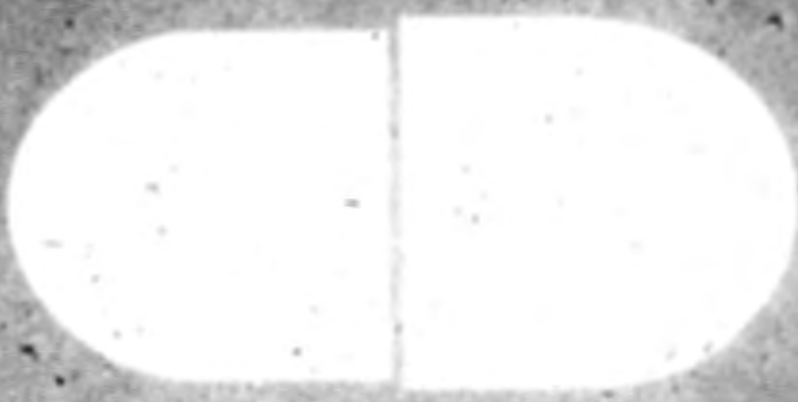


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0499-K-03

CUMULATIVE
SUPPLEMENT 10
JAN'98-OCT'98



APPROVED DRUG PRODUCTS

WITH
THERAPEUTIC EQUIVALENCE EVALUATIONS

18TH EDITION

U.S. DEPARTMENT OF HEALTH AND HUMAN SERVICES

PUBLIC HEALTH SERVICE
FOOD AND DRUG ADMINISTRATION
CENTER FOR DRUG EVALUATION AND RESEARCH
OFFICE OF RESEARCH AND TRAINING
DIVISION OF DATA MANAGEMENT AND SERVICES

99-013177

2:

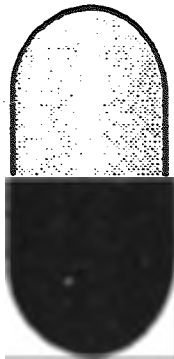
Prepared By
Division of Data Management and Services
Office of Information Technology
Center for Drug Evaluation and Research, FDA

C-1

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APPROVED DRUG PRODUCTS

**WITH
THERAPEUTIC EQUIVALENCE EVALUATIONS**

**19TH EDITION
1999**

CONTENTS

- Prescription Drug Product List
- OTC Drug Product List
- Drug Products with Approval under Section 505 of the Act Administered by the Center for Biologics Evaluation and Research List
- Discontinued Drug Product List
- Orphan Drug Product Designations
- Drug Products Which Must Demonstrate in vivo Bioavailability Only if Product Fails to Achieve Adequate Dissolution
- Patent and Exclusivity Information

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**APPROVED DRUG PRODUCTS
with
THERAPEUTIC EQUIVALENCE EVALUATIONS**

18TH EDITION

Cumulative Supplement 10

OCTOBER 1998

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**APPROVED DRUG PRODUCTS
with
THERAPEUTIC EQUIVALENCE EVALUATIONS**

18TH EDITION

**CUMULATIVE SUPPLEMENT 10
OCTOBER 1998**

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, 18th 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. Patent and Exclusivity information for a specific drug is indicated by an abbreviation followed by the date upon which the patent and/or exclusivity expires. Refer to the Patent and Exclusivity Terms section in the Patent and Exclusivity Information Addendum for an explanation of all codes and abbreviations. Information regarding drug patents and exclusivity for an approved product will appear in this addendum as it is received and may not correspond with the month the drug product is approved and published in the Rx/OTC sections of the Cumulative Supplement.

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

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

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

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

New deletions to the Prescription Drug Product List, OTC Drug Product List, and the Patent and Exclusivity Data are indicated by the symbol >DLT> (DELETE) to the left of the line. 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 (hard copy only) for this edition. However, the overstruck data in the Patent and Exclusivity Data is dropped in subsequent Cumulative Supplements (hard copy).

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 18th Edition List will then be added to the "Discontinued Drug Product List" appearing in the 19th Edition.

1.2 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 Whitworth Towne PLSN [New Abbreviated Name], the name change will appear in this section and will be identified with an asterisk.

APPLICANT NAME CHANGES

<u>FORMER APPLICANT NAME</u> <u>(FORMER ABBREVIATED NAME)</u>	<u>NEW APPLICANT NAME</u> <u>(NEW ABBREVIATED NAME)</u>
ASTRA MERCK INC (ASTRA MERCK)	ASTRA PHARMACEUTICALS LP (ASTRA PHARMS)
ASTRA USA INC (ASTRA)	ASTRA PHARMACEUTICALS LP (ASTRA PHARMS)
DUPONT RADIOPHARMACEUTICALS DIV (DUPONT)	DUPONT PHARMACEUTICALS COMPANY (DUPONT PHARMS)
FUJISAWA USA INC (FUJISAWA)	AMERICAN PHARMACEUTICAL PARTNERS INC (AM PHARM PARTNERS)
GENSIA INC (GENSIA)	GENSIA SICOR PHARMACEUTICALS INC (GENSIA SICOR PHARMS)
GENSIA LABORATORIES LTD (GENSIA)	GENSIA SICOR PHARMACEUTICALS INC (GENSIA SICOR PHARMS)
JONES MEDICAL INDUSTRIES INC (JONES MEDCL INDS)	JONES PHARMA INC (JONES PHARMA)
MERCK RESEARCH LABORATORIES DIV MERCK AND CO INC (MERCK)	MERCK AND CO INC (MERCK)
MERCK SHARP AND DOHME DIV MERCK AND CO INC (MERCK)	MERCK AND CO INC (MERCK)

1.3 ACYCLOVIR 200MG TABLET-REFERENCE LISTED DRUG

Novopharm's single source acyclovir tablets have been declared to be a reference listed drug for the 200 mg tablet in addition to the acyclovir (Zovirax) 800 mg tablet of the innovator. A generic firm wishing to submit an ANDA for a duplicate of the 200 mg acyclovir tablet will be eligible for a waiver of the *in vivo* determination of bioequivalence (1) if their product is proportionally similar in its active and inactive ingredients to their own 800 mg acyclovir tablet and (2) by doing an acceptable comparative dissolution test (dissolution profile) against Novopharm's 200 mg acyclovir reference listed drug.

Before a waiver of the *in vivo* determination of bioequivalence can be granted for the 200 mg acyclovir tablet, the generic firm must have completed an acceptable fasting and fed study comparing their acyclovir 800 mg tablet against the Zovirax 800 mg tablet.

For further information on the study designs, you should contact the Division of Bioequivalence, Office of Generic Drugs.

1.4 DICLOFENAC SODIUM OPHTHALMIC SOLUTION 0.1%

Two NDAs have been approved for diclofenac sodium ophthalmic solution 0.1% (DSOS), (1) Ciba's NDA 20-037 for Voltaren and (2) Alcon's NDA 20-809 for DSOS. Alcon was required to do a study comparing their DSOS to Voltaren and to a placebo control in post cataract surgical inflammation. This study was necessary to demonstrate that the different formulation of the Alcon drug product did not affect the safety and/or effectiveness of the proposed drug product for this indication. Prior to the approval of Alcon's DSOS Ciba did clinical studies and was approved for two additional indications for the temporary relief of pain and photophobia in patients undergoing corneal refractive surgery. Three years of Waxman-Hatch marketing exclusivity was granted to Ciba for these two new uses.

Since the treatment of pain has a different site of action than the anti-inflammatory or photophobia indications the Agency did not have information to support a recommendation that the Alcon and Ciba DSOS are therapeutically equivalent for the treatment of pain. The designation of therapeutic equivalence at this time applies only to the anti-inflammatory indication. The therapeutic equivalence designation will apply to the photophobia indication upon expiration of Ciba's marketing exclusivity.

1.5 RIBAVIRIN

Indicated for use and comarketed with interferon alfa-2b, recombinant (Intron A), as Rebetrone Combination Therapy.

1.6 FOLLITROPIN ALFA AND BETA

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

1.7 AVAILABILITY OF THE PUBLICATION AND UPDATING PROCEDURES

The *Approved Drug Products with Therapeutic Equivalence Evaluations* files are available on Internet. There is more than one media users may select to access these files.

Preface and ASCII Text Files:

The Preface may be accessed using this URL: <http://www.fda.gov/cder/orange/adp.htm>. Users who wish to download the Prescription Drug Product List; OTC Drug Products and Discontinued Drug Products lists may access the ASCII text files using this URL: <http://www.fda.gov/cder/orange/obreadme.htm>. The drug product lists and the zipobtxt.exe files are updated concurrently with the publication of the annual edition or monthly cumulative supplements. Appendix A and B are updated twice a year.

Preface and Searchable Query Database:

The Preface may be accessed using this URL: <http://www.fda.gov/cder/ob/docs/preface/ectablec18.htm>. Users who wish to query on a specific drug product may access the database using this URL: <http://www.fda.gov/cder/ob>. The Query enables searching of the database by active ingredient, proprietary name, applicant holder or applicant number. The data is updated concurrently with the publication of the annual edition or monthly cumulative supplements.

1.8 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 section 505 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 1997) 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 1997</u>	<u>MAR 1998</u>	<u>JUN 1998</u>	<u>SEP 1998</u>
DRUG PRODUCTS LISTED	9624	9711	9768	9798
SINGLE SOURCE	2462 (25.6%)	2484 (25.6%)	2494 (25.6%)	2479 (25.3%)
MULTISOURCE	7052 (73.3%)	7117 (73.3%)	7164 (73.3%)	7208 (73.6%)
THERAPEUTICALLY EQUIVALENT	6673 (69.3%)	6746 (69.5%)	6790 (69.5%)	6829 (69.7%)
NOT THERAPEUTICALLY EQUIVALENT	379 (4.0%)	371 (3.8%)	374 (3.8%)	379 (3.9%)
EXCEPTIONS ¹	110 (1.1%)	110 (1.1%)	110 (1.1%)	111 (1.1%)
NEW MOLECULAR ENTITIES APPROVED	--	8	9	4
NUMBER OF APPLICANTS	551	529	538	551

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

ACETAMINOPHEN, HYDROCODONE BITARTRATE

TABLET, ORAL
HYDROCODONE BITARTRATE AND ACETAMINOPHEN

AA	WATSON LABS	50MG/2.5MG	MAR 04, 1996
AA		50MG/5MG	MAR 04, 1996
AA		50MG/7.5MG	MAR 04, 1996
AA		50MG/7.5MG	MAR 04, 1996
AA		50MG/10MG	MAR 04, 1996
AA		75MG/7.5MG	MAR 04, 1996

ACETAMINOPHEN, OXYCODONE HYDROCHLORIDE

TABLET, ORAL
OXYCODONE AND ACETAMINOPHEN

AA	DURAMED	325MG/5MG	MAR 04, 1996
AA	WATSON LABS	325MG/5MG	MAR 04, 1996

ACETAMINOPHEN, PROPXYPHENE HYDROCHLORIDE

TABLET, ORAL
PROPXYPHENE HCL AND ACETAMINOPHEN

AA	WATSON LABS	550MG/5MG	MAR 04, 1996
----	-------------	-----------	--------------

ACETIC ACID, GLACIAL

SOLUTION, IRRIGATION, URETHRAL
ACETIC ACID 0.25% IN PLASTIC CONTAINER
B. BRAUN

AT		250MG/100ML	MAR 04, 1996
----	--	-------------	--------------

ACETAMINOPHEN, OXYCODONE

CAPSULE, ORAL
OXYCODONE AND ACETAMINOPHEN

AA	HALSET	50MG/5MG	MAR 04, 1996
----	--------	----------	--------------

ACETIC ACID, GLACIAL, HYDROCORTISONE

SOLUTION/DROPS, OTIC
HYDROCORTISONE AND ACETIC ACID

		2g/1g	MAR 04, 1996
--	--	-------	--------------

ACETAMINOPHEN, OXYCODONE HYDROCHLORIDE

CAPSULE, ORAL
OXYCODONE AND ACETAMINOPHEN

AA	WALLINGCROFT	50MG/5MG	MAR 04, 1996
AA	WATSON LABS	50MG/5MG	MAR 04, 1996

ACYCLOVIR

CAPSULE, ORAL
ACYCLOVIR

AA	CHELSEA LABS	200MG	MAR 04, 1996
AA	GENPHARM	200MG	MAR 04, 1996

AA		200MG	MAR 04, 1996
----	--	-------	--------------

AA		200MG	MAR 04, 1996
----	--	-------	--------------

ALLOPURINOL SODIUM

INJECTABLE; INJECTION
STYOPHIN

ALPRAZOLAM

TABLET; ORAL
ALPRAZOLAM
GENEVA PHARMS

2MG

1MG

0.5MG

0.25MG

NATSON LABS

0.25MG

0.5MG

1MG

ALPROSTADIL

INJECTABLE; INJECTION
ALPROSTADIL
BEDFORD

0.5MG/ML

PROCTER & KENDRICK
+ PHARMACIA AND UNION

0.5MG/ML

AMANTADINE HYDROCHLORIDE

CAPSULE; ORAL
AMANTADINE HCL

100MG

50MG

AMANTADINE HYDROCHLORIDE

CAPSULE; ORAL
AMANTADINE HCL
ENDO PHARMS

100MG

SYRUP; ORAL

AMANTADINE HCL
+ ENDO PHARMS

50MG/5ML

TABLET; ORAL

SYNTHREL
+ NATSON LABS

100MG

AMCINONIDE

OINTMENT; TOPICAL

CYCLOCORT
+ WYETH AVERST

0.1%

ANILORIDE HYDROCHLORIDE

TABLET; ORAL

ANILORIDE HCL
+ MERCK

5MG

ANILORIDE HYDROCHLORIDE; HYDROCHLOROTHIAZIDE

TABLET; ORAL

ANILORIDE HCL AND HYDROCHLOROTHIAZIDE
+ NATSON LABS

50 50MG ANILORIDE; 50MG

MOORETIC S-50
+ MERCK

50 50MG ANILORIDE; 50MG

AMINO ACIDS

INJECTABLE; INJECTION
PROSOL 204 SULFITE FREE IN PLASTIC CONTAINER
+ BAXTER MLTICARE 204

N30849 001
AUG 26, 1998

AMINOCAPROIC ACID

SYRUP; ORAL

AMICAR

2A + IMMUNE

1.25GM/5ML

N15230 002

AMINOCAPROIC ACID

2A NICART

1.25GM/5ML

N74759 001
SEP 02, 1998

AMIODARONE HYDROCHLORIDE

TABLET; ORAL

CORDARONE

2A + WYETH AYERST

200MG

N10972 001
DEC 24, 1985

2A PACTONE

200MG

N75135 001
APR 30, 1998

ANITRIPTYLINE HYDROCHLORIDE

TABLET; ORAL

ANITRIPTYLINE HCL

2A

10MG

N72539 001

10MG; 2MG

N72540 001

10MG; 4MG

N72541 001

25MG; 2MG

N72134 001

25MG; 4MG

N72135 001

50MG; 4MG

10MG

N72539 001
FEB 15, 1989

N72540 001
FEB 15, 1989

N72541 001
FEB 15, 1989

N72134 001
FEB 15, 1989

N72135 001
FEB 15, 1989

ANITRIPTYLINE HYDROCHLORIDE

TABLET; ORAL

ANITRIPTYLINE HCL

2A DANBURY PHARMA

25MG

50MG

75MG

100MG

150MG

N88621 001
MAR 02, 1984

N88622 001
MAR 02, 1984

N88633 001
MAR 02, 1984

N88634 001
MAR 02, 1984

N88635 001
MAR 02, 1984

ANITRIPTYLINE HYDROCHLORIDE; PERPHENAZINE

TABLET; ORAL

PERPHENAZINE AND ANITRIPTYLINE HCL

2A

10MG; 2MG

10MG; 4MG

25MG; 2MG

25MG; 4MG

50MG; 4MG

10MG

N72539 001

FEB 15, 1989

N72540 001

FEB 15, 1989

N72541 001

FEB 15, 1989

N72134 001

FEB 15, 1989

N72135 001

FEB 15, 1989

ATENOLOL

TABLET; ORAL

ATENOLOL

CEPHALAM

25MG

M74136 003

AUG 26, 1998

ATENOLOL; CHLORTHALIDONE

TABLET; ORAL

ATENOLOL AND CHLORTHALIDONE

25MG; 25MG

M74404 001

MAY 14, 1998

100MG; 25MG

M74404 002

MAY 14, 1998

ATORVASTATIN CALCIUM

TABLET; ORAL

LIPITOR

ATORVASTATIN

20MG

M74404 001

MAY 14, 1998

+

WARNER LANBERT EXFOR EQ 10MG BASE

EQ 20MG BASE

M20702 002

DEC 17, 1996

EQ 40MG BASE

M20702 003

DEC 17, 1996

ATRACURTIUM BESYLATE

INJECTABLE; INJECTION

ATRACURTIUM BESYLATE

MADEAN

10MG/ML

M74945 001

JUL 26, 1998

ATRACURTIUM BESYLATE PRESERVATIVE FREE

10MG/ML

M74944 001

JUL 26, 1998

BACITRACIN

TABLET; ORAL

BACITRACIN

CEPHALAM

5,000,000 UNITS/BOT

M62456 001

JUL 27, 1993

BACITRACIN ZINC; HYDROCORTISONE; NEOMYCIN SULFATE; POLYMYXIN B SULFATE

OINTMENT; OPHTHALMIC

CORTISPORIN

AT + MONARCH PHARMS

M50416 002

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

10,000 UNITS/GM

BACITRACIN ZINC; NEOMYCIN SULFATE; POLYMYXIN B SULFATE

OINTMENT; OPHTHALMIC

NEOSPORIN

AT + MONARCH PHARMS

M50417 001

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

10,000 UNITS/GM

BACITRACIN ZINC; POLYMYXIN B SULFATE

OINTMENT; OPHTHALMIC

BACITRACIN ZINC AND POLYMYXIN B SULFATE

AT

M50417 001

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

10,000 UNITS/GM

AT

AKORN

500 UNITS/GM;

10,000 UNITS/GM

M54028 001

JAN 30, 1995

AT

AKORN

500 UNITS/GM;

10,000 UNITS/GM

M61229 001

JAN 30, 1995

NOTES

TALENT: ONLY

1990

WATSON LABS

22

MF73092 001
 JAN 28, 1994
 MF73093 001
 JAN 28, 1994

ANALYSTS TO GO

INJECTABLE, INJECTION

PLATE POLYCYCLOPOLYESTERS

INJECTABLE; INJECTION

104-104

+

69-1201-23

FOOTNOTES

BYRON

SUSPENSION/DROPS: OPHTHALMIC

1307

MOJO + ALCON

APR 01, 1998
N20816 001

REVOLUTION

TABLIST: ORAL

vasco

30003

40002

N19002 003
 DEC 28, 1990
 N19002 003
 DEC 28, 1990

MONOCLIPTRIN M331AT3

TABULET: GRAY

WATKINS INTERNATIONAL

NEW YORK

LEX PHARM 80 2 2003 PAGE 22

W74631 001
JAN 13, 1998

PROTIV

27 HOWARTIS

NOVARTIS
NO. 3, 5th Ave. NYC

17962 001

PROFESSIONAL VALUERS

RESEARCHER VALUE

CREAM: TOPICAL

STYLING: JESSICA WILSON

BO 0.19 2257

W70033 001
JUN 10. 1986

BUPROPION HYDROCHLORIDE

"TABLET- EXTENDED RELEASE: ORAL.

NEW!

EUTROPIUM HYDROCHLORIDE

TABLET, EXTENDED RELEASE, ORAL
EUTRAM
6 GLAND WELLCOME 100MG

870711 002
MAY 14, 1997

EUTROPIUM TARTARATE

INJECTABLE; INJECTION
EUTROPIUM TARTARATE
BEDFORD

2MG/ML

875046 001
AUG 12, 1998

EUTROPIUM TARTARATE PRESERVATIVE FREE

BEDFORD

1MG/ML

875045 001
AUG 12, 1998

2MG/ML

875045 002
AUG 12, 1998

1MG/ML

875170 001
SEP 28, 1998

2MG/ML

875170 002
SEP 28, 1998

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

INJECTABLE; INJECTION
ISOLYTE R IN DEXTROSE 5% IN PLASTIC CONTAINER
B BRAUN

37MG/100ML; 5GM/100ML; 31MG/100ML;
120MG/100ML; 330MG/100ML;
88MG/100ML

819864 001
JUN 10, 1993

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

INJECTABLE; INJECTION
ISOLYTE R IN DEXTROSE 5% IN PLASTIC CONTAINER
B BRAUN

35MG/100ML; 5GM/100ML; 30MG/100ML;
74MG/100ML; 640MG/100ML; 500MG/100ML;
74MG/100ML

819867 001
DEC 20, 1993

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

INJECTABLE; INJECTION
ISOLYTE R IN DEXTROSE 5% IN PLASTIC CONTAINER
B BRAUN

CALCIUM CHLORIDE; DEXTROSE; POTASSIUM CHLORIDE; SODIUM CHLORIDE

INJECTABLE; INJECTION

DEXTROSE 5% IN RINGER'S IN PLASTIC CONTAINER

B BRAUN

31MG/100ML; 5GM/100ML; 30MG/100ML;
88MG/100ML

818356 001
31MG/100ML; 5GM/100ML; 30MG/100ML;
88MG/100ML

826000 001
APR 17, 1992

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

INJECTABLE; INJECTION

DEXTROSE 5% IN LACTATED RINGER'S IN PLASTIC CONTAINER

B BRAUN

20MG/100ML; 5GM/100ML; 30MG/100ML;
500MG/100ML; 310MG/100ML

817510 001

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

INJECTABLE; INJECTION

ISOLYTE R IN PLASTIC CONTAINER

B BRAUN

35MG/100ML; 30MG/100ML; 74MG/100ML;
640MG/100ML; 500MG/100ML;
74MG/100ML

819718 001
SEP 29, 1989

**CALCIUM CHLORIDE, MAGNESIUM CHLORIDE, POTASSIUM CHLORIDE, SODIUM
LACTATE, SODIUM CHLORIDE, SODIUM CITRATE**

INJECTABLE; INJECTION
ISOLATE 5 IN PLASTIC CONTAINER

17.6MG/100ML; 325.3MG/100ML;
119.3MG/100ML; 643MG/100ML; 100ML; 100ML; 100ML;
FEB 26, 1982

**CALCIUM CHLORIDE, MAGNESIUM CHLORIDE, POTASSIUM CHLORIDE, SODIUM
CHLORIDE**

SOLUTION; PERUSION, CARDIAC
FLUIDS; IN PLASTIC CONTAINER

17.6MG/100ML; 325.3MG/100ML;
119.3MG/100ML; 643MG/100ML; 100ML; 100ML; 100ML;
FEB 26, 1982

CALCIUM CHLORIDE, POTASSIUM CHLORIDE, SODIUM CHLORIDE

INJECTABLE; INJECTION
ISOLATE 5 IN PLASTIC CONTAINER

17.6MG/100ML; 325.3MG/100ML;
119.3MG/100ML; 643MG/100ML; 100ML; 100ML; 100ML;
FEB 26, 1982

17.6MG/100ML; 325.3MG/100ML;
119.3MG/100ML; 643MG/100ML; 100ML; 100ML; 100ML;
FEB 26, 1982

SOLUTION; IRRIGATION

17.6MG/100ML; 325.3MG/100ML;
119.3MG/100ML; 643MG/100ML; 100ML; 100ML; 100ML;
FEB 26, 1982

**CALCIUM CHLORIDE, POTASSIUM CHLORIDE, SODIUM CHLORIDE, SODIUM
LACTATE**

INJECTABLE; INJECTION
ISOLATE 5 IN PLASTIC CONTAINER

17.6MG/100ML; 325.3MG/100ML;
119.3MG/100ML; 643MG/100ML; 100ML; 100ML; 100ML;
FEB 29, 1988

**CALCIUM CHLORIDE, POTASSIUM CHLORIDE, SODIUM CHLORIDE, SODIUM
CHLORIDE**

SOLUTION; IRRIGATION
ISOLATE 5 IN PLASTIC CONTAINER

17.6MG/100ML; 325.3MG/100ML;
119.3MG/100ML; 643MG/100ML; 100ML; 100ML; 100ML;
FEB 27, 1982

CALCIUM

INJECTABLE; INJECTION
ISOLATE 5 IN PLASTIC CONTAINER

17.6MG/100ML; 325.3MG/100ML;
119.3MG/100ML; 643MG/100ML; 100ML; 100ML; 100ML;
FEB 27, 1982

**CALCIUM CHLORIDE, POTASSIUM CHLORIDE, SODIUM CHLORIDE, SODIUM
LACTATE**

INJECTABLE; INJECTION
ISOLATE 5 IN PLASTIC CONTAINER

17.6MG/100ML; 325.3MG/100ML;
119.3MG/100ML; 643MG/100ML; 100ML; 100ML; 100ML;
FEB 27, 1982

JUN 04, 1998

CAPTORIL: HYDROCHLOROTHIAZIDE

TABLET; ORAL
CAPTORIL AND HYDROCHLOROTHIAZIDE
SMITH GOLDLINE 15MG; 25MG
N178033 002
JUN 18, 1998
N178035 004
JUN 18, 1998
N178035 003
JUN 18, 1998

