

APPROVED DRUG PRODUCTS

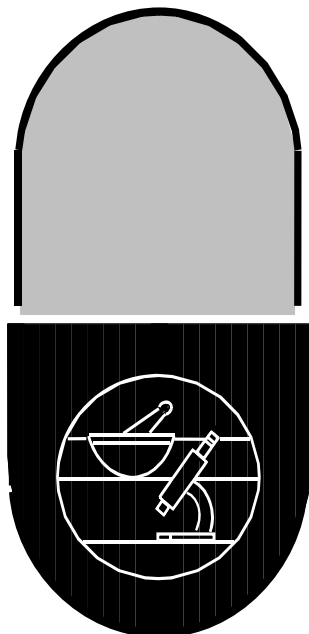
With Therapeutic Equivalence Evaluations



The "Orange Book"

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**CUMULATIVE
SUPPLEMENT 12
December 2006**



**APPROVED
DRUG PRODUCTS**

**WITH
THERAPEUTIC EQUIVALENCE EVALUATIONS**

26th EDITION

Department of Health and Human Services

**Food and Drug Administration
Center for Drug Evaluation and Research
Office of Generic Drugs**

2006

Prepared By
Office of Generic Drugs
Center for Drug Evaluation and Research
Food and Drug Administration

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THERAPEUTIC EQUIVALENCE EVALUATIONS**

26th EDITION

Cumulative Supplement 12

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CONTENTS

	<i>PAGE</i>
1.0 INTRODUCTION	iii
1.1 How to use the Cumulative Supplement	iii
1.2 Cumulative Supplement Content	iv
1.3 Applicant Name Changes.....	v
1.4 Availability of the Edition	vi
1.5 Report of Counts for the Prescription Drug Product List	vii
1.6 Zocor (simvastatin) patent relisting	viii
1.7 Levothyroxine Sodium.....	viii
1.8 Cumulative Supplement Legend	x
 DRUG PRODUCT LISTS	
Prescription Drug Product List	1-1
OTC Drug Product List	2-1
Drug Products with Approval under Section 505 of the Act	
Administered by the Center for Biologics Evaluation and Research List.....	3-1
Orphan Product Designations and Approvals List	4-1
Drug Products Which Must Demonstrate in vivo Bioavailability	
Only if Product Fails to Achieve Adequate Dissolution	5-1
 PATENT AND EXCLUSIVITY INFORMATION ADDENDUM	
A. Patent and Exclusivity Lists	A-1
B. Patent and Exclusivity Terms	B-1

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26th EDITION

**CUMULATIVE SUPPLEMENT 12
December 2006**

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, 25th 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, are for exportation, are for military use, 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.

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 mark 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.

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.)

Products that have never been marketed, are for exportation, are for military use, or 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 25th Edition List will then be added to the "Discontinued Drug Product List" appearing in the 26th Edition. The current edition

Section 2. How To Use The Drug Product Lists describes the layout and usage of the List.

New additions to the Prescription Drug Product List and OTC Drug Product List are indicated by the symbol >A>. The Patent and Exclusivity List new additions are indicated by the symbol >A> to the left of Patent Number or Exclusivity Code. The >A> symbol is then dropped in subsequent Cumulative Supplements for that item.

New deletions to the Prescription Drug Product List and OTC Drug Product List are indicated by the symbol >D> (DELETE) to the left of the line. The information line with the >D> symbol is dropped in subsequent Cumulative Supplements for that item.

The Patent and Exclusivity List is arranged in alphabetical order by active ingredient name(s) and trade name. The trade name will follow the active ingredient name separated by a dash symbol. 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. Drug substance and drug product patents are indicated as such with DS or DP in the Patent codes column. Use patents are indicated with the symbol "U" followed by a number representing a specific use. Exclusivity information for a specific drug is indicated by an abbreviation followed by the date upon which the exclusivity expires. Refer to the Exclusivity Terms, Section B, in the Patent and Exclusivity Information Addendum for an explanation of all codes and abbreviations. Refer to Section 1.3 for internet access to the most current list of Patent and Exclusivity terms.

1.2 CUMULATIVE SUPPLEMENT CONTENT

Since February 2005, we have been providing daily Electronic Orange Book (EOB) product information for new generic drug approvals. Daily generic updates provide the consumer with the current list of approved generic products which is important for substitution purposes. Previously, a first-time-generic product approved early in the month would not be published in the Cumulative Supplement (CS) for several weeks.

The CS monthly update publish goal is by the end of the following month's second work week (e.g., November's supplement will be updated by the end of the second full work week in December).

Currently, the monthly PDF CS includes:

- Generic product ANDA (Abbreviated New Drug Approval) approvals as of the date of publication.
- All product changes received and processed as of the date of publication.
 - o Refer to CS Section 1.8 Cumulative Supplement Legend for types of changes
 - o Discontinued products will be processed as of the date of publication. There will be circumstances where a product is discontinued in one month, however, it will be reported in a different month's CS. For example, the Orange Book Staff received a letter November 7 that the product has been discontinued from manufacturing and marketing. The Orange Book subsequently publishes the October CS on November 14. The product will show in the October CS that it is discontinued even though the date of discontinuance is the

day that the Orange Book Staff receives notification (November 7).

- New Drug Application (NDA) approvals (20,000 and 50,000 series) appear in the CS month they were approved.
- Patent information, also updated daily in the EOB, is current to the date of publication.
- Exclusivity information is updated monthly and current to the date of publication.

1.3 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 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. The Electronic Orange Book Query, updated monthly, will contain the most current applicant holder name.

<u>FORMER APPLICANT NAME</u> <u>(FORMER ABBREVIATED NAME)</u>	<u>NEW APPLICANT NAME</u> <u>(NEW ABBREVIATED NAME)</u>
AMERICAN PHARMACEUTICAL PARTNERS INC (AM PHARM PARTNERS)	ABRAXIS PHARMACEUTICAL PRODUCTS (ABRAXIS PHARM)
AMERICAN PHARMACEUTICAL CO INC SUB BURR CORP (AM PHARM)	ABRAXIS PHARMACEUTICAL PRODUCTS (ABRAXIS PHARM)
AMIDE PHARMACEUTICAL INC (AMIDE PHARM)	ACTAVIS TOTOWA LLC (ACTAVIS TOTOWA)
APOTEX CORP (APOTEX)	APOTEX INC ETOBICOKE SITE (APOTEX INC)
APOTEX CORP (APOTEX)	APOTEX INC RICHMOND HILL (APOTEX INC)
APOTEX INC (APOTEX)	APOTEX INC ETOBICOKE SITE (APOTEX INC)
APOTEX INC (APOTEX)	APOTEX INC RICHMOND HILL (APOTEX INC)
APOTEX INC (APOTEX INC)	APOTEX INC ETOBICOKE SITE (APOTEX INC)
APOTEX INC (APOTEX INC)	APOTEX INC RICHMOND HILL (APOTEX INC)
AVENTIS PHARMACEUTICALS INC (AVENTIS)	SANOFI AVENTIS US LLC (SANOFI AVENTIS US)
AVENTIS PHARMACEUTICAL PRODUCTS INC (AVENTIS PHARMS)	SANOFI AVENTIS US LLC (SANOFI AVENTIS US)
CLAY PARK LABORATORIES INC	PERRIGO NEW YORK INC

(CLAY PARK)	(PERRIGO NEW YORK)
CLAY PARK LABS INC	PERRIGO NEW YORK INC
(CLAY PARK)	(PERRIGO NEW YORK)
DERMIK LABORATORIES INC	SANOFI AVENTIS US LLC
(DERMIK LABS)	(SANOFI AVENTIS US)
DERMIK LABORATORIES INC SUB RORER	SANOFI AVENTIS US LLC
(DERMIK LABS)	(SANOFI AVENTIS US)
FIRST HORIZON PHARMACEUTICAL COMPANY	SCIELE PHARMA INC
(FIRST HORIZON)	(SCIELE PHARMA INC)
LOREX PHARMACEUTICALS	SANOFI AVENTIS US LLC
(LOREX)	(SANOFI AVENTIS US)
MARTEC PHARMACEUTICALS	MARTEC USA LLC
(MARTEC)	(MARTEC USA LLC)
MARTEC SCIENTIFIC INC	MARTEC USA LLC
(MARTEC)	(MARTEC USA LLC)
MCNEIL CONSUMER AND SPECIALTY	MCNEIL PEDIATRICS
PHARMACEUTICALS DIV MCNEIL PCC IN	
(MCNEIL CONS SPECLT)	(MCNEIL PED)
ORPHAN MEDICAL INC	JAZZ PHARMACEUTICALS
(ORPHAN MEDCL)	(JAZZ)
PHARMACEUTICAL FORMULATIONS INC	LEINER HEALTH PRODUCTS INC
(PHARM FORM)	(LEINER HLTH PRODS)
PHARMA TEK INC	X GEN PHARMACEUTICALS INC
(PHARMA TEK)	(X GEN)
PRIVATE FORMULATIONS INC	LEINER HEALTH PRODUCTS INC
(PRIVATE FMLTNS)	(LEINER HLTH PRODS)
SANKYO PHARMA INC	DAIICHI SANKYO INC
(SANKYO)	(DAIICHI SANKYO)
SANOFI AVENTIS US INC	SANOFI AVENTIS US LLC
(SANOFI AVENTIS US)	(SANOFI AVENTIS US)
SANOFI-AVENTIS US INC	SANOFI AVENTIS US LLC
(SANOFI AVENTIS)	(SANOFI AVENTIS US)
SANOFI INC	SANOFI AVENTIS US LLC
(SANOFI)	(SANOFI AVENTIS US)
SANOFI SYNTHELABO INC	SANOFI AVENTIS US LLC
(SANOFI SYNTHELABO)	(SANOFI AVENTIS US)
SANOFI SYNTHELABO RESEARCH DIV SANOFI	SANOFI AVENTIS US LLC
SYNTHELABO INC	
(SANOFI SYN RES)	(SANOFI AVENTIS US)
STERIS LABORATORIES INC	WATSON LABORATORIES INC
(STERIS)	(WATSON)
TRIGEN LABORATORIES INC	JUBILANT PHARMACEUTICALS INC
(TRIGEN)	(JUBILANT PHARMS)
UCB PHARMA INC	UCB INC
(UCB PHARMA)	(UCB INC)

1.4 AVAILABILITY OF THE EDITION

Since 1997, the Electronic Orange Book Query (EOBQ) <http://www.fda.gov/cder/ob/default.htm>, has been available on the internet and has become the updated-every-month Orange Book. The Query provides searching of the approved drug list by active ingredient, proprietary name, applicant holder, applicant number or patent number. Product search categories are: prescription, over-the-counter, discontinued drugs. There are links to patent and exclusivity information that may be applicable to each product.

Commencing with the 25th edition, the Annual Edition and monthly Cumulative Supplements have been provided in downloadable Portable Document Format (PDF) at the EOB home page by clicking on Annual Edition. The PDF annual and cumulative supplements duplicate previous paper versions. Over time, there will be an archive for the annuals and each year's December Cumulative Supplement.

The downloaded Annual Edition and Cumulative Supplements are also available in a paper version (Approved Drug Products with Therapeutic Equivalence Evaluations, ADP) from the U.S. Government Printing Office: <http://www.bookstore.gpo.gov/>; toll free 866-512-1800.

There are historical lists of Orange Book cumulative supplement product monthly changes at <http://www.fda.gov/cder/rxotcdpl/pdplarchive.htm>

There are ASCII text files of the Orange Book drug product, patent, and exclusivity data at <http://www.fda.gov/cder/orange/obreadme.htm>. The drug product text files are provided in eobzip.exe and eobzip.zip format. The files are updated concurrently with the monthly cumulative supplements. The annual Orange Book Edition Appendices A, B, and C in PDF format are updated quarterly.

Effective August 18, 2003, patent submissions for publication in the Orange Book and Docket *95S-0117 need to be submitted on form FDA-3542 which may be downloaded from the FDA Forms List, <http://www.fda.gov/opacom/morechoices/fdaforms/default.html>.

The current listing of the Orphan Product Designations and Approvals is available at <http://www.fda.gov/orphan/designat/list.htm>.

1.5 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 2004) 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>MAR 2006</u>	<u>JUN 2006</u>	<u>SEPT 2006</u>	<u>DEC 2006</u>
DRUG PRODUCTS LISTED	11487	11636	11741	11896
SINGLE SOURCE	2461 (21.4%)	2447 (21.0%)	2456 (20.9%)	2471 (20.8%)
MULTISOURCE	8937 (77.8%)	9000 (78.2%)	9196 (78.3%)	9336 (78.5%)
THERAPEUTICALLY EQUIVALENT	8730 (76.0%)	8900 (76.5%)	8997 (76.6%)	9139 (76.8%)
NOT THERAPEUTICALLY EQUIVALENT	207 (1.8%)	200 (1.7%)	199 (1.7%)	197 (1.7%)
EXCEPTIONS ¹	89 (0.8%)	89 (0.8%)	89 (0.8%)	89 (0.7%)
NEW MOLECULAR ENTITIES APPROVED	6	8	1	10
NUMBER OF APPLICANTS	629	642	649	666

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

1.6 ZOCOR (SIMVASTATIN) PATENT RELISTING

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

1.7 LEVOTHYROXINE SODIUM

The Description of Special Situations, Levothyroxine Sodium, published in the 26th Annual Edition of the Orange Book, has been modified in the Cumulative Supplement to include information on a supplemental approval for Genpharm ANDA 76752 approved in 2006. The full discussion as published in the 26th Annual Edition is repeated in the Cumulative Supplement and includes recent approval information on Levothyroxine Sodium.

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

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

Levo-T (Alara NDA 021342), Levothyroxine Sodium (Mylan ANDA 76187), Unithroid (Jerome Stevens NDA 021210) and Levothyroxine Sodium (Genpharm

ANDA 76752) tablets have been determined to be therapeutically equivalent to corresponding strengths of Synthroid (Abbott NDA 021402) tablets.

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

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

Novothyrox (Genpharm NDA 021292) requires further investigation and review to establish therapeutic equivalence to corresponding strengths of any other Levothyroxine Sodium drug products and is rated BX.

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

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

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

*Revised September 2006 **Revised December 2006

1.8 CUMULATIVE SUPPLEMENT LEGEND

The List is sorted by Ingredient(s) and, within each grouping, by the Dosage Form; Route and then by trade name.

The individual product record contains the Therapeutic Equivalence Code, Reference Listed Drug symbol, applicant holder, strength(s), New Drug Application number, product number, and approval date. The last two columns describe the action. The Action Month is the CS month the action occurred. The OB Action is the type of change that has occurred.

New ingredient(s), new dosage form; route(s), new trade names, and new product additions are preceded by >A> during the action month. The change month is the current CS month; the change code for new approvals is NEWA. Following months will display the same information without the >A>.

Changes to currently listed products will list two records. The deleted product record will be preceded by >D>. The product record change addition being made will be preceded by >A>. Following months will display only the >A> record without the >A>. All changes that occur to the product through the Annual year will be listed. The change month and change code will document the change.

The change code and description:

NEWA	New drug product approval usually in the supplement month.
CAHN	Applicant holder firm name has changed.
CAIN	Change. There has been a change in the Ingredient(s) name. All products will be deleted under the old name and all products will be added under the changed ingredient(s) name.
CDFR	Change. Dosage Form; Route of Administration.
CFTG	Change. A first time generic for the innovator product. A TE Code is added.
CMFD	Change. The product is moved from the Discontinued Section due to a change in marketing status.
CMS1	Change. Miscellaneous addition to list.
CMS2	Change. Miscellaneous deletion from list.
CPOT	Change. Potency amount/unit.
CRLD	Change. Reference Listed Drug.
CTEC	Change. Therapeutic Equivalence Code.
CTNA	Change. Trade Name.
DISC	Discontinued. The Rx or OTC listed product is not being marketed and will be moved to the discontinued section in the next edition.

PRESCRIPTION DRUG PRODUCT LIST - 26TH EDITION

RX DRUG PRODUCT LIST - CUMULATIVE SUPPLEMENT 12 - December 2006

1-1

ABARELIX

INJECTABLE; INTRAMUSCULAR

PLENAXIS

@ PRAECIS

100MG/VIAL

N21320 001 Nov 25, 2003 Sep DISC

ACEBUTOLOL HYDROCHLORIDE

CAPSULE; ORAL

SECTRAL

AB DR REDDYS LABS INC

EQ 200MG BASE

N18917 001 Dec 28, 1984 May CAHN

AB +

EQ 400MG BASE

N18917 003 Dec 28, 1984 May CAHN

ACETAMINOPHEN; BUTALBITAL

TABLET; ORAL

BUTAPAP

AB + MIKART

650MG;50MG

N89988 001 Oct 26, 1992 Jan CRLD

SEDAPAP

@ MAYRAND

650MG;50MG

N88944 001 Oct 17, 1985 Jan DISC

ACETAMINOPHEN; CAFFEINE; DIHYDROCODEINE BITARTRATE

TABLET; ORAL

ACETAMINOPHEN, CAFFEINE, AND DIHYDROCODEINE BITARTRATE

AB + MIKART

712.8MG;60MG;32MG

N40316 001 Apr 28, 1999 Sep CFTG

ACETAMINOPHEN, CAFFEINE, AND DIHYDROCODEINE BITATRATE

AB WEST WARD

712.8MG;60MG;32MG

N40637 001 Sep 22, 2006 Sep NEWA

ACETAMINOPHEN; CODEINE PHOSPHATE

SOLUTION; ORAL

ACETAMINOPHEN AND CODEINE PHOSPHATE

AA + ACTAVIS MID ATLANTIC

120MG/5ML;12MG/5ML

N85861 001

Jul CAHN

@ CLONMEL

120MG/5ML;12MG/5ML

N40098 001

Sep 20, 1996 Jan DISC

SUSPENSION; ORAL

CAPITAL AND CODEINE

AA ACTAVIS MID ATLANTIC

120MG/5ML;12MG/5ML

N85883 001

Jul CAHN

TABLET; ORAL

ACETAMINOPHEN AND CODEINE PHOSPHATE

AA + TEVA

300MG;60MG

N88629 001

Mar 06, 1985 Apr CRLD

ACETAMINOPHEN W/ CODEINE NO. 3

@ ROXANE

300MG;30MG

N84656 001

Jul DISC

ACETAMINOPHEN; HYDROCODONE BITARTRATE

TABLET; ORAL

HYDROCODONE BITARTRATE AND ACETAMINOPHEN

@ ENDO PHARMS

500MG;7.5MG

N40280 001

Sep 30, 1998 Feb DISC

@

650MG;7.5MG

N40280 002

Sep 30, 1998 Feb DISC

@

650MG;10MG

N40280 003

Sep 30, 1998 Feb DISC

@

750MG;7.5MG

N40281 002

Sep 30, 1998 Feb DISC

AA INTERPHARM

325MG;5MG

N40736 001

Aug 25, 2006 Aug NEWA

AA

325MG;10MG

N40746 001

Aug 25, 2006 Aug NEWA

AA

500MG;5MG

N40729 001

Aug 25, 2006 Aug NEWA

AA

500MG;7.5MG

N40748 001

Aug 25, 2006 Aug NEWA

AA

650MG;7.5MG

N40754 001

Aug 25, 2006 Aug NEWA

AA

650MG;10MG

N40757 001

Aug 25, 2006 Aug NEWA

AA

750MG;7.5MG

N40769 001

Aug 28, 2006 Aug NEWA

TABLET; ORAL

HYDROCODONE BITARTRATE AND ACETAMINOPHEN

	MIKART	300MG;5MG	N40658	001	Jan 19, 2006	Jan	NEWA
	+	300MG;7.5MG	N40556	002	Mar 24, 2006	Mar	NEWA
AA	VINTAGE PHARMS	325MG;5MG	N40655	001	Jan 19, 2006	Jan	NEWA
AA		325MG;7.5MG	N40656	001	Jan 19, 2006	Jan	NEWA
	HY-PHEN						
	@ ASCHER	500MG;5MG	N87677	001	May 03, 1982	Mar	DISC
	ZYDONE						
	+	ENDO PHARMS 400MG;5MG	N40288	001	Nov 27, 1998	May	CTNA
	+	400MG;7.5MG	N40288	002	Nov 27, 1998	May	CTNA
	+	400MG;10MG	N40288	003	Nov 27, 1998	May	CTNA

ACETAMINOPHEN; OXYCODONE HYDROCHLORIDE

SOLUTION; ORAL

OXYCODONE AND ACETAMINOPHEN

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

TABLET; ORAL

OXYCODONE AND ACETAMINOPHEN

	+	MIKART 400MG;2.5MG	N40679	001	May 16, 2006	May	NEWA
	+	400MG;5MG	N40687	001	Apr 27, 2006	Apr	NEWA
	+	400MG;7.5MG	N40698	001	Apr 27, 2006	Apr	NEWA
	+	400MG;10MG	N40692	001	Apr 27, 2006	Apr	NEWA
		500MG;10MG	N40676	001	Apr 19, 2006	Apr	NEWA

ACETAMINOPHEN; PROPOXYPHENE NAPSYLATE

TABLET; ORAL

PROPOXYPHENE NAPSYLATE AND ACETAMINOPHEN

AB	ACTAVIS ELIZABETH	650MG;100MG	N70910	001	Jan 02, 1987	Jun	CAHN
	CORNERSTONE	325MG;100MG	N76743	001	May 07, 2004	Jul	CAHN
AB		500MG;100MG	N76750	001	Jun 28, 2004	Jul	CAHN

ACETAMINOPHEN; TRAMADOL HYDROCHLORIDE

TABLET; ORAL

TRAMADOL HYDROCHLORIDE AND ACETAMINOPHEN

AB	BARR	325MG;37.5MG	N76914	001	Jul 26, 2006	Jul	NEWA
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ACETAZOLAMIDE

TABLET; ORAL

ACETAZOLAMIDE

AB	+	TARO 250MG	N40195	002	May 28, 1997	Mar	CRLD
	DIAMOX						
	@ DURAMED PHARMS BARR	125MG	N08943	001		Mar	DISC
	@	250MG	N08943	002		Mar	DISC

ACETIC ACID, GLACIAL

SOLUTION/DROPS; OTIC

ACETASOL

AT	ACTAVIS MID ATLANTIC	2%	N87146	001		Jul	CAHN
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ACETIC ACID, GLACIAL; HYDROCORTISONE

SOLUTION/DROPS; OTIC

ACETASOL HC

AT	ACTAVIS MID ATLANTIC	2%;1%	N87143	001	Jan 13, 1982	Jul	CAHN
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HYDROCORTISONE AND ACETIC ACID

AT	VINTAGE	2%;1%	N40609	001	Feb 06, 2006	Jan	NEWA
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ACETYLCHOLINE CHLORIDE

FOR SOLUTION; OPHTHALMIC

MIOCHOL

@	NOVARTIS	20MG/VIAL	N16211	001		Sep	DISC
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ACYCLOVIR

CAPSULE; ORAL

ACYCLOVIR

AB	ACTAVIS ELIZABETH	200MG	N74906	001	Aug 26, 1997	Jun	CAHN
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AB	CLONMEL HLTHCARE	200MG	N74833	001	Apr 22, 1997	May	CAHN
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SUSPENSION; ORAL

ACYCLOVIR

AB	ACTAVIS MID ATLANTIC	200MG/5ML	N74738	001	Apr 28, 1997	Jul	CAHN
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TABLET; ORAL

ACYCLOVIR

AB	ACTAVIS ELIZABETH	400MG	N74870	001	Jun 05, 1997	Jun	CAHN
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AB		800MG	N74870	002	Jun 05, 1997	Jun	CAHN
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AB	CLONMEL HLTHCARE	400MG	N74946	001	Nov 19, 1997	May	CAHN
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AB		800MG	N74946	002	Nov 19, 1997	May	CAHN
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ACYCLOVIR SODIUM

INJECTABLE; INJECTION

ACYCLOVIR SODIUM

@	HOSPIRA	EQ 500MG BASE/VIAL	N74663	001	Apr 22, 1997	Jun	DISC
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@		EQ 1GM BASE/VIAL	N74663	002	Apr 22, 1997	Jun	DISC
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ALBUTEROL

AEROSOL, METERED; INHALATION

ALBUTEROL

@	GENPHARM	0.09MG/INH	N73045	001	Aug 19, 1997	Feb	DISC
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@	PLIVA	0.09MG/INH	N74072	001	Aug 01, 1996	Feb	DISC
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ALBUTEROL SULFATE

SOLUTION; INHALATION

ALBUTEROL SULFATE

AN	ACTAVIS MID ATLANTIC	EQ 0.083% BASE	N73533	001	Sep 26, 1995	Jul	CAHN
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AN	RXELITE	EQ 0.083% BASE	N77569	001	Apr 04, 2006	Mar	NEWA
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SYRUP; ORAL

ALBUTEROL SULFATE

AA	ACTAVIS MID ATLANTIC	EQ 2MG BASE/5ML	N74454	001	Sep 25, 1995	Jul	CAHN
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AA		EQ 2MG BASE/5ML	N75262	001	Mar 30, 1999	Jul	CAHN
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>A>	AA	VINTAGE	EQ 2MG BASE/5ML	N78105	001	Dec 27, 2006	Dec	NEWA
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TABLET, EXTENDED RELEASE; ORAL

VOSPIRE ER

	DAVA PHARMS INC	EQ 4MG BASE	N76130	002	Sep 26, 2002	Sep	CAHN
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+		EQ 8MG BASE	N76130	003	Sep 26, 2002	Sep	CAHN
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ALBUTEROL SULFATE; IPRATROPIUM BROMIDE

SOLUTION; INHALATION

>A> ALBUTEROL SULFATE AND IPRATROPIUM BROMIDE

>A> AN SANDOZ EQ 0.083% BASE;0.017% N76867 001 Dec 21, 2006 Dec NEWA

DUONEB

>D> + DEY EQ 0.083% BASE;0.017% N20950 001 Mar 21, 2001 Dec CFTG

>A> AN + EQ 0.083% BASE;0.017% N20950 001 Mar 21, 2001 Dec CFTG

ALITRETINOIN

GEL; TOPICAL

PANRETIN

+ EISAI MEDCL RES EQ 0.1% BASE N20886 001 Feb 02, 1999 Oct CAHN

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

INJECTABLE; INJECTION

INFUVITE ADULT

+ SANDOZ 2 IU/ML;40MG/ML;12UGM/ML;40 IU/ML;1UGM/ML;3MG/ML;120UGM/ML;8MG/ML;1.2MG/ML;0.72MG/ML;1.2MG/ML;660 IU/ML;0.03MG/ML N21163 001 May 18, 2000 Jan CAHN

INJECTABLE; IV (INFUSION)

INFUVITE ADULT

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

ALPRAZOLAM

TABLET; ORAL

ALPRAZOLAM

AB ACTAVIS ELIZABETH 0.25MG N74342 001 Oct 31, 1993 Jun CAHN

AB 0.5MG N74342 002 Oct 31, 1993 Jun CAHN

AB 1MG N74342 003 Oct 31, 1993 Jun CAHN

AB 2MG N74342 004 Oct 31, 1993 Jun CAHN

TABLET, EXTENDED RELEASE; ORAL

ALPRAZOLAM

AB BARR 0.5MG N77725 001 Jul 31, 2006 Jul NEWA

AB 1MG N77725 002 Jul 31, 2006 Jul NEWA

AB 2MG N77725 004 Jul 31, 2006 Aug NEWA

AB 3MG N77725 003 Jul 31, 2006 Jul NEWA

AB MYLAN 0.5MG N77391 002 Jan 26, 2006 Jan NEWA

AB 1MG N77391 003 Jan 26, 2006 Jan NEWA

AB 2MG N77391 004 Jan 26, 2006 Jan NEWA

AB 3MG N77391 001 Jan 26, 2006 Jan NEWA

AB SANDOZ 0.5MG N77777 001 Jun 30, 2006 Jun NEWA

AB 1MG N77777 002 Jun 30, 2006 Jun NEWA

AB 2MG N77777 003 Jun 30, 2006 Jun NEWA

AB 3MG N77777 004 Jun 30, 2006 Jun NEWA

XANAX XR

AB PHARMACIA AND UPJOHN 0.5MG N21434 001 Jan 17, 2003 Jan CFTG

AB 1MG N21434 002 Jan 17, 2003 Jan CFTG

AB 2MG N21434 003 Jan 17, 2003 Jan CFTG

AB + 3MG N21434 004 Jan 17, 2003 Jan CFTG

AMANTADINE HYDROCHLORIDE

CAPSULE; ORAL

AMANTADINE HYDROCHLORIDE

AB		AMIDE PHARM	100MG	N77659 001	Feb 23, 2006	Feb	NEWA
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SYRUP; ORAL

AMANTADINE HYDROCHLORIDE

AA	+	ACTAVIS MID ATLANTIC	50MG/5ML	N72655 001	Oct 30, 1990	Jul	CAHN
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AA	+	VINTAGE	50MG/5ML	N77992 001	Dec 12, 2006	Nov	NEWA
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AMCINONIDE

CREAM; TOPICAL

AMCINONIDE

AB	+	ALTANA	0.1%	N76065 001	May 15, 2003	Sep	CRLD
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CYCLOCORT

@ ASTELLAS

0.1%

N18116 002		Sep	DISC
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LOTION; TOPICAL

AMCINONIDE

+	ALTANA	0.1%
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N76329 001	Nov 06, 2002	Sep	CRLD
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CYCLOCORT

@ ASTELLAS

0.1%

N19729 001	Jun 13, 1988	Sep	DISC
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OINTMENT; TOPICAL

AMCINONIDE

AB	+	ALTANA	0.1%	N76096 001	Nov 19, 2002	Sep	CRLD
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CYCLOCORT

@ ASTELLAS

0.1%

N18498 001		Sep	DISC
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AMILORIDE HYDROCHLORIDE

TABLET; ORAL

AMILORIDE HYDROCHLORIDE

+	PAR PHARM	5MG
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N70346 001	Jan 22, 1986	Jun	CRLD
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MIDAMOR

@ MERCK

5MG

N18200 001		Jun	DISC
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AMILORIDE HYDROCHLORIDE; HYDROCHLOROTHIAZIDE

TABLET; ORAL

AMILORIDE HYDROCHLORIDE AND HYDROCHLOROTHIAZIDE

AB	+	MYLAN	EQ 5MG ANHYDROUS;50MG	N73209 001	Oct 31, 1991	Jun	CRLD
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MODURETIC 5-50

@ MERCK

EQ 5MG ANHYDROUS;50MG

N18201 001		Jun	DISC
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AMINOPHYLLINE

SOLUTION; ORAL

AMINOPHYLLINE DYE FREE

AA	+	ACTAVIS MID ATLANTIC	105MG/5ML	N87727 001	Apr 16, 1982	Jul	CAHN
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AMIODARONE HYDROCHLORIDE

INJECTABLE; INJECTION

AMIODARONE HYDROCHLORIDE

AP	+	AKORN	50MG/ML	N76232 001	Jul 05, 2006	Jun	NEWA
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>D>	AP	+	APOTEX INC	50MG/ML	N77161 001	Apr 20, 2005	Dec	CAHN
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>A>	AP	+	GLAND PHARMA LTD	50MG/ML	N77161 001	Apr 20, 2005	Dec	CAHN
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AMITRIPTYLINE HYDROCHLORIDE; PERPHENAZINE

TABLET; ORAL

PERPHENAZINE AND AMITRIPTYLINE HYDROCHLORIDE

	MYLAN	10MG;2MG	N70336	001	Nov 10, 1988	Oct	CTEC
		10MG;4MG	N71442	001	Nov 10, 1988	Oct	CTEC
		25MG;2MG	N70337	001	Nov 10, 1988	Oct	CTEC
+		25MG;4MG	N70338	001	Nov 10, 1988	Oct	CTEC
+		50MG;4MG	N71443	001	Nov 10, 1988	Oct	CTEC
	@ SANDOZ	10MG;2MG	N71062	001	Nov 27, 1987	Oct	DISC
	@	10MG;4MG	N71862	001	Dec 21, 1987	Oct	DISC
	@	25MG;2MG	N71063	001	Nov 27, 1987	Oct	DISC
	@	25MG;4MG	N71064	001	Nov 27, 1987	Oct	DISC
	@	50MG;4MG	N71863	001	Dec 21, 1987	Oct	DISC
	@ WATSON LABS	10MG;2MG	N73007	001	Oct 17, 1991	Oct	DISC
	@	10MG;4MG	N73009	001	Oct 17, 1991	Oct	DISC
	@	25MG;2MG	N73008	001	Oct 17, 1991	Oct	DISC
	@	25MG;4MG	N73010	001	Oct 17, 1991	Oct	DISC

AMLODIPINE BESYLATE; BENAZEPRIL HYDROCHLORIDE

CAPSULE; ORAL

LOTREL

	NOVARTIS	EQ 5MG BASE;40MG	N20364	007	Apr 11, 2006	Apr	NEWA
		EQ 10MG BASE;20MG	N20364	005	Jun 20, 2002	Apr	CRLD
+		EQ 10MG BASE;40MG	N20364	006	Apr 11, 2006	Apr	NEWA

AMMONIUM LACTATE

CREAM; TOPICAL

AMMONIUM LACTATE

AB	PADDOCK	EQ 12% BASE	N76829	001	Feb 07, 2006	Jan	NEWA
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AMOXICILLIN

CAPSULE; ORAL

AMOXICILLIN

AB	AM ANTIBIOTICS	250MG	N62058	001		Jan	CAHN
AB		500MG	N62058	002		Jan	CAHN

FOR SUSPENSION; ORAL

AMOXICILLIN

	AB	AM ANTIBIOTICS	125MG/5ML	N62059	001		Jan	CAHN
	AB		250MG/5ML	N62059	002		Jan	CAHN
>A>	AB	AUROBINDO	200MG/5ML	N65334	001	Dec 28, 2006	Dec	NEWA
>A>	AB		400MG/5ML	N65334	002	Dec 28, 2006	Dec	NEWA
	AB	HIKMA	125MG/5ML	N65322	002	Jun 19, 2006	Jun	NEWA
	AB		200MG/5ML	N65325	002	Jun 19, 2006	Jun	NEWA
	AB		250MG/5ML	N65322	001	Jun 19, 2006	Jun	NEWA
	AB		400MG/5ML	N65325	001	Jun 19, 2006	Jun	NEWA

TABLET; ORAL

AMOXICILLIN

AB	HIKMA	875MG	N65255	001	Mar 29, 2006	Mar	NEWA
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AMPHETAMINE ASPARTATE; AMPHETAMINE SULFATE; DEXTROAMPHETAMINE SACCHARATE; DEXTROAMPHETAMINE SULFATE

TABLET; ORAL

DEXTROAMP SACCHARATE, AMP ASPARTATE, DEXTROAMP SULFATE AND AMP SULFATE

AB	TEVA PHARMS	1.25MG;1.25MG;1.25MG;1.25MG	N40472	001	Sep 30, 2003	May	CAHN
AB		2.5MG;2.5MG;2.5MG;2.5MG	N40472	002	Sep 30, 2003	May	CAHN

TABLET; ORAL

DEXTROAMP SACCHARATE, AMP ASPARTATE, DEXTROAMP SULFATE AND AMP SULFATE

AB	TEVA PHARMS	5MG;5MG;5MG;5MG	N40472 003	Sep 30, 2003	May	CAHN
AB		7.5MG;7.5MG;7.5MG;7.5MG	N40472 004	Sep 30, 2003	May	CAHN

AMPHOTERICIN B

INJECTABLE; INJECTION

AMPHOTERICIN B

AP	X GEN PHARMS	50MG/VIAL	N63206 001	Apr 29, 1992	Jun	CAHN
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LOTION; TOPICAL

FUNGIZONE

@	APOTHECON	3%	N60570 001		Nov	DISC
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AMPICILLIN SODIUM; SULBACTAM SODIUM

INJECTABLE; INJECTION

AMPICILLIN AND SULBACTAM

AP	INSTITUTO BIOCHIMICO	EQ 10GM BASE/VIAL;EQ 5GM BASE/VIAL	N65314 001	Nov 27, 2006	Nov	NEWA
AP	SANDOZ	EQ 1GM BASE/VIAL;EQ 500MG BASE/VIAL	N65241 001	Jul 25, 2006	Jul	NEWA
AP		EQ 1GM BASE/VIAL;EQ 500MG BASE/VIAL	N65310 001	Jul 25, 2006	Jul	NEWA
AP		EQ 2GM BASE/VIAL;EQ 1GM BASE/VIAL	N65241 002	Jul 25, 2006	Jul	NEWA
AP		EQ 2GM BASE/VIAL;EQ 1GM BASE/VIAL	N65310 002	Jul 25, 2006	Jul	NEWA
AP		EQ 10GM BASE/VIAL;EQ 5GM BASE/VIAL	N65240 001	Jul 25, 2006	Jul	NEWA

AMPICILLIN/AMPICILLIN TRIHYDRATE

CAPSULE; ORAL

AMPICILLIN TRIHYDRATE

@ AM ANTIBIOTICS

		EQ 250MG BASE	N61602 001		Jan	CAHN
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		EQ 500MG BASE	N61602 002		Jan	CAHN
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>D>	AB	CLONMEL	EQ 250MG BASE	N62883 001	Feb 25, 1988	Dec	CTEC
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>A>			EQ 250MG BASE	N62883 001	Feb 25, 1988	Dec	CTEC
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>D>	AB		EQ 500MG BASE	N62882 001	Feb 25, 1988	Dec	CRLD
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>A>	+		EQ 500MG BASE	N62882 001	Feb 25, 1988	Dec	CRLD
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>D>		PRINCIPEN					
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>D>	AB	APOTHECON	EQ 250MG BASE	N61392 001		Dec	DISC
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>A>		@	EQ 250MG BASE	N61392 001		Dec	DISC
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		@	EQ 250MG BASE	N62888 001	Mar 04, 1988	Nov	DISC
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>D>	AB	+	EQ 500MG BASE	N61392 002		Dec	DISC
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>A>		@	EQ 500MG BASE	N61392 002		Dec	DISC
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	AB	+	EQ 500MG BASE	N61392 002		Nov	CRLD
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		@	EQ 500MG BASE	N62888 002	Mar 04, 1988	Nov	DISC
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FOR SUSPENSION; ORAL

AMPICILLIN TRIHYDRATE

@ AM ANTIBIOTICS

		EQ 125MG BASE/5ML	N61601 001		Jan	CAHN
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		EQ 250MG BASE/5ML	N61601 002		Jan	CAHN
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PRINCIPEN

>D>		APOTHECON	EQ 100MG BASE/ML	N61394 001		Dec	DISC
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>A>		@	EQ 100MG BASE/ML	N61394 001		Dec	DISC
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ANAGRELIDE HYDROCHLORIDE

CAPSULE; ORAL

ANAGRELIDE HYDROCHLORIDE

AB	ALPHAPHARM	EQ 0.5MG BASE	N77613 001	Jun 27, 2006	Jun	NEWA
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AB		EQ 1MG BASE	N77613 002	Jun 27, 2006	Jun	NEWA
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ANIDULAFUNGIN

INJECTABLE; IV (INFUSION)

ERAXIS

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

ANISINDIONE

TABLET; ORAL

MIRADON

@	SCHERING	50MG	N10909 003		Jan	DISC
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APOMORPHINE HYDROCHLORIDE

INJECTABLE; SUBCUTANEOUS

APOKYN

@	VERNALIS	20MG/2ML (10MG/ML)	N21264 001	Apr 20, 2004	Oct	DISC
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APREPITANT

CAPSULE; ORAL

EMEND

	MERCK	40MG	N21549 003	Jun 30, 2006	Jun	NEWA
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ARFORMOTEROL TARTRATE

SOLUTION; INHALATION

BROVANA

+	SEPRACOR	EQ 0.015MG BASE/2ML	N21912 001	Oct 06, 2006	Oct	NEWA
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ARIPIRAZOLE

INJECTABLE; INTRAMUSCULAR

ABILIFY

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

ABILIFY

	OTSUKA	2MG	N21436 006	Nov 15, 2002	Jul	CMFD
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+		10MG	N21436 001	Nov 15, 2002	Sep	CRLD
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TABLET, ORALLY DISINTEGRATING; ORAL

ABILIFY

	OTSUKA	10MG	N21729 002	Jun 07, 2006	Jun	NEWA
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		15MG	N21729 003	Jun 07, 2006	Jun	NEWA
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		20MG	N21729 004	Jun 07, 2006	Jun	NEWA
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+		30MG	N21729 005	Jun 07, 2006	Jun	NEWA
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ARTICAINE HYDROCHLORIDE; EPINEPHRINE

INJECTABLE; INJECTION

SEPTOCAINE

	DEPROCO	4%;EQ 0.005MG BASE/ML	N22010 001	Mar 30, 2006	Mar	NEWA
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+		4%; EQ 0.017MG BASE/1.7ML (4%; EQ 0.01MG BASE/ML)	N20971 001	Apr 03, 2000	Mar	CPOT
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ARTICAINE HYDROCHLORIDE; EPINEPHRINE BITARTRATE

INJECTABLE; INJECTION

SEPTOCAINE

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

INJECTABLE; IV (INFUSION)

INFUVITE PEDIATRIC

+	SANDOZ	80MG/VIAL;0.02MG/VIAL;400 IU/VIAL;0.001MG/VIAL;5MG/VIAL;0.1 4MG/VIAL;17MG/VIAL;1MG/VIAL;1.4MG /VIAL;1.2MG/VIAL;7 IU/VIAL;2,300 IU/VIAL;0.2MG/VIAL	N21265 001	Feb 21, 2001	Jan	CAHN
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INFUVITE PEDIATRIC (PHARMACY BULK PACKAGE)

+	SANDOZ	80MG/VIAL;0.02MG/VIAL;400 IU/VIAL;0.001MG/VIAL;5MG/VIAL;0.1 4MG/VIAL;17MG/VIAL;1MG/VIAL;1.4MG /VIAL;1.2MG/VIAL;7 IU/VIAL;2,300 IU/VIAL;0.2MG/VIAL	N21646 001	Jan 29, 2004	Jan	CAHN
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ASCORBIC ACID; POLYETHYLENE GLYCOL 3350; POTASSIUM CHLORIDE; SODIUM ASCORBATE; SODIUM
 CHLORIDE; SODIUM SULFATE

FOR SOLUTION; ORAL

MOVIPREP

+	NORGINE B V	4.7GM;100GM;1.015GM;5.9GM;2.691GM ;7.5GM	N21881 001	Aug 02, 2006	Aug	NEWA
+	SALIX PHARMS	4.7GM;100GM;1.015GM;5.9GM;2.691GM ;7.5GM	N21881 001	Aug 02, 2006	Sep	CAHN

ASPIRIN; BUTALBITAL; CAFFEINE

TABLET; ORAL

ASPIRIN AND CAFFEINE W/ BUTALBITAL

AB	ACTAVIS ELIZABETH	325MG;50MG;40MG	N86710 002	Aug 23, 1983	Jun	CAHN
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ASPIRIN; BUTALBITAL; CAFFEINE; CODEINE PHOSPHATE

CAPSULE; ORAL

BUTALBITAL, ASPIRIN, CAFFEINE, AND CODEINE PHOSPHATE

@	ENDO PHARMS	325MG;50MG;40MG;30MG	N75351 001	Mar 05, 1999	Feb	DISC
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FIORINAL W/CODEINE

AB	+	WATSON PHARMS	325MG;50MG;40MG;30MG	N19429 003	Oct 26, 1990	Apr	CTNA
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ASPIRIN; OXYCODONE HYDROCHLORIDE

TABLET; ORAL

ASPIRIN AND OXYCODONE

+	ENDO PHARMS	325MG;4.8355MG	N07337 007	Aug 05, 2005	Jun	NEWA
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ASPIRIN; OXYCODONE HYDROCHLORIDE; OXYCODONE TEREPHTHALATE

TABLET; ORAL

PERCODAN-DEMI

@	ENDO PHARMS	325MG;2.25MG;0.19MG	N07337 005		Feb	DISC
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ATAZANAVIR SULFATE

CAPSULE; ORAL

REYATAZ

BRISTOL MYERS SQUIBB EQ 200MG BASE

+		EQ 300MG BASE	N21567 003	Jun 20, 2003	Oct	CRLD
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+			N21567 004	Oct 16, 2006	Oct	NEWA
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ATENOLOL

TABLET; ORAL

ATENOLOL

>A>	AB	OHM LABS	25MG	N77877 001	Dec 27, 2006	Dec	NEWA
>A>	AB		50MG	N77877 002	Dec 27, 2006	Dec	NEWA
>A>	AB		100MG	N77877 003	Dec 27, 2006	Dec	NEWA

TABLET; ORALATENOLOL

AB	UNIQUE PHARM LABS	25MG	N77443 001	Sep 13, 2006	Aug	NEWA
AB		50MG	N77443 002	Sep 13, 2006	Aug	NEWA
AB		100MG	N77443 003	Sep 13, 2006	Aug	NEWA

ATOVAQUONETABLET; ORALMEPRON

@ GLAXOSMITHKLINE	250MG	N20259 001	Nov 25, 1992	May	DISC
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ATROPINE; PRALIDOXIME CHLORIDEINJECTABLE; INTRAMUSCULARDUODOTE

+	MERIDIAN MEDCL	2.1MG/0.7ML;600MG/2ML	N21983 001	Sep 28, 2006	Sep	NEWA
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AZITHROMYCINFOR SUSPENSION; ORALAZITHROMYCIN

AB	PLIVA	EQ 100MG BASE/5ML	N65246 002	Jul 05, 2006	Jun	NEWA
AB		EQ 200MG BASE/5ML	N65246 001	Jul 05, 2006	Jun	NEWA
AB	SANDOZ	EQ 100MG BASE/5ML	N65297 001	Sep 18, 2006	Sep	NEWA
AB		EQ 200MG BASE/5ML	N65297 002	Sep 18, 2006	Sep	NEWA

ZITHROMAX

AB	PFIZER	EQ 100MG BASE/5ML	N50710 001	Oct 19, 1995	Jun	CFTG
AB	+	EQ 200MG BASE/5ML	N50710 002	Oct 19, 1995	Jun	CFTG

TABLET; ORALAZITHROMYCIN

>A>	AB	MYLAN	EQ 600MG BASE	N65360 001	Jan 08, 2007	Dec	NEWA
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AZITHROMYCIN HYDROGENCITRATEINJECTABLE; INJECTIONAZITHROMYCIN

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

BACAMPICILLIN HYDROCHLORIDEFOR SUSPENSION; ORALSPECTROBID

@ PFIZER	125MG/5ML	N50556 001	Mar 23, 1982	Feb	DISC
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TABLET; ORALSPECTROBID

@ PFIZER	400MG	N50520 001		Feb	DISC
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BACLOFENTABLET; ORALBACLOFEN

AB	CARACO	10MG	N77984 001	Aug 14, 2006	Aug	NEWA
AB		20MG	N77862 002	Aug 14, 2006	Aug	NEWA

BENAZEPRIL HYDROCHLORIDETABLET; ORALBENAZEPRIL HYDROCHLORIDE

AB	APOTEX INC	5MG	N77128 001	Mar 08, 2006	Feb	NEWA
AB		10MG	N77128 002	Mar 08, 2006	Feb	NEWA

TABLET; ORAL

BENAZEPRIL HYDROCHLORIDE

AB	APOTEX INC	20MG	N77128 003	Mar 08, 2006	Feb	NEWA
AB		40MG	N77128 004	Mar 08, 2006	Feb	NEWA
AB	BIOKEY	5MG	N76820 001	Feb 03, 2006	Jan	NEWA
AB		10MG	N76820 002	Feb 03, 2006	Jan	NEWA
AB		20MG	N76820 003	Feb 03, 2006	Jan	NEWA
AB		40MG	N76820 004	Feb 03, 2006	Jan	NEWA

BENDROFLUMETHIAZIDE

TABLET; ORAL

NATURETIN-5

@ APOTHECON	5MG	N12164 002		Jun	DISC
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BENZPHETAMINE HYDROCHLORIDE

TABLET; ORAL

BENZPHETAMINE HYDROCHLORIDE

AA	PADDOCK	50MG	N40578 001	Apr 17, 2006	Apr	NEWA
AA	+ DIDREX					
AA	+ PHARMACIA AND UPJOHN	50MG	N12427 002		Apr	CFTG

BENZQUINAMIDE HYDROCHLORIDE

INJECTABLE; INJECTION

EMETE-CON

@ PFIZER	EQ 50MG BASE/VIAL	N16820 001		Mar	DISC
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BETAINE, ANHYDROUS

FOR SOLUTION; ORAL

CYSTADANE

+ JAZZ	1GM/SC00PFUL	N20576 001	Oct 25, 1996	Feb	CAHN
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BETAMETHASONE DIPROPIONATE

CREAM; TOPICAL

ALPHATREX

@ SAVAGE LABS	EQ 0.05% BASE	N19138 001	Jun 26, 1984	Jun	DISC
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BETAMETHASONE DIPROPIONATE

AB	ACTAVIS MID ATLANTIC	EQ 0.05% BASE	N70885 001	Feb 03, 1987	Jun	CAHN
AB	+ FOUGERA	EQ 0.05% BASE	N19137 001	Jun 26, 1984	Jun	CRLD

GEL, AUGMENTED; TOPICAL

BETAMETHASONE DIPROPIONATE

AB	+ ALTANA	EQ 0.05% BASE	N75276 001	May 13, 2003	Aug	CRLD
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DIPROLENE

@ SCHERING	EQ 0.05% BASE	N19408 002	Nov 22, 1991	Aug	DISC
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LOTION; TOPICAL

ALPHATREX

@ SAVAGE LABS	EQ 0.05% BASE	N70273 001	Aug 12, 1985	Jun	DISC
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BETAMETHASONE DIPROPIONATE

AB	ACTAVIS MID ATLANTIC	EQ 0.05% BASE	N70281 001	Jul 31, 1985	Jul	CAHN
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OINTMENT; TOPICAL

BETAMETHASONE DIPROPIONATE

AB	ACTAVIS MID ATLANTIC	EQ 0.05% BASE	N71012 001	Feb 03, 1987	Jun	CAHN
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OINTMENT, AUGMENTED; TOPICAL

BETAMETHASONE DIPROPIONATE

AB	ACTAVIS MID ATLANTIC	EQ 0.05% BASE	N74304 001	Aug 31, 1995	Jun	CAHN
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BETAMETHASONE DIPROPIONATE; CALCIPOTRIENE HYDRATE

OINTMENT; TOPICAL

TACLONEX

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

CREAM; TOPICAL

CLOTRIMAZOLE AND BETAMETHASONE DIPROPIONATE

AB	ACTAVIS MID ATLANTIC	EQ 0.05% BASE;1%	N76002	001	Aug 02, 2002	Jun	CAHN
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BETAMETHASONE VALERATE

CREAM; TOPICAL

VALNAC

AB	ACTAVIS MID ATLANTIC	EQ 0.1% BASE	N70050	001	Oct 10, 1984	Jun	CAHN
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LOTION; TOPICAL

BETAMETHASONE VALERATE

AB	ACTAVIS MID ATLANTIC	EQ 0.1% BASE	N70052	001	Jul 31, 1985	Jul	CAHN
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OINTMENT; TOPICAL

BETAMETHASONE VALERATE

AB	ACTAVIS MID ATLANTIC	EQ 0.1% BASE	N70051	001	Oct 10, 1984	Jun	CAHN
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BETHANECHOL CHLORIDE

TABLET; ORAL

BETHANECHOL CHLORIDE

AA	IMPAX LABS	5MG	N40739	001	Nov 01, 2006	Oct	NEWA
AA		10MG	N40741	001	Nov 01, 2006	Oct	NEWA
AA		25MG	N40740	001	Nov 01, 2006	Oct	NEWA
AA		50MG	N40721	004	Nov 01, 2006	Oct	NEWA

BEXAROTENE

CAPSULE; ORAL

TARGRETIN

+	EISAI MEDCL RES	75MG	N21055	001	Dec 29, 1999	Oct	CAHN
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GEL; TOPICAL

TARGRETIN

+	EISAI MEDCL RES	1%	N21056	001	Jun 28, 2000	Oct	CAHN
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BISKALCITRATE; METRONIDAZOLE; TETRACYCLINE

CAPSULE; ORAL

PYLERA

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

TABLET; ORAL

BISOPROLOL FUMARATE

>A>	AB	AUROBINDO PHARMA	5MG	N77910	001	Dec 27, 2006	Dec	NEWA
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>A>	AB		10MG	N77910	002	Dec 27, 2006	Dec	NEWA
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BISOPROLOL FUMARATE; HYDROCHLOROTHIAZIDE

TABLET; ORAL

BISOPROLOL FUMARATE AND HYDROCHLOROTHIAZIDE

AB	ACTAVIS ELIZABETH	2.5MG;6.25MG	N75672	001	Sep 25, 2000	Jun	CAHN
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AB		5MG;6.25MG	N75672	002	Sep 25, 2000	Jun	CAHN
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AB		10MG;6.25MG	N75672	003	Sep 25, 2000	Jun	CAHN
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BLEOMYCIN SULFATE

INJECTABLE; INJECTION

BLEOMYCIN SULFATE

@ SICOR PHARMS

EQ 15 UNITS BASE/VIAL

N64084 001 Jun 01, 1996 Jun DISC

@

EQ 30 UNITS BASE/VIAL

N64084 002 Jun 01, 1996 Jun DISC

BRIMONIDINE TARTRATE

SOLUTION/DROPS; OPHTHALMIC

ALPHAGAN P

AT + ALLERGAN 0.15%

N21262 001 Mar 16, 2001 May CTEC

BRIMONIDINE TARTRATE

AT AKORN 0.2%

N76439 001 Mar 14, 2006 Feb NEWA

AT ALCON RES 0.15%

N21764 001 May 22, 2006 May NEWA

BRINZOLAMIDE

SUSPENSION/DROPS; OPHTHALMIC

AZOPT

+ ALCON 1%

N20816 001 Apr 01, 1998 Feb CAHN

BROMODIPHENHYDRAMINE HYDROCHLORIDE; CODEINE PHOSPHATE

SYRUP; ORAL

AMBENYL

@ FOREST LABS

12.5MG/5ML;10MG/5ML

N09319 006 Jan 10, 1984 Aug DISC

MYBANIL

+ MORTON GROVE 12.5MG/5ML;10MG/5ML

N88626 001 Oct 12, 1984 Aug CRLD

BUDESONIDE

POWDER, METERED; INHALATION

BUDESONIDE

ASTRAZENECA 0.08MG/INH

N21949 001 Jul 12, 2006 Jul NEWA

+ 0.16MG/INH

N21949 002 Jul 12, 2006 Jul NEWA

SPRAY, METERED; NASAL

RHINOCORT

+ ASTRAZENECA 0.032MG/INH

N20746 001 Oct 01, 1999 Mar CRLD

@ 0.064MG/INH

N20746 002 Oct 01, 1999 Mar DISC

BUDESONIDE; FORMOTEROL FUMARATE

SPRAY, METERED; INHALATION

SYMBICORT

ASTRAZENECA 0.08MG/INH;EQ 0.045MG BASE

N21929 001 Jul 21, 2006 Jul NEWA

+ 0.016MG/INH;EQ 0.045MG BASE

N21929 002 Jul 21, 2006 Jul NEWA

BUMETANIDE

INJECTABLE; INJECTION

BUMETANIDE

AP + BEDFORD 0.25MG/ML

N74441 001 Jan 27, 1995 Feb CRLD

BUMEX

@ ROCHE 0.25MG/ML

N18226 001 Feb 28, 1983 Feb DISC

BUPIVACAINE HYDROCHLORIDE; EPINEPHRINE BITARTRATE

INJECTABLE; INJECTION

BUPIVACAINE HYDROCHLORIDE

+ HOSPIRA 0.5%;EQ 0.009MG BASE/ML

N22046 001 Jul 13, 1983 Aug CTNA

INJECTABLE; INJECTION

BUPIVACAINE HYDROCHLORIDE AND EPINEPHRINE							
AP	SEPTODONT	0.5%;0.0091MG/ML	N77250	001	Sep 27, 2006	Sep	NEWA
BUPIVACAINE HYDROCHLORIDE W/EPINEPHRINE							
AP	+ HOSPIRA	0.5%;0.0091MG/ML	N22046	001	Jul 13, 1983	Oct	CTEC
	+	0.5%;EQ 0.009MG BASE/ML	N22046	001	Jul 13, 1983	Apr	NEWA

BUPROPION HYDROCHLORIDE

TABLET; ORAL

BUPROPION HYDROCHLORIDE							
AB	APOTEX INC	75MG	N76143	001	Jan 17, 2006	Jan	NEWA
AB		100MG	N76143	002	Jan 17, 2006	Jan	NEWA

TABLET, EXTENDED RELEASE; ORAL

BUPROPION HYDROCHLORIDE								
>A>	AB3	ANCHEN PHARMS	150MG	N77284	001	Dec 14, 2006	Dec	NEWA
>A>	AB3		300MG	N77284	002	Dec 14, 2006	Dec	NEWA
>A>	AB3	IMPAX PHARMS	300MG	N77415	002	Dec 15, 2006	Dec	NEWA
WELLBUTRIN XL								
>D>		+ SMITHKLINE BEECHAM	150MG	N21515	001	Aug 28, 2003	Dec	CFTG
>A>	AB3	+	150MG	N21515	001	Aug 28, 2003	Dec	CFTG
>D>			300MG	N21515	002	Aug 28, 2003	Dec	CFTG
>A>	AB3		300MG	N21515	002	Aug 28, 2003	Dec	CFTG

BUSPIRONE HYDROCHLORIDE

TABLET; ORAL

BUSPIRONE HYDROCHLORIDE							
AB	ACTAVIS TOTOWA	5MG	N75388	001	May 09, 2002	Aug	CAHN
AB		10MG	N75388	002	May 09, 2002	Aug	CAHN
AB		15MG	N75388	003	May 09, 2002	Aug	CAHN

BUSULFAN

INJECTABLE; INJECTION

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

CAFFEINE CITRATE

SOLUTION; INTRAVENOUS

		CAFCIT							
AP	+	MEAD JOHNSON	EQ 30MG BASE/3ML (EQ 10MG BASE/ML)	N20793	001	Sep 21, 1999	Sep	CFTG	
		CAFFEINE CITRATE							
AP		PHARMAFORCE	EQ 30MG BASE/3ML (EQ 10MG BASE/ML)	N77233	001	Sep 21, 2006	Sep	NEWA	

SOLUTION; ORAL

		CAFCIT							
AA	+	MEAD JOHNSON	EQ 30MG BASE/3ML (EQ 10MG BASE/ML)	N20793	002	Apr 12, 2000	Sep	CFTG	
		CAFFEINE CITRATE							
AA		PHARMAFORCE	EQ 30MG BASE/3ML (EQ 10MG BASE/ML)	N77304	001	Sep 21, 2006	Sep	NEWA	

CAFFEINE; ERGOTAMINE TARTRATE

SUPPOSITORY; RECTAL

CAFERGOT								
	@	NOVARTIS	100MG;2MG	N09000	002		Feb	DISC
MIGERGOT								
+	G	AND W LABS	100MG;2MG	N86557	001	Oct 04, 1983	Feb	CRLD

CALCIPOTRIENECREAM; TOPICAL
DOVONEX

+ LEO PHARM 0.005% N20554 001 Jul 22, 1996 Feb CAHN

OINTMENT; TOPICAL
DOVONEX

+ LEO PHARM 0.005% N20273 001 Dec 29, 1993 Feb CAHN

SOLUTION; TOPICAL
DOVONEX

+ LEO PHARM 0.005% N20611 001 Mar 03, 1997 Feb CAHN

CALCITONIN SALMON RECOMBINANTSPRAY, METERED; NASAL
FORTICAL

+ UPSHER SMITH 200 IU/SPRAY N21406 001 Aug 12, 2005 Jun CAHN

CALCITONIN, SALMONINJECTABLE; INJECTION
MIACALCIN

+ NOVARTIS 200 IU/ML N17808 002 Mar 29, 1991 Jan CTEC

CALCITRIOLCAPSULE; ORAL
CALCITRIOL

AB ROXANE 0.25UGM N76917 001 Mar 27, 2006 Mar NEWA

INJECTABLE; INJECTION
CALCITRIOL

AP FRESENIUS MEDCL 0.001MG/ML N75766 001 Feb 20, 2003 Nov CAHN

AP 0.002MG/ML N75766 002 Feb 20, 2003 Nov CAHN

AP GENIX THERAP 0.001MG/ML N77102 001 Feb 08, 2006 Jan NEWA

CALCIUM ACETATECAPSULE; ORAL
PHOSLO

@ FRESENIUS MEDCL EQ 84.5MG CALCIUM N21160 001 Apr 02, 2001 Nov CAHN

@ EQ 169MG CALCIUM N21160 002 Apr 02, 2001 Nov CAHN

PHOSLO GELCAPS

+ FRESENIUS MEDCL EQ 169MG CALCIUM N21160 003 Apr 02, 2001 Nov CAHN

TABLET; ORAL
PHOSLO

@ FRESENIUS MEDCL EQ 169MG CALCIUM N19976 001 Dec 10, 1990 Nov CAHN

@ NABI EQ 169MG CALCIUM N19976 001 Dec 10, 1990 Jun DISC

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

INJECTABLE; INJECTION

PRISMASOL BGK 0/2.5 IN PLASTIC CONTAINER

+ GAMBRO RENAL PRODS 3.68GM/1000ML;20GM/1000ML;5.4GM/1000ML;3.05GM/1000ML;N/A/1000ML;3.09GM/1000ML;6.46GM/1000ML N21703 006 Oct 25, 2006 Oct NEWA

