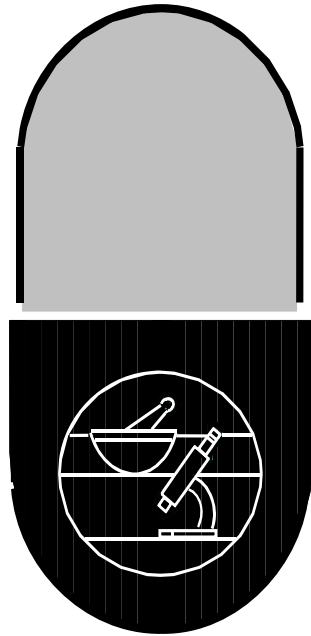


**CUMULATIVE
SUPPLEMENT 01
January 2008**



**APPROVED
DRUG PRODUCTS**

**WITH
THERAPEUTIC EQUIVALENCE EVALUATIONS**

28th EDITION

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

2008

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

**APPROVED DRUG PRODUCTS
with
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28th EDITION

Cumulative Supplement 01

January 2008

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**APPROVED DRUG PRODUCTS
with
THERAPEUTIC EQUIVALENCE EVALUATIONS**

28th EDITION

**CUMULATIVE SUPPLEMENT 01
December 2008**

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, 27th 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are for exportation, are for military use, have been discontinued from marketing or that have had their approvals withdrawn for other than safety or efficacy reasons.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 27th Edition List will then be added to the "Discontinued Drug Product List" appearing in the 28th 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.
 - Refer to CS Section 1.8 Cumulative Supplement Legend for types of changes
 - 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@cderr.fda.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>
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None

1.4 AVAILABILITY OF THE EDITION

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

Commencing with the 25th edition, the Annual Edition and monthly Cumulative Supplements have been provided in downloadable Portable Document Format (PDF) at the EOB home page by clicking on Annual Edition. The PDF annual and cumulative supplements duplicate

previous paper versions. Over time, there will be an archive for the annuals and each year's December Cumulative Supplement.

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

There are historical lists of Orange Book cumulative supplement product monthly changes at

<http://www.fda.gov/cder/rxotcdpl/pdplarchive.htm>

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

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

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

1.5 REPORT OF COUNTS FOR THE PRESCRIPTION DRUG PRODUCT LIST

DESCRIPTION OF REPORT

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

The baseline column (Dec 2004) refers to the products in the Prescription Drug Product List. For each three-month period, a column of quarterly data is added which incorporates counts of product activity from the previous quarter(s) with those in the baseline count.

DEFINITIONS

Drug Product

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

New Molecular Entity

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

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

<u>CATEGORIES COUNTED</u>	<u>DEC 2007</u>	<u>MAR 2007</u>	<u>JUN 2007</u>	<u>SEPT 2007</u>
DRUG PRODUCTS LISTED	12302	12063	11900	12130
SINGLE SOURCE	2489 (20.2%)	2471 (20.5%)	2483 (20.9%)	2494 (20.6%)
MULTISOURCE	9724 (79.0%)	9503 (78.8%)	9328 (78.4%)	9547 (78.7%)
THERAPEUTICALLY EQUIVALENT	9571 (77.8%)	9320 (77.3%)	9148 (76.9%)	9394 (77.4%)
NOT THERAPEUTICALLY EQUIVALENT	153 (1.2%)	183 (1.5%)	180 (1.5%)	153 (1.3%)
EXCEPTIONS ¹	89 (0.7%)	89 (0.7%)	89 (0.7%)	89 (0.7%)
NEW MOLECULAR ENTITIES				
APPROVED	7	4	7	10
NUMBER OF APPLICANTS	693	675	679	683

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 >A>. All changes that occur to the product through the Annual year will be listed. The change month and change code will document the change.

The change code and description:

NEWA	New drug product approval usually in the supplement month.
CAHN	Applicant holder firm name has changed.
CAIN	Change. There has been a change in the Ingredient(s) name. All products will be deleted under the old name and all products will be added under the changed ingredient(s) name.
CDFR	Change. Dosage Form; Route of Administration.
CFTG	Change. A first time generic for the innovator product. A TE Code is added.
CMFD	Change. The product is moved from the Discontinued Section due to a change in marketing status.
CMS1	Change. Miscellaneous addition to list.

CMS2	Change. Miscellaneous deletion from list.
CPOT	Change. Potency amount/unit.
CRLD	Change. Reference Listed Drug.
CTEC	Change. Therapeutic Equivalence Code.
CTNA	Change. Trade Name.
DISC	Discontinued. The Rx or OTC listed product is not being marketed and will be moved to the discontinued section in the next edition.

