

APPROVED DRUG PRODUCTS

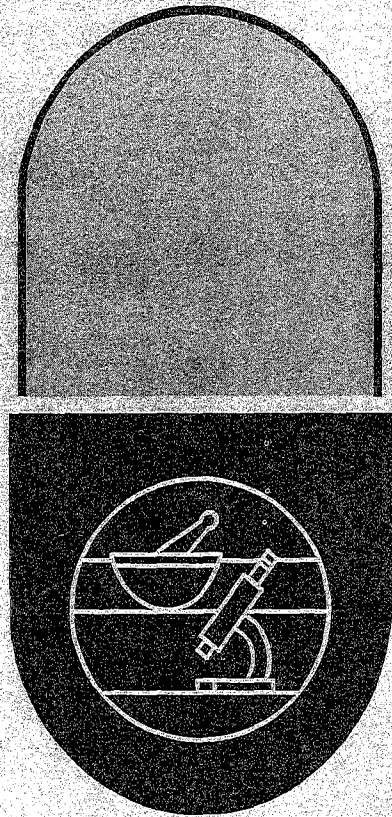
With Therapeutic Equivalence Evaluations



The "Orange Book"

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**CUMULATIVE
SUPPLEMENT 12
JAN'96-DEC'96**



APPROVED DRUG PRODUCTS

**WITH
THERAPEUTIC EQUIVALENCE EVALUATIONS**

16TH EDITION

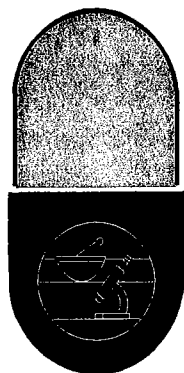
U.S. DEPARTMENT OF HEALTH AND HUMAN SERVICES

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APPROVED DRUG PRODUCTS

WITH
THERAPEUTIC EQUIVALENCE EVALUATIONS

**17TH EDITION
1997**

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**APPROVED DRUG PRODUCTS
with
THERAPEUTIC EQUIVALENCE EVALUATIONS**

16TH EDITION

Cumulative Supplement 12

DECEMBER 1996

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**APPROVED DRUG PRODUCTS
with
THERAPEUTIC EQUIVALENCE EVALUATIONS**

16TH EDITION

**CUMULATIVE SUPPLEMENT 12
DECEMBER 1996**

1.0 INTRODUCTION

1.1 HOW TO USE THE CUMULATIVE SUPPLEMENT

This Cumulative Supplement is one of a series of monthly updates to the Approved Drug Products with Therapeutic Equivalence Evaluations, 16th Edition (the List). The List is composed of four parts: approved prescription drug products with therapeutic equivalence evaluations, over-the-counter (OTC) drug products that require approved applications as a condition of marketing, drug products with approval under Section 505 of the Act administered by the Center for Biologics Evaluation and Research and products that have never been marketed, have been discontinued from marketing or that have had their approvals withdrawn for other than safety or efficacy reasons.

The Cumulative Supplement provides, among other things, information on newly approved drugs and, if necessary, revised therapeutic equivalence evaluations and updated patent and exclusivity data. The Addendum contains appropriate drug patent and exclusivity information required of the Agency by the "Drug Price Competition and Patent Term Restoration Act of 1984" for the Prescription, OTC, and Drug Products with Approval under Section 505 of the Act Administered by the Center for Biologics Evaluation and Research Lists.

The Patent and Exclusivity Lists are arranged in alphabetical order by active ingredient name. For those products with multiple active ingredients, only the first active ingredient (in alphabetical sort) will appear. In addition, the trade name will be displayed to the right of the active ingredient name for each product. Also shown is the application number and product number (FDA's internal file number) for reference purposes. All patents with their expiration dates are displayed for each application number. Use patents are indicated with the symbol "U" followed by a number representing a specific use. Exclusivity information for a specific drug is indicated by an abbreviation followed by the date upon which the exclusivity expires. Refer to the Exclusivity Terms section in the Patent and Exclusivity Information Addendum for an explanation of all codes and abbreviations.

Because all parts of the publication are subject to changes, additions, or deletions, the List must be used in conjunction with the most current Cumulative Supplement. Users may wish to place an asterisk (*) to the left of the ingredient(s) in the List to indicate that changes to that entry appear in the Cumulative Supplement.

Drug product information is provided in each Cumulative Supplement for completeness to assist in locating the proper place in the List for the revision. [Strength(s) which already exist in the List will not be repeated for context.]

The presence of any therapeutic equivalence code indicates that the drug product is multisource; the deletion of a therapeutic equivalence code indicates that the drug product has become single source. (An infrequent exception exists when a therapeutic equivalence code is revised. In that case the deletion of the therapeutic equivalence code is followed immediately by the addition of the revised one.)

New additions new to the Prescription Drug Product List, OTC Drug Product List, and the Patent and Exclusivity Data are indicated by the symbol >ADD> to the left of the line on which new information exists. The >ADD> symbol is then dropped in subsequent Cumulative Supplements for that item.

New deletions to the Prescription Drug Product List, OTC Drug Product List, and the Patent and Exclusivity Data are indicated by the symbol >DLT> (DELETE) to the left of the line containing shaded print. The >DLT> symbol is dropped in subsequent Cumulative Supplements for that item. The shaded print remains in the Prescription Drug Product List and OTC Drug Product Lists in all Cumulative Supplements for this edition. However, the shaded print in the Patent and Exclusivity Data is dropped in subsequent Cumulative Supplements.

Products that have never been marketed, have been discontinued from marketing or that have had their approvals withdrawn for other than safety or efficacy reasons, will be flagged in this Cumulative Supplement with the "@" symbol to designate their non-marketed status. All products having a "@" symbol in the 12th Cumulative Supplement of the 16th Edition List will then be added to the "Discontinued Drug Product List" appearing in the 17th Edition.

1.2 PRODUCTS REQUIRING REVISED LABELING FOR FULL APPROVAL

Drug products in this category (1) initially received approval only on the basis of safety before effectiveness studies were required, or (2) were conditionally approved under the temporary exemption that allowed these products to be marketed while effectiveness studies were being conducted. Listed below are those drugs which are now required to revise their labeling and provide additional information necessary for full approval

on the basis of requirements listed in the Federal Register. As approval is granted by the Agency for a specific product, based on additional information submitted by the applicant, the product will be included in the appropriate Drug Product List.

<u>Products</u>	<u>Federal Register Reference</u>
Nitroglycerin (capsule, controlled release;oral)	SEP 07, 1984 (49 FR 35428)
Nitroglycerin (film, extended release;transdermal*)	JUL 15, 1993 (58 FR 38129)
Nitroglycerin (tablet, controlled release;oral)	SEP 07, 1984 (49 FR 35428)
Nitroglycerin (tablet, controlled release;buccal)	JUL 05, 1985 (50 FR 27688)

*The Federal Register of July 15, 1993 (58 FR 38129) announced that the FDA was revoking the temporary exemption for nitroglycerin in a transdermal delivery system. Marketing of a drug product that is the subject of a conditionally approved ANDA may continue by meeting the requirements listed in the Federal Register. Firms wishing to submit a new ANDA before a drug product is approved (NDA or ANDA) and appears in the List should submit a 505(b)(2) application following the directions contained in the Federal Register. Nitro-Dur has been selected as the reference listed drug. The preamble to the final rule (57 FR 17958) states if there are multiple NDA's, the reference listed drug generally will be the market leader. This is the basis upon which Nitro-Dur was selected. In addition, the preamble states that, in multiple NDA situations, 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. This is the case with Summit's Transderm-Nitro. The Office of Generic Drugs (OGD) has been requested to have a second listed drug as provided for in the Final Rule. OGD has granted this request. Therefore, at the time that Schering's and Summit's supplements are fully approved and their products are entered into the List, we will have two reference listed drugs for the nitroglycerin transdermal systems. Firms may, therefore, elect to conduct bioequivalence studies against either of these products. It is conceivable that a non-referenced listed drug may be fully approved and appear in the List; in this case, the Agency's referenced listed drug will not change and a 505(b)(2) application will be appropriate until the reference listed drugs are fully approved. Once they are fully approved, a 505(j) application will be the appropriate mechanism for an ANDA submission.

1.3 CHANGE OF A THERAPEUTIC EQUIVALENT CODE FOR A DRUG ENTITY

Propantheline Bromide

In Cumulative Supplement 1 of the *Approved Drug Products with Therapeutic Equivalence Evaluations*, 16th Edition, (Orange Book), the Agency proposed to change the therapeutic equivalence code for propantheline bromide oral tablets from a drug product not presenting a bioequivalence problem (**AA**) to a drug product with a potential bioequivalence problem (**BP**).

The Agency solicited comments from interested persons to be received no later than 60 days from the first day of the month following the publication of Cumulative Supplement 1. The proposal did not elicit any comments from the readers. In addition, the two firms who hold an active ANDA and are marketing the drug product were contacted to inform them that the codes for their propantheline drug products were going to be changed. Since there were no comments submitted by the readers or by the two firms who hold an active ANDA, the therapeutic equivalence code for propantheline bromide tablets will be changed to one reflecting a potential bioequivalence problem. Therefore, all oral propantheline bromide tablets will be changed in this month's Cumulative Supplement from **(AA)** to **(BP)** to reflect it has a potential for a bioequivalence problem.

An acceptable *in vivo* bioequivalence study, among other information, will be required to change the code from **(BP)** to **(AB)** for an already approved ANDA listed in the Orange Book. Any ANDA submission must contain an acceptable *in vivo* bioequivalence study for filing purposes.

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 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 multiple NDAs 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 an NDA listed drug that is not designated as the reference listed 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 NDA 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* of the *Introduction* to the *Approved Drug Products with Therapeutic Equivalence Evaluations* publication explains the coding system for multisource drug products listed under the same heading with two reference listed drugs.

The concept of having only one reference listed drug was intended to apply to drug products in which bioequivalence is demonstrated through *in vivo* methodology. It was not intended to apply to two NDA drug products in which the *in vivo* determination of bioequivalence is self evident and a waiver of *in vivo* bioequivalence is granted by the agency. These types of drug products are assigned therapeutic equivalence codes, e.g., of **AN**, **AT**, **AA**. Therefore, drug products that do not represent a bioequivalence problem with two or more NDAs will have the reference listed drug designation assigned to each NDA.

The reference listed drug is identified by the symbol "+" in the Prescription Drug Product List. These identified reference listed drugs represent the best judgement of the Division of Bioequivalence at this time. The prescription Drug Product List identifies 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 COURT ORDER REGARDING ABBOTT U.S. PATENT NO. 4112097, (TERAZOSIN HCL)

On April 9, 1996, the United States District Court for the Northern District of Illinois (Eastern Division) issued an order in the case of Abbott Labs v. Geneva Pharmaceuticals, Inc., directing Abbott to remove U.S. Patent No. 4112097 from the Orange Book. To comply with that order, Abbott has requested that FDA remove patent 4112097 from the Orange Book. The FDA complied with this request in the March 1996 cumulative supplement. On April 9, 1996, Abbott appealed the district court's decision to the U.S. Court of Appeals for the Federal Circuit.

1.6 COURT ORDER AFFECTING URUGUAY ROUND AGREEMENTS ACT-EXTENDED PATENTS

As a result of the April 4, 1996, decision of the United States Court of Appeals for the Federal Circuit in Merck, et al. v. Kessler, patent expiration dates for certain patents subject to patent term extensions under the Uruguay Round Agreements Act and to the patent term extension provisions at 35 U.S.C. § 156 may be changed. FDA is publishing a notice in the *Federal Register* advising NDA and NADA applicants that patent expiration dates changed by the Merck decision must be submitted within 60 days. Because there may be changes in listed patents as a result of the Merck decision, users of this publication should consult the most recent supplement, and are encouraged to confirm that patent information upon which they intend to rely is current. (See the *Patent and Exclusivity Addendum* to the *Approved Drug Products with Therapeutic Equivalence Evaluations*, 16th Edition that explains the background information on this court decision).

1.7 APPLICANT NAME CHANGES

It is not practical to identify in the Cumulative Supplement each and every product involved when an applicant transfers its entire line of approved drug products to another applicant, or when an applicant changes its name. Therefore, the cumulation of these transfers and name changes will be identified in this section only. Where only partial lines of approved products are transferred between applicants, each approved product involved will appear as an applicant name change entry in the Cumulative Supplement.

It is also not practical to identify each and every product involved when an applicant name is changed to meet internal publication standards (e.g., MSD or Zenith [Former Abbreviated Names] are changed, respectively, to Merck Sharp Dohme or Zenith Labs

[New Abbreviated Names]]. When this occurs, each product involved (either currently in the Cumulative Supplement or in the following year's edition) will automatically reflect the new abbreviated name. Consequently, it will not appear as an applicant name change entry in the Cumulative Supplement nor will the cumulation of these name changes appear in this section. However, when the applicant name change is one which may not be easily recognized or located in the listing (e.g., White Towne Paulsen [Former Abbreviated Name] is changed to Whiteworth Towne [New Abbreviated Name], the name change will appear in this section and will be identified with an asterisk.

APPLICANT NAME CHANGES

FORMER APPLICANT NAME
(FORMER ABBREVIATED NAME)

NEW APPLICANT NAME
(NEW ABBREVIATED NAME)

1ST TEXAS PHARMACEUTICALS INC
SUB SCHERER LABORATORIES
(1ST TX)

SCHERER LABORATORIES, INC
(SCHERER)

BARRE NATIONAL INC
(BARRE)

ALPHARMA USPD INC
(ALPHARMA)

BEN VENUE
(BEN VENUE LABORATORIES INC)

BEDFORD
(BEDFORD LABORATORIES DIV
BEN VENUE LABORATORIES INC)

BIOCRAFT
(BIOCRAFT LABORATORIES INC)

TEVA
(TEVA PHARMACEUTICALS USA)

BOEHRINGER MANNHEIM PHARMACEUTICALS CORP
(BOEHRINGER MANNHEIM)

BOEHRINGER MANNHEIM CORPORATION
THERAPEUTICS DIVISION
(BOEHRINGER MANNHEIM)

CETUS BEN VENUE
(CETUS BEN VENUE THERAPEUTICS)

BEDFORD
(BEDFORD LABORATORIES DIV
BEN VENUE LABORATORIES INC)

DAVID BULL LABORATORIES PARTY LTD
(BULL D)

FH FAULDING AND CO LTD
(FAULDING)
THEN CHANGED TO
FAULDING PHARMACEUTICAL CO
(FAULDING)

HOECHST ROUSSEL PHARMACEUTICALS INC
(HOECHST ROUSSEL)

HOECHST MARION ROUSSEL INC
(HOECHST MARION RSSL)

KM LEE LABORATORIES INC
(KM LEE)

KM LEE LABORATORIES INC
(LEE KM)

APPLICANT NAME CHANGES

FORMER APPLICANT NAME
(FORMER ABBREVIATED NAME)

NEW APPLICANT NAME
(NEW ABBREVIATED NAME)

PHARMACIA INC
(PHARMACIA)

PHARMACIA AND UPJOHN CO
(PHARMACIA AND UPJOHN)

SCHWARZ PHARMA KREMERS
URBAN CO SUB SCHWARZ PHARMA AG
(SPKU)

SCHWARZ PHARMA INC
(SCHWARZ PHARMA)

UPJOHN CO
(UPJOHN)

PHARMACIA AND UPJOHN CO
(PHARMACIA AND UPJOHN)

UPJOHN MANUFACTURING CO
(UPJOHN)

PHARMACIA AND UPJOHN CARIBE INC
(PHARMACIA AND UPJOHN)

1.8 AVAILABILITY OF THE PUBLICATION AND UPDATING PROCEDURES

The *Approved Drug Products with Therapeutic Equivalence Evaluations* (Prescription Drug Products, OTC Drug Products and the Discontinued Drug Product Lists) is now available on diskette, on a quarterly basis, from the National Technical Information Service. The telephone number for the Subscription Department is (703) 487-6430. Written inquiries regarding this subscription may be forwarded to 5285 Port Royal Road Springfield, VA 22161.

The following *Approved Drug Products with Therapeutic Equivalence Evaluations* files are now available on Internet and are updated each October and April: Prescription Drug Product List; OTC Drug Products; Discontinued Drug Products; and Appendices. The update in October will include drug products that have been approved through August and the update in April will include drug products that have been approved through December.

These files may be accessed on the Internet's World Wide Web. FDA's Internet site replaces the Agency's electronic bulletin board and offers more information, in a more user-friendly form. To access the CDER Home Page, use this Uniform Resource Locator (URL): <http://www.fda.gov/cder>. You do not need an Internet connection to reach the FDA Home Page; you can use the free dial-up connection (800) 222-0185 for text based, non-graphical use only. For further assistance, please call (301) 443-4908.

The Prescription Drug Products and OTC Drug Product files will be available on a monthly basis in the near future.

1.9 REPORT OF COUNTS FOR THE PRESCRIPTION DRUG PRODUCT LIST

DESCRIPTION OF REPORT

This report provides summary counts derived from the product information in the Prescription Drug Product List and the current Cumulative Supplement. Products included in the counts are domestically marketed drug products approved for both safety and effectiveness under sections 505 and 507 of the Federal Food, Drug, and Cosmetic Act. Excluded are approved drug products marketed by distributors; those marketed solely abroad; and those now regarded as medical devices, biologics or foods.

The baseline column (Dec 1995) refers to the products in the Prescription Drug Product List. For each three-month period, a column of quarterly data is added which incorporates counts of product activity from the previous quarter(s) with those in the baseline count.

DEFINITIONS

Drug Product

For this report, a drug product is the representation in the Prescription Drug Product List of an active moiety (molecular entity and its salts, esters and derivatives) either as a single ingredient or as a combination product provided in a specific dosage form and strength for a given route of administration with approval for marketing by a firm under a particular generic or trade name.

New Molecular Entity

A new molecular entity is considered an active moiety that has not previously been approved (either as the parent compound or as a salt, ester or derivative of the parent compound) in the United States for use in a drug product either as a single ingredient or as part of a combination.

REPORT OF COUNTS FOR THE PRESCRIPTION DRUG PRODUCT LIST

COUNTS CUMULATIVE BY QUARTER

<u>CATEGORIES COUNTED</u>	<u>DEC 1995</u>	<u>JUN 1996</u>	<u>SEP 1996</u>	<u>DEC 1996</u>
DRUG PRODUCTS LISTED	9286	9384	9403	9392
SINGLE SOURCE	2217 (23.9%)	2323 (24.8%)	2331 (24.8%)	2383 (25.4%)
MULTISOURCE	7069 (76.1%)	7061 (75.2%)	7072 (75.2%)	7009 (74.6%)
THERAPEUTICALLY EQUIVALENT	6437 (69.3%)	6490 (69.2%)	6519 (69.3%)	6463 (68.8%)
NOT THERAPEUTICALLY EQUIVALENT	440 (4.7%)	468 (5.0%)	449 (4.8%)	442 (4.7%)
EXCEPTIONS ¹	192 (2.1%)	103 (1.0%)	104 (1.1%)	104 (1.1%)
NEW MOLECULAR ENTITIES APPROVED	--	15	14	19
NUMBER OF APPLICANTS	586	621	636	650

<u>CATEGORIES COUNTED</u>	<u>DEC 1996*</u>
DRUG PRODUCTS LISTED	9392
SINGLE SOURCE	2383 (25.4%)
MULTISOURCE	6905 (73.5%)
THERAPEUTICALLY EQUIVALENT	6463 (68.8%)
NOT THERAPEUTICALLY EQUIVALENT	442 (4.7%)
EXCEPTIONS ¹	104 (1.1%)
NEW MOLECULAR ENTITIES APPROVED	19
NUMBER OF APPLICANTS	650

¹Amino acid-containing products of varying composition (see Introduction, page xvi of the List).

*Exceptions were originally included in the total count of the Multisource Drug Products. Beginning with December 1996, exceptions will no longer be included in the Multisource Drug Products total count, but will be included in the total count of the Drug Products Listed.

PRESCRIPTION DRUG PRODUCT LIST
16TH EDITION
RX DRUG PRODUCT LIST / CUMULATIVE SUPPLEMENT NUMBER 12 / JAN'96 - DEC'96

1

ACETAMINOPHEN; BUTALBITAL

TABLET; ORAL			
<u>BUTALBITAL AND ACETAMINOPHEN</u>			
<u>AB</u>	<u>HALSEY</u>	<u>325MG;50MG</u>	<u>N89568 001</u>
			OCT 05, 1988
@		325MG;50MG	N89568 001
			OCT 05, 1988

ACETAMINOPHEN; BUTALBITAL; CAFFEINE

CAPSULE; ORAL			
ESGIC-PLUS			
+	MIKART	500MG;50MG;40MG	N40085 001
			MAR 28, 1996

TABLET; ORAL			
BUTALBITAL, ACETAMINOPHEN AND CAFFEINE			
	MIKART	500MG;50MG;40MG	N89451 001
			MAY 23, 1988
+		500MG;50MG;40MG	N89451 001
			MAY 23, 1988

ACETAMINOPHEN; CAFFEINE; DIHYDROCODEINE BITARTRATE

CAPSULE; ORAL			
DHC PLUS			
<u>AA</u>	<u>BURDUE FREDERICK</u>	<u>356.4MG;30MG;16MG</u>	<u>N88584 001</u>
			MAR 04, 1986
+		356.4MG;30MG;16MG	N88584 001
			MAR 04, 1986
<u>SYNALGOS-DC-A</u>			
<u>AA</u>	<u>* WYETH AYERST</u>	<u>356.4MG;30MG;16MG</u>	<u>N89166 001</u>
			MAY 14, 1986
@		356.4MG;30MG;16MG	N89166 001
			MAY 14, 1986

ACETAMINOPHEN; CODEINE PHOSPHATE

SOLUTION; ORAL			
<u>ACETAMINOPHEN AND CODEINE PHOSPHATE</u>			
<u>AA</u>	<u>HI TECH PHARMA</u>	<u>120MG/5ML;12MG/5ML</u>	<u>N40119 001</u>
			APR 26, 1996
<u>AA</u>	<u>MOVA</u>	<u>120MG/5ML;12MG/5ML</u>	<u>N40098 001</u>
			SEP 20, 1996
<u>TYLENOL W/ CODEINE</u>			
<u>AA</u>	<u>JOHNSON RW</u>	<u>120MG/5ML;12MG/5ML</u>	<u>N85057 001</u>

ACETAMINOPHEN; CODEINE PHOSPHATE

SOLUTION; ORAL			
<u>TYLENOL W/ CODEINE</u>			
<u>AA</u>	<u>+ JOHNSON RW</u>	<u>120MG/5ML;12MG/5ML</u>	<u>N85057 001</u>
SUSPENSION; ORAL			
<u>ACETAMINOPHEN AND CODEINE PHOSPHATE</u>			
> ADD >	<u>AA</u>	<u>CARNRICK</u>	<u>120MG/5ML;12MG/5ML</u>
> DLT >			
> DLT >	<u>AA</u>	<u>BARRE</u>	<u>120MG/5ML;12MG/5ML</u>
			N85883 001
<u>CAPITAL AND CODEINE</u>			
> ADD >	<u>AA</u>	<u>ALPHARMA</u>	<u>120MG/5ML;12MG/5ML</u>
> DLT >	<u>AA</u>	<u>CARNRICK</u>	<u>120MG/5ML;12MG/5ML</u>
			N85883 001

TABLET; ORAL			
<u>ACETAMINOPHEN AND CODEINE PHOSPHATE</u>			
<u>AA</u>	<u>BARR</u>	<u>300MG;15MG</u>	<u>N85795 001</u>
<u>AA</u>		<u>300MG;30MG</u>	<u>N85794 001</u>
@		300MG;15MG	N85795 001
@		300MG;30MG	N85794 001
<u>AA</u>	<u>GENEVA PHARMS</u>	<u>300MG;30MG</u>	<u>N85291 002</u>
<u>AA</u>		<u>300MG;60MG</u>	<u>N85964 001</u>
@		300MG;30MG	N85291 002
@		300MG;60MG	N85964 001
<u>AA</u>	<u>HALSEY</u>	<u>300MG;60MG</u>	<u>N86549 001</u>
@		300MG;60MG	N86549 001
<u>AA</u>	<u>LEMMON</u>	<u>300MG;15MG</u>	<u>N88627 001</u>
			MAR 06, 1985
<u>AA</u>		<u>300MG;30MG</u>	<u>N88628 001</u>
			MAR 06, 1985
<u>AA</u>		<u>300MG;60MG</u>	<u>N88629 001</u>
			MAR 06, 1985
<u>AA</u>	<u>MIKART</u>	<u>300MG;30MG</u>	<u>N89238 001</u>
			FEB 25, 1986
<u>AA</u>		<u>300MG;60MG</u>	<u>N89244 001</u>
			FEB 25, 1986
@		300MG;60MG	N89244 001
			FEB 25, 1986
<u>AA</u>	<u>TEVA</u>	<u>300MG;15MG</u>	<u>N88627 001</u>
			MAR 06, 1985
<u>AA</u>		<u>300MG;30MG</u>	<u>N88628 001</u>
			MAR 06, 1985
<u>AA</u>		<u>300MG;60MG</u>	<u>N88629 001</u>
			MAR 06, 1985
<u>ACETAMINOPHEN AND CODEINE PHOSPHATE #3</u>			
<u>AA</u>	<u>MIKART</u>	<u>300MG;30MG</u>	<u>N89238 001</u>
			FEB 25, 1986

ACETAMINOPHEN; CODEINE PHOSPHATE

TABLET; ORAL

<u>ACETAMINOPHEN AND CODEINE PHOSPHATE #3</u>		
AA	SUPERPHARM	300MG;30MG
	@	300MG;30MG
<u>ACETAMINOPHEN AND CODEINE PHOSPHATE #4</u>		
AA	SUPERPHARM	300MG;60MG
	@	300MG;60MG
<u>ACETAMINOPHEN W/ CODEINE</u>		
AA	BARR	300MG;60MG
	@	300MG;60MG
<u>ACETAMINOPHEN W/ CODEINE PHOSPHATE</u>		
AA	HALSEY	300MG;15MG
AA		300MG;30MG
	@	300MG;15MG
	@	300MG;30MG

ACETAMINOPHEN; HYDROCODONE BITARTRATE

TABLET; ORAL

<u>ANEXSIA</u>		
AA	KING PHARMS	500MG;5MG
AA	MALLINCKRODT CHEM	500MG;5MG
<u>ANEXSIA 10/660</u>		
	MALLINCKRODT	660MG;10MG
AA	+ MALLINCKRODT CHEM	660MG;10MG
<u>ANEXSIA 7.5/650</u>		
AA	KING PHARMS	650MG;7.5MG
AA	MALLINCKRODT CHEM	650MG;7.5MG
<u>HYDROCODONE BITARTRATE AND ACETAMINOPHEN</u>		
AA	KING PHARMS	500MG;5MG
AA		750MG;7.5MG

ACETAMINOPHEN; HYDROCODONE BITARTRATE

TABLET; ORAL

<u>HYDROCODONE BITARTRATE AND ACETAMINOPHEN</u>		
AA	MALLINCKRODT CHEM	500MG;5MG
AA		750MG;7.5MG
AA	ROYCE LABS	500MG;2.5MG
AA		500MG;5MG
AA		500MG;7.5MG
AA		650MG;7.5MG
AA		650MG;10MG
AA		750MG;7.5MG
AA	UCB	650MG;7.5MG
AA	VINTAGE PHARMS	500MG;7.5MG
AA		650MG;10MG
AA		750MG;7.5MG
	LORTAB	
	+ GRAHAM	500MG;10MG
	* UCB	500MG;10MG
<u>VICODIN HP</u>		
AA	KNOLL PHARM	660MG;10MG

ACETAMINOPHEN; OXYCODONE HYDROCHLORIDE

CAPSULE; ORAL

<u>OXYCODONE AND ACETAMINOPHEN</u>		
AA	HALSEY	500MG;5MG
	@	500MG;5MG
AA	VINTAGE PHARMS	500MG;5MG

> DLT >
> DLT >
> ADD >
> ADD >

N89994 001
MAY 04, 1989
N89994 001
MAY 04, 1989
N40106 001
JUL 30, 1996

ACETAMINOPHEN; OXYCODONE HYDROCHLORIDE

TABLET; ORAL
OXYCODONE AND ACETAMINOPHEN
AA VINTAGE PHARMS 325MG;5MG N40105 001
 JUL 30, 1996

ACETAMINOPHEN; PROPOXYPHENE HYDROCHLORIDE

TABLET; ORAL
PROPOXYPHENE HCL AND ACETAMINOPHEN
 > ADD > AA ROYCE LABS 650MG;65MG N40139 001
 > ADD > DEC 16, 1996

ACETAMINOPHEN; PROPOXYPHENE NAPSYLATE

TABLET; ORAL
PROPOXYPHENE NAPSYLATE AND ACETAMINOPHEN
AB SUPERPHARM 650MG;100MG N71319 001
 @ 650MG;100MG JAN 05, 1987
 N71319 001
 JAN 06, 1987

ACETIC ACID, GLACIAL

SOLUTION/DROPS; OTIC
ACETIC ACID
AT MORTON GROVE 2% N40166 001
 JUL 26, 1996

ACETIC ACID, GLACIAL; HYDROCORTISONE

SOLUTION/DROPS; OTIC
HYDROCORTISONE AND ACETIC ACID
AT MORTON GROVE 2%;1% N40168 001
 AUG 30, 1996

ACITRETIN

CAPSULE; ORAL
 SORIATANE
 ROCHE 10MG N19821 001
 OCT 28, 1996

ACITRETIN

CAPSULE; ORAL
 SORIATANE
 + ROCHE 25MG N19821 002
 OCT 28, 1996

ACRIVASTINE; PSEUDOEPHEDRINE HYDROCHLORIDE

CAPSULE; ORAL
 SEMPRES-D
 + GLAXO WELLCOME 8MG;60MG N19806 001
 + MEDEVA 8MG;60MG MAR 25, 1994
 N19806 001
 MAR 25, 1994

ADAPALENE

GEL; TOPICAL
 DIFFERIN
 + GALDERMA 0.1% N20380 001
 MAY 31, 1996

SOLUTION; TOPICAL

DIFFERIN
 + GALDERMA 0.1% N20338 001
 MAY 31, 1996

ALBENDAZOLE

TABLET; ORAL
 ALBENZA
 + SMITHKLINE BEECHAM 200MG N20666 001
 JUN 11, 1996

ALBUTEROLAEROSOL, METERED; INHALATION

ALBUTEROL
AB ARMSTRONG PHARMS 0.09MG/INH N72273 001
 AUG 14, 1996
AB MEDISOL 0.09MG/INH N74072 001
 AUG 01, 1996

ALBUTEROL SULFATEAEROSOL, METERED; INHALATION
PROVENTIL-HFA+ 3M EQ 0.09MG BASE/INH N20503 001
AUG 15, 1996

SOLUTION; INHALATION

VENTOLIN

<u>AN</u>	GLAXO WELLCOME	<u>EQ 0.083% BASE</u>	N19773 001 APR 23, 1992
<u>AN</u>	+	<u>EQ 0.083% BASE</u>	N19773 001 APR 23, 1992
<u>AN</u>		<u>EQ 0.5% BASE</u>	N19269 002 JAN 16, 1987
<u>AN</u>	+	<u>EQ 0.5% BASE</u>	N19269 002 JAN 16, 1987

SYRUP; ORAL
PROVENTIL

<u>AA</u>	SCHERING	<u>EQ 2MG BASE/5ML</u>	N18062 001 JAN 19, 1983
<u>AA</u>	+	<u>EQ 2MG BASE/5ML</u>	N18062 001 JAN 19, 1983

TABLET; ORAL

ALBUTEROL SULFATE

<u>AB</u>	LEMMON	<u>EQ 2MG BASE</u>	N72938 001 MAR 30, 1990
<u>AB</u>		<u>EQ 4MG BASE</u>	N72939 001 MAR 30, 1990
<u>AB</u>	TEVA	<u>EQ 2MG BASE</u>	N72938 001 MAR 30, 1990
<u>AB</u>		<u>EQ 4MG BASE</u>	N72939 001 MAR 30, 1990

ALBUTEROL SULFATE; IPRATROPIUM BROMIDEAEROSOL, METERED; INHALATION
COMBIVENT+ BOEHRINGER INGELHEIM EQ 0.09MG BASE/INH;
0.018MG/INH N20291 001
OCT 24, 1996ALLOPURINOL

TABLET; ORAL

ALLOPURINOL

<u>AB</u>	BARR	<u>100MG</u>	N70466 001 DEC 24, 1985
<u>AB</u>		<u>300MG</u>	N70467 001 DEC 24, 1985
	@	100MG	N70466 001 DEC 24, 1985
	@	300MG	N70467 001 DEC 24, 1985

ALLOPURINOL SODIUMINJECTABLE; INJECTION
ZYLOPRIM+ GLAXO WELLCOME EQ 500MG BASE/VIAL N20298 001
MAY 17, 1996ALPRAZOLAM

TABLET; ORAL

ALPRAZOLAM

<u>AB</u>	NOVOPHARM	<u>2MG</u>	N74085 004 FEB 26, 1996
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ALPROSTADIL

INJECTABLE; INJECTION

CAVERJECT

PHARMACIA AND UPJOHN 0.005MG/VIAL N20379 003
JUN 27, 1996

SUPPOSITORY; URETHRAL

MUSE

VIVUS

0.125MG	N20700 001 NOV 19, 1996
0.25MG	N20700 002 NOV 19, 1996
0.5MG	N20700 003 NOV 19, 1996
1MG	N20700 004 NOV 19, 1996

AMINO ACIDS

INJECTABLE; INJECTION

AMINOSYN-HF 8%

ABBOTT 8%

N20345 001
APR 04, 1996

CLINISOL 15% SULFITE FREE IN PLASTIC CONTAINER

CLINTEC NUTR 15%

N20512 001
AUG 30, 1996

HEPATASOL 8%

BAXTER HLTHCARE 8%

N20360 001
APR 04, 1996AMINOPHYLLINE

TABLET; ORAL

AMINOPHYLLINE

@ PHOENIX LABS NY 100MG

N85409 001

@ 200MG

N85410 001

@ VINTAGE PHARMS 100MG

N85409 001

@ 200MG

N85410 001

AMITRIPTYLINE HYDROCHLORIDE

TABLET; ORAL

AMITRIPTYLINE HCL

BP HALSEY 25MG

N85922 001

@ 25MG

N85922 001

ENDEP

AB ROCHE 150MG

N85303 001

@ 150MG

N85303 001

AMITRIPTYLINE HYDROCHLORIDE; CHLORDIAZEPOXIDE

TABLET; ORAL

CHLORDIAZEPOXIDE AND AMITRIPTYLINE HCL

AB BARR EQ 12.5MG BASE;5MG

N70765 001

DEC 10, 1986

AB EQ 25MG BASE;10MG

N70766 001

DEC 10, 1986

@ EQ 12.5MG BASE;5MG

N70765 001

DEC 10, 1986

@ EQ 25MG BASE;10MG

N70766 001

DEC 10, 1986

AMITRIPTYLINE HYDROCHLORIDE; PERPHENAZINE

TABLET; ORAL

PERPHENAZINE AND AMITRIPTYLINE HCL

AB BARR 10MG;2MG

N71077 001

NOV 12, 1986

AB 10MG;4MG

N71078 001

NOV 12, 1986

AB 25MG;2MG

N70297 001

NOV 12, 1986

AB 25MG;4MG

N71079 001

NOV 12, 1986

@ 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

> ADD > AMLEXANOX> ADD > PASTE; DENTAL> ADD > APHTHASOL> ADD > + BLOCK DRUG 5%

N20511 001

DEC 17, 1996

> ADD >AMMONIUM LACTATE

CREAM; TOPICAL

LAC-HYDRIN

BRISTOL MYERS EQ 12% BASE

N20508 001

AUG 29, 1986

WESTWOOD SQUIBB EQ 12% BASE

N20508 001

AUG 29, 1996

AMOXICILLIN

CAPSULE; ORAL

AMOXIL

AB SMITHKLINE BEECHAM 500MG

N62216 004

@ 250MG

N62216 003

LACTID

AB SMITHKLINE BEECHAM 250MG

N62216 003

AB 500MG

N62216 004

AMOXICILLIN

TABLET, CHEWABLE; ORAL

<u>AB</u>	<u>AMOXICILLIN</u>	<u>125MG</u>	N64131 001
	APOTHECON		MAY 06, 1996
<u>AB</u>		<u>250MG</u>	N64131 002
			MAY 06, 1996
<u>AB</u>	CLONMEL HLTH CARE	<u>125MG</u>	N64139 001
			JAN 29, 1996
<u>AB</u>		<u>250MG</u>	N64139 002
			JAN 29, 1996
> <u>ADD</u> >	<u>AB</u>	<u>125MG</u>	N64031 001
> <u>ADD</u> >	NOVOPHARM		DEC 19, 1996
> <u>ADD</u> >	<u>AB</u>	<u>250MG</u>	N64031 002
> <u>ADD</u> >			DEC 19, 1996

AMOXICILLIN; CLAVULANATE POTASSIUMPOWDER FOR RECONSTITUTION; ORAL
AUGMENTIN '200'

+	SMITHKLINE BEECHAM	200MG/5ML; EQ 28.5MG BASE/5ML	N50725 001
			MAY 31, 1996

	AUGMENTIN '400'		
+	SMITHKLINE BEECHAM	400MG/5ML; EQ 57MG BASE/5ML	N50725 002
			MAY 31, 1996

TABLET; ORAL

	AUGMENTIN '875'		
+	SMITHKLINE BEECHAM	875MG;EQ 125MG BASE	N50720 001
			FEB 13, 1996

TABLET, CHEWABLE; ORAL
AUGMENTIN '200'

+	SMITHKLINE BEECHAM	200MG;EQ 28.5MG BASE	N50726 001
			MAY 31, 1996

	AUGMENTIN '400'		
+	SMITHKLINE BEECHAM	400MG;EQ 57MG BASE	N50726 002
			MAY 31, 1996

AMPHETAMINE ASPARTATE; AMPHETAMINE SULFATE; DEXTROAMPHETAMINE
SACCHARATE; DEXTROAMPHETAMINE SULFATE

TABLET; ORAL

ADDERALL 10			
RICHWOOD PHARM	2.5MG;2.5MG;2.5MG;2.5MG	N11522 007	
		FEB 13, 1996	

ADDERALL 20			
+ RICHWOOD PHARM	5MG;5MG;5MG;5MG	N11522 008	
		FEB 13, 1996	

AMPHETAMINE RESIN COMPLEX; DEXTROAMPHETAMINE RESIN COMPLEX

CAPSULE, EXTENDED RELEASE; ORAL

BIPHETAMINE 12.5			
@ FISOXS	EQ 6.25MG BASE;		
	EQ 6.25MG BASE	N10093 007	
@ MEDEVA PHARMS	EQ 6.25MG BASE;		
	EQ 6.25MG BASE	N10093 007	

BIPHETAMINE 20			
@ FISOXS	EQ 10MG BASE;EQ 10MG BASE	N10093 003	
@ MEDEVA PHARMS	EQ 10MG BASE;EQ 10MG BASE	N10093 003	

BIPHETAMINE 7.5			
@ FISOXS	EQ 3.75MG BASE;		
	EQ 3.75MG BASE	N10093 009	
@ MEDEVA PHARMS	EQ 3.75MG BASE;		
	EQ 3.75MG BASE	N10093 009	

AMPHOTERICIN B

INJECTABLE; INJECTION

<u>AMPHOTERICIN B</u>			
SANOFI WINTHROP	<u>50MG/VIAL</u>	N64141 001	
		DEC 23, 1996	

INJECTABLE, LIPID COMPLEX; INJECTION
AMPHOTEC

+	SEQUUS PHARMS	50MG/VIAL	N50729 001
			NOV 22, 1996
+		100MG/VIAL	N50729 002
			NOV 22, 1996

SUSPENSION; ORAL

FUNGIZONE			
+ BRISTOL MYERS SQUIBB	100MG/ML	N50341 003	

AMPICILLIN SODIUM

INJECTABLE; INJECTION

OMNIPEN-N

AP	WYETH AYERST	EQ 2GM BASE/VIAL	N60626 005
AP	+	EQ 2GM BASE/VIAL	N60626 005
AP	<u>PENERGIN-S</u>		
AP	WYETH AYERST	EQ 125MG BASE/VIAL	N50072 001
AP		EQ 250MG BASE/VIAL	N50072 002
AP		EQ 500MG BASE/VIAL	N50072 003
AP		EQ 1GM BASE/VIAL	N50072 004
AP	*	EQ 2GM BASE/VIAL	N50072 005
	@	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

AMPICILLIN/AMPICILLIN TRIHYDRATE

CAPSULE; ORAL

PFIZERPEN-A

AB	PFIZER	EQ 250MG BASE	N62050 001
AB		EQ 500MG BASE	N62050 002
	@	EQ 250MG BASE	N62050 001
	@	EQ 500MG BASE	N62050 002

POWDER FOR RECONSTITUTION; ORAL

PFIZERPEN-A

AB	PFIZER	EQ 125MG BASE/5ML	N62049 001
AB		EQ 250MG BASE/5ML	N62049 002
	@	EQ 125MG BASE/5ML	N62049 001
	@	EQ 250MG BASE/5ML	N62049 002

AMPICILLIN/AMPICILLIN TRIHYDRATE; PROBENECID

POWDER FOR RECONSTITUTION; ORAL

POLYCILLIN-PRE

AB	* APOTHECON	EQ 3.5GM BASE/BOT;1GM/BOT	N61898 001
	@	EQ 3.5GM BASE/BOT;1GM/BOT	N61898 001
AB	<u>PROBAMPACIN</u>		
AB	BIOCRIFT	EQ 3.5GM BASE/BOT;1GM/BOT	N61741 001
	+	EQ 3.5GM BASE/BOT;1GM/BOT	N61741 001

ASPIRIN; BUTALBITAL; CAFFEINE

TABLET; ORAL

BUTALBITAL, ASPIRIN & CAFFEINE

AB	HALSEY	325MG;50MG;40MG	N89448 001
	@	325MG;50MG;40MG	DEC 01, 1986
			N89448 001
			DEC 01, 1986

ASPIRIN; CAFFEINE; ORPHENADRINE CITRATE

TABLET; ORAL

INVAGESIC

AB	INVAMED	385MG;30MG;25MG	N74817 001
			NOV 27, 1996

INVAGESIC FORTE

AB	INVAMED	770MG;60MG;50MG	N74817 002
			NOV 27, 1996

NORGESIC

AB	3M	385MG;30MG;25MG	N13416 003
			OCT 27, 1982

NORGESIC FORTE

AB	+ 3M	770MG;60MG;50MG	N13416 004
			OCT 27, 1982

ORPHENADRINE CITRATE, ASPIRIN, AND CAFFEINE

AB	EON LABS MFG	385MG;30MG;25MG	N74654 001
			DEC 31, 1996
AB		770MG;60MG;50MG	N74654 002
			DEC 31, 1996

ASPIRIN; CARISOPRODOL

TABLET; ORAL

CARISOPRODOL AND ASPIRIN

AB	EON LABS MFG	325MG;200MG	N40116 001
			APR 25, 1996

ASPIRIN; CARISOPRODOL; CODEINE PHOSPHATE

TABLET; ORAL

CARISOPRODOL, ASPIRIN AND CODEINE PHOSPHATE

AB	EON LABS MFG	325MG;200MG;16MG	N40118 001
			APR 16, 1996

SOMA COMPOUND W/ CODEINE

AB	+ WALLACE PHARMS	325MG;200MG;16MG	N12366 002
			JUL 11, 1983

> ADD > ATORVASTATIN CALCIUM

> ADD > TABLET; ORAL
 > ADD > LIPITOR
 > ADD > PARKE DAVIS EQ 10MG BASE N20702 001
 > ADD > DEC 17, 1996
 > ADD > EQ 20MG BASE N20702 002
 > ADD > DEC 17, 1996
 > ADD > + EQ 40MG BASE N20702 003
 > ADD > DEC 17, 1996