CARBIDOPA: LEVODOPA

TABLET, EXTENDED RELEASE; ORAL
SINEMET CR
N19856 002
DEC 24, 1992
N19856 001
MAY 30, 1991

CARBAMAZEPINE

CAPSULE, EXTENDED RELEASE; ORAL
CARBATROL
N178033 002
JUN 18, 1998
N178035 004
JUN 18, 1998
N178035 003
JUN 18, 1998

CARISOPRODOL

TABLET; ORAL
CARISOPRODOL
N178033 002
JUN 18, 1998
N178035 004
JUN 18, 1998
N178035 003
JUN 18, 1998

CETACLOF

POWDER FOR RECONSTITUTION; ORAL

CETACLOF
N178033 002
JUN 18, 1998
N178035 004
JUN 18, 1998
N178035 003
JUN 18, 1998

CARBIDOPA

TABLET; ORAL
LODOVYN
N178033 002
JUN 18, 1998
N178035 004
JUN 18, 1998
N178035 003
JUN 18, 1998

CARBIDOPA: LEVODOPA

TABLET; ORAL
SINEMET CR
N19856 002
DEC 24, 1992
N19856 001
MAY 30, 1991

CETAFOLIN SODIUM

INJECTABLE; INJECTION
CETAFOLIN SODIUM
AM PHARM PARTNERS

N178033 002
JUN 18, 1998
N178035 004
JUN 18, 1998
N178035 003
JUN 18, 1998

EQ 1MG BASE/VIAL
EQ 500MG BASE/VIAL
EQ 100MG BASE/VIAL
EQ 200MG BASE/VIAL

N64169 002
AUG 14, 1998
N64169 001
AUG 14, 1998
N64170 001
MAR 18, 1998
N64170 002
MAR 18, 1998

N64204 001
FEB 18, 1998
N64205 001
FEB 18, 1998
N64206 001
FEB 18, 1998
N64207 001
FEB 18, 1998

N40152 001
DEC 03, 1996

CETROXIME SODIUM

INJECTABLE; INJECTION

CETROXIME

ASTRA PHARMS

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CERIVALEXIN

CAPSULE; ORAL

CERIVALEXIN

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CETROXIME SODIUM

INJECTABLE; INJECTION

CETROXIME

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CERIVALEXIN

CAPSULE; ORAL

CERIVALEXIN

SMITH GOLDLINE

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CETROXIME SODIUM

INJECTABLE; INJECTION

CETROXIME

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CERIVALEXIN

CAPSULE; ORAL

CERIVALEXIN

SMITH GOLDLINE

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CHLORAMPHENICOL; HYDROCORTISONE ACETATE

POWDER FOR RECONSTITUTION; OPHTHALMIC

CHLORAMPHENICOL HYDROCORTISONE

+ PARKEDALE 12.5MG/VIAL; 25MG/VIAL

NS0202 001

CHLORAMPHENICOL; HYDROCORTISONE ACETATE; POLYMYXIN B SULFATE

OPHTHALMIC

OPHTHOCORT

+ PARKEDALE 10MG/GM; 5MG/GM;

10,000 UNITS/GM

NS0201 002

CHLORAMPHENICOL SODIUM SUCCEINATE

INJECTABLE; INJECTION

CHLORAMPHENICOL

+ PARKEDALE

NO 1GM BULK/VIAL

NS0155 001

CHLORDIAEPORIDE

TABLET; ORAL

LIMITABS

+ ICH

5MG

10MG

25MG

NS5482 001

NS5481 001

NS5483 001

CHLORDIAEPORIDE HYDROCHLORIDE

CAPSULE; ORAL

LIMITABS

+ ICH

5MG

10MG

25MG

NS5461 001

NS5472 001

NS5473 001

CHLORHEXIDINE GLUCONATE

SOLUTION; DENTAL

PERIOCHIP

AS + ZILA 9.123

NS19028 001

AUG 13, 1986

TABLET; DENTAL

PERIOCHIP

+ PERIO PRODS

2.5MG

NS0774 001

MAY 15, 1998

CHLOROPROCAINE HYDROCHLORIDE

INJECTABLE; INJECTION

CHLOROPROCAINE HCL

AS BEDFORD

23

NS0273 001

SEP 09, 1998

NS0273 002

SEP 09, 1998

CHLOROCQUINE PHOSPHATE

TABLET; ORAL

CHLOROCQUINE PHOSPHATE

AS WEST WARD

NO 150MG PAST

NS3022 001

CHLOROTHALIDE

SUSPENSION; ORAL

DIURIL

+ MERCK

250MG/5ML

NS11870 001

TABLET; ORAL

DIURIL

+ MERCK

250MG

NS1145 004

NS1145 002

CHLOROTHIAZIDE, RESERPINE

TABLET, ORAL

DIUPRES-250

MERCK

250MG; 0.125MG

N11635 003
AUG 26, 1987

DIUPRES-500

BP + MERCK

500MG; 0.125MG

N11635 006
AUG 26, 1987

CHLOROTHIAZIDE SODIUM

INJECTABLES; INJECTION

DIURIL

+ MERCK

EQ 500MG BASE/VIAL
EQ 100MG BASE/VIAL

N11145 005
MAY 28, 1998

CHLORPHENIRAMINE MALEATE

TABLET, ORAL

CHLORPHENIRAMINE MALEATE

4MG

N80696 001
MAY 28, 1998

CHLORPROMAZINE HYDROCHLORIDE

INJECTABLES; INJECTION

CHLORPROMAZINE HCL

25MG/ML

N89563 001
APR 15, 1988

CHLORZOXAZONE

TABLET, ORAL

CHLORZOXAZONE

250MG

250MG

N86901 001
MAY 28, 1998

CHLORZOXAZONE

TABLET, ORAL

CHLORZOXAZONE

300MG

300MG

N71040 001
AUG 22, 1989

CHOLESTYRAMINE

POWDER, ORAL

CHOLESTYRAMINE

NOVOPHARM

EQ 4GM RESIN/PACKET

N74347 001
MAY 28, 1998

EQ 4GM RESIN/SCOOPFUL

N74347 002
MAY 28, 1998

CHOLESTYRAMINE LIGHT

COSLEY PHARM

EQ 4GM RESIN/PACKET

N74555 001
SEP 30, 1998

EQ 4GM RESIN/PACKET

N74348 001
MAY 28, 1998

EQ 4GM RESIN/SCOOPFUL

N74348 002
MAY 28, 1998

LOCHOLEST

EQN

EQ 4GM RESIN/PACKET

N74561 001
AUG 15, 1996

EQ 4GM RESIN/SCOOPFUL

N74561 002
AUG 15, 1996

EQ 4GM RESIN/PACKET

N74561 003
AUG 15, 1996

EQ 4GM RESIN/SCOOPFUL

N74561 004
AUG 15, 1996

LOCHOLEST LIGHT

EQN

EQ 4GM RESIN/PACKET

N74562 001
AUG 15, 1996

EQ 4GM RESIN/SCOOPFUL

N74562 002
AUG 15, 1996

EQ 4GM RESIN/PACKET

N74562 003
AUG 15, 1996

EQ 4GM RESIN/SCOOPFUL

N74562 004
AUG 15, 1996

CIMETIDINE

SOLUTION; ORAL

CIMETIDINE HCL

DURAMED

300MG/5ML

N75110 001
JUN 18, 1998

CIMETIDINE

TABLET; ORAL

CIMETIDINE

200MG

200MG

200MG

200MG

ZENITH LABS

200MG

200MG

400MG

800MG

JUL 28, 1995

N74424 002

JUL 28, 1995

N74424 003

JUL 28, 1995

N74424 004

JUL 28, 1995

CIMETIDINE HYDROCHLORIDE

SOLUTION; ORAL

CIMETIDINE HCL

+ COPLEY PHARM

EQ 300MG BASE/5ML

N74859 001

JUL 09, 1998

CIPROFLOXACIN HYDROCHLORIDE

OINTMENT; OPHTHALMIC

CILLOXAN

+ ALCON

EQ 0.3% BASE

N20369 001

MAR 30, 1998

CIPROFLOXACIN HYDROCHLORIDE, HYDROCORTISONE

SUSPENSION/DROPS; OTIC

CIPRO HC

+ ALCON

EQ 0.2% BASE; 1%

N20805 001

FEB 10, 1998

CISAPRIDE NONHYDRATE

TABLET; ORAL

PROPULSID QUICKSOLV

+ JANSSEN

EQ 20MG BASE

N20767 001

NOV 07, 1997

TABLET, ORALLY DISINTEGRATING; ORAL

PROPULSID QUICKSOLV

+ JANSSEN

EQ 20MG BASE

N20767 001

CITALOPRAM HYDROBROMIDE

TABLET; ORAL

CELEXA

FOREST LABS

EQ 20MG BASE

N20822 002

JUL 17, 1998

N20822 003

JUL 17, 1998

N20822 004

JUL 17, 1998

CLEMASTINE FUMARATE

SYRUP; ORAL

CLEMASTINE FUMARATE

+ MORTON GROVE

EQ 0.5MG BASE/5ML

N74863 001

MAR 13, 1998

CLIDINIUM BROMIDE

TABLET; ORAL

CLIDINIUM BROMIDE

+ JANSSEN

EQ 2.5MG

N10355 001

N10355 002

CLINDAMYCIN HYDROCHLORIDE

CAPSULE; ORAL

CLINDAMYCIN HCL

+ JANSSEN

EQ 2.5MG

N10355 001

N10355 002

CLONIDINE HYDROCHLORIDE

CAPSULE, ORAL
CLONIDINE HCL
O DANDERY PHARMA

EQ 75MG BASE

N63082 001
JUL 31, 1991

CLONIDINE PROPIONATE

CREAM, VAGINAL
CLONIDINE

EQ 2t BASE

N50680 001
AUG 11, 1992
N50680 002
MAR 02, 1998

> DLT >
> DLT >
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INJECTABLE, INJECTION
CLONIDINE PROPIONATE

EQ 150MG BASE/ML

N62913 001
OCT 20, 1988

SOLUTION, TOPICAL
CLONIDINE T

EQ 1t BASE

N62363 001
FEB 08, 1982

CLONITASOL PROPIONATE

ointment, TOPICAL
CLONITASOL PROPIONATE
STIEFEL

0.05t

N75057 001
AUG 12, 1998

CLONIPRAMINE HYDROCHLORIDE

CAPSULE, ORAL
CLONIPRAMINE HCL
CHELSEA LABS

25MG

N74751 001
SEP 30, 1998

CLONIPRAMINE HYDROCHLORIDE

CAPSULE, ORAL
CLONIPRAMINE HCL
CHELSEA LABS

50MG

N74751 002
SEP 30, 1998
N74751 003
SEP 30, 1998
N74947 001
APR 30, 1998
N74947 002
APR 30, 1998
N74947 003
APR 30, 1998

MYLAN

25MG

50MG

75MG

CLONAZEPAM

TABLET, ORAL
CLONAZEPAM
MYLAN

0.5MG

N75150 001
OCT 05, 1998
N75150 002
OCT 05, 1998
N75150 003
OCT 05, 1998
N74920 001
AUG 04, 1998
N74920 002
AUG 04, 1998
N74920 003
AUG 04, 1998

NOVOPHARM

0.5MG

1MG

2MG

2MG

TABLET, ORALLY DISINTEGRATING, ORAL
KLOXOPIN RAPIDLY DISINTEGRATING
+ ROCHE
0.125MG

N20813 001
DEC 23, 1997

CLONAZEPAM

TABLET, ORALLY DISINTEGRATING; ORAL
KLOXONIN RAPIDLY DISINTEGRATING
ROCHE

0.25MG
0.5MG
1MG
2MG

N20813 002
DEC 23, 1997
N20813 003
DEC 23, 1997
N20813 004
DEC 23, 1997
N20813 005
DEC 23, 1997

+

CLONIDINE HYDROCHLORIDE

INJECTABLE; INJECTION
DURACION

0.1MG/ML

+

CODEINE PHOSPHATE; PSEUDOPHEDRINE HYDROCHLORIDE; TRIPROLIDINE
HYDROCHLORIDE

SYRUP; ORAL

10MG/5ML; 30MG/5ML;
1.25MG/5ML

N12575 003
APR 04, 1984

TRIPROLIDINE HCL; PSEUDOPHEDRINE HCL AND CODEINE PHOSPHATE

10MG/5ML; 30MG/5ML;
1.25MG/5ML

N88833 001
NOV 16, 1984

AA +

COLISTIMETHATE SODIUM

INJECTABLE; INJECTION
COLY-MYCIN M

EQ 150MG BASE/VIAL

N50108 002

COLISTIMETHATE SODIUM

INJECTABLE; INJECTION
COLY-MYCIN M
+ PARKEDALE

COLISTIN SULFATE; HYDROCORTISONE ACETATE; NEOMYCIN SULFATE;
THIOTHIONIN BROMIDE

SUSPENSION/DROPS; OTIC
COLY-MYCIN S

EQ 3MG BASE/ML; 10MG/ML;
EQ 3.3MG BASE/ML; 0.5MG/ML N50356 001

+

CORTICOTROPIN

INJECTABLE; INJECTION
ACTH

25 UNITS/VIAL
25 UNITS/VIAL
40 UNITS/VIAL

N08317 002
N08317 004

CRONOLYN SODIUM

SOLUTION/DROPS; OPHTHALMIC
CHOLON
BAUSCH AND LOMB

20MG

N16990 001

AT

> DLT >
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N74443 001
JAN 30, 1995

CRONOLYN SODIUM

AT
AKORN
OFFICIN
+ ALLEGAN

N74706 001
APR 29, 1998
N18193 001
OCT 03, 1984

CROMOLIN SODIUM

SOLUTION/DROPS; OPHTHALMIC
CORTICOSTEROID

CYCLOHEXAPRINE HYDROCHLORIDE

TABLET; ORAL

CYCLOHEXAPRINE HCL

AB WATSON LABS

10MG

CYCLOSPORINE

CAPSULE; ORAL
NEURAL

BX NOVARTIS

25MG

BX

50MG

BX +

100MG

SANDIMUNE
BX NOVARTIS

25MG

BX

50MG

BX +

100MG

CYCLOSPORINE

SOLUTION; ORAL
NEURAL

> ADD > AB + NOVARTIS

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> DLT >

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SANDIMUNE
BX + NOVARTIS

100MG/ML

100MG/ML

> ADD > AB

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SANCTA
SANGSTAT

100MG/ML

CYCLOHEPTADINE HYDROCHLORIDE

TABLET; ORAL

CYCLOHEPTADINE HCL

4MG

DACARBAZINE

INJECTABLE; INJECTION

DACARBAZINE

AP GENSLA SICOR PHARMS

200MG/VIAL

AP + BAYER

200MG/VIAL

DACTINOMYCIN

INJECTABLE; INJECTION

DACTINOMYCIN

AP + MERCK

0.5MG/VIAL

MS0716 001

JUL 14, 1995

MS0574 001

NOV 14, 1983

MS4195 001

OCT 31, 1998

MS0580 001

MS0529 002

AUG 27, 1998

MS17875 002

MS0682 001

DEXTROSE

INJECTABLE; INJECTION

DEXTROSE 5% IN PLASTIC CONTAINER

AP	B BRAUN	5GM/100ML; 31MG/100ML; 130MG/100ML;	N18738 001
		26MG/100ML; 320MG/100ML	N18738 002

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

INJECTABLE; INJECTION

ISOLYTE P IN DEXTROSE 5% IN PLASTIC CONTAINER

B BRAUN

5GM/100ML; 31MG/100ML; 130MG/100ML;	N19873 001
26MG/100ML; 320MG/100ML	

JUN 10, 1993

5GM/100ML; 31MG/100ML; 130MG/100ML;	N19873 001
26MG/100ML; 320MG/100ML	

JUN 10, 1993

DEXTROSE; MAGNESIUM CHLORIDE; POTASSIUM CHLORIDE; SODIUM ACETATE; SODIUM CHLORIDE

INJECTABLE; INJECTION

ISOLYTE H IN DEXTROSE 5% IN PLASTIC CONTAINER

B BRAUN

5GM/100ML; 30MG/100ML; 97MG/100ML;	N19844 001
220MG/100ML; 140MG/100ML	

JUN 10, 1993

5GM/100ML; 30MG/100ML; 97MG/100ML;	N19844 001
220MG/100ML; 140MG/100ML	

JUN 10, 1993

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

INJECTABLE; INJECTION

ISOLYTE S IN DEXTROSE 5% IN PLASTIC CONTAINER

B BRAUN

5GM/100ML; 30MG/100ML; 37MG/100ML;	N19843 001
370MG/100ML; 530MG/100ML;	
500MG/100ML	

AUG 09, 1993

5GM/100ML; 30MG/100ML; 37MG/100ML;	N19843 001
370MG/100ML; 530MG/100ML;	
500MG/100ML	

AUG 09, 1993

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

INJECTABLE; INJECTION

ISOLYTE S W/ DEXTROSE 5% IN PLASTIC CONTAINER

AP	B BRAUN	5GM/100ML; 30MG/100ML; 37MG/100ML;	
		370MG/100ML; 530MG/100ML;	
		500MG/100ML	N18274 001

DEXTROSE; POTASSIUM CHLORIDE

INJECTABLE; INJECTION

POTASSIUM CHLORIDE 0.075% IN DEXTROSE 5% IN PLASTIC CONTAINER

AP	B BRAUN	5GM/100ML; 75MG/100ML	N18744 001
			NOV 09, 1982
AP	MCCANN	5GM/100ML; 75MG/100ML	N18744 001
			NOV 09, 1982

DEXTROSE; POTASSIUM CHLORIDE; POTASSIUM PHOSPHATE, DIBASIC; SODIUM ACETATE; SODIUM CHLORIDE

INJECTABLE; INJECTION

ISOLYTE M IN DEXTROSE 5% IN PLASTIC CONTAINER

B BRAUN

5GM/100ML; 150MG/100ML; 130MG/100ML;	N19870 001
280MG/100ML; 91MG/100ML	

JUN 10, 1993

5GM/100ML; 150MG/100ML; 130MG/100ML;	N19870 001
280MG/100ML; 91MG/100ML	

JUN 10, 1993

DEXTROSE; SODIUM CHLORIDE

INJECTABLE; INJECTION

DEXTROSE 10% AND SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER

⊙ B BRAUN	10GM/100ML; 900MG/100ML	N18047 001
⊙ MCCANN	10GM/100ML; 900MG/100ML	N18047 001

DEXTROSE 5% AND SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER

⊙ B BRAUN	5GM/100ML; 900MG/100ML	N18026 001
⊙ MCCANN	5GM/100ML; 900MG/100ML	N18026 001

DILTIAZEM HYDROCHLORIDE

CAPSULE, EXTENDED RELEASE; ORAL

TIANAC

BC +

180MG

240MG

300MG

360MG

420MG

BC +

300MG

360MG

420MG

BC +

300MG

360MG

420MG

BC +

300MG

360MG

420MG

BC +

300MG

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420MG

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BC +

300MG

360MG

420MG

BC +

300MG

360MG

420MG

BC +

300MG

360MG

420MG

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INJECTABLE; INJECTION

DILTIAZEM HCL

ABBOTT

AP

TAYLOR PHARMA

5MG/ML

5MG/ML

5MG/ML

5MG/ML

5MG/ML

5MG/ML

5MG/ML

5MG/ML

5MG/ML

5MG/ML

5MG/ML

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DILTIAZEM MALATE

TABLET, EXTENDED RELEASE; ORAL

TIANAC

+ HOSCHST MARION RSSL

EQ 180MG HCL

EQ 240MG HCL

EQ 300MG HCL

EQ 360MG HCL

EQ 420MG HCL

EQ 480MG HCL

EQ 540MG HCL

EQ 600MG HCL

EQ 660MG HCL

EQ 720MG HCL

EQ 780MG HCL

EQ 840MG HCL

EQ 900MG HCL

EQ 960MG HCL

EQ 1020MG HCL

EQ 1080MG HCL

EQ 1140MG HCL

EQ 1200MG HCL

EQ 1260MG HCL

EQ 1320MG HCL

EQ 1380MG HCL

EQ 1440MG HCL

EQ 1500MG HCL

EQ 1560MG HCL

EQ 1620MG HCL

EQ 1680MG HCL

EQ 1740MG HCL

EQ 1800MG HCL

EQ 1860MG HCL

EQ 1920MG HCL

EQ 1980MG HCL

EQ 2040MG HCL

EQ 2100MG HCL

EQ 2160MG HCL

EQ 2220MG HCL

EQ 2280MG HCL

EQ 2340MG HCL

EQ 2400MG HCL

EQ 2460MG HCL

EQ 2520MG HCL

EQ 2580MG HCL

EQ 2640MG HCL

EQ 2700MG HCL

EQ 2760MG HCL

EQ 2820MG HCL

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N74941 001

APR 15, 1998

N75086 001

APR 09, 1998

N74084 001

FEB 25, 1994

N74084 002

FEB 25, 1994

N74084 003

FEB 25, 1994

N74084 004

FEB 25, 1994

N74084 005

FEB 25, 1994

N74084 006

FEB 25, 1994

N74084 007

FEB 25, 1994

N74084 008

FEB 25, 1994

N74084 009

FEB 25, 1994

N74084 010

FEB 25, 1994

N74084 011

FEB 25, 1994

5MG/ML

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DISCLAIMER

TABLET; ORAL
DIPHENIDOL
O POREPAC PHARM

5552

001 254298
 JUL 12, 1990

NO 2309 0221

W72986 001
MAR 29, 1991
W72987 001
MAR 29, 1991

DISOPYRANIDE PHOSPHATE

[REDACTED]



DOBUTAMINE HYDROCHLORIDE

**INJECTABLES; INJECTION
DOUBTING WCL**

WITTPOLD

NO 13.540 PAGE 2/22

W74545 001
JUN 25, 1998
W74279 001
FEB 18, 1998
W74995 001
MAR 31, 1998

LONG/VIA?

LONG/VIA?

LONG/VIA?

LONG/VIA?

