

APPROVED DRUG PRODUCTS

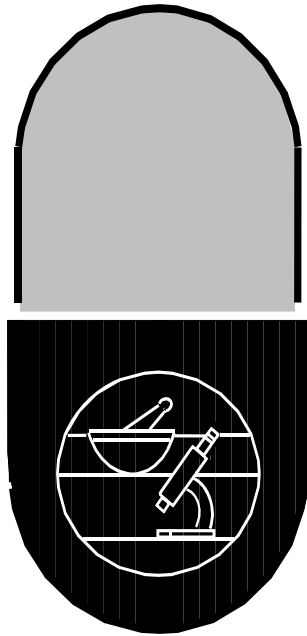
With Therapeutic Equivalence Evaluations



The "Orange Book"

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**CUMULATIVE
SUPPLEMENT 08
August 2009**



**APPROVED
DRUG PRODUCTS**

**WITH
THERAPEUTIC EQUIVALENCE EVALUATIONS**

29th EDITION

**Department of Health and Human Services
Food and Drug Administration
Center for Drug Evaluation and Research
Office of Generic Drugs**

2009

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

**APPROVED DRUG PRODUCTS
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THERAPEUTIC EQUIVALENCE EVALUATIONS**

29th EDITION

Cumulative Supplement 8

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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	vi
1.6 Cumulative Supplement Legend	vii
 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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29th EDITION

**CUMULATIVE SUPPLEMENT 8
August 2009**

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, 29th 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 29th Edition List will then be added to the "Discontinued Drug Product List" appearing in the 30th 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.

Every effort is made to ensure the Cumulative Supplement is current and accurate. Applicant holders are requested to inform the FDA Orange Book Staff (OBS) of any changes or corrections. The OBS can be contacted by email at drugproducts@fda.hhs.gov. Send Changes by FAX: 240-276-8974; mail to:

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

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>
ABLE LABORATORIES INC (ABLE)	IVAX PHARMACEUTICALS INC SUB TEVA PHARMACEUTICALS USA (IVAX SUB TEVA PHARM)
AXIOM PHARMACEUTICAL CORP (AXIOM PHARM)	IVAX PHARMACEUTICALS INC SUB TEVA PHARMACEUTICALS USA (IVAX SUB TEVA PHARM)
DABUR ONCOLOGY PLC (DABUR ONCOLOGY PLC)	FRESENIUS KABI ONCOLOGY PLC (FRESENIUS KABI ONCOL)
HALSEY DRUG CO INC (HALSEY)	IVAX PHARMACEUTICALS INC SUB TEVA PHARMACEUTICALS USA (IVAX SUB TEVA PHARM)
IVAX PHARMACEUTICALS INC SUB TEVA PHARMACEUTICALS (IVAX PAHRMS)	IVAX PHARMACEUTICALS INC SUB TEVA PHARMACEUTICALS USA (IVAX SUB TEVA PHARM)

NORTON WATERFORD LTD
(NORTON WATERFORD)

IVAX PHARMACEUTICALS INC SUB TEVA
PHARMACEUTICALS USA
(IVAX SUB TEVA PHARM)

TORPHARM INC
(TORPHARM)

APOTEX INC
(APOTEX)

ZENITH GOLDLINE
(ZENTH GOLDLINE)

IVAX PHARMACEUTICALS INC SUB TEVA
PHARMACEUTICALS USA
(IVAX SUB TEVA PHARM)

1.4 AVAILABILITY OF THE EDITION

Since 1997, the Electronic Orange Book Query (EOBQ) <http://www.accessdata.fda.gov/scripts/cder/ob/default.cfm>, 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 Publications. 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://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/Drugs/InformationOnDrugs/ucm086229.htm>. There are ASCII text files of the Orange Book drug product, patent, and exclusivity data at <http://www.fda.gov/Drugs/InformationOnDrugs/ucm129689.htm>. The drug product text files are provided in 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 2008) refers to the products in the Prescription Drug Product List. For each three-month period, a column of quarterly data is added which incorporates counts of product activity from the previous quarter(s) with those in the baseline count.

DEFINITIONS

Drug Product

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

New Molecular Entity

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

REPORT OF COUNTS FOR THE PRESCRIPTION DRUG PRODUCT LIST COUNTS CUMULATIVE BY QUARTER

<u>CATEGORIES COUNTED</u>	<u>DEC 2008</u>	<u>MAR 2009</u>	<u>JUN 2009</u>	<u>SEPT 2009</u>
DRUG PRODUCTS LISTED	12751	12910	13027	
SINGLE SOURCE	2433 (19.1%)	2449 (19.0%)	2451 (18.8%)	
MULTISOURCE	10229 (80.2%)	10372 (80.3%)	10487 (80.5%)	
THERAPEUTICALLY EQUIVALENT	10072 (79.0%)	10216 (79.1%)	10333 (79.3%)	
NOT THERAPEUTICALLY EQUIVALENT	157 (1.2%)	156 (1.2%)	154 (1.2%)	
EXCEPTIONS ¹	89 (0.7%)	89 (0.7%)	89 (0.7%)	
NEW MOLECULAR ENTITIES APPROVED	15	5	6	
NUMBER OF APPLICANTS	719	724	732	

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

1.6 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 >D>. 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 - 28TH EDITION

RX DRUG PRODUCT LIST - CUMULATIVE SUPPLEMENT 8 - August 2009

1-1

ACARBOSE

TABLET; ORAL

ACARBOSE

AB	IMPAX LABS	25MG	N78441 001	May 14, 2009	May	NEWA
AB		50MG	N78441 002	May 14, 2009	May	NEWA
AB		100MG	N78441 003	May 14, 2009	May	NEWA

PRECOSE

>D>	AB	BAYER HLTHCARE	25MG	N20482 004	May 29, 1997	Aug	CRLD
>A>	AB	+	25MG	N20482 004	May 29, 1997	Aug	CRLD

ACETAMINOPHEN; BUTALBITAL; CAFFEINE

TABLET; ORAL

BUTALBITAL, ACETAMINOPHEN AND CAFFEINE

@	MUTUAL PHARM	325MG;50MG;40MG	N40601 001	Jul 29, 2005	Jul	DISC
---	--------------	-----------------	------------	--------------	-----	------

ACETAMINOPHEN; CODEINE PHOSPHATE

TABLET; ORAL

ACETAMINOPHEN AND CODEINE PHOSPHATE

@	SANDOZ	300MG;30MG	N81250 001	Jul 16, 1992	Mar	DISC
---	--------	------------	------------	--------------	-----	------

@		300MG;60MG	N81249 001	Jul 16, 1992	Mar	DISC
---	--	------------	------------	--------------	-----	------

TYLENOL W/ CODEINE NO. 4

AA	ORTHO MCNEIL JANSSEN	300MG;60MG	N85055 004		Mar	CMFD
----	----------------------	------------	------------	--	-----	------

ACETAMINOPHEN; HYDROCODONE BITARTRATE

CAPSULE; ORAL

HYDROCODONE BITARTRATE AND ACETAMINOPHEN

@	MALLINCKRODT	500MG;5MG	N89006 001	Aug 09, 1985	Feb	CTNA
---	--------------	-----------	------------	--------------	-----	------

>D>		LORCET-HD				
-----	--	-----------	--	--	--	--

>D>	AA	+	MALLINCKRODT	500MG;5MG	N87336 001	Jul 08, 1982	Aug	DISC
-----	----	---	--------------	-----------	------------	--------------	-----	------

>A>		@		500MG;5MG	N87336 001	Jul 08, 1982	Aug	DISC
-----	--	---	--	-----------	------------	--------------	-----	------

ACETAMINOPHEN; OXYCODONE HYDROCHLORIDE

TABLET; ORAL

PERCOCET

AA	+	ENDO PHARMS	325MG;2.5MG	N40330 001	Jun 25, 1999	May	CTEC
----	---	-------------	-------------	------------	--------------	-----	------

ACETAMINOPHEN; PROPOXYPHENE HYDROCHLORIDE

TABLET; ORAL

PROPOXYPHENE HYDROCHLORIDE AND ACETAMINOPHEN

@	SANDOZ	650MG;65MG	N89959 001	Jul 18, 1989	Mar	DISC
---	--------	------------	------------	--------------	-----	------

>D>	AA	+	WATSON LABS	650MG;65MG	N40139 001	Dec 16, 1996	Aug	DISC
-----	----	---	-------------	------------	------------	--------------	-----	------

>A>		@		650MG;65MG	N40139 001	Dec 16, 1996	Aug	DISC
-----	--	---	--	------------	------------	--------------	-----	------

ACETAMINOPHEN; PROPOXYPHENE NAPSYLATE

TABLET; ORAL

PROPOXYPHENE NAPSYLATE AND ACETAMINOPHEN

@	SANDOZ	650MG;100MG	N70443 001	Jan 23, 1986	Mar	DISC
---	--------	-------------	------------	--------------	-----	------

ACETAMINOPHEN; TRAMADOL HYDROCHLORIDE

TABLET; ORAL

TRAMADOL HYDROCHLORIDE AND ACETAMINOPHEN

AB	MYLAN	325MG;37.5MG	N77858 001	Sep 26, 2008	Apr	CAHN
----	-------	--------------	------------	--------------	-----	------

ACETAZOLAMIDE SODIUM

INJECTABLE; INJECTION

ACETAZOLAMIDE SODIUM

@ HOSPIRA

EQ 500MG BASE/VIAL

N40108 001 Oct 30, 1995 Apr DISC

ACETIC ACID, GLACIAL

SOLUTION/DROPS; OTIC

VOSOL

AT HI TECH PHARMA

2%

N12179 001

Jul CMFD

ACETIC ACID, GLACIAL; HYDROCORTISONE

SOLUTION/DROPS; OTIC

HYDROCORTISONE AND ACETIC ACID

>D> AT MORTON GROVE

2%;1%

N40168 001 Aug 30, 1996 Aug DISC

>A> @

2%;1%

N40168 001 Aug 30, 1996 Aug DISC

ACYCLOVIR

CAPSULE; ORAL

ACYCLOVIR

@ BELCHER PHARMS

200MG

N74889 001 Oct 31, 1997 Jun CAHN

TABLET; ORAL

ACYCLOVIR

@ BELCHER PHARMS

400MG

N74891 001 Oct 31, 1997 Jun CAHN

@

800MG

N74891 002 Oct 31, 1997 Jun CAHN

ACYCLOVIR SODIUM

INJECTABLE; INJECTION

ACYCLOVIR SODIUM

@ HOSPIRA

EQ 500MG BASE/VIAL

N74758 001 Apr 22, 1997 Apr DISC

@

EQ 1GM BASE/VIAL

N74758 002 Apr 22, 1997 Apr DISC

ACYCLOVIR; HYDROCORTISONE

CREAM; TOPICAL

ACYCLOVIR AND HYDROCORTISONE

+ MEDIVIR

5%;1%

N22436 001 Jul 31, 2009 Jul NEWA

ADENOSINE

INJECTABLE; INJECTION

ADENOSINE

AP LUITPOLD

3MG/ML

N90010 001 Apr 28, 2009 Apr NEWA

AP STRIDES ARCOLAB LTD

3MG/ML

N78686 001 May 13, 2009 May NEWA

AP WOCKHARDT

3MG/ML

N90220 001 Jul 20, 2009 Jun NEWA

ALBUTEROL

AEROSOL, METERED; INHALATION

ALBUTEROL

@ ARMSTRONG PHARMS

0.09MG/INH

N72273 001 Aug 14, 1996 Jan DISC

ALBUTEROL SULFATE

SOLUTION; INHALATION

ALBUTEROL SULFATE

@ BAUSCH AND LOMB

EQ 0.083% BASE

N75358 001 Mar 29, 2000 Apr DISC

AN HOLOPACK INTL

EQ 0.083% BASE

N77839 001 Dec 16, 2008 Jan CAHN

AN TEVA PARENTERAL

EQ 0.083% BASE

N75343 001 Nov 09, 1999 Apr CAHN

TABLET; ORAL

ALBUTEROL SULFATE

@ SANDOZ

EQ 2MG BASE

N72151 001 Dec 05, 1989 Mar DISC

@

EQ 4MG BASE

N72152 001 Dec 05, 1989 Mar DISC

ALBUTEROL SULFATE; IPRATROPIUM BROMIDE

SOLUTION; INHALATION

ALBUTEROL SULFATE AND IPRATROPIUM BROMIDE

AN TEVA PARENTERAL EQ 0.083% BASE;0.017%

N76724 001 Dec 31, 2007 Apr CAHN

ALCLOMETASONE DIPROPIONATE

CREAM; TOPICAL

ALCLOMETASONE DIPROPIONATE

AB GLENMARK GENERICS 0.05%

N79061 001 Jun 23, 2009 Jun NEWA

OINTMENT; TOPICAL

ALCLOMETASONE DIPROPIONATE

AB GLENMARK GENERICS 0.05%

N79227 001 Jul 30, 2009 Jul NEWA

ALENDRONATE SODIUM

TABLET; ORAL

ALENDRONATE SODIUM

AB SANDOZ EQ 5MG BASE

N75871 001 Apr 22, 2009 Apr NEWA

AB EQ 10MG BASE

N75871 002 Apr 22, 2009 Apr NEWA

AB EQ 35MG BASE

N75871 004 Apr 22, 2009 Apr NEWA

AB EQ 40MG BASE

N75871 003 Apr 22, 2009 Apr NEWA

AB EQ 70MG BASE

N75871 005 Apr 22, 2009 Apr NEWA

ALISKIREN FUMARATE; HYDROCHLOROTHIAZIDE

TABLET; ORAL

TEKTURNA HCT

>A> + NOVARTIS EQ 300MG BASE;12.5MG

N22107 003 Jan 18, 2008 Aug CRLD

ALISKIREN HEMIFUMARATE; HYDROCHLOROTHIAZIDE

TABLET; ORAL

TEKTURNA HCT

>D> NOVARTIS EQ 300MG BASE;12.5MG

N22107 003 Jan 18, 2008 Aug CRLD

ALLOPURINOL

TABLET; ORAL

ALLOPURINOL

@ SANDOZ

100MG

N70268 001 Dec 31, 1985 Mar DISC

@

300MG

N70269 001 Dec 31, 1985 Mar DISC

AMIKACIN SULFATE

INJECTABLE; INJECTION

AMIKACIN SULFATE

>D> HOSPIRA EQ 62.5MG BASE/ML

N63283 001 Oct 31, 1994 Aug DISC

>A> @ EQ 62.5MG BASE/ML

N63283 001 Oct 31, 1994 Aug DISC

AMILORIDE HYDROCHLORIDE

TABLET; ORAL

AMILORIDE HYDROCHLORIDE

AB + PAR PHARM 5MG

N70346 001 Jan 22, 1986 Jan CTEC

AB SIGMAPHARM LABS LLC 5MG

N79133 001 Jan 30, 2009 Jan NEWA

MIDAMOR

AB PADDOCK LABS 5MG

N18200 001 May CMFD

AMILORIDE HYDROCHLORIDE; HYDROCHLOROTHIAZIDE

TABLET; ORAL

AMILORIDE HYDROCHLORIDE AND HYDROCHLOROTHIAZIDE
@ SANDOZ EQ 5MG ANHYDROUS;50MG

N73357 001 Nov 27, 1991 Mar DISC

AMINOCAPROIC ACID

INJECTABLE; INJECTION

AMINOCAPROIC ACID
@ HOSPIRA 250MG/ML

N70888 001 Jun 16, 1988 Apr DISC

TABLET; ORAL

AMICAR
XANODYNE PHARM 1GM

N15197 002 Jun 24, 2004 Jan NEWA

AMINOPHYLLINE

INJECTABLE; INJECTION

AMINOPHYLLINE
AP + HOSPIRA 25MG/ML
AMINOPHYLLINE IN SODIUM CHLORIDE 0.45%
@ HOSPIRA 100MG/100ML
@ 200MG/100ML

N87242 001 Oct 26, 1983 Jun CRLD

N88147 002 May 03, 1983 Apr DISC

N88147 003 May 03, 1983 Apr DISC

SUPPOSITORY; RECTAL

TRUPHYLLINE
@ G AND W LABS 250MG

N85498 001 Mar 23, 1983 Jul DISC

AMIODARONE HYDROCHLORIDE

INJECTABLE; INJECTION

AMIODARONE HYDROCHLORIDE
@ INTL MEDICATION SYS 50MG/ML

N21594 001 Feb 04, 2004 Jun DISC

TABLET; ORAL

AMIODARONE HYDROCHLORIDE
AB MYLAN 200MG

N75188 001 Feb 24, 1999 Apr CAHN

AMLODIPINE BESYLATE

TABLET; ORAL

AMLODIPINE BESYLATE

AB	ALKEM	EQ 2.5MG BASE	N78925 001	May 04, 2009	Apr	NEWA
AB		EQ 5MG BASE	N78925 002	May 04, 2009	Apr	NEWA
AB		EQ 10MG BASE	N78925 003	May 04, 2009	Apr	NEWA
AB	GLENMARK GENERICS	EQ 2.5MG BASE	N78552 001	Apr 08, 2009	Mar	NEWA
AB		EQ 5MG BASE	N78552 002	Apr 08, 2009	Mar	NEWA
AB		EQ 10MG BASE	N78552 003	Apr 08, 2009	Mar	NEWA
AB	ORCHID HLTHCARE	EQ 2.5MG BASE	N78453 001	Jul 02, 2009	Jun	NEWA
AB		EQ 5MG BASE	N78453 002	Jul 02, 2009	Jun	NEWA
AB		EQ 10MG BASE	N78453 003	Jul 02, 2009	Jun	NEWA
AB	SYNTHON PHARMS	EQ 2.5MG BASE	N77080 001	Jun 27, 2007	Jan	CAHN
AB		EQ 5MG BASE	N77080 002	Jun 27, 2007	Jan	CAHN
AB		EQ 10MG BASE	N77080 003	Jun 27, 2007	Jan	CAHN

AMLODIPINE; HYDROCHLOROTHIAZIDE; VALSARTAN

TABLET; ORAL

EXFORGE HCT
NOVARTIS 5MG;12.5MG;160MG
5MG;25MG;160MG
10MG;12.5MG;160MG
+ 10MG;25MG;320MG

N22314 001 Apr 30, 2009 Apr NEWA

N22314 002 Apr 30, 2009 Apr NEWA

N22314 003 Apr 30, 2009 Apr NEWA

N22314 005 Apr 30, 2009 Apr NEWA

TABLET; ORAL
EXFORGE HCT
NOVARTIS

10MG;25MG;160MG

N22314 004 Apr 30, 2009 Apr NEWA

AMOXICILLIN

CAPSULE; ORAL
AMOXIL

>D> AB GLAXOSMITHKLINE 250MG N62216 001 Aug DISC
>A> @ 250MG N62216 001 Aug DISC

AMOXICILLIN; CLAVULANATE POTASSIUM

TABLET; ORAL
AMOXICILLIN AND CLAVULANATE POTASSIUM

AB APOTEX 250MG;EQ 125MG BASE N65333 001 Feb 24, 2009 Feb NEWA
AB 500MG;EQ 125MG BASE N65333 002 Feb 24, 2009 Feb NEWA

AMPHETAMINE ASPARTATE; AMPHETAMINE SULFATE; DEXTROAMPHETAMINE SACCHARATE; DEXTROAMPHETAMINE SULFATE

TABLET; ORAL

DEXTROAMP SACCHARATE, AMP ASPARTATE, DEXTROAMP SULFATE AND AMP SULFATE

>D> AB WATSON LABS 1.25MG;1.25MG;1.25MG;1.25MG N40456 001 May 06, 2003 Aug DISC
>A> @ 1.25MG;1.25MG;1.25MG;1.25MG N40456 001 May 06, 2003 Aug DISC
>D> AB 2.5MG;2.5MG;2.5MG;2.5MG N40456 002 May 06, 2003 Aug DISC
>A> @ 2.5MG;2.5MG;2.5MG;2.5MG N40456 002 May 06, 2003 Aug DISC
>D> AB 5MG;5MG;5MG;5MG N40456 003 May 06, 2003 Aug DISC
>A> @ 5MG;5MG;5MG;5MG N40456 003 May 06, 2003 Aug DISC
>D> AB 7.5MG;7.5MG;7.5MG;7.5MG N40456 004 May 06, 2003 Aug DISC
>A> @ 7.5MG;7.5MG;7.5MG;7.5MG N40456 004 May 06, 2003 Aug DISC

APRACLOINIDINE HYDROCHLORIDE

SOLUTION/DROPS; OPHTHALMIC
APRACLOINIDINE HYDROCHLORIDE

AT AKORN INC EQ 0.5% BASE N77764 001 Mar 12, 2009 Feb NEWA
AT + ALCON IOPIDINE EQ 0.5% BASE N20258 001 Jul 30, 1993 Feb CFTG

ARMODAFINIL

TABLET; ORAL
NUVIGIL
CEPHALON

100MG
200MG

N21875 002 Mar 26, 2009 Mar CMFD
N21875 005 Mar 26, 2009 Mar NEWA

ARTEMETHER; LUMEFANTRINE

TABLET; ORAL
COARTEM
NOVARTIS

20MG;120MG

N22268 001 Apr 07, 2009 Apr NEWA

>A> ASENAPINE

>A> TABLET; SUBLINGUAL
>A> SAPHRIS

>A> ORGANON USA INC 5MG N22117 001 Aug 13, 2009 Aug NEWA
>A> + 10MG N22117 002 Aug 13, 2009 Aug NEWA

ASPIRIN; BUTALBITAL; CAFFEINE

TABLET; ORAL

BUTALBITAL, ASPIRIN AND CAFFEINE

@ SANDOZ 325MG;50MG;40MG

N86398 002 Apr 06, 1984 Mar DISC

ASPIRIN; CAFFEINE; ORPHENADRINE CITRATE

TABLET; ORAL

ORPHENGESIC

@ SOLCO HLTHCARE 385MG;30MG;25MG

N75141 001 May 29, 1998 Jan CAHN

ORPHENGESIC FORTE

@ SOLCO HLTHCARE 770MG;60MG;50MG

N75141 002 May 29, 1998 Jan CAHN

ASPIRIN; DIPYRIDAMOLE

CAPSULE, EXTENDED RELEASE; ORAL

AGGRENOX

>D> + BOEHRINGER INGELHEIM 25MG;200MG

N20884 001 Nov 22, 1999 Aug CFTG

>A> AB + 25MG;200MG

N20884 001 Nov 22, 1999 Aug CFTG

>A> ASPIRIN AND DIPYRIDAMOLE

>A> AB BARR 25MG;200MG

N78804 001 Aug 14, 2009 Aug NEWA

ATENOLOL

TABLET; ORAL

ATENOLOL

AB IPCA LABS LTD 25MG

N77877 001 Dec 27, 2006 Jun CAHN

AB 50MG

N77877 002 Dec 27, 2006 Jun CAHN

AB 100MG

N77877 003 Dec 27, 2006 Jun CAHN

ATRACURIUM BESYLATE

INJECTABLE; INJECTION

ATRACURIUM BESYLATE

>D> AP BAXTER HLTHCARE CORP 10MG/ML

N74753 001 Jan 23, 1997 Aug DISC

>A> @ 10MG/ML

N74753 001 Jan 23, 1997 Aug DISC

>D> AP + BEDFORD 10MG/ML

N74901 001 Jul 18, 1997 Aug CTEC

>A> + 10MG/ML

N74901 001 Jul 18, 1997 Aug CTEC

@ HOSPIRA 10MG/ML

N74740 001 Mar 28, 1997 Jan DISC

>D> AP MARSAM PHARMS LLC 10MG/ML

N74945 001 Jul 28, 1998 Aug DISC

>A> @ 10MG/ML

N74945 001 Jul 28, 1998 Aug DISC

>D> AP TEVA PARENTERAL 10MG/ML

N74784 001 Jun 11, 1997 Aug DISC

>A> @ 10MG/ML

N74784 001 Jun 11, 1997 Aug DISC

ATRACURIUM BESYLATE PRESERVATIVE FREE

>D> AP BAXTER HLTHCARE CORP 10MG/ML

N74768 001 Jan 23, 1997 Aug DISC

>A> @ 10MG/ML

N74768 001 Jan 23, 1997 Aug DISC

>D> AP + BEDFORD 10MG/ML

N74900 001 Jul 18, 1997 Aug CTEC

>A> + 10MG/ML

N74900 001 Jul 18, 1997 Aug CTEC

@ HOSPIRA 10MG/ML

N74741 001 Mar 28, 1997 Jan DISC

>D> AP MARSAM PHARMS LLC 10MG/ML

N74944 001 Jul 28, 1998 Aug DISC

>A> @ 10MG/ML

N74944 001 Jul 28, 1998 Aug DISC

ATROPINE SULFATE; DIFENOXIN HYDROCHLORIDE

TABLET; ORAL

MOTOFEN

+ VALEANT 0.025MG;1MG

N17744 002 Jun CMFD

ATROPINE SULFATE; DIPHENOXYLATE HYDROCHLORIDE

TABLET; ORAL

>D>		DIPHENOXYLATE HYDROCHLORIDE W/ ATROPINE SULFATE					
>D>	AA	SANDOZ	0.025MG;2.5MG	N86173 001		Aug	DISC
>A>		@	0.025MG;2.5MG	N86173 001		Aug	DISC

ATROPINE SULFATE; EDROPHONIUM CHLORIDE

INJECTABLE; INJECTION

ENLON-PLUS

+	BIONICHE PHARMA	0.14MG/ML;10MG/ML	N19677 001	Nov 06, 1991	Jul	CMFD
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AZACITIDINE

INJECTABLE; IV-SC

VIDAZA

+	CELGENE	100MG/VIAL	N50794 001	May 19, 2004	Mar	CAHN
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AZELASTINE HYDROCHLORIDE

SOLUTION/DROPS; OPHTHALMIC

AZELASTINE HYDROCHLORIDE

AT		APOTEX INC	0.05%	N78621 001	Aug 03, 2009	Jul	NEWA
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OPTIVAR

AT	+	MEDA PHARMS	0.05%	N21127 001	May 22, 2000	Jul	CFTG
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SPRAY, METERED; NASAL

ASTEPRO

>A>	+	MEDA PHARMS	EQ 0.1876MG BASE/SPRAY	N22371 001	Aug 31, 2009	Aug	NEWA
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AZELASTINE HYDROCHLORIDE

@	APOTEX INC	EQ 0.125MG BASE/SPRAY	N77954 001	Apr 30, 2009	Apr	DISC
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AZITHROMYCIN

INJECTABLE; INJECTION

AZITHROMYCIN

AP		HOSPIRA	EQ 500MG BASE/VIAL	N65500 001	Jun 26, 2009	Jun	NEWA
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AP			EQ 500MG BASE/VIAL	N65511 001	Jun 26, 2009	Jun	NEWA
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AP		SAGENT STRIDES	EQ 500MG BASE/VIAL	N65506 001	Mar 24, 2009	Mar	NEWA
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BACLOFEN

TABLET; ORAL

BACLOFEN

AB		MYLAN	10MG	N77181 001	Jul 29, 2005	Apr	CAHN
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AB			20MG	N77121 002	Jul 29, 2005	Apr	CAHN
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BENDAMUSTINE HYDROCHLORIDE

POWDER; IV (INFUSION)

TREANDA

+	CEPHALON	25MG/VIAL	N22249 002	May 01, 2009	May	NEWA
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BENZOYL PEROXIDE; CLINDAMYCIN PHOSPHATE

GEL; TOPICAL

BENZACLIN

AB	+	SANOFI AVENTIS US	5%;EQ 1% BASE	N50756 001	Dec 21, 2000	Jul	CFTG
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CLINDAMYCIN PHOSPHATE AND BENZOYL PEROXIDE

AB		DOW PHARM SCIENCES	5%;EQ 1% BASE	N65443 001	Aug 11, 2009	Jul	NEWA
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BENZTROPINE MESYLATE

INJECTABLE; INJECTION

BENZTROPINE MESYLATE

>A>	AP	HIKMA FARMACEUTICA	1MG/ML	N90287 001	Aug 31, 2009	Aug	NEWA
	AP	NEXUS PHARMS	1MG/ML	N90233 001	Jul 28, 2009	Jul	NEWA
		COGENTIN					
	AP	+ LUNDBECK INC	1MG/ML	N12015 001		Jul	CFTG
		+	1MG/ML	N12015 001		Apr	CAHN

BENZYL ALCOHOL

LOTION; TOPICAL

BENZYL ALCOHOL

	+	SCIELE PHARMA INC	5%	N22129 001	Apr 09, 2009	Apr	NEWA
		ULESFIA					
	+	SCIELE PHARMA INC	5%	N22129 001	Apr 09, 2009	May	CTNA

BESIFLOXACIN HYDROCHLORIDE

SUSPENSION/DROPS; OPHTHALMIC

BESIVANCE

	+	BAUSCH AND LOMB	EQ 0.6% BASE	N22308 001	May 28, 2009	May	NEWA
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BETAMETHASONE ACETATE; BETAMETHASONE SODIUM PHOSPHATE

INJECTABLE; INJECTION

BETAMETHASONE ACETATE AND BETAMETHASONE SODIUM PHOSPHATE

AB		PHARMAFORCE	3MG/ML;EQ 3MG BASE/ML	N90747 001	Jul 31, 2009	Jul	NEWA
		CELESTONE SOLUSPAN					
AB	+	SCHERING	3MG/ML;EQ 3MG BASE/ML	N14602 001		Jul	CFTG

BETAMETHASONE DIPROPIONATE; CALCIPOTRIENE HYDRATE

OINTMENT; TOPICAL

TACLONEX

	+	LEO PHARM	0.064%;0.005%	N21852 001	Jan 09, 2006	Mar	CAHN
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BETAMETHASONE VALERATE

AEROSOL, FOAM; TOPICAL

LUXIQ

	+	CONNECTICS	EQ 0.12% BASE	N20934 001	Feb 28, 1999	May	CAHN
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BETHANECHOL CHLORIDE

TABLET; ORAL

BETHANECHOL CHLORIDE

AA		SUN PHARM INDS INC	5MG	N40897 001	Apr 22, 2009	Apr	NEWA
AA			10MG	N40897 002	Apr 22, 2009	Apr	NEWA
AA			25MG	N40897 003	Apr 22, 2009	Apr	NEWA
AA			50MG	N40897 004	Apr 22, 2009	Apr	NEWA

BICALUTAMIDE

TABLET; ORAL

BICALUTAMIDE

	AB	ACCORD HLTHCARE INC	50MG	N78917 001	Jul 06, 2009	Jun	NEWA
>A>	AB	ACTAVIS TOTOWA	50MG	N78634 001	Aug 28, 2009	Aug	NEWA
	AB	KUDCO IRELAND	50MG	N77995 001	Jul 06, 2009	Jun	NEWA
	AB	MYLAN	50MG	N79185 001	Jul 06, 2009	Jun	NEWA
	AB	SANDOZ	50MG	N78575 001	Jul 06, 2009	Jun	NEWA
	AB	SUN PHARMA GLOBAL	50MG	N79110 001	Jul 06, 2009	Jun	NEWA

TABLET; ORAL

BICALUTAMIDE

AB	SYNTHON PHARMS	50MG	N77973 001	Jul 06, 2009	Jun	NEWA
AB	TEVA	50MG	N76932 001	Jul 06, 2009	Jun	NEWA
AB	ZYDUS PHARMS USA INC	50MG	N79089 001	Jul 06, 2009	Jun	NEWA
CASODEX						
AB	+ ASTRAZENECA	50MG	N20498 001	Oct 04, 1995	Jun	CFTG

BISOPROLOL FUMARATE

TABLET; ORAL

BISOPROLOL FUMARATE

>A>	AB	UNICHEM PHARMS (USA)	5MG	N78635 001	Aug 18, 2009	Aug	NEWA
>A>	AB		10MG	N78635 002	Aug 18, 2009	Aug	NEWA

BRIMONIDINE TARTRATE

SOLUTION/DROPS; OPHTHALMIC

BRIMONIDINE TARTRATE

AT	TEVA PARENTERAL	0.2%	N76372 001	Sep 10, 2004	Apr	CAHN
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BROMOCRIPTINE MESYLATE

TABLET; ORAL

CYCLOSET

	+ VEROSCIENCE	EQ 0.8MG BASE	N20866 001	May 05, 2009	May	NEWA
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BROMPHENIRAMINE MALEATE

>D>	INJECTABLE; INJECTION					
>D>	BROMPHENIRAMINE MALEATE					
>D>	+ WATSON LABS	10MG/ML	N83821 001		Aug	DISC
>A>	@	10MG/ML	N83821 001		Aug	DISC

BROMPHENIRAMINE MALEATE; DEXTROMETHORPHAN HYDROBROMIDE; PSEUDOEPHEDRINE HYDROCHLORIDE

SYRUP; ORAL

BROMFED-DM

	+ MORTON GROVE	2MG/5ML;10MG/5ML;30MG/5ML	N88811 001	Jun 07, 1985	Jun	CTNA
	@ WOCKHARDT EU	2MG/5ML;10MG/5ML;30MG/5ML	N89681 001	Dec 22, 1988	Jun	DISC
AA		2MG/5ML;10MG/5ML;30MG/5ML	N89681 001	Dec 22, 1988	Apr	CAHN

BUDESONIDE

SUSPENSION; INHALATION

BUDESONIDE

AN	APOTEX	0.25MG/2ML	N78202 001	Mar 30, 2009	Mar	NEWA
AN		0.5MG/2ML	N78202 002	Mar 30, 2009	Mar	NEWA
AN	TEVA PARENTERAL	0.25MG/2ML	N77519 001	Nov 18, 2008	Apr	CAHN
AN		0.5MG/2ML	N77519 002	Nov 18, 2008	Apr	CAHN

BUPIVACAINE HYDROCHLORIDE

INJECTABLE; INJECTION

SENSORCAINE

AP	APP PHARMS	0.25%	N70552 001	May 21, 1986	Jun	CAHN
AP		0.5%	N70553 001	May 21, 1986	Jun	CAHN
AP		0.75%	N70554 001	May 21, 1986	Jun	CAHN

INJECTABLE; SPINAL

SENSORCAINE

AP	APP PHARMS	0.75%	N71202 001	Apr 15, 1987	Jun	CAHN
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BUPIVACAINE HYDROCHLORIDE; EPINEPHRINE BITARTRATE

INJECTABLE; INJECTION

SENSORCAINE

AP	APP PHARMS	0.25%;0.0091MG/ML	N70966 001	Oct 13, 1987	Jun	CAHN
AP		0.25%;0.0091MG/ML	N70967 001	Oct 13, 1987	Jun	CAHN
AP		0.5%;0.0091MG/ML	N70968 001	Oct 13, 1987	Jun	CAHN

BUPROPION HYDROCHLORIDE

TABLET, EXTENDED RELEASE; ORAL

BUPROPION HYDROCHLORIDE

AB1	WATSON LABS	100MG	N79095 001	Mar 24, 2009	Mar	NEWA
AB2		150MG	N79094 001	Mar 24, 2009	Mar	NEWA
AB1		150MG	N79095 002	Mar 24, 2009	Mar	NEWA
AB1		200MG	N79095 003	Mar 24, 2009	Mar	NEWA
	WELLBUTRIN XL					
AB3 +	BIOVAIL LABS INTL	150MG	N21515 001	Aug 28, 2003	Jul	CAHN
AB3		300MG	N21515 002	Aug 28, 2003	Jul	CAHN

BUSPIRONE HYDROCHLORIDE

TABLET; ORAL

BUSPAR

AB	+ BRISTOL MYERS SQUIBB	15MG	N18731 003	Apr 22, 1996	Feb	CRLD
	@	30MG	N18731 004	Apr 22, 1996	Feb	DISC

BUSPIRONE HYDROCHLORIDE

AB	DR REDDYS LABS LTD	5MG	N78246 001	Feb 27, 2009	Feb	NEWA
AB		10MG	N78246 002	Feb 27, 2009	Feb	NEWA
AB		15MG	N78246 003	Feb 27, 2009	Feb	NEWA
AB		30MG	N78246 004	Feb 27, 2009	Feb	NEWA
AB	MYLAN	5MG	N75467 001	Feb 28, 2002	Apr	CAHN
AB		7.5MG	N75467 002	Mar 28, 2001	Apr	CAHN
AB		10MG	N75467 003	Feb 28, 2002	Apr	CAHN
AB		15MG	N75467 004	Feb 28, 2002	Apr	CAHN
	@ SANDOZ	5MG	N75413 001	Mar 19, 2002	Mar	DISC
	@	10MG	N75413 002	Mar 19, 2002	Mar	DISC
	@	15MG	N75413 003	Mar 19, 2002	Mar	DISC

BUTORPHANOL TARTRATE

INJECTABLE; INJECTION

BUTORPHANOL TARTRATE

AP	HIKMA FARMACEUTICA	1MG/ML	N78400 001	May 01, 2009	Apr	NEWA
AP		2MG/ML	N78247 001	Apr 29, 2009	Apr	NEWA
AP		2MG/ML	N78400 002	May 01, 2009	Apr	NEWA

CAFFEINE CITRATE

SOLUTION; INTRAVENOUS

CAFFEINE CITRATE

AP	PADDOCK LABS	EQ 30MG BASE/3ML (EQ 10MG BASE/ML)	N77233 001	Sep 21, 2006	Apr	CAHN
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CALCITONIN, SALMON

SPRAY, METERED; NASAL

CALCITONIN-SALMON

AB	MDRNA	200 IU/SPRAY	N76979 001	Jun 08, 2009	May	NEWA
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CALCITRIOL

CAPSULE; ORAL

ROCALTROL

AB	VALIDUS PHARMS	0.25UGM	N18044 001		Mar	CAHN
AB	+	0.5UGM	N18044 002		Mar	CAHN

OINTMENT; TOPICAL

VECTICAL

+	GALDERMA LABS LP	3UGM/GM	N22087 001	Jan 23, 2009	Jan	NEWA
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SOLUTION; ORAL

ROCALTROL

AA	+	VALIDUS PHARMS	1UGM/ML	N21068 001	Nov 20, 1998	Mar	CAHN
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CAPECITABINE

TABLET; ORAL

XELODA

	HOFFMANN LA ROCHE	150MG	N20896 001	Apr 30, 1998	Jul	CAHN
+		500MG	N20896 002	Apr 30, 1998	Jul	CAHN

CAPREOMYCIN SULFATE

INJECTABLE; INJECTION

CAPASTAT SULFATE

>A>	+	AKORN	EQ 1GM BASE/VIAL	N50095 001		Aug	CAHN
>D>	+	LILLY	EQ 1GM BASE/VIAL	N50095 001		Aug	CAHN

CARBAMAZEPINE

TABLET; ORAL

CARBAMAZEPINE

	TORRENT PHARMS	100MG	N77272 001	Dec 07, 2005	Apr	CAHN
AB		200MG	N77272 002	Dec 07, 2005	Apr	CAHN
		300MG	N77272 003	Dec 07, 2005	Apr	CAHN
		400MG	N77272 004	Dec 07, 2005	Apr	CAHN

TABLET, CHEWABLE; ORAL

CARBAMAZEPINE

AB	TORRENT PHARMS	100MG	N75712 001	Jul 05, 2001	Jan	CAHN
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TABLET, EXTENDED RELEASE; ORAL

CARBAMAZEPINE

AB	TARO	100MG	N78115 001	Mar 31, 2009	Mar	NEWA
AB		200MG	N78115 002	Mar 31, 2009	Mar	NEWA
AB		400MG	N78115 003	Mar 31, 2009	Mar	NEWA

TEGRETOL -XR

AB	NOVARTIS	100MG	N20234 001	Mar 25, 1996	Mar	CFTG
AB		200MG	N20234 002	Mar 25, 1996	Mar	CFTG
AB	+	400MG	N20234 003	Mar 25, 1996	Mar	CFTG

CARBENICILLIN INDANYL SODIUM

TABLET; ORAL

GEOCILLIN

@	PFIZER	EQ 382MG BASE	N50435 001		Mar	DISC
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CARBIDOPA; LEVODOPA

TABLET; ORAL

CARBIDOPA AND LEVODOPA

@	SANDOZ	10MG;100MG	N73586 001	Jun 29, 1995	Mar	DISC
@		25MG;100MG	N73587 001	Jun 29, 1995	Mar	DISC
@		25MG;250MG	N73620 001	Jun 29, 1995	Mar	DISC

TABLET, EXTENDED RELEASE; ORAL

CARBIDOPA AND LEVODOPA

@ KV PHARM	50MG;200MG	N76663 001	Jun 24, 2004	Apr	DISC
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TABLET, FOR SUSPENSION; ORAL

CARBILEV

@ RANBAXY	10MG;100MG	N76643 001	Jun 10, 2005	Jul	DISC
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@	25MG;100MG	N76643 002	Jun 10, 2005	Jul	DISC
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@	25MG;250MG	N76643 003	Jun 10, 2005	Jul	DISC
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TABLET, ORALLY DISINTEGRATING; ORAL

CARBIDOPA AND LEVODOPA

AB	SUN PHARM INDS	10MG;100MG	N78690 001	Jul 31, 2009	Jul	NEWA
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AB		25MG;100MG	N78690 002	Jul 31, 2009	Jul	NEWA
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AB		25MG;250MG	N78690 003	Jul 31, 2009	Jul	NEWA
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CARBOPLATIN

INJECTABLE; INJECTION

CARBOPLATIN

AP	+	WATSON LABS	50MG/VIAL	N76162 001	Oct 14, 2004	Apr	CAHN
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AP	+		150MG/VIAL	N76162 002	Oct 14, 2004	Apr	CAHN
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AP	+		450MG/VIAL	N76162 003	Oct 14, 2004	Apr	CAHN
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INJECTABLE; IV (INFUSION)

CARBOPLATIN

	@	ABRAXIS PHARM	50MG/5ML (10MG/ML)	N77247 001	Oct 21, 2004	Jul	CPOT
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AP			50MG/5ML (10MG/ML)	N77266 001	Feb 15, 2006	Jul	CPOT
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	@		150MG/15ML (10MG/ML)	N77247 002	Oct 21, 2004	Jul	CPOT
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AP			150MG/15ML (10MG/ML)	N77266 002	Feb 15, 2006	Jul	CPOT
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AP			450MG/45ML (10MG/ML)	N77247 003	Oct 21, 2004	Jul	CPOT
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AP			450MG/45ML (10MG/ML)	N77266 003	Feb 15, 2006	Jul	CPOT
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AP			600MG/60ML (10MG/ML)	N77266 004	Feb 15, 2006	Jul	CPOT
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AP		AKORN	50MG/5ML (10MG/ML)	N90475 001	Jul 29, 2009	Jul	NEWA
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AP			150MG/15ML (10MG/ML)	N90475 002	Jul 29, 2009	Jul	NEWA
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AP			450MG/45ML (10MG/ML)	N90475 003	Jul 29, 2009	Jul	NEWA
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AP		BEDFORD LABS	50MG/5ML (10MG/ML)	N77244 001	Oct 15, 2004	Jul	CPOT
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AP			150MG/15ML (10MG/ML)	N77244 002	Oct 15, 2004	Jul	CPOT
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AP			450MG/45ML (10MG/ML)	N77244 003	Oct 15, 2004	Jul	CPOT
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AP			600MG/60ML (10MG/ML)	N77244 004	Jan 20, 2006	Jul	CPOT
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AP		EBEWE PHARMA	50MG/5ML (10MG/ML)	N78280 001	May 08, 2008	Jul	CPOT
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AP			150MG/15ML (10MG/ML)	N78280 002	May 08, 2008	Jul	CPOT
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AP			450MG/45ML (10MG/ML)	N78280 003	May 08, 2008	Jul	CPOT
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AP		FRESENIUS KABI ONCOL	450MG/45ML (10MG/ML)	N77432 003	Sep 29, 2006	Jul	CPOT
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AP			50MG/5ML (10MG/ML)	N77432 001	Sep 29, 2006	Jul	CPOT
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AP			150MG/15ML (10MG/ML)	N77432 002	Sep 29, 2006	Jul	CPOT
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AP		GENERAMEDIX	50MG/5ML (10MG/ML)	N77998 001	Apr 24, 2007	Jul	CPOT
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AP			150MG/15ML (10MG/ML)	N77998 002	Apr 24, 2007	Jul	CPOT
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AP			450MG/45ML (10MG/ML)	N77998 003	Apr 24, 2007	Jul	CPOT
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AP		HOSPIRA	50MG/5ML (10MG/ML)	N76517 001	Oct 14, 2004	Jul	CPOT
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AP			150MG/15ML (10MG/ML)	N76517 002	Oct 14, 2004	Jul	CPOT
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AP			450MG/45ML (10MG/ML)	N76517 003	Oct 14, 2004	Jul	CPOT
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AP			600MG/60ML (10MG/ML)	N77059 001	Nov 23, 2004	Jul	CPOT
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AP	+	PHARMACHEMIE	50MG/5ML (10MG/ML)	N77269 001	Oct 14, 2004	Jul	CPOT
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AP	+		150MG/15ML (10MG/ML)	N77269 002	Oct 14, 2004	Jul	CPOT
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AP	+		450MG/45ML (10MG/ML)	N77269 003	Oct 14, 2004	Jul	CPOT
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AP	+		600MG/60ML (10MG/ML)	N77269 004	Dec 28, 2007	Jul	CPOT
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AP		PHARMACHEMIE BV	50MG/5ML (10MG/ML)	N77679 001	Feb 25, 2009	Jul	CPOT
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AP			EQ 50MG/5ML (10MG/ML)	N77679 001	Feb 25, 2009	Feb	NEWA
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AP			150MG/15ML (10MG/ML)	N77679 002	Feb 25, 2009	Jul	CPOT
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INJECTABLE; IV (INFUSION)

CARBOPLATIN

AP	PHARMACHEMIE BV	EQ 150MG/15ML (10MG/ML)	N77679 002	Feb 25, 2009	Feb	NEWA
AP		450MG/45ML (10MG/ML)	N77679 003	Feb 25, 2009	Jul	CPOT
AP		EQ 450MG/45ML (10MG/ML)	N77679 003	Feb 25, 2009	Feb	NEWA
AP	PLIVA LACHEMA	50MG/5ML (10MG/ML)	N78631 001	Dec 02, 2008	Jul	CPOT
AP		150MG/15ML (10MG/ML)	N78631 002	Dec 02, 2008	Jul	CPOT
AP		450MG/45ML (10MG/ML)	N78631 003	Dec 02, 2008	Jul	CPOT
AP		600MG/60ML (10MG/ML)	N78631 004	Dec 02, 2008	Jul	CPOT
AP	SPECTRUM PHARMS	50MG/5ML (10MG/ML)	N77096 001	Jun 14, 2005	Jul	CPOT
AP		150MG/15ML (10MG/ML)	N77096 002	Jun 14, 2005	Jul	CPOT
AP		450MG/45ML (10MG/ML)	N77096 003	Jun 14, 2005	Jul	CPOT
AP	SUN PHARM INDS	50MG/5ML (10MG/ML)	N77926 001	Sep 19, 2008	Jul	CPOT
AP		150MG/15ML (10MG/ML)	N77926 002	Sep 19, 2008	Jul	CPOT
AP		450MG/45ML (10MG/ML)	N77926 003	Sep 19, 2008	Jul	CPOT
AP	+ TEVA PARENTERAL	50MG/5ML (10MG/ML)	N77139 001	Sep 21, 2005	Jul	CPOT
AP		50MG/5ML (10MG/ML)	N77389 001	Mar 30, 2007	Jul	CPOT
AP	+	150MG/15ML (10MG/ML)	N77139 002	Sep 21, 2005	Jul	CPOT
AP		150MG/15ML (10MG/ML)	N77389 002	Mar 30, 2007	Jul	CPOT
AP	+	450MG/45ML (10MG/ML)	N77139 003	Sep 21, 2005	Jul	CPOT
AP		450MG/45ML (10MG/ML)	N77389 003	Mar 30, 2007	Jul	CPOT
AP	+	600MG/60ML (10MG/ML)	N77139 004	Sep 21, 2005	Jul	CPOT
AP	WATSON LABS	50MG/5ML (10MG/ML)	N77861 001	Jan 18, 2007	Jul	CPOT
AP		150MG/15ML (10MG/ML)	N77861 002	Jan 18, 2007	Jul	CPOT
AP		450MG/45ML (10MG/ML)	N77861 003	Jan 18, 2007	Jul	CPOT
AP		600MG/60ML (10MG/ML)	N77861 004	Jan 18, 2007	Jul	CPOT
PARAPLATIN						
AP	+ BRISTOL MYERS SQUIBB	50MG/5ML (10MG/ML)	N20452 001	Jul 14, 2003	Jul	CPOT
AP	+	150MG/15ML (10MG/ML)	N20452 002	Jul 14, 2003	Jul	CPOT
AP	+	450MG/45ML (10MG/ML)	N20452 003	Jul 14, 2003	Jul	CPOT
AP	+	600MG/60ML (10MG/ML)	N20452 004	Jan 15, 2004	Jul	CPOT

CARISOPRODOL

TABLET; ORAL

CARISOPRODOL

AA	AUROBINDO PHARMA	350MG	N40792 001	Aug 06, 2009	Jul	NEWA
	@ SANDOZ	350MG	N81025 001	Apr 13, 1989	Mar	DISC
>D>	AA	WATSON LABS	N40152 001	Dec 03, 1996	Aug	DISC
>A>	@	350MG	N40152 001	Dec 03, 1996	Aug	DISC
>D>	AA	350MG	N86179 001		Aug	DISC
>A>	@	350MG	N86179 001		Aug	DISC

CEFADROXIL ANHYDROUS

>D> CAPSULE; ORAL

>D> CEFADROXIL

>D>	AB	+ IVAX SUB TEVA PHARMS	EQ 500MG BASE	N62766 001	Mar 03, 1987	Aug	DISC
>A>	@		EQ 500MG BASE	N62766 001	Mar 03, 1987	Aug	DISC