ATRACURIUM BESYLATE

INJECTABLE; INJECTION
 > ADD > ATRACURIUM BESYLATE
 > ADD > AP ABBOTT 10MG/ML N74632 001
 > ADD > DEC 23, 1996
 > ADD > AP 10MG/ML N74633 001
 > ADD > DEC 23, 1996
 > ADD > TRACRIUM
 > ADD > AP + GLAXO WELLCOME 10MG/ML N18831 001
 > ADD > NOV 23, 1983

ATROPINE SULFATE; DIPHENOXYLATE HYDROCHLORIDE

TABLET; ORAL
DIPHENOXYLATE HCL AND ATROPINE SULFATE
AA BARR 0.025MG; 2.5MG N85506 001
 @ 0.025MG; 2.5MG N85506 001
AA LOGEN 0.025MG; 2.5MG N88962 001
 SUPERPHARM MAY 10, 1985
 @ 0.025MG; 2.5MG N88962 001
 MAY 10, 1985
AA LOW-QUEL 0.025MG; 2.5MG N85211 001
 HALSEY 0.025MG; 2.5MG N85211 001
 @

AZATHIOPRINE

TABLET; ORAL
AZATHIOPRINE
AB ROXANE 50MG N74069 001
 FEB 16, 1996
IMURAN
AB + GLAXO WELLCOME 50MG N16324 001

AZELASTINE HYDROCHLORIDE

SPRAY, METERED; NASAL
 ASTELIN
 + WALLACE EQ 0.125MG BASE/INH N20114 001
 NOV 01, 1996

AZITHROMYCIN DIHYDRATE

TABLET; ORAL
 ZITHROMAX
 + PFIZER EQ 250MG BASE N50711 001
 JUL 18, 1996
 + EQ 600MG BASE N50730 001
 JUN 12, 1996

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
 * SQUIBB 500MG/VIAL N50580 001
 DEC 31, 1986
 * 1GM/VIAL N50580 002
 DEC 31, 1986
 * 2GM/VIAL N50580 003
 DEC 31, 1986

BACITRACIN

OINTMENT; OPHTHALMIC
BACITRACIN
AT ALTANA 500 UNITS/GM N61212 001
 + 500 UNITS/GM N61212 001
AT * LILLY 500 UNITS/GM N60687 001
 @ 500 UNITS/GM N60687 001

BACLOFEN

INJECTABLE; INTRATHECAL
LIORESAL
+ MEDTRONIC

0.05MG/ML

N20075 003
NOV 07, 1996

TABLET; ORAL

BACLOFEN

AB CHELSEA LABS

10MG

N74698 001
AUG 20, 1996

AB 20MG

N74698 002
AUG 20, 1996

AB ROSEMONT 10MG

N74584 001
AUG 19, 1996

AB 20MG

N74584 002
AUG 19, 1996

> ADD >> ADD >> ADD >> ADD >> ADD >> ADD >BECLOMETHASONE DIPROPIONATE

AEROSOL, METERED; INHALATION
VANCERIL DOUBLE STRENGTH

+ SCHERING 0.084MG/INH

N20486 001
DEC 24, 1996

> ADD >> ADD >> ADD >BECLOMETHASONE DIPROPIONATE MONOHYDRATE

SPRAY, METERED; NASAL
BECONASE AQ

BN + GLAXO WELLCOME

EQ 0.042MG DIPROP/INH

N19389 001
JUL 27, 1987

BN EQ 0.042MG DIPROP/INH

N19389 001
JUL 27, 1987

VANCENASE AQ

BN SCHERING

EQ 0.042MG DIPROP/INH

N19589 001
DEC 23, 1987

BN + EQ 0.042MG DIPROP/INH

N19589 001
DEC 23, 1987

+ EQ 0.084MG DIPROP/INH

N20469 001
JUN 26, 1996

BENTIROMIDE

SOLUTION; ORAL

CHYMEX

SAVAGE LABS

500MG/7.5ML

N18366 001
DEC 29, 1983

BENTIROMIDE

SOLUTION; ORAL

CHYMEX

@ SAVAGE LABS

500MG/7.5ML

N18366 001
DEC 29, 1983

BENZTROPINE MESYLATE

TABLET; ORAL

BENZTROPINE MESYLATE

ROSEMONT

0.5MG

N40103 001
DEC 12, 1996

> ADD >> ADD >> ADD >> ADD >> ADD >> ADD >

AA

AA

AA

AA

1MG2MG

N40103 002
DEC 12, 1996

N40103 003
DEC 12, 1996

BETA-CAROTENE

CAPSULE; ORAL

SOLATENE

+ ROCHE

30MG

N17589 001

@

30MG

N17589 001

BETAINE, ANHYDROUS

POWDER FOR RECONSTITUTION; ORAL

CYSTADANE

+ ORPHAN MEDCL

1GM/SCOOPFUL

N20576 001
OCT 25, 1996

BETAMETHASONE BENZOATE

CREAM; TOPICAL

UTICORT

+ PARKE DAVIS

0.025%

N16998 002

@

0.025%

N16998 002

BISMUTH SUBSALICYLATE; METRONIDAZOLE; TETRACYCLINE HYDROCHLORIDETABLET, CHEWABLE, TABLET, CAPSULE; ORAL
HELIDAC+ PROCTER AND GAMBLE 262.4MG;250MG;500MG N50719 001
AUG 15, 1996BLEOMYCIN SULFATE

INJECTABLE; INJECTION

BLENOXANEAP + BRISTOL MYERS SQUIBB EQ 15 UNITS BASE/VIAL N50443 001
EQ 15 UNITS BASE/VIAL N50443 001BLEOMYCIN SULFATEAP PHARMACIA AND UPJOHN EQ 15 UNITS BASE/VIAL N64084 001
JUN 01, 1996
+ N64084 002
EQ 30 UNITS BASE/VIAL JUN 01, 1996BRIMONIDINE TARTRATESOLUTION/DROPS; OPHTHALMIC
ALPHAGAN+ ALLERGAN 0.2% N20613 001
SEP 06, 1996BROMPHENIRAMINE MALEATE

TABLET; ORAL

DIMETANEAA + ROBINS AH 4MG N10799 003
@ WHITEHALL ROBINS 4MG N10799 003BUCLIZINE HYDROCHLORIDETABLET; ORAL
BUCLADIN-SSTUART 50MG N10911 006
@ 50MG N10911 006BUMETANIDE

TABLET; ORAL

BUMETANIDE

AB EON LABS MFG 0.5MG

AB 1MG

AB 2MG

N74700 001
NOV 21, 1996
N74700 002
NOV 21, 1996
N74700 003
NOV 21, 1996BUPRENORPHINE HYDROCHLORIDE

INJECTABLE; INJECTION

BUPRENEX

AP + RECKITT AND COLMAN EQ 0.3MG BASE/ML N18401 001

AP BUPRENORPHINE HCL
SANOFI WINTHROP EQ 0.3MG BASE/ML N74137 001
JUN 03, 1996BUPROPION HYDROCHLORIDETABLET, EXTENDED RELEASE; ORAL
WELLBUTRIN

+ GLAXO WELLCOME 50MG N20358 001

+ 100MG OCT 04, 1996

+ 150MG N20358 002
OCT 04, 1996N20358 003
OCT 04, 1996BUSPIRONE HYDROCHLORIDE

TABLET; ORAL

BUSPAR

+ BRISTOL MYERS SQUIBB 10MG

N18731 002
SEP 29, 1986

10MG N18731 002

SEP 29, 1986
N18731 003

+ 15MG

APR 22, 1996
N18731 004

@ 30MG

APR 22, 1996

BUTABARBITAL SODIUM

<u>ELIXIR, ORAL</u>			
<u>AA</u>	<u>SARISOL</u>		
	<u>HALSEY</u>	<u>30MG/5ML</u>	<u>N84723 001</u>
	@	30MG/5ML	N84723 001
<u>TABLET, ORAL</u>			
<u>AA</u>	<u>SARISOL NO. 1</u>		
	<u>HALSEY</u>	<u>15MG</u>	<u>N84719 001</u>
	@	15MG	N84719 001
<u>AA</u>	<u>SARISOL NO. 2</u>		
	<u>HALSEY</u>	<u>30MG</u>	<u>N84719 002</u>
	@	30MG	N84719 002

BUTENAFINE HYDROCHLORIDE

CREAM; TOPICAL		
MENTAX		
+ PENEDERM	1%	N20524 001
		OCT 18, 1996

> ADD > CABERGOLINE

> <u>ADD</u> >	TABLET; ORAL		
> <u>ADD</u> >	DOSTINEX		
> <u>ADD</u> >	+ PHARMACIA AND UPJOHN 0.5MG		
> <u>ADD</u> >		N20664 001	
		DEC 23, 1996	

CALCIPOTRIENE

CREAM; TOPICAL		
DOVONEX		
+ BRISTOL MYERS SQUIBB 0.005%		N20554 001
		JUL 22, 1996

CALCITONIN, HUMAN

<u>INJECTABLE, INJECTION</u>			
	<u>CIBACALCIN</u>		
	* CIBA	<u>0.5MG/VIAL</u>	<u>N18470 001</u>
			OCT 31, 1986
	@	0.5MG/VIAL	N18470 001
			OCT 31, 1986

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

<u>SOLUTION; INTRAPERITONEAL</u>			
<u>AT</u>	<u>DELFLX W/ DEXTROSE 1.5% IN PLASTIC CONTAINER</u>		
	FRESENIUS	<u>25.7MG/100ML; 1.5GM/100ML;</u> <u>15.2MG/100ML; 567MG/100ML;</u> <u>392MG/100ML</u>	N18379 002
<u>AT</u>	<u>DELFLX W/ DEXTROSE 2.5% IN PLASTIC CONTAINER</u>		
	FRESENIUS	<u>25.7MG/100ML; 2.5GM/100ML;</u> <u>15.2MG/100ML; 567MG/100ML;</u> <u>392MG/100ML</u>	N18379 003
<u>AT</u>	<u>DELFLX W/ DEXTROSE 3.5% IN PLASTIC CONTAINER</u>		
	FRESENIUS	<u>25.7MG/100ML; 3.5GM/100ML;</u> <u>15.2MG/100ML; 567MG/100ML;</u> <u>392MG/100ML</u>	N18379 007
			JUN 24, 1988
<u>AT</u>	<u>DELFLX W/ DEXTROSE 4.25% IN PLASTIC CONTAINER</u>		
	FRESENIUS	<u>25.7MG/100ML; 4.25GM/100ML;</u> <u>15.2MG/100ML; 567MG/100ML;</u> <u>392MG/100ML</u>	N18379 001
<u>AT</u>	<u>DELFLX-LM W/ DEXTROSE 1.5% IN PLASTIC CONTAINER</u>		
	FRESENIUS	<u>25.7MG/100ML; 1.5GM/100ML;</u> <u>5.08MG/100ML; 538MG/100ML;</u> <u>448MG/100ML</u>	N18379 004
			JUL 07, 1982
<u>AT</u>	<u>DELFLX-LM W/ DEXTROSE 2.5% IN PLASTIC CONTAINER</u>		
	FRESENIUS	<u>25.7MG/100ML; 2.5GM/100ML;</u> <u>5.08MG/100ML; 538MG/100ML;</u> <u>448MG/100ML</u>	N18379 005
			JUL 07, 1982
<u>AT</u>	<u>DELFLX-LM W/ DEXTROSE 3.5% IN PLASTIC CONTAINER</u>		
	FRESENIUS	<u>25.7MG/100ML; 3.5GM/100ML;</u> <u>5.08MG/100ML; 538MG/100ML;</u> <u>448MG/100ML</u>	N18379 008
			JUN 24, 1988
<u>AT</u>	<u>DELFLX-LM W/ DEXTROSE 4.25% IN PLASTIC CONTAINER</u>		
	FRESENIUS	<u>25.7MG/100ML; 4.25GM/100ML;</u> <u>5.08MG/100ML; 538MG/100ML;</u> <u>448MG/100ML</u>	N18379 006
			JUL 07, 1982
<u>AT</u>	<u>INPERSOL W/ DEXTROSE 1.5% IN PLASTIC CONTAINER</u>		
	FRESENIUS	<u>25.7MG/100ML; 1.5GM/100ML;</u> <u>15.2MG/100ML; 567MG/100ML;</u> <u>392MG/100ML</u>	N18379 002
<u>AT</u>	<u>INPERSOL W/ DEXTROSE 2.5% IN PLASTIC CONTAINER</u>		
	FRESENIUS	<u>25.7MG/100ML; 2.5GM/100ML;</u> <u>15.2MG/100ML; 567MG/100ML;</u> <u>392MG/100ML</u>	N18379 003

CALCIUM CHLORIDE; DEXTROSE; MAGNESIUM CHLORIDE; SODIUM CHLORIDE;
SODIUM LACTATESOLUTION; INTRAPERITONEAL

AT INPERSOL W/ DEXTROSE 3.5% IN PLASTIC CONTAINER
FRESENIUS 25.7MG/100ML; 3.5GM/100ML;
 15.2MG/100ML; 567MG/100ML;
 392MG/100ML N18379 007

JUN 24, 1988

AT INPERSOL W/ DEXTROSE 4.25% IN PLASTIC CONTAINER
FRESENIUS 25.7MG/100ML; 4.25GM/100ML;
 15.2MG/100ML; 567MG/100ML;
 392MG/100ML N18379 001

AT INPERSOL-LM W/ DEXTROSE 1.5% IN PLASTIC CONTAINER
FRESENIUS 25.7MG/100ML; 1.5GM/100ML;
 5.08MG/100ML; 538MG/100ML;
 448MG/100ML N18379 004

JUL 07, 1982

AT INPERSOL-LM W/ DEXTROSE 2.5% IN PLASTIC CONTAINER
FRESENIUS 25.7MG/100ML; 2.5GM/100ML;
 5.08MG/100ML; 538MG/100ML;
 448MG/100ML N18379 005

JUL 07, 1982

AT INPERSOL-LM W/ DEXTROSE 3.5% IN PLASTIC CONTAINER
FRESENIUS 25.7MG/100ML; 3.5GM/100ML;
 5.08MG/100ML; 538MG/100ML;
 448MG/100ML N18379 008

JUN 24, 1988

AT INPERSOL-LM W/ DEXTROSE 4.25% IN PLASTIC CONTAINER
FRESENIUS 25.7MG/100ML; 4.25GM/100ML;
 5.08MG/100ML; 538MG/100ML;
 448MG/100ML N18379 006

JUL 07, 1982

CALCIUM CHLORIDE; DEXTROSE; MAGNESIUM SULFATE; POTASSIUM
CHLORIDE; SODIUM BICARBONATE; SODIUM CHLORIDE; SODIUM
PHOSPHATE, DIBASICINJECTABLE; INTRATHECALELLIOTTS B SOLUTION

+ ORPHAN MEDCL 0.2MG/ML; 0.8MG/ML; 0.3MG/ML; 0.3MG/ML;
 1.9MG/ML; 7.3MG/ML;
 0.2MG/ML N20577 001

SEP 27, 1996

CAPTOPRILTABLET; ORALCAPOTEN

AB BRISTOL MYERS SQUIBB 100MG N18343 003
 AB + 100MG N18343 003
 75MG N18343 007

* 150MG JUN 13, 1995

@ 75MG N18343 004

@ 150MG N18343 007

JUN 13, 1995

N18343 004

JUN 13, 1995

N18343 004

JUN 13, 1995

CAPTOPRILAB BAKER NORTON 12.5MG N74590 004

AUG 30, 1996

AB 25MG N74590 002

AUG 30, 1996

AB 50MG N74590 001

AUG 30, 1996

AB 100MG N74590 003

AUG 30, 1996

AB BIOCRAFT 12.5MG N74433 001

FEB 13, 1996

AB 25MG N74433 002

FEB 13, 1996

AB 50MG N74433 003

FEB 13, 1996

AB 100MG N74433 004

FEB 13, 1996

AB CHELSEA LABS 12.5MG N74576 001

APR 23, 1996

AB 25MG N74576 002

APR 23, 1996

AB 50MG N74576 003

APR 23, 1996

AB 100MG N74576 004

APR 23, 1996

AB COPLEY PHARM 12.5MG N74462 001

FEB 13, 1996

AB 25MG N74462 002

FEB 13, 1996

AB 50MG N74462 003

FEB 13, 1996

AB 100MG N74462 004

FEB 13, 1996

AB DANBURY PHARMA 12.5MG N74386 001

MAY 23, 1996

CAPTOPRILTABLET; ORAL
CAPTOPRIL

<u>AB</u>	DANBURY PHARMA	<u>25MG</u>
<u>AB</u>		<u>50MG</u>
<u>AB</u>		<u>100MG</u>
<u>AB</u>	ENDO LABS	<u>12.5MG</u>
<u>AB</u>		<u>25MG</u>
<u>AB</u>		<u>50MG</u>
<u>AB</u>		<u>100MG</u>
<u>AB</u>	EON LABS MFG	<u>12.5MG</u>
<u>AB</u>		<u>25MG</u>
<u>AB</u>		<u>50MG</u>
<u>AB</u>		<u>100MG</u>
<u>AB</u>	HALLMARK PHARMS	<u>12.5MG</u>
<u>AB</u>		<u>25MG</u>
<u>AB</u>		<u>50MG</u>
<u>AB</u>		<u>100MG</u>
<u>AB</u>	INVAMED	<u>12.5MG</u>
<u>AB</u>		<u>25MG</u>
<u>AB</u>		<u>50MG</u>
<u>AB</u>		<u>100MG</u>
<u>AB</u>	LEMMON	<u>12.5MG</u>
<u>AB</u>		<u>25MG</u>
<u>AB</u>		<u>50MG</u>

N74386 002
 MAY 23, 1996
 N74386 003
 MAY 23, 1996
 N74386 004
 MAY 23, 1996
 N74418 001
 FEB 13, 1996
 N74418 002
 FEB 13, 1996
 N74418 003
 FEB 13, 1996
 N74418 004
 FEB 13, 1996
 N74519 001
 FEB 13, 1996
 N74519 002
 FEB 13, 1996
 N74519 003
 FEB 13, 1996
 N74519 004
 FEB 13, 1996
 N74477 001
 FEB 13, 1996
 N74477 002
 FEB 13, 1996
 N74477 003
 FEB 13, 1996
 N74477 004
 FEB 13, 1996
 N74481 001
 FEB 13, 1996
 N74481 002
 FEB 13, 1996
 N74481 003
 FEB 13, 1996
 N74481 004
 FEB 13, 1996
 N74483 001
 FEB 13, 1996
 N74483 002
 FEB 13, 1996
 N74483 003
 FEB 13, 1996

CAPTOPRILTABLET; ORAL
CAPTOPRIL

<u>AB</u>	LEMMON	<u>100MG</u>
<u>AB</u>	MOVA	<u>12.5MG</u>
<u>AB</u>		<u>25MG</u>
<u>AB</u>		<u>50MG</u>
<u>AB</u>		<u>100MG</u>
<u>AB</u>	MYLAN	<u>12.5MG</u>
<u>AB</u>		<u>25MG</u>
<u>AB</u>		<u>50MG</u>
<u>AB</u>		<u>100MG</u>
<u>AB</u>	NOVOPHARM	<u>12.5MG</u>
<u>AB</u>		<u>25MG</u>
<u>AB</u>		<u>50MG</u>
<u>AB</u>		<u>100MG</u>
<u>AB</u>	PAR PHARM	<u>12.5MG</u>
<u>AB</u>		<u>25MG</u>
<u>AB</u>		<u>50MG</u>
<u>AB</u>		<u>100MG</u>
<u>AB</u>	ROYCE LABS	<u>12.5MG</u>
<u>AB</u>		<u>25MG</u>
<u>AB</u>		<u>50MG</u>
<u>AB</u>		<u>100MG</u>
<u>AB</u>	WEST WARD	<u>12.5MG</u>

N74483 004
 FEB 13, 1996
 N74423 001
 FEB 13, 1996
 N74423 002
 FEB 13, 1996
 N74423 003
 FEB 13, 1996
 N74423 004
 FEB 13, 1996
 N74434 001
 FEB 13, 1996
 N74434 002
 FEB 13, 1996
 N74434 003
 FEB 13, 1996
 N74434 004
 FEB 13, 1996
 N74322 001
 FEB 13, 1996
 N74322 002
 FEB 13, 1996
 N74322 003
 FEB 13, 1996
 N74322 004
 FEB 13, 1996
 N74493 001
 FEB 13, 1996
 N74493 002
 FEB 13, 1996
 N74493 003
 FEB 13, 1996
 N74493 004
 FEB 13, 1996
 N74451 001
 FEB 13, 1996
 N74451 002
 FEB 13, 1996
 N74451 003
 FEB 13, 1996
 N74451 004
 FEB 13, 1996
 N74505 001
 FEB 13, 1996

CAPTOPRIL

TABLET; ORAL			
<u>CAPTOPRIL</u>			
<u>AB</u>	WEST WARD	<u>25MG</u>	N74505 002 FEB 13, 1996
<u>AB</u>		<u>50MG</u>	N74505 003 FEB 13, 1996
<u>AB</u>		<u>100MG</u>	N74505 004 FEB 13, 1996

CARBAMAZEPINE

TABLET; ORAL			
<u>CARBAMAZEPINE</u>			
<u>AB</u>	TARO	<u>200MG</u>	N74649 001 OCT 03, 1996
TABLET, EXTENDED RELEASE; ORAL			
TEGRETOL-XR			
+	CIBA GEIGY	100MG	N20234 001 MAR 25, 1996
+		200MG	N20234 002 MAR 25, 1996
+		400MG	N20234 003 MAR 25, 1996

CARISOPRODOL

TABLET; ORAL			
<u>CARISOPRODOL</u>			
> <u>ADD</u> >	<u>AA</u> ROYCE LABS	<u>350MG</u>	N40152 001 DEC 03, 1996
> <u>ADD</u> >	<u>AA</u> WEST WARD	<u>350MG</u>	N40124 001 JAN 24, 1996

CARMUSTINE

IMPLANT; INTRACRANIAL			
GLIADEL			
+	GUILFORD PHARMS	7.7MG	N20637 001 SEP 23, 1996

CEFACLOL

CAPSULE; ORAL			
<u>CEFACLOL</u>			
<u>AB</u>	BIOCRAFT	<u>EQ 250MG BASE</u>	N64081 001 SEP 16, 1996
<u>AB</u>		<u>EQ 500MG BASE</u>	N64081 002 SEP 16, 1996
<u>AB</u>	MARSAM	<u>EQ 250MG BASE</u>	N64148 001 MAY 23, 1996
<u>AB</u>		<u>EQ 500MG BASE</u>	N64148 002 MAY 23, 1996
<u>AB</u>	NOVOPHARM	<u>EQ 250MG BASE</u>	N64145 001 JUN 24, 1996
<u>AB</u>		<u>EQ 500MG BASE</u>	N64145 002 JUN 24, 1996

TABLET, EXTENDED RELEASE; ORAL
CECLOR CD

+	LILLY	EQ 375MG BASE	N50673 001 JUN 28, 1996
+		EQ 500MG BASE	N50673 002 JUN 28, 1996

CEFAZOLIN SODIUM

INJECTABLE; INJECTION

ANCEP IN DEXTROSE 5% IN PLASTIC CONTAINER			
*	BAXTER	EQ 10MG BASE/ML	N50566 003 JUN 08, 1983
*		EQ 20MG BASE/ML	N50566 004 JUN 08, 1983
@	BAXTER HLTHCARE	EQ 10MG BASE/ML	N50566 003 JUN 08, 1983
@		EQ 20MG BASE/ML	N50566 004 JUN 08, 1983

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

CEFTAZIDIME (ARGININE FORMULATION)

INJECTABLE; INJECTION

CEPTAZ

* GLAXO WELLCOME 500MG/VIAL

N50646 001

SEP 27, 1990

@ 500MG/VIAL

N50646 001

SEP 27, 1990

CEFTAZIDIME SODIUM

INJECTABLE; INJECTION

CEFTAZIDIME SODIUM IN PLASTIC CONTAINER

AP BAXTER EQ 10MG BASE/ML

N63221 001

APR 29, 1993

+ BAXTER HLTHCARE EQ 10MG BASE/ML

N63221 001

APR 29, 1993

FORTAZ IN PLASTIC CONTAINER

AP * GLAXO WELLCOME EQ 10MG BASE/ML

N50634 001

APR 28, 1989

@ EQ 10MG BASE/ML

N50634 001

APR 28, 1989

CEFTIZOXIME SODIUM

INJECTABLE; INJECTION

CEFIZOX IN DEXTROSE 5% IN PLASTIC CONTAINER

* FUJISAWA EQ 20MG BASE/ML

N50589 003

APR 13, 1995

* EQ 40MG BASE/ML

N50589 004

APR 13, 1995

CEFIZOX IN PLASTIC CONTAINER

+ FUJISAWA EQ 20MG BASE/ML

N50589 003

APR 13, 1995

+ EQ 40MG BASE/ML

N50589 004

APR 13, 1995

CEFTRIAXONE SODIUM

INJECTABLE; INJECTION

ROCEPHIN

ROCHE

EQ 500MG BASE/VIAL

N62654 001

APR 30, 1987

@ EQ 500MG BASE/VIAL

N62654 001

APR 30, 1987

CEPHALEXIN

CAPSULE; ORAL

CEPANEX

AB APOTHECON

EQ 250MG BASE

N63063 001

SEP 29, 1989

AB EQ 500MG BASE

N63063 002

SEP 29, 1989

AB CEPHALEXIN
APOTHECON

EQ 250MG BASE

N63063 001

SEP 29, 1989

AB EQ 500MG BASE

N63063 002

SEP 29, 1989

AB YOSHITOMI EQ 250MG BASE

N62872 001

JUN 20, 1988

AB EQ 500MG BASE

N62871 001

JUL 05, 1988

@ EQ 250MG BASE

N62872 001

JUN 20, 1988

@ EQ 500MG BASE

N62871 001

JUL 05, 1988

TABLET; ORAL

CEPHALEXIN

AB BARR

EQ 250MG BASE

N62826 001

AUG 17, 1987

AB EQ 500MG BASE

N62827 001

AUG 17, 1987

@ EQ 250MG BASE

N62826 001

AUG 17, 1987

@ EQ 500MG BASE

N62827 001

AUG 17, 1987

CEPHALEXIN HYDROCHLORIDE

TABLET; ORAL

KEFTAB

* LILLY

EQ 250MG BASE

N50614 001

OCT 29, 1987

* EQ 333MG BASE

N50614 003

MAY 16, 1988

@ EQ 250MG BASE

N50614 001

OCT 29, 1987

@ EQ 333MG BASE

N50614 003

MAY 16, 1988

CEPHALOTHIN SODIUM

INJECTABLE; INJECTION

CEPHALOTHIN SODIUM W/ DEXTROSE IN PLASTIC CONTAINER

* BAXTER	EQ 20MG BASE/ML	N62422 003
		JAN 31, 1984
	EQ 20MG BASE/ML	N62422 005
		JUL 16, 1991
	EQ 20MG BASE/ML	N62730 001
		MAR 05, 1987
*	EQ 40MG BASE/ML	N62422 004
		JAN 31, 1984
	EQ 40MG BASE/ML	N62422 006
		JUL 16, 1991
	EQ 40MG BASE/ML	N62730 002
		MAR 05, 1987
@ BAXTER HLTHCARE	EQ 20MG BASE/ML	N62422 003
@	EQ 20MG BASE/ML	JAN 31, 1984
		N62422 005
@	EQ 20MG BASE/ML	JUL 16, 1991
		N62730 001
@	EQ 40MG BASE/ML	MAR 05, 1987
		N62422 004
@	EQ 40MG BASE/ML	JAN 31, 1984
		N62422 006
@	EQ 40MG BASE/ML	JUL 16, 1991
		N62730 002
		MAR 05, 1987
<u>CEPHALOTHIN SODIUM W/ SODIUM CHLORIDE IN PLASTIC CONTAINER</u>		
* BAXTER	EQ 20MG BASE/ML	N62422 001
		JAN 31, 1984
*	EQ 40MG BASE/ML	N62422 002
		JAN 31, 1984
@ BAXTER HLTHCARE	EQ 20MG BASE/ML	N62422 001
@	EQ 40MG BASE/ML	JAN 31, 1984
		N62422 002
		JAN 31, 1984

CEPHRADINE

CAPSULE; ORAL

CEPHRADINE

AB BARR	250MG	N62850 001
		APR 22, 1988
AB	500MG	N62851 001
		APR 22, 1988
@	250MG	N62850 001
		APR 22, 1988

CEPHRADINE

CAPSULE; ORAL

CEPHRADINE

@ BARR	500MG	N62851 001
		APR 22, 1988

POWDER FOR RECONSTITUTION; ORAL

CEPHRADINE

AB BARR	125MG/5ML	N62858 001
		MAY 19, 1988
AB	250MG/5ML	N62859 001
		MAY 19, 1988
@	125MG/5ML	N62858 001
		MAY 19, 1988
@	250MG/5ML	N62859 001
		MAY 19, 1988

CETIRIZINE HYDROCHLORIDE

SYRUP; ORAL

ZYRTEC

+ PFIZER	5MG/5ML	N20346 001
		SEP 27, 1996

CHLORDIAZEPOXIDE HYDROCHLORIDE

CAPSULE; ORAL

CHLORDIAZEPOXIDE HCL

AB HALSEY	5MG	N85340 001
AB	10MG	N85339 001
AB	25MG	N84685 001
@	5MG	N85340 001
@	10MG	N85339 001
@	25MG	N84685 001

CHLORHEXIDINE GLUCONATE

SOLUTION; DENTAL

CHLORHEXIDINE GLUCONATE

AT HI TECH PHARMA	0.12%	N74356 001
		MAY 07, 1996

CHLORPROTHIXENE

TABLET, ORAL

TARACTAN

@ ROCHE

@

50MG

N12486 003

100MG

N12486 001

CHLORTHALIDONE

TABLET, ORAL

CHLORTHALIDONE

BARR

AB25MG

N87292 001

AB50MG

N87293 001

@

25MG

N87292 001

@

50MG

N87293 001

AB

SUPERPHARM

50MG

N87247 001

@

50MG

FEB 09, 1983

N87247 001

FEB 09, 1983

CHLORZOXAZONE

TABLET, ORAL

CHLORZOXAZONE

CHELSEA LABS

AA500MG

N40137 001

AUG 09, 1996

CHOLESTYRAMINE

POWDER, ORAL

CHOLESTYRAMINE

COPLEY PHARM

ABEQ 4GM RESIN/PACKET

N74554 001

OCT 02, 1996

ABEQ 4GM RESIN/SCOOPFUL

N74554 002

OCT 02, 1996

AB

EON LABS MFG

EQ 4GM RESIN/PACKET

N74557 001

AUG 15, 1996

ABEQ 4GM RESIN/SCOOPFUL

N74557 002

AUG 15, 1996

CHOLESTYRAMINE LIGHTAB

EON LABS MFG

EQ 4GM RESIN/PACKET

N74558 001

AUG 15, 1996

ABEQ 4GM RESIN/SCOOPFUL

N74558 002

AUG 15, 1996

LOCHOLESTAB

EON LABS

EQ 4GM RESIN/PACKET

N74557 001

AUG 15, 1996

CHOLESTYRAMINE

POWDER, ORAL

LOCHOLEST

EON LABS

ABABAB

@ EON LABS MFG

@

LOCHOLEST LIGHT

EON LABS

ABABABAB

@ EON LABS MFG

@

PREVALITE

UPSHER SMITH

ABQUESTRAN

+ BRISTOL MYERS

ABABQUESTRAN LIGHT

* BRISTOL MYERS

ABABABCHROMIC CHLORIDE

INJECTABLE, INJECTION

CHROMIC CHLORIDE

FUJISAWA

APEQ 4GM RESIN/SCOOPFUL

N74557 002

AUG 15, 1996

EQ 4GM RESIN/PACKET

N74561 001

AUG 15, 1996

EQ 4GM RESIN/SCOOPFUL

N74561 002

AUG 15, 1996

EQ 4GM RESIN/PACKET

N74561 001

AUG 15, 1996

EQ 4GM RESIN/SCOOPFUL

N74561 002

AUG 15, 1996

EQ 4GM RESIN/PACKET

N74558 001

AUG 15, 1996

EQ 4GM RESIN/SCOOPFUL

N74558 002

AUG 15, 1996

EQ 4GM RESIN/PACKET

N74562 001

AUG 15, 1996

EQ 4GM RESIN/SCOOPFUL

N74562 002

AUG 15, 1996

EQ 4GM RESIN/PACKET

N74562 001

AUG 15, 1996

EQ 4GM RESIN/SCOOPFUL

N74562 002

AUG 15, 1996

EQ 4GM RESIN/PACKET

N73263 001

FEB 22, 1996

EQ 4GM RESIN/PACKET

N16640 001

EQ 4GM RESIN/SCOOPFUL

N16640 003

EQ 4GM RESIN/PACKET

N19669 001

DEC 05, 1988

EQ 4GM RESIN/PACKET

N19669 001

DEC 05, 1988

EQ 4GM RESIN/SCOOPFUL

N19669 003

DEC 05, 1988

EQ 0.004MG CHROMIUM/ML

N19271 001

MAY 05, 1987

CHROMIC CHLORIDE

INJECTABLE; INJECTION

CHROMIC CHLORIDE

@ FUJISAWA

EQ 0.004MG CHROMIUM/ML N19271 001
MAY 05, 1987CHROMIC CHLORIDE IN PLASTIC CONTAINER

AP * ABBOTT

EQ 0.004MG CHROMIUM/ML N18961 001
JUN 26, 1986

+

EQ 0.004MG CHROMIUM/ML N18961 001
JUN 26, 1986CIDOFIVIRINJECTABLE; INJECTION
VISTIDE

+ GILEAD

EQ 75MG BASE/ML N20638 001
JUN 26, 1996CIMETIDINE

TABLET; ORAL

CIMETIDINE

AB DANBURY PHARMA

200MG

N74349 001
AUG 30, 1996

AB 300MG

N74349 002
AUG 30, 1996

AB 400MG

N74349 003
AUG 30, 1996

AB 800MG

N74316 001
FEB 28, 1996

AB INVAMED 200MG

N74506 001
JAN 24, 1996

AB 300MG

N74506 002
JAN 24, 1996

AB 400MG

N74506 003
JAN 24, 1996

AB 800MG

N74506 004
JAN 24, 1996

AB ZENITH GOLDLINE 200MG

N74401 001
MAY 30, 1995

AB 300MG

N74401 002
MAY 30, 1995

AB 400MG

N74401 003
MAY 30, 1995

AB ZENITH LABS 200MG

N74401 001
MAY 30, 1995CIMETIDINE

TABLET; ORAL

CIMETIDINE

AB ZENITH LABS

300MG

N74401 002

AB 400MG

MAY 30, 1995

N74401 003

MAY 30, 1995

CIMETIDINE HYDROCHLORIDE

INJECTABLE; INJECTION

CIMETIDINE HCL

AP MOVA

EQ 300MG BASE/2ML

N74428 001

APR 25, 1996

SOLUTION; ORAL

CIMETIDINE HCL

AA LEMMON

EQ 300MG BASE/5ML

N74610 001

SEP 26, 1996

TAGAMET

AA SMITHKLINE BEECHAM

EQ 300MG BASE/5ML

N17924 001

AA + EQ 300MG BASE/5ML

N17924 001

CIPROFLOXACIN HYDROCHLORIDE

TABLET; ORAL

CIPRO

BAYER

EQ 100MG BASE

N19537 001

APR 08, 1996

CLINDAMYCIN PHOSPHATE

SOLUTION; TOPICAL

CLINDAMYCIN PHOSPHATE

AT STIEFEL

EQ 1% BASE

N64108 001

SEP 27, 1996

SWAB; TOPICAL

CLEOCIN

AT PHARMACIA AND UPJOHN EQ 1% BASE

N50537 002

FEB 22, 1994

CLINDETS

AT STIEFEL

EQ 1% BASE

N64136 001

SEP 30, 1996

CLOBETASOL PROPIONATE

CREAM; TOPICAL

CLOBETASOL PROPIONATE

AB FOUGERA 0.05%

AB TARO 0.05%

OINTMENT; TOPICAL

CLOBETASOL PROPIONATE

AB FOUGERA 0.05%

AB TARO 0.05%

CLOCORTOLONE PIVALATE

CREAM; TOPICAL

CLODERM

> ADD > + CTR LABS 0.1%

> DLT > + HERMAL PHARM 0.1%

N74392 001
SEP 30, 1996
N74249 001
JUL 08, 1996

N74407 001
FEB 23, 1996
N74248 001
JUL 12, 1996

N17765 001
N17765 001

CLOMIPRAMINE HYDROCHLORIDE

CAPSULE; ORAL

ANAFRANIL

AB CIBA GEIGY 25MG

AB + 50MG

AB 75MG

CLOMIPRAMINE HCL

AB CIRCA 25MG

AB 50MG

AB 75MG

AB GENEVA PHARMS 25MG

AB 50MG

AB 75MG

N19906 001
DEC 29, 1989
N19906 002
DEC 29, 1989
N19906 003
DEC 29, 1989

N74600 001
NOV 27, 1996
N74600 002
NOV 27, 1996
N74600 003
NOV 27, 1996
N74364 001
MAR 29, 1996
N74364 002
MAR 29, 1996
N74364 003
MAR 29, 1996

CLOMIPRAMINE HYDROCHLORIDE

CAPSULE; ORAL

CLOMIPRAMINE HCL

AB TARO 25MG

AB 50MG

AB 75MG

CLONAZEPAM

TABLET; ORAL

CLONAZEPAM

AB LEMMON 0.5MG

AB 1MG

AB 2MG

AB PUREPAC PHARM 0.5MG

AB 1MG

AB 2MG

KLONOPIN

AB ROCHE 0.5MG

AB 1MG

AB + 2MG

CLONIDINE HYDROCHLORIDE

INJECTABLE; INJECTION

DURACLON

+ FUJISAWA 0.1MG/ML

TABLET; ORAL

CLONIDINE HCL

AB BARE 0.2MG

AB 0.3MG

@ 0.2MG

N74694 001
DEC 31, 1996
N74694 002
DEC 31, 1996
N74694 003
DEC 31, 1996

N74569 001
SEP 10, 1996
N74569 002
SEP 10, 1996
N74569 003
SEP 10, 1996
N74869 001
OCT 31, 1996
N74869 002
OCT 31, 1996
N74869 003
OCT 31, 1996

N17533 001
N17533 002
N17533 003

N20615 001
OCT 02, 1996

N70924 001
SEP 04, 1987
N70923 001
SEP 04, 1987
N70924 001
SEP 04, 1987

CLONIDINE HYDROCHLORIDE

<u>TABLET; ORAL</u>			
	<u>CLONIDINE HCL</u>		
	@ BARR	0.3MG	N70923 001
			SEP 04, 1987
<u>AB</u>	WARNER CHILCOTT	<u>0.1MG</u>	N72138 001
			JUN 13, 1988
<u>AB</u>		<u>0.2MG</u>	N72139 001
			JUN 13, 1988
<u>AB</u>		<u>0.3MG</u>	N72140 001
			JUN 13, 1988
	@	0.1MG	N72138 001
	@	0.2MG	N72139 001
	@	0.3MG	N72140 001
			JUN 13, 1988

CLOTTRIMAZOLE

<u>SOLUTION; TOPICAL</u>			
	<u>CLOTTRIMAZOLE</u>		
<u>AT</u>	LEMMON	<u>1%</u>	N73306 001
			FEB 28, 1995
<u>AT</u>	TARO	<u>1%</u>	N74580 001
			JUL 29, 1996
<u>AT</u>	TEVA	<u>1%</u>	N73306 001
			FEB 28, 1995

CLOZAPINE

<u>TABLET; ORAL</u>			
	<u>CLOZAPINE</u>		
<u>AB</u>	CREIGHTON	<u>25MG</u>	N74546 001
			AUG 30, 1996
<u>AB</u>		<u>100MG</u>	N74546 002
			AUG 30, 1996
	<u>CLOZARIL</u>		
<u>AB</u>	+ SANDOZ	<u>25MG</u>	N19758 001
			SEP 26, 1989
<u>AB</u>		<u>100MG</u>	N19758 002
			SEP 26, 1989

CODEINE PHOSPHATE; PHENYLEPHRINE HYDROCHLORIDE; PROMETHAZINE HYDROCHLORIDE

<u>SYRUP; ORAL</u>			
	<u>PHERAZINE VC W/ CODEINE</u>		
<u>AA</u>	HALSEY	<u>10MG/5ML; 5MG/5ML;</u> <u>6.25MG/5ML</u>	N88870 001
			MAR 02, 1987
	@	10MG/5ML; 5MG/5ML; 6.25MG/5ML	N88870 001
			MAR 02, 1987
	<u>PROMETHAZINE VC W/ CODEINE</u>		
<u>AA</u>	CENCI	<u>10MG/5ML; 5MG/5ML;</u> <u>6.25MG/5ML</u>	N88816 001
			NOV 22, 1985
	@	10MG/5ML; 5MG/5ML; 6.25MG/5ML	N88816 001
			NOV 22, 1985

CODEINE PHOSPHATE; PROMETHAZINE HYDROCHLORIDE

<u>SYRUP; ORAL</u>			
	<u>PROMETHAZINE W/ CODEINE</u>		
<u>AA</u>	CENCI	<u>10MG/5ML; 6.25MG/5ML</u>	N88814 001
			NOV 22, 1985
	@	10MG/5ML; 6.25MG/5ML	N88814 001
			NOV 22, 1985

CODEINE PHOSPHATE; PSEUDOEPHEDRINE HYDROCHLORIDE; TRIPROLIDINE HYDROCHLORIDE

<u>SYRUP; ORAL</u>			
	<u>TRIPROLIDINE AND PSEUDOEPHEDRINE HYDROCHLORIDES W/ CODEINE</u>		
<u>AA</u>	CENCI	<u>10MG/5ML; 30MG/5ML;</u> <u>1.25MG/5ML</u>	N89018 001
			JUL 23, 1986
	@	10MG/5ML; 30MG/5ML; 1.25MG/5ML	N89018 001
			JUL 23, 1986

CORTICORELIN OVINE TRIFLUTATE

<u>INJECTABLE; INJECTION</u>			
	ACTHREL		
	+ FERRING LABS	EQ 0.1MG BASE/VIAL	N20162 001
			MAY 23, 1996

CROMOLYN SODIUMCAPSULE; ORAL
GASTROCROM

* FISOXS	100MG	N19188 001
		DEC 22, 1989
+ MEDEVA PHARMS	100MG	N19188 001
		DEC 22, 1989

CONCENTRATE; ORAL
GASTROCROM

* FISOXS	100MG/5ML	N20479 001
		FEB 29, 1996
+ MEDEVA PHARMS	100MG/5ML	N20479 001
		FEB 29, 1996

SOLUTION/DROPS; OPHTHALMIC

CROLOM

AT	BAUSCH AND LOMB	4%	N74443 001
			JAN 30, 1995
+		4%	N74443 001
			JAN 30, 1995

OPTICROM

AT	* FISOXS	4%	N18155 001
			OCT 03, 1984
@		4%	N18155 001
			OCT 03, 1984

CYANOCOBALAMINGEL, METERED; NASAL
NASCOBAL

+ NASTECH	0.5MG/INH	N19722 001
		NOV 05, 1996

INJECTABLE; INJECTION

BETALIN 12

AP	LILLY	0.1MG/ML	N80855 001
AP		1MG/ML	N80855 002
@		0.1MG/ML	N80855 001
@		1MG/ML	N80855 002

CYANOCOBALAMIN

AP	DELL LABS	0.1MG/ML	N80689 002
AP		1MG/ML	N80689 003
		0.03MG/ML	N80689 001
@		0.03MG/ML	N80689 001
@		0.1MG/ML	N80689 002
@		1MG/ML	N80689 003

CYCLOSPORINE

CAPSULE; ORAL

BP	NEORAL	25MG	N50715 001
	SANDOZ		JUL 14, 1995
BP		50MG	N50715 003
			JUL 14, 1995
BP	*	100MG	N50715 002
			JUL 14, 1995

BP SANDIMMUNE
SANDOZ

BP		25MG	N50625 001
			MAR 02, 1990
BP		50MG	N50625 003
			NOV 23, 1992
BP	*	100MG	N50625 002
			MAR 02, 1990
		25MG	N50625 001
			MAR 02, 1990
		50MG	N50625 003
			NOV 23, 1992
+		100MG	N50625 002
			MAR 02, 1990