PRISMASOL BGK 2/0 IN PLASTIC CONTAINER

+ GAMBRO RENAL PRODS N/A/1000ML;20GM/1000ML;5.4GM/1000ML;2.03GM/1000ML;0.157GM/1000ML;3.09GM/1000ML;6.46GM/1000ML N21703 002 Oct 25, 2006 Oct NEWA

INJECTABLE; INJECTION

PRISMASOL BGK 2/3.5 IN PLASTIC CONTAINER

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

+	GAMBRO RENAL PRODS	N/A/1000ML;20GM/1000ML;5.4GM/1000ML;3.05GM/1000ML;0.314GM/1000ML;3.09GM/1000ML;6.46GM/1000ML	N21703 005	Oct 25, 2006	Oct	NEWA
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PRISMASOL BGK 4/2.5 IN PLASTIC CONTAINER

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

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

+	GAMBRO RENAL PRODS	5.15GM/1000ML;N/A/1000ML;5.4GM/1000ML;2.03GM/1000ML;N/A/1000ML;3.09GM/1000ML;6.46GM/1000ML	N21703 001	Oct 25, 2006	Oct	NEWA
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CALCIUM CHLORIDE; DEXTROSE; MAGNESIUM CHLORIDE; SODIUM CHLORIDE; SODIUM LACTATE

SOLUTION; INTRAPERITONEAL

DIANEAL LOW CALCIUM W/ DEXTROSE 2.5% IN PLASTIC CONTAINER

	BAXTER HLTHCARE	18.3MG/100ML;2.5GM/100ML;5.08MG/100ML;538MG/100ML;448MG/100ML	N20183 002	Dec 04, 1992	Nov	CTEC
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DIANEAL LOW CALCIUM W/ DEXTROSE 3.5% IN PLASTIC CONTAINER

	BAXTER HLTHCARE	18.3MG/100ML;3.5GM/100ML;5.08MG/100ML;538MG/100ML;448MG/100ML	N20183 003	Dec 04, 1992	Nov	CTEC
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DIANEAL LOW CALCIUM W/ DEXTROSE 4.25% IN PLASTIC CONTAINER

	BAXTER HLTHCARE	18.3MG/100ML;4.25GM/100ML;5.08MG/100ML;538MG/100ML;448MG/100ML	N20183 004	Dec 04, 1992	Nov	CTEC
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INPERSOL-LC/LM W/ DEXTROSE 3.5% IN PLASTIC CONTAINER

	FRESENIUS	18.4MG/100ML;3.5GM/100ML;5.08MG/100ML;538MG/100ML;448MG/100ML	N20374 003	Jun 13, 1994	Nov	CTEC
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CALCIUM CHLORIDE; MAGNESIUM CHLORIDE; POTASSIUM CHLORIDE; SODIUM ACETATE; SODIUM CHLORIDE; SODIUM CITRATE

SOLUTION; IRRIGATION

BALANCED SALT SOLUTION

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

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

TABLET; ORAL

ATACAND

	ASTRAZENECA	8MG	N20838 002	Jun 04, 1998	May	CRLD
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CAPTOPRIL

TABLET; ORAL

CAPTOPRIL

@	APOTHECON	12.5MG	N74472 001	Mar 31, 1995	Nov	DISC
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@		25MG	N74472 002	Mar 31, 1995	Nov	DISC
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@		50MG	N74472 003	Mar 31, 1995	Nov	DISC
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@		100MG	N74472 004	Mar 31, 1995	Nov	DISC
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@	CLONMEL HLTHCARE	12.5MG	N74423 001	Feb 13, 1996	Jan	DISC
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@		25MG	N74423 002	Feb 13, 1996	Jan	DISC
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@		50MG	N74423 003	Feb 13, 1996	Jan	DISC
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TABLET; ORAL

CAPTOPRIL

@ CLONMEL HLTHCARE	100MG	N74423 004	Feb 13, 1996	Jan	DISC
@ ENDO LABS	12.5MG	N74418 001	Feb 13, 1996	Feb	DISC
@	25MG	N74418 002	Feb 13, 1996	Feb	DISC
@	50MG	N74418 003	Feb 13, 1996	Feb	DISC
@	100MG	N74418 004	Feb 13, 1996	Feb	DISC

CAPTAPRIL; HYDROCHLOROTHIAZIDE

TABLET; ORAL

CAPTOPRIL AND HYDROCHLOROTHIAZIDE

@ ENDO LABS	25MG;15MG	N74788 001	Dec 29, 1997	Feb	DISC
@	25MG;25MG	N74788 002	Dec 29, 1997	Feb	DISC
@	50MG;15MG	N74788 004	Dec 29, 1997	Feb	DISC
@	50MG;25MG	N74788 003	Dec 29, 1997	Feb	DISC

CARBAMAZEPINE

TABLET, CHEWABLE; ORAL

CARBAMAZEPINE

AB	JUBILANT PHARMS	100MG	N71940 001	Feb 01, 1988	Jul	CAHN
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CARBIDOPA; LEVODOPA

TABLET; ORAL

CARBIDOPA AND LEVODOPA

AB	ACTAVIS ELIZABETH	10MG;100MG	N74260 001	Sep 03, 1993	Jun	CAHN
AB		25MG;100MG	N74260 002	Sep 03, 1993	Jun	CAHN
AB		25MG;250MG	N74260 003	Sep 03, 1993	Jun	CAHN

CARBOPLATIN

INJECTABLE; INJECTION

CARBOPLATIN

AP	WATSON LABS	50MG/VIAL	N77383 001	Jan 27, 2006	Jan	NEWA
AP		150MG/VIAL	N77383 002	Jan 27, 2006	Jan	NEWA
AP		450MG/VIAL	N77383 003	Jan 27, 2006	Jan	NEWA

INJECTABLE; IV (INFUSION)

CARBOPLATIN

	@ AM PHARM	EQ 50MG/5ML (10MG/ML)	N77247 001	Oct 21, 2004	Feb	DISC
AP		EQ 50MG/5ML (10MG/ML)	N77266 001	Feb 15, 2006	Jan	NEWA
	@	EQ 150MG/15ML (10MG/ML)	N77247 002	Oct 21, 2004	Feb	DISC
AP		EQ 150MG/15ML (10MG/ML)	N77266 002	Feb 15, 2006	Jan	NEWA
AP		EQ 450MG/45ML (10MG/ML)	N77266 003	Feb 15, 2006	Jan	NEWA
AP		EQ 600MG/60ML (10MG/ML)	N77266 004	Feb 15, 2006	Jan	NEWA
AP	BEDFORD LABS	EQ 600MG/60ML (10MG/ML)	N77244 004	Jan 20, 2006	Jan	NEWA
AP	DABUR ONCOLOGY PLC	EQ 450MG/45ML (10MG/ML)	N77432 003	Sep 29, 2006	Sep	NEWA
AP		EQ 50MG/5ML (10MG/ML)	N77432 001	Sep 29, 2006	Sep	NEWA
AP		EQ 150MG/15ML (10MG/ML)	N77432 002	Sep 29, 2006	Sep	NEWA

CARVEDILOL

TABLET; ORAL

COREG

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

CARVEDILOL PHOSPHATECAPSULE, EXTENDED RELEASE; ORAL
COREG CR

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

CASPOFUNGIN ACETATEINJECTABLE; IV (INFUSION)
CANCIDAS

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

CEFACLORTABLET, EXTENDED RELEASE; ORAL
CEFACLOR

AB	+	PAR PHARM	EQ 500MG BASE	N65057 001	Jan 05, 2001	May	CAHN
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CEFADROXIL/CEFADROXIL HEMIHYDRATE

CAPSULE; ORAL

CEFADROXIL

AB	+	IVAX PHARMS	EQ 500MG BASE	N62766 001	Mar 03, 1987	Mar	CRLD
AB		ORCHID HLTHCARE	EQ 500MG BASE	N65309 001	Sep 18, 2006	Sep	NEWA
AB		SANDOZ	EQ 500MG BASE	N62291 001		Aug	CAHN
AB		TEVA PHARMS	EQ 500MG BASE	N65282 001	Jan 20, 2006	Jan	NEWA
AB		WESTWARD	EQ 500MG BASE	N65311 001	Feb 07, 2006	Jan	NEWA

DURICEF

@ WARNER CHILCOTT

EQ 500MG BASE

N50512 001 Jan DISC

FOR SUSPENSION; ORAL

CEFADROXIL

AB		ORCHID HLTHCARE	EQ 250MG BASE/5ML	N65307 002	Oct 16, 2006	Oct	NEWA
AB			EQ 500MG BASE/5ML	N65307 003	Oct 16, 2006	Oct	NEWA
	+	RANBAXY	EQ 125MG BASE/5ML	N65115 001	Mar 26, 2003	Jul	CRLD
			EQ 125MG BASE/5ML	N65115 001	Mar 26, 2003	Feb	CTEC
AB		TEVA PHARMS	EQ 250MG BASE/5ML	N65278 001	Jan 20, 2006	Jan	NEWA
AB			EQ 500MG BASE/5ML	N65278 002	Jan 20, 2006	Jan	NEWA

DURICEF

@ WARNER CHILCOTT

EQ 125MG BASE/5ML

N50527 002 Feb DISC

TABLET; ORAL

CEFADROXIL

AB		HIKMA	EQ 1GM BASE	N65260 001	Mar 30, 2006	Mar	NEWA
AB	+	IVAX PHARMS	EQ 1GM BASE	N62774 001	Apr 08, 1987	Feb	CRLD
AB		ORCHID HLTHCARE	EQ 1GM BASE	N65301 001	Sep 18, 2006	Sep	NEWA

DURICEF

@ WARNER CHILCOTT

EQ 1GM BASE

N50528 001 Jan DISC

CEFAZOLIN SODIUMINJECTABLE; INJECTION
CEFAZOLIN

AP		ORCHID HLTHCARE	EQ 1GM BASE/VIAL	N65244 001	Aug 12, 2005	Nov	CPOT
AP			EQ 10GM BASE/VIAL	N65247 001	Aug 12, 2005	Nov	CPOT
		CEFAZOLIN AND DEXTROSE					
		@ B BRAUN	EQ 500MG BASE/VIAL	N50779 001	Jul 27, 2000	May	DISC

INJECTABLE; INJECTION

CEFAZOLIN SODIUM

+	SAMSON MEDCL	EQ 100GM BASE/VIAL	N65141 001	Nov 29, 2006	Nov	NEWA
+		EQ 300GM BASE/VIAL	N65141 002	Nov 29, 2006	Nov	NEWA

CEFDINIR

CAPSULE; ORAL

CEFDINIR

AB	LUPIN	300MG	N65264 001	May 19, 2006	May	NEWA
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OMNICEF

AB	+ ABBOTT	300MG	N50739 001	Dec 04, 1997	May	CFTG
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FOR SUSPENSION; ORAL

CEFDINIR

AB	LUPIN	125MG/5ML	N65259 001	May 31, 2006	Jul	CAHN
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AB	LUPIN (USA)	125MG/5ML	N65259 001	May 31, 2006	May	NEWA
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OMNICEF

AB	ABBOTT	125MG/5ML	N50749 001	Dec 04, 1997	May	CFTG
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CEFDITOREN PIVOXIL

TABLET; ORAL

SPECTRACEF

>A>	+	CORNERSTONE	200MG	N21222 001	Aug 29, 2001	Dec	CAHN
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>D>	+	PURDUE PHARMA LP	200MG	N21222 001	Aug 29, 2001	Dec	CAHN
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CEFOTAXIME SODIUM

INJECTABLE; INJECTION

CEFOTAXIME

AP	WOCKHARDT	EQ 1GM BASE/VIAL	N65197 001	Aug 29, 2006	Aug	NEWA
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CEFOTAXIME AND DEXTROSE 2.4% IN PLASTIC CONTAINER

	@ B BRAUN	EQ 2GM BASE	N50792 001	Jul 29, 2004	May	DISC
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CEFOTAXIME AND DEXTROSE 3.9% IN PLASTIC CONTAINER

	@ B BRAUN	EQ 1GM BASE	N50792 002	Jul 29, 2004	May	DISC
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CEFOTAXIME SODIUM

AP	ORCHID HLTHCARE	EQ 500MG BASE/VIAL	N65290 001	Aug 11, 2006	Aug	NEWA
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AP		EQ 1GM BASE/VIAL	N65290 002	Aug 11, 2006	Aug	NEWA
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AP		EQ 1GM BASE/VIAL	N65293 001	Aug 10, 2006	Aug	NEWA
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AP		EQ 2GM BASE/VIAL	N65290 003	Aug 11, 2006	Aug	NEWA
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AP		EQ 2GM BASE/VIAL	N65293 002	Aug 10, 2006	Aug	NEWA
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AP		EQ 10GM BASE/VIAL	N65292 001	Aug 10, 2006	Aug	NEWA
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CEFOTETAN DISODIUM

INJECTABLE; INJECTION

CEFOTAN

	@ ASTRAZENECA	EQ 1GM BASE/VIAL	N50588 001	Dec 27, 1985	Oct	DISC
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	@	EQ 1GM BASE/VIAL	N63293 001	Apr 29, 1993	Jun	DISC
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	@	EQ 2GM BASE/VIAL	N50588 002	Dec 27, 1985	Oct	DISC
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	@	EQ 2GM BASE/VIAL	N63293 002	Apr 29, 1993	Jun	DISC
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	@	EQ 10GM BASE/VIAL	N50588 003	Apr 25, 1988	Oct	DISC
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CEFOTAN IN PLASTIC CONTAINER

	@ ASTRAZENECA	EQ 20MG BASE/ML	N50694 002	Jul 30, 1993	Oct	DISC
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	@	EQ 40MG BASE/ML	N50694 001	Jul 30, 1993	Oct	DISC
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CEFOXITIN SODIUMINJECTABLE; INJECTION
CEFOXITIN

AP	ORCHID HLTHCARE	EQ 1GM BASE/VIAL	N65313 001	Jan 23, 2006	Jan	NEWA
AP		EQ 2GM BASE/VIAL	N65313 002	Jan 23, 2006	Jan	NEWA
AP		EQ 10GM BASE/VIAL	N65312 001	Feb 13, 2006	Jan	NEWA
CEFOXITIN AND DEXTROSE IN DUPLEX CONTAINER						
AP	B BRAUN	EQ 1GM BASE/VIAL	N65214 001	Mar 10, 2006	Feb	NEWA
AP		EQ 2GM BASE/VIAL	N65214 002	Mar 10, 2006	Feb	NEWA

CEFPROZILFOR SUSPENSION; ORAL
CEFPROZIL

AB	RANBAXY	125MG/5ML	N65202 001	Jun 30, 2006	Jun	NEWA
AB		250MG/5ML	N65202 002	Jun 30, 2006	Jun	NEWA

TABLET; ORAL
CEFPROZIL

AB	RANBAXY	250MG	N65198 001	Dec 13, 2006	Nov	NEWA
AB		500MG	N65198 002	Dec 13, 2006	Nov	NEWA

CEFTAZIDIME (ARGININE FORMULATION)INJECTABLE; INJECTION
CEPTAZ

@	GLAXOSMITHKLINE	1GM/VIAL	N50646 002	Sep 27, 1990	May	DISC
@		2GM/VIAL	N50646 003	Sep 27, 1990	May	DISC
@		10GM/VIAL	N50646 004	Sep 27, 1990	May	DISC

CEFTAZIDIME SODIUMINJECTABLE; INJECTION
CEFTAZIDIME SODIUM IN PLASTIC CONTAINER

@	BAXTER HLTHCARE	EQ 10MG BASE/ML	N63221 001	Apr 29, 1993	Aug	DISC
@		EQ 20MG BASE/ML	N63221 002	Apr 29, 1993	Aug	DISC
@		EQ 40MG BASE/ML	N63221 003	Apr 29, 1993	Aug	DISC

FORTAZ IN PLASTIC CONTAINER

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

CEFTRIAZONE SODIUMINJECTABLE; IM-IV
CEFTRIAZONE

AP	AM PHARM PARTNERS	EQ 250MG BASE/VIAL	N65245 001	Feb 15, 2006	Jan	NEWA
AP		EQ 500MG BASE/VIAL	N65245 002	Feb 15, 2006	Jan	NEWA
AP		EQ 1GM BASE/VIAL	N65245 003	Feb 15, 2006	Jan	NEWA
AP		EQ 2GM BASE/VIAL	N65245 004	Feb 15, 2006	Jan	NEWA

CEFTRIAZONE SODIUM

AP	TEVA	EQ 1GM BASE/VIAL	N65262 001	Jun 29, 2006	Jun	NEWA
AP		EQ 2GM BASE/VIAL	N65262 002	Jun 29, 2006	Jun	NEWA

INJECTABLE; INJECTION
CEFTRIAZONE

AP	AM PHARM	EQ 10GM BASE/VIAL	N65252 001	Feb 15, 2006	Jan	NEWA
AP	LUPIN	EQ 10GM BASE/VIAL	N65263 001	Sep 12, 2006	Aug	NEWA
AP	WOCKHARDT	EQ 1GM BASE/VIAL	N65180 001	May 12, 2006	May	NEWA

CEFTRIAZONE SODIUM

AP	TEVA	EQ 10GM BASE/VIAL	N65274 001	May 01, 2006	Apr	NEWA
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CEFUROXIME AXETIL

TABLET; ORAL

CEFUROXIME AXETIL

AB	AUROBINDO PHARMA LTD	EQ 125MG BASE	N65308 001	Mar 29, 2006	Mar	NEWA
AB		EQ 250MG BASE	N65308 002	Mar 29, 2006	Mar	NEWA
AB		EQ 500MG BASE	N65308 003	Mar 29, 2006	Mar	NEWA

CEPHALEXIN

CAPSULE; ORAL

CEPHALEXIN

AB	HIKMA	EQ 250MG BASE	N65215 001	Jan 24, 2006	Jan	NEWA
AB		EQ 500MG BASE	N65215 002	Jan 24, 2006	Jan	NEWA

KEFLEX

	ADVANCIS PHARM	EQ 333MG BASE	N50405 004	May 12, 2006	May	NEWA
AB		EQ 500MG BASE	N50405 003		May	CRLD
+		EQ 750MG BASE	N50405 005	May 12, 2006	May	NEWA

FOR SUSPENSION; ORAL

CEPHALEXIN

AB	ORCHID HLTHCARE	EQ 125MG BASE/5ML	N65326 001	Jul 10, 2006	Jun	NEWA
AB		EQ 250MG BASE/5ML	N65326 002	Jul 10, 2006	Jun	NEWA

CEPHRADINE

CAPSULE; ORAL

ANSPOR

@ GLAXOSMITHKLINE

250MG

N61859 001

Mar DISC

@

500MG

N61859 002

Mar DISC

CEPHRADINE

>D>	AB	TEVA	250MG	N62683 001	Jan 09, 1987	Dec	CTEC
>A>			250MG	N62683 001	Jan 09, 1987	Dec	CTEC
>D>	AB		500MG	N62683 002	Jan 09, 1987	Dec	CRLD
>A>	+		500MG	N62683 002	Jan 09, 1987	Dec	CRLD
>D>		VELOSEF					
>D>	AB	+	APOTHECON	500MG	N61764 002	Dec	DISC
>A>		@	500MG	N61764 002		Dec	DISC

CETYL ALCOHOL; COLFOSKERIL PALMITATE; TYLOXAPOL

FOR SUSPENSION; INTRATRACHEAL

EXOSURF NEONATAL

@ GLAXOSMITHKLINE

12MG/VIAL;108MG/VIAL;8MG/VIAL

N20044 001 Aug 02, 1990 May DISC

CHLORHEXIDINE GLUCONATE

SOLUTION; DENTAL

CHLORHEXIDINE GLUCONATE

AT	ACTAVIS MID ATLANTIC	0.12%	N74291 001	Dec 28, 1995	Jul	CAHN
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CHLOROPROCAINE HYDROCHLORIDE

INJECTABLE; INJECTION

NESACAINE

+	ABRAXIS BIOSCIENCE	1%	N09435 001		Jul	CAHN	
AP		2%	N09435 002		Jul	CAHN	
	NESACAINE-MPF						
AP	+	ABRAXIS BIOSCIENCE	2%	N09435 006	May 02, 1996	Jul	CAHN
	@	2%	N09435 003		Jul	CAHN	
	@	3%	N09435 004		Jul	CAHN	

INJECTABLE; INJECTION
NESACAINE-MPF

AP + ABRAXIS BIOSCIENCE 3% N09435 007 May 02, 1996 Jul CAHN

CHLOROTHIAZIDE; RESERPINE

TABLET; ORAL
DIUPRES-250

@ MERCK 250MG;0.125MG N11635 003 Aug 26, 1987 Jun DISC

DIUPRES-500

@ MERCK 500MG;0.125MG N11635 006 Aug 26, 1987 Jun DISC

CHLORPHENIRAMINE POLISTIREX; HYDROCODONE POLISTIREX

SUSPENSION, EXTENDED RELEASE; ORAL

TUSSIONEX PENNKINETIC

+ UCB INC EQ 8MG MALEATE/5ML;EQ 10MG
BITARTRATE/5ML N19111 001 Dec 31, 1987 Nov CTNA

CHLORPROMAZINE

SUPPOSITORY; RECTAL

THORAZINE

@ GLAXOSMITHKLINE 25MG N09149 024 May DISC

@ 100MG N09149 033 May DISC

CHLORPROMAZINE HYDROCHLORIDE

CAPSULE, EXTENDED RELEASE; ORAL

THORAZINE

@ GLAXOSMITHKLINE 200MG N11120 019 May DISC

@ 300MG N11120 020 May DISC

CONCENTRATE; ORAL

CHLORPROMAZINE HYDROCHLORIDE

AA ACTAVIS MID ATLANTIC 100MG/ML N86863 001 Jul CAHN

CICLESONIDE

SPRAY, METERED; NASAL

OMNARIS

+ ALTANA PHARMA 0.05MG/INH N22004 001 Oct 20, 2006 Oct NEWA

CICLOPIROX

CREAM; TOPICAL

CICLOPIROX

AB PERRIGO 0.77% N77364 001 Mar 03, 2006 Mar CAHN

AB PERRIGO NEW YORK 0.77% N77364 001 Mar 03, 2006 Feb NEWA

SUSPENSION; TOPICAL

CICLOPIROX

>A> AB PERRIGO NEW YORK 0.77% N77676 001 Dec 15, 2006 Dec NEWA

CILASTATIN SODIUM; IMIPENEM

INJECTABLE; INJECTION

PRIMAXIN

@ MERCK EQ 750MG BASE/VIAL;750MG/VIAL N50630 002 Dec 14, 1990 Jun DISC

CILOSTAZOL

TABLET; ORAL

CILOSTAZOL

AB MUTUAL PHARM 50MG N77208 002 Mar 29, 2006 Mar NEWA

AB 100MG N77208 001 Mar 29, 2006 Mar NEWA

TABLET; ORAL

CLOSTAZOL

AB	MYLAN	50MG	N77323 002	Apr 20, 2006	Apr	NEWA
AB		100MG	N77323 001	Apr 20, 2006	Apr	NEWA

CIMETIDINE

TABLET; ORAL

CIMETIDINE

@ ENDO PHARMS

200MG

N74281 001 May 17, 1994 Feb DISC

@

300MG

N74281 002 May 17, 1994 Feb DISC

@

400MG

N74281 003 May 17, 1994 Feb DISC

@

800MG

N74329 001 May 17, 1994 Feb DISC

CIMETIDINE HYDROCHLORIDE

INJECTABLE; INJECTION

CIMETIDINE HYDROCHLORIDE

@ ENDO PHARMS

EQ 300MG BASE/2ML

N74005 001 Aug 31, 1994 Feb DISC

SOLUTION; ORAL

CIMETIDINE HYDROCHLORIDE

AA + ACTAVIS MID ATLANTIC EQ 300MG BASE/5ML

N74176 001 Jun 01, 1994 Jul CAHN

@ ENDO PHARMS

EQ 300MG BASE/5ML

N74251 001 Dec 22, 1994 Feb DISC

CIPROFLOXACIN

INJECTABLE; INJECTION

CIPRO

AP	+	BAYER PHARMS	400MG/40ML (10MG/ML)	N19847 001	Dec 26, 1990	Aug	CFTG
		@	1200MG/120ML (10MG/ML)	N19847 003	Dec 26, 1990	Aug	DISC
AP	+		200MG/20ML (10MG/ML)	N19847 002	Dec 26, 1990	Aug	CFTG

CIPROFLOXACIN

AP		ABRAXIS PHARM	400MG/40ML (10MG/ML)	N76484 002	Aug 28, 2006	Aug	NEWA
AP			200MG/20ML (10MG/ML)	N76484 001	Aug 28, 2006	Aug	NEWA
AP		BEDFORD LABS	200MG/20ML (10MG/ML)	N76992 001	Aug 28, 2006	Aug	NEWA
AP			400MG/40ML (10MG/ML)	N76992 002	Aug 28, 2006	Aug	NEWA
			1200MG/120ML (10MG/ML)	N76993 001	Aug 28, 2006	Aug	NEWA
AP		HOSPIRA	200MG/20ML (10MG/ML)	N77245 001	Aug 28, 2006	Aug	NEWA
AP			400MG/40ML (10MG/ML)	N77245 002	Aug 28, 2006	Aug	NEWA
AP		SICOR PHARMS	400MG/40ML (10MG/ML)	N77782 002	Aug 28, 2006	Aug	NEWA
AP			200MG/20ML (10MG/ML)	N77782 001	Aug 28, 2006	Aug	NEWA

SOLUTION/DROPS; OPHTHALMIC

CIPROFLOXACIN

AT		NEXUS PHARMS	EQ 0.3% BASE	N77689 001	Dec 13, 2006	Nov	NEWA
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CITALOPRAM HYDROBROMIDE

SOLUTION; ORAL

CITALOPRAM HYDROBROMIDE

AA		AUROBINDO PHARMA LTD	EQ 10MG BASE/5ML	N77812 001	Aug 28, 2006	Aug	NEWA
AA		SILARX	EQ 10MG BASE/5ML	N77629 001	Jun 15, 2006	May	NEWA

TABLET; ORAL

CITALOPRAM HYDROBROMIDE

AB		ACTAVIS ELIZABETH	EQ 10MG BASE	N77033 001	Oct 28, 2004	Jun	CAHN
AB			EQ 20MG BASE	N77033 002	Oct 28, 2004	Jun	CAHN
AB			EQ 40MG BASE	N77033 003	Oct 28, 2004	Jun	CAHN
AB		INTERPHARM	EQ 10MG BASE	N77289 001	Nov 30, 2006	Nov	NEWA
AB			EQ 20MG BASE	N77289 002	Nov 30, 2006	Nov	NEWA
AB			EQ 40MG BASE	N77289 003	Nov 30, 2006	Nov	NEWA
AB		INVAGEN PHARMS	EQ 10MG BASE	N77534 001	Oct 03, 2006	Sep	NEWA

TABLET; ORAL

CITALOPRAM HYDROBROMIDE

AB	INVAGEN PHARMS	EQ 20MG BASE	N77534 002	Oct 03, 2006	Sep	NEWA
AB		EQ 40MG BASE	N77534 003	Oct 03, 2006	Sep	NEWA
AB	MUTUAL PHARM	EQ 10MG BASE	N77052 001	Jul 03, 2006	Jun	NEWA
AB		EQ 20MG BASE	N77052 002	Jul 03, 2006	Jun	NEWA
AB		EQ 40MG BASE	N77052 003	Jul 03, 2006	Jun	NEWA
AB	TARO	EQ 10MG BASE	N77278 001	Mar 22, 2006	Mar	NEWA
AB		EQ 20MG BASE	N77278 002	Mar 22, 2006	Mar	NEWA
AB		EQ 40MG BASE	N77278 003	Mar 22, 2006	Mar	NEWA
AB	TEVA PHARMS	EQ 10MG BASE	N77213 001	Mar 31, 2006	Mar	NEWA
AB		EQ 20MG BASE	N77213 002	Mar 31, 2006	Mar	NEWA
AB		EQ 40MG BASE	N77213 003	Mar 31, 2006	Mar	NEWA

TABLET, ORALLY DISINTEGRATING; ORAL

CITALOPRAM HYDROBROMIDE

@	BIOVAIL LABS INTL	EQ 10MG BASE	N21763 001	Dec 20, 2005	Oct	DISC
@		EQ 20MG BASE	N21763 002	Dec 20, 2005	Oct	DISC
@		EQ 40MG BASE	N21763 003	Dec 20, 2005	Oct	DISC

CITRIC ACID; MAGNESIUM OXIDE; SODIUM CARBONATE

SOLUTION; IRRIGATION

IRRIGATING SOLUTION G IN PLASTIC CONTAINER

@	BAXTER HLTHCARE	3.24GM/100ML;380MG/100ML;430MG/100ML	N18519 001	Jun 22, 1982	Jun	DISC
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UROLOGIC G IN PLASTIC CONTAINER

+	HOSPIRA	3.24GM/100ML;380MG/100ML;430MG/100ML	N18904 001	May 27, 1983	Jun	CTEC
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CITRIC ACID; UREA, C-13

FOR SOLUTION, TABLET, FOR SOLUTION; ORAL

IDKIT:HP

@	BREATHID 2006	N/A, 4GM; 75MG, N/A	N21314 001	Dec 17, 2002	Sep	CAHN
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CLARITHROMYCIN

TABLET; ORAL

CLARITHROMYCIN

AB	WOCKHARDT	250MG	N65266 001	May 31, 2006	May	NEWA
AB		500MG	N65266 002	May 31, 2006	May	NEWA

TABLET, EXTENDED RELEASE; ORAL

CLARITHROMYCIN

+	RANBAXY	1GM	N65210 001	Jan 26, 2005	Apr	CRLD
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CLEMASTINE FUMARATE

SYRUP; ORAL

CLEMASTINE FUMARATE

AA	ACTAVIS MID ATLANTIC	EQ 0.5MG BASE/5ML	N74075 001	Oct 31, 1993	Jul	CAHN
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CLINDAMYCIN PHOSPHATE

SOLUTION; TOPICAL

CLINDAMYCIN PHOSPHATE

AT	ACTAVIS MID ATLANTIC	EQ 1% BASE	N62811 001	Sep 01, 1988	Jul	CAHN
AT	ALTANA	EQ 1% BASE	N65254 001	Feb 14, 2006	Jan	NEWA

CLINDAMYCIN PHOSPHATE; TRETINOIN

GEL; TOPICAL

ZIANA

+	MEDICIS	1.2%;0.025%	N50802 001	Nov 07, 2006	Nov	NEWA
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CLOBETASOL PROPIONATE

CREAM; TOPICAL

CLOBETASOL PROPIONATE

AB1	ACTAVIS MID ATLANTIC	0.05%	N74139 001	Aug 03, 1994	Jun	CAHN
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OINTMENT; TOPICAL

CLOBETASOL PROPIONATE

AB	ACTAVIS MID ATLANTIC	0.05%	N74128 001	Aug 03, 1994	Jun	CAHN
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SOLUTION; TOPICAL

CLOBETASOL PROPIONATE

AT	ACTAVIS MID ATLANTIC	0.05%	N74331 001	Dec 15, 1995	Jul	CAHN
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@ ALTANA

0.05%

N75391 001	Feb 08, 1999	May	DISC
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SPRAY; TOPICAL

CLOBEX

+	GALDERMA LABS LP	0.05%	N21835 001	Oct 27, 2005	Feb	CAHN
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CLOFAZIMINE

CAPSULE; ORAL

LAMPRENE

+	NOVARTIS	50MG	N19500 002	Dec 15, 1986	Jun	CMFD
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CLONAZEPAM

TABLET; ORAL

CLONAZEPAM

AB	ACTAVIS ELIZABETH	0.5MG	N74869 001	Oct 31, 1996	Jun	CAHN
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AB		1MG	N74869 002	Oct 31, 1996	Jun	CAHN
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AB		2MG	N74869 003	Oct 31, 1996	Jun	CAHN
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AB	VINTAGE PHARMS	0.5MG	N77856 001	Jun 28, 2006	Jun	NEWA
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AB		1MG	N77856 002	Jun 28, 2006	Jun	NEWA
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AB		2MG	N77856 003	Jun 28, 2006	Jun	NEWA
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CLONIDINE HYDROCHLORIDE

TABLET; ORAL

CLONIDINE HYDROCHLORIDE

AB	ACTAVIS ELIZABETH	0.1MG	N70974 001	Dec 16, 1986	Jun	CAHN
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AB		0.2MG	N70975 001	Dec 16, 1986	Jun	CAHN
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AB		0.3MG	N70976 001	Dec 16, 1986	Jun	CAHN
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CLOPIDOGREL BISULFATE

TABLET; ORAL

CLOPIDOGREL BISULFATE

AB	APOTEX	EQ 75MG BASE	N76274 001	Jan 20, 2006	Jan	NEWA
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PLAVIX

AB	+	SANOFI SYNTHELABO	EQ 75MG BASE	N20839 001	Nov 17, 1997	Jan	CFTG
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CLOZAPINE

TABLET, ORALLY DISINTEGRATING; ORAL

FAZACLO ODT

AVANIR PHARMS

25MG

N21590 001	Feb 10, 2004	Jul	CAHN
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50MG

N21590 003	Jun 03, 2005	Jul	CAHN
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TABLET, ORALLY DISINTEGRATING; ORAL

FAZACLO ODT

+	AVANIR PHARMS	100MG	N21590 002	Feb 10, 2004	Jul	CAHN
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COBALT CHLORIDE, CO-57; CYANOCOBALAMIN; CYANOCOBALAMIN, CO-57; INTRINSIC FACTOR

N/A; N/A

RUBRATOPE-57 KIT

@ BRACCO	N/A;N/A;N/A;N/A	N16089 001		Jun	DISC
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CODEINE PHOSPHATE; PHENYLEPHRINE HYDROCHLORIDE; PROMETHAZINE HYDROCHLORIDE

SYRUP; ORAL

PROMETH HYDROCHLORIDE, PHENYLEPHRINE HYDROCHLORIDE W/CODEINE PHOSPHATE

AA	VINTAGE	10MG/5ML;5MG/5ML;6.25MG/5ML	N40660 001	Dec 07, 2006	Nov	NEWA
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PROMETH VC W/ CODEINE

AA	+	ACTAVIS MID ATLANTIC	10MG/5ML;5MG/5ML;6.25MG/5ML	N88764 001	Oct 31, 1984	Nov	CFTG
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+			10MG/5ML;5MG/5ML;6.25MG/5ML	N88764 001	Oct 31, 1984	Jul	CAHN
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+	ALPHARMA US PHARMS		10MG/5ML;5MG/5ML;6.25MG/5ML	N88764 001	Oct 31, 1984	Jan	CTEC
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PROMETHAZINE VC W/ CODEINE

@ MORTON GROVE	10MG/5ML;5MG/5ML;6.25MG/5ML	N88896 001	Jan 04, 1985	Jan	DISC
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CODEINE PHOSPHATE; PROMETHAZINE HYDROCHLORIDE

SYRUP; ORAL

PROMETH W/ CODEINE

AA	+	ACTAVIS MID ATLANTIC	10MG/5ML;6.25MG/5ML	N88763 001	Oct 31, 1984	Jul	CAHN
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PROMETHAZINE WITH CODEINE SYRUP

AA	VINTAGE	10MG/5ML;6.25MG/5ML	N40650 001	Jan 31, 2006	Jan	NEWA
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CODEINE PHOSPHATE; PSEUDOEPHEDRINE HYDROCHLORIDE; TRIPROLIDINE HYDROCHLORIDE

SYRUP; ORAL

TRIACIN-C

AA	ACTAVIS MID ATLANTIC	10MG/5ML;30MG/5ML;1.25MG/5ML	N88704 001	Mar 22, 1985	Jul	CAHN
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COLESTIPOL HYDROCHLORIDE

GRANULE; ORAL

COLESTID

AB	+	PHARMACIA AND UPJOHN	5GM/PACKET	N17563 004	Sep 22, 1995	Apr	CFTG
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AB			5GM/SCOOPFUL	N17563 003	Sep 22, 1995	Apr	CFTG
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COLESTIPOL HYDROCHLORIDE

AB	IMPAX LABS		5GM/SCOOPFUL	N77277 001	May 02, 2006	Apr	NEWA
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AB			5GM/PACKET	N77277 002	May 02, 2006	Apr	NEWA
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FLAVORED COLESTID

PHARMACIA AND UPJOHN	5GM/PACKET	N17563 001		Apr	CFTG
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TABLET; ORAL

COLESTID

AB	+	PHARMACIA AND UPJOHN	1GM	N20222 001	Jul 19, 1994	Oct	CFTG
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COLESTIPOL HYDROCHLORIDE

AB	IMPAX LABS		1GM	N77510 001	Oct 24, 2006	Oct	NEWA
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COPPER

INTRAUTERINE DEVICE; INTRAUTERINE

PARAGARD T 380A

+	DURAMED PHARMS BARR	309MG/COPPER	N18680 001	Nov 15, 1984	Oct	CTNA
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CROMOLYN SODIUM

CONCENTRATE; ORAL					
GASTROCROM					
	+ AZUR PHARMA	100MG/5ML	N20479 001	Feb 29, 1996	May CDFR
SOLUTION; INHALATION					
CROMOLYN SODIUM					
AN	ACTAVIS MID ATLANTIC	10MG/ML	N75067 001	Jul 19, 1999	Jul CAHN
SOLUTION, CONCENTRATE; ORAL					
GASTROCROM					
	+ AZUR PHARMA	100MG/5ML	N20479 001	Feb 29, 1996	Feb CAHN

CYANOCOBALAMIN

GEL, METERED; NASAL					
NASCOBAL					
	@ QOL MEDCL	0.5MG/INH	N19722 001	Nov 05, 1996	Mar DISC
INJECTABLE; INJECTION					
CYANOCOBALAMIN					
	@ ABRAXIS PHARM	0.1MG/ML	N80557 002		Nov DISC
AP	+ BIONICHE PHARMA	1MG/ML	N40451 001	Sep 23, 2003	Nov CRLD
AP	+ LUITPOLD	1MG/ML	N80737 001		Nov CRLD
	@ WATSON LABS	0.1MG/ML	N80573 002		Nov DISC
	@	1MG/ML	N80573 001		Nov DISC
RUBRAMIN PC					
	@ BRISTOL MYERS SQUIBB	1MG/ML	N06799 010	Apr 28, 1988	Nov DISC
VIBISONE					
AP	+ ABRAXIS PHARM	1MG/ML	N80557 003		Nov CRLD

CYCLOBENZAPRINE HYDROCHLORIDE

TABLET; ORAL					
CYCLOBENZAPRINE HYDROCHLORIDE					
AB	AMIDE PHARM	5MG	N77291 001	Feb 03, 2006	Jan NEWA
AB	JUBILANT PHARMS	5MG	N77563 001	Apr 19, 2006	Apr NEWA
AB		10MG	N77563 002	Apr 19, 2006	Apr NEWA
AB	MUTUAL PHARM	5MG	N73541 002	Apr 06, 2006	Mar NEWA
AB	MYLAN	5MG	N73144 002	Feb 03, 2006	Jan NEWA
AB	SANDOZ	5MG	N72854 002	Feb 03, 2006	Jan NEWA
AB	WATSON LABS	5MG	N71611 002	Feb 03, 2006	Jan NEWA
		7.5MG	N71611 003	Feb 03, 2006	Jan NEWA
FLEXERIL					
AB	MCNEIL CONS SPECLT	5MG	N17821 001		Jan CFTG

CYPROHEPTADINE HYDROCHLORIDE

SYRUP; ORAL					
CYPROHEPTADINE HYDROCHLORIDE					
AA	+ ACTAVIS MID ATLANTIC	2MG/5ML	N86833 001		Jul CAHN
AA	+ ALPHARMA US PHARMS	2MG/5ML	N86833 001		Jun CTEC
AA	LYNE	2MG/5ML	N40668 001	Jun 28, 2006	Jun NEWA
TABLET; ORAL					
CYPROHEPTADINE HYDROCHLORIDE					
	@ TG UNITED LABS	4MG	N88212 001	May 26, 1983	May CAHN
CYPROHEPTADINE HYDROCHLORIDE					
AA	STASON PHARMS	4MG	N40644 001	May 30, 2006	May NEWA

DAPTOMYCIN

INJECTABLE; IV (INFUSION)

CUBICIN

@ CUBIST

250MG/VIAL

N21572 001 Sep 12, 2003 Jun DISC

DARUNAVIR ETHANOLATE

TABLET; ORAL

PREZISTA

+ TIBOTEC

EQ 300MG BASE

N21976 001 Jun 23, 2006 Jun NEWA

DASATINIB

TABLET; ORAL

SPRYCEL

BRISTOL MYERS SQUIBB

20MG

N21986 001 Jun 28, 2006 Jun NEWA

50MG

N21986 002 Jun 28, 2006 Jun NEWA

+

70MG

N21986 003 Jun 28, 2006 Jun NEWA

DECITABINE

INJECTABLE; INTRAVENOUS

DACOGEN

+ MGI PHARMA INC

50MG/VIAL

N21790 001 May 02, 2006 May NEWA

DEFEROXAMINE MESYLATE

INJECTABLE; INJECTION

DEFEROXAMINE MESYLATE

AP SICOR PHARMS

500MG/VIAL

N76806 001 Mar 31, 2006 Mar NEWA

AP

2GM/VIAL

N76806 002 Mar 31, 2006 Mar NEWA

DEMECLOCYCLINE HYDROCHLORIDE

TABLET; ORAL

DECLOMYCIN

@ GLADES PHARMS LLC

75MG

N50261 001

Mar CAHN

AB

150MG

N50261 002

Mar CAHN

AB

+

300MG

N50261 003

Mar CAHN

@ PROTEIN DESIGN LABS

75MG

N50261 001

Feb CAHN

AB

150MG

N50261 002

Feb CAHN

AB

+

300MG

N50261 003

Feb CAHN

DESLORATADINE; PSEUDOEPHEDRINE SULFATE

TABLET, EXTENDED RELEASE; ORAL

CLARINEX-D 12 HOUR

+ SCHERING

2.5MG;120MG

N21313 001 Feb 01, 2006 Feb NEWA

DESMOPRESSIN ACETATE

INJECTABLE; INJECTION

DESMOPRESSIN ACETATE

@ BEDFORD

0.004MG/ML

N74575 001 Feb 18, 2000 Jan DISC

DESMOPRESSIN ACETATE PRESERVATIVE FREE

@ BEDFORD

0.004MG/ML

N74574 001 Feb 18, 2000 Jan DISC

TABLET; ORAL

DESMOPRESSIN ACETATE

AB APOTEX

0.1MG

N77414 001 Mar 07, 2006 Feb NEWA

AB

0.2MG

N77414 002 Mar 07, 2006 Feb NEWA

AB

TEVA PHARMS

0.1MG

N77122 001 Jan 25, 2006 Jan NEWA

TABLET; ORAL

DESMOPRESSIN ACETATE

AB		TEVA PHARMS	0.2MG	N77122 002	Jan 25, 2006	Jan	NEWA
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DESOGESTREL; ETHINYL ESTRADIOL

TABLET; ORAL-28

MIRCETTE

AB	+	DURAMED	0.15MG, N/A; 0.02MG, 0.01MG	N20713 001	Apr 22, 1998	Feb	CAHN
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AB	+		0.15MG, N/A; 0.02MG, 0.01MG	N20713 001	Apr 22, 1998	Feb	CAHN
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DESONIDE

AEROSOL, FOAM; TOPICAL

VERDESO

	+	CONNETICS	0.05%	N21978 001	Sep 19, 2006	Sep	NEWA
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GEL; TOPICAL

DESONATE

	+	SKINMEDICA	0.05%	N21844 001	Oct 20, 2006	Oct	NEWA
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DEXAMETHASONE

ELIXIR; ORAL

DEXAMETHASONE

AA	+	ACTAVIS MID ATLANTIC	0.5MG/5ML	N84754 001		Jul	CAHN
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AA	+	ALPHARMA US PHARMS	0.5MG/5ML	N84754 001		Jun	CRLD
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HEXADROL

@ ORGANON USA INC

			0.5MG/5ML	N12674 001		Jun	DISC
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MYMETHASONE

AA	+	MORTON GROVE	0.5MG/5ML	N88254 001	Jul 27, 1983	Jun	CRLD
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TABLET; ORAL

DECADRON

@ MERCK

			0.5MG	N11664 001		Aug	DISC
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			0.75MG	N11664 002		Aug	DISC
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DEXAMETHASONE

BP		ROXANE	0.5MG	N84611 001		Aug	CTEC
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BP			0.75MG	N84613 001		Aug	CTEC
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HEXADROL

@ ORGANON USA INC

			4MG	N12675 010		Jun	DISC
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DEXAMETHASONE SODIUM PHOSPHATE

INJECTABLE; INJECTION

DEXAMETHASONE

AP	+	BAXTER HLTHCARE	EQ 10MG PHOSPHATE/ML	N87702 001	Sep 07, 1982	Oct	CRLD
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AP			EQ 10MG PHOSPHATE/ML	N87702 001	Sep 07, 1982	Mar	CAHN
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DEXMETHYLPHENIDATE HYDROCHLORIDE

CAPSULE, EXTENDED RELEASE; ORAL

FOCALIN XR

		NOVARTIS	15MG	N21802 004	Aug 01, 2006	Aug	NEWA
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DEXTROAMPHETAMINE SULFATE

TABLET; ORAL

DEXTROAMPHETAMINE SULFATE

		@ ENDO PHARMS	5MG	N40299 001	May 13, 1999	Feb	DISC
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DEXTROMETHORPHAN HYDROBROMIDE; PROMETHAZINE HYDROCHLORIDE

SYRUP; ORAL

PROMETH W/ DEXTROMETHORPHAN

AA	+	ACTAVIS MID ATLANTIC	15MG/5ML;6.25MG/5ML	N88762	001	Oct 31, 1984	Jul	CAHN
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PROMETHAZINE DM

AA		VINTAGE	15MG/5ML;6.25MG/5ML	N40649	001	Feb 14, 2006	Jan	NEWA
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DIAZEPAM

TABLET; ORAL

DIAZEPAM

AB		ACTAVIS ELIZABETH	2MG	N70781	001	Mar 19, 1986	Jun	CAHN
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AB			5MG	N70706	001	Mar 19, 1986	Jun	CAHN
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AB			10MG	N70707	001	Mar 19, 1986	Jun	CAHN
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AB		VINTAGE PHARMS	2MG	N77749	001	Mar 31, 2006	Mar	NEWA
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AB			5MG	N77749	002	Mar 31, 2006	Mar	NEWA
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AB			10MG	N77749	003	Mar 31, 2006	Mar	NEWA
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DIAZOXIDE

INJECTABLE; INJECTION

HYPERSTAT

@ SCHERING

15MG/ML

N16996	001		Oct	DISC
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DICLOFENAC SODIUM

TABLET, DELAYED RELEASE; ORAL

DICLOFENAC SODIUM

AB		ACTAVIS ELIZABETH	50MG	N74514	001	Mar 26, 1996	Jun	CAHN
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AB			75MG	N74514	002	Mar 26, 1996	Jun	CAHN
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AB		SANDOZ	75MG	N74394	001	Nov 30, 1995	May	CRLD
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VOLTAREN

AB	+	NOVARTIS	75MG	N19201	003	Jul 28, 1988	May	CMFD
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TABLET, EXTENDED RELEASE; ORAL

DICLOFENAC SODIUM

AB		ACTAVIS ELIZABETH	100MG	N75910	001	Jan 07, 2002	Jun	CAHN
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DIETHYLPROPION HYDROCHLORIDE

TABLET; ORAL

DIETHYLPROPION HYDROCHLORIDE

@ TG UNITED LABS

25MG

N88267	001	Aug 25, 1983	May	CAHN
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@

25MG

N88268	001	Aug 25, 1983	May	CAHN
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TENUATE

>D>	+	SANOFI AVENTIS US	25MG	N11722	002		Dec	CAHN
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>A>	+	WATSON PHARMS	25MG	N11722	002		Dec	CAHN
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TABLET, EXTENDED RELEASE; ORAL

TENUATE DOSPAN

>D>	+	SANOFI AVENTIS US	75MG	N12546	001		Dec	CAHN
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>A>	+	WATSON PHARMS	75MG	N12546	001		Dec	CAHN
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DIFLORASONE DIACETATE

CREAM; TOPICAL

DIFLORASONE DIACETATE

BX	+	ALTANA	0.05%	N76263	001	Dec 20, 2002	Jan	CRLD
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FLORONE

@ PHARMACIA AND UPJOHN 0.05%

N17741	001		Jan	DISC
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CREAM; TOPICAL

FLORONE E

@ PHARMACIA AND UPJOHN 0.05%

N19259 001 Aug 28, 1985 Jan DISC

OINTMENT; TOPICAL

DIFLORASONE DIACETATE

AB + TARO 0.05%

N75331 001 May 14, 1999 Jan CRLD

FLORONE

@ PHARMACIA AND UPJOHN 0.05%

N17994 001 Jan DISC

PSORCON

@ PHARMACIA AND UPJOHN 0.05%

N19260 001 Aug 28, 1985 Jan DISC

DIFLUNISAL

TABLET; ORAL

DIFLUNISAL

AB + TEVA 500MG

N73673 001 Jul 31, 1992 Jun CRLD

DOLOBID

@ MERCK 250MG

N18445 001 Apr 19, 1982 Jun DISC

@ 500MG

N18445 002 Apr 19, 1982 Jun DISC

DIGOXIN

INJECTABLE; INJECTION

DIGOXIN

AP SANDOZ 0.25MG/ML

N40481 001 Aug 21, 2003 Jan CAHN

DILTIAZEM HYDROCHLORIDE

CAPSULE, EXTENDED RELEASE; ORAL

DILTIAZEM HYDROCHLORIDE

AB3 ACTAVIS ELIZABETH 120MG

N74984 001 Dec 20, 1999 Jun CAHN

AB3 180MG

N74984 002 Dec 20, 1999 Jun CAHN

AB3 240MG

N74984 003 Dec 20, 1999 Jun CAHN

AB3 300MG

N74984 004 Dec 20, 1999 Jun CAHN

AB4 KV PHARM 120MG

N76563 002 Sep 12, 2006 Aug NEWA

AB4 180MG

N76563 003 Sep 12, 2006 Aug NEWA

AB4 240MG

N76563 004 Sep 12, 2006 Aug NEWA

AB4 300MG

N76563 005 Sep 12, 2006 Aug NEWA

AB4 360MG

N76563 006 Sep 12, 2006 Aug NEWA

AB4 420MG

N76563 001 Sep 12, 2006 Aug NEWA

DILTZAC

AB4 APOTEX INC 120MG

N76395 001 Feb 01, 2006 Jan NEWA

AB4 180MG

N76395 002 Feb 01, 2006 Jan NEWA

AB4 240MG

N76395 003 Feb 01, 2006 Jan NEWA

AB4 300MG

N76395 004 Feb 01, 2006 Jan NEWA

AB4 360MG

N76395 005 Feb 01, 2006 Jan NEWA

TIAZAC

AB4 + BIOVAIL 420MG

N20401 006 Oct 16, 1998 Aug CFTG

DILTIAZEM MALATE; ENALAPRIL MALEATE

TABLET, EXTENDED RELEASE; ORAL

TECZEM

@ BIOVAIL EQ 180MG HCL;5MG

N20507 001 Oct 04, 1996 Oct DISC

DIMETHYL SULFOXIDE

SOLUTION; INTRAVESICAL

RIMSO-50

AT + BIONICHE PHARMA 50%

N17788 001 Jul CAHN

DIPHENHYDRAMINE HYDROCHLORIDE

CAPSULE; ORAL

BENADRYL

>A>	@ MCNEIL CONS	25MG	N05845 007	Dec	CAHN
>A>	@	50MG	N05845 001	Dec	CAHN
>D>	@ PARKE DAVIS	25MG	N05845 007	Dec	CAHN
>D>	@	50MG	N05845 001	Dec	CAHN

ELIXIR; ORAL

BENADRYL

>A>	@ MCNEIL CONS	12.5MG/5ML	N05845 004	Dec	CAHN
>D>	@ PARKE DAVIS	12.5MG/5ML	N05845 004	Dec	CAHN

DIPYRIDAMOLE

TABLET; ORAL

DIPYRIDAMOLE

AB	AMIDE PHARM	25MG	N40542 001	Apr 21, 2006	Apr	NEWA
AB		50MG	N40542 002	Apr 21, 2006	Apr	NEWA
AB		75MG	N40542 003	Apr 21, 2006	Apr	NEWA
AB	GLENMARK PHARMA	25MG	N88999 001	Feb 05, 1991	Oct	CMFD
	@	25MG	N88999 001	Feb 05, 1991	Jul	CAHN
AB		50MG	N89000 001	Feb 05, 1991	Jul	CAHN
AB		75MG	N89001 001	Feb 05, 1991	Oct	CMFD
	@	75MG	N89001 001	Feb 05, 1991	Jul	CAHN

DISOPYRAMIDE PHOSPHATE

CAPSULE; ORAL

DISOPYRAMIDE PHOSPHATE

@	IVAX PHARMS	EQ 100MG BASE	N70186 001	Nov 18, 1985	Jan	DISC
@		EQ 150MG BASE	N70187 001	Nov 18, 1985	Jan	DISC
@	SANDOZ	EQ 100MG BASE	N70470 001	Dec 10, 1985	Jan	DISC
@		EQ 150MG BASE	N70471 001	Dec 10, 1985	Jan	DISC

DISULFIRAM

TABLET; ORAL

ANTABUSE

ODYSSEY PHARMS	250MG	N88482 001	Dec 08, 1983	Oct	CRLD
+	500MG	N88483 001	Dec 08, 1983	Oct	CMFD

DOLASETRON MESYLATE

INJECTABLE; INJECTION

ANZEMET

+	SANOFI AVENTIS US	EQ 12.5MG BASE/0.625ML (EQ 20MG BASE/ML)	N20624 002	Sep 11, 1997	Mar	CAIN
+		EQ 100MG BASE/5ML (EQ 20MG BASE/ML)	N20624 001	Sep 11, 1997	Mar	CAIN
+		EQ 500MG BASE/25ML (EQ 20MG BASE/ML)	N20624 003	Dec 11, 2001	Mar	CAIN

TABLET; ORAL

ANZEMET

SANOFI AVENTIS US	EQ 50MG BASE	N20623 001	Sep 11, 1997	Mar	CAIN
+	EQ 100MG BASE	N20623 002	Sep 11, 1997	Mar	CAIN

DOPAMINE HYDROCHLORIDE

INJECTABLE; INJECTION

DOPAMINE HYDROCHLORIDE

@	HOSPIRA	40MG/ML	N74403 001	May 23, 1996	Apr	DISC
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INJECTABLE; INJECTION

DOPAMINE HYDROCHLORIDE

@ INTL MEDICATION

40MG/ML

N18014 001

Apr DISC

@ SICOR PHARMS

40MG/ML

N72999 001 Oct 23, 1991 Apr DISC

@

80MG/ML

N73000 001 Oct 23, 1991 Apr DISC

DOXAZOSIN MESYLATE

TABLET; ORAL

DOXAZOSIN MESYLATE

AB ACTAVIS ELIZABETH

EQ 1MG BASE

N75574 001 Oct 18, 2000 Jun CAHN

AB

EQ 2MG BASE

N75574 002 Oct 18, 2000 Jun CAHN

AB

EQ 4MG BASE

N75574 003 Oct 18, 2000 Jun CAHN

AB

EQ 8MG BASE

N75574 004 Oct 18, 2000 Jun CAHN

DOXYCYCLINE

CAPSULE; ORAL

DOXYCYCLINE

@ PAR PHARM

EQ 75MG BASE

N65055 004 Apr 18, 2005 Mar DISC

@

EQ 150MG BASE

N65055 003 Jul 15, 2005 Mar DISC

RANBAXY

EQ 75MG BASE

N65053 003 Sep 10, 2003 Mar CTEC

CAPSULE, DELAYED RELEASE; ORAL

ORACEA

+ COLLAGENEX PHARMS

40MG

N50805 001 May 26, 2006 May NEWA

TABLET; ORAL

DOXYCYCLINE

AB MYLAN

EQ 50MG BASE

N65377 001 Nov 07, 2006 Oct NEWA

AB

EQ 75MG BASE

N65377 002 Nov 07, 2006 Oct NEWA

AB

EQ 100MG BASE

N65377 003 Nov 07, 2006 Oct NEWA

AB

PAR PHARM

EQ 75MG BASE

N65070 003 Dec 30, 2002 May CFTG

EQ 75MG BASE

N65070 003 Dec 30, 2002 Mar CMFD

AB

RANBAXY

EQ 50MG BASE

N65356 001 May 31, 2006 May NEWA

AB

EQ 75MG BASE

N65356 002 May 31, 2006 May NEWA

AB

EQ 100MG BASE

N65356 003 May 31, 2006 May NEWA

AB

SANDOZ

EQ 50MG BASE

N65353 001 Nov 27, 2006 Nov NEWA

AB

EQ 75MG BASE

N65353 002 Nov 27, 2006 Nov NEWA

AB

EQ 100MG BASE

N65353 003 Nov 27, 2006 Nov NEWA

DOXYCYCLINE HYCLATE

CAPSULE, DELAYED RELEASE; ORAL

DORYX

@ FH FAULDING CO LTD

EQ 75MG BASE

N50582 002 Aug 13, 2001 Apr DISC

@

EQ 100MG BASE

N50582 001 Jul 22, 1985 Apr DISC

@ MAYNE PHARMA INTL

EQ 75MG BASE

N50582 002 Aug 13, 2001 Sep CAHN

@

EQ 100MG BASE

N50582 001 Jul 22, 1985 Sep CAHN

@ WARNER CHILCOTT

EQ 100MG BASE

N62653 001 Oct 30, 1985 Apr DISC

DOXYCYCLINE HYCLATE

SANDOZ

EQ 75MG BASE

N65281 001 Dec 21, 2005 Apr CTEC

+

EQ 100MG BASE

N65281 002 Dec 21, 2005 Apr CRLD

INJECTABLE; INJECTION

DOXY 100

AP + ABRAXIS PHARM

EQ 100MG BASE/VIAL

N62475 001 Dec 09, 1983 Oct CTEC

DOXYCYCLINE

AP + BEDFORD

EQ 100MG BASE/VIAL

N62569 001 Mar 09, 1988 Oct CMFD

TABLET; ORAL

DOXYCYCLINE HYCLATE

AB PAR PHARM

EQ 20MG BASE

N65287 001 Feb 28, 2006 Feb NEWA

TABLET, DELAYED RELEASE; ORAL

DORYX

	MAYNE PHARMA INTL	EQ 75MG BASE	N50795 001	May 06, 2005	Oct	CAHN
+		EQ 100MG BASE	N50795 002	May 06, 2005	Oct	CAHN

DROSPIRENONE; ETHINYL ESTRADIOL

TABLET; ORAL

YAZ

+	BERLEX	3MG;0.02MG	N21676 001	Mar 16, 2006	Jun	CAHN
+	BERLEX LABS	3MG;0.02MG	N21676 001	Mar 16, 2006	Mar	NEWA

DYPHYLLINE

TABLET; ORAL

DILOR

	@ SAVAGE LABS	200MG	N84514 001		Jun	DISC
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DILOR-400

	@ SAVAGE LABS	400MG	N84751 001		Jun	DISC
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LUFYLLIN

	MEDPOINTE PHARM HLC	200MG	N84566 001		Jun	CTEC
+		400MG	N84566 002		Jun	CRLD

EFAVIRENZ; EMTRICITABINE; TENOFOVIR DISOPROXIL FUMARATE

TABLET; ORAL

ATRIPLA

+	GILEAD	600MG;200MG;300MG	N21937 001	Jul 12, 2006	Jul	NEWA
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EPINEPHRINE; LIDOCAINE HYDROCHLORIDE