TABLET; ORAL

CEFADROXIL

>D>	AB	+ IVAX SUB TEVA PHARMS	EQ 1GM BASE	N62774 001	Apr 08, 1987	Aug	CTEC
>A>	+		EQ 1GM BASE	N62774 001	Apr 08, 1987	Aug	CTEC

CEFAZOLIN SODIUM

INJECTABLE; INJECTION

ANCEF

	@ GLAXOSMITHKLINE	EQ 1GM BASE/VIAL	N50461 003		Jun	DISC
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INJECTABLE; INJECTION

ANCEF

@ GLAXOSMITHKLINE

EQ 10GM BASE/VIAL

N50461 005

Jun DISC

CEFAZOLIN SODIUM

AP CEPHAZONE PHARMA

EQ 500MG BASE/VIAL

N65280 001 Mar 18, 2009 Mar NEWA

AP EQ 1GM BASE/VIAL

N65280 002 Mar 18, 2009 Mar NEWA

AP EQ 10GM BASE/VIAL

N65295 001 Mar 18, 2009 Mar NEWA

AP EQ 20GM BASE/VIAL

N65296 001 Mar 18, 2009 Mar NEWA

@ GLAXOSMITHKLINE

EQ 1GM BASE/VIAL

N64033 001 Oct 31, 1993 Mar DISC

>D> AP HANFORD GC

EQ 500MG BASE/VIAL

N63214 001 Dec 27, 1991 Aug DISC

>A> @

EQ 500MG BASE/VIAL

N63214 001 Dec 27, 1991 Aug DISC

>D> AP

EQ 500MG BASE/VIAL

N63216 001 Dec 27, 1991 Aug DISC

>A> @

EQ 500MG BASE/VIAL

N63216 001 Dec 27, 1991 Aug DISC

CEFOXITIN SODIUM

INJECTABLE; INJECTION

CEFOXITIN

AP ACS DOBFAR

EQ 1GM BASE/VIAL

N65414 001 Jun 12, 2009 Jun NEWA

AP EQ 2GM BASE/VIAL

N65414 002 Jun 12, 2009 Jun NEWA

CEFTAZIDIME

INJECTABLE; INJECTION

TAZICEF

AP HOSPIRA

1GM/VIAL

N64032 001 Oct 31, 1993 Jul CMFD

AP 2GM/VIAL

N64032 002 Oct 31, 1993 Jul CMFD

CEFTRIAXONE SODIUM

INJECTABLE; IM-IV

CEFTRIAXONE

AP + SANDOZ

EQ 250MG BASE/VIAL

N65169 001 May 09, 2005 Jul CRLD

AP + EQ 500MG BASE/VIAL

N65169 002 May 09, 2005 Jul CRLD

AP + EQ 1GM BASE/VIAL

N65169 003 May 09, 2005 Jul CRLD

AP + EQ 2GM BASE/VIAL

N65169 004 May 09, 2005 Mar CRLD

ROCEPHIN

@ HOFFMANN LA ROCHE

EQ 250MG BASE/VIAL

N50585 001 Dec 21, 1984 Jul CAHN

@ EQ 500MG BASE/VIAL

N50585 002 Dec 21, 1984 Jul CAHN

@ EQ 1GM BASE/VIAL

N50585 003 Dec 21, 1984 Jul CAHN

@ EQ 2GM BASE/VIAL

N50585 004 Dec 21, 1984 Jul CAHN

INJECTABLE; INJECTION

ROCEPHIN

@ HOFFMANN LA ROCHE

EQ 10GM BASE/VIAL

N50585 005 Dec 21, 1984 Jul CAHN

ROCEPHIN W/ DEXTROSE IN PLASTIC CONTAINER

@ HOFFMANN LA ROCHE

EQ 10MG BASE/ML

N50624 001 Feb 11, 1987 Jul CAHN

@ EQ 20MG BASE/ML

N50624 002 Feb 11, 1987 Jul CAHN

@ EQ 40MG BASE/ML

N50624 003 Feb 11, 1987 Jul CAHN

CEFTRIAXONE SODIUM; LIDOCAINE

INJECTABLE; INJECTION

ROCEPHIN KIT

@ HOFFMANN LA ROCHE

EQ 500MG BASE/VIAL, N/A; N/A, 1%

N50585 007 May 08, 1996 Jul CAHN

@ EQ 1GM BASE/VIAL, N/A; N/A, 1%

N50585 006 May 08, 1996 Jul CAHN

CEPHALEXIN

FOR SUSPENSION; ORAL

CEPHALEXIN

>A> AB HIKMA PHARMS

EQ 125MG BASE/5ML

N65444 001 Aug 28, 2009 Aug NEWA

		FOR SUSPENSION; ORAL					
		CEPHALEXIN					
>A>	AB	HIKMA PHARMS	EQ 250MG BASE/5ML	N65444 002	Aug 28, 2009	Aug	NEWA
<u>CETIRIZINE HYDROCHLORIDE</u>							
		SYRUP; ORAL					
		CETIRIZINE HYDROCHLORIDE					
	AA	DR REDDYS LABS LTD	5MG/5ML	N78870 001	Apr 27, 2009	Apr	NEWA
>A>	AA	WOCKHARDT	5MG/5ML	N78757 001	Aug 28, 2009	Aug	NEWA
<u>CHLORHEXIDINE GLUCONATE</u>							
		SOLUTION; DENTAL					
		CHLORHEXIDINE GLUCONATE					
	AT	XTTRIUM	0.12%	N77789 001	Jun 18, 2009	Jun	NEWA
<u>CHLOROTHIAZIDE</u>							
		TABLET; ORAL					
		DIURIL					
		@ LUNDBECK INC	250MG	N11145 004		Apr	CAHN
		@	500MG	N11145 002		Apr	CAHN
<u>CHLOROTHIAZIDE SODIUM</u>							
		INJECTABLE; INJECTION					
		DIURIL					
		+ LUNDBECK INC	EQ 500MG BASE/VIAL	N11145 005		Apr	CAHN
<u>CHLORPROPAMIDE</u>							
		TABLET; ORAL					
		CHLORPROPAMIDE					
		@ SANDOZ	100MG	N88725 001	Aug 31, 1984	Mar	DISC
		@	250MG	N88726 001	Aug 31, 1984	Mar	DISC
<u>CHLORZOXAZONE</u>							
		TABLET; ORAL					
		CHLORZOXAZONE					
		@ SANDOZ	250MG	N89852 001	May 04, 1988	Mar	DISC
		@	500MG	N89853 001	May 04, 1988	Mar	DISC
	AA	WATSON LABS	500MG	N89859 001	May 04, 1988	Apr	CAHN
<u>CHOLINE FENOFIBRATE</u>							
		CAPSULE, DELAYED RELEASE; ORAL					
		TRILIPIX					
		ABBOTT LABS	EQ 45MG FENOFIBRIC ACID	N22224 001	Dec 15, 2008	Jan	CTNA
		+	EQ 135MG FENOFIBRIC ACID	N22224 002	Dec 15, 2008	Jan	CTNA
<u>CHYMOPAPAIN</u>							
		INJECTABLE; INJECTION					
		CHYMODIACTIN					
		@ CHART MEDCL	4,000 UNITS/VIAL	N18663 002	Aug 21, 1984	Mar	CAHN
		@	10,000 UNITS/VIAL	N18663 001	Nov 10, 1982	Mar	CAHN
<u>CIMETIDINE</u>							
		TABLET; ORAL					
		CIMETIDINE					
>D>	AB	LEK PHARMS	300MG	N74250 002	Jun 29, 1995	Aug	DISC
>A>		@	300MG	N74250 002	Jun 29, 1995	Aug	DISC

TABLET; ORAL

CIMETIDINE

>D>	AB	LEK PHARMS	400MG	N74250 003	Jun 29, 1995	Aug	DISC
>A>		@	400MG	N74250 003	Jun 29, 1995	Aug	DISC
>D>	AB		800MG	N74250 004	Jun 29, 1995	Aug	DISC
>A>		@	800MG	N74250 004	Jun 29, 1995	Aug	DISC
		@ SANDOZ	200MG	N74100 001	Jan 31, 1995	Mar	DISC
		@	300MG	N74100 002	Jan 31, 1995	Mar	DISC
		@	400MG	N74100 003	Jan 31, 1995	Mar	DISC
		@	800MG	N74100 004	Jan 31, 1995	Mar	DISC

CIPROFLOXACIN

INJECTABLE; INJECTION

CIPROFLOXACIN

>D>	AP	BEDFORD LABS	200MG/20ML (10MG/ML)	N76992 001	Aug 28, 2006	Aug	DISC
>D>	AP		400MG/40ML (10MG/ML)	N76992 002	Aug 28, 2006	Aug	DISC
>A>		@	400MG/40ML (10MG/ML)	N76992 002	Aug 28, 2006	Aug	DISC
>A>		@	200MG/20ML (10MG/ML)	N76992 001	Aug 28, 2006	Aug	DISC
>D>			1200MG/120ML (10MG/ML)	N76993 001	Aug 28, 2006	Aug	DISC
>A>		@	1200MG/120ML (10MG/ML)	N76993 001	Aug 28, 2006	Aug	DISC

CIPROFLOXACIN IN DEXTROSE 5% IN PLASTIC CONTAINER

>D>	AP	BEDFORD	200MG/100ML	N78114 001	Mar 18, 2008	Aug	DISC
>A>		@	200MG/100ML	N78114 001	Mar 18, 2008	Aug	DISC
>D>	AP		400MG/200ML	N78114 002	Mar 18, 2008	Aug	DISC
>A>		@	400MG/200ML	N78114 002	Mar 18, 2008	Aug	DISC

CIPROFLOXACIN HYDROCHLORIDE

SOLUTION/DROPS; OTIC

CETRAXAL

+	LABORATORIOS SALVAT	EQ 0.2% BASE	N21918 001	May 01, 2009	May	NEWA
+	WRASER PHARMS	EQ 0.2% BASE	N21918 001	May 01, 2009	Jul	CAHN

TABLET; ORAL

CIPROFLOXACIN HYDROCHLORIDE

AB	MYLAN	EQ 250MG BASE	N75685 002	Jun 09, 2004	Mar	CMFD
AB		EQ 500MG BASE	N75685 003	Jun 09, 2004	Mar	CMFD
AB		EQ 750MG BASE	N75685 001	Jun 09, 2004	Mar	CMFD
	@ TEVA	EQ 250MG BASE	N76136 001	Jun 09, 2004	Jan	DISC
	@	EQ 500MG BASE	N76136 002	Jun 09, 2004	Jan	DISC
	@	EQ 750MG BASE	N76136 003	Jun 09, 2004	Jan	DISC

CITALOPRAM HYDROBROMIDE

TABLET; ORAL

CITALOPRAM HYDROBROMIDE

>D>	AB	AMNEAL PHARMS	EQ 10MG BASE	N77045 003	Apr 29, 2005	Aug	DISC
>A>		@	EQ 10MG BASE	N77045 003	Apr 29, 2005	Aug	DISC
	AB		EQ 10MG BASE	N77045 003	Apr 29, 2005	Feb	CAHN
>D>	AB		EQ 20MG BASE	N77045 002	Apr 29, 2005	Aug	DISC
>A>		@	EQ 20MG BASE	N77045 002	Apr 29, 2005	Aug	DISC
	AB		EQ 20MG BASE	N77045 002	Apr 29, 2005	Feb	CAHN
>D>	AB		EQ 40MG BASE	N77045 001	Apr 29, 2005	Aug	DISC
>A>		@	EQ 40MG BASE	N77045 001	Apr 29, 2005	Aug	DISC
	AB		EQ 40MG BASE	N77045 001	Apr 29, 2005	Feb	CAHN
	AB	GLENMARK GENERICS	EQ 10MG BASE	N77654 001	Feb 27, 2009	Feb	NEWA
	AB		EQ 20MG BASE	N77654 002	Feb 27, 2009	Feb	NEWA
	AB		EQ 40MG BASE	N77654 003	Feb 27, 2009	Feb	NEWA

CLARITHROMYCIN

TABLET, EXTENDED RELEASE; ORAL

CLARITHROMYCIN

@ RANBAXY

1GM

N65210 001 Jan 26, 2005 Jul DISC

CLINDAMYCIN HYDROCHLORIDE

CAPSULE; ORAL

CLINDAMYCIN HYDROCHLORIDE

>A> AB AUROBINDO PHARMA EQ 150MG BASE

N65442 001 Aug 26, 2009 Aug NEWA

>A> AB EQ 300MG BASE

N65442 002 Aug 26, 2009 Aug NEWA

CLOCORTOLONE PIVALATE

CREAM; TOPICAL

CLODERM

+ DOW PHARM SCIENCES 0.1%

N17765 001 Jun CAHN

>D> CLOFIBRATE

>D> CAPSULE; ORAL

>D> CLOFIBRATE

>D> + USL PHARMA 500MG

N70531 001 Jun 16, 1986 Aug DISC

>A> @ 500MG

N70531 001 Jun 16, 1986 Aug DISC

CLOMIPRAMINE HYDROCHLORIDE

CAPSULE; ORAL

CLOMIPRAMINE HYDROCHLORIDE

>D> AB SANDOZ 25MG

N74953 001 Jun 25, 1997 Aug DISC

>A> @ 25MG

N74953 001 Jun 25, 1997 Aug DISC

>D> AB 50MG

N74953 002 Jun 25, 1997 Aug DISC

>A> @ 50MG

N74953 002 Jun 25, 1997 Aug DISC

>D> AB 75MG

N74953 003 Jun 25, 1997 Aug DISC

>A> @ 75MG

N74953 003 Jun 25, 1997 Aug DISC

CLONAZEPAM

TABLET; ORAL

CLONAZEPAM

@ KALI LABS

0.5MG

N77147 001 May 02, 2005 Jul DISC

@

1MG

N77147 002 May 02, 2005 Jul DISC

@

2MG

N77147 003 May 02, 2005 Jul DISC

CLONIDINE

FILM, EXTENDED RELEASE; TRANSDERMAL

CATAPRES-TTS-1

>D> BOEHRINGER INGELHEIM 0.1MG/24HR

N18891 001 Oct 10, 1984 Aug CFTG

>A> AB 0.1MG/24HR

N18891 001 Oct 10, 1984 Aug CFTG

CATAPRES-TTS-2

>D> BOEHRINGER INGELHEIM 0.2MG/24HR

N18891 002 Oct 10, 1984 Aug CFTG

>A> AB 0.2MG/24HR

N18891 002 Oct 10, 1984 Aug CFTG

CATAPRES-TTS-3

>D> + BOEHRINGER INGELHEIM 0.3MG/24HR

N18891 003 Oct 10, 1984 Aug CFTG

>A> AB + 0.3MG/24HR

N18891 003 Oct 10, 1984 Aug CFTG

CLONIDINE

>A> AB AVEVA 0.1MG/24HR

N76157 001 Aug 18, 2009 Aug NEWA

>A> AB 0.2MG/24HR

N76157 002 Aug 18, 2009 Aug NEWA

>A> AB 0.3MG/24HR

N76157 003 Aug 18, 2009 Aug NEWA

CLONIDINE HYDROCHLORIDE

TABLET; ORAL

CLONIDINE HYDROCHLORIDE

>A>	AB	IMPAX LABS	0.1MG	N78099 001	Aug 27, 2009	Aug	NEWA
>A>	AB		0.2MG	N78099 002	Aug 27, 2009	Aug	NEWA
>A>	AB		0.3MG	N78099 003	Aug 27, 2009	Aug	NEWA
		@ SANDOZ	0.1MG	N70887 001	Aug 31, 1988	Mar	DISC
		@	0.2MG	N70886 001	Aug 31, 1988	Mar	DISC
		@	0.3MG	N71294 001	Aug 31, 1988	Mar	DISC
>A>	AB	UNICHEM	0.1MG	N78895 001	Aug 26, 2009	Aug	NEWA
>A>			0.2MG	N78895 002	Aug 26, 2009	Aug	NEWA
>A>	AB		0.3MG	N78895 003	Aug 26, 2009	Aug	NEWA
>D>	AB	WATSON LABS	0.1MG	N70965 001	Jul 08, 1986	Aug	DISC
>A>		@	0.1MG	N70965 001	Jul 08, 1986	Aug	DISC
>D>	AB		0.2MG	N70964 001	Jul 08, 1986	Aug	DISC
>A>		@	0.2MG	N70964 001	Jul 08, 1986	Aug	DISC

CLOPIDOGREL BISULFATE

TABLET; ORAL

CLOPIDOGREL BISULFATE

	@ DR REDDYS LABS INC	EQ 75MG BASE	N76273 001	Jan 14, 2008	Jun	DISC
	PLAVIX					
	SANOFI AVENTIS US	EQ 75MG BASE	N20839 001	Nov 17, 1997	Jun	CTEC

CLORAZEPATE DIPOTASSIUM

CAPSULE; ORAL

CLORAZEPATE DIPOTASSIUM

	@ SANDOZ	3.75MG	N72219 001	Aug 26, 1988	Mar	DISC
	TRANXENE					
	@ LUNDBECK INC	3.75MG	N17105 001		Apr	CAHN
	@	7.5MG	N17105 002		Apr	CAHN
	@	15MG	N17105 003		Apr	CAHN

TABLET; ORAL

CLORAZEPATE DIPOTASSIUM

	@ SANDOZ	7.5MG	N72513 001	May 11, 1990	Mar	DISC
	@	15MG	N72514 001	May 11, 1990	Mar	DISC
	TRANXENE					
AB	LUNDBECK INC	3.75MG	N17105 006		Apr	CAHN
AB		7.5MG	N17105 007		Apr	CAHN
AB	+	15MG	N17105 008		Apr	CAHN
	TRANXENE SD					
	@ LUNDBECK INC	11.25MG	N17105 005		May	DISC
		11.25MG	N17105 005		Apr	CAHN
	@	22.5MG	N17105 004		May	DISC
	+	22.5MG	N17105 004		Apr	CAHN

CLOXACILLIN SODIUM

CAPSULE; ORAL

CLOXAPEN

	@ GLAXOSMITHKLINE	EQ 250MG BASE	N61806 001		Jun	DISC
	@	EQ 500MG BASE	N61806 002		Jun	DISC

CODEINE PHOSPHATE; PSEUDOEPHEDRINE HYDROCHLORIDE; TRIPROLIDINE HYDROCHLORIDE

SYRUP; ORAL

TRIACIN-C

+	STI PHARMA LLC	10MG/5ML;30MG/5ML;1.25MG/5ML	N88704 001	Mar 22, 1985	Apr	CAHN
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CODEINE SULFATE

TABLET; ORAL

CODEINE SULFATE

ROXANE

		15MG	N22402 001	Jul 16, 2009	Jul	NEWA
		30MG	N22402 002	Jul 16, 2009	Jul	NEWA
+		60MG	N22402 003	Jul 16, 2009	Jul	NEWA

COLCHICINE

TABLET; ORAL

COLCRYS

+	AR HOLDING CO INC	0.6MG	N22352 001	Jul 29, 2009	Jul	CAHN
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COLISTIMETHATE SODIUM

INJECTABLE; INJECTION

COLISTIMETHATE SODIUM

AP	PADDOCK	EQ 150MG BASE/VIAL	N65177 001	Mar 19, 2004	Apr	CTNA
AP	X GEN PHARMS	EQ 150MG BASE/VIAL	N64216 001	Feb 26, 1999	Apr	CTNA

CONIVAPTAN HYDROCHLORIDE

INJECTABLE; IV (INFUSION)

>D> VAPRISOL

>D>	+	ASTELLAS	20MG/4ML (5MG/ML)	N21697 001	Dec 29, 2005	Aug	DISC
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>A>	@		20MG/4ML (5MG/ML)	N21697 001	Dec 29, 2005	Aug	DISC
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CROMOLYN SODIUM

SOLUTION; INHALATION

CROMOLYN SODIUM

>D>	AN	PHARMASCIENCE INC	10MG/ML	N75437 001	Apr 21, 2000	Aug	DISC
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>A>	@		10MG/ML	N75437 001	Apr 21, 2000	Aug	DISC
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AN	+	TEVA PARENTERAL	10MG/ML	N75271 001	Jan 18, 2000	Apr	CAHN
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CYANOCOBALAMIN

GEL, METERED; NASAL

NASCOBAL

@ PAR PHARM

0.5MG/INH

N19722 001	Nov 05, 1996	Mar	CAHN
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SPRAY, METERED; NASAL

NASCOBAL

+	PAR PHARM CO	0.5MG/SPRAY	N21642 001	Jan 31, 2005	May	CAHN
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CYCLOBENZAPRINE HYDROCHLORIDE

TABLET; ORAL

CYCLOBENZAPRINE HYDROCHLORIDE

>D>	AB	AMNEAL PHARM	10MG	N78218 001	Apr 18, 2008	Aug	CAHN
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>A>	AB	ORIT LABS LLC	10MG	N78218 001	Apr 18, 2008	Aug	CAHN
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CYCLOPHOSPHAMIDE

INJECTABLE; INJECTION

CYTOXAN

@ BAXTER HLTHCARE

100MG/VIAL

N12142 001	Feb	CAHN
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200MG/VIAL

N12142 002	Feb	CAHN
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INJECTABLE; INJECTION

CYTOXAN

@ BAXTER HLTHCARE	500MG/VIAL	N12142 003		Feb	CAHN
@	1GM/VIAL	N12142 004	Aug 30, 1982	Feb	CAHN
@	2GM/VIAL	N12142 005	Aug 30, 1982	Feb	CAHN

LYOPHILIZED CYTOXAN

+	BAXTER HLTHCARE	100MG/VIAL	N12142 006	Dec 05, 1985	Feb	CAHN
+		200MG/VIAL	N12142 007	Dec 10, 1985	Feb	CAHN
AP	+	500MG/VIAL	N12142 008	Jan 04, 1984	Feb	CAHN
AP	+	1GM/VIAL	N12142 010	Sep 24, 1985	Feb	CAHN
AP	+	2GM/VIAL	N12142 009	Dec 10, 1984	Feb	CAHN

TABLET; ORAL

CYTOXAN

@ BAXTER HLTHCARE	25MG	N12141 002		Feb	CAHN
@	50MG	N12141 001		Feb	CAHN

DACTINOMYCININJECTABLE; INJECTION

COSMEGEN

+	LUNDBECK INC	0.5MG/VIAL	N50682 001		Apr	CAHN
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DARUNAVIR ETHANOLATETABLET; ORAL

PREZISTA

	CENTOCOR ORTHO	EQ 75MG BASE	N21976 004	Dec 18, 2008	Jul	CAHN
		EQ 150MG BASE	N21976 005	Dec 18, 2008	Jul	CAHN
		EQ 300MG BASE	N21976 001	Jun 23, 2006	Jul	CAHN
		EQ 400MG BASE	N21976 003	Oct 21, 2008	Jul	CAHN
+		EQ 600MG BASE	N21976 002	Feb 25, 2008	Jul	CAHN
	TIBOTEC	EQ 150MG BASE	N21976 005	Dec 18, 2008	Jun	CMFD

DAUNORUBICIN CITRATEINJECTABLE, LIPOSOMAL; INJECTION

DAUNOXOME

@ DIATOS	EQ 2MG BASE/ML	N50704 002	Apr 08, 1996	Jul	DISC
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DEFEROXAMINE MESYLATEINJECTABLE; INJECTION

DEFEROXAMINE MESYLATE

AP	WATSON LABS	500MG/VIAL	N76806 001	Mar 31, 2006	Apr	CAHN
AP		2GM/VIAL	N76806 002	Mar 31, 2006	Apr	CAHN

DEGARELIX ACETATEPOWDER; SUBCUTANEOUS

FIRMAGON

	FERRING	EQ 80MG BASE/VIAL	N22201 001	Dec 24, 2008	Jun	CTNA
+		EQ 120MG BASE/VIAL	N22201 002	Dec 24, 2008	Jun	CTNA

DESOGESTREL; ETHINYL ESTRADIOLTABLET; ORAL-21

DESOGESTREL AND ETHINYL ESTRADIOL

@ DURAMED PHARMS BARR	0.15MG;0.03MG	N75256 001	Aug 12, 1999	Feb	DISC
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DESVENLAFAXINE SUCCINATE

TABLET, EXTENDED RELEASE; ORAL
PRISTIQ

+ WYETH PHARMS INC EQ 50MG BASE N21992 001 Feb 29, 2008 Jul CRLD

DEXAMETHASONE

ELIXIR; ORAL
DEXAMETHASONE

AA + STI PHARMA LLC 0.5MG/5ML N84754 001 Apr CAHN

IMPLANT; INTRAVITREAL
OZURDEX

+ ALLERGAN 0.7MG N22315 001 Jun 17, 2009 Jun NEWA

DEXAMETHASONE; TOBRAMYCIN

SUSPENSION/DROPS; OPHTHALMIC
TOBRADEX ST

+ ALCON 0.05%;0.3% N50818 001 Feb 13, 2009 Feb NEWA

DEXLANSOPRAZOLE

CAPSULE, DELAYED RELEASE; ORAL
KAPIDEX

+ TAKEDA PHARMS 30MG N22287 001 Jan 30, 2009 Jan NEWA

+ 60MG N22287 002 Jan 30, 2009 Jan NEWA

DEXTROAMPHETAMINE SULFATE

TABLET; ORAL
DEXTROAMPHETAMINE SULFATE

AA + BARR 10MG N40361 002 Jan 31, 2001 Mar CRLD

DEXTROSTAT

@ SHIRE 5MG N84051 001 Mar DISC

@ 10MG N84051 002 Mar DISC

DEXTROSE

INJECTABLE; INJECTION
DEXTROSE 60% IN PLASTIC CONTAINER
@ HOSPIRA 60GM/100ML

N19346 001 Jan 25, 1985 Apr DISC

DIATRIZOATE MEGLUMINE

INJECTABLE; INJECTION
RENO-60

@ BRACCO 60% N10040 016 Jun DISC

RENO-DIP

@ BRACCO 30% N10040 012 Jun DISC

SOLUTION; URETERAL

RENO-30

@ BRACCO 30% N10040 021 Jun DISC

DIATRIZOATE MEGLUMINE; DIATRIZOATE SODIUM

INJECTABLE; INJECTION
RENOGRAFIN-60

@ BRACCO 52%;8% N10040 006 Jun DISC

DIAZEPAM

TABLET; ORAL

DIAZEPAM

@ SANDOZ	2MG	N70302 001	Dec 20, 1985	Mar	DISC
@	5MG	N70303 001	Dec 20, 1985	Mar	DISC
@	10MG	N70304 001	Dec 20, 1985	Mar	DISC

DICLOFENAC POTASSIUM

CAPSULE; ORAL

ZIPSOR

+ XANODYNE PHARM	25MG	N22202 001	Jun 18, 2009	Jun	NEWA
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FOR SOLUTION; ORAL

CAMBIA

+ KOWA PHARMS	50MG	N22165 001	Jun 17, 2009	Jun	NEWA
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DIETHYLPROPION HYDROCHLORIDE

TABLET; ORAL

DIETHYLPROPION HYDROCHLORIDE

AA	COREPHARMA	25MG	N40828 001	Nov 05, 2008	Feb	CTEC
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TENUATE

AA	+ WATSON PHARMS	25MG	N11722 002		Feb	CTEC
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DIFLUNISAL

TABLET; ORAL

DIFLUNISAL

@ SANDOZ	500MG	N74604 001	Jun 10, 1996	Mar	DISC
+ TEVA	500MG	N73673 001	Jul 31, 1992	Mar	CTEC

DIGOXIN

INJECTABLE; INJECTION

DIGOXIN

@ HOSPIRA	0.25MG/ML	N40206 001	Aug 28, 1998	Apr	DISC
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TABLET; ORAL

DIGOXIN

AB	IMPAX LABS	0.125MG	N78556 001	Jul 20, 2009	Jun	NEWA
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AB		0.25MG	N78556 002	Jul 20, 2009	Jun	NEWA
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DILTIAZEM HYDROCHLORIDE

CAPSULE, EXTENDED RELEASE; ORAL

DILT-CD

AB3	APOTEX	300MG	N76151 004	May 20, 2004	Apr	CAHN
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DILTIAZEM HYDROCHLORIDE

BC	+ MYLAN	120MG	N74910 003	May 02, 1997	Jul	CTEC
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DIPHENHYDRAMINE HYDROCHLORIDE

CAPSULE; ORAL

DIPHENHYDRAMINE HYDROCHLORIDE

+ BARR	50MG	N80738 001		Mar	CRLD
@ LNK	25MG	N87977 001	Jan 27, 1983	Mar	DISC
@	50MG	N87978 001	Jan 27, 1983	Mar	DISC
@ SANDOZ	25MG	N80832 001		Mar	DISC
@	50MG	N80832 002		Mar	DISC
@ VALEANT PHARM INTL	50MG	N80592 001		Mar	DISC
@ WATSON LABS	25MG	N80728 001		Mar	DISC
@	50MG	N80727 001		Mar	DISC

DIPYRIDAMOLE

TABLET; ORAL

DIPYRIDAMOLE

@ SANDOZ	25MG	N86944 002	Apr 16, 1991	Mar	DISC
@	50MG	N87562 001	Feb 25, 1992	Mar	DISC
@	75MG	N87561 001	Feb 25, 1992	Mar	DISC

DIVALPROEX SODIUM

CAPSULE, DELAYED REL PELLETS; ORAL

DEPAKOTE

AB	+	ABBOTT	EQ 125MG VALPROIC ACID	N19680 001	Sep 12, 1989	Jan	CFTG
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DIVALPROEX SODIUM

AB		DR REDDYS LABS LTD	EQ 125MG VALPROIC ACID	N78979 001	Jan 23, 2009	Jan	NEWA
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AB		ZYDUS PHARMS USA INC	EQ 125MG VALPROIC ACID	N78919 001	Jan 27, 2009	Jan	NEWA
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TABLET, DELAYED RELEASE; ORAL

DIVALPROEX SODIUM

AB		MYLAN	EQ 125MG VALPROIC ACID	N90062 001	Mar 17, 2009	Mar	NEWA
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AB			EQ 250MG VALPROIC ACID	N90062 002	Mar 17, 2009	Mar	NEWA
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AB			EQ 500MG VALPROIC ACID	N90062 003	Mar 17, 2009	Mar	NEWA
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AB		ZYDUS PHARMS USA INC	EQ 125MG VALPROIC ACID	N77100 001	Mar 05, 2009	Feb	NEWA
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AB			EQ 250MG VALPROIC ACID	N77100 002	Mar 05, 2009	Feb	NEWA
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AB			EQ 500MG VALPROIC ACID	N77100 003	Mar 05, 2009	Feb	NEWA
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TABLET, EXTENDED RELEASE; ORAL

DEPAKOTE ER

AB		ABBOTT	EQ 250MG VALPROIC ACID	N21168 002	May 31, 2002	Jan	CFTG
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AB	+		EQ 500MG VALPROIC ACID	N21168 001	Aug 04, 2000	Jan	CFTG
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DIVALPROEX SODIUM

AB		ANCHEN PHARMS	EQ 250MG VALPROIC ACID	N78445 001	Feb 26, 2009	Feb	NEWA
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AB			EQ 500MG VALPROIC ACID	N78445 002	Aug 04, 2009	Jul	NEWA
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AB		IMPAX LABS	EQ 250MG VALPROIC ACID	N78791 001	May 06, 2009	Apr	NEWA
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AB			EQ 500MG VALPROIC ACID	N78791 002	Aug 04, 2009	Jul	NEWA
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AB		MYLAN	EQ 250MG VALPROIC ACID	N77567 001	Jan 29, 2009	Jan	NEWA
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AB			EQ 500MG VALPROIC ACID	N77567 002	Jan 29, 2009	Jan	NEWA
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AB		TEVA PHARMS	EQ 500MG VALPROIC ACID	N78700 001	Aug 03, 2009	Jul	NEWA
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AB		WOCKHARDT	EQ 250MG VALPROIC ACID	N78705 002	Feb 10, 2009	Jan	NEWA
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AB			EQ 500MG VALPROIC ACID	N78705 001	Aug 04, 2009	Jul	NEWA
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AB		ZYDUS PHARMS USA INC	EQ 250MG VALPROIC ACID	N78239 001	Feb 27, 2009	Feb	NEWA
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AB			EQ 500MG VALPROIC ACID	N78239 002	Aug 04, 2009	Jul	NEWA
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DOBUTAMINE HYDROCHLORIDE

INJECTABLE; INJECTION

DOBUTAMINE HYDROCHLORIDE

>D>	AP	LUITPOLD	EQ 12.5MG BASE/ML	N74545 001	Jun 25, 1998	Aug	DISC
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>A>		@	EQ 12.5MG BASE/ML	N74545 001	Jun 25, 1998	Aug	DISC
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DORIPENEM

INJECTABLE; IV (INFUSION)

DORIBAX

	+	ORTHO MCNEIL JANSSEN	500MG/VIAL	N22106 001	Oct 12, 2007	Jun	CAHN
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DORZOLAMIDE HYDROCHLORIDE

SOLUTION/DROPS; OPHTHALMIC

DORZOLAMIDE HYDROCHLORIDE

AT		ALCON	EQ 2% BASE	N78981 001	Apr 13, 2009	Mar	NEWA
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AT		BAUSCH AND LOMB	EQ 2% BASE	N90143 001	Jun 25, 2009	Jun	NEWA
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DORZOLAMIDE HYDROCHLORIDE; TIMOLOL MALEATE

SOLUTION/DROPS; OPHTHALMIC

DORZOLAMIDE HYDROCHLORIDE AND TIMOLOL MALEATE

AT	BAUSCH AND LOMB	EQ 2% BASE;EQ 0.5% BASE	N90037 001	Jul 14, 2009	Jun	NEWA
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DOXAZOSIN MESYLATE

TABLET; ORAL

DOXAZOSIN MESYLATE

>D>	AB	SANDOZ	EQ 1MG BASE	N75646 001	Oct 18, 2000	Aug	DISC
>A>		@	EQ 1MG BASE	N75646 001	Oct 18, 2000	Aug	DISC
>D>	AB		EQ 2MG BASE	N75646 002	Oct 18, 2000	Aug	DISC
>A>		@	EQ 2MG BASE	N75646 002	Oct 18, 2000	Aug	DISC
>D>	AB		EQ 4MG BASE	N75646 003	Oct 18, 2000	Aug	DISC
>A>		@	EQ 4MG BASE	N75646 003	Oct 18, 2000	Aug	DISC
>D>	AB		EQ 8MG BASE	N75646 004	Oct 18, 2000	Aug	DISC
>A>		@	EQ 8MG BASE	N75646 004	Oct 18, 2000	Aug	DISC

DOXERCALCIFEROL

CAPSULE; ORAL

HECTOROL

GENZYME

1UGM

N20862 003	Jul 13, 2009	Jul	NEWA
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DOXYCYCLINE

CAPSULE; ORAL

DOXYCYCLINE

+ PAR PHARM

EQ 150MG BASE

N65055 003	Jul 15, 2005	Jan	CRLD
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TABLET; ORAL

DOXYCYCLINE

AB MUTUAL PHARM

EQ 50MG BASE

N65471 001	Apr 17, 2009	Mar	NEWA
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AB

EQ 75MG BASE

N65471 002	Apr 17, 2009	Mar	NEWA
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AB

EQ 100MG BASE

N65471 003	Apr 17, 2009	Mar	NEWA
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DOXYCYCLINE HYCLATE

CAPSULE; ORAL

DOXYCYCLINE HYCLATE

>D>	AB	WATSON LABS	EQ 50MG BASE	N61717 001		Aug	DISC
>A>		@	EQ 50MG BASE	N61717 001		Aug	DISC
>D>	AB		EQ 100MG BASE	N61717 002		Aug	DISC
>A>		@	EQ 100MG BASE	N61717 002		Aug	DISC

DRONABINOL

CAPSULE; ORAL

DRONABINOL

AB SVC PHARMA

2.5MG

N78292 001	Jun 27, 2008	Mar	CAHN
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AB

5MG

N78292 002	Jun 27, 2008	Mar	CAHN
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AB

10MG

N78292 003	Jun 27, 2008	Mar	CAHN
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DRONEDARONE HYDROCHLORIDE

TABLET; ORAL

MULTAQ

+ SANOFI AVENTIS US

EQ 400MG BASE

N22425 001	Jul 01, 2009	Jul	NEWA
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DROSPIRENONE; ETHINYL ESTRADIOL

TABLET; ORAL

DROSPIRENONE AND ETHINYL ESTRADIOL

AB BARR 3MG;0.02MG N78515 001 Mar 30, 2009 Mar NEWA

YAZ

AB + BAYER HLTHCARE 3MG;0.02MG N21676 001 Mar 16, 2006 Mar CFTG

EDROPHONIUM CHLORIDE

INJECTABLE; INJECTION

ENLON

AP BIONICHE PHARMA 10MG/ML N88873 001 Aug 06, 1985 Jun CMFD

EFAVIRENZ

CAPSULE; ORAL

SUSTIVA

@ BRISTOL MYERS SQUIBB 100MG

N20972 002 Sep 17, 1998 Jun DISC

ENALAPRIL MALEATE

TABLET; ORAL

ENALAPRIL MALEATE

@ SANDOZ 2.5MG

N75048 001 Aug 22, 2000 Mar DISC

@ 5MG

N75048 002 Aug 22, 2000 Mar DISC

@ 10MG

N75048 003 Aug 22, 2000 Mar DISC

@ 20MG

N75048 004 Aug 22, 2000 Mar DISC

ENALAPRIL MALEATE; FELODIPINE

TABLET, EXTENDED RELEASE; ORAL

LEXXEL

@ ASTRAZENECA 5MG;2.5MG

N20668 002 Oct 28, 1998 May DISC

ENALAPRILAT

INJECTABLE; INJECTION

ENALAPRILAT

AP + HOSPIRA 1.25MG/ML N75458 001 Aug 22, 2000 Feb CRLD

VASOTEC

@ BIOVAIL LABS INTL 1.25MG/ML

N19309 001 Feb 09, 1988 Feb DISC

EPINEPHRINE

INJECTABLE; IM-SC

TWINJECT 0.15

+ SCIELE PHARMA INC EQ 0.15MG /DELIVERY

N20800 002 May 28, 2004 Jun CAHN

TWINJECT 0.3

+ SCIELE PHARMA INC EQ 0.3MG /DELIVERY

N20800 001 May 30, 2003 Jun CAHN

EPINEPHRINE BITARTRATE; PRILOCAINE HYDROCHLORIDE

INJECTABLE; INJECTION

CITANEST FORTE DENTAL

+ DENTSPLY PHARM 0.005MG/ML;4%

N21383 001 Apr CTNA

EPINEPHRINE; LIDOCAINE HYDROCHLORIDE

INJECTABLE; INJECTION

XYLOCAINE DENTAL WITH EPINEPHRINE

AP + DENTSPLY PHARM 0.01MG/ML;2%

N21381 001 Apr CTNA

AP + 0.02MG/ML;2%

N21381 002 Apr CTNA

INJECTABLE; INJECTION

XYLOCAINE W/ EPINEPHRINE

AP	+	APP PHARMS	0.02MG/ML;2%	N06488 005		Apr	CMFD
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EPIRUBICIN HYDROCHLORIDE

INJECTABLE; INJECTION

EPIRUBICIN HYDROCHLORIDE

AP		AKORN INC	50MG/25ML (2MG/ML)	N90163 001	Jun 24, 2009	Jun	NEWA
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EPOPROSTENOL SODIUM

INJECTABLE; INJECTION

EPOPROSTENOL SODIUM

	+	ACTELION	EQ 1.5MG BASE/VIAL	N22260 001	Jun 27, 2008	Apr	CAHN
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ERGOCALCIFEROL

CAPSULE; ORAL

ERGOCALCIFEROL

AA		ORIT LABS LLC	50,000 IU	N40833 001	May 20, 2009	May	NEWA
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ERYTHROMYCIN

OINTMENT; OPHTHALMIC

ERYTHROMYCIN

@ AKORN 0.5%

N64030 001	Jul 18, 1996	Jul	DISC
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OINTMENT; TOPICAL

AKNE-MYCIN

	+	DOW PHARM SCIENCES	2%	N50584 001	Jan 10, 1985	Jun	CAHN
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SOLUTION; TOPICAL

ERYDERM

>D>							
>D>	AT	ABBOTT	2%	N62290 001		Aug	DISC

>A>		@	2%	N62290 001		Aug	DISC
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SANSAC

	@	DOW PHARM SCIENCES	2%	N62522 001	Jan 24, 1985	Jul	CAHN
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TABLET, DELAYED RELEASE; ORAL

E-MYCIN

>D>							
>D>	AB	ABBOTT	250MG	N60272 001		Aug	DISC

>A>		@	250MG	N60272 001		Aug	DISC
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>D>	AB		333MG	N60272 002		Aug	DISC
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>A>		@	333MG	N60272 002		Aug	DISC
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ERYTHROMYCIN ESTOLATE

SUSPENSION; ORAL

ERYTHROMYCIN ESTOLATE

@ ALPHARMA US PHARMS EQ 125MG BASE/5ML

N62353 001	Nov 18, 1982	Apr	DISC
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@ EQ 250MG BASE/5ML

N62409 001	Dec 16, 1982	Apr	DISC
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ERYTHROMYCIN ETHYLSUCCINATE

SUSPENSION; ORAL

ERYTHROMYCIN ETHYLSUCCINATE

@ ALPHARMA US PHARMS EQ 200MG BASE/5ML

N62200 001		Apr	DISC
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@ EQ 400MG BASE/5ML

N62200 002		Apr	DISC
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ERYTHROMYCIN ETHYLSUCCINATE; SULFISOXAZOLE ACETYL

GRANULE; ORAL

ERYTHROMYCIN ETHYLSUCCINATE AND SULFISOXAZOLE ACETYL

	+	BARR	EQ 200MG BASE/5ML;EQ 600MG BASE/5ML	N62759 001	May 20, 1988	Jun	CMFD
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GRANULE; ORAL

ERYZOLE

@ ALRA

EQ 200MG BASE/5ML;EQ 600MG
BASE/5ML

N62758 001 Jun 15, 1988 Jan DISC

PEDIAZOLE

@ ROSS LABS

EQ 200MG BASE/5ML;EQ 600MG
BASE/5ML

N50529 001 Jan DISC

ERYTHROMYCIN LACTOBIONATE

INJECTABLE; INJECTION

ERYTHROCIN

@ HOSPIRA

EQ 1GM BASE/VIAL

N50182 003 Feb DISC

AP +

EQ 1GM BASE/VIAL

N62638 002 Oct 31, 1986 Feb CRLD

ERYTHROMYCIN LACTOBIONATE

>D> AP

BAXTER HLTHCARE

EQ 500MG BASE/VIAL

N62993 001 May 09, 1989 Aug DISC

>A>

@

EQ 500MG BASE/VIAL

N62993 001 May 09, 1989 Aug DISC

>D> AP

@

EQ 1GM BASE/VIAL

N62993 002 May 09, 1989 Aug DISC

>A>

@

EQ 1GM BASE/VIAL

N62993 002 May 09, 1989 Aug DISC

ESCITALOPRAM OXALATE

TABLET; ORAL

LEXAPRO

FOREST LABS

EQ 5MG BASE

N21323 001 Aug 14, 2002 Jul CTEC

EQ 10MG BASE

N21323 002 Aug 14, 2002 Jul CTEC

+

EQ 20MG BASE

N21323 003 Aug 14, 2002 Jul CTEC

ESTRADIOL

TABLET; ORAL

INNOFEM

@ NOVO NORDISK INC

0.5MG

N40312 001 Nov 19, 1999 Apr DISC

@

1MG

N40312 002 Nov 19, 1999 Apr DISC

@

2MG

N40312 003 Nov 19, 1999 Apr DISC

ESTRADIOL; NORGESTIMATE

TABLET; ORAL

PREFEST

>D> AB

+

DURAMED

1MG, 1MG; N/A, 0.09MG

N21040 001 Oct 22, 1999 Aug CAHN

>A> AB

+

DURAMED RES

1MG, 1MG; N/A, 0.09MG

N21040 001 Oct 22, 1999 Aug CAHN

ETHAMBUTOL HYDROCHLORIDE

TABLET; ORAL

ETHAMBUTOL HYDROCHLORIDE

AB

LUPIN

100MG

N78939 001 Jun 17, 2009 Jun NEWA

AB

@

400MG

N78939 002 Jun 17, 2009 Jun NEWA

MYAMBUTOL

AB

STI PHARMA LLC

100MG

N16320 001 Mar CAHN

@

200MG

N16320 002 Mar CAHN

AB

@

400MG

N16320 003 Mar CAHN

@

500MG

N16320 004 Mar CAHN

ETHINYL ESTRADIOL; ETHYNODIOL DIACETATE

TABLET; ORAL-21

DEMULEN 1/35-21

@ GD SEARLE LLC

0.035MG; 1MG

N18168 001 Apr DISC

DEMULEN 1/50-21

@ GD SEARLE LLC

0.05MG; 1MG

N16927 001 Apr DISC

TABLET; ORAL-21

ZOVIA 1/35E-21

@ WATSON LABS

0.035MG;1MG

N72720 001 Dec 30, 1991 Apr DISC

ZOVIA 1/50E-21

@ WATSON LABS

0.05MG;1MG

N72722 001 Dec 30, 1991 Apr DISC

TABLET; ORAL-28

DEMULEN 1/35-28

@ GD SEARLE LLC

0.035MG;1MG

N18160 001 Apr DISC

DEMULEN 1/50-28

@ GD SEARLE LLC

0.05MG;1MG

N16936 001 Apr DISC

ZOVIA 1/50E-28

+ WATSON LABS

0.05MG;1MG

N72723 001 Dec 30, 1991 Apr CRLD

ETHINYL ESTRADIOL; LEVONORGESTREL

TABLET; ORAL-21

TRIPHASIL-21

@ AKRIMAX PHARMS

0.03MG,0.04MG,0.03MG;0.05MG,0.075
MG,0.125MG

N19192 001 Nov 01, 1984 Apr CAHN

TABLET; ORAL-28

LEVLITE

>D>

>D> AB2 + BAYER HLTHCARE

0.02MG;0.1MG

N20860 002 Jul 13, 1998 Aug DISC

>A>

@

0.02MG;0.1MG

N20860 002 Jul 13, 1998 Aug DISC

LEVONORGESTREL AND ETHINYL ESTRADIOL

>D>

AB2 WATSON LABS

0.02MG;0.1MG

N77681 001 May 31, 2006 Aug CRLD

>A>

AB2 +

0.02MG;0.1MG

N77681 001 May 31, 2006 Aug CRLD

TRIPHASIL-28

@ WYETH PHARMS INC

0.03MG,0.04MG,0.03MG;0.05MG,0.075
MG,0.125MG

N19190 001 Nov 01, 1984 Apr DISC

ETHINYL ESTRADIOL; NORETHINDRONE

TABLET; ORAL-21

BALZIVA-21

>D>

>D> BARR

0.035MG;0.4MG

N76198 001 Apr 22, 2004 Aug DISC

>A>

@

0.035MG;0.4MG

N76198 001 Apr 22, 2004 Aug DISC

NORETHINDRONE AND ETHINYL ESTRADIOL (7/14)

WATSON LABS

0.035MG,0.035MG;0.5MG,1MG

N71041 001 Sep 24, 1991 Apr CRLD

TABLET; ORAL-28

NORETHINDRONE AND ETHINYL ESTRADIOL (10/11)

WATSON LABS

0.035MG,0.035MG;0.5MG,1MG

N71044 001 Apr 01, 1988 Apr CTEC

ETHINYL ESTRADIOL; NORETHINDRONE ACETATE

TABLET; ORAL

LOESTRIN 24 FE

>D>

+ WARNER CHILCOTT

0.02MG;1MG

N21871 001 Feb 17, 2006 Aug CFTG

>A>

AB

+

0.02MG;1MG

N21871 001 Feb 17, 2006 Aug CFTG

TABLET; ORAL-21

LOESTRIN 21 1.5/30

AB

WARNER CHILCOTT

0.03MG;1.5MG

N17875 001 Apr CRLD

LOESTRIN 21 1/20

AB

WARNER CHILCOTT

0.02MG;1MG

N17876 001 Apr CRLD

TABLET; ORAL-28

LOESTRIN FE 1/20

AB

WARNER CHILCOTT

0.02MG;1MG

N17354 001 Apr CRLD

>A>

NORETHINDRONE ACETATE AND ETHINYL ESTRADIOL AND FERROUS FUMARATE

>A>

AB

WATSON LABS

0.02MG;1MG

N78267 001 Sep 01, 2009 Aug NEWA

ETHINYL ESTRADIOL; NORGESTIMATE

TABLET; ORAL-28

ORTHO TRI-CYCLEN LO

AB	+	ORTHO MCNEIL JANSSEN	0.025MG,0.025MG,0.025MG;0.18MG,0.215MG,0.25MG	N21241 001	Aug 22, 2002	Jun	CFTG
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TRI LO SPRINTEC

AB		BARR	0.025MG,0.025MG,0.025MG;0.18MG,0.215MG,0.25MG	N76784 001	Jun 29, 2009	Jun	NEWA
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ETHINYL ESTRADIOL; NORGESTREL

TABLET; ORAL-21

OGESTREL 0.5/50-21

@ WATSON LABS

0.05MG;0.5MG

N75406 001 Dec 15, 1999 Apr DISC

OVRAL

@ AKRIMAX PHARMS

0.05MG;0.5MG

N16672 001 Mar CAHN

ETHOTOIN

TABLET; ORAL

PEGANONE

+ LUNDBECK INC

250MG

N10841 001 Apr CAHN

@

500MG

N10841 003 Apr CAHN

ETOPOSID

CAPSULE; ORAL

ETOPOSID

+ GENPHARM

50MG

N75635 001 Sep 19, 2001 Feb CRLD

VEPESID

@ BRISTOL MYERS SQUIBB

50MG

N19557 001 Dec 30, 1986 Feb DISC

INJECTABLE; INJECTION

ETOPOSID

>D>	AP	HOSPIRA	20MG/ML	N74320 001	Aug 30, 1995	Aug	DISC
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>A>		@	20MG/ML	N74320 001	Aug 30, 1995	Aug	DISC
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>D>	AP		20MG/ML	N74351 001	Aug 30, 1995	Aug	DISC
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>A>		@	20MG/ML	N74351 001	Aug 30, 1995	Aug	DISC
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EVEROLIMUS

