May 10, 2019 Oct 09, 2018 Apr 09, 2019 Oct 09, 2018 Apr 09, 2019	U-108 U-108 U-108 U-108		

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
OMEPRAZOLE MAGNESIUM; PRILOSEC OTC					
021229 001	4786505	Apr 20, 2007			
	4786505*PED	Oct 20, 2007			
	4853230	Apr 20, 2007			
	4853230*PED	Oct 20, 2007			
	5690960	Nov 25, 2014			
	5753265	Jun 07, 2015			
	5817338	Oct 06, 2015			
	5900424	May 04, 2016			
	6403616	Nov 15, 2019			
	6428810	Nov 03, 2019			
OMEPRAZOLE; SODIUM BICARBONATE; ZEGERID					
021636 001	5840737	Jul 16, 2016		U-588	
	6489346	Jul 16, 2016	DS DP	U-588	
	6645988	Jul 16, 2016	DS DP		
	6699885	Jul 16, 2016		U-588	
	6780882	Jul 16, 2016	DS DP		
OMEPRAZOLE; SODIUM BICARBONATE; ZEGERID					
021706 001	5840737	Jul 16, 2016	DS	U-623	NP Dec 21, 2007
	5840737	Jul 16, 2016	DS	U-624	
	6489346	Jul 16, 2016	DS DP	U-624	
	6489346	Jul 16, 2016	DS DP	U-623	
	6645988	Jul 16, 2016	DS DP		
	6699885	Jul 16, 2016		U-623	
	6699885	Jul 16, 2016		U-624	
	6780882	Jul 16, 2016	DS DP		
OMEPRAZOLE; SODIUM BICARBONATE; ZEGERID					
021849 001	6489346	Jul 16, 2016	DS DP	U-588	
	6645988	Jul 16, 2016	DS DP		
	6699885	Jul 16, 2016		U-588	
OMEPRAZOLE; SODIUM BICARBONATE; ZEGERID					
021849 002	6489346	Jul 16, 2016	DS DP	U-588	
	6489346	Jul 16, 2016	DS DP	U-623	
	6645988	Jul 16, 2016	DS DP		
	6699885	Jul 16, 2016		U-623	
	6699885	Jul 16, 2016		U-588	
ONDANSETRON; ZOFRAN ODT					
020781 001	4753789*PED	Dec 24, 2006			
	5955488	Nov 14, 2015			
	5955488*PED	May 14, 2016			
	6063802	Nov 14, 2015			
	6063802*PED	May 14, 2016			
ONDANSETRON; ZOFRAN ODT					
020781 002	4753789*PED	Dec 24, 2006			
	5955488	Nov 14, 2015			
	5955488*PED	May 14, 2016			
	6063802	Nov 14, 2015			
	6063802*PED	May 14, 2016			
ONDANSETRON HYDROCHLORIDE; ONDANSETRON HYDROCHLORIDE					
076960 001				PC	Jun 25, 2007
ONDANSETRON HYDROCHLORIDE; ZOFRAN					
020007 001	4753789*PED	Dec 24, 2006		D-98	Mar 25, 2008
				D-97	Mar 25, 2008
				PED	Sep 25, 2008
				PED	Sep 25, 2008
ONDANSETRON HYDROCHLORIDE; ZOFRAN					
020103 001	4753789*PED	Dec 24, 2006			
	5344658	Sep 06, 2011			
	5344658*PED	Mar 06, 2012			
ONDANSETRON HYDROCHLORIDE; ZOFRAN					
020103 002	4753789*PED	Dec 24, 2006			
	5344658	Sep 06, 2011			
	5344658*PED	Mar 06, 2012			

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE		PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE	
ONDANSETRON HYDROCHLORIDE; ZOFTRAN							
020103 003	4753789*PED	Dec	24, 2006	DP U-44			
	5344658	Sep	06, 2011				
	5344658*PED	Mar	06, 2012				
ONDANSETRON HYDROCHLORIDE; ZOFTRAN							
020605 001	4753789*PED	Dec	24, 2006	DP U-44			
	5854270	Nov	20, 2015				
	5854270*PED	May	20, 2016				
ONDANSETRON HYDROCHLORIDE; ZOFTRAN IN PLASTIC CONTAINER							
020403 001	4753789*PED	Dec	24, 2006				
ONDANSETRON HYDROCHLORIDE; ZOFTRAN PRESERVATIVE FREE							
020007 003	4753789*PED	Dec	24, 2006		D-98	Mar	25, 2008
					D-97	Mar	25, 2008
					PED	Sep	25, 2008
					PED	Sep	25, 2008
ORLISTAT; XENICAL							
020766 001	4598089	Jun	18, 2009		M-29	Dec	12, 2006
	4598089*PED	Dec	18, 2009		PED	Jun	12, 2007
	6004996	Jan	06, 2018				
	6004996*PED	Jul	06, 2018				
OSELTAMIVIR PHOSPHATE; TAMIFLU							
021087 001	5763483	Dec	27, 2016		I-480	Dec	21, 2008
	5763483*PED	Jun	27, 2017		PED	Jun	21, 2009
	5866601	Feb	02, 2016				
	5866601*PED	Aug	02, 2016				
	5952375	Feb	02, 2016				
	5952375*PED	Aug	02, 2016				
OSELTAMIVIR PHOSPHATE; TAMIFLU							
021246 001	5763483	Dec	27, 2016	U-376	I-480	Dec	21, 2008
	5763483*PED	Jun	27, 2017		PED	Jun	21, 2009
	5866601	Feb	02, 2016				
	5866601*PED	Aug	02, 2016				
	5952375	Feb	02, 2016				
	5952375*PED	Aug	02, 2016				
OXALIPLATIN; ELOXATIN							
021492 001	5290961	Jan	12, 2013	DS	I-441	Nov	04, 2007
	5290961*PED	Jul	12, 2013		I-425	Jan	09, 2007
	5338874	Apr	07, 2013		NCE	Aug	09, 2007
	5338874*PED	Oct	07, 2013		PED	Jul	09, 2007
	5420319	Aug	09, 2016		PED	Feb	09, 2008
	5420319*PED	Aug	09, 2016		PED	May	04, 2008
OXALIPLATIN; ELOXATIN							
021492 002	5290961	Jan	12, 2013	DS	I-441	Nov	04, 2007
	5290961*PED	Jul	12, 2013		I-425	Jan	09, 2007
	5338874	Apr	07, 2013		NCE	Aug	09, 2007
	5338874*PED	Oct	07, 2013		PED	Jul	09, 2007
	5420319	Aug	09, 2016		PED	Feb	09, 2008
	5420319*PED	Aug	09, 2016		PED	May	04, 2008
OXALIPLATIN; ELOXATIN							
021759 001	5290961	Jan	12, 2013	DS	I-441	Nov	04, 2007
	5290961*PED	Jul	12, 2013		NCE	Aug	09, 2007
	5338874	Apr	07, 2013		PED	Feb	09, 2008
	5338874*PED	Oct	07, 2013		PED	May	04, 2008
	5420319	Aug	08, 2016				
	5420319*PED	Feb	08, 2017				
	5716988	Aug	07, 2015				
	5716988*PED	Feb	07, 2016				

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE		PATENT CODES		EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE	
OXALIPLATIN; ELOXATIN								
021759 002	5290961	Jan 12, 2013	DS		I-441	Nov 04, 2007		
	5290961*PED	Jul 12, 2013			NCE	Aug 09, 2007		
	5338874	Apr 07, 2013	DS		PED	Feb 09, 2008		
	5338874*PED	Oct 07, 2013			PED	May 04, 2008		
	5420319	Aug 08, 2016	DS					
	5420319*PED	Feb 08, 2017						
	5716988	Aug 07, 2015	DP					
5716988*PED	Feb 07, 2016							
OXALIPLATIN; ELOXATIN								
021759 003					I-441	Nov 04, 2007		
					NCE	Aug 09, 2007		
					PED	Feb 09, 2008		
					PED	May 04, 2008		
OXANDROLONE; OXANDRIN								
013718 001	5872147	Dec 05, 2017		U-585	M-42	Jun 20, 2008		
	6090799	Jul 18, 2017		U-585				
	6576659	Dec 05, 2017		U-585				
	6670351	Oct 20, 2012		U-585				
	6828313	Dec 05, 2017		U-585				
OXANDROLONE; OXANDRIN								
013718 002	5872147	Dec 05, 2017		U-585	M-42	Jun 20, 2008		
	6090799	Jul 18, 2017		U-585				
	6576659	Dec 05, 2017		U-585				
	6670351	Oct 20, 2012		U-585				
	6828313	Dec 05, 2017		U-585				
OXAPROZIN POTASSIUM; DAYPRO ALTA								
020776 001	6030643	May 16, 2017		U-497				
OXCARBAZEPINE; TRILEPTAL								
021014 001	7037525	Feb 12, 2018		U-724	I-478	Oct 28, 2008		
					PED	Apr 28, 2009		
OXCARBAZEPINE; TRILEPTAL								
021014 002	7037525	Feb 12, 2018		U-724	I-478	Oct 28, 2008		
					PED	Apr 28, 2009		
OXCARBAZEPINE; TRILEPTAL								
021014 003	7037525	Feb 12, 2018		U-724	I-478	Oct 28, 2008		
					PED	Apr 28, 2009		
OXCARBAZEPINE; TRILEPTAL								
021285 001	7037525	Feb 12, 2018		U-724	I-478	Oct 28, 2008		
					PED	Apr 28, 2009		
OXYBUTYNIN; OXYTROL								
021351 002	5164190	Dec 11, 2010						
	5601839	Apr 26, 2015						
	5834010	Apr 26, 2015						
	6743441	Apr 26, 2020	DP	U-318				
OXYBUTYNIN CHLORIDE; DITROPAN								
017577 001					PED	Oct 15, 2006		
OXYBUTYNIN CHLORIDE; DITROPAN								
018211 001					PED	Oct 15, 2006		
OXYBUTYNIN CHLORIDE; DITROPAN XL								
020897 001	5674895	May 22, 2015			PED	Oct 15, 2006		
	5674895*PED	Nov 22, 2015						
	5840754	May 22, 2015						
	5840754*PED	Nov 22, 2015						
	5912268	May 22, 2015						
	5912268*PED	Nov 22, 2015						
	6124355	May 22, 2015		U-378				
	6124355*PED	Nov 22, 2015		U-378				
	6262115	May 22, 2015		U-393				
	6262115*PED	Nov 22, 2015		U-393				
	6919092	May 22, 2015	DP	U-667				
	6919092*PED	Nov 22, 2015						

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE		PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE	
<u>OXYBUTYNIN CHLORIDE; DITROPAN XL</u>							
020897 002	5674895	May	22, 2015		PED	Oct	15, 2006
	5674895*PED	Nov	22, 2015				
	5840754	May	22, 2015				
	5840754*PED	Nov	22, 2015				
	5912268	May	22, 2015				
	5912268*PED	Nov	22, 2015				
	6124355	May	22, 2015	U-378			
	6124355*PED	Nov	22, 2015	U-378			
	6262115	May	22, 2015	U-393			
	6262115*PED	Nov	22, 2015	U-393			
	6919092	May	22, 2015	DP U-667			
	6919092*PED	Nov	22, 2015				
<u>OXYBUTYNIN CHLORIDE; DITROPAN XL</u>							
020897 003	5674895	May	22, 2015		PED	Oct	15, 2006
	5674895*PED	Nov	22, 2015				
	5840754	May	22, 2015				
	5840754*PED	Nov	22, 2015				
	5912268	May	22, 2015				
	5912268*PED	Nov	22, 2015				
	6124355	May	22, 2015	U-378			
	6124355*PED	Nov	22, 2015	U-378			
	6262115	May	22, 2015	U-393			
	6262115*PED	Nov	22, 2015	U-393			
	6919092	May	22, 2015	DP U-667			
	6919092*PED	Nov	22, 2015				
<u>OXYBUTYNIN CHLORIDE; OXYBUTYNIN CHLORIDE</u>							
076644 001				PC	May	09, 2007	
<u>OXYBUTYNIN CHLORIDE; OXYBUTYNIN CHLORIDE</u>							
076702 001				PC	May	09, 2007	
<u>OXYBUTYNIN CHLORIDE; OXYBUTYNIN CHLORIDE</u>							
076745 001				PC	May	09, 2007	
<u>OXYCODONE HYDROCHLORIDE; OXYCONTIN</u>							
020553 001	4861598	Aug	29, 2006				
	4970075	Aug	29, 2006				
	5266331	Oct	26, 2007	DP			
	5508042	Apr	16, 2013		U-443		
	5549912	Oct	26, 2007	DP			
	5656295	Oct	26, 2007		U-443		
<u>OXYCODONE HYDROCHLORIDE; OXYCONTIN</u>							
020553 002	4861598	Aug	29, 2006				
	4970075	Aug	29, 2006				
	5266331	Oct	26, 2007	DP			
	5508042	Apr	16, 2013		U-443		
	5549912	Oct	26, 2007	DP			
	5656295	Oct	26, 2007		U-443		
<u>OXYCODONE HYDROCHLORIDE; OXYCONTIN</u>							
020553 003	4861598	Aug	29, 2006				
	4970075	Aug	29, 2006				
	5266331	Oct	26, 2007	DP			
	5508042	Apr	16, 2013		U-443		
	5549912	Oct	26, 2007	DP			
	5656295	Oct	26, 2007		U-443		
<u>OXYCODONE HYDROCHLORIDE; OXYCONTIN</u>							
020553 004	4861598	Aug	29, 2006				
	4970075	Aug	29, 2006				
	5266331	Oct	26, 2007	DP			
	5508042	Apr	16, 2013		U-443		
	5549912	Oct	26, 2007	DP			
	5656295	Oct	26, 2007		U-443		

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>OXYCODONE HYDROCHLORIDE; OXYCONTIN</u>					
020553 005	4861598	Aug 29, 2006			
	4970075	Aug 29, 2006			
	5266331	Oct 26, 2007	DP		
	5508042	Apr 16, 2013		U-443	
	5549912	Oct 26, 2007	DP		
	5656295	Oct 26, 2007		U-443	
<u>OXYCODONE HYDROCHLORIDE; OXYCONTIN</u>					
020553 006	5266331	Oct 26, 2007	DP		
	5508042	Apr 16, 2013		U-443	
	5549912	Oct 26, 2007	DP		
	5656295	Oct 26, 2007	DP	U-443	
<u>OXYCODONE HYDROCHLORIDE; OXYCONTIN</u>					
020553 007	5266331	Oct 26, 2007	DP		
	5508042	Apr 16, 2013		U-443	
	5549912	Oct 26, 2007	DP		
	5656295	Oct 26, 2007	DP	U-443	
<u>OXYCODONE HYDROCHLORIDE; OXYCONTIN</u>					
020553 008	5266331	Oct 26, 2007	DP		
	5508042	Apr 16, 2013		U-443	
	5549912	Oct 26, 2007	DP		
	5656295	Oct 26, 2007	DP	U-443	
<u>OXYMORPHONE HYDROCHLORIDE; OPANA</u>					
021611 001				NDF	Jun 22, 2009
<u>OXYMORPHONE HYDROCHLORIDE; OPANA</u>					
021611 002				NDF	Jun 22, 2009
<u>OXYMORPHONE HYDROCHLORIDE; OPANA ER</u>					
021610 001	5128143	Sep 19, 2008	DP	NDF	Jun 22, 2009
<u>OXYMORPHONE HYDROCHLORIDE; OPANA ER</u>					
021610 002	5128143	Sep 19, 2008	DP	NDF	Jun 22, 2009
<u>OXYMORPHONE HYDROCHLORIDE; OPANA ER</u>					
021610 003	5128143	Sep 19, 2008	DP	NDF	Jun 22, 2009
<u>OXYMORPHONE HYDROCHLORIDE; OPANA ER</u>					
021610 004	5128143	Sep 19, 2008	DP	NDF	Jun 22, 2009
<u>PACLITAXEL; ABRAXANE</u>					
021660 001	5439686	Feb 22, 2013	DP		
	5498421	Mar 12, 2013	DP	U-634	
	6096331	Feb 22, 2013	DP	U-633	
	6506405	Feb 22, 2013	DP	U-633	
	6537579	Feb 22, 2013		U-632	
	6749868	Feb 22, 2013	DP		
	6753006	Feb 22, 2013	DP		
<u>PALIPERIDONE; INVEGA</u>					
021999 001				NCE	Dec 19, 2011
<u>PALIPERIDONE; INVEGA</u>					
021999 002				NCE	Dec 19, 2011
<u>PALIPERIDONE; INVEGA</u>					
021999 003				NCE	Dec 19, 2011
<u>PALIPERIDONE; INVEGA</u>					
021999 004				NCE	Dec 19, 2011
<u>PALONOSETRON HYDROCHLORIDE; ALOXI</u>					
021372 001	5202333	Apr 13, 2010		U-528	NCE
<u>PANTOPRAZOLE SODIUM; PROTONIX</u>					
020987 001	4758579	Jul 19, 2010			
	5997903	Dec 07, 2016			
<u>PANTOPRAZOLE SODIUM; PROTONIX</u>					
020987 002	4758579	Jul 19, 2010			
	5997903	Dec 07, 2016			

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>PANTOPRAZOLE SODIUM; PROTONIX IV</u>					
020988 001	4758579	Jul 19, 2010		I-444	Dec 06, 2007
	6780881	Nov 17, 2021	DP		
<u>PARICALCITOL; ZEMPLAR</u>					
020819 001	5246925	Apr 17, 2012	U-314	M-31	Mar 31, 2007
	5246925*PED	Oct 17, 2012		PED	Oct 01, 2007
	5587497	Dec 24, 2013			
	5587497*PED	Jun 24, 2014			
	6136799	Apr 08, 2018			
	6136799*PED	Oct 08, 2019			
	6361758	Apr 08, 2018	DP		
	6361758*PED	Oct 08, 2018			
<u>PARICALCITOL; ZEMPLAR</u>					
020819 002	5246925	Apr 17, 2012	U-314	M-31	Mar 31, 2007
	5246925*PED	Oct 17, 2012		PED	Oct 01, 2007
	5587497	Dec 24, 2013			
	5587497*PED	Jun 24, 2014			
	6136799	Apr 08, 2018			
	6136799*PED	Oct 08, 2019			
	6361758	Apr 08, 2018	DP		
	6361758*PED	Oct 08, 2018			
<u>PARICALCITOL; ZEMPLAR</u>					
021606 001	5246925	Apr 17, 2012	U-671	NDF	May 26, 2008
	5246925*PED	Oct 17, 2012			
	5587497	Dec 24, 2013	DS		
	5587497*PED	Jun 24, 2014			
<u>PARICALCITOL; ZEMPLAR</u>					
021606 002	5246925	Apr 17, 2012	U-671	NDF	May 26, 2008
	5246925*PED	Oct 17, 2012			
	5587497	Dec 24, 2013	DS		
	5587497*PED	Jun 24, 2014			
<u>PARICALCITOL; ZEMPLAR</u>					
021606 003	5246925	Apr 17, 2012	U-671	NDF	May 26, 2008
	5246925*PED	Oct 17, 2012			
	5587497	Dec 24, 2013	DS		
	5587497*PED	Jun 24, 2014			
<u>PAROXETINE HYDROCHLORIDE; PAXIL</u>					
020031 001	4721723	Dec 29, 2006	U-12		
	4721723*PED	Jun 29, 2007	U-12		
	5789449	Jan 06, 2009	U-285		
	5789449*PED	Jul 06, 2009	U-285		
	5872132	May 19, 2015			
	5872132*PED	Nov 19, 2015			
	5900423	May 19, 2015			
	5900423*PED	Nov 19, 2015			
	6113944	Dec 14, 2014			
	6113944*PED	Jun 14, 2015			
	6121291	Mar 17, 2017	U-431		
	6121291	Mar 17, 2017	U-286		
	6121291*PED	Sep 17, 2017	U-431		
	6121291*PED	Sep 17, 2017	U-286		
	6133289	May 19, 2015	U-358		
	6133289*PED	Nov 19, 2015	U-358		

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>PAROXETINE HYDROCHLORIDE; PAXIL</u>					
020031 002	4721723	Dec 29, 2006		U-12	
	4721723*PED	Jun 29, 2007		U-12	
	5789449	Jan 06, 2009		U-285	
	5789449*PED	Jul 06, 2009		U-285	
	5872132	May 19, 2015			
	5872132*PED	Nov 19, 2015			
	5900423	May 19, 2015			
	5900423*PED	Nov 19, 2015			
	6113944	Dec 14, 2014			
	6113944*PED	Jun 14, 2015			
	6121291	Mar 17, 2017		U-431	
	6121291	Mar 17, 2017		U-286	
	6121291*PED	Sep 17, 2017		U-431	
	6121291*PED	Sep 17, 2017		U-286	
	6133289	May 19, 2015		U-358	
	6133289*PED	Nov 19, 2015		U-358	
<u>PAROXETINE HYDROCHLORIDE; PAXIL</u>					
020031 003	4721723	Dec 29, 2006		U-12	
	4721723*PED	Jun 29, 2007		U-12	
	5789449	Jan 06, 2009		U-285	
	5789449*PED	Jul 06, 2009		U-285	
	5872132	May 19, 2015			
	5872132*PED	Nov 19, 2015			
	5900423	May 19, 2015			
	5900423*PED	Nov 19, 2015			
	6113944	Dec 14, 2014			
	6113944*PED	Jun 14, 2015			
	6121291	Mar 17, 2017		U-431	
	6121291	Mar 17, 2017		U-286	
	6121291*PED	Sep 17, 2017		U-286	
	6121291*PED	Sep 17, 2017		U-431	
	6133289	May 19, 2015		U-358	
	6133289*PED	Nov 19, 2015		U-358	
<u>PAROXETINE HYDROCHLORIDE; PAXIL</u>					
020031 004	4721723	Dec 29, 2006		U-12	
	4721723*PED	Jun 29, 2007		U-12	
	5789449	Jan 06, 2009		U-285	
	5789449*PED	Jul 06, 2009		U-285	
	5872132	May 19, 2015			
	5872132*PED	Nov 19, 2015			
	5900423	May 19, 2015			
	5900423*PED	Nov 19, 2015			
	6113944	Dec 14, 2014			
	6113944*PED	Jun 14, 2015			
	6121291	Mar 17, 2017		U-286	
	6121291	Mar 17, 2017		U-431	
	6121291*PED	Sep 17, 2017		U-431	
	6121291*PED	Sep 17, 2017		U-286	
	6133289	May 19, 2015		U-358	
	6133289*PED	Nov 19, 2015		U-358	
<u>PAROXETINE HYDROCHLORIDE; PAXIL</u>					
020031 005	4721723	Dec 29, 2006		U-12	
	4721723*PED	Jun 29, 2007		U-12	
	5789449	Jan 06, 2009		U-285	
	5789449*PED	Jul 06, 2009		U-285	
	5872132	May 19, 2015			
	5872132*PED	Nov 19, 2015			
	5900423	May 19, 2015			
	5900423*PED	Nov 19, 2015			
	6113944	Dec 14, 2014			
	6113944*PED	Jun 14, 2015			
	6121291	Mar 17, 2017		U-431	
	6121291	Mar 17, 2017		U-286	
	6121291*PED	Sep 17, 2017		U-431	
	6121291*PED	Sep 17, 2017		U-286	
	6133289	May 19, 2015		U-358	
	6133289*PED	Nov 19, 2015		U-358	

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>PAROXETINE HYDROCHLORIDE; PAXIL</u>					
020710 001	4721723	Dec 29, 2006			
	4721723*PED	Jun 29, 2007			
	5789449	Jan 06, 2009		U-285	
	5789449*PED	Jul 06, 2009		U-285	
	5811436	Sep 22, 2015			
	5811436*PED	Mar 22, 2016			
	5872132	May 19, 2015			
	5872132*PED	Nov 19, 2015			
	5900423	May 19, 2015			
	5900423*PED	Nov 19, 2015			
	6121291	Mar 17, 2017		U-431	
	6121291	Mar 17, 2017		U-286	
	6121291*PED	Sep 17, 2017		U-286	
	6121291*PED	Sep 17, 2017		U-431	
	6133289	May 19, 2015		U-358	
	6133289*PED	Nov 19, 2015		U-358	
<u>PAROXETINE HYDROCHLORIDE; PAXIL</u>					
020885 001	4721723	Dec 29, 2006		U-12	
	4721723*PED	Jun 29, 2007		U-12	
	5789449	Jan 06, 2009		U-285	
	5789449*PED	Jul 06, 2009		U-285	
	5872132	May 19, 2015			
	5872132*PED	Nov 19, 2015			
	5900423	May 19, 2015			
	5900423*PED	Nov 19, 2015			
	6063927	Apr 23, 2019			
	6063927*PED	Oct 23, 2019			
	6080759	May 19, 2015			
	6080759*PED	Nov 19, 2015			
	6121291	Mar 17, 2017		U-431	
	6121291	Mar 17, 2017		U-286	
	6121291*PED	Sep 17, 2017		U-431	
	6121291*PED	Sep 17, 2017		U-286	
	6133289	May 19, 2015		U-358	
	6133289*PED	Nov 19, 2015		U-358	
	6172233	Jan 15, 2018			
	6172233*PED	Jul 15, 2018			
<u>PAROXETINE HYDROCHLORIDE; PAXIL</u>					
020885 002	4721723	Dec 29, 2006		U-12	
	4721723*PED	Jun 29, 2007		U-12	
	5789449	Jan 06, 2009		U-285	
	5789449*PED	Jul 06, 2009		U-285	
	5872132	May 19, 2015			
	5872132*PED	Nov 19, 2015			
	5900423	May 19, 2015			
	5900423*PED	Nov 19, 2015			
	6063927	Apr 23, 2019			
	6063927*PED	Oct 23, 2019			
	6080759	May 19, 2015			
	6080759*PED	Nov 19, 2015			
	6121291	Mar 17, 2017		U-431	
	6121291	Mar 17, 2017		U-286	
	6121291*PED	Sep 17, 2017		U-286	
	6121291*PED	Sep 17, 2017		U-431	
	6133289	May 19, 2015		U-358	
	6133289*PED	Nov 19, 2015		U-358	
	6172233	Jan 15, 2018			
	6172233*PED	Jul 15, 2018			

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>PAROXETINE HYDROCHLORIDE; PAXIL</u>					
020885 003	4721723	Dec 29, 2006		U-12	
	4721723*PED	Jun 29, 2007		U-12	
	5789449	Jan 06, 2009		U-285	
	5789449*PED	Jul 06, 2009		U-285	
	5872132	May 19, 2015			
	5872132*PED	Nov 19, 2015			
	5900423	May 19, 2015			
	5900423*PED	Nov 19, 2015			
	6063927	Apr 23, 2019			
	6063927*PED	Oct 23, 2019			
	6080759	May 19, 2015			
	6080759*PED	Nov 19, 2015			
	6121291	Mar 17, 2017		U-431	
	6121291	Mar 17, 2017		U-286	
	6121291*PED	Sep 17, 2017		U-286	
	6121291*PED	Sep 17, 2017		U-431	
	6133289	May 19, 2015		U-358	
	6133289*PED	Nov 19, 2015		U-358	
	6172233	Jan 15, 2018			
	6172233*PED	Jul 15, 2018			
<u>PAROXETINE HYDROCHLORIDE; PAXIL</u>					
020885 004	4721723	Dec 29, 2006		U-12	
	4721723*PED	Jun 29, 2007		U-12	
	5789449	Jan 06, 2009		U-285	
	5789449*PED	Jul 06, 2009		U-285	
	5872132	May 19, 2015			
	5872132*PED	Nov 19, 2015			
	5900423	May 19, 2015			
	5900423*PED	Nov 19, 2015			
	6062927	Apr 23, 2019			
	6063927*PED	Oct 23, 2019			
	6080759	May 19, 2015			
	6080759*PED	Nov 19, 2015			
	6121291	Mar 17, 2017		U-431	
	6121291	Mar 17, 2017		U-286	
	6121291*PED	Sep 17, 2017		U-286	
	6121291*PED	Sep 17, 2017		U-431	
	6133289	May 19, 2015		U-358	
	6133289*PED	Nov 19, 2015		U-358	
	6172233	Jan 15, 2018			
	6172233*PED	Jul 15, 2018			
<u>PAROXETINE HYDROCHLORIDE; PAXIL CR</u>					
020936 001	4721723	Dec 29, 2006		D-91	Jan 27, 2007
	4721723*PED	Jun 29, 2007		I-405	Aug 28, 2006
	4839177*PED	Dec 13, 2006			
	5422123	Jun 06, 2012			
	5422123*PED	Dec 06, 2012			
	5789449	Jan 06, 2009		U-286	
	5789449*PED	Jul 06, 2009		U-286	
	5872132	May 19, 2015			
	5872132*PED	Nov 19, 2015			
	5900423	May 19, 2015			
	5900423*PED	Nov 19, 2015			
	6121291	Mar 17, 2017		U-286	
	6121291*PED	Sep 17, 2017		U-286	
	6133289	May 19, 2015		U-286	
	6133289*PED	Nov 19, 2015		U-286	
	6548084	Jul 19, 2016			
	6548084*PED	Jan 19, 2017			

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>PAROXETINE HYDROCHLORIDE; PAXIL CR</u>					
020936 002	4721723	Dec 29, 2006		D-91	Jan 27, 2007
	4721723*PED	Jun 29, 2007		I-405	Aug 28, 2006
	4839177*PED	Dec 13, 2006			
	5422123	Jun 06, 2012			
	5422123*PED	Dec 06, 2012			
	5789449	Jan 06, 2009	U-286		
	5789449*PED	Jul 06, 2009	U-286		
	5872132	May 19, 2015			
	5872132*PED	Nov 19, 2015			
	5900423	May 19, 2015			
	5900423*PED	Nov 19, 2015			
	6121291	Mar 17, 2017	U-286		
	6121291*PED	Sep 17, 2017	U-286		
	6133289	May 19, 2015	U-286		
	6133289*PED	Nov 19, 2015	U-286		
	6548084	Jul 19, 2016			
	6548084*PED	Jan 19, 2017			
<u>PAROXETINE HYDROCHLORIDE; PAXIL CR</u>					
020936 003	4721723	Dec 29, 2006		D-91	Jan 27, 2007
	4721723*PED	Jun 29, 2007		I-405	Aug 28, 2006
	4839177*PED	Dec 13, 2006			
	5422123	Jun 06, 2012			
	5422123*PED	Dec 06, 2012			
	5789449	Jan 06, 2009	U-286		
	5789449*PED	Jul 06, 2009	U-286		
	5872132	May 19, 2015			
	5872132*PED	Nov 19, 2015			
	5900423	May 19, 2015			
	5900423*PED	Nov 19, 2015			
	6121291	Mar 17, 2017	U-286		
	6121291*PED	Sep 17, 2017	U-286		
	6133289	May 19, 2015	U-286		
	6133289*PED	Nov 19, 2015	U-286		
	6548084	Jul 19, 2016			
	6548084*PED	Jan 19, 2017			
<u>PAROXETINE MESYLATE; PEXEVA</u>					
021299 001	5874447	Jun 10, 2017	U-286		
	5874447	Jun 10, 2017	U-518		
	5874447	Jun 10, 2017	U-46		
	6703408	Oct 21, 2022	DP		
<u>PAROXETINE MESYLATE; PEXEVA</u>					
021299 002	5874447	Jun 10, 2017	U-286		
	5874447	Jun 10, 2017	U-518		
	5874447	Jun 10, 2017	U-46		
	6703408	Oct 21, 2022	DP		
<u>PAROXETINE MESYLATE; PEXEVA</u>					
021299 003	5874447	Jun 10, 2017	U-286		
	5874447	Jun 10, 2017	U-518		
	5874447	Jun 10, 2017	U-46		
	6703408	Oct 21, 2022	DP		
<u>PAROXETINE MESYLATE; PEXEVA</u>					
021299 004	5874447	Jun 10, 2017	U-518		
	5874447	Jun 10, 2017	U-286		
	5874447	Jun 10, 2017	U-46		
	6703408	Oct 21, 2022	DP		
<u>PEGAPTANIB SODIUM; MACUGEN</u>					
021756 001	5919455	Oct 27, 2013	DS	NCE	Dec 17, 2009
	5932462	Aug 03, 2016	DS		
	6011020	Jan 04, 2017	DS		
	6051698	Oct 17, 2012	DS		
	6113906	Oct 27, 2013	DS		
	6147204	Jun 11, 2010	DS		
	6426335	Jun 11, 2010	U-622		

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>PEGVISOMANT; SOMAVERT</u>					
021106 001	5350836	Sep 27, 2011	U-507	NCE	Mar 25, 2008
	5681809	Sep 27, 2011	U-507	ODE	Mar 25, 2010
	5849535	Sep 21, 2015			
	5958879	Sep 27, 2011	U-507		
	6057292	Sep 21, 2015	U-507		
	6583115	Sep 27, 2011	U-507		
<u>PEGVISOMANT; SOMAVERT</u>					
021106 002	5350836	Sep 27, 2011	U-507	NCE	Mar 25, 2008
	5681809	Sep 27, 2011	U-507	ODE	Mar 25, 2010
	5849535	Sep 21, 2015			
	5958879	Sep 27, 2011	U-507		
	6057292	Sep 21, 2015	U-507		
	6583115	Sep 27, 2011	U-507		
<u>PEGVISOMANT; SOMAVERT</u>					
021106 003	5350836	Sep 27, 2011	U-507	NCE	Mar 25, 2008
	5681809	Sep 27, 2011	U-507	ODE	Mar 25, 2010
	5849535	Sep 21, 2015			
	5958879	Sep 27, 2011	U-507		
	6057292	Sep 21, 2015	U-507		
	6583115	Sep 27, 2011	U-507		
<u>PEMETREXED DISODIUM; ALIMTA</u>					
021462 001	5217974	Mar 29, 2011		NCE	Feb 04, 2009
	5344932	Sep 06, 2011	DS DP	ODE	Feb 04, 2011
<u>PEMIROLAST POTASSIUM; ALAMAST</u>					
021079 001	5034230	Dec 23, 2008		U-184	
	5034230*PED	Jun 23, 2009		U-184	
<u>PENCICLOVIR SODIUM; DENAVIR</u>					
020629 001	5075445	Sep 24, 2010		NPP	Oct 10, 2006
	5840763	Sep 01, 2015			
	5866581	Sep 04, 2014	U-501		
	5916893	Sep 01, 2015	U-501		
	6124304	Sep 04, 2014	U-501		
	6469015	Oct 22, 2019	U-501		
	6573378	Sep 24, 2010			
	6579981	Jun 17, 2020	U-501		
<u>PENTETATE CALCIUM TRISODIUM; PENTETATE CALCIUM TRISODIUM</u>					
021749 001				NCE	Aug 11, 2009
				ODE	Aug 11, 2011
<u>PENTETATE ZINC TRISODIUM; PENTETATE ZINC TRISODIUM</u>					
021751 001				NCE	Aug 11, 2009
				ODE	Aug 11, 2011
<u>PENTOSAN POLYSULFATE SODIUM; ELMIRON</u>					
020193 001	5180715	Jan 19, 2010		U-159	
<u>PERFLUOROPOLYMETHYLISOPROPYL ETHER; POLYTETRAFLUOROETHYLENE; SKIN EXPOSURE REDUCTION PASTE AGAINST CHEMICAL WARFARE AGENTS</u>					
021084 001	5607979	May 30, 2015			
<u>PERFLUTREN; DEFINITY</u>					
021064 001	5547656	Apr 05, 2011		NCE	Jul 31, 2006
	5769080	Jul 20, 2010			
	6033645	Jun 19, 2016			
	6146657	Dec 22, 2009	DS	U-665	
	6528039	Apr 05, 2011	DS		
	6773696	Apr 05, 2011	DS		
<u>PERGOLIDE MESYLATE; PERMAX</u>					
019385 001	4797405	Oct 26, 2007			
	5114948	Oct 19, 2009			
<u>PERGOLIDE MESYLATE; PERMAX</u>					
019385 002	4797405	Oct 26, 2007			
	5114948	Oct 19, 2009			
<u>PERGOLIDE MESYLATE; PERMAX</u>					
019385 003	4797405	Oct 26, 2007			
	5114948	Oct 19, 2009			

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>PERINDOPRIL ERBUMINE; ACEON</u>					
020184 001	4508729	Aug 21, 2006		I-468	Aug 23, 2008
	5162362	Nov 10, 2009	DS DP U-531		
<u>PERINDOPRIL ERBUMINE; ACEON</u>					
020184 002	4508729	Aug 21, 2006		I-468	Aug 23, 2008
	5162362	Nov 10, 2009	DS DP U-531		
<u>PERINDOPRIL ERBUMINE; ACEON</u>					
020184 003	4508729	Aug 21, 2006		I-468	Aug 23, 2008
	5162362	Nov 10, 2009	DS DP U-531		
<u>PIMECROLIMUS; ELIDEL</u>					
021302 001	5912238	Jun 15, 2016		NCE	Dec 13, 2006
	5912238*PED	Dec 15, 2016		PED	Jun 13, 2007
	6352998	Oct 26, 2015			
	6352998*PED	Apr 26, 2016			
	6423722	Jun 26, 2018			
	6423722*PED	Dec 26, 2018			
<u>PIOGLITAZONE HYDROCHLORIDE; ACTOS</u>					
021073 001	4687777	Jan 17, 2011		M-35	Nov 26, 2006
	5965584	Jun 19, 2016	U-417		
	6150383	Jun 19, 2016	U-418		
	6150384	Jun 19, 2016	U-419		
	6166042	Jun 19, 2016	U-414		
	6166043	Jun 19, 2016	U-415		
	6172090	Jun 19, 2016	U-416		
	6211205	Jun 19, 2016	U-410		
	6271243	Jun 19, 2016	U-411		
	6303640	Aug 09, 2016	U-425		
	6329404	Jun 19, 2016	U-430		
<u>PIOGLITAZONE HYDROCHLORIDE; ACTOS</u>					
021073 002	4687777	Jan 17, 2011		M-35	Nov 26, 2006
	5965584	Jun 19, 2016	U-417		
	6150383	Jun 19, 2016	U-418		
	6150384	Jun 19, 2016	U-419		
	6166042	Jun 19, 2016	U-414		
	6166043	Jun 19, 2016	U-415		
	6172090	Jun 19, 2016	U-416		
	6211205	Jun 19, 2016	U-410		
	6271243	Jun 19, 2016	U-411		
	6303640	Aug 09, 2016	U-425		
	6329404	Jun 19, 2016	U-430		
<u>PIOGLITAZONE HYDROCHLORIDE; ACTOS</u>					
021073 003	4687777	Jan 17, 2011		M-35	Nov 26, 2006
	5965584	Jun 19, 2016	U-417		
	6150383	Jun 19, 2016	U-418		
	6150384	Jun 19, 2016	U-419		
	6166042	Jun 19, 2016	U-414		
	6166043	Jun 19, 2016	U-415		
	6172090	Jun 19, 2016	U-416		
	6211205	Jun 19, 2016	U-410		
	6271243	Jun 19, 2016	U-411		
	6303640	Aug 09, 2016	U-425		
	6329404	Jun 19, 2016	U-430		
<u>PODOFILOX; CONDYLOX</u>					
020529 001	5057616	Oct 15, 2008			
<u>POLYETHYLENE GLYCOL 3350; MIRALAX</u>					
022015 001				NP	Oct 06, 2009
<u>PORFIMER SODIUM; PHOTOFRIN</u>					
020451 001	4932934	Jun 12, 2007	U-128	I-391	Aug 01, 2006
	5145863	Dec 15, 2009	U-129	ODE	Aug 01, 2010
	5438071	Aug 01, 2012			
<u>POSACONAZOLE; NOXAFIL</u>					
022003 001	5661151	Aug 26, 2014	DS DP U-760	NCE	Sep 15, 2011
	5703079	Dec 30, 2014	DS DP U-760		
	6958337	Sep 25, 2020	DS DP U-760		

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE		PATENT CODES		EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE		
POTASSIUM CHLORIDE; K-DUR 10									
019439 002	4863743	Sep 05, 2006			U-99				
POTASSIUM CHLORIDE; K-DUR 20									
019439 001	4863743	Sep 05, 2006			U-99				
PRAMIPEXOLE DIHYDROCHLORIDE; MIRAPEX									
020667 001	4886812	Mar 25, 2011	DS	DP	I-517	Nov 07, 2009			
	6001861	Jan 16, 2018							U-784
	6194445	Jan 16, 2018							U-784
PRAMIPEXOLE DIHYDROCHLORIDE; MIRAPEX									
020667 002	4886812	Mar 25, 2011			I-517	Nov 07, 2009			
	6001861	Jan 16, 2018							U-784
	6194445	Jan 16, 2018							U-784
PRAMIPEXOLE DIHYDROCHLORIDE; MIRAPEX									
020667 003	4886812	Mar 25, 2011	DS	DP	I-517	Nov 07, 2007			
	6001861	Jan 16, 2018							U-784
	6194445	Jan 16, 2018							U-784
PRAMIPEXOLE DIHYDROCHLORIDE; MIRAPEX									
020667 004	4886812	Mar 25, 2011							
PRAMIPEXOLE DIHYDROCHLORIDE; MIRAPEX									
020667 005	4886812	Mar 25, 2011	DS	DP	I-517	Nov 07, 2009			
	6001861	Jan 16, 2018							U-784
	6194445	Jan 16, 2018							U-784
PRAMIPEXOLE DIHYDROCHLORIDE; MIRAPEX									
020667 006	4886812	Mar 25, 2011	DS	DP	I-517	Nov 07, 2009			
	6001861	Jan 16, 2018							U-784
	6194445	Jan 16, 2018							U-784
PRAMLINTIDE ACETATE; SYMLIN									
021332 001	5175145	Dec 29, 2009	DS	DP	NCE	Mar 16, 2010			
	5686411	Nov 11, 2014							U-637
	5814600	Sep 29, 2015	U-638						
	5998367	Mar 08, 2011	U-639						
	6114304	Sep 05, 2017	DP	U-640					
	6410511	Jan 09, 2018							
	6608029	Sep 07, 2013		U-641					
	6610824	Mar 08, 2011	DS						
PRAVASTATIN SODIUM; PRAVACHOL									
019898 002	5030447	Jul 09, 2008							
	5030447*PED	Jan 09, 2009							
	5180589	Jul 09, 2008							
	5180589*PED	Jan 09, 2009							
	5622985	Apr 22, 2014							U-335
	5622985*PED	Oct 22, 2014							U-335
PRAVASTATIN SODIUM; PRAVACHOL									
019898 003	5030447	Jul 09, 2008							
	5030447*PED	Jan 09, 2009							
	5180589	Jul 09, 2008							
	5180589*PED	Jan 09, 2009							
	5622985	Apr 22, 2014							U-335
	5622985*PED	Oct 22, 2014							U-335
PRAVASTATIN SODIUM; PRAVACHOL									
019898 004	5030447	Jul 09, 2008							
	5030447*PED	Jan 09, 2009							
	5180589	Jul 09, 2008							
	5180589*PED	Jan 09, 2009							
	5622985	Apr 22, 2014							U-335
	5622985*PED	Oct 22, 2014							U-335
PRAVASTATIN SODIUM; PRAVACHOL									
019898 008	5030447	Jul 09, 2008							
	5030447*PED	Jan 09, 2009							
	5180589	Jul 09, 2008							
	5180589*PED	Jan 09, 2009							
	5622985	Apr 22, 2014							U-335
	5622985*PED	Oct 22, 2014							U-335

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>PRAVASTATIN SODIUM; PRAVASTATIN SODIUM</u>					
076056 001				PC	Oct 21, 2006
<u>PRAVASTATIN SODIUM; PRAVASTATIN SODIUM</u>					
076056 002				PC	Oct 21, 2006
<u>PRAVASTATIN SODIUM; PRAVASTATIN SODIUM</u>					
076056 003				PC	Oct 21, 2006
<u>PREDNISOLONE SODIUM PHOSPHATE; ORAPRED ODT</u>					
021959 001	5178878	Jan 12, 2010	DP		
<u>PREDNISOLONE SODIUM PHOSPHATE; ORAPRED ODT</u>					
021959 002	5178878	Jan 12, 2010	DP		
<u>PREDNISOLONE SODIUM PHOSPHATE; ORAPRED ODT</u>					
021959 003	5178878	Jan 12, 2010	DP		
<u>PREGABALIN; LYRICA</u>					
021446 001	5563175 6001876 6197819	Oct 08, 2013 Jul 16, 2017 Mar 06, 2018	DS DP	U-661 U-55	NCE Dec 30, 2009
<u>PREGABALIN; LYRICA</u>					
021446 002	5563175 6001876 6197819	Oct 08, 2013 Jul 16, 2017 Mar 06, 2018	DS DP	U-661 U-55	NCE Dec 30, 2009
<u>PREGABALIN; LYRICA</u>					
021446 003	5563175 6001876 6197819	Oct 08, 2013 Jul 16, 2017 Mar 06, 2018	DS DP	U-661 U-55	NCE Dec 30, 2009
<u>PREGABALIN; LYRICA</u>					
021446 004	5563175 6001876 6197819	Oct 08, 2013 Jul 16, 2017 Mar 06, 2018	DS DP	U-661 U-55	NCE Dec 30, 2009
<u>PREGABALIN; LYRICA</u>					
021446 005	5563175 6001876 6197819	Oct 08, 2013 Jul 16, 2017 Mar 06, 2018	DS DP	U-661 U-55	NCE Dec 30, 2009
<u>PREGABALIN; LYRICA</u>					
021446 006	5563175 6001876 6197819	Oct 08, 2013 Jul 16, 2017 Mar 06, 2018	DS DP	U-661 U-55	NCE Dec 30, 2009
<u>PREGABALIN; LYRICA</u>					
021446 007	5563175 6001876 6197819	Oct 08, 2013 Jul 16, 2017 Mar 06, 2018	DS DP	U-661 U-55	NCE Dec 30, 2009
<u>PREGABALIN; LYRICA</u>					
021446 008	5563175 6001876 6197819	Oct 08, 2013 Jul 16, 2017 Mar 06, 2018	DS DP	U-661 U-55	NCE Dec 30, 2009
<u>PROCAINAMIDE HYDROCHLORIDE; PROCANBID</u>					
020545 001	5656296	Aug 12, 2014			
<u>PROCAINAMIDE HYDROCHLORIDE; PROCANBID</u>					
020545 002	5656296	Aug 12, 2014			
<u>PROGESTERONE; CRINONE</u>					
020701 001	5543150	Sep 15, 2013		U-209	
<u>PROGESTERONE; CRINONE</u>					
020701 002	5543150	Sep 15, 2013		U-209	
<u>PROPAFENONE HYDROCHLORIDE; RYTHMOL SR</u>					
021416 001	5681588	Oct 28, 2014		NDF	Sep 04, 2006
<u>PROPAFENONE HYDROCHLORIDE; RYTHMOL SR</u>					
021416 002	5681588	Oct 28, 2014		NDF	Sep 04, 2006
<u>PROPAFENONE HYDROCHLORIDE; RYTHMOL SR</u>					
021416 003	5681588	Oct 28, 2014		NDF	Sep 04, 2006

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>PROPOFOL; DIPRIVAN</u>					
019627 002	5714520	Mar 22, 2015			
	5714520*PED	Sep 22, 2015			
	5731355	Mar 22, 2015		U-217	
	5731355*PED	Sep 22, 2015		U-217	
	5731356	Mar 22, 2015		U-218	
	5731356*PED	Sep 22, 2015		U-218	
	5908869	Mar 22, 2015		U-270	
	5908869*PED	Sep 22, 2015		U-270	
<u>PROPRANOLOL HYDROCHLORIDE; INNOPRAN XL</u>					
021438 001	6500454	Dec 31, 2022			
<u>PROPRANOLOL HYDROCHLORIDE; INNOPRAN XL</u>					
021438 002	6500454	Dec 31, 2022			
<u>QUETIAPINE FUMARATE; SEROQUEL</u>					
020639 001	4879288	Sep 26, 2011	DS DP	U-550	
				I-503	Oct 20, 2009
				I-418	Jan 12, 2007
				I-419	Jan 12, 2007
<u>QUETIAPINE FUMARATE; SEROQUEL</u>					
020639 002	4879288	Sep 26, 2011	DS DP	U-550	
				I-503	Oct 20, 2009
				I-418	Jan 12, 2007
				I-419	Jan 12, 2007
<u>QUETIAPINE FUMARATE; SEROQUEL</u>					
020639 003	4879288	Sep 26, 2011	DS DP	U-550	
				I-503	Oct 20, 2009
				I-418	Jan 12, 2007
				I-419	Jan 12, 2007
<u>QUETIAPINE FUMARATE; SEROQUEL</u>					
020639 004	4879288	Sep 26, 2011	DS DP	U-550	
				I-503	Oct 20, 2009
				I-418	Jan 12, 2007
				I-419	Jan 12, 2007
<u>QUETIAPINE FUMARATE; SEROQUEL</u>					
020639 005	4879288	Sep 26, 2011	DS DP	U-550	
				I-503	Oct 20, 2009
				I-418	Jan 12, 2007
				I-419	Jan 12, 2007
<u>QUETIAPINE FUMARATE; SEROQUEL</u>					
020639 006	4879288	Sep 26, 2011	DS DP	U-550	
				I-503	Oct 20, 2009
				I-419	Jan 12, 2007
				I-418	Jan 12, 2007
<u>QUETIAPINE FUMARATE; SEROQUEL</u>					
020639 007	4879288	Sep 26, 2011	DS DP	U-550	
				I-503	Oct 20, 2009
				I-419	Jan 12, 2007
				I-418	Jan 12, 2007
<u>QUINAPRIL HYDROCHLORIDE; ACCUPRIL</u>					
019885 001	4743450	Feb 24, 2007			
	4743450*PED	Aug 24, 2007			
	5684016	Nov 04, 2014		U-210	
	5684016*PED	May 04, 2015		U-210	
<u>QUINAPRIL HYDROCHLORIDE; ACCUPRIL</u>					
019885 002	4743450	Feb 24, 2007			
	4743450*PED	Aug 24, 2007			
	5684016	Nov 04, 2014		U-210	
	5684016*PED	May 04, 2015		U-210	
<u>QUINAPRIL HYDROCHLORIDE; ACCUPRIL</u>					
019885 003	4743450	Feb 24, 2007			
	4743450*PED	Aug 24, 2007			
	5684016	Nov 04, 2014		U-210	
	5684016*PED	May 04, 2015		U-210	