INJECTABLE; INJECTION

XYLOCAINE W/ EPINEPHRINE

AP	+	ABRAXIS BIOSCIENCE	0.005MG/ML;0.5%	N06488 012		Jul	CAHN
AP	+		0.005MG/ML;1%	N06488 018	Nov 13, 1986	Jul	CAHN
AP	+		0.005MG/ML;1.5%	N06488 017		Jul	CAHN
AP	+		0.005MG/ML;2%	N06488 019	Nov 13, 1986	Jul	CAHN
AP	+		0.01MG/ML;1%	N06488 004		Jul	CAHN
		@	0.01MG/ML;2%	N06488 003		Jul	CAHN
		@	0.02MG/ML;2%	N06488 005		Jul	CAHN
	+	DENTSPLY PHARM	0.02MG/ML;2%	N21381 002		Mar	CRLD

PATCH; IONTOPHORESIS, TOPICAL

LIDOSITE TOPICAL SYSTEM KIT

+	VYTERIS	1.05MG/PATCH;100MG/PATCH	N21504 001	May 06, 2004	May	CDFR
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SOLUTION; IONTOPHORESIS, TOPICAL

LIDOCAINE HYDROCHLORIDE AND EPINEPHRINE

+	EMPI	0.01MG/ML;2%	N21486 001	Oct 26, 2004	May	CDFR
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EPIRUBICIN HYDROCHLORIDE

INJECTABLE; IV (INFUSION)

EPIRUBICIN HYDROCHLORIDE

+	MAYNE PHARMA USA	50MG/VIAL	N50807 001	Sep 15, 2006	Sep	NEWA
+		200MG/VIAL	N50807 002	Sep 15, 2006	Sep	NEWA

ERYTHROMYCIN

SOLUTION; TOPICAL

A/T/S

AT		TARO PHARMS NORTH	2%	N62405 001	Nov 18, 1982	Feb	CAHN
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SWAB; TOPICAL									
ERYTHROMYCIN									
AT	ALTANA	2%		N65320 001	Jul 25, 2006	Jul	NEWA		
<u>ERYTHROMYCIN ETHYLSUCCINATE</u>									
TABLET, CHEWABLE; ORAL									
E.E.S.									
	@ ABBOTT	EQ 200MG BASE		N50297 002		Aug	DISC		
ERYPED									
	@ ABBOTT	EQ 200MG BASE		N50297 003	Jul 05, 1988	Aug	DISC		
<u>ESCITALOPRAM OXALATE</u>									
TABLET; ORAL									
ESCITALOPRAM OXALATE									
AB	IVAX PHARMS	EQ 5MG BASE		N76765 001	May 22, 2006	Oct	CPOT		
AB		5MG		N76765 001	May 22, 2006	May	NEWA		
AB		EQ 10MG BASE		N76765 002	May 22, 2006	Oct	CPOT		
AB		10MG		N76765 002	May 22, 2006	May	NEWA		
AB		EQ 20MG BASE		N76765 003	May 22, 2006	Oct	CPOT		
AB		20MG		N76765 003	May 22, 2006	May	NEWA		
LEXAPRO									
AB	FOREST LABS	EQ 5MG BASE		N21323 001	Aug 14, 2002	Oct	CPOT		
AB		5MG		N21323 001	Aug 14, 2002	May	CFTG		
AB		EQ 10MG BASE		N21323 002	Aug 14, 2002	Oct	CPOT		
AB		10MG		N21323 002	Aug 14, 2002	May	CFTG		
AB	+	EQ 20MG BASE		N21323 003	Aug 14, 2002	Oct	CPOT		
AB	+	20MG		N21323 003	Aug 14, 2002	May	CFTG		
<u>ESOMEPRAZOLE MAGNESIUM</u>									
FOR SUSPENSION, DELAYED RELEASE; ORAL									
NEXIUM									
	ASTRAZENECA	EQ 20MG BASE/PACKET		N21957 001	Oct 20, 2006	Oct	NEWA		
	+	EQ 40MG BASE/PACKET		N21957 002	Oct 20, 2006	Oct	NEWA		
<u>ESTAZOLAM</u>									
TABLET; ORAL									
ESTAZOLAM									
AB	PAR PHARM	1MG		N74826 001	Jul 03, 1997	Apr	CAHN		
AB		2MG		N74826 002	Jul 03, 1997	Apr	CAHN		
<u>ESTRADIOL</u>									
FILM, EXTENDED RELEASE; TRANSDERMAL									
CLIMARA									
AB2	BERLEX	0.025MG/24HR		N20375 004	Mar 05, 1999	Aug	CRLD		
AB		0.0375MG/24HR		N20375 005	May 27, 2003	Aug	CRLD		
AB	+	0.0375MG/24HR		N20375 005	May 27, 2003	Jul	CFTG		
AB2		0.05MG/24HR		N20375 001	Dec 22, 1994	Aug	CRLD		
AB		0.06MG/24HR		N20375 006	May 27, 2003	Aug	CRLD		
AB	+	0.06MG/24HR		N20375 006	May 27, 2003	Jul	CFTG		
AB2		0.075MG/24HR		N20375 003	Mar 23, 1998	Aug	CRLD		
ESTRADERM									
BX	NOVARTIS	0.05MG/24HR		N19081 002	Sep 10, 1986	Aug	CRLD		
ESTRADIOL									
AB	MYLAN TECHNOLOGIES	0.0375MG/24HR		N75182 004	Jul 20, 2006	Jul	NEWA		
AB		0.06MG/24HR		N75182 005	Jul 20, 2006	Jul	NEWA		

FILM, EXTENDED RELEASE; TRANSDERMAL										
MENOSTAR										
	+	BERLEX	0.014MG/24HR		N21674	001	Jun 08, 2004	Jun	CAHN	
VIVELLE-DOT										
BX		NOVARTIS	0.025MG/24HR		N20538	009	May 03, 2002	Aug	CRLD	
BX			0.0375MG/24HR		N20538	005	Jan 08, 1999	Aug	CRLD	
AB1			0.05MG/24HR		N20538	006	Jan 08, 1999	Aug	CRLD	
BX			0.075MG/24HR		N20538	007	Jan 08, 1999	Aug	CRLD	
GEL; TOPICAL										
ESTROGEL										
	@	ASCEND	0.06%		N21166	001	Feb 09, 2004	Jan	CAHN	
>D>	GEL, METERED; TOPICAL									
>D>	ESTROGEL									
>D>	+	ASCEND	0.06%		N21166	002	Feb 09, 2004	Dec	CDFR	
	+		0.06%		N21166	002	Feb 09, 2004	Jan	CAHN	
GEL, METERED; TRANSDERMAL										
>A>	ELESTRIN									
>A>	BX	+	BIOSANTE	0.06%		N21813	001	Dec 15, 2006	Dec	NEWA
>A>	ESTROGEL									
>A>	BX	+	ASCEND	0.06%		N21166	002	Feb 09, 2004	Dec	CDFR
TABLET; ORAL										
ESTRADIOL										
	@	RADIUS PHARMS	0.5MG		N40275	001	Dec 29, 1998	Aug	CAHN	
	@		1MG		N40275	002	Dec 29, 1998	Aug	CAHN	
	@		2MG		N40275	003	Dec 29, 1998	Aug	CAHN	
<u>ESTRADIOL HEMIHYDRATE</u>										
EMULSION; TOPICAL										
ESTRASORB										
	+	ESPRIT PHARMA	0.25%		N21371	001	Oct 09, 2003	Feb	CAHN	
<u>ESTRADIOL; NORETHINDRONE ACETATE</u>										
TABLET; ORAL										
ACTIVELLA										
>A>		NOVO NORDISK INC	0.5MG;0.1MG		N20907	002	Dec 28, 2006	Dec	NEWA	
<u>ESTROGENS, CONJUGATED SYNTHETIC B</u>										
TABLET; ORAL										
ENJUVIA										
		DURAMED	0.3MG		N21443	001	Dec 20, 2004	Mar	CMFD	
			0.45MG		N21443	002	Dec 20, 2004	Mar	CMFD	
			0.625MG		N21443	003	May 10, 2004	Mar	CMFD	
	+		1.25MG		N21443	004	May 10, 2004	Mar	CMFD	
<u>ETHACRYNATE SODIUM</u>										
INJECTABLE; INJECTION										
EDECIN										
	+	ATON	EQ 50MG BASE/VIAL		N16093	001		Jul	CAHN	
<u>ETHACRYNIC ACID</u>										
TABLET; ORAL										
EDECIN										
	+	ATON	25MG		N16092	001		Oct	CAHN	
	@		50MG		N16092	002		Oct	CAHN	

ETHINYL ESTRADIOL; LEVONORGESTREL

TABLET; ORAL									
QUASENSE									
AB	WATSON LABS	0.03MG;0.15MG	N77101	001	Sep 06, 2006	Aug	NEWA		
SEASONALE									
AB	+ DURAMED	0.03MG;0.15MG	N21544	001	Sep 05, 2003	Aug	CFTG		
SEASONIQUE									
	+ DURAMED RES	0.03MG,0.01MG;0.15MG,N/A	N21840	001	May 25, 2006	May	NEWA		
TABLET; ORAL-28									
LEVONORGESTREL AND ETHINYL ESTRADIOL									
AB2	WATSON LABS	0.02MG;0.1MG	N77681	001	May 31, 2006	May	NEWA		

ETHINYL ESTRADIOL; NORETHINDRONE

TABLET; ORAL-21									
BALZIVA-21									
	BARR	0.035MG;0.4MG	N76198	001	Apr 22, 2004	Mar	CRLD		
TABLET; ORAL-28									
OVCON-35									
AB	+ WARNER CHILCOTT	0.035MG;0.4MG	N17716	001		Mar	CRLD		

ETHINYL ESTRADIOL; NORETHINDRONE ACETATE

TABLET; ORAL									
LOESTRIN 24 FE									
	+ WARNER CHILCOTT	0.02MG;1MG	N21871	001	Feb 17, 2006	Feb	NEWA		

ETHINYL ESTRADIOL; NORGESTIMATE

TABLET; ORAL-28									
NORGESTIMATE AND ETHINYL ESTRADIOL									
AB	WATSON LABS	0.035MG,0.035MG,0.035MG;0.18MG,0.215MG,0.25MG	N76626	001	Aug 17, 2006	Aug	NEWA		
AB		0.035MG,0.035MG,0.035MG;0.18MG,0.215MG,0.25MG	N76626	001	Aug 17, 2006	Aug	NEWA		
AB		0.035MG,0.035MG,0.035MG;0.18MG,0.215MG,0.25MG	N76626	001	Aug 17, 2006	Aug	NEWA		
AB		0.035MG;0.25MG	N76627	001	Aug 17, 2006	Aug	NEWA		

ETODOLAC

CAPSULE; ORAL									
ETODOLAC									
	@ ENDO PHARMS	200MG	N74842	001	Jul 17, 1997	Feb	DISC		
	@	300MG	N74842	002	Jul 17, 1997	Feb	DISC		
AB	+ TARO	300MG	N75078	002	Apr 30, 1998	Mar	CRLD		
LODINE									
	@ WYETH PHARMS INC	300MG	N18922	003	Jan 31, 1991	Mar	DISC		
TABLET; ORAL									
ETODOLAC									
AB	ACTAVIS ELIZABETH	400MG	N74819	001	Feb 28, 1997	Jun	CAHN		
	@ ENDO PHARMS	400MG	N74841	001	Jun 27, 1997	Feb	DISC		

ETONOGESTREL

IMPLANT; IMPLANTATION									
IMPLANON									
	+ ORGANON USA INC	68MG/IMPLANT	N21529	001	Jul 17, 2006	Jul	NEWA		

ETOPOSIDE

INJECTABLE; INJECTION
TOPOSAR

@ SICOR PHARMS 20MG/ML N74166 001 Feb 27, 1995 Jun DISC

FAMOTIDINE

TABLET; ORAL
FAMOTIDINE

AB ACTAVIS ELIZABETH 20MG N75650 001 Sep 14, 2001 Jun CAHN

AB 40MG N75650 002 Sep 14, 2001 Jun CAHN

FENOFIBRATE

CAPSULE; ORAL
ANTARA (MICRONIZED)
OSCIENT

43MG N21695 001 Nov 30, 2004 Aug CAHN

@ 87MG N21695 002 Nov 30, 2004 Aug CAHN

+ 130MG N21695 003 Nov 30, 2004 Aug CAHN

LIPOFEN

CIPHER 50MG N21612 001 Jan 11, 2006 Jan NEWA

100MG N21612 002 Jan 11, 2006 Jan NEWA

+ 150MG N21612 003 Jan 11, 2006 Jan NEWA

TABLET; ORAL
FENOFIBRATE

AB + TEVA 160MG N76433 002 May 13, 2005 Jan CRLD

TRICOR

@ ABBOTT 54MG N21203 001 Sep 04, 2001 Jan DISC

@ 160MG N21203 003 Sep 04, 2001 Jan DISC

FENOLDOPAM MESYLATE

INJECTABLE; INJECTION
FENOLDOPAM MESYLATE

AP SANDOZ EQ 10MG BASE/ML N77155 001 Feb 15, 2005 Jan CAHN

FENOPROFEN CALCIUM

TABLET; ORAL
FENOPROFEN CALCIUM

AB ACTAVIS ELIZABETH EQ 600MG BASE N72274 001 May 02, 1988 Jun CAHN

@ CLONMEL HLTHCARE EQ 600MG BASE N72326 001 Aug 17, 1988 Jan DISC

FENTANYL

FILM, EXTENDED RELEASE; TRANSDERMAL
FENTANYL

AB LAVIPHARM LABS 25UGM/HR N77051 001 Aug 04, 2006 Jul NEWA

AB 50UGM/HR N77051 002 Aug 04, 2006 Jul NEWA

AB 75UGM/HR N77051 003 Aug 04, 2006 Jul NEWA

AB 100UGM/HR N77051 004 Aug 04, 2006 Jul NEWA

FENTANYL CITRATE

TABLET; BUCCAL
FENTORA

CEPHALON EQ 0.1MG BASE N21947 001 Sep 25, 2006 Sep NEWA

EQ 0.2MG BASE N21947 002 Sep 25, 2006 Sep NEWA

+ EQ 0.4MG BASE N21947 003 Sep 25, 2006 Sep NEWA

EQ 0.6MG BASE N21947 004 Sep 25, 2006 Sep NEWA

TABLET; BUCCAL

FENTORA

CEPHALON

EQ 0.8MG BASE

N21947 005 Sep 25, 2006 Sep NEWA

TROCHE/LOZENGE; ORAL

FENTANYL

@ CEPHALON

EQ 0.1MG BASE

N20195 007 Oct 30, 1995 Jul CAHN

@

EQ 0.2MG BASE

N20195 001 Oct 04, 1993 Jul CAHN

@

EQ 0.3MG BASE

N20195 002 Oct 04, 1993 Jul CAHN

@

EQ 0.4MG BASE

N20195 003 Oct 04, 1993 Jul CAHN

TROCHE/LOZENGE; TRANSMUCOSAL

ACTIQ

CEPHALON

EQ 0.2MG BASE

N20747 001 Nov 04, 1998 Oct CTNA

+

EQ 0.4MG BASE

N20747 002 Nov 04, 1998 Oct CTNA

EQ 0.6MG BASE

N20747 003 Nov 04, 1998 Oct CTNA

EQ 0.8MG BASE

N20747 004 Nov 04, 1998 Oct CTNA

EQ 1.2MG BASE

N20747 005 Nov 04, 1998 Oct CTNA

EQ 1.6MG BASE

N20747 006 Nov 04, 1998 Oct CTNA

ACTIQ (SUGAR-FREE)

CEPHALON

EQ 0.2MG BASE

N20747 001 Nov 04, 1998 Mar CTNA

+

EQ 0.4MG BASE

N20747 002 Nov 04, 1998 Mar CTNA

EQ 0.6MG BASE

N20747 003 Nov 04, 1998 Mar CTNA

EQ 0.8MG BASE

N20747 004 Nov 04, 1998 Mar CTNA

EQ 1.2MG BASE

N20747 005 Nov 04, 1998 Mar CTNA

EQ 1.6MG BASE

N20747 006 Nov 04, 1998 Mar CTNA

FEXOFENADINE HYDROCHLORIDE

SUSPENSION; ORAL

ALLEGRA

+ SANOFI AVENTIS US

30MG/5ML

N21963 001 Oct 16, 2006 Oct NEWA

TABLET; ORAL

FEXOFENADINE HYDROCHLORIDE

AB DR REDDYS LABS LTD

30MG

N76502 001 Apr 11, 2006 Mar NEWA

AB

60MG

N76502 002 Apr 11, 2006 Mar NEWA

AB

180MG

N76502 003 Apr 11, 2006 Mar NEWA

FINASTERIDE

TABLET; ORAL

FINASTERIDE

AB DR REDDYS LABS INC

1MG

N76436 001 Jul 28, 2006 Jul NEWA

>A>

AB GEDEON RICHTER USA

5MG

N77251 001 Dec 22, 2006 Dec NEWA

AB IVAX PHARMS

5MG

N76340 001 Jun 19, 2006 Jun NEWA

>A>

AB MYLAN

5MG

N77578 001 Dec 18, 2006 Dec NEWA

>A>

AB TEVA

5MG

N76511 001 Dec 15, 2006 Dec NEWA

PROPECIA

AB + MERCK

1MG

N20788 001 Dec 19, 1997 Jul CFTG

PROSCAR

AB + MERCK

5MG

N20180 001 Jun 19, 1992 Jun CFTG

FLUCONAZOLE

FOR SUSPENSION; ORAL

FLUCONAZOLE

>A> AB TARO PHARM INDS

50MG/5ML

N76918 001 Dec 18, 2006 Dec NEWA

>A> AB

200MG/5ML

N76918 002 Dec 18, 2006 Dec NEWA

TABLET; ORAL

FLUCONAZOLE

@ GEDEON RICHTER USA

50MG

N76432 001 Jul 29, 2004 Nov DISC

INJECTABLE; INJECTION

FLUOROURACIL

AP	+	SICOR PHARMS	50MG/ML	N40333	001	Jan 27, 2000	Mar	CRLD
AP	+		50MG/ML	N40334	001	Feb 25, 2000	Mar	CRLD
AP	+	STERIS	50MG/ML	N87792	001	Oct 13, 1982	Mar	CRLD
		@ WATSON LABS	50MG/ML	N87792	001	Oct 13, 1982	Apr	DISC

FLUOXETINE HYDROCHLORIDE

SOLUTION; ORAL

FLUOXETINE

AA		ACTAVIS MID ATLANTIC	EQ 20MG BASE/5ML	N75690	001	Jan 31, 2002	Jul	CAHN
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TABLET; ORAL

FLUOXETINE HYDROCHLORIDE

	+	IVAX PHARMS	EQ 40MG BASE	N75865	003	Aug 30, 2004	Aug	CRLD
>D>		PROZAC						
>D>	AB	+	LILLY	EQ 10MG BASE	N20974	001	Mar 09, 1999	Dec DISC
>A>		@		EQ 10MG BASE	N20974	001	Mar 09, 1999	Dec DISC
		SARAFEM						
		WARNER CHILCOTT	EQ 10MG BASE	N21860	001	May 19, 2006	May	NEWA
			EQ 15MG BASE	N21860	002	May 19, 2006	May	NEWA
	+		EQ 20MG BASE	N21860	003	May 19, 2006	May	NEWA

FLUPHENAZINE HYDROCHLORIDE

CONCENTRATE; ORAL

FLUPHENAZINE HYDROCHLORIDE

	+	PHARM ASSOC	5MG/ML	N74725	001	Sep 16, 1996	Aug	CRLD
		@ TEVA PHARMS	5MG/ML	N73058	001	Aug 30, 1991	Aug	DISC
		PROLIXIN						
		@ APOTHECON	5MG/ML	N70533	001	Nov 07, 1985	Jul	DISC

ELIXIR; ORAL

FLUPHENAZINE HYDROCHLORIDE

	+	PHARM ASSOC	2.5MG/5ML	N40146	001	Aug 21, 1996	Apr	CRLD
		@ TEVA PHARMS	2.5MG/5ML	N81310	001	Apr 29, 1993	Apr	DISC
		PROLIXIN						
		@ APOTHECON	2.5MG/5ML	N12145	003		Apr	DISC

FLURBIPROFEN SODIUM

SOLUTION/DROPS; OPHTHALMIC

FLURBIPROFEN SODIUM

AT		BAUSCH AND LOMB	0.03%	N74447	001	Jan 04, 1995	Aug	CAHN
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FLUTAMIDE

CAPSULE; ORAL

EULEXIN

		@ SCHERING	125MG	N18554	001	Jan 27, 1989	May	DISC
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FLUTAMIDE

AB		PAR PHARM	125MG	N75298	001	Sep 18, 2001	Apr	CAHN
AB	+	SANDOZ	125MG	N75818	001	Sep 18, 2001	May	CRLD

FLUTICASONE PROPIONATE

AEROSOL, METERED; INHALATION

FLOVENT

		@ GLAXOSMITHKLINE	0.044MG/INH	N20548	001	Mar 27, 1996	Oct	DISC
		@	0.11MG/INH	N20548	002	Mar 27, 1996	Oct	DISC
		@	0.22MG/INH	N20548	003	Mar 27, 1996	Oct	DISC

CREAM; TOPICAL

AB	FLUTICASONE PROPIONATE G AND W LABS	0.05%	N77055 001	Jun 30, 2006	Jun	NEWA
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OINTMENT; TOPICAL

AB	FLUTICASONE PROPIONATE G AND W LABS	0.005%	N77168 001	Mar 03, 2006	Feb	NEWA
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SPRAY, METERED; NASAL

AB	FLONASE + GLAXOSMITHKLINE	0.05MG/SPRAY	N20121 001	Oct 19, 1994	Feb	CFTG
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AB	FLUTICASONE PROPIONATE ROXANE	0.05MG/SPRAY	N76504 001	Feb 22, 2006	Feb	NEWA
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FLUTICASONE PROPIONATE; SALMETEROL XINAFOATE

AEROSOL, METERED; INHALATION

ADVAIR HFA

+	GLAXOSMITHKLINE	0.045MG/INH;EQ 0.021MG BASE/INH	N21254 001	Jun 08, 2006	Jun	NEWA
+		0.115MG/INH;EQ 0.021MG BASE/INH	N21254 002	Jun 08, 2006	Jun	NEWA
+		0.23MG/INH;EQ 0.021MG BASE/INH	N21254 003	Jun 08, 2006	Jun	NEWA

FLUVOXAMINE MALEATE

TABLET; ORAL

AB	FLUVOXAMINE MALEATE ACTAVIS ELIZABETH	25MG	N75901 001	Dec 28, 2000	Jun	CAHN
AB		50MG	N75901 002	Dec 28, 2000	Jun	CAHN
AB		100MG	N75901 003	Dec 28, 2000	Jun	CAHN
AB	CARACO	25MG	N75900 001	Feb 23, 2006	Feb	NEWA
AB		50MG	N75900 002	Feb 23, 2006	Feb	NEWA
AB		100MG	N75900 003	Feb 23, 2006	Feb	NEWA

FOLLITROPIN ALFA/BETA

INJECTABLE; SUBCUTANEOUS

FOLLISTIM AQ

@	ORGANON USA INC	150 IU/0.18ML	N21211 003	Feb 11, 2004	Jul	DISC
+		300 IU/0.36ML	N21211 001	Mar 23, 2004	Jul	CMFD
@		300 IU/0.36ML	N21211 001	Mar 23, 2004	Jun	DISC

FORMOTEROL FUMARATE

POWDER; INHALATION

>A>	FORADIL CERTIHALER					
>A>	NOVARTIS	EQ 0.0085MG BASE/INH	N21592 001	Dec 15, 2006	Dec	NEWA

FOSCARNET SODIUM

INJECTABLE; INJECTION

AP	FOSCARNET SODIUM HOSPIRA	2.4GM/100ML	N77174 001	May 31, 2005	Feb	CAHN
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FOSINOPRIL SODIUM

TABLET; ORAL

AB	FOSINOPRIL SODIUM COBALT	10MG	N77531 001	Aug 31, 2006	Aug	NEWA
AB		20MG	N77531 002	Aug 31, 2006	Aug	NEWA
AB		40MG	N77531 003	Aug 31, 2006	Aug	NEWA

FOSINOPRIL SODIUM; HYDROCHLOROTHIAZIDE

TABLET; ORAL

FOSINOPRIL SODIUM AND HYDROCHLOROTHIAZIDE

AB	GENPHARM	10MG;12.5MG	N77705 001	Aug 14, 2006	Aug	NEWA
AB		20MG;12.5MG	N77705 002	Aug 14, 2006	Aug	NEWA
AB	TEVA	10MG;12.5MG	N76945 001	Jul 05, 2006	Jun	NEWA
AB		20MG;12.5MG	N76945 002	Jul 05, 2006	Jun	NEWA

FUROSEMIDE

TABLET; ORAL

FUROSEMIDE

AB	OHM LABS	20MG	N78010 001	Sep 18, 2006	Sep	NEWA
AB		40MG	N78010 002	Sep 18, 2006	Sep	NEWA
AB		80MG	N78010 003	Sep 18, 2006	Sep	NEWA

GABAPENTIN

CAPSULE; ORAL

GABAPENTIN

AB	ACTAVIS ELIZABETH	100MG	N75350 001	Sep 12, 2003	Jun	CAHN
AB		300MG	N75350 002	Sep 12, 2003	Jun	CAHN
AB		400MG	N75350 003	Sep 12, 2003	Jun	CAHN
AB	SANDOZ	100MG	N75428 001	Jan 24, 2006	Jan	NEWA
AB		300MG	N75428 002	Jan 24, 2006	Jan	NEWA
AB		400MG	N75428 003	Jan 24, 2006	Jan	NEWA
AB	SUN PHARM INDS LTD	100MG	N77242 001	Aug 24, 2006	Aug	NEWA
AB		300MG	N77242 002	Aug 24, 2006	Aug	NEWA
AB		400MG	N77242 003	Aug 24, 2006	Aug	NEWA

TABLET; ORAL

GABAPENTIN

AB	ACTAVIS ELIZABETH	600MG	N75694 001	Oct 21, 2004	Jun	CAHN
AB		800MG	N75694 002	Oct 21, 2004	Jun	CAHN
AB	APOTEX INC	100MG	N77894 001	Oct 10, 2006	Sep	NEWA
AB		300MG	N77894 002	Oct 10, 2006	Sep	NEWA
AB		400MG	N77894 003	Oct 10, 2006	Sep	NEWA
AB		600MG	N77661 004	Sep 13, 2006	Aug	NEWA
AB		800MG	N77661 005	Sep 13, 2006	Aug	NEWA
AB	GLENMARK PHARMS	600MG	N77662 001	Aug 18, 2006	Aug	NEWA
AB		800MG	N77662 002	Aug 18, 2006	Aug	NEWA
AB	IVAX PHARMS	100MG	N76017 001	Apr 28, 2004	Sep	CFTG
AB		300MG	N76017 002	Apr 28, 2004	Sep	CFTG
AB		400MG	N76017 003	Apr 28, 2004	Sep	CFTG
AB	SANDOZ	600MG	N76120 001	Jan 27, 2006	Jan	NEWA
AB		600MG	N76877 001	Jul 06, 2006	Jun	NEWA
AB		800MG	N76120 002	Jan 27, 2006	Jan	NEWA
AB		800MG	N76877 002	Jul 06, 2006	Jun	NEWA
AB	SUN PHARM INDS LTD	600MG	N77525 001	Aug 24, 2006	Aug	NEWA
AB		800MG	N77525 002	Aug 24, 2006	Aug	NEWA

GADOVERSETAMIDE

INJECTABLE; INJECTION

OPTIMARK

+	MALLINCKRODT	1654.5MG/5ML (330.9MG/ML)	N20937 001	Dec 08, 1999	Jan	CPOT
+		3309MG/10ML (330.9MG/ML)	N20937 002	Dec 08, 1999	Jan	NEWA
+		4963.5MG/15ML (330.9MG/ML)	N20937 003	Dec 08, 1999	Jan	NEWA
+		6618MG/20ML (330.9MG/ML)	N20937 004	Dec 08, 1999	Jan	NEWA

INJECTABLE; INJECTION

OPTIMARK

+	MALLINCKRODT	16.545GM/50ML (330.9MG/ML)	N20975 001	Dec 08, 1999	Jan	CPOT
OPTIMARK IN PLASTIC CONTAINER						
+	MALLINCKRODT	1654.5MG/5ML (330.9MG/ML)	N20976 001	Dec 08, 1999	Jan	CPOT
+		3309MG/10ML (330.9MG/ML)	N20976 002	Dec 08, 1999	Jan	NEWA
+		4963.5MG/15ML (330.9MG/ML)	N20976 003	Dec 08, 1999	Jan	NEWA
+		6618MG/20ML (330.9MG/ML)	N20976 004	Dec 08, 1999	Jan	NEWA

GALANTAMINE HYDROBROMIDE

CAPSULE, EXTENDED RELEASE; ORAL

RAZADYNE ER

+	JANSSEN	EQ 8MG BASE	N21615 001	Apr 01, 2005	May	CMS2
		EQ 16MG BASE	N21615 002	Apr 01, 2005	May	CMS2
		EQ 24MG BASE	N21615 003	Apr 01, 2005	May	CMS2

GANCICLOVIR

CAPSULE; ORAL

CYTOVENE

	@ ROCHE PALO	250MG	N20460 001	Dec 22, 1994	Jun	DISC
	@	500MG	N20460 002	Dec 12, 1997	Jun	DISC
GANCICLOVIR						
	RANBAXY	250MG	N76457 001	Jun 27, 2003	Jun	CTEC
+		500MG	N76457 002	Jun 27, 2003	Jun	CRLD

GATIFLOXACIN

INJECTABLE; INJECTION

TEQUIN

	@ BRISTOL MYERS SQUIBB	400MG/40ML(10MG/ML)	N21062 004	Dec 17, 1999	May	DISC
TEQUIN IN DEXTROSE 5% IN PLASTIC CONTAINER						
	@ BRISTOL MYERS SQUIBB	200MG/100ML(2MG/ML)	N21062 001	Dec 17, 1999	May	DISC
	@	400MG/200ML(2MG/ML)	N21062 002	Dec 17, 1999	May	DISC

SUSPENSION; ORAL

TEQUIN

	@ BRISTOL MYERS SQUIBB	200MG/5ML	N21678 001	Aug 27, 2004	Jun	DISC
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TABLET; ORAL

TEQUIN

	@ BRISTOL MYERS SQUIBB	200MG	N21061 001	Dec 17, 1999	May	DISC
	@	400MG	N21061 002	Dec 17, 1999	May	DISC

GEMFIBROZIL

TABLET; ORAL

GEMFIBROZIL

AB	INVAGEN PHARMS	600MG	N77836 001	Jul 27, 2006	Jul	NEWA
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GENTAMICIN SULFATE

INJECTABLE; INJECTION

GENTAMICIN SULFATE IN SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER

AP	+	B BRAUN	EQ 0.8MG BASE/ML	N62814 001	Aug 28, 1987	Nov	CRLD
AP	+		EQ 1.2MG BASE/ML	N62814 002	Aug 28, 1987	Nov	CRLD
AP	+		EQ 1.4MG BASE/ML	N62814 003	Aug 28, 1987	Nov	CRLD
AP	+		EQ 1.6MG BASE/ML	N62814 004	Aug 28, 1987	Nov	CRLD
AP	+		EQ 1.8MG BASE/ML	N62814 005	Aug 28, 1987	Nov	CRLD
AP	+		EQ 2MG BASE/ML	N62814 006	Aug 28, 1987	Nov	CRLD
AP	+		EQ 2.4MG BASE/ML	N62814 007	Aug 28, 1987	Nov	CRLD

INJECTABLE; INJECTION

GENTAMICIN SULFATE IN SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER

AP	+	B BRAUN	EQ 40MG BASE/100ML	N62814 008	Aug 28, 1987	Nov	CRLD
AP	+		EQ 60MG BASE/100ML	N62814 009	Aug 28, 1987	Nov	CRLD
AP	+		EQ 70MG BASE/100ML	N62814 010	Aug 28, 1987	Nov	CRLD
AP	+		EQ 80MG BASE/100ML	N62814 011	Aug 28, 1987	Nov	CRLD
AP	+		EQ 90MG BASE/100ML	N62814 012	Aug 28, 1987	Nov	CRLD
AP	+		EQ 100MG BASE/100ML	N62814 013	Aug 28, 1987	Nov	CRLD
AP	+		EQ 120MG BASE/100ML	N62814 014	Aug 28, 1987	Nov	CRLD

SOLUTION/DROPS; OPHTHALMIC

GENTACIDIN

@ NOVARTIS

EQ 0.3% BASE

N62480 001 Mar 30, 1984 Jun DISC

GLIMEPIRIDE

TABLET; ORAL

GLIMEPIRIDE

AB		COBALT	1MG	N77280 001	Feb 03, 2006	Jan	NEWA
AB			2MG	N77280 002	Feb 03, 2006	Jan	NEWA
AB			4MG	N77280 003	Feb 03, 2006	Jan	NEWA
AB		GENPHARM	1MG	N77486 001	Feb 10, 2006	Jan	NEWA
AB			2MG	N77486 002	Feb 10, 2006	Jan	NEWA
AB			4MG	N77486 003	Feb 10, 2006	Jan	NEWA

GLIMEPIRIDE; PIOGLITAZONE HYDROCHLORIDE

TABLET; ORAL

DUETACT

TAKEDA GLOBAL

2MG;30MG

N21925 001 Jul 28, 2006 Jul NEWA

+

4MG;30MG

N21925 002 Jul 28, 2006 Sep CRLD

4MG;30MG

N21925 002 Jul 28, 2006 Jul NEWA

GLIPIZIDE

TABLET; ORAL

GLIPIZIDE

AB		CARACO	5MG	N77820 001	Jul 11, 2006	Jun	NEWA
AB			10MG	N77820 002	Jul 11, 2006	Jun	NEWA
		@ ENDO PHARMS	5MG	N74378 001	Nov 28, 1994	Feb	DISC
		@	10MG	N74378 002	Nov 28, 1994	Feb	DISC

TABLET, EXTENDED RELEASE; ORAL

GLIPIZIDE

AB		WATSON LABS	2.5MG	N76467 003	Mar 27, 2006	Mar	NEWA
		GLUCOTROL XL					
AB		PFIZER	2.5MG	N20329 003	Aug 10, 1999	Mar	CFTG

GLYBURIDE

TABLET; ORAL

DIABETA

BX	+	SANOFI AVENTIS US	5MG	N17532 003	May 01, 1984	Feb	CRLD
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GLYBURIDE; METFORMIN HYDROCHLORIDE

TABLET; ORAL

GLYBURIDE AND METFORMIN HYDROCHLORIDE

AB		ACTAVIS ELIZABETH	1.25MG;250MG	N76716 001	Jun 28, 2005	Jun	CAHN
AB			2.5MG;500MG	N76716 002	Jun 28, 2005	Jun	CAHN
AB			5MG;500MG	N76716 003	Jun 28, 2005	Jun	CAHN

GLYCOPYRROLATE

TABLET; ORAL

GLYCOPYRROLATE

AA	KALI LABS	1MG	N40653 001	Aug 31, 2006	Aug	NEWA
AA		2MG	N40653 002	Aug 31, 2006	Aug	NEWA
ROBINUL						
AA	+ SCIELE PHARMA INC	1MG	N12827 001		Jun	CAHN
ROBINUL FORTE						
AA	+ SCIELE PHARMA INC	2MG	N12827 002		Jun	CAHN

GUANFACINE HYDROCHLORIDE

TABLET; ORAL

TENEX

AB	DR REDDYS LABS INC	EQ 1MG BASE	N19032 001	Oct 27, 1986	May	CAHN
AB	+ @	EQ 2MG BASE	N19032 002	Nov 07, 1988	May	CAHN
		EQ 3MG BASE	N19032 003	Nov 07, 1988	May	CAHN

HALOBETASOL PROPIONATE

CREAM; TOPICAL

HALOBETASOL PROPIONATE

AB	PERRIGO ISRAEL	0.05%	N77123 001	Dec 16, 2004	Apr	CAHN
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OINTMENT; TOPICAL

HALOBETASOL PROPIONATE

AB	ACTAVIS MID ATLANTIC	0.05%	N77109 001	Jun 14, 2005	Jun	CAHN
AB	G AND W LABS	0.05%	N77721 001	Sep 07, 2006	Aug	NEWA
AB	PERRIGO	0.05%	N76872 001	Dec 16, 2004	Mar	CAHN

HALOPERIDOL DECANOATE

INJECTABLE; INJECTION

HALOPERIDOL DECANOATE

AO	SANDOZ	EQ 50MG BASE/ML	N76463 001	Jun 24, 2005	Jan	CAHN
AO		EQ 100MG BASE/ML	N76463 002	Jun 24, 2005	Jan	CAHN

HALOPERIDOL LACTATE

INJECTABLE; INJECTION

HALOPERIDOL

>D>	AP	APOTEX INC	EQ 5MG BASE/ML	N76774 001	Aug 25, 2004	Dec	CAHN
>A>	AP	GLAND PHARMA LTD	EQ 5MG BASE/ML	N76774 001	Aug 25, 2004	Dec	CAHN
	AP	SANDOZ	EQ 5MG BASE/ML	N76464 001	Sep 29, 2004	Jan	CAHN

SOLUTION; ORAL

HALOPERIDOL LACTATE

ACTAVIS MID ATLANTIC EQ 1MG BASE/ML

N74536 001 Nov 02, 1995 Jul CAHN

>D> HALOTHANE

>D> LIQUID; INHALATION

>D> HALOTHANE

>D>	AN	HALOCARBON	99.99%	N80810 001		Dec	DISC
>A>		@	99.99%	N80810 001		Dec	DISC
		@ HOSPIRA	99.99%	N83254 001		Aug	DISC

HOMATROPINE METHYLBROMIDE; HYDROCODONE BITARTRATE

SYRUP; ORAL

HYDROCODONE COMPOUND

AA	ACTAVIS MID ATLANTIC	1.5MG/5ML;5MG/5ML	N88017 001	Jul 05, 1983	Jul	CAHN
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TABLET; ORAL

TUSSIGON

>D>	AA	JONES PHARMA	1.5MG;5MG	N88508 001	Jul 30, 1985	Dec	CAHN
>A>	AA	KING PHARMS	1.5MG;5MG	N88508 001	Jul 30, 1985	Dec	CAHN

HYDRALAZINE HYDROCHLORIDE

INJECTABLE; INJECTION

HYDRALAZINE HYDROCHLORIDE

@	SICOR PHARMS	20MG/ML	N40373 001	Feb 23, 2000	Aug	DISC
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TABLET; ORAL

APRESOLINE

@	NOVARTIS	10MG	N08303 004		Feb	DISC
@		25MG	N08303 001		Feb	DISC
@		50MG	N08303 002		Feb	DISC
@		100MG	N08303 005		Feb	DISC

HYDRALAZINE HYDROCHLORIDE

AA	+	PLIVA	10MG	N89097 001	Dec 18, 1985	Feb	CRLD
AA	+		25MG	N88467 001	May 01, 1984	Feb	CRLD
AA	+		50MG	N88468 001	May 01, 1984	Feb	CRLD
AA	+		100MG	N89098 001	Dec 18, 1985	Feb	CRLD
	@	RADIUS PHARMS	25MG	N86243 001		Aug	CAHN
	@		50MG	N86242 002		Aug	CAHN
	@	TG UNITED LABS	10MG	N88846 001	Feb 26, 1985	May	CAHN
	@		25MG	N88847 001	Feb 26, 1985	May	CAHN
	@		50MG	N88848 001	Feb 26, 1985	May	CAHN
	@		100MG	N88849 001	Feb 26, 1985	May	CAHN

HYDRALAZINE HYDROCHLORIDE; HYDROCHLOROTHIAZIDE; RESERPINE

TABLET; ORAL

CAM-AP-ES

@	TG UNITED LABS	25MG;15MG;0.1MG	N84897 001		May	CAHN
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HYDROCHLOROTHIAZIDE

TABLET; ORAL

HYDROCHLOROTHIAZIDE

AB		ACTAVIS ELIZABETH	25MG	N85054 002		Jun	CAHN
AB			50MG	N85208 001		Jun	CAHN
	@	TG UNITED LABS	25MG	N85683 001		May	CAHN
	@		50MG	N83965 001		May	CAHN
AB		WEST WARD	25MG	N84878 002	Jul 12, 2006	Jun	NEWA

HYDROCHLOROTHIAZIDE; LISINOPRIL

TABLET; ORAL

LISINOPRIL AND HYDROCHLOROTHIAZIDE

AB		ACTAVIS ELIZABETH	12.5MG;10MG	N76230 001	Jul 01, 2002	Jun	CAHN
AB			12.5MG;20MG	N76230 002	Jul 01, 2002	Jun	CAHN
AB			25MG;20MG	N76230 003	Jul 01, 2002	Jun	CAHN
AB		AUROBINDO	12.5MG;10MG	N77606 001	Mar 14, 2006	Feb	NEWA
AB			12.5MG;20MG	N77606 002	Mar 14, 2006	Feb	NEWA
AB			25MG;20MG	N77606 003	Mar 14, 2006	Feb	NEWA
AB		LUPIN	12.5MG;10MG	N77912 001	Sep 27, 2006	Sep	NEWA
AB			12.5MG;20MG	N77912 002	Sep 27, 2006	Sep	NEWA
AB			25MG;20MG	N77912 003	Sep 27, 2006	Sep	NEWA

HYDROCHLOROTHIAZIDE; METHYLDOPA

TABLET; ORAL

ALDORIL 15

@ MERCK	15MG;250MG	N13402 001		Jun	DISC
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ALDORIL 25

@ MERCK	25MG;250MG	N13402 002		Jun	DISC
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ALDORIL D30

@ MERCK	30MG;500MG	N13402 003		Jun	DISC
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ALDORIL D50

@ MERCK	50MG;500MG	N13402 004		Jun	DISC
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METHYLDOPA AND HYDROCHLOROTHIAZIDE

MYLAN

15MG;250MG	N70264 001	Jan 23, 1986	Jun	CTEC
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+

25MG;250MG	N70265 001	Jan 23, 1986	Jun	CRLD
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@ PAR PHARM

15MG;250MG	N70616 001	Feb 02, 1987	Jun	DISC
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@

25MG;250MG	N70612 001	Feb 02, 1987	Jun	DISC
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@

30MG;500MG	N70613 001	Feb 02, 1987	Jun	DISC
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@

50MG;500MG	N70614 001	Feb 02, 1987	Jun	DISC
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@ SANDOZ

15MG;250MG	N70182 001	Jan 15, 1986	Jun	DISC
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@

25MG;250MG	N70183 001	Jan 15, 1986	Jun	DISC
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@

30MG;500MG	N70543 001	Jan 15, 1986	Jun	DISC
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@

50MG;500MG	N70544 001	Jan 15, 1986	Jun	DISC
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@ WATSON LABS

15MG;250MG	N70958 001	Feb 06, 1989	Jun	DISC
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@

25MG;250MG	N70959 001	Jan 19, 1989	Jun	DISC
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@

30MG;500MG	N71069 001	Jan 19, 1989	Jun	DISC
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@

50MG;500MG	N70960 001	Feb 06, 1989	Jun	DISC
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HYDROCHLOROTHIAZIDE; METOPROLOL SUCCINATE

TABLET, EXTENDED RELEASE; ORAL

DUTOPROL

ASTRAZENECA

12.5MG;EQ 25MG TARTRATE	N21956 001	Aug 28, 2006	Aug	NEWA
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12.5MG;EQ 50MG TARTRATE

N21956 002	Aug 28, 2006	Aug	NEWA
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+

12.5MG;EQ 100MG TARTRATE	N21956 003	Aug 28, 2006	Aug	NEWA
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HYDROCHLOROTHIAZIDE; PROPRANOLOL HYDROCHLORIDE

TABLET; ORAL

PROPRANOLOL HYDROCHLORIDE AND HYDROCHLOROTHIAZIDE

AB	ACTAVIS ELIZABETH	25MG;40MG	N70851 001	May 15, 1986	Jun	CAHN
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AB		25MG;80MG	N70852 001	May 15, 1986	Jun	CAHN
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HYDROCHLOROTHIAZIDE; TIMOLOL MALEATE

TABLET; ORAL

TIMOLIDE 10-25

@ MERCK	25MG;10MG	N18061 001		Jun	DISC
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HYDROCHLOROTHIAZIDE; TRIAMTERENE

TABLET; ORAL

TRIAMTERENE AND HYDROCHLOROTHIAZIDE

AB	APOTEX INC	25MG;37.5MG	N71251 002	May 05, 1998	Nov	CAHN
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AB		50MG;75MG	N71251 001	Apr 17, 1988	Nov	CAHN
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HYDROCHLOROTHIAZIDE; VALSARTAN

TABLET; ORAL

DIOVAN HCT

NOVARTIS

12.5MG;320MG	N20818 004	Apr 28, 2006	Apr	NEWA
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TABLET; ORAL

DIOVAN HCT

NOVARTIS

25MG;160MG

N20818 003 Jan 17, 2002 Apr CRLD

+

25MG;320MG

N20818 005 Apr 28, 2006 Apr NEWA

HYDROCODONE BITARTRATE; IBUPROFEN

TABLET; ORAL

HYDROCODONE BITARTRATE AND IBUPROFEN

AB	+	INTERPHARM	5MG;200MG	N76642 002	Mar 18, 2004	Oct	CTEC
AB		VINTAGE PHARMS	5MG;200MG	N77727 001	Nov 06, 2006	Oct	NEWA
AB			7.5MG;200MG	N77723 001	Nov 06, 2006	Oct	NEWA
			10MG;200MG	N77723 002	Nov 06, 2006	Oct	NEWA

HYDROCORTISONE

CREAM; TOPICAL

HYDROCORTISONE

AT		ACTAVIS MID ATLANTIC	1%	N87795 001	May 03, 1983	Jun	CAHN
AT			2.5%	N89682 001	Mar 10, 1988	Jun	CAHN

PENECORT

@ ALLERGAN HERBERT

1%

N88216 001 Jun 06, 1984 Oct DISC

OINTMENT; TOPICAL

HYDROCORTISONE

AT		ACTAVIS MID ATLANTIC	1%	N87796 001	Oct 13, 1982	Jun	CAHN
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POWDER; FOR RX COMPOUNDING

HYDRO-RX

+		X GEN PHARMS	100%	N85982 001		Jun	CTNA
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SOLUTION; TOPICAL

PENECORT

@ ALLERGAN HERBERT

1%

N88214 001 Jun 06, 1984 Oct DISC

TABLET; ORAL

CORTEF

@ PHARMACIA AND UPJOHN

10MG

N08697 001 Jun CTEC

HYDROCORTONE

@ MERCK

10MG

N08506 007 Jun DISC

@

20MG

N08506 011 Jun DISC

HYDROCORTISONE SODIUM SUCCINATE

INJECTABLE; INJECTION

A-HYDROCORT

AP		HOSPIRA	EQ 100MG BASE/VIAL	N40666 001	Apr 06, 2006	Mar	NEWA
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HYDROCORTISONE; NEOMYCIN; POLYMYXIN B SULFATE

SUSPENSION/DROPS; OTIC

NEOMYCIN AND POLYMYXIN B SULFATES AND HYDROCORTISONE

AT		PHARMAFORCE	1%;EQ 3.5MG BASE;10,000 UNITS	N65219 001	May 01, 2006	Apr	NEWA
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HYDROXOCOBALAMIN

INJECTABLE; INJECTION

CYANOKIT

>A>

>A>

+

EMD PHARMS

2.5GM/VIAL (5GM/KIT)

N22041 002 Dec 15, 2006 Dec NEWA

HYDROXYPROPYL CELLULOSE

INSERT; OPHTHALMIC

LACRISERT

+		ATON	5MG	N18771 001		Oct	CAHN
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HYDROXYZINE HYDROCHLORIDE

SYRUP; ORAL

ATARAX

@ ROERIG

10MG/5ML

N10485 001

Jun DISC

HYDROXYZINE HYDROCHLORIDE

AA + ACTAVIS MID ATLANTIC 10MG/5ML

N86880 001

Jul CAHN

AA + ALPHARMA US PHARMS 10MG/5ML

N86880 001

Jun CRLD

AA + HI TECH PHARMA 10MG/5ML

N40010 001 Oct 28, 1994 Jun CRLD

AA + MORTON GROVE 10MG/5ML

N87294 001 Apr 12, 1982 Jun CRLD

AA + VINTAGE PHARMS 10MG/5ML

N40391 001 Apr 10, 2002 Jun CRLD

HYDROXYZINE PAMOATE

CAPSULE; ORAL

HYDROXYZINE PAMOATE

BARR

EQ 100MG HCL

N88488 001 Jun 15, 1984 Mar CTEC

VISTARIL

@ PFIZER

EQ 100MG HCL

N11459 006

Mar DISC

IBANDRONATE SODIUM

INJECTABLE; INTRAVENOUS

BONIVA

+ ROCHE EQ 3MG BASE/3ML

N21858 001 Jan 06, 2006 Jan NEWA

IBUPROFEN

SUSPENSION; ORAL

IBUPROFEN

AB ACTAVIS MID ATLANTIC 100MG/5ML

N74978 001 Mar 25, 1998 Jul CAHN

TABLET; ORAL

IBU

@ BASF

400MG

N18197 001

Jun DISC

@

400MG

N70083 001 Feb 22, 1985 Jun DISC

@

600MG

N70099 001 Mar 29, 1985 Jun DISC

@

800MG

N70745 001 Jul 23, 1986 Jun DISC

IBUPROFEN LYSINE

INJECTABLE; INTRAVENOUS

NEOPROFEN

+ FARMACON IL EQ 20MG BASE/2ML (EQ 10MG BASE/ML)

N21903 001 Apr 13, 2006 Apr NEWA

IDARUBICIN HYDROCHLORIDE

INJECTABLE; INJECTION

IDARUBICIN HYDROCHLORIDE

>A> AP BEDFORD LABS 1MG/ML

N65275 001 Dec 14, 2006 Dec NEWA

IMATINIB MESYLATE

CAPSULE; ORAL

GLEEVEC

@ NOVARTIS

EQ 50MG BASE

N21335 001 May 10, 2001 Nov CPOT

@

EQ 100MG BASE

N21335 002 May 10, 2001 Nov CPOT

TABLET; ORAL

GLEEVEC

NOVARTIS

EQ 100MG BASE

N21588 001 Apr 18, 2003 Nov CPOT

+ EQ 400MG BASE

N21588 002 Apr 18, 2003 Nov CPOT

INDAPAMIDE

TABLET; ORAL

INDAPAMIDE

AB	ACTAVIS ELIZABETH	1.25MG	N74722 001	Jun 17, 1996	Jun	CAHN
AB		2.5MG	N74722 002	Jun 17, 1996	Jun	CAHN

INDIUM IN-111 OXYQUINOLINE

INJECTABLE; INJECTION

INDIUM IN-111 OXYQUINOLINE

+	GE HEALTHCARE	1mCi/ML	N19044 001	Dec 24, 1985	Aug	CRLD
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INDIUM IN-111 PENTETATE DISODIUM

INJECTABLE; INTRATHECAL

MPI INDIUM DTPA IN 111

+	GE HEALTHCARE	1mCi/ML	N17707 001	Feb 18, 1982	Aug	CRLD
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INDIUM IN-111 PENTETREOTIDE KIT

INJECTABLE; INJECTION

OCTREOSCAN

+	MALLINCKRODT	3mCi/ML	N20314 001	Jun 02, 1994	Aug	CRLD
---	--------------	---------	------------	--------------	-----	------

INDOMETHACIN

CAPSULE; ORAL

INDOCIN

@	MERCK	25MG	N16059 001		Jun	DISC
---	-------	------	------------	--	-----	------

@		50MG	N16059 002		Jun	DISC
---	--	------	------------	--	-----	------

INDO-LEMMON

@	TEVA	25MG	N70266 001	Nov 07, 1985	Jun	DISC
---	------	------	------------	--------------	-----	------

@		50MG	N70267 001	Nov 07, 1985	Jun	DISC
---	--	------	------------	--------------	-----	------

INDOMETHACIN

@	CLONMEL HLTHCARE	25MG	N18851 001	May 18, 1984	Jun	DISC
---	------------------	------	------------	--------------	-----	------

@		50MG	N18851 002	May 18, 1984	Jun	DISC
---	--	------	------------	--------------	-----	------

@	IVAX PHARMS	25MG	N70719 001	Feb 12, 1986	Jun	DISC
---	-------------	------	------------	--------------	-----	------

@		50MG	N70756 001	Feb 12, 1986	Jun	DISC
---	--	------	------------	--------------	-----	------

@	MUTUAL PHARM	25MG	N70899 001	Feb 09, 1987	Jun	DISC
---	--------------	------	------------	--------------	-----	------

@		50MG	N70900 001	Feb 09, 1987	Jun	DISC
---	--	------	------------	--------------	-----	------

	MYLAN	25MG	N18858 001	Apr 20, 1984	Jun	CTEC
--	-------	------	------------	--------------	-----	------

AB	+		50MG	N18858 002	Apr 20, 1984	Jun	CRLD
----	---	--	------	------------	--------------	-----	------

@	PAR PHARM	25MG	N18829 002	Aug 06, 1984	Jun	DISC
---	-----------	------	------------	--------------	-----	------

@		50MG	N18829 001	Aug 06, 1984	Jun	DISC
---	--	------	------------	--------------	-----	------

@		50MG	N70651 001	Mar 05, 1986	Jun	DISC
---	--	------	------------	--------------	-----	------

@	PLIVA	25MG	N71148 001	Mar 18, 1987	Jun	DISC
---	-------	------	------------	--------------	-----	------

@		50MG	N71149 001	Mar 18, 1987	Jun	DISC
---	--	------	------------	--------------	-----	------

@	RADIUS PHARMS	25MG	N18851 001	May 18, 1984	Aug	CAHN
---	---------------	------	------------	--------------	-----	------

@		50MG	N18851 002	May 18, 1984	Aug	CAHN
---	--	------	------------	--------------	-----	------

@	SANDOZ	25MG	N70673 001	Apr 29, 1987	Jun	DISC
---	--------	------	------------	--------------	-----	------

@		50MG	N70674 001	Apr 29, 1987	Jun	DISC
---	--	------	------------	--------------	-----	------

@	TEVA	25MG	N71342 001	Apr 18, 1988	Jun	DISC
---	------	------	------------	--------------	-----	------

@		50MG	N71343 001	Apr 18, 1988	Jun	DISC
---	--	------	------------	--------------	-----	------

CAPSULE, EXTENDED RELEASE; ORAL

INDOCIN SR

+	SANDOZ	75MG	N74464 001	May 28, 1998	Jun	CTEC
---	--------	------	------------	--------------	-----	------

INDOMETHACIN

@	INWOOD LABS	75MG	N72410 001	Mar 15, 1989	Jun	DISC
---	-------------	------	------------	--------------	-----	------

SUPPOSITORY; RECTAL
INDOMETHACIN

+ G AND W LABS 50MG N73314 001 Aug 31, 1992 Apr CTNA

INSULIN GLULISINE RECOMBINANT

INJECTABLE; SUBCUTANEOUS
APIDRA

+ SANOFI AVENTIS US 1000 UNITS/10ML (100 UNITS/ML) N21629 001 Apr 16, 2004 Mar CMFD

+ 300 UNITS/3ML (100 UNITS/ML) N21629 002 Dec 20, 2005 Mar CMFD

INSULIN PURIFIED PORK

INJECTABLE; INJECTION
ILETIN II

@ LILLY 500 UNITS/ML N18344 002 Jun DISC

INSULIN RECOMBINANT HUMAN

POWDER; INHALATION
EXUBERA

PFIZER 1MG/INH N21868 001 Jan 27, 2006 Jan NEWA

+ 3MG/INH N21868 002 Jan 27, 2006 Jan NEWA

IPRATROPIUM BROMIDE

AEROSOL, METERED; INHALATION
ATROVENT

@ BOEHRINGER INGELHEIM 0.018MG/INH N19085 001 Dec 29, 1986 May DISC

SOLUTION; INHALATION
ATROVENT

@ BOEHRINGER INGELHEIM 0.02% N20228 001 Sep 29, 1993 May DISC

IPRATROPIUM BROMIDE

AN ACTAVIS MID ATLANTIC 0.02% N75111 001 Apr 22, 1999 Jul CAHN

AN + DEY 0.02% N74755 001 Jan 10, 1997 May CRLD

@ ROXANE 0.02% N75867 001 Jul 22, 2002 May DISC

ISONIAZID

INJECTABLE; INJECTION
ISONIAZID

AP SANDOZ 100MG/ML N40648 001 Jul 05, 2005 Jan CAHN

ISOSORBIDE MONONITRATE

TABLET; ORAL
ISMO

AB DR REDDYS LABS INC 20MG N19091 001 Dec 30, 1991 May CAHN

ISOSORBIDE MONONITRATE

AB ACTAVIS ELIZABETH 10MG N75037 002 Oct 30, 1998 Jun CAHN

AB 20MG N75037 001 Oct 30, 1998 Jun CAHN

TABLET, EXTENDED RELEASE; ORAL
ISOSORBIDE MONONITRATE

AB ACTAVIS ELIZABETH 30MG N75306 001 Dec 31, 1998 Jun CAHN

AB 60MG N75306 002 Dec 31, 1998 Jun CAHN

AB WEST WARD 30MG N76813 002 Mar 30, 2006 Mar NEWA

ISOTRETINOIN

CAPSULE; ORAL
CLARAVIS

AB BARR 30MG N76135 003 May 11, 2006 May NEWA

CAPSULE; ORAL
SOTRET

AB	RANBAXY	30MG	N76503 001	Jun 20, 2003	May	CTEC
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ISRADIPINECAPSULE; ORAL
DYNACIRC

@ RELIANT PHARMS

2.5MG

N19546 001 Dec 20, 1990 May DISC

ISRADIPINE

AB + ABRIKA PHARMS

5MG

N77317 002 Jan 05, 2006 Jun CRLD

AB 5MG

N77317 002 Jan 05, 2006 Apr CTEC

AB AMIDE PHARM 2.5MG

N77169 001 Apr 24, 2006 Apr NEWA

AB 5MG

N77169 002 Apr 24, 2006 Apr NEWA

IVERMECTINTABLET; ORAL
STROMECTOL

+ MERCK

3MG

N50742 002 Oct 08, 1998 Jun CRLD

@ 6MG

N50742 001 Nov 22, 1996 Jun DISC

KETOCONAZOLEGEL; TOPICAL
XOLEGEL

+ BARRIER THERAP

2%

N21946 001 Jul 28, 2006 Jul NEWA

KETOPROFENCAPSULE; ORAL
KETOPROFEN

AB RADIUS PHARMS

25MG

N74014 001 Jan 29, 1993 Jul CAHN

AB 50MG

N74014 002 Jan 29, 1993 Jul CAHN

AB 75MG

N74014 003 Jan 29, 1993 Jul CAHN

KETOROLAC TROMETHAMINE

INJECTABLE; INJECTION

KETOROLAC TROMETHAMINE

>D> AP APOTEX INC 15MG/ML

N76722 001 Jul 27, 2004 Dec CAHN

>D> AP 30MG/ML

N76722 002 Jul 27, 2004 Dec CAHN

>A> AP GLAND PHARMA LTD 15MG/ML

N76722 001 Jul 27, 2004 Dec CAHN

>A> AP 30MG/ML

N76722 002 Jul 27, 2004 Dec CAHN

AP SANDOZ 15MG/ML

N76271 001 Oct 06, 2004 Jan CAHN

AP 30MG/ML

N76271 002 Oct 06, 2004 Jan CAHN

KETOTIFEN FUMARATESOLUTION/DROPS; OPHTHALMIC
KETOTIFEN FUMARATE

AT APOTEX EQ 0.025% BASE

N77354 001 May 09, 2006 Apr NEWA

APOTEX INC EQ 0.025% BASE

N77354 001 May 09, 2006 Oct CTEC

KUNECATECHINSOINTMENT; TOPICAL
VEREGEN

+ MEDIGENE 15%

N21902 001 Oct 31, 2006 Oct NEWA

LACTULOSESOLUTION; ORAL
CONSTULOSE

AA + ACTAVIS MID ATLANTIC 10GM/15ML N70288 001 Aug 15, 1988 Jul CAHN

SOLUTION; ORAL, RECTAL
ENULOSE

AA + ACTAVIS MID ATLANTIC 10GM/15ML N71548 001 Aug 15, 1988 Jul CAHN

LAMOTRIGINETABLET; ORAL
LAMICTAL

AB + GLAXOSMITHKLINE 25MG N20241 005 Dec 27, 1994 Aug CFTG
 AB 100MG N20241 001 Dec 27, 1994 Aug CFTG
 AB 150MG N20241 002 Dec 27, 1994 Aug CFTG
 AB 200MG N20241 003 Dec 27, 1994 Aug CFTG

