

APPROVED DRUG PRODUCTS

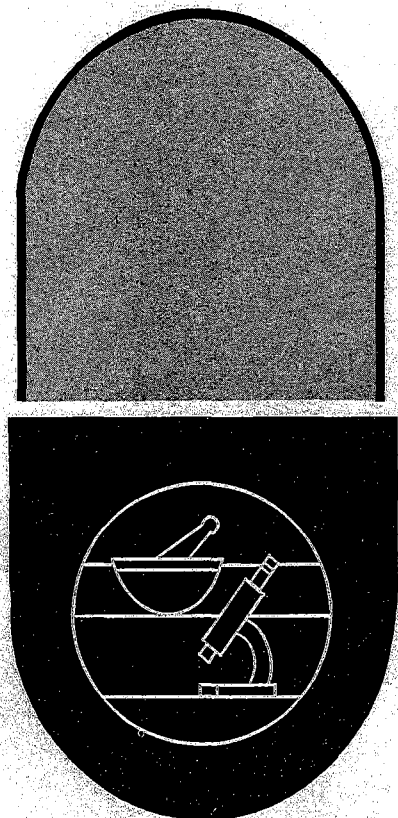
With Therapeutic Equivalence Evaluations



The "Orange Book"

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**CUMULATIVE
SUPPLEMENT 12
JAN'95-DEC'95**



APPROVED DRUG PRODUCTS

**WITH
THERAPEUTIC EQUIVALENCE EVALUATIONS**

15TH EDITION

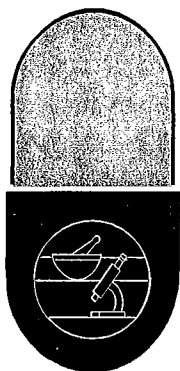
U.S. DEPARTMENT OF HEALTH AND HUMAN SERVICES

**PUBLIC HEALTH SERVICE
FOOD AND DRUG ADMINISTRATION
CENTER FOR DRUG EVALUATION AND RESEARCH
OFFICE OF MANAGEMENT
DIVISION OF DRUG INFORMATION RESOURCES**

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APPROVED DRUG PRODUCTS

WITH
THERAPEUTIC EQUIVALENCE EVALUATIONS

**16TH EDITION
1996**

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**APPROVED DRUG PRODUCTS
with
THERAPEUTIC EQUIVALENCE EVALUATIONS**

15TH EDITION

Cumulative Supplement 12

DECEMBER 1995

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**APPROVED DRUG PRODUCTS
with
THERAPEUTIC EQUIVALENCE EVALUATIONS**

15TH EDITION

**CUMULATIVE SUPPLEMENT 12
DECEMBER 1995**

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, 15th 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and products that have never been marketed, have been discontinued from marketing or that have had their approvals withdrawn for other than safety or efficacy reasons.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.)

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.

Deletions new 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 15th Edition List will then be added to the "Discontinued Drug Product List" appearing in the 16th Edition.

1.2 COURT ORDER AFFECTING URUGUAY ROUND AGREEMENTS ACT-EXTENDED PATENTS

The Uruguay Round Agreements Act (URAA), Public Law 103-465, extended the term of patents issued on or after June 8, 1995, from 17 years from date of issue to 20 years from date of filing. Patents in effect on, or based upon applications filed by, June 8, 1995, are entitled to 17 years from date of issue or 20 years from date of filing, whichever is greater. On June 7, 1995, the Patent and Trademark Office (PTO) published a notice in the *Federal Register* (60 FR 30069) that established the method for calculating the patent term expiration date for any patent subject to both the terms of the URAA and the patent term extension provisions at title 35, U.S.C. § 156. FDA published a notice in the *Federal Register* on July 21, 1995, (60 FR 37652) announcing that it would not publish

in this publication patent expiration dates that the NDA applicant submitting the information stated were not calculated in accordance with the PTO method for determining the correct patent expiration date.

Both PTO's determination of the correct relationship between the extension of patents under the URAA and patent term extensions under title 35, U.S.C. § 156, and FDA's refusal to publish patent expiration dates that are not consistent with the PTO determination have been challenged. On October 16, 1995, the U.S. District Court for the Eastern District of Virginia issued an Opinion and Order finding that PTO had misinterpreted the Patent Code, and that PTO's determination of June 7, 1995, is invalid and unenforceable. The court ordered FDA to publish the patent dates it had, by its July 21, 1995, notice, refused to publish. Therefore, because of the court's order, and pending final resolution of appeals from the October 16, 1995, decision, FDA is publishing the patent term expiration dates that NDA applicants have told FDA are not consistent with PTO's June 7, 1995, determination. Because the district court decision may be reversed upon appeal, 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.

1.3 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.

1.4 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)

BOOTS PHARMACEUTICALS INC
(BOOTS)

KNOLL PHARMACEUTICAL COMPANY
SUB BASF CORPORATION
(KNOLL PHARM)

BRIAN PHARMACEUTICALS INC
(BRIAN)

HYGENICS PHARMACEUTICALS INC
(HYGENICS)

APPLICANT NAME CHANGES

FORMER APPLICANT NAME
(FORMER ABBREVIATED NAME)

NEW APPLICANT NAME
(NEW ABBREVIATED NAME)

BURROUGHS WELLCOME CO
(BURROUGHS WELLCOME)

GLAXO WELLCOME INC
(GLAXO WELLCOME)

DORSEY LABORATORIES DIV
SANDOZ WANDER INC
(DORSEY)

SANDOZ CONSUMER HEALTH CARE
GROUP DIV SANDOZ
PHARMACEUTICALS CORP
(SANDOZ)

FAULDING HOSPITAL PRODUCTS INC
(FAULDING)

FAULDING PHARMACEUTICAL CO
(FAULDING)

GLAXO INC
(GLAXO)

GLAXO WELLCOME INC
(GLAXO WELLCOME)

MARION MERRELL DOW INC
(MARION MERRELL DOW)

HOECHST MARION ROUSSEL INC
(HOECHST MARION RSSL)

MERRELL DOW PHARMACEUTICALS
INC SUB DOW CHEMICAL CO
(MERRELL DOW)

HOECHST MARION ROUSSEL INC
(HOECHST MARION RSSL)

MILES PHARMACEUTICAL DIV
MILES INC
(MILES)

BAYER CORPORATION
(BAYER)

PENNEX PHARMACEUTICALS INC
(PENNEX)

MORTON GROVE PHARMACEUTICALS INC
(MORTON GROVE)

TAP PHARMACEUTICALS INC
(TAP PHARMS)

TAP HOLDINGS INC
(TAP HOLDINGS)

1.5 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.

1.6 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 1994) 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 1994</u>	<u>JUN 1995</u>	<u>SEP 1995</u>	<u>DEC 1995</u>
DRUG PRODUCTS LISTED	9141	9221	9221	9286
SINGLE SOURCE	2178 (23.8%)	2186 (23.7%)	2168 (23.5%)	2217 (23.9%)
MULTISOURCE	6963 (76.2%)	7035 (76.3%)	7053 (76.5%)	7069 (76.1%)
THERAPEUTICALLY EQUIVALENT	6330 (69.2%)	6399 (69.4%)	6427 (69.7%)	6437 (69.3%)
NOT THERAPEUTICALLY EQUIVALENT	453 (5.0%)	452 (4.9%)	444 (4.8%)	440 (4.7%)
EXCEPTIONS ¹	180 (2.0%)	184 (2.0%)	182 (2.0%)	192 (2.1%)
NEW MOLECULAR ENTITIES APPROVED	--	10	6	14
NUMBER OF APPLICANTS	534	559	553	586

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

PRESCRIPTION DRUG PRODUCT LIST
15TH EDITION
RX DRUG PRODUCT LIST / CUMULATIVE SUPPLEMENT NUMBER 12 / JAN'95 - DEC'95

1

ACARBOSE

TABLET; ORAL
PRECOSE
BAYER

50MG

N20482 001
SEP 06, 1995
N20482 002
SEP 06, 1995

+

100MG

ACEBUTOLOL HYDROCHLORIDE

CAPSULE; ORAL
ACEBUTOLOL HCL

AB MYLAN

EQ 200MG BASE

N74288 001
APR 24, 1995

AB

EQ 400MG BASE

N74288 002
APR 24, 1995

AB

WATSON LABS

EQ 200MG BASE

N74007 001
OCT 18, 1995

AB

EQ 400MG BASE

N74007 002
OCT 18, 1995

SECTRAL

AB WYETH AYERST

EQ 200MG BASE

N18917 001
DEC 28, 1984

AB

+

EQ 400MG BASE

N18917 003
DEC 28, 1984

ACETAMINOPHEN; BUTALBITAL; CAFFEINE

TABLET; ORAL

BUTALBITAL, ACETAMINOPHEN AND CAFFEINE

AB WEST WARD PHARM

325MG; 50MG; 40MG

N89718 001
JUN 12, 1995

ACETAMINOPHEN; CODEINE PHOSPHATE

SOLUTION; ORAL

ACETAMINOPHEN AND CODEINE PHOSPHATE

AA BARRE

120MG/5ML; 12MG/5ML

N85861 001

AA

APAP W/ CODEINE

120MG/5ML; 12MG/5ML

N85861 001

TABLET; ORAL

ACETAMINOPHEN AND CODEINE PHOSPHATE

AA KV PHARM

300MG; 30MG

N85288 001

AA

300MG; 60MG

N85365 001

ACETAMINOPHEN; CODEINE PHOSPHATE

TABLET; ORAL

ACETAMINOPHEN AND CODEINE PHOSPHATE

KV PHARM

325MG; 15MG

N85364 001

@

325MG; 45MG

N85363 001

@

300MG; 30MG

N85288 001

@

300MG; 60MG

N85365 001

@

325MG; 15MG

N85364 001

@

325MG; 45MG

N85363 001

LEMMON

300MG; 15MG

N88627 001

> ADD > AA

> ADD >

> ADD > AA

> ADD >

> ADD > AA

> ADD >

> ADD > AA

MIKART

650MG; 30MG

N89231 001

+

650MG; 30MG

N89231 001

ROXANE

500MG; 15MG

N89511 001

+

500MG; 30MG

N89512 001

+

500MG; 60MG

N89513 001

@

500MG; 15MG

N89511 001

@

500MG; 30MG

N89512 001

@

500MG; 60MG

N89513 001

> DLT >

> DLT > AA

> DLT >

> DLT > AA

> DLT >

> DLT > AA

> DLT >

> DLT > AA

> DLT >

> DLT > AA

ACETAMINOPHEN W/ CODEINE #2

LEMMON

300MG; 15MG

N88627 001

ACETAMINOPHEN W/ CODEINE #3

LEMMON

300MG; 30MG

N88628 001

ACETAMINOPHEN W/ CODEINE #4

LEMMON

300MG; 60MG

N88629 001

PHENAPHEN-650 W/ CODEINE

AA ROBINS PH

650MG; 30MG

N85856 001

@

650MG; 30MG

N85856 001

ACETAMINOPHEN; HYDROCODONE BITARTRATE

ELIXIR; ORAL

HYDROCODONE BITARTRATE AND ACETAMINOPHEN

MIKART

500MG/15ML; 5MG/15ML N89557 001

APR 29, 1992

500MG/15ML; 7.5MG/15ML N81051 001

AUG 28, 1992

+

500MG/15ML; 5MG/15ML N89557 001

APR 29, 1992

+

500MG/15ML; 7.5MG/15ML N81051 001

AUG 28, 1992

TABLET; ORAL

ANEXSIA

AA BOEHRINGER MANNHEIM 500MG; 5MG

N89160 001

APR 23, 1987

AA KING PHARMS 500MG; 5MG

N89160 001

APR 23, 1987

ANEXSIA 7.5/650

AA BOEHRINGER MANNHEIM 650MG; 7.5MG

N89725 001

SEP 30, 1987

AA KING PHARMS 650MG; 7.5MG

N89725 001

SEP 30, 1987

HYDROCODONE BITARTRATE AND ACETAMINOPHEN

AA HALSEY 500MG; 5MG

N89554 001

JUN 12, 1987

@

500MG; 5MG N89554 001

JUN 12, 1987

AA KING PHARMS 500MG; 5MG

N40084 002

JUN 01, 1995

AA 750MG; 7.5MG

N40084 001

JUN 01, 1995

AA + MIKART 650MG; 10MG

N81223 001

MAY 29, 1992

AA WATSON LABS 650MG; 7.5MG

N40094 001

SEP 29, 1995

AA 650MG; 10MG

N40094 002

SEP 29, 1995

ACETAMINOPHEN; OXYCODONE HYDROCHLORIDE

CAPSULE; ORAL

ROXILOX

AA ROXANE 500MG; 5MG

N40061 001

JUL 03, 1995

ACETAMINOPHEN; OXYCODONE HYDROCHLORIDE

TABLET; ORAL

OXYCET

AA HALSEY 325MG; 5MG

N87463 001

DEC 07, 1983

AA MALLINCKRODT 325MG; 5MG

N87463 001

DEC 07, 1983

ACETAZOLAMIDE

CAPSULE, EXTENDED RELEASE; ORAL

DIAMOX

* STORZ OPHTHALM 500MG

N12945 001

@ 500MG

N12945 001

+ 500MG

N12945 003

APR 25, 1995

ACETAZOLAMIDE SODIUM

INJECTABLE; INJECTION

ACETAZOLAMIDE SODIUM

AP BEDFORD EQ 500MG BASE/VIAL

N40089 001

FEB 28, 1995

AP SANOFI WINTHROP EQ 500MG BASE/VIAL

N40108 001

OCT 30, 1995

DIAMOX

AP + STORZ OPHTHALM EQ 500MG BASE/VIAL

N09388 001

DEC 05, 1990

ACETYLCYSTEINE

SOLUTION; INHALATION, ORAL

ACETYLCYSTEINE

AN DUPONT MERCK 10%

N71364 001

MAY 01, 1989

AN 20%

N71365 001

MAY 01, 1989

AN FAULDING 10%

N71364 001

MAY 01, 1989

AN 20%

N71365 001

MAY 01, 1989

AN LUITPOLD 10%

N72489 001

JUL 28, 1995

AN 20%

N72547 001

JUL 28, 1995

ADENOSINE

INJECTABLE; INJECTION
ADENOSCAN
+ MEDCO RES

3MG/ML

N20059 001
MAY 18, 1995

ALBUTEROL

AEROSOL, METERED; INHALATION

> ADD >
> ADD > AB ALBUTEROL
> ADD > NORTON WATERFORD 0.09MG/INH

N73272 001
DEC 28, 1995

> DLT > BN * GLAXO 0.09MG/INH
> ADD > AB + GLAXO WELLCOME 0.09MG/INH

N18473 001
N18473 001

ALBUTEROL SULFATE

SOLUTION; INHALATION

> ADD > AN ALBUTEROL SULFATE
> ADD > BARRE EQ 0.083% BASE
> DLT > AN PACO EQ 0.083% BASE
> DLT >

N73533 001
SEP 26, 1995
N73533 001
SEP 26, 1995

SYRUP; ORAL

AA ALBUTEROL SULFATE
BARRE EQ 2MG BASE/5ML

N74454 001
SEP 25, 1995

ALENDRONATE SODIUM

TABLET; ORAL
FOSAMAX
MERCK

EQ 10MG BASE

N20560 001
SEP 29, 1995
N20560 002
SEP 29, 1995

+

EQ 40MG BASE

ALPRAZOLAM

TABLET; ORAL

AB ALPRAZOLAM
CHELSEA LABS 0.25MG

N74456 001
AUG 31, 1995

ALPRAZOLAM

TABLET; ORAL

AB ALPRAZOLAM
CHELSEA LABS 0.5MG

N74456 002
AUG 31, 1995

AB 1MG

N74456 003
AUG 31, 1995

> ADD > AB GENEVA PHARMS 0.25MG

> ADD >
> ADD > AB 0.5MG

N74112 001
DEC 29, 1995

> ADD >
> ADD > AB 1MG

N74112 002
DEC 29, 1995
N74112 003
DEC 29, 1995

ALPROSTADIL

INJECTABLE; INJECTION
CAVERJECT
UPJOHN

0.01MG/VIAL

N20379 001
JUL 06, 1995

+

0.02MG/VIAL

N20379 002
JUL 06, 1995

ALSEROXYLON

TABLET; ORAL

RAUWILOID
+ 3M 2MG
@ 2MG

N08867 001
N08867 001

AMANTADINE HYDROCHLORIDE

SYRUP; ORAL

AA AMANTADINE HCL
PHARM ASSOC 50MG/5ML

N74509 001
JUL 17, 1995

> ADD > AMIFOSTINE

> ADD > INJECTABLE; INJECTION

> ADD > ETHYOL
> ADD > + US BIOSCIENCE 500MG/VIAL

N20221 001
DEC 08, 1995

AMIKACIN SULFATE

INJECTABLE; INJECTION

AMIKACIN

AP	DUPONT MERCK	EQ 50MG BASE/ML	N63350 001
			JUL 30, 1993
AP		EQ 250MG BASE/ML	N63350 002
			JUL 30, 1993
AP	* ELKINS SINN	EQ 50MG BASE/ML	N63274 001
			MAY 18, 1992
AP	*	EQ 250MG BASE/ML	N63275 001
			MAY 18, 1992
<u>AMIKACIN SULFATE</u>			
> ADD >	AP	ASTRA	EQ 50MG BASE/ML
> ADD >			N63167 001
> ADD >			DEC 14, 1995
> ADD >	AP		EQ 250MG BASE/ML
			N63169 001
			DEC 14, 1995
AP	ELKINS SINN	EQ 50MG BASE/ML	N63274 001
			MAY 18, 1992
AP		EQ 250MG BASE/ML	N63275 001
			MAY 18, 1992
AP	FAULDING	EQ 50MG BASE/ML	N63350 001
			JUL 30, 1993
AP		EQ 250MG BASE/ML	N63350 002
			JUL 30, 1993
AP	SANOFI WINTHROP	EQ 250MG BASE/ML	N64098 001
			JUN 26, 1995
AP		EQ 250MG BASE/ML	N64099 001
			JUN 20, 1995
<u>AMIKIN</u>			
AP	APOTHECON	EQ 50MG BASE/ML	N62311 001
AP	+	EQ 50MG BASE/ML	N62311 001
AP		EQ 250MG BASE/ML	N62311 002
AP	+	EQ 250MG BASE/ML	N62311 002
	@	EQ 50MG BASE/ML	N62562 001
			SEP 20, 1984
	@	EQ 250MG BASE/ML	N62562 002
			SEP 20, 1984
	@ BRISTOL	EQ 50MG BASE/ML	N62562 001
			SEP 20, 1984
	@	EQ 250MG BASE/ML	N62562 002
			SEP 20, 1984
AMIKIN IN SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER			
	@ APOTHECON	EQ 5MG BASE/ML	N50618 002
			NOV 30, 1987
	@	EQ 10MG BASE/ML	N50618 001
			NOV 30, 1987
	@ BRISTOL	EQ 5MG BASE/ML	N50618 002
			NOV 30, 1987

AMIKACIN SULFATE

INJECTABLE; INJECTION

AMIKIN IN SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER

@ BRISTOL	EQ 10MG BASE/ML	N50618 001
		NOV 30, 1987

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

INJECTABLE; INJECTION

TRAVASOL 2.75% SULFITE FREE W/ ELECTROLYTES IN DEXTROSE 10% IN PLASTIC CONTAINER

BAXTER	2.75%; 10GM/100ML; 51MG/100ML; 261MG/100ML; 216MG/100ML; 112MG/100ML	N20147 002
		OCT 23, 1995

TRAVASOL 2.75% SULFITE FREE W/ ELECTROLYTES IN DEXTROSE 15% IN PLASTIC CONTAINER

BAXTER	2.75%; 15GM/100ML; 51MG/100ML; 261MG/100ML; 216MG/100ML; 112MG/100ML	N20147 003
		OCT 23, 1995

TRAVASOL 2.75% SULFITE FREE W/ ELECTROLYTES IN DEXTROSE 20% IN PLASTIC CONTAINER

BAXTER	2.75%; 20GM/100ML; 51MG/100ML; 261MG/100ML; 216MG/100ML; 112MG/100ML	N20147 004
		OCT 23, 1995

TRAVASOL 2.75% SULFITE FREE W/ ELECTROLYTES IN DEXTROSE 25% IN PLASTIC CONTAINER

BAXTER	2.75%; 25GM/100ML; 51MG/100ML; 261MG/100ML; 216MG/100ML; 112MG/100ML	N20147 005
		OCT 23, 1995

TRAVASOL 2.75% SULFITE FREE W/ ELECTROLYTES IN DEXTROSE 5% IN PLASTIC CONTAINER

BAXTER	2.75%; 5GM/100ML; 51MG/100ML; 261MG/100ML; 216MG/100ML; 112MG/100ML	N20147 001
		OCT 23, 1995

TRAVASOL 4.25% SULFITE FREE W/ ELECTROLYTES IN DEXTROSE 10% IN PLASTIC CONTAINER

BAXTER	4.25%; 10GM/100ML; 51MG/100ML; 261MG/100ML; 297MG/100ML; 77MG/100ML	N20147 007
		OCT 23, 1995

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

INJECTABLE; INJECTION

TRAVASOL 4.25% SULFITE FREE W/ ELECTROLYTES IN DEXTROSE 15%
IN PLASTIC CONTAINER
BAXTER

4.25%;15GM/100ML;51MG/100ML;
261MG/100ML;297MG/100ML;
77MG/100ML N20147 008
OCT 23, 1995

TRAVASOL 4.25% SULFITE FREE W/ ELECTROLYTES IN DEXTROSE 20%
IN PLASTIC CONTAINER
BAXTER

4.25%;20GM/100ML;51MG/100ML;
261MG/100ML;297MG/100ML;
77MG/100ML N20147 009
OCT 23, 1995

TRAVASOL 4.25% SULFITE FREE W/ ELECTROLYTES IN DEXTROSE 25%
IN PLASTIC CONTAINER
BAXTER

4.25%;25GM/100ML;51MG/100ML;
261MG/100ML;297MG/100ML;
77MG/100ML N20147 010
OCT 23, 1995

TRAVASOL 4.25% SULFITE FREE W/ ELECTROLYTES IN DEXTROSE 5%
IN PLASTIC CONTAINER
BAXTER

4.25%;5GM/100ML;51MG/100ML;
261MG/100ML;297MG/100ML;
77MG/100ML N20147 006
OCT 23, 1995

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

INJECTABLE; INJECTION

TRAVASOL 3.5% SULFITE FREE W/ ELECTROLYTES IN PLASTIC
CONTAINER
BAXTER

3.5%;51MG/100ML;131MG/100ML;
218MG/100ML;35MG/100ML N20177 001
OCT 23, 1995

TRAVASOL 5.5% SULFITE FREE W/ ELECTROLYTES IN PLASTIC
CONTAINER
BAXTER

5.5%;102MG/100ML;522MG/100ML;
431MG/100ML;224MG/100ML N20173 001
OCT 27, 1995

TRAVASOL 8.5% SULFITE FREE W/ ELECTROLYTES IN PLASTIC
CONTAINER
BAXTER

8.5%;102MG/100ML;522MG/100ML;
594MG/100ML;154MG/100ML N20173 002
OCT 27, 1995

AMINOPHYLLINE

INJECTABLE; INJECTION

AMINOPHYLLINE

AP FUJISAWA 25MG/ML N87200 001
AP 25MG/ML N88407 001
JAN 25, 1984

@ 25MG/ML N87200 001
@ 25MG/ML N88407 001
JAN 25, 1984

AP KING PHARMS 25MG/ML N86606 001
@ 25MG/ML N86606 001
AMINOPHYLLINE IN SODIUM CHLORIDE 0.45%
ABBOTT 100MG/100ML N88147 002
MAY 03, 1983

200MG/100ML N88147 003
MAY 03, 1982
+ 100MG/100ML N88147 002
MAY 03, 1983
+ 200MG/100ML N88147 003
MAY 03, 1983

TABLET; ORAL

AMINOPHYLLINE

BD PHOENIX LABS NY 100MG N85409 001
BD 200MG N85410 001
@ 100MG N85409 001
@ 200MG N85410 001

AMINOSALICYLATE SODIUM

POWDER; ORAL

P.A.S. SODIUM

AA CENTURY PHARMS 4GM/PACKET N80947 001
@ 4GM/PACKET N80947 001

AA SODIUM AMINOSALICYLATE 100% N80097 001
@ 100% N80097 001

AMINOSALICYLATE SODIUM; AMINOSALICYLIC ACID

TABLET; ORAL

NEOPASALATE

* WALLACE 846MG;112MG N80059 002
@ 846MG;112MG N80059 002

AMIODARONE HYDROCHLORIDEINJECTABLE; INJECTION
CORDARONE

+ WYETH AYERST 50MG/ML

N20377 001
AUG 03, 1995AMITRIPTYLINE HYDROCHLORIDE

TABLET; ORAL

AMITRIPTYLINE HCLAB ROXANE 150MG
@ 150MGN86090 001
N86090 001AMLODIPINE BESYLATE; BENAZEPRIL HYDROCHLORIDE

CAPSULE; ORAL

LOTREL

CIBA GEIGY

EQ 2.5MG BASE;10MG

N20364 002
MAR 03, 1995

EQ 5MG BASE;10MG

N20364 003

MAR 03, 1995

N20364 004

MAR 03, 1995

+ EQ 5MG BASE;20MG

> ADD >

> ADD >

> ADD >

AMMONIUM LACTATE

LOTION; TOPICAL

LAC-HYDRIN

* WESTWOOD SQUIRE

EQ 12% ACID

N19155 001
APR 24, 1985

+ EQ 12% BASE

N19155 001
APR 24, 1985

> DLT >

> DLT >

> DLT >

> ADD >

> ADD >

> ADD >

AMOXICILLIN

CAPSULE; ORAL

AMOXICILLINAB CONSOLIDATED PHARM 250MG
AB 500MGN62058 001
N62058 002COMOXAB COPANOS 250MG
AB 500MGN62058 001
N62058 002POLYMOXAB APOTHECON 250MGN63099 001
MAR 20, 1992

> DLT >

> DLT >

> DLT >

> ADD >

> ADD >

> ADD >

> ADD >

> DLT >

> DLT >

AMOXICILLIN

CAPSULE; ORAL

POLYMOXAB APOTHECON 500MGN63099 002
MAR 20, 1992TRIMOXAB APOTHECON 250MGN63099 001
MAR 20, 1992AB 500MGN63099 002
MAR 20, 1992

POWDER FOR RECONSTITUTION; ORAL

AMOXICILLINAB CONSOLIDATED PHARM 125MG/5ML
AB 250MG/5MLN62059 001
N62059 002AMOXICILLIN TRIHYDRATEAB COPANOS 125MG/5ML
AB 250MG/5MLN62059 001
N62059 002

TABLET, CHEWABLE; ORAL

AMOXICILLINAB BIOCRAFT 125MGN64013 002
SEP 11, 1995AMOXILAB + SMITHKLINE BEECHAM 125MG

N50542 002

AMOXICILLIN; CLAVULANATE POTASSIUM

POWDER FOR RECONSTITUTION; ORAL

AUGMENTIN '125'

* SMITHKLINE BEECHAM 125MG/5ML;

EQ 31.25MG ACID/5ML

N50575 001
AUG 06, 1984

+ 125MG/5ML;

EQ 31.25MG BASE/5ML

N50575 001
AUG 06, 1984

AUGMENTIN '250'

* SMITHKLINE BEECHAM 250MG/5ML;

EQ 62.5MG ACID/5ML

N50575 002
AUG 06, 1984

+ 250MG/5ML;

EQ 62.5MG BASE/5ML

N50575 002
AUG 06, 1984

TABLET; ORAL

AUGMENTIN '250'

* SMITHKLINE BEECHAM 250MG; EQ 125MG ACIDN50564 001
AUG 06, 1984

AMOXICILLIN; CLAVULANATE POTASSIUM

TABLET; ORAL

> ADD > AUGMENTIN '250'
 + SMITHKLINE BEECHAM 250MG;EQ 125MG BASE N50564 001
 > ADD > AUG 06, 1984
 > DLT > AUGMENTIN '500'
 * SMITHKLINE BEECHAM 500MG;EQ 125MG ACID N50564 002
 > DLT > AUG 06, 1984
 > ADD > + 500MG;EQ 125MG BASE N50564 002
 > ADD > AUG 06, 1984

TABLET, CHEWABLE; ORAL

> DLT > AUGMENTIN '125'
 * SMITHKLINE BEECHAM 125MG;EQ 31.25MG ACID N50597 001
 > DLT > JUL 22, 1985
 > ADD > + 125MG;EQ 31.25MG BASE N50597 001
 > ADD > JUL 22, 1985
 > DLT > AUGMENTIN '250'
 * SMITHKLINE BEECHAM 250MG;EQ 62.5MG ACID N50597 002
 > DLT > JUL 22, 1985
 > ADD > + 250MG;EQ 62.5MG BASE N50597 002
 > ADD > JUL 22, 1985

AMPHOTERICIN B

INJECTABLE; INJECTION

> DLT > ABELCET
 > DLT > * LIPOSOME 5MG/ML N50724 001
 > DLT > NOV 20, 1995

AMPHOTERICIN B

AP GENSLA 50MG/VIAL N64062 001
 MAR 31, 1995

> ADD > INJECTABLE, LIPID COMPLEX; INJECTION
 > ADD > ABELCET
 > ADD > + LIPOSOME 5MG/ML N50724 001
 > ADD > NOV 20, 1995

AMPICILLIN SODIUM

INJECTABLE; INJECTION

AP AMPICILLIN SODIUM
 AP APOTHECON EQ 125MG BASE/VIAL N61395 001
 EQ 125MG BASE/VIAL N62860 001
 FEB 05, 1988
 AP EQ 250MG BASE/VIAL N61395 002

AMPICILLIN SODIUM

INJECTABLE; INJECTION

AP AMPICILLIN SODIUM
 APOTHECON EQ 250MG BASE/VIAL N62860 002
 FEB 05, 1988
 AP EQ 500MG BASE/VIAL N61395 003
 AP EQ 500MG BASE/VIAL N62860 003
 FEB 05, 1988
 AP EQ 1GM BASE/VIAL N61395 004
 AP EQ 1GM BASE/VIAL N62738 001
 FEB 19, 1987
 AP EQ 1GM BASE/VIAL N62860 004
 FEB 05, 1988
 AP EQ 2GM BASE/VIAL N61395 005
 AP EQ 2GM BASE/VIAL N62738 002
 FEB 19, 1987
 AP EQ 2GM BASE/VIAL N62860 005
 FEB 05, 1988
 AP + EQ 10GM BASE/VIAL N61395 006
 @ CONSOLIDATED PHARM EQ 125MG BASE/VIAL N61936 005
 @ EQ 250MG BASE/VIAL N61936 001
 @ EQ 500MG BASE/VIAL N61936 002
 @ EQ 1GM BASE/VIAL N61936 003
 @ EQ 2GM BASE/VIAL N61936 004
 @ COPANOS EQ 125MG BASE/VIAL N61936 005
 @ EQ 250MG BASE/VIAL N61936 001
 @ EQ 500MG BASE/VIAL N61936 002
 @ EQ 1GM BASE/VIAL N61936 003
 @ EQ 2GM BASE/VIAL N61936 004
 AP PRINCIPEN
 APOTHECON EQ 125MG BASE/VIAL N61395 001
 AP EQ 125MG BASE/VIAL N62860 001
 FEB 05, 1988
 AP EQ 250MG BASE/VIAL N61395 002
 AP EQ 250MG BASE/VIAL N62860 002
 FEB 05, 1988
 AP EQ 500MG BASE/VIAL N61395 003
 AP EQ 500MG BASE/VIAL N62860 003
 FEB 05, 1988
 AP EQ 1GM BASE/VIAL N61395 004
 AP EQ 1GM BASE/VIAL N62738 001
 FEB 19, 1987
 AP EQ 1GM BASE/VIAL N62860 004
 FEB 05, 1988
 AP EQ 2GM BASE/VIAL N61395 005
 AP EQ 2GM BASE/VIAL N62738 002
 FEB 19, 1987

AMPICILLIN SODIUM

INJECTABLE; INJECTION

PRINCIPENAP APOTHECONEQ 2CM BASE/VIALN62860 005

FEB 05, 1988

AP *EQ 10CM BASE/VIALN61395 006AMPICILLIN/AMPICILLIN TRIHYDRATE

CAPSULE; ORAL

AMPICILLIN TRIHYDRATEAB BIOCHEMIEEQ 250MG BASEN64082 001

AUG 29, 1995

ABEQ 500MG BASEN64082 002

AUG 29, 1995

AB CONSOLIDATED PHARMEQ 250MG BASEN61602 001ABEQ 500MG BASEN61602 002AB COPANOSEQ 250MG BASEN61602 003ABEQ 500MG BASEN61602 002

POWDER FOR RECONSTITUTION; ORAL

AMPICILLIN TRIHYDRATEAB CONSOLIDATED PHARMEQ 125MG BASE/5MLN61601 001ABEQ 250MG BASE/5MLN61601 002AB COPANOSEQ 125MG BASE/5MLN61601 003ABEQ 250MG BASE/5MLN61601 002POLYCILLINAB * BRISTOLEQ 125MG BASE/5MLN50308 001AB *EQ 250MG BASE/5MLN50308 002AB *EQ 500MG BASE/5MLN50308 003AB *EQ 125MG BASE/5MLN50308 001AB *EQ 250MG BASE/5MLN50308 002AB *EQ 500MG BASE/5MLN50308 003> ADD > ANASTROZOLE> ADD > TABLET; ORAL> ADD > ARIMIDEX> ADD > + ZENECA1MGN20541 001

DEC 27, 1995

ANTAZOLINE PHOSPHATE; NAPHAZOLINE HYDROCHLORIDE

SOLUTION/DROPS; OPHTHALMIC

VASOCON-A* CIBA VISION0.5%;0.05%N18746 001

APR 30, 1990

ASPIRIN; BUTALBITAL; CAFFEINE; CODEINE PHOSPHATE

CAPSULE; ORAL

BUTALBITAL, ASPIRIN, CAFFEINE, AND CODEINE PHOSPHATEAB WATSON LABS325MG;50MG;40MG;30MGN74359 001

AUG 31, 1995

FIORINAL W/CODEINE NO 3AB + SANDOZ325MG;50MG;40MG;30MGN19429 003

OCT 26, 1990

ASPIRIN; METHOCARBAMOL

TABLET; ORAL

METHOCARBAMOL AND ASPIRINAB STEVENS J325MG;400MGN81145 001

JAN 31, 1995

ATENOLOL

TABLET; ORAL

ATENOLOLAB COPLY PHARM50MGN74120 001

FEB 24, 1995

AB100MGN74120 002

FEB 24, 1995

ABLEMMON50MGN74056 001

JAN 18, 1995

AB100MGN74056 002

JAN 18, 1995

ABMARTEC50MGN74127 001

FEB 21, 1995

AB100MGN74127 002

FEB 21, 1995

ATOVAQUONE

SUSPENSION; ORAL
MEPRON
+ GLAXO WELLCOME

750MG/5ML

N20500 001
FEB 08, 1995

ATROPINE

INJECTABLE; INJECTION
ATROPEN

AP * SURVIVAL TECH
+

EQ 2MG SULFATE/0.7ML
EQ 2MG SULFATE/0.7ML

N17106 001
N17106 001

AP ATROPINE
KALI DUPHAR

EQ 2MG SULFATE/0.7ML

N71295 001
JAN 30, 1987
N71295 001
JAN 30, 1987

© SOLVAY

EQ 2MG SULFATE/0.7ML

ATROPINE SULFATE; DIPHENOXYLATE HYDROCHLORIDE

TABLET; ORAL

DI-ATRO

AA MD PHARM
©

0.025MG;2.5MG
0.025MG;2.5MG

N85266 001
N85266 001

AZATHIOPRINE SODIUM

INJECTABLE; INJECTION
AZATHIOPRINE SODIUM

AP BEDFORD

EQ 100MG BASE/VIAL

N74419 001
MAR 31, 1995

IMURAN

AP + GLAXO WELLCOME

EQ 100MG BASE/VIAL

N17391 001

AZELAIC ACID

CREAM; TOPICAL

AZELEX

+ ALLERGAN

20%

N20428 001
SEP 13, 1995

AZITHROMYCIN DIHYDRATE

POWDER FOR RECONSTITUTION; ORAL

ZITHROMAX

+ PFIZER

EQ 100MG BASE/5ML

N50710 001

+

EQ 200MG BASE/5ML

OCT 19, 1995
N50710 002
OCT 19, 1995

BACITRACIN

INJECTABLE; INJECTION

BACITRACIN

AP * PFIZER

50,000 UNITS/VIAL

N60282 001

@

50,000 UNITS/VIAL

N60282 001

AP UPJOHN

50,000 UNITS/VIAL

N60733 002

+

50,000 UNITS/VIAL

N60733 002

POWDER; FOR RX COMPOUNDING

BACITRACIN

> ADD >

AA

APOTHEKERNES

5,000,000 UNITS/BOT

N61699 001

> DLT >

AA

BRAE

5,000,000 UNITS/BOT

N61699 001

BACITRACIN ZINC; HYDROCORTISONE; NEOMYCIN SULFATE; POLYMYXIN B SULFATE

OINTMENT; OPHTHALMIC

CORTISPORIN

AT + GLAXO WELLCOME

400 UNITS/GM;1%;EQ 3.5MG BASE/GM;

10,000 UNITS/GM

N50416 002

NEOMYCIN AND POLYMYXIN B SULFATES, BACITRACIN ZINC AND

AT HYDROCORTISONE

400 UNITS/GM;1%;EQ 3.5MG BASE/GM;

10,000 UNITS/GM

N64068 001

OCT 30, 1995

BACITRACIN ZINC; NEOMYCIN SULFATE; POLYMYXIN B SULFATE

OINTMENT; OPHTHALMIC

NEOMYCIN AND POLYMYXIN B SULFATES AND BACITRACIN ZINC

AT BAUSCH AND LOMB

400 UNITS/GM;EQ 3.5MG BASE/GM;

10,000 UNITS/GM

N64064 001

OCT 30, 1995

BACITRACIN ZINC; POLYMYXIN B SULFATE

OINTMENT; OPHTHALMIC

BACITRACIN ZINC AND POLYMYXIN B SULFATE

<u>AT</u>	ADV REMEDIES	500 UNITS/GM; 10,000 UNITS/GM	N64028 001 JAN 30, 1995
<u>AT</u>	BAUSCH AND LOMB	500 UNITS/GM; 10,000 UNITS/GM	N64046 001 JAN 26, 1995
<u>AT</u>	<u>POLYSPORIN</u> + GLAXO WELLCOME	500 UNITS/GM; 10,000 UNITS/GM	N61229 001

BENDROFLUMETHIAZIDETABLET; ORAL
NATURETIN-10

+	APOTHECON	10MG	N12164 003
*	SQUIBB	10MG	N12164 003
	NATURETIN-2.5		
@	APOTHECON	2.5MG	N12164 001
@	SQUIBB	2.5MG	N12164 001
	NATURETIN-5		
	APOTHECON	5MG	N12164 002
	SQUIBB	5MG	N12164 002

BENTONITE; SULFUR

POWDER; TOPICAL

	BENSULPOID		
@	POYTHRESS	66.64%; 33.32%	N62918 001

BETAMETHASONE BENZOATE

GEL; TOPICAL

	UTICORT		
+	PARKE DAVIS	0.025%	N17244 001
@		0.025%	N17244 001

LOTION; TOPICAL

	UTICORT		
+	PARKE DAVIS	0.025%	N17528 001
@		0.025%	N17528 001

BETAMETHASONE DIPROPIONATE

OINTMENT, AUGMENTED; TOPICAL

BETAMETHASONE DIPROPIONATE

<u>AB</u>	NMC	EQ 0.05% BASE	N74304 001 AUG 31, 1995
<u>AB</u>	<u>DIPROLENE</u> + SCHERING	EQ 0.05% BASE	N18741 001 JUL 27, 1983

BICALUTAMIDE

TABLET; ORAL

	CASODEX		
+	ZENECA	50MG	N20498 001 OCT 04, 1995

BRETYLIUM TOSYLATE

INJECTABLE; INJECTION

BRETYLOL

<u>AP</u>	DUPONT MERCK	50MG/ML	N17954 001
<u>AP</u>	FAULDING	50MG/ML	N17954 001

BROMPHENIRAMINE MALEATE; CODEINE PHOSPHATE; PHENYLPROPANOLAMINE HYDROCHLORIDE

SYRUP; ORAL

DIMETANE-DC

<u>AA</u>	ROBINS AM	2MG/5ML; 10MG/5ML; 12.5MG/5ML	N11694 006 MAR 29, 1984
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<u>AA</u>	+	2MG/5ML; 10MG/5ML; 12.5MG/5ML	N11694 006 MAR 29, 1984
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BUMETANIDE

INJECTABLE; INJECTION

BUMETANIDE

<u>AP</u>	BEDFORD	0.25MG/ML	N74441 001 JAN 27, 1995
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TABLET; ORAL

BUMETANIDE

<u>AB</u>	ZENITH LABS	0.5MG	N74225 001 APR 24, 1995
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BUMETANIDE

TABLET; ORAL

BUMETANIDE

<u>AB</u>	ZENITH LABS	<u>1MG</u>	N74225 002
			APR 24, 1995
<u>AB</u>		<u>2MG</u>	N74225 003
			APR 24, 1995
<u>AB</u>	<u>BUMEX</u>	<u>0.5MG</u>	N18225 002
	ROCHE		FEB 28, 1983
<u>AB</u>		<u>1MG</u>	N18225 001
			FEB 28, 1983
<u>AB</u>	+	<u>2MG</u>	N18225 003
			JUN 14, 1985

BUPIVACAINE HYDROCHLORIDE

INJECTABLE; INJECTION

MARCAINE HCL

<u>AP</u>	+	SANOFI WINTHROP	<u>0.25%</u>	N16964 001
<u>AP</u>	+		<u>0.5%</u>	N16964 006
<u>AP</u>	+	STERLING WINTHROP	<u>0.25%</u>	N16964 001
<u>AP</u>	+		<u>0.5%</u>	N16964 006
<u>AP</u>	+		<u>0.75%</u>	N16964 009
		<u>MARCAINE HCL PRESERVATIVE FREE</u>		
<u>AP</u>	+	SANOFI WINTHROP	<u>0.25%</u>	N16964 012
<u>AP</u>	+		<u>0.5%</u>	N16964 005
<u>AP</u>	+		<u>0.75%</u>	N16964 009