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>QUINAPRIL HYDROCHLORIDE; ACCUPRIL</u>					
019885 004	4743450	Feb 24, 2007			
	4743450*PED	Aug 24, 2007			
	5684016	Nov 04, 2014		U-210	
	5684016*PED	May 04, 2015		U-210	
<u>QUININE SULFATE; QUININE SULFATE</u>					
021799 001				ODE	Aug 12, 2012
<u>RABEPRAZOLE SODIUM; ACIPHEX</u>					
020973 001	5035899	Apr 04, 2009		U-385	
	5045552	May 08, 2013		U-385	
<u>RABEPRAZOLE SODIUM; ACIPHEX</u>					
020973 002	5035899	Apr 04, 2009		U-385	
	5045552	May 08, 2013		U-385	
<u>RALOXIFENE HYDROCHLORIDE; EVISTA</u>					
020815 001	5393763	Jul 28, 2012		U-114	
	5457117	Jul 28, 2012		U-114	
	5478847	Mar 02, 2014		U-114	
	5811120	Mar 02, 2014			
	5972383	Mar 02, 2014		U-287	
	6458811	Mar 10, 2017			
	6797719	Mar 10, 2017			
	6894064	Mar 10, 2017	DP	U-657	
	6906086	Jul 28, 2012	DP	U-657	
	6906086	Jul 28, 2012		U-662	
	RE38968	Jul 28, 2012		U-657	
	RE38968	Jul 28, 2012		U-662	
	RE39049	Jul 28, 2012		U-657	
	RE39049	Jul 28, 2012		U-662	
	RE39050	Mar 02, 2014		U-657	
	RE39050	Mar 02, 2014		U-662	
<u>RAMELTEON; ROZEREM</u>					
021782 001	6034239	Mar 06, 2017	DS DP	U-674 NCE	Jul 22, 2010
<u>RAMIPRIL; ALTACE</u>					
019901 001	5061722	Oct 19, 2008			
	5403856	Apr 04, 2012		U-71	
<u>RAMIPRIL; ALTACE</u>					
019901 002	5061722	Oct 19, 2008			
	5403856	Apr 04, 2012		U-71	
<u>RAMIPRIL; ALTACE</u>					
019901 003	5061722	Oct 19, 2008			
	5403856	Apr 04, 2012		U-71	
<u>RAMIPRIL; ALTACE</u>					
019901 004	5061722	Oct 19, 2008			
	5403856	Apr 04, 2012		U-71	
<u>RANITIDINE BISMUTH CITRATE; TRITEC</u>					
020559 001	5008256	Jul 17, 2009			
	5256684	Oct 26, 2010		U-199	
	5403830	Apr 04, 2012		U-200	
	5407688	Apr 04, 2012		U-201	
	5456925	Oct 10, 2012			
	5601848	Feb 11, 2014		U-202	
	5629297	May 13, 2014		U-186	
<u>RANITIDINE HYDROCHLORIDE; ZANTAC</u>					
019675 001	5068249	Nov 26, 2008			
	5068249*PED	May 26, 2009			
<u>RANITIDINE HYDROCHLORIDE; ZANTAC 150</u>					
020095 001	5028432	Feb 22, 2010			
	5028432*PED	Aug 22, 2010			
<u>RANITIDINE HYDROCHLORIDE; ZANTAC 150</u>					
020251 001	5102665	Jun 23, 2009			
	5102665*PED	Dec 23, 2009			

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>RANITIDINE HYDROCHLORIDE; ZANTAC 150</u>					
020251 002	5102665	Jun 23, 2009			
	5102665*PED	Dec 23, 2009			
<u>RANITIDINE HYDROCHLORIDE; ZANTAC 150</u>					
021698 001				NP	Aug 31, 2007
<u>RANITIDINE HYDROCHLORIDE; ZANTAC 300</u>					
020095 002	5028432	Feb 22, 2010			
	5028432*PED	Aug 22, 2010			
<u>RANOLAZINE; RANEXA</u>					
021526 002	4567264	May 18, 2007	DS	NCE	Jan 27, 2011
	6303607	May 27, 2019		U-705	
	6369062	May 27, 2019	DP		
	6479496	May 27, 2019		U-705	
	6503911	May 27, 2019	DP		
	6525057	May 27, 2019		U-705	
	6562826	May 27, 2019		U-705	
	6617328	May 27, 2019	DP		
	6620814	May 27, 2019		U-705	
	6852724	May 27, 2019		U-705	
	6864258	May 27, 2019		U-705	
<u>RAPACURONIUM BROMIDE; RAPLON</u>					
020984 001	5418226	Apr 14, 2013			
<u>RAPACURONIUM BROMIDE; RAPLON</u>					
020984 002	5418226	Apr 14, 2013			
<u>RASAGILINE MESYLATE; AZILECT</u>					
021641 001	5387612	Feb 07, 2012		U-219	May 16, 2011
	5453446	Feb 07, 2012		U-219	
	5457133	Feb 07, 2012	DS DP		
	5532415	Jul 02, 2013	DS		
	5786390	Feb 07, 2012		DP	
	6126968	Sep 18, 2016		DP	
<u>RASAGILINE MESYLATE; AZILECT</u>					
021641 002	5387612	Feb 07, 2012		U-219	May 16, 2011
	5453446	Feb 07, 2012		U-219	
	5457133	Feb 07, 2012	DS DP		
	5532415	Jul 02, 2013	DS		
	5786390	Feb 07, 2012		DP	
	6126968	Sep 18, 2016		DP	
<u>REMIFENTANIL HYDROCHLORIDE; ULTIVA</u>					
020630 001	5019583	Feb 15, 2009			
	5019583*PED	Aug 15, 2009			
	5466700	Aug 30, 2013		U-156	
	5466700*PED	Mar 01, 2014		U-156	
<u>REMIFENTANIL HYDROCHLORIDE; ULTIVA</u>					
020630 002	5019583	Feb 15, 2009			
	5019583*PED	Aug 15, 2009			
	5466700	Aug 30, 2013		U-156	
	5466700*PED	Mar 01, 2014			
<u>REMIFENTANIL HYDROCHLORIDE; ULTIVA</u>					
020630 003	5019583	Feb 15, 2009			
	5019583*PED	Aug 15, 2009			
	5466700	Aug 30, 2013		U-156	
	5466700*PED	Mar 01, 2014		U-156	
<u>REPAGLINIDE; PRANDIN</u>					
020741 001	5312924	Sep 05, 2006		U-214	
	6143769	Sep 05, 2006			
	6677358	Jun 12, 2018	DS DP	U-546	
	RE37035	Mar 14, 2009			
<u>REPAGLINIDE; PRANDIN</u>					
020741 002	5312924	Sep 05, 2006		U-214	
	6143769	Sep 05, 2006			
	6677358	Jun 12, 2018	DS DP	U-546	
	RE37035	Mar 14, 2009			

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE		PATENT CODES			EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE	
REPAGLINIDE; PRANDIN									
020741 003	5312924	Sep 05, 2006				U-214			
	6143769	Sep 05, 2006							
	6677358	Jun 12, 2018		DS	DP	U-546			
	RE37035	Mar 14, 2009							
RIBAVIRIN; COPEGUS									
021511 001							I-447	Feb 25, 2008	
RIBAVIRIN; COPEGUS									
021511 002							I-447	Feb 25, 2008	
RIBAVIRIN; REBETOL									
020903 001	5767097	Jan 23, 2016				U-235	ODE	Jul 29, 2010	
	5767097*PED	Jul 23, 2016				U-235	PED	Jan 29, 2011	
	5914128	Dec 22, 2017							
	5914128*PED	Jun 22, 2018							
	6051252	Dec 22, 2017							
	6051252*PED	Jun 22, 2018							
	6063772	Jan 23, 2016				U-375			
	6063772*PED	Jul 23, 2016				U-375			
	6172046	Sep 21, 2017				U-377			
	6172046*PED	Mar 21, 2018				U-377			
	6177074	Nov 01, 2016				U-454			
	6177074*PED	May 01, 2017				U-454			
	6335032	Dec 22, 2017							
	6335032*PED	Jun 22, 2018							
	6337090	Dec 22, 2017							
	6337090*PED	Jun 22, 2018							
	6461605	Nov 01, 2016				U-478			
	6461605*PED	May 01, 2017				U-478			
	6472373	Sep 21, 2017				U-479			
	6472373*PED	Mar 21, 2018				U-479			
	6524570	Nov 01, 2016				U-499			
	6524570*PED	May 01, 2017				U-499			
RIBAVIRIN; REBETOL									
020903 002	5767097	Jan 23, 2016				U-235	ODE	Jul 29, 2010	
	5767097*PED	Jul 23, 2016				U-235	PED	Jan 29, 2011	
	5914128	Dec 22, 2017							
	5914128*PED	Jun 22, 2018							
	6051252	Dec 22, 2017							
	6051252*PED	Jun 22, 2018							
	6063772	Jan 23, 2016				U-375			
	6063772*PED	Jul 23, 2016				U-375			
	6172046	Sep 21, 2017				U-377			
	6172046*PED	Mar 21, 2018				U-377			
	6177074	Nov 01, 2016				U-454			
	6177074*PED	May 01, 2017				U-454			
	6335032	Dec 22, 2017							
	6335032*PED	Jun 22, 2018							
	6337090	Dec 22, 2017							
	6337090*PED	Jun 22, 2018							
	6461605	Nov 01, 2016				U-478			
	6461605*PED	May 01, 2017				U-478			
	6472373	Sep 21, 2017				U-479			
	6472373*PED	Mar 21, 2018				U-479			
	6524570	Nov 01, 2016				U-499			
	6524570*PED	May 01, 2017				U-499			
RIBAVIRIN; REBETOL									
021546 001	5767097	Jan 23, 2016				U-521	NDF	Jul 29, 2006	
	5767097*PED	Jul 23, 2016				U-521	ODE	Jul 29, 2010	
	6063772	Jan 23, 2016				U-521	PED	Jan 29, 2011	
	6063772*PED	Jul 23, 2016				U-521	PED	Jan 29, 2007	
	6172046	Sep 21, 2017				U-521			
	6172046*PED	Mar 21, 2018				U-521			
	6461605	Nov 01, 2016				U-521			
	6461605*PED	May 01, 2017				U-521			
	6472373	Sep 21, 2017				U-521			
	6472373*PED	Mar 21, 2018				U-521			
RIBAVIRIN; VIRAZOLE									
018859 001	6150337	Nov 21, 2017				U-400			

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>RIFAXIMIN; XIFAXAN</u>					
021361 001	7045620	May 22, 2024	DS	NCE	May 25, 2009
<u>RILUZOLE; RILUTEK</u>					
020599 001	5527814	Jun 18, 2013			
<u>RIMEXOLONE; VEXOL</u>					
020474 001	4686214	Jul 22, 2008	U-100		
<u>RISEDRONATE SODIUM; ACTONEL</u>					
020835 001	5583122	Dec 10, 2013	U-222	M-52	Jan 24, 2009
	6096342	Nov 22, 2011			
	6165513	Jun 10, 2018			
<u>RISEDRONATE SODIUM; ACTONEL</u>					
020835 002	5583122	Dec 10, 2013	U-222	M-52	Jan 24, 2009
	6096342	Nov 22, 2011			
	6165513	Jun 10, 2018			
<u>RISEDRONATE SODIUM; ACTONEL</u>					
020835 003	5583122	Dec 10, 2013	DS DP U-222	I-309	Aug 11, 2009
	5583122	Dec 10, 2013	DS DP U-756	M-52	Jan 24, 2009
	5994329	Jul 17, 2018			
	6015801	Jul 17, 2018			
	6096342	Nov 22, 2011	DP		
	6165513	Jun 10, 2018	DP		
	6432932	Jul 17, 2018			
	6465443	Aug 14, 2018	DP	U-595	
<u>RISPERIDONE; RISPERDAL</u>					
020272 001	4804663	Dec 29, 2007	U-90	I-509 I-413 I-412	Oct 06, 2009 Dec 04, 2006 Dec 04, 2006
<u>RISPERIDONE; RISPERDAL</u>					
020272 002	4804663	Dec 29, 2007	U-90	I-509 I-413 I-412	Oct 06, 2009 Dec 04, 2006 Dec 04, 2006
<u>RISPERIDONE; RISPERDAL</u>					
020272 003	4804663	Dec 29, 2007	U-90	I-509 I-413 I-412	Oct 06, 2009 Dec 04, 2006 Dec 04, 2006
<u>RISPERIDONE; RISPERDAL</u>					
020272 004	4804663	Dec 29, 2007	U-90	I-509 I-413 I-412	Oct 06, 2009 Dec 04, 2006 Dec 04, 2006
<u>RISPERIDONE; RISPERDAL</u>					
020272 005	4804663	Dec 29, 2007	U-90	I-413 I-412	Dec 04, 2006 Dec 04, 2006
<u>RISPERIDONE; RISPERDAL</u>					
020272 007	4804663	Dec 29, 2007	U-90	I-509 I-413 I-412	Oct 06, 2009 Dec 04, 2006 Dec 04, 2006
<u>RISPERIDONE; RISPERDAL</u>					
020272 008	4804663	Dec 29, 2007	U-90	I-509 I-413 I-412	Oct 06, 2009 Dec 04, 2006 Dec 04, 2006
<u>RISPERIDONE; RISPERDAL</u>					
020588 001	4804663	Dec 29, 2007	U-90	I-509	Oct 06, 2009
	5453425	Jul 11, 2014		I-413	Dec 04, 2006
	5616587	Jul 11, 2014		I-412	Dec 04, 2006
	RE39181	Jul 11, 2014	DP		

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE		PATENT CODES			EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE	
RISPERIDONE; RISPERDAL									
021444 001	4804663	Dec	29, 2007	DS	DP	U-516	I-509	Oct	06, 2009
	5648093	Jul	15, 2014				I-413	Dec	04, 2006
	6224905	Jun	10, 2017				I-412	Dec	04, 2006
RISPERIDONE; RISPERDAL									
021444 002	4804663	Dec	29, 2007	DS	DP	U-516	I-509	Oct	06, 2009
	5648093	Jul	15, 2014				I-413	Dec	04, 2006
	6224905	Jun	10, 2017				I-412	Dec	04, 2006
RISPERIDONE; RISPERDAL									
021444 003	4804663	Dec	29, 2007	DS	DP	U-516	I-509	Oct	06, 2006
	5648093	Jul	15, 2014				I-413	Dec	04, 2006
	6224905	Jun	10, 2017				I-412	Dec	04, 2006
RISPERIDONE; RISPERDAL									
021444 004	4804663	Dec	29, 2007	DS	DP	U-543	I-509	Oct	06, 2006
	5648093	Jul	15, 2014				I-413	Dec	04, 2006
	6224905	Jun	10, 2017				I-412	Dec	04, 2006
RISPERIDONE; RISPERDAL									
021444 005	4804663	Dec	29, 2007	DS	DP	U-543	I-509	Oct	06, 2009
	5648093	Jul	15, 2014				I-413	Dec	04, 2006
	6224905	Jun	10, 2017				I-412	Dec	04, 2006
RISPERIDONE; RISPERDAL CONSTA									
021346 001	4804663	Dec	29, 2007				NDF	Oct	29, 2006
	5688801	Nov	18, 2014						
	5770231	Nov	19, 2013						
	5792477	May	02, 2017						
	5916598	May	02, 2017						
	5965168	Nov	19, 2013						
	6110503	May	02, 2017						
	6110921	Nov	19, 2013						
	6194006	Dec	30, 2018						
	6264987	May	19, 2020						
	6368632	Nov	19, 2013						
	6379703	Dec	30, 2018						
	6379704	May	19, 2020						
	6403114	May	02, 2017						
	6534092	May	19, 2020						
	6596316	Dec	30, 2008						
RISPERIDONE; RISPERDAL CONSTA									
021346 002	4804663	Dec	29, 2007				NDF	Oct	29, 2006
	5688801	Nov	18, 2014						
	5770231	Nov	19, 2013						
	5792477	May	02, 2017						
	5916598	May	02, 2017						
	5965168	Nov	19, 2013						
	6110503	May	02, 2017						
	6110921	Nov	19, 2013						
	6194006	Dec	30, 2018						
	6264987	May	19, 2020						
	6368632	Nov	19, 2013						
	6379703	Dec	30, 2018						
	6379704	May	19, 2020						
	6403114	May	02, 2017						
	6534092	May	19, 2020						
	6596316	Dec	30, 2008						

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE		PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>RISPERIDONE; RISPERDAL CONSTA</u>						
021346 003	4804663	Dec 29, 2007			NDF	Oct 29, 2006
	5688801	Nov 18, 2014				
	5770231	Nov 19, 2013				
	5792477	May 02, 2017				
	5916598	May 02, 2017				
	5965168	Nov 19, 2013				
	6110503	May 02, 2017				
	6110921	Nov 19, 2013				
	6194006	Dec 30, 2018				
	6264987	May 19, 2020				
	6368632	Nov 19, 2013		U-543		
	6379703	Dec 30, 2018	DP			
	6379704	May 19, 2020	DP			
	6403114	May 02, 2017				
	6534092	May 19, 2020	DP			
	6596316	Dec 30, 2008	DP			
<u>RITONAVIR; NORVIR</u>						
020659 001	5484801	Jan 28, 2014				
	5484801*PED	Jul 28, 2014				
	5541206	Jul 30, 2013		U-140		
	5541206*PED	Jan 30, 2014				
	5635523	Jun 03, 2014		U-190		
	5635523*PED	Dec 03, 2014				
	5648497	Jul 15, 2014				
	5648497*PED	Jan 15, 2015				
	5674882	Oct 07, 2014				
	5674882*PED	Apr 07, 2015				
	5846987	Dec 29, 2012		U-190		
	5846987*PED	Jun 29, 2013				
	5948436	Sep 13, 2013	DP			
	5948436*PED	Mar 13, 2014				
	6037157	Jun 26, 2016				
	6037157*PED	Dec 26, 2016				
	6703403	Jun 26, 2016		U-564		
	6703403*PED	Dec 26, 2016				
<u>RITONAVIR; NORVIR</u>						
020680 001	5541206	Jul 30, 2013		U-140		
	5541206*PED	Jan 30, 2014				
	5635523	Jun 03, 2014		U-190		
	5635523*PED	Dec 03, 2014				
	5648497	Jul 15, 2014				
	5648497*PED	Jan 15, 2015				
	5846987	Dec 29, 2012		U-190		
	5846987*PED	Jun 29, 2013				
	5948436	Sep 13, 2013				
	5948436*PED	Mar 13, 2014				
<u>RITONAVIR; NORVIR</u>						
020945 001	5541206	Jul 30, 2013		U-348		
	5541206*PED	Jan 30, 2014				
	5635523	Jun 03, 2014		U-347		
	5635523*PED	Dec 03, 2014				
	5648497	Jul 15, 2014				
	5648497*PED	Jan 15, 2015				
	5846987	Dec 29, 2012		U-347		
	5846987*PED	Jun 29, 2013				
	5948436	Sep 13, 2013	DP			
	5948436*PED	Mar 13, 2014				
	6232333	Nov 07, 2017				
	6232333*PED	May 07, 2018				
	6703403	Jun 26, 2016		U-564		
	6703403*PED	Dec 26, 2016				
	7141593	May 22, 2020	DP			
	7141593*PED	Nov 22, 2020				
<u>RIVASTIGMINE TARTRATE; EXELON</u>						
020823 003	4948807	Aug 14, 2012	DS	U-322		
	5602176	Feb 11, 2014		U-322		
<u>RIVASTIGMINE TARTRATE; EXELON</u>						
020823 004	4948807	Aug 14, 2012	DS	U-322		
	5602176	Feb 11, 2014		U-322		

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>RIVASTIGMINE TARTRATE; EXELON</u>					
020823 005	4948807	Aug 14, 2012	DS	U-322	
	5602176	Feb 11, 2014		U-322	
<u>RIVASTIGMINE TARTRATE; EXELON</u>					
020823 006	4948807	Aug 14, 2012	DS	U-322	
	5602176	Feb 11, 2014		U-322	
<u>RIVASTIGMINE TARTRATE; EXELON</u>					
021025 001	4948807	Aug 14, 2012	DS	U-322	
	5602176	Feb 11, 2014		U-322	
<u>RIZATRIPTAN BENZOATE; MAXALT</u>					
020864 001	5298520	Jun 29, 2012		U-240	
	5602162	Feb 11, 2014			
<u>RIZATRIPTAN BENZOATE; MAXALT</u>					
020864 002	5298520	Jun 29, 2012		U-240	
	5602162	Feb 11, 2014			
<u>RIZATRIPTAN BENZOATE; MAXALT-MLT</u>					
020865 001	5298520	Jun 29, 2012		U-240	
	5457895	Oct 01, 2013			
	5602162	Feb 11, 2014		U-240	
<u>RIZATRIPTAN BENZOATE; MAXALT-MLT</u>					
020865 002	5298520	Jun 29, 2012		U-240	
	5457895	Oct 01, 2013			
	5602162	Feb 11, 2014		U-240	
<u>ROCURONIUM BROMIDE; ZEMURON</u>					
020214 001	4894369	Apr 13, 2008			
<u>ROCURONIUM BROMIDE; ZEMURON</u>					
020214 002	4894369	Apr 13, 2008			
<u>ROCURONIUM BROMIDE; ZEMURON</u>					
020214 003	4894369	Apr 13, 2008			
<u>ROFECOXIB; VIOXX</u>					
021042 001	5474995	Jun 24, 2013	DS	DP	U-602
	5474995*PED	Dec 24, 2013			
	5691374	May 18, 2015			
	5691374*PED	Nov 18, 2015			
	6063811	May 06, 2017			U-602
	6063811*PED	Nov 06, 2017			
	6239173	Jun 24, 2013	DS	DP	U-602
	6239173*PED	Dec 24, 2013			
<u>ROFECOXIB; VIOXX</u>					
021042 002	5474995	Jun 24, 2013	DS	DP	U-602
	5474995*PED	Dec 24, 2013			
	5691374	May 18, 2015			
	5691374*PED	Nov 18, 2015			
	6063811	May 06, 2017			U-602
	6063811*PED	Nov 06, 2017			
	6239173	Jun 24, 2013	DS	DP	U-602
	6239173*PED	Dec 24, 2013			
<u>ROFECOXIB; VIOXX</u>					
021042 003	5474995	Jun 24, 2013	DS	DP	U-602
	5474995*PED	Dec 24, 2013			
	5691374	May 18, 2015			
	5691374*PED	Nov 18, 2015			
	6063811	May 06, 2017			U-602
	6063811*PED	Nov 06, 2017			
	6239173	Jun 24, 2013	DS	DP	U-602
	6239173*PED	Dec 24, 2013			

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATON DATE		PATENT CODES			EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE	
ROFECOXIB; VIOXX									
021052 001	5474995	Jun 24, 2013	U-266	U-266	M-27 PED	Aug 06, 2006 Feb 06, 2007			
	5474995*PED	Dec 24, 2013							
	5691374	May 18, 2015							
	5691374*PED	Nov 18, 2015							
	6063811	May 06, 2017							
	6063811*PED	Nov 06, 2017							
	6239173	Jun 24, 2013							
	6239173*PED	Dec 24, 2013							
ROFECOXIB; VIOXX									
021052 002	5474995	Jun 24, 2013	U-266	U-266	M-27 PED	Aug 06, 2006 Feb 06, 2007			
	5474995*PED	Dec 24, 2013							
	5691374	May 18, 2015							
	5691374*PED	Nov 18, 2015							
	6063811	May 06, 2017							
	6063811*PED	Nov 06, 2017							
	6239173	Jun 24, 2013							
	6239173*PED	Dec 24, 2013							
ROPINIROLE HYDROCHLORIDE; REQUIP									
020658 001	4452808	Dec 07, 2007	DS DP	U-212	I-464	May 04, 2008			
	4824860	May 19, 2008							
ROPINIROLE HYDROCHLORIDE; REQUIP									
020658 002	4452808	Dec 07, 2007	DS DP	U-212	I-464	May 04, 2008			
	4824860	May 19, 2008							
ROPINIROLE HYDROCHLORIDE; REQUIP									
020658 003	4452808	Dec 07, 2007	DS DP	U-212	I-464	May 04, 2008			
	4824860	May 19, 2008							
ROPINIROLE HYDROCHLORIDE; REQUIP									
020658 004	4452808	Dec 07, 2007	DS DP	U-212	I-464	May 04, 2008			
	4824860	May 19, 2008							
ROPINIROLE HYDROCHLORIDE; REQUIP									
020658 005	4452808	Dec 07, 2007	DS DP	U-212	I-464	May 04, 2008			
	4824860	May 19, 2008							
ROPINIROLE HYDROCHLORIDE; REQUIP									
020658 006	4452808	Dec 07, 2007	DS DP		I-464	May 04, 2008			
	4824860	May 19, 2008							
ROPINIROLE HYDROCHLORIDE; REQUIP									
020658 007	4452808	Dec 07, 2007	DS DP		I-464	May 04, 2008			
	4824860	May 19, 2008							
ROPIVACAINE HYDROCHLORIDE MONOHYDRATE; NAROPIN									
020533 001	4870086	Sep 24, 2010							
ROPIVACAINE HYDROCHLORIDE MONOHYDRATE; NAROPIN									
020533 003	4870086	Sep 24, 2010							
ROPIVACAINE HYDROCHLORIDE MONOHYDRATE; NAROPIN									
020533 004	4870086	Sep 24, 2010							
ROPIVACAINE HYDROCHLORIDE MONOHYDRATE; NAROPIN									
020533 005	4870086	Sep 24, 2010							
ROSIGLITAZONE MALEATE; AVANDIA									
021071 002	5002953	Sep 17, 2011	DS DP	U-628	I-453 PED	Feb 28, 2008 Aug 27, 2006			
	5002953	Sep 17, 2011							
	5002953*PED	Mar 17, 2012	DS DP						
	5741803	Apr 21, 2015							
	5741803	Apr 21, 2015	DS DP						
	5741803*PED	Oct 21, 2015							
	6288095	Feb 11, 2017	U-420						
	6288095*PED	Aug 11, 2017							

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>ROSIGLITAZONE MALEATE; AVANDIA</u>					
021071 003	5002953	Sep 17, 2011	DS DP U-329	I-453	Feb 28, 2008
	5002953	Sep 17, 2011	DS DP U-628	PED	Aug 27, 2006
	5002953*PED	Mar 17, 2012			
	5741803	Apr 21, 2015	DS DP U-329		
	5741803	Apr 21, 2015	DS DP U-628		
	5741803*PED	Oct 21, 2015			
	6288095	Feb 11, 2017		U-420	
	6288095*PED	Aug 11, 2017			
<u>ROSIGLITAZONE MALEATE; AVANDIA</u>					
021071 004	5002953	Sep 17, 2011	DS DP U-329	I-453	Feb 28, 2008
	5002953	Sep 17, 2011	DS DP U-628	PED	Aug 27, 2006
	5002953*PED	Mar 17, 2012			
	5741803	Apr 21, 2015	DS DP U-329		
	5741803	Apr 21, 2015	DS DP U-628		
	5741803*PED	Oct 21, 2015			
	6288095	Feb 11, 2017		U-420	
	6288095*PED	Aug 11, 2017			
<u>ROSUVASTATIN CALCIUM; CRESTOR</u>					
021366 002	6316460	Aug 04, 2020		NCE	Aug 12, 2008
	6589959	Dec 23, 2019			
	6858618	Dec 17, 2021	U-618		
	RE37314	Jun 12, 2012			
<u>ROSUVASTATIN CALCIUM; CRESTOR</u>					
021366 003	6316460	Aug 04, 2020		NCE	Aug 12, 2008
	6589959	Dec 23, 2019			
	6858618	Dec 17, 2021	U-618		
	RE37314	Jun 12, 2012			
<u>ROSUVASTATIN CALCIUM; CRESTOR</u>					
021366 004	6316460	Aug 04, 2020		NCE	Aug 12, 2008
	6589959	Dec 23, 2019			
	6858618	Dec 17, 2021	U-618		
	RE37314	Jun 12, 2012			
<u>ROSUVASTATIN CALCIUM; CRESTOR</u>					
021366 005	6316460	Aug 04, 2020		NCE	Aug 12, 2008
	6589959	Dec 23, 2019			
	6858618	Dec 17, 2021	U-618		
	RE37314	Jun 12, 2012			
<u>SALMETEROL XINAFOATE; SEREVENT</u>					
020236 001	4992474	Feb 12, 2008			
	4992474*PED	Aug 12, 2008			
	5126375	Feb 12, 2008			
	5126375*PED	Aug 12, 2008			
	5225445	Feb 12, 2008	U-182		
	5225445*PED	Aug 12, 2008			
<u>SALMETEROL XINAFOATE; SEREVENT</u>					
020692 001	4992474	Feb 12, 2008			
	4992474*PED	Aug 12, 2008			
	5126375	Feb 12, 2008			
	5126375*PED	Aug 12, 2008			
	5225445	Feb 12, 2008	U-211		
	5225445*PED	Aug 12, 2008			
	6536427	Mar 01, 2011	DP		
<u>SAMARIUM SM 153 LEXIDRONAM PENTASODIUM; QUADRAMET</u>					
020570 001	4898724	Feb 06, 2007		U-187	
<u>SAQUINAVIR; FORTOVASE</u>					
020828 001	5196438	Nov 19, 2010			
	6008228	Jun 06, 2015			
	6352717	Nov 16, 2019			
<u>SAQUINAVIR MESYLATE; INVIRASE</u>					
020628 001	5196438	Nov 19, 2010			
<u>SECRETIN SYNTHETIC HUMAN; CHIRHOSTIM</u>					
021256 001				NCE ODE	Apr 09, 2009 Apr 04, 2009

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>SECRETIN SYNTHETIC PORCINE; SECREFOLO</u>					
021136 001				ODE	Apr 04, 2009
<u>SELEGILINE; EMSAM</u>					
021336 001	7070808 RE34579	May 10, 2018 Aug 18, 2007	DS DP DS DP U-711	NDF	Feb 27, 2009
<u>SELEGILINE; EMSAM</u>					
021336 002	RE34579	Aug 18, 2007	DS DP U-711	NDF	Feb 27, 2009
<u>SELEGILINE; EMSAM</u>					
021336 003	RE34579	Aug 18, 2007	DS DP U-711	NDF	Feb 27, 2009
<u>SELEGILINE HYDROCHLORIDE; ZELAPAR</u>					
021479 001	5648093 6423342	Jul 15, 2014 Mar 01, 2016	DP DP	NDF	Jun 14, 2009
<u>SERTACONAZOLE NITRATE; ERTACZO</u>					
021385 001	5135943	May 31, 2014	DS DP U-786	NCE	Dec 10, 2008
<u>SERTRALINE HYDROCHLORIDE; SERTRALINE HYDROCHLORIDE</u>					
075719 001				PC	Feb 06, 2007
<u>SERTRALINE HYDROCHLORIDE; SERTRALINE HYDROCHLORIDE</u>					
075719 002				PC	Feb 06, 2007
<u>SERTRALINE HYDROCHLORIDE; SERTRALINE HYDROCHLORIDE</u>					
075719 003				PC	Feb 06, 2007
<u>SERTRALINE HYDROCHLORIDE; SERTRALINE HYDROCHLORIDE</u>					
076934 001				PC	Feb 03, 2007
<u>SERTRALINE HYDROCHLORIDE; ZOLOFT</u>					
019839 001	4962128 4962128*PED 5248699 5248699*PED 5744501 5789449	Nov 02, 2009 May 02, 2010 Aug 13, 2012 Feb 13, 2013 Jan 06, 2009 Jan 06, 2009	U-152 U-152 U-461 U-460		
<u>SERTRALINE HYDROCHLORIDE; ZOLOFT</u>					
019839 002	4962128 4962128*PED 5248699 5248699*PED 5744501 5789449	Nov 02, 2009 May 02, 2010 Aug 13, 2012 Feb 13, 2013 Jan 06, 2009 Jan 06, 2009	U-152 U-152 U-461 U-460		
<u>SERTRALINE HYDROCHLORIDE; ZOLOFT</u>					
019839 003	4962128 4962128*PED 5248699 5248699*PED 5744501 5789449	Nov 02, 2009 May 02, 2010 Aug 13, 2012 Feb 13, 2013 Jan 06, 2009 Jan 06, 2009	U-152 U-152 U-461 U-460		
<u>SERTRALINE HYDROCHLORIDE; ZOLOFT</u>					
019839 004	4962128 4962128*PED 5248699 5248699*PED 5744501 5789449	Nov 02, 2009 May 02, 2010 Aug 13, 2012 Feb 13, 2013 Jan 06, 2009 Jan 06, 2009	U-152 U-152 U-461 U-460		
<u>SERTRALINE HYDROCHLORIDE; ZOLOFT</u>					
019839 005	4962128 4962128*PED 5248699 5248699*PED 5744501 5789449	Nov 02, 2009 May 02, 2010 Aug 13, 2012 Feb 13, 2013 Jan 06, 2009 Jan 06, 2009	U-152 U-152 U-461 U-460		

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>SERTRALINE HYDROCHLORIDE; ZOLOFT</u>					
020990 001	5248699	Aug 13, 2012			
	5248699*PED	Feb 13, 2013			
	5744501	Jan 06, 2009		U-461	
	5789449	Jan 06, 2009		U-460	
	6727283	Oct 11, 2019	DP	U-580	
	6727283*PED	Apr 11, 2020			
	7067555	Nov 10, 2019	DP		
	7067555*PED	May 10, 2020			
<u>SEVELAMER HYDROCHLORIDE; RENAGEL</u>					
020926 001	5496545	Aug 11, 2013		U-246	
	5667775	Sep 16, 2014		U-246	
	6509013	Aug 11, 2013			
<u>SEVELAMER HYDROCHLORIDE; RENAGEL</u>					
021179 001	5496545	Aug 11, 2013		U-246	
	5667775	Sep 16, 2014		U-246	
	6509013	Aug 11, 2013			
	6733780	Oct 18, 2020	DP		
	7014846	Aug 11, 2013	DP	U-246	
<u>SEVELAMER HYDROCHLORIDE; RENAGEL</u>					
021179 002	5496545	Aug 11, 2013		U-246	
	5667775	Sep 16, 2014		U-246	
	6509013	Aug 13, 2013			
	6733780	Oct 18, 2020	DP		
	7014846	Aug 11, 2013	DP	U-246	
<u>SEVOFLURANE; ULTANE</u>					
020478 001	5990176	Jan 27, 2017			
	5990176*PED	Jul 27, 2017			
	6074668	Jan 09, 2018			
	6074668*PED	Jul 09, 2018			
	6288127	Jan 27, 2017			
	6288127*PED	Jul 27, 2017			
	6444859	Jan 27, 2017			
	6444859*PED	Jul 27, 2017			
<u>SIBUTRAMINE HYDROCHLORIDE; MERIDIA</u>					
020632 001	4746680	Jun 11, 2007			
	4746680*PED	Dec 11, 2007			
	4929629	May 29, 2007			
	4929629*PED	Nov 29, 2007			
	5436272	Jul 25, 2012		U-439	
	5436272*PED	Jan 25, 2013			
<u>SIBUTRAMINE HYDROCHLORIDE; MERIDIA</u>					
020632 002	4746680	Jun 11, 2007			
	4746680*PED	Dec 11, 2007			
	4929629	May 29, 2007			
	4929629*PED	Nov 29, 2007			
	5436272	Jul 25, 2012		U-439	
	5436272*PED	Jan 25, 2013			
<u>SIBUTRAMINE HYDROCHLORIDE; MERIDIA</u>					
020632 003	4746680	Jun 11, 2007			
	4746680*PED	Dec 11, 2007			
	4929629	May 29, 2007			
	4929629*PED	Nov 29, 2007			
	5436272	Jul 25, 2012		U-439	
	5436272*PED	Jan 25, 2013			
<u>SILDENAFIL CITRATE; REVATIO</u>					
021845 001	5250534	Mar 27, 2012	DS DP	NP	Jun 03, 2008
<u>SILDENAFIL CITRATE; VIAGRA</u>					
020895 001	5250534	Mar 27, 2012			
	6469012	Oct 22, 2019		U-155	
<u>SILDENAFIL CITRATE; VIAGRA</u>					
020895 002	5250534	Mar 27, 2012			
	6469012	Oct 22, 2019		U-155	

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>SILDENAFIL CITRATE; VIAGRA</u>					
020895 003	5250534	Mar 27, 2012			
	6469012	Oct 22, 2019	U-155		
<u>SIMVASTATIN; SIMVASTATIN</u>					
076052 001				PC	Dec 20, 2006
<u>SIMVASTATIN; SIMVASTATIN</u>					
076052 002				PC	Dec 20, 2006
<u>SIMVASTATIN; SIMVASTATIN</u>					
076052 003				PC	Dec 20, 2006
<u>SIMVASTATIN; SIMVASTATIN</u>					
076052 004				PC	Dec 20, 2006
<u>SIMVASTATIN; SIMVASTATIN</u>					
076285 005				PC	Dec 20, 2006
<u>SINCALIDE; KINEVAC</u>					
017697 001	6803046	Aug 16, 2022	DP		
<u>SIROLIMUS; RAPAMUNE</u>					
021083 001	5100899	Jul 07, 2013	U-290	NPP	Mar 11, 2008
	5100899*PED	Jan 07, 2014		PED	Sep 11, 2008
	5212155	May 18, 2010	U-291	PED	Oct 11, 2006
	5212155*PED	Nov 18, 2010			
	5308847	May 03, 2011	U-292		
	5308847*PED	Nov 03, 2011			
	5403833	Apr 04, 2012	U-293		
	5403833*PED	Oct 04, 2012			
	5536729	Sep 30, 2013	DP		
	5536729*PED	Mar 30, 2014			
<u>SIROLIMUS; RAPAMUNE</u>					
021110 001	5100899	Jul 07, 2013	U-290	NPP	Mar 11, 2008
	5100899*PED	Jan 07, 2014		PED	Sep 11, 2008
	5212155	May 18, 2010	U-291	PED	Oct 11, 2006
	5212155*PED	Nov 18, 2010			
	5308847	May 03, 2011	U-292		
	5308847*PED	Nov 03, 2011			
	5403833	Apr 04, 2012	U-293		
	5403833*PED	Oct 04, 2012			
	5989591	Mar 11, 2018	DP		
	5989591*PED	Sep 11, 2018			
<u>SIROLIMUS; RAPAMUNE</u>					
021110 002	5100899	Jul 07, 2013	U-290	NPP	Mar 11, 2008
	5100899*PED	Jan 07, 2014		PED	Sep 11, 2008
	5212155	May 18, 2010	U-291	PED	Oct 11, 2006
	5212155*PED	Nov 18, 2010			
	5308847	May 03, 2011	U-292		
	5308847*PED	Nov 03, 2011			
	5403833	Apr 04, 2012	U-293		
	5403833*PED	Oct 04, 2012			
	5989591	Mar 11, 2018	DP		
	5989591*PED	Sep 11, 2018			
<u>SIROLIMUS; RAPAMUNE</u>					
021110 003	5100899	Jul 07, 2013	U-290	NPP	Mar 11, 2008
	5100899*PED	Jan 07, 2014		PED	Sep 11, 2008
	5212155	May 18, 2010	U-291	PED	Oct 11, 2006
	5212155*PED	Nov 18, 2010			
	5403833	Apr 04, 2012	U-293		
	5403833*PED	Oct 04, 2012			
	5989591	Mar 11, 2018	DP		
	5989591*PED	Sep 11, 2018			
<u>SITAGLIPTIN PHOSPHATE; JANUVIA</u>					
021995 001	6303661	Apr 24, 2017		NCE	Oct 16, 2011
	6699871	Jul 26, 2022	DS DP	U-774	
	6890898	Feb 02, 2019		U-775	
	7078381	Feb 02, 2019		U-775	
	7125873	Jul 26, 2022		U-775	

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE		PATENT CODES		EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE	
<u>SITAGLIPTIN PHOSPHATE; JANUVIA</u>								
021995 002	6303661	Apr 24, 2017	DS	DP	U-774	NCE	Oct 16, 2011	
	6699871	Jul 26, 2022			U-774			
	6890898	Feb 02, 2019			U-775			
	7078381	Feb 02, 2019			U-775			
	7125873	Jul 26, 2022			U-775			
<u>SITAGLIPTIN PHOSPHATE; JANUVIA</u>								
021995 003	6303661	Apr 24, 2017	DS	DP	U-774	NCE	Oct 16, 2011	
	6699871	Jul 26, 2022			U-774			
	6890898	Feb 02, 2019			U-775			
	7078381	Feb 02, 2019			U-775			
	7125873	Jul 26, 2022			U-775			
<u>SODIUM BENZOATE; SODIUM PHENYLACETATE; AMMONUL</u>								
020645 001						NDF	Feb 17, 2008	
						ODE	Feb 17, 2012	
<u>SODIUM FERRIC GLUCONATE COMPLEX; FERRLECIT</u>								
020955 001						NPP	Aug 13, 2007	
						PED	Feb 13, 2008	
<u>SODIUM FLUORIDE; TRICLOSAN; COLGATE TOTAL</u>								
020231 001	4894220	Jan 30, 2007		DP				
	5037635	Jan 30, 2007		DP				
	5156835	Oct 20, 2009		DP				
	5288480	Jan 30, 2007		DP				
	5344641	Jan 30, 2007		DP				
	5538715	Jan 30, 2007		DP				
	5776435	Jan 30, 2007		DP				
<u>SODIUM OXYBATE; XYREM</u>								
021196 001	6780889	Dec 22, 2019		DP		NCE	Jul 17, 2007	
						ODE	Nov 18, 2012	
						ODE	Jul 17, 2009	
<u>SODIUM PHOSPHATE, DIBASIC, ANHYDROUS; SODIUM PHOSPHATE, MONOBASIC, MONOHYDRATE; OSMOPREP</u>								
021892 001	5616346	May 18, 2013		DP	U-715	NP	Mar 16, 2009	
<u>SODIUM PHOSPHATE, DIBASIC, ANHYDROUS; SODIUM PHOSPHATE, MONOBASIC, MONOHYDRATE; VISICOL</u>								
021097 001	5616346	May 18, 2013			U-359			
<u>SOLIFENACIN SUCCINATE; VESICARE</u>								
021518 001	6017927	Dec 27, 2015	DS	DP		NCE	Nov 19, 2009	
<u>SOLIFENACIN SUCCINATE; VESICARE</u>								
021518 002	6017927	Dec 27, 2015	DS	DP		NCE	Nov 19, 2009	
<u>SOMATROPIN RECOMBINANT; GENOTROPIN</u>								
020280 006	4968299	Jun 28, 2008		DP		I-496	Apr 27, 2009	
						ODE	Jul 25, 2008	
						ODE	Jun 20, 2007	
<u>SOMATROPIN RECOMBINANT; GENOTROPIN</u>								
020280 007	4968299	Jun 28, 2008		DP		I-496	Apr 27, 2009	
						ODE	Jul 25, 2008	
						ODE	Jun 20, 2007	
<u>SOMATROPIN RECOMBINANT; GENOTROPIN PRESERVATIVE FREE</u>								
020280 001	4968299	Jun 28, 2008		DP		I-496	Apr 27, 2009	
	5435076	Apr 16, 2013		DP		ODE	Jul 25, 2008	
	5716338	Feb 10, 2015		DP		ODE	Jun 20, 2007	
	6152897	Nov 20, 2018		DP				
<u>SOMATROPIN RECOMBINANT; GENOTROPIN PRESERVATIVE FREE</u>								
020280 002	4968299	Jun 28, 2008		DP		I-496	Apr 27, 2009	
	5435076	Apr 16, 2013		DP		ODE	Jul 25, 2008	
	5716338	Feb 10, 2015		DP		ODE	Jun 20, 2007	
	6152897	Nov 20, 2018		DP				

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>SOMATROPIN RECOMBINANT; GENOTROPIN PRESERVATIVE FREE</u>					
020280 003	4968299	Jun 28, 2008	DP	I-496	Apr 27, 2009
	5435076	Apr 16, 2013	DP	ODE	Jul 25, 2008
	5716338	Feb 10, 2015	DP	ODE	Jun 20, 2007
	6152897	Nov 20, 2018	DP		
<u>SOMATROPIN RECOMBINANT; GENOTROPIN PRESERVATIVE FREE</u>					
020280 004	4968299	Jun 28, 2008	DP	I-496	Apr 27, 2009
				ODE	Jul 25, 2008
				ODE	Jun 20, 2007
<u>SOMATROPIN RECOMBINANT; GENOTROPIN PRESERVATIVE FREE</u>					
020280 005	4968299	Jun 28, 2008	DP	I-496	Apr 27, 2009
	5435076	Apr 16, 2013	DP	ODE	Jul 25, 2008
	5716338	Feb 10, 2015	DP	ODE	Jun 20, 2007
	6152897	Nov 20, 2018	DP		
<u>SOMATROPIN RECOMBINANT; GENOTROPIN PRESERVATIVE FREE</u>					
020280 008	4968299	Jun 28, 2008	DP	I-496	Apr 27, 2009
	5435076	Apr 16, 2013	DP	ODE	Jul 25, 2008
	5716338	Feb 10, 2015	DP	ODE	Jun 20, 2007
	6152897	Nov 20, 2018	DP		
<u>SOMATROPIN RECOMBINANT; GENOTROPIN PRESERVATIVE FREE</u>					
020280 009	4968299	Jun 28, 2008	DP	I-496	Apr 27, 2009
	5435076	Apr 16, 2013	DP	ODE	Jul 25, 2008
	5716338	Feb 10, 2015	DP	ODE	Jun 20, 2007
	6152897	Nov 20, 2018	DP		
<u>SOMATROPIN RECOMBINANT; GENOTROPIN PRESERVATIVE FREE</u>					
020280 010	4968299	Jun 28, 2008	DP	I-496	Apr 27, 2009
	5435076	Apr 16, 2013	DP	ODE	Jul 25, 2008
	5716338	Feb 10, 2015	DP	ODE	Jun 20, 2007
<u>SOMATROPIN RECOMBINANT; GENOTROPIN PRESERVATIVE FREE</u>					
020280 011	4968299	Jun 28, 2008	DP	I-496	Apr 27, 2009
	5435076	Apr 16, 2013	DP	ODE	Jul 25, 2008
	5716338	Feb 10, 2015	DP	ODE	Jun 20, 2007
<u>SOMATROPIN RECOMBINANT; GENOTROPIN PRESERVATIVE FREE</u>					
020280 012	4968299	Jun 28, 2008	DP	I-496	Apr 27, 2009
	5435076	Apr 16, 2013	DP	ODE	Jul 25, 2008
	5716338	Feb 10, 2015	DP	ODE	Jun 20, 2007
<u>SOMATROPIN RECOMBINANT; GENOTROPIN PRESERVATIVE FREE</u>					
020280 013	4968299	Jun 28, 2008	DP	I-496	Apr 27, 2009
	5435076	Apr 16, 2013	DP	ODE	Jul 25, 2008
	5716338	Feb 10, 2015	DP	ODE	Jun 20, 2007
<u>SOMATROPIN RECOMBINANT; HUMATROPE</u>					
019640 001				I-398	Jul 25, 2006
				M-45	Oct 12, 2008
<u>SOMATROPIN RECOMBINANT; HUMATROPE</u>					
019640 004				I-518	Nov 01, 2009
				I-398	Jul 25, 2006
				M-45	Oct 12, 2008
<u>SOMATROPIN RECOMBINANT; HUMATROPE</u>					
019640 005				I-518	Nov 01, 2009
				I-398	Jul 25, 2006
				M-45	Oct 12, 2008
<u>SOMATROPIN RECOMBINANT; HUMATROPE</u>					
019640 006				I-518	Nov 01, 2009
				I-398	Jul 25, 2006
				M-45	Oct 12, 2008

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>SOMATROPIN RECOMBINANT; HUMATROPE</u>					
019640	007			I-518 I-398 M-45	Nov 01, 2009 Jul 25, 2006 Oct 12, 2008
<u>SOMATROPIN RECOMBINANT; NORDITROPIN</u>					
021148	001	5849700 5849704	Dec 15, 2015 Dec 15, 2015	U-340 I-439	Nov 01, 2007
<u>SOMATROPIN RECOMBINANT; NORDITROPIN</u>					
021148	002	5849700 5849704	Dec 15, 2015 Dec 15, 2015	U-340 I-439	Nov 01, 2007
<u>SOMATROPIN RECOMBINANT; NORDITROPIN</u>					
021148	003	5633352 5849700 5849704	May 27, 2014 Dec 15, 2015 Dec 15, 2015	I-439 U-340	Nov 01, 2007
<u>SOMATROPIN RECOMBINANT; NORDITROPIN NORDIFLEX</u>					
021148	004			I-439	Nov 01, 2007
<u>SOMATROPIN RECOMBINANT; NORDITROPIN NORDIFLEX</u>					
021148	005			I-439	Nov 01, 2007
<u>SOMATROPIN RECOMBINANT; NORDITROPIN NORDIFLEX</u>					
021148	006			I-439	Nov 01, 2007
<u>SOMATROPIN RECOMBINANT; NUTROPIN</u>					
020168	001	5096885	Mar 17, 2009	DP I-462	Jun 28, 2008
<u>SOMATROPIN RECOMBINANT; NUTROPIN</u>					
020168	002	5096885	Mar 17, 2009	DP I-462	Jun 28, 2008
<u>SOMATROPIN RECOMBINANT; NUTROPIN AQ</u>					
020522	001	5763394	Jun 09, 2015	DP I-462 M-50	Jun 28, 2008 Nov 18, 2008
<u>SOMATROPIN RECOMBINANT; NUTROPIN AQ PEN</u>					
020522	002	5763394	Jun 09, 2015	DP I-462 M-50	Jun 28, 2008 Nov 18, 2008
<u>SOMATROPIN RECOMBINANT; NUTROPIN DEPOT</u>					
021075	001	5654010 5656297 5912015 6051259	Aug 05, 2014 Jul 25, 2014 Mar 12, 2012 Dec 02, 2012	U-340	
<u>SOMATROPIN RECOMBINANT; NUTROPIN DEPOT</u>					
021075	002	5654010 5656297 5912015 6051259	Aug 05, 2014 Jul 25, 2014 Mar 12, 2012 Dec 02, 2012	U-340	
<u>SOMATROPIN RECOMBINANT; NUTROPIN DEPOT</u>					
021075	003	5654010 5656297 5912015 6051259	Aug 05, 2014 Jul 25, 2014 Mar 12, 2012 Dec 02, 2012	U-340	
<u>SOMATROPIN RECOMBINANT; OMNITROPE</u>					
021426	001			NP	May 30, 2009
<u>SOMATROPIN RECOMBINANT; OMNITROPE</u>					
021426	002			NP	May 30, 2009
<u>SOMATROPIN RECOMBINANT; SAIZEN</u>					
019764	001			I-440	Aug 26, 2007
<u>SOMATROPIN RECOMBINANT; SAIZEN</u>					
019764	002			I-440	Aug 26, 2007
<u>SOMATROPIN RECOMBINANT; SAIZEN</u>					
019764	003			I-440	Aug 26, 2007

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>SOMATROPIN RECOMBINANT; ZORBTIVE</u>					
021597 001				NP ODE	Dec 01, 2006 Dec 01, 2010
<u>SOMATROPIN RECOMBINANT; ZORBTIVE</u>					
021597 002				NP ODE	Dec 01, 2006 Dec 01, 2010
<u>SOMATROPIN RECOMBINANT; ZORBTIVE</u>					
021597 003				NP ODE	Dec 01, 2006 Dec 01, 2010
<u>SOMATROPIN RECOMBINANT; ZORBTIVE</u>					
021597 004				NP ODE	Dec 01, 2006 Dec 01, 2010
<u>SORAFENIB TOSYLATE; NEXAVAR</u>					
021923 001				NCE ODE	Dec 20, 2010 Dec 20, 2012
<u>SPARFLOXACIN; ZAGAM</u>					
020677 001	4795751	Feb 04, 2010	U-160		
<u>STAVUDINE; ZERIT</u>					
020412 001	4978655 4978655*PED	Jun 24, 2008 Dec 24, 2008	U-94 U-94		
<u>STAVUDINE; ZERIT</u>					
020412 002	4978655 4978655*PED	Jun 24, 2008 Dec 24, 2008	U-94 U-94		
<u>STAVUDINE; ZERIT</u>					
020412 003	4978655 4978655*PED	Jun 24, 2008 Dec 24, 2008	U-94 U-94		
<u>STAVUDINE; ZERIT</u>					
020412 004	4978655 4978655*PED	Jun 24, 2008 Dec 24, 2008	U-94 U-94		
<u>STAVUDINE; ZERIT</u>					
020412 005	4978655 4978655*PED	Jun 24, 2008 Dec 24, 2008	U-94 U-94		
<u>STAVUDINE; ZERIT</u>					
020413 001	4978655 4978655*PED	Jun 24, 2008 Dec 24, 2008			
<u>STAVUDINE; ZERIT XR</u>					
021453 001	4978655 4978655*PED 7135465 7135465*PED	Jun 24, 2008 Dec 24, 2008 Feb 18, 2023 Aug 18, 2023	U-248 U-248 U-167	PED DP	Jul 01, 2006
<u>STAVUDINE; ZERIT XR</u>					
021453 002	4978655 4978655*PED 7135465 7135465*PED	Jun 24, 2008 Dec 24, 2008 Feb 18, 2023 Aug 18, 2023	U-248 U-248 U-167	PED DP	Jul 01, 2006
<u>STAVUDINE; ZERIT XR</u>					
021453 003	4978655 4978655*PED 7135465 7135465*PED	Jun 24, 2008 Dec 24, 2008 Feb 18, 2023 Aug 18, 2023	U-248 U-167	PED DP	Jul 01, 2006
<u>STAVUDINE; ZERIT XR</u>					
021453 004	4978655 4978655*PED 7135465 7135465*PED	Jun 24, 2008 Dec 24, 2008 Feb 18, 2023 Aug 18, 2023	U-248 U-248 U-167	PED DP	Jul 01, 2006