TABLET; ORAL

AFINITOR

NOVARTIS

5MG

N22334 001 Mar 30, 2009 Mar NEWA

+

10MG

N22334 002 Mar 30, 2009 Mar NEWA

FAMOTIDINE

TABLET; ORAL

FAMOTIDINE

AB		ALEMBIC LTD	20MG	N78916 001	May 22, 2009	May	NEWA
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AB			40MG	N78916 002	May 22, 2009	May	NEWA
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		@ SANDOZ	20MG	N75302 001	Apr 16, 2001	Mar	DISC
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		@	40MG	N75302 002	Apr 16, 2001	Mar	DISC
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FEBUXOSTAT

TABLET; ORAL

ULORIC

TAKEDA PHARMS

40MG

N21856 001 Feb 13, 2009 Feb NEWA

+

80MG

N21856 002 Feb 13, 2009 Feb NEWA

FELODIPINE

TABLET, EXTENDED RELEASE; ORAL

FELODIPINE

>D>	AB	MYLAN	10MG	N78855 003	Apr 17, 2008	Aug	CRLD
>A>	AB	+	10MG	N78855 003	Apr 17, 2008	Aug	CRLD

FENOFIBRATE

CAPSULE; ORAL

LIPOFEN

>D>		CIPHER	50MG	N21612 001	Jan 11, 2006	Aug	CAHN
>D>			100MG	N21612 002	Jan 11, 2006	Aug	CAHN
>D>		+	150MG	N21612 003	Jan 11, 2006	Aug	CAHN
>A>		CIPHER PHARMS INC	50MG	N21612 001	Jan 11, 2006	Aug	CAHN
>A>			100MG	N21612 002	Jan 11, 2006	Aug	CAHN
>A>		+	150MG	N21612 003	Jan 11, 2006	Aug	CAHN

FENOFIBRIC ACID

TABLET; ORAL

FIBRICOR

>A>		AR HOLDING CO INC	35MG	N22418 001	Aug 14, 2009	Aug	NEWA
>A>		+	105MG	N22418 002	Aug 14, 2009	Aug	NEWA

FENOPROFEN CALCIUM

CAPSULE; ORAL

NALFON

+	PEDINOL	EQ 200MG BASE	N17604 003		Jul	CTNA
@		EQ 300MG BASE	N17604 002		Feb	DISC
		EQ 400MG BASE	N17604 004	Jul 21, 2009	Jul	NEWA

NALFON 200

+	PEDINOL	EQ 200MG BASE	N17604 003		Feb	CRLD
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TABLET; ORAL

FENOPROFEN CALCIUM

>D>	AB	ACTAVIS ELIZABETH	EQ 600MG BASE	N72274 001	May 02, 1988	Aug	DISC
>A>		@	EQ 600MG BASE	N72274 001	May 02, 1988	Aug	DISC
		@ SANDOZ	EQ 600MG BASE	N72396 001	Oct 17, 1988	Mar	DISC

FENTANYL CITRATE

INJECTABLE; INJECTION

SUBLIMAZE PRESERVATIVE FREE

AP	+	AKORN	EQ 0.05MG BASE/ML	N16619 001		May	CAHN
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FERUMOXYTOL

SOLUTION; INTRAVENOUS

FERAHEME

+	AMAG PHARMS INC	EQ 510MG IRON/17ML (EQ 30MG IRON/ML)	N22180 001	Jun 30, 2009	Jun	NEWA
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FLECAINIDE ACETATE

TABLET; ORAL

FLECAINIDE ACETATE

AB		APOTEX INC	50MG	N79164 001	Jul 09, 2009	Jun	NEWA
AB			100MG	N79164 002	Jul 09, 2009	Jun	NEWA
AB			150MG	N79164 003	Jul 09, 2009	Jun	NEWA
		@ SANDOZ	50MG	N76030 001	Oct 28, 2002	Mar	DISC
		@	100MG	N76030 002	Oct 28, 2002	Mar	DISC

TABLET; ORAL

FLECAINIDE ACETATE

@ SANDOZ

150MG

N76030 003 Oct 28, 2002 Mar DISC

FLUCONAZOLE

INJECTABLE; INJECTION

FLUCANAZOLE

AP ACS DOBFAR INFO SA 200MG/100ML (2MG/ML)

N79104 001 Jul 30, 2009 Jul NEWA

AP 400MG/200ML (2MG/ML)

N79104 002 Jul 30, 2009 Jul NEWA

FLUCONAZOLE IN SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER

>D> AP APOTEX INC 200MG/100ML (2MG/ML)

N76889 001 Mar 25, 2005 Aug DISC

>A> @ 200MG/100ML (2MG/ML)

N76889 001 Mar 25, 2005 Aug DISC

>D> AP 400MG/200ML (2MG/ML)

N76889 002 Mar 25, 2005 Aug DISC

>A> @ 400MG/200ML (2MG/ML)

N76889 002 Mar 25, 2005 Aug DISC

FLUDARABINE PHOSPHATE

INJECTABLE; INJECTION

FLUDARA

AP + GENZYME 50MG/VIAL

N20038 001 Apr 18, 1991 Jun CAHN

FLUDARABINE PHOSPHATE

AP ACTAVIS TOTOWA 50MG/VIAL

N78610 001 Feb 11, 2009 Feb NEWA

TABLET; ORAL

FLUDARABINE PHOSPHATE

+ SANOFI AVENTIS US 10MG

N22273 001 Dec 18, 2008 Jun CAHN

OFORTA

+ SANOFI AVENTIS US 10MG

N22273 001 Dec 18, 2008 Jul CTNA

FLUMAZENIL

INJECTABLE; INJECTION

FLUMAZENIL

AP HIKMA FARMACEUTICA 0.5MG/5ML (0.1MG/ML)

N78527 001 Mar 23, 2009 Mar NEWA

AP 1MG/10ML (0.1MG/ML)

N78527 002 Mar 23, 2009 Mar NEWA

FLUOCINOLONE ACETONIDE

CREAM; TOPICAL

FLUOCINOLONE ACETONIDE

>D> AT TARO 0.01%

N87102 001 Apr 27, 1982 Aug DISC

>A> @ TARO PHARMS USA INC 0.01%

N87102 001 Apr 27, 1982 Aug DISC

OIL; TOPICAL

DERMA-SMOOTH/FS

+ HILL DERMAC 0.01%

N19452 002 Nov 09, 2005 Feb NEWA

OIL/DROPS; OTIC

DERMOTIC

+ HILL DERMAC 0.01%

N19452 003 Nov 09, 2005 Jul CTNA

SHAMPOO; TOPICAL

CAPEX

+ GALDERMA LABS LP 0.01%

N20001 001 Aug 27, 1990 Jun CTNA

FLUOROMETHOLONE ACETATE; TOBRAMYCIN

SUSPENSION/DROPS; OPHTHALMIC

TOBRASONE

@ ALCON 0.1%;0.3%

N50628 001 Jul 21, 1989 Jun DISC

FLUOROURACIL

SOLUTION; TOPICAL

FLUOROPLEX

@ ELORAC

1%

N16765 001

Feb CAHN

FLUOXETINE HYDROCHLORIDE

CAPSULE; ORAL

FLUOXETINE HYDROCHLORIDE

AB1 ALEMBIC LTD EQ 10MG BASE

N90223 001 Mar 19, 2009 Mar NEWA

AB1 EQ 20MG BASE

N90223 002 Mar 19, 2009 Mar NEWA

AB EQ 40MG BASE

N90223 003 Mar 19, 2009 Mar NEWA

AB1 BEIJING DOUBLE CRANE EQ 10MG BASE

N76165 001 Feb 01, 2002 Mar CAHN

AB1 EQ 20MG BASE

N76165 002 Feb 01, 2002 Mar CAHN

AB1 LANDELA PHARM EQ 10MG BASE

N75464 001 Jan 30, 2002 Jul CAHN

AB1 EQ 20MG BASE

N75464 002 Jan 30, 2002 Jul CAHN

SOLUTION; ORAL

FLUOXETINE HYDROCHLORIDE

AA AUROBINDO PHARM EQ 20MG BASE/5ML

N79209 001 Mar 20, 2009 Mar NEWA

>D> AA HI TECH PHARMA EQ 20MG BASE/5ML

N75525 001 Jun 27, 2002 Aug DISC

>A> @ EQ 20MG BASE/5ML

N75525 001 Jun 27, 2002 Aug DISC

>D> AA PHARM ASSOC EQ 20MG BASE/5ML

N76015 001 Jan 30, 2002 Aug CRLD

>A> AA + EQ 20MG BASE/5ML

N76015 001 Jan 30, 2002 Aug CRLD

PROZAC

@ LILLY

EQ 20MG BASE/5ML

N20101 001 Apr 24, 1991 Jun DISC

TABLET; ORAL

FLUOXETINE HYDROCHLORIDE

AB MYLAN EQ 10MG BASE

N75755 001 Aug 02, 2001 Apr CAHN

+ EQ 20MG BASE

N75755 002 Aug 02, 2001 Apr CAHN

>D> AB SANDOZ EQ 10MG BASE

N76024 001 Jan 29, 2002 Aug DISC

>A> @ EQ 10MG BASE

N76024 001 Jan 29, 2002 Aug DISC

FLURAZEPAM HYDROCHLORIDE

CAPSULE; ORAL

FLURAZEPAM HYDROCHLORIDE

@ SANDOZ

15MG

N71716 001 Jul 31, 1991 Mar DISC

@

30MG

N71717 001 Jul 31, 1991 Mar DISC

FLURBIPROFEN

TABLET; ORAL

FLURBIPROFEN

@ SANDOZ

50MG

N74448 001 Jul 28, 1995 Mar DISC

@

100MG

N74448 002 Jul 28, 1995 Mar DISC

FLUTICASONE PROPIONATE

OINTMENT; TOPICAL

FLUTICASONE PROPIONATE

>D> AB TARO PHARM INDS 0.005%

N77145 001 Jun 14, 2005 Aug DISC

>A> @ 0.005%

N77145 001 Jun 14, 2005 Aug DISC

FLUVOXAMINE MALEATE

TABLET; ORAL

FLUVOXAMINE MALEATE

>D> AB GENPHARM 50MG

N75950 001 Oct 15, 2001 Aug DISC

>A> @ 50MG

N75950 001 Oct 15, 2001 Aug DISC

>D> AB 100MG

N75950 002 Oct 15, 2001 Aug DISC

TABLET; ORAL

FLUVOXAMINE MALEATE

>A>	@ GENPHARM	100MG	N75950 002	Oct 15, 2001	Aug	DISC
	@ IVAX PHARMS	25MG	N75898 001	Mar 12, 2001	Mar	DISC
	@	50MG	N75898 002	Mar 12, 2001	Mar	DISC
	@	100MG	N75898 003	Mar 12, 2001	Mar	DISC

FOLIC ACID

TABLET; ORAL

FOLIC ACID

AA	INVAGEN PHARMS	1MG	N90035 001	Jun 09, 2009	Jun	NEWA
>D>	AA	FOLICET				
	MISSION PHARMA	1MG	N87438 001		Aug	DISC
>A>	@	1MG	N87438 001		Aug	DISC

FOMEPIZOLE

INJECTABLE; INJECTION

FOMEPIZOLE

AP	GENERAMEDIX	1.5GM/1.5ML (1GM/ML)	N79033 001	Apr 07, 2009	Mar	NEWA
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FOSFOMYCIN TROMETHAMINE

FOR SUSPENSION; ORAL

MONUROL

+	ZAMBON SPA	EQ 3GM BASE/PACKET	N50717 001	Dec 19, 1996	Jun	CAHN
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FOSINOPRIL SODIUM

TABLET; ORAL

FOSINOPRIL SODIUM

	@ SANDOZ	10MG	N76188 001	Oct 08, 2004	Mar	DISC
	@	20MG	N76188 002	Oct 08, 2004	Mar	DISC
	@	40MG	N76188 003	Oct 08, 2004	Mar	DISC
AB	+ TEVA	40MG	N76139 003	Nov 25, 2003	Jun	CRLD

FOSINOPRIL SODIUM; HYDROCHLOROTHIAZIDE

TABLET; ORAL

FOSINOPRIL SODIUM AND HYDROCHLOROTHIAZIDE

AB	AUROBINDO PHARMA	10MG;12.5MG	N79245 001	Jul 09, 2009	Jun	NEWA
AB		20MG;12.5MG	N79245 002	Jul 09, 2009	Jun	NEWA
AB	INVAGEN PHARMS	10MG;12.5MG	N90228 001	Jul 09, 2009	Jun	NEWA
AB		20MG;12.5MG	N90228 002	Jul 09, 2009	Jun	NEWA
>D>	AB	TEVA	N76945 001	Jul 05, 2006	Aug	DISC
>A>	@	10MG;12.5MG	N76945 001	Jul 05, 2006	Aug	DISC
>D>	AB	+	N76945 002	Jul 05, 2006	Aug	DISC
>A>	@	20MG;12.5MG	N76945 002	Jul 05, 2006	Aug	DISC
AB	+	20MG;12.5MG	N76945 002	Jul 05, 2006	Jun	CRLD
	MONOPRIL-HCT					
	@ BRISTOL MYERS SQUIBB	10MG;12.5MG	N20286 002	Nov 30, 1994	Feb	DISC
	@	20MG;12.5MG	N20286 001	Nov 30, 1994	Feb	DISC

FUROSEMIDE

INJECTABLE; INJECTION

FUROSEMIDE

@ BAXTER HLTHCARE

10MG/ML

N71439 001	Sep 14, 1990	Jul	DISC
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TABLET; ORAL

FUROSEMIDE

AB	IPCA LABS LTD	20MG	N78010 001	Sep 18, 2006	Jun	CAHN
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TABLET; ORAL

FUROSEMIDE

AB	IPCA LABS LTD	40MG	N78010 002	Sep 18, 2006	Jun	CAHN
AB		80MG	N78010 003	Sep 18, 2006	Jun	CAHN

GABAPENTIN

CAPSULE; ORAL

GABAPENTIN

@ SANDOZ	100MG	N75428 001	Jan 24, 2006	Jul	DISC
@	300MG	N75428 002	Jan 24, 2006	Jul	DISC
@	400MG	N75428 003	Jan 24, 2006	Jul	DISC

TABLET; ORAL

GABAPENTIN

@ RANBAXY	600MG	N76605 001	Sep 14, 2005	Jul	DISC
@	800MG	N76605 002	Sep 14, 2005	Jul	DISC
@ SANDOZ	600MG	N76120 001	Jan 27, 2006	Jul	DISC
@	800MG	N76120 002	Jan 27, 2006	Jul	DISC

GALANTAMINE HYDROBROMIDE

CAPSULE, EXTENDED RELEASE; ORAL

GALANTAMINE HYDROBROMIDE

AB	IMPAX LABS	EQ 8MG BASE	N78484 001	May 27, 2009	May	NEWA
AB		EQ 16MG BASE	N78484 002	May 27, 2009	May	NEWA
AB		EQ 24MG BASE	N78484 003	May 27, 2009	May	NEWA

SOLUTION; ORAL

GALANTAMINE HYDROBROMIDE

AA	ROXANE	4MG/ML	N78185 001	Jan 30, 2009	Jan	NEWA
AA	+ ORTHO MCNEIL JANSSEN	4MG/ML	N21224 001	Jun 22, 2001	Jan	CFTG

TABLET; ORAL

GALANTAMINE HYDROBROMIDE

>A>	AB	ACTAVIS ELIZABETH	EQ 4MG BASE	N77585 001	Sep 15, 2009	Aug	NEWA
>A>	AB		EQ 8MG BASE	N77585 002	Sep 15, 2009	Aug	NEWA
>A>	AB		EQ 12MG BASE	N77585 003	Sep 15, 2009	Aug	NEWA
	AB	BEJING YABAO	EQ 4MG BASE	N77604 001	Feb 06, 2009	Apr	CAHN
	AB		EQ 8MG BASE	N77604 002	Feb 06, 2009	Apr	CAHN
	AB		EQ 12MG BASE	N77604 003	Feb 06, 2009	Apr	CAHN
	AB	MYLAN	EQ 4MG BASE	N77590 001	May 29, 2009	May	NEWA
	AB		EQ 4MG BASE	N77603 001	Aug 28, 2008	Apr	CAHN
	AB		EQ 8MG BASE	N77590 002	May 29, 2009	May	NEWA
	AB		EQ 8MG BASE	N77603 002	Aug 28, 2008	Apr	CAHN
	AB		EQ 12MG BASE	N77590 003	May 29, 2009	May	NEWA
	AB		EQ 12MG BASE	N77603 003	Aug 28, 2008	Apr	CAHN
	AB	PAR PHARM	EQ 4MG BASE	N77604 001	Feb 06, 2009	Jan	NEWA
	AB		EQ 8MG BASE	N77604 002	Feb 06, 2009	Jan	NEWA
	AB		EQ 12MG BASE	N77604 003	Feb 06, 2009	Jan	NEWA
	AB	ROXANE	EQ 4MG BASE	N77608 001	Feb 11, 2009	Jan	NEWA
	AB		EQ 8MG BASE	N77608 002	Feb 11, 2009	Jan	NEWA
	AB		EQ 12MG BASE	N77608 003	Feb 11, 2009	Jan	NEWA
	AB	SANDOZ	EQ 4MG BASE	N77589 001	Jun 22, 2009	Jun	NEWA
	AB		EQ 8MG BASE	N77589 002	Jun 22, 2009	Jun	NEWA
	AB		EQ 12MG BASE	N77589 003	Jun 22, 2009	Jun	NEWA
	AB	TEVA PHARMS	EQ 4MG BASE	N77587 001	Jul 09, 2009	Jun	NEWA
	AB		EQ 8MG BASE	N77587 002	Jul 09, 2009	Jun	NEWA
	AB		EQ 12MG BASE	N77587 003	Jul 09, 2009	Jun	NEWA

GANCICLOVIR SODIUM

INJECTABLE; INJECTION

CYTOVENE

@ ROCHE PALO

EQ 500MG BASE/VIAL

N19661 001 Jun 23, 1989 Jun DISC

GANCICLOVIR SODIUM

+ BEDFORD

EQ 500MG BASE/VIAL

N76222 001 Jul 16, 2003 Jun CRLD

GLIMEPIRIDE

TABLET; ORAL

GLIMEPIRIDE

@ RANBAXY

3MG

N77366 001 Oct 06, 2005 Jul DISC

@

6MG

N77366 002 Oct 06, 2005 Jul DISC

GLIMEPIRIDE; PIOGLITAZONE HYDROCHLORIDE

TABLET; ORAL

DUETACT

+ TAKEDA GLOBAL

2MG;30MG

N21925 001 Jul 28, 2006 May CRLD

4MG;30MG

N21925 002 Jul 28, 2006 May CRLD

GLIPIZIDE

TABLET; ORAL

GLIPIZIDE

>D> AB

SANDOZ

5MG

N74542 001 Jun 20, 1995 Aug DISC

>A>

@

5MG

N74542 001 Jun 20, 1995 Aug DISC

>D> AB

10MG

N74542 002 Jun 20, 1995 Aug DISC

>A>

@

10MG

N74542 002 Jun 20, 1995 Aug DISC

GLYBURIDE

TABLET; ORAL

GLYBURIDE

AB

+ TEVA

5MG

N74388 003 Aug 29, 1995 Mar CRLD

GLYBURIDE (MICRONIZED)

>D> AB

SANDOZ

1.5MG

N75174 001 Jun 22, 1998 Aug DISC

>A>

@

1.5MG

N75174 001 Jun 22, 1998 Aug DISC

>D> AB

3MG

N75174 002 Jun 22, 1998 Aug DISC

>A>

@

3MG

N75174 002 Jun 22, 1998 Aug DISC

MICRONASE

@ PHARMACIA AND UPJOHN

1.25MG

N17498 001 May 01, 1984 Mar DISC

@

2.5MG

N17498 002 May 01, 1984 Mar DISC

@

5MG

N17498 003 May 01, 1984 Mar DISC

GLYBURIDE; METFORMIN HYDROCHLORIDE

TABLET; ORAL

GLUCOVANCE

AB

+ BRISTOL MYERS SQUIBB

2.5MG;500MG

N21178 002 Jul 31, 2000 May CRLD

AB

5MG;500MG

N21178 003 Jul 31, 2000 May CRLD

GLYBURIDE AND METFORMIN HYDROCHLORIDE

AB

DR REDDYS LABS INC

1.25MG;250MG

N79009 001 Jun 03, 2009 May NEWA

AB

2.5MG;500MG

N79009 002 Jun 03, 2009 May NEWA

AB

5MG;500MG

N79009 003 Jun 03, 2009 May NEWA

GLYCOPYRROLATE

TABLET; ORAL

GLYCOPYRROLATE

>A> AA

RANBAXY

1MG

N40844 001 Aug 18, 2009 Aug NEWA

TABLET; ORAL

GLYCOPYRROLATE

>A>	AA	RANBAXY	2MG	N40844 002	Aug 18, 2009	Aug	NEWA
	AA	WEST WARD	1MG	N40836 001	Mar 05, 2009	Feb	NEWA
	AA		2MG	N40836 002	Mar 05, 2009	Feb	NEWA

GRANISETRON HYDROCHLORIDE

INJECTABLE; INJECTION

GRANISETRON HYDROCHLORIDE

>A>	AP	AKORN INC	EQ 0.1MG BASE/ML (EQ 0.1MG BASE/ML)	N79119 001	Sep 10, 2009	Aug	NEWA
>A>	AP		EQ 4MG BASE/4ML (EQ 1MG BASE/ML)	N79078 002	Sep 14, 2009	Aug	NEWA
>A>	AP		EQ 1MG BASE/ML (EQ 1MG BASE/ML)	N79078 001	Sep 14, 2009	Aug	NEWA
	AP	SANDOZ	EQ 0.1MG BASE/ML (EQ 0.1MG BASE/ML)	N78534 001	Apr 30, 2009	Apr	NEWA
	AP		EQ 1MG BASE/ML (EQ 1MG BASE/ML)	N78531 001	Apr 30, 2009	Apr	NEWA
	AP		EQ 4MG BASE/4ML (EQ 1MG BASE/ML)	N78531 002	Apr 30, 2009	Apr	NEWA

TABLET; ORAL

GRANISETRON HYDROCHLORIDE

	AB	DR REDDYS LABS LTD	EQ 1MG BASE	N78846 001	Feb 27, 2009	Feb	NEWA
	AB	NATCO PHARMA	EQ 1MG BASE	N78969 001	Jun 22, 2009	Jun	NEWA

GUANFACINE HYDROCHLORIDE

TABLET; ORAL

GUANFACINE HYDROCHLORIDE

>D>	AB	WATSON LABS	EQ 1MG BASE	N74762 001	Jun 25, 1997	Aug	DISC
>A>	@		EQ 1MG BASE	N74762 001	Jun 25, 1997	Aug	DISC
>D>	AB		EQ 2MG BASE	N74762 002	Jun 25, 1997	Aug	DISC
>A>	@		EQ 2MG BASE	N74762 002	Jun 25, 1997	Aug	DISC

HALOPERIDOL

TABLET; ORAL

HALOPERIDOL

>A>	AB	MYLAN	0.5MG	N70278 006	Jun 10, 1986	Aug	NEWA
>A>	AB		1MG	N70278 004	Jun 10, 1986	Aug	NEWA
>A>	AB		5MG	N70278 005	Jun 10, 1986	Aug	NEWA
	AB		10MG	N70278 002	Jul 16, 2009	Jun	NEWA
	AB		20MG	N70278 003	Jul 16, 2009	Jun	NEWA

HALOPERIDOL DECANOATE

INJECTABLE; INJECTION

HALOPERIDOL DECANOATE

	@	SANDOZ	EQ 50MG BASE/ML	N76463 001	Jun 24, 2005	Apr	DISC
	@		EQ 100MG BASE/ML	N76463 002	Jun 24, 2005	Apr	DISC

HALOPERIDOL LACTATE

INJECTABLE; INJECTION

HALOPERIDOL

	@	SANDOZ	EQ 5MG BASE/ML	N76464 001	Sep 29, 2004	Apr	DISC
>D>	AP	WATSON LABS	EQ 5MG BASE/ML	N70714 001	May 17, 1988	Aug	DISC
>A>	@		EQ 5MG BASE/ML	N70714 001	May 17, 1988	Aug	DISC

>D>		SOLUTION; ORAL					
>D>		HALOPERIDOL LACTATE					
>D>		ACTAVIS MID ATLANTIC	EQ 1MG BASE/ML	N74536 001	Nov 02, 1995	Aug	DISC
>A>	@		EQ 1MG BASE/ML	N74536 001	Nov 02, 1995	Aug	DISC

HEPARIN SODIUM

INJECTABLE; INJECTION

HEPARIN SODIUM

>A>	AP	HOSPIRA INC	1,000 UNITS/ML	N90571 001	Aug 31, 2009	Aug	NEWA
>A>	AP		5,000 UNITS/ML	N90571 002	Aug 31, 2009	Aug	NEWA
>A>	AP		10,000 UNITS/ML	N90571 003	Aug 31, 2009	Aug	NEWA

HISTRELIN ACETATE

IMPLANT; SUBCUTANEOUS

SUPPRELIN LA

+	ENDO PHARM	50MG	N22058 001	May 03, 2007	Apr	CAHN
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VANTAS

+	ENDO PHARM	50MG	N21732 001	Oct 12, 2004	Apr	CAHN
+	ENDO PHARMS	50MG	N21732 001	Oct 12, 2004	Mar	CAHN

HOMATROPINE METHYLBROMIDE; HYDROCODONE BITARTRATE

SYRUP; ORAL

HYCODAN

@	ENDO PHARMS	1.5MG/5ML;5MG/5ML	N05213 002	Jul 26, 1988	Feb	DISC
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HYDROCODONE BITARTRATE AND HOMATROPINE METHYLBROMIDE

AA	+	HI TECH PHARMA	1.5MG/5ML;5MG/5ML	N40613 001	Feb 08, 2008	Feb	CRLD
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TABLET; ORAL

HYCODAN

@	ENDO PHARMS	1.5MG;5MG	N05213 001	Jul 26, 1988	Feb	DISC
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TUSSIGON

AA	+	KING PHARMS	1.5MG;5MG	N88508 001	Jul 30, 1985	Feb	CRLD
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HYDRALAZINE HYDROCHLORIDE

INJECTABLE; INJECTION

HYDRALAZINE HYDROCHLORIDE

AP		AKORN	20MG/ML	N40730 001	Apr 21, 2009	Mar	NEWA
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TABLET; ORAL

HYDRALAZINE HYDROCHLORIDE

AA		GLENMARK PHARMS LTD	10MG	N90527 001	May 27, 2009	May	NEWA
AA			25MG	N90527 002	May 27, 2009	May	NEWA
AA			50MG	N90527 003	May 27, 2009	May	NEWA
AA			100MG	N90527 004	May 27, 2009	May	NEWA
		@ SANDOZ	10MG	N83241 001		Mar	DISC
		@	25MG	N83560 001		Mar	DISC
		@	50MG	N83561 001		Mar	DISC

HYDROCHLOROTHIAZIDE

CAPSULE; ORAL

HYDROCHLOROTHIAZIDE

AB		IPCA LABS LTD	12.5MG	N79237 001	Apr 02, 2009	Mar	NEWA
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SOLUTION; ORAL

HYDROCHLOROTHIAZIDE

@	ROXANE	50MG/5ML	N88587 001	Jul 02, 1984	Mar	DISC
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TABLET; ORAL

HYDROCHLOROTHIAZIDE

@	SANDOZ	25MG	N87565 001	Mar 09, 1982	Mar	DISC
@		50MG	N84912 001		Mar	DISC

HYDROCHLOROTHIAZIDE; PROPRANOLOL HYDROCHLORIDE

TABLET; ORAL

PROPRANOLOL HYDROCHLORIDE AND HYDROCHLOROTHIAZIDE

>D>	AB	ACTAVIS ELIZABETH	25MG;40MG	N70851 001	May 15, 1986	Aug	DISC
>A>		@	25MG;40MG	N70851 001	May 15, 1986	Aug	DISC
>D>	AB		25MG;80MG	N70852 001	May 15, 1986	Aug	DISC
>A>		@	25MG;80MG	N70852 001	May 15, 1986	Aug	DISC
		@ SANDOZ	25MG;40MG	N71060 001	Aug 26, 1987	Mar	DISC
		@	25MG;80MG	N71061 001	Aug 26, 1987	Mar	DISC

HYDROCHLOROTHIAZIDE; QUINAPRIL HYDROCHLORIDE

TABLET; ORAL

QUINAPRIL HYDROCHLORIDE AND HYDROCHLOROTHIAZIDE

AB		RANBAXY	12.5MG;EQ 10MG BASE	N78211 001	Mar 04, 2009	Feb	NEWA
AB			12.5MG;EQ 20MG BASE	N78211 002	Mar 04, 2009	Feb	NEWA
AB			25MG;EQ 20MG BASE	N78211 003	Mar 04, 2009	Feb	NEWA

HYDROCHLOROTHIAZIDE; SPIRONOLACTONE

TABLET; ORAL

SPIRONOLACTONE AND HYDROCHLOROTHIAZIDE

		@ SANDOZ	25MG;25MG	N86881 001		Mar	DISC
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HYDROCORTISONE

LOTION; TOPICAL

CETACORT

		@ CORIA	0.5%	N80426 002		Jun	DISC
		@	1%	N80426 001		Jun	DISC

HYDROCORTISONE ACETATE; OXYTETRACYCLINE HYDROCHLORIDE

SUSPENSION; OPHTHALMIC

TERRA-CORTRIL

		@ PFIZER	1.5%;EQ 5MG BASE/ML	N61016 001		Feb	DISC
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HYDROCORTISONE ACETATE; PRAMOXINE HYDROCHLORIDE

AEROSOL, METERED; TOPICAL

PROCTOFOAM HC

BX		UCB INC	1%;1%	N86195 001		Apr	CAHN
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HYDROCORTISONE ACETATE; UREA

CREAM; TOPICAL

CARMOL HC

AT	+	NYCOMED US	1%;10%	N80505 001		Apr	CAHN
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HYDROMORPHONE HYDROCHLORIDE

INJECTABLE; INJECTION

DILAUDID

+	PURDUE PHARM PRODS	1MG/ML	N19034 003	Apr 30, 2009	May	CMFD
		1MG/ML	N19034 003	Apr 30, 2009	Apr	NEWA
+		2MG/ML	N19034 004	Apr 30, 2009	May	CRLD
		2MG/ML	N19034 004	Apr 30, 2009	Apr	NEWA
+		4MG/ML	N19034 005	Apr 30, 2009	May	CRLD
		4MG/ML	N19034 005	Apr 30, 2009	Apr	NEWA

TABLET; ORAL

HYDROMORPHONE HYDROCHLORIDE

AB		ROXANE	4MG	N74597 003	May 29, 2009	May	NEWA
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HYDROXYUREACAPSULE; ORAL
HYDROXYUREA

AB	BARR	500MG	N75143 001	Oct 16, 1998	Feb	CMFD
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HYDROXYZINE HYDROCHLORIDEINJECTABLE; INJECTION
HYDROXYZINE HYDROCHLORIDE

AP	+	ABRAXIS PHARM	25MG/ML	N87329 001		Jun	CRLD
AP	+		50MG/ML	N87329 002		Jun	CRLD

		VISTARIL					
		@ PFIZER	25MG/ML	N11111 001		Jun	DISC
		@	50MG/ML	N11111 002		Jun	DISC

TABLET; ORAL

HYDROXYZINE HYDROCHLORIDE

		@ SANDOZ	10MG	N87869 001	Dec 20, 1982	Mar	DISC
		@	25MG	N87870 001	Dec 20, 1982	Mar	DISC
		@	50MG	N87871 001	Dec 20, 1982	Mar	DISC

HYDROXYZINE PAMOATECAPSULE; ORAL
HYDROXYZINE PAMOATE
@ SANDOZ

EQ 25MG HCL	N81127 001	Jun 28, 1991	Mar	DISC
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IBANDRONATE SODIUM

TABLET; ORAL

BONIVA

@ ROCHE

EQ 2.5MG BASE	N21455 001	May 16, 2003	Jul	DISC
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IBUPROFEN

SOLUTION; INTRAVENOUS

CALDOLOR

CUMBERLAND PHARMS

400MG/4ML (100MG/ML)	N22348 001	Jun 11, 2009	Jun	NEWA
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+	800MG/8ML (100MG/ML)	N22348 002	Jun 11, 2009	Jun	NEWA
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TABLET; ORAL

IBUPROFEN

AB	+	DR REDDYS LA	800MG	N75682 003	Nov 14, 2001	Feb	CRLD
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	@ SANDOZ	300MG	N70734 001	Jun 12, 1986	Mar	DISC
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	@	400MG	N70735 001	Jun 12, 1986	Mar	DISC
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	@	600MG	N70736 001	Jun 12, 1986	Mar	DISC
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	@	800MG	N72169 001	Dec 11, 1987	Mar	DISC
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AB		SHASUN USA	400MG	N78329 001	Feb 05, 2009	Jan	NEWA
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AB			600MG	N78329 002	Feb 05, 2009	Jan	NEWA
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AB			800MG	N78329 003	Feb 05, 2009	Jan	NEWA
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>D>	AB	WATSON LABS	800MG	N71547 001	Jul 02, 1987	Aug	DISC
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>A>		@	800MG	N71547 001	Jul 02, 1987	Aug	DISC
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MOTRIN

@ MCNEIL CONSUMER

300MG	N17463 003		Apr	DISC
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@	N17463 002		Feb	DISC
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@	N17463 004		Feb	DISC
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@	N17463 005	May 22, 1985	Feb	DISC
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IBUPROFEN LYSINE

INJECTABLE; INTRAVENOUS
NEOPROFEN

+	LUNDBECK INC	EQ 20MG BASE/2ML (EQ 10MG BASE/ML)	N21903 001	Apr 13, 2006	Apr	CAHN
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IDARUBICIN HYDROCHLORIDE

INJECTABLE; INJECTION
IDARUBICIN HYDROCHLORIDE

AP	APP PHARMS	1MG/ML	N65440 001	Aug 04, 2009	Jul	NEWA
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IFOSFAMIDE

INJECTABLE; INJECTION
IFEX

@	BAXTER HLTHCARE	1GM/VIAL	N19763 001	Dec 30, 1988	Feb	CAHN
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@		3GM/VIAL	N19763 002	Dec 30, 1988	Feb	CAHN
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IFOSFAMIDE; MESNA

INJECTABLE; INJECTION
IFEX/MESNEX KIT

+	BAXTER HLTHCARE	1GM/VIAL;100MG/ML	N19763 003	Oct 10, 1992	Feb	CAHN
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+		3GM/VIAL;100MG/ML	N19763 004	Oct 10, 1992	Feb	CAHN
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INJECTABLE; INTRAVENOUS
IFOSFAMIDE/MESNA KIT

>D>	+	TEVA PARENTERAL	EQ 1GM /20ML(50MG/ML);EQ 1GM /10ML(100MG/ML)	N75874 001	Feb 26, 2002	Aug	CPOT
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>A>	+		EQ 1GM/20ML;EQ 1GM/10ML (50MG/ML; 100MG/ML)	N75874 001	Feb 26, 2002	Aug	CPOT
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ILOPERIDONE

TABLET; ORAL
FANAPT

	VANDA PHARMS INC	1MG	N22192 001	May 06, 2009	May	NEWA
		2MG	N22192 002	May 06, 2009	May	NEWA
		4MG	N22192 003	May 06, 2009	May	NEWA
		6MG	N22192 004	May 06, 2009	May	NEWA
		8MG	N22192 005	May 06, 2009	May	NEWA
		10MG	N22192 006	May 06, 2009	May	NEWA
+		12MG	N22192 007	May 06, 2009	May	NEWA

INAMRINONE LACTATE

INJECTABLE; INJECTION
AMRINONE LACTATE

>D>	AP	BEDFORD	EQ 5MG BASE/ML	N75513 001	May 09, 2000	Aug	CRLD
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>A>	AP	+	EQ 5MG BASE/ML	N75513 001	May 09, 2000	Aug	CRLD
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>D>	AP	+	HOSPIRA	EQ 5MG BASE/ML	N74616 001	Aug 03, 1998	Aug	DISC
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>A>		@	EQ 5MG BASE/ML	N74616 001	Aug 03, 1998	Aug	DISC
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INDINAVIR SULFATE

CAPSULE; ORAL
CRIXIVAN

@	MERCK	EQ 333MG BASE	N20685 005	Dec 17, 1998	Jun	DISC
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INDOMETHACIN

		CAPSULE, EXTENDED RELEASE; ORAL			
		INDOCIN SR			
AB	+	SANDOZ	75MG	N74464 001	May 28, 1998 Feb CTEC
		INDOMETHACIN			
AB		AVANTHI INC	75MG	N79175 001	Mar 06, 2009 Feb NEWA
		SUSPENSION; ORAL			
		INDOCIN			
	+	IROKO PHARMS	25MG/5ML	N18332 001	Oct 10, 1985 Mar CAHN

INDOMETHACIN SODIUM

		INJECTABLE; INJECTION			
		INDOCIN			
AP	+	LUNDBECK INC	EQ 1MG BASE/VIAL	N18878 001	Jan 30, 1985 Apr CAHN

INSULIN GLULISINE RECOMBINANT

		INJECTABLE; SUBCUTANEOUS			
		APIDRA SOLOSTAR			
		SANOFI AVENTIS US	300 UNITS/3ML	N21629 003	Feb 24, 2009 Feb NEWA

INSULIN RECOMBINANT HUMAN

		POWDER; INHALATION			
		EXUBERA			
	@	PFIZER	1MG/INH	N21868 001	Jan 27, 2006 Jun DISC
	@		3MG/INH	N21868 002	Jan 27, 2006 Jun DISC

IPRATROPIUM BROMIDE

		SOLUTION; INHALATION			
		IPRATROPIUM BROMIDE			
>D>	AN	PHARMASCIENCE INC	0.02%	N75507 001	Jan 19, 2001 Aug DISC
>A>		@	0.02%	N75507 001	Jan 19, 2001 Aug DISC
	AN	TEVA PARENTERAL	0.02%	N75313 001	Feb 07, 2000 Apr CAHN

IRINOTECAN HYDROCHLORIDE

		INJECTABLE; INJECTION			
		IRINOTECAN HYDROCHLORIDE			
AP		PHARMAFORCE	40MG/2ML (20MG/ML)	N90016 001	Jan 28, 2009 Mar NEWA
AP			40MG/2ML (20MG/ML)	N90016 001	Jan 28, 2009 Jan NEWA
AP			100MG/5ML (20MG/ML)	N90016 002	Jan 28, 2009 Mar NEWA
AP			100MG/5ML (20MG/ML)	N90016 002	Jan 28, 2009 Jan NEWA

ISOPROTERENOL HYDROCHLORIDE

		INJECTABLE; INJECTION			
		ISOPROTERENOL HYDROCHLORIDE			
>D>	AP	HOSPIRA	0.2MG/ML	N83346 001	Aug DISC
>A>		@	0.2MG/ML	N83346 001	Aug DISC

ISOSORBIDE DINITRATE

		TABLET; SUBLINGUAL			
		ISOSORBIDE DINITRATE			
	@	SANDOZ	2.5MG	N86225 001	Feb 19, 1988 Mar DISC
	@		5MG	N86222 001	Feb 19, 1988 Mar DISC

ISOTRETINOIN

CAPSULE; ORAL

ACCUTANE

@ HOFFMANN LA ROCHE

10MG

N18662 002 May 07, 1982 Jul DISC

@

20MG

N18662 004 Mar 28, 1983 Jul DISC

@

40MG

N18662 003 May 07, 1982 Jul DISC

KETOCONAZOLE

GEL; TOPICAL

XOLEGEL

+ STIEFEL LABS INC

2%

N21946 001 Jul 28, 2006 Jan CAHN

KETOPROFEN

CAPSULE; ORAL

KETOPROFEN

@ SANDOZ

50MG

N74024 001 Dec 29, 1995 Mar DISC

@

75MG

N74024 002 Dec 29, 1995 Mar DISC

KETOROLAC TROMETHAMINE

SOLUTION/DROPS; OPHTHALMIC

ACUVAIL

+ ALLERGAN

0.45%

N22427 001 Jul 22, 2009 Jul NEWA

LAMOTRIGINE

TABLET; ORAL

LAMOTRIGINE

AB APOTEX INC

25MG

N78625 001 Jan 27, 2009 Jan NEWA

AB

100MG

N78625 002 Jan 27, 2009 Jan NEWA

AB

150MG

N78625 003 Jan 27, 2009 Jan NEWA

AB

200MG

N78625 004 Jan 27, 2009 Jan NEWA

AB

AUROBINDO PHARMA

25MG

N78956 001 Jan 27, 2009 Jan NEWA

AB

100MG

N78956 002 Jan 27, 2009 Jan NEWA

AB

150MG

N78956 003 Jan 27, 2009 Jan NEWA

AB

200MG

N78956 004 Jan 27, 2009 Jan NEWA

AB

CADISTA PHARMS

25MG

N79132 001 Jan 27, 2009 Jan NEWA

AB

100MG

N79132 002 Jan 27, 2009 Jan NEWA

AB

150MG

N79132 003 Jan 27, 2009 Jan NEWA

AB

200MG

N79132 004 Jan 27, 2009 Jan NEWA

AB

DR REDDYS LABS LTD

25MG

N76708 001 Jan 27, 2009 Jan NEWA

AB

100MG

N76708 002 Jan 27, 2009 Jan NEWA

AB

150MG

N76708 003 Jan 27, 2009 Jan NEWA

AB

200MG

N76708 004 Jan 27, 2009 Jan NEWA

AB

GENPHARM ULC

25MG

N77428 001 Jan 27, 2009 Jan NEWA

AB

100MG

N77428 002 Jan 27, 2009 Jan NEWA

AB

150MG

N77428 003 Jan 27, 2009 Jan NEWA

AB

200MG

N77428 004 Jan 27, 2009 Jan NEWA

AB

MATRIX LABS LTD

25MG

N78443 001 Feb 11, 2009 Jan NEWA

AB

100MG

N78443 002 Feb 11, 2009 Jan NEWA

AB

150MG

N78443 003 Feb 11, 2009 Jan NEWA

AB

200MG

N78443 004 Feb 11, 2009 Jan NEWA

AB

MYLAN

25MG

N77420 001 Jan 27, 2009 Jan NEWA

AB

100MG

N77420 002 Jan 27, 2009 Jan NEWA

AB

150MG

N77420 003 Jan 27, 2009 Jan NEWA

AB

200MG

N77420 004 Jan 27, 2009 Jan NEWA

AB

ROXANE

25MG

N77392 001 Jan 27, 2009 Jan NEWA

TABLET; ORAL

LAMOTRIGINE

AB	ROXANE	100MG	N77392 002	Jan 27, 2009	Jan	NEWA
AB		150MG	N77392 003	Jan 27, 2009	Jan	NEWA
AB		200MG	N77392 004	Jan 27, 2009	Jan	NEWA
AB	SANDOZ	25MG	N78645 001	Jan 27, 2009	Jan	NEWA
AB		100MG	N78645 002	Jan 27, 2009	Jan	NEWA
AB		150MG	N78645 003	Jan 27, 2009	Jan	NEWA
AB		200MG	N78645 004	Jan 27, 2009	Jan	NEWA
AB	TARO PHARM INDS	25MG	N78525 001	Jan 27, 2009	Jan	NEWA
AB		100MG	N78525 002	Jan 27, 2009	Jan	NEWA
AB		150MG	N78525 003	Jan 27, 2009	Jan	NEWA
AB		200MG	N78525 004	Jan 27, 2009	Jan	NEWA
AB	TORRENT PHARMS	25MG	N78947 001	Jan 27, 2009	Jan	NEWA
AB		100MG	N78947 002	Jan 27, 2009	Jan	NEWA
AB		150MG	N78947 003	Jan 27, 2009	Jan	NEWA
AB		200MG	N78947 004	Jan 27, 2009	Jan	NEWA
AB	UPSHER SMITH	25MG	N78310 001	Feb 04, 2009	Jan	NEWA
AB		100MG	N78310 002	Feb 04, 2009	Jan	NEWA
AB		150MG	N78310 003	Feb 04, 2009	Jan	NEWA
AB		200MG	N78310 004	Feb 04, 2009	Jan	NEWA
AB	WOCKHARDT	25MG	N78982 001	Jan 27, 2009	Jan	NEWA
AB		100MG	N78982 002	Jan 27, 2009	Jan	NEWA
AB		150MG	N78982 003	Jan 27, 2009	Jan	NEWA
AB		200MG	N78982 004	Jan 27, 2009	Jan	NEWA
AB	ZYDUS PHARMS USA	25MG	N77633 001	Jan 27, 2009	Jan	NEWA
		50MG	N77633 002	Jan 27, 2009	Jan	NEWA
AB		100MG	N77633 003	Jan 27, 2009	Jan	NEWA
AB		150MG	N77633 004	Jan 27, 2009	Jan	NEWA
AB		200MG	N77633 005	Jan 27, 2009	Jan	NEWA
		250MG	N77633 006	Jan 27, 2009	Jan	NEWA

TABLET, CHEWABLE; ORAL

LAMOTRIGINE

AB	GLENMARK GENERICS	5MG	N79099 001	Feb 19, 2009	Feb	NEWA
AB		25MG	N79099 002	Feb 19, 2009	Feb	NEWA
AB	TARO	5MG	N79204 001	Feb 04, 2009	Jan	NEWA
AB		25MG	N79204 002	Feb 04, 2009	Jan	NEWA

TABLET, EXTENDED RELEASE; ORAL

LAMICTAL XR

	SMITHKLINE BEECHAM	25MG	N22115 001	May 29, 2009	May	NEWA
+		50MG	N22115 002	May 29, 2009	Jul	CRLD
		50MG	N22115 002	May 29, 2009	May	NEWA
		100MG	N22115 003	May 29, 2009	May	NEWA
		200MG	N22115 004	May 29, 2009	Jul	CRLD
+		200MG	N22115 004	May 29, 2009	May	NEWA

TABLET, ORALLY DISINTEGRATING; ORAL

LAMICTAL ODT

	SMITHKLINE BEECHAM	25MG	N22251 001	May 08, 2009	May	NEWA
+		50MG	N22251 002	May 08, 2009	Jul	CRLD
		50MG	N22251 002	May 08, 2009	May	NEWA
		100MG	N22251 003	May 08, 2009	May	NEWA
		200MG	N22251 004	May 08, 2009	Jul	CRLD
+		200MG	N22251 004	May 08, 2009	May	NEWA

LANSOPRAZOLE

FOR SUSPENSION, DELAYED RELEASE; ORAL

PREVACID

@ TAKEDA PHARMS NA 15MG/PACKET

N21281 001 May 03, 2001 May DISC

@ 30MG/PACKET

N21281 002 May 03, 2001 May DISC

>D> INJECTABLE; INTRAVENOUS

>D> PREVACID IV

>D> + TAKEDA PHARMS NA 30MG/VIAL

N21566 001 May 27, 2004 Aug DISC

>A> @ 30MG/VIAL

N21566 001 May 27, 2004 Aug DISC

>D> LANSOPRAZOLE; NAPROXEN

>D> CAPSULE, DELAYED REL PELLETS; TABLET; ORAL

>D> PREVACID NAPRAPAC 500 (COPACKAGED)

>A> @ TAKEDA PHARMS NA 15MG,N/A;N/A,500MG

N21507 004 Nov 14, 2003 Aug DISC

>D> CAPSULE, DELAYED REL PELLETS, TABLET; ORAL

>D> PREVACID NAPRAPAC 500 (COPACKAGED)