PRESCRIPTION DRUG PRODUCT LIST - 28TH EDITION

RX DRUG PRODUCT LIST - CUMULATIVE SUPPLEMENT 1 - January 2008

1-1

ACETAMINOPHEN; PROPOXYPHENE NAPSYLATE

TABLET; ORAL

PROPOXYPHENE NAPSYLATE AND ACETAMINOPHEN

>A> AB CONCORD LABS NJ 650MG;100MG N77821 001 Feb 11, 2008 Jan NEWA

ACETOHEXAMIDE

TABLET; ORAL

ACETOHEXAMIDE

>D> AB + BARR 500MG N70870 001 Feb 09, 1987 Jan DISC

>A> @ 500MG N70870 001 Feb 09, 1987 Jan DISC

>D> AB WATSON LABS 500MG N71894 001 Nov 25, 1987 Jan CRLD

>A> + 500MG N71894 001 Nov 25, 1987 Jan CRLD

ALENDRONATE SODIUM

TABLET; ORAL

ALENDRONATE SODIUM

>A> AB BARR EQ 70MG BASE N76184 001 Feb 06, 2008 Jan NEWA

>A> AB TEVA PHARMS EQ 5MG BASE N75710 001 Feb 06, 2008 Jan NEWA

>A> AB EQ 10MG BASE N75710 002 Feb 06, 2008 Jan NEWA

>A> AB EQ 35MG BASE N75710 003 Feb 06, 2008 Jan NEWA

>A> AB EQ 40MG BASE N75710 004 Feb 06, 2008 Jan NEWA

>A> AB EQ 70MG BASE N75710 005 Feb 06, 2008 Jan NEWA

FOSAMAX

>D> MERCK EQ 5MG BASE N20560 003 Apr 25, 1997 Jan CFTG

>D> EQ 10MG BASE N20560 001 Sep 29, 1995 Jan CFTG

>D> EQ 35MG BASE N20560 004 Oct 20, 2000 Jan CFTG

>D> EQ 40MG BASE N20560 002 Sep 29, 1995 Jan CFTG

>D> + EQ 70MG BASE N20560 005 Oct 20, 2000 Jan CFTG

>A> AB MERCK AND CO INC EQ 5MG BASE N20560 003 Apr 25, 1997 Jan CFTG

>A> AB EQ 10MG BASE N20560 001 Sep 29, 1995 Jan CFTG

>A> AB EQ 35MG BASE N20560 004 Oct 20, 2000 Jan CFTG

>A> AB EQ 40MG BASE N20560 002 Sep 29, 1995 Jan CFTG

>A> AB + EQ 70MG BASE N20560 005 Oct 20, 2000 Jan CFTG

>A> ALISKIREN HEMIFUMARATE; HYDROCHLOROTHIAZIDE

>A> TABLET; ORAL

>A> TEKTURNA HCT

>A> NOVARTIS EQ 150MG BASE;12.5MG N22107 001 Jan 18, 2008 Jan NEWA

>A> EQ 150MG BASE;25MG N22107 002 Jan 18, 2008 Jan NEWA

>A> EQ 300MG BASE;12.5MG N22107 003 Jan 18, 2008 Jan NEWA

>A> + EQ 300MG BASE;25MG N22107 004 Jan 18, 2008 Jan NEWA

AMLODIPINE BESYLATE

TABLET; ORAL

AMLODIPINE BESYLATE

>A> AB INTERPHARM EQ 2.5MG BASE N78477 001 Jan 16, 2008 Jan NEWA

>A> AB EQ 5MG BASE N78477 002 Jan 16, 2008 Jan NEWA

>A> AB EQ 10MG BASE N78477 003 Jan 16, 2008 Jan NEWA

>A> AB MUTUAL PHARMA EQ 2.5MG BASE N78081 001 Jan 31, 2008 Jan NEWA

>A> AB EQ 5MG BASE N78081 002 Jan 31, 2008 Jan NEWA

>A> AB EQ 10MG BASE N78081 003 Jan 31, 2008 Jan NEWA

AMOXICILLIN

>A>		TABLET, EXTENDED RELEASE; ORAL							
>A>		MOXATAG							
>A>		+ MIDDLEBROOK PHARMS	775MG		N50813	001	Jan 23, 2008	Jan	NEWA

AZITHROMYCIN

		TABLET; ORAL							
		AZITHROMYCIN							
>A>	AB	WOCKHARDT	EQ 250MG BASE		N65404	001	Feb 11, 2008	Jan	NEWA
>A>	AB		EQ 500MG BASE		N65405	001	Feb 11, 2008	Jan	NEWA
>A>	AB		EQ 600MG BASE		N65302	003	Feb 11, 2008	Jan	NEWA

AZTREONAM

		INJECTABLE; INJECTION							
		AZACTAM							
>D>		+ BRISTOL MYERS SQUIBB	500MG/VIAL		N50580	001	Dec 31, 1986	Jan	DISC
>A>		@	500MG/VIAL		N50580	001	Dec 31, 1986	Jan	DISC

BENZONATATE

		CAPSULE; ORAL							
		BENZONATATE							
>D>	AA	AMNEAL PHARM	100MG		N40682	001	Jul 30, 2007	Jan	CAHN
>D>	AA		200MG		N40682	002	Jul 30, 2007	Jan	CAHN
>A>	AA	ORIT LABS LLC	100MG		N40682	001	Jul 30, 2007	Jan	CAHN
>A>	AA		200MG		N40682	002	Jul 30, 2007	Jan	CAHN

BENZTROPINE MESYLATE

		TABLET; ORAL							
		BENZTROPINE MESYLATE							
>A>	AA	ACTAVIS TOTOWA	0.5MG		N40699	001	Feb 14, 2008	Jan	NEWA
>A>	AA		1MG		N40705	001	Feb 14, 2008	Jan	NEWA
>A>	AA		2MG		N40706	001	Feb 14, 2008	Jan	NEWA

BLEOMYCIN SULFATE

		INJECTABLE; INJECTION							
		BLEOMYCIN							
>D>	AP	BEDFORD	EQ 15 UNITS BASE/VIAL		N65042	002	Oct 17, 2001	Jan	CTNA
>D>	AP		EQ 30 UNITS BASE/VIAL		N65042	001	Oct 17, 2001	Jan	CTNA
>D>	AP	HOSPIRA	EQ 15 UNITS BASE/VIAL		N65031	001	Mar 10, 2000	Jan	CTNA
>D>	AP		EQ 30 UNITS BASE/VIAL		N65031	002	Mar 10, 2000	Jan	CTNA
>D>	AP	PHARMACHEMIE BV	EQ 15 UNITS BASE/VIAL		N65201	001	Dec 13, 2007	Jan	CTNA
>D>	AP	TEVA PARENTERAL	EQ 15 UNITS BASE/VIAL		N65033	001	Jun 27, 2000	Jan	CTNA
>D>	AP		EQ 30 UNITS BASE/VIAL		N65033	002	Jun 27, 2000	Jan	CTNA
		BLEOMYCIN SULFATE							
>A>	AP	ABRAXIS PHARM	EQ 15 UNITS BASE/VIAL		N65185	001	Jan 28, 2008	Jan	NEWA
>A>	AP		EQ 30 UNITS BASE/VIAL		N65185	002	Jan 28, 2008	Jan	NEWA
>A>	AP	BEDFORD	EQ 15 UNITS BASE/VIAL		N65042	002	Oct 17, 2001	Jan	CTNA
>A>	AP		EQ 30 UNITS BASE/VIAL		N65042	001	Oct 17, 2001	Jan	CTNA
>A>	AP	HOSPIRA	EQ 15 UNITS BASE/VIAL		N65031	001	Mar 10, 2000	Jan	CTNA
>A>	AP		EQ 30 UNITS BASE/VIAL		N65031	002	Mar 10, 2000	Jan	CTNA
>A>	AP	PHARMACHEMIE BV	EQ 15 UNITS BASE/VIAL		N65201	001	Dec 13, 2007	Jan	CTNA
>A>	AP	TEVA PARENTERAL	EQ 15 UNITS BASE/VIAL		N65033	001	Jun 27, 2000	Jan	CTNA
>A>	AP		EQ 30 UNITS BASE/VIAL		N65033	002	Jun 27, 2000	Jan	CTNA

BRIMONIDINE TARTRATESOLUTION/DROPS; OPHTHALMIC
BRIMONIDINE TARTRATE

>A>	AT	SANDOZ	0.2%	N78075 001	Jan 30, 2008	Jan	NEWA
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CAFFEINE CITRATESOLUTION; ORAL
CAFFEINE CITRATE

>A>	AA	ABRAXIS PHARM PRODS	EQ 30MG BASE/3ML (EQ 10MG BASE/ML)	N78002 001	Jan 31, 2008	Jan	NEWA
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CALCITRIOLINJECTABLE; INJECTION
CALCITRIOL

>A>	AP	AKORN	0.001MG/ML	N78066 001	Jan 29, 2008	Jan	NEWA
>A>	AP		0.002MG/ML	N78066 002	Jan 29, 2008	Jan	NEWA

CALCIUM ACETATE

>A>		TABLET; ORAL					
>A>		CALCIUM ACETATE					
>A>		+ ROXANE	EQ 169MG CALCIUM	N77693 001	Jan 30, 2008	Jan	NEWA

CEFADROXIL/CEFADROXIL HEMIHYDRATECAPSULE; ORAL
CEFADROXIL

>A>	AB	HIKMA	EQ 500MG BASE	N65311 001	Feb 07, 2006	Jan	CAHN
>D>	AB	WESTWARD	EQ 500MG BASE	N65311 001	Feb 07, 2006	Jan	CAHN