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>SUMATRIPTAN; IMITREX</u>					
020626 001	4816470	Dec 28, 2006	U-72		
	4816470*PED	Jun 28, 2007			
	5037845	Aug 06, 2008			
	5037845*PED	Feb 06, 2009			
	5307953	Dec 02, 2012			
	5307953*PED	Jun 02, 2013			
	5554639	Sep 10, 2013		U-232	
	5554639*PED	Mar 10, 2014			
	5705520	Dec 10, 2011		U-232	
	5705520*PED	Jun 10, 2012			
<u>SUMATRIPTAN; IMITREX</u>					
020626 002	4816470	Dec 28, 2006	U-72		
	4816470*PED	Jun 28, 2007			
	5037845	Aug 06, 2008			
	5037845*PED	Feb 06, 2009			
	5307953	Dec 02, 2012			
	5307953*PED	Jun 02, 2013			
	5554639	Sep 10, 2013		U-232	
	5554639*PED	Mar 10, 2014			
	5705520	Dec 10, 2011		U-232	
	5705520*PED	Jun 10, 2012			
<u>SUMATRIPTAN; IMITREX</u>					
020626 003	4816470	Dec 28, 2006	U-72		
	4816470*PED	Jun 28, 2007			
	5037845	Aug 06, 2008			
	5037845*PED	Feb 06, 2009			
	5307953	Dec 02, 2012			
	5307953*PED	Jun 02, 2013			
	5554639	Sep 10, 2013		U-232	
	5554639*PED	Mar 10, 2014			
	5705520	Dec 10, 2011		U-232	
	5705520*PED	Jun 10, 2012			
<u>SUMATRIPTAN SUCCINATE; IMITREX</u>					
020080 001	4816470	Dec 28, 2006		U-72	
	4816470*PED	Jun 28, 2007			
	5037845	Aug 06, 2008		U-72	
	5037845*PED	Feb 06, 2009			
<u>SUMATRIPTAN SUCCINATE; IMITREX</u>					
020132 001	4816470	Dec 28, 2006		U-72	
	4816470*PED	Jun 28, 2007			
	5037845	Aug 06, 2008		U-72	
	5037845*PED	Feb 06, 2009			
	5863559	Jan 26, 2016		U-72	
	5863559*PED	Jul 26, 2016			
	6020001	Mar 02, 2012		U-444	
	6020001*PED	Sep 02, 2012			
	6368627	Mar 02, 2012		U-444	
	6368627*PED	Sep 02, 2012			
<u>SUMATRIPTAN SUCCINATE; IMITREX</u>					
020132 002	4816470	Dec 28, 2006		U-72	
	4816470*PED	Jun 28, 2007			
	5037845	Aug 06, 2008		U-72	
	5037845*PED	Feb 06, 2009			
	5863559	Jan 26, 2016		U-72	
	5863559*PED	Jul 26, 2016			
	6020001	Mar 02, 2012		U-444	
	6020001*PED	Sep 02, 2012			
	6368627	Mar 02, 2012		U-444	
	6368627*PED	Sep 02, 2012			

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE		PATENT CODES			EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE			
SUMATRIPTAN SUCCINATE; IMITREX											
020132 003	4816470	Dec	28,	2006	U-72						
	4816470*PED	Jun	28,	2007							
	5037845	Aug	06,	2008	U-72						
	5037845*PED	Feb	06,	2009							
	5863559	Jan	26,	2016	U-72						
	5863559*PED	Jul	26,	2016							
	6020001	Mar	02,	2012	U-444						
	6020001*PED	Sep	02,	2012							
	6368627	Mar	02,	2012	U-444						
6368627*PED	Sep	02,	2012								
SUMATRIPTAN SUCCINATE; IMITREX STATDOSE											
020080 003	4816470	Dec	28,	2006	U-72						
	4816470*PED	Jun	28,	2007							
	5037845	Aug	06,	2008	U-72						
	5037845*PED	Feb	06,	2009							
SUNITINIB MALATE; SUTENT											
021938 001	6573293	Feb	15,	2021	DS	DP	U-703	NCE	Jan	26,	2011
	7125905	Feb	15,	2021	DS	DP					
SUNITINIB MALATE; SUTENT											
021938 002	6573293	Feb	15,	2021	DS	DP	U-703	NCE	Jan	26,	2011
	7125905	Feb	15,	2021	DS	DP					
SUNITINIB MALATE; SUTENT											
021938 003	6573293	Feb	15,	2021	DS	DP	U-703	NCE	Jan	26,	2011
	7125905	Feb	15,	2021	DS	DP					
TACRINE HYDROCHLORIDE; COGNEX											
020070 001	4816456	Sep	09,	2007			U-82				
TACRINE HYDROCHLORIDE; COGNEX											
020070 002	4816456	Sep	09,	2007			U-82				
TACRINE HYDROCHLORIDE; COGNEX											
020070 003	4816456	Sep	09,	2007			U-82				
TACRINE HYDROCHLORIDE; COGNEX											
020070 004	4816456	Sep	09,	2007			U-82				
TACROLIMUS; PROGRAF											
050708 001								ODE	Mar	29,	2013
TACROLIMUS; PROGRAF											
050708 002								ODE	Mar	29,	2013
TACROLIMUS; PROGRAF											
050708 003								ODE	Mar	29,	2013
TACROLIMUS; PROGRAF											
050709 001								ODE	Mar	29,	2013
TADALAFIL; CIALIS											
021368 001	5859006	Jan	12,	2016	DS	DP		NCE	Nov	21,	2008
	6140329	Jul	11,	2016	DP	U-155					
	6821975	Nov	19,	2020	DS	DP	U-614				
	6821975	Nov	19,	2020	DS	DP	U-533				
	6943166	Apr	26,	2020			U-614				
TADALAFIL; CIALIS											
021368 002	5859006	Jan	12,	2016	DS	DP		NCE	Nov	21,	2008
	6140329	Jul	11,	2016	DP	U-155					
	6821975	Nov	19,	2020	DS	DP	U-614				
	6821975	Nov	19,	2020	DS	DP	U-533				
	6943166	Apr	26,	2020			U-614				
TADALAFIL; CIALIS											
021368 003	5859006	Jan	12,	2016	DS	DP		NCE	Nov	21,	2008
	6140329	Jul	11,	2016	DP	U-155					
	6821975	Nov	19,	2020	DS	DP	U-533				
	6821975	Nov	19,	2020	DS	DP	U-614				
	6943166	Apr	26,	2020			U-614				
TAMOXIFEN CITRATE; SOLTAMOX											
021807 001	6127425	Jun	26,	2018		DP					

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>TAMSULOSIN HYDROCHLORIDE; FLOMAX</u>					
020579 001	4703063	Oct 27, 2009			
	4868216	Sep 19, 2006	U-181		
<u>TAZAROTENE; AVAGE</u>					
021184 003	5089509	Jun 13, 2011	U-481		
<u>TAZAROTENE; TAZORAC</u>					
020600 001	5089509	Jun 13, 2011	U-481		
	5089509	Jun 13, 2011	U-193		
	5914334	Jun 07, 2014	U-517		
	6258830	Jun 07, 2014	U-517		
<u>TAZAROTENE; TAZORAC</u>					
020600 002	5089509	Jun 13, 2011	U-481		
	5089509	Jun 13, 2011	U-193		
	5914334	Jun 07, 2014	U-517		
	6258830	Jun 07, 2014	U-517		
<u>TAZAROTENE; TAZORAC</u>					
021184 001	5089509	Jun 13, 2011	U-481		
<u>TAZAROTENE; TAZORAC</u>					
021184 002	5089509	Jun 13, 2011	U-481		
<u>TECHNETIUM TC-99M APCITIDE; ACUTECT</u>					
020887 001	5443815	Aug 22, 2012			
	5508020	Apr 16, 2013			
	5645815	Jul 08, 2014			
<u>TECHNETIUM TC-99M BICISATE KIT; NEUROLITE</u>					
020256 001	5279811	Nov 23, 2008	U-101		
	5431900	Jul 11, 2012	U-336		
<u>TECHNETIUM TC-99M SESTAMIBI KIT; CARDIOLITE</u>					
019785 001	4885100	Sep 11, 2007			
	4894445	Jan 16, 2007	U-337		
	4988827	Jan 29, 2008			
	5324824	Jan 16, 2007			
<u>TECHNETIUM TC-99M SESTAMIBI KIT; MIRALUMA</u>					
019785 003	4885100	Sep 11, 2007			
	4894445	Jan 16, 2007	U-337		
	4988827	Jan 29, 2008			
	5324824	Jan 16, 2007			
<u>TECHNETIUM TC-99M TEBOROXIME KIT; CARDIOTEC</u>					
019928 001	6056941	Jul 28, 2019	DP		
<u>TECHNETIUM TC-99M TETROFOSMIN KIT; MYOVUE</u>					
020372 001	5045302	Feb 09, 2010			
<u>TEGASEROD MALEATE; ZELNORM</u>					
021200 001	5510353	Apr 26, 2013	U-466	I-435 NCE	Aug 21, 2007 Jul 24, 2007
<u>TEGASEROD MALEATE; ZELNORM</u>					
021200 002	5510353	Apr 26, 2013	U-466	I-435 NCE	Aug 21, 2007 Jul 24, 2007
<u>TELBIVUDINE; TYZEKA</u>					
022011 001	6395716	Aug 10, 2019	U-782	NCE	Oct 25, 2011
	6444652	Aug 10, 2019	U-782		
	6566344	Aug 10, 2019	U-782		
	6569837	Aug 10, 2019	U-782		
<u>TELITHROMYCIN; KETEK</u>					
021144 001	5635485	Apr 21, 2015	DS DP	NCE	Apr 01, 2009
	D459798	Sep 24, 2015			
<u>TELITHROMYCIN; KETEK</u>					
021144 002	5635485	Apr 21, 2015	DS DP	NCE	Apr 01, 2009
	D459798	Sep 24, 2015			
<u>TELMISARTAN; MICARDIS</u>					
020850 001	5591762	Jan 07, 2014	U-3		
	6358986	Jan 10, 2020			

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE		PATENT CODES			EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE	
<u>TELMISARTAN; MICARDIS</u>									
020850 002	5591762 6358986	Jan 07, 2014 Jan 10, 2020	U-3						
<u>TELMISARTAN; MICARDIS</u>									
020850 003	5591762 6358986	Jan 07, 2014 Jan 10, 2020	U-3						
<u>TEMAZEPAM; RESTORIL</u>									
018163 003	5030632 5211954 5326758	Jul 09, 2008 May 18, 2010 Jul 09, 2008	U-70 U-70						
<u>TEMOZOLOMIDE; TEMODAR</u>									
021029 001	5260291 5260291*PED	Aug 11, 2013 Feb 11, 2014	DS	DP	U-619	I-450 ODE ODE PED	Mar 15, 2008 Aug 11, 2006 Mar 15, 2012 Feb 11, 2007		
<u>TEMOZOLOMIDE; TEMODAR</u>									
021029 002	5260291 5260291*PED	Aug 11, 2013 Feb 11, 2014	DS	DP	U-619	I-450 ODE ODE PED	Mar 15, 2008 Aug 11, 2006 Mar 15, 2012 Feb 11, 2007		
<u>TEMOZOLOMIDE; TEMODAR</u>									
021029 003	5260291 5260291*PED	Aug 11, 2013 Feb 11, 2014	DS	DP	U-619	I-450 ODE ODE PED	Mar 15, 2008 Aug 11, 2006 Mar 15, 2012 Feb 11, 2007		
<u>TEMOZOLOMIDE; TEMODAR</u>									
021029 004	5260291 5260291*PED	Aug 11, 2013 Feb 11, 2014	DS	DP	U-619	I-450 ODE ODE PED	Mar 15, 2008 Aug 11, 2006 Mar 15, 2012 Feb 11, 2007		
<u>TEMOZOLOMIDE; TEMODAR</u>									
021029 005						I-450 ODE ODE PED	Mar 15, 2008 Mar 15, 2012 Aug 11, 2006 Feb 11, 2007		
<u>TEMOZOLOMIDE; TEMODAR</u>									
021029 006						I-450 ODE ODE PED	Mar 15, 2008 Mar 15, 2012 Aug 11, 2006 Feb 11, 2007		
<u>TENOFOVIR DISOPROXIL FUMARATE; VIREAD</u>									
021356 001	5922695 5935946 5977089 6043230	Jul 25, 2017 Jul 25, 2017 Jul 25, 2017 Jul 25, 2017	DS DS DS	DP DP	U-248 U-248 U-248 U-248	NPP NCE	Aug 15, 2006 Oct 26, 2006		
<u>TERAZOSIN HYDROCHLORIDE; HYTRIN</u>									
019057 001	5212176 5294615 5294615 5412095	Jun 29, 2010 Apr 29, 2013 Apr 29, 2013 Apr 29, 2013	U-3 U-165						
<u>TERAZOSIN HYDROCHLORIDE; HYTRIN</u>									
019057 002	5212176 5294615 5294615 5412095	Jun 29, 2010 Apr 29, 2013 Apr 29, 2013 Apr 29, 2013	U-165 U-3						
<u>TERAZOSIN HYDROCHLORIDE; HYTRIN</u>									
019057 003	5212176 5294615 5294615 5412095	Jun 29, 2010 Apr 29, 2013 Apr 29, 2013 Apr 29, 2013	U-3 U-165						

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>TERAZOSIN HYDROCHLORIDE; HYTRIN</u>					
019057 004	5212176	Jun 29, 2010			
	5294615	Apr 29, 2013	U-3		
	5294615	Apr 29, 2013	U-165		
	5412095	Apr 29, 2013			
<u>TERAZOSIN HYDROCHLORIDE; HYTRIN</u>					
020347 001	5212176	Jun 29, 2010			
	5294615	Apr 29, 2013	U-3		
	5294615	Apr 29, 2013	U-165		
	5412095	Apr 29, 2013			
<u>TERAZOSIN HYDROCHLORIDE; HYTRIN</u>					
020347 002	5212176	Jun 29, 2010			
	5294615	Apr 29, 2013	U-3		
	5294615	Apr 29, 2013	U-165		
	5412095	Apr 29, 2013			
<u>TERAZOSIN HYDROCHLORIDE; HYTRIN</u>					
020347 003	5212176	Jun 29, 2010			
	5294615	Apr 29, 2013	U-3		
	5294615	Apr 29, 2013	U-165		
	5412095	Apr 29, 2013			
<u>TERAZOSIN HYDROCHLORIDE; HYTRIN</u>					
020347 004	5212176	Jun 29, 2010			
	5294615	Apr 29, 2013	U-165		
	5294615	Apr 29, 2013	U-3		
	5412095	Apr 29, 2013			
<u>TERBINAFINE; LAMISIL</u>					
020846 001	4755534	Dec 30, 2006		U-445	
	4755534*PED	Jun 30, 2007			
	5681849	Oct 28, 2014	DP		
	5681849*PED	Apr 28, 2015			
	5856355	May 18, 2012	DP	U-540	
	5856355	May 18, 2012	DP	U-504	
	5856355	May 18, 2012	DP	U-502	
	5856355*PED	Nov 18, 2012			
	6005001	May 18, 2012	DP	U-502	
	6005001	May 18, 2012	DP	U-540	
	6005001	May 18, 2012	DP	U-504	
	6005001*PED	Nov 18, 2012			
	6121314	May 18, 2012	DP	U-540	
	6121314	May 18, 2012	DP	U-502	
	6121314	May 18, 2012	DP	U-504	
	6121314*PED	Nov 18, 2012			
<u>TERBINAFINE HYDROCHLORIDE; LAMISIL</u>					
020192 001	4755534	Dec 30, 2006		U-73	
	4755534*PED	Jun 30, 2007			
<u>TERBINAFINE HYDROCHLORIDE; LAMISIL</u>					
020539 001	4755534	Dec 30, 2006		U-73	
	4755534*PED	Jun 30, 2007			
<u>TERBINAFINE HYDROCHLORIDE; LAMISIL</u>					
020749 001	4755534	Dec 30, 2006			
	4755534*PED	Jun 30, 2007			
	6121314	May 18, 2012		U-502	
	6121314*PED	Nov 18, 2012			
<u>TERBINAFINE HYDROCHLORIDE; LAMISIL</u>					
020980 001	4755534	Dec 30, 2006		U-73	
	4755534*PED	Jun 30, 2007			
<u>TERBINAFINE HYDROCHLORIDE; LAMISIL AT</u>					
021124 001	4755534	Dec 30, 2006		U-73	
	4755534*PED	Jun 30, 2007			
	5681849	Oct 28, 2014			
	5681849*PED	Apr 28, 2015			
	6121314	May 18, 2012		U-504	
	6121314*PED	Nov 18, 2012			

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
TERBINAFINE HYDROCHLORIDE; LAMISIL AT					
021124 002	4755534	Dec 30, 2006	U-73	U-504	
	4755534*PED	Jun 30, 2007			
	5681849	Oct 28, 2014			
	5681849*PED	Apr 28, 2015			
	6121314	May 18, 2012			
	6121314*PED	Nov 18, 2012			
TERIPARATIDE RECOMBINANT HUMAN; FORTEO					
021318 001	6770623	Dec 08, 2018	DP	U-597	U-597
	6977077	Aug 19, 2019	DP		
	7144861	Dec 08, 2018			
TESTOSTERONE; ANDRODERM					
020489 001	4849224	Nov 12, 2007	U-490		
	4855294	Sep 06, 2008			
	4863970	Nov 14, 2006			
	4983395	Nov 12, 2007			
	5152997	Dec 11, 2010			
	5164190	Dec 11, 2010			
TESTOSTERONE; ANDRODERM					
020489 002	4849224	Nov 12, 2007	U-490		
	4855294	Sep 06, 2008			
	4863970	Nov 14, 2006			
	4983395	Nov 12, 2007			
	5152997	Dec 11, 2010			
	5164190	Dec 11, 2010			
TESTOSTERONE; ANDROGEL					
021015 001	6503894	Aug 30, 2020		U-490	
TESTOSTERONE; STRIANT					
021543 001	6248358	Aug 23, 2019		U-527	
TESTOSTERONE; TESTIM					
021454 001	5023252	Jun 11, 2008		U-483	
TESTOSTERONE; TESTODERM					
019762 001	5840327	Aug 15, 2016			
TESTOSTERONE; TESTODERM					
019762 002	5840327	Aug 15, 2016			
TESTOSTERONE; TESTODERM TTS					
020791 001	6348210	Nov 10, 2019		U-440	
THALIDOMIDE; THALOMID					
020785 001	5629327	May 13, 2014	U-731	ODE	May 25, 2013
	6045501	Aug 28, 2018	U-731		
	6045501	Aug 28, 2018	U-371		
	6235756	Mar 01, 2013	U-731		
	6315720	Oct 23, 2020	U-442		
	6315720	Oct 23, 2020	U-731		
	6561976	Aug 28, 2018	U-731		
	6561976	Aug 28, 2018	U-371		
	6561977	Oct 23, 2020	U-371		
	6561977	Oct 23, 2020	U-731		
	6755784	Oct 23, 2020	U-371		
	6755784	Oct 23, 2020	U-731		
	6869399	Oct 23, 2020	U-732		
	6869399	Oct 23, 2020	U-733		
	6869399	Oct 23, 2020	U-371		
	6869399	Oct 23, 2020	U-731		
	6908432	Aug 28, 2018	U-731		
	6908432	Aug 28, 2018	U-371		
	7141018	Oct 23, 2020	U-731		
	7141018	Oct 23, 2020	U-371		
	7141018	Oct 23, 2020	U-732		
	7141018	Oct 23, 2020	U-733		

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE		PATENT CODES			EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE	
THALIDOMIDE; THALOMID									
020785 002	5629327	May	13,	2014	U-731		ODE	May	25, 2013
	6045501	Aug	28,	2018	U-371				
	6045501	Aug	28,	2018	U-731				
	6235756	Mar	01,	2013	U-731				
	6315720	Oct	23,	2020	U-731				
	6315720	Oct	23,	2020	U-442				
	6561976	Aug	28,	2018	U-731				
	6561976	Aug	28,	2018	U-371				
	6561977	Oct	23,	2020	U-371				
	6561977	Oct	23,	2020	U-731				
	6755784	Oct	23,	2020	U-371				
	6755784	Oct	23,	2020	U-731				
	6869399	Oct	23,	2020	U-371				
	6869399	Oct	23,	2020	U-732				
	6869399	Oct	23,	2020	U-731				
	6869399	Oct	23,	2020	U-733				
	6908432	Aug	28,	2018	U-371				
	6908432	Aug	28,	2018	U-731				
	7141018	Oct	23,	2020	U-731				
	7141018	Oct	23,	2020	U-371				
	7141018	Oct	23,	2020	U-733				
	7141018	Oct	23,	2020	U-732				
THALIDOMIDE; THALOMID									
020785 003	5629327	May	13,	2014	U-731		ODE	May	25, 2013
	6045501	Aug	28,	2018	U-731				
	6045501	Aug	28,	2018	U-371				
	6235756	Mar	01,	2013	U-731				
	6315720	Oct	23,	2020	U-731				
	6315720	Oct	23,	2020	U-442				
	6561976	Aug	28,	2018	U-371				
	6561976	Aug	28,	2018	U-731				
	6561977	Oct	23,	2020	U-371				
	6561977	Oct	23,	2020	U-731				
	6755784	Oct	23,	2020	U-731				
	6755784	Oct	23,	2020	U-371				
	6869399	Oct	23,	2020	U-732				
	6869399	Oct	23,	2020	U-733				
	6869399	Oct	23,	2020	U-731				
	6869399	Oct	23,	2020	U-371				
	6908432	Aug	28,	2018	U-371				
	6908432	Aug	28,	2018	U-731				
	7141018	Oct	23,	2020	U-371				
	7141018	Oct	23,	2020	U-731				
	7141018	Oct	23,	2020	U-732				
	7141018	Oct	23,	2020	U-733				
THYROTROPIN ALFA; THYROGEN									
020898 001	5840566	Nov	24,	2015	DS DP U-556		M-53	Jan	23, 2009
	6365127	Nov	24,	2015					
TIAGABINE HYDROCHLORIDE; GABITRIL									
020646 001	5010090	Sep	30,	2011					
	5354760	Mar	24,	2012					
	5866590	Apr	29,	2016					
	5958951	Jun	10,	2017					
TIAGABINE HYDROCHLORIDE; GABITRIL									
020646 002	5010090	Sep	30,	2011					
	5354760	Mar	24,	2012					
	5866590	Apr	29,	2016					
	5958951	Jun	10,	2017					
TIAGABINE HYDROCHLORIDE; GABITRIL									
020646 003	5010090	Sep	30,	2011					
	5354760	Mar	24,	2012					
	5866590	Apr	29,	2016					
	5958951	Jun	10,	2017					
TIAGABINE HYDROCHLORIDE; GABITRIL									
020646 004	5010090	Sep	30,	2011					
	5354760	Mar	24,	2012					
	5866590	Apr	29,	2016					
	5958951	Jun	10,	2017					

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>TIAGABINE HYDROCHLORIDE; GABITRIL</u>					
020646 005	5010090	Sep 30, 2011			
	5354760	Mar 24, 2012			
	5866590	Apr 29, 2016			
	5958951	Jun 10, 2017			
<u>TIGECYCLINE; TYGACIL</u>					
021821 001	5494903	Aug 13, 2012	DS DP	NCE	Jun 15, 2010
	5529990	Jun 25, 2013	U-282		
<u>TILUDRONATE DISODIUM; SKELID</u>					
020707 001	4876248	Jan 30, 2010			
	4980171	Apr 06, 2009			
<u>TIMOLOL; BETIMOL</u>					
020439 001	5231095	Jul 27, 2010			
<u>TIMOLOL; BETIMOL</u>					
020439 002	5231095	Jul 27, 2010			
<u>TIMOLOL MALEATE; TIMOLOL MALEATE</u>					
020963 001	6174524	Mar 26, 2019			
<u>TIMOLOL MALEATE; TIMOLOL MALEATE</u>					
020963 002	6174524	Mar 26, 2019			
<u>TIMOLOL MALEATE; TIMOPTIC-XE</u>					
020330 001	4861760	Sep 25, 2006			
<u>TIMOLOL MALEATE; TIMOPTIC-XE</u>					
020330 002	4861760	Sep 25, 2006			
<u>TINIDAZOLE; TINDAMAX</u>					
021618 001				NCE	May 17, 2009
				ODE	May 17, 2011
				ODE	May 17, 2011
<u>TINIDAZOLE; TINDAMAX</u>					
021618 002				NCE	May 17, 2009
				ODE	May 17, 2011
				ODE	May 17, 2011
<u>TIOTROPIUM BROMIDE MONOHYDRATE; SPIRIVA</u>					
021395 001	5610163	Mar 11, 2014	DS DP	U-566	Jan 30, 2009
	6777423	Sep 24, 2021	DS DP		
	6908928	Nov 25, 2022	DS DP	U-566	
	7070800	Jan 22, 2022	DP	U-566	
	RE38912	Oct 11, 2021	DP		
<u>TIPRANAVIR; APTIVUS</u>					
021814 001	5852195	Dec 22, 2015	DS		Jun 22, 2010
	6147095	Oct 29, 2019		U-670	
	6169181	May 06, 2014	DS		
	6231887	Jul 27, 2018	DP		
<u>TIROFIBAN HYDROCHLORIDE; AGGRASTAT</u>					
020912 001	5292756	May 14, 2012		U-230	
	5658929	Sep 27, 2010			
	5733919	Oct 23, 2016			
	5880136	Sep 27, 2010		U-254	
	5965581	Oct 23, 2016			
	5972967	Oct 23, 2016			
	5978698	Oct 08, 2017			
	6136794	Jan 29, 2019			
<u>TIROFIBAN HYDROCHLORIDE; AGGRASTAT</u>					
020913 001	5292756	May 14, 2012		U-230	
	5658929	Sep 27, 2010			
	5733919	Oct 23, 2016			
	5880136	Sep 27, 2010		U-254	
	5965581	Oct 23, 2016			
	5972967	Oct 23, 2016			
	5978698	Oct 08, 2017			
	6136794	Jan 29, 2019			

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>TIZANIDINE HYDROCHLORIDE; ZANAFLEX</u>					
021447 001	6455557	Nov 28, 2021			
<u>TIZANIDINE HYDROCHLORIDE; ZANAFLEX</u>					
021447 002	6455557	Nov 28, 2021			
<u>TIZANIDINE HYDROCHLORIDE; ZANAFLEX</u>					
021447 003	6455557	Nov 28, 2021			
<u>TOLCAPONE; TASMAR</u>					
020697 001	5236952	Jan 29, 2012			
	5476875	Dec 19, 2012		U-219	
<u>TOLCAPONE; TASMAR</u>					
020697 002	5236952	Jan 29, 2012			
	5476875	Dec 19, 2012		U-219	
<u>TOLTERODINE TARTRATE; DETROL</u>					
020771 001	5382600	Mar 25, 2012			
	5382600*PED	Sep 25, 2012			
<u>TOLTERODINE TARTRATE; DETROL</u>					
020771 002	5382600	Mar 25, 2012			
	5382600*PED	Sep 25, 2012			
<u>TOLTERODINE TARTRATE; DETROL LA</u>					
021228 001	5382600	Mar 25, 2012		M-46	Oct 21, 2008
	5382600*PED	Sep 25, 2012			
	6630162	Nov 11, 2019	DP	U-544	
	6630162*PED	May 11, 2020			
	6770295	Aug 26, 2019	DP	U-544	
	6770295*PED	Feb 26, 2020			
	6911217	Aug 26, 2019	DP	U-544	
	6911217*PED	Feb 26, 2020	DP	U-544	
<u>TOLTERODINE TARTRATE; DETROL LA</u>					
021228 002	5382600	Mar 25, 2012		M-46	Oct 21, 2008
	5382600*PED	Sep 25, 2012			
	6630162	Nov 11, 2019	DP	U-544	
	6630162*PED	May 11, 2020			
	6770295	Aug 26, 2019	DP	U-544	
	6770295*PED	Feb 26, 2020			
	6911217	Aug 26, 2019	DP	U-544	
	6911217*PED	Feb 26, 2020	DP	U-544	
<u>TOPIRAMATE; TOPAMAX</u>					
020505 001	4513006	Sep 26, 2008	DS	DP	U-83
	5998380	Oct 13, 2015			U-598
	6503884	Oct 13, 2015			U-598
	7018983	Oct 13, 2015			U-723
				D-88	Dec 16, 2006
				I-467	Jun 29, 2008
				I-41	Aug 11, 2007
				ODE	Aug 28, 2008
<u>TOPIRAMATE; TOPAMAX</u>					
020505 002	4513006	Sep 26, 2008	DS	DP	U-83
	5998380	Oct 13, 2015			U-598
	6503884	Oct 13, 2015			U-598
	7018983	Oct 13, 2015			U-723
				D-88	Dec 16, 2006
				I-467	Jun 29, 2008
				I-41	Aug 11, 2007
				ODE	Aug 28, 2008
<u>TOPIRAMATE; TOPAMAX</u>					
020505 003	4513006	Sep 26, 2008	DS	DP	U-83
	5998380	Oct 13, 2015			U-598
	6503884	Oct 13, 2015			U-598
	7018983	Oct 13, 2015			U-723
				D-88	Dec 16, 2006
				I-467	Jun 29, 2008
				I-41	Aug 11, 2007
				ODE	Aug 28, 2008
<u>TOPIRAMATE; TOPAMAX</u>					
020505 004	4513006	Sep 26, 2008	DS	DP	U-83
	5998380	Oct 13, 2015			U-598
	6503884	Oct 13, 2015			U-598
	7018983	Oct 13, 2015			U-723
				D-88	Dec 16, 2006
				I-467	Jun 29, 2008
				I-41	Aug 11, 2007
				ODE	Aug 28, 2008

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE		PATENT CODES		EXCLUSIVITY CODE(S)		EXCLUSIVITY EXPIRATION DATE	
TOPIRAMATE; TOPAMAX									
020505 005	4513006	Sep 26, 2008	DS	DP	U-83	D-88	Dec 16, 2006		
	5998380	Oct 13, 2015			U-598	I-467	Jun 29, 2008		
	6503884	Oct 13, 2015			U-598	I-41	Aug 11, 2007		
	7018983	Oct 13, 2015			U-723	ODE	Aug 28, 2008		
TOPIRAMATE; TOPAMAX									
020505 006	4513006	Sep 26, 2008	DS	DP	U-83	D-88	Dec 16, 2006		
	5998380	Oct 13, 2015			U-598	I-467	Jun 29, 2008		
	6503884	Oct 13, 2015			U-598	I-41	Aug 11, 2007		
	7018983	Oct 13, 2015			U-723	ODE	Aug 28, 2008		
TOPIRAMATE; TOPAMAX SPRINKLE									
020844 001	4513006	Sep 26, 2008	DS	DP	U-83	D-88	Dec 16, 2006		
	5998380	Oct 13, 2015			U-598	I-467	Jun 29, 2008		
	6503884	Oct 13, 2015			U-598	I-41	Aug 11, 2007		
	7018983	Oct 13, 2015			U-723	ODE	Aug 28, 2008		
	7125560	Mar 01, 2019			U-766				
TOPIRAMATE; TOPAMAX SPRINKLE									
020844 002	4513006	Sep 26, 2008	DS	DP	U-83	D-88	Dec 16, 2006		
	5998380	Oct 13, 2015			U-598	I-467	Jun 29, 2008		
	6503884	Oct 13, 2015			U-598	I-41	Aug 11, 2007		
	7018983	Oct 13, 2015			U-723	ODE	Aug 28, 2008		
	7125560	Mar 01, 2019			U-766				
TOPIRAMATE; TOPAMAX SPRINKLE									
020844 003	4513006	Sep 26, 2008	DS	DP	U-83	D-88	Dec 16, 2006		
	5998380	Oct 13, 2015			U-598	I-467	Jun 29, 2008		
	6503884	Oct 13, 2015			U-598	I-41	Aug 11, 2007		
	7018983	Oct 13, 2015			U-723	ODE	Aug 28, 2008		
	7125560	Mar 01, 2019			U-766				
TOPOTECAN HYDROCHLORIDE; HYCAMTIN									
020671 001	5004758	May 28, 2010	DS	DP	U-741				
	5004758*PED	Nov 28, 2010							
TOREMIFENE CITRATE; FARESTON									
020497 001	4696949	Sep 29, 2009			U-196				
TORSEMIDE; DEMADEX									
020136 001	RE34672	Aug 11, 2006							
TORSEMIDE; DEMADEX									
020136 002	RE34672	Aug 11, 2006							
TORSEMIDE; DEMADEX									
020136 003	RE34672	Aug 11, 2006							
TORSEMIDE; DEMADEX									
020136 004	RE34672	Aug 11, 2006							
TORSEMIDE; DEMADEX									
020137 002	4861786	Jul 08, 2007							
	RE34672	Aug 11, 2006							
TRAMADOL HYDROCHLORIDE; TRAMADOL HYDROCHLORIDE									
021692 001	6254887	May 10, 2014		DP		NP	Sep 08, 2008		
TRAMADOL HYDROCHLORIDE; TRAMADOL HYDROCHLORIDE									
021692 002	6254887	May 10, 2014		DP		NP	Sep 08, 2008		
TRAMADOL HYDROCHLORIDE; TRAMADOL HYDROCHLORIDE									
021692 003	6254887	May 10, 2014		DP		NP	Sep 08, 2008		
TRAMADOL HYDROCHLORIDE; TRAMADOL HYDROCHLORIDE									
021693 001	5464632	Mar 22, 2013		DP					
	6106861	Dec 05, 2017							
TRAMADOL HYDROCHLORIDE; ULTRAM									
020281 001	6339105	Oct 12, 2019			U-435				
	6339105*PED	Apr 12, 2020			U-435				
TRAMADOL HYDROCHLORIDE; ULTRAM									
020281 002	6339105	Oct 12, 2019			U-435				
	6339105*PED	Apr 12, 2020			U-435				

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>TRANDOLAPRIL; MAVIK</u>					
020528 001	4933361 5744496	Jun 12, 2007 Apr 28, 2015		U-229	
<u>TRANDOLAPRIL; MAVIK</u>					
020528 002	4933361 5744496	Jun 12, 2007 Apr 28, 2015		U-229	
<u>TRANDOLAPRIL; MAVIK</u>					
020528 003	4933361 5744496	Jun 12, 2007 Apr 28, 2015		U-229	
<u>TRANDOLAPRIL; VERAPAMIL HYDROCHLORIDE; TARKA</u>					
020591 001	4933361 5721244	Jun 12, 2007 Feb 24, 2015			
<u>TRANDOLAPRIL; VERAPAMIL HYDROCHLORIDE; TARKA</u>					
020591 002	4933361 5721244	Jun 12, 2007 Feb 24, 2015			
<u>TRANDOLAPRIL; VERAPAMIL HYDROCHLORIDE; TARKA</u>					
020591 003	4933361 5721244	Jun 12, 2007 Feb 24, 2015			
<u>TRANDOLAPRIL; VERAPAMIL HYDROCHLORIDE; TARKA</u>					
020591 004	4933361 5721244	Jun 12, 2007 Feb 24, 2015			
<u>TRAVOPROST; TRAVATAN</u>					
021257 001	5510383 5631287 5849792 5889052 6011062	Aug 03, 2013 Dec 22, 2014 Dec 22, 2014 Dec 02, 2014 Dec 22, 2014	DP DP DP DP	U-383 U-382 U-383 U-383	
<u>TRAVOPROST; TRAVATAN Z</u>					
021994 001	5510383 5889052 6503497 6849253	Aug 03, 2013 Dec 02, 2014 May 06, 2012 May 06, 2012	DP DP DP DP	U-383 U-383	
<u>TREPROSTINIL SODIUM; REMODULIN</u>					
021272 001	5153222 6765117	Oct 06, 2014 Oct 24, 2017	DS	U-455 NCE ODE	May 21, 2007 May 21, 2009
<u>TREPROSTINIL SODIUM; REMODULIN</u>					
021272 002	5153222 6765117	Oct 06, 2014 Oct 24, 2017	DS	U-455 NCE ODE	May 21, 2007 May 21, 2009
<u>TREPROSTINIL SODIUM; REMODULIN</u>					
021272 003	5153222 6765117	Oct 06, 2014 Oct 24, 2017	DS	U-455 NCE ODE	May 21, 2007 May 21, 2009
<u>TREPROSTINIL SODIUM; REMODULIN</u>					
021272 004	5153222 6765117	Oct 06, 2014 Oct 24, 2017	DS	U-455 NCE ODE	May 21, 2007 May 21, 2009
<u>TRETINOIN; AVITA</u>					
020404 003	4971800 5045317	Nov 20, 2007 Sep 03, 2008		U-178 U-179	
<u>TRETINOIN; RENOVA</u>					
021108 001	6531141	Mar 07, 2020			
<u>TRETINOIN; RETIN-A MICRO</u>					
020475 001	5955109	Sep 21, 2016	DP	U-134	
<u>TRETINOIN; RETIN-A MICRO</u>					
020475 002	5955109	Sep 21, 2016		U-134	
<u>TRIAMCINOLONE ACETONIDE; NASACORT</u>					
019798 001	4767612	Jan 23, 2007		U-85	
<u>TRIAMCINOLONE ACETONIDE; NASACORT AQ</u>					
020468 001	5976573 6143329	Jul 03, 2016 Jul 03, 2016		U-295	

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>TRIMETHOPRIM HYDROCHLORIDE; PRIMSO</u>					
074973 001	5763449	Aug 07, 2016			
	5962461	Aug 07, 2016			
<u>TRIMETREXATE GLUCURONATE; NEUTREXIN</u>					
020326 001	6017922	May 18, 2018			
<u>TRIMETREXATE GLUCURONATE; NEUTREXIN</u>					
020326 002	6017922	May 18, 2018			
<u>TRIPTORELIN PAMOATE; TRELSTAR</u>					
021288 001	5225205	Jul 20, 2010			
<u>TRIPTORELIN PAMOATE; TRELSTAR DEPOT</u>					
020715 001	5225205	Jul 20, 2010			
	5776885	Jul 07, 2015			
<u>TROGLITAZONE; PRELAY</u>					
020719 001	4572912	Nov 09, 2008			
	5602133	Sep 15, 2013		U-173	
	5859037	Nov 13, 2017		U-251	
	6011049	Nov 13, 2017		U-301	
	6046202	Sep 15, 2013		U-317	
<u>TROGLITAZONE; PRELAY</u>					
020719 002	4572912	Nov 09, 2008			
	5602133	Sep 15, 2013		U-173	
	5859037	Nov 13, 2017		U-251	
	6011049	Nov 13, 2017		U-301	
	6046202	Sep 15, 2013		U-317	
<u>TROGLITAZONE; PRELAY</u>					
020719 003	4572912	Nov 09, 2008			
	5602133	Sep 15, 2013		U-173	
	5859037	Nov 13, 2017		U-251	
	6011049	Nov 13, 2017		U-301	
	6046202	Sep 15, 2013		U-317	
<u>TROGLITAZONE; REZULIN</u>					
020720 001	4572912	Nov 09, 2008			
	5602133	Sep 15, 2013		U-173	
	5859037	Nov 13, 2017		U-251	
	6011049	Nov 13, 2017		U-301	
	6046202	Sep 15, 2013		U-317	
<u>TROGLITAZONE; REZULIN</u>					
020720 002	4572912	Nov 09, 2008			
	5602133	Sep 15, 2013		U-173	
	5859037	Nov 13, 2017		U-251	
	6011049	Nov 13, 2017		U-301	
	6046202	Sep 15, 2013		U-317	
<u>TROGLITAZONE; REZULIN</u>					
020720 003	4572912	Nov 09, 2008			
	5602133	Sep 15, 2013		U-173	
	5859037	Nov 13, 2017		U-251	
	6011049	Nov 13, 2017		U-301	
	6046202	Sep 15, 2013		U-317	
<u>TROSPIUM CHLORIDE; SANCTURA</u>					
021595 001				NCE	May 28, 2009
<u>TROVAFLOXACIN MESYLATE; TROVAN</u>					
020759 001	5164402	Dec 18, 2011		U-282	
	5763454	Jun 15, 2015		U-282	
	6187341	Jan 20, 2019			
<u>TROVAFLOXACIN MESYLATE; TROVAN</u>					
020759 002	5164402	Dec 18, 2011		U-282	
	5763454	Jun 15, 2015		U-282	
	6187341	Jan 20, 2019			
<u>TRYPAN BLUE; VISIONBLUE</u>					
021670 001				NCE	Dec 16, 2009

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>UNOPROSTONE ISOPROPYL; RESCULA</u>					
021214 001	5001153	Sep 19, 2008			
	5151444	Mar 19, 2008	U-333		
	5166178	Nov 24, 2009	U-333		
	5208256	May 21, 2011	U-333		
	5212200	May 18, 2010	U-333		
	5221763	Jun 22, 2010			
<u>UREA, C-13; BREATHTEK UBT FOR H-PYLORI</u>					
020586 002	4830010	Oct 27, 2009	U-147		
	5140993	Aug 24, 2009	U-148		
<u>UREA, C-13; MERETEK UBT KIT (W/ PRANACTIN)</u>					
020586 001	4830010	Oct 27, 2009	U-147		
	5140993	Aug 24, 2009	U-148		
<u>UROFOLLITROPIN; FERTINEX</u>					
019415 004	4845077	Jul 04, 2006	U-408		
	5767067	Jun 16, 2015			
<u>UROFOLLITROPIN; FERTINEX</u>					
019415 005	4845077	Jul 04, 2006	U-408		
	5767067	Jun 16, 2015			
<u>URSODIOL; URSO 250</u>					
020675 001	4859660	Nov 19, 2007		D-93	Jul 21, 2007
<u>URSODIOL; URSO FORTE</u>					
020675 002	4859660	Nov 19, 2007	U-740	D-93	Jul 21, 2007
<u>VALACYCLOVIR HYDROCHLORIDE; VALTREX</u>					
020487 001	4957924	Jun 23, 2009	U-530	I-403	Aug 29, 2006
	5879706	Jan 19, 2016	U-530		
	6107302	Jan 19, 2016	U-530		
<u>VALACYCLOVIR HYDROCHLORIDE; VALTREX</u>					
020487 002	4957924	Jun 23, 2009	U-530	I-403	Aug 29, 2006
	5879706	Jan 19, 2016	U-530		
	6107302	Jan 19, 2016	U-530		
<u>VALDECOXIB; BEXTRA</u>					
021341 002	5633272	Feb 13, 2015	U-462	NCE	Nov 16, 2006
	7135489	Aug 12, 2017	DS DP		
<u>VALDECOXIB; BEXTRA</u>					
021341 003	5633272	Feb 13, 2015	U-462	NCE	Nov 16, 2006
	7135489	Aug 12, 2017	DS DP		
<u>VALGANCICLOVIR HYDROCHLORIDE; VALCYTE</u>					
021304 001	6083953	Jul 28, 2014	U-384	I-406	Sep 12, 2006
<u>VALSARTAN; DIOVAN</u>					
020665 001	5399578	Mar 21, 2012	U-3		
<u>VALSARTAN; DIOVAN</u>					
020665 002	5399578	Mar 21, 2012	U-3		
<u>VALSARTAN; DIOVAN</u>					
021283 001	5399578	Mar 21, 2012		I-463	Aug 03, 2008
	5972990	Oct 26, 2016	U-692		
	6294197	Jun 18, 2017	U-3		
<u>VALSARTAN; DIOVAN</u>					
021283 002	5399578	Mar 21, 2012		I-463	Aug 03, 2008
	5972990	Oct 26, 2016	U-692		
	6294197	Jun 18, 2017	U-3		
<u>VALSARTAN; DIOVAN</u>					
021283 003	5399578	Mar 21, 2012		I-463	Aug 03, 2008
	5972990	Oct 26, 2016	U-692		
	6294197	Jun 18, 2017	U-3		
<u>VALSARTAN; DIOVAN</u>					
021283 004	5399578	Mar 21, 2012		I-463	Aug 03, 2008
	5972990	Oct 26, 2016	U-692		
	6294197	Jun 18, 2017	U-3		

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE		PATENT CODES			EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE	
<u>VARDENAFIL HYDROCHLORIDE; LEVITRA</u>									
021400 001	6362178	Oct	31, 2018	DS	DP	U-533	NCE	Aug	19, 2008
<u>VARDENAFIL HYDROCHLORIDE; LEVITRA</u>									
021400 002	6362178	Oct	31, 2018	DS	DP	U-533	NCE	Aug	19, 2008
<u>VARDENAFIL HYDROCHLORIDE; LEVITRA</u>									
021400 003	6362178	Oct	31, 2018	DS	DP	U-533	NCE	Aug	19, 2008
<u>VARDENAFIL HYDROCHLORIDE; LEVITRA</u>									
021400 004	6362178	Oct	31, 2018	DS	DP	U-533	NCE	Aug	19, 2008
<u>VARENICLINE TARTRATE; CHANTIX</u>									
021928 001	6410550	Nov	13, 2018	DS	DP	U-56	NCE	May	10, 2011
	6890927	May	06, 2022	DS	DP	U-56			
<u>VARENICLINE TARTRATE; CHANTIX</u>									
021928 002	6410550	Nov	13, 2018	DS	DP	U-56	NCE	May	10, 2011
	6890927	May	06, 2022	DS	DP	U-56			
<u>VENLAFAXINE HYDROCHLORIDE; EFFEXOR</u>									
020151 001	4535186	Dec	13, 2007						
	4535186*PED	Jun	13, 2008						
<u>VENLAFAXINE HYDROCHLORIDE; EFFEXOR</u>									
020151 002	4535186	Dec	13, 2007						
	4535186*PED	Jun	13, 2008						
<u>VENLAFAXINE HYDROCHLORIDE; EFFEXOR</u>									
020151 003	4535186	Dec	13, 2007						
	4535186*PED	Jun	13, 2008						
<u>VENLAFAXINE HYDROCHLORIDE; EFFEXOR</u>									
020151 004	4535186	Dec	13, 2007						
	4535186*PED	Jun	13, 2008						
<u>VENLAFAXINE HYDROCHLORIDE; EFFEXOR</u>									
020151 005	4535186	Dec	13, 2007						
	4535186*PED	Jun	13, 2008						
<u>VENLAFAXINE HYDROCHLORIDE; EFFEXOR</u>									
020151 006	4535186	Dec	13, 2007						
	4535186*PED	Jun	13, 2008						
<u>VENLAFAXINE HYDROCHLORIDE; EFFEXOR XR</u>									
020699 001	4535186	Dec	13, 2007						
	4535186*PED	Jun	13, 2008						
	5916923	Jun	28, 2013						
	5916923*PED	Dec	28, 2013						
	6274171	Mar	20, 2017						
	6274171*PED	Sep	20, 2017						
	6403120	Mar	20, 2017						
	6403120	Mar	20, 2017						
	6403120*PED	Sep	20, 2017						
	6403120*PED	Sep	20, 2017						
	6419958	Mar	20, 2017						
	6419958	Mar	20, 2017						
	6419958*PED	Sep	20, 2017						
	6419958*PED	Sep	20, 2017						
	6444708	Jun	28, 2013						
	6444708*PED	Dec	28, 2013						

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>VENLAFAXINE HYDROCHLORIDE; EFFEXOR XR</u>					
020699 002	4535186	Dec 13, 2007			
	4535186*PED	Jun 13, 2008			
	5916923	Jun 28, 2013		U-398	
	5916923*PED	Dec 28, 2013		U-398	
	6274171	Mar 20, 2017			
	6274171*PED	Sep 20, 2017			
	6403120	Mar 20, 2017		U-451	
	6403120	Mar 20, 2017		U-535	
	6403120*PED	Sep 20, 2017		U-535	
	6403120*PED	Sep 20, 2017		U-451	
	6419958	Mar 20, 2017		U-535	
	6419958	Mar 20, 2017		U-459	
	6419958*PED	Sep 20, 2017		U-535	
	6419958*PED	Sep 20, 2017		U-459	
	6444708	Jun 28, 2013		U-398	
	6444708*PED	Dec 28, 2013		U-398	
<u>VENLAFAXINE HYDROCHLORIDE; EFFEXOR XR</u>					
020699 003	4535186	Dec 13, 2007			
	4535186*PED	Jun 13, 2008			
	5916923	Jun 28, 2013		U-398	
	5916923*PED	Dec 28, 2013		U-398	
	6274171	Mar 20, 2017			
	6274171*PED	Sep 20, 2017			
	6403120	Mar 20, 2017		U-451	
	6403120	Mar 20, 2017		U-535	
	6403120*PED	Sep 20, 2017		U-451	
	6403120*PED	Sep 20, 2017		U-535	
	6419958	Mar 20, 2017		U-459	
	6419958	Mar 20, 2017		U-535	
	6419958*PED	Sep 20, 2017		U-459	
	6419958*PED	Sep 20, 2017		U-535	
	6444708	Jun 28, 2013		U-398	
	6444708*PED	Dec 28, 2013		U-398	
<u>VENLAFAXINE HYDROCHLORIDE; EFFEXOR XR</u>					
020699 004	4535186	Dec 13, 2007			
	4535186*PED	Jun 13, 2008			
	5916923	Jun 28, 2013		U-398	
	5916923*PED	Dec 28, 2013		U-398	
	6274171	Mar 20, 2017			
	6274171*PED	Sep 20, 2017			
	6403120	Mar 20, 2017		U-451	
	6403120	Mar 20, 2017		U-535	
	6403120*PED	Sep 20, 2017		U-535	
	6403120*PED	Sep 20, 2017		U-451	
	6419958	Mar 20, 2017		U-535	
	6419958	Mar 20, 2017		U-459	
	6419958*PED	Sep 20, 2017		U-459	
	6419958*PED	Sep 20, 2017		U-535	
	6444708	Jun 28, 2013		U-398	
	6444708*PED	Dec 28, 2013		U-398	
<u>VENLAFAXINE HYDROCHLORIDE; VENLAFAXINE HYDROCHLORIDE</u>					
076690 001				PC	Jan 30, 2007
<u>VENLAFAXINE HYDROCHLORIDE; VENLAFAXINE HYDROCHLORIDE</u>					
076690 002				PC	Jan 30, 2007
<u>VENLAFAXINE HYDROCHLORIDE; VENLAFAXINE HYDROCHLORIDE</u>					
076690 003				PC	Jan 30, 2007
<u>VENLAFAXINE HYDROCHLORIDE; VENLAFAXINE HYDROCHLORIDE</u>					
076690 004				PC	Jan 30, 2007
<u>VENLAFAXINE HYDROCHLORIDE; VENLAFAXINE HYDROCHLORIDE</u>					
076690 005				PC	Jan 30, 2007