LAMOTRIGINE

AB TEVA 25MG N76388 001 Aug 30, 2006 Aug NEWA
 AB 100MG N76388 002 Aug 30, 2006 Aug NEWA
 AB 150MG N76388 003 Aug 30, 2006 Aug NEWA
 AB 200MG N76388 004 Aug 30, 2006 Aug NEWA

TABLET, CHEWABLE; ORAL
LAMICTAL CD

AB GLAXOSMITHKLINE 5MG N20764 001 Aug 24, 1998 Jun CFTG
 AB + 25MG N20764 002 Aug 24, 1998 Jun CFTG
 LAMOTRIGINE
 AB TEVA 5MG N76420 001 Jun 21, 2006 Jun NEWA
 AB 25MG N76420 002 Jun 21, 2006 Jun NEWA

LANSOPRAZOLE; NAPROXEN

CAPSULE, DELAYED REL PELLETS; TABLET; ORAL

>D> PREVACID NAPRAPAC 375 (COPACKAGED)
 >A> @ TAP PHARM 15MG, N/A; N/A, 375MG N21507 003 Nov 14, 2003 Dec DISC

CAPSULE, DELAYED REL PELLETS, TABLET; ORAL

PREVACID NAPRAPAC 250 (COPACKAGED)

@ TAP PHARM 15MG, N/A; N/A, 250MG N21507 002 Nov 14, 2003 Oct DISC
 15MG, N/A; N/A, 250MG N21507 002 Nov 14, 2003 Feb CTNA

PREVACID NAPRAPAC 375 (COPACKAGED)

>D> TAP PHARM 15MG, N/A; N/A, 375MG N21507 003 Nov 14, 2003 Dec DISC
 15MG, N/A; N/A, 375MG N21507 003 Nov 14, 2003 Feb CTNA

PREVACID NAPRAPAC 500 (COPACKAGED)

+ TAP PHARM 15MG, N/A; N/A, 500MG N21507 004 Nov 14, 2003 Feb CTNA

LENALIDOMIDECAPSULE; ORAL
REVLIMID

CELGENE 10MG N21880 002 Dec 27, 2005 Jul CRLD
 15MG N21880 003 Jun 29, 2006 Jul NEWA
 + 25MG N21880 004 Jun 29, 2006 Jul NEWA

LEVETIRACETAMINJECTABLE; IV (INFUSION)
KEPPRA

+ UCB INC 500MG/5ML (100MG/ML) N21872 001 Jul 31, 2006 Jul NEWA

TABLET; ORAL

KEPPRA

UCB INC

750MG

N21035 003 Nov 30, 1999 Mar CRLD

+

1GM

N21035 004 Jan 06, 2006 Mar NEWA

LEVOBETAXOLOL HYDROCHLORIDE

SUSPENSION/DROPS; OPHTHALMIC

BETAXON

@ ALCON

EQ 0.5% BASE

N21114 001 Feb 23, 2000 Feb CAHN

LEVOCABASTINE HYDROCHLORIDE

SUSPENSION/DROPS; OPHTHALMIC

LIVOSTIN

@ NOVARTIS

EQ 0.05% BASE

N20219 001 Nov 10, 1993 May DISC

LEVONORGESTREL

TABLET; ORAL

PLAN B

+ DURAMED

0.75MG

N21045 002 Aug 24, 2006 Aug NEWA

LEVOTHYROXINE SODIUM**

**Refer to Preface Section 1.7 Levothyroxine Sodium for amplifying information

CAPSULE; ORAL

TIROSINT

INST BIOCHIMIQUE

0.025MG

N21924 002 Oct 13, 2006 Oct NEWA

0.05MG

N21924 003 Oct 13, 2006 Oct NEWA

0.075MG

N21924 004 Oct 13, 2006 Oct NEWA

0.1MG

N21924 005 Oct 13, 2006 Oct NEWA

0.125MG

N21924 006 Oct 13, 2006 Oct NEWA

+

0.15MG

N21924 007 Oct 13, 2006 Oct NEWA

TABLET; ORAL

LEVOTHROID

>D>	BX	LLOYD	0.025MG	N21116 001	Oct 24, 2002	Dec	CTEC
>A>	AB4		0.025MG	N21116 001	Oct 24, 2002	Dec	CTEC
>D>	BX		0.05MG	N21116 002	Oct 24, 2002	Dec	CTEC
>A>	AB4		0.05MG	N21116 002	Oct 24, 2002	Dec	CTEC
>D>	BX		0.075MG	N21116 003	Oct 24, 2002	Dec	CTEC
>A>	AB4		0.075MG	N21116 003	Oct 24, 2002	Dec	CTEC
>D>	BX		0.088MG	N21116 010	Oct 24, 2002	Dec	CTEC
>A>	AB4		0.088MG	N21116 010	Oct 24, 2002	Dec	CTEC
>D>	BX		0.1MG	N21116 004	Oct 24, 2002	Dec	CTEC
>A>	AB4		0.1MG	N21116 004	Oct 24, 2002	Dec	CTEC
>D>	BX		0.112MG	N21116 011	Oct 24, 2002	Dec	CTEC
>A>	AB4		0.112MG	N21116 011	Oct 24, 2002	Dec	CTEC
>D>	BX		0.125MG	N21116 005	Oct 24, 2002	Dec	CTEC
>A>	AB4		0.125MG	N21116 005	Oct 24, 2002	Dec	CTEC
>D>	BX		0.137MG	N21116 012	Dec 07, 2004	Dec	CTEC
>A>	AB4		0.137MG	N21116 012	Dec 07, 2004	Dec	CTEC
>D>	BX		0.15MG	N21116 006	Oct 24, 2002	Dec	CTEC
>A>	AB4		0.15MG	N21116 006	Oct 24, 2002	Dec	CTEC
>D>	BX		0.175MG	N21116 007	Oct 24, 2002	Dec	CTEC
>A>	AB4		0.175MG	N21116 007	Oct 24, 2002	Dec	CTEC
>D>	BX		0.2MG	N21116 008	Oct 24, 2002	Dec	CTEC
>A>	AB4		0.2MG	N21116 008	Oct 24, 2002	Dec	CTEC
>D>	BX	+	0.3MG	N21116 009	Oct 24, 2002	Dec	CTEC
>A>	AB4	+	0.3MG	N21116 009	Oct 24, 2002	Dec	CTEC

TABLET; ORAL

LEVOTHYROXINE SODIUM

	AB2, AB3	GENPHARM	0.025MG	N76752 001	Jun 16, 2005	Sep	CTEC
	AB2, AB3		0.05MG	N76752 002	Jun 16, 2005	Sep	CTEC
	AB2, AB3		0.075MG	N76752 003	Jun 16, 2005	Sep	CTEC
	AB2, AB3		0.088MG	N76752 004	Jun 16, 2005	Sep	CTEC
	AB2, AB3		0.1MG	N76752 005	Jun 16, 2005	Sep	CTEC
	AB2, AB3		0.112MG	N76752 006	Jun 16, 2005	Sep	CTEC
	AB2, AB3		0.125MG	N76752 007	Jun 16, 2005	Sep	CTEC
	AB2, AB3		0.15MG	N76752 008	Jun 16, 2005	Sep	CTEC
	AB2, AB3		0.175MG	N76752 009	Jun 16, 2005	Sep	CTEC
	AB2, AB3		0.2MG	N76752 010	Jun 16, 2005	Sep	CTEC
	AB2, AB3		0.3MG	N76752 011	Jun 16, 2005	Sep	CTEC
>D>	AB1, AB2, AB3	MYLAN	0.025MG	N76187 001	Jun 05, 2002	Dec	CTEC
>A>	AB1, AB2, AB3, AB4		0.025MG	N76187 001	Jun 05, 2002	Dec	CTEC
>D>	AB1, AB2, AB3		0.05MG	N76187 002	Jun 05, 2002	Dec	CTEC
>A>	AB1, AB2, AB3, AB4		0.05MG	N76187 002	Jun 05, 2002	Dec	CTEC
>D>	AB1, AB2, AB3		0.075MG	N76187 003	Jun 05, 2002	Dec	CTEC
>A>	AB1, AB2, AB3, AB4		0.075MG	N76187 003	Jun 05, 2002	Dec	CTEC
>D>	AB1, AB2, AB3		0.088MG	N76187 004	Jun 05, 2002	Dec	CTEC
>A>	AB1, AB2, AB3, AB4		0.088MG	N76187 004	Jun 05, 2002	Dec	CTEC
>D>	AB1, AB2, AB3		0.1MG	N76187 005	Jun 05, 2002	Dec	CTEC
>A>	AB1, AB2, AB3, AB4		0.1MG	N76187 005	Jun 05, 2002	Dec	CTEC
>D>	AB1, AB2, AB3		0.112MG	N76187 006	Jun 05, 2002	Dec	CTEC
>A>	AB1, AB2, AB3, AB4		0.112MG	N76187 006	Jun 05, 2002	Dec	CTEC
>D>	AB1, AB2, AB3		0.125MG	N76187 007	Jun 05, 2002	Dec	CTEC

TABLET; ORAL

LEVOTHYROXINE SODIUM

>A>	AB1, AB2, AB3, AB4	MYLAN	0.125MG	N76187 007	Jun 05, 2002	Dec	CTEC
>D>	AB1, AB2, AB3		0.137MG	N76187 012	Dec 13, 2006	Dec	CTEC
>A>	AB1, AB2, AB3, AB4		0.137MG	N76187 012	Dec 13, 2006	Dec	CTEC
	AB1, AB2, AB3		0.137MG	N76187 012	Dec 13, 2006	Nov	NEWA
>D>	AB1, AB2, AB3		0.15MG	N76187 008	Jun 05, 2002	Dec	CTEC
>A>	AB1, AB2, AB3, AB4		0.15MG	N76187 008	Jun 05, 2002	Dec	CTEC
>D>	AB1, AB2, AB3		0.175MG	N76187 009	Jun 05, 2002	Dec	CTEC
>A>	AB1, AB2, AB3, AB4		0.175MG	N76187 009	Jun 05, 2002	Dec	CTEC
>D>	AB1, AB2, AB3		0.2MG	N76187 010	Jun 05, 2002	Dec	CTEC
>A>	AB1, AB2, AB3, AB4		0.2MG	N76187 010	Jun 05, 2002	Dec	CTEC
>D>	AB1, AB2, AB3		0.3MG	N76187 011	Jun 05, 2002	Dec	CTEC
>A>	AB1, AB2, AB3, AB4		0.3MG	N76187 011	Jun 05, 2002	Dec	CTEC
	AB1, AB3	LEVOXYL JONES PHARMA	0.137MG	N21301 008	May 25, 2001	Nov	CTEC
	AB1, AB2	SYNTHROID ABBOTT	0.137MG	N21402 008	Jul 24, 2002	Nov	CTEC

LIDOCAINE HYDROCHLORIDE

INJECTABLE; INJECTION

XYLOCAINE

AP	+	ABRAXIS BIOSCIENCE	0.5%	N06488 008		Jul	CAHN
AP	+		1%	N06488 007		Jul	CAHN
AP	+		1.5%	N06488 010		Jul	CAHN
	@		2%	N06488 002		Jul	CAHN
		XYLOCAINE 4% PRESERVATIVE FREE					
AP	+	ABRAXIS BIOSCIENCE	4%	N10417 001		Jul	CAHN
		XYLOCAINE PRESERVATIVE FREE					
AP		ABRAXIS BIOSCIENCE	1%	N16801 005	Jan 19, 1988	Jul	CAHN
AP	+		2%	N16801 001		Jul	CAHN
AP			4%	N16801 002		Jul	CAHN
AP	+		10%	N16801 003		Jul	CAHN
AP	+		20%	N16801 004		Jul	CAHN

INJECTABLE; SPINAL

XYLOCAINE 1.5% W/ DEXTROSE 7.5%

@ ABRAXIS BIOSCIENCE 1.5%

N16297 001

Jul CAHN

JELLY; TOPICAL

ANESTACON

AT + POLYMEDICA 2%

N80429 001

Jan CDFR

XYLOCAINE

AT + ABRAXIS BIOSCIENCE 2%

N08816 001

Jul CAHN

SOLUTION; ORAL

LIDOCAINE HYDROCHLORIDE VISCOUS

AT ACTAVIS MID ATLANTIC 2%

N86578 001

Jul CAHN

XYLOCAINE VISCOUS

AT + ABRAXIS BIOSCIENCE 2%

N09470 001

Jul CAHN

SOLUTION; TOPICAL

XYLOCAINE 4% PRESERVATIVE FREE

AT + ABRAXIS BIOSCIENCE 4%

N10417 002

Jul CAHN

LIDOCAINE; PRILOCAINE

CREAM; TOPICAL

EMLA

AB + ABRAXIS BIOSCIENCE 2.5%;2.5%

N19941 001 Dec 30, 1992 Jul CAHN

LIDOCAINE; TETRACAINE

CREAM; TOPICAL

LIDOCAINE AND TETRACAINE

+ ZARS 7%;7%

N21717 001 Jun 29, 2006 Jun NEWA

PATCH; TOPICAL

SYNERA

+ ENDO PHARMS 70MG;70MG

N21623 001 Jun 23, 2005 Feb CAHN

LINDANE

LOTION; TOPICAL

LINDANE

AT + ACTAVIS MID ATLANTIC 1%

N87313 001

Jul CAHN

SHAMPOO; TOPICAL

LINDANE

AT + ACTAVIS MID ATLANTIC 1%

N87266 001

Jul CAHN

LIOTHYRONINE SODIUM

INJECTABLE; INJECTION

LIOTHYRONINE SODIUM

AP X GEN PHARMS EQ 0.01MG BASE/ML

N76923 001 Aug 17, 2005 Sep CAHN

LISINOPRIL

TABLET; ORAL

LISINOPRIL

AB ACTAVIS ELIZABETH 2.5MG

N76180 001 Jul 01, 2002 Jun CAHN

AB 5MG

N76180 002 Jul 01, 2002 Jun CAHN

AB 10MG

N76180 003 Jul 01, 2002 Jun CAHN

AB 20MG

N76164 001 Jul 01, 2002 Jun CAHN

AB 30MG

N76164 002 Jul 01, 2002 Jun CAHN

AB 40MG

N76164 003 Jul 01, 2002 Jun CAHN

AB AUROBINDO 2.5MG

N77622 001 Feb 22, 2006 Feb NEWA

AB 5MG

N77622 002 Feb 22, 2006 Feb NEWA

AB 10MG

N77622 003 Feb 22, 2006 Feb NEWA

TABLET; ORAL

LISINOPRIL

AB	AUROBINDO	20MG	N77622 004	Feb 22, 2006	Feb	NEWA
AB		30MG	N77622 005	Feb 22, 2006	Feb	NEWA
AB		40MG	N77622 006	Feb 22, 2006	Feb	NEWA
	PRINIVIL					
	@ MERCK	2.5MG	N19558 006	Jan 28, 1994	Jun	DISC

LITHIUM CARBONATE

TABLET; ORAL

LITHIUM CARBONATE

	@ PFIZER	300MG	N16834 001		Aug	DISC
+	ROXANE	300MG	N18558 001	Jan 29, 1982	Aug	CRLD
	TABLET, EXTENDED RELEASE; ORAL					
	ESKALITH CR					
	@ GLAXOSMITHKLINE	450MG	N18152 001	Mar 29, 1982	Jul	DISC
	LITHIUM CARBONATE					
	@ BARR	450MG	N76366 001	Aug 21, 2003	Jul	DISC
AB	+ ROXANE	450MG	N76691 001	Jan 05, 2004	Jul	CRLD

>D> LORACARBEF

>D> CAPSULE; ORAL

>D> LORABID

>D>	KING PHARMS	200MG	N50668 001	Dec 31, 1991	Dec	DISC
>A>	@	200MG	N50668 001	Dec 31, 1991	Dec	DISC
>D>	+	400MG	N50668 002	Apr 05, 1996	Dec	DISC
>A>	@	400MG	N50668 002	Apr 05, 1996	Dec	DISC
>D>	FOR SUSPENSION; ORAL					
>D>	LORABID					
>D>	KING PHARMS	100MG/5ML	N50667 001	Dec 31, 1991	Dec	DISC
>A>	@	100MG/5ML	N50667 001	Dec 31, 1991	Dec	DISC
>D>	+	200MG/5ML	N50667 002	Dec 31, 1991	Dec	DISC
>A>	@	200MG/5ML	N50667 002	Dec 31, 1991	Dec	DISC

LORAZEPAM

TABLET; ORAL

LORAZEPAM

AB	ACTAVIS ELIZABETH	0.5MG	N71403 001	Apr 21, 1987	Jun	CAHN
AB		1MG	N71404 001	Apr 21, 1987	Jun	CAHN
AB		2MG	N71141 001	Apr 21, 1987	Jun	CAHN
AB	IVAX PHARMS	0.5MG	N77396 001	Dec 13, 2006	Nov	NEWA
AB		1MG	N77396 002	Dec 13, 2006	Nov	NEWA
AB		2MG	N77396 003	Dec 13, 2006	Nov	NEWA
AB	MYLAN	0.5MG	N77657 001	Mar 16, 2006	Mar	NEWA
AB		1MG	N77657 002	Mar 16, 2006	Mar	NEWA
AB		2MG	N77657 003	Mar 16, 2006	Mar	NEWA
AB	VINTAGE PHARMS	0.5MG	N77754 001	May 10, 2006	Apr	NEWA
AB		1MG	N77754 002	May 10, 2006	Apr	NEWA
AB		2MG	N77754 003	May 10, 2006	Apr	NEWA

LOVASTATIN

TABLET; ORAL

LOVASTATIN

AB	ACTAVIS ELIZABETH	10MG	N75828 001	Dec 17, 2001	Jun	CAHN
AB		20MG	N75828 002	Dec 17, 2001	Jun	CAHN
AB		40MG	N75828 003	Dec 17, 2001	Jun	CAHN

TABLET; ORALLOVASTATIN

AB	MUTUAL PHARM	10MG	N77520 001	Apr 14, 2006	Apr	NEWA
AB		20MG	N77520 002	Apr 14, 2006	Apr	NEWA
AB		40MG	N77520 003	Apr 14, 2006	Apr	NEWA
	MEVACOR					
	@ MERCK	10MG	N19643 002	Mar 28, 1991	Aug	DISC

LOVASTATIN; NIACINTABLET, EXTENDED RELEASE; ORALADVICOR

+	KOS LIFE	20MG;750MG	N21249 002	Dec 17, 2001	Feb	CMFD
+		40MG;1GM	N21249 004	Apr 27, 2006	Jul	NEWA

LUBIPROSTONECAPSULE; ORALAMITIZA

+	SUCAMPO PHARMS	24UGM	N21908 001	Jan 31, 2006	Jan	NEWA
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MAGNESIUM CHLORIDE; SODIUM BICARBONATE; SODIUM CHLORIDESOLUTION; INJECTIONNORMOCARB HF 25

+	DIALYSIS SUPS	0.21GM/100ML;2.8GM/100ML;9.07GM/100ML	N21910 001	Jul 26, 2006	Jul	NEWA
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NORMOCARB HF 35

+	DIALYSIS SUPS	0.21GM/100ML;3.97GM/100ML;8.3GM/100ML	N21910 002	Jul 26, 2006	Jul	NEWA
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MAGNESIUM HYDROXIDE; OMEPRAZOLE; SODIUM BICARBONATETABLET, CHEWABLE; ORALZEGERID

	SANTARUS	700MG;20MG;600MG	N21850 001	Mar 24, 2006	Mar	NEWA
+		700MG;40MG;600MG	N21850 002	Mar 24, 2006	Mar	NEWA

MEDROXYPROGESTERONE ACETATEINJECTABLE; SUBCUTANEOUSDEPO-SUBQ PROVERA 104

+	PHARMACIA AND UPJOHN	104MG/0.65ML	N21583 001	Dec 17, 2004	Jan	CAHN
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MEGESTROL ACETATESUSPENSION; ORALMEGESTROL ACETATE

AB	APOTEX	40MG/ML	N77404 001	Feb 16, 2006	Jan	NEWA
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MELOXICAMTABLET; ORALMELOXICAM

AB	ACTAVIS TOTOWA	7.5MG	N77938 001	Jul 19, 2006	Jul	NEWA
AB		15MG	N77938 002	Jul 19, 2006	Jul	NEWA
AB	APOTEX INC	7.5MG	N77882 001	Jul 20, 2006	Jul	NEWA
AB		15MG	N77882 002	Jul 20, 2006	Jul	NEWA
AB	AUROBINDO PHARMA	7.5MG	N78008 001	Oct 02, 2006	Sep	NEWA
AB		15MG	N78008 002	Oct 02, 2006	Sep	NEWA
AB	BRECKENRIDGE PHARM	7.5MG	N77920 001	Jul 19, 2006	Jul	NEWA
AB		15MG	N77920 002	Jul 19, 2006	Jul	NEWA
AB	CARACO	7.5MG	N77937 001	Jul 19, 2006	Jul	NEWA

TABLET; ORAL

MELOXICAM

	AB	CARACO	15MG		N77937 002	Jul 19, 2006	Jul	NEWA
	AB	CARLSBAD	7.5MG		N77918 001	Dec 07, 2006	Nov	NEWA
	AB		15MG		N77918 002	Dec 07, 2006	Nov	NEWA
	AB	COREPHARMA	7.5MG		N77930 001	Jul 19, 2006	Jul	NEWA
	AB		15MG		N77930 002	Jul 19, 2006	Jul	NEWA
	AB	DR REDDYS LABS INC	7.5MG		N77931 001	Jul 25, 2006	Jul	NEWA
	AB		15MG		N77931 002	Jul 25, 2006	Jul	NEWA
	AB	GENPHARM	7.5MG		N77934 001	Jul 20, 2006	Jul	NEWA
	AB		15MG		N77934 002	Jul 20, 2006	Jul	NEWA
	AB	GLENMARK PHARMS LTD	7.5MG		N77932 001	Jul 19, 2006	Jul	NEWA
	AB		15MG		N77932 002	Jul 19, 2006	Jul	NEWA
	AB	LUPIN PHARMS	7.5MG		N77944 001	Jul 19, 2006	Jul	NEWA
	AB		15MG		N77944 002	Jul 19, 2006	Jul	NEWA
	AB	MUTUAL PHARM	7.5MG		N77935 001	Jul 19, 2006	Jul	NEWA
	AB		15MG		N77935 002	Jul 19, 2006	Jul	NEWA
	AB	MYLAN	7.5MG		N77923 001	Jul 19, 2006	Jul	NEWA
	AB		15MG		N77923 002	Jul 19, 2006	Jul	NEWA
	AB	PAR PHARM	7.5MG		N77933 001	Jul 19, 2006	Jul	NEWA
	AB		15MG		N77933 002	Jul 19, 2006	Jul	NEWA
	AB	RANBAXY	7.5MG		N78039 001	Dec 14, 2006	Nov	NEWA
	AB		15MG		N78039 002	Dec 14, 2006	Nov	NEWA
	AB	ROXANE	7.5MG		N77925 001	Jul 19, 2006	Jul	NEWA
	AB		15MG		N77925 002	Jul 19, 2006	Jul	NEWA
	AB	TARO	7.5MG		N78102 001	Nov 07, 2006	Oct	NEWA
	AB		15MG		N78102 002	Nov 07, 2006	Oct	NEWA
	AB	TEVA PHARMS	7.5MG		N77936 001	Jul 19, 2006	Jul	NEWA
	AB		15MG		N77936 002	Jul 19, 2006	Jul	NEWA
>A>	AB	UNICHEM	7.5MG		N77927 001	Dec 20, 2006	Dec	NEWA
>A>	AB		15MG		N77927 002	Dec 20, 2006	Dec	NEWA
	AB	WATSON LABS	7.5MG		N77929 001	Jul 19, 2006	Jul	NEWA
	AB		15MG		N77929 002	Jul 19, 2006	Jul	NEWA
	AB	ZYDUS PHARMS USA	7.5MG		N77921 001	Jul 19, 2006	Jul	NEWA
	AB		15MG		N77921 002	Jul 19, 2006	Jul	NEWA
		MOBIC						
	AB	BOEHRINGER INGELHEIM	7.5MG		N20938 001	Apr 13, 2000	Jul	CFTG
	AB	+	15MG		N20938 002	Aug 23, 2000	Jul	CFTG

MEPROBAMATE

TABLET; ORAL

MEPROBAMATE

	@	ROXANE	600MG	N84332 001	Jan	DISC
	@	SANDOZ	200MG	N14547 002	Jan	DISC
	@		400MG	N14547 001	Jan	DISC
	@		400MG	N80655 001	Jan	DISC
	@	SCHERER LABS	400MG	N83343 001	Jan	DISC
	@	TABLICAPS	400MG	N83494 001	Jan	DISC
AA	+	WATSON LABS	200MG	N83304 001	Jan	CRLD
	@		200MG	N85720 001	Jan	DISC
	+		400MG	N83308 001	Jan	CRLD
	@		400MG	N85721 001	Jan	DISC
	MILTOWN					
	@	MEDPOINTE PHARM HLC	200MG	N09698 004	Jan	DISC
	@		400MG	N09698 002	Jan	DISC

TABLET; ORAL

TRANMEP

@ SOLVAY

400MG

N16249 001

Jan DISC

MESALAMINE

ENEMA; RECTAL

ROWASA

AB + ALAVEN PHARM 4GM/60ML

N19618 001 Dec 24, 1987 Apr CAHN

SUPPOSITORY; RECTAL

CANASA

@ AXCAN SCANDIPHARM

500MG

N21252 001 Jan 05, 2001 May DISC

ROWASA

@ ALAVEN PHARM

500MG

N19919 001 Dec 18, 1990 Jul CAHN

METARAMINOL BITARTRATE

INJECTABLE; INJECTION

ARAMINE

@ MERCK

EQ 10MG BASE/ML

N09509 002 Dec 22, 1987 Jun DISC

METARAMINOL BITARTRATE

+ ABRAXIS PHARM

EQ 10MG BASE/ML

N80722 001 Jun CRLD

METAXALONE

TABLET; ORAL

SKELAXIN

>D> @ JONES PHARMA INC 400MG

N13217 001 Dec CAHN

>D> + 800MG

N13217 003 Aug 30, 2002 Dec CAHN

>A> @ KING PHARMS 400MG

N13217 001 Dec CAHN

>A> + 800MG

N13217 003 Aug 30, 2002 Dec CAHN

METFORMIN HYDROCHLORIDE

TABLET; ORAL

METFORMIN HYDROCHLORIDE

AB ACTAVIS ELIZABETH 500MG

N76033 001 Jan 24, 2002 Jun CAHN

AB 850MG

N76033 002 Jan 24, 2002 Jun CAHN

AB 1GM

N76033 003 Jan 24, 2002 Jun CAHN

AB AMNEAL PHARM 500MG

N77853 001 Jul 28, 2006 Jul NEWA

AB 850MG

N77853 002 Jul 28, 2006 Jul NEWA

AB 1GM

N77853 003 Jul 28, 2006 Jul NEWA

AB DR REDDYS LABS INC 500MG

N77787 001 Aug 23, 2006 Aug NEWA

AB 850MG

N77787 002 Aug 23, 2006 Aug NEWA

AB 1GM

N77787 003 Aug 23, 2006 Aug NEWA

AB INTERPHARM 500MG

N77880 001 Jun 05, 2006 May NEWA

AB 850MG

N77880 002 Jun 05, 2006 May NEWA

AB 1GM

N77880 003 Jun 05, 2006 May NEWA

TABLET, EXTENDED RELEASE; ORAL

GLUMETZA

BX DEPOMED INC 500MG

N21748 001 Jun 03, 2005 Jan CAHN

BX + 1GM

N21748 002 Jun 03, 2005 Aug CRLD

BX 1GM

N21748 002 Jun 03, 2005 Jan CAHN

METFORMIN HYDROCHLORIDE

AB ACTAVIS ELIZABETH 500MG

N76450 001 Oct 01, 2004 Jun CAHN

AB 750MG

N76878 001 Apr 13, 2005 Jun CAHN

AB NOSTRUM 500MG

N76756 001 Jul 26, 2006 Jul NEWA

AB SUN PHARM INDS (IN) 500MG

N77336 001 Feb 09, 2006 Jan NEWA

AB 750MG

N77336 002 Feb 09, 2006 Jan NEWA

METHIMAZOLE

TABLET; ORAL

METHIMAZOLE

	+	CEDAR PHARMS	20MG	N40547 004	Feb 18, 2005	May	CRLD
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TAPAZOLE

>D>	AB	JONES PHARMA	5MG	N40320 001	Mar 31, 2000	Dec	CAHN
	AB		5MG	N40320 001	Mar 31, 2000	May	CTNA
>D>	AB		10MG	N40320 002	Mar 31, 2000	Dec	CAHN
	AB		10MG	N40320 002	Mar 31, 2000	May	CTNA
>A>	AB	KING PHARMS	5MG	N40320 001	Mar 31, 2000	Dec	CAHN
>A>	AB		10MG	N40320 002	Mar 31, 2000	Dec	CAHN

METHOTREXATE SODIUM

INJECTABLE; INJECTION

METHOTREXATE

AP	+	ABRAXIS PHARM	EQ 50MG BASE/2ML (25MG/ML)	N40263 001	Feb 26, 1999	Jun	CMFD
AP	+		EQ 250MG BASE/10ML (25MG/ML)	N40263 002	Feb 26, 1999	Jul	CRLD
AP			EQ 250MG BASE/10ML (25MG/ML)	N40263 002	Feb 26, 1999	Jun	NEWA

METHOTREXATE PRESERVATIVE FREE

	+	BEDFORD	EQ 1GM BASE/VIAL	N40632 001	Aug 12, 2005	Apr	CAIN
	+	MAYNE PHARMA USA	EQ 1GM BASE/40ML (25MG/ML)	N11719 012	Apr 13, 2005	Jun	CPOT
	+		EQ 1GM BASE/40ML (25 MG/ML)	N11719 012	Apr 13, 2005	Apr	CPOT

METHOTREXATE SODIUM

AP	+	BEDFORD	EQ 50MG BASE/2ML (25MG/ML)	N89340 001	Sep 16, 1986	Apr	CPOT
	+		EQ 100MG BASE/4ML (25MG/ML)	N89341 001	Sep 16, 1986	Apr	CTEC
	+		EQ 200MG BASE/8ML (25MG/ML)	N89342 001	Sep 16, 1986	Apr	CTEC
	+		EQ 250MG BASE/10ML (25 MG/ML)	N89343 001	Sep 16, 1986	Apr	CTEC

TABLET; ORAL

METHOTREXATE SODIUM

AB	+	STADA PHARMS	EQ 2.5MG BASE	N08085 002		Aug	CAHN
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METHSCOPOLAMINE BROMIDE

TABLET; ORAL

METHSCOPOLAMINE BROMIDE

>A>	AA	BOCA PHARMA	2.5MG	N40624 001	Dec 28, 2006	Dec	NEWA
>A>	AA		5MG	N40624 002	Dec 28, 2006	Dec	NEWA

PAMINE

>D>	+	BRADLEY PHARMS	2.5MG	N08848 001		Dec	CTEC
>A>	AA	+	2.5MG	N08848 001		Dec	CTEC

PAMINE FORTE

>D>	+	BRADLEY PHARMS	5MG	N08848 002	Mar 25, 2003	Dec	CFTG
>A>	AA	+	5MG	N08848 002	Mar 25, 2003	Dec	CFTG

METHYLPHENIDATE

FILM, EXTENDED RELEASE; TRANSDERMAL

DAYTRANA

	+	SHIRE	10MG/9HR (1.1MG/HR)	N21514 001	Apr 06, 2006	Apr	NEWA
	+		15MG/9HR (1.6MG/HR)	N21514 002	Apr 06, 2006	Apr	NEWA
	+		20MG/9HR (2.2MG/HR)	N21514 003	Apr 06, 2006	Apr	NEWA
	+		30MG/9HR (3.3MG/HR)	N21514 004	Apr 06, 2006	Apr	NEWA

METHYLPHENIDATE HYDROCHLORIDECAPSULE, EXTENDED RELEASE; ORAL
METADATE CD

BX	UCB INC	40MG	N21259 004	Feb 19, 2006	Feb	NEWA
		50MG	N21259 005	Feb 19, 2006	Feb	NEWA
	+	60MG	N21259 006	Feb 19, 2006	Feb	NEWA

RITALIN LA

BX	+ NOVARTIS	40MG	N21284 003	Jun 05, 2002	Feb	CTEC
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TABLET; ORAL

METHYLPHENIDATE HYDROCHLORIDE

AB	ACTAVIS ELIZABETH	5MG	N40321 001	Feb 05, 2002	Jun	CAHN
AB		10MG	N40321 002	Feb 05, 2002	Jun	CAHN
AB		20MG	N40321 003	Feb 05, 2002	Jun	CAHN

TABLET, EXTENDED RELEASE; ORAL

METHYLPHENIDATE HYDROCHLORIDE

AB	ACTAVIS ELIZABETH	20MG	N75450 001	Dec 21, 2001	Jun	CAHN
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METHYLPREDNISOLONE ACETATE

INJECTABLE; INJECTION

METHYLPREDNISOLONE ACETATE

AB	SICOR PHARMS	40MG/ML	N40620 001	Oct 27, 2006	Oct	NEWA
AB		80MG/ML	N40620 002	Oct 27, 2006	Oct	NEWA

METOCLOPRAMIDE HYDROCHLORIDE

SOLUTION; ORAL

METOCLOPRAMIDE

AA	+ JVL	EQ 5MG BASE/5ML	N74703 001	Oct 31, 1997	Aug	CRLD
	@ VISTAPHARM	EQ 5MG BASE/5ML	N75051 001	Jan 26, 2001	Aug	DISC

METOCLOPRAMIDE HYDROCHLORIDE

AA	ACTAVIS MID ATLANTIC	EQ 5MG BASE/5ML	N71340 001	Aug 18, 1988	Jul	CAHN
	@ ROXANE	EQ 5MG BASE/5ML	N72038 001	Dec 05, 1988	Aug	DISC
	@ TEVA	EQ 5MG BASE/5ML	N70819 001	Jul 10, 1987	Aug	DISC
	@	EQ 5MG BASE/5ML	N71315 001	Jun 30, 1993	Aug	DISC

TABLET; ORAL

METOCLOPRAMIDE

AB	VINTAGE PHARMS	EQ 5MG BASE	N77878 001	Aug 28, 2006	Aug	NEWA
AB		EQ 10MG BASE	N77878 002	Aug 28, 2006	Aug	NEWA

METOCLOPRAMIDE HYDROCHLORIDE

AB	ACTAVIS ELIZABETH	EQ 10MG BASE	N70581 001	Oct 17, 1985	Jun	CAHN
AB	MUTUAL PHARM	EQ 5MG BASE	N71536 002	Jan 16, 1997	Apr	CMFD
AB		EQ 10MG BASE	N71536 001	Apr 28, 1993	Apr	CMFD

TABLET, ORALLY DISINTEGRATING; ORAL

REGLAN ODT

@ SCHWARZ PHARMA

@

EQ 5MG BASE

EQ 10MG BASE

N21793 001 Jun 10, 2005 May DISC

N21793 002 Jun 10, 2005 May DISC

METOPROLOL SUCCINATE

TABLET, EXTENDED RELEASE; ORAL

METOPROLOL SUCCINATE

AB	SANDOZ	EQ 25MG TARTRATE	N76969 001	Jul 31, 2006	Jul	NEWA
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TOPROL-XL

AB	ASTRAZENECA	EQ 25MG TARTRATE	N19962 004	Feb 05, 2001	Jul	CFTG
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METOPROLOL TARTRATE

TABLET; ORAL

METOPROLOL TARTRATE

@ APOTHECON

50MG

N74258 001 Jan 27, 1994 Nov DISC

@

100MG

N74258 002 Jan 27, 1994 Nov DISC

METRONIDAZOLE

GEL; TOPICAL

METROGEL

AB + GALDERMA LABS LP

0.75%

N19737 001 Nov 22, 1988 May CFTG

METRONIDAZOLE

AB ALTANA

0.75%

N77018 001 Jun 06, 2006 May NEWA

AB QLT USA

0.75%

N77547 001 Jul 13, 2006 Jun NEWA

AB TARO

0.75%

N77819 001 Jul 18, 2006 Jul NEWA

GEL; VAGINAL

METROGEL-VAGINAL

AB + 3M

0.75%

N20208 001 Aug 17, 1992 Oct CFTG

METRONIDAZOLE

AB QLT USA

0.75%

N77264 001 Oct 31, 2006 Oct NEWA

LOTION; TOPICAL

METROLOTION

AB + GALDERMA LABS LP

0.75%

N20901 001 Nov 24, 1998 May CFTG

METRONIDAZOLE

AB ALTANA

0.75%

N77197 001 May 24, 2006 May NEWA

METYROSINE

CAPSULE; ORAL

DEMSEER

+ ATON

250MG

N17871 001 Oct CAHN

MICAFUNGIN SODIUM

INJECTABLE; IV (INFUSION)

MYCAMINE

ASTELLAS

100MG/VIAL

N21506 003 Jun 27, 2006 Jun NEWA

MICONAZOLE NITRATE

SUPPOSITORY; VAGINAL

MICONAZOLE NITRATE

AB ACTAVIS MID ATLANTIC 200MG

N73508 001 Nov 19, 1993 Jun CAHN

MONISTAT 3

AB + PERSONAL PRODS 200MG

N18888 001 Aug 15, 1984 Aug CAHN

MICONAZOLE NITRATE; PETROLATUM, WHITE; ZINC OXIDE

OINTMENT; TOPICAL

VUSION

+ BARRIER

0.25%;81.35%;15%

N21026 001 Feb 16, 2006 Feb NEWA

MIDAZOLAM HYDROCHLORIDE

INJECTABLE; INJECTION

MIDAZOLAM HYDROCHLORIDE

AP + HOSPIRA EQ 5MG BASE/ML

N75293 002 Jun 20, 2000 Mar CRLD

MIDODRINE HYDROCHLORIDE

TABLET; ORAL

MIDODRINE HYDROCHLORIDE

AB	APOTEX INC	2.5MG	N77746 001	Sep 12, 2006	Aug	NEWA
AB		5MG	N77746 002	Sep 12, 2006	Aug	NEWA
AB		10MG	N77746 003	Sep 12, 2006	Aug	NEWA

MILRINONE LACTATE

INJECTABLE; INJECTION

MILRINONE LACTATE

AP	GLAND PHARMA LTD	EQ 1MG BASE/ML	N77190 001	Oct 31, 2006	Oct	NEWA
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MINOCYCLINE HYDROCHLORIDE

CAPSULE; ORAL

MINOCIN

AB	TRIAx PHARMS	EQ 50MG BASE	N50649 001	May 31, 1990	Feb	CAHN
	@	EQ 75MG BASE	N50649 003	Feb 12, 2001	Feb	CAHN
AB	+	EQ 100MG BASE	N50649 002	May 31, 1990	Feb	CAHN

TABLET, EXTENDED RELEASE; ORAL

SOLODYN

MEDICIS

		EQ 45MG BASE	N50808 001	May 08, 2006	May	NEWA
		EQ 90MG BASE	N50808 002	May 08, 2006	May	NEWA
	+	EQ 135MG BASE	N50808 003	May 08, 2006	May	NEWA

MIRTAZAPINE

TABLET; ORAL

MIRTAZAPINE

AB	ACTAVIS ELIZABETH	15MG	N76308 001	Jun 20, 2003	Jun	CAHN
AB		30MG	N76308 002	Jun 20, 2003	Jun	CAHN
AB		45MG	N76308 003	Jun 20, 2003	Jun	CAHN

TABLET, ORALLY DISINTEGRATING; ORAL

MIRTAZAPINE

AB	AUROBINDO PHARMA LTD	45MG	N77376 004	Feb 28, 2006	Feb	NEWA
AB	BARR	45MG	N76307 003	Feb 28, 2006	Feb	NEWA

MITOXANTRONE HYDROCHLORIDE

INJECTABLE; INJECTION

MITOXANTRONE

AP	AM PHARM	EQ 20MG BASE/10ML (2MG/ML)	N77496 001	Apr 11, 2006	Mar	NEWA
AP		EQ 25MG BASE/12.5ML (2MG/ML)	N77496 002	Apr 11, 2006	Mar	NEWA
AP		EQ 30MG BASE/15ML (2MG/ML)	N77496 003	Apr 11, 2006	Mar	NEWA
AP	BEDFORD	EQ 20MG BASE/10ML (2MG/ML)	N76611 001	Apr 11, 2006	Mar	NEWA
AP		EQ 25MG BASE/12.5ML (2MG/ML)	N76611 002	Apr 11, 2006	Mar	NEWA
AP		EQ 30MG BASE/15ML (2MG/ML)	N76611 003	Apr 11, 2006	Mar	NEWA
AP	MAYNE PHARMA USA	EQ 20MG BASE/10ML (2MG/ML)	N76871 001	Apr 11, 2006	Mar	NEWA
AP		EQ 25MG BASE/12.5ML (2MG/ML)	N76871 002	Apr 11, 2006	Mar	NEWA
AP		EQ 30MG BASE/15ML (2MG/ML)	N76871 003	Apr 11, 2006	Mar	NEWA
AP	SICOR PHARMS	EQ 20MG BASE/10ML (2MG/ML)	N77356 001	Apr 11, 2006	Mar	NEWA
AP		EQ 25MG BASE/12.5ML (2MG/ML)	N77356 002	Apr 11, 2006	Mar	NEWA
AP		EQ 30MG BASE/15ML (2MG/ML)	N77356 003	Apr 11, 2006	Mar	NEWA

NOVANTRONE

AP	+	SERONO INC	EQ 20MG BASE/10ML (2MG/ML)	N19297 001	Dec 23, 1987	Mar	CFTG
AP	+		EQ 25MG BASE/12.5ML (2MG/ML)	N19297 002	Dec 23, 1987	Mar	CFTG
AP	+		EQ 30MG BASE/15ML (2MG/ML)	N19297 003	Dec 23, 1987	Mar	CFTG

MIVACURIUM CHLORIDE

INJECTABLE; INJECTION

MIVACRON

@ ABBOTT

EQ 2MG BASE/ML

N20098 001 Jan 22, 1992 Sep DISC

MIVACRON IN DEXTROSE 5% IN PLASTIC CONTAINER

@ ABBOTT

EQ 50MG BASE/100ML

N20098 003 Jan 22, 1992 Sep DISC

MOEXIPRIL HYDROCHLORIDE

TABLET; ORAL

MOEXIPRIL HYDROCHLORIDE

AB PADDOCK 7.5MG

N77536 001 Nov 30, 2006 Nov NEWA

AB 15MG

N77536 002 Nov 30, 2006 Nov NEWA

MOMETASONE FUROATE

CREAM; TOPICAL

MOMETASONE FUROATE

AB G AND W LABS 0.1%

N77447 001 May 22, 2006 May NEWA

LOTION; TOPICAL

MOMETASONE FUROATE

AB PERRIGO 0.1%

N77180 001 Apr 06, 2005 Mar CAHN

AB TARO 0.1%

N76788 001 Mar 15, 2006 Feb NEWA

OINTMENT; TOPICAL

MOMETASONE FUROATE

AB G AND W LABS 0.1%

N77401 001 Jun 20, 2006 Jun NEWA

MORPHINE SULFATE

CAPSULE, EXTENDED RELEASE; ORAL

KADIAN

ALPHARMA US PHARMS 80MG

N20616 006 Oct 27, 2006 Oct NEWA

INJECTABLE; INJECTION

MORPHINE SULFATE

+ HOSPIRA 5MG/ML

N19916 002 Oct 27, 2006 Oct NEWA

NABILONE

CAPSULE; ORAL

CESAMET

+ VALEANT 1MG

N18677 001 Dec 26, 1985 May CMFD

NABUMETONE

TABLET; ORAL

NABUMETONE

@ COPLEY PHARM

750MG

N75179 001 Jun 06, 2000 Jun DISC

AB PAR PHARM 500MG

N76009 001 Jan 24, 2003 Apr CAHN

AB 750MG

N76009 002 Jan 24, 2003 Apr CAHN

AB + TEVA 750MG

N75189 002 Sep 24, 2001 Jul CRLD

RELAFEN

@ SMITHKLINE BEECHAM

500MG

N19583 001 Dec 24, 1991 Jul DISC

@

750MG

N19583 002 Dec 24, 1991 Jul DISC

NADOLOL

TABLET; ORAL

CORCARD

AB KING PHARMS 20MG

N18063 005 Oct 28, 1986 Jun CAHN

AB 40MG

N18063 001 Jun CAHN

TABLET; ORAL

CORGARD

AB	KING PHARMS	80MG	N18063 002	Jun	CAHN
AB		120MG	N18063 003	Jun	CAHN
AB	+	160MG	N18063 004	Jun	CAHN

NAFCILLIN SODIUM

INJECTABLE; INJECTION

NAFCILLIN SODIUM

AP	+	SANDOZ	EQ 1GM BASE/VIAL	N62527 002	Aug 02, 1984	Apr	CRLD
AP	+		EQ 1GM BASE/VIAL	N62732 001	Dec 23, 1986	Apr	CRLD
AP	+		EQ 2GM BASE/VIAL	N62527 003	Aug 02, 1984	Apr	CRLD
AP	+		EQ 2GM BASE/VIAL	N62732 002	Dec 23, 1986	Apr	CRLD
AP	+		EQ 10GM BASE/VIAL	N62527 004	Aug 02, 1984	Apr	CRLD

NAFTIFINE HYDROCHLORIDE

CREAM; TOPICAL

NAFTIN

+	MERZ PHARMS	1%	N19599 001	Feb 29, 1988	Sep	CRLD
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NALTREXONE

FOR SUSPENSION, EXTENDED RELEASE; INTRAMUSCULAR

VIVITROL

+	ALKERMES	380MG/VIAL	N21897 001	Apr 13, 2006	Apr	NEWA
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NAPROXEN

TABLET, DELAYED RELEASE; ORAL

NAPROXEN

AB	ACTAVIS ELIZABETH	375MG	N74936 001	Feb 24, 1998	Jun	CAHN
AB		500MG	N74936 002	Feb 24, 1998	Jun	CAHN

NEOMYCIN SULFATE

TABLET; ORAL

NEOMYCIN SULFATE

	@	BRISTOL MYERS SQUIBB	500MG	N60365 001		Jul	CPOT
	@	LANNETT	500MG	N60607 001		Jul	CPOT
	@	LILLY	500MG	N60385 001		Jul	CPOT
	@	ROXANE	500MG	N62173 001		Jul	CPOT
	@	SANDOZ	500MG	N61586 001		Jul	CPOT
AB	+	TEVA	500MG	N60304 001		Jul	CFTG
AB		X GEN PHARMS	500MG	N65220 001	Jul 28, 2006	Jul	NEWA

NEOMYCIN SULFATE; POLYMYXIN B SULFATE

SOLUTION; IRRIGATION

NEOMYCIN AND POLYMYXIN B SULFATE

AT	WATSON LABS	EQ 40MG BASE/ML;200,000 UNITS/ML	N62664 001	Apr 08, 1986	Jul	CMFD	
AT	X GEN PHARMS	EQ 40MG BASE/ML;200,000 UNITS/ML	N65106 001	Jan 31, 2006	Jun	CTNA	
AT		EQ 800MG BASE/20ML;4,000,000 UNITS/20ML	N65108 001	Jan 31, 2006	Jun	CTNA	
		NEOSPORIN AND POLYMYXIN B SULFATE					
AT	X GEN PHARMS	EQ 40MG BASE/ML;200,000 UNITS/ML	N65106 001	Jan 31, 2006	Jan	NEWA	
AT		EQ 800MG BASE/20ML;4,000,000 UNITS/20ML (EQ 40MG BASE/ML;200,000 UNITS/ML)	N65108 001	Jan 31, 2006	Jan	NEWA	
		NEOSPORIN G.U. IRRIGANT					
AT	+	MONARCH PHARMS	EQ 40MG BASE/ML;200,000 UNITS/ML	N60707 001		Jan	CTEC

SOLUTION; IRRIGATION

NEOSPORIN G.U. IRRIGANT

AT	+	MONARCH PHARMS	EQ 800MG BASE/20ML;4,000,000 UNITS/20ML (EQ 40MG BASE/ML;200,000 UNITS/ML)	N60707 002		Jan	NEWA
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NIACIN

TABLET, EXTENDED RELEASE; ORAL

NIACIN

@	BARR	500MG	N76378 001	Apr 26, 2005	Sep	DISC
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@		750MG	N76378 002	Apr 26, 2005	Sep	DISC
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@		1GM	N76250 001	Apr 14, 2005	Sep	DISC
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NIASPAN

+	KOS LIFE	500MG	N20381 002	Jul 28, 1997	Sep	CTEC
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+		750MG	N20381 003	Jul 28, 1997	Sep	CTEC
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+		1GM	N20381 004	Jul 28, 1997	Sep	CTEC
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NICARDIPINE HYDROCHLORIDE

CAPSULE; ORAL

CARDENE

>A>	AB	PDL BIOPHARMA INC	20MG	N19488 001	Dec 21, 1988	Dec	CAHN
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>A>	AB	+	30MG	N19488 002	Dec 21, 1988	Dec	CAHN
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>D>	AB	ROCHE PALO	20MG	N19488 001	Dec 21, 1988	Dec	CAHN
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>D>	AB	+	30MG	N19488 002	Dec 21, 1988	Dec	CAHN
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NICARDIPINE HYDROCHLORIDE

AB	BARR	20MG	N74439 001	Dec 10, 1996	Apr	CAHN
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AB		30MG	N74439 002	Dec 10, 1996	Apr	CAHN
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CAPSULE, EXTENDED RELEASE; ORAL

CARDENE SR

>A>	+	PDL BIOPHARMA INC	30MG	N20005 001	Feb 21, 1992	Dec	CAHN
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>A>	+		45MG	N20005 002	Feb 21, 1992	Dec	CAHN
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>A>	+		60MG	N20005 003	Feb 21, 1992	Dec	CAHN
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>D>	+	ROCHE PALO	30MG	N20005 001	Feb 21, 1992	Dec	CAHN
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>D>	+		45MG	N20005 002	Feb 21, 1992	Dec	CAHN
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>D>	+		60MG	N20005 003	Feb 21, 1992	Dec	CAHN
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INJECTABLE; INJECTION

CARDENE

+	PDL BIOPHARMA INC	2.5MG/ML	N19734 001	Jan 30, 1992	Jan	CAHN
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NICOTINE

FILM, EXTENDED RELEASE; TRANSDERMAL

NICOTINE

AVEVA	7MG/24HR	N74645 001	Oct 20, 1997	Oct	CAHN
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	14MG/24HR	N74611 001	Oct 20, 1997	Oct	CAHN
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	21MG/24HR	N74612 001	Oct 20, 1997	Oct	CAHN
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SPRAY, METERED; NASAL

NICOTROL

+	PFIZER INC	0.5MG/SPRAY	N20385 001	Mar 22, 1996	Nov	CAHN
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NIFEDIPINE

CAPSULE; ORAL

NIFEDIPINE

AB	ACTAVIS ELIZABETH	10MG	N72579 001	Jan 08, 1991	Jun	CAHN
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AB		20MG	N72556 001	Sep 20, 1990	Jun	CAHN
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TABLET, EXTENDED RELEASE; ORAL

AFEDITAB CR

AB1	WATSON LABS	30MG	N75128 001	Mar 10, 2000	Jan	CAHN
AB1		60MG	N75659 001	Oct 26, 2001	Jan	CAHN
NIFEDIPINE						
AB1	ABRIKA PHARMS	30MG	N77899 001	Dec 13, 2006	Nov	NEWA
AB1		60MG	N77899 002	Dec 13, 2006	Nov	NEWA

NISOLDIPINE

TABLET, EXTENDED RELEASE; ORAL

SULAR

+	SCIELE PHARMA INC	10MG	N20356 001	Feb 02, 1995	Jun	CAHN
		20MG	N20356 002	Feb 02, 1995	Jun	CAHN
+		30MG	N20356 003	Feb 02, 1995	Jun	CAHN
+		40MG	N20356 004	Feb 02, 1995	Jun	CAHN

NITROGLYCERIN

AEROSOL; SUBLINGUAL

NITROLINGUAL

@	POHL BOSKAMP	0.4MG/SPRAY	N18705 001	Oct 31, 1985	Jun	CAHN
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AEROSOL, METERED; SUBLINGUAL

NITROMIST

+	NOVADEL	0.4MG/SPRAY	N21780 001	Nov 02, 2006	Nov	NEWA
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SPRAY, METERED; SUBLINGUAL

NITROLINGUAL PUMPSPRAY

+	POHL BOSKAMP	0.4MG/SPRAY	N18705 002	Jan 10, 1997	Jun	CAHN
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NOREPINEPHRINE BITARTRATE

INJECTABLE; INJECTION

NOREPINEPHRINE BITARTRATE

AP	METRICS PHARM	EQ 1MG BASE/ML	N40522 001	Sep 30, 2004	May	CAHN
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NORTRIPTYLINE HYDROCHLORIDE

SOLUTION; ORAL

NORTRIPTYLINE HYDROCHLORIDE

AA	TARO	EQ 10MG BASE/5ML	N77965 001	Jun 20, 2006	Jun	NEWA
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NYSTATIN

CREAM; TOPICAL

NYSTATIN

AT	ACTAVIS MID ATLANTIC	100,000 UNITS/GM	N62949 001	Jun 13, 1988	Jun	CAHN
	@ TARO	100,000 UNITS/GM	N62457 001	Jul 28, 1983	May	DISC
AT	VINTAGE	100,000 UNITS/GM	N65315 001	May 31, 2006	May	NEWA

OINTMENT; TOPICAL

NYSTATIN

AT	ACTAVIS MID ATLANTIC	100,000 UNITS/GM	N62840 001	Nov 13, 1987	Jun	CAHN
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POWDER; ORAL

NILSTAT

@	STADA PHARMS	100%	N50576 001	Dec 22, 1983	Sep	CAHN
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NYSTATIN

@	PADDOCK	100%	N62613 001	Nov 26, 1985	Oct	DISC
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POWDER; TOPICAL

NYSTATIN

AT	KV PHARM	100,000 UNITS/GM	N65321 001	Aug 18, 2006	Aug	NEWA
AT	PHARMAFORCE	100,000 UNITS/GM	N65203 001	Jul 15, 2004	Nov	CAHN

SUSPENSION; ORAL

NYSTATIN

AA	ACTAVIS MID ATLANTIC	100,000 UNITS/ML	N62349 001	Jul 14, 1982	Jul	CAHN
AA	TARO	100,000 UNITS/ML	N62876 001	Feb 29, 1988	May	CMFD

NYSTATIN; TRIAMCINOLONE ACETONIDE

CREAM; TOPICAL

MYKACET

AT	ACTAVIS MID ATLANTIC	100,000 UNITS/GM;0.1%	N62367 001	May 28, 1985	Jun	CAHN
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OINTMENT; TOPICAL

MYKACET

AT	ACTAVIS MID ATLANTIC	100,000 UNITS/GM;0.1%	N62733 001	Mar 09, 1987	Jun	CAHN
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OCTREOTIDE ACETATE

INJECTABLE; INJECTION

OCTREOTIDE ACETATE

AP	AM PHARM	EQ 0.2MG BASE/ML	N77450 001	Feb 10, 2006	Jan	NEWA
AP		EQ 1MG BASE/ML	N77450 002	Feb 10, 2006	Jan	NEWA

OCTREOTIDE ACETATE (PRESERVATIVE FREE)

AP	AM PHARM	EQ 0.05MG BASE/ML	N77457 001	Feb 10, 2006	Jan	NEWA
AP		EQ 0.1MG BASE/ML	N77457 002	Feb 10, 2006	Jan	NEWA
AP		EQ 0.5MG BASE/ML	N77457 003	Feb 10, 2006	Jan	NEWA

OFLOXACIN

SOLUTION/DROPS; OTIC

FLOXIN OTIC

+	DAIICHI	0.3%	N20799 001	Dec 16, 1997	Oct	CTEC
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TABLET; ORAL

OFLOXACIN

AB	DR REDDYS LABS LTD	200MG	N77098 001	Feb 10, 2006	Jan	NEWA
AB		300MG	N77098 002	Feb 10, 2006	Jan	NEWA
AB		400MG	N77098 003	Feb 10, 2006	Jan	NEWA

OLOPATADINE HYDROCHLORIDE

SOLUTION/DROPS; OPHTHALMIC

PATADAY

+	ALCON	0.2%	N21545 001	Dec 22, 2004	Oct	CTNA
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OMEPRAZOLE; SODIUM BICARBONATE

CAPSULE; ORAL

ZEGERID

	SANTARUS	20MG;1.1GM	N21849 001	Feb 27, 2006	Feb	NEWA
+		40MG;1.1GM	N21849 002	Feb 27, 2006	Feb	NEWA

FOR SUSPENSION; ORAL

ZEGERID

	SANTARUS	20MG/PACKET;1.68GM/PACKET	N21636 001	Jun 15, 2004	Feb	CAIN
+		40MG/PACKET;1.68GM/PACKET	N21706 001	Dec 21, 2004	Feb	CAIN

ONDANSETRON

TABLET, ORALLY DISINTEGRATING; ORAL

ONDANSETRON

>A>						
>A>	AB	KALI LABS	4MG	N76506 001	Dec 26, 2006	Dec NEWA
>A>	AB		8MG	N76506 002	Dec 26, 2006	Dec NEWA
>A>			16MG	N77406 001	Dec 26, 2006	Dec NEWA
>A>			24MG	N77406 002	Dec 26, 2006	Dec NEWA

TABLET, ORALLY DISINTEGRATING; ORAL
ZOFRAN ODT

>D>		GLAXOSMITHKLINE	4MG	N20781	001	Jan 27, 1999	Dec	CFTG
>A>	AB		4MG	N20781	001	Jan 27, 1999	Dec	CFTG
			4MG	N20781	001	Jan 27, 1999	Nov	CPOT
>D>	+		8MG	N20781	002	Jan 27, 1999	Dec	CFTG
>A>	AB	+	8MG	N20781	002	Jan 27, 1999	Dec	CFTG
		+	8MG	N20781	002	Jan 27, 1999	Nov	CPOT