BUPIVACAINE HYDROCHLORIDE; EPINEPHRINE BITARTRATE

INJECTABLE; INJECTION

MARCAINE HCL W/ EPINEPHRINE

<u>AP</u>	+	SANOFI WINTHROP	<u>0.25%; 0.0091MG/ML</u>	N16964 004
<u>AP</u>	+		<u>0.5%; 0.0091MG/ML</u>	N16964 008
<u>AP</u>	+	STERLING WINTHROP	<u>0.25%; 0.0091MG/ML</u>	N16964 004
<u>AP</u>	+		<u>0.5%; 0.0091MG/ML</u>	N16964 008
<u>AP</u>	+		<u>0.75%; 0.0091MG/ML</u>	N16964 010
		<u>MARCAINE HCL W/ EPINEPHRINE PRESERVATIVE FREE</u>		
<u>AP</u>	+	SANOFI WINTHROP	<u>0.25%; 0.0091MG/ML</u>	N16964 013
<u>AP</u>	+		<u>0.5%; 0.0091MG/ML</u>	N16964 007
<u>AP</u>	+		<u>0.75%; 0.0091MG/ML</u>	N16964 010

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

BUTORPHANOL TARTRATE

INJECTABLE; INJECTION

STADOL

+	APOTHECON	<u>1MG/ML</u>	N17857 001
+		<u>2MG/ML</u>	N17857 002
+		<u>2MG/ML</u>	N17857 004
	<u>STADOL PRESERVATIVE FREE</u>		
+	APOTHECON	<u>1MG/ML</u>	N17857 001
+		<u>2MG/ML</u>	N17857 002

CAFFEINE; ERGOTAMINE TARTRATE

TABLET; ORAL

CAFERGOT

<u>AA</u>	+	SANDOZ	<u>100MG; 1MG</u>	N06620 001
		@	<u>100MG; 1MG</u>	N06620 001
		<u>ERCATAB</u>		
<u>AA</u>		GENEVA PHARMS	<u>100MG; 1MG</u>	N84294 001
<u>AA</u>	+		<u>100MG; 1MG</u>	N84294 001

CALCITONIN, SALMON

INJECTABLE; INJECTION

CALCITONIN-SALMON

<u>AP</u>		ASTRA	<u>200 IU/ML</u>	N73690 001
				APR 14, 1995

SPRAY, METERED; NASAL

MIACALCIN

+	SANDOZ	<u>200 IU/INH</u>	N20313 002
			AUG 17, 1995

CAPTOPRIL

TABLET; ORAL

CAPOTEN

<u>AB</u>		BRISTOL MYERS SQUIBB	<u>12.5MG</u>	N18343 005
				JAN 17, 1985
<u>AB</u>	+		<u>25MG</u>	N18343 002
<u>AB</u>			<u>50MG</u>	N18343 001
<u>AB</u>			<u>100MG</u>	N18343 003
			<u>150MG</u>	N18343 004
				JUN 13, 1995
			<u>75MG</u>	N18343 007
				JUN 13, 1995

CAPTOPRIL

TABLET; ORAL
CAPTOPRIL
AB APOTHECON 12.5MG
AB 25MG
AB 50MG
AB 100MG
AB GENEVA PHARMS 12.5MG
AB 25MG
AB 50MG
AB 100MG

N74472 001 > DLT >
MAR 31, 1995 > DLT >
N74472 002 > ADD >
MAR 31, 1995 > ADD >
N74472 003
MAR 31, 1995
N74472 004
MAR 31, 1995
N74363 001
NOV 09, 1995
N74363 002
NOV 09, 1995
N74363 003 > DLT >
NOV 09, 1995 > ADD >
N74363 004
NOV 09, 1995

CARBAMAZEPINE

SUSPENSION; ORAL
TEGRETOL
* BASEL PHARMS 100MG/5ML N18927 001
DEC 18, 1987
+ CIBA GEIGY 100MG/5ML N18927 001
DEC 18, 1987
TABLET; ORAL
TEGRETOL
AB * BASEL PHARMS 200MG N16608 001
AB + CIBA GEIGY 200MG N16608 001
TABLET, CHEWABLE; ORAL
TEGRETOL
AB * BASEL PHARMS 100MG N18281 001
AB + CIBA GEIGY 100MG N18281 001

CAPTOPRIL; HYDROCHLOROTHIAZIDE

TABLET; ORAL
CAPOZIDE 25/25
SQUIBB 25MG; 25MG
+ 25MG; 25MG
CAPOZIDE 50/15
SQUIBB 50MG; 15MG
+ 50MG; 15MG
CAPOZIDE 50/25
* SQUIBB 50MG; 25MG
50MG; 25MG

N18709 002
OCT 12, 1984
N18709 002
OCT 12, 1984
N18709 004
OCT 12, 1984
N18709 004
OCT 12, 1984
N18709 003
OCT 12, 1984
N18709 003
OCT 12, 1984

CARBIDOPA; LEVODOPA

TABLET; ORAL
CARBIDOPA AND LEVODOPA
AB GENEVA PHARMS 10MG; 100MG N73586 001
JUN 29, 1995
AB 25MG; 100MG N73587 001
JUN 29, 1995
AB 25MG; 250MG N73620 001
JUN 29, 1995

CARTEOLOL HYDROCHLORIDE

SOLUTION/DROPS; OPHTHALMIC
OCUPRESS
+ OTSUKA 1% N19972 001
MAY 23, 1990
OPTIPRESS
* OTSUKA 1% N19972 001
MAY 23, 1990

CARBACHOL

SOLUTION; INTRAOCULAR
CARBASTAT
AT CIBA 0.01%
MIOSTAT
AT + ALCON 0.01%

N73677 001
APR 28, 1995
N16968 001

CARVEDILOL

TABLET; ORAL
COREG
SMITHKLINE BEECHAM 6.25MG N20297 003
SEP 14, 1995

CARVEDILOLTABLET; ORAL
COREGSMITHKLINE BEECHAM 12.5MG
+ 25MGN20297 002
SEP 14, 1995
N20297 001
SEP 14, 1995CEFACLOL

CAPSULE; ORAL

CECLOR

AB + LILLY

EQ 250MG BASE

N50521 001

AB

EQ 250MG BASE

N62205 001

AB +

EQ 500MG BASE

N50521 002

AB

EQ 500MG BASE

N62205 002

CEFACLOL

AB LEDERLE

EQ 250MG BASEN64107 001
APR 27, 1995

AB

EQ 500MG BASE

N64107 002

AB ZENITH LABS

EQ 250MG BASE

APR 27, 1995

AB

EQ 500MG BASEN64061 001
APR 27, 1995
N64061 002
APR 27, 1995

POWDER FOR RECONSTITUTION; ORAL

CECLOR

AB + LILLY

EQ 125MG BASE/5ML

N50522 001

AB

EQ 125MG BASE/5ML

N62206 001

AB +

EQ 187MG BASE/5ML

N62206 003

AB +

EQ 250MG BASE/5ML

APR 20, 1988

AB

EQ 250MG BASE/5ML

N50522 002

AB +

EQ 375MG BASE/5ML

N62206 002

CEFACLOL

AB LEDERLE

EQ 125MG BASE/5MLN64114 001
APR 28, 1995

AB

EQ 187MG BASE/5ML

N64115 001

AB

EQ 250MG BASE/5ML

APR 28, 1995

AB

EQ 375MG BASE/5ML

N64116 001

AB ZENITH LABS

EQ 125MG BASE/5MLAPR 28, 1995
N64110 001
APR 28, 1995
N64087 001
APR 28, 1995CEFACLOL

POWDER FOR RECONSTITUTION; ORAL

CEFACLOL

AB ZENITH LABS

EQ 187MG BASE/5ML

N64086 001

AB

EQ 250MG BASE/5ML

APR 28, 1995

AB

EQ 375MG BASE/5ML

N64085 001

APR 28, 1995

N64070 001

APR 28, 1995

CEFAMANDOLE NAFATE

INJECTABLE; INJECTION

MANDOL

+ LILLY

EQ 500MG BASE/VIAL

N50504 001

@

EQ 500MG BASE/VIAL

N50504 001

CEFAZOLIN SODIUM

INJECTABLE; INJECTION

ANCEF

AP + SMITHKLINE BEECHAM

EQ 250MG BASE/VIAL

N50461 001

AP

EQ 250MG BASE/VIAL

N50461 001

CEFAZOLIN SODIUM

AP APOTHECON

EQ 500MG BASE/VIAL

N62831 001

AP

EQ 1GM BASE/VIAL

DEC 09, 1988

AP

EQ 10GM BASE/VIAL

N62831 002

AP ELKINS SINN

EQ 250MG BASE/VIAL

DEC 09, 1988

AP

EQ 500MG BASE/VIAL

N62831 003

AP

EQ 1GM BASE/VIAL

SEP 25, 1992

AP

EQ 250MG BASE/VIAL

N62807 001

AP

EQ 500MG BASE/VIAL

JAN 12, 1988

AP

EQ 1GM BASE/VIAL

N62807 002

AP

EQ 5GM BASE/VIAL

JAN 12, 1988

AP

EQ 10GM BASE/VIAL

N62807 003

AP

EQ 20GM BASE/VIAL

JAN 12, 1988

@

EQ 250MG BASE/VIAL

N62807 004

@

EQ 500MG BASE/VIAL

JAN 12, 1988

N62807 005

JAN 12, 1988

N62807 006

JAN 12, 1988

N62807 007

JAN 12, 1988

N62807 008

JAN 12, 1988

CEFAZOLIN SODIUM

INJECTABLE; INJECTION
CEFAZOLIN SODIUM
 @ ELKINS SINN

EQ 1GM BASE/VIAL N62807 003
 JAN 12, 1988
 EQ 5GM BASE/VIAL N62807 004
 JAN 12, 1988
 EQ 10GM BASE/VIAL N62807 005
 JAN 12, 1988
 EQ 20GM BASE/VIAL N62807 006
 JAN 12, 1988

KEFZOL

LILLY

> DLT > AP EQ 250MG BASE/VIAL N61773 001
 > ADD > AP + EQ 250MG BASE/VIAL N61773 001
 > DLT > AP EQ 20GM BASE/VIAL N61773 005
 > DLT > SEP 08, 1987
 > ADD > AP + EQ 20GM BASE/VIAL N61773 005
 > ADD > SEP 08, 1987

ZOLICEF

AP APOTHECON

EQ 500MG BASE/VIAL N62831 001
 DEC 09, 1988
 EQ 1GM BASE/VIAL N62831 002
 DEC 09, 1988
 EQ 10GM BASE/VIAL N62831 003
 SEP 25, 1992

CEFOPERAZONE SODIUM

INJECTABLE; INJECTION
 CEFOBID
 PFIZER

EQ 1GM BASE/VIAL N63333 001
 MAR 31, 1995
 EQ 2GM BASE/VIAL N63333 002
 MAR 31, 1995

CEFORANIDE

INJECTABLE; INJECTION
 PRECEF
 APOTHECON

500MG/VIAL N62579 001
 NOV 26, 1984
 1GM/VIAL N62579 002
 NOV 26, 1984
 2GM/VIAL N62579 003
 NOV 26, 1984

CEFORANIDE

INJECTABLE; INJECTION
 PRECEF
 APOTHECON

10GM/VIAL N62579 004
 NOV 26, 1984
 20GM/VIAL N62579 005
 NOV 26, 1984
 500MG/VIAL N62579 001
 NOV 26, 1984
 1GM/VIAL N62579 002
 NOV 26, 1984
 2GM/VIAL N62579 003
 NOV 26, 1984
 10GM/VIAL N62579 004
 NOV 26, 1984
 20GM/VIAL N62579 005
 NOV 26, 1984

BRISTOLCEFOXITIN SODIUM

INJECTABLE; INJECTION
 MEFOXIN IN SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER

* MERCK SHARP DOHME EQ 20MG BASE/ML N50581 002
 SEP 20, 1984
 * EQ 40MG BASE/ML N50581 001
 SEP 20, 1984
 @ EQ 20MG BASE/ML N50581 002
 SEP 20, 1984
 @ EQ 40MG BASE/ML N50581 001
 SEP 20, 1984

CEFTAZIDIME (ARGININE FORMULATION)

INJECTABLE; INJECTION

PENTACEF

AP SMITHKLINE BEECHAM 1GM/VIAL
 AP 2GM/VIAL

N63322 001
 NOV 07, 1995
 N63322 002
 NOV 07, 1995

CEFTIBUTEN DIHYDRATE

CAPSULE; ORAL

CEDAX

+ SCHERING PLOUGH EQ 400MG BASE

N50685 002
 DEC 20, 1995

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

> ADD > CEFTIBUTEN DIHYDRATE> ADD > POWDER FOR RECONSTITUTION; ORAL> ADD > CEDAX> ADD > + SCHERING PLOUGH EQ 90MG BASE/5ML N50686 001> ADD > DEC 20, 1995> ADD > + EQ 180MG BASE/5ML N50686 002> ADD > DEC 20, 1995CEFTRIAZONE SODIUM

INJECTABLE; INJECTION

ROCEPHINAP * ROCHE EQ 250MG BASE/VIAL N50585 001

DEC 21, 1984

AP * EQ 500MG BASE/VIAL N50585 002

DEC 21, 1984

AP * EQ 1GM BASE/VIAL N50585 003

DEC 21, 1984

+ EQ 250MG BASE/VIAL N50585 001

DEC 21, 1984

+ EQ 500MG BASE/VIAL N50585 002

DEC 21, 1984

+ EQ 1GM BASE/VIAL N50585 003

DEC 21, 1984

CEFUROXIME SODIUM

INJECTABLE; INJECTION

KEFUROXAP * LILLY EQ 7.5GM BASE/VIAL N62591 003

DEC 17, 1987

AP EQ 7.5GM BASE/VIAL N62591 003

DEC 17, 1987

AP ZINACEF EQ 7.5GM BASE/VIAL N50558 004

OCT 23, 1986

AP GLAXO EQ 7.5GM BASE/VIAL N50558 004

OCT 23, 1986

AP + GLAXO WELLCOME EQ 7.5GM BASE/VIAL N50558 004

OCT 23, 1986

CEPHALEXIN

POWDER FOR RECONSTITUTION; ORAL

CEPHALEXINAB APOTHECON EQ 125MG BASE/5ML N62986 001

APR 18, 1991

CEPHALEXIN

POWDER FOR RECONSTITUTION; ORAL

CEPHALEXINAB APOTHECON EQ 250MG BASE/5ML N62987 001

JUL 25, 1989

AB SQUIBB MARK EQ 125MG BASE/5ML N62986 001

APR 18, 1991

AB EQ 250MG BASE/5ML N62987 001

JUL 25, 1989

CEPHRADINE

CAPSULE; ORAL

VELOSEFAB + APOTHECON 250MG N61764 001AB + 500MG N61764 002AB + ERSANA 250MG N61764 001AB + 500MG N61764 002

INJECTABLE; INJECTION

VELOSEF> DLT > + APOTHECON 250MG/VIAL N61976 001> DLT > + 500MG/VIAL N61976 002> DLT > + 1GM/VIAL N61976 004> DLT > + 2GM/VIAL N61976 003> DLT > + 4GM/VIAL N61976 005> ADD > @ 250MG/VIAL N61976 001> ADD > @ 500MG/VIAL N61976 002> ADD > @ 1GM/VIAL N61976 004> ADD > @ 2GM/VIAL N61976 003> ADD > @ 4GM/VIAL N61976 005

POWDER FOR RECONSTITUTION; ORAL

VELOSEF '125'AB + APOTHECON 125MG/5ML N61763 001AB + ERSANA 125MG/5ML N61763 001VELOSEF '250'AB + APOTHECON 250MG/5ML N61763 002AB + ERSANA 250MG/5ML N61763 002> ADD > CETIRIZINE HYDROCHLORIDE> ADD > TABLET; ORAL> ADD > ZYRTEC> ADD > PFIZER 5MG N19835 001> ADD > DEC 08, 1995

> ADD > CETIRIZINE HYDROCHLORIDE

> ADD > TABLET; ORAL
 > ADD > ZYRTEC
 > ADD > + PFIZER 10MG N19835 002
 > ADD > DEC 08, 1995

CHLORAMPHENICOL

> DLT > CAPSULE; ORAL
 > ADD > AMPHICOL
 > ADD > @ FERRANTE 100MG N60058 001
 > ADD > 250MG N60058 002
 > DLT > AB MK LABS 100MG N60058 001
 > DLT > AB 250MG N60058 002
 > DLT > AB * PARKE DAVIS 100MG N60591 003
 > ADD > + 100MG N60591 003
 > ADD > MYCHEL
 > ADD > AB ARMENPHARM 250MG N60851 001
 > ADD > AB RACHELLE 250MG N60851 001

SOLUTION/DROPS; OPHTHALMIC

> ADD > OPTOMYCIN
 > ADD > AT OPTOPICS 0.5% N62171 001
 > ADD > @ MAR 31, 1982
 > ADD > 0.5% N62171 001
 > ADD > @ MAR 31, 1982

CHLORAMPHENICOL SODIUM SUCCINATE

> ADD > INJECTABLE; INJECTION
 > ADD > CHLORAMPHENICOL
 > ADD > AP ELKINS SINN EQ 1GM BASE/VIAL N62406 001
 > ADD > @ NOV 09, 1982
 > ADD > EQ 1GM BASE/VIAL N62406 001
 > ADD > @ NOV 09, 1982

CHLORDIAZEPOXIDE; ESTROGENS, ESTERIFIED

> ADD > TABLET; ORAL
 > ADD > MENRIUM 5-2
 > ADD > @ ROCHE 10MG; 0.4MG N14740 006
 > ADD > 10MG; 0.4MG N14740 006
 > ADD > @ MENRIUM 5-2
 > ADD > @ ROCHE 5MG; 0.2MG N14740 002

CHLORDIAZEPOXIDE; ESTROGENS, ESTERIFIED

> ADD > TABLET; ORAL
 > ADD > MENRIUM 5-2
 > ADD > @ ROCHE 5MG; 0.2MG N14740 002
 > ADD > MENRIUM 5-4
 > ADD > * ROCHE 5MG; 0.4MG N14740 004
 > ADD > @ 5MG; 0.4MG N14740 004

CHLORHEXIDINE GLUCONATE

> ADD > SOLUTION; DENTAL
 > ADD > CHLORHEXIDINE GLUCONATE
 > ADD > AT BARRE 0.12% N74291 001
 > ADD > DEC 28, 1995
 > ADD > AT LEMMON 0.12% N74522 001
 > ADD > DEC 15, 1995

CHLORPHENIRAMINE MALEATE

> ADD > INJECTABLE; INJECTION
 > ADD > CHLORPHENIRAMINE MALEATE
 > ADD > AP STERIS 10MG/ML N83593 001
 > ADD > @ 100MG/ML N86095 001
 > ADD > @ 10MG/ML N83593 001
 > ADD > @ 100MG/ML N86095 001

CHLORPROMAZINE HYDROCHLORIDE

> ADD > INJECTABLE; INJECTION
 > ADD > CHLORPROMAZINE HCL
 > ADD > AP STERIS 25MG/ML N85591 001
 > ADD > @ 25MG/ML N85591 001

CHLORPROPAMIDE

> ADD > TABLET; ORAL
 > ADD > CHLORPROPAMIDE
 > ADD > AB LEMMON 100MG N88768 001
 > ADD > @ 100MG OCT 11, 1984
 > ADD > N88768 001
 > ADD > @ OCT 11, 1984
 > ADD > GLUCAMIDE
 > ADD > AB LEMMON 250MG N88641 001
 > ADD > @ OCT 11, 1984

CHLORPROPAMIDE

TABLET; ORAL
GLUCAMIDE
 @ LEMMON

250MG

N88641 001
 OCT 11, 1984

CHLORPROTHIXENE

CONCENTRATE; ORAL
 TARACTAN
 ROCHE
 @

100MG/5ML
 100MG/5ML

N16149 002
 N16149 002

CHLORTHALIDONE

TABLET; ORAL
CHLORTHALIDONE
AB MUTUAL PHARM

25MG

N89738 001
 SEP 19, 1988

AB

50MG

N89739 001
 SEP 19, 1988

@

25MG

N89738 001
 SEP 19, 1988

@

50MG

N89739 001
 SEP 19, 1988

CHLORZOXAZONE

TABLET; ORAL
CHLORZOXAZONE
AA AMIDE PHARM

500MG

N40113 001
 SEP 29, 1995

CHOLESTYRAMINE

BAR, CHEWABLE; ORAL
 CHOLYBAR
 * PARKE DAVIS

EQ 4GM RESIN/BAR

N71621 001
 MAY 26, 1988

EQ 4GM RESIN/BAR

N71739 001
 MAY 26, 1988

@

EQ 4GM RESIN/BAR

N71621 001
 MAY 26, 1988

CHOLESTYRAMINE

BAR, CHEWABLE; ORAL
 CHOLYBAR
 @ PARKE DAVIS

EQ 4GM RESIN/BAR

N71739 001
 MAY 26, 1988

TABLET; ORAL
 QUESTRAN

* BRISTOL MYERS SQUIBB EQ 1GM RESIN

N73403 001
 APR 28, 1994

@

EQ 1GM RESIN

N73403 001
 APR 28, 1994

CIMETIDINE

TABLET; ORAL
CIMETIDINE

AB BAKER NORTON

200MG

N74424 001
 JUL 28, 1995

AB

300MG

N74424 002
 JUL 28, 1995

AB

400MG

N74424 003
 JUL 28, 1995

AB

800MG

N74424 004
 JUL 28, 1995

AB

GENEVA PHARMS

200MG

N74100 001
 JAN 31, 1995

AB

300MG

N74100 002
 JAN 31, 1995

AB

400MG

N74100 003
 JAN 31, 1995

AB

800MG

N74100 004
 JAN 31, 1995

AB

LEK LJUBLJANA

200MG

N74250 001
 JUN 29, 1995

AB

300MG

N74250 002
 JUN 29, 1995

AB

400MG

N74250 003
 JUN 29, 1995

AB

800MG

N74250 004
 JUN 29, 1995

AB

LEMMON

200MG

N74365 001
 FEB 28, 1995

AB

300MG

N74365 002
 FEB 28, 1995

AB

400MG

N74365 003
 FEB 28, 1995

CIMETIDINE

TABLET; ORAL		
<u>CIMETIDINE</u>		
<u>AB</u>	LEMMON	800MG
		N74365 004
		FEB 28, 1995
<u>AB</u>	MOVA	300MG
		N74340 001
		JUN 23, 1995
<u>AB</u>		400MG
		N74340 002
		JUN 23, 1995
<u>AB</u>		800MG
		N74339 001
		JUN 23, 1995
<u>AB</u>	ZENITH LABS	200MG
		N74401 001
		MAY 30, 1995
<u>AB</u>		300MG
		N74401 002
		MAY 30, 1995
<u>AB</u>		400MG
		N74401 003
		MAY 30, 1995
<u>AB</u>		800MG
		N74402 001
		MAY 30, 1995

CIMETIDINE HYDROCHLORIDE

INJECTABLE; INJECTION		
<u>CIMETIDINE HCL</u>		
<u>AP</u>	ABBOTT	EQ 300MG BASE/2ML
		N74344 001
		JAN 31, 1995
<u>AP</u>		EQ 300MG BASE/2ML
		N74345 001
		JAN 31, 1995
<u>AP</u>		EQ 300MG BASE/2ML
		N74422 001
		JAN 31, 1995

CIPROFLOXACIN

INJECTABLE; INJECTION		
<u>CIPRO IN SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER</u>		
> DLT >	@ BAYER	200MG/100ML
> ADD >		N19858 001
> ADD >		DEC 26, 1990
> DLT >	* MILES	200MG/100ML
> DLT >		N19858 001
		DEC 26, 1990

CISAPRIDE MONOHYDRATE

SUSPENSION; ORAL		
PROPULSID		
+	JANSSEN	EQ 1MG BASE/ML
		N20398 001
		SEP 15, 1995

> ADD > CISATRACURIUM BESYLATE

> ADD >	INJECTABLE; INJECTION	
> ADD >	NIMBEX	
> ADD >	+ GLAXO WELLCOME	EQ 2MG BASE/ML
> ADD >		N20551 001
> ADD >		DEC 15, 1995
> ADD >	NIMBEX PRESERVATIVE FREE	
> ADD >	+ GLAXO WELLCOME	EQ 2MG BASE/ML
> ADD >		N20551 003
> ADD >		DEC 15, 1995
> ADD >	+	EQ 10MG BASE/ML
> ADD >		N20551 002
		DEC 15, 1995

CLAVULANATE POTASSIUM; TICARCILLIN DISODIUM

INJECTABLE; INJECTION		
TIMENTIN		
> DLT >	* SMITHKLINE BEECHAM	EQ 100MG ACID/VIAL;
> DLT >		EQ 3GM BASE/VIAL
> DLT >		N50590 001
> DLT >		APR 01, 1985
> DLT >		EQ 100MG ACID/VIAL;
> DLT >		EQ 3GM BASE/VIAL
> DLT >		N62691 001
> DLT >		DEC 19, 1986
> DLT >	*	EQ 200MG ACID/VIAL;
> DLT >		EQ 3GM BASE/VIAL
> DLT >		N50590 002
> DLT >		APR 01, 1985
> DLT >	*	EQ 1GM ACID/VIAL;
> DLT >		EQ 30GM BASE/VIAL
> DLT >		N50590 003
> DLT >		AUG 18, 1987
> ADD >	+	EQ 100MG BASE/VIAL;
> ADD >		EQ 3GM BASE/VIAL
> ADD >		N50590 001
> ADD >		APR 01, 1985
> ADD >		EQ 100MG BASE/VIAL;
> ADD >		EQ 3GM BASE/VIAL
> ADD >		N62691 001
> ADD >		DEC 19, 1986
> ADD >	+	EQ 200MG BASE/VIAL;
> ADD >		EQ 3GM BASE/VIAL
> ADD >		N50590 002
> ADD >		APR 01, 1985
> ADD >	+	EQ 1GM BASE/VIAL;
> ADD >		EQ 30GM BASE/VIAL
> ADD >		N50590 003
> ADD >		AUG 18, 1987
TIMENTIN IN PLASTIC CONTAINER		
> DLT >	* SMITHKLINE BEECHAM	EQ 100MG ACID/100ML;
> DLT >		EQ 3GM BASE/100ML
> DLT >		N50658 001
> DLT >		DEC 15, 1989
> ADD >	+	EQ 100MG BASE/100ML;
> ADD >		EQ 3GM BASE/100ML
> ADD >		N50658 001
> ADD >		DEC 15, 1989

CLINDAMYCIN PHOSPHATESOLUTION; TOPICAL
CLEOCIN
UPJOHN

EQ 1% BASE

N50537 002
FEB 22, 1994CLINDAMYCIN PHOSPHATE

AT CLAY PARK

EQ 1% BASE

N64050 001
NOV 30, 1995SWAB; TOPICAL
CLEOCIN
UPJOHN

EQ 1% BASE

N50537 002
FEB 22, 1994CLOBETASOL PROPIONATECREAM; TOPICAL
TEMOVATE E

BX + GLAXO WELLCOME

0.05%

N20340 001
JUN 17, 1994

TEMOVATE EMOLLIENT

BX + GLAXO

0.05%

N20340 001
JUN 17, 1994

OINTMENT; TOPICAL

EMBELINE

AB DPT

0.05%

N74221 001
MAR 31, 1995

SOLUTION; TOPICAL

CLOBETASOL PROPIONATE

> ADD >

AT NMC

0.05%

N74331 001
DEC 15, 1995

> ADD >

EMBELINE

> ADD >

AT DPT

0.05%

N74222 001
DEC 06, 1995

> ADD >

TEMOVATE

> ADD >

AT + GLAXO WELLCOME

0.05%

N19966 001
FEB 22, 1990

> ADD >

CLOFIBRATECAPSULE; ORAL
CLOFIBRATE

AB GENEVA PHARMS

500MG

N72191 001
MAY 02, 1988CLOFIBRATE

CAPSULE; ORAL

CLOFIBRATE

@ GENEVA PHARMS

500MG

N72191 001
MAY 02, 1988CLOMIPRAMINE HYDROCHLORIDE

CAPSULE; ORAL

ANAFRANIL

BASEL PHARMS

50MG

N19906 002
DEC 29, 1989

+

75MG

N19906 003
DEC 29, 1989

+

50MG

N19906 002
DEC 29, 1989

75MG

N19906 003
DEC 29, 1989CLOTRIMAZOLE

SOLUTION; TOPICAL

CLOTRIMAZOLE

AT LEMMON

1%

N73306 001
FEB 28, 1995CLOXACILLIN SODIUM

CAPSULE; ORAL

CLOXACILLIN SODIUM

AB + APOTHECON

EQ 250MG BASE

N61452 001

AB +

EQ 500MG BASE

N61452 002

TEGOPEN

AB + APOTHECON

EQ 250MG BASE

N61452 001

AB +

EQ 500MG BASE

N61452 002

POWDER FOR RECONSTITUTION; ORAL

TEGOPEN

AA @ APOTHECON

EQ 125MG BASE/5ML

N50192 001

BRISTOL

EQ 125MG BASE/5ML

N50192 001

CLOZAPINE

TABLET; ORAL
CLOZARIL
SANDOZ

25MG	N19758 001
	SEP 26, 1989
+	N19758 002
	SEP 26, 1989
+	N19758 001
	SEP 26, 1989
	N19758 002
100MG	SEP 26, 1989

COLESTIPOL HYDROCHLORIDE

GRANULE; ORAL
COLESTID
UPJOHN

5GM/SCOOPFUL	N17563 003
	SEP 22, 1995
+	N17563 004
5GM/PACKET	SEP 22, 1995

COLISTIN SULFATE

SUSPENSION; ORAL
COLY-MYCIN S
PARKE DAVIS

@	EQ 25MG BASE/5ML	N50355 001
	EQ 25MG BASE/5ML	N50355 001

CORTICOTROPIN

INJECTABLE; INJECTION

AP	ACTH	
	PARKE DAVIS	40 UNITS/VIAL
	@	40 UNITS/VIAL
		N08317 004
		N08317 004

CROMOLYN SODIUM

SOLUTION/DROPS; OPHTHALMIC

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

AT	OPTICROM	
	+ FISONS	4%
		N18155 001
		OCT 03, 1984

CUPRIC SULFATE

INJECTABLE; INJECTION
CUPRIC SULFATE

+	FUJISAWA	EQ 0.4MG COPPER/ML	N19350 001
	@	EQ 0.4MG COPPER/ML	MAY 05, 1987
			N19350 001
			MAY 05, 1987

CYANOCOBALAMIN

INJECTABLE; INJECTION
CYANOCOBALAMIN

AP	AKORN	1MG/ML	N87969 001
	@	1MG/ML	NOV 10, 1983
	@ WARNER CHILCOTT	1MG/ML	N87969 001
	RUBRAMIN PC		NOV 10, 1983
AP	* SQUIBB	0.1MG/ML	N07085 002
	@	0.1MG/ML	N06799 002
	SYTOBEX		N06799 002
AP	PARKE DAVIS	1MG/ML	N07085 002

CYCLACILLIN

TABLET; ORAL
CYCLACILLIN

AB	ETOCRAFT	250MG	N62895 001
			AUG 04, 1988
AB		500MG	N62895 002
			AUG 04, 1988
	+	250MG	N62895 001
	+	500MG	AUG 04, 1988
			N62895 002
			AUG 04, 1988

AB	CYCLAPEN-W	250MG	N50509 001
AB	+ WYETH AYERST	500MG	N50509 002
	@	250MG	N50509 001
	@	500MG	N50509 002

CYCLOBENZAPRINE HYDROCHLORIDE

TABLET; ORAL

CYCLOBENZAPRINE HCL

AB BARR 10MG
 AB SIDMAK LABS NJ 10MG

N73541 001 > DLT >
 MAY 23, 1995 > DLT >
 N74421 001 > ADD >
 SEP 29, 1995 > ADD >

CYCLOPHOSPHAMIDE

TABLET; ORAL

CYTOXAN

BRISTOL 25MG
 * BRISTOL 50MG
 BRISTOL MYERS SQUIBB 25MG
 + 50MG

N12141 002
 N12141 001
 N12141 002
 N12141 001

CYCLOPHOSPHAMIDE

INJECTABLE; INJECTION

CYTOXAN

> DLT > AP + BRISTOL 100MG/VIAL
 > DLT > AP + 200MG/VIAL
 > DLT > AP + 500MG/VIAL
 > DLT > AP + 1GM/VIAL
 > DLT > AP +
 > DLT > AP + 2GM/VIAL
 > DLT > AP +
 > ADD > AP + BRISTOL MYERS SQUIBB 100MG/VIAL
 > ADD > AP + 200MG/VIAL
 > ADD > AP + 500MG/VIAL
 > ADD > AP + 1GM/VIAL
 > ADD > AP +
 > ADD > AP + 2GM/VIAL
 > ADD > AP +

N12142 001
 N12142 002
 N12142 003
 N12142 004
 AUG 30, 1982
 N12142 005
 AUG 30, 1982
 N12142 001
 N12142 002
 N12142 003
 N12142 004
 AUG 30, 1982
 N12142 005
 AUG 30, 1982

LYOPHILIZED CYTOXAN

> DLT > AP + BRISTOL 100MG/VIAL
 > DLT > AP +
 > DLT > AP + 200MG/VIAL
 > DLT > AP + 500MG/VIAL
 > DLT > AP +
 > DLT > AP + 1GM/VIAL
 > DLT > AP + 2GM/VIAL
 > DLT > AP +
 > ADD > AP + BRISTOL MYERS SQUIBB 100MG/VIAL
 > ADD > AP + 200MG/VIAL
 > ADD > AP + 500MG/VIAL
 > ADD > AP + 1GM/VIAL
 > ADD > AP + 2GM/VIAL
 > ADD > AP +

N12142 006
 DEC 05, 1985
 N12142 007
 DEC 10, 1985
 N12142 008
 JAN 04, 1984
 N12142 010
 SEP 24, 1985
 N12142 009
 DEC 10, 1984
 N12142 006
 DEC 05, 1985
 N12142 007
 DEC 10, 1985
 N12142 008
 JAN 04, 1984
 N12142 010
 SEP 24, 1985
 N12142 009
 DEC 10, 1984

CYCLOSPORINE

CAPSULE; ORAL

NEORAL

BP SANDOZ 25MG
 BP 50MG
 BP + 100MG
 BP + 25MG
 * 50MG
 * 100MG
 BP SANDIMMUNE SANDOZ 25MG
 BP 50MG
 BP + 100MG
 BP 25MG
 BP 50MG
 * 100MG

N50715 001
 JUL 14, 1995
 N50715 003
 JUL 14, 1995
 N50715 002
 JUL 14, 1995
 N50715 001
 JUL 14, 1995
 N50715 003
 JUL 14, 1995
 N50715 002
 JUL 14, 1995

SOLUTION; ORAL

NEORAL

BP + SANDOZ 100MG/ML
 * 100MG/ML
 BP SANDIMMUNE SANDOZ 100MG/ML

N50625 001
 MAR 02, 1990
 N50625 003
 NOV 23, 1992
 N50625 002
 MAR 02, 1990
 N50625 001
 MAR 02, 1990
 N50625 003
 NOV 23, 1992
 N50625 002
 MAR 02, 1990

N50716 001
 JUL 14, 1995
 N50716 001
 JUL 14, 1995

N50574 001
 NOV 14, 1983

CYCLOSPORINESOLUTION; ORAL
SANDIMMUNE

* SANDOZ 100MG/ML

N50574 001
NOV 14, 1993CYPROHEPTADINE HYDROCHLORIDETABLET; ORAL
CYPROHEPTADINE HCL

AA ASCOT 4MG

N87685 001
OCT 25, 1982
N87685 001
OCT 25, 1982

@ 4MG

DAUNORUBICIN HYDROCHLORIDEINJECTABLE; INJECTION
DAUNORUBICIN HCL

AP CETUS BEN VENUE EQ 20MG BASE/VIAL

N64103 001
FEB 03, 1995DESMOPRESSIN ACETATEINJECTABLE; INJECTION
DDAVP

+ RHONE POULENC 0.015MG/ML

N18938 002
APR 25, 1995SPRAY, METERED; NASAL
STIMATE

+ ARMOUR PHARM 0.15MG/INH

N20355 001
MAR 07, 1994

* RHONE POULENC RORER 0.15MG/INH

N20355 001
MAR 07, 1994TABLET; ORAL
DDAVP

RHONE POULENC RORER 0.1MG

N19955 001
SEP 06, 1995

+ 0.2MG

N19955 002
SEP 06, 1995DESOGESTREL; ETHINYL ESTRADIOL

TABLET; ORAL-21

DESOREN

AB * ORGANOX 0.15MG;0.03MG

N20071 001

@

0.15MG;0.03MG

DEC 10, 1992
N20071 001
DEC 10, 1992

ORTHO-CEPT

AB JOHNSON RW 0.15MG;0.03MG

N20301 001

0.15MG;0.03MG

DEC 14, 1992
N20301 001
DEC 14, 1992DEXAMETHASONEAEROSOL; TOPICAL
DECASPRAY

* MERCK SHARP DOHME 0.4%

N12731 002

+ 0.04%

N12731 002

TABLET; ORAL

HEXADROL

BP ORGANOX 0.5MG

N12675 004

BP 0.75MG

N12675 007

BP 1.5MG

N12675 009

@ 0.5MG

N12675 004

@ 0.75MG

N12675 007

@ 1.5MG

N12675 009

DEXAMETHASONE; NEOMYCIN SULFATE; POLYMYXIN B SULFATE

SUSPENSION/DROPS; OPHTHALMIC

DEXASPORIN

AT BAUSCH AND LOMB 0.1%;EQ 3.5MG BASE/ML;
10,000 UNITS/MLN64135 001
SEP 13, 1995DEXAMETHASONE SODIUM PHOSPHATE

INJECTABLE; INJECTION

DEXAMETHASONE

AP FUJISAWA EQ 4MG PHOSPHATE/ML

N88448 001

@

EQ 4MG PHOSPHATE/ML

JAN 25, 1984
N88448 001
JAN 25, 1984

DEXAMETHASONE SODIUM PHOSPHATE

INJECTABLE; INJECTION

DEXAMETHASONE SODIUM PHOSPHATE

<u>AP</u>	<u>AKORN</u>	<u>EQ 4MG PHOSPHATE/ML</u>	<u>N84493 001</u>
	@	EQ 4MG PHOSPHATE/ML	N84493 001

DEXAMETHASONE SODIUM PHOSPHATE; NEOMYCIN SULFATE

SOLUTION/DROPS; OPHTHALMIC

NEOMYCIN SULFATE AND DEXAMETHASONE SODIUM PHOSPHATE

<u>AT</u>	<u>BAUSCH AND LOMB</u>	<u>EQ 0.1% PHOSPHATE;</u>	
		<u>EQ 3.5MG BASE/ML</u>	<u>N64055 001</u>
			OCT 30, 1995

DEXRAZOXANE HYDROCHLORIDE

INJECTABLE; INJECTION

ZINECARD

+	PHARMACIA	EQ 250MG BASE/VIAL	N20212 001
			MAY 26, 1995
+		EQ 500MG BASE/VIAL	N20212 002
			MAY 26, 1995

DEXTROSE

INJECTABLE; INJECTION

DEXTROSE 2.5% IN PLASTIC CONTAINER

<u>MCGAW</u>	<u>2.5GM/100ML</u>	<u>N19626 001</u>
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@	2.5GM/100ML	N19626 001
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		FEB 02, 1988
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		N19626 001
--	--	------------

		FEB 02, 1988
--	--	--------------

<u>AP</u>	<u>DHL</u>	<u>5GM/100ML</u>	<u>N19971 001</u>
			SEP 28, 1995

DEXTROSE 7.7% IN PLASTIC CONTAINER

<u>MCGAW</u>	<u>7.7GM/100ML</u>	<u>N19626 003</u>
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@	7.7GM/100ML	N19626 003
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		FEB 02, 1988
--	--	--------------

		N19626 003
--	--	------------

		FEB 02, 1988
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DIATRIZOATE MEGLUMINE

INJECTABLE; INJECTION

ANGIOVIST 282

> DLT >	<u>AP</u>	<u>BERLEX</u>	<u>60%</u>	<u>N87726 001</u>
> DLT >				SEP 23, 1982
> DLT >		@	60%	N87726 001
> ADD >				SEP 23, 1982
> ADD >				
> DLT >	<u>AP</u>	<u>UROVIST MEGLUMINE DIV/CT</u>	<u>30%</u>	<u>N87739 001</u>
> DLT >		BERLEX		SEP 23, 1982
> ADD >	@		30%	N87739 001
> ADD >				SEP 23, 1982

SOLUTION; URETERAL

RENO-30

> ADD >		BRACCO	30%	N10040 021
> DLT >	<u>AT</u>	<u>BRACCO DXS</u>	<u>30%</u>	<u>N10040 021</u>
> DLT >				
> DLT >	<u>AT</u>	<u>UROVIST CYSTO</u>	<u>30%</u>	<u>N87729 001</u>
> DLT >		BERLEX		SEP 23, 1982
> ADD >	@		30%	N87729 001
> ADD >				SEP 23, 1982
> DLT >	<u>AT</u>	<u>UROVIST CYSTO PEDIATRIC</u>	<u>30%</u>	<u>N87731 001</u>
> DLT >		BERLEX		SEP 23, 1982
> ADD >	@		30%	N87731 001
> ADD >				SEP 23, 1982

DIATRIZOATE MEGLUMINE; DIATRIZOATE SODIUM

INJECTABLE; INJECTION

ANGIOVIST 292

> DLT >	<u>AP</u>	<u>BERLEX</u>	<u>52%;8%</u>	<u>N87724 001</u>
> DLT >				SEP 23, 1982
> DLT >		@	52%;8%	N87724 001
> ADD >				SEP 23, 1982
> ADD >				
> DLT >	<u>AP</u>	<u>ANGIOVIST 370</u>	<u>66%;10%</u>	<u>N87723 001</u>
> DLT >		BERLEX		SEP 23, 1982
> ADD >	@		66%;10%	N87723 001
> ADD >				SEP 23, 1982
> ADD >				
> ADD >		<u>RENOGRAFIN-60</u>		
> ADD >		BRACCO	52%;8%	N10040 006
> DLT >	<u>AP</u>	<u>BRACCO DXS</u>	<u>52%;8%</u>	<u>N10040 006</u>

SOLUTION; ORAL, RECTAL

GASTROVIST

> DLT >	<u>AA</u>	<u>BERLEX</u>	<u>56%;10%</u>	<u>N87728 001</u>
> DLT >				SEP 23, 1982
> DLT >				

DIATRIZOATE MEGLUMINE; DIATRIZOATE SODIUM

SOLUTION; ORAL, RECTAL

> <u>DLT</u> >	<u>GASTROVIST</u>		
> <u>ADD</u> >	@ BERLEX	66%;10%	N87728 001
> <u>ADD</u> >			SEP 23, 1982