CEFTRIAZONE SODIUMINJECTABLE; IM-IV
CEFTRIAZONE

>A>	AP	LUITPOLD	EQ 250MG BASE/VIAL	N65305 001	Jan 11, 2008	Jan	NEWA
>A>	AP		EQ 500MG BASE/VIAL	N65305 002	Jan 11, 2008	Jan	NEWA
>A>	AP		EQ 1GM BASE/VIAL	N65305 003	Jan 11, 2008	Jan	NEWA
>A>	AP		EQ 2GM BASE/VIAL	N65305 004	Jan 11, 2008	Jan	NEWA

CEFUROXIME AXETILFOR SUSPENSION; ORAL
CEFTIN

>D>		GLAXOSMITHKLINE	EQ 125MG BASE/5ML	N50672 001	Jun 30, 1994	Jan	CFTG
>A>	AB		EQ 125MG BASE/5ML	N50672 001	Jun 30, 1994	Jan	CFTG
>D>		+	EQ 250MG BASE/5ML	N50672 002	Apr 29, 1997	Jan	CFTG
>A>	AB	+	EQ 250MG BASE/5ML	N50672 002	Apr 29, 1997	Jan	CFTG
>A>		CEFUROXIME AXETIL					
>A>	AB	RANBAXY	EQ 125MG BASE/5ML	N65323 001	Feb 05, 2008	Jan	NEWA
>A>	AB		EQ 250MG BASE/5ML	N65323 002	Feb 05, 2008	Jan	NEWA

TABLET; ORAL

CEFUROXIME AXETIL

>A>	AB	ORCHID HLTHCARE	EQ 125MG BASE	N65359 001	Feb 15, 2008	Jan	NEWA
>A>	AB		EQ 250MG BASE	N65359 002	Feb 15, 2008	Jan	NEWA
>A>	AB		EQ 500MG BASE	N65359 003	Feb 15, 2008	Jan	NEWA

CETIRIZINE HYDROCHLORIDE

SYRUP; ORAL
ZYRTEC

>A>	+	MCNEIL CONSUMER	5MG/5ML	N20346 001	Sep 27, 1996	Jan	CAHN
>D>	+	PFIZER	5MG/5ML	N20346 001	Sep 27, 1996	Jan	CAHN

CICLESONIDE

>A>		AEROSOL, METERED; INHALATION					
>A>		ALVESCO					
>A>		NYCOMED US	0.08MG/INH	N21658 002	Jan 10, 2008	Jan	NEWA
>A>	+		0.16MG/INH	N21658 003	Jan 10, 2008	Jan	NEWA

CIPROFLOXACIN HYDROCHLORIDE

SOLUTION/DROPS; OPHTHALMIC
CIPROFLOXACIN HYDROCHLORIDE

>A>	AT	PHARMAFORCE	EQ 0.3% BASE	N78598 001	Jan 16, 2008	Jan	NEWA
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CLOPIDOGREL BISULFATE

TABLET; ORAL
CLOPIDOGREL BISULFATE

>A>	AB	DR REDDYS LABS INC	EQ 75MG BASE	N76273 001	Jan 14, 2008	Jan	NEWA
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CYCLOPHOSPHAMIDE

INJECTABLE; INJECTION
CYCLOPHOSPHAMIDE

>D>	AP	BAXTER HLTHCARE	100MG/VIAL	N88371 001	Jul 03, 1986	Jan	DISC
>A>		@	100MG/VIAL	N88371 001	Jul 03, 1986	Jan	DISC
>D>	AP		200MG/VIAL	N88372 001	Jul 03, 1986	Jan	DISC
>A>		@	200MG/VIAL	N88372 001	Jul 03, 1986	Jan	DISC
>D>	AP		500MG/VIAL	N88373 001	Jul 03, 1986	Jan	DISC
>A>		@	500MG/VIAL	N88373 001	Jul 03, 1986	Jan	DISC
>D>	AP		1GM/VIAL	N88374 001	Sep 24, 1986	Jan	DISC
>A>		@	1GM/VIAL	N88374 001	Sep 24, 1986	Jan	DISC

LYOPHILIZED CYTOXAN

>D>	AP	+	BRISTOL MYERS SQUIBB	100MG/VIAL	N12142 006	Dec 05, 1985	Jan	CTEC
>A>		+		100MG/VIAL	N12142 006	Dec 05, 1985	Jan	CTEC
>D>	AP	+		200MG/VIAL	N12142 007	Dec 10, 1985	Jan	CTEC
>A>		+		200MG/VIAL	N12142 007	Dec 10, 1985	Jan	CTEC
>D>	AP	+		500MG/VIAL	N12142 008	Jan 04, 1984	Jan	CTEC
>A>		+		500MG/VIAL	N12142 008	Jan 04, 1984	Jan	CTEC
>D>	AP	+		1GM/VIAL	N12142 010	Sep 24, 1985	Jan	CTEC
>A>		+		1GM/VIAL	N12142 010	Sep 24, 1985	Jan	CTEC
>D>	AP	+		2GM/VIAL	N12142 009	Dec 10, 1984	Jan	CTEC
>A>		+		2GM/VIAL	N12142 009	Dec 10, 1984	Jan	CTEC

NEOSAR

>D>	AP		TEVA PARENTERAL	100MG/VIAL	N87442 001	Feb 16, 1982	Jan	DISC
>A>		@		100MG/VIAL	N87442 001	Feb 16, 1982	Jan	DISC
>D>	AP			200MG/VIAL	N87442 002	Feb 16, 1982	Jan	DISC
>A>		@		200MG/VIAL	N87442 002	Feb 16, 1982	Jan	DISC
>D>	AP			500MG/VIAL	N87442 003	Feb 16, 1982	Jan	DISC
>A>		@		500MG/VIAL	N87442 003	Feb 16, 1982	Jan	DISC
>D>	AP			1GM/VIAL	N87442 004	Jul 08, 1983	Jan	DISC
>A>		@		1GM/VIAL	N87442 004	Jul 08, 1983	Jan	DISC
>D>	AP			2GM/VIAL	N87442 005	Mar 30, 1989	Jan	DISC

INJECTABLE; INJECTION

>D>		NEOSAR							
>A>		@ TEVA PARENTERAL	2GM/VIAL	N87442	005	Mar 30, 1989	Jan	DISC	
		TABLET; ORAL							
		CYCLOPHOSPHAMIDE							
>D>	AB	ROXANE	25MG	N40032	001	Aug 17, 1999	Jan	CTEC	
>A>			25MG	N40032	001	Aug 17, 1999	Jan	CTEC	
>D>	AB		50MG	N40032	002	Aug 17, 1999	Jan	CRLD	
>A>		+	50MG	N40032	002	Aug 17, 1999	Jan	CRLD	
>D>		CYTOXAN							
>D>	AB	BRISTOL MYERS SQUIBB	25MG	N12141	002		Jan	DISC	
>A>		@	25MG	N12141	002		Jan	DISC	
>D>	AB	+	50MG	N12141	001		Jan	DISC	
>A>		@	50MG	N12141	001		Jan	DISC	

DEXTROAMPHETAMINE SULFATE

>A>		SOLUTION; ORAL							
>A>		DEXTROAMPHETAMINE SULFATE							
>A>		+	OUTLOOK PHARMS	5MG/5ML	N40776	001	Jan 29, 2008	Jan	NEWA