ONDANSETRON HYDROCHLORIDE

INJECTABLE; INJECTION

		ONDANSETRON							
	AP	TEVA	EQ 2MG BASE/ML	N76876	001	Nov 22, 2006	Nov	NEWA	
>D>		ONDANSETRON AND DEXTROSE IN PLASTIC CONTAINER							
>D>	AP	SICOR PHARMS	EQ 0.64MG BASE/ML	N77480	001	Nov 22, 2006	Dec	CTNA	
	AP		EQ 0.64MG BASE/ML	N77480	001	Nov 22, 2006	Nov	NEWA	
		ONDANSETRON HYDROCHLORIDE							
>A>	AP	ABRAXIS PHARM	EQ 2MG BASE/ML	N76974	001	Dec 26, 2006	Dec	NEWA	
>A>	AP	APOTEX	EQ 2MG BASE/ML	N77368	001	Dec 26, 2006	Dec	NEWA	
>A>	AP	BAXTER HLTHCARE	EQ 2MG BASE/ML	N77365	001	Dec 26, 2006	Dec	NEWA	
>A>	AP	BEDFORD	EQ 2MG BASE/ML	N76967	001	Dec 26, 2006	Dec	NEWA	
>A>	AP	HIKMA FARMACEUTICA	EQ 2MG BASE/ML	N76781	001	Dec 26, 2006	Dec	NEWA	
>A>	AP	HOSPIRA	EQ 2MG BASE/ML	N77473	001	Dec 26, 2006	Dec	NEWA	
>A>	AP	MAYNE PHARMA USA	EQ 2MG BASE/ML	N76695	001	Dec 26, 2006	Dec	NEWA	
>A>	AP	PHARMAFORCE	EQ 2MG BASE/ML	N77582	001	Dec 26, 2006	Dec	NEWA	
>A>	AP	PLIVA HRVATSKA DOO	EQ 2MG BASE/ML	N77544	001	Dec 26, 2006	Dec	NEWA	
>A>	AP	SUN PHARM INDS (IN)	EQ 2MG BASE/ML	N77172	001	Dec 26, 2006	Dec	NEWA	
>A>	AP	TEVA	EQ 2MG BASE/ML	N76876	001	Nov 22, 2006	Dec	NEWA	
>A>	AP	WOCKHARDT	EQ 2MG BASE/ML	N77577	001	Dec 26, 2006	Dec	NEWA	
>A>		ONDANSETRON HYDROCHLORIDE AND DEXTROSE IN PLASTIC CONTAINER							
>A>	AP	SICOR PHARMS	EQ 0.64MG BASE/ML	N77480	001	Nov 22, 2006	Dec	CTNA	
>A>		ONDANSETRON HYDROCHLORIDE AND SODIUM CHLORIDE IN PLASTIC CONTAINER							
>A>	AP	+ BAXTER HLTHCARE	EQ 0.64MG BASE/ML	N21915	002	Dec 27, 2006	Dec	NEWA	
		ONDANSETRON HYDROCHLORIDE PRESERVATIVE FREE							
>A>	AP	ABRAXIS PHARM	EQ 2MG BASE/ML	N76972	001	Dec 26, 2006	Dec	NEWA	
>A>	AP	APOTEX INC	EQ 2MG BASE/ML	N77343	001	Dec 26, 2006	Dec	NEWA	
>A>	AP	BAXTER HLTHCARE	EQ 2MG BASE/ML	N77541	001	Dec 26, 2006	Dec	NEWA	
>A>	AP	BEDFORD LABS	EQ 2MG BASE/ML	N77011	001	Dec 26, 2006	Dec	NEWA	
>A>	AP	HIKMA FARMACEUTICA	EQ 2MG BASE/ML	N76780	001	Dec 26, 2006	Dec	NEWA	
>A>	AP	HOSPIRA	EQ 2MG BASE/ML	N77548	001	Dec 26, 2006	Dec	NEWA	
>A>	AP	MAYNE PHARMA USA	EQ 2MG BASE/ML	N76696	001	Dec 26, 2006	Dec	NEWA	
>A>	AP	PHARMAFORCE	EQ 2MG BASE/ML	N77387	001	Dec 26, 2006	Dec	NEWA	
>A>	AP	SUN PHARM INDS LTD	EQ 2MG BASE/ML	N77173	001	Dec 26, 2006	Dec	NEWA	
>A>	AP	TEVA	EQ 2MG BASE/ML	N76759	001	Nov 22, 2006	Dec	CTNA	
>A>	AP	WOCKHARDT	EQ 2MG BASE/ML	N77716	001	Dec 26, 2006	Dec	NEWA	
>D>		ONDANSETRON PRESERVATIVE FREE							
>D>	AP	TEVA	EQ 2MG BASE/ML	N76759	001	Nov 22, 2006	Dec	CTNA	
	AP		EQ 2MG BASE/ML	N76759	001	Nov 22, 2006	Nov	NEWA	
		ZOFRAN							
	AP	+ GLAXOSMITHKLINE	EQ 2MG BASE/ML	N20007	001	Jan 04, 1991	Nov	CFTG	
		ZOFRAN IN PLASTIC CONTAINER							
	AP	+ GLAXOSMITHKLINE	EQ 0.64MG BASE/ML	N20403	001	Jan 31, 1995	Nov	CFTG	
		ZOFRAN PRESERVATIVE FREE							
	AP	+ GLAXOSMITHKLINE	EQ 2MG BASE/ML	N20007	003	Dec 10, 1993	Nov	CFTG	

SOLUTION; ORAL

>A>		ONDANSETRON HYDROCHLORIDE							
>A>	AA	ROXANE	EQ 4MG BASE/5ML	N76960	001	Dec 26, 2006	Dec	NEWA	
		ZOFRAN							
>D>	+	GLAXOSMITHKLINE	EQ 4MG BASE/5ML	N20605	001	Jan 24, 1997	Dec	CFTG	
>A>	AA	+	EQ 4MG BASE/5ML	N20605	001	Jan 24, 1997	Dec	CFTG	
		<u>TABLET; ORAL</u>							
>A>		ONDANSETRON HYDROCHLORIDE							
>A>	AB	DR REDDYS LABS LTD	EQ 4MG BASE	N76183	003	Dec 26, 2006	Dec	NEWA	
>A>	AB		EQ 8MG BASE	N76183	002	Dec 26, 2006	Dec	NEWA	
>A>			EQ 16MG BASE	N76559	001	Dec 26, 2006	Dec	NEWA	
>A>	AB		EQ 24MG BASE	N76183	001	Dec 26, 2006	Dec	NEWA	
		ZOFRAN							
>D>		GLAXOSMITHKLINE	EQ 4MG BASE	N20103	001	Dec 31, 1992	Dec	CFTG	
>A>	AB		EQ 4MG BASE	N20103	001	Dec 31, 1992	Dec	CFTG	
>D>			EQ 8MG BASE	N20103	002	Dec 31, 1992	Dec	CFTG	
>A>	AB		EQ 8MG BASE	N20103	002	Dec 31, 1992	Dec	CFTG	
>D>	+		EQ 24MG BASE	N20103	003	Aug 27, 1999	Dec	CFTG	
>A>	AB	+	EQ 24MG BASE	N20103	003	Aug 27, 1999	Dec	CFTG	

ORPHENADRINE CITRATEINJECTABLE; INJECTIONORPHENADRINE CITRATE

AP		AKORN	30MG/ML	N40484	001	May 24, 2006	May	NEWA	
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OXALIPLATININJECTABLE; IV (INFUSION)ELOXATIN

@ SANOFI AVENTIS US

50MG/VIAL

N21492 001 Aug 09, 2002 Jun DISC

@

100MG/VIAL

N21492 002 Aug 09, 2002 Jun DISC

+

200MG/40ML (5MG/ML)

N21759 003 Nov 17, 2006 Nov NEWA

OXANDROLONETABLET; ORALOXANDRIN

AB		SAVIENT PHARMS	2.5MG	N13718	001		Nov	CFTG	
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AB	+		10MG	N13718	002	Nov 05, 2001	Nov	CFTG	
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OXANDROLONE

AB		SANDOZ	2.5MG	N76897	001	Dec 01, 2006	Nov	NEWA	
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AB			10MG	N76897	002	Dec 01, 2006	Nov	NEWA	
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AB		UPSHER SMITH	2.5MG	N76761	001	Dec 01, 2006	Nov	NEWA	
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OXAPROZINTABLET; ORALOXAPROZIN

AB		ACTAVIS ELIZABETH	600MG	N75843	001	Oct 03, 2001	Jun	CAHN	
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OXAZEPAMCAPSULE; ORALOXAZEPAM

AB		ACTAVIS ELIZABETH	10MG	N72251	001	Apr 14, 1988	Jun	CAHN	
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AB			15MG	N72252	001	Apr 14, 1988	Jun	CAHN	
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AB			30MG	N72253	001	Apr 14, 1988	Jun	CAHN	
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OXYBUTYNIN CHLORIDE

TABLET, EXTENDED RELEASE; ORAL
DITROPAN XL

AB	ALZA	5MG	N20897 001	Dec 16, 1998	Oct	CFTG
AB		10MG	N20897 002	Dec 16, 1998	Oct	CFTG
AB	+	15MG	N20897 003	Jun 22, 1999	Oct	CFTG
<u>OXYBUTYNIN CHLORIDE</u>						
AB	IMPAX PHARMS	15MG	N76745 001	Nov 09, 2006	Oct	NEWA
AB	MYLAN	5MG	N76702 001	Nov 09, 2006	Oct	NEWA
AB		10MG	N76644 001	Nov 09, 2006	Oct	NEWA

OXYCODONE HYDROCHLORIDE

TABLET, EXTENDED RELEASE; ORAL
OXYCONTIN

PURDUE PHARMA LP	15MG	N20553 006	Sep 18, 2006	Sep	NEWA
	30MG	N20553 007	Sep 18, 2006	Sep	NEWA
	60MG	N20553 008	Sep 18, 2006	Sep	NEWA

OXYMETHOLONE

TABLET; ORAL
ANADROL-50

+	ALAVEN PHARM	50MG	N16848 001		Apr	CAHN
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OXYMORPHONE HYDROCHLORIDE

TABLET; ORAL
OPANA

ENDO PHARMS	5MG	N21611 001	Jun 22, 2006	Jun	NEWA
+	10MG	N21611 002	Jun 22, 2006	Jun	NEWA

TABLET, EXTENDED RELEASE; ORAL
OPANA ER

ENDO PHARMS	5MG	N21610 001	Jun 22, 2006	Jun	NEWA
	10MG	N21610 002	Jun 22, 2006	Jun	NEWA
	20MG	N21610 003	Jun 22, 2006	Jun	NEWA
+	40MG	N21610 004	Jun 22, 2006	Jun	NEWA

PACLITAXEL

FOR SUSPENSION; IV (INFUSION)
ABRAXANE

+	ABRAXIS BIOSCIENCE	100MG/VIAL	N21660 001	Jan 07, 2005	Apr	CAHN
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INJECTABLE; INJECTION
PACLITAXEL

AP	DABUR ONCOLOGY PLC	6MG/ML	N77574 001	Nov 27, 2006	Nov	NEWA
AP	SICOR PHARMS	6MG/ML	N75184 001	Jan 25, 2002	Nov	CAHN
AP		6MG/ML	N75297 001	Jan 25, 2002	Nov	CAHN

>A> PALIPERIDONE

>A> TABLET, EXTENDED RELEASE; ORAL
>A> INVEGA

>A>	JANSSEN LP	3MG	N21999 001	Dec 19, 2006	Dec	NEWA
>A>		6MG	N21999 002	Dec 19, 2006	Dec	NEWA
>A>	+	9MG	N21999 003	Dec 19, 2006	Dec	NEWA
>A>	@	12MG	N21999 004	Dec 19, 2006	Dec	DISC

PARICALCITOL

INJECTABLE; INJECTION

ZEMPLAR

+	ABBOTT	0.002MG/ML	N20819	002	Feb 01, 2000	Aug	CRLD
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PAROXETINE HYDROCHLORIDE

SUSPENSION; ORAL

PAROXETINE HYDROCHLORIDE

AB	APOTEX INC	EQ 10MG BASE/5ML	N77395	001	Dec 05, 2006	Nov	NEWA
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PAXIL

AB	+	GLAXOSMITHKLINE	EQ 10MG BASE/5ML	N20710	001	Jun 25, 1997	Nov	CFTG
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PAROXETINE MESYLATE

TABLET; ORAL

PEXEVA

JDS PHARMS

EQ 10MG BASE

N21299	001	Jul 03, 2003	Apr	CAHN
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EQ 20MG BASE

N21299	002	Jul 03, 2003	Apr	CAHN
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EQ 30MG BASE

N21299	003	Jul 03, 2003	Apr	CAHN
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+

EQ 40MG BASE

N21299	004	Jul 03, 2003	Apr	CAHN
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PEGAPTANIB SODIUM

INJECTABLE; INTRAVITREAL

MACUGEN

+	OSI EYTECH	EQ 0.3MG ACID/0.09ML	N21756	001	Dec 17, 2004	May	CAHN
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PENICILLAMINE

CAPSULE; ORAL

CUPRIMINE

@ ATON

125MG

N19853	002		Aug	CAHN
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+

250MG

N19853	001		Aug	CAHN
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@ MERCK

125MG

N19853	002		Jun	DISC
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PENICILLIN V POTASSIUM

FOR SOLUTION; ORAL

PENICILLIN V POTASSIUM

AA	AM ANTIBIOTICS	EQ 125MG BASE/5ML	N61529	001		Jan	CAHN
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AA		EQ 250MG BASE/5ML	N61529	002		Jan	CAHN
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TABLET; ORAL

PENICILLIN V POTASSIUM

@ AM ANTIBIOTICS

EQ 250MG BASE

N61528	001		Jan	CAHN
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@

EQ 500MG BASE

N61528	002		Jan	CAHN
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VEETIDS

@ APOTHECON

EQ 250MG BASE

N61411	001		Nov	DISC
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@

EQ 500MG BASE

N61411	002		Nov	DISC
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PENTOSTATIN

INJECTABLE; INJECTION

NIPENT

+	MAYNE PHARMA USA	10MG/VIAL	N20122	001	Oct 11, 1991	Aug	CAHN
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PENTOXIFYLLINE

TABLET, EXTENDED RELEASE; ORAL

PENTOXIFYLLINE

AB	ACTAVIS ELIZABETH	400MG	N74878	001	Jul 09, 1997	Jun	CAHN
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TABLET, EXTENDED RELEASE; ORAL

PENTOXIFYLLINE

AB	RADIUS PHARMS	400MG	N74877 001	Jul 08, 1997	Jul	CAHN
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PERGOLIDE MESYLATE

TABLET; ORAL

PERGOLIDE MESYLATE

AB	PAR PHARM	EQ 0.05MG BASE	N76061 001	Nov 27, 2002	Apr	CAHN
AB		EQ 0.25MG BASE	N76061 002	Nov 27, 2002	Apr	CAHN
AB		EQ 1MG BASE	N76061 003	Nov 27, 2002	Apr	CAHN
PERMAX						
AB	VALEANT	EQ 0.05MG BASE	N19385 001	Dec 30, 1988	Jan	CRLD
AB	+	EQ 0.25MG BASE	N19385 002	Dec 30, 1988	Jan	CRLD

PERMETHRIN

CREAM; TOPICAL

PERMETHRIN

AB	ACTAVIS MID ATLANTIC	5%	N74806 001	Jan 23, 1998	Jun	CAHN
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PHENDIMETRAZINE TARTRATE

TABLET; ORAL

CAM-METRAZINE

	@ TG UNITED LABS	35MG	N83922 001		May	CAHN
	@	35MG	N85318 001		May	CAHN
	@	35MG	N85320 001		May	CAHN
	@	35MG	N85321 001		May	CAHN

PHENDIMETRAZINE TARTRATE

	@ TG UNITED LABS	35MG	N85761 001		May	CAHN
	@	35MG	N85941 001	Jun 27, 1983	May	CAHN

PHENTERMINE HYDROCHLORIDE

CAPSULE; ORAL

PHENTERMINE HYDROCHLORIDE

AA	+	SANDOZ	15MG	N87190 002		Nov	CMS1
		@	15MG	N87301 001		Nov	DISC
		@	30MG	N87208 001		Nov	DISC
		@	30MG	N87223 001		Nov	DISC
		@ TG UNITED LABS	18.75MG	N88576 001	May 23, 1984	May	CAHN
		@	30MG	N85417 001		May	CAHN
		@	30MG	N86732 002		May	CAHN
		@	30MG	N87215 001		May	CAHN
		@	37.5MG	N87915 001	Dec 22, 1983	May	CAHN
		@	37.5MG	N87918 001	Dec 22, 1983	May	CAHN
		@	37.5MG	N87930 001	Oct 14, 1983	May	CAHN
		@	37.5MG	N88610 001	Jun 04, 1984	May	CAHN
		@	37.5MG	N88611 001	Jun 04, 1984	May	CAHN
		@	37.5MG	N88625 001	Aug 23, 1984	May	CAHN

TABLET; ORAL

PHENTERMINE HYDROCHLORIDE

AA		ACTAVIS ELIZABETH	37.5MG	N40276 001	Nov 25, 1998	Jun	CAHN
		@ TG UNITED LABS	8MG	N83923 001		May	CAHN
		@	8MG	N85319 001		May	CAHN
		@	37.5MG	N87805 001	Dec 06, 1982	May	CAHN
		@	37.5MG	N88596 001	Apr 04, 1984	May	CAHN

PHENYLEPHRINE HYDROCHLORIDE; PROMETHAZINE HYDROCHLORIDE

SYRUP; ORAL

PHENYLEPHRINE HYDROCHLORIDE AND PROMETHAZINE HYDROCHLORIDE

AA	VINTAGE	5MG/5ML;6.25MG/5ML	N40654	001	Dec 07, 2006	Nov	NEWA
PROMETH VC PLAIN							
AA	+	ACTAVIS MID ATLANTIC	5MG/5ML;6.25MG/5ML	N88761	001	Nov 08, 1984	Nov CFTG
	+		5MG/5ML;6.25MG/5ML	N88761	001	Nov 08, 1984	Jul CAHN
	+	ALPHARMA US PHARMS	5MG/5ML;6.25MG/5ML	N88761	001	Nov 08, 1984	Jan CTEC
PROMETHAZINE VC PLAIN							
	@	MORTON GROVE	5MG/5ML;6.25MG/5ML	N88897	001	Jan 04, 1985	Jan DISC

PHENYTOIN

SUSPENSION; ORAL

PHENYTOIN

AB	ACTAVIS MID ATLANTIC	125MG/5ML	N89892	001	Sep 25, 1992	Jul	CAHN
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PHENYTOIN SODIUM

CAPSULE; ORAL

EXTENDED PHENYTOIN SODIUM

AB	SUN PHARM INDS (IN)	100MG EXTENDED	N40621	001	Dec 11, 2006	Nov	NEWA
AB	TARO	100MG EXTENDED	N40684	001	Sep 05, 2006	Aug	NEWA

INJECTABLE; INJECTION

PHENYTOIN SODIUM

AP	HIKMA FARMACEUTICA	50MG/ML	N40573	001	Sep 13, 2006	Aug	NEWA
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PHYTONADIONE

INJECTABLE; INJECTION

AQUAMEPHYTON

@ ATON

1MG/0.5ML

N12223 002

Oct CAHN

@

10MG/ML

N12223 001

Oct CAHN

@ MERCK

1MG/0.5ML

N12223 002

Feb DISC

@

10MG/ML

N12223 001

Feb DISC

VITAMIN K1

BP	+	HOSPIRA	1MG/0.5ML	N87954	001	Jul 25, 1983	Feb CRLD
	+		10MG/ML	N87955	001	Jul 25, 1983	Feb CRLD

TABLET; ORAL

MEPHYTON

	+	ATON	5MG	N10104	003		Oct CAHN
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PILOCARPINE HYDROCHLORIDE

TABLET; ORAL

PILOCARPINE HYDROCHLORIDE

AB	IMPAX LABS	5MG	N77248	001	Mar 31, 2006	Mar	NEWA
AB		7.5MG	N77248	002	Mar 31, 2006	Mar	NEWA
SALAGEN							
AB	+	MGI PHARMA INC	7.5MG	N20237	002	Apr 18, 2003	Mar CFTG

PIOGLITAZONE HYDROCHLORIDE

TABLET; ORAL

ACTOS

TAKEDA GLOBAL

EQ 15MG BASE

N21073 001 Jul 15, 1999 Oct CAHN

EQ 30MG BASE

N21073 002 Jul 15, 1999 Oct CAHN

+ EQ 45MG BASE

N21073 003 Jul 15, 1999 Oct CAHN

POLYETHYLENE GLYCOL 3350

FOR SOLUTION; ORAL

POLYETHYLENE GLYCOL 3350

AA	COASTAL PHARMS	17GM/SC00PFUL	N77893 001	May 26, 2006	May	NEWA
AA	KALI LABS	17GM/SC00PFUL	N77736 001	May 26, 2006	May	NEWA
AA	TEVA PHARMS	17GM/SC00PFUL	N77445 001	May 04, 2006	Apr	NEWA
AA	YVR THERAP	17GM/SC00PFUL	N77706 001	Sep 27, 2006	Sep	NEWA

POSACONAZOLE

SUSPENSION; ORAL

NOXAFIL

+	SCHERING	40MG/ML	N22003 001	Sep 15, 2006	Sep	NEWA
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POTASSIUM CHLORIDE

INJECTABLE; INJECTION

POTASSIUM CHLORIDE 10MEQ IN PLASTIC CONTAINER

AP	+	BAXTER HLTHCARE	14.9MG/ML	N19904 001	Dec 26, 1989	Jun	CTEC
AP	+		746MG/100ML	N19904 005	Dec 17, 1990	Jun	CTEC

POTASSIUM CHLORIDE 20MEQ IN PLASTIC CONTAINER

AP	+	BAXTER HLTHCARE	1.49GM/100ML	N19904 006	Dec 17, 1990	Jun	CTEC
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TABLET, EXTENDED RELEASE; ORAL

KLOR-CON

AB		UPSHER SMITH	8MEQ	N19123 001	Apr 17, 1986	Jan	CRLD
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POTASSIUM CHLORIDE

AB	+	COPLEY PHARM	8MEQ	N70618 001	Sep 09, 1987	Jan	CRLD
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POTASSIUM CHLORIDE; SODIUM CHLORIDE

INJECTABLE; INJECTION

SODIUM CHLORIDE 0.45% AND POTASSIUM CHLORIDE 0.15% IN PLASTIC CONTAINER

+	BAXTER HLTHCARE	450MG/100ML;150MG/100ML	N17648 005	Nov 26, 2002	Oct	NEWA
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POTASSIUM CITRATE

TABLET, EXTENDED RELEASE; ORAL

POTASSIUM CITRATE

AB		COREPHARMA	5MEQ	N77440 001	Jun 09, 2006	May	NEWA
AB			10MEQ	N77440 002	Jun 09, 2006	May	NEWA

UROCIT-K

AB		MISSION PHARMA	5MEQ	N19071 001	Aug 30, 1985	May	CFTG
AB	+		10MEQ	N19071 002	Aug 31, 1992	May	CFTG

PRAVASTATIN SODIUM

TABLET; ORAL

PRAVACHOL

AB		BRISTOL MYERS SQUIBB	10MG	N19898 002	Oct 31, 1991	Apr	CFTG
AB			20MG	N19898 003	Oct 31, 1991	Apr	CFTG
AB			40MG	N19898 004	Mar 22, 1993	Apr	CFTG

PRAVASTATIN SODIUM

AB		APOTEX	10MG	N76341 001	Oct 23, 2006	Oct	NEWA
AB			20MG	N76341 002	Oct 23, 2006	Oct	NEWA
AB			40MG	N76341 003	Oct 23, 2006	Oct	NEWA
AB		COBALT	10MG	N76939 004	Oct 23, 2006	Oct	NEWA
AB			20MG	N76939 003	Oct 23, 2006	Oct	NEWA
AB			40MG	N76939 002	Oct 23, 2006	Oct	NEWA
AB		DR REDDYS LABS INC	10MG	N76714 001	Oct 23, 2006	Oct	NEWA

TABLET; ORAL

PRAVASTATIN SODIUM

AB	DR REDDYS LABS INC	20MG	N76714 002	Oct 23, 2006	Oct	NEWA
AB		40MG	N76714 003	Oct 23, 2006	Oct	NEWA
AB	GENPHARM	10MG	N77013 001	Oct 23, 2006	Oct	NEWA
AB		20MG	N77013 002	Oct 23, 2006	Oct	NEWA
AB		40MG	N77013 003	Oct 23, 2006	Oct	NEWA
AB	LEK PHARMS DD	10MG	N76397 003	Oct 23, 2006	Oct	NEWA
AB		20MG	N76397 002	Oct 23, 2006	Oct	NEWA
AB		40MG	N76397 001	Oct 23, 2006	Oct	NEWA
AB	PLIVA HRVATSKA DOO	10MG	N77730 001	Nov 21, 2006	Nov	NEWA
AB		20MG	N77730 002	Nov 21, 2006	Nov	NEWA
		30MG	N77730 003	Nov 21, 2006	Nov	NEWA
AB		40MG	N77730 005	Nov 21, 2006	Nov	NEWA
AB	TEVA	10MG	N76056 001	Apr 24, 2006	Apr	NEWA
AB		20MG	N76056 002	Apr 24, 2006	Apr	NEWA
AB		40MG	N76056 003	Apr 24, 2006	Apr	NEWA
AB	WATSON LABS	10MG	N77904 001	Oct 23, 2006	Oct	NEWA
AB		20MG	N77904 002	Oct 23, 2006	Oct	NEWA
AB		40MG	N77904 003	Oct 23, 2006	Oct	NEWA

PRAZOSIN HYDROCHLORIDE

CAPSULE; ORAL

PRAZOSIN HYDROCHLORIDE

@ CLONMEL HLTHCARE

EQ 1MG BASE

N72705 001 May 16, 1989 Jan DISC

@

EQ 5MG BASE

N72707 001 May 16, 1989 Jan DISC

PREDNICARBATE

CREAM; TOPICAL

DERMATOP E EMOLLIENT

AB	+	SANOFI AVENTIS US	0.1%	N20279 001	Oct 29, 1993	Sep	CFTG
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PREDNICARBATE

AB		ALTANA	0.1%	N77287 001	Sep 19, 2006	Sep	NEWA
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PREDNISOLONE ACETATE

INJECTABLE; INJECTION

PREDNISOLONE ACETATE

@ STERIS

25MG/ML

N83398 001 Mar DISC

@

50MG/ML

N83764 001 Mar DISC

SUSPENSION/DROPS; OPHTHALMIC

OMNIPRED

AB		ALCON	1%	N17469 001		May	CTNA
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PREDNISOLONE SODIUM PHOSPHATE

TABLET, ORALLY DISINTEGRATING; ORAL

ORAPRED ODT

BIOMARIN PHARM

EQ 10MG BASE

N21959 001 Jun 01, 2006 Jun NEWA

EQ 15MG BASE

N21959 002 Jun 01, 2006 Jun NEWA

+

EQ 30MG BASE

N21959 003 Jun 01, 2006 Jun NEWA

MEDICIS

EQ 10MG BASE

N21959 001 Jun 01, 2006 Oct CAHN

EQ 15MG BASE

N21959 002 Jun 01, 2006 Oct CAHN

+

EQ 30MG BASE

N21959 003 Jun 01, 2006 Oct CAHN

PRIMIDONE

TABLET; ORAL

PRIMIDONE

AB	WEST WARD	50MG	N40667 001	Jul 27, 2006	Jul	NEWA
AB		250MG	N40667 002	Jul 27, 2006	Jul	NEWA

PROCHLORPERAZINE

SUPPOSITORY; RECTAL

COMPAZINE

@ GLAXOSMITHKLINE

2.5MG

N11127 003

May DISC

@

5MG

N11127 001

May DISC

PROCHLORPERAZINE EDISYLATE

SYRUP; ORAL

COMPAZINE

@ GLAXOSMITHKLINE

EQ 5MG BASE/5ML

N11188 001

May DISC

PROCHLORPERAZINE MALEATE

CAPSULE, EXTENDED RELEASE; ORAL

COMPAZINE

@ GLAXOSMITHKLINE

EQ 10MG BASE

N21019 001 Oct 06, 1999 May DISC

PROGESTERONE

GEL; VAGINAL

CRINONE

COLUMBIA LABS

4%

N20701 001 Jul 31, 1997 May CAHN

+

8%

N20701 002 Jul 31, 1997 May CAHN

PROMETHAZINE HYDROCHLORIDE

INJECTABLE; INJECTION

PROMETHAZINE HYDROCHLORIDE

AP SANDOZ 25MG/ML

N40593 001 Nov 08, 2006 Oct NEWA

AP 50MG/ML

N40593 002 Nov 08, 2006 Oct NEWA

SUPPOSITORY; RECTAL

PHENERGAN

@ WYETH PHARMS INC

12.5MG

N10926 002

Mar DISC

@

25MG

N10926 001

Mar DISC

PROMETHAZINE HYDROCHLORIDE

AB + G AND W LABS 25MG

N40428 001 Feb 05, 2002 Mar CRLD

AB TARO 12.5MG

N40603 001 Oct 26, 2006 Oct NEWA

AB 25MG

N40603 002 Oct 26, 2006 Oct NEWA

PROMETHEGAN

+ G AND W LABS

50MG

N87165 001 Aug 14, 1987 Jan CRLD

SYRUP; ORAL

PROMETH PLAIN

@ ACTAVIS MID ATLANTIC 6.25MG/5ML

N85953 001

Jul CAHN

PROMETHAZINE HYDROCHLORIDE

AA VINTAGE 6.25MG/5ML

N40643 001 Apr 26, 2006 Apr NEWA

TABLET; ORAL

PROMETHAZINE HYDROCHLORIDE

AB KVK-TECH INC 25MG

N40712 001 Jul 31, 2006 Jul NEWA

AB 50MG

N40713 001 Jul 31, 2006 Jul NEWA

AB VINTAGE PHARMS 12.5MG

N40622 001 Jul 18, 2006 Jul NEWA

AB 25MG

N40622 002 Jul 18, 2006 Jul NEWA

TABLET; ORAL

PROMETHAZINE HYDROCHLORIDE

AB	VINTAGE PHARMS	50MG	N40622 003	Jul 18, 2006	Jul	NEWA
AB		50MG	N40622 003	Jul 18, 2006	Jun	NEWA
AB	ZYDUS PHARMS USA	12.5MG	N40596 001	Nov 18, 2005	Jul	CTEC

PROPOFOL

INJECTABLE; INJECTION

DIPRIVAN

AB	+ ABRAXIS BIOSCIENCE	10MG/ML	N19627 002	Jun 11, 1996	Jul	CAHN
	@	10MG/ML	N19627 001	Oct 02, 1989	Jul	CAHN
AB	PROPOFOL					
AB	HOSPIRA	10MG/ML	N77908 001	Mar 17, 2006	Mar	NEWA

PROPOXYPHENE HYDROCHLORIDE

CAPSULE; ORAL

DOLENE

	@ RADIUS PHARMS	65MG	N80530 001		Aug	CAHN
AA	PAR PHARM	65MG	N80269 001		Mar	CAHN

PROPRANOLOL HYDROCHLORIDE

INJECTABLE; INJECTION

PROPRANOLOL HYDROCHLORIDE

AP	SANDOZ	1MG/ML	N76400 001	Feb 26, 2003	Jan	CAHN
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PROPYLTHIOURACIL

TABLET; ORAL

PROPYLTHIOURACIL

BD	ACTAVIS ELIZABETH	50MG	N80172 001		Jun	CAHN
	@ IMPAX LABS	50MG	N80159 001		Nov	DISC
BD	+ STADA PHARMS	50MG	N06188 001		Sep	CAHN

PYRIDOSTIGMINE BROMIDE

INJECTABLE; INJECTION

REGONOL

AP	SANDOZ	5MG/ML	N17398 001		Jan	CAHN
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QUAZEPAM

TABLET; ORAL

DORAL

	@ QUESTCOR PHARMS	7.5MG	N18708 003	Feb 26, 1987	May	CAHN
	+	15MG	N18708 001	Dec 27, 1985	May	CAHN

QUINAPRIL HYDROCHLORIDE

TABLET; ORAL

QUINAPRIL

AB	LUPIN	EQ 5MG BASE	N77690 001	Jun 20, 2006	Jun	NEWA
AB		EQ 10MG BASE	N77690 002	Jun 20, 2006	Jun	NEWA
AB		EQ 20MG BASE	N77690 003	Jun 20, 2006	Jun	NEWA
AB		EQ 40MG BASE	N77690 004	Jun 20, 2006	Jun	NEWA

QUINAPRIL HYDROCHLORIDE

AB	TORPHARM	EQ 5MG BASE	N76240 001	Jan 26, 2006	Jan	NEWA
AB		EQ 10MG BASE	N76240 002	Jan 26, 2006	Jan	NEWA
AB		EQ 20MG BASE	N76240 003	Jan 26, 2006	Jan	NEWA

TABLET; ORAL

QUINAPRIL HYDROCHLORIDE

AB	TORPHARM	EQ 40MG BASE	N76240 004	Jan 26, 2006	Jan	NEWA
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QUINIDINE GLUCONATE

TABLET, EXTENDED RELEASE; ORAL

QUINIDINE GLUCONATE

BX	+	MUTUAL PHARM	324MG	N89338 001	Feb 11, 1987	Jan	CTEC
BX		WATSON LABS	324MG	N87810 001	Sep 29, 1982	Jan	CMFD

QUINIDINE SULFATE

TABLET; ORAL

QUINIDINE SULFATE

	@	CLONMEL HLTHCARE	200MG	N87011 001		Jan	DISC
	@	LANNETT	200MG	N83743 001		Jan	DISC
	@	MUTUAL PHARM	100MG	N81029 001	Apr 14, 1989	Jan	DISC
AB			300MG	N81031 001	Apr 14, 1989	Jul	CMFD
	@	PHARM FORM	200MG	N83808 001		Jan	DISC
	@	SANDOZ	200MG	N84631 001		Jan	DISC
	@		200MG	N84914 001		Jan	DISC
AB			200MG	N88072 002		Jan	NEWA
	@		300MG	N89839 001	Sep 29, 1988	Jan	DISC
AB		WATSON LABS	200MG	N83288 001		Jul	CMFD
	@		200MG	N83288 001		Jan	DISC
	@		200MG	N85140 002		Jan	DISC

QUININE SULFATE

CAPSULE; ORAL

QUININE SULFATE

+	AR HOLDING CO INC	324MG	N21799 001	Aug 12, 2005	Sep	CAHN
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RANITIDINE

INJECTABLE; INJECTION

RANITIDINE

AP	BEDFORD	EQ 25MG BASE/ML	N77458 001	Feb 16, 2006	Feb	NEWA
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RANITIDINE HYDROCHLORIDE

TABLET; ORAL

RANITIDINE HYDROCHLORIDE

AB	INTERPHARM	EQ 150MG BASE	N77824 001	Oct 13, 2006	Oct	NEWA
AB		EQ 300MG BASE	N77824 002	Oct 13, 2006	Oct	NEWA

RANOLAZINE

TABLET, EXTENDED RELEASE; ORAL

RANEXA

+	CV THERAP	500MG	N21526 002	Jan 27, 2006	Jan	NEWA
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RASAGILINE MESYLATE

TABLET; ORAL

AZILECT

TEVA

+		EQ 0.5MG BASE	N21641 001	May 16, 2006	May	NEWA
		EQ 1MG BASE	N21641 002	May 16, 2006	May	NEWA

RIBAVIRIN

TABLET; ORAL							
COPEGUS							
AB	ROCHE	200MG	N21511	001	Dec 03, 2002	Nov	CRLD
RIBAVIRIN							
AB	SANDOZ	200MG	N77743	001	Oct 03, 2006	Sep	NEWA

RISPERIDONE

TABLET, ORALLY DISINTEGRATING; ORAL
RISPERDAL

JANSSEN PHARMA	3MG	N21444	004	Dec 23, 2004	Mar	CMFD
	4MG	N21444	005	Dec 23, 2004	Mar	CMFD

ROPIVACAINE HYDROCHLORIDE MONOHYDRATE

INJECTABLE; INJECTION
NAROPIN

ABRAXIS BIOSCIENCE	2MG/ML	N20533	001	Sep 24, 1996	Jul	CAHN
	5MG/ML	N20533	003	Sep 24, 1996	Jul	CAHN
	7.5MG/ML	N20533	004	Sep 24, 1996	Jul	CAHN
+	10MG/ML	N20533	005	Sep 24, 1996	Jul	CAHN

SALMETEROL XINAFOATE

AEROSOL, METERED; INHALATION
SEREVENT

@ GLAXOSMITHKLINE	EQ 0.021MG BASE/INH	N20236	001	Feb 04, 1994	Oct	DISC
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SAQUINAVIR

CAPSULE; ORAL
FORTOVASE

@ HLR	200MG	N20828	001	Nov 07, 1997	Nov	DISC
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SECRETIN SYNTHETIC PORCINE

FOR SOLUTION; INTRAVENOUS
SECREFLO

+	CHIRHOCLIN	16UGM/VIAL	N21136	001	Apr 04, 2002	May	CTNA
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SELEGILINE

FILM, EXTENDED RELEASE; TRANSDERMAL
EMSAM

SOMERSET	6MG/24HR	N21336	001	Feb 27, 2006	Mar	CAIN
	9MG/24HR	N21336	002	Feb 27, 2006	Mar	CAIN
+	12MG/24HR	N21336	003	Feb 27, 2006	Mar	CAIN

SELEGILINE HYDROCHLORIDE

CAPSULE; ORAL
SELEGILINE HYDROCHLORIDE

@ AAIPHARMA LLC	5MG	N75145	001	Sep 15, 2003	Sep	DISC
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FILM, EXTENDED RELEASE; TRANSDERMAL
EMSAM

SOMERSET	6MG/24HR	N21336	001	Feb 27, 2006	Feb	NEWA
	9MG/24HR	N21336	002	Feb 27, 2006	Feb	NEWA
+	12MG/24HR	N21336	003	Feb 27, 2006	Feb	NEWA

TABLET, ORALLY DISINTEGRATING; ORAL
ZELAPAR

+ VALEANT PHARM INTL 1.25MG N21479 001 Jun 14, 2006 Jun NEWA

SELENIUM SULFIDE

LOTION/SHAMPOO; TOPICAL
SELENIUM SULFIDE

AT ACTAVIS MID ATLANTIC 2.5% N84394 001 Jul CAHN

SERTRALINE HYDROCHLORIDE

CONCENTRATE; ORAL

SERTRALINE HYDROCHLORIDE

AB ROXANE EQ 20MG BASE/ML N76934 001 Jun 30, 2006 Jun NEWA

ZOLOFT

AB + PFIZER EQ 20MG BASE/ML N20990 001 Dec 07, 1999 Jun CFTG

TABLET; ORAL

SERTRALINE HYDROCHLORIDE

AB IVAX PHARMS EQ 25MG BASE N75719 003 Jun 30, 2006 Jun NEWA

AB EQ 50MG BASE N75719 001 Jun 30, 2006 Jun NEWA

AB EQ 100MG BASE N75719 002 Jun 30, 2006 Jun NEWA

AB TEVA EQ 25MG BASE N76465 001 Aug 11, 2006 Aug NEWA

AB EQ 50MG BASE N76465 002 Aug 11, 2006 Aug NEWA

AB EQ 100MG BASE N76465 003 Aug 11, 2006 Aug NEWA

ZOLOFT

AB PFIZER EQ 25MG BASE N19839 005 Mar 06, 1996 Jun CFTG

AB EQ 50MG BASE N19839 001 Dec 30, 1991 Jun CFTG

AB + EQ 100MG BASE N19839 002 Dec 30, 1991 Jun CFTG

SIMVASTATIN

TABLET; ORAL

SIMVASTATIN

>A> AB AUROBINDO PHARMA 5MG N77691 001 Dec 20, 2006 Dec NEWA

>A> AB 10MG N77691 002 Dec 20, 2006 Dec NEWA

>A> AB 20MG N77691 003 Dec 20, 2006 Dec NEWA

>A> AB 40MG N77691 004 Dec 20, 2006 Dec NEWA

>A> AB 80MG N77691 005 Dec 20, 2006 Dec NEWA

>A> AB COBALT 5MG N76685 001 Dec 20, 2006 Dec NEWA

>A> AB 10MG N76685 002 Dec 20, 2006 Dec NEWA

>A> AB 20MG N76685 003 Dec 20, 2006 Dec NEWA

>A> AB 40MG N76685 004 Dec 20, 2006 Dec NEWA

>A> AB 80MG N76685 005 Dec 20, 2006 Dec NEWA

>A> AB DR REDDYS LABS INC 10MG N77752 001 Dec 20, 2006 Dec NEWA

>A> AB 20MG N77752 002 Dec 20, 2006 Dec NEWA

>A> AB 40MG N77752 003 Dec 20, 2006 Dec NEWA

>A> AB 80MG N77752 004 Dec 20, 2006 Dec NEWA

AB IVAX PHARMS 5MG N76052 001 Jun 23, 2006 Jun NEWA

AB 10MG N76052 002 Jun 23, 2006 Jun NEWA

AB 20MG N76052 003 Jun 23, 2006 Jun NEWA

AB 40MG N76052 004 Jun 23, 2006 Jun NEWA

>A> AB 80MG N76052 005 Dec 20, 2006 Dec NEWA

>A> AB PERRIGO R AND D 5MG N78034 001 Dec 20, 2006 Dec NEWA

>A> AB 10MG N78034 002 Dec 20, 2006 Dec NEWA

>A> AB 20MG N78034 003 Dec 20, 2006 Dec NEWA

>A> AB 40MG N78034 004 Dec 20, 2006 Dec NEWA

>A> AB 80MG N78034 005 Dec 20, 2006 Dec NEWA

>A> AB RANBAXY 5MG N76285 001 Dec 20, 2006 Dec NEWA

TABLET; ORAL
SIMVASTATIN

>A>	AB	RANBAXY	10MG	N76285 002	Dec 20, 2006	Dec	NEWA
>A>	AB		20MG	N76285 003	Dec 20, 2006	Dec	NEWA
>A>	AB		40MG	N76285 004	Dec 20, 2006	Dec	NEWA
	AB		80MG	N76285 005	Jun 23, 2006	Jun	NEWA
>A>	AB	SANDOZ	5MG	N77766 001	Dec 20, 2006	Dec	NEWA
>A>	AB		10MG	N77766 002	Dec 20, 2006	Dec	NEWA
>A>	AB		20MG	N77766 003	Dec 20, 2006	Dec	NEWA
>A>	AB		40MG	N77766 004	Dec 20, 2006	Dec	NEWA
>A>	AB		80MG	N77766 005	Dec 20, 2006	Dec	NEWA
>A>	AB	ZYDUS PHARMS USA	5MG	N77837 001	Dec 20, 2006	Dec	NEWA
>A>	AB		10MG	N77837 002	Dec 20, 2006	Dec	NEWA
>A>	AB		20MG	N77837 003	Dec 20, 2006	Dec	NEWA
>A>	AB		40MG	N77837 004	Dec 20, 2006	Dec	NEWA
>A>	AB		80MG	N77837 005	Dec 20, 2006	Dec	NEWA
		ZOCOR					
	AB	MERCK	5MG	N19766 001	Dec 23, 1991	Jun	CFTG
	AB		10MG	N19766 002	Dec 23, 1991	Jun	CFTG
	AB		20MG	N19766 003	Dec 23, 1991	Jun	CFTG
	AB		40MG	N19766 004	Dec 23, 1991	Jun	CFTG
	AB	+	80MG	N19766 005	Jul 10, 1998	Jun	CFTG

SITAGLIPTIN PHOSPHATE

TABLET; ORAL
JANUVIA

		MERCK CO INC	EQ 25MG BASE	N21995 001	Oct 16, 2006	Oct	NEWA
			EQ 50MG BASE	N21995 002	Oct 16, 2006	Oct	NEWA
		+	EQ 100MG BASE	N21995 003	Oct 16, 2006	Oct	NEWA

SODIUM CHLORIDE

INJECTABLE; INJECTION

SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER

AP	+	ABRAXIS PHARM	9MG/ML	N88912 001	Jan 10, 1985	Oct	CRLD
AP	+	B BRAUN	900MG/100ML	N17464 001		May	CRLD
AP	+		900MG/100ML	N19635 002	Mar 09, 1988	May	CRLD
AP	+	BAXTER HLTHCARE	900MG/100ML	N16677 001		May	CRLD
AP	+		900MG/100ML	N20178 001	Dec 07, 1992	May	CRLD
AP	+	HOSPIRA	9MG/ML	N18803 001	Oct 29, 1982	Oct	CRLD
AP	+		900MG/100ML	N16366 001		May	CRLD
AP	+		900MG/100ML	N19465 001	Jul 15, 1985	May	CRLD
AP	+		900MG/100ML	N19480 001	Sep 17, 1985	May	CRLD
	+	MALLINCKRODT	45MG/50ML (9MG/ML)	N21569 001	Jul 27, 2006	Aug	CDFR
			112.5MG/125ML (9MG/ML)	N21569 002	Jul 27, 2006	Aug	CDFR
AP	+	TARO PHARMS IRELAND	9MG/ML	N77407 001	Aug 11, 2006	Oct	CRLD
AP			9MG/ML	N77407 001	Aug 11, 2006	Aug	NEWA

INJECTABLE; INTRAVASCULAR

SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER

	+	MALLINCKRODT	45MG/50ML (0.9MG/ML)	N21569 001	Jul 27, 2006	Jul	NEWA
			112.5MG/125ML (0.9MG/ML)	N21569 002	Jul 27, 2006	Jul	NEWA

SODIUM IODIDE, I-123

CAPSULE; ORAL

SODIUM IODIDE I 123

AA	+	GE HEALTHCARE	100uCi	N17630 001		Sep	CRLD
AA	+	SYNCOR PHARMS	100uCi	N18671 001	May 27, 1982	Sep	CRLD

CAPSULE; ORAL

SODIUM IODIDE I 123

AA + SYNCOR PHARMS 200uCi N18671 002 May 27, 1982 Sep CRLD

SOLUTION; ORAL

SODIUM IODIDE I 123

+ GE HEALTHCARE 2mCi/ML N17630 002 Sep CRLD

SODIUM IODIDE, I-131

CAPSULE; ORAL

SODIUM IODIDE I 131

>D> DRAXIMAGE 100mCi N21305 004 Nov 18, 2004 Dec CPOT

>A> 100uCi N21305 004 Nov 18, 2004 Dec CPOT

100mCi N21305 004 Nov 18, 2004 Jun NEWA

SODIUM PHOSPHATE, DIBASIC, ANHYDROUS; SODIUM PHOSPHATE, MONOBASIC, MONOHYDRATE

TABLET; ORAL

OSMOPREP

+ SALIX PHARMS 0.398GM;1.102GM N21892 001 Mar 16, 2006 Mar NEWA

SOMATROPIN RECOMBINANT

INJECTABLE; INJECTION

GENOTROPIN

BX + PHARMACIA AND UPJOHN 5.8MG/VIAL N20280 006 Aug 24, 1995 May CTEC

GENOTROPIN PRESERVATIVE FREE

BX PHARMACIA AND UPJOHN 1.5MG/VIAL N20280 004 Aug 24, 1995 May CTEC

NUTROPIN DEPOT

@ GENENTECH 13.5MG/VIAL N21075 001 Dec 22, 1999 Jul DISC

@ 18MG/VIAL N21075 002 Dec 22, 1999 Jul DISC

@ 22.5MG/VIAL N21075 003 Dec 22, 1999 Jul DISC

OMNITROPE

BX SANDOZ 1.5MG/VIAL N21426 002 May 30, 2006 May NEWA

BX 5.8MG/VIAL N21426 001 May 30, 2006 May NEWA

SOYBEAN OIL

INJECTABLE; INJECTION

SOYACAL 10%

@ ALPHA THERA 10% N18465 001 Jun 29, 1983 Nov DISC

SOYACAL 20%

@ ALPHA THERA 20% N18786 001 Jun 29, 1983 Nov DISC

SPIRONOLACTONE

TABLET; ORAL

SPIRONOLACTONE

AB ACTAVIS ELIZABETH 25MG N40353 003 Mar 15, 2006 Jun CAHN

AB 50MG N40353 001 Jul 29, 1999 Jun CAHN

AB 100MG N40353 002 Jul 29, 1999 Jun CAHN

AB PUREPAC PHARM 25MG N40353 003 Mar 15, 2006 Feb NEWA

AB VINTAGE 25MG N40750 001 Aug 29, 2006 Aug NEWA

AB 50MG N40750 002 Aug 29, 2006 Aug NEWA

AB 100MG N40750 003 Aug 29, 2006 Aug NEWA

SUCCINYLCHOLINE CHLORIDE

INJECTABLE; INJECTION

ANECTINE

AP + SANDOZ 20MG/ML N08453 002 Jan CAHN

INJECTABLE; INJECTION

ANECTINE

@ SANDOZ 50MG/ML
 @ 500MG/VIAL
 @ 1GM/VIAL

N08453 003 Jan CAHN
 N08453 001 Jan CAHN
 N08453 004 Jan CAHN

SUCCINYLCHOLINE CHLORIDE

@ ORGANON USA INC 20MG/ML

N80997 001 Jun DISC

SULCONAZOLE NITRATESOLUTION; TOPICAL
EXELDERM

+ WESTWOOD SQUIBB 1%

N18738 001 Aug 30, 1985 Jun CMFD

SULFACETAMIDE SODIUMLOTION; TOPICAL
KLARON

AB + SANOFI AVENTIS US 10%

N19931 001 Dec 23, 1996 Nov CFTG

SULFACETAMIDE SODIUM

AB ALTANA 10%

N77015 001 Nov 17, 2006 Nov NEWA

SOLUTION/DROPS; OPHTHALMIC

SULFACETAMIDE SODIUM

AT ALCON 30%

N89068 001 May 05, 1987 Apr CAHN

SULFAMETHOXAZOLE; TRIMETHOPRIM

SUSPENSION; ORAL

BACTRIM PEDIATRIC

AB MUTUAL PHARM 200MG/5ML;40MG/5ML

N17560 002 Apr CMFD

SULFAMETHOXAZOLE AND TRIMETHOPRIM

AB TEVA PHARMS 200MG/5ML;40MG/5ML

N77612 001 Nov 13, 2006 Oct NEWA

SULFATRIM

AB ACTAVIS MID ATLANTIC 200MG/5ML;40MG/5ML

N18615 002 Jan 07, 1983 Jul CAHN

SULFATRIM PEDIATRIC

AB ACTAVIS MID ATLANTIC 200MG/5ML;40MG/5ML

N18615 001 Jan 07, 1983 Jul CAHN

SULFANILAMIDECREAM; VAGINAL
AVC

+ PHARMELLE 15%

N06530 003 Jan 27, 1987 Oct CMFD

SULFASALAZINE

TABLET; ORAL

SULFASALAZINE

@ RADIUS PHARMS 500MG

N80197 001 Aug CAHN

SULINDAC

TABLET; ORAL

CLINORIL

@ MERCK 150MG

N17911 001 Jun DISC

SULINDAC

@ RADIUS PHARMS 150MG

N73261 001 Sep 06, 1991 Jul CAHN

@ 200MG

N73262 001 Sep 06, 1991 Jul CAHN

SUMATRIPTAN SUCCINATE

INJECTABLE; SUBCUTANEOUS

IMITREX

+	GLAXOSMITHKLINE	EQ 6MG BASE/0.5ML (12MG/ML)	N20080 001	Dec 28, 1992	Feb	CDFR
	IMITREX STATDOSE					
+	GLAXOSMITHKLINE	EQ 4MG BASE/0.5ML (8MG/ML)	N20080 002	Feb 01, 2006	Feb	NEWA
+		EQ 6MG BASE/0.5ML (12MG/ML)	N20080 003	Dec 23, 1996	Feb	NEWA

SUNITINIB MALATE

CAPSULE; ORAL

SUTENT

	CPPI CV	EQ 12.5MG BASE	N21938 001	Jan 26, 2006	Jul	CAHN
		EQ 25MG BASE	N21938 002	Jan 26, 2006	Jul	CAHN
+		EQ 50MG BASE	N21938 003	Jan 26, 2006	Jul	CAHN
	PFIZER	EQ 12.5MG BASE	N21938 001	Jan 26, 2006	Jun	CPOT
		12.5MG	N21938 001	Jan 26, 2006	Jan	NEWA
		EQ 25MG BASE	N21938 002	Jan 26, 2006	Jun	CPOT
		25MG	N21938 002	Jan 26, 2006	Jan	NEWA
+		EQ 50MG BASE	N21938 003	Jan 26, 2006	Jun	CPOT
+		50MG	N21938 003	Jan 26, 2006	Jan	NEWA

TAMOXIFEN CITRATE

SOLUTION; ORAL

SOLTAMOX

+	ROSEMONT	EQ 10MG BASE/5ML	N21807 001	Oct 29, 2005	Nov	CAHN
+	SAVIENT PHARMA	EQ 10MG BASE/5ML	N21807 001	Oct 29, 2005	Jun	CAHN
+	SAVIENT PHARMS	EQ 10MG BASE/5ML	N21807 001	Oct 29, 2005	Jul	CAHN

TECHNETIUM TC-99M APCITIDE

INJECTABLE; INJECTION

ACUTECT

	@ CIS BIO INTL SA	N/A	N20887 001	Sep 14, 1998	May	DISC
		N/A	N20887 001	Sep 14, 1998	Apr	CAHN

TECHNETIUM TC-99M DEPREOTIDE

INJECTABLE; INJECTION

NEO TECT KIT

	@ CIS BIO INTL SA	N/A	N21012 001	Aug 03, 1999	May	DISC
+		N/A	N21012 001	Aug 03, 1999	Apr	CAHN

TECHNETIUM TC-99M MEBROFENIN KIT

INJECTABLE; INJECTION

CHOLETEC

+	BRACCO	N/A	N18963 001	Jan 21, 1987	Aug	CRLD
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TECHNETIUM TC-99M TETROFOSMIN KIT

INJECTABLE; INJECTION

MYOVUE 30ML

	@ GE HEALTHCARE	N/A	N20372 002	Jul 07, 2005	May	DISC
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TELBIVUDINE

TABLET; ORAL

TYZEKA

+	IDENIX	600MG	N22011 001	Oct 25, 2006	Oct	NEWA
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TEMAZEPAM

CAPSULE; ORAL

TEMAZEPAM

AB	ACTAVIS ELIZABETH	15MG	N71638 001	Aug 07, 1987	Jun	CAHN
AB		30MG	N71620 001	Aug 07, 1987	Jun	CAHN

TEMOZOLOMIDE

CAPSULE; ORAL

TEMODAR

SCHERING

140MG

N21029 005 Oct 19, 2006 Oct NEWA

180MG

N21029 006 Oct 19, 2006 Oct NEWA

TERBUTALINE SULFATE

INJECTABLE; INJECTION

BRETHINE

@ AAIPHARMA LLC

1MG/ML

N18571 001

Aug DISC

TERBUTALINE SULFATE

AP + BEDFORD 1MG/ML

N76770 001 Apr 23, 2004 Aug CRLD

TERCONAZOLE

CREAM; VAGINAL

TERAZOL 3

AB + ORTHO MCNEIL PHARM 0.8%

N19964 001 Feb 21, 1991 Sep CMFD

@ 0.8%

N19964 001 Feb 21, 1991 Jul DISC

TERAZOL 7

AB + ORTHO MCNEIL PHARM 0.4%

N19579 001 Dec 31, 1987 Sep CMFD

@ 0.4%

N19579 001 Dec 31, 1987 Jul DISC

TERCONAZOLE

AB TARO 0.4%

N76043 001 Jan 19, 2005 Sep CRLD

AB + 0.4%

N76043 001 Jan 19, 2005 Jul CRLD

AB 0.8%

N75953 001 Apr 06, 2004 Sep CRLD

BX + 0.8%

N75953 001 Apr 06, 2004 Jul CRLD

SUPPOSITORY; VAGINAL

TERAZOL 3

AB + ORTHO MCNEIL PHARM 80MG

N19641 001 May 24, 1988 Mar CFTG

TERCONAZOLE

AB ALTANA 80MG

N76850 001 Jul 12, 2006 Jun NEWA

AB PERRIGO NEW YORK 80MG

N77149 001 Mar 17, 2006 Mar NEWA

TESTOSTERONE

GEL; TRANSDERMAL

ANDROGEL

AB + UNIMED PHARMS 1%

N21015 001 Feb 28, 2000 Jan CTEC

TESTOSTERONE

AB WATSON LABS 1%

N76737 001 Jan 27, 2006 Jan NEWA

TESTOSTERONE CYPIONATE

INJECTABLE; INJECTION

TESTOSTERONE CYPIONATE

AO SANDOZ 100MG/ML

N40615 001 Aug 10, 2006 Aug NEWA

AO 200MG/ML

N40615 002 Aug 10, 2006 Aug NEWA

AO SYNERX PHARMA 200MG/ML

N40652 001 Dec 11, 2006 Nov NEWA

TESTOSTERONE ENANTHATE

	INJECTABLE; INJECTION					
	DELATESTRYL					
AO	+	@ INDEVUS PHARMS	200MG/ML	N09165 001	Jan	CAHN
			200MG/ML	N09165 003	Jan	CAHN
	TESTOSTERONE ETHANATE					
AO		PADDOCK	200MG/ML	N40575 001	Jun 14, 2006	May NEWA

TETRACYCLINE HYDROCHLORIDE

CAPSULE; ORAL					
ACHROMYCIN V					
@	RADIUS PHARMS	250MG	N50278 003	May	CAHN
@		500MG	N50278 001	May	CAHN
@	SCIREG INTL INC	250MG	N50278 003	Apr	CAHN
@		500MG	N50278 001	Apr	CAHN
SUSPENSION; ORAL					
SUMYCIN					
+	PAR PHARM	125MG/5ML	N60400 001	Aug	CAHN

THALLOUS CHLORIDE, TL-201

INJECTABLE; INJECTION					
THALLOUS CHLORIDE TL 201					
AP		TRACE RADIOCHEMICALS	1mCi/ML	N75569 001	Nov 21, 2001 Feb CAHN
INJECTABLE; INTRAVENOUS					
THALLOUS CHLORIDE TL 201					
AP	+	BRISTOL MYERS SQUIBB	2mCi/ML	N17806 002	Oct 09, 1998 Oct CFTG
AP		TCI MEDCL	2mCi/ML	N77698 001	Nov 09, 2006 Oct NEWA

THEOPHYLLINE

ELIXIR; ORAL					
THEOPHYLLINE					
AA		ACTAVIS MID ATLANTIC	80MG/15ML	N85863 001	Jul CAHN
SOLUTION; ORAL					
THEOLAIR					
		@ 3M	80MG/15ML	N86107 001	Aug DISC
THEOPHYLLINE					
	+	ROXANE	80MG/15ML	N87449 001	Sep 15, 1983 Aug CRLD
TABLET, EXTENDED RELEASE; ORAL					
THEOPHYLLINE					
AB		NOSTRUM	400MG	N40595 001	Apr 21, 2006 Apr NEWA
AB			600MG	N40560 002	Apr 21, 2006 Apr NEWA
T-PHYL					
		@ PURDUE FREDERICK	200MG	N88253 001	Aug 17, 1983 Jun DISC
UNIPHYL					
AB		PURDUE FREDERICK	400MG	N87571 001	Sep 01, 1982 Nov CRLD
AB	+		400MG	N87571 001	Sep 01, 1982 Apr CTEC
AB	+		600MG	N40086 001	Apr 15, 1996 Apr CTEC

THIABENDAZOLE

SUSPENSION; ORAL					
MINTEZOL					
		@ MERCK	500MG/5ML	N16097 001	Jun DISC

THIORIDAZINE HYDROCHLORIDE

CONCENTRATE; ORAL

THIORIDAZINE HYDROCHLORIDE

AA	ACTAVIS MID ATLANTIC	100MG/ML	N88229 001	Aug 23, 1983	Jul	CAHN
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TICARCILLIN DISODIUM

INJECTABLE; INJECTION

TICAR

@ GLAXOSMITHKLINE

EQ 1GM BASE/VIAL

N50497 001

May DISC

@

EQ 3GM BASE/VIAL

N50497 002

May DISC

@

EQ 20GM BASE/VIAL

N50497 004

May DISC

@

EQ 30GM BASE/VIAL

N50497 005 Apr 04, 1984 May DISC

TICLOPIDINE HYDROCHLORIDE

TABLET; ORAL

TICLOPIDINE HYDROCHLORIDE

AB	ACTAVIS ELIZABETH	250MG	N75253 001	Aug 20, 1999	Jun	CAHN
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TIMOLOL MALEATE

TABLET; ORAL

BLOCADREN

@ MERCK

5MG

N18017 001

Jun DISC

@

10MG

N18017 002

Jun DISC

@

20MG

N18017 004

Jun DISC

TIMOLOL MALEATE

AB	+	MYLAN	20MG	N72668 001	Jun 08, 1990	Jun	CRLD
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TINIDAZOLE

TABLET; ORAL

TINDAMAX

MISSION PHARMA

250MG

N21618 001 May 17, 2004 Jan CAHN

+

500MG

N21618 002 May 17, 2004 Jan CAHN

TIROFIBAN HYDROCHLORIDE

INJECTABLE; INJECTION

AGGRASTAT

+ MEDICURE

EQ 0.05MG BASE/ML

N20913 001 May 14, 1998 Aug CAHN

+

EQ 0.25MG BASE/ML

N20912 001 May 14, 1998 Aug CAHN

TIZANIDINE HYDROCHLORIDE

TABLET; ORAL

TIZANIDINE HYDROCHLORIDE

AB	ACTAVIS ELIZABETH	EQ 2MG BASE	N76283 001	Jul 12, 2002	Jun	CAHN
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AB		EQ 4MG BASE	N76283 002	Jul 12, 2002	Jun	CAHN
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TOBRAMYCIN

SOLUTION; INHALATION

TOBI

+ NOVARTIS PHARMS

300MG/5ML

N50753 001 Dec 22, 1997 Jul CAHN

TOBRAMYCIN SULFATE

INJECTABLE; INJECTION

TOBRAMYCIN

AP	+	ABRAXIS PHARM	EQ 40MG BASE/ML	N65122 002	Nov 29, 2002	Jul	CRLD
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TOLMETIN SODIUM

CAPSULE; ORAL

TOLMETIN SODIUM

AB	ACTAVIS ELIZABETH	EQ 400MG BASE	N73308 001	Jan 24, 1992	Jun	CAHN
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TABLET; ORAL

TOLMETIN SODIUM

AB	ACTAVIS ELIZABETH	EQ 600MG BASE	N73527 001	Jun 30, 1992	Jun	CAHN
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TOPIRAMATE

TABLET; ORAL

TOPAMAX

AB	+	ORTHO MCNEIL PHARM	25MG	N20505 004	Dec 24, 1996	Aug	CFTG
AB			100MG	N20505 001	Dec 24, 1996	Aug	CFTG
AB			200MG	N20505 002	Dec 24, 1996	Aug	CFTG