DIATRIZOATE SODIUM

INJECTABLE; INJECTION

> <u>DLT</u> >	<u>AP</u>	<u>HYPAAQUE</u>	<u>50%</u>	<u>N09561 001</u>
> <u>ADD</u> >		NYCONED	50%	N09561 001
> <u>DLT</u> >		<u>UROVIST SODIUM 300</u>		
> <u>DLT</u> >	<u>AP</u>	BERLEX	<u>50%</u>	<u>N87725 001</u>
> <u>DLT</u> >				SEP 23, 1982
> <u>ADD</u> >	@		50%	N87725 001
> <u>ADD</u> >				SEP 23, 1982

DIAZEPAM

INJECTABLE; INJECTION

<u>AP</u>	<u>DIAZEPAM</u>		
	FUJISAWA	<u>5MG/ML</u>	<u>N70662 001</u>
	@	5MG/ML	JUN 25, 1986
			N70662 001
			JUN 25, 1986

DICLOFENAC POTASSIUM

TABLET; ORAL

CATAFLAM	
GEIGY	<u>25MG</u>
@	25MG

<u>N20142 001</u>
NOV 24, 1993
<u>N20142 001</u>
NOV 24, 1993

DICLOFENAC SODIUM

TABLET, DELAYED RELEASE; ORAL

<u>AB</u>	<u>DICLOFENAC SODIUM</u>	
	GENEVA PHARMS	<u>25MG</u>
<u>AB</u>		<u>50MG</u>

<u>N74376 001</u>
SEP 28, 1995
<u>N74376 002</u>
SEP 28, 1995

DICLOFENAC SODIUM

TABLET, DELAYED RELEASE; ORAL

<u>AB</u>	<u>DICLOFENAC SODIUM</u>	
	GENEVA PHARMS	<u>75MG</u>
<u>AB</u>	ROXANE	<u>25MG</u>
<u>AB</u>		<u>50MG</u>
<u>AB</u>		<u>75MG</u>
	<u>VOLTAREN</u>	
<u>AB</u>	+ GEIGY	<u>25MG</u>
<u>AB</u>	+	<u>50MG</u>
<u>AB</u>	+	<u>75MG</u>

<u>N74394 001</u>
NOV 30, 1995
<u>N74391 001</u>
JUN 29, 1995
<u>N74391 002</u>
JUN 29, 1995
<u>N74391 003</u>
JUN 29, 1995
<u>N19201 001</u>
JUL 28, 1988
<u>N19201 002</u>
JUL 28, 1988
<u>N19201 003</u>
JUL 28, 1988

DICLOXACILLIN SODIUM

CAPSULE; ORAL

<u>AB</u>	<u>DICLOXACILLIN SODIUM</u>	
	APOTHECON	<u>EQ 250MG BASE</u>
<u>AB</u>		<u>EQ 500MG BASE</u>
		<u>EQ 125MG BASE</u>
	<u>DYNAPEN</u>	
<u>AB</u>	APOTHECON	<u>EQ 250MG BASE</u>
<u>AB</u>		<u>EQ 500MG BASE</u>
		<u>EQ 125MG BASE</u>

<u>N61454 001</u>
<u>N61454 003</u>
<u>N61454 002</u>
<u>N61454 001</u>
<u>N61454 003</u>
<u>N61454 002</u>

POWDER FOR RECONSTITUTION; ORAL

<u>AB</u>	<u>DICLOXACILLIN SODIUM</u>	
	APOTHECON	<u>EQ 62.5MG BASE/5ML</u>
<u>AB</u>	<u>DYNAPEN</u>	
	APOTHECON	<u>EQ 62.5MG BASE/5ML</u>
	@	<u>EQ 62.5MG BASE/5ML</u>
	@ BRISTOL	<u>EQ 62.5MG BASE/5ML</u>

<u>N61455 001</u>
<u>N61455 001</u>
<u>N50337 002</u>
<u>N50337 002</u>

DIDANOSINE

POWDER FOR RECONSTITUTION; ORAL

VIDEX	
BRISTOL MYERS SQUIBB	<u>250MG/PACKET</u>

<u>N20155 005</u>
OCT 09, 1991

DIDANOSINEPOWDER FOR RECONSTITUTION; ORAL
VIDEX

* BRISTOL MYERS SQUIBB	375MG/PACKET	N20155 006
		OCT 09, 1991
+	250MG/PACKET	N20155 005
		OCT 09, 1991
@	375MG/PACKET	N20155 006
		OCT 09, 1991

DIENESTROLCREAM; VAGINAL
DIENESTROL

AT	* JOHNSON RW	0.01%	N06110 005
	+	0.01%	N06110 005
	DV		
	@ HOECHST MARION RSSL	0.01%	N83518 001
AT	MERRELL DOW	0.01%	N83518 001

DIFLORASONE DIACETATE

CREAM; TOPICAL

DIFLORASONE DIACETATE
@ UPJOHN

		0.05%	N19259 001
			AUG 28, 1985
BX	FLORONE E	0.05%	N19259 001
	+ UPJOHN		AUG 28, 1985
BX	PSORCON	0.05%	N20205 001
	DERMIK		NOV 20, 1992
BX	+	0.05%	N20205 001
			NOV 20, 1992

DILTIAZEM HYDROCHLORIDECAPSULE, EXTENDED RELEASE; ORAL
CARDIZEM CD

BC	+ CARDERM	300MG	N20062 004
			DEC 27, 1991
BC	DILACOR XR	120MG	N20092 001
	RHONE POULENC RORER		MAY 29, 1992

DILTIAZEM HYDROCHLORIDECAPSULE, EXTENDED RELEASE; ORAL
DILACOR XR

BC	+ RHONE POULENC RORER	120MG	N20092 001
			MAY 29, 1992
BC		180MG	N20092 002
			MAY 29, 1992
BC	+	180MG	N20092 002
			MAY 29, 1992
BC		240MG	N20092 003
			MAY 29, 1992
BC	+	240MG	N20092 003
			MAY 29, 1992
	TIAZAC		
BC	BIOVAIL	120MG	N20401 001
			SEP 11, 1995
BC		180MG	N20401 002
			SEP 11, 1995
BC		240MG	N20401 003
			SEP 11, 1995
BC		300MG	N20401 004
			SEP 11, 1995
	+	360MG	N20401 005
			SEP 11, 1995

INJECTABLE; INJECTION
CARDIZEM

+	HOECHST MARION RSSL	25MG/VIAL	N20027 003
			AUG 18, 1995

TABLET; ORAL

DILTIAZEM HCL
LEMMON

AB		30MG	N74185 001
			MAY 31, 1995
AB		60MG	N74185 002
			MAY 31, 1995
AB		90MG	N74185 003
			MAY 31, 1995
AB		120MG	N74185 004
			MAY 31, 1995
AB	ZENITH LABS	30MG	N74168 001
			MAR 03, 1995
AB		60MG	N74168 002
			MAR 03, 1995
AB		90MG	N74168 003
			MAR 03, 1995
AB		120MG	N74168 004
			MAR 03, 1995

DIMENHYDRINATE

INJECTABLE; INJECTION
DIMENHYDRINATE
 AP STERIS 50MG/ML N83531 001
 @ 50MG/ML N83531 001

DINOPROSTONE

INSERT, EXTENDED RELEASE; VAGINAL
 CERVIDIL
 + CONTROLLED THERAP 10MG N20411 001
 MAR 30, 1995

DIPHENHYDRAMINE HYDROCHLORIDE

CAPSULE; ORAL
DIPHENHYDRAMINE HCL
 AA WEST WARD PHARM 50MG N83567 001
 @ 50MG N83567 001

DIPIVEFRIN HYDROCHLORIDE

SOLUTION/DROPS; OPHTHALMIC
AKPRO
 AT AKORN 0.1% N74382 001
 SEP 29, 1995
DIPIVEFRIN HCL
 AT BAUSCH AND LOMB 0.1% N74188 001
 MAY 19, 1995

DIRITHROMYCIN

TABLET, DELAYED RELEASE; ORAL
 DYNABAC
 + LILLY 250MG N50678 001
 JUN 19, 1995

DIVALPROEX SODIUM

CAPSULE, DELAYED REL PELLETS; ORAL
 DEPAKOTE
 * ABBOTT EQ 125MG BASE N19680 001
 SEP 12, 1989

DIVALPROEX SODIUM

CAPSULE, DELAYED REL PELLETS; ORAL
 DEPAKOTE
 + ABBOTT EQ 125MG VALPROIC ACID N19680 001
 SEP 12, 1989

TABLET, DELAYED RELEASE; ORAL
 DEPAKOTE
 ABBOTT EQ 125MG BASE N18723 003
 OCT 26, 1984
 EQ 250MG BASE N18723 001
 MAR 10, 1983
 EQ 500MG BASE N18723 002
 MAR 10, 1983
 EQ 125MG VALPROIC ACID N18723 003
 OCT 26, 1984
 EQ 250MG VALPROIC ACID N18723 001
 MAR 10, 1983
 EQ 500MG VALPROIC ACID N18723 002
 MAR 10, 1983

DOBUTAMINE HYDROCHLORIDE

INJECTABLE; INJECTION
DOBUTAMINE HCL
 AP ASTRA EQ 12.5MG BASE/ML N74098 001
 FEB 21, 1995
 AP SANOFI WINTHROP EQ 12.5MG BASE/ML N74292 001
 FEB 16, 1995

DOPAMINE HYDROCHLORIDE

INJECTABLE; INJECTION
DOPAMINE HCL
 AP ABBOTT 40MG/ML N70656 001
 JAN 24, 1989
 AP 80MG/ML N70657 001
 JAN 24, 1989
 @ 40MG/ML N70656 001
 JAN 24, 1989
 @ 80MG/ML N70657 001
 JAN 24, 1989
DOPAMINE HCL IN DEXTROSE 5%
 AP + ABBOTT 1.6MG/ML N20542 001
 AUG 30, 1995
INTROPIN
 AP * DUPONT MERCK 40MG/ML N17395 001

DOPAMINE HYDROCHLORIDE

INJECTABLE; INJECTION

INTROPIN

AP	+	DUPONT MERCK	80MG/ML	N17395 002
AP	+		160MG/ML	N17395 003
AP	+	FAULDING	40MG/ML	N17395 001
AP	+		80MG/ML	N17395 002
AP	+		160MG/ML	N17395 003

DOXORUBICIN HYDROCHLORIDE

INJECTABLE; INJECTION

> DLT >	DOXIL			
> DLT >	+	SEQUUS PHARM	2MG/ML	N50718 001
> DLT >				NOV 17, 1995

DOXORUBICIN HCL

AP		FUJISAWA	2MG/ML	N63277 001
				OCT 26, 1995
AP		GENSIA	2MG/ML	N64140 001
				JUL 28, 1995
AP			200MG/100ML	N64140 002
				JUL 28, 1995
AP		PHARMACHEMIE (NL)	2MG/ML	N63336 001
				FEB 28, 1995
AP			200MG/100ML	N63336 004
				FEB 28, 1995

> ADD >	INJECTABLE, LIPOSOMAL; INJECTION			
> ADD >	DOXIL			
> ADD >	+	SEQUUS PHARM	2MG/ML	N50718 001
> ADD >				NOV 17, 1995

DOXYCYCLINE

CAPSULE; ORAL

DOXYCYCLINE MONOHYDRATE

+	VINTAGE PHARMS	EQ 100MG BASE	N50641 001
			DEC 29, 1989

MONODOX

OCLASSEN

		EQ 50MG BASE	N50641 002
			FEB 10, 1992
+		EQ 100MG BASE	N50641 001
			DEC 29, 1989

DOXYCYCLINE HYCLATE

CAPSULE; ORAL

DOXYCYCLINE HYCLATE

AB	HALSEY	EQ 50MG BASE	N62119 002
			MAY 24, 1985
AB		EQ 100MG BASE	N62119 001
			MAY 24, 1985
@		EQ 50MG BASE	N62119 002
			MAY 24, 1985
@		EQ 100MG BASE	N62119 001
			MAY 24, 1985
AB	PVT FORM	EQ 50MG BASE	N62631 001
			JUL 24, 1986
AB		EQ 100MG BASE	N62631 002
			JUL 24, 1986
@		EQ 50MG BASE	N62631 001
			JUL 24, 1986
@		EQ 100MG BASE	N62631 002
			JUL 24, 1986

DROPERIDOL

INJECTABLE; INJECTION

DROPERIDOL

AP	DUPONT MERCK	2.5MG/ML	N71645 001
			APR 07, 1988
AP	FAULDING	2.5MG/ML	N71645 001
			APR 07, 1988
AP	SANOFI WINTHROP	2.5MG/ML	N72272 001
			AUG 31, 1995

EDETATE DISODIUM

INJECTABLE; INJECTION

SODIUM VERSENATE

+	3M	200MG/ML	N10573 001
@		200MG/ML	N10573 001

ENALAPRIL MALEATE; HYDROCHLOROTHIAZIDE

TABLET; ORAL

VASERETIC

MERCK

5MG;12.5MG

N19221 003
JUL 12, 1995

EPINEPHRINE

INJECTABLE; INJECTION

> DLT > EPIPEN
 > DLT > * SURVIVAL TECH 1MG/ML N19410 001
 > DLT > DEC 22, 1987
 > DLT > EPIPEN JR.
 > DLT > * SURVIVAL TECH 0.5MG/ML N19430 002
 > DLT > DEC 22, 1987

> ADD > INJECTABLE; INTRAMUSCULAR
 > ADD > EPIPEN
 > ADD > + SURVIVAL TECH 0.3MG/DELIVERY N19430 001
 > ADD > DEC 22, 1987
 > ADD > EPIPEN JR.
 > ADD > + SURVIVAL TECH 0.15MG/DELIVERY N19430 002
 > ADD > DEC 22, 1987

EPINEPHRINE; LIDOCAINE HYDROCHLORIDE

> ADD > SOLUTION; IONTOPHORESIS
 > ADD > IONTOCAINE
 > ADD > + IOMED 0.01MG/ML; 2% N20530 001
 > ADD > DEC 21, 1995

EPOPROSTENOL SODIUM

INJECTABLE; INJECTION

FLOLAN
 + GLAXO WELLCOME EQ 0.5MG BASE/VIAL N20444 001
 SEP 20, 1995
 + EQ 1.5MG BASE/VIAL N20444 002
 SEP 20, 1995

ERYTHROMYCIN

CAPSULE, DELAYED REL PELLETS; ORAL

ERYC

AB FAULDING 250MG N50536 001
AB 250MG N50536 001
AB * PARKE DAVIS 250MG N62338 001
AB * 250MG N62618 001
SEP 25, 1985

@ 250MG N62338 001
 @ 250MG N62618 001
 SEP 25, 1985

ERYTHROMYCIN

SWAB; TOPICAL

C-SOLVE-2
 > DLT > AT SYOSSET 2% N62751 001
 > DLT > JUL 30, 1993
 > ADD > AT ZENITH GOLDLINE 2% N62751 001
 > ADD > JUL 30, 1993

TABLET, DELAYED RELEASE; ORAL

ROBIMYCIN
AB ROBINS AM 250MG N61633 001
 @ 250MG N61633 001

ERYTHROMYCIN ESTOLATE

DROPS; ORAL

ILOSONE
 * DISTA EQ 100MG BASE/ML N61894 003

SUSPENSION/DROPS; ORAL

ILOSONE
 + DISTA EQ 100MG BASE/ML N61894 003

ERYTHROMYCIN ETHYLSUCCINATE

DROPS; ORAL

PEDIAMYCIN
 + ROSS LABS EQ 100MG BASE/2.5ML N62305 002

SUSPENSION; ORAL

WYAMYCIN E
AB WYETH AYERST EQ 200MG BASE/5ML N62123 002
AB EQ 400MG BASE/5ML N62123 001
 @ EQ 200MG BASE/5ML N62123 002
 @ EQ 400MG BASE/5ML N62123 001

SUSPENSION/DROPS; ORAL

PEDIAMYCIN
 + ROSS LABS EQ 100MG BASE/2.5ML N62305 002

ERYTHROMYCIN STEARATE

TABLET; ORAL

ETHRIL 250
AB SQUIBB EQ 250MG BASE N61605 001

ERYTHROMYCIN STEARATE

TABLET; ORAL		
	<u>ETHRIL 250</u>	
	@ SQUIBB	EQ 250MG BASE
	<u>ETHRIL 500</u>	N61605 001
<u>AB</u>	SQUIBB	<u>EQ 500MG BASE</u>
	@	EQ 500MG BASE
		N61605 002

ESTRADIOL

FILM, EXTENDED RELEASE; TRANSDERMAL		
CLIMARA		
BX	* 3M	0.05MG/24HR
		N20375 001
		DEC 22, 1994
BX	*	0.1MG/24HR
		N20375 002
		DEC 22, 1994
BX	+ BERLEX	0.05MG/24HR
		N20375 001
		DEC 22, 1994
BX	+	0.1MG/24HR
		N20375 002
		DEC 22, 1994
VIVELLE		
BX		0.05MG/24HR
		N20323 002
		OCT 28, 1994
BX		0.1MG/24HR
		N20323 004
		OCT 28, 1994
		0.0375MG/24HR
		N20323 001
		OCT 28, 1994
		0.075MG/24HR
		N20323 003
		OCT 28, 1994
BX	+ CIBA GEIGY	0.05MG/24HR
		N20323 002
		OCT 28, 1994
BX	+	0.1MG/24HR
		N20323 004
		OCT 28, 1994
	+	0.0375MG/24HR
		N20323 001
		OCT 28, 1994
	+	0.075MG/24HR
		N20323 003
		OCT 28, 1994

ESTRADIOL VALERATE

INJECTABLE; INJECTION		
<u>DELESTROGEN</u>		
<u>AO</u>	* SQUIBB	<u>10MG/ML</u>
	+	10MG/ML
	<u>ESTRADIOL VALERATE</u>	N09402 002
		N09402 002
<u>AO</u>	STERIS	<u>10MG/ML</u>
		N83546 001

ESTRADIOL VALERATE

INJECTABLE; INJECTION		
<u>ESTRADIOL VALERATE</u>		
@ STERIS	10MG/ML	N83546 001

ESTROGENS, CONJUGATED; MEDROXYPROGESTERONE ACETATE

TABLET; ORAL-28		
PREMPHASE 14/14		
+ WYETH AYERST	0.625MG, 0.625MG; N/A, 5MG	N20527 002
		NOV 17, 1995
PREMPRO 14/14		
+ WYETH AYERST	0.625MG, 0.625MG; 2.5MG, 2.5MG	N20527 001
		NOV 17, 1995

ESTRONE

INJECTABLE; INJECTION		
ESTROGENIC SUBSTANCE		
BP	WYETH AYERST	2MG/ML
BP	+	2MG/ML
	THELIN	N83488 001
BP	* PARKE DAVIS	2MG/ML
	@	2MG/ML
		N03977 002
		N03977 002

ETHACRYNATE SODIUM

INJECTABLE; INJECTION		
EDECIN		
* MERCK SHARP DOHME	EQ 50MG ACID/VIAL	N16093 001
+	EQ 50MG BASE/VIAL	N16093 001

ETHANOLAMINE OLEATE

INJECTABLE; INJECTION		
ETHAMOLIN		
* REED AND CARNRICK	50MG/ML	N19357 001
		DEC 22, 1988
+	SPKU	50MG/ML
		N19357 001
		DEC 22, 1988

ETHINYL ESTRADIOL; FERROUS FUMARATE; NORETHINDRONE

TABLET; ORAL-28

NORQUEST FE

@ SEARLE

0.035MG;75MG;1MG

N18926 001

JUL 18, 1986

@ SYNTEX

0.035MG;75MG;1MG

N18926 001

JUL 18, 1986

ETHINYL ESTRADIOL; LEVONORGESTREL

TABLET; ORAL-21

LEVORA 0.15/30-21

> ADD >

AB

SEARLE

0.03MG;0.15MG

N73592 001

> ADD >

AB

SYNTEX

0.03MG;0.15MG

N73592 001

> DLT >

AB

SYNTEX

0.03MG;0.15MG

N73592 001

> DLT >

AB

SYNTEX

0.03MG;0.15MG

N73592 001

TABLET; ORAL-28

LEVORA 0.15/30-28

> ADD >

AB

SEARLE

0.03MG;0.15MG

N73594 001

> ADD >

AB

SYNTEX

0.03MG;0.15MG

N73594 001

> DLT >

AB

SYNTEX

0.03MG;0.15MG

N73594 001

> DLT >

AB

SYNTEX

0.03MG;0.15MG

N73594 001

ETHINYL ESTRADIOL; NORETHINDRONE

TABLET; ORAL-21

BREVICON 21-DAY

AB

SEARLE

0.035MG;0.5MG

N17566 001

AB

SYNTEX

0.035MG;0.5MG

N17566 001

NORINYL 1+35 21-DAY

AB

SEARLE

0.035MG;1MG

N17565 001

AB

SYNTEX

0.035MG;1MG

N17565 001

OVCON-35

+ BRISTOL MYERS SQUIBB

0.035MG;0.4MG

N18127 001

* MEAD JOHNSON

0.035MG;0.4MG

N18127 001

OVCON-50

@ BRISTOL MYERS SQUIBB

0.05MG;1MG

N18128 001

* MEAD JOHNSON

0.05MG;1MG

N18128 001

TRI-NORINYL 21-DAY

+ SEARLE

0.035MG;0.035MG;0.5MG;1MG N18977 001

APR 13, 1984

* SYNTEX

0.035MG;0.035MG;0.5MG;1MG N18977 001

APR 13, 1984

TABLET; ORAL-28

BREVICON 28-DAY

AB

SEARLE

0.035MG;0.5MG

N17743 001

ETHINYL ESTRADIOL; NORETHINDRONE

TABLET; ORAL-28

BREVICON 28-DAY

AB

SYNTEX

0.035MG;0.5MG

N17743 001

NORINYL 1+35 28-DAY

AB

SEARLE

0.035MG;1MG

N17565 002

AB

SYNTEX

0.035MG;1MG

N17565 002

OVCON-35

BRISTOL MYERS SQUIBB

0.035MG;0.4MG

N17716 001

MEAD JOHNSON

0.035MG;0.4MG

N17716 001

OVCON-50

BRISTOL MYERS SQUIBB

0.05MG;1MG

N17576 001

MEAD JOHNSON

0.05MG;1MG

N17576 001

TRI-NORINYL 28-DAY

SEARLE

0.035MG;0.035MG;0.5MG;1MG N18977 002

APR 13, 1984

SYNTEX

0.035MG;0.035MG;0.5MG;1MG N18977 002

APR 13, 1984

ETHOPROPAZINE HYDROCHLORIDE

TABLET; ORAL

PARSIDOL

PARKE DAVIS

10MG

N09078 003

50MG

N09078 006

100MG

N09078 008

*

10MG

N09078 003

@

50MG

N09078 006

@

100MG

N09078 008

ETOPOSIDE

INJECTABLE; INJECTION

ETOPOSIDE

AP

ABBOTT

20MG/ML

N74320 001

AP

ABBOTT

20MG/ML

AUG 30, 1995

AP

ABBOTT

20MG/ML

N74351 001

AP

ABBOTT

20MG/ML

AUG 30, 1995

AP

BEN VENUE

20MG/ML

N74290 001

AP

BEN VENUE

20MG/ML

JUL 17, 1995

AP

GENSIA

20MG/ML

N74510 001

AP

GENSIA

20MG/ML

JUN 29, 1995

TOPOSAR

AP

PHARMACIA

20MG/ML

N74166 001

FEB 27, 1995

FAMCICLOVIR

TABLET; ORAL
FAMVIR
SMITHKLINE BEECHAM 125MG N20363 003
> ADD > DEC 11, 1995
> ADD >

FENOFIBRATE

CAPSULE; ORAL
LIPIDIL
+ LABS FOURNIER 100MG N19304 001
@ 100MG DEC 31, 1993
N19304 001
DEC 31, 1993

FENTANYL CITRATE

TROCHE/LOZENGE; ORAL
FENTANYL
ANESTA EQ 0.1MG BASE N20195 007
OCT 30, 1995

FLUDROCORTISONE ACETATE

TABLET; ORAL
FLORINEF
+ APOTHECON 0.1MG N10060 001
+ SQUIBB 0.1MG N10060 001

FLUNISOLIDE

SPRAY, METERED; NASAL
NASALIDE
EX + SYNTEX 0.025MG/INH N18148 001
NASAREL
BX + SYNTEX 0.025MG/INH N20409 001
MAR 08, 1995

FLUOCINOLONE ACETONIDE

CREAM; TOPICAL
FLUOCINOLONE ACETONIDE
AT + HAMILTON PHARMA CA 0.01% N12787 004

FLUOCINOLONE ACETONIDE

CREAM; TOPICAL
FLUOCINOLONE ACETONIDE
AT + HAMILTON PHARMA CA 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
SYNALAR-HP
+ SYNTEX 0.2% N16161 002
SYNEMOL
AT + SYNTEX 0.025% N12787 005

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 + PHARMADERM 0.01% N88048 001
@ 0.01% DEC 16, 1982
N88048 001
DEC 16, 1982
SYNALAR
AT + SYNTEX 0.01% N15296 001

FLUOCINONIDE

CREAM; TOPICAL
FLUOCINONIDE
AB + HAMILTON PHARMA CA 0.05% N16908 002
FLUOCINONIDE EMOLLIENT BASE
AB HAMILTON PHARMA CA 0.05% N16908 003
FLUOCINONIDE EMULSIFIED BASE
AB NMC 0.05% N74204 001
JUN 13, 1995
LIDEX
AB + SYNTEX 0.05% N16908 002
LIDEX-E
AB SYNTEX 0.05% N16908 003

GEL; TOPICAL
FLUOCINONIDE
AB + HAMILTON PHARMA CA 0.05% N17373 001

FLUOCINONIDE

GEL; TOPICAL

LIDEXAB * SYNTEX 0.05%N17373 001

OINTMENT; TOPICAL

FLUOCINONIDEAB + HAMILTON PHARMA CA 0.05%N16909 002LIDEXAB * SYNTEX 0.05%N16909 002

SOLUTION; TOPICAL

FLUOCINONIDEAT FOUGERA 0.05%N72934 001
FEB 27, 1995AT + HAMILTON PHARMA CA 0.05%N18849 001
APR 06, 1984LIDEXAT * SYNTEX 0.05%N18849 001
APR 06, 1984FLUOROURACIL

INJECTABLE; INJECTION

ADRUCIL> DLT > AP PHARMACIA 50MG/MLN81222 001> DLT > JUN 28, 1991> ADD > @ 50MG/MLN81222 001> ADD >

JUN 28, 1991

FLUPHENAZINE HYDROCHLORIDE

CONCENTRATE; ORAL

PERMITILAA SCHERING 5MG/MLN15008 001AA + 5MG/MLN16008 001FLURBIPROFEN

TABLET; ORAL

FLURBIPROFENAB GENEVA PHARMS 50MGN74448 001
JUL 28, 1995AB 100MGN74448 002
JUL 28, 1995FLURBIPROFEN

TABLET; ORAL

FLURBIPROFENAB LEMMON 100MGN74431 001AB NOVOPHARM 50MG

MAY 31, 1995

AB 100MGN74405 002AB ZENITH LABS 50MG

MAY 24, 1995

AB 100MGN74405 001

MAY 24, 1995

N74411 001

MAY 31, 1995

N74411 002

MAY 31, 1995

FLURBIPROFEN SODIUM

SOLUTION/DROPS; OPHTHALMIC

FLURBIPROFEN SODIUMAT BAUSCH AND LOMB 0.03%N74447 001

JAN 04, 1995

OCUFENAT + ALLERGAN 0.03%N19404 001

DEC 31, 1986

FOSCARNET SODIUM

INJECTABLE; INJECTION

FOSCAVIR> DLT > * ASTRA 24MG/MLN20068 001> DLT > + 2.4GM/100ML

SEP 27, 1991

> ADD >N20068 001> ADD >

SEP 27, 1991

FOSINOPRIL SODIUM

TABLET; ORAL

MONOPRIL* BRISTOL MYERS SQUIBB 20MGN19915 003

20MG

MAY 16, 1991

N19915 003

MAY 16, 1991

N19915 004

MAR 28, 1995

+ 40MG

GALLIUM NITRATEINJECTABLE; INJECTION
GANITE

+	FUJISAWA	25MG/ML
+	SOLOPAK	25MG/ML

N19961 002
JAN 17, 1991
N19961 002
JAN 17, 1991

GEMFIBROZIL

CAPSULE; ORAL

GEMFIBROZIL

<u>AB</u>	+	MYLAN	<u>300MG</u>
	@		300MG
<u>AB</u>		PUREPAC PHARM	<u>300MG</u>
	@		300MG

N73466 001
JAN 25, 1993
N73466 001
JAN 25, 1993
N72929 001
JAN 29, 1993
N72929 001
JAN 29, 1993

LOPID

<u>AB</u>	+	PARKE DAVIS	<u>300MG</u>
	@		300MG

N18422 002
N18422 002

TABLET; ORAL

GEMFIBROZIL

<u>AB</u>		CHELSEA LABS	<u>600MG</u>
<u>AB</u>		INVAMED	<u>600MG</u>
<u>AB</u>		MYLAN	<u>600MG</u>

N74442 001
APR 28, 1995
N74615 001
SEP 29, 1995
N74452 001
FEB 16, 1995

GENTAMICIN SULFATE

INJECTABLE; INJECTION

GENTAFATE

<u>AP</u>		PHARMAFAIR	<u>EQ 40MG BASE/ML</u>
	@		EQ 40MG BASE/ML

N62493 001
AUG 28, 1985
N62493 001
AUG 28, 1985

GENTAMICIN

<u>AP</u>		INTL MEDICATION	<u>EQ 1MG BASE/ML</u>
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N62325 003
JUN 23, 1982
N62325 004
JUN 23, 1982

<u>AP</u>			<u>EQ 100MG BASE/100ML</u>
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GENTAMICIN SULFATE

INJECTABLE; INJECTION

GENTAMICIN

<u>AP</u>		INTL MEDICATION	<u>EQ 40MG BASE/ML</u>	N62325 001
	@		EQ 1MG BASE/ML	N62325 003
	@		EQ 100MG BASE/100ML	JUN 23, 1982
	@		EQ 40MG BASE/ML	N62325 004
	@			JUN 23, 1982
	@			N62325 001

OINTMENT; OPHTHALMIC

GENTAMICIN SULFATE

<u>AT</u>		ADV REMEDIES	<u>EQ 0.3% BASE</u>	N64093 001
				AUG 31, 1995

OINTMENT; TOPICAL

GENTAMICIN SULFATE

<u>AT</u>		PHARMADERM	<u>EQ 0.1% BASE</u>	N62534 001
	@		EQ 0.1% BASE	OCT 10, 1984
	@			N62534 001
	@			OCT 10, 1984

SOLUTION/DROPS; OPHTHALMIC

GENTAMICIN SULFATE

<u>AT</u>		ALCON	<u>EQ 0.3% BASE</u>	N62196 001
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GLIMEPIRIDE

TABLET; ORAL

AMARYL

	HOECHST ROUSSEL	1MG	N20496 001
		2MG	NOV 30, 1995
		2MG	N20496 002
		4MG	NOV 30, 1995
		4MG	N20496 003
			NOV 30, 1995

GLIPIZIDE

TABLET; ORAL

GLIPIZIDE

<u>AB</u>		ALPHAPHARM	<u>5MG</u>	N74438 001
				JUN 20, 1995
<u>AB</u>			<u>10MG</u>	N74438 002
				JUN 20, 1995

GLIPIZIDETABLET; ORAL
GLIPIZIDE

> DLT >	AB	BAKER NORTON	5MG
> DLT >			
> DLT >	AB		10MG
> DLT >			
	AB	GENEVA PHARMS	5MG
	AB		10MG
	AB	INVAMED	5MG
	AB		10MG
	AB	WATSON LABS	5MG
	AB		10MG
> ADD >	AB	ZENITH GOLDLINE	5MG
> ADD >			
> ADD >	AB		10MG
> ADD >			

GLUCAGON HYDROCHLORIDEINJECTABLE; INJECTION
GLUCAGON

> DLT >		LILLY	EQ 10MG BASE/VIAL
> ADD >	@		EQ 10MG BASE/VIAL

GLYBURIDE

TABLET; ORAL

GLIBENCLAMIDE

@ HOECHST ROUSSEL

1.5MG

@

3MG

GLYBURIDE

AB NOVOPHARM

1.25MG

AB

2.5MG

AB

5MG

N74497 001
AUG 31, 1995
N74497 002
AUG 31, 1995
N74305 001
APR 07, 1995
N74305 002
APR 07, 1995
N74542 001
JUN 20, 1995
N74542 002
JUN 20, 1995
N74223 001
FEB 27, 1995
N74223 002
FEB 27, 1995
N74497 001
AUG 31, 1995
N74497 002
AUG 31, 1995

GLYBURIDE

TABLET; ORAL

GLYBURIDE (MICRONIZED)

AB HOECHST ROUSSEL 1.5MG

AB 3MG

AB GLYNASE
UPJOHN 1.5MG

AB 3MG

AB MICRONASE
UPJOHN 1.25MG

AB 2.5MG

AB + 5MG

GLYCINE

SOLUTION; IRRIGATION

GLYCINE 1.5% IN PLASTIC CONTAINER

AT BAXTER 1.5GM/100ML

@ 1.5GM/100ML

N20055 001
APR 17, 1992
N20055 002
APR 17, 1992

N20051 001
MAR 04, 1992
N20051 002
MAR 04, 1992

N17498 001
MAY 01, 1984
N17498 002
MAY 01, 1984
N17498 003
MAY 01, 1984

N18522 001
FEB 19, 1982
N18522 001
FEB 19, 1982

GRANISETRON HYDROCHLORIDE

TABLET; ORAL

KYTRIL

+ SMITHKLINE BEECHAM EQ 1MG BASE

N20305 001
MAR 16, 1995

GUANABENZ ACETATE

TABLET; ORAL

GUANABENZ ACETATE

AB ZENITH LABS EQ 4MG BASE

AB EQ 8MG BASE

N74149 001
APR 07, 1995
N74149 002
APR 07, 1995

GUANFACINE HYDROCHLORIDE

TABLET; ORAL

GUANFACINE HCL

AB WATSON LABS EQ 1MG BASE

N74145 001

OCT 17, 1995

AB EQ 2MG BASE

N74145 002

OCT 17, 1995

TENEX

AB ROBINS AH EQ 1MG BASE

N19032 001

OCT 27, 1986

AB + EQ 2MG BASE

N19032 002

NOV 07, 1988

EQ 1MG BASE

N19032 003

OCT 27, 1986

* EQ 2MG BASE

N19032 002

NOV 07, 1988

@ 3MG

N19032 003

NOV 07, 1988

@ EQ 3MG BASE

N19032 003

NOV 07, 1988

HALCINONIDE

CREAM; TOPICAL

HALOG

AT * WESTWOOD SQUIBB 0.1%

N17556 001

+ 0.1%

N17556 001

HALOG-E

AT WESTWOOD SQUIBB 0.1%

N18234 001

0.1%

N18234 001

HALOPERIDOL LACTATE

SOLUTION; ORAL

HALOPERIDOL LACTATE

UDL

EQ 1MG BASE/ML

N74536 001

NOV 02, 1995

HALOPROGIN

SOLUTION; TOPICAL

HALOTEX

* WESTWOOD SQUIBB 1%

N16943 001

@ 1%

N16943 001

HEPARIN CALCIUM

INJECTABLE; INJECTION

CALCIPARINE

* CHOAY

25,000 UNITS/ML

N18237 001

@ SANOFI WINTHROP

25,000 UNITS/ML

N18237 001

HEPARIN SODIUM

INJECTABLE; INJECTION

HEPARIN LOCK FLUSH

AP

SANOFI WINTHROP

10 UNITS/ML

N40082 001

AP

100 UNITS/ML

FEB 28, 1995

N40082 002

FEB 28, 1995

HEPARIN SODIUM

AP

* ABBOTT

2,500 UNITS/ML

N05264 014

APR 07, 1986

*

2,000 UNITS/ML

N05264 013

APR 07, 1986

AP

ELKINS SINN

10,000 UNITS/ML

N17037 013

APR 07, 1986

AP

5,000 UNITS/0.5ML

N17037 013

APR 07, 1986

AP

MARSAM

1,000 UNITS/ML

N40008 001

OCT 10, 1995

AP

PHARMA SERVE NY

1,000 UNITS/ML

N86129 001

AP

WYETH AYERST

2,500 UNITS/ML

N17007 007

AP

+ 2,500 UNITS/ML

N17007 007

HEPARIN SODIUM 1000 UNITS AND DEXTROSE 5% IN PLASTICCONTAINER

AP

MCGAW

200 UNITS/100ML

N19130 001

DEC 31, 1984

@ 200 UNITS/100ML

N19130 001

DEC 31, 1984

HEPARIN SODIUM 1000 UNITS IN SODIUM CHLORIDE 0.9% INPLASTIC CONTAINER

AP

MCGAW

200 UNITS/100ML

N19042 001

MAR 29, 1985

@ 200 UNITS/100ML

N19042 001

MAR 29, 1985

HEPARIN SODIUM 12,500 UNITS IN SODIUM CHLORIDE 0.45% INPLASTIC CONTAINER

AP

ABBOTT

5,000 UNITS/100ML

N18916 006

JAN 31, 1984

5,000 UNITS/100ML

N18916 006

JAN 31, 1984

HEPARIN SODIUMINJECTABLE; INJECTIONHEPARIN SODIUM 12500 UNITS IN SODIUM CHLORIDE 0.45% IN

<u>PLASTIC CONTAINER</u>		
AP	MCGAW	5,000 UNITS/100ML N19802 001 JUL 20, 1992
@		5,000 UNITS/100ML N19802 001 JUL 20, 1992

HEPARIN SODIUM 2000 UNITS IN DEXTROSE 5% IN PLASTIC

<u>CONTAINER</u>		
AP	MCGAW	200 UNITS/100ML N19130 003 DEC 31, 1984
@		200 UNITS/100ML N19130 003 DEC 31, 1984

HEPARIN SODIUM 2000 UNITS IN SODIUM CHLORIDE 0.9% IN

<u>PLASTIC CONTAINER</u>		
AP	MCGAW	200 UNITS/100ML N19042 002 MAR 29, 1985
@		200 UNITS/100ML N19042 002 MAR 29, 1985

HEPARIN SODIUM 25,000 UNITS IN SODIUM CHLORIDE 0.45% IN

<u>PLASTIC CONTAINER</u>		
AP	ABBOTT	5,000 UNITS/100ML N18916 007 JAN 31, 1984
AP		10,000 UNITS/100ML N18916 008 JAN 31, 1984
		5,000 UNITS/100ML N18916 007 JAN 31, 1984
		10,000 UNITS/100ML N18916 008 JAN 31, 1984

HEPARIN SODIUM 25000 UNITS IN SODIUM CHLORIDE 0.45% IN

<u>PLASTIC CONTAINER</u>		
AP	MCGAW	5,000 UNITS/100ML N19802 005 JUL 20, 1992
AP		10,000 UNITS/100ML N19802 002 JUL 20, 1992
@		5,000 UNITS/100ML N19802 005 JUL 20, 1992
@		10,000 UNITS/100ML N19802 002 JUL 20, 1992

HEPARIN SODIUM 25000 UNITS IN SODIUM CHLORIDE 0.9% IN

<u>PLASTIC CONTAINER</u>		
AP	MCGAW	5,000 UNITS/100ML N19802 003 JUL 20, 1992
@		5,000 UNITS/100ML N19802 003 JUL 20, 1992

HEPARIN SODIUMINJECTABLE; INJECTIONHEPARIN SODIUM 5000 UNITS IN DEXTROSE 5% IN PLASTIC

<u>CONTAINER</u>		
AP	MCGAW	1,000 UNITS/100ML N19130 002 DEC 31, 1984
@		1,000 UNITS/100ML N19130 002 DEC 31, 1984

HEPARIN SODIUM PRESERVATIVE FREE

AP	+	ABBOTT	2,500 UNITS/ML	N05264 014 APR 07, 1986
	+		2,000 UNITS/ML	N05264 013 APR 07, 1986
AP		FUJISAWA	1,000 UNITS/ML	N17029 010 APR 28, 1986
AP	+		1,000 UNITS/ML	N17029 010 APR 28, 1986
AP		PHARMA SERVE NY	1,000 UNITS/ML	N86129 001
AP		STERLING WINTHROP	10,000 UNITS/ML	N89522 001 MAY 04, 1987
AP	+		10,000 UNITS/ML	N89522 001 MAY 04, 1987

<u>LIQUAEMIN LOCK PLUSH</u>			
<u>AP</u>	<u>ORGANON</u>	<u>100 UNITS/ML</u>	<u>N00552 007</u>
	@	100 UNITS/ML	N00552 007
<u>LIQUAEMIN SODIUM</u>			
<u>AP</u>	<u>ORGANON</u>	<u>1,000 UNITS/ML</u>	<u>N00552 004</u>
<u>AP</u>		<u>5,000 UNITS/ML</u>	<u>N00552 003</u>
<u>AP</u>		<u>10,000 UNITS/ML</u>	<u>N00552 005</u>
	@	1,000 UNITS/ML	N00552 004
	@	5,000 UNITS/ML	N00552 003
	@	10,000 UNITS/ML	N00552 005

HYDRALAZINE HYDROCHLORIDETABLET; ORAL

<u>DRALZINE</u>				
<u>AA</u>	<u>LEMON</u>	<u>25MG</u>		<u>N84301 001</u>
	@	25MG		N84301 001
<u>HYDRALAZINE HCL</u>				
<u>AA</u>	<u>HALSEY</u>	<u>50MG</u>		<u>N89222 001</u>
	@	50MG		N89222 001
				JAN 22, 1986
				JAN 22, 1986