DICLOFENAC SODIUM

		SOLUTION/DROPS; OPHTHALMIC							
		DICLOFENAC SODIUM							
>A>	AT	ALCON	0.1%	N78031	001	Feb 06, 2008	Jan	NEWA	

DICLOXACILLIN SODIUM

>D>		FOR SUSPENSION; ORAL							
>D>		DICLOXACILLIN SODIUM							
>D>		+	APOTHECON	EQ 62.5MG BASE/5ML	N61455	001		Jan	DISC
>A>		@		EQ 62.5MG BASE/5ML	N61455	001		Jan	DISC

DIPYRIDAMOLE

		TABLET; ORAL							
		DIPYRIDAMOLE							
>A>	AB	ZYDUS PHARMS USA INC	25MG	N40874	001	Jan 28, 2008	Jan	NEWA	
>A>	AB		50MG	N40874	002	Jan 28, 2008	Jan	NEWA	
>A>	AB		75MG	N40874	003	Jan 28, 2008	Jan	NEWA	

EDETATE DISODIUM

		INJECTABLE; INJECTION							
		EDETATE DISODIUM							
>D>	AP	APOTEX INC	150MG/ML	N40376	001	Nov 04, 2002	Jan	DISC	
>A>		@	150MG/ML	N40376	001	Nov 04, 2002	Jan	DISC	

ESTRADIOL

		GEL, METERED; TRANSDERMAL							
		ELESTRIN							
>D>	BX	+	BRADLEY PHARMS	0.06% (0.87GM/ACTIVATION)	N21813	001	Dec 15, 2006	Jan	CTEC
>A>		+		0.06% (0.87GM/ACTIVATION)	N21813	001	Dec 15, 2006	Jan	CTEC
		ESTROGEL							
>D>	BX	+	ASCEND	0.06% (1.25GM/ACTIVATION)	N21166	002	Feb 09, 2004	Jan	CTEC
>A>		+		0.06% (1.25GM/ACTIVATION)	N21166	002	Feb 09, 2004	Jan	CTEC

ETHAMBUTOL HYDROCHLORIDE

TABLET; ORAL

MYAMBUTOL

>D>	AB	STAT TRADE	100MG	N16320 001	Jan	CAHN
>D>		@	200MG	N16320 002	Jan	CAHN
>D>	AB		400MG	N16320 003	Jan	CAHN
>D>		@	500MG	N16320 004	Jan	CAHN
>A>	AB	STAT-TRADE	100MG	N16320 001	Jan	CAHN
>A>		@	200MG	N16320 002	Jan	CAHN
>A>	AB		400MG	N16320 003	Jan	CAHN
>A>		@	500MG	N16320 004	Jan	CAHN

>A> ETRAVIRINE

>A> TABLET; ORAL

>A> INTELENCE

>A>	+	TIBOTEC	100MG	N22187 001	Jan 18, 2008	Jan	NEWA
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FLUOXETINE HYDROCHLORIDE

CAPSULE; ORAL

FLUOXETINE

>A>	AB	AUROBINDO PHARMA	EQ 10MG BASE	N78619 001	Jan 31, 2008	Jan	NEWA
>A>	AB		EQ 20MG BASE	N78619 002	Jan 31, 2008	Jan	NEWA
>A>	AB		EQ 40MG BASE	N78619 003	Jan 31, 2008	Jan	NEWA
>A>	AB	WOCKHARDT	EQ 10MG BASE	N78143 001	Jan 16, 2008	Jan	NEWA
>A>	AB		EQ 20MG BASE	N78143 002	Jan 16, 2008	Jan	NEWA
>A>	AB		EQ 40MG BASE	N78143 003	Jan 16, 2008	Jan	NEWA

FLUTICASONE PROPIONATE

SPRAY, METERED; NASAL

FLUTICASONE PROPIONATE

>A>	AB	HI TECH PHARMA	0.05MG/SPRAY	N77570 001	Jan 16, 2008	Jan	NEWA
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>A> FOSAPREPITANT DIMEGLUMINE

>A> POWDER; INTRAVENOUS

>A> EMEND

>A>	+	MERCK AND CO INC	EQ 115MG BASE/VIAL	N22023 001	Jan 25, 2008	Jan	NEWA
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GABAPENTIN

CAPSULE; ORAL

GABAPENTIN

>A>	AB	AUROBINDO PHARM	100MG	N78787 001	Jan 31, 2008	Jan	NEWA
>A>	AB		300MG	N78787 002	Jan 31, 2008	Jan	NEWA
>A>	AB		400MG	N78787 003	Jan 31, 2008	Jan	NEWA

GLIPIZIDE; METFORMIN HYDROCHLORIDE

TABLET; ORAL

GLIPIZIDE AND METFORMIN HYDROCHLORIDE

>A>	AB	CARACO	2.5MG;250MG	N77620 001	Jan 11, 2008	Jan	NEWA
>A>	AB		2.5MG;500MG	N77620 002	Jan 11, 2008	Jan	NEWA
>A>	AB		5MG;500MG	N77620 003	Jan 11, 2008	Jan	NEWA

GRANISETRON HYDROCHLORIDE

TABLET; ORAL

GRANISETRON HYDROCHLORIDE

>A>	AB	MYLAN	EQ 1MG BASE	N78725 001	Jan 30, 2008	Jan	NEWA
>A>	AB	ORCHID HLTHCARE	EQ 1MG BASE	N78678 001	Feb 13, 2008	Jan	NEWA

HEPARIN SODIUM

INJECTABLE; INJECTION

HEPARIN SODIUM PRESERVATIVE FREE

>D>	AP	+	HOSPIRA	2,500 UNITS/ML	N05264 014	Apr 07, 1986	Jan	CTEC
>A>		+		2,500 UNITS/ML	N05264 014	Apr 07, 1986	Jan	CTEC

HOMATROPINE METHYLBROMIDE; HYDROCODONE BITARTRATE

SYRUP; ORAL

HYDROCODONE BITARTRATE AND HOMATROPINE METHYLBROMIDE

>A>	AA	HI TECH PHARMA	1.5MG/5ML;5MG/5ML	N40613 001	Feb 08, 2008	Jan	NEWA
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HYDROCHLOROTHIAZIDE

CAPSULE; ORAL

HYDROCHLOROTHIAZIDE

>A>	AB	CADISTA PHARMS	12.5MG	N78391 001	Feb 11, 2008	Jan	NEWA
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HYDROXYCHLOROQUINE SULFATE

TABLET; ORAL

HYDROXYCHLOROQUINE SULFATE

>D>		@	IPCA	200MG	N40766 001	Jun 14, 2007	Jan	CMFD
>A>	AB		IPCA LABS LTD	200MG	N40766 001	Jun 14, 2007	Jan	CMFD

HYDROXYUREA

>D>			TABLET; ORAL					
>D>			HYDROXYUREA					
>D>		+	BARR	1GM	N75734 001	Aug 29, 2000	Jan	DISC
>A>		@		1GM	N75734 001	Aug 29, 2000	Jan	DISC

IMIPRAMINE PAMOATE

CAPSULE; ORAL

TOFRANIL-PM

>D>			TYCO HLTHCARE	EQ 75MG HCL	N17090 001		Jan	CRLD
>A>		+		EQ 75MG HCL	N17090 001		Jan	CRLD
>D>		+		EQ 150MG HCL	N17090 002		Jan	CRLD
>A>				EQ 150MG HCL	N17090 002		Jan	CRLD