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>VERAPAMIL HYDROCHLORIDE; COVERA-HS</u>					
020552 001	4946687	Oct 02, 2007			
	5030456	Nov 07, 2008			
	5082668	Jan 21, 2009			
	5160744	Jun 27, 2011			
	5190765	Aug 14, 2007			
	5200196	Jan 22, 2008			
	5232705	Aug 31, 2010			
	5252338	Jun 27, 2011			
	5785994	Oct 22, 2009		U-315	
	6096339	Apr 04, 2017		U-365	
	6146662	Aug 14, 2007		U-366	
<u>VERAPAMIL HYDROCHLORIDE; COVERA-HS</u>					
020552 002	4946687	Oct 02, 2007			
	5030456	Nov 07, 2008			
	5082668	Jan 21, 2009			
	5160744	Jun 27, 2011			
	5190765	Aug 14, 2007			
	5200196	Jan 22, 2008			
	5232705	Aug 31, 2010			
	5252338	Jun 27, 2011			
	5785994	Oct 22, 2009		U-315	
	6096339	Apr 04, 2017		U-365	
	6146662	Aug 14, 2007		U-366	
<u>VERAPAMIL HYDROCHLORIDE; VERELAN</u>					
019614 001	4863742	Jun 19, 2007		U-3	
<u>VERAPAMIL HYDROCHLORIDE; VERELAN</u>					
019614 002	4863742	Jun 19, 2007		U-3	
<u>VERAPAMIL HYDROCHLORIDE; VERELAN</u>					
019614 003	4863742	Jun 19, 2007		U-3	
<u>VERAPAMIL HYDROCHLORIDE; VERELAN</u>					
019614 004	4863742	Jun 19, 2007			
<u>VERAPAMIL HYDROCHLORIDE; VERELAN PM</u>					
020943 001	4863742	Jun 19, 2007			
<u>VERAPAMIL HYDROCHLORIDE; VERELAN PM</u>					
020943 002	4863742	Jun 19, 2007			
<u>VERAPAMIL HYDROCHLORIDE; VERELAN PM</u>					
020943 003	4863742	Jun 19, 2007			
<u>VERTEPORFIN; VISUDYNE</u>					
021119 001	4883790	Jan 20, 2007			
	4920143	Apr 24, 2007			
	5095030	Sep 09, 2011	DS		
	5214036	May 25, 2010			
	5283255	Jan 20, 2007		U-357	
	5707608	Aug 02, 2015			
	5756541	Mar 11, 2016		U-357	
	5770619	Jan 06, 2015		U-357	
	5798349	Aug 25, 2015		U-357	
	6074666	Feb 05, 2012			
<u>VORICONAZOLE; VFEND</u>					
021266 001	5116844	Aug 11, 2009		DP U-540	I-442 Dec 21, 2007
	5364938	Nov 15, 2011	DS		I-409 Nov 14, 2006
	5567817	May 24, 2016	DS DP	U-540	NCE May 24, 2007
	5773443	Jan 25, 2011	DS DP	U-540	
<u>VORICONAZOLE; VFEND</u>					
021266 002	5116844	Aug 11, 2009		DP U-540	I-442 Dec 21, 2007
	5364938	Nov 15, 2011	DS		I-409 Nov 14, 2006
	5567817	May 24, 2016	DS DP	U-540	NCE May 24, 2007
	5773443	Jan 25, 2011	DS DP	U-540	

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE		PATENT CODES			EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE	
VORICONAZOLE; VFEND									
021267 001	5116844	Aug 11, 2009		DP	U-540	I-442	Dec 21, 2007		
	5134127	Jan 23, 2010		DP		I-409	Nov 14, 2006		
	5364938	Nov 15, 2011	DS			NCE	May 24, 2007		
	5376645	Jan 23, 2010		DP					
	5567817	May 24, 2016	DS	DP	U-540				
	5773443	Jan 25, 2011	DS	DP	U-540				
	6632803	Jun 02, 2018		DP					
VORICONAZOLE; VFEND									
021630 001	5116844	Aug 11, 2009		DP	U-540	I-442	Dec 21, 2007		
	5364938	Nov 15, 2011	DS			I-409	Nov 14, 2006		
	5567817	May 24, 2016	DS	DP	U-540	NCE	May 24, 2007		
	5773443	Jan 25, 2011	DS	DP	U-540				
VORINOSTAT; ZOLINZA									
021991 001	6087367	Oct 04, 2011			U-776	NCE	Oct 06, 2011		
	RE38506	Nov 29, 2011	DS	DP		ODE	Oct 06, 2013		
ZAFIRLUKAST; ACCOLATE									
020547 001	4859692	Sep 26, 2010							
	5294636	Dec 11, 2011							
	5319097	Dec 11, 2011							
	5482963	Jan 09, 2013							
	5583152	Sep 26, 2010							
	5612367	Mar 18, 2014			U-189				
	6143775	Dec 11, 2011							
ZAFIRLUKAST; ACCOLATE									
020547 003	4859692	Sep 26, 2010							
	5294636	Dec 11, 2011							
	5319097	Dec 11, 2011							
	5482963	Jan 09, 2013							
	5583152	Sep 26, 2010							
	5612367	Mar 18, 2014			U-189				
	6143775	Dec 11, 2011							
ZALCITABINE; HIVID									
020199 001	4879277	Nov 07, 2006			U-65				
	5028595	Jul 02, 2008			U-65				
ZALCITABINE; HIVID									
020199 002	4879277	Nov 07, 2006			U-65				
	5028595	Jul 02, 2008			U-65				
ZALEPLON; SONATA									
020859 001	4626538	Jun 06, 2008							
ZALEPLON; SONATA									
020859 002	4626538	Jun 06, 2008							
ZANAMIVIR; RELENZA									
021036 001	5360817	Jul 26, 2013	DS	DP		I-491	Mar 29, 2009		
	5648379	Jul 15, 2014			U-721				
	5648379	Jul 15, 2014			U-722				
	5648379	Jul 15, 2014			U-274				
	6294572	Dec 15, 2014	DS	DP					
ZICONOTIDE; PRIALT									
021060 001	5364842	Dec 30, 2011			U-55	NCE	Dec 28, 2009		
	5364842	Dec 30, 2011			U-48				
	5795864	Jun 27, 2015		DP					
	5859186	Dec 30, 2011			U-55				
	5859186	Dec 30, 2011			U-48				
ZICONOTIDE; PRIALT									
021060 002	5364842	Dec 30, 2011			U-48	NCE	Dec 28, 2009		
	5364842	Dec 30, 2011			U-55				
	5795864	Jun 27, 2015		DP					
	5859186	Dec 30, 2011			U-55				
	5859186	Dec 30, 2011			U-48				

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE		PATENT CODES		EXCLUSIVITY CODE(S)		EXCLUSIVITY EXPIRATION DATE	
<u>ZICONOTIDE; PRIALT</u>									
021060 003	5364842	Dec 30, 2011	DP	U-55 U-48	NCE		Dec 28, 2009		
	5364842	Dec 30, 2011							
	5795864	Jun 27, 2015							
	5859186	Dec 30, 2011							
	5859186	Dec 30, 2011							
<u>ZICONOTIDE; PRIALT</u>									
021060 004	5364842	Dec 30, 2011	DP	U-48 U-55 U-55 U-48	NCE		Dec 28, 2009		
	5364842	Dec 30, 2011							
	5795864	Jun 27, 2015							
	5859186	Dec 30, 2011							
	5859186	Dec 30, 2011							
<u>ZILEUTON; ZYFLO</u>									
020471 001	4873259	Dec 10, 2010		U-168					
<u>ZILEUTON; ZYFLO</u>									
020471 003	4873259	Dec 10, 2010		U-168					
<u>ZIPRASIDONE HYDROCHLORIDE; GEODON</u>									
020825 001	4831031	Mar 02, 2012	DS DS	DP DP DP	I-492		Aug 19, 2007		
	5312925	Sep 01, 2012							
	6150366	May 27, 2019							
	6245766	Dec 18, 2018							
<u>ZIPRASIDONE HYDROCHLORIDE; GEODON</u>									
020825 002	4831031	Mar 02, 2012	DS DS	DP DP DP	I-492		Aug 19, 2007		
	5312925	Sep 01, 2012							
	6150366	May 27, 2019							
	6245766	Dec 18, 2018							
<u>ZIPRASIDONE HYDROCHLORIDE; GEODON</u>									
020825 003	4831031	Mar 02, 2012	DS DS	DP DP DP	I-492		Aug 19, 2007		
	5312925	Sep 01, 2012							
	6150366	May 27, 2019							
	6245766	Dec 18, 2018							
<u>ZIPRASIDONE HYDROCHLORIDE; GEODON</u>									
020825 004	4831031	Mar 02, 2012	DS DS	DP DP DP	I-492		Aug 19, 2007		
	5312925	Sep 01, 2012							
	6150366	May 27, 2019							
	6245766	Dec 18, 2018							
<u>ZIPRASIDONE HYDROCHLORIDE; GEODON</u>									
021483 001	4831031	Mar 02, 2012	DS DS	DP DP DP	I-492		Aug 19, 2007		
	5312925	Sep 01, 2012							
	6150366	May 27, 2019							
	6245766	Dec 18, 2018							
<u>ZIPRASIDONE MESYLATE; GEODON</u>									
020919 001	4831031	Mar 02, 2012	DS	DP	U-720				
	6110918	Mar 26, 2017							
	6232304	Apr 01, 2017							
	6399777	Apr 01, 2017							
<u>ZOLEDRONIC ACID; ZOMETA</u>									
021223 001	4777163	Jul 24, 2007	DS	DP	U-53	NCE ODE	Aug 20, 2006 Aug 20, 2008		
	4939130	Sep 02, 2012							
<u>ZOLEDRONIC ACID; ZOMETA</u>									
021223 002	4777163	Jul 24, 2007	DS	DP	U-53	NCE ODE	Aug 20, 2006 Aug 20, 2008		
	4939130	Sep 02, 2012							
<u>ZOLMITRIPTAN; ZOMIG</u>									
020768 001	5466699	Nov 14, 2012							
	5466699*PED	May 14, 2013							
	5863935	Nov 14, 2012							
	5863935*PED	May 14, 2013							
<u>ZOLMITRIPTAN; ZOMIG</u>									
020768 002	5466699	Nov 14, 2012							
	5466699*PED	May 14, 2013							
	5863935	Nov 14, 2012							
	5863935*PED	May 14, 2013							

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE		PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE	
<u>ZOLMITRIPTAN; ZOMIG</u>							
021450 004	5466699	Nov 14, 2012	U-436	NDF	Sep 26, 2006		
	5466699*PED	May 14, 2013					
	6750237	Nov 28, 2020					
	6750237*PED	May 28, 2021					
<u>ZOLMITRIPTAN; ZOMIG-ZMT</u>							
021231 001	5466699	Nov 14, 2012					
	5466699*PED	May 14, 2013					
<u>ZOLMITRIPTAN; ZOMIG-ZMT</u>							
021231 002	5466699	Nov 14, 2012					
	5466699*PED	May 14, 2013					
<u>ZOLPIDEM TARTRATE; AMBIEN</u>							
019908 001	4382938	Oct 21, 2006	U-74				
	4382938*PED	Apr 21, 2007					
<u>ZOLPIDEM TARTRATE; AMBIEN</u>							
019908 002	4382938	Oct 21, 2006	U-74				
	4382938*PED	Apr 21, 2007					
<u>ZOLPIDEM TARTRATE; AMBIEN CR</u>							
021774 001	4382938	Oct 21, 2006	DS	U-682	NDF	Sep 02, 2008	
	4382938*PED	Apr 21, 2007					
	6514531	Dec 01, 2019					DP
	6514531*PED	Jun 01, 2020					
<u>ZOLPIDEM TARTRATE; AMBIEN CR</u>							
021774 002	4382938	Oct 21, 2006	DS	U-682	NDF	Sep 02, 2008	
	4382938*PED	Apr 21, 2007					
	6514531	Dec 01, 2019					DP
	6514531*PED	Jun 01, 2020					

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnote for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
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Footnote:

1. Patents are published upon receipt by the Orange Book Staff and may not reflect the official receipt date as described in 21 CFR 314.53(d)(5).
2. Patents submitted on FDA Form 3542 and listed after August 18, 2003 will have one to three patent codes indicating specific patent claims as submitted by the sponsor:
 DS = Drug Substance claim
 DP = Drug Product claim
 U and number = Method of Use claim (may be multiple). Specific Method of use claims are listed at <http://www.fda.gov/cder/orange/patex.htm>
3. Patents listed prior to August 18, 2003 are flagged with method of use claims only as applicable and submitted by the sponsor. They may not be flagged with respect to other claims which may apply.
4. *PED and PED represent pediatric exclusivity. Patents with pediatric exclusivity granted after August 18, 2003 will be indicated with *PED as was done prior to August 18, 2003. Patents with *PED added after August 18, 2003 will not contain any information relative to the patent itself other than the *PED extension. Information related specifically to the patent will be conveyed on the original patent only.
5. U.S. Patent Nos. RE 36481 and RE 36520 were relisted for Zocor (NDA 19-766) pursuant to the decision and related order in *Ranbaxy Labs. v. Leavitt*, No. 05-1838 (D.D.C. April 30, 2006). The '481 and '520 patents remained listed in Approved Drug Products with Therapeutic Equivalence Evaluations until any applicable periods of exclusivity pursuant to section 505(j)(5)(B)(iv) of the Federal Food, Drug, and Cosmetic Act were triggered and run. For additional information on this matter, please refer to Docket Nos. 2005P-0008 and 2005P-0046. Patents were subsequently delisted in the December 2006 Orange Book update as the exclusivity periods have triggered and run to expiration.

PATENT AND EXCLUSIVITY TERMS

ADB 1 of 35

PATENT & EXCLUSIVITY ABBREVIATIONS

D	NEW DOSING SCHEDULE (SEE INDIVIDUAL REFERENCES)
I	NEW INDICATION (SEE INDIVIDUAL REFERENCES)
M	MISCELLANEOUS EXCLUSIVITY CODES (SEE INDIVIDUAL REFERENCES)
NC	NEW COMBINATION
NCE	NEW CHEMICAL ENTITY
NDF	NEW DOSAGE FORM
NE	NEW ESTER OR SALT OF AN ACTIVE INGREDIENT
NP	NEW PRODUCT
NP*	NEW PRODUCT (MINT FLAVORED)
NPP	NEW PATIENT POPULATION
NR	NEW ROUTE
NS	NEW STRENGTH
ODE	ORPHAN DRUG EXCLUSIVITY
PC	PATENT CHALLENGE
PED	PEDIATRIC EXCLUSIVITY
RTO	RX TO OTC SWITCH OR OTC USE
U	PATENT USE CODE (SEE INDIVIDUAL REFERENCES)
W	EXCLUSIVITY ON THIS APPLICATION EXPIRING ON THIS DATE HAS BEEN WAIVED BY SPONSOR - SEE SECTION 1.8 OF ORANGE BOOK PREFACE WAIVED EXCLUSIVITY

EXCLUSIVITY DOSING SCHEDULE

D-1	ONCE A DAY APPLICATION
D-2	ONCE DAILY DOSING
D-3	SEVEN DAYS/SEVEN DAYS/SEVEN DAYS DOSING SCHEDULE
D-4	SEVEN DAYS/FOURTEEN DAYS DOSING SCHEDULE
D-5	TEN DAYS/ELEVEN DAYS DOSING SCHEDULE
D-6	SEVEN DAYS/NINE DAYS/FIVE DAYS DOSING SCHEDULE
D-7	BID DOSING
D-8	INTRAVENOUS, EPIDURAL AND INTRATHECAL DOSING
D-9	NARCOTIC OVERDOSE IN ADULTS
D-10	NARCOTIC OVERDOSE IN CHILDREN
D-11	POSTOPERATIVE NARCOTIC DEPRESSION IN CHILDREN
D-12	BEDTIME DOSING OF 800MG FOR TREATMENT OF ACTIVE DUODENAL ULCER
D-13	INCREASED MAXIMUM DAILY DOSAGE RECOMMENDATION
D-14	BEDTIME DOSING OF 800MG FOR TREATMENT OF ACTIVE BENIGN GASTRIC ULCER
D-15	SINGLE DAILY DOSE OF 25MG/37.5MG
D-16	CONTINUOUS INTRAVENOUS INFUSION
D-17	400MG EVERY 12 HOURS FOR THREE DAYS FOR UNCOMPLICATED URINARY TRACT INFECTIONS
D-18	LOWER RECOMMENDED STARTING DOSE GUIDELINES
D-19	BOLUS DOSING GUIDELINES
D-20	SINGLE 32MG DOSE
D-21	ALTERNATIVE DOSAGE OF 300MG ONCE DAILY AFTER THE EVENING MEAL
D-22	REDUCTION IN INFUSION TIME FROM 24 TO 4 HOURS FOR THE 60MG DOSE
D-23	INCREASE MAXIMUM DOSE AND VARIATIONS IN THE DOSING REGIMEN
D-24	FOR OVARIAN CANCER THE RECOMMENDED REGIMEN IS 135MG/M2 OR 175MG/M2 INTRAVENOUSLY OVER THREE HOURS EVERY THREE WEEKS
D-25	ADDITIONAL DOSAGE REGIMEN EQUAL TO HALF THE ORIGINAL DOSING REGIMEN
D-26	ONCE WEEKLY APPLICATION
D-27	BID DOSING IN PATIENTS 12 YEARS OF AGE AND OLDER FOR PREVENTION OF NAUSEA AND VOMITING ASSOCIATED WITH MODERATE EMETOGENIC CANCER CHEMOTHERAPY
D-28	USE OF ISOVUE-370 IN EXCRETORY UROGRAPHY AT EQUIVALENT GRAMS OF IODINE TO THE CURRENTLY APPROVED ISOVUE-250 AND ISOVUE-300
D-29	INCREASE OF CUMULATIVE DOSE TO 0.3MMOL/KG FOR MRI OF CNS IN ADULTS
D-30	5000 IU DOSE FOR PHOPHYLAXIX AGAINST DEEP VEIN THROMBOSIS
D-31	CHANGE IN RECOMMENDED TOTAL DAILY DOSE TO 80MG (40MG BID)
D-32	REMOVAL OF THE RESTRICTIONS LIMITING TREATMENT TO TWO CONSECUTIVE WEEKS AND TO SMALL AREAS
D-33	ONCE DAILY DOSING FOR PLAQUE PSORIASIS
D-34	EVERY FOUR MONTHS DOSAGE REGIMEN
D-35	FOR A ONE WEEK DOSING OF INTERDIGITAL TINEA PEDIS
D-36	FOR A SINGLE 2MG DOSE AS AN ALTERNATIVE TO THE 1MG DOSE GIVEN TWICE DAILY

PATENT AND EXCLUSIVITY TERMS

ADB 2 of 35

EXCLUSIVITY DOSING SCHEDULE

- D-37 DOSING REGIMEN FOR ADMINISTRATION EITHER ONCE DAILY (QD) OR TWICE DAILY (BID)
- D-38 CONTINUOUS INFUSION AS AN ALTERNATE METHOD OF ADMINISTRATION
- D-39 CHANGE IN TIME TO TAKE THE DRUG PRIOR TO A MEAL TO PREVENT MEAL-INDUCED HEARTBURN SYMPTOMS FROM "...1/2 TO 1 HOUR BEFORE EATING" TO "... RIGHT BEFORE EATING OR UP TO 60MIN BEFORE CONSUMING..."
- D-40 ONCE-A-DAY DOSING REGIMEN
- D-41 DRUG MAY BE DOSED RIGHT BEFORE A MEAL OR ANY TIME UP TO 30MIN BEFORE EATING OR DRINKING FOOD AND BEVERAGES THAT WOULD BE EXPECTED TO CAUSE SYMPTOMS
- D-42 TEN DAY DOSING REGIMEN FOR TRIPLE THERAPY, PREVACID IN COMBINATION WITH CLARITHROMYCIN AND AMOXICILLIN, FOR THE ERADICATION OF H.PYLORI IN PATIENTS WITH DUODENAL ULCER DISEASE
- D-43 INITIATION OF TREATMENT WITH 900MG/DAY BY DELETION OF THE REQUIREMENT TO TITRATE TO 900MG/DAY OVER A 3-DAY PERIOD
- D-44 IN A CLINICAL TRIAL, FEWER DISCONTINUATIONS DUE TO ADVERSE EVENTS, ESPECIALLY DIZZINESS AND VERTIGO, WERE OBSERVED WHEN TITRATING THE DOSE IN INCREMENTS OF 50MG/DAY EVERY 3 DAYS UNTIL AN EFFECTIVE DOSE (NOT EXCEEDING 400MG/DAY) WAS REACHED
- D-45 ONCE DAILY DOSING FOR MAINTENANCE ONLY
- D-46 NEW DOSING REGIMEN OF 80MG DAILY
- D-47 PREVENTION OF HEARTBURN SYMPTOMS WHEN ADMINISTERED FROM 15 MINUTES UP TO, BUT NOT INCLUDING, 1 HOUR PRIOR TO A PROVOCATIVE MEAL
- D-48 ADMINISTRATION OF CISATRICURIUM A NEUROMUSCULAR BLOCKING AGENT AT DOSES OF 3 AND 4X THE ED95 OF CISATRICURIUM FOLLOWING INDUCTION WITH THIOPENTAL
- D-49 PEDIATRIC DOSING GUIDELINES
- D-50 INFORMATION FOR USE OF CORVERT IN POST-CARDIAC SURGERY PATIENTS
- D-51 OPTIONAL STARTING DOSE OF 40MG/DAY
- D-52 ALTERNATE DOSING REGIMEN OF 1250MG TWICE DAILY
- D-53 USE IN PEDIATRIC PATIENTS FROM 1 MONTH TO 16 YEARS OF AGE
- D-54 USE OF ZYBAN FOR MAINTENANCE THERAPY. TREATMENT UP TO 6 MONTHS WAS SHOWN EFFICACIOUS
- D-55 ADDITION OF A HIGHER DOSE OF NUTROPIN FOR PUBERTAL PATIENTS (PUBERTAL DOSE LESS THAN OR EQUAL TO 0.7MG/KG/WEEK)
- D-56 ADDITION OF POSTPRANDIAL DOSING
- D-57 3-HOUR INFUSION OF TAXOL GIVEN EVERY THREE WEEKS AT A DOSE OF 175MG/M2 FOLLOWED BY CISPLATIN AT A DOSE OF 75MG/M2 FOR THE FIRST-LINE TREATMENT OF ADVANCED OVARIAN CANCER
- D-58 CHANGE IN DOSING INTERVAL TO ONCE-DAILY ADMINISTRATION
- D-59 REDUCTION OF ELEVATED LDL-C IN A NEW, HIGHER STRENGTH TABLET, 0.8MG, AND FOR EXTENSION OF THE DOSAGE RANGE TO 0.8MG DAILY
- D-60 ADDITION OF A POST-OPERATIVE DOSING REGIMEN
- D-61 ONCE WEEKLY DOSING FOR THE TREATMENT OF POSTMENOPAUSAL OSTEOPOROSIS
- D-62 ONCE WEEKLY DOSING FOR THE PREVENTION OF POSTMENOPAUSAL OSTEOPOROSIS
- D-63 TO ALLOW A TITRATION DOSING REGIMEN USING A 25MG DOSE
- D-64 INCREASING DOSAGE FOR NERVE BLOCK ANESTHESIA USING NAROPIN 7.5MG/ML AND FOR EXTENDING THE DURATION OF TREATMENT FOR POSTOPERATIVE ANALGESIA USING NAROPIN 2MG/ML
- D-65 CHANGE DOSING AND ADMINISTRATION TO INDICATE MAINTENANCE OF WEIGHT LOSS OVER AN 18 MONTH PERIOD THUS EXTENDING THE USE OF THIS DRUG FROM ONE TO TWO YEARS
- D-66 DOSING RECOMMENDATIONS FOR PATIENTS UNDERGOING PCI
- D-67 SHORTER TREATMENT COURSE OF THREE DAYS IN THE TREATMENT OF RECURRENT EPISODES OF GENITAL HERPES
- D-68 CHANGE OF ADMIN RATE FOR INFUSION OF AREDIA FOR TREATMENT OF MODERATE AND SEVERE HYPERCALCEMIA OF MALIGNANCY FROM 24 HOURS TO 2 HOURS UP TO BUT NOT INCLUDING 24 HOURS
- D-69 SHORTENED DOSING REGIMEN TO 5 DAYS FOR THE TREATMENT OF ACUTE EXACERBATION OF CHRONIC BRONCHITIS
- D-70 80MG ONCE DAILY DOSING REGIMEN
- D-71 EIGHT WEEK DOSING REGIMEN
- D-72 INFORMATION REGARDING INCREASED RATE OF INFUSION FOR DEPACON
- D-73 ONCE A WEEK DOSING FOR THE TREATMENT OF POSTMENOPAUSAL OSTEOPOROSIS
- D-74 ONCE A WEEK DOSING FOR THE PREVENTION OF POSTMENOPAUSAL OSTEOPOROSIS
- D-75 INTERMITTENT DOSING REGIMEN, STARTING DAILY DOSE 14 DAYS PRIOR TO THE ANTICIPATED ONSET OF MENSTRUATION THROUGH THE FIRST FULL DAY OF MENSES AND REPEATING WITH EACH NEW CYCLE
- D-76 FOR USE ON AN "AS NEEDED" OR PRN BASIS FOR THE MANAGEMENT OF NASAL SYMPTOMS IN PATIENTS FOR WHOM THE DRUG IS INDICATED
- D-77 ADDITION OF 20MG AND 40MG DAILY AS OPTIONAL STARTING DOSES WITH 40MG INTENDED FOR PATIENTS WHO REQUIRE A LARGE REDUCTION IN LDL-C (MORE THAN 45%)
- D-78 USE OF FLEXERIL 5MG FOR THE RELIEF OF MUSCLE SPASM ASSOCIATED WITH ACUTE, PAINFUL, MUSCULOSKELETAL CONDITIONS
- D-79 NEW LOWER STARTING DOSE FOR TREATMENT OF MODERATE TO SEVERE VASOMOTOR SYMPTOMS AND/OR MODERATE TO SEVERE SYMPTOMS OF VULVAR AND VAGINAL ATROPHY ASSOCIATED W/ THE MENOPAUSE

PATENT AND EXCLUSIVITY TERMS

ADB 3 of 35

EXCLUSIVITY DOSING SCHEDULE

- D-80 CHANGE OF DOSING SCHEDULE FOR LANTUS FROM ONCE DAILY AT BEDTIME TO FLEXIBLE DAILY DOSING
- D-81 NEW LOWER STARTING DOSE FOR THE PREVENTION OF POSTMENOPAUSAL OSTEOPOROSIS
- D-82 USE OF PREMARIN 0.3 MG AND 0.45 MG FOR THE PREVENTION OF POSTMENOPAUSAL OSTEOPOROSIS
- D-83 750 MG, ONCE DAILY FOR 5 DAYS FOR COMMUNITY ACQUIRED PNEUMONIA (CAP)
- D-84 ONCE-A-DAY DOSING OF FLOXACIN OTIC FOR THE TREATMENT OF ADULTS AND PEDIATRIC PATIENTS(AGES 6 MO & OLDER) W/ OTITIS EXTERNA CAUSED BY SUSCEPTIBLE STRAINS OF E.COLI, P.AERUGINOSA AND S.AUREUS
- D-85 LOWER RECOMMENDED STARTING DOSE GUIDELINES FOR TREATMENT OF MODERATE TO SEVERE VASOMOTOR SYMPTOMS ASSOCIATED WITH THE MENOPAUSE
- D-86 FOR USE IN SELECT EXTERNAL INSULIN PUMPS
- D-87 ADDITION OF ONCE-WEEKLY DOSING FOR THE TREATMENT TO INCREASE BONE MASS IN MEN WITH OSTEOPOROSIS
- D-88 NEW DOSING RANGE OF 200-400MG PER DAY IN TWO DIVIDED DOSES FOR ADULTS WITH PARTIAL SEIZURES
- D-89 USE OF REYATAZ 300 MG/RITONAVIR 100 MG ONCE DAILY FOR TREATMENT IN HIV-INFECTED ANTIRETROVIRAL-EXPERIENCED PATIENTS
- D-90 ADDITION OF DAYTIME ADMINISTRATION TO TREAT VULVOVAGINAL CANDIDIASIS
- D-91 ALTERNATE INTERMITTENT DOSING REGIMEN
- D-92 ALTERNATIVE DOSAGE OF 1000MG ONCE DAILY AT BEDTIME
- D-93 ALTERNATE TWO OR THREE TIMES DAILY DOSING REGIMENS
- D-94 NEW MAXIMUM DOSAGE OF 72 MG/DAY IN ADOLESCENTS 13-17 YEARS OF AGE WITH ATTENTION DEFECIT HYPERACTIVITY DISORDER (ADHD)
- D-95 BROADENED INITIAL STARTING DOSE FOR HYPERTENSION FROM 50 MG TO 100 MG TO 25 MG TO 100 MG DOSE RANGE
- D-96 ONCE-MONTHLY TREATMENT OF POSTMENOPAUSAL OSTEOPOROSIS WITH BONIVA (IBANDRONATE SODIUM) 150 MG TABLETS
- D-97 PED CANCER PT POPULATION EXPANDED TO INCLUDE PTS 6 MOS UP TO BUT NOT INCLUDING 4 YRS AND DOSING INSTRUCTIONS TO ADMIN 30 MIN BEFPRE CHEMO WITH SECOND AND THIRD DOSES 4 & 8 HOURS AFTER FIRST DOSE
- D-98 DOSING FOR PED SURGICAL PTS EXPANDED TO INCLUDE PTS 1 MONTH UP TO BUT NOT INCLUDING 2 YEARS OF AGE
- D-99 ONCE DAILY ADMINISTRATION FOR THE TREATMENT OF HIV INFECTION IN THERAPY NAIVE ADULT PATIENTS
- D-100 750 MG ONCE DAILY FOR FIVE DAYS FOR THE TREATMENT OF ACUTE BACTERIAL SINUSITIS
- D-101 ONCE DAILY IN CHRONIC IDIOPATHIC UTICARIA FOR ADULTS AND CHILDREN 12 YEARS OF AGE AND OLDER
- D-102 NEW DOSING REGIMEN OF ONE SPRAY TWICE DAILY FOR SEASONAL ALLERIC RHINITIS IN PATIENTS 12 YRS OF AGE AND OLDER
- D-103 NEW DOSING RECOMMENDATION FOR THE TREATMENT OF RECURRENT GENITAL HERPES IN IMMUNOCOMPETENT PATIENTS, SPECIFICALLY A REDUCTION IN COURSE OF THERAPY FROM FAMCICLOVIR 125 MG TWICE-A-DAY FOR 5 DAYS TO 1000 MG TWICE-A-DAY FOR 1 DAY.
- D-104 0.5MG/0.1MG FOR THE TREATMENT OF MODERATE TO SEVERE VASOMOTOR SYMPTOMS ASSOCIATED WITH MENOPAUSE IN WOMEN WHO HAVE A UTERUS

EXCLUSIVITY INDICATION

- I-1 DYSMENORRHEA
- I-2 CHOLANGIOPANCREATOGRAPHY
- I-3 INTRAVENOUS DIGITAL SUBTRACTION ANGIOGRAPHY
- I-4 PERIPHERAL VENOGRAPHY (PHLEBOGRAPHY)
- I-5 HYSTEROSALPINGOGRAPHY
- I-6 TREATMENT OF JUVENILE ARTHRITIS
- I-7 BIOPSY PROVEN MINIMAL CHANGE NEPHROTIC SYNDROME IN CHILDREN
- I-8 ADULT INTRAVENOUS CONTRAST-ENHANCED COMPUTED TOMOGRAPHY OF THE HEAD AND BODY
- I-9 PREVENTION OF POSTOPERATIVE NAUSEA AND VOMITING
- I-10 PREVENTION OF POSTOPERATIVE DEEP VENOUS THROMBOSIS AND PULMONARY EMBOLISM IN TOTAL HIP REPLACEMENT SURGERY
- I-11 RELIEF OF MILD TO MODERATE PAIN
- I-12 TREATMENT OF CUTANEOUS CANDIDIASIS
- I-13 URINARY TRACT INFECTION (UTI) PREVENTION FOR PERIODS UP TO FIVE MONTHS IN WOMEN WITH A HISTORY OF RECURRENT UTI
- I-14 SEBORRHEIC DERMATITIS
- I-15 PHOTOPHERESIS IN THE PALLIATIVE TREATMENT OF SKIN MANIFESTATIONS OF CUTANEOUS T-CELL LYMPHOMA IN PERSONS NOT RESPONSIVE TO OTHER TREATMENT

PATENT AND EXCLUSIVITY TERMS

ADB 4 of 35

EXCLUSIVITY INDICATION

- I-16 STIMULATE THE DEVELOPMENT OF MULTIPLE FOLLICLES/OOCYTES IN OVULATORY PATIENTS PARTICIPATING IN AN IN VITRO FERTILIZATION PROGRAM
- I-17 MANAGEMENT OF CONGESTIVE HEART FAILURE
- I-18 ENDOSCOPIC RETROGRADE PANCREATOGRAPHY
- I-19 HERNIOGRAPHY
- I-20 KNEE ARTHROGRAPHY
- I-21 HIGH DOSE METHOTREXATE WITH LEUCOVORIN RESCUE IN COMBINATION WITH OTHER CHEMOTHERAPEUTIC AGENTS TO DELAY RECURRENCE IN PATIENTS WITH NONMETASTATIC OSTEOSARCOMA WHO HAVE UNDERGONE SURGICAL RESECTION OR AMPUTATION FOR THE PRIMARY TUMOR
- I-22 RESCUE AFTER HIGH-DOSE METHOTREXATE THERAPY IN OSTEOSARCOMA
- I-23 SHORT-TERM TREATMENT OF ACTIVE BENIGN GASTRIC ULCER
- I-24 TREATMENT OF RHEUMATOID ARTHRITIS
- I-25 ADULT INTRA-ARTERIAL DIGITAL SUBTRACTION ANGIOGRAPHY OF THE HEAD, NECK, ABDOMINAL, RENAL AND PERIPHERAL VESSELS
- I-26 TREATMENT OF LIVER FLUKES
- I-27 ADJUNCTIVE THERAPY TO DIET TO REDUCE THE RISK OF CORONARY ARTERY DISEASE
- I-28 SELECTIVE ADULT VISCERAL ARTERIOGRAPHY
- I-29 METASTATIC BREAST CANCER IN PREMENOPAUSAL WOMEN AS AN ALTERNATIVE TO OOPHORECTOMY OR OVARIAN IRRADIATION
- I-30 TREATMENT OF TINEA PEDIS
- I-31 CONTRAST ENHANCEMENT AGENT TO FACILITATE VISUALIZATION OF LESIONS IN THE SPINE AND ASSOCIATED TISSUES
- I-32 PEDIATRIC MYELOGRAPHY
- I-33 ORAL USE OF DILUTED OMNIPAQUE INJECTION IN ADULTS FOR CONTRAST ENHANCED COMPUTED TOMOGRAPHY OF THE ABDOMEN
- I-34 ORAL USE IN ADULTS FOR PASS-THROUGH EXAMINATION OF THE GASTROINTESTINAL TRACT
- I-35 PEDIATRIC CONTRAST ENHANCEMENT OF COMPUTED TOMOGRAPHIC HEAD IMAGING
- I-36 ARTHROGRAPHY OF THE SHOULDER JOINTS IN ADULTS
- I-37 RADIOGRAPHY OF THE TEMPOROMANDIBULAR JOINT IN ADULTS
- I-38 CONTRAST ENHANCEMENT AGENT TO FACILITATE VISUALIZATION OF LESIONS OF THE CENTRAL NERVOUS SYSTEM IN CHILDREN (2 YEARS OF AGE AND OLDER)
- I-39 TREATMENT OF ACUTE MYOCARDIAL INFARCTION
- I-40 PRIMARY NOCTURNAL ENURESIS
- I-41 MIGRAINE HEADACHE PROPHYLAXIS
- I-42 HERPES ZOSTER
- I-43 HERPES SIMPLEX ENCEPHALITIS
- I-44 MAINTENANCE THERAPY IN HEALED DUODENAL ULCER PATIENTS AT DOSE OF 1 GRAM TWICE DAILY
- I-45 ACUTE TREATMENT OF VARICELLA ZOSTER VIRUS
- I-46 USE IN PEDIATRIC COMPUTED TOMOGRAPHIC HEAD AND BODY IMAGING
- I-47 TREATMENT OF PEDIATRIC PATIENTS WITH SYMPTOMATIC HUMAN IMMUNODEFICIENCY VIRUS (HIV) DISEASE
- I-48 PEDIATRIC ANGIOCARDIOGRAPHY
- I-49 TREATMENT OF TRAVELERS' DIARRHEA DUE TO SUSCEPTIBLE STRAINS OF ENTEROTOXIGENIC ESCHERICHIA COLI
- I-50 FOR USE IN WOMEN WITH AXILLARY NODE-NEGATIVE BREAST CANCER
- I-51 TREATMENT OF PRIMARY DYSMENORRHEA AND FOR THE TREATMENT OF IDIOPATHIC HEAVY MENSTRUAL BLOOD LOSS
- I-52 PEDIATRIC EXCRETORY UROGRAPHY
- I-53 TREATMENT OF PANIC DISORDER, WITH OR WITHOUT AGORAPHOBIA
- I-54 RENAL CONCENTRATION CAPACITY TEST
- I-55 HYPERTENSION
- I-56 EROSIIVE GASTROESOPHAGEAL REFLUX DISEASE
- I-57 SHORT-TERM TREATMENT OF ACTIVE DUODENAL ULCER
- I-58 INITIAL TREATMENT OF ADVANCED OVARIAN CARCINOMA IN COMBINATION WITH OTHER APPROVED CHEMOTHERAPEUTIC AGENTS
- I-59 ENDOSCOPICALLY DIAGNOSED ESOPHAGITIS, INCLUDING EROSIIVE AND ULCERATIVE ESOPHAGITIS, AND ASSOCIATED HEARTBURN DUE TO GASTROESOPHAGEAL REFLUX DISEASE
- I-60 SINGLE APPLICATION TREATMENT OF HEAD LICE IN CHILDREN TWO MONTHS TO TWO YEARS IN AGE
- I-61 FEMALE ANDROGENETIC ALOPECIA
- I-62 PREVENTION AND TREATMENT OF POSTMENOPAUSAL OSTEOPOROSIS
- I-63 ONCE DAILY TREATMENT AS INITIAL THERAPY IN THE TREATMENT OF HYPERTENSION
- I-64 PREVENTION OF SUPRAVENTRICULAR TACHYCARDIAS
- I-65 PREVENTION OF UPPER GASTROINTESTINAL BLEEDING IN CRITICALLY ILL PATIENTS
- I-66 UNCOMPLICATED GONORRHEA

PATENT AND EXCLUSIVITY TERMS

ADB 5 of 35

EXCLUSIVITY INDICATION

- I-67 TREATMENT OF ACUTE ASTHMATIC ATTACKS IN CHILDREN SIX YEARS OF AGE AND OLDER
- I-68 CENTRAL PRECOCIOUS PUBERTY
- I-69 SHORT TERM TREATMENT OF PATIENTS WITH SYMPTOMS OF GASTROESOPHAGEAL REFLUX DISEASE (GERD), AND FOR THE SHORT TERM TREATMENT OF ESOPHAGITIS DUE TO GERD INCLUDING ULCERATIVE DISEASE DIAGNOSED BY ENDOSCOPY
- I-70 USE IN COMBINATION WITH 5-FLUOROURACIL TO PROLONG SURVIVAL IN THE PALLIATIVE TREATMENT OF PATIENTS WITH ADVANCED COLORECTAL CANCER
- I-71 VARICELLA INFECTIONS (CHICKENPOX)
- I-72 PREVENTION OF CMV DISEASE IN TRANSPLANT PATIENTS AT RISK FOR CMV DISEASE
- I-73 INITIATE AND MAINTAIN MONITORED ANESTHESIA CARE (MAC) SEDATION DURING DIAGNOSTIC PROCEDURES
- I-74 INTRAVENOUS DIGITAL SUBTRACTION ANGIOGRAPHY
- I-75 TREATMENT OF ENDOSCOPICALLY DIAGNOSED EROSIIVE ESOPHAGITIS
- I-76 PREVENTION OF OSTEOPOROSIS
- I-77 DERMAL INFECTIONS-TINEA PEDIS, TINEA CORPORIS, TINEA CRURIS DUE TO EPIDERMOPHYTON FLOCCOSUM
- I-78 CONTRAST ENHANCED COMPUTED TOMOGRAPHIC IMAGING OF THE HEAD AND BODY AND INTRAVENOUS EXCRETORY UROGRAPHY
- I-79 MANAGEMENT OF CHRONIC STABLE ANGINA AND ANGINA DUE TO CORONARY ARTERY SPASM
- I-80 DIAGNOSIS AND LOCALIZATION OF ISCHEMIA AND CORONARY HEART DISEASE
- I-81 PROPHYLAXIS IN DESIGNATED IMMUNOCOMPROMISED CONDITIONS TO REDUCE THE INCIDENCE OF OROPHARYNGEAL CANDIDIASIS
- I-82 TREATMENT OF TRAVELERS' DIARRHEA
- I-83 ANGIOCARDIOGRAPHY, CONTRAST ENHANCED COMPUTED TOMOGRAPHIC IMAGING OF THE HEAD AND BODY, AND INTRAVENOUS EXCRETORY UROGRAPHY IN CHILDREN
- I-84 INTRAOPERATIVE AND POSTOPERATIVE TACHYCARDIA AND/OR HYPERTENSION
- I-85 TREATMENT OF ANOREXIA ASSOCIATED WITH WEIGHT LOSS IN PATIENTS WITH AIDS
- I-86 TREATMENT OF SECONDARY CARNITINE DEFICIENCY
- I-87 RENAL IMAGING AGENT FOR USE IN CHILDREN
- I-88 MANAGEMENT OF ENDOMETRIOSIS
- I-89 EPIDURAL USE IN LABOR AND DELIVERY AS AN ANALGESIC ADJUNCT TO BUPIVACAINE
- I-90 INTENSIVE CARE UNIT SEDATION
- I-91 MONOTHERAPY USE FOR HYPERTENSION
- I-92 ADJUNCTIVE THERAPY IN THE MANAGEMENT OF HEART FAILURE
- I-93 PREVENTION OF EXERCISE-INDUCED BRONCHOSPASM IN CHILDREN AGES 4-11 YEARS
- I-94 USE WITH MRI IN ADULTS TO PROVIDE CONTRAST ENHANCEMENT AND FACILITATE VISUALIZATION OF LESIONS IN THE BODY [EXCLUDING THE HEART]
- I-95 TREATMENT OF LEFT VENTRICULAR DYSFUNCTION FOLLOWING MYOCARDIAL INFARCTION
- I-96 TREATMENT OF SYMPTOMATIC BENIGN PROSTATIC HYPERPLASIA
- I-97 ORAL OR RECTAL USE IN CHILDREN FOR THE EXAMINATION OF THE GASTROINTESTINAL TRACT
- I-98 TREATMENT OF CHILDREN WHO HAVE GROWTH FAILURE ASSOCIATED WITH CHRONIC RENAL INSUFFICIENCY
- I-99 PEDIATRIC ANESTHESIA IN CHILDREN 3 YEARS AND OLDER
- I-100 TO DECREASE THE INCIDENCE OF CANDIDIASIS IN PATIENTS UNDERGOING BONE MARROW TRANSPLANTATION WHO RECEIVE CYTOTOXIC CHEMOTHERAPY AND/OR RADIATION THERAPY
- I-101 TREATMENT OF DIABETIC NEPHROPATHY IN PATIENTS WITH TYPE I INSULIN-DEPENDENT DIABETES MELLITUS AND RETINOPATHY
- I-102 TREATMENT OF OBSESSIVE-COMPULSIVE DISORDER
- I-103 PROPHYLAXIS AGAINST PNEUMOCYSTIS CARINII PNEUMONIA IN INDIVIDUALS WHO ARE IMMUNOCOMPROMISED AND CONSIDERED TO BE AT RISK OF DEVELOPING PNEUMOCYSTIS CARINII PNEUMONIA
- I-104 TREATMENT OF PULMONARY AND EXTRAPULMONARY ASPERGILLOSIS IN PATENTS WHO ARE INTOLERANT OF OR WHO ARE REFRACTORY TO AMPHOTERICIN B THERAPY
- I-105 TREATMENT OF METASTATIC CARCINOMA OF THE BREAST AFTER FAILURE OF FIRST-LINE OR SUBSEQUENT CHEMOTHERAPY
- I-106 TREATMENT OF ACROMEGALY
- I-107 VAGINAL CANDIDIASIS
- I-108 EXPANDED USE-FOR ICU PATIENTS UNDERGOING LONG-TERM INFUSION DURING MECHANICAL VENTILATION
- I-109 TYPHOID FEVER
- I-110 PREVENTION OF NAUSEA AND VOMITING ASSOCIATED WITH RADIOTHERAPY
- I-111 TREATMENT OF PAGET'S DISEASE OF BONE
- I-112 MANAGEMENT OF MODERATE TO SEVERE PAIN
- I-113 TREATMENT OF PROSTATITIS
- I-114 USE IN CHILDREN TO VISUALIZE LESIONS WITH ABNORMAL VASCULARITY IN THE BRAIN (INTRACRANIAL LESIONS), SPINE, AND ASSOCIATED TISSUE

PATENT AND EXCLUSIVITY TERMS

ADB 6 of 35

EXCLUSIVITY INDICATION

- I-115 USE IN MRI IN ADULTS TO VISUALIZE LESIONS IN THE HEAD AND NECK
- I-116 MAINTENANCE OF HEALING OF EROSIIVE ESOPHAGITIS
- I-117 TO SLOW THE PROGRESSION FO CORONARY ATHEROSCLEROSIS IN PATIENTS WITH CORONARY HEART DISEASE
- I-118 PREVENTION OF DEEP VEIN THROMBOSIS, WHICH MAY LEAD TO PULMONARY EMBOLISM FOLLOWING KNEE REPLACEMENT SURGERY
- I-119 TREATMENT OF ANEMIA CAUSED BY UTERINE LEIOMYOMATA IN WOMEN WHO FAIL IRON THERAPY
- I-120 MAINTENANCE THERAPY FOR GASTRIC ULCER PATIENTS AT REDUCED DOSAGE AFTER HEALING ACUTE ULCERS
- I-121 EXPANDED PATIENT POPULATION -- USE IN ICU PATIENTS
- I-122 PSORIASIS OF THE SCALP
- I-123 RELIEF OF MILD TO MODERATE PAIN IN PATIENTS AGED 6 MONTHS AND OLDER
- I-124 LEUCOCYTE LABELED SCINTIGRAPHY AS AN ADJUNCT IN THE LOCALIZATION OF INTRA-ABDOMINAL INFECTION AND INFLAMMATORY BOWEL DISEASE
- I-125 EXPANSION OF CONSCIOUS SEDATION INDICATION TO INCLUDE SHORT THERAPEUTIC PROCEDURES
- I-126 ADJUNCT TO THALLIUM- 201 MYOCARDIAL PERFUSION IN PATIENTS UNABLE TO EXERCISE ADEQUATELY
- I-127 TREATMENT OF ACYCLOVIR-RESISTANT HERPES IN IMMUNOCOMPROMISED PATIENTS
- I-128 IN PT W/ CH DISEASE AND HYPERCHOLESTEROLEMIA: REDUCE RISK TOTAL MORTALITY BY REDUCING CORONARY DEATH; REDUCE RISK NON-FATAL MI; REDUCE RISK UNDERGOING MYOCARDIAL REVASCULARIZATION PROCEDURES; REDUCTION ELEVATED TOTAL AND LDL CHOL LEVELS...
- I-129 TREATMENT OF ALCOHOL DEPENDENCE
- I-130 MAINTENANCE OF HEALING OF EROSIIVE ESOPHAGITIS
- I-131 PERIPHERAL ARTERIOGRAPHY
- I-132 TREATMENT OF MANIC PHASE OF BIPOLAR DISORDER
- I-133 MANAGEMENT OF CHRONIC STABLE ANGINA
- I-134 HEART FAILURE POST MYOCARDIAL INFARCTION
- I-135 BONE METASTASES ASSOCIATED WITH MULTIPLE MYELOMA
- I-136 IDIOPATHIC CHRONIC URTICARIA
- I-137 PREVENTION OF METAL-INDUCED HEART BURN, ACID INDIGESTION, AND SOUR STOMACH WHEN TAKEN 30 MINUTES PRIOR TO CONSUMING FOOD OR BEVERAGES
- I-138 TREATMENT OF ACUTE RECURRENT GENITAL HERPES
- I-139 PALLIATIVE TREATMENT OF ADVANCED BREAST CANCER IN PRE- AND PERIMENOPAUSAL WOMEN
- I-140 PREVENTION OF CYTOMEGALOVIRUS (CMV) DISEASE IN INDIVIDUALS WITH HIV INFECTION AT RISK FOR DEVELOPING CMV DISEASE
- I-141 TREATMENT OF HEMODYNAMICALLY STABLE PATIENTS WITHIN 24 HOURS OF ACUTE MYOCARDIAL INFARCTION TO IMPROVE SURVIVAL
- I-142 LOCALIZE MYOCARDIAL ISCHEMIA(REVERSIBLE DEFECT) AND INFARCTION (NON-REVERSIBLE DEFECTS) IN EVALUATING MYOCARDIAL FUNCTION
- I-143 EPISODIC TREATMENT OF RECURRENT GENITAL HERPES IN IMMUNOCOMPETENT ADULTS
- I-144 ENHANCEMENT OF MRI OF THE ADULT BODY INTERNAL ORGANS
- I-145 0.1MMOL/KG AS A SINGLE INTRAVEOUS BOLUS FOR MRI OF THE CNS IN CHILDREN
- I-146 CONTRAST ENHANCEMENT AND FACILITATION OF VISUALIZATION OF EXTRACRANIAL HEAD AND NECK LESIONS
- I-147 PREVENTION OF GALLSTONE FORMATION IN OBESE PATIENTS EXPERIENCING RAPID WEIGHT LOSS
- I-148 TREATMENT OF ACUTE PNEUMOCYSTIC CARINI PNEUMONIA (PCP) IN HIV-INFECTED PATIENTS WHOSE ALVEOLAR-ARTERIAL OXYGEN DIFFERENCE (AaDO2) IS LESS THAN OR EQUAL TO 55 TORR
- I-149 TREATMENT OF PATIENTS WITH NON-SMALL CELL LUNG CANCER
- I-150 TREATMENT OF OBSESSIVE COMPULSIVE DISORDER AND PANIC DISORDER
- I-151 PREVENTION OF AND PREVENTION OF FURTHER POSTOPERATIVE NAUSEA AND VOMITING IN PEDIATRIC PATIENTS RECEIVING GENERAL ANESTHESIA
- I-152 SLOWING THE PROGRESSION OF CORONARY ATHEROSCLEROSIS AND REDUCING THE RISK OF ACUTE CORONARY EVENTS
- I-153 MANAGEMENT OF SEVERE SPASTICITY [ENCOMPASES SPINAL AND CEREBRAL ORIGIN]
- I-154 PATIENT POPULATION ALTERED TO INCLUDE PEDIATRIC USE
- I-155 TREATMENT OF ONCHOMYCOSIS DUE TO DERMATOPHYTES (TINEA UNGUIUM) OF THE TOENAIL WITH OR WITHOUT FINGERNAIL INVOLVEMENT
- I-156 ADDITIONAL DATA REGARDING THE SAFE USE OF NORVASC IN PATIENTS WITH HEART FAILURE
- I-157 TREATMENT OF ACUTE UNCOMPLICATED CYSTITIS IN FEMALES
- I-158 TREATMENT OF OSTEOLYTIC BONE METASTASES OF BREAST CANCER
- I-159 FOR HYPERCHOLESTEROLEMIC PATIENTS WITHOUT CLINICALLY EVIDENT HEART DISEASE REDUCE THE RISK OF MYOCARDIAL INFARCTION, REVASCULARIZATION, AND DEATH DUE TO CARDIOVASCULAR CAUSES WITH NO INCREASE IN DEATH FROM NON-CARDIOVASCULAR CAUSES
- I-160 TREATMENT OF BACTERIAL CORNEAL ULCERS
- I-161 TREATMENT OF ADULT-ONSET OR CHILDHOOD-ONSET ADULT GROWTH HORMONE DEFICIENCY
- I-162 FOR USE IN PATIENTS 6-11 YEARS OF AGE