CAPSULE, MICROEMULSION; ORAL

	NEORAL	25MG	N50715 001
	SANDOZ		JUL 14, 1995
		50MG	N50715 003
			JUL 14, 1995
+		100MG	N50715 002
			JUL 14, 1995

SOLUTION; ORAL

BP	NEORAL	100MG/ML	N50716 001
	SANDOZ		JUL 14, 1995

BP SANDIMMUNE
SANDOZ

BP	*	100MG/ML	N50574 001
			NOV 14, 1983
+		100MG/ML	N50574 001
			NOV 14, 1983

SOLUTION, MICROEMULSION; ORAL

	NEORAL		
+	SANDOZ	100MG/ML	N50716 001
			JUL 14, 1995

CYCLOTHIAZIDE

TABLET; ORAL
ANHYDRON
* LILLY
@

2MG
2MG

N13157 002
N13157 002

CYPROHEPTADINE HYDROCHLORIDE

SYRUP; ORAL
CYPROHEPTADINE HCL
AA RALSEY

2MG/5ML

N89199 001
JUL 03, 1986
N89199 001
JUL 03, 1986

2MG/5ML

DALTEPARIN SODIUM

INJECTABLE; INJECTION
FRAGMIN

+ PHARMACIA AND UPJOHN 5,000 IU/0.2ML

N20287 003
MAR 18, 1996

> ADD > DANAPAROID SODIUM

> ADD > INJECTABLE; INJECTION

> ADD > ORGARAN

> ADD > + ORGANON

> ADD >

750 UNITS/0.6ML

N20430 001
DEC 24, 1996

DANAZOL

CAPSULE; ORAL

AB DANAZOL
BARR

200MG

N74582 001
AUG 09, 1996

AB DANOCRINE

+ SANOFI WINTHROP

200MG

N17557 002

DAUNORUBICIN CITRATE

INJECTABLE, LIPOSOMAL; INJECTION

DAUNOXOME

+ NEXSTAR

EQ 2MG BASE/ML

N50704 002
APR 08, 1996

DESIPRAMINE HYDROCHLORIDE

TABLET; ORAL
DESIPRAMINE HCL

AB EON LABS MFG

10MG

AB

150MG

N74430 001
FEB 09, 1996
N74430 002
FEB 09, 1996

DESMOPRESSIN ACETATE

SPRAY, METERED; NASAL
STIMATE

* ARMOUR PHARM

0.15MG/INH

+ CENTEON

0.15MG/INH

N20355 001
MAR 07, 1994
N20355 001
MAR 07, 1994

DESOGESTREL; ETHINYL ESTRADIOL

TABLET; ORAL-21

ORTHO-CEPT

JOHNSON RW

0.15MG; 0.03MG

+

0.15MG; 0.03MG

N20301 001
DEC 14, 1992
N20301 001
DEC 14, 1992

DESOXIMETASONE

OINTMENT; TOPICAL

DESOXIMETASONE

AB TARO

0.25%

N74286 001
JUN 07, 1996

AB TOPICORT

+ HOECHST MARION RSSL

0.25%

N18763 001
SEP 30, 1983

DEXAMETHASONE

CONCENTRATE; ORAL

DEXAMETHASONE INTENSOL

+ ROXANE

1MG/ML

N88252 001
SEP 01, 1983

DEXAMETHASONE

SOLUTION; ORAL

DEXAMETHASONE INTENSOL

ROXANE

0.5MG/0.5ML

N88252 001

SEP 01, 1983

DEXAMETHASONE; NEOMYCIN SULFATE; POLYMYXIN B SULFATE

OINTMENT; OPHTHALMIC

DEXACIDIN

AT CIBA

0.1%;EQ 3.5MG BASE/GM;
10,000 UNITS/GM

N62566 001

FEB 22, 1985

AT IOLAB

0.1%;EQ 3.5MG BASE/GM;
10,000 UNITS/GM

N62566 001

FEB 22, 1985

SUSPENSION/DROPS; OPHTHALMIC

DEXACIDIN

AT CIBA

0.1%;EQ 3.5MG BASE/ML;
10,000 UNITS/ML

N62544 001

OCT 29, 1984

AT IOLAB

0.1%;EQ 3.5MG BASE/ML;
10,000 UNITS/ML

N62544 001

OCT 29, 1984

DEXAMETHASONE SODIUM PHOSPHATE

SOLUTION/DROPS; OPHTHALMIC, OTIC

DEXAMETHASONE SODIUM PHOSPHATE

AT BAUSCH AND LOMB

EQ 0.1% PHOSPHATE

N40069 001

JUL 26, 1996

DEXFENFLURAMINE HYDROCHLORIDE

CAPSULE; ORAL

REDUX

+ INTERNEURON

15MG

N20344 001

APR 29, 1996

DEXTROAMPHETAMINE SULFATE

TABLET; ORAL

DEXTROAMPHETAMINE SULFATE

AA HALSEY

10MG

N83930 001

DEXTROAMPHETAMINE SULFATE

TABLET; ORAL

DEXTROAMPHETAMINE SULFATE

@ HALSEY

10MG

N83930 001

AA MM MAST

5MG

N86521 001

@

5MG

N86521 001

AA * REXAR

10MG

N84051 002

+

10MG

N84051 002

DEXTROMETHORPHAN HYDROBROMIDE; PROMETHAZINE HYDROCHLORIDE

SYRUP; ORAL

PROMETHAZINE DM

AA HALSEY

15MG/5ML; 6.25MG/5ML

N88913 001

@

15MG/5ML; 6.25MG/5ML

MAR 02, 1987

N88913 001

MAR 02, 1987

PROMETHAZINE HCL AND DEXTROMETHORPHAN HYDROBROMIDE

AA HI TECH PHARMA

15MG/5ML; 6.25MG/5ML

N40027 001

JUL 31, 1996

DEXTROSE

INJECTABLE; INJECTION

DEXTROSE 60% IN PLASTIC CONTAINER

AP BAXTER

60GM/100ML

N20047 002

JUL 02, 1991

@ BAXTER HLTHCARE

60GM/100ML

N20047 002

JUL 02, 1991

DEXTROSE; SODIUM CHLORIDE

INJECTABLE; INJECTION

DEXTROSE 10% AND SODIUM CHLORIDE 0.2% IN PLASTIC CONTAINER

MCGAW

10GM/100ML; 200MG/100ML

N18386 001

@

10GM/100ML; 200MG/100ML

N18386 001

DEXTROSE 10% AND SODIUM CHLORIDE 0.45% IN PLASTIC CONTAINER

MCGAW

10GM/100ML; 450MG/100ML

N18229 001

@

10GM/100ML; 450MG/100ML

N18229 001

DEXTROSE 2.5% AND SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER

MCGAW

2.5GM/100ML; 900MG/100ML

N18376 001

@

2.5GM/100ML; 900MG/100ML

N18376 001

DIATRIZOATE MEGLUMINE; DIATRIZOATE SODIUM

INJECTABLE; INJECTION

<u>AP</u>	<u>MD-76</u> <u>MALLINCKRODT</u>	<u>65%;10%</u>	N19292 001 SEP 29, 1989
<u>AP</u>	<u>MD-76R</u> + <u>MALLINCKRODT MEDCL</u>	<u>66%;10%</u>	N19292 001 SEP 29, 1989

DIATRIZOATE SODIUM

SOLUTION; URETERAL

<u>AT</u>	<u>HYPAAQUE SODIUM 20%</u> <u>NYCOMED</u>	<u>20%</u> 20%	N09561 002 N09561 002
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DIAZEPAM

TABLET; ORAL

<u>AB</u>	<u>DIAZEPAM</u> <u>ZENITH GOLDLINE</u>	<u>2MG</u>	N71307 001 DEC 10, 1986
<u>AB</u>		<u>5MG</u>	N71321 001 DEC 10, 1986
<u>AB</u>		<u>10MG</u>	N70362 001 SEP 04, 1985
<u>AB</u>		<u>10MG</u>	N71322 001 DEC 10, 1986
<u>AB</u>	<u>ZENITH LABS</u>	<u>2MG</u>	N71307 001 DEC 10, 1986
<u>AB</u>		<u>5MG</u>	N71321 001 DEC 10, 1986
<u>AB</u>		<u>10MG</u>	N70362 001 SEP 04, 1985
<u>AB</u>		<u>10MG</u>	N71322 001 DEC 10, 1986

DICLOFENAC SODIUM

TABLET, DELAYED RELEASE; ORAL

<u>AB</u>	<u>DICLOFENAC SODIUM</u> <u>NOVOPHARM</u>	<u>75MG</u>	N74390 001 AUG 15, 1996
<u>AB</u>	<u>PUREPAC PHARM</u>	<u>50MG</u>	N74514 001 MAR 26, 1996

DICLOFENAC SODIUM

TABLET, DELAYED RELEASE; ORAL

<u>AB</u>	<u>DICLOFENAC SODIUM</u> <u>PUREPAC PHARM</u>	<u>75MG</u>	N74514 002 MAR 26, 1996
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TABLET, EXTENDED RELEASE; ORAL

	<u>VOLTAREN-XR</u> + <u>GEIGY</u>	<u>100MG</u>	N20254 001 MAR 08, 1996
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DICLOXACILLIN SODIUM

CAPSULE; ORAL

<u>AB</u>	<u>DICLOXACILLIN SODIUM</u> <u>APOTHECON</u>	<u>EQ 250MG BASE</u>	N61454 001
<u>AB</u>	+	<u>EQ 250MG BASE</u>	N61454 001
<u>AB</u>		<u>EQ 500MG BASE</u>	N61454 003
<u>AB</u>	+	<u>EQ 500MG BASE</u>	N61454 003
<u>AB</u>	<u>* PATHOCIL</u> <u>* WYETH AYERST</u>	<u>EQ 250MG BASE</u>	N50011 002
<u>AB</u>		<u>EQ 250MG BASE</u>	N50011 002
<u>AB</u>	*	<u>EQ 500MG BASE</u>	N50011 003
			MAR 28, 1983
<u>AB</u>		<u>EQ 500MG BASE</u>	N50011 003
			MAR 28, 1983

POWDER FOR RECONSTITUTION; ORAL

<u>AB</u>	<u>DICLOXACILLIN SODIUM</u> <u>APOTHECON</u>	<u>EQ 62.5MG BASE/5ML</u>	N61455 001
<u>AB</u>	+	<u>EQ 62.5MG BASE/5ML</u>	N61455 001
<u>AB</u>	<u>* PATHOCIL</u> <u>* WYETH AYERST</u>	<u>EQ 62.5MG BASE/5ML</u>	N50092 001
<u>AB</u>		<u>EQ 62.5MG BASE/5ML</u>	N50092 001

DICYCLOMINE HYDROCHLORIDE

CAPSULE; ORAL

<u>AB</u>	<u>DICYCLOMINE HCL</u> <u>LANNETT</u>	<u>10MG</u>	N84285 001
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INJECTABLE; INJECTION

<u>AP</u>	<u>BENTYL PRESERVATIVE FREE</u> <u>HOECHST MARION RSSL</u>	<u>10MG/ML</u>	N08370 002 OCT 15, 1984
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DICYCLOMINE HYDROCHLORIDE

TABLET; ORAL
DICYCLOMINE HCL
AB WEST WARD 20MG N40161 001
OCT 01, 1996

DIETHYLCARBAMAZINE CITRATE

> DLT > TABLET; ORAL
> DLT > HETRAZAN
> DLT > LEDERLE 50MG N06459 001
> ADD > @ 50MG N06459 001

DIFLUNISAL

TABLET; ORAL
DIFLUNISAL
AB GENEVA PHARMS 500MG N74604 001
JUN 10, 1996
AB PUREPAC PHARM 250MG N74285 001
MAY 07, 1996
AB 500MG N74285 002
MAY 07, 1996

DIGOXIN

INJECTABLE; INJECTION
DIGOXIN
AP SANOFI WINTHROP 0.25MG/ML N40093 001
MAY 16, 1996
AP DIGOXIN PEDIATRIC
SANOFI WINTHROP 0.1MG/ML N40092 001
APR 25, 1996
LANOXIN
* GLAXO-WELLCOME 0.1MG/ML N09330 004
AP LANOXIN PEDIATRIC
+ GLAXO WELLCOME 0.1MG/ML N09330 004

DILTIAZEM HYDROCHLORIDE

INJECTABLE; INJECTION
CARDIZEM
AP + HOECHST MARION RSSL 5MG/ML N20027 001
OCT 24, 1991

DILTIAZEM HYDROCHLORIDE

INJECTABLE; INJECTION
DILTIAZEM HCL
AP BEDFORD 5MG/ML N74617 001
FEB 28, 1996

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
+ MERCK EQ 180MG HCL;5MG N20507 001
OCT 04, 1996

DIPHENHYDRAMINE HYDROCHLORIDE

CAPSULE; ORAL
DIPHENHYDRAMINE HCL
AA HALSEY 50MG N87914 001
JUN 04, 1984
@ 50MG N87914 001
JUN 04, 1984

ELIXIR; ORAL
BELIX
AA HALSEY 12.5MG/5ML N86586 001
OCT 03, 1983
@ 12.5MG/5ML N86586 001
OCT 03, 1983
DIBENIL
AA CENCEL 12.5MG/5ML N88304 001
DEC 16, 1983
@ 12.5MG/5ML N88304 001
DEC 16, 1983

DIPHENHYDRAMINE HYDROCHLORIDE

ELIXIR; ORAL
DIPHENHYDRAMINE HCL
AP CENCT 12.5MG/5ML N87941 001
 DEC 17, 1982
 @ 12.5MG/5ML N87941 001
 DEC 17, 1982

DIPYRIDAMOLE

INJECTABLE; INJECTION
DIPYRIDAMOLE
AP ELKINS SINN 5MG/ML N74521 001
 OCT 18, 1996
IV PERSANTINE
AP + BOEHRINGER INGELHEIM 5MG/ML N19817 001
 DEC 13, 1990

DISOPYRAMIDE PHOSPHATE

CAPSULE; ORAL
DISOPYRAMIDE PHOSPHATE
AB BARR EQ 100MG BASE N70351 001
 DEC 17, 1985
AB EQ 150MG BASE N70352 001
 DEC 17, 1985
 @ EQ 100MG BASE N70351 001
 DEC 17, 1985
 @ EQ 150MG BASE N70352 001
 DEC 17, 1985
 CAPSULE, EXTENDED RELEASE; ORAL
DISOPYRAMIDE PHOSPHATE
E* KV PHARM EQ 100MG BASE N71929 001
 AUG 13, 1988
 @ EQ 100MG BASE N71929 001
 AUG 19, 1988
NORPACE CR
AB SEARLE EQ 100MG BASE N18655 001
 JUL 20, 1982
 EQ 100MG BASE N18655 001
 JUL 20, 1982

DOBUTAMINE HYDROCHLORIDE

INJECTABLE; INJECTION
DOBUTAMINE HCL
AP ABBOTT EQ 1.25GM BASE/100ML N74634 001
 SEP 27, 1996
AP ELKINS SINN EQ 12.5MG BASE/ML N74381 001
 SEP 26, 1996

DOCETAXEL

INJECTABLE; INJECTION
 TAXOTERE
 + RHONE POULENC EQ 40MG BASE/ML N20449 001
 MAY 14, 1996

DONEPEZIL HYDROCHLORIDE

TABLET; ORAL
 ARICEPT
 EISAI 5MG N20690 002
 NOV 25, 1996
 + 10MG N20690 001
 NOV 25, 1996

DOPAMINE HYDROCHLORIDE

INJECTABLE; INJECTION
DOPAMINE HCL
AP SANOFI WINTHROP 40MG/ML N74403 001
 MAY 23, 1996

DOXYCYCLINE HYCLATE

TABLET; ORAL
DOXYCYCLINE HYCLATE
AB SUPERPHARM EQ 100MG BASE N62494 001
 FEB 20, 1985
 @ EQ 100MG BASE N62494 001
 FEB 20, 1985

DROPERIDOL; FENTANYL CITRATE

INJECTABLE; INJECTION

FENTANYL CITRATE AND DROPERIDOL

AP	ABBOTT	2.5MG/ML; EQ 0.05MG BASE/ML	N71982 001 MAY 04, 1988
AP	+	2.5MG/ML; EQ 0.05MG BASE/ML	N71982 001 MAY 04, 1988
AP	INNOVAR * JANSSEN	2.5MG/ML; EQ 0.05MG BASE/ML	N16049 001
	@	2.5MG/ML; EQ 0.05MG BASE/ML	N16049 001

EDROPHONIUM CHLORIDE

INJECTABLE; INJECTION

EDROPHONIUM CHLORIDE

AP	STERIS	10MG/ML	N40044 001 MAR 20, 1996
AP	<u>EDROPHONIUM CHLORIDE PRESERVATIVE FREE</u> STERIS	10MG/ML	N40043 001 MAR 20, 1996
AP	+	<u>TENSILON PRESERVATIVE FREE</u> ROCHE	10MG/ML N07959 002

> ADD > ENALAPRIL MALEATE; FELODIPINE> ADD > TABLET, EXTENDED RELEASE; ORAL> ADD > LEXXEL> ADD > + ASTRA MERCK 5MG;5MG> ADD > N20668 001
DEC 27, 1996ENCAINIDE HYDROCHLORIDE

CAPSULE; ORAL

ENKAID

@ BRISTOL 25MG

@ 35MG

BRISTOL MYERS SQUIBB 25MG

N18981 002
DEC 24, 1986
N18981 003
DEC 24, 1986
N18981 002
DEC 24, 1986

> DLT >> DLT >> ADD >> ADD >ENCAINIDE HYDROCHLORIDE

CAPSULE; ORAL

ENKAID

BRISTOL MYERS SQUIBB 35MG

N18981 003
DEC 24, 1986

EPINEPHRINE; LIDOCAINE HYDROCHLORIDE

INJECTABLE; INJECTION

LIDOCAINE HCL W/ EPINEPHRINE

AP	ABBOTT	0.01MG/ML; 1%	N83154 001
	@	0.01MG/ML; 1%	N83154 001

ERGOLOID MESYLATES

TABLET; ORAL

ERGOLOID MESYLATES

AE	BARR	1MG	N88891 001 NOV 01, 1985
	@	1MG	N88891 001 NOV 01, 1985

TABLET; SUBLINGUAL

ERGOLOID MESYLATES

AA	BARR	0.5MG	N87407 001
AA		1MG	N87552 001
	@	0.5MG	N87407 001
	@	1MG	N87552 001
AA	KV PHARM	0.5MG	N85899 001
AA		1MG	N85900 001
	@	0.5MG	N85899 001
	@	1MG	N85900 001

ERYTHROMYCIN

OINTMENT; OPHTHALMIC

ERYTHROMYCIN

AT	ADV REMEDIES	0.5%	N64030 001 JUL 18, 1996
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OINTMENT; TOPICAL

AKNE-MYCIN

+ CTR LABS 2%

N50584 001
JAN 10, 1985

ERYTHROMYCIN

OINTMENT; TOPICAL
 AKNE-MYCIN
 > DLT > * HERMAL PHARM 2% N50584 001
 > DLT > JAN 10, 1985

 SOLUTION; TOPICAL
 ERYTHRO-STATIN
 AT HI TECH PHARMA 2% N64101 001
 OCT 22, 1996

 SWAB; TOPICAL
 ERYCETTE
 AT + J AND J 2% N50594 001
 FEB 15, 1985
 AT * JOHNSON RW 2% N50594 001
 FEB 15, 1985
 AT ERYTHROMYCIN 2% N64126 001
 STIEFEL JUL 03, 1996
 AT 2% N64128 001
 JUL 03, 1996

ERYTHROMYCIN ESTOLATE

CAPSULE; ORAL
 ERYTHROMYCIN ESTOLATE
 AB BARR EQ 125MG BASE N62162 001
 @ EQ 125MG BASE N62162 001
 ILOSONE
 > DLT > AB * DISTA EQ 250MG BASE N61897 002
 > DLT > EQ 125MG BASE N61897 001
 > ADD > AB + LILLY EQ 250MG BASE N61897 002
 > ADD > EQ 125MG BASE N61897 001

 SUSPENSION; ORAL
 ERYTHROMYCIN ESTOLATE
 AB BARR EQ 125MG BASE/5ML N62169 001
 OCT 17, 1990
 AB EQ 250MG BASE/5ML N62169 002
 OCT 17, 1990
 @ EQ 125MG BASE/5ML N62169 001
 OCT 17, 1990
 @ EQ 250MG BASE/5ML N62169 002
 OCT 17, 1990
 ILOSONE
 > DLT > @ DISTA EQ 125MG BASE/5ML N61894 001

ERYTHROMYCIN ESTOLATE

SUSPENSION; ORAL
 ILOSONE
 > DLT > @ DISTA EQ 250MG BASE/5ML N61894 002
 AB LILLY EQ 125MG BASE/5ML N50010 001
 AB + EQ 250MG BASE/5ML N50010 002
 > ADD > @ EQ 125MG BASE/5ML N61894 001
 > ADD > @ EQ 250MG BASE/5ML N61894 002

 SUSPENSION/DROPS; ORAL
 ILOSONE
 > DLT > @ DISTA EQ 100MG BASE/ML N61894 003
 > ADD > + LILLY EQ 100MG BASE/ML N61894 003

 TABLET; ORAL
 ILOSONE
 > DLT > * DISTA EQ 500MG BASE N61896 001
 > ADD > + LILLY EQ 500MG BASE N61896 001

ERYTHROMYCIN LACTOBIONATE

INJECTABLE; INJECTION
 ERYTHROCIN
 AP ABBOTT EQ 500MG BASE/VIAL N62586 001
 JAN 04, 1988
 AP EQ 1GM BASE/VIAL N62586 002
 JAN 04, 1988
 @ EQ 500MG BASE/VIAL N62586 001
 JAN 04, 1988
 @ EQ 1GM BASE/VIAL N62586 002
 JAN 04, 1988

ESTRADIOL

FILM, EXTENDED RELEASE; TRANSDERMAL
 ALORA
 > ADD > BX THERATECH 0.05MG/24HR N20655 001
 > ADD > DEC 20, 1996
 > ADD > BX 0.075MG/24HR N20655 002
 > ADD > DEC 20, 1996
 > ADD > BX 0.1MG/24HR N20655 003
 > ADD > DEC 20, 1996
 AB ESTRADIOL 0.0375MG/24HR N20538 001
 MENOREST JUL 31, 1996

ESTRADIOL

FILM, EXTENDED RELEASE; TRANSDERMAL

ESTRADIOL

<u>AB</u>	MENOREST	<u>0.05MG/24HR</u>	N20538 003 JUL 31, 1996
<u>AB</u>		<u>0.075MG/24HR</u>	N20538 002 JUL 31, 1996
<u>AB</u>		<u>0.1MG/24HR</u>	N20538 004 JUL 31, 1996

> ADD >
> ADD >
> ADD >

	FEMPATCH		
	+ PARKE DAVIS	0.025MG/24HR	N20417 001 DEC 03, 1996

VIVELLE

<u>AB</u>	+ CIBA GEIGY	<u>0.0375MG/24HR</u>	N20323 001 OCT 28, 1994
<u>AB</u>	+	<u>0.05MG/24HR</u>	N20323 002 OCT 28, 1994
<u>AB</u>	+	<u>0.075MG/24HR</u>	N20323 003 OCT 28, 1994
<u>AB</u>	+	<u>0.1MG/24HR</u>	N20323 004 OCT 28, 1994

INSERT, EXTENDED RELEASE; VAGINAL
ESTRING

	+ PHARMACIA AND UPJOHN	0.0075MG/24HR	N20472 001 APR 26, 1996
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TABLET; ORAL

ESTRACE

<u>AB</u>	BRISTOL MYERS SQUIBB	<u>0.5MG</u>	N81295 001 JUN 30, 1993
<u>AB</u>		<u>1MG</u>	N84499 001
<u>AB</u>	+	<u>2MG</u>	N84500 001
<u>AB</u>	<u>ESTRADIOL</u>		
<u>AB</u>	WATSON LABS	<u>0.5MG</u>	N40114 003 MAR 14, 1996
<u>AB</u>		<u>1MG</u>	N40114 001 MAR 14, 1996
<u>AB</u>		<u>2MG</u>	N40114 002 MAR 14, 1996

ESTRONEINJECTABLE; INJECTION
ESTROGENIC SUBSTANCE

<u>BP</u>	* WYETH AYERST	2MG/ML	N83488 003
	@	2MG/ML	N83488 001

ESTRONE

INJECTABLE; INJECTION

NATURAL ESTROGENIC SUBSTANCE-ESTRONE

<u>BP</u>	* STERIS	2MG/ML	N85237 001 NOV 23, 1982
	+	2MG/ML	N85237 001 NOV 23, 1982

ESTROPIPATE

TABLET; ORAL

ESTROPIPATE

<u>AB</u>	BARR	<u>0.75MG</u>	N40135 001 NOV 27, 1996
<u>AB</u>		<u>1.5MG</u>	N40135 002 NOV 27, 1996
<u>AB</u>		<u>3MG</u>	N40135 003 NOV 27, 1996

ETHANOLAMINE OLEATE

INJECTABLE; INJECTION

ETHAMOLIN

	+ CYPROS	50MG/ML	N19357 001 DEC 22, 1988
<u>*</u>	<u>SPKU</u>	<u>50MG/ML</u>	N19357 001 DEC 22, 1988

ETHINYL ESTRADIOL; ETHYNODIOL DIACETATE

TABLET; ORAL-21

ETHYNODIOL DIACETATE AND ETHINYL ESTRADIOL 1/35-21

<u>AB</u>	WATSON LABS	<u>0.035MG; 1MG</u>	N72720 001 DEC 30, 1991
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ETHYNODIOL DIACETATE AND ETHINYL ESTRADIOL 1/50-21

<u>AB</u>	WATSON LABS	<u>0.05MG; 1MG</u>	N72722 001 DEC 30, 1991
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ZOVIA 1/35E-21

<u>AB</u>	WATSON LABS	<u>0.035MG; 1MG</u>	N72720 001 DEC 30, 1991
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ZOVIA 1/50E-21

<u>AB</u>	WATSON LABS	<u>0.05MG; 1MG</u>	N72722 001 DEC 30, 1991
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ETHINYL ESTRADIOL; ETHYNODIOL DIACETATE

TABLET; ORAL-28			
<u>ETHYNODIOL DIACETATE AND ETHINYL ESTRADIOL 1/35-28</u>			
<u>AB</u>	<u>WATSON LABS</u>	<u>0.035MG;1MG</u>	<u>N72721 001</u>
			DEC 30, 1991
<u>ETHYNODIOL DIACETATE AND ETHINYL ESTRADIOL 1/50-28</u>			
<u>AB</u>	<u>WATSON LABS</u>	<u>0.05MG;1MG</u>	<u>N72723 001</u>
			DEC 30, 1991
<u>AB</u>	<u>ZOVIA 1/35E-28</u>	<u>0.035MG;1MG</u>	<u>N72721 001</u>
	<u>WATSON LABS</u>		DEC 30, 1991
<u>AB</u>	<u>ZOVIA 1/50E-28</u>	<u>0.05MG;1MG</u>	<u>N72723 001</u>
	<u>WATSON LABS</u>		DEC 30, 1991

ETHINYL ESTRADIOL; FERROUS FUMARATE; NORETHINDRONE ACETATE

TABLET; ORAL-28			
<u>LOESTRIN FE 1.5/30</u>			
*	<u>PARKE DAVIS</u>	<u>0.03MG;75MG;1.5MG</u>	<u>N17355 001</u>
<u>LOESTRIN FE 1/20</u>			
*	<u>PARKE DAVIS</u>	<u>0.02MG;75MG;1MG</u>	<u>N17354 001</u>

ETHINYL ESTRADIOL; NORETHINDRONE

TABLET; ORAL-21			
<u>NORETHINDRONE AND ETHINYL ESTRADIOL (7/14)</u>			
<u>AB</u>	<u>WATSON LABS</u>	<u>0.035MG;0.035MG;0.5MG;1MG</u>	<u>N71041 001</u>
			SEP 24, 1991
<u>AB</u>	<u>+</u>	<u>0.035MG;0.035MG;0.5MG;1MG</u>	<u>N71041 001</u>
			SEP 24, 1991
<u>ORTHO-NOVUM 7/14-21</u>			
<u>AB</u>	<u>* JOHNSON RW</u>	<u>0.035MG;0.035MG;0.5MG;1MG</u>	<u>N19004 001</u>
			APR 04, 1984
	<u>@</u>	<u>0.035MG;0.035MG;0.5MG;1MG</u>	<u>N19004 001</u>
			APR 04, 1984

ETHINYL ESTRADIOL; NORETHINDRONE ACETATE

TABLET; ORAL-21			
<u>ESTROSTEP 21</u>			
<u>+</u>	<u>PARKE DAVIS</u>	<u>0.02MG;0.03MG;0.035MG;1MG;1MG;</u>	<u>1MG</u>
			N20130 001
			OCT 09, 1996

ETHINYL ESTRADIOL; NORETHINDRONE ACETATE

TABLET; ORAL-28			
<u>ESTROSTEP FE</u>			
<u>+</u>	<u>PARKE DAVIS</u>	<u>0.02MG;0.03MG;0.035MG;1MG;1MG;</u>	<u>1MG</u>
			N20130 002
			OCT 09, 1996
<u>LOESTRIN FE 1.5/30</u>			
<u>+</u>	<u>PARKE DAVIS</u>	<u>0.03MG;1.5MG</u>	<u>N17355 001</u>
<u>LOESTRIN FE 1/20</u>			
<u>+</u>	<u>PARKE DAVIS</u>	<u>0.02MG;1MG</u>	<u>N17354 001</u>

ETODOLAC

TABLET; ORAL			
<u>LODINE</u>			
*	<u>WYETH AYERST</u>	<u>400MG</u>	<u>N18922 004</u>
			JUL 29, 1993
		<u>400MG</u>	<u>N18922 004</u>
			JUL 29, 1993
<u>+</u>		<u>500MG</u>	<u>N18922 005</u>
			JUN 28, 1996

TABLET, EXTENDED RELEASE; ORAL

<u>LODINE XL</u>			
<u>+</u>	<u>WYETH AYERST</u>	<u>400MG</u>	<u>N20584 001</u>
			OCT 25, 1996
<u>+</u>		<u>600MG</u>	<u>N20584 002</u>
			OCT 25, 1996

ETOMIDATE

INJECTABLE; INJECTION			
<u>AMIDATE</u>			
<u>AP</u>	<u>+</u>	<u>ABBOTT</u>	<u>2MG/ML</u>
			N18227 001
			SEP 07, 1982
<u>ETOMIDATE</u>			
<u>AP</u>		<u>BEDFORD</u>	<u>2MG/ML</u>
			N74593 001
			NOV 04, 1996

ETOPOSIDE

INJECTABLE; INJECTION			
<u>ETOPOSIDE</u>			
<u>AP</u>		<u>GENSIA</u>	<u>20MG/ML</u>
			N74529 001
			JUL 24, 1996

ETOPOSIDE

INJECTABLE; INJECTION

<u>AP</u>	<u>IMMUNEX</u>	<u>20MG/ML</u>	N74513 001
			MAR 14, 1996
<u>AP</u>	<u>LEDERLE</u>	<u>20MG/ML</u>	N74513 001
			MAR 14, 1996
<u>AP</u>	PHARMACHEMIE (NL)	<u>20MG/ML</u>	N74227 001
			FEB 22, 1996
<u>AP</u>	STERIS	<u>20MG/ML</u>	N74228 001
			OCT 15, 1996

ETOPOSIDE PHOSPHATE

INJECTABLE; INJECTION

ETOPOPHOS

+ BRISTOL MYERS SQUIBB EQ 100MG BASE/VIAL

N20457 001
MAY 17, 1996> ADD >EVANS BLUE

INJECTABLE; INJECTION

EVANS BLUE

@ PARKE DAVIS 0.5%

N08041 001

> ADD >> ADD >> ADD >> ADD >FAMCICLOVIR

TABLET; ORAL

FAMVIR

SMITHKLINE BEECHAM 250MG

N20363 001
APR 26, 1996FAMOTIDINE

INJECTABLE; INJECTION

PEPCID IV PRESERVATIVE FREE

* MERCK 10MG/ML

N19510 004
NOV 04, 1986

PEPCID PRESERVATIVE FREE

+ MERCK 10MG/ML

N19510 004
NOV 04, 1986FENTANYL CITRATE

INJECTABLE; INJECTION

FENTANYL CITRATE

<u>AP</u>	<u>STERIS</u>	<u>EQ 0.05MG BASE/ML</u>	N73488 001
			JUN 30, 1992
@		EQ 0.05MG BASE/ML	N73488 001
			JUN 30, 1992

FERUMOXIDES

INJECTABLE; INJECTION

FERIDEX I.V.

+ ADV MAGNETICS

EQ 11.2MG IRON/ML

N20416 001
AUG 30, 1996FERUMOXASIL

SUSPENSION; ORAL

GASTROMARK

+ ADV MAGNETICS

EQ 0.175MG IRON/ML

N20410 001
DEC 06, 1996FEXOFENADINE HYDROCHLORIDE

CAPSULE; ORAL

ALLEGRA

+ HOECHST MARION RSSL 60MG

N20625 001
JUL 25, 1996FLECAINIDE ACETATE

TABLET; ORAL

TAMBOCOR

3M

50MG

N18830 004

100MG

AUG 23, 1988

N18830 001

OCT 31, 1985

N18830 003

JUN 03, 1988

50MG

N18830 004

AUG 23, 1988

100MG

N18830 001

OCT 31, 1985

FLECAINIDE ACETATETABLET, ORAL
TAMBOCOR

@ 3M

150MG

N18830 003

JUN 03, 1988

@

200MG

N18830 002

OCT 31, 1985

FLUOCINOLONE ACETONIDE

CREAM; TOPICAL

FLUOCINOLONE ACETONIDE

AT	+	HAMILTON PHARMA CA	0.01%	N12787 004
AT	+		0.025%	N12787 002
AT	+		0.025%	N12787 005
	+		0.2%	N16161 002

SYNALAR

AT	+	SYNTEX	0.01%	N12787 004
AT	+		0.025%	N12787 002
AT	+		0.025%	N12787 005

SYNALAR-HP

+ SYNTEX 0.2%

N16161 002

OINTMENT; TOPICAL

FLUOCINOLONE ACETONIDE

AT	+	HAMILTON PHARMA CA	0.025%	N13960 001
AT	+	SYNTEX	0.025%	N13960 001

SOLUTION; TOPICAL

FLUOCINOLONE ACETONIDE

AT	+	HAMILTON PHARMA CA	0.01%	N15296 001
AT	+	SYNTEX	0.01%	N15296 001

FLUOCINOLONE ACETONIDE; NEOMYCIN SULFATE

CREAM; TOPICAL

NEO-SYNALAR

> DLT >	+	HAMILTON PHARMA CA	0.025%;EQ 3.5MG BASE/GM	N60700 001
> ADD >	+	SYNTEX	0.025%;EQ 3.5MG BASE/GM	N60700 001

FLUOCINONIDE

CREAM; TOPICAL

FLUOCINONIDE

AB	+	HAMILTON PHARMA CA	0.05%	N16908 002
AB		TARO	0.05%	N72494 001

JAN 19, 1989

N72494 001

JAN 19, 1989

AB		TICAN	0.05%	
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FLUOCINONIDE EMOLLIENT BASE

AB		HAMILTON PHARMA CA	0.05%	N16908 003
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LIDEX

AB	+	SYNTEX	0.05%	N16908 002
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LIDEX-E

AB		SYNTEX	0.05%	N16908 003
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GEL; TOPICAL

FLUOCINONIDE

AB	+	HAMILTON PHARMA CA	0.05%	N17373 001
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LIDEX

AB	+	SYNTEX	0.05%	N17373 001
----	---	--------	-------	------------

OINTMENT; TOPICAL

FLUOCINONIDE

AB	+	HAMILTON PHARMA CA	0.05%	N16909 002
----	---	--------------------	-------	------------

LIDEX

AB	+	SYNTEX	0.05%	N16909 002
----	---	--------	-------	------------

SOLUTION; TOPICAL

FLUOCINONIDE

AT	+	HAMILTON PHARMA CA	0.05%	N18849 001
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APR 06, 1984

N74799 001

DEC 31, 1996

TARO

AT			0.05%	
----	--	--	-------	--

LIDEX

AT	+	SYNTEX	0.05%	N18849 001
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APR 06, 1984

FLUOROURACIL

INJECTABLE; INJECTION

ADRUCIL

AP	+	PHARMACIA	50MG/ML	N40023 001
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OCT 18, 1991

N40023 001

OCT 18, 1991

AP		PHARMACIA AND UPJOHN	50MG/ML	
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FLUOROURACIL

AP		ROCHE	50MG/ML	N12209 001
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FLUOROURACIL

INJECTABLE; INJECTION

FLUOROURACILAP + ROCHE50MG/ML

N12209 001

FLUPHENAZINE DECANOATE

INJECTABLE; INJECTION

FLUPHENAZINE DECANOATEAO BEDFORD25MG/ML

N74531 001

AUG 30, 1996

AO GENSLA25MG/ML

N74795 001

SEP 10, 1996

FLUPHENAZINE HYDROCHLORIDE

CONCENTRATE; ORAL

FLUPHENAZINE HCLAA PHARM ASSOC5MG/ML

N74725 001

SEP 16, 1996

ELIXIR; ORAL

FLUPHENAZINE HCLAA PHARM ASSOC2.5MG/5ML

N40146 001

AUG 21, 1996

FLURAZEPAM HYDROCHLORIDE

CAPSULE; ORAL

FLURAZEPAM HCLAB BARR15MG

N70454 001

AUG 04, 1986

AB30MG

N70455 001

AUG 04, 1986

@

15MG

N70454 001

AUG 04, 1986

@

30MG

N70455 001

AUG 04, 1986

AB SUPERPHARM15MG

N71659 001

AUG 04, 1988

AB30MG

N71660 001

AUG 04, 1988

@

15MG

N71659 001

AUG 04, 1988

FLURAZEPAM HYDROCHLORIDE

CAPSULE; ORAL

FLURAZEPAM HCL

@ SUPERPHARM

30MG

N71660 001

AUG 04, 1988

AB WARNER CHILCOTT15MG

N71767 001

DEC 04, 1987

AB30MG

N71768 001

DEC 04, 1987

@

15MG

N71767 001

DEC 04, 1987

@

30MG

N71768 001

DEC 04, 1987

FLUTICASONE PROPIONATE

AEROSOL, METERED; INHALATION

FLOVENT

GLAXO WELLCOME

0.044MG/INH

N20548 001

MAR 27, 1996

0.11MG/INH

N20548 002

MAR 27, 1996

+

0.22MG/INH

N20548 003

MAR 27, 1996

FOLIC ACID

TABLET; ORAL

FOLIC ACIDAA HALSEY1MG

N83598 001

@

1MG

N83598 001

FOSFOMYCIN TROMETHAMINE

GRANULE, FOR RECONSTITUTION; ORAL

MONUROL

+ ZAMBON

EQ 3GM BASE/PACKET

N50717 001

DEC 19, 1996

FOSPHENYTOIN SODIUM

INJECTABLE; INJECTION

CEREBYX

+ PARKE DAVIS

EQ 50MG PHENYTOIN NA/ML

N20450 001

AUG 05, 1996

FUROSEMIDE

TABLET; ORAL			
<u>FUROSEMIDE</u>			
<u>AB</u>	<u>BARR</u>	<u>20MG</u>	<u>N70043 001</u>
			<u>SEP 26, 1985</u>
<u>AB</u>		<u>40MG</u>	<u>N18790 001</u>
			<u>NOV 29, 1983</u>
<u>AB</u>		<u>80MG</u>	<u>N70100 001</u>
			<u>JAN 26, 1988</u>
	@	20MG	N70043 001
			SEP 26, 1985
	@	40MG	N18790 001
			NOV 29, 1983
	@	80MG	N70100 001
			JAN 26, 1988
<u>AB</u>	<u>SUPERPHARM</u>	<u>20MG</u>	<u>N18370 002</u>
			<u>JUN 26, 1984</u>
<u>AB</u>		<u>40MG</u>	<u>N18370 001</u>
			<u>FEB 10, 1983</u>
	@	20MG	N18370 002
			JUN 26, 1984
	@	40MG	N18370 001
			FEB 10, 1983
<u>AB</u>	<u>ZENITH GOLDLINE</u>	<u>20MG</u>	<u>N18413 001</u>
			<u>NOV 30, 1983</u>
<u>AB</u>		<u>40MG</u>	<u>N18413 002</u>
			<u>NOV 30, 1983</u>
<u>AB</u>	<u>ZENITH LABS</u>	<u>20MG</u>	<u>N18413 001</u>
			<u>NOV 30, 1983</u>
<u>AB</u>		<u>40MG</u>	<u>N18413 002</u>
			<u>NOV 30, 1983</u>

GANCICLOVIR

IMPLANT; IMPLANTATION			
VITRASERT			
*	CHIRON VISION	4.5-6.4MG	N20569 001
			MAR 04, 1996
+		4.5MG	N20569 001
			MAR 04, 1996

GANCICLOVIR SODIUM

INJECTABLE; INJECTION			
CYTOVENE IV			
+	ROCHE	EQ 500MG BASE/VIAL	N19661 001
			JUN 23, 1989

GANCICLOVIR SODIUM

INJECTABLE; INJECTION			
CYTOVENE IV			
*	SYNTEX	EQ 500MG BASE/VIAL	N19661 001
			JUN 23, 1989

GEMCITABINE HYDROCHLORIDE

INJECTABLE; INJECTION			
GEMZAR			
+	LILLY	EQ 200MG BASE/VIAL	N20509 001
			MAY 15, 1996
+		EQ 1GM BASE/VIAL	N20509 002
			MAY 15, 1996

GENTAMICIN SULFATE

INJECTABLE; INJECTION			
<u>GENTAMICIN SULFATE</u>			
<u>AP</u>	<u>ABBOTT</u>	<u>EQ 50MG BASE/100ML</u>	<u>N62413 006</u>
			<u>AUG 11, 1983</u>
<u>AP</u>		<u>EQ 70MG BASE/100ML</u>	<u>N62413 007</u>
			<u>AUG 11, 1983</u>
<u>AP</u>		<u>EQ 80MG BASE/100ML</u>	<u>N62413 008</u>
			<u>AUG 11, 1983</u>
<u>AP</u>		<u>EQ 90MG BASE/100ML</u>	<u>N62413 009</u>
			<u>AUG 11, 1983</u>
<u>AP</u>		<u>EQ 100MG BASE/100ML</u>	<u>N62413 010</u>
			<u>AUG 11, 1983</u>
<u>AP</u>		<u>EQ 1.2MG BASE/ML</u>	<u>N62413 001</u>
			<u>AUG 11, 1983</u>
<u>AP</u>		<u>EQ 1.4MG BASE/ML</u>	<u>N62413 002</u>
			<u>AUG 11, 1983</u>
<u>AP</u>		<u>EQ 1.6MG BASE/ML</u>	<u>N62413 003</u>
			<u>AUG 11, 1983</u>
<u>AP</u>		<u>EQ 1.8MG BASE/ML</u>	<u>N62413 004</u>
			<u>AUG 11, 1983</u>
<u>AP</u>		<u>EQ 2MG BASE/ML</u>	<u>N62413 005</u>
			<u>AUG 11, 1983</u>
@		EQ 60MG BASE/100ML	N62413 006
			AUG 11, 1983
@		EQ 70MG BASE/100ML	N62413 007
			AUG 11, 1983
@		EQ 80MG BASE/100ML	N62413 008
			AUG 11, 1983

GENTAMICIN SULFATE

INJECTABLE; INJECTION

GENTAMICIN SULFATE

@ ABBOTT	EQ 90MG BASE/100ML	N62413 009	
		AUG 11, 1983	
@	EQ 100MG BASE/100ML	N62413 010	
		AUG 11, 1983	
@	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	

OINTMENT; OPHTHALMIC

GENTACIDIN

<u>AT</u> CIBA	<u>EQ 0.3% BASE</u>	N62501 001	
		JUL 26, 1984	
<u>AT</u> IOLAB	<u>EQ 0.3% BASE</u>	N62501 001	
		JUL 26, 1984	