INDOMETHACIN

CAPSULE; ORAL

INDOMETHACIN

>D>		@	IVAX PHARMS	25MG	N70719 001	Feb 12, 1986	Jan	CMFD
>A>	AB			25MG	N70719 001	Feb 12, 1986	Jan	CMFD
>D>		@		50MG	N70756 001	Feb 12, 1986	Jan	CMFD
>A>	AB			50MG	N70756 001	Feb 12, 1986	Jan	CMFD

ISOSORBIDE DINITRATE

TABLET; SUBLINGUAL

ISOSORBIDE DINITRATE

>D>	AB	WATSON LABS	5MG	N86031 001	Sep 29, 1987	Jan	CRLD
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TABLET; SUBLINGUAL
ISOSORBIDE DINITRATE

>A> AB + WATSON LABS 5MG N86031 001 Sep 29, 1987 Jan CRLD

KETOROLAC TROMETHAMINE

INJECTABLE; INJECTION
KETOROLAC TROMETHAMINE

>A> AP LUITPOLD 15MG/ML N78145 001 Jan 14, 2008 Jan NEWA

>A> AP 30MG/ML N78145 002 Jan 14, 2008 Jan NEWA

LEVOCETIRIZINE DIHYDROCHLORIDE

>A> SOLUTION; ORAL

>A> XYZAL

>A> + UCB INC 2.5MG/5ML N22157 001 Jan 28, 2008 Jan NEWA

LEVONORDEFRIN; MEPIVACAINE HYDROCHLORIDE

INJECTABLE; INJECTION
POLOCAINE W/ LEVONORDEFRIN

>D> AP DENTSPLY PHARM 0.05MG/ML;2% N89517 001 Apr 14, 1988 Jan CRLD

>A> AP + 0.05MG/ML;2% N89517 001 Apr 14, 1988 Jan CRLD

LEVOTHYROXINE SODIUM**

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

TABLET; ORAL

LEVOXYL

>D> AB1, KING PHARMS 0.2MG N21301 011 May 25, 2001 Jan CRLD
AB3

>A> AB1, + 0.2MG N21301 011 May 25, 2001 Jan CRLD
AB3

>D> AB1, + 0.3MG N21301 012 May 25, 2001 Jan DISC
AB3

>A> @ 0.3MG N21301 012 May 25, 2001 Jan DISC

LITHIUM CARBONATE

CAPSULE; ORAL

ESKALITH

>D> AB JDS PHARMS 300MG N16860 001 Jan CAHN

>A> AB NOVEN THERAP 300MG N16860 001 Jan CAHN

TABLET, EXTENDED RELEASE; ORAL

LITHOBID

>D> AB + JDS PHARMS 300MG N18027 001 Jan CAHN

>A> AB + NOVEN THERAP 300MG N18027 001 Jan CAHN

MINOXIDIL

TABLET; ORAL

LONITEN

>D> AB PHARMACIA AND UPJOHN 2.5MG N18154 001 Jan DISC

>A> @ 2.5MG N18154 001 Jan DISC

>D> AB + 10MG N18154 003 Jan DISC

>A> @ 10MG N18154 003 Jan DISC

NIMODIPINE

CAPSULE; ORAL

NIMODIPINE

>A> AB BANNER PHARMACAPS 30MG N76740 001 Jan 17, 2008 Jan NEWA

NISOLDIPINE

TABLET, EXTENDED RELEASE; ORAL
SULAR

>A>	+	SCIELE PHARMA INC	8.5MG	N20356 008	Jan 02, 2008	Jan	NEWA
>A>	+		17MG	N20356 007	Jan 02, 2008	Jan	NEWA
>A>			25.5MG	N20356 006	Jan 02, 2008	Jan	NEWA
>A>	+		34MG	N20356 005	Jan 02, 2008	Jan	NEWA

>D> NITROFUZAZONE

OINTMENT; TOPICAL
NITROFUZAZONE

>D>	+	TARO	0.2%	N86156 001		Jan	DISC
>A>	@		0.2%	N86156 001		Jan	DISC

NYSTATIN

TABLET; ORAL
NYSTATIN

>D>	AA	TEVA	500,000 UNITS	N62506 001	Jan 16, 1984	Jan	CRLD
>A>	AA	+	500,000 UNITS	N62506 001	Jan 16, 1984	Jan	CRLD

OCTREOTIDE ACETATE

INJECTABLE; INJECTION
OCTREOTIDE ACETATE

>D>	AP	BEDFORD	EQ 0.2MG BASE/ML	N76330 001	Apr 08, 2005	Jan	CRLD
>A>	AP	+	EQ 0.2MG BASE/ML	N76330 001	Apr 08, 2005	Jan	CRLD
>D>	AP		EQ 1MG BASE/ML	N76330 002	Apr 08, 2005	Jan	CRLD
>A>	AP	+	EQ 1MG BASE/ML	N76330 002	Apr 08, 2005	Jan	CRLD

OXYCODONE HYDROCHLORIDE

TABLET, EXTENDED RELEASE; ORAL
OXYCONTIN

>D>	@	PURDUE PHARMA LP	15MG	N20553 006	Sep 18, 2006	Jan	CMFD
>A>			15MG	N20553 006	Sep 18, 2006	Jan	CMFD
>D>	@		30MG	N20553 007	Sep 18, 2006	Jan	CMFD
>A>			30MG	N20553 007	Sep 18, 2006	Jan	CMFD
>D>	@		60MG	N20553 008	Sep 18, 2006	Jan	CMFD
>A>			60MG	N20553 008	Sep 18, 2006	Jan	CMFD

OXYTOCIN

INJECTABLE; INJECTION
OXYTOCIN

>D>	+	APP PHARMS	100USP UNITS/10 ML (10USP UNITS/ML)	N18248 002		Jan	CFTG
>A>	AP	+	100USP UNITS/10 ML (10USP UNITS/ML)	N18248 002		Jan	CFTG
>A>	AP	TEVA PARENTERAL	10USP UNITS/ML (10USP UNITS/ML)	N77453 001	Jan 24, 2008	Jan	NEWA
>A>	AP		100USP UNITS/10ML (10USP UNITS/ML)	N77453 002	Jan 24, 2008	Jan	NEWA

PALIPERIDONE

TABLET, EXTENDED RELEASE; ORAL
INVEGA

>D>		JANSSEN LP	6MG	N21999 002	Dec 19, 2006	Jan	CRLD
>A>	+		6MG	N21999 002	Dec 19, 2006	Jan	CRLD
>D>	+		9MG	N21999 003	Dec 19, 2006	Jan	CRLD
>A>			9MG	N21999 003	Dec 19, 2006	Jan	CRLD

PHENDIMETRAZINE TARTRATE

TABLET; ORAL

PHENDIMETRAZINE TARTRATE

>A>	AA	AMIDE PHARM	35MG	N40762 001	Jan 28, 2008	Jan	NEWA
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PHENYTOIN SODIUM

CAPSULE; ORAL

EXTENDED PHENYTOIN SODIUM

>A>	AB	WOCKHARDT USA	100MG EXTENDED	N40732 001	Jan 30, 2008	Jan	NEWA
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POTASSIUM CHLORIDE