PATENT AND EXCLUSIVITY TERMS

ADB 7 of 35

EXCLUSIVITY INDICATION

- I-163 TREATMENT OF PHOTOPHOBIA
- I-164 CHRONIC BACTERIAL PROSTATITIS
- I-165 MANAGEMENT OF ADULTS WITH ACTIVE, CLASSIC AND DEFINITIVE RHEUMATOID ARTHRITIS WHO HAVE HAD INSUFFICIENT THERAPEUTIC RESPONSE TO OR ARE INTOLERANT OF AN ADEQUATE TRIAL OF FULL DOSES OF ONE OR MORE NON-STEROIDAL ANTI-INFLAMMATORY DRUGS
- I-166 TREATMENT OF BULIMIA
- I-167 COMPLICATED INTRA-ABDOMINAL INFECTIONS (USED IN COMBINATION WITH METRONIDAZOLE) CAUSED BY MIXED AEROBIC/ANAEROBIC PATHOGENS
- I-168 MANAGEMENT OF LOCALLY CONFINED STAGE B2-C METASTATIC CARCINOMA OF THE PROSTATE (IN COMBINATION WITH LHRH AGONISTS)
- I-169 USE IN COMBINATION WITH CORTICOSTEROIDS AS INITIAL CHEMOTHERAPY FOR THE TREATMENT OF PATIENTS WITH PAIN RELATED TO ADVANCED HORMONE-REFRACTORY PROSTATE CANCER
- I-170 PROPHYLACTIC USE DURING HEAD LICE EPIDEMICS
- I-171 RELIEF OF SYMPTOMS OF THE COMMON COLD
- I-172 TREATMENT OF INITIAL EPISODE OF GENITAL HERPES
- I-173 PREOPERATIVELY FOR THE PREVENTION OF INFECTION IN TRANSRECTAL PROSTATE BIOPSY
- I-174 PELVIC INFLAMMATORY DISEASE
- I-175 TREATMENT OF TINEA CORPORA AND TINEA CRURIS
- I-176 TREATMENT OF POSTOPERATIVE INFLAMMATION IN PATIENTS WHO HAVE UNDERGONE CATARACT EXTRACTION
- I-177 TX OF MODERATE ACNE VULGARIS IN FEMALES, GREATER OR EQUAL TO 15YRS OF AGE, WHO HAVE NO KNOWN CONTRAINDICATIONS TO ORAL CONTRACEPTIVE THERAPY, DESIRE CONTRACEPTION, HAVE ACHIEVED MENARCHE AND ARE UNRESPONSIVE TO TOPICAL ANTI-ACNE MEDICATIONS
- I-178 TREATMENT OF ONCHOMYCOSIS OF THE FINGERNAIL WITHOUT CONCOMITANT ONCHOMYCOSIS OF THE TOENAIL WITH A PULSE DOSING REGIMEN
- I-179 NOSOCOMIAL PNEUMONIA-MILD TO MODERATE AND SEVERE CAUSED BY HAEMOPHILUS INFLUENZAE OR KLEBSIELLA PNEUMONIAE
- I-180 TREATMENT OF PLANTAR TINEA PEDIS (MOCCASIN TYPE)
- I-181 TREATMENT OF PATIENTS WITH COMPLEX PARTIAL SEIZURES WITH AND WITHOUT SECONDARY GENERALIZATION
- I-182 TREATMENT OF GROWTH FAILURE ASSOCIATED WITH TURNER SYNDROME
- I-183 MAINTENANCE THERAPY IN THE MANAGEMENT OF MILD TO MODERATE ASTHMA IN PEDIATRIC PATIENTS AGES 6-11
- I-184 TREATMENT OF PANIC DISORDER AT A RECOMMENDED DOSE RANGE OF 1 TO 2MG/DAY (MAXIMUM OF 4MG)
- I-185 PREVENTION OF OSTEOPOROSIS IN POSTMENOPAUSAL WOMEN
- I-186 TREATMENT OF TINEA (PITYRIASIS) VERSICOLOR CAUSED BY OR PRESUMED TO BE CAUSED BY PITYROSPORUM ORBICULARE (ALSO KNOWN AS MALASSEZIA FURFUR OR M. ORBICULARE)
- I-187 PREVENTION OF FRACTURES IN THE TREATMENT OF POSTMENOPAUSAL OSTEOPOROSIS
- I-188 TREATMENT OF ACUTE SINUSITIS AND ACUTE EXACERBATION OF CHRONIC SINUSITIS
- I-189 TREATMENT OF ACUTE OTITIS MEDIA IN PEDIATRIC PATIENTS
- I-190 PLANAR IMAGING AS A SECOND LINE DIAGNOSTIC DRUG AFTER MAMMOGRAPHY TO ASSIST IN THE EVALUATION OF BREAST LESIONS IN PATIENTS WITH AN ABNORMAL MAMMOGRAM OR A PALPABLE BREAST MASS
- I-191 ENDOMETRIAL THINNING AGENT PRIOR TO ENDOMETRIAL ABLATION FOR DYSFUNCTIONAL UTERINE BLEEDING
- I-192 THE PREVENTION OF DEEP VEIN THROMBOSIS, WHICH MAY LEAD TO PULMONARY EMBOLISM, IN PATIENTS UNDERGOING ABDOMINAL SURGERY WHO ARE AT RISK FOR THROMBOEMBOLIC COMPLICATIONS AND A NEW DOSAGE REGIMEN, 40MG ONCE DAILY, FOR THIS INDICATION
- I-193 TREATMENT OF PANIC DISORDER IN A RECOMMENDED DOSE RANGE OF 50 TO 200MG/DAY
- I-194 CONGESTIVE HEART FAILURE
- I-195 FOR USE OF LANSOPRAZOLE IN COMBINATION WITH CLARITHROMYCIN AND AMOXICILLIN FOR THE ERADICATION OF HELICOBACTER PYLORI IN PATIENTS WITH ACTIVE DUODENAL ULCER DISEASE OR A ONE-YEAR HISTORY OF DUODENAL ULCER
- I-196 ACUTE TREATMENT OF ACTIVE BENIGN GASTRIC ULCER
- I-197 MAINTENANCE OF HEALING OF DUODENAL ULCER
- I-198 FOR THE USE OF LANSOPRAZOLE IN COMBINATION WITH AMOXICILLIN FOR THE ERADICATION OF HELICOBACTER PYLORI IN PATIENTS WITH ACTIVE DUODENAL ULCER DISEASE OR A ONE-YEAR HISTORY OF A DUODENAL ULCER
- I-199 MONOTHERAPY AND COMBINATION THERAPY WITH SULFONYL UREAS IN THE TREATMENT OF TYPE II DIABETES
- I-200 TREATMENT OF TINEA (PITYRIASIS) VERSICOLOR
- I-201 EMPIRICAL THERAPY FOR FEBRILE NEUTROPENIC PATIENTS
- I-202 SECOND-LINE TREATMENT OF AIDS-RELATED KAPOSI'S SARCOMA
- I-203 MAINTENANCE OF REMISSION OF ULCERATIVE COLITIS
- I-204 USE IN PEDIATRIC PATIENTS BETWEEN THE AGES OF 6 AND 11 FOR THE TREATMENT OF THE NASAL SYMPTOMS OF SEASONAL AND PERENNIAL ALLERGIC RHINITIS

PATENT AND EXCLUSIVITY TERMS

ADB 8 of 35

EXCLUSIVITY INDICATION

- I-205 INITIAL ANTICONVULSANT TREATMENT OF STATUS EPILEPTICUS
- I-206 TREATMENT OF EDEMA ASSOCIATED WITH CHRONIC RENAL FAILURE
- I-207 FOR THE SUPPRESSION OF RECURRENT EPISODES OF GENITAL HERPES IN IMMUNOCOMPETENT ADULTS
- I-208 TREATMENT OF OBSESSIVE COMPULSIVE DISORDER IN THE PEDIATRIC POPULATION
- I-209 PAROXYSMAL SUPRAVENTRICULAR TACHYCARDIA (PSVT)
- I-210 TO SLOW THE PROGRESSION OF CORONARY ATHEROSCLEROSIS IN PATIENTS WITH CORONARY HEART DISEASE AS PART OF A TREATMENT STRATEGY TO LOWER TOTAL AND LDL CHOLESTEROL TO TARGET LEVELS
- I-211 FOR USE IN PEDIATRIC POPULATION
- I-212 TREATMENT OF SYMPTOMS OF DRY MOUTH IN PATIENTS WITH SJOGREN'S SYNDROME
- I-213 TEMPORARY RELIEF OF PAIN AND PHOTOPHOBIA IN PATIENTS UNDERGOING CORNEAL REFRACTIVE SURGERY
- I-214 TREATMENT OF OSTEOPOROSIS
- I-215 PRE-PROCEDURAL APPLICATION TO ADULT MALE GENITAL SKIN PRIOR TO SITE-SPECIFIC SUBCUTANEOUS INFILTRATION WITH LIDOCAINE FOR THE REMOVAL OF GENITAL WARTS
- I-216 FOR THE LONG-TERM TWICE-DAILY (MORNING AND EVENING) ADMINISTRATION IN THE MAINTENANCE TREATMENT OF BRONCHOSPASM ASSOCIATED WITH COPD, INCLUDING CHRONIC BRONCHITIS AND EMPHYSEMA
- I-217 PREVENTION (DURING AND FOLLOWING HOSPITALIZATION) OF DEEP VEIN THROMBOSIS, WHICH MAY LEAD TO PULMONARY EMBOLISM, IN PATIENTS UNDERGOING HIP REPLACEMENT SURGERY
- I-218 USE OF LIPITOR AS AN ADJUNCTIVE THERAPY TO DIET FOR THE TREATMENT OF PATIENTS WITH ELEVATED SERUM TRIGLYCERIDE LEVELS (FREDERICKSON TYPE IV)
- I-219 USE OF LIPITOR BY PATIENTS WITH PRIMARY DYSBETALIPOPROTEINEMIA (FREDERICKSON TYPE III) WHO DO NOT RESPOND ADEQUATELY TO DIET
- I-220 TREATMENT OF EPISODIC- HEARTBURN, ACID INDIGESTION AND SOUR STOMACH
- I-221 TREATMENT OF BENIGN PROSTATIC HYPERPLASIA (BPH) IN MEN WITH AN ENLARGED PROSTATE TO IMPROVE SYMPTOMS, REDUCE THE RISK OF ACUTE URINARY RETENTION AND REDUCE THE RISK OF THE NEED OF SURGERY
- I-222 PREVENTION OF ISCHEMIC COMPLICATIONS OF UNSTABLE ANGINA AND NON-Q-WAVE MYOCARDIAL INFARCTION, WHEN CONCURRENTLY ADMINISTERED WITH ASPIRIN
- I-223 USE IN THE SYMPTOMATIC RELIEF OF RHINORRHEA ASSOCIATED WITH ALLERGIC AND NONALLERGIC-PERENNIAL RHINITIS IN CHILDREN AGE 6-11 YEARS
- I-224 FOR THE USE IN PEDIATRIC PATIENTS 4 TO 11 YEARS OF AGE FOR THE MANAGEMENT OF THE NASAL SYMPTOMS OF SEASONAL AND PERENNIAL ALLERGIC RHINITIS
- I-225 USE IN PATIENTS WITH PREVIOUS MI AND NORMAL CHOLESTEROL LEVELS, TO REDUCE RISK OF RECURRENT MI, MYOCARDIAL REVASCULARIZATION, AND CEREBROVASCULAR DISEASE EVENTS
- I-226 FIRST-LINE THERAPY FOR THE TREATMENT OF ADVANCED CARCINOMA OF THE OVARY IN COMBINATION WITH CISPLATIN
- I-227 SHORT-TERM TREATMENT OF SYMPTOMATIC GASTROESOPHAGEAL REFLUX DISEASE (GERD)
- I-228 PREVENTION OF MEAL INDUCED HEARTBURN AT A DOSE OF 75MG TAKEN 30-60MIN PRIOR TO A MEAL
- I-229 PRILOSEC (OMEPRazole), AMOXICILLIN, AND CLARITHROMYCIN FOR THE ERADICATION OF H. PYLORI IN PATIENTS WITH DUODENAL ULCER DISEASE
- I-230 IN COMBINATION WITH CIS-PLATIN, FOR THE FIRST LINE TREATMENT OF NON-SMALL CELL LUNG CANCER IN PATIENTS WHO ARE NOT CANDIDATES FOR POTENTIALLY CURATIVE SURGERY AND/OR RADIATION
- I-231 TREATMENT OF PATIENTS WITH LOCALLY ADVANCED OR METASTATIC BREAST CANCER AFTER FAILURE OF PRIOR CHEMOTHERAPY
- I-232 TREATMENT OF RECURRENT MUCOCUTANEOUS HERPES SIMPLEX INFECTIONS IN HIV-AFFECTED PATIENTS AT A DOSE OF 500MG TWICE DAILY
- I-233 PROPHYLACTIC USE TO REDUCE PERIOPERATIVE BLOOD LOSS AND THE NEED FOR BLOOD TRANSFUSION IN PATIENTS UNDERGOING CARDIOPULMONARY BYPASS IN THE COURSE OF CORONARY ARTERY BYPASS GRAFT SURGERY
- I-234 FOR USE IN COMBINATION WITH CISPLATIN FOR THE FIRST-LINE TREATMENT OF PATIENTS WITH INOPERABLE LOCALLY ADVANCED (STAGE IIIA OR IIIB) OR METASTATIC (STAGE IV) NON-SMALL CELL LUNG CANCER
- I-235 PREVENTION OF EXERCISE-INDUCED BRONCHOSPASM IN PATIENTS 12 YEARS OF AGE AND OLDER
- I-236 PREVENTION OF EXERCISE-INDUCED BRONCHOSPASM IN PATIENTS 4 YEARS OF AGE AND OLDER
- I-237 MAINTENANCE TREATMENT OF ASTHMA AND PREVENTION OF BRONCHOSPASM IN PATIENTS 4 YEARS OF AGE AND OLDER
- I-238 ADJUNCTIVE TREATMENT OF LENNOX-GASTAUT SYNDROME IN PEDIATRIC AND ADULT PATIENTS
- I-239 TREATMENT OF PATIENTS WITH HOMOZYGOUS FAMILIAL HYPERCHOLESTEROLEMIA
- I-240 MANAGEMENT OF SECONDARY HYPERPARATHYROIDISM AND RESULTANT METABOLIC BONE DISEASE IN PATIENTS WITH MODERATE TO SEVERE CHRONIC RENAL FAILURE (CCR 15 TO 55ML/MIN) NOT YET ON DIALYSIS
- I-241 USE IN PHOTODYNAMIC THERAPY (PDT) FOR REDUCTION OF OBSTRUCTION AND PALLIATION OF SYMPTOMS IN PATIENTS WITH COMPLETELY OR PARTIALLY OBSTRUCTING ENDOBRONCHIAL NONSMALL CELL LUNG CANCER

PATENT AND EXCLUSIVITY TERMS

ADB 9 of 35

EXCLUSIVITY INDICATION

- I-242 TREATMENT OF MODERATE TO SEVERE VASOMOTOR SYMPTOMS ASSOCIATED WITH THE MENOPAUSE AND IN THE TREATMENT OF VULVAR AND VAGINAL ATROPHY IN WOMEN WITH AN INTACT UTERUS
- I-243 USE IN THE SYMPTOMATIC RELIEF OF RHINORRHEA ASSOCIATED WITH THE COMMON COLD IN CHILDREN AGE 5 TO 11 YEARS
- I-244 REDUCE THE INCIDENCE OF BREAST CANCER IN WOMEN AT HIGH RISK FOR BREAST CANCER
- I-245 TREATMENT OF ACUTE SINUSITIS
- I-246 TREATMENT OF UNCOMPLICATED URINARY TRACT INFECTIONS
- I-247 USE IN CONVERSION TO MONOTHERAPY IN ADULTS WITH PARTIAL SEIZURES WHO ARE RECEIVING TREATMENT WITH A SINGLE ENZYME-INDUCING ANTIEPILEPTIC DRUG
- I-248 INPATIENT TREATMENT OF ACUTE DEEP VEIN THROMBOSIS WITH/WITHOUT PULMONARY EMBOLISM WHEN ADMIN WITH WARFARIN SODIM AND OUTPATIENT TREATMENT OF ACUTE DEEP VEIN THROMBOSIS WITHOUT PULMONARY EMBOLISM WHEN ADMIN WITH WARFARIN SODIUM
- I-249 TREATMENT OF CHRONIC HEPATITIS C IN PATIENTS WITH COMPENSATED LIVER DISEASE PREVIOUSLY UNTREATED WITH ALPHA INTERFERON THERAPY
- I-250 PRIMARY PREVENTION OF CORONARY HEART DISEASE IN PATIENTS WITHOUT SYMPATOMATIC CARDIOVASCULAR DISEASE WHO HAVE AVERAGE TO MODERATELY ELEVATED TOTAL-C AND LDL-C AND BELOW AVERAGE HDL-C
- I-251 TREATMENT OF GENERALIZED ANXIETY DISORDER
- I-252 NEW COMBINATION USE OF PRECOSE FOR PATIENTS WITH TYPE 2 DIABETES TREATED WITH DIET PLUS METFORMIN
- I-253 COMBINATION USE OF PRECOSE FOR PATIENTS WITH TYPE 2 DIABETES TREATED WITH DIET PLUS INSULIN
- I-254 PREVENTION OF POSTMENOPAUSAL OSTEOPOROSIS (LOSS OF BONE MASS)
- I-255 PREVENTION OF PNEUMOCYSTIS CARINII PNEUMONIA (PCP)
- I-256 USE IN TREATMENT OF SMALL CELL LUNG CANCER SENSITIVE DISEASE AFTER FAILURE OF FIRST-LINE CHEMOTHERAPY
- I-257 TREATMENT OF CHRONIC HEPATITIS B ASSOCIATED WITH EVIDENCE OF HEPATITIS B VIRAL REPLICATION AND ACTIVE LIVER INFLAMATION
- I-258 FOR PERENNIAL NONALLERGIC RHINITIS FOR AGES 4 AND ABOVE
- I-259 PROPHYLAXIS OF DEEP VEIN THROMBOSIS (DVT), WHICH MAY LEAD TO PULMONARY EMBOLISM, IN PATIENTS UNDERGOING HIP REPLACEMENT SURGERY
- I-260 EXPANDED PEDIATRIC USE IN CHILDREN YOUNGER THAN ONE MONTH OF AGE TO BIRTH (WITH A GESTATIONAL AGE OF 37 WEEKS OR GREATER)
- I-261 TREATMENT OF SOCIAL ANXIETY DISORDER
- I-262 TREATMENT OR PREVENTION OF BRONCHOSPASM WITH REVERSIBLE OBSTRUCTIVE AIRWAY DISEASE AND FOR THE PREVENTION OF EXERCISE INDUCED BRONCHOSPASM IN CHILDREN AGES 4-12
- I-263 TREATMENT OF UNSTABLE ANGINA AND NON-Q-WAVE MYOCARDIAL INFARCTION FOR THE PREVENTION OF ISCHEMIC COMPLICATIONS IN PATIENTS ON CONCURRENT ASPIRIN THERAPY
- I-264 PREVENTION OF NAUSEA AND VOMITING ASSOCIATED WITH RADIATION, INCLUDING TOTAL BODY IRRADIATION (TBI) AND FRACTIONATED ABDOMINAL RADIATION
- I-265 TREATMENT OF ATOPIC DERMATITIS IN PEDIATRIC PATIENTS 6 YEARS AND OLDER
- I-266 USE OF TOPAMAX AS ADJUNCTIVE THERAPY IN PEDIATRIC PATIENTS AGES 2-16 YEARS WITH PARTIAL ONSET SEIZURES
- I-267 USE IN PEDIATRIC PATIENTS 3 MONTHS OLD AND OLDER - FOR CORTICOSTEROID-RESPONSIVE DERMATOSES
- I-268 PROPHYLAXIS AND CHRONIC TREATMENT OF ASTHMA IN PATIENTS 7-11 YEARS OF AGE
- I-269 PREVENTION OF NAUSEA AND VOMITING ASSOCIATED WITH HIGHLY EMETOGENIC CANCER CHEMOTHERAPY, INCLUDING CISPLATIN
- I-270 ADJUVANT TREATMENT OF NODE-POSITIVE BREAST CANCER ADMINSTRERED SEQUENTIALLY TO STANDARD DOXORUBICIN-CONTAINING COMBINATION CHEMOTHERAPY
- I-271 TREATMENT OF OSTEOPOROSIS IN POSTMENOPAUSAL WOMEN
- I-272 TREATMENT OF GLUCOCORTICOID-INDUCED OSTEOPOROSIS IN MEN AND WOMEN RECEIVING GLUCOCORTICIDS IN A DAILY DOSE EQUIVALENT TO 7.5MG OR GREATER OF PREDNISONE AND WHO HAVE LOW BONE MINERAL DENSITY
- I-273 ADJUNCT TO DIET TO INCREASE HDL-C IN PATIENTS WITH PRIMARY HYPERCHOLESTEROLEMIA (HETEROZYGEOUS FAMILIAL AND NON FAMILIAL) AND MIXED DYSLIPIDEMIA (FREDERICKSON TYPES IIA AND IIB)
- I-274 USE OF TOPAMAX AS ADJUNCTIVE THERAPY IN THE TREATMENT OF PRIMARY GENERALIZED TONIC-CLONIC SEIZURES
- I-275 USE IN COMBINATION WITH METFORMIN AND SULFONYLUREA IN PATIENTS WITH TYPE 2 DIABETES
- I-276 USE OF REZULIN IN COMBINATION WITH METFORMIN AND SULFONYLUREAS IN PATIENTS WITH TYPE 2 DIABETES
- I-277 TREATMENT OF TYPE III HYPERLIPOPROTEINEMIA
- I-278 TREATMENT OF PATIENTS WITH ISOLATED HYPERTRIGLYCERIDEMIA (FREDERICKSON TYPE IV)
- I-279 TREATMENT OF POST-TRAUMATIC STRESS DISORDER
- I-280 USE OF CARNITOR INJECTION FOR THE PREVENTION AND TREATMENT OF CARNITINE DEFICIENCY IN PATIENTS WITH END STAGE RENAL DISEASE WHO ARE UNDERGOING DIALYSIS

PATENT AND EXCLUSIVITY TERMS

ADB 10 of 35

EXCLUSIVITY INDICATION

- I-281 INCREASING HDL-C IN PATIENTS WITH PRIMARY HYPERCHOLESTEROLEMIA (HETEROZYGOUS FAMILIAL AND NONFAMILIAL) AND MIXED DYSLIPIDEMIA (FREDERICKSON TYPES IIA AND IIB)
- I-282 TREATMENT OF PATIENTS WITH LOCALLY ADVANCED OR METASTATIC NON-SMALL CELL LUNG CANCER AFTER FAILURE OF PRIOR PLATINUM-BASED CHEMOTHERAPY
- I-283 TO REDUCE THE INCIDENCE OF MODERATE TO SEVERE XEROSTOMIA IN PATIENTS UNDERGOING POST-OPERATIVE RADIATION TREATMENT FOR HEAD AND NECK CANCER, WHERE THE RADIATION PORT INCLUDES A SUBSTANTIAL PORTION OF THE PAROTID GLANDS
- I-284 TO REDUCE THE NUMBER OF ADENOMATOUS COLORECTAL POLYPS IN FAMILIAL ADENOMATOUS POLYPOSIS PATIENTS AS AN ADJUNCT TO USUAL CARE
- I-285 TREATMENT OF NASAL SYMPTOMS OF SEASONAL AND PERENNIAL RHINITIS IN ADULTS AND CHILDREN 3 YEARS OF AGE AND OLDER
- I-286 TREATMENT OF PATIENTS WITH FREDERICKSON TYPE III
- I-287 USE OF PRAVASTATIN IN PATIENTS WITH EVIDENT CORONARY HEART DISEASE TO REDUCE THE RISK OF TOTAL MORTALITY BY REDUCING CORONARY DEATH
- I-288 CHANGES IN SEVERAL SECTIONS OF THE INSERT TO INCORPORATE STATEMENTS CONCERNING THE USE OF HIGH DOSES OF LISINAPRIL TO REDUCE THE RISK OF THE COMBINED OUTCOMES OF MORTALITY AND HOSPITALIZATION IN PATIENTS WITH CONGESTIVE HEART FAILURE
- I-289 USE OF AVANDIA IN COMBINATION WITH A SULFONYLUREA IN PATIENTS WITH TYPE 2 DIABETES MELLITUS WHEN DIET AND EXERCISE WITH EITHER SINGLE AGENT DOES NOT ACHIEVE ADEQUATE GLYCEMIC CONTROL
- I-290 PREVENTION OF CORTICOSTEROID-INDUCED OSTEOPOROSIS
- I-291 PREVENTION OF POSTMENOPAUSAL OSTEOPOROSIS
- I-292 TREATMENT OF POSTMENOPAUSAL OSTEOPOROSIS
- I-293 TREATMENT OF CORTICOSTEROID-INDUCED OSTEOPOROSIS
- I-294 TREATMENT OF UNCOMPLICATED ACUTE ILLNESS DUE TO INFLUENZA A AND B IN PEDIATRIC PATIENTS 7 YEARS AND OLDER WHO HAVE BEEN SYMPTOMATIC FOR NO MORE THAN 2 DAYS
- I-295 PREVENTION OF POSTMENOPAUSAL OSTEOPOROSIS FOR WOMEN WITH AN INTACT UTERUS
- I-296 LONG-TERM INTRAVENOUS TREATMENT OF PULMONARY HYPERTENSION ASSOCIATED WITH THE SCLERODERMA SPECTRUM OF DISEASE IN NYHA CLASS III AND CLASS IV PATIENTS WHO DO NOT RESPOND TO CONVENTIONAL THERAPY
- I-297 SHORT-TERM TREATMENT OF ACUTE MANIC EPISODES ASSOCIATED WITH BIPOLAR I DISORDER
- I-298 TREATMENT OF PATIENTS WITH FREDERICKSON TYPE IIA AND IIB HYPERLIPOPROTEINEMIA
- I-299 USE OF CAMPTOSAR AS A COMPONENT OF FIRST-LINE THERAPY IN COMBINATION WITH 5-FLUOROURACIL AND LEUCOVARIN FOR PATIENTS WITH METASTATIC CARCINOMA OF THE COLON OR RECTUM
- I-300 PROPHYLAXIS FOR ASTHMA IN CHILDREN 2-5 YEARS OF AGE
- I-301 TREATMENT OF SIGNS AND SYMPTOMS OF ALLERGIC CONJUNCTIVITIS
- I-302 TREATMENT OF PEDIATRIC PATIENTS WITH PRADER-WILLI SYNDROME
- I-303 INCREASING HDL-CHOLESTEROL IN PATIENTS WITH PRIMARY HYPERCHOLESTEROLEMIA AND MIXED DYSLIPIDEMIAS
- I-304 TREATMENT OF PATIENTS WITH FREDERICKSON TYPE IV
- I-305 TREATMENT OF LEVOFLOXACIN SUSCEPTIBLE STRAINS OF PENICILLIN-RESISTANT STREPTOCOCCUS PNEUMONIAE IN PATIENTS WITH COMMUNITY ACQUIRED PNEUMONIA
- I-306 INDUCTION OF SPERMATOGENESIS IN MEN WITH PRIMARY AND SECONDARY HYPOGONADOTROPIC HYPOGONADISM IN WHOM THE CAUSE OF INFERTILITY IS NOT DUE TO PRIMARY TESTICULAR FAILURE
- I-307 NEW COMBINATION USE OF METFORMIN AND INSULIN IN TYPE 2 DIABETES
- I-308 TREATMENT OF PEDIATRIC PATIENTS WITH POLYARTICULAR COURSE JUVENILE RHEUMATOID ARTHRITIS WHO RESPONDED INADEQUATELY TO SALICYLATES OR OTHER NSAIDS
- I-309 USE OF ACTONEL 35MG ONCE A WEEK TO INCREASE BONE MASS IN MEN WITH OSTEOPOROSIS
- I-310 REDUCTION IN RISK OF MYOCARDIAL INFARCTION, STROKE, AND DEATH FROM CARDIOVASCULAR CAUSES
- I-311 ADJUNCTIVE THERAPY IN THE TREATMENT OF PARTIAL SEIZURES IN PEDIATRIC PATIENTS AGE 3 TO 12 YEARS
- I-312 FIRST LINE TREATMENT OF POSTMENOPAUSAL WOMEN WITH HORMONE RECEPTOR POSITIVE OR HORMONE RECEPTOR UNKNOWN LOCALLY ADVANCED OR METASTATIC BREAST CANCER
- I-313 EXTENSION OF INDICATION TO PROVIDE FOR MAINTENANCE OF RESPONSE
- I-314 TOPICAL ANESTHETIC FOR SUPERFICIAL MINOR SURGERY OF GENITAL MUCOUS MEMBRANES AND AS AN ADJUNCT FOR LOCAL INFILTRATION ANESTHESIA IN GENITAL MUCOUS MEMBRANES
- I-315 THROMBOPROPHYLAXIS OF DEEP VEIN THROMBOSIS, WHICH MAY LEAD TO PULMONARY EMBOLISM, IN MEDICAL PATIENTS WHO ARE AT RISK FOR THROMBOEMBOLIC COMPLICATIONS DUE TO SEVERELY RESTRICTED MOBILITY DURING ACUTE ILLNESS
- I-316 TREATMENT OF NSAID-ASSOCIATED GASTRIC ULCER PATIENTS WHO CONTINUE NSAID USE AND REDUCING RISK OF NSAID-ASSOCIATED GASTRIC ULCERS IN PATIENTS WITH HISTORY OF DOCUMENTED GASTRIC ULCER WHO REQUIRE USE OF AN NSAID
- I-317 PROPHYLAXIS OF INFLUENZA IN ADULTS AND ADOLESCENTS 13 YEARS AND OLDER
- I-318 FIRSTLINE TREATMENT OF POSTMENOPAUSAL WOMEN WITH HORMONE RECEPTOR POSITIVE OR HORMONE RECEPTOR UNKNOWN LOCALLY ADVANCED OR METASTATIC BREAST CANCER
- I-319 USE FOR SUSPECTED OR CONFIRMED METHANOL POISONING, EITHER ALONE OR IN COMBINATION WITH HEMODIALYSIS

PATENT AND EXCLUSIVITY TERMS

ADB 11 of 35

EXCLUSIVITY INDICATION

- I-320 TREATMENT OF TYPE 2 DIABETES IN PEDIATRIC PATIENTS (AGES 10-16 YEARS)
- I-321 JUVENILE RHEUMATOID ARTHRITIS
- I-322 USE OF DIPRIVAN IN PATIENTS 3 MONTHS TO 16 YEARS
- I-323 COLORECTAL CANCER
- I-324 REDUCING NEUROLOGIC DISABILITY AND/OR FREQUENCY OF CLINICAL RELAPSES IN PATIENTS WITH SECONDARY (CHRONIC) PROGRESSIVE, PROGRESSIVE RELAPSING, OR WORSENING RELAPSING-REMITTING MULTIPLE SCLEROSIS
- I-325 PREVENTION OF RELAPSE AND RECURRENCE OF DEPRESSION
- I-326 GENERALIZED ANXIETY DISORDER
- I-327 SYMPTOMATIC RELIEF OF RHINORRHEA ASSOCIATED WITH SEASONAL ALLERGIC RHINITIS IN PATIENTS 5 YEARS AND OLDER
- I-328 PROPHYLAXIS AND CHRONIC TREATMENT OF ASTHMA IN PATIENTS 5-6 YEARS OF AGE
- I-329 UNCOMPLICATED SKIN AND SKIN STRUCTURE INFECTIONS
- I-330 MAINTENANCE OF HEALING OF EROSIIVE ESOPHAGITIS AND CONTROL OF DAYTIME AND NIGHTTIME HEARTBURN SYSTOMS IN PATIENTS WITH GERD
- I-331 TREATMENT OF MODERATE ACNE VULGARIS
- I-332 EMPIRIC THERAPY IN FEBRILE NEUTROPENIC PATIENTS WITH SUSPECTED FUNGAL INFECTIONS (EFTN)
- I-333 TOPICAL TREATMENT OF TINEA (PITYRIASIS) VERSICOLOR DUE TO MALASSEZIA FURFUR (FORMERLY PITYROSPORUM ORBICULARE)
- I-334 LONG-TERM TREATMENT OF GROWTH FAILURE IN CHILDREN BORN SMALL FOR GESTATIONAL AGE WHO FAIL TO MANIFEST CATCH-UP GROWTH BY TWO YEARS OF AGE
- I-335 ADJUNCTIVE THERAPY IN PATIENTS TWO YEARS AND OLDER WITH SEIZURES ASSOCIATED WITH LENNOX-GASTAUT SYNDROME
- I-336 EXPANSION OF INDICATION TO INCLUDE THE TREATMENT OF PATIENTS WITH PREDOMINATELY CLASSIC SUBFOVEAL CHOROIDAL NEOVASCULARIZATION DUE TO PATHOLOGIC MYOPIA OR PRESUMED OCULAR HISTOPLASMOSIS
- I-337 PATHOLOGICAL HYPERSECRETION ASSOCIATED WITH ZOLLINGER-ELLISON SNYDROME
- I-338 MANAGEMENT OF ACUTE PAIN IN ADULTS AND TREATMENT OF PRIMARY DYSMENORRHEA
- I-339 TREATMENT OF HEPATITIS B IN PEDIATRIC PATIENTS AGES 2-17 YEARS
- I-340 ATOPIC DERMATITIS IN PEDIATRIC PATIENTS AGES 2-5
- I-341 BREAST CANCER COMBINATION THERAPY
- I-342 USE OF FORADIL FOR LONG-TERM, TWICE DAILY (MORNING AND EVENING) ADMINISTRATION IN THE MAINTENANCE TREATMENT OF BRONCHO-CONSTRICTION IN PATIENTS WITH COPD INCLUDING CHRONIC BRONCHITIS AND EMPHYSEMA
- I-343 USE OF COREG FOR SEVERE HEART FAILURE
- I-344 ACNE VULGARIS
- I-345 TREATMENT OF POSTTRAUMATIC STRESS DISORDER
- I-346 TREATMENT OF SYMPTOMATIC GASTRO ESOPHAGEAL REFLUX DISEASE (GERD)
- I-347 TREATMENT OR PREVENTION OF BRONCHOSPASM IN CHILDREN 6 YEARS OF AGE AND OLDER WITH OBSTRUCTIVE AIRWAY DISEASE
- I-348 LONG-TERM, TWICE-DAILY (MORNING AND EVENING) ADMINISTRATION IN THE MAINTENANCE TREATMENT OF BRONCHOSPASM ASSOCIATED WITH COPD (INCLUDING EMPHYSEMA AND CHRONIC BRONCHITIS)
- I-349 ACUTE CORONARY SYNDROME
- I-350 TREATMENT OF HETEROZYGOUS FAMILIAL HYPERCHOLESTEROLEMIA IN ADOLESCENT BOYS AND GIRLS AT LEAST ONE YEAR POSTMENARCHAL, AGES 10 TO 17 YEARS, WITH A RECOMMENDED DOSING RANGE OF 10 TO 40MG ONCE DAILY
- I-351 PREVENTION OF POSTMENOPAUSAL OSTEOPOROSIS FOR ALL STRENGTHS
- I-352 ANTICOAGULANT IN PATIENTS WITH OR AT RISK FOR HEPARIN-INDUCED THROMBOCYTOPENIA UNDERGOING PERCUTANEOUS CORONARY INTERVENTIONS (PCI)
- I-353 TREATMENT OF SIGNS AND SYMPTOMS OF RHEUMATOID ARTHRITIS
- I-354 MANAGEMENT OF POST HERPETIC NEURALGIA
- I-355 PREMENSTRUAL DYSPHORIC DISORDER
- I-356 TREATMENT OF PATHOLOGICAL HYPERSECRETORY CONDITIONS, INCLUDING ZOLLINGER-ELLISON SYNDROME
- I-357 TREATMENT OF COMPLICATED SKIN AND SKIN STRUCTURE INFECTIONS
- I-358 TREATMENT OF PANIC DISORDER
- I-359 TREATMENT OF VULVAR AND VAGINAL ATROPHY ASSOCIATED WITH THE MENOPAUSE
- I-360 TREATMENT OF NASAL SYMPTOMS OF SEASONAL AND PERENNIAL RHINITIS IN CHILDREN AGES TWO UP TO AGE THREE
- I-361 TREATMENT OF MULTIPLE MYELOMA AND DOCUMENTED BONE METASTASES FROM SOLID TUMORS, IN CONJUNCTION WITH STANDARD ANTINEOPLASTIC THERAPY. PROSTATE CANCER SHOULD HAVE PROGRESSED AFTER TREATMENT WITH AT LEAST ONE HORMONAL THERAPY
- I-362 TREATMENT OF PANIC DISORDER, WITH OR WITHOUT AGORAPHOBIA
- I-363 ADJUVANT TREATMENT OF POST MENOPAUSAL WOMEN WITH HORMONE RECEPTOR POSITIVE EARLY BREAST CANCER
- I-364 TREATMENT OF COMMUNITY-ACQUIRED PNEUMONIA IN ADULTS

PATENT AND EXCLUSIVITY TERMS

ADB 12 of 35

EXCLUSIVITY INDICATION

- I-365 TREATMENT OF HEART FAILURE (NYHA CLASS II-IV) IN PATIENTS WHO ARE INTOLERANT TO AN ACE INHIBITOR
- I-366 PREVENTION OF RELAPSE FOLLOWING LONG-TERM TREATMENT OF MAJOR DEPRESSIVE DISORDER
- I-367 COMBINATION THERAPY WITH THIAZOLIDINEDIONE TO LOWER BLOOD GLUCOSE IN PTS WHOSE HYPERGLYCEMIA CANNOT BE CONTROLLED BY DIET/EXERCISE PLUS MONOTHERAPY WITH ANY OF THE FOLLOWING AGENTS:METFORMIN,SULFONYLUREAS,REPAGLINIDE,OR THIAZOLIDINEDIONES
- I-368 USE OF GLUCOVANCE WITH A THIAZOLIDINEDIONE WHEN GLYCEMIC CONTROL IS NOT OBTAINED WITH GLUCOVANCE ALONE
- I-369 PREVENTION AND TREATMENT OF POSTOPERATIVE NAUSEA AND VOMITING
- I-370 TREATMENT OF HETEROZYGOUS FAMILIAL HYPERCHOLESTEROLEMIA IN CHILDREN, AGES 8-13 YEARS, WITH RECOMMENDED DOSE OF 20MG ONCE DAILY AND IN ADOLESCENTS, AGES 14-18 WITH A RECOMMENDED DOSE OF 40MG ONCE DAILY
- I-371 HELICOBACTER PYLORI ERADICATION TO REDUCE THE RISK OF DUODENAL ULCER RECURRENCE
- I-372 NOSOCOMIAL PNEUMONIA
- I-373 TREATMENT OF TYPE 2 DIABETIC NEPHROPATHY
- I-374 SHORT TERM TOPICAL TREATMENT OF MILD TO MODERATE PLAQUE-TYPE PSORIASIS OF NON SCALP REGIONS
- I-375 FIRST LINE THERAPY FOR THE REDUCTION OF INTRAOCULAR PRESSURE IN PATIENTS WITH OPEN-ANGLE GLAUCOMA OR OCULAR HYPERTENSION
- I-376 TREATMENT OF NEWLY DIAGNOSED ADULT PATIENTS WITH PHILADELPHIA CHROMOSOME POSITIVE CHRONIC MYELOID LEUKEMIA (CML)
- I-377 USE OF BRAVELLE FOR MULTIPLE FOLLICULAR DEVELOPMENT (CONTROLLED OVARIAN STIMULATION) DURING ASSISTED REPRODUCTIVE TECHNOLOGY CYCLES IN PATIENTS WHO HAVE PREVIOUSLY RECEIVED PITUITARY SUPPRESSION
- I-378 RELIEF OF SYMPTOMS OF SEASONAL ALLERGIC RHINITIS IN ADULTS AND PEDIATRIC PATIENTS 2 YEARS OF AGE AND OLDER
- I-379 USE TAXOTERE IN COMBINATION WITH CISPLATIN FOR THE TREATMENT OF PATIENTS WITH UNRESECTABLE, LOCALLY ADVANCED OR METASTATIC NON-SMALL CELL LUNG CANCER WHO HAVE NOT PREVIOUSLY RECEIVED CHEMOTHERAPY FOR THIS CONDITION
- I-380 TO TREAT PATIENTS WITH SCHIZOPHRENIA OR SCHIZOAFFECTIVE DISORDER AT RISK FOR EMERGENT SUICIDAL BEHAVIOR
- I-381 TREATMENT OF COLD SORES (HERPES LABIALIS) IN ADULT AND ADOLESCENT PATIENTS 12 YEARS OF AGE AND OLDER
- I-382 FOR NEWLY-DIAGNOSED HIGH GRADE MALIGNANT GLIOMA PATIENTS AS AN ADJUNCT TO SURGERY AND RADIATION
- I-383 TREATMENT OF TYPE 2 DIABETIC NEPHROPATHY
- I-384 USE IN COMBINATION WITH INSULIN FOR THE TREATMENT OF PATIENTS WITH TYPE 2 DIABETES MELLITUS
- I-385 MODIFICATION OF THE INDICATION FOR COMMUNITY ACQUIRED PNEUMONIA TO ADD"INCLUDING PENICILLIN-RESISTANT STRAINS, MIC PENICILLIN>=2MCG/ML TO STREPTOCOCCUS PNEUMONIAE
- I-386 RAPAMUNE (SIROLIMUS) WITHIN AN IMMUNOSUPPRESSIVE REGIMEN THAT WOULD ALLOW FOR THE WITHDRAWAL OF CYCLOSPORINE 2 TO 4 MONTHS AFTER RENAL TRANSPLANTATION IN PATIENTS CONSIDERED AT LOW TO MODERATE IMMUNOLOGIC RISK FOR RENAL TRANSPLANT REJECTION
- I-387 ADJUNCTIVE THERAPY OF PARTIAL SEIZURES IN PEDIATRIC PATIENTS GREATER THAT OR EQUAL TO 2 YEARS OF AGE
- I-388 TREATMENT OF PATIENTS WITH LEFT VENTRICULAR DYSFUNCTION FOLLOWING MYOCARDIAL INFARCTION
- I-389 SUPPRESSION OF RECURRENT GENITAL HERPES IN HIV-INFECTED INDIVIDUALS
- I-390 USE IN PTS AT HIGH RISK CORONARY EVENTS DUE TO EXISTING CORONARY HEART DISEASE,DIABETES,PERIPHERAL VESSEL DISEASE,STROKE HISTORY,OTHER CV DISEASE TO REDUCE RISK TOTAL MORTALITY BY REDUCING CORONARY DEATH,REDUCE NONFATAL MI & STROKE.....
- I-391 ABLATION OF HIGH-GRADE DYSPLASIA IN BARRETT'S ESOPHAGUS PATIENTS WHO DO NOT UNDERGO ESOPHAGECTOMY
- I-392 TX OF PED PATIENTS W/PH+ CHRONIC PHASE CML DISEASE RECUR AFTER STEM CELL TRNSPLT OR RESIST TO INTERFERON ALPHA THERAPY.NO CONTROLLED TRIALS DEMONSTRATING A CLINICAL BENEFIT SUCH AS IMPROVE IN DISEASE RELATED SX OR INCREASED SURVIVAL
- I-393 CHRONIC BACTERIAL PROSTATITIS
- I-394 USE IN PATIENTS WITH CORONARY HEART DISEASE TO REDUCE THE RISK OF UNDERGOING CORONARY REVASCULARIZATION PROCEDURES
- I-395 TO IMPROVE PHYSICAL FUNCTION
- I-396 EXPANDED INDICATION TO INCLUDE THE ASSESSMENT OF VENTRICULAR FUNCTION IN SUBJECTS BEING EVALUATED FOR HEART DISEASE AND/OR VENTRICULAR FUNCTION
- I-397 EXTENDED PROPHYLAXIS IN PATIENTS UNDERGOING HIP FRACTURE SURGERY
- I-398 IDIOPATHIC SHORT STATURE
- I-399 TREATMENT OF CANDIDEMIA AND THE FOLLOWING CANDIDA INFECTIONS: INTRA-ABDOMINAL ABSCESSSES, PERITONITIS AND PLEURAL SPACE INFECTIONS
- I-400 USE OF OLANZAPINE IN COMBINATION WITH LITHIUM OR VALPROATE FOR THE TREATMENT OF ACUTE MANIC EPISODES ASSOCIATED WITH BIPOLAR DISORDER

PATENT AND EXCLUSIVITY TERMS

ADB 13 of 35

EXCLUSIVITY INDICATION

- I-401 LONGER-TERM EFFICACY OF ARIPIPRAZOLE IN THE TREATMENT OF SCHIZOPHRENIA
- I-402 DIABETIC FOOT INFECTIONS WITHOUT CONCOMITANT OSTEOMYELITIS
- I-403 USE OF VALTREX IN COMBINATION WITH SAFER SEX PRACTICES FOR THE REDUCTION OF THE RISK OF TRANSMISSION OF GENITAL HERPES DURING SUPPRESSIVE THERAPY OF THE SOURCE PARTNER IN A HETEROSEXUAL COUPLE
- I-404 MAINTENANCE TREATMENT OF BIPOLAR I DISORDER TO DELAY THE TIME TO OCCURRENCE OF MOOD EPISODES (DEPRESSION, MANIA, HYPOMANIA, MIXED EPISODES) IN PATIENTS TREATED FOR ACUTE MOOD EPISODES WITH STANDARD THERAPY
- I-405 TREATMENT OF PREMENSTRUAL DYSPHORIC DISORDER (PMDD) USING AN INTERMITTENT DOSING REGIMEN
- I-406 PREVENTION OF CYTOMEGALOVIRUS DISEASE IN KIDNEY, HEART, AND KIDNEY-PANCREAS TRANSPLANT PATIENTS AT HIGH RISK (DONOR CMV SEROPOSITIVE/RECIPIENT CMV SERONEGATIVE)
- I-407 IMPROVE SURVIVAL OF STABLE PATIENTS WITH LEFT VENTRICULAR SYSTOLIC DYSFUNCTION (EJECTION FRACTION<=40%) AND CLINICAL EVIDENCE OF CONGESTIVE HEART FAILURE AFTER AN ACUTE MYOCARDIAL INFARCTION
- I-408 STIMULATION OF PANCREATIC SECRETIONS TO FACILITATE THE IDENTIFICATION OF THE AMPULLA OF VATER AND ACCESSORY PAPILLA DURING ENDOSCOPIC RETROGRADE CHOLANGIO-PANCREATOGRAPHY (ERCP)
- I-409 ESOPHAGEAL CANDIDIASIS
- I-410 USE OF ADVAIR DISKUS 250/50 FOR CHRONIC OBSTRUCTIVE PULMONARY DISEASE (COPD) ASSOCIATED WITH CHRONIC BRONCHITIS
- I-411 EXPANDED INDICATION FOR USE IN COMBINATION WITH ANTIDIABETIC DRUGS IN THE THIAZOLIDINEDIONE CLASS
- I-412 MONOTHERAPY FOR THE SHORT TERM TREATMENT OF ACUTE MANIC OR MIXED EPISODES ASSOCIATED WITH BIPOLAR I DISORDER
- I-413 ADJUNCTIVE THERAPY FOR THE SHORT TERM TREATMENT OF ACUTE MANIC OR MIXED EPISODES ASSOCIATED WITH BIPOLAR I DISORDER
- I-414 PROPHYLAXIS OF DEEP VEIN THROMBOSIS (DVT), WHICH MAY LEAD TO PULMONARY EMBOLISM (PE) IN MEDICAL PATIENTS WHO ARE AT RISK FOR THROMBOEMBOLIC COMPLICATIONS DUE TO SEVERELY RESTRICTED MOBILITY DURING ACUTE ILLNESS
- I-415 SEVERE HYPERTENSION WHEN THE VALUE OF ACHIEVING PROMPT BLOOD PRESSURE CONTROL EXCEEDS THE RISK OF INITIATING COMBINATION THERAPY
- I-416 THE USE OF CIPRO XR FOR COMPLICATED URINARY TRACT INFECTIONS AND ACUTE UNCOMPLICATED PYELONEPHRITIS
- I-417 USE IN THE LONG TERM TREATMENT OF BIPOLAR I DISORDER
- I-418 ADJUNCTIVE THERAPY W/ MOOD STABILIZERS (LITHIUM OR DIVALPROEX) IN THE TREATMENT OF ACUTE MANIC EPISODES ASSOCIATED WITH BIPOLAR I DISORDERS
- I-419 MONOTHERAPY IN THE TREATMENT OF ACUTE MANIC EPISODES ASSOCIATED WITH BIPOLAR I DISORDER
- I-420 TOPICAL TREATMENT OF CLINICALLY TYPICAL, NONHYPERKERATOTIC, NONHYPERTROPHIC ACTINIC KERATOSES ON THE FACE OR SCALP IN IMMUNOCOMPETENT ADULTS
- I-421 TREATMENT OF COMPLICATED URINARY TRACT INFECTIONS AND PYELONEPHRITIS DUE TO E.COLI FOR PED PATIENTS (1-17) NOT AS FIRST CHOICE
- I-422 INDICATED FOR THE IN-HOSPITAL SHORT-TERM (UP TO 4 HOURS) REDUCTION IN BLOOD PRESSURE IN PEDIATRIC PATIENTS
- I-423 ACUTE TREATMENT OF MIGRAINE ATTACKS WITH OR WITHOUT AURA IN ADULTS
- I-424 MANAGEMENT OF SECONDARY HYPERPARATHYROIDISM IN PATIENTS WITH MODERATE TO SEVERE CHRONIC RENAL INSUFFICIENCY NOT YET ON DIALYSIS
- I-425 FLOXATIN IN COMBINATION WITH INFUSIONAL 5-FLUOROURACIL (5-FU) AND LEUCOVORIN (LV) FOR THE TREATMENT OF PATIENTS PREVIOUSLY UNTREATED FOR ADVANCED COLORECTAL CANCER
- I-426 TREATMENT OF ACUTE PULMONARY EMBOLISM WHEN ADMINISTERED IN CONJUNCTION WITH WARFARIN SODIUM
- I-427 TREATMENT OF ACUTE DEEP VEIN THROMBOSIS WITHOUT PULMONARY EMBOLISM WHEN ADMINISTERED IN CONJUNCTION WITH WARFARIN SODIUM
- I-428 FOR USE IN COMBINATION WITH PACLITAXEL FOR THE FIRST-LINE TREATMENT OF PATIENTS WITH METASTATIC BREAST CANCER AFTER FAILURE OF PRIOR ANTHRACYCLINE CONTAINING ADJUVANT CHEMOTHERAPY UNLESS ANTHRACYCLINES WERE CLINICALLY CONTRAINDICATED
- I-429 FOR USE IN COMBINATION WITH PREDNISONE FOR THE TREATMENT OF PATIENTS WITH ANDROGEN INDEPENDENT (HORMONE REFRACTORY) METASTATIC PROSTATE CANCER
- I-430 FOR USE IN THE RELIEF OF THE SIGNS AND SYMPTOMS OF RHEUMATOID ARTHRITIS IN ADULTS
- I-431 NOSOCOMIAL PNEUMONIA AND COMMUNITY-ACQUIRED PNEUMONIA CAUSED BY STREPTOCOCCUS PNEUMONIAE INDICATION EXPANDED TO INCLUDE MULTI-DRUG RESISTANT STRAINS
- I-432 TREATMENT OF COMMUNITY ACQUIRED PNEUMONIA CAUSED BY MULTI-DRUG RESISTANT STREPTOCOCCUS PNEUMONIAE
- I-433 TREATMENT OF BIOPSY-CONFIRMED, PRIMARY SUPERFICIAL BASAL CELL CARCINOMA IN IMMUNOCOMPETENT ADULTS, WITH A MAXIMUM TUMOR DIAMETER OF 2.0CM, LOCATED ON THE TRUNK (EXCLUDING ANOGENITAL SKIN), NECK, OR EXTREMITIES (EXCLUDING HANDS AND FEET)
- I-434 PREVENTION OF CARDIOVASCULAR DISEASE IN ADULT PATIENTS WITHOUT CLINICALLY EVIDENT HEART DISEASE, BUT WITH MULTIPLE RISK FACTORS FOR CORONARY HEART DISEASE TO REDUCE RISK OF MI AND RISK FOR REVASCULARIZATION PROCEDURES AND ANGINA