SOLUTION/DROPS; OPHTHALMIC

GENTACIDIN

<u>AT</u> CIBA	<u>EQ 0.3% BASE</u>	N62480 001	
		MAR 30, 1984	
<u>AT</u> IOLAB	<u>EQ 0.3% BASE</u>	N62480 001	
		MAR 30, 1984	

> ADD > GLATIRAMER ACETATE> ADD > INJECTABLE; INJECTION> ADD > COPAXONE> ADD > + TEVA 20MG/VIAL> ADD > N20622 001
DEC 20, 1996GLIPIZIDE

TABLET; ORAL

GLIPIZIDE

<u>AB</u> NOVOPHARM	<u>5MG</u>	N74387 001	
		MAR 04, 1996	

GLIPIZIDE

TABLET; ORAL

GLIPIZIDE

<u>AB</u> NOVOPHARM	<u>10MG</u>	N74387 002	
		MAR 04, 1996	

GLUTETHIMIDE

TABLET; ORAL

GLUTETHIMIDE

<u>HALSEY</u>	<u>250MG</u>	N89458 001	
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@	250MG	N89458 001	
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OCT 10, 1986

OCT 10, 1986

GLYBURIDE

TABLET; ORAL

GLYNASE

<u>AB</u> PHARMACIA AND UPJOHN	<u>3MG</u>	N20051 002	
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+	6MG	N20051 004	
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<u>AB</u> * UPJOHN	<u>3MG</u>	N20051 002	
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@	6MG	N20051 004	
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MAR 04, 1992

SEP 24, 1993

MAR 04, 1992

SEP 24, 1993

GLYCOPYRROLATE

INJECTABLE; INJECTION

GLYCOPYRROLATE

<u>AP</u> ABBOTT	<u>0.2MG/ML</u>	N89393 001	
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@	0.2MG/ML	N89393 001	
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JUN 15, 1988

JUN 15, 1988

GOSERELIN ACETATE

IMPLANT; IMPLANTATION

ZOLADEX

+ ZENECA	EQ 10.8MG BASE	N20578 001	
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JAN 11, 1996

GRAMICIDIN; NEOMYCIN SULFATE; POLYMYXIN B SULFATE

SOLUTION/DROPS; OPHTHALMIC

NEOMYCIN AND POLYMYXIN B SULFATES AND GRAMICIDIN

AT	BAUSCH AND LOMB	0.025MG/ML;EQ 1.75MG BASE/ML; 10,000 UNITS/ML	N64047 001 JAN 31, 1996
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HALOPERIDOL

TABLET; ORAL

HALOPERIDOL

AB	BARR	0.5MG	N71156 001 JAN 02, 1987
AB		1MG	N71157 001 JAN 02, 1987
AB		2MG	N71172 001 JAN 02, 1987
AB		5MG	N71212 001 JAN 07, 1988
AB		10MG	N71173 001 JAN 07, 1988
AB		20MG	N71177 001 JAN 07, 1988
@		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

HALOPERIDOL DECANOATE

INJECTABLE; INJECTION

HALDOL DECANOATE 100

+	JOHNSON RW	EQ 100MG BASE/ML	N18701 002 OCT 31, 1989
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HALOPROGIN

CREAM; TOPICAL

HALOTEX

+	WESTWOOD SQUIBB	1%	N16942 001
@		1%	N16942 001

HEPARIN SODIUM

INJECTABLE; INJECTION

HEPARIN LOCK FLUSH

AP	SOLOPAK	100 UNITS/ML	N87905 001 APR 20, 1983
@		100 UNITS/ML	N87905 001 APR 20, 1983

HEPARIN SODIUM

AP	LILLY	1,000 UNITS/ML	N05521 001
AP		20,000 UNITS/ML	N05521 004
@		1,000 UNITS/ML	N05521 001
@		20,000 UNITS/ML	N05521 004
AP	MARSAM	1,000 UNITS/ML	N40007 001 JUN 07, 1996
AP	ORGANON	1,000 UNITS/ML	N00552 008
AP		5,000 UNITS/ML	N00552 009
AP		10,000 UNITS/ML	N00552 010
AP	PHARM SPEC ASSOC	20,000 UNITS/ML	N17780 004
@		1,000 UNITS/ML	N17780 001
@		5,000 UNITS/ML	N17780 002
@		10,000 UNITS/ML	N17780 003
@		20,000 UNITS/ML	N17780 004
AP	PHARM SPECILTS ASSOC	1,000 UNITS/ML	N17780 001
AP		5,000 UNITS/ML	N17780 002
AP		10,000 UNITS/ML	N17780 003
AP	SANOFI WINTHROP	10,000 UNITS/ML	N40095 001 JUL 26, 1996
AP	SOLOPAK	1,000 UNITS/ML	N87043 001
AP		5,000 UNITS/ML	N87077 001
AP		10,000 UNITS/ML	N87107 001
AP		5,000 UNITS/0.5ML	N87395 001
@		1,000 UNITS/ML	N87043 001
@		5,000 UNITS/ML	N87077 001
@		10,000 UNITS/ML	N87107 001
@		5,000 UNITS/0.5ML	N87395 001
AP	WYETH AYERST	5,000 UNITS/0.5ML	N17007 010

HEPARIN SODIUM

INJECTABLE; INJECTION

HEPARIN SODIUM 25,000 UNITS IN DEXTROSE 5% IN PLASTICCONTAINER

AP	MCGAW	5,000 UNITS/100ML	N19134 001
			MAR 29, 1985
@		5,000 UNITS/100ML	N19134 001
			MAR 29, 1985

HEXACHLOROPHENE

SPONGE; TOPICAL

E-Z SCRUB

+	BECTON DICKINSON	450MG	N17452 001
*	DESERT	450MG	N17452 001

HYDRALAZINE HYDROCHLORIDE

TABLET; ORAL

HYDRALAZINE HCL

AA	BARR	10MG	N88728 001
			APR 11, 1985
AA		25MG	N84106 002
AA		50MG	N84107 002
AA		100MG	N88729 001
			APR 11, 1985
@		10MG	N88728 001
			APR 11, 1985
@		25MG	N84106 002
@		50MG	N84107 002
@		100MG	N88729 001
			APR 11, 1985
AA	HALSEY	10MG	N89218 001
			JAN 22, 1986
@		10MG	N89218 001
			JAN 22, 1986

HYDRALAZINE HYDROCHLORIDE; HYDROCHLOROTHIAZIDE; RESERPINE

TABLET; ORAL

RESERPINE, HYDRALAZINE HCL AND HYDROCHLOROTHIAZIDE

BP	BARR	25MG;15MG;0.1MG	N88570 001
			APR 10, 1984
@		25MG;15MG;0.1MG	N88570 001
			APR 10, 1984

HYDROCHLOROTHIAZIDE

CAPSULE; ORAL

MICROZIDE

+ WATSON LABS

12.5MG

N20504 001
DEC 27, 1996

TABLET; ORAL

HYDRO-D

AB	HALSEY	25MG	N86504 001
AB		50MG	N83891 002
@		25MG	N86504 001
@		50MG	N83891 002

HYDROCHLOROTHIAZIDE

AB	BARR	50MG	N84771 001
AB		100MG	N83972 003
@		50MG	N84771 001
@		100MG	N83972 003
AB	SUPERPHARM	25MG	N88827 001
			DEC 28, 1984
AB		50MG	N88828 001
			DEC 28, 1984
AB		100MG	N88829 001
			DEC 28, 1984

@		25MG	N88827 001
@		50MG	DEC 28, 1984
@		100MG	N88828 001
			DEC 28, 1984
@			N88829 001
			DEC 28, 1984

HYDROCHLOROTHIAZIDE; METHYLDOPA

TABLET; ORAL

METHYLDOPA AND HYDROCHLOROTHIAZIDE

AB	NOVOPHARM	15MG;250MG	N71819 001
			APR 08, 1988
AB		25MG;250MG	N71820 001
			APR 08, 1988
AB		30MG;500MG	N71821 001
			APR 08, 1988
AB		50MG;500MG	N71822 001
			APR 08, 1988
@		15MG;250MG	N71819 001
@		25MG;250MG	APR 08, 1988
			N71820 001
			APR 08, 1988

HYDROCHLOROTHIAZIDE; METHYLDOPA

TABLET; ORAL			
<u>METHYLDOPA AND HYDROCHLOROTHIAZIDE</u>			
AB	@ NOVOPHARM	30MG;500MG	N71821 001 APR 08, 1988
	@	50MG;500MG	N71822 001 APR 08, 1988
AB	PARKE DAVIS	15MG;250MG	N71897 001 NOV 23, 1987
AB		25MG;250MG	N71898 001 NOV 23, 1987
AB		30MG;500MG	N71899 001 NOV 23, 1987
AB		50MG;500MG	N71900 001 NOV 23, 1987
	@	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

HYDROCHLOROTHIAZIDE; PROPRANOLOL HYDROCHLORIDE

TABLET; ORAL			
<u>PROPRANOLOL HCL AND HYDROCHLOROTHIAZIDE</u>			
AB	WARNER CHILCOTT	25MG;40MG	N71771 001 JAN 26, 1988
	@	25MG;40MG	N71771 001 JAN 26, 1988

HYDROCHLOROTHIAZIDE; TRIAMTERENE

CAPSULE; ORAL			
<u>DYAZIDE</u>			
AB	+ SMITHKLINE BEECHAM	25MG;37.5MG	N16042 003 MAR 03, 1994
<u>TRIAMTERENE AND HYDROCHLOROTHIAZIDE</u>			
AB	MYLAN	25MG;37.5MG	N74701 001 JUN 07, 1996
TABLET; ORAL			
<u>TRIAMTERENE AND HYDROCHLOROTHIAZIDE</u>			
AB	SIDMAK LABS NJ	25MG;37.5MG	N74026 001 APR 26, 1996

HYDROCHLOROTHIAZIDE; TRIAMTERENE

TABLET; ORAL			
<u>TRIAMTERENE AND HYDROCHLOROTHIAZIDE</u>			
AB	SIDMAK LABS NJ	50MG;75MG	N73467 001 JAN 31, 1996

HYDROCORTISONE

CREAM; TOPICAL			
<u>CORT-DOME</u>			
AT	BAYER	0.5%	N09585 003
AT		1%	N09585 001
	@	0.5%	N09585 003
	@	1%	N09585 001
<u>HYDROCORTISONE</u>			
AT	AMBIX	1%	N86080 001
AT		2.5%	N86271 001
	@	1%	N86080 001
	@	2.5%	N86271 001
AT	STOSSET	1%	N85733 001
AT	ZENITH GOLDLINE	1%	N85733 001
<u>NOGENIC HC</u>			
AT	STOSSET	1%	N87427 001
			APR 04, 1988
AT	ZENITH GOLDLINE	1%	N87427 001
			APR 04, 1988

OINTMENT; TOPICAL

<u>HYDROCORTISONE</u>			
AT	AMBIX	1%	N86079 001
AT		2.5%	N86272 001
	@	1%	N86079 001
	@	2.5%	N86272 001
<u>PENECORT</u>			
AT	ALLERGAN HERBERT	2.5%	N88217 001
			JUN 06, 1984
	@	2.5%	N88217 001
			JUN 06, 1984

HYDROCORTISONE; NEOMYCIN SULFATE; POLYMYXIN B SULFATE

SUSPENSION/DROPS; OTIC			
<u>OTICAIR</u>			
AT	BAUSCH AND LOMB	1%;EQ 3.5MG BASE/ML; 10,000 UNITS/ML	N64065 001 AUG 28, 1996

HYDROCORTISONE; POLYMYXIN B SULFATE

SOLUTION/DROPS; OTIC

OTOBiotic

<u>AT</u>	* SCHERING	<u>5MG/ML;</u>		
		<u>EQ 10,000 UNITS BASE/ML</u>	<u>N62302 001</u>	
	+	5MG/ML;		
		<u>EQ 10,000 UNITS BASE/ML</u>	<u>N62302 001</u>	
<u>AT</u>	<u>PYOCIDIN</u>	<u>5MG/ML;</u>		
	FOREST LABS	<u>EQ 10,000 UNITS BASE/ML</u>	<u>N61606 001</u>	
	@	5MG/ML;		
		<u>EQ 10,000 UNITS BASE/ML</u>	<u>N61606 001</u>	

HYDROCORTISONE ACETATE

CREAM; TOPICAL

HYDROCORTISONE ACETATE

<u>AT</u>	* CENCE	<u>1%</u>	<u>N80419 001</u>	
			<u>JAN 25, 1982</u>	
	@	1%	<u>N80419 001</u>	
			<u>JAN 25, 1982</u>	
<u>AT</u>	PUREPAC PHARM	<u>1%</u>	<u>N86052 001</u>	
<u>AT</u>	+	<u>1%</u>	<u>N86052 001</u>	

PASTE; TOPICAL

ORABASE HCA

COLGATE

HOYT LABS

0.5%

0.5%

N83205 001N83205 001> DLT >> DLT >> ADD >> ADD >HYDROCORTISONE SODIUM SUCCINATE

INJECTABLE; INJECTION

A-HYDROCORT

<u>AP</u>	ABBOTT	<u>EQ 100MG BASE/VIAL</u>	<u>N85928 001</u>	
	@	<u>EQ 100MG BASE/VIAL</u>	<u>N85928 001</u>	

HYDROXOCOBALAMIN

INJECTABLE; INJECTION

ALPHAREDISOL

<u>AP</u>	* MERCK SHARP DOHME	<u>1MG/ML</u>	<u>N80778 001</u>	
	@	1MG/ML	<u>N80778 001</u>	

HYDROXOCOBALAMIN

STERIS

<u>AP</u>		<u>1MG/ML</u>	<u>N85528 001</u>	
<u>AP</u>		<u>1MG/ML</u>	<u>N85998 001</u>	

HYDROXOCOBALAMIN

INJECTABLE; INJECTION

HYDROXOCOBALAMIN

@ STERIS

1MG/ML

N85528 001

+

1MG/ML

N85998 001HYDROXYCHLOROQUINE SULFATE

TABLET; ORAL

HYDROXYCHLOROQUINE SULFATE

AB INVAMED

200MGN40150 001

JAN 27, 1996

HYDROXYZINE HYDROCHLORIDE

INJECTABLE; INJECTION

HYDROXYZINE HCL

AP SOLOPAK

25MG/MLN86822 001

AP

50MG/MLN87310 001

@

25MG/ML

N86822 001

@

50MG/ML

N87310 001

SYRUP; ORAL

HYDROXYZINE HCL

AA KV PHARM

10MG/5MLN87730 001

AA

@

10MG/5ML

JUL 01, 1982N87730 001JUL 01, 1982

TABLET; ORAL

HYDROXYZINE HCL

AB HALSEY

10MGN89366 001

AB

@

10MG

MAY 02, 1988N89366 001

AB

SUPERPHARM

10MGMAY 02, 1988N88794 001

AB

25MGDEC 05, 1984N88795 001

AB

50MGDEC 05, 1984N88796 001

@

10MG

DEC 05, 1984N88794 001

@

25MG

DEC 05, 1984N88795 001DEC 05, 1984

HYDROXYZINE HYDROCHLORIDE

TABLET; ORAL

HYDROXYZINE HCL

@ SUPERPHARM

50MG

N88796 001

DEC 05, 1984

HYDROXYZINE PAMOATE

CAPSULE; ORAL

HYDROXYZINE PAMOATEAB CHELSEA LABSEQ 25MG HCL

N40156 001

JUL 15, 1996

ABEQ 50MG HCL

N40156 002

JUL 15, 1996

IBUPROFEN

SUSPENSION; ORAL

CHILDREN'S ADVIL

BX AM HOME PRODS

100MG/5ML

N19833 002

SEP 19, 1989

BX WHITEHALL LABS

100MG/5ML

N19833 002

SEP 19, 1989

BX CHILDREN'S MOTRIN

* MCNEIL CONS PRODS

100MG/5ML

N19842 001

SEP 19, 1989

BX IBU

KNOLL PHARM

100MG/5ML

N19784 001

DEC 18, 1989

BX MOTRIN

+ MCNEIL CONS PRODS

100MG/5ML

N19842 001

SEP 19, 1989

BX RUFEN

KNOLL PHARM

100MG/5ML

N19784 001

DEC 18, 1989

TABLET; ORAL

IBUAB KNOLL PHARM400MG

N18197 001

AB400MG

N70083 001

FEB 22, 1985

AB600MG

N70088 001

FEB 08, 1985

AB600MG

N70099 001

MAR 29, 1985

IBUPROFEN

TABLET; ORAL

IBUAB KNOLL PHARM800MG

N70745 001

JUL 23, 1986

AB IBUPROFENBARR400MG

N70079 001

JUL 24, 1985

AB600MG

N70080 001

JUL 24, 1985

AB800MG

N71448 001

FEB 18, 1987

@

400MG

N70079 001

@

600MG

JUL 24, 1985

@

800MG

N71448 001

FEB 18, 1987

ABHALSEY300MG

N71028 001

MAR 23, 1987

AB400MG

N71029 001

MAR 23, 1987

AB600MG

N71030 001

MAR 23, 1987

AB800MG

N72137 001

FEB 05, 1988

@

300MG

N71028 001

@

400MG

MAR 23, 1987

@

600MG

N71030 001

@

800MG

N72137 001

FEB 05, 1988

ABKNOLL PHARM400MG

N70083 001

FEB 22, 1985

ABRUFEN400MG

N18197 001

ABKNOLL PHARM600MG

N70088 001

FEB 08, 1985

AB600MG

N70099 001

MAR 29, 1985

AB800MG

N70745 001

JUL 23, 1986

IDARUBICIN HYDROCHLORIDE

INJECTABLE; INJECTION
IDAMYCIN
+ PHARMACIA AND UPJOHN 20MG/VIAL

N50661 003
APR 25, 1995

IMIPRAMINE HYDROCHLORIDE

TABLET; ORAL

IMIPRAMINE HCL

<u>AB</u>	EON LABS	<u>10MG</u>
<u>AB</u>		<u>25MG</u>
<u>AB</u>		<u>50MG</u>
@	EON LABS MFG	10MG
@		25MG
@		50MG

N85200 001
N84869 002
N85133 001
N85200 001
N84869 002
N85133 001

INDAPAMIDE

TABLET; ORAL

INDAPAMIDE

<u>AB</u>	INVAMED	<u>1.25MG</u>
<u>AB</u>		<u>2.5MG</u>
<u>AB</u>	LEMMON	<u>2.5MG</u>
<u>AB</u>	MYLAN	<u>2.5MG</u>
<u>AB</u>	PUREPAC PHARM	<u>1.25MG</u>
<u>AB</u>		<u>2.5MG</u>
<u>AB</u>	WATSON LABS	<u>1.25MG</u>
<u>AB</u>		<u>2.5MG</u>
<u>AB</u>	ZENITH GOLDLINE	<u>1.25MG</u>
<u>AB</u>		<u>2.5MG</u>
<u>AB</u>	ZENITH LABS	<u>2.5MG</u>
<u>AB</u>		<u>2.5MG</u>
<u>AB</u>	LOZOL	
<u>AB</u>	RHONE POULENC RORER	<u>1.25MG</u>

N74594 001
MAY 23, 1996
N74594 002
MAY 23, 1996
N74498 001
OCT 31, 1996
N74461 001
MAR 27, 1996
N74722 001
JUN 17, 1996
N74722 002
JUN 17, 1996
N74585 001
SEP 26, 1996
N74585 002
SEP 26, 1996
N74299 002
APR 29, 1996
N74299 001
JUL 27, 1995
N74299 001
JUL 27, 1995
N18538 002
APR 29, 1993

INDINAVIR SULFATE

CAPSULE; ORAL
CRIXIVAN
MERCK

EQ 200MG BASE
EQ 400MG BASE

N20685 003
MAR 13, 1996
N20685 001
MAR 13, 1996

INDOMETHACIN

CAPSULE; ORAL

INDOMETHACIN

<u>AB</u>	BARR	<u>25MG</u>
<u>AB</u>		<u>50MG</u>
@		25MG
@		50MG
<u>AB</u>	HALSEY	<u>25MG</u>
<u>AB</u>		<u>50MG</u>
@		25MG
@		50MG
<u>AB</u>	PARKE DAVIS	<u>25MG</u>
<u>AB</u>		<u>50MG</u>
@		25MG
@		50MG

N70067 001
OCT 03, 1986
N70068 001
OCT 03, 1986
N70067 001
OCT 03, 1986
N70068 001
OCT 03, 1986
N70782 001
JUN 03, 1987
N70635 001
JUN 03, 1987
N70782 001
JUN 03, 1987
N70635 001
JUN 03, 1987
N18806 001
NOV 23, 1984
N18806 002
NOV 23, 1984
N18806 001
NOV 23, 1984
N18806 002
NOV 23, 1984

INSULIN LISPRO

INJECTABLE; INJECTION
HUMALOG
+ LILLY

100 UNITS/ML

N20563 001
JUN 14, 1996

IODIXANOL

INJECTABLE; INJECTION
VISIPAQUE 270
+ NYCOMED

55%

N20351 001
MAR 22, 1996

VISIPAQUE 320
+ NYCOMED

65.2%

N20351 002
MAR 22, 1996

IOPAMIDOL

INJECTABLE; INJECTION
IOPAMIDOL

AP ELKINS SINN

41%

N74629 001
NOV 06, 1996

AP

61%

N74629 002
NOV 06, 1996

AP

76%

N74629 003
NOV 06, 1996

> ADD > AP FAULDING
> ADD >
> ADD > AP
> ADD >

61%

N74734 001
DEC 10, 1996

76%

N74734 002
DEC 10, 1996

ISOVUE-200

AP + BRACCO

41%

N18735 006
JUL 07, 1987

@

41%

N20327 001
OCT 12, 1994

@

41%

N20327 001
OCT 12, 1994

ISOVUE-250

BRACCO

51%

N20327 002
OCT 12, 1994

+

51%

N20327 002
OCT 12, 1994

ISOVUE-300

AP + BRACCO

61%

N18735 002
DEC 31, 1985

AP +

61%

N20327 003
OCT 12, 1994

*

61%

N20327 003
OCT 12, 1994

ISOVUE-370

AP + BRACCO

76%

N18735 003
DEC 31, 1985

AP +

76%

N20327 004
OCT 12, 1994

IOPAMIDOL

INJECTABLE; INJECTION
ISOVUE-370

* BRACCO

76%

N20327 004
OCT 12, 1994

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

ISONIAZID

TABLET; ORAL

ISONIAZID

AA DANBURY PHARMA

50MG

N80522 001

AA

100MG

N80523 001

@

50MG

N80522 001

@

100MG

N80523 001

AA

DURAMED

100MG

N88231 001

@

100MG

MAR 17, 1983

@

100MG

N88231 001

@

100MG

MAR 17, 1983

@ GLOBAL PHARM

100MG

N80153 001

AA

GLOBAL PHARMS

100MG

N80153 001

AA

HALSEY

50MG

N83632 001

@

50MG

N83632 001

AA

PHOENIX LABS NY

50MG

N80368 001

AA

100MG

N80368 002

@

50MG

N80368 001

@

100MG

N80368 002

AA

ZENITH LABS

100MG

N80270 001

AA

300MG

N83610 001

@

100MG

N80270 001

@

300MG

N83610 001

LANIAZID

AA

KANNETT

50MG

N80140 001

ISONIAZID

TABLET; ORAL
LANIAZID
 LANNETT

50MG

N80140 001

ISOSORBIDE DINITRATE

TABLET; ORAL
ISOSORBIDE DINITRATE

AB BARR 5MG

N86166 002

AB 10MG

SEP 19, 1986

AB 20MG

N86169 001

AB 30MG

SEP 19, 1986

N86167 001

SEP 19, 1986

N87564 001

SEP 18, 1986

@ 5MG

N86166 002

@ 10MG

SEP 19, 1986

@ 20MG

N86169 001

@ 30MG

SEP 19, 1986

N86167 001

SEP 19, 1986

N87564 001

SEP 18, 1986

BX SORBITRATE
 ZENECA 5MG

N16192 001

BX 10MG

APR 01, 1996

N16192 002

APR 01, 1996

TABLET; SUBLINGUAL
ISORDIL

AB * NYETH AYERST 10MG

N12940 005

+ 10MG

JUL 29, 1988

N12940 005

JUL 29, 1988

ISOSORBIDE DINITRATE

AB BARR 2.5MG

N84204 001

AB 5MG

SEP 18, 1986

N86168 001

AB 10MG

SEP 18, 1986

N87545 001

SEP 18, 1986

N84204 001

SEP 18, 1986

@ 2.5MG

ISOSORBIDE DINITRATE

TABLET; SUBLINGUAL
ISOSORBIDE DINITRATE
 @ BARR

5MG

N86168 001

SEP 18, 1986

@ 10MG

N87545 001

SEP 18, 1986

BX SORBITRATE
 ZENECA 2.5MG

N16191 002

APR 01, 1996

BX 5MG

N16191 001

APR 01, 1996

TABLET, CHEWABLE; ORAL
 SORBITRATE
 ZENECA

5MG

N16776 002

APR 01, 1996

+ 10MG

N16776 003

APR 01, 1996

ISOSORBIDE MONONITRATE

TABLET; ORAL
 MONOKET
 * SCHWARZ

10MG

N20215 002

JUN 30, 1993

10MG

N20215 002

JUN 30, 1993

TABLET, EXTENDED RELEASE; ORAL
 IMDUR
 @ SCHERING

30MG

N20225 001

AUG 12, 1993

+ 30MG

N20225 001

AUG 12, 1993

IVERMECTIN

TABLET; ORAL
 STROMECTOL
 + MERCK

6MG

N50742 001

NOV 22, 1996

KETAMINE HYDROCHLORIDE

INJECTABLE; INJECTION

KETALAR

<u>AP</u>	+	PARKE DAVIS	<u>EQ 50MG BASE/ML</u>	N16812 002
<u>AP</u>	+		<u>EQ 100MG BASE/ML</u>	N16812 003
<u>KETAMINE HCL</u>				
<u>AP</u>		BEDFORD	<u>EQ 50MG BASE/ML</u>	N74524 001
				MAR 22, 1996
<u>AP</u>			<u>EQ 100MG BASE/ML</u>	N74524 002
				MAR 22, 1996
<u>AP</u>		SANOFI WINTHROP	<u>EQ 50MG BASE/ML</u>	N74549 001
				JUN 27, 1996
<u>AP</u>			<u>EQ 100MG BASE/ML</u>	N74549 002
				JUN 27, 1996

KETOPROFEN

CAPSULE; ORAL

KETOPROFEN

> <u>ADD</u> >	<u>AB</u>	MYLAN	<u>50MG</u>	N74035 002
> <u>ADD</u> >				DEC 31, 1996
> <u>ADD</u> >	<u>AB</u>		<u>75MG</u>	N74035 003
> <u>ADD</u> >				DEC 31, 1996

LACTULOSE

SOLUTION; ORAL

LACTULOSE

<u>AA</u>		MORTON GROVE	<u>10GM/15ML</u>	N74602 001
				NOV 14, 1996
<u>AA</u>		PHARM ASSOC	<u>10GM/15ML</u>	N74623 001
				JUL 30, 1996

SOLUTION; ORAL, RECTAL

CEPHULAC

<u>AA</u>		HOECHST MARION RSSI	<u>10GM/15ML</u>	N17657 001
<u>AA</u>	+		<u>10GM/15ML</u>	N17657 001
<u>GENERLAC</u>				
<u>AA</u>		MORTON GROVE	<u>10GM/15ML</u>	N74603 001
				OCT 31, 1996

LATANOPROST

SOLUTION/DROPS; OPHTHALMIC

XALATAN

+	PHARMACIA AND UPJOHN	0.005%	N20597 001
			JUN 05, 1996

LEUCOVORIN CALCIUM

POWDER FOR RECONSTITUTION; ORAL

LEUCOVORIN CALCIUMIMMUNEX

		<u>EQ 60MG BASE/VIAL</u>	N08107 003
			JAN 30, 1987
@		<u>EQ 60MG BASE/VIAL</u>	N08107 003
			JAN 30, 1987

LEVOBUNOLOL HYDROCHLORIDE

SOLUTION/DROPS; OPHTHALMIC

AKBETA

<u>AT</u>		AKORN	<u>0.25%</u>	N74779 001
				OCT 29, 1996
<u>AT</u>			<u>0.5%</u>	N74780 001
				OCT 29, 1996

LEVOBUNOLOL HCL

<u>AT</u>		ALCON	<u>0.25%</u>	N74851 001
				OCT 28, 1996
<u>AT</u>			<u>0.5%</u>	N74850 001
				OCT 28, 1996

> ADD > LEVOFLOXACIN> ADD > INJECTABLE; INJECTION

> <u>ADD</u> >		LEVAQUIN		
> <u>ADD</u> >	+	JOHNSON RW	25MG/ML	N20635 001
> <u>ADD</u> >				DEC 20, 1996
> <u>ADD</u> >		LEVAQUIN IN DEXTROSE 5% IN PLASTIC CONTAINER		
> <u>ADD</u> >	+	JOHNSON RW	5MG/ML	N20635 002
> <u>ADD</u> >				DEC 20, 1996
> <u>ADD</u> >	+		500MG/100ML	N20635 003
> <u>ADD</u> >				DEC 20, 1996

> ADD > TABLET; ORAL

> <u>ADD</u> >		LEVAQUIN		
> <u>ADD</u> >		JOHNSON RW	250MG	N20634 001
> <u>ADD</u> >				DEC 20, 1996

> ADD > LEVOFLOXACIN

> ADD > TABLET; ORAL
 > ADD > LEVAQUIN
 > ADD > + JOHNSON RW 500MG N20634 002
 > ADD > DEC 20, 1996

LEVONORGESTREL

IMPLANT; IMPLANTATION
 LEVONORGESTREL
 BX + WYETH AYERST 75MG/IMPLANT N20627 001
 AUG 15, 1996
 NORPLANT II
 BX POPULATION COUNCIL 75MG/IMPLANT N20544 001
 NOV 01, 1996
 NORPLANT SYSTEM
 WYETH AYERST 36MG/IMPLANT N20088 001
 DEC 10, 1990
 + 36MG/IMPLANT N20088 001
 DEC 10, 1990

LIDOCAINE

FILM, EXTENDED RELEASE; BUCCAL
 LIDOCAINE
 + NOVEN 23MG/PATCH N20575 001
 MAY 21, 1996
 + 46.1MG/PATCH N20575 002
 MAY 21, 1996

OINTMENT; TOPICAL

ALPHACAINE
 AT CARLSLE 5% N84944 001
 AT 5% N84946 001
 @ 5% N84944 001
 @ 5% N84946 001

LIDOCAINE HYDROCHLORIDE

INJECTABLE; INJECTION

ALPHACAINE HCL
 AP CARLSLE 2% N84721 001
 @ 2% N84721 001
LIDOCAINE HCL
 AP ABBOTT 20% N89362 001
 MAY 25, 1988

LIDOCAINE HYDROCHLORIDE

INJECTABLE; INJECTION

LIDOCAINE HCL
 @ ABBOTT 20% N89362 001
 MAY 25, 1988
 AP DELL LABS 1% N83387 001
 AP 2% N83388 001
 @ 1% N83387 001
 @ 2% N83388 001
 AP FUJISAWA 2% N17508 001
 AP 4% N17508 002
 AP 20% N17508 004
 @ 2% N17508 001
 @ 4% N17508 002
 @ 20% N17508 004
 AP INTL MEDICATION 1% N17701 002
 AP 2% N17701 001
 @ 1% N17701 002
 @ 2% N17701 001

SOLUTION; ORAL

LIDOCAINE HCL
 AT MORTON GROVE 2% N87872 001
 NOV 18, 1982
LIDOCAINE HCL VISCOUS
 AT INTL MEDICATION 2% N86389 001
 FEB 02, 1982
 @ 2% N86389 001
 FEB 02, 1982
MYLOCAINE
 AT MORTON GROVE 2% N87872 001
 NOV 18, 1982

SOLUTION; TOPICAL

ANESTACON
 AT ALCON 2% N80429 001
 AT + POLYMEDICA 2% N80429 001
LIDOCAINE HCL
 AT MORTON GROVE 4% N87881 001
 NOV 18, 1982
MYLOCAINE
 AT MORTON GROVE 4% N87881 001
 NOV 18, 1982

LINDANECREAM; TOPICAL
KWELL

* REED AND CARNRICK

1%

N06309 001

1%

N84218 001

1%

N06309 001

+

N84218 001

LOTION; TOPICAL

KWELL

* REED AND CARNRICK

1%

N06309 003

1%

N84218 002

+

N84218 002

1%

N06309 003

SHAMPOO; TOPICAL

KWELL

* REED AND CARNRICK

1%

N10718 001

1%

N84219 001

+

N84219 001

1%

N10718 001

LISINAPRIL

TABLET; ORAL

ZESTRIL

AB ZENECA

2.5MG

N19777 005

APR 29, 1993

AB +

2.5MG

N19777 005

APR 29, 1993

LITHIUM CITRATE

SYRUP; ORAL

CIBALITH-S

AA SOLVAY

EQ 100MG CARBONATE/5ML

N17672 001

LITHONATE

@ SOLVAY

EQ 300MG CARBONATE/5ML

N17672 001

LORACARBEF

CAPSULE; ORAL

LORABID

* LILLY

200MG

N50668 001

DEC 31, 1991

LORACARBEF

CAPSULE; ORAL

LORABID

LILLY

200MG

N50668 001

+

400MG

DEC 31, 1991

N50668 002

APR 05, 1996

LORATADINE

SYRUP; ORAL

CLARITIN

+ SCHERING

1MG/ML

N20641 001

OCT 10, 1996

TABLET; ORAL

CLARITIN REDITABS

SCHERING

10MG

N20704 001

DEC 23, 1996

LORATADINE; PSEUDOEPHEDRINE SULFATE

TABLET, EXTENDED RELEASE; ORAL

CLARITIN-D 24 HOUR

+ SCHERING

10MG; 240MG

N20470 001

AUG 23, 1996

LORAZEPAM

INJECTABLE; INJECTION

LORAZEPAM

AP

MARSAM

2MG/ML

N74535 001

SEP 12, 1996

AP

2MG/ML

N74551 001

SEP 12, 1996

AP

1MG/0.5ML

N74551 003

SEP 12, 1996

AP

4MG/ML

N74535 002

SEP 12, 1996

AP

4MG/ML

N74551 002

SEP 12, 1996

TABLET; ORAL

LORAZEPAM

AB

BARR

0.5MG

N70472 001

DEC 10, 1985

LORAZEPAM

TABLET; ORAL

LORAZEPAM

<u>AB</u>	<u>BARR</u>	<u>1MG</u>	N70473 001
			DEC 10, 1985
<u>AB</u>		<u>2MG</u>	N70474 001
			DEC 10, 1985
	@	0.5MG	N70472 001
			DEC 10, 1985
	@	1MG	N70473 001
			DEC 10, 1985
	@	2MG	N70474 001
			DEC 10, 1985
<u>AB</u>	<u>HALSEY</u>	<u>0.5MG</u>	N71434 001
			SEP 01, 1987
<u>AB</u>		<u>1MG</u>	N71435 001
			SEP 01, 1987
<u>AB</u>		<u>2MG</u>	N71436 001
			SEP 01, 1987
	@	0.5MG	N71434 001
			SEP 01, 1987
	@	1MG	N71435 001
			SEP 01, 1987
	@	2MG	N71436 001
			SEP 01, 1987
<u>AB</u>	<u>SUPERPHARM</u>	<u>0.5MG</u>	N71245 001
			FEB 09, 1987
<u>AB</u>		<u>1MG</u>	N71246 001
			FEB 09, 1987
<u>AB</u>		<u>2MG</u>	N71247 001
			FEB 09, 1987
	@	0.5MG	N71245 001
			FEB 09, 1987
	@	1MG	N71246 001
			FEB 09, 1987
	@	2MG	N71247 001
			FEB 09, 1987

MANGANESE CHLORIDE

INJECTABLE; INJECTION

MANGANESE CHLORIDE IN PLASTIC CONTAINER

@	ABBOTT	EQ 0.1MG MANGANESE/ML	N18962 001
			JUN 26, 1986
		EQ 0.1MG MANGANESE/ML	N18962 001
			JUN 26, 1986

MANGANESE SULFATE

INJECTABLE; INJECTION

MANGANESE SULFATE

	FUJISAWA	EQ 0.1MG MANGANESE/ML	N19228 001
			MAY 05, 1987
@		EQ 0.1MG MANGANESE/ML	N19228 001
			MAY 05, 1987

MECLOCYCLINE SULFOSALICYLATE

CREAM; TOPICAL

MECLAN

+	J AND J	1%	N50518 001
+	JOHNSON RN	1%	N50518 001

MECLOFENAMATE SODIUM

CAPSULE; ORAL

MECLOFENAMATE SODIUM

<u>AB</u>	<u>BARR</u>	<u>EQ 50MG BASE</u>	N72848 001
			MAR 20, 1989
<u>AB</u>		<u>EQ 100MG BASE</u>	N72809 001
			MAR 20, 1989
	@	EQ 50MG BASE	N72848 001
			MAR 20, 1989
	@	EQ 100MG BASE	N72809 001
			MAR 20, 1989

MEDROXYPROGESTERONE ACETATE

TABLET; ORAL

MEDROXYPROGESTERONE ACETATE

<u>AB</u>	<u>BARR</u>	<u>2.5MG</u>	N40159 001
			AUG 09, 1996
<u>AB</u>		<u>5MG</u>	N40159 002
			AUG 09, 1996
<u>AB</u>		<u>10MG</u>	N40159 003
			AUG 09, 1996

MEGESTROL ACETATE

TABLET; ORAL

MEGESTROL ACETATE

<u>AB</u>	<u>BARR</u>	<u>20MG</u>	N74621 002
			AUG 16, 1996

MEPERIDINE HYDROCHLORIDE

INJECTABLE; INJECTION

DEMEROL

AP	+	SANOFI WINTHROP	25MG/ML	N05010 007
AP	+		50MG/ML	N05010 002
AP	+		75MG/ML	N05010 009
AP	+		100MG/ML	N05010 003
AP	+	STERLING WINTHROP	25MG/ML	N05010 007
AP	+		50MG/ML	N05010 002
AP	+		75MG/ML	N05010 009
AP	+		100MG/ML	N05010 003

SYRUP; ORAL

DEMEROL

AA	+	SANOFI WINTHROP	50MG/5ML	N05010 005
AA	+	STERLING WINTHROP	50MG/5ML	N05010 005

TABLET; ORAL

DEMEROL

AA	+	SANOFI WINTHROP	50MG	N05010 001
AA	+		100MG	N05010 004
AA	+	STERLING WINTHROP	50MG	N05010 001
AA	+		100MG	N05010 004

MEPROBAMATE

CAPSULE, EXTENDED RELEASE; ORAL

MEPROSPAN

		WALLACE PHARMS	200MG	N11284 001
*			400MG	N11284 002
@			200MG	N11284 001
@			400MG	N11284 002

TABLET; ORAL

MEPROBAMATE

AA		BARR	200MG	N80699 001
AA			600MG	N84230 001
	@		200MG	N80699 001
	@		600MG	N84230 001
AA		MK LABS	200MG	N14368 004
AA			400MG	N14368 002
	@		200MG	N14368 004
	@		400MG	N14368 002
		<u>NEURAMATE</u>		
AA		HALSEY	200MG	N14359 002
AA			400MG	N14359 001
	@		200MG	N14359 002

MEPROBAMATE

TABLET; ORAL

NEURAMATE

@	HALSEY	400MG	N14359 001
---	--------	-------	------------

MEROPENEM

INJECTABLE; INJECTION

MERREM I.V.

