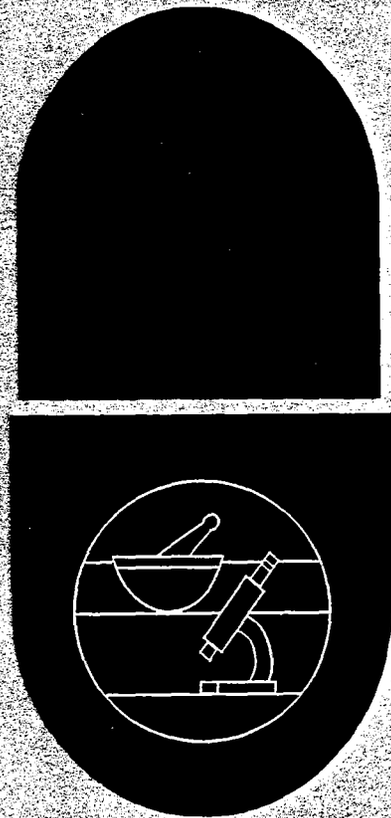


**CUMULATIVE  
SUPPLEMENT 12**

**JAN'94-DEC'94**



# **APPROVED DRUG PRODUCTS**

**WITH  
THERAPEUTIC EQUIVALENCE EVALUATIONS**

**14<sup>TH</sup> EDITION**

**U.S. DEPARTMENT OF HEALTH AND HUMAN SERVICES**

**PUBLIC HEALTH SERVICE**

**FOOD AND DRUG ADMINISTRATION**

**CENTER FOR DRUG EVALUATION AND RESEARCH**

**OFFICE OF MANAGEMENT**

**DIVISION OF DRUG INFORMATION RESOURCES**

**APPROVED DRUG PRODUCTS  
with  
THERAPEUTIC EQUIVALENCE EVALUATIONS**

**14TH EDITION**

**Cumulative Supplement 12**

**DECEMBER 1994**

**CONTENTS**

	<i><b>PAGE</b></i>
1.0 INTRODUCTION .....	iii
1.1 How to Use the Cumulative Supplement .....	iii
1.2 Products Requiring Revised Labeling for Full Approval .....	v
1.3 Applicant Name Changes .....	vi
1.4 New Indications for Previously Approved Drug Products .....	vii
1.5 USP Monograph Title Additions or Changes .....	ix
1.6 Report of Counts for the Prescription Drug Product List .....	x
2.0 DRUG PRODUCT LISTS .....	
2.1 Prescription Drug Product List .....	1
2.2 OTC Drug Product List .....	61
2.3 Drug Products with Approval under Section 505 of the Act Administered by the Center for Biologics Evaluation and Research List .....	63
2.4 Orphan Drug Product Designations .....	64
2.5 Drug Products Which Must Demonstrate in vivo Bioavailability Only if Product Fails to Achieve Adequate Dissolution .....	70
2.6 Biopharmaceutical Guidance Availability .....	71
2.7 ANDA Suitability Petitions .....	72
PATENT AND EXCLUSIVITY INFORMATION ADDENDUM	
A. Exclusivity Terms .....	74
B. Patent and Exclusivity Lists .....	76



**APPROVED DRUG PRODUCTS  
with  
THERAPEUTIC EQUIVALENCE EVALUATIONS**

**14TH EDITION**

**CUMULATIVE SUPPLEMENT 12**

**DECEMBER 1994**

**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, 14th Edition (the List). The List is composed of four parts: approved prescription drug products with therapeutic equivalence evaluations, over-the-counter (OTC) drug products that require approved applications as a condition of marketing, drug products with approval under Section 505 of the Act administered by the Center for Biologics Evaluation and Research and products that have never been marketed, have been discontinued from marketing or that have had their approvals withdrawn for other than safety or efficacy reasons.

The Cumulative Supplement provides, among other things, information on newly approved drugs and, if necessary, revised therapeutic equivalence evaluations and updated patent and exclusivity data. The Addendum contains appropriate drug patent and exclusivity information required of the Agency by the "Drug Price Competition and Patent Term Restoration Act of 1984" for the Prescription, OTC, and Drug Products with Approval under Section 505 of the Act Administered by the Center for Biologics Evaluation and Research Lists.

The Patent and Exclusivity Lists are arranged in alphabetical order by active ingredient name. For those products with multiple active ingredients, only the first active ingredient (in alphabetical sort) will appear. In addition, the trade name will be displayed to the right of the active ingredient name for each product. Also shown is the application number and product number (FDA's internal file number) for reference purposes. All patents with their expiration dates are displayed for each application number. Use patents are indicated with the symbol "U" followed by a number representing a specific use. Exclusivity information for a specific drug is indicated by an abbreviation followed by the date upon which the exclusivity expires. Refer to the Exclusivity Terms section in the Patent and Exclusivity Information Addendum for an explanation of all codes and abbreviations.

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

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

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

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

Deletions new to the Prescription Drug Product List, OTC Drug Product List, and the Patent and Exclusivity Data are indicated by the symbol >DLT> (DELETE) to the left of the line containing shaded print. The >DLT> symbol is dropped in subsequent Cumulative Supplements for that item. The shaded print remains in the Prescription Drug Product List and OTC Drug Product Lists in all Cumulative Supplements for this edition. However, the shaded print in the Patent and Exclusivity Data is dropped in subsequent Cumulative Supplements.

Products that have never been marketed, have been discontinued from marketing or that have had their approvals withdrawn for other than safety or efficacy reasons, will be flagged in this Cumulative Supplement with the "@" symbol to designate their non-marketed status. All products having a "@" symbol in the 12th Cumulative Supplement of the 14th Edition List will then be added to the "Discontinued Drug Product List" appearing in the 15th Edition.

## 1.2 PRODUCTS REQUIRING REVISED LABELING FOR FULL APPROVAL

Drug products in this category (1) initially received approval only on the basis of safety before effectiveness studies were required, or (2) were conditionally approved under the temporary exemption that allowed these products to be marketed while effectiveness studies were being conducted. Listed below are those drugs which are now required to revise their labeling and provide additional information necessary for full approval on the basis of requirements listed in the Federal Register. As approval is granted by the Agency for a specific product, based on additional information submitted by the applicant, the product will be included in the appropriate Drug Product List.

<u>Products</u>	<u>Federal Register Reference</u>
Nitroglycerin (capsule, controlled release;oral)	SEP 07, 1984 (49 FR 35428)
Nitroglycerin (film, extended release; transdermal*)	JUL 15, 1993 (58 FR 38129)
Nitroglycerin (tablet, controlled release;oral)	SEP 07, 1984 (49 FR 35428)
Nitroglycerin (tablet, controlled release;buccal)	JUL 05, 1985 (50 FR 27688)

\*The Federal Register of July 15, 1993 (58 FR 38129) announced that the FDA was revoking the temporary exemption for nitroglycerin in a transdermal delivery system. Marketing of a drug product that is the subject of a conditionally approved ANDA may continue by meeting the requirements listed in the Federal Register. Firms wishing to submit a new ANDA before a drug product is approved (NDA or ANDA) and appears in the List should submit a 505(b)(2) application following the directions contained in the Federal Register. Nitro-Dur has been selected as the reference listed drug. The preamble to the final rule (57 FR 17958) states if there are multiple NDA's, the reference listed drug generally will be the market leader. This is the basis upon which Nitro-Dur was selected. In addition, the preamble states that, in multiple NDA situations, a product not designated as the reference listed drug and not shown to be bioequivalent to the reference listed drug may be shielded from generic competition. This is the case with Summit's Transderm-Nitro. The Office of Generic Drugs (OGD) has been requested to have a second listed drug as provided for in the Final Rule. OGD has granted this request. Therefore, at the time that Schering's and Summit's supplements are fully approved and their products are entered into the List, we will have two reference listed drugs for the nitroglycerin transdermal systems. Firms may, therefore, elect to conduct bioequivalence studies against either of these products. It is conceivable that a non-referenced listed drug may be fully approved and appear in the List; in this case, the Agency's referenced listed drug will not change and a 505(b)(2) application will be appropriate until the reference listed drugs are fully approved. Once they are fully approved, a 505(j) application will be the appropriate mechanism for an ANDA submission.

## 1.3 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 automatically reflect the new abbreviated name. Consequently, it will not appear as an applicant name change entry in the Cumulative Supplement nor will the cumulation of these name changes appear in this section. However, when the applicant name change is one which may not be easily recognized or located in the listing (e.g., White Towne Paulsen [Former Abbreviated Name] is changed to Whiteworth Towne [New Abbreviated Name], the name change will appear in this section and will be identified with an asterisk.

APPLICANT NAME CHANGES

<u>FORMER APPLICANT NAME</u> <u>(FORMER ABBREVIATED NAME)</u>	<u>NEW APPLICANT NAME</u> <u>(NEW ABBREVIATED NAME)</u>
ADRIA LABORATORIES DIV ERBAMONT INC (ADRIA)	PHARMACIA INC (PHARMACIA)
BANNER GELATIN PRODUCTS CORP (BANNER GELATIN)	BANNER PHARMACAPS INC (BANNER PHARMACAPS)
*BROCADES PHARMA BV DIV YAMANOUCHI GROUP (BROCADES PHARMA)	*YAMANOUCHI EUROPE BV (YAMANOUCHI)
CLONMEL CHEMICALS CO LTD (CLONMEL)	CLONMEL HEALTH CARE (CLONMEL HLTH CARE)
DUPONT PHARMACEUTICALS (DUPONT)	DUPONT MERCK PHARMACEUTICAL CO (DUPONT MERCK)
*G POHL BOSKAMP GMBH AND CO (BOSKAMP)	*G POHL BOSKAMP GMBH AND CO (POHL BOSKAMP)
GYNEX INC (GYNEX)	BTG PHARMACEUTICALS CORP SUB BIOTECHNOLOGY GENERAL CORP (BTG PHARMS)
KABI PHARMACIA INC (KABI)	PHARMACIA INC (PHARMACIA)
*KM LEE LABORATORIES (LEE LABS)	*KM LEE LABORATORIES INC (KM LEE)

## APPLICANT NAME CHANGES

<u>FORMER APPLICANT NAME</u> <u>(FORMER ABBREVIATED NAME)</u>	<u>NEW APPLICANT NAME</u> <u>(NEW ABBREVIATED NAME)</u>
*LABORATOIRES FOURNIER SA (FOURNIER)	*LABORATOIRES FOURNIER SCA (LABS FOURNIER)
*LABORATORIOS ATRAL SARL (ATRAL)	*LABORATORIOS ATRAL SARL (LABS ATRAL)
MALLINCKRODT SPECIALTY CHEMICALS CO (MALLINCKRODT)	MALLINCKRODT CHEMICAL INC (MALLINCKRODT)
*MM MAST AND CO (MAST)	*MM MAST AND CO (MM MAST)
NORTH AMERICAN CHEMICAL CORP (NORTH AM CHEM)	GOLDEN PHARMACEUTICALS INC (GOLDEN PHARMS)
PHARMACEUTICAL BASICS INC (PHARM BASICS)	ROSEMONT PHARMACEUTICAL CORP (ROSEMONT)
*PRIVATE FORMULATIONS INC (PRIVATE FORM)	*PRIVATE FORMULATIONS INC (PVT FORM)
RICHLYN LABORATORIES INC (RICHLYN)	GLOBAL PHARMACEUTICAL CORP (GLOBAL PHARMS)
*SCHWARZ PHARMA DIV KREMERS URBAN CO (SCHWARZ PHARMA)	*SCHWARZ PHARMA KREMERS URBAN CO SUB SCHWARZ PHARMA AG (SPKU)
SQUIBB DIAGNOSTICS (SQUIBB)	BRACCO DIAGNOSTICS INC (BRACCO DXS)
*WHITEWORTH TOWNE PAULSEN INC (WHITE TOWNE PAULSEN)	*WHITEWORTH TOWNE PAULSEN INC (WHITEWORTH TOWNE)

### 1.4 NEW INDICATIONS FOR PREVIOUSLY APPROVED DRUG PRODUCTS

When an application is submitted to FDA for a new indication for a drug product that duplicates a drug product (same active moiety, same salt, same formulation, or same combination) already approved or marketed in the United States by the same firm, the application is either submitted as a supplement to the original NDA (if the clinical expertise for the review of the new indication resides in the same division that reviewed the original NDA), or as a "Type 6 NDA" and assigned a new NDA number

(if the clinical expertise for the review of the new indication resides in another review division). When an application is submitted to FDA for a new indication for a drug product that duplicates a drug product (same active moiety, same salt, same formulation, or same combination) already approved or marketed in the United States by a different firm, the application is classified as "Type 6" and assigned a new NDA number. For administrative purposes, FDA has been listing all "Type 6 NDA's" in the *Approved Drug Products with Therapeutic Equivalence Evaluations*, (ADP), even when the application was submitted by the original NDA holder. However, FDA has determined that the practice of listing a separate "Type 6 NDA" number in the ADP when the applicant is the original NDA holder may cause confusion to the ADP reader.

Accordingly, to prevent confusion and to eliminate duplicity of data, the approval of an application for a new indication for a previously approved drug product submitted by the original NDA holder will no longer be listed in the ADP. Any exclusivity awarded for that approval will be shown in the Patent and Exclusivity Information Addendum under the original NDA number. However, approval of an application for a new indication submitted by an applicant other than the original NDA holder will be shown in the appropriate drug product list of the ADP. Any exclusivity awarded will be shown under the NDA number of the new applicant.

All approvals of "Type 6" applications submitted by the original NDA holder currently in the ADP are listed in the table below. For reference purposes, the original NDA number is listed next to the corresponding "Type 6 NDA Number". This data ("Type 6 NDA Number") will continue to be listed in the remaining Cumulative Supplements to the 14th Edition of the ADP; but it will not appear in the 15th Edition of the ADP.

<u>TYPE 6 NDA NUMBER</u>	<u>ORIGINAL NDA NUMBER</u>	<u>ACTIVE INGREDIENT (TRADE NAME)</u>	<u>DOSAGE FORM (ROUTE)</u>
17-117	16-020	AMANTADINE HCL (SYMMETREL)	CAPSULE (ORAL)
17-118	16-023	AMANTADINE HCL (SYMMETREL)	SYRUP (ORAL)
50-697	50-662	CLARITHROMYCIN (BIAXIN)	TABLET (ORAL)
19-576	19-084	KETOCONAZOLE (NIZORAL)	CREAM (TOPICAL)
19-648	19-084	KETOCONAZOLE (NIZORAL)	CREAM (TOPICAL)
18-064	18-063	NADOLOL (CORGARD)	TABLET (ORAL)
20-109	19-886	NAFARELIN ACETATE (SYNAREL)	SPRAY, METERED (NASAL)
20-223	19-057	TERAZOSIN HCL (HYTRIN)	TABLET (ORAL)

## 1.5 USP MONOGRAPH TITLE ADDITIONS OR CHANGES

The U.S. Pharmacopeia (USP) periodically makes additions to or changes in monograph titles. Some of these additions or changes may affect dosage form terms listed in *Approved Drug Products with Therapeutic Equivalence Evaluations* (ADP). Instead of making the change in each affected product, the Cumulative Supplement (CS) will list applicable monograph title and dosage form additions or changes in this section. These will appear as soon as the modified USP monograph title is official. It is possible for these additions or changes to be listed in this section before all applicant holders have made labeling modifications.

The monograph title additions or changes shown below will remain in this section in each succeeding supplement of this edition. Once the next edition of the ADP is published, the products affected by the title additions or changes will be displayed with the new dosage form in the appropriate drug list. As notification to the reader, these monograph title additions or changes will also be listed in a special section of the ADP.

### USP MONOGRAPH TITLE ADDITIONS OR CHANGES

FORMER USP MONOGRAPH TITLE  
(FORMER ADP DOSAGE FORM; ROUTE)

NEW USP MONOGRAPH TITLE  
(NEW ADP DOSAGE FORM; ROUTE)

THERE WERE NO USP MONOGRAPH TITLE ADDITIONS OR CHANGES DURING THE MONTH OF  
DECEMBER 1994.

## 1.6 REPORT OF COUNTS FOR THE PRESCRIPTION DRUG PRODUCT LIST

### DESCRIPTION OF REPORT

This report provides summary counts derived from the product information in the Prescription Drug Product List and the current Cumulative Supplement. Products included in the counts are domestically marketed drug products approved for both safety and effectiveness under sections 505 and 507 of the Federal Food, Drug, and Cosmetic Act. Excluded are approved drug products marketed by distributors; those marketed solely abroad; and those now regarded as medical devices, biologics or foods.

The baseline column (Dec 1993) 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 1993</u>	<u>JUN 1994</u>	<u>SEP 1994</u>	<u>DEC 1994</u>
DRUG PRODUCTS LISTED	9140	9079	9092	9141
SINGLE SOURCE	2144 (23.5%)	2150 (23.7%)	2127 (23.4%)	2178 (23.8%)
MULTISOURCE	6996 (76.5%)	6929 (76.3%)	6965 (76.6%)	6963 (76.2%)
THERAPEUTICALLY EQUIVALENT	6292 (68.8%)	6290 (69.3%)	6329 (69.6%)	6330 (69.2%)
NOT THERAPEUTICALLY EQUIVALENT	527 ( 5.8%)	458 ( 5.0%)	453 ( 5.0%)	453 ( 5.0%)
EXCEPTIONS <sup>1</sup>	177 ( 1.9%)	181 ( 2.0%)	183 ( 2.0%)	180 ( 2.0%)
NEW MOLECULAR ENTITIES APPROVED	--	5	2	10
NUMBER OF APPLICANTS	526	490	539	534

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



PRESCRIPTION DRUG PRODUCT LIST  
14TH EDITION  
RX DRUG PRODUCT LIST / CUMULATIVE SUPPLEMENT NUMBER 12 / JAN'94 - DEC'94

1

ACETAMINOPHEN; BUTALBITAL; CAFFEINE

CAPSULE; ORAL  
MEDICESTIC PLUS  
AB US CHEM 325MG;50MG;40MG N89115 001  
JAN 14, 1986  
@ 325MG;50MG;40MG N89115 001  
JAN 14, 1986

ACETAMINOPHEN; CAFFEINE; DIHYDROCODEINE BITARTRATE

CAPSULE; ORAL  
COMPAL  
AA PURDUE FREDERICK 356.4MG;30MG;16MG N88584 001  
MAR 04, 1986  
DHC PLUS  
AA PURDUE FREDERICK 356.4MG;30MG;16MG N88584 001  
MAR 04, 1986

ACETAMINOPHEN; CODEINE PHOSPHATE

TABLET; ORAL  
ACETAMINOPHEN AND CODEINE PHOSPHATE  
> ADD > AA ROYCE LABS 300MG;15MG N89997 001  
> ADD > DEC 28, 1994  
> ADD > AA 300MG;30MG N89998 001  
> ADD > DEC 28, 1994  
> ADD > AA 300MG;60MG N89999 001  
> ADD > DEC 28, 1994

ACETAMINOPHEN; HYDROCODONE BITARTRATE

CAPSULE; ORAL  
CO-CESSIC  
AA CENT PHARMS 500MG;5MG N89360 001  
MAR 02, 1988  
@ 500MG;5MG N89360 001  
MAR 02, 1988

ACETAMINOPHEN; PENTAZOCINE HYDROCHLORIDE

TABLET; ORAL  
TALACEN  
+ SANOFI WINTHROP 650MG;EQ 25MG BASE N18458 001  
SEP 23, 1982

ACETAMINOPHEN; PENTAZOCINE HYDROCHLORIDE

TABLET; ORAL  
TALACEN  
\* STERLING WINTHROP 650MG;EQ 25MG BASE N18458 001  
SEP 23, 1982

ACETAMINOPHEN; PROPOXYPHENE NAPSYLATE

TABLET; ORAL  
PROPOXYPHENE NAPSYLATE AND ACETAMINOPHEN  
> ADD > AB LEMMON 650MG;100MG N74119 001  
> ADD > DEC 19, 1994

ACETAZOLAMIDE

CAPSULE, EXTENDED RELEASE; ORAL  
DIAMOX  
> DLT > \* LEDERLE 500MG N12945 001  
> ADD > + STORZ OPHTHALM 500MG N12945 001

TABLET; ORAL

DIAMOX  
> DLT > AB LEDERLE 125MG N08943 001  
> DLT > AB \* 250MG N08943 002  
> ADD > AB STORZ OPHTHALM 125MG N08943 001  
> ADD > AB + 250MG N08943 002

ACETAZOLAMIDE SODIUM

INJECTABLE; INJECTION  
DIAMOX  
> DLT > \* LEDERLE EQ 500MG BASE/VIAL N09388 001  
> DLT > DEC 05, 1990  
> ADD > + STORZ OPHTHALM EQ 500MG BASE/VIAL N09388 001  
> ADD > DEC 05, 1990

ACETIC ACID, GLACIAL; ALUMINUM ACETATE

SOLUTION/DROPS; OTIC  
ACETIC ACID 2% IN AQUEOUS ALUMINUM ACETATE  
AT BAUSCH AND LOMB 2%;0.79% N40063 001  
FEB 25, 1994  
BOROFAIR  
AT PHARMAFAIR 2%;0.79% N88606 001  
AUG 21, 1985

ACETIC ACID, GLACIAL; ALUMINUM ACETATE

SOLUTION/DROPS; OTIC

BOROPAIN

@ PHARMAFAIR 2%;0.79%

N88606 001  
AUG 21, 1985

DOMEBORO

AT MILES

2%;0.79%

N84476 001

AT +

2%;0.79%

N84476 001

ACETIC ACID, GLACIAL; HYDROCORTISONE

SOLUTION/DROPS; OTIC

HYDROCORTISONE AND ACETIC ACID

AT BAUSCH AND LOMB 2%;1%

N40097 001  
OCT 31, 1994

ACETYLCHOLINE CHLORIDE

POWDER FOR RECONSTITUTION; OPHTHALMIC

MIOCHOL

+ CIBA VISION 20MG/VIAL

N16211 001

\* IOLAB 20MG/VIAL

N16211 001

MIOCHOL-E

+ CIBA VISION 20MG/VIAL

N20213 001

\* IOLAB 20MG/VIAL

SEP 22, 1993

N20213 001

SEP 22, 1993

ACETYLCYSTEINE

SOLUTION; INHALATION, ORAL

ACETYLCYSTEINE

AN ABBOTT 10%

N73664 001

AUG 30, 1994

AN 20%

N74037 001

AUG 30, 1994

ACRIVASTINE; PSEUDOEPHEDRINE HYDROCHLORIDE

CAPSULE; ORAL

SEMPREX-D

+ BURROUGHS WELLCOME 8MG;60MG

N19806 001

MAR 25, 1994

ALBUTEROL SULFATE

SYRUP; ORAL

ALBUTEROL SULFATE

AA MOVA

EQ 2MG BASE/5ML

N74302 001  
SEP 30, 1994

TABLET; ORAL

ALBUTEROL SULFATE

AB WARNER CHILCOTT

EQ 2MG BASE

N72817 001

JAN 09, 1990

AB

EQ 4MG BASE

N72818 001

JAN 09, 1990

@

EQ 2MG BASE

N72817 001

JAN 09, 1990

@

EQ 4MG BASE

N72818 001

JAN 09, 1990

ALPRAZOLAM

TABLET; ORAL

ALPRAZOLAM

AB MYLAN

0.25MG

N74215 001

JAN 27, 1994

AB

0.5MG

N74215 002

JAN 27, 1994

AB

1MG

N74215 003

JAN 27, 1994

AB

2MG

N74215 004

JAN 27, 1994

AB NOVOPHARM

0.25MG

N74085 001

FEB 16, 1994

AB

0.5MG

N74085 002

FEB 16, 1994

AB

1MG

N74085 003

FEB 16, 1994

AB ZENITH LABS

0.25MG

N74294 001

JUL 29, 1994

AB

0.5MG

N74294 002

JUL 29, 1994

AB

1MG

N74294 003

JUL 29, 1994

AB

2MG

N74294 004

JUL 29, 1994

AMANTADINE HYDROCHLORIDE

SYRUP; ORAL

AMANTADINE HCL  
AA HI TECH PHARMA 50MG/5ML N74170 001  
 OCT 28, 1994

AMBENONIUM CHLORIDE

TABLET; ORAL  
 MYTELASE

SANOFI WINTHROP 10MG N10155 002  
 STERLING WINTHROP 10MG N10155 002

AMIKACIN SULFATE

INJECTABLE; INJECTION

AMIKACIN  
AP ABBOTT EQ 50MG BASE/ML N63263 001  
 NOV 30, 1994  
AP EQ 250MG BASE/ML N63264 001  
 NOV 30, 1994  
AP EQ 250MG BASE/ML N63265 001  
 NOV 30, 1994  
AP EQ 250MG BASE/ML N63266 001  
 OCT 31, 1994  
EQ 62.5MG BASE/ML N63283 001  
 OCT 31, 1994  
AP BEDFORD EQ 50MG BASE/ML N63313 001  
 APR 11, 1994  
AP EQ 250MG BASE/ML N63315 001  
 APR 11, 1994  
AP ELKINS SINN EQ 50MG BASE/ML N63274 001  
 MAY 18, 1992  
AP + EQ 50MG BASE/ML N63274 001  
 MAY 18, 1992  
AP EQ 250MG BASE/ML N63275 001  
 MAY 18, 1992  
AP + EQ 250MG BASE/ML N63275 001  
 MAY 18, 1992  
AMIKIN  
 @ APOTHECON EQ 50MG BASE/ML N50495 001  
EQ 250MG BASE/ML N50495 002  
AP \* BRISTOL EQ 50MG BASE/ML N62562 001  
AP \* EQ 50MG BASE/ML N62562 001  
 SEP 20, 1984  
AP \* EQ 250MG BASE/ML N50495 002

AMIKACIN SULFATE

INJECTABLE; INJECTION

AMIKIN  
AP BRISTOL EQ 250MG BASE/ML N62562 002  
 SEP 20, 1984  
 @ EQ 50MG BASE/ML N62562 001  
 SEP 20, 1984  
 @ EQ 250MG BASE/ML N62562 002  
 SEP 20, 1984

AMINO ACIDS; DEXTROSE

INJECTABLE; INJECTION

AMINOSYN II 5% IN DEXTROSE 25% IN PLASTIC CONTAINER  
 ABBOTT 5%;25GM/100ML N19565 001  
 DEC 17, 1986  
 @ 5%;25GM/100ML N19565 001  
 DEC 17, 1986

AMINOPHYLLINE

INJECTABLE; INJECTION

AMINOPHYLLINE  
AP FUJISAWA 25MG/ML N87886 001  
 AUG 30, 1983  
 @ 25MG/ML N87886 001  
 AUG 30, 1983  
AP KING PHARMS 25MG/ML N86606 001  
AP SMITHKLINE BEECHAM 25MG/ML N86606 001

TABLET; ORAL

AMINOPHYLLINE  
AB GLOBAL PHARMS 100MG N84574 001  
AB 200MG N84576 001  
ED LANNETT 100MG N84588 001  
ED 200MG N84588 002  
 @ 100MG N84588 001  
 @ 200MG N84588 002  
ED 100MG N84574 001  
ED RICHLYN 200MG N84576 001

AMINOSALICYLIC ACID

GRANULE, DELAYED RELEASE; ORAL  
PAGER

+ JACOBUS 4GM/PACKET N74346 001  
JUN 30, 1994

AMITRIPTYLINE HYDROCHLORIDE

TABLET; ORAL

AMITRIPTYLINE HCL

<u>AB</u>	<u>LEMON</u>	<u>10MG</u>	<u>N86610 001</u>
<u>AB</u>		<u>25MG</u>	<u>N86859 001</u>
<u>AB</u>		<u>50MG</u>	<u>N86857 001</u>
<u>AB</u>		<u>75MG</u>	<u>N86860 001</u>
<u>AB</u>		<u>100MG</u>	<u>N86854 001</u>
<u>AB</u>		<u>150MG</u>	<u>N86853 001</u>
@		10MG	N86610 001
@		25MG	N86859 001
@		50MG	N86857 001
@		75MG	N86860 001
@		100MG	N86854 001
@		150MG	N86853 001

AMITRIPTYLINE HYDROCHLORIDE; PERPHENAZINE

TABLET; ORAL

TRIAVIL 2-10

> <u>ADD</u> >	<u>AB</u>	+ LOTUS	<u>10MG; 2MG</u>	<u>N14715 004</u>
> <u>DLT</u> >	<u>AB</u>	* MSD	<u>10MG; 2MG</u>	<u>N14715 004</u>
> <u>ADD</u> >	<u>AB</u>	+ LOTUS	<u>25MG; 2MG</u>	<u>N14715 002</u>
> <u>DLT</u> >	<u>AB</u>	* MSD	<u>25MG; 2MG</u>	<u>N14715 002</u>
> <u>ADD</u> >	<u>AB</u>	+ LOTUS	<u>10MG; 4MG</u>	<u>N14715 003</u>
> <u>DLT</u> >	<u>AB</u>	* MSD	<u>10MG; 4MG</u>	<u>N14715 003</u>
> <u>ADD</u> >	<u>AB</u>	+ LOTUS	<u>25MG; 4MG</u>	<u>N14715 005</u>
> <u>DLT</u> >	<u>AB</u>	* MSD	<u>25MG; 4MG</u>	<u>N14715 005</u>
> <u>ADD</u> >	<u>AB</u>	+ LOTUS	<u>50MG; 4MG</u>	<u>N14715 006</u>
> <u>DLT</u> >	<u>AB</u>	* MSD	<u>50MG; 4MG</u>	<u>N14715 006</u>

AMOXICILLIN

CAPSULE; ORAL  
AMOXICILLIN

<u>AB</u>	<u>BIOCHEMIE</u>	<u>250MG</u>	<u>N64076 001</u>
			<u>SEP 30, 1994</u>
<u>AB</u>		<u>500MG</u>	<u>N64076 002</u>
			<u>SEP 30, 1994</u>
<u>AB</u>	<u>POLYMOX</u>	<u>250MG</u>	<u>N63099 001</u>
	<u>APOTHECON</u>		<u>MAR 20, 1992</u>
<u>AB</u>		<u>500MG</u>	<u>N63099 002</u>
			<u>MAR 20, 1992</u>
<u>AB</u>	<u>BRISTOL MYERS</u>	<u>250MG</u>	<u>N63099 001</u>
			<u>MAR 20, 1992</u>
<u>AB</u>		<u>500MG</u>	<u>N63099 002</u>
			<u>MAR 20, 1992</u>

POWDER FOR RECONSTITUTION; ORAL

POLYMOX

@	<u>APOTHECON</u>	<u>125MG/5ML</u>	<u>N61851 001</u>
@		<u>125MG/5ML</u>	<u>N62323 001</u>
@		<u>250MG/5ML</u>	<u>N61851 002</u>
@		<u>250MG/5ML</u>	<u>N62323 002</u>
<u>AB</u>	<u>BRISTOL</u>	<u>125MG/5ML</u>	<u>N61851 001</u>
<u>AB</u>		<u>125MG/5ML</u>	<u>N62323 001</u>
<u>AB</u>		<u>250MG/5ML</u>	<u>N61851 002</u>
<u>AB</u>		<u>250MG/5ML</u>	<u>N62323 002</u>

AMPHETAMINE SULFATE

TABLET; ORAL

AMPHETAMINE SULFATE  
LANNETT

		<u>5MG</u>	<u>N83901 001</u>
			<u>AUG 31, 1984</u>
		<u>10MG</u>	<u>N83901 002</u>
			<u>AUG 31, 1984</u>
@		<u>5MG</u>	<u>N83901 001</u>
			<u>AUG 31, 1984</u>
@		<u>10MG</u>	<u>N83901 002</u>
			<u>AUG 31, 1984</u>

AMPHOTERICIN B

INJECTABLE; INJECTION

AMPHOTERICIN B

<u>AP</u>	<u>FUJISAWA</u>	<u>50MG/VIAL</u>	<u>N62728 001</u>
			<u>APR 13, 1987</u>

AMPHOTERICIN B

INJECTABLE; INJECTION  
AMPHOTERICIN B  
 @ FUJISAWA

50MG/VIAL

N62728 001  
 APR 13, 1987

AMPICILLIN/AMPICILLIN TRIHYDRATE

POWDER FOR RECONSTITUTION; ORAL

POLYCILLIN  
 @ APOTHECON  
 @

EQ 125MG BASE/5ML  
 EQ 250MG BASE/5ML  
~~EQ 125MG BASE/5ML~~  
~~EQ 250MG BASE/5ML~~

N62297 001  
 N62297 002  
~~N62297 001~~  
~~N62297 002~~

~~AB~~  
~~AB~~

~~BRISTOL~~

AMRINONE LACTATE

INJECTABLE; INJECTION  
 INOCOR

SANOFI WINTHROP

EQ 5MG BASE/ML

N18700 001  
 JUL 31, 1984

~~STERLING WINTHROP~~

~~EQ 5MG BASE/ML~~

~~N18700 001~~  
~~JUL 31, 1984~~

ANTAZOLINE PHOSPHATE; NAPHAZOLINE HYDROCHLORIDE

SOLUTION/DROPS; OPHTHALMIC  
 VASOCON-A

+ CIBA VISION

0.5%;0.05%

N18746 001  
 APR 30, 1990

~~\* TOLAB~~

~~0.5%;0.05%~~

~~N18746 001~~  
~~APR 30, 1990~~

ASCORBIC ACID; BIOTIN; CYANOCOBALAMIN; DEXPANTHENOL;  
 ERGOCALCIFEROL; FOLIC ACID; NIACINAMIDE; PYRIDOXINE  
 HYDROCHLORIDE; RIBOFLAVIN PHOSPHATE SODIUM; THIAMINE  
 HYDROCHLORIDE; VITAMIN A; VITAMIN E

INJECTABLE; INJECTION

M.V.C. 9+3  
 FUJISAWA

~~10MG/ML;0.006MG/ML;0.5 UGM/ML;  
 1.5MG/ML;20 IU/ML;0.04MG/ML;4MG/ML;  
 0.4MG/ML;0.36MG/ML;0.3MG/ML;  
 330 UNITS/ML;1 IU/ML~~

~~N18440 002~~  
~~AUG 08, 1985~~

~~AP~~

ASCORBIC ACID; BIOTIN; CYANOCOBALAMIN; DEXPANTHENOL;  
 ERGOCALCIFEROL; FOLIC ACID; NIACINAMIDE; PYRIDOXINE  
 HYDROCHLORIDE; RIBOFLAVIN PHOSPHATE SODIUM; THIAMINE  
 HYDROCHLORIDE; VITAMIN A; VITAMIN E

INJECTABLE; INJECTION

M.V.C. 9+3  
 @ FUJISAWA

10MG/ML;0.006MG/ML;0.5 UGM/ML;  
 1.5MG/ML;20 IU/ML;0.04MG/ML;4MG/ML;  
 0.4MG/ML;0.36MG/ML;0.3MG/ML;  
 330 UNITS/ML;1 IU/ML