APR 13, 1989
N62926 002
APR 13, 1989
N62926 003
APR 13, 1989

DORZOLANIDE HYDROCHLORIDE: TIMOLOL MALATE

**SOLUTION/DROPS; OPHTHALMIC
CISOPT
+ MERCK**

EQ 23 EAS; EQ 0.54 EASR 12069 001
APR 07, 1998

ESYB DM02 03

**N50744 001
SEP 30, 1998**

DOXEPIN HYDROCHLORIDE

CAPSULE; ORAL
DOXEPIN HCL

Mar 29, 1991

DA/ENCL 3

N16796 001

PROPERIDOL

INJECTABLE; INJECTION

AKORN MFG

10/1/96

N16796 001

ESTRADIOL

TABLET, ORAL
ESTRADIOL
BIOBEAVER

AA 0.5MG M00138 001
JAN 30, 1998
AA 1MG M00138 002
JAN 30, 1998
AA 2MG M00138 003
JAN 30, 1998

ESTRADIOL: NORTHYNDRONE ACETATE

FILM, EXTENDED RELEASE; TRANSDERMAL
COMBIPATCH
RHONE POULENC RORER

0.05MG/24HR; 0.14MG/24HR M20870 001
AUG 07, 1998
+ 0.05MG/24HR; 0.25MG/24HR M20870 002
AUG 07, 1998

ESTRADIOL CYPIONATE

INJECTABLE; INJECTION
DEPO-ESTRADIOL

1MG/ML M05470 001
3MG/ML M05470 002

ESTRADIOL VALERATE

INJECTABLE; INJECTION
DELESTROGEN

AO + BRISTOL MYERS SQUIBB 20MG/ML M09402 004
AO + 40MG/ML M09402 003
AO + 10MG/ML M09402 002

ESTRADIOL VALERATE: TESTOSTERONE ENANTHATE

INJECTABLE; INJECTION
DELAUDONE

• BRISTOL MYERS SQUIBB 4MG/ML; 90MG/ML M09545 001

ESTRADIOL VALERATE: TESTOSTERONE ENANTHATE

INJECTABLE; INJECTION
DELAUDONE

• BRISTOL MYERS SQUIBB 8MG/ML; 180MG/ML M09545 002
• BRISTOL MYERS SQUIBB 4MG/ML; 90MG/ML M09545 001

ESTROGENS, COMBINATION: MEDROXYPROGESTERONE ACETATE

TABLET, ORAL-28
PREMPRO 14/14

+ WYETH AYERST 0.625MG, 0.625MG; 5MG, 5MG M20527 003
JAN 09, 1998

ESTROGENS, ESTERIFIED

TABLET, ORAL
MENEST

BS MONARCH PHARMS 0.3MG M04951 001
BS 0.625MG M04948 001
BS 2.5MG M04949 001
BS 1.25MG M04950 001

ESTRONE

INJECTABLE; INJECTION
THRELIN

• BRISTOL MYERS SQUIBB 1MG/ML M03977 001
• BRISTOL MYERS SQUIBB 2MG/ML M03977 002
• BRISTOL MYERS SQUIBB 5MG/ML M03977 003

ETHACRYNATE SODIUM

INJECTABLE; INJECTION
EDSCRIN

+ MERCK BQ 50MG BASE/VIAL M16093 001

ETHACRYNATE SODIUM

INJECTABLE, INJECTION
EDCCLIN

25MG
50MG

W73594 001
DEC 13, 1993

ETHACRYNIC ACID

TABLET, ORAL

EDCCLIN
MERCK

25MG
50MG

W73594 001
DEC 13, 1993

ETHINYL ESTRADIOL, LEVONORGESTREL

TABLET, ORAL

PREVEN EMERGENCY CONTRACEPTIVE KIT
+ GYNETICS

N20946 001
SEP 01, 1998

TABLET, ORAL-21

ALESSE
BX + WYETH AYERST

0.02MG;0.1MG

N20683 001
MAR 27, 1997

LEVILITE

BX BERLEX LABS

0.02MG;0.1MG

N20860 001
JUL 13, 1998

LEVORA 0.15/30-21

BX WATSON LABS

0.03MG;0.15MG

N73592 001
DEC 13, 1993

TABLET, ORAL-28

ALESSE
BX WYETH AYERST

0.02MG;0.1MG

N20683 002
MAR 27, 1997

LEVILITE

BX BERLEX LABS

0.02MG;0.1MG

N20860 002
JUL 13, 1998

ETHINYL ESTRADIOL, LEVONORGESTREL

TABLET, ORAL-28

LEVILITE

W73594 001
DEC 13, 1993

LEVORA 0.15/30-28

BX WATSON LABS

0.03MG;0.15MG

ETHINYL ESTRADIOL, NORETHINDRONE

TABLET, ORAL-21

NORSTIN 1/35R-21

W71480 001
APR 12, 1988

BX WATSON LABS

0.035MG;1MG

TABLET, ORAL-28

NORSTIN 1/35R-28

W71481 001
APR 12, 1988

BX WATSON LABS

0.035MG;1MG

ETODOLAC

CAPSULE, ORAL

ETODOLAC
AESGEN

300MG

N74929 001
JAN 30, 1998

BX GENPHARM

200MG

N75071 001
SEP 30, 1998

BX TARO

300MG

N75071 002
SEP 30, 1998

BX TARO

200MG

N75078 001
APR 30, 1998

BX TARO

300MG

N75078 002
APR 30, 1998

TABLET, ORAL

ETODOLAC
CHELSEA LABS

400MG

N75069 001
APR 16, 1998

RESEARCH CREATS

INTEGRAL INSPECTION

ZENTANT, CITRATE PRESERVATIVE FREE
ABBOTT **NO. 0.95MG BASE/ML**

AP	ELKINS SING	NO 0.05MG BASE/ML
AP <td>NARSON <th>NO 0.05MG BASE/ML</th> </td>	NARSON <th>NO 0.05MG BASE/ML</th>	NO 0.05MG BASE/ML

NO. 0.05% BAST/IL

SUBSTITUTE PRESERVATIVE FREE

♦ JANSSEN

ITERIC AMMONIUM CITRATE

POWDER FOR RECONSTITUTION; ORAL

FERRISILTS
+ NYCOMED AMERSHAM
600MG/PACKET

ADD ADD DLT DLT

REVIEWS

Age Group	50MG	75MG	100MG	150MG
18-24	~10%	~10%	~10%	~10%
25-34	~10%	~10%	~10%	~10%
35-44	~10%	~10%	~10%	~10%
45-54	~10%	~10%	~10%	~10%

DEC 30, 1992
W19960 002
DEC 30, 1992
W19960 003
DEC 30, 1992

DEC 30, 1992

1996 003

DEC 30, 1992

11

PHOTOCHROMIC ACTINOPOLYMER

CREAM; TOPICAL

ADD A
ADD A
DLY A
AT

TARO

0.019

FLUOCINONIDE

CREAM: TOPICAL

AD 0.059

FLUOROURACIL

INJECTABLE; INJECTION

AP	+	50000 / YL
AP		50000 / YL
AP		50000 / YL

AP	AN PHARM PARTNERS	50MG/ML
AP		50MG/ML

24/0865

FLUPHENAZINE DICHLORIDE

INJECTABLES; INJECTION

NO. 2500/DT.
FLUORESCENT DECAONATE
KING PHARMS
INDUSTRIALS, INC.

FLURANDENOLOL; NITROXYN SULFATE

0.054:20 3.5MG BASE/GM

FLUPYRROXIDE, NEOTICIN SULFATE

TABLET, ORAL
LOVOX
+
0.05% BQ 3.5MG BASE/CM

N20345 001
DEC 05, 1994

> ADD >
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GANCICLOVIR

IMPLANT; IMPLANTATION
VITRASERT
+ BAUSCH AND LOMB
4.5MG

N20569 001
MAR 04, 1996

FLAVOXANINE MALEATE

TABLET, ORAL
LOVOX
+
25MG

N20243 001
DEC 05, 1994

GENFIBROIL

CAPSULE; ORAL
LOPID
+
200MG
300MG
PARKE DAVIS PHARMS

N18422 001
N18422 002

PONIVIRSEN SODIUM

INJECTABLE; INJECTION
VITRAVENE PRESERVATIVE FREE
+ CIBA VISION OPHTHLMC 6.6MG/ML

N20961 001
AUG 26, 1998

TABLET, ORAL
GENFIBROIL
TORPHARM
600MG

N75034 001
JUL 20, 1998

GABAPENTIN

CAPSULE; ORAL
NEUROMTIN
+
100MG
400MG

N20235 001
DEC 30, 1993
N20235 003
DEC 30, 1993

GENTAMICIN SULFATE

INJECTABLE; INJECTION
GENTAMICIN SULFATE IN SODIUM CHLORIDE 0.9% IN PLASTIC
CONTAINER
B BRAUN

AP EQ 40MG BASE/100ML M62814 008
AUG 28, 1987
AP EQ 60MG BASE/100ML M62814 009
AUG 28, 1987
AP EQ 70MG BASE/100ML M62814 010
AUG 28, 1987
AP EQ 0.8MG BASE/ML M62814 001
AUG 28, 1987
AP EQ 80MG BASE/100ML M62814 011
AUG 28, 1987
AP EQ 90MG BASE/100ML M62814 012
AUG 28, 1987
AP EQ 100MG BASE/100ML M62814 013
AUG 28, 1987

> DLT >
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TABLET, ORAL
NEUROMTIN
PARKE DAVIS
+
600MG
800MG

N20882 001
OCT 09, 1998
N20882 002
OCT 09, 1998

> ADD >
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> ADD >
> ADD >

GENTAMICIN SULFATE

INJECTABLE; INJECTION

GENTAMICIN SULFATE IN SODIUM CHLORIDE 0.9% IN PLASTIC

CONTAINER

B Braun

AP	EQ 1.2MG BASE/ML	N62814 002	AP
AP	EQ 1.2MG BASE/100ML	N62814 014	AP
AP	EQ 1.4MG BASE/ML	N62814 003	AP
AP	EQ 1.6MG BASE/ML	N62814 004	AP
AP	EQ 1.8MG BASE/ML	N62814 005	AP
AP	EQ 2MG BASE/ML	N62814 006	AP
AP	EQ 2.4MG BASE/ML	N62814 007	AP

> DLT >
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> ADD >
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GENTAMICIN SULFATE

ointment; ophthalmic

GENTAMICIN SULFATE

AP	AT	AKORN	EQ 0.3% BASE	N64093 001	AP
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GLUCAGON HYDROCHLORIDE RECOMBINANT

INJECTABLE; INJECTION

GLUCAGON

+ NOVO NORDISK

EQ 1MG BASE/VIAL

N20918 001
JUN 22, 1998

GLUCAGON. BIOSYNTHETIC

INJECTABLE; INJECTION

GLUCAGON

+ LILLY

1MG/VIAL

N20928 001
SEP 11, 1998

GLYBURIDE

TABLET; ORAL

GLYBURIDE (MICRONIZED)

INVAIED

1.5MG

N75174 001
JUN 22, 1998

3MG

N75174 002
JUN 22, 1998

AP

1.5MG

N74792 001
JUN 26, 1998

AP

3MG

N74792 002
JUN 26, 1998

GLYCOPYRROLATE

TABLET; ORAL

GLYCOPYRROLATE

1MG

N86902 001

2MG

N86900 001

AP

3MG

N86900 002

GLYCOPHOLATE

TABLET; ORAL

ROBINS AM
 ROBINSON PHARMS

1MG

M13827 001

2MG

M13827 002

M74517 002
 SEP 30, 1998

GRAMICIDIN; NEOMYCIN SULFATE; POLYMYXIN B SULFATE

SOLUTION/DROPS; OPHTHALMIC

NEOPHARM

AT + MONARCH PHARMS

0.025MG/ML, EQ 1.75MG BASE/ML;
 10.000 UNITS/ML

M6882 001

M13553 001

GRANISTRON HYDROCHLORIDE

TABLET; ORAL

KYTRIL
 SMITHKLINE BEECHAM

BQ 2MG BASE

M20305 002
 JUN 15, 1998

M74762 001
 JUN 25, 1997
 M74762 002
 JUN 25, 1997

GRAPFLOXACIN HYDROCHLORIDE

TABLET; ORAL

RAYAR

BQ 200MG BASE

M20695 001

BQ 400MG BASE

M20695 002

BQ 600MG BASE

M20695 003

MAY 14, 1998

HALOPERIDOL

TABLET; ORAL

HALOPERIDOL

10MG

M72113 001

20MG

M72353 001

GUANABENE ACETATE

TABLET; ORAL

GUANABENE ACETATE

BQ 4MG BASE

M74517 001
 SEP 30, 1998

GUANABENE ACETATE

TABLET; ORAL

GUANABENE ACETATE

BQ 8MG BASE

M74517 002
 SEP 30, 1998

GUANETHIDINE MONOSULFATE; HYDROCHLOROTHIAZIDE

10MG;25MG

M13553 001

GUANFACINE HYDROCHLORIDE

TABLET; ORAL

GUANFACINE HCL

BQ 1MG BASE

BQ 1MG BASE

BQ 2MG BASE

M74762 001
 JUN 25, 1997
 M74762 002
 JUN 25, 1997

HEPARIN SODIUM**INJECTABLE; INJECTION****HEPARIN SODIUM**

© AM PHARM PARTNERS

©

XXXXXXXXXX

10,000 UNITS/ML

20,000 UNITS/ML

20,000 UNITS/ML

20,000 UNITS/ML

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20,000 UNITS/ML

N17651 003

N17651 008

N17651 001

N17651 002

N17651 004

N17651 003

N17651 002

N17651 004

N17651 003

N17651 002

N17651 003

N17651 002

N17651 003

N17651 002

HEPARIN SODIUM 1,000 UNITS IN SODIUM CHLORIDE 0.9% IN**PLASTIC CONTAINER**

AP

B BRAUN

200 UNITS/100ML

N19953 001

JUL 20, 1992

©

200 UNITS/100ML

N19042 001

MAR 29, 1985

AP

MCNAM

200 UNITS/100ML

N19953 001

JUL 20, 1992

©

200 UNITS/100ML

N19042 001

MAR 29, 1985

HEPARIN SODIUM 10,000 UNITS IN DEXTROSE 5% IN**PLASTIC CONTAINER**

AP

ABBOTT

10,000 UNITS/100ML

N18911 004

JAN 30, 1985

©

10,000 UNITS/100ML

N18911 006

JAN 30, 1985

HEPARIN SODIUM 12,500 UNITS IN DEXTROSE 5% IN**PLASTIC CONTAINER**

AP

ABBOTT

5,000 UNITS/100ML

N18911 007

JAN 30, 1985

©

5,000 UNITS/100ML

N18911 007

JAN 30, 1985

HEPARIN SODIUM 12,500 UNITS IN DEXTROSE 5% IN PLASTIC**CONTAINER**

AP

ABBOTT

5,000 UNITS/100ML

N19339 001

MAR 27, 1985

5,000 UNITS/100ML

N19339 001

MAR 27, 1985

HEPARIN SODIUM 12,500 UNITS IN SODIUM CHLORIDE 0.45% IN**PLASTIC CONTAINER**

© B BRAUN

5,000 UNITS/100ML

N19802 001

JUL 20, 1992

© MCNAM

5,000 UNITS/100ML

N19802 001

JUL 20, 1992

HEPARIN SODIUM**INJECTABLE; INJECTION****HEPARIN SODIUM 2,000 UNITS IN SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER**

© B BRAUN

200 UNITS/100ML

N19042 002

MAR 29, 1985

© MCNAM

200 UNITS/100ML

N19042 002

MAR 29, 1985

HEPARIN SODIUM 20,000 UNITS IN DEXTROSE 5% IN PLASTIC**CONTAINER**

AP

B BRAUN

4,000 UNITS/100ML

N19952 001

JUL 20, 1992

AP

MCNAM

4,000 UNITS/100ML

N19952 001

JUL 20, 1992

HEPARIN SODIUM 25,000 UNITS IN DEXTROSE 5% IN**PLASTIC CONTAINER**

AP

ABBOTT

5,000 UNITS/100ML

N18911 009

JAN 30, 1985

AP

MCNAM

10,000 UNITS/100ML

N18911 009

JAN 30, 1985

HEPARIN SODIUM 25,000 UNITS IN DEXTROSE 5% IN PLASTIC**CONTAINER**

AP

B BRAUN

5,000 UNITS/100ML

N19952 004

JUL 20, 1992

AP

MCNAM

10,000 UNITS/100ML

N19952 005

JUL 20, 1992

HEPARIN SODIUM 5,000 UNITS IN SODIUM CHLORIDE 0.45% IN**PLASTIC CONTAINER**

AP

B BRAUN

5,000 UNITS/100ML

N19134 001

MAR 29, 1985

AP

MCNAM

5,000 UNITS/100ML

N19134 001

JUL 20, 1992

HEPARIN SODIUM 10,000 UNITS IN SODIUM CHLORIDE 0.45% IN**PLASTIC CONTAINER**

AP

B BRAUN

10,000 UNITS/100ML

N19134 001

MAR 29, 1985

AP

MCNAM

10,000 UNITS/100ML

N19134 001

JUL 20, 1992

HEPARIN SODIUM 25,000 UNITS IN SODIUM CHLORIDE 0.45% IN**PLASTIC CONTAINER**

AP

B BRAUN

5,000 UNITS/100ML

N19802 005

JUL 20, 1992

HEPARIN SODIUM 10,000 UNITS IN SODIUM CHLORIDE 0.45% IN**PLASTIC CONTAINER**

AP

B BRAUN

10,000 UNITS/100ML

N19802 002

JUL 20, 1992

HEPARIN SODIUM 5,000 UNITS IN SODIUM CHLORIDE 0.45% IN**PLASTIC CONTAINER**

AP

B BRAUN

5,000 UNITS/100ML

N19802 003

JUL 20, 1992

HEPARIN SODIUM 10,000 UNITS IN SODIUM CHLORIDE 0.45% IN**PLASTIC CONTAINER**

AP

B BRAUN

10,000 UNITS/100ML

N19802 004

JUL 20, 1992

HEPARIN SODIUM

INJECTABLE; INJECTION

HEPARIN SODIUM 25,000 UNITS IN SODIUM CHLORIDE 0.9% IN

PLASTIC CONTAINER

• B BRAUN	5,000 UNITS/100ML	N19135 001
		MAR 29, 1985
•	5,000 UNITS/100ML	N19802 003
		JUL 20, 1992
• HOGAN	5,000 UNITS/100ML	N19135 001
		MAR 29, 1985
•	5,000 UNITS/100ML	N19802 003
		JUL 20, 1992

HEPARIN SODIUM 5,000 UNITS IN SODIUM CHLORIDE 0.9% IN

PLASTIC CONTAINER

• B BRAUN	1,000 UNITS/100ML	N19042 004
		MAR 29, 1985
• HOGAN	1,000 UNITS/100ML	N19042 004
		MAR 29, 1985

HEPARIN SODIUM IN PLASTIC CONTAINER

AP	AM PHARM PARTNERS	1,000 UNITS/ML	N17029 013
			DEC 05, 1985
AP		5,000 UNITS/ML	N17029 014
			DEC 05, 1985
AP		10,000 UNITS/ML	N17029 015
			DEC 05, 1985
AP		20,000 UNITS/ML	N17029 016
			DEC 05, 1985
AP	FUJISAWA	1,000 UNITS/ML	N17029 013
			DEC 05, 1985
AP		5,000 UNITS/ML	N17029 014
			DEC 05, 1985
AP		10,000 UNITS/ML	N17029 015
			DEC 05, 1985
AP		20,000 UNITS/ML	N17029 016
			DEC 05, 1985

HEPARIN SODIUM PRESERVATIVE FREE

AP	+ AM PHARM PARTNERS	1,000 UNITS/ML	N17029 010
			APR 28, 1986
AP	+ FUJISAWA	1,000 UNITS/ML	N17029 010
			APR 28, 1986

HEPFLUSH-10

AP	AM PHARM PARTNERS	10 UNITS/ML	N17651 009
			JUN 26, 1984
AP	FUJISAWA	10 UNITS/ML	N17651 009
			JUN 26, 1984

HYDRALAZINE HYDROCHLORIDE

INJECTABLE; INJECTION

AP	• NOVARTIS	20MG/ML	N08303 003
			N08303 003
AP	HYDRALAZINE HCL	20MG/ML	N40136 001
			JUN 30, 1997
AP	+	20MG/ML	N40136 001
			JUN 30, 1997
AP	MOLOMAX	20MG/ML	N08517 001
			AUG 22, 1985
AP	+	20MG/ML	N08517 001
			AUG 22, 1985

HYDROCHLOROTHIAZIDE; IRBESARTAN

TABLET; ORAL

	AVAPRO HCT		
	• SANOFI	12.5MG;75MG	N20758 001
			SEP 30, 1997
+		12.5MG;150MG	N20758 002
			SEP 30, 1997
	IRBESARTAN-HYDROCHLOROTHIAZIDE		
	• SANOFI	12.5MG;75MG	N20758 001
			SEP 30, 1997
+		12.5MG;150MG	N20758 002
			SEP 30, 1997

HYDROCHLOROTHIAZIDE; METHYLDOPA

TABLET; ORAL

	METHYLDOPA AND HYDROCHLOROTHIAZIDE		
AP	INVASEO	15MG;250MG	N70829 001
			MAR 09, 1987
AP		25MG;250MG	N70830 001
			MAR 09, 1987
	•	15MG;250MG	N70829 001
			MAR 09, 1987
	•	25MG;250MG	N70830 001
			MAR 09, 1987

HYDROCHLOROTHIAZIDE; PROPRANOLOL HYDROCHLORIDE

TABLET; ORAL

PROPRANOLOL HCL AND HYDROCHLOROTHIAZIDE

AB	PARKE DAVIS	25MG;25MG	N71498 001
			DEC 18, 1991
AB		25MG;25MG	N71501 001
			DEC 18, 1991
0		25MG;40MG	N71498 001
			DEC 18, 1991
0		25MG;80MG	N71501 001
			DEC 18, 1991

HYDROCHLOROTHIAZIDE; RESERPINE

TABLET; ORAL

HYDOPRES 25

BP	MERCK	25MG;0.125MG	N11958 002
BP	MERCK SHARE DOWNE	25MG;0.125MG	N11958 002
BP	HYDOPRES 50	50MG;0.125MG	N11958 003
BP	MERCK SHARE DOWNE	50MG;0.125MG	N11958 003

HYDROCHLOROTHIAZIDE; SPIRONOLACTONE

TABLET; ORAL

SPIRONOLACTONE AND HYDROCHLOROTHIAZIDE

AB	GENEVA PHARMS	25MG;25MG	N86881 001
AB	SPIRONOLACTONE W/ HYDROCHLOROTHIAZIDE	25MG;25MG	N86881 001

HYDROCHLOROTHIAZIDE; TIMOLOL MALEATE

TABLET; ORAL

TIMOLIDE 10-25

+	MERCK	25MG;10MG	N18061 001
+	MERCK SHARE DOWNE	25MG;10MG	N18061 001

HYDROCHLOROTHIAZIDE; TRIAMTERENE

CAPSULE; ORAL

TRIAMTERENE AND HYDROCHLOROTHIAZIDE

AB	BARR	25MG;37.5MG	N74970 001
			JAN 06, 1998

HYDROCHLOROTHIAZIDE; TRIAMTERENE

TABLET; ORAL

TRIAMTERENE AND HYDROCHLOROTHIAZIDE

AB	BARR	25MG;37.5MG	N71251 002
			MAY 05, 1998

HYDROCHLOROTHIAZIDE; VALSARTAN

TABLET; ORAL

DIOVAN HCT

NOVARTIS

		12.5MG;80MG	N20818 001
+		12.5MG;160MG	N20818 002
			MAR 06, 1998

HYDROCORTISONE

AEROSOL; TOPICAL

AEROCOR-HC

+ ALLERGAN HEPBENT

> DLT >		0.5%	N85805 001
> DLT >		0.5%	N85805 001
> DLT >		0.5%	N85805 001
> ADD >		0.5%	N85805 001

CREAM; TOPICAL

AMUSOL HC

PARKE DAVIS

AT		2.5%	N88250 001
AT	PARKEDALE	2.5%	N88250 001
			JUN 06, 1984

ELDECORT

0.1%

0.1%

0.1%

0.1%

> DLT >		1%	N80459 001
> DLT >		2.5%	N80459 001
> ADD >		1%	N80459 001
> ADD >		2.5%	N80459 001

LOTION; TOPICAL

CETACORT

CALDERMA

> DLT >	AT	0.5%	N80426 002
> DLT >	AT	1%	N80426 002
> ADD >	AT	0.5%	N80426 002
> ADD >	AT	1%	N80426 002

SOLUTION; TOPICAL

TEXACORT

+ CHAMBERLAIN

+ MEDICIS

AT		1%	N80425 001
AT		1%	N80425 001

HYDROCORTISONE; NEOMYCIN SULFATE; POLYMYXIN B SULFATESUSPENSION/DROPS; OPHTHALMIC
CORTISPORIN

AB	* CLARKO PHARM	14, EQ 3.5MG BASE/ML; 10,000 UNITS/ML	N50169 001
AT	+ MONARCH PHARMS	14, EQ 3.5MG BASE/ML; 10,000 UNITS/ML	N50169 001

SUSPENSION/DROPS; OTIC
CORTISPORIN

AB	* CLARKO PHARM	14, EQ 3.5MG BASE/ML; 10,000 UNITS/ML	N60613 001
AT	+ MONARCH PHARMS	14, EQ 3.5MG BASE/ML; 10,000 UNITS/ML	N60613 001

HYDROCORTISONE VALERATE

CREAM; TOPICAL

AB	<u>HYDROCORTISONE VALERATE</u> COPLEY PHARM	0.2%	N74489 001 AUG 12, 1998
AB	TARO	0.2%	N75042 001 AUG 25, 1998
AB	<u>WESTCORT</u> + WESTWOOD SQUIBB	0.2%	N17950 001

OINTMENT; TOPICAL

AB	<u>HYDROCORTISONE VALERATE</u> TARO	0.2%	N75043 001 AUG 25, 1998
AB	<u>WESTCORT</u> + WESTWOOD SQUIBB	0.2%	N18726 001 AUG 08, 1983

HYDROMORPHONE HYDROCHLORIDE

SOLUTION; ORAL

AA	<u>DILAUDID</u> + KNOLL PHARM	5MG/5ML	N19891 001 DEC 07, 1992
AA	<u>HYDROMORPHONE HCL</u> ROXANE	5MG/5ML	N74653 001 JUL 29, 1998

TABLET; ORAL

AB	<u>DILAUDID</u> + KNOLL PHARM	8MG	N19892 001 DEC 07, 1992
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HYDROMORPHONE HYDROCHLORIDE

TABLET; ORAL

AB	<u>HYDROMORPHONE HCL</u> ROXANE	8MG	N74597 001 JUL 29, 1998
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HYDROXYAMPHETAMINE HYDROBROMIDE; TROPICAMIDE

SOLUTION/DROPS; OPHTHALMIC

	PAREMYD		N19261 001 JAN 30, 1992
+	AKORN	1%; 0.25%	
*	MYLAN	1%; 0.25%	N19261 001 JAN 30, 1992

HYDROXYCHLOROQUINE SULFATE

TABLET; ORAL

AB	<u>HYDROXYCHLOROQUINE SULFATE</u> MYLAN	200MG	N40274 001 MAY 29, 1998
AB	<u>HYDROXYCHLOROQUINE SULFATE</u> MYLAN	200MG	N40133 001 NOV 30, 1995
AB	WATSON LABS	200MG	N40133 001 NOV 30, 1995

HYDROXYUREA

CAPSULE; ORAL

	DROXIA		
	BRISTOL MYERS SQUIBB	300MG	N16295 002 FEB 25, 1998
		200MG	N16295 002
		300MG	N16295 003 FEB 25, 1998
		400MG	N16295 004 FEB 25, 1998

HYDREA

AB	+ BRISTOL MYERS SQUIBB	500MG	N16295 001
AB	* BRISTOL MYERS SQUIBB	500MG	N16295 001

HYDROXYUREA

AB	BARR	500MG	N75143 001 OCT 16, 1998
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> ADD >
> ADD >

HYDROXYUREA

CAPSULE; ORAL

HYDROXYUREA
DURAMED500MGN75020 001
JUL 30, 1998HYDROXYZINE HYDROCHLORIDE

TABLET; ORAL

HYDROXYZINE HCL
ROCHE LABS25MG25MG25MGAB WATSON LABS 10MGAB 25MGAB 50MGMAR 18, 1994
N81150 001
MAR 18, 1994
N81151 001
MAR 18, 1994HYDROXYZINE PAMOATE