TABLET, EXTENDED RELEASE; ORAL

>D>		K-DUR 10					
>D>	AB	KEY PHARMS	10MEQ	N19439 002	Jun 13, 1986	Jan	CTNA
>D>		K-DUR 20					
>D>	AB	+ KEY PHARMS	20MEQ	N19439 001	Jun 13, 1986	Jan	CTNA
>A>		POTASSIUM CHLORIDE					
>A>	AB	SCHERING	10MEQ	N19439 002	Jun 13, 1986	Jan	CTNA
>A>	AB	+	20MEQ	N19439 001	Jun 13, 1986	Jan	CTNA

PRAVASTATIN SODIUM

TABLET; ORAL

PRAVASTATIN SODIUM

>A>	AB	LEK PHARMS DD	80MG	N77491 001	Feb 11, 2008	Jan	NEWA
>A>	AB	TEVA PHARMS	80MG	N77793 001	Jan 15, 2008	Jan	NEWA

PREDNISOLONE ACETATE

>A>		SUSPENSION; ORAL					
>A>		FLO-PRED					
>A>		TARO	EQ 5MG BASE/5ML	N22067 001	Jan 17, 2008	Jan	NEWA
>A>		+	EQ 15MG BASE/5ML	N22067 002	Jan 17, 2008	Jan	NEWA

PRIMIDONE

TABLET; ORAL

PRIMIDONE

>A>	AB	IMPAX LABS	50MG	N40717 001	Feb 12, 2008	Jan	NEWA
>A>	AB		250MG	N40717 002	Feb 12, 2008	Jan	NEWA

PROCHLORPERAZINE EDISYLATE

INJECTABLE; INJECTION

PROCHLORPERAZINE EDISYLATE

>D>	AP	TEVA PARENTERAL	EQ 5MG BASE/ML	N40505 001	May 30, 2003	Jan	DISC
>A>		@	EQ 5MG BASE/ML	N40505 001	May 30, 2003	Jan	DISC

PROMETHAZINE HYDROCHLORIDE

TABLET; ORAL

PROMETHAZINE HYDROCHLORIDE

>A>	AB	IMPAX LABS	12.5MG	N40724 001	Feb 12, 2008	Jan	NEWA
>A>	AB		25MG	N40724 002	Feb 12, 2008	Jan	NEWA

PROPRANOLOL HYDROCHLORIDE

INJECTABLE; INJECTION

PROPRANOLOL HYDROCHLORIDE

>A>	AP	HIKMA FARMACEUTICA	1MG/ML	N77760 001	Jan 31, 2008	Jan	NEWA
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SERTRALINE HYDROCHLORIDE

TABLET; ORAL

SERTRALINE HYDROCHLORIDE

>A>	AB	MATRIX LABS LTD	EQ 25MG BASE	N78626 001	Jan 31, 2008	Jan	NEWA
>A>	AB		EQ 50MG BASE	N78626 002	Jan 31, 2008	Jan	NEWA
>A>	AB		EQ 100MG BASE	N78626 003	Jan 31, 2008	Jan	NEWA

SOMATROPIN RECOMBINANT

INJECTABLE; INJECTION

ACCRETROPIN

>A>	+	CANGENE	5MG/ML (5MG/ML)	N21538 001	Jan 23, 2008	Jan	NEWA
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NORDITROPIN

>D>		NOVO NORDISK INC	5MG/1.5ML	N21148 001	Jun 20, 2000	Jan	CTEC
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>A>	BX		5MG/1.5ML	N21148 001	Jun 20, 2000	Jan	CTEC
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NORDITROPIN NORDIFLEX

>D>		NOVO NORDISK INC	5MG/1.5ML	N21148 004	Oct 01, 2004	Jan	CTEC
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>A>	BX		5MG/1.5ML	N21148 004	Oct 01, 2004	Jan	CTEC
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NUTROPIN AQ

>A>	+	GENENTECH	5MG/2ML (2.5MG/ML)	N20522 003	Jan 03, 2008	Jan	NEWA
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>D>	+		5MG/ML	N20522 001	Dec 29, 1995	Jan	CPOT
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>A>	+		10MG/2ML (5MG/ML)	N20522 001	Dec 29, 1995	Jan	CPOT
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>A>	+		20MG/2ML (10MG/ML)	N20522 004	Jan 03, 2008	Jan	NEWA
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NUTROPIN AQ PEN

>D>	BX	+	GENENTECH	5MG/ML	N20522 002	Apr 22, 2002	Jan	CPOT
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>A>	+		10MG/2ML (5MG/ML)	N20522 002	Apr 22, 2002	Jan	CPOT
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OMNITROPE

>A>	BX	SANDOZ	5MG/1.5ML	N21426 003	Jan 16, 2008	Jan	NEWA
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TEV-TROPIN

>D>	BX	+	FERRING	5MG/ML	N19774 002	Jan 04, 2002	Jan	CPOT
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>A>	BX	+		5MG/VIAL	N19774 002	Jan 04, 2002	Jan	CPOT
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TADALAFIL

TABLET; ORAL

CIALIS

>A>		LILLY	2.5MG	N21368 004	Jan 07, 2008	Jan	NEWA
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TECHNETIUM TC-99M MEBROFENIN KIT

INJECTABLE; INJECTION

CHOLETEC

>D>	+	BRACCO	N/A	N18963 001	Jan 21, 1987	Jan	CFTG
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>A>	AP	+		N/A	N18963 001	Jan 21, 1987	Jan	CFTG
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TECHNETIUM TC-99M MEBROFENIN

>A>	AP	CIS	N/A	N78242 001	Jan 29, 2008	Jan	NEWA
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VARDENAFIL HYDROCHLORIDE

TABLET; ORAL

LEVITRA

>A>		BAYER HLTHCARE	2.5MG	N21400 003	Aug 19, 2003	Jan	CAHN
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>A>			5MG	N21400 001	Aug 19, 2003	Jan	CAHN
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>A>			10MG	N21400 002	Aug 19, 2003	Jan	CAHN
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>A>	+		20MG	N21400 004	Aug 19, 2003	Jan	CAHN
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>D>		BAYER PHARMS	2.5MG	N21400 003	Aug 19, 2003	Jan	CAHN
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>D>			5MG	N21400 001	Aug 19, 2003	Jan	CAHN
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>D>			10MG	N21400 002	Aug 19, 2003	Jan	CAHN
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TABLET; ORAL