+	ZENECA	500MG/VIAL	N50706 003
			JUN 21, 1996
+		1GM/VIAL	N50706 001
			JUN 21, 1996

METHADONE HYDROCHLORIDE

CONCENTRATE; ORAL

METHADOSE

AA		MALLINCKRODT	10MG/ML	N17116 002
AA	+	MALLINCKRODT CHEM	10MG/ML	N17116 002

METHAZOLAMIDE

TABLET; ORAL

METHAZOLAMIDE

AB		INVAMED	25MG	N40102 001
				AUG 28, 1996
AB			50MG	N40102 002
				AUG 28, 1996

METHOCARBAMOL

TABLET; ORAL

METHOCARBAMOL

AA		BARR	750MG	N84486 001
----	--	------	-------	------------

METHOCARBAMOL

TABLET; ORAL

METHOCARBAMOL

> DLT >	AA	@ BARR	750MG	N84486 001
> DLT >	AA	KV PHARM	500MG	N85660 001
> ADD >	@		750MG	N85658 001
> ADD >	@		500MG	N85660 001
			750MG	N85658 001

METHYLDOPA

TABLET; ORAL

METHYLDOPA

AB	BARR	125MG	N70073 001
			OCT 09, 1986
AB		250MG	N70060 001
			OCT 09, 1986
AB		500MG	N70074 001
			OCT 09, 1986
@		125MG	N70073 001
@		250MG	N70060 001
@		500MG	N70074 001
			OCT 09, 1986
AB	HALSEY	500MG	N71753 001
			MAR 28, 1988
@		500MG	N71753 001
			MAR 28, 1988
AB	NOVOPHARM	125MG	N71105 001
			DEC 05, 1986
AB		250MG	N71106 001
			DEC 05, 1986
AB		500MG	N71067 001
			DEC 05, 1986
@		125MG	N71105 001
@		250MG	N71106 001
@		500MG	N71067 001
			DEC 05, 1986

METHYLDOPATE HYDROCHLORIDE

INJECTABLE; INJECTION

METHYLDOPATE HCL

AP	MARSAM	50MG/ML	N71812 001
			DEC 22, 1987
@		50MG/ML	N71812 001
			DEC 22, 1987

METHYLTESTOSTERONE

TABLET; ORAL

ORETON METHYL

BP	SCHERING	25MG	N03158 002
@		25MG	N03158 002

METOCLOPRAMIDE HYDROCHLORIDE

INJECTABLE; INJECTION

METOCLOPRAMIDE HCL

AP	ABBOTT	EQ 10MG BASE/2ML	N70505 001
			JUN 23, 1989
AP		EQ 5MG BASE/ML	N70505 001
			JUN 23, 1989
AP		EQ 10MG BASE/2ML	N70506 001
			JUN 22, 1989
AP		EQ 5MG BASE/ML	N70506 001
			JUN 22, 1989
AP		EQ 10MG BASE/2ML	N73117 001
			JAN 17, 1991
AP		EQ 5MG BASE/ML	N73117 001
			JAN 17, 1991
AP		EQ 10MG BASE/2ML	N73118 001
			JAN 17, 1991
AP		EQ 5MG BASE/ML	N73118 001
			JAN 17, 1991
AP	BULL D	EQ 10MG BASE/2ML	N71990 001
			JAN 18, 1989
AP	CETUS BEN VENUE	EQ 10MG BASE/2ML	N72155 001
			MAR 30, 1992
AP		EQ 5MG BASE/ML	N72155 001
			MAR 30, 1992
AP		EQ 10MG BASE/2ML	N72244 001
			MAR 30, 1992
AP		EQ 5MG BASE/ML	N72244 001
			MAR 30, 1992

METOCLOPRAMIDE HYDROCHLORIDEINJECTABLE; INJECTION
METOCLOPRAMIDE HCL

AP	CETUS BEN VENUE	EQ 10MG BASE/2ML	N72247 001 MAY 18, 1992
AP		EQ 5MG BASE/ML	N72247 001 MAY 18, 1992
AP	FAULDING	EQ 10MG BASE/2ML	N70847 001 NOV 07, 1988
AP		EQ 5MG BASE/ML	N70847 001 NOV 07, 1988
AP		EQ 10MG BASE/2ML	N71291 001 MAR 03, 1989
AP		EQ 5MG BASE/ML	N71291 001 MAR 03, 1989
AP		EQ 5MG BASE/ML	N71990 001 JAN 18, 1989
AP	GENSIA	EQ 10MG BASE/2ML	N73135 001 NOV 27, 1991
AP		EQ 5MG BASE/ML	N73135 001 NOV 27, 1991
AP	SANOFI WINTHROP	EQ 5MG BASE/ML	N74147 001 AUG 02, 1996
AP	SMITH AND NEPHEW	EQ 10MG BASE/2ML	N70623 001 MAR 02, 1987
AP		EQ 5MG BASE/ML	N70623 001 MAR 02, 1987
AP	REGLAN		
AP	* ROBINSON	EQ 10MG BASE/2ML	N17862 001
AP	+	EQ 5MG BASE/ML	N17862 001

TABLET; ORAL
METOCLOPRAMIDE HCL

AB	HALSEY	EQ 10MG BASE	N70906 001 OCT 28, 1986
@		EQ 10MG BASE	N70906 001 OCT 28, 1986
AB	SCHERING	EQ 10MG BASE	N70598 001 FEB 02, 1987
@		EQ 10MG BASE	N70598 001 FEB 02, 1987
AB	SUPERPHARM	EQ 10MG BASE	N70926 001 JUN 26, 1987
@		EQ 10MG BASE	N70926 001 JUN 26, 1987

METOLAZONETABLET; ORAL
MYKROX

	FISONS	0.5MG	N19532 001 OCT 30, 1987
	MEDEVA PHARMS	0.5MG	N19532 001 OCT 30, 1987
	ZAROXOLYN		
	FISONS	2.5MG	N17386 001
		5MG	N17386 002
*		10MG	N17386 003
	MEDEVA PHARMS	2.5MG	N17386 001
		5MG	N17386 002
+		10MG	N17386 003

METOPROLOL TARTRATETABLET; ORAL
METOPROLOL TARTRATE

> ADD >	AB	CARACO	50MG	N74644 001 DEC 10, 1996
> ADD >				
> ADD >	AB		100MG	N74644 002 DEC 10, 1996
> ADD >				

METRONIDAZOLEGEL; VAGINAL
METROGEL

*	CURATEK	0.75%	N20208 001 AUG 17, 1992
	METROGEL-VAGINAL		
+	CURATEK	0.75%	N20208 001 AUG 17, 1992

TABLET; ORAL
METROMIDOL

AB	LABS AF	250MG	N74523 001 OCT 24, 1996
AB		500MG	N74523 002 OCT 24, 1996

METRONIDAZOLE

AB	BARR	250MG	N18818 001 FEB 16, 1983
AB		500MG	N18818 002 FEB 16, 1983

METRONIDAZOLE

TABLET; ORAL

METRONIDAZOLE

@ BARR

250MG

N18818 001

FEB 16, 1983

@

500MG

N18818 002

FEB 16, 1983

AB HALSEY250MG

N70021 001

APR 02, 1985

AB500MG

N70593 001

FEB 27, 1986

@

250MG

N70021 001

APR 02, 1985

@

500MG

N70593 001

FEB 27, 1986

> ADD > AB LEMMON250MG

N70035 001

DEC 20, 1984

> ADD > AB500MG

N70044 001

FEB 08, 1985

AB ZENITH GOLDLINE250MG

N18517 001

MAY 05, 1982

AB500MG

N18517 002

MAY 05, 1982

AB ZENITH LABS250MG

N18517 001

MAY 05, 1982

AB500MG

N18517 002

MAY 05, 1982

> DLT > AB250MG

N70035 001

DEC 20, 1984

> DLT > AB500MG

N70044 001

FEB 08, 1985

> DLT > AB500MG

N70044 001

FEB 08, 1985

METRAPONE

CAPSULE; ORAL

METOPIRONE

+ CIBA

250MG

N12911 002

AUG 09, 1996

MEXILETINE HYDROCHLORIDE

CAPSULE; ORAL

MEXILETINE HCLAB GENEVA PHARMS150MG

N74450 001

MAY 16, 1996

MEXILETINE HYDROCHLORIDE

CAPSULE; ORAL

MEXILETINE HCLAB

GENEVA PHARMS

200MG

N74450 002

MAY 16, 1996

AB250MG

N74450 003

MAY 16, 1996

MICONAZOLE NITRATE

CREAM, SUPPOSITORY, TOPICAL, VAGINAL

MONISTAT DUAL PAK

* JOHNSON RW

2%, 200MG

N18889 002

OCT 17, 1988

MIDODRINE HYDROCHLORIDE

TABLET; ORAL

PROAMATINE

ROBERTS LABS

2.5MG

N19815 001

SEP 06, 1996

+

5MG

N19815 002

SEP 06, 1996

> ADD >

MIGLITOL

> ADD >

TABLET; ORAL

> ADD >

GLYSET

> ADD >

BAYER

25MG

N20682 001

DEC 18, 1996

> ADD >

50MG

N20682 002

DEC 18, 1996

> ADD >

100MG

N20682 003

DEC 18, 1996

> ADD >

+

MINOCYCLINE HYDROCHLORIDE

TABLET; ORAL

MINOCYCLINE HCL

LEDERLE

EQ 100MG BASE

N50451 002

AUG 10, 1982

+

EQ 100MG BASE

N50451 002

AUG 10, 1982

MINOXIDILSOLUTION; TOPICAL

ROGAINE
* UPJOHN

2%

N19501 001
AUG 17, 1988

MIRTAZAPINE

TABLET; ORAL
REMERON
ORGANON

15MG

N20415 001
JUN 14, 1996

+

30MG

N20415 002
JUN 14, 1996

MORPHINE SULFATE

CAPSULE, EXTENDED RELEASE; ORAL
KADIAN

+ FAULDING SVCS

20MG

N20616 001
JUL 03, 1996

+

50MG

N20616 002
JUL 03, 1996

+

100MG

N20616 003
JUL 03, 1996

INJECTABLE; INJECTION

MORPHINE SULFATE

AP MALLINCKRODT CHEM

1MG/ML

N20631 001
JUL 03, 1996

+

2MG/ML

N20631 002
JUL 03, 1996

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

@

EQ 250MG BASE/VIAL

N50550 001

@

EQ 500MG BASE/VIAL

N50550 002

@

EQ 1GM BASE/VIAL

N50550 003

MOXALACTAM DISODIUM

INJECTABLE; INJECTION

MOXAM

@ LILLY

@

EQ 2GM BASE/VIAL
EQ 10GM BASE/VIAL

N50550 004
N50550 008

NADOLOL

TABLET; ORAL

NADOLOL

AB

ZENITH GOLDLINE

20MG

N74229 001

AB

40MG

AUG 30, 1996

N74229 002

AB

80MG

AUG 30, 1996

N74255 001

AB

120MG

JAN 24, 1996

N74255 002

AB

160MG

JAN 24, 1996

N74255 003

AB

ZENITH LABS

80MG

JAN 24, 1996

N74255 001

AB

120MG

JAN 24, 1996

N74255 002

AB

160MG

JAN 24, 1996

N74255 003

JAN 24, 1996

NALBUPHINE HYDROCHLORIDE

INJECTABLE; INJECTION

NALBUPHINE HCL

+ ABBOTT

1.5MG/ML

N20200 001
MAR 12, 1993

NALBUPHINE HYDROCHLORIDE

ABBOTT

1.5MG/ML

N20200 001
MAR 12, 1993

NALIDIXIC ACID

TABLET; ORAL

NALIDIXIC ACID

AB

BARR

250MG

N70270 001

AB

500MG

JUN 29, 1988

N70271 001

JUN 29, 1988

NALIDIXIC ACIDTABLET; ORAL
NALIDIXIC ACID

<u>AB</u>	<u>BARR</u>	<u>1GM</u>	<u>N70272 001</u>
			<u>JUN 29, 1988</u>
	@	250MG	<u>N70270 001</u>
			<u>JUN 29, 1988</u>
	@	500MG	<u>N70271 001</u>
			<u>JUN 29, 1988</u>
	@	1GM	<u>N70272 001</u>
			<u>JUN 29, 1988</u>

NALOXONE HYDROCHLORIDEINJECTABLE; INJECTION
NALOXONE HCL

<u>AP</u>	<u>SMITH AND NEPHEW</u>	<u>0.4MG/ML</u>	<u>N71682 001</u>
			<u>NOV 17, 1987</u>
	@	0.4MG/ML	<u>N71682 001</u>
			<u>NOV 17, 1987</u>

NAPROXENTABLET; ORAL
NAPROXEN

<u>AB</u>	<u>BIOCRAFT</u>	<u>250MG</u>	<u>N74216 001</u>
			<u>APR 11, 1996</u>
<u>AB</u>		<u>375MG</u>	<u>N74216 002</u>
			<u>APR 11, 1996</u>
<u>AB</u>		<u>500MG</u>	<u>N74216 003</u>
			<u>APR 11, 1996</u>
<u>AB</u>	<u>SIDMAK LABS NJ</u>	<u>250MG</u>	<u>N74182 001</u>
			<u>JUN 27, 1996</u>
<u>AB</u>		<u>375MG</u>	<u>N74182 002</u>
			<u>JUN 27, 1996</u>
<u>AB</u>		<u>500MG</u>	<u>N74182 003</u>
			<u>JUN 27, 1996</u>

NAPROXEN SODIUMTABLET; ORAL
NAPROXEN SODIUM

<u>AB</u>	<u>AL HIKMA</u>	<u>EQ 500MG BASE</u>	<u>N74480 001</u>
			<u>MAY 14, 1996</u>

NAPROXEN SODIUMTABLET; ORAL
NAPROXEN SODIUM

<u>AB</u>	<u>SIDMAK LABS NJ</u>	<u>EQ 250MG BASE</u>	<u>N74242 001</u>
			<u>JUN 20, 1996</u>
<u>AB</u>		<u>EQ 500MG BASE</u>	<u>N74242 002</u>
			<u>JUN 20, 1996</u>
<u>AB</u>	<u>WEST WARD PHARM</u>	<u>EQ 500MG BASE</u>	<u>N74480 001</u>
			<u>MAY 14, 1996</u>

TABLET, EXTENDED RELEASE; ORAL
NAPRELAN

+	<u>ELAN PHARM</u>	<u>EQ 375MG BASE</u>	<u>N20353 001</u>
			<u>JAN 05, 1996</u>
+		<u>EQ 500MG BASE</u>	<u>N20353 002</u>
			<u>JAN 05, 1996</u>
+		<u>EQ 750MG BASE</u>	<u>N20353 003</u>
			<u>JAN 05, 1996</u>

NEOMYCIN SULFATEINJECTABLE; INJECTION
MYCIFRADIN

<u>AP</u>	<u>* UPJOHN</u>	<u>EQ 350MG BASE/VIAL</u>	<u>N60477 001</u>
<u>AP</u>	<u>NEOMYCIN SULFATE</u>	<u>EQ 350MG BASE/VIAL</u>	<u>N61084 001</u>
<u>AP</u>	<u>PFIZER</u>	<u>EQ 350MG BASE/VIAL</u>	<u>N61084 001</u>
<u>AP</u>	<u>SQUIBB</u>	<u>EQ 350MG BASE/VIAL</u>	<u>N60366 001</u>

POWDER; FOR RX COMPOUNDING

NEO-RX

<u>AA</u>	<u>PHARMA TEK</u>	<u>100%</u>	<u>N61579 001</u>
		<u>100%</u>	<u>N61579 001</u>

NEOMYCIN SULFATE

<u>AA</u>	<u>* ELKINS SINN</u>	<u>100%</u>	<u>N61698 001</u>
		<u>100%</u>	<u>N62385 001</u>
	<u>PADDOCK</u>	<u>100%</u>	<u>JUN 01, 1982</u>

NEVIRAPINETABLET; ORAL
VIRAMUNE

+	<u>BOEHRINGER INGELHEIM 200MG</u>	<u>N20636 001</u>
		<u>JUN 21, 1996</u>

NICARDIPINE HYDROCHLORIDE

CAPSULE; ORAL

CARDENE

AB SYNTEX 20MG

N19488 001

DEC 21, 1988

AB + 30MG

N19488 002

DEC 21, 1988

NICARDIPINE HCL

AB CHELSEA LABS 20MG

N74670 001

OCT 28, 1996

AB 30MG

N74670 002

OCT 28, 1996

AB LEMMON 20MG

N74540 001

OCT 28, 1996

AB 30MG

N74540 002

OCT 28, 1996

AB MYLAN 20MG

N74642 001

JUL 18, 1996

AB 30MG

N74642 002

JUL 18, 1996

> ADD > AB ZENITH GOLDLINE 20MG

N74439 001

DEC 10, 1996

> ADD > AB 30MG

N74439 002

DEC 10, 1996

> ADD >

NICLOSAMIDE

TABLET, CHEWABLE, ORAL

NICLOCIDE

* BAYER 500MG

N18669 001

MAY 14, 1982

@ 500MG

N18669 001

MAY 14, 1982

NICOTINE

FILM, EXTENDED RELEASE; TRANSDERMAL

HABITROL

BC * CIBA 7MG/24HR

N20076 001

NOV 27, 1991

BC * 14MG/24HR

N20076 002

NOV 27, 1991

BC * 21MG/24HR

N20076 003

NOV 27, 1991

+ 7MG/24HR

N20076 001

NOV 27, 1991

NICOTINE

FILM, EXTENDED RELEASE; TRANSDERMAL

HABITROL

+ CIBA 14MG/24HR

N20076 002

NOV 27, 1991

+ 21MG/24HR

N20076 003

NOV 27, 1991

NICODERM

BC * HOECHST MARION ROSS 7MG/24HR

N20165 001

NOV 07, 1991

BC * 14MG/24HR

N20165 002

NOV 07, 1991

BC * 21MG/24HR

N20165 003

NOV 07, 1991

NICOTROL

* PHARMACIA 5MG/16HR

N20150 001

APR 22, 1992

* 10MG/16HR

N20150 002

APR 22, 1992

* 15MG/16HR

N20150 003

APR 22, 1992

SPRAY, METERED; NASAL

NICOTROL

+ PHARMACIA AND UPJOHN 0.5MG/INH

N20385 001

MAR 22, 1996

NICOTINE POLACRILEX

GUM, CHEWING; BUCCAL

NICORETTE

* SMITHKLINE BEECHAM EQ 2MG BASE

N18612 001

JAN 13, 1984

NICORETTE DS

* SMITHKLINE BEECHAM EQ 4MG BASE

N20066 001

JUN 08, 1992

NILUTAMIDE

TABLET; ORAL

NILANDRON

+ ROUSSEL UCLAF 50MG

N20169 001

SEP 19, 1996

NITROFURANTOINSUSPENSION; ORAL
FURADANTIN

+	DURA	25MG/5ML	N09175 001
*	PROCTER AND GAMBLE	25MG/5ML	N09175 001

NITROFURAZONE

OINTMENT; TOPICAL

NITROFURAZONE

<u>AT</u>	AMBITX	0.2%	N86077 001
@		0.2%	N86077 001

NITROGLYCERIN

FILM, EXTENDED RELEASE; TRANSDERMAL

MINITRAN

<u>AB1</u>	3M	0.1MG/HR	N89771 001
			AUG 30, 1996
<u>AB1</u>		0.2MG/HR	N89772 001
			AUG 30, 1996
<u>AB1</u>		0.4MG/HR	N89773 001
			AUG 30, 1996
<u>AB1</u>		0.6MG/HR	N89774 001
			AUG 30, 1996

NITRO-DUR

<u>BX</u>	*	KEY PHARMS	0.1MG/HR	N20145 001
				APR 04, 1995
<u>BX</u>	*		0.2MG/HR	N20145 002
				APR 04, 1995
<u>BX</u>	*		0.4MG/HR	N20145 004
				APR 04, 1995
<u>BX</u>	*		0.6MG/HR	N20145 005
				APR 04, 1995
<u>AB1</u>	+	KEY PHARMS	0.1MG/HR	N20145 001
				APR 04, 1995
<u>AB1</u>	+		0.2MG/HR	N20145 002
				APR 04, 1995
<u>AB1</u>	+		0.4MG/HR	N20145 004
				APR 04, 1995
<u>AB1</u>	+		0.6MG/HR	N20145 005
				APR 04, 1995
<u>BX</u>	+		0.8MG/HR	N20145 006
				APR 04, 1995

NITROGLYCERIN

<u>AB2</u>	MYLAN	0.2MG/HR	N74609 001
			AUG 30, 1996

NITROGLYCERIN

FILM, EXTENDED RELEASE; TRANSDERMAL

NITROGLYCERIN

<u>AB2</u>	MYLAN	0.4MG/HR	N74607 001
			AUG 30, 1996
<u>AB2</u>		0.6MG/HR	N74559 001
			AUG 30, 1996

TRANSDERM-NITRO

<u>BX</u>	*	CIBA	0.2MG/HR	N20144 002
				FEB 27, 1996
<u>BX</u>	*		0.4MG/HR	N20144 003
				FEB 27, 1996
<u>BX</u>	*		0.6MG/HR	N20144 004
				FEB 27, 1996
<u>AB2</u>	+	CIBA	0.2MG/HR	N20144 002
				FEB 27, 1996
<u>AB2</u>	+		0.4MG/HR	N20144 003
				FEB 27, 1996
<u>AB2</u>	+		0.6MG/HR	N20144 004
				FEB 27, 1996
<u>BX</u>			0.1MG/HR	N20144 001
				FEB 27, 1996
<u>BX</u>			0.8MG/HR	N20144 005
				FEB 27, 1996
<u>BX</u>	+		0.8MG/HR	N20144 005
				FEB 27, 1996

NORETHINDRONE

TABLET; ORAL

NOR-Q.D.

SEARLE

NOR-QD

+ SEARLE

0.35MG	N17060 001
0.35MG	N17060 001

NORTRIPTYLINE HYDROCHLORIDE

CAPSULE; ORAL

NORTRIPTYLINE HCL

BIOCRAFT

<u>AB</u>	EQ 10MG BASE	N73667 001
		APR 11, 1996
<u>AB</u>	EQ 25MG BASE	N73667 002
		APR 11, 1996

NORTRIPTYLINE HYDROCHLORIDE

CAPSULE; ORAL
NORTRIPTYLINE HCL
AB BIOCRAFT EQ 50MG BASE N73667 003
 APR 11, 1996
AB EQ 75MG BASE N73667 004
 APR 11, 1996

NYSTATIN

POWDER; TOPICAL
MYCOSTATIN
AT + WESTWOOD SQUIBB 100,000 UNITS/GM N60578 001
NYSTOP
AT PADDOCK 100,000 UNITS/GM N64118 001
 AUG 16, 1996

NYSTATIN; TRIAMCINOLONE ACETONIDE

CREAM; TOPICAL
MYCOLOG-II
AT * APOTHECON 100,000 UNITS/GM; 0.1% N60576 002
 MAY 01, 1985
AT 100,000 UNITS/GM; 0.1% N62606 001
 MAY 15, 1985
AT + 100,000 UNITS/GM; 0.1% N62606 001
 MAY 15, 1985
 @ 100,000 UNITS/GM; 0.1% N60576 002
 MAY 01, 1985

OLANZAPINE

TABLET; ORAL
 ZYPREXA
 @ LILLY 2.5MG N20592 001
 SEP 30, 1996
 5MG N20592 002
 SEP 30, 1996
 7.5MG N20592 003
 SEP 30, 1996
 + 10MG N20592 004
 SEP 30, 1996

> ADD > OLOPATADINE HYDROCHLORIDE

> ADD > SOLUTION/DROPS; OPHTHALMIC
 > ADD > PATANOL
 > ADD > + ALCON EQ 0.1% BASE N20688 001
 > ADD > DEC 18, 1996

ONDANSETRON HYDROCHLORIDE

INJECTABLE; INJECTION
 ZOFRAN PRESERVATIVE FREE
 + GLAXO WELLCOME EQ 2MG BASE/ML N20007 003
 DEC 10, 1993

OXAZEPAM

CAPSULE; ORAL

OXAZEPAM
AB BARR 10MG N70957 001
 AUG 10, 1987
AB 15MG N71025 001
 AUG 10, 1987
AB 30MG N71026 001
 AUG 10, 1987
 @ 10MG N70957 001
 AUG 10, 1987
 @ 15MG N71025 001
 AUG 10, 1987
 @ 30MG N71026 001
 AUG 10, 1987

TABLET; ORAL

OXAZEPAM
AB BARR 15MG N70683 001
 JAN 16, 1987
 @ 15MG N70683 001
 JAN 16, 1987
AB PARKE DAVIS 15MG N71508 001
 FEB 02, 1987
 @ 15MG N71508 001
 FEB 02, 1987

OXTRIPHYLLINE

<u>SOLUTION; ORAL</u>			
<u>CHOLEDYL</u>			
	PARKE DAVIS	100MG/5ML	N09268 012
			NOV 27, 1984
@		100MG/5ML	N09268 012
			NOV 27, 1984
<u>SYRUP; ORAL</u>			
<u>CHOLEDYL</u>			
	PARKE DAVIS	50MG/5ML	N09268 011
@		50MG/5ML	N09268 011
<u>TABLET, DELAYED RELEASE; ORAL</u>			
<u>CHOLEDYL</u>			
*	PARKE DAVIS	100MG	N09268 003
*		200MG	N09268 007
@		100MG	N09268 003
@		200MG	N09268 007

OXYBUTYNIN CHLORIDE

<u>SYRUP; ORAL</u>			
<u>DITROPAN</u>			
AA	+	HOECHST MARION RSSL	5MG/5ML
			N18211 001
AA		<u>OXYBUTYNIN CHLORIDE</u>	
		SILARX	5MG/5ML
			N74520 001
			MAR 29, 1996
<u>TABLET; ORAL</u>			
<u>OXYBUTYNIN CHLORIDE</u>			
AB		ROSEMONT	5MG
			N74625 001
			JUL 31, 1996

> ADD > PANCRELIPASE (AMYLASE;LIPASE;PROTEASE)

> <u>ADD</u> >	<u>CAPSULE; ORAL</u>		
> <u>ADD</u> >	<u>COTAZYM</u>		
> <u>ADD</u> >	+	ORGANON	30,000 USP UNITS;8,000 USP UNITS;
> <u>ADD</u> >			30,000 USP UNITS
> <u>ADD</u> >			N20580 001
> <u>ADD</u> >			DEC 09, 1996

PAROXETINE HYDROCHLORIDE

<u>TABLET; ORAL</u>			
<u>PAXIL</u>			
@	SMITHKLINE BEECHAM	EQ 10MG BASE	N20031 001
			DEC 29, 1992
+		EQ 30MG BASE	N20031 003
			DEC 29, 1992
@		EQ 40MG BASE	N20031 005
			DEC 29, 1992
		EQ 10MG BASE	N20031 001
			DEC 29, 1992
		EQ 30MG BASE	N20031 003
			DEC 29, 1992
+		EQ 40MG BASE	N20031 005
			DEC 29, 1992

PENCICLOVIR

<u>CREAM; TOPICAL</u>			
<u>DENAVIR</u>			
+	SMITHKLINE BEECHAM	1%	N20629 001
			SEP 24, 1996

PENICILLIN G POTASSIUM

<u>POWDER FOR RECONSTITUTION; ORAL</u>			
<u>PENICILLIN</u>			
AA		<u>BIOCRIFT</u>	400,000 UNITS/5ML
			400,000 UNITS/5ML
			N60307 004
			N60307 004
AA		<u>PEIZERPEN G</u>	400,000 UNITS/5ML
AA		PEIZER	400,000 UNITS/5ML
	@		N60587 001
			N60587 001

TABLET; ORAL

<u>PENICILLIN G POTASSIUM</u>			
AB		<u>MYLAN</u>	800,000 UNITS
	+		800,000 UNITS
			N60781 004
			N60781 004
AB		<u>PEIZERPEN G</u>	200,000 UNITS
AB		PEIZER	250,000 UNITS
AB			400,000 UNITS
AB			800,000 UNITS
	+		50,000 UNITS
			100,000 UNITS
	@		50,000 UNITS
	@		100,000 UNITS
			N60075 003
			N60075 004
			N60075 005
			N60075 006
			N60075 001
			N60075 002
			N60075 001
			N60075 002

PENICILLIN G POTASSIUM

TABLET; ORAL

PFIZERPEN G

@ PFIZER	200,000 UNITS	N60075 003
@	250,000 UNITS	N60075 004
@	400,000 UNITS	N60075 005
@	800,000 UNITS	N60075 006

PENICILLIN V POTASSIUM

POWDER FOR RECONSTITUTION; ORAL

PFIZERPEN VK

AA PFIZER	EQ 125MG BASE/5ML	N61815 001
AA	EQ 250MG BASE/5ML	N61815 002
@	EQ 125MG BASE/5ML	N61815 001
@	EQ 250MG BASE/5ML	N61815 002

TABLET; ORAL

PFIZERPEN VK

AE PFIZER	EQ 250MG BASE	N61836 001
AE	EQ 500MG BASE	N61836 002
@	EQ 250MG BASE	N61836 001
@	EQ 500MG BASE	N61836 002

PENTAMIDINE ISETHIONATE

POWDER FOR RECONSTITUTION; INHALATION

NEBUPENT

FUJISAWA

600MG/VIAL

N19887 002
MAR 22, 1996

PENTOBARBITAL SODIUM

CAPSULE; ORAL

SODIUM PENTOBARBITAL

AA HALSEY	100MG	N84677 001
@	100MG	N84677 001

PENTOSAN POLYSULFATE SODIUM

CAPSULE; ORAL

ELMIRON

+ BAKER NORTON

100MG

N20193 001
SEP 26, 1996

PENTOSTATIN

INJECTABLE; INJECTION

NIPENT

* PARKE DAVIS 10MG/VIAL

N20122 001

OCT 11, 1991

+ SUPERGEN 10MG/VIAL

N20122 001

OCT 11, 1991

PERFLUBRON

LIQUID; ORAL

IMAGENT

* ALLIANCE PHARM 100%

N20091 001

AUG 13, 1993

@ 100%

N20091 001

AUG 13, 1993

PERINDOPRIL ERBUMINE

TABLET; ORAL

ACEON

AMARIC

2MG

N20184 001

DEC 30, 1993

4MG

N20184 002

DEC 30, 1993

* 8MG

N20184 003

DEC 30, 1993

RHONE POULENC RORER 2MG

N20184 001

DEC 30, 1993

4MG

N20184 002

DEC 30, 1993

+ 8MG

N20184 003

DEC 30, 1993

PHENSUXIMIDE

CAPSULE; ORAL

MILONTIN

* PARKE DAVIS 500MG

N08855 004

@ 500MG

N08855 004

PHENTERMINE HYDROCHLORIDE

TABLET; ORAL

UMI-PEX 30

FERNDAL LABS 30MG

N88605 001

SEP 28, 1987

+ 30MG

N88605 001

SEP 28, 1987

PHENTERMINE RESIN COMPLEX

CAPSULE, EXTENDED RELEASE; ORAL

IONAMIN

PISONS EQ 15MG BASE

N11613 004

* EQ 30MG BASE

N11613 002

MEDEVA PHARMS EQ 15MG BASE

N11613 004

+ EQ 30MG BASE

N11613 002

PHENYLBUTAZONE

CAPSULE; ORAL

PHENYLBUTAZONE

* BARR 100MG

N88994 001

DEC 04, 1985

@ 100MG

N88994 001

DEC 04, 1985

TABLET; ORAL

PHENYLBUTAZONE

* BARR 100MG

N88863 001

DEC 04, 1985

@ 100MG

N88863 001

DEC 04, 1985

PHENYLEPHRINE HYDROCHLORIDE; PROMETHAZINE HYDROCHLORIDE

SYRUP; ORAL

PHERAZINE VC

AA HALSEY 5MG/5ML; 6.25MG/5ML

N88868 001

MAR 02, 1987

@ 5MG/5ML; 6.25MG/5ML

N88868 001

MAR 02, 1987

PROMETHAZINE VC PLAIN

AA CENCI 5MG/5ML; 6.25MG/5ML

N88815 001

NOV 22, 1985

PHENYLEPHRINE HYDROCHLORIDE; PROMETHAZINE HYDROCHLORIDE

SYRUP; ORAL

PROMETHAZINE VC PLAIN

@ CENCI

5MG/5ML; 6.25MG/5ML

N88815 001

NOV 22, 1985

PHENYTOIN SODIUM

INJECTABLE; INJECTION

PHENYTOIN SODIUM

AP FUJISAWA 50MG/ML

N89003 001

MAY 31, 1985

@ 50MG/ML

N89003 001

MAY 31, 1985

AP MARSAM 50MG/ML

N89501 001

OCT 13, 1987

AP 50MG/ML

N89779 001

NOV 27, 1992

@ 50MG/ML

N89501 001

OCT 13, 1987

@ 50MG/ML

N89779 001

NOV 27, 1992

AP SOLOPAK 50MG/ML

N88520 001

DEC 17, 1984

@ 50MG/ML

N88520 001

DEC 17, 1984

PIMOZIDE

TABLET; ORAL

ORAP

* LEMMON 2MG

N17473 001

JUL 31, 1984

+ TEVA 2MG

N17473 001

JUL 31, 1984

PINDOLOL

TABLET; ORAL

PINDOLOL

AB MARTEC 5MG

N74474 001

OCT 28, 1996

AB 10MG

N74474 002

OCT 28, 1996

PIROXICAM

<u>CAPSULE; ORAL</u>			
<u>PIROXICAM</u>			
<u>AB</u>	DANBURY PHARMA	<u>10MG</u>	N74287 001 MAY 16, 1996
<u>AB</u>		<u>20MG</u>	N74287 002 MAY 16, 1996
<u>AB</u>	ZENITH GOLDLINE	<u>10MG</u>	N74148 001 JUN 03, 1996
<u>AB</u>		<u>20MG</u>	N74148 002 JUN 03, 1996

POLYESTRADIOL PHOSPHATE

<u>INJECTABLE; INJECTION</u>			
<u>ESTRADURIN</u>			
	NYETH AVERST	<u>40MG/AMP</u>	N10753 001
@		<u>40MG/AMP</u>	N10753 001

POTASSIUM CHLORIDE

<u>CAPSULE, EXTENDED RELEASE; ORAL</u>			
<u>POTASSIUM CHLORIDE</u>			
<u>AB</u>	BIOCRAFT	<u>8MEQ</u>	N73531 001 APR 26, 1996
<u>AB</u>		<u>10MEQ</u>	N73532 001 APR 26, 1996
<u>INJECTABLE; INJECTION</u>			
<u>POTASSIUM CHLORIDE</u>			
<u>AP</u>	FUTISAWA	<u>2MEQ/ML</u>	N87787 001 APR 20, 1982
@		<u>2MEQ/ML</u>	N87787 001 APR 20, 1982
<u>TABLET, EXTENDED RELEASE; ORAL</u>			
<u>K-DUR 10</u>			
BC	+ KEY PHARMS	<u>10MEQ</u>	N19439 002 JUN 13, 1986
BC	+ SCHERING	<u>10MEQ</u>	N19439 002 JUN 13, 1986
<u>K-DUR 20</u>			
	+ KEY PHARMS	<u>20MEQ</u>	N19439 001 JUN 13, 1986
	+ SCHERING	<u>20MEQ</u>	N19439 001 JUN 13, 1986

PREDNISOLONE

<u>SYRUP; ORAL</u>			
<u>PRELONE</u>			
	MURO	<u>5MG/5ML</u>	N89654 001 JAN 17, 1989
		<u>15MG/5ML</u>	N89081 001 FEB 04, 1986
+		<u>5MG/5ML</u>	N89654 001 JAN 17, 1989
+		<u>15MG/5ML</u>	N89081 001 FEB 04, 1986

PREDNISOLONE ACETATE

<u>SUSPENSION; OPHTHALMIC</u>			
<u>PRED FORTE</u>			
BX	+ ALLERGAN	<u>1%</u>	N17011 001
<u>SUSPENSION/DROPS; OPHTHALMIC</u>			
<u>PRED FORTE</u>			
<u>AB</u>	+ ALLERGAN	<u>1%</u>	N17011 001

PREDNISOLONE SODIUM PHOSPHATE; SULFACETAMIDE SODIUM

<u>SOLUTION/DROPS; OPHTHALMIC</u>			
<u>SULSTER</u>			
<u>AT</u>	+ AKORN	<u>EQ 0.23% PHOSPHATE; 10%</u>	N74511 001 JUL 30, 1996

PREDNISONE

<u>TABLET; ORAL</u>			
<u>PREDNISONE</u>			
<u>AB</u>	BARR	<u>5MG</u>	N80701 001
<u>AB</u>		<u>20MG</u>	N84634 001
@		<u>5MG</u>	N80701 001
@		<u>20MG</u>	N84634 001
<u>AB</u>	INTERPHARM	<u>5MG</u>	N89597 001 OCT 05, 1987
<u>AB</u>		<u>10MG</u>	N89598 001 OCT 05, 1987
<u>AB</u>		<u>20MG</u>	N89599 001 OCT 05, 1987
@		<u>5MG</u>	N89597 001 OCT 05, 1987

PREDNISONE

TABLET; ORAL			
<u>PREDNISONE</u>			
@	INTERPHARM	10MG	N89598 001
			OCT 05, 1987
@		20MG	N89599 001
			OCT 05, 1987
<u>AB</u>	<u>SUPERPHARM</u>	<u>5MG</u>	<u>N88865 001</u>
			OCT 25, 1984
<u>AB</u>		<u>10MG</u>	<u>N88866 001</u>
			OCT 25, 1984
<u>AB</u>		<u>20MG</u>	<u>N88867 001</u>
			OCT 25, 1984
@		5MG	N88865 001
			OCT 25, 1984
@		10MG	N88866 001
			OCT 25, 1984
@		20MG	N88867 001
			OCT 25, 1984

PROCAINAMIDE HYDROCHLORIDE

TABLET, EXTENDED RELEASE; ORAL			
<u>PROCAINAMIDE HCL</u>			
> ADD >	<u>AB</u> COPLEY PHARM	<u>1GM</u>	N40111 001
> ADD >			DEC 13, 1996
<u>PROCAN SR</u>			
> ADD >	<u>AB</u> + PARKE DAVIS	<u>1GM</u>	N88489 001
> ADD >			JAN 16, 1985
PROCANBID			
	+ PARKE DAVIS	500MG	N20545 001
			JAN 31, 1996
			N20545 002
		1GM	JAN 31, 1996

PROCHLORPERAZINE MALEATE

TABLET; ORAL			
<u>COMPazine</u>			
<u>AB</u>	SMITHKLINE BEECHAM	<u>EQ 5MG BASE</u>	N10571 001
<u>AB</u>		<u>EQ 10MG BASE</u>	N10571 002
<u>AB</u>		<u>EQ 25MG BASE</u>	N10571 003
<u>PROCHLORPERAZINE MALEATE</u>			
<u>AB</u>	COPLEY PHARM	<u>EQ 5MG BASE</u>	N40120 001
			JUL 11, 1996

PROCHLORPERAZINE MALEATE

TABLET; ORAL			
<u>PROCHLORPERAZINE MALEATE</u>			
<u>AB</u>	COPLEY PHARM	<u>EQ 10MG BASE</u>	N40120 002
			JUL 11, 1996
<u>AB</u>	INVAMED	<u>EQ 5MG BASE</u>	N40101 001
			JUL 19, 1996
<u>AB</u>		<u>EQ 10MG BASE</u>	N40101 002
			JUL 19, 1996
<u>AB</u>		<u>EQ 25MG BASE</u>	N40101 003
			JUL 19, 1996
<u>AB</u>	MYLAN	<u>EQ 5MG BASE</u>	N40185 002
			OCT 28, 1996
<u>AB</u>		<u>EQ 10MG BASE</u>	N40185 001
			OCT 28, 1996

PROMAZINE HYDROCHLORIDE

TABLET; ORAL			
SPARINE			
	WYETH AYERST	50MG	N10348 002
		100MG	N10348 003
	+	50MG	N10348 002
	@	100MG	N10348 003

PROMETHAZINE HYDROCHLORIDE

SYRUP; ORAL			
<u>PHENERGAN FORTIS</u>			
<u>AA</u>	WYETH AYERST	<u>25MG/5ML</u>	N08381 003
	+	25MG/5ML	N08381 003
<u>AA</u>	<u>PROMETH</u>	<u>6.25MG/5ML</u>	N85953 001
<u>AA</u>	BARRE	<u>25MG/5ML</u>	N84772 001
	PROMETH FORTIS		
	@ ALPHARMA	25MG/5ML	N84772 001
<u>AA</u>	<u>PROMETH PLAIN</u>	<u>6.25MG/5ML</u>	N85953 001
<u>AA</u>	<u>PROMETHAZINE</u>	<u>6.25MG/5ML</u>	N89013 001
	CENCI		SEP 20, 1985
			N89013 001
	@	6.25MG/5ML	SEP 20, 1985

PROPANTHELINE BROMIDE

TABLET; ORAL			
<u>PRO-BANTHINE</u>			
AA	* ROBERTS LABS	7.5MG	N08732 003
AA	*	15MG	N08732 002
BP	+	7.5MG	N08732 003
BP	+	15MG	N08732 002
<u>PROPANTHELINE BROMIDE</u>			
AA	FAR PHARM	15MG	N88377 001
			DEC 08, 1983
BP		15MG	N88377 001
			DEC 08, 1983
AA	ROXANE	7.5MG	N80927 001
AA		15MG	N80927 002
BP		7.5MG	N80927 001
BP		15MG	N80927 002

PROPOFOL

INJECTABLE; INJECTION
DIPRIVAN

*	ZENECA	10MG/ML	N19627 001
			OCT 02, 1989
@		10MG/ML	N19627 001
			OCT 02, 1989
+		10MG/ML	N19627 002
			JUN 11, 1996

PROPOXYPHENE HYDROCHLORIDE

CAPSULE; ORAL

<u>PROPHENE 65</u>			
AA	HALSEY	65MG	N83538 002
@		65MG	N83538 002

PROPRANOLOL HYDROCHLORIDE

TABLET; ORAL
PROPRANOLOL HCL

AB	BARR	10MG	N70319 001
			OCT 22, 1985
AB		20MG	N70320 001
			OCT 22, 1985
AB		40MG	N70103 001
			OCT 22, 1985

PROPRANOLOL HYDROCHLORIDE

TABLET; ORAL
PROPRANOLOL HCL

AB	BARR	60MG	N70321 001
			SEP 24, 1986
AB		80MG	N70322 001
			AUG 04, 1986
@		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

PROPYLTHIOURACIL

TABLET; ORAL
PROPYLTHIOURACIL

BD	BARR	50MG	N83982 001
@		50MG	N83982 001
BD	HALSEY	50MG	N80015 001
@		50MG	N80015 001

PROTIRELIN

INJECTABLE; INJECTION

<u>THYPINONE</u>			
AP	* ABBOTT	0.5MG/ML	N17638 001
@		0.5MG/ML	N17638 001
<u>THYREL TRH</u>			
AP	FERRING LABS	0.5MG/ML	N18087 001
+		0.5MG/ML	N18087 001

PSEUDOEPHEDRINE HYDROCHLORIDE

CAPSULE, EXTENDED RELEASE; ORAL

NOVAFED			
*	DOW PHARM	120MG	N17603 001
+	HOECHST MARION RSSL	120MG	N17603 001

PSEUDOEPHEDRINE HYDROCHLORIDE; TRIPROLIDINE HYDROCHLORIDE

SYRUP; ORAL

<u>AA</u>	<u>ACTAHIST</u>	<u>30MG/5ML; 1.25MG/5ML</u>	<u>N88344 001</u>
	CENCI		FEB 09, 1984
	@	30MG/5ML; 1.25MG/5ML	N88344 001
			FEB 09, 1984
<u>AA</u>	<u>TRILITRON</u>	<u>30MG/5ML; 1.25MG/5ML</u>	<u>N88474 001</u>
	NEWTRON PHARMS		FEB 12, 1985
	+	30MG/5ML; 1.25MG/5ML	N88474 001
			FEB 12, 1985

PYRIDOXINE HYDROCHLORIDE

INJECTABLE; INJECTION

<u>AP</u>	<u>PYRIDOXINE HCL</u>	<u>100MG/ML</u>	<u>N83772 001</u>
	BELL LABS		
	@	50MG/ML	N83771 001
		50MG/ML	N83771 001
	@	100MG/ML	N83772 001

QUINIDINE GLUCONATE

TABLET, EXTENDED RELEASE; ORAL

<u>AB</u>	<u>QUINIDINE GLUCONATE</u>	<u>324MG</u>	<u>N89476 001</u>
	HALSEY		APR 10, 1987
	@	324MG	N89476 001
			APR 10, 1987

QUINIDINE SULFATE

CAPSULE; ORAL

<u>AB</u>	<u>CIN-QUIN</u>	<u>200MG</u>	<u>N85296 001</u>
	SOLVAY		N85297 001
		300MG	N85296 001
		200MG	N85296 001
		300MG	N85297 001
<u>AB</u>	<u>QUINIDINE SULFATE</u>	<u>200MG</u>	<u>N85103 001</u>
	* LILLY		N85103 001
	@	200MG	

TABLET; ORAL

<u>AB</u>	<u>QUINIDINE SULFATE</u>	<u>200MG</u>	<u>N85068 001</u>
	1ST TX		

QUINIDINE SULFATE

TABLET; ORAL

<u>AB</u>	<u>QUINIDINE SULFATE</u>	<u>200MG</u>	<u>N84177 001</u>
	BARR		
	@	200MG	N84177 001
<u>AB</u>	<u>HALSEY</u>	<u>200MG</u>	<u>N83583 001</u>
		200MG	N83583 001
<u>AB</u>	<u>ICN</u>	<u>200MG</u>	<u>N83393 001</u>
	@	200MG	N83393 001
<u>AB</u>	<u>KV PHARM</u>	<u>200MG</u>	<u>N85276 001</u>
	@	200MG	N85276 001
<u>AB</u>	<u>LILLY</u>	<u>200MG</u>	<u>N85038 001</u>
	@	200MG	N85038 001
<u>AB</u>	<u>ROXANE</u>	<u>200MG</u>	<u>N83640 001</u>
<u>AB</u>	+	<u>200MG</u>	<u>N83640 001</u>
<u>AB</u>		<u>300MG</u>	<u>N85632 001</u>
<u>AB</u>	+	<u>300MG</u>	<u>N85632 001</u>
	@ SCHERER	200MG	N85068 001
<u>AB</u>	<u>QUINORA</u>	<u>300MG</u>	<u>N85222 001</u>
	* KEY PHARMS		
	@ SCHERING	300MG	N85222 001

RAMIPRIL

CAPSULE; ORAL

ALTACE

HOECHST MARION RSSL	1.25MG	N19901 001
		JAN 28, 1991
	2.5MG	N19901 002
		JAN 28, 1991
	5MG	N19901 003
		JAN 28, 1991
+	10MG	N19901 004
		JAN 28, 1991
HOECHST ROUSSEL	1.25MG	N19901 001
		JAN 28, 1991
	2.5MG	N19901 002
		JAN 28, 1991
	5MG	N19901 003
		JAN 28, 1991
*	10MG	N19901 004
		JAN 28, 1991