>D> + TAKEDA PHARMS NA 15MG,N/A;N/A,500MG

N21507 004 Nov 14, 2003 Aug DISC

LEUCOVORIN CALCIUM

INJECTABLE; INJECTION

LEUCOVORIN CALCIUM PRESERVATIVE FREE

>D> AP + BEDFORD EQ 10MG BASE/ML

N40347 001 Apr 25, 2000 Aug CTEC

>A> + EQ 10MG BASE/ML

N40347 001 Apr 25, 2000 Aug CTEC

>D> AP TEVA PARENTERAL EQ 10MG BASE/ML

N40332 001 Jun 28, 1999 Aug DISC

>A> @ EQ 10MG BASE/ML

N40332 001 Jun 28, 1999 Aug DISC

TABLET; ORAL

LEUCOVORIN CALCIUM

@ COREPHARMA EQ 5MG BASE

N74544 001 Aug 28, 1997 Jun CAHN

@ EQ 25MG BASE

N74544 002 Aug 28, 1997 Jun CAHN

ROXANE EQ 10MG BASE

N72734 001 Feb 22, 1993 Jun CTEC

@ XANODYNE PHARM EQ 5MG BASE

N18459 001 Jan 30, 1986 May DISC

@ EQ 10MG BASE

N71962 001 Nov 19, 1987 Jun DISC

LEUPROLIDE ACETATE

IMPLANT; IMPLANTATION

VIADUR

@ ORTHO MCNEIL JANSSEN EQ 65MG BASE

N21088 001 Mar 03, 2000 Jun DISC

INJECTABLE; INJECTION

LEUPROLIDE ACETATE

AP SUN PHARMA GLOBAL 1MG/0.2ML

N78885 001 Mar 09, 2009 Feb NEWA

LEVALBUTEROL HYDROCHLORIDE

SOLUTION; INHALATION

LEVALBUTEROL HYDROCHLORIDE

AN DEY EQ 0.25% BASE

N78309 001 Mar 20, 2009 Mar NEWA

XOPENEX

AN + SEPRACOR EQ 0.25% BASE

N20837 004 Jul 18, 2003 Mar CFTG

LEVETIRACETAM

SOLUTION; ORAL

LEVETIRACETAM

AA SILARX 100MG/ML

N90263 001 Apr 03, 2009 Mar NEWA

AA TARO 100MG/ML

N78774 001 Feb 10, 2009 Jan NEWA

TABLET; ORAL

LEVETIRACETAM

AB	APOTEX INC	250MG	N78869 001	Mar 13, 2009	Mar	NEWA
AB		500MG	N78869 002	Mar 13, 2009	Mar	NEWA
AB		750MG	N78869 003	Mar 13, 2009	Mar	NEWA
AB		1GM	N78869 004	Mar 13, 2009	Mar	NEWA
AB	CIPLA LTD	250MG	N77319 001	Mar 20, 2009	Mar	NEWA
AB		500MG	N77319 002	Mar 20, 2009	Mar	NEWA
AB		750MG	N77319 003	Mar 20, 2009	Mar	NEWA
AB	GENPHARM ULC	250MG	N78731 001	Feb 10, 2009	Jan	NEWA
AB		500MG	N78731 002	Feb 10, 2009	Jan	NEWA
AB		750MG	N78731 003	Feb 10, 2009	Jan	NEWA
AB		1GM	N78731 004	Feb 10, 2009	Jan	NEWA
AB	SOLCO HLTHCARE	250MG	N78106 001	Feb 10, 2009	Jan	NEWA
AB		500MG	N78106 002	Feb 10, 2009	Jan	NEWA
AB		750MG	N78106 003	Feb 10, 2009	Jan	NEWA
AB		1GM	N78106 004	Feb 10, 2009	Jan	NEWA
AB	WATSON LABS FLORIDA	250MG	N77408 001	Mar 02, 2009	Feb	NEWA
AB		500MG	N77408 002	Mar 02, 2009	Feb	NEWA
AB		750MG	N77408 003	Mar 02, 2009	Feb	NEWA
AB	ZYDUS PHARMS USA INC	250MG	N78918 001	Apr 29, 2009	Apr	NEWA
AB		1GM	N78918 002	Apr 29, 2009	Apr	NEWA

TABLET, EXTENDED RELEASE; ORAL

KEPPRA XR

	UCB INC	500MG	N22285 001	Sep 12, 2008	Feb	CRLD
+		750MG	N22285 002	Feb 12, 2009	Feb	NEWA

LEVOFLOXACIN

TABLET; ORAL

LEVAQUIN

	ORTHO MCNEIL JANSSEN	250MG	N20634 001	Dec 20, 1996	Jun	CTEC
AB		250MG	N20634 001	Dec 20, 1996	Apr	CFTG
		500MG	N20634 002	Dec 20, 1996	Jun	CTEC
AB		500MG	N20634 002	Dec 20, 1996	Apr	CFTG
+		750MG	N20634 003	Sep 08, 2000	Jun	CTEC
AB	+	750MG	N20634 003	Sep 08, 2000	Apr	CFTG

LEVONORDEFIN; MEPIVACAINE HYDROCHLORIDE

INJECTABLE; INJECTION

MEPIVACAINE HYDROCHLORIDE W/ LEVONORDEFIN

>D>	AP	GRAHAM CHEM	0.05MG/ML; 2%	N84850 002	Oct 21, 1983	Aug	DISC
>A>		@	0.05MG/ML; 2%	N84850 002	Oct 21, 1983	Aug	DISC
>D>		POLOCAINE W/ LEVONORDEFIN					
>D>	AP	+ DENTSPLY PHARM	0.05MG/ML; 2%	N89517 001	Apr 14, 1988	Aug	DISC
>A>		@	0.05MG/ML; 2%	N89517 001	Apr 14, 1988	Aug	DISC

LEVONORGESTREL

IMPLANT; IMPLANTATION

NORPLANT II

	@ POPULATION COUNCIL	75MG/IMPLANT	N20544 001	Nov 01, 1996	Jul	DISC
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TABLET; ORAL

LEVONORGESTREL

>A>	AB	WATSON LABS	0.75MG	N78665 001	Aug 28, 2009	Aug	NEWA
	AB		0.75MG	N78666 001	Jun 24, 2009	Jun	NEWA
		PLAN B ONE-STEP					
	+	DURAMED	1.5MG	N21998 001	Jul 10, 2009	Jul	NEWA

LEVOTHYROXINE SODIUM**

**Refer to Annual Edition Preface Section 1.8 Levothyroxine Sodium for amplifying information

TABLET; ORAL

LEVOTHYROXINE SODIUM

AB2, AB3	MERCK KGAA	0.025MG	N76752 001	Jun 16, 2005	Apr	CAHN
AB2, AB3		0.05MG	N76752 002	Jun 16, 2005	Apr	CAHN
AB2, AB3		0.075MG	N76752 003	Jun 16, 2005	Apr	CAHN
AB2, AB3		0.088MG	N76752 004	Jun 16, 2005	Apr	CAHN
AB2, AB3		0.1MG	N76752 005	Jun 16, 2005	Apr	CAHN
AB2, AB3		0.112MG	N76752 006	Jun 16, 2005	Apr	CAHN
AB2, AB3		0.125MG	N76752 007	Jun 16, 2005	Apr	CAHN
AB2, AB3		0.15MG	N76752 008	Jun 16, 2005	Apr	CAHN
AB2, AB3		0.175MG	N76752 009	Jun 16, 2005	Apr	CAHN
AB2, AB3		0.2MG	N76752 010	Jun 16, 2005	Apr	CAHN
AB2, AB3		0.3MG	N76752 011	Jun 16, 2005	Apr	CAHN

LIDOCAINE HYDROCHLORIDE

INJECTABLE; INJECTION

LIDOCAINE HYDROCHLORIDE IN PLASTIC CONTAINER

	@ HOSPIRA	10%	N88367 001	Jul 31, 1984	Apr	DISC
>D>	AP	20%	N88368 001	Jul 31, 1984	Aug	DISC
>A>	@	20%	N88368 001	Jul 31, 1984	Aug	DISC
	XYLOCAINE					
AP	+ APP PHARMS	2%	N06488 002		Apr	CMFD
	XYLOCAINE DENTAL					
AP	+ DENTSPLY PHARM	2%	N21380 001		Apr	CTNA
	SYSTEM; INTRADERMAL					
	ZINGO					
	@ ANESIVA	0.5MG	N22114 001	Aug 16, 2007	May	DISC

LINDANE

LOTION; TOPICAL

LINDANE

@ OLTA PHARMS

1%

N87313 001

Apr CAHN

SHAMPOO; TOPICAL

LINDANE

AT + OLTA PHARMS

1%

N87266 001

Jan CAHN

LIOTHYRONINE SODIUM

TABLET; ORAL

CYTOMEL

AB KING PHARMS

EQ 0.005MG BASE

N10379 001

Mar CFTG

AB

EQ 0.025MG BASE

N10379 002

Mar CTEC

AB +

EQ 0.05MG BASE

N10379 003

Mar CTEC

LIOTHYRONINE SODIUM

AB COASTAL PHARMS

EQ 0.005MG BASE

N90097 001 Mar 20, 2009 Mar NEWA

AB

EQ 0.025MG BASE

N90097 002 Mar 20, 2009 Mar NEWA

AB

EQ 0.05MG BASE

N90097 003 Mar 20, 2009 Mar NEWA

TABLET; ORAL

LIOTHYRONINE SODIUM

AB	MYLAN	EQ 0.005MG BASE	N90326 001	Jul 14, 2009	Jun	NEWA
AB		EQ 0.025MG BASE	N90326 002	Jul 14, 2009	Jun	NEWA
AB		EQ 0.05MG BASE	N90326 003	Jul 14, 2009	Jun	NEWA

LITHIUM CARBONATE

CAPSULE; ORAL

LITHIUM CARBONATE

AB	GLENMARK GENERICS	150MG	N79139 001	Feb 03, 2009	Jan	NEWA
AB		300MG	N79139 002	Feb 03, 2009	Jan	NEWA
AB		600MG	N79139 003	Feb 03, 2009	Jan	NEWA

LOPERAMIDE HYDROCHLORIDE

CAPSULE; ORAL

LOPERAMIDE HYDROCHLORIDE

@ SANDOZ 2MG

N72993 001 Aug 28, 1992 Mar DISC

LORAZEPAM

CONCENTRATE; ORAL

LORAZEPAM

AA	PADDOCK LABS	2MG/ML	N79244 001	Apr 28, 2009	Apr	NEWA
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LORAZEPAM INTENSOL

AA	+ ROXANE	2MG/ML	N72755 001	Jun 28, 1991	Apr	CFTG
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SOLUTION; ORAL

LORAZEPAM

@ ROXANE 0.5MG/5ML

N74648 001 Mar 18, 1997 Mar DISC

TABLET; ORAL

LORAZEPAM

>D>	AB	WATSON LABS	0.5MG	N71117 001	Jul 24, 1986	Aug	DISC
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>A>	@		0.5MG	N71117 001	Jul 24, 1986	Aug	DISC
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>D>	AB		2MG	N71110 001	Jul 24, 1986	Aug	DISC
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>A>	@		2MG	N71110 001	Jul 24, 1986	Aug	DISC
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MALATHION

LOTION; TOPICAL

MALATHION

AT	SYNERX PHARMA	0.5%	N78743 001	Mar 06, 2009	Feb	NEWA
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OVIDE

AT	+ TARO PHARMS NORTH	0.5%	N18613 001	Aug 02, 1982	Feb	CFTG
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>D> MECAMYLAMINE HYDROCHLORIDE

>D> TABLET; ORAL

>D> INVERSINE

>D>	+ TARGACEPT	2.5MG	N10251 001		Aug	DISC
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>A>	@	2.5MG	N10251 001		Aug	DISC
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MECHLORETHAMINE HYDROCHLORIDE

INJECTABLE; INJECTION

MUSTARGEN

	+ LUNDBECK INC	10MG/VIAL	N06695 001		Apr	CAHN
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MECLIZINE HYDROCHLORIDE

TABLET; ORAL

ANTIVERT

AA	+ PFIZER	12.5MG	N10721 006		Jan	CAHN
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TABLET; ORAL									
ANTIVERT									
AA	+	PFIZER	25MG	N10721	004		Jan	CAHN	
AA	+		50MG	N10721	001	Jan 20, 1982	Jan	CAHN	
<u>MECLOFENAMATE SODIUM</u>									
CAPSULE; ORAL									
MECLOFENAMATE SODIUM									
	@	SANDOZ	EQ 50MG BASE	N72262	001	Nov 29, 1988	Mar	DISC	
	@		EQ 100MG BASE	N72263	001	Nov 29, 1988	Mar	DISC	
<u>MEDROXYPROGESTERONE ACETATE</u>									
INJECTABLE; INJECTION									
MEDROXYPROGESTERONE ACETATE									
AB		SANDOZ	150MG/ML	N78711	001	May 20, 2009	May	NEWA	
<u>MELOXICAM</u>									
TABLET; ORAL									
MELOXICAM									
AB		BEIJING YABAO	7.5MG	N77933	001	Jul 19, 2006	Apr	CAHN	
AB			15MG	N77933	002	Jul 19, 2006	Apr	CAHN	
AB		STRIDES ARCOLAB LTD	7.5MG	N77928	001	May 13, 2009	May	NEWA	
AB			15MG	N77928	002	May 13, 2009	May	NEWA	
<u>MELPHALAN HYDROCHLORIDE</u>									
INJECTABLE; INJECTION									
ALKERAN									
AP	+	GLAXOSMITHKLINE	EQ 50MG BASE/VIAL	N20207	001	Nov 18, 1992	Jun	CFTG	
MELPHALAN HYDROCHLORIDE									
AP		SYNERX	EQ 50MG BASE/VIAL	N90270	001	Jun 09, 2009	Jun	NEWA	
<u>MEPERIDINE HYDROCHLORIDE</u>									
TABLET; ORAL									
MEPERIDINE HYDROCHLORIDE									
AA		MIKART	50MG	N40893	001	Jun 24, 2009	Jun	NEWA	
			75MG	N40893	002	Jun 24, 2009	Jun	NEWA	
AA			100MG	N40893	003	Jun 24, 2009	Jun	NEWA	
			150MG	N40893	004	Jun 24, 2009	Jun	NEWA	
<u>MEPIVACAINE HYDROCHLORIDE</u>									
INJECTABLE; INJECTION									
POLOCAINE									
AP		APP PHARMS	1%	N89407	001	Dec 01, 1986	Jun	CAHN	
AP			2%	N89410	001	Dec 01, 1986	Jun	CAHN	
POLOCAINE-MPF									
AP		APP PHARMS	1%	N89406	001	Dec 01, 1986	Jun	CAHN	
AP			1.5%	N89408	001	Dec 01, 1986	Jun	CAHN	
AP			2%	N89409	001	Dec 01, 1986	Jun	CAHN	
<u>MEPROBAMATE</u>									
TABLET; ORAL									
MEPROBAMATE									
AA		ALEMBIC LTD	200MG	N90122	001	Feb 18, 2009	Feb	NEWA	
AA			400MG	N90122	002	Feb 18, 2009	Feb	NEWA	
	@	IVC INDS	400MG	N84153	001		Apr	CAHN	

MEQUINOL; TRETINOIN

SOLUTION; TOPICAL

SOLAGE

+	STIEFEL LABS INC	2%;0.01%	N20922 001	Dec 10, 1999	Jan	CAHN
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MESALAMINE

TABLET, DELAYED RELEASE; ORAL

ASACOL

+	PROCTER AND GAMBLE	800MG	N21830 001	May 29, 2008	May	CTNA
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ASACOL HD

+	PROCTER AND GAMBLE	800MG	N21830 001	May 29, 2008	Jul	CTNA
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MESTRANOL; NORETHINDRONE

TABLET; ORAL-21

NORETHIN 1/50M-21

@	WATSON LABS	0.05MG;1MG	N71539 001	Apr 12, 1988	Apr	DISC
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NORETHINDRONE AND MESTRANOL

@	WATSON LABS	0.05MG;1MG	N70758 001	Jul 01, 1988	Apr	DISC
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NORINYL 1+50 21-DAY

@	WATSON LABS	0.05MG;1MG	N13625 002		Apr	DISC
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TABLET; ORAL-28

NORETHIN 1/50M-28

@	WATSON LABS	0.05MG;1MG	N71540 001	Apr 12, 1988	Apr	DISC
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NORETHINDRONE AND MESTRANOL

@	WATSON LABS	0.05MG;1MG	N70759 001	Jul 01, 1988	Apr	DISC
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NORINYL 1+50 28-DAY

+	WATSON LABS	0.05MG;1MG	N16659 001		Apr	CRLD
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ORTHO-NOVUM 1/50 28

@	ORTHO MCNEIL JANSSEN	0.05MG;1MG	N16709 001		Apr	DISC
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METAPROTERENOL SULFATE

SOLUTION; INHALATION

ALUPENT

@	BOEHRINGER INGELHEIM	0.4%	N18761 002	Oct 10, 1986	Feb	DISC
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@		0.6%	N18761 001	Jun 30, 1983	Feb	DISC
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METFORMIN HYDROCHLORIDE

TABLET; ORAL

METFORMIN HYDROCHLORIDE

AB	ALVOGEN	500MG	N76033 001	Jan 24, 2002	Jan	CAHN
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AB		850MG	N76033 002	Jan 24, 2002	Jan	CAHN
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AB		1GM	N76033 003	Jan 24, 2002	Jan	CAHN
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AB	PROVIDENT PHARM	500MG	N77853 001	Jul 28, 2006	Apr	CAHN
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AB		850MG	N77853 002	Jul 28, 2006	Apr	CAHN
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AB		1GM	N77853 003	Jul 28, 2006	Apr	CAHN
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TABLET, EXTENDED RELEASE; ORAL

METFORMIN HYDROCHLORIDE

@	SANDOZ	500MG	N76223 001	Dec 14, 2004	Mar	DISC
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METFORMIN HYDROCHLORIDE; PIOGLITAZONE HYDROCHLORIDE

TABLET, EXTENDED RELEASE; ORAL

ACTOPLUS MET XR

	TAKEDA GLOBAL	1GM;EQ 15MG BASE	N22024 001	May 12, 2009	May	NEWA
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+		1GM;EQ 30MG BASE	N22024 002	May 12, 2009	May	NEWA
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METHAMPHETAMINE HYDROCHLORIDE

TABLET; ORAL

DESOXYN

+	LUNDBECK INC	5MG	N05378 002		Apr	CAHN
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TABLET, EXTENDED RELEASE; ORAL

DESOXYN

@	LUNDBECK INC	5MG	N05378 004		Apr	CAHN
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@		10MG	N05378 003		Apr	CAHN
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@		15MG	N05378 005		Apr	CAHN
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METHIMAZOLE

TABLET; ORAL

METHIMAZOLE

>D>		CEDAR PHARMS	15MG	N40619 003	Jul 12, 2005	Aug	DISC
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>A>		@	15MG	N40619 003	Jul 12, 2005	Aug	DISC
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>D>	+		20MG	N40547 004	Feb 18, 2005	Aug	DISC
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>A>		@	20MG	N40547 004	Feb 18, 2005	Aug	DISC
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>D>	AB	GENPHARM	10MG	N40350 002	Mar 29, 2000	Aug	CRLD
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>A>	AB	+	10MG	N40350 002	Mar 29, 2000	Aug	CRLD
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	AB		10MG	N40350 002	Mar 29, 2000	May	CRLD
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METHOCARBAMOL

TABLET; ORAL

METHOCARBAMOL

@	SOLCO HLTHCARE	500MG	N86989 001		Jan	CAHN
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@		750MG	N86988 001		Jan	CAHN
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METHOTREXATE SODIUM

INJECTABLE; INJECTION

METHOTREXATE SODIUM PRESERVATIVE FREE

AP		EBEWE PARENTA	EQ 50MG BASE/2ML (EQ 25MG BASE/ML)	N90039 001	Mar 31, 2009	Mar	NEWA
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AP			EQ 250MG BASE/10ML (EQ 25MG BASE/ML)	N90039 002	Mar 31, 2009	Mar	NEWA
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AP			EQ 1GM BASE/40ML (EQ 25MG BASE/ML)	N90029 001	Mar 31, 2009	Mar	NEWA
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METHYLCLOTHIAZIDE

TABLET; ORAL

ENDURON

ABBOTT	2.5MG	N12524 001		Mar	CTEC
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METHYLCLOTHIAZIDE

@ SANDOZ	2.5MG	N89835 001	Aug 18, 1988	Mar	DISC
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@	5MG	N89837 001	Aug 18, 1988	Mar	DISC
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METHYLDOPA

TABLET; ORAL

METHYLDOPA

@ SANDOZ	125MG	N71700 001	Mar 02, 1988	Mar	DISC
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@	250MG	N18934 001	Jun 29, 1984	Mar	DISC
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@	500MG	N18934 002	Jun 29, 1984	Mar	DISC
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>D>	AB	WATSON LABS	250MG	N70703 001	Jun 06, 1986	Aug	DISC
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>A>		@	250MG	N70703 001	Jun 06, 1986	Aug	DISC
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METHYLPREDNISOLONE

TABLET; ORAL

MEDROL

@ PHARMACIA AND UPJOHN 24MG

N11153 005

Jun DISC

METHYLPREDNISOLONE ACETATE

INJECTABLE; INJECTION

METHYLPREDNISOLONE ACETATE

AB SANDOZ 40MG/ML

N40719 001 Jan 29, 2009 Jan NEWA

AB 40MG/ML

N40794 001 Mar 05, 2009 Feb NEWA

AB 80MG/ML

N40719 002 Jan 29, 2009 Jan NEWA

AB 80MG/ML

N40794 002 Mar 05, 2009 Feb NEWA

METOCLOPRAMIDE HYDROCHLORIDE

SOLUTION; ORAL

METOCLOPRAMIDE HYDROCHLORIDE

AA VISTAPHARM EQ 5MG BASE/5ML

N75051 001 Jan 26, 2001 Mar CMFD

TABLET; ORAL

METOCLOPRAMIDE HYDROCHLORIDE

@ SANDOZ EQ 5MG BASE

N74478 001 Oct 05, 1995 Mar DISC

@ EQ 10MG BASE

N72215 001 Jan 30, 1990 Mar DISC

METOPROLOL SUCCINATE

TABLET, EXTENDED RELEASE; ORAL

METOPROLOL SUCCINATE

AB WATSON LABS FLORIDA EQ 25MG TARTRATE

N77118 001 Aug 03, 2009 Jul NEWA

AB EQ 50MG TARTRATE

N76862 001 Aug 03, 2009 Jul NEWA

METOPROLOL TARTRATE

TABLET; ORAL

METOPROLOL TARTRATE

AB MUTUAL PHARM 25MG

N73654 002 Jul 15, 2009 Jun NEWA

@ SOLCO HLTHCARE 50MG

N74453 001 Apr 27, 1995 Jan CAHN

@ 100MG

N74453 002 Apr 27, 1995 Jan CAHN

METRONIDAZOLE

CAPSULE; ORAL

METRONIDAZOLE

AB ALEMBIC LTD 375MG

N79065 001 Jun 23, 2009 Jun NEWA

TABLET; ORAL

METRONIDAZOLE

AB ALEMBIC LTD 250MG

N79067 001 Mar 13, 2009 Mar NEWA

AB 500MG

N79067 002 Mar 13, 2009 Mar NEWA

@ SANDOZ 250MG

N18740 001 Oct 22, 1982 Mar DISC

@ 500MG

N18740 002 Oct 22, 1982 Mar DISC

AB WATSON LABS 250MG

N70035 001 Dec 20, 1984 Jun CAHN

MEXILETINE HYDROCHLORIDE

CAPSULE; ORAL

MEXILETINE HYDROCHLORIDE

@ SANDOZ 150MG

N74450 001 May 16, 1996 Mar DISC

@ 200MG

N74450 002 May 16, 1996 Mar DISC

@ 250MG

N74450 003 May 16, 1996 Mar DISC

MICONAZOLE NITRATE

CREAM; TOPICAL

MONISTAT-DERM

@ ORTHONEUTROGENA 2%

N17494 001

Jun DISC

MICONAZOLE NITRATE; PETROLATUM, WHITE; ZINC OXIDE

OINTMENT; TOPICAL

VUSION

+ STIEFEL LABS INC 0.25%;81.35%;15%

N21026 001 Feb 16, 2006 Jan CAHN

MILNACIPRAN HYDROCHLORIDE

TABLET; ORAL

SAVELLA

CYPRESS BIOSCIENCE 12.5MG

N22256 001 Jan 14, 2009 Jan NEWA

25MG

N22256 002 Jan 14, 2009 Jan NEWA

50MG

N22256 003 Jan 14, 2009 Jan NEWA

+ 100MG

N22256 004 Jan 14, 2009 Jan NEWA

MILRINONE LACTATE

INJECTABLE; INJECTION

MILRINONE LACTATE

>D> AP BAXTER HLTHCARE CORP EQ 1MG BASE/ML

N75852 001 May 28, 2002 Aug DISC

>A> @ EQ 1MG BASE/ML

N75852 001 May 28, 2002 Aug DISC

AP + BEDFORD EQ 1MG BASE/ML

N75660 001 May 28, 2002 Jun CRLD

>D> AP BIONICHE PHARMA EQ 1MG BASE/ML

N76428 001 Jun 16, 2003 Aug DISC

>A> @ EQ 1MG BASE/ML

N76428 001 Jun 16, 2003 Aug DISC

MINOCYCLINE HYDROCHLORIDE

INJECTABLE; INJECTION

MINOCIN

+ TRIAX PHARMS LLC EQ 100MG BASE/VIAL

N50444 001

May CMFD

TABLET, EXTENDED RELEASE; ORAL

MINOCYCLINE HYDROCHLORIDE

AB BARR EQ 45MG BASE

N65485 001 Mar 17, 2009 Mar NEWA

AB EQ 90MG BASE

N65485 002 Mar 17, 2009 Mar NEWA

AB EQ 135MG BASE

N65485 003 Mar 17, 2009 Mar NEWA

AB IMPAX LABS INC EQ 45MG BASE

N90024 001 Feb 03, 2009 Jan NEWA

AB EQ 90MG BASE

N90024 002 Feb 03, 2009 Jan NEWA

AB EQ 135MG BASE

N90024 003 Feb 03, 2009 Jan NEWA

>A> AB SANDOZ EQ 45MG BASE

N90422 001 Aug 13, 2009 Aug NEWA

>A> AB EQ 90MG BASE

N90422 002 Aug 13, 2009 Aug NEWA

>A> AB EQ 135MG BASE

N90422 003 Aug 13, 2009 Aug NEWA

SOLODYN

AB MEDICIS EQ 45MG BASE

N50808 001 May 08, 2006 Jan CFTG

EQ 65MG BASE

N50808 004 Jul 23, 2009 Jul NEWA

AB EQ 90MG BASE

N50808 002 May 08, 2006 Jan CFTG

EQ 115MG BASE

N50808 005 Jul 23, 2009 Jul NEWA

AB + EQ 135MG BASE

N50808 003 May 08, 2006 Jan CFTG

MITOMYCIN

INJECTABLE; INJECTION

MITOMYCIN

>A> AP ACCORD HLTHCARE 40MG/VIAL

N64144 003 Aug 11, 2009 Aug NEWA

MUTAMYCIN

AP + BRISTOL MYERS 40MG/VIAL

N62336 003 Mar 10, 1988 Jul CTEC

MITOXANTRONE HYDROCHLORIDE

INJECTABLE; INJECTION

MITOXANTRONE HYDROCHLORIDE

AP	GENERAMEDIX	EQ 20MG BASE/10ML (EQ 2MG BASE/ML)	N78980 001	Apr 13, 2009	Mar	NEWA
AP		EQ 30MG BASE/15ML (EQ 2MG BASE/ML)	N78980 002	Apr 13, 2009	Mar	NEWA

MIVACURIUM CHLORIDE

INJECTABLE; INJECTION

MIVACURIUM CHLORIDE

+	EBEWE PARENTA	EQ 2MG BASE/ML	N78562 001	Apr 30, 2009	Apr	NEWA
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MORPHINE SULFATE

CAPSULE, EXTENDED RELEASE; ORAL

KADIAN

+	ACTAVIS ELIZABETH	10MG	N20616 008	Apr 20, 2007	Feb	CRLD
+		80MG	N20616 006	Oct 27, 2006	Feb	CRLD

INJECTABLE; INJECTION

ASTRAMORPH PF

AP	APP PHARMS	0.5MG/ML	N71050 001	Oct 07, 1986	Jun	CAHN
AP		0.5MG/ML	N71051 001	Oct 07, 1986	Jun	CAHN
AP		1MG/ML	N71052 001	Oct 07, 1986	Jun	CAHN
AP		1MG/ML	N71053 001	Oct 07, 1986	Jun	CAHN

TABLET, EXTENDED RELEASE; ORAL

MORPHINE SULFATE

@	AB GENERICS	15MG	N74862 001	Jul 07, 1998	Apr	DISC
@		30MG	N74862 002	Jul 07, 1998	Apr	DISC
@		60MG	N74862 003	Jul 07, 1998	Apr	DISC
@		100MG	N74769 001	Jul 02, 1998	Apr	DISC
@		200MG	N74769 002	Jul 02, 1998	Apr	DISC

>A> MORPHINE SULFATE; NALTREXONE HYDROCHLORIDE

>A> CAPSULE, EXTENDED RELEASE; ORAL

>A> EMBEDA

>A>	ALPHARMA KING	20MG;0.8MG	N22321 001	Aug 13, 2009	Aug	NEWA
>A>		30MG;1.2MG	N22321 002	Aug 13, 2009	Aug	NEWA
>A>		50MG;2MG	N22321 003	Aug 13, 2009	Aug	NEWA
>A>		60MG;2.4MG	N22321 004	Aug 13, 2009	Aug	NEWA
>A>		80MG;3.2MG	N22321 005	Aug 13, 2009	Aug	NEWA
>A>	+	100MG;4MG	N22321 006	Aug 13, 2009	Aug	NEWA

MOXIFLOXACIN HYDROCHLORIDE

INJECTABLE; IV (INFUSION)

AVELOX IN SODIUM CHLORIDE 0.8% IN PLASTIC CONTAINER

+	BAYER HLTHCARE	160MG/100ML	N21277 001	Nov 30, 2001	Jun	CAHN
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TABLET; ORAL

AVELOX

+	BAYER HLTHCARE	EQ 400MG BASE	N21085 001	Dec 10, 1999	Jun	CAHN
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MYCOPHENOLATE MOFETIL

CAPSULE; ORAL

MYCOPHENOLATE MOFETIL

AB	ACCORD HLTHCARE INC	250MG	N90253 001	May 04, 2009	Apr	NEWA
AB	APOTEX CORP	250MG	N90419 001	Apr 22, 2009	Apr	NEWA

CAPSULE; ORAL

MYCOPHENOLATE MOFETIL

AB	MYLAN	250MG	N65520 001	May 04, 2009	Apr	NEWA
AB	TEVA PHARMS	250MG	N65491 001	May 06, 2009	Apr	NEWA
AB	ZYDUS PHARMS USA INC	250MG	N65433 001	May 04, 2009	Apr	NEWA

TABLET; ORAL

MYCOPHENOLATE MOFETIL

AB	ACCORD HLTHCARE	500MG	N65416 001	May 04, 2009	Apr	NEWA
AB	APOTEX	500MG	N90499 001	Apr 22, 2009	Apr	NEWA
AB	MYLAN	500MG	N65521 001	May 04, 2009	Apr	NEWA
AB	TEVA PHARMS	500MG	N65457 001	May 04, 2009	Apr	NEWA
AB	ZYDUS PHARMS USA INC	500MG	N65477 001	May 04, 2009	Apr	NEWA

NABUMETONE

TABLET; ORAL

NABUMETONE

AB	ACTAVIS ELIZABETH	500MG	N79093 001	Feb 27, 2009	Feb	NEWA
AB		750MG	N79093 002	Feb 27, 2009	Feb	NEWA
	@ SANDOZ	500MG	N75590 001	Feb 25, 2002	Mar	DISC
	@	750MG	N75590 002	Feb 25, 2002	Mar	DISC

NALIDIXIC ACID

TABLET; ORAL

NALIDIXIC ACID

>D>	AB	WATSON LABS	250MG	N71936 001	Jun 29, 1988	Aug	DISC
>A>		@	250MG	N71936 001	Jun 29, 1988	Aug	DISC
>D>	AB		500MG	N72061 001	Jun 29, 1988	Aug	DISC
>A>		@	500MG	N72061 001	Jun 29, 1988	Aug	DISC
>D>	AB		1GM	N71919 001	Jun 29, 1988	Aug	DISC
>A>		@	1GM	N71919 001	Jun 29, 1988	Aug	DISC

NEGGRAM

>D>	AB	SANOFI AVENTIS US	250MG	N14214 002		Aug	CTEC
>A>			250MG	N14214 002		Aug	CTEC
>D>	AB		500MG	N14214 004		Aug	CTEC
>A>			500MG	N14214 004		Aug	CTEC
>D>	AB	+	1GM	N14214 005		Aug	CTEC
>A>		+	1GM	N14214 005		Aug	CTEC

NALTREXONE HYDROCHLORIDE

TABLET; ORAL

NALTREXONE HYDROCHLORIDE

MALLINCKRODT 100MG

N76264 003 Mar 22, 2002 May CRLD

NAPROXEN

TABLET; ORAL

NAPROXEN

>D>	AB	WATSON LABS	250MG	N74163 001	Feb 10, 1995	Aug	DISC
>A>		@	250MG	N74163 001	Feb 10, 1995	Aug	DISC
>D>	AB		375MG	N74163 002	Feb 10, 1995	Aug	DISC
>A>		@	375MG	N74163 002	Feb 10, 1995	Aug	DISC
>D>	AB		500MG	N74163 003	Feb 10, 1995	Aug	DISC
>A>		@	500MG	N74163 003	Feb 10, 1995	Aug	DISC

NAPROXEN SODIUM

TABLET; ORAL

NAPROXEN SODIUM

>D>	AB	WATSON LABS	EQ 250MG BASE	N74195 001	Dec 21, 1993	Aug	DISC
>A>		@	EQ 250MG BASE	N74195 001	Dec 21, 1993	Aug	DISC
>D>	AB		EQ 500MG BASE	N74195 002	Dec 21, 1993	Aug	DISC
>A>		@	EQ 500MG BASE	N74195 002	Dec 21, 1993	Aug	DISC

TABLET, EXTENDED RELEASE; ORAL

NAPRELAN

+	STAT TRADE	EQ 375MG BASE	N20353 001	Jan 05, 1996	Mar	CTEC
+		EQ 500MG BASE	N20353 002	Jan 05, 1996	Mar	CTEC

NAPROXEN SODIUM

@	WATSON LABS FLORIDA	EQ 375MG BASE	N75416 002	Apr 23, 2003	Mar	DISC
@		EQ 500MG BASE	N75416 001	Aug 27, 2002	Mar	DISC

NATEGLINIDE

TABLET; ORAL

NATEGLINIDE

>A>	AB	DR REDDYS LABS LTD	60MG	N77461 001	Sep 09, 2009	Aug	NEWA
>A>	AB		120MG	N77461 002	Sep 09, 2009	Aug	NEWA
>A>	AB	PAR PHARM	60MG	N77463 001	Sep 09, 2009	Aug	NEWA
>A>	AB		120MG	N77463 002	Sep 09, 2009	Aug	NEWA
>A>	AB	TEVA PHARMS	60MG	N77467 001	Sep 09, 2009	Aug	NEWA
>A>	AB		120MG	N77467 002	Sep 09, 2009	Aug	NEWA

STARLIX

>D>		NOVARTIS	60MG	N21204 001	Dec 22, 2000	Aug	CFTG
>A>	AB		60MG	N21204 001	Dec 22, 2000	Aug	CFTG
>D>		+	120MG	N21204 002	Dec 22, 2000	Aug	CFTG
>A>	AB	+	120MG	N21204 002	Dec 22, 2000	Aug	CFTG

NIFEDIPINE

CAPSULE; ORAL

NIFEDIPINE

>D>	AB	ACTAVIS ELIZABETH	20MG	N72556 001	Sep 20, 1990	Aug	CTEC
>A>			20MG	N72556 001	Sep 20, 1990	Aug	CTEC
>D>	AB	CATALENT	20MG	N74045 001	Apr 30, 1992	Aug	DISC
>A>		@	20MG	N74045 001	Apr 30, 1992	Aug	DISC

PROCARDIA

AB	+	PFIZER	10MG	N18482 001		Apr	CRLD
		@	20MG	N18482 002	Jul 24, 1986	Apr	DISC

TABLET, EXTENDED RELEASE; ORAL

ADALAT CC

AB1		BAYER HLTHCARE	30MG	N20198 001	Apr 21, 1993	Jun	CAHN
AB1	+		60MG	N20198 002	Apr 21, 1993	Jun	CAHN
AB1	+		90MG	N20198 003	Apr 21, 1993	Jun	CAHN

NIMODIPINE

CAPSULE; ORAL

NIMODIPINE

AB	+	BARR	30MG	N77811 001	May 02, 2007	Mar	CRLD
		NIMOTOP					
		@ BAYER PHARMS	30MG	N18869 001	Dec 28, 1988	Feb	DISC

NITISINONE

CAPSULE; ORAL

ORFADIN

	RARE DIS	2MG	N21232 001	Jan 18, 2002	Mar	CAHN
		5MG	N21232 002	Jan 18, 2002	Mar	CAHN
+		10MG	N21232 003	Jan 18, 2002	Mar	CAHN

NITROFURANTOIN, MACROCRYSTALLINE

CAPSULE; ORAL

NITROFURANTOIN

@	SANDOZ	25MG	N74336 001	Jan 25, 1995	Mar	DISC
@		50MG	N74336 002	Jan 25, 1995	Mar	DISC
@		100MG	N74336 003	Jan 25, 1995	Mar	DISC

NORTRIPTYLINE HYDROCHLORIDE

CAPSULE; ORAL

NORTRIPTYLINE HYDROCHLORIDE

>D>	AB	SANDOZ	EQ 10MG BASE	N74835 001	Jun 30, 1997	Aug	DISC
>A>		@	EQ 10MG BASE	N74835 001	Jun 30, 1997	Aug	DISC
>D>	AB		EQ 25MG BASE	N74835 002	Jun 30, 1997	Aug	DISC
>A>		@	EQ 25MG BASE	N74835 002	Jun 30, 1997	Aug	DISC
>D>	AB		EQ 50MG BASE	N74835 003	Jun 30, 1997	Aug	DISC
>A>		@	EQ 50MG BASE	N74835 003	Jun 30, 1997	Aug	DISC
>D>	AB		EQ 75MG BASE	N74835 004	Jun 30, 1997	Aug	DISC
>A>		@	EQ 75MG BASE	N74835 004	Jun 30, 1997	Aug	DISC

OFLOXACIN

SOLUTION/DROPS; OPHTHALMIC

OFLOXACIN

AT	FDC LTD	0.3%	N78559 001	Feb 25, 2009	Feb	NEWA
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SOLUTION/DROPS; OTIC

OFLOXACIN

AT	PHARMAFORCE	0.3%	N90395 001	Aug 11, 2009	Jul	NEWA
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TABLET; ORAL

FLOXIN

@	ORTHO MCNEIL JANSSEN	200MG	N19735 001	Dec 28, 1990	Mar	DISC
@		300MG	N19735 002	Dec 28, 1990	Mar	DISC
@		400MG	N19735 003	Dec 28, 1990	Mar	DISC

OFLOXACIN

@	RANBAXY	200MG	N76220 001	Sep 02, 2003	Jul	DISC
@		300MG	N76220 002	Sep 02, 2003	Jul	DISC
@		400MG	N76220 003	Sep 02, 2003	Jul	DISC

OLSALAZINE SODIUM

CAPSULE; ORAL

DIPENTUM

+	ALAVEN PHARM	250MG	N19715 001	Jul 31, 1990	Apr	CAHN
+	UCB INC	250MG	N19715 001	Jul 31, 1990	Jun	CAHN

OMEPRazole

CAPSULE, DELAYED REL PELLETS; ORAL

OMEPRazole

AB	DR REDDYS LABS	40MG	N78490 001	Apr 17, 2009	Mar	NEWA
AB	DR REDDYS LABS LTD	10MG	N78693 001	Mar 16, 2009	Mar	NEWA
AB		20MG	N78693 002	Mar 16, 2009	Mar	NEWA

CAPSULE, DELAYED REL PELLETS; ORAL

OMEPRAZOLE

AB	KREMERS URBAN DEV	40MG	N75410 003	Jan 23, 2009	Jan	NEWA
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OMEPRAZOLE MAGNESIUM

CAPSULE, DELAYED RELEASE; ORAL

OMEPRAZOLE MAGNESIUM

	DR REDDYS LABS LTD	20MG	N78878 001	Jun 05, 2009	May	NEWA
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ONDANSETRON HYDROCHLORIDE

INJECTABLE; INJECTION

ONDANSETRON HYDROCHLORIDE AND DEXTROSE IN PLASTIC CONTAINER

AP	BEDFORD LABS	EQ 0.64MG BASE/ML	N78291 001	Apr 13, 2009	Mar	NEWA
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ORLISTAT

CAPSULE; ORAL

XENICAL

+	HOFFMANN LA ROCHE	120MG	N20766 001	Apr 23, 1999	Jul	CAHN
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OXACILLIN SODIUM

INJECTABLE; INJECTION

OXACILLIN SODIUM

@	WATSON LABS	EQ 250MG BASE/VIAL	N62856 001	Oct 26, 1988	Apr	DISC
@		EQ 500MG BASE/VIAL	N62856 002	Oct 26, 1988	Apr	DISC
@		EQ 1GM BASE/VIAL	N62856 003	Oct 26, 1988	Apr	DISC
@		EQ 2GM BASE/VIAL	N62856 004	Oct 26, 1988	Apr	DISC
@		EQ 4GM BASE/VIAL	N62856 005	Oct 26, 1988	Apr	DISC

OXALIPLATIN

INJECTABLE; IV (INFUSION)

ELOXATIN

AP	+	SANOFI AVENTIS US	50MG/10ML (5MG/ML)	N21759 001	Jan 31, 2005	Jul	CFTG
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AP	+		100MG/20ML (5MG/ML)	N21759 002	Jan 31, 2005	Jul	CFTG
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OXALIPLATIN

AP		EBEWE PHARMA	50MG/10ML (5MG/ML)	N78812 001	Aug 07, 2009	Jul	NEWA
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AP			100MG/20ML (5MG/ML)	N78812 002	Aug 07, 2009	Jul	NEWA
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AP		FRESENIUS KABI ONCOL	50MG/VIAL	N78810 001	Aug 07, 2009	Jul	NEWA
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AP			100MG/VIAL	N78810 002	Aug 07, 2009	Jul	NEWA
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AP		HOSPIRA WORLDWIDE	50MG/10ML (5MG/ML)	N78813 001	Aug 07, 2009	Jul	NEWA
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AP			100MG/20ML (5MG/ML)	N78813 002	Aug 07, 2009	Jul	NEWA
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AP		SUN PHARMA GLOBAL	50MG/VIAL	N78818 001	Aug 07, 2009	Jul	NEWA
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AP			100MG/VIAL	N78818 002	Aug 07, 2009	Jul	NEWA
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>A>	AP	TEVA PARENTERAL	50MG/10ML (5MG/ML)	N22160 001	Aug 07, 2009	Aug	NEWA
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>A>	AP		100MG/20ML (5MG/ML)	N22160 002	Aug 07, 2009	Aug	NEWA
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OXAPROZIN

TABLET; ORAL

OXAPROZIN

@	SANDOZ	600MG	N75850 001	Apr 27, 2001	Mar	DISC
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OXCARBAZEPINE

SUSPENSION; ORAL

OXCARBAZEPINE

AB		RANBAXY	300MG/5ML	N78734 001	Jun 26, 2009	Jun	NEWA
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TRILEPTAL

AB	+	NOVARTIS	300MG/5ML	N21285 001	May 25, 2001	Jun	CFTG
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OXTRIPHYLLINE

TABLET, EXTENDED RELEASE; ORAL

CHOLEDYL SA

+	WARNER CHILCOTT	400MG	N87863	001	May 24, 1983	Jun	CAHN
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OXYBUTYNIN CHLORIDE

GEL; TRANSDERMAL

GELNIQUE

+	WATSON LABS	10%(100MG/PACKET)	N22204	001	Jan 27, 2009	Mar	CTNA
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OXYBUTYNIN CHLORIDE

+	WATSON LABS	10%(100MG/PACKET)	N22204	001	Jan 27, 2009	Jan	NEWA
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TABLET, EXTENDED RELEASE; ORAL

>D> OXYBUTYNIN CHLORIDE

>D>	AB	OSMOTICA PHARM	5MG	N78503	001	Feb 04, 2009	Aug	CTNA
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	AB		5MG	N78503	001	Feb 04, 2009	Jan	NEWA
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>D>	AB		10MG	N78503	002	Feb 04, 2009	Aug	CTNA
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	AB		10MG	N78503	002	Feb 04, 2009	Jan	NEWA
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>D>	AB		15MG	N78503	003	Feb 04, 2009	Aug	CTNA
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	AB		15MG	N78503	003	Feb 04, 2009	Jan	NEWA
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>A> OXYBUTYNIN CHLORIDE

>A>	AB	OSMOTICA PHARM	5MG	N78503	001	Feb 04, 2009	Aug	CTNA
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>A>	AB		10MG	N78503	002	Feb 04, 2009	Aug	CTNA
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>A>	AB		15MG	N78503	003	Feb 04, 2009	Aug	CTNA
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OXYCODONE HYDROCHLORIDE

TABLET; ORAL

OXYCODONE HYDROCHLORIDE

>A>	AB	AVANTHI INC	5MG	N91393	001	Aug 31, 2009	Aug	NEWA
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>A>	AB		10MG	N91393	002	Aug 31, 2009	Aug	NEWA
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>A>	AB		15MG	N91393	003	Aug 31, 2009	Aug	NEWA
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>A>	AB		20MG	N91393	004	Aug 31, 2009	Aug	NEWA
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>A>	AB		30MG	N91393	005	Aug 31, 2009	Aug	NEWA
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>A>	AB	COREPHARMA	5MG	N90895	001	Aug 24, 2009	Aug	NEWA
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>A>	AB		15MG	N90895	002	Aug 24, 2009	Aug	NEWA
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>A>	AB		30MG	N90895	003	Aug 24, 2009	Aug	NEWA
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>D>		KV PHARM	10MG	N77290	002	Dec 08, 2005	Aug	CTEC
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>A>	AB		10MG	N77290	002	Dec 08, 2005	Aug	CTEC
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>D>			20MG	N77290	004	Dec 08, 2005	Aug	CTEC
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>A>	AB		20MG	N77290	004	Dec 08, 2005	Aug	CTEC
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	AB	SUN PHARM INDS INC	5MG	N90659	001	Apr 10, 2009	Mar	NEWA
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	AB		15MG	N90659	002	Apr 10, 2009	Mar	NEWA
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	AB		30MG	N90659	003	Apr 10, 2009	Mar	NEWA
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	AB	VINTAGE PHARMS	5MG	N77712	003	Mar 02, 2009	Mar	NEWA
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ROXICODONE

AB		XANODYNE PHARMS	5MG	N21011	003	May 15, 2009	May	NEWA
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PALIPERIDONE PALMITATE

SUSPENSION, EXTENDED RELEASE; INTRAMUSCULAR

INVEGA SUSTENNA

JOHNSON AND JOHNSON 39MG/0.25ML (39MG/0.25ML)

N22264	001	Jul 31, 2009	Jul	NEWA
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78MG/0.5ML (78MG/0.5ML)

N22264	002	Jul 31, 2009	Jul	NEWA
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117MG/0.75ML (117MG/0.75ML)

N22264	003	Jul 31, 2009	Jul	NEWA
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156MG/ML (156MG/ML)

N22264	004	Jul 31, 2009	Jul	NEWA
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234MG/1.5ML (156MG/ML)

N22264	005	Jul 31, 2009	Jul	NEWA
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PAMIDRONATE DISODIUM

INJECTABLE; INJECTION

PAMIDRONATE DISODIUM

AP	GENERAMEDIX	30MG/VIAL	N78300 001	Mar 10, 2009	Feb	NEWA
AP		90MG/VIAL	N78300 002	Mar 10, 2009	Feb	NEWA

PANCRELIPASE (AMYLASE;LIPASE;PROTEASE)

CAPSULE; ORAL

CREON

SOLVAY

		30,000USP UNITS;6,000USP UNITS;19,000USP UNITS	N20725 001	Apr 30, 2009	Apr	NEWA
		60,000USP UNITS;12,000USP UNITS;38,000USP UNITS	N20725 002	Apr 30, 2009	Apr	NEWA
+		120,000USP UNITS;24,000USP UNITS;76,000USP UNITS	N20725 003	Apr 30, 2009	Apr	NEWA

CAPSULE, DELAYED RELEASE; ORAL

CREON

SOLVAY

		30,000USP UNITS;6,000USP UNITS;19,000USP UNITS	N20725 001	Apr 30, 2009	Jul	CDFR
		60,000USP UNITS;12,000USP UNITS;38,000USP UNITS	N20725 002	Apr 30, 2009	Jul	CDFR
+		120,000USP UNITS;24,000USP UNITS;76,000USP UNITS	N20725 003	Apr 30, 2009	Jul	CDFR

>A> ZENPEP

>A> EURAND

		27,000USP UNITS;5,000USP UNITS;17,000USP UNITS	N22210 001	Aug 27, 2009	Aug	NEWA
>A>		55,000USP UNITS;10,000USP UNITS;34,000USP UNITS	N22210 002	Aug 27, 2009	Aug	NEWA
>A>		82,000USP UNITS;15,000USP UNITS;51,000USP UNITS	N22210 003	Aug 27, 2009	Aug	NEWA
>A>	+	109,000USP UNITS;20,000USP UNITS;68,000USP UNITS	N22210 004	Aug 27, 2009	Aug	NEWA

PANCURONIUM BROMIDE

INJECTABLE; INJECTION

PANCURONIUM BROMIDE

@ HOSPIRA

2MG/ML

N72321 001 Jan 19, 1989 Apr DISC

PANTOPRAZOLE SODIUM

TABLET, DELAYED RELEASE; ORAL

PANTOPRAZOLE SODIUM

AB	KUDCO IRELAND	EQ 20MG BASE	N78281 001	Mar 17, 2009	Mar	NEWA
AB		EQ 40MG BASE	N78281 002	Mar 17, 2009	Mar	NEWA

PAROMOMYCIN SULFATE

CAPSULE; ORAL

PAROMOMYCIN SULFATE

AA	HERITAGE PHARMS INC	EQ 250MG BASE	N65173 001	Dec 14, 2007	Jul	CAHN
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PENICILLIN G POTASSIUM

INJECTABLE; INJECTION

PENICILLIN G POTASSIUM

>A>	AP	APP PHARMS	5,000,000 UNITS/VIAL	N65448 001	Aug 18, 2009	Aug	NEWA
>A>	AP		20,000,000 UNITS/VIAL	N65448 002	Aug 18, 2009	Aug	NEWA
		HANFORD GC	1,000,000 UNITS/VIAL	N65149 001	Jul 23, 2009	Jul	NEWA
	AP		5,000,000 UNITS/VIAL	N65149 002	Jul 23, 2009	Jul	NEWA
	AP		20,000,000 UNITS/VIAL	N65149 003	Jul 23, 2009	Jul	NEWA