PATENT AND EXCLUSIVITY TERMS

ADB 14 of 35

EXCLUSIVITY INDICATION

- I-435 CHRONIC IDIOPATHIC CONSTIPATION
- I-436 FOR USE IN COMBINATION WITH DOXORUBICIN AND CYCLOPHOSPHAMIDE FOR THE ADJUVANT TREATMENT OF PATIENTS WITH OPERABLE NODE-POSITIVE BREAST CANCER
- I-437 TREATMENT OF ACUTE MANIC AND MIXED EPISODES ASSOCIATED WITH BIPOLAR DISORDER
- I-438 EMPIRICAL THERAPY FOR PRESUMED FUNGAL INFECTIONS IN FEBRILE, NEUTROPENIC PATIENTS
- I-439 USED TO TREAT ADULTS WITH GROWTH HORMONE DEFICIENCY
- I-440 FOR THE REPLACEMENT OF ENDOGENOUS GROWTH HORMONE IN ADULTS WITH GROWTH HORMONE DEFICIENCY
- I-441 USE COMBINATION WITH INFUSIONAL 5-FU/LV FOR ADJUVANT TREATMENT STAGE III COLON CANCER PTS WHO HAVE UNDERGONE COMPLETE RESECTION PRIMARY TUMOR-BASED ON IMPROVEMENT IN DISEASE FREE SURVIVAL, NO DEMONSTRATED BENEFIT OVERALL SURVIVAL AFTER 4YRS
- I-442 USED FOR CANDIDEMIA IN NONNEUTROPENIC PATIENTS AND THE FOLLOWING CANDIDA INFECTIONS: DISSEMINATED INFECTIONS IN SKIN & INFECTIONS IN ABDOMEN, KIDNEY, BLADDER WALL, AND WOUNDS
- I-443 TREATMENT OF NASAL POLYPS IN PATIENTS 18 YEARS OF AGE AND OLDER
- I-444 USE OF PROTONIX IV FOR INJECTION AS STAND ALONE THERAPY FOR THE SHORT-TERM TREATMENT OF PATIENTS HAVING GASTROESOPHAGEAL REFLUX (GERD) WITH A HISTORY OF EROSIIVE ESOPHAGITIS
- I-445 TO IMPROVE (COMPARED TO 4.25% DEXTROSE) LONG-DWELL ULTRAFILTRATION AND CLEARANCE OF CREATININE AND UREA NITROGEN IN PATIENTS WITH HIGH AVERAGE OR GREATER TRANSPORT CHARACTERISTICS, AS DEFINED USING THE PERITONEAL EQUILIBRATION TEST (PET)
- I-446 EXTENDED ADJUVANT TREATMENT OF EARLY BREAST CANCER IN POSTMENOPAUSAL WOMEN WHO HAVE RECEIVED 5 YRS ADJUVANT TAMOXIFEN THERAPY-EFFECTIVENESS BASED ON AN ANALYSIS OF DISEASE FREE SURVIVAL IN PATIENTS TREATED FOR A MEDIAN 24 MONTHS
- I-447 USE OF COPEGUS (RIBAVIRIN) FOR TREATMENT OF CHRONIC HEPATITIS C IN ADULT PATIENTS COINFECTED WITH HIV IN COMBINATION WITH PEGASYS (PEGINTERFERON ALFA-2A)
- I-448 TREATMENT OF HEART FAILURE (NYHA CLASS II-IV AND EJECTION FRACTION <=40%) TO REDUCE THE RISK OF DEATH FROM CARDIOVASCULAR CAUSES AND TO REDUCE HOSPITALIZATIONS FOR HEART FAILURE
- I-449 TO IMPROVE WAKEFULNESS IN TWO NEW PATIENT POPULATIONS WITH EXCESSIVE SLEEPINESS: THOSE WITH OBSTRUCTIVE SLEEP APNEA/HYPOPNEA SYNDROME AND THOSE WITH SHIFT WORK SLEEP DISORDER
- I-450 TREATMENT OF PATIENTS WITH NEWLY DIAGNOSED HIGH GRADE GLIOMAS CONCOMITANTLY WITH RADIOTHERAPY AND THEN AS ADJUVANT TREATMENT
- I-451 MANAGEMENT OF ENDOMETRIOSIS ASSOCIATED PAIN
- I-452 EXPANDED INDICATION TO INCLUDE TREATMENT OF MULTIPLE MYELOMA PATIENTS WHO HAVE RECEIVED AT LEAST 1 PRIOR THERAPY
- I-453 USE IN COMBINATION WITH A SULFONYLUREA PLUS METFORMIN WHEN DIET, EXERCISE AND BOTH AGENTS DO NOT RESULT IN ADEQUATE GLYCEMIC CONTROL (TRIPLE THERAPY)
- I-454 MAINTENANCE OF CLINICAL REMISSION OF MILD TO MODERATE CHRON'S DISEASE INVOLVING THE ILEUM AND/OR THE ASCENDING COLON FOR UP TO 3 MONTHS
- I-455 MODIFIED HEART FAILURE INDICATION TO INCLUDE TREATMENT OF HEART FAILURE IN PATIENTS WITH LEFT VENTRICULAR SYSTOLIC DYSFUNCTION (NYHA CLASS II-IV; EJECTION FRACTION LESS THAN OR EQUAL TO 40%)
- I-456 TO REDUCE CARDIOVASCULAR DEATH AND TO REDUCE HEART FAILURE HOSPITALIZATIONS. INCLUDES ADDITIONAL INFORMATION ON THE ADDED EFFECT ON THESE OUTCOMES WHEN USED WITH AN ACE INHIBITOR
- I-457 TREATMENT OF PATIENTS UNDERGOING ABDOMINAL SUREGERY WHO ARE AT RISK FOR THROMBOEMBOLIC COMPLICATIONS
- I-458 USE OF BIVALIRUDIN FOR INJECTION WITH PROVISIONAL USE OF GLYCOPROTEIN IIB/IIA INHIBITOR (GPI) AS LISTED IN THE CLINICAL TRIALS REPLACE-2 SECTION FOR USE AS AN ANTICOAGULANT IN PATIENTS UNDERGOING PERCUTANEOUS CORONARY INTERVENTION (PCI)
- I-459 NON-DIALYSIS DEPENDENT CHRONIC KIDNEY DISEASE (NDD-CKD) PATIENTS RECEIVING OR NOT RECEIVING AN ERYTHROPOIETIN
- I-460 TREATMENT OF DIARRHEA CAUSED BY CRYPTOSPORIDIUM PARVUM IN NON-HIV INFECTED PATIENTS 12 YEARS OF AGE AND OLDER
- I-461 USE AS A SINGLE AGENT FOR ADJUVANT TREATMENT IN PATIENTS WITH DUKES' C COLON CANCER WHO HAVE UNDERGONE COMPLETE RESECTION OF THE PRIMARY TUMOR WHEN TREATMENT WITH FLUOROPYRIMIDINE THERAPY ALONE IS PREFERRED
- I-462 LONG TERM TREATMENT OF IDIOPATHIC SHORT STATURE
- I-463 TREATMENT OF PATIENTS POST MYOCARDIAL INFARCTION
- I-464 TREATMENT OF MODERATE TO SEVERE PRIMARY RESTLESS LEGS SYNDROME
- I-465 PERENNIAL ALLERGIC RHINITIS IN ADULTS AND PEDIATRIC PATIENTS 6 MONTHS OF AGE AND OLDER
- I-466 FOR RELIEF OF THE SIGNS AND SYMPTOMS OF ANKYLOSING SPONDYLITIS
- I-467 USE OF TOPIRAMATE AS INITIAL MONOTHERAPY IN PATIENTS 10 YEARS OF AGE AND OLDER WITH PARTIAL ONSET OR PRIMARY GENERALIZED TONIC CLONIC SEIZURES
- I-468 USE IN PATIENTS WITH STABLE CORONARY ARTERY DISEASE TO REDUSE THE RISK OF CARDIOVASCULAR MORTALITY OR NON-FATAL MYOCARDIAL INFECTION
- I-469 RELIEF OF THE SIGNS AND SYMPTOMS OF PAUCIARTICULAR OR POLYARTICULAR COURSE JUVENILE RHEUMATOID ARTHRITIS IN PATIENTS 2 YEARS OF AGE AND OLDER
- I-470 DIABETIC PERIPHERAL NEUROPATHIC PAIN

PATENT AND EXCLUSIVITY TERMS

ADB 15 of 35

EXCLUSIVITY INDICATION

- I-471 INDICATED TO REDUCE THE RISK OF MYOCARDIAL INFARCTION AND STROKE IN PATIENTS WITH TYPE 2 DIABETES AND WITHOUT CLINICALLY EVIDENT CORONARY HEART DISEASE BUT WITH MULTIPLE RISK FACTORS FOR CORONARY HEART DISEASE
- I-472 USE IN PATIENTS WITH ANGIOGRAPHICALLY DOCUMENTED CORONARY ARTERY DISEASE
- I-473 USE IN COMBINATION WITH GEMCITABINE FOR THE FIRST LINE TREATMENT OF PATIENTS WITH LOCALLY ADVANCED UNRESECTABLE OR METASTATIC PANCREATIC CANCER
- I-474 TREATMENT OF IRON DEFICIENCY ANEMIA IN PERITONEAL DIALYSIS DEPENDANT CHRONIC KIDNEY DISEASE IN PATIENTS RECIEVING AN ERYTHROPOIETIN
- I-475 PREVENTION OF NAUSEA AND VOMITTING ASSOCIATED WITH INITIAL AND REPEAT COURSES OF MODERATELY EMETOGENIC CANCER CHEMOTHERAPY
- I-476 TREATMENT OF DIABETIC FOOT INFECTIONS WITHOUT OSTEOMYELITIS
- I-477 TREATMENT OF COMPLICATED SKIN AND SKIN STRUCTURE INFECTIONS CAUSED BY METHICILLIN SUSCEPTIBLE STAPHYLOCOCCUS AUREUS, ESCHERICHIA COLI, KLEBSIELLA PNEUMONIAE, OR ENTEROBACTER CLOACAE
- I-478 FOR USE AS ADJUNCTIVE THERAPY IN THE TREATMENT OF PARTIAL SEIZURES IN CHILDREN WITH EPILEPSY AGED 2-4 YEARS
- I-479 TREATMENT OF COMPLICATED INTRA-ABDOMINAL INFECTIONS CAUSED BY E.COLI, B. FRAGILIS, S.ANGINOSUS, S.CONSTELLATUS, E. FAECALIS, P. MIRABILIS, C. PERFRINGENS, B. THETAIOAOMICRON OR PEPTOSTREPTOCOCCUS SPECIES
- I-480 PROPHYLAXIS OF INFLUENZA FOR PATIENTS BETWEEN 1-12 YEARS OF AGE
- I-481 INDICATED FOR THE ADJUVANT TREATMENT OF POSTMENOPAUSAL WOMEN WITH HORMONE RECEPTOR POSITIVE EARLY BREAST CANCER
- I-482 TREATMENT OF ACUTE MANIC OR MIXED EPISODES ASSOCIATED WITH BIPOLAR I DISORDER WITH OR WITHOUT PSYCHOTIC FEATURES
- I-483 PREVENTION OF POSTMENOPAUSAL OSTEOPOROSIS
- I-484 FOR THE RISK REDUCTION OF NSAID-ASSOCIATED GASTRIC ULCERS
- I-485 TREATMENT OF POSTOPERATIVE INFLAMMATION AND REDUCTION OF OCULAR PAIN IN PATIENTS WHO HAVE UNDERGONE CATARACT EXTRACTION
- I-486 ANGIOMAX IS INDICATED FOR PATIENTS WITH, OR AT RISK OF, HIT/HITTS UNDERGOING PCI
- I-487 INDICATED FOR THE RELIEF OF THE INFAMMATORY AND PRURITIC MANIFESTATIONS OF CORTICOSTEROID RESPONSIVE DERMATOSES IN PATIENTS 12 YRS OF AGE OR OLDER
- I-488 MAINTENANCE THERAPY IN BIPOLAR I DISORDER
- I-489 FOR USE IN PEDIATRIC PATIENTS WITH TYPE I DIABETES
- I-490 FOR USE IN COMBINATION WITH CISPLATIN AND FLUOROURACIL FOR THE TREATMENT OF PATIENTS WITH ADVANCED GASTRIC ADENOCARCINOMA, INCLUDING ADENOCARCINOMA OF GASTROESOPHAGEAL JUNCTION, WHO HAVE NOT RECEIVED PRIOR CHEMOTHERAPY FOR ADVANCED DISEASE
- I-491 INFLUENZA PROPHYLAXIS
- I-492 MONOTHERAPY IN THE TREATMENT OF ACUTE MANIC OR MIXED EPISODES IN BIPOLAR I DISORDER, WITH OR WITHOUT PSYCHOTIC FEATURES
- I-493 ADMINISTERED IN COMBINATION WITH FENOFIBRATE, AS ADJUNCTIVE THERAPY TO DIET FOR THE REDUCTION OF ELEVATED TOTAL-C, LDL-C, APO B, AND NON-HDL-C IN PATIENTS WITH MIXED HYPERLIPIDEMIA
- I-494 CLINICAL DATA IN SUPPORT OF AVANDAMET AS AN ADJUNCT TO DIET AND EXERCISE TO IMPROVE GLYCEMIC CONTROL IN PATIENTS WITH TYPE 2 DIABETES MELLITUS WHEN TREATMENT WITH DUAL ROSIGLITAZONE AND METFORMIN THERAPY IS APPROPRIATE
- I-495 ADJUVANT TX OF POSTMENOPAUSAL WOMEN WITH ESTROGEN-RECEPTOR POSITIVE EARLY BREAST CANCER WHO HAVE RECEIVED 2 TO 3 YRS OF TAMOXIFEN AND ARE SWITCHED TO AROMASIN FOR COMPLETION OF A TOTAL OF 5 CONSECUTIVE YRS OF ADJUVANT HORMONAL THERAPY
- I-496 LONG TERM TREATMENT OF GROWTH FAILURE ASSOCIATED WITH TURNER SYNDROME IN PATIENTS WHO HAVE OPEN EPIPHYSES
- I-497 PREVENTION OF SEASONAL MAJOR DEPRESSIVE EPISODES IN PATIENTS WITH SEASONAL AFFECTIVE DISORDER
- I-498 PREVENTION OF POSTOPERATIVE NAUSEA AND VOMITING
- I-499 USE OF GEMZAR IN COMBINATION WITH CARBOPLATIN FOR THE TREATMENT OF PATIENTS WITH ADVANCED OVARIAN CANCER THAT HAS RELAPSED AT LEAST 6 MONTHS AFTER COMPLETION OF PLATINUM-BASED THERAPY
- I-500 FOR USE IN COMBINATION WITH DEXAMETHASONE FOR THE TREATMENT OF MULTIPLE MYELOMA PATIENTS WHO HAVE RECEIVED AT LEAST ONE PRIOR THERAPY
- I-501 TREATMENT OF RECURRENT HERPES LABIALIS (COLD SORES) IN IMMUNOCOMPETANT PATIENTS WITH A SINGLE DOSE OF FAMCICLOVIR 1500 MG.
- I-502 FOR PTS WITH ST-SEGMENT ELEVATION ACUTE MYOCARDIAL INFARCTION, PLAVIX TO REDUCE RATE OF DEATH FROM ANY CAUSE AND THE RATE OF A COMBINED ENDPOINT OF DEATH, REINFARCTION OR STROKE. NOT KNOWN TO PERTAIN TO PTS WHO RECEIVE PRIMARY ANGIOPLASTY
- I-503 TREATMENT OF MAJOR DEPRESSIVE EPISODES ASSOCIATED WITH BIPOLAR DISORDER
- I-504 TREATMENT OF PATHOLOGICAL HYPERSECRETORY CONDITIONS INCLUDING ZOLLINGER-ELLISON SYNDROME

PATENT AND EXCLUSIVITY TERMS

ADB 16 of 35

EXCLUSIVITY INDICATION

- I-505 TREATMENT OF STAPHYLOCOCCUS AUREUS BLOODSTREAM INFECTIONS (BACTEREMIA), INCLUDING THOSE WITH RIGHT SIDED INFECTIVE ENDOCARDITIS, CAUSED BY METHICILLIN-SUSCEPTIBLE AND METHICILLIN-RESISTANT ISOLATES
- I-506 ADJUNCTIVE THERAPY OF MYOCLONIC SEIZURES IN ADULTS AND ADOLESCENTS AGE 12 AND OVER WITH JUVENILE MYOCLONIC EPILEPSY
- I-507 ADJUNCT TO DIET TO REDUCE TOTAL-C, LDL-C AND APO B LEVELS IN ADOLESCENT BOYS AND GIRLS WHO ARE AT LEAST ONE YEAR POST-MENARCHE, 10-16 YEARS OF AGE, WITH HETEROZYGOUS FAMILIAL HYPERCHOLESTEROLEMIA
- I-508 PREMENSTRUAL DYSPHONIC DISORDER
- I-509 TREATMENT OF IRRITABILITY ASSOCIATED WITH AUTISTIC DISORDER
- I-510 ADULT DERMAFIBROSARCOMA PROTUBERANS (DFSP)
- I-511 ADULT MYELODYSPLASTIC SYNDROME/MYELOPROLIFERATIVE DISEASES (MDS/MDP)
- I-512 ADULT PH+ ACUTE LYMPHOBLASTIC LEUKEMIA (ALL) MONOTHERAPY
- I-513 ADULT AGGRESSIVE SYSTEMIC MASTOCYTOSIS (ASM)
- I-514 ADULT HYPEREOSINOPHILIC SYNDROME/CHRONIC EOSINOPHILIC LEUKEMIA (HES/CEL)
- I-515 PROPHYLAXIS OF SURGICAL SITE INFECTION FOLLOWING ELECTIVE COLORECTAL SURGERY
- I-516 PRIMARY GENERALIZED TONIC CLONIC SEIZURES IN ADULTS AND PEDIATRIC PATIENTS 2 YEARS OF AGE AND OLDER
- I-517 TREATMENT OF MODERATE TO SEVERE PRIMARY RESTLESS LEG SYNDROME (RLS)
- I-518 TREATMENT OF SHORT STATURE OR GROWTH FAILURE IN CHILDREN WITH SHOX (SHORT STATURE HOMEBOX CONTAINING GENE) DEFICIENCY WHOSE EPIPHYSES ARE NOT CLOSED
- I-519 USE OF TAXOTERE (DOCETAXEL) INJECTION CONCENTRATE IN COMBINATION WITH CISPLATIN AND FLUOROURACIL FOR THE INDUCTION OF PATIENTS WITH INOPERABLE LOCALLY ADVANCED SQUAMOUS CELL CARCINOMA OF THE HEAD AND NECK (SCCHN)
- I-520 USE OF EXENATIDE IN PATIENTS WITH TYPE 2 DIABETES MELLITUS WHO ARE USING A THIAZOLIDINEDIONE ALONE OR IN COMBINATION WITH METFORMIN BUT HAVE NOT ACHIEVED ADEQUATE GLYCEMIC CONTROL
- I-521 TREATMENT OF PATIENTS WITH MANTLE CELL LYMPHOMA WHO HAVE RECEIVED AT LEAST 1 YEAR PRIOR THERAPY

EXCLUSIVITY MISCELLANEOUS

- M-1 INFORMATION REGARDING SUPERIORITY CLAIM OVER RANITIDINE FOR DAY AND NIGHT HEARTBURN ADDED TO CLINICAL STUDIES SECTION
- M-2 APPROVAL FOR ADDITION TO CLINICAL PHARMACOLOGY SECTION OF THE LABEL REGARDING (1) IMPROVEMENT IN BONE MINERAL DENSITY IN CHILDHOOD-ONSET ADULT GROWTH HORMONE DEFICIENT PATIENTS AND (2) INCREASES IN SERUM ALKALINE PHOSPHATASE
- M-3 ADDITION OF EFFICACY AND SAFETY INFORMATION IN WHICH FOSAMAX WAS USED CONCOMITANTLY WITH ESTROGEN ALONE OR WITH ESTROGEN PLUS PROGESTIN
- M-4 CHANGES TO PEDIATRIC USE SECTION TO PROVIDE INFORMATION REGARDING SAFETY AND EFFICACY IN PEDIATRIC PATIENTS AS YOUNG AS 2 YEARS OLD
- M-5 INFORMATION REGARDING EFFECTS IN PATIENTS WITH ASTHMA ON CONCOMITANT INHALED CORTICOSTEROIDS IN CLINICAL PHARMACOLOGY SECTION
- M-6 ADDITIONAL INFORMATION REGARDING CLINICAL STUDIES DONE WITH GLUCOPHAGE/GLYBURIDE COMBINATION ADDED TO CLINICAL PHARMACOLOGY AND DOSING AND ADMINISTRATION
- M-7 CLINICAL PHARMACOLOGY IN PEDIATRIC PATIENTS; DOSAGE AND ADMINISTRATION INFORMATION
- M-8 ADDITIONAL INFORMATION FOR THE USE OF SONATA CAPSULES FOR UP TO 5 WEEKS (35 NIGHTS) OF TREATMENT IN A CONTROLLED TRIAL SETTING
- M-9 ADDITION TO THE CLINICAL STUDIES SECTION OF THE LABELING OF TEXT AND TWO TABLES CONTAINING INFORMATION FOR THE PRESCRIBING PHYSICIAN ON BLOOD PRESSURE, HEART RATE, AND HEART RATE VARIABILITY
- M-10 INFORMATION REGARDING MAINTENANCE OF AN ANTIDEPRESSANT EFFECT UP TO 1 YEAR OF DOSING
- M-11 USE FOR LONG-TERM TREATMENT OF POSTTRAUMATIC STRESS DISORDER
- M-12 NEW LANGUAGE FOR PEDIATRIC USE
- M-13 INFORMATION FROM PEDIATRIC STUDIES ADDED TO CLINICAL PHARMACOLOGY, PRECAUTIONS, AND DOSAGE AND ADMINISTRATION
- M-14 ADDITIONAL CLINICAL TRIAL INFORMATION ADDED TO PEDIATRIC USE SUBSECTION
- M-15 LONGER TERM EFFICACY INFORMATION FOR RISPERIDONE IN THE TREATMENT OF SCHIZOPHRENIA
- M-16 CHANGE IN WORDING OF THE PEDIATRIC SECTION OF THE PACKAGE INSERT
- M-17 INFORMATION REGARDING USE OF ULTANE IN PEDIATRIC PATIENTS WITH CONGENITAL HEART DISEASE
- M-18 INFORMATION DENOTING THE EFFICACY OF REMERON IN MAINTAINING A RESPONSE IN PATIENTS WITH MAJOR DEPRESSIVE DISORDER (MDD)
- M-19 INFORMATION REGARDING USE IN PEDIATRIC PATIENTS TWO YEARS OF AGE AND OLDER
- M-20 LABELING REVISIONS RELATED TO MCCUNE ALBRIGHT SYNDROME
- M-21 COMPARISON DATA ON THE ANTIHYPERTENSIVE EFFECTS OF ATACAND AND COZAAR
- M-22 CHANGE IN TIME TO ONSET OF ACTION

PATENT AND EXCLUSIVITY TERMS

ADB 17 of 35

EXCLUSIVITY MISCELLANEOUS

- M-23 INFORMATION REGARDING ELIMINATION ADDED TO CLINICAL PHARMACOLOGY, STUDY RESULTS IN PATIENTS WITH HEPATIC AND RENAL IMPAIRMENT
- M-24 INFORMATION ON RESULTS OF A LONG TERM LONGITUDINAL GROWTH STUDY AND PEDIATRIC SAFETY INFORMATION
- M-25 ADDITIONAL SAFETY & PK INFORMATION IN CHILDREN 6 MONTHS TO LESS THAN 6 YEARS OF AGE ADDED TO PKG INSERT
- M-26 INCORPORATION OF INFORMATION CONTAINED IN THE PEG-INTRON PACKAGE INSERT INTO THE REBETOL PACKAGE INSERT AND MEDGUIDE-PEG-INTRON WAS APPROVED FOR USE IN COMBINATION WITH REBETOL FOR TREATMENT OF CHRONIC HEPATITIS C VIRUS INFECTION ON 8/7/01
- M-27 INFORMATION DESCRIBING ASPIRIN ENDOSCOPY STUDY AND THE MAXIMUM RECOMMENDED DOSE FOR PATIENTS WITH MODERATE HEPATIC INSUFFICIENCY
- M-28 INFORMATION FROM A STUDY IN PEDIATRIC PATIENTS IN ASSOCIATION WITH A NEUROLOGICAL CONDITION
- M-29 LABELING CHANGES TO PROVIDE INFORMATION IN THE MANAGEMENT OF OBESITY IN ADOLESCENTS AGED 12 TO 16 YEARS
- M-30 CHANGES TO CLINICAL PHARMACOLOGY, PRECAUTIONS, AND DOSAGE AND ADMINISTRATION SECTIONS OF LABELING CONCERNING USE OF LOTENSIN IN PEDIATRIC PATIENTS WITH HYPERTENSION
- M-31 INFORMATION FOR USE IN PEDIATRIC PATIENTS WITH CHRONIC KIDNEY DISEASE STAGE 5 (END-STAGE RENAL DISEASE)
- M-32 ADDITIONAL LANGUAGE TO CLINICAL PHARMACOLOGY AND CLINICAL STUDIES
- M-33 INFORMATION FOR USE OF ADVAIR DISKUS 100/50 IN CHILDREN 4 TO 11 YEARS OF AGE WITH ASTHMA
- M-34 EXPANDED INFORMATION TO PEDIATRIC USE SUBSECTION OF LABELING IN RESPONSE TO PEDIATRIC WRITTEN REQUEST
- M-35 ADDITIONAL INFORMATION REGARDING CLINICAL STUDIES DONE WITH ACTOS IN COMBINATION WITH METFORMIN, A SULFONYLUREA, OR INSULIN ADDED TO CLINICAL PHARMACOLOGY
- M-36 ADDITION OF INFORMATION TO CLINICAL STUDIES REGARDING PREVENTION OF CARDIOVASCULAR DISEASE
- M-37 INFORMATION ADDED TO THE LABELING THAT DETAILS INFORMATION RELATIVE TO STUDIES DONE IN PEDIATRIC POPULATIONS IN THE CLINICAL PHARMACOLOGY AND PEDIATRIC USE SUBSECTIONS
- M-38 SAFETY AND IOP-LOWERING EFFECTS OF TRUSOPT HAVE BEEN DEMONSTRATED IN PEDIATRIC PATIENTS IN A 3 MONTH, MULTI-CENTER DOUBLE MASKED ACTIVE-TREATMENT-CONTROLLED TRIAL
- M-39 FOR LABELING CHANGES BASED ON RESULTS OF THE SPD422-202 CLINICAL STUDY REPORT (CSR) SUBMITTED IN RESPONSE TO THE WRITTEN REQUEST
- M-40 ADDITIONAL INFORMATION REGARDING CLINICAL STUDIES PERFORMED IN PEDIATRIC PATIENTS WITH LEUKEMIA ADDED TO PRECAUTIONS
- M-41 REVISION TO THE PEDIATRIC USE PRECAUTIONS OF THE PRESCRIBING INFORMATION TO INCORPORATE THE RESULTS FROM THE CAPPS-169 STUDY ENTITLED "THE EFFECT OF ORTHO TRICYCLEN ON BONE MINERAL DENSITY IN PEDIATRIC SUBJECTS WITH ANOREXIA NERVOSA"
- M-42 ADDITION OF A GERIATRIC USE SUBSECTION TO THE PRECAUTIONS SECTION OF THE PACKAGE INSERT AND GERIATRIC DOSING INFORMATION
- M-43 INCLUSION OF RESULTS OF STUDY-"PLACEBO-CONTROLLED STUDY TO EVALUATE SAFETY AND PILOT EFFICACY OF ILOPROST AS ADD ON THERAPY WITH BOSENTAN IN SUBJECTS WITH PULMONARY ARTERIAL HYPERTENSION
- M-44 CLINICAL INFORMATION ADDED TO THE PEDIATRIC USE SUBSECTION OF PRECAUTIONS REGARDING THE USE OF NOVOLOG IN ADOLESCENTS WITH TYPE I DIABETES AGE 6 TO 18
- M-45 INFORMATION ADDED TO CLINICAL TRIALS SECTION OF LABELING - EFFECTS OF HUMATROPE TREATMENT IN ADULTS WITH GROWTH HORMONE DEFICIENCY
- M-46 PROVISION OF RESULTS OF STUDY AND PROPOSED REVISIONS TO PACKAGE INSERT SEE SECTION ON CARDIAC ELECTROPHYSIOLOGY
- M-47 PROVIDES FOR USE OF ANTARA WITHOUT REGARD TO MEALS
- M-48 CHANGES TO THE LABELING DESCRIBING THE RESULTS OF A STUDY OF THE USE OF NOVOLOG MIX 70/30 WITH ORAL ANTIDIABETIC AGENTS IN PATIENTS WITH TYPE 2 DIABETES
- M-49 CLINICAL DATA ADDED TO THE CLINICAL PHARMACOLOGY SECTION REGARDING EFFECT OF SINGULAIR ON GROWTH RATES IN PREPUBERTAL CHILDREN
- M-50 NEW INFO TO THE CLINICAL STUDIES, ADULT GROWTH HORMONE DEFICIENCY (GHD) SUBSECTION OF THE NUTROPIN AQ PACKAGE INSERT DESCRIBING THE EFFECTS OF SOMATROPIN ON VISCERAL ADIPOSE TISSUE IN THE ADULT GROWTH HORMONE DEFICIENT PATIENT POPULATION
- M-51 INFORMATION ADDED TO LABELING REGARDING OSTEOGENESIS IMPERFECTA STUDY
- M-52 INFORMATION ADDED TO THE CLINICAL PHARMACOLOGY/CLINICAL STUDIES SECTION REGARDING THE USE OF RISEDRONATE ADMINISTERED ONCE A WEEK IN THE PREVENTION OF OSTEOPOROSIS IN POSTMENOPAUSAL WOMEN
- M-53 FOR LABELING CHANGES TO THE QUALITY OF LIFE (QOL) STATEMENT IN THE APPROVED PACKAGE INSERT
- M-54 INFORMATION FROM PEDIATRIC STUDIES ADDED TO LABEL
- M-55 INFORMATION ON RESULTS OF A STUDY OF THE USE OF SANDOSTATIN LAR DEPOT IN PEDIATRIC PATIENTS WITH HYPOTHALAMIC OBESITY.
- M-56 INFORMATION ADDED TO CLINICAL TRIAL SECTION WITH INFORMATION ON "GEMINI" TRIAL

PATENT AND EXCLUSIVITY TERMS

ADB 18 of 35

EXCLUSIVITY MISCELLANEOUS

- M-57 CLINICAL DATA ADDED TO THE CLINICAL PHARMACOLOGY SECTION REGARDING THE PHARMACOKINETICS OF EZETIMIBE IN ASIAN SUBJECTS
- M-58 CHANGES TO THE CLINICAL STUDIES, PRIMARY HYPERCHOLESTEROLEMIA, VYTORIN SUBSECTION OF THE PACKAGE INSERT TO ADD EFFICACY DATA FOR THE EZETIMIBE/SIMVASTATIN COMBINATION PRODUCT AND FOR AN ATORVASTATIN PRODUCT ON LDL-C AND OTHER LIPID PMTRS
- M-59 RESULTS OF THE T20-310 STUDY WHICH EVALUATED THE PHARMACOKINETICS, SAFETY, AND ANTIVIRAL ACTIVITY OF FUZEON IN TREATMENT EXPERIENCED PEDIATRIC SUBJECTS AND ADOLSCENTS WAS ADDED TO THE PEDIATRIC SUBSECTION OF PRECAUTIONS
- M-60 CHANGES TO CLINICAL STUDIES, PRIMARY HYPERCHOLESTEROLEMIA, TO ADD EFFICACY DATA FOR THE EZETIMIBE/SIMVASTATIN COMBINATION PRODUCT AND FOR A ROSUVASTATIN PRODUCT ON LDL-C AND OTHER LIPID PARAMETERS IN PATIENTS WTH HYPERCHOLESTEROLEMIA

PATENT USE

- U-1 PREVENTION OF PREGNANCY
- U-2 TREATMENT OR PROPHYLAXIS OF ANGINA PECTORIS AND ARRHYTHMIA
- U-3 TREATMENT OF HYPERTENSION
- U-4 PROVIDING PREVENTION AND TREATMENT OF EMESIS AND NAUSEA IN MAMMALS
- U-5 METHOD OF PRODUCING BRONCHODILATION
- U-6 METHOD OF PRODUCING SYMPATHOMIMETIC EFFECTS
- U-7 INCREASING CARDIAC CONTRACTILITY
- U-8 ACUTE MYOCARDIAL INFARCTION
- U-9 CONTROL OF EMESIS ASSOCIATED WITH ANY CANCER CHEMOTHERAPY AGENT
- U-10 DIAGNOSTIC METHOD FOR DISTINGUISHING BETWEEN HYPOTHALMIC MALFUNCTIONS OR LESIONS IN HUMANS
- U-11 TREATMENT OR PROPHYLAXIS OF CARDIAC DISORDERS
- U-12 METHOD OF TREATING [A] HUMAN SUFFERING FROM DEPRESSION
- U-13 A METHOD FOR TREATING ANXIETY IN A HUMAN SUBJECT IN NEED OF SUCH TREATMENT
- U-14 ADJUNCTIVE THERAPY FOR THE PREVENTION AND TREATMENT OF HYPERAMMONEMIA IN THE CHRONIC MANAGEMENT OF PATIENTS WITH UREA CYCLE ENZYMOPATHIES
- U-15 METHOD OF LOWERING INTRAOCULAR PRESSURE
- U-16 USE IN LUNG SCANNING PROCEDURES
- U-17 TREATMENT OF VENTRICULAR AND SUPRAVENTRICULAR ARRHYTHMIAS
- U-18 METHOD FOR INHIBITING GASTRIC SECRETION IN MAMMALS
- U-19 TREATMENT OF INFLAMMATION
- U-20 A PROCESS FOR TREATING A PATIENT SUFFERING FROM PARKINSON'S SYNDROME AND IN NEED OF TREATMENT
- U-21 TREATMENT OF HUMANS SUFFERING UNDESIRE UROTOXIC SIDE EFFECTS CAUSED BY CYTOSTATICALLY ACTIVE ALKYLATING AGENTS
- U-22 METHOD OF COMBATting PATHOLOGICALLY REDUCED CEREBRAL FUNCTIONS AND PERFORMANCE WEAKNESSES, CEREBRAL INSUFFICIENCY AND DISORDERS IN CEREBRAL CIRCULATION AND METABOLISM IN WARM-BLOODED ANIMALS
- U-23 METHOD FOR TREATING PROSTATIC CARCINOMA COMPRISING ADMINISTERING FLUTAMIDE
- U-24 METHOD FOR TREATING PROSTATE ADENOCARCINOMA COMPRISING ADMINISTERING AN ANTIANDROGEN INCLUDING FLUTAMIDE AND AN LHRH AGONIST
- U-25 REDUCING CHOLESTEROL IN CHOLELITHIASIS PATIENTS
- U-26 REDUCING CHOLESTEROL GALLSTONES AND/OR FRAGMENTS THEREOF
- U-27 DISSOLVING CHOLESTEROL GALLSTONES AND/OR FRAGMENTS THEREOF
- U-28 CEREBRAL, CORONARY, PERIPHERAL, VISCERAL AND RENAL ARTERIOGRAPHY, AORTOGRAPHY AND LEFT VENTRICULOGRAPHY
- U-29 CT IMAGING OF THE HEAD AND BODY, AND INTRAVENOUS EXCRETORY UROGRAPHY
- U-30 CEREBRAL ANGIOGRAPHY, AND VENOGRAPHY
- U-31 INTRA-ARTERIAL DIGITAL SUBTRACTION ANGIOGRAPHY
- U-32 PALLIATIVE TREATMENT OF PATIENTS WITH OVARIAN CARCINOMA RECURRENT AFTER PRIOR CHEMOTHERAPY, INCLUDING PATIENTS WHO HAVE BEEN PREVIOUSLY TREATED WITH CISPLATIN
- U-33 TREATING VIRAL INFECTIONS IN A MAMMAL
- U-34 TREATING VIRAL INFECTIONS IN A WARM-BLOODED ANIMAL
- U-35 TREATING CYTOMEGALOVIRUS IN A HUMAN WITH AN INJECTABLE COMPOSITION
- U-36 METHODS OF TREATING BACTERIAL ILLNESSES
- U-37 METHOD OF TREATING GASTROINTESTINAL DISEASE
- U-38 TREATMENT OF PAROXYSMAL SUPRAVENTRICULAR TACHYCARDIA
- U-39 ANGINA PECTORIS
- U-40 METHOD OF TREATMENT OF BURNS
- U-41 METHOD OF TREATING CARDIAC ARRHYTHMIAS

PATENT AND EXCLUSIVITY TERMS

ADB 19 of 35

PATENT USE

- U-42 ADJUVANT TREATMENT IN COMBINATION WITH FLUOROURACIL AFTER SURGICAL RESECTION IN PATIENTS WITH DUKES' STAGE C COLON CANCER
- U-43 MANAGEMENT OF CHRONIC PAIN IN PATIENTS REQUIRING OPIOID ANALGESIA
- U-44 RELIEF OF NAUSEA AND VOMITING
- U-45 TREATMENT OF INFLAMMATION AND ANALGESIA
- U-46 TREATMENT OF PANIC DISORDER
- U-47 STIMULATION OF THE RELEASE OF GROWTH HORMONE
- U-48 ANALGESIA
- U-49 SYMPTOMATIC CANCER-RELATED HYPERCALCEMIA
- U-50 USE IN TREATING INFLAMMATORY DERMATOSES
- U-51 BLOOD POOL IMAGING, INCLUDING CARDIAC FIRST PASS AND GATED EQUILIBRIUM IMAGING AND FOR DETECTION OF SITES OF GASTROINTESTINAL BLEEDING
- U-52 TREATMENT OF ADULT AND PEDIATRIC PATIENTS(OVER SIX MONTHS OF AGE) WITH ADVANCED HIV INFECTION
- U-53 HYPERCALCEMIA OF MALIGNANCY
- U-54 REVERSAL AGENT OR ANTAGONIST OF NONDEPOLARIZING NEUROMUSCULAR BLOCKING AGENTS
- U-55 TREATMENT OF PAIN
- U-56 AID TO SMOKING CESSATION
- U-57 OPHTHALMIC USE OF NORFLOXACIN
- U-58 METHOD OF TREATING INFLAMMATORY INTESTINAL DISEASES
- U-59 METHOD OF TREATING HYPERCHOLESTEROLEMIA
- U-60 NASAL ADMINISTRATION OF BUTORPHANOL
- U-61 CEREBRAL AND PERIPHERAL ARTERIOGRAPHY AND CT IMAGING OF THE HEAD
- U-62 CORONARY ARTERIOGRAPHY, LEFT VENTRICULOGRAPHY, CT IMAGING OF THE BODY, INTRAVENOUS EXCRETORY UROGRAPHY, INTRAVENOUS DIGITAL SUBTRACTION ANGIOGRAPHY AND VENOGRAPHY
- U-63 ISOPRENALINE ANTAGONISM ON THE HEART RATE OR BLOOD PRESSURE
- U-64 TREATMENT OF VIRAL INFECTIONS
- U-65 METHOD OF TREATMENT OF A PATIENT INFECTED WITH HIV
- U-66 TRIPHASIC REGIMEN
- U-67 METHOD OF INDUCING ANESTHESIA IN A WARM BLOODED ANIMAL
- U-68 TREATMENT OF ACTINIC KERATOSIS
- U-69 TREATMENT OF PNEUMOCYSTIS CARINII INFECTIONS
- U-70 TREATMENT OF TRANSIENT INSOMNIA
- U-71 METHOD OF TREATMENT OF HEART FAILURE
- U-72 TREATMENT OF MIGRAINE
- U-73 METHOD OF TREATING DISEASES OR INFECTIONS CAUSED BY MYCETES
- U-74 METHOD OF PROVIDING HYPNOTIC EFFECT
- U-75 RELIEF OF OCULAR ITCHING DUE TO SEASONAL ALLERGIC CONJUNCTIVITIS
- U-76 USE TO IMAGE A SUBJECT WITH A MAGNETIC RESONANCE IMAGING SYSTEM
- U-77 TREATMENT OF SYMPTOMS OF SEASONAL ALLERGIC RHINITIS
- U-78 ULCERATIVE COLITIS
- U-79 SYMPTOMATIC TREATMENT OF PATIENTS WITH NOCTURNAL HEARTBURNDUE TO GERD
- U-80 METHOD OF TREATING OCULAR BACTERIAL INFECTIONS
- U-81 RELIEF OF SYMPTOMS ASSOCIATED WITH SEASONAL ALLERGIC RHINITIS
- U-82 TREATMENT FOR DEMENTIA IN PATIENTS WITH ALZHEIMER'S DISEASE
- U-83 TREATMENT OF SEIZURES
- U-84 A METHOD OF BLOCKING THE UPTAKE OF MONOAMINES BY BRAIN NEURONS IN ANIMALS
- U-85 NASAL TREATMENT OF SEASONAL AND PERENNIAL ALLERGIC RHINITIS SYMPTOMS
- U-86 METHOD OF TREATING CERTAIN FORMS OF EPILEPSY
- U-87 METHOD FOR NONINVASIVE ADMINISTRATION OF SEDATIVES, ANALGESICS, AND ANESTHETICS
- U-88 TREATMENT OF MODERATE PLAQUE PSORIASIS
- U-89 TREATMENT OR PROPHYLAXIS OF EMESIS
- U-90 TREATMENT OF PYSCHOTIC DISORDERS
- U-91 ALTERNATIVE THERAPY TO TRIMETHOPRIM-SULFAMETHOXAZOLE FOR TREATMENT OF MODERATE-TO-SEVERE PNEUMOCYSTIS CARINII PNEUMONIA IN IMMUNOCOMPROMISED AND AIDS PATIENTS
- U-92 TREATMENT OF DIABETIC NEPHROPATHY IN PATIENTS WITH TYPE I INSULIN DEPENDENT DIABETES MELLITUS AND RETINOPATY
- U-93 USE AS AN ANTIHISTAMINE/DECONGESTANT
- U-94 TREATMENT-ADULTS W/ ADVANCED HIV,INTOLERANT OF APPROVED THERAPIES,INTOLERANT OF APPROVED THERAPIES W/PROVEN BENEFIT OR HAVE EXPERIENCED CLINICAL/IMMUNOLOGICAL DETERIORATION WHILE RECEIVING..OR FOR WHOM SUCH THERAPIES-CONTRAINDICATED
- U-95 SHORT TERM MANAGEMENT OF MODERATE PRURITIS IN ADULTS WITH ATOPIC DERMATITIS AND LICHEN SIMPLEX CHRONICUS

PATENT AND EXCLUSIVITY TERMS

ADB 20 of 35

PATENT USE

U-96 METHOD OF TREATING VARICELLA ZOSTER (SHINGLES) INFECTIONS
U-97 A METHOD OF TREATING A PATIENT IN NEED OF MEMORY ENHANCEMENT
U-98 A METHOD OF INDUCING REGRESSION OF LEUKEMIA CELL GROWTH IN A MAMMAL
U-99 METHOD OF PROVIDING POTASSIUM TO A SUBJECT IN NEED OF POTASSIUM
U-100 METHOD OF TREATING OCULAR INFLAMMATION
U-101 ADJUNCT TO CONVENTIONAL CT OR MRI IMAGING IN THE LOCALIZATION OF STROKE IN PATIENTS IN WHOM STROKE HAS ALREADY BEEN DIAGNOSED
U-102 METHOD OF HORMONALLY TREATING MENOPAUSAL OR POST-MENOPAUSAL DISORDERS IN WOMEN
U-103 TREATMENT OF OCULAR HYPERTENSION
U-104 TREATMENT OF AQUEOUS HUMOR FORMATION AND INTRAOCULAR PRESSURE
U-105 EMESIS
U-106 TREATMENT OF EPILEPSY
U-107 TREATMENT OF HYPERTENSION AND ANGINA PECTORIS
U-108 SHORT-TERM TREATMENT OF ACTIVE DUODENAL ULCER, GASTROESOPHAGEAL REFLUX DISEASE (GERD), SEVERE EROSIIVE ESOPHAGITIS, POORLY RESPONSIVE SYMPTOMATIC GERD AND PATHOLOGICAL HYPERSECRETORY CONDITIONS AND MAINTENANCE HEALING OF EROSIIVE ESOPHAGITIS
U-109 ADJUNCT DIET IN THE TX OF ELEVATED TOTAL CHOLESTEROL AND LDL-C LEVELS IN PTS W/PRIMARY HYPERCHOLESTEROLEMIA WHOSE RESPONSE TO DIETARY RESTRICTION OF SAT FAT AND CHOLESTEROL AND OTHER NONPHARMACOLOGICAL MEASURES HAS NOT BEEN ADEQUATE
U-110 USE AS A RETRIEVABLE PERSSARY
U-111 DIABETES
U-112 CONTRACEPTION
U-113 METHOD OF CONDUCTING RADIOLOGICAL EXAMINATION OF A PATIENT BY ADMINISTERING TO SAID PATIENT A RADIOPAQUE AMOUNT OF IOPROMIDE
U-114 USE FOR INHIBITING BONE RESORPTION
U-115 USE OF VASODILATORS TO EFFECT AND ENHANCE AN ERECTION (AND THUS TREAT ERECTILE DYSFUNCTION), BY INJECTION INTO THE PENIS
U-116 METHOD OF MYOCARDIAL IMAGING
U-117 TREATMENT OF OCULAR ALLERGIC RESPONSE IN HUMAN EYES
U-118 METHOD OF LOWERING BLOOD SUGAR LEVEL
U-119 TREATMENT OF NASAL HYPERSECRETION
U-120 CONTROLLING OR PREVENTING POST-OPERATIVE INTRAOCULAR PRESSURE RISES ASSOCIATED WITH OPHTHALMIC LASER SURGICAL PROCEDURES
U-121 METHOD OF TREATING CONDITIONS MEDIATED THROUGH HISTAMINE H2-RECEPTORS
U-122 A THERAPEUTIC METHOD FOR CONTROLLING THROMBOSIS
U-123 METHOD FOR CONTROLLING THROMBOSIS AND DECREASING BLOOD HYPERCOAGULATION AND HEMORRHAGING RISKS
U-124 TREATMENT OF ACNE
U-125 TREATMENT NEUROGENERATIVE DISEASES
U-126 TREATMENT OF GASTRITIS
U-127 METHOD OF PRODUCING NEUROMUSCULAR BLOCKADE
U-128 METHOD FOR TREATMENT OF TUMORS
U-129 METHOD TO DESTROY OR IMPAIR TARGET CELLS
U-130 MANAGEMENT OF PATIENTS WITH MASTOCYTOSIS
U-131 PHOTODAMAGED SKIN
U-132 INHIBITING HIV PROTEASE
U-133 MANAGEMENT OF OBESITY INCLUDING WEIGHT LOSS AND MAINTENANCE IN PATIENTS ON A REDUCED-CALORIE DIET
U-134 TREATMENT OF ACNE VULGARIS
U-135 ANTITUMOR AGENT
U-136 PROCESS FOR WASTE NITROGEN REMOVAL
U-137 METHOD OF TREATING BACTERIAL VAGINOSIS
U-138 TREATMENT OF ALLERGIC RHINITIS
U-139 TREATMENT OF ALLERGIC REACTIONS
U-140 USE OF NORVIR TO INHIBIT HIV PROTEASE OR TO INHIBIT AN HIV INFECTION
U-141 TREATMENT OF ULCERATIVE COLITIS
U-142 METHOD OF TREATING ALLERGIC REACTIONS IN A MAMMAL BY USING THIS ACTIVE METABOLITE
U-143 BIODEGRADABLE SUPERPARAMAGNETIC METAL OXIDES AS CONTRAST AGENTS FOR MR IMAGING
U-144 BIOLOGICALLY DEGRADABLE SUPERPARAMAGNETIC MATERIALS FOR USE IN CLINICAL APPLICATIONS
U-145 BIOLOGICALLY DEGRADABLE SUPERPARAMAGNETIC PARTICLES FOR USE AS NUCLEAR MAGNETIC RESONANCE IMAGING AGENTS
U-146 METHOD OF TREATING SUSCEPTIBLE NEOPLASMS IN MAMMALS
U-147 DETECTION OF GASTROINTESTINAL DISORDERS AND THE SUBSEQUENT BREATH COLLECTION AND MEASUREMENT OF 13C02