RANITIDINE BISMUTH CITRATE

TABLET; ORAL
TRITEC
+ GLAXO WELLCOME 400MG

N20559 001
AUG 08, 1996

RAUWOLFIA SERPENTINA

TABLET; ORAL
RAUVAL
BP PAL PAK 50MG
BP + 100MG
BP VALK 50MG
BP 100MG

N09108 002
N09108 004
N09108 002
N09108 004

REMIFENTANIL HYDROCHLORIDE

INJECTABLE; INJECTION
ULTIVA
GLAXO WELLCOME EQ 1MG BASE/VIAL
EQ 2MG BASE/VIAL
+ EQ 5MG BASE/VIAL

N20630 001
JUL 12, 1996
N20630 002
JUL 12, 1996
N20630 003
JUL 12, 1996

RIBAVIRIN

POWDER FOR RECONSTITUTION; INHALATION
VIRAZOLE
+ ICN 6GM/VIAL

> ADD >
> ADD >
> DLT >
> DLT >

* VIRATEK 5GM/VIAL

N18859 001
DEC 31, 1985
N18859 001
DEC 31, 1985

RISPERIDONE

SOLUTION; ORAL
RISPERDAL
+ JANSSEN 1MG/ML

N20588 001
JUN 10, 1996

RITONAVIR

CAPSULE; ORAL
NORVIR
+ ABBOTT 100MG

N20680 001
MAR 01, 1996

SOLUTION; ORAL
NORVIR
ABBOTT 80MG/ML

N20659 001
MAR 01, 1996

ROPIVACAINE HYDROCHLORIDE MONOHYDRATE

INJECTABLE; INJECTION
NAROPIN
ASTRA 2MG/ML
5MG/ML
7.5MG/ML
+ 10MG/ML

N20533 001
SEP 24, 1996
N20533 003
SEP 24, 1996
N20533 004
SEP 24, 1996
N20533 005
SEP 24, 1996

SELEGILINE HYDROCHLORIDE

CAPSULE; ORAL
ELDEPRYL
+ SOMERSET 5MG

N20647 001
MAY 15, 1996

TABLET; ORAL
ELDEPRYL
* SOMERSET 5MG

N19334 001
JUN 05, 1989
N19334 001
JUN 05, 1989

@ 5MG

AB SELEGILINE HCL
ENDO LABS 5MG

N74565 001
AUG 02, 1996

AB LEDERLE 5MG

N74641 001
AUG 02, 1996

AB NOVOPHARM 5MG

N74537 001
AUG 02, 1996

SELENIUM SULFIDE

LOTION/SHAMPOO; TOPICAL

SELENIUM SULFIDE

AT	SYOCSET	2.5%	N85777 001
AT	ZENITH GOLDLINE	2.5%	N85777 001

SERTRALINE HYDROCHLORIDE

TABLET; ORAL

ZOLOFT

PFIZER

EQ 25MG BASE

N19839 005
MAR 06, 1996

SODIUM CHLORIDE

INJECTABLE; INJECTION

SODIUM CHLORIDE 0.45% IN PLASTIC CONTAINER

AP	MCGAW	450MG/100ML	N18184 001
	@	450MG/100ML	N18184 001

SODIUM LACTATE

INJECTABLE; INJECTION

SODIUM LACTATE 0.167 MOLAR IN PLASTIC CONTAINER

AP	MCGAW	1.87GM/100ML	N18186 001
	@	1.87GM/100ML	N18186 001

SODIUM NITROPRUSSIDE

INJECTABLE; INJECTION

NIPRIDE

AP	* ROCHE	50MG/VIAL	N17546 001
	@	50MG/VIAL	N17546 001

SODIUM NITROPRUSSIDE

AP	ELKINS SINN	50MG/VIAL	N18581 001
			JUL 28, 1982

AP	+	50MG/VIAL	N18581 001
			JUL 28, 1982

SODIUM PHENYLBUTYRATE

POWDER; ORAL

BUPHENYL

+ UCYCLYD

3GM/TEASPOONFUL

N20573 001
APR 30, 1996

TABLET; ORAL

BUPHENYL

+ UCYCLYD

500MG

N20572 001
MAY 13, 1996

SOMATROPIN, BIOSYNTHETIC

INJECTABLE; INJECTION

SAIZEN

BX SERONO

5MG/VIAL

N19764 002
OCT 08, 1996

BX

6MG/VIAL

N19764 001
OCT 08, 1996

BX SEROSTIM

5MG/VIAL

N20604 002
AUG 23, 1996

BX +

6MG/VIAL

N20604 001
AUG 23, 1996

*

6MG/VIAL

N20604 001
AUG 23, 1996

SOYBEAN OIL

INJECTABLE; INJECTION

INTRALIPID 20%

AP PHARMACIA AND UPJOHN 20%

N20248 001
AUG 07, 1996

> ADD >

SPARFLOXACIN

> ADD >

TABLET; ORAL

> ADD >

ZAGAM

> ADD >

+ RHONE POULENC RORER 200MG

> ADD >

N20677 001
DEC 19, 1996

SPIRAPRIL HYDROCHLORIDE

TABLET; ORAL
RENORMAX
SANDOZ

3MG

6MG

12MG

24MG

© SCHERING

3MG

©

6MG

©

12MG

©

24MG

N20240 001

DEC 29, 1994

N20240 002

DEC 29, 1994

N20240 003

DEC 29, 1994

N20240 004

DEC 29, 1994

N20240 001

DEC 29, 1994

N20240 002

DEC 29, 1994

N20240 003

DEC 29, 1994

N20240 004

DEC 29, 1994

SPIRONOLACTONE

TABLET; ORAL
ALDACTONE

AB * SEARLE

25MG

AB

25MG

100MG

+

100MG

N12151 009

DEC 30, 1983

N12151 009

DEC 30, 1983

N12151 010

DEC 30, 1983

N12151 010

DEC 30, 1983

SUCCIMER

CAPSULE; ORAL
CHEMET

* ROCK PHARMA

100MG

+ SANOFI WINTHROP

100MG

N19998 002

JAN 30, 1991

N19998 002

JAN 30, 1991

SUCCINYLCHOLINE CHLORIDE

INJECTABLE; INJECTION

QUELICIN

AP * ABBOTT

20MG/ML

N08845 001

AP +

20MG/ML

N08845 006

AP +

50MG/ML

N08845 002

AP *

100MG/ML

N08845 004

QUELICIN PRESERVATIVE FREE

AP + ABBOTT

20MG/ML

N08845 001

AP +

50MG/ML

N08845 002

AP +

100MG/ML

N08845 004

SUCRALFATE

TABLET; ORAL

CARAFATE

AB + BLUE RIDGE

1GM

N18333 001

AB SUCRALFATE

BIOCRAFT

1GM

N70848 001

MAR 29, 1996

SUFENTANIL CITRATE

INJECTABLE; INJECTION

SULFENTANIL CITRATE

AP ABBOTT

EQ 0.05MG BASE/ML

N74534 001

DEC 11, 1996

SULFACETAMIDE SODIUM

LOTION; TOPICAL

KLARON

+ DERMIK LABS

10%

N19931 001

DEC 23, 1996

SOLUTION/DROPS; OPHTHALMIC

OCUSULF-30

AT OPTOPICS

30%

N80660 002

@

30%

N80660 002

SULFAMETHOXAZOLE; TRIMETHOPRIM

TABLET; ORAL

SULFAMETHOXAZOLE AND TRIMETHOPRIM

<u>AB</u>	<u>BARR</u>	<u>400MG; 80MG</u>	<u>N70006 001</u>
			NOV 14, 1984
@		400MG; 80MG	N70006 001
			NOV 14, 1984

SULFAMETHOXAZOLE AND TRIMETHOPRIM DOUBLE STRENGTH

<u>AB</u>	<u>BARR</u>	<u>800MG; 160MG</u>	<u>N70007 001</u>
			NOV 14, 1984
@		800MG; 160MG	N70007 001
			NOV 14, 1984

SULPATRIM-DS

<u>AB</u>	<u>SUPERPHARM</u>	<u>800MG; 160MG</u>	<u>N70066 001</u>
			JUN 24, 1985
@		800MG; 160MG	N70066 001
			JUN 24, 1985

SULPATRIM-SS

<u>AB</u>	<u>SUPERPHARM</u>	<u>400MG; 80MG</u>	<u>N70065 002</u>
			JUN 24, 1985
@		400MG; 80MG	N70065 002
			JUN 24, 1985

SULFISOXAZOLE

TABLET; ORAL

GANTRISIN

<u>AB</u>	<u>* ROCHE</u>	<u>500MG</u>	<u>N06525 001</u>
		500MG	N06525 001

SULFISOXAZOLE

<u>AB</u>	<u>ZENITH LABS</u>	<u>500MG</u>	<u>N80142 001</u>
<u>AB</u>	<u>+</u>	<u>500MG</u>	<u>N80142 001</u>

TAMOXIFEN CITRATE

TABLET; ORAL

NOLVADEXZENECA

@		EQ 20MG BASE	N17970 002
			MAR 21, 1994
+		EQ 20MG BASE	N17970 002
			MAR 21, 1994

TECHNETIUM TC-99M SULFUR COLLOID

SOLUTION; ORAL

TECHNETIUM TC 99M SULFUR COLLOID

> <u>DLT</u> >			
> <u>DLT</u> >			
> <u>DLT</u> >	<u>MALLINCKRODT</u>	<u>3mCi/ML</u>	<u>N17724 001</u>
> <u>ADD</u> >	@ <u>MALLINCKRODT MEDCL</u>	<u>3mCi/ML</u>	<u>N17724 001</u>

TECHNETIUM TC-99M TETROFOSMIN KIT

INJECTABLE; INJECTION

MYOVUEMEDI PHYSICS

N/A

N20372 001
FEB 09, 1996

TEMAZEPAM

CAPSULE; ORAL

TEMAZEPAM

<u>AB</u>	<u>BARR</u>	<u>15MG</u>	<u>N71174 001</u>
			JUL 10, 1986
<u>AB</u>		<u>30MG</u>	<u>N71175 001</u>
			JUL 10, 1986
@		15MG	N71174 001
			JUL 10, 1986
@		30MG	N71175 001
			JUL 10, 1986

TERAZOSIN HYDROCHLORIDE

TABLET; ORAL

HYTRIN* ABBOTT

@		EQ 1MG BASE	N19057 001
			AUG 07, 1987
@		EQ 2MG BASE	N19057 002
			AUG 07, 1987
@		EQ 5MG BASE	N19057 003
			AUG 07, 1987
@		EQ 10MG BASE	N19057 004
			AUG 07, 1987
+		EQ 1MG BASE	N19057 001
			AUG 07, 1987
		EQ 2MG BASE	N19057 002
			AUG 07, 1987
		EQ 5MG BASE	N19057 003
			AUG 07, 1987

TERAZOSIN HYDROCHLORIDE

TABLET; ORAL
HYTRIN
ABBOTT

EQ 10MG BASE

N19057 004
AUG 07, 1987

TERBINAFINE HYDROCHLORIDE

TABLET; ORAL
LAMISIL
+ SANDOZ

EQ 250MG BASE

N20539 001
MAY 10, 1996

TETRACYCLINE HYDROCHLORIDE

CAPSULE; ORAL

TETRACYCLINE HCL

AB SUPERPHARM 250MG

N62540 001
MAR 21, 1985

AB 500MG

N62540 002
MAR 21, 1985

@ 250MG

N62540 001
MAR 21, 1985

@ 500MG

N62540 002
MAR 21, 1985

THALLOUS CHLORIDE, TL-201

INJECTABLE; INJECTION
THALLOUS CHLORIDE TL 201
MEDI PHYSICS

2mCi/ML

N18110 001
FEB 01, 1982

1mCi/ML

N18110 002
FEB 27, 1996

THEOPHYLLINE

CAPSULE, EXTENDED RELEASE; ORAL

THEOVENT

BC SCHERING 125MG

N87010 001
JAN 31, 1985

BC 250MG

N87910 001
JAN 31, 1985

THEOPHYLLINE

CAPSULE, EXTENDED RELEASE; ORAL

THEOVENT

@ SCHERING

125MG

N87010 001

JAN 31, 1985

@

250MG

N87910 001

JAN 31, 1985

ELIXIR; ORAL

ELIXOMINAA

CENCI

80MG/15MLN88303 001JAN 25, 1984

@

80MG/15ML

N88303 001

JAN 25, 1984

THEOPHYLLINEAAHALSEY80MG/15MLN85169 001

@

80MG/15ML

N85169 001

INJECTABLE; INJECTION

THEOPHYLLINE 0.04% AND DEXTROSE 5% IN PLASTIC CONTAINERAPMCGAW40MG/100MLN19083 001NOV 07, 1984

@

40MG/100ML

N19083 001

NOV 07, 1984

THEOPHYLLINE 0.08% AND DEXTROSE 5% IN PLASTIC CONTAINERAPMCGAW80MG/100MLN19083 002NOV 07, 1984

@

80MG/100ML

N19083 002

NOV 07, 1984

THEOPHYLLINE 0.16% AND DEXTROSE 5% IN PLASTIC CONTAINERAPMCGAW160MG/100MLN19083 003NOV 07, 1984

@

160MG/100ML

N19083 003

NOV 07, 1984

THEOPHYLLINE 0.2% AND DEXTROSE 5% IN PLASTIC CONTAINERAPMCGAW200MG/100MLN19826 004AUG 14, 1992

@

200MG/100ML

N19826 004

AUG 14, 1992

THEOPHYLLINE 0.4% AND DEXTROSE 5% IN PLASTIC CONTAINERAPMCGAW400MG/100MLN19826 005AUG 14, 1992

@

400MG/100ML

N19826 005

AUG 14, 1992

SUSPENSION; ORAL

ELIXICON

* FOREST LABS

100MG/5ML

N85502 001

THEOPHYLLINESUSPENSION; ORALELIXICON

@ FOREST LABS 100MG/5ML

N85502 001

TABLET, EXTENDED RELEASE; ORALUNI-DUR

BC * KEY PHARMS 400MG

N89822 001

BC * 600MG

JAN 04, 1995

N89823 001

BC + SCHERING 400MG

JAN 04, 1995

N89822 001

BC + 600MG

JAN 04, 1995

N89823 001

JAN 04, 1995

UNIPHYL

BC PURDUE FREDERICK 600MG

N40086 001

APR 15, 1996

THIAMINE HYDROCHLORIDEINJECTABLE; INJECTIONTHIAMINE HCL

AP DELL LABS 100MG/ML

N88375 001

@ 100MG/ML

N88375 001

AP SANOFI WINTHROP 100MG/ML

N40079 001

MAY 03, 1996

THIORIDAZINE HYDROCHLORIDECONCENTRATE; ORALMELLARIL

AA SANDOZ 30MG/ML

N11808 012

AA + 30MG/ML

N11808 012

AA 100MG/ML

N11808 018

AA + 100MG/ML

N11808 018

THIORIDAZINE HCL

AA HI TECH PHARMA 30MG/ML

N40125 001

AUG 16, 1996

AA 100MG/ML

N40126 001

AUG 16, 1996

TABLET; ORALTHIORIDAZINE HCL

AB BARR 10MG

N88375 001

NOV 18, 1983

THIORIDAZINE HYDROCHLORIDETABLET; ORALTHIORIDAZINE HCL

AB BARR 15MG

N88461 001

NOV 18, 1983

AB 25MG

N87264 001

NOV 18, 1983

AB 100MG

N88379 001

NOV 18, 1983

AB 150MG

N88737 001

SEP 26, 1984

AB 200MG

N88738 001

OCT 16, 1984

@ 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

AB SUPERPHARM 10MG

N89103 001

JUL 02, 1985

AB 25MG

N89104 001

JUL 02, 1985

AB 50MG

N89105 001

JUL 02, 1985

@ 10MG

N89103 001

JUL 02, 1985

@ 25MG

N89104 001

JUL 02, 1985

@ 50MG

N89105 001

JUL 02, 1985

TIMOLOL MALEATESOLUTION, GEL FORMING/DROPS; OPHTHALMICTIMOPTIC-XE

+ MERCK EQ 0.25% BASE

N20330 001

NOV 04, 1993

+ EQ 0.5% BASE

N20330 002

NOV 04, 1993

TIMOLOL MALEATE

SOLUTION/DROPS; OPHTHALMIC

TIMOPTIC-XE

* MERCK

EQ 0.25% BASE

N20330 001

NOV 04, 1993

†

EQ 0.5% BASE

N20330 002

NOV 04, 1993

TIZANIDINE HYDROCHLORIDE

TABLET; ORAL

ZANAFLEX

+ ATHENA

EQ 4MG BASE

N20397 001

NOV 27, 1996

TOBRAMYCIN

SOLUTION/DROPS; OPHTHALMIC

AKTOB

AT AKORN

0.3%

N64096 001

JAN 31, 1996

TOLAZAMIDE

TABLET; ORAL

TOLAZAMIDE

AB BARR

100MG

N70162 001

JAN 14, 1986

AB

250MG

N70163 001

JAN 14, 1986

AB

500MG

N70164 001

JAN 14, 1986

@

100MG

N70162 001

JAN 14, 1986

@

250MG

N70163 001

JAN 14, 1986

@

500MG

N70164 001

JAN 14, 1986

AB

ZENITH GOLDLINE

100MG

N18894 001

NOV 02, 1984

AB

250MG

N18894 002

NOV 02, 1984

AB

500MG

N18894 003

NOV 02, 1984

> ADD >

> ADD >

> ADD >

> ADD >

> ADD >

> ADD >

> ADD >

> ADD >

> ADD >

> ADD >

TOLAZAMIDE

TABLET; ORAL

TOLAZAMIDE

AB

ZENITH LABS

100MG

N18894 001

NOV 02, 1984

AB

250MG

N18894 002

NOV 02, 1984

AB

500MG

N18894 003

NOV 02, 1984

TOLBUTAMIDE

TABLET; ORAL

ORINASE

AB

@ PHARMACIA AND UPJOHN

500MG

N10670 001

AB

* UPJOHN

500MG

N10670 001

TOLBUTAMIDE

AB

BARR

500MG

N87121 001

@

500MG

N87121 001

AB

EON LABS

500MG

N12678 001

AB

+ EON LABS MFG

500MG

N12678 001

AB

SUPERPHARM

500MG

N88893 001

NOV 19, 1984

@

500MG

N88893 001

NOV 19, 1984

TOLMETIN SODIUM

TABLET; ORAL

TOLMETIN SODIUM

AB

BAKER NORTON

EQ 600MG BASE

N74399 001

MAR 28, 1996

TOPIRAMATE

TABLET; ORAL

TOPAMAX

JOHNSON RW

25MG

N20505 004

DEC 24, 1996

@

50MG

N20505 005

DEC 24, 1996

100MG

N20505 001

DEC 24, 1996

+

200MG

N20505 002

DEC 24, 1996

> ADD > TOPIRAMATE

> ADD > TABLET; ORAL
 > ADD > TOPAMAX
 > ADD > @ JOHNSON RW 300MG N20505 003
 > ADD > DEC 24, 1996

TOPOTECAN HYDROCHLORIDE

INJECTABLE; INJECTION
 HYCAMTIN
 + SMITHKLINE BEECHAM EQ 4MG BASE/VIAL N20671 001
 MAY 28, 1996

TRANDOLAPRIL

TABLET; ORAL
 MAVIK
 KNOLL PHARM 1MG N20528 001
 APR 26, 1996
 2MG N20528 002
 APR 26, 1996
 + 4MG N20528 003
 APR 26, 1996

TRANDOLAPRIL; VERAPAMIL HYDROCHLORIDE

TABLET, EXTENDED RELEASE; ORAL
 TARKA
 + KNOLL PHARM 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

TRANEXAMIC ACID

TABLET, ORAL
 CYKLOKAPRON
 * PHARMACIA 500MG N19280 001
 DEC 30, 1986

TRANEXAMIC ACID

TABLET, ORAL
 CYKLOKAPRON
 @ PHARMACIA AND UPJOHN 500MG N19280 001
 DEC 30, 1986

TRETINOIN

CREAM; TOPICAL
 RENOVA
 J AND J 0.05% N19963 001
 DEC 29, 1995

TRIAMCINOLONE ACETONIDE

CREAM; TOPICAL

FLUTEX
AT SYOSSET 0.025% N85539 001
AT 0.1% N85539 002
AT 0.5% N85539 003
AT ZENITH GOLDLINE 0.025% N85539 001
AT 0.1% N85539 002
AT 0.5% N85539 003
AT KENALOG-H
AT APOTHECON 0.1% N86240 001
 @ 0.1% N86240 001
AT TRIALEX
AT SYOSSET 0.025% N87430 001
AT 0.1% N87429 001
AT 0.5% N87428 001
AT ZENITH GOLDLINE 0.025% N87430 001
AT 0.1% N87429 001
AT 0.5% N87428 001
 NOV 01, 1988

OINTMENT; TOPICAL

ARISTOCORT A
AT LEBERLE 0.5% N80745 003
AT + 0.5% N80745 003
AT FLUTEX
AT SYOSSET 0.025% N87375 001
 NOV 01, 1988

TRIAMCINOLONE ACETONIDE

OINTMENT; TOPICAL			
<u>FLUTEX</u>			
<u>AT</u>	<u>SYOSSET</u>	<u>0.1%</u>	<u>N87377 001</u>
			NOV 01, 1988
<u>AT</u>		<u>0.5%</u>	<u>N87376 001</u>
			NOV 01, 1988
<u>AT</u>	ZENITH GOLDLINE	<u>0.025%</u>	<u>N87375 001</u>
			NOV 01, 1988
<u>AT</u>		<u>0.1%</u>	<u>N87377 001</u>
			NOV 01, 1988
<u>AT</u>		<u>0.5%</u>	<u>N87376 001</u>
			NOV 01, 1988
<u>KENALOG</u>			
<u>AT</u>	* <u>APOTHECON</u>	<u>0.5%</u>	<u>N83944 001</u>
	@	<u>0.5%</u>	<u>N83944 001</u>
SPRAY, METERED; NASAL			
NASACORT AQ			
	+ RHONE POULENC RORER	0.055MG/INH	<u>N20468 001</u>
			MAY 20, 1996

TRIFLUOPERAZINE HYDROCHLORIDE

TABLET; ORAL			
<u>TRIFLUOPERAZINE HCL</u>			
<u>AB</u>	INVAMED	<u>EQ 1MG BASE</u>	<u>N40153 001</u>
			OCT 25, 1996
<u>AB</u>		<u>EQ 2MG BASE</u>	<u>N40153 002</u>
			OCT 25, 1996
<u>AB</u>		<u>EQ 5MG BASE</u>	<u>N40153 003</u>
			OCT 25, 1996
<u>AB</u>		<u>EQ 10MG BASE</u>	<u>N40153 004</u>
			OCT 25, 1996

TRIMETHOPRIM

TABLET; ORAL			
<u>TRIMETHOPRIM</u>			
<u>AB</u>	<u>BARR</u>	<u>100MG</u>	<u>N70494 001</u>
			JAN 22, 1986
	@	100MG	<u>N70494 001</u>
			JAN 22, 1986

TRIPLE SULFA (SULFABENZAMIDE;SULFACETAMIDE;SULFATHIAZOLE)

CREAM; VAGINAL			
<u>TRIPLE SULFA</u>			
<u>AT</u>	ALPHARMA	<u>3.7%;2.86%;3.42%</u>	<u>N87864 001</u>
			SEP 01, 1982
<u>AT</u>	NMC	<u>3.7%;2.86%;3.42%</u>	<u>N87864 001</u>
			SEP 01, 1982

TRIPROLIDINE HYDROCHLORIDE

SYRUP; ORAL			
<u>TRIPROLIDINE HCL</u>			
	HALSEY	<u>1.25MG/5ML</u>	<u>N88735 001</u>
			JAN 17, 1985
	@	1.25MG/5ML	<u>N88735 001</u>
			JAN 17, 1985

TRISULFAPYRIMIDINES (SULFADIAZINE;SULFAMERAZINE;SULFAMETHAZINE)

TABLET; ORAL			
<u>TRIPLE SULFOID</u>			
<u>AB</u>	PAL PAK	<u>167MG;167MG;167MG</u>	<u>N80094 001</u>
<u>AB</u>	VALE	<u>167MG;167MG;167MG</u>	<u>N80094 001</u>

UREA, C-13

POWDER FOR RECONSTITUTION; ORAL			
MERETEK UBT KIT (W/ PRANACTIN)			
	+ MERETEK	125MG/VIAL	<u>N20586 001</u>
			SEP 17, 1996

UROFOLLITROPIN

INJECTABLE; INJECTION			
<u>METRODIN</u>			
	* SERONO	<u>75 IU/AMP</u>	<u>N19415 002</u>
			SEP 18, 1986
		<u>150 IU/AMP</u>	<u>N19415 003</u>
			SEP 18, 1986

INJECTABLE; INTRAMUSCULAR

METRODIN			
	+ SERONO	75 IU/AMP	<u>N19415 002</u>
			SEP 18, 1986

UROFOLLITROPIN

INJECTABLE; INTRAMUSCULAR

METRODIN

+ SERONO 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> ADD > VALPROATE SODIUM

INJECTABLE; INJECTION

DEPACON

+ ABBOTT EQ 100MG BASE/ML

N20593 001
DEC 30, 1996> ADD > VALSARTAN

CAPSULE; ORAL

DIOVAN

CIBA GEIGY 80MG

N20665 001
DEC 23, 1996

+ 160MG

N20665 002
DEC 23, 1996VERAPAMIL HYDROCHLORIDE

CAPSULE, EXTENDED RELEASE; ORAL

VERELAN

* ELAN PHARM 120MG

N19614 001
MAY 29, 1990

* 180MG

N19614 003
JAN 09, 1992

* 240MG

N19614 002
MAY 29, 1990

+ LEDERLE 120MG

N19614 001
MAY 29, 1990

+ 180MG

N19614 003
JAN 09, 1992

+ 240MG

N19614 002
MAY 29, 1990VERAPAMIL HYDROCHLORIDE

CAPSULE, EXTENDED RELEASE; ORAL

VERELAN

+ LEDERLE 360MG

N19614 004
MAY 10, 1996

TABLET; ORAL

VERAPAMIL HCLAB BARR80MGN70482 001
SEP 24, 1986AB120MGN70483 001
SEP 24, 1986

@ 80MG

N70482 001
SEP 24, 1986

@ 120MG

N70483 001
SEP 24, 1986AB SIDMAK LABS NJ 40MGN72751 001
FEB 23, 1996

TABLET, EXTENDED RELEASE; ORAL

COVERA-HS

BC SEARLE 180MG

N20552 001
FEB 26, 1996

BC 240MG

N20552 002
FEB 26, 1996VERAPAMIL HCLAB MYLAN240MGN74587 001
MAR 23, 1996AB SIDMAK LABS NJ 240MGN72922 001
MAR 01, 1996VIDARABINEINJECTABLE; INJECTIONVIRA A

* PARKE DAVIS EQ 187.4MG BASE/ML

N50523 001

@ EQ 187.4MG BASE/ML

N50523 001

WARFARIN SODIUM

TABLET; ORAL

COUMADIN

DUPONT MERCK 3MG

N09218 025
NOV 18, 1996

6MG

N09218 026
NOV 18, 1996

ZAFIRLUKASTTABLET; ORAL
ACCOLATE
+ ZENECA

20MG

N20547 001
SEP 26, 1996ZIDOVUDINETABLET; ORAL
RETROVIR
+ GLAXO WELLCOME

300MG

N20518 002
OCT 04, 1996> ADD >ZILEUTON> ADD >

TABLET; ORAL

> ADD >

ZYFLO

> ADD >

ABBOTT

> ADD >

300MG

> ADD >

+

600MG

> ADD >N20471 001
DEC 09, 1996
N20471 003
DEC 09, 1996

ASPIRIN

TABLET, EXTENDED RELEASE; ORAL

8-HOUR BAYER

+ BAYER	650MG	N16030 001
* STERLING	650MG	N16030 001
MEASURIN		
+ BAYER	650MG	N16030 002
* STERLING	650MG	N16030 002

BENTOQUATAM

LOTION; TOPICAL

IVY BLOCK

+ ENVIRODERM	5%	N20532 001
		AUG 26, 1996

BROMPHENIRAMINE MALEATE

TABLET, EXTENDED RELEASE; ORAL

DIMETANE

* ROBINS AH	8MG	N10799 010
		JUN 10, 1983
*	12MG	N10799 011
		JUN 10, 1983
@ WHITEHALL ROBINS	8MG	N10799 010
		JUN 10, 1983
DIMETAPP		
+ WHITEHALL ROBINS	12MG	N10799 011
		JUN 10, 1983

BROMPHENIRAMINE MALEATE; PHENYLPROPANOLAMINE HYDROCHLORIDE

ELIXIR; ORAL

DIMETAPP

* ROBINS AH	2MG/5ML; 12.5MG/5ML	N13087 003
		MAR 29, 1984
+ WHITEHALL ROBINS	2MG/5ML; 12.5MG/5ML	N13087 003
		MAR 29, 1984

TABLET, EXTENDED RELEASE; ORAL

DIMETAPP

* ROBINS AH	12MG; 75MG	N12436 003
		MAY 14, 1985
+ WHITEHALL ROBINS	12MG; 75MG	N12436 003
		MAY 14, 1985

BROMPHENIRAMINE MALEATE; PSEUDOEPHEDRINE HYDROCHLORIDE

TABLET, EXTENDED RELEASE; ORAL

EFIDAC 24 PSEUDOEPHEDRINE HCL/BROMPHENIRAMINE MALEATE

+ ALZA	16MG; 240MG	N19672 001
		MAR 29, 1996

CHLORPHENIRAMINE MALEATE; PSEUDOEPHEDRINE HYDROCHLORIDE

CAPSULE, EXTENDED RELEASE; ORAL

CODIMAL-L.A. 12

CENT PHARMS	12MG; 120MG	N18935 001
		APR 15, 1985
+	12MG; 120MG	N18935 001
		APR 15, 1985

PSEUDOEPHEDRINE HCL AND CHLORPHENIRAMINE MALEATE

+ CENT PHARMS	8MG; 120MG	N19428 001
		AUG 02, 1988

PSEUDOEPHEDRINE HCL/CHLORPHENIRAMINE MALEATE

* GRAHAM	8MG; 120MG	N18844 001
		MAR 20, 1985
*	12MG; 120MG	N18843 001
		MAR 20, 1985
@	8MG; 120MG	N18844 001
		MAR 20, 1985
@	12MG; 120MG	N18843 001
		MAR 18, 1985
PSEUDOEPHEDRINE HYDROCHLORIDE AND CHLORPHENIRAMINE MALEATE		
CENT PHARMS	8MG; 120MG	N19428 001
		AUG 02, 1988

CHLORPHENIRAMINE POLISTIREX; PHENYLPROPANOLAMINE POLISTIREX

SUSPENSION, EXTENDED RELEASE; ORAL

CORSYM

@ FISOXS

EQ 4MG MALEATE/5ML;	
EQ 37.5MG HCL/5ML	N18050 001
	JAN 04, 1984

@ MEDEVA PHARMS

EQ 4MG MALEATE/5ML;	
EQ 37.5MG HCL/5ML	N18050 001
	JAN 04, 1984

CIMETIDINE

TABLET; ORAL
TAGAMET HB

* SMITHKLINE BEECHAM 100MG

100MG

+

200MG

N20238 001

JUN 19, 1995

N20238 001

JUN 19, 1995

N20238 002

AUG 21, 1996

CLEMASTINE FUMARATE; PHENYLPROPANOLAMINE HYDROCHLORIDE

TABLET, EXTENDED RELEASE; ORAL
TAVIST-D

SANDOZ

1.34MG; 75MG

N20640 001

AUG 09, 1996

CLOTRIMAZOLE

CREAM, SUPPOSITORY; TOPICAL, VAGINAL
GYNE-LOTTRIMIN 3 COMBINATION PACK

+ SCHERING PLOUGH 1%, 200MG

N20526 002

JUL 29, 1996

SUPPOSITORY; VAGINAL
GYNE-LOTTRIMIN

+ SCHERING PLOUGH 100MG

N17717 002

NOV 30, 1990

GYNE-LOTTRIMIN 3

+ SCHERING PLOUGH 200MG

N20525 001

JUL 29, 1996

MYCELEX-7

BAYER

100MG

N18182 002

DEC 26, 1991

TABLET, VAGINAL
GYNE-LOTTRIMIN

* SCHERING PLOUGH 100MG

N17717 002

NOV 30, 1990

MYCELEX-7

BAYER

100MG

N18182 002

DEC 26, 1991

DEXTROMETHORPHAN POLISTIREX

SUSPENSION, EXTENDED RELEASE; ORAL
DELSYM

* FISONS

EQ 30MG HBR/5ML

N18658 001

+ MEDEVA PHARMS

EQ 30MG HBR/5ML

N18658 001

DOXYLAMINE SUCCINATE

TABLET; ORAL

DOXYLAMINE SUCCINATE

PERRIGO

25MG

N40167 001

SEP 18, 1996

IBUPROFEN

CAPSULE; ORAL

MIDOL

@ BAYER

200MG

N70626 001

SEP 02, 1987

@

200MG

N71002 001

SEP 02, 1987

@ WINTHROP

200MG

N70626 001

SEP 02, 1987

@

200MG

N71002 001

SEP 02, 1987

PROVEL

* SANDOZ

200MG

N20402 001

APR 20, 1995

@

200MG

N20402 001

APR 20, 1995

SUSPENSION; ORAL

CHILDREN'S ADVIL

WHITEHALL ROBINS

100MG/5ML

N20589 001

JUN 27, 1996

SUSPENSION/DROPS; ORAL

CHILDREN'S MOTRIN

+ MCNEIL CONS PRODS

40MG/ML

N20603 001

JUN 10, 1996

TABLET; ORAL

IBUPROFEN

BARR

200MG

N70493 001

DEC 24, 1985

IBUPROFEN

TABLET, CHEWABLE; ORAL
JUNIOR STRENGTH MOTRIN
MCNEIL CONS PRODS 100MG

N20601 002
NOV 15, 1996

BARR	200MG	N70908 001
		SEP 26, 1986
	200MG	N71462 001
		OCT 02, 1986
@	200MG	N70493 001
		DEC 24, 1985
@	200MG	N70908 001
		SEP 26, 1986
@	200MG	N71462 001
		OCT 02, 1986
HALSEY	200MG	N71027 001
		SEP 29, 1987
@	200MG	N71027 001
		SEP 29, 1987
@ LEMMON	200MG	N73141 001
		MAY 29, 1992
MCNEIL CONS PRODS	200MG	N73019 001
		MAR 30, 1994
+	200MG	N73019 001
		MAR 30, 1994
TAG PHARMS	200MG	N73141 001
		MAY 29, 1992

$$\begin{array}{r} > \frac{\text{ADD}}{\text{ADD}} > \\ > \frac{\text{ADD}}{\text{ADD}} > \\ > \text{ADD} > \end{array}$$

JUNIOR STRENGTH ADVIL			
WHITEHALL ROBINS	100MG		N20267 002
			DEC 13, 1996
JUNIOR STRENGTH MOTRIN			
MCNEIL CONS PRODS	100MG		N20602 001
			JUN 10, 1996
MIDOL			
@ BAYER	200MG		N70591 001
			SEP 02, 1987
@	200MG		N71001 001
			SEP 02, 1987
@ WINTHROP	200MG		N70591 001
			SEP 02, 1987
@	200MG		N71001 001
			SEP 02, 1987
NUPRIN			
* BRISTOL MYERS	200MG		N72035 001
			FEB 16, 1988
	200MG		N72035 001
			FEB 16, 1988

TABLET, CHEWABLE; ORAL
CHILDREN'S MOTRIN
MCNEIL CONS PRODS 50MG N20601 001
NOV 15, 1996

INJECTABLE; INJECTION
REGULAR ILETIN II
+ LILLY
@

N18478 001
N18478 001

INJECTABLE, INJECTION
NOVOLIN N
* NOVO NORDISK

N19065 001
JAN 23, 1985
N19065 001
JAN 23, 1985

INJECTABLE; INJECTION
PROTAMINE ZINC INSULIN
+ SQUIBB
+
@
@

N17928 001
N17928 003
N17928 001
N17928 003

```
> DLT > INJECTABLE; INJECTION
> DLT > LENTE INSULIN
> DLT > * NOVIO NORDISK
> ADD > @
```

N17998 003
N17998 003

INJECTABLE, INJECTION
ULTRALENTE
+ NOVO NORDISK

N18385 001

INSULIN ZINC SUSP EXTENDED PURIFIED BEEF

INJECTABLE; INJECTION

ULTRALENTE

@ NOVO NORDISK 100 UNITS/ML

N18385 001 > ADD >
> ADD >INSULIN ZINC SUSP PROMPT PURIFIED PORK

INJECTABLE; INJECTION

SEMILENTE

* NOVO NORDISK 100 UNITS/ML
@ 100 UNITS/MLN18382 001
N18382 001INSULIN ZINC SUSP PURIFIED BEEF

INJECTABLE; INJECTION

LENTE ILETIN II

@ LILLY 100 UNITS/ML
+ 100 UNITS/MLN18477 001
N18477 001MICONAZOLE NITRATE

CREAM; VAGINAL

MICONAZOLE 7

NMC 2%

N74164 001
MAR 29, 1996

MICONAZOLE NITRATE

GW LABS 2%

N74366 001
FEB 22, 1996

CREAM, SUPPOSITORY; TOPICAL, VAGINAL

MONISTAT-3 COMBINATION PACK

+ ADV CARE 2%, 200MG

N20670 002
APR 16, 1996MINOXIDIL

SOLUTION; TOPICAL

MINOXIDIL (FOR MEN)

ALPHARMA 2%

N74588 001
APR 05, 1996

BAUSCH AND LOMB 2%

N74643 001
APR 09, 1996

COPLEY PHARM 2%

N74500 001
MAY 23, 1996MINOXIDIL

SOLUTION; TOPICAL

MINOXIDIL (FOR MEN)

HI TECH PHARMA 2%

N74731 001
DEC 24, 1996

LEMMON 2%

N74589 001
APR 05, 1996

SIGHT PHARMS 2%

N74743 002
OCT 18, 1996

MINOXIDIL (FOR WOMEN)

SIGHT PHARMS 2%

N74743 001
OCT 18, 1996

ROGAINE (FOR MEN)

+ PHARMACIA AND UPJOHN 2%

N19501 002
FEB 09, 1996

ROGAINE (FOR WOMEN)

+ PHARMACIA AND UPJOHN 2%

N19501 003
FEB 09, 1996NAPHAZOLINE HYDROCHLORIDE; PHENIRAMINE MALEATE

SOLUTION/DROPS; OPHTHALMIC

OCUHIST

AKORN 0.025%; 0.3%

N20485 001
JAN 31, 1996

OPCON-A

* BAUSCH AND LOMB 0.027%; 0.315%

N20065 001
JUN 08, 1994

+ 0.02675%; 0.315%

N20065 001
JUN 08, 1994NICOTINE

FILM, EXTENDED RELEASE; TRANSDERMAL

NICODERM CQ

+ HOECHST MARION RSSL 7MG/24HR

N20165 006
AUG 02, 1996

+ 14MG/24HR

N20165 005
AUG 02, 1996

+ 21MG/24HR

N20165 004
AUG 02, 1996

NICOTROL

+ MCNEIL CONS PRODS 15MG/16HR

N20536 001
JUL 03, 1996

NICOTINE POLACRILEXGUM, CHEWING; BUCCAL
NICORETTE

+ SMITHKLINE BEECHAM	EQ 2MG BASE	N18612 002
		FEB 09, 1996
+	EQ 4MG BASE	N20066 002
		FEB 09, 1996

NIZATIDINETABLET; ORAL
AXID AR

+ WHITEHALL ROBINS	75MG	N20555 001
		MAY 09, 1996

PSEUDOEPHEDRINE HYDROCHLORIDETABLET, EXTENDED RELEASE; ORAL
EFIDAC 24 PSEUDOEPHEDRINE HCL

+ ALZA	240MG	N20021 002
		DEC 15, 1992
* CIBA	240MG	N20021 002
		DEC 15, 1992

PSEUDOEPHEDRINE POLISTIREXSUSPENSION, EXTENDED RELEASE; ORAL
PSEUDO-12

@ Fisons	EQ 60MG HCL/5ML	N19401 001
		JUN 19, 1987
@ MEDEVA PHARMS	EQ 60MG HCL/5ML	N19401 001
		JUN 19, 1987

DRUG PRODUCTS WITH APPROVAL UNDER SECTION 505 OF THE ACT ADMINISTERED BY THE CENTER FOR BIOLOGICS EVALUATION AND RESEARCH LIST
CUMULATIVE SUPPLEMENT NUMBER 12/ DEC '96

NO DECEMBER 1996 APPROVALS

LIST of ORPHAN PRODUCTS DESIGNATIONS and APPROVALS

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January 1, 1996 thru December, 1996

NAME Generic Name TN=Trade Name	INDICATION DESIGNATED	SPONSOR & ADDRESS DD=Date Designated MA=Marketing Approval
2'-deoxycytidine TN=	As a host-protective agent in the treatment of acute myelogenous leukemia.	Grant, Steven M.D. Massey Cancer Center, VCU P.O. Box 980230 Richmond, VA 23298 DD=09/09/96 MA= / /
9-nitro-20-(S)-camptothecin (9-NC) TN=	Treatment of pancreatic cancer.	Stehlin Foundation for Cancer Research 1315 Calhoun, Suite 1818 Houston, TX 77002 DD=09/16/96 MA= / /
Albendazole TN= Albenza	Treatment of hydatid disease (cystic echinococcosis due to E. granulosus larvae or alveolar echinococcosis due to E. multilocularis larvae).	SmithKline Beecham Pharmaceuticals One Franklin Plaza P.O. Box 7929 Philadelphia, PA 19101 DD=01/17/96 MA=06/11/96
Albendazole TN= Albenza	Treatment of neurocysticercosis due to Taenia solium as: 1) chemotherapy of parenchymal, subarachnoidal and racemose (cysts in spinal fluid) neurocysticercosis in symptomatic cases and 2) prophylaxis of epilepsy and other sequelae in asymptomatic neurocysticercosis.	SmithKline Beecham Pharmaceuticals One Franklin Plaza P.O. Box 7929 Philadelphia, PA 19101 DD=01/18/96 MA=06/11/96
Allopurinol sodium TN= Zylprim for Injection	Management of patients with leukemia, lymphoma, and solid tumor malignancies who are receiving cancer therapy which causes elevations of serum and urinary uric acid levels and who cannot tolerate oral therapy.	Glaxo Wellcome Inc. Five Moore Drive P.O. Box 13398 Research Triangle Park, NC 27709 DD=10/16/92 MA=05/17/96
Amphotericin B lipid complex TN= Abelcet	Treatment of invasive fungal infections.	The Liposome Company, Inc. One Research Way Princeton, NJ 08540 DD=12/03/96 MA=10/18/96
Amphotericin B lipid complex TN= Abelcet	Treatment of invasive candidiasis.	The Liposome Company, Inc. One Research Way Princeton, NJ 08540 DD=06/27/96 MA= / /
Amphotericin B lipid complex TN= Abelcet	Treatment of invasive zygomycosis.	The Liposome Company, Inc. One Research Way Princeton, NJ 08540 DD=05/06/96 MA= / /
Amphotericin B lipid complex TN= Abelcet	Treatment of invasive coccidioidomycosis.	The Liposome Company, Inc. One Research Way Princeton, NJ 08540 DD=05/06/96 MA= / /
Amphotericin B lipid complex TN= Abelcet	Treatment of invasive sporotrichosis.	The Liposome Company, Inc. One Research Way Princeton, NJ 08540 DD=09/23/96 MA= / /

LIST of ORPHAN PRODUCTS DESIGNATIONS and APPROVALS

January 1, 1996 thru December, 1996

NAME Generic Name TN=Trade Name	INDICATION DESIGNATED	SPONSOR & ADDRESS DD=Date Designated MA=Marketing Approval
Amphotericin B lipid complex TN= Abelcet	Treatment of invasive protothecosis.	The Liposome Company, Inc. One Research Way Princeton, NJ 08540 DD=08/21/96 MA= / /
Antihemophilic factor (human) TN= Alphanate	Treatment of von Willebrand's disease.	Alpha Therapeutic Corporation 5555 Valley Boulevard Los Angeles, CA 90032 DD=01/05/96 MA= / /
Arcitumomab TN= 99m Tc-labeled CEA-Scan	Diagnosis and localization of primary, residual, recurrent and metastatic medullary thyroid carcinoma.	Immunomedics, Inc. 300 American Road Morris Plains, NJ 07950 DD=05/10/96 MA= / /
Betaine TN= Cystadane	Treatment of homocystinuria.	Orphan Medical 13911 Ridgedale Drive, Suite 475 Minnetonka, MN 55305 DD=05/16/94 MA=10/25/96
Bleomycin sulfate TN= Bleoxane	Treatment of malignant pleural effusion.	Bristol-Myers Squibb P.O. Box 4000 Princeton, NJ 08543 DD=09/17/93 MA=02/20/96
Buffered intrathecal electrolyte/dextrose injection TN= Elliotts B Solution	For use as a diluent in the intrathecal administration of methotrexate and cytarabine for the prevention or treatment of meningeal leukemia and lymphocytic lymphoma.	Orphan Medical 13911 Ridgedale Drive Minnetonka, MN 55305 DD=08/24/94 MA=09/27/96
C1 esterase inhibitor (human) TN=	Treatment and prevention of angioedema caused by C1-esterase inhibitor deficiency.	Alpha Therapeutic Corporation 5555 Valley Boulevard Los Angeles, CA 90032 DD=08/21/96 MA= / /
Clonidine TN= Duraclon	For continuous epidural administration as adjunctive therapy with intraspinal opiates for the treatment of pain in cancer patients tolerant to, or unresponsive to, intraspinal opiates.	Fujisawa USA, Inc. 3 Parkway North Center Deerfield, IL 60015 DD=01/24/89 MA=10/02/96
Clostridial collagenase TN=	Treatment of advanced (involutional or residual stage) Dupuytren's disease.	Hurst, L. M.D. & Badalamente, M. Ph.D. State University of New York at Stony Brook School of Medicine, Health Sciences Center T18-020 Stony Brook, NY 11794 DD=05/23/96 MA= / /
Collagenase (lyophilized) for injection TN= Plaques	Treatment of Peyronie's disease.	Advance Biofactures Corporation 35 Wilbur Street Lynbrook, NY 11563 DD=03/12/96 MA= / /