HYDRALAZINE HYDROCHLORIDE; HYDROCHLOROTHIAZIDE

TABLET; ORAL

APRESOLINE-ESIDRIX

* CIBA

@

25MG;15MG

25MG;15MG

N12026 002

N12026 002

HYDROCHLOROTHIAZIDE

TABLET; ORAL

HYDROCHLOROTHIAZIDEAB

ASCOT

50MG

N87540 001

FEB 03, 1982

@

50MG

N87540 001

FEB 03, 1982

AB

INWOOD LABS

25MG

N85067 001

@

25MG

N85067 001

HYDROCHLOROTHIAZIDE; LOSARTAN POTASSIUM

TABLET; ORAL

HYZAAR

+ MERCK

12.5MG;50MG

N20387 001

APR 28, 1995

HYDROCHLOROTHIAZIDE; METHYLDOPA

TABLET; ORAL

METHYLDOPA AND HYDROCHLOROTHIAZIDEAB

ZENITH LABS

15MG;250MG

N71458 001

MAR 08, 1988

AB25MG;250MG

N71459 001

MAR 08, 1988

AB30MG;500MG

N71460 001

MAR 08, 1988

AB50MG;500MG

N71461 001

MAR 08, 1988

@

15MG;250MG

N71458 001

MAR 08, 1988

@

25MG;250MG

N71459 001

MAR 08, 1988

@

30MG;500MG

N71460 001

MAR 08, 1988

@

50MG;500MG

N71461 001

MAR 08, 1988

HYDROCHLOROTHIAZIDE; METOPROLOL TARTRATE

TABLET; ORAL

LOPRESSOR HCT

CIBA

25MG;50MG

N18303 001

DEC 31, 1984

25MG;100MG

N18303 002

DEC 31, 1984

+

50MG;100MG

N18303 003

DEC 31, 1984

LOPRESSOR HCT 100/25

CIBA

25MG;100MG

N18303 002

DEC 31, 1984

LOPRESSOR HCT 100/50

* CIBA

50MG;100MG

N18303 003

DEC 31, 1984

LOPRESSOR HCT 50/25

CIBA

25MG;50MG

N18303 001

DEC 31, 1984

HYDROCHLOROTHIAZIDE; PROPRANOLOL HYDROCHLORIDE

TABLET; ORAL

PROPRANOLOL HCL AND HYDROCHLOROTHIAZIDEAB

WARNER CHILCOTT

25MG;80MG

N71772 001

JAN 26, 1988

@

25MG;80MG

N71772 001

JAN 26, 1988

HYDROCHLOROTHIAZIDE; TRIAMTERENE

CAPSULE; ORAL

DIAZIDE

SMITHKLINE BEECHAM

25MG;37.5MG

N16042 003

MAR 03, 1994

+

25MG;37.5MG

N16042 003

MAR 03, 1994

TRIAMTERENE AND HYDROCHLOROTHIAZIDEAB

ZENITH LABS

25MG;50MG

N74259 001

MAR 30, 1995

HYDROCORTISONE

CREAM; TOPICAL

FLEXICORTAT

NESTWOOD SQUIBB

0.5%

N87136 003

APR 08, 1982

HYDROCORTISONE

CREAM; TOPICAL

FLEXICORT

AT	WESTWOOD SQUIBB	1%	N87136 002	> ADD >
			APR 08, 1982	> ADD >
AT		2.5%	N87136 001	> ADD >
			APR 08, 1982	> ADD >
@		0.5%	N87136 003	
@		1%	N87136 002	
@		2.5%	N87136 001	
			APR 08, 1982	
AT	<u>HYDROCORTISONE</u>	0.5%	N84970 002	
AT	CLAY PARK	1%	N85026 001	
@		0.5%	N84970 002	
@		1%	N85026 001	

ENEMA; RECTAL
CORTENEMA

BR	+	SOLVAY	100MG/60ML	N16199 001
			100MG/60ML	N16199 001
BR	+	HYDROCORTISONE		
		COPLEY PHARM	100MG/60ML	N74171 001
				MAY 27, 1994
@			100MG/60ML	N74171 001
				MAY 27, 1994

LOTION; TOPICAL

HYDROCORTISONE

AT	CLAY PARK	0.5%	N85662 001	
@		0.5%	N85662 001	

OINTMENT; TOPICAL
HYDROCORTISONE

@	CLAY PARK	0.5%	N84969 003	
@		0.5%	N84969 003	

HYDROCORTISONE; NEOMYCIN SULFATE; POLYMYXIN B SULFATESOLUTION/DROPS; OTIC
CORTISPORIN

AT	+	BURROUGHS WELLCOME	10.1MG/ML; EQ 3.5MG BASE/ML;	
			12,000 UNITS/ML	N50479 001
AT	+	GLAXO WELLCOME	1%; EQ 3.5MG BASE/ML;	
			10,000 UNITS/ML	N50479 001

HYDROCORTISONE; NEOMYCIN SULFATE; POLYMYXIN B SULFATE

SOLUTION/DROPS; OTIC

NEOMYCIN AND POLYMYXIN B SULFATES AND HYDROCORTISONE

AT	BAUSCH AND LOMB	1%; EQ 3.5MG BASE/ML;		
		10,000 UNITS/ML		N64053 001
				DEC 29, 1995

HYDROCORTISONE; UREA

CREAM; TOPICAL

ALPHADERM

AT	VIVAN	1%; 10%		N85008 001
@		1%; 10%		N86008 001
AT	* CALMURID HC	1%; 10%		N83947 001
@	* PHARMACIA	1%; 10%		N83947 001

HYDROCORTISONE ACETATE

AEROSOL; RECTAL

CORTIFOAM

+	REED AND CARNRICK	10%		N17351 001
				FEB 10, 1982
+	SPKU	10%		N17351 001
				FEB 10, 1982

HYDROCORTISONE ACETATE; PRAMOXINE HYDROCHLORIDE

AEROSOL; TOPICAL

EPIFOAM

BX	REED AND CARNRICK	1%; 1%		N86457 001
BX	SPKU	1%; 1%		N86457 001
BX	PROCTOFOAM HC	1%; 1%		N86195 001
BX	REED AND CARNRICK	1%; 1%		N86195 001
BX	SPKU	1%; 1%		N86195 001

HYDROCORTISONE BUTYRATE

OINTMENT; TOPICAL

LOCOID

@	YAMANOUCHI	0.1%		N18652 001
				OCT 29, 1982
+		0.1%		N18652 001
				OCT 29, 1982

HYDROCORTISONE SODIUM SUCCINATE

INJECTABLE; INJECTION

A-HYDROCORT

> DLT >	AP	ABBOTT	EQ 100MG BASE/VIAL	N89577 001
> DLT >				APR 11, 1989
> ADD >	@		EQ 100MG BASE/VIAL	N89577 001
> ADD >				APR 11, 1989

HYDROMORPHONE HYDROCHLORIDE

INJECTABLE; INJECTION

DILAUDID-HP

AP	+	KNOLL PHARM	10MG/ML	N19034 001
				JAN 11, 1984

HYDROMORPHONE HCL

AP		STERIS	10MG/ML	N74317 001
				AUG 23, 1995

HYDROXYCHLOROQUINE SULFATE

TABLET; ORAL

HYDROXYCHLOROQUINE SULFATE

AB		GENEVA PHARMS	200MG	N40104 001
				NOV 30, 1995
AB		ROYCE LABS	200MG	N40133 001
				NOV 30, 1995

HYDROXYPROGESTERONE CAPROATE

INJECTABLE; INJECTION

HYDROXYPROGESTERONE CAPROATE

*	AKORN	125MG/ML	N18004 001
@		125MG/ML	N18004 001

HYDROXYUREA

CAPSULE; ORAL

HYDREA

AB	+	SQUIBB	500MG	N16295 001
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HYDROXYUREA

AB		ROXANE	500MG	N74476 001
				AUG 18, 1995

HYDROXYZINE HYDROCHLORIDE

INJECTABLE; INJECTION

HYDROXYZINE HCL

AP		PHARMAFATR	50MG/ML	N88881 001
				FEB 14, 1986
	@		50MG/ML	N88881 001
				FEB 14, 1986
AP		STERIS	25MG/ML	N87274 001
AP			50MG/ML	N87274 002
	@		25MG/ML	N87274 001
	@		50MG/ML	N87274 002

SYRUP; ORAL

HYDROXYZINE HCL

AA		BARRE	10MG/5ML	N88785 001
				FEB 03, 1988
	@		10MG/5ML	N88785 001
				FEB 03, 1988

TABLET; ORAL

HYDROXYZINE HCL

AB		HAUSEY	50MG	N89396 001
				MAY 02, 1988
	@		50MG	N89396 001
				MAY 02, 1988
AB		SIDMAK LABS NJ	100MG	N81054 001
				SEP 25, 1995

IBUPROFEN

SUSPENSION; ORAL

CHILDREN'S MOTRIN

BX	+	MCNEIL CONS PRODS	100MG/5ML	N19842 001
				SEP 19, 1989

PEDIA PROFEN

BX	+	MCNEIL CONS PRODS	100MG/5ML	N19842 001
				SEP 19, 1989

SUSPENSION/DROPS; ORAL

MOTRIN

+	MCNEIL CONS PRODS	40MG/ML	N20476 001
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MAY 25, 1995

TABLET; ORAL

IBU-TAB

AB		ALRA	800MG	N71965 001
				AUG 11, 1988

IBUPROFEN

TABLET; ORAL

IBU-TAB

@ ALRA

800MG

N71965 001
AUG 11, 1988IBUPROFENLEMMON400MG

N73343 001

JUN 30, 1992

N73344 001

JUN 30, 1992

N73345 001

JUN 30, 1992

N73343 001

JUN 30, 1992

N73344 001

JUN 30, 1992

N73345 001

JUN 30, 1992

> DLT > AB
> DLT >
> DLT > AB
> DLT >
> DLT > AB
> DLT >
> ADD > @
> ADD >
> ADD > @
> ADD >
> ADD > @
> ADD >

> ADD > IBUTILIDE FUMARATE

> ADD > INJECTABLE; INJECTION
> ADD > CORVERT
> ADD > + UPJOHN
> ADD >

0.1MG/ML

N20491 001
DEC 28, 1995INDAPAMIDE

TABLET; ORAL

INDAPAMIDEAB

ZENITH LABS

2.5MGN74299 001
JUL 27, 1995LOZOLAB

+ RHONE POULENC RORER

2.5MGN18538 001
JUL 06, 1983INDECAINIDE HYDROCHLORIDE

TABLET, EXTENDED RELEASE, ORAL

DECABIDLILLYEQ 50MG BASE

N19693 001

DEC 29, 1989

EQ 75MG BASE

N19693 002

DEC 29, 1989

INDECAINIDE HYDROCHLORIDE

TABLET, EXTENDED RELEASE, ORAL

DECABID* LILLYEQ 100MG BASE

N19693 003

DEC 29, 1989

@

EQ 50MG BASE

N19693 001

DEC 29, 1989

@

EQ 75MG BASE

N19693 002

DEC 29, 1989

@

EQ 100MG BASE

N19693 003

DEC 29, 1989

INULIN

INJECTABLE; INJECTION

INULIN AND SODIUM CHLORIDE

+ CYPROS

100MG/ML

N02282 001

* ISO TEX

100MG/ML

N02282 001

IOCTETAMIC ACID

TABLET, ORAL

CHOLEBRINE* MALLINCKRODT750MG

N17129 001

@

750MG

N17129 001

IOHEXOL

SOLUTION; INJECTION, ORAL

OMNIPAQUE 350

NYCOMED

75.5%

N20608 003

OCT 24, 1995

SOLUTION; INJECTION, ORAL, RECTAL

OMNIPAQUE 240

NYCOMED

51.8%

N20608 001

OCT 24, 1995

OMNIPAQUE 300

NYCOMED

64.7%

N20608 002

OCT 24, 1995

IOPAMIDOL

INJECTABLE; INTRAVASCULAR

ISOVUE-200

@ BRACCO 41%

BRACCO DXS 41%

N20327 001

OCT 12, 1994

N20327 001

OCT 12, 1994

IOPROMIDE

INJECTABLE; INJECTION

ULTRAVIST

* BERLEX

EQ 150MG IODINE/ML

N20220 004

MAY 10, 1995

*

EQ 240MG IODINE/ML

N20220 003

MAY 10, 1995

*

EQ 300MG IODINE/ML

N20220 002

MAY 10, 1995

*

EQ 370MG IODINE/ML

N20220 001

MAY 10, 1995

ULTRAVIST 150

+ BERLEX

31.2%

N20220 004

MAY 10, 1995

ULTRAVIST 240

+ BERLEX

49.9%

N20220 003

MAY 10, 1995

ULTRAVIST 300

+ BERLEX

62.3%

N20220 002

MAY 10, 1995

ULTRAVIST 370

+ BERLEX

76.9%

N20220 001

MAY 10, 1995

IOTHALAMATE SODIUM

INJECTABLE; INJECTION

ANGIO-CONRAY

* MALLINCKRODT

80%

N13319 001

@

80%

N13319 001

IOTHALAMATE SODIUM, I-125

INJECTABLE; INJECTION

GLOFIL-125

CYPROS

250-300 uCi/ML

N17279 001

IOTHALAMATE SODIUM, I-125

INJECTABLE; INJECTION

GLOFIL-125

ISO-TEX

250-300 uCi/ML

N17279 001

IOTROLAN

INJECTABLE; INTRATHECAL

OSMOVIST

BERLEX

EQ 150MG IODINE/ML

N19580 001

EQ 240MG IODINE/ML

DEC 07, 1989

OSMOVIST 190

BERLEX

40.6%

N19580 001

OSMOVIST 240

BERLEX

51.3%

N19580 002

DEC 07, 1989

IOXILAN

INJECTABLE; INJECTION

OXILAN-300

COOK IMAGING

62%

N20316 001

OXILAN-350

COOK IMAGING

73%

N20316 002

DEC 21, 1995

IPODATE SODIUM

CAPSULE; ORAL

BILIVIST

AA * BERLEX

500MG

N87768 001

@

500MG

AUG 11, 1982

ORAGRAFIN SODIUM

+ BRACCO

BRACCO DXS

500MG

N12967 001

500MG

N12967 001

IPRATROPIUM BROMIDESPRAY, METERED; NASAL
ATROVENT+ BOEHRINGER INGELHEIM 0.021MG/INH
+ 0.042MG/INHN20393 001
OCT 20, 1995
N20394 001
OCT 20, 1995ISOETHARINE HYDROCHLORIDESOLUTION; INHALATION
ISOETHARINE HCL S/FAN * DEX 1%
@ 1%N89252 001
SEP 15, 1986
N89252 001
SEP 15, 1986ISOFLURANELIQUID; INHALATION
ISOFLURANEAN MARSAM 99.9%
AN RHONE POULENC 99.9%N74393 001
MAY 12, 1995
N74502 001
JUN 27, 1995ISOSORBIDE DINITRATECAPSULE, EXTENDED RELEASE; ORAL
DILATRATE-SRBC REED AND CARRICK 40MG
BC SPKU 40MGN19790 001
SEP 02, 1988
N19790 001
SEP 02, 1988ISOSORBIDE MONONITRATETABLET, EXTENDED RELEASE; ORAL
IMDUR@ SCHERING 30MG
+ 60MGN20225 001
AUG 12, 1993
N20225 002
AUG 12, 1993ISOSORBIDE MONONITRATETABLET, EXTENDED RELEASE; ORAL
IMDUR+ SCHERING 120MG
@ SCHERING PLOUGH 30MG
60MGN20225 003
MAR 30, 1995
N20225 001
AUG 12, 1993
N20225 002
AUG 12, 1993KANAMYCIN SULFATE

INJECTABLE; INJECTION

KANAMYCIN
AP ELKINS SINNAP EQ 75MG BASE/2ML
AP EQ 500MG BASE/2ML
AP EQ 1GM BASE/3ML
@ EQ 75MG BASE/2ML
@ EQ 500MG BASE/2ML
@ EQ 1GM BASE/3MLN62324 001
N62324 002
N62324 003
N62324 001
N62324 002
N62324 003KETOPROFEN

CAPSULE; ORAL

KETOPROFEN

> ADD > AB GENEVA PHARMS 50MG
> ADD >
> ADD > AB 75MG
> ADD >N74024 001
DEC 29, 1995
N74024 002
DEC 29, 1995CAPSULE, EXTENDED RELEASE; ORAL
ORUVAIL+ WYETH AYERST 100MG
+ 150MGN19816 003
FEB 08, 1995
N19816 002
FEB 08, 1995LACTULOSE

SOLUTION; ORAL

LACTULOSE

AA HI TECH PHARMA 10GM/15ML

N74076 001
JUL 03, 1995

LACTULOSE

SOLUTION; ORAL, RECTAL
LACTULOSE
AA HI TECH PHARMA 10GM/15ML N74077 001
 JUL 03, 1995

LAMIVUDINE

SOLUTION; ORAL
 EPIVIR
 GLAXO WELLCOME 10MG/ML N20596 001
 NOV 17, 1995

TABLET; ORAL
 EPIVIR
 + GLAXO WELLCOME 150MG N20564 001
 NOV 17, 1995

LANSOPRAZOLE

CAPSULE, DELAYED REL GRANULES; ORAL
 PREVACID
 TAP HOLDINGS 15MG N20406 001
 MAY 10, 1995
 + 30MG N20406 002
 MAY 10, 1995

LEUCOVORIN CALCIUM

INJECTABLE; INJECTION
 LEUCOVORIN CALCIUM
 + CETUS BEN VENUE EQ 200MG BASE/VIAL N40056 001
 MAY 23, 1995

WELLCOVORIN
AP BURROUGHS WELLCOME EQ 50MG BASE/VIAL N89465 001
 JAN 23, 1989
 * EQ 25MG BASE/VIAL N89833 001
 JAN 23, 1989
 @ GLAXO WELLCOME EQ 25MG BASE/VIAL N89833 001
 JAN 23, 1989
 @ EQ 50MG BASE/VIAL N89465 001
 JAN 23, 1989

LEUPROLIDE ACETATE

INJECTABLE; INJECTION
 LUPRON
 TAP HOLDINGS 5MG/ML N20263 001
 APR 16, 1993
 + 1MG/0.2ML N19010 001
 APR 09, 1985
 * TAP PHARMS 5MG/ML N19010 001
 APR 09, 1985
 LUPRON DEPOT
 + TAP HOLDINGS 22.5MG/VIAL N20517 001
 DEC 22, 1995

LIDOCAINE HYDROCHLORIDE

INJECTABLE; INJECTION
LIDOCAINE HCL
AP AKORN 1% N85037 001
AP 2% N85037 002
 @ 1% N85037 001
 @ 2% N85037 002
AP FUJISAWA 2% N80420 004
 @ 2% N80420 004
AP SANOFI WINTHROP 1% N40013 001
 JUN 23, 1995
AP 2% N40078 001
 JUN 23, 1995

SOLUTION; ORAL
LIDOCAINE HCL
AT HI TECH PHARMA 2% N40014 001
 JUL 10, 1995

LINDANE

LOTION; TOPICAL
SCABENE
AT STIEFEL 1% N86769 001
 @ 1% N86769 001
 SHAMPOO; TOPICAL
SCABENE
AT STIEFEL 1% N87940 001
 APR 08, 1983
 @ 1% N87940 001
 APR 08, 1983

LISINOPRIL

TABLET; ORAL

AB PRINIVIL 2.5MG
MERCK
2.5MG

N19558 006
JAN 28, 1994
N19558 006
JAN 28, 1994

AB ZESTRIL 2.5MG
ZENECA
2.5MG

N19777 005
APR 29, 1993
N19777 005
APR 29, 1993

LITHIUM CARBONATE

TABLET; ORAL

AB LITHOTABS 300MG
SOLVAY
AB + 300MG

N16980 001
N16980 001

TABLET, EXTENDED RELEASE; ORAL

LITHOBID 300MG
@ SOLVAY 300MG

N18027 001
N18027 001

LOSARTAN POTASSIUM

TABLET; ORAL

COZAAR 25MG
MERCK

N20386 001
APR 14, 1995

+ 50MG

N20386 002
APR 14, 1995

MAGNESIUM CHLORIDE; POTASSIUM CHLORIDE; SODIUM ACETATE; SODIUM CHLORIDE; SODIUM GLUCONATE

SOLUTION; IRRIGATION

PHYSIOLYTE IN PLASTIC CONTAINER

AT MCGAW 30MG/100ML; 37MG/100ML; 370MG/100ML;
530MG/100ML; 500MG/100ML N19024 001
JUN 08, 1984

MAGNESIUM CHLORIDE; POTASSIUM CHLORIDE; SODIUM ACETATE; SODIUM CHLORIDE; SODIUM GLUCONATE

SOLUTION; IRRIGATION

PHYSIOLYTE IN PLASTIC CONTAINER

MCGAW 30MG/100ML; 37MG/100ML; 370MG/100ML;
530MG/100ML; 500MG/100ML N19024 001
JUN 08, 1984

PHYSIOSOL IN PLASTIC CONTAINER

AT ABBOTT 30MG/100ML; 37MG/100ML; 222MG/100ML;
526MG/100ML; 502MG/100ML N17637 002
JUL 08, 1982
30MG/100ML; 37MG/100ML; 222MG/100ML;
526MG/100ML; 502MG/100ML N17637 002
JUL 08, 1982

SYNOVALYTE IN PLASTIC CONTAINER

AT BAXTER 30MG/100ML; 37MG/100ML; 368MG/100ML;
526MG/100ML; 502MG/100ML N19326 001
JAN 25, 1985
30MG/100ML; 37MG/100ML; 368MG/100ML;
526MG/100ML; 502MG/100ML N19326 001
JAN 25, 1985

MAGNESIUM SULFATE

INJECTABLE; INJECTION

MAGNESIUM SULFATE IN DEXTROSE 5% IN PLASTIC CONTAINER

ABBOTT 1GM/100ML N20488 001
JUL 11, 1995
+ 2GM/100ML N20488 002
JUL 11, 1995

MANNITOL

INJECTABLE; INJECTION

MANNITOL 10%

AP ABBOTT 10GM/100ML N16269 002
@ 10GM/100ML N16269 002

MANNITOL 15%

AP ABBOTT 15GM/100ML N16269 003
@ 15GM/100ML N16269 003

MANNITOL 20%

AP ABBOTT 20GM/100ML N16269 004
@ 20GM/100ML N16269 004

MANNITOL 25%

AP ABBOTT 12.5GM/50ML N16269 005

MANNITOL

INJECTABLE; INJECTION

<u>AP</u>	<u>MANNITOL 25%</u>	<u>12.5GM/50ML</u>	N16269 006
	ABBOTT		AUG 25, 1994
	@	12.5GM/50ML	N16269 005
<u>AP</u>	<u>MANNITOL 5%</u>	<u>5GM/100ML</u>	N16269 001
	ABBOTT	5GM/100ML	N16269 001
	@		

MASOPROCOL

CREAM; TOPICAL

ACTINEX

* BLOCK DRUG 10%

+ SPKU 10%

N19940 001 > DLT >
 SEP 04, 1992 > ADD >
 N19940 001 > DLT >
 SEP 04, 1992 > ADD >

MEGESTROL ACETATE

TABLET; ORAL

MEGESTROL ACETATE

<u>AB</u>	BARR	<u>40MG</u>	N74621 001
			NOV 30, 1995
<u>AB</u>	ROXANE	<u>20MG</u>	N74458 001
			SEP 29, 1995
<u>AB</u>		<u>40MG</u>	N74458 002
			SEP 29, 1995

MEPERIDINE HYDROCHLORIDE

TABLET; ORAL

DEMEROL

STERLING WINTHROP 50MG
 + 50MG
100MG
 + 100MG

N05010 001
 N05010 001
 N05010 004
 N05010 004

MEBENDAZOLE

TABLET, CHEWABLE; ORAL

MEBENDAZOLEAB COPLEY PHARM 100MG

N73580 001
 JAN 04, 1995

VERMOXAB + JANSSEN 100MG

N17481 001

MESTRANOL; NORETHINDRONE

TABLET; ORAL-20

NORINYL

@ SEARLE 0.1MG; 2MG
 @ SYNTEX 0.1MG; 2MG

N13625 004
 N13625 004

TABLET; ORAL-21

NORINYL 1+50 21-DAY

AB SEARLE 0.05MG; 1MG
AB SYNTEX 0.05MG; 1MG

N13625 002
 N13625 002

NORINYL 1+80 21-DAY

@ SEARLE 0.08MG; 1MG
 @ SYNTEX 0.08MG; 1MG

N16724 001
 N16724 001

TABLET; ORAL-28

NORINYL 1+50 28-DAY

AB SEARLE 0.05MG; 1MG
AB SYNTEX 0.05MG; 1MG

N16659 001
 N16659 001

NORINYL 1+80 28-DAY

@ SEARLE 0.08MG; 1MG
 @ SYNTEX 0.08MG; 1MG

N16725 001
 N16725 001

MEDROXYPROGESTERONE ACETATE

INJECTABLE; INJECTION

DEPO-PROVERA

* UPJOHN 100MG/ML
 @ 100MG/ML

N12541 002
 N12541 002

MEGESTROL ACETATE

TABLET; ORAL

MEGACEAB BRISTOL MYERS SQUIBB 20MG

N16979 001

AB + 40MG

N16979 002

AB MEAD JOHNSON 20MG

N16979 001

AB * 40MG

N16979 002

METAPROTERENOL SULFATE

SOLUTION; INHALATION
METAPROTERENOL SULFATE
 AN BARRE 0.4%
 AN 0.6%
 AN PACO 0.4%
 AN 0.6%

N71855 001
 JUL 14, 1988
 N71726 001
 JUL 14, 1988
 N71855 001
 JUL 14, 1988
 N71726 001
 JUL 14, 1988

SYRUP; ORAL

ALUPENT
 AA BOEHRINGER INGELHEIM 10MG/5ML
 AA + 10MG/5ML

N17571 001
 N17571 001

METFORMIN HYDROCHLORIDE

TABLET; ORAL
 GLUCOPHAGE
 BRISTOL MYERS SQUIBB 500MG
 + 850MG
 LIPHA 500MG
 * 850MG

N20357 001
 DEC 29, 1994
 N20357 002
 DEC 29, 1994
 N20357 001
 DEC 29, 1994
 N20357 002
 DEC 29, 1994

METHADONE HYDROCHLORIDE

POWDER; FOR RX COMPOUNDING

METHADONE HCL
 MALLINCKRODT 50GM/BOT
 100GM/BOT
 500GM/BOT

N06383 002
 N06383 003
 N06383 004

TABLET, DISPERSIBLE; ORAL

METHADONE HCL
 AA ROXANE 40MG

N74081 001
 APR 28, 1995

METHICILLIN SODIUM

INJECTABLE; INJECTION

STAPHICILLIN
 @ APOTHECON

EQ 900MG BASE/VIAL
 EQ 3.6GM BASE/VIAL
 EQ 5.4GM BASE/VIAL
 EQ 900MG BASE/VIAL
 EQ 3.6GM BASE/VIAL
 EQ 5.4GM BASE/VIAL

N50117 001
 N50117 002
 N50117 003
 N50117 001
 N50117 002
 N50117 003

METHOTRIMEPAZINE

INJECTABLE; INJECTION

LEVOPROME

+ IMMUNEX
 * LEDERLE

20MG/ML
 20MG/ML

N15865 001
 N15865 001

METHYLDOPATE HYDROCHLORIDE

INJECTABLE; INJECTION

METHYLDOPATE HCL

AP DUPONT MERCK

50MG/ML

N70691 001

AP

50MG/ML

JUN 19, 1987

AP

FAULDING

50MG/ML

N70849 001

AP

50MG/ML

JUN 19, 1987

N70691 001

JUN 19, 1987

N70849 001

JUN 19, 1987

METHYLPREDNISOLONE ACETATE

INJECTABLE; INJECTION

METHYLPREDNISOLONE ACETATE

BP AKORN

40MG/ML

N86903 001

BP

80MG/ML

OCT 20, 1982

N86903 002

@

40MG/ML

OCT 20, 1982

N86903 001

@

80MG/ML

OCT 20, 1982

N86903 002

OCT 20, 1982

METOCLOPRAMIDE HYDROCHLORIDE

INJECTABLE; INJECTION

METOCLOPRAMIDE HCL

<u>AP</u>	DUPONT MERCK	<u>EQ 10MG BASE/2ML</u>	N70847 001 NOV 07, 1988
<u>AP</u>		<u>EQ 10MG BASE/2ML</u>	N71291 001 MAR 03, 1989
<u>AP</u>	FAULDING	<u>EQ 10MG BASE/2ML</u>	N70847 001 NOV 07, 1988
<u>AP</u>		<u>EQ 10MG BASE/2ML</u>	N71291 001 MAR 03, 1989

TABLET; ORAL

METOCLOPRAMIDE HCL

<u>AB</u>	INVAMED	<u>EQ 5MG BASE</u>	N74478 001 OCT 05, 1995
<u>AB</u>		<u>EQ 10MG BASE</u>	N74478 002 OCT 05, 1995
> <u>ADD</u> >	<u>AB</u>	<u>EQ 5MG BASE</u>	N72750 001 DEC 28, 1995
> <u>ADD</u> >	SIDMAK LABS NJ		

METOPROLOL FUMARATE

TABLET, EXTENDED RELEASE; ORAL

LOPRESSOR

GEIGY

		<u>EQ 100MG TARTRATE</u>	N19786 001 DEC 27, 1989
		<u>EQ 200MG TARTRATE</u>	N19786 002 DEC 27, 1989
		<u>EQ 300MG TARTRATE</u>	N19786 003 DEC 27, 1989
		<u>EQ 400MG TARTRATE</u>	N19786 004 DEC 27, 1989
+			
@		<u>EQ 100MG TARTRATE</u>	N19786 001 DEC 27, 1989
@		<u>EQ 200MG TARTRATE</u>	N19786 002 DEC 27, 1989
@		<u>EQ 300MG TARTRATE</u>	N19786 003 DEC 27, 1989
@		<u>EQ 400MG TARTRATE</u>	N19786 004 DEC 27, 1989

METOPROLOL TARTRATE

TABLET; ORAL

METOPROLOL TARTRATE

<u>AB</u>	LEMMON	<u>50MG</u>	N74141 001 JAN 31, 1995
<u>AB</u>		<u>100MG</u>	N74141 002 JAN 31, 1995
<u>AB</u>	PAR PHARM	<u>50MG</u>	N74453 001 APR 27, 1995
<u>AB</u>		<u>100MG</u>	N74453 002 APR 27, 1995

METRONIDAZOLE

CAPSULE; ORAL

FLAGYL

+ SEARLE 375MG

N20334 001
MAY 03, 1995

CREAM; TOPICAL

METROCREAM

+ GALDERMA 0.75%

N20531 001
SEP 20, 1995METYRAPONE

TABLET, ORAL

METOPIRONE+ CIBA 250MG
@ 250MGN12911 001
N12911 001MEXILETINE HYDROCHLORIDE

CAPSULE; ORAL

MEXILETINE HCLAB NOVOPHARM 150MGN74377 001
MAY 16, 1995AB 200MGN74377 002
MAY 16, 1995AB 250MGN74377 003
MAY 16, 1995MEXITILAB BOEHRINGER INGELHEIM 150MGN18873 002
DEC 30, 1985

MEXILETINE HYDROCHLORIDE

CAPSULE; ORAL

MEXITILAB BOEHRINGER INGELHEIM 200MGN18873 003
DEC 30, 1985
N18873 004
DEC 30, 1985AB + 250MGMICONAZOLE NITRATE

SUPPOSITORY; VAGINAL

MICONAZOLE NITRATEAB ABLE 200MGN73508 001
NOV 19, 1993
N73508 001
NOV 19, 1993AB NMC 200MGMINOXIDIL

TABLET; ORAL

MINOXIDIL> ADD > AB MUTUAL PHARM 2.5MG
> ADD >
> ADD > AB 10MG
> ADD >N72708 001
DEC 14, 1995
N72709 001
DEC 14, 1995> DLT >
> DLT >
> ADD >
> ADD >MITOMYCIN

INJECTABLE; INJECTION

MITOMYCINAP CETUS BEN VENUE 5MG/VIALN64117 001
APR 19, 1995
N64117 002
APR 19, 1995AP 20MG/VIAL20MG/VIALAP FAULDING 20MG/VIALN64117 002
APR 19, 1995
N64106 001
NOV 29, 1995MOEXIPRIL HYDROCHLORIDE

TABLET; ORAL

UNIVASC

SPKU

7.5MG

N20312 001
APR 19, 1995MOEXIPRIL HYDROCHLORIDE

TABLET; ORAL

UNIVASC

+ SPKU

15MG

N20312 002
APR 19, 1995MORPHINE SULFATE

TABLET, EXTENDED RELEASE; ORAL

MS CONTIN

BC + PURDUE FREDERICK 15MG

N19516 003
SEP 12, 1989BC ORAMORPH SR 15MG
ROXANEN19977 004
NOV 23, 1994MUPIROCIN CALCIUM

OINTMENT; NASAL

BACTROBAN

* SMITHKLINE BEECHAM

EQ 2% ACID

N50703 001
SEP 18, 1995
N50703 001
SEP 18, 1995

+

EQ 2% BASE

MYCOPHENOLATE MOFETIL

CAPSULE; ORAL

CELLCEPT

+ SYNTEX

250MG

N50722 001
MAY 03, 1995NADOLOL

TABLET; ORAL

NADOLOLAB INVAMED 20MGN74501 001
NOV 09, 1995AB 40MGN74501 002
NOV 09, 1995AB 80MGN74501 003
NOV 09, 1995

NAFARELIN ACETATESPRAY, METERED; NASAL
SYNAREL

+ SEARLE

EQ 0.2MG BASE/INH

N19886 001

FEB 13, 1990

* SYNTEX

EQ 0.2MG BASE/INH

N19886 001

FEB 13, 1990

NAFCILLIN SODIUM

INJECTABLE; INJECTION

NAFCILAP APOTHECONEQ 500MG BASE/VIAL

N61984 001

APEQ 500MG BASE/VIAL

N62527 001

AUG 02, 1984

APEQ 1GM BASE/VIAL

N61984 002

APEQ 1GM BASE/VIAL

N62527 002

AUG 02, 1984

APEQ 1GM BASE/VIAL

N62732 001

DEC 23, 1986

APEQ 2GM BASE/VIAL

N61984 003

APEQ 2GM BASE/VIAL

N62527 003

AUG 02, 1984

APEQ 2GM BASE/VIAL

N62732 002

DEC 23, 1986

APEQ 4GM BASE/VIAL

N61984 005

APEQ 10GM BASE/VIAL

N62527 004

AUG 02, 1984

NAFCILLIN SODIUMAP APOTHECONEQ 500MG BASE/VIAL

N61984 001

APEQ 500MG BASE/VIAL

N62527 001

AUG 02, 1984

APEQ 1GM BASE/VIAL

N61984 002

APEQ 1GM BASE/VIAL

N62527 002

AUG 02, 1984

APEQ 1GM BASE/VIAL

N62732 001

DEC 23, 1986

APEQ 2GM BASE/VIAL

N61984 003

APEQ 2GM BASE/VIAL

N62527 003

AUG 02, 1984

APEQ 2GM BASE/VIAL

N62732 002

DEC 23, 1986

APEQ 4GM BASE/VIAL

N61984 005

APEQ 10GM BASE/VIAL

N62527 004

AUG 02, 1984

NALMEFENE HYDROCHLORIDEINJECTABLE; INJECTION
REVEX

OHMEDA

EQ 0.1MG BASE/ML

N20459 001

+

EQ 1MG BASE/ML

APR 17, 1995

N20459 002

APR 17, 1995

NALOXONE HYDROCHLORIDE

INJECTABLE; INJECTION

NALOXONE HCLAP FUJISAWA0.02MG/ML

N70648 001

AP0.4MG/ML

NOV 17, 1986

AP0.4MG/ML

N70649 001

@

0.02MG/ML

NOV 17, 1986

@

0.4MG/ML

N70648 001

AP0.4MG/ML

NOV 17, 1986

AP0.4MG/ML

N70649 001

AP0.4MG/ML

NOV 17, 1986

AP0.4MG/ML

N71811 001

AP0.4MG/ML

JUL 19, 1988

AP0.4MG/ML

N71811 001

AP0.4MG/ML

JUL 19, 1988

NANDROLONE DECANOATE

INJECTABLE; INJECTION

NANDROLONE DECANOATEAO AKORN100MG/ML

N87519 001

AO100MG/ML

SEP 28, 1983

@

100MG/ML

N87519 001

SEP 28, 1983

NAPROXEN

TABLET; ORAL

NAPROXENAB CHELSEA LABS250MG

N74457 001

AB375MG

MAY 31, 1995

AB375MG

N74457 002

AB500MG

MAY 31, 1995

N74457 003

MAY 31, 1995

NAPROXEN

TABLET; ORAL

NAPROXEN

<u>AB</u>	DANBURY PHARMA	<u>250MG</u>
<u>AB</u>		<u>375MG</u>
<u>AB</u>		<u>500MG</u>
> <u>DLT</u> >	<u>AB</u> HAMILTON PHARMS	<u>250MG</u>
> <u>DLT</u> >		
> <u>DLT</u> >	<u>AB</u>	<u>375MG</u>
> <u>DLT</u> >		
> <u>DLT</u> >	<u>AB</u>	<u>500MG</u>
> <u>DLT</u> >		
> <u>ADD</u> >	@	250MG
> <u>ADD</u> >		
> <u>ADD</u> >	@	375MG
> <u>ADD</u> >		
> <u>ADD</u> >	@	500MG
> <u>ADD</u> >		
<u>AB</u>	MOVA	<u>250MG</u>
<u>AB</u>		<u>375MG</u>
<u>AB</u>		<u>500MG</u>
<u>AB</u>	ZENITH LABS	<u>250MG</u>
<u>AB</u>		<u>375MG</u>
<u>AB</u>		<u>500MG</u>

NAPROXEN SODIUM

TABLET; ORAL

NAPROXEN SODIUM

<u>AB</u>	CHELSEA LABS	<u>EQ 250MG BASE</u>
<u>AB</u>		<u>EQ 500MG BASE</u>
> <u>DLT</u> >	<u>AB</u> HAMILTON PHARMS	<u>EQ 250MG BASE</u>
> <u>DLT</u> >		
> <u>DLT</u> >	<u>AB</u>	<u>EQ 500MG BASE</u>
> <u>DLT</u> >		

N74163 001	> <u>ADD</u> >
FEB 10, 1995	> <u>ADD</u> >
N74163 002	> <u>ADD</u> >
FEB 10, 1995	> <u>ADD</u> >
N74163 003	
FEB 10, 1995	
N74110 001	
OCT 30, 1992	
N74110 002	
OCT 30, 1992	
N74110 003	
OCT 30, 1992	
N74110 001	
OCT 30, 1992	
N74110 002	
OCT 30, 1992	
N74110 003	
OCT 30, 1992	
N74410 001	
APR 28, 1995	
N74410 002	
APR 28, 1995	
N74410 003	
APR 28, 1995	
N74111 001	
FEB 28, 1995	
N74111 002	
FEB 28, 1995	
N74111 003	
FEB 28, 1995	

NAPROXEN SODIUM

TABLET; ORAL

NAPROXEN SODIUM

@ HAMILTON PHARMS	EQ 250MG BASE
@	EQ 500MG BASE
<u>AB</u> PUREPAC PHARM	<u>EQ 250MG BASE</u>
<u>AB</u>	<u>EQ 500MG BASE</u>
<u>AB</u> ZENITH LABS	<u>EQ 250MG BASE</u>
<u>AB</u>	<u>EQ 500MG BASE</u>

NEOMYCIN SULFATE

TABLET; ORAL

NEOMYCIN SULFATE

<u>AA</u> BIOCRAFT	<u>EQ 350MG BASE</u>	N50304 001
	EQ 350MG BASE	N60304 001
<u>AA</u> LILLY	<u>EQ 350MG BASE</u>	N60385 001
@	EQ 350MG BASE	N60385 001

NICOTINE

FILM, EXTENDED RELEASE; TRANSDERMAL

HABITROL

BC *	BASEL PHARMS	7MG/24HR	N20076 001
BC *		14MG/24HR	NOV 27, 1991
BC *		21MG/24HR	N20076 002
BC *			NOV 27, 1991
BC *			N20076 003
BC *			NOV 27, 1991
BC +	CIBA	7MG/24HR	N20076 001
BC +		14MG/24HR	NOV 27, 1991
BC +		21MG/24HR	N20076 002
			NOV 27, 1991
			N20076 003
			NOV 27, 1991

NICOTINE POLACRILEX

GUM, CHEWING; BUCCAL

NICORETTE

* MERRELL DOW EQ 2MG BASE

N18612 001

JAN 13, 1984

+ SMITHKLINE BEECHAM EQ 2MG BASE

N18612 001

JAN 13, 1984

NICORETTE DS

* MERRELL DOW EQ 4MG BASE

N20066 001

JUN 08, 1992

+ SMITHKLINE BEECHAM EQ 4MG BASE

N20066 001

JUN 08, 1992

NIFEDIPINE

TABLET, EXTENDED RELEASE; ORAL

ADALAT CC

BC + BAYER 30MG

N20198 001

APR 21, 1993

BC + 60MG

N20198 002

APR 21, 1993

BC + 90MG

N20198 003

APR 21, 1993

BC MILES 30MG

N20198 001

APR 21, 1993

BC 60MG

N20198 002

APR 21, 1993

BC 90MG

N20198 003

APR 21, 1993

NISOLDIPINE

TABLET, EXTENDED RELEASE; ORAL

NISOCOR

> DLT >

+ ZENECA 10MG

N20356 001

> DLT >

+ 20MG

N20356 002

> DLT >

+ 30MG

N20356 003

> DLT >

+ 40MG

N20356 004

> ADD >

SULAR

> ADD >

+ ZENECA 10MG

N20356 001

FEB 02, 1995

> ADD >

+ 20MG

N20356 002

FEB 02, 1995

> ADD >

+ 30MG

N20356 003

FEB 02, 1995

NISOLDIPINE

TABLET, EXTENDED RELEASE; ORAL

SULAR

> ADD >

> ADD >

> ADD >

+ ZENECA 40MG

N20356 004

FEB 02, 1995

NITROFURANTOIN, MACROCRYSTALLINE

CAPSULE; ORAL

NITROFURANTOIN

AB GENEVA PHARMS

25MG

N74336 001

JAN 25, 1995

AB 50MG

N74336 002

JAN 25, 1995

AB 100MG

N74336 003

JAN 25, 1995

NITROGLYCERIN

FILM, EXTENDED RELEASE; TRANSDERMAL

NITRO-DUR

+ KEY PHARMS 0.1MG/HR

N20145 001

APR 04, 1995

+ 0.2MG/HR

N20145 002

APR 04, 1995

+ 0.3MG/HR

N20145 003

APR 04, 1995

+ 0.4MG/HR

N20145 004

APR 04, 1995

+ 0.6MG/HR

N20145 005

APR 04, 1995

+ 0.8MG/HR

N20145 006

APR 04, 1995

INJECTABLE; INJECTION

NITROGLYCERIN

AP FUJISAWA

5MG/ML

N70077 001

DEC 13, 1985

@ 5MG/ML

N70077 001

DEC 13, 1985

NITROSTAT

AP PARKE DAVIS

5MG/ML

N18588 002

DEC 23, 1983

* 0.8MG/ML

N18588 001

@ 0.8MG/ML

N18588 001

NITROGLYCERIN

INJECTABLE; INJECTION

NITROSTAT
@ PARKE DAVIS

5MG/ML

N18588 002
DEC 23, 1983TRIDILAP DUPONT MERCK5MG/MLN18537 001

*

0.5MG/MLN18537 002
JUN 16, 1983AP FAULDING5MG/MLN18537 001

+

0.5MG/MLN18537 002
JUN 16, 1983NORETHINDRONE

TABLET; ORAL

NOR-Q.D.