TOPIRAMATE

AB	MYLAN	25MG	N76314 001	Sep 11, 2006	Aug	NEWA
AB		100MG	N76314 002	Sep 11, 2006	Aug	NEWA
AB		200MG	N76314 003	Sep 11, 2006	Aug	NEWA

TRAMADOL HYDROCHLORIDE

TABLET; ORAL

TRAMADOL HYDROCHLORIDE

AB	ACTAVIS ELIZABETH	50MG	N75960 001	Jun 19, 2002	Jun	CAHN
	@ IVAX PHARMS	50MG	N75963 001	Jul 03, 2002	Jan	DISC

TRANDOLAPRIL

TABLET; ORAL

MAVIK

AB	ABBOTT	1MG	N20528 001	Apr 26, 1996	Nov	CFTG
AB		2MG	N20528 002	Apr 26, 1996	Nov	CFTG
AB	+	4MG	N20528 003	Apr 26, 1996	Nov	CFTG

TRANDOLAPRIL

AB	TEVA PHARMS	1MG	N77489 001	Dec 12, 2006	Nov	NEWA
AB		2MG	N77489 002	Dec 12, 2006	Nov	NEWA
AB		4MG	N77489 003	Dec 12, 2006	Nov	NEWA

TRANLYCYPROMINE SULFATE

TABLET; ORAL

PARNATE

AB	+	GLAXOSMITHKLINE	EQ 10MG BASE	N12342 003	Aug 16, 1985	Jun	CFTG
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TRANLYCYPROMINE SULFATE

AB	KALI LABS	EQ 10MG BASE	N40640 001	Jun 29, 2006	Jun	NEWA
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TRAVOPROST

SOLUTION/DROPS; OPHTHALMIC

TRAVATAN Z

+	ALCON	0.004%	N21994 001	Sep 21, 2006	Sep	NEWA
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TRAZODONE HYDROCHLORIDE

TABLET; ORAL

DESYREL

@ APOTHECON

@

@

50MG

100MG

150MG

N18207 001		Aug	DISC
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N18207 002		Aug	DISC
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N18207 003	Mar 25, 1985	Aug	DISC
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TABLET; ORAL

DESYREL

@ APOTHECON

300MG

N18207 004 Nov 07, 1988 Aug DISC

TRAZODONE HYDROCHLORIDE

AB ACTAVIS ELIZABETH

50MG

N71636 001 Apr 18, 1988 Jun CAHN

AB 100MG

N71514 001 Apr 18, 1988 Jun CAHN

AB APOTEX INC 100MG

N71196 001 Mar 25, 1987 Nov CAHN

AB 150MG

N71196 002 Apr 26, 1999 Nov CAHN

AB + 300MG

N71196 003 Apr 26, 1999 Nov CAHN

AB + BARR 300MG

N71196 003 Apr 26, 1999 Sep CRLD

TRETINOIN

CREAM; TOPICAL

TRETINOIN

AB TRIAX PHARMS LLC

0.025%

N75264 001 Dec 24, 1998 Nov CAHN

AB1 0.05%

N75265 001 Dec 24, 1998 Nov CAHN

AB 0.1%

N75213 001 Dec 24, 1998 Nov CAHN

TRIAMCINOLONE

TABLET; ORAL

ARISTOCORT

@ ASTELLAS

4MG

N11161 007 Sep DISC

>D> KENACORT

>D> + BRISTOL MYERS SQUIBB 4MG

N11283 006 Dec DISC

>A> @ 4MG

N11283 006 Dec DISC

+ 4MG

N11283 006 Sep CRLD

TRIAMCINOLONE ACETONIDE

CREAM; TOPICAL

ARISTOCORT A

@ ASTELLAS

0.025%

N88818 001 Oct 16, 1984 Sep DISC

@ 0.1%

N88819 001 Oct 16, 1984 Sep DISC

@ 0.5%

N88820 001 Oct 16, 1984 Sep DISC

TRIAMCINOLONE ACETONIDE

AT ACTAVIS MID ATLANTIC 0.1%

N87798 001 Jun 04, 1982 Jun CAHN

AT VINTAGE 0.025%

N40671 001 Jun 09, 2006 May NEWA

AT 0.1%

N40671 002 Jun 09, 2006 May NEWA

LOTION; TOPICAL

TRIAMCINOLONE ACETONIDE

AT VINTAGE 0.1%

N40672 002 Dec 13, 2006 Nov NEWA

OINTMENT; TOPICAL

TRIAMCINOLONE ACETONIDE

AT ACTAVIS MID ATLANTIC 0.1%

N87799 001 Jun 07, 1982 Jun CAHN

SPRAY, METERED; NASAL

NASACORT HFA

@ SANOFI AVENTIS US

0.055MG/SPRAY

N20784 001 Apr 07, 2004 Nov DISC

TRIAMCINOLONE DIACETATE

INJECTABLE; INJECTION

ARISTOCORT

@ SANDOZ

25MG/ML

N11685 003 Jan CAHN

@ 40MG/ML

N12802 001 Jan CAHN

TRIAMCINOLONE HEXACETONIDE

INJECTABLE; INJECTION
ARISTOSPAN

+	SANDOZ	5MG/ML	N16466 001	Jan	CAHN
+		20MG/ML	N16466 002	Jan	CAHN

TRICHLORMETHIAZIDE

TABLET; ORAL
TRICHLORMETHIAZIDE
@ TG UNITED LABS

4MG

N85568 001

May CAHN

TRIENTINE HYDROCHLORIDE

CAPSULE; ORAL
SYPRINE

+	ATON	250MG	N19194 001	Nov 08, 1985	Oct	CAHN
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TRIFLUOPERAZINE HYDROCHLORIDE

INJECTABLE; INJECTION
STELAZINE

@ GLAXOSMITHKLINE EQ 2MG BASE/ML

N11552 005

May DISC

TRIMIPRAMINE MALEATE

CAPSULE; ORAL
SURMONTIL

AB	ODYSSEY PHARMS	EQ 25MG BASE	N16792 001	Jul	CFTG	
AB		EQ 50MG BASE	N16792 002	Jul	CFTG	
AB	+	EQ 100MG BASE	N16792 003	Sep 15, 1982	Jul	CFTG
	TRIMIPRAMINE MALEATE					
AB	ACTAVIS TOTOWA	EQ 25MG BASE	N77361 001	Aug 02, 2006	Jul	NEWA
AB		EQ 50MG BASE	N77361 002	Aug 02, 2006	Jul	NEWA
AB		EQ 100MG BASE	N77361 003	Aug 02, 2006	Jul	NEWA

TRIPLENNAMINE HYDROCHLORIDE

TABLET; ORAL
PBZ

@ NOVARTIS 50MG

N05914 002

Jan DISC

TROLEANDOMYCIN

CAPSULE; ORAL
TAO

@ PFIZER EQ 250MG BASE

N50336 002

Mar DISC

UNOPROSTONE ISOPROPYL

SOLUTION/DROPS; OPHTHALMIC
RESCULA

@ R TECH UENO LTD 0.15%

N21214 001	Aug 03, 2000	Jun	DISC
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+
 0.15% |

N21214 001	Aug 03, 2000	Feb	CAHN
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UREA

INJECTABLE; INJECTION
UREAPHIL

@ HOSPIRA 40GM/VIAL

N12154 001

Jun DISC

UROKINASE

INJECTABLE; INJECTION

ABBOKINASE

@ IMARX THERAP

5,000 IU/VIAL

N21846 003

Apr CAHN

@

9,000 IU/VIAL

N21846 002

Apr CAHN

+

250,000 IU/VIAL

N21846 001

Apr CAHN

URSODIOL

CAPSULE; ORAL

URSODIOL

AB COREPHARMA

300MG

N77895 001 Jul 27, 2006 Jul NEWA

VALPROIC ACID

SOLUTION; ORAL

VALPROIC ACID

AA VINTAGE

250MG/5ML

N77960 001 Oct 13, 2006 Oct NEWA

VALRUBICIN

SOLUTION; INTRAVESICAL

VALSTAR PRESERVATIVE FREE

+ VALERA

40MG/ML

N20892 001 Sep 25, 1998 Jul CAHN

VARENICLINE TARTRATE

TABLET; ORAL

CHANTIX

CPPI CV

EQ 0.5MG BASE

N21928 001 May 10, 2006 Jul CAHN

+

EQ 1MG BASE

N21928 002 May 10, 2006 Jul CAHN

PFIZER

EQ 0.5MG BASE

N21928 001 May 10, 2006 May NEWA

+

EQ 1MG BASE

N21928 002 May 10, 2006 May NEWA

PFIZER INC

EQ 0.5MG BASE

N21928 001 May 10, 2006 Sep CAHN

+

EQ 1MG BASE

N21928 002 May 10, 2006 Sep CAHN

VECURONIUM BROMIDE

INJECTABLE; INJECTION

VECURONIUM BROMIDE

AP + BEDFORD

20MG/VIAL

N75549 002 Jun 13, 2000 Jan CRLD

VENLAFAXINE HYDROCHLORIDE

TABLET; ORAL

EFFEXOR

AB WYETH PHARMS INC

EQ 25MG BASE

N20151 002 Dec 28, 1993 Jul CFTG

AB

EQ 37.5MG BASE

N20151 006 Dec 28, 1993 Jul CFTG

AB +

EQ 50MG BASE

N20151 003 Dec 28, 1993 Jul CFTG

AB

EQ 75MG BASE

N20151 004 Dec 28, 1993 Jul CFTG

AB

EQ 100MG BASE

N20151 005 Dec 28, 1993 Jul CFTG

VENLAFAXINE HYDROCHLORIDE

AB TEVA

EQ 25MG BASE

N76690 001 Aug 03, 2006 Jul CTNA

AB

EQ 37.5MG BASE

N76690 002 Aug 03, 2006 Jul NEWA

AB

EQ 50MG BASE

N76690 003 Aug 03, 2006 Jul NEWA

AB

EQ 75MG BASE

N76690 004 Aug 03, 2006 Jul CTNA

AB

EQ 100MG BASE

N76690 005 Aug 03, 2006 Jul NEWA

VERAPAMIL HYDROCHLORIDE

TABLET; ORAL

VERAPAMIL HYDROCHLORIDE

AB	ACTAVIS ELIZABETH	80MG	N71019 001	Sep 24, 1986	Jun	CAHN
AB		120MG	N70468 001	Sep 24, 1986	Jun	CAHN
	@ RADIUS PHARMS	80MG	N71880 001	Apr 05, 1988	Jul	CAHN
	@	120MG	N71881 001	Apr 05, 1988	Jul	CAHN

TABLET, EXTENDED RELEASE; ORAL

ISOPTIN SR

AB	+ FSC	120MG	N19152 003	Mar 06, 1991	Nov	CAHN
AB	+	180MG	N19152 002	Dec 15, 1989	Nov	CAHN
AB	+	240MG	N19152 001	Dec 16, 1986	Nov	CAHN

VORINOSTAT

CAPSULE; ORAL

ZOLINZA

+	MERCK	100MG	N21991 001	Oct 06, 2006	Oct	NEWA
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WARFARIN SODIUM

TABLET; ORAL

WARFARIN SODIUM

AB	PLIVA	1MG	N40616 009	Jul 05, 2006	Jun	NEWA
AB		2MG	N40616 001	Jul 05, 2006	Jun	NEWA
AB		2.5MG	N40616 002	Jul 05, 2006	Jun	NEWA
AB		3MG	N40616 003	Jul 05, 2006	Jun	NEWA
AB		4MG	N40616 004	Jul 05, 2006	Jun	NEWA
AB		5MG	N40616 005	Jul 05, 2006	Jun	NEWA
AB		6MG	N40616 006	Jul 05, 2006	Jun	NEWA
AB		7.5MG	N40616 007	Jul 05, 2006	Jun	NEWA
AB		10MG	N40616 008	Jul 05, 2006	Jun	NEWA
AB	ZYDUS PHARMS USA	1MG	N40663 001	May 30, 2006	May	NEWA
AB		2MG	N40663 002	May 30, 2006	May	NEWA
AB		2.5MG	N40663 003	May 30, 2006	May	NEWA
AB		3MG	N40663 004	May 30, 2006	May	NEWA
AB		4MG	N40663 005	May 30, 2006	May	NEWA
AB		5MG	N40663 006	May 30, 2006	May	NEWA
AB		6MG	N40663 007	May 30, 2006	May	NEWA
AB		7.5MG	N40663 008	May 30, 2006	May	NEWA
AB		10MG	N40663 009	May 30, 2006	May	NEWA

WATER FOR INJECTION, STERILE

LIQUID; N/A

STERILE WATER FOR INJECTION IN PLASTIC CONTAINER

AP	TARO PHARMS IRELAND	100%	N77393 001	Aug 11, 2006	Aug	NEWA
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ZALEPLON

CAPSULE; ORAL

SONATA

>D>	JONES PHARMA	5MG	N20859 001	Aug 13, 1999	Dec	CAHN
>D>	+	10MG	N20859 002	Aug 13, 1999	Dec	CAHN
>A>	KING PHARMS	5MG	N20859 001	Aug 13, 1999	Dec	CAHN
>A>	+	10MG	N20859 002	Aug 13, 1999	Dec	CAHN

ZIDOVUDINECAPSULE; ORAL
RETROVIR

AB	+	GLAXOSMITHKLINE	100MG	N19655 001	Mar 19, 1987	Mar	CFTG
ZIDOVUDINE							
AB		AUROBINDO PHARMA LTD	100MG	N78128 001	Mar 27, 2006	Mar	NEWA

ZIPRASIDONE HYDROCHLORIDESUSPENSION; ORAL
GEODON

+	PFIZER INC	EQ 10MG BASE/ML	N21483 001	Mar 29, 2006	Mar	NEWA
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ZONISAMIDECAPSULE; ORAL
ZONISAMIDE

AB		BANNER PHARMACAPS	25MG	N77813 001	Aug 16, 2006	Aug	NEWA
AB			50MG	N77813 002	Aug 16, 2006	Aug	NEWA
AB			100MG	N77813 003	Aug 16, 2006	Aug	NEWA
AB		DR REDDYS LABS LTD	25MG	N77891 001	Sep 29, 2006	Sep	NEWA
AB			50MG	N77891 002	Sep 29, 2006	Sep	NEWA
AB		GLENMARK PHARMS	25MG	N77651 001	Jan 30, 2006	Jan	NEWA
AB			50MG	N77651 002	Jan 30, 2006	Jan	NEWA
AB			100MG	N77651 003	Jan 30, 2006	Jan	NEWA
AB		INVAGEN PHARMS	25MG	N77869 001	May 31, 2006	May	NEWA
AB			50MG	N77869 002	May 31, 2006	May	NEWA
AB			100MG	N77869 003	May 31, 2006	May	NEWA
AB		SUN PHARM INDS (IN)	25MG	N77634 001	Mar 17, 2006	Mar	NEWA
AB			50MG	N77634 002	Mar 17, 2006	Mar	NEWA
AB			100MG	N77634 003	Mar 17, 2006	Mar	NEWA
AB		WATSON LABS	25MG	N77650 001	Apr 20, 2006	Apr	NEWA
AB			50MG	N77650 002	Apr 20, 2006	Apr	NEWA
AB			100MG	N77650 003	Apr 20, 2006	Apr	NEWA
AB		WOCKHARDT	25MG	N77636 003	Jul 27, 2006	Jul	NEWA
AB			50MG	N77636 002	Jul 27, 2006	Jul	NEWA
AB		ZYDUS PHARMS USA	25MG	N77625 001	Oct 16, 2006	Oct	NEWA
AB			50MG	N77625 002	Oct 16, 2006	Oct	NEWA
AB			100MG	N77625 003	Oct 16, 2006	Oct	NEWA

OTC DRUG PRODUCT LIST - 26TH EDITION

OTC DRUG PRODUCT LIST - CUMULATIVE SUPPLEMENT 12 - December 2006

2-1

ACETAMINOPHEN

SUPPOSITORY; RECTAL

ACETAMINOPHEN

ACTAVIS MID ATLANTIC	120MG	N18337 003	Sep 12, 1983	Jun	CAHN
	325MG	N18337 002		Jun	CAHN
+	650MG	N18337 001		Jun	CAHN
INFANTS' FEVERALL					
ACTAVIS MID ATLANTIC	80MG	N18337 004	Aug 26, 1992	Jun	CAHN
TYLENOL					
@ MCNEIL CONS	120MG	N17756 002		Jun	CAHN
@	650MG	N17756 001		Jun	CAHN
TABLET, EXTENDED RELEASE; ORAL					
TYLENOL (CAPLET)					
+	MCNEIL CONS 650MG	N19872 001	Jun 08, 1994	Jun	CAHN
TYLENOL (GELTAB)					
+	MCNEIL CONS 650MG	N19872 002	Jan 11, 2001	Jun	CAHN

AVOBENZONE; ECAMSULE; OCTOCRYLENE

CREAM; TOPICAL

ANTHELIOS SX

+	LOREAL USA	2%;2%;10%	N21502 001	Jul 21, 2006	Jul	NEWA
CAPITAL SOLEIL 15						
+	LOREAL USA	2%;3%;10%	N21501 001	Oct 02, 2006	Oct	NEWA

AVOBENZONE; ECAMSULE; OCTOCRYLENE; TITANIUM DIOXIDE

CREAM; TOPICAL

ANTHELIOS 20

+	LOREAL USA	2%;2%;10%;2%	N21471 001	Oct 05, 2006	Oct	NEWA
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CHLORHEXIDINE GLUCONATE

CLOTH; TOPICAL

CHLORHEXIDINE GLUCONATE

+	SAGE PRODS	2%	N21669 001	Apr 25, 2005	Oct	CTNA
HALO						
+	SAGE PRODS	2%	N21669 001	Apr 25, 2005	Aug	CTNA

SPONGE; TOPICAL

CHLORHEXIDINE GLUCONATE

	BECTON DICKINSON	4%	N72525 001	Oct 24, 1989	Aug	CAHN
@	KENDALL IL	4%	N19490 001	Mar 27, 1987	Aug	DISC

CHLORHEXIDINE GLUCONATE; ISOPROPYL ALCOHOL

SPONGE; TOPICAL

CHLORAPREP ONE-STEP

+	MEDI FLEX INC	2%;70% (26ML)	N20832 006	Nov 21, 2006	Nov	NEWA
CHLORAPREP WITH TINT						
+	MEDI FLEX INC	2%;70% (10.5ML)	N20832 005	Apr 03, 2006	Apr	NEWA

CLOTRIMAZOLE

CREAM; VAGINAL

CLOTRIMAZOLE

	ACTAVIS MID ATLANTIC	1%	N74165 001	Jul 16, 1993	Jun	CAHN
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CROMOLYN SODIUM

SPRAY, METERED; NASAL

CROMOLYN SODIUM

ACTAVIS MID ATLANTIC 5.2MG/SPRAY

N74800 001 Jul 26, 2001 Jul CAHN

ALPHARMA US PHARMS 5.2MG/SPRAY

N74800 001 Jul 26, 2001 Jan CPOT

+ BAUSCH AND LOMB 5.2MG/SPRAY

N75702 001 Jul 03, 2001 Jan CRLD

NASALCROM

>A> @ MCNEIL CONS 5.2MG/SPRAY

N20463 001 Jan 03, 1997 Dec CAHN

>D> @ PHARMACIA UPJOHN 5.2MG/SPRAY

N20463 001 Jan 03, 1997 Dec CAHN

@ 5.2MG/SPRAY

N20463 001 Jan 03, 1997 Jan DISC

DEXTROMETHORPHAN HYDROBROMIDE; GUAIFENESIN

TABLET, EXTENDED RELEASE; ORAL

MUCINEX DM

ADAMS RESP THERAP 30MG;600MG

N21620 002 Apr 29, 2004 May CAHN

+ 60MG;1.2GM

N21620 001 Apr 29, 2004 May CAHN

DEXTROMETHORPHAN POLISTIREX

SUSPENSION, EXTENDED RELEASE; ORAL

DELSYM

+ ADAMS RESP THERAP EQ 30MG HBR/5ML

N18658 001 Oct 08, 1982 Jun CAHN

FAMOTIDINE

TABLET; ORAL

FAMOTIDINE

DR REDDYS LABS LTD 20MG

N77367 001 Sep 25, 2006 Sep NEWA

PERRIGO 20MG

N77351 001 Sep 25, 2006 Sep NEWA

GUAIFENESIN

TABLET, EXTENDED RELEASE; ORAL

HUMABID

+ ADAMS RESP THERAP 1.2GM

N21282 002 Dec 18, 2002 Aug CTNA

MUCINEX

ADAMS RESP THERAP 600MG

N21282 001 Jul 12, 2002 May CAHN

+ 1.2GM

N21282 002 Dec 18, 2002 May CAHN

GUAIFENESIN; PSEUDOEPHEDRINE HYDROCHLORIDE

TABLET, EXTENDED RELEASE; ORAL

MUCINEX D

ADAMS RESP THERAP 600MG;60MG

N21585 001 Jun 22, 2004 May CAHN

+ 1.2GM;120MG

N21585 002 Jun 22, 2004 May CAHN

IBUPROFEN

SUSPENSION/DROPS; ORAL

CHILDREN'S MOTRIN

+ MCNEIL CONS 40MG/ML

N20603 001 Jun 10, 1996 Jun CAHN

SUSPENSION; ORAL

CHILDREN'S ELIXSURE

ALTERNA-TCHP LLC 100MG/5ML

N21604 001 Jan 07, 2004 Jul CAHN

IBUPROFEN

ACTAVIS MID ATLANTIC 100MG/5ML

N74916 001 Apr 30, 1999 Jul CAHN

TABLET, CHEWABLE; ORAL

CHILDREN'S MOTRIN

MCNEIL CONS 50MG

N20601 001 Nov 15, 1996 Jun CAHN

TABLET, CHEWABLE; ORAL									
JUNIOR STRENGTH MOTRIN									
+	MCNEIL CONS	100MG	N20601	003	Nov 15, 1996	Jun	CAHN		
TABLET; ORAL									
JUNIOR STRENGTH MOTRIN									
	MCNEIL CONS	100MG	N20602	001	Jun 10, 1996	Jun	CAHN		
<u>IBUPROFEN; PSEUDOEPHEDRINE HYDROCHLORIDE</u>									
SUSPENSION; ORAL									
CHILDREN'S MOTRIN COLD									
+	MCNEIL CONS	100MG/5ML;15MG/5ML	N21128	001	Aug 01, 2000	Jun	CAHN		
TABLET; ORAL									
IBUPROFEN AND PSEUDOEPHEDRINE HYDROCHLORIDE									
	DR REDDYS LABS LTD	200MG;30MG	N77628	001	Aug 14, 2006	Aug	NEWA		
SINE-AID IB									
	MCNEIL CONS	200MG;30MG	N19899	001	Dec 31, 1992	Jun	CAHN		
<u>INSULIN PURIFIED PORK</u>									
INJECTABLE; INJECTION									
REGULAR ILETIN II (PORK)									
	@ LILLY	100 UNITS/ML	N18344	001		Jun	DISC		
<u>INSULIN RECOMBINANT HUMAN</u>									
INJECTABLE; INJECTION									
VELOSULIN BR									
	@ NOVO NORDISK INC	100 UNITS/ML	N21028	001	Jul 19, 1999	Nov	DISC		
<u>INSULIN SUSP ISOPHANE BEEF/PORK</u>									
INJECTABLE; INJECTION									
NPH ILETIN I (BEEF-PORK)									
	@ LILLY	100 UNITS/ML	N17936	002		Jun	DISC		
<u>INSULIN SUSP ISOPHANE PURIFIED PORK</u>									
INJECTABLE; INJECTION									
NPH ILETIN II (PORK)									
	@ LILLY	100 UNITS/ML	N18345	001		Jun	DISC		
<u>INSULIN ZINC SUSP EXTENDED RECOMBINANT HUMAN</u>									
INJECTABLE; INJECTION									
HUMULIN U									
	@ LILLY	100 UNITS/ML	N19571	002	Jun 10, 1987	Jun	DISC		
<u>INSULIN ZINC SUSP PURIFIED PORK</u>									
INJECTABLE; INJECTION									
LENTE ILETIN II (PORK)									
	@ LILLY	100 UNITS/ML	N18347	001		Jun	DISC		
<u>INSULIN ZINC SUSP RECOMBINANT HUMAN</u>									
INJECTABLE; INJECTION									
NOVOLIN L									
	@ NOVO NORDISK INC	100 UNITS/ML	N19965	001	Jun 25, 1991	Nov	DISC		
<u>IODINE POVACRYLEX; ISOPROPYL ALCOHOL</u>									
SPONGE; TOPICAL									
DURAPREP									
+	3M	EQ 0.7% IODINE;74% (26ML)	N21586	002	Sep 29, 2006	Oct	CRLD		

SPONGE; TOPICAL

DURAPREP

+ 3M	EQ 0.7% IODINE; 74% (6ML)	N21586 001	Sep 29, 2006	Sep	NEWA
	EQ 0.7% IODINE;74% (26ML)	N21586 002	Sep 29, 2006	Sep	NEWA

KETOCONAZOLE

SHAMPOO; TOPICAL

NIZORAL A-D

+ MCNEIL CONS	1%	N20310 001	Oct 10, 1997	Jun	CAHN
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KETOPROFEN

TABLET; ORAL

ACTRON

@ BAYER

12.5MG

N20499 001	Oct 06, 1995	Feb	DISC
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ORUDIS KT

@ WYETH CONS

12.5MG

N20429 001	Oct 06, 1995	Feb	DISC
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KETOTIFEN FUMARATE

SOLUTION/DROPS; OPHTHALMIC

ALAWAY

>A>

+ ALIMERA SCIENCES INC	EQ 0.025% BASE	N21996 001	Dec 01, 2006	Dec	NEWA
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ZADITOR

+ NOVARTIS	EQ 0.025% BASE	N21066 002	Oct 19, 2006	Oct	NEWA
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>A>

LEVONORGESTREL

TABLET; ORAL

PLAN B

+ DURAMED	0.75MG	N21045 002	Aug 24, 2006	Aug	NEWA
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LOPERAMIDE HYDROCHLORIDE

SOLUTION; ORAL

IMODIUM A-D

+ MCNEIL CONS	1MG/5ML	N19487 001	Mar 01, 1988	Jun	CAHN
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SUSPENSION; ORAL

IMODIUM A-D

+ MCNEIL	1MG/7.5ML	N19487 002	Jul 08, 2004	Mar	CDFR
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+ MCNEIL CONS	1MG/7.5ML	N19487 002	Jul 08, 2004	Jun	CAHN
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TABLET; ORAL

IMODIUM A-D

+ MCNEIL CONS	2MG	N19860 001	Nov 22, 1989	Jun	CAHN
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LOPERAMIDE HYDROCHLORIDE; SIMETHICONE

TABLET; ORAL

IMODIUM ADVANCED

+ MCNEIL CONS	2MG;125MG	N21140 001	Nov 30, 2000	Jun	CAHN
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LOPERAMIDE HYDROCHLORIDE AND SIMETHICONE

RANBAXY

2MG;125MG

N77500 001	Sep 06, 2006	Aug	NEWA
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LORATADINE

SYRUP; ORAL

LORATADINE

SILARX

1MG/ML

N77421 001	Jun 29, 2006	Jun	NEWA
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TABLET, CHEWABLE; ORAL

CHILDREN'S CLARITIN

+ SCHERING PLOUGH	5MG	N21891 001	Aug 23, 2006	Aug	NEWA
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TABLET, ORALLY DISINTEGRATING; ORAL						
>A>	CLARITIN REDITABS					
>A>	+ SCHERING PLOUGH	5MG	N21993	001	Dec 12, 2006	Dec NEWA
TABLET; ORAL						
	LORATADINE					
	APOTEX	10MG	N76471	001	Feb 14, 2006	Jan NEWA

MICONAZOLE NITRATE

CREAM, SUPPOSITORY; TOPICAL, VAGINAL

M-ZOLE 3 COMBINATION PACK

ACTAVIS MID ATLANTIC	2%, 200MG	N74926	001	Apr 16, 1999	Jun CAHN
	2%, 200MG	N74926	001	Apr 16, 1999	Jun CAHN
	2%, 200MG	N74926	001	Apr 16, 1999	Jun CAHN
	2%, 200MG	N74926	001	Apr 16, 1999	Jun CAHN
	2%, 200MG	N74926	001	Apr 16, 1999	Jun CAHN
	2%, 200MG	N74926	001	Apr 16, 1999	Jun CAHN
	2%, 200MG	N74926	001	Apr 16, 1999	Jun CAHN
	2%, 200MG	N74926	001	Apr 16, 1999	Jun CAHN

M-ZOLE 7 DUAL PACK

ACTAVIS MID ATLANTIC	2%, 100MG	N74586	001	Jul 17, 1997	Jun CAHN
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CREAM; VAGINAL

MICONAZOLE 7

ACTAVIS MID ATLANTIC	2%	N74164	001	Mar 29, 1996	Jun CAHN
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SUPPOSITORY; VAGINAL

MICONAZOLE NITRATE

ACTAVIS MID ATLANTIC	100MG	N73507	001	Nov 19, 1993	Jun CAHN
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MINOXIDIL

AEROSOL, FOAM; TOPICAL

MEN'S ROGAINE

>A>	+ JOHNSON AND JOHNSON	5%	N21812	001	Jan 20, 2006	Dec CAHN
>D>	+ PHARMACIA AND UPJOHN	5%	N21812	001	Jan 20, 2006	Dec CAHN
	+	5%	N21812	001	Jan 20, 2006	Jan NEWA

SOLUTION; TOPICAL

MINOXIDIL (FOR MEN)

ACTAVIS MID ATLANTIC	2%	N74588	001	Apr 05, 1996	Jul CAHN
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MINOXIDIL EXTRA STRENGTH (FOR MEN)

ACTAVIS MID ATLANTIC	5%	N75518	001	Nov 17, 2000	Jul CAHN
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ROGAINE (FOR MEN)

>A>	+ JOHNSON AND JOHNSON	2%	N19501	002	Feb 09, 1996	Dec CAHN
>D>	+ PHARMACIA AND UPJOHN	2%	N19501	002	Feb 09, 1996	Dec CAHN

ROGAINE (FOR WOMEN)

>A>	+ JOHNSON AND JOHNSON	2%	N19501	003	Feb 09, 1996	Dec CAHN
>D>	+ PHARMACIA AND UPJOHN	2%	N19501	003	Feb 09, 1996	Dec CAHN

ROGAINE EXTRA STRENGTH (FOR MEN)

>A>	+ JOHNSON AND JOHNSON	5%	N20834	001	Nov 14, 1997	Dec CAHN
>D>	+ PHARMACIA AND UPJOHN	5%	N20834	001	Nov 14, 1997	Dec CAHN

NAPHAZOLINE HYDROCHLORIDE; PHENIRAMINE MALEATE

SOLUTION/DROPS; OPHTHALMIC

VISINE-A

>A>	JOHNSON AND JOHNSON	0.025%;0.3%	N20485	001	Jan 31, 1996	Dec CAHN
>D>	PFIZER	0.025%;0.3%	N20485	001	Jan 31, 1996	Dec CAHN

NAPROXEN SODIUM

CAPSULE; ORAL

NAPROXEN SODIUM

+	BANNER PHARMACAPS	EQ 200MG BASE	N21920 001	Feb 17, 2006	Oct	CRLD
		EQ 200MG BASE	N21920 001	Feb 17, 2006	Sep	CMFD
@		EQ 200MG BASE	N21920 001	Feb 17, 2006	Jul	DISC
+		EQ 200MG BASE	N21920 001	Feb 17, 2006	Feb	NEWA

NAPROXEN SODIUM; PSEUDOEPHEDRINE HYDROCHLORIDE

TABLET, EXTENDED RELEASE; ORAL

NAPROXEN SODIUM AND PSEUDOEPHEDRINE HYDROCHLORIDE

DR REDDYS LABS INC	EQ 220MG BASE;120MG	N77381 001	Sep 27, 2006	Sep	NEWA
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NICOTINE

FILM, EXTENDED RELEASE; TRANSDERMAL

NICOTROL

>A>	@ MCNEIL CONS	15MG/16HR	N20536 001	Jul 03, 1996	Dec	CAHN
>D>	@ PHARMACIA AND UPJOHN	15MG/16HR	N20536 001	Jul 03, 1996	Dec	CAHN

NICOTINE POLACRILEX

GUM, CHEWING; BUCCAL

NICORETTE

GLAXOSMITHKLINE	EQ 2MG BASE	N18612 004	Sep 25, 2000	Mar	CRLD
	EQ 2MG BASE	N18612 003	Dec 23, 1998	Mar	CRLD
	EQ 4MG BASE	N20066 003	Dec 23, 1998	Mar	CRLD
	EQ 4MG BASE	N20066 004	Sep 25, 2000	Mar	CRLD

NICORETTE (MINT)

GLAXOSMITHKLINE	EQ 2MG BASE	N18612 003	Dec 23, 1998	Apr	CTNA
	EQ 4MG BASE	N20066 003	Dec 23, 1998	Apr	CTNA

NICOTINE POLACRILEX

PERRIGO	EQ 2MG BASE	N76776 001	Sep 16, 2004	Apr	CTNA
	EQ 2MG BASE	N76777 001	Sep 16, 2004	Apr	CTNA
	EQ 4MG BASE	N76778 001	Sep 16, 2004	Apr	CTNA
	EQ 4MG BASE	N76779 001	Sep 16, 2004	Apr	CTNA
PERRIGO R AND D	EQ 2MG BASE	N78325 001	Oct 30, 2006	Oct	NEWA
	EQ 4MG BASE	N78326 001	Oct 30, 2006	Oct	NEWA
WATSON LABS	EQ 2MG BASE	N76569 001	Jul 29, 2004	Apr	CTNA
	EQ 4MG BASE	N76568 002	Jul 29, 2004	Apr	CTNA

TROCHE/LOZENGE; ORAL

NICOTINE POLACRILEX

PERRIGO R AND D	EQ 2MG BASE	N77007 001	Jan 31, 2006	Jan	NEWA
	EQ 4MG BASE	N77007 002	Jan 31, 2006	Jan	NEWA

OXYMETAZOLINE HYDROCHLORIDE

SOLUTION/DROPS; OPHTHALMIC

VISINE L.R.

>A>	+	JOHNSON AND JOHNSON	0.025%	N19407 001	Mar 31, 1989	Dec	CAHN
>D>	+	PFIZER	0.025%	N19407 001	Mar 31, 1989	Dec	CAHN

PERMETHRIN

LOTION; TOPICAL

PERMETHRIN

ACTAVIS MID ATLANTIC	1%	N75014 001	Mar 28, 2000	Jun	CAHN
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POLYETHYLENE GLYCOL 3350

FOR SOLUTION; ORAL

MIRALAX

+	BRAINTREE	17GM/SC00PFUL	N22015	001	Oct 06, 2006	Oct	NEWA
+	SCHERING PLOUGH	17GM/SC00PFUL	N22015	001	Oct 06, 2006	Nov	CAHN

POVIDONE-IODINE

SOLUTION; TOPICAL

E-Z PREP

@	CLINIPAD	10%	N19382	001	Jul 25, 1989	Aug	DISC
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SPONGE; TOPICAL

E-Z PREP

@	CLINIPAD	5%	N19382	002	Jul 25, 1989	Aug	DISC
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E-Z PREP 220

@	CLINIPAD	5%	N19382	003	Jul 25, 1989	Aug	DISC
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RANITIDINE HYDROCHLORIDE

TABLET, EFFERVESCENT; ORAL

ZANTAC 75

>A>	@	MCNEIL CONS	EQ 75MG BASE	N20745	001	Feb 26, 1998	Dec	CAHN
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>D>	@	WARNER LAMBERT	EQ 75MG BASE	N20745	001	Feb 26, 1998	Dec	CAHN
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TABLET; ORAL

RANITIDINE

WOCKHARDT	EQ 75MG BASE	N76760	001	Feb 24, 2006	Feb	NEWA
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ZANTAC 150

>A>	+	MCNEIL CONS	EQ 150MG BASE	N21698	001	Aug 31, 2004	Dec	CAHN
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>D>	+	PFIZER CONS HLTHCARE	EQ 150MG BASE	N21698	001	Aug 31, 2004	Dec	CAHN
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ZANTAC 75

>A>		MCNEIL CONS	EQ 75MG BASE	N20520	001	Dec 19, 1995	Dec	CAHN
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>D>		WARNER LAMBERT	EQ 75MG BASE	N20520	001	Dec 19, 1995	Dec	CAHN
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TERBINAFINE

GEL; TOPICAL

LAMISIL AT

+	NOVARTIS	1%	N21958	001	Jul 24, 2006	Jul	NEWA
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**DRUG PRODUCTS WITH APPROVAL UNDER SECTION 505 OF THE ACT
ADMINISTERED BY THE CENTER FOR BIOLOGICS EVALUATION AND RESEARCH LIST**

CUMULATIVE SUPPLEMENT NUMBER 12 DECEMBER 2006

NO DECEMBER 2006 APPROVALS

ORPHAN PRODUCT DESIGNATIONS AND APPROVALS LIST

The list of List of Orphan Designations and Approvals is available at:

<http://www.fda.gov/orphan/designat/list.htm>

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

NO DECEMBER 2006 ADDITIONS

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnotes for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>ABACAVIR SULFATE; LAMIVUDINE - EPZICOM</u>					
021652 001	5034394	Dec 18, 2011	DS DP	D-40	Aug 02, 2007
	5034394*PED	Jun 18, 2012			
	5047407	Nov 17, 2009	DS DP	U-257	
	5047407*PED	May 17, 2010			
	5089500	Jun 26, 2009		U-257	
	5089500*PED	Dec 26, 2009			
	5905082	May 18, 2016	DS DP		
	5905082*PED	Nov 18, 2016			
	6294540	May 14, 2018	DS DP	U-257	
	6294540*PED	Nov 14, 2018			
<u>ABACAVIR SULFATE; LAMIVUDINE; ZIDOVUDINE - TRIZIVIR</u>					
021205 001	5034394	Dec 18, 2011	DS DP		
	5034394*PED	Jun 18, 2012			
<u>ACETAMINOPHEN; TRAMADOL HYDROCHLORIDE - ULTRACET</u>					
021123 001	RE39221	Aug 09, 2011	DS DP	U-55	
<u>ALBUTEROL SULFATE - PROAIR HFA</u>					
021457 001				I-235	Feb 03, 2009
<u>ALBUTEROL SULFATE - VENTOLIN HFA</u>					
020983 001	6131566	Apr 14, 2015	DP	U-716	
	6131566	Apr 14, 2015	DP	U-589	
	6532955	Apr 14, 2015	DP	U-716	
	6532955	Apr 14, 2015	DP	U-590	
<u>ALENDRONATE SODIUM; CHOLECALCIFEROL - FOSAMAX PLUS D</u>					
021762 001				NC	Apr 07, 2008
<u>ALFUZOSIN HYDROCHLORIDE - UROXATRAL</u>					
021287 001	4661491	May 27, 2007		U-706	
<u>ALITRETINOIN - PANRETIN</u>					
020886 001	7056954	Aug 02, 2012	DP		
<u>AMLODIPINE BESYLATE; BENAZEPRIL HYDROCHLORIDE - LOTREL</u>					
020364 006	4879303	Mar 25, 2007	DS DP	NS	Apr 11, 2009
	6162802	Dec 19, 2017	DS DP	U-185	
<u>AMLODIPINE BESYLATE; BENAZEPRIL HYDROCHLORIDE - LOTREL</u>					
020364 007	4879303	Mar 25, 2007	DS DP	NS	Apr 11, 2009
	6162802	Dec 19, 2017	DS DP	U-185	
<u>ANIDULAFUNGIN - ERAXIS</u>					
021632 001	5965525	Oct 12, 2016	DS DP	U-540	NCE
	6384013	Mar 19, 2012	DS		
	6743777	Mar 19, 2012	DP	U-540	
	6960564	Apr 12, 2021	DP	U-540	
<u>ANIDULAFUNGIN - ERAXIS</u>					
021632 002				>A> NCE	Feb 17, 2011
<u>APREPITANT - EMEND</u>					
021549 001	5145684	Jan 25, 2011	DP		
	5719147	Jun 29, 2012	DS DP		
	6096742	Jul 01, 2018	DS DP	U-745	
	6235735	Jun 29, 2012		U-746	
	6235735	Jun 29, 2012		U-747	
<u>APREPITANT - EMEND</u>					
021549 002	5145684	Jan 25, 2011	DP		
	5719147	Jun 29, 2012	DS DP		
	6096742	Jul 01, 2018	DS DP	U-745	
	6235735	Jun 29, 2012		U-747	
	6235735	Jun 29, 2012		U-746	

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

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APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>APREPITANT - EMEND</u>					
021549 003	5145684	Jan 25, 2011	DP	I-498	Jun 30, 2009
	5719147	Jun 29, 2012	DS DP	NS	Jun 30, 2009
	6096742	Jul 01, 2018	DS DP U-745	NCE	Mar 26, 2008
	6235735	Jun 29, 2012	U-747		
	6235735	Jun 29, 2012	U-746		
<u>ARFORMOTEROL TARTRATE - BROVANA</u>					
021912 001				NP	Oct 06, 2009
<u>ARIPIRAZOLE - ABILIFY</u>					
021436 001	5006528	Oct 20, 2014	DS DP U-761	I-488	Mar 01, 2008
<u>ARIPIRAZOLE - ABILIFY</u>					
021436 002	5006528	Oct 20, 2014	DS DP U-761	I-488	Mar 01, 2008
<u>ARIPIRAZOLE - ABILIFY</u>					
021436 003	5006528	Oct 20, 2014	DS DP U-761	I-488	Mar 01, 2008
<u>ARIPIRAZOLE - ABILIFY</u>					
021436 004	5006528	Oct 20, 2014	DS DP U-761	I-488	Mar 01, 2008
<u>ARIPIRAZOLE - ABILIFY</u>					
021436 005	5006528	Oct 20, 2014	DS DP U-761	I-488	Mar 01, 2008
<u>ARIPIRAZOLE - ABILIFY</u>					
021436 006	5006528	Oct 20, 2014	DS DP U-761	I-488	Mar 01, 2008
<u>ARIPIRAZOLE - ABILIFY</u>					
021713 001	5006528	Oct 20, 2014	DS DP U-761	I-488	Mar 01, 2008
	6977257	Apr 24, 2022	DS DP		
<u>ARIPIRAZOLE - ABILIFY</u>					
021729 002	5006528	Oct 20, 2014	DS DP U-761	I-488 I-437 NCE	Mar 01, 2008 Sep 29, 2007 Nov 15, 2007
<u>ARIPIRAZOLE - ABILIFY</u>					
021729 003	5006528	Oct 20, 2014	DS DP U-761	I-488 I-437 NCE	Mar 01, 2008 Sep 29, 2007 Nov 15, 2007
<u>ARIPIRAZOLE - ABILIFY</u>					
021729 004	5006528	Oct 20, 2014	DS DP U-761	I-488 I-437 NCE	Mar 01, 2008 Sep 29, 2007 Nov 15, 2007
<u>ARIPIRAZOLE - ABILIFY</u>					
021729 005	5006528	Oct 20, 2014	DS DP U-761	I-488 I-437 NCE	Mar 01, 2008 Sep 29, 2007 Nov 15, 2007
<u>ARIPIRAZOLE - ABILIFY</u>					
021866 001	5006528	Oct 20, 2014	DS DP U-763	NCE	Nov 15, 2007
	7115587	Jul 21, 2024	DS DP U-764	NDF	Sep 20, 2009
<u>ARTICAINE HYDROCHLORIDE; EPINEPHRINE BITARTRATE - SEPTOCAINE</u>					
022010 001				NP	Mar 30, 2009
<u>ASCORBIC ACID; POLYETHYLENE GLYCOL 3350; POTASSIUM CHLORIDE; SODIUM ASCORBATE; SODIUM CHLORIDE; SODIUM SULFATE - MOVIPREP</u>					
021881 001				NP	Aug 02, 2009
<u>ATAZANAVIR SULFATE - REYATAZ</u>					
021567 001	5849911	Apr 09, 2017	DS DP U-167		
	6087383	Dec 21, 2018	DS DP		
<u>ATAZANAVIR SULFATE - REYATAZ</u>					
021567 002	5849911	Apr 09, 2017	DS DP U-167		
	6087383	Dec 21, 2018	DS DP		
<u>ATAZANAVIR SULFATE - REYATAZ</u>					
021567 003	5849911	Apr 09, 2017	DS DP U-167		
	6087383	Dec 21, 2018	DS DP		

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

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APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE		PATENT CODES			EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE	
ATAZANAVIR SULFATE - REYATAZ									
021567 004	5849911	Apr	09, 2017	DS	DP	U-167	D-89 NCE	Jul 06, 2007	
	6087383	Dec	21, 2018	DS	DP			Jun 20, 2008	
ATOMOXETINE HYDROCHLORIDE - STRATTERA									
021411 001	5658590	Nov	26, 2016			U-494			
	5658590*PED	May	26, 2017			U-494			
ATOMOXETINE HYDROCHLORIDE - STRATTERA									
021411 002	5658590	Nov	26, 2016			U-494			
	5658590*PED	May	26, 2017			U-494			
ATOMOXETINE HYDROCHLORIDE - STRATTERA									
021411 003	5658590	Nov	26, 2016			U-494			
	5658590*PED	May	26, 2017			U-494			
ATOMOXETINE HYDROCHLORIDE - STRATTERA									
021411 004	5658590	Nov	26, 2016			U-494			
	5658590*PED	May	26, 2017			U-494			
ATOMOXETINE HYDROCHLORIDE - STRATTERA									
021411 005	5658590	Nov	26, 2016			U-494			
	5658590*PED	May	26, 2017			U-494			
ATOMOXETINE HYDROCHLORIDE - STRATTERA									
021411 006	5658590	Nov	26, 2016			U-494			
	5658590*PED	May	26, 2017			U-494			
ATOMOXETINE HYDROCHLORIDE - STRATTERA									
021411 007	5658590	Nov	26, 2016			U-494			
	5658590*PED	May	26, 2017			U-494			
ATOMOXETINE HYDROCHLORIDE - STRATTERA									
021411 008	5658590	Nov	26, 2016			U-494			
	5658590*PED	May	26, 2017			U-494			
ATROPINE; PRALIDOXIME CHLORIDE - DUODOTE									
021983 001	5092843	Apr	12, 2010		DP				
AVOBENZONE; ECAMSULE; OCTOCRYLENE - ANTHELIOS SX									
021502 001	4585597	Jun	16, 2007	DS	DP	U-752	NC	Jul 21, 2009	
	5587150	Dec	24, 2013		DP	U-752			
AZELASTINE HYDROCHLORIDE - ASTELIN									
020114 001							D-102	Feb 17, 2009	
BALSALAZIDE DISODIUM - COLAZAL									
020610 001	4412992*PED	Jan	08, 2007				>A> NPP >A> PED	Dec 20, 2009 Jun 20, 2010	
BETAMETHASONE DIPROPIONATE; CALCIPOTRIENE HYDRATE - TACLONEX									
021852 001	4866048	Dec	29, 2007	DS	DP	U-88	NC	Jan 09, 2009	
	4866048	Dec	29, 2007	DS	DP	U-193			
	5763426	Jun	09, 2015	DS	DP				
	6753013	Jan	27, 2020		DP	U-88			
	6753013	Jan	27, 2020		DP	U-193			
BETAMETHASONE VALERATE - LUXIQ									
020934 001	7078058	Mar	01, 2016	DS	DP				
BETAXOLOL HYDROCHLORIDE - BETOPTIC S									
019845 001	4911920	Mar	27, 2007						
	4911920*PED	Sep	27, 2007						
BIMATOPROST - LUMIGAN									
021275 001	5688819	Aug	19, 2014			U-446			
	6403649	Sep	21, 2012	DS	DP	U-446			
BIVALIRUDIN - ANGIOMAX									
020873 001							I-486	Nov 30, 2008	
BORTEZOMIB - VELCADE									
021602 001							>A> I-521 ODE	Dec 08, 2009 Mar 25, 2012	

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

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APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
BRIMONIDINE TARTRATE - ALPHAGAN P					
021770 001	5424078	Jun 13, 2012	DP		
	5424078*PED	Dec 13, 2012			
	6562873	Jul 10, 2021	DP		
	6562873*PED	Jan 10, 2022			
	6627210	Jul 18, 2021	DP		
	6627210*PED	Jan 18, 2022			
	6641834	Jul 28, 2021	DP		
	6641834*PED	Jan 28, 2022			
	6673337	Jul 26, 2021	DP		
6673337*PED	Jan 26, 2022				
BRINZOLAMIDE - AZOPT					
020816 001	5240923	Aug 31, 2010	DS DP	U-224	M-54 PED
	5240923*PED	Mar 01, 2011			
	5378703	Apr 01, 2012	DS DP	U-224	
	5378703*PED	Oct 01, 2012			
	5461081	Oct 24, 2012	DP	U-225	
	5461081*PED	Apr 24, 2013			
BROMFENAC SODIUM - XIBROM					
021664 001				I-485	Jan 27, 2009
BUDESONIDE - BUDESONIDE					
021949 001	>A> 4907583	Mar 13, 2007	DP		NP
	>A> 6027714	Jan 09, 2018	DP	U-787	
	>A> 6142145	May 08, 2018	DP		
	>A> 6287540	Jan 09, 2018	DP		
	>A> 7143764	Mar 13, 2018	DP		
BUDESONIDE - BUDESONIDE					
021949 002	>A> 4907583	Mar 13, 2007	DP		NP
	>A> 6027714	Jan 09, 2018	DP	U-787	
	>A> 6142145	May 08, 2018	DP		
	>A> 6287540	Jan 09, 2018	DP		
	>A> 7143764	Mar 13, 2018	DP		
BUDESONIDE - PULMICORT RESPULES					
020929 001	6598603*PED	Jun 23, 2019			
	6899099	Dec 23, 2018		U-751	
	6899099*PED	Jun 23, 2019			
BUDESONIDE - PULMICORT RESPULES					
020929 002	6899099	Dec 23, 2018		U-751	
	6899099*PED	Jun 23, 2019			
BUDESONIDE - RHINOCORT					
020746 001	6986904	Apr 29, 2017	DP	U-699	
BUDESONIDE - RHINOCORT					
020746 002	6986904	Apr 29, 2017	DP	U-699	
BUDESONIDE; FORMOTEROL FUMARATE - SYMBICORT					
021929 001	5674860	Oct 07, 2014	DP	U-754	NC
	5972919	Dec 17, 2012	DP	U-754	
	6123924	Sep 26, 2017	DP		
	6641800	Sep 26, 2017	DP		
BUDESONIDE; FORMOTEROL FUMARATE - SYMBICORT					
021929 002	5674860	Oct 07, 2014	DP	U-754	NC
	5972919	Dec 17, 2012	DP	U-754	
	6123924	Sep 26, 2017	DP		
	6641800	Sep 26, 2017	DP		
BUPROPION HYDROCHLORIDE - WELLBUTRIN XL					
021515 001				I-497	Jun 12, 2009
BUPROPION HYDROCHLORIDE - WELLBUTRIN XL					
021515 002				I-497	Jun 12, 2009

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnotes for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>BUSULFAN - BUSULFEX</u>					
020954 001	5559148	Sep 30, 2013	U-264		
	5559148*PED	Mar 30, 2014	U-264		
<u>CALCIUM CARBONATE; RISEDRONATE SODIUM - ACTONEL WITH CALCIUM (COPACKAGED)</u>					
021823 001				M-52	Jan 24, 2009
<u>CARBIDOPA; ENTACAPONE; LEVODOPA - STALEVO 100</u>					
021485 002	4963590	Nov 27, 2007	DP	U-219	
	5112861	May 12, 2009		U-219	
	5135950	Oct 31, 2010	DS DP	U-219	
	6500867	Jun 29, 2020	DP	U-219	
<u>CARBIDOPA; ENTACAPONE; LEVODOPA - STALEVO 150</u>					
021485 003	4963590	Nov 27, 2007	DP	U-219	
	5112861	May 12, 2009		U-219	
	5135950	Oct 31, 2010	DS DP	U-219	
	6500867	Jun 29, 2020	DP	U-219	
<u>CARBIDOPA; ENTACAPONE; LEVODOPA - STALEVO 50</u>					
021485 001	4963590	Nov 27, 2007	DP	U-219	
	5112861	May 12, 2009		U-219	
	5135950	Oct 31, 2010	DS DP	U-219	
	6500867	Jun 29, 2020	DP	U-219	
<u>CARVEDILOL - COREG</u>					
020297 001	4503067	Mar 05, 2007	U-3	M-56	Aug 28, 2009
	4503067*PED	Sep 05, 2007		PED	Feb 28, 2010
	5760069	Jun 07, 2015	U-233		
	5760069*PED	Dec 07, 2015			
	5902821	Feb 07, 2016	U-313		
	5902821*PED	Aug 07, 2016			
<u>CARVEDILOL - COREG</u>					
020297 002	4503067	Mar 05, 2007	U-3	M-56	Aug 28, 2009
	4503067*PED	Sep 05, 2007		PED	Feb 28, 2010
	5760069	Jun 07, 2015	U-233		
	5760069*PED	Dec 07, 2015			
	5902821	Feb 07, 2016	U-313		
	5902821*PED	Aug 07, 2016			
<u>CARVEDILOL - COREG</u>					
020297 003	4503067	Mar 05, 2007	U-3	M-56	Aug 28, 2009
	4503067*PED	Sep 05, 2007		PED	Feb 28, 2010
	5760069	Jun 07, 2015	U-233		
	5760069*PED	Dec 07, 2015			
	5902821	Feb 07, 2016	U-313		
	5902821*PED	Aug 07, 2016			
<u>CARVEDILOL - COREG</u>					
020297 004	4503067	Mar 05, 2007	U-3	M-56	Aug 28, 2009
	4503067*PED	Sep 05, 2007		PED	Feb 28, 2010
	5760069	Jun 07, 2015	U-233		
	5760069*PED	Dec 07, 2015			
	5902821	Feb 07, 2016	U-313		
	5902821*PED	Aug 07, 2016			

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnotes for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>CARVEDILOL PHOSPHATE - COREG CR</u>					
022012 001	>A> 4503067	Mar 05, 2007	DS DP U-3	>A> M-56	Aug 28, 2009
	>A> 4503067*PED	Sep 05, 2007		>A> NDF	Oct 20, 2009
	>A> 5760069	Jun 07, 2015		>A> PED	Feb 28, 2010
	>A> 5760069*PED	Dec 07, 2015		>A> PED	Apr 20, 2010
	>A> 5902821	Feb 07, 2016			
	>A> 5902821	Feb 07, 2016		U-313	
	>A> 5902821*PED	Aug 07, 2016		U-777	
	>A> 6022562	Oct 17, 2015	DP		
	>A> 6022562*PED	Apr 17, 2016			
<u>CARVEDILOL PHOSPHATE - COREG CR</u>					
022012 002	>A> 4503067	Mar 05, 2007	DS DP U-3	>A> M-56	Aug 28, 2009
	>A> 4503067*PED	Sep 05, 2007		>A> NDF	Oct 20, 2009
	>A> 5760069	Jun 07, 2015		>A> PED	Feb 28, 2010
	>A> 5760069*PED	Dec 07, 2015		>A> PED	Apr 20, 2010
	>A> 5902821	Feb 07, 2016			
	>A> 5902821	Feb 07, 2016		U-777	
	>A> 5902821*PED	Aug 07, 2016		U-313	
	>A> 6022562	Oct 17, 2015	DP		
	>A> 6022562*PED	Apr 17, 2016			
<u>CARVEDILOL PHOSPHATE - COREG CR</u>					
022012 003	>A> 4503067	Mar 05, 2007	DS DP U-3	>A> M-56	Aug 28, 2009
	>A> 4503067*PED	Sep 05, 2007		>A> NDF	Oct 20, 2009
	>A> 5760069	Jun 07, 2015		>A> PED	Feb 28, 2010
	>A> 5760069*PED	Dec 07, 2015		>A> PED	Apr 20, 2010
	>A> 5902821	Feb 07, 2016			
	>A> 5902821	Feb 07, 2016		U-777	
	>A> 5902821*PED	Aug 07, 2016		U-313	
	>A> 6022562	Oct 17, 2015	DP		
	>A> 6022562*PED	Apr 17, 2016			
<u>CARVEDILOL PHOSPHATE - COREG CR</u>					
022012 004	>A> 4503067	Mar 05, 2007	DS DP U-3	>A> M-56	Aug 28, 2009
	>A> 4503067*PED	Sep 05, 2007		>A> NDF	Oct 20, 2009
	>A> 5760069	Jun 07, 2015		>A> PED	Feb 28, 2010
	>A> 5760069*PED	Dec 07, 2015		>A> PED	Apr 20, 2010
	>A> 5902821	Feb 07, 2016			
	>A> 5902821	Feb 07, 2016		U-777	
	>A> 5902821*PED	Aug 07, 2016		U-313	
	>A> 6022562	Oct 17, 2015	DP		
	>A> 6022562*PED	Apr 17, 2016			
<u>CASPOFUNGIN ACETATE - CANCIDAS</u>					
021227 001	5514650	Jan 26, 2015	DP U-607		
<u>CASPOFUNGIN ACETATE - CANCIDAS</u>					
021227 002	5514650	Jan 26, 2015	DP U-607		
<u>CEFDITOREN PIVOXIL - SPECTRACEF</u>					
021222 001	4839350	Apr 14, 2009	DS DP		
<u>CELECOXIB - CELEBREX</u>					
020998 001	5466823	Nov 30, 2013	DS	I-466	Jul 29, 2008
	5466823*PED	May 30, 2014		>A> NPP	Dec 15, 2009
	5563165	Nov 30, 2013	DP	>A> PED	Jun 15, 2010
	5563165*PED	May 30, 2014		PED	Jan 29, 2009
	5760068	Jun 02, 2015			
	5760068	Jun 02, 2015		U-672	
	5760068*PED	Dec 02, 2015		U-299	
	5972986	Oct 14, 2017			
	5972986*PED	Apr 14, 2018		U-299	

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnotes for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>CELECOXIB - CELEBREX</u>					
020998 002	5466823	Nov 30, 2013	DS	I-466	Jul 29, 2008
	5466823*PED	May 30, 2014		>A> NPP	Dec 15, 2009
	5563165	Nov 30, 2013	DP	>A> PED	Jun 15, 2010
	5563165*PED	May 30, 2014		PED	Jan 29, 2009
	5760068	Jun 02, 2015		U-672	
	5760068	Jun 02, 2015		U-299	
	5760068*PED	Dec 02, 2015			
	5972986	Oct 14, 2017		U-299	
	5972986*PED	Apr 14, 2018			
<u>CELECOXIB - CELEBREX</u>					
020998 003	5466823	Nov 30, 2013	DS	I-466	Jul 29, 2008
	5466823*PED	May 30, 2014		>A> NPP	Dec 15, 2009
	5563165	Nov 30, 2013	DP	>A> PED	Jun 15, 2010
	5563165*PED	May 30, 2014		PED	Jan 29, 2009
	5760068	Jun 02, 2015		U-672	
	5760068	Jun 02, 2015		U-299	
	5760068*PED	Dec 02, 2015			
	5972986	Oct 14, 2017		U-299	
	5972986*PED	Apr 14, 2018			
<u>CETIRIZINE HYDROCHLORIDE - ZYRTEC</u>					
019835 001	4525358	Jun 25, 2007	DS DP	U-565	
	4525358*PED	Dec 25, 2007			
<u>CETIRIZINE HYDROCHLORIDE - ZYRTEC</u>					
019835 002	4525358	Jun 25, 2007	DS DP	U-565	
	4525358*PED	Dec 25, 2007			
<u>CETIRIZINE HYDROCHLORIDE; PSEUDOEPHEDRINE HYDROCHLORIDE - ZYRTEC-D 12 HOUR</u>					
021150 001	7014867	Jun 10, 2022	DP		
<u>CHLORHEXIDINE GLUCONATE - CHLORHEXIDINE GLUCONATE</u>					
021669 001	7066916	Feb 17, 2024		U-737	
<u>CHLORHEXIDINE GLUCONATE; ISOPROPYL ALCOHOL - CHLORAPREP ONE-STEP</u>					
020832 004	5690958	Sep 30, 2016	DP		
	6536975	Nov 10, 2020	DP		
<u>CHLORHEXIDINE GLUCONATE; ISOPROPYL ALCOHOL - CHLORAPREP ONE-STEP FREPP</u>					
020832 003	5538353	Aug 25, 2015	DP		
	5690958	Sep 30, 2016	DP		
	5752363	Apr 22, 2017	DP		
	5772346	Apr 22, 2017	DP		
	D386849	Nov 25, 2011	DP		
	D396911	Aug 11, 2012	DP		
<u>CHLORHEXIDINE GLUCONATE; ISOPROPYL ALCOHOL - CHLORAPREP SINGLE SWABSTICK</u>					
021555 002	5690958	Sep 30, 2016	DP		
<u>CHLORHEXIDINE GLUCONATE; ISOPROPYL ALCOHOL - CHLORAPREP WITH TINT</u>					
020832 002	6991393	Mar 14, 2023	DP		
<u>CHLORHEXIDINE GLUCONATE; ISOPROPYL ALCOHOL - CHLORAPREP WITH TINT</u>					
020832 005	5690958	Sep 30, 2016	DP		
	6536975	Nov 10, 2020	DP		
	6729786	Mar 14, 2023	DP		
	6991393	Jan 31, 2024	DP		
<u>CICLESONIDE - OMNARIS</u>					
022004 001				NCE	Oct 20, 2011
<u>CICLOPIROX - LOPROX</u>					
020519 001	7018656	Sep 05, 2018	DP		
	7026337	Apr 02, 2018		U-714	
<u>CIPROFLOXACIN - CIPRO</u>					
019847 002				I-421 PED	Mar 25, 2007 Sep 25, 2007