N18440 002  
 AUG 08, 1985

ASPIRIN; PENTAZOCINE HYDROCHLORIDE

TABLET; ORAL

TALWIN COMPOUND

+ SANOFI WINTHROP

325MG;EQ 12.5MG BASE

N16891 001

\* ~~STERLING WINTHROP~~

~~325MG;EQ 12.5MG BASE~~

~~N16891 001~~

ATENOLOL

TABLET; ORAL

ATENOLOL

~~AB~~ GENPHARM

50MG

N74126 001  
 MAR 23, 1994

~~AB~~

100MG

N74126 002  
 MAR 23, 1994

~~AB~~ INVAMED

25MG

N74265 001  
 FEB 28, 1994

~~AB~~

50MG

N74265 002  
 FEB 28, 1994

~~AB~~

100MG

N74265 003  
 FEB 28, 1994

ATROPINE SULFATE; DIPHENOXYLATE HYDROCHLORIDE

TABLET; ORAL

LOFENE

~~AA~~ LANNETT

0.025MG;2.5MG

N85372 001

@

0.025MG;2.5MG

N85372 001

AZITHROMYCIN DIHYDRATE

POWDER FOR RECONSTITUTION; ORAL

ZITHROMAX

+ PFIZER

EQ 1GM BASE/PACKET

N50693 001  
SEP 28, 1994

BACITRACIN

OINTMENT; OPHTHALMIC

BACITRACIN

AT PHARMADERM

500 UNITS/GM

N62158 001

@

500 UNITS/GM

N62158 001

AT PHARMAFAIR

500 UNITS/GM

N62453 001

@

500 UNITS/GM

MAR 28, 1984  
N62453 001  
MAR 28, 1984

BACITRACIN ZINC; POLYMYXIN B SULFATE

OINTMENT; OPHTHALMIC

OCUMYCIN

> DLT >

AT PHARMAFAIR

500 UNITS/GM;

N62430 001

10,000 UNITS/GM

APR 08, 1983

> DLT >

@

500 UNITS/GM;

N62430 001

10,000 UNITS/GM

APR 08, 1983

> ADD >

POLYSPORIN

> DLT >

AT \* BURROUGHS WELLCOME

500 UNITS/GM;

N61229 001

10,000 UNITS/GM

> ADD >

+

500 UNITS/GM;

N61229 001

10,000 UNITS/GM

BACLOFEN

TABLET; ORAL

BACLOFEN

AB ROYCE LABS

10MG

N73092 001  
JAN 28, 1994

AB

20MG

N73093 001  
JAN 28, 1994

BENZYL BENZOATE

EMULSION; TOPICAL

BENZYL BENZOATE

\* LANNETT

50%  
50%

N84535 001  
N84535 001

BETAMETHASONE DIPROPIONATE

CREAM, AUGMENTED; TOPICAL

DIPROLENE

\* SCHERING

EQ 0.05% BASE

N19408 001

@

EQ 0.05% BASE

N19408 001

GEL; TOPICAL

DIPROLENE

+ SCHERING

EQ 0.05% BASE

N19408 002

NOV 22, 1991

LOTION; TOPICAL

BETAMETHASONE DIPROPIONATE

AB TARO PHARMS

EQ 0.05% BASE

N74272 001

SEP 30, 1994

OINTMENT; TOPICAL

BETAMETHASONE DIPROPIONATE

AB TARO PHARMS

EQ 0.05% BASE

N74271 001

SEP 15, 1994

BETAMETHASONE VALERATE

LOTION; TOPICAL

BETAMETHASONE VALERATE

AB PHARMAFAIR

EQ 0.1% BASE

N70484 001

MAY 29, 1987

@

EQ 0.1% BASE

N70484 001

MAY 29, 1987

OINTMENT; TOPICAL

BETAMETHASONE VALERATE

E\* CLAY PARK

EQ 0.1% BASE

N71478 001

DEC 23, 1987

@

EQ 0.1% BASE

N71478 001

DEC 23, 1987

BETHANECHOL CHLORIDE

TABLET; ORAL  
BETHANECHOL CHLORIDE  
AA LANNETT 5MG N84702 001  
AA 10MG N84712 001  
AA 25MG N84074 001  
 @ 5MG N84702 001  
 @ 10MG N84712 001  
 @ 25MG N84074 001

BITOLTEROL MESYLATE

AEROSOL, METERED; INHALATION  
 TORNALATE  
 + SANOFI WINTHROP 0.37MG/INH N18770 001  
 DEC 28, 1984  
 \* STERLING WINTHROP 0.37MG/INH N18770 001  
 DEC 28, 1984  
 SOLUTION; INHALATION  
 TORNALATE  
 SANOFI WINTHROP 0.2% N19548 001  
 FEB 19, 1992  
 STERLING WINTHROP 0.2% N19548 001  
 FEB 19, 1992

BROMPHENIRAMINE MALEATE; DEXTROMETHORPHAN HYDROBROMIDE;  
 PSEUDOEPHEDRINE HYDROCHLORIDE

SYRUP; ORAL  
DIMETANE-DX  
AA ROBINSON AH 2MG/5ML; 10MG/5ML; 30MG/5ML N11694 007  
 MAR 29, 1984

BUDESONIDE

AEROSOL, METERED; NASAL  
 RHINOCORT  
 + ASTRA 0.05MG/INH N20233 001  
 FEB 14, 1994

BUMETANIDE

INJECTABLE; INJECTION  
BUMETANIDE  
AP SANOFI WINTHROP 0.25MG/ML N74332 001  
 OCT 31, 1994  
BUMEX  
AP + ROCHE 0.25MG/ML N18226 001  
 FEB 28, 1983

BUTABARBITAL SODIUM

ELIXIR; ORAL  
BUTALAN  
 LANNETT 33.3MG/5ML N85880 001  
 @ 33.3MG/5ML N85880 001  
 TABLET; ORAL  
BUTISOL SODIUM  
AA WALLACE 100MG N00793 005  
 + 100MG N00793 005  
AA SODIUM BUTABARBITAL  
AA LANNETT 100MG N85881 001  
 @ 100MG N85881 001  
AA ZENITH 30MG N84040 001  
 @ ZENITH LABS 30MG N84040 001

CAFFEINE; ERGOTAMINE TARTRATE

SUPPOSITORY; RECTAL  
 MIGERGOT  
 BR G AND W LABS 100MG;2MG N86557 001  
 OCT 04, 1983  
WIGRAINE  
 BR ORGANON 100MG;2MG N86557 001  
 OCT 04, 1983

CALCIFEDIOL

CAPSULE; ORAL  
 CALDEROL  
 ORGANON 0.02MG N18312 001  
 + 0.05MG N18312 002

CALCIFEDIOL, ANHYDROUS

CAPSULE, ORAL  
CALDEROL  
ORGANON

0.02MG N18312 001  
 0.05MG N18312 002

CALCIUM; MEGLUMINE; METRIZOIC ACID

INJECTABLE; INJECTION  
ISOPAQUE 280  
@ NYCOMED

0.35MG/ML;140.1MG/ML;  
 461.8MG/ML N17506 001  
~~0.35MG/ML;140.1MG/ML;  
 461.8MG/ML N17506 001~~

CALCIUM CHLORIDE; DEXTROSE; MAGNESIUM CHLORIDE; SODIUM CHLORIDE;  
 SODIUM LACTATE

SOLUTION; INTRAPERITONEAL

DELFLX W/ DEXTROSE 1.5% LOW MAGNESIUM LOW CALCIUM IN  
 PLASTIC CONTAINER

AT FRESENIUS 18.4MG/100ML;1.5GM/100ML;  
5.08MG/100ML;538MG/100ML;  
448MG/100ML N20171 001  
 AUG 19, 1992

DELFLX W/ DEXTROSE 2.5% LOW MAGNESIUM LOW CALCIUM IN  
 PLASTIC CONTAINER

AT FRESENIUS 18.4MG/100ML;2.5GM/100ML;  
5.08MG/100ML;538MG/100ML;  
448MG/100ML N20171 002  
 AUG 19, 1992

DELFLX W/ DEXTROSE 4.25% LOW MAGNESIUM LOW CALCIUM IN  
 PLASTIC CONTAINER

AT FRESENIUS 18.4MG/100ML;4.25GM/100ML;  
5.08MG/100ML;538MG/100ML;  
448MG/100ML N20171 003  
 AUG 19, 1992

DIANEAL LOW CALCIUM W/ DEXTROSE 1.5% IN PLASTIC CONTAINER

AT BAXTER 18.3MG/100ML;1.5GM/100ML;  
5.08MG/100ML;538MG/100ML;  
448MG/100ML N20183 001  
 DEC 04, 1992

CALCIUM CHLORIDE; DEXTROSE; MAGNESIUM CHLORIDE; SODIUM CHLORIDE;  
 SODIUM LACTATE

SOLUTION; INTRAPERITONEAL

DIANEAL LOW CALCIUM W/ DEXTROSE 2.5% IN PLASTIC CONTAINER

AT BAXTER 18.3MG/100ML;2.5GM/100ML;  
5.08MG/100ML;538MG/100ML;  
448MG/100ML N20183 002  
 DEC 04, 1992

DIANEAL LOW CALCIUM W/ DEXTROSE 3.5% IN PLASTIC CONTAINER

AT BAXTER 18.3MG/100ML;3.5GM/100ML;  
5.08MG/100ML;538MG/100ML;  
448MG/100ML N20183 003  
 DEC 04, 1992

DIANEAL LOW CALCIUM W/ DEXTROSE 4.25% IN PLASTIC CONTAINER

AT BAXTER 18.3MG/100ML;4.25GM/100ML;  
5.08MG/100ML;538MG/100ML;  
448MG/100ML N20183 004  
 DEC 04, 1992

DIANEAL PD-2 W/ DEXTROSE 1.5% IN PLASTIC CONTAINER

AT BAXTER 18.3MG/100ML;1.5GM/100ML;  
5.08MG/100ML;538MG/100ML;  
448MG/100ML N17512 004

INPERSOL W/ DEXTROSE 1.5% IN PLASTIC CONTAINER

AT ABBOTT 25.7MG/100ML;1.5GM/100ML;  
15.2MG/100ML;567MG/100ML;  
392MG/100ML N18379 002

AT FRESENIUS 25.7MG/100ML;1.5GM/100ML;  
15.2MG/100ML;567MG/100ML;  
392MG/100ML N18379 002

INPERSOL W/ DEXTROSE 2.5% IN PLASTIC CONTAINER

AT ABBOTT 25.7MG/100ML;2.5GM/100ML;  
15.2MG/100ML;567MG/100ML;  
392MG/100ML N18379 003

AT FRESENIUS 25.7MG/100ML;2.5GM/100ML;  
15.2MG/100ML;567MG/100ML;  
392MG/100ML N18379 003

INPERSOL W/ DEXTROSE 3.5% IN PLASTIC CONTAINER

AT ABBOTT 25.7MG/100ML;3.5GM/100ML;  
15.2MG/100ML;567MG/100ML;  
392MG/100ML N18379 007  
 JUN 24, 1988

AT FRESENIUS 25.7MG/100ML;3.5GM/100ML;  
15.2MG/100ML;567MG/100ML;  
392MG/100ML N18379 007  
 JUN 24, 1988

CALCIUM CHLORIDE; DEXTROSE; MAGNESIUM CHLORIDE; SODIUM CHLORIDE;  
SODIUM LACTATE

SOLUTION; INTRAPERITONEAL

INPERSOL W/ DEXTROSE 4.25% IN PLASTIC CONTAINER

AT ABBOTT 25.7MG/100ML; 4.25GM/100ML;  
15.2MG/100ML; 567MG/100ML;  
392MG/100ML N18379 001  
AT FRESENIUS 25.7MG/100ML; 4.25GM/100ML;  
15.2MG/100ML; 567MG/100ML;  
392MG/100ML N18379 001

INPERSOL-LC/LM W/ DEXTROSE 1.5% IN PLASTIC CONTAINER

> DLT > AT ABBOTT 18.4MG/100ML; 1.5GM/100ML;  
> DLT > 5.08MG/100ML; 538MG/100ML;  
> DLT > 448MG/100ML N20374 001  
> DLT > JUN 13, 1994  
> ADD > AT FRESENIUS 18.4MG/100ML; 1.5GM/100ML;  
> ADD > 5.08MG/100ML; 538MG/100ML;  
> ADD > 448MG/100ML N20374 001  
> ADD > JUN 13, 1994

INPERSOL-LC/LM W/ DEXTROSE 2.5% IN PLASTIC CONTAINER

> DLT > AT ABBOTT 18.4MG/100ML; 2.5GM/100ML;  
> DLT > 5.08MG/100ML; 538MG/100ML;  
> DLT > 448MG/100ML N20374 002  
> DLT > JUN 13, 1994  
> ADD > AT FRESENIUS 18.4MG/100ML; 2.5GM/100ML;  
> ADD > 5.08MG/100ML; 538MG/100ML;  
> ADD > 448MG/100ML N20374 002  
> ADD > JUN 13, 1994

INPERSOL-LC/LM W/ DEXTROSE 3.5% IN PLASTIC CONTAINER

> DLT > AT ABBOTT 18.4MG/100ML; 3.5GM/100ML;  
> DLT > 5.08MG/100ML; 538MG/100ML;  
> DLT > 448MG/100ML N20374 003  
> DLT > JUN 13, 1994  
> ADD > AT FRESENIUS 18.4MG/100ML; 3.5GM/100ML;  
> ADD > 5.08MG/100ML; 538MG/100ML;  
> ADD > 448MG/100ML N20374 003  
> ADD > JUN 13, 1994

INPERSOL-LC/LM W/ DEXTROSE 4.25% IN PLASTIC CONTAINER

> DLT > AT ABBOTT 25.7MG/100ML; 4.25GM/100ML;  
> DLT > 5.08MG/100ML; 538MG/100ML;  
> DLT > 448MG/100ML N20374 004  
> DLT > JUN 13, 1994  
> ADD > AT FRESENIUS 25.7MG/100ML; 4.25GM/100ML;  
> ADD > 5.08MG/100ML; 538MG/100ML;  
> ADD > 448MG/100ML N20374 004  
> ADD > JUN 13, 1994

CALCIUM CHLORIDE; DEXTROSE; MAGNESIUM CHLORIDE; SODIUM CHLORIDE;  
SODIUM LACTATE

SOLUTION; INTRAPERITONEAL

INPERSOL-LM W/ DEXTROSE 1.5% IN PLASTIC CONTAINER

AT ABBOTT 25.7MG/100ML; 1.5GM/100ML;  
5.08MG/100ML; 538MG/100ML;  
448MG/100ML N18379 004  
JUL 07, 1982  
AT FRESENIUS 25.7MG/100ML; 1.5GM/100ML;  
5.08MG/100ML; 538MG/100ML;  
448MG/100ML N18379 004  
JUL 07, 1982

INPERSOL-LM W/ DEXTROSE 2.5% IN PLASTIC CONTAINER

AT ABBOTT 25.7MG/100ML; 2.5GM/100ML;  
5.08MG/100ML; 538MG/100ML;  
448MG/100ML N18379 005  
JUL 07, 1982  
AT FRESENIUS 25.7MG/100ML; 2.5GM/100ML;  
5.08MG/100ML; 538MG/100ML;  
448MG/100ML N18379 005  
JUL 07, 1982

INPERSOL-LM W/ DEXTROSE 3.5% IN PLASTIC CONTAINER

AT ABBOTT 25.7MG/100ML; 3.5GM/100ML;  
5.08MG/100ML; 538MG/100ML;  
448MG/100ML N18379 008  
JUN 24, 1988  
AT FRESENIUS 25.7MG/100ML; 3.5GM/100ML;  
5.08MG/100ML; 538MG/100ML;  
448MG/100ML N18379 008  
JUN 24, 1988

INPERSOL-LM W/ DEXTROSE 4.25% IN PLASTIC CONTAINER

AT ABBOTT 25.7MG/100ML; 4.25GM/100ML;  
5.08MG/100ML; 538MG/100ML;  
448MG/100ML N18379 006  
JUL 07, 1982  
AT FRESENIUS 25.7MG/100ML; 4.25GM/100ML;  
5.08MG/100ML; 538MG/100ML;  
448MG/100ML N18379 006  
JUL 07, 1982

CALCIUM CHLORIDE; DEXTROSE; SODIUM CHLORIDE; SODIUM LACTATE

SOLUTION; INTRAPERITONEAL

INPERSOL-ZM W/ DEXTROSE 1.5% IN PLASTIC CONTAINER

@ ABBOTT 25.7MG/100ML; 1.5GM/100ML;  
538MG/100ML; 448MG/100ML N19395 001  
MAR 26, 1986

CALCIUM CHLORIDE; DEXTROSE; SODIUM CHLORIDE; SODIUM LACTATE

SOLUTION; INTRAPERITONEAL  
 INPERSOL-ZM W/ DEXTROSE 1.5% IN PLASTIC CONTAINER  
 © FRESenius 25.7MG/100ML;1.5GM/100ML;  
 538MG/100ML;448MG/100ML N19395 001  
 MAR 26, 1986

INPERSOL-ZM W/ DEXTROSE 2.5% IN PLASTIC CONTAINER  
 © ABBOTT 25.7MG/100ML;2.5GM/100ML;  
 538MG/100ML;448MG/100ML N19395 002  
 MAR 26, 1986

© FRESenius 25.7MG/100ML;2.5GM/100ML;  
 538MG/100ML;448MG/100ML N19395 002  
 MAR 26, 1986

INPERSOL-ZM W/ DEXTROSE 4.25% IN PLASTIC CONTAINER  
 © ABBOTT 25.7MG/100ML;4.25GM/100ML;  
 538MG/100ML;448MG/100ML N19395 003  
 MAR 26, 1986

© FRESenius 25.7MG/100ML;4.25GM/100ML;  
 538MG/100ML;448MG/100ML N19395 003  
 MAR 26, 1986

CALCIUM METRIZOATE; MEGLUMINE METRIZOATE; METRIZOATE MAGNESIUM;  
 METRIZOATE SODIUM

INJECTABLE; INJECTION  
 ISOPAQUE 440  
 © NYCOMED 0.78MG/ML;75.9MG/ML;0.15MG/ML;  
 16.6MG/ML N16847 001

© STERLING WINTHROP 0.78MG/ML;75.9MG/ML;0.15MG/ML;  
 16.6MG/ML N16847 001

CARBACHOL

INJECTABLE; INJECTION  
 CARBACHOL  
 © PHARMAFAIR 0.01% N70292 001  
 MAY 21, 1986

MIOSTAT  
 \* ALCON 0.01% N16968 001

SOLUTION; INTRAOCULAR  
 CARBACHOL  
 © PHARMAFAIR 0.01% N70292 001  
 MAY 21, 1986

MIOSTAT  
 + ALCON 0.01% N16968 001

CARBAMAZEPINE

SUSPENSION; ORAL  
 TEGRETOL  
 + BASEL PHARMS 100MG/5ML N18927 001  
 DEC 18, 1987

\* GEIGY 100MG/5ML N18927 001  
 DEC 18, 1987

TABLET; ORAL  
 TEGRETOL  
 AB + BASEL PHARMS 200MG N16608 001  
 AB \* GEIGY 200MG N16608 001

TABLET, CHEWABLE; ORAL  
 TEGRETOL  
 AB + BASEL PHARMS 100MG N18281 001  
 AB \* GEIGY 100MG N18281 001

CARBIDOPA; LEVODOPA

TABLET; ORAL  
 CARBIDOPA AND LEVODOPA  
 AB SCS 10MG;100MG N74080 001  
 MAR 25, 1994

AB 25MG;100MG N74080 002  
 MAR 25, 1994

AB 25MG;250MG N74080 003  
 MAR 25, 1994

TABLET, EXTENDED RELEASE; ORAL  
 SINEMET CR  
 MERCK SHARP DOHME 25MG;100MG N19856 002  
 DEC 24, 1992

CEFADROXIL/CEFADROXIL HEMIHYDRATE

CAPSULE; ORAL  
 CEFADROXIL  
 AB APOTHECON EQ 500MG BASE N62291 001  
 AB ZENITH LABS EQ 500MG BASE N62766 001  
 MAR 03, 1987

© EQ 500MG BASE N62766 001  
 MAR 03, 1987

ULTRACEF  
 AB BRISTOL EQ 500MG BASE N62291 001

CEFADROXIL/CEFADROXIL HEMIHYDRATE

POWDER FOR RECONSTITUTION; ORAL

<u>AB</u>	<u>CEFADROXIL</u>	<u>EQ 125MG BASE/5ML</u>	N62334 001
<u>AB</u>	APOTHECON	<u>EQ 250MG BASE/5ML</u>	N62334 002
<u>AB</u>		<u>EQ 500MG BASE/5ML</u>	N62334 003
<u>AB</u>	<u>ULTRACEF</u>	<u>EQ 125MG BASE/5ML</u>	N62334 001
<u>AB</u>	BRISTON	<u>EQ 250MG BASE/5ML</u>	N62334 002
<u>AB</u>		<u>EQ 500MG BASE/5ML</u>	N62334 003

TABLET; ORAL

<u>AB</u>	<u>CEFADROXIL</u>	<u>EQ 1GM BASE</u>	N62774 001
	ZENITH LABS		APR 08, 1987
	@	<u>EQ 1GM BASE</u>	N62774 001
			APR 08, 1987
<u>AB</u>	<u>ULTRACEF</u>	<u>EQ 1GM BASE</u>	N62390 001
	APOTHECON		JUN 10, 1982
<u>AB</u>	BRISTON	<u>EQ 1GM BASE</u>	N62390 001
			JUN 10, 1982

CEFAZOLIN SODIUM

INJECTABLE; INJECTION

<u>AP</u>	<u>CEFAZOLIN SODIUM</u>	<u>EQ 500MG BASE/VIAL</u>	N62688 002
	FUJISAWA		NOV 17, 1986
<u>AP</u>		<u>EQ 1GM BASE/VIAL</u>	N62688 003
			NOV 17, 1986
<u>AP</u>		<u>EQ 10GM BASE/VIAL</u>	N62688 004
			NOV 17, 1986
<u>AP</u>		<u>EQ 20GM BASE/VIAL</u>	N62688 005
			AUG 03, 1987
	@	<u>EQ 500MG BASE/VIAL</u>	N62688 002
			NOV 17, 1986
	@	<u>EQ 1GM BASE/VIAL</u>	N62688 003
			NOV 17, 1986
	@	<u>EQ 10GM BASE/VIAL</u>	N62688 004
			NOV 17, 1986
	@	<u>EQ 20GM BASE/VIAL</u>	N62688 005
			AUG 03, 1987

CEFTIZOXIME SODIUM

INJECTABLE; INJECTION

	<u>CEFIZOX</u>	<u>EQ 1GM BASE/VIAL</u>	N63294 002
	FUJISAWA		MAR 31, 1994
		<u>EQ 2GM BASE/VIAL</u>	N63294 003
			MAR 31, 1994

CEFUROXIME AXETIL

POWDER FOR RECONSTITUTION; ORAL

	<u>CEFTIN</u>	<u>EQ 125MG BASE/5ML</u>	N50672 001
	+ GLAXO		JUN 30, 1994

CEPHALEXIN

CAPSULE; ORAL

	<u>CEPHALEXIN</u>	<u>EQ 250MG BASE</u>	N63186 001
	APOTHECON		DEC 30, 1994
> ADD >	<u>AB</u>	<u>EQ 500MG BASE</u>	N63186 002
> ADD >	<u>AB</u>		DEC 30, 1994

CEPHALOTHIN SODIUM

INJECTABLE; INJECTION

<u>AP</u>	<u>CEPHALOTHIN SODIUM</u>	<u>EQ 1GM BASE/VIAL</u>	N62666 002
	FUJISAWA		JUN 10, 1987
<u>AP</u>		<u>EQ 2GM BASE/VIAL</u>	N62666 001
			JUN 10, 1987
	@	<u>EQ 1GM BASE/VIAL</u>	N62666 002
			JUN 10, 1987
	@	<u>EQ 2GM BASE/VIAL</u>	N62666 001
			JUN 10, 1987
<u>AP</u>	<u>SEFFIN</u>	<u>EQ 1GM BASE/VIAL</u>	N62435 001
	GLAXO		NOV 15, 1983
<u>AP</u>		<u>EQ 2GM BASE/VIAL</u>	N62435 002
			NOV 15, 1983
	*	<u>EQ 10GM BASE/VIAL</u>	N62435 003
			NOV 15, 1983
	@	<u>EQ 1GM BASE/VIAL</u>	N62435 001
			NOV 15, 1983

CEPHALOTHIN SODIUM

INJECTABLE; INJECTION

SEFEIN

@ GLAXO

EQ 2GM BASE/VIAL

N62435 002

NOV 15, 1983

@

EQ 10GM BASE/VIAL

N62435 003

NOV 15, 1983

CHENODIOL

TABLET; ORAL

CHENIX

\* SOLVAY

250MG

N18513 002

JUL 28, 1983

@

250MG

N18513 002

JUL 28, 1983

CHLORAMPHENICOL

CREAM; TOPICAL

CHLOROMYCETIN

PARKE DAVIS

1%

N50183 001

@

1%

N50183 001

OINTMENT; OPHTHALMIC

CHLOROPAIR

PHARMAFAIR

1%

N62439 001

APR 21, 1983

@

1%

N62439 001

APR 21, 1983

SOLUTION/DROPS; OPHTHALMIC

CHLOROPAIR

PHARMAFAIR

0.5%

N62437 001

APR 14, 1983

@

0.5%

N62437 001

APR 14, 1983

CHLORHEXIDINE GLUCONATE

SOLUTION; DENTAL

PERIDEX

AT + PROCTER AND GAMBLE

0.12%

N19028 001

AUG 13, 1986

CHLORHEXIDINE GLUCONATE

SOLUTION; DENTAL

PERIOGARD

AT

COLGATE PALMOLIVE

0.12%

N73695 001

JAN 14, 1994

CHLORMERODRIN, HG-197

INJECTABLE; INJECTION

CHLORMERODRIN HG 197

@ BRACCO DXS

0.6-1.4mCi/ML

N17269 001

@ SQUIBB

0.6-1.4mCi/ML

N17269 001

CHLORMEZANONE

TABLET; ORAL

TRANCOPAL

SANOFI WINTHROP

100MG

N11467 003

200MG

N11467 005

STERLING WINTHROP

100MG

N11467 003

200MG

N11467 005

CHLOROQUINE HYDROCHLORIDE

INJECTABLE; INJECTION

ARALEN HCL

+ SANOFI WINTHROP

EQ 40MG BASE/ML

N06002 002

\* STERLING WINTHROP

EQ 40MG BASE/ML

N06002 002

CHLOROQUINE PHOSPHATE

TABLET; ORAL

ARALEN

AA + SANOFI WINTHROP

EQ 300MG BASE

N06002 001

AA STERLING WINTHROP

EQ 300MG BASE

N06002 001

CHLOROQUINE PHOSPHATE; PRIMAQUINE PHOSPHATE

TABLET; ORAL

ARALEN PHOSPHATE W/ PRIMAQUINE PHOSPHATE

@ SANOFI WINTHROP

EQ 300MG BASE;

N14860 002

EQ 45MG BASE

CHLOROQUINE PHOSPHATE; PRIMAQUINE PHOSPHATE

TABLET; ORAL  
 ARALEN PHOSPHATE W/ PRIMAQUINE PHOSPHATE  
 @ STERLING WINTHROP EQ 300MG BASE  
 EQ 45MG BASE N14860 002

CHLORPHENIRAMINE MALEATE

TABLET; ORAL  
CHLORPHENIRAMINE MALEATE  
 @ ZENITH ZENITH 4MG N80779 001  
 @ ZENITH LABS 4MG N80779 001

CHLORTHALIDONE

TABLET; ORAL  
 THALITONE  
 HORUS THERAP 15MG N19574 001  
 + 15MG DEC 20, 1988  
 N19574 001  
 DEC 20, 1988

CHOLESTYRAMINE

TABLET; ORAL  
 QUESTRAN  
 + BRISTOL MYERS SQUIBB EQ 1GM RESIN N73403 001  
 APR 28, 1994

CHYMOTRYPSIN

POWDER FOR RECONSTITUTION; OPHTHALMIC  
 CATARASE  
 @ CIBA VISION 150 UNITS/VIAL N18121 001  
 + 300 UNITS/VIAL N16938 001  
 @ IOLAB 150 UNITS/VIAL N18121 001  
 \* 300 UNITS/VIAL N16938 001

CIMETIDINE

TABLET; ORAL  
CIMETIDINE  
 AB ENDO LABS 200MG N74281 001  
 MAY 17, 1994

CIMETIDINE

TABLET; ORAL  
CIMETIDINE

AB ENDO LABS 300MG N74281 002  
 MAY 17, 1994  
 AB 400MG N74281 003  
 MAY 17, 1994  
 AB 800MG N74329 001  
 MAY 17, 1994  
 AB MYLAN 200MG N74246 001  
 MAY 17, 1994  
 AB 300MG N74246 002  
 MAY 17, 1994  
 AB 400MG N74246 003  
 MAY 17, 1994  
 AB 800MG N74246 004  
 MAY 17, 1994  
 AB NOVOPHARM 200MG N74151 001  
 MAY 17, 1994  
 AB 300MG N74151 002  
 MAY 17, 1994  
 AB 400MG N74151 003  
 MAY 17, 1994  
 AB 800MG N74463 001  
 MAY 17, 1994  
 AB ROXANE 300MG N74361 001  
 DEC 23, 1994  
 AB 400MG N74361 002  
 DEC 23, 1994  
 AB 800MG N74371 001  
 DEC 23, 1994

TAGAMET

AB SMITHKLINE BEECHAM 200MG N17920 002  
 AB 300MG N17920 003  
 AB 400MG N17920 004  
 AB + 800MG DEC 14, 1983  
 N17920 005  
 APR 30, 1986

CIMETIDINE HYDROCHLORIDE

INJECTABLE; INJECTION  
CIMETIDINE HCL

AP ENDO LABS EQ 300MG BASE/2ML N74005 001  
 AUG 31, 1994  
 AP LUITPOLD EQ 300MG BASE/2ML N74353 001  
 DEC 20, 1994

CIMETIDINE HYDROCHLORIDE

INJECTABLE; INJECTION

> ADD >  
 > ADD > AP CIMETIDINE HCL IN SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER  
ABBOTT EQ 6MG BASE/ML N74269 001  
 DEC 27, 1994  
 > ADD > + N74468 005  
 EQ 90MG BASE/100ML  
 DEC 29, 1994  
 > ADD > + N74468 006  
 EQ 120MG BASE/100ML  
 DEC 29, 1994  
 > ADD > + N74468 003  
 EQ 180MG BASE/100ML  
 DEC 29, 1994  
 > ADD > + N74468 004  
 EQ 240MG BASE/100ML  
 DEC 29, 1994  
 > ADD > + N74468 001  
 EQ 360MG BASE/100ML  
 DEC 29, 1994  
 > ADD > + N74468 002  
 EQ 480MG BASE/100ML  
 DEC 29, 1994

TAGAMET

AP + SMITHKLINE BEECHAM EQ 300MG BASE/2ML N17939 002  
TAGAMET HCL IN SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER  
 > ADD > AP + SMITHKLINE BEECHAM EQ 6MG BASE/ML N19434 001  
 OCT 31, 1985

SOLUTION; ORAL

CIMETIDINE HCL

AA BARRE EQ 300MG BASE/5ML N74176 001  
 JUN 01, 1994  
 > ADD > AA ENDO LABS EQ 300MG BASE/5ML N74251 001  
 DEC 22, 1994

TAGAMET

AA SMITHKLINE BEECHAM EQ 300MG BASE/5ML N17924 001

CLEMASTINE FUMARATE

SYRUP; ORAL

CLEMASTINE FUMARATE

AA LEMMON EQ 0.5MG BASE/5ML N73399 001  
 JUN 30, 1994

CLINDAMYCIN PHOSPHATE

INJECTABLE; INJECTION

CLINDAMYCIN PHOSPHATE

AP BEDFORD EQ 150MG BASE/ML N63163 001  
 JUN 30, 1994

CLINDAMYCIN PHOSPHATE

INJECTABLE; INJECTION

CLINDAMYCIN PHOSPHATE

AP DUPONT EQ 150MG BASE/ML N62908 001  
 FEB 01, 1989  
 @ DUPONT MERCK EQ 150MG BASE/ML N62908 001  
 FEB 01, 1989  
AP FUJISAWA EQ 150MG BASE/ML N62747 001  
 JUN 03, 1988  
 @ EQ 150MG BASE/ML N62747 001  
 JUN 03, 1988  
 SOLUTION; TOPICAL  
CLEOCIN  
UPJOHN EQ 1% BASE N50537 002  
 FEB 22, 1994

CLOBETASOL PROPIONATE

CREAM; TOPICAL

CLOBETASOL PROPIONATE

AB COPELY PHARM 0.05% N74087 001  
 FEB 16, 1994  
AB NMC 0.05% N74139 001  
 AUG 03, 1994  
AB + TEMOVATE 0.05% N19322 001  
 DEC 27, 1985  
AB \* 0.05% N20340 001  
 JUN 17, 1994  
BX + GLAXO 0.05% N20340 001  
 JUN 17, 1994

GEL; TOPICAL

TEMOVATE

+ GLAXO 0.05% N20337 001  
 APR 29, 1994

OINTMENT; TOPICAL

CLOBETASOL PROPIONATE

AB COPELY PHARM 0.05% N74089 001  
 FEB 16, 1994  
AB NMC 0.05% N74128 001  
 AUG 03, 1994  
AB + TEMOVATE 0.05% N19323 001  
 DEC 27, 1985

CLOMIPRAMINE HYDROCHLORIDE

CAPSULE; ORAL  
ANAFRANIL

BASEL PHARMS	25MG	N19906 001
		DEC 29, 1989
	50MG	N19906 002
		DEC 29, 1989
+	75MG	N19906 003
		DEC 29, 1989
CIBA	25MG	N19906 001
		DEC 29, 1989
	50MG	N19906 002
		DEC 29, 1989
*	75MG	N19906 003
		DEC 29, 1989

CODEINE PHOSPHATE; PHENYLEPHRINE HYDROCHLORIDE; PROMETHAZINE HYDROCHLORIDE

SYRUP; ORAL  
PROMETHAZINE VC W/ CODEINE

AA	PENNEX	10MG/5ML; 5MG/5ML; 6.25MG/5ML	N88896 001
			JAN 04, 1985
@	PENNEX PHARMS	10MG/5ML; 5MG/5ML; 6.25MG/5ML	N88896 001
			JAN 04, 1985

CODEINE PHOSPHATE; PROMETHAZINE HYDROCHLORIDE

SYRUP; ORAL  
PROMETHAZINE W/ CODEINE

AA	PENNEX	10MG/5ML; 6.25MG/5ML	N88875 001
			DEC 17, 1984
@	PENNEX PHARMS	10MG/5ML; 6.25MG/5ML	N88875 001
			DEC 17, 1984

COLESTIPOL HYDROCHLORIDE

TABLET; ORAL  
COLESTID  
UPJOHN

1GM	N20222 001
	JUL 19, 1994

CORTISONE ACETATE

INJECTABLE; INJECTION  
CORTISONE ACETATE

BP	STERIS	25MG/ML	N83147 003
BP		25MG/ML	N85677 001
BP		50MG/ML	N83147 004
BP		50MG/ML	N85677 002
@		25MG/ML	N83147 003
@		25MG/ML	N85677 001
@		50MG/ML	N83147 004
@		50MG/ML	N85677 002
BP	UPJOHN	25MG/ML	N08126 002
@		25MG/ML	N08126 002
	CORTONE		
	MERCK SHARP DOHME	25MG/ML	N07110 002
		50MG/ML	N07110 003
BP	MSD	25MG/ML	N07110 002
BP		50MG/ML	N07110 003