PATENT AND EXCLUSIVITY TERMS

ADB 21 of 35

PATENT USE

- U-148 DEVICE FOR COLLECTING A BREATH SAMPLE
- U-149 METHOD OF TREATING AN ANIMAL, INCLUDING A HUMAN SUFFERING FROM OR SUSCEPTIBLE TO PSYCHOSIS, ACUTE MANIA OR MILD ANXIETY STATES
- U-150 METHOD OF USE FOR CONTROLLING HYPERGLYCEMIA BY ADMINISTRATION OF THIS SUSTAINED RELEASE DOSAGE FORM OF GLIPIZIDE
- U-151 RELIEF OF SYMPTOMS OF THE COMMON COLD
- U-152 METHOD OF TREATING ANXIETY RELATED DISORDERS INCLUDING OBSESSIVE COMPULSIVE DISORDER
- U-153 TREATMENT OF INITIAL EPISODE GENITAL HERPES
- U-154 METHOD OF TREATING ANIMALS SUFFERING FROM AN APPETITE DISORDER
- U-155 TREATMENT OF ERECTILE DYSFUNCTION
- U-156 METHOD OF PROVIDING ANESTHESIA
- U-157 TREATMENT OF A HUMAN SUFFERING FROM VITAMIN B12 DEFICIENCY
- U-158 ANGINA
- U-159 TREATMENT OF INTERSTITIAL CYSTITIS
- U-160 TREATMENT OF BACTERIAL INFECTIOUS DISEASE
- U-161 METHOD OF INHIBITING CHOLESTEROL BIOSYNTHESIS IN A PATIENT
- U-162 METHOD OF USE TO INHIBIT CHOLESTEROL SYNTHESIS IN A HUMAN SUFFERING FROM HYPERCHOLESTEROLEMIA
- U-163 METHOD OF USING TROGLITAZONE TO TREAT IMPAIRED GLUCOSE TOLERANCE TO PREVENT OR DELAY THE ONSET OF NONINSULIN-DEPENDENT DIABETES MELLITUS
- U-164 METHOD OF USING TROGLITAZONE TO PREVENT OR DELAY THE ONSET OF NONINSULIN-DEPENDENT DIABETES MELLITUS IN A DEFINED POPULATION OF PATIENTS
- U-165 TREATMENT OF SYMPTOMATIC BENIGN PROSTATIC HYPERPLASIA
- U-166 TREATMENT OF H.PYLORI-ASSOCIATED DUODENAL ULCER
- U-167 METHOD FOR TREATING HIV-1 INFECTION
- U-168 METHOD OF INHIBITING LIPOXYGENASE ACTIVITY IN A MAMMAL WHICH IS THE MODE OF ACTION IN THE TREATMENT OF ASTHMA
- U-169 METHODS OF USING THE COMPOUND/DRUG PRODUCT AS A CONTRAST AGENT IN MAGNETIC RESONANCE IMAGING
- U-170 METHOD OF OBTAINING AN MR IMAGE USING THE COMPOSITION/DRUG PRODUCT AS A CONTRAST AGENT
- U-171 METHODS OF USING THE COMPOUND/DRUG PRODUCT AS AN ORAL CONTRAST AGENT IN MAGNETIC RESONANCE IMAGING OF THE GASTROINTESTINAL TRACT
- U-172 TREATMENT OF GENITAL WARTS
- U-173 ADMINISTRATION TO A HOST SUFFERING FROM GESTATIONAL DIABETES
- U-174 USE AS AN ANTIHISTAMINE AGENT
- U-175 METHOD OF TREATING MALIGNANT TUMORS
- U-176 METHOD OF TREATING A PATIENT SUFFERING FROM LISTED CONDITIONS, INCLUDING SPECIFIC PSYCHOSES
- U-177 FUNGICIDE
- U-178 FACILITATED ADHERENCE OF AGENTS TO SKIN
- U-179 ENHANCED CUTANEOUS PENETRATION OF A DERMALLY-APPLIED PHARMACOLOGICALLY ACTIVE AGENT
- U-180 TREATMENT OF ADULT AND PEDIATRIC PATIENTS (OVER 6 MONTHS OF AGE) WITH ADVANCED HIV INFECTION
- U-181 PRODUCING ALPHA ADRENERGIC ANTAGONISTIC ACTION IN A HOST
- U-182 USE OF SALMETEROL IN PATIENTS WITH REVERSIBLE AIRWAY OBSTRUCTION
- U-183 TREATMENT OF CONDITIONS CAUSED BY DISTURBANCE OF NEURONAL 5HT FUNCTION
- U-184 TREATING ALLERGIC EYE DISEASES IN HUMANS
- U-185 METHOD OF TREATING HYPERTENSION
- U-186 METHOD FOR TREATING GI DISORDERS CAUSED BY H.PYLORI WHICH COMPRISES ADMINISTRATION OF RANITIDINE BISMUTH CITRATE AND CLARITHROMYCIN FOR A GREATER THAN ADDITIVE EFFECT
- U-187 THERAPEUTIC TREATMENT OF CALCIFIC TUMORS
- U-188 TREATMENT OF H.PYLORI ASSOCIATED DUODENAL ULCER
- U-189 ENHANCEMENT OF THE BIOAVAILABILITY OF THE DRUG SUBSTANCE
- U-190 USE OF RITONAVIR IN COMBINATION WITH ANY REVERSE TRANSCRIPTASE INHIBITOR
- U-191 METHOD OF TREATMENT FOR CONTROLLING AND LOWERING INTRAOCULAR PRESSURE IN A HUMAN
- U-192 USE IN TREATING ALLERGIC REACTIONS
- U-193 PSORIASIS
- U-194 TREATING ANGINA PECTORIS AND HIGH BLOOD PRESSURE
- U-195 METHOD FOR THE DIAGNOSIS OF GASTROINTESTINAL DISORDERS BY UREA ISOTOAC OR NITROGEN LABELED CARBON
- U-196 TREATMENT OF METASTATIC BREAST CANCER IN POSTMENOPAUSAL WOMEN WITH ESTROGEN RECEPTOR POSITIVE TUMORS
- U-197 USE IN COMBINATION WITH CERTAIN LHRH ANALOGUES FOR THE TREATMENT OF ADVANCED PROSTATE CANCER

PATENT AND EXCLUSIVITY TERMS

ADB 22 of 35

PATENT USE

- U-198 TREATMENT METASTATIC CARCINOMA OF OVARY AFTER 1ST LINE FAILURE OR SUBSEQUENT CHEMOTHERAPY, TREATMENT OF BREAST CANCER AFTER FAILURE OF COMBINATION CHEMOTHERAPY FOR METASTATIC DISEASE AND 2ND LINE TREATMENT OF AIDS RELATED KAPOSI'S SARCOMA
- U-199 METHOD OF TREATING INFECTIOUS UPPER GI TRACT DISORDERS CAUSED BY CAMPYLOBACTER PYLORIDIS INFECTION COMPRISING ADMINISTRATION OF A BISMUTH AGENT AND AN ANTIMICROBIAL AGENT
- U-200 METHOD OF TREATING GI DISORDERS COMPRISING ADMINISTRATION OF A BISMUTH-CONTAINING AGENT AND H2 RECEPTOR BLOCKING ANTI-SECRETORY AGENT
- U-201 METHOD OF TREATING GI DISORDERS COMPRISING ADMINISTRATION OF CAMPYLOBACTER-INHIBITING ANTIMICROBIAL AGENT AND H2 RECEPTOR BLOCKING ANTI-SECRETORY AGENT
- U-202 METHOD OF TREATING PEPTIC ULCER DISEASE CAUSED BY CAMPYLOBACTER PYLORIDIS COMPRISING ORAL ADMINISTRATION OF 50 TO 5,000MG BISMUTH DAILY FOR 3-56 DAYS
- U-203 TREATMENT OF ADVANCED BREAST CANCER IN POSTMENOPAUSAL WOMEN WITH DISEASE PROGRESSION FOLLOWING ANTIESTROGEN THERAPY
- U-204 USE OF TAXOL IN COMBINATION WITH G-CSF FOR TREATMENT OF PATIENTS WITH AIDS-RELATED KAPOSI'S SARCOMA
- U-205 METHOD FOR TREATING HEARTBURN
- U-206 METHOD OF USING FSH ALONE, WITHOUT THE PRESENCE OF EXOGENEOUS LH, IN IN VITRO FERTILIZATION
- U-207 USE AS NASAL SPRAY
- U-208 VAGINAL ADMINISTRATION USING SPECIFIED FORMULATION
- U-209 VAGINAL ADMINISTRATION OF PROGESTERONE USING SPECIFIED FORMULATION
- U-210 METHOD OF TREATING CONGESTIVE HEART FAILURE
- U-211 USE IN PATIENTS WITH REVERSIBLE AIRWAY OBSTRUCTION
- U-212 METHOD OF TREATMENT OF PARKINSON'S DISEASE
- U-213 METHOD OF INHIBITING CHOLESTEROL BIOSYNTHESIS AND TREATING HYPERCHOLESTEROLEMIA AND METHOD FOR TREATING HYPERLIPIDEMIA
- U-214 USE AS A BLOOD GLUCOSE-LOWERING AGENT
- U-215 TREATMENT OF EPILEPSY TWICE DAILY. TREATING A PATIENT BY ADMINISTERING CARBAMAZEPINE IN A DOSAGE FORM CAPABLE OF MAINTAINING BLOOD CONCENTRATION FROM 4-12MCG/ML OVER 12 HOURS
- U-216 TREATMENT OF ADENOCARCINOMA, INCLUDING STAGE B2-C BY ADMINISTERING AN AGONIST OF LH-RH AND FLUTAMIDE
- U-217 METHOD OF PRODUCING ANESTHESIA
- U-218 METHOD FOR LIMITING THE POTENTIAL FOR MICROBIAL GROWTH IN THE DRUG PRODUCT
- U-219 TREATMENT OF PARKINSON'S DISEASE
- U-220 METHOD OF DIAGNOSIS
- U-221 SELECTIVE VASODILATION BY CONTINUOUS ADENOSINE INFUSION
- U-222 METHOD OF TREATING PAGET'S DISEASE USING ACTONEL
- U-223 TREATMENT OF BACTERIAL CONJUNCTIVITIS CAUSED BY SUSCEPTIBLE STRAINS OF MICROORGANISMS
- U-224 CONTROLLING INTRAOCULAR PRESSURE
- U-225 MEHTOD FOR DELIVERY
- U-226 METHOD OF ENHANCING THE DISSOLUTION PROFILE OF A PHARMACEUTICAL FROM A SOLID DOSAGE FORM CONTAINING THE PHARMACEUTICAL AND SIMETHICONE
- U-227 NASAL ADMINISTRATION
- U-228 ASTHMA
- U-229 CARDIAC INSUFFICIENCY (CONGESTIVE HEART FAILURE)
- U-230 PREVENTION OF ACUTE CARDIAC ISCHEMIC EVENTS
- U-231 USE IN PARKINSON'S DISEASE
- U-232 METHOD OF TREATING MIGRAINE
- U-233 DECREASING MORTALITY CAUSED BY CONGESTIVE HEART FAILURE
- U-234 METHOD OF USING RIBAVIRIN TO TREAT VIRAL INFECTIONS IN MAMMALS
- U-235 METHOD OF MODULATING TH1 AND TH2 RESPONSE IN ACTIVATED T CELLS OF A HUMAN COMPRISING ADMINISTERING RIBAVIRIN TO THE T CELLS IN A DOSAGE WHICH PROMOTES THE TH1 RESPONSE AND SUPPRESSES THE TH2 RESPONSE
- U-236 TREATING MALE PATTERN BALDNESS WITH 0.05 TO 3.0MG/DAY
- U-237 METHOD OF PERFORMING NMR IMAGING WITH A PATIENT COMPRISING ADMINISTERING TO THE PATIENT AN EFFECTIVE AMOUNT OF CONTRAST AGENT DISCLOSED IN THE CLAIMS
- U-238 IMAGING A BODY TISSUE AND SUBJECTING TO NMR TOMOGRAPHY, ADMINISTERING AN AMOUNT OF PHARMACEUTICAL AGENT FOR AFFECTING THE RELAXATION TIMES OF ATOMS IN BODY TISSUES UNDERGOING NMR DIAGNOSIS, WHEREBY THE IMAGE CONTRAST IN ENHANCED....
- U-239 TREATING OR CONTROLLING OCULAR INFLAMATION WHICH COMPRISES TOPICALLY ADMINISTERING TO AFFECTED EYE A COMPOSITION COMPRISING AN NSAID, A POLYMERIC QUATERNARY AMMONIUM COMPOUND AND BORIC ACID
- U-240 TREATMENT OF ACUTE MIGRAINE ATTACKS

PATENT AND EXCLUSIVITY TERMS

ADB 23 of 35

PATENT USE

- U-241 FOR SHORT-TERM TREATMENT ACTIVE DUODENAL ULCER, MAINTENANCE THERAPY FOR DUODENAL ULCER PATIENTS AT REDUCED DOSAGE AFTER HEALING OF ACTIVE ULCER, SHORT-TERM TREATMENT ACTIVE BENIGN GASTRIC ULCER & GERD, PATHOLOGICAL HYPERSECRETORY CONDITIONS
- U-242 USE OF FOLLITROPIN ALPHA ALONE IN IN-VITRO FERTILIZATION
- U-243 TOPICAL ADMINISTRATION
- U-244 PLATELET AGGREGATION INHIBITORS
- U-245 TREATMENT OF SEBORRHEA DERMATITIS IN HUMANS
- U-246 PHOSPHATE BINDING
- U-247 TREATMENT OF RHEUMATOID ARTHRITIS
- U-248 TREATMENT OF HIV
- U-249 METHOD OF TREATING ALLERGIC OR NON-ALLERGIC RHINITIS IN PATIENTS BY ADMINISTERING AEROSOLIZED PARTICLES OF MOMETASONE FUROATE
- U-250 TREATMENT OF HEPATITIS B INFECTION
- U-251 USE OF TROGLITAZONE IN COMBINATION WITH SULFONYLUREAS IN THE TREATMENT OF TYPE II DIABETES
- U-252 METHOD OF TREATING A HUMAN SUBJECT HAVING GAUCHER'S DISEASE
- U-253 ORAL TRANSMUCOSAL USE
- U-254 USE OF AGGRASTAT IN COMBINATION WITH HEPARIN
- U-255 IMPROVED WAKEFULNESS IN PATIENTS WITH EXCESSIVE DAYTIME SLEEPINESS ASSOCIATED WITH NARCOLEPSY
- U-256 TREATMENT OF HIV INFECTION IN COMBINATION WITH ONE OR MORE ADDITIONAL HIV ANTIVIRAL AGENTS
- U-257 TREATMENT OF HIV INFECTION
- U-258 TREATMENT OF NEURODEGENERATIVE DISEASES
- U-259 TREATMENT OF ANDROGENIC ALOPECIA BY ORAL ADMINISTRATION DRUG SUBSTANCE
- U-260 REDUCTION OF INTRAOCULAR PRESSURE IN PATIENTS WITH OPEN ANGLE GLAUCOMA AND OCULAR HYPERTENSION WHO ARE INTOLERANT OF OTHER IOP LOWERING MEDICATIONS OR INSUFFICIENTLY RESPONSIVE TO ANOTHER IOP LOWERING MEDICATION
- U-261 TREATING BENIGN PROSTATIC HYPERPLASIA WITH A GENUS OF COMPOUNDS, INCLUDING FINASTERIDE
- U-262 TREATING BENIGN PROSTATIC HYPERTROPHY WITH FINASTERIDE
- U-263 METHOD OF TREATING A MALIGNANT CONDITION THROUGH INTRAVASCULAR ADMINISTRATION OF BUSULFAN. METHOD FOR TREATING LEUKEMIA OR LYMPHOMA IN A PATIENT UNDERGOING A BONE MARROW TRANSPLANT THROUGH INTRAVENOUS ADMINISTRATION OF BUSULFAN
- U-264 METHOD OF TREATING A MALIGNANT DISEASE THROUGH PARENTERAL ADMINISTRATION OF BUSULFAN. METHOD FOR TREATING A PATIENT UNDERGOING A BONE MARROW TRANSPLANT THROUGH INTRAVASCULAR ADMINISTRATION OF BUSULFAN
- U-265 USE AS LAXATIVE
- U-266 RELIEF OF THE SIGNS AND SYMPTOMS OF OSTEOARTHRITIS; RELIEF OF THE SIGNS AND SYMPTOMS OF RHEUMATOID ARTHRITIS IN ADULTS; MANAGEMENT OF ACUTE PAIN IN ADULTS; TREATMENT OF PRIMARY DYSMENORRHEA; ACUTE TREATMENT OF MIGRAINE ATTACKS IN ADULTS
- U-267 PREVENTING HEARTBURN EPISODES FOLLOWING INGESTION OF HEARTBURN-INDUCING FOOD/BEVERAGE, COMPRISING ADMIN TO PT, 30 MIN PRIOR TO CONSUMPTION BY THE PT THE FOOD/BEVERAGE, A COMPOSITION COMPRISING 10MG FAMOTIDINE
- U-268 ACROMEGALY
- U-269 EXCESS GH-SECRETION OR GASTRO-INTESTINAL DISORDERS
- U-270 METHOD OF IMPROVING THE TIME FOR ADMINISTRATION OR THE TIME BETWEEN CHANGES OF GIVING SETS FOR THE DRUG PRODUCT
- U-271 METHOD OF TREATING TUMORS
- U-272 METHOD OF TREATING CARCINOMA
- U-273 CUTANEOUS T-CELL LYMPHOMA
- U-274 ZANAMIVIR FOR INHALATION
- U-275 METHOD OF USE OF THE DRUG SUBSTANCE
- U-276 METHOD OF USE OF LEVOBUPIVACAINE
- U-277 NEUROLOGICAL AND OTHER DISORDERS (TREATMENT OF EPILEPSY, BID ORAL DOSING)
- U-278 METHOD OF USE OF THE INDICATION OF THE DRUG PRODUCT
- U-279 METHOD OF USE OF THE APPROVED PRODUCT
- U-280 TREATING PRECIPITATED ACUTE URINARY RETENTION WITH FINASTERIDE
- U-281 ANTIMYCOTIC USES, SPECIFICALLY TREATMENT OF ONYCHOMYCOSIS
- U-282 METHOD OF TREATING BACTERIAL INFECTIONS
- U-283 METHOD FOR TREATING MENOPAUSAL SYMPTOMS IN A POSTMENOPAUSAL FEMALE
- U-284 MENOPAUSAL AND POSTMENOPAUSAL DISORDERS (INCLUDING VASOMOTOR SYMPTOMS ASSOCIATED WITH MENOPAUSE, AND VULVAR AND VAGINAL ATROPHY) AND OSTEOPOROSIS
- U-285 DEPRESSION AND SOCIAL ANXIETY DISORDER/SOCIAL PHOBIA
- U-286 DEPRESSION
- U-287 TREATMENT OR PREVENTION OF OSTEOPOROSIS

PATENT AND EXCLUSIVITY TERMS

ADB 24 of 35

PATENT USE

- U-288 THERAPY OF INFLUENZA
- U-289 TREATMENT OF NON-HYPERKERATOTIC ACTINIC KERATOSES OF FACE AND SCALP
- U-290 INHIBITING TRANSPLANT REJECTION USING RAPAMYCIN (SIROLIMUS)
- U-291 INHIBITING TRANSPLANT REJECTION USING RAPAMYCIN (SIROLIMUS) IN COMBINATION WITH CYCLOSPORIN
- U-292 INHIBITING TRANSPLANT REJECTION USING RAPAMYCIN (SIROLIMUS) IN COMBINATION WITH AZATHIOPRINE
- U-293 INHIBITING TRANSPLANT REJECTION USING RAPAMYCIN (SIROLIMUS) IN COMBINATION WITH A CORTICOSTEROID
- U-294 TREATMENT OF HYPERPIGMENTARY DISORDERS
- U-295 TREATMENT OF SEASONAL AND PERENNIAL ALLERGIC RHINITIS SYMPTOMS
- U-296 TREATING MIGRAINE PAIN AND ONE OR MORE OF A CLUSTER OF SYMPTOMS CHARACTERISTIC OF A MIGRAINE ATTACK SYMPTOMS BEING SELECTED FROM PHOTOPHOBIA, PHONOPHOBIA NAUSEA AND FUNCTIONAL DISABILITY
- U-297 PREVENTION OR TREATMENT OF REVERSIBLE VASOCONSTRICTION BY THE INHALATION OF NITRIC OXIDE WITH AN OXYGEN CONTAINING GAS
- U-298 METHOD OF COMBATING BACTERIA IN A PATIENT
- U-299 TREATMENT OF ADENOMATOUS POLYPS
- U-300 INDICATED FOR THE REDUCTION OF ELEVATED TOTAL AND LDL CHOLESTEROL LEVELS IN PATIENTS WITH PRIMARY HYPERCHOLESTEROLEMIA
- U-301 USE OF TROGLITAZONE IN COMBINATION WITH SULFONYLUREAS AND BIGUANIDES IN THE TREATMENT OF TYPE II DIABETES
- U-302 TO REDUCE THE RISK OF STROKE IN PATIENTS WHO HAVE HAD TRANSIENT ISCHEMIA OF THE BRAIN OR COMPLETED ISCHEMIC STROKE DUE TO THROMBOSIS
- U-303 METHOD OF USE PATENT-PRODUCT APPROVED FOR TREATMENT OF OSTEOPOROSIS, PAGET'S DISEASE, PREVENTION AND TREATMENT OF GLUCOCORTICOID INDUCED OSTEOPOROSIS
- U-304 A METHOD OF TREATMENT OF A CONDITION INVOLVING AN ANTIBODY ANTIGEN REACTION
- U-305 METHODS FOR USING THE DRUG PRODUCT
- U-306 TREATMENT OF POST-MENOPAUSAL UROGENITAL SYMPTOMS ASSOCIATED WITH ESTROGEN DEFICIENCY
- U-307 CLAIMS AN OLANZAPINE POLYMORPH USEFUL FOR TREATING ANY NUMBER OF LISTED CONDITIONS, INCLUDING SPECIFIC PSYCHOSES, EMPLOYING OLANZAPINE AS PER THE INDICATION OF THIS NDA
- U-308 CLAIMS A SOLID ORAL FORMULATION INCLUDING TABLETS AND GRANULES OF OLANZAPINE USEFUL FOR TREATING ANY NUMBER OF LISTED CONDITIONS, INCLUDING SPECIFIC PSYCHOSES, EMPLOYING OLANZAPINE AS PER THE INDICATIONS OF THIS NDA
- U-309 TREATING SJOEGREN SYNDROME
- U-310 TREATMENT OF XEROSTOMIA
- U-311 HORMONE REPLACEMENT
- U-312 PANIC DISORDER, OBSESSIVE-COMPULSIVE DISORDER, POSTTRAUMATIC STRESS DISORDER
- U-313 TREATMENT OF CONGESTIVE HEART FAILURE
- U-314 METHOD FOR TREATING HYPERPARATHYROIDISM WHICH COMPRISES SUPPRESSING PARATHYROID ACTIVITY
- U-315 METHOD FOR ADMINISTERING DRUG TO GASTROINTESTINAL TRACT
- U-316 METHOD OF TREATING A SUBJECT SUFFERING FROM PROSTATE CANCER
- U-317 METHOD OF USING TROGLITAZONE TO TREAT PATIENTS HAVING INSULIN RESISTANCE
- U-318 TREATMENT OF PATIENTS WITH AN OVERACTIVE BLADDER WITH SYMPTOMS OF URINARY FREQUENCY, URGENCY, OR URGE INCONTINENCE
- U-319 TREATMENT OF MICROBIAL INFECTIONS
- U-320 INHIBITING OR ELIMINATING ACUTE MYELOID LEUKEMIA
- U-321 REDUCTION OF ELEVATED IPTH LEVELS IN THE MGT OF SECONDARY HYPERPARATHYROIDISM IN PATIENTS UNDERGONG CHRONIC RENAL DIALYSIS
- U-322 TREATMENT OF ALZHEIMER'S DEMENTIA
- U-323 USE AS A BILE ACID SEQUESTRANT
- U-324 METHOD OF TREATING AN ANIMAL, INCLUDING A HUMAN, SUFFERING FROM OR SUSCEPTIBLE TO PSYCHOSIS OR ACUTE MANIA EMPLOYING OLANZAPINE
- U-325 METHOD OF TREATING A PATIENT SUFFERING FROM ANY OF A NUMBER OF LISTED CONDITIONS, INCLUDING "BIPOLAR DISORDER NOS" EMPLOYING OLANZAPINE
- U-326 METHOD OF TREATING SCHIZOPHRENIA AND BIPOLAR DISORDER
- U-327 METHOD OF TREATING A PATIENT SUFFERING FROM ANY OF A NUMBER OF LISTED PSYCHOTIC CONDITONS EMPLOYING OLANZAPINE
- U-328 METHOD OF TREATING A PATIENT SUFFERING FROM ANY OF A NUMBER OF LISTED CONDITIONS INCLUDING "A PSYCHOTIC CONDITION" EMPLOYING AN OLANZAPINE POLYMORPH
- U-329 USE OF AVANDIA AS MONOTHERAPY, IN COMBINATION WITH METFORMIN, AND IN COMBINATION WITH SULFONYLUREAS TO IMPROVE GLYCEMIC CONTROL IN PATIENTS WITH TYPE 2 DIABETES MELLITUS
- U-330 TREATMENT OF NAUSEA AND VOMITING
- U-331 METHOD OF TREATING HYPERLIPIDEMIA WITH NICOTINIC ACID BY DOSING ONCE PER DAY IN THE EVENING OR AT NIGHT

PATENT AND EXCLUSIVITY TERMS

ADB 25 of 35

PATENT USE

- U-332 TREATMENT OR PREVENTION OF BRONCHOSPASM
- U-333 METHOD OF TREATING OCULAR HYPERTENSION
- U-334 TREATMENT OF EXCESSIVE FEMALE FACIAL HAIR
- U-335 USE OF PRAVASTATIN SODIUM FOR SECONDARY PREVENTION OF CORONARY EVENTS IN MEN AND WOMEN WHO HAVE HAD A MYOCARDIAL INFARCTION AND HAVE NORMAL CHOLESTEROL LEVELS
- U-336 DIAGNOSTIC RADIOIMAGING
- U-337 USE OF CARDIOLITE/MIRALUMA KIT FOR THE PREPARATION OF TC99M SESTAMIBI
- U-338 METHODS FOR TREATING DISTURBANCES OF MOOD, DISTURBANCES OF APPETITE, DEPRESSED MOOD, OR CARBOHYDRATE CRAVING ALL ASSOCIATED WITH PREMENSTRUAL SYNDROME
- U-339 PREVENTION OF CARDIO-TOXICITY CAUSED BY THE ADMINISTRATION OF DOXORUBICIN
- U-340 THE LONG TERM TREATMENT OF GROWTH FAILURE DUE TO LACK OF ADEQUATE ENDOGENOUS GROWTH HORMONE SECRETION IN CHILDREN
- U-341 METHOD FOR ENHANCING THE TREATMENT OF ... LATE LUTEAL PHASE DYSPHORIC DISORDER
- U-342 METHOD FOR TREATMENT OF LATE LUTEAL PHASE DYSPHORIC DISORDER
- U-343 REDUCTION OF INTESTINAL GAS, CRAMPING AND ANORECTAL IRRITATION
- U-344 METHOD FOR INHIBITING HIV INFECTION BY ADMINISTERING RITONAVIR IN COMBINATION WITH ANOTHER HIV PROTEASE INHIBITOR
- U-345 RITONAVIR AND ANOTHER HIV PROTEASE INHIBITOR FOR CONCOMITANT ADMINISTRATION FOR THE TREATMENT OF AN HIV INFECTION
- U-346 METHOD FOR INHIBITING CYTOCHROME P450 MONOOXYGENASE WITH RITONAVIR AND A METHOD FOR IMPROVING THE PHARMACOKINETICS OF A DRUG THAT IS METABOLIZED BY CYTOCHROME P450 MONOOXYGENASE BY ADMINISTERING THE DRUG AND RITONAVIR
- U-347 METHOD OF USE IN COMBINATION WITH REVERSE TRANSCRIPTASE INHIBITORS
- U-348 METHOD OF USE FOR INHIBITING HIV INFECTION
- U-349 METHOD OF USE WHICH IS SUBJECT OF THE APPLICATION
- U-350 PREPARATION OF A PHARMACEUTICAL COMPOSITION FOR CONCOMITANT ADMINISTRATION WITH A REVERSE TRANSCRIPTASE INHIBITOR
- U-351 INHIBITING PROTEASE WITH LOPINAVIR AND INHIBITING AN HIV INFECTION WITH LOPINAVIR
- U-352 INHIBITING HIV INFECTION BY ADMINISTERING RITONAVIR IN COMBINATION WITH A REVERSE TRANSCRIPTASE INHIBITOR
- U-353 PREVENTION AND TREATMENT OF OSTEOPOROSIS
- U-354 METHOD OF TREATING HYPERLIPIDEMIA WITH NICOTINIC ACID WITHOUT CAUSING TREATMENT-LIMITING ELEVATIONS IN URIC ACID OR GLUCOSE LEVELS OR CAUSING LIVER DAMAGE, BY DOSING ONCE PER DAY IN THE EVENING OR AT NIGHT
- U-355 METHOD OF ASSISTING PERSON TO QUIT SMOKING...TRANSDERMALLY ADMINISTERING NICOTINE VIA...PATCH ADHERED TO SKIN AT DOSING RATE APPROX SAME AS ABSORBED FROM SMOKING
- U-356 DELIVERING A MEDICINAL AEROSOL FORMULATION USING CFC-FREE PROPELLANT 134A.
- U-357 USE OF THE DRUG PRODUCT IN PHOTODYNAMIC THERAPEUTIC PROTOCOLS FOR THE TREATMENT OF AGE-RELATED MACULAR DEGENERATION AND RELATED CONDITIONS INVOLVING UNWANTED NEOVASCULATURE IN THE EYE
- U-358 DEPRESSION, OBSESSIVE COMPULSIVE DISORDER, PANIC DISORDER AND SOCIAL ANXIETY DISORDER
- U-359 METHOD OF USE OF VISICOL
- U-360 METHOD OF TREATING A PATIENT SUFFERING FROM ANY OF A NUMBER OF PATHOLOGICAL PSYCHOLOGICAL CONDITIONS INCLUDING MENTAL DISORDERS EMPLOYING OLANZAPINE AS PER THE INDICATION WHICH IS THE SUBJECT MATTER OF THIS SNDA-011
- U-361 MANAGEMENT OF ANXIETY DISORDERS AND THE SHORT-TERM RELIEF OF THE SYMPTOMS OF ANXIETY
- U-362 USE OF APPROVED FORMULATIONS TO TREAT ALL APPROVED DISEASE INDICATIONS
- U-363 METHOD OF TREATING A PATIENT SUFFERING FROM ANY OF A NUMBER OF PATHOLOGICAL PSYCHOLOGICAL CONDITIONS THAT RELATE TO THE USE OF A PSYCHOACTIVE SUBSTANCE EMPLOYING OLANZAPINE AS PER THE INDICATION THE SUBJECT MATTER OF SUPPLEMENT 011
- U-364 TREATING A PATIENT SUFFERING FROM OR SUSCEPTIBLE TO ANY NUMBER OF LISTED CONDITIONS INCLUDING PSYCHOSIS, EMPLOYING OLANZAPINE AS PER THE INDICATION WHICH IS THE SUBJECT MATTER OF THIS SNDA-011
- U-365 METHOD FOR THE TREATMENT OF CARDIOVASCULAR DISEASE THROUGH THE ADMINISTRATION OF A CALCIUM BLOCKING VASODILATOR IN OUR EXTENDED, CONTROLLED RELEASE FORMULATION
- U-366 METHOD FOR THE TREATMENT OF CARDIOVASCULAR DISEASE THROUGH THE ADMINISTRATION OF A CALCIUM BLOCKING VASODILATOR IN A DELAYED RELEASE FORMULATION
- U-367 TREATMENT OF CARDIOVASCULAR DISORDERS
- U-368 HEARTBURN
- U-369 METHOD OF CONTROLLING AND LOWERING INTRAOCULAR PRESSURE
- U-370 INTRAVAGINAL TREATMENT OF VAGINAL INFECTIONS WITH BUFFERED METRONIDAZOLE COMPOSITIONS
- U-371 APPROVAL FOR MARKETING ONLY UNDER A SPECIAL RESTRICTION PROGRAM APPROVED BY FDA CALLED "SYSTEM FOR THALIDOMIDE EDUCATION AND PRESCRIBING SAFETY" (S.T.E.P.S.)
- U-372 METHOD FOR ADMINISTERING A BENEFICIAL DRUG TO THE GI TRACT OF AN ANIMAL, WHICH METHOD COMPRISES ADMITTING AN OSMOTIC DEVICE ORALLY INTO THE ANIMAL...

PATENT AND EXCLUSIVITY TERMS

ADB 26 of 35

PATENT USE

- U-373 GENERAL USE CLAIM SUBMITTED FOR 12 NEXIUM PATIENTS STATING "PERTINENT TO THE CAPSULE FORMULATION FOR NEXIUM AND ITS INDICATIONS FOR THE TREATMENT OF GERD AND ERADICATION OF H.PYLORI TO REDUCE THE RISK OF DUODENAL ULCER RECURRENCE
- U-374 KIT ADAPTED AND DESIGNED TO PROVIDE BOTH DATA ON THE CURRENT REPRODUCTIVE STATUS OF A PATIENT AND CONTRACEPTION FOR THOSE WHO ARE NOT PREGNANT, BUT RECENTLY ENGAGED IN UNPROTECTED SEX
- U-375 METHOD OF USING RIBAVIRIN FOR TREATING A DISEASE RESPONSIVE TO RIBAVIRIN, E.G. HEPATITIS C
- U-376 TREATMENT OF INFLUENZA
- U-377 METHOD OF TREATING PT WITH CHRONIC HEPATITIS C HAVING HCV GENOTYPE 1 AND VIRAL LOAD GREATER THAN 2 MILLION COPIES/ML TO ERADICATE DETECTABLE HCV-RNA BY ADMIN COMBINATION OF RIBAVIRIN AND INTERFERON ALFA-2B FOR A LEAST 24 WEEKS
- U-378 METHOD FOR TREATING INCONTINENCE
- U-379 METHOD OF TREATING ONYCHROMYCOSIS
- U-380 COMBINATIONS OF TAXOL (PACLITAXEL) AND CISPLATIN WHICH ARE SUITABLE FOR THE TREATMENT OF OVARIAN AND NON-SMALL CELL LUNG CARCINOMAS
- U-381 TREATMENT OF HYPERPHOSPHATEMIA
- U-382 METHOD OF STABILIZING PROSTAGLANDIN
- U-383 METHOD FOR TREATING GLAUCOMA AND OCULAR HYPERTENSION
- U-384 TREATMENT OF CMV RETINITIS
- U-385 TREATMENT OF PEPTIC ULCERS
- U-386 TREATMENT OF PATIENTS SUFFERING FROM A LATE ASTHMATIC REACTION OR LATE PHASE ASTHMA
- U-387 TREATMENT OF PATIENTS WITH RESPIRATORY DISORDERS
- U-388 SMOKING CESSATION AID APPLIED TO THE SKIN
- U-389 SMOKING CESSATION AID APPLIED TO THE SKIN ON WAKING AND REMOVED PRIOR TO SLEEP AFTER ABOUT 16 HOURS
- U-390 METHOD OF USING THE DRUG TO TREAT NEUROIMMUNOLOGIC DISEASES (INCLUDING MULTIPLE SCLEROSIS)
- U-391 USE OF CASODEX IN COMBINATION WITH LHRH AGONISTS FOR THE TREATMENT OF PROSTATE CANCER
- U-392 TREATMENT OF PATIENTS FOR INFLAMMATION
- U-393 MANAGEMENT OF INCONTINENCE, MGT OF HORMONE REPLACEMENT THERAPY, TREATMENT OF INVOLUNTARY INCONTINENCE, MGT OVERACTIVE BLADDER AND INCREASING COMPLIANCE IN SUCH PT
- U-394 METHOD OF USE OF ALPHAGAN
- U-395 METHOD OF USE OF ALPHAGAN P
- U-396 METHOD OF TREATING PEOPLE SUFFERING FROM DEPRESSION
- U-397 METHOD OF TREATING PEOPLE SUFFERING FROM DEPRESSION WITHOUT AN INCREASE IN NAUSEA
- U-398 TREATMENT OF GENERALIZED ANXIETY DISORDER
- U-399 IN-THE-EYE USE OF CHLORINE DIOXIDE CONTAINING COMPOSITIONS
- U-400 USE OF RIBAVIRIN TO INCREASE TYPE 1 CYTOKINE RESPONSE AND SUPPRESS TYPE 2 CYTOKINE RESPONSE TO LYMPHOCYTES, INCLUDING METHODS THAT TAKE ADVANTAGE OF SUCH MODULATION TO TREAT INFECTIONS AND INFESTATIONS
- U-401 USE OF LOPINAVIR IN COMBINATION WITH REVERSE TRANSCRIPTASE INHIBITORS FOR TREATING HIV INFECTION AND IN COMBO WITH OTHER HIV PROTEASE INHIBITORS
- U-402 TREATMENT OF ACTINIC KERATOSES
- U-403 ANTI-ALLERGIC FOR VARIOUS ALLERGIC DISEASES
- U-404 TREATMENT OF ALLERGIC CONJUNCTIVITIS
- U-405 FOR WOMEN WITH SEVERE DIARRHEA-PREDOMINANT IRRITABLE BOWEL SYNDROME (IBS)
- U-406 METHOD OF USE OF ATOVAQUONE AND PROGUANIL
- U-407 METHOD OF TREATING OTOPATHY
- U-408 FOR INDUCING OVULATION IN CONJUNCTION WITH A GONADOTROPIN RELEASING FACTOR ANTAGONIST AND RECRUITING OOCYTES FOR IN-VITRO FERTILIZATION
- U-409 METHOD OF TREATING INFLAMMATION USING DRUG SUBSTANCE
- U-410 METHOD OF REDUCING AMOUNT OF RESPECTIVE ACTIVE COMPONENTS ADMINISTERED TO A DIABETIC PATIENT BY ADMINISTERING A CHEMICAL COMPOUND HAVING A PARTICULAR FORMULA (INCLUDING PIOGLITAZONE) IN COMBINATION WITH AN INSULIN SECRETION ENHANCER
- U-411 METHOD OF REDUCING THE SIDE EFFECTS OF ACTIVE COMPONENTS ADMINISTERED TO A DIABETIC PATIENT BY ADMINISTERING A CHEMICAL COMPOUND HAVING A PARTICULAR FORMULA (WHICH INCLUDES PIOGLITAZONE) IN COMBINATION WITH AN INSULIN PREPARATION
- U-412 TREATMENT OF TYPE 2 DIABETES
- U-413 USE OF THE ACTIVE INGREDIENT FOR INHIBITING THE BIOSYNTHESIS OF CHOLESTEROL AND TREATMENT OF ATHEROSCLEROSIS
- U-414 A METHOD OF TREATING GLYCOMETABOLISM DISORDERS BY ADMINISTERING AN INSULIN SENSITIVITY ENHANCER (INCLUDING PIOGLITAZONE) IN COMBINATION WITH A BIGUANIDE
- U-415 A METHOD FOR REDUCING THE AMOUNT OF ACTIVE COMPONENTS ADMINISTERED TO A DIABETIC PATIENT BY ADMINISTERING AN INSULIN SENSITIVITY ENHANCER (INCLUDING PIOGLITAZONE) IN COMBINATION WITH A BIGUANIDE AS SAID ACTIVE COMPONENTS

PATENT AND EXCLUSIVITY TERMS

ADB 27 of 35

PATENT USE

- U-416 A METHOD FOR REDUCING SIDE EFFECTS OF ACTIVE COMPONENTS ADMINISTERED TO A DIABETIC PATIENT BY ADMINISTERING AN INSULIN SENSITIVITY ENHANCER (INCLUDING PIOGLITAZONE) IN COMBINATION WITH A BIGUANIDE AS SAID ACTIVE COMPONENTS
- U-417 COMBINATION USE OF AD-4833 WITH A BIGUANIDE
- U-418 A METHOD OF TREATING LIPID METABOLISM DISORDERS BY ADMINISTERING A CHEMICAL COMPOUND HAVING A PARTICULAR FORMULA (WHICH INCLUDES PIOGLITAZONE) IN COMBINATION WITH AN INSULIN SECRETION ENHANCER
- U-419 A METHOD OF TREATING LIPID METABOLISM DISORDERS BY ADMINISTERING AN INSULIN SENSITIVITY ENHANCER (INCLUDING PIOGLITAZONE) IN COMBINATION WITH A BIGUANIDE
- U-420 METHOD OF TREATMENT OF TYPE II DIABETES
- U-421 USE FOR SEDATION
- U-422 METHOD OF TREATING AT LEAST ONE OF ATTENTION DEFICIT DISORDER AND ATTENTION DEFICIT HYPERACTIVITY DISORDER
- U-423 METHOD OF TREATING AT LEAST ONE OF ATTENTION DEFICIT DISORDER, ATTENTION DEFICIT HYPERACTIVITY DISORDER, OR AIDS RELATED DEMENTIA
- U-424 FOR ONCE DAILY, BOLUS ADMINISTRATION TO A PATIENT IN ORDER TO ENGENDER TREATMENT FOR A NERVOUS DISORDER FOR SUBSTANTIALLY AN ENTIRE DAY ON A CHRONIC BASIS
- U-425 METHOD OF REDUCING SIDE EFFECTS OF ACTIVE COMPONENTS ADMIN TO A DIABETIC BY ADMIN A CHEMICAL COMPOUND HAVING FORMULA (INCL PIOGLITAZONE) IN COMBINATION WITH AN INSULIN SECRETION ENHANCER
- U-426 PREVENTION OF PREMATURE LH SURGES IN WOMEN UNDERGOING CONTROLLED OVARIAN STIMULATION
- U-427 METHOD OF TREATING ALLERGIC REACTIONS IN MAMMALS
- U-428 METHOD OF TREATING ALLERGY IN A MAMMAL USING THIS ACTIVE METABOLITE
- U-429 METHOD OF USING DESLORATADINE TO TREAT ALLERGIC RHINITIS
- U-430 METHOD OF TREATING A DIABETIC BY ADMINISTERING AN INSULIN SENSITIZER IN COMBINATION WITH AN INSULIN SECRETION ENHANCER, AND A DRUG PRODUCT COMPRISING AN INSULIN SENSITIZER AND AN INSULIN SECRETION ENHANCER
- U-431 POSTTRAUMATIC STRESS DISORDER
- U-432 REDUCTION OF ATHEROSCLEROTIC EVENTS (MYOCARDIAL INFARCTION, STROKE, AND VASCULAR DEATH) IN PATIENTS WITH ATHEROSCLEROSIS DOCUMENTED BY RECENT STROKE, RECENT MYOCARDIAL INFARCTION OR ESTABLISHED PERIPHERAL ARTERIAL DISEASE
- U-433 USE OF LEVOCARITINE IN PREVENTION AND TREATMENT OF CARNITINE DEFICIENCY IN PATIENTS WITH END STAGE RENAL DISEASE WHO ARE UNDERGOING DIALYSIS
- U-434 CONTROLLED SYMPTOMS OF DIARRHEA, BLOATING PRESSURE AND CRAMPS, COMMONLY REFERRED TO AS GAS
- U-435 A TITRATION DOSING REGIMEN FOR THE TREATMENT OF PAIN USING AN INITIAL DOSE OF ABOUT 25MG
- U-436 ACUTE TREATMENT OF MIGRAINE ATTACKS WITH OR WITHOUT AURA IN ADULTS
- U-437 METHOD OF USE EQUAL TO PROCESS OF PREPARATION
- U-438 TREATMENT/PREVENTION OF NEURODEGENERATIVE DISEASE
- U-439 TREATMENT OF OBESITY
- U-440 METHOD FOR TRANSDERMAL ADMINISTRATION OF A DRUG THROUGH NON-SCROTAL SKIN USING A TRANSDERMAL DRUG DELIVERY DEVICE CONTAINING THE DRUG AND HAVING AN ADHESIVE SURFACE
- U-441 METHOD OF TREATING MS BY ADMINISTERING COPAXONE
- U-442 METHOD FOR DELIVERING A DRUG TO A PATIENT IN NEED OF THE DRUG, WHILE AVOIDING THE OCCURENCE OF AN ADVERSE SIDE EFFECT KNOWN OR SUSPECTED OF BEING CAUSED BY SAID DRUG
- U-443 MANAGEMENT OF MODERATE TO SEVERE PAIN WHEN A CONTINUOUS, AROUND-THE-CLOCK ANALGESIC IS NEEDED FOR AN EXTENDED PERIOD OF TIME
- U-444 TREATMENT OF MIGRAINE
- U-445 USE AS AN ANTIMYCOTIC AGENT
- U-446 TOPICAL TREATMENT OF OCULAR HYPERTENSION AND GLAUCOMA
- U-447 METHOD OF TREATING HYPERLIPIDEMIA WITH NICOTINIC ACID BY DOSING ONCE PER DAY IN THE EVENING OR AT NIGHT
- U-448 METHOD OF TREATING HYPERLIPIDEMIA WITH NICOTINIC ACID WITHOUT CAUSING TREATMENT-LIMITING ELEVATIONS IN URIC ACID OR GLUCOSE LEVELS OR CAUSING LIVER DAMAGE, BY DOSING ONCE PER DAY IN THE EVENING OR AT NIGHT
- U-449 USE IN COMBINATION WITH 5-FLUOROURACIL AND LEUCOVORIN FOR THE TREATMENT OF METASTATIC COLORECTAL CANCER WHERE THE DOSE OF LEUCOVORIN IS AT LEAST 200MG PER SQUARE METER
- U-450 INTERMEDIATE REL NICOTINIC ACID FORMULATIONS HAVING UNIQUE URINARY METAB PROFILES RESULTING FROM ABSORPTION PROFILES OF NICOTINIC ACID FROM THE INTERMEDIATE NICOTINIC ACID FORMULATIONS, SUITABLE FOR TX HYPERLIPIDEMIA FOLLOWING QD DOSING
- U-451 TREATMENT OF DEPRESSION AND GENERALIZED ANXIETY DISORDER
- U-452 USE OF LANSOPRAZOLE FOR COMBATTING DISEASES CAUSED BY THE GENUS CAMPYLOBACTER (C.PYLORI=H.PYLORI)
- U-453 TREATMENT OF PLATELET ASSOCIATED ISCHEMIC DISORDERS
- U-454 METHOD OF TX A PT SUSPECTED OF HAVING HEPATITIS C BY ADMIN, IN COMBINATION, A CONJUGATE COMPRISING PEG 12000 & INTERFERON ALFA-2B IN AN AMT OF FROM 0.5MCG/KG TO 2MCG/KG, ONCE WEEKLY, AND RIBAVIRIN

PATENT AND EXCLUSIVITY TERMS

ADB 28 of 35

PATENT USE

U-455 TREATMENT OF PULMONARY HYPERTENSION WITH UT-15
U-456 METHOD OF DECREASING THE PRODUCTION OF A-BETA USING A COMPOSITION WHICH DECREASES BLOOD CHOLESTEROL IN PATIENTS AT RISK OF OR EXHIBITING SYMPTOMS OF ALZHEIMER'S DISEASE
U-457 METHOD OF TREATING A VAGINAL FUNGAL INFECTION IN A FEMALE HUMAN
U-458 METHOD OF USE OF IMAGENT
U-459 TREATMENT OF DEPRESSION AND GENERALIZED ANXIETY DISORDER
U-460 METHOD OF TREATING PSYCHIATRIC SYMPTOMS ASSOCIATED WITH PREMENSTRUAL DISORDERS USING SERTRALINE
U-461 METHOD OF TREATMENT OF LATE LUTEAL PHASE DYSPHORIC DISORDER (PMDD) USING SERTRALINE
U-462 SIGNS AND SYMPTOMS OF OSTEOARTHRITIS AND ADULT RHEUMATOID ARTHRITIS AND TREATMENT OF PRIMARY DYSMENORRHEA
U-463 VENOGRAPHY
U-464 PERIPHERAL ARTERIOGRAPHY
U-465 CT IMAGING OF THE HEAD
U-466 TREATMENT OF IRRITABLE BOWEL SYNDROME
U-467 USE OF EPLERENONE IN COMBINATION WITH AN ANGIOTENSIN CONVERTING ENZYME (ACE) INHIBITOR FOR TREATING HYPERTENSION
U-468 METHOD OF USING FEXOFENADINE HCL IN TREATING ALLERGIC RHINITIS
U-469 TREATMENT OF GASTROESOPHAGEAL REFLEX DISEASE (GERD) AND ERADICATION OF H.PYLORI TO REDUCE RISK OF DUODENAL ULCER RECURRENCE
U-470 THERAPY IN CHRONIC HEPATITIS B VIRUS INFECTION
U-471 METHOD OF TREATING A PATIENT SUFFERING FROM DIABETES MELLITUS
U-472 TREATMENT OF ATTENTION DEFICIT HYPERACTIVITY DISORDER USING METHYLPHENIDATE BI-MODAL RELEASE PROFILE EXTENDED-RELEASE CAPSULES
U-473 TO REDUCE PLASMA CHOLESTEROL LEVELS IN A MAMMAL
U-474 TO REDUCE PLASMA CHOLESTEROL LEVELS BY ADMIN EZETIMIBE IN COMBO WITH CHOLESTEROL BIOSYNTHESIS INHIB SELECTED FROM GROUP CONSISTING OF HMG COA REDUCTASE INHIBITORS INCL SIMVASTATIN
U-475 TREATMENT OF CUTANEOUS MANIFESTATIONS OF CUTANEOUS T-CELL LYMPHOMA IN PATIENTS WHO ARE REFRACTORY TO AT LEAST ONE PRIOR SYSTEMIC THERAPY
U-476 METHOD OF TREATING ANDROGEN RESPONSIVE/MEDIATED CONDITION IN MAMMAL BY ADMIN A SAFE, EFFECTIVE AMOUNT OF DUTASTERIDE OR PHARMACEUTICALLY ACCEPTABLE DERIVATIVE THEREOF..CONDITIONS INCLUDE BENIGN PROSTATIC HYPERTROPHY
U-477 METHOD OF INHIBITING 5 ALPHA TESTOSTERONE REDUCTASE ENZYME WITH DUTASTERIDE OR ITS DERIVATIVE AND TREATING ANDROGEN RESPONSIVE/MEDIATED DISEASE INCLUDING BENIGN PROSTATIC HYPERPLASIA
U-478 METHOD OF TREATING HEPATITIS C VIRAL INFECTION BY CONTINUOUS PARENTERAL ADMIN INTERFERON ALPHA 2-10 MILLION IU WEEKLY, SUBCUTANEOUSLY, INJECTION OF POLYMER-INTERFERON ALPHA CONJUGATE-POLYMER IS PEG-INTERFERON IS ALPHA 2B
U-479 METHOD OF USING PEG-INTRON/REBETOL COMBINATION THERAPY AND INTRON/REBETOL COMBINATION THERAPY
U-480 CONTRAST AGENT FOR MRI
U-481 DISUBSTITUTED ACETYLENES BEARING HETEROAROMATIC AND HETEROBICYCLIC GROUPS HAVING RETINOID-LIKE ACTIVITY
U-482 METHOD OF IN VITRO FERTILIZATION THERAPY INCLUDING MEANS FOR INDUCING OVULATION....
U-483 METHOD FOR THE ADMINISTRATION OF DRUGS USING THAT COMPOUND
U-484 METHOD OF TREATING A SKIN DISEASE WITH A CORTICOSTEROID-CONTAINING PHARMACEUTICAL COMPOSITION
U-485 METHOD AND COMPOSITION FOR REDUCING NERVE INJURY PAIN ASSOCIATED WITH SHINGLES (HERPES ZOSTER AND POST-HERPETIC NEURALGIA)
U-486 EXTERNAL PREPARATION FOR APPLICATION TO THE SKIN CONTAINING LIDOCAINE-DRUG RETAINING LAYER PLACED ON SUPPORT AND COMPRISES ADHESIVE GEL BASE 1-10% BY WEIGHT OF LIDOCAINE
U-487 METHOD AND COMPOSITION FOR REDUCING NERVE INJURY PAIN ASSOCIATED WITH SHINGLES (HERPES ZOSTER AND POST-HERPETIC NEURALGIA)
U-488 METHOD FOR REDUCING THE PAIN ASSOCIATED WITH HERPES-ZOSTER AND POST-HERPETIC NEURALGIA
U-489 EXPECTORANT
U-490 TESTOSTERONE REPLACEMENT THERAPY IN MALES FOR CONDITIONS ASSOCIATED WITH A DEFICIENCY OR ABSENCE OF ENDOGENOUS TESTOSTERONE
U-491 METHOD OF DELIVERING A DRUG TO THE LUNG
U-492 METHOD FOR THE TREATMENT OF SKIN, SUFFERING FROM A CONDITION SELECTED FROM A GROUP CONSISTING OF NONACNE INFLAMMATORY DERMATOSES... COMPRISING APPLYING TO AFFECTED AREA. A THERAPEUTICALLY EFFECTIVE AMT AZELAIC ACID
U-493 TREATMENT OF TYPE 2 DIABETES MELLITUS
U-494 TREATMENT OF ATTENTION-DEFICIT HYPERACTIVITY DISORDER
U-495 PERITONEAL DIALYSIS SOLUTION
U-496 METHOD FOR TREATING CHRONIC RENAL FAILURE