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnotes for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>CIPROFLOXACIN - CIPRO</u>					
019847 003				I-421 PED	Mar 25, 2007 Sep 25, 2007
<u>CLOBETASOL PROPIONATE - CLOBEX</u>					
021835 001	5972920	Feb 12, 2018	DP	NDF	Oct 27, 2008
	5990100	Mar 24, 2018	DP	U-742	
<u>CLOPIDOGREL BISULFATE - CLOPIDOGREL BISULFATE</u>					
076274 001				PC	Feb 04, 2007
<u>CLOPIDOGREL BISULFATE - PLAVIX</u>					
020839 001				I-502	Aug 17, 2009
<u>COLESEVELAM HYDROCHLORIDE - WELCHOL</u>					
021176 001	5693675	Dec 02, 2014	DS		
	5917007	Apr 29, 2014	DS	U-323	
	5919832	Jun 10, 2014	DS		
	6066678	Jun 10, 2014	DS	U-323	
	6433026	Jun 10, 2014	DS		
	6784254	Jun 10, 2014	DS	DP	
	7101960	Apr 29, 2014	DS	DP	U-757
<u>CONIVAPTAN HYDROCHLORIDE - VAPRISOL</u>					
021697 001	5723606	Mar 03, 2015	DS	DP	U-698
<u>DAPSONE - ACZONE</u>					
021794 001	5863560	Sep 11, 2016	DP		
	6620435	Sep 11, 2016	DP		
<u>DAPTOMYCIN - CUBICIN</u>					
021572 001	6468967	Sep 24, 2019		U-282	
	RE39071	Jun 15, 2016	DS	DP	U-728
<u>DAPTOMYCIN - CUBICIN</u>					
021572 002	6468967	Sep 24, 2019		U-282	I-505
	RE39071	Jun 15, 2016	DS	DP	U-728
<u>DARUNAVIR ETHANOLATE - PREZISTA</u>					
021976 001	5843946	Dec 01, 2015		DP	U-744
	6248775	Aug 25, 2012	DS		
	6335460	Aug 25, 2012	DS	DP	U-744
<u>DASATINIB - SPRYCEL</u>					
021986 001	6596746	Apr 13, 2020	DS	DP	U-748
	7125875	Apr 13, 2020		U-780	NCE
	7125875	Apr 13, 2020		U-779	ODE
<u>DASATINIB - SPRYCEL</u>					
021986 002	6596746	Apr 13, 2020	DS	DP	U-748
	7125875	Apr 13, 2020		U-780	NCE
	7125875	Apr 13, 2020		U-779	ODE
<u>DASATINIB - SPRYCEL</u>					
021986 003	6596746	Apr 13, 2020	DS	DP	U-748
	7125875	Apr 13, 2020		U-780	NCE
	7125875	Apr 13, 2020		U-779	ODE
<u>DECITABINE - DACOGEN</u>					
021790 001				NCE	May 02, 2011
<u>DEFERASIROX - EXJADE</u>					
021882 001	6465504	Jun 24, 2017	DS	DP	
	6596750	Jun 24, 2017	DS		U-735
<u>DEFERASIROX - EXJADE</u>					
021882 002	6465504	Jun 24, 2017	DS	DP	
	6596750	Jun 24, 2017	DS		U-735
<u>DEFERASIROX - EXJADE</u>					
021882 003	6465504	Jun 24, 2017	DS	DP	
	6596750	Jun 24, 2017	DS		U-735

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnotes for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>DESFLURANE - SUPRANE</u>					
020118 001	4762856	Feb 02, 2007	U-67		
	4762856*PED	Aug 02, 2007			
	5617906	Apr 08, 2014	DP		
	5617906*PED	Oct 08, 2014			
<u>DESIRUDIN RECOMBINANT - IPRIVASK</u>					
021271 001	4745177	May 17, 2010	DS DP		
<u>DESLORATADINE - CLARINEX</u>					
021165 001	4659716	Mar 31, 2007	DP	U-725	
	4659716	Mar 31, 2007	DP	U-427	
	4659716*PED	Oct 01, 2007		U-427	
<u>DESLORATADINE - CLARINEX</u>					
021300 001	4659716	Mar 31, 2007	DP	U-725	
	4659716	Mar 31, 2007	DP	U-611	
	4659716*PED	Oct 01, 2007			
<u>DESLORATADINE - CLARINEX</u>					
021312 001	4659716	Mar 31, 2007	DP	U-725	
	4659716	Mar 31, 2007	DP	U-427	
	4659716*PED	Oct 01, 2007		U-427	
	5178878	Jan 12, 2010	DP		
	5607697	Jun 07, 2015	DP		
<u>DESLORATADINE - CLARINEX</u>					
021312 002	4659716	Mar 31, 2007	DP	U-725	
	4659716	Mar 31, 2007	DP	U-427	
	4659716*PED	Oct 01, 2007	DP		
	5178878	Jan 12, 2010	DP		
	5607697	Jun 07, 2015	DP		
<u>DESLORATADINE; PSEUDOEPHEDRINE SULFATE - CLARINEX D 24 HOUR</u>					
021605 001	4659716	Mar 31, 2007	DP	U-726	
	4659716	Mar 31, 2007	DP	U-644	
	4659716*PED	Oct 01, 2007			
	6979463	Mar 28, 2022	DP		
<u>DESLORATADINE; PSEUDOEPHEDRINE SULFATE - CLARINEX-D 12 HOUR</u>					
021313 001	4659716	Mar 31, 2007	DP	U-707	
	4659716*PED	Oct 01, 2007			
	6100274	Jul 07, 2019	DP		
	6100274*PED	Jan 07, 2020			
	6709676	Feb 18, 2021	DP	U-707	
<u>DESMOPRESSIN ACETATE - DDAVP</u>					
019955 001	7022340	Apr 30, 2023	DP		
<u>DESMOPRESSIN ACETATE - DDAVP</u>					
019955 002	7022340	Apr 30, 2023	DP		
<u>DESONIDE - DESONATE</u>					
021844 001	6387383	Aug 03, 2020	DS DP	U-783	
<u>DESONIDE - VERDESO</u>					
021978 001	6730288	Sep 08, 2019	DP		
	7029659	Sep 08, 2019	DP		
<u>DEXMEDETOMIDINE - PRECEDEX</u>					
021038 001	4910214	Jul 15, 2013	DS DP	U-421	
<u>DEXMETHYLPHENIDATE HYDROCHLORIDE - FOCALIN XR</u>					
021802 004					
				NDF	May 26, 2008
<u>DIAZEPAM - DIASTAT</u>					
020648 001	5462740	Sep 17, 2013	DP		
<u>DIAZEPAM - DIASTAT</u>					
020648 002	5462740	Sep 17, 2013	DP		

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnotes for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>DIAZEPAM - DIASTAT</u>					
020648 003	5462740	Sep 17, 2013	DP		
<u>DIAZEPAM - DIASTAT</u>					
020648 004	5462740	Sep 17, 2013	DP		
<u>DIAZEPAM - DIASTAT</u>					
020648 005	5462740	Sep 17, 2013	DP		
<u>DIAZEPAM - DIASTAT ACUDIAL</u>					
020648 006	5462740	Sep 17, 2013	DP		
<u>DIAZEPAM - DIASTAT ACUDIAL</u>					
020648 007	5462740	Sep 17, 2013	DP		
<u>DILTIAZEM HYDROCHLORIDE - CARDIZEM LA</u>					
021392 001	7108866	Dec 17, 2019	DP U-107		
<u>DILTIAZEM HYDROCHLORIDE - CARDIZEM LA</u>					
021392 002	7108866	Dec 17, 2019	DP U-107		
<u>DILTIAZEM HYDROCHLORIDE - CARDIZEM LA</u>					
021392 003	7108866	Dec 17, 2019	DP U-107		
<u>DILTIAZEM HYDROCHLORIDE - CARDIZEM LA</u>					
021392 004	7108866	Dec 17, 2019	DP U-107		
<u>DILTIAZEM HYDROCHLORIDE - CARDIZEM LA</u>					
021392 005	7108866	Dec 17, 2019	DP U-107		
<u>DILTIAZEM HYDROCHLORIDE - CARDIZEM LA</u>					
021392 006	7108866	Dec 17, 2019	DP U-107		
<u>DOCETAXEL - TAXOTERE</u>					
020449 001	4814470	May 14, 2010	DS DP	I-519	Oct 17, 2009
	5438072	Nov 22, 2013	DP	I-490	Mar 22, 2009
	5698582	Jul 03, 2012	DP		
	5714512	Jul 03, 2012	DP		
	5750561	Jul 03, 2012	DP		
<u>DONEPEZIL HYDROCHLORIDE - ARICEPT ODT</u>					
021720 001	4895841	Nov 25, 2010	DS DP U-713		
<u>DONEPEZIL HYDROCHLORIDE - ARICEPT ODT</u>					
021720 002	4895841	Nov 25, 2010	DS DP U-713		
<u>DROSPIRENONE; ETHINYL ESTRADIOL - YAZ</u>					
021676 001	5569652	Oct 29, 2013	U-758	I-508	Oct 04, 2009
	5569652	Oct 29, 2013	U-1	NP	Mar 16, 2009
	5798338	Jul 10, 2015	DP		
	6787531	Aug 31, 2020	DP		
	6933395	Aug 11, 2017	DP		
	6958326	Dec 20, 2021	DP		
	6987101	Dec 22, 2017	U-758		
	6987101	Dec 22, 2017	U-1		
	RE37564	Jun 30, 2014	DP		
	RE37838	Jun 30, 2014	DP		
	RE38253	Jun 30, 2014	DP		
<u>EFAVIRENZ - SUSTIVA</u>					
020972 001	5519021	May 21, 2013	DS DP		
	5663169	Sep 02, 2014	U-257		
	5811423	Aug 07, 2012	DS DP U-256		
	6238695	Apr 06, 2019	DP		
<u>EFAVIRENZ - SUSTIVA</u>					
020972 002	5519021	May 21, 2013	DS DP		
	5663169	Sep 02, 2014	U-257		
	5811423	Aug 07, 2012	DS DP U-256		
	6238695	Apr 06, 2019	DP		

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnotes for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE			PATENT CODES		EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE	
EFAVIRENZ - SUSTIVA									
020972 003	5519021	May	21, 2013	DS	DP				
	5663169	Sep	02, 2014			U-257			
	5811423	Aug	07, 2012	DS	DP	U-256			
	6238695	Apr	06, 2019		DP				
EFAVIRENZ - SUSTIVA									
021360 002	5519021	May	21, 2013	DS	DP				
EFAVIRENZ; EMTRICITABINE; TENOFOVIR DISOPROXIL FUMARATE - ATRIPLA									
021937 001	5210085	May	11, 2010			U-750			
	5210085*PED	Nov	11, 2010						
	5519021	May	21, 2013	DS	DP				
	5663169	Sep	02, 2014			U-750			
	5811423	Aug	07, 2012			U-750			
	5814639	Sep	29, 2015	DS	DP				
	5814639*PED	Mar	29, 2016						
	5914331	Sep	29, 2015	DS					
	5914331*PED	Mar	29, 2016						
	5922695	Jul	25, 2017	DS		U-750			
	5935946	Jul	25, 2017	DS	DP	U-750			
	5977089	Jul	25, 2017	DS	DP	U-750			
	6043230	Jul	25, 2017			U-750			
	6238695	Apr	06, 2019		DP				
	6555133	Apr	06, 2019			U-750			
	6639071	Nov	13, 2021	DS					
	6642245	Nov	04, 2020			U-750			
	6642245*PED	May	04, 2021						
	6703396	Mar	09, 2021	DS	DP				
	6703396*PED	Sep	09, 2021						
	6939964	Jan	20, 2018	DS					
EMTRICITABINE - EMTRIVA									
021500 001	5210085	May	11, 2010				NCE	Jul	02, 2008
	5210085*PED	Nov	11, 2010				PED	Jan	02, 2009
	5814639	Sep	29, 2015						
	5814639*PED	Mar	29, 2016						
	5914331	Sep	29, 2015						
	5914331*PED	Mar	29, 2016						
	6642245	Nov	04, 2020			U-541			
	6642245*PED	May	04, 2021						
	6703396	Mar	09, 2021	DS	DP				
	6703396*PED	Sep	09, 2021						
EMTRICITABINE - EMTRIVA									
021896 001	5210085	May	11, 2010			U-257	>A> NPP NDF NCE	Dec	22, 2009
	5210085*PED	Nov	11, 2010					Sep	27, 2008
	5814639	Sep	29, 2015	DS	DP			Jul	02, 2008
	5814639*PED	Mar	29, 2016				>A> PED	Jun	22, 2010
	5914331	Sep	29, 2015	DS			PED	Mar	27, 2009
	5914331*PED	Mar	29, 2016				PED	Jan	02, 2009
	6642245	Nov	04, 2020			U-257			
	6642245*PED	May	04, 2021						
	6703396	Mar	09, 2021	DS	DP				
	6703396*PED	Sep	09, 2021						

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnotes for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE		PATENT CODES		EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE	
EMTRICITABINE; TENOFOVIR DISOPROXIL FUMARATE - TRUVADA								
021752 001	5210085	May	11, 2010		U-248	NCE PED	Jul	02, 2008
	5210085	May	11, 2010		U-541		Jan	02, 2009
	5210085*PED	Nov	11, 2010					
	5814639	Sep	29, 2015	DS	DP			
	5814639*PED	Mar	29, 2016					
	5914331	Sep	29, 2015	DS	DP	U-248		
	5914331*PED	Mar	29, 2016					
	6642245	Nov	04, 2020			U-248		
	6642245	Nov	04, 2020			U-541		
	6642245*PED	May	04, 2021					
	6703396	Mar	09, 2021	DS	DP			
	6703396*PED	Sep	09, 2021					
ENFUVIRTIDE - FUZEON								
021481 001						M-59	Sep	29, 2009
ENTACAPONE - COMTAN								
020796 001	4963590	Nov	27, 2007		DP	U-219		
	5112861	May	12, 2009			U-219		
	5135950	Oct	31, 2010	DS	DP	U-219		
	6599530	Sep	14, 2018		DP	U-219		
EPLERENONE - INSPRA								
021437 001	4559332	Apr	09, 2007	DS	DP	U-537		
EPLERENONE - INSPRA								
021437 002	4559332	Apr	09, 2007	DS	DP	U-537		
EPLERENONE - INSPRA								
021437 003	4559332	Apr	09, 2007	DS	DP	U-537		
ERTAPENEM SODIUM - INVANZ								
021337 001						I-515	Aug	10, 2009
ESCITALOPRAM OXALATE - LEXAPRO								
021323 001	RE34712	Sep	14, 2011	DS	DP			
	RE34712*PED	Mar	14, 2012					
ESCITALOPRAM OXALATE - LEXAPRO								
021323 002	RE34712	Sep	14, 2011	DS	DP			
	RE34712*PED	Mar	14, 2012					
ESCITALOPRAM OXALATE - LEXAPRO								
021323 003	RE34712	Sep	14, 2011	DS	DP			
	RE34712*PED	Mar	14, 2012					
ESCITALOPRAM OXALATE - LEXAPRO								
021365 001	RE34712	Sep	14, 2011	DS	DP			
	RE34712*PED	Mar	14, 2012					

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnotes for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE		PATENT CODES		EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE	
ESOMEPRAZOLE MAGNESIUM - NEXIUM								
021153 001	4738974	Apr	19, 2007	DS	DP	U-770	I-504	Oct 11, 2009
	4738974	Apr	19, 2007	DS	DP	U-635	I-484	Nov 24, 2007
	4738974	Apr	19, 2007	DS	DP	U-373	NPP	Apr 28, 2009
	4738974*PED	Oct	19, 2007			U-373		
	4786505	Apr	20, 2007		DP	U-373		
	4786505	Apr	20, 2007		DP	U-729		
	4786505	Apr	20, 2007		DP	U-770		
	4853230	Apr	20, 2007		DP	U-373		
	4853230	Apr	20, 2007		DP	U-770		
	4853230	Apr	20, 2007		DP	U-729		
	5690960	Nov	25, 2014		DP	U-770		
	5690960	Nov	25, 2014		DP	U-729		
	5690960	Nov	25, 2014		DP	U-373		
	5714504	Feb	03, 2015		DP	U-729		
	5714504	Feb	03, 2015		DP	U-770		
	5714504	Feb	03, 2015		DP	U-373		
	5877192	May	27, 2014		DP	U-373		
	5877192	May	27, 2014		DP	U-770		
	5877192	May	27, 2014		DP	U-729		
	5900424	May	04, 2016	DS		U-770		
	5900424	May	04, 2016	DS		U-729		
	5900424	May	04, 2016	DS		U-373		
	6369085	May	25, 2018	DS	DP	U-729		
	6369085	May	25, 2018	DS	DP	U-770		
	6428810	Nov	03, 2019		DP	U-469		
	6428810	Nov	03, 2019		DP	U-770		
	6428810	Nov	03, 2019		DP	U-729		
ESOMEPRAZOLE MAGNESIUM - NEXIUM								
021153 002	4738974	Apr	19, 2007	DS	DP	U-635	I-504	Oct 11, 2009
	4738974	Apr	19, 2007	DS	DP	U-770	I-484	Nov 24, 2007
	4738974	Apr	19, 2007	DS	DP	U-373	NPP	Apr 28, 2009
	4738974*PED	Oct	19, 2007			U-373		
	4786505	Apr	20, 2007		DP	U-373		
	4786505	Apr	20, 2007		DP	U-770		
	4786505	Apr	20, 2007		DP	U-729		
	4853230	Apr	20, 2007		DP	U-373		
	4853230	Apr	20, 2007		DP	U-729		
	4853230	Apr	20, 2007		DP	U-770		
	5690960	Nov	25, 2014		DP	U-770		
	5690960	Nov	25, 2014		DP	U-729		
	5690960	Nov	25, 2014		DP	U-373		
	5714504	Feb	03, 2015		DP	U-729		
	5714504	Feb	03, 2015		DP	U-770		
	5714504	Feb	03, 2015		DP	U-373		
	5877192	May	27, 2014		DP	U-373		
	5877192	May	27, 2014		DP	U-770		
	5877192	May	27, 2014		DP	U-729		
	5900424	May	04, 2016	DS		U-729		
	5900424	May	04, 2016	DS		U-770		
	5900424	May	04, 2016	DS		U-373		
	6369085	May	25, 2018	DS	DP	U-729		
	6369085	May	25, 2018	DS	DP	U-770		
	6428810	Nov	03, 2019		DP	U-469		
	6428810	Nov	03, 2019		DP	U-770		
	6428810	Nov	03, 2019		DP	U-729		

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnotes for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>ESOMEPRAZOLE MAGNESIUM - NEXIUM</u>					
021957 001	4738974	Apr 19, 2007	DS DP	U-773	I-504
	4738974	Apr 19, 2007	DS DP	U-729	I-484
	4783974*PED	Oct 19, 2007			NPP
	4786505	Apr 20, 2007		DP U-729	
	4786505	Apr 20, 2007		DP U-773	
	4786505*PED	Oct 20, 2007			
	4853230	Apr 20, 2007		DP U-729	
	4853230	Apr 20, 2007		DP U-773	
	4853230*PED	Oct 20, 2007			
	5690960	Nov 25, 2014		DP U-729	
	5690960	Nov 25, 2014		DP U-773	
	5690960*PED	May 25, 2015			
	5714504	Feb 03, 2015		DP U-773	
	5714504	Feb 03, 2015		DP U-729	
	5714504*PED	Aug 03, 2015			
	5877192	May 27, 2014		U-773	
	5877192	May 27, 2014		U-729	
	5877192*PED	Nov 27, 2014			
	5900424	May 04, 2016	DS	U-729	
	5900424	May 04, 2016	DS	U-773	
	5900424*PED	Nov 04, 2016			
	6369085	May 25, 2018	DS DP	U-729	
	6369085	May 25, 2018	DS DP	U-773	
	6369085*PED	Nov 25, 2018			
	6428810	Nov 03, 2019		DP U-729	
	6428810	Nov 03, 2019		DP U-773	
	6428810*PED	May 03, 2020			
	6875872	May 27, 2014	DS		
	6875872*PED	Nov 27, 2014			
<u>ESOMEPRAZOLE MAGNESIUM - NEXIUM</u>					
021957 002	4738974	Apr 19, 2007	DS DP	U-729	I-504
	4738974	Apr 19, 2007	DS DP	U-773	I-484
	4783974*PED	Oct 19, 2007			NPP
	4786505	Apr 20, 2007		DP U-773	
	4786505	Apr 20, 2007		DP U-729	
	4786505*PED	Oct 20, 2007			
	4853230	Apr 20, 2007		DP U-773	
	4853230	Apr 20, 2007		DP U-729	
	4853230*PED	Oct 20, 2007			
	5690960	Nov 25, 2014		DP U-773	
	5690960	Nov 25, 2014		DP U-729	
	5690960*PED	May 25, 2015			
	5714504	Feb 03, 2015		DP U-773	
	5714504	Feb 03, 2015		DP U-729	
	5714504*PED	Aug 03, 2015			
	5877192	May 27, 2014		U-729	
	5877192	May 27, 2014		U-773	
	5877192*PED	Nov 27, 2014			
	5900424	May 04, 2016	DS	U-773	
	5900424	May 04, 2016	DS	U-729	
	5900424*PED	Nov 04, 2016			
	6369085	May 25, 2018	DS DP	U-773	
	6369085	May 25, 2018	DS DP	U-729	
	6369085*PED	Nov 25, 2018			
	6428810	Nov 03, 2019		DP U-773	
	6428810	Nov 03, 2019		DP U-729	
	6428810*PED	May 03, 2020			
	6875872	May 27, 2014	DS		
	6875872*PED	Nov 27, 2014			

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnotes for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>ESTRADIOL - ELESTRIN</u> 021813 001				>A> NP	Dec 15, 2009
<u>ESTRADIOL - ESTRADIOL</u> 075182 004				PC	Feb 06, 2007
<u>ESTRADIOL - ESTRADIOL</u> 075182 005				PC	Feb 06, 2007
<u>ESTRADIOL; NORETHINDRONE ACETATE - ACTIVELLA</u> 020907 002				>A> D-104 >A> NS	Dec 29, 2009 Dec 28, 2009
<u>ETHINYL ESTRADIOL; LEVONORGESTREL - SEASONIQUE</u> 021840 001				NP	May 25, 2009
<u>ETHINYL ESTRADIOL; NORELGESTROMIN - ORTHO EVRA</u> 021180 001	5876746	Nov 20, 2015	DP U-514		
<u>ETHINYL ESTRADIOL; NORETHINDRONE - OVCON-35</u> 021490 001	6667050	Apr 06, 2019	DP U-1		
<u>ETHINYL ESTRADIOL; NORETHINDRONE ACETATE - LOESTRIN 24 FE</u> 021871 001	5552394	Jul 22, 2014	U-1	NP	Feb 17, 2009
<u>ETONOGESTREL - IMPLANON</u> 021529 001	4957119	Aug 05, 2008	DP	NDF	Jul 17, 2009
	5150718	Sep 29, 2009	U-749		
<u>EXEMESTANE - AROMASIN</u> 020753 001				I-495	Oct 05, 2008
<u>EXENATIDE SYNTHETIC - BYETTA</u> 021773 001				>A> I-520	Dec 22, 2009
<u>EXENATIDE SYNTHETIC - BYETTA</u> 021773 002				>A> I-520	Dec 22, 2009
<u>EZETIMIBE - ZETIA</u> 021445 001	7030106	Jan 25, 2022	DP	I-493	May 23, 2009
	RE37721	Oct 25, 2016	DS DP U-473	M-57	Mar 16, 2009
<u>EZETIMIBE; SIMVASTATIN - VYTORIN</u> 021687 001	RE37721	Oct 25, 2016	DS DP U-473	M-60 M-58	Oct 03, 2009 Mar 16, 2009
<u>EZETIMIBE; SIMVASTATIN - VYTORIN</u> 021687 002	RE37721	Oct 25, 2016	DS DP U-473	M-60 M-58	Oct 03, 2009 Mar 16, 2009
<u>EZETIMIBE; SIMVASTATIN - VYTORIN</u> 021687 003	RE37721	Oct 25, 2016	DS DP U-473	M-60 M-58	Oct 03, 2009 Mar 16, 2009
<u>EZETIMIBE; SIMVASTATIN - VYTORIN</u> 021687 004	RE37721	Oct 25, 2016	DS DP U-473	M-60 M-58	Oct 03, 2009 Mar 16, 2009
<u>FAMCICLOVIR - FAMVIR</u> 020363 001				D-103 I-501	Jul 28, 2009 Jul 28, 2009
<u>FAMCICLOVIR - FAMVIR</u> 020363 002				D-103 I-501	Jul 28, 2009 Jul 28, 2009
<u>FAMCICLOVIR - FAMVIR</u> 020363 003				D-103 I-501	Jul 28, 2009 Jul 28, 2009
<u>FENOFIBRATE - ANTARA (MICRONIZED)</u> 021695 001	7101574	Aug 20, 2020	DS DP	M-47	Oct 21, 2008
<u>FENOFIBRATE - ANTARA (MICRONIZED)</u> 021695 003	7101574	Aug 20, 2020	DS DP	M-47	Oct 21, 2008
<u>FENOFIBRATE - FENOFIBRATE</u> 076433 001				PC	May 22, 2006

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnotes for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>FENOFIBRATE - FENOFIBRATE</u>					
076433 002				PC	May 22, 2006
<u>FENOFIBRATE - LIPOFEN</u>					
021612 001	5545628	Jan 10, 2015	U-701		
<u>FENOFIBRATE - LIPOFEN</u>					
021612 002	5545628	Jan 10, 2015	U-701		
<u>FENOFIBRATE - LIPOFEN</u>					
021612 003	5545628	Jan 10, 2015	U-701		
<u>FENOFIBRATE - TRICOR</u>					
021203 001	>A> 7037529	Jan 09, 2018	DP		
	>A> 7041319	Jan 09, 2018	DP		
<u>FENOFIBRATE - TRICOR</u>					
021203 003	>A> 7037529	Jan 09, 2018	DP		
	>A> 7041319	Jan 09, 2018	DP		
<u>FENOFIBRATE - TRICOR</u>					
021656 001	7037529	Jan 09, 2018	DP		
	7041319	Jan 09, 2018	DP		
<u>FENOFIBRATE - TRICOR</u>					
021656 002	7037529	Jan 09, 2018	DP		
	7041319	Jan 09, 2018	DP		
<u>FENTANYL CITRATE - FENTORA</u>					
021947 001	6200604	Mar 26, 2019	U-767	NDF	Sep 25, 2009
	6974590	Mar 26, 2019	U-767		
<u>FENTANYL CITRATE - FENTORA</u>					
021947 002	6200604	Mar 26, 2019	U-767	NDF	Sep 25, 2009
	6974590	Mar 26, 2019	U-767		
<u>FENTANYL CITRATE - FENTORA</u>					
021947 003	6200604	Mar 26, 2019	U-767	NDF	Sep 25, 2009
	6974590	Mar 26, 2019	U-767		
<u>FENTANYL CITRATE - FENTORA</u>					
021947 004	6200604	Mar 26, 2019	U-767	NDF	Sep 25, 2009
	6974590	Mar 26, 2019	U-767		
<u>FENTANYL CITRATE - FENTORA</u>					
021947 005	6200604	Mar 26, 2019	U-767	NDF	Sep 25, 2009
	6974590	Mar 26, 2019	U-767		
<u>FENTANYL HYDROCHLORIDE - IONSYS</u>					
021338 001	5232438	Oct 03, 2008	DP	U-736	NDF
	5445606	Dec 11, 2011	DP		
	5697896	Dec 16, 2014	DP		
	5843014	Dec 01, 2015	DP		
	6169920	Jan 02, 2018	DP		
	6171294	Jun 05, 2015		U-736	
	6181963	Nov 02, 2019	DP		
	6195582	Jan 28, 2019	DP	U-736	
	6216033	Jun 05, 2015	DP		
	6317629	Jun 02, 2012	DP		
	6425892	Jun 05, 2015		U-736	
	6842640	Jun 02, 2015	DP		
	6881208	Apr 19, 2022		U-736	
	6975902	Apr 01, 2024	DP		
	7018370	Jun 05, 2015		U-736	
	7027859	Sep 26, 2014	DP		

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnotes for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>FEXOFENADINE HYDROCHLORIDE - ALLEGRA</u>					
020625 001	>A> 7135571	May 18, 2014	DS		
	>A> 7135571*PED	Nov 18, 2014			
	>A> 7138524	May 18, 2014	DS		
	>A> 7138524*PED	Nov 18, 2014			
<u>FEXOFENADINE HYDROCHLORIDE - ALLEGRA</u>					
020872 001	>A> 7135571	May 18, 2014	DS		
	>A> 7135571*PED	Nov 18, 2014			
	>A> 7138524	May 18, 2014	DS		
	>A> 7138524*PED	Nov 18, 2014			
<u>FEXOFENADINE HYDROCHLORIDE - ALLEGRA</u>					
020872 002	>A> 7135571	May 18, 2014	DS		
	>A> 7135571*PED	Nov 18, 2014			
	>A> 7138524	May 18, 2014	DS		
	>A> 7138524*PED	Nov 18, 2014			
<u>FEXOFENADINE HYDROCHLORIDE - ALLEGRA</u>					
020872 004	>A> 7135571	May 18, 2014	DS		
	>A> 7135571*PED	Nov 18, 2014			
	>A> 7138524	May 18, 2014	DS		
	>A> 7138524*PED	Nov 18, 2014			
<u>FEXOFENADINE HYDROCHLORIDE - ALLEGRA</u>					
021963 001	5578610	Nov 26, 2013	DS DP	U-772	
	6037353	Mar 14, 2017		U-772	
	6187791	May 11, 2012		U-772	
	6399632	May 11, 2012		U-772	
	>A> 7138524	May 18, 2014	DS		
	>A> 7138524*PED	Nov 18, 2014			
<u>FEXOFENADINE HYDROCHLORIDE; PSEUDOEPHEDRINE HYDROCHLORIDE - ALLEGRA D 24 HOUR</u>					
021704 001	>A> 7138524	May 18, 2014	DS		
	>A> 7138524*PED	Nov 18, 2014			
	RE39069	May 29, 2018	DP		
<u>FEXOFENADINE HYDROCHLORIDE; PSEUDOEPHEDRINE HYDROCHLORIDE - ALLEGRA-D 12 HOUR</u>					
020786 001	>A> 7135571	May 18, 2014	DS		
	>A> 7135571*PED	Nov 18, 2014			
	>A> 7138524	May 18, 2014	DS		
	>A> 7138524*PED	Nov 18, 2014			
<u>FINASTERIDE - FINASTERIDE</u>					
076340 001				PC	Dec 16, 2006
<u>FLUNISOLIDE - AEROSPAN HFA</u>					
021247 001				NP	Jan 27, 2009
<u>FLUOCINOLONE ACETONIDE - RETISERT</u>					
021737 001	6217895	Mar 22, 2019	DP	U-708	
	6548078	Mar 22, 2019	DP	U-708	
<u>FLUOCINONIDE - VANOS</u>					
021758 001				I-487	Mar 02, 2009
<u>FLUOXETINE HYDROCHLORIDE - PROZAC WEEKLY</u>					
021235 001	RE39030	May 29, 2017	DP	U-397	
	RE39030	May 29, 2017	DP	U-396	
<u>FLUTICASONE PROPIONATE - FLOVENT HFA</u>					
021433 001	5658549	Sep 19, 2014	DP	U-710	NPP
	5674472	Oct 07, 2014	DP		
	6251368	Dec 04, 2012	DP		
<u>FLUTICASONE PROPIONATE - FLOVENT HFA</u>					
021433 002	5658549	Sep 19, 2014	DP	U-710	NPP
	5674472	Oct 07, 2014	DP		
	6251368	Dec 04, 2012	DP		

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnotes for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
FLUTICASONE PROPIONATE - FLOVENT HFA					
021433 003	5658549	Sep 19, 2014	DP U-710	NPP	Feb 28, 2009
	5674472	Oct 07, 2014	DP		
	6251368	Dec 04, 2012	DP		
FLUTICASONE PROPIONATE; SALMETEROL XINAFOATE - ADVAIR DISKUS 100/50					
021077 001	4992474	Feb 12, 2008	U-211		
	4992474*PED	Aug 12, 2008	U-211		
	5126375	Feb 12, 2008			
	5126375*PED	Aug 12, 2008			
	5225445	Feb 12, 2008	U-211		
	5225445*PED	Aug 12, 2008	U-211		
	6536427	Mar 01, 2011	DP		
	6536427*PED	Sep 01, 2011			
FLUTICASONE PROPIONATE; SALMETEROL XINAFOATE - ADVAIR DISKUS 250/50					
021077 002	4992474	Feb 12, 2008	U-211		
	4992474*PED	Aug 12, 2008	U-211		
	5126375	Feb 12, 2008			
	5126375*PED	Aug 12, 2008			
	5225445	Feb 12, 2008	U-211		
	5225445*PED	Aug 12, 2008	U-211		
	6536427	Mar 01, 2011	DP		
	6536427*PED	Sep 01, 2011			
FLUTICASONE PROPIONATE; SALMETEROL XINAFOATE - ADVAIR DISKUS 500/50					
021077 003	4992474	Feb 12, 2008	U-211		
	4992474*PED	Aug 12, 2008	U-211		
	5126375	Feb 12, 2008			
	5126375*PED	Aug 12, 2008			
	5225445	Feb 12, 2008	U-211		
	5225445*PED	Aug 12, 2008	U-211		
	6536427	Mar 01, 2011	DP		
	6536427*PED	Sep 01, 2011			
FLUTICASONE PROPIONATE; SALMETEROL XINAFOATE - ADVAIR HFA					
021254 001	4992474	Feb 12, 2008	DS DP U-738	NP	Jun 08, 2009
	4992474*PED	Aug 12, 2008			
	5126375	Feb 12, 2008	DS DP		
	5126375*PED	Aug 12, 2008			
	5225445	Feb 12, 2008	U-738		
	5225445*PED	Aug 12, 2008			
	5270305	Sep 07, 2010	U-738		
	5658549	Aug 19, 2014	DP U-738		
	5674472	Oct 07, 2014	DP		
	6143277	Apr 14, 2015	DP U-738		
	6251368	Dec 04, 2012	DP		
	6253762	Apr 14, 2015	DP U-738		
	6315173	Dec 23, 2017	DP		
	6510969	Dec 23, 2017	DP		
	6524555	Apr 14, 2015	DP		
	6546928	Apr 14, 2015	DP		

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>FLUTICASONE PROPIONATE; SALMETEROL XINAFOATE - ADVAIR HFA</u>					
021254 002	4992474	Feb 12, 2008	DS DP	NP	Jun 08, 2009
	4992474*PED	Aug 12, 2008			
	5126375	Feb 12, 2008	DS DP		
	5126375*PED	Aug 12, 2008			
	5225445	Feb 12, 2008		U-738	
	5225445*PED	Aug 12, 2008			
	5270305	Sep 07, 2010		U-738	
	5658549	Aug 19, 2014	DP	U-738	
	5674472	Oct 07, 2014	DP		
	6143277	Apr 14, 2015	DP	U-738	
	6251368	Dec 04, 2012	DP		
	6253762	Apr 14, 2015	DP	U-738	
	6315173	Dec 23, 2017	DP		
	6510969	Dec 23, 2017	DP		
	6524555	Apr 14, 2015	DP		
	6546928	Apr 14, 2015	DP		
<u>FLUTICASONE PROPIONATE; SALMETEROL XINAFOATE - ADVAIR HFA</u>					
021254 003	4992474	Feb 12, 2008	DS DP	NP	Jun 08, 2009
	4992474*PED	Aug 12, 2008			
	5126375	Feb 12, 2008	DS DP		
	5126375*PED	Aug 12, 2008			
	5225445	Feb 12, 2008		U-738	
	5225445*PED	Aug 12, 2008			
	5270305	Sep 07, 2010		U-738	
	5658549	Aug 19, 2014	DP	U-738	
	5674472	Oct 07, 2014	DP		
	6143277	Apr 14, 2015	DP	U-738	
	6251368	Dec 04, 2012	DP		
	6253762	Apr 14, 2015	DP	U-738	
	6315173	Dec 23, 2017	DP		
	6510969	Dec 23, 2017	DP		
	6524555	Apr 14, 2015	DP		
	6546928	Apr 14, 2015	DP		
<u>FLUVASTATIN SODIUM - LESCOL</u>					
020261 001				I-507 PED	Apr 10, 2009 Oct 10, 2009
<u>FLUVASTATIN SODIUM - LESCOL</u>					
020261 002				I-507 PED	Apr 10, 2009 Oct 10, 2009
<u>FLUVASTATIN SODIUM - LESCOL XL</u>					
021192 001				I-507 PED	Apr 10, 2009 Oct 10, 2009
<u>FORMOTEROL FUMARATE - FORADIL</u>					
020831 001	6887459	Nov 28, 2020		U-762	
<u>FROVATRIPTAN SUCCINATE - FROVA</u>					
021006 001	5464864	Nov 07, 2015		U-436	
<u>FULVESTANT - FASLODEX</u>					
021344 001	4659516	Dec 11, 2007	DS DP	U-596	
<u>GALANTAMINE HYDROBROMIDE - RAZADYNE ER</u>					
021615 001				NDF	Apr 01, 2008
<u>GALANTAMINE HYDROBROMIDE - RAZADYNE ER</u>					
021615 002				NDF	Apr 01, 2008
<u>GALANTAMINE HYDROBROMIDE - RAZADYNE ER</u>					
021615 003				NDF	Apr 01, 2008
<u>GANIRELIX ACETATE - GANIRELIX ACETATE INJECTION</u>					
021057 001	4801577	Feb 05, 2012	DS DP		

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnotes for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>GEMCITABINE HYDROCHLORIDE - GEMZAR</u>					
020509 001				I-499	Jul 14, 2009
<u>GEMCITABINE HYDROCHLORIDE - GEMZAR</u>					
020509 002				I-499	Jul 14, 2009
<u>GEMIFLOXACIN MESYLATE - FACTIVE</u>					
021158 001	5776944	Apr 04, 2017	DS DP		
<u>GLIMEPIRIDE - AMARYL</u>					
020496 001				M-54 PED	Nov 28, 2008 May 28, 2009
<u>GLIMEPIRIDE - AMARYL</u>					
020496 002				M-54 PED	Nov 28, 2008 May 28, 2009
<u>GLIMEPIRIDE - AMARYL</u>					
020496 003				M-54 PED	Nov 28, 2008 May 28, 2009
<u>GLIMEPIRIDE; PIOGLITAZONE HYDROCHLORIDE - DUETACT</u>					
021925 001	4687777	Jan 17, 2011	DS		
	6150383	Jun 19, 2016		U-753	
	6211205	Jun 19, 2016		U-753	
	6303640	Aug 09, 2016		U-753	
	6329404	Jun 19, 2016	DP	U-753	
<u>GLIMEPIRIDE; PIOGLITAZONE HYDROCHLORIDE - DUETACT</u>					
021925 002	4687777	Jan 17, 2011	DS		
	6150383	Jun 19, 2016		U-753	
	6211205	Jun 19, 2016		U-753	
	6303640	Aug 09, 2016		U-753	
	6329404	Jun 19, 2016	DP	U-753	
<u>GLIMEPIRIDE; ROSIGLITAZONE MALEATE - AVANDARYL</u>					
021700 001	5002953	Sep 17, 2011	DS DP	U-781	
	5002953	Sep 17, 2011	DS DP	U-690	
	5002953*PED	Mar 17, 2012		U-781	
	5741803	Apr 21, 2015	DS DP	U-781	
	5741803	Apr 21, 2015	DS DP	U-690	
	5741803*PED	Oct 21, 2015			
<u>GLIMEPIRIDE; ROSIGLITAZONE MALEATE - AVANDARYL</u>					
021700 002	5002953	Sep 17, 2011	DS DP	U-781	
	5002953	Sep 17, 2011	DS DP	U-690	
	5002953*PED	Mar 17, 2012			
	5741803	Apr 21, 2015	DS DP	U-781	
	5741803	Apr 21, 2015	DS DP	U-690	
	5741803*PED	Oct 21, 2015			
<u>GLIMEPIRIDE; ROSIGLITAZONE MALEATE - AVANDARYL</u>					
021700 003	5002953	Sep 17, 2011	DS DP	U-781	
	5002953	Sep 17, 2011	DS DP	U-690	
	5002953*PED	Mar 17, 2012			
	5741803	Apr 21, 2015	DS DP	U-781	
	5741803	Apr 21, 2015	DS DP	U-690	
	5741803*PED	Oct 21, 2015			
<u>HYALURONIDASE RECOMBINANT HUMAN - HYLENEX RECOMBINANT</u>					
021859 001				NCE	Dec 02, 2010
<u>HYDROCHLOROTHIAZIDE; METOPROLOL SUCCINATE - DUTOPROL</u>					
021956 001				NC	Aug 28, 2009
<u>HYDROCHLOROTHIAZIDE; METOPROLOL SUCCINATE - DUTOPROL</u>					
021956 002				NC	Aug 28, 2009
<u>HYDROCHLOROTHIAZIDE; METOPROLOL SUCCINATE - DUTOPROL</u>					
021956 003				NC	Aug 28, 2009

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnotes for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE		PATENT CODES			EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE	
<u>HYDROCHLOROTHIAZIDE; VALSARTAN - DIOVAN HCT</u>									
020818 004							NS	Apr	28, 2009
<u>HYDROCHLOROTHIAZIDE; VALSARTAN - DIOVAN HCT</u>									
020818 005							NS	Apr	28, 2009
<u>HYDROXOCOBALAMIN - CYANOKIT</u>									
022041 002							>A> NP	Dec	15, 2009
<u>IBANDRONATE SODIUM - BONIVA</u>									
021455 001	4927814	Jul	09, 2007	DS	DP	U-642			
	6143326	Apr	21, 2017			U-642			
<u>IBANDRONATE SODIUM - BONIVA</u>									
021858 001	4927814	Jul	09, 2007	DS	DP	U-700	NDF	Jan	06, 2009
	5662918	Sep	02, 2014		DP		NCE	May	16, 2008
<u>IBUPROFEN LYSINE - NEOPROFEN</u>									
021903 001							NE	Apr	13, 2009
							ODE	Apr	13, 2013
<u>ICODEXTRIN - EXTRANEAL</u>									
021321 001	4761237	Aug	09, 2009			U-495			
<u>IMATINIB MESYLATE - GLEEVEC</u>									
021335 001	5521184	Jan	04, 2015	DS	DP		I-392	May	20, 2006
	5521184*PED	Jul	04, 2015				I-376	Dec	20, 2005
	6894051	May	23, 2019	DS	DP	U-649	NCE	May	10, 2006
	6894051*PED	Nov	23, 2019				ODE	Feb	01, 2009
	6958335	Dec	19, 2021	DS	DP		ODE	May	10, 2008
	6958335*PED	Jun	19, 2022				PED	Nov	10, 2008
							PED	Nov	20, 2006
							PED	Jun	20, 2006
							PED	Aug	01, 2009
							PED	Nov	10, 2006
<u>IMATINIB MESYLATE - GLEEVEC</u>									
021335 002	5521184	Jan	04, 2015				I-392	May	20, 2006
	5521184*PED	Jul	04, 2015				I-376	Dec	20, 2005
	6894051	May	23, 2019	DS	DP	U-649	NCE	May	10, 2006
	6894051*PED	Nov	23, 2019				ODE	Feb	01, 2009
	6958335	Dec	19, 2021	DS	DP		ODE	May	10, 2008
	6958335*PED	Jun	19, 2022				PED	Nov	10, 2008
							PED	Nov	20, 2006
							PED	Jun	20, 2006
							PED	Aug	01, 2009
							PED	Nov	10, 2006
<u>IMATINIB MESYLATE - GLEEVEC</u>									
021588 001	5521184	Jan	04, 2015	DS	DP		I-514	Oct	19, 2009
	5521184*PED	Jul	04, 2015				I-510	Oct	19, 2009
	6894051	May	23, 2019	DS	DP	U-649	I-511	Oct	19, 2009
	6894051*PED	Nov	23, 2019				I-513	Oct	19, 2009
	6958335	Dec	19, 2021	DS	DP		I-512	Oct	19, 2009
	6958335*PED	Jun	19, 2022				I-376	Dec	20, 2005
							I-392	May	20, 2006
							NCE	May	10, 2006
							ODE	Oct	19, 2013
							ODE	Oct	19, 2013
							ODE	Oct	19, 2013
							ODE	Oct	19, 2013
							ODE	Oct	19, 2013
							ODE	Oct	19, 2013
							ODE	Feb	01, 2009
							ODE	May	10, 2008
							PED	Aug	01, 2009
							PED	Nov	10, 2008
							PED	Nov	20, 2006
							PED	Jun	20, 2006
							PED	Nov	10, 2006

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnotes for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE		PATENT CODES			EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE	
<u>IMATINIB MESYLATE - GLEEVEC</u>									
021588 002	5521184	Jan	04, 2015				I-514	Oct	19, 2009
	5521184*PED	Jul	04, 2015				I-512	Oct	19, 2009
	6894051	May	23, 2019	DS	DP	U-649	I-513	Oct	19, 2009
	6894051*PED	Nov	23, 2019				I-511	Oct	19, 2009
	6958335	Dec	19, 2021	DS	DP		I-510	Oct	19, 2009
	6958335*PED	Jun	19, 2022				I-376	Dec	20, 2005
							I-392	May	20, 2006
							NCE	May	10, 2006
							ODE	Oct	19, 2013
							ODE	Oct	19, 2013
							ODE	Oct	19, 2013
							ODE	Oct	19, 2013
							ODE	Oct	19, 2013
							ODE	Feb	01, 2009
							ODE	May	10, 2008
							PED	Nov	10, 2006
							PED	Aug	01, 2009
							PED	Jun	20, 2006
							PED	Nov	10, 2008
							PED	Nov	20, 2006
<u>IMIQUIMOD - ALDARA</u>									
020723 001	4689338	Aug	25, 2009			U-172	I-433	Jul	14, 2007
	4689338*PED	Feb	25, 2010				I-420	Mar	02, 2007
	5238944	Aug	24, 2010				PED	Sep	02, 2007
	5238944*PED	Feb	24, 2011				PED	Jan	14, 2008
<u>INSULIN DETEMIR RECOMBINANT - LEVEMIR</u>									
021536 001							I-489	Oct	19, 2008
<u>INSULIN GLARGINE RECOMBINANT - LANTUS</u>									
021081 001	6100376	Nov	06, 2009	DS	DP	U-771			
	6100376*PED	May	06, 2010						
<u>INSULIN RECOMBINANT HUMAN - EXUBERA</u>									
021868 001	5740794	Apr	21, 2015		DP		NP	Jan	27, 2009
	5997848	Mar	07, 2014			U-704			
	6051256	Mar	07, 2014		DP				
	6257233	May	14, 2019			U-704			
	6423344	Mar	07, 2014		DP				
	6543448	Sep	21, 2014		DP				
	6546929	May	14, 2019			U-704			
	6582728	Jun	24, 2020		DP				
	6592904	Mar	07, 2014		DP				
	6685967	Sep	11, 2018		DP				
	6737045	Mar	07, 2014			U-704			
	RE37872	Feb	12, 2010		DP				
	RE38385	Feb	12, 2010		DP				
<u>INSULIN RECOMBINANT HUMAN - EXUBERA</u>									
021868 002	5740794	Apr	21, 2015		DP		NP	Jan	27, 2009
	5997848	Mar	07, 2014			U-704			
	6051256	Mar	07, 2014		DP				
	6257233	May	14, 2019			U-704			
	6423344	Mar	07, 2014		DP				
	6543448	Sep	21, 2014		DP				
	6546929	May	14, 2019			U-704			
	6582728	Jun	24, 2020		DP				
	6592904	Mar	07, 2014		DP				
	6685967	Sep	11, 2018		DP				
	6737045	Mar	07, 2014			U-704			
	RE37872	Feb	12, 2010		DP				
	RE38385	Feb	12, 2010		DP				
<u>IODINE POVACRYLEX; ISOPROPYL ALCOHOL - DURAPREP</u>									
021586 001							NC	Sep	29, 2009

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnotes for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>IODINE POVACRYLEX; ISOPROPYL ALCOHOL - DURAPREP</u>					
021586 002				NC	Sep 29, 2009
<u>IPRATROPIUM BROMIDE - ATROVENT HFA</u>					
021527 001	6983743	May 26, 2020	DP		
<u>KETOCONAZOLE - XOLEGEL</u>					
021946 001				NDF	Jul 28, 2009
<u>KETOTIFEN FUMARATE - KETOTIFEN FUMARATE</u>					
077354 001				PC	Dec 30, 2006
<u>KUNECATECHINS - VEREGEN</u>					
021902 001	5795911	Apr 10, 2017	U-172	NP	Oct 31, 2009
	5968973	Apr 10, 2017	U-172		
<u>LAMOTRIGINE - LAMICTAL</u>					
020241 001				I-516	Sep 22, 2009
<u>LAMOTRIGINE - LAMICTAL</u>					
020241 002				I-516	Sep 22, 2009
<u>LAMOTRIGINE - LAMICTAL</u>					
020241 003				I-516	Sep 22, 2009
<u>LAMOTRIGINE - LAMICTAL</u>					
020241 004				I-516	Sep 22, 2009
<u>LAMOTRIGINE - LAMICTAL</u>					
020241 005				I-516	Sep 22, 2009
<u>LAMOTRIGINE - LAMICTAL</u>					
020241 006				I-516	Sep 22, 2009
<u>LAMOTRIGINE - LAMICTAL CD</u>					
020764 001				I-516	Sep 22, 2009
<u>LAMOTRIGINE - LAMICTAL CD</u>					
020764 002				I-516	Sep 22, 2009
<u>LAMOTRIGINE - LAMICTAL CD</u>					
020764 003				I-516	Sep 22, 2009
<u>LAMOTRIGINE - LAMICTAL CD</u>					
020764 004				I-516	Sep 22, 2009
<u>LAMOTRIGINE - LAMOTRIGINE</u>					
076420 001				PC	Feb 25, 2007
<u>LAMOTRIGINE - LAMOTRIGINE</u>					
076420 002				PC	Dec 25, 2006
<u>LANSOPRAZOLE - PREVACID</u>					
020406 001	6749864	Feb 13, 2007	DP		
<u>LANSOPRAZOLE - PREVACID</u>					
020406 002	6749864	Feb 13, 2007	DP		
<u>LANSOPRAZOLE - PREVACID</u>					
021281 001	6749864	Feb 13, 2007	DP		
<u>LANSOPRAZOLE - PREVACID</u>					
021281 002	6749864	Feb 13, 2007	DP		
<u>LANSOPRAZOLE - PREVACID</u>					
021428 001	6749864	Feb 13, 2007	DP		
<u>LANSOPRAZOLE - PREVACID</u>					
021428 002	6749864	Feb 13, 2007	DP		
<u>LANTHANUM CARBONATE - FOSRENOL</u>					
021468 003	5968976	Mar 19, 2016	DP	U-613	
<u>LANTHANUM CARBONATE - FOSRENOL</u>					
021468 004	5968976	Mar 19, 2016	DP	U-613	
<u>LATANOPROST - XALATAN</u>					
020597 001	5296504	Mar 22, 2011	DP	U-778	
	5422368	Mar 22, 2011	DP	U-778	
	6429226	Sep 06, 2009	DP	U-778	

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnotes for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE		PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE	
<u>LENALIDOMIDE - REVLIMID</u>							
021880 001	5635517	Jul	24, 2016	DS		ODE	Jun 29, 2013
	6045501	Aug	28, 2018		U-694	ODE	Dec 27, 2012
	6281230	Jul	24, 2016		U-769		
	6315720	Oct	23, 2020		U-694		
	6555554	Jul	24, 2016	DP			
	6561976	Aug	28, 2018		U-694		
	6561977	Oct	23, 2020		U-694		
	6755784	Oct	23, 2020		U-694		
	6908432	Aug	28, 2018		U-694		
	7119106	Jul	24, 2016	DP			
<u>LENALIDOMIDE - REVLIMID</u>							
021880 002	5635517	Jul	24, 2016	DS		ODE	Jun 29, 2013
	6045501	Aug	28, 2018		U-694	ODE	Dec 27, 2012
	6281230	Jul	24, 2016		U-769		
	6315720	Oct	23, 2020		U-694		
	6555554	Jul	24, 2016	DP			
	6561976	Aug	28, 2018		U-694		
	6561977	Oct	23, 2020		U-694		
	6755784	Oct	23, 2020		U-694		
	6908432	Aug	28, 2018		U-694		
	7119106	Jul	24, 2016	DP			
<u>LENALIDOMIDE - REVLIMID</u>							
021880 003	5635517	Jul	24, 2016	DS		I-500	Jun 29, 2009
	6045501	Aug	28, 2018		U-694	NCE	Dec 27, 2010
	6281230	Jul	24, 2016		U-769	ODE	Jun 29, 2013
	6315720	Oct	23, 2020		U-694		
	6555554	Jul	24, 2016	DP			
	6561976	Aug	28, 2018		U-694		
	6561977	Oct	23, 2020		U-694		
	6755784	Oct	23, 2020		U-694		
	6908432	Aug	28, 2018		U-694		
	7119106	Jul	24, 2016	DP			
<u>LENALIDOMIDE - REVLIMID</u>							
021880 004	5635517	Jul	24, 2016	DS		I-500	Jun 29, 2009
	6045501	Aug	28, 2018		U-694	NCE	Dec 27, 2010
	6281230	Jul	24, 2016		U-769	ODE	Jun 29, 2013
	6315720	Oct	23, 2020		U-694		
	6555554	Jul	24, 2016	DP			
	6561976	Aug	28, 2018		U-694		
	6561977	Oct	23, 2020		U-694		
	6755784	Oct	23, 2020		U-694		
	6908432	Aug	28, 2018		U-694		
	7119106	Jul	24, 2016	DP			
<u>LEVALBUTEROL TARTRATE - XOPENEX HFA</u>							
021730 001	5362755	Mar	25, 2013		U-636		
<u>LEVETIRACETAM - KEPPRA</u>							
021035 001						I-506	Aug 15, 2009
<u>LEVETIRACETAM - KEPPRA</u>							
021035 002						I-506	Aug 15, 2009
<u>LEVETIRACETAM - KEPPRA</u>							
021035 003						I-506	Aug 15, 2009
<u>LEVETIRACETAM - KEPPRA</u>							
021035 004	4943639	Jul	14, 2008	DS		I-506 NPP	Aug 15, 2009 Jun 21, 2008
<u>LEVETIRACETAM - KEPPRA</u>							
021505 001						I-506	Aug 15, 2009

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnotes for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>LEVETIRACETAM - KEPRA</u>					
021872 001	4943639	Jul 14, 2008	DS	NDF	Jul 31, 2009
<u>LEVOBETAXOLOL HYDROCHLORIDE - BETAXON</u>					
021114 001	4911920	Mar 27, 2007	DS DP	M-54	Sep 28, 2009
	4911920	Mar 27, 2007	DS DP	PED	Mar 28, 2010
	4911920*PED	Sep 27, 2007			
	5540918	Jul 30, 2013	DP		
	5540918*PED	Jan 30, 2014			
<u>LEVONORGESTREL - PLAN B</u>					
021045 002				NP	Aug 24, 2009
<u>LEVOTHYROXINE SODIUM - LEVOXYL</u>					
021301 001	7067148	Feb 15, 2022	DP		
	7101569	Aug 14, 2022		U-759	
<u>LEVOTHYROXINE SODIUM - LEVOXYL</u>					
021301 002	7067148	Feb 15, 2022	DP		
	7101569	Aug 14, 2022		U-759	
<u>LEVOTHYROXINE SODIUM - LEVOXYL</u>					
021301 003	7067148	Feb 15, 2022	DP		
	7101569	Aug 14, 2022		U-759	
<u>LEVOTHYROXINE SODIUM - LEVOXYL</u>					
021301 004	7067148	Feb 15, 2022	DP		
	7101569	Aug 14, 2022		U-759	
<u>LEVOTHYROXINE SODIUM - LEVOXYL</u>					
021301 005	7067148	Feb 15, 2022	DP		
	7101569	Aug 14, 2022		U-759	
<u>LEVOTHYROXINE SODIUM - LEVOXYL</u>					
021301 006	7067148	Feb 15, 2022	DP		
	7101569	Aug 14, 2022		U-759	
<u>LEVOTHYROXINE SODIUM - LEVOXYL</u>					
021301 007	7067148	Feb 15, 2022	DP		
	7101569	Aug 14, 2022		U-759	
<u>LEVOTHYROXINE SODIUM - LEVOXYL</u>					
021301 008	7067148	Feb 15, 2022	DP		
	7101569	Aug 14, 2022		U-759	
<u>LEVOTHYROXINE SODIUM - LEVOXYL</u>					
021301 009	7067148	Feb 15, 2022	DP		
	7101569	Aug 14, 2022		U-759	
<u>LEVOTHYROXINE SODIUM - LEVOXYL</u>					
021301 010	7067148	Feb 15, 2022	DP		
	7101569	Aug 14, 2022		U-759	
<u>LEVOTHYROXINE SODIUM - LEVOXYL</u>					
021301 011	7067148	Feb 15, 2022	DP		
	7101569	Aug 14, 2022		U-759	
<u>LEVOTHYROXINE SODIUM - LEVOXYL</u>					
021301 012	7067148	Feb 15, 2022	DP		
	7101569	Aug 14, 2022		U-759	
<u>LIDOCAINE; TETRACAINE - LIDOCAINE AND TETRACAINE</u>					
021717 001	5919479	Jul 28, 2015	DP	NP	Jun 29, 2009
	6528086	Sep 28, 2019	DP		
<u>LIDOCAINE; TETRACAINE - SYNERA</u>					
021623 001				NC	Jun 23, 2008
<u>LOPERAMIDE HYDROCHLORIDE; SIMETHICONE - LOPERAMIDE HYDROCHLORIDE AND SIMETHICONE</u>					
077500 001				PC	May 12, 2007