TABLET; ORAL  
CORTISONE ACETATE

BP	INWOOD	25MG	N80731 001
@	INWOOD LABS	25MG	N80731 001

CROMOLYN SODIUM

SOLUTION; INHALATION  
CROMOLYN SODIUM

AN	DEY	10MG/ML	N74209 001
			APR 26, 1994
AN	INTAL		
AN	+ FISOXS	10MG/ML	N18596 001
			MAY 28, 1982

CYANOCOBALAMIN

INJECTABLE; INJECTION

> DLT >	BERUBIGEN		
> DLT >	BP	1MG/ML	N06798 001
> ADD >	UPJOHN	1MG/ML	N06798 001
	@		

CYANOCOBALAMIN, CO-57

CAPSULE; ORAL  
RUBRATOPE-57  
BRACCO DXS

0.5-1 uCi	N16089 002
-----------	------------

CYANOCOBALAMIN, CO-57

CAPSULE; ORAL  
RUBRATOPE-57  
SQUIBB 0.5-1 uCi N16089 002

CYANOCOBALAMIN, CO-60

CAPSULE; ORAL  
RUBRATOPE-60  
@ BRACCO DXS 0.5-1 uCi N16090 002  
@ SQUIBB 0.5-1 uCi N16090 002

CYCLOBENZAPRINE HYDROCHLORIDE

TABLET; ORAL  
CYCLOBENZAPRINE HCL  
AB ROYCE LABS 10MG N74436 001  
NOV 30, 1994

CYCLOPENTOLATE HYDROCHLORIDE

SOLUTION/DROPS; OPHTHALMIC  
CYCLOGYL  
AT \* RECON 0.5% N84109 001  
+ 0.5% N84109 001  
PENTOLAIR  
AT BAUSCH AND LOMB 1% N40075 001  
APR 29, 1994  
AT PHARMARAIR 0.5% N88643 001  
FEB 09, 1987  
AT 1% N88150 001  
FEB 25, 1983  
@ 0.5% N88643 001  
FEB 09, 1987  
@ 1% N88150 001  
FEB 25, 1983

CYCLOSPORINE

CAPSULE; ORAL  
SANDIMMUNE  
SANDOZ 50MG N50625 003  
NOV 23, 1992  
> ADD >  
> ADD >

CYPROHEPTADINE HYDROCHLORIDE

TABLET; ORAL  
CYPROHEPTADINE HCL  
AA CHELSEA 4MG N86165 001  
@ CHELSEA LABS 4MG N86165 001

CYSTEAMINE BITARTRATE

CAPSULE; ORAL  
CYSTAGON  
MYLAN EQ 50MG BASE N20392 001  
AUG 15, 1994  
+ EQ 150MG BASE N20392 002  
AUG 15, 1994

CYTARABINE

INJECTABLE; INJECTION  
CYTARABINE  
+ BULL D 20MG/ML N72945 001  
FEB 28, 1994  
AP CETUS BEN VENUE 1GM/VIAL N74245 001  
AUG 31, 1994  
AP 2GM/VIAL N74245 002  
AUG 31, 1994  
CYTOSAR-U  
AP + UPJOHN 1GM/VIAL N16793 003  
DEC 21, 1987  
AP + 2GM/VIAL N16793 004  
DEC 21, 1987

DACTINOMYCIN

INJECTABLE; INJECTION  
COSMEGEN  
+ MERCK SHARP DOHME 0.5MG/VIAL N50682 001  
\* MSD 0.5MG/VIAL N60467 001

> ADD > DALTEPARIN SODIUM

> ADD >  
> ADD > INJECTABLE; INJECTION  
> ADD > FRAGMIN  
> ADD > + PHARMACIA 2,500 IU/0.2ML N20287 001  
> ADD > DEC 22, 1994

DANAZOL

CAPSULE; ORAL  
DANOCRINE

SANOFI WINTHROP	50MG	N17557 003
	100MG	N17557 004
	200MG	N17557 002
+ STERLING WINTHROP	50MG	N17557 003
	100MG	N17557 004
	200MG	N17557 002

DESMOPRESSIN ACETATE

SPRAY, METERED; NASAL  
DESMOPRESSIN ACETATE

+ RHONE POULENC RORER	0.15MG/INH	N20355 001
		MAR 07, 1994

DESONIDE

OINTMENT; TOPICAL  
DESONIDE

AB TARO	0.05%	N74254 001
		AUG 03, 1994

DEXAMETHASONE

TABLET; ORAL  
DECADRON

BP MERCK SHARP DOHME	0.25MG	N11664 004
MSD	0.25MG	N11664 004

DEXAMETHASONE; NEOMYCIN SULFATE; POLYMYXIN B SULFATE

OINTMENT; OPHTHALMIC  
DEXASPORIN

AT BAUSCH AND LOMB	0.1%; EQ 3.5MG BASE/GM; 10,000 UNITS/GM	N64063 001
		JUL 25, 1994

DEXAMETHASONE SODIUM PHOSPHATE

AEROSOL; NASAL  
DEXACORT

* ADAMS LABS	EQ 0.1MG PHOSPHATE/INH	N14242 001
--------------	------------------------	------------

DEXAMETHASONE SODIUM PHOSPHATE

AEROSOL; NASAL  
DEXACORT

+ MEDEVA	EQ 0.1MG PHOSPHATE/INH	N14242 001
----------	------------------------	------------

AEROSOL, METERED; INHALATION

DECADRON		
* MSD	EQ 0.1MG PHOSPHATE/INH	N13413 001
DEXACORT		
+ MEDEVA	EQ 0.1MG PHOSPHATE/INH	N13413 001

OINTMENT; OPHTHALMIC

AT DEXAIR	EQ 0.05% PHOSPHATE	N88071 001
PHARMAFAIR		DEC 28, 1982
@	EQ 0.05% PHOSPHATE	N88071 001
		DEC 28, 1982

SOLUTION/DROPS; OPHTHALMIC

DEXAIR		
PHARMAFAIR	EQ 0.1% PHOSPHATE	N88433 001
		DEC 15, 1983
@	EQ 0.1% PHOSPHATE	N88433 001
		DEC 15, 1983

DEXAMETHASONE SODIUM PHOSPHATE; NEOMYCIN SULFATE

SOLUTION/DROPS; OPHTHALMIC

AT NEOMYCIN SULFATE-DEXAMETHASONE SODIUM PHOSPHATE		
PHARMAFAIR	EQ 0.1% PHOSPHATE; EQ 3.5MG BASE/ML	N62539 001
		JAN 10, 1985
@	EQ 0.1% PHOSPHATE; EQ 3.5MG BASE/ML	N62539 001
		JAN 10, 1985

DEXTROAMPHETAMINE SULFATE

ELIXIR; ORAL  
DEXEDRINE

SMITHKLINE BEECHAM	5MG/5ML	N83902 001
@	5MG/5ML	N83902 001

TABLET; ORAL

AA DEXTROAMPHETAMINE SULFATE		
LANNETT	5MG	N83903 001

DEXTROAMPHETAMINE SULFATE

TABLET; ORAL			
<u>DEXTROAMPHETAMINE SULFATE</u>			
<u>AA</u>	<u>LANNETT</u>	<u>10MG</u>	<u>N83903 003</u>
		<u>15MG</u>	<u>N85652 001</u>
	@	<u>5MG</u>	<u>N83903 001</u>
	@	<u>10MG</u>	<u>N83903 003</u>
	@	<u>15MG</u>	<u>N85652 001</u>

DEXTROMETHORPHAN HYDROBROMIDE; PROMETHAZINE HYDROCHLORIDE

SYRUP; ORAL			
<u>PROMETHAZINE H/ DEXTROMETHORPHAN</u>			
<u>AA</u>	<u>PENNEX</u>	<u>15MG/5ML; 6.25MG/5ML</u>	<u>N88864 001</u>
			<u>JAN 04, 1985</u>
	@	<u>PENNEX PHARMS</u>	<u>15MG/5ML; 6.25MG/5ML</u>
			<u>N88864 001</u>
			<u>JAN 04, 1985</u>

DEXTROSE

INJECTABLE; INJECTION			
<u>DEXTROSE 5% IN PLASTIC CONTAINER</u>			
<u>AP</u>	<u>MCGAW</u>	<u>50MG/ML</u>	<u>N16730 002</u>

DEXTROTHYROXINE SODIUM

TABLET; ORAL			
<u>CHOLOXIN</u>			
	<u>BOOTS</u>	<u>4MG</u>	<u>N12302 004</u>
		<u>6MG</u>	<u>N12302 006</u>
	+	<u>4MG</u>	<u>N12302 004</u>
	@	<u>6MG</u>	<u>N12302 006</u>

DIATRIZOATE MEGLUMINE

INJECTABLE; INJECTION			
<u>CARDIOGRAFIN</u>			
	@	<u>BRACCO DXS</u>	<u>85%</u>
	@	<u>SQUIBB</u>	<u>85%</u>
		<u>HYPAQUE</u>	
<u>AP</u>		<u>NYCOMED</u>	<u>30%</u>
<u>AP</u>			<u>N16403 002</u>
<u>AP</u>			<u>N16403 001</u>
<u>AP</u>		<u>STERLING WINTHROP</u>	<u>30%</u>
<u>AP</u>			<u>N16403 002</u>
<u>AP</u>			<u>N16403 001</u>

DIATRIZOATE MEGLUMINE

INJECTABLE; INJECTION			
<u>RENO-60</u>			
<u>AP</u>		<u>BRACCO DXS</u>	<u>60%</u>
		<u>RENO-DIP</u>	
<u>AP</u>		<u>BRACCO DXS</u>	<u>30%</u>
		<u>RENO-M-60</u>	
<u>AP</u>		<u>SQUIBB</u>	<u>60%</u>
		<u>RENO-M-DIP</u>	
<u>AP</u>		<u>SQUIBB</u>	<u>30%</u>
			<u>N10040 016</u>
			<u>N10040 012</u>
			<u>N10040 016</u>
			<u>N10040 012</u>
SOLUTION; URETERAL			
<u>RENO-30</u>			
<u>AT</u>		<u>BRACCO DXS</u>	<u>30%</u>
		<u>RENO-N-30</u>	
<u>AT</u>		<u>SQUIBB</u>	<u>30%</u>
			<u>N10040 021</u>
			<u>N10040 021</u>
SOLUTION; URETHRAL			
<u>HYPAQUE-CYSTO</u>			
<u>AT</u>		<u>NYCOMED</u>	<u>30%</u>
<u>AT</u>		<u>STERLING WINTHROP</u>	<u>30%</u>
			<u>N16403 003</u>
			<u>N16403 003</u>

DIATRIZOATE MEGLUMINE; DIATRIZOATE SODIUM

INJECTABLE; INJECTION			
<u>HYPAQUE-76</u>			
<u>AP</u>		<u>NYCOMED</u>	<u>66%;10%</u>
<u>AP</u>		<u>STERLING WINTHROP</u>	<u>66%;10%</u>
		<u>HYPAQUE-M, 75%</u>	
	@	<u>NYCOMED</u>	<u>50%;25%</u>
	@	<u>STERLING WINTHROP</u>	<u>50%;25%</u>
		<u>HYPAQUE-M, 90%</u>	
	@	<u>NYCOMED</u>	<u>60%;30%</u>
	@	<u>STERLING WINTHROP</u>	<u>60%;30%</u>
		<u>ND-60</u>	
<u>AP</u>		<u>MALLINCKRODT</u>	<u>52%;8%</u>
	@		<u>52%;8%</u>
			<u>N86505 001</u>
			<u>N86505 001</u>
			<u>N10220 003</u>
			<u>N10220 003</u>
			<u>N10220 002</u>
			<u>N10220 002</u>
			<u>N87074 001</u>
			<u>N87074 001</u>

DIATRIZOATE MEGLUMINE; IODIPAMIDE MEGLUMINE

SOLUTION; INTRAUTERINE			
<u>SINOGRAFIN</u>			
		<u>BRACCO DXS</u>	<u>52.7%;26.8%</u>
		<u>SQUIBB</u>	<u>52.7%;26.8%</u>
			<u>N11324 002</u>
			<u>N11324 002</u>

DIATRIZOATE SODIUM

INJECTABLE; INJECTION

<u>AP</u>	<u>HYPAAQUE</u>			
	NYCOMED	50%	N09561 001	
		25%	N09561 003	
<u>AP</u>	STERLING WINTHROP	50%	N09561 001	
		25%	N09561 003	

POWDER FOR RECONSTITUTION; ORAL, RECTAL

	<u>HYPAAQUE</u>		
	NYCOMED	100%	N11386 001
	STERLING WINTHROP	100%	N11386 003

SOLUTION; ORAL, RECTAL

	<u>HYPAAQUE</u>		
	NYCOMED	40%	N11386 003
	STERLING WINTHROP	40%	N11386 003

SOLUTION; URETERAL

	<u>HYPAAQUE SODIUM 20%</u>		
	NYCOMED	20%	N09561 002
	@ STERLING WINTHROP	20%	N09561 002

DICLOXACILLIN SODIUM

CAPSULE; ORAL

<u>AB</u>	<u>DYNAPEN</u>		
	APOTHECON	<u>EQ 500MG BASE</u>	N61454 003

DICUMAROL

TABLET; ORAL

	<u>DICUMAROL</u>		
	ABBOTT	25MG	N05545 003
	+	25MG	N05545 003

DIENESTROL

CREAM; VAGINAL

<u>AT</u>	<u>ESTRAGUARD</u>		
	SOLVAY	0.01%	N84436 001
	@	0.01%	N84436 001

DIETHYLPROPION HYDROCHLORIDE

TABLET; ORAL

<u>AA</u>	<u>TEPANIL</u>		
	3M	25MG	N11673 001
	@	25MG	N11673 001

DIETHYLSTILBESTROL

TABLET; ORAL

	<u>DIETHYLSTILBESTROL</u>		
BP	LILLY	1MG	N04041 004
BP *		5MG	N04041 005
		1MG	N04041 004
	+	5MG	N04041 005
	<u>STILBESTROL</u>		
BP	TABLICAPS	1MG	N83002 001
BP		5MG	N83006 001
	@	1MG	N83002 001
	@	5MG	N83006 001

TABLET, DELAYED RELEASE; ORAL

	<u>DIETHYLSTILBESTROL</u>		
BE *	LILLY	1MG	N04039 004
BE *		5MG	N04039 006
	@	1MG	N04039 004
	@	5MG	N04039 006
	<u>STILBESTROL</u>		
BE	TABLICAPS	0.5MG	N83003 001
BE		1MG	N83005 001
BE		5MG	N83007 001
	@	0.5MG	N83003 001
	@	1MG	N83005 001
	@	5MG	N83007 001
	<u>STILRETIN</u>		
BE	SQUIBB	0.5MG	N04056 012
BE		1MG	N04056 013
BE		5MG	N04056 014
	@	0.5MG	N04056 012
	@	1MG	N04056 013
	@	5MG	N04056 014

DIETHYLSTILBESTROL; METHYLTESTOSTERONE

TABLET; ORAL

	<u>TYLOSTERONE</u>		
*	LILLY	0.25MG; 5MG	N07661 001

DIETHYLSTILBESTROL; METHYLTESTOSTERONE

> DLT > TABLET; ORAL  
 > DLT > MYLESTERONE  
 > ADD > @ LILLY 0.25MG;5MG N07661 001

DILTIAZEM HYDROCHLORIDE

TABLET; ORAL  
AB DILTIAZEM HCL 30MG N74084 001  
 NOVOPHARM FEB 25, 1994  
AB 60MG N74084 002  
 FEB 25, 1994

DIPHENHYDRAMINE HYDROCHLORIDE

CAPSULE; ORAL  
AA DIPHENHYDRAMINE HCL 25MG N80868 001  
AA LASHNET 50MG N80868 002  
 @ 25MG N80868 001  
 @ 50MG N80868 002  
AA ROXANE 50MG N80635 001  
 @ 50MG N80635 001

DIPIVEFRIN HYDROCHLORIDE

SOLUTION/DROPS; OPHTHALMIC  
AT DIPIVEFRIN HCL 0.1% N73636 001  
 ALCON JUN 30, 1994  
AT + PROPINE 0.1% N18239 001  
 ALLERGAN

DOBUTAMINE HYDROCHLORIDE

INJECTABLE; INJECTION  
AP DOBUTAMINE HCL EQ 12.5MG BASE/ML N74277 001  
 BEDFORD OCT 31, 1994  
AP DOBUTAMINE HCL IN DEXTROSE 5% IN PLASTIC CONTAINER EQ 400MG BASE/100ML N20201 006  
 ABBOTT JUL 07, 1994

DOBUTAMINE HYDROCHLORIDE

INJECTABLE; INJECTION  
AP + DOBUTAMINE HCL IN DEXTROSE 5% IN PLASTIC CONTAINER EQ 400MG BASE/100ML N20255 005  
 BAXTER OCT 19, 1993

> ADD > DORZOLAMIDE HYDROCHLORIDE

> ADD > SOLUTION/DROPS; OPHTHALMIC  
 > ADD > TRUSOPT  
 > ADD > + MERCK EQ 2% BASE N20408 001  
 > ADD > DEC 09, 1994

DOXEPIN HYDROCHLORIDE

CAPSULE; ORAL  
 DOXEPIN HCL  
 @ RISOXS EQ 10MG BASE N16987 001  
 @ EQ 25MG BASE N16987 002  
 @ EQ 50MG BASE N16987 003  
 @ EQ 75MG BASE N16987 006  
 @ EQ 100MG BASE N16987 004  
 @ EQ 150MG BASE N16987 007  
 @ APR 13, 1987  
 @ LOTUS BIOCHEM EQ 10MG BASE N16987 001  
 @ EQ 25MG BASE N16987 002  
 @ EQ 50MG BASE N16987 003  
 @ EQ 75MG BASE N16987 006  
 @ EQ 100MG BASE N16987 004  
 @ EQ 150MG BASE N16987 007  
 @ APR 13, 1987

CREAM; TOPICAL

ZONALON  
 + GENDERM 5% N20126 001  
 APR 01, 1994

DOXORUBICIN HYDROCHLORIDE

INJECTABLE; INJECTION  
AP + ADRIAMYCIN PFS 200MG/100ML N50629 002  
 PHARMACIA MAY 03, 1988  
AP 200MG/100ML N63165 002  
 JAN 30, 1991

DOXORUBICIN HYDROCHLORIDE

INJECTABLE; INJECTION  
ADRIAMYCIN PFS

* PHARMACIA	200MG/100ML	N50629 002
		MAY 03, 1988
	200MG/100ML	N63165 002
		JAN 30, 1991

<u>AP</u>	<u>DOXORUBICIN HCL</u> CETUS BEN VENUE	200MG/100ML	N64097 001
			SEP 13, 1994

ENFLURANE

LIQUID; INHALATION  
ENFLURANE

<u>AN</u>	INHALON	99.9%	N74396 001
			JUL 29, 1994

EPINEPHRINE; LIDOCAINE HYDROCHLORIDE

INJECTABLE; INJECTION  
LIDOCAINE HCL AND EPINEPHRINE

<u>AP</u>	EASTMAN KODAK	0.01MG/ML; 2%	N40057 002
			FEB 26, 1993
		0.02MG/ML; 2%	N40057 001
			FEB 26, 1993

<u>AP</u>	STERLING WINTHROP	0.01MG/ML; 2%	N40057 002
			FEB 26, 1993
		0.02MG/ML; 2%	N40057 001
			FEB 26, 1993

ERGOCALCIFEROL

CAPSULE; ORAL

<u>AA</u>	<u>DELTALIN</u> LILLY	50,000 IU	N80884 001
	@	50,000 IU	N80884 001

<u>AA</u>	+ SANOFI WINTHROP	50,000 IU	N03444 001
<u>AA</u>	STERLING WINTHROP	50,000 IU	N03444 001

<u>AA</u>	<u>VITAMIN D</u> LANNETT	50,000 IU	N80825 001
	@	50,000 IU	N80825 001
<u>AA</u>	WEST WARD	50,000 IU	N83102 001
	@ WEST WARD PHARM	50,000 IU	N83102 001

ERYTHROMYCIN

CAPSULE, DELAYED REL PELLETS; ORAL  
ERYC

<u>AB</u>	FARKE DAVIS	250MG	N62546 001
	@	250MG	JUL 25, 1985
			N62546 001
			JUL 25, 1985

OINTMENT; OPHTHALMIC  
ERYTHROMYCIN

<u>AT</u>	BAUSCH AND LOMB	0.5%	N64067 001
			JUL 29, 1994
<u>AT</u>	FOUGERA	0.5%	N62447 001
			SEP 26, 1983
<u>AT</u>		5MG/GM	N62447 001
			SEP 26, 1983

ILOTYCIN

<u>AT</u>	+ DISTA	0.5%	N50368 001
<u>AT</u>	*	5MG/GM	N50368 001

SOLUTION; TOPICAL  
ERYTHROMYCIN

<u>AT</u>	BARRE	2%	N62957 001
	@	2%	JUL 21, 1988
			N62957 001
			JUL 21, 1988
<u>AT</u>	BAUSCH AND LOMB	2%	N64039 001
			JAN 27, 1994
<u>AT</u>	PHARMAFAIR	2%	N62616 001
	@	2%	JUL 25, 1985
			N62616 001
			JUL 25, 1985

TABLET, COATED PARTICLES; ORAL  
PCE

	ABBOTT	333MG	N50611 001
			SEP 09, 1986
			N50611 001
			SEP 09, 1986

ERYTHROMYCIN ETHYLSUCCINATE

GRANULE; ORAL  
E.E.S.

<u>AB</u>	* ABBOTT	EQ 200MG BASE/5ML	N50207 001
<u>AB</u>		EQ 200MG BASE/5ML	N50207 001
	ERYPED		
	ABBOTT	EQ 400MG BASE/5ML	N50207 002

ERYTHROMYCIN ETHYLSUCCINATE

GRANULE; ORAL  
ERYPED  
AB + ABBOTT EQ 200MG BASE/5ML N50207 003  
 MAR 30, 1987  
 + EQ 400MG BASE/5ML N50207 002

SUSPENSION; ORAL

ERYTHROMYCIN ETHYLSUCCINATE

AB DISTA EQ 200MG BASE/5ML N62177 001  
AB EQ 400MG BASE/5ML N62177 002  
 @ EQ 200MG BASE/5ML N62177 001  
 @ EQ 400MG BASE/5ML N62177 002  
AB PHARMAPAIR EQ 200MG BASE/5ML N62559 001  
 MAR 15, 1985  
AB EQ 400MG BASE/5ML N62558 001  
 MAR 15, 1985  
 @ EQ 200MG BASE/5ML N62559 001  
 MAR 15, 1985  
 @ EQ 400MG BASE/5ML N62558 001  
 MAR 15, 1985

TABLET; ORAL

E.E.S. 400

AB \* ABBOTT EQ 400MG BASE N61905 001  
AB EQ 400MG BASE N61905 002  
 AUG 12, 1982  
AB + EQ 400MG BASE N61905 002  
 AUG 12, 1982  
 @ EQ 400MG BASE N61905 001

ERYTHROMYCIN STEARATE

TABLET; ORAL

ERYPAR

AB PARKS DAVIS EQ 250MG BASE N62322 001  
 @ WARNER CHILCOTT EQ 250MG BASE N62322 001  
AB ERYTHROMYCIN STEARATE  
AB PUREPAC EQ 250MG BASE N61743 001  
 @ PUREPAC PHARM EQ 250MG BASE N61743 001

ESTRADIOL

FILM, EXTENDED RELEASE; TRANSDERMAL

> ADD >  
 > ADD > BX CLIMARA 3M 0.05MG/24HR N20375 001  
 > ADD > DEC 22, 1994

ESTRADIOL

FILM, EXTENDED RELEASE; TRANSDERMAL

> ADD >  
 > ADD > BX CLIMARA 3M 0.1MG/24HR N20375 002  
 > ADD > DEC 22, 1994  
 > ADD >  
 > ADD > BX + ESTRADERM CIBA 0.05MG/24HR N19081 002  
 > ADD > SEP 10, 1986  
 > ADD > BX + 0.1MG/24HR N19081 003  
 > ADD > SEP 10, 1986  
 > ADD >  
 > ADD > BX VIVELLE NOVEN 0.05MG/24HR N20323 002  
 > ADD > OCT 28, 1994  
 > ADD > BX 3 dm 0.1MG/24HR N20323 004  
 > ADD > OCT 28, 1994  
 > ADD > 0.0375MG/24HR N20323 001  
 > ADD > OCT 28, 1994  
 > ADD > 0.075MG/24HR N20323 003  
 > ADD > OCT 28, 1994

ESTRADIOL VALERATE; TESTOSTERONE ENANTHATE

INJECTABLE; INJECTION

DELADUMONE

AO \* SQUIBB 4MG/ML; 90MG/ML N09545 001  
 @ 4MG/ML; 90MG/ML N09545 001  
AO \* TESTOSTERONE ENANTHATE AND ESTRADIOL VALERATE  
STERIS 4MG/ML; 90MG/ML N85865 001  
 + 4MG/ML; 90MG/ML N85865 001

> ADD > ESTROGENS, CONJUGATED; MEDROXYPROGESTERONE ACETATE

> ADD > TABLET; ORAL-28

> ADD > PREMPHASE (PREMARIN; CYCRIN 14/14)  
 > ADD > + WYETH AYERST 0.625MG, 0.625MG; N/A, 5MG N20303 002  
 > ADD > DEC 30, 1994  
 > ADD >  
 > ADD > PREMPRO (PREMARIN; CYCRIN 14/14)  
 > ADD > WYETH AYERST 0.625MG, 0.625MG; 2.5MG,  
 > ADD > 2.5MG N20303 001  
 > ADD > DEC 30, 1994

ESTROGENS, ESTERIFIED

TABLET; ORAL

ESTRATAB

BS SOLVAY 0.625MG N83209 001

ESTROGENS, ESTERIFIED

TABLET; ORAL  
 ESTRATAB  
 BS + SOLVAY 0.625MG N83209 001  
 BS 2.5MG N83857 001  
 BS + 2.5MG N83857 001  
 MENEST  
 BS \* SMITHKLINE BEECHAM 0.625MG N84948 001  
 BS 0.625MG N84948 001  
 BS \* 2.5MG N84949 001  
 BS 2.5MG N84949 001

ESTRONE

INJECTABLE; INJECTION  
 ESTRONE  
 BS STERIS 2MG/ML N83397 001  
 @ 2MG/ML N83397 001

ETHAMBUTOL HYDROCHLORIDE

TABLET; ORAL  
 MYAMBUTOL  
 LEDEBIE 200MG N16320 002  
 400MG N16320 003  
 \* 500MG N16320 004  
 @ 200MG N16320 002  
 + 400MG N16320 003  
 @ 500MG N16320 004

ETHINAMATE

CAPSULE; ORAL  
 VALMID  
 DISTA 500MG N09750 001  
 @ 500MG N09750 001

ETHINYL ESTRADIOL; FERROUS FUMARATE; NORETHINDRONE

TABLET; ORAL-28  
 NORQUEST FE N18926 001  
 \* SYNTEX 0.035MG;75MG;1MG JUL 18, 1986

ETHINYL ESTRADIOL; FERROUS FUMARATE; NORETHINDRONE

TABLET; ORAL-28  
 NORQUEST FE N18926 001  
 @ SYNTEX 0.035MG;75MG;1MG JUL 18, 1986

ETHINYL ESTRADIOL; NORETHINDRONE

TABLET; ORAL-21  
 NORETHINDRONE AND ETHINYL ESTRADIOL (7/14)  
 WATSON 0.035MG,0.035MG,0.5MG,1MG N71041 001  
 SEP 24, 1991  
 AB WATSON LABS 0.035MG,0.035MG;0.5MG,1MG N71041 001  
 SEP 24, 1991  
 ORTHO-NOVUM 7/14-21  
 AB + JOHNSON RW 0.035MG,0.035MG;0.5MG,1MG N19004 001  
 APR 04, 1984  
 @ 0.035MG,0.035MG,0.5MG,1MG N19004 001  
 APR 04, 1984

TABLET; ORAL-28  
 NORETHINDRONE AND ETHINYL ESTRADIOL (7/14)  
 WATSON 0.035MG,0.035MG,0.5MG,1MG N71042 001  
 SEP 24, 1991  
 AB WATSON LABS 0.035MG,0.035MG;0.5MG,1MG N71042 001  
 SEP 24, 1991  
 ORTHO-NOVUM 7/14-28  
 AB + JOHNSON RW 0.035MG,0.035MG;0.5MG,1MG N19004 002  
 APR 04, 1984  
 @ 0.035MG,0.035MG,0.5MG,1MG N19004 002  
 APR 04, 1984

ETODOLAC

TABLET; ORAL  
 LODINE N18922 004  
 + WYETH AYERST 400MG JUL 29, 1993

ETOPOSIDE

INJECTABLE; INJECTION  
 ETOPOSIDE N74284 001  
 AP GENSIA 20MG/ML FEB 10, 1994

ETOPOSID

INJECTABLE; INJECTION  
VEPESID  
 AP + BRISTOL 20MG/ML N18768 001  
 NOV 10, 1983

FAMCICLOVIR

TABLET; ORAL  
 FAMVIR  
 + SMITHKLINE BEECHAM 500MG N20363 002  
 JUN 29, 1994

FAMOTIDINE

INJECTABLE; INJECTION  
 PEPCID IN PLASTIC CONTAINER  
 + MERCK 0.4MG/ML N20249 001  
 FEB 18, 1994

FELODIPINE

TABLET, EXTENDED RELEASE; ORAL  
 PLENDIL  
 + ASTRA MERCK 2.5MG N19834 004  
 SEP 22, 1994  
 + 5MG N19834 001  
 JUL 25, 1991  
 + 10MG N19834 002  
 JUL 25, 1991  
~~MERCK 5MG N19834 001~~  
~~JUL 25, 1991~~  
~~10MG N19834 002~~  
~~JUL 25, 1991~~

FLUCONAZOLE

TABLET; ORAL  
 DIFLUCAN  
 \* PFIZER 50MG N19949 001  
 JAN 29, 1990  
 \* 100MG N19949 002  
 JAN 29, 1990

FLUCONAZOLE

TABLET; ORAL  
 DIFLUCAN  
 PFIZER 50MG N19949 001  
 100MG N19949 002  
 150MG N20322 001  
 JUN 30, 1994

FLUDEOXYGLUCOSE, F-18

INJECTABLE; INJECTION  
 FLUDEOXYGLUCOSE F 18  
 + DOWNSTATE CLINCL 6.8-35.7mCi/ML N20306 001  
 AUG 19, 1994

FLUMAZENIL

INJECTABLE; INJECTION  
~~MAZICON~~  
~~\* ROCHE 0.1MG/ML N20073 001~~  
~~DEC 20, 1991~~  
 ROMAZICON  
 + ROCHE 0.1MG/ML N20073 001  
 DEC 20, 1991

FLUOCINOLONE ACETONIDE

CREAM; TOPICAL  
FLUOCINOLONE ACETONIDE  
 AT TARO PHARMS 0.025% N40042 001  
 OCT 31, 1994  
 AT 0.1% N40035 001  
 OCT 31, 1994

OINTMENT; TOPICAL

FLUOCINOLONE ACETONIDE  
 AT TARO PHARMS 0.025% N40041 001  
 SEP 15, 1994

SOLUTION; TOPICAL

FLUOCINOLONE ACETONIDE  
 AT PHARMAFAIR 0.01% N88449 001  
 FEB 08, 1984

FLUOCINOLONE ACETONIDE

SOLUTION; TOPICAL  
FLUOCINOLONE ACETONIDE  
@ PHARMAFAIR 0.01%

N88449 001  
FEB 08, 1984

FLUOCINONIDE

CREAM; TOPICAL  
FLUOCINONIDE  
B\* CLAY PARK 0.05%  
@ 0.05%  
AB FOUGERA 0.05%

N71790 001  
JUL 11, 1988  
N71790 001  
JUL 13, 1988  
N73030 001  
OCT 17, 1994

GEL; TOPICAL  
FLUOCINONIDE  
> ADD > AB FOUGERA 0.05%  
> ADD >

N72933 001  
DEC 30, 1994

FLUORESCEIN SODIUM

INJECTABLE; INJECTION  
FUNDUSCEIN-25  
+ CIBA VISION 25%  
\* IOLAB 25%

N17869 001  
N17869 001

FLUOROMETHOLONE

SUSPENSION/DROPS; OPHTHALMIC  
FLUOR-OP  
AB CIBA VISION 0.1%  
AB IOLAB 0.1%

N70185 001  
FEB 27, 1986  
N70185 001  
FEB 27, 1986

FLUOROURACIL

INJECTABLE; INJECTION  
ADRUCIL  
AP \* ADRIA 50MG/ML

N17959 001

FLUOROURACIL

INJECTABLE; INJECTION  
ADRUCIL

AP ADRIA 50MG/ML  
AP + PHARMACIA 50MG/ML  
@ 50MG/ML

N40023 001  
OCT 18, 1991  
N40023 001  
OCT 18, 1991  
N17959 001

FLURANDRENOLIDE

CREAM; TOPICAL  
CORDRAN SP  
\* DISTA 0.025%  
\* 0.05%  
+ LILLY 0.025%  
+ 0.05%

N12806 003  
N12806 002  
N12806 003  
N12806 002

LOTION; TOPICAL

CORDRAN  
AT \* DISTA 0.05%  
AT + LILLY 0.05%

N13790 001  
N13790 001

OINTMENT; TOPICAL

CORDRAN  
\* DISTA 0.025%  
\* 0.05%  
+ LILLY 0.025%  
+ 0.05%

N12806 004  
N12806 001  
N12806 004  
N12806 001

TAPE; TOPICAL

CORDRAN  
\* DISTA 0.004MG/SQ CM  
+ LILLY 0.004MG/SQ CM

N16455 001  
N16455 001

FLURBIPROFEN

TABLET; ORAL  
ANSAID  
AB UPJOHN 50MG  
AB + 100MG  
AB FLURBIPROFEN  
MYLAN 50MG

N18766 002  
OCT 31, 1988  
N18766 003  
OCT 31, 1988  
N74358 001  
JUN 20, 1994

FLURBIPROFEN

TABLET; ORAL  
FLURBIPROFEN  
AB MYLAN 100MG N74358 002  
 JUN 20, 1994

FLUTICASONE PROPIONATE

SPRAY, METERED; NASAL  
 FLONASE  
 + GLAXO 0.05MG/INH N20121 001  
 OCT 19, 1994

> ADD > FLUVOXAMINE MALEATE

> ADD > TABLET; ORAL  
 > ADD > LUVOX  
 > ADD > @ SOLVAY 25MG N20243 001  
 > ADD > DEC 05, 1994  
 > ADD > 50MG N20243 002  
 > ADD > DEC 05, 1994  
 > ADD > + N20243 003  
 > ADD > 100MG DEC 05, 1994  
 > ADD > @ N20243 004  
 > ADD > 150MG DEC 05, 1994

FOLIC ACID

TABLET; ORAL  
FOLIC ACID  
AA LANNETT 1MG N80816 001  
 @ 1MG N80816 001  
AA PUREPAC 1MG N80784 001  
 @ PUREPAC PHARM 1MG N80784 001