LEVITRA

>D> + BAYER PHARMS 20MG N21400 004 Aug 19, 2003 Jan CAHN

VINORELBINE TARTRATE

INJECTABLE; INJECTION

VINORELBINE TARTRATE

>A> AP EBEWE PHARMA EQ 10MG BASE/ML N78408 001 Feb 13, 2008 Jan NEWA

ZIDOVUDINE

TABLET; ORAL

ZIDOVUDINE

>A> AB MATRIX LABS LTD 300MG N78922 001 Feb 14, 2008 Jan NEWA

OTC DRUG PRODUCT LIST - 28TH EDITION

OTC DRUG PRODUCT LIST - CUMULATIVE SUPPLEMENT 1 - January 2008

2-1

CALCIUM CARBONATE; FAMOTIDINE; MAGNESIUM HYDROXIDE

TABLET, CHEWABLE; ORAL

CALCIUM CARBONATE, FAMOTIDINE AND MAGNESIUM HYDROXIDE

>A> PERRIGO R AND D 800MG;10MG;165MG N77355 001 Feb 06, 2008 Jan NEWA

CETIRIZINE HYDROCHLORIDE

SYRUP; ORAL

CHILDREN'S ZYRTEC ALLERGY

>A> + MCNEIL CONSUMER 1MG/ML N22155 002 Nov 16, 2007 Jan CAHN

>D> + PFIZER 1MG/ML N22155 002 Nov 16, 2007 Jan CAHN

CHILDREN'S ZYRTEC HIVES RELIEF

>A> + MCNEIL CONSUMER 1MG/ML N22155 001 Nov 16, 2007 Jan CAHN

>D> + PFIZER 1MG/ML N22155 001 Nov 16, 2007 Jan CAHN

TABLET, CHEWABLE; ORAL

CETIRIZINE HYDROCHLORIDE ALLERGY

>A> SANDOZ 5MG N78692 001 Feb 14, 2008 Jan NEWA

>A> 10MG N78692 002 Feb 14, 2008 Jan NEWA

TABLET; ORAL

CETIRIZINE HYDROCHLORIDE ALLERGY

>A> DR REDDYS LABS LTD 5MG N78343 004 Jan 15, 2008 Jan NEWA

>A> 10MG N78343 003 Jan 15, 2008 Jan NEWA

CETIRIZINE HYDROCHLORIDE HIVES

>A> DR REDDYS LABS LTD 5MG N78343 001 Jan 15, 2008 Jan NEWA

>A> 10MG N78343 002 Jan 15, 2008 Jan NEWA

ZYRTEC ALLERGY

>A> MCNEIL CONSUMER 5MG N19835 003 Nov 16, 2007 Jan CAHN

>A> + 10MG N19835 004 Nov 16, 2007 Jan CAHN

>D> PFIZER 5MG N19835 003 Nov 16, 2007 Jan CAHN

>D> + 10MG N19835 004 Nov 16, 2007 Jan CAHN

ZYRTEC HIVES RELIEF

>A> MCNEIL CONSUMER 5MG N19835 005 Nov 16, 2007 Jan CAHN

>A> + 10MG N19835 006 Nov 16, 2007 Jan CAHN

>D> PFIZER 5MG N19835 005 Nov 16, 2007 Jan CAHN

>D> + 10MG N19835 006 Nov 16, 2007 Jan CAHN

CETIRIZINE HYDROCHLORIDE; PSEUDOEPHEDRINE HYDROCHLORIDE

TABLET, EXTENDED RELEASE; ORAL

ZYRTEC-D 12 HOUR

>A> + MCNEIL 5MG;120MG N21150 002 Nov 09, 2007 Jan CAHN

>D> + PFIZER 5MG;120MG N21150 002 Nov 09, 2007 Jan CAHN

POTASSIUM IODIDE

TABLET; ORAL

THYROSAFE

>D> + R R REGISTRATIONS 65MG N76350 001 Sep 10, 2002 Jan CAHN

>A> + RECIP 65MG N76350 001 Sep 10, 2002 Jan CAHN

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

CUMULATIVE SUPPLEMENT NUMBER 01 JANUARY 2008

NO JANUARY 2008 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 JANUARY 2008 ADDITIONS

PATENT & EXCLUSIVITY DRUG PRODUCT LIST - CUMULATIVE SUPPLEMENT 1 - January 2008

See List footnote for information regarding List content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>ADEFOVIR DIPIVOXIL - HEPSERA</u>					
021449	001			>A> NPP	Dec 19, 2010
<u>ALISKIREN HEMIFUMARATE; HYDROCHLOROTHIAZIDE - TEKTRUNA HCT</u>					
022107	001			>A> NCE >A> NC	Mar 05, 2012 Jan 18, 2011
<u>ALISKIREN HEMIFUMARATE; HYDROCHLOROTHIAZIDE - TEKTRUNA HCT</u>					
022107	002			>A> NCE >A> NC	Mar 05, 2012 Jan 18, 2011
<u>ALISKIREN HEMIFUMARATE; HYDROCHLOROTHIAZIDE - TEKTRUNA HCT</u>					
022107	003			>A> NCE >A> NC	Mar 05, 2012 Jan 18, 2011
<u>ALISKIREN HEMIFUMARATE; HYDROCHLOROTHIAZIDE - TEKTRUNA HCT</u>					
022107	004			>A> NCE >A> NC	Mar 05, 2012 Jan 18, 2011
<u>ARFORMOTEROL TARTRATE - BROVANA</u>					
021912	001	>A> 6040344	Nov 12, 2016	DS	
		>A> 6472563	Nov 09, 2021	DS	
		>A> 6720453	Nov 09, 2021	DS	
		>A> 7145036	Nov 09, 2021	DS	
<u>CARVEDILOL - COREG</u>					
020297	001	>A> RE40000	Jun 07, 2015	U-233	
		>A> RE40000*PED	Dec 07, 2015		
<u>CARVEDILOL - COREG</u>					
020297	002	>A> RE40000	Jun 07, 2015	U-233	
		>A> RE40000*PED	Dec 07, 2015		
<u>CARVEDILOL - COREG</u>					
020297	003	>A> RE40000	Jun 07, 2015	U-233	
		>A> RE40000*PED	Dec 07, 2015		
<u>CARVEDILOL - COREG</u>					
020297	004	>A> RE40000	Jun 07, 2015	U-233	
		>A> RE40000*PED	Dec 07, 2015		
<u>CARVEDILOL PHOSPHATE - COREG CR</u>					
022012	001	>A> RE40000	Jun 07, 2015	U-777	
		>A> RE40000*PED	Dec 07, 2015		
<u>CARVEDILOL PHOSPHATE - COREG CR</u>					
022012	002	>A> RE40000	Jun 07, 2015	U-777	
		>A> RE40000*PED	Dec 07, 2015		
<u>CARVEDILOL PHOSPHATE - COREG CR</u>					
022012	003	>A> RE40000	Jun 07, 2015	U-777	
		>A> RE40000*PED	Dec 07, 2015		
<u>CARVEDILOL PHOSPHATE - COREG CR</u>					
022012	004	>A> RE40000	Jun 07, 2015	U-777	
		>A> RE40000*PED	Dec 07, 2015		
<u>COLESEVELAM HYDROCHLORIDE - WELCHOL</u>					
021176	001			>A> I-553	Jan 18, 2011
<u>DARIFENACIN HYDROBROMIDE - ENABLEX</u>					
021513	001	>A> 5096890	Mar 13, 2015	DS DP	U-631
<u>DARIFENACIN HYDROBROMIDE - ENABLEX</u>					
021513	002	>A> 5096890	Mar 13, 2015	DS DP	U-631
<u>EPLERENONE - INSPRA</u>					
021437	001			>A> M-72 >A> PED	Jan 31, 2011 Jul 31, 2011