PENICILLIN V POTASSIUM

FOR SOLUTION; ORAL

PENICILLIN-VK

AA	+	TEVA	EQ 250MG BASE/5ML	N60456 002	Apr	CRLD
		VEETIDS				
		@ APOTHECON	EQ 125MG BASE/5ML	N61410 001	Mar	DISC
		@	EQ 250MG BASE/5ML	N61410 002	Mar	DISC

PENTAMIDINE ISETHIONATE

FOR SOLUTION; INHALATION

NEBUPENT

@ APP PHARMS	600MG/VIAL	N19887 002	Mar 22, 1996	May	DISC
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PENTOBARBITAL SODIUM

INJECTABLE; INJECTION

NEMBUTAL SODIUM

+	LUNDBECK INC	50MG/ML	N83246 001	Jun	CAHN
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PENTOXIFYLLINE

TABLET, EXTENDED RELEASE; ORAL

PENTOXIFYLLINE

>D>	AB	ACTAVIS ELIZABETH	400MG	N74878 001	Jul 09, 1997	Aug	DISC
>A>		@	400MG	N74878 001	Jul 09, 1997	Aug	DISC

PHENDIMETRAZINE TARTRATE

CAPSULE; ORAL

PHENDIMETRAZINE TARTRATE

@ SANDOZ	35MG	N85695 001	May	DISC
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PHENTERMINE HYDROCHLORIDE

TABLET; ORAL

PHENTERMINE HYDROCHLORIDE

>A>	AA	BARR	37.5MG	N90470 001	Aug 31, 2009	Aug	NEWA
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PHENYLEPHRINE HYDROCHLORIDE; PROMETHAZINE HYDROCHLORIDE

SYRUP; ORAL

PROMETHAZINE HYDROCHLORIDE AND PHENYLEPHRINE HYDROCHLORIDE

>A>	AA	AMNEAL PHARMS	5MG/5ML;6.25MG/5ML	N40902 001	Aug 25, 2009	Aug	NEWA
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PHENYTOIN

SUSPENSION; ORAL

PHENYTOIN

AB		WOCKHARDT EU	125MG/5ML	N40420 001	Apr 19, 2002	Apr	CAHN
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PHENYTOIN SODIUM

CAPSULE; ORAL

PROMPT PHENYTOIN SODIUM

@ IVAX PHARMS	100MG PROMPT	N80259 001	Jan	DISC
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PILOCARPINE HYDROCHLORIDE

TABLET; ORAL

PILOCARPINE HYDROCHLORIDE

AB		LANNETT	7.5MG	N77220 002	May 06, 2009	Apr	NEWA
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PINDOLOL

TABLET; ORAL

PINDOLOL

@ SANDOZ

5MG

N73608 001 Mar 29, 1993 Mar DISC

@

10MG

N73609 001 Mar 29, 1993 Mar DISC

>A> PITAVASTATIN CALCIUM

>A> TABLET; ORAL

>A> LIVALO

>A> KOWA

EQ 1MG BASE

N22363 001 Aug 03, 2009 Aug NEWA

>A> EQ 2MG BASE

N22363 002 Aug 03, 2009 Aug NEWA

>A> + EQ 4MG BASE

N22363 003 Aug 03, 2009 Aug NEWA

POLYETHYLENE GLYCOL 3350

FOR SOLUTION; ORAL

POLYETHYLENE GLYCOL 3350

AA BRECKENRIDGE PHARM 17GM/SC00PFUL

N77736 001 May 26, 2006 Jun CAHN

AA GAVIS PHARMS 17GM/SC00PFUL

N77736 001 May 26, 2006 Jan CAHN

@ TEVA PHARMS 17GM/SC00PFUL

N77445 001 May 04, 2006 Jan DISC

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

FOR SOLUTION; ORAL

PEG-3350;POTASSIUM CHLORIDE;SODIUM BICARBONATE;SODIUM CHLORIDE

AA NOVEL LABS INC 420GM/BOT;1.48GM/BOT;5.72GM/BOT;1
1.2GM/BOT

N90019 001 May 27, 2009 May NEWA

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

FOR SOLUTION; ORAL

GOLYTELY

+ BRAINTREE 227.1GM/PACKET;2.82GM/PACKET;6.36
GM/PACKET;5.53GM/PACKET;21.5GM/PA
CKET

N19011 002 Jun 02, 1992 May CTEC

AA + 236GM/BOT;2.97GM/BOT;6.74GM/BOT;5
.86GM/BOT;22.74GM/BOT

N19011 001 Jul 13, 1984 May CFTG

+ 236GM/BOT;2.97GM/BOT;6.74GM/BOT;5
.86GM/BOT;22.74GM/BOT

N19011 001 Jul 13, 1984 Mar CTEC

PEG 3350 AND ELECTROLYTES

AA NOVEL LABS INC 236GM/BOT;2.97GM/BOT;6.74GM/BOT;5
.86GM/BOT;22.74GM/BOT

N90231 001 Jun 01, 2009 May NEWA

AA 240GM/BOT;2.98GM/BOT;6.72GM/BOT;5
.84GM/BOT;22.72GM/BOT

N90186 001 Jun 01, 2009 May NEWA

FOR SUSPENSION; ORAL

CO-LAV

@ BOCA PHARMA

240GM/BOT;2.98GM/BOT;6.72GM/BOT;5
.84GM/BOT;22.72GM/BOT

N73428 001 Jan 28, 1992 Mar DISC

GO-EVAC

@ BOCA PHARMA

236GM/BOT;2.97GM/BOT;6.74GM/BOT;5
.86GM/BOT;22.74GM/BOT

N73433 001 Apr 28, 1992 Mar DISC

PORACTANT ALFA

SUSPENSION; INTRATRACHEAL

CUROSURF

>A> + CORNERSTONE THERAP 80MG/ML

N20744 001 Nov 18, 1999 Aug CAHN

>D> + DEY 80MG/ML

N20744 001 Nov 18, 1999 Aug CAHN

POTASSIUM CHLORIDE

CAPSULE, EXTENDED RELEASE; ORAL

MICRO-K

@ KV PHARM	8MEQ	N18238 001		Feb	DISC
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MICRO-K 10

@ KV PHARM	10MEQ	N18238 002	May 14, 1984	Feb	DISC
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POTASSIUM CHLORIDE

@ KV PHARM	10MEQ	N70980 001	Feb 17, 1987	Feb	DISC
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WATSON LABS FLORIDA

8MEQ	N77419 001	Jun 02, 2008	Feb	CTEC
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+	10MEQ	N77419 002	Jun 02, 2008	Feb	CRLD
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INJECTABLE; INJECTION

POTASSIUM CHLORIDE

@ HOSPIRA	1.5MEQ/ML	N83345 001		Apr	DISC
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>D> AP	LUITPOLD	2MEQ/ML	N87584 001		Aug	DISC
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>A>	@	2MEQ/ML	N87584 001		Aug	DISC
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>D> AP		2MEQ/ML	N87585 001		Aug	DISC
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>A>	@	2MEQ/ML	N87585 001		Aug	DISC
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TABLET, EXTENDED RELEASE; ORAL

K+8

>D> AB	ALRA	8MEQ	N70998 001	Jan 25, 1993	Aug	DISC
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>A>	@	8MEQ	N70998 001	Jan 25, 1993	Aug	DISC
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KLOR-CON

>D> AB	UPSHER SMITH	8MEQ	N19123 001	Apr 17, 1986	Aug	CRLD
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>A>	+	8MEQ	N19123 001	Apr 17, 1986	Aug	CRLD
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POTASSIUM CHLORIDE

>D> AB	+	COPLEY PHARM	8MEQ	N70618 001	Sep 09, 1987	Aug	DISC
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>A>	@	8MEQ	N70618 001	Sep 09, 1987	Aug	DISC
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AB	WATSON LABS FLORIDA	10MEQ	N75604 001	Apr 10, 2002	May	CRLD
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AB	+	10MEQ	N75604 001	Apr 10, 2002	Jan	CRLD
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PRASUGREL HYDROCHLORIDE

TABLET; ORAL

EFFIENT

ELI LILLY AND CO

EQ 5MG BASE	N22307 001	Jul 10, 2009	Jul	NEWA
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+	EQ 10MG BASE	N22307 002	Jul 10, 2009	Jul	NEWA
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PRAZQUANTEL

TABLET; ORAL

BILTRICIDE

+ BAYER HLTHCARE

600MG

N18714 001	Dec 29, 1982	Jun	CAHN
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PRAZOSIN HYDROCHLORIDE

CAPSULE; ORAL

PRAZOSIN HYDROCHLORIDE

@ SANDOZ	EQ 1MG BASE	N72576 001	May 16, 1989	Mar	DISC
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@	EQ 2MG BASE	N72577 001	May 16, 1989	Mar	DISC
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@	EQ 5MG BASE	N72578 001	May 16, 1989	Mar	DISC
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>D> AB	WATSON LABS	EQ 5MG BASE	N72609 001	May 16, 1989	Aug	DISC
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>A>	@	EQ 5MG BASE	N72609 001	May 16, 1989	Aug	DISC
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PREDNISOLONE SODIUM PHOSPHATE

SOLUTION; ORAL

ORAPRED

AA + SCIELE PHARMA INC

EQ 15MG BASE/5ML

N75117 001	Dec 14, 2000	Jul	CAHN
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SOLUTION; ORAL

PREDNISOLONE SODIUM PHOSPHATE

AA	AMNEAL PHARMS	EQ 15MG BASE/5ML	N78345 001	Mar 10, 2009	Feb	NEWA
AA	VINTAGE	EQ 15MG BASE/5ML	N79010 001	May 26, 2009	May	NEWA

SOLUTION/DROPS; OPHTHALMIC

PREDNISOLONE SODIUM PHOSPHATE

@ BAUSCH AND LOMB	EQ 0.11% PHOSPHATE	N40065 001	Jul 29, 1994	Apr	DISC
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TABLET, ORALLY DISINTEGRATING; ORAL

ORAPRED ODT

SCIELE PHARMA INC	EQ 10MG BASE	N21959 001	Jun 01, 2006	Jul	CAHN
	EQ 15MG BASE	N21959 002	Jun 01, 2006	Jul	CAHN
+	EQ 30MG BASE	N21959 003	Jun 01, 2006	Jul	CAHN

PREDNISONE

TABLET; ORAL

PREDNISONE

@ SANDOZ	10MG	N89983 001	Jan 12, 1989	Mar	DISC
@	50MG	N89984 001	Jan 12, 1989	Mar	DISC
>D> AB	WATSON LABS	20MG	N86813 001	Aug	DISC
>A> @	20MG	N86813 001		Aug	DISC

PRILOCAINE HYDROCHLORIDE

INJECTABLE; INJECTION

CITANEST PLAIN DENTAL

+ DENTSPLY PHARM	4%	N21382 001		Apr	CTNA
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PROGESTERONE

INJECTABLE; INJECTION

PROGESTERONE

AO	PHARMAFORCE	50MG/ML	N90845 001	Jun 22, 2009	Jul	CTEC
AP		50MG/ML	N90845 001	Jun 22, 2009	Jun	NEWA

PROMETHAZINE HYDROCHLORIDE

SYRUP; ORAL

PROMETHAZINE HYDROCHLORIDE

AA	SUN PHARM INDS INC	6.25MG/5ML	N40891 001	Mar 13, 2009	Mar	NEWA
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TABLET; ORAL

PROMETHAZINE HYDROCHLORIDE

AB	@ SANDOZ	12.5MG	N84176 002	May 22, 2009	Jul	DISC
		12.5MG	N84176 002	May 22, 2009	May	NEWA

PROPANTHELINE BROMIDE

TABLET; ORAL

PROPANTHELINE BROMIDE

>D>	@ ROXANE	7.5MG	N80927 001		Jul	DISC
>A>	+	15MG	N80927 002		Aug	DISC
	@	15MG	N80927 002		Aug	DISC

PROPOFOL

INJECTABLE; INJECTION

DIPRIVAN

AB	@ APP PHARMS	10MG/ML	N19627 001	Oct 02, 1989	Jun	CAHN
	+	10MG/ML	N19627 002	Jun 11, 1996	Jun	CAHN

PROPOXYPHENE HYDROCHLORIDE

CAPSULE; ORAL

DOLENE

AA	HERITAGE PHARMS INC	65MG	N80530 001		Mar	CMFD
	PROPOXYPHENE HYDROCHLORIDE					
AA	VINTAGE PHARMS	65MG	N40908 001	Jul 17, 2009	Jun	NEWA

PROPRANOLOL HYDROCHLORIDE

CAPSULE, EXTENDED RELEASE; ORAL

INDERAL LA

>A>	AB	AKRIMAX PHARMS	60MG	N18553 004	Mar 18, 1987	Aug	CAHN
>A>	AB		80MG	N18553 002	Apr 19, 1983	Aug	CAHN
>A>	AB		120MG	N18553 003	Apr 19, 1983	Aug	CAHN
>A>	AB	+	160MG	N18553 001	Apr 19, 1983	Aug	CAHN
>D>	AB	WYETH PHARMS INC	60MG	N18553 004	Mar 18, 1987	Aug	CAHN
>D>	AB		80MG	N18553 002	Apr 19, 1983	Aug	CAHN
>D>	AB		120MG	N18553 003	Apr 19, 1983	Aug	CAHN
>D>	AB	+	160MG	N18553 001	Apr 19, 1983	Aug	CAHN

PROPRANOLOL HYDROCHLORIDE

AB	UPSHER SMITH	60MG	N78311 001	Mar 06, 2009	Feb	NEWA
AB		80MG	N78311 002	Mar 06, 2009	Feb	NEWA
AB		120MG	N78311 003	Mar 06, 2009	Feb	NEWA
AB		160MG	N78311 004	Mar 06, 2009	Feb	NEWA

TABLET; ORAL

PROPRANOLOL HYDROCHLORIDE

@	SANDOZ	10MG	N70663 001	Jun 13, 1986	Mar	DISC
@		20MG	N70664 001	Jun 13, 1986	Mar	DISC
@		40MG	N70665 001	Jun 13, 1986	Mar	DISC
@		60MG	N70666 001	Oct 10, 1986	Mar	DISC
@		80MG	N70667 001	Jun 13, 1986	Mar	DISC

PSEUDOEPHEDRINE HYDROCHLORIDE; TRIPROLIDINE HYDROCHLORIDE

TABLET; ORAL

CORPHED

@	SANDOZ	60MG;2.5MG	N88602 001	Apr 11, 1985	Mar	DISC
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PSEUDOEPHEDRINE HYDROCHLORIDE AND TRIPROLIDINE HYDROCHLORIDE

+	SANDOZ	60MG;2.5MG	N88193 001	May 17, 1983	Mar	CTEC
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QUINAPRIL HYDROCHLORIDE

TABLET; ORAL

QUINAPRIL HYDROCHLORIDE

AB	SUN PHARM INDS LTD	EQ 5MG BASE	N90800 001	Jun 18, 2009	Jun	NEWA
AB		EQ 10MG BASE	N90800 002	Jun 18, 2009	Jun	NEWA
AB		EQ 20MG BASE	N90800 003	Jun 18, 2009	Jun	NEWA
AB		EQ 40MG BASE	N90800 004	Jun 18, 2009	Jun	NEWA

RAMIPRIL

CAPSULE; ORAL

RAMIPRIL

AB	RANBAXY	5MG	N78849 001	Mar 06, 2009	Feb	NEWA
AB		10MG	N78849 002	Mar 06, 2009	Feb	NEWA

RANITIDINE HYDROCHLORIDE

INJECTABLE; INJECTION

RANITIDINE HYDROCHLORIDE

@ BEDFORD

EQ 25MG BASE/ML

N74764 001 Nov 19, 2004 Apr DISC

SYRUP; ORAL

RANITIDINE HYDROCHLORIDE

AA AMNEAL PHARMS EQ 15MG BASE/ML

N78312 001 Sep 02, 2008 Feb CTNA

>A> AA CYPRESS PHARM EQ 15MG BASE/ML

N78684 001 Aug 27, 2009 Aug NEWA

AA DR REDDYS LABS LTD EQ 15MG BASE/ML

N90102 001 May 26, 2009 May NEWA

AA WOCKHARDT EQ 15MG BASE/ML

N79211 001 May 26, 2009 May NEWA

AA EQ 15MG BASE/ML

N79212 001 Feb 23, 2009 Feb NEWA

TABLET; ORAL

RANITIDINE HYDROCHLORIDE

@ RANBAXY

EQ 150MG BASE

N75439 001 Apr 19, 2000 Jul DISC

@

EQ 300MG BASE

N75439 002 Apr 19, 2000 Jul DISC

RANOLAZINE

TABLET, EXTENDED RELEASE; ORAL

RANEXA

GILEAD

500MG

N21526 002 Jan 27, 2006 Jun CAHN

+

1GM

N21526 001 Feb 12, 2007 Jun CAHN

RISEDRONATE SODIUM

TABLET; ORAL

ACTONEL

@ PROCTER AND GAMBLE

75MG

N20835 004 Apr 16, 2007 May DISC

RISPERIDONE

SOLUTION; ORAL

RISPERDAL

AA + ORTHO MCNEIL JANSSEN 1MG/ML

N20588 001 Jun 10, 1996 Jan CFTG

RISPERIDONE

AA APOTEX INC 1MG/ML

N77719 001 Jul 29, 2009 Jul NEWA

>A> AA AUROBINDO PHARMA 1MG/ML

N78452 001 Sep 04, 2009 Aug NEWA

AA DR REDDYS LABS LTD 1MG/ML

N78909 001 Jul 29, 2009 Jul NEWA

AA ROXANE 1MG/ML

N76904 001 Jul 29, 2009 Jul NEWA

AA TEVA 1MG/ML

N76440 001 Jan 30, 2009 Jan NEWA

TABLET; ORAL

RISPERIDONE

AB ACTAVIS TOTOWA 0.25MG

N78071 001 Jun 17, 2009 Jun NEWA

AB 0.5MG

N78071 002 Jun 17, 2009 Jun NEWA

AB 1MG

N78071 003 Jun 17, 2009 Jun NEWA

AB 2MG

N78071 004 Jun 17, 2009 Jun NEWA

AB 3MG

N78071 005 Jun 17, 2009 Jun NEWA

AB 4MG

N78071 006 Jun 17, 2009 Jun NEWA

AB CADISTA PHARMS 0.25MG

N78828 001 Mar 23, 2009 Mar NEWA

AB 0.5MG

N78828 002 Mar 23, 2009 Mar NEWA

AB 1MG

N78828 003 Mar 23, 2009 Mar NEWA

AB 2MG

N78828 004 Mar 23, 2009 Mar NEWA

AB 3MG

N78828 005 Mar 23, 2009 Mar NEWA

AB 4MG

N78828 006 Mar 23, 2009 Mar NEWA

AB WEST WARD PHARMS 0.25MG

N78740 001 May 29, 2009 May NEWA

AB 0.5MG

N78740 002 May 29, 2009 May NEWA

AB 1MG

N78740 003 May 29, 2009 May NEWA

AB 2MG

N78740 004 May 29, 2009 May NEWA

TABLET; ORAL

RISPERIDONE

AB	WEST WARD PHARMS	3MG	N78740 005	May 29, 2009	May	NEWA
AB		4MG	N78740 006	May 29, 2009	May	NEWA

TABLET, ORALLY DISINTEGRATING; ORAL

RISPERDAL

AB	ORTHO MCNEIL JANSSEN	0.5MG	N21444 001	Apr 02, 2003	Feb	CFTG
AB		2MG	N21444 003	Apr 02, 2003	Feb	CFTG
AB		3MG	N21444 004	Dec 23, 2004	Apr	CFTG
AB		4MG	N21444 005	Dec 23, 2004	Apr	CFTG

RISPERIDONE

AB	DR REDDYS LABS LTD	0.5MG	N77328 001	Feb 24, 2009	Feb	NEWA
AB		2MG	N77328 003	Feb 24, 2009	Feb	NEWA
	KALI LABS	0.25MG	N77494 001	Apr 30, 2009	Apr	NEWA
AB		0.5MG	N77494 002	Apr 30, 2009	Apr	NEWA
AB		2MG	N77494 004	Apr 30, 2009	Apr	NEWA
AB		3MG	N77494 005	Apr 30, 2009	Apr	NEWA
AB		4MG	N77494 006	Apr 30, 2009	Apr	NEWA
	PAR PHARM	0.25MG	N77494 001	Apr 30, 2009	Jun	CAHN
AB		0.5MG	N77494 002	Apr 30, 2009	Jun	CAHN
AB		2MG	N77494 004	Apr 30, 2009	Jun	CAHN
AB		3MG	N77494 005	Apr 30, 2009	Jun	CAHN
AB		4MG	N77494 006	Apr 30, 2009	Jun	CAHN
AB	ZYDUS PHARMS USA	0.5MG	N78516 001	May 01, 2009	Apr	NEWA
AB		2MG	N78516 003	May 01, 2009	Apr	NEWA

RITODRINE HYDROCHLORIDE

INJECTABLE; INJECTION

RITODRINE HYDROCHLORIDE

@	HOSPIRA	10MG/ML	N71618 001	Feb 28, 1991	Apr	DISC
@		15MG/ML	N71619 001	Feb 28, 1991	Apr	DISC
	RITODRINE HYDROCHLORIDE	IN DEXTROSE 5% IN PLASTIC CONTAINER				
@	HOSPIRA	30MG/100ML	N71438 001	Jan 22, 1991	Apr	DISC

ROPINIROLE HYDROCHLORIDE

TABLET; ORAL

ROPINIROLE HYDROCHLORIDE

AB	HUAHAI US INC	EQ 0.25MG BASE	N78110 001	May 05, 2008	Apr	CAHN
AB		EQ 0.5MG BASE	N78110 002	May 05, 2008	Apr	CAHN
AB		EQ 1MG BASE	N78110 003	May 05, 2008	Apr	CAHN
AB		EQ 2MG BASE	N78110 004	May 05, 2008	Apr	CAHN
AB		EQ 3MG BASE	N78110 005	May 05, 2008	Apr	CAHN
AB		EQ 4MG BASE	N78110 006	May 05, 2008	Apr	CAHN
AB	ZYDUS PHARMS USA INC	EQ 0.25MG BASE	N90411 001	Jun 01, 2009	May	NEWA
AB		EQ 0.5MG BASE	N90411 002	Jun 01, 2009	May	NEWA
AB		EQ 1MG BASE	N90411 003	Jun 01, 2009	May	NEWA
AB		EQ 2MG BASE	N90411 004	Jun 01, 2009	May	NEWA
AB		EQ 3MG BASE	N90411 005	Jun 01, 2009	May	NEWA
AB		EQ 4MG BASE	N90411 006	Jun 01, 2009	May	NEWA
AB		EQ 5MG BASE	N90411 007	Jun 01, 2009	May	NEWA

TABLET, EXTENDED RELEASE; ORAL

REQUIP XL

	SMITHKLINE BEECHAM	EQ 6MG BASE	N22008 006	Apr 10, 2009	Apr	NEWA
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SAMARIUM SM 153 LEXIDRONAM PENTASODIUM

INJECTABLE; INJECTION

QUADRAMET

+	EUSA PHARMA USA	50mCi/ML	N20570 001	Mar 28, 1997	May	CAHN
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SAXAGLIPTIN HYDROCHLORIDE

TABLET; ORAL

ONGLYZA

	BRISTOL MYERS SQUIBB	EQ 2.5MG BASE	N22350 001	Jul 31, 2009	Jul	NEWA
+		EQ 5MG BASE	N22350 002	Jul 31, 2009	Jul	NEWA

SELEGILINE HYDROCHLORIDE

TABLET; ORAL

SELEGILINE HYDROCHLORIDE

@	ENDO PHARMS	5MG	N74565 001	Aug 02, 1996	Apr	DISC
@	SIEGFRIED	5MG	N74672 001	Apr 01, 1997	Apr	DISC

SERTACONAZOLE NITRATE

CREAM; TOPICAL

ERTACZO

+	ORTHO DERMATOLOGICS	2%	N21385 001	Dec 10, 2003	Jul	CAHN
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SERTRALINE HYDROCHLORIDE

TABLET; ORAL

SERTRALINE HYDROCHLORIDE

AB	AUSTARPHARMA LLC	EQ 25MG BASE	N78677 001	Mar 04, 2009	Feb	NEWA
AB		EQ 50MG BASE	N78677 002	Mar 04, 2009	Feb	NEWA
AB		EQ 100MG BASE	N78677 003	Mar 04, 2009	Feb	NEWA
AB	HIKMA PHARMS	EQ 25MG BASE	N77864 001	Aug 10, 2009	Jul	NEWA
AB		EQ 50MG BASE	N77864 002	Aug 10, 2009	Jul	NEWA
AB		EQ 100MG BASE	N77864 003	Aug 10, 2009	Jul	NEWA

SEVELAMER CARBONATE

>A> FOR SUSPENSION; ORAL

>A> RENVELA

>A>	GENZYME	800MG/PACKET	N22318 001	Aug 12, 2009	Aug	NEWA
>A>	+	2.4GM/PACKET	N22318 002	Feb 18, 2009	Aug	NEWA

SIMVASTATIN

TABLET; ORAL

SIMVASTATIN

AB	LUPIN	5MG	N78103 005	Apr 14, 2009	Mar	NEWA
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SODIUM PHOSPHATE, P-32

SOLUTION; INJECTION, ORAL

SODIUM PHOSPHATE P 32

@	MALLINCKRODT	0.67mCi/ML	N11777 001		May	DISC
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SODIUM POLYSTYRENE SULFONATE

POWDER; ORAL, RECTAL

KALEXATE

AA	KVK TECH	454GM/BOT	N40905 001	Mar 30, 2009	Mar	NEWA
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SOMATROPIN RECOMBINANT

INJECTABLE; INJECTION

ACCRETROPIN

@ CANGENE

5MG/ML (5MG/ML)

N21538 001 Jan 23, 2008 Jul DISC

NORDITROPIN NORDIFLEX

NOVO NORDISK INC

30MG/3ML

N21148 007 Mar 10, 2009 Mar NEWA

SEROSTIM

@ EMD SERONO

8.8MG/VIAL

N20604 004 Sep 06, 2001 Jul DISC

SORAFENIB TOSYLATE

TABLET; ORAL

NEXAVAR

+ BAYER HLTHCARE

EQ 200MG BASE

N21923 001 Dec 20, 2005 Jun CAHN

SOTALOL HYDROCHLORIDE

SOLUTION; INTRAVENOUS

SOTALOL HYDROCHLORIDE

+ ACADEMIC PHARMS

150MG/10ML (15MG/ML)

N22306 001 Jul 02, 2009 Jul NEWA

SPIRONOLACTONE

TABLET; ORAL

SPIRONOLACTONE

>D> AB MUTUAL PHARM

25MG

N89424 001 Jul 23, 1986 Aug DISC

>A> @

25MG

N89424 001 Jul 23, 1986 Aug DISC

>D> AB

50MG

N89424 002 Aug 11, 1999 Aug DISC

>A> @

50MG

N89424 002 Aug 11, 1999 Aug DISC

>D> AB

100MG

N89424 003 Aug 11, 1999 Aug DISC

>A> @

100MG

N89424 003 Aug 11, 1999 Aug DISC

STANOZOLOL

TABLET; ORAL

WINSTROL

@ LUNDBECK INC

2MG

N12885 001 May 14, 1984 Mar CAHN

STAVUDINE

FOR SOLUTION; ORAL

STAVUDINE

AA CIPLA LTD

1MG/ML

N78030 001 Mar 20, 2009 Mar NEWA

SUCCIMER

CAPSULE; ORAL

CHEMET

+ LUNDBECK INC

100MG

N19998 002 Jan 30, 1991 Mar CAHN

SULFACETAMIDE SODIUM

LOTION; TOPICAL

SULFACETAMIDE SODIUM

AB PERRIGO CO TENNESSEE

10%

N78649 001 Mar 23, 2009 Mar NEWA

AB TARO

10%

N78668 001 May 20, 2009 May NEWA

SOLUTION/DROPS; OPHTHALMIC

ISOPTO CETAMIDE

>D>

>D> AT

+ ALCON

15%

N80020 002

Aug DISC

>A> @

15%

N80020 002

Aug DISC

SULF-10

@ NOVARTIS

10%

N80025 001

Apr DISC

SOLUTION/DROPS; OPHTHALMIC

>D>		SULFACEL -15							
>D>	AT	OPTOPICS	15%	N80024	001		Aug	DISC	
>A>		@	15%	N80024	001		Aug	DISC	
		SULFACETAMIDE SODIUM							
>D>	AT	AKORN	10%	N40215	001	May 25, 1999	Aug	DISC	
>A>		@	10%	N40215	001	May 25, 1999	Aug	DISC	
		@ ALCON	30%	N89068	001	May 05, 1987	Apr	DISC	

SULFAMETHOXAZOLE; TRIMETHOPRIM

TABLET; ORAL

		SULFAMETHOPRIM							
AB		PAR PHARM	400MG;80MG	N70022	001	Feb 15, 1985	Mar	CMFD	
		SULFAMETHOPRIM-DS							
AB		PAR PHARM	800MG;160MG	N70032	001	Feb 15, 1985	Mar	CMFD	
		SULFAMETHOXAZOLE AND TRIMETHOPRIM							
		@ SANDOZ	800MG;160MG	N70890	001	Nov 13, 1986	Mar	DISC	
>D>	AB	WATSON LABS	400MG;80MG	N18852	001	May 09, 1983	Aug	DISC	
>A>		@	400MG;80MG	N18852	001	May 09, 1983	Aug	DISC	
		SULFAMETHOXAZOLE AND TRIMETHOPRIM DOUBLE STRENGTH							
>D>	AB	WATSON LABS	800MG;160MG	N18854	001	May 09, 1983	Aug	DISC	
>A>		@	800MG;160MG	N18854	001	May 09, 1983	Aug	DISC	

SULFISOXAZOLE ACETYL

SUSPENSION; ORAL

GANTRISIN PEDIATRIC

@ ROCHE	EQ 500MG BASE/5ML	N09182	004		Jun	DISC	
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SULINDAC

TABLET; ORAL

SULINDAC

@ SANDOZ	150MG	N72712	001	Aug 30, 1991	Mar	DISC	
@	200MG	N72713	001	Aug 30, 1991	Mar	DISC	

SUMATRIPTAN SUCCINATE

INJECTABLE; SUBCUTANEOUS

IMITREX

AP	+	GLAXOSMITHKLINE	EQ 6MG BASE/0.5ML (EQ 12MG BASE/ML)	N20080	001	Dec 28, 1992	Jan	CFTG	
		SUMATRIPTAN SUCCINATE							
AP		APP PHARMS	EQ 6MG BASE/0.5ML (EQ 12MG BASE/ML)	N79242	001	Mar 02, 2009	Feb	NEWA	
AP		BEDFORD	EQ 6MG BASE/0.5ML (EQ 12MG BASE/ML)	N79123	001	Feb 06, 2009	Jan	NEWA	
AP		JHP PHARMS	EQ 6MG BASE/0.5ML (EQ 12MG BASE/ML)	N77871	001	Jul 09, 2009	Jun	NEWA	
AP		SANDOZ	EQ 4MG BASE/0.5ML (EQ 8MG BASE/ML)	N78067	002	Feb 06, 2009	Jan	NEWA	
AP			EQ 6MG BASE/0.5ML (EQ 12MG BASE/ML)	N78067	001	Feb 06, 2009	Jan	NEWA	
AP		TEVA PARENTERAL	EQ 4MG BASE/0.5ML (EQ 8MG BASE/ML)	N78318	001	Feb 06, 2009	Jan	NEWA	
AP			EQ 6MG BASE/0.5ML (EQ 12MG BASE/ML)	N78318	002	Feb 06, 2009	Jan	NEWA	
AP			EQ 6MG BASE/0.5ML (EQ 12MG BASE/ML)	N77907	001	Feb 06, 2009	Jan	NEWA	
AP		WOCKHARDT	EQ 6MG BASE/0.5ML (EQ 12MG BASE/ML)	N78593	001	Feb 06, 2009	Jan	NEWA	

INJECTABLE; SUBCUTANEOUS

SUMAVEL DOSEPRO

+	ZOGENIX INC	EQ 6MG BASE/0.5ML (EQ 6MG BASE/0.5ML)	N22239 001	Jul 15, 2009	Jul	NEWA
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TABLET; ORAL

IMITREX

AB	GLAXOSMITHKLINE	EQ 25MG BASE	N20132 002	Jun 01, 1995	Jan	CFTG
AB		EQ 50MG BASE	N20132 003	Jun 01, 1995	Jan	CFTG
AB	+	EQ 100MG BASE	N20132 001	Jun 01, 1995	Jan	CFTG

SUMATRIPTAN SUCCINATE

AB	AUROBINDO PHARMA	EQ 25MG BASE	N78327 001	Aug 10, 2009	Jul	NEWA
AB		EQ 50MG BASE	N78327 002	Aug 10, 2009	Jul	NEWA
AB		EQ 100MG BASE	N78327 003	Aug 10, 2009	Jul	NEWA
AB	COBALT LABS INC	EQ 25MG BASE	N76933 001	Aug 10, 2009	Jul	NEWA
AB		EQ 50MG BASE	N76933 002	Aug 10, 2009	Jul	NEWA
AB		EQ 100MG BASE	N76933 003	Aug 10, 2009	Jul	NEWA
AB	DR REDDYS LABS INC	EQ 25MG BASE	N76847 001	Aug 10, 2009	Jul	NEWA
AB		EQ 50MG BASE	N76847 002	Aug 10, 2009	Jul	NEWA
AB		EQ 100MG BASE	N76847 003	Aug 10, 2009	Jul	NEWA
AB	MYLAN	EQ 25MG BASE	N77744 001	Aug 10, 2009	Jul	NEWA
AB		EQ 50MG BASE	N77744 002	Aug 10, 2009	Jul	NEWA
AB		EQ 100MG BASE	N77744 003	Aug 10, 2009	Jul	NEWA
AB	ORCHID HLTHCARE	EQ 25MG BASE	N78284 001	Aug 10, 2009	Jul	NEWA
AB		EQ 50MG BASE	N78284 002	Aug 10, 2009	Jul	NEWA
AB		EQ 100MG BASE	N78284 003	Aug 10, 2009	Jul	NEWA
AB	RANBAXY	EQ 25MG BASE	N76554 001	Aug 10, 2009	Jul	NEWA
AB		EQ 50MG BASE	N76554 002	Aug 10, 2009	Jul	NEWA
AB		EQ 100MG BASE	N76572 001	Feb 09, 2009	Jan	NEWA
AB	ROXANE	EQ 25MG BASE	N78241 001	Aug 10, 2009	Jul	NEWA
AB		EQ 50MG BASE	N78241 002	Aug 10, 2009	Jul	NEWA
AB		EQ 100MG BASE	N78241 003	Aug 10, 2009	Jul	NEWA
AB	SANDOZ	EQ 25MG BASE	N76976 001	Aug 10, 2009	Jul	NEWA
AB		EQ 50MG BASE	N76976 002	Aug 10, 2009	Jul	NEWA
AB		EQ 100MG BASE	N76976 003	Aug 10, 2009	Jul	NEWA
AB	SUN PHARM INDS	EQ 25MG BASE	N78295 001	Aug 10, 2009	Jul	NEWA
AB		EQ 50MG BASE	N78295 002	Aug 10, 2009	Jul	NEWA
AB		EQ 100MG BASE	N78295 003	Aug 10, 2009	Jul	NEWA
AB	TEVA	EQ 25MG BASE	N76840 001	Feb 09, 2009	Jan	NEWA
AB		EQ 50MG BASE	N76840 002	Feb 09, 2009	Jan	NEWA
AB		EQ 100MG BASE	N76840 003	Feb 09, 2009	Jan	NEWA

SUNITINIB MALATE

CAPSULE; ORAL

SUTENT

CPPI CV

EQ 37.5MG BASE	N21938 004	Mar 31, 2009	Mar	NEWA
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TACROLIMUS

CAPSULE; ORAL

PROGRAF

AB	ASTELLAS	EQ 0.5MG BASE	N50708 003	Aug 24, 1998	Jul	CFTG
AB		EQ 1MG BASE	N50708 001	Apr 08, 1994	Jul	CFTG
AB	+	EQ 5MG BASE	N50708 002	Apr 08, 1994	Jul	CFTG

TACROLIMUS

AB	SANDOZ	EQ 0.5MG BASE	N65461 001	Aug 10, 2009	Jul	NEWA
AB		EQ 1MG BASE	N65461 002	Aug 10, 2009	Jul	NEWA
AB		EQ 5MG BASE	N65461 003	Aug 10, 2009	Jul	NEWA

TADALAFIL

TABLET; ORAL

ADCIRCA

+	ELI LILLY CO	20MG	N22332 001	May 22, 2009	May	NEWA
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TAMOXIFEN CITRATE

TABLET; ORAL

TAMOXIFEN CITRATE

@	ROXANE	EQ 10MG BASE	N76027 001	Feb 20, 2003	Mar	DISC
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@		EQ 20MG BASE	N76027 002	Feb 20, 2003	Mar	DISC
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TAPENTADOL HYDROCHLORIDE

TABLET; ORAL

TAPENTADOL HYDROCHLORIDE

	ORTHO MCNEIL JANSSEN	EQ 50MG BASE	N22304 001	Nov 20, 2008	Jun	CPOT
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		EQ 75MG BASE	N22304 002	Nov 20, 2008	Jun	CPOT
--	--	--------------	------------	--------------	-----	------

+		EQ 100MG BASE	N22304 003	Nov 20, 2008	Jun	CPOT
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TECHNETIUM TC-99M BICISATE KIT

INJECTABLE; INJECTION

NEUROLITE

	LANTHEUS MEDCL	N/A	N20256 001	Nov 23, 1994	Apr	CPOT
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TECHNETIUM TC-99M MEDRONATE KIT

INJECTABLE; INJECTION

DRAXIMAGE MDP-10

+	DRAXIMAGE	N/A	N18035 001		Apr	CTEC
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DRAXIMAGE MDP-25

@	DRAXIMAGE	N/A	N18035 002	Feb 27, 2004	Jul	DISC
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+		N/A	N18035 002	Feb 27, 2004	Jun	CMFD
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@		N/A	N18035 002	Feb 27, 2004	Apr	DISC
---	--	-----	------------	--------------	-----	------

TECHNETIUM TC 99M MPI MDP

@	GE HEALTHCARE	N/A	N18141 001		Apr	DISC
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@		N/A	N18141 002	Jun 12, 1989	Apr	DISC
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TECHNETIUM TC-99M PENTETATE KIT

INJECTABLE; INJECTION

MPI DTPA KIT - CHELATE

>D>						
>D>	AP	GE HEALTHCARE	N/A	N17255 001		Aug DISC

>A>		@	N/A	N17255 001		Aug DISC
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TECHNETIUM TC-99M SESTAMIBI KIT

INJECTABLE; INJECTION

TECHNETIUM TC 99M SESTAMIBI

AP	CARDINAL HEALTH 414	N/A	N78809 001	Apr 28, 2009	Apr	NEWA
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AP	DRAXIMAGE	N/A	N78806 001	Apr 29, 2009	Apr	NEWA
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AP	PHARMALUCENCE	N/A/VIAL	N79157 001	Jul 10, 2009	Jun	NEWA
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TECHNETIUM TC-99M TETROFOSMIN KIT

INJECTABLE; INJECTION

MYOVUE 30ML

+	GE HEALTHCARE	N/A	N20372 002	Jul 07, 2005	Jul	CMFD
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TELBIVUDINE

SOLUTION; ORAL

TYZEKA

+	NOVARTIS	100MG/5ML	N22154 001	Apr 28, 2009	Apr	NEWA
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TEMAZEPAM

CAPSULE; ORAL

RESTORIL

>D>		TYCO HLTHCARE	7.5MG	N18163 003	Oct 25, 1991	Aug	CFTG
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>A>	AB		7.5MG	N18163 003	Oct 25, 1991	Aug	CFTG
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	AB		22.5MG	N18163 004	Nov 02, 2004	Jun	CFTG
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TEMAZEPAM

>A>	AB	MUTUAL PHARM	7.5MG	N78581 001	Sep 08, 2009	Aug	NEWA
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>A>	AB		22.5MG	N71175 002	Sep 14, 2009	Aug	NEWA
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	AB	MYLAN	22.5MG	N70920 003	Jun 12, 2009	Jun	NEWA
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	@	NOVEL LABS INC	30MG	N71457 001	Apr 21, 1987	Mar	CAHN
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TEMOZOLOMIDE

POWDER; INTRAVENOUS

TEMODAR

+	SCHERING	100MG/VIAL	N22277 001	Feb 27, 2009	Feb	NEWA
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TERAZOSIN HYDROCHLORIDE

CAPSULE; ORAL

TERAZOSIN HYDROCHLORIDE

AB		IVAX PHARMS	EQ 1MG BASE	N75614 002	Jan 30, 2001	May	CMFD
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AB			EQ 2MG BASE	N75614 001	Jan 30, 2001	May	CMFD
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AB			EQ 5MG BASE	N75614 003	Jan 30, 2001	May	CMFD
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AB			EQ 10MG BASE	N75614 004	Jan 30, 2001	May	CMFD
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TABLET; ORAL

HYTRIN

	ABBOTT	EQ 1MG BASE	N19057 001	Aug 07, 1987	Mar	CTEC
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+		EQ 2MG BASE	N19057 002	Aug 07, 1987	Mar	CTEC
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		EQ 5MG BASE	N19057 003	Aug 07, 1987	Mar	CTEC
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		EQ 10MG BASE	N19057 004	Aug 07, 1987	Mar	CTEC
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TERAZOSIN HYDROCHLORIDE

@	SANDOZ	EQ 1MG BASE	N74315 001	Dec 31, 1998	Mar	DISC
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@		EQ 2MG BASE	N74315 002	Dec 31, 1998	Mar	DISC
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@		EQ 5MG BASE	N74315 003	Dec 31, 1998	Mar	DISC
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@		EQ 10MG BASE	N74315 004	Dec 31, 1998	Mar	DISC
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TERBUTALINE SULFATE

INJECTABLE; INJECTION

TERBUTALINE SULFATE

AP		HIKMA FARMACEUTICA	1MG/ML	N78630 001	May 20, 2009	May	NEWA
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TERIPARATIDE RECOMBINANT HUMAN

INJECTABLE; SUBCUTANEOUS

FORTEO

	LILLY	0.6MG/2.4ML (0.25MG/ML)	N21318 002	Jun 25, 2008	Feb	NEWA
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+		0.75MG/3ML (0.25MG/ML)	N21318 001	Nov 26, 2002	Feb	CPOT
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TESTOSTERONE

		GEL; TRANSDERMAL							
		ANDROGEL							
		UNIMED PHARMS	1% (2.5GM/PACKET)	N21015	001	Feb 28, 2000	Mar	CTEC	
BX	+		1% (5GM/PACKET)	N21015	002	Feb 28, 2000	Mar	CTEC	
		TESTOSTERONE							
		@ WATSON LABS	1% (5GM/PACKET)	N76737	002	Jan 27, 2006	Mar	DISC	
		@	1% (2.5GM/PACKET)	N76737	001	Jan 27, 2006	Mar	DISC	

TESTOSTERONE ENANTHATE

		INJECTABLE; INJECTION							
		DELATESTYL							
AO	+	ENDO PHARM	200MG/ML	N09165	003		Apr	CAHN	
		@	200MG/ML	N09165	001		Apr	CAHN	
		@ ENDO PHARMS	200MG/ML	N09165	001		Mar	CAHN	
AO	+		200MG/ML	N09165	003		Mar	CAHN	

TETRAHYDROZOLINE HYDROCHLORIDE

		SOLUTION; NASAL							
		TYZINE							
	+	NYCOMED US	0.05%	N86576	002		Apr	CAHN	
			0.1%	N86576	001		Apr	CAHN	
		SPRAY; NASAL							
		TYZINE							
	+	NYCOMED US	0.1%	N86576	003		Apr	CAHN	

THEOPHYLLINE

		ELIXIR; ORAL							
		ELIXOPHYLLIN							
	+	FOREST LABS	80MG/15ML	N85186	001		Jun	CRLD	
		THEOPHYLLINE							
		@ MORTON GROVE	80MG/15ML	N86748	001		Jun	DISC	
		@ PRECISION DOSE	80MG/15ML	N85863	001		Jun	DISC	
AA			80MG/15ML	N85863	001		Apr	CAHN	
		@ TARO	80MG/15ML	N89626	001	Oct 28, 1988	Jun	DISC	
		SOLUTION; ORAL							
		THEOPHYLLINE							
		@ ROXANE	80MG/15ML	N87449	001	Sep 15, 1983	Mar	DISC	

THIABENDAZOLE

		TABLET, CHEWABLE; ORAL							
		MINTEZOL							
		@ MERCK	500MG	N16096	001		Jun	DISC	

THIAMINE HYDROCHLORIDE

		INJECTABLE; INJECTION							
		THIAMINE HYDROCHLORIDE							
>D>	AP	BAXTER HLTHCARE	100MG/ML	N80575	001		Aug	DISC	
>A>		@	100MG/ML	N80575	001		Aug	DISC	

THIORIDAZINE HYDROCHLORIDE

>D>	CONCENTRATE; ORAL									
>D>	THIORIDAZINE HYDROCHLORIDE									
>D>	+	ACTAVIS MID ATLANTIC	100MG/ML	N88229	001	Aug 23, 1983	Aug	DISC		
>A>		@	100MG/ML	N88229	001	Aug 23, 1983	Aug	DISC		

TABLET; ORAL

THIORIDAZINE HYDROCHLORIDE

>A>	AB	MYLAN	10MG	N88004 002	Mar 15, 1983	Aug	NEWA
>A>	AB		25MG	N88004 003	Mar 15, 1983	Aug	NEWA
>A>	AB		50MG	N88004 004	Mar 15, 1983	Aug	NEWA

THIOTHIXENE

CAPSULE; ORAL

NAVANE

@ PFIZER

20MG

N16584 005

Jun DISC

THIOTHIXENE

>A>	AB	MYLAN	1MG	N71093 002	Jun 23, 1987	Aug	NEWA
>A>	AB		2MG	N71093 003	Jun 23, 1987	Aug	NEWA
>A>	AB		5MG	N71093 004	Jun 23, 1987	Aug	NEWA
>D>	AB	WATSON LABS	1MG	N70600 001	Jun 05, 1987	Aug	DISC
>A>		@	1MG	N70600 001	Jun 05, 1987	Aug	DISC
>D>	AB		10MG	N70603 001	Jun 05, 1987	Aug	DISC
>A>		@	10MG	N70603 001	Jun 05, 1987	Aug	DISC

TICLOPIDINE HYDROCHLORIDE

TABLET; ORAL

TICLID

@ ROCHE PALO

250MG

N19979 002 Oct 31, 1991 Jun DISC

TICLOPIDINE HYDROCHLORIDE

AB	+	TEVA	250MG	N75149 001	Aug 20, 1999	Jun	CRLD
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TIMOLOL MALEATE

SOLUTION/DROPS; OPHTHALMIC

TIMOPTIC

AT	+	ATON	EQ 0.25% BASE	N18086 001		Feb	CAHN
AT	+		EQ 0.5% BASE	N18086 002		Feb	CAHN

TIMOPTIC IN OCUDOSE

+	ATON	EQ 0.25% BASE	N19463 001	Nov 05, 1986	Feb	CAHN
+		EQ 0.5% BASE	N19463 002	Nov 05, 1986	Feb	CAHN

SOLUTION, GEL FORMING/DROPS; OPHTHALMIC

TIMOPTIC-XE

AB	+	ATON	EQ 0.25% BASE	N20330 001	Nov 04, 1993	Feb	CAHN
AB	+		EQ 0.5% BASE	N20330 002	Nov 04, 1993	Feb	CAHN

TABLET; ORAL

TIMOLOL MALEATE

MYLAN

5MG

N72666 001 Jun 08, 1990 Mar CTEC

10MG

N72667 001 Jun 08, 1990 Mar CTEC

+

20MG

N72668 001 Jun 08, 1990 Mar CTEC

@ SANDOZ

5MG

N72550 001 Apr 13, 1989 Mar DISC

@

10MG

N72551 001 Apr 13, 1989 Mar DISC

@

20MG

N72552 001 Apr 13, 1989 Mar DISC

TOLAZAMIDE

TABLET; ORAL

TOLAZAMIDE

IVAX PHARMS

100MG

N18894 001 Nov 02, 1984 Mar CTEC

@ IVAX SUB TEVA PHARMS

100MG

N18894 001 Nov 02, 1984 Jul DISC

@

250MG

N18894 002 Nov 02, 1984 Jul DISC

@

500MG

N18894 003 Nov 02, 1984 Jul DISC

AB	+	MYLAN	500MG	N70913 001	Mar 17, 1986	Jul	CRLD
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@ SANDOZ