FOSINOPRIL SODIUM; HYDROCHLOROTHIAZIDE

TABLET; ORAL  
 MONOPRIL-HCT  
 BRISTOL MYERS SQUIBB 10MG;12.5MG N20286 002  
 NOV 30, 1994  
 + 20MG;12.5MG N20286 001  
 NOV 30, 1994

FUROSEMIDE

INJECTABLE; INJECTION  
FUROSEMIDE  
AP MARSAM 10MG/ML N74017 001  
 JUN 30, 1994  
AP ORGANON 10MG/ML N70017 001  
 DEC 15, 1986  
 @ N70017 001  
 10MG/ML DEC 15, 1986  
AP SANOFI WINTHROP 10MG/ML N74337 001  
 OCT 31, 1994  
AP WARNER CHILCOTT 10MG/ML N18420 001  
 FEB 26, 1982  
 @ N18420 001  
 10MG/ML FEB 26, 1982

GADODIAMIDE

INJECTABLE; INJECTION  
 OMNISCAN  
 NYCOMED 287MG/ML N20123 001  
 JAN 08, 1993  
 STERLING WINTHROP 287MG/ML N20123 001  
 JAN 08, 1993

GALLIUM CITRATE, GA-67

INJECTABLE; INJECTION  
 GALLIUM CITRATE GA 67  
 BS DUPONT 2mCi/ML N17478 001  
 BS DUPONT MERCK 2mCi/ML N17478 001

> ADD > GANCICLOVIR

> ADD > CAPSULE; ORAL  
 > ADD > CYTOVENE  
 > ADD > + SYNTEX 250MG N20460 001  
 > ADD > DEC 22, 1994

GEMFIBROZIL

TABLET; ORAL  
GEMFIBROZIL  
AB PUREPAC PHARM 600MG N74360 001  
 AUG 31, 1994

GEMFIBROZIL

TABLET; ORAL  
GEMFIBROZIL  
AB WATSON LABS 600MG N74156 001  
 OCT 24, 1994

GENTAMICIN SULFATE

CREAM; TOPICAL  
GARAMYCIN  
AT + SCHERING EQ 0.1% BASE N60462 001  
AT \* EQ 1MG BASE/GM N60462 001  
AT GENTAFAIR  
AT PHARMAFAIR EQ 1MG BASE/GM N62458 001  
 SEP 01, 1983  
 @ EQ 0.1% BASE N62458 001  
 SEP 01, 1983  
GENTAMICIN  
AT CLAY PARK EQ 0.1% BASE N62307 001  
AT EQ 1MG BASE/GM N62307 001  
AT GENTAMICIN SULFATE  
 BAUSCH AND LOMB EQ 0.1% BASE N64056 001  
 APR 29, 1994  
AT FOUGERA EQ 0.1% BASE N62531 001  
 JUL 05, 1984  
AT EQ 1MG BASE/GM N62531 001  
 JUL 05, 1984  
AT NMC EQ 0.1% BASE N62471 001  
 SEP 27, 1983  
AT EQ 1MG BASE/GM N62471 001  
 SEP 27, 1983  
AT THAMES EQ 0.1% BASE N62427 001  
 MAY 26, 1983  
AT EQ 1MG BASE/GM N62427 001  
 MAY 26, 1983

INJECTABLE; INJECTION

APOGEN  
AP KING PHARMS EQ 10MG BASE/ML N62289 001  
AP EQ 40MG BASE/ML N62289 002  
AP SMITHKLINE BEECHAM EQ 10MG BASE/ML N62289 001  
AP EQ 40MG BASE/ML N62289 002  
AT \* GARAMYCIN  
 SCHERING EQ 2MG BASE/ML N50505 001

INJECTABLE; INTRATHECAL

GARAMYCIN  
 + SCHERING EQ 2MG BASE/ML N50505 001

GENTAMICIN SULFATE

OINTMENT; OPHTHALMIC  
GARAMYCIN  
AT + SCHERING EQ 0.3% BASE N50425 001  
AT \* EQ 3MG BASE/GM N50425 001  
AT GENTACIDIN  
 IOLAB EQ 0.3% BASE N62501 001  
AT EQ 3MG BASE/GM N62501 001  
 JUL 26, 1984  
 JUL 26, 1984

OINTMENT; TOPICAL

GARAMYCIN  
AT + SCHERING EQ 0.1% BASE N60463 001  
AT \* EQ 1MG BASE/GM N60463 001  
AT GENTAFAIR  
 PHARMAFAIR EQ 0.1% BASE N62444 001  
 @ EQ 0.1% BASE N62444 001  
 MAY 26, 1983  
 MAY 26, 1983  
GENTAMICIN  
AT CLAY PARK EQ 0.1% BASE N62351 001  
AT EQ 1MG BASE/GM N62351 001  
 FEB 18, 1982  
 FEB 18, 1982

GENTAMICIN SULFATE

BAUSCH AND LOMB EQ 0.1% BASE N64054 001  
 APR 29, 1994  
AT FOUGERA EQ 0.1% BASE N62533 001  
AT EQ 1MG BASE/GM N62533 001  
 OCT 05, 1984  
 OCT 05, 1984  
AT NMC EQ 0.1% BASE N62496 001  
AT EQ 1MG BASE/GM N62496 001  
 MAR 14, 1984  
 MAR 14, 1984  
AT PHARMADERM EQ 0.1% BASE N62534 001  
AT EQ 1MG BASE/GM N62534 001  
 OCT 10, 1984  
 OCT 10, 1984  
AT THAMES EQ 0.1% BASE N62477 001  
AT EQ 1MG BASE/GM N62477 001  
 DEC 23, 1983  
 DEC 23, 1983

SOLUTION/DROPS; OPHTHALMIC

GARAMYCIN  
AT + SCHERING EQ 0.3% BASE N50039 002

GENTAMICIN SULFATE

SOLUTION/DROPS; OPHTHALMIC

<u>GARAMYCIN</u>			
<u>AT</u>	* SCHERING	<u>EQ 3MG BASE/ML</u>	N50039 002
<u>AT</u>	GENOPTIC	<u>EQ 0.3% BASE</u>	N62452 001
	ALLERGAN		OCT 10, 1984
<u>AT</u>		<u>EQ 3MG BASE/ML</u>	N62452 001
			OCT 10, 1984
<u>GENTACIDIN</u>			
<u>AT</u>	IOLAB	<u>EQ 0.3% BASE</u>	N62480 001
			MAR 30, 1984
<u>AT</u>		<u>EQ 3MG BASE/ML</u>	N62480 001
			MAR 30, 1984
GENTAFAIR			
	@ PHARMAFAIR	EQ 0.3% BASE	N62440 001
			MAY 03, 1983
<u>GENTAMICIN SULFATE</u>			
<u>AT</u>	AKORN	<u>EQ 0.3% BASE</u>	N62635 001
			JAN 08, 1987
<u>AT</u>		<u>EQ 3MG BASE/ML</u>	N62635 001
			JAN 08, 1987
<u>AT</u>	BAUSCH AND LOMB	<u>EQ 0.3% BASE</u>	N64048 001
			MAY 11, 1994
<u>AT</u>	STERIS	<u>EQ 0.3% BASE</u>	N62523 001
			NOV 25, 1985
<u>AT</u>		<u>EQ 3MG BASE/ML</u>	N62523 001
			NOV 25, 1985

GLIPIZIDE

TABLET; ORAL

<u>GLIPIZIDE</u>			
<u>AB</u>	CIRCA	<u>5MG</u>	N74370 001
			NOV 22, 1994
<u>AB</u>		<u>10MG</u>	N74370 002
			NOV 22, 1994
<u>AB</u>	ENDO LABS	<u>5MG</u>	N74378 001
			NOV 28, 1994
<u>AB</u>		<u>10MG</u>	N74378 002
			NOV 28, 1994
<u>AB</u>	MYLAN	<u>5MG</u>	N74226 001
			MAY 10, 1994
<u>AB</u>		<u>10MG</u>	N74226 002
			MAY 10, 1994
<u>AB</u>	<u>GLUCOTROL</u>	<u>5MG</u>	N17783 001
	PFIZER		MAY 08, 1984

GLIPIZIDE

TABLET; ORAL

<u>GLUCOTROL</u>			
<u>AB</u>	+ PFIZER	<u>10MG</u>	N17783 002
			MAY 08, 1984
TABLET, EXTENDED RELEASE; ORAL			
<u>GLUCOTROL XL</u>			
	+ PFIZER	5MG	N20329 001
			APR 26, 1994
		10MG	N20329 002
			APR 26, 1994
<u>GLUTETHIMIDE</u>			
TABLET; ORAL			
<u>GLUTETHIMIDE</u>			
<u>AA</u>	HALESY	<u>250MG</u>	N89458 001
			OCT 10, 1986
		250MG	N89458 001
			OCT 10, 1986
<u>AA</u>	LANNETT	<u>250MG</u>	N83475 001
<u>AA</u>		<u>500MG</u>	N85571 001
	@	250MG	N83475 001
	@	500MG	N85571 001

GLYBURIDE

TABLET; ORAL

<u>GLYNASE</u>			
	HEJOHN	<u>4.5MG</u>	N20051 003
			SEP 24, 1993
	@	4.5MG	N20051 003
			SEP 24, 1993

GLYCOPYRROLATE

INJECTABLE; INJECTION

<u>GLYCOPYRROLATE</u>			
<u>AP</u>	FOJISAWA	<u>0.2MG/ML</u>	N88475 001
			JUN 12, 1984
	@	0.2MG/ML	N88475 001
			JUN 12, 1984

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

GONADOTROPIN, CHORIONIC

INJECTABLE; INJECTION

A.P.L.

<u>AP</u>	* WYETH AYERST	<u>20,000 UNITS/VIAL</u>	N17055 003
	+	20,000 UNITS/VIAL	N17055 003
<u>AP</u>	<u>CHORIONIC GONADOTROPIN</u>	<u>5,000 UNITS/VIAL</u>	N17067 001
<u>AP</u>	FUKESAWA	<u>20,000 UNITS/VIAL</u>	N17067 003
	@	5,000 UNITS/VIAL	N17067 001
	@	20,000 UNITS/VIAL	N17067 003
<u>AP</u>	STERIS	<u>20,000 UNITS/VIAL</u>	N17016 004
	*	<u>15,000 UNITS/VIAL</u>	N17016 010
	@	15,000 UNITS/VIAL	FEB 15, 1985 N17016 010
	@	20,000 UNITS/VIAL	FEB 15, 1985 N17016 004

GRAMICIDIN; NEOMYCIN SULFATE; POLYMYXIN B SULFATE

SOLUTION/DROPS; OPHTHALMIC

NEOMYCIN SULFATE AND POLYMYXIN B SULFATE GRAMICIDIN

<u>AT</u>	PHARMAFAIR	<u>0.025MG/ML; EQ 1.75MG BASE/ML;</u>	N62383 001
		<u>10,000 UNITS/ML</u>	AUG 31, 1982
	@	0.025MG/ML; EQ 1.75MG BASE/ML;	N62383 001
		10,000 UNITS/ML	AUG 31, 1982

GRANISETRON HYDROCHLORIDE

INJECTABLE; INJECTION

KYTRIL

*	SMITHKLINE BEECHAM	<u>EQ 3MG BASE/ML</u>	N20239 001
			DEC 29, 1993
	+	EQ 1MG BASE/ML	N20239 002
			MAR 11, 1994
	@	EQ 3MG BASE/ML	N20239 001
			DEC 29, 1993

GUANABENZ ACETATE

TABLET; ORAL

GUANABENZ ACETATE

<u>AB</u>	COPLEY PHARM	<u>EQ 4MG BASE</u>	N74267 001
			JUN 01, 1994

GUANABENZ ACETATE

TABLET; ORAL

GUANABENZ ACETATE

<u>AB</u>	COPLEY PHARM	<u>EQ 8MG BASE</u>	N74267 002
			JUN 01, 1994
<u>AB</u>	WATSON LABS	<u>EQ 4MG BASE</u>	N74025 001
			FEB 28, 1994
<u>AB</u>		<u>EQ 8MG BASE</u>	N74025 002
			FEB 28, 1994
<u>AB</u>	<u>WYTENSIN</u>	<u>EQ 4MG BASE</u>	N18587 001
	WYETH AYERST		SEP 07, 1982
<u>AB</u>	+	<u>EQ 8MG BASE</u>	N18587 002
			SEP 07, 1982

HEPARIN CALCIUM

INJECTABLE; INJECTION

CALCIPARINE

+ CHOAY

\* DUPONT

25,000 UNITS/ML	N18237 001
25,000 UNITS/ML	N18237 001

HEPARIN SODIUM

INJECTABLE; INJECTION

HEPARIN SODIUM

<u>AP</u>	AKORN	<u>1,000 UNITS/ML</u>	N17486 001
<u>AP</u>		<u>5,000 UNITS/ML</u>	N17486 002
<u>AP</u>		<u>10,000 UNITS/ML</u>	N17486 003
<u>AP</u>		<u>20,000 UNITS/ML</u>	N17486 004
<u>AP</u>		<u>40,000 UNITS/ML</u>	N17486 005
	@	1,000 UNITS/ML	N17486 001
	@	5,000 UNITS/ML	N17486 002
	@	10,000 UNITS/ML	N17486 003
	@	20,000 UNITS/ML	N17486 004
	@	40,000 UNITS/ML	N17486 005

HEXACHLOROPHENE

EMULSION; TOPICAL

PHISOEX

+ SANOFI WINTHROP

@

\* STERLING WINTHROP

@

3%	N06882 001
3%	N08402 001
3%	N06882 001
3%	N08402 001

HEXACHLOROPHENE

SPONGE; TOPICAL  
PHISO-SCRUB  
@ SANOFI WINTHROP 3% N17446 001  
@ STERLING WINTHROP 3% N17446 001

HYDRALAZINE HYDROCHLORIDE

TABLET; ORAL  
HYDRALAZINE HCL  
AA AMIDE PHARM 25MG N88560 001  
OCT 04, 1984  
AA 50MG N88649 001  
OCT 18, 1984  
@ 25MG N88560 001  
OCT 04, 1984  
@ 50MG N88649 001  
OCT 18, 1984

HYDRALAZINE HYDROCHLORIDE; HYDROCHLOROTHIAZIDE; RESERPINE

TABLET; ORAL  
RESERPINE, HYDRALAZINE HCL AND HYDROCHLOROTHIAZIDE  
BP DANBURY PHARMA 25MG;15MG;0.1MG N87556 001  
@ 25MG;15MG;0.1MG N87556 001

HYDROCHLOROTHIAZIDE

TABLET; ORAL  
HYDROCHLOROTHIAZIDE  
AB WEST WARD 25MG N84899 001  
@ WEST WARD PHARM 25MG N84899 001  
AB ZENITH 50MG N84658 001  
@ ZENITH LABS 50MG N84658 001

HYDROCHLOROTHIAZIDE; LABETALOL HYDROCHLORIDE

TABLET; ORAL  
NORMOZIDE  
SCHERING 25MG;100MG N19046 001  
APR 05, 1987  
25MG;200MG N19046 002  
APR 06, 1987

HYDROCHLOROTHIAZIDE; LABETALOL HYDROCHLORIDE

TABLET; ORAL  
NORMOZIDE  
\* SCHERING 25MG;300MG N19046 003  
APR 06, 1987  
@ 25MG;100MG N19046 001  
APR 06, 1987  
@ 25MG;200MG N19046 002  
APR 06, 1987  
@ 25MG;300MG N19046 003  
APR 06, 1987

HYDROCHLOROTHIAZIDE; RESERPINE

TABLET; ORAL  
HYDROCHLOROTHIAZIDE W/ RESERPINE  
BP ROXANE 50MG;0.125MG N84603 001  
@ 50MG;0.125MG N84603 001

HYDROCHLOROTHIAZIDE; TRIAMTERENE

CAPSULE; ORAL  
DYAZIDE  
AB \* SMITHKLINE BEECHAM 25MG;50MG N16042 002  
25MG;37.5MG N16042 003  
MAR 03, 1994  
@ 25MG;50MG N16042 002  
AB TRIAMTERENE AND HYDROCHLOROTHIAZIDE  
GENEVA PHARMS 25MG;50MG N73191 001  
JUL 31, 1991  
+ 25MG;50MG N73191 001  
JUL 31, 1991

HYDROCORTISONE

CREAM; TOPICAL  
HYDROCORTISONE  
AT BENSON 1% N85191 001  
@ 1% N85191 001  
AT PHARMAFAIR 1% N87838 001  
JUL 28, 1982  
@ 1% N87838 001  
JUL 28, 1982  
ENEMA; RECTAL  
CORTENEMA  
+ SOLVAY 100MG/60ML N16199 001

HYDROCORTISONE  
 ENEMA; RECTAL  
HYDROCORTISONE  
 AT COPLEY PHARM 100MG/60ML N74171 001  
 MAY 27, 1994

GEL; TOPICAL  
HYTRACORT  
 AT \* GALDERMA 1% N84698 001  
 @ 1% N84698 001

PENECORT  
 AT ALLERGAN HERBERT 1% N88215 001  
 JUN 06, 1984  
 + 1% N88215 001  
 JUN 06, 1984

LOTION; TOPICAL  
HYDROCORTISONE  
 AT CLAY PARK 1% N85663 001  
 @ 1% N85663 001

OINTMENT; TOPICAL  
HYDROCORTISONE  
 AT CLAY PARK 1% N85028 001  
 @ 1% N85028 001

TABLET; ORAL  
 HYDROCORTISONE  
 BP DANBURY PHARMA 20MG N80355 001  
 @ 20MG N80355 001  
 BP INWOOD 20MG N80732 001  
 @ INWOOD LABS 20MG N80732 001

HYDROCORTISONE; NEOMYCIN SULFATE; POLYMYXIN B SULFATE  
 SOLUTION/DROPS; OTIC  
NEOMYCIN SULFATE-POLYMYXIN B SULFATE-HYDROCORTISONE  
 AT PHARMAFAIR 1%;EQ 3.5MG BASE/ML; N62394 001  
 10,000 UNITS/ML SEP 29, 1982  
 @ 1%;EQ 3.5MG BASE/ML; N62394 001  
 10,000 UNITS/ML SEP 29, 1982

SUSPENSION; OTIC  
OTICAIR  
 AT PHARMAFAIR 1%;EQ 3.5MG BASE/ML; N62399 001  
 10,000 UNITS/ML NOV 18, 1982

HYDROCORTISONE; NEOMYCIN SULFATE; POLYMYXIN B SULFATE  
 SUSPENSION; OTIC  
OTICAIR  
 @ PHARMAFAIR 1%;EQ 3.5MG BASE/ML; N62399 001  
 10,000 UNITS/ML NOV 18, 1982

HYDROCORTISONE ACETATE  
 CREAM; TOPICAL  
HYDROCORTISONE ACETATE  
 AT PARKE DAVIS 1% N89914 001  
 @ 1% N89914 001  
 JAN 03, 1989  
 N89914 001  
 JAN 03, 1989

INJECTABLE; INJECTION  
 HYDROCORTISONE ACETATE  
 BP STERIS 50MG/ML N85214 001  
 @ 50MG/ML N85214 001

HYDROCORTISONE ACETATE; UREA  
 CREAM; TOPICAL  
CARMOL HC  
 AT + KENWOOD LABS 1%;10% N80505 001  
 AT \* SYNTEX 1%;10% N80505 001

HYDROCORTISONE BUTYRATE  
 CREAM; TOPICAL  
 LOCID  
 \* BROCADES PHARMA 0.1% N18514 001  
 MAR 31, 1982  
 + YAMANOUCHI 0.1% N18514 001  
 MAR 31, 1982

OINTMENT; TOPICAL  
 LOCID  
 @ BROCADES PHARMA 0.1% N18652 001  
 OCT 29, 1982  
 @ YAMANOUCHI 0.1% N18652 001  
 OCT 29, 1982

HYDROCORTISONE BUTYRATE

SOLUTION; TOPICAL  
LOCOID

* BROCADES PHARMA	0.1%	N19116 001
		FEB 25, 1987
+ YAMANOUCHI	0.1%	N19116 001
		FEB 25, 1987

HYDROCORTISONE SODIUM SUCCINATE

INJECTABLE; INJECTION

A-HYDROCORT

<u>AP</u>	ABBOTT	<u>EQ 250MG BASE/VIAL</u>	N89578 001
			APR 11, 1989
<u>AP</u>		<u>EQ 500MG BASE/VIAL</u>	N89579 001
			APR 11, 1989
<u>AP</u>		<u>EQ 1GM BASE/VIAL</u>	N89580 001
			APR 11, 1989
@		EQ 250MG BASE/VIAL	N89578 001
			APR 11, 1989
@		EQ 500MG BASE/VIAL	N89579 001
			APR 11, 1989
@		EQ 1GM BASE/VIAL	N89580 001
			APR 11, 1989

HYDROMORPHONE HYDROCHLORIDE

INJECTABLE; INJECTION  
DILAUDID-HP

+ KNOLL PHARM	250MG/VIAL	N19034 002
		AUG 04, 1994

HYDROXYCHLOROQUINE SULFATE

TABLET; ORAL

HYDROXYCHLOROQUINE SULFATE

<u>AB</u>	COPLBY PHARM	<u>200MG</u>	N40081 001
			SEP 30, 1994
<u>AB</u>	+ SANOFI WINTHROP	<u>200MG</u>	N09768 001
<u>AB</u>	* STERLING WINTHROP	<u>200MG</u>	N09768 001

HYDROXYPROGESTERONE CAPROATE

INJECTABLE; INJECTION

DELALGIN

<u>AO</u>	* SQUIBB	<u>125MG/ML</u>	N10347 004
<u>AO</u>		<u>125MG/ML</u>	N16911 001
<u>AO</u>	*	<u>250MG/ML</u>	N10347 002
<u>AO</u>		<u>250MG/ML</u>	N16911 002
@		125MG/ML	N10347 004
@		125MG/ML	N16911 001
@		250MG/ML	N10347 002
@		250MG/ML	N16911 002

HYDROXYPROGESTERONE CAPROATE

<u>AO</u>	AKORN	<u>125MG/ML</u>	N18004 001
		125MG/ML	N18004 001
<u>AO</u>	* STERIS	<u>125MG/ML</u>	N17439 001
<u>AO</u>		<u>250MG/ML</u>	N17439 002
@		125MG/ML	N17439 001
+		250MG/ML	N17439 002

HYDROXYZINE HYDROCHLORIDE

INJECTABLE; INJECTION

HYDROXYZINE HCL

<u>AP</u>	SMITH KEPHEW SOLOPAK	<u>25MG/ML</u>	N87591 001
<u>AP</u>		<u>50MG/ML</u>	N87593 001
<u>AP</u>		<u>50MG/ML</u>	N87595 001
<u>AP</u>	SOLOPAK	<u>25MG/ML</u>	N87591 001
<u>AP</u>		<u>50MG/ML</u>	N87593 001
<u>AP</u>		<u>50MG/ML</u>	N87595 001
<u>AP</u>	<u>ORGATEX</u>	<u>25MG/ML</u>	N87014 001
<u>AP</u>	ORGANON	<u>50MG/ML</u>	N87014 002
@		25MG/ML	N87014 001
@		50MG/ML	N87014 002

SYRUP; ORAL

HYDROXYZINE HCL

<u>AA</u>	HI TECH PHARMA	<u>10MG/5ML</u>	N40010 001
			OCT 28, 1994

TABLET; ORAL

HYDROXYZINE HCL

<u>AB</u>	AMIDE PHARM	<u>10MG</u>	N89071 001
			JUL 22, 1986
<u>AB</u>		<u>25MG</u>	N89072 001
			JUL 22, 1986

HYDROXYZINE HYDROCHLORIDE

TABLET; ORAL			
<u>HYDROXYZINE HCL</u>			
<u>AB</u>	<u>AMIDE PHARM</u>	<u>50MG</u>	<u>N89073 001</u>
	@	10MG	JUL 22, 1986
	@	25MG	N89071 001
	@	50MG	JUL 22, 1986
<u>AB</u>	<u>ROYCE LABS</u>	<u>10MG</u>	N89072 001
			JUL 22, 1986
<u>AB</u>		<u>25MG</u>	N89073 001
			JUL 22, 1986
<u>AB</u>		<u>50MG</u>	N81149 001
			MAR 18, 1994
			N81150 001
			MAR 18, 1994
			N81151 001
			MAR 18, 1994

IBUPROFEN

TABLET; ORAL			
<u>MOTRIN</u>			
	<u>MCNEIL CONS PRODS</u>	<u>100MG</u>	<u>N20418 001</u>
			NOV 16, 1994
TABLET, CHEWABLE; ORAL			
<u>MOTRIN</u>			
	<u>MCNEIL CONS PRODS</u>	<u>50MG</u>	<u>N20135 001</u>
			NOV 16, 1994
	<u>+</u>	<u>100MG</u>	<u>N20135 002</u>
			NOV 16, 1994

IMIGLUCERASE

INJECTABLE; INJECTION			
<u>CEREZYME</u>			
	<u>+ GENZYME</u>	<u>200 UNITS/VIAL</u>	<u>N20367 001</u>
			MAY 23, 1994

INDIUM IN-111 PENTETREOTIDE KIT

INJECTABLE; INJECTION			
<u>OCTREOSCAN</u>			
	<u>MALLINCKRODT</u>	<u>3mCi/ML</u>	<u>N20314 001</u>
			JUN 02, 1994

INDOMETHACIN

CAPSULE; ORAL			
<u>INDOMETHACIN</u>			
<u>AB</u>	<u>CHELSEA</u>	<u>25MG</u>	<u>N18690 001</u>
			JUL 31, 1984
<u>AB</u>		<u>50MG</u>	<u>N18690 002</u>
			JUL 31, 1984
	@ <u>CHELSEA LABS</u>	25MG	N18690 001
	@	50MG	JUL 31, 1984
			N18690 002
			JUL 31, 1984

INSULIN BIOSYNTHETIC HUMAN

INJECTABLE; INJECTION			
<u>HUMULIN R</u>			
	<u>+ LILLY</u>	<u>500 UNITS/ML</u>	<u>N18780 004</u>
			MAR 31, 1994

INVERT SUGAR

INJECTABLE INJECTION			
<u>TRAVERT 10% IN PLASTIC CONTAINER</u>			
	<u>BAXTER</u>	<u>10GM/100ML</u>	<u>N16717 001</u>
	@	<u>10GM/100ML</u>	<u>N16717 001</u>

IOBENGUANE SULFATE I 131

INJECTABLE; INJECTION			
<u>IOBENGUANE SULFATE I 131</u>			
	<u>CIS</u>	<u>2.3mCi/ML</u>	<u>N20084 001</u>
			MAR 25, 1994

IODAMIDE MEGLUMINE

INJECTABLE; INJECTION			
<u>RENOVUE-65</u>			
	<u>+ BRACCO DXS</u>	<u>65%</u>	<u>N17902 001</u>
	<u>* SQUIBB</u>	<u>65%</u>	<u>N17902 001</u>

IODIPAMIDE MEGLUMINE

INJECTABLE; INJECTION  
 CHOLOGRAFIN MEGLUMINE  
 + BRACCO DXS  
 + SQUIBB

10.3%  
 52%  
 10.3%  
 52%

N09321 007  
 N09321 003  
 N09321 007  
 N09321 003

IODIPAMIDE SODIUM

INJECTABLE; INJECTION  
 CHOLOGRAFIN SODIUM  
 @ BRACCO DXS  
 @ SQUIBB

20%  
 20%

N09321 001  
 N09321 001

IODOHIPPURATE SODIUM, I-123

INJECTABLE; INJECTION  
 NEPHROFLOW  
 MEDI PHYSICS  
 @

1mCi/ML  
 1mCi/ML

N18289 001  
 DEC 28, 1984  
 N18289 001  
 DEC 28, 1984

IODOXAMATE MEGLUMINE

INJECTABLE; INJECTION  
 CHOLOVUE  
 @ BRACCO DXS  
 @ SQUIBB

9.9%  
 40.3%  
 9.9%  
 40.3%

N18077 001  
 N18076 001  
 N18077 001  
 N18076 001

IOHEXOL

INJECTABLE; INJECTION  
 OMNIPAQUE 140  
 + NYCOMED  
 \* STERLING WINTHROP  
 OMNIPAQUE 210  
 + NYCOMED

30.2%  
 30.2%  
 45.3%

N18956 005  
 NOV 30, 1988  
 N18956 005  
 NOV 30, 1988  
 N18956 006  
 JUN 30, 1989

IOHEXOL

INJECTABLE; INJECTION  
 OMNIPAQUE 210  
 \* STERLING WINTHROP

45.3%

N18956 006  
 JUN 30, 1989

SOLUTION; INJECTION, ORAL  
 OMNIPAQUE 350  
 + NYCOMED

75.5%

N18956 004  
 DEC 26, 1985  
 N18956 004  
 DEC 26, 1985

\* STERLING WINTHROP

75.5%

SOLUTION; INJECTION, ORAL, RECTAL  
 OMNIPAQUE 180  
 + NYCOMED

38.8%

N18956 001  
 DEC 26, 1985  
 N18956 001  
 DEC 26, 1985

\* STERLING WINTHROP

38.8%

OMNIPAQUE 240  
 + NYCOMED

51.8%

N18956 002  
 DEC 26, 1985  
 N18956 002  
 DEC 26, 1985

\* STERLING WINTHROP

51.8%

OMNIPAQUE 300  
 + NYCOMED

64.7%

N18956 003  
 DEC 26, 1985  
 N18956 003  
 DEC 26, 1985

\* STERLING WINTHROP

64.7%

SOLUTION; URETHRAL  
 OMNIPAQUE 70  
 + NYCOMED

15.1%

N18956 007  
 JUN 01, 1994  
 N18956 007  
 JUN 01, 1994

\* STERLING WINTHROP

15.1%

IOPAMIDOL

INJECTABLE; INTRAVASCULAR  
 ISOVUE-200  
 BRACCO DXS  
 ISOVUE-250  
 BRACCO DXS

41%  
 51%

N20327 001  
 OCT 12, 1994  
 N20327 002  
 OCT 12, 1994

IOPAMIDOL

INJECTABLE; INTRAVASCULAR

ISOVUE-300

BRACCO DXS 61%

N20327 003  
OCT 12, 1994

ISOVUE-370

BRACCO DXS 76%

N20327 004  
OCT 12, 1994

IOPANOIC ACID

TABLET; ORAL

TELEPAQUE

NYCOMED 500MG

STERLING WINTHROP 500MG

N08032 001  
N08032 001

IPODATE CALCIUM

GRANULE; ORAL

ORAGRAFIN CALCIUM

BRACCO DXS 3GM/PACKET

SQUIBB 3GM/PACKET

N12968 001  
N12968 001

IPODATE SODIUM

CAPSULE; ORAL

ORAGRAFIN SODIUM

AA BRACCO DXS 500MG

AA SQUIBB 500MG

N12967 001  
N12967 001

ISOETHARINE HYDROCHLORIDE

SOLUTION; INHALATION

BRONKOSOL

AN + SANOFI WINTHROP 1%

@ 0.25%

AN + STERLING WINTHROP 1%

@ 0.25%

N12339 008  
N12339 009  
N12339 008  
N12339 009

> DLT >

> DLT >

> ADD >

ISOETHARINE MESYLATE

AEROSOL, METERED; INHALATION

BRONKOMETER

+ SANOFI WINTHROP 0.34MG/INH

N12339 007

ISOETHARINE MESYLATE

AEROSOL, METERED; INHALATION

BRONKOMETER

\* STERLING WINTHROP 0.34MG/INH

N12339 007

ISOFLURANE

LIQUID; INHALATION

ISOFLURANE

AN INHALON 99.9%

N74416 001  
SEP 30, 1994

ISONIAZID; PYRAZINAMIDE; RIFAMPIN

TABLET; ORAL

RIFATER

+ MARION MERRELL DOW 50MG;300MG;120MG

N50705 001  
MAY 31, 1994

ISOPROTERENOL HYDROCHLORIDE

AEROSOL, METERED; INHALATION

ISUPREL

+ SANOFI WINTHROP 0.131MG/INH

\* STERLING WINTHROP 0.131MG/INH

N11178 001  
N11178 001

INJECTABLE; INJECTION

ISUPREL

AP + SANOFI WINTHROP 0.2MG/ML

AE \* STERLING WINTHROP 0.2MG/ML

N10515 001  
N10515 001

SOLUTION; INHALATION

AEROLONE

\* LILLY 0.25%

@ 0.25%

ISUPREL

+ SANOFI WINTHROP 0.5%

+ 1%

\* STERLING WINTHROP 0.5%

\* 1%

N07245 001  
N07245 001

N06327 002  
N06327 003  
N06327 002  
N06327 003

TABLET, RECTAL, SUBLINGUAL

ISUPREL

@ SANOFI WINTHROP 10MG

@ 15MG

N06328 001  
N06328 002



> ADD > LAMOTRIGINE  
 > ADD > TABLET; ORAL  
 > ADD > LAMICTAL  
 > ADD > @ BURROUGHS WELLCOME 50MG N20241 006  
 DEC 27, 1994  
 > ADD > 100MG N20241 001  
 DEC 27, 1994  
 > ADD > 150MG N20241 002  
 DEC 27, 1994  
 > ADD > + 200MG N20241 003  
 DEC 27, 1994  
 > ADD > @ 250MG N20241 004  
 DEC 27, 1994

LEUCOVORIN CALCIUM

INJECTABLE; INJECTION  
LEUCOVORIN CALCIUM  
 AP ELKINS SINN EQ 100MG BASE/VIAL N81224 001  
 JUN 03, 1994  
 \* IMMUMEX EQ 3MG BASE/ML N08107 001  
 @ EQ 3MG BASE/ML N08107 001

LEUPROLIDE ACETATE

INJECTABLE; INJECTION  
 LUPRON DEPOT-PED  
 > DLT > \* TAP 3.75MG/VIAL, 7.5MG/VIAL N20263 003  
 APR 16, 1993  
 > DLT > \* 7.5MG/VIAL, 7.5MG/VIAL N20263 004  
 APR 16, 1993  
 > ADD > @ TAP PHARMS 3.75MG/VIAL, 7.5MG/VIAL N20263 003  
 APR 16, 1993  
 > ADD > @ 7.5MG/VIAL, 7.5MG/VIAL N20263 004  
 APR 16, 1993  
 > ADD > + 11.25MG/VIAL N20263 005  
 JAN 21, 1994  
 > ADD > + 15MG/VIAL N20263 006  
 JAN 21, 1994

LEVOBUNOLOL HYDROCHLORIDE

SOLUTION/DROPS; OPHTHALMIC  
BETAGAN  
 AT + ALLERGAN 0.25% N19814 001  
 JUN 28, 1989

LEVOBUNOLOL HYDROCHLORIDE

SOLUTION/DROPS; OPHTHALMIC  
BETAGAN  
 AT + ALLERGAN 0.5% N19219 002  
 DEC 19, 1985  
LEVOBUNOLOL HCL  
 AT BAUSCH AND LOMB 0.25% N74307 001  
 MAR 04, 1994  
 AT 0.5% N74326 001  
 MAR 04, 1994

LEVOCABASTINE HYDROCHLORIDE

SUSPENSION/DROPS; OPHTHALMIC  
 LIVOSTIN  
 + CIBA VISION EQ 0.05% BASE N20219 001  
 NOV 10, 1993  
 \* IOLAB EQ 0.05% BASE N20219 001  
 NOV 10, 1993