PATENT AND EXCLUSIVITY TERMS

ADB 29 of 35

PATENT USE

- U-497 RELIEF OF THE SIGNS AND SYMPTOMS OF OSTEOARTHRITIS AND RHEUMATOID ARTHRITIS
- U-498 INTRA-ARTERIAL AND INTRAVENOUS USES OF ULTRAVIST
- U-499 METHOD OF USING REBETOL CAPSULES IN COMBINATION WITH A CONJUGATE COMPRISING POLYETHYLENE GLYCOL(PEG) AND AN ALPHA INTERFERON, INCLUDING, FOR EXAMPLE, PEG-INTRON POWDER FOR INJECTION
- U-500 USE AS AN ANTIHYPERTENSIVE AGENT
- U-501 TREATMENT OF RECURRENT HERPES LABIALIS (COLD SORES) IN ADULTS
- U-502 PITYRIASIS VERSICOLOR
- U-503 GENERATOR MUST BE USED WITH INFUSION SYSTEM SPECIFICALLY LABELED FOR USE WITH GENERATOR
- U-504 TINEA PEDIS, TINEA CRURIS, TINEA CORPORIS
- U-505 ULTRASOUND CONTRAST AGENT
- U-506 PHARM PRODUCT CONTAINER 1ST CHAMBER IS DISPOSED AQUEOUS DILUENT SOL 2ND CHAMBER PHARM ACTIVE AGENT COMPRISING ACETYLCHOLINE,BUFFER IN 1ST CHAM IS SUFFICIENT TO BUFFER PH OF MIXED SOL RESULTING MIXTURE OF AQUEOUS DILUENT SOL & PHARM ACTIVE..
- U-507 ACROMEGALY IN PATIENTS W/INADEQUATE RESPONSE TO SURGERY AND/OR RADIATION THERAPY AND/OR MEDICAL THERAPIES, OR FOR WHOM THESE THERAPIES ARE NOT APPROPRIATE
- U-508 METHOD OF RELEASING 17-BETA OESTRADIOL PRECURSOR IN A SUBSTANTIALLY ZERO ORDER PATTERN FOR AT LEAST THREE WEEKS
- U-509 TREATMENT OF CUTANEOUS MANIFESTATIONS OF CUTANEOUS T-CELL LYMPHOMA IN PATIENTS WHO ARE REFRACTORY TO AT LEAST ONE PRIOR SYSTEMIC THERAPY
- U-510 TOPICAL TREATMENT OF CUTANEOUS LESIONS IN PATIENTS WITH CUTANEOUS T-CELL LYMPHOMA (STAGE IA AND IB) WHO HAVE REFRACTORY OR PERSISTENT DISEASE AFTER OTHER THERAPIES OR WHO HAVE NOT TOLERATED OTHER THERAPIES
- U-511 USE OF QUINOLONE COMPOUNDS AGAINST ANAEROBIC PATHOGENIC BACTERIA
- U-512 USE OF QUINOLONE COMPOUNDS AGAINST ATYPICAL UPPER RESPIRATORY PATHOGENIC BACTERIA
- U-513 METHODS OF USE OF ANTIMICROBIAL COMPOUNDS AGAINST PATHOGENIC MYCOPLASMA BACTERIA
- U-514 PREVENTION OF OVULATION IN A WOMAN
- U-515 TREATMENT OF MULTIPLE MYELOMA PATIENTS WHO HAVE RECEIVED AT LEAST TWO PRIOR THERAPIES AND HAVE DEMONSTRATED DISEASE PROGRESSION ON THE LAST THERAPY
- U-516 METHOD OF TREATING A PSYCHOTIC DISEASE
- U-517 STABLE GEL FORMULATION FOR TOPICAL TREATMENT OF SKIN CONDITIONS
- U-518 OBSESSIVE COMPULSIVE DISORDER
- U-519 POST OPERATIVE NAUSEA AND VOMITING
- U-520 PREMENOPAUSAL OSTEOPOROSIS
- U-521 METHOD OF USING RIBAVIRIN IN COMBINATION WITH INTRON A (INTERFERON ALPHA-2 B RECOMBINANT) INJECTION TO TREAT PATIENTS WITH CHRONIC HEPATITIS C
- U-522 TREATMENT OF CMV RETINITIS BY INTRAVITREAL ADMIN OF A PHOSPHOROTHIOATE OLIGONUCLEOTIDE CAPABLE OF HYBRIDIZING WITH CMV MRNA
- U-523 METHOD OF TREATING INFECTION BY CRYPTOSPORIDIUM PARVUM IN AN IMMUNOCOMPROMISED MAMMAL
- U-524 METHOD OF TREATING DIARRHEA
- U-525 METHOD OF TREATING PARASITIC INFECTIONS
- U-526 METHOD OF PROVIDING CONTROLLED RELEASE OF A TREATING AGENT USING A CONTROLLED RELEASE COMPOSITION
- U-527 METHOD OF DELIVERING AN ACTIVE INGREDIENT USING A PROGRESSIVE HYDRATION BIOADHESIVE
- U-528 PREVENTION OF CHEMOTHERAPY-INDUCED NAUSEA AND VOMITING
- U-529 ONCE DAILY TREATMENT OF ASTHMA WITH NEBULIZED BUDESONIDE
- U-530 TREATMENT OF HERPES ZOSTER, TREATMENT OF GENITAL HERPES, TREATMENT OF COLD SORES, SUPPRESSION OF GENITAL HERPES IN IMMUNOCOMPETENT AND HIV-INFECTED INDIVIDUALS, REDUCTION OF RISK OF HETEROSEXUAL TRANSMISSION OF GENITAL HERPES
- U-531 TREATMENT OF PATIENTS WITH ESSENTIAL HYPERTENSION. MAY BE USED ALONE OR GIVEN WITH OTHER CLASSES OF ANTIHYPERTENSIVES, ESPECIALLY THIAZIDE DERIVATIVES
- U-532 TREATMENT OF BRONCHOSPASM ASSOCIATED WITH COPD IN PATIENTS REQUIRING MORE THAN ONE BRONCHO DILATOR
- U-533 ERECTILE DYSFUNCTION
- U-534 HUMALOG IS AN INSULIN ANALOG THAT IS INDICATED IN THE TREATMENT OF PATIENTS WITH DIABETES MELLITUS FOR THE CONTROL OF HYPERGLYCEMIA
- U-535 TREATMENT OF SOCIAL ANXIETY DISORDER
- U-536 CONTRAST AGENT FOR MAGNETIC RESONANCE IMAGING
- U-537 TREATMENT OF CONDITIONS RELATED TO HYPERALDOSTERONISM SUCH AS HYPERTENSION AND CARDIAC INSUFFICIENCY, WITH EPLERENONE
- U-538 FIRST LINE TREATMENT OF SEVERE HYPERTENSION, IN PATIENTS WITH HYPERTENSION SEVERE ENOUGH THAT THE VALUE OF ACHIEVING PROMPT BLOOD PRESSURE CONTROL EXCEEDS THE RISK OF INITIATING COMBINATION THERAPY IN THESE PATIENTS
- U-539 TREATMENT OF MODERATE TO SEVERE DEMENTIA OF THE ALZHEIMER'S TYPE
- U-540 TREATMENT OF FUNGAL INFECTIONS

PATENT AND EXCLUSIVITY TERMS

ADB 30 of 35

PATENT USE

- U-541 METHOD OF TREATMENT OF ADULTS INFECTED WITH HIV-1
- U-542 METHOD OF TREATING PATIENT WITH TYPE 2 DIABETES BY ONCE DAILY ADMINISTRATION
- U-543 TREATMENT OF SCHIZOPHRENIA
- U-544 TREATMENT OF OVERACTIVE BLADDER. TREATMENT OF URINARY INCONTINENCE.
- U-545 METHOD FOR THE PREVENTION AND/OR TREATMENT OF THROMBOTIC EPISODES, SUCH AS MYOCARDIAL INFARCTION, IN A HUMAN PATIENT AND METHOD FOR THE PREVENTION OF VENOUS THROMBOSIS IN A POSTOPERATIVE HUMAN PATIENT
- U-546 USE OF REPAGLINIDE IN COMBINATION WITH METFORMIN TO LOWER BLOOD GLUCOSE
- U-547 MAINTENANCE MONOTHERAPY FOR BIPOLAR DISORDER
- U-548 A METHOD OF REDUCING FLUSH IN AN INDIVIDUAL BEING TREATED FOR A LIPIDEMIC DISORDER AND EFFECTIVELY TREATING THE LIPIDEMIC DISORDER
- U-549 USE IN THE TREATMENT OF MEN WITH ADVANCED SYMPTOMATIC PROSTATE CANCER
- U-550 TREATMENT OF BIPOLAR MANIA AND SCHIZOPHRENIA
- U-551 METHOD FOR REDUCING TOXICITY OF ALIMTA TREATED PATIENTS BY ADMINISTERING FOLIC ACID
- U-552 TREATMENT OF HYPERTENSION AND HYPERLIPIDEMIA WITH A SINGLE COMPOSITION
- U-553 MANAGEMENT OF PAIN AND DISCOMFORT ASSOCIATED WITH PERIDONTAL SCALING AND ROOT PLANNING PROCEDURES BY APPLICATION OF AN EUTECTIC MIXTURE OF LOCAL ANESTHETICS TO PERIDONTAL POCKETS
- U-554 TREATING HIV INFECTION WITH INDINAVIR SULFATE IN COMBINATION WITH ANTIRETROVIRAL AGENTS
- U-555 TREATMENT OF COMPLICATED URINARY TRACT INFECTIONS AND ACUTE UNCOMPLICATED PYELONEPHRITIS
- U-556 USE AS ADJUNCT DIAGNOSTIC FOR SERUM THYROGLOBULIN (TG) TESTING
- U-557 NASAL TREATMENT OF SEASONAL AND PERENNIAL ALLERGIC RHINITIS SYMPTOMS
- U-558 INDICATED FOR THE RELIEF OF BRONCHOSPASM IN PATIENTS 2-12 YEARS OF AGE WITH ASTHMA (REVERSIBLE OBSTRUCTIVE AIRWAY DISEASE)
- U-559 METHOD OF DECREASING OR REDUCING PARATHYROID HORMONE LEVEL; METHOD OF MODULATING PARATHYROID HORMONE SECRETION;METHOD OF TREATING HYPERPARATHYROIDISM; METHOD OF REDUCING SERUM IONIZED CALCIUM LEVEL
- U-560 METHOD OF DECREASING PARATHYROID HORMONE LEVEL;METHOD OF TREATING HYPERPARATHYROIDISM
- U-561 COSOPT IS INDICATED FOR THE REDUCTION OF ELEVATED INTRAOCULAR PRESSURE IN PATIENTS WITH OPEN-ANGLE GLAUCOMA OR OCULAR HYPERTENSION WHO ARE INSUFFICIENTLY RESPONSIVE TO BETA BLOCKERS
- U-562 TOPICAL TREATMENT OF CUTANEOUS LESIONS IN PATIENTS WITH AIDS-RELATED KAPOSI'S SARCOMA
- U-563 MARINOL IS INDICATED FOR, INTER ALIA, ANOREXIA ASSOCIATED WITH WEIGHT LOSS IN PATIENTS WITH AIDS
- U-564 TREATMENT OF HIV IN CONCOMITANT THERAPY
- U-565 TREATMENT OF SEASONAL AND PERENNIAL ALLERGIC RHINITIS SYMPTOMS, AND CHRONIC URTICARIA
- U-566 FOR THE LONG-TERM, ONCE-DAILY, MAINTENANCE TREATMENT OF BRONCHOSPASM ASSOCIATED WITH CHRONIC OBSTRUCTIVE PULMONARY DISEASE (COPD), INCLUDING CHRONIC BRONCHITIS AND EMPHYSEMA
- U-567 METHOD OF TREATING INFERTILITY
- U-568 METHOD OF USING FSH ALONE (WITHOUT EXOGENOUS LH) IN IN VITRO FERTILIZATION
- U-569 METHOD OF USING FSH ALONE (WITHOUT EXOGENOUS LH) IN IN VITRO FERTILIZATION AND WHEREIN THEREAFTER AN OVULATORY INDUCING AMOUNT OF HCG IS ADMINISTERED
- U-570 METHOD OF USING FSH ALONE (WITHOUT EXOGENOUS LH) IN IN VITRO FERTILIZATION AND WHEREIN THE DAILY AMOUNT OF FSH IS ABOUT 5-10 IU/KG
- U-571 TREATMENT OF AGITATION ASSOCIATED WITH SCHIZOPHRENIA AND BIPOLAR I MANIA
- U-572 INTENSIVE CARE UNIT SEDATION
- U-573 TREATMENT OF ACUTE PROMYELOGENOUS LEUKEMIA (APL)
- U-574 PROPHYLAXIS AND TREATMENT OF THE NASAL SYMPTOMS OF SEASONAL ALLERGIC RHINITIS AND TREATMENT OF THE NASAL SYMPTOMS OF PERENNIAL ALLERGIC RHINITIS IN ADULTS AND PEDIATRIC PATIENTS 12 YEARS OF AGE AND OLDER
- U-575 LOTEMAX OPHTHALMIC SUSPENSION IS INDICATED FOR THE TREATMENT OF STEROID RESPONSIVE CONDITIONS OF THE PALPEBRAL BULBAR CONJUNCTIVA, CORNEA AND ANTERIOR SEGMENT OF THE GLOBE.
- U-576 ALREX OPHTHALMIC SUSPENSION IS INDICATED FOR THE TEMPORARY RELIEF OF THE SIGNS AND SYMPTOMS OF SEASONAL ALLERGIC CONJUNCTIVITIS.
- U-577 TREATMENT OF BENIGN PROSTATIC HYPERPLASIA WITH FINASTERIDE IN COMBINATION WITH DOXAZOSIN
- U-578 TREATMENT OF COMMUNITY ACQUIRED PNEUMONIA, ACUTE EXACERBATION OF CHRONIC BRONCHITIS, AND ACUTE BACTERIAL SINUSITIS CAUSED BY SUSCEPTIBLE STRAINS OF DESIGNATED MICROORGANISMS IN PATIENTS 18 YEARS AND OLDER.
- U-579 TREATMENT OF EPILEPSY AND/OR MIGRAINE.
- U-580 TREATMENT OF DISORDERS OF THE SEROTONERGIC SYSTEM SUCH AS DEPRESSION AND ANXIETY-RELATED DISORDERS
- U-581 METHOD OF TREATING A CONDITION CAPABLE OF TREATMENT BY INHALATION, E.G. ASTHMA, COMPRISING ADMINISTRATION OF A FORMULATION CLAIMED IN US PATENT NO. 6743413
- U-582 METHOD FOR THE TREATMENT OF A RESPIRATORY DISORDER, E.G. ASTHMA, COMPRISING ADMINISTERING AN EFFECTIVE AMOUNT OF AN AEROSOL COMPOSITION TO A PATIENT FROM A METERED DOSE INHALER SYSTEM AS CLAIMED IN US PATENT NO. 6253762

PATENT AND EXCLUSIVITY TERMS

ADB 31 of 35

PATENT USE

- U-583 METHOD FOR THE TREATMENT OF A RESPIRATORY DISORDER, E.G. ASTHMA, COMPRISING ADMINISTERING TO A PATIENT BY INHALATION, A METERED AEROSOL DOSE OF A DRUG FORMULATION FROM THE METERED DOSE INHALER SYSTEM CLAIMED IN US 6546928
- U-584 SINGLE-DOSE ADMINISTRATION BY THE EPIDURAL ROUTE, AT THE LUMBAR LEVEL, FOR THE TREATMENT OF PAIN FOLLOWING MAJOR SURGERY
- U-585 TO PROMOTE WEIGHT GAIN AFTER WEIGHT LOSS IN CERTAIN TYPES OF PATIENTS
- U-586 AN INTERMEDIATE RELEASE NICOTINIC ACID FORMULATION SUITABLE FOR ORAL ADMINISTRATION ONCE-A-DAY AS A SINGLE DOSE FOR TREATING HYPERLIPIDEMIA WITHOUT CAUSING DRUG-INDUCED HEPATOTOXICITY OR ELEVATIONS IN URIC ACID OR GLUCOSE OR BOTH
- U-587 USE OF EPLERENONE IN COMBINATION WITH AN ANGIOTENSIN CONVERTING ENZYME (ACE) INHIBITOR (AND OPTIONALLY A DIURETIC) FOR TREATING CONGESTIVE HEART FAILURE AND HYPERTENSION
- U-588 SHORT-TERM TREATMENT OF ACTIVE DUODENAL ULCER; TREATMENT OF HEARTBURN AND OTHER SYMPTOMS ASSOCIATED WITH GERD; SHORT-TERM TREATMENT OF EROSIIVE ESOPHAGITIS; MAINTENANCE OF HEALING OF EROSIIVE ESOPHAGITIS
- U-589 METHOD FOR TREATMENT OF A RESPIRATORY DISORDER, E.G., BRONCHOSPASM, COMPRISING ADMINISTERING AN EFFECTIVE AMOUNT OF AN AEROSOL COMPOSITION TO A PATIENT FROM A METERED DOSE INHALER SYSTEM AS CLAIMED IN U.S. PATENT NO. 6131966
- U-590 METHOD FOR TREATMENT OF A RESPIRATORY DISORDER, E.G., BRONCHOSPASM, COMPRISING ADMINISTERING TO A PATIENT BY ORAL OR NASAL INHALATION A DRUG FORMULATION BY USING THE METERED DOSE INHALER SYSTEM AS CLAIMED IN US PATENT NO. 6532955
- U-591 TREATMENT OF ATTENTION DEFICIT HYPERACTIVITY DISORDER USING A DOSAGE FORM WHICH PROVIDES ONCE-DAILY ORAL ADMINISTRATION OF A PHENIDATE DRUG
- U-592 TREATMENT OF PRIMARY HYPERCHOLESTEROLEMIA, MIXED HYPERLIPIDEMIA AND/OR HOMOZYGOUS FAMILIAL HYPERCHOLESTEROLEMIA (HOFA)
- U-593 TREATMENT OF PRIMARY HYPERCHOLESTEROLEMIA, MIXED HYPERLIPIDEMIA AND/OR HOMOZYGOUS FAMILIAL HYPERCHOLESTEROLEMIA (HOFH)
- U-594 PREVENTION OF POSTMENOPAUSAL OSTEOPOROSIS
- U-595 35 MG ORALLY ONCE A WEEK FOR PREVENTION OF OSTEOPOROSIS IN POSTMENOPAUSAL WOMEN; 35 MG ORALLY ONCE A WEEK FOR TREATMENT OF OSTEOPOROSIS IN POSTMENOPAUSAL WOMEN
- U-596 TREATMENT OF HORMONE RECEPTOR POSITIVE METASTATIC BREAST CANCER IN POSTMENOPAUSAL WOMEN WITH DISEASE PROGRESSION FOLLOWING ANTIESTROGEN THERAPY
- U-597 FORTEO IS INDICATED FOR THE TREATMENT OF POST MENOPAUSAL WOMEN WITH OSTEOPOROSIS WHO ARE AT HIGH RISK FOR FRACTURE
- U-598 PROPHYLACTIC TREATMENT OF MIGRAINE
- U-599 METHOD FOR TREATING ALLERGIC CONJUNCTIVITIS
- U-600 A METHOD OF TREATING A PATIENT IN NEED OF OPHTHALMIC ANTIMICROBIAL THERAPY WITH LEVOFLOXACIN
- U-601 TREATMENT OF BIPOLAR DISORDERS
- U-602 SIGNS AND SYMPTOMS OF OSTEOARTHRITIS, RHEUMATOID ARTHRITIS IN ADULTS, AND/OR PAUCIARTICULAR OR POLYARTICULAR COURSE JUVENILE RHEUMATOID ARTHRITIS, ACUTE PAIN IN ADULTS; PRIMARY DYSMENORRHEA; AND/OR ACUTE MIGRAINE ATTACKS IN ADULTS
- U-603 METHOD OF TREATING INFECTIONS COMPRISING ORALLY ADMINISTERING AN EFFECTIVE AMOUNT OF THE FDA APPROVED ORAL SUSPENSION
- U-604 METHOD OF LOWERING BLOOD GLUCOSE BY ONCE DAILY ADMINISTRATION
- U-605 TREATMENT OF MAJOR DEPRESSIVE DISORDER(MDD);ALTHOUGH THE MECHANISM OF THE ANTIDEPRESSANT ACTION OF DULOXETINE IN HUMANS IS UNKNOWN, IT IS BELIEVED TO BE RELATED TO ITS POTENTIATION OF SERATONERGIC AND NORADRENERGIC ACTIVITY IN THE CNS
- U-606 USE OF IRINOTECAN IN COMBINATION WITH 5-FLUOROURACIL AND LEUCOVORIN FOR THE TREATMENT OF METASTATIC COLRECTAL CANCER
- U-607 CANCIDAS IS INDICATED FOR EMPIRICAL THERAPY FOR PRESUMED FUNGAL INFECTIONS IN FEBRILE, NEUTROPENIC PATIENTS.
- U-608 USE OF QUINOLONE COMPOUNDS AGAINST PNEUMOCOCCAL PATHOGENIC BACTERIA
- U-609 USE OF QUINOLONE COMPOUNDS AGAINST QUINOLONE-RESISTANT PNEUMOCOCCAL PATHOGENIC BACTERIA
- U-610 ATROVENT HFA (IPRATROPIUM BROMIDE HFA) INHALATION AEROSOL IS INDICATED AS A BRONCHODILATOR FOR MAINTENANCE TREATMENT OF BRONCHOSPASM ASSOCIATED WITH CHRONIC OBSTRUCTIVE PULMONARY DISEASE, INCLUDING CHRONIC BRONCHITIS AND EMPHYSEMA.
- U-611 METHOD OF USING DESLORATADINE TO TREAT SEASONAL AND PERENNIAL ALLERGIC RHINITIS, PRURITIS, AND CHRONIC IDIOPATHIC URTICARIA IN PATIENTS 2 YEARS OF AGE AND OLDER
- U-612 TREATMENT OF SEASONAL ALLERGY SYMPTOMS WITH NASAL CONGESTION IN ADULTS AND CHILDREN 12 YEARS OF AGE AND OLDER
- U-613 REDUCTION OF SERUM PHOSPHATE
- U-614 TREATMENT OF SEXUAL DYSFUNCTION
- U-615 ADJUNCTIVE THERAPY TO DIET IN ADULTS TO REDUCE LDL-C, TOTAL-C, TRIGLYCERIDES AND APO B, AND INCREASE HDL-C IN PATIENTS WITH PRIMARY HYPERCHOLESTEROLEMIA OR MIXED DYSLIPIDEMIA (TYPES IIA, IIB) AND TO TREAT HYPERTRIGLYCERIDEMIA (TYPES IV, V)
- U-616 MANAGEMENT OF PERSISTENT, MODERATE TO SEVERE PAIN IN PATIENTS REQUIRING CONTINUOUS, AROUND-THE-CLOCK ANALGESIA WITH A HIGH POTENCY OPIOID FOR AN EXTENDED PERIOD OF TIME GENERALLY WEEKS TO MONTHS OR LONGER

PATENT AND EXCLUSIVITY TERMS

ADB 32 of 35

PATENT USE

U-617 TREATMENT OF ACUTE PROMYELOGENOUS LEUKEMIA (APL)
U-618 USE OF ROSUVASTATIN CALCIUM TO REDUCE ELEVATED TOTAL-C, LDL-C, APOB, NONHDL-C OR TG LEVELS; TO INCREASE HDL-C; AND TO TREAT PATIENTS WITH PRIMARY HYPERCHOLESTOLEMIA
U-619 TREATMENT OF MALIGNANT NEOPLASM
U-620 TREATMENT OF INSOMNIA
U-621 METHOD OF TREATING CANCER
U-622 TREATMENT OF VEGF MEDIATED OCULAR DISEASE.
U-623 SHORT TERM TREATMENT OF ACTIVE BENIGN GASTRIC ULCER
U-624 REDUCTION OF RISK OF UPPER GASTROINTESTINAL BLEEDING IN CRITICALLY ILL PATIENTS
U-625 ALLERGIC RHINITIS OR NASAL POLYPS
U-626 CLOLAR IS INDICATED FOR THE TREATMENT OF PEDIATRIC PATIENTS 1 TO 21 YEARS OLD WITH RELAPSED OR REFRACTORY ACUTE LYMPHOBLASTIC LEUKEMIA AFTER AT LEAST TWO PRIOR REGIMENS
U-627 TREATMENT OF PATIENTS USING EXTENDED-RELEASE CARBAMAZEPINE
U-628 USE OF AVANDIA IN COMBINATION WITH A SULFONYLUREA, AND IN COMBINATION WITH METFORMIN AND A SULFONYLUREA TO IMPROVE GLYCEMIC CONTROL IN PATIENTS WITH TYPE 2 DIABETES MELLITUS
U-629 METHOD OF INDUCING A HYPNOTIC OR SEDATIVE EFFECT IN A HUMAN BY ADMINISTERING ESZOPICLONE
U-630 TREATING URINARY INCONTINENCE BY ADMINISTERING AN EXTENDED-RELEASE FORM OF DARIFENACIN
U-631 TREATING A DISEASE OF ALTERED MOTILITY OR TONE OF SMOOTH MUSCLE BY ADMINISTERING A MUSCARINIC RECEPTOR ANTAGONIZING AMOUNT OF DARIFENACIN
U-632 METHOD OF TREATMENT OF CANCER BY ADMINISTERING PARTICLES OF PACLITAXEL THAT HAVE A PROTEIN COATING
U-633 METHOD FOR TREATMENT OF TUMORS BY ADMINISTERING PACLITAXEL AT A DOSE IN THE RANGE OF ABOUT 30MG/METER SQUARE TO ABOUT 100MG/METER SQUARE IN A PHARMACEUTICALLY ACCEPTABLE FORMULATION THAT DOES NOT CONTAIN CREMOPHOR
U-634 METHOD FOR DELIVERY OF A BIOLOGIC (INCLUDING ANTINEOPLASTIC AGENTS) BY ADMINISTERING TO A PATIENT AN EFFECTIVE AMOUNT OF A BIOLOGIC AS A SOLID OR LIQUID WITH A POLYMERIC BIOCOMPATIBLE MATERIAL
U-635 TREATMENT OF GERD, MAINTENANCE OF HEALING OF EROSIIVE ESOPHAGITIS AND RISK REDUCTION OF NSAID ASSOCIATED GASTRIC ULCERS
U-636 TREATMENT OR PREVENTION OF BRONCHOSPASM OR ASTHMATIC SYMPTOMS
U-637 TREATMENT OF DIABETES WITH AN AMYLIN AGONIST
U-638 TREATMENT OF DIABETES WITH AN AMYLIN AGONIST, INCLUDING WITH INSULIN
U-639 TREATMENT OF A MAMMAL HAVING A NEED OF OR REDUCED ABILITY TO PRODUCE INSULIN WITH AN INSULIN AND AN AMYLIN SUCH AS PRAMLINTIDE
U-640 USE OF AN AMYLIN AGONIST TO REDUCE GASTRIC MOTILITY AND TREAT POST PRANDIAL HYPERGLYCEMIA
U-641 USE OF AN AMYLIN AGONIST HAVING SPECIFIED BINDING ACTIVITY TO REDUCE GASTRIC MOTILITY, INCLUDING USE THROUGH PARENTERAL ADMINISTRATION
U-642 TREATMENT AND PREVENTION OF OSTEOPOROSIS
U-643 THE SHORT TERM TREATMENT (UP TO 10 DAYS) IN PTS HAVING GASTROESOPHAGEAL REFLUX DISEASE (GERD) AS AN ALTERNATIVE TO ORAL THERAPY IN PTS WHEN THERAPY WITH NEXIUM CAPSULES IS NOT POSSIBLE OR APPROPRIATE
U-644 TREATMENT OF SEASONAL ALLERGIC RHINITIS
U-645 TREATMENT OF ASTHMA
U-646 METHOD OF TREATING OTITIS
U-647 TREATMENT OF OSTEOPOROSIS IN POST MENOPAUSAL WOMEN AND/OR THE TREATMENT TO INCREASE BONE MASS IN MEN WITH OSTEOPOROSIS
U-648 THE TREATMENT OF OSTEOPOROSIS IN POSTMENOPAUSAL WOMEN AND/OR THE TREATMENT TO INCREASE BONE MASS IN MEN
U-649 A METHOD FOR TREATING A TUMOR DISEASE
U-650 TREATMENT OF ESOPHAGEAL CANDIDIASIS AND PROPHYLAXIS OF CANDIDA INFECTIONS IN HSCT PATIENTS
U-651 TREATMENT OF ACUTE PROMYELOCYTIC LEUKEMIA (APL)
U-652 TREATMENT OF CARDIAC ARRHYTHMIA
U-653 STIMULATING INSULIN RELEASE BY ADMINISTERING EXENATIDE
U-654 LOWERING PLASMA GLUCAGON IN A SUBJECT IN NEED THEREOF, INCLUDING ONE WITH TYPE 2 DIABETES, BY ADMINISTERING AN EXENDIN OR ANALOG, SUCH AS EXENDIN-4
U-655 TREATMENT OF MILD TO MODERATE ACTIVE CROHN'S DISEASE INVOLVING THE ILEUM AND/OR THE ASCENDING COLON AND THE MAINTENANCE OF CLINICAL REMISSION OF MILD TO MODERATE CROHN'S DISEASE INVOLVING THE ILEUM AND/OR ASCENDING COLON FOR UP TO 3 MONTHS
U-656 REDUCING GASTRIC MOTILITY OR DELAYING GASTRIC EMPTYING BY ADMINISTERING AN EXENDIN, SUCH AS EXENDIN-4
U-657 PREVENTION OF OSTEOPOROSIS IN POSTMENOPAUSAL WOMEN
U-658 TREATMENT OF ADVANCED HORMONE-DEPENDENT BREAST CANCER
U-659 TREATMENT OF LOCALLY ADVANCED OR METASTATIC NON SMALL-CELL LUNG CANCER (NSCLC) AFTER FAILURE OF AT LEAST ONE PRIOR CHEMOTHERAPY REGIMEN

PATENT AND EXCLUSIVITY TERMS

ADB 33 of 35

PATENT USE

- U-660 TREATMENT OF HYPERTENSION AND TREATMENT OF HEART FAILURE
- U-661 TREATMENT OF SEIZURE DISORDER
- U-662 TREATMENT OF OSTEOPOROSIS IN POSTMENOPAUSAL WOMEN
- U-663 THE TREATMENT OF UNCOMPLICATED URINARY TRACT INFECTIONS
- U-664 TREATMENT OF CONDITIONS FOR WHICH AN ALDOSTERONE RECEPTOR BLOCKER IS INDICATED, SUCH AS HYPERTENSION, HEART FAILURE, AND POST-MYOCARDIAL INFARCTION
- U-665 METHOD OF USING THE DRUG SUBSTANCE/DRUG PRODUCT FOR ULTRASOUND IMAGING
- U-666 METHOD OF TREATING ADHD
- U-667 MANAGEMENT OF INCONTINENCE; METHOD FOR TREATING INCONTINENCE
- U-668 LEVEMIR IS A LONG-ACTING BASAL INSULIN ANALOG THAT IS INDICATED IN THE TREATMENT OF PATIENTS WITH DIABETES MELLITUS
- U-669 INDICATION OF TYPE II DIABETES
- U-670 TREATMENT OF HIV-1 INFECTION BY THE CO-ADMINISTRATION OF TIPRANAVIR AND RITONAVIR.
- U-671 PREVENTION AND TREATMENT OF SECONDARY HYPERPARATHYROIDISM ASSOCIATED WITH CHRONIC KIDNEY DISEASE (CKD) STAGE 3 AND 4
- U-672 TREATMENT OF INFLAMMATION OR AN INFLAMMATION-ASSOCIATED DISORDER
- U-673 METHOD OF TREATMENT WITH ONCE-DAILY DOSES OF 625MG/5ML
- U-674 METHOD OF TREATING INSOMNIA CHARACTERIZED BY DIFFICULTY WITH SLEEP ONSET
- U-675 PROPHYLAXIS AND CHRONIC TREATMENT OF ASTHMA; RELIEF OF SYMPTOMS OF ALLERGIC RHINITIS
- U-676 METHOD OF TREATING ATTENTION DEFICIT DISORDER USING ORAL ADMINISTRATION OF A BI-MODAL OR PULSATILE RELEASE COMPOSITION
- U-677 A METHOD OF TREATING DISEASE AMENABLE TO TREATMENT WITH A PHENIDATE DRUG BY ONCE DAILY ORAL ADMINISTRATION OF AN EXTENDED RELEASE DOSAGE FORM
- U-678 METHOD OF TREATING ATTENTION DEFICIT DISORDER AND/OR ATTENTION DEFICIT HYPERACTIVITY DISORDER
- U-679 ADJUNCT TO DIET AND EXERCISE TO IMPROVE GLYCEMIC CONTROL IN PATIENTS WITH TYPE 2 DIABETES WHO ARE ALREADY TREATED WITH A PIOGLITAZONE AND METFORMIN
- U-680 A METHOD OF TREATING DYSLIPIDEMIA AND DYSLIPOPROTEINEMIA USING A DOSAGE FORM THAT CAN PROVIDE AN EFFECTIVE AMOUNT OF FENOFIBRATE TO A PATIENT IN A FASTED STATE WHICH IS AT LEAST 90% OF THE AUC AMOUNT PROVIDED BY THE DOSAGE FORM
- U-681 TREATMENT OF PRIMARY IGF-1 DEFICIENCY
- U-682 NON-BENZODIAZEPINE HYPNOTIC AGENT INDICATED FOR TREATMENT OF INSOMNIA, CHARACTERIZED BY DIFFICULTIES WITH SLEEP ONSET AND/OR SLEEP MAINTENANCE
- U-683 PREVENTION OR TREATMENT OF ISCHEMIC HEART DISEASE
- U-684 TREATMENT OF UNCOMPLICATED SKIN MANIFESTATIONS OF CHRONIC IDIOPATHIC URTICARIA IN ADULTS AND CHILDREN 6 YEARS OF AGE AND OLDER
- U-685 EXPECTORANT AND COUGH SUPPRESSANT
- U-686 EXPECTORANT AND NASAL DECONGESTANT
- U-687 REDUCING FOOD INTAKE IN A SUBJECT WITH TYPE 2 DIABETES BY ADMINISTERING AN EXENDIN, SUCH AS EXENDIN-4
- U-688 TREATMENT OF HIV-INFECTION IN COMBINATION WITH OTHER ANTIRETROVIRAL AGENTS
- U-689 TREATMENT OF PATIENTS WITH T-CELL ACUTE LYMPHOBLASTIC LEUKEMIA WHOSE DISEASE HAS NOT RESPONDED TO OR HAS RELAPSED FOLLOWING TREATMENT WITH AT LEAST TWO CHEMOTHERAPY REGIMENS
- U-690 TO IMPROVE GLYCEMIC CONTROL IN PATIENTS WITH TYPE 2 DIABETES MELLITUS
- U-691 USE AS A MONOTHERAPY, IN COMBINATION WITH A SULFONYLUREA, METFORMIN OR INSULIN OR IN COMBINATION WITH A SULFONYLUREA PLUS METFORMIN TO IMPROVE GLYCEMIC CONTROL IN PATIENTS WITH TYPE 2 DIABETES MELLITUS
- U-692 USE OF VALSARTAN TO REDUCE CARDIOVASCULAR MORTALITY IN CLINICALLY STABLE PATIENTS WITH LEFT VENTRICULAR FAILURE OR LEFT VENTRICULAR DYSFUNCTION FOLLOWING MYOCARDIAL INFARCTION
- U-693 THE RECOMMENDED INITIAL DOSE OF EQUETRO IS 400MG/DAY GIVEN IN DIVIDED DOSES, TWICE DAILY. THE DOSE SHOULD BE ADJUSTED IN 200MG DAILY INCREMENTS TO ACHIEVE OPTIMAL CLINICAL RESPONSE.
- U-694 LENALIDOMIDE IS AN ANALOGUE OF THALIDOMIDE. THALIDOMIDE IS A KNOWN HUMAN TERATOGEN THAT CAUSES SEVERE LIFE-THREATENING HUMAN BIRTH DEFECTS. IF LENALIDOMIDE IS TAKEN DURING PREGNANCY, IT MAY CAUSE BIRTH DEFECTS OR DEATH TO AN UNBORN BABY.
- U-695 TREATMENT OF PATIENTS WITH T-CELL ACUTE LYMPHOBLASTIC LEUKEMIA AND T-CELL LYMPHOBLASTIC LYMPHOMA WHOSE DISEASE HAS NOT RESPONDED TO OR HAS RELAPSED FOLLOWING TREATMENT WITH AT LEAST TWO CHEMOTHERAPY REGIMENS
- U-696 TREATMENT OF PATIENTS WITH T-CELL LYMPHOBLASTIC LYMPHOMA WHOSE DISEASE HAS NOT RESPONDED TO OR HAS RELAPSED FOLLOWING TREATMENT WITH AT LEAST TWO CHEMOTHERAPY REGIMENS
- U-697 A METHOD OF USING RINFABATE RECOMBINANT (RHIGFBP-3) WITH MECASERMIN RECOMBINANT (RHIGF-1) TO PROMOTE LINEAR GROWTH IN THE TRATMENT OF PRIMARY IGF-1 DEFICIENCY
- U-698 METHOD OF USING ANTAGONIST OF ARGININE VASOPRESSIN (AVA) V1A AND V2 RECEPTORS FOR INTRAVENOUS TREATMENT OF PATEINTS WITH EUVOLEMIC HYPONATREMIA
- U-699 NASAL TREATMENT OF SEASONAL AND PERENNIAL ALLERGIC RHINITIS SYMPTOMS
- U-700 TREATMENT AND PREVENTION OF OSTEOPOROSIS IN POSTMENOPAUSAL WOMEN

PATENT AND EXCLUSIVITY TERMS

ADB 34 of 35

PATENT USE

U-701 TREATMENT OF HYPERCHOLESTEROLEMIA AND/OR HYPERTRIGLYCERIDEMIA
U-702 TOPICAL AEROSOL HAIR REGROWTH TREATMENT
U-703 TREATMENT OF PROTEIN KINASE RELATED DISORDERS, SUCH AS GASTROINTESTINAL STROMAL TUMOR AND RENAL CELL CARCINOMA WITH SUNITINIB
U-704 METHOD OF ADMINISTERING INSULIN VIA INHALATION
U-705 TREATING CHRONIC ANGINA BY ADMINISTERING AN EXTENDED RELEASE FORM OF RANOLAZINE
U-706 TREATMENT OF BENIGN PROSTATIC HYPERPLASIA
U-707 ALLERGIC RHINITIS
U-708 TREATMENT OF CHRONIC NON-INFECTIOUS UVEITIS AFFECTING THE POSTERIOR SEGMENT OF THE EYE
U-709 METHOD OF COMBATING BACTERIA IN A PATIENT
U-710 A METHOD OF TREATING RESPIRATORY DISORDERS, E.G., ASTHMA, WHICH COMPRISES ADMINISTRATION BY INHALATION OF AN EFFECTIVE AMOUNT OF A PHARMACEUTICAL FORMULATION AS CLAIMED IN US PATENT NO. 5658549
U-711 ACUTE AND LONGER-TERM TREATMENT OF MAJOR DEPRESSIVE DISORDER
U-712 A METHOD OF USING A NICOTINIC ACID FORMULATION TO REDUCE ELEVATED TC, LDL-C AND TG LEVELS, AND RAISE HDL-C LEVELS IN PATIENTS WITH HYPERLIPIDEMIA
U-713 TREATMENT OF MILD TO MODERATE DEMENTIA OF THE ALZHEIMER'S TYPE
U-714 TOPICAL TREATMENT OF INTERDIGITAL TINEA PEDIS AND TINEA CORPORIS DUE TO TRICHOPHYTON RUBRUM, TRICHOPHYTON MENTAGROPHYTES OR EPIDERMOPHYTON FLOCCOSUM
U-715 FOR CLEANSING THE BOWEL IN PREPARATION FOR COLONOSCOPY, IN ADULTS 18 YEARS OF AGE OR OLDER
U-716 THE TREATMENT OR PREVENTION OF BRONCHOSPASM IN ADULTS AND CHILDREN 4 YEARS OF AGE AND OLDER WITH REVERSIBLE OBSTRUCTIVE AIRWAYS DISEASE AND THE PREVENTION OF EXERCISED-INDUCED BRONCHOSPASM IN PATIENTS 4 YEARS OF AGE AND OLDER
U-717 METHOD OF RELIEVING OR PREVENTING CONSTIPATION IN A HUMAN CONSTIPATED PATIENT
U-718 TREATMENT OF FUNGAL INFECTIONS
U-719 TREATMENT OF PSYCHOSIS
U-720 TREATMENT OF NEUROLEPTIC DISEASES
U-721 TREATMENT OF INFLUENZA
U-722 PROPHYLAXIS OF INFLUENZA
U-723 PROPHYLACTIC TREATMENT OF MIGRAINE
U-724 METHOD OF TREATING SEIZURES
U-725 ALLERGIC RHINITIS AND URTICARIA
U-726 ALLERGIC RHINITIS
U-727 FOR THE TREATMENT OF ATTENTION DEFICIT HYPERACTIVITY DISORDER (ADHD)
U-728 METHOD FOR TREATING BACTERIAL INFECTION
U-729 TREATMENT OF GASTROESOPHAGEAL REFLUX DISEASE (GERD), RISK-REDUCTION OF NSAID-ASSOCIATED GASTRIC ULCER, H. PYLORI ERADICATION TO REDUCE THE RISK OF DUODENAL ULCER RECURRENCE
U-730 USE AS A NASAL SPRAY FOR TREATMENT OF THE SYMPTOMS OF SEASONAL ALLERGIC RHINITIS AND VASOMOTOR RHINITIS
U-731 USE IN COMBINATION WITH DEXAMETHASONE IS INDICATED FOR THE TREATMENT OF PATIENTS WITH NEWLY DIAGNOSED MULTIPLE MYELOMA
U-732 ACUTE TREATMENT OF THE CUTANEOUS MANIFESTATIONS OF MODERATE TO SEVERE ERYTHEMA NODOSUM LEPROSUM (ENL)
U-733 MAINTENANCE THERAPY FOR PREVENTION AND SUPPRESSION OF THE CUTANEOUS MANIFESTATIONS OF ENL RECURRENCE
U-734 FIRST LINE THERAPY FOR TYPE 2 DIABETES MELLITUS
U-735 METHOD OF TREATING CHRONIC IRON OVERLOAD
U-736 METHOD FOR IONTOPHORETIC TRANSDERMAL DELIVERY OF FENTANYL HYDROCHLORIDE
U-737 DISINFECTION OF PATIENT SKIN PRIOR TO AN INVASIVE PROCEDURE
U-738 INDICATED FOR THE LONG-TERM, TWICE-DAILY MAINTENANCE TREATMENT OF ASTHMA IN PATIENTS 12 YEARS OF AGE OR OLDER
U-739 METHOD FOR TREATING CONSTIPATION BY OPENING CIC CHANNELS IN A MAMALIAN SUBJECT
U-740 FOR THE TREATMENT OF PATIENTS WITH PRIMARY BILIARY CIRRHOSIS
U-741 COMBINATION THERAPY WITH CISPLATIN FOR THE TREATMENT OF LATE STAGE CERVICAL CANCER
U-742 TWICE DAILY TOPICAL TREATMENT OF MODERATE TO SEVERE PLAQUE PSORIASIS.
U-743 ONCE A DAY TOPICAL TREATMENT OF THE INFLAMMATORY LESIONS OF ROSACEA
U-744 TREATMENT OF HIV INFECTION IN ANTIRETROVIRAL TREATMENT-EXPERIENCED ADULT PATIENTS
U-745 TREATMENT OR PREVENTION OF EMESIS
U-746 PREVENTION OR TREATMENT OF NAUSEA OR EMESIS INDUCED BY A CANCER CHEMOTHERAPEUTIC AGENT
U-747 PREVENTION OR TREATMENT OF POST-OPERATIVE NAUSEA AND VOMITING
U-748 A METHOD FOR THE TREATMENT OF A PROTEIN TYROSINE KINASE-ASSOCIATED DISORDER
U-749 METHOD OF CONTRACEPTION
U-750 TREATMENT OF HIV-1 INFECTION IN ADULTS

PATENT AND EXCLUSIVITY TERMS

ADB 35 of 35

PATENT USE

U-751 ONCE DAILY DOSING OF BUDESONIDE VIA NEBULIZER FOR THE TREATMENT OF ASTHMA
U-752 SUNSCREEN
U-753 AS AN ADJUNCT TO DIET AND EXERCISE TO IMPROVE GLYCEMIC CONTROL IN PATIENTS WITH TYPE 2 DIABETES
U-754 USE FOR THE LONG-TERM MAINTENANCE TREATMENT OF ASTHMA
U-755 TREATMENT OF ANOREXIA, CACHEXIA, OR AN UNEXPLAINED, SIGNIFICANT WEIGHT LOSS IN PATIENTS WITH A DIAGNOSIS OF ACQUIRED IMMUNODEFICIENCY SYNDROME (AIDS)
U-756 ADDITION OF ONCE-WEEKLY DOSING FOR THE TREATMENT TO INCREASE BONE MASS IN MEN WITH OSTEOPOROSIS
U-757 USE AS A BILE ACID SEQUESTRANT FOR LOWERING CHOLESTEROL
U-758 TREATMENT OF SYMPTOMS OF PREMENSTRUAL DYSPHORIC DISORDER
U-759 METHOD OF USE OF ADMINISTERING LEVOTHYROXINE
U-760 PROPHYLAXIS OF INVASIVE ASPERGILLUS AND CANDIDA INFECTIONS AND TREATMENT OF OROPHARYNGEAL CANDIDIASIS
U-761 TREATMENT OF SCHIZOPHRENIA INCLUDING MAINTAINING STABILITY IN PATIENTS WITH SCHIZOPHRENIA
U-762 TREATMENT OF CHRONIC OBSTRUCTIVE PULMONARY DISEASE
U-763 ADMINISTRATION OF ARIPIPRAZOLE BY INJECTION
U-764 TREATMENT OF SCHIZOPHRENIA
U-765 METHOD OF TREATING ALLERGIC CONJUNCTIVITIS
U-766 TREATMENT OF SEIZURES
U-767 MANAGEMENT OF BREAKTHROUGH PAIN IN PATIENTS WITH CANCER
U-768 A METHOD OF REDUCING THE CAPACITY OF EXTENDED RELEASE NICOTINIC ACID TO PROVOKE A FLUSHING REACTION BY PRETREATING AN INDIVIDUAL WITH A FLUSH INHIBITING AGENT PRIOR TO THE ADMINISTRATION OF THE EXTENDED RELEASE NICOTINIC ACID
U-769 REVLIMID (LENALIDOMIDE) IN COMBINATION WITH DEXAMETHASONE IS INDICATED FOR THE TREATMENT OF MULTIPLE MYELOMA PATIENTS WHO HAVE RECEIVED AT LEAST ONE PRIOR THERAPY
U-770 LONG-TERM TREATMENT OF PATHOLOGICAL HYPERSECRETORY CONDITIONS
U-771 METHOD FOR THE TREATMENT OF DIABETES MELLITUS, SUCH AS TYPE 1 DIABETES MELLITUS OR TYPE 2 DIABETES MELLITUS, IN A HUMAN PATIENT
U-772 RELIEF OF SYMPTOMS ASSOCIATED WITH SEASONAL ALLERGIC RHINITIS IN CHILDREN 2 TO 11 YEARS AND FOR THE RELIEF OF SYMPTOMS ASSOCIATED WITH UNCOMPLICATED SKIN MANIFESTATIONS OF CHRONIC IDIOPATHIC URTICARIA IN CHILDREN 6 MONTHS TO 11 YEARS
U-773 PATHOLOGICAL HYPERSECRETORY CONDITIONS
U-774 METHOD OF TREATING TYPE 2 DIABETES MELLITUS BY ADMINISTERING A DIPEPTIDYL PEPTIDASE-IV INHIBITOR
U-775 METHOD OF TREATING TYPE 2 DIABETES MELLITUS BY ADMINISTERING A DIPEPTIDYL PEPTIDASE-IV INHIBITOR IN COMBINATION WITH METFORMIN OR A PPAR-GAMMA AGONIST
U-776 TREATMENT OF CUTANEOUS MANIFESTATION IN PATIENTS WITH CUTANEOUS T-CELL LYMPHOMA (CTCL) WHO HAVE PROGRESSIVE, PERSISTENT OR RECURRENT DISEASE ON OR FOLLOWING TWO SYSTEMIC THERAPIES.
U-777 DECREASING MORTALITY CAUSED BY CONGESTIVE HEART FAILURE
U-778 REDUCTION OF ELEVATED INTRAOCULAR PRESSURE IN PATIENTS WITH OPEN ANGLE GLAUCOMA OR OCULAR HYPERTENSION
U-779 A METHOD FOR TREATMENT OF A CANCER, WHEREIN THE CANCER IS CHRONIC MYELOGENOUS LEUKEMIA
U-780 A METHOD FOR THE TREATMENT OF CANCER
U-781 FOR TREATMENT OF ADULT PATIENTS WITH TYPE 2 DIABETES MELLITUS WHO ARE NAIVE TO PHARMACOLOGIC THERAPY
U-782 TREATMENT OF CHRONIC HEPATITIS B IN ADULT PATIENTS WITH EVIDENCE OF VIRAL REPLICATION AND EITHER EVIDENCE OF PERSISTENT ELEVATIONS IN SERUM AMINOTRANSFERASES (ALT OR AST) OR HISTOLOGICALLY ACTIVE DISEASE
U-783 DESONATE GEL IS INDICATED FOR THE TREATMENT OF MILD TO MODERATE ATOPIC DERMATITIS IN PATIENTS 3 MONTHS OF AGE AND OLDER
U-784 TREATMENT OF MODERATE TO SEVERE PRIMARY RESTLESS LEGS SYNDROME (RLS)
U-785 USE AS REPLACEMENT SOLUTION, HEMOFILTRATION SOLUTION OR HEMODIAFILTRATION SOLUTION IN CONTINUOUS RENAL REPLACEMENT THERAPY
U-786 PRODUCT IS APPROVED FOR THE TOPICAL TREATMENT OF TINEA PEDIS
U-787 MAINTENANCE TREATMENT OF ASTHMA AS PROPHYLACTIC THERAPY IN ADULT AND PEDIATRIC PATIENTS SIX YEARS OF AGE OR OLDER, INCLUDING PATIENTS REQUIRING ORAL CORTICOSTEROID THERAPY FOR ASTHMA