CAPSULE; ORAL

HYDROXYZINE PAMOATE
LANEY PHARMEQ 50MG HCLEQ 100MG HCL

EQ 50MG HCL

EQ 100MG HCL

N87767 001
AUG 16, 1982
N87768 001
AUG 16, 1982
N87767 001
AUG 16, 1982
N87790 001
AUG 16, 1982IBUPROFEN

SUSPENSION; ORAL

CHILDREN'S ADVIL
LANEY PHARM100MG/5MLBX WHITEHALL ROBINS 100MG/5MLN19833 002
SEP 19, 1989
N19833 002
SEP 19, 1989IBUPROFEN

SUSPENSION; ORAL

IBUPROFEN
ALPHARMA100MG/5MLMOTRIM

AB + MCNEIL

100MG/5MLBX 100MG/5MLN74978 001
MAR 25, 1998N19842 001
SEP 19, 1989
N19842 001
SEP 19, 1989

TABLET; ORAL

IBUPROFEN
INVAMED400MG400MG400MG

EQ 400MG

EQ 600MG

EQ 800MG

N72144 001
JAN 14, 1988
N72065 001
JAN 14, 1988
N71938 001
JAN 14, 1988INDAPAMIDE

TABLET; ORAL

INDAPAMIDE
ALPHAPHARM1.25MGAB 2.5MGAB TEVA 1.25MGN75105 001
JUL 23, 1998
N75105 002
JUL 23, 1998
N74498 002
FEB 12, 1998INDOMETHACIN

CAPSULE; ORAL

INDOMETHACIN
LANEY PHARM25MG25MGN72894 001
JUL 31, 1981
N72897 001
JUL 31, 1981

INDOMETHACIN

CAPSULE; ORAL
INDOMETHACIN
• DAINBURY PHARMA

25MG

W7296 001
JUL 31, 1991

40.6¢

•

W7297 001
JUL 31, 1991

40.6¢

AP SON

75¢

W7466 001
MAY 28, 1998

51.3¢

INSULIN LISPRO

INJECTABLE; INJECTION
HUMALOG PEN
+ LILLY

100 UNITS/ML

W20563 002
AUG 06, 1998

IOVERSOL

INJECTABLE; INJECTION
OPTIRAY 240
+ WALLINGCROFT

51¢

W20923 001
MAY 28, 1998

INJECTABLE; INJECTION

OPTIRAY 320
+ WALLINGCROFT

68¢

W20923 002
MAY 29, 1998

•

W17903 001

74¢

W20923 003
MAY 28, 1998

IOPANIDOL

INJECTABLE; INJECTION
IOPANIDOL
ELKINS SINN

51¢

W74629 004
MAR 31, 1998

AEROSOL, METERED; INHALATION
ATROVENT
+

0.018MG/INH

W19085 001
DEC 29, 1986

W75005 001
FEB 24, 1998

0.018MG/INH

AP ABBOTT

51¢

W75005 002
FEB 24, 1998

ISOPROTERENOL HYDROCHLORIDE

AP ABBOTT

76¢

W75005 003
FEB 24, 1998

AEROSOL, METERED; INHALATION
ISUPREL
+

0.103MG/INH

W11178 001

LABETALOL HYDROCHLORIDE

TABLET; ORAL

LABETALOL HCL

AB ZENITH GOLDLINE 100MG
 AB 200MG
 AB 300MG

N74787 001 > DLT >
 AUG 03, 1998 > DLT >
 N74787 002
 AUG 03, 1998
 N74787 003
 AUG 03, 1998

LACTULOSE

POWDER FOR RECONSTITUTION; ORAL

LACTULOSE

10GM/PACKET
 + 10GM/PACKET

N74712 001
 DEC 10, 1997
 N74712 001
 DEC 10, 1997

LANOTRIGINE

TABLET, CHEWABLE; ORAL

LAMICTAL CD

GLAXO WELLCOME

5MG
 25MG
 + 100MG

N20764 001
 AUG 24, 1998
 N20764 002
 AUG 24, 1998
 N20764 003
 AUG 24, 1998

LEFLUNOMIDE

TABLET; ORAL

ARAVA

HOECHST MARION RSSL

> ADD > 10MG
 > ACC >
 > ADD > 20MG
 > ADD >
 > ADD > 100MG
 > ADD >
 > DLT > QUINTELAR 10MG
 > DLT >
 > DLT >
 > DLT >

N20905 001
 SEP 10, 1998
 N20905 002
 SEP 10, 1998
 N20905 003
 SEP 10, 1998
 N20905 001
 SEP 10, 1998
 N20905 002
 SEP 10, 1998

LEFLUNOMIDE

TABLET; ORAL

ARAVA

* QUINTELAR 100MG

N20905 001
 SEP 10, 1998

LEPIRUDIN

INJECTABLE; INJECTION

REFLUDAN

+ HOECHST MARION RSSL 50MG/VIAL

N20807 001
 MAR 06, 1998

LEUPROLIDE ACETATE

INJECTABLE; INJECTION

LEUPROLIDE ACETATE

A) BEDFORD LABS 1MG/0.2ML

N74728 001
 AUG 04, 1998

AP LUPRON
 + TAP HOLDINGS 1MG/0.2ML

N19010 001
 APR 09, 1985

LEVODOPA

CAPSULE; ORAL

DOPAR

ROBERTS LABS

100MG
 250MG
 500MG
 100MG
 250MG
 500MG

N16913 001
 N16913 001
 N16913 001
 N16913 003
 N16913 001
 N16913 002

LEVORPHANOL TARTRATE

INJECTABLE; INJECTION

LEVO-DROMORAN

+ ICN 2MG/ML

* ROCHE 2MG/ML

N08719 001
 DEC 19, 1991
 N08719 001
 DEC 19, 1991

LEVORPHANOL TARTRATE

TABLET; ORAL
LEVO-DRONORAN
+ ICH

2MG
[REDACTED]

M08720 001
DEC 19, 1991
[REDACTED]

LIDOCAINE

AEROSOL; ORAL
XYLOCAINE
+ ASTRA PHARMS

10t

M14394 001

DISC; ORAL
XYLOCAINE
[REDACTED]

[REDACTED]

LIDOCAINE; PRILLOCAINE

AEROSOL; TOPICAL
ENLA
+ ASTRA PHARMS

2.5t;2.5t

N209c2 001
FEB 04, 1998

LIDOCAINE HYDROCHLORIDE

INJECTABLE; INJECTION
[REDACTED]

[REDACTED]

[REDACTED]
[REDACTED]

[REDACTED]

INTL MEDICATION

LIDOCAINE HCL PRESERVATIVE FREE
ABBOTT

1t
1.5t
4t

M00408 001
M00408 002
M00295 001
MAY 17, 1984

LIDOCAINE HYDROCHLORIDE

INJECTABLE; INJECTION

LIDOCAINE HCL PRESERVATIVE FREE

AM PHARM PARTNERS

1t

M00404 002

AP

1t

M17584 001

AP

1t

M00404 003

AP

1t

M17584 002

AP

1t

M04625 001

AP

1t

M04625 002

AP

1t

M17702 001

AP

1t

M0302 001

AP

1t

M0302 002

AP

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M0302 002

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LOXAPINE

TABLET, ORALLY DISINTEGRATING; ORAL
CLARITIN PEDITABS
+ SCHERING

10MG
N20704 001
DEC 23, 1996

LOXAPINE

INJECTABLE; INJECTION

LOXAPINE C

AP 2MG/ML
AP ELKINS SINN 2MG/ML
AP 4MG/ML
AP TAYLOR 2MG/ML

W74974 001
JUL 23, 1998
W74496 001
SEP 28, 1998
W74496 002
SEP 28, 1998
W75025 001
JUL 23, 1998

TABLET; ORAL

LOXAPINE

AP 1MG
AP 2MG
AP 0.5MG
AP 1MG
AP 2MG

W72927 001
OCT 31, 1991
W72928 001
OCT 31, 1991

LOTEPREDNOL ETABONATE

SUSPENSION/DROPS; OPHTHALMIC
ALREX
+ PHARMOS

N20803 001
MAR 09, 1998

LOTEPREDNOL ETABONATE

SUSPENSION/DROPS; OPHTHALMIC
LOTENAX
+ PHARMOS
0.5%
0.5%

N20583 001
MAR 09, 1998
N20841 001
MAR 09, 1998

LOXAPINE HYDROCHLORIDE

CONCENTRATE; ORAL

LOXITANE C

+ WATSON LABS

EQ 25MG BASE/ML

N17658 001

INJECTABLE; INJECTION

LOXITANE IN

+ WATSON LABS

EQ 50MG BASE/ML

N18039 001

LOXAPINE SUCCINATE

CAPSULE; ORAL

LOXITANE

+ WATSON LABS

W17525 001
W17525 002
W17525 003
W17525 004

TABLET; ORAL

LOXITANE

+ WATSON LABS

EQ 10MG BASE
EQ 25MG BASE
EQ 50MG BASE

N17525 006
N17525 007
N17525 008

MAFENIDE ACETATECREAM; TOPICAL
SULFANYLON

+ BERTEK	EQ 85MG BASE/GM	N16763 001
+ MON HICKMAN	EQ 85MG BASE/GM	N16763 001

POWDER FOR RECONSTITUTION; TOPICAL
SULFANYLON

+ NYLAN	5g	N19832 003
		JUN 05, 1998

MAGNESIUM CHLORIDE; POTASSIUM CHLORIDE; POTASSIUM PHOSPHATE;
MONOBASIC; SODIUM ACETATE; SODIUM CHLORIDE; SODIUM GLUCONATE;
SODIUM PHOSPHATE, DIBASIC

INJECTABLE; INJECTION

ISOLYTE S PH 7.4 IN PLASTIC CONTAINER

B BRAUN	30MG/100ML; 37MG/100ML; 0.82MG/100ML; 370MG/100ML; 530MG/100ML; 500MG/100ML; 12MG/100ML	N19696 001
		SEP 29, 1989

MCGRAN	30MG/100ML; 37MG/100ML; 0.82MG/100ML; 370MG/100ML; 530MG/100ML; 500MG/100ML; 12MG/100ML	N19696 001
		SEP 29, 1989

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

INJECTABLE; INJECTION

ISOLYTE S IN PLASTIC CONTAINER

AP	B BRAUN	30MG/100ML; 37MG/100ML; 370MG/100ML; 530MG/100ML; 500MG/100ML	N18252 001
AP		30MG/100ML; 37MG/100ML; 370MG/100ML; 530MG/100ML; 500MG/100ML	N19711 001
			SEP 29, 1989

AP	MCGRAN	30MG/100ML; 37MG/100ML; 370MG/100ML; 530MG/100ML; 500MG/100ML	N18252 001
AP		30MG/100ML; 37MG/100ML; 370MG/100ML; 530MG/100ML; 500MG/100ML	N19711 001
			SEP 29, 1989

MALATHIONLOTION; TOPICAL
OVIDE

© GUNDBERN	0.5g	N18613 001
		AUG 02, 1982
© MEDICIS	0.5g	N18613 001
		AUG 02, 1982

MANNITOL

INJECTABLE; INJECTION

MANNITOL 10% IN PLASTIC CONTAINER

AP	B BRAUN	10GM/100ML	N20006 002
			JUL 26, 1993
AP	MCGRAN	10GM/100ML	N20006 002
			JUL 26, 1993

MANNITOL 15% IN PLASTIC CONTAINER

AP	B BRAUN	15GM/100ML	N20006 003
			JUL 26, 1993
AP	MCGRAN	15GM/100ML	N20006 003
			JUL 26, 1993

MANNITOL 20%

AP	B BRAUN	20GM/100ML	N14738 001
AP	MCGRAN	20GM/100ML	N14738 001

MANNITOL 20% IN PLASTIC CONTAINER

AP	B BRAUN	20GM/100ML	N20006 004
			JUL 26, 1993
AP	MCGRAN	20GM/100ML	N20006 004
			JUL 26, 1993

MANNITOL 5% IN PLASTIC CONTAINER

AP	B BRAUN	5GM/100ML	N20006 001
			JUL 26, 1993
AP	MCGRAN	5GM/100ML	N20006 001
			JUL 26, 1993

SOLUTION; IRRIGATION

RESECTISOL IN PLASTIC CONTAINER

B BRAUN	5GM/100ML	N16772 002
MCGRAN	5GM/100ML	N16772 002

MECAMYLAMINE HYDROCHLORIDE

TABLET; ORAL

INVERSINE

+ LAYTON	2.5MG	N10251 001
+ MERCK SHARP DOHME	2.5MG	N10251 001

MEGESTROL ACETATE

TABLET; ORAL
MEGESTROL ACETATE
 AB PHARMACHEMIE 40MG N74745 001
 FEB 27, 1998

NEPERIDINE HYDROCHLORIDE

TABLET; ORAL
NEPERIDINE HCL
 AA ROXANE 25MG N40186 001
 JUN 30, 1997
 AA WATSON LABS 50MG N40186 001
 JUN 30, 1997

NEPROBAMATE

TABLET; ORAL
NEPROBAMATE
 AA 600MG N84274 001

NESALAMINE

CAPSULE, EXTENDED RELEASE; ORAL
 PENTASA
 + ROBERTS LABS 250MG N20049 001
 MAY 10, 1993

MESTRANOL; NORETHINDRONE

TABLET; ORAL-21
NORETHIN 1/50M-21
 AB 0.05MG;1MG N71539 001
 APR 12, 1988
 AB WATSON LABS 0.05MG;1MG N71539 001
 APR 12, 1988

TABLET; ORAL-28
NORETHIN 1/50M-28
 AB 0.05MG;1MG N71540 001
 APR 12, 1988

MESTRANOL; NORETHINDRONE

TABLET; ORAL-28
NORETHIN 1/50M-28
 AB WATSON LABS 0.05MG;1MG N71540 001
 APR 12, 1988

METHADONE HYDROCHLORIDE

CONCENTRATE; ORAL
METHADONE HCL
 AA ROXANE 10MG/ML N40180 001
 APR 30, 1998

TABLET; ORAL
METHADONE HCL
 AA SON 5MG N40241 001
 MAY 29, 1998
 AA 10MG N40241 002
 MAY 29, 1998

TABLET, DISPERSIBLE; ORAL
METHADONE HCL
 AA SON 40MG N75082 001
 MAR 25, 1998

METHOCARBAMOL

INJECTABLE; INJECTION
METHOCARBAMOL
 AA 100MG/ML N89849 001
 DEC 27, 1991

TABLET; ORAL
METHOCARBAMOL
 AA 750MG N85136 001
 N85137 001

METOCLOPRAMIDE HYDROCHLORIDE

TABLET; ORAL
METOCLOPRAMIDE HCL
SANTALINI

EQ 5MG BASE

N72839 001
JUN 21, 1989

EQ 10MG BASE

N72840 001
JUN 21, 1989

EQ 5MG BASE

N72436 001
JUN 23, 1989

EQ 10MG BASE

N70850 001
FEB 03, 1987

METOPROLOL TARTRATE

INJECTABLE; INJECTION
METOPROLOL TARTRATE
ABBOTT

1MG/ML

N75160 001
JUL 05, 1998

MEKILITINE HYDROCHLORIDE

CAPSULE; ORAL
MEKILITINE HCL
DANBURY PHARMA

150MG

N74865 001
APR 13, 1998

200MG

N74865 002
APR 13, 1998

250MG

N74865 003
APR 13, 1998

MIDAZOLAM HYDROCHLORIDE

SYRUP; ORAL
VERSED
+ ROCHE

EQ 2MG BASE/ML

N20942 001
OCT 15, 1998

MITOMYCIN

INJECTABLE; INJECTION
MITOMYCIN
SUPERNEN

5MG/VIAL

N64144 001
APR 30, 1998

MITOMYCIN

INJECTABLE; INJECTION
MITOMYCIN
SUPERNEN

20MG/VIAL

N64144 002
APR 30, 1998

MONTILUKAST SODIUM

TABLET; ORAL
SINGULAIR
+ MERCK

EQ 10MG BASE

N20829 002
FEB 20, 1998

TABLET, CHEWABLE; ORAL
SINGULAIR
+ MERCK

EQ 5MG BASE

N20830 001
FEB 20, 1998

MORPHINE SULFATE

CAPSULE, EXTENDED RELEASE; ORAL
KADIAN

+ FAULDING

20MG

N20616 001
JUL 03, 1996

+

50MG

N20616 002
JUL 03, 1996

+

100MG

N20616 003
JUL 03, 1996

+

200MG

N20616 004
JUL 03, 1996

+

300MG

N20616 005
JUL 03, 1996

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

TABLET, EXTENDED RELEASE; ORAL
MORPHINE SULFATE
AB GENERICS

AB

15MG

N74862 001
JUL 07, 1998

AB

30MG

N74862 002
JUL 07, 1998

AB

60MG

N74862 003
JUL 07, 1998

AB

100MG

N74769 001
JUL 02, 1998

MORPHINE SULFATE

TABLET, EXTENDED RELEASE; ORAL

MORPHINE SULFATE

AB GENERICS

AB 200MG

> ADD > 200MG

> ADD > 15MG

> ADD > 30MG

> ADD > 60MG

> ADD > 15MG

AB 15MG

AB 30MG

AB 60MG

AB 100MG

AB 200MG

AB 300MG

AB 400MG

AB 500MG

AB 600MG

AB 700MG

AB 800MG

AB 900MG

AB 1000MG

AB 1100MG

AB 1200MG

AB 1300MG

AB 1400MG

AB 1500MG

MYCOPHENOLATE MOFETIL HYDROCHLORIDE

SUSPENSION; ORAL

CELLCEPT

+ ROCHE GLOBAL

> ADD >

> ADD >

> ADD >

200MG/ML

MYCOPHENOLATE MOFETIL HYDROCHLORIDE

INJECTABLE; INJECTION

CELLCEPT

+ ROCHE GLOBAL

500MG/VIAL

NADOLOL

TABLET; ORAL

CONCARD

BRISTOL MYERS SQUIBB

AB 20MG

AB 40MG

AB 80MG

AB 120MG

AB 160MG

AB 200MG

AB 240MG

AB 280MG

AB 320MG

AB 360MG

AB 400MG

AB 440MG

AB 480MG

AB 520MG

AB 560MG

AB 600MG

AB 640MG

AB 680MG

AB 720MG

AB 760MG

AB 800MG

AB 840MG

AB 880MG

AB 920MG

AB 960MG

AB 1000MG

AB 1040MG

AB 1080MG

AB 1120MG

AB 1160MG

AB 1200MG

AB 1240MG

AB 1280MG

AB 1320MG

AB 1360MG

AB 1400MG

AB 1440MG

AB 1480MG

AB 1520MG

AB 1560MG

AB 1600MG

AB 1640MG

AB 1680MG

AB 1720MG

AB 1760MG

AB 1800MG

AB 1840MG

AB 1880MG

AB 1920MG

AB 1960MG

AB 2000MG

AB 2040MG

AB 2080MG

AB 2120MG

AB 2160MG

W7479 002

JUL 02, 1998

W7329 001

OCT 28, 1998

W7329 002

OCT 28, 1998

W7329 003

OCT 28, 1998

W7329 004

SEP 12, 1998

W7329 005

MAY 29, 1987

W7329 006

APR 08, 1988

W7329 007

JAN 16, 1990

W7329 008

SEP 12, 1998

W7329 009

MAY 29, 1987

W7329 010

APR 08, 1988

W7329 011

JAN 16, 1990

W7329 012

SEP 12, 1998

W7329 013

MAY 29, 1987

W7329 014

APR 08, 1988

W7329 015

JAN 16, 1990

W7329 016

SEP 12, 1998

W7329 017

MAY 29, 1987

W7329 018

APR 08, 1988

W7329 019

JAN 16, 1990

W7329 020

SEP 12, 1998

W7329 021

MAY 29, 1987

W7329 022

APR 08, 1988

W7329 023

JAN 16, 1990

W7329 024

SEP 12, 1998

W7329 025

MAY 29, 1987

W7329 026

APR 08, 1988

W7329 027

JAN 16, 1990

W7329 028

SEP 12, 1998

W7329 029

MAY 29, 1987

W7329 030

APR 08, 1988

W7329 031

W7479 001

JUL 02, 1998

W7329 001

OCT 28, 1998

W7329 002

OCT 28, 1998

W7329 003

OCT 28, 1998

W7329 004

SEP 12, 1998

W7329 005

MAY 29, 1987

W7329 006

APR 08, 1988

W7329 007

JAN 16, 1990

W7329 008

SEP 12, 1998

W7329 009

MAY 29, 1987

W7329 010

APR 08, 1988

W7329 011

JAN 16, 1990

W7329 012

SEP 12, 1998

W7329 013

MAY 29, 1987

W7329 014

APR 08, 1988

W7329 015

JAN 16, 1990

W7329 016

SEP 12, 1998

W7329 017

MAY 29, 1987

W7329 018

APR 08, 1988

W7329 019

JAN 16, 1990

W7329 020

SEP 12, 1998

W7329 021

MAY 29, 1987

W7329 022

APR 08, 1988

W7329 023

JAN 16, 1990

W7329 024

SEP 12, 1998

W7329 025

MAY 29, 1987

W7329 026

APR 08, 1988

W7329 027

JAN 16, 1990

W7329 028

SEP 12, 1998

W7329 029

MAY 29, 1987

W7329 030

APR 08, 1988

W7329 031

W7479 001

JUL 02, 1998

W7329 001

OCT 28, 1998

W7329 002

OCT 28, 1998

W7329 003

OCT 28, 1998

W7329 004

SEP 12, 1998

W7329 005

MAY 29, 1987

W7329 006

APR 08, 1988

W7329 007

JAN 16, 1990

W7329 008

SEP 12, 1998

W7329 009

MAY 29, 1987

W7329 010

APR 08, 1988

W7329 011

JAN 16, 1990

W7329 012

SEP 12, 1998

W7329 013

MAY 29, 1987

W7329 014

APR 08, 1988

W7329 015

JAN 16, 1990

W7329 016

SEP 12, 1998

W7329 017

MAY 29, 1987

W7329 018

APR 08, 1988

W7329 019

JAN 16, 1990

W7329 020

SEP 12, 1998

NALTREXONE HYDROCHLORIDE

TABLET; ORAL
NALTREXONE HCL
AB BARR 50MG N74918 001
MAY 08, 1998

REZIA
AB + DUPONT MERCK 50MG N18932 001
NOV 20, 1984

NAPROXEN

TABLET, DELAYED RELEASE; ORAL
EC-NAPROSYN
12 + SYNTEX 375MG N20067 002
OCT 14, 1994

AB + 500MG N20067 003
OCT 14, 1994

NAPROXEN
AB INVAMED 375MG N75061 001
FEB 18, 1998

AB 500MG N75061 002
FEB 18, 1998

AB PUREPAC PHARM 375MG N74936 001
FEB 24, 1998

AB 500MG N74936 002
FEB 24, 1998

AB TEVA 375MG N75227 001
JUN 30, 1998

AB 500MG N75227 002
JUN 30, 1998

NAPROXEN SODIUM

TABLET; ORAL
NAPROXEN SODIUM
AB AL HIKMA EQ 250MG BASE N74480 002
FEB 18, 1998

> ADD > AB INVAMED EQ 250MG BASE N74495 001
> ADD > DEC 05, 1994

> DLT > AB EQ 250MG BASE N74495 002
> DLT > DEC 05, 1994

> ADD > AB EQ 500MG BASE N74495 002
> ADD > DEC 05, 1994

> DLT > AB EQ 500MG BASE N74495 002
> DLT > DEC 05, 1994

NARATRIPTAN HYDROCHLORIDE

TABLET; ORAL
AMERGE
GLAXO WELLCOME EQ 1MG BASE N20763 002
FEB 10, 1998

+ EQ 2.5MG BASE N20763 001
FEB 10, 1998

NEOMYCIN SULFATE

TABLET; ORAL
NEOMYCIN SULFATE
TEVA EQ 150MG BASE N60304 002
+ EQ 350MG BASE N60304 001

NEOMYCIN SULFATE; POLYMYXIN B SULFATE

SOLUTION; IRRIGATION
NEOSPORIN G.U. IRRIGANT
AT GLAXO WELLCOME EQ 40MG BASE/ML; 200,000 UNITS/ML N60707 002
AT MONARCH PHARMS EQ 40MG BASE/ML; 200,000 UNITS/ML N60707 001