SEARLE
SYNTEX0.35MG
0.35MGN17060 001
N17060 001NORTRIPTYLINE HYDROCHLORIDE

CAPSULE; ORAL

NORTRIPTYLINE HCLAB LEMMONEQ 10MG BASEN74132 001
MAR 27, 1995ABEQ 25MG BASEN74132 002
MAR 27, 1995ABEQ 50MG BASEN74132 003
MAR 27, 1995ABEQ 75MG BASEN74132 004
MAR 27, 1995NYSTATIN

TABLET; ORAL

MYCOSTATINAA + APOTHECON500,000 UNITSN60574 001AA * SQUIBB500,000 UNITSN60574 001

TABLET; VAGINAL

NYSTATINAT LEMMON100,000 UNITSN62502 001
DEC 23, 1983NYSTATIN

TABLET; VAGINAL

NYSTATIN
@ LEMMON

100,000 UNITS

N62502 001
DEC 23, 1983NYSTATIN; TRIAMCINOLONE ACETONIDE

OINTMENT; TOPICAL

NYSTATIN AND TRIAMCINOLONE ACETONIDEAT PHARMAFAIR100,000 UNITS/GM; 0.1%N62656 001

@

100,000 UNITS/GM; 0.1%

JUL 30, 1986
N62656 001
JUL 30, 1986OMEPRAZOLE

CAPSULE, DELAYED REL PELLETS; ORAL

PRILOSEC

+ ASTRA MERCK

10MG

N19810 003
OCT 05, 1995ONDANSETRON HYDROCHLORIDE

INJECTABLE; INJECTION

ZOFRAN IN PLASTIC CONTAINER

+ GLAXO WELLCOME

EQ 0.64MG BASE/ML

N20403 001
JAN 31, 1995OXACILLIN SODIUM

CAPSULE; ORAL

OXACILLIN SODIUMAB APOTHECONEQ 250MG BASEN61450 002AB +EQ 500MG BASEN61450 001AB PROSTAPHLINEQ 250MG BASEN61450 002AB APOTHECONEQ 500MG BASEN61450 001AB *EQ 500MG BASEN50118 002

@

EQ 500MG BASEN50118 002

@ BRISTOL

INJECTABLE; INJECTION

OXACILLIN SODIUMAP IBIEQ 250MG BASE/VIALN62798 004
DEC 11, 1995> ADD >
> ADD >

OXACILLIN SODIUM

INJECTABLE; INJECTION

> <u>ADD</u> >	<u>AP</u>	<u>IBI</u>	<u>EQ 500MG BASE/VIAL</u>	N62798 005
> <u>ADD</u> >				DEC 11, 1995
> <u>ADD</u> >	<u>AP</u>		<u>EQ 1GM BASE/VIAL</u>	N62798 001
> <u>ADD</u> >				DEC 11, 1995
> <u>ADD</u> >	<u>AP</u>		<u>EQ 2GM BASE/VIAL</u>	N62798 002
> <u>ADD</u> >				DEC 11, 1995
> <u>ADD</u> >			<u>EQ 125MG BASE/VIAL</u>	N62798 003
> <u>ADD</u> >				DEC 11, 1995

POWDER FOR RECONSTITUTION; ORAL

<u>AA</u>	<u>APOTHECON</u>	<u>EQ 250MG BASE/5ML</u>	N61457 001
<u>AA</u>	<u>PROSTAPHLIN</u>		
<u>AA</u>	<u>APOTHECON</u>	<u>EQ 250MG BASE/5ML</u>	N61457 001
	@	<u>EQ 250MG BASE/5ML</u>	N50194 001
	@	<u>BRISTOL</u>	N50194 001

> ADD > OXYCODONE HYDROCHLORIDE

> <u>ADD</u> >	TABLET, EXTENDED RELEASE; ORAL		
> <u>ADD</u> >	OXYCONTIN		
> <u>ADD</u> >	+	PURDUE FREDERICK	10MG
> <u>ADD</u> >			N20553 001
> <u>ADD</u> >			DEC 12, 1995
> <u>ADD</u> >	+		20MG
> <u>ADD</u> >			N20553 002
> <u>ADD</u> >			DEC 12, 1995
> <u>ADD</u> >	+		40MG
> <u>ADD</u> >			N20553 003
> <u>ADD</u> >			DEC 12, 1995

OXYPHENCYCLIMINE HYDROCHLORIDE

TABLET; ORAL

	<u>DARICON</u>		
*	<u>PFIZER</u>	10MG	N11612 001
@		10MG	N11612 001

OXYTETRACYCLINE HYDROCHLORIDE

CAPSULE; ORAL

> <u>DLT</u> >	<u>OXY-KESSO-TETRA</u>		
> <u>ADD</u> >	@	FERRANTE	EQ 250MG BASE
> <u>DLT</u> >	<u>AB</u>	MK LABS	EQ 250MG BASE
			N60179 001
			N60179 001

PENBUTOLOL SULFATE

TABLET; ORAL

	<u>LEVATOL</u>		
@	<u>REED AND CARNICK</u>	10MG	N18976 001
			DEC 30, 1987
*		20MG	N18976 004
			JAN 05, 1989
@	<u>SPKU</u>	10MG	N18976 001
			DEC 30, 1987
+		20MG	N18976 004
			JAN 05, 1989

PENICILLAMINE

TABLET; ORAL

	<u>DEPEN</u>		
+	<u>WALLACE</u>	250MG	N19854 001
	<u>DEPEN 250</u>		
+	<u>WALLACE</u>	250MG	N19854 001

PENICILLIN G POTASSIUM

INJECTABLE; INJECTION

	<u>PENICILLIN G POTASSIUM</u>		
@	<u>CONSOLIDATED PHARM</u>	500,000 UNITS/VIAL	N60806 001
@		1,000,000 UNITS/VIAL	N60806 002
@		5,000,000 UNITS/VIAL	N60806 003
@		10,000,000 UNITS/VIAL	N60806 004
@	<u>COPANOS</u>	500,000 UNITS/VIAL	N60806 001
@		1,000,000 UNITS/VIAL	N60806 002
@		5,000,000 UNITS/VIAL	N60806 003
@		10,000,000 UNITS/VIAL	N60806 004
<u>AP</u>	+	<u>LILLY</u>	1,000,000 UNITS/VIAL
<u>AP</u>	+		5,000,000 UNITS/VIAL
<u>AP</u>	+		20,000,000 UNITS/VIAL
<u>AP</u>	+		20,000,000 UNITS/VIAL
			N60601 001
	+		200,000 UNITS/VIAL
	+		500,000 UNITS/VIAL
@		200,000 UNITS/VIAL	N60384 004
@		500,000 UNITS/VIAL	N60384 003
@		1,000,000 UNITS/VIAL	N60384 002
@		5,000,000 UNITS/VIAL	N60384 001
@		20,000,000 UNITS/VIAL	N60384 005
@		20,000,000 UNITS/VIAL	N60601 001
<u>AP</u>	<u>PFIZERPEN</u>		
<u>AP</u>	<u>PFIZER</u>	1,000,000 UNITS/VIAL	N60657 001

PENICILLIN G POTASSIUM

INJECTABLE; INJECTION

PFIZERPEN

<u>AP</u>	+	PFIZER	<u>1,000,000 UNITS/VIAL</u>	N60657 001
<u>AP</u>			<u>5,000,000 UNITS/VIAL</u>	N60657 002
<u>AP</u>	+		<u>5,000,000 UNITS/VIAL</u>	N60657 002
<u>AP</u>			<u>20,000,000 UNITS/VIAL</u>	N60657 003
<u>AP</u>	+		<u>20,000,000 UNITS/VIAL</u>	N60657 003

TABLET; ORAL

PENICILLIN G POTASSIUM

<u>AB</u>		<u>DISTA</u>	<u>250,000 UNITS</u>	N60403 001
		@ LILLY	250,000 UNITS	N60403 001

PENICILLIN G PROCAINE

INJECTABLE; INJECTION

PENICILLIN G PROCAINE

@ CONSOLIDATED PHARM

@

@ COPANOS

@

300,000 UNITS/ML	N60800 001
600,000 UNITS/1.2ML	N60800 002
300,000 UNITS/ML	N60800 001
600,000 UNITS/1.2ML	N60800 002

PENICILLIN V POTASSIUM

POWDER FOR RECONSTITUTION; ORAL

PENICILLIN V POTASSIUM

<u>AA</u>		CONSOLIDATED PHARM	<u>EQ 125MG BASE/5ML</u>	N61529 001
<u>AA</u>			<u>EQ 250MG BASE/5ML</u>	N61529 002
<u>AA</u>		COPANOS	<u>EQ 125MG BASE/5ML</u>	N61529 001
<u>AA</u>			<u>EQ 250MG BASE/5ML</u>	N61529 002

TABLET; ORAL

BETAPEN-VK

<u>AB</u>		APOTHECON	<u>EQ 250MG BASE</u>	N61411 001
<u>AB</u>			<u>EQ 500MG BASE</u>	N61411 002

PENICILLIN V POTASSIUM

<u>AB</u>		BIOCHEMIE	<u>EQ 250MG BASE</u>	N64071 001
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<u>AB</u>			<u>EQ 500MG BASE</u>	N64071 002
-----------	--	--	----------------------	------------

<u>AB</u>		CONSOLIDATED PHARM	<u>EQ 250MG BASE</u>	N61528 001
-----------	--	--------------------	----------------------	------------

<u>AB</u>			<u>EQ 500MG BASE</u>	N61528 002
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<u>AB</u>		COPANOS	<u>EQ 250MG BASE</u>	N61528 001
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<u>AB</u>			<u>EQ 500MG BASE</u>	N61528 002
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VEETIDS

<u>AB</u>		APOTHECON	<u>EQ 250MG BASE</u>	N61411 001
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PENICILLIN V POTASSIUM

TABLET; ORAL

VEETIDS

<u>AB</u>		APOTHECON	<u>EQ 500MG BASE</u>	N61411 002
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PENTAMIDINE ISETHIONATE

INJECTABLE; INJECTION

PENTACARINAT

<u>AP</u>		ARMOUR	<u>300MG/VIAL</u>	N73447 001
-----------	--	--------	-------------------	------------

<u>AP</u>		RHONE-POULENC RORER	<u>300MG/VIAL</u>	N73447 001
-----------	--	---------------------	-------------------	------------

				APR 28, 1994
--	--	--	--	--------------

				APR 28, 1994
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PENTAMIDINE ISETHIONATE

> ADD >	<u>AP</u>	ELKINS SINN	<u>300MG/VIAL</u>	N73617 001
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> ADD >	<u>AP</u>		<u>300MG/VIAL</u>	DEC 18, 1995
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<u>AP</u>		STERIS	<u>300MG/VIAL</u>	N74303 001
-----------	--	--------	-------------------	------------

				AUG 17, 1995
--	--	--	--	--------------

PERINDOPRIL ERBUMINE

TABLET; ORAL

ACEON

AMARIC

2MG

N20184 001

4MG

DEC 30, 1993

8MG

N20184 002

DEC 30, 1993

JOHNSON RW

2MG

N20184 003

DEC 30, 1993

4MG

N20184 001

DEC 30, 1993

8MG

N20184 002

DEC 30, 1993

N20184 003

DEC 30, 1993

PHENDIMETRAZINE TARTRATE

CAPSULE, EXTENDED RELEASE; ORAL

MELFIAT-105

@ NUMARK

105MG

N87487 001

@ SOLVAY

105MG

OCT 13, 1982

N87487 001

OCT 13, 1982

PHENDIMETRAZINE TARTRATE

CAPSULE, EXTENDED RELEASE; ORAL

SPRX-105

@ NUMARK 105MG

@ SOLVAY 105MG

N88024 001

DEC 22, 1982

N88024 001

DEC 22, 1982

TABLET; ORAL

MELFIAT

@ NUMARK 35MG

@ SOLVAY 35MG

N83790 002

N83790 002

PHENDIMETRAZINE TARTRATE

@ NUMARK 35MG

@ SOLVAY 35MG

N83790 001

N83790 001

PHENTERMINE HYDROCHLORIDE

CAPSULE; ORAL

PHENTERMINE HCL

AA COMMON 30MG

@ 30MG

N87777 001

NOV 01, 1985

N87777 001

NOV 01, 1985

PHENTERMINE RESIN COMPLEX

CAPSULE, EXTENDED RELEASE; ORAL

IONAMIN

FISONS EQ 15MG BASE

+ EQ 30MG BASE

N11613 004

N11613 002

IONAMIN-15

FISONS EQ 15MG BASE

N11613 004

IONAMIN-30

+ FISONS EQ 30MG BASE

N11613 002

PINACIDIL

CAPSULE, EXTENDED RELEASE; ORAL

PINDAC

+ LEO PHARM 12.5MG

+ 25MG

N19456 001

DEC 28, 1989

N19456 002

DEC 28, 1989

PINACIDIL

CAPSULE, EXTENDED RELEASE; ORAL

PINDAC

@ LEO PHARM 12.5MG

@ 25MG

N19456 001

DEC 28, 1989

N19456 002

DEC 28, 1989

PINDOLOL

TABLET; ORAL

PINDOLOL

AB ROYCE LABS 5MG

AB 10MG

N74437 001

FEB 27, 1995

N74437 002

FEB 27, 1995

PIROXICAM

CAPSULE; ORAL

PIROXICAM

AB ROYCE LABS 10MG

AB 20MG

N74460 001

SEP 29, 1995

N74460 002

SEP 29, 1995

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

POWDER FOR RECONSTITUTION; ORAL

NULYTELY-FLAVORED

BRAINTREE

420GM/BOT; 1.48GM/BOT; 5.72GM/BOT;

11.2GM/BOT

N19797 002

NOV 18, 1994

POLYETHYLENE GLYCOL 3350; POTASSIUM CHLORIDE; SODIUM BICARBONATE; SODIUM CHLORIDE; SODIUM SULFATE, ANHYDROUS

POWDER FOR RECONSTITUTION; ORAL

COLYTE

AA KREMERS URBAN

227.1GM/BOT; 2.82GM/BOT; 6.36GM/BOT;

5.53GM/BOT; 21.5GM/BOT

N18983 010

JAN 31, 1989

POLYETHYLENE GLYCOL 3350; POTASSIUM CHLORIDE; SODIUM
BICARBONATE; SODIUM CHLORIDE; SODIUM SULFATE, ANHYDROUS

POWDER FOR RECONSTITUTION; ORAL

COLYTE

AA	KREMERS URBAN	240GM/BOT; 2.98GM/BOT; 6.72GM/BOT; 5.84GM/BOT; 22.72GM/BOT	N18983 007
		JUN 12, 1987	
@	REED AND CARRICK	120GM/PACKET; 1.49GM/PACKET; 3.36GM/PACKET; 2.92GM/PACKET; 11.36GM/PACKET	N18983 005
		OCT 26, 1984	
@		227.1GM/PACKET; 2.82GM/PACKET; 6.36GM/PACKET; 5.53GM/PACKET; 21.5GM/PACKET	N18983 004
		OCT 26, 1984	
@		360GM/PACKET; 4.47GM/PACKET; 10.08GM/PACKET; 8.76GM/PACKET; 34.08GM/PACKET	N18983 006
		OCT 26, 1984	
AA	SPKU	227.1GM/BOT; 2.82GM/BOT; 6.36GM/BOT; 5.53GM/BOT; 21.5GM/BOT	N18983 010
		JAN 31, 1989	
AA		240GM/BOT; 2.98GM/BOT; 6.72GM/BOT; 5.84GM/BOT; 22.72GM/BOT	N18983 007
		JUN 12, 1987	
@		120GM/PACKET; 1.49GM/PACKET; 3.36GM/PACKET; 2.92GM/PACKET; 11.36GM/PACKET	N18983 005
		OCT 26, 1984	
@		227.1GM/PACKET; 2.82GM/PACKET; 6.36GM/PACKET; 5.53GM/PACKET; 21.5GM/PACKET	N18983 004
		OCT 26, 1984	
@		360GM/PACKET; 4.47GM/PACKET; 10.08GM/PACKET; 8.76GM/PACKET; 34.08GM/PACKET	N18983 006
		OCT 26, 1984	
AA	<u>COLYTE-FLAVORED</u> KREMERS URBAN	227.1GM/BOT; 2.82GM/BOT; 6.36GM/BOT; 5.53GM/BOT; 21.5GM/BOT	N18983 008
		NOV 14, 1991	
AA		240GM/BOT; 2.98GM/BOT; 6.72GM/BOT; 5.84GM/BOT; 22.72GM/BOT	N18983 009
		NOV 14, 1991	
AA	SPKU	227.1GM/BOT; 2.82GM/BOT; 6.36GM/BOT; 5.53GM/BOT; 21.5GM/BOT	N18983 008
		NOV 14, 1991	

POLYETHYLENE GLYCOL 3350; POTASSIUM CHLORIDE; SODIUM
BICARBONATE; SODIUM CHLORIDE; SODIUM SULFATE, ANHYDROUS

POWDER FOR RECONSTITUTION; ORAL

COLYTE-FLAVORED

AA	SPKU	240GM/BOT; 2.98GM/BOT; 6.72GM/BOT; 5.84GM/BOT; 22.72GM/BOT	N18983 009
		NOV 14, 1991	
AA	<u>GOLYTELY</u> BRAINTREE	227.1GM/PACKET; 2.82GM/PACKET; 6.36GM/PACKET; 5.53GM/PACKET; 21.5GM/PACKET	N19011 002
		JUN 02, 1992	

POLYTHIAZIDE; PRAZOSIN HYDROCHLORIDE

CAPSULE; ORAL

MINIZIDE

PFIZER

	0.5MG; 1MG	N17986 001
	0.5MG; 2MG	N17986 002
	0.5MG; 5MG	N17986 003
+	0.5MG; EQ 1MG BASE	N17986 001
	0.5MG; EQ 2MG BASE	N17986 002
+	0.5MG; EQ 5MG BASE	N17986 003

> ADD > PORFIMER SODIUM

> ADD > INJECTABLE; INJECTION

> ADD > PHOTOFRIN

> ADD >	+ QLT	75MG/VIAL	N20451 001
> ADD >			DEC 27, 1995

POTASSIUM CHLORIDE

INJECTABLE; INJECTION

POTASSIUM CHLORIDE

AP	AKORN	2MEQ/ML	N88286 001
			SEP 05, 1985
@		2MEQ/ML	N88286 001
			SEP 05, 1985

TABLET, EXTENDED RELEASE; ORAL

KAON CL

SAVAGE LABS

	SAVAGE LABS	6.7MEQ	N17046 001
@		6.7MEQ	N17046 001
	KAON CL-10		
BC	SAVAGE LABS	10MEQ	N17046 002

POTASSIUM CHLORIDE

TABLET, EXTENDED RELEASE; ORAL
KACON CL-10
@ SAVAGE LABS 10MEQ

N17046 002 > ADD >
> ADD >

PREDNISOLONE

TABLET; ORAL
CORTALONE

BX HALSEY 1MG
BX 2.5MG
BX 5MG
@ 1MG
@ 2.5MG
@ 5MG
PREDNISOLONE
@ FERRANTE 2.5MG
@ 5MG
BX MARSHALL PHARMA 2.5MG
BX 5MG
BX SPERTI 1MG
1MG

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

N80304 003
N80304 002
N80304 001
N80304 003
N80304 002
N80304 001
N80562 001
N80562 002
N80562 001
N80562 002
N80358 001
N80358 001

PREDNISOLONE ACETATE

SUSPENSION/DROPS; OPHTHALMIC
ECONOPRED PLUS
ALCON 1%
EX 1%

N17469 001
N17469 001

PREDNISOLONE SODIUM PHOSPHATE

INJECTABLE; INJECTION
HYDELTRASOL

AP + MERCK SHARP DOHME EQ 20MG PHOSPHATE/ML
+ EQ 20MG PHOSPHATE/ML
PREDNISOLONE SODIUM PHOSPHATE
AP STERIS EQ 20MG PHOSPHATE/ML
@ EQ 20MG PHOSPHATE/ML

N11583 002
N11583 002
N80517 001
N80517 001

PREDNISOLONE SODIUM PHOSPHATE; SULFACETAMIDE SODIUM

SOLUTION/DROPS; OPHTHALMIC
SULFACETAMIDE SODIUM AND PREDNISOLONE SODIUM PHOSPHATE
BAUSCH AND LOMB EQ 0.23% PHOSPHATE;10% N74449 001
DEC 29, 1995

PREDNISONE

TABLET; ORAL

CORTAN
BX HALSEY 20MG
@ 20MG N87480 001
N87480 001

PROBUCOL

TABLET; ORAL

LORELCO
@ HOECHST MARION RSSL 250MG
@ 500MG N17535 001
N17535 002
JUL 06, 1988
MERRELL DOW 250MG
+ 500MG N17535 001
N17535 002
JUL 06, 1988

PROCAINAMIDE HYDROCHLORIDE

TABLET; ORAL

PRONESTYL
APOTHECON 250MG
375MG N17371 001
500MG N17371 002
+ 500MG N17371 003
SQUIBB 250MG
375MG N17371 001
500MG N17371 002
500MG N17371 003

TABLET, EXTENDED RELEASE; ORAL

PROCAN SR
AP PARKE DAVIS 250MG
@ 250MG N86468 001
N86468 001
PRONESTYL-SR
BC APOTHECON 500MG
BC BRISTOL MYERS SQUIBB 500MG
N87361 001
N87361 001

PROMETHAZINE HYDROCHLORIDE

INJECTABLE; INJECTION

PROMETHAZINE HCL

<u>AP</u>	<u>AKORN</u>	<u>25MG/ML</u>
<u>AP</u>		<u>50MG/ML</u>
	@	25MG/ML
	@	50MG/ML

N83955 002	> <u>ADD</u> >
N83955 001	> <u>ADD</u> >
N83955 002	> <u>ADD</u> >
N83955 001	> <u>ADD</u> >

PROPANTHELINE BROMIDE

TABLET; ORAL

PROPANTHELINE BROMIDE

<u>AA</u>	<u>DANBURY PHARMA</u>	<u>15MG</u>
	@	15MG
<u>AA</u>	<u>GLOBAL PHARMS</u>	<u>15MG</u>
	@	15MG
<u>AA</u>	<u>TABLICAPS</u>	<u>15MG</u>
	@	15MG

N83029 002
N83029 002
N84541 002
N84541 002
N84428 001
N84428 001

PROPACARCAINE HYDROCHLORIDE

SOLUTION/DROPS; OPHTHALMIC

OPHTHAINE

<u>AT</u>	<u>+ APOTHECON</u>	<u>0.5%</u>
<u>AT</u>	<u>* SQUIBB</u>	<u>0.5%</u>
	<u>PROPARACCAINE HCL</u>	
<u>AT</u>	<u>BAUSCH AND LOMB</u>	<u>0.5%</u>

N08883 001
N08883 001
N40074 001
SEP 29, 1995

PROPRANOLOL HYDROCHLORIDE

TABLET; ORAL

PROPRANOLOL HCL

> <u>DLT</u> >	<u>AB</u>	<u>INTERPHARM</u>	<u>10MG</u>
> <u>DLT</u> >	<u>AB</u>		<u>20MG</u>
> <u>DLT</u> >	<u>AB</u>		<u>40MG</u>
> <u>DLT</u> >	<u>AB</u>		<u>80MG</u>
> <u>ADD</u> >	@		10MG
> <u>ADD</u> >	@		20MG

N71368 001
MAY 05, 1987
N71369 001
MAY 05, 1987
N71370 001
MAY 05, 1987
N71371 001
MAY 05, 1987
N71368 001
MAY 05, 1987
N71369 001
MAY 05, 1987

PROPRANOLOL HYDROCHLORIDE

TABLET; ORAL

PROPRANOLOL HCL

@	<u>INTERPHARM</u>	40MG
@		80MG
<u>AB</u>	<u>PARKE DAVIS</u>	<u>20MG</u>
<u>AB</u>		<u>40MG</u>
<u>AB</u>		<u>60MG</u>
<u>AB</u>		<u>80MG</u>
<u>AB</u>	<u>WARNER CHILCOTT</u>	<u>10MG</u>
@		10MG
@		20MG
@		40MG
@		60MG
@		80MG

N71370 001
MAY 05, 1987
N71371 001
MAY 05, 1987
N70439 001
SEP 15, 1986
N70440 001
SEP 15, 1986
N70441 001
SEP 24, 1986
N70442 001
SEP 15, 1986
N70438 001
SEP 15, 1986
N70439 001
SEP 15, 1986
N70440 001
SEP 15, 1986
N70441 001
SEP 24, 1986
N70442 001
SEP 15, 1986

PROPYLTHIOURACIL

TABLET; ORAL

PROPYLTHIOURACIL

<u>BD</u>	<u>LILLY</u>	<u>50MG</u>
@		50MG

N06213 001
N06213 001

PROTRIPTYLINE HYDROCHLORIDE

TABLET; ORAL

PROTRIPTYLINE HCL

<u>AB</u>	<u>SIDMAK LABS NJ</u>	<u>5MG</u>
<u>AB</u>		<u>10MG</u>
<u>AB</u>	<u>VIVACTIL</u>	<u>5MG</u>
<u>AB</u>	<u>+ MERCK SHARP DOHME</u>	<u>10MG</u>

N73644 001
AUG 24, 1995
N73645 001
AUG 24, 1995
N16012 001
N16012 002

PYRIDOXINE HYDROCHLORIDE

INJECTABLE; INJECTION
PYRIDOXINE HCL
 AP AKORN 100MG/ML
 @ 100MG/ML

N87967 001
 OCT 01, 1982
 N87967 001
 OCT 01, 1982

RAUWOLFIA SERPENTINA

TABLET; ORAL
 RAUWOLFIA SERPENTINA
 BP HALSEY 100MG
 @ 50MG
 @ 100MG

N80498 002
 N80498 001
 N80498 002

QUINESTROL

TABLET; ORAL
 ESTROVIS
 * PARKE DAVIS 0.1MG
 @ 0.1MG

N16768 002
 N16768 002

> ADD >

RILUZOLE

> ADD >

TABLET; ORAL
 RILUTEK
 + RHONE POULENC 50MG

> ADD >

> ADD >

> ADD >

N20599 001
 DEC 12, 1995

QUINIDINE GLUCONATE

TABLET, EXTENDED RELEASE; ORAL
 QUINALAN
 BC LANNETT 324MG
 @ 324MG

N88081 001
 FEB 10, 1986
 N88081 001
 FEB 10, 1986

> ADD >

RITODRINE HYDROCHLORIDE

TABLET; ORAL
 YUTOPAR
 * ASTRA 10MG
 @ 10MG

N18555 001
 N18555 001

QUINIDINE SULFATE

TABLET; ORAL
QUINIDINE SULFATE
 AB PHOENIX LABS NY 200MG
 @ VINTAGE PHARMS 200MG

N83963 001
 N83963 001

> ADD >

> ADD >

> ADD >

> ADD >

SAQUINAVIR MESYLATE

CAPSULE; ORAL
 INVIRASE
 + ROCHE EQ 200MG BASE

N20628 001
 DEC 06, 1995

SECOBARBITAL SODIUM

CAPSULE; ORAL
SECOBARBITAL SODIUM
 AA ZENITH LABS 100MG
 @ 100MG

N85869 001
 N85869 001

RAUWOLFIA SERPENTINA

TABLET; ORAL
 RAUDIXIN
 BP APOTHECON 50MG
 BP 100MG
 @ 50MG
 @ 100MG
 RAUVAL
 BP VALE 100MG
 BP + 100MG
 RAUWOLFIA SERPENTINA
 BP HALSEY 50MG

N08842 001
 N08842 002
 N08842 001
 N08842 002

N09108 004
 N09108 004

N80498 001

SEVOFLURANE

LIQUID; INHALATION
 ULTANE
 ABBOTT 100%

N20478 001
 JUN 07, 1995

SILVER SULFADIAZINE

CREAM; TOPICAL
SSD AF
BX KNOLL PHARM 1%

N18578 003
JUL 11, 1990

SODIUM CHLORIDE

INJECTABLE; INJECTION
SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER
AP BAXTER 9MG/ML

N16677 004
OCT 30, 1985
N16677 004
OCT 30, 1985

AP + 9MG/ML

SOLUTION; IRRIGATION
SODIUM CHLORIDE 0.45% IN PLASTIC CONTAINER
AT BAXTER 450MG/100ML

N18497 001
FEB 19, 1982
N18497 001
FEB 19, 1982

@ 450MG/100ML

SODIUM CHROMATE, CR-51

INJECTABLE; INJECTION
CHROMITOPHE SODIUM

BRACCO 200 uCi/ML
BRACCO DXS 250 uCi/VIAL
1mCi/VIAL

N13993 001
N13993 001
N13993 001

SOMATROPIN, BIOSYNTHETIC

INJECTABLE; INJECTION
BIO-TROPIN

+ BIO TECH GEN 4.8MG/VIAL

N19774 001
MAY 25, 1995

GENOTROPIN

+ PHARMACIA 1.5MG/VIAL

N20280 004
AUG 24, 1995

+ 5.8MG/VIAL

N20280 006
AUG 24, 1995

NORDITROPIN

+ NOVO NORDISK 4MG/VIAL

N19721 001
MAY 08, 1995

SOMATROPIN, BIOSYNTHETIC

INJECTABLE; INJECTION
NORDITROPIN

+ NOVO NORDISK 8MG/VIAL

N19721 002
MAY 08, 1995

> ADD >

> ADD >

> ADD >

NUTROPIN AQ

+ GENENTECH 5MG/ML

N20522 001
DEC 29, 1995

SOTALOL HYDROCHLORIDE

TABLET; ORAL

BETAPACE

BERLEX

120MG

N19865 005
APR 20, 1994

@

120MG

N19865 005
APR 20, 1994

STREPTOMYCIN SULFATE

INJECTABLE; INJECTION
STREPTOMYCIN SULFATE

AP LILLY
AP

EQ 1GM BASE/VIAL

N60107 001

EQ 5GM BASE/VIAL

N60107 002

EQ 1GM BASE/2ML

N60404 001

EQ 1GM BASE/2ML

N60404 001

EQ 1GM BASE/VIAL

N60107 001

EQ 5GM BASE/VIAL

N60107 002

EQ 1GM BASE/VIAL

N60076 001

EQ 5GM BASE/VIAL

N60076 002

EQ 1GM BASE/VIAL

N60076 001

EQ 5GM BASE/VIAL

N60076 002

SUCCIMER

CAPSULE; ORAL
CHEMET

+ BOCK PHARMA 100MG

N19998 002
JAN 30, 1991

+ MCNEIL CONS PRODS 100MG

N19998 002
JAN 30, 1991

SUCCINYLCHOLINE CHLORIDE

INJECTABLE; INJECTION

<u>AP</u>	<u>SUCOSTRIN</u>		
	APOTHECON	<u>20MG/ML</u>	N08847 001
	@	100MG/ML	N08847 003
<u>AP</u>	<u>SQUIBB</u>	<u>20MG/ML</u>	N08847 001
	@	100MG/ML	N08847 003

SUFENTANIL CITRATE

INJECTABLE; INJECTION

> ADD >	<u>AP</u>	+	JANSSEN	<u>EQ 0.05MG BASE/ML</u>	N19050 001
> ADD >					MAY 04, 1984
> ADD >			<u>SUFENTANIL CITRATE</u>		
> ADD >	<u>AP</u>		ELKINS SINN	<u>EQ 0.05MG BASE/ML</u>	N74413 001
> ADD >					DEC 15, 1995
> ADD >	<u>AP</u>		STERIS	<u>EQ 0.05MG BASE/ML</u>	N74406 001
> ADD >					DEC 15, 1995

SULFACETAMIDE SODIUM

SOLUTION/DROPS; OPHTHALMIC

<u>AT</u>	<u>SULF-15</u>		
	CIBA	<u>15%</u>	N89047 001
			OCT 31, 1995

SULFAMETHOXAZOLE; TRIMETHOPRIM

SUSPENSION; ORAL

<u>AP</u>	<u>TRIMETH/SULFA</u>		
	BARRE	<u>200MG/5ML; 40MG/5ML</u>	N72289 001
			MAY 23, 1988
	@	200MG/5ML; 40MG/5ML	N72289 001
			MAY 23, 1988

SULFUR

POWDER; TOPICAL

	BENSULFOID		
	@ POYTHRESS	33.32%	N02918 001

SUMATRIPTAN SUCCINATE

TABLET; ORAL

	IMITREX		
	GLAXO WELLCOME	EQ 25MG BASE	N20132 002
			JUN 01, 1995
+		EQ 50MG BASE	N20132 003
			JUN 01, 1995
@		EQ 100MG BASE	N20132 001
			JUN 01, 1995

TECHNETIUM TC-99M SODIUM PERTECHNETATE GENERATOR

SOLUTION; INJECTION, ORAL

	TECHNELITE		
	DUPONT	0.0083-2.7 CI/GENERATOR	N17771 001
	TECHNETIUM TC-99M GENERATOR		
	DUPONT	0.0083-2.7 CI/GENERATOR	N17771 001

TESTOSTERONE

FILM, EXTENDED RELEASE; TRANSDERMAL

	ANDRODERM		
+	THERATECH	2.5MG/24HR	N20489 001
			SEP 29, 1995

TETRACYCLINE HYDROCHLORIDE

CAPSULE; ORAL

TETRACYCLINE HCL

> ADD >	@ FERRANTE	125MG	N60173 001
> ADD >	@	250MG	N60173 002
> DLT >	<u>AB</u>	<u>250MG</u>	N60173 002
> DLT >		125MG	N60173 001
	<u>AB</u>	<u>250MG</u>	N62686 001
			JUL 24, 1986
	<u>AB</u>	<u>500MG</u>	N62686 002
			JUL 24, 1986
	@	250MG	N62686 001
			JUL 24, 1986
	@	500MG	N62686 002
			JUL 24, 1986

FIBER, EXTENDED RELEASE; PERIODONTAL

	ACTISITE		
+	ON SITE	12.7MG/FIBER	N50653 001
			MAR 25, 1994

TETRACYCLINE HYDROCHLORIDE

FIBER, EXTENDED RELEASE; PERIODONTAL
ACTISITE
+ ON SITE ALZA 12.7MG/FIBER

N50653 001
MAR 25, 1994

ointment, Ophthalmic

ACHROMYCIN
* LEDERLE 10MG/GM
@ STORZ OPHTHALM 10MG/GM

N50266 001
N50266 001

SUSPENSION; ORAL
ACHROMYCIN V

AB + LEDERLE 125MG/5ML

N50263 002

AB SUMYCIN
APOTHECON 125MG/5ML

N60400 001

AB TETRACYCLINE HCL
BARRE 125MG/5ML

N60633 001

@ FERRANTE 125MG/5ML

N60174 001

AB MR LABS 125MG/5ML

N60174 001

@ PROTER 125MG/5ML

N60446 001

AB PUREPAC PHARM 125MG/5ML

N60291 001

AB TETRACYN
PFIPHARMECS 125MG/5ML

N60095 001

AB TETRAMED
ZENITH LABS 125MG/5ML

N61468 001

SUSPENSION/DROPS; Ophthalmic

ACHROMYCIN

* LEDERLE 1%

@ STORZ OPHTHALM 1%

N50268 001

N50268 001

Syrup; Oral

AB ACHROMYCIN V
* LEDERLE 125MG/5ML

N50263 002

AB SUMYCIN
SQUIBB 125MG/5ML

N60400 001

AB TETRACYCLINE HCL
BARRE 125MG/5ML

N60633 001

AB PUREPAC PHARM 125MG/5ML

N60291 001

AB TETRACYN
PFIPHARMECS 125MG/5ML

N60095 001

AB TETRAMED
ZENITH LABS 125MG/5ML

N61468 001

THEOPHYLLINE

CAPSULE, EXTENDED RELEASE; ORAL

THEOPHYLLINE
BC FAULDING 100MG

N89976 001

BC 200MG

JAN 04, 1995

BC 300MG

N89977 001

JAN 04, 1995

N89932 001

JAN 04, 1995

TABLET, EXTENDED RELEASE; ORAL

LABID
BC * PROCTER AND GAMBLE 250MG

N87225 001

@ 250MG

N87225 001

BC THEOLAIR-SR 250MG

N86363 002

JUL 16, 1987

BC 250MG

N86363 002

JUL 16, 1987

AB THEOPHYLLINE
INWOOD LABS 450MG

N40034 001

APR 28, 1995

UNI-DUR
BC + KEY PHARMS 400MG

N89822 001

JAN 04, 1995

+ 600MG

N89823 001

JAN 04, 1995

UNIPHYL
BC PURDUE FREDERICK 400MG

N87571 001

SEP 01, 1982

THEOPHYLLINE SODIUM GLYCINATE

TABLET, ORAL

ASERON
* DORSEY EQ 150MG BASE

N85148 001

@ SANDOZ EQ 150MG BASE

N85148 001

THIAMINE HYDROCHLORIDE

INJECTABLE; INJECTION

THIAMINE HCL
AP AKORN 100MG/ML

N87968 001

OCT 01, 1982

@ 100MG/ML

N87968 001

OCT 01, 1982

> ADD >
> DLT >

THIOTEPA

INJECTABLE; INJECTION

THIOPLEX

IMMUNEX

15MG/VIAL

N20058 001

DEC 22, 1994

AP LEADERLE

15MG/VIAL

N20058 001

DEC 22, 1994

AP * THIOTEPA

IMMUNEX

15MG/VIAL

N11683 001

+

15MG/VIAL

N11683 001

THIOTHIXENE HYDROCHLORIDE

CONCENTRATE; ORAL

THIOTHIXENE HCL

LEMMON

EQ 5MG BASE/ML

N71184 001

JUN 22, 1987

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

@

EQ 5MG BASE/ML

N71184 001

JUN 22, 1987

TICARCILLIN DISODIUM

INJECTABLE; INJECTION

TICAR

* SMITHKLINE BEECHAM

EQ 6GM BASE/VIAL

N50497 003

@

EQ 6GM BASE/VIAL

N50497 003

TIMOLOL

SOLUTION/DROPS; OPHTHALMIC

BETIMOL

+ LEIRAS

EQ 0.25% BASE

N20439 001

MAR 31, 1995

+

EQ 0.5% BASE

N20439 002

MAR 31, 1995

> DLT >> DLT >> ADD >> ADD >TIMOLOL MALEATE

SOLUTION/DROPS; OPHTHALMIC

TIMOLOL MALEATE

ALCON

EQ 0.25% BASE

N74261 001

APR 28, 1995

AT

EQ 0.5% BASE

N74262 001

APR 28, 1995

TIMOLOL MALEATE

SOLUTION/DROPS; OPHTHALMIC

TIMOPTIC

AT + MERCK

EQ 0.25% BASE

N18086 001

AT +

EQ 0.5% BASE

N18086 002

TIOCONAZOLE

OINTMENT; VAGINAL

VAGISTAT-1

+ BRISTOL MYERS

6.5%

N19355 001

DEC 30, 1986

* BRISTOL MYERS SQUIBB

6.5%

N19355 001

DEC 30, 1986

TOCAINIDE HYDROCHLORIDE

TABLET; ORAL

TONOCARD

ASTRA MERCK

400MG

N18257 001

NOV 09, 1984

+

600MG

N18257 002

NOV 09, 1984

MERCK SHARP DOHME

400MG

N18257 001

NOV 09, 1984

*

600MG

N18257 002

NOV 09, 1984

TOLMETIN SODIUM

CAPSULE; ORAL

TOLMETIN SODIUM

BAKER NORTON

EQ 400MG BASE

N73392 001

JAN 24, 1992

AB

ZENITH GOLDLINE

EQ 400MG BASE

N73392 001

JAN 24, 1992

TRAMADOL HYDROCHLORIDE

TABLET; ORAL

ULTRAM

+ JOHNSON RW

50MG

N20281 002

MAR 03, 1995

TRAMADOL HYDROCHLORIDE

TABLET; ORAL
ULTRAM
@ JOHNSON RW

100MG

N20281 001
MAR 03, 1995

TRAZODONE HYDROCHLORIDE

TABLET; ORAL
TRAZODONE HCL
MUTUAL PHARM

150MG

N73138 001
DEC 22, 1995

> ADD >
> ADD >

AB SIDMAK LABS NJ

150MG

N71525 001
MAR 09, 1988

AB TRAZON-150

SIDMAK LABS NJ

150MG

N71525 001
MAR 09, 1988

TRETINOIN

CAPSULE; ORAL
VESANOID
+ ROCHE

10MG

N20438 001
NOV 22, 1995

TRIAMCINOLONE ACETONIDE

CREAM; TOPICAL
KENALOG-H

AT APOTHECON

0.1%

N86240 001

AT WESTWOOD SQUIBB

0.1%

N86240 001

INJECTABLE; INJECTION
KENALOG-10

+ APOTHECON

10MG/ML

N12041 001

* WESTWOOD SQUIBB

10MG/ML

N12041 001

KENALOG-40

BP APOTHECON

40MG/ML

N14901 001

BP WESTWOOD SQUIBB

40MG/ML

N14901 001

LOTION; TOPICAL
KENALOG

AT APOTHECON

0.025%

N84343 001

AT +

0.1%

N84343 002

@

0.025%

N11602 003

TRIAMCINOLONE ACETONIDE

LOTION; TOPICAL
KENALOG

@ APOTHECON

0.1%

N11602 001

AT WESTWOOD SQUIBB

0.025%

N84343 001

AT *

0.1%

N84343 002

@

0.025%

N11602 001

@

0.1%

N11602 001

TRIAMCINOLONE ACETONIDE

AT BARRE

0.025%

N87191 001

@

0.025%

SEP 08, 1982

N87191 001

SEP 08, 1982

OINTMENT; TOPICAL
TRIAMCINOLONE ACETONIDE IN ABSORBASE

+ CAROLINA MEDCL

0.05%

N89595 001
MAR 23, 1995

PASTE; DENTAL
KENALOG IN ORABASE

AT + APOTHECON

0.1%

N12097 001

AT * SQUIBB

0.1%

N12097 001

TRIAMCINOLONE DIACETATE

INJECTABLE; INJECTION
ARISTOCORT

BP * LEDERLE

25MG/ML

N11685 001

+

25MG/ML

N11685 003

TRIAMCINOLONE DIACETATE

BP AKORN

25MG/ML

N85122 001

BP 40MG/ML

40MG/ML

N86394 001

@

25MG/ML

N85122 001

@

40MG/ML

N86394 001

TRIAZOLAM

TABLET; ORAL
TRIAZOLAM

AB CHELSEA LABS

0.125MG

N74445 001

AB

0.25MG

OCT 20, 1995

N74445 002

OCT 20, 1995

TRICHLORMETHIAZIDE

TABLET; ORAL			
NAQUA			
BP	SCHERING	2MG	N12265 001
@		2MG	N12265 001

TRIFLURIDINE

SOLUTION/DROPS; OPHTHALMIC			
<u>TRIFLURIDINE</u>			
AT	STERIS	1%	N74311 001
			OCT 06, 1995
<u>VIROPTIC</u>			
AT	+ GLAXO WELLCOME	1%	N18299 001