PATENT & EXCLUSIVITY DRUG PRODUCT LIST - CUMULATIVE SUPPLMENT 1 - January 2008

See List footnote for information regarding List content

APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>EPLERENONE - INSPRA</u>					
021437 002				>A> M-72 >A> PED	Jan 31, 2011 Jul 31, 2011
<u>ESTRADIOL; NORGESTIMATE - PREFEST</u>					
021040 001	>A> 7320970	Mar 30, 2020	DP U-844		
<u>ETHINYL ESTRADIOL; LEVONORGESTREL - SEASONIQUE</u>					
021840 001	>A> 7320969	Jan 30, 2024	U-828		
<u>ETRAVIRINE - INTELENCE</u>					
022187 001				>A> NCE	Jan 18, 2013
<u>FENOFIBRATE - TRICOR</u>					
021656 001	>A> 7320802	Feb 21, 2023	U-847		
<u>FENOFIBRATE - TRICOR</u>					
021656 002	>A> 7320802	Feb 21, 2023	U-847		
<u>FENTANYL CITRATE - FENTORA</u>					
021947 006	>A> 6200604	Mar 26, 2019	U-767		
	>A> 6974590	Mar 26, 2019	U-767		
<u>GLIMEPIRIDE; ROSIGLITAZONE MALEATE - AVANDARYL</u>					
021700 004	>A> 5002953	Sep 17, 2011	DS DP U-840		
<u>GLIMEPIRIDE; ROSIGLITAZONE MALEATE - AVANDARYL</u>					
021700 005	>A> 5002953	Sep 17, 2011	DS DP U-840		
<u>GRANISETRON HYDROCHLORIDE - GRANISETRON HYDROCHLORIDE</u>					
077177 001				>A> PC	Jun 28, 2008
<u>GRANISETRON HYDROCHLORIDE - GRANISETRON HYDROCHLORIDE PRESERVATIVE FREE</u>					
077165 001				>A> PC	Jun 28, 2008
<u>LISDEXAMFETAMINE DIMESYLATE - VYVANSE</u>					
021977 004	>A> 7105486	Jun 29, 2023	U-842		
	>A> 7223735	Jun 29, 2023	DP		
<u>LISDEXAMFETAMINE DIMESYLATE - VYVANSE</u>					
021977 005	>A> 7105486	Jun 29, 2023	U-842		
	>A> 7223735	Jun 29, 2023	DP		
<u>LISDEXAMFETAMINE DIMESYLATE - VYVANSE</u>					
021977 006	>A> 7105486	Jun 29, 2023	U-842		
	>A> 7223735	Jun 29, 2023	DP		
<u>MICAFUNGIN SODIUM - MYCAMINE</u>					
021506 002	>A> 6107458	Sep 29, 2015	DS DP U-845	>A> I-554	Jan 22, 2011
	>A> 6107458	Sep 29, 2015	DS DP U-650		
	>A> 6265536	Sep 29, 2015	DS DP U-845		
	>A> 6265536	Sep 29, 2015	DS DP U-650		
	>A> 6774104	Jan 08, 2021	DP U-845		
	>A> 6774104	Jan 08, 2021	DP U-650		
<u>MICAFUNGIN SODIUM - MYCAMINE</u>					
021506 003	>A> 5376634	Dec 27, 2011	DS DP	>A> I-554	Jan 22, 2011
	>A> 6107458	Sep 29, 2015	DS DP U-845	>A> NCE	Mar 16, 2010
	>A> 6107458	Sep 29, 2015	DS DP U-650		
	>A> 6265536	Sep 29, 2015	DS DP U-845		
	>A> 6265536	Sep 29, 2015	DS DP U-650		
	>A> 6774104	Jan 08, 2021	DP U-845		
	>A> 6774104	Jan 08, 2021	DP U-650		
<u>MODAFINIL - PROVIGIL</u>					
020717 001	>A> 7297346	Nov 29, 2023	DP		
<u>MODAFINIL - PROVIGIL</u>					
020717 002	>A> 7297346	Nov 29, 2023	DP		

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APPL/PROD NO	PATENT NO	PATENT EXPIRATION DATE	PATENT CODES	EXCLUSIVITY CODE(S)	EXCLUSIVITY EXPIRATION DATE
<u>NEBIVOLOL HYDROCHLORIDE - BYSTOLIC</u>					
021742 002	>A> 5759580	Jun 02, 2015	DP		
	>A> 6545040	Apr 08, 2020	DP	U-3	
<u>NEBIVOLOL HYDROCHLORIDE - BYSTOLIC</u>					
021742 003	>A> 5759580	Jun 02, 2015	DP		
	>A> 6545040	Apr 08, 2020	DP	U-3	
<u>NEBIVOLOL HYDROCHLORIDE - BYSTOLIC</u>					
021742 004	>A> 5759580	Jun 02, 2015	DP		
	>A> 6545040	Apr 08, 2020	DP	U-3	
<u>OXCARBAZEPINE - OXCARBAZEPINE</u>					
078069 001				>A> PC	Apr 06, 2008
<u>OXCARBAZEPINE - OXCARBAZEPINE</u>					
078069 002				>A> PC	Apr 06, 2008
<u>OXCARBAZEPINE - OXCARBAZEPINE</u>					
078069 003				>A> PC	Apr 06, 2008
<u>PANTOPRAZOLE SODIUM - PANTOPRAZOLE SODIUM</u>					
077056 001				>A> PC	Jun 18, 2008
<u>PANTOPRAZOLE SODIUM - PANTOPRAZOLE SODIUM</u>					
077056 002				>A> PC	Jun 18, 2008
<u>PANTOPRAZOLE SODIUM - PANTOPRAZOLE SODIUM</u>					
077058 001				>A> PC	Jun 18, 2008
<u>PANTOPRAZOLE SODIUM - PANTOPRAZOLE SODIUM</u>					
077058 002				>A> PC	Jun 18, 2008
<u>PEMETREXED DISODIUM - ALIMTA</u>					
021462 002	>A> 5217974	Mar 29, 2011		U-551	
<u>PRAMIPEXOLE DIHYDROCHLORIDE - MIRAPEX</u>					
020667 007	>A> 4886812	Mar 25, 2011	DS DP	>A> I-517	Nov 07, 2009
	>A> 6001861	Jan 16, 2018		U-784	
	>A> 6194445	Jan 16, 2018		U-784	
<u>SEVELAMER CARBONATE - RENVELA</u>					
022127 001	>A> 5667775	Sep 16, 2014		U-246	
<u>SOMATROPIN RECOMBINANT - ACCRETROPIN</u>					
021538 001				>A> NP	Jan 23, 2011
<u>SOMATROPIN RECOMBINANT - OMNITROPE</u>					
021426 003				>A> NP	May 30, 2009
<u>SUMATRIPTAN SUCCINATE - IMITREX STATDOSE</u>					
020080 002	>A> 5037845	Aug 06, 2008	DS DP	U-848	
	>A> 5037845*PED	Feb 06, 2009			
<u>TECHNETIUM TC-99M SESTAMIBI KIT - CARDIOLITE</u>					
019785 001	>A> 4988827	Jan 29, 2008			
	>A> 4988827*PED	Jul 29, 2008			
<u>TERBINAFINE - LAMISIL AT</u>					
021958 001	>A> 6121314	May 18, 2012		U-540	
	>A> 6121314	May 18, 2012		U-504	
<u>TESTOSTERONE - TESTIM</u>					
021454 001	>A> 7320968	Jan 18, 2025		U-843	
<u>TIOTROPIUM BROMIDE MONOHYDRATE - SPIRIVA</u>					
021395 001	>A> 5478578	Dec 26, 2012		DP	
	>A> 7309707	Mar 10, 2023	DS DP		
<u>TRIAMCINOLONE ACETONIDE - TRIESENCE</u>					
022048 001	>A> 6395294	Jan 13, 2020		DP	U-846

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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, 28th 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>