NEVIRAPINE

SUSPENSION; ORAL
VIRAMUNE
+ BOEHRINGER INGELHEIM 50MG/5ML N20933 001
SEP 11, 1998

NIACIN

TABLET; ORAL
NIACIN
AB GENPHARM 500MG N83305 001
500MG

NICARDIPINE HYDROCHLORIDE

CAPSULE; ORAL
NICARDIPINE HCL
AB GENPHARM 20MG N74928 001
MAR 19, 1998

NORTHINDOLINE ACETATE

TABLET, ORAL

AVANTIN

5MG

AB

AVANTIN

5MG

AVANTIN

5MG

MYSTATIN

SUSPENSION, ORAL

MYSTATIN

UOL

100,000 UNITS/ML

N64142 001

JUN 25, 1998

OLANZAPINE

TABLET, ORAL

ZYPRIA

ZYPRIA

2.5MG

15MG

20MG

N20592 001

SEP 30, 1996

N20592 005

SEP 09, 1997

N20592 006

SEP 09, 1997

ONEPRAZOLE

CAPSULE, DELAYED REL PELLETS, ORAL

PRIOSEC

PRIOSEC

40MG

N19810 002

JAN 15, 1998

ORPHENDRINE CITRATE

TABLET, EXTENDED RELEASE, ORAL

ORPHEN

3H

100MG

N12157 001

ORPHENDRINE CITRATE

TABLET, EXTENDED RELEASE, ORAL

ORPHENDRINE CITRATE

100MG

N40284 001

JUN 19, 1998

OXAZEPAM

TABLET, ORAL

OXAZEPAM

OXAZEPAM

OXAZEPAM

OXAZEPAM

OXAZEPAM

OXAZEPAM

OXAZEPAM

OXYBUTYRIN CHLORIDE

SYRUP, ORAL

OXYBUTYRIN CHLORIDE

OXYBUTYRIN CHLORIDE

OXYBUTYRIN CHLORIDE

OXYBUTYRIN CHLORIDE

OXYBUTYRIN CHLORIDE

OXYBUTYRIN CHLORIDE

OXYBUTYRIN CHLORIDE

OXYBUTYRIN CHLORIDE

OXYCODONE HYDROCHLORIDE

TABLET, EXTENDED RELEASE, ORAL

OXYCONTIN

OXYCONTIN

OXYCONTIN

OXYCONTIN

OXYCONTIN

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OXYCONTIN

OXYCONTIN

OXYCONTIN

N20553 001

DEC 12, 1995

N20932 001

OCT 26, 1998

N20932 002

OCT 26, 1998

N20932 003

OCT 26, 1998

N20932 004

OCT 26, 1998

N20932 005

OCT 26, 1998

N20932 006

OCT 26, 1998

N20932 007

OCT 26, 1998

N20932 008

OCT 26, 1998

N20932 009

OCT 26, 1998

N20932 010

OCT 26, 1998

N10211 001

SEP 10, 1996

N17577 001

SEP 10, 1996

N17577 002

SEP 10, 1996

N17577 003

SEP 10, 1996

N17577 004

SEP 10, 1996

N17577 005

SEP 10, 1996

N17577 006

SEP 10, 1996

N17577 007

SEP 10, 1996

N17577 008

SEP 10, 1996

N17577 009

SEP 10, 1996

N17577 010

SEP 10, 1996

N17577 011

SEP 10, 1996

N17577 012

SEP 10, 1996

N17577 013

SEP 10, 1996

N17577 014

SEP 10, 1996

N17577 015

SEP 10, 1996

OXITETRACYCLINE

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

OXITETRACYCLINE
CAPSULE, ORAL
FAXIL
+ SMITHKLINE BEECHAM

250MG

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

CAPSULE, ORAL
FAXIL
+ SMITHKLINE BEECHAM

EQ 40MG BASE

N20885 004
SEP 09, 1998

OXYTOCIN

INJECTABLE, INJECTION

PITOCIN

AP + PARKEDALE

10 USP UNITS/ML

N18261 001

PENTOSAN POLYSULFATE SODIUM

CAPSULE, ORAL

ELMIRON

+ ALZA

100MG

N20193 001
SEP 26, 1996

PARICALCITOL

INJECTABLE, INJECTION

ZEMPLAR

+ ABBOTT

0.005MG/ML

N20819 001
APR 17, 1998

PAROMYCIN SULFATE

CAPSULE, ORAL

MONATIN

KING PHARMS

+ PARKEDALE

EQ 250MG BASE

EQ 250MG BASE

EQ 250MG BASE

EQ 250MG BASE

N62310 001

N60521 001

N64171 001

JUN 30, 1997

PENTOXIFYLLINE

TABLET, EXTENDED RELEASE, ORAL

PENTOXIFYLLINE

BIOVAIL

400MG

400MG

N75028 001
JUL 20, 1998
N75107 001
SEP 04, 1998

PERMETHRIN

CREAM, TOPICAL

ELIMITE

AB + ALLERGAN

5g

N19855 001
AUG 25, 1989

PERMETHRIN

ALPHA

5g

N74806 001
JAN 23, 1998

PAROXETINE HYDROCHLORIDE

CAPSULE, ORAL

FAXIL

SMITHKLINE BEECHAM

EQ 10MG BASE

EQ 20MG BASE

EQ 30MG BASE

N20885 001

OCT 09, 1998

N20885 002

OCT 09, 1998

N20885 003

OCT 09, 1998

PHENAZOPRINE HYDROCHLORIDE, SULFAMETHOXAZOLE

CAPSULE, ORAL

FAXIL

SMITHKLINE BEECHAM

100MG/500MG

100MG/500MG

N13294 001
SEP 10, 1987
N13294 001
SEP 10, 1987

PHENAZOPRINE HYDROCHLORIDE, SULFISOXAZOLE

~~XXXXXXXXXX~~
~~XXXXXXXXXX~~
~~XXXXXXXXXX~~
•
50MG;500MG

~~XXXXXXXXXX~~
~~XXXXXXXXXX~~
N19158 001
AUG 31, 1990

> ADD >
> DEL >

PILOCARPINE

DRUG DELIVERY SYSTEM; OPHTHALMIC
OCUSERT PILO-40

11MG
~~XXXXXXXXXX~~

N17548 001
~~XXXXXXXXXX~~

PHENTERMINE HYDROCHLORIDE

CAPSULE; ORAL
PHENTERMINE HCL

37.5MG
37.5MG

~~XXXXXXXXXX~~
~~XXXXXXXXXX~~
N88414 001
OCT 19, 1983

TABLET; ORAL
PHENTERMINE HCL

30MG
30MG

~~XXXXXXXXXX~~
~~XXXXXXXXXX~~
N88605 001
SEP 26, 1987

PHENTOLAMINE MESYLATE

INJECTABLE; INJECTION
PHENTOLAMINE MESYLATE

5MG/VIAL

N40335 001
MAR 11, 1998

REGITINE
AP + NOVARTIS

5MG/VIAL

N08278 003

PHENYTOIN SODIUM, PROMPT

CAPSULE; ORAL
PROMPT PHENYTOIN SODIUM

100MG
100MG

~~XXXXXXXXXX~~
~~XXXXXXXXXX~~
N80905 001
N80259 001

PIPERACILLIN SODIUM; TAZOBACTAM SODIUM

INJECTABLE; INJECTION
ZOSYN IN PLASTIC CONTAINER
+ LEDERLE

EQ 40MG BASE/ML;
EQ 5MG BASE/ML

N50750 001
FEB 24, 1998

EQ 4GM BASE/100ML;
EQ 500MG BASE/100ML

N50750 003
FEB 24, 1998

EQ 60MG BASE/ML;
EQ 7.5MG BASE/ML

N50750 002
FEB 24, 1998

POTASSIUM CHLORIDE; SODIUM CHLORIDEINJECTABLE; INJECTIONPOTASSIUM CHLORIDE 0.01% IN SODIUM CHLORIDE 0.9% INPLASTIC CONTAINER

B BRAUN 37MG/100ML;900MG/100ML N19708 001

SEP 29, 1989

● 37MG/100ML;900MG/100ML N19708 001

SEP 29, 1989

POTASSIUM CHLORIDE 0.075% IN SODIUM CHLORIDE 0.9% INPLASTIC CONTAINER

B BRAUN 75MG/100ML;900MG/100ML N19708 002

SEP 29, 1989

● 75MG/100ML;900MG/100ML N19708 002

SEP 29, 1989

POTASSIUM CHLORIDE 0.11% IN SODIUM CHLORIDE 0.9% IN PLASTICCONTAINER

B BRAUN 110MG/100ML;900MG/100ML N19708 003

SEP 29, 1989

● 110MG/100ML;900MG/100ML N19708 003

SEP 29, 1989

POTASSIUM CHLORIDE 0.22% IN SODIUM CHLORIDE 0.9% IN PLASTICCONTAINER

B BRAUN 220MG/100ML;900MG/100ML N19708 005

SEP 29, 1989

● 220MG/100ML;900MG/100ML N19708 005

SEP 29, 1989

POTASSIUM CHLORIDE 0.3% IN SODIUM CHLORIDE 0.9% IN PLASTICCONTAINER

B BRAUN 300MG/100ML;900MG/100ML N19708 006

SEP 29, 1989

● 300MG/100ML;900MG/100ML N19708 006

SEP 29, 1989

SODIUM CHLORIDE 0.9% AND POTASSIUM CHLORIDE 0.075% INPLASTIC CONTAINER

● B BRAUN 75MG/100ML;900MG/100ML N18722 001

NOV 09, 1982

● MCNAM 75MG/100ML;900MG/100ML N18722 001

NOV 09, 1982

SODIUM CHLORIDE 0.9% AND POTASSIUM CHLORIDE 0.15% INPLASTIC CONTAINER

● B BRAUN 150MG/100ML;900MG/100ML N18722 002

NOV 09, 1982

● MCNAM 150MG/100ML;900MG/100ML N18722 002

NOV 09, 1982

SODIUM CHLORIDE 0.9% AND POTASSIUM CHLORIDE 0.22% INPLASTIC CONTAINER

● B BRAUN 220MG/100ML;900MG/100ML N18722 003

NOV 09, 1982

POTASSIUM CHLORIDE; SODIUM CHLORIDEINJECTABLE; INJECTIONSODIUM CHLORIDE 0.9% AND POTASSIUM CHLORIDE 0.22% INPLASTIC CONTAINER

● MCNAM 220MG/100ML;900MG/100ML N18722 003

NOV 09, 1982

SODIUM CHLORIDE 0.9% AND POTASSIUM CHLORIDE 0.3% IN PLASTICCONTAINER

● B BRAUN 300MG/100ML;900MG/100ML N18722 004

NOV 09, 1982

● MCNAM 300MG/100ML;900MG/100ML N18722 004

NOV 09, 1982

PRAMIPEXOLE DIHYDROCHLORIDETABLET; ORALMIRAPEX

PHARMACIA AND UPJOHN 0.5MG

N20667 006

FEB 12, 1998

PREDNISOLONESYRUP; ORALPRE-PRED

AA WE PHARMS 15MG/5ML

N40192 001

MAY 28, 1998

AA PREDNISOLONE

WE PHARMS 15MG/5ML

N40192 001

MAY 28, 1998

AA PRELONE

+ NURO 15MG/5ML

N89081 001

FEB 04, 1986

TABLET; ORALPREDNISOLONE

BX DARBONT PHARMA 5MG

N80354 001

BX + 5MG

N80354 001

BX GENOVA PHARMA 5MG

N80339 001

● 5MG

N80339 001

PREDNISOLONE SODIUM PHOSPHATE; SULFACETAMIDE SODIUMSOLUTION/DROPS; OPHTHALMICAA GENOVA

10 0.1% PHOSPHATE 101

N74113 001

JUN 29, 1998

PREDNISOLONE SODIUM PHOSPHATE, SULFACETAMIDE SODIUM

SOLUTION/DROPS; OPHTHALMIC

EQ 0.23% PHOSPHATE:10%
N74511 001
JUL 30, 1996

PREDNISONE

TABLET; ORAL

PREDNISONE

5MG
10MG
20MG

PRINIDONE

SUSPENSION; ORAL

MYSOLINE

+ ELAN PHARMA

250MG/5ML

TABLET; ORAL

MYSOLINE

+ ELAN PHARMA

250MG
500MG

PROCAINAMIDE HYDROCHLORIDE

TABLET, EXTENDED RELEASE; ORAL

PROCAINAMIDE HCL

500MG
250MG

PROCAIN SR

PROCAINAMIDE HYDROCHLORIDE

TABLET, EXTENDED RELEASE; ORAL

PROCAIN SR

PARKE-DAVIS

750MG

AB + PARKEDALE

500MG

750MG

100MG

PROCHLORPERAZINE EDISYLATE

INJECTABLE; INJECTION

PROCHLORPERAZINE EDISYLATE

PARKE-DAVIS

EQ 5MG BASE/ML

PROCHLORPERAZINE MALEATE

TABLET; ORAL

PROCHLORPERAZINE MALEATE

EQ 5MG BASE

EQ 10MG BASE

EQ 5MG BASE

EQ 10MG BASE

EQ 5MG BASE

EQ 10MG BASE

PROGESTERONE

CAPSULE; ORAL

PROMETRIUM

+ SCHERING PLOUGH

100MG

N77510 001
APR 01, 1982
N77510 001
JAN 16, 1985
N76065 001
N77510 001
APR 01, 1982
N78489 001
JAN 16, 1985

N77510 001
APR 01, 1982
N77510 001
JAN 16, 1985
N76065 001
N77510 001
APR 01, 1982
N78489 001
JAN 16, 1985

N40268 001
FEB 27, 1998
N40268 002
FEB 27, 1998
N40162 001
JAN 20, 1998
N40162 002
JAN 20, 1998

N19781 001
MAY 14, 1998

PROMETHAZINE HYDROCHLORIDE

INJECTABLE; INJECTION

PROMETHAZINE HCL

25MG/ML

N89463 001

50MG/ML

N89477 001

25MG/ML

N89463 001

50MG/ML

N89477 001

SYRUP; ORAL

PROMETHAZINE HCL

5.25MG

N89463 001

TABLET; ORAL

PROMETHAZINE HCL

12.5MG

N83712 001

PROPORYPHENE HYDROCHLORIDE

CAPSULE; ORAL

PROPORYPHENE HCL

65MG

N80908 002

65MG

N83278 001

PROPRANOLOL HYDROCHLORIDE

TABLET; ORAL

PROPRANOLOL HCL

60MG

N71098 001

PROPRANOLOL HYDROCHLORIDE

TABLET; ORAL

PROPRANOLOL HCL

10MG

N71418 001

20MG

N71687 001

40MG

N71688 001

60MG

N72197 001

80MG

N71689 001

90MG

N72198 001

QUINIDINE SULFATE

TABLET; ORAL

QUINIDINE SULFATE

100MG

N85584 001

200MG

N84003 001

TABLET, EXTENDED RELEASE; ORAL

QUINIDINE

ROBINS PH

300MG

N12796 002

RANITIDINE HYDROCHLORIDE

SYRUP; ORAL

ZANTAC

EQ 15MG BASE/ML

N19675 001

EQ 15MG BASE/ML

EQ 15MG BASE/ML

N19675 001

DEC 30, 1988

† SEE SECTION 1.5 OF INTRODUCTION

RIENATRIPTAN BENZOATE

TABLET, ORALLY DISINTEGRATING; ORAL

MAVOLT
[REDACTED]

MAVOLT-NLT
MERCK

+

BQ 5MG BASE
BQ 10MG BASE

> DLT >
> DLT >

[REDACTED]
JUN 29, 1998
N20865 001
N20865 002
JUN 29, 1998

N20895 001
MAR 27, 1998
N20895 002
MAR 27, 1998
N20895 003
MAR 27, 1998

SACROSIDASE

SOLUTION; ORAL

SUCRAID
+ CRPHAN MEDCL

8,500 IU/ML

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

N20772 001
APR 09, 1998

N20773 001
OCT 29, 1998

SACQUINAVIR

CAPSULE; ORAL

FORTOVASE
[REDACTED]

+

[REDACTED]
200MG

[REDACTED]
N20828 001
NOV 07, 1997

[REDACTED]
N19766 001
DEC 23, 1991
N19766 004
DEC 23, 1991
N19766 005
JUL 10, 1998

SELEGILINE HYDROCHLORIDE

TABLET; ORAL

SELEGILINE HCL
ESI LEDEBIE

AP

5MG

[REDACTED]

STASON

AP

5MG

N74641 001
AUG 02, 1996
N74912 001
APR 30, 1998

SODIUM CHLORIDE

INJECTABLE; INJECTION

SODIUM CHLORIDE 0.45% IN PLASTIC CONTAINER

AP

450MG/100ML

450MG/100ML

N19635 001
MAR 09, 1988
N18184 001
MAR 09, 1988
N17464 001

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

SEVFLAMER HYDROCHLORIDE

CAPSULE; ORAL

RENAGEL
+ GELTEX

403MG

SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER

AP

900MG/100ML

SODIUM CHLORIDEINJECTABLE; INJECTIONSODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER

AP	B BRAUN	900MG/100ML	N19635 002
			MAR 09, 1988
AP	MCCAW	900MG/100ML	N17484 001
AP		900MG/100ML	N19635 002
			MAR 09, 1988

SODIUM CHLORIDE 3% IN PLASTIC CONTAINER

	B BRAUN	3GM/100ML	N19635 003
			MAR 09, 1988
	MCCAW	3GM/100ML	N19635 003
			MAR 09, 1988

SODIUM CHLORIDE 5% IN PLASTIC CONTAINER

	B BRAUN	5GM/100ML	N19635 004
			MAR 09, 1988
	MCCAW	5GM/100ML	N19635 004
			MAR 09, 1988

SODIUM LACTATEINJECTABLE; INJECTIONSODIUM LACTATE 0.167 MOLAR IN PLASTIC CONTAINER

	B BRAUN	1.87GM/100ML	N18186 001
			N18186 001
	MCCAW	1.87GM/100ML	N18186 001

SODIUM LACTATE 1/6 MOLAR IN PLASTIC CONTAINER

AP	B BRAUN	1.87GM/100ML	N20004 001
			APR 21, 1992
AP	MCCAW	1.87GM/100ML	N20004 001
			APR 21, 1992

SODIUM POLYSTYRENE SULFONATEPOWDER; ORAL, RECTALKAYEALATE

AA	SAROF	453.6GM/BOT	N11287 001
AA		453.6GM/BOT	N11287 001

KIONEX

AA	PADDOCK	454GM/BOT	N40029 001
			FEB 06, 1998

SONATROPIN, BIOSYNTHETICINJECTABLE; INJECTION

GENOTROPIN
+ PHARMACIA AND UPJOHN 13.8MG/VIAL

N20280 007
OCT 23, 1996

SOTALOL HYDROCHLORIDETABLET; ORAL

BETAPACE
+ BERLEX LABS 120MG

120MG

160MG

240MG

120MG

160MG

240MG

N19865 005
APR 20, 1994
N19865 002
OCT 30, 1992
N19865 003
OCT 30, 1992

SOYBEAN OILINJECTABLE; INJECTION

INTRALIPID 30%
AP + PHARMACIA AND UPJOHN 30%

N19942 001
DEC 30, 1993

LIPOSYN III 30%
AP + ABBOTT 30%

N20181 001
JAN 13, 1998

NUTRILIPID 10%
AP + B BRAUN 10%

N19531 001
MAY 28, 1993

AP + MCCAW 10%

N19531 001
MAY 28, 1993

NUTRILIPID 20%
AP + B BRAUN 20%

N19531 002
MAY 28, 1993

AP + MCCAW 20%

N19531 002
MAY 28, 1993

SPIRIFLOXACIN

TABLET; ORAL

200MG

WYLAN

200MG

20077 001
DEC 19, 1995

EQ 0.05MG BASE/ML

M74406 001
DEC 15, 1995

STREPTOMYCIN SULFATE

INJECTABLE; INJECTION

STREPTOMYCIN SULFATE

AP + PFIZER

EQ 1GM BASE/VIAL

260076 001

260111 001

264210 001

JUN 30, 1998

104

M80025 001

AP + PHARMA TEK

EQ 1GM BASE/VIAL

SULFAMETHOZOLE; TRIMETHOPRIM

INJECTABLE; INJECTION

10MG/ML; 10MG/ML

M18452 001

SUCCINYLCHOLINE CHLORIDE

INJECTABLE; INJECTION

SUCCINYLCHOLINE CHLORIDE

AP +

20MG/ML

M08847 001

20MG/ML; 40MG/5ML

M17598 001

SUCRALFATE

TABLET; ORAL

SUCRALFATE

AP + RATIOPHARM

1GM

M74415 001

JUN 08, 1998

20MG/5ML; 40MG/5ML

M17598 001

SUFENTANIL CITRATE

INJECTABLE; INJECTION

SUFENTANIL CITRATE

AP + AKORN

EQ 0.05MG BASE/ML

M19050 001

MAY 04, 1994

200MG/5ML; 40MG/5ML

M18812 001

AP +

EQ 0.05MG BASE/ML

M19050 001

MAY 04, 1994

200MG/5ML; 40MG/5ML

M18812 002

SUFENTANIL CITRATE

TABLET; ORAL

SUFENTANIL CITRATE

AP +

EQ 0.05MG BASE/ML

M19050 001

MAY 04, 1994

200MG/5ML; 40MG/5ML

M18812 001

AP +

EQ 0.05MG BASE/ML

M19050 001

MAY 04, 1994

200MG/5ML; 40MG/5ML

M18812 001

AP +

EQ 0.05MG BASE/ML

M19050 001

MAY 04, 1994

200MG/5ML; 40MG/5ML

M18812 001

TECHNETIUM TC-99M APCITIDE

INJECTABLE; INJECTION
ACUTECT
DIATIDE

N/A

N20887 001
SEP 14, 1998

TECHNETIUM TC-99M DISOFENIN KIT

INJECTABLE; INJECTION
HEPATOLITE

N/A

DUPONT PHARMS

N/A

N18467 001
MAR 16, 1982

TECHNETIUM TC-99M GLUCEPTATE KIT

INJECTABLE; INJECTION

TECHNISCAN
GLUCEPTATE

N/A

TECHNISCAN
GLUCEPTATE

N/A

N17907 001
JAN 27, 1982

N18272 001
JAN 27, 1982

TECHNETIUM TC-99M LIDOFENIN KIT

INJECTABLE; INJECTION

TECHNISCAN HIDA
DRAXIMAGE

N/A

N/A

N/A

N18489 001
OCT 31, 1986

TECHNETIUM TC-99M MEDRONATE KIT

INJECTABLE; INJECTION

TECHNISCAN MDP KIT

DRAXIMAGE

N/A

DRAXIMAGE

N/A

N18035 001
N18035 001

TECHNETIUM TC-99M PENTETATE KIT

INJECTABLE; INJECTION

DRAXIMAGE

N/A

N/A

N/A

N/A

N18511 001
DEC 29, 1989

TECHNETIUM TC-99M SULFUR COLLOID KIT

SOLUTION; INJECTION, ORAL

AM-SULFUR COLLOID

N/A

CIS

N/A

N/A

BRAND

N/A

N/A

N17858 001
N18922 001
N18923 001

TERAZOSIN HYDROCHLORIDE

CAPSULE; ORAL

HYTRIN

N/A

ABBOTT

EQ 1MG BASE

N/A

+

EQ 2MG BASE

N/A

EQ 5MG BASE

N/A

EQ 10MG BASE

N20347 001
DEC 14, 1994
N20347 002
DEC 14, 1994
N20347 003
DEC 14, 1994
N20347 004
DEC 14, 1994

N/A

TERAZOSIN HCL
GENEVA PHARMS

EQ 1MG BASE

N/A

EQ 2MG BASE

N/A

EQ 5MG BASE

N/A

EQ 10MG BASE

N74823 001
MAR 30, 1998
N74823 002
MAR 30, 1998
N74823 003
MAR 30, 1998
N74823 004
MAR 30, 1998

TERBINAFINE

GEL; TOPICAL

LAMISIL

+ NOVARTIS

14

N20846 001
APR 29, 1998

RX DRUG PRODUCT LIST / CUMULATIVE SUPPLEMENT NUMBER 10 / JAN'98 - OCT'98

TESTOSTERONE

FILM, EXTENDED RELEASE; TRANSDERMAL

ANDRODERM
NDC [REDACTED]
[REDACTED]

2.5MG/24HR

NDC [REDACTED]
N20489 001
SEP 29, 1995TETRACYCLINE HYDROCHLORIDECAPSULE; ORAL
ACROMICIN V
ESTI LEDEBKE250MG
500MG
[REDACTED]N50278 003
N50278 001
N50278 002
N50278 004TABLET; ORAL
SUMYCIN50MG
100MG
100MGN61147 003
N61147 002
N61147 001THALIDOMIDECAPSULE; ORAL
THALONID
+ CELGENE

50MG

N20785 001
JUL 16, 1998THEOPHYLLINECAPSULE; ORAL
[REDACTED][REDACTED]
[REDACTED]
100MG
200MGN85545 001
JUL 31, 1984
N83921 001
JUL 31, 1984THEOPHYLLINE

CAPSULE, EXTENDED RELEASE; ORAL

NDC [REDACTED]
[REDACTED]125MG
250MGN86826 001
JAN 29, 1985
N86826 002
JAN 29, 1985

TABLET; ORAL

QUIBRON-T
+ MONARCH PHARMS300MG
[REDACTED]N88656 001
AUG 22, 1985
N88656 002
AUG 22, 1985

TABLET, EXTENDED RELEASE; ORAL

NDC [REDACTED]
QUIBRON-T/SR
[REDACTED]

300MG

N87563 001
JUN 21, 1983
N87563 002
JUN 21, 1983THIAMYAL SODIUM[REDACTED]
[REDACTED]1GM/VIAL
5GM/VIAL
10GM/VIALN07600 003
N07600 005
N07600 009THIORIDAZINE HYDROCHLORIDE