TRIHEXYPHENIDYL HYDROCHLORIDE

ELIXIR; ORAL			
<u>ARTANE</u>			
AA	LEDERLE	2MG/5ML	N06773 009
AA	+	2MG/5ML	N06773 009

TRILOSTANE

CAPSULE; ORAL			
MODRASTANE			
	SANOFI WINTHROP	30MG	N18719 002
			DEC 31, 1984
*		60MG	N18719 001
			DEC 31, 1984
@		30MG	N18719 002
			DEC 31, 1984
@		60MG	N18719 001
			DEC 31, 1984

TRIMETHOPRIM HYDROCHLORIDE

SOLUTION; ORAL			
PRIMSOL			
	ASCENT	EQ 25MG BASE/5ML	N74374 001
			JUN 23, 1995

TRIPROLIDINE HYDROCHLORIDE

TABLET; ORAL			
TRIPROLIDINE HCL			
+	DANBURY PHARMA	2.5MG	N85094 001
@		2.5MG	N85094 001

TRISULFAPYRIMIDINES (SULFADIAZINE;SULFAMERAZINE;SULFAMETHAZINE)

SUSPENSION; ORAL			
<u>TERFONYL</u>			
+	SQUIBB	167MG/5ML;167MG/5ML;	
		167MG/5ML	N06904 002
@		167MG/5ML;167MG/5ML;	
		167MG/5ML	N06904 002

TABLET; ORAL			
<u>SULFA-TRIPLE #2</u>			
AB	GLOBAL PHARMS	167MG;167MG;167MG	N80079 001
AB	+	167MG;167MG;167MG	N80079 001
<u>TERFONYL</u>			
AB	+ SQUIBB	167MG;167MG;167MG	N06904 001
@		167MG;167MG;167MG	N06904 001

TUBOCURARINE CHLORIDE

INJECTABLE; INJECTION			
<u>TUBOCURARINE CHLORIDE</u>			
AP	+ APOTHECON	3MG/ML	N05657 001
AP	+ SQUIBB	3MG/ML	N05657 001

UROFOLLITROPIN

INJECTABLE; INJECTION			
METRODIN			
	SERONO	150 IU/AMP	N19415 003
			SEP 18, 1986

VALACYCLOVIR HYDROCHLORIDE

TABLET; ORAL			
VALTREX			
+	GLAXO WELLCOME	EQ 500MG BASE	N20487 001
			JUN 23, 1995

VALACYCLOVIR HYDROCHLORIDETABLET; ORAL
VALTREX

@ GLAXO WELLCOME

EQ 1GM BASE

N20487 002
JUN 23, 1995VALPROIC ACID

SYRUP; ORAL

VALPROIC ACID

AA HIGH TECH PHARMA

250MG/5MLN74060 001
JAN 13, 1995VANCOMYCIN HYDROCHLORIDE

POWDER FOR RECONSTITUTION; ORAL

VANOCIN HCL

AA LILLY

EQ 250MG BASE/5MLN61667 002
JUL 13, 1983

AA +

EQ 500MG BASE/6ML

N61667 001

AB

EQ 250MG BASE/5ML

N61667 002

+

EQ 500MG BASE/6ML

JUL 13, 1983

N61667 001

VANCOLED

AA LEDERLE

EQ 250MG BASE/5ML

N63321 002

OCT 15, 1993

AA

EQ 500MG BASE/6ML

N63321 003

OCT 15, 1993

AB

EQ 250MG BASE/5ML

N63321 002

OCT 15, 1993

@

EQ 500MG BASE/6ML

N63321 003

OCT 15, 1993

VECURONIUM BROMIDE

INJECTABLE; INJECTION

NORCURON

AP + ORGANON

10MG/VIAL

N18776 002

APR 30, 1984

AP +

20MG/VIAL

N18776 003

JAN 03, 1992

VECURONIUM BROMIDE

AP STERIS

10MG/VIAL

N74334 001

AUG 31, 1995

VECURONIUM BROMIDE

INJECTABLE; INJECTION

VECURONIUM BROMIDE

AP

STERIS

20MG/VIALN74334 002
AUG 31, 1995VERAPAMIL HYDROCHLORIDE

INJECTABLE; INJECTION

VERAPAMIL HCL

AP

BEDFORD

2.5MG/MLN72888 001
JUL 28, 1995VITAMIN A

CAPSULE; ORAL

VITAMIN A

AA

BANNER PHARMACAPS

50,000 USP UNITS

N83973 001

@

50,000 USP UNITS

N83973 001

VITAMIN A PALMITATE

CAPSULE; ORAL

VITAMIN A

AA

BANNER PHARMACAPS

EQ 50,000 UNITS BASE

N80702 001

@

EQ 50,000 UNITS BASE

N80702 001

VITAMIN A PALMITATE

AA

BANNER PHARMACAPS

EQ 50,000 UNITS BASE

N83948 001

@

EQ 50,000 UNITS BASE

N83948 001

WARFARIN SODIUM

INJECTABLE; INJECTION

COUMADIN

+ DUPONT MERCK

5MG/VIAL

N09218 024
FEB 07, 1995WATER FOR INJECTION, STERILE

LIQUID; N/A

BACTERIOSTATIC WATER FOR INJECTION IN PLASTIC CONTAINER

AP

FUJISAWA

100%

N89099 001

DEC 29, 1987

WATER FOR INJECTION, STERILE

LIQUID; N/A

BACTERIOSTATIC WATER FOR INJECTION IN PLASTIC CONTAINER

<u>AP</u>	<u>FUJISAWA</u>	<u>100%</u>	<u>N89100 001</u>
			<u>DEC 29, 1987</u>
	@	100%	N89099 001
			DEC 29, 1987
	@	100%	N89100 001
			DEC 29, 1987

ZIDOVUDINE

> <u>ADD</u> >	TABLET; ORAL		
> <u>ADD</u> >	RETROVIR		
> <u>ADD</u> >	@ GLAXO WELLCOME	200MG	N20518 001
> <u>ADD</u> >			DEC 19, 1995

ACETAMINOPHENSUPPOSITORY; RECTAL
ACETAMINOPHEN
ABLE

120MG

325MG

650MG

N73106 001
FEB 27, 1995
N73107 001
FEB 27, 1995
N73108 001
FEB 27, 1995ANTAZOLINE PHOSPHATE; NAPHAZOLINE HYDROCHLORIDESOLUTION/DROPS; OPHTHALMIC
VASOCON-A

+ CIBA 0.5%;0.05%

N18746 002
JUL 11, 1994AVOBENZONE; PADIMATE OLOTION; TOPICAL
PHOTOPLEX

* ALLERGAN HERBERT 3%;7%

@ 3%;7%

N19459 001
SEP 30, 1988
N19459 001
SEP 30, 1988BACITRACIN ZINC; POLYMYXIN B SULFATEAEROSOL; TOPICAL
POLYSPORIN
@ GLAXO WELLCOME10,000 UNITS/GM;
2,000,000 UNITS/GMN50167 002
MAR 01, 1985> ADD > BUTOCONAZOLE NITRATE> ADD > CREAM; VAGINAL
> ADD > FEMSTAT 3
> ADD > + SYNTEX

2%

N20421 001
DEC 21, 1995CIMETIDINETABLET; ORAL
TAGAMET HB

+ SMITHKLINE BEECHAM 100MG

N20238 001
JUN 19, 1995CLEMASTINE FUMARATETABLET; ORAL
CLEMASTINE FUMARATE
PERRIGO

1.34MG

N74512 001
NOV 22, 1995CLOTRIMAZOLECREAM; VAGINAL
CLOTRIMAZOLE
TARO

1%

N72641 001
DEC 04, 1995DEXBROMPHENIRAMINE MALEATE; PSEUDOEPHEDRINE SULFATE

TABLET, EXTENDED RELEASE; ORAL

DISOBROM

GENEVA PHARMS 6MG;120MG

N70770 001
SEP 30, 1991

@ 6MG;120MG

N70770 001
SEP 30, 1991DIPHENHYDRAMINE HYDROCHLORIDESYRUP; ORAL
ANTITUSSIVE
PERRIGO

12.5MG/5ML

N71292 001
APR 10, 1987

@ 12.5MG/5ML

N71292 001
APR 10, 1987

BELDIN

HALSEY 12.5MG/5ML

N89179 001
JUN 05, 1986

@ 12.5MG/5ML

N89179 001
JUN 05, 1986

BENYLIN

* PARKE DAVIS 12.5MG/5ML

N06514 004

DIPHENHYDRAMINE HYDROCHLORIDE

SYRUP; ORAL

BENYLIN

@ PARKE DAVIS

12.5MG/5ML

N06514 004

DIPHEN

@ MORTON GROVE

12.5MG/5ML

N70118 001
OCT 01, 1985

PENNEX

12.5MG/5ML

N70118 001
OCT 01, 1985

DIPHENHYDRAMINE HCL

CUMBERLAND SWAN

12.5MG/5ML

N73611 001
AUG 20, 1992

@

12.5MG/5ML

N73611 001
AUG 20, 1992

HI TECH PHARMA

12.5MG/5ML

N72416 001
SEP 28, 1990

@

12.5MG/5ML

N72416 001
SEP 28, 1990

HYDRAMINE

BARRE

12.5MG/5ML

N70205 001
JAN 28, 1986

@

12.5MG/5ML

N70205 001
JAN 28, 1986

SILPHEN

SILARX

12.5MG/5ML

N72646 001
FEB 27, 1992

@

12.5MG/5ML

N72646 001
FEB 27, 1992

VICKS FORMULA 44

@ PROCTER AND GAMBLE

12.5MG/5ML

N70524 001
JAN 14, 1987

VICKS HLTH CARE

12.5MG/5ML

N70524 001
JAN 14, 1987FAMOTIDINE

TABLET; ORAL

PEPCID AC

+ MERCK

10MG

N20325 001
APR 28, 1995

> DLT >

> DLT >

> DLT >

> ADD >

IBUPROFEN

CAPSULE; ORAL

MIDOL

* WINTHROP

200MG

N70626 001
SEP 02, 1987IBUPROFEN

CAPSULE; ORAL

MIDOL

WINTHROP

200MG

N71002 001

SEP 02, 1987

@

200MG

N70626 001

SEP 02, 1987

@

200MG

N71002 001

SEP 02, 1987

PROVEL

+ SANDOZ

200MG

N20402 001

APR 20, 1995

SUSPENSION; ORAL

CHILDREN'S MOTRIN

+ MCNEIL CONS PRODS

100MG/5ML

N20516 001

JUN 16, 1995

TABLET; ORAL

IBUPROFEN

INVAMED

200MG

N74525 001

DEC 15, 1995

> ADD >

> ADD >

> ADD >

> ADD >

200MG

N74533 001

DEC 15, 1995

MIDOL

WINTHROP

200MG

N70591 001

SEP 02, 1987

200MG

N71001 001

SEP 02, 1987

@

200MG

N70591 001

SEP 02, 1987

@

200MG

N71001 001

SEP 02, 1987

INSULIN PORK

INJECTABLE; INJECTION

REGULAR INSULIN

NOVO NORDISK

100 UNITS/ML

N17926 003

@

100 UNITS/ML

N17926 003

INSULIN PURIFIED PORK

INJECTABLE; INJECTION

VELOSULIN

NOVO NORDISK

100 UNITS/ML

N18193 001

INSULIN PURIFIED PORK

INJECTABLE; INJECTION

VELOSULIN

@ NOVO NORDISK	100 UNITS/ML	N18193 001
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INSULIN PURIFIED PORK; INSULIN SUSP ISOPHANE PURIFIED PORK

INJECTABLE; INJECTION

INSULIN NORDISK MIXTARD (PORK)

* NOVO NORDISK	30 UNITS/ML; 70 UNITS/ML	N18195 001
@	30 UNITS/ML; 70 UNITS/ML	N18195 001

INSULIN SEMISYNTHETIC PURIFIED HUMAN

INJECTABLE; INJECTION

NOVOLIN R

* NOVO NORDISK	100 UNITS/ML	N18778 001
		AUG 30, 1983
@	100 UNITS/ML	N18778 001
		AUG 30, 1983

INSULIN SEMISYNTHETIC PURIFIED HUMAN; INSULIN SUSP ISOPHANE SEMISYNTHETIC PURIFIED HUMAN

INJECTABLE; INJECTION

MIXTARD HUMAN 70/30

* NOVO NORDISK	30 UNITS/ML; 70 UNITS/ML	N19585 001
		MAR 11, 1988
@	30 UNITS/ML; 70 UNITS/ML	N19585 001
		MAR 11, 1988

NOVOLIN 70/30

* NOVO NORDISK	30 UNITS/ML; 70 UNITS/ML	N19441 001
		JUL 11, 1986
@	30 UNITS/ML; 70 UNITS/ML	N19441 001
		JUL 11, 1986

INSULIN SUSP ISOPHANE PURIFIED BEEF

INJECTABLE; INJECTION

NPH ILETIN II

* LILLY	100 UNITS/ML	N18479 001
@	100 UNITS/ML	N18479 001

INSULIN SUSP ISOPHANE PURIFIED PORK

INJECTABLE; INJECTION

INSULIN INSULATARD NPH NORDISK

* NOVO NORDISK	100 UNITS/ML	N18194 001
@	100 UNITS/ML	N18194 001
NPH ILETIN II (PORK)		
@ LILLY	100 UNITS/ML	N18345 001
+	100 UNITS/ML	N18345 001

INSULIN SUSP ISOPHANE SEMISYNTHETIC PURIFIED HUMAN

INJECTABLE; INJECTION

INSULATARD NPH HUMAN

* NOVO NORDISK	100 UNITS/ML	N19449 001
@	100 UNITS/ML	MAY 30, 1986
		N19449 001
		MAY 30, 1986

INSULIN SUSP PROTAMINE ZINC PURIFIED BEEF

INJECTABLE; INJECTION

PROTAMINE ZINC AND ILETIN II

* LILLY	100 UNITS/ML	N18476 001
@	100 UNITS/ML	N18476 001
PROTAMINE ZINC INSULIN		
SQUIBB	100 UNITS/ML	N17928 003
+	100 UNITS/ML	N17928 003

INSULIN ZINC SUSP EXTENDED BEEF

INJECTABLE; INJECTION

ULTRALENTE INSULIN

* NOVO NORDISK	100 UNITS/ML	N17997 003
@	100 UNITS/ML	N17997 003

INSULIN ZINC SUSP PROMPT BEEF

INJECTABLE; INJECTION

SEMILENTE INSULIN

* NOVO NORDISK	100 UNITS/ML	N17996 003
@	100 UNITS/ML	N17996 003

INSULIN ZINC SUSP SEMISYNTHETIC PURIFIED HUMANINJECTABLE; INJECTION
NOVOLIN L

* NOVO NORDISK	100 UNITS/ML	N18777 001 AUG 30, 1983
@	100 UNITS/ML	N18777 001 AUG 30, 1983

KETOPROFEN

TABLET; ORAL

ACTRON
BAYER

12.5MG	N20499 001 OCT 06, 1995
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ORUDIS KT

+ WHITEHALL ROBINS

12.5MG	N20429 001 OCT 06, 1995
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LOPERAMIDE HYDROCHLORIDE

SOLUTION; ORAL

LOPERAMIDE HCL

HI TECH PHARMA

1MG/5ML	N74352 001 NOV 17, 1995
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LEMMON

1MG/5ML	N73478 001 JUN 23, 1995
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MICONAZOLE NITRATE

CREAM; VAGINAL

MICONAZOLE NITRATE

LEMMON

2%	N74136 001 JAN 04, 1995
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SUPPOSITORY; VAGINAL

MICONAZOLE NITRATE

ABLE

100MG	N71507 001 NOV 19, 1993
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NMC

100MG	N73507 001 NOV 19, 1993
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NAPROXEN SODIUM

TABLET; ORAL

ALEVE

HAMILTON PHARMS

EQ 200MG BASE

N20204 002

JAN 11, 1994

+

EQ 200MG BASE

N20204 002

JAN 11, 1994

NEOMYCIN SULFATE; POLYMYXIN B SULFATE

CREAM; TOPICAL

NEOSPORIN

@ GLAXO WELLCOME

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

N50176 002

JAN 14, 1985

NONOXYNOL-9

AEROSOL; VAGINAL

DELFIN

@ ORTHO

12.5%

N14349 002

SPONGE; VAGINAL

TODAY

@ WHITEHALL LABS

1GM

N18583 001

APR 01, 1983

@ WHITEHALL ROBINS

1GM

N18583 001

APR 01, 1983

POTASSIUM IODIDE

SOLUTION; ORAL

POTASSIUM IODIDE

* ROXANE

1GM/ML

N18551 001

FEB 19, 1982

@

1GM/ML

N18551 001

FEB 19, 1982

PYRITHIONE ZINC

LOTION; TOPICAL

HEAD & SHOULDERS CONDITIONER

* PROCTER AND GAMBLE 0.3%

N19412 002

MAR 10, 1986

PYRITHIONE ZINC

LOTION, TOPICAL
HEAD & SHOULDERS CONDITIONER
© PROCTER AND GAMBLE 0.3%

N19412 002
MAR 10, 1986

> ADD > RANITIDINE HYDROCHLORIDE

> ADD > TABLET; ORAL
> ADD > ZANTAC 75
> ADD > + GLAXO WELLCOME EQ 75MG BASE
> ADD >

N20520 001
DEC 19, 1995

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 '95

HETASTARCH 6% IN SODIUM CHLORIDE 0.9%

INJECTABLE; INJECTION

6% HETASTARCH IN SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER

ABBOTT 6GM/100ML;0.9GM/100ML N74193

JAN 30, 1995

LIST OF ORPHAN PRODUCT DESIGNATIONS & APPROVALS
[January - December, 1995]

NAME Generic/Chemical TN=Trade Name	INDICATION DESIGNATED	SPONSOR & ADDRESS DD=Date Designated MA=Marketing Approval
ADENO-AS'TED VIRAL-BASED VECTOR CYSTIC FIBROSIS GENE THERAPY TN=	TREATMENT OF CYSTIC FIBROSIS.	TARGETED GENETICS CORPORATION 1100 OLIVE WAY, SUITE 100 SEATTLE WA 98101 DD 02/15/95 MA / /
ALGLUCERASE INJECTION TN= CEREDASE	REPLACEMENT THERAPY IN PATIENTS WITH TYPE II AND III GAUCHER'S DISEASE.	GENZYME CORPORATION ONE KENDALL SQUARE CAMBRIDGE MA 02139-1562 DD 07/21/95 MA / /
AMINOCAPROIC ACID TN=	FOR THE TOPICAL TREATMENT OF TRAUMATIC HYPHEMA OF THE EYE.	ORPHAN MEDICAL 13911 RIDGEDALE DRIVE MINNETONKA MN 55305 DD 01/06/95 MA / /
APL 400-020 TN=	TREATMENT OF CUTANEOUS T CELL LYMPHOMA.	APOLLON, INC. ONE GREAT VALLEY PARKWAY MALVERN PA 19355 DD 03/08/95 MA / /
APOMORPHINE HCL TN=	TREATMENT OF ON-OFF FLUCTUATIONS ASSOCIATED WITH LATE-STAGE PARKINSON'S DISEASE.	PENTECH PHARMACEUTICALS, INC. 417 HARVESTER COURT WHEELING IL 60090 DD 07/17/95 MA / /
BROMODEOXYURIDINE TN=	RADIATION SENSITIZER IN THE TREATMENT OF PRIMARY BRAIN TUMORS.	NEOPHARM, INC. 225 EAST DEERPATH, SUITE 250 LAKE FOREST IL 60045 DD 09/18/95 MA / /
CHIMERIC A2 (HUMAN-MURINE) IGG MONOCLONAL ANTI-TNF ANTIBODY (CA2) TN=	TREATMENT OF CROHN'S DISEASE.	CENTOCOR, INC 200 GREAT VALLEY PARKWAY MALVERN, PA 19355 DD 11/14/95 MA / /
CHONDROITINASE TN=	TREATMENT OF PATIENTS UNDERGOING VITRECTOMY.	STORZ OPHTHALMICS AMERICAN CYANAMID COMPANY PEARL RIVER NY 10965 DD 02/09/95 MA / /
CLOTRIMIDAZOLE TN=	TREATMENT OF SICKLE CELL DISEASE.	BRUGNARA, CARLO M.D. THE CHILDREN'S HOSPITAL BOSTON MA 02115 DD 04/24/95 MA / /
CYSTIC FIBROSIS TR GENE THERAPY (RECOMBINANT ADENOVIRUS) TN= ADgvcFTR.10	TREATMENT OF CYSTIC FIBROSIS.	GENVAC, INCORPORATED 12111 PARKLAWN DRIVE ROCKVILLE MD 20852 DD 03/09/95 MA / /
ELCATONIN TN=	INTRATHECAL TREATMENT OF INTRACTABLE PAIN.	INNAPHARMA, INCORPORATED 75 MONTEBELLO ROAD SUFFERN NY 10901 DD 09/25/95 MA / /
ENCAPSULATED PORCINE ISLET PREPARATION TN= BETARX	TREATMENT OF TYPE I DIABETIC PATIENTS WHO ARE ALREADY ON IMMUNOSUPPRESSION.	VIVORX 3212 NEBRASKA AVENUE SANTA MONICA CA 90404 DD 07/05/95 MA / /

CUMULATIVE LIST OF DESIGNATIONS & APPROVALS

75

NAME Generic/Chemical TN=Trade Name	INDICATION DESIGNATED	SPONSOR & ADDRESS DD=Date Designated MA=Marketing Approval
ETIOCHOLANEDIONE TN=	TREATMENT OF APLASTIC ANEMIA.	SUPERGEN, INC 3158 DES PLAINES AVENUE SUITE 10 DES PLAINES, IL 60018 DD 11/03/95 MA / /
FIBRINOGEN (HUMAN) TN=	FOR THE CONTROL OF BLEEDING AND PROPHYLACTIC TREATMENT OF PATIENTS DEFICIENT IN FIBRINOGEN.	ALPHA THERAPEUTIC CORPORATION 5555 VALLEY BOULEVARD LOS ANGELES CA 90032 DD 08/23/95 MA / /
FILGRASTIM TN= NEUPOGEN	FOR USE IN THE MOBILIZATION OF PERIPHERAL BLOOD PROGENITOR CELLS FOR COLLECTION IN PATIENTS WHO WILL RECEIVE MYELOABLATIVE OR MYELOSUPPRESSIVE CHEMOTHERAPY.	AMGEN, INCORPORATED 1840 DEHAVILLAND DRIVE THOUSAND OAKS CA 91320-1789 DD 07/17/95 MA / /
GABAPENTIN TN= NEURONTIN	TREATMENT OF AMYOTROPHIC LATERAL SCLEROSIS.	WARNER-LAMBERT COMPANY PARKE-DAVIS PHARMACEUTICAL RESEARCH DIV. ANN ARBOR MI 48105-2430 DD 07/05/95 MA / /
GLUTAMINE TN=	FOR USE WITH HUMAN GROWTH HORMONE IN THE TREATMENT OF SHORT BOWEL SYNDROME (NUTRIENT MALABSORPTION FROM THE GASTROINTESTINAL TRACT RESULTING FROM AN INADEQUATE ABSORPTIVE SURFACE).	RESEARCH TRIANGLE PHARMACEUTICALS 4364 SOUTH ALSTON AVENUE DURHAM NC 27713 DD 03/06/95 MA / /
GLYCERYL TRIOLEATE AND GLYCERYL TRIERUCATE TN= LORENZO'S OIL	TREATMENT OF ADRENOLEUKODYSTROPHY.	MOSER, HUGO W. M.D. JOHNS HOPKINS UNIVERSITY BALTIMORE MD 21205 DD 02/14/95 MA / /
HEPATITIS B IMMUNE GLOBULIN, INTRAVENOUS TN= H-BIGIV	PROPHYLAXIS AGAINST HEPATITIS B VIRUS REINFECTION IN LIVER TRANSPLANT PATIENTS.	NORTH AMERICAN BIOLOGICS, INC. 16500 N.W. 15th AVENUE MIAMI FL 33169 DD 03/08/95 MA / /
HUMAN GROWTH HORMONE TN=	FOR USE WITH GLUTAMINE IN THE TREATMENT OF SHORT BOWEL SYNDROME (NUTRIENT MALABSORPTION FROM THE GASTROINTESTINAL TRACT RESULTING FROM AN INADEQUATE ABSORPTIVE SURFACE).	RESEARCH TRIANGLE PHARMACEUTICALS 4364 SOUTH ALSTON AVENUE DURHAM NC 27713 DD 03/06/95 MA / /
HUMAN IMMUNODEFICIENCY VIRUS IMMUNE GLOBULIN TN= HIVIG	TREATMENT OF HIV-INFECTED PEDIATRIC PATIENTS.	NORTH AMERICAN BIOLOGICS, INC. 16500 N.W. 15TH AVENUE MIAMI FL 33169 DD 01/04/95 MA / /
INTERFERON GAMMA-1B TN=ACTIMMUNE	TREATMENT OF RENAL CELL CARCINOMA.	GENENTECH, INC. 460 POINT SAN BRUNO BOULEVARD SOUTH SAN FRANCISCO, CA 94080 DD 12/04/95 MA / /
INTRAVITREAL GANCICLOVIR FREE ACID IMPLANT TN= VITRASERT IMPLANT	TREATMENT OF CYTOMEGALOVIRUS RETINITIS.	CHIRON VISION 500 IOLAB DRIVE CLAREMONT CA 91711 DD 06/07/95 MA / /
KL4-SURFACTANT TN=	TREATMENT OF ACUTE RESPIRATORY DISTRESS SYNDROME IN ADULTS.	R.W.JOHNSON RESEARCH INSTITUTE ROUTE 202, PO BOX 300 RARITAN NJ 08869-0602 DD 07/17/95 MA / /

CUMULATIVE LIST OF DESIGNATIONS & APPROVALS

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NAME Generic/Chemical TN=Trade Name	INDICATION DESIGNATED	SPONSOR & ADDRESS DD=Date Designated MA=Marketing Approval
KL4-SURFACTANT TN=	TREATMENT OF RESPIRATORY DISTRESS SYNDROME IN PREMATURE INFANTS.	COCHRANE, CHARLES G. M.D. THE SCRIPPS RESEARCH INSTITUTE 10666 NORTH TORREY PINES ROAD IMM 12 LA JOLLA, CA 92037 DD 10/18/95 MA / /
LAMOTRIGINE TN= LAMICTAL	TREATMENT OF LENNOX-GASTAUT SYNDROME.	BURROUGHS-WELLCOME COMPANY 3030 CORNWALLIS ROAD, P.O. BOX 12700 RESEARCH TRIANGLE PK NC 27709 DD 08/23/95 MA / /
LIDOCAINE PATCH 5% TN=LIDOCAINE PATCH	TREATMENT OF POST-HERPETIC NEURALGIA RESULTING FROM HERPES ZOSTER INFECTIONS.	HIND HEALTH CARE, INC 165 GIBRALTAR COURT SUNNYVALE, CA 94089 DD 10/24/95 MA / /
MECASERMIN TN=	TREATMENT OF GROWTH HORMONE INSUFFICIENCY SYNDROME.	GENENTECH, INC. 460 POINT SAN BRUNO BOULEVARD SOUTH SAN FRANCISCO, CA 94080 DD 12/12/95 MA / /
MITOLACTOL TN=	AS ADJUVANT THERAPY IN THE TREATMENT OF PRIMARY BRAIN TUMORS.	BIOPHARMACEUTICS, INC. 990 STATION ROAD BELLPORT NY 11713 DD 07/12/95 MA / /
MYCOBACTERIUM AVIUM SENSITIN RS-10 TN=	FOR USE IN THE DIAGNOSIS OF INVASIVE MYCOBACTERIUM AVIUM DISEASE IN IMMUNOCOMPETENT INDIVIDUALS.	STATENS SERUMINSTITUT 5 ARTILLERIVEJ DK-2300 COPENHAGEN S DENMARK DD 10/11/95 MA / /
NITRIC OXIDE TN=	TREATMENT OF ACUTE RESPIRATORY DISTRESS SYNDROME IN ADULTS.	OHMEDA PHARMACEUTICAL PRODUCTS DIVISION 110 ALLEN ROAD LIBERTY CORNER NJ 07938-0804 DD 07/10/95 MA / /
NTBC TN=	TREATMENT OF TYROSINEMIA TYPE 1.	SWEDISH ORPHAN AB ORPHAN PHARMACEUTICAL, USA, INC. NASHVILLE TN 37217 DD 05/16/95 MA / /
OMEGA-3 (N-3) POLYUNSATURATED FATTY ACID WITH ALL DOUBLE BONDS IN THE CIS CONFIGURATION TN=	PREVENTION OF ORGAN GRAFT REJECTION.	RESEARCH TRIANGLE PHARMACEUTICAL 4364 SOUTH ALSTON AVENUE DURHAM, NC 27713 DD 11/22/95 MA / /
PHENYLALANINE AMMONIA-LYASE TN= PHENYLASE	TREATMENT OF HYPERPHENYLALANINEMIA.	IBEX TECHNOLOGIES, INC. 5485 PARE MONTREAL, QUEBEC DD 03/08/95 MA / /
PORFIROMYCIN TN=	TREATMENT OF HEAD AND NECK CANCER.	ONCORX INC. 4 SCIENCE PARK NEW HAVEN CT 06511 DD 09/19/95 MA / /
PURIFIED TYPE II COLLAGEN TN= COLLOLAL	TREATMENT OF JUVENILE RHEUMATOID ARTHRITIS.	AUTOIMMUNE, INCORPORATED 128 SPRING STREET LEXINGTON MA 02173 DD 02/09/95 MA / /

CUMULATIVE LIST OF DESIGNATIONS & APPROVALS

77

NAME Generic/Chemical TN=Trade Name	INDICATION DESIGNATED	SPONSOR & ADDRESS DD=Date Designated MA=Marketing Approval
RECOMBINANT HUMAN GELSOLIN TN=	TREATMENT OF ACUTE AND CHRONIC RESPIRATORY SYMPTOMS OF BRONCHIECTASIS.	BIOGEN, INCORPORATED 14 CAMBRIDGE CENTER CAMBRIDGE MA 02142 DD 03/06/95 MA / /
RECOMBINANT HUMAN INSULIN-LIKE GROWTH FACTOR I TN=	TREATMENT OF POST-POLIOMYELITIS SYNDROME.	CEPHALON, INC 145 BRANDYWINE PARKWAY WEST CHESTER, PA 19380 DD 10/13/95 MA / /
RECOMBINANT HUMAN INSULIN-LIKE GROWTH FACTOR I TN= IGEF	TREATMENT OF GROWTH HORMONE RECEPTOR DEFICIENCY.	PHARMACIA, INC. PO BOX 16529 COLUMBUS OH 43216-6529 DD 06/07/95 MA / /
RECOMBINANT HUMAN INSULIN-LIKE GROWTH FACTOR I TN= IGEF	TREATMENT OF ANTIBODY-MEDIATED GROWTH HORMONE RESISTANCE IN PATIENTS WITH ISOLATED GROWTH HORMONE DEFICIENCY IA.	PHARMACIA, INC. P.O. BOX 16529 COLUMBUS OH 43216-6529 DD 06/07/95 MA / /
RECOMBINANT HUMAN RELAXIN TN=	TREATMENT OF PROGRESSIVE SYSTEMIC SCLEROSIS.	CONNECTIVE THERAPEUTICS, INC. 3400 WEST BAYSHORE ROAD PALO ALTO, CA 94303 DD 11/03/95 MA / /
RECOMBINANT METHIONYL HUMAN STEM CELL FACTOR TN=	FOR USE IN COMBINATION WITH FILGRASTIM TO DECREASE THE NUMBER OF PHERESES REQUIRED TO COLLECT PERIPHERAL BLOOD PROGENITOR CELLS CAPABLE OF PROVIDING RAPID MULTI-LINEAGE HEMATOPOIETIC RECONSTITUTION FOLLOWING MYELOSUPPRESSIVE OR MYELOABLATIVE THERAPY.	AMGEN, INCORPORATED 1840 DEHAVILLAND DRIVE THOUSAND OAKS CA 91320-1789 DD 07/05/95 MA / /
RECOMBINANT METHIONYL HUMAN STEM CELL FACTOR TN=	TREATMENT OF PRIMARY BONE MARROW FAILURE.	AMGEN, INCORPORATED 1840 DEHAVILLAND DRIVE THOUSAND OAKS CA 91320-1789 DD 11/22/95 MA / /
RGG0853, E1A LIPID COMPLEX TN=	TREATMENT OF ADVANCED OVARIAN CANCER THAT OVEREXPRESSES THE HER-2/neu ONCOGENE.	RGENE THERAPEUTICS, INC. 2170 BUCKTHORNE PLACE, SUITE 230 THE WOODLANDS TX 77380 DD 09/19/95 MA / /
RIFAPENTINE TN=	TREATMENT OF PULMONARY TUBERCULOSIS.	MARION MERRELL DOW INC. PO BOX 9627 (PARK A) KANSAS CITY MO 94137 DD 06/09/95 MA / /
RIFAPENTINE TN=	TREATMENT OF MYCOBACTERIUM AVIUM COMPLEX IN PATIENTS WITH ACQUIRED IMMUNODEFICIENCY SYNDROME.	MARION MERRELL DOW INC. PO BOX 9627 (PARK A) KANSAS CITY MO 64137 DD 06/09/95 MA / /
SARGRAMOSTIM TN= LEUKINE	TO REDUCE NEUTROPENIA AND LEUKOPENIA AND DECREASE THE INCIDENCE OF DEATH DUE TO INFECTION IN PATIENTS WITH ACUTE MYELOGENOUS LEUKEMIA.	IMMUNEX CORPORATION 51 UNIVERSITY STREET SEATTLE WA 98101 DD 03/06/95 MA / /
SORIVUDINE TN= BRAVAVIR	TREATMENT OF HERPES ZOSTER (SHINGLES) IN IMMUNOCOMPROMISED PATIENTS.	BRISTOL MYERS SQUIBB 5 RESEARCH PARKWAY P.O. BOX 5100 WALLINGFORD, CT 06492 DD 11/09/95 MA / /
STERILE AEROSOL TALC TN=	TREATMENT OF MALIGNANT PLEURAL EFFUSION.	BRYAN CORPORATION 4 PLYMPTON STREET WOBBURN MA 01801 DD 09/18/95 MA / /

CUMULATIVE LIST OF DESIGNATIONS & APPROVALS

78

NAME Generic/Chemical TN=Trade Name	INDICATION DESIGNATED	SPONSOR & ADDRESS DD=Date Designated MA=Marketing Approval
SU-101 TN=	TREATMENT OF MALIGNANT GLIOMA.	SUGEN, INC. 515 GALVESTON DRIVE REDWOOD CITY CA 94063-4720 DD 05/25/95 MA / /
SYNSORB PK TN=	TREATMENT OF VEROCYTOTOXOGENIC E. COLI INFECTIONS.	SYNSORB BIOTECH INC. FOURTH FLOOR, 140 4TH AVENUE SW CALGARY, ALBERTA DD 07/17/95 MA / /
THALIDOMIDE TN=	TREATMENT OF SEVERE RECURRENT APHTHOUS STOMATITIS IN SEVERELY, TERMINALLY IMMUNOCOMPROMISED PATIENTS.	CELGENE CORPORATION P.O. BOX 4914 WARREN NJ 07059 DD 05/01/95 MA / /
THALIDOMIDE TN=	TREATMENT AND PREVENTION OF RECURRENT APHTHOUS ULCERS IN SEVERELY, TERMINALLY IMMUNOCOMPROMISED PATIENTS.	ANDRULIS RESEARCH CORPORATION 11800 BALTIMORE AVENUE, SUITE 113 BELTSVILLE MD 20705 DD 05/15/95 MA / /
THALIDOMIDE TN= SYNOVIR	TREATMENT OF ERYTHEMA NODOSUM LEPROSUM.	CELGENE CORPORATION 7 POWDER HORN DRIVE, PO BOX 4914 WARREN NJ 07059 DD 07/26/95 MA / /
TRISODIUM CITRATE CONCENTRATION TN= HEMOCITRATE	FOR USE IN LEUKAPHERESIS PROCEDURES.	HEMOTEC MEDICAL PRODUCTS, INC. BOX 19255 JOHNSTON RI 02919 DD 06/15/95 MA / /
TYLOXAPOL TN=	TREATMENT OF CYSTIC FIBROSIS.	KENNEDY & HOIDAL, MDs 50 NORTH MEDICAL DRIVE, U OF UTAH SALT LAKE CITY UT 84132 DD 03/08/95 MA / /
URIDINE 5'-TRIPHOSPHATE TN=	TREATMENT OF CYSTIC FIBROSIS.	INSPIRE PHARMACEUTICALS, INC P.O. BOX 14285 RESEARCH TRIANGLE PARK, NC 27709 DD 12/04/95 MA / /

CUMULATIVE LIST OF DESIGNATIONS & APPROVALS

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NAME Generic/Chemical TN=Trade Name	INDICATION DESIGNATED	SPONSOR & ADDRESS DD=Date Designated MA=Marketing Approval
Approved Orphan Products in 1995		
AMIFOSTINE TN= ETHYOL	FOR USE AS A CHEMOPROTECTIVE AGENT FOR CISPLATIN IN THE TREATMENT OF ADVANCED OVARIAN CARCINOMA.	U.S. BIOSCIENCE, INC. ONE TOWER BRIDGE 100 FRONT STREET WEST CONSHOHOCKEN, PA 19428 DD 05/30/90 MA 12/08/95
AMIODARONE HCL TN= CORDARONE	FOR THE ACUTE TREATMENT AND PROPHYLAXIS OF LIFE-THREATENING VENTRICULAR TACHYCARDIA OR VENTRICULAR FIBRILLATION.	WYETH-AYERST LABORATORIES P.O. BOX 8299 PHILADELPHIA PA 19101-1245 DD 03/16/94 MA 08/03/95
DEXRAZOXANE TN= ZINECARD	FOR THE PREVENTION OF CARDIOMYOPATHY ASSOCIATED WITH DOXORUBICIN ADMINISTRATION.	PHARMACIA, INC. P.O. BOX 16529 COLUMBUS OH 43216-6529 DD 12/17/91 MA 05/26/95
EPOPROSTENOL TN= FLOLAN	LONG-TERM TREATMENT OF PRIMARY PULMONARY HYPERTENSION IN NEW YORK HEART ASSOCIATION CLASS III AND CLASS IV PATIENTS.	BURROUGHS WELLCOME COMPANY 3030 CORNWALLIS ROAD RESEARCH TRIANGLE PK NC 27709 DD 09/25/85 MA 09/20/95
INTERFERON ALPHA-2A TN= ROFERON	TREATMENT OF CHRONIC MYELOGENOUS LEUKEMIA (CML).	HOFFMAN-LA ROCHE 340 KINGSLAND STREET NUTLEY, NJ 07110-1199 DD 06/06/89 MA 10/19/95
PORFIMER SODIUM TN= PHOTOFRIN	FOR THE PHOTODYNAMIC THERAPY OF PATIENTS WITH PRIMARY OR RECURRENT OBSTRUCTING (EITHER PARTIALLY OR COMPLETELY) ESOPHAGEAL CARCINOMA.	QLT PHOTOTHERAPEUTICS, INC LEDERLE LABORATORIES 401 NORTH MIDDLETOWN ROAD PEARL RIVER, NY 10965 DD 06/06/89 MA 12/27/95
Rho (D) IMMUNE GLOBULIN INTRAVENOUS (HUMAN) TN= WinRho SD	TREATMENT OF IMMUNE THROMBOCYTOPENIC PURPURA.	RH PHARMACEUTICALS, INC. 104 CHANCELLOR MATHESON ROAD WINNIPEG, MANITOBA DD 11/09/93 MA 03/24/95
RILUZOLE TN=RILUTEK	TREATMENT OF AMYOTROPHIC LATERAL SCLEROSIS.	RHONE-POULENC RORER PHARM. 500 ARCOLA ROAD, PO BOX 1200 COLLEGEVILLE PA 19426-0107 DD 03/16/93 MA 12/12/95
TRETINOIN TN= VESANOID	INDUCTION OF REMISSION IN PATIENTS WITH ACUTE PROMYELOCYTIC LEUKEMIA WHO ARE REFRACTORY TO OR UNABLE TO TOLERATE ANTHRACYCLINE BASED CYTOTOXIC CHEMOTHERAPEUTIC REGIMENS.	HOFFMANN-LA ROCHE, INC. 340 KINGSLAND STREET NUTLEY NJ 07110 DD 10/24/90 MA 11/22/95

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

NO DECEMBER 1995 ADDITIONS

BIOPHARMACEUTIC GUIDANCE AVAILABILITY

DRUG NAME (DOSAGE FORM)	DATE	REVISED DATE
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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) 5600 FISHERS LANE, ROCKVILLE, MD 20857.

REFER BACK TO THE APPROVED DRUG PRODUCTS WITH THERAPEUTIC EQUIVALENCE EVALUATIONS, 15TH 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> INTERIM (TABLET)	NOV 15, 1995	
CORTICOSTEROIDS, DERMATOLOGIC <i>IN VIVO</i> (TOPICAL)	JUN 02, 1995	
FLURBIPROFEN (TABLET)	DEC 24, 1992	JUN 08, 1995
HYDROXYCHLOROQUINE SULFATE (TABLET)	DEC 28, 1995	
NAPROXEN (TABLET)	JUN 12, 1992	JUN 08, 1995
PENTOXIFYLLINE (EXTENDED-RELEASE TABLET)	DEC 22, 1995	
SELEGILINE HYDROCHLORIDE (TABLET)	DEC 22, 1995	
TERFENADINE (TABLET)	JUN 12, 1992	SEP 11, 1995

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, 15TH EDITION FOR A FULL LISTING OF ANDA SUITABILITY PETITIONS DATA. ONLY NEW DATA WILL BE ADDED TO THE CUMULATIVE SUPPLEMENT.