100MG

N71633 001 Dec 09, 1987 Mar DISC

TABLET; ORAL

TOLAZAMIDE

@ SANDOZ

250MG

N70289 001 Mar 13, 1986 Mar DISC

@

500MG

N70290 001 Mar 13, 1986 Mar DISC

TOLBUTAMIDE

TABLET; ORAL

TOLBUTAMIDE

@ SANDOZ

500MG

N86574 001 Mar DISC

TOLMETIN SODIUM

TABLET; ORAL

TOLMETIN SODIUM

>D> AB ACTAVIS ELIZABETH

EQ 600MG BASE

N73527 001 Jun 30, 1992 Aug DISC

>A>

@

EQ 600MG BASE

N73527 001 Jun 30, 1992 Aug DISC

@ SANDOZ

EQ 200MG BASE

N73588 001 Jul 31, 1992 Mar DISC

@

EQ 600MG BASE

N74002 001 Sep 27, 1993 Mar DISC

TOLVAPTAN

TABLET; ORAL

SAMSCA

OTSUKA AMERICA PHARM

15MG

N22275 001 May 19, 2009 May NEWA

+

30MG

N22275 002 May 19, 2009 May NEWA

@

60MG

N22275 003 May 19, 2009 May DISC

TOPIRAMATE

CAPSULE; ORAL

TOPAMAX

AB ORTHO MCNEIL JANSSEN

15MG

N20844 001 Oct 26, 1998 Mar CFTG

AB +

25MG

N20844 002 Oct 26, 1998 Mar CFTG

TOPIRAMATE

AB BARR

15MG

N76448 001 Apr 15, 2009 Mar NEWA

AB

25MG

N76448 002 Apr 15, 2009 Mar NEWA

AB COBALT LABS INC

15MG

N77868 001 Apr 15, 2009 Mar NEWA

AB

25MG

N77868 002 Apr 15, 2009 Mar NEWA

AB TEVA

15MG

N76575 001 Apr 17, 2009 Mar NEWA

AB

25MG

N76575 002 Apr 17, 2009 Mar NEWA

TABLET; ORAL

TOPAMAX

AB + ORTHO MCNEIL JANSSEN

25MG

N20505 004 Dec 24, 1996 Mar CFTG

AB

50MG

N20505 005 Dec 24, 1996 Mar CFTG

AB

100MG

N20505 001 Dec 24, 1996 Mar CFTG

AB

200MG

N20505 002 Dec 24, 1996 Mar CFTG

TOPIRAMATE

AB APOTEX INC

25MG

N77733 001 Mar 27, 2009 Mar NEWA

AB

50MG

N77733 002 Mar 27, 2009 Mar NEWA

AB

100MG

N77733 003 Mar 27, 2009 Mar NEWA

AB

200MG

N77733 004 Mar 27, 2009 Mar NEWA

AB AUROBINDO PHARMA

25MG

N78462 001 Mar 27, 2009 Mar NEWA

AB

50MG

N78462 002 Mar 27, 2009 Mar NEWA

AB

100MG

N78462 003 Mar 27, 2009 Mar NEWA

AB

200MG

N78462 004 Mar 27, 2009 Mar NEWA

AB BARR

25MG

N76315 001 Mar 27, 2009 Mar NEWA

AB

100MG

N76315 002 Mar 27, 2009 Mar NEWA

AB

200MG

N76315 003 Mar 27, 2009 Mar NEWA

AB CIPLA LTD

25MG

N76343 001 Mar 27, 2009 Mar NEWA

TABLET; ORAL

TOPIRAMATE

AB	CIPLA LTD	50MG	N76343 002	Mar 27, 2009	Mar	NEWA
AB		100MG	N76343 003	Mar 27, 2009	Mar	NEWA
AB		200MG	N76343 004	Mar 27, 2009	Mar	NEWA
AB	COBALT LABS INC	25MG	N77643 001	Mar 27, 2009	Mar	NEWA
AB		50MG	N77643 002	Mar 27, 2009	Mar	NEWA
AB		100MG	N77643 003	Mar 27, 2009	Mar	NEWA
AB		200MG	N77643 004	Mar 27, 2009	Mar	NEWA
AB	GLENMARK GENERICS	25MG	N77627 001	Mar 27, 2009	Mar	NEWA
AB		50MG	N77627 002	Mar 27, 2009	Mar	NEWA
AB		100MG	N77627 003	Mar 27, 2009	Mar	NEWA
AB		200MG	N77627 004	Mar 27, 2009	Mar	NEWA
AB	INVAGEN PHARMS	25MG	N79162 001	Mar 27, 2009	Mar	NEWA
AB		50MG	N79162 002	Mar 27, 2009	Mar	NEWA
AB		100MG	N79162 003	Mar 27, 2009	Mar	NEWA
AB		200MG	N79162 004	Mar 27, 2009	Mar	NEWA
AB	MYLAN	25MG	N76314 001	Mar 27, 2009	Mar	NEWA
AB		50MG	N76314 002	Mar 27, 2009	Mar	NEWA
AB		100MG	N76314 003	Mar 27, 2009	Mar	NEWA
AB		200MG	N76314 004	Mar 27, 2009	Mar	NEWA
AB	PAR PHARM	25MG	N76311 001	Mar 27, 2009	Mar	NEWA
AB		50MG	N76311 002	Mar 27, 2009	Mar	NEWA
AB		100MG	N76311 003	Mar 27, 2009	Mar	NEWA
AB		200MG	N76311 004	Mar 27, 2009	Mar	NEWA
AB	PLIVA HRVATSKA DOO	25MG	N77905 001	Mar 30, 2009	Mar	NEWA
AB		50MG	N77905 002	Mar 30, 2009	Mar	NEWA
AB		100MG	N77905 003	Mar 30, 2009	Mar	NEWA
AB		200MG	N77905 004	Mar 30, 2009	Mar	NEWA
AB	RANBAXY	25MG	N76327 001	Mar 27, 2009	Mar	NEWA
AB		100MG	N76327 002	Mar 27, 2009	Mar	NEWA
AB		200MG	N76327 003	Mar 27, 2009	Mar	NEWA
AB	ROXANE	25MG	N76306 001	Mar 27, 2009	Mar	NEWA
AB		50MG	N76306 002	Mar 27, 2009	Mar	NEWA
AB		100MG	N76306 003	Mar 27, 2009	Mar	NEWA
AB		200MG	N76306 004	Mar 27, 2009	Mar	NEWA
AB	SUN PHARM INDS LTD	25MG	N90278 001	Mar 27, 2009	Mar	NEWA
AB		50MG	N90278 002	Mar 27, 2009	Mar	NEWA
AB		100MG	N90278 003	Mar 27, 2009	Mar	NEWA
AB		200MG	N90278 004	Mar 27, 2009	Mar	NEWA
AB	TEVA	25MG	N76317 001	Mar 27, 2009	Mar	NEWA
AB		50MG	N76317 002	Mar 27, 2009	Mar	NEWA
AB		100MG	N76317 003	Mar 27, 2009	Mar	NEWA
AB		200MG	N76317 004	Mar 27, 2009	Mar	NEWA
AB	TORRENT PHARMS	25MG	N79153 001	Mar 27, 2009	Mar	NEWA
AB		50MG	N79153 002	Mar 27, 2009	Mar	NEWA
AB		100MG	N79153 003	Mar 27, 2009	Mar	NEWA
AB		200MG	N79153 004	Mar 27, 2009	Mar	NEWA
AB	UNICHEM	25MG	N90162 001	Mar 27, 2009	Mar	NEWA
AB		50MG	N90162 002	Mar 27, 2009	Mar	NEWA
AB		100MG	N90162 003	Mar 27, 2009	Mar	NEWA
AB	ZYDUS PHARMS USA INC	25MG	N78235 001	Mar 27, 2009	Mar	NEWA
AB		50MG	N78235 002	Mar 27, 2009	Mar	NEWA
AB		100MG	N78235 003	Mar 27, 2009	Mar	NEWA
AB		200MG	N78235 004	Mar 27, 2009	Mar	NEWA

TORSEMIDE

TABLET; ORAL

TORSEMIDE

AB	HETERO DRUGS	5MG	N79234 001	Jan 27, 2009	Jan	NEWA
AB		10MG	N79234 002	Jan 27, 2009	Jan	NEWA
AB		20MG	N79234 003	Jan 27, 2009	Jan	NEWA
AB		100MG	N79234 004	Jan 27, 2009	Jan	NEWA

TRAMADOL HYDROCHLORIDE

TABLET, EXTENDED RELEASE; ORAL

RYZOLT

BC	+	PURDUE PHARMA	100MG	N21745 001	Dec 30, 2008	Mar	CAHN
BC			200MG	N21745 002	Dec 30, 2008	Mar	CAHN
BC			300MG	N21745 003	Dec 30, 2008	Mar	CAHN

TABLET, ORALLY DISINTEGRATING; ORAL

TRAMADOL HYDROCHLORIDE

@ ETHYPHARM NORTH

50MG

N21693 001 May 05, 2005 Mar CAHN

TRAZODONE HYDROCHLORIDE

TABLET; ORAL

TRAZODONE HYDROCHLORIDE

AB		ALVOGEN	50MG	N71636 001	Apr 18, 1988	Feb	CAHN
AB			100MG	N71514 001	Apr 18, 1988	Feb	CAHN
AB		APOTEX INC	300MG	N71196 003	Apr 26, 1999	May	CTEC
AB		MATRIX LABS LTD	50MG	N90514 001	Jun 02, 2009	May	NEWA
AB			100MG	N90514 002	Jun 02, 2009	May	NEWA
AB			150MG	N90514 003	Jun 02, 2009	May	NEWA
AB			300MG	N90514 004	Jun 02, 2009	May	NEWA
		@ SANDOZ	50MG	N72484 001	Apr 30, 1990	Mar	DISC
		@	100MG	N72483 001	Apr 30, 1990	Mar	DISC

TREPROSTINIL SODIUM

SOLUTION; INHALATION

TYVASO

+ UNITED THERAP

EQ 0.6MG BASE/ML

N22387 001 Jul 30, 2009 Jul NEWA

TRETINOIN

CREAM; TOPICAL

RENOVA

	+	ORTHO DERMATOLOGICS	0.02%	N21108 001	Aug 31, 2000	Jul	CAHN
AB2	+		0.05%	N19963 001	Dec 29, 1995	Jul	CAHN

GEL; TOPICAL

ATRALIN

	+	DOW PHARM SCIENCES	0.05%	N22070 001	Jul 26, 2007	Jun	CAHN
		RETIN-A MICRO					
	+	ORTHO DERMATOLOGICS	0.04%	N20475 002	May 10, 2002	Jul	CAHN
	+		0.1%	N20475 001	Feb 07, 1997	Jul	CAHN

TRETINOIN

>D>	AB	SPEAR PHARMS	0.01%	N75589 001	Jun 11, 2002	Aug	CAHN
>D>	AB		0.025%	N75529 001	Feb 22, 2000	Aug	CAHN
>A>	AB	TRIAX PHARMS LLC	0.01%	N75589 001	Jun 11, 2002	Aug	CAHN
>A>	AB		0.025%	N75529 001	Feb 22, 2000	Aug	CAHN

TRIAMCINOLONE ACETONIDE

CREAM; TOPICAL

TRIAMCINOLONE ACETONIDE

>D>	AT	G AND W LABS	0.1%	N89798 001	May 31, 1991	Aug	DISC
>A>		@	0.1%	N89798 001	May 31, 1991	Aug	DISC

INJECTABLE; INJECTION

KENALOG-10

AB		APOTHECON	10MG/ML	N12041 001		May	CFTG
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KENALOG-40

AB	+	APOTHECON	40MG/ML	N14901 001		May	CFTG
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TRIAMCINOLONE ACETONIDE

AB		SANDOZ	10MG/ML	N90166 001	May 27, 2009	May	NEWA
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AB			40MG/ML	N90164 001	May 27, 2009	May	NEWA
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SPRAY, METERED; NASAL

NASACORT AQ

AB	+	SANOFI AVENTIS US	0.055MG/SPRAY	N20468 001	May 20, 1996	Jul	CFTG
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TRIAMCINOLONE ACETONIDE

AB		BARR	0.055MG/SPRAY	N78104 001	Jul 30, 2009	Jul	NEWA
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TROSPIDIUM CHLORIDE

CAPSULE, EXTENDED RELEASE; ORAL

SANCTURA XR

+	ALLERGAN	60MG	N22103 001	Aug 03, 2007	Apr	CAHN
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+	ENDO PHARMS	60MG	N22103 001	Aug 03, 2007	Mar	CAHN
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TABLET; ORAL

SANCTURA

+	ALLERGAN	20MG	N21595 001	May 28, 2004	Apr	CAHN
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+	ENDO PHARMS	20MG	N21595 001	May 28, 2004	Mar	CAHN
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TRYPAN BLUE

SOLUTION; OPHTHALMIC

MEMBRANEBLUE

+	DORC	0.15%	N22278 001	Feb 20, 2009	Feb	NEWA
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UNOPROSTONE ISOPROPYL

SOLUTION/DROPS; OPHTHALMIC

RESCULA

>D>		@ R TECH UENO LTD	0.15%	N21214 001	Aug 03, 2000	Aug	CMFD
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>A>		+	0.15%	N21214 001	Aug 03, 2000	Aug	CMFD
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>D> UROKINASE

>D> INJECTABLE; INJECTION

>D> KINLYTIC

>D>	+	MICROBIX BIOSYSTEMS	250,000 IU/VIAL	N21846 001		Aug	DISC
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>A>		@	250,000 IU/VIAL	N21846 001		Aug	DISC
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URSODIOL

TABLET; ORAL

URSO 250

AB		AXCAN	250MG	N20675 001	Dec 10, 1997	May	CFTG
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URSO FORTE

AB	+	AXCAN	500MG	N20675 002	Jul 21, 2004	May	CFTG
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URSODIOL

AB		TEVA PHARMS	250MG	N79184 001	May 13, 2009	May	NEWA
----	--	-------------	-------	------------	--------------	-----	------

AB			500MG	N79184 002	May 13, 2009	May	NEWA
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VALGANCICLOVIR HYDROCHLORIDE

>A> FOR SOLUTION; ORAL

>A> VALCYTE

>A> + ROCHE PALO 50MG/ML N22257 001 Aug 28, 2009 Aug NEWA

VALRUBICIN

SOLUTION; INTRAVESICAL

VALSTAR PRESERVATIVE FREE

+ ENDO PHARM 40MG/ML N20892 001 Sep 25, 1998 Apr CAHN

+ ENDO PHARMS 40MG/ML N20892 001 Sep 25, 1998 Mar CAHN

VANCOMYCIN HYDROCHLORIDE

INJECTABLE; INJECTION

VANCOMYCIN HYDROCHLORIDE

AP + ABRAXIS PHARM EQ 10GM BASE/VIAL N62663 004 Nov 28, 1997 Apr CTEC

AP HOSPIRA EQ 750MG BASE/VIAL N62912 002 Jan 07, 2009 May CTEC

AP EQ 750MG BASE/VIAL N62933 002 May 27, 2009 May NEWA

AP HOSPIRA INC EQ 10GM BASE/VIAL N65455 001 Apr 29, 2009 Apr NEWA

VECURONIUM BROMIDE

INJECTABLE; INJECTION

VECURONIUM BROMIDE

AP SUN PHARMA GLOBAL 10MG/VIAL N79001 001 Jun 17, 2009 Jun NEWA

AP 20MG/VIAL N79001 002 Jun 17, 2009 Jun NEWA

VENLAFAXINE HYDROCHLORIDE

TABLET; ORAL

VENLAFAXINE HYDROCHLORIDE

>D> AB SANDOZ EQ 25MG BASE N77515 001 Jun 13, 2008 Aug DISC

>A> @ EQ 25MG BASE N77515 001 Jun 13, 2008 Aug DISC

>D> AB EQ 37.5MG BASE N77515 002 Jun 13, 2008 Aug DISC

>A> @ EQ 37.5MG BASE N77515 002 Jun 13, 2008 Aug DISC

>D> AB EQ 50MG BASE N77515 003 Jun 13, 2008 Aug DISC

>A> @ EQ 50MG BASE N77515 003 Jun 13, 2008 Aug DISC

>D> AB EQ 75MG BASE N77515 004 Jun 13, 2008 Aug DISC

>A> @ EQ 75MG BASE N77515 004 Jun 13, 2008 Aug DISC

>D> AB EQ 100MG BASE N77515 005 Jun 13, 2008 Aug DISC

>A> @ EQ 100MG BASE N77515 005 Jun 13, 2008 Aug DISC

VERAPAMIL HYDROCHLORIDE

INJECTABLE; INJECTION

VERAPAMIL HYDROCHLORIDE

@ BEDFORD 2.5MG/ML N72888 001 Jul 28, 1995 Apr DISC

TABLET; ORAL

VERAPAMIL HYDROCHLORIDE

@ SANDOZ 40MG N73168 001 Jul 31, 1992 Mar DISC

@ 80MG N71423 001 May 24, 1988 Mar DISC

@ 120MG N71424 001 May 25, 1988 Mar DISC

>A> VIGABATRIN

>A> FOR SOLUTION; ORAL

>A> SABRIL

>A> + LUNDBECK INC 500MG/PACKET N22006 001 Aug 21, 2009 Aug NEWA

>A>	TABLET; ORAL						
>A>	SABRIL						
>A>	+ LUNDBECK INC	500MG		N20427 001	Aug 21, 2009	Aug	NEWA

VINORELBINE TARTRATE

	INJECTABLE; INJECTION						
	VINORELBINE TARTRATE						
AP	ACTAVIS TOTOWA	EQ 10MG BASE/ML		N78011 001	Jul 22, 2009	Jul	NEWA

ZIDOVUDINE

	TABLET; ORAL						
	ZIDOVUDINE						
	@ AUROBINDO PHARMA	60MG		N22294 001	Jul 23, 2009	Jul	DISC

ZOLPIDEM TARTRATE

	TABLET; ORAL						
	ZOLPIDEM TARTRATE						
>D> AB	MUTUAL PHARMA	5MG		N77288 001	Apr 23, 2007	Aug	DISC
>A>	@	5MG		N77288 001	Apr 23, 2007	Aug	DISC
>D> AB		10MG		N77288 002	Apr 23, 2007	Aug	DISC
>A>	@	10MG		N77288 002	Apr 23, 2007	Aug	DISC

	TABLET; SUBLINGUAL						
	EDLUAR						

	MEDA PHARMS	5MG		N21997 001	Mar 13, 2009	Jul	CAHN
+		10MG		N21997 002	Mar 13, 2009	Jul	CAHN
	OREXO AB	5MG		N21997 001	Mar 13, 2009	Jun	CAHN
		5MG		N21997 001	Mar 13, 2009	Mar	NEWA
+		10MG		N21997 002	Mar 13, 2009	Jun	CAHN
+		10MG		N21997 002	Mar 13, 2009	Mar	NEWA

ZONISAMIDE

	CAPSULE; ORAL						
	ZONISAMIDE						

	@ MUTUAL PHARM	25MG		N77635 001	Dec 22, 2005	May	DISC
	@	50MG		N77635 002	Dec 22, 2005	May	DISC
	@	100MG		N77635 003	Dec 22, 2005	May	DISC
>D> AB	WATSON LABS	25MG		N77650 001	Apr 20, 2006	Aug	DISC
>A>	@	25MG		N77650 001	Apr 20, 2006	Aug	DISC
>D> AB		50MG		N77650 002	Apr 20, 2006	Aug	DISC
>A>	@	50MG		N77650 002	Apr 20, 2006	Aug	DISC
>D> AB		100MG		N77650 003	Apr 20, 2006	Aug	DISC
>A>	@	100MG		N77650 003	Apr 20, 2006	Aug	DISC

OTC DRUG PRODUCT LIST - 29TH EDITION

OTC DRUG PRODUCT LIST - CUMULATIVE SUPPLEMENT 8 - August 2009

2-1

ACETAMINOPHEN

SUPPOSITORY; RECTAL

ACEPHEN

>D>	G AND W LABS	120MG	N72218 001	Mar 27, 1992	Aug	DISC
>A>	@	120MG	N72218 001	Mar 27, 1992	Aug	DISC

TABLET, EXTENDED RELEASE; ORAL

ACETAMINOPHEN

OHM LABS	650MG	N76200 001	Mar 19, 2002	Mar	CAHN
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CETIRIZINE HYDROCHLORIDE

CAPSULE; ORAL

CETIRIZINE HYDROCHLORIDE ALLERGY

BANNER PHARMACAPS	5MG	N22429 001	Jul 23, 2009	Jul	NEWA
	10MG	N22429 004	Jul 23, 2009	Jul	NEWA

CETIRIZINE HYDROCHLORIDE HIVES

BANNER PHARMACAPS	5MG	N22429 003	Jul 23, 2009	Jul	NEWA
+	10MG	N22429 002	Jul 23, 2009	Jul	NEWA

SYRUP; ORAL

CHILDREN'S CETIRIZINE HYDROCHLORIDE ALLERGY

DR REDDYS LABS LTD	5MG/5ML	N90474 002	Mar 30, 2009	Mar	NEWA
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CHILDREN'S CETIRIZINE HYDROCHLORIDE HIVES RELIEF

DR REDDYS LABS LTD	5MG/5ML	N90474 001	Mar 30, 2009	Mar	NEWA
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TABLET, CHEWABLE; ORAL

CETIRIZINE HYDROCHLORIDE ALLERGY

CARACO	5MG	N77631 004	Jan 11, 2008	Jul	NEWA
	10MG	N77631 003	Jan 11, 2008	Jul	NEWA

CETIRIZINE HYDROCHLORIDE HIVES RELIEF

CARACO	5MG	N77631 001	Jan 11, 2008	Jul	CDFR
	10MG	N77631 002	Jan 11, 2008	Jul	CDFR

TABLET; ORAL

CETIRIZINE HYDROCHLORIDE ALLERGY

ORCHID HLTHCARE	5MG	N78862 001	Feb 19, 2009	Feb	NEWA	
	10MG	N78862 002	Feb 19, 2009	Feb	NEWA	
TARO	5MG	N78072 001	Jul 22, 2009	Jul	NEWA	
	5MG	N78072 003	Jul 22, 2009	Jul	NEWA	
UNICHEM	5MG	N78680 003	Jun 26, 2009	Jun	NEWA	
	10MG	N78680 004	Jun 26, 2009	Jun	NEWA	
>A>	UNIQUE PHARM LABS	5MG	N77829 001	Aug 26, 2009	Aug	NEWA
>A>		10MG	N77829 004	Aug 26, 2009	Aug	NEWA

CETIRIZINE HYDROCHLORIDE HIVES

ORCHID HLTHCARE	5MG	N78862 003	Feb 19, 2009	Feb	NEWA	
	10MG	N78862 004	Feb 19, 2009	Feb	NEWA	
UNICHEM	5MG	N78680 001	Jun 26, 2009	Jun	NEWA	
	10MG	N78680 002	Jun 26, 2009	Jun	NEWA	
>A>	UNIQUE PHARM LABS	5MG	N77829 003	Aug 26, 2009	Aug	NEWA
>A>		10MG	N77829 002	Aug 26, 2009	Aug	NEWA

CETIRIZINE HYDROCHLORIDE HIVES RELIEF

TARO	10MG	N78072 002	Jul 22, 2009	Jul	NEWA
	10MG	N78072 004	Jul 22, 2009	Jul	NEWA

CHLORPHENIRAMINE MALEATE

TABLET, EXTENDED RELEASE; ORAL
 CHLORPHENIRAMINE MALEATE
 AVANTHI INC 12MG

N40829 001 May 13, 2009 Apr NEWA

CIMETIDINE

TABLET; ORAL
 CIMETIDINE

>D> APOTEX 100MG
 >A> @ 100MG
 >D> 200MG
 >A> @ 200MG
 >D> LEK PHARMS 200MG
 >A> @ 200MG

N74948 001 Jun 19, 1998 Aug DISC
 N74948 001 Jun 19, 1998 Aug DISC
 N74948 002 Jul 26, 2002 Aug DISC
 N74948 002 Jul 26, 2002 Aug DISC
 N75122 002 Jun 19, 1998 Aug DISC
 N75122 002 Jun 19, 1998 Aug DISC

DEXTROMETHORPHAN POLISTIREX

SUSPENSION, EXTENDED RELEASE; ORAL
 DELSYM
 + RECKITT BENCKISER EQ 30MG HBR/5ML

N18658 001 Oct 08, 1982 Feb CAHN

DIPHENHYDRAMINE CITRATE; IBUPROFEN

TABLET; ORAL
 IBUPROFEN AND DIPHENHYDRAMINE CITRATE
 DR REDDYS LABS LTD 38MG;200MG

N90619 001 Jul 08, 2009 Jun NEWA

EPINEPHRINE

AEROSOL, METERED; INHALATION
 PRIMATENE MIST
 @ WYETH CONS 0.2MG/INH

N16126 001 Jun DISC

EPINEPHRINE BITARTRATE

AEROSOL, METERED; INHALATION
 BRONITIN MIST
 @ WYETH CONS 0.3MG/INH

N16126 002 Jun DISC

FAMOTIDINE

TABLET; ORAL
 FAMOTIDINE

>D> APOTEX 10MG
 >A> @ 10MG

N75610 001 Mar 12, 2002 Aug DISC
 N75610 001 Mar 12, 2002 Aug DISC

IBUPROFEN

CAPSULE; ORAL
 IBUPROFEN

BANNER PHARMACAPS EQ 200MG FREE ACID AND POTASSIUM
 SALT
 DR REDDYS LABS LTD EQ 200MG FREE ACID AND POTASSIUM
 SALT
 EQ 200MG FREE ACID AND POTASSIUM
 MARKSANS PHARMA EQ 200MG FREE ACID AND POTASSIUM
 SALT

N78682 001 Mar 24, 2009 Mar NEWA
 N77338 001 Jul 10, 2009 Jul NEWA
 N77338 001 Jul 10, 2009 Jun NEWA
 N79205 001 Jun 26, 2009 Jun NEWA

MIDOL LIQUID GELS
 + BANNER PHARMACAPS 200MG
 SUSPENSION/DROPS; ORAL

N21472 001 Oct 18, 2002 Feb CTNA

IBUPROFEN

>A> TRIS PHARMA INC 40MG/ML

N79058 001 Aug 31, 2009 Aug NEWA

TABLET; ORAL

IBUPROFEN

@ SANDOZ	200MG	N70733 001	Sep 19, 1986	Mar	DISC
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MEDIPREN

@ MCNEIL	200MG	N70475 001	Feb 06, 1986	Apr	DISC
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@	200MG	N71215 001	Jun 26, 1986	Apr	DISC
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INSULIN RECOMBINANT HUMAN; INSULIN SUSP ISOPHANE RECOMBINANT HUMAN

INJECTABLE; INJECTION

HUMULIN 50/50

@ LILLY	50 UNITS/ML;50 UNITS/ML	N20100 001	Apr 29, 1992	Jan	DISC
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LANSOPRAZOLE

CAPSULE, DELAYED REL PELLETS; ORAL

PREVACID 24 HR

+ NOVARTIS	15MG	N22327 001	May 18, 2009	May	NEWA
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LEVONORGESTREL

TABLET; ORAL

LEVONORGESTREL

>A>	WATSON LABS	0.75MG	N78665 001	Aug 28, 2009	Aug	NEWA
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PLAN B ONE-STEP

+ DURAMED	1.5MG	N21998 001	Jul 10, 2009	Jul	NEWA
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NICOTINE

FILM, EXTENDED RELEASE; TRANSDERMAL

PROSTEP

@ AVEVA	11MG/24HR	N19983 003	Dec 23, 1998	Jun	DISC
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@	22MG/24HR	N19983 004	Dec 23, 1998	Jun	DISC
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NICOTINE POLACRILEX

GUM, CHEWING; BUCCAL

NICOTINE POLACRILEX

IVAX PHARMS

EQ 2MG BASE	N76880 001	Feb 18, 2009	Feb	NEWA
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EQ 4MG BASE	N77850 001	Feb 18, 2009	Feb	NEWA
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WATSON LABS

EQ 2MG BASE	N79044 001	Jul 08, 2009	Jun	NEWA
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EQ 2MG BASE	N79216 001	Jul 08, 2009	Jun	NEWA
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EQ 4MG BASE	N79038 001	Jul 08, 2009	Jun	NEWA
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EQ 4MG BASE	N79219 001	Jul 08, 2009	Jun	NEWA
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TROCHE/LOZENGE; ORAL

NICORETTE

GLAXOSMITHKLINE CONS

EQ 2MG BASE	N22360 001	May 18, 2009	May	NEWA
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+ EQ 4MG BASE	N22360 002	May 18, 2009	May	NEWA
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NICOTINE POLACRILEX

PERRIGO R AND D

EQ 2MG BASE	N90711 001	Jul 10, 2009	Jun	NEWA
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EQ 2MG BASE	N90821 001	Jul 10, 2009	Jun	NEWA
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EQ 4MG BASE	N90711 002	Jul 10, 2009	Jun	NEWA
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EQ 4MG BASE	N90821 002	Jul 10, 2009	Jun	NEWA
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POTASSIUM IODIDE

SOLUTION; ORAL

POTASSIUM IODIDE

@ ROXANE	1GM/ML	N18551 001	Feb 19, 1982	Jul	DISC
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THYROSHIELD

+ FLEMING	65MG/ML	N77218 001	Jan 12, 2005	Apr	CRLD
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RANITIDINE HYDROCHLORIDE

TABLET; ORAL

RANITIDINE

@ RANBAXY

EQ 75MG BASE

N75132 001 Jan 14, 2000 Jul DISC

@ SANDOZ

EQ 75MG BASE

N75519 001 Sep 26, 2002 Mar DISC

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

CUMULATIVE SUPPLEMENT NUMBER 08 AUGUST 2009

NO AUGUST 2009 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 AUGUST 2009 ADDITIONS

PRESCRIPTION AND OTC DRUG PRODUCT PATENT AND EXCLUSIVITY LIST - 29TH EDITION

A - 1

PATENT & EXCLUSIVITY DRUG PRODUCT LIST - CUMULATIVE SUPPLEMENT 8 - August 2009

See List footnote for information regarding List content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	PATENT DELIST REQUESTED	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>ACYCLOVIR; HYDROCORTISONE - ACYCLOVIR AND HYDROCORTISONE</u>						
022436 001					>A> NC	Jul 31, 2012
<u>ADAPALENE - DIFFERIN</u>						
021753 001	>A> 7579377	Feb 23, 2025	U-818			
<u>ADAPALENE; BENZOYL PEROXIDE - EPIDUO</u>						
022320 001	4717720	May 31, 2010	DS DP			
	RE34440	May 31, 2010	U-818			
<u>ALBUTEROL SULFATE - VENTOLIN HFA</u>						
020983 001	6161724	Jan 16, 2018	DP			
	6161724*PED	Jul 16, 2018				
	6170717	Dec 23, 2017	DP			
	6170717*PED	Jun 23, 2018				
	6431168	Jun 08, 2018	DP			
	6431168*PED	Dec 08, 2018				
	6435372	Jan 16, 2018	DP			
	6435372*PED	Jul 16, 2018				
	6596260	Aug 10, 2014	DP			
	6596260*PED	Feb 10, 2015				
	6938796	Jan 16, 2018	DP			
	6938796*PED	Jul 16, 2018				
	6966467	Dec 23, 2017	DP			
	6966467*PED	Jun 23, 2018				
	6997349	Jan 16, 2018	DP			
	6997349*PED	Jul 16, 2018				
	7107986	Jun 08, 2018	DP			
	7107986*PED	Dec 06, 2018				
	7143908	Jan 16, 2018	DP			
	7143908*PED	Jul 16, 2018				
	7350676	Aug 24, 2018	DP			
	7350676*PED	Feb 24, 2019				
	7500444	Jan 04, 2025	DP			
	7500444*PED	Jul 04, 2025				
<u>ALISKIREN FUMARATE; HYDROCHLOROTHIAZIDE - TEKTRUNA HCT</u>						
022107 003					I-600	Jul 16, 2012
<u>ALISKIREN HEMIFUMARATE; HYDROCHLOROTHIAZIDE - TEKTRUNA HCT</u>						
022107 001					I-600	Jul 16, 2012
<u>ALISKIREN HEMIFUMARATE; HYDROCHLOROTHIAZIDE - TEKTRUNA HCT</u>						
022107 002					I-600	Jul 16, 2012
<u>ALISKIREN HEMIFUMARATE; HYDROCHLOROTHIAZIDE - TEKTRUNA HCT</u>						
022107 004					I-600	Jul 16, 2012
<u>ALMOTRIPTAN MALATE - AXERT</u>						
021001 001	5565447	May 07, 2015	DS DP U-969			
	5565447*PED	Nov 07, 2015				
<u>ALMOTRIPTAN MALATE - AXERT</u>						
021001 002	5565447	May 07, 2015	DS DP U-969			
	5565447*PED	Nov 07, 2015				

PATENT & EXCLUSIVITY DRUG PRODUCT LIST - CUMULATIVE SUPPLEMENT 8 - August 2009

See List footnote for information regarding List content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	PATENT DELIST REQUESTED	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>ALPRAZOLAM - ALPRAZOLAM</u>						
078088 001					PC	Jul 13, 2009
<u>ALPRAZOLAM - ALPRAZOLAM</u>						
078088 002					PC	Jul 13, 2009
<u>ALPRAZOLAM - ALPRAZOLAM</u>						
078088 003					PC	Jul 13, 2009
<u>ALPRAZOLAM - ALPRAZOLAM</u>						
078088 004					PC	Jul 13, 2009
<u>AMIODARONE HYDROCHLORIDE - NEXTERONE</u>						
022325 001	5134127	Jan 23, 2010	DP			
	5376645	Jan 23, 2010	DP			
	6869939	May 04, 2022	DP			
<u>AMLODIPINE BESYLATE; ATORVASTATIN CALCIUM - CADUET</u>						
021540 001	RE40667	Dec 28, 2010	DS DP U-162			
	RE40667*PED	Jun 28, 2011				
<u>AMLODIPINE BESYLATE; ATORVASTATIN CALCIUM - CADUET</u>						
021540 002	RE40667	Dec 28, 2010	DS DP U-162			
	RE40667*PED	Jun 28, 2011				
<u>AMLODIPINE BESYLATE; ATORVASTATIN CALCIUM - CADUET</u>						
021540 003	RE40667	Dec 28, 2010	DS DP U-162			
	RE40667*PED	Jun 28, 2011				
<u>AMLODIPINE BESYLATE; ATORVASTATIN CALCIUM - CADUET</u>						
021540 004	RE40667	Dec 28, 2010	DS DP U-162			
	RE40667*PED	Jun 28, 2011				
<u>AMLODIPINE BESYLATE; ATORVASTATIN CALCIUM - CADUET</u>						
021540 005	RE40667	Dec 28, 2010	DS DP U-162			
	RE40667*PED	Jun 28, 2011				
<u>AMLODIPINE BESYLATE; ATORVASTATIN CALCIUM - CADUET</u>						
021540 006	RE40667	Dec 28, 2010	DS DP U-162			
	RE40667*PED	Jun 28, 2011				
<u>AMLODIPINE BESYLATE; ATORVASTATIN CALCIUM - CADUET</u>						
021540 007	RE40667	Dec 28, 2010	DS DP U-162			
	RE40667*PED	Jun 28, 2011				
<u>AMLODIPINE BESYLATE; ATORVASTATIN CALCIUM - CADUET</u>						
021540 008	RE40667	Dec 28, 2010	DS DP U-162			
	RE40667*PED	Jun 28, 2011				
<u>AMLODIPINE BESYLATE; ATORVASTATIN CALCIUM - CADUET</u>						
021540 009	RE40667	Dec 28, 2010	DS DP U-162			
	RE40667*PED	Jun 28, 2011				
<u>AMLODIPINE BESYLATE; ATORVASTATIN CALCIUM - CADUET</u>						
021540 010	RE40667	Dec 28, 2010	DS DP U-162			
	RE40667*PED	Jun 28, 2011				

PATENT & EXCLUSIVITY DRUG PRODUCT LIST - CUMULATIVE SUPPLEMENT 8 - August 2009

See List footnote for information regarding List content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	PATENT DELIST REQUESTED	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>AMLODIPINE BESYLATE; ATORVASTATIN CALCIUM - CADUET</u>						
021540 011	RE40667	Dec 28, 2010	DS DP U-162			
	RE40667*PED	Jun 28, 2011				
<u>AMLODIPINE; HYDROCHLOROTHIAZIDE; VALSARTAN - EXFORGE HCT</u>						
022314 001	5399578	Mar 21, 2012	DS DP U-3		NC	Apr 30, 2012
	5399578*PED	Sep 21, 2012				
	6294197	Jun 18, 2017	DP U-3			
	6294197*PED	Dec 18, 2017				
<u>AMLODIPINE; HYDROCHLOROTHIAZIDE; VALSARTAN - EXFORGE HCT</u>						
022314 002	5399578	Mar 21, 2012	DS DP U-3		NC	Apr 30, 2012
	5399578*PED	Sep 21, 2012				
	6294197	Jun 18, 2017	DP U-3			
	6294197*PED	Dec 18, 2017				
<u>AMLODIPINE; HYDROCHLOROTHIAZIDE; VALSARTAN - EXFORGE HCT</u>						
022314 003	5399578	Mar 21, 2012	DS DP U-3		NC	Apr 30, 2012
	5399578*PED	Sep 21, 2012				
	6294197	Jun 18, 2017	DP U-3			
	6294197*PED	Dec 18, 2017				
<u>AMLODIPINE; HYDROCHLOROTHIAZIDE; VALSARTAN - EXFORGE HCT</u>						
022314 004	5399578	Mar 21, 2012	DS DP U-3		NC	Apr 30, 2012
	5399578*PED	Sep 21, 2012				
	6294197	Jun 18, 2017	DP U-3			
	6294197*PED	Dec 18, 2017				
<u>AMLODIPINE; HYDROCHLOROTHIAZIDE; VALSARTAN - EXFORGE HCT</u>						
022314 005	5399578	Mar 21, 2012	DS DP U-3		NC	Apr 30, 2012
	5399578*PED	Sep 21, 2012				
	6294197	Jun 18, 2017	DP U-3			
	6294197*PED	Dec 18, 2017				
<u>AMMONIA, N-13 - AMMONIA N 13</u>						
022119 001					NCE W	Aug 23, 2012 Aug 23, 2012
<u>AMOXICILLIN; CLARITHROMYCIN; LANSOPRAZOLE - PREVPAC</u>						
050757 001	4628098	May 10, 2009	DS			
	4628098*PED	Nov 10, 2009				
	5013743	Feb 12, 2010			U-452	
	5013743*PED	Aug 12, 2010				
	5045321	Sep 03, 2008	DP			
	5045321*PED	Mar 03, 2009				
	5093132	Sep 03, 2008	DP			
	5093132*PED	Mar 03, 2009				
	5433959	Sep 03, 2008	DP			
	5433959*PED	Mar 03, 2009				
<u>ANASTROZOLE - ARIMIDEX</u>						
020541 001	RE36617	Dec 27, 2009	DS DP U-946			
<u>ANIDULAFUNGIN - ERAXIS</u>						
021632 001	5965525	Feb 17, 2020	DS DP U-540			

PATENT & EXCLUSIVITY DRUG PRODUCT LIST - CUMULATIVE SUPPLEMENT 8 - August 2009

See List footnote for information regarding List content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	PATENT DELIST REQUESTED	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>ANIDULAFUNGIN - ERAXIS</u>						
021632 002	5965525	Feb 17, 2020	DS DP U-540			
<u>ARIPIRAZOLE - ABILIFY</u>						
021866 001	7550445	Jul 21, 2024	DP			
	7550445*PED	Jan 21, 2025				
<u>ARMODAFINIL - NUVIGIL</u>						
021875 001	4927855	Apr 22, 2010	DS DP U-820			
	4927855*PED	Oct 22, 2010				
<u>ARMODAFINIL - NUVIGIL</u>						
021875 002	4927855	Apr 22, 2010	DS DP U-820		NP	Jun 15, 2010
	4927855*PED	Oct 22, 2010				
	7297346	Nov 29, 2023	DP			
	7297346*PED	May 29, 2024				
<u>ARMODAFINIL - NUVIGIL</u>						
021875 003	4927855	Apr 22, 2010	DS DP U-820			
	4927855*PED	Oct 22, 2010				
<u>ARMODAFINIL - NUVIGIL</u>						
021875 004	4927855	Apr 22, 2010	DS DP U-820			
	4927855*PED	Oct 22, 2010				
<u>ARMODAFINIL - NUVIGIL</u>						
021875 005	4927855	Apr 22, 2010	DS DP U-820		NP	Jun 15, 2010
	4927855*PED	Oct 22, 2010				
	7132570	Dec 18, 2023	DS DP			
	7132570*PED	Jun 18, 2024				
	7297346	Nov 29, 2023	DP			
	7297346*PED	May 29, 2024				
	RE37516	Oct 06, 2014	DP U-820			
	RE37516*PED	Apr 06, 2015				
<u>ARSENIC TRIOXIDE - TRISENOX</u>						
021248 001	6982096	Nov 10, 2018	U-651			
<u>ARTEMETHER; LUMEFANTRINE - COARTEM</u>						
022268 001	5677331	Oct 14, 2014	DP U-977		NCE ODE	Apr 07, 2014 Apr 07, 2016
<u>ASENAPINE - SAPHRIS</u>						
022117 001	>A> 5763476	Jun 09, 2015	DP U-326		>A> NCE	Aug 13, 2014
<u>ASENAPINE - SAPHRIS</u>						
022117 002	>A> 5763476	Jun 09, 2015	DP U-326		>A> NCE	Aug 13, 2014
<u>ATORVASTATIN CALCIUM - LIPITOR</u>						
020702 001	RE40667	Dec 28, 2010	DS DP U-162			
	RE40667*PED	Jun 28, 2011				
<u>ATORVASTATIN CALCIUM - LIPITOR</u>						
020702 002	RE40667	Dec 28, 2010	DS DP U-162			
	RE40667*PED	Jun 28, 2011				

PATENT & EXCLUSIVITY DRUG PRODUCT LIST - CUMULATIVE SUPPLEMENT 8 - August 2009

See List footnote for information regarding List content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	PATENT DELIST REQUESTED	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>ATORVASTATIN CALCIUM - LIPITOR</u>						
020702 003	RE40667	Dec 28, 2010	DS DP U-162			
	RE40667*PED	Jun 28, 2011				
<u>ATORVASTATIN CALCIUM - LIPITOR</u>						
020702 004	RE40667	Dec 28, 2010	DS DP U-162			
	RE40667*PED	Jun 28, 2011				
<u>ATOVAQUONE; PROGUANIL HYDROCHLORIDE - MALARONE</u>						
021078 001	5998449	Nov 25, 2013	U-990			
	5998449*PED	May 25, 2014				
<u>ATOVAQUONE; PROGUANIL HYDROCHLORIDE - MALARONE PEDIATRIC</u>						
021078 002	5998449	Nov 25, 2013	U-990			
	5998449*PED	May 25, 2014				
<u>AZELASTINE HYDROCHLORIDE - ASTEPRO</u>						
022371 001					>A> NP	Aug 31, 2012
<u>AZITHROMYCIN - AZASITE</u>						
050810 001	5192535	Mar 09, 2010	DP U-709			
	6159458	Nov 04, 2017	DP U-709			
	6239113	Mar 31, 2019	U-709			
	6569443	Mar 31, 2019	DP U-709			
	6861411	Nov 25, 2018	U-709			
	7056893	Mar 31, 2019	DP U-709			
<u>BENDAMUSTINE HYDROCHLORIDE - TREANDA</u>						
022249 002					I-580 NCE ODE	Oct 31, 2011 Mar 20, 2013 Mar 20, 2015
<u>BENZOYL PEROXIDE; CLINDAMYCIN PHOSPHATE - ACANYA</u>						
050819 001	5733886	Mar 31, 2015	DP U-124			
	6117843	Feb 18, 2012	DP			
<u>BENZOYL PEROXIDE; CLINDAMYCIN PHOSPHATE - DUAC</u>						
050741 001	5466446	Feb 16, 2014	DS DP			
<u>BENZYL ALCOHOL - ULESFIA</u>						
022129 001	5858383	Aug 11, 2017	U-970		NCE	Apr 09, 2014
	6139859	Aug 11, 2017	U-970			
	6793931	Jul 11, 2022	DP U-970			
	7294342	May 19, 2024	U-970			
<u>BESIFLOXACIN HYDROCHLORIDE - BESIVANCE</u>						
022308 001	5447926	Sep 05, 2012	DS DP U-80		NCE	May 28, 2014
	6685958	Jun 29, 2021	DP U-80			
	6699492	Mar 31, 2019	DP U-80			
<u>BETAMETHASONE VALERATE - LUXIQ</u>						
020934 001	7078058	May 24, 2017	DP			
<u>BIMATOPROST - LATISSE</u>						
022369 001					NP	Dec 24, 2011

PATENT & EXCLUSIVITY DRUG PRODUCT LIST - CUMULATIVE SUPPLEMENT 8 - August 2009

See List footnote for information regarding List content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	PATENT DELIST REQUESTED	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>BISMUTH SUBCITRATE POTASSIUM; METRONIDAZOLE; TETRACYCLINE - PYLERA</u>						
050786 001	5196205	Mar 23, 2010	U-933			
	5476669	Mar 23, 2010	U-933			
	6350468	Dec 14, 2018	U-956			
	6350468	Dec 14, 2018	U-932			
<u>BIVALIRUDIN - ANGIOMAX</u>						
020873 001	5196404	Mar 23, 2010				
	5196404*PED	Sep 23, 2010				
	>A> 7582727	Jul 27, 2028	DP			
	>A> 7582727*PED	Jan 27, 2029				
<u>BOSENTAN - TRACLEER</u>						
021290 001					>A> I-607	Aug 07, 2012
<u>BOSENTAN - TRACLEER</u>						
021290 002					>A> I-607	Aug 07, 2012
<u>BRIMONIDINE TARTRATE - BRIMONIDINE TARTRATE</u>						
021764 001	>A> 7265117	Aug 19, 2025	DP			
<u>BRIMONIDINE TARTRATE; TIMOLOL MALEATE - COMBIGAN</u>						
021398 001	7323463	Jan 19, 2023	DP			
<u>BROMOCRIPTINE MESYLATE - CYCLOSET</u>						
020866 001	5468755	Nov 21, 2012	U-976		NP	May 05, 2012
	5679685	Oct 21, 2014	DP			
	5716957	Feb 10, 2015	U-976			
	5756513	Nov 21, 2012	U-976			
	5866584	Nov 21, 2012	U-976			
<u>BUDESONIDE - PULMICORT RESPULES</u>						
020929 001	7524834	Nov 11, 2018	DP U-966			
	7524834*PED	May 11, 2019				
<u>BUDESONIDE - PULMICORT RESPULES</u>						
020929 002	7524834	Nov 11, 2018	DP U-966			
	7524834*PED	May 11, 2019				
<u>BUDESONIDE - PULMICORT RESPULES</u>						
020929 003	7524834	Nov 11, 2018	DP U-966			
	7524834*PED	May 11, 2019				
<u>BUDESONIDE; FORMOTEROL FUMARATE DIHYDRATE - SYMBICORT</u>						
021929 001					I-582	Feb 27, 2012
<u>BUDESONIDE; FORMOTEROL FUMARATE DIHYDRATE - SYMBICORT</u>						
021929 002					I-582	Feb 27, 2012
<u>BUPROPION HYDROBROMIDE - APLENZIN</u>						
022108 001	>A> 7569610	Jun 27, 2026	U-997			
	>A> 7572935	Jun 27, 2026	DP			
<u>BUPROPION HYDROBROMIDE - APLENZIN</u>						
022108 002	>A> 7569610	Jun 27, 2026	U-997			
	>A> 7572935	Jun 27, 2026	DP			

PATENT & EXCLUSIVITY DRUG PRODUCT LIST - CUMULATIVE SUPPLEMENT 8 - August 2009

See List footnote for information regarding List content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	PATENT DELIST REQUESTED	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>BUPROPION HYDROBROMIDE - APLENZIN</u>						
022108 003	>A> 7569610	Jun 27, 2026	U-997			
	>A> 7572935	Jun 27, 2026	DP			
<u>CALCITONIN SALMON RECOMBINANT - FORTICAL</u>						
021406 001	RE40812	Feb 02, 2021	DP			
<u>CALCITRIOL - VECTICAL</u>						
022087 001					NDF	Jan 23, 2012
<u>CALCIUM CARBONATE; FAMOTIDINE; MAGNESIUM HYDROXIDE - PEPICID COMPLETE</u>						
020958 001	5075114	May 23, 2010	DP			
	5075114*PED	Nov 23, 2010				
	6814978	Aug 26, 2021	DP			
	6814978*PED	Feb 26, 2022				
<u>CALCIUM CARBONATE; RISEDRONATE SODIUM - ACTONEL WITH CALCIUM (COPACKAGED)</u>						
021823 001	5583122	Dec 10, 2013	DS DP U-353			
	5583122*PED	Jun 10, 2014				
	5994329	Jul 17, 2018	U-353			
	5994329*PED	Jan 17, 2019				
	6015801	Jul 17, 2018	U-353			
	6015801*PED	Jan 17, 2019				
	6096342	Nov 21, 2011	DP			
	6096342*PED	May 21, 2012				
	6165513	Jun 10, 2018	DP			
	6165513*PED	Dec 10, 2018				
	6432932	Jul 17, 2018	U-595			
	6432932*PED	Jan 17, 2019				
	6465443	Aug 14, 2018	DP			
	6465443*PED	Feb 14, 2019				
<u>CANDESARTAN CILEXETIL - ATACAND</u>						
020838 001	5196444	Jun 04, 2012	DS DP U-660			
	5196444	Jun 04, 2012	DS DP U-3			
	5196444*PED	Dec 04, 2012				
	5534534	Jul 09, 2013	DP			
	5534534*PED	Jan 09, 2014				
	5705517	Apr 18, 2011	DS DP U-660			
	5705517*PED	Oct 18, 2011				
	7538133	Apr 18, 2011	DS			
	7538133*PED	Oct 18, 2011				
<u>CANDESARTAN CILEXETIL - ATACAND</u>						
020838 002	5196444	Jun 04, 2012	DS DP U-660			
	5196444	Jun 04, 2012	DS DP U-3			
	5196444*PED	Dec 04, 2012				
	5534534	Jul 09, 2013	DP			
	5534534*PED	Jan 09, 2014				
	5705517	Apr 18, 2011	DS DP U-660			
	5705517*PED	Oct 18, 2011				
	7538133	Apr 18, 2011	DS			
	7538133*PED	Oct 18, 2011				