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnotes for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>LOPINAVIR; RITONAVIR - KALETRA</u>					
021226 001	>A> 5886036	Nov 19, 2013	DS DP		
	>A> 5886036*PED	May 19, 2014			
	>A> 5948436	Sep 13, 2013	DP		
	>A> 5948436*PED	Mar 13, 2014			
	>A> 7141593	May 22, 2020	DP		
	>A> 7141593*PED	Nov 22, 2020			
<u>LOPINAVIR; RITONAVIR - KALETRA</u>					
021251 001	>A> 5886036	Nov 19, 2013	DS DP		
	>A> 5886036*PED	May 19, 2014			
	>A> 5948436	Sep 13, 2013	DP		
	>A> 5948436*PED	Mar 13, 2014			
<u>LOPINAVIR; RITONAVIR - KALETRA</u>					
021906 001	5541206	Jul 30, 2013	DS DP	U-688	
	5541206*PED	Jan 30, 2014			
	5635523	Jun 03, 2014		U-688	
	5635523*PED	Dec 03, 2014			
	5648497	Jul 15, 2014	DS DP		
	5648497*PED	Jan 15, 2015			
	5674882	Oct 07, 2014		U-688	
	5674882*PED	Apr 07, 2015			
	5846987	Dec 29, 2012		U-688	
	5846987*PED	Jun 29, 2013			
	>A> 5886036	Nov 19, 2013	DS DP		
	>A> 5886036*PED	May 19, 2014			
	6037157	Jun 26, 2016		U-688	
	6037157*PED	Dec 26, 2016			
	6703403	Jun 26, 2016		U-688	
	6703403*PED	Dec 26, 2016			
<u>LOVASTATIN; NIACIN - ADVICOR</u>					
021249 001	6469035	Mar 15, 2018		U-768	
	7011848	Sep 20, 2013		U-712	
<u>LOVASTATIN; NIACIN - ADVICOR</u>					
021249 002	6469035	Mar 15, 2018		U-768	
	7011848	Sep 20, 2013		U-712	
<u>LOVASTATIN; NIACIN - ADVICOR</u>					
021249 003	6469035	Mar 15, 2018		U-768	
	7011848	Sep 20, 2013		U-712	
<u>LOVASTATIN; NIACIN - ADVICOR</u>					
021249 004	6469035	Mar 15, 2018		U-768	
<u>LUBIPROSTONE - AMITIZA</u>					
021908 001	5284858	Feb 08, 2011	DS DP	NCE	Jan 31, 2011
	5317032	May 31, 2011	DS DP	U-717	
	6414016	Sep 05, 2020	DS DP	U-717	
	6583174	Oct 16, 2020	DS DP		
	7064148	Aug 30, 2022	DS DP	U-739	
<u>MAGNESIUM CHLORIDE; SODIUM BICARBONATE; SODIUM CHLORIDE - NORMOCARB HF 25</u>					
021910 001	>A> 5945449	Oct 31, 2017	DP	U-785	ODE Jul 26, 2013
<u>MAGNESIUM CHLORIDE; SODIUM BICARBONATE; SODIUM CHLORIDE - NORMOCARB HF 35</u>					
021910 002	>A> 5945449	Oct 31, 2017	DP	U-785	ODE Jul 26, 2013
<u>MAGNESIUM HYDROXIDE; OMEPRAZOLE; SODIUM BICARBONATE - ZEGEID</u>					
021850 001	6489346	Jul 16, 2016	DS DP	U-623	
	6489346	Jul 16, 2016	DS DP	U-588	
	6645988	Jul 16, 2016	DS DP	U-623	
	6645988	Jul 16, 2016	DS DP	U-588	
	6699885	Jul 16, 2016		U-623	
	6699885	Jul 16, 2016		U-588	

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnotes for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>MAGNESIUM HYDROXIDE; OMEPRAZOLE; SODIUM BICARBONATE - ZEGERID</u>					
021850 002	6489346	Jul 16, 2016	DS DP	U-623	
	6489346	Jul 16, 2016	DS DP	U-588	
	6645988	Jul 16, 2016	DS DP	U-623	
	6645988	Jul 16, 2016	DS DP	U-588	
	6699885	Jul 16, 2016		U-623	
	6699885	Jul 16, 2016		U-588	
<u>MECASERMIN RINFABATE RECOMBINANT - IPLEX</u>					
021884 001	5200509	Apr 06, 2010	DS		
	5681818	Oct 28, 2014		U-697	
<u>MEGESTROL ACETATE - MEGACE ES</u>					
021778 001	7101576	Apr 22, 2024		U-755	
<u>MELOXICAM - MOBIC</u>					
020938 001				ODE PED	Aug 11, 2012 Feb 11, 2013
<u>MELOXICAM - MOBIC</u>					
020938 002				ODE PED	Aug 11, 2012 Feb 11, 2013
<u>MELOXICAM - MOBIC</u>					
021530 001				I-469 ODE PED PED	Aug 11, 2008 Aug 11, 2012 Feb 11, 2013 Feb 11, 2009
<u>MEQUINOL; TRETINOIN - SOLAGE</u>					
020922 001	5194247	Dec 10, 2013	DP	U-294	
<u>METFORMIN HYDROCHLORIDE; ROSIGLITAZONE MALEATE - AVANDAMET</u>					
021410 001	5002953	Sep 17, 2011	DS DP	U-690	I-494 May 19, 2009
	5002953	Sep 17, 2011	DS DP	U-734	
	5002953	Sep 17, 2011	DS DP	U-691	
	5002953	Sep 17, 2011	DS DP	U-493	
	5741803	Apr 21, 2015	DS DP	U-734	
	5741803	Apr 21, 2015	DS DP	U-493	
<u>METFORMIN HYDROCHLORIDE; ROSIGLITAZONE MALEATE - AVANDAMET</u>					
021410 002	5002953	Sep 17, 2011	DS DP	U-690	I-494 May 19, 2009
	5002953	Sep 17, 2011	DS DP	U-734	
	5002953	Sep 17, 2011	DS DP	U-691	
	5002953	Sep 17, 2011	DS DP	U-493	
	5741803	Apr 21, 2015	DS DP	U-734	
	5741803	Apr 21, 2015	DS DP	U-493	
<u>METFORMIN HYDROCHLORIDE; ROSIGLITAZONE MALEATE - AVANDAMET</u>					
021410 003	5002953	Sep 17, 2011	DS DP	U-691	I-494 May 19, 2009
	5002953	Sep 17, 2011	DS DP	U-734	
	5002953	Sep 17, 2011	DS DP	U-690	
	5002953	Sep 17, 2011	DS DP	U-493	
	5741803	Apr 21, 2015	DS DP	U-734	
	5741803	Apr 21, 2015	DS DP	U-493	
<u>METFORMIN HYDROCHLORIDE; ROSIGLITAZONE MALEATE - AVANDAMET</u>					
021410 004	5002953	Sep 17, 2011	DS DP	U-691	I-494 May 19, 2009
	5002953	Sep 17, 2011	DS DP	U-734	
	5002953	Sep 17, 2011	DS DP	U-690	
	5002953	Sep 17, 2011	DS DP	U-493	
	5741803	Apr 21, 2015	DS DP	U-734	
	5741803	Apr 21, 2015	DS DP	U-493	

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnotes for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
METFORMIN HYDROCHLORIDE; ROSIGLITAZONE MALEATE - AVANDAMET					
021410 005	5002953	Sep 17, 2011	DS DP	I-494	May 19, 2009
	5002953	Sep 17, 2011	DS DP		
	5002953	Sep 17, 2011	DS DP		
	5002953	Sep 17, 2011	DS DP		
	5741803	Apr 21, 2015	DS DP		
	5741803	Apr 21, 2015	DS DP		
	5741803*PED	Oct 21, 2015			
METHYLPHENIDATE - DAYTRANA					
021514 001	5958446	Dec 12, 2012	DP	NDF	Apr 06, 2009
	6210705	Sep 30, 2018	DP		
	6348211	Sep 30, 2018	DP		
METHYLPHENIDATE - DAYTRANA					
021514 002	5958446	Dec 12, 2012	DP	NDF	Apr 06, 2009
	6210705	Sep 30, 2018	DP		
	6348211	Sep 30, 2018	DP		
METHYLPHENIDATE - DAYTRANA					
021514 003	5958446	Dec 12, 2012	DP	NDF	Apr 06, 2009
	6210705	Sep 30, 2018	DP		
	6348211	Sep 30, 2018	DP		
METHYLPHENIDATE - DAYTRANA					
021514 004	5958446	Dec 12, 2012	DP	NDF	Apr 06, 2009
	6210705	Sep 30, 2018	DP		
	6348211	Sep 30, 2018	DP		
METHYLPHENIDATE HYDROCHLORIDE - METADATE CD					
021259 001	6344215	Oct 27, 2020	DP		
METHYLPHENIDATE HYDROCHLORIDE - METADATE CD					
021259 002	6344215	Oct 27, 2020	DP		
METHYLPHENIDATE HYDROCHLORIDE - METADATE CD					
021259 003	6344215	Oct 27, 2020	DP		
METHYLPHENIDATE HYDROCHLORIDE - METADATE CD					
021259 004	6344215	Oct 27, 2020	DP		
METOPROLOL SUCCINATE - TOPROL-XL					
019962 001	4927640	May 22, 2007	DP	D-95 PED	Feb 15, 2008 Aug 15, 2008
	4927640*PED	Nov 22, 2007			
	4957745	Sep 18, 2007	DP		
	4957745*PED	Mar 18, 2008			
	5001161	Sep 18, 2007	DP		
	5001161*PED	Mar 18, 2008			
	5081154	Sep 18, 2007	DS		
	5081154*PED	Mar 18, 2008			
	>A> 5246714	Sep 21, 2010			
	>A> 5246714*PED	Mar 21, 2011			
METOPROLOL SUCCINATE - TOPROL-XL					
019962 002	4927640	May 22, 2007	DP	D-95 PED	Feb 15, 2008 Aug 15, 2008
	4927640*PED	Nov 22, 2007			
	4957745	Sep 18, 2007	DP		
	4957745*PED	Mar 18, 2008			
	5001161	Sep 18, 2007	DP		
	5001161*PED	Mar 18, 2008			
	5081154	Sep 18, 2007	DS		
	5081154*PED	Mar 18, 2008			
	>A> 5246714	Sep 21, 2010			
	>A> 5246714*PED	Mar 21, 2011			

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnotes for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE		PATENT CODES		EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE	
<u>METOPROLOL SUCCINATE - TOPROL-XL</u>								
019962 003	4927640	May	22, 2007	DP		D-95 PED	Feb 15, 2008	
	4927640*PED	Nov	22, 2007				Aug 15, 2008	
	4957745	Sep	18, 2007	DP	U-107			
	4957745*PED	Mar	18, 2008					
	5001161	Sep	18, 2007	DP				
	5001161*PED	Mar	18, 2008					
	5081154	Sep	18, 2007	DS				
	5081154*PED	Mar	18, 2008					
	>A> 5246714	Sep	21, 2010					
	>A> 5246714*PED	Mar	21, 2011					
<u>METOPROLOL SUCCINATE - TOPROL-XL</u>								
019962 004	4927640	May	22, 2007	DP		D-95 PED	Feb 15, 2008	
	4927640*PED	Nov	22, 2007				Aug 15, 2008	
	4957745	Sep	18, 2007	DP	U-107			
	4957745*PED	Mar	18, 2008					
	5001161	Sep	18, 2007	DP	U-107			
	5001161*PED	Mar	18, 2008					
	5081154	Sep	18, 2007	DS	U-107			
	5081154*PED	Mar	18, 2008					
	>A> 5246714	Sep	21, 2010					
	>A> 5246714*PED	Mar	21, 2011					
<u>METRONIDAZOLE - METROGEL</u>								
021789 001	6881726	Feb	21, 2022	DP	U-743			
<u>MICONAZOLE NITRATE; PETROLATUM, WHITE; ZINC OXIDE - VUSION</u>								
021026 001	4911932	Mar	27, 2007	DP	U-718	NP	Feb 16, 2009	
<u>MINOXIDIL - MEN'S ROGAINE</u>								
021812 001	6946120	Apr	20, 2019	DP	U-702	NDF	Jan 20, 2009	
<u>MODAFINIL - PROVIGIL</u>								
020717 001	4927855	May	22, 2007		U-255	I-449 ODE PED PED	Jan 23, 2007	
	4927855*PED	Nov	22, 2007				Dec 24, 2005	
	RE37516	Oct	06, 2014		U-255		Jul 23, 2007	
	RE37516*PED	Apr	06, 2015				Jun 24, 2006	
<u>MODAFINIL - PROVIGIL</u>								
020717 002	4927855	May	22, 2007		U-255	I-449 ODE PED PED	Jan 23, 2007	
	4927855*PED	Nov	22, 2007				Dec 24, 2005	
	RE37516	Oct	06, 2014		U-255		Jul 23, 2007	
	RE37516*PED	Apr	06, 2015				Jun 24, 2006	
<u>MORPHINE SULFATE - KADIAN</u>								
020616 004	5378474	Mar	23, 2010					
<u>MORPHINE SULFATE - KADIAN</u>								
020616 005	5202128	Apr	13, 2010					
	5378474	Mar	23, 2010					
<u>MOXIFLOXACIN HYDROCHLORIDE - AVELOX</u>								
021085 001	4990517	Dec	08, 2011	DS	DP	U-298		
	6610327	Oct	29, 2019		DP	U-298		
<u>MOXIFLOXACIN HYDROCHLORIDE - AVELOX IN SODIUM CHLORIDE 0.8% IN PLASTIC CONTAINER</u>								
021277 001	4990517	Dec	08, 2011	DS	DP	U-298		
	6548079	Jul	25, 2020		DP	U-298		
<u>MOXIFLOXACIN HYDROCHLORIDE - VIGAMOX</u>								
021598 001	4990517	Dec	08, 2011	DS	DP	U-709		
	4990517*PED	Jun	08, 2012					

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnotes for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>NALTREXONE - VIVITROL</u>					
021897 001	5792477	May 02, 2017	DP	NDF	Apr 13, 2009
	5916598	May 02, 2017	DP		
	6110503	May 02, 2017	DP		
	6194006	Dec 30, 2018	DP		
	6264987	May 19, 2020	DP		
	6331317	Nov 12, 2019	DP		
	6379703	Dec 30, 2018	DP		
	6379704	May 19, 2020	DP		
	6395304	Nov 12, 2019	DP		
	6403114	May 02, 2017	DP		
	6495164	May 25, 2020	DP		
	6495166	Nov 12, 2019	DP		
	6534092	May 19, 2020	DP		
	6537586	Nov 12, 2019	DP		
	6596316	Dec 30, 2018	DP		
	6667061	May 25, 2020	DP		
	6713090	Nov 12, 2019	DP		
	6939033	Nov 12, 2019	DP		
	<u>NELARABINE - ARRANON</u>				
021877 001	5747472	Feb 20, 2013	U-696		
	5747472	Feb 20, 2013	U-695		
	5747472	Feb 20, 2013	U-689		
	5821236	Feb 20, 2013	U-695		
<u>NIACIN - NIASPAN</u>					
020381 001	7011848	Sep 20, 2013	U-712		
<u>NIACIN - NIASPAN</u>					
020381 002	6469035	Mar 15, 2018	U-768		
	7011848	Sep 20, 2013	U-712		
<u>NIACIN - NIASPAN</u>					
020381 003	6469035	Mar 15, 2018	U-768		
	7011848	Sep 20, 2013	U-712		
<u>NIACIN - NIASPAN</u>					
020381 004	6469035	Mar 15, 2018	U-768		
	7011848	Sep 20, 2013	U-712		
<u>NIACIN - NIASPAN TITRATION STARTER PACK</u>					
020381 005	7011848	Sep 20, 2013	U-712		
<u>NICOTINE POLACRILEX - NICOTINE POLACRILEX</u>					
077007 001				PC	Aug 21, 2006
<u>NICOTINE POLACRILEX - NICOTINE POLACRILEX</u>					
077007 002				PC	Aug 21, 2006
<u>NITROGLYCERIN - NITROMIST</u>					
021780 001				NP	Nov 02, 2009
<u>OCTREOTIDE ACETATE - SANDOSTATIN</u>					
019667 001	5753618	May 19, 2015			
	5753618*PED	Nov 19, 2015			
<u>OCTREOTIDE ACETATE - SANDOSTATIN</u>					
019667 002	5753618	May 19, 2015			
	5753618*PED	Nov 19, 2015			
<u>OCTREOTIDE ACETATE - SANDOSTATIN</u>					
019667 003	5753618	May 19, 2015			
	5753618*PED	Nov 19, 2015			
<u>OCTREOTIDE ACETATE - SANDOSTATIN</u>					
019667 004	5753618	May 19, 2015			
	5753618*PED	Nov 19, 2015			

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnotes for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>OCTREOTIDE ACETATE - SANDOSTATIN</u>					
019667 005	5753618	May 19, 2015			
	5753618*PED	Nov 19, 2015			
<u>OCTREOTIDE ACETATE - SANDOSTATIN LAR</u>					
021008 001	5538739	Jul 23, 2013	DP	M-55	May 10, 2009
	5538739*PED	Jan 23, 2014		ODE	Nov 25, 2005
	5639480	Jun 17, 2014	DP	PED	Nov 10, 2009
	5639480*PED	Dec 17, 2014		PED	May 25, 2006
	5688530	Nov 18, 2014	U-268		
	5688530*PED	May 18, 2015			
	5922338	Jul 13, 2016	DP		
	5922338*PED	Jan 13, 2017			
	5922682	Jul 13, 2016	DP		
	5922682*PED	Jan 13, 2017			
<u>OCTREOTIDE ACETATE - SANDOSTATIN LAR</u>					
021008 002	5538739	Jul 23, 2013		M-55	May 10, 2009
	5538739*PED	Jan 23, 2014		ODE	Nov 25, 2005
	5639480	Jun 17, 2014	DP	PED	Nov 10, 2009
	5639480*PED	Dec 17, 2014		PED	May 25, 2006
	5688530	Nov 18, 2014	U-268		
	5688530*PED	May 18, 2015			
	5922338	Jul 13, 2016	DP		
	5922338*PED	Jan 13, 2017			
	5922682	Jul 13, 2016	DP		
	5922682*PED	Jan 13, 2017			
<u>OCTREOTIDE ACETATE - SANDOSTATIN LAR</u>					
021008 003	5538739	Jul 23, 2013		M-55	May 25, 2009
	5538739*PED	Jan 23, 2014		ODE	Nov 25, 2005
	5639480	Jun 17, 2014	DP	PED	Nov 25, 2009
	5639480*PED	Dec 17, 2014		PED	May 25, 2006
	5688530	Nov 18, 2014	U-268		
	5688530*PED	May 18, 2015			
	5922338	Jul 13, 2016	DP		
	5922338*PED	Jan 13, 2017			
	5922682	Jul 13, 2016	DP		
	5922682*PED	Jan 13, 2017			
<u>OLOPATADINE HYDROCHLORIDE - PATADAY</u>					
021545 001	5116863	Dec 18, 2010	DS DP		
	5641805	Jun 06, 2015		U-765	
	6995186	Nov 12, 2023	DP	U-765	
<u>OMEPRazole; SODIUM BICARBONATE - ZEGERID</u>					
021849 001	6489346	Jul 16, 2016	DS DP	U-588	
	6645988	Jul 16, 2016	DS DP		
	6699885	Jul 16, 2016		U-588	
<u>OMEPRazole; SODIUM BICARBONATE - ZEGERID</u>					
021849 002	6489346	Jul 16, 2016	DS DP	U-623	
	6489346	Jul 16, 2016	DS DP	U-588	
	6645988	Jul 16, 2016	DS DP		
	6699885	Jul 16, 2016		U-623	
	6699885	Jul 16, 2016		U-588	
<u>ONDANSETRON HYDROCHLORIDE - ONDANSETRON HYDROCHLORIDE</u>					
076960 001				>A> PC	Jun 25, 2007

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnotes for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>OXALIPLATIN - ELOXATIN</u>					
021492 001	5290961	Jan 12, 2013		I-441	Nov 04, 2007
	5290961*PED	Jul 12, 2013		I-425	Jan 09, 2007
	5338874	Apr 07, 2013	DS	NCE	Aug 09, 2007
	5338874*PED	Oct 07, 2013		PED	Jul 09, 2007
	5420319	Aug 09, 2016	DS	PED	Feb 09, 2008
	5420319*PED	Aug 09, 2016		PED	May 04, 2008
<u>OXALIPLATIN - ELOXATIN</u>					
021492 002	5290961	Jan 12, 2013		I-441	Nov 04, 2007
	5290961*PED	Jul 12, 2013		I-425	Jan 09, 2007
	5338874	Apr 07, 2013	DS	NCE	Aug 09, 2007
	5338874*PED	Oct 07, 2013		PED	Jul 09, 2007
	5420319	Aug 09, 2016	DS	PED	Feb 09, 2008
	5420319*PED	Aug 09, 2016		PED	May 04, 2008
<u>OXALIPLATIN - ELOXATIN</u>					
021759 001	5290961	Jan 12, 2013	DS	I-441	Nov 04, 2007
	5290961*PED	Jul 12, 2013		NCE	Aug 09, 2007
	5338874	Apr 07, 2013	DS	PED	Feb 09, 2008
	5338874*PED	Oct 07, 2013		PED	May 04, 2008
	5420319	Aug 08, 2016	DS		
	5420319*PED	Feb 08, 2017			
	5716988	Aug 07, 2015	DP		
	5716988*PED	Feb 07, 2016			
<u>OXALIPLATIN - ELOXATIN</u>					
021759 002	5290961	Jan 12, 2013	DS	I-441	Nov 04, 2007
	5290961*PED	Jul 12, 2013		NCE	Aug 09, 2007
	5338874	Apr 07, 2013	DS	PED	Feb 09, 2008
	5338874*PED	Oct 07, 2013		PED	May 04, 2008
	5420319	Aug 08, 2016	DS		
	5420319*PED	Feb 08, 2017			
	5716988	Aug 07, 2015	DP		
	5716988*PED	Feb 07, 2016			
<u>OXALIPLATIN - ELOXATIN</u>					
021759 003				>A> I-441	Nov 04, 2007
				>A> NCE	Aug 09, 2007
				>A> PED	Feb 09, 2008
				>A> PED	May 04, 2008
<u>OXCARBAZEPINE - TRILEPTAL</u>					
021014 001	7037525	Feb 12, 2018	U-724		
<u>OXCARBAZEPINE - TRILEPTAL</u>					
021014 002	7037525	Feb 12, 2018	U-724		
<u>OXCARBAZEPINE - TRILEPTAL</u>					
021014 003	7037525	Feb 12, 2018	U-724		
<u>OXCARBAZEPINE - TRILEPTAL</u>					
021285 001	7037525	Feb 12, 2018	U-724		
<u>OXYBUTYNIN CHLORIDE - OXYBUTYNIN CHLORIDE</u>					
076644 001				PC	May 09, 2007
<u>OXYBUTYNIN CHLORIDE - OXYBUTYNIN CHLORIDE</u>					
076702 001				PC	May 09, 2007
<u>OXYBUTYNIN CHLORIDE - OXYBUTYNIN CHLORIDE</u>					
076745 001				PC	May 09, 2007
<u>OXYCODONE HYDROCHLORIDE - OXYCONTIN</u>					
020553 001	5266331	Oct 26, 2007	DP		
	5549912	Oct 26, 2007	DP		
<u>OXYCODONE HYDROCHLORIDE - OXYCONTIN</u>					
020553 002	5266331	Oct 26, 2007	DP		
	5549912	Oct 26, 2007	DP		

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnotes for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>OXYCODONE HYDROCHLORIDE - OXYCONTIN</u>					
020553 003	5266331	Oct 26, 2007	DP		
	5549912	Oct 26, 2007	DP		
<u>OXYCODONE HYDROCHLORIDE - OXYCONTIN</u>					
020553 004	5266331	Oct 26, 2007	DP		
	5549912	Oct 26, 2007	DP		
<u>OXYCODONE HYDROCHLORIDE - OXYCONTIN</u>					
020553 005	5266331	Oct 26, 2007	DP		
	5549912	Oct 26, 2007	DP		
<u>OXYCODONE HYDROCHLORIDE - OXYCONTIN</u>					
020553 006	5266331	Oct 26, 2007	DP		
	5508042	Apr 16, 2013		U-443	
	5549912	Oct 26, 2007	DP		
	5656295	Oct 26, 2007	DP	U-443	
<u>OXYCODONE HYDROCHLORIDE - OXYCONTIN</u>					
020553 007	5266331	Oct 26, 2007	DP		
	5508042	Apr 16, 2013		U-443	
	5549912	Oct 26, 2007	DP		
	5656295	Oct 26, 2007	DP	U-443	
<u>OXYCODONE HYDROCHLORIDE - OXYCONTIN</u>					
020553 008	5266331	Oct 26, 2007	DP		
	5508042	Apr 16, 2013		U-443	
	5549912	Oct 26, 2007	DP		
	5656295	Oct 26, 2007	DP	U-443	
<u>OXYMORPHONE HYDROCHLORIDE - OPANA</u>					
021611 001				NDF	Jun 22, 2009
<u>OXYMORPHONE HYDROCHLORIDE - OPANA</u>					
021611 002				NDF	Jun 22, 2009
<u>OXYMORPHONE HYDROCHLORIDE - OPANA ER</u>					
021610 001	5128143	Sep 19, 2008	DP	NDF	Jun 22, 2009
<u>OXYMORPHONE HYDROCHLORIDE - OPANA ER</u>					
021610 002	5128143	Sep 19, 2008	DP	NDF	Jun 22, 2009
<u>OXYMORPHONE HYDROCHLORIDE - OPANA ER</u>					
021610 003	5128143	Sep 19, 2008	DP	NDF	Jun 22, 2009
<u>OXYMORPHONE HYDROCHLORIDE - OPANA ER</u>					
021610 004	5128143	Sep 19, 2008	DP	NDF	Jun 22, 2009
<u>PALIPERIDONE - INVEGA</u>					
021999 001				>A> NCE	Dec 19, 2011
<u>PALIPERIDONE - INVEGA</u>					
021999 002				>A> NCE	Dec 19, 2011
<u>PALIPERIDONE - INVEGA</u>					
021999 003				>A> NCE	Dec 19, 2011
<u>PALIPERIDONE - INVEGA</u>					
021999 004				>A> NCE	Dec 19, 2011
<u>PAROXETINE HYDROCHLORIDE - PAXIL</u>					
020031 004	6133289	May 19, 2015		U-358	
<u>PAROXETINE HYDROCHLORIDE - PAXIL</u>					
020031 005	6133289	May 19, 2015		U-358	
<u>POLYETHYLENE GLYCOL 3350 - MIRALAX</u>					
022015 001				NP	Oct 06, 2009
<u>POSACONAZOLE - NOXAFIL</u>					
022003 001	5661151	Aug 26, 2014	DS DP	U-760	
	5703079	Dec 30, 2014	DS DP	U-760	
	6958337	Sep 25, 2020	DS DP	U-760	

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnotes for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>PRAMIPEXOLE DIHYDROCHLORIDE - MIRAPEX</u>					
020667 001	4843086	Jun 27, 2006	U-231	I-517	Nov 07, 2009
	6001861	Jan 16, 2018	U-784		
	6194445	Jan 16, 2018	U-784		
<u>PRAMIPEXOLE DIHYDROCHLORIDE - MIRAPEX</u>					
020667 002	4843086	Jun 27, 2006	U-231	I-517	Nov 07, 2009
	6001861	Jan 16, 2018	U-784		
	6194445	Jan 16, 2018	U-784		
<u>PRAMIPEXOLE DIHYDROCHLORIDE - MIRAPEX</u>					
020667 003	4843086	Jun 27, 2006	U-231	I-517	Nov 07, 2007
	6001861	Jan 16, 2018	U-784		
	6194445	Jan 16, 2018	U-784		
<u>PRAMIPEXOLE DIHYDROCHLORIDE - MIRAPEX</u>					
020667 004	4843086	Jun 27, 2006	U-231		
<u>PRAMIPEXOLE DIHYDROCHLORIDE - MIRAPEX</u>					
020667 005	4843086	Jun 27, 2006	U-231	I-517	Nov 07, 2009
	6001861	Jan 16, 2018	U-784		
	6194445	Jan 16, 2018	U-784		
<u>PRAMIPEXOLE DIHYDROCHLORIDE - MIRAPEX</u>					
020667 006	4843086	Jun 27, 2006	U-231	I-517	Nov 07, 2009
	6001861	Jan 16, 2018	U-784		
	6194445	Jan 16, 2018	U-784		
<u>PRAVASTATIN SODIUM - PRAVASTATIN SODIUM</u>					
076056 001				PC	Oct 21, 2006
<u>PRAVASTATIN SODIUM - PRAVASTATIN SODIUM</u>					
076056 002				PC	Oct 21, 2006
<u>PRAVASTATIN SODIUM - PRAVASTATIN SODIUM</u>					
076056 003				PC	Oct 21, 2006
<u>PREDNISOLONE SODIUM PHOSPHATE - ORAPRED ODT</u>					
021959 001	5178878	Jan 12, 2010	DP		
<u>PREDNISOLONE SODIUM PHOSPHATE - ORAPRED ODT</u>					
021959 002	5178878	Jan 12, 2010	DP		
<u>PREDNISOLONE SODIUM PHOSPHATE - ORAPRED ODT</u>					
021959 003	5178878	Jan 12, 2010	DP		
<u>QUETIAPINE FUMARATE - SEROQUEL</u>					
020639 001				I-503	Oct 20, 2009
<u>QUETIAPINE FUMARATE - SEROQUEL</u>					
020639 002				I-503	Oct 20, 2009
<u>QUETIAPINE FUMARATE - SEROQUEL</u>					
020639 003				I-503	Oct 20, 2009
<u>QUETIAPINE FUMARATE - SEROQUEL</u>					
020639 004				I-503	Oct 20, 2009
<u>QUETIAPINE FUMARATE - SEROQUEL</u>					
020639 005				I-503	Oct 20, 2009
<u>QUETIAPINE FUMARATE - SEROQUEL</u>					
020639 006	4879288	Sep 26, 2011	DS DP U-550	I-503	Oct 20, 2009
<u>QUETIAPINE FUMARATE - SEROQUEL</u>					
020639 007	4879288	Sep 26, 2011	DS DP U-550	I-503	Oct 20, 2009
<u>RALOXIFENE HYDROCHLORIDE - EVISTA</u>					
020815 001	RE38968	Jul 28, 2012	U-662		
	RE38968	Jul 28, 2012	U-657		
	RE39049	Jul 28, 2012	U-662		
	RE39049	Jul 28, 2012	U-657		
	RE39050	Mar 02, 2014	U-662		
	RE39050	Mar 02, 2014	U-657		

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnotes for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>RAMIPRIL - ALTACE</u>					
019901 001	5061722	Oct 19, 2008			
<u>RANOLAZINE - RANEXA</u>					
021526 002	4567264	May 18, 2007	DS	NCE	Jan 27, 2011
	6303607	May 27, 2019		U-705	
	6369062	May 27, 2019	DP		
	6479496	May 27, 2019		U-705	
	6503911	May 27, 2019	DP		
	6525057	May 27, 2019		U-705	
	6562826	May 27, 2019		U-705	
	6617328	May 27, 2019	DP		
	6620814	May 27, 2019		U-705	
	6852724	May 27, 2019		U-705	
	6864258	May 27, 2019		U-705	
<u>RASAGILINE MESYLATE - AZILECT</u>					
021641 001	5387612	Feb 07, 2012		U-219	NCE
	5453446	Feb 07, 2012		U-219	
	5457133	Feb 07, 2012	DS	DP	
	5532415	Jul 02, 2013	DS		
	5786390	Feb 07, 2012		DP	
	6126968	Sep 18, 2016		DP	
<u>RASAGILINE MESYLATE - AZILECT</u>					
021641 002	5387612	Feb 07, 2012		U-219	NCE
	5453446	Feb 07, 2012		U-219	
	5457133	Feb 07, 2012	DS	DP	
	5532415	Jul 02, 2013	DS		
	5786390	Feb 07, 2012		DP	
	6126968	Sep 18, 2016		DP	
<u>RIFAXIMIN - XIFAXAN</u>					
021361 001	7045620	May 22, 2024	DS		
<u>RISEDRONATE SODIUM - ACTONEL</u>					
020835 001				M-52	Jan 24, 2009
<u>RISEDRONATE SODIUM - ACTONEL</u>					
020835 002				M-52	Jan 24, 2009
<u>RISEDRONATE SODIUM - ACTONEL</u>					
020835 003	5583122	Dec 10, 2013	DS	DP	I-309
	5583122	Dec 10, 2013	DS	DP	M-52
	6096342	Nov 22, 2011		DP	
<u>RISPERIDONE - RISPERDAL</u>					
020272 001				I-509	Oct 06, 2009
<u>RISPERIDONE - RISPERDAL</u>					
020272 002				I-509	Oct 06, 2009
<u>RISPERIDONE - RISPERDAL</u>					
020272 003				I-509	Oct 06, 2009
<u>RISPERIDONE - RISPERDAL</u>					
020272 004				I-509	Oct 06, 2009
<u>RISPERIDONE - RISPERDAL</u>					
020272 007				I-509	Oct 06, 2009
<u>RISPERIDONE - RISPERDAL</u>					
020272 008				I-509	Oct 06, 2009
<u>RISPERIDONE - RISPERDAL</u>					
020588 001	RE39181	Jul 11, 2014		DP	I-509
<u>RISPERIDONE - RISPERDAL</u>					
021444 001				I-509	Oct 06, 2009
<u>RISPERIDONE - RISPERDAL</u>					
021444 002				I-509	Oct 06, 2009

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnotes for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>RISPERIDONE - RISPERDAL</u>					
021444 003				I-509	Oct 06, 2006
<u>RISPERIDONE - RISPERDAL</u>					
021444 004				I-509	Oct 06, 2006
<u>RISPERIDONE - RISPERDAL</u>					
021444 005				I-509	Oct 06, 2009
<u>RITONAVIR - NORVIR</u>					
020659 001	>A> 5948436	Sep 13, 2013	DP		
	>A> 5948436*PED	Mar 13, 2014			
<u>RITONAVIR - NORVIR</u>					
020945 001	>A> 5948436	Sep 13, 2013	DP		
	>A> 5948436*PED	Mar 13, 2014			
	>A> 7141593	May 22, 2020	DP		
	>A> 7141593*PED	Nov 22, 2020			
<u>RIVASTIGMINE TARTRATE - EXELON</u>					
020823 003	4948807	Aug 14, 2012	DS	U-322	
<u>RIVASTIGMINE TARTRATE - EXELON</u>					
020823 004	4948807	Aug 14, 2012	DS	U-322	
<u>RIVASTIGMINE TARTRATE - EXELON</u>					
020823 005	4948807	Aug 14, 2012	DS	U-322	
<u>RIVASTIGMINE TARTRATE - EXELON</u>					
020823 006	4948807	Aug 14, 2012	DS	U-322	
<u>RIVASTIGMINE TARTRATE - EXELON</u>					
021025 001	4948807	Aug 14, 2012	DS	U-322	
<u>ROCURONIUM BROMIDE - ZEMURON</u>					
020214 003	4894369	Apr 13, 2008			
<u>SALMETEROL XINAFOATE - SEREVENT</u>					
020236 001	4992474	Feb 12, 2008			
	4992474*PED	Aug 12, 2008			
	5126375	Feb 12, 2008			
	5126375*PED	Aug 12, 2008			
	5225445	Feb 12, 2008		U-182	
	5225445*PED	Aug 12, 2008			
<u>SALMETEROL XINAFOATE - SEREVENT</u>					
020692 001	4992474	Feb 12, 2008			
	4992474*PED	Aug 12, 2008			
	5126375	Feb 12, 2008			
	5126375*PED	Aug 12, 2008			
	5225445	Feb 12, 2008		U-211	
	5225445*PED	Aug 12, 2008			
	6536427	Mar 01, 2011	DP		
<u>SELEGILINE - EMSAM</u>					
021336 001	7070808	May 10, 2018	DS	DP	
	RE34579	Aug 18, 2007	DS	DP	U-711
<u>SELEGILINE - EMSAM</u>					
021336 002	RE34579	Aug 18, 2007	DS	DP	U-711
<u>SELEGILINE - EMSAM</u>					
021336 003	RE34579	Aug 18, 2007	DS	DP	U-711
<u>SELEGILINE HYDROCHLORIDE - ZELAPAR</u>					
021479 001	5648093	Jul 15, 2014		DP	
	6423342	Mar 01, 2016		DP	
<u>SERTACONAZOLE NITRATE - ERTACZO</u>					
021385 001	>A> 5135943	May 31, 2014	DS	DP	U-786
<u>SERTRALINE HYDROCHLORIDE - SERTRALINE HYDROCHLORIDE</u>					
075719 001				PC	Feb 06, 2007

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnotes for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>SERTRALINE HYDROCHLORIDE - SERTRALINE HYDROCHLORIDE</u>					
075719 002				PC	Feb 06, 2007
<u>SERTRALINE HYDROCHLORIDE - SERTRALINE HYDROCHLORIDE</u>					
075719 003				PC	Feb 06, 2007
<u>SERTRALINE HYDROCHLORIDE - SERTRALINE HYDROCHLORIDE</u>					
076934 001				PC	Feb 03, 2007
<u>SERTRALINE HYDROCHLORIDE - ZOLOFT</u>					
020990 001	6727283	Oct 11, 2019	DP U-580		
	6727283*PED	Apr 11, 2020			
	7067555	Nov 10, 2019	DP		
	7067555*PED	May 10, 2020			
<u>SEVELAMER HYDROCHLORIDE - RENAGEL</u>					
021179 001	7014846	Aug 11, 2013	DP U-246		
<u>SEVELAMER HYDROCHLORIDE - RENAGEL</u>					
021179 002	7014846	Aug 11, 2013	DP U-246		
<u>SIMVASTATIN - SIMVASTATIN</u>					
076052 001				PC	Dec 20, 2006
<u>SIMVASTATIN - SIMVASTATIN</u>					
076052 002				PC	Dec 20, 2006
<u>SIMVASTATIN - SIMVASTATIN</u>					
076052 003				PC	Dec 20, 2006
<u>SIMVASTATIN - SIMVASTATIN</u>					
076052 004				PC	Dec 20, 2006
<u>SIMVASTATIN - SIMVASTATIN</u>					
076285 005				PC	Dec 20, 2006
<u>SIROLIMUS - RAPAMUNE</u>					
021083 001	5100899	Jul 07, 2013	U-290		
	5100899*PED	Jan 07, 2014			
<u>SIROLIMUS - RAPAMUNE</u>					
021110 001	5100899	Jul 07, 2013	U-290		
	5100899*PED	Jan 07, 2014			
<u>SIROLIMUS - RAPAMUNE</u>					
021110 002	5100899	Jul 07, 2013	U-290		
	5100899*PED	Jan 07, 2014			
<u>SIROLIMUS - RAPAMUNE</u>					
021110 003	5100899	Jul 07, 2013	U-290		
	5100899*PED	Jan 07, 2014			
<u>SITAGLIPTIN PHOSPHATE - JANUVIA</u>					
021995 001	6303661	Apr 24, 2017	U-774	NCE	Oct 16, 2011
	6699871	Jul 26, 2022	DS DP U-774		
	6890898	Feb 02, 2019	U-775		
	7078381	Feb 02, 2019	U-775		
	7125873	Jul 26, 2022	U-775		
<u>SITAGLIPTIN PHOSPHATE - JANUVIA</u>					
021995 002	6303661	Apr 24, 2017	U-774	NCE	Oct 16, 2011
	6699871	Jul 26, 2022	DS DP U-774		
	6890898	Feb 02, 2019	U-775		
	7078381	Feb 02, 2019	U-775		
	7125873	Jul 26, 2022	U-775		
<u>SITAGLIPTIN PHOSPHATE - JANUVIA</u>					
021995 003	6303661	Apr 24, 2017	U-774	NCE	Oct 16, 2011
	6699871	Jul 26, 2022	DS DP U-774		
	6890898	Feb 02, 2019	U-775		
	7078381	Feb 02, 2019	U-775		
	7125873	Jul 26, 2022	U-775		

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnotes for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
SODIUM PHOSPHATE, DIBASIC, ANHYDROUS; SODIUM PHOSPHATE, MONOBASIC, MONOHYDRATE - OSMOPREP					
021892 001	5616346	May 18, 2013	DP U-715	NP	Mar 16, 2009
SOMATROPIN RECOMBINANT - GENOTROPIN					
020280 006	4968299	Jun 28, 2008	DP	I-496	Apr 27, 2009
SOMATROPIN RECOMBINANT - GENOTROPIN					
020280 007	4968299	Jun 28, 2008	DP	I-496	Apr 27, 2009
SOMATROPIN RECOMBINANT - GENOTROPIN PRESERVATIVE FREE					
020280 001	5435076	Apr 16, 2013	DP	I-496	Apr 27, 2009
	5716338	Feb 10, 2015	DP		
SOMATROPIN RECOMBINANT - GENOTROPIN PRESERVATIVE FREE					
020280 002	5435076	Apr 16, 2013	DP	I-496	Apr 27, 2009
	5716338	Feb 10, 2015	DP		
SOMATROPIN RECOMBINANT - GENOTROPIN PRESERVATIVE FREE					
020280 003	5435076	Apr 16, 2013	DP	I-496	Apr 27, 2009
	5716338	Feb 10, 2015	DP		
SOMATROPIN RECOMBINANT - GENOTROPIN PRESERVATIVE FREE					
020280 004				I-496	Apr 27, 2009
SOMATROPIN RECOMBINANT - GENOTROPIN PRESERVATIVE FREE					
020280 005	5435076	Apr 16, 2013	DP	I-496	Apr 27, 2009
	5716338	Feb 10, 2015	DP		
SOMATROPIN RECOMBINANT - GENOTROPIN PRESERVATIVE FREE					
020280 008	5435076	Apr 16, 2013	DP	I-496	Apr 27, 2009
	5716338	Feb 10, 2015	DP		
SOMATROPIN RECOMBINANT - GENOTROPIN PRESERVATIVE FREE					
020280 009	5435076	Apr 16, 2013	DP	I-496	Apr 27, 2009
	5716338	Feb 10, 2015	DP		
SOMATROPIN RECOMBINANT - GENOTROPIN PRESERVATIVE FREE					
020280 010	5435076	Apr 16, 2013	DP	I-496	Apr 27, 2009
	5716338	Feb 10, 2015	DP		
SOMATROPIN RECOMBINANT - GENOTROPIN PRESERVATIVE FREE					
020280 011	5435076	Apr 16, 2013	DP	I-496	Apr 27, 2009
	5716338	Feb 10, 2015	DP		
SOMATROPIN RECOMBINANT - GENOTROPIN PRESERVATIVE FREE					
020280 012	5435076	Apr 16, 2013	DP	I-496	Apr 27, 2009
	5716338	Feb 10, 2015	DP		
SOMATROPIN RECOMBINANT - GENOTROPIN PRESERVATIVE FREE					
020280 013	5435076	Apr 16, 2013	DP	I-496	Apr 27, 2009
	5716338	Feb 10, 2015	DP		
SOMATROPIN RECOMBINANT - HUMATROPE					
019640 004				I-518	Nov 01, 2009
SOMATROPIN RECOMBINANT - HUMATROPE					
019640 005				I-518	Nov 01, 2009
SOMATROPIN RECOMBINANT - HUMATROPE					
019640 006				I-518	Nov 01, 2009
SOMATROPIN RECOMBINANT - HUMATROPE					
019640 007				I-518	Nov 01, 2009
SOMATROPIN RECOMBINANT - OMNITROPE					
021426 001				NP	May 30, 2009
SOMATROPIN RECOMBINANT - OMNITROPE					
021426 002				NP	May 30, 2009
SORAFENIB TOSYLATE - NEXAVAR					
021923 001				ODE	Dec 20, 2012
STAVUDINE - ZERIT XR					
021453 001	7135465	Feb 18, 2023	DP U-167		
	7135465*PED	Aug 18, 2023			

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnotes for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>STAVUDINE - ZERIT XR</u>					
021453 002	7135465	Feb 18, 2023	DP U-167		
	7135465*PED	Aug 18, 2023			
<u>STAVUDINE - ZERIT XR</u>					
021453 003	7135465	Feb 18, 2023	DP U-167		
	7135465*PED	Aug 18, 2023			
<u>STAVUDINE - ZERIT XR</u>					
021453 004	7135465	Feb 18, 2023	DP U-167		
	7135465*PED	Aug 18, 2023			
<u>SUMATRIPTAN SUCCINATE - IMITREX STATDOSE</u>					
020080 003	4816470	Dec 28, 2006	U-72		
	4816470*PED	Jun 28, 2007			
	5037845	Aug 06, 2008	U-72		
	5037845*PED	Feb 06, 2009			
<u>SUNITINIB MALATE - SUTENT</u>					
021938 001	6573293	Feb 15, 2021	DS DP U-703	NCE	Jan 26, 2011
	7125905	Feb 15, 2021	DS DP		
<u>SUNITINIB MALATE - SUTENT</u>					
021938 002	6573293	Feb 15, 2021	DS DP U-703	NCE	Jan 26, 2011
	7125905	Feb 15, 2021	DS DP		
<u>SUNITINIB MALATE - SUTENT</u>					
021938 003	6573293	Feb 15, 2021	DS DP U-703	NCE	Jan 26, 2011
	7125905	Feb 15, 2021	DS DP		
<u>TACROLIMUS - PROGRAF</u>					
050708 001				ODE	Mar 29, 2013
<u>TACROLIMUS - PROGRAF</u>					
050708 002				ODE	Mar 29, 2013
<u>TACROLIMUS - PROGRAF</u>					
050708 003				ODE	Mar 29, 2013
<u>TACROLIMUS - PROGRAF</u>					
050709 001				ODE	Mar 29, 2013
<u>TELBIVUDINE - TYZEKA</u>					
022011 001	6395716	Aug 10, 2019	U-782	NCE	Oct 25, 2011
	6444652	Aug 10, 2019	U-782		
	6566344	Aug 10, 2019	U-782		
	6569837	Aug 10, 2019	U-782		
<u>TEMOZOLOMIDE - TEMODAR</u>					
021029 005				I-450 ODE ODE PED	Mar 15, 2008 Mar 15, 2012 Aug 11, 2006 Feb 11, 2007
<u>TEMOZOLOMIDE - TEMODAR</u>					
021029 006				I-450 ODE ODE PED	Mar 15, 2008 Mar 15, 2012 Aug 11, 2006 Feb 11, 2007

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnotes for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>TERBINAFAINE - LAMISIL</u>					
020846 001	4755534	Dec 30, 2006		U-445	
	4755534*PED	Jun 30, 2007			
	5681849	Oct 28, 2014	DP		
	5681849*PED	Apr 28, 2015			
	5856355	May 18, 2012	DP	U-502	
	5856355	May 18, 2012	DP	U-504	
	5856355	May 18, 2012	DP	U-540	
	5856355*PED	Nov 18, 2012			
	6005001	May 18, 2012	DP	U-502	
	6005001	May 18, 2012	DP	U-504	
	6005001	May 18, 2012	DP	U-540	
	6005001*PED	Nov 18, 2012			
	6121314	May 18, 2012	DP	U-504	
	6121314	May 18, 2012	DP	U-540	
	6121314	May 18, 2012	DP	U-502	
	6121314*PED	Nov 18, 2012			
<u>TERBINAFAINE HYDROCHLORIDE - LAMISIL</u>					
020192 001	4755534	Dec 30, 2006		U-73	
	4755534*PED	Jun 30, 2007			
<u>TERBINAFAINE HYDROCHLORIDE - LAMISIL</u>					
020539 001	4755534	Dec 30, 2006		U-73	
	4755534*PED	Jun 30, 2007			
<u>TERBINAFAINE HYDROCHLORIDE - LAMISIL</u>					
020749 001	4755534	Dec 30, 2006			
	4755534*PED	Jun 30, 2007			
	6121314	May 18, 2012		U-502	
	6121314*PED	Nov 18, 2012			
<u>TERBINAFAINE HYDROCHLORIDE - LAMISIL</u>					
020980 001	4755534	Dec 30, 2006		U-73	
	4755534*PED	Jun 30, 2007			
<u>TERBINAFAINE HYDROCHLORIDE - LAMISIL AT</u>					
021124 001	4755534	Dec 30, 2006		U-73	
	4755534*PED	Jun 30, 2007			
	5681849	Oct 28, 2014			
	5681849*PED	Apr 28, 2015			
	6121314	May 18, 2012		U-504	
	6121314*PED	Nov 18, 2012			
<u>TERBINAFAINE HYDROCHLORIDE - LAMISIL AT</u>					
021124 002	4755534	Dec 30, 2006		U-73	
	4755534*PED	Jun 30, 2007			
	5681849	Oct 28, 2014			
	5681849*PED	Apr 28, 2015			
	6121314	May 18, 2012		U-504	
	6121314*PED	Nov 18, 2012			
<u>TERIPARATIDE RECOMBINANT HUMAN - FORTEO</u>					
021318 001	6977077	Aug 19, 2019		U-597	
	7144861	Dec 08, 2018	DP		

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnotes for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>THALIDOMIDE - THALOMID</u>					
020785 001	>A> 5629327	May 13, 2014	U-731	ODE	May 25, 2013
	6045501	Aug 28, 2018	U-731		
	6045501	Aug 28, 2018	U-371		
	6235756	Mar 01, 2013	U-731		
	6315720	Oct 23, 2020	U-442		
	6315720	Oct 23, 2020	U-731		
	6561976	Aug 28, 2018	U-371		
	6561976	Aug 28, 2018	U-731		
	6561977	Oct 23, 2020	U-371		
	6561977	Oct 23, 2020	U-731		
	6755784	Oct 23, 2020	U-731		
	6755784	Oct 23, 2020	U-371		
	6869399	Oct 23, 2020	U-732		
	6869399	Oct 23, 2020	U-733		
	6869399	Oct 23, 2020	U-731		
	6869399	Oct 23, 2020	U-371		
	6908432	Aug 28, 2018	U-371		
	6908432	Aug 28, 2018	U-731		
	>A> 7141018	Oct 23, 2020	U-732		
	>A> 7141018	Oct 23, 2020	U-371		
	>A> 7141018	Oct 23, 2020	U-731		
	>A> 7141018	Oct 23, 2020	U-733		
<u>THALIDOMIDE - THALOMID</u>					
020785 002	>A> 5629327	May 13, 2014	U-731	ODE	May 25, 2013
	6045501	Aug 28, 2018	U-731		
	6045501	Aug 28, 2018	U-371		
	6235756	Mar 01, 2013	U-731		
	6315720	Oct 23, 2020	U-442		
	6315720	Oct 23, 2020	U-731		
	6561976	Aug 28, 2018	U-371		
	6561976	Aug 28, 2018	U-731		
	6561977	Oct 23, 2020	U-371		
	6561977	Oct 23, 2020	U-731		
	6755784	Oct 23, 2020	U-731		
	6755784	Oct 23, 2020	U-371		
	6869399	Oct 23, 2020	U-733		
	6869399	Oct 23, 2020	U-732		
	6869399	Oct 23, 2020	U-731		
	6869399	Oct 23, 2020	U-371		
	6908432	Aug 28, 2018	U-371		
	6908432	Aug 28, 2018	U-731		
	>A> 7141018	Oct 23, 2020	U-733		
	>A> 7141018	Oct 23, 2020	U-371		
	>A> 7141018	Oct 23, 2020	U-731		
	>A> 7141018	Oct 23, 2020	U-732		

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnotes for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>THALIDOMIDE - THALOMID</u>					
020785 003	>A> 5629327	May 13, 2014	U-731	ODE	May 25, 2013
	6045501	Aug 28, 2018	U-731		
	6045501	Aug 28, 2018	U-371		
	6235756	Mar 01, 2013	U-731		
	6315720	Oct 23, 2020	U-442		
	6315720	Oct 23, 2020	U-731		
	6561976	Aug 28, 2018	U-371		
	6561976	Aug 28, 2018	U-731		
	6561977	Oct 23, 2020	U-371		
	6561977	Oct 23, 2020	U-731		
	6755784	Oct 23, 2020	U-731		
	6755784	Oct 23, 2020	U-371		
	6869399	Oct 23, 2020	U-732		
	6869399	Oct 23, 2020	U-733		
	6869399	Oct 23, 2020	U-731		
	6869399	Oct 23, 2020	U-371		
	6908432	Aug 28, 2018	U-371		
	6908432	Aug 28, 2018	U-731		
	>A> 7141018	Oct 23, 2020	U-733		
	>A> 7141018	Oct 23, 2020	U-371		
	>A> 7141018	Oct 23, 2020	U-731		
	>A> 7141018	Oct 23, 2020	U-732		
<u>THYROTROPIN ALFA - THYROGEN</u>					
020898 001				M-53	Jan 23, 2009
<u>TIOTROPIUM BROMIDE MONOHYDRATE - SPIRIVA</u>					
021395 001	7070800	Jan 22, 2022	DP U-566		
	RE38912	Oct 11, 2021	DP		
<u>TOPIRAMATE - TOPAMAX</u>					
020505 001	7018983	Oct 13, 2015	U-723		
<u>TOPIRAMATE - TOPAMAX</u>					
020505 002	7018983	Oct 13, 2015	U-723		
<u>TOPIRAMATE - TOPAMAX</u>					
020505 003	7018983	Oct 13, 2015	U-723		
<u>TOPIRAMATE - TOPAMAX</u>					
020505 004	7018983	Oct 13, 2015	U-723		
<u>TOPIRAMATE - TOPAMAX</u>					
020505 005	7018983	Oct 13, 2015	U-723		
<u>TOPIRAMATE - TOPAMAX</u>					
020505 006	7018983	Oct 13, 2015	U-723		
<u>TOPIRAMATE - TOPAMAX SPRINKLE</u>					
020844 001	7018983	Oct 13, 2015	U-723		
	7125560	Mar 01, 2019	U-766		
<u>TOPIRAMATE - TOPAMAX SPRINKLE</u>					
020844 002	7018983	Oct 13, 2015	U-723		
	7125560	Mar 01, 2019	U-766		
<u>TOPIRAMATE - TOPAMAX SPRINKLE</u>					
020844 003	7018983	Oct 13, 2015	U-723		
	7125560	Mar 01, 2019	U-766		
<u>TOPOTECAN HYDROCHLORIDE - HYCAMTIN</u>					
020671 001	5004758	May 28, 2010	DS DP U-741		
<u>TRAVOPROST - TRAVATAN Z</u>					
021994 001	5510383	Aug 03, 2013	DP U-383		
	5889052	Dec 02, 2014	DP U-383		
	6503497	May 06, 2012	DP		
	6849253	May 06, 2012	DP		

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnotes for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>TREPROSTINIL SODIUM - REMODULIN</u>					
021272 001	5153222	Oct 06, 2014	U-455		
<u>TREPROSTINIL SODIUM - REMODULIN</u>					
021272 002	5153222	Oct 06, 2014	U-455		
<u>TREPROSTINIL SODIUM - REMODULIN</u>					
021272 003	5153222	Oct 06, 2014	U-455		
<u>TREPROSTINIL SODIUM - REMODULIN</u>					
021272 004	5153222	Oct 06, 2014	U-455		
<u>URSODIOL - URSO FORTE</u>					
020675 002	4859660	Nov 19, 2007	U-740		
<u>VALDECOXIB - BEXTRA</u>					
021341 002	7135489	Aug 12, 2017	DS DP		
<u>VALDECOXIB - BEXTRA</u>					
021341 003	7135489	Aug 12, 2017	DS DP		
<u>VARENICLINE TARTRATE - CHANTIX</u>					
021928 001	6410550	Nov 13, 2018	DS DP	U-56	NCE
	6890927	May 06, 2022	DS DP	U-56	May 10, 2011
<u>VARENICLINE TARTRATE - CHANTIX</u>					
021928 002	6410550	Nov 13, 2018	DS DP	U-56	NCE
	6890927	May 06, 2022	DS DP	U-56	May 10, 2011
<u>VENLAFAXINE HYDROCHLORIDE - VENLAFAXINE HYDROCHLORIDE</u>					
076690 001				PC	Jan 30, 2007
<u>VENLAFAXINE HYDROCHLORIDE - VENLAFAXINE HYDROCHLORIDE</u>					
076690 002				PC	Jan 30, 2007
<u>VENLAFAXINE HYDROCHLORIDE - VENLAFAXINE HYDROCHLORIDE</u>					
076690 003				PC	Jan 30, 2007
<u>VENLAFAXINE HYDROCHLORIDE - VENLAFAXINE HYDROCHLORIDE</u>					
076690 004				PC	Jan 30, 2007
<u>VENLAFAXINE HYDROCHLORIDE - VENLAFAXINE HYDROCHLORIDE</u>					
076690 005				PC	Jan 30, 2007
<u>VORINOSTAT - ZOLINZA</u>					
021991 001	6087367	Oct 04, 2011	U-776	NCE	Oct 06, 2011
	RE38506	Nov 29, 2011	DS DP	>A> ODE	Oct 06, 2013
<u>ZANAMIVIR - RELENZA</u>					
021036 001	5648379	Jul 15, 2014	U-722	I-491	Mar 29, 2009
	5648379	Jul 15, 2014	U-721		
	5648379	Jul 15, 2014	U-274		
	6294572	Dec 15, 2014	DS DP		
<u>ZIPRASIDONE HYDROCHLORIDE - GEODON</u>					
020825 001	4831031	Mar 02, 2012	DS DP	U-720	I-492
<u>ZIPRASIDONE HYDROCHLORIDE - GEODON</u>					
020825 002	4831031	Mar 02, 2012	DS DP	U-720	I-492
<u>ZIPRASIDONE HYDROCHLORIDE - GEODON</u>					
020825 003	4831031	Mar 02, 2012	DS DP	U-720	I-492
<u>ZIPRASIDONE HYDROCHLORIDE - GEODON</u>					
020825 004	4831031	Mar 02, 2012	DS DP	U-720	I-492
<u>ZIPRASIDONE HYDROCHLORIDE - GEODON</u>					
021483 001	4831031	Mar 02, 2012	DS DP	U-720	I-492
	5312925	Sep 01, 2012	DS DP	U-720	
	6150366	May 27, 2019	DP	U-719	
	6245766	Dec 18, 2018		U-601	
<u>ZIPRASIDONE MESYLATE - GEODON</u>					
020919 001	4831031	Mar 02, 2012	DS DP	U-720	

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST

See report footnotes for information regarding report content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE			PATENT CODES		EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE	
<u>ZOLEDRONIC ACID - ZOMETA</u>									
021223 001	4939130	Sep 02, 2012	DS	DP	U-53				
<u>ZOLEDRONIC ACID - ZOMETA</u>									
021223 002	4939130	Sep 02, 2012	DS	DP	U-53				
<u>ZOLPIDEM TARTRATE - AMBIEN</u>									
019908 001	4382938	Oct 21, 2006			U-74				
	4382938*PED	Apr 21, 2007							
<u>ZOLPIDEM TARTRATE - AMBIEN</u>									
019908 002	4382938	Oct 21, 2006			U-74				
	4382938*PED	Apr 21, 2007							
<u>ZOLPIDEM TARTRATE - AMBIEN CR</u>									
021774 001	4382938	Oct 21, 2006	DS		U-682	NDF	Sep 02, 2008		
	4382938*PED	Apr 21, 2007				PED	Mar 02, 2009		
	6514531	Dec 01, 2019		DP					
	6514531*PED	Jun 01, 2020							
<u>ZOLPIDEM TARTRATE - AMBIEN CR</u>									
021774 002	4382938	Oct 21, 2006	DS		U-682	NDF	Sep 02, 2008		
	4382938*PED	Apr 21, 2007				PED	Mar 02, 2009		
	6514531	Dec 01, 2019		DP					
	6514531*PED	Jun 01, 2020							

Footnotes:

- Patents are published upon receipt by the Orange Book Staff and may not reflect the official receipt date as described in 21 CFR 314.53(d)(5).
- Patents submitted on FDA Form 3542 and listed after August 18, 2003 will have one to three patent codes indicating specific patent claims as submitted by the sponsor:
DS = Drug Substance claim
DP = Drug Product claim
U and number = Method of Use claim (may be multiple). Specific Method of use claims are listed at <http://www.fda.gov/cder/orange/patex.htm>
- Patents listed prior to August 18, 2003 are flagged with method of use claims only as applicable and submitted by the sponsor. They may not be flagged with respect to other claims which may apply.
- *PED and PED represent pediatric exclusivity. Patents with pediatric exclusivity granted after August 18, 2003 will be indicated with *PED as was done prior to August 18, 2003. Patents with *PED added after August 18, 2003 will not contain any information relative to the patent itself other than the *PED extension. Information related specifically to the patent will be conveyed on the original patent only.
- *** U.S. Patent Nos. RE 36481 and RE 36520 were relisted for Zocor (NDA 19-766) pursuant to the decision and related order in *Ranbaxy Labs. v. Leavitt*, No. 05-1838 (D.D.C. April 30, 2006). The '481 and '520 patents remained listed in Approved Drug Products with Therapeutic Equivalence Evaluations until any applicable periods of exclusivity pursuant to section 505(j)(5)(B)(iv) of the Federal Food, Drug, and Cosmetic Act were triggered and run. For additional information on this matter, please refer to Docket Nos. 2005P-0008 and 2005P-0046. Patents were subsequently delisted in the December 2006 Orange Book update as the exclusivity periods have triggered and run to expiration.

PATENT AND EXCLUSIVITY TERMS

Due to space limitations in the patent and exclusivity columns, abbreviations and references have been developed. Refer to the APPROVED DRUG PRODUCTS WITH THERAPEUTIC EQUIVALENCE EVALUATIONS, 25th Edition for a full listing of patent and exclusivity terms (Abbreviations, Dosing Schedule, Indications, and Patent Use Codes).

The current complete list of patent terms is available at
<http://www.accessdata.fda.gov/scripts/cder/ob/docs/pattermsall.cfm>

The current complete list of exclusivity terms is available at
<http://www.accessdata.fda.gov/scripts/cder/ob/docs/excltermsall.cfm>