LIST of ORPHAN PRODUCTS DESIGNATIONS and APPROVALS

January 1, 1996 thru December, 1996

NAME Generic Name TN=Trade Name	INDICATION DESIGNATED	SPONSOR & ADDRESS DD=Date Designated MA=Marketing Approval
Corticotropin ovine trifluate TN= Acthrel	For use in differentiating pituitary and ectopic production of ACTH in patients with ACTH-dependent Cushing's syndrome.	Ferring Laboratories, Inc. 400 Rella Boulevard, Suite 201 Suffern, NY 10901 DD=11/24/89 MA=05/23/96
DAB389IL-2 TN=	Treatment of cutaneous T-cell lymphoma.	Scragen, Inc. 97 South Street Hopkinton, MA 01748 DD=08/21/96 MA= / /
DMP 777 TN=	Therapeutic management of patients with lung disease attributable to cystic fibrosis.	DuPont Merck Pharmaceutical Company Dupont Merck Plaza, Maple Run 2110 Wilmington, DE 19805 DD=06/04/96 MA= / /
Doxorubicin citrate liposome injection TN= DoxoXomic	Treatment of patients with advanced HIV-associated Kaposi's sarcoma.	NeXstar Pharmaceuticals, Inc. 650 Cliffside Drive San Dimas, CA 91773 DD=05/14/93 MA=04/08/96
Dihydrotestosterone TN= Androgel -DHT	Treatment of weight loss in AIDS patients with HIV-associated wasting.	Unimed Pharmaceuticals, Inc. 2150 East Lake Cook Road, Suite 210 Buffalo Grove, IL 60089 DD=02/05/96 MA= / /
Etiololanedione TN=	Treatment of Prader-Willi syndrome.	Supergen, Inc. 3158 Des Plaines Avenue Suite 10 Des Plaines, IL 60018 DD=05/07/96 MA= / /
Filgrastim TN= Neupogen	Reduction in the duration of neutropenia, fever, antibiotic use, and hospitalization, following induction and consolidation treatment for acute myeloid leukemia.	Amgen, Inc. 1840 DeHavilland Drive Thousand Oaks, CA 91320 DD=11/07/96 MA= / /
Fosphenytoin TN=	For the acute treatment of patients with status epilepticus of the grand mal type.	Warner-Lambert Company 2800 Plymouth Road Ann Arbor, MI 48106 DD=06/04/91 MA=08/05/96
Ganciclovir intravitreal implant TN= Vitrasert Implant	Treatment of cytomegalovirus retinitis.	Chiron Vision 9342 Jeronimo Road Irvine, CA 92718 DD=06/07/95 MA=03/04/96
Glatiramer acetate TN= Copaxone	Treatment of multiple sclerosis.	Teva Pharmaceuticals USA 1510 Delp Drive Kulpsville, PA 19443 DD=11/09/87 MA=12/20/96

LIST of ORPHAN PRODUCTS DESIGNATIONS and APPROVALS

January 1, 1996 thru December, 1996

NAME Generic Name TN=Trade Name	INDICATION DESIGNATED	SPONSOR & ADDRESS DD=Date Designated MA=Marketing Approval
Gusperimus TN= Spanidin	Treatment of acute renal graft rejection episodes.	Bristol-Myers Squibb Company 5 Research Parkway P.O. Box 5100 Wallingford, CT 06492 DD=06/27/96 MA= / /
Human acid alpha-glucosidase TN=	Treatment of glycogen storage disease type II.	Pharming B.V. Niels Bohrweg 11-13 2333 CA Leiden The Netherlands, DD=09/10/96 MA= / /
Ibuprofen i.v. solution TN= Salprofen	Prevention of patent ductus arteriosus.	Farnacon, Inc. 90 Grove Street, Suite 109 Ridgefield, CT 06877 DD=10/29/96 MA= / /
Ibuprofen i.v. solution TN=	Treatment of patent ductus arteriosus.	Farnacon, Inc. 90 Grove Street Ridgefield, CT 06877 DD=10/29/96 MA= / /
Idoxuridine TN=	Treatment of nonparenchymatous sarcomas.	NeoPharm, Inc. 225 East Decrpath, Suite 250 Lake Forest, IL 60045 DD=04/08/96 MA= / /
Imexon TN=	Treatment of multiple myeloma.	Amplimed Corporation 2321 Camino La Zorrera Tucson, AZ 85718 DD=11/08/96 MA= / /
Interferon beta-1a TN= Avonex	Treatment of multiple sclerosis.	Biogen, Inc. 14 Cambridge Center Cambridge, MA 02142 DD=12/16/91 MA=05/17/96
Interferon beta-1a TN= Rebif	Treatment of patients with secondary progressive multiple sclerosis.	Serono Laboratories, Inc. 100 Longwater Circle Norwell, MA 02061 DD=03/11/96 MA= / /
Interferon gamma-1b TN= Actimmune	Treatment of severe congenital osteopetrosis.	Genentech, Inc. 460 Point San Bruno Boulevard South San Francisco, CA 94080 DD=09/30/96 MA= / /
KL4-surfactant TN=	Treatment of meconium aspiration syndrome in newborn infants.	Acute Therapeutics, Inc. 3359 Durham Road Doylestown, PA 18901 DD=07/30/96 MA= / /

LIST of ORPHAN PRODUCTS DESIGNATIONS and APPROVALS

January 1, 1996 thru December, 1996

NAME Generic Name TN=Trade Name	INDICATION DESIGNATED	SPONSOR & ADDRESS DD=Date Designated MA=Marketing Approval
L-2-oxothiazolidine-4-carboxylic acid TN= Procyteine	Treatment of amyotrophic lateral sclerosis.	Transcend Therapeutics, Inc. 640 Memorial Drive, 3rd Floor West Cambridge, MA 02139 DD=07/30/96 MA= / /
Leflunomide TN=	Prevention of acute and chronic rejection in patients who have received solid organ transplants.	James W. Williams, M.D. 655 Superior Oak Park, IL 60302 DD=10/18/96 MA= / /
Lipid/DNA human cystic fibrosis gene TN=	Treatment of cystic fibrosis.	Genzyme Corporation One Kendall Square Cambridge, MA 02139 DD=04/08/96 MA= / /
Liposomal amphotericin B TN= AmBisome	Treatment of cryptococcal meningitis.	Fujisawa USA, Inc. 3 Parkway North Center Deerfield, IL 60015 DD=12/10/96 MA= / /
Liposomal amphotericin B TN= AmBisome	Treatment of visceral leishmaniasis.	Fujisawa USA, Inc. 3 Parkway North Center Deerfield, IL 60015 DD=12/06/96 MA= / /
Liposomal amphotericin B TN= AmBisome	Treatment of histoplasmosis.	Fujisawa USA, Inc. 3 Parkway North Center Deerfield, IL 60015 DD=12/10/96 MA= / /
Liposomal prostaglandin E1 injection TN=	Treatment of acute respiratory distress syndrome.	The Liposome Company, Inc. One Research Way Princeton, NJ 08540 DD=04/25/96 MA= / /
Methionine/L-methionine TN=	Treatment of AIDS myelopathy.	Di Rocco, Alessandro M.D. The Mount Sinai Medical Center One Gustave L. Levy Place, Box 1139 New York, NY 10029 DD=08/21/96 MA= / /
Methylnaltrexone TN=	Treatment of chronic opioid-induced constipation unresponsive to conventional therapy.	The University of Chicago 5841 South Maryland Avenue MC 4028 Chicago, IL 60637 DD=06/17/96 MA= / /
Midodrine HCl TN= Amatine	Treatment of patients with symptomatic orthostatic hypotension.	Roberts Pharmaceutical Corp. Meridian Center III 6 Industrial Way West Eatontown, NJ 07724 DD=12/05/96 MA=09/06/96

LIST of ORPHAN PRODUCTS DESIGNATIONS and APPROVALS

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January 1, 1996 thru December, 1996

NAME Generic Name TN=Trade Name	INDICATION DESIGNATED	SPONSOR & ADDRESS DD=Date Designated MA=Marketing Approval
Mitoxantrone TN= Novantrone	Treatment of hormone refractory prostate cancer.	Immunex Corporation 51 University Street Seattle, WA 98101 DD=08/21/96 MA=11/13/96
Monoclonal antibody-B43.13 TN= Ovarex MAb-B43.13	Treatment of epithelial ovarian cancer.	AltaRex, Inc. 1134 Dentistry-Pharmacy Building University of Alberta Edmonton, Alberta, Canada, DD=11/25/96 MA= / /
N-acetyl-procainamide TN=	Prevention of life-threatening ventricular arrhythmias in patients with documented procainamide-induced lupus.	NAPA of the Bahamas 3560 Pennsylvania Avenue, Suite 7 Dubuque, IA 52002 DD=12/10/96 MA= / /
Nitazoxanide TN=	Treatment of immunocompromised patients with cryptosporidiosis.	Unimed Pharmaceuticals, Inc. 2150 East Lake Cook Road, Suite 210 Buffalo Grove, IL 60089 DD=12/12/96 MA= / /
Ofloxacin TN= Ocuflax Ophthalmic Solution	Treatment of bacterial corneal ulcers.	Allergan, Inc. 2525 Dupont Drive P.O. Box 19534 Irvine, CA 92713 DD=04/18/91 MA=05/22/96
Pentosan polysulfate sodium TN= Elmiron	Treatment of interstitial cystitis.	Baker Norton Pharmaceuticals 4400 Biscayne Boulevard Miami, FL 33137 DD=08/07/85 MA=09/26/96
Polifeprosan 20 with carmustine TN= Gliadel	Treatment of malignant glioma.	Guilford Pharmaceuticals, Inc. 6611 Tributary Street Baltimore, MD 21224 DD=12/13/89 MA=09/23/96
Porcine fetal neural dopaminergic cells and/or precursors aseptically prepared and coated with anti-MHC-I Ab for intracerebral implantation TN= NeuroCell-PD	Treatment of Hoehn and Yahr stage 4 and 5 Parkinson's disease.	Diacrin, Inc. Building 39, 13th Street Charlestown, MA 02129 DD=12/17/96 MA= / /
Porcine fetal neural dopaminergic cells and/or precursors aseptically prepared for intracerebral implantation. TN= NeuroCell-PD	Treatment of Hoehn and Yahr stage 4 and 5 Parkinson's disease.	Diacrin, Inc. Building 96, 13th Street Charlestown Navy Yard Charlestown, MA 02129 DD=12/17/96 MA= / /

LIST of ORPHAN PRODUCTS DESIGNATIONS and APPROVALS

January 1, 1996 thru December, 1996

NAME Generic Name TN=Trade Name	INDICATION DESIGNATED	SPONSOR & ADDRESS DD=Date Designated MA=Marketing Approval
Porcine fetal neural gabaergic cells and/or precursors aseptically prepared and coated with anti-MHC-1 Ab for intracerebral implantation TN= NeuroCell-HD	Treatment of Huntington's disease.	Diacrin, Inc. Building 96, 13th Street Charlestown, MA 02129 DD=12/10/96 MA= / /
Porcine fetal neural gabaergic cells and/or precursors aseptically prepared for intracerebral implantation for Huntington's disease. TN= NeuroCell-HD	Treatment of Huntington's disease.	Diacrin, Inc. Building 96, 13th Street Charlestown Navy Yard Charlestown, MA 02129 DD=12/10/96 MA= / /
Prostaglandin E1 in lipid emulsion TN=	Treatment of ischemic ulceration of the lower limbs due to peripheral arterial disease.	Alpha Therapeutic Corporation 5555 Valley Boulevard Los Angeles, CA 90032 DD=09/10/96 MA= / /
R-VIII SQ TN= REFECTO	For long-term and/or hospital treatment of hemophilia A or for treatment of patients with hemophilia A in connection with surgical procedures.	Pharmacia & Upjohn 7000 Portage Road Kalamazoo, MI 49001 DD=02/08/96 MA= / /
Recombinant human interleukin-11 TN= Neumega rhIL-11 Growth Factor	Prevention of severe chemotherapy-induced thrombocytopenia.	Genetics Institute, Inc. 87 Cambridge Park Drive Cambridge, MA 02140 DD=12/17/96 MA= / /
Respiratory syncytial virus immune globulin (human) TN= Respigam	Prophylaxis of respiratory syncytial virus lower respiratory tract infections in infants and young children at high risk of RSV disease.	MedImmune, Inc. 35 West Watkins Mill Road Gaithersburg, MD 20878 DD=09/27/90 MA=01/18/96
Rifapentine TN=	Prophylactic treatment of Mycobacterium avium complex in patients with acquired immunodeficiency syndrome and a CD4+ count less than or equal to 75/mm3.	Marion Merrell Dow Inc. P.O. Box 9627 (Park A) Kansas City, MO 64137 DD=03/12/96 MA= / /
Riluzole TN= Rilutek	Treatment of Huntington's disease.	Rhone-Poulenc Rorer Pharmaceuticals, Inc. 500 Arcola Road Collegeville, PA 19426 DD=10/15/96 MA= / /
SU101 TN=	Treatment of ovarian cancer.	Sugen, Inc. 515 Galveston Drive Redwood City, CA 94063 DD=03/12/96 MA= / /
Sodium phenylbutyrate TN= Buphenyl	Treatment of urea cycle disorders: carbamylphosphate synthetase deficiency, ornithine transcarbamylase deficiency, and argininosuccinic acid synthetase deficiency.	Ucyclyd Pharma 10819 Gilroy Road Suite 100 Hunt Valley, MD 21031 DD=11/22/93 MA=04/30/96

LIST of ORPHAN PRODUCTS DESIGNATIONS and APPROVALS

January 1, 1996 thru December, 1996

NAME Generic Name TN=Trade Name		INDICATION DESIGNATED	SPONSOR & ADDRESS DD=Date Designated MA=Marketing Approval
Somatropin for injection TN= Nutropin		Treatment of short stature associated with Turner's syndrome.	Genentech, Inc. 460 Point San Bruno Boulevard South San Francisco, CA 94080 DD=03/23/89 MA=12/30/96
Somatropin for injection TN= Serostim		Treatment of AIDS-associated catabolism/weight loss.	Serono Laboratories, Inc. 100 Longwater Circle Norwell, MA 02061 DD=11/15/91 MA=08/23/96
Somatropin for injection TN= Serostim		Treatment of children with AIDS-associated failure-to-thrive including AIDS-associated wasting.	Serono Laboratories, Inc. 100 Longwater Circle Norwell, MA 02061 DD=03/26/96 MA= / /
Somatropin for injection TN= Nutropin		As replacement therapy for growth hormone deficiency in adults after epiphyseal closure.	Genentech, Inc. 460 Point San Bruno Boulevard South San Francisco, CA 94080 DD=11/18/96 MA= / /
Testosterone TN= Androgel		Treatment of weight loss in AIDS patients with HIV-associated wasting.	Unimed Pharmaceuticals, Inc. 2150 East Lake Cook Road Buffalo Grove, IL 60089 DD=02/05/96 MA= / /
Thalidomide TN= Synovir		Treatment of HIV-associated wasting syndrome.	Celgene Corporation P.O. Box 4914 7 Powder Horn Drive Warren, NJ 07059 DD=03/11/96 MA= / /
Uridine 5'-triphosphate TN=		To facilitate the removal of lung secretions in the treatment of patients with primary ciliary dyskinesia.	Inspire Pharmaceuticals, Inc. 4222 Emperor Boulevard, Suite 470 Durham, NC 27703 DD=06/26/96 MA= / /
Valine, isoleucine and leucine TN= VIL		Treatment of hyperphenylalaninemia.	Leas Research Products 4 Brookview Lane Troy, NY 12180 DD=01/05/96 MA= / /

DRUG PRODUCTS WHICH MUST DEMONSTRATE *IN VIVO* BIOAVAILABILITY ONLY
IF PRODUCT FAILS TO ACHIEVE ADEQUATE DISSOLUTION

NO DECEMBER 1996 ADDITIONS

BIOPHARMACEUTIC GUIDANCE AVAILABILITY

<u>DRUG NAME (DOSAGE FORM)</u>	<u>DATE</u>	<u>REVISED DATE</u>
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THE FOLLOWING IS A LIST OF GUIDANCES AVAILABLE FOR *IN VIVO* BIOEQUIVALENCE STUDIES AND *IN VITRO* DISSOLUTION TESTING. COMMENTS AND SUGGESTIONS CONCERNING THESE GUIDANCES ARE ENCOURAGED AND SHOULD BE SENT TO THE DIVISION OF BIOEQUIVALENCE (HFD-650, MPN-2 ROOM 279) 7500 STANDISH PLACE, ROCKVILLE, MD 20855. COPIES OF THESE GUIDANCES MAY ALSO BE OBTAINED FROM THE DIVISION OF COMMUNICATION MANAGEMENT, CENTER FOR DRUG EVALUATION AND RESEARCH, FDA, 5600 FISHERS LANE (HFD-210) ROCKVILLE, MD 20857 OR BY CALLING (301) 827-4573, FAX: (301) 827-4577.

REFER BACK TO THE APPROVED DRUG PRODUCTS WITH THERAPEUTIC EQUIVALENCE EVALUATIONS, 16TH EDITION FOR A FULL LISTING OF BIOPHARMACEUTIC GUIDANCE AVAILABILITY DATA. ONLY NEW DATA WILL BE ADDED TO THE CUMULATIVE SUPPLEMENT.

CLOZAPINE <i>IN VITRO</i> AND <i>IN VIVO</i> (TABLET)

NOV 15, 1995

NOV 15, 1996

ANDA SUITABILITY PETITIONS

PETITIONS APPROVED

DRUG NAME DOSAGE FORM; ROUTE	STRENGTH (CONTAINER SIZE)	DOCKET NUMBER	PETITIONER	REASON FOR PETITION	STATUS
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THE FOLLOWING ARE TWO LISTS OF PETITIONS FILED UNDER SECTION 505(j)(2)(C) OF THE ACT WHERE THE AGENCY HAS DETERMINED THAT THE REFERENCED PRODUCT: (1) IS SUITABLE FOR SUBMISSION AS AN ANDA (PETITIONS APPROVED) OR (2) IS NOT SUITABLE FOR SUBMISSION AS AN ANDA (PETITIONS DENIED). THE DETERMINATION THAT AN ANDA WILL BE APPROVED IS NOT MADE UNTIL THE ANDA ITSELF IS SUBMITTED AND REVIEWED BY THE AGENCY. A COPY OF EACH PETITION IS LISTED BY DOCKET NUMBER ON PUBLIC DISPLAY IN FDA'S DOCKETS MANAGEMENT BRANCH, HFA-305, ROOM 1-23, PARK BUILDING, 5600 FISHERS LANE, ROCKVILLE, MD 20857.

REFER BACK TO THE APPROVED DRUG PRODUCTS WITH THERAPEUTIC EQUIVALENCE EVALUATIONS, 16TH EDITION FOR A FULL LISTING OF ANDA SUITABILITY PETITIONS DATA. ONLY NEW DATA WILL BE ADDED TO THE CUMULATIVE SUPPLEMENT.

ACETAMINOPHEN; BUTALBITAL; CAFFEINE SOLUTION; ORAL	325MG/15ML 50MG/15ML 40MG/15ML	96 P-0208/ CP1	MIKART	NEW DOSAGE FORM	APPROVED OCT 10, 1996
ACETAMINOPHEN; BUTALBITAL; CAFFEINE; HYDROCODONE BITARTRATE CAPSULE; ORAL	325MG 50MG 40MG 7.5MG	95 P-0279/ CP2	MIKART	NEW COMBINATION NEW DOSAGE FORM NEW STRENGTH	APPROVED MAR 08, 1996
ACETAMINOPHEN; BUTALBITAL; CAFFEINE; HYDROCODONE BITARTRATE CAPSULE; ORAL	325MG 50MG 40MG 10MG	95 P-0279/ CP1	MIKART	NEW COMBINATION NEW DOSAGE FORM NEW STRENGTH	APPROVED MAR 08, 1996
ACETAMINOPHEN; BUTALBITAL; CAFFEINE; HYDROCODONE BITARTRATE CAPSULE; ORAL	500MG 50MG 40MG 7.5MG	95 P-0279/ CP4	MIKART	NEW COMBINATION NEW DOSAGE FORM NEW STRENGTH	APPROVED MAR 08, 1996
ACETAMINOPHEN; BUTALBITAL; CAFFEINE; HYDROCODONE BITARTRATE CAPSULE; ORAL	500MG 50MG 40MG 10MG	95 P-0279/ CP3	MIKART	NEW COMBINATION NEW DOSAGE FORM NEW STRENGTH	APPROVED MAR 08, 1996
ACETAMINOPHEN; BUTALBITAL; CAFFEINE; HYDROCODONE BITARTRATE TABLET; ORAL	325MG 50MG 40MG 7.5MG	95 P-0279/ CP2	MIKART	NEW COMBINATION NEW DOSAGE FORM NEW STRENGTH	APPROVED MAR 08, 1996
ACETAMINOPHEN; BUTALBITAL; CAFFEINE; HYDROCODONE BITARTRATE TABLET; ORAL	325MG 50MG 40MG 10MG	95 P-0279/ CP1	MIKART	NEW COMBIANTION NEW DOSAGE FORM NEW STRENGTH	APPROVED MAR 08. 1996
ACETAMINOPHEN; BUTALBITAL; CAFFEINE; HYDROCODONE BITARTRATE TABLET; ORAL	500MG 50MG 40MG 7.5MG	95 P-0279/ CP4	MIKART	NEW COMBINATION NEW DOSAGE FORM NEW STRENGTH	APPROVED MAR 08, 1996

ANDA SUITABILITY PETITIONS

PETITIONS APPROVED

DRUG NAME DOSAGE FORM; ROUTE	STRENGTH (CONTAINER SIZE)	DOCKET NUMBER	PETITIONER	REASON FOR PETITION	STATUS
ACETAMINOPHEN; BUTALBITAL; CAFFEINE; HYDROCODONE BITARTRATE TABLET; ORAL	500MG 50MG 40MG 10MG	95 P-0279/ CP3	MIKART	NEW COMBINATION NEW DOSAGE FORM NEW STRENGTH	APPROVED MAR 08, 1996
ACETAMINOPHEN; HYDROCODONE BITARTRATE CAPSULE; ORAL	325MG 5MG	95 P-0278/ CP1	MIKART	NEW STRENGTH	APPROVED MAY 28, 1996
ACYCLOVIR SODIUM INJECTABLE; INJECTION	EQ 5MG BASE/ML (100ML/CONTAINER) (200ML/CONTAINER)	95 P-0268/ CP1	WILMER, CUTLER, & PICKERING	NEW DOSAGE FORM NEW STRENGTH	APPROVED FEB 27, 1996
ACYCLOVIR SODIUM INJECTABLE; INJECTION	EQ 250MG/VIAL	96 P-0284/ CP1	ABBOTT	NEW STRENGTH	APPROVED OCT 10, 1996
ASPIRIN; BUTALBITAL CAPSULE; ORAL	650MG 50MG	96 P-0021/ CP1	SAVAGE	NEW DOSAGE FORM	APPROVED APR 19, 1996
ATRAURIUM BESYLATE INJECTABLE; INJECTION	0.5MG/ML 1MG/ML (100ML CONTAINER)	95 P-0372/ CP1	ABBOTT	NEW STRENGTH	APPROVED MAR 08, 1996
CARBIDOPA; LEVODOPA POWDER FOR RECONSTITUTION; ORAL	25MG/PACKET 100MG/PACKET	95 P-0100/ CP1	ATHENA	NEW DOSAGE FORM	APPROVED MAY 28, 1996
CARBIDOPA; LEVODOPA POWDER FOR RECONSTITUTION; ORAL	25MG/PACKET 250MG/PACKET	95 P-0100/ CP1	ATHENA	NEW DOSAGE FORM	APPROVED MAY 28, 1996
CHOLESTYRAMINE TABLET, CHEWABLE; ORAL	EQ 2GM RESIN	95 P-0277/ CP1	MAYRAND	NEW DOSAGE FORM NEW STRENGTH	APPROVED FEB 27, 1996
CYTARABINE INJECTABLE; INJECTION	100MG/ML (1ML/VIAL) (5ML/VIAL)	92 P-0183/ CP1	FAULDING	NEW DOSAGE FORM NEW STRENGTH	APPROVED JUL 26, 1996
CYTARABINE INJECTABLE; INJECTION	100MG/ML (10ML/VIAL) (20ML/VIAL)	92 P-0184/ CP1	FAULDING	NEW DOSAGE FORM	APPROVED JUL 26, 1996
DILTIAZEM HYDROCHLORIDE INJECTABLE, INJECTION	5MG/ML (25ML/SYRINGE) (50ML/SYRINGE)	95 P-0196/ CP1	INTL MEDICATION	NEW STRENGTH	APPROVED FEB 27, 1996
EPINEPHRINE INJECTABLE; SUBCUTANEOUS	0.3MG/DELIVERY	95 P-0190/ CP1	SENETCK PLC	NEW ROUTE OF ADMINISTRATION	APPROVED FEB 15, 1996
ETOPOSIDE CAPSULE; ORAL	25MG	95 P-0142/ CP1	GUIDELINES	NEW STRENGTH	APPROVED SEP 12, 1996

ANDA SUITABILITY PETITIONS

PETITIONS APPROVED

DRUG NAME DOSAGE FORM; ROUTE	STRENGTH (CONTAINER SIZE)	DOCKET NUMBER	PETITIONER	REASON FOR PETITION	STATUS
HYDROCORTISONE BUTYRATE LOTION; TOPICAL	0.1%	95 P-0223/ CP1	MCKENNA & CUNEO	NEW DOSAGE FORM	APPROVED FEB 21, 1996
LACTULOSE CRYSTALS, FOR RECONSTITUTION; ORAL	20GM/PACKET	95 P-0287/ CP1	BENNETT	NEW DOSAGE FORM NEW STRENGTH	APPROVED APR 19, 1996
MEPERIDINE HYDROCHLORIDE INJECTABLE; INJECTION	10MG/ML (60ML/SYRINGE)	95 P-0348/ CP1	MALLINCKRODT	NEW STRENGTH	APPROVED MAR 08, 1996
METRONIDAZOLE LOTION; TOPICAL	0.75%	95 P-0328/ CP1	RNB PHARM	NEW DOSAGE FORM	APPROVED FEB 23, 1996
NIFEDIPINE CAPSULE, EXTENDED RELEASE; ORAL	30MG 60MG 90MG	95-P-0326/ CP1	KV	NEW DOSAGE FORM	APPROVED FEB 23, 1996
PACLITAXEL INJECTABLE; INJECTION	6MG/ML (16.7ML/VIAL) (33.3ML/VIAL) (50ML/VIAL)	95 P-0360/ CP1	ABBOTT	NEW STRENGTH	APPROVED APR 29, 1996
PENTOXIFYLLINE SUSPENSION, EXTENDED RELEASE; ORAL	400MG/PACKET	96 P-0079/ CP1	KV PHARM	NEW DOSAGE FORM	APPROVED AUG 13, 1996
POTASSIUM CHLORIDE CAPSULE, EXTENDED RELEASE; ORAL	20MEQ	96 P-0018/ CP1	KV PHARM	NEW DOSAGE FORM	APPROVED AUG 19, 1996
POTASSIUM CHLORIDE SUSPENSION, EXTENDED RELEASE; ORAL	10MEQ	96 P-0054/ CP1	KV PHARM	NEW DOSAGE FORM	APPROVED AUG 19, 1996

ANDA SUITABILITY PETITIONS

PETITIONS DENIED

DRUG NAME DOSAGE FORM; ROUTE	STRENGTH (CONTAINER SIZE)	DOCKET NUMBER	PETITIONER	REASON FOR PETITION	STATUS
HYDROCODONE BITRATRATE; PHENYLEPHRINE HYDROCHLORIDE SOLUTION; ORAL	2.5MG/5ML 5MG/5ML	95 P-0336/ CP1	BOCK PHARMA	NEW COMBINATION	DENIED AUG 19, 1996
HYDROCODONE BITRATRATE; PHENYLEPHRINE HYDROCHLORIDE SOLUTION; ORAL	5MG/5ML 10MG/5ML	95 P-0336/ CP2	BOCK PHARMA	NEW COMBINATION	DENIED AUG 19, 1996

PRESCRIPTION AND OTC DRUG PRODUCT
PATENT AND EXCLUSIVITY DATA

APPL/PROD NUMBER	INGREDIENT NAME; TRADE NAME	PATENT NUMBER	PATENT EXPIRES	USE CODE	EXCLUS CODE	EXCLUS EXPIRES
19821 001	ACITRETIN; SORIATANE				NE	OCT 28, 1999
19821 002	ACITRETIN; SORIATANE				NE	OCT 28, 1999
19806 001	ACRIVASTINE; SEMPREX-D	4650807	MAR 26, 2008	U-93		
20338 001	ADAPALENE; DIFFERIN	4717720	APR 10, 2006	U-134	NCE	MAY 31, 2001
20380 001	ADAPALENE; DIFFERIN	4717720	APR 10, 2006	U-134	NCE	MAY 31, 2001
20666 001	ALBENDAZOLE; ALBENZA				ODE	JUN 11, 2003
					NCE	JUN 11, 2001
					NC	OCT 24, 1999
20291 001	ALBUTEROL SULFATE; COMBIVENT					
20503 001	ALBUTEROL SULFATE; PROVENTIL-HFA	5225183	JUL 06, 2010			
20298 001	ALLOPURINOL SODIUM; ZYLOPRIM				NDF	MAY 17, 1999
					ODE	MAY 17, 2003
20700 001	ALPROSTADIL; MUSE	5474535	DEC 12, 2012	U-155	NDF	NOV 19, 1999
		5242391	SEP 07, 2010	U-155		
		4801587	JAN 31, 2007	U-155		
20700 002	ALPROSTADIL; MUSE	5474535	DEC 12, 2012	U-155	NDF	NOV 19, 1999
		5242391	SEP 07, 2010	U-155		
		4801587	JAN 31, 2007	U-155		
20700 003	ALPROSTADIL; MUSE	5474535	DEC 12, 2012	U-155	NDF	NOV 19, 1999
		5242391	SEP 07, 2010	U-155		
		4801587	JAN 31, 2007	U-155		
20700 004	ALPROSTADIL; MUSE	5474535	DEC 12, 2012	U-155	NDF	NOV 19, 1999
		5242391	SEP 07, 2010	U-155		
		4801587	JAN 31, 2007	U-155		
20221 001	AMIFOSTINE; ETHYOL				I-149	MAR 15, 1999
>ADD>	20511 001	AMLEXANOX; APHTHASOL			NCE	DEC 17, 2001
	19787 001	AMLODIPINE BESYLATE; NORVASC	4879303	MAR 25, 2007	I-156	JUN 14, 1999
	19787 002	AMLODIPINE BESYLATE; NORVASC	4879303	MAR 25, 2007	I-156	JUN 14, 1999
	19787 003	AMLODIPINE BESYLATE; NORVASC	4879303	MAR 25, 2007	I-156	JUN 14, 1999
	20508 001	AMMONIUM LACTATE; LAC-HYDRIN			NDF	AUG 29, 1999
>ADD>	50724 001	AMPHOTERICIN B; ABELCET			ODE	OCT 18, 2003
	20541 001	ANASTROZOLE; ARIMIDEX	4935437	JUN 10, 2008		
>ADD>	20702 001	ATORVASTATIN CALCIUM; LIPITOR			NCE	DEC 17, 2001
>ADD>	20702 002	ATORVASTATIN CALCIUM; LIPITOR			NCE	DEC 17, 2001
>ADD>	20702 003	ATORVASTATIN CALCIUM; LIPITOR			NCE	DEC 17, 2001
	20428 001	AZELAIC ACID; AZELEX	4386104	MAY 31, 2000	U-124	
	20114 001	AZELASTINE HYDROCHLORIDE; ASTELIN			NCE	NOV 01, 2001
	20075 001	BACLOFEN; LIORESAL			I-153	JUN 14, 1999
	20075 002	BACLOFEN; LIORESAL			I-153	JUN 14, 1999
	20469 001	BECLOMETHASONE DIPROPIONATE MONOHYDRATE; VANCENASE AQ			NP	JUN 26, 1999
	20532 001	BENTOQUATAM; IVY BLOCK			NCE	AUG 26, 2001
	20576 001	BETAINE, ANHYDROUS; CYSTADANE			NCE	OCT 25, 2001
					ODE	OCT 25, 2003

PRESCRIPTION AND OTC DRUG PRODUCT
PATENT AND EXCLUSIVITY DATA

APPL/PROD NUMBER	INGREDIENT NAME; TRADE NAME	PATENT NUMBER	PATENT EXPIRES	USE CODE	EXCLUS CODE	EXCLUS EXPIRES
20498 001	BICALUTAMIDE; CASODEX	4636505	JAN 13, 2004			
50443 001	BLEOMYCIN SULFATE; BLENOXANE				ODE	FEB 20, 2003
20613 001	BRIMONIDINE TARTRATE; ALPHAGAN				NCE	SEP 06, 2001
19672 001	BROMPHENIRAMINE MALEATE; EFIDAC 24	4810502	MAR 14, 2006			
		4801461	MAR 14, 2006			
		4673405	MAR 18, 2003			
		4662880	MAR 14, 2006		NP	MAR 29, 1999
20358 001	BUPROPION HYDROCHLORIDE; WELLBUTRIN	5427798	AUG 12, 2013			
		5358970	AUG 12, 2013			
		RE33994	AUG 18, 2004			
20358 002	BUPROPION HYDROCHLORIDE; WELLBUTRIN	5427798	AUG 12, 2013			
		5358970	AUG 12, 2013			
		RE33994	AUG 18, 2004			
20358 003	BUPROPION HYDROCHLORIDE; WELLBUTRIN	5427798	AUG 12, 2013			
		5358970	AUG 12, 2013			
		RE33994	AUG 18, 2004			
18731 001	BUSPIRONE HYDROCHLORIDE; BUSPAR	5015646	MAY 14, 2008	U-13		
18731 002	BUSPIRONE HYDROCHLORIDE; BUSPAR	4182763	MAY 22, 2000	U-13		
		5015646	MAY 14, 2008	U-13		
		4182763	MAY 22, 2000	U-13		
18731 003	BUSPIRONE HYDROCHLORIDE; BUSPAR	5015646	MAY 14, 2008	U-13		
		4182763	MAY 22, 2000	U-13		
18731 004	BUSPIRONE HYDROCHLORIDE; BUSPAR	5015646	MAY 14, 2008	U-13		
		4182763	MAY 22, 2000	U-13		
20524 001	BUTENAFINE HYDROCHLORIDE; MENTAX				NCE	OCT 18, 2001
>ADD>					I-175	DEC 31, 1999
19215 001	BUTOCONAZOLE NITRATE; FEMSTAT	4078071	MAR 07, 1997			
19359 001	BUTOCONAZOLE NITRATE; FEMSTAT	4078071	MAR 07, 1997			
20421 001	BUTOCONAZOLE NITRATE; FEMSTAT 3	4078071	MAR 07, 1997		NP	DEC 21, 1998
>ADD>					NCE	DEC 23, 2001
20664 001	CABERGOLINE; DOSTINEX				NCE	DEC 29, 1998
20273 001	CALCIPOTRIENE; DOVONEX	4866048	DEC 29, 2007	U-88	NDF	JUL 22, 1999
20554 001	CALCIPOTRIENE; DOVONEX	4866048	SEP 12, 2006			
18874 001	CALCITRIOL; CALCIJEX	4308264	JAN 28, 2001			
18874 002	CALCITRIOL; CALCIJEX	4308264	JAN 28, 2001			
20577 001	CALCIUM CHLORIDE; ELLIOTTS B SOLUTION				ODE	SEP 27, 2003
18343 004	CAPTOPRIL; CAPOTEN				I-95	SEP 23, 1996
					I-101	JAN 28, 1997
18343 007	CAPTOPRIL; CAPOTEN				I-95	SEP 23, 1996
					I-101	JAN 28, 1997
20234 001	CARBAMAZEPINE; TEGRETOL-XR	5284662	FEB 08, 2011			
		RE34990	JUL 29, 2007			

PRESCRIPTION AND OTC DRUG PRODUCT
PATENT AND EXCLUSIVITY DATA

APPL/PROD NUMBER	INGREDIENT NAME; TRADE NAME	PATENT NUMBER	PATENT EXPIRES	USE CODE	EXCLUS CODE	EXCLUS EXPIRES
20234 002	CARBAMAZEPINE; TEGRETOL-XR	5284662	FEB 08, 2011			
		RE34990	JUL 29, 2007			
20234 003	CARBAMAZEPINE; TEGRETOL-XR	5284662	FEB 08, 2011			
		RE34990	JUL 29, 2007			
19880 001	CARBOPLATIN; PARAPLATIN	4140707	AUG 24, 1998			
19880 002	CARBOPLATIN; PARAPLATIN	4140707	AUG 24, 1998			
19880 003	CARBOPLATIN; PARAPLATIN	4140707	AUG 24, 1998			
>ADD> >ADD> >DLT>	20637 001	CARMUSTINE; GLIADEL	5179189	JAN 19, 2010	NP	SEP 23, 1999
			4789724	JUL 12, 2005	ODE	SEP 23, 2003
			4789724	JUL 12, 2005	ODE	AUG 23, 1999
	19835 001	CETIRIZINE HYDROCHLORIDE; ZYRTEC	4525358	JUN 25, 2002		
	19835 002	CETIRIZINE HYDROCHLORIDE; ZYRTEC	4525358	JUN 25, 2002		
	20346 001	CETIRIZINE HYDROCHLORIDE; ZYRTEC	4525358	JUN 25, 2002	NDF	SEP 27, 1999
					NCE	DEC 08, 2000
	20638 001	CIDOFOVIR; VISTIDE	5142051	AUG 25, 2009	NCE	JUN 26, 2001
	20238 002	CIMETIDINE; TAGAMET HB			NS	JUN 19, 1998
					I-137	NOV 15, 1998
	19847 001	CIPROFLOXACIN; CIPRO	4670444	DEC 09, 2003	I-167	OCT 10, 1999
	19857 001	CIPROFLOXACIN; CIPRO IN DEXTROSE 5%	4957922	SEP 18, 2007	I-167	OCT 10, 1999
	19858 001	CIPROFLOXACIN; CIPRO IN SODIUM CHLORIDE 0.9%	4670444	DEC 09, 2003	I-167	OCT 10, 1999
	19537 001	CIPROFLOXACIN HYDROCHLORIDE; CIPRO			I-157	APR 08, 1999
					I-164	SEP 11, 1999
					I-167	OCT 10, 1999
	19537 002	CIPROFLOXACIN HYDROCHLORIDE; CIPRO			I-157	APR 08, 1999
					I-164	SEP 11, 1999
					I-167	OCT 10, 1999
	19537 003	CIPROFLOXACIN HYDROCHLORIDE; CIPRO			I-157	APR 08, 1999
					I-164	SEP 11, 1999
					I-167	OCT 10, 1999
	19537 004	CIPROFLOXACIN HYDROCHLORIDE; CIPRO			I-157	APR 08, 1999
					I-164	SEP 11, 1999
					I-167	OCT 10, 1999
					NCE	JUL 29, 1998
	20398 001	CISAPRIDE MONOHYDRATE; PROPULSID	4962115	OCT 09, 2007	U-79	
	20551 001	CISATRACURIUM BESYLATE; NIMBEX	5453510	SEP 26, 2012	U-127	
			4179507	DEC 18, 1996	U-127	
	20551 002	CISATRACURIUM BESYLATE; NIMBEX PRESERVATIVE FREE	5453510	SEP 26, 2012	U-127	
			4179507	DEC 18, 1996	U-127	
	20551 003	CISATRACURIUM BESYLATE; NIMBEX PRESERVATIVE FREE	5453510	SEP 26, 2012	U-127	
			4179507	DEC 18, 1996	U-127	
	18057 001	CISPLATIN; PLATINOL	5562925	MAY 08, 2012		
	18057 002	CISPLATIN; PLATINOL	5562925	MAY 08, 2012		
	18057 003	CISPLATIN; PLATINOL-AQ	5562925	MAY 08, 2012		

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PATENT AND EXCLUSIVITY DATA

APPL/PROD NUMBER	INGREDIENT NAME; TRADE NAME	PATENT NUMBER	PATENT EXPIRES	USE CODE	EXCLUS CODE	EXCLUS EXPIRES
	18057 004 CISPLATIN; PLATINOL-AQ	5562925	MAY 08, 2012			
>ADD>	17661 003 CLEMASTINE FUMARATE; TAVIST-1				I-171	OCT 31, 1999
>DLT>	17661 003 CLEMASTINE FUMARATE; TAVIST-1				I-151	OCT 31, 1999
>ADD>	18298 002 CLEMASTINE FUMARATE; TAVIST-D				I-171	OCT 31, 1999
>DLT>	18298 002 CLEMASTINE FUMARATE; TAVIST-D				I-151	OCT 31, 1999
	20340 001 CLOBETASOL PROPIONATE; TEMOVATE E				D-32	MAY 03, 1999
	20615 001 CLONIDINE HYDROCHLORIDE; DURACLON				NDF	OCT 02, 1999
					ODE	OCT 02, 2003
	20525 001 CLOTTRIMAZOLE; GYNE-LOTTRIMIN 3				NP	JUL 29, 1999
	20526 002 CLOTTRIMAZOLE; GYNE-LOTTRIMIN 3 COMBINATION PACK				NP	JUL 29, 1999
>ADD>	20162 001 CORTICORELIN OVINE TRIFLUTATE; ACTHREL	4415558	JUN 08, 2001		NCE	MAY 23, 2001
	20479 001 CROMOLYN SODIUM; GASTROCROM	4515805	MAY 07, 2002	U-130		
		4421762	DEC 20, 2000	U-130		
>ADD>	19722 001 CYANOCOBALAMIN; NASCOBAL	4724231	APR 16, 2005	U-157		
	20287 001 DALTEPARIN SODIUM; FRAGMIN	4303651	JAN 04, 2005		NCE	DEC 22, 1999
					D-30	MAR 18, 1999
	20287 003 DALTEPARIN SODIUM; FRAGMIN	4303651	JAN 04, 2005		NCE	DEC 22, 1999
					D-30	MAR 18, 1999
>ADD>	20430 001 DANAPAROID SODIUM; ORGARAN				NCE	DEC 24, 2001
	50704 002 DAUNORUBICIN CITRATE; DAUNOXOME				ODE	APR 08, 2003
	20118 001 DESFLURANE; SUPRANE	4762856	FEB 02, 2007	U-67	NCE	SEP 18, 1997
	19955 001 DESMOPRESSIN ACETATE; DDAVP	5047398	SEP 10, 2008			
	19955 002 DESMOPRESSIN ACETATE; DDAVP	5047398	SEP 10, 2008			
	20071 001 DESOGESTREL; DESOGEN	3927046	NOV 19, 1995			
	20071 002 DESOGESTREL; DESOGEN	3927046	NOV 19, 1995			
	20301 001 DESOGESTREL; ORTHO-CEPT	3927046	NOV 19, 1995			
	20301 002 DESOGESTREL; ORTHO-CEPT	3927046	NOV 19, 1995			
	20344 001 DEXFENFLURAMINE HYDROCHLORIDE; REDUX	4309445	JUN 16, 2000	U-133		
	20037 001 DICLOFENAC SODIUM; VOLTAREN	4960799	OCT 02, 2007		I-163	JUL 23, 1999
	20254 001 DICLOFENAC SODIUM; VOLTAREN-XR				NDF	MAR 08, 1999
	20092 001 DILTIAZEM HYDROCHLORIDE; DILACOR XR	5422123	JUN 06, 2012			
	20092 002 DILTIAZEM HYDROCHLORIDE; DILACOR XR	5422123	JUN 06, 2012			
	20092 003 DILTIAZEM HYDROCHLORIDE; DILACOR XR	5422123	JUN 06, 2012			
	20401 001 DILTIAZEM HYDROCHLORIDE; TIAZAC	5529791	JUN 25, 2013			
	20401 002 DILTIAZEM HYDROCHLORIDE; TIAZAC	5529791	JUN 25, 2013			
	20401 003 DILTIAZEM HYDROCHLORIDE; TIAZAC	5529791	JUN 25, 2013			
	20401 004 DILTIAZEM HYDROCHLORIDE; TIAZAC	5529791	JUN 25, 2013			
	20401 005 DILTIAZEM HYDROCHLORIDE; TIAZAC	5529791	JUN 25, 2013		NS	SEP 11, 1998
	20506 001 DILTIAZEM MALATE; TIAMATE	4968507	JUL 25, 2006		NE	OCT 04, 1999
		4880631	SEP 24, 2007			
	20506 002 DILTIAZEM MALATE; TIAMATE	4968507	JUL 25, 2006		NE	OCT 04, 1999
		4880631	SEP 24, 2007			