LEVONORDEFRIN; MEPIVACAINE HYDROCHLORIDE

INJECTABLE; INJECTION  
ARESTOCAINE HCL W/ LEVONORDEFRIN  
 AP CARLISLE 0.05MG/ML, 2% N85010 001  
 @ SOLVAY 0.05MG/ML, 2% N85010 001  
CARBOCAINE W/ NEO-COBEFRIN  
 AP \* COOK WHITE 0.05MG/ML, 2% N12125 002  
 AP + EASTMAN KODAK 0.05MG/ML, 2% N12125 002

LEVONORDEFRIN; PROCAINE HYDROCHLORIDE; PROPOXYCAINE HYDROCHLORIDE

INJECTABLE; INJECTION  
 RAVOCAINE AND NOVOCAIN W/ NEO-COBEFRIN  
 @ EASTMAN KODAK 0.05MG/ML, 2%; 0.4% N08592 007  
 \* STERLING WINTHROP 0.05MG/ML, 2%; 0.4% N08592 007

LIDOCAINE HYDROCHLORIDE

SOLUTION; TOPICAL  
MYLOCAINE  
 AT PENNEX 4% N87881 001  
 NOV 18, 1982

LIDOCAINE HYDROCHLORIDE

SOLUTION; TOPICAL  
MYLOCAINE  
 © PENNEX PHARMS 4% N87881 001  
 NOV 18, 1982  
PEDIATRIC LTA KIT  
 AT ABBOTT 2% N88572 001  
 JUL 31, 1984  
 © N88572 001  
 JUL 31, 1984

LIOTRIX (T4;T3)

TABLET; ORAL  
EUTHROID-0.5  
 PARKE DAVIS 0.03MG;0.0075MG N16680 001  
 © 0.03MG;0.0075MG N16680 001  
EUTHROID-1  
 PARKE DAVIS 0.06MG;0.015MG N16680 002  
 © 0.06MG;0.015MG N16680 002  
EUTHROID-2  
 PARKE DAVIS 0.12MG;0.03MG N16680 003  
 © 0.12MG;0.03MG N16680 003  
EUTHROID-3  
 \* PARKE DAVIS 0.18MG;0.045MG N16680 004  
 © 0.18MG;0.045MG N16680 004

LISINAPRIL

TABLET; ORAL  
PRINIVIL  
 AB MERCK 2.5MG N19558 006  
 JAN 28, 1994  
ZESTRIL  
 AB ZENECA 2.5MG N19777 005  
 APR 29, 1993

LITHIUM CARBONATE

TABLET, EXTENDED RELEASE; ORAL  
LITHOBID  
 © CIBA 300MG N18027 001  
 © SOLVAY 300MG N18027 001

LITHIUM CITRATE

SYRUP; ORAL  
CIBALITH-S  
 AP CIBA EQ 300MG CARBONATE/5ML N17672 001  
 AA SOLVAY EQ 300MG CARBONATE/5ML N17672 001

LORATADINE; PSEUDOEPHEDRINE SULFATE

TABLET, EXTENDED RELEASE; ORAL  
 CLARITIN-D  
 + SCHERING 5MG;120MG N19670 001  
 NOV 14, 1994

LORAZEPAM

INJECTABLE; INJECTION  
ATIVAN  
 AP + WYETH AYERST 2MG/ML N18140 001  
 AP + 4MG/ML N18140 002  
LORAZEPAM  
 AP ABBOTT 2MG/ML N74280 001  
 MAY 27, 1994  
 AP 2MG/ML N74282 001  
 MAY 27, 1994  
 AP 4MG/ML N74280 002  
 MAY 27, 1994  
 AP 4MG/ML N74282 002  
 MAY 27, 1994  
 AP 2MG/ML N74276 001  
 APR 15, 1994  
 AP 4MG/ML N74276 002  
 APR 15, 1994  
 AP 2MG/ML N74243 001  
 APR 12, 1994  
 AP 2MG/ML N74300 001  
 APR 12, 1994  
 AP 4MG/ML N74243 002  
 APR 12, 1994

TABLET; ORAL

LORAZEPAM  
 AB WARNER CHILCOTT 1MG N71038 001  
 AB 2MG N71039 001  
 JAN 12, 1988

LORAZEPAM

TABLET; ORAL  
LORAZEPAM  
 @ WARNER CHILCOTT 1MG N71038 001  
 JAN 12, 1988  
 @ 2MG N71039 001  
 JAN 12, 1988

MAGNESIUM SULFATE

INJECTABLE; INJECTION  
 MAGNESIUM SULFATE IN PLASTIC CONTAINER  
 ABBOTT 4GM/100ML N20309 001  
 JUN 24, 1994  
 80MG/ML N20309 002  
 JUN 24, 1994

MEDROXYPROGESTERONE ACETATE

INJECTABLE; INJECTION  
 DEPO-PROVERA  
 UPJOHN 150MG/ML N20246 001  
 OCT 29, 1992  
 + 150MG/ML N20246 001  
 OCT 29, 1992

TABLET; ORAL  
CYCRIN

AB ESI 2.5MG N81239 001  
 OCT 30, 1992  
AB 5MG N81240 001  
 OCT 30, 1992  
AB 10MG N89386 001  
 SEP 09, 1987  
AB WYETH AYERST 2.5MG N81239 001  
 OCT 30, 1992  
AB 5MG N81240 001  
 OCT 30, 1992  
AB 10MG N89386 001  
 SEP 09, 1987

MENOTROPINS (FSH;LH)

INJECTABLE; INJECTION

HUMEGON  
AB ORGANON 75 IU/VIAL;75 IU/VIAL N20328 001  
 SEP 01, 1994  
AB 150 IU/VIAL;150 IU/VIAL N20328 002  
 SEP 01, 1994  
~~75 IU/VIAL;75 IU/VIAL N20328 001~~  
~~150 IU/VIAL;150 IU/VIAL N20328 002~~  
~~SEP 01, 1994~~

PERGONAL  
AB + SERONO 75 IU/AMP;75 IU/AMP N17646 001  
AB + 150 IU/AMP;150 IU/AMP N17646 002  
 MAY 20, 1985  
 + SERONO LABS 75 IU/AMP;75 IU/AMP N17646 001  
 \* 150 IU/AMP;150 IU/AMP N17646 002  
 MAY 20, 1985

MEPIVACAINE HYDROCHLORIDE

INJECTABLE; INJECTION

ARESTOCAINE HCL  
AP CARLISLE 3% N84777 002  
 APR 18, 1982  
 @ SOLVAY 3% N84777 002  
 APR 18, 1982

CARBOCAINE  
AP \* COOK WALTER 3% N12125 003  
AP + EASTMAN KODAK 3% N12125 003

MEPROBAMATE

TABLET; ORAL

MEPRIAM  
AP LEMMON 400MG N16069 001  
 @ 400MG N16069 001

METAPROTERENOL SULFATE

SOLUTION; INHALATION

METAPROTERENOL SULFATE  
AN DEX 5% N78805 001  
 > DLT > AUG 17, 1987  
 > DLT >



METHYLPREDNISOLONE SODIUM SUCCINATE

INJECTABLE; INJECTION

A-METHAPRED

<u>AP</u>	<u>ABBOTT</u>	<u>EQ 125MG BASE/VIAL</u>	<u>N89574 001</u>
			FEB 22, 1991
<u>AP</u>		<u>EQ 500MG BASE/VIAL</u>	<u>N89575 001</u>
			FEB 22, 1991
	@	EQ 40MG BASE/VIAL	N89573 001
			FEB 22, 1991
	@	EQ 125MG BASE/VIAL	N89574 001
			FEB 22, 1991
	@	EQ 500MG BASE/VIAL	N89575 001
			FEB 22, 1991

METHYLPREDNISOLONE

<u>AP</u>	<u>ORGANON</u>	<u>EQ 500MG BASE/VIAL</u>	<u>N87535 001</u>
			JUN 25, 1982
<u>AP</u>		<u>EQ 1GM BASE/VIAL</u>	<u>N87535 002</u>
			JUN 25, 1982
	@	EQ 500MG BASE/VIAL	N87535 001
			JUN 25, 1982
	@	EQ 1GM BASE/VIAL	N87535 002
			JUN 25, 1982

METHYLTESTOSTERONE

TABLET; BUCCAL/SUBLINGUAL

METHYLTESTOSTERONE

		<u>PRIVATE FORM</u>	<u>5MG</u>	<u>N83836 003</u>
	@	PVT FORM	5MG	N83836 001
<u>BP</u>		<u>TABLICAPS</u>	<u>10MG</u>	<u>N85125 001</u>
	@		10MG	N85125 001

TABLET; ORAL

ANDROID 25

<u>AB</u>	<u>ICN</u>	<u>25MG</u>	<u>N87147 001</u>
<u>AB</u>	+	<u>25MG</u>	<u>N87147 001</u>
		<u>METHYLTESTOSTERONE</u>	
<u>BP</u>	<u>DANBURY PHARMA</u>	<u>10MG</u>	<u>N80933 001</u>
<u>BP</u>		<u>25MG</u>	<u>N80931 001</u>
	@	10MG	N80933 001
	@	25MG	N80931 001
<u>BP</u>	<u>PUREPAC</u>	<u>10MG</u>	<u>N80475 002</u>
<u>BP</u>		<u>25MG</u>	<u>N80475 003</u>
	@	10MG	N80475 002
	@	25MG	N80475 003
<u>BP</u>	<u>TABLICAPS</u>	<u>10MG</u>	<u>N80313 001</u>
<u>BP</u>		<u>25MG</u>	<u>N85270 001</u>

METHYLTESTOSTERONE

TABLET; ORAL

METHYLTESTOSTERONE

	@	<u>TABLICAPS</u>	<u>10MG</u>	<u>N80313 001</u>
			<u>25MG</u>	<u>N85270 001</u>
<u>BP</u>		<u>WEST WARD</u>	<u>10MG</u>	<u>N84331 001</u>
<u>BP</u>			<u>25MG</u>	<u>N84331 002</u>
	@	WEST WARD PHARM	10MG	N84331 001
	@		25MG	N84331 002
		<u>ORETON METHYL</u>		
<u>BP</u>	*	<u>SCHERING</u>	<u>25MG</u>	<u>N03158 002</u>
	@		25MG	N03158 002

METOCLOPRAMIDE HYDROCHLORIDE

TABLET; ORAL

MAXOLON

<u>AB</u>	<u>KING PHARMS</u>	<u>EQ 10MG BASE</u>	<u>N70106 001</u>
			MAR 04, 1986
	@	EQ 10MG BASE	N70106 001
			MAR 04, 1986

METOPROLOL SUCCINATE

TABLET, EXTENDED RELEASE; ORAL

TOPROL XL

*	<u>HASSLE AB</u>	<u>EQ 50MG TARTRATE</u>	<u>N19962 001</u>
			JAN 10, 1992
*		<u>EQ 100MG TARTRATE</u>	<u>N19962 002</u>
			JAN 10, 1992
*		<u>EQ 200MG TARTRATE</u>	<u>N19962 003</u>
			JAN 10, 1992
		<u>TOPROL-XL</u>	
+	<u>ASTRA</u>	<u>EQ 50MG TARTRATE</u>	<u>N19962 001</u>
			JAN 10, 1992
+		<u>EQ 100MG TARTRATE</u>	<u>N19962 002</u>
			JAN 10, 1992
+		<u>EQ 200MG TARTRATE</u>	<u>N19962 003</u>
			JAN 10, 1992

METOPROLOL TARTRATE

TABLET; ORAL

METOPROLOL TARTRATE

<u>AB</u>	<u>APOTHECON</u>	<u>50MG</u>	<u>N74258 001</u>
			JAN 27, 1994

METOPROLOL TARTRATE

TABLET; ORAL			
<u>METOPROLOL TARTRATE</u>			
<u>AB</u>	APOTHECON	<u>100MG</u>	N74258 002 JAN 27, 1994
<u>AB</u>	COPLEY PHARM	<u>50MG</u>	N74333 001 JAN 27, 1994
<u>AB</u>		<u>100MG</u>	N74333 002 JAN 27, 1994
<u>AB</u>	GENEVA PHARMS	<u>50MG</u>	N73288 001 MAR 25, 1994
<u>AB</u>		<u>100MG</u>	N73289 001 MAR 25, 1994
<u>AB</u>	NOVOPHARM	<u>50MG</u>	N74143 001 SEP 30, 1994
<u>AB</u>		<u>100MG</u>	N74143 002 SEP 30, 1994
<u>AB</u>	PUREPAC PHARM	<u>50MG</u>	N74380 001 JUL 29, 1994
<u>AB</u>		<u>100MG</u>	N74380 002 JUL 29, 1994
<u>AB</u>	WATSON LABS	<u>50MG</u>	N74217 001 MAY 27, 1994
<u>AB</u>		<u>100MG</u>	N74217 002 MAY 27, 1994

METRIZAMIDE

INJECTABLE; INJECTION			
AMIPAQUE			
	@ NYCOMED	2.5GM/VIAL	N17982 003 SEP 12, 1983
+		3.75GM/VIAL	N17982 001
+		6.75GM/VIAL	N17982 002
@		13.5GM/VIAL	N17982 004 SEP 12, 1983
@	STERLING WINTHROP	2.5GM/VIAL	N17982 003 SEP 12, 1983
*		3.75GM/VIAL	N17982 001
*		6.75GM/VIAL	N17982 002
@		13.5GM/VIAL	N17982 004 SEP 12, 1983

MILRINONE LACTATE

INJECTABLE; INJECTION			
PRIMACOR			
+	SANOFI WINTHROP	EQ 1MG BASE/ML	N19436 001 DEC 31, 1987
*	STERLING WINTHROP	EQ 1MG BASE/ML	N19436 001 DEC 31, 1987
PRIMACOR IN DEXTROSE 5% IN PLASTIC CONTAINER			
@	SANOFI WINTHROP	EQ 10MG BASE/100ML	N20343 001 AUG 09, 1994
@		EQ 15MG BASE/100ML	N20343 002 AUG 09, 1994
+		EQ 20MG BASE/100ML	N20343 003 AUG 09, 1994
@	STERLING WINTHROP	EQ 10MG BASE/100ML	N20343 001 AUG 09, 1994
@		EQ 15MG BASE/100ML	N20343 002 AUG 09, 1994
*		EQ 20MG BASE/100ML	N20343 003 AUG 09, 1994

MINOCYCLINE HYDROCHLORIDE

TABLET; ORAL			
MINOCIN			
@	LEDERLE	EQ 50MG BASE	N50451 003 AUG 10, 1982
@		EQ 100MG BASE	N50451 002 AUG 10, 1982
MINOCYCLINE HCL			
	LEDERLE	EQ 50MG BASE	N50451 003 AUG 10, 1982
		EQ 100MG BASE	N50451 002 AUG 10, 1982

MITOMYCIN

INJECTABLE; INJECTION			
<u>MUTAMYCIN</u>			
<u>AP</u>	* BRISTOL	<u>5MG/VIAL</u>	N50450 001
<u>AP</u>	*	<u>20MG/VIAL</u>	N50450 002
	@	5MG/VIAL	N50450 001
	@	20MG/VIAL	N50450 002
<u>AP</u>	BRISTOL MYERS	<u>5MG/VIAL</u>	N62336 001
<u>AP</u>	+	<u>5MG/VIAL</u>	N62336 001
<u>AP</u>		<u>20MG/VIAL</u>	N62336 002



NAPHAZOLINE HYDROCHLORIDE

SOLUTION/DROPS; OPHTHALMIC

NAFAZOLIN

AT BAUSCH AND LOMB 0.1%

AT PHARMAFAIR 0.1%

@ 0.1%

VASOCON

AT CIBA VISION 0.1%

AT LOIAB 0.1%

N40073 001

MAY 25, 1994

N88101 001

APR 15, 1983

N88101 001

APR 15, 1983

N80235 002

MAR 24, 1983

N80235 002

MAR 24, 1983

> ADD >

> ADD >

> ADD >

> ADD >

NAPROXEN SODIUM

TABLET; ORAL

NAPROXEN SODIUM

AB COPLEY PHARM EQ 250MG BASE

AB EQ 500MG BASE

AB INVAMED EQ 250MG BASE

AB EQ 500MG BASE

AB MYLAN EQ 250MG BASE

AB EQ 500MG BASE

N74289 001

JAN 27, 1994

N74289 002

JAN 27, 1994

N74495 001

DEC 05, 1994

N74495 002

DEC 05, 1994

N74367 001

AUG 31, 1994

N74367 002

AUG 31, 1994

NAPROXEN

SUSPENSION; ORAL

NAPROSYN

AB + SYNTEX 25MG/ML

NAPROXEN

AB ROXANE 25MG/ML

TABLET; ORAL

NAPROSYN

AB SYNTEX 500MG

AB + 500MG

NAPROXEN

AB ROXANE 250MG

AB 375MG

AB 500MG

N18965 001

MAR 23, 1987

N74190 001

MAR 30, 1994

N17581 004

APR 15, 1982

N17581 004

APR 15, 1982

N74211 001

FEB 28, 1994

N74211 002

FEB 28, 1994

N74211 003

FEB 28, 1994

> ADD >

NEFAZODONE HYDROCHLORIDE

TABLET; ORAL

SERZONE

@ BRISTOL MYERS SQUIBB 50MG

100MG

150MG

200MG

+

250MG

@

300MG

N20152 001

DEC 22, 1994

N20152 002

DEC 22, 1994

N20152 003

DEC 22, 1994

N20152 004

DEC 22, 1994

N20152 005

DEC 22, 1994

N20152 006

DEC 22, 1994

NEOMYCIN SULFATE

TABLET; ORAL

NEOMYCIN SULFATE

AA LANNETT EQ 350MG BASE

@ EQ 350MG BASE

N60607 001

N60607 001

NICOTINE

FILM, EXTENDED RELEASE; TRANSDERMAL

HABITROL

BC BASEL PHARMS 7MG/24HR

N20076 001

NOV 27, 1994

TABLET, DELAYED RELEASE; ORAL

EC-NAPROSYN

+ SYNTEX 375MG

+ 500MG

N20067 002

OCT 14, 1994

N20067 003

OCT 14, 1994

NICOTINE

FILM, EXTENDED RELEASE; TRANSDERMAL  
HABITROL

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

NITROFURANTOIN

TABLET; ORAL  
FURALAN

AB	LANBETT	50MG	N80017 001
AB		100MG	N80017 002
	@	50MG	N80017 001
	@	100MG	N80017 002

NITROFURAZONE

OINTMENT; TOPICAL  
NITROFURAZONE

AT	LANBETT	0.2%	N84393 001
	@	0.2%	N84393 001

NITROGLYCERIN

INJECTABLE; INJECTION  
NITROGLYCERIN

AP	INTL MEDICATION	5MG/ML	N70026 001
			SEP 10, 1985
	@	5MG/ML	N70026 001
			SEP 10, 1985
AP	ABBOTT	0.1MG/ML	N74083 001
			OCT 26, 1994
	NITRONAL		
	* BOSKAMP	1MG/ML	N18672 001
			AUG 30, 1983

NITROGLYCERIN

INJECTABLE; INJECTION  
NITRONAL

	@ POHL BOSKAMP	1MG/ML	N18672 001
			AUG 30, 1983
AP	PARKE DAVIS	5MG/ML	N70863 001
			JAN 08, 1987
		10MG/ML	N70871 001
			JAN 08, 1987
		10MG/ML	N70872 001
			JAN 08, 1987
	@	5MG/ML	N70863 001
			JAN 08, 1987
	@	10MG/ML	N70871 001
			JAN 08, 1987
	@	10MG/ML	N70872 001
			JAN 08, 1987

NOREPINEPHRINE BITARTRATE

INJECTABLE; INJECTION  
LEVOPHED

+	SANOPI WINTHROP	EQ 1MG BASE/ML	N07513 001
*	STERLING WINTHROP	EQ 1MG BASE/ML	N07513 001

NOREPINEPHRINE BITARTRATE; PROCAINE HYDROCHLORIDE; PROPOXYCAINE HYDROCHLORIDE

INJECTABLE; INJECTION  
RAVOCAINE AND NOVOCAIN W/ LEVOPHED

	EASTMAN KODAK	EQ 0.033MG BASE/ML; 2%;	
		0.4%	N08592 003
	STERLING WINTHROP	EQ 0.033MG BASE/ML; 2%;	
		0.4%	N08592 003

NORETHINDRONE

TABLET; ORAL  
NORLUTIN

	* PARKE DAVIS	5MG	N10895 002
	@	5MG	N10895 002

NYSTATIN

SUSPENSION; ORAL

<u>AA</u>	<u>NYSTATIN</u>	<u>100,000 UNITS/ML</u>	<u>N64042 001</u>
	BAUSCH AND LOMB		FEB 28, 1994
<u>AA</u>	<u>PHARMAFAIR</u>	<u>100,000 UNITS/ML</u>	<u>N62541 001</u>
			JAN 16, 1985
	@	100,000 UNITS/ML	N62541 001
			JAN 16, 1985

NYSTATIN; TRIAMCINOLONE ACETONIDE

CREAM; TOPICAL

<u>AT</u>	<u>MYCOLOG-II</u>	<u>100,000 UNITS/GM;0.1%</u>	<u>N60576 002</u>
	+ APOTHECON		MAY 01, 1985
<u>AT</u>		<u>100,000 UNITS/GM;0.1%</u>	<u>N62606 001</u>
			MAY 15, 1985
<u>AT</u>	* SQUIBB	<u>100,000 UNITS/GM;0.1%</u>	<u>N60576 002</u>
			MAY 01, 1985
<u>AT</u>		<u>100,000 UNITS/GM;0.1%</u>	<u>N62606 001</u>
			MAY 15, 1985
<u>AT</u>	<u>NYSTATIN AND TRIAMCINOLONE ACETONIDE</u>	<u>100,000 UNITS/GM;0.1%</u>	<u>N63010 001</u>
	BARRE		DEC 20, 1988
	@	100,000 UNITS/GM;0.1%	N63010 001
			DEC 20, 1988
<u>EA</u>	CLAY PARK	<u>100,000 UNITS/GM;0.1%</u>	<u>N62186 002</u>
			JUN 06, 1985
	@	100,000 UNITS/GM;0.1%	N62186 002
			JUN 06, 1985
<u>AT</u>	<u>PHARMAFAIR</u>	<u>100,000 UNITS/GM;0.1%</u>	<u>N62657 001</u>
			JUL 30, 1986
	@	100,000 UNITS/GM;0.1%	N62657 001
			JUL 30, 1986

OINTMENT; TOPICAL

<u>AT</u>	<u>MYCOLOG-II</u>	<u>100,000 UNITS/GM;0.1%</u>	<u>N60572 001</u>
	+ APOTHECON		JUN 28, 1985
<u>AT</u>	* WESTWOOD SQUIBB	<u>100,000 UNITS/GM;0.1%</u>	<u>N60572 001</u>
			JUN 28, 1985
<u>AT</u>	<u>NYSTATIN-TRIAMCINOLONE ACETONIDE</u>	<u>100,000 UNITS/GM;0.1%</u>	<u>N62603 001</u>
	PHARMADERM		OCT 09, 1985
	@	100,000 UNITS/GM;0.1%	N62603 001
			OCT 09, 1985

OMEPRAZOLE

CAPSULE, DELAYED REL PELLETS; ORAL

	<u>PRIOSECC</u>		
	+ ASTRA MERCK	20MG	N19810 001
			SEP 14, 1989
	* MERCK	20MG	N19810 001
			SEP 14, 1989

ORPHENADRINE HYDROCHLORIDE

TABLET; ORAL

	<u>DISIPAL</u>		
	3M	50MG	N10653 001
	@	50MG	N10653 001

PAROXETINE HYDROCHLORIDE

TABLET; ORAL

	<u>PAXIL</u>		
	SMITHKLINE BEECHAM	EQ 30MG BASE	N20031 001
			DEC 29, 1992
	+	EQ 30MG BASE	N20031 001
			DEC 29, 1992

PENICILLIN G POTASSIUM

INJECTABLE; INJECTION

PENICILLIN G POTASSIUM

	@ APOTHECON	1,000,000 UNITS/VIAL	N60362 001
	@	5,000,000 UNITS/VIAL	N60362 003
	@	10,000,000 UNITS/VIAL	N60362 004
	@	20,000,000 UNITS/VIAL	N60362 002
<u>AP</u>	<u>SQUIBB</u>	<u>1,000,000 UNITS/VIAL</u>	<u>N60362 001</u>
<u>AP</u>		<u>5,000,000 UNITS/VIAL</u>	<u>N60362 003</u>
<u>AP</u>		<u>10,000,000 UNITS/VIAL</u>	<u>N60362 004</u>
<u>AP</u>		<u>20,000,000 UNITS/VIAL</u>	<u>N60362 002</u>

PENICILLIN V POTASSIUM

POWDER FOR RECONSTITUTION; ORAL

BETAPEN-VK

	@ APOTHECON	EQ 125MG BASE/5ML	N61149 001
	@	EQ 250MG BASE/5ML	N61149 002
<u>AA</u>	<u>BRISTOL</u>	<u>EQ 125MG BASE/5ML</u>	<u>N61149 001</u>

PENICILLIN V POTASSIUM

POWDER FOR RECONSTITUTION; ORAL

<u>AA</u>	<u>BETAPEN-VK</u>	<u>EQ 250MG BASE/5ML</u>	<u>N61149 002</u>
	BRISTOL		
<u>AA</u>	<u>VEETIDS</u>	<u>EQ 125MG BASE/5ML</u>	<u>N61410 001</u>
	APOTHECON		
<u>AA</u>		<u>EQ 250MG BASE/5ML</u>	<u>N61410 002</u>

TABLET; ORAL

<u>AB</u>	<u>UTICILLIN VK</u>	<u>EQ 500MG BASE</u>	<u>N61651 002</u>
	URJOHN		
	@	<u>EQ 500MG BASE</u>	<u>N61651 002</u>

PENTAMIDINE ISETHIONATE

INJECTABLE; INJECTION

<u>AP</u>	<u>PENTACARINAT</u>	<u>300MG/VIAL</u>	<u>N73447 001</u>
	RHONE POULENC RORER		<u>APR 28, 1994</u>

PENTAZOCINE LACTATE

INJECTABLE; INJECTION

	TALWIN		
+	SANOFI WINTHROP	<u>EQ 30MG BASE/ML</u>	<u>N16194 001</u>
*	STERLING WINTHROP	<u>EQ 30MG BASE/ML</u>	<u>N16194 001</u>

PENTOBARBITAL SODIUM

CAPSULE; ORAL

<u>AA</u>	<u>PENTOBARBITAL SODIUM</u>	<u>50MG</u>	<u>N85937 001</u>
	LANNETT		
<u>AA</u>		<u>100MG</u>	<u>N85915 001</u>
	@	50MG	<u>N85937 001</u>
	@	100MG	<u>N85915 001</u>

PHENDIMETRAZINE TARTRATE

TABLET; ORAL

<u>AA</u>	<u>PHENDIMETRAZINE TARTRATE</u>	<u>35MG</u>	<u>N85611 001</u>
	ZENITH		
	@ ZENITH LABS	35MG	<u>N85611 001</u>

PHENTERMINE HYDROCHLORIDE

CAPSULE; ORAL

<u>AA</u>	<u>PHENTERMINE HCL</u>	<u>30MG</u>	<u>N87022 001</u>
	LANNETT		<u>FEB 03, 1983</u>
	@	30MG	<u>N87022 001</u>
			<u>FEB 03, 1983</u>
<u>AA</u>	<u>LEMMON</u>	<u>30MG</u>	<u>N88612 001</u>
			<u>APR 04, 1984</u>
<u>AA</u>		<u>30MG</u>	<u>N88613 001</u>
			<u>APR 09, 1984</u>
<u>AA</u>		<u>30MG</u>	<u>N88614 001</u>
			<u>APR 09, 1984</u>
	@	30MG	<u>N88612 001</u>
	@	30MG	<u>APR 04, 1984</u>
	@	30MG	<u>N88613 001</u>
	@	30MG	<u>APR 09, 1984</u>
	@	30MG	<u>N88614 001</u>
	@	30MG	<u>APR 09, 1984</u>

PHENYLEPHRINE HYDROCHLORIDE; PROMETHAZINE HYDROCHLORIDE

SYRUP; ORAL

<u>AA</u>	<u>PROMETHAZINE VC PLAIN</u>	<u>5MG/5ML; 6.25MG/5ML</u>	<u>N88897 001</u>
	PENNEX		<u>JAN 04, 1985</u>
	@ PENNEX PHARMS	<u>5MG/5ML; 6.25MG/5ML</u>	<u>N88897 001</u>
	@		<u>JAN 04, 1985</u>

PHENYTOIN SODIUM, PROMPT

CAPSULE; ORAL

<u>EX</u>	<u>DIPHENYLAN SODIUM</u>	<u>100MG</u>	<u>N80857 002</u>
	LANNETT		
	@	30MG	<u>N80857 001</u>
	@	30MG	<u>N80857 001</u>
	@	100MG	<u>N80857 002</u>

PHYTONADIONE

INJECTABLE; INJECTION

<u>BF</u>	<u>PHYTONADIONE</u>	<u>1MG/0.5ML</u>	<u>N84060 001</u>
	SMITHKLINE BEECHAM		
<u>BF</u>		<u>10MG/ML</u>	<u>N84060 002</u>
	@	<u>1MG/0.5ML</u>	<u>N84060 001</u>

PHYTONADIONE

INJECTABLE; INJECTION  
PHYTONADIONE  
@ SMITHKLINE BEECHAM 10MG/ML N84060 002

PILOCARPINE HYDROCHLORIDE

TABLET; ORAL  
SALAGEN  
+ MGI 5MG N20237 001  
MAR 22, 1994

PINDOLOL

TABLET; ORAL  
PINDOLOL  
AB MUTUAL PHARM 5MG N74063 001  
JAN 27, 1994  
AB 10MG N74063 002  
JAN 27, 1994

PIPERAZINE CITRATE

SYRUP; ORAL  
BRYREL  
@ SANOFI WINTHROP EQ 500MG BASE/5ML N17796 001  
@ STERLING WINTHROP EQ 500MG BASE/5ML N17796 001  
AA VERMIDOL  
@ SOLVAY EQ 500MG BASE/5ML N80992 001  
EQ 500MG BASE/5ML N80992 001

PIROXICAM

CAPSULE; ORAL  
PIROXICAM  
AB NOVOPHARM 10MG N73637 001  
JAN 28, 1994  
AB 20MG N73638 001  
JAN 28, 1994

PLICAMYCIN

INJECTABLE; INJECTION  
MITHRACIN  
\* MILES 2.5MG/VIAL N50109 001  
+ PFIZER 2.5MG/VIAL N50109 001

POLYMYXIN B SULFATE

INJECTABLE; INJECTION  
AEROSPORIN  
AP \* BURROUGHS WELLCOME 500,000 UNITS/VIAL N62036 001  
AP + EQ 500,000 U BASE/VIAL N62036 001  
AP POLYMYXIN B SULFATE  
AP PFIZER 500,000 UNITS/VIAL N60716 001  
EQ 500,000 U BASE/VIAL N60716 001  
AP POLYMYXIN B SULFATE  
PHARMA TEK EQ 500,000 U BASE/VIAL N63000 001  
SEP 30, 1994

POTASSIUM CHLORIDE

INJECTABLE; INJECTION  
POTASSIUM CHLORIDE  
AP FUJISAWA 2MEQ/ML N87885 001  
@ 2MEQ/ML FEB 03, 1983  
N87885 001  
FEB 03, 1983  
TABLET, EXTENDED RELEASE; ORAL  
K-DUR 20  
SCHERING 20MEQ N19439 001  
JUN 13, 1986  
+ 20MEQ N19439 001  
JUN 13, 1986

PREDNISOLONE

TABLET; ORAL  
PREDNISOLONE  
BX BUNDY 5MG N83675 001  
@ 5MG N83675 001  
BX ICN 5MG N80236 001  
@ 5MG N80236 001  
BX INWOOD 5MG N80748 001  
@ INWOOD LABS 5MG N80748 001

*Bob Roehlf*

PREDNISOLONE

TABLET; ORAL  
 PREDNISOLONE  
 BX TABLICAPS 5MG N85170 001  
 @ 5MG 85170 001

PREDNISOLONE ACETATE

SUSPENSION/DROPS; OPHTHALMIC

> DLT > AT ECONOPRED PLUS 1% N17469 001  
 > ADD > BX ALCON 1% N17469 001  
 > DLT > AT \* PRED FORTE 1% N17011 001  
 > ADD > BX + ALLERGAN 1% N17011 001

PREDNISOLONE ACETATE; SULFACETAMIDE SODIUM

OINTMENT; OPHTHALMIC

817  
551  
8512

CETAPRED  
 ALCON 0.25%;10% ? N87771 001  
 AUG 06, 1993  
 + 0.25%;10% ' N87771 001  
 AUG 06, 1993  
AT + METIMYD 0.5%;10% N10210 002  
 + SCHERING  
 VASOCIDIN  
 CIBA VISION 0.5%;10% N88791 001  
 OCT 05, 1984  
 IOLAB 0.5%;10% N88791 001  
 OCT 05, 1984

SUSPENSION/DROPS; OPHTHALMIC

BLEPHAMIDE  
 ALLERGAN 0.2%;10% ? N12813 002  
 + 0.2%;10% N12813 002

PREDNISOLONE SODIUM PHOSPHATE

SOLUTION/DROPS; OPHTHALMIC

INFLAMASE FORTE  
AT + CIBA VISION EQ 0.9% PHOSPHATE N80751 002  
AT \* IOLAB EQ 0.9% PHOSPHATE N80751 002  
INFLAMASE MILD  
AT + CIBA VISION EQ 0.11% PHOSPHATE N80751 001

PREDNISOLONE SODIUM PHOSPHATE

SOLUTION/DROPS; OPHTHALMIC

INFLAMASE MILD  
AT \* IOLAB EQ 0.11% PHOSPHATE N80751 001  
PREDAIR FORTE  
AT PHARMAFAIR EQ 0.9% PHOSPHATE N88165 001  
 MAR 28, 1983  
 @ EQ 0.9% PHOSPHATE N88165 001  
 MAR 28, 1983  
PREDNISOLONE SODIUM PHOSPHATE  
AT BAUSCH AND LOMB EQ 0.11% PHOSPHATE N40065 001  
 JUL 29, 1994  
AT EQ 0.9% PHOSPHATE N40070 001  
 JUL 29, 1994

PREDNISOLONE SODIUM PHOSPHATE; SULFACETAMIDE SODIUM

SOLUTION/DROPS; OPHTHALMIC

VASOCIDIN  
AT + CIBA VISION EQ 0.23% PHOSPHATE;10% N18988 001  
 AUG 26, 1988  
AT \* IOLAB EQ 0.23% PHOSPHATE;10% N18988 001  
 AUG 26, 1988

PREDNISONE

TABLET; ORAL

PREDNISONE  
 @ 1ST TX 5MG N80371 001  
 BX BUNDY 5MG N83676 001  
 @ 5MG N83676 001  
 BX DANBURY PHARMA 50MG N86867 001  
 @ 50MG N86867 001  
 @ FERRANTE 2.5MG N80563 001  
 @ 5MG N80563 002  
 BX FIRST TX 5MG N80371 001  
 BX ICN 5MG N80237 001  
 @ 5MG N80237 001  
 BX INWOOD 1MG N80328 001  
 BX 2.5MG N80306 001  
 @ INWOOD LABS 1MG N80328 001  
 @ 2.5MG N80306 001  
 BX MK 2.5MG N80563 001  
 BX 5MG N80563 002  
 BX NYLOS 5MG N85115 001  
 @ 5MG N85115 001