ACETAMINOPHEN; ASPIRIN; CODEINE PHOSPHATE TABLET; ORAL	150MG 180MG 15MG	94 P-0212/ CP1	MIKART	NEW DOSAGE FORM	APPROVED JAN 19, 1995
ACETAMINOPHEN; ASPIRIN; CODEINE PHOSPHATE TABLET; ORAL	150MG 180MG 30MG	94 P-0211/ CP1	MIKART	NEW DOSAGE FORM	APPROVED JAN 19, 1995
ACETAMINOPHEN; ASPIRIN; CODEINE PHOSPHATE TABLET; ORAL	150MG 180MG 60MG	94 P-0210/ CP1	MIKART	NEW DOSAGE FORM	APPROVED JAN 19, 1995
ACETAMINOPHEN; CAFFEINE; DIHYDROCODEINE BITARTRATE TABLET; ORAL	712.8MG 60MG 32MG	93 P-0484/ CP1	MIKART	NEW DOSAGE FORM NEW STRENGTH	APPROVED JAN 19, 1995
ACETAMINOPHEN; CODEINE PHOSPHATE TABLET, CHEWABLE; ORAL	120MG 12MG	94 P-0182/ CP1	WE PHARMS	NEW DOSAGE FORM	APPROVED JAN 19, 1995
ACETAMINOPHEN HYDROCODONE BITARTRATE TABLET; ORAL	650MG 5MG	95 P-0222/ CP2	MIKART	NEW STRENGTH	APPROVED NOV 21, 1995
ALBUTEROL SULFATE TABLET, CHEWABLE; ORAL	EQ 2MG BASE EQ 4MG BASE	92 P-0335/ CP1	WE PHARMS	NEW DOSAGE FORM	APPROVED JAN 19, 1995
ATRAURIUM BESYLATE INJECTABLE; INJECTION	25MG/ML	94 P-0314/ CP1	ABBOTT	NEW STRENGTH	APPROVED MAY 02, 1995
CALCITONIN, SALMON INJECTABLE; INJECTION	100 IU/ML (0.5ML/AMP) 1ML/AMP)	95 P-0080/ CP1	FERRING	NEW STRENGTH	APPROVED AUG 07, 1995
CAPTOPRIL SOLUTION; ORAL	25MG/ML	95 P-0008/ CP1	ROXANE	NEW DOSAGE FORM NEW STRENGTH	APPROVED AUG 07, 1995
FLUOROURACIL GEL; TOPICAL	5%	94 P-0263/ CP1	BRADLEY PHARMS	NEW DOSAGE FORM	APPROVED SEP 12, 1995
IOPAMIDOL INJECTABLE; INJECTION	61% (250ML/CONTAINER) (500ML/CONTAINER)	95 P-0073/ CP1	ABBOTT	NEW STRENGTH	APPROVED AUG 23, 1995
IOPAMIDOL INJECTABLE; INJECTION	76% (250ML/CONTAINER) (500ML/CONTAINER)	95 P-0073/ CP1	ABBOTT	NEW STRENGTH	APPROVED AUG 23, 1995

ANDA SUITABILITY PETITIONS

PETITIONS APPROVED

DRUG NAME DOSAGE FORM; ROUTE	STRENGTH (CONTAINER SIZE)	DOCKET NUMBER	PETITIONER	REASON FOR PETITION	STATUS
LEUCOVORIN CALCIUM INJECTABLE; INJECTION	10MG/ML (5ML/VIAL)	94 P-0433/ CP2	LEDERLE	NEW DOSAGE FORM	APPROVED AUG 07, 1995
LEUCOVORIN CALCIUM INJECTABLE; INJECTION	10MG/ML (20ML/VIAL)	94 P-0433/ CP3	LEDERLE	NEW DOSAGE FORM NEW STRENGTH	APPROVED AUG 07, 1995
LEUCOVORIN CALCIUM INJECTABLE; INJECTION	10MG/ML (35ML/VIAL)	94 P-0433/ CP1	LEDERLE	NEW DOSAGE FORM	APPROVED AUG 07, 1995
LEUCOVORIN CALCIUM INJECTABLE; INJECTION	EQ 10MG BASE/ML (100MG/VIAL)	93 P-0427/ CP3	ABBOTT	NEW DOSAGE FORM	APPROVED JAN 19, 1995
LEUCOVORIN CALCIUM INJECTABLE; INJECTION	EQ 10MG BASE/ML (250MG/VIAL)	93 P-0427/ CP2	ABBOTT	NEW DOSAGE FORM NEW STRENGTH	APPROVED JAN 19, 1995
LORAZEPAM SOLUTION; ORAL	0.5MG/5ML	94 P-0199/ CP1	ROXANE	NEW DOSAGE FORM	APPROVED FEB 07, 1995
MEDROXYPROGESTERONE ACETATE TABLET; ORAL	2MG 4MG 8MG	92 P-0452/ CP1	CARNRICK	NEW STRENGTH	APPROVED AUG 07, 1995
METHYLPREDNISOLONE TABLET, CHEWABLE; ORAL	4MG 16MG 24MG 32MG	94 P-0432/ CP1	DURA PHARMS	NEW DOSAGE FORM	APPROVED AUG 07, 1995
SULFAMETHOXAZOLE; TRIMETHOPRIM TABLET, CHEWABLE; ORAL	200MG 40MG	94 P-0186/ CP1	DURA PHARMS	NEW DOSAGE FORM	APPROVED JAN 19, 1995
THIORIDAZINE HYDROCHLORIDE SOLUTION; ORAL	25MG/5ML	92 P-0283/ CP1	UDL LABS	NEW STRENGTH	APPROVED JAN 19, 1995

ANDA SUITABILITY PETITIONS

PETITIONS DENIED

DRUG NAME DOSAGE FORM; ROUTE	STRENGTH (CONTAINER SIZE)	DOCKET NUMBER	PETITIONER	REASON FOR PETITION	STATUS
HYDROMORPHONE HYDROCHLORIDE INJECTABLE; INJECTION	0.2MG/ML (50ML PREFILLED SYRINGE)	95 P-0022/ CP1	ASTRA	NEW INDICATION NEW ROUTE OF ADMINISTRATION NEW STRENGTH	DENIED SEP 07, 1995
MEFLOQUINE HYDROCHLORIDE TABLET; ORAL	275MG	94 P-0329/ CP1	LACASSE	NEW DOSING REGIMEN NEW STRENGTH	DENIED SEP 07, 1995
NICOTINE POLACRILEX LOLLIPOP; ORAL	2MG	93 P-0414/ CP1	SAVAGE	NEW DOSAGE FORM	DENIED MAY 02, 1995
NORETHINDRONE ACETATE TABLET; ORAL	1MG 2.5MG	94 P-0446/ CP1	APOTHECON	NEW INGREDIENT NEW STRENGTH	DENIED SEP 07, 1995
SELEGILINE HYDROCHLORIDE TABLET, EXTENDED RELEASE; ORAL	10MG	94 P-0387/ CP1	PHARMAVENE	NEW DOSAGE FORM NEW STRENGTH	DENIED MAY 08, 1995
TERFENADINE TABLET, CHEWABLE; ORAL	60MG	94 P-0119/ CP1	DURA PHARMS	NEW DOSAGE FORM	DENIED AUG 23, 1995

EXCLUSIVITY TERMS

DUE TO SPACE LIMITATIONS IN THE EXCLUSIVITY COLUMN, ABBREVIATIONS AND REFERENCES HAVE BEEN DEVELOPED. PLEASE REFER BACK TO THE APPROVED DRUG PRODUCTS WITH THERAPEUTIC EQUIVALENCE EVALUATIONS, 15TH EDITION FOR A FULL LISTING OF EXCLUSIVITY TERMS (ABBREVIATIONS, NEW DOSING SCHEDULE, NEW INDICATIONS AND PATENT USE CODES). ONLY NEW CODES WILL BE ADDED TO THE CUMULATIVE SUPPLEMENT.

REFERENCES

NEW DOSING SCHEDULE

- D-26 ONCE WEEKLY APPLICATION
- D-27 BID DOSING IN PATIENTS 12 YEARS OF AGE AND OLDER FOR PREVENTION OF NAUSEA AND VOMITING ASSOCIATED WITH MODERATELY EMETOGENIC CANCER CHEMOTHERAPY
- D-28 USE OF ISOVUE-370 IN EXCRETORY UROGRAPHY AT EQUIVALENT GRAMS OF IODINE TO THE CURRENTLY APPROVED ISOVUE-250 AND ISOVUE-300

REFERENCES

NEW INDICATION

- I-117 TO SLOW THE PROGRESSION OF CORONARY ATHEROSCLEROSIS IN PATIENTS WITH CORONARY HEART DISEASE
- I-118 PREVENTION OF DEEP VEIN THROMBOSIS, WHICH MAY LEAD TO PULMONARY EMBOLISM, FOLLOWING KNEE REPLACEMENT SURGERY
- I-119 TREATMENT OF ANEMIA CAUSED BY UTERINE LEIOMYOMATA IN WOMEN WHO FAIL IRON THERAPY
- I-120 MAINTENANCE THERAPY FOR GASTRIC ULCER PATIENTS AT REDUCED DOSAGE AFTER HEALING ACUTE ULCERS
- I-121 EXPANDED PATIENT POPULATION - USE IN ICU PATIENTS
- I-122 PSORIASIS OF THE SCALP
- I-123 RELIEF OF MILD TO MODERATE PAIN IN PATIENTS AGED 6 MONTHS AND OLDER
- I-124 LEUCOCYTE LABELED SCINTIGRAPHY AS AN ADJUNCT IN THE LOCALIZATION OF INTRA-ABDOMINAL INFECTION AND INFLAMMATORY BOWEL DISEASE
- I-125 EXPANSION OF CONSCIOUS SEDATION INDICATION TO INCLUDE SHORT THERAPEUTIC PROCEDURES
- I-126 ADJUNCT TO THALLIUM-201 MYOCARDIAL PERFUSION IN PATIENTS UNABLE TO EXERCISE ADEQUATELY
- I-127 TREATMENT OF ACYCLOVIR-RESISTANT HERPES IN IMMUNOCOMPROMISED PATIENTS
- I-128 IN PATIENTS WITH CORONARY HEART DISEASE AND HYPERCHOLESTEROLEMIA: TO REDUCE THE RISK OF TOTAL MORTALITY BY REDUCING CORONARY DEATH; REDUCE THE RISK OF NON-FATAL MYOCARDIAL INFARCTION; REDUCE THE RISK FOR UNDERGOING MYOCARDIAL REVASCLARIZATION PROCEDURES; REDUCTION OF ELEVATED TOTAL AND LDL CHOLESTEROL LEVELS IN PATIENTS WITH PRIMARY HYPERCHOLESTEROLEMIA (TYPES IIA AND IIB)
- I-129 TREATMENT OF ALCOHOL DEPENDENCE
- I-130 MAINTENANCE OF HEALING OF EROSIIVE ESOPHAGITIS
- I-131 PERIPHERAL ARTERIOGRAPHY
- I-132 TREATMENT OF MANIC PHASE OF BIPOLAR DISORDER
- I-133 MANAGEMENT OF CHRONIC STABLE ANGINA
- I-134 HEART FAILURE POST MYOCARDIAL INFARCTION
- I-135 BONE METASTASES ASSOCIATED WITH MULTIPLE MYELOMA
- I-136 IDIOPATHIC CHRONIC URTICARIA
- I-137 PREVENTION OF MEAL-INDUCED HEARTBURN, ACID INDIGESTION, AND SOUR STOMACH WHEN TAKEN 30 MINUTES PRIOR TO CONSUMING FOOD OR BEVERAGES
- I-138 TREATMENT OF ACUTE RECURRENT GENITAL HERPES
- I-139 PALLIATIVE TREATMENT OF ADVANCED BREAST CANCER IN PRE- AND PERIMENOPAUSAL WOMEN
- I-140 PREVENTION OF CYTOMEGALOVIRUS (CMV) DISEASE IN INDIVIDUALS WITH HIV INFECTION AT RISK FOR DEVELOPING CMV DISEASE

REFERENCES

PATENT USE CODE

- U-102 METHOD OF HORMONALLY TREATING MENOPAUSAL OR POST-MENOPAUSAL DISORDERS IN WOMEN
- U-103 TREATMENT OF OCULAR HYPERTENSION
- U-104 TREATMENT OF AQUEOUS HUMOR FORMATION AND INTRAOCULAR PRESSURE
- U-105 EMESIS
- U-106 TREATMENT OF EPILEPSY
- U-107 TREATMENT OF HYPERTENSION AND ANGINA PECTORIS
- U-108 SHORT-TERM TREATMENT OF ACTIVE DUODENAL ULCER, GASTROESOPHAGEAL REFLUX DISEASE (GERD), SEVERE EROSIIVE ESOPHAGITIS, POORLY RESPONSIVE SYMPTOMATIC GERD, PATHOLOGICAL HYPERSECRETORY CONDITIONS AND MAINTENANCE OF HEALING OF EROSIIVE ESOPHAGITIS

EXCLUSIVITY TERMS**REFERENCES**
PATENT USE CODE

U-109 USE AS AN ADJUNCT TO DIET IN THE TREATMENT OF ELEVATED TOTAL CHOLESTEROL AND LDL-C LEVELS IN PATIENTS WITH PRIMARY HYPERCHOLESTEROLEMIA WHOSE RESPONSE TO DIETARY RESTRICTION OF SATURATED FAT AND CHOLESTEROL AND OTHER NONPHARMACOLOGICAL MEASURES HAS NOT BEEN ADEQUATE

U-110 USE AS A RETRIEVABLE PESSARY

U-111 DIABETES

U-112 CONTRACEPTION

U-113 METHOD OF CONDUCTING A RADIOLOGICAL EXAMINATION OF A PATIENT BY ADMINISTERING TO SAID PATIENT A RADIOPAQUE AMOUNT OF IOPROMIDE

U-114 USE FOR INHIBITING BONE RESORPTION

U-115 USE OF VASODILATORS TO EFFECT AND ENHANCE AN ERECTION (AND THUS TREAT ERECTILE DYSFUNCTION), BY INJECTION INTO THE PENIS

U-116 METHOD OF MYOCARDIAL IMAGING

U-117 TREATMENT OF OCULAR ALLERGIC RESPONSE IN HUMAN EYES

U-118 METHOD OF LOWERING BLOOD SUGAR LEVEL

U-119 TREATMENT OF NASAL HYPERSECRETION

U-120 CONTROLLING OR PREVENTING POST-OPERATIVE INTRAOCULAR PRESSURE RISES ASSOCIATED WITH OPHTHALMIC LASER SURGICAL PROCEDURES

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
20482 001	ACARBOSE; PRECOSE	4904769	FEB 27, 2007		NCE	SEP 06, 2000
20482 002	ACARBOSE; PRECOSE	4904769	FEB 27, 2007		NCE	SEP 06, 2000
19872 001	ACETAMINOPHEN; TYLENOL	5004613	JUL 27, 2007			
		4968509	NOV 06, 2007			
		4820522	JUL 27, 2007		NDF	JUN 08, 1997
19806 001	ACRIVASTINE; SEMPREX-D	4501893	FEB 01, 2003			
20059 001	ADENOSINE; ADENOSCAN	5070877	DEC 10, 2008	U-116	I-126	MAY 18, 1998
18062 001	ALBUTEROL SULFATE; PROVENTIL	4499108	JUN 08, 2003			
19489 001	ALBUTEROL SULFATE; VENTOLIN ROTACAPS	4353365	APR 24, 1998			
		4206758	APR 24, 1998			
18702 001	ALCLOMETASONE DIPROPIONATE; ACLOVATE	4124707	DEC 12, 1996			
18707 001	ALCLOMETASONE DIPROPIONATE; ACLOVATE	4124707	DEC 12, 1996			
20560 001	ALENDRONATE SODIUM; FOSAMAX	5358941	DEC 02, 2012			
		4621077	NOV 04, 2003	U-114	NCE	SEP 29, 2000
20560 002	ALENDRONATE SODIUM; FOSAMAX	5358941	DEC 02, 2012			
		4621077	NOV 04, 2003	U-114	NCE	SEP 29, 2000
19353 001	ALFENTANIL HYDROCHLORIDE; ALFENTA	4167574	MAY 05, 1999			
20379 001	ALPROSTADIL; CAVERJECT	4127118	MAR 16, 1997	U-115	NP	JUL 06, 1998
20379 002	ALPROSTADIL; CAVERJECT	4127118	MAR 16, 1997	U-115	NP	JUL 06, 1998
>ADD> >ADD>	20221 001	AMIFOSTINE; ETHYOL			ODE	DEC 08, 2002
					NCE	DEC 08, 2000
	20377 001	AMIODARONE HYDROCHLORIDE; CORDARONE			ODE	AUG 03, 2002
					NDF	AUG 03, 1998
19787 001	AMLODIPINE BESYLATE; NORVASC	4879303	MAR 25, 2007			
19787 002	AMLODIPINE BESYLATE; NORVASC	4879303	MAR 25, 2007			
19787 003	AMLODIPINE BESYLATE; NORVASC	4879303	MAR 25, 2007			
20364 002	AMLODIPINE BESYLATE; LOTREL	4879303	MAR 25, 2007		NC	MAR 03, 1998
		4572909	AUG 01, 2006		NCE	JUN 25, 1996
		4410520	AUG 11, 2003		NCE	JUL 31, 1997
20364 003	AMLODIPINE BESYLATE; LOTREL	4879303	MAR 25, 2007		NC	MAR 03, 1998
		4572909	AUG 01, 2006		NCE	JUN 25, 1996
		4410520	AUG 11, 2003		NCE	JUL 31, 1997
20364 004	AMLODIPINE BESYLATE; LOTREL	4879303	MAR 25, 2007		NC	MAR 03, 1998
		4572909	AUG 01, 2006		NCE	JUN 25, 1996
		4410520	AUG 11, 2003		NCE	JUL 31, 1997
>ADD>	19155 001	AMMONIUM LACTATE; LAC-HYDRIN	4105783	JAN 15, 1997		
	20541 001	ANASTROZOLE; ARIMIDEX			NCE	DEC 27, 2000

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
>ADD>	19779 001 APRACLONIDINE HYDROCHLORIDE; IOPIDINE	5212196	MAY 18, 2010	U-120		
	19402 001 ASTEMIZOLE; HISMANAL	4219559	APR 03, 2000			
	20259 001 ATOVAQUONE; MEPRON	4981874	AUG 15, 2009	U-69		
	20500 001 ATOVAQUONE; MEPRON	5053432	OCT 01, 2008		NCE	NOV 25, 1997
		4981874	AUG 15, 2009	U-69	NDF	FEB 08, 1998
					NCE	SEP 13, 2000
	20428 001 AZELAIC ACID; AZELEX					
	19851 001 BENAZEPRIL HYDROCHLORIDE; LOTENSIN	4410520	AUG 11, 2003			
	19851 002 BENAZEPRIL HYDROCHLORIDE; LOTENSIN	4410520	AUG 11, 2003			
	19851 003 BENAZEPRIL HYDROCHLORIDE; LOTENSIN	4410520	AUG 11, 2003			
	19851 004 BENAZEPRIL HYDROCHLORIDE; LOTENSIN	4410520	AUG 11, 2003			
	20033 001 BENAZEPRIL HYDROCHLORIDE; LOTENSIN HCT	4410520	AUG 11, 2003			
	20033 002 BENAZEPRIL HYDROCHLORIDE; LOTENSIN HCT	4410520	AUG 11, 2003			
	20033 003 BENAZEPRIL HYDROCHLORIDE; LOTENSIN HCT	4410520	AUG 11, 2003			
	20033 004 BENAZEPRIL HYDROCHLORIDE; LOTENSIN HCT	4410520	AUG 11, 2003			
	18827 001 BETAMETHASONE DIPROPIONATE; LOTRISONE	4298604	OCT 06, 2000			
	19555 001 BETAMETHASONE DIPROPIONATE; DIPROLENE AF	4489071	DEC 09, 2003			
	19716 001 BETAMETHASONE DIPROPIONATE; DIPROLENE	4775529	MAY 21, 2007			
>ADD>	19270 001 BETAXOLOL HYDROCHLORIDE; BETOPTIC	4342783	JUN 30, 2000			
>DLT>	19270 001 BETAXOLOL HYDROCHLORIDE; BETOPTIC	4342783	AUG 03, 1999			
>ADD>	19845 001 BETAXOLOL HYDROCHLORIDE; BETOPTIC S	4342783	JUN 30, 2000			
>DLT>	19845 001 BETAXOLOL HYDROCHLORIDE; BETOPTIC S	4342783	AUG 03, 1999			
	20498 001 BICALUTAMIDE; CASODEX				NCE	OCT 04, 2000
	18644 001 BUPROPION HYDROCHLORIDE; WELLBUTRIN	4507323	JUL 25, 2004			
		4438138	DEC 06, 2002			
		4435449	MAY 14, 2001			
		4425363	MAY 14, 2001			
		4393078	MAR 15, 2002			
		4347257	OCT 09, 1999			
	18644 002 BUPROPION HYDROCHLORIDE; WELLBUTRIN	4507323	JUL 25, 2004			
		4438138	DEC 06, 2002			
		4435449	MAY 14, 2001			
		4425363	MAY 14, 2001			
		4393078	MAR 15, 2002			
		4347257	OCT 09, 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
18644 003	BUPROPION HYDROCHLORIDE; WELLBUTRIN	4507323	JUL 25, 2004			
		4438138	DEC 06, 2002			
		4435449	MAY 14, 2001			
		4425363	MAY 14, 2001			
		4393078	MAR 15, 2002			
		4347257	OCT 09, 1999			
18731 001	BUSPIRONE HYDROCHLORIDE; BUSPAR	4182763	MAY 22, 2000			
18731 002	BUSPIRONE HYDROCHLORIDE; BUSPAR	4182763	MAY 22, 2000			
19215 001	BUTOCONAZOLE NITRATE; FEMSTAT	4078071	JUL 28, 1997			
19359 001	BUTOCONAZOLE NITRATE; FEMSTAT	4078071	JUL 28, 1997			
20313 002	CALCITONIN, SALMON; MIACALCIN				NDF	AUG 17, 1998
>ADD>	18469 001	4443432	OCT 05, 2001			
>DLT>	18469 001	4443432	APR 07, 2001			
	18343 001	4105776	FEB 13, 1996			
	18343 002	4105776	FEB 13, 1996			
	18343 003	4105776	FEB 13, 1996			
>ADD>	18343 004	5238924	AUG 24, 2010	U-92		
>ADD>		4258027	MAR 26, 1999			
>ADD>		4215104	MAR 26, 1999			
>ADD>		4105776	FEB 13, 1996			
	18343 005	4105776	FEB 13, 1996			
	18343 006	4105776	FEB 13, 1996			
>ADD>	18343 007	5238924	AUG 24, 2010	U-92		
>ADD>		4258027	MAR 26, 1999			
>ADD>		4215104	MAR 26, 1999			
>ADD>		4105776	FEB 13, 1996			
	18709 001	4217347	DEC 27, 1997			
		4105776	FEB 13, 1996			
	18709 002	4217347	DEC 27, 1997			
		4105776	FEB 13, 1996			
	18709 003	4217347	DEC 27, 1997			
		4105776	FEB 13, 1996			
	18709 004	4217347	DEC 27, 1997			
		4105776	FEB 13, 1996			
	19856 001	4900755	JUN 16, 2006			
		4832957	JUN 16, 2006			

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APPL/PROD NUMBER	INGREDIENT NAME; TRADE NAME	PATENT NUMBER	PATENT EXPIRES	USE CODE	EXCLUS CODE	EXCLUS EXPIRES
19856 002	CARBIDOPA; SINEMET CR	4900755	JUN 16, 2006			
		4832957	JUN 16, 2006			
>ADD>	20297 001 CARVEDILOL; COREG	4503067	MAR 05, 2002	U-3	NCE	SEP 14, 2000
>ADD>	20297 002 CARVEDILOL; COREG	4503067	MAR 05, 2002	U-3	NCE	SEP 14, 2000
>ADD>	20297 003 CARVEDILOL; COREG	4503067	MAR 05, 2002	U-3	NCE	SEP 14, 2000
>ADD>	19835 001 CETIRIZINE HYDROCHLORIDE; ZYRTEC				NCE	DEC 08, 2000
>ADD>	19835 002 CETIRIZINE HYDROCHLORIDE; ZYRTEC				NCE	DEC 08, 2000
	20044 001 CETYL ALCOHOL; EXOSURF NEONATAL	5110806	MAY 02, 2006			
		4312860	AUG 20, 2003			
	18663 001 CHYMOPAPAIN; CHYMODIACTIN	4439423	MAY 13, 2001			
	18663 002 CHYMOPAPAIN; CHYMODIACTIN	4439423	MAY 13, 2001			
	20238 001 CIMETIDINE; TAGAMET HB				NS	JUN 19, 1998
					I-137	NOV 15, 1998
	19847 001 CIPROFLOXACIN; CIPRO	4670444	DEC 09, 2003			
	19857 001 CIPROFLOXACIN; CIPRO IN DEXTROSE 5%	4670444	DEC 09, 2003			
	19858 001 CIPROFLOXACIN; CIPRO IN SODIUM CHLORIDE 0.9%	4670444	DEC 09, 2003			
	19537 002 CIPROFLOXACIN HYDROCHLORIDE; CIPRO	5286754	FEB 15, 2011			
		4670444	DEC 09, 2003	U-36		
	19537 003 CIPROFLOXACIN HYDROCHLORIDE; CIPRO	5286754	FEB 15, 2011			
		4670444	DEC 09, 2003	U-36		
	19537 004 CIPROFLOXACIN HYDROCHLORIDE; CIPRO	5286754	FEB 15, 2011			
		4670444	DEC 09, 2003	U-36		
		4670444	DEC 09, 2003			
>ADD>	19992 001 CIPROFLOXACIN HYDROCHLORIDE; CILOXAN	4670444	DEC 09, 2003			
>DLT>	19992 001 CIPROFLOXACIN HYDROCHLORIDE; CILOXAN	4670444	OCT 01, 2002			
	20398 001 CISAPRIDE MONOHYDRATE; PROPULSID				NCE	JUL 29, 1998
>ADD>	20551 001 CISATRACURIUM BESYLATE; NIMBEX	5453510	SEP 26, 2012		NE	DEC 15, 1998
>ADD>	20551 002 CISATRACURIUM BESYLATE; NIMBEX PRESERVATIVE FREE	5453510	SEP 26, 2012		NE	DEC 15, 1998
>ADD>	20551 003 CISATRACURIUM BESYLATE; NIMBEX PRESERVATIVE FREE	5453510	SEP 26, 2012		NE	DEC 15, 1998
	18891 001 CLONIDINE; CATAPRES-TTS-1	4559222	MAY 04, 2003			
		4201211	JUL 12, 1997			
	18891 002 CLONIDINE; CATAPRES-TTS-2	4559222	MAY 04, 2003			
		4201211	JUL 12, 1997			
	18891 003 CLONIDINE; CATAPRES-TTS-3	4559222	MAY 04, 2003			
		4201211	JUL 12, 1997			
	20222 001 COLESTIPOL HYDROCHLORIDE; COLESTID				NDF	JUL 19, 1997
	12142 006 CYCLOPHOSPHAMIDE; LYOPHILIZED CYTOXAN	4537883	NOV 12, 2002			
	12142 007 CYCLOPHOSPHAMIDE; LYOPHILIZED CYTOXAN	4537883	NOV 12, 2002			

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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
12142 008	CYCLOPHOSPHAMIDE; LYOPHILIZED CYTOXAN	4537883	NOV 12, 2002			
12142 009	CYCLOPHOSPHAMIDE; LYOPHILIZED CYTOXAN	4537883	NOV 12, 2002			
12142 010	CYCLOPHOSPHAMIDE; LYOPHILIZED CYTOXAN	4537883	NOV 12, 2002			
20287 001	DALTEPARIN SODIUM; FRAGMIN	4303651	JAN 04, 2000		NCE	DEC 22, 1999
19849 001	DAPIPRAZOLE HYDROCHLORIDE; REV-EYES	4252721	FEB 07, 2003			
20071 001	DESOGESTREL; DESOGEN	3927046	NOV 06, 1996			
20071 002	DESOGESTREL; DESOGEN	3927046	NOV 06, 1996			
20301 001	DESOGESTREL; ORTHO-CEPT	3927046	NOV 06, 1996			
20301 002	DESOGESTREL; ORTHO-CEPT	3927046	NOV 06, 1996			
20212 001	DEXRAZOXANE HYDROCHLORIDE; ZINECARD				NCE	MAY 26, 2000
					ODE	MAY 26, 2002
20212 002	DEXRAZOXANE HYDROCHLORIDE; ZINECARD				NCE	MAY 26, 2000
					ODE	MAY 26, 2002
20062 001	DILTIAZEM HYDROCHLORIDE; CARDIZEM CD	5470584	MAY 20, 2011			
		5439689	AUG 08, 2012	U-107		
		5286497	MAY 20, 2011			
20062 002	DILTIAZEM HYDROCHLORIDE; CARDIZEM CD	5470584	MAY 20, 2011			
		5439689	AUG 08, 2012	U-107		
		5286497	MAY 20, 2011			
20062 003	DILTIAZEM HYDROCHLORIDE; CARDIZEM CD	5470584	MAY 20, 2011			
		5439689	AUG 08, 2012	U-107		
		5286497	MAY 20, 2011			
20062 004	DILTIAZEM HYDROCHLORIDE; CARDIZEM CD	5470584	MAY 20, 2011			
		5439689	AUG 08, 2012	U-107		
		5286497	MAY 20, 2011			
20092 001	DILTIAZEM HYDROCHLORIDE; DILACOR XR	4839177	DEC 09, 2006		I-133	OCT 15, 1995
20092 002	DILTIAZEM HYDROCHLORIDE; DILACOR XR	4839177	DEC 09, 2006		I-133	OCT 15, 1995
20092 003	DILTIAZEM HYDROCHLORIDE; DILACOR XR	4839177	DEC 09, 2006		I-133	OCT 15, 1995
20401 005	DILTIAZEM HYDROCHLORIDE; TIAZAC				NS	SEP 11, 1998
20411 001	DINOPROSTONE; CERVIDIL	5269321	DEC 14, 2010	U-110		
		4931288	JAN 16, 2007		NDF	MAR 30, 1998
18723 001	DIVALPROEX SODIUM; DEPAKOTE	4988731	JAN 29, 2008		I-132	MAY 26, 1998
18723 002	DIVALPROEX SODIUM; DEPAKOTE	4988731	JAN 29, 2008		I-132	MAY 26, 1998
18723 003	DIVALPROEX SODIUM; DEPAKOTE	4988731	JAN 29, 2008		I-132	MAY 26, 1998
20408 001	DORZOLAMIDE HYDROCHLORIDE; TRUSOPT	4797413	DEC 12, 2004	U-103	NCE	DEC 09, 1999
		4619939	OCT 28, 2003	U-104		
19946 001	DOXACURIUM CHLORIDE; NUROMAX				I-121	DEC 08, 1997

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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
19668 001	DOXAZOSIN MESYLATE; CARDURA	4188390	OCT 18, 2000		I-96	FEB 06, 1998
19668 002	DOXAZOSIN MESYLATE; CARDURA	4188390	OCT 18, 2000		I-96	FEB 06, 1998
19668 003	DOXAZOSIN MESYLATE; CARDURA	4188390	OCT 18, 2000		I-96	FEB 06, 1998
19668 004	DOXAZOSIN MESYLATE; CARDURA	4188390	OCT 18, 2000		I-96	FEB 06, 1998
20126 001	DOXEPIN HYDROCHLORIDE; ZONALON	4395420	DEC 09, 2001	U-95		
19221 003	ENALAPRIL MALEATE; VASERETIC	4472380	SEP 18, 2001			
		4374829	FEB 22, 2000		NS	JUL 12, 1998
19616 004	ENOXACIN; PENETREX	4442101	FEB 04, 2002			
19616 005	ENOXACIN; PENETREX	4442101	FEB 04, 2002			
>ADD>	20164 001	ENOXAPARIN SODIUM; LOVENOX			I-118	MAR 09, 1998
	20530 001	EPINEPHRINE; IONTOCAINE			NP	DEC 21, 1998
>ADD>	20444 001	EPOPROSTENOL SODIUM; FLOLAN			NCE	SEP 20, 2000
					ODE	SEP 20, 2002
>ADD>	20444 002	EPOPROSTENOL SODIUM; FLOLAN			NCE	SEP 20, 2000
					ODE	SEP 20, 2002
18418 001	ERGOLOID MESYLATES; HYDERGINE	4138565	MAY 26, 1996			
18706 001	ERGOLOID MESYLATES; HYDERGINE LC	4366145	JUN 24, 2001			
19081 002	ESTRADIOL; ESTRADERM	4379454	FEB 17, 2001			
19081 003	ESTRADIOL; ESTRADERM	4379454	FEB 17, 2001			
20323 001	ESTRADIOL; VIVELLE	5300291	APR 05, 2011		NS	OCT 28, 1997
		4994278	MAR 04, 2008			
		4994267	MAR 04, 2008			
		4814168	MAR 04, 2008			
20323 002	ESTRADIOL; VIVELLE	5300291	APR 05, 2011			
		4994278	MAR 04, 2008			
		4994267	MAR 04, 2008			
		4814168	MAR 04, 2008			
20323 003	ESTRADIOL; VIVELLE	5300291	APR 05, 2011		NS	OCT 28, 1997
		4994278	MAR 04, 2008			
		4994267	MAR 04, 2008			
		4814168	MAR 04, 2008			
20323 004	ESTRADIOL; VIVELLE	5300291	APR 05, 2011			
		4994278	MAR 04, 2008			
		4994267	MAR 04, 2008			
		4814168	MAR 04, 2008			
20375 001	ESTRADIOL; CLIMARA	5223261	JUN 29, 2010		D-26	DEC 22, 1997
20375 002	ESTRADIOL; CLIMARA	5223261	JUN 29, 2010		D-26	DEC 22, 1997

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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
86069 001	ESTRADIOL; ESTRACE	4436738	MAR 15, 2002			
20303 001	ESTROGENS, CONJUGATED; PREMPRO (PREMARIN;CYCRIN 14/14)	4826831	MAY 02, 2006	U-102	NP	DEC 30, 1997
20527 001	ESTROGENS, CONJUGATED; PREMPRO 14/14	4826831	MAY 02, 2006	U-102		
18977 001	ETHINYL ESTRADIOL; TRI-NORINYL 21-DAY	4390531	AUG 10, 2001			
18977 002	ETHINYL ESTRADIOL; TRI-NORINYL 28-DAY	4390531	AUG 10, 2001			
18985 001	ETHINYL ESTRADIOL; ORTHO-NOVUM 7/7/7-21	4628051	SEP 26, 2003			
		4616006	SEP 26, 2003			
		4544554	SEP 26, 2003			
		4530839	SEP 26, 2003			
18985 002	ETHINYL ESTRADIOL; ORTHO-NOVUM 7/7/7-28	4628051	SEP 26, 2003			
		4616006	SEP 26, 2003			
		4544554	SEP 26, 2003			
		4530839	SEP 26, 2003			
19697 001	ETHINYL ESTRADIOL; ORTHO TRI-CYCLEN	4628051	SEP 26, 2003	U-66		
		4616006	SEP 26, 2003			
		4544554	SEP 26, 2003	U-66		
		4530839	SEP 26, 2003	U-112		
19697 002	ETHINYL ESTRADIOL; ORTHO TRI-CYCLEN	4628051	SEP 26, 2003	U-66		
		4616006	SEP 26, 2003			
		4544554	SEP 26, 2003	U-66		
		4530839	SEP 26, 2003	U-112		
>ADD>	20363 003	FAMCICLOVIR; FAMVIR			I-138	DEC 11, 1998
>ADD>					NCE	JUN 29, 1999
	19462 001	FAMOTIDINE; PEPCID	4283408	OCT 15, 2000		
	19462 002	FAMOTIDINE; PEPCID	4283408	OCT 15, 2000		
	19510 001	FAMOTIDINE; PEPCID	4283408	OCT 15, 2000		
	19527 001	FAMOTIDINE; PEPCID	4283408	OCT 15, 2000		
	20249 001	FAMOTIDINE; PEPCID	4283408	OCT 15, 2000		
	20325 001	FAMOTIDINE; PEPCID AC	4283408	OCT 15, 2000	NS	APR 28, 1998
	20189 001	FELBAMATE; FELBATOL	5082861	SEP 26, 2009	U-83	
		4978680	SEP 26, 2009	U-83		
	20189 002	FELBAMATE; FELBATOL	5082861	SEP 26, 2009	U-83	
		4978680	SEP 26, 2009	U-83		
	20189 003	FELBAMATE; FELBATOL	5082861	SEP 26, 2009	U-83	
		4978680	SEP 26, 2009	U-83		
	19834 001	FELODIPINE; PLENDIL	4803081	APR 03, 2007		
		4264611	JUN 19, 2001	U-3	NCE	JUL 25, 1996

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APPL/PROD NUMBER	INGREDIENT NAME; TRADE NAME	PATENT NUMBER	PATENT EXPIRES	USE CODE	EXCLUS CODE	EXCLUS EXPIRES
19834 002	FELODIPINE; PLENDIL	4803081	APR 03, 2007			
19834 004	FELODIPINE; PLENDIL	4264611	JUN 19, 2001	U-3	NCE	JUL 25, 1996
		4803081	APR 03, 2007			
		4264611	JUN 19, 2001	U-3	NCE	JUL 25, 1996
19813 001	FENTANYL; DURAGESIC	4588580	JUL 23, 2004	U-43		
19813 002	FENTANYL; DURAGESIC	4588580	JUL 23, 2004	U-43		
19813 003	FENTANYL; DURAGESIC	4588580	JUL 23, 2004	U-43		
19813 004	FENTANYL; DURAGESIC	4588580	JUL 23, 2004	U-43		
19960 001	FLOSEQUINAN; MANOPLAX	4302460	MAR 24, 2000			
19960 002	FLOSEQUINAN; MANOPLAX	4302460	MAR 24, 2000			
19960 003	FLOSEQUINAN; MANOPLAX	4302460	MAR 24, 2000			
19960 004	FLOSEQUINAN; MANOPLAX	4302460	MAR 24, 2000			
19949 001	FLUCONAZOLE; DIFLUCAN	4416682	JUN 02, 2001			
		4404216	JAN 29, 2004			
19949 002	FLUCONAZOLE; DIFLUCAN	4416682	JUN 02, 2001			
		4404216	JAN 29, 2004			
19949 003	FLUCONAZOLE; DIFLUCAN	4416682	JUN 02, 2001			
		4404216	JAN 29, 2004			
19950 001	FLUCONAZOLE; DIFLUCAN	4416682	JUN 02, 2001			
		4404216	JAN 29, 2004			
20090 001	FLUCONAZOLE; DIFLUCAN	4416682	JUN 02, 2001			
		4404216	JAN 29, 2004			
20090 002	FLUCONAZOLE; DIFLUCAN	4416682	JUN 02, 2001			
		4404216	JAN 29, 2004			
20322 001	FLUCONAZOLE; DIFLUCAN	4416682	JUN 02, 2001			
		4404216	JAN 29, 2004			
20073 001	FLUMAZENIL; ROMAZICON	4316839	OCT 10, 2004			
18148 001	FLUNISOLIDE; NASALIDE	4933168	JUN 12, 2007			
18340 001	FLUNISOLIDE; AEROBID	4933168	JUN 12, 2007			
20409 001	FLUNISOLIDE; NASAREL	4983595	MAY 22, 2006			
		4933168	JUN 12, 2007			
		4782047	MAY 22, 2006			
19452 001	FLUOCINOLONE ACETONIDE; DERMA-SMOOTH/FS				I-122	FEB 16, 1998
19957 001	FLUTICASONE PROPIONATE; CUTIVATE	4335121	NOV 14, 2003			
19958 001	FLUTICASONE PROPIONATE; CUTIVATE	4335121	NOV 14, 2003			
20121 001	FLUTICASONE PROPIONATE; FLONASE	4335121	NOV 14, 2003		NCE NDF	DEC 14, 1995 OCT 19, 1997

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APPL/PROD NUMBER	INGREDIENT NAME; TRADE NAME	PATENT NUMBER	PATENT EXPIRES	USE CODE	EXCLUS CODE	EXCLUS EXPIRES	
20261 001	FLUVASTATIN SODIUM; LESCOL	5354772	OCT 11, 2011	U-109	NCE	DEC 31, 1998	
20261 002	FLUVASTATIN SODIUM; LESCOL	5354772	OCT 11, 2011	U-109	NCE	DEC 31, 1998	
20068 001	FOSCARNET SODIUM; FOSCAVIR	4339445	JUL 29, 1997	U-64	I-127	JUN 16, 1998	
19915 002	FOSINOPRIL SODIUM; MONOPRIL	5006344	JUL 10, 2009				
		4384123	DEC 04, 2000		I-92	MAY 02, 1998	
		4337201	DEC 04, 2002				
19915 003	FOSINOPRIL SODIUM; MONOPRIL	5006344	JUL 10, 2009				
		4384123	DEC 04, 2000		I-92	MAY 02, 1998	
		4337201	DEC 04, 2002				
19915 004	FOSINOPRIL SODIUM; MONOPRIL	5006344	JUL 10, 2009				
		4384123	DEC 04, 2000		I-92	MAY 02, 1998	
		4337201	DEC 04, 2002		NCE	MAY 16, 1996	
20286 001	FOSINOPRIL SODIUM; MONOPRIL-HCT	5006344	JUL 10, 2009				
		4384123	DEC 04, 2000				
		4337201	DEC 04, 2002				
20286 002	FOSINOPRIL SODIUM; MONOPRIL-HCT	5006344	JUL 10, 2009				
		4384123	DEC 04, 2000				
		4337201	DEC 04, 2002				
20235 001	GABAPENTIN; NEURONTIN	4087544	MAY 02, 1996	U-86			
20235 002	GABAPENTIN; NEURONTIN	4087544	MAY 02, 1996	U-86			
20235 003	GABAPENTIN; NEURONTIN	4087544	MAY 02, 1996	U-86			
19596 001	GADOPENTETATE DIMEGLUMINE; MAGNEVIST	5362475	NOV 08, 2011				
20460 001	GANCICLOVIR; CYTOVENE	4507305	OCT 19, 1999	U-64	NDF	DEC 22, 1997	
>ADD>		4355032	JUN 23, 2003	U-64	I-140	OCT 27, 1998	
19661 001	GANCICLOVIR SODIUM; CYTOVENE	4507305	MAY 21, 2001	U-35			
		4423050	MAY 21, 2001	U-34			
		4355032	JUN 23, 2003	U-64			
>ADD>	20496 001	GLIMEPIRIDE; AMARYL	4379785	DEC 17, 2000	U-118	NCE	NOV 30, 2000
>ADD>	20496 002	GLIMEPIRIDE; AMARYL	4379785	DEC 17, 2000	U-118	NCE	NOV 30, 2000
>ADD>	20496 003	GLIMEPIRIDE; AMARYL	4379785	DEC 17, 2000	U-118	NCE	NOV 30, 2000
	20329 001	GLIPIZIDE; GLUCOTROL XL	5091190	JUN 18, 2008	U-111		
		5082668	SEP 16, 2003				
		5024843	SEP 05, 2009				
		4612008	SEP 16, 2003				
		4327725	NOV 25, 2000		NDF	APR 26, 1997	