PATENT & EXCLUSIVITY DRUG PRODUCT LIST - CUMULATIVE SUPPLEMENT 8 - August 2009

See List footnote for information regarding List content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	PATENT DELIST REQUESTED	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>CANDESARTAN CILEXETIL - ATACAND</u>						
020838 003	5196444	Jun 04, 2012	DS DP U-660			
	5196444	Jun 04, 2012	DS DP U-3			
	5196444*PED	Dec 04, 2012				
	5534534	Jul 09, 2013	DP			
	5534534*PED	Jan 09, 2014				
	5705517	Apr 18, 2011	DS DP U-660			
	5705517*PED	Oct 18, 2011				
	7538133	Apr 18, 2011	DS			
	7538133*PED	Oct 18, 2011				
<u>CANDESARTAN CILEXETIL - ATACAND</u>						
020838 004	5196444	Jun 04, 2012	DS DP U-660			
	5196444	Jun 04, 2012	DS DP U-3			
	5196444*PED	Dec 04, 2012				
	5534534	Jul 09, 2013	DP			
	5534534*PED	Jan 09, 2014				
	5705517	Apr 18, 2011	DS DP U-660			
	5705517*PED	Oct 18, 2011				
	7538133	Apr 18, 2011	DS			
	7538133*PED	Oct 18, 2011				
<u>CANDESARTAN CILEXETIL; HYDROCHLOROTHIAZIDE - ATACAND HCT</u>						
021093 001	5196444	Jun 04, 2012	DS DP U-3			
	5196444*PED	Dec 04, 2012				
	5534534	Jul 09, 2013	DP			
	5534534*PED	Jan 09, 2014				
	5705517	Apr 18, 2011	DS DP U-3			
	5705517*PED	Oct 18, 2011				
	7538133	Apr 18, 2011	DS			
	7538133*PED	Oct 18, 2011				
<u>CANDESARTAN CILEXETIL; HYDROCHLOROTHIAZIDE - ATACAND HCT</u>						
021093 002	5196444	Jun 04, 2012	DS DP U-3			
	5196444*PED	Dec 04, 2012				
	5534534	Jul 09, 2013	DP			
	5534534*PED	Jan 09, 2014				
	5705517	Apr 18, 2011	DS DP U-3			
	5705517*PED	Oct 18, 2011				
	7538133	Apr 18, 2011	DS			
	7538133*PED	Oct 18, 2011				
<u>CANDESARTAN CILEXETIL; HYDROCHLOROTHIAZIDE - ATACAND HCT</u>						
021093 003	5196444	Jun 04, 2012	DS DP U-3			
	5196444*PED	Dec 04, 2012				
	5534534	Jul 09, 2013	DP			
	5534534*PED	Jan 09, 2014				
	5705517	Apr 18, 2011	DS DP U-3			
	5705517*PED	Oct 18, 2011				
	7538133	Apr 18, 2011	DS			
	7538133*PED	Oct 18, 2011				

PATENT & EXCLUSIVITY DRUG PRODUCT LIST - CUMULATIVE SUPPLEMENT 8 - August 2009

See List footnote for information regarding List content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	PATENT DELIST REQUESTED	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>CICLESONIDE - ALVESCO</u>						
021658 002	>A> 5482934	Oct 24, 2017	DS DP U-1002		NDF	Jan 10, 2011
	>A> 5695743	Jul 06, 2010	DP U-1001		NCE	Oct 20, 2011
<u>CICLESONIDE - ALVESCO</u>						
021658 003	>A> 5482934	Oct 24, 2017	DS DP U-1002		NDF	Jan 10, 2011
	>A> 5695743	Jul 06, 2010	DP U-1001		NCE	Oct 20, 2011
<u>CICLESONIDE - OMNARIS</u>						
022004 001	5482934	Oct 24, 2017	DS DP U-557			
<u>CIPROFLOXACIN HYDROCHLORIDE - CETRAXAL</u>						
021918 001					NDF	May 01, 2012
<u>CIPROFLOXACIN HYDROCHLORIDE; HYDROCORTISONE - CIPRO HC</u>						
020805 001	5843930	Jun 06, 2015	U-646			
<u>CLARITHROMYCIN - BIAVIN XL</u>						
050775 001	6551616	Jul 15, 2017	U-924			
<u>CLOBETASOL PROPIONATE - OLUX</u>						
021142 001	6126920	Mar 01, 2016	U-484			
<u>CLOPIDOGREL BISULFATE - PLAVIX</u>						
020839 002	4847265	Nov 17, 2011	DS DP			
	6429210	Jun 10, 2019	DS DP			
	6504030	Jun 10, 2019	DS			
<u>COLCHICINE - COLCRYS</u>						
022352 001					I-603 >A> ODE	Jul 30, 2012 Jul 29, 2016
<u>COLESEVELAM HYDROCHLORIDE - WELCHOL</u>						
021141 001	5607669	Jun 10, 2014	U-323			
	5607669*PED	Dec 10, 2014				
	5679717	Apr 29, 2014	U-323			
	5679717*PED	Oct 29, 2014				
	5693675	Dec 02, 2014				
	5693675*PED	Jun 02, 2015				
	5917007	Apr 29, 2014	U-323			
	5917007*PED	Oct 29, 2014				
	5919832	Jun 10, 2014				
	5919832*PED	Dec 10, 2014				
	6066678	Jun 10, 2014	U-323			
	6066678*PED	Dec 10, 2014				
	6433026	Jun 10, 2014				
	6433026*PED	Dec 10, 2014				

PATENT & EXCLUSIVITY DRUG PRODUCT LIST - CUMULATIVE SUPPLEMENT 8 - August 2009

See List footnote for information regarding List content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	PATENT DELIST REQUESTED	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>COLESEVELAM HYDROCHLORIDE - WELCHOL</u>						
021176 001	5607669	Jun 10, 2014	U-323		I-553	Jan 18, 2011
	5607669*PED	Dec 10, 2014			PED	Jul 18, 2011
	5679717	Apr 29, 2014	U-323			
	5679717*PED	Oct 29, 2014				
	5693675	Dec 02, 2014	DS			
	5693675*PED	Jun 02, 2015				
	5917007	Apr 29, 2014	DS U-323			
	5917007*PED	Oct 29, 2014				
	5919832	Apr 29, 2014	DS			
	5919832*PED	Oct 29, 2014				
	6066678	Apr 29, 2014	DS U-323			
	6066678*PED	Oct 29, 2014				
	6433026	Apr 29, 2014	DS			
	6433026*PED	Oct 29, 2014				
	6784254	Apr 29, 2014	DS DP			
	6784254*PED	Oct 29, 2014				
	7101960	Apr 29, 2014	DS DP U-757			
	7101960*PED	Oct 29, 2014				
	7229613	Apr 17, 2022	U-851			
	7229613*PED	Oct 17, 2022				
<u>CYCLOBENZAPRINE HYDROCHLORIDE - AMRIX</u>						
021777 001	7544372	Nov 14, 2023	U-979			
<u>CYCLOBENZAPRINE HYDROCHLORIDE - AMRIX</u>						
021777 002	7544372	Nov 14, 2023	U-979			
<u>CYCLOSPORINE - SANDIMMUNE</u>						
050625 001	7511014	Feb 16, 2010	DP			
<u>CYCLOSPORINE - SANDIMMUNE</u>						
050625 002	7511014	Feb 16, 2010	DP			
<u>CYCLOSPORINE - SANDIMMUNE</u>						
050625 003	7511014	Feb 16, 2010	DP			
<u>DASATINIB - SPRYCEL</u>						
021986 001	7491725	Oct 13, 2025	DS DP		D-120	May 21, 2012
<u>DASATINIB - SPRYCEL</u>						
021986 002	7491725	Oct 13, 2025	DS DP		D-120	May 21, 2012
<u>DASATINIB - SPRYCEL</u>						
021986 003	7491725	Oct 13, 2025	DS DP		D-120	May 21, 2012
<u>DASATINIB - SPRYCEL</u>						
021986 004	6596746	Jun 28, 2020	DS DP U-780		D-120	May 21, 2012
	6596746	Jun 28, 2020	DS DP U-748			
	7125875	Apr 13, 2020	U-780			
	7125875	Apr 13, 2020	U-779			
	7153856	Apr 28, 2020	U-780			
	7491725	Oct 13, 2025	DS DP			
<u>DEGARELIX ACETATE - FIRMAGON</u>						
022201 001	5925730	Apr 11, 2017	DS DP U-943			

PATENT & EXCLUSIVITY DRUG PRODUCT LIST - CUMULATIVE SUPPLEMENT 8 - August 2009

See List footnote for information regarding List content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	PATENT DELIST REQUESTED	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>DEGARELIX ACETATE - FIRMAGON</u>						
022201 002	5925730	Apr 11, 2017	DS DP U-943			
<u>DESONIDE - DESONATE</u>						
021844 001					NDF	Oct 20, 2009
<u>DEXAMETHASONE - OZURDEX</u>						
022315 001	6726918	Oct 20, 2020	DP U-985		NDF	Jun 17, 2012
	6899717	Nov 01, 2023	U-985			
	7033605	Oct 20, 2020	DP			
<u>DEXAMETHASONE; TOBRAMYCIN - TOBRADEX ST</u>						
050818 001	5149694	Sep 22, 2009	U-953			
<u>DEXLANSOPRAZOLE - KAPIDEX</u>						
022287 001	5045321	Sep 03, 2008	DP		NP	Jan 30, 2012
	5045321*PED	Mar 03, 2009			PED	Jul 30, 2012
	5093132	Sep 03, 2008	DP U-951			
	5093132	Sep 03, 2008	DP U-949			
	5093132	Sep 03, 2008	DP U-950			
	5093132*PED	Mar 03, 2009				
	5433959	Sep 03, 2008	DP U-951			
	5433959	Sep 03, 2008	DP U-950			
	5433959	Sep 03, 2008	DP U-949			
	5433959*PED	Mar 03, 2009				
	6462058	Jun 15, 2020	DS DP U-949			
	6462058	Jun 15, 2020	DS DP U-951			
	6462058	Jun 15, 2020	DS DP U-950			
	6462058*PED	Dec 15, 2020				
	6664276	Jun 15, 2020	DS DP U-950			
	6664276	Jun 15, 2020	DS DP U-949			
	6664276	Jun 15, 2020	DS DP U-951			
	6664276*PED	Dec 15, 2020				
	6939971	Jun 15, 2020	U-951			
	6939971	Jun 15, 2020	U-950			
	6939971	Jun 15, 2020	U-949			
	6939971*PED	Dec 15, 2020				
	7285668	Jun 15, 2020	DS			
	7285668*PED	Dec 15, 2020				

PATENT & EXCLUSIVITY DRUG PRODUCT LIST - CUMULATIVE SUPPLEMENT 8 - August 2009

See List footnote for information regarding List content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	PATENT DELIST REQUESTED	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>DEXLANSOPRAZOLE - KAPIDEX</u>						
022287 002	5045321	Sep 03, 2008	DP		NP	Jan 30, 2012
	5045321*PED	Mar 03, 2009			PED	Jul 30, 2012
	5093132	Sep 03, 2008	DP U-949			
	5093132	Sep 03, 2008	DP U-950			
	5093132	Sep 03, 2008	DP U-951			
	5093132*PED	Mar 03, 2009				
	5433959	Sep 03, 2008	DP U-949			
	5433959	Sep 03, 2008	DP U-950			
	5433959	Sep 03, 2008	DP U-951			
	5433959*PED	Mar 03, 2009				
	6462058	Jun 15, 2020	DS DP U-951			
	6462058	Jun 15, 2020	DS DP U-950			
	6462058	Jun 15, 2020	DS DP U-949			
	6462058*PED	Dec 15, 2020				
	6664276	Jun 15, 2020	DS DP U-949			
	6664276	Jun 15, 2020	DS DP U-950			
	6664276	Jun 15, 2020	DS DP U-951			
	6664276*PED	Dec 15, 2020				
	6939971	Jun 15, 2020	U-949			
	6939971	Jun 15, 2020	U-950			
	6939971	Jun 15, 2020	U-951			
	6939971*PED	Dec 15, 2020				
	7285668	Jun 15, 2020	DS			
	7285668*PED	Dec 15, 2020				
<u>DICLOFENAC POTASSIUM - CAMBIA</u>						
022165 001	6974595	May 15, 2017	U-436			
<u>DICLOFENAC POTASSIUM - ZIPSOR</u>						
022202 001	6365180	Jul 16, 2019	DP U-980		NDF	Jun 16, 2012
<u>DILTIAZEM HYDROCHLORIDE - CARDIZEM CD</u>						
020062 005	5286497	May 20, 2011	DP			
	5439689	Aug 08, 2012	DP U-107			
	5470584	May 20, 2011	DP			
<u>DINOPROSTONE - CERVIDIL</u>						
020411 001	5269321	Jul 14, 2012	DP U-110			
<u>DIVALPROEX SODIUM - DIVALPROEX SODIUM</u>						
077567 002					PC	Aug 01, 2009
<u>DOXERCALCIFEROL - HECTOROL</u>						
020862 001	5602116	Feb 11, 2014	U-987			
	5602116	Feb 11, 2014	U-278			
	6903083	Jul 18, 2021	DS DP	Y		
<u>DOXERCALCIFEROL - HECTOROL</u>						
020862 002	5602116	Feb 11, 2014	U-987			
	5602116	Feb 11, 2014	U-278			
	6903083	Jul 18, 2021	DS DP	Y		
<u>DOXERCALCIFEROL - HECTOROL</u>						
020862 003	5602116	Feb 11, 2014	U-987			

PATENT & EXCLUSIVITY DRUG PRODUCT LIST - CUMULATIVE SUPPLEMENT 8 - August 2009

See List footnote for information regarding List content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	PATENT DELIST REQUESTED	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>DOXERCALCIFEROL - HECTOROL</u>						
021027 001	5707980	Aug 17, 2010	U-321	Y		
	6903083	Jul 18, 2021	DS DP	Y		
<u>DRONEDARONE HYDROCHLORIDE - MULTAQ</u>						
022425 001	5223510	Jul 26, 2011	DS DP U-992		NCE	Jul 01, 2014
	7323493	Jun 19, 2018	DP			
<u>ELTROMBOPAG OLAMINE - PROMACTA</u>						
022291 001	>A> 7452874	May 24, 2021	DS DP			
	>A> 7473686	May 24, 2021	DS DP U-930			
	7547719	Mar 04, 2024	DS DP U-930			
<u>ELTROMBOPAG OLAMINE - PROMACTA</u>						
022291 002	>A> 7452874	May 24, 2021	DS DP			
	>A> 7473686	May 24, 2021	DS DP U-930			
	7547719	Mar 04, 2024	DS DP U-930			
<u>EPINEPHRINE - EPIPEN</u>						
019430 001	7449012	Sep 11, 2025	DP			
<u>EPINEPHRINE - EPIPEN JR.</u>						
019430 002	7449012	Sep 11, 2025	DP			
<u>ESCITALOPRAM OXALATE - LEXAPRO</u>						
021323 001					NPP	Mar 19, 2012
<u>ESCITALOPRAM OXALATE - LEXAPRO</u>						
021323 002					NPP	Mar 19, 2012
<u>ESCITALOPRAM OXALATE - LEXAPRO</u>						
021323 003					NPP	Mar 19, 2012
<u>ESCITALOPRAM OXALATE - LEXAPRO</u>						
021365 001					NPP	Mar 19, 2012
<u>ESOMEPRAZOLE MAGNESIUM - NEXIUM</u>						
021153 001					NPP PED	Apr 28, 2009 Oct 28, 2009
<u>ESOMEPRAZOLE MAGNESIUM - NEXIUM</u>						
021153 002					NPP PED	Apr 28, 2009 Oct 28, 2009
<u>ESOMEPRAZOLE MAGNESIUM - NEXIUM</u>						
021957 001					M-86 NPP PED PED	Jun 18, 2012 Apr 28, 2009 Dec 18, 2012 Oct 28, 2009
<u>ESOMEPRAZOLE MAGNESIUM - NEXIUM</u>						
021957 002					M-86 NPP PED PED	Jun 18, 2012 Apr 28, 2009 Dec 18, 2012 Oct 28, 2009
<u>ESOMEPRAZOLE MAGNESIUM - NEXIUM</u>						
022101 001					NPP PED	Feb 27, 2011 Aug 27, 2011
<u>ESTRADIOL - ELESTRIN</u>						
021813 001	7470433	Aug 03, 2021	DP			

PATENT & EXCLUSIVITY DRUG PRODUCT LIST - CUMULATIVE SUPPLEMENT 8 - August 2009

See List footnote for information regarding List content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	PATENT DELIST REQUESTED	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>ESTRADIOL - EVAMIST</u>						
022014 001	6818226	Feb 19, 2017	DP U-889			
	6818226	Feb 19, 2017	DP U-888			
	6923983	Feb 19, 2017	DP U-889			
	6923983	Feb 19, 2017	DP U-888			
	6978945	Nov 30, 2021	DP			
<u>ETHINYL ESTRADIOL; NORGESTIMATE - TRI LO SPRINTec</u>						
076784 001					PC	Dec 29, 2009
<u>EVEROLIMUS - AFINITOR</u>						
022334 001	5665772	Sep 09, 2014	DS DP		NCE	Mar 30, 2014
	6004973	Jul 12, 2016	DP			
	7297703	Dec 06, 2019	DP			
<u>EVEROLIMUS - AFINITOR</u>						
022334 002	5665772	Sep 09, 2014	DS DP		NCE	Mar 30, 2014
	6004973	Jul 12, 2016	DP			
	7297703	Dec 06, 2019	DP			
<u>EXENATIDE SYNTHETIC - BYETTA</u>						
021773 001	7521423	Oct 15, 2017	DP			
<u>EXENATIDE SYNTHETIC - BYETTA</u>						
021773 002	7521423	Oct 15, 2017	DP			
<u>FEBUXOSTAT - ULORIC</u>						
021856 001	5614520	Mar 25, 2014	DS DP U-954		NCE	Feb 13, 2014
	6225474	Jun 18, 2019	DS			
	7361676	Mar 08, 2024	DP			
<u>FEBUXOSTAT - ULORIC</u>						
021856 002	5614520	Mar 25, 2014	DS DP U-954		NCE	Feb 13, 2014
	6225474	Jun 18, 2019	DS			
	7361676	Mar 08, 2024	DP			
<u>FENOFIBRIC ACID - FIBRICOR</u>						
022418 001	>A> 7569612	Aug 20, 2027	U-1000			
<u>FENOFIBRIC ACID - FIBRICOR</u>						
022418 002	>A> 7569612	Aug 20, 2027	U-1000			
<u>FENTANYL CITRATE - ONSOLIS</u>						
022266 001	>A> 6159498	Oct 18, 2016	DP		NP	Jul 16, 2012
<u>FENTANYL CITRATE - ONSOLIS</u>						
022266 002	>A> 6159498	Oct 18, 2016	DP		NP	Jul 16, 2012
<u>FENTANYL CITRATE - ONSOLIS</u>						
022266 003	>A> 6159498	Oct 18, 2016	DP		NP	Jul 16, 2012
<u>FENTANYL CITRATE - ONSOLIS</u>						
022266 004	>A> 6159498	Oct 18, 2016	DP		NP	Jul 16, 2012
<u>FENTANYL CITRATE - ONSOLIS</u>						
022266 005	>A> 6159498	Oct 18, 2016	DP		NP	Jul 16, 2012

PATENT & EXCLUSIVITY DRUG PRODUCT LIST - CUMULATIVE SUPPLEMENT 8 - August 2009

See List footnote for information regarding List content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	PATENT DELIST REQUESTED	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>FERUMOXYTOL - FERAHEME</u>						
022180 001	6599498	Mar 08, 2020	DS DP		NP	Jun 30, 2012
	7553479	Mar 08, 2020	DS DP			
<u>FLUDARABINE PHOSPHATE - OFORTA</u>						
022273 001	7148207	Dec 20, 2022	DP U-944		NDF	Dec 18, 2011
<u>FLUOXETINE HYDROCHLORIDE - PROZAC</u>						
018936 001	6960577	Nov 01, 2017	U-963		I-589	Mar 19, 2012
<u>FLUOXETINE HYDROCHLORIDE - PROZAC</u>						
018936 003	6960577	Nov 01, 2017	U-963		I-589	Mar 19, 2012
<u>FLUOXETINE HYDROCHLORIDE - PROZAC</u>						
018936 006	6960577	Nov 01, 2017	U-963		I-589	Mar 19, 2012
<u>FLUOXETINE HYDROCHLORIDE; OLANZAPINE - SYMBYAX</u>						
021520 001	6960577	Nov 01, 2017	U-962		I-593	Mar 19, 2012
<u>FLUOXETINE HYDROCHLORIDE; OLANZAPINE - SYMBYAX</u>						
021520 002	6960577	Nov 01, 2017	U-962		I-593	Mar 19, 2012
<u>FLUOXETINE HYDROCHLORIDE; OLANZAPINE - SYMBYAX</u>						
021520 003	6960577	Nov 01, 2017	U-962		I-593	Mar 19, 2012
<u>FLUOXETINE HYDROCHLORIDE; OLANZAPINE - SYMBYAX</u>						
021520 004	6960577	Nov 01, 2017	U-962		I-593	Mar 19, 2012
<u>FLUOXETINE HYDROCHLORIDE; OLANZAPINE - SYMBYAX</u>						
021520 005	6960577	Nov 01, 2017	U-962		I-593	Mar 19, 2012
<u>FLUTICASONE FUROATE - VERAMYST</u>						
022051 001	7541350	Aug 03, 2021	DP U-988			
<u>FLUTICASONE PROPIONATE - FLOVENT DISKUS 100</u>						
020833 002	5590645	Mar 01, 2011	DP			
	5590645*PED	Sep 01, 2011				
	5860419	Mar 01, 2011	DP			
	5860419*PED	Sep 01, 2011				
	5873360	Feb 23, 2016	DP			
	5873360*PED	Aug 23, 2016				
	6032666	Mar 01, 2011	DP			
	6032666*PED	Sep 01, 2011				
	6378519	Mar 01, 2011	DP			
	6378519*PED	Sep 01, 2011				
	6536427	Mar 01, 2011	DP			
	6536427*PED	Sep 01, 2011				
	6792945	Mar 01, 2011	DP			
	6792945*PED	Sep 01, 2011				
	7225808	Mar 01, 2011	DP			
	7225808*PED	Sep 01, 2011				
	7389775	Mar 01, 2011	DP			
	7389775*PED	Sep 01, 2011				

PATENT & EXCLUSIVITY DRUG PRODUCT LIST - CUMULATIVE SUPPLEMENT 8 - August 2009

See List footnote for information regarding List content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	PATENT DELIST REQUESTED	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>FLUTICASONE PROPIONATE - FLOVENT DISKUS 250</u>						
020833 003	5590645	Mar 01, 2011	DP			
	5590645*PED	Sep 01, 2011				
	5860419	Mar 01, 2011	DP			
	5860419*PED	Sep 01, 2011				
	5873360	Feb 23, 2016	DP			
	5873360*PED	Aug 23, 2016				
	6032666	Mar 01, 2011	DP			
	6032666*PED	Sep 01, 2011				
	6378519	Mar 01, 2011	DP			
	6378519*PED	Sep 01, 2011				
	6536427	Mar 01, 2011	DP			
	6536427*PED	Sep 01, 2011				
	6792945	Mar 01, 2011	DP			
	6792945*PED	Sep 01, 2011				
	7225808	Mar 01, 2011	DP			
	7225808*PED	Sep 01, 2011				
	7389775	Mar 01, 2011	DP			
	7389775*PED	Sep 01, 2011				
<u>FLUTICASONE PROPIONATE - FLOVENT DISKUS 50</u>						
020833 001	5590645	Mar 01, 2011	DP			
	5590645*PED	Sep 01, 2011				
	5860419	Mar 01, 2011	DP			
	5860419*PED	Sep 01, 2011				
	5873360	Feb 23, 2016	DP			
	5873360*PED	Aug 23, 2016				
	6032666	Mar 01, 2011	DP			
	6032666*PED	Sep 01, 2011				
	6378519	Mar 01, 2011	DP			
	6378519*PED	Sep 01, 2011				
	6536427*PED	Sep 01, 2011				
	6792945	Mar 01, 2011	DP			
	6792945*PED	Sep 01, 2011				
	7225808	Mar 01, 2011	DP			
	7225808*PED	Sep 01, 2011				
	7389775	Mar 01, 2011	DP			
	7389775*PED	Sep 01, 2011				

PATENT & EXCLUSIVITY DRUG PRODUCT LIST - CUMULATIVE SUPPLEMENT 8 - August 2009

See List footnote for information regarding List content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	PATENT DELIST REQUESTED	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
FLUTICASONE PROPIONATE - FLOVENT HFA						
021433 001	5658549	Aug 19, 2014	DP U-710			
	5658549*PED	Feb 19, 2015				
	5674472	Oct 07, 2014	DP			
	5674472*PED	Apr 07, 2015				
	6161724	Jan 16, 2018	DP			
	6161724*PED	Jul 16, 2018				
	6170717	Dec 23, 2017	DP			
	6170717*PED	Jun 23, 2018				
	6251368	Dec 04, 2012	DP			
	6251368*PED	Jun 04, 2013				
	6253762	Apr 14, 2015	DP U-582			
	6253762*PED	Oct 14, 2015				
	6315173	Jun 23, 2017	DP			
	6315173*PED	Jun 23, 2018				
	6431168	Jun 08, 2018	DP			
	6431168*PED	Dec 08, 2018				
	6435372	Jan 16, 2018	DP			
	6435372*PED	Jul 16, 2018				
	6510969	Dec 23, 2017	DP			
	6510969*PED	Jun 23, 2018				
	6546928	Apr 14, 2015	DP U-583			
	6546928*PED	Oct 14, 2015				
	6596260	Aug 10, 2014	DP			
	6596260*PED	Feb 10, 2015				
	6743413	Jun 01, 2021	U-581			
	6743413*PED	Dec 01, 2021				
	6938796	Jan 16, 2018	DP			
	6938796*PED	Jul 16, 2018				
	6966467	Dec 23, 2017	DP			
	6966467*PED	Jun 23, 2018				
	6997349	Jan 16, 2018	DP			
	6997349*PED	Jul 16, 2018				
	7107986	Jun 08, 2018	DP			
	7107986*PED	Dec 08, 2019				
	7143908	Jan 16, 2018	DP			
	7143908*PED	Jul 16, 2018				
	7350676	Aug 24, 2018	DP			
	7350676*PED	Feb 24, 2019				
	7500444	Jan 04, 2025	DP			
	7500444*PED	Jul 04, 2025				

PATENT & EXCLUSIVITY DRUG PRODUCT LIST - CUMULATIVE SUPPLEMENT 8 - August 2009

See List footnote for information regarding List content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	PATENT DELIST REQUESTED	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
FLUTICASONE PROPIONATE - FLOVENT HFA						
021433 002	5658549	Aug 19, 2014	DP U-710			
	5658549*PED	Feb 19, 2015				
	5674472	Oct 07, 2014	DP			
	5674472*PED	Apr 07, 2015				
	6161724	Jan 16, 2018	DP			
	6161724*PED	Jul 16, 2018				
	6170717	Dec 23, 2017	DP			
	6170717*PED	Jun 23, 2018				
	6251368	Dec 04, 2012	DP			
	6251368*PED	Jun 04, 2013				
	6253762	Apr 14, 2015	DP U-582			
	6253762*PED	Oct 14, 2015				
	6315173	Dec 23, 2017	DP			
	6315173*PED	Jun 23, 2018				
	6431168	Jun 08, 2018	DP			
	6431168*PED	Dec 08, 2018				
	6435372	Jan 16, 2018	DP			
	6435372*PED	Jul 16, 2018				
	6510969	Dec 23, 2017	DP			
	6510969*PED	Jun 23, 2018				
	6546928	Apr 14, 2015	DP U-583			
	6546928*PED	Oct 14, 2015				
	6596260	Aug 10, 2014	DP			
	6596260*PED	Feb 10, 2015				
	6743413	Jun 01, 2021	U-581			
	6743413*PED	Dec 01, 2021				
	6938796	Jan 16, 2018	DP			
	6938796*PED	Jul 16, 2018				
	6966467	Dec 23, 2017	DP			
	6966467*PED	Jun 23, 2018				
	6997349	Jan 16, 2018	DP			
	6997349*PED	Jul 16, 2018				
	7107986	Jun 08, 2018	DP			
	7107986*PED	Dec 08, 2019				
	7143908	Jan 16, 2018	DP			
	7143908*PED	Jul 16, 2018				
	7350676	Aug 24, 2018	DP			
	7350676*PED	Feb 24, 2019				
	7500444	Jan 04, 2025	DP			
	7500444*PED	Jul 04, 2025				

PATENT & EXCLUSIVITY DRUG PRODUCT LIST - CUMULATIVE SUPPLEMENT 8 - August 2009

See List footnote for information regarding List content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	PATENT DELIST REQUESTED	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
FLUTICASONE PROPIONATE - FLOVENT HFA						
021433 003	5658549	Aug 19, 2014	DP U-710			
	5658549*PED	Feb 19, 2015	U-710			
	5674472	Oct 07, 2014	DP			
	5674472*PED	Apr 07, 2015				
	6161724	Jan 16, 2018	DP			
	6161724*PED	Jul 16, 2018				
	6170717	Dec 23, 2017	DP			
	6170717*PED	Jun 23, 2018				
	6251368	Dec 04, 2012	DP			
	6251368*PED	Jun 04, 2013				
	6253762	Apr 14, 2015	DP U-582			
	6253762*PED	Oct 14, 2015	U-582			
	6315173	Dec 23, 2017	DP			
	6315173*PED	Jun 23, 2018	DP			
	6431168	Jun 08, 2018	DP			
	6431168*PED	Dec 08, 2018				
	6435372	Jan 16, 2018	DP			
	6435372*PED	Jul 16, 2018				
	6510969	Dec 23, 2017	DP			
	6510969*PED	Jun 23, 2018	DP			
	6546928	Apr 14, 2015	DP U-583			
	6546928*PED	Oct 14, 2015	U-583			
	6596260	Aug 10, 2014	DP			
	6596260*PED	Feb 10, 2015				
	6743413	Jun 01, 2021	U-581			
	6743413*PED	Dec 01, 2021	U-581			
	6938796	Jan 16, 2018	DP			
	6938796*PED	Jul 16, 2018				
	6966467	Dec 23, 2017	DP			
	6966467*PED	Jun 23, 2018				
	6997349	Jan 16, 2018	DP			
	6997349*PED	Jul 16, 2018				
	7107986	Jun 08, 2018	DP			
	7107986*PED	Dec 08, 2019				
	7143908	Jan 16, 2018	DP			
	7143908*PED	Jul 16, 2018				
	7350676	Aug 24, 2018	DP			
	7350676*PED	Feb 24, 2019				
	7500444	Jan 04, 2025	DP			
	7500444*PED	Jul 04, 2025				

PATENT & EXCLUSIVITY DRUG PRODUCT LIST - CUMULATIVE SUPPLEMENT 8 - August 2009

See List footnote for information regarding List content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	PATENT DELIST REQUESTED	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>FLUTICASONE PROPIONATE; SALMETEROL XINAFOATE - ADVAIR DISKUS 100/50</u>						
021077 001	5590645	Mar 01, 2011	DP		M-84	Mar 31, 2012
	5590645*PED	Sep 01, 2011				
	5860419	Mar 01, 2011	DP			
	5860419*PED	Sep 01, 2011				
	5873360	Feb 23, 2016	DP			
	5873360*PED	Aug 23, 2016				
	6032666	Mar 01, 2011	DP			
	6032666*PED	Sep 01, 2011				
	6378519	Mar 01, 2011	DP			
	6378519*PED	Sep 01, 2011				
	6792945	Mar 01, 2011	DP			
	6792945*PED	Sep 01, 2011				
	7225808	Mar 01, 2011	DP			
	7225808*PED	Sep 01, 2011				
	7389775	Mar 01, 2011	DP			
	7389775*PED	Sep 01, 2011				
<u>FLUTICASONE PROPIONATE; SALMETEROL XINAFOATE - ADVAIR DISKUS 250/50</u>						
021077 002	5590645	Mar 01, 2011	DP		M-84	Mar 31, 2012
	5590645*PED	Sep 01, 2011				
	5860419	Mar 01, 2011	DP			
	5860419*PED	Sep 01, 2011				
	5873360	Feb 23, 2016	DP			
	5873360*PED	Aug 23, 2016				
	6032666	Mar 01, 2011	DP			
	6032666*PED	Sep 01, 2011				
	6378519	Mar 01, 2011	DP			
	6378519*PED	Sep 01, 2011				
	6792945	Mar 01, 2011	DP			
	6792945*PED	Sep 01, 2011				
	7225808	Mar 01, 2011	DP			
	7225808*PED	Sep 01, 2011				
	7389775	Mar 01, 2011	DP			
	7389775*PED	Sep 01, 2011				

PATENT & EXCLUSIVITY DRUG PRODUCT LIST - CUMULATIVE SUPPLEMENT 8 - August 2009

See List footnote for information regarding List content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	PATENT DELIST REQUESTED	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>FLUTICASONE PROPIONATE; SALMETEROL XINAFOATE - ADVAIR DISKUS 500/50</u>						
021077 003	5590645	Mar 01, 2011	DP		M-84	Mar 31, 2012
	5590645*PED	Sep 01, 2011				
	5860419	Mar 01, 2011	DP			
	5860419*PED	Sep 01, 2011				
	5873360	Feb 23, 2016	DP			
	5873360*PED	Aug 23, 2016				
	6032666	Mar 01, 2011	DP			
	6032666*PED	Sep 01, 2011				
	6378519	Mar 01, 2011	DP			
	6378519*PED	Sep 01, 2011				
	6792945	Mar 01, 2011	DP			
	6792945*PED	Sep 01, 2011				
	7225808	Mar 01, 2011	DP			
	7225808*PED	Sep 01, 2011				
	7389775	Mar 01, 2011	DP			
	7389775*PED	Sep 01, 2011				

PATENT & EXCLUSIVITY DRUG PRODUCT LIST - CUMULATIVE SUPPLEMENT 8 - August 2009

See List footnote for information regarding List content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	PATENT DELIST REQUESTED	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
FLUTICASONE PROPIONATE; SALMETEROL XINAFOATE - ADVAIR HFA						
021254 001	5658549	Aug 19, 2014	DP		U-738	
	5658549*PED	Feb 19, 2015			U-738	
	5674472	Oct 07, 2014	DP			
	5674472*PED	Apr 07, 2015				
	6161724	Jan 16, 2018	DP			
	6161724*PED	Jul 16, 2018				
	6170717	Dec 23, 2017	DP			
	6170717*PED	Jun 23, 2018				
	6251368	Dec 04, 2012	DP			
	6251368*PED	Jun 04, 2013				
	6253762	Apr 14, 2015	DP		U-738	
	6253762*PED	Oct 14, 2015			U-738	
	6315173	Dec 23, 2017	DP			
	6315173*PED	Jun 23, 2018				
	6431168	Jun 08, 2018	DP			
	6431168*PED	Dec 08, 2018				
	6435372	Jan 16, 2018	DP			
	6435372*PED	Jul 16, 2018				
	6510969	Dec 23, 2017				
	6510969*PED	Jun 23, 2018				
	6546928	Apr 14, 2015	DP			
	6546928*PED	Oct 14, 2015				
	6596260	Aug 10, 2014	DP			
	6596260*PED	Feb 10, 2015				
	6743413	Jun 01, 2021			U-841	
	6743413*PED	Dec 01, 2021			U-841	
	6938796	Jan 16, 2018	DP			
	6938796*PED	Jul 16, 2018				
	6966467	Dec 23, 2017	DP			
	6966467*PED	Jun 23, 2018				
	6997349	Jan 16, 2018	DP			
	6997349*PED	Jul 16, 2018				
	7107986	Jun 08, 2018	DP			
	7107986*PED	Dec 08, 2018				
	7143908	Jan 16, 2018	DP			
	7143908*PED	Jul 16, 2018				
	7350676	Aug 24, 2018	DP			
	7350676*PED	Feb 24, 2019				
	7500444	Jan 04, 2025	DP			
	7500444*PED	Jul 04, 2025				

PATENT & EXCLUSIVITY DRUG PRODUCT LIST - CUMULATIVE SUPPLEMENT 8 - August 2009

See List footnote for information regarding List content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	PATENT DELIST REQUESTED	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
FLUTICASONE PROPIONATE; SALMETEROL XINAFOATE - ADVAIR HFA						
021254 002	5658549	Aug 19, 2014	DP U-738			
	5658549*PED	Feb 19, 2015	DP U-738			
	5674472	Oct 07, 2014	DP			
	5674472*PED	Apr 07, 2015	DP			
	6161724	Jan 16, 2018	DP			
	6161724*PED	Jul 16, 2018				
	6170717	Dec 23, 2017	DP			
	6170717*PED	Jun 23, 2018				
	6251368	Dec 04, 2012	DP			
	6251368*PED	Jun 04, 2013	DP			
	6253762	Apr 14, 2015	DP U-738			
	6253762*PED	Oct 14, 2015	DP U-738			
	6315173	Dec 23, 2017	DP			
	6315173*PED	Jun 23, 2018	DP			
	6431168	Jun 08, 2018	DP			
	6431168*PED	Dec 08, 2018				
	6435372	Jan 16, 2018	DP			
	6435372*PED	Jul 16, 2018				
	6510969	Dec 23, 2017	DP			
	6510969*PED	Jun 23, 2018	DP			
	6546928	Apr 14, 2015	DP			
	6546928*PED	Oct 14, 2015	DP			
	6596260	Aug 10, 2014	DP			
	6596260*PED	Feb 10, 2015				
	6743413	Jun 01, 2021	U-841			
	6743413*PED	Dec 01, 2021	U-841			
	6938796	Jan 16, 2018	DP			
	6938796*PED	Jul 16, 2018				
	6966467	Dec 23, 2017	DP			
	6966467*PED	Jun 23, 2018				
	6997349	Jan 16, 2018	DP			
	6997349*PED	Jul 16, 2018				
	7107986	Jun 08, 2018	DP			
	7107986*PED	Dec 08, 2018				
	7143908	Jan 16, 2018	DP			
	7143908*PED	Jul 16, 2018				
	7350676	Aug 24, 2018	DP			
	7350676*PED	Feb 24, 2019				
	7500444	Jan 04, 2025	DP			
	7500444*PED	Jul 04, 2025				

PATENT & EXCLUSIVITY DRUG PRODUCT LIST - CUMULATIVE SUPPLEMENT 8 - August 2009

See List footnote for information regarding List content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	PATENT DELIST REQUESTED	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>FLUTICASONE PROPIONATE; SALMETEROL XINAFOATE - ADVAIR HFA</u>						
021254 003	5658549	Aug 19, 2014	DP U-738			
	5658549*PED	Feb 19, 2015	DP U-738			
	5674472	Oct 07, 2014	DP			
	5674472*PED	Apr 07, 2015	DP			
	6161724	Jan 16, 2018	DP			
	6161724*PED	Jul 16, 2018				
	6170717	Dec 23, 2017	DP			
	6170717*PED	Jun 23, 2018				
	6251368	Dec 04, 2012	DP			
	6251368*PED	Jun 04, 2013	DP			
	6253762	Apr 14, 2015	DP U-738			
	6253762*PED	Oct 14, 2015	DP U-738			
	6315173	Dec 23, 2017	DP			
	6315173*PED	Jun 23, 2018	DP			
	6431168	Jun 08, 2018	DP			
	6431168*PED	Dec 08, 2018				
	6435372	Jan 16, 2018	DP			
	6435372*PED	Jul 16, 2018				
	6510969	Dec 23, 2017	DP			
	6510969*PED	Jun 23, 2018	DP			
	6546928	Apr 14, 2015	DP			
	6546928*PED	Oct 14, 2015	DP			
	6596260	Aug 10, 2014	DP			
	6596260*PED	Feb 10, 2015				
	6743413	Jun 01, 2021	U-841			
	6743413*PED	Dec 01, 2021	U-841			
	6938796	Jan 16, 2018	DP			
	6938796*PED	Jul 16, 2018				
	6966467	Dec 23, 2017	DP			
	6966467*PED	Jun 23, 2018				
	6997349	Jan 16, 2018	DP			
	6997349*PED	Jul 16, 2018				
	7107986	Jun 08, 2018	DP			
	7107986*PED	Dec 08, 2018				
	7143908	Jan 16, 2018	DP			
	7143908*PED	Jul 16, 2018				
	7350676	Aug 24, 2018	DP			
	7350676*PED	Feb 24, 2019				
	7500444	Jan 04, 2025	DP			
	7500444*PED	Jul 04, 2025				
<u>FLUVOXAMINE MALEATE - LUVOX</u>						
021519 001					M-83	Apr 14, 2011
<u>FLUVOXAMINE MALEATE - LUVOX</u>						
021519 002					M-83	Apr 14, 2011
<u>FLUVOXAMINE MALEATE - LUVOX</u>						
021519 003					M-83	Apr 14, 2011
<u>FOLLITROPIN ALFA/BETA - FOLLISTIM AQ</u>						
021211 001	>A> 7563763	Aug 23, 2019	U-993			

PATENT & EXCLUSIVITY DRUG PRODUCT LIST - CUMULATIVE SUPPLEMENT 8 - August 2009

See List footnote for information regarding List content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	PATENT DELIST REQUESTED	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>FOLLITROPIN ALFA/BETA - FOLLISTIM AQ</u>						
021211 002	>A> 7563763	Aug 23, 2019	U-993			
<u>FOLLITROPIN ALFA/BETA - FOLLISTIM AQ</u>						
021211 003	>A> 7563763	Aug 23, 2019	U-993			
<u>FOLLITROPIN ALFA/BETA - FOLLISTIM AQ</u>						
021211 004	>A> 7563763	Aug 23, 2019	U-993			
<u>FOLLITROPIN ALFA/BETA - GONAL-F</u>						
020378 004	7563763	Aug 23, 2019	DP			
<u>FOLLITROPIN ALFA/BETA - GONAL-F</u>						
020378 005	7563763	Aug 23, 2019	DP			
<u>FOLLITROPIN ALFA/BETA - GONAL-F RFF PEN</u>						
021684 001	7446090	Aug 23, 2019	DP			
<u>FOLLITROPIN ALFA/BETA - GONAL-F RFF PEN</u>						
021684 002	7446090	Aug 23, 2019	DP			
<u>FOLLITROPIN ALFA/BETA - GONAL-F RFF PEN</u>						
021684 003	7446090	Aug 23, 2019	DP			
<u>FOMEPIZOLE - ANTIZOL</u>						
020696 001	>A> 7553863	Jun 30, 2027	DS DP			
<u>FOSPROPOFOL DISODIUM - LUSEDRA</u>						
022244 001	6204257	Jun 07, 2018	DS DP U-945		NCE	Dec 12, 2013
<u>GADODIAMIDE - OMNISCAN</u>						
022066 002	5362475	Nov 08, 2011	DS			
<u>GATIFLOXACIN - ZYMAR</u>						
021493 001	4980470	Dec 15, 2009	DS DP			
	4980470*PED	Jun 15, 2010				
	5880283	Dec 05, 2015				
	5880283*PED	Jun 05, 2016				
	6333045	Aug 20, 2019	DP			
	6333045*PED	Feb 20, 2020				
<u>GLATIRAMER ACETATE - COPAXONE</u>						
020622 001					I-594	Feb 27, 2012
<u>GLATIRAMER ACETATE - COPAXONE</u>						
020622 002					I-594	Feb 27, 2012
<u>GLIMEPIRIDE; PIOGLITAZONE HYDROCHLORIDE - DUETACT</u>						
021925 001	7538125	Jun 19, 2016	DP			
<u>GLIMEPIRIDE; PIOGLITAZONE HYDROCHLORIDE - DUETACT</u>						
021925 002	7538125	Jun 19, 2016	DP			
<u>GOSERELIN ACETATE - ZOLADEX</u>						
019726 001	7500964	Feb 26, 2021	DP			
<u>GOSERELIN ACETATE - ZOLADEX</u>						
020578 001	7500964	Feb 26, 2021	DP			

PATENT & EXCLUSIVITY DRUG PRODUCT LIST - CUMULATIVE SUPPLEMENT 8 - August 2009

See List footnote for information regarding List content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	PATENT DELIST REQUESTED	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>HYDROCHLOROTHIAZIDE; TELMISARTAN - MICARDIS HCT</u>						
021162 003	5591762	Jan 07, 2014	DS DP U-3			
<u>IBUPROFEN - CALDOLOR</u>						
022348 001	6727286	Nov 27, 2021	DP U-981		NP	Jun 11, 2012
<u>IBUPROFEN - CALDOLOR</u>						
022348 002	6727286	Nov 27, 2021	DP U-981		NP	Jun 11, 2012
<u>ILOPERIDONE - FANAPT</u>						
022192 001	RE39198	Nov 15, 2011	DS DP U-971		NCE	May 06, 2014
<u>ILOPERIDONE - FANAPT</u>						
022192 002	RE39198	Nov 15, 2011	DS DP U-971		NCE	May 06, 2014
<u>ILOPERIDONE - FANAPT</u>						
022192 003	RE39198	Nov 15, 2011	DS DP U-971		NCE	May 06, 2014
<u>ILOPERIDONE - FANAPT</u>						
022192 004	RE39198	Nov 15, 2011	DS DP U-971		NCE	May 06, 2014
<u>ILOPERIDONE - FANAPT</u>						
022192 005	RE39198	Nov 15, 2011	DS DP U-971		NCE	May 06, 2014
<u>ILOPERIDONE - FANAPT</u>						
022192 006	RE39198	Nov 15, 2011	DS DP U-971		NCE	May 06, 2014
<u>ILOPERIDONE - FANAPT</u>						
022192 007	RE39198	Nov 15, 2011	DS DP U-971		NCE	May 06, 2014
<u>IMATINIB MESYLATE - GLEEVEC</u>						
021335 002	>A> 5521184	Jan 04, 2015	DS DP U-649			
<u>IMATINIB MESYLATE - GLEEVEC</u>						
021588 001	>A> 7544799	Jan 16, 2019	DS DP		I-583	Dec 19, 2011
	>A> 7544799*PED	Jul 16, 2019				
<u>IMATINIB MESYLATE - GLEEVEC</u>						
021588 002	>A> 7544799	Jan 16, 2019	DS DP		I-583	Dec 19, 2011
	>A> 7544799*PED	Jul 16, 2019				
<u>INSULIN DETEMIR RECOMBINANT - LEVEMIR</u>						
021536 001	5750497	May 16, 2019	DS DP U-668			
<u>INSULIN GLARGINE RECOMBINANT - LANTUS</u>						
021081 001	5656722	Aug 12, 2014	DS DP U-948			
	5656722*PED	Feb 12, 2015				
	7476652	Jul 23, 2023	DP			
	7476652*PED	Jan 23, 2024				
<u>INSULIN GLULISINE RECOMBINANT - APIDRA</u>						
021629 002					NCE	Apr 16, 2009
<u>INSULIN GLULISINE RECOMBINANT - APIDRA SOLOSTAR</u>						
021629 003					NPP NCE	Oct 24, 2011 Apr 16, 2009
<u>IODIXANOL - VISIPAQUE 270</u>						
020351 001	5366722	Nov 22, 2011	DP			
	RE36418	Jul 12, 2011	DP			

PATENT & EXCLUSIVITY DRUG PRODUCT LIST - CUMULATIVE SUPPLEMENT 8 - August 2009

See List footnote for information regarding List content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	PATENT DELIST REQUESTED	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>IODIXANOL - VISIPAQUE 270</u>						
020808 001	5366722	Nov 22, 2011	DP			
	RE36418	Jul 12, 2011	DP			
<u>IODIXANOL - VISIPAQUE 320</u>						
020351 002	RE36418	Jul 12, 2011	DP			
<u>IODIXANOL - VISIPAQUE 320</u>						
020808 002	RE36418	Jul 12, 2011	DP			
<u>IXABEPILONE - IXEMPRA KIT</u>						
022065 001	6605599	May 26, 2018	DS DP U-961			
	6670384	Jan 23, 2022	DP U-960			
	6670384	Jan 23, 2022	DP U-959			
	7022330	Jan 23, 2022	DP U-958			
	7125899	May 26, 2018	DS DP U-957			
	7312237	Aug 21, 2024	U-965			
<u>IXABEPILONE - IXEMPRA KIT</u>						
022065 002	6605599	May 26, 2018	DS DP U-961			
	6670384	Jan 23, 2022	DP U-960			
	6670384	Jan 23, 2022	DP U-959			
	7022330	Jan 23, 2022	DP U-958			
	7125899	May 26, 2018	DS DP U-957			
	7312237	Aug 21, 2024	U-965			
<u>KETOCONAZOLE - EXTINA</u>						
021738 001	7553835	Oct 19, 2018	DP U-245			
<u>KETOROLAC TROMETHAMINE - ACUVAIL</u>						
022427 001					NP	Jul 22, 2012
<u>LANSOPRAZOLE - PREVACID</u>						
020406 001					M-85 PED	Oct 28, 2011 Apr 28, 2012
<u>LANSOPRAZOLE - PREVACID</u>						
020406 002					M-85 PED	Oct 28, 2011 Apr 28, 2012
<u>LANSOPRAZOLE - PREVACID</u>						
021281 001					M-85 PED	Oct 28, 2011 Apr 28, 2012
<u>LANSOPRAZOLE - PREVACID</u>						
021281 002					M-85 PED	Oct 28, 2011 Apr 28, 2012
<u>LANSOPRAZOLE - PREVACID</u>						
021428 001	7431942	May 17, 2019	DP		M-85 PED	Oct 28, 2011 Apr 28, 2012
	7431942*PED	Nov 17, 2019				
<u>LANSOPRAZOLE - PREVACID</u>						
021428 002	7431942	May 17, 2019	DP		M-85 PED	Oct 28, 2011 Apr 28, 2012
	7431942*PED	Nov 17, 2019				
<u>LANSOPRAZOLE - PREVACID 24 HR</u>						
022327 001					NP	May 18, 2012