PREDNISONONE

TABLET; ORAL			
PREDNISONONE			
BX	REXALL	5MG	N80232 001
	@	5MG	N80232 001
BX	SPERTI	1MG	N80359 001
BX		2.5MG	N80359 002
BX		5MG	N80359 003
	@	1MG	N80359 001
	@	2.5MG	N80359 002
	@	5MG	N80359 003
BX	WHITE TOWNE PAULSEN	20MG	N84913 002
	@ WHITEWORTH TOWNE	20MG	N84913 002

PRIMAQUINE PHOSPHATE

TABLET; ORAL			
PRIMAQUINE			
	SANOPI WINTHROP	EQ 15MG BASE	N08316 001
	STERLING WINTHROP	EQ 15MG BASE	N08316 001

PROMETHAZINE HYDROCHLORIDE

SYRUP; ORAL			
<u>PROMETHAZINE PLAIN</u>			
AA	PENNEX	6.25MG/5ML	N87953 001
	@ PENNEX PHARMS	6.25MG/5ML	NOV 15, 1982
			N87953 001
			NOV 15, 1982
TABLET; ORAL			
PROMETHAZINE HCL			
BP	TABLICAPS	12.5MG	N84080 001
BP		25MG	N84027 001
	@	12.5MG	N84080 001
	@	25MG	N84027 001
REMSED			
BP	DUPONT	25MG	N83176 002
	@ DUPONT MERCK	25MG	N83176 002

PROPANTHELINE BROMIDE

TABLET; ORAL			
<u>PRO-BANTHINE</u>			
AA	+ ROBERTS LABS	7.5MG	N08732 003

PROPANTHELINE BROMIDE

TABLET; ORAL			
<u>PRO-BANTHINE</u>			
AA	+ ROBERTS LABS	15MG	N08732 002
AA	SCS PHARMS	7.5MG	N08732 003
AA		15MG	N08732 002

PROPARACAINE HYDROCHLORIDE

SOLUTION/DROPS; OPHTHALMIC			
<u>KATAIR</u>			
AT	PHARMAFAIR	0.5%	N88087 001
	@	0.5%	JUN 07, 1983
			N88087 001
			JUN 07, 1983

PROPIOLACTONE

> DLT >	SOLUTION, IRRIGATION		
> DLT >	BETAPRONE		
> DLT >	FOREST LABS	N/A	N11657 001
> ADD >	@	N/A	N11657 001

PROPYLTHIOURACIL

TABLET; ORAL			
<u>PROPYLTHIOURACIL</u>			
BD	DANBURY PHARMA	50MG	N80932 001
	@	50MG	N80932 001
BD	LANNETT	50MG	N80016 001
	@	50MG	N80016 001
BD	TABLICAPS	50MG	N80840 001
	@	50MG	N80840 001

PROTIRELIN

INJECTABLE; INJECTION			
<u>RELEFACT TRH</u>			
AP	FERRING LABS	0.5MG/ML	N18087 001
<u>THYREL TRH</u>			
AP	FERRING LABS	0.5MG/ML	N18087 001

PSEUDOEPHEDRINE HYDROCHLORIDE

CAPSULE, EXTENDED RELEASE; ORAL

NOVAFED

DOW PHARMS 120MG N17603 001  
 + 120MG N17603 001

PSEUDOEPHEDRINE HYDROCHLORIDE; TERFENADINE

TABLET, EXTENDED RELEASE; ORAL

SELDANE-D

+ MARION MERRELL DOW 120MG;60MG N19664 001  
 AUG 19, 1991  
 MERRELL DOW 120MG;60MG N19664 001  
 AUG 19, 1991

PSEUDOEPHEDRINE HYDROCHLORIDE; TRIPROLIDINE HYDROCHLORIDE

TABLET, ORAL

ALLENFED

AA PRIVATE FORM 60MG;2.5MG N88860 001  
 JAN 31, 1985  
 @ PVT FORM 60MG;2.5MG N88860 001  
 JAN 31, 1985

TRIPHED

AA LEVMON 60MG;2.5MG N88630 001  
 MAY 17, 1984  
 @ 60MG;2.5MG N88630 001  
 MAY 17, 1984

PYRIDOSTIGMINE BROMIDE

TABLET, ORAL

PYRIDOSTIGMINE BROMIDE

KALI DUPHAR 30MG N89572 001  
 NOV 27, 1990  
 @ SOLVAY 30MG N89572 001  
 NOV 27, 1990

QUINIDINE SULFATE

TABLET, EXTENDED RELEASE; ORAL

QUINIDEX

AB + ROBINS AH 300MG N12796 002

QUINIDINE SULFATE

TABLET, EXTENDED RELEASE; ORAL

QUINIDINE SULFATE

AB COPLEY PHARM 300MG N40045 001  
 JUN 30, 1994

RANITIDINE HYDROCHLORIDE

CAPSULE; ORAL

ZANTAC 150

GLAXO EQ 150MG BASE N20095 001  
 MAR 08, 1994

ZANTAC 300

+ GLAXO EQ 300MG BASE N20095 002  
 MAR 08, 1994

GRANULE, EFFERVESCENT; ORAL

ZANTAC 150

+ GLAXO EQ 150MG BASE/PACKET N20251 002  
 MAR 31, 1994

TABLET, EFFERVESCENT; ORAL

ZANTAC 150

+ GLAXO EQ 150MG BASE N20251 001  
 MAR 31, 1994

RAUWOLFIA SERPENTINA

TABLET, ORAL

RAUWOLFIA

> DLT > BP BOWMAN 50MG N09276 005  
 > DLT > BP BOWMAN 50MG N09276 005  
 > ADD > @ BOWMAN PHARMS 50MG N09276 005

RESERPINE

ELIXIR, ORAL

SERPASIL

CIBA 0.2MG/4ML N09115 005  
 @ 0.2MG/4ML N09115 005

TABLET, ORAL

RESERPINE

BP EOW LABS 0.25MG N09838 002  
 BP + 0.25MG N09838 002

SERPASIL

BP CIBA 0.1MG N09115 001

RESERPINE

TABLET; ORAL  
 SERPASTIL  
 BP \* CIBA 0.25MG N09115 003  
 @ 0.1MG N09115 001  
 @ 0.25MG N09115 003  
 > DLT >  
 > DLT > SERPIVITE  
 > ADD > BP \* VITARINE 0.25MG N09645 002  
 @ 0.25MG N09645 002

> ADD > RIMEXOLONE  
 > ADD > SUSPENSION/DROPS; OPHTHALMIC  
 > ADD > VEXOL  
 > ADD > + ALCON 1% N20474 001  
 > ADD > DEC 30, 1994

ROCURONIUM BROMIDE

INJECTABLE; INJECTION  
 ZEMURON  
 + ORGANON 10MG/ML N20214 002  
 MAR 17, 1994  
 ZEMURON (P/F)  
 + ORGANON 10MG/ML N20214 001  
 MAR 17, 1994

ROSE BENGAL SODIUM, I-131

INJECTABLE; INJECTION  
 ROBENGATOPE  
 @ BRACCO DXS 0.5mCi/VIAL N16224 001  
 @ 1mCi/VIAL N16224 002  
 @ 2mCi/VIAL N16224 003  
 @ SQUIBB 0.5mCi/VIAL N16224 001  
 @ 1mCi/VIAL N16224 002  
 @ 2mCi/VIAL N16224 003

SALMETEROL XINAFOATE

AEROSOL, METERED; INHALATION  
 SEREVENT  
 + GLAXO EQ 0.021MG BASE/INH N20236 001  
 FEB 04, 1994

SECOBARBITAL SODIUM

CAPSULE; ORAL  
 SECOBARBITAL SODIUM  
 AA LANNETT 50MG N85909 001  
 AA 100MG N85903 001  
 @ 50MG N85909 001  
 @ 100MG N85903 001  
 RA SECONAL SODIUM 50MG N86101 001  
 ELLI  
 + 50MG N86101 001  
 OCT 03, 1983  
 N86101 001  
 OCT 03, 1983

SELENOMETHIONINE, SE-75

INJECTABLE; INJECTION  
 SETHOTOPE  
 @ BRACCO DXS 85-550 uCi/ML N17047 001  
 @ SQUIBB 85-550 uCi/ML N17047 001

SILVER SULFADIAZINE

DRESSING; TOPICAL  
 SILDIMAC  
 @ BIOPLASTY 1% N19608 001  
 NOV 30, 1989  
 @ ENQUAY 1% N19608 001  
 NOV 30, 1989

SODIUM CHLORIDE

INJECTABLE; INJECTION  
 BACTERIOSTATIC SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER  
 > DLT > AP ABBOTT 9MG/ML N18800 001  
 > DLT > OCT 29, 1982  
 > ADD > AP + 9MG/ML N18800 001  
 > ADD > OCT 29, 1982  
 SODIUM CHLORIDE  
 MCGAW 20GM/100ML N17038 001  
 @ 20GM/100ML N17038 001

SODIUM CHROMATE, CR-51

INJECTABLE; INJECTION  
CHROMIOTOPE SODIUM  
BRACCO DXS

	250 uCi/VIAL	N13993 001
	1mCi/VIAL	N13993 003
	2mCi/VIAL	N13993 002
@	250 uCi/VIAL	N13993 001
SQUIBB	1mCi/VIAL	N13993 003
@	2mCi/VIAL	N13993 002

SODIUM IODIDE, I-123

CAPSULE; ORAL  
SODIUM IODIDE I 123

AA	GOLDEN PHARMS	100 uCi	N18671 001
			MAY 27, 1982
AA		200 uCi	N18671 002
			MAY 27, 1982
AA	NORTH AN CHEM	100 uCi	N18671 001
			MAY 27, 1982
AA		200 uCi	N18671 002
			MAY 27, 1982

SODIUM IODIDE, I-131

CAPSULE; ORAL  
IODOTOPE  
BRACCO DXS

	8-100 uCi	N10929 001
	1-50mCi	N10929 003
SQUIBB	8-100 uCi	N10929 001
	1-50mCi	N10929 003

SOLUTION; ORAL  
IODOTOPE

	7-106mCi/BOT	N10929 002
BRACCO DXS	7-106mCi/BOT	N10929 002
SQUIBB		

SODIUM NITROPRUSSIDE

INJECTABLE; INJECTION  
NITROPRESS

AP	ABBOTT	50MG/VIAL	N18450 001
@		50MG/VIAL	N18450 001

SODIUM PHOSPHATE, P-32

SOLUTION; INJECTION, ORAL  
PHOSPHOTOPE

@	BRACCO DXS	1-8mCi/VIAL	N10927 001
@	SQUIBB	1-8mCi/VIAL	N10927 001

SODIUM POLYSTYRENE SULFONATE

POWDER; ORAL, RECTAL

KAYEXALATE

AA	SANOFI WINTHROP	453.6GM/BOT	N11287 001
AA	STERLING WINTHROP	453.6GM/BOT	N11287 001

SOTALOL HYDROCHLORIDE

TABLET; ORAL

	BETAPACE		
	BERLEX	120MG	N19865 005
			APR 20, 1994

> ADD >

SPIRAPRIL HYDROCHLORIDE

> ADD >

TABLET; ORAL

> ADD >

	RENORMAX		
	SANDOZ	3MG	N20240 001

> ADD >

+		24MG	N20240 004
			DEC 29, 1994

STANZOLOL

TABLET; ORAL

WINSTROL

+	SANOFI WINTHROP	2MG	N12885 001
			MAY 14, 1984

*	STERLING WINTHROP	2MG	N12885 001
			MAY 14, 1984

STAVUDINE

CAPSULE; ORAL  
ZERIT

@ BRISTOL MYERS SQUIBB	5MG	N20412 001	JUN 24, 1994
	15MG	N20412 002	JUN 24, 1994
	20MG	N20412 003	JUN 24, 1994
	30MG	N20412 004	JUN 24, 1994
+	40MG	N20412 005	JUN 24, 1994

STREPTOMYCIN SULFATE

INJECTABLE; INJECTION  
STREPTOMYCIN SULFATE

<u>AP</u> <u>PFIZER</u>	<u>EQ 1GM BASE/2ML</u>	<u>N60111 001</u>
	EQ 1GM BASE/2.5ML	N60111 001

SUCCINYLCHOLINE CHLORIDE

INJECTABLE; INJECTION  
QUELICIN

> <u>DLT</u> >	<u>AP</u> *	<u>ABBOTT</u>	<u>100MG/ML</u>	<u>N08845 004</u>
> <u>ADD</u> >		+	100MG/ML	N08845 004
> <u>DLT</u> >	<u>AP</u>	<u>SUCOSTRIN</u>	<u>100MG/ML</u>	<u>N08847 003</u>
> <u>ADD</u> >		@	100MG/ML	N08847 003

SULFACETAMIDE SODIUM

SOLUTION/DROPS; OPHTHALMIC

	<u>AT</u>	<u>SULF-10</u>	<u>10%</u>	<u>N80025 001</u>
	<u>AT</u>	<u>CIBA VISION</u>	<u>10%</u>	<u>N80025 001</u>
	<u>AT</u>	<u>IOLAB</u>	<u>10%</u>	<u>N40066 001</u>
> <u>ADD</u> >		<u>BAUSCH AND LOMB</u>	<u>10%</u>	DEC 28, 1994
> <u>ADD</u> >		<u>PHARMAFAIR</u>	<u>10%</u>	<u>N88947 001</u>
		@	10%	MAY 17, 1985
				N88947 001
				MAY 17, 1985

SULFACETAMIDE SODIUM

SOLUTION/DROPS; OPHTHALMIC

<u>AT</u>	<u>SULFAIR 10</u>	<u>10%</u>	<u>N87949 001</u>
	<u>PHARMAFAIR</u>		DEC 13, 1982
	@	10%	N87949 001
			DEC 13, 1982
<u>AT</u>	<u>SULFAIR FORTE</u>	<u>30%</u>	<u>N88385 001</u>
	<u>PHARMAFAIR</u>		OCT 13, 1983
	@	30%	N88385 001
			OCT 13, 1983
<u>AT</u>	<u>SULFAIR-15</u>	<u>15%</u>	<u>N88186 001</u>
	<u>PHARMAFAIR</u>		MAY 25, 1983
	@	15%	N88186 001
			MAY 25, 1983

SULFADIAZINE

TABLET; ORAL  
SULFADIAZINE

<u>AB</u>	<u>+ EON LABS</u>	<u>500MG</u>	<u>N40091 001</u>
			JUL 29, 1994
<u>AB</u>	<u>EON MANUFACTURE LABS</u>	<u>500MG</u>	<u>N40091 001</u>
			JUL 29, 1994
<u>AB</u>	* <u>LILLY</u>	<u>500MG</u>	<u>N04122 002</u>
	@	500MG	N04122 002

SULFADIAZINE; SULFAMERAZINE

> <u>DLT</u> >	<u>SUSPENSION, ORAL</u>		
> <u>DLT</u> >	<u>SULFONAMIDES DUPLEX</u>		
> <u>DLT</u> >	<u>LILLY</u>	<u>250MG/5ML; 250MG/5ML</u>	<u>N06317 007</u>
> <u>ADD</u> >	@	250MG/5ML; 250MG/5ML	N06317 007

SULFAMETHOXAZOLE; TRIMETHOPRIM

INJECTABLE; INJECTION

<u>AP</u>	<u>SULFAMETHOXAZOLE AND TRIMETHOPRIM</u>	<u>80MG/ML; 16MG/ML</u>	<u>N70223 001</u>
	<u>FUJISAWA</u>		DEC 29, 1987
	@	80MG/ML; 16MG/ML	N70223 001
			DEC 29, 1987

SULFAMETHOXAZOLE; TRIMETHOPRIM

SUSPENSION; ORAL  
BACTRIM  
AB \* ROCHE 200MG/5ML; 40MG/5ML N17560 001  
 @ 200MG/5ML; 40MG/5ML N17560 001  
BACTRIM PEDIATRIC  
AB ROCHE 200MG/5ML; 40MG/5ML N17560 002  
AB + 200MG/5ML; 40MG/5ML N17560 002  
TRIMETH/SULFA  
AB BARRE 200MG/5ML; 40MG/5ML N72398 001  
 @ 200MG/5ML; 40MG/5ML N72398 001  
 MAY 23, 1988  
 N72398 001  
 MAY 23, 1988

TABLET; ORAL

SULFAMETHOXAZOLE AND TRIMETHOPRIM  
AB EON LABS 400MG; 80MG N18598 003  
 @ 400MG; 80MG MAY 19, 1982  
 N18598 003  
 MAY 19, 1982  
UROPLUS DS  
AB SHIONOGI 800MG; 160MG N71816 001  
 @ 800MG; 160MG SEP 28, 1987  
 N71816 001  
 SEP 28, 1987  
UROPLUS SS  
AB SHIONOGI 400MG; 80MG N71815 001  
 @ 400MG; 80MG SEP 28, 1987  
 N71815 001  
 SEP 28, 1987

SULFASALAZINE

SUSPENSION; ORAL

AZULFIDINE  
 \* KABI 250MG/5ML N18605 001  
 250MG/5ML N86983 001  
 @ PHARMACIA 250MG/5ML N18605 001  
 + 250MG/5ML N86983 001

SULFISOXAZOLE

TABLET; ORAL

SULFISOXAZOLE  
AB LANNETT 500MG N80085 001  
 @ 500MG N80085 001

TACROLIMUS

CAPSULE; ORAL

PROGRAF  
 + FUJISAWA EQ 1MG BASE N50708 001  
 APR 08, 1994  
 + EQ 5MG BASE N50708 002  
 APR 08, 1994  
 INJECTABLE; INJECTION  
 PROGRAF  
 + FUJISAWA EQ 5MG BASE/ML N50709 001  
 APR 08, 1994

TALBUTAL

TABLET; ORAL

LOTUSATE  
 @ SANOFI WINTHROP 120MG N09410 005  
 @ STERLING WINTHROP 120MG N09410 005

TAMOXIFEN CITRATE

TABLET; ORAL

NOLVADEX  
 \* ZENECA EQ 20MG BASE N17970 002  
 MAR 21, 1994  
 @ EQ 20MG BASE N17970 002  
 MAR 21, 1994

TECHNETIUM TC-99M ALBUMIN AGGREGATED KIT

INJECTABLE; INJECTION

MACROTEC  
 BS BRACCO DXS N/A N17833 001  
 BS SQUIBB N/A N17833 001

TECHNETIUM TC-99M BICISATE KIT

INJECTABLE; INJECTION

NEUROLITE  
 DUPONT MERCK N/A N20256 001  
 NOV 23, 1994

TECHNETIUM TC-99M FERSENTETATE KIT

INJECTABLE; INJECTION  
 RENOtec  
 © BRACCO DXS  
 © SQUIBB

N/A  
 N/A

N17045 001  
 N17045 001

TECHNETIUM TC-99M MEDRONATE KIT

INJECTABLE; INJECTION  
 MDP-SQUIBB

AP BRACCO DXS  
 AP SQUIBB

N/A  
 N/A

N18107 001  
 N18107 001

TECHNETIUM TC-99M OXIDRONATE KIT

INJECTABLE; INJECTION  
 OSTEOSCAN HDP  
 MALLINCKRODT  
 TECHNISCAN HDP  
 MALLINCKRODT

N/A  
 N/A

N18321 001  
 N18321 001

TECHNETIUM TC-99M PENTETATE KIT

INJECTABLE; INJECTION  
 DTPA

AP MERCK

N/A

N18511 001  
 DEC 29, 1989

TECHNISCAN DTPA KIT

AP MERCK

N/A

N18511 001  
 DEC 29, 1989

TECHNETIUM TC-99M SESTAMIBI KIT

INJECTABLE; INJECTION  
 CARDIOLITE  
 DUPONT

N/A

N19785 001  
 DEC 21, 1990

DUPONT MERCK

N/A

N19785 001  
 DEC 21, 1990

TECHNETIUM TC-99M SULFUR COLLOID KIT

SOLUTION; INJECTION, ORAL

TESULOID  
 SQUIBB

AP

N/A

N16923 001

SOLUTION; INTRAVENOUS, ORAL

TESULOID  
 BRACCO DXS

AP

N/A

N16923 001

TERAZOSIN HYDROCHLORIDE

CAPSULE; ORAL  
 HYTRIN  
 ABBOTT

EQ 1MG BASE

N20347 001  
 DEC 14, 1994

> ADD >  
 > ADD >

+

EQ 2MG BASE

N20347 002  
 DEC 14, 1994

EQ 5MG BASE

N20347 003  
 DEC 14, 1994

EQ 10MG BASE

N20347 004  
 DEC 14, 1994

TETRACYCLINE

SYRUP, ORAL

ACHROMYCIN V  
 LEDERLE

AB

EQ 125MG HCL/5ML

N50263 002

SUNYCIN  
 SQUIBB

AB

EQ 125MG HCL/5ML

N60400 001

TETRACYCLINE  
 BARRE

AB

EQ 125MG HCL/5ML

N60633 001

MP

AB

EQ 125MG HCL/5ML

N60174 001

TETRACYCLINE HCL

AB

EQ 125MG HCL/5ML

N60291 001

PUREPAC

AB

EQ 125MG HCL/5ML

N60095 001

TETRACYCIN

AB

EQ 125MG HCL/5ML

N61468 001

TETRAMED

AB

EQ 125MG HCL/5ML

N61468 001

ZENITH

TETRACYCLINE HYDROCHLORIDE

CAPSULE; ORAL

TETRACYCLINE HCL  
 ICN

AB

500MG  
 500MG

N60471 002  
 N60471 002

TETRACYCLINE HYDROCHLORIDE

FIBER, EXTENDED RELEASE; PERIODONTAL  
ACTISITE  
+ ON SITE 12.7MG/FIBER N50653 001  
MAR 25, 1994

SYRUP; ORAL  
ACHROMYCIN V  
AB + LEDERLE 125MG/5ML N50263 002  
SUMYCIN  
AB SQUIBB 125MG/5ML N60400 001  
TETRACYCLINE HCL  
AB BARRE 125MG/5ML N60633 001  
AB MK LABS 125MG/5ML N60174 001  
AB PUREPAC PHARM 125MG/5ML N60291 001  
TETRACYN  
AB PFIPHARMECS 125MG/5ML N60095 001  
TETRAMED  
AB ZENITH LABS 125MG/5ML N61468 001

TABLET; ORAL  
PANMYCIN  
AB URJOHN 500MG N61705 002  
@ 250MG N61705 001  
@ 500MG N61705 002  
SUMYCIN  
AB \* APOTHECON 500MG N61147 004  
+ 500MG N61147 004

TETRAHYDROZOLINE HYDROCHLORIDE

SOLUTION; NASAL  
TYZINE  
+ KENWOOD LABS 0.05% N86576 002  
0.1% N86576 001  
\* KEY PHARMS 0.05% N86576 002  
0.1% N86576 001

SPRAY; NASAL  
TYZINE  
+ KENWOOD LABS 0.1% N86576 003  
\* KEY PHARMS 0.1% N86576 003

THEOPHYLLINE

CAPSULE, EXTENDED RELEASE; ORAL  
SLO-BID  
AB RHONE POULENC RORER 100MG N87892 001  
JAN 31, 1985  
AB 125MG N89540 001  
MAY 10, 1989  
AB 200MG N87893 001  
JAN 31, 1985  
AB + 300MG N87894 001  
JAN 31, 1985

THEOPHYLLINE  
AB INWOOD LABS 100MG N40052 001  
FEB 14, 1994  
AB 125MG N40052 002  
FEB 14, 1994  
AB 200MG N40052 003  
FEB 14, 1994  
AB 300MG N40052 004  
FEB 14, 1994

ELIXIR; ORAL  
LANOPHYLLIN  
AA LANNETT 80MG/15ML N84578 001  
@ 80MG/15ML N84578 001  
THEOPHYLLINE  
AA BARRE 80MG/15ML N89223 001  
MAY 27, 1988  
@ N89223 001  
MAY 27, 1988  
@ CENCI 80MG/15ML N87679 001  
APR 15, 1982  
AA LIFE LABS 80MG/15ML N87679 001  
APR 15, 1982

INJECTABLE; INJECTION  
THEOPHYLLINE 0.2% AND DEXTROSE 5% IN PLASTIC CONTAINER  
AB MCGAW 200MG/100ML N19212 001  
NOV 07, 1984  
@ 200MG/100ML N19212 001  
NOV 07, 1984  
THEOPHYLLINE 0.4% AND DEXTROSE 5% IN PLASTIC CONTAINER  
AB MCGAW 400MG/100ML N19212 002  
NOV 07, 1984  
AB 4MG/ML N19212 003  
NOV 07, 1984  
@ 400MG/100ML N19212 002  
NOV 07, 1984

THEOPHYLLINE

INJECTABLE; INJECTION

THEOPHYLLINE 0.4% AND DEXTROSE 5% IN PLASTIC CONTAINER  
 @ MCGAW 4MG/ML N19212 003  
 NOV 07, 1984

THEOPHYLLINE AND DEXTROSE 5% IN PLASTIC CONTAINER  
AP BAXTER 4MG/ML N18649 007  
 JUL 26, 1982

THEOPHYLLINE IN DEXTROSE 5% IN PLASTIC CONTAINER  
AP ABBOTT 4MG/ML N19211 007  
 DEC 14, 1984

SYRUP; ORAL

ACCURBRON  
 MERRELL DOW 150MG/15ML N88746 001  
 NOV 22, 1985  
 @ 150MG/15ML N88746 001  
 NOV 22, 1985

THEOPHYLLINE  
AA BARRE 80MG/15ML N86001 001  
 @ 80MG/15ML N86001 001

THIOTEPA

INJECTABLE; INJECTION

> ADD >  
 > ADD > AP THIOPLEX  
 > ADD > LEDERLE 15MG/VIAL N20058 001  
 DEC 22, 1994

> ADD > AP + THIOTEPA  
 + IMMUNEX 15MG/VIAL N11683 001

THYROGLOBULIN

TABLET; ORAL  
 PROLOID

BB \* FARKE DAVIS 65MG N02245 002  
 32MG N02245 005  
 100MG N02245 008  
 130MG N02245 010  
 200MG N02245 007  
 @ 32MG N02245 005  
 @ 65MG N02245 002  
 @ 100MG N02245 008  
 @ 130MG N02245 010  
 @ 200MG N02245 007  
 THYROGLOBULIN  
 + GLOBAL PHARMS 64.8MG N80151 001

THYROGLOBULIN

TABLET; ORAL  
 THYROGLOBULIN

BB RICHLYN 64.8MG N80151 001

TOBRAMYCIN

SOLUTION/DROPS; OPHTHALMIC

AT TOBRAMYCIN  
 STERIS 0.3% N63176 001  
 MAY 25, 1994

TOBRAMYCIN SULFATE

INJECTABLE; INJECTION

TOBRAMYCIN SULFATE  
AP APOTHECON EQ 10MG BASE/ML N64021 001  
 MAY 31, 1994  
AP EQ 40MG BASE/ML N64021 002  
 MAY 31, 1994  
AP EQ 40MG BASE/ML N64026 001  
 MAY 31, 1994  
AP ASTRA EQ 10MG BASE/ML N63119 001  
 OCT 31, 1994  
AP EQ 40MG BASE/ML N63120 001  
 OCT 31, 1994  
AP EQ 40MG BASE/ML N63121 001  
 OCT 31, 1994  
AP EQ 40MG BASE/ML N63122 001  
 OCT 31, 1994

TOBRAMYCIN SULFATE IN SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER

+ ABBOTT EQ 1.6MG BASE/ML N63081 006  
 JUN 02, 1993

TOLBUTAMIDE

TABLET; ORAL  
 ORINASE

> DLT >  
 > ADD > BB URJOHN 250MG N10670 002  
 @ 250MG N10670 002

TOLMETIN SODIUM

TABLET; ORAL  
TOLMETIN SODIUM  
AB MYLAN EQ 600MG BASE N74473 001  
 AUG 30, 1994

TRIAMCINOLONE

TABLET; ORAL  
TRIAMCINOLONE  
SP MYLAN 2MG N84406 001  
 @ 2MG N84406 001

TRIAMCINOLONE ACETONIDE

AEROSOL; TOPICAL  
 KENALOG  
 + APOTHECON 0.147MG/GM N12104 001  
 \* WESTWOOD SQUIBB 0.147MG/GM N12104 001

CREAM; TOPICAL  
KENALOG  
AT + APOTHECON 0.025% N11601 003  
AT + 0.1% N11601 006  
AT + 0.5% N83943 001  
AT \* WESTWOOD SQUIBB 0.025% N11601 003  
AT \* 0.1% N11601 006  
AT \* 0.5% N83943 001

TRIAMCINOLONE ACETONIDE  
AT TARO PHARMS 0.025% N40038 001  
 OCT 26, 1994

OINTMENT; TOPICAL  
KENALOG

AT + APOTHECON 0.025% N11600 003  
AT + 0.1% N11600 001  
AT + 0.5% N83944 001  
AT \* WESTWOOD SQUIBB 0.025% N11600 003  
AT \* 0.1% N11600 001  
AT \* 0.5% N83944 001

TRIAMCINOLONE ACETONIDE  
AT TARO PHARMS 0.025% N40040 001  
 SEP 30, 1994  
AT 0.1% N40037 001  
 SEP 30, 1994

TRIAZOLAM

TABLET; ORAL  
HALCION  
AB UPJOHN 0.125MG N17892 003  
 APR 26, 1985  
AB 0.25MG N17892 001  
 NOV 15, 1982

TRIAZOLAM  
AB ALPHAPHARM 0.125MG N74031 001  
 MAR 25, 1994  
AB 0.25MG N74031 002  
 MAR 25, 1994  
AB ROXANE 0.125MG N74224 001  
 JUN 01, 1994  
AB 0.25MG N74224 002  
 JUN 01, 1994

TRIHEXYPHENIDYL HYDROCHLORIDE

CAPSULE, EXTENDED RELEASE; ORAL  
 ARTANE  
 \* LEDEBEE 5MG N06773 010  
 5MG N12947 001  
 @ 5MG N06773 010  
 + 5MG N12947 001

TRILOSTANE

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

TRISULFAPYRIMIDINES (SULFADIAZINE;SULFAMERAZINE;SULFAMETHAZINE)

SUSPENSION; ORAL  
LANTREISUL  
AB LANNETT 167MG/5ML; 167MG/5ML; N80123 002  
 167MG/5ML

TRISULFAPYRIMIDINES (SULFADIAZINE;SULFAMERAZINE;SULFAMETHAZINE)

SUSPENSION; ORAL

LANTERISUL

@ LANNETT

167MG/5ML;167MG/5ML;  
167MG/5ML N80123 002

> DLT > NEOTRIZINE

> DLT > AB \* ELLYL

167MG/5ML;167MG/5ML;  
167MG/5ML N06317 012

> DLT >

> ADD > @

167MG/5ML;167MG/5ML;  
167MG/5ML N06317 012

> ADD >

TERFONYL

SQUIBB

> DLT > AB

167MG/5ML;167MG/5ML;  
167MG/5ML N06904 002

> DLT >

> ADD > +

167MG/5ML;167MG/5ML;  
167MG/5ML N06904 002

> ADD >

TABLET; ORAL

NEOTRIZINE

> DLT >

> DLT > AB \* ELLYL

167MG;167MG;167MG  
167MG;167MG;167MG N06317 011

> ADD > @

TERFONYL

SQUIBB

> DLT > AB

167MG;167MG;167MG  
167MG;167MG;167MG N06904 001

> ADD > AB +

TROPICAMIDE

SOLUTION/DROPS; OPHTHALMIC

MYDRIAFAIR

AT PHARMAFAIR

0.5% N88274 001

AT

1% N88230 001

@

0.5% N88274 001

@

1% N88230 001

TROPICAMIDE

AT BAUSCH AND LOMB

0.5% N40067 001

AT

1% N40064 001

JUL 27, 1994

JUL 27, 1994

TYROPANOATE SODIUM

CAPSULE; ORAL

BILOPAQUE

NYCOMED

750MG N13731 001

TYROPANOATE SODIUM

CAPSULE; ORAL

BILOPAQUE

STERLING WINTHROP 750MG

N13731 001

VERAPAMIL HYDROCHLORIDE

TABLET, EXTENDED RELEASE; ORAL

ISOPTIN SR

AB + KNOLL PHARM 180MG

N19152 002

DEC 15, 1989

VERAPAMIL HCL

AB BAKER NORTON 180MG

N74330 001

JAN 31, 1994

VINBLASTINE SULFATE

INJECTABLE; INJECTION

VINBLASTINE SULFATE

AP BULL 10MG/VIAL

N89565 001

AUG 18, 1987

AP FAULDING 10MG/VIAL

N89565 001

AUG 18, 1987

VINCRISTINE SULFATE

INJECTABLE; INJECTION

VINCRISTINE SULFATE

AP FUJISAWA 1MG/ML

N70411 001

SEP 10, 1986

@ 1MG/ML

N70411 001

SEP 10, 1986

VINCRISTINE SULFATE PFS

AP BULL 1MG/ML

N71484 001

APR 19, 1988

AP FAULDING 1MG/ML

N71484 001

APR 19, 1988

> ADD > VINOURELBINE TARTRATE

> ADD > INJECTABLE; INJECTION

> ADD > NAVELBINE

> ADD > + BURROUGHS WELLCOME EQ 10MG BASE/ML

N20388 001

> ADD > DEC 23, 1994

ACETAMINOPHEN

SUPPOSITORY; RECTAL  
 INFANTS' FEVERALL  
 UPSHER SMITH 80MG  
 N18337 004  
 AUG 26, 1992

TABLET, EXTENDED RELEASE; ORAL  
 TYLENOL  
 + MCNEIL CONS PRODS 650MG  
 N19872 001  
 JUN 08, 1994

ASPIRIN

TABLET, EXTENDED RELEASE; ORAL  
 8-HOUR BAYER  
 + STERLING 650MG N16030 001  
 \* STERLING WINTHROP 650MG N16030 001  
 MEASURIN  
 + STERLING 650MG N16030 002  
 \* STERLING WINTHROP 650MG N16030 002

BACITRACIN

OINTMENT; TOPICAL  
 BACITRACIN  
 COMBE 500 UNITS/GM N62799 001  
 MAY 14, 1987  
 @ 500 UNITS/GM N62799 001  
 MAY 14, 1987

CHLORPHENIRAMINE MALEATE

TABLET, EXTENDED RELEASE; ORAL  
 EFIDAC 24 CHLORPHENIRAMINE  
 ALZA 16MG  
 N19746 002  
 NOV 18, 1994

CLOTRIMAZOLE

CREAM, SUPPOSITORY; TOPICAL, VAGINAL  
 MYCELEX-7 COMBINATION PACK  
 MILES 1%, 100MG  
 N20389 002  
 JUN 23, 1994

DIPHENHYDRAMINE HYDROCHLORIDE

SYRUP; ORAL  
 DIPHENHYDRAMINE HCL  
 BARRE 12.5MG/5ML N70497 001  
 APR 25, 1989  
 @ 12.5MG/5ML N70497 001  
 APR 25, 1989