PRESCRIPTION AND OTC DRUG PRODUCT
PATENT AND EXCLUSIVITY DATA

APPL/PROD NUMBER	INGREDIENT NAME; TRADE NAME	PATENT NUMBER	PATENT EXPIRES	USE CODE	EXCLUS CODE	EXCLUS EXPIRES	
20506 003	DILTIAZEM MALATE; TIAMATE	4968507	JUL 25, 2006		NE	OCT 04, 1999	
		4880631	SEP 24, 2007				
20507 001	DILTIAZEM MALATE; TECZEM	4983598	JAN 08, 2008				
		4968507	JUL 25, 2006				
		4880631	SEP 24, 2007				
		4472380	SEP 18, 2001		NC	OCT 04, 1999	
		4374829	FEB 22, 2000	U-3	NE	OCT 04, 1999	
18723 001	DIVALPROEX SODIUM; DEPAKOTE	5212326	JAN 29, 2008		I-41	MAR 18, 1999	
18723 002	DIVALPROEX SODIUM; DEPAKOTE	5212326	JAN 29, 2008		I-41	MAR 18, 1999	
18723 003	DIVALPROEX SODIUM; DEPAKOTE	5212326	JAN 29, 2008		I-41	MAR 18, 1999	
20449 001	DOCETAXEL; TAXOTERE	5403858	JUL 03, 2012				
		4814470	JUL 14, 2007		NCE	MAY 14, 2001	
20690 001	DONEPEZIL HYDROCHLORIDE; ARICEPT	4895841	JUN 20, 2008		NCE	NOV 25, 2001	
20690 002	DONEPEZIL HYDROCHLORIDE; ARICEPT	4895841	JUN 20, 2008		NCE	NOV 25, 2001	
>ADD>	20668 001	ENALPRIL MALEATE; LEXXEL	4264611	JUN 19, 2001	U-3	NC	DEC 27, 1999
>ADD>			4803081	APR 03, 2007			
	20164 001	ENOXAPARIN SODIUM; LOVENOX	5389618	FEB 14, 2012			
		4692435	DEC 24, 2004	U-123			
		4486420	DEC 04, 2001	U-122			
>ADD>	20655 002	ESTRADIOL; ALORA			NS	OCT 28, 1997	
	20472 001	ESTRADIOL; ESTRING			NDF	APR 26, 1999	
	20538 001	ESTRADIOL; ESTRADIOL	5300291	APR 05, 2011	NS	OCT 28, 1997	
		4994278	MAR 04, 2008				
		4994267	MAR 04, 2008				
		4814168	MAR 04, 2008				
	20538 002	ESTRADIOL; ESTRADIOL	5300291	APR 05, 2011			
		4994278	MAR 04, 2008				
		4994267	MAR 04, 2008				
		4814168	MAR 04, 2008				
	20538 003	ESTRADIOL; ESTRADIOL	5300291	APR 05, 2011	NS	OCT 28, 1997	
		4994278	MAR 04, 2008				
		4994267	MAR 04, 2008				
		4814168	MAR 04, 2008				
	20538 004	ESTRADIOL; ESTRADIOL	5300291	APR 05, 2011			
		4994278	MAR 04, 2008				
		4994267	MAR 04, 2008				
		4814168	MAR 04, 2008				
>ADD>	20130 001	ETHINYL ESTRADIOL; ESTROSTEP 21	4962098	OCT 09, 2007	U-112	NP	OCT 09, 1999
>ADD>	20130 002	ETHINYL ESTRADIOL; ESTROSTEP FE	4962098	OCT 09, 2007	U-112	NP	OCT 09, 1999
	18922 002	ETODOLAC; LODINE			I-24	JUN 28, 1999	
	18922 003	ETODOLAC; LODINE			I-24	JUN 28, 1999	
	18922 004	ETODOLAC; LODINE			I-24	JUN 28, 1999	

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PATENT AND EXCLUSIVITY DATA

APPL/PROD NUMBER	INGREDIENT NAME; TRADE NAME	PATENT NUMBER	PATENT EXPIRES	USE CODE	EXCLUS CODE	EXCLUS EXPIRES
20584 001	ETODOLAC; LODINE XL	4966768	OCT 30, 2007		NDF	OCT 25, 1999
20584 002	ETODOLAC; LODINE XL	4966768	OCT 30, 2007		NDF	OCT 25, 1999
20457 001	ETOPOSIDE PHOSPHATE; ETOPOPHOS	5041424	AUG 20, 2008	U-135		
		4904768	FEB 27, 2007		NE	MAY 17, 1999
20195 001	FENTANYL CITRATE; FENTANYL	4671953	MAY 01, 2005	U-87	NDF	OCT 04, 1996
20195 002	FENTANYL CITRATE; FENTANYL	4671953	MAY 01, 2005	U-87	NDF	OCT 04, 1996
20195 003	FENTANYL CITRATE; FENTANYL	4671953	MAY 01, 2005	U-87	NDF	OCT 04, 1996
20195 007	FENTANYL CITRATE; FENTANYL	4671953	MAY 01, 2005	U-87	NDF	OCT 04, 1996
20416 001	FERUMOXIDES; FERIDEX I.V.	5248492	SEP 28, 2010		NCE	AUG 30, 2001
		5219554	JUN 15, 2010			
		5102652	FEB 06, 2009			
		5055288	OCT 08, 2008			
		4951675	AUG 28, 2007	U-143		
		4827945	MAY 09, 2006	U-144		
		4770183	SEP 13, 2005	U-145		
>ADD> 20410 001	FERUMOXIL; GASTROMARK				NCE	DEC 06, 2001
20625 001	FEXOFENADINE HYDROCHLORIDE; ALLEGRA	5375693	AUG 03, 2012	U-138	NCE	JUL 25, 2001
		4254129	APR 10, 1999	U-139		
		4760071	JUN 19, 2006			
20180 001	FINASTERIDE; PROSCAR	4626549	DEC 02, 2003	U-154	I-166	NOV 21, 1999
18936 001	FLUOXETINE HYDROCHLORIDE; PROZAC	4626549	DEC 02, 2003	U-154	I-166	NOV 21, 1999
18936 006	FLUOXETINE HYDROCHLORIDE; PROZAC	4626549	DEC 02, 2003	U-154	I-166	NOV 21, 1999
18554 001	FLUTAMIDE; EULEXIN	4329364	MAY 11, 2001	U-23	I-168	JUN 21, 1999
20548 001	FLUTICASONE PROPIONATE; FLOVENT	4335121	NOV 14, 2003		NP	MAR 27, 1999
20548 002	FLUTICASONE PROPIONATE; FLOVENT	4335121	NOV 14, 2003		NP	MAR 27, 1999
20548 003	FLUTICASONE PROPIONATE; FLOVENT	4335121	NOV 14, 2003		NP	MAR 27, 1999
20261 001	FLUVASTATIN SODIUM; LESCOL				D-31	MAR 20, 1999
20261 002	FLUVASTATIN SODIUM; LESCOL				D-31	MAR 20, 1999
20450 001	FOSPHENYTOIN SODIUM; CEREBYX	4925860	AUG 05, 2007		ODE	AUG 05, 2003
		4260769	APR 07, 1998		NCE	AUG 05, 2001
20235 001	GABAPENTIN; NEURONTIN	5084479	JAN 02, 2010	U-125		
		4894476	MAY 02, 2008			
		4087544	JAN 17, 2001	U-86	NCE	DEC 30, 1998
20235 002	GABAPENTIN; NEURONTIN	5084479	JAN 02, 2010	U-125		
		4894476	MAY 02, 2008			
		4087544	JAN 17, 2001	U-86	NCE	DEC 30, 1998
20235 003	GABAPENTIN; NEURONTIN	5084479	JAN 02, 2010	U-125		
		4894476	MAY 02, 2008			
		4087544	JAN 17, 2001	U-86	NCE	DEC 30, 1998
20123 001	GADODIAMIDE; OMNISCAN	4687659	MAY 04, 2007		D-29	FEB 05, 1999
					I-145	FEB 05, 1999
					I-144	FEB 05, 1999
19596 001	GADOPENTETATE DIMEGLUMINE; MAGNEVIST	4647447	MAR 03, 2004		I-146	FEB 28, 1999

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PATENT AND EXCLUSIVITY DATA

APPL/PROD NUMBER	INGREDIENT NAME; TRADE NAME	PATENT NUMBER	PATENT EXPIRES	USE CODE	EXCLUS CODE	EXCLUS EXPIRES
20569 001	GANCICLOVIR; VITRASERT	5378475	JAN 03, 2012		NP ODE	MAR 04, 1999 MAR 04, 2003
20509 001	GEMCITABINE HYDROCHLORIDE; GEMZAR	5464826	NOV 07, 2012	U-146		
		4808614	FEB 28, 2006	U-146	NCE	MAY 15, 2001
20509 002	GEMCITABINE HYDROCHLORIDE; GEMZAR	5464826	NOV 07, 2012	U-146		
		4808614	FEB 28, 2006	U-146	NCE	MAY 15, 2001 DEC 20, 2001
>ADD> 20622 001	GLATIRAMER ACETATE; COPAXONE					
20329 001	GLIPIZIDE; GLUCOTROL XL	5545413	JUL 02, 2008	U-150		
		5082668	SEP 16, 2003			
20329 002	GLIPIZIDE; GLUCOTROL XL	5545413	JUL 02, 2008	U-150		
		5082668	SEP 16, 2003			
19726 001	GOSERELIN ACETATE; ZOLADEX	5366734	NOV 22, 2011			
		4767628	AUG 30, 2005		I-88	FEB 02, 1996
20578 001	GOSERELIN ACETATE; ZOLADEX	5366734	NOV 22, 2011			
		4767628	AUG 30, 2005			
20239 001	GRANISETRON HYDROCHLORIDE; KYTRIL	4886808	DEC 29, 2007	U-89		
20305 001	GRANISETRON HYDROCHLORIDE; KYTRIL	4886808	DEC 29, 2007	U-105		
19836 001	HISTRELIN ACETATE; SUPPRELIN	4244946	JAN 13, 2000		NCE	DEC 24, 1996
19836 002	HISTRELIN ACETATE; SUPPRELIN	4244946	JAN 13, 2000		NCE	DEC 24, 1996
19836 003	HISTRELIN ACETATE; SUPPRELIN	4244946	JAN 13, 2000		NCE	DEC 24, 1996
20589 001	IBUPROFEN; CHILDREN'S ADVIL	4788220	JUL 08, 2007		NP	JUN 16, 1998
20601 001	IBUPROFEN; CHILDREN'S MOTRIN	5215755	JUN 01, 2010		NP	JUN 16, 1998
20601 002	IBUPROFEN; JUNIOR STRENGTH MOTRIN	5215755	JUN 01, 2010		NP	JUN 16, 1998
20602 001	IBUPROFEN; JUNIOR STRENGTH MOTRIN				NP	JUN 16, 1998
20603 001	IBUPROFEN; CHILDREN'S MOTRIN	5374659	DEC 20, 2011		NP	JUN 16, 1998
20685 001	INDINAVIR SULFATE; CRIXIVAN	5413999	MAY 07, 2013	U-132	NCE	MAR 13, 2001
20685 003	INDINAVIR SULFATE; CRIXIVAN	5413999	MAY 07, 2013	U-132	NCE	MAR 13, 2001
20563 001	INSULIN LISPRO; HUMALOG	5514646	MAY 07, 2013	U-111	NCE	JUN 14, 2001
		5474978	JUN 16, 2014			
20351 001	IODIXANOL; VISIPAQUE 270	5349085	SEP 20, 2011		NCE	MAR 22, 2001
		4396597	JUL 03, 1999			
		4278654	JUL 03, 1999			
20351 002	IODIXANOL; VISIPAQUE 320	5349085	SEP 20, 2011		NCE	MAR 22, 2001
		4396597	JUL 03, 1999			
		4278654	JUL 03, 1999			
18735 001	IOPAMIDOL; ISOVUE-M 200	4001323	JAN 04, 1996			
18735 002	IOPAMIDOL; ISOVUE-300	4001323	JAN 04, 1996			
18735 003	IOPAMIDOL; ISOVUE-370	4001323	JAN 04, 1996		D-28	MAY 15, 1998
18735 004	IOPAMIDOL; ISOVUE-M 300	4001323	JAN 04, 1996			
18735 007	IOPAMIDOL; ISOVUE-250	4001323	JAN 04, 1996			
20327 001	IOPAMIDOL; ISOVUE-200	4001323	JAN 04, 1996			
20327 002	IOPAMIDOL; ISOVUE-250	4001323	JAN 04, 1996			

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PATENT AND EXCLUSIVITY DATA

APPL/PROD NUMBER	INGREDIENT NAME; TRADE NAME	PATENT NUMBER	PATENT EXPIRES	USE CODE	EXCLUS CODE	EXCLUS EXPIRES
20327 003	IOPAMIDOL; ISOVUE-300	4001323	JAN 04, 1996			
20327 004	IOPAMIDOL; ISOVUE-370	4001323	JAN 04, 1996			
20220 001	IOPROMIDE; ULTRAVIST 370	4364921	MAR 06, 2005	U-113	NCE	MAY 10, 2000
20220 002	IOPROMIDE; ULTRAVIST 300	4364921	MAR 06, 2005	U-113	NCE	MAY 10, 2000
20220 003	IOPROMIDE; ULTRAVIST 240	4364921	MAR 06, 2005	U-113	NCE	MAY 10, 2000
20220 004	IOPROMIDE; ULTRAVIST 150	4364921	MAR 06, 2005	U-113	NCE	MAY 10, 2000
20571 001	IRINOTECAN HYDROCHLORIDE; CAMPTOSAR	4604463	JUL 05, 2004		NCE	JUN 14, 2001
20083 001	ITRACONAZOLE; SPORANOX				I-155	SEP 28, 1998
19645 001	KETOROLAC TROMETHAMINE; TORADOL	4089969	MAY 16, 1997	U-55		
19698 001	KETOROLAC TROMETHAMINE; TORADOL	4089969	MAY 16, 1997	U-55	NR	DEC 07, 1997
19698 002	KETOROLAC TROMETHAMINE; TORADOL	4089969	MAY 16, 1997	U-55	NR	DEC 07, 1997
>ADD>	19700 001	4089969	MAY 16, 1997	U-75	I-176	DEC 31, 1999
	20564 001	5047407	FEB 08, 2009			
	20596 001	5047407	FEB 08, 2009			
	20241 001	4602017	JUL 22, 2008	U-106	NCE	DEC 27, 1999
	20241 002	4602017	JUL 22, 2008	U-106	NCE	DEC 27, 1999
	20241 003	4602017	JUL 22, 2008	U-106	NCE	DEC 27, 1999
	20241 004	4602017	JUL 22, 2008	U-106	NCE	DEC 27, 1999
	20241 005	4602017	JUL 22, 2008	U-106	NCE	DEC 27, 1999
	20241 006	4602017	JUL 22, 2008	U-106	NCE	DEC 27, 1999
>ADD>	20406 001	5433959	SEP 03, 2008			
		4689333	JUL 29, 2005	U-126	I-116	APR 08, 1999
>ADD>		4628098	MAY 10, 2009		NCE	MAY 10, 2000
>DEL>		4628098	JUL 29, 2005		NCE	MAY 10, 2000
>ADD>	20406 002	5433959	SEP 03, 2008			
		4689333	JUL 29, 2005	U-126	I-116	APR 08, 1999
>ADD>		4628098	MAY 10, 2009		NCE	MAY 10, 2000
>DEL>		4628098	JUL 29, 2005		NCE	MAY 10, 2000
	20597 001				NCE	JUN 05, 2001
>ADD>	19732 001	5575987	NOV 19, 2013			
>ADD>	20011 001	5575987	NOV 19, 2013			
>ADD>	19943 001	5575987	NOV 19, 2013			
>ADD>	20263 001	5575987	NOV 19, 2013			
>ADD>	20263 002	5575987	NOV 19, 2013			
>ADD>	20263 003	5575987	NOV 19, 2013			
>ADD>	20263 004	5575987	NOV 19, 2013			
>ADD>	20263 005	5575987	NOV 19, 2013			
>ADD>	20263 006	5575987	NOV 19, 2013			
>ADD>	20517 001	5575987	NOV 19, 2013			
		5480656	JAN 02, 2013			
	20219 001	4369184	DEC 02, 2004		NCE	NOV 10, 1998

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PATENT AND EXCLUSIVITY DATA

APPL/PROD NUMBER	INGREDIENT NAME; TRADE NAME	PATENT NUMBER	PATENT EXPIRES	USE CODE	EXCLUS CODE	EXCLUS EXPIRES
>ADD>	20634 001 LEVOFLOXACIN; LEVAQUIN	4382892	SEP 02, 2001			
>ADD>		5053407	OCT 01, 2008			
>ADD>	20634 002 LEVOFLOXACIN; LEVAQUIN	4382892	SEP 02, 2001			
>ADD>		5053407	OCT 01, 2008			
>ADD>	20635 001 LEVOFLOXACIN; LEVAQUIN	4382892	SEP 02, 2001			
>ADD>		5053407	OCT 01, 2008			
>ADD>	20635 002 LEVOFLOXACIN; LEVAQUIN	4382892	SEP 02, 2001			
>ADD>		5053407	OCT 01, 2008			
>ADD>	20635 003 LEVOFLOXACIN; LEVAQUIN	4382892	SEP 02, 2001			
>ADD>		5053407	OCT 01, 2008			
>ADD>	20544 001 LEVONORGESTREL; NORPLANT II				NP	NOV 01, 1999
	20627 001 LEVONORGESTREL; LEVONORGESTREL				NP	AUG 15, 1999
	20575 001 LIDOCAINE; LIDOCAINE	5446070	FEB 27, 2011			
		5332576	JUL 26, 2011			
		5234957	FEB 27, 2011		NDF	MAY 21, 1999
	20575 002 LIDOCAINE; LIDOCAINE	5446070	FEB 27, 2011			
		5332576	JUL 26, 2011			
		5234957	FEB 27, 2011		NDF	MAY 21, 1999
	19558 001 LISINOPRIL; PRINIVIL				I-141	NOV 24, 1998
	19558 002 LISINOPRIL; PRINIVIL				I-141	NOV 24, 1998
	19558 003 LISINOPRIL; PRINIVIL				I-141	NOV 24, 1998
	19558 004 LISINOPRIL; PRINIVIL				I-141	NOV 24, 1998
	19558 006 LISINOPRIL; PRINIVIL				I-141	NOV 24, 1998
	19777 001 LISINOPRIL; ZESTRIL				I-141	NOV 24, 1998
	19777 002 LISINOPRIL; ZESTRIL				I-141	NOV 24, 1998
	19777 003 LISINOPRIL; ZESTRIL				I-141	NOV 24, 1998
	19777 004 LISINOPRIL; ZESTRIL				I-141	NOV 24, 1998
	19777 005 LISINOPRIL; ZESTRIL				I-141	NOV 24, 1998
>ADD>	20013 001 LOMEFLOXACIN HYDROCHLORIDE; MAXAQUIN				I-173	OCT 09, 1999
	19658 001 LORATADINE; CLARITIN	4659716	APR 21, 2004	U-142	I-136	SEP 20, 1998
	19670 001 LORATADINE; CLARITIN-D	4659716	APR 21, 2004	U-142	NC	NOV 14, 1997
	20470 001 LORATADINE; CLARITIN-D 24 HOUR	5314697	OCT 23, 2012			
		4659716	APR 21, 2004	U-142	NP	AUG 23, 1999
		4282233	JUL 19, 2002	U-77	NCE	APR 12, 1998
		4282233	JUN 19, 2002	U-77	NDF	OCT 10, 1999
>ADD>	20641 001 LORATADINE; CLARITIN	4282233	JUN 19, 2002	U-77		
>ADD>	20704 001 LORATADINE; CLARITIN REDITABS	4371516	FEB 01, 2000			
	19940 001 MASOPROCOL; ACTINEX	4695590	APR 17, 2008			
	19651 001 MESALAMINE; ASACOL	5541171	JUL 30, 2013	U-141		
		5541170	JUL 30, 2013	U-141		
	20208 001 METRONIDAZOLE; METROGEL-VAGINAL	5536743	JUL 16, 2013	U-137		
	20670 002 MICONAZOLE NITRATE; MONISTAT-3 COMBINATION PACK				NP	APR 16, 1999

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PATENT AND EXCLUSIVITY DATA

APPL/PROD NUMBER	INGREDIENT NAME; TRADE NAME	PATENT NUMBER	PATENT EXPIRES	USE CODE	EXCLUS CODE	EXCLUS EXPIRES
19815 001	MIDODRINE HYDROCHLORIDE; PROAMATINE				NCE	SEP 06, 2001
19815 002	MIDODRINE HYDROCHLORIDE; PROAMATINE				NCE	SEP 06, 2001
>ADD>	20682 001 MIGLITOL; GLYSET	4639436	JAN 27, 2004	U-111	NCE	DEC 18, 2001
>ADD>	20682 002 MIGLITOL; GLYSET	4639436	JAN 27, 2004	U-111	NCE	DEC 18, 2001
>ADD>	20682 003 MIGLITOL; GLYSET	4639436	JAN 27, 2004	U-111	NCE	DEC 18, 2001
	20415 001 MIRTAZAPINE; REMERON				NCE	JUN 14, 2001
	20415 002 MIRTAZAPINE; REMERON				NCE	JUN 14, 2001
	19297 001 MITOXANTRONE HYDROCHLORIDE; NOVANTRONE	4197249	APR 08, 1997		I-169	NOV 13, 1999
>ADD>		4138415	FEB 06, 1996		ODE	NOV 13, 2003
	20312 001 MOEXIPRIL HYDROCHLORIDE; UNIVASC	4344949	OCT 03, 2000			
	20312 002 MOEXIPRIL HYDROCHLORIDE; UNIVASC	4344949	OCT 03, 2000			
	20616 001 MORPHINE SULFATE; KADIAN	5378474	MAR 23, 2010			
		5202128	APR 13, 2010		NDF	JUL 03, 1999
	20616 002 MORPHINE SULFATE; KADIAN	5378474	MAR 23, 2010			
		5202128	APR 13, 2010		NDF	JUL 03, 1999
	20616 003 MORPHINE SULFATE; KADIAN	5378474	MAR 23, 2010			
		5202128	APR 13, 2010		NDF	JUL 03, 1999
	19886 001 NAFARELIN ACETATE; SYNAREL	4234571	JUN 11, 2011			
	20353 001 NAPROXEN SODIUM; NAPRELAN				NDF	JAN 05, 1999
	20353 002 NAPROXEN SODIUM; NAPRELAN				NDF	JAN 05, 1999
	20353 003 NAPROXEN SODIUM; NAPRELAN				NDF	JAN 05, 1999
	20152 001 NEFAZODONE HYDROCHLORIDE; SERZONE	4338317	MAR 16, 2003	U-12	NCE	DEC 22, 1999
	20152 002 NEFAZODONE HYDROCHLORIDE; SERZONE	4338317	MAR 16, 2003	U-12	NCE	DEC 22, 1999
	20152 003 NEFAZODONE HYDROCHLORIDE; SERZONE	4338317	MAR 16, 2003	U-12	NCE	DEC 22, 1999
	20152 004 NEFAZODONE HYDROCHLORIDE; SERZONE	4338317	MAR 16, 2003	U-12	NCE	DEC 22, 1999
	20152 005 NEFAZODONE HYDROCHLORIDE; SERZONE	4338317	MAR 16, 2003	U-12	NCE	DEC 22, 1999
	20152 006 NEFAZODONE HYDROCHLORIDE; SERZONE	4338317	MAR 16, 2003	U-12	NCE	DEC 22, 1999
	20636 001 NEVIRAPINE; VIRAMUNE	5366972	NOV 22, 2011		NCE	JUN 21, 2001
	19488 001 NICARDIPINE HYDROCHLORIDE; CARDENE	3985758	OCT 12, 1995			
	19488 002 NICARDIPINE HYDROCHLORIDE; CARDENE	3985758	OCT 12, 1995			
	19734 001 NICARDIPINE HYDROCHLORIDE; CARDENE	3985758	OCT 12, 1995			
	20005 001 NICARDIPINE HYDROCHLORIDE; CARDENE SR	3985758	OCT 12, 1995			
	20005 002 NICARDIPINE HYDROCHLORIDE; CARDENE SR	3985758	OCT 12, 1995			
	20005 003 NICARDIPINE HYDROCHLORIDE; CARDENE SR	3985758	OCT 12, 1995			
>DLT>	20165 001 NICOTINE; NICODERM CQ	5508038	APR 16, 2013			
>DLT>	20165 002 NICOTINE; NICODERM CQ	5508038	APR 16, 2013			
>DLT>	20165 003 NICOTINE; NICODERM CQ	5508038	APR 16, 2013			
>ADD>	20165 004 NICOTINE; NICODERM CQ	5508038	APR 16, 2013		NP	AUG 02, 1999
>ADD>	20165 005 NICOTINE; NICODERM CQ	5508038	APR 16, 2013		NP	AUG 02, 1999
>ADD>	20165 006 NICOTINE; NICODERM CQ	5508038	APR 16, 2013		NP	AUG 02, 1999
	20385 001 NICOTINE; NICOTROL				NP	MAR 22, 1999
>ADD>	20536 001 NICOTINE; NICOTROL				NP	JUL 03, 1999

PRESCRIPTION AND OTC DRUG PRODUCT
PATENT AND EXCLUSIVITY DATA

APPL/PROD NUMBER	INGREDIENT NAME; TRADE NAME	PATENT NUMBER	PATENT EXPIRES	USE CODE	EXCLUS CODE	EXCLUS EXPIRES
18612 002	NICOTINE POLACRILEX; NICORETTE				NP	FEB 09, 1999
20066 002	NICOTINE POLACRILEX; NICORETTE				NP	FEB 09, 1999
20169 001	NILUTAMIDE; NILANDRON				NCE	SEP 19, 2001
>ADD>	20144 001	4954344	DEC 04, 2001	U-158		
>ADD>		4849226	DEC 04, 2001	U-158		
>ADD>		4812313	DEC 04, 2001	U-158		
>ADD>	20144 002	4954344	DEC 04, 2001	U-158		
>ADD>		4849226	DEC 04, 2001	U-158		
>ADD>		4812313	DEC 04, 2001	U-158		
>ADD>	20144 003	4954344	DEC 04, 2001	U-158		
>ADD>		4849226	DEC 04, 2001	U-158		
>ADD>		4812313	DEC 04, 2001	U-158		
>ADD>	20144 004	4954344	DEC 04, 2001	U-158		
>ADD>		4849226	DEC 04, 2001	U-158		
>ADD>		4812313	DEC 04, 2001	U-158		
>ADD>	20144 005	4954344	DEC 04, 2001	U-158		
>ADD>		4849226	DEC 04, 2001	U-158		
>ADD>		4812313	DEC 04, 2001	U-158		
	20555 001	4375547	OCT 02, 2002		NS	MAY 09, 1999
>ADD>	19735 001	4382892	SEP 02, 2003		I-174	DEC 19, 1999
>ADD>	19735 002	4382892	SEP 02, 2003		I-174	DEC 19, 1999
>ADD>	19735 003	4382892	SEP 02, 2003		I-174	DEC 19, 1999
	19921 001				ODE	MAY 22, 2003
					I-160	MAY 22, 1999
>ADD>	20087 001	4382892	SEP 02, 2001		I-174	DEC 19, 1999
>ADD>	20087 002	4382892	SEP 02, 2001		I-174	DEC 19, 1999
>ADD>	20087 003	4382892	SEP 02, 2001		I-174	DEC 19, 1999
>ADD>	20087 004	4382892	SEP 02, 2001		I-174	DEC 19, 1999
>ADD>	20087 005	4382892	SEP 02, 2001		I-174	DEC 19, 1999
	20592 001	5229382	APR 23, 2011	U-149	NCE	SEP 30, 2001
	20592 002	5229382	APR 23, 2011	U-149	NCE	SEP 30, 2001
	20592 003	5229382	APR 23, 2011	U-149	NCE	SEP 30, 2001
	20592 004	5229382	APR 23, 2011	U-149	NCE	SEP 30, 2001
>ADD>	20688 001				NCE	DEC 18, 2001
	19810 001	4255431	APR 05, 2001	U-108	I-23	MAR 22, 1999
	19810 003	4853230	APR 20, 2007	U-108	I-23	MAR 22, 1999
>ADD>	20007 001	5578628	JUN 24, 2006	U-44	I-151	MAY 16, 1999
>ADD>	20007 003	5578628	JUN 24, 2006	U-44		
>ADD>	20103 001	5578628	JUN 24, 2006	U-44	I-110	SEP 26, 1997
>ADD>	20103 002	5578628	JUN 24, 2006	U-44	I-110	SEP 26, 1997
>ADD>	20403 001	5578628	JUN 24, 2006	U-44	I-9	AUG 13, 1996

PRESCRIPTION AND OTC DRUG PRODUCT
PATENT AND EXCLUSIVITY DATA

APPL/PROD NUMBER	INGREDIENT NAME; TRADE NAME	PATENT NUMBER	PATENT EXPIRES	USE CODE	EXCLUS CODE	EXCLUS EXPIRES	
18841 004	OXAPROZIN; DAYPRO				NCE	OCT 29, 1999*	
20553 001	OXYCODONE HYDROCHLORIDE; OXYCONTIN	5549912	FEB 05, 2008				
		5266331	FEB 05, 2008		NDF	DEC 12, 1998	
20553 002	OXYCODONE HYDROCHLORIDE; OXYCONTIN	5549912	FEB 05, 2008				
		5266331	FEB 05, 2008		NDF	DEC 12, 1998	
20553 003	OXYCODONE HYDROCHLORIDE; OXYCONTIN	5549912	FEB 05, 2008				
		5266331	FEB 05, 2008		NDF	DEC 12, 1998	
20036 001	PAMIDRONATE DISODIUM; AREDIA	3962432	JUL 16, 1996		I-158	JUL 16, 1999	
20036 003	PAMIDRONATE DISODIUM; AREDIA	3962432	JUL 16, 1996		I-158	JUL 16, 1999	
20036 004	PAMIDRONATE DISODIUM; AREDIA	3962432	JUL 16, 1996		I-158	JUL 16, 1999	
20031 001	PAROXETINE HYDROCHLORIDE; PAXIL				I-150	MAR 07, 1999	
20031 002	PAROXETINE HYDROCHLORIDE; PAXIL				I-150	MAR 07, 1999	
20031 003	PAROXETINE HYDROCHLORIDE; PAXIL				I-150	MAR 07, 1999	
20031 004	PAROXETINE HYDROCHLORIDE; PAXIL				I-150	MAR 07, 1999	
20031 005	PAROXETINE HYDROCHLORIDE; PAXIL				I-150	MAR 07, 1999	
20629 001	PENCICLOVIR; DENAVIR	5075445	DEC 24, 2008		NCE	SEP 24, 2001	
19887 002	PENTAMIDINE ISETHIONATE; NEBUPENT				I-148	MAR 05, 1999	
>ADD>	20193 001	PENTOSAN POLYSULFATE SODIUM; ELMIRON	5180715	JAN 19, 2010	U-159	NCE	SEP 26, 2001
	20184 001	PERINDOPRIL ERBUMINE; ACEON	4508729	AUG 21, 2006			
	20184 002	PERINDOPRIL ERBUMINE; ACEON	4508729	AUG 21, 2006			
	20184 003	PERINDOPRIL ERBUMINE; ACEON	4508729	AUG 21, 2006			
	19918 001	PERMETHRIN; NIX	4024163	MAY 17, 1996		I-170	NOV 01, 1999
	20451 001	PORFIMER SODIUM; PHOTOFRIN	5438071	AUG 01, 2012			
		5145863	JUN 12, 2007	U-129	ODE	DEC 27, 2002	
		5028621	MAR 10, 2004				
		4932934	JUN 12, 2007	U-128	NCE	DEC 27, 2000	
		4866168	MAR 10, 2004				
		4649151	MAR 10, 2004				
>ADD>	19898 002	PRAVASTATIN SODIUM; PRAVACHOL	5180589	JUL 09, 2008		I-152	MAR 22, 1999
>ADD>			5030447	JUL 09, 2008		I-159	JUL 02, 1999
>ADD>			4346227	OCT 20, 2005			
>ADD>	19898 003	PRAVASTATIN SODIUM; PRAVACHOL	5180589	JUL 09, 2008		I-152	MAR 22, 1999
>ADD>			5030447	JUL 09, 2008		I-159	JUL 02, 1999
>ADD>			4346227	OCT 20, 2005			
>ADD>	19898 004	PRAVASTATIN SODIUM; PRAVACHOL	5180589	JUL 09, 2008		I-152	MAR 22, 1999
>ADD>			5030447	JUL 09, 2008		I-159	JUL 02, 1999
>ADD>			4346227	OCT 20, 2005			

* - In accordance with section 2105(c) of the FDA Export Reform and Enhancement Act of 1996 (Public Law 104-134)

PRESCRIPTION AND OTC DRUG PRODUCT
PATENT AND EXCLUSIVITY DATA

APPL/PROD NUMBER	INGREDIENT NAME; TRADE NAME	PATENT NUMBER	PATENT EXPIRES	USE CODE	EXCLUS CODE	EXCLUS EXPIRES
>DLT>	19898 005 PRAVASTATIN SODIUM; PRAVACHOL	5180589	JUL 09, 2008		I-152	MAR 22, 1999
>DLT>		5030447	JUL 09, 2008		I-159	JUL 02, 1999
>DLT>		4346227	OCT 20, 2005		NCE	OCT 31, 1996
>DLT>	19898 006 PRAVASTATIN SODIUM; PRAVACHOL	5180589	JUL 09, 2008		I-152	MAR 22, 1999
>DLT>		5030447	JUL 09, 2008		I-159	JUL 02, 1999
>DLT>		4346227	OCT 20, 2005		NCE	OCT 31, 1996
>DLT>	19898 007 PRAVASTATIN SODIUM; PRAVACHOL	5180589	JUL 09, 2008		I-152	MAR 22, 1999
>DLT>		5030447	JUL 09, 2008		I-159	JUL 02, 1999
>DLT>		4346227	OCT 20, 2005		NCE	OCT 31, 1996
	20279 001 PREDNICARBATE; DERMATOP				I-154	MAY 03, 1999
	20545 001 PROCAINAMIDE HYDROCHLORIDE; PROCANBID				NP	JAN 31, 1999
	20545 002 PROCAINAMIDE HYDROCHLORIDE; PROCANBID				NP	JAN 31, 1999
	19627 001 PROPOFOL; DIPRIVAN	4056635	NOV 01, 1996		I-90	MAR 08, 1996
	19885 001 QUINAPRIL HYDROCHLORIDE; ACCUPRIL	4344949	OCT 03, 2002			
	19885 002 QUINAPRIL HYDROCHLORIDE; ACCUPRIL	4344949	OCT 03, 2002			
	19885 003 QUINAPRIL HYDROCHLORIDE; ACCUPRIL	4344949	OCT 03, 2002			
	19885 004 QUINAPRIL HYDROCHLORIDE; ACCUPRIL	4344949	OCT 03, 2002			
	20559 001 RANITIDINE BISMUTH CITRATE; TRITEC	5008256	JUL 17, 2009		NE	AUG 08, 1999
	19593 001 RANITIDINE HYDROCHLORIDE; ZANTAC	4585790	MAY 11, 2004			
		4521431	JUN 04, 2002	U-121		
		4128658	JUL 25, 1997	U-121		
	19593 002 RANITIDINE HYDROCHLORIDE; ZANTAC	4585790	MAY 11, 2004			
		4521431	JUN 04, 2002	U-121		
		4128658	JUL 25, 1997	U-121		
	20520 001 RANITIDINE HYDROCHLORIDE; ZANTAC 75	4880636	MAY 13, 2008			
		4521431	JUN 04, 2002	U-121		
		4128658	JUL 25, 1997	U-121		
	20630 001 REMIFENTANIL HYDROCHLORIDE; ULTIVA	5466700	AUG 30, 2013	U-156		
		5019583	FEB 15, 2009		NCE	JUL 12, 2001
	20630 002 REMIFENTANIL HYDROCHLORIDE; ULTIVA	5466700	AUG 30, 2013	U-156		
		5019583	FEB 15, 2009		NCE	JUL 12, 2001
	20630 003 REMIFENTANIL HYDROCHLORIDE; ULTIVA	5466700	AUG 30, 2013	U-156		
		5019583	FEB 15, 2009		NCE	JUL 12, 2001
	20272 001 RISPERIDONE; RISPERDAL	4804663	DEC 29, 2007	U-90		
	20272 002 RISPERIDONE; RISPERDAL	4804663	DEC 29, 2007	U-90		
	20272 003 RISPERIDONE; RISPERDAL	4804663	DEC 29, 2007	U-90		
	20272 004 RISPERIDONE; RISPERDAL	4804663	DEC 29, 2007	U-90		
	20272 005 RISPERIDONE; RISPERDAL	4804663	DEC 29, 2007	U-90		
	20588 001 RISPERIDONE; RISPERDAL	4804663	DEC 29, 2007	U-90	NCE	DEC 29, 1998
	20659 001 RITONAVIR; NORVIR	5541206	JUL 30, 2013	U-140		
		5484801	JAN 28, 2014		NCE	MAR 01, 2001
	20680 001 RITONAVIR; NORVIR	5541206	JUL 30, 2013	U-140	NCE	MAR 01, 2001

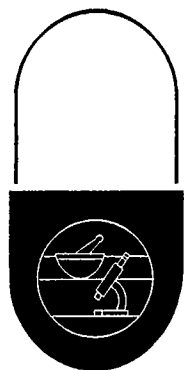
PRESCRIPTION AND OTC DRUG PRODUCT
PATENT AND EXCLUSIVITY DATA

APPL/PROD NUMBER	INGREDIENT NAME; TRADE NAME	PATENT NUMBER	PATENT EXPIRES	USE CODE	EXCLUS CODE	EXCLUS EXPIRES	
20533 001	ROPIVACAINE HYDROCHLORIDE MONOHYDRATE; NAROPIN	4870086	SEP 26, 2006		NCE	SEP 24, 2001	
		4695576	SEP 22, 2004				
20533 003	ROPIVACAINE HYDROCHLORIDE MONOHYDRATE; NAROPIN	4870086	SEP 26, 2006		NCE	SEP 24, 2001	
		4695576	SEP 22, 2004				
20533 004	ROPIVACAINE HYDROCHLORIDE MONOHYDRATE; NAROPIN	4870086	SEP 26, 2006		NCE	SEP 24, 2001	
		4695576	SEP 22, 2004				
20533 005	ROPIVACAINE HYDROCHLORIDE MONOHYDRATE; NAROPIN	4870086	SEP 26, 2006		NCE	SEP 24, 2001	
		4695576	SEP 22, 2004				
20628 001	SAQUINAVIR MESYLATE; INVIRASE	5196438	NOV 19, 2010				
19839 001	SERTRALINE HYDROCHLORIDE; ZOLOFT	4962128	NOV 02, 2009	U-152			
		4536518	DEC 31, 2005	U-152	I-102	OCT 25, 1999	
19839 002	SERTRALINE HYDROCHLORIDE; ZOLOFT	4962128	NOV 02, 2009	U-152			
		4536518	DEC 31, 2005	U-152	I-102	OCT 25, 1999	
19839 003	SERTRALINE HYDROCHLORIDE; ZOLOFT	4962128	NOV 02, 2009	U-152			
		4536518	DEC 31, 2005	U-152	I-102	OCT 25, 1999	
19839 004	SERTRALINE HYDROCHLORIDE; ZOLOFT	4962128	NOV 02, 2009	U-152			
		4536518	DEC 31, 2005	U-152	I-102	OCT 25, 1999	
19839 005	SERTRALINE HYDROCHLORIDE; ZOLOFT	4962128	NOV 02, 2009	U-152			
		4536518	DEC 31, 2005	U-152	I-102	OCT 25, 1999	
20572 001	SODIUM PHENYLBUTYRATE; BUPHENYL	4457942	AUG 20, 2002	U-136	NCE	APR 30, 2001	
20573 001	SODIUM PHENYLBUTYRATE; BUPHENYL	4457942	AUG 20, 2002	U-136	NCE	APR 30, 2001	
					ODE	APR 30, 2003	
>ADD>	19931 001	SODIUM SULFACETAMIDE; KLARON			NDF	DEC 23, 1999	
	19640 001	SOMATROPIN, BIOSYNTHETIC; HUMATROPE			I-161	AUG 01, 1999	
	19640 004	SOMATROPIN, BIOSYNTHETIC; HUMATROPE			I-161	AUG 01, 1999	
	20280 004	SOMATROPIN, BIOSYNTHETIC; GENOTROPIN			NS	AUG 24, 1998	
	20280 006	SOMATROPIN, BIOSYNTHETIC; GENOTROPIN			NS	AUG 24, 1998	
	20604 001	SOMATROPIN, BIOSYNTHETIC; SEROSTIM			ODE	AUG 23, 2003	
					NP	AUG 23, 1999	
	20604 002	SOMATROPIN, BIOSYNTHETIC; SEROSTIM			ODE	AUG 23, 2003	
					NP	AUG 23, 1999	
>ADD>	20677 001	SPARFLOXACIN; ZAGAM	4795751	OCT 28, 2006	U-160	NCE	DEC 19, 2001
	20240 001	SPIRAPRIL HYDROCHLORIDE; RENORMAX	4470972	SEP 11, 2003	U-3	NCE	DEC 29, 1999
	20240 002	SPIRAPRIL HYDROCHLORIDE; RENORMAX	4470972	SEP 11, 2003	U-3	NCE	DEC 29, 1999
	20240 003	SPIRAPRIL HYDROCHLORIDE; RENORMAX	4470972	SEP 11, 2003	U-3	NCE	DEC 29, 1999
	20240 004	SPIRAPRIL HYDROCHLORIDE; RENORMAX	4470972	SEP 11, 2003	U-3	NCE	DEC 29, 1999
	20412 001	STAVUDINE; ZERIT	4978655	JUN 25, 2008	U-94		
	20412 002	STAVUDINE; ZERIT	4978655	JUN 25, 2008	U-94		
	20412 003	STAVUDINE; ZERIT	4978655	JUN 25, 2008	U-94		
	20412 004	STAVUDINE; ZERIT	4978655	JUN 25, 2008	U-94		
	20412 005	STAVUDINE; ZERIT	4978655	JUN 25, 2008	U-94		
	07073 002	SULFASALAZINE; AZULFIDINE EN-TABS				I-165	OCT 17, 1999

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PATENT AND EXCLUSIVITY DATA

APPL/PROD NUMBER	INGREDIENT NAME; TRADE NAME	PATENT NUMBER	PATENT EXPIRES	USE CODE	EXCLUS CODE	EXCLUS EXPIRES
20547 001	ZAFIRLUKAST; ACCOLATE	5482963	JAN 09, 2013			
		5319097	DEC 11, 2011			
		5294636	DEC 11, 2011			
		4859692	AUG 22, 2006			
>ADD>	20471 001	ZILEUTON; ZYFLO			NCE	SEP 26, 2001
>ADD>	20471 003	ZILEUTON; ZYFLO			NCE	DEC 09, 2001
					NCE	DEC 09, 2001

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