CONCENTRATE; ORAL

NDC [REDACTED]
THIORIDAZINE HCL
PHARM ASSOC

100MG/ML

N40213 001
MAY 29, 1998

TABLET; ORAL

NDC [REDACTED]
THIORIDAZINE HCL
[REDACTED]N40213 001
MAY 29, 1998

TORSEMIDEINJECTABLE; INJECTION
DEMADEX* SCHERING-KLEBER
+ ROCHE 10MG/MLN20137 001
AUG 23, 1993
N20137 002
AUG 23, 1993TABLET; ORAL
DEMADEXSCHERING-KLEBER
5MG
10MG
20MG
40MG
ROCHE 5MG
10MG
20MG
100MGN20136 001
AUG 23, 1993
N20136 002
AUG 23, 1993
N20136 003
AUG 23, 1993
N20136 004
AUG 23, 1993TRETINOINCREAM; TOPICAL
AVITA* SCHERING-KLEBER
AB 0.025%N20404 001
JAN 14, 1997
N20404 003
JAN 14, 1997GEL; TOPICAL
AVITABT PENEDERM 0.025%
* SCHERING-KLEBER 0.025%N20400 001
JAN 29, 1998
N20400 001
JAN 29, 1998

RETIN-A

* SCHERING-KLEBER
BX + JOHNSON AND JOHNSON 0.025%N17579 001
N17579 002

SOLUTION; TOPICAL

RETIN-A

* SCHERING-KLEBER

N16921 001

TRETINOIN

SOLUTION; TOPICAL

* SCHERING-KLEBER
AT + JOHNSON AND JOHNSON 0.05%
* SCHERING-KLEBER
AT COPLEY PHARM 0.05%N16921 001
N74873 001
JUN 19, 1998TRIAMCINOLONE ACETONIDE

CREAM; TOPICAL

* ALPHARMA

* ALPHARMA 0.025%
AT ROC 0.025%N87797 001
JUN 07, 1982
N87797 001
JUN 07, 1982TRIAMCINOLONE DIACETATE

SYRUP; ORAL

KENACORT

* SCHERING-KLEBER

EQ 4MG BASE/5ML

EQ 4MG BASE/5ML
EQ 4MG BASE/5MLN12515 001
N12515 001TRICHLORMETHIAZIDE

TABLET; ORAL

TRICHLORMETHIAZIDE

* DANNIOTT PHARMA 2MG
* DANNIOTT PHARMA 4MG
* DANNIOTT PHARMA 2MG
* DANNIOTT PHARMA 4MGN83847 001
N83847 001
N83847 001
N83855 001TRIFLUOPERAZINE HYDROCHLORIDE

TABLET; ORAL

* ZENITH GOLDLINE

AB ZENITH GOLDLINE EQ 1MG BASE
AB ZENITH GOLDLINE EQ 2MG BASE
AB ZENITH GOLDLINE EQ 1MG BASEN87612 001
NOV 19, 1982
N87613 001
NOV 19, 1982
N87613 001
NOV 19, 1982

APPENDIX

OPTICALLY ACTIVE

VIET-1

PARKEDALZ



001 006030

VINCISTIB SULFATE

INJECTABLE; INJECTION



TVIA/ENT

2025/02/21

SMO/VIAL

APR 11, 1988
N71560 001
APR 11, 1988
N71561 001
APR 11, 1988

WATER FOR INJECTION, STERILE

V/M ; CREDIT

STERILIZING WATER FOR INJECTION IN PLASTIC CONTAINER

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100-44363-001

723 29, 1988

ACETAMINOPHEN; ASPIRIN; CAFFEINE

TABLET; ORAL
EXCEDRIN (MIGRAINE)
+ BRISTOL MYERS 250MG;250MG;65MG N20802 001
JAN 14, 1998

ALUMINUM HYDROXIDE; MAGNESIUM TRISILICATE

TABLET, CHEWABLE; ORAL
GAVISCON 80MG;20MG N18685 001
DEC 09, 1983
+ 160MG;40MG N18685 002
DEC 09, 1983
GAVISCON-2 80MG;20MG N18685 002
DEC 09, 1983

CHLORHEXIDINE GLUCONATE

SOLUTION; TOPICAL
CHG SCRUB
ECOLAB 4% N19258 002
JUL 22, 1986
HUNTINGTON LABS 4% N19258 002
JUL 22, 1986
CIDA-STAT
ECOLAB 2% N19258 001
JUL 22, 1986
HUNTINGTON LABS 2% N19258 001
JUL 22, 1986

CIMETIDINE

TABLET; ORAL
CIMETIDINE
LEK PHARM 100MG N75122 001
JUN 19, 1998
200MG N75122 002
JUN 19, 1998
NOVOPHARM 200MG N74961 001
JUN 19, 1998

CIMETIDINE

TABLET; ORAL
CIMETIDINE
PERRIGO 100MG N74972 001
JUN 19, 1998
200MG N75285 001
OCT 29, 1998
PHARM FORM 200MG N74963 001
JUN 19, 1998
TORPHARM 100MG N74948 001
JUN 19, 1998

CLOTRIMAZOLE

TABLET; VAGINAL
GYNIX
COPLEY PHARM 100MG N73249 001
FEB 13, 1998

FAMOTIDINE

TABLET, CHEWABLE; ORAL
PEPCID AC
+ MERCK 10MG N20801 001
SEP 24, 1998

IBUPROFEN

SUSPENSION; ORAL
CHILDREN'S ADVIL-FLAVORED
WHITEHALL ROBINS 100MG/5ML N20589 001
NOV 07, 1997
100MG/5ML N20589 002
NOV 07, 1997

SUSPENSION/DROPS; ORAL
PEDIATRIC ADVIL
+ WHITEHALL ROBINS 100MG/2.5ML N20812 001
JAN 30, 1998

TABLET; ORAL
IBUPRIN 200MG N71773 001
JUN 19, 1998

IBUPROFEN

TABLET; ORAL

IBUPRAIN
© SINDAK LABS NJ

200MG

N71773 001
JUL 16, 1987

IBUPROFEN

IBUPROFEN

200MG

N71905 001
MAR 08, 1988

200MG

N73019 001
MAR 30, 1994

200MG

N74931 001
JUL 20, 1998

200MG

N74782 001
JUL 06, 1998

NOVOPHARM

200MG

PHARM FORM

200MG

IBUPROFEN

IBUPROFEN

200MG

200MG

200MG

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200MG

200MG

INSULIN BIOSYNTHETIC HUMAN

INJECTABLE; INJECTION

HUMULIN R PEN

100 UNITS/ML

N18780 005
AUG 06, 1998

+ LILLY

INSULIN BIOSYNTHETIC HUMAN; INSULIN SUSP ISOPHANE BIOSYNTHETIC HUMAN

INJECTABLE; INJECTION

HUMULIN 70/30 PEN

30 UNITS/ML; 70 UNITS/ML

N19717 002
AUG 06, 1998

+ LILLY

NICOTIAZOLE NITRATE

CREAM; VAGINAL

MONISTAT 3

+ ADVANCED CARE PRODS 4t

N20827 001
MAR 30, 1998**MINOXIDIL**

SOLUTION; TOPICAL

MINOXIDIL (FOR MEN)

NOVEX

2t

N74924 001
APR 29, 1998

+ NOVEX

+ NOVEX

+ NOVEX

+ NOVEX

+ NOVEX

+ NOVEX

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NAPROXEN SODIUM

TABLET; ORAL

NAPROXEN SODIUM

PAR PHARM

EQ 200MG BASE

N75169 001
JUL 28, 1998

NICOTINE

FILE, EXTENDED RELEASE; TRANSDERMAL

NICOTROL

WATSON

+ PHARMACIA AND UPJOHN 15MG/16HR

XXXXXXXXXX

N20536 001

JUL 03, 1996

RANITIDINE HYDROCHLORIDE

TABLET, EFFERVESCENT; ORAL

ZANTAC 75

+ GLAXO WELLCOME

EQ 75MG BASE

N20745 001

FEB 26, 1998

**DRUG PRODUCTS WITH APPROVAL UNDER SECTION 505 OF THE ACT ADMINISTERED BY THE
CENTER FOR BIOLOGICS EVALUATION AND RESEARCH LIST**

CUMULATIVE SUPPLEMENT NUMBER 10 OCT '98

NO OCTOBER 1998 APPROVALS

This data is provided to the Division of Data Management and Services from the Office of Orphan Products Development and it is not edited prior to publication.

**Orphan Product Designations and Approvals List
January 1998 through October 1998**

Name Generic Name TN=Trade Name	Indication Designated	Sponsor & Address DD= Date Designated MA=Marketing Approval
1,5-(Butylimino) Treatment of Fabry's disease. -1,5 dideoxy,D-glucit ol TN=		Oxford GlycoSciences 10, The Quadrant Abington Science Park, Abington Oxfordshire OX14 3YS UK, DD=05/12/1998
1,5-(Butylimino) Treatment of Gaucher disease. -1,5 dideoxy,D-glucit ol TN=		Oxford GlycoSciences 10, The Quadrant Abington Science Park, Abington Oxfordshire OX14 3YS UK, DD=05/29/1998
3-(3,5-dimethyl- Treatment of Kaposi's sarcoma. 1H-2ylmethylene) -1,3-dihydro-ind ol-2-one TN=		Sugen, Inc. 230 East Grand Ave. South San Francisco, CA 94080 DD=09/11/1998
Aldesleukin Treatment of metastatic TN= Proleukin melanoma.		Chiron Corporation 4560 Horton Street Emeryville, CA 94608 DD=09/10/1996 MA=01/09/1998
Aldesleukin Treatment of acute myelogenous TN= Proleukin leukemia.		Chiron Corporation 4560 Horton St. Emeryville, CA 94608 DD=07/31/1998

Orphan Product Designations and Approvals List

January 1998 through October 1998

Name Generic Name TN=Trade Name	Indication Designated	Sponsor & Address DD= Date Designated MA=Marketing Approval
Aliperetinate TN= Panretin	For the topical treatment of cutaneous lesions in patients with AIDS-related Kaposi's sarcoma.	Ligand Pharmaceuticals Inc. 10275 Science Center Drive San Diego, CA 92121 DD=03/24/1998
Alpha-galactosidase A TN=	Long-term enzyme replacement therapy for the treatment of Fabry disease.	Transkaryotic Therapies Inc. 195 Albany St. Cambridge, MA 02139 DD=06/22/1998
Amifostine TN= Ethyol	Reduction of the incidence and severity of radiation-induced xerostomia.	U.S. Bioscience, Inc. One Tower Bridge 100 Front Street, Suite 400 Conshohocken, PA 19428 DD=05/12/1998
Arsenic trioxide TN=	Treatment of acute promyelocytic leukemia.	Polarx, Inc. 787 7th Ave., 48th Floor New York, NY 10019 DD=03/03/1998
Basiliximab TN= Simulect	Prophylaxis of solid organ rejection.	Novartis Pharmaceuticals Corporation 59 Route 10 East Hanover, NJ 07936 DD=12/12/1997 MA=05/12/1998

Orphan Product Designations and Approvals List

January 1998 through October 1998

Name Generic Name TN=Trade Name	Indication Designated	Sponsor & Address DD= Date Designated MA=Marketing Approval
Beclomethasone dipropionate TN=	For oral administration in the treatment of intestinal graft-versus-host disease.	George B. McDonald, M.D. Fred Hutchinson Cancer Research Center 1100 Fairview Avenue North (SC-113); PO Box 19024 Seattle, WA 98109 DD=03/27/1998
Benzydamine hydrochloride TN= Tantom	Prophylactic treatment of oral mucositis resulting from radiation therapy for head and neck cancer.	Angelini Pharmaceuticals, Inc. 70 Grand Avenue River Edge, NJ 07661 DD=05/18/1998
Bindarit TN=	Treatment of lupus nephritis.	Angelini Pharmaceuticals, Inc. 70 Grand Avenue River Edge, NJ 07661 DD=02/03/1998
Botulinum toxin type A TN= Dysport	Treatment of spasmodic torticollis (cervical dystonia).	Ipsen Limited 1 Bath Road Maidenhead, Berkshire U.K. SL6 4UH DD=08/12/1998
Carbamylglutamic acid TN=	Treatment of N-acetylglutamate synthetase deficiency.	Orphan Europe Immeuble "Le Guillaumet" 60 avenue du President Wilson 92046 Paris France, DD=01/20/1998
Corticotropin-re leasing factor, human TN= Xerecept	Treatment of peritumoral brain edema.	Neurobiological Technologies, Inc. 1387 Marina Way South Richmond, CA 94804 DD=04/06/1998

Orphan Product Designations and Approvals List January 1998 through October 1998

Name Generic Name TN=Trade Name	Indication Designated	Sponsor & Address DD= Date Designated MA=Marketing Approval
Dexamethasone TN=	For use in posterior segment drug delivery system in the treatment of idiopathic intermediate uveitis.	Oculex Pharmaceuticals 639 N. Pastoria Avenue Sunnyvale, CA 94086 DD=09/11/1998
Dimethylsulfoxide TN=	Treatment of palmar-plantar erythrodysesthesia syndrome.	Cancer Technologies, Inc. 7301 East 22nd Street Suite 10E Tucson, AZ 85710 DD=04/06/1998
Filgrastim TN= Neupogen	Reduction in the duration of neutropenia, fever, antibiotic use, and hospitalization, following induction and consolidation treatment for acute myeloid leukemia.	Amgen, Inc. 1840 DeHavilland Drive Thousand Oaks, CA 91320 DD=11/07/1996 MA=04/02/1998
Fructose-1,6-diphosphate TN=	Treatment of painful vaso-occlusive episodes associated with sickle cell disease.	Cypros Pharmaceutical Corporation 2714 Loker Avenue West Carlsbad, CA 92008 DD=05/29/1998
Humanized anti-human CD2 MAb TN= MEDI-507	For the induction of donor-specific immunologic unresponsiveness resulting in prophylaxis of organ rejection without the need for chronic immunosuppressive therapy, in patients receiving allogeneic renal transplants.	Biotransplant, Inc. Building 75, 3rd. Ave. Charlestown Navy Yard Charlestown, MA 02129 DD=09/17/1998
Hydroxyurea TN= Droxia	Treatment of patients with sickle cell anemia as shown by the presence of hemoglobin S.	Bristol-Myers Squibb Pharmaceutical Research Institute P.O. Box 4000 Princeton, NJ 08543 DD=10/01/1990 MA=02/25/1998

Orphan Product Designations and Approvals List

January 1998 through October 1998

Name Generic Name TN=Trade Name	Indication Designated	Sponsor & Address DD= Date Designated MA=Marketing Approval
Infliximab TN= Remicade	Treatment of moderately to severely active Crohn's disease for the reduction of the signs and symptoms, in patients who have an inadequate response to conventional therapy; and treatment of patients with fistulizing Crohn's disease for the reduction in the number of draining enterocutaneous fistula(s).	Centocor, Inc. 200 Great Valley Parkway Malvern, PA 19355 DD=11/14/1995 MA=08/24/1998
Interferon Beta-1a TN= Avonex	Treatment of juvenile rheumatoid arthritis.	Biogen, Inc. 14 Cambridge Center Cambridge, MA 02142 DD=10/14/1998
L-baclofen TN=	Treatment of trigeminal neuralgia.	Pharmascience, Inc. 8400 Darnley Road Montreal, Quebec Canada H4T 1M4, DD=01/06/1998
Lamotrigine TN= Lamictal	Treatment of Lennox-Gastaut syndrome.	Glaxo Wellcome Research and Development 5 Moore Drive P.O. Box 13398 Research Triangle Park, NC 27709 DD=08/23/1995 MA=08/24/1998
Lepirudin TN= Refluden	Treatment of heparin-associated thrombocytopenia type II.	Hoechst Marion Roussel Frankfurt am Main Germany DD=02/13/1997 MA=03/06/1998

Orphan Product Designations and Approvals List January 1998 through October 1998

Name Generic Name TN=Trade Name	Indication Designated	Sponsor & Address DD= Date Designated MA=Marketing Approval
Liposomal Cyclosporin A TN= Cyclospire	For aerosolized administration in the prevention and treatment of lung allograft rejection and pulmonary rejection events associated with bone marrow transplantation.	Vernon Knight, M.D. Baylor College of Medicine, Dept. of Molecular Physiology One Baylor Plaza Houston, TX 77030 DD=04/30/1998
Liposomal N-Acetylglucosmi nyl-N-Acetylmura mly-L-Ala-D-isoG ln-L-Ala -gylcerolidpalmi toyl TN= ImmTher	Treatment of osteosarcoma.	Endorex Corp. 900 North Shore Drive Lake Bluff, IL 60044 DD=06/10/1998
Liposomal N-Acetylglucosmi nyl-N-Acetylmura mly-L-Ala-D-isoG ln-L-Ala -gylcerolidpalmi toyl TN= ImmTher	Treatment of Ewing's sarcoma.	Endorex Corp. 900 North Shore Drive Lake Bluff, IL 60044 DD=06/10/1998
MN14 monoclonal antibody to carcinoembryonic antigen TN= CEA-CIDE	For the treatment of small cell lung cancer.	Immunomedics, Inc. 300 American Rd. Morris Plains, NJ 07950 DD=09/18/1998
Mafenide acetate solution TN= Sulfamylon solution	For use as an adjunctive topical antimicrobial agent to control bacterial infection when used under moist dressings over meshed autografts on excised burn wounds.	Mylan Laboratories, Inc. 781 Chestnut Ridge Road P.O. Box 4310 Morgantown, WV 26504 DD=07/18/1990 MA=06/05/1998

Orphan Product Designations and Approvals List January 1998 through October 1998

Name Generic Name TN=Trade Name	Indication Designated	Sponsor & Address DD= Date Designated MA=Marketing Approval
Mecamylamine TN= Inversine	Treatment of Tourette's syndrome.	Layton Bioscience, Inc. 105 Reservoir Rd. Atherton, CA 94027 DD=10/14/1998
Methoxsalen TN= Uvadex	For use in conjunction with the UVAR photopheresis system to treat graft versus host disease.	Therakos, Inc. 437 Creamery Way Exton, PA 19331 DD=10/14/1998
Octreotide TN= Sandostatin LAR	Treatment of acromegaly.	Novartis Pharmaceuticals Corporation 59 Route 10 East Hanover, NJ 07936 DD=08/24/1998
Octreotide TN= Sandostatin LAR	Treatment of severe diarrhea and flushing associated with malignant carcinoid tumors.	Novartis Pharmaceuticals Corporation 59 Route 10 East Hanover, NJ 07936 DD=08/24/1998
Octreotide TN= Sandostatin LAR	Treatment of diarrhea associated with vasoactive intestinal peptide tumors (VIPoma).	Novartis Pharmaceuticals Corporation 59 Route 10 East Hanover, NJ 07936 DD=08/24/1998
P1, P4-Di(uridine 5'-tetraphosphat e), tetrasodium salt TN=	Treatment of cystic fibrosis.	Inspire Pharmaceuticals, Inc. 4222 Emperor Blvd., Suite 470 Durham, NC 27703 DD=10/27/1998

Orphan Product Designations and Approvals List January 1998 through October 1998

Name Generic Name TN=Trade Name	Indication Designated	Sponsor & Address DD= Date Designated MA=Marketing Approval
PEGASYS TN=	Treatment of renal cell carcinoma.	Hoffman-La Roche Inc. 340 Kingsland St. Nutley, NJ 07110 DD=07/13/1998
Pentostatin TN=	Treatment of cutaneous T-cell lymphoma.	SuperGen, Inc. Two Annbel Lane, Suite 220 San Ramon, CA 94583 DD=03/27/1998
Phenylacetate TN=	For use as an adjunct to surgery, radiation therapy and chemotherapy for the treatment of patients with primary or recurrent malignant glioma.	Targon Corporation 307 College Road East Princeton, NJ 08540 DD=03/06/1998
Pilocarpine HCl TN= Salagen	Treatment of xerostomia and keratoconjunctivitis sicca in Sjogren's syndrome patients.	MGI Pharma, Inc. 9900 Bren Road East Suite 300E Minneapolis, MN 55343 DD=02/28/1992 MA=02/11/1998
Prostaglandin E1 enol ester (AS-013) TN=	Treatment of Fontaine Stage IV chronic critical limb ischemia.	Alpha Therapeutic Corp. 5555 Valley Blvd. Los Angeles, CA 90032 DD=06/12/1998
Radiolabeled monoclonal antibody to CD22 antigen on B-cells TN= LymphoCIDE	Treatment of non-Hodgkin's lymphoma.	Immunomedics, Inc. 300 American Rd. Morris Plains, NJ 07950 DD=07/13/1998
Recombinant bactericidal/permeability-increasing protein TN= Neuprex	Treatment of severe meningococcal disease.	Xoma Corporation 2910 Seventh Street Berkeley, CA 94710 DD=06/22/1998

Orphan Product Designations and Approvals List

January 1998 through October 1998

Name Generic Name TN=Trade Name	Indication Designated	Sponsor & Address DD= Date Designated MA=Marketing Approval
Recombinant human Clara Cell 10kDa protein TN=	Prevention of neonatal bronchopulmonary dysplasia in premature neonates with respiratory distress syndrome.	Claragen, Inc. 335 Paint Branch Drive College Park, MD 20742 DD=07/13/1998
Recombinant human tumor necrosis factor receptor fusion protein TN= Enbrel	Treatment of juvenile rheumatoid arthritis.	Immunex Corporation 51 University St. Seattle, WA 98101 DD=10/27/1998
Recombinant humanized MAb 5c8 TN=	Prevention and treatment of Factor VIII/Factor IX inhibitors in patients with hemophilia A or B.	Biogen, Inc. 14 Cambridge Center Cambridge, MA 02142 DD=10/14/1998
Recombinant humanized monoclonal antibody 5c8 TN=	Treatment of immune thrombocytopenic purpura.	Biogen, Inc. 14 Cambridge Center Cambridge, MA 02142 DD=02/03/1998
Recombinant humanized monoclonal antibody 5c8 TN=	Treatment of systemic lupus erythematosus.	Biogen, Inc. 14 Cambridge Center Cambridge, MA 02142 DD=02/18/1998
Rifapentine TN= Priftin	Treatment of pulmonary tuberculosis.	Hoechst Marion Roussel P.O. Box 9627 Mail Station: H3-M2516 Kansas City, MO 64134 DD=06/09/1995 MA=06/22/1998
Rifaximin TN= Normix	Treatment of hepatic encephalopathy.	Salix Pharmaceuticals, Inc. 3600 W. Bayshore Road Palo Alto, CA 94303 DD=02/10/1998

Orphan Product Designations and Approvals List

January 1998 through October 1998

Name Generic Name TN=Trade Name	Indication Designated	Sponsor & Address DD= Date Designated MA=Marketing Approval
S-adenosylmethio nine TN=	Treatment of AIDS-myelopathy.	Di Rocco, Alessandro M.D. Beth Israel Medical Center, Phillips Ambulatory Care Center Philips Building, Suite 2Q; 10 Union Square East New York, NY 10003 DD=04/30/1998
Sacrosidase TN= Sucraid	Treatment of congenital sucrase-isomaltase deficiency.	Orphan Medical, Inc. 13911 Ridgedale Drive Suite 475 Minnetonka, MN 55305 DD=12/10/1993 MA=04/09/1998
Sodium phenylbutyrate TN=	For use as an adjunct to surgery, radiation therapy and chemotherapy for the treatment of patients with primary or recurrent malignant glioma.	Targon Corporation 307 College Road East Princeton, NJ 08540 DD=04/24/1998
TAK-603 TN=	Treatment of Crohn's disease.	TAP Holdings Inc. 2355 Waukegan Road Deerfield, IL 60015 DD=05/13/1998
Tacrolimus TN= Prograf	Prophylaxis of graft-versus-host-disease.	Fujisawa USA, Inc. 3 Parkway North Center Deerfield, IL 60015 DD=04/06/1998
Temozolomide TN= Temodal	Treatment of recurrent malignant glioma.	Shering-Plough Research Institute 2000 Galloping Hill Rd. Kenilworth, NJ 07033 DD=10/05/1998

Orphan Product Designations and Approvals List

January 1998 through October 1998

Name Generic Name TN=Trade Name	Indication Designated	Sponsor & Address DD= Date Designated MA=Marketing Approval
Temozolomide TN= Temodal	Treatment of advanced metastatic melanoma.	Schering-Plough Research Institute 2000 Galloping Hill Rd. Kenilworth, NJ 07033 DD=10/14/1998
Tetrabenazine TN=	Treatment for moderate/severe tardive dyskinesia.	Lifehealth Limited Richmond House, Old Brewery Court, Sandyford Road Newcastle upon Tyne NE2 1XG England DD=05/12/1998
Thalidomide TN= Thalomid	Treatment of erythema nodosum leprosum.	Celgene Corporation P.O. Box 4914 7 Powder Horn Drive Warren, NJ 07059 DD=07/26/1995 MA=07/16/1998
Thalidomide TN=	Treatment of primary brain malignancies.	EntreMed, Inc. 9610 Medical Center Drive, Suite 200 Rockville, MD 20850 DD=02/27/1998
Thalidomide TN=	Treatment of Kaposi's sarcoma.	EntreMed, Inc. 9610 Medical Center Dr., Suite 200 Rockville, MD 20850 DD=07/29/1998
Thalidomide TN= Thalomid	Treatment of multiple myeloma.	Celgene Corporation 7 Powder Horn Dr. Warren, NJ 07059 DD=10/14/1998

Orphan Product Designations and Approvals List January 1998 through October 1998

Name Generic Name TN=Trade Name	Indication Designated	Sponsor & Address DD= Date Designated MA=Marketing Approval
Thymalfasin TN= Zadaxin	Treatment of DiGeorge anomaly with immune defects.	SciClone Pharmaceuticals, Inc. 901 Mariner's Island Blvd. San Mateo, CA 94404 DD=01/08/1998
Tiapride TN=	Treatment of Tourette's syndrome.	Synthelabo Research, Inc. 400 Plaza Drive Secaucus, NJ 07094 DD=04/21/1998
Transgenic human alpha 1 antitrypsin TN=	Treatment of cystic fibrosis.	PPL Therapeutics (Scotland) Limited Roslin, Edinburgh EH25 9PP Scotland U.K., DD=03/06/1998
Valrubicin TN= Valstar	Treatment of carcinoma in situ of the urinary bladder.	Anthra Pharmaceuticals, Inc. 103 Carnegie Center, Suite 102 Princeton, NJ 08540 DD=05/23/1994 MA=09/25/1998

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

NO OCTOBER 1998 ADDITIONS

PATENT AND EXCLUSIVITY TERMS PATENT AND EXCLUSIVITY TERMS

DUE TO SPACE LIMITATIONS IN THE PATENT AND EXCLUSIVITY COLUMNS, ABBREVIATIONS AND REFERENCES HAVE BEEN DEVELOPED. PLEASE REFER BACK TO THE APPROVED DRUG PRODUCTS WITH THERAPEUTIC EQUIVALENCE EVALUATIONS, 18TH EDITION FOR A FULL LISTING OF PATENT AND EXCLUSIVITY TERMS (ABBREVIATIONS, NEW DOSING SCHEDULE, NEW INDICATIONS AND PATENT USE CODES). ONLY NEW CODES WILL BE ADDED TO THE CUMULATIVE SUPPLEMENT.