EPINEPHRINE

AEROSOL, METERED; INHALATION  
 BRONKAID MIST  
 + STERLING 0.25MG/INH N16803 001  
 \* STERLING WINTHROP 0.25MG/INH N16803 001

IBUPROFEN

TABLET; ORAL  
 IBUPROFEN  
 MCNEIL CONS PRODS 200MG N73019 001  
 MAR 30, 1994  
 PVT FORM 200MG N73691 001  
 FEB 25, 1994

INSULIN BIOSYNTHETIC HUMAN

INJECTABLE; INJECTION  
 HUMULIN BR  
 \* LILLY 100 UNITS/ML N19529 001  
 APR 28, 1986  
 @ 100 UNITS/ML N19529 001  
 APR 28, 1986

INSULIN SUSP ISOPHANE BEEF

INJECTABLE; INJECTION  
 NPH INSULIN  
 \* NOVO NORDISK 40 UNITS/ML N17929 001  
 @ 40 UNITS/ML N17929 001

INSULIN SUSP ISOPHANE BEEF/PORK

INJECTABLE; INJECTION  
 NPH ILETIN I (BEEF-PORK)  
 \* LILLY 40 UNITS/ML N17936 001  
 @ 40 UNITS/ML N17936 001

PSEUDOEPHEDRINE HYDROCHLORIDE

TABLET, EXTENDED RELEASE; ORAL  
 SUDAFED 12 HOUR  
 + WARNER WELLCOME 120MG

N73585 001  
 OCT 31, 1991

NAPHAZOLINE HYDROCHLORIDE; PHENIRAMINE MALEATE

SOLUTION/DROPS; OPHTHALMIC  
 NAPHCN-A  
 + ALCON 0.025%;0.3% N20226 001  
 JUN 08, 1994  
 OPCN-A  
 + BAUSCH AND LOMB 0.027%;0.315% N20065 001  
 JUN 08, 1994

NAPROXEN SODIUM

TABLET; ORAL  
 ALEVE  
 HAMILTON PHARMS EQ 200MG BASE N20204 002  
 JAN 11, 1994

PERMETHRIN

LOTION; TOPICAL  
 NIX  
 \* BURROUGHS WELLCOME 1% N19918 001  
 MAY 02, 1990  
 + WARNER WELLCOME 1% N19918 001  
 MAY 02, 1990

PSEUDOEPHEDRINE HYDROCHLORIDE

TABLET, EXTENDED RELEASE; ORAL  
 EFIDAC/24  
 + CIBA CONS 240MG N20021 002  
 DEC 15, 1992  
 PSEUDOEPHEDRINE HCL  
 ALZA 240MG N20021 002  
 DEC 15, 1992  
 SUDAFED 12 HOUR  
 \* BURROUGHS WELLCOME 120MG N73585 001  
 OCT 31, 1991

DRUG PRODUCTS WITH APPROVAL UNDER SECTION 505 OF THE ACT ADMINISTERED BY THE CENTER FOR BIOLOGICS EVALUATION AND RESEARCH LIST  
CUMULATIVE SUPPLEMENT NUMBER 12 / DECEMBER '94

ANTICOAGULANT CITRATE PHOSPHATE DEXTROSE ADENINE SOLUTION USP

INJECTABLE; INJECTION  
BLOOD PACK UNIT CPDA-1 IN PLASTIC CONTAINER  
BAXTER HLTHCARE

N940404  
JUL 28, 1994

INDIUM IN<sup>111</sup> CHLORIDE STERILE SOLUTION

INJECTABLE; INJECTION  
ULTRAPURE  
MALLINCKRODT      N/A

N19841  
SEP 27, 1994

**LIST OF ORPHAN PRODUCT DESIGNATIONS & APPROVALS**  
**[January-December, 1994]**

<b>NAME</b> Generic/Chemical TN= Trade Name	<b>INDICATION DESIGNATED</b>	<b>SPONSOR &amp; ADDRESS</b> DD= Date Designated MA= Marketing Approval
8-METHOXSALEN TN= UVADEX	FOR THE PREVENTION OF ACUTE REJECTION OF CARDIAC ALLOGRAFTS.	THERAKOS, INCORPORATED 201 BRANDYWINE PARKWAY WEST CHESTER PA 19380 DD 05/12/94 MA / /
AMINOSALICYLIC ACID TN= PASER GRANULES	TREATMENT OF TUBERCULOSIS INFECTIONS.	JACOBUS PHARMACEUTICAL COMPANY 37 CLEVELAND LANE PRINCETON NJ 08540 DD 02/19/92 MA 06/30/94
AMINOSIDINE TN= PAROMOMYCIN	TREATMENT OF VISCERAL LEISHMANIASIS (KALA-AZAR).	KANYOK, THOMAS P. PHARM.D. UNIV. OF ILLINOIS AT CHICAGO CHICAGO IL 60612 DD 09/09/94 MA / /
AMIODARONE HCL TN= CORDARONE	FOR THE ACUTE TREATMENT AND PROPHYLAXIS OF LIFE-THREATENING VENTRICULAR TACHYCARDIA OR VENTRICULAR FIBRILLATION.	WYETH-AYERST LABORATORIES P.O. BOX 8299 PHILADELPHIA PA 19101-1245 DD 03/16/94 MA / /
AMMONIUM TETRATHIOMOLYBDATE TN=	TREATMENT OF WILSON'S DISEASE.	BREWER, GEORGE J. M.D. UNIVERSITY OF MICHIGAN MEDICAL SCHOOL ANN ARBOR MI 48109-0618 DD 01/31/94 MA / /
ANTIVENIN, POLYVALENT CROTALID (OVINE) FAB TN= CROTAB	TREATMENT OF ENVENOMATIONS INFLICTED BY NORTH AMERICAN CROTALID SNAKES.	THERAPEUTIC ANTIBODIES INC. 1500 21ST AVENUE SOUTH, SUITE 310 NASHVILLE TN 37212 DD 01/12/94 MA / /
ARGININE BUTYRATE TN=	TREATMENT OF SICKLE CELL DISEASE AND BETA THALASSEMIA.	VERTEX PHARMACEUTICALS INC. 40 ALLSTON STREET CAMBRIDGE MA 02139-4211 DD 05/25/94 MA / /
AUTOLYMPHOCYTE THERAPY TN=	TREATMENT OF RENAL CELL CARCINOMA.	CELLCOR INCORPORATED 200 WELLS AVENUE NEWTON MA 02159 DD 07/12/94 MA / /
BACLOFEN TN= LIORESAL INTRATHECAL	TREATMENT OF SPASTICITY ASSOCIATED WITH CEREBRAL PALSY.	MEDTRONIC, INCORPORATED 800 53RD AVENUE N.E. MINNEAPOLIS MN 55432 DD 09/26/94 MA / /
BETAINE TN=	TREATMENT OF HOMOCYSTINURIA.	ORPHAN MEDICAL 13911 RIDGEDALE DRIVE MINNETONKA MN 55305 DD 05/16/94 MA / /
BOVINE IMMUNOGLOBULIN CONCENTRATE, CRYPTOSPORIDIUM PARVUM TN= SPORIDIN-G	TREATMENT AND SYMPTOMATIC RELIEF OF CRYPTOSPORIDIUM PARVUM INFECTION OF THE GASTROINTESTINAL TRACT IN IMMUNOCOMPROMISED PATIENTS.	GALAGEN, INCORPORATED 4001 LEXINGTON AVENUE NORTH ARDEN HILLS MN 55126-2998 DD 03/01/94 MA / /
BUPRENORPHINE HYDROCHLORIDE TN=	TREATMENT OF OPIATE ADDICTION IN OPIATE USERS.	RECKITT & COLMAN PHARMACEUTICALS, INC. 1901 HUGUENOT ROAD RICHMOND VA 23235 DD 06/15/94 MA / /

**CUMULATIVE LIST OF DESIGNATIONS & APPROVALS**

<b>NAME</b> <i>Generic/Chemical</i> <i>TN= Trade Name</i>	<b>INDICATION DESIGNATED</b>	<b>SPONSOR &amp; ADDRESS</b> <i>DD=Date Designated</i> <i>MA=Marketing Approval</i>
BUPRENORPHINE IN COMBINATION WITH NALOXONE TN=	TREATMENT OF OPIATE ADDICTION IN OPIATE USERS.	RECKITT & COLMAN PHARMACEUTICALS INC. 1901 HUGUENOT ROAD RICHMOND VA 23235 DD 10/27/94 MA / /
BUSULFAN TN=	FOR USE AS PREPARATIVE THERAPY FOR MALIGNANCIES TREATED WITH BONE MARROW TRANSPLANTATION.	SPARTA PHARMACEUTICALS, INC. P.O. BOX 13288 RESEARCH TRIANGLE PK NC 27709 DD 04/21/94 MA / /
BUSULFAN TN=	AS PREPARATIVE THERAPY IN THE TREATMENT OF MALIGNANCIES WITH BONE MARROW TRANSPLANTATION.	ORPHAN MEDICAL 13911 RIDGEDALE DRIVE MINNETONKA MN 55305 DD 07/28/94 MA / /
CCD 1042 TN=	TREATMENT OF INFANTILE SPASMS.	COCENSYS, INC. 213 TECHNOLOGY DRIVE IRVINE CA 92718 DD 05/25/94 MA / /
CHIMERIC (MURINE VARIABLE, HUMAN CONSTANT) MAB TO CD20 TN=	TREATMENT OF NON-HODGKIN'S B-CELL LYMPHOMA.	IDEC PHARMACEUTICALS CORPORATION 11011 TORREYANA ROAD SAN DIEGO CA 92121 DD 06/13/94 MA / /
CHOLINE CHLORIDE TN=	TREATMENT OF CHOLINE DEFICIENCY, SPECIFICALLY THE CHOLINE DEFICIENCY, HEPATIC STEATOSIS, AND CHOLESTASIS, ASSOCIATED WITH LONG-TERM PARENTERAL NUTRITION.	BUCHMAN, ALAN M.D. 6550 FANNIN, SUITE 1122 HOUSTON TX 77030 DD 02/10/94 MA / /
CLADRIBINE TN= LEUSTATIN	TREATMENT OF THE CHRONIC PROGRESSIVE FORM OF MULTIPLE SCLEROSIS.	R.W. JOHNSON RESEARCH INSTITUTE 700 ROUTE 200 SOUTH, P.O. BOX 670 RARITAN NJ 08869-0670 DD 04/19/94 MA / /
CLONAZEPAM TN= KLOPIN	TREATMENT OF HYPEREKPLEXIA (STARTLE DISEASE).	HOFFMAN-LA ROCHE, INCORPORATED 340 KINGSLAND STREET NUTLEY NJ 07110-1199 DD 08/04/94 MA / /
COAGULATION FACTOR IX (RECOMBINANT) TN=	TREATMENT OF HEMOPHILIA B.	GENETICS INSTITUTE, INCORPORATED 87 CAMBRIDGE PARK DRIVE CAMBRIDGE MA 02140 DD 10/03/94 MA / /
COUMARIN TN= ONCOSTATE	TREATMENT OF RENAL CELL CARCINOMA.	SCHAPER AND BRUMMER GmbH & CO., KG 1425 BROAD STREET CLIFTON NJ 07013 DD 12/22/94 MA / /
CY-1899 TN=	TREATMENT OF CHRONIC ACTIVE HEPATITIS B INFECTION IN HLA-A2 POSITIVE PATIENTS.	CYTEL CORPORATION 3525 JOHN HOPKINS COURT SAN DIEGO CA 92121 DD 03/16/94 MA / /
CYSTEAMINE TN= CYSTAGON	TREATMENT OF NEPHROPATHIC CYSTINOSIS.	MYLAN LABORATORIES, INC 781 CHESTNUT RIDGE ROAD, PO BOX 4310 MORGANTOWN WV 26504-4310 DD 01/25/91 MA 08/15/94

**CUMULATIVE LIST OF DESIGNATIONS & APPROVALS**

<b>NAME</b> <i>Generic/Chemical</i> <i>TN= Trade Name</i>	<b>INDICATION DESIGNATED</b>	<b>SPONSOR &amp; ADDRESS</b> <i>DD=Date Designated</i> <i>MA=Marketing Approval</i>
DAPSONE TN=	PROPHYLAXIS OF TOXOPLASMOSIS IN SEVERELY IMMUNOCOMPROMISED PATIENTS WITH CD4 COUNTS BELOW 100.	JACOBUS PHARMACEUTICAL COMPANY 37 CLEVELAND LANE, PO BOX 5290 PRINCETON NJ 08540 DD 11/07/94 MA / /
DEHYDROEPIANDROSTERONE TN=	TREATMENT OF SYSTEMIC LUPUS ERYTHEMATOSUS (SLE) AND THE REDUCTION IN THE USE OF STEROIDS IN STEROID-DEPENDENT SLE PATIENTS.	GENELABS TECHNOLOGIES, INC. 505 PENOBSCOT DRIVE REDWOOD CITY CA 94063 DD 07/13/94 MA / /
DESMOPRESSIN ACETATE TN=	TREATMENT OF MILD HEMOPHILIA A AND VON WILLEBRAND'S DISEASE.	RHONE-POULENC RORER PHARM. 500 ARCOLA ROAD COLLEGEVILLE PA 19426 DD 01/22/91 MA 03/07/94
DIMETHYL SULFOXIDE TN=	TREATMENT OF INCREASED INTRACRANIAL PRESSURE IN PATIENTS WITH SEVERE, CLOSED-HEAD INJURY, ALSO KNOWN AS TRAUMATIC BRAIN COMA, FOR WHOM NO OTHER EFFECTIVE TREATMENT IS AVAILABLE.	PHARMA 21 4850 S.W. SCHOLLS FERRY ROAD, SUITE 301 PORTLAND OR 97225-1686 DD 11/22/94 MA / /
ELLIOTT'S B SOLUTION TN=	TREATMENT OF ACUTE LYMPHOCYTIC LEUKEMIAS AND ACUTE LYMPHOBLASTIC LYMPHOMAS.	ORPHAN MEDICAL 13911 RIDGEDALE DRIVE MINNETONKA MN 55305 DD 08/24/94 MA / /
FGN-1 TN=	FOR THE SUPPRESSION AND CONTROL OF COLONIC ADENOMATOUS POLYPS IN THE INHERITED DISEASE ADENOMATOUS POLYPOSIS COLI.	CELL PATHWAYS, INC. 1700 BROADWAY, SUITE 2000 DENVER CO 80290 DD 02/14/94 MA / /
FILGRASTIM TN= NEUPOGEN	TREATMENT OF NEUTROPENIA ASSOCIATED WITH BONE MARROW TRANSPLANTS.	AMGEN, INC. 1840 DEHAVILLAND DRIVE THOUSAND OAKS CA 91320-1789 DD 10/01/90 MA 06/15/94
GAMMA-HYDROXYBUTYRATE TN=	TREATMENT OF NARCOLEPSY.	ORPHAN MEDICAL 13911 RIDGEDALE DRIVE MINNETONKA MN 55305 DD 11/07/94 MA / /
GAMMALINOLENIC ACID TN=	TREATMENT OF JUVENILE RHEUMATOID ARTHRITIS.	ZURIER, ROBERT B. M.D. 55 LAKE AVE. UNIV. OF MASS. MED. CTR. WORCESTER MA 01655 DD 07/27/94 MA / /
HEME ARGINATE TN= NORMOSANG	TREATMENT OF MYELODYSPLASTIC SYNDROMES.	LEIRAS, INCORPORATED 1850 CENTENNIAL PARK DRIVE, SUITE 450 RESTON VA 22091 DD 03/01/94 MA / /
I-131 RADIOLABELED B1 MONOCLONAL ANTIBODY TN=	TREATMENT OF NON-HODGKIN'S B-CELL LYMPHOMA.	COULTER CORPORATION 11800 S.W. 147 AVENUE, P.O. BOX 169015 MIAMI FL 33116-9015 DD 05/16/94 MA / /
IMIGLUCERASE TN= CEREZYME	FOR REPLACEMENT THERAPY IN PATIENTS WITH TYPES I, II, AND III GAUCHER'S DISEASE.	GENZYME CORPORATION ONE KENDALL SQUARE CAMBRIDGE MA 02139 DD 11/05/91 MA 05/23/94

**CUMULATIVE LIST OF DESIGNATIONS & APPROVALS**

<b>NAME</b> Generic/Chemical TN= Trade Name	<b>INDICATION DESIGNATED</b>	<b>SPONSOR &amp; ADDRESS</b> DD=Date Designated MA=Marketing Approval
IN-111 MURINE MAB(2B8-MX-DTPA) & Y-90 MURINE MAB(2B8-MXDTPA) TN= MELIMMUNE	TREATMENT OF B-CELL NON-HODGKIN'S LYMPHOMA.	IDEC PHARMACEUTICALS CORPORATION 11011 TORREYANA ROAD SAN DIEGO CA 92121 DD 09/06/94 MA / /
ISOBUTYRAMIDE TN=	TREATMENT OF SICKLE CELL DISEASE AND BETA THALASSEMIA.	VERTEX PHARMACEUTICALS INC. 40 ALLSTON STREET CAMBRIDGE MA 02139-4211 DD 05/25/94 MA / /
L-2-OXOTHIAZOLIDINE-4-CARBOXYLIC ACID TN= PROCYSTEINE	TREATMENT OF ADULT RESPIRATORY DISTRESS SYNDROME.	FREE RADICAL SCIENCES, INC. 245 FIRST STREET CAMBRIDGE MA 02142 DD 06/14/94 MA / /
L-CYSTEINE TN=	FOR THE PREVENTION AND LESSENING OF PHOTSENSITIVITY IN ERYTHROPOIETIC PROTOPORPHYRIA.	TYSON AND ASSOCIATES 12832 SOUTH CHADRON AVENUE HAWTHORNE CA 90250 DD 05/16/94 MA / /
LIPOSOME ENCAPSULATED RECOMBINANT INTERLEUKIN-2 TN=	TREATMENT OF CANCERS OF THE KIDNEY AND RENAL PELVIS.	ONCOTHERAPEUTICS, INC. 1002 EASTPARK BOULEVARD CRANBURY NJ 08512 DD 06/20/94 MA / /
MELANOMA CELL VACCINE TN=	TREATMENT OF INVASIVE MELANOMA.	MORTON, DONALD L. M.D. JOHN WAYNE CANCER INSTITUTE SANTA MONICA CA 90404 DD 10/13/94 MA / /
MITOGUAZONE TN=	TREATMENT OF DIFFUSE NON-HODGKIN'S LYMPHOMA, INCLUDING AIDS-RELATED DIFFUSE NON-HODGKIN'S LYMPHOMA.	CTRC RESEARCH FOUNDATION 11812 BECKET STREET POTOMAC MD 20854 DD 03/18/94 MA / /
MONOCLONAL AB(MURINE) ANTI-IDIOTYPE MELANOMA ASS'TED ANTIGEN TN= MELIMMUNE	TREATMENT OF INVASIVE CUTANEOUS MELANOMA.	IDEC PHARMACEUTICALS CORPORATION 11011 TORREYANA ROAD SAN DIEGO CA 92121 DD 09/19/94 MA / /
N-TRIFLUOROACETYLDRIAMYCIN-14- VALERATE TN=	TREATMENT OF CARCINOMA IN SITU OF THE URINARY BLADDER.	ANTHRA PHARMACEUTICALS, INC. 19 CARSON ROAD PRINCETON NJ 08540 DD 05/23/94 MA / /
NEUROTROPHIN-1 TN=	TREATMENT OF MOTOR NEURON DISEASE/AMYOTROPHIC LATERAL SCLEROSIS.	ERICSSON, ARTHUR DALE, M.D. 6560 FANNIN, SCURLOCK TOWER, SUITE 720 HOUSTON TX 77303 DD 09/13/94 MA / /
ORGOTEIN FOR INJECTION TN=	TREATMENT OF FAMILIAL AMYOTROPHIC LATERAL SCLEROSIS ASSOCIATED WITH A MUTATION OF THE GENE (ON CHROMOSOME 21q) FOR COPPER, ZINC SUPEROXIDE DISMUTASE.	OXIS INTERNATIONAL, INC. 6040 N. CUTTER CIRCLE, SUITE 317 PORTLAND OR 97217 DD 12/22/94 MA / /
OXANDROLONE TN= HEPANDRIN	TREATMENT OF MODERATE/SEVERE ACUTE ALCOHOLIC HEPATITIS IN THE PRESENCE OF MODERATE PROTEIN CALORIE MALNUTRITION.	BIO-TECHNOLOGY GENERAL CORP. 70 WOOD AVENUE SOUTH ISELIN NJ 08830 DD 03/18/94 MA / /

**CUMULATIVE LIST OF DESIGNATIONS & APPROVALS**

<b>NAME</b> Generic/Chemical TN= Trade Name	<b>INDICATION DESIGNATED</b>	<b>SPONSOR &amp; ADDRESS</b> DD=Date Designated MA=Marketing Approval
PEGASPARGASE TN= ONCASPAR	TREATMENT OF ACUTE LYMPHOCYTIC LEUKEMIA (ALL).	ENZON, INC. 40 KINGSBRIDGE ROAD PISCATAWAY NJ 08854-3998 DD 10/20/89 MA 02/01/94
PILOCARPINE TN= SALAGEN	TREATMENT OF XEROSTOMIA INDUCED BY RADIATION THERAPY FOR HEAD AND NECK CANCER.	MGI PHARMA, INC. SUITE 300 E, 9900 BREN ROAD EAST MINNEAPOLIS MN 55343-9667 DD 09/24/90 MA 03/22/94
PROGESTERONE TN=	ESTABLISHMENT AND MAINTENANCE OF PREGNANCY IN WOMEN UNDERGOING IN VITRO FERTILIZATION OR EMBRYO TRANSFER PROCEDURES.	GYNOPHARMA, INC. 50 DIVISION STREET SOMERVILLE NJ 08876 DD 12/22/94 MA / /
RECOMBINANT HUMAN GELSOLIN TN=	TREATMENT OF THE RESPIRATORY SYMPTOMS OF CYSTIC FIBROSIS.	BIOGEN, INC. 14 CAMBRIDGE CENTER CAMBRIDGE MA 02124 DD 01/12/94 MA / /
RECOMBINANT HUMAN LUTEINIZING HORMONE TN=	FOR USE IN ASSOCIATION WITH RECOMBINANT HUMAN FOLLICLE STIMULATING HORMONE FOR THE TREATMENT OF WOMEN WITH CHRONIC ANOVULATION DUE TO HYPOGONADOTROPIC HYPOGONADISM.	SERONO LABORATORIES, INCORPORATED 100 LONGWATER CIRCLE NORWELL MA 02061 DD 10/07/94 MA / /
RECOMBINANT METHIONYL BRAIN-DERIVED NEUROTROPHIC FACTOR TN=	TREATMENT OF AMYOTROPHIC LATERAL SCLEROSIS.	AMGEN, INCORPORATED 1840 DEHAVILLAND DRIVE THOUSAND OAKS CA 91320-1789 DD 11/28/94 MA / /
RECOMBINANT VACCINIA (HUMAN PAPILLOMAVIRUS) TN= TA-HPV	TREATMENT OF CERVICAL CANCER.	CANTAB PHARMACEUTICALS RESEARCH, LTD. 184 CAMBRIDGE SCIENCE PARK CAMBRIDGE CB4 4GN UK DD 08/24/94 MA / /
REDUCED L-GLUTATHIONE TN= CACHEXON	TREATMENT OF AIDS-ASSOCIATED CACHEXIA.	TELLURIDE PHARMACEUTICAL CORPORATION 146 FLANDERS DRIVE HILLSBOROUGH NJ 08876-4656 DD 02/14/94 MA / /
RICIN (BLOCKED) CONJUGATED MURINE MOAB (CD6) TN=	TREATMENT OF CUTANEOUS T-CELL LYMPHOMAS, ACUTE T-CELL LEUKEMIA-LYMPHOMA, AND RELATED MATURE T-CELL MALIGNANCIES.	IMMUNOGEN, INC. 148 SIDNEY STREET CAMBRIDGE MA 02139-4239 DD 09/06/94 MA / /
RIFAMPIN, ISONIAZID, PYRAZINAMIDE TN= RIFATER	SHORT-COURSE TREATMENT OF TUBERCULOSIS.	MARION MERRELL DOW, INC. P.O. BOX 9627 KANSAS CITY MO 64134-0627 DD 09/12/85 MA 05/31/94
SODIUM DICHOROACETATE TN=	TREATMENT OF LACTIC ACIDOSIS IN PATIENTS WITH SEVERE MALARIA.	STACPOOLE, PETER W., PH.D., M.D. UNIVERSITY OF FLORIDA GAINESVILLE FL 32610-0277 DD 11/10/94 MA / /
SOLUBLE RECOMBINANT HUMAN COMPLEMENT RECEPTOR TYPE 1 TN=	PREVENTION OR REDUCTION OF ADULT RESPIRATORY DISTRESS SYNDROME.	T CELL SCIENCES, INC. 38 SIDNEY STREET CAMBRIDGE MA 02139-4135 DD 11/21/94 MA / /

**CUMULATIVE LIST OF DESIGNATIONS & APPROVALS**

<b>NAME</b> <i>Generic/Chemical</i> <i>TN- Trade Name</i>	<b>INDICATION DESIGNATED</b>	<b>SPONSOR &amp; ADDRESS</b> <i>DD-Date Designated</i> <i>MA-Marketing Approval</i>
SOMATOSTATIN TN=	TREATMENT OF BLEEDING ESOPHAGEAL VARICES.	UCB PHARMACEUTICALS, INC. PO BOX 4410 HAMPTON VA 23664-0410 DD 12/22/94 MA / /
SOMATROPIN TN= NUTROPIN (EXCLUSIVITY NOT APPLICABLE)	FOR USE IN THE LONG-TERM TREATMENT OF CHILDREN WHO HAVE GROWTH FAILURE DUE TO A LACK OF ADEQUATE ENDOGENOUS GROWTH HORMONE SECRETION.	GENENTECH, INC. 460 POINT SAN BRUNO BOULEVARD SOUTH SAN FRANCISCO CA 94080 DD 03/06/87 MA 03/09/94
SOMATROPIN TN= GENOTROPIN/GENOTONORM	TREATMENT OF ADULTS WITH GROWTH HORMONE DEFICIENCY.	PHARMACIA, INC. P.O. BOX 16529 COLUMBUS OH 43216-6529 DD 09/06/94 MA / /
SULFADIAZINE TN=	FOR USE IN COMBINATION WITH PYRIMETHAMINE FOR THE TREATMENT OF TOXOPLASMA GONDII ENCEPHALITIS IN PATIENTS WITH AND WITHOUT ACQUIRED IMMUNODEFICIENCY SYNDROME.	EON LABS MANUFACTURING, INC. 227-15 NORTH CONDUIT AVENUE LAURELTON NY 11413 DD 03/14/94 MA 07/29/94
TIZANIDINE HCL TN= ZANAFLEX	TREATMENT OF SPASTICITY ASSOCIATED WITH MULTIPLE SCLEROSIS AND SPINAL CORD INJURY.	ATHENA NEUROSCIENCES, INC. 800F GATEWAY BOULEVARD SOUTH SAN FRANCISCO CA 94080 DD 01/31/94 MA / /
TOBRAMYCIN FOR INHALATION TN=	TREATMENT OF BRONCHOPULMONARY INFECTIONS OF PSEUDOMONAS AERUGINOSA IN CYSTIC FIBROSIS PATIENTS.	PATHOGENESIS CORPORATION 201 ELLIOTT AVENUE WEST, SUITE 150 SEATTLE WA 98119 DD 10/13/94 MA / /
TREOSULFAN TN= OVASTAT	TREATMENT OF OVARIAN CANCER.	MEDAC GmbH c/o PRINCETON REG. ASSOC. 65 SOUTH MAIN STREET PENNINGTON NJ 08534 DD 05/16/94 MA / /

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

---

NO DECEMBER 1994 ADDITIONS

## BIOPHARMACEUTIC GUIDANCE AVAILABILITY

DRUG NAME (DOSAGE FORM)	DATE	REVISED DATE
-------------------------	------	--------------

THE FOLLOWING IS A LIST OF GUIDANCES AVAILABLE FOR *IN VIVO* BIOEQUIVALENCE STUDIES AND *IN VITRO* DISSOLUTION TESTING. COMMENTS AND SUGGESTIONS CONCERNING THESE GUIDANCES ARE ENCOURAGED AND SHOULD BE SENT TO THE DIVISION OF BIOEQUIVALENCE (HFD-650, MPN-2 ROOM 279) 5600 FISHERS LANE, ROCKVILLE, MD 20857.

REFER BACK TO THE APPROVED DRUG PRODUCTS WITH THERAPEUTIC EQUIVALENCE EVALUATIONS, 14TH EDITION FOR A FULL LISTING OF BIOPHARMACEUTIC GUIDANCE AVAILABILITY DATA. ONLY NEW DATA WILL BE ADDED TO THE CUMULATIVE SUPPLEMENT.

ALBUTEROL (METERED DOSE INHALER - <i>IN VIVO</i> )	JAN 27, 1994	OCT 06, 1994
DICLOFENAC SODIUM (TABLET)	DEC 24, 1992	FEB 04, 1994
FLURBIPROFEN (TABLET)	DEC 24, 1992	
PHENYTOIN (SUSPENSION AND CHEWABLE TABLET)	MAR 04, 1994	
PHENYTOIN SODIUM (CAPSULE, EXTENDED AND PROMPT)	MAR 04, 1994	
TOLMETIN SODIUM (CAPSULE AND TABLET)	APR 20, 1989	OCT 06, 1994

## ANDA SUITABILITY PETITIONS

## PETITIONS APPROVED

DRUG NAME DOSAGE FORM; ROUTE	STRENGTH (CONTAINER SIZE)	DOCKET NUMBER	PETITIONER	REASON FOR PETITION	STATUS
---------------------------------	------------------------------	---------------	------------	------------------------	--------

THE FOLLOWING ARE TWO LISTS OF PETITIONS FILED UNDER SECTION 505(j)(2)(C) OF THE ACT WHERE THE AGENCY HAS DETERMINED THAT THE REFERENCED PRODUCT: (1) IS SUITABLE FOR SUBMISSION AS AN ANDA (PETITIONS APPROVED) OR (2) IS NOT SUITABLE FOR SUBMISSION AS AN ANDA (PETITIONS DENIED). THE DETERMINATION THAT AN ANDA WILL BE APPROVED IS NOT MADE UNTIL THE ANDA ITSELF IS SUBMITTED AND REVIEWED BY THE AGENCY. A COPY OF EACH PETITION IS LISTED BY DOCKET NUMBER ON PUBLIC DISPLAY IN FDA'S DOCKETS MANAGEMENT BRANCH, HFA-305, ROOM 4-62, 5600 FISHERS LANE, ROCKVILLE, MD 20857.

REFER BACK TO THE APPROVED DRUG PRODUCTS WITH THERAPEUTIC EQUIVALENCE EVALUATIONS, 13TH EDITION FOR A FULL LISTING OF ANDA SUITABILITY PETITIONS DATA. ONLY NEW DATA WILL BE ADDED TO THE CUMULATIVE SUPPLEMENT.

ACETAMINOPHEN; BUTALBITAL; CAFFEINE; HYDROCODONE BITARTRATE CAPSULE; ORAL	325MG 50MG 40MG 5MG	93 P-0346/ CP1	MIKART	NEW COMBINATION	APPROVED NOV 15, 1994
ACETAMINOPHEN; BUTALBITAL; CAFFEINE; HYDROCODONE BITARTRATE TABLET; ORAL	325MG 50MG 40MG 5MG	93 P-0346/ CP1	MIKART	NEW COMBINATION NEW DOSAGE FORM	APPROVED NOV 15, 1994
ACETAMINOPHEN; BUTALBITAL; CAFFEINE; HYDROCODONE BITARTRATE TABLET; ORAL	500MG 50MG 40MG 5MG	93 P-0298/ CP1	ARNOLD & PORTER	NEW COMBINATION NEW DOSAGE FORM NEW STRENGTH	APPROVED NOV 15, 1994
ACYCLOVIR TABLET; ORAL	200MG	93 P-0339/ CP1	NOVOPHARM	NEW DOSAGE FORM	APPROVED FEB 08, 1994
ACYCLOVIR SODIUM INJECTABLE; INJECTION	25MG/ML (20ML/VIAL) (40ML/VIAL)	93 P-0469/ CP1	FAULDING	NEW DOSAGE FORM	APPROVED JUN 09, 1994
ESTRADIOL TABLET; ORAL	1.5MG	93 P-0344/ CP1	BRISTOL MYERS SQIBB	NEW STRENGTH	APPROVED JUN 08, 1994
LOPERAMIDE HYDROCHLORIDE TABLET, EFFERVESCENT; ORAL	1MG	93 P-0332/ CP1	ELLIS PHARM CONSULTING	NEW DOSAGE FORM	APPROVED FEB 08, 1994
METHYLTESTOSTERONE CAPSULE; ORAL	25MG	93 P-0459/ CP1	ICN	NEW DOSAGE FORM	APPROVED JUN 08, 1994
MORPHINE SULFATE CAPSULE, EXTENDED RELEASE; ORAL	15MG 60MG 100MG	93 P-0446/ CP1	PHARMA CONSULT	NEW DOSAGE FORM	APPROVED JUN 08, 1994
MORPHINE SULFATE CAPSULE, EXTENDED RELEASE; ORAL	90MG	93 P-0446/ CP1	PHARMA CONSULT	NEW DOSAGE FORM NEW STRENGTH	APPROVED JUN 08, 1994
PREDNISON TABLET, CHEWABLE; ORAL	1MG 2.5MG 20MG 50MG	93 P-0333/ CP1	DURA	NEW DOSAGE FORM	APPROVED JUN 08, 1994
PSEUDOEPHEDRINE HYDROCHLORIDE; TERFENADINE CAPSULE, EXTENDED RELEASE; ORAL	120MG 60MG	93 P-0367/ CP1	EURAND AMERICA	NEW DOSAGE FORM	APPROVED FEB 08, 1994

## ANDA SUITABILITY PETITIONS

## PETITIONS APPROVED

DRUG NAME DOSAGE FORM; ROUTE	STRENGTH (CONTAINER SIZE)	DOCKET NUMBER	PETITIONER	REASON FOR PETITION	STATUS
VALPROIC ACID CAPSULE; ORAL	500MG	93 P-0452/ CP1	KROSS	NEW STRENGTH	APPROVED DEC 19, 1994

## \*\*\*ERRATA\*\*\*

CIMETIDINE TABLET, EFFERVESCENT; ORAL	200MG 300MG 400MG 800MG	93 P-0048/ CP1*	FLEMINGTON PHARM	NEW DOSAGE FORM	APPROVED SEP 18, 1993
--	----------------------------------	--------------------	------------------	--------------------	--------------------------

\*Not previously published

## EXCLUSIVITY TERMS

DUE TO SPACE LIMITATIONS IN THE EXCLUSIVITY COLUMN, ABBREVIATIONS AND REFERENCES HAVE BEEN DEVELOPED. PLEASE REFER BACK TO THE APPROVED DRUG PRODUCTS WITH THERAPEUTIC EQUIVALENCE EVALUATIONS, 14TH EDITION FOR A FULL LISTING OF EXCLUSIVITY TERMS (ABBREVIATIONS, NEW DOSING SCHEDULE, NEW INDICATIONS AND PATENT USE CODES). ONLY NEW CODES WILL BE ADDED TO THE CUMULATIVE SUPPLEMENT.