PATENT & EXCLUSIVITY DRUG PRODUCT LIST - CUMULATIVE SUPPLEMENT 8 - August 2009

See List footnote for information regarding List content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	PATENT DELIST REQUESTED	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>LANSOPRAZOLE - PREVACID IV</u>						
021566 001	4628098	May 10, 2009	DS			
	4628098*PED	Nov 10, 2009				
	7396841	Aug 17, 2021	DP U-947			
	7396841*PED	Feb 17, 2022				
<u>LANSOPRAZOLE; NAPROXEN - PREVACID NAPRAPAC 500 (COPACKAGED)</u>						
021507 004	4628098	May 10, 2009	DS			
	4628098*PED	Nov 10, 2009				
	5045321	Sep 03, 2008	DP			
	5045321*PED	Mar 03, 2009				
	5093132	Sep 03, 2008	DP			
	5093132*PED	Mar 03, 2009				
	5433959	Sep 03, 2008	DP			
	5433959*PED	Mar 03, 2009				
<u>LANTHANUM CARBONATE - FOSRENOL</u>						
021468 001	5968976	Oct 26, 2018	DP U-613			
<u>LANTHANUM CARBONATE - FOSRENOL</u>						
021468 002	5968976	Oct 26, 2018	DP U-613			
<u>LANTHANUM CARBONATE - FOSRENOL</u>						
021468 003	5968976	Oct 26, 2018	DP U-613			
<u>LANTHANUM CARBONATE - FOSRENOL</u>						
021468 004	5968976	Oct 26, 2018	DP U-613			
<u>LEVETIRACETAM - KEPPRA XR</u>						
022285 002					NDF	Sep 12, 2011
<u>LEVOCETIRIZINE DIHYDROCHLORIDE - XYZAL</u>						
022064 001					>A> NPP	Aug 21, 2012
					>A> PED	Feb 21, 2013
<u>LEVOCETIRIZINE DIHYDROCHLORIDE - XYZAL</u>						
022157 001					>A> NPP	Aug 21, 2012
					>A> PED	Feb 21, 2013
<u>LEVONORGESTREL - PLAN B ONE-STEP</u>						
021998 001					NP	Jul 10, 2012
<u>LIDOCAINE HYDROCHLORIDE - ZINGO</u>						
022114 001					NPP	Jan 08, 2012
<u>MALATHION - OVIDE</u>						
018613 001	7560445	Feb 01, 2027	DS DP U-986			
<u>MARAVIROC - SELZENTRY</u>						
022128 001	>A> 7576097	May 25, 2021	DS			
<u>MARAVIROC - SELZENTRY</u>						
022128 002	>A> 7576097	May 25, 2021	DS			
<u>MEMANTINE HYDROCHLORIDE - NAMENDA</u>						
021487 001	5061703	Apr 11, 2015	U-539			
<u>MEMANTINE HYDROCHLORIDE - NAMENDA</u>						
021487 002	5061703	Apr 11, 2015	U-539			

PATENT & EXCLUSIVITY DRUG PRODUCT LIST - CUMULATIVE SUPPLEMENT 8 - August 2009

See List footnote for information regarding List content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	PATENT DELIST REQUESTED	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>MEMANTINE HYDROCHLORIDE - NAMENDA</u>						
021627 001	5061703	Apr 11, 2015	U-539			
<u>METFORMIN HYDROCHLORIDE; PIOGLITAZONE HYDROCHLORIDE - ACTOPLUS MET XR</u>						
022024 001	4687777	Jan 17, 2011	DS			
	5965584	Jun 19, 2016	DP U-973			
	6099859	Mar 20, 2018	DP			
	6166042	Jun 19, 2016	U-973			
	6166043	Jun 19, 2016	U-973			
	6172090	Jun 19, 2016	U-973			
	6495162	Mar 20, 2018	DP			
	6790459	Mar 17, 2021	U-974			
	6866866	Mar 17, 2021	DP			
<u>METFORMIN HYDROCHLORIDE; PIOGLITAZONE HYDROCHLORIDE - ACTOPLUS MET XR</u>						
022024 002	4687777	Jan 17, 2011	DS			
	5965584	Jun 19, 2016	DP U-973			
	6099859	Mar 20, 2018	DP			
	6166042	Jun 19, 2016	U-973			
	6166043	Jun 19, 2016	U-973			
	6172090	Jun 19, 2016	U-973			
	6495162	Mar 20, 2018	DP			
	6790459	Mar 17, 2021	U-974			
	6866866	Mar 17, 2021	DP			
<u>MICONAZOLE NITRATE - MONISTAT 1 COMBINATION PACK</u>						
021308 001	5514698	Mar 21, 2014		Y		
	6153635	Nov 28, 2020		Y		
<u>MILNACIPRAN HYDROCHLORIDE - SAVELLA</u>						
022256 001	6602911	Nov 05, 2021	U-882		NCE	Jan 14, 2014
	6992110	Nov 05, 2021	U-882			
<u>MILNACIPRAN HYDROCHLORIDE - SAVELLA</u>						
022256 002	6602911	Nov 05, 2021	U-882		NCE	Jan 14, 2014
	6992110	Nov 05, 2021	U-882			
<u>MILNACIPRAN HYDROCHLORIDE - SAVELLA</u>						
022256 003	6602911	Nov 05, 2021	U-882		NCE	Jan 14, 2014
	6992110	Nov 05, 2021	U-882			
<u>MILNACIPRAN HYDROCHLORIDE - SAVELLA</u>						
022256 004	6602911	Nov 05, 2021	U-882		NCE	Jan 14, 2014
	6992110	Nov 05, 2021	U-882			
<u>MINOCYCLINE HYDROCHLORIDE - SOLODYN</u>						
050808 002	7541347	Apr 02, 2027	U-917			
	7544373	Apr 02, 2027	DP			
<u>MINOCYCLINE HYDROCHLORIDE - SOLODYN</u>						
050808 004	5908838	Feb 19, 2018	U-917			
<u>MINOCYCLINE HYDROCHLORIDE - SOLODYN</u>						
050808 005	5908838	Feb 19, 2018	U-917			

PATENT & EXCLUSIVITY DRUG PRODUCT LIST - CUMULATIVE SUPPLEMENT 8 - August 2009

See List footnote for information regarding List content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	PATENT DELIST REQUESTED	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>MOMETASONE FUROATE - ASMANEX TWISTHALER</u>						
021067 002	5394868	Jun 25, 2012	DP		NPP	Feb 01, 2011
	5394868*PED	Dec 25, 2012				
	5687710	Nov 18, 2014	DP			
	5687710*PED	May 18, 2015				
	5829434	Nov 03, 2015	DP			
	5829434*PED	May 03, 2016				
	5889015	Jan 27, 2014	U-645			
	5889015*PED	Jul 27, 2014				
	6057307	Jan 27, 2014	DP U-645			
	6057307*PED	Jul 27, 2014				
	6240918	Feb 20, 2017	DP			
	6240918*PED	Aug 20, 2017				
	6365581	Jan 27, 2014	U-645			
	6365581*PED	Jul 27, 2014				
	6503537	Mar 17, 2018	DP			
	6503537*PED	Sep 17, 2018				
	6677322	Jan 27, 2014	U-645			
	6677322*PED	Jul 27, 2014				
	6949532	Jan 27, 2014	U-645			
	6949532*PED	Jul 27, 2014				
<u>MORPHINE SULFATE - AVINZA</u>						
021260 005	6066339	Nov 25, 2017	DP			
<u>MORPHINE SULFATE - AVINZA</u>						
021260 006	6066339	Nov 25, 2017	DP			
<u>MORPHINE SULFATE; NALTREXONE HYDROCHLORIDE - EMBEDA</u>						
022321 001	>A> 5202128	Apr 13, 2010	DP U-43		>A> NC	Aug 13, 2012
	>A> 5378474	Mar 23, 2010	DP			
<u>MORPHINE SULFATE; NALTREXONE HYDROCHLORIDE - EMBEDA</u>						
022321 002	>A> 5202128	Apr 13, 2010	DP U-43		>A> NC	Aug 13, 2012
	>A> 5378474	Mar 23, 2010	DP			
<u>MORPHINE SULFATE; NALTREXONE HYDROCHLORIDE - EMBEDA</u>						
022321 003	>A> 5202128	Apr 13, 2010	DP U-43		>A> NC	Aug 13, 2012
	>A> 5378474	Mar 23, 2010	DP			
<u>MORPHINE SULFATE; NALTREXONE HYDROCHLORIDE - EMBEDA</u>						
022321 004	>A> 5202128	Apr 13, 2010	DP U-43		>A> NC	Aug 13, 2012
	>A> 5378474	Mar 23, 2010	DP			
<u>MORPHINE SULFATE; NALTREXONE HYDROCHLORIDE - EMBEDA</u>						
022321 005	>A> 5202128	Apr 13, 2010	DP U-43		>A> NC	Aug 13, 2012
	>A> 5378474	Mar 23, 2010	DP			
<u>MORPHINE SULFATE; NALTREXONE HYDROCHLORIDE - EMBEDA</u>						
022321 006	>A> 5202128	Apr 13, 2010	DP U-43		>A> NC	Aug 13, 2012
	>A> 5378474	Mar 23, 2010	DP			
<u>NICOTINE POLACRILEX - NICORETTE</u>						
022360 001	5110605	Aug 21, 2010	DP			

PATENT & EXCLUSIVITY DRUG PRODUCT LIST - CUMULATIVE SUPPLEMENT 8 - August 2009

See List footnote for information regarding List content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	PATENT DELIST REQUESTED	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>NICOTINE POLACRILEX - NICORETTE</u>						
022360 002	5110605	Aug 21, 2010	DP			
<u>NITROGLYCERIN - NITROMIST</u>						
021780 001	5869082	Apr 16, 2016	DP			
<u>OLANZAPINE - ZYPREXA</u>						
020592 001	6960577	Nov 01, 2017	U-963		I-591	Mar 19, 2012
<u>OLANZAPINE - ZYPREXA</u>						
020592 002	6960577	Nov 01, 2017	U-963		I-591	Mar 19, 2012
<u>OLANZAPINE - ZYPREXA</u>						
020592 003	6960577	Nov 01, 2017	U-963		I-591	Mar 19, 2012
<u>OLANZAPINE - ZYPREXA</u>						
020592 004	6960577	Nov 01, 2017	U-963		I-591	Mar 19, 2012
<u>OLANZAPINE - ZYPREXA</u>						
020592 005	6960577	Nov 01, 2017	U-963		I-591	Mar 19, 2012
<u>OLANZAPINE - ZYPREXA</u>						
020592 006	6960577	Nov 01, 2017	U-963		I-591	Mar 19, 2012
<u>OLANZAPINE - ZYPREXA ZYDIS</u>						
021086 001	6960577	Nov 01, 2017	U-964		I-591	Mar 19, 2012
<u>OLANZAPINE - ZYPREXA ZYDIS</u>						
021086 002	6960577	Nov 01, 2017	U-964		I-591	Mar 19, 2012
<u>OLANZAPINE - ZYPREXA ZYDIS</u>						
021086 003	6960577	Nov 01, 2017	U-964		I-591	Mar 19, 2012
<u>OLANZAPINE - ZYPREXA ZYDIS</u>						
021086 004	6960577	Nov 01, 2017	U-964		I-591	Mar 19, 2012
<u>OLOPATADINE HYDROCHLORIDE - PATADAY</u>						
021545 001	5116863	Dec 18, 2010	DS DP			
	5116863*PED	Jun 18, 2011				
	5641805	Jun 06, 2015	U-765			
	5641805*PED	Dec 06, 2015				
	6995186	Nov 12, 2023	DP U-765			
	6995186*PED	May 12, 2024				
	7402609	Jun 19, 2022	DP			
	7402609*PED	Dec 19, 2022				
<u>OLOPATADINE HYDROCHLORIDE - PATANASE</u>						
021861 001	5116863	Dec 18, 2010	DS DP		NDF	Apr 15, 2011
	5116863*PED	Jun 18, 2011			PED	Oct 15, 2011
<u>OLOPATADINE HYDROCHLORIDE - PATANOL</u>						
020688 001	5116863	Dec 18, 2010				
	5116863*PED	Jun 18, 2011				
	5641805	Jun 06, 2015	U-184			
	5641805*PED	Dec 06, 2015				
<u>OMEGA-3-ACID ETHYL ESTERS - LOVAZA</u>						
021654 001	5656667	Apr 10, 2017	DS DP U-822			

PATENT & EXCLUSIVITY DRUG PRODUCT LIST - CUMULATIVE SUPPLEMENT 8 - August 2009

See List footnote for information regarding List content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	PATENT DELIST REQUESTED	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>OXYBUTYNIN CHLORIDE - GELNIQUE</u>						
022204 001	7029694	Apr 26, 2020	DP U-318		NDF	Jan 27, 2012
	7179483	Apr 26, 2020	U-318			
<u>PALIPERIDONE - INVEGA</u>						
021999 001	>A> 5158952	Apr 09, 2012	DP U-90		>A> I-606 >A> I-605	Jul 31, 2012 Jul 31, 2012
<u>PALIPERIDONE - INVEGA</u>						
021999 002	>A> 5158952	Apr 09, 2012	DP U-90		>A> I-606 >A> I-605	Jul 31, 2012 Jul 31, 2012
<u>PALIPERIDONE - INVEGA</u>						
021999 003	>A> 5158952	Apr 09, 2012	DP U-90		>A> I-606 >A> I-605	Jul 31, 2012 Jul 31, 2012
<u>PALIPERIDONE - INVEGA</u>						
021999 004	>A> 5158952	Apr 09, 2012	DP U-90			
<u>PALIPERIDONE - INVEGA</u>						
021999 006	>A> 5158952	Apr 09, 2012	DP U-90		>A> I-606 >A> I-605	Jul 31, 2012 Jul 31, 2012
<u>PALIPERIDONE PALMITATE - INVEGA SUSTENNA</u>						
022264 001	>A> 5254556	Oct 27, 2009	DS DP U-543		>A> NDF	Jul 31, 2012
	>A> 5352459	Dec 16, 2012	DP		>A> NCE	Dec 19, 2011
	>A> 6077843	May 12, 2017	DP U-543			
	>A> 6555544	Nov 10, 2018	DP U-543			
<u>PALIPERIDONE PALMITATE - INVEGA SUSTENNA</u>						
022264 002	>A> 5254556	Oct 27, 2009	DS DP U-543		>A> NDF	Jul 31, 2012
	>A> 5352459	Dec 16, 2012	DP		>A> NCE	Dec 19, 2011
	>A> 6077843	May 12, 2017	DP U-543			
	>A> 6555544	Nov 10, 2018	DP U-543			
<u>PALIPERIDONE PALMITATE - INVEGA SUSTENNA</u>						
022264 003	>A> 5254556	Oct 27, 2009	DS DP U-543		>A> NDF	Jul 31, 2012
	>A> 5352459	Dec 16, 2012	DP		>A> NCE	Dec 19, 2011
	>A> 6077843	May 12, 2017	DP U-543			
	>A> 6555544	Nov 10, 2018	DP U-543			
<u>PALIPERIDONE PALMITATE - INVEGA SUSTENNA</u>						
022264 004	>A> 5254556	Oct 27, 2009	DS DP U-543		>A> NDF	Jul 31, 2012
	>A> 5352459	Dec 16, 2012	DP		>A> NCE	Dec 19, 2011
	>A> 6077843	May 12, 2017	DP U-543			
	>A> 6555544	Nov 10, 2018	DP U-543			
<u>PALIPERIDONE PALMITATE - INVEGA SUSTENNA</u>						
022264 005	>A> 5254556	Oct 27, 2009	DS DP U-543		>A> NDF	Jul 31, 2012
	>A> 5352459	Dec 16, 2012	DP		>A> NCE	Dec 19, 2011
	>A> 6077843	May 12, 2017	DP U-543			
	>A> 6555544	Nov 10, 2018	DP U-543			
<u>PANCRELIPASE (AMYLASE;LIPASE;PROTEASE) - CREON</u>						
020725 001					>A> NCE	Apr 30, 2014
<u>PANCRELIPASE (AMYLASE;LIPASE;PROTEASE) - CREON</u>						
020725 002					>A> NCE	Apr 30, 2014

PATENT & EXCLUSIVITY DRUG PRODUCT LIST - CUMULATIVE SUPPLEMENT 8 - August 2009

See List footnote for information regarding List content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	PATENT DELIST REQUESTED	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>PANCRELIPASE (AMYLASE;LIPASE;PROTEASE) - CREON</u>						
020725 003					>A> NCE	Apr 30, 2014
<u>PANCRELIPASE (AMYLASE;LIPASE;PROTEASE) - ZENPEP</u>						
022210 001					>A> NCE	Aug 27, 2014
<u>PANCRELIPASE (AMYLASE;LIPASE;PROTEASE) - ZENPEP</u>						
022210 002					>A> NCE	Aug 27, 2014
<u>PANCRELIPASE (AMYLASE;LIPASE;PROTEASE) - ZENPEP</u>						
022210 003					>A> NCE	Aug 27, 2014
<u>PANCRELIPASE (AMYLASE;LIPASE;PROTEASE) - ZENPEP</u>						
022210 004					>A> NCE	Aug 27, 2014
<u>PANTOPRAZOLE SODIUM - PROTONIX</u>						
020987 001	4758579	Jul 19, 2010				
	4758579*PED	Jan 19, 2011				
	5997903	Dec 07, 2016				
	5997903*PED	Jun 07, 2017				
<u>PANTOPRAZOLE SODIUM - PROTONIX</u>						
020987 002	4758579	Jul 19, 2010				
	4758579*PED	Jan 19, 2011				
	5997903	Dec 07, 2016				
	5997903*PED	Jun 07, 2017				
<u>PANTOPRAZOLE SODIUM - PROTONIX</u>						
022020 001	4758579	Jul 19, 2010	DS DP U-859			
	4758579*PED	Jan 19, 2011				
	7544370	Feb 07, 2026	DP			
	7544370*PED	Aug 07, 2026				
	7550153	Sep 30, 2024		U-859		
	7550153*PED	Mar 30, 2025				
	7553498	Sep 30, 2024		U-859		
	7553498*PED	Mar 30, 2025				
<u>PANTOPRAZOLE SODIUM - PROTONIX IV</u>						
020988 001	4758579	Jul 19, 2010				
	4758579*PED	Jan 19, 2011				
	6780881	Nov 17, 2021	DP			
	6780881*PED	May 17, 2022				
	7351723	Nov 17, 2021	DP			
	7351723*PED	May 17, 2022				
<u>PARICALCITOL - ZEMPLAR</u>						
021606 001					I-599	Jun 29, 2012
<u>PARICALCITOL - ZEMPLAR</u>						
021606 002					I-599	Jun 29, 2012
<u>PARICALCITOL - ZEMPLAR</u>						
021606 003					I-599	Jun 29, 2012
<u>PEMETREXED DISODIUM - ALIMTA</u>						
021462 001					I-601	Jul 02, 2012
<u>PEMETREXED DISODIUM - ALIMTA</u>						
021462 002					I-601	Jul 02, 2012

PATENT & EXCLUSIVITY DRUG PRODUCT LIST - CUMULATIVE SUPPLEMENT 8 - August 2009

See List footnote for information regarding List content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	PATENT DELIST REQUESTED	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>PHENTOLAMINE MESYLATE - ORAVERSE</u>						
022159 001	6764678	May 11, 2021	U-967			
	6872390	May 11, 2021	DP			
	7229630	Jun 20, 2023	DP			
	>A> 7569230	Oct 17, 2023	U-967			
	>A> 7575757	Apr 21, 2025	DP			
<u>PITAVASTATIN CALCIUM - LIVALO</u>						
022363 001	>A> 5753675	May 19, 2015	DS DP U-998		>A> NCE	Aug 03, 2014
	>A> 5854259	Dec 29, 2015	DP			
	>A> 5856336	Jan 05, 2016	DS U-998			
	>A> 6465477	Dec 20, 2016	DP			
	>A> 7022713	Feb 19, 2024	U-998			
<u>PITAVASTATIN CALCIUM - LIVALO</u>						
022363 002	>A> 5753675	May 19, 2015	DS DP U-998		>A> NCE	Aug 03, 2014
	>A> 5854259	Dec 29, 2015	DP			
	>A> 5856336	Jan 05, 2016	DS U-998			
	>A> 6465477	Dec 20, 2016	DP			
	>A> 7022713	Feb 19, 2024	U-998			
<u>PITAVASTATIN CALCIUM - LIVALO</u>						
022363 003	>A> 5753675	May 19, 2015	DS DP U-998		>A> NCE	Aug 03, 2014
	>A> 5854259	Dec 29, 2015	DP			
	>A> 5856336	Jan 05, 2016	DS U-998			
	>A> 6465477	Dec 20, 2016	DP			
	>A> 7022713	Feb 19, 2024	U-998			
<u>PRAMLintide ACETATE - SYMLIN</u>						
021332 002	5814600	Sep 29, 2015	U-639			
	5814600	Sep 29, 2015	U-638			
	5814600	Sep 29, 2015	U-637			
	5998367	Mar 08, 2011	DS DP			
	6114304	Sep 05, 2017	U-640			
	6114304	Sep 05, 2017	U-637			
	6608029	Sep 07, 2013	U-641			
	6608029	Sep 07, 2013	U-640			
	6608029	Sep 07, 2013	U-637			
	6610824	Mar 03, 2011	DS			
	7407934	Mar 08, 2011	U-640			
	7407934	Mar 08, 2011	U-637			

PATENT & EXCLUSIVITY DRUG PRODUCT LIST - CUMULATIVE SUPPLEMENT 8 - August 2009

See List footnote for information regarding List content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	PATENT DELIST REQUESTED	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>PRAMLINTIDE ACETATE - SYMLIN</u>						
021332 003	5814600	Sep 29, 2015	U-639			
	5814600	Sep 29, 2015	U-638			
	5814600	Sep 29, 2015	U-637			
	5998367	Mar 08, 2011	DS DP			
	6114304	Sep 05, 2017	U-640			
	6114304	Sep 05, 2017	U-637			
	6608029	Sep 07, 2013	U-641			
	6608029	Sep 07, 2013	U-640			
	6608029	Sep 07, 2013	U-637			
	6610824	Mar 03, 2011	DS			
	7407934	Mar 08, 2011	U-640			
	7407934	Mar 08, 2011	U-637			
<u>PRASUGREL HYDROCHLORIDE - EFFIENT</u>						
022307 001	5288726	Sep 08, 2012	DS DP U-991		NCE	Jul 10, 2014
	6693115	Jul 03, 2021	DS DP U-991			
<u>PRASUGREL HYDROCHLORIDE - EFFIENT</u>						
022307 002	5288726	Sep 08, 2012	DS DP U-991		NCE	Jul 10, 2014
	6693115	Jul 03, 2021	DS DP U-991			
<u>QUETIAPINE FUMARATE - SEROQUEL</u>						
020639 001	4879288	Sep 26, 2011	DS DP U-550		I-560	May 13, 2011
	4879288*PED	Mar 26, 2012			I-503	Oct 20, 2009
					PED	Nov 13, 2011
					PED	Apr 20, 2010
<u>QUETIAPINE FUMARATE - SEROQUEL</u>						
020639 002	4879288	Sep 26, 2011	DS DP U-550		I-560	May 13, 2011
	4879288*PED	Mar 26, 2012			I-503	Oct 20, 2009
					PED	Nov 13, 2011
					PED	Apr 20, 2010
<u>QUETIAPINE FUMARATE - SEROQUEL</u>						
020639 003	4879288	Sep 26, 2011	DS DP U-550		I-560	May 13, 2011
	4879288*PED	Mar 26, 2012			I-503	Oct 20, 2009
					PED	Nov 13, 2011
					PED	Apr 20, 2010
<u>QUETIAPINE FUMARATE - SEROQUEL</u>						
020639 004	4879288	Sep 26, 2011	DS DP U-550		I-560	May 13, 2011
	4879288*PED	Mar 26, 2012			I-503	Oct 20, 2009
					PED	Nov 13, 2011
					PED	Apr 20, 2010
<u>QUETIAPINE FUMARATE - SEROQUEL</u>						
020639 005	4879288	Sep 26, 2011	DS DP U-550		I-560	May 13, 2011
	4879288*PED	Mar 26, 2012			I-503	Oct 20, 2009
					PED	Nov 13, 2011
					PED	Apr 20, 2010
<u>QUETIAPINE FUMARATE - SEROQUEL</u>						
020639 006	4879288	Sep 26, 2011	DS DP U-550		I-560	May 13, 2011
	4879288*PED	Mar 26, 2012			I-503	Oct 20, 2009
					PED	Nov 13, 2011
					PED	Apr 20, 2010

PATENT & EXCLUSIVITY DRUG PRODUCT LIST - CUMULATIVE SUPPLEMENT 8 - August 2009

See List footnote for information regarding List content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	PATENT DELIST REQUESTED	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>QUETIAPINE FUMARATE - SEROQUEL</u>						
020639 007	4879288	Sep 26, 2011	DS DP U-550		I-560	May 13, 2011
	4879288*PED	Mar 26, 2012			I-503 PED PED	Oct 20, 2009 Nov 13, 2011 Apr 20, 2010
<u>QUETIAPINE FUMARATE - SEROQUEL XR</u>						
022047 001	4879288	Sep 26, 2011	DS DP U-814		D-117	Oct 08, 2011
	4879288	Sep 26, 2011	DS DP U-601		I-576	Oct 08, 2011
	4879288*PED	Mar 26, 2012			I-575	Oct 08, 2011
	5948437	May 28, 2017	DP U-814		I-574	Oct 08, 2011
	5948437	May 28, 2017	DP U-601		NDF	May 17, 2010
	5948437*PED	Nov 28, 2017			PED	Apr 08, 2012
					PED	Nov 17, 2010
<u>QUETIAPINE FUMARATE - SEROQUEL XR</u>						
022047 002	4879288	Sep 26, 2011	DS DP U-814		D-117	Oct 08, 2011
	4879288	Sep 26, 2011	DS DP U-601		I-576	Oct 08, 2011
	4879288*PED	Mar 26, 2012			I-575	Oct 08, 2011
	5948437	May 28, 2017	DP U-814		I-574	Oct 08, 2011
	5948437	May 28, 2017	DP U-601		NDF	May 17, 2010
	5948437*PED	Nov 28, 2017			PED	Apr 08, 2012
					PED	Nov 17, 2010
<u>QUETIAPINE FUMARATE - SEROQUEL XR</u>						
022047 003	4879288	Sep 26, 2011	DS DP U-814		D-117	Oct 08, 2011
	4879288	Sep 26, 2011	DS DP U-601		I-576	Oct 08, 2011
	4879288*PED	Mar 26, 2012			I-575	Oct 08, 2011
	5948437	May 28, 2017	DP U-814		I-574	Oct 08, 2011
	5948437	May 28, 2017	DP U-601		NDF	May 17, 2010
	5948437*PED	Nov 28, 2017			PED	Apr 08, 2012
					PED	Nov 17, 2010
<u>QUETIAPINE FUMARATE - SEROQUEL XR</u>						
022047 004	4879288	Sep 26, 2011	DS DP U-814		D-117	Oct 08, 2011
	4879288	Sep 26, 2011	DS DP U-601		I-576	Oct 08, 2011
	4879288*PED	Mar 26, 2012			I-575	Oct 08, 2011
	5948437	May 28, 2017	DP U-814		I-574	Oct 08, 2011
	5948437	May 28, 2017	DP U-601		NDF	May 17, 2010
	5948437*PED	Nov 28, 2017			PED	Apr 08, 2012
					PED	Nov 17, 2010
<u>QUETIAPINE FUMARATE - SEROQUEL XR</u>						
022047 005	4879288	Sep 26, 2011	DS DP U-814		D-117	Oct 08, 2011
	4879288	Sep 26, 2011	DS DP U-601		I-576	Oct 08, 2011
	4879288*PED	Mar 26, 2012			I-575	Oct 08, 2011
	5948437	May 28, 2017	DP U-814		I-574	Oct 08, 2011
	5948437	May 28, 2017	DP U-601		NDF	May 17, 2010
	5948437*PED	Nov 28, 2017			PED	Apr 08, 2012
					PED	Nov 17, 2010
<u>RASAGILINE MESYLATE - AZILECT</u>						
021641 001	5453446	Feb 07, 2017	U-219			
	>A> 7572834	Dec 05, 2026	DP			
<u>RASAGILINE MESYLATE - AZILECT</u>						
021641 002	5453446	Feb 07, 2017	U-219			
	>A> 7572834	Dec 05, 2026	DP			

PATENT & EXCLUSIVITY DRUG PRODUCT LIST - CUMULATIVE SUPPLEMENT 8 - August 2009

See List footnote for information regarding List content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	PATENT DELIST REQUESTED	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>REMIFENTANIL HYDROCHLORIDE - ULTIVA</u>						
020630 001	5019583	Jul 12, 2010	DS DP U-952			
	5019583*PED	Jan 12, 2011				
<u>REMIFENTANIL HYDROCHLORIDE - ULTIVA</u>						
020630 002	5019583	Jul 12, 2010	DS DP U-952			
	5019583*PED	Jan 12, 2011				
<u>REMIFENTANIL HYDROCHLORIDE - ULTIVA</u>						
020630 003	5019583	Jul 12, 2010	DS DP U-952			
	5019583*PED	Jan 12, 2011				
<u>REPAGLINIDE - PRANDIN</u>						
020741 001	6677358	Jun 12, 2018	DS DP U-968			
<u>REPAGLINIDE - PRANDIN</u>						
020741 002	6677358	Jun 12, 2018	DS DP U-968			
<u>REPAGLINIDE - PRANDIN</u>						
020741 003	6677358	Jun 12, 2018	DS DP U-968			
<u>RISEDRONATE SODIUM - ACTONEL</u>						
020835 001	5583122	Dec 10, 2013	U-222		M-61	Jul 23, 2012
	5583122*PED	Jun 10, 2014			PED	Jan 23, 2013
	6096342	Nov 22, 2011				
	6096342*PED	May 22, 2012				
	6165513	Jun 10, 2018				
	6165513*PED	Dec 10, 2018				
<u>RISEDRONATE SODIUM - ACTONEL</u>						
020835 002	5583122	Dec 10, 2013	U-222		M-61	Jul 23, 2012
	5583122*PED	Jun 10, 2014			PED	Jan 23, 2013
	6096342	Nov 22, 2011				
	6096342*PED	May 22, 2012				
	6165513	Jun 10, 2018				
	6165513*PED	Dec 10, 2018				
<u>RISEDRONATE SODIUM - ACTONEL</u>						
020835 003	5583122	Dec 10, 2013	DS DP U-756		I-309	Aug 11, 2009
	5583122	Dec 10, 2013	DS DP U-222		M-61	Jul 23, 2012
	5583122*PED	Jun 10, 2014			PED	Jan 23, 2013
	5994329	Jul 17, 2018	U-353		PED	Feb 11, 2010
	5994329*PED	Jan 17, 2019				
	6015801	Jul 17, 2018	U-353			
	6015801*PED	Jan 17, 2019				
	6096342	Nov 22, 2011	DP			
	6096342*PED	May 22, 2012				
	6165513	Jun 10, 2018	DP			
	6165513*PED	Dec 10, 2018				
	6432932	Jul 17, 2018	U-595			
	6432932*PED	Jan 17, 2019				
	6465443	Aug 14, 2018	DP			
	6465443*PED	Feb 14, 2019				

PATENT & EXCLUSIVITY DRUG PRODUCT LIST - CUMULATIVE SUPPLEMENT 8 - August 2009

See List footnote for information regarding List content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE		PATENT CODES			PATENT DELIST REQUESTED	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE			
<u>RISEDRONATE SODIUM - ACTONEL</u>												
020835 004	5583122	Dec	10,	2013	DS	DP	U-353	D-105 M-61 PED PED	Apr Jul Jan Oct	16, <td>2010</td>	2010	
	5583122*PED	Jun	10,	2014							23,	2012
	6096342	Nov	22,	2011	DP	U-353	23,				2013	
	6096342*PED	May	22,	2012			16,				2010	
	6165513	Jun	10,	2018	DP							
	6165513*PED	Dec	10,	2018								
<u>RISEDRONATE SODIUM - ACTONEL</u>												
020835 005	5583122	Dec	10,	2013	DS	DP	U-353	M-61 NS PED PED	Jul Apr Jan Oct	23, <td>2012</td>	2012	
	5583122*PED	Jun	10,	2014							22,	2011
	6165513	Jun	10,	2018	DP						23,	2013
	6165513*PED	Dec	10,	2018		22,	2011					
	7192938	May	06,	2023	U-353							
	7192938*PED	Nov	06,	2023								
<u>RISPERIDONE - RISPERDAL CONSTA</u>												
021346 001	5688801	Nov	18,	2014			U-972	I-597 I-596	May May	15, <td>2012</td>	2012	
	5688801	Nov	18,	2014							15,	2012
	>A> 6667061	May	25,	2020	DP							
	>A> 6667061*PED	Nov	25,	2020								
	7547452	Nov	19,	2013	DP							
	7547452*PED	May	19,	2014								
<u>RISPERIDONE - RISPERDAL CONSTA</u>												
021346 002	5688801	Nov	18,	2014			U-972	I-597 I-596	May May	15, <td>2012</td>	2012	
	5688801	Nov	18,	2014							15,	2012
	>A> 6667061	May	25,	2020	DP							
	>A> 6667061*PED	Nov	25,	2020								
	7547452	Nov	19,	2013	DP							
	7547452*PED	May	19,	2014								
<u>RISPERIDONE - RISPERDAL CONSTA</u>												
021346 003	5688801	Nov	18,	2014			U-972	I-597 I-596	May May	15, <td>2012</td>	2012	
	5688801	Nov	18,	2014							15,	2012
	>A> 6667061	May	25,	2020	DP							
	>A> 6667061*PED	Nov	25,	2020								
	7547452	Nov	19,	2013	DP							
	7547452*PED	May	19,	2014								
<u>RISPERIDONE - RISPERDAL CONSTA</u>												
021346 004	5688801	Nov	18,	2014			U-972	I-597 I-596	May May	15, <td>2012</td>	2012	
	5688801	Nov	18,	2014							15,	2012
	>A> 6667061	May	25,	2020	DP							
	>A> 6667061*PED	Nov	25,	2020								
	7547452	Nov	19,	2013	DP							
	7547452*PED	May	19,	2014								
<u>RISPERIDONE - RISPERIDONE</u>												
076440 001								PC	Jul	29,	2009	
<u>ROPINIROLE HYDROCHLORIDE - REQUIP XL</u>												
022008 006	5422123	Jun	06,	2012		DP		NDF	Jun	13,	2011	

PATENT & EXCLUSIVITY DRUG PRODUCT LIST - CUMULATIVE SUPPLEMENT 8 - August 2009

See List footnote for information regarding List content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	PATENT DELIST REQUESTED	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>ROSUVASTATIN CALCIUM - CRESTOR</u>						
021366 002	6316460	Aug 04, 2020	DP		I-573	Nov 06, 2011
	6316460*PED	Feb 04, 2021			I-547	Nov 08, 2010
	6858618	Dec 17, 2021	U-618		PED	May 06, 2012
	6858618*PED	Jun 17, 2022			PED	May 08, 2011
	RE37314	Jan 08, 2016	DS			
	RE37314*PED	Jul 08, 2016				
<u>ROSUVASTATIN CALCIUM - CRESTOR</u>						
021366 003	6316460	Aug 04, 2020	DP		I-573	Nov 06, 2011
	6316460*PED	Feb 04, 2021			I-547	Nov 08, 2010
	6858618	Dec 17, 2021	U-618		PED	May 06, 2012
	6858618*PED	Jun 17, 2022			PED	May 08, 2011
	RE37314	Jan 08, 2016	DS			
	RE37314*PED	Jul 08, 2016				
<u>ROSUVASTATIN CALCIUM - CRESTOR</u>						
021366 004	6316460	Aug 04, 2020	DP		I-573	Nov 06, 2011
	6316460*PED	Feb 04, 2021			I-547	Nov 08, 2010
	6858618	Dec 17, 2021	U-618		PED	May 06, 2012
	6858618*PED	Jun 17, 2022			PED	May 08, 2011
	RE37314	Jan 08, 2016	DS			
	RE37314*PED	Jul 08, 2016				
<u>ROSUVASTATIN CALCIUM - CRESTOR</u>						
021366 005	6316460	Aug 04, 2020	DP		I-573	Nov 06, 2011
	6316460*PED	Feb 04, 2021			I-547	Nov 08, 2010
	6858618	Dec 17, 2021	U-618		PED	May 06, 2012
	6858618*PED	Jun 17, 2022			PED	May 08, 2011
	RE37314	Jan 08, 2016	DS			
	RE37314*PED	Jul 08, 2016				
<u>SALMETEROL XINAFOATE - SEREVENT</u>						
020692 001	5590645	Mar 01, 2011	DP			
	5590645*PED	Sep 01, 2011				
	5860419	Mar 01, 2011	DP			
	5860419*PED	Sep 01, 2011				
	5873360	Feb 23, 2016	DP			
	5873360*PED	Aug 23, 2016				
	6032666	Mar 01, 2011	DP			
	6032666*PED	Sep 01, 2011				
	6378519	Mar 01, 2011	DP			
	6378519*PED	Sep 01, 2011				
	6536427	Mar 01, 2011	DP			
	6536427*PED	Sep 01, 2011				
	6792945	Mar 01, 2011	DP			
	6792945*PED	Sep 01, 2011				
	7225808	Mar 01, 2011	DP			
	7225808*PED	Sep 01, 2011				
	7389775	Mar 01, 2011	DP			
	7389775*PED	Sep 01, 2011				

PATENT & EXCLUSIVITY DRUG PRODUCT LIST - CUMULATIVE SUPPLEMENT 8 - August 2009

See List footnote for information regarding List content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	PATENT DELIST REQUESTED	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>SAPROPTERIN DIHYDROCHLORIDE - KUVAN</u>						
022181 001	7566462	Nov 16, 2025	DP			
	7566714	Nov 17, 2024	U-989			
<u>SAXAGLIPTIN HYDROCHLORIDE - ONGLYZA</u>						
022350 001	>A> 6395767	Feb 16, 2021	DS DP U-995		NCE	Jul 31, 2014
<u>SAXAGLIPTIN HYDROCHLORIDE - ONGLYZA</u>						
022350 002	>A> 6395767	Feb 16, 2021	DS DP U-995		NCE	Jul 31, 2014
<u>SEVELAMER CARBONATE - RENVELA</u>						
022318 001					>A> NE >A> NDF	Oct 19, 2010 Aug 12, 2012
<u>SEVELAMER CARBONATE - RENVELA</u>						
022318 002					>A> NE >A> NDF	Oct 19, 2010 Aug 12, 2012
<u>SILDENAFIL CITRATE - REVATIO</u>						
021845 001					I-598	May 07, 2012
<u>SILODOSIN - RAPAFLO</u>						
022206 001	5780485	Nov 13, 2012	U-902			
<u>SILODOSIN - RAPAFLO</u>						
022206 002	5780485	Nov 13, 2012	U-902			
<u>SINECATECHINS - VEREGEN</u>						
021902 001	5795911	Oct 31, 2020	U-172			
<u>SOMATROPIN RECOMBINANT - HUMATROPE</u>						
019640 001					I-585	Mar 12, 2012
<u>SOMATROPIN RECOMBINANT - HUMATROPE</u>						
019640 004	5612315	Mar 18, 2014	DP		I-585	Mar 12, 2012
<u>SOMATROPIN RECOMBINANT - HUMATROPE</u>						
019640 005	5612315	Mar 18, 2014	DP		I-585	Mar 12, 2012
<u>SOMATROPIN RECOMBINANT - HUMATROPE</u>						
019640 006	5612315	Mar 18, 2014	DP		I-585	Mar 12, 2012
<u>SOMATROPIN RECOMBINANT - HUMATROPE</u>						
019640 007	5612315	Mar 18, 2014	DP		I-585	Mar 12, 2012
<u>SOMATROPIN RECOMBINANT - NORDITROPIN NORDIFLEX</u>						
021148 004	5849700	Dec 15, 2015	U-340			
	5849704	Dec 15, 2015	DP U-340			
<u>SOMATROPIN RECOMBINANT - NORDITROPIN NORDIFLEX</u>						
021148 005	5849700	Dec 15, 2015	U-340			
	5849704	Dec 15, 2015	DP U-340			
<u>SOMATROPIN RECOMBINANT - NORDITROPIN NORDIFLEX</u>						
021148 006	5849700	Dec 15, 2015	U-340			
	5849704	Dec 15, 2015	DP U-340			
<u>SOMATROPIN RECOMBINANT - NORDITROPIN NORDIFLEX</u>						
021148 007	5849700	Dec 15, 2015	U-340		I-572	Oct 31, 2011
	5849704	Dec 15, 2015	DP U-340		I-551 I-536 ODE	Sep 20, 2010 May 31, 2010 May 31, 2014

PATENT & EXCLUSIVITY DRUG PRODUCT LIST - CUMULATIVE SUPPLEMENT 8 - August 2009

See List footnote for information regarding List content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	PATENT DELIST REQUESTED	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>SOMATROPIN RECOMBINANT - SAIZEN</u>						
019764 002	5898030	Apr 27, 2016	DP			
<u>SOMATROPIN RECOMBINANT - SAIZEN</u>						
019764 003	5898030	Apr 27, 2016	DP			
<u>SOMATROPIN RECOMBINANT - SEROSTIM</u>						
020604 001	5898030	Apr 27, 2016	DP			
<u>SOMATROPIN RECOMBINANT - SEROSTIM</u>						
020604 002	5898030	Apr 27, 2016	DP			
<u>SOMATROPIN RECOMBINANT - SEROSTIM</u>						
020604 003	5898030	Apr 27, 2016	DP			
<u>SOMATROPIN RECOMBINANT - SEROSTIM</u>						
020604 004	5898030	Apr 27, 2016	DP			
<u>SOMATROPIN RECOMBINANT - ZORBTIVE</u>						
021597 004	5288703	Oct 07, 2011		U-898		
	5898030	Apr 27, 2016	DP			
<u>SORAFENIB TOSYLATE - NEXAVAR</u>						
021923 001	>A> 7235576	Jan 12, 2020	DS DP			
<u>SOTALOL HYDROCHLORIDE - SOTALOL HYDROCHLORIDE</u>						
022306 001					>A> ODE	Jul 02, 2016
<u>SUMATRIPTAN SUCCINATE - SUMATRIPTAN SUCCINATE</u>						
076572 001					PC	Aug 08, 2009
<u>SUMATRIPTAN SUCCINATE - SUMATRIPTAN SUCCINATE</u>						
076840 001					PC	Aug 08, 2009
<u>SUMATRIPTAN SUCCINATE - SUMATRIPTAN SUCCINATE</u>						
076840 002					PC	Aug 08, 2009
<u>SUMATRIPTAN SUCCINATE - SUMATRIPTAN SUCCINATE</u>						
076840 003					PC	Aug 08, 2009
<u>SUMATRIPTAN SUCCINATE - SUMAVEL DOSEPRO</u>						
022239 001	5891086	Jul 27, 2014	DP			
	5957886	Mar 08, 2016	DP			
	6135979	Mar 21, 2017	DP			
<u>SUNITINIB MALATE - SUTENT</u>						
021938 004	6573293	Feb 15, 2021	DS DP	U-703	NCE	Jan 26, 2011
	7125905	Feb 15, 2021	DS DP			
	7211600	Dec 22, 2020		U-883		
<u>TADALAFIL - ADCIRCA</u>						
022332 001	5859006	Nov 21, 2017	DS DP	U-975	NP	May 22, 2012
	6821975	Nov 19, 2020	DS DP		ODE	May 22, 2016
	7182958	Apr 26, 2020	DP			
<u>TELBIVUDINE - TYZEKA</u>						
022154 001	>A> 6395716	Aug 10, 2019		U-999	NCE	Oct 25, 2011
	>A> 6444652	Aug 10, 2019		U-999		
	>A> 6566344	Aug 10, 2019		U-999		
	>A> 6569837	Aug 10, 2019		U-999		

PATENT & EXCLUSIVITY DRUG PRODUCT LIST - CUMULATIVE SUPPLEMENT 8 - August 2009

See List footnote for information regarding List content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	PATENT DELIST REQUESTED	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>TEMOZOLOMIDE - TEMODAR</u>						
022277 001	5260291	Aug 11, 2013	DS DP U-619			
	5260291*PED	Feb 11, 2014				
	6987108	Sep 08, 2023	DP			
<u>TERIPARATIDE RECOMBINANT HUMAN - FORTEO</u>						
021318 001	7550434	Dec 08, 2018	DP U-982		I-602	Jul 22, 2012
<u>TERIPARATIDE RECOMBINANT HUMAN - FORTEO</u>						
021318 002	6770623	Dec 08, 2018	DP U-982		I-602	Jul 22, 2012
	>A> 6977077	Aug 19, 2019	U-994			
	>A> 6977077	Aug 19, 2019	U-982			
	7144861	Dec 08, 2018	DP			
	>A> 7163684	Aug 19, 2019	U-994			
	>A> 7163684	Aug 19, 2019	U-983			
	>A> 7351414	Aug 19, 2019	U-994			
	>A> 7351414	Aug 19, 2019	U-984			
	7550434	Dec 08, 2018	DP U-982			
<u>TIGECYCLINE - TYGACIL</u>						
021821 001					I-588 I-587 I-586	Mar 20, 2012 Mar 20, 2012 Mar 20, 2012
<u>TOLVAPTAN - SAMSCA</u>						
022275 001	5258510	Nov 02, 2010	DS		NCE	May 19, 2014
	5753677	May 19, 2015	U-978			
<u>TOLVAPTAN - SAMSCA</u>						
022275 002	5258510	Nov 02, 2010	DS		NCE	May 19, 2014
	5753677	May 19, 2015	U-978			
<u>TOLVAPTAN - SAMSCA</u>						
022275 003	5258510	Nov 02, 2010	DS		NCE	May 19, 2014
	5753677	May 19, 2015	U-978			
<u>TOPIRAMATE - TOPAMAX</u>						
020505 001	7498311	Oct 13, 2015	U-955			
	7498311*PED	Apr 13, 2016				
<u>TOPIRAMATE - TOPAMAX</u>						
020505 002	7498311	Oct 13, 2015	U-955			
	7498311*PED	Apr 13, 2016				
<u>TOPIRAMATE - TOPAMAX</u>						
020505 003	7498311	Oct 13, 2015	U-955			
	7498311*PED	Apr 13, 2016				
<u>TOPIRAMATE - TOPAMAX</u>						
020505 004	7498311	Oct 13, 2015	U-955			
	7498311*PED	Apr 13, 2016				
<u>TOPIRAMATE - TOPAMAX</u>						
020505 005	7498311	Oct 13, 2015	U-955			
	7498311*PED	Apr 13, 2016				

PATENT & EXCLUSIVITY DRUG PRODUCT LIST - CUMULATIVE SUPPLEMENT 8 - August 2009

See List footnote for information regarding List content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	PATENT DELIST REQUESTED	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
TOPIRAMATE - TOPAMAX						
020505 006	7498311	Oct 13, 2015	U-955			
	7498311*PED	Apr 13, 2016				
TOPIRAMATE - TOPAMAX						
020844 001	7498311	Oct 13, 2015	U-955			
	7498311*PED	Apr 13, 2016				
TOPIRAMATE - TOPAMAX						
020844 002	7498311	Oct 13, 2015	U-955			
	7498311*PED	Apr 13, 2016				
TOPIRAMATE - TOPAMAX SPRINKLE						
020844 003	7498311	Oct 13, 2015	U-955			
	7498311*PED	Apr 13, 2016				
TOPIRAMATE - TOPIRAMATE						
076448 001					PC	Oct 12, 2009
TOPIRAMATE - TOPIRAMATE						
076448 002					PC	Oct 12, 2009
TOPIRAMATE - TOPIRAMATE						
077868 001					PC	Oct 12, 2009
TOPIRAMATE - TOPIRAMATE						
077868 002					PC	Oct 12, 2009
TRAMADOL HYDROCHLORIDE - RYZOLT						
021745 001					NP	Dec 30, 2011
TRAMADOL HYDROCHLORIDE - RYZOLT						
021745 002					NP	Dec 30, 2011
TRAMADOL HYDROCHLORIDE - RYZOLT						
021745 003					NP	Dec 30, 2011
TREPROSTINIL SODIUM - TYVASO						
022387 001					NDF	Jul 30, 2012
TRYPAN BLUE - MEMBRANEBLUE						
022278 001					NCE ODE	Dec 16, 2009 Dec 16, 2011
UNOPROSTONE ISOPROPYL - RESCULA						
021214 001	>A> 5221763	Jul 15, 2012	DS U-333			
	>A> 6458836	Jul 09, 2021				
VALGANCICLOVIR HYDROCHLORIDE - VALCYTE						
021304 001					>A> I-604 >A> PED	Aug 28, 2012 Feb 28, 2013
VALGANCICLOVIR HYDROCHLORIDE - VALCYTE						
022257 001					>A> NDF >A> PED	Aug 28, 2012 Feb 28, 2013
VIGABATRIN - SABRIL						
020427 001					>A> NCE	Aug 21, 2014
VIGABATRIN - SABRIL						
022006 001					>A> NCE >A> ODE	Aug 21, 2014 Aug 21, 2016

PATENT & EXCLUSIVITY DRUG PRODUCT LIST - CUMULATIVE SUPPLEMENT 8 - August 2009

See List footnote for information regarding List content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	PATENT DELIST REQUESTED	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>ZOLEDRONIC ACID - RECLAST</u>						
021817 001					I-595	May 29, 2012
					I-584	Mar 15, 2012
					I-581	Dec 19, 2011
<u>ZOLPIDEM TARTRATE - EDLUAR</u>						
021997 001	6761910	Sep 24, 2018	DP U-674			
<u>ZOLPIDEM TARTRATE - EDLUAR</u>						
021997 002	6761910	Sep 24, 2018	DP U-674			

Footnote:

1. Patents are published upon receipt by the Orange Book Staff and may not reflect the official receipt date as described in 21 CFR 314.53(d)(5).
2. Patents 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.

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, 29th 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>