ABBREVIATIONS

PED PEDIATRIC EXCLUSIVITY

REFERENCES NEW DOSING SCHEDULE

- D-38 CONTINUOUS INFUSION AS AN ALTERNATE METHOD OF ADMINISTRATION**
- D-39 CHANGE IN TIME TO TAKE THE DRUG PRIOR TO A MEAL TO PREVENT MEAL-INDUCED HEARTBURN SYMPTOMS FROM "...1/2-1 HOUR BEFORE EATING..." TO "...RIGHT BEFORE EATING OR UP TO 60 MIN BEFORE CONSUMING..."**
- D-40 ONCE-A-DAY DOSING REGIMEN**
- D-41 DRUG MAY BE DOSED RIGHT BEFORE A MEAL OR ANY TIME UP TO 30 MIN BEFORE EATING OR DRINKING FOOD AND BEVERAGES THAT WOULD BE EXPECTED TO CAUSE SYMPTOMS**
- D-42 TEN DAY DOSING REGIMEN FOR TRIPLE THERAPY. PREVACID IN COMBINATION WITH CLARITHROMYCIN AND AMOXICILIN, FOR THE ERADICATION OF H. PYLORI IN PATIENTS WITH DUODENAL ULCER DISEASE**
- D-43 INITIATION OF TREATMENT WITH 900MG/DAY BY DELETION OF THE REQUIREMENT TO TITRATE TO 900MG/DAY OVER A 3-DAY PERIOD**
- D-44 IN A CLINICAL TRIAL, FEWER DISCONTINUATIONS DUE TO ADVERSE EVENTS, ESPECIALLY DIZZINESS AND VERTIGO, WERE OBSERVED WHEN TITRATING THE DOSE IN INCREMENTS OF 50MG/DAY EVERY THREE DAYS UNTIL AN EFFECTIVE DOSE (NOT EXCEEDING 400MG/DAY) WAS REACHED**
- D-45 ONCE DAILY DOSING FOR MAINTENANCE ONLY**
- D-46 NEW DOSING REGIMEN OF 80MG DAILY**
- D-47 TAKE DRUG "15 MINUTES TO 1 HOUR" PRIOR TO A MEAL TO PREVENT MEAL-INDUCED HEARTBURN**
- D-48 ADMINISTRATION OF CISATRICURIUM, A NEUROMUSCULAR BLOCKING AGENT AT DOSES OF 3 AND 4X THE ED95 OF CISATRICURIUM FOLLOWING INDUCTION WITH THIOPENTAL**
- D-49 PEDIATRIC DOSING GUIDELINES**

NEW INDICATION

- I-212 TREATMENT OF SYMPTOMS OF DRY MOUTH IN PATIENTS WITH SJOGREN'S SYNDROME**
- I-213 TEMPORARY RELIEF OF PAIN AND PHOTOPHOBIA IN PATIENTS UNDERGOING CORNEAL REFRACTIVE SURGERY**
- I-214 TREATMENT OF OSTEOPOROSIS**
- I-215 PRE-PROCEDURAL APPLICATION TO ADULT MALE GENITAL SKIN PRIOR TO SITE-SPECIFIC SUBCUTANEOUS INFILTRATION WITH LIDOCAINE FOR THE REMOVAL OF GENITAL WARTS**
- I-216 FOR THE LONG-TERM TWICE-DAILY (MORNING AND EVENING) ADMINISTRATION IN THE MAINTENANCE TREATMENT OF BRONCHOSPASM ASSOCIATED WITH COPD, INCLUDING CHRONIC BRONCHITIS AND EMPHYSEMA**
- I-217 PREVENTION (DURING AND FOLLOWING HOSPITALIZATION) OF DEEP VEIN THROMBOSIS, WHICH MAY LEAD TO PULMONARY EMBOLISM, IN PATIENTS UNDERGOING HIP REPLACEMENT SURGERY**
- I-218 USE OF LIPITOR AS AN ADJUNCTIVE THERAPY TO DIET FOR THE TREATMENT OF PATIENTS WITH ELEVATED SERUM TRIGLYCERIDE LEVELS (FREDERICKSON TYPE IV)**

PATENT AND EXCLUSIVITY TERMS

NEW INDICATION

- I-219 USE OF LIPITOR BY PATIENTS WITH PRIMARY DYSBETALIPOPROTEINEMIA (FREDERICKSON TYPE III) WHO DO NOT RESPOND ADEQUATELY TO DIET
- I-220 TREATMENT OF EPISODIC HEARTBURN, ACID INDIGESTION AND SOUR STOMACH
- I-221 TREATMENT OF BENIGN PROSTATIC HYPERPLASIA (BPH) IN MEN WITH AN ENLARGED PROSTATE TO IMPROVE SYMPTOMS, REDUCE THE RISK OF ACUTE URINARY RETENTION AND REDUCE THE RISK OF THE NEED OF SURGERY
- I-222 PREVENTION OF ISCHEMIC COMPLICATIONS OF UNSTABLE ANGINA AND NON-Q-WAVE MYOCARDIAL INFARCTION, WHEN CONCURRENTLY ADMINISTERED WITH ASPIRIN
- I-223 USE IN THE SYMPTOMATIC RELIEF OF RHINORRHEA ASSOCIATED WITH ALLERGIC AND NONALLERGIC PERENNIAL RHINITIS IN CHILDREN AGE 6-11 YEARS
- I-224 FOR THE USE IN PEDIATRIC PATIENTS 4 TO 11 YEARS OF AGE FOR THE MANAGEMENT OF THE NASAL SYMPTOMS OF SEASONAL AND PERENNIAL ALLERGIC RHINITIS
- I-225 USE IN PATIENTS WITH PREVIOUS MI AND NORMAL CHOLESTEROL LEVELS, TO REDUCE RISK OF RECURRENT MI, MYOCARDIAL REVASCULARIZATION, AND CEREBROVASCULAR DISEASE EVENTS
- I-226 FIRST-LINE THERAPY FOR THE TREATMENT OF ADVANCED CARCINOMA OF THE OVARY IN COMBINATION WITH CISPLATIN
- I-227 SHORT-TERM TREATMENT OF SYMPTOMATIC GASTROESOPHAGEAL REFLUX DISEASE (GERD)
- I-228 PREVENTION OF MEAL INDUCED HEARTBURN AT A DOSE OF 75MG TAKEN 30-60 MIN PRIOR TO A MEAL
- I-229 PRILOSEC (OMEPRazole), AMOXICILLIN AND CLARITHROMYCIN FOR THE ERADICATION OF H. PYLORI IN PATIENTS WITH DUODENAL ULCER DISEASE
- I-230 IN COMBINATION WITH CISPLATIN, FOR THE FIRST-LINE TREATMENT OF NON-SMALL CELL LUNG CANCER IN PATIENTS WHO ARE NOT CANDIDATES FOR POTENTIALLY CURATIVE SURGERY AND/OR RADIATION
- I-231 TREATMENT OF PATIENTS WITH LOCALLY ADVANCED OR METASTATIC BREAST CANCER AFTER FAILURE OF PRIOR CHEMOTHERAPY
- I-232 TREATMENT OF RECURRENT MUCOCUTANEOUS HERPES SIMPLEX INFECTIONS IN HIV-AFFECTED PATIENTS AT A DOSE OF 500MG TWICE DAILY
- I-233 PROPHYLACTIC USE TO REDUCE PERIOPERATIVE BLOOD LOSS AND THE NEED FOR BLOOD TRANSFUSION IN PATIENTS UNDERGOING CARDIOPULMONARY BYPASS IN THE COURSE OF CORONARY ARTERY BYPASS GRAFT SURGERY
- I-234 FOR USE IN COMBINATION WITH CISPLATIN FOR THE FIRST-LINE TREATMENT OF PATIENTS WITH INOPERABLE LOCALLY ADVANCED (STAGE IIIA OR IIIB) OR METASTATIC (STAGE IV) NON-SMALL CELL LUNG CANCER
- I-235 PREVENTION OF EXERCISE-INDUCED BRONCHOSPASM IN PATIENTS 12 YEARS OF AGE AND OLDER
- I-236 PREVENTION OF EXERCISE-INDUCED BRONCHOSPASM IN PATIENTS 4 YEARS OF AGE AND OLDER
- I-237 MAINTENANCE TREATMENT OF ASTHMA AND PREVENTION OF BRONCHOSPASM IN PATIENTS 4 YEARS OF AGE AND OLDER
- I-238 ADJUNCTIVE TREATMENT OF LENNOX-GASTAUT SYNDROME IN PEDIATRIC AND ADULT PATIENTS
- I-239 TREATMENT OF PATIENTS WITH HOMOZYGOUS FAMILIAL HYPERCHOLESTEROLEMIA
- I-240 MANAGEMENT OF SECONDARY HYPERPARATHYROIDISM AND RESULTANT METABOLIC BONE DISEASE IN PATIENTS WITH MODERATE TO SEVERE CHRONIC RENAL FAILURE (Cr 15 TO 55ML MIN) NOT YET ON DIALYSIS
- I-241
- I-242 TREATMENT OF MODERATE TO SEVERE VASOMOTOR SYMPTOMS ASSOCIATED WITH THE MENOPAUSE AND IN THE TREATMENT OF VULVAR AND VAGINAL ATROPHY IN WOMEN WITH AN INTACT UTERUS
- I-243 USE IN THE SYMPTOMATIC RELIEF OF RHINORRHEA ASSOCIATED WITH THE COMMON COLD IN CHILDREN AGE 5 TO 11 YEARS
- I-244 REDUCE THE INCIDENCE OF BREAST CANCER IN WOMEN AT HIGH RISK FOR BREAST CANCER
- I-245 TREATMENT OF ACUTE SINUSITIS

PATENT AND EXCLUSIVITY TERMS

PATENT USE CODE

U-215	TREATMENT OF EPILEPSY TWICE DAILY. TREATING A PATIENT BY ADMINISTERING CARBAMAZEPINE IN A DOSAGE FORM CAPABLE OF MAINTAINING BLOOD CONCENTRATION FROM 4-12MCG/ML OVER 12 HOURS
U-216	TREATMENT OF ADENOCARCINOMA, INCLUDING STAGE B2-C, BY ADMINISTERING AN AGONIST OF LR-RH AND FLUTAMIDE
U-217	METHOD OF PRODUCING ANESTHESIA
U-218	METHOD FOR LIMITING THE POTENTIAL FOR MICROBIAL GROWTH IN THE DRUG PRODUCT
U-219	TREATMENT OF PARKISON'S DISEASE
U-220	METHOD OF DIAGNOSIS
U-221	SELECTIVE VASODILATION BY CONTINUOUS ADENOSINE INFUSION
U-222	METHOD OF TREATING PAGETS DISEASE USING ACTONEL
U-223	TREATMENT OF BACTERIAL CONJUNCTIVITIS CAUSED BY SUSCEPTIBLE STRAINS OF MICROORGANISMS
U-224	CONTROLLING INTRAOCULAR PRESSURE
U-225	METHOD FOR DELIVERY
U-226	METHOD OF ENHANCING THE DISSOLUTION PROFILE OF A PHARMACEUTICAL FROM A SOLID DOSAGE FORM CONTAINING THE PHARMACEUTICAL AND SIMETHICONE
U-227	NASAL ADMINISTRATION
U-228	ASTHMA
U-229	CARDIAC INSUFFICIENCY (CONGESTIVE HEART FAILURE)
U-230	PREVENTION OF ACUTE CARDIAC ISCHEMIC EVENTS
U-231	USE IN PARKINSON'S DISEASE
U-232	METHOD OF TREATING MIGRAINE
U-233	DECREASING MORTALITY CAUSED BY CONGESTIVE HEART FAILURE
U-234	METHOD OF USING RIBAVIRIN TO TREAT VIRAL INFECTIONS IN MAMMALS
U-235	METHOD OF MODULATING TH1 AND TH2 RESPONSE IN ACTIVATED T CELLS OF A HUMAN COMPRISING ADMINISTERING RIBAVIRIN TO THE T CELLS IN A DOSAGE WHICH PROMOTES THE TH1 RESPONSE AND SUPPRESSES THE TH2 RESPONSE
U-236	TREATING MALE PATTERN BALDNESS WITH 0.05 TO 3 MG/DAY
U-237	METHOD OF PERFORMING NMR IMAGING WITH A PATIENT COMPRISING ADMINISTERING TO THE PATIENT AN EFFECTIVE AMOUNT OF CONTRAST AGENT DISCLOSED IN THE CLAIMS
U-238	IMAGING A BODY TISSUE AND SUBJECTING TO NMR TOMOGRAPHY, ADMINISTERING AN AMOUNT OF PHARMACEUTICAL AGENT FOR AFFECTING THE RELAXATION TIMES OF ATOMS IN BODY TISSUES UNDERGOING NMR DIAGNOSIS, WHEREBY THE IMAGE CONTRAST IS ENHANCED....
U-239	TREATING OR CONTROLLING OCULAR INFLAMMATION WHICH COMPRISES TOPICALLY ADMINISTERING TO THE AFFECTED EYE A COMPOSITION COMPRISING A NSAID, A POLYMERIC QUATERNARY AMMONIUM COMPOUND AND BORIC ACID
U-240	TREATMENT OF ACUTE MIGRAINE ATTACKS
U-241	FOR SHORT-TERM TREATMENT ACTIVE DUODENAL ULCER, MAINTENANCE THERAPY FOR DUODENAL ULCER PATIENTS AT REDUCED DOSAGE AFTER HEALING OF ACTIVE ULCER, SHORT-TERM TREATMENT ACTIVE BENIGN GASTRIC ULCER & GERD, PATHOLOGICAL HYPERSECRETORY CONDITIONS
U-242	USE OF FOLLITROPIN ALPHA ALONE IN IN-VITRO FERTILIZATION
U-243	TOPICAL ADMINISTRATION
U-244	PLATELET AGGREGATION INHIBITORS
U-245	TREATMENT OF SEBORRHEA DERMATITIS IN HUMANS
U-246	PHOSPHATE BINDING
U-247	TREATMENT OF RHEUMATOID ARTHRITIS

**PRESCRIPTION AND OTC DRUG PRODUCT
PATENT AND EXCLUSIVITY DATA**
*PED and PED represent Pediatric Exclusivity

APPL/PROD NUMBER	INGREDIENT NAME; TRADE NAME	PATENT NUMBER	PATENT/PED EXCL EXPIRES	USE CODE	EXCLUS CODE	EXCLUS EXPIRES
020838 002	CANDESARTAN CILEXETIL; ATACAND	5196444 5534534 5703110	APR 18, 2011 JUL 09, 2013 APR 18, 2011	U-3	NCE	JUN 04, 2003
020838 003	CANDESARTAN CILEXETIL; ATACAND	5705517 5196444 5534534 5703110	APR 18, 2011 APR 18, 2011 JUL 09, 2013 APR 18, 2011	U-3	NCE	JUN 04, 2003
020838 004	CANDESARTAN CILEXETIL; ATACAND	5705517 5196444 5534534 5703110	APR 18, 2011 APR 18, 2011 JUL 09, 2013 APR 18, 2011	U-3	NCE	JUN 04, 2003
020896 001	CAPECITABINE; XELODA				NCE	APR 30, 2003
020896 002	CAPECITABINE; XELODA				NCE	APR 30, 2003
020712 001	CARBAMAZEPINE; CARBATROL	5326570	JUL 05, 2011	U-215		
020712 002	CARBAMAZEPINE; CARBATROL	5326570	JUL 05, 2011	U-215		
019972 001	CARTEOLOL HYDROCHLORIDE; OCUPRESS	4309432	JAN 02, 2000			
020297 001	CARVEDILOL; COREG	4503067 5760069	MAR 05, 2007 JUN 07, 2015	U-3 U-233		
020297 002	CARVEDILOL; COREG	4503067 5760069	MAR 05, 2007 JUN 07, 2015	U-3 U-233		
020297 003	CARVEDILOL; COREG	4503067 5760069	MAR 05, 2007 JUN 07, 2015	U-3 U-233		
020297 004	CARVEDILOL; COREG	4503067 5760069	MAR 05, 2007 JUN 07, 2015	U-3 U-233		
020774 001	CHLORHEXIDINE GLUCONATE; PERIOCHIP				NP	MAY 15, 2001
020238 002	CINETIDINE; TAGAMET HB				D-41	JUN 05, 2001
020369 001	CIPROFLOXACIN HYDROCHLORIDE; CILLOXAN	4670444	JUN 02, 2004	U-223	NDF	MAR 30, 2001
020805 001	CIPROFLOXACIN HYDROCHLORIDE; CIPRO HC	4670444 4844902	DEC 09, 2003 FEB 11, 2008		NC	FEB 10, 2001
019847 001	CIPROFLOXACIN; CIPRO				1-164 1-245	SEP 11, 1999 JUN 03, 2000
020780 001	CIPROFLOXACIN; CIPRO	4670444	DEC 09, 2003			
020780 002	CIPROFLOXACIN; CIPRO	4670444	DEC 09, 2003			
019857 001	CIPROFLOXACIN; CIPRO IN DEXTROSE 5%				1-164 1-245	SEP 11, 1999 JUN 03, 2000
019858 001	CIPROFLOXACIN; CIPRO IN SODIUM CHLORIDE 0.9%				1-164 1-245	SEP 11, 1999 JUN 03, 2000
020551 001	CISATRACURIUM BESYLATE; NIMBEX				D-48	OCT 15, 2001
020551 002	CISATRACURIUM BESYLATE; NIMBEX PRESERVATIVE FREE				D-48	OCT 15, 2001
020551 003	CISATRACURIUM BESYLATE; NIMBEX PRESERVATIVE FREE				D-48	OCT 15, 2001
020822 002	CITALOPRAM HYDROBROMIDE; CELEXA				NCE	JUL 17, 2003
020822 003	CITALOPRAM HYDROBROMIDE; CELEXA				NCE	JUL 17, 2003
020822 004	CITALOPRAM HYDROBROMIDE; CELEXA				NCE	JUL 17, 2003
020839 001	CLOPIDOGREL BISULFATE; PLAVIX	4529596 4847265 5576328	JUL 05, 2003 FEB 12, 2008 JAN 31, 2014			
017922 001	DESMOPRESSIN ACETATE; DDAVP	5763407	DEC 23, 2013			
017922 002	DESMOPRESSIN ACETATE; DDAVP	5763407	DEC 23, 2013			

**PRESCRIPTION AND OTC DRUG PRODUCT
PATENT AND EXCLUSIVITY DATA**
*PED and PED represent Pediatric Exclusivity

APPL/PROD NUMBER	INGREDIENT NAME; TRADE NAME	PATENT NUMBER	PATENT/PED EXCL EXPIRES	USE CODE	EXCLUS CODE	EXCLUS EXPIRES
017922 003	DESMOPRESSIN ACETATE;DDAVP	5763407	DEC 23, 2013			
018938 001	DESMOPRESSIN ACETATE;DDAVP	5763407	DEC 23, 2013			
018938 002	DESMOPRESSIN ACETATE;DDAVP	5763407	DEC 23, 2013			
019955 001	DESMOPRESSIN ACETATE;DDAVP	5763407	DEC 23, 2013		1-40	MAR 25, 2001
019955 002	DESMOPRESSIN ACETATE;DDAVP	5763407	DEC 23, 2013		1-40	MAR 25, 2001
020713 001	DESOGESTREL;MIRCETTE				NP	APR 22, 2001
020344 001	DEXFENFLURAMINE HYDROCHLORIDE;REDUX	4309445	FEB 19, 2004	U-133		
020809 001	DICLOFENAC SODIUM;DICLOFENAC SOD;UM	5603929	NOV 16, 2014	U-239		
		5653972	NOV 16, 2014	U-239		
020037 001	DICLOFENAC SODIUM;VOLTAREN				1-213	FEB 25, 2001
020148 001	DINHYDROERGOTAMINE MESYLATE;MIGRANAL	4758423	JUL 31, 2001			
		4462983	JUL 31, 2001	U-227		
		5169849	DEC 08, 2009	U-227		
020401 001	DILTIAZEM HYDROCHLORIDE;TIAZAC				1-133	JAN 30, 2001
020401 002	DILTIAZEM HYDROCHLORIDE;TIAZAC				1-133	JAN 30, 2001
020401 003	DILTIAZEM HYDROCHLORIDE;TIAZAC				1-133	JAN 30, 2001
020401 004	DILTIAZEM HYDROCHLORIDE;TIAZAC				1-133	JAN 30, 2001
020401 005	DILTIAZEM HYDROCHLORIDE;TIAZAC				1-133	JAN 30, 2001
020449 001	DOCETAXEL;TAXOTERE				1-231	JUN 22, 2001
020690 001	DONEPEZIL HYDROCHLORIDE;ARICEPT	4895841	NOV 25, 2010			
020690 002	DONEPEZIL HYDROCHLORIDE;ARICEPT	4895841	NOV 25, 2010			
020869 001	DORZOLAMIDE HYDROCHLORIDE;COSOPT				NC	APR 07, 2001
020972 001	EFAVIRENZ;SUSTIVA				NCE	SEP 17, 2003
020972 002	EFAVIRENZ;SUSTIVA				NCE	SEP 17, 2003
020972 003	EFAVIRENZ;SUSTIVA				NCE	SEP 17, 2003
020164 001	ENOXAPARIN SODIUM;LOVENOX				1-217	JAN 30, 2001
					1-222	MAR 27, 2001
020164 002	ENOXAPARIN SODIUM;LOVENOX				1-222	MAR 27, 2001
					1-217	JAN 30, 2001
020738 004	EPROSARTAN MESYLATE;TEVETEN	5185351	FEB 09, 2010	U-3	D-40	OCT 28, 2001
020738 005	EPROSARTAN MESYLATE;TEVETEN	5185351	FEB 09, 2010	U-3	D-40	OCT 28, 2001
020718 001	EPTIFIBATIDE;INTEGRILIN	5756451	NOV 11, 2014		NCE	MAY 18, 2003
		5686570	NOV 11, 2014			
		5807825	SEP 15, 2015	U-244		
020718 002	EPTIFIBATIDE;INTEGRILIN	5756451	NOV 11, 2014		NCE	MAY 18, 2003
		5686570	NOV 11, 2014			
		5807825	SEP 15, 2015	U-244		
020907 001	ESTRADIOL;ACTIVELLE				NC	NOV 18, 2001
					1-242	NOV 18, 2001
020375 003	ESTRADIOL;CLIMARA	5223261	JUN 29, 2010			
020870 001	ESTRADIOL;COMBIPATCH				NP	AUG 07, 2001
020870 002	ESTRADIOL;COMBIPATCH				NP	AUG 07, 2001
020527 002	ESTROGENS, CONJUGATED;PREMPHASE 14/14	5547948	JAN 17, 2015			
020527 001	ESTROGENS, CONJUGATED;PREMPRO 14/14	5547948	JAN 17, 2015			
083209 001	ESTROGENS, ESTERIFIED;ESTRATAB				1-214	MAR 10, 2001
086715 001	ESTROGENS, ESTERIFIED;ESTRATAB				1-214	MAR 10, 2001
020363 001	FANCICLOVIR;FAMVIR				NCE	JUN 29, 1999
					1-232	JUN 12, 2001