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APPL/PROD NUMBER	INGREDIENT NAME; TRADE NAME	PATENT NUMBER	PATENT EXPIRES	USE CODE	EXCLUS CODE	EXCLUS EXPIRES
20329 002	GLIPIZIDE; GLUCOTROL XL	5091190	JUN 18, 2008	U-111		
		5082668	SEP 16, 2003			
		5024843	SEP 05, 2009			
		4612008	SEP 16, 2003			
		4327725	NOV 25, 2000		NDF	APR 26, 1997
>ADD> 19726 001	GOSERELIN ACETATE; ZOLADEX	4100274	APR 22, 1999		I-139	DEC 18, 1998
20305 001	GRANISETRON HYDROCHLORIDE; KYTRIL	4886808	DEC 12, 2006	U-105	NCE	DEC 29, 1998
					NDF	MAR 16, 1998
19059 001	HYDROCHLOROTHIAZIDE; INDERIDE LA 80/50	4138475	SEP 14, 1997			
19059 002	HYDROCHLOROTHIAZIDE; INDERIDE LA 120/50	4138475	SEP 14, 1997			
19059 003	HYDROCHLOROTHIAZIDE; INDERIDE LA 160/50	4138475	SEP 14, 1997			
19129 001	HYDROCHLOROTHIAZIDE; MAXZIDE	4444769	JUL 27, 2002			
19129 003	HYDROCHLOROTHIAZIDE; MAXZIDE-25	4444769	JUL 27, 2002			
20387 001	HYDROCHLOROTHIAZIDE; HYZAAR	5153197	OCT 06, 2009	U-3	NC	APR 28, 1998
		5138069	AUG 11, 2009		NCE	APR 14, 2000
19842 001	IBUPROFEN; CHILDREN'S MOTRIN	5374659	DEC 20, 2011		I-123	MAR 24, 1998
20135 001	IBUPROFEN; MOTRIN	5320855	JUN 14, 2011		I-123	MAR 24, 1998
		5215755	JUN 01, 2010		NDF	NOV 16, 1997
20135 002	IBUPROFEN; MOTRIN	5320855	JUN 14, 2011		I-123	MAR 24, 1998
		5215755	JUN 01, 2010		NDF	NOV 16, 1997
20418 001	IBUPROFEN; MOTRIN				I-123	MAR 24, 1998
20516 001	IBUPROFEN; CHILDREN'S MOTRIN	5374659	DEC 20, 2011		NP	JUN 16, 1998
>ADD> 20491 001	IBUTILIDE FUMARATE; CORVERT	5155268	OCT 13, 2009		NCE	DEC 28, 2000
18185 001	INDOMETHACIN; INDOCIN SR	4173626	DEC 11, 1998			
20084 001	IOBENGUANE SULFATE I 131; IOBENGUANE SULFATE I 131				ODE	MAR 25, 2001
18956 007	IOHEXOL; OMNIPaque 70	4396597	JUL 14, 1998			
		4250113	DEC 26, 1999			
18735 001	IOPAMIDOL; ISOVUE-M 200	4001323	NOV 24, 1997			
18735 002	IOPAMIDOL; ISOVUE-300	4001323	NOV 24, 1997			
18735 003	IOPAMIDOL; ISOVUE-370	4001323	NOV 24, 1997		D-28	MAY 15, 1998
18735 004	IOPAMIDOL; ISOVUE-M 300	4001323	NOV 24, 1997			
18735 007	IOPAMIDOL; ISOVUE-250	4001323	NOV 24, 1997			
20327 001	IOPAMIDOL; ISOVUE-200	4001323	NOV 24, 1997			
20327 002	IOPAMIDOL; ISOVUE-250	4001323	NOV 24, 1997			
20327 003	IOPAMIDOL; ISOVUE-300	4001323	NOV 24, 1997			
20327 004	IOPAMIDOL; ISOVUE-370	4001323	NOV 24, 1997			
20220 001	IOPROMIDE; ULTRAVIST 370	4364921	DEC 21, 1999	U-113	NCE	MAY 10, 2000

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APPL/PROD NUMBER	INGREDIENT NAME; TRADE NAME	PATENT NUMBER	PATENT EXPIRES	USE CODE	EXCLUS CODE	EXCLUS EXPIRES
20220 002	IOPROMIDE; ULTRAVIST 300	4364921	DEC 21, 1999	U-113	NCE	MAY 10, 2000
20220 003	IOPROMIDE; ULTRAVIST 240	4364921	DEC 21, 1999	U-113	NCE	MAY 10, 2000
20220 004	IOPROMIDE; ULTRAVIST 150	4364921	DEC 21, 1999	U-113	NCE	MAY 10, 2000
19710 005	IOVERSOL; OPTIRAY 350				I-131	JUN 21, 1998
18905 002	IOXAGLATE MEGLUMINE; HEXABRIX	4014986	MAY 20, 1997			
>ADD>	20316 001				NCE	DEC 21, 2000
>ADD>	20316 002				NCE	DEC 21, 2000
>ADD>	20393 001	4385048	MAY 24, 2000	U-119	NDF	OCT 20, 1998
>ADD>	20394 001	4385048	MAY 24, 2000	U-119	NDF	OCT 20, 1998
	20225 003				NDF	AUG 12, 1996
					NCE	DEC 30, 1996
	20336 001	4816263	OCT 02, 2007	U-3		
	20336 002	4816263	OCT 02, 2007	U-3		
	20083 001	4267179	JUN 23, 2000			
	19816 002				NDF	SEP 24, 1996
	19816 003				NDF	SEP 24, 1996
>ADD>	20429 001				NS	OCT 06, 1998
>ADD>	20499 001				NS	OCT 06, 1998
	19645 001	4089969	JUL 14, 1998	U-55		
	19698 001	4089969	JUL 14, 1998	U-55	NR	DEC 07, 1997
	19698 002	4089969	JUL 14, 1998	U-55	NR	DEC 07, 1997
	19700 001	4454151	MAR 22, 2002	U-75		
		4089969	JUL 14, 1998	U-55		
		4328213	NOV 28, 1999			
	18686 001				NCE	NOV 17, 2000
	20564 001				NCE	NOV 17, 2000
	20596 001				NCE	NOV 17, 2000
	20241 001	4602017	JUL 22, 2003	U-106	NCE	DEC 27, 1999
	20241 002	4602017	JUL 22, 2003	U-106	NCE	DEC 27, 1999
	20241 003	4602017	JUL 22, 2003	U-106	NCE	DEC 27, 1999
	20241 004	4602017	JUL 22, 2003	U-106	NCE	DEC 27, 1999
	20241 005	4602017	JUL 22, 2003	U-106	NCE	DEC 27, 1999
	20241 006	4602017	JUL 22, 2003	U-106	NCE	DEC 27, 1999
	20406 001	5093132	SEP 03, 2008			
		5045321	SEP 03, 2008			
		5026560	JUN 25, 2008			
		4689333	JUL 29, 2005			
		4628098	JUL 29, 2005		NCE	MAY 10, 2000

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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
20406 002	LANSOPRAZOLE; PREVACID	5093132	SEP 03, 2008			
		5045321	SEP 03, 2008			
		5026560	JUN 25, 2008			
		4689333	JUL 29, 2005			
		4628098	JUL 29, 2005		NCE	MAY 10, 2000
>ADD> 19732 001	LEUPROLIDE ACETATE; LUPRON DEPOT	5476663	APR 17, 2007			
		5330767	NOV 01, 2004			
		4917893	NOV 01, 2004			
		4728721	MAY 01, 2006			
		4677191	JUL 03, 2005			
		4652441	NOV 01, 2004			
>ADD> 20011 001	LEUPROLIDE ACETATE; LUPRON DEPOT	5476663	APR 17, 2007			
		5330767	NOV 01, 2004			
		4917893	NOV 01, 2004			
		4728721	MAY 01, 2006			
		4677191	JUL 03, 2005			
		4652441	NOV 01, 2004			
		4005063	JAN 25, 1996		I-119	MAR 30, 1998
>ADD> 20263 001	LEUPROLIDE ACETATE; LUPRON	5476663	APR 17, 2007			
		5330767	NOV 01, 2004			
		4917893	NOV 01, 2004			
		4728721	MAY 01, 2006			
		4677191	JUL 03, 2005			
		4652441	NOV 01, 2004			
>ADD> 20263 002	LEUPROLIDE ACETATE; LUPRON DEPOT-PED	5476663	APR 17, 2007			
		5330767	NOV 01, 2004			
		4917893	NOV 01, 2004			
		4728721	MAY 01, 2006			
		4677191	JUL 03, 2005			
		4652441	NOV 01, 2004			
>ADD> 20263 003	LEUPROLIDE ACETATE; LUPRON DEPOT-PED	5476663	APR 17, 2007			
		5330767	NOV 01, 2004			
		4917893	NOV 01, 2004			
		4728721	MAY 01, 2006			
		4677191	JUL 03, 2005			
		4652441	NOV 01, 2004			

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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>	20263 004 LEUPROLIDE ACETATE; LUPRON DEPOT-PED	5476663	APR 17, 2007			
		5330767	NOV 01, 2004			
		4917893	NOV 01, 2004			
		4728721	MAY 01, 2006			
		4677191	JUL 03, 2005			
		4652441	NOV 01, 2004			
>ADD>	20263 005 LEUPROLIDE ACETATE; LUPRON DEPOT-PED	5476663	APR 17, 2007			
		5330767	NOV 01, 2004			
		4917893	NOV 01, 2004			
		4849228	JUL 18, 2006			
		4728721	MAY 01, 2006			
		4677191	JUL 03, 2005			
		4652441	NOV 01, 2004			
		4005063	JAN 25, 1996			
>ADD>	20263 006 LEUPROLIDE ACETATE; LUPRON DEPOT-PED	5476663	APR 17, 2007			
		5330767	NOV 01, 2004			
		4917893	NOV 01, 2004			
		4849228	JUL 18, 2006			
		4728721	MAY 01, 2006			
		4677191	JUL 03, 2005			
		4652441	NOV 01, 2004			
		4005063	JAN 25, 1996			
>ADD>	20517 001 LEUPROLIDE ACETATE; LUPRON DEPOT	5476663	APR 17, 2007		NP	DEC 22, 1998
>ADD>		5330767	NOV 01, 2004			
>ADD>		4954298	NOV 01, 2004			
>ADD>		4917893	NOV 01, 2004			
>ADD>		4849228	JUL 18, 2006			
>ADD>		4728721	MAY 01, 2006			
>ADD>		4677191	JUL 03, 2005			
>ADD>		4652441	NOV 01, 2004			
	18027 001 LITHIUM CARBONATE; LITHOBID	4264573	MAY 21, 1999			
>ADD>	20191 001 LODOXAMIDE TROMETHAMINE; ALOMIDE	5457126	OCT 10, 2012	U-117	NCE	SEP 23, 1998
	20013 001 LOMEFLOXACIN HYDROCHLORIDE; MAXAQUIN	4528287	FEB 21, 2006	U-36		
	19658 001 LORATADINE; CLARITIN	4282233	JUN 19, 2002	U-77	I-136	SEP 20, 1998
	19670 001 LORATADINE; CLARITIN-D	4282233	JUN 19, 2002		NCE	APR 12, 1998
	20386 001 LOSARTAN POTASSIUM; COZAAR	5153197	OCT 06, 2009	U-3		
		5138069	AUG 11, 2009		NCE	APR 14, 2000

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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
20386 002	LOSARTAN POTASSIUM; COZAAR	5153197	OCT 06, 2009	U-3		
		5138069	AUG 11, 2009		NCE	APR 14, 2000
19643 002	LOVASTATIN; MEVACOR	4231938	JUN 15, 2001		I-117	FEB 08, 1998
19643 003	LOVASTATIN; MEVACOR	4231938	JUN 15, 2001		I-117	FEB 08, 1998
19643 004	LOVASTATIN; MEVACOR	4231938	JUN 15, 2001		I-117	FEB 08, 1998
19940 001	MASOPROCOL; ACTINEX	4695590	SEP 04, 2006		NCE	SEP 04, 1997
19591 001	MEFLOQUINE HYDROCHLORIDE; LARIAM	4579855	OCT 01, 2004			
20207 001	MELPHALAN HYDROCHLORIDE; ALKERAN	4997651	NOV 18, 2008			
19884 001	MESNA; MESNEX	4220660	MAR 06, 2001			
19600 001	METHOXSALEN; OXSORALEN-ULTRA	4454152	DEC 21, 2001			
18029 001	METHYLPHENIDATE HYDROCHLORIDE; RITALIN-SR	4137300	AUG 20, 1996			
19962 001	METOPROLOL SUCCINATE; TOPROL-XL	5081154	JAN 14, 2009			
		5001161	MAR 19, 2008			
		4957745	SEP 18, 2007	U-107	NE	JAN 10, 1995
19962 002	METOPROLOL SUCCINATE; TOPROL-XL	5081154	JAN 14, 2009			
		5001161	MAR 19, 2008			
		4957745	SEP 18, 2007	U-107	NE	JAN 10, 1995
19962 003	METOPROLOL SUCCINATE; TOPROL-XL	5081154	JAN 14, 2009			
		5001161	MAR 19, 2008			
		4957745	SEP 18, 2007	U-107	NE	JAN 10, 1995
20531 001	METRONIDAZOLE; METROCREAM				NDF	SEP 20, 1998
18654 001	MIDAZOLAM HYDROCHLORIDE; VERSED	4280957	DEC 20, 1999		I-125	APR 26, 1997
18654 002	MIDAZOLAM HYDROCHLORIDE; VERSED	4280957	DEC 20, 1999		I-125	APR 26, 1997
19268 001	MISOPROSTOL; CYTOTEC	4301146	JUL 29, 2000			
19268 003	MISOPROSTOL; CYTOTEC	4301146	JUL 29, 2000			
20312 001	MOEXIPRIL HYDROCHLORIDE; UNIVASC	4743450	FEB 24, 2007		NCE	APR 19, 2000
20312 002	MOEXIPRIL HYDROCHLORIDE; UNIVASC	4743450	FEB 24, 2007		NCE	APR 19, 2000
19625 001	MOMETASONE FUROATE; ELOCON	4808610	OCT 02, 2006			
19796 001	MOMETASONE FUROATE; ELOCON	4775529	MAY 21, 2007			
20459 001	NALMEFENE HYDROCHLORIDE; REVEX				NCE	APR 17, 2000
20459 002	NALMEFENE HYDROCHLORIDE; REVEX				NCE	APR 17, 2000
18932 001	NALTREXONE HYDROCHLORIDE; REVIA				I-129	DEC 30, 1997
20152 001	NEFAZODONE HYDROCHLORIDE; SERZONE	4338317	MAR 16, 2001			
20152 002	NEFAZODONE HYDROCHLORIDE; SERZONE	4338317	MAR 16, 2001			
20152 003	NEFAZODONE HYDROCHLORIDE; SERZONE	4338317	MAR 16, 2001			
20152 004	NEFAZODONE HYDROCHLORIDE; SERZONE	4338317	MAR 16, 2001			
20152 005	NEFAZODONE HYDROCHLORIDE; SERZONE	4338317	MAR 16, 2001			

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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
20152 006	NEFAZODONE HYDROCHLORIDE; SERZONE	4338317	MAR 16, 2001			
19488 001	NICARDIPINE HYDROCHLORIDE; CARDENE	3985758	FEB 15, 1996			
19488 002	NICARDIPINE HYDROCHLORIDE; CARDENE	3985758	FEB 15, 1996			
19734 001	NICARDIPINE HYDROCHLORIDE; CARDENE	5164405	NOV 17, 2009			
		4880823	NOV 14, 2006			
		3985758	FEB 15, 1996			
20005 001	NICARDIPINE HYDROCHLORIDE; CARDENE SR	5198226	MAR 30, 2010			
		3985758	FEB 15, 1996			
20005 002	NICARDIPINE HYDROCHLORIDE; CARDENE SR	5198226	MAR 30, 2010			
		3985758	FEB 15, 1996			
20005 003	NICARDIPINE HYDROCHLORIDE; CARDENE SR	5198226	MAR 30, 2010			
		3985758	FEB 15, 1996			
20076 001	NICOTINE; HABITROL	4597961	JAN 23, 2005	U-56		
20076 002	NICOTINE; HABITROL	4597961	JAN 23, 2005	U-56		
20076 003	NICOTINE; HABITROL	4597961	JAN 23, 2005	U-56		
20150 001	NICOTINE; NICOTROL	4915950	FEB 12, 2008			
20150 002	NICOTINE; NICOTROL	4915950	FEB 12, 2008			
20150 003	NICOTINE; NICOTROL	4915950	FEB 12, 2008			
20165 001	NICOTINE; NICODERM	5462745	JUN 14, 2008			
		5364630	JUN 14, 2008			
		5344656	JUN 14, 2008			
		5342623	JUN 14, 2008			
		5004610	JUN 14, 2008			
20165 002	NICOTINE; NICODERM	5462745	JUN 14, 2008			
		5364630	JUN 14, 2008			
		5344656	JUN 14, 2008			
		5342623	JUN 14, 2008			
		5004610	JUN 14, 2008			
20165 003	NICOTINE; NICODERM	5462745	JUN 14, 2008			
		5364630	JUN 14, 2008			
		5344656	JUN 14, 2008			
		5342623	JUN 14, 2008			
		5004610	JUN 14, 2008			
19684 001	NIFEDIPINE; PROCARDIA XL	4327725	NOV 25, 2000			
19684 002	NIFEDIPINE; PROCARDIA XL	4327725	NOV 25, 2000			
19684 003	NIFEDIPINE; PROCARDIA XL	4327725	NOV 25, 2000			

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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
20198 001	NIFEDIPINE; ADALAT CC	5264446	NOV 23, 2010			
		4892741	JUN 08, 2008			
20198 002	NIFEDIPINE; ADALAT CC	5264446	NOV 23, 2010			
		4892741	JUN 08, 2008			
20198 003	NIFEDIPINE; ADALAT CC	5264446	NOV 23, 2010			
		4892741	JUN 08, 2008			
20356 001	NISOLDIPINE; SULAR	4892741	JUN 08, 2008			
		4154839	NOV 02, 1996		NCE	FEB 02, 2000
20356 002	NISOLDIPINE; SULAR	4892741	JUN 08, 2008			
		4154839	NOV 02, 1996		NCE	FEB 02, 2000
20356 003	NISOLDIPINE; SULAR	4892741	JUN 08, 2008			
		4154839	NOV 02, 1996		NCE	FEB 02, 2000
20356 004	NISOLDIPINE; SULAR	4892741	JUN 08, 2008			
		4154839	NOV 02, 1996		NCE	FEB 02, 2000
20064 001	NITROFURANTOIN; MACROBID	4798725	JUN 16, 2006			
		4772473	JUN 16, 2006			
20145 001	NITROGLYCERIN; NITRO-DUR	5186938	FEB 16, 2010			
20145 002	NITROGLYCERIN; NITRO-DUR	5186938	FEB 16, 2010			
20145 003	NITROGLYCERIN; NITRO-DUR	5186938	FEB 16, 2010			
20145 004	NITROGLYCERIN; NITRO-DUR	5186938	FEB 16, 2010			
20145 005	NITROGLYCERIN; NITRO-DUR	5186938	FEB 16, 2010			
20145 006	NITROGLYCERIN; NITRO-DUR	5186938	FEB 16, 2010			
19508 001	NIZATIDINE; AXID	4375547	APR 12, 2002			
19508 002	NIZATIDINE; AXID	4375547	APR 12, 2002			
19384 002	NORFLOXACIN; NOROXIN	4639458	JAN 22, 2005			
		4146719	FEB 16, 2000			
19757 001	NORFLOXACIN; CHIBROXIN	4551456	NOV 14, 2003			
		4146719	FEB 16, 2000			
19667 001	OCTREOTIDE ACETATE; SANDOSTATIN	4395403	OCT 21, 2002			
19667 002	OCTREOTIDE ACETATE; SANDOSTATIN	4395403	OCT 21, 2002			
19667 003	OCTREOTIDE ACETATE; SANDOSTATIN	4395403	OCT 21, 2002			
19667 004	OCTREOTIDE ACETATE; SANDOSTATIN	4395403	OCT 21, 2002			
19667 005	OCTREOTIDE ACETATE; SANDOSTATIN	4395403	OCT 21, 2002			
19735 001	OFLOXACIN; FLOXIN	4382892	SEP 02, 2003			
19735 002	OFLOXACIN; FLOXIN	4382892	SEP 02, 2003			
19735 003	OFLOXACIN; FLOXIN	4382892	SEP 02, 2003			
20087 001	OFLOXACIN; FLOXIN IN DEXTROSE 5%	4382892	SEP 02, 2001			

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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
20087 002	OFLOXACIN; FLOXIN	4382892	SEP 02, 2001			
20087 003	OFLOXACIN; FLOXIN	4382892	SEP 02, 2001			
20087 004	OFLOXACIN; FLOXIN IN DEXTROSE 5%	4382892	SEP 02, 2001			
20087 005	OFLOXACIN; FLOXIN IN DEXTROSE 5%	4382892	SEP 02, 2001			
19810 001	OMEPRAZOLE; PRILOSEC	4853230	APR 20, 2007	U-108		
		4786505	APR 20, 2007	U-108	I-130	JUN 22, 1998
		4255431	APR 05, 2001	U-108		
19810 003	OMEPRAZOLE; PRILOSEC	4853230	APR 20, 2007	U-108		
		4786505	APR 20, 2007	U-108		
		4255431	APR 05, 2001	U-108		
20007 001	ONDANSETRON HYDROCHLORIDE; ZOFRAN	4753789	JUN 24, 2006	U-44		
		4695578	JAN 25, 2005		D-20	FEB 02, 1996
20103 001	ONDANSETRON HYDROCHLORIDE; ZOFRAN	4753789	JUN 24, 2006	U-44	D-27	APR 10, 1998
		4695578	JAN 25, 2005		NCE	JAN 04, 1996
					I-9	APR 19, 1998
20103 002	ONDANSETRON HYDROCHLORIDE; ZOFRAN	4753789	JUN 24, 2006	U-44	D-27	APR 10, 1998
		4695578	JAN 25, 2005		NCE	JAN 04, 1996
					I-9	APR 19, 1998
20403 001	ONDANSETRON HYDROCHLORIDE; ZOFRAN	4753789	JUN 24, 2006	U-44	D-20	FEB 02, 1996
		4695578	JAN 25, 2005		NCE	JAN 04, 1996
					I-9	AUG 13, 1996
19828 001	OXICONAZOLE NITRATE; OXISTAT	4124767	DEC 27, 1996			
20209 001	OXICONAZOLE NITRATE; OXISTAT	4124767	DEC 27, 1996			
>ADD>	20553 001	OXYCODONE HYDROCHLORIDE; OXYCONTIN	5266331	FEB 05, 2008		
>ADD>			4970075	NOV 13, 2007		
>ADD>			4861598	AUG 29, 2006		
>ADD>	20553 002	OXYCODONE HYDROCHLORIDE; OXYCONTIN	5266331	FEB 05, 2008		
>ADD>			4970075	NOV 13, 2007		
>ADD>			4861598	AUG 29, 2006		
>ADD>	20553 003	OXYCODONE HYDROCHLORIDE; OXYCONTIN	5266331	FEB 05, 2008		
>ADD>			4970075	NOV 13, 2007		
>ADD>			4861598	AUG 29, 2006		
>ADD>	20036 001	PAMIDRONATE DISODIUM; ARELIA	4711880	JUL 29, 2005	I-135	SEP 01, 1998
>ADD>			3962432	JAN 09, 1998	U-53	NCE OCT 31, 1996
>DLT>			3962432	JUL 16, 1996	U-53	NCE OCT 31, 1996

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APPL/PROD NUMBER	INGREDIENT NAME; TRADE NAME	PATENT NUMBER	PATENT EXPIRES	USE CODE	EXCLUS CODE	EXCLUS EXPIRES
20036 003	PAMIDRONATE DISODIUM; AREDIA	4711880	JUL 29, 2005		I-135	SEP 01, 1998
>ADD>		3962432	JAN 09, 1998	U-53	NCE	OCT 31, 1996
>DLT>		3962432	JUL 16, 1996	U-53	NCE	OCT 31, 1996
20036 004	PAMIDRONATE DISODIUM; AREDIA	4711880	JUL 29, 2005		I-135	SEP 01, 1998
>ADD>		3962432	JAN 09, 1998	U-53	NCE	OCT 31, 1996
>DLT>		3962432	JUL 16, 1996	U-53	NCE	OCT 31, 1996
20031 001	PAROXETINE HYDROCHLORIDE; PAXIL	4721723	SEP 24, 2008	U-12		
20031 002	PAROXETINE HYDROCHLORIDE; PAXIL	4721723	SEP 24, 2008	U-12		
20031 003	PAROXETINE HYDROCHLORIDE; PAXIL	4721723	SEP 24, 2008	U-12		
20031 004	PAROXETINE HYDROCHLORIDE; PAXIL	4721723	SEP 24, 2008	U-12		
20031 005	PAROXETINE HYDROCHLORIDE; PAXIL	4721723	SEP 24, 2008	U-12		
19385 001	PERGOLIDE MESYLATE; PERMAX	4797405	OCT 26, 2007			
		4166182	FEB 08, 2000			
19385 002	PERGOLIDE MESYLATE; PERMAX	4797405	OCT 26, 2007			
		4166182	FEB 08, 2000			
19385 003	PERGOLIDE MESYLATE; PERMAX	4797405	OCT 26, 2007			
		4166182	FEB 08, 2000			
>ADD>	20451 001	PORFIMER SODIUM; PHOTOFRIN			NCE	DEC 27, 2000
	17850 001	POTASSIUM CHLORIDE; KLOTRIX	4140756	JUN 10, 1996		
	18238 001	POTASSIUM CHLORIDE; MICRO-K	4259315	JUN 13, 2000		
	18238 002	POTASSIUM CHLORIDE; MICRO-K 10	4259315	JUN 13, 2000		
	19561 003	POTASSIUM CHLORIDE; MICRO-K LS	4259315	JUN 13, 2000		
	19898 002	PRAVASTATIN SODIUM; PRAVACHOL	4346227	OCT 20, 2005		
	19898 003	PRAVASTATIN SODIUM; PRAVACHOL	4346227	OCT 20, 2005		
	19898 004	PRAVASTATIN SODIUM; PRAVACHOL	4346227	OCT 20, 2005		
	19775 001	PRazosin HYDROCHLORIDE; MINIPRESS XL	4327725	NOV 25, 2000		
	19775 002	PRazosin HYDROCHLORIDE; MINIPRESS XL	4327725	NOV 25, 2000		
	19627 001	PROPOFOL; DIPRIVAN	4798846	MAR 19, 1997		
		4056635	MAR 19, 1997			
	18553 001	PROPRANOLOL HYDROCHLORIDE; INDERAL LA	4138475	SEP 14, 1997		
	18553 002	PROPRANOLOL HYDROCHLORIDE; INDERAL LA	4138475	SEP 14, 1997		
	18553 003	PROPRANOLOL HYDROCHLORIDE; INDERAL LA	4138475	SEP 14, 1997		
	18553 004	PROPRANOLOL HYDROCHLORIDE; INDERAL LA	4138475	SEP 14, 1997		
	19536 001	PROPRANOLOL HYDROCHLORIDE; INDERAL	4600708	JUL 19, 2005		
	19664 001	PSEUDOEPHEDRINE HYDROCHLORIDE; SELDANE-D	4929605	OCT 07, 2007	U-81	
		4254129	APR 10, 1999	U-81		
20021 002	PSEUDOEPHEDRINE HYDROCHLORIDE; EFIDAC/24	4801461	MAR 14, 2006			

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APPL/PROD NUMBER	INGREDIENT NAME; TRADE NAME	PATENT NUMBER	PATENT EXPIRES	USE CODE	EXCLUS CODE	EXCLUS EXPIRES
19885 001	QUINAPRIL HYDROCHLORIDE; ACCUPRIL	4743450	FEB 24, 2007			
		4344949	AUG 17, 2001	U-3		
19885 002	QUINAPRIL HYDROCHLORIDE; ACCUPRIL	4743450	FEB 24, 2007			
		4344949	AUG 17, 2001	U-3		
19885 003	QUINAPRIL HYDROCHLORIDE; ACCUPRIL	4743450	FEB 24, 2007			
		4344949	AUG 17, 2001	U-3		
19885 004	QUINAPRIL HYDROCHLORIDE; ACCUPRIL	4743450	FEB 24, 2007			
		4344949	AUG 17, 2001	U-3		
19901 001	RAMIPRIL; ALTACE	5061722	OCT 29, 2008	U-3	I-134	AUG 22, 1998
19901 002	RAMIPRIL; ALTACE	5061722	OCT 29, 2008	U-3	I-134	AUG 22, 1998
19901 003	RAMIPRIL; ALTACE	5061722	OCT 29, 2008	U-3	I-134	AUG 22, 1998
19901 004	RAMIPRIL; ALTACE	5061722	OCT 29, 2008	U-3	I-134	AUG 22, 1998
18703 001	RANITIDINE HYDROCHLORIDE; ZANTAC 150	4880636	MAY 13, 2008			
		4128658	JUL 25, 1997		I-120	MAR 29, 1998
18703 002	RANITIDINE HYDROCHLORIDE; ZANTAC 300	4880636	MAY 13, 2008			
		4128658	JUL 25, 1997		I-120	MAR 29, 1998
19090 001	RANITIDINE HYDROCHLORIDE; ZANTAC	4585790	MAY 11, 2004			
		4128658	JUL 25, 1997			
19593 001	RANITIDINE HYDROCHLORIDE; ZANTAC	4128658	JUL 25, 1997			
19593 002	RANITIDINE HYDROCHLORIDE; ZANTAC	4585790	APR 29, 2003			
		4521431	JUN 04, 2002			
		4128658	JUL 25, 1997			
>ADD> 19675 001	RANITIDINE HYDROCHLORIDE; ZANTAC	5068249	NOV 26, 2008			
		4585790	MAY 11, 2004		I-120	MAR 29, 1998
		4128658	JUL 25, 1997			
20095 001	RANITIDINE HYDROCHLORIDE; ZANTAC 150	5028432	FEB 22, 2010		I-120	MAR 29, 1998
		4128658	JUL 25, 1997			
20095 002	RANITIDINE HYDROCHLORIDE; ZANTAC 300	5028432	FEB 22, 2010		I-120	MAR 29, 1998
		4128658	JUL 25, 1997			
20251 001	RANITIDINE HYDROCHLORIDE; ZANTAC 150	5102665	JUN 23, 2009		I-120	MAR 29, 1998
		4128658	JUL 25, 1997			
20251 002	RANITIDINE HYDROCHLORIDE; ZANTAC 150	5102665	JUN 23, 2009		I-120	MAR 29, 1998
		4128658	JUL 25, 1997			
>ADD> 20520 001	RANITIDINE HYDROCHLORIDE; ZANTAC 75				NS	DEC 19, 1998
>ADD> 20599 001	RILUZOLE; RILUTEK				ODE	DEC 12, 2002
>ADD>					NCE	DEC 12, 2000

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APPL/PROD NUMBER	INGREDIENT NAME; TRADE NAME	PATENT NUMBER	PATENT EXPIRES	USE CODE	EXCLUS CODE	EXCLUS EXPIRES
>ADD> 20474 001	RIMEXOLONE; VEXOL	4686214	OCT 30, 2005	U-100	NCE	DEC 30, 1999
>DLT> 20474 001	RIMEXOLONE; VEXOL	4686214	AUG 11, 2004	U-100	NCE	DEC 30, 1999
20214 001	ROCURONIUM BROMIDE; ZEMURON (P/F)	4894369	APR 13, 2008			
20214 002	ROCURONIUM BROMIDE; ZEMURON	4894369	APR 13, 2008			
20236 001	SALMETEROL XINAFOATE; SEREVENT	5380922	MAY 14, 2013			
>ADD> 20628 001	SAQUINAVIR MESYLATE; INVIRASE				NCE	DEC 06, 2000
19839 001	SERTRALINE HYDROCHLORIDE; ZOLOFT	5248699	AUG 13, 2012	U-12		
		4962128	NOV 02, 2009	U-12		
19839 002	SERTRALINE HYDROCHLORIDE; ZOLOFT	5248699	AUG 13, 2012	U-12		
		4962128	NOV 02, 2009	U-12		
19839 003	SERTRALINE HYDROCHLORIDE; ZOLOFT	5248699	AUG 13, 2012	U-12		
		4962128	NOV 02, 2009	U-12		
19839 004	SERTRALINE HYDROCHLORIDE; ZOLOFT	5248699	AUG 13, 2012	U-12		
		4962128	NOV 02, 2009	U-12		
20478 001	SEVOFLURANE; ULTANE				NCE	JUN 07, 2000
19766 001	SIMVASTATIN; ZOCOR				I-128	JUN 30, 1998
19766 002	SIMVASTATIN; ZOCOR				I-128	JUN 30, 1998
19766 003	SIMVASTATIN; ZOCOR				I-128	JUN 30, 1998
19766 004	SIMVASTATIN; ZOCOR				I-128	JUN 30, 1998
19721 001	SOMATROPIN, BIOSYNTHETIC; NORDITROPIN				NS	MAY 08, 1998
19721 002	SOMATROPIN, BIOSYNTHETIC; NORDITROPIN				NS	MAY 08, 1998
>ADD> 20522 001	SOMATROPIN, BIOSYNTHETIC; NUTROPIN AQ				NP	DEC 29, 1998
20240 001	SPIRAPRIL HYDROCHLORIDE; RENORMAX	4470972	SEP 11, 2001	U-3	NCE	DEC 29, 1999
20240 002	SPIRAPRIL HYDROCHLORIDE; RENORMAX	4470972	SEP 11, 2001	U-3	NCE	DEC 29, 1999
20240 003	SPIRAPRIL HYDROCHLORIDE; RENORMAX	4470972	SEP 11, 2001	U-3	NCE	DEC 29, 1999
20240 004	SPIRAPRIL HYDROCHLORIDE; RENORMAX	4470972	SEP 11, 2001	U-3	NCE	DEC 29, 1999
20080 001	SUMATRIPTAN SUCCINATE; IMITREX	4816470	DEC 28, 2006	U-72		
20132 001	SUMATRIPTAN SUCCINATE; IMITREX	5037845	AUG 06, 2008	U-72		
		4816470	DEC 28, 2006	U-72		
20132 002	SUMATRIPTAN SUCCINATE; IMITREX	5037845	AUG 06, 2008	U-72		
		4816470	DEC 28, 2006	U-72		
20132 003	SUMATRIPTAN SUCCINATE; IMITREX	5037845	AUG 06, 2008	U-72		
		4816470	DEC 28, 2006	U-72		
>DLT> 19387 001	SUPROFEN; PROFENAL	4035376	JUL 12, 1994			
>DLT> 19387 001	SUPROFEN; PROFENAL	4035376	JUL 12, 1996			
20070 001	TACRINE HYDROCHLORIDE; COGNEX	4816456	OCT 01, 2006	U-82		
		4631286	OCT 25, 2004	U-97		

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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
20070 002	TACRINE HYDROCHLORIDE; COGNEX	4816456	OCT 01, 2006	U-82		
		4631286	OCT 25, 2004	U-97		
20070 003	TACRINE HYDROCHLORIDE; COGNEX	4816456	OCT 01, 2006	U-82		
		4631286	OCT 25, 2004	U-97		
20070 004	TACRINE HYDROCHLORIDE; COGNEX	4816456	OCT 01, 2006	U-82		
		4631286	OCT 25, 2004	U-97		
19829 001	TECHNETIUM TC-99M EXAMETAZIME KIT; CERETEC	4789736	DEC 06, 2005		I-124	APR 07, 1998
		4615876	OCT 07, 2003			
19981 001	TECHNETIUM TC-99M RED BLOOD CELL KIT; ULTRATAG	4755375	JUL 05, 2005	U-51		
19057 001	TERAZOSIN HYDROCHLORIDE; HYTRIN	5412095	APR 29, 2013			
		5294615	APR 29, 2013			
		5212176	JUN 29, 2010			
		4112097	JAN 21, 1997	U-3		
19057 002	TERAZOSIN HYDROCHLORIDE; HYTRIN	5412095	APR 29, 2013			
		5294615	APR 29, 2013			
		5212176	JUN 29, 2010			
		4112097	JAN 21, 1997	U-3		
19057 003	TERAZOSIN HYDROCHLORIDE; HYTRIN	5412095	APR 29, 2013			
		5294615	APR 29, 2013			
		5212176	JUN 29, 2010			
		4112097	JAN 21, 1997	U-3		
19057 004	TERAZOSIN HYDROCHLORIDE; HYTRIN	5412095	APR 29, 2013			
		5294615	APR 29, 2013			
		5212176	JUN 29, 2010			
		4112097	JAN 21, 1997	U-3		
20347 001	TERAZOSIN HYDROCHLORIDE; HYTRIN	5412095	APR 29, 2013			
		5294615	APR 29, 2013			
		5212176	JUN 29, 2010			
		4112097	JAN 21, 1997	U-3		
20347 002	TERAZOSIN HYDROCHLORIDE; HYTRIN	5412095	APR 29, 2013			
		5294615	APR 29, 2013			
		5212176	JUN 29, 2010			
		4112097	JAN 21, 1997	U-3		
20347 003	TERAZOSIN HYDROCHLORIDE; HYTRIN	5412095	APR 29, 2013			
		5294615	APR 29, 2013			
		5212176	JUN 29, 2010			
		4112097	JAN 21, 1997	U-3		

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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
20347 004	TERAZOSIN HYDROCHLORIDE; HYTRIN	5412095	APR 29, 2013			
		5294615	APR 29, 2013			
		5212176	JUN 29, 2010			
		4112097	JAN 21, 1997			
		4254129	APR 10, 1999	U-81		
18949 001	TERFENADINE; SELDANE				NS	SEP 29, 1998
20489 001	TESTOSTERONE; ANDRODERM					
19979 001	TICLOPIDINE HYDROCHLORIDE; TICLID	4591592	MAY 27, 2003			
19979 002	TICLOPIDINE HYDROCHLORIDE; TICLID	4591592	MAY 27, 2003			
20439 001	TIMOLOL; BETIMOL	5231095	JUL 27, 2010		NP	MAR 31, 1998
20439 002	TIMOLOL; BETIMOL	5231095	JUL 27, 2010		NP	MAR 31, 1998
20136 001	TORSEMIDE; DEMADEX	RE34672	AUG 11, 2006			
20136 002	TORSEMIDE; DEMADEX	RE34672	AUG 11, 2006			
20136 003	TORSEMIDE; DEMADEX	RE34672	AUG 11, 2006			
20136 004	TORSEMIDE; DEMADEX	RE34672	AUG 11, 2006			
20137 002	TORSEMIDE; DEMADEX	4861786	JUL 08, 2007			
		RE34672	AUG 11, 2006			
20281 001	TRAMADOL HYDROCHLORIDE; ULTRAM				NCE	MAR 03, 2000
20281 002	TRAMADOL HYDROCHLORIDE; ULTRAM				NCE	MAR 03, 2000
18207 003	TRAZODONE HYDROCHLORIDE; DESYREL	4258027	MAR 26, 1999			
		4215104	MAR 26, 1999			
18207 004	TRAZODONE HYDROCHLORIDE; DESYREL	4258027	MAR 26, 1999			
		4215104	MAR 26, 1999			
>ADD> 20438 001	TRETINOIN; VESANOID				ODE	NOV 22, 2002
19798 001	TRIAMCINOLONE ACETONIDE; NASACORT	4767612	JAN 23, 2007	U-85		
20326 001	TRIMETREXATE GLUCURONATE; NEUTREXIN	4694007	MAY 20, 2006	U-91		
		4376858	OCT 31, 2000			
20487 001	VALACYCLOVIR HYDROCHLORIDE; VALTREX	4957924	SEP 18, 2007		NE	JUN 23, 1998
20487 002	VALACYCLOVIR HYDROCHLORIDE; VALTREX	4957924	SEP 18, 2007		NE	JUN 23, 1998
18776 002	VECURONIUM BROMIDE; NORCURON	4297351	AUG 20, 1999			
		4237126	AUG 20, 1999			
18776 003	VECURONIUM BROMIDE; NORCURON	4297351	AUG 20, 1999			
		4237126	AUG 20, 1999			
20151 001	VENLAFAXINE HYDROCHLORIDE; EFFEXOR	4535186	DEC 13, 2002			
20151 002	VENLAFAXINE HYDROCHLORIDE; EFFEXOR	4535186	DEC 13, 2002			
20151 003	VENLAFAXINE HYDROCHLORIDE; EFFEXOR	4535186	DEC 13, 2002			
20151 004	VENLAFAXINE HYDROCHLORIDE; EFFEXOR	4535186	DEC 13, 2002			
20151 005	VENLAFAXINE HYDROCHLORIDE; EFFEXOR	4535186	DEC 13, 2002			

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APPL/PROD NUMBER	INGREDIENT NAME; TRADE NAME	PATENT NUMBER	PATENT EXPIRES	USE CODE	EXCLUS CODE	EXCLUS EXPIRES
20151 006	VENLAFAXINE HYDROCHLORIDE; EFFEXOR	4535186	DEC 13, 2002			
19614 001	VERAPAMIL HYDROCHLORIDE; VERELAN	4863742	JUN 19, 2007	U-3		
19614 002	VERAPAMIL HYDROCHLORIDE; VERELAN	4863742	JUN 19, 2007	U-3		
19614 003	VERAPAMIL HYDROCHLORIDE; VERELAN	4863742	JUN 19, 2007	U-3		
20388 001	VINORELBINE TARTRATE; NAVELBINE	4307100	AUG 20, 1999		NCE	DEC 23, 1999
19655 001	ZIDOVUDINE; RETROVIR	4837208	SEP 17, 2005			
		4833130	SEP 17, 2005			
		4828838	SEP 17, 2005			
		4818538	SEP 17, 2005			
		4724232	SEP 17, 2005			
19910 001	ZIDOVUDINE; RETROVIR	4837208	SEP 17, 2005			
		4833130	SEP 17, 2005			
		4818538	SEP 17, 2005			
		4724232	SEP 17, 2005			
19951 001	ZIDOVUDINE; RETROVIR	4837208	SEP 17, 2005			
		4833130	SEP 17, 2005			
		4818538	SEP 17, 2005			
		4724232	SEP 17, 2005			
>ADD>	20518 001	ZIDOVUDINE; RETROVIR	4837208	SEP 17, 2005		
>ADD>			4833130	SEP 17, 2005		
>ADD>			4828838	SEP 17, 2005		
>ADD>			4818538	SEP 17, 2005		
>ADD>			4724232	SEP 17, 2005		
	19908 001	ZOLPIDEM TARTRATE; AMBIEN	4382938	OCT 21, 2006	U-74	
	19908 002	ZOLPIDEM TARTRATE; AMBIEN	4382938	OCT 21, 2006	U-74	

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