**PRESCRIPTION AND OTC DRUG PRODUCT
PATENT AND EXCLUSIVITY DATA**
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APPL/PROD NUMBER	INGREDIENT NAME; TRADE NAME	PATENT NUMBER	PATENT/PED EXCL EXPIRES	USE CODE	EXCLUS CODE	EXCLUS EXPIRES
020363 002	FAMCICLOVIR; FAMVIR				1-232	JUN 12, 2001
020363 003	FAMCICLOVIR; FAMVIR				1-232	JUN 12, 2001
020325 001	FAMOTIDINE; PEPICID AC				D-47	NOV 09, 2001
019510 004	FAMOTIDINE; PEPICID PRESERVATIVE FREE	4283408	OCT 15, 2000			
020752 001	FAMOTIDINE; PEPICID RPD	4283408	OCT 15, 2000			
		4305502	DEC 15, 1998			
		4371516	JAN 31, 2000	U-241		
020752 002	FAMOTIDINE; PEPICID RPD	4283408	OCT 15, 2000			
		4305502	DEC 15, 1998			
		4371516	JAN 31, 2000	U-241		
020625 001	FEXOFENADINE HYDROCHLORIDE; ALLEGRA	4254129	FEB 18, 2001	U-139		
020786 001	FEXOFENADINE HYDROCHLORIDE; ALLEGRA-D	4254129	FEB 18, 2001		NCE	JUL 25, 2001
		5375693	AUG 03, 2012			
		5578610	NOV 26, 2013			
020788 001	FINASTERIDE; PROPECIA	5547957	OCT 15, 2013	U-236		
020180 001	FINASTERIDE; PROSCAR				1-221	MAR 20, 2001
018830 001	FLECAINIDE ACETATE; TANBOCOR	4642384	FEB 10, 2004			
018830 002	FLECAINIDE ACETATE; TANBOCOR	4642384	FEB 10, 2004			
018830 003	FLECAINIDE ACETATE; TANBOCOR	4642384	FEB 10, 2004			
018830 004	FLECAINIDE ACETATE; TANBOCOR	4642384	FEB 10, 2004			
018554 001	FLUTAMIDE; EULEXIN	4472382	SEP 18, 2001	U-24		
		5712251	SEP 18, 2001	U-216		
020121 001	FLUTICASONE PROPIONATE; FLOMASE				1-224	OCT 31, 2000
020378 001	FOLLITROPIN ALFA/BETA; GONAL-F	4589402	JUL 26, 2004	U-242		
		5767251	JUN 16, 2015			
020378 002	FOLLITROPIN ALFA/BETA; GONAL-F	4589402	JUL 26, 2004	U-242		
		5767251	JUN 16, 2015			
020961 001	FOMIVIRSEN SODIUM; VITRAVENE PRESERVATIVE FREE				NCE	AUG 26, 2003
020450 001	FOSPHENYTOIN SODIUM; CEREBYX	4260769	APR 07, 2003			
020235 001	GABAPENTIN; NEURONTIN				D-43	SEP 29, 2001
020235 002	GABAPENTIN; NEURONTIN				D-43	SEP 29, 2001
020235 003	GABAPENTIN; NEURONTIN				D-43	SEP 29, 2001
020882 001	GABAPENTIN; NEURONTIN	4087544	JAN 16, 2000			
		4894476	MAY 02, 2008			
		5084479	JAN 02, 2010			
020882 002	GABAPENTIN; NEURONTIN	4087544	JAN 16, 2000			
		4894476	MAY 02, 2008			
		5084479	JAN 02, 2010			
019596 001	GADOPENTETATE DIMEGLUMINE; MAGNEVIST	5560903	OCT 01, 2013			
020460 001	GANCICLOVIR; CYTOVENE	4423050	MAY 21, 2001	U-64		
		4642346	JUN 24, 2005			
		4507305	MAY 12, 2001	U-64		
020509 001	GEMCITABINE HYDROCHLORIDE; GENZAR				1-234	AUG 26, 2001
020509 002	GEMCITABINE HYDROCHLORIDE; GENZAR				1-234	AUG 26, 2001
020695 001	GREPAFLOXACIN HYDROCHLORIDE; RAXAR	5563138	OCT 08, 2013			
020818 001	HYDROCHLOROTHIAZIDE; DIOVAN HCT	5399578	MAR 21, 2012	U-3	NCE	DEC 23, 2001
					NC	MAR 06, 2001
020818 002	HYDROCHLOROTHIAZIDE; DIOVAN HCT	5399578	MAR 21, 2012	U-3	NCE	DEC 23, 2001
					NC	MAR 06, 2001

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APPL/PROD NUMBER	INGREDIENT NAME; TRADE NAME	PATENT NUMBER	PATENT/PED EXCL EXPIRES	USE CODE	EXCLUS CODE	EXCLUS EXPIRES
020716 001	HYDROCODONE BITARTRATE; VICOPROFEN	4587252	DEC 18, 2004	U-55		
016295 002	HYDROXYUREA; DROXIA				ODE	FEB 25, 2005
016295 003	HYDROXYUREA; DROXIA				ODE	FEB 25, 2005
016295 004	HYDROXYUREA; DROXIA				ODE	FEB 25, 2005
019771 001	IBUPROFEN; ADVIL COLD AND SINUS	4552899	NOV 12, 2002			
		4552899*PED	MAY 12, 2003			
019833 002	IBUPROFEN; CHILDREN'S ADVIL	4788220	NOV 29, 2005			
		4788220*PED	MAY 29, 2006			
020589 001	IBUPROFEN; CHILDREN'S ADVIL	4788220	JUL 08, 2007		NP	JUN 16, 1998
		4788220*PED	JAN 08, 2008		PED	DEC 16, 1998
020516 001	IBUPROFEN; CHILDREN'S MOTRIN	5374659	DEC 20, 2011		NP	JUN 16, 1998
		5374659*PED	JUN 20, 2012		PED	DEC 16, 1998
020601 001	IBUPROFEN; CHILDREN'S MOTRIN	5215755	JUN 01, 2010		NP	NOV 15, 1999
		5215755*PED	DEC 01, 2010		PED	MAY 15, 2000
020603 001	IBUPROFEN; CHILDREN'S MOTRIN	5374659	DEC 20, 2011		NP	JUN 16, 1998
		5374659*PED	JUN 20, 2012		PED	DEC 16, 1998
020267 002	IBUPROFEN; JUNIOR STRENGTH ADVIL				NP	JUN 16, 1998
					PED	DEC 16, 1998
020601 003	IBUPROFEN; JUNIOR STRENGTH MOTRIN	5215755	JUN 01, 2010		NP	NOV 15, 1999
		5215755*PED	DEC 01, 2010		PED	MAY 15, 2000
020602 001	IBUPROFEN; JUNIOR STRENGTH MOTRIN				NP	JUN 16, 1998
					PED	DEC 16, 1998
019842 001	IBUPROFEN; MOTRIN	5374659	DEC 20, 2011			
		5374659*PED	JUN 20, 2012			
020135 001	IBUPROFEN; MOTRIN	5215755	JUN 01, 2010			
		5320855	JUN 14, 2011			
		5215755*PED	DEC 01, 2010			
		5320855*PED	DEC 14, 2011			
020135 002	IBUPROFEN; MOTRIN	5215755	JUN 01, 2010			
		5320855	JUN 14, 2011			
		5215755*PED	DEC 01, 2010			
		5320855*PED	DEC 14, 2011			
020812 001	IBUPROFEN; PEDIATRIC ADVIL				NP	JUN 16, 1998
					PED	DEC 16, 1998
020903 001	INTERFERON ALFA-2B; REBETRON	4530901	JUL 23, 2002		NP	JUN 03, 2001
		4211771	JUL 08, 1999	U-234		
		5767097	JAN 23, 2016	U-235		
020923 001	IOVERSOL; OPTIRAY 240	4396598	DEC 30, 2002			
020923 002	IOVERSOL; OPTIRAY 320	4396598	DEC 30, 2002			
020923 003	IOVERSOL; OPTIRAY 350	4396598	DEC 30, 2002			
020393 001	IPRATROPIUM BROMIDE; ATROVENT				1-223	APR 01, 2001
020394 001	IPRATROPIUM BROMIDE; ATROVENT				1-243	NOV 09, 2001
020571 001	IRINOTECAN HYDROCHLORIDE; CAMPTOSAR	4604463	AUG 20, 2007			
020083 001	ITRACONAZOLE; SPORANOX	5633015	MAY 27, 2014			
020657 001	ITRACONAZOLE; SPORANOX	4267179	JUN 23, 2000			
		5707975	JAN 13, 2015			
		4727064	FEB 23, 2005			
019927 001	KETOCONAZOLE; NIZORAL	4942162	FEB 11, 2003			
020310 001	KETOCONAZOLE; NIZORAL A-D	4942162	FEB 11, 2003	U-245		
		4335125	JUN 15, 1999			
		5456851	APR 07, 2014			

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020241 001	LANOTRIGINE; LANICTAL				ODE 1-238	AUG 24, 2005 AUG 24, 2001
020241 002	LANOTRIGINE; LANICTAL				ODE 1-238	AUG 24, 2005 AUG 24, 2001
020241 003	LANOTRIGINE; LANICTAL				ODE 1-238	AUG 24, 2005 AUG 24, 2001
020241 004	LANOTRIGINE; LANICTAL				ODE 1-238	AUG 24, 2005 AUG 24, 2001
020241 005	LANOTRIGINE; LANICTAL				ODE 1-238	AUG 24, 2005 AUG 24, 2001
020241 006	LANOTRIGINE; LANICTAL				ODE 1-238	AUG 24, 2005 AUG 24, 2001
020764 001	LANOTRIGINE; LANICTAL CD	5698226 4602017	JAN 29, 2012 JUL 22, 2008	U-106	ODE 1-238	AUG 24, 2005 AUG 24, 2001
020764 002	LANOTRIGINE; LANICTAL CD	5698226 4602017	JAN 29, 2012 JUL 22, 2008	U-106	ODE 1-238	AUG 24, 2005 AUG 24, 2001
020764 003	LANOTRIGINE; LANICTAL CD	5698226 4602017	JAN 29, 2012 JUL 22, 2008	U-106	ODE 1-238	AUG 24, 2005 AUG 24, 2001
020406 001	LANSOPRAZOLE; PREVACID				1-227 D-42	MAR 12, 2001 JUL 20, 2001
020406 002	LANSOPRAZOLE; PREVACID				1-227 D-42	MAR 12, 2001 JUL 20, 2001
020905 001	LEFLUNOMIDE; ARAVA	4284786 4351841 5679709	DEC 13, 1999 DEC 13, 1999 OCT 21, 2014	U-247 U-247 U-247	NCE	SEP 10, 2003
020905 002	LEFLUNOMIDE; ARAVA	4284786 4351841 5679709	DEC 13, 1999 DEC 13, 1999 OCT 21, 2014	U-247 U-247 U-247	NCE	SEP 10, 2003
020905 003	LEFLUNOMIDE; ARAVA	4284786 4351841 5679709	DEC 13, 1999 DEC 13, 1999 OCT 21, 2014	U-247 U-247 U-247	NCE	SEP 10, 2003
020807 001	LEPIRUDIN; REFLUDAN	5180668	JAN 19, 2010		ODE NCE	MAR 06, 2005 MAR 06, 2003
019732 001	LEUPROLIDE ACETATE; LUPRON DEPOT	5716640	SEP 02, 2013			
020011 001	LEUPROLIDE ACETATE; LUPRON DEPOT	5716640	SEP 02, 2013			
020517 001	LEUPROLIDE ACETATE; LUPRON DEPOT	5716640	SEP 02, 2013			
020263 002	LEUPROLIDE ACETATE; LUPRON DEPOT-PED	5716640	SEP 02, 2013			
020263 003	LEUPROLIDE ACETATE; LUPRON DEPOT-PED	5716640	SEP 02, 2013			
020263 004	LEUPROLIDE ACETATE; LUPRON DEPOT-PED	5716640	SEP 02, 2013			
020263 005	LEUPROLIDE ACETATE; LUPRON DEPOT-PED	5716640	SEP 02, 2013			
020263 006	LEUPROLIDE ACETATE; LUPRON DEPOT-PED	5716640	SEP 02, 2013			
020708 001	LEUPROLIDE ACETATE; LUPRON DEPOT-3	5716640	SEP 02, 2013			
020517 002	LEUPROLIDE ACETATE; LUPRON DEPOT-4	5716640	SEP 02, 2013			
019941 001	LIDOCAINE; ENLA				1-215 NP	FEB 04, 2001 FEB 04, 2001
020962 001	LIDOCAINE; ENLA					
020606 001	LOPERAMIDE HYDROCHLORIDE; IMODIUM ADVANCED	5716641	MAY 21, 2012	U-226		
020803 001	LOTEPREDNOL ETABONATE; ALREX	4996335 5540930	FEB 26, 2008 OCT 25, 2013		NCE	MAR 09, 2003
020583 001	LOTEPREDNOL ETABONATE; LOTEMAX	4996335 5540930	FEB 26, 2008 OCT 25, 2013		NCE	MAR 09, 2003

**PRESCRIPTION AND OTC DRUG PRODUCT
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APPL/PROD NUMBER	INGREDIENT NAME; TRADE NAME	PATENT NUMBER	PATENT/PED EXCL EXPIRES	USE CODE	EXCLUS CODE	EXCLUS EXPIRES
020841 001	LOTEPREDNOL ETABONATE; LOTEMAX	4996335	FEB 26, 2008		NCE	MAR 09, 2003
019832 003	MAFENIDE ACETATE; SULFAMYLON	5540930	OCT 25, 2013		NDF ODE	JUN 05, 2001 JUN 05, 2005
020652 001	MANGAFODIPIR TRISODIUM; TESLASCAN	4933456 4992554 5091169 5223243 4647447 4657900 RE33239	JUN 12, 2007 FEB 12, 2008 FEB 25, 2009 JUN 29, 2010 MAR 03, 2004 APR 14, 2004 MAY 12, 2004	U-237 U-238		
019618 001	MESALAMINE; ROMASA					
020208 001	METRONIDAZOLE; METROGEL-VAGINAL				D-40	MAY 16, 2000
020901 001	METRONIDAZOLE; METROLOTION				NDF	NOV 24, 2001
020827 001	MICONAZOLE NITRATE; MONISTAT 3				NP	MAR 30, 2001
018654 001	MIDAZOLAM HYDROCHLORIDE; VERSED	4280957	DEC 20, 1999			
018654 002	MIDAZOLAM HYDROCHLORIDE; VERSED	4280957*PED	JUN 20, 2000			
020942 001	MIDAZOLAM HYDROCHLORIDE; VERSED	4280957 4280957*PED	DEC 20, 1999 JUN 20, 2000		PED NDF NCE	APR 15, 2002 OCT 15, 2001 JUN 14, 2001
020415 003	MIRTAZAPINE; REMERON					
020762 001	MONETASONE FURATE MONOHYDRATE; NASONEX	4472393	SEP 18, 2001			
020829 002	MONTELUKAST SODIUM; SINGULAIR	5565473	NOV 30, 2010	U-228	NCE	FEB 20, 2003
020830 001	MONTELUKAST SODIUM; SINGULAIR	5565473	NOV 30, 2010	U-228	NCE	FEB 20, 2003
020763 001	NARATRIPTAN HYDROCHLORIDE; AMERGE				NCE	FEB 10, 2003
020763 002	NARATRIPTAN HYDROCHLORIDE; AMERGE				NCE	FEB 10, 2003
020636 001	NEVIRAPINE; VIRAMUNE				D-49	SEP 11, 2001
020933 001	NEVIRAPINE; VIRAMUNE				NDF	SEP 11, 2001
020536 001	NICOTINE; NICOTROL				NCE	JUN 21, 2001
020555 001	NIZATIDINE; AXID AR	4915950	FEB 12, 2008		1-220 D-39 NDF	APR 01, 2001 APR 01, 2001 DEC 16, 2000
020799 001	OFLOXACIN; FLOXIN					
020688 001	OLOPATADINE HYDROCHLORIDE; PATANOL	5116863	DEC 18, 2010			
019810 001	OMEPRazole; PRILOSEC				1-229	JUN 29, 2001
019810 002	OMEPRazole; PRILOSEC				1-229	JUN 29, 2001
020932 002	OXycodone HYDROCHLORIDE; ROXICODONE				NS	OCT 26, 2001
020262 001	PACLITAXEL; TAXOL				1-226	APR 09, 2001
020819 001	PARICALCITOL; ZEMPLAR				1-230	JUN 30, 2001
020710 001	PAROXETINE HYDROCHLORIDE; PAXIL				NCE	APR 17, 2003
020237 001	PILOCARPINE HYDROCHLORIDE; SALAGEN	5811436	SEP 22, 2015		ODE 1-212	FEB 11, 2005 FEB 11, 2001
020451 001	PORFIMER SODIUM; PHOTOFRIN	5145863	DEC 15, 2009	U-129		
020667 001	PRAMIPEXOLE DIHYDROCHLORIDE; MIRAPEX	4886812 4843086	DEC 12, 2006 JUN 27, 2006	U-231		
020667 002	PRAMIPEXOLE DIHYDROCHLORIDE; MIRAPEX	4886812 4843086	DEC 12, 2006 JUN 27, 2006	U-231		
020667 003	PRAMIPEXOLE DIHYDROCHLORIDE; MIRAPEX	4886812 4843086	DEC 12, 2006 JUN 27, 2006	U-231		

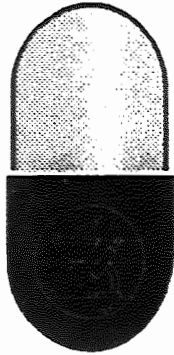
PRESCRIPTION AND OTC DRUG PRODUCT
PATENT AND EXCLUSIVITY DATA
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APPL/PROD NUMBER	INGREDIENT NAME; TRADE NAME	PATENT NUMBER	PATENT/PED EXCL EXPIRES	USE CODE	EXCLUS CODE	EXCLUS EXPIRES
020667 004	PRAMIPEXOLE DIHYDROCHLORIDE; MIRAPEX	4886812	DEC 12, 2006			
		4843086	JUN 27, 2006	U-231		
020667 005	PRAMIPEXOLE DIHYDROCHLORIDE; MIRAPEX	4886812	DEC 12, 2006			
		4843086	JUN 27, 2006	U-231		
019898 002	PRAVASTATIN SODIUM; PRAVACHOL				1-225	MAR 27, 2001
019898 003	PRAVASTATIN SODIUM; PRAVACHOL				1-225	MAR 27, 2001
019898 004	PRAVASTATIN SODIUM; PRAVACHOL				1-225	MAR 27, 2001
019781 001	PROGESTERONE; PROMETRIUM				NP	MAY 14, 2001
019627 002	PROPOFOL; DIPRIVAK	5731355	MAR 22, 2015	U-217		
		5731356	MAR 22, 2015	U-218		
020815 001	RALOXIFENE HYDROCHLORIDE; EVISTA	4418068	APR 03, 2001			
		5393763	JUL 28, 2012	U-114		
		5457117	JUL 28, 2012	U-114		
		5478847	MAR 02, 2014	U-114		
020520 001	RANITIDINE HYDROCHLORIDE; ZANTAC 75				1-228	JUN 08, 2001
021024 001	RIFAPENTINE; PRIFTIN				NCE	JUN 22, 2003
					ODE	JUN 22, 2005
020835 001	RISEDROMATE SODIUM; ACTONEL	5583122	DEC 10, 2013	U-222	NCE	MAR 27, 2003
020272 005	RISPERIDONE; RISPERDAL	5158952	OCT 27, 2009		D-37	OCT 17, 2000
020864 001	RIZATRIPTAN BENZOATE; MAXALT	5298520	JAN 28, 2012	U-240	NCE	JUN 29, 2003
		5602162	MAY 10, 2015	U-240		
020864 002	RIZATRIPTAN BENZOATE; MAXALT	5298520	JAN 28, 2012	U-240	NCE	JUN 29, 2003
		5602162	MAY 10, 2015	U-240		
020865 001	RIZATRIPTAN BENZOATE; MAXALT-MLT	4305502	DEC 15, 1998		NCE	JUN 29, 2003
		5298520	JAN 28, 2012	U-240		
		4758598	DEC 15, 1998			
		4371516	FEB 01, 2000			
		5602162	MAY 10, 2015	U-240		
020865 002	RIZATRIPTAN BENZOATE; MAXALT-MLT	4305502	DEC 15, 1998		NCE	JUN 29, 2003
		5298520	JAN 28, 2012	U-240		
		4758598	DEC 15, 1998			
		4371516	FEB 01, 2000			
		5602162	MAY 10, 2015	U-240		
020533 001	ROPIVACAINE HYDROCHLORIDE MONOHYDRATE; NAROPIN	4870086	JUL 28, 2010			
020533 003	ROPIVACAINE HYDROCHLORIDE MONOHYDRATE; NAROPIN	4870086	JUL 28, 2010			
020533 004	ROPIVACAINE HYDROCHLORIDE MONOHYDRATE; NAROPIN	4870086	JUL 28, 2010			
020533 005	ROPIVACAINE HYDROCHLORIDE MONOHYDRATE; NAROPIN	4870086	JUL 28, 2010			
020772 001	SACROSIDASE; SUCRAID				ODE	APR 09, 2005
					NCE	APR 09, 2003
020236 001	SALMETEROL XINAFOATE; SEREVENT	5126375	FEB 12, 2008		1-216	FEB 05, 2001
020692 001	SALMETEROL XINAFOATE; SEREVENT	5225445	FEB 12, 2008	U-211	1-237	SEP 25, 2001
		5380922	JAN 10, 2012		1-236	SEP 25, 2001
		5590645	MAR 01, 2011			
		5126375	FEB 12, 2008			
		0342994	JAN 04, 2008			
020828 001	SAQUINAVIR; FORTOVASE	5196438	NOV 19, 2012			
020443 001	SERMORELIN ACETATE; GEREf	4517181	MAY 14, 2002			
		4703035	DEC 28, 2004			
020443 002	SERMORELIN ACETATE; GEREf	4517181	MAY 14, 2002			
		4703035	DEC 28, 2004			

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APPL/PROD NUMBER	INGREDIENT NAME; TRADE NAME	PATENT NUMBER	PATENT/PED EXCL EXPIRES	USE CODE	EXCLUS CODE	EXCLUS EXPIRES
020697 002	TOLCAPONE;TASMAR	5236952	AUG 17, 2010		NCE	JAN 29, 2003
		5476875	DEC 19, 2012	U-219		
020771 001	TOLTERODINE TARTRATE;DETROL	5382600	JAN 17, 2012		NCE	MAR 25, 2003
020771 002	TOLTERODINE TARTRATE;DETROL	5382600	JAN 17, 2012		NCE	MAR 25, 2003
020844 001	TOPIRAMATE;TOPAMAX SPRINKLE				NCE	DEC 24, 2001
020844 002	TOPIRAMATE;TOPAMAX SPRINKLE				NCE	DEC 24, 2001
020844 003	TOPIRAMATE;TOPAMAX SPRINKLE				NCE	DEC 24, 2001
020671 001	TOPOTECAN HYDROCHLORIDE;HYCANTIN	5004758	MAY 28, 2010			
020137 002	TORSEMIDE;DEMADEX				D-38	FEB 13, 2001
020281 001	TRAMADOL HYDROCHLORIDE;ULTRAM				D-44	AUG 21, 2001
020281 002	TRAMADOL HYDROCHLORIDE;ULTRAM				D-44	AUG 21, 2001
020528 001	TRANDOLAPRIL;MAVIK	5744496	APR 28, 2015	U-229		
020528 002	TRANDOLAPRIL;MAVIK	5744496	APR 28, 2015	U-229		
020528 003	TRANDOLAPRIL;MAVIK	5744496	APR 28, 2015	U-229		
020719 001	TROGLITAZONE;PRELAY	4572912	NOV 09, 2008			
020719 002	TROGLITAZONE;PRELAY	4572912	NOV 09, 2008			
020719 003	TROGLITAZONE;PRELAY	4572912	NOV 09, 2008			
020720 001	TROGLITAZONE;REZULIN	4572912	NOV 09, 2008			
020720 002	TROGLITAZONE;REZULIN	4572912	NOV 09, 2008			
020720 003	TROGLITAZONE;REZULIN	4572912	NOV 09, 2008			
020586 001	UREA, C-13;MERETEK UBT KIT (W/ PRAMACTIN)	4830010	OCT 27, 2009	U-147		
020675 001	URSODIOL;URSO	4859660	AUG 22, 2006			
020892 001	VALRUBICIN;VALSTAR PRESERVATIVE FREE				NCE ODE	SEP 25, 2003 SEP 25, 2005
020699 001	VENLAFAXINE HYDROCHLORIDE;EFFEXOR XR	4535186	DEC 13, 2007			
020699 002	VENLAFAXINE HYDROCHLORIDE;EFFEXOR XR	4535186	DEC 13, 2007			
020699 003	VENLAFAXINE HYDROCHLORIDE;EFFEXOR XR	4535186	DEC 13, 2007			
020699 004	VENLAFAXINE HYDROCHLORIDE;EFFEXOR XR	4535186	DEC 13, 2007			
020388 001	VINORELBINE TARTRATE;NAVELBINE	4307100	JUL 08, 2002			
020547 001	ZAFIRLUKAST;ACCOLATE	4859692	SEP 27, 2010			
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