### REFERENCES NEW DOSING SCHEDULE

D-21 ALTERNATIVE DOSAGE OF 300MG ONCE DAILY AFTER THE EVENING MEAL  
 D-22 REDUCTION IN INFUSION TIME FROM 24 TO 4 HOURS FOR THE 60MG DOSE  
 D-23 INCREASE MAXIMUM DOSE AND VARIATIONS IN THE DOSING REGIMEN  
 D-24 FOR OVARIAN CANCER THE RECOMMENDED REGIMEN IS 135MG/M<sup>2</sup> OR 175MG/M<sup>2</sup> INTRAVENOUSLY OVER THREE HOURS EVERY THREE WEEKS  
 D-25 ADDITIONAL DOSAGE REGIMEN EQUAL TO HALF OF THE ORIGINAL DOSING REGIMEN

### REFERENCES NEW INDICATION

I-99 PEDIATRIC ANESTHESIA IN CHILDREN 3 YEARS AND OLDER  
 I-100 TO DECREASE THE INCIDENCE OF CANDIDIASIS IN PATIENTS UNDERGOING BONE MARROW TRANSPLANTATION WHO RECEIVE CYTOTOXIC CHEMOTHERAPY AND/OR RADIATION THERAPY  
 I-101 TREATMENT OF DIABETIC NEPHROPATHY IN PATIENTS WITH TYPE I INSULIN-DEPENDENT DIABETES MELLITUS AND RETINOPATHY  
 I-102 TREATMENT OF OBSESSIVE-COMPULSIVE DISORDER  
 I-103 PROPHYLAXIS AGAINST PNEUMOCYSTIS CARINII PNEUMONIA IN INDIVIDUALS WHO ARE IMMUNOCOMPROMISED AND CONSIDERED TO BE AT RISK OF DEVELOPING PNEUMOCYSTIS CARINII PNEUMONIA  
 I-104 TREATMENT OF PULMONARY AND EXTRAPULMONARY ASPERGILLOSIS IN PATIENTS WHO ARE INTOLERANT OF OR WHO ARE REFRACTORY TO AMPHOTERICIN B THERAPY  
 I-105 TREATMENT OF METASTATIC CARCINOMA OF THE BREAST AFTER FAILURE OF FIRST-LINE OR SUBSEQUENT CHEMOTHERAPY  
 I-106 TREATMENT OF ACROMEGALY  
 I-107 VAGINAL CANDIDIASIS  
 I-108 EXPANDED USE-FOR ICU PATIENTS UNDERGOING LONG-TERM INFUSION DURING MECHANICAL VENTILATION  
 I-109 TYPHOID FEVER  
 I-110 PREVENTION OF NAUSEA AND VOMITING ASSOCIATED WITH RADIOTHERAPY  
 I-111 TREATMENT OF PAGET'S DISEASE OF BONE  
 I-112 MANAGEMENT OF MODERATE TO SEVERE PAIN  
 I-113 TREATMENT OF PROSTATITIS  
 I-114 USE IN CHILDREN TO VISUALIZE LESIONS WITH ABNORMAL VASCULARITY IN THE BRAIN (INTRACRANIAL LESIONS), SPINE AND ASSOCIATED TISSUE  
 I-115 USE IN MRI IN ADULTS TO VISUALIZE LESIONS IN THE HEAD AND NECK  
 I-116 MAINTENANCE OF HEALING OF EROSIIVE ESOPHAGITIS

### REFERENCES PATENT USE CODE

U-91 ALTERNATIVE THERAPY TO TRIMETHOPRIM-SULFAMETHOXAZOLE FOR TREATMENT OF MODERATE-TO-SEVERE PNEUMOCYSTIS CARINII PNEUMONIA IN IMMUNOCOMPROMISED AND AIDS PATIENTS  
 U-92 TREATMENT OF DIABETIC NEPHROPATHY IN PATIENTS WITH TYPE I INSULIN DEPENDENT DIABETES MELLITUS AND RETINOPATHY  
 U-93 USE AS AN ANTIHISTAMINE/DECONGESTANT  
 U-94 TREATMENT OF ADULTS WITH ADVANCED HIV INFECTION WHO ARE INTOLERANT OF APPROVED THERAPIES WITH PROVEN CLINICAL BENEFIT OR WHO HAVE EXPERIENCED SIGNIFICANT CLINICAL OR IMMUNOLOGIC DETERIORATION WHILE RECEIVING THESE THERAPIES OR FOR WHOM SUCH THERAPIES ARE CONTRAINDICATED  
 U-95 SHORT TERM MANAGEMENT OF MODERATE PRURITIS IN ADULTS WITH ATOPIC DERMATITIS AND LICHEN SIMPLEX CHRONICUS

**EXCLUSIVITY TERMS****REFERENCES**  
*PATENT USE CODE*

U-96 METHOD OF TREATING VARICELLA ZOSTER (SHINGLES) INFECTIONS  
U-97 A METHOD OF TREATING A PATIENT IN NEED OF MEMORY ENHANCEMENT  
U-98 A METHOD OF INDUCING REGRESSION OF LEUKEMIA CELL GROWTH IN A MAMMAL  
U-99 METHOD OF PROVIDING POTASSIUM TO A SUBJECT IN NEED OF POTASSIUM  
U-100 METHOD OF TREATING OCULAR INFLAMMATION  
U-101 ADJUNCT TO CONVENTIONAL CT OR MRI IMAGING IN THE LOCALIZATION OF STROKE IN PATIENTS IN WHOM STROKE HAS ALREADY BEEN  
DIAGNOSED

PRESCRIPTION AND OTC DRUG PRODUCT  
PATENT AND EXCLUSIVITY DATA

APPL/PROD NUMBER	INGREDIENT NAME; TRADE NAME	PATENT NUMBER	PATENT EXPIRES	USE CODE	EXCLUS CODE	EXCLUS EXPIRES
19872 001	ACETAMINOPHEN; TYLENOL				NDF	JUN 08, 1997
19806 001	ACRIVASTINE; SEMPREX-D	4650807	MAR 17, 2004	U-93		
		4501893	FEB 26, 2002		NC	MAR 25, 1997
74346 001	AMINOSALICYLIC ACID; PASER				ODE	JUN 30, 2001
18700 001	AMRINONE LACTATE; INOCOR	4072746	JUL 31, 1998	U-7	NCE	JUL 31, 1994
20304 001	APROTININ BOVINE; TRASYLOL				ODE	DEC 29, 2000
					D-25	OCT 12, 1997
18831 001	ATRACURIUM BESYLATE; TRACRIUM	4179507	DEC 18, 1996		I-108	JUN 06, 1997
20233 001	BUDESONIDE; RHINOCORT				NCE	FEB 14, 1999
18731 001	BUSPIRONE HYDROCHLORIDE; BUSPAR	5015646	MAY 14, 2008	U-13		
18731 002	BUSPIRONE HYDROCHLORIDE; BUSPAR	5015646	MAY 14, 2008	U-13		
18343 001	CAPTAPRIL; CAPOTEN	5238924	AUG 24, 2010	U-92	I-101	JAN 28, 1997
18343 002	CAPTAPRIL; CAPOTEN	5238924	AUG 24, 2010	U-92	I-101	JAN 28, 1997
18343 003	CAPTAPRIL; CAPOTEN	5238924	AUG 24, 2010	U-92	I-101	JAN 28, 1997
18343 005	CAPTAPRIL; CAPOTEN	5238924	AUG 24, 2010	U-92	I-101	JAN 28, 1997
18343 006	CAPTAPRIL; CAPOTEN	5238924	AUG 24, 2010	U-92	I-95	SEP 23, 1996
		4105776	AUG 08, 1995		I-101	JAN 28, 1997
>ADD>	19746 002	CHLORPHENIRAMINE MALEATE; EFIDAC 24	4857330	AUG 15, 2006		
>ADD>		CHLORPHENIRAMINE	4673405	MAR 18, 2003		
>ADD>			4576604	MAR 18, 2003		
>ADD>	19574 001	CHLORTHALIDONE; THALITONE	4933360	JUN 12, 2007		
	19537 002	CIPROFLOXACIN HYDROCHLORIDE; CIPRO	4670444	OCT 01, 2002	U-36	I-66 JUL 21, 1997
					I-109	JUL 21, 1997
	19537 003	CIPROFLOXACIN HYDROCHLORIDE; CIPRO	4670444	OCT 01, 2002	U-36	I-66 JUL 21, 1997
					I-109	JUL 21, 1997
	19537 004	CIPROFLOXACIN HYDROCHLORIDE; CIPRO	4670444	OCT 01, 2002	U-36	I-66 JUL 21, 1997
					I-109	JUL 21, 1997
	20392 001	CYSTEAMINE BITARTRATE; CYSTAGON			NCE	AUG 15, 1999
					ODE	AUG 15, 2001
	20392 002	CYSTEAMINE BITARTRATE; CYSTAGON			NCE	AUG 15, 1999
					ODE	AUG 15, 2001
>ADD>	20287 001	DALTEPARIN SODIUM; FRAGMIN			NCE	DEC 22, 1999
	20355 001	DESMOPRESSIN ACETATE; DESMOPRESSIN ACETATE			NP	MAR 07, 1997
					ODE	MAR 07, 2001

PRESCRIPTION AND OTC DRUG PRODUCT  
PATENT AND EXCLUSIVITY DATA

APPL/PROD NUMBER	INGREDIENT NAME; TRADE NAME	PATENT NUMBER	PATENT EXPIRES	USE CODE	EXCLUS CODE	EXCLUS EXPIRES
20062 001	DILTIAZEM HYDROCHLORIDE; CARDIZEM CD	5364620	NOV 14, 2011	U-3		
		5286497	FEB 14, 2011			
20062 002	DILTIAZEM HYDROCHLORIDE; CARDIZEM CD	5364620	NOV 14, 2011	U-3		
		5286497	FEB 14, 2011			
20062 003	DILTIAZEM HYDROCHLORIDE; CARDIZEM CD	5364620	NOV 14, 2011	U-3		
		5286497	FEB 14, 2011			
20062 004	DILTIAZEM HYDROCHLORIDE; CARDIZEM CD	5364620	NOV 14, 2011	U-3		
		5286497	FEB 14, 2011			
>ADD>	20408 001	DORZOLAMIDE HYDROCHLORIDE; TRUSOPT			NCE	DEC 09, 1999
	20126 001	DOXEPIN HYDROCHLORIDE; ZONALON	4395420	JUL 26, 2000	U-95	NDF APR 01, 1997
	20323 001	ESTRADIOL; VIVELLE			NS	OCT 28, 1997
	20323 003	ESTRADIOL; VIVELLE			NS	OCT 28, 1997
	81295 001	ESTRADIOL; ESTRACE			I-76	SEP 08, 1995
>ADD>	20303 001	ESTROGENS, CONJUGATED; PREMPRO (PREMARIN/CYCRIN)			NP	DEC 30, 1997
>ADD>	20303 002	ESTROGENS, CONJUGATED; PREMPHASE (PREMARIN/CYCRIN)			NP	DEC 30, 1997
	18922 004	ETODOLAC; LODINE	4076831	FEB 28, 1997	U-45	NCE JAN 31, 1996
	20363 002	FAMCICLOVIR; FAMVIR	5246937	SEP 21, 2010	U-96	NCE JUN 29, 1999
	20249 001	FAMOTIDINE; PEPCID	4283408	AUG 11, 2000		I-69 DEC 10, 1994
	19834 004	FELODIPINE; PLENDIL	4264611	APR 28, 1998		NCE JUL 25, 1996
	19304 001	FENOFIBRATE; LIPIDIL	4058552	NOV 15, 1994		NCE DEC 31, 1998
	19960 001	FLOSEQUINAN; MANOPLAX	4552884	DEC 31, 2006	U-71	
	19960 002	FLOSEQUINAN; MANOPLAX	4552884	DEC 31, 2006	U-71	
	19960 003	FLOSEQUINAN; MANOPLAX	4552884	DEC 31, 2006	U-71	
	19960 004	FLOSEQUINAN; MANOPLAX	4552884	DEC 31, 2006	U-71	
	19949 001	FLUCONAZOLE; DIFLUCAN	4416682	NOV 22, 2000		I-100 DEC 30, 1996
	19949 002	FLUCONAZOLE; DIFLUCAN	4416682	NOV 22, 2000		I-100 DEC 30, 1996
	19949 003	FLUCONAZOLE; DIFLUCAN	4416682	NOV 22, 2000		I-100 DEC 30, 1996
	19950 001	FLUCONAZOLE; DIFLUCAN	4416682	NOV 22, 2000		I-100 DEC 30, 1996
	20322 001	FLUCONAZOLE; DIFLUCAN	4416682	NOV 22, 2000		NCE JAN 29, 1995
			4404216	OCT 16, 2003		I-100 DEC 30, 1996
						NS JUN 30, 1997
						I-107 JUN 30, 1997
	18936 001	FLUOXETINE HYDROCHLORIDE; PROZAC	4018895	APR 19, 1994	U-12	I-102 FEB 28, 1997
	18936 006	FLUOXETINE HYDROCHLORIDE; PROZAC	4314081	FEB 02, 2001		I-102 FEB 28, 1997
	20101 001	FLUOXETINE HYDROCHLORIDE; PROZAC	4314081	FEB 02, 2001		I-102 FEB 28, 1997
>ADD>	20243 001	FLUVOXAMINE MALEATE; LUVOX	4085225	APR 18, 1995		NCE DEC 05, 1999
>ADD>	20243 002	FLUVOXAMINE MALEATE; LUVOX	4085225	APR 18, 1995		NCE DEC 05, 1999

PRESCRIPTION AND OTC DRUG PRODUCT  
PATENT AND EXCLUSIVITY DATA

APPL/PROD NUMBER	INGREDIENT NAME; TRADE NAME	PATENT NUMBER	PATENT EXPIRES	USE CODE	EXCLUS CODE	EXCLUS EXPIRES
>ADD>	20243 003 FLUVOXAMINE MALEATE; LUVOX	4085225	APR 18, 1995		NCE	DEC 05, 1999
>ADD>	20243 004 FLUVOXAMINE MALEATE; LUVOX	4085225	APR 18, 1995		NCE	DEC 05, 1999
>ADD>	19915 002 FOSINOPRIL SODIUM; MONOPRIL	5006344	APR 09, 2008			
>ADD>	19915 003 FOSINOPRIL SODIUM; MONOPRIL	5006344	APR 09, 2008			
	20286 001 FOSINOPRIL SODIUM; MONOPRIL-HCT	5006344	APR 09, 2008			
		4384123	MAY 17, 2000			
		4337201	JUN 29, 2001		NC	NOV 30, 1997
	20286 002 FOSINOPRIL SODIUM; MONOPRIL-HCT	5006344	APR 09, 2008			
		4384123	MAY 17, 2000			
		4337201	JUN 29, 2001		NC	NOV 30, 1997
	20235 001 GABAPENTIN; NEURONTIN	4894476	JAN 16, 2007			
	20235 002 GABAPENTIN; NEURONTIN	4894476	JAN 16, 2007			
	20235 003 GABAPENTIN; NEURONTIN	4894476	JAN 16, 2007			
	20123 001 GADODIAMIDE; OMNISCAN	4687659	JAN 08, 2007	U-76	NCE	JAN 08, 1998
	20131 001 GADOTERIDOL; PROHANCE				I-115	NOV 21, 1997
					I-114	NOV 21, 1997
>ADD>	20460 001 GANCICLOVIR; CYTOVENE	4642346	FEB 10, 2004			
>ADD>		4507305	OCT 19, 1999	U-64		
>ADD>		4423050	OCT 19, 1999	U-64		
>ADD>		4355032	MAR 16, 2003			
	20329 001 GLIPIZIDE; GLUCOTROL XL				NDF	APR 26, 1997
	20329 002 GLIPIZIDE; GLUCOTROL XL				NDF	APR 26, 1997
	19778 003 HYDROCHLOROTHIAZIDE; PRINZIDE 10-12.5	4472380	SEP 18, 2001			
		4374829	DEC 30, 2001	U-3	NS	NOV 18, 1996
	19888 001 HYDROCHLOROTHIAZIDE; ZESTORETIC 20-12.5	4472380	SEP 18, 2001			
		4374829	DEC 30, 2001	U-3		
	19888 002 HYDROCHLOROTHIAZIDE; ZESTORETIC 20-25	4374829	DEC 30, 2001	U-3		
	19888 003 HYDROCHLOROTHIAZIDE; ZESTORETIC 10-12.5	4472380	SEP 18, 2001		NS	NOV 18, 1996
		4374829	DEC 30, 2001	U-3		
	19771 001 IBUPROFEN; ADVIL COLD AND SINUS	4552899	NOV 12, 2002			
	20135 001 IBUPROFEN; MOTRIN				NDF	NOV 16, 1997
	20135 002 IBUPROFEN; MOTRIN				NDF	NOV 16, 1997
	20367 001 IMIGLUCERASE; CEREZYME				NCE	MAY 23, 1999
					ODE	MAY 23, 2001
	20314 001 INDIUM IN-111 PENTETREOTIDE KIT; OCTREOSCAN				NCE	JUN 02, 1999
	20084 001 IOBENGUANE SULFATE I 131; IOBENGUANE SULFATE I 131	4584187	APR 22, 2003		NCE	MAR 25, 1999

PRESCRIPTION AND OTC DRUG PRODUCT  
PATENT AND EXCLUSIVITY DATA

APPL/PROD NUMBER	INGREDIENT NAME; TRADE NAME	PATENT NUMBER	PATENT EXPIRES	USE CODE	EXCLUS CODE	EXCLUS EXPIRES
20327 001	IOPAMIDOL; ISOVUE-200	4001323	JAN 04, 1996			
20327 002	IOPAMIDOL; ISOVUE-250	4001323	JAN 04, 1996			
20327 003	IOPAMIDOL; ISOVUE-300	4001323	JAN 04, 1996			
20327 004	IOPAMIDOL; ISOVUE-370	4001323	JAN 04, 1996			
50705 001	ISONIAZID; RIFATER				ODE	MAY 31, 2001
20336 001	ISRADIPINE; DYNACIRC CR	5030456	JUL 09, 2008			
		4950486	AUG 21, 2007		NDF	JUN 01, 1997
		4946687	AUG 07, 2007		NCE	DEC 20, 1995
		4816263	MAR 28, 2006	U-3		
		4783337	SEP 16, 2003	U-3		
		4466972	AUG 21, 2003	U-3		
20336 002	ISRADIPINE; DYNACIRC CR	5030456	JUL 09, 2008			
		4950486	AUG 21, 2007		NDF	JUN 01, 1997
		4946687	AUG 07, 2007		NCE	DEC 20, 1995
		4816263	MAR 28, 2006	U-3		
		4783337	SEP 16, 2003	U-3		
		4466972	AUG 21, 2003	U-3		
20083 001	ITRACONAZOLE; SPORANOX				I-104	MAR 29, 1997
18754 001	KETOPROFEN; ORUDIS				I-112	JAN 06, 1997
18754 002	KETOPROFEN; ORUDIS				I-112	JAN 06, 1997
18754 003	KETOPROFEN; ORUDIS				I-112	JAN 06, 1997
19816 001	KETOPROFEN; ORUVAIL				NDF	SEP 24, 1996
>ADD>	20241 001 LAMOTRIGINE; LAMICTAL				NCE	DEC 27, 1999
>ADD>	20241 002 LAMOTRIGINE; LAMICTAL				NCE	DEC 27, 1999
>ADD>	20241 003 LAMOTRIGINE; LAMICTAL				NCE	DEC 27, 1999
>ADD>	20241 004 LAMOTRIGINE; LAMICTAL				NCE	DEC 27, 1999
>ADD>	20241 005 LAMOTRIGINE; LAMICTAL				NCE	DEC 27, 1999
>ADD>	20241 006 LAMOTRIGINE; LAMICTAL				NCE	DEC 27, 1999
	20219 001 LEVOCABASTINE HYDROCHLORIDE; LIVOSTIN	4369184	JAN 18, 2000		NCE	NOV 10, 1998
	19558 006 LISINOPRIL; PRINIVIL	4374829	DEC 30, 2001		I-92	JUN 09, 1996
	19658 001 LORATADINE; CLARITIN	4282233	AUG 04, 2000	U-77	NCE	APR 12, 1998
>ADD>	19670 001 LORATADINE; CLARITIN-D				NC	NOV 14, 1997
	20264 001 MEGESTROL ACETATE; MEGACE				NDF	SEP 10, 1997
>ADD>	20357 001 METFORMIN HYDROCHLORIDE; GLUCOPHAGE				NCE	DEC 29, 1999
>ADD>	20357 002 METFORMIN HYDROCHLORIDE; GLUCOPHAGE				NCE	DEC 29, 1999
>ADD>	19737 001 METRONIDAZOLE; METROGEL	4837378	JUN 06, 2006		ODE	NOV 22, 1995
>ADD>	20208 001 METRONIDAZOLE; METROGEL	4837378	JUN 06, 2006		NDF	AUG 17, 1995

PRESCRIPTION AND OTC DRUG PRODUCT  
PATENT AND EXCLUSIVITY DATA

APPL/PROD NUMBER	INGREDIENT NAME; TRADE NAME	PATENT NUMBER	PATENT EXPIRES	USE CODE	EXCLUS CODE	EXCLUS EXPIRES
20343 001	MILRINONE LACTATE; PRIMACOR IN DEXTROSE 5%	4313951	FEB 02, 2001			
20343 002	MILRINONE LACTATE; PRIMACOR IN DEXTROSE 5%	4313951	FEB 02, 2001			
20343 003	MILRINONE LACTATE; PRIMACOR IN DEXTROSE 5%	4313951	FEB 02, 2001			
19297 001	MITOXANTRONE HYDROCHLORIDE; NOVANTRONE	4820738	APR 11, 2006	U-98		
20065 001	NAPHAZOLINE HYDROCHLORIDE; OPCON-A				NC	JUN 08, 1997
20226 001	NAPHAZOLINE HYDROCHLORIDE; NAPHCN-A				NC	JUN 08, 1997
20067 002	NAPROXEN; EC-NAPROSYN				NDF	OCT 14, 1997
20067 003	NAPROXEN; EC-NAPROSYN				NDF	OCT 14, 1997
20204 002	NAPROXEN SODIUM; ALEVE				NS	JAN 11, 1997
19660 001	NEDOCROMIL SODIUM; TILADE	4474787	OCT 02, 2006		NCE	DEC 30, 1997
>ADD>	20152 001 NEFAZODONE HYDROCHLORIDE; SERZONE	4338317	JUL 06, 1999	U-12	NCE	DEC 22, 1999
>ADD>	20152 002 NEFAZODONE HYDROCHLORIDE; SERZONE	4338317	JUL 06, 1999	U-12	NCE	DEC 22, 1999
>ADD>	20152 003 NEFAZODONE HYDROCHLORIDE; SERZONE	4338317	JUL 06, 1999	U-12	NCE	DEC 22, 1999
>ADD>	20152 004 NEFAZODONE HYDROCHLORIDE; SERZONE	4338317	JUL 06, 1999	U-12	NCE	DEC 22, 1999
>ADD>	20152 005 NEFAZODONE HYDROCHLORIDE; SERZONE	4338317	JUL 06, 1999	U-12	NCE	DEC 22, 1999
>ADD>	20152 006 NEFAZODONE HYDROCHLORIDE; SERZONE	4338317	JUL 06, 1999	U-12	NCE	DEC 22, 1999
20165 001	NICOTINE; NICODERM	5364630	APR 02, 2010			
		5344656	APR 02, 2008			
		5342623	APR 02, 2008			
20165 002	NICOTINE; NICODERM	5364630	APR 02, 2010			
		5344656	APR 02, 2008			
		5342623	APR 02, 2008			
20165 003	NICOTINE; NICODERM	5364630	APR 02, 2010			
		5344656	APR 02, 2008			
		5342623	APR 02, 2008			
19684 001	NIFEDIPINE; PROCARDIA XL	5264446	NOV 23, 2010			
19684 002	NIFEDIPINE; PROCARDIA XL	5264446	NOV 23, 2010			
19684 003	NIFEDIPINE; PROCARDIA XL	5264446	NOV 23, 2010			
19384 002	NORFLOXACIN; NOROXIN				I-113	OCT 24, 1997
19667 001	OCTREOTIDE ACETATE; SANDOSTATIN	4395403	JUL 26, 2002		I-106	MAY 03, 1997
19667 002	OCTREOTIDE ACETATE; SANDOSTATIN	4395403	JUL 26, 2002		I-106	MAY 03, 1997
19667 003	OCTREOTIDE ACETATE; SANDOSTATIN	4395403	JUL 26, 2002		I-106	MAY 03, 1997
19667 004	OCTREOTIDE ACETATE; SANDOSTATIN	4395403	JUL 26, 2002		I-106	MAY 03, 1997
19667 005	OCTREOTIDE ACETATE; SANDOSTATIN	4395403	JUL 26, 2002		I-106	MAY 03, 1997
20103 001	ONDANSETRON HYDROCHLORIDE; ZOFTRAN				I-110	SEP 26, 1997
20103 002	ONDANSETRON HYDROCHLORIDE; ZOFTRAN				I-110	SEP 26, 1997

PRESCRIPTION AND OTC DRUG PRODUCT  
PATENT AND EXCLUSIVITY DATA

APPL/PROD NUMBER	INGREDIENT NAME; TRADE NAME	PATENT NUMBER	PATENT EXPIRES	USE CODE	EXCLUS CODE	EXCLUS EXPIRES
20262 001	PACLITAXEL; TAXOL					I-105 APR 13, 1997
20036 001	PAMIDRONATE DISODIUM; AREDIA	4711880	DEC 08, 2004			D-24 JUN 22, 1997
20036 003	PAMIDRONATE DISODIUM; AREDIA	4711880	DEC 08, 2004			D-22 APR 15, 1997
20036 004	PAMIDRONATE DISODIUM; AREDIA	4711880	DEC 08, 2004			I-111 SEP 21, 1997
20031 001	PAROXETINE HYDROCHLORIDE; PAXIL	4721723	DEC 29, 2006	U-12		D-22 APR 15, 1997
20031 002	PAROXETINE HYDROCHLORIDE; PAXIL	4721723	DEC 29, 2006	U-12	NCE	I-111 SEP 21, 1997
20031 003	PAROXETINE HYDROCHLORIDE; PAXIL	4721723	DEC 29, 2006	U-12	NCE	D-22 APR 15, 1997
20031 004	PAROXETINE HYDROCHLORIDE; PAXIL	4721723	DEC 29, 2006	U-12	NCE	I-111 SEP 21, 1997
20031 005	PAROXETINE HYDROCHLORIDE; PAXIL	4721723	DEC 29, 2006	U-12	NCE	D-22 APR 15, 1997
20184 001	PERINDOPRIL ERBUMINE; ACEON	4508729	APR 02, 2002			I-111 SEP 21, 1997
20184 002	PERINDOPRIL ERBUMINE; ACEON	4508729	APR 02, 2002			D-22 APR 15, 1997
20184 003	PERINDOPRIL ERBUMINE; ACEON	4508729	APR 02, 2002			I-111 SEP 21, 1997
20237 001	PILOCARPINE HYDROCHLORIDE; SALAGEN					D-22 APR 15, 1997
19439 001	POTASSIUM CHLORIDE; K-DUR 20	4863743	SEP 05, 2006	U-99		I-111 SEP 21, 1997
19439 002	POTASSIUM CHLORIDE; K-DUR 10	4863743	SEP 05, 2006	U-99		D-22 APR 15, 1997
19898 002	PRAVASTATIN SODIUM; PRAVACHOL	4346227	JAN 09, 2004		NCE	DEC 29, 1997
19898 003	PRAVASTATIN SODIUM; PRAVACHOL	4346227	JAN 09, 2004		NCE	DEC 29, 1997
19898 004	PRAVASTATIN SODIUM; PRAVACHOL	4346227	JAN 09, 2004		NCE	DEC 29, 1997
20279 001	PREDNICARBATE; DERMATOP					NCE DEC 29, 1997
19627 001	PROPOFOL; DIPRIVAN					NCE DEC 30, 1998
20021 002	PSEUDOEPHEDRINE HYDROCHLORIDE; EFIDAC/24	4801461	MAY 05, 2004			NCE DEC 30, 1998
		4576604	MAR 18, 2003			NCE DEC 30, 1998
		4128658	DEC 05, 1995			NCE DEC 30, 1998
>ADD>	18703 001					ODE MAR 22, 2001
						NDF MAR 22, 1997
	18703 002					
>ADD>	19675 001					
	20095 001					
		4521431	JUN 04, 2002			
		5028432	JUL 02, 2008			
		4521431	JUN 04, 2002			
		4128658	DEC 05, 1995			
						I-75 MAY 19, 1995
						D-21 FEB 28, 1997

PRESCRIPTION AND OTC DRUG PRODUCT  
PATENT AND EXCLUSIVITY DATA

APPL/PROD NUMBER	INGREDIENT NAME; TRADE NAME	PATENT NUMBER	PATENT EXPIRES	USE CODE	EXCLUS CODE	EXCLUS EXPIRES
20095 002	RANITIDINE HYDROCHLORIDE; ZANTAC 300	5028432	JUL 02, 2008			
		4521431	JUN 04, 2002		I-75	MAY 19, 1995
		4128658	DEC 05, 1995		D-21	FEB 28, 1997
20251 001	RANITIDINE HYDROCHLORIDE; ZANTAC 150	5102665	APR 07, 2009			
		4521431	JUN 04, 2002		I-75	MAY 19, 1995
		4128658	DEC 05, 1995		D-21	FEB 28, 1997
20251 002	RANITIDINE HYDROCHLORIDE; ZANTAC 150	5102665	APR 07, 2009			
		4521431	JUN 04, 2002		I-75	MAY 19, 1995
		4128658	DEC 05, 1995		D-21	FEB 28, 1997
>ADD>	20474 001 RIMEXOLONE; VEXOL	4686214	AUG 11, 2004	U-100	NCE	DEC 30, 1999
	20214 001 ROCURONIUM BROMIDE; ZEMURON (P/F)	4894369	JAN 16, 2007		NCE	MAR 17, 1999
	20214 002 ROCURONIUM BROMIDE; ZEMURON	4894369	JAN 16, 2007		NCE	MAR 17, 1999
	20236 001 SALMETEROL XINAFOATE; SEREVENT	4992474	FEB 12, 2008		NCE	FEB 04, 1999
>ADD>	19839 001 SERTRALINE HYDROCHLORIDE; ZOLOFT	4962128	OCT 09, 2007	U-12		
>ADD>	19839 002 SERTRALINE HYDROCHLORIDE; ZOLOFT	4962128	OCT 09, 2007	U-12		
>ADD>	19839 003 SERTRALINE HYDROCHLORIDE; ZOLOFT	4962128	OCT 09, 2007	U-12		
>ADD>	19839 004 SERTRALINE HYDROCHLORIDE; ZOLOFT	4962128	OCT 09, 2007	U-12		
	19766 001 SIMVASTATIN; ZOCOR	4444784	DEC 24, 2005	U-59	NCE	DEC 23, 1996
	19766 002 SIMVASTATIN; ZOCOR	4444784	DEC 24, 2005	U-59	NCE	DEC 23, 1996
	19766 003 SIMVASTATIN; ZOCOR	4444784	DEC 24, 2005	U-59	NCE	DEC 23, 1996
	19766 004 SIMVASTATIN; ZOCOR	4444784	DEC 24, 2005	U-59	NCE	DEC 23, 1996
	19640 001 SOMATROPIN, BIOSYNTHETIC; HUMATROPE				D-23	APR 15, 1997
	19640 004 SOMATROPIN, BIOSYNTHETIC; HUMATROPE				D-23	APR 15, 1997
	19865 005 SOTALOL HYDROCHLORIDE; BETAPACE				NCE	OCT 30, 1997
					ODE	OCT 30, 1999
>ADD>	20240 001 SPIRAPRIL HYDROCHLORIDE; RENORMAX				NCE	DEC 29, 1999
>ADD>	20240 002 SPIRAPRIL HYDROCHLORIDE; RENORMAX				NCE	DEC 29, 1999
>ADD>	20240 003 SPIRAPRIL HYDROCHLORIDE; RENORMAX				NCE	DEC 29, 1999
>ADD>	20240 004 SPIRAPRIL HYDROCHLORIDE; RENORMAX				NCE	DEC 29, 1999
	20412 001 STAVUDINE; ZERIT	4978655	DEC 18, 2007	U-94	NCE	JUN 24, 1999
	20412 002 STAVUDINE; ZERIT	4978655	DEC 18, 2007	U-94	NCE	JUN 24, 1999
	20412 003 STAVUDINE; ZERIT	4978655	DEC 18, 2007	U-94	NCE	JUN 24, 1999
	20412 004 STAVUDINE; ZERIT	4978655	DEC 18, 2007	U-94	NCE	JUN 24, 1999
	20412 005 STAVUDINE; ZERIT	4978655	DEC 18, 2007	U-94	NCE	JUN 24, 1999
	40091 001 SULFADIAZINE; SULFADIAZINE				ODE	JUL 29, 2001
	17376 001 SULFAMETHOXAZOLE; SEPTRA	4209513	JUN 24, 1997		I-103	JAN 07, 1997
	17376 002 SULFAMETHOXAZOLE; SEPTRA DS	4209513	JUN 24, 1997		I-103	JAN 07, 1997

PRESCRIPTION AND OTC DRUG PRODUCT  
PATENT AND EXCLUSIVITY DATA

APPL/PROD NUMBER	INGREDIENT NAME; TRADE NAME	PATENT NUMBER	PATENT EXPIRES	USE CODE	EXCLUS CODE	EXCLUS EXPIRES
17377 001	SULFAMETHOXAZOLE; BACTRIM				I-103	JAN 07, 1997
17377 002	SULFAMETHOXAZOLE; BACTRIM DS				I-103	JAN 07, 1997
17560 002	SULFAMETHOXAZOLE; BACTRIM PEDIATRIC				I-103	JAN 07, 1997
17598 001	SULFAMETHOXAZOLE; SEPTRA				I-103	JAN 07, 1997
17598 002	SULFAMETHOXAZOLE; SEPTRA GRAPE				I-103	JAN 07, 1997
20080 001	SUMATRIPTAN SUCCINATE; IMITREX	4816470	DEC 29, 2006	U-72		
20070 001	TACRINE HYDROCHLORIDE; COGNEX	4631286	DEC 23, 2003	U-97		
20070 002	TACRINE HYDROCHLORIDE; COGNEX	4631286	DEC 23, 2003	U-97		
20070 003	TACRINE HYDROCHLORIDE; COGNEX	4631286	DEC 23, 2003	U-97		
20070 004	TACRINE HYDROCHLORIDE; COGNEX	4631286	DEC 23, 2003	U-97		
17970 002	TAMOXIFEN CITRATE; NOLVADEX	4536516	AUG 20, 2002			
>ADD>	20256 001	TECHNETIUM TC-99M BICISATE KIT; NEUROLITE	5279811	MAR 18, 2008	U-101	NCE NOV 23, 1999
	18163 003	TEMAZEPAM; RESTORIL	5326758	JUL 09, 2008	U-70	
>ADD>	20347 001	TERAZOSIN HYDROCHLORIDE; HYTRIN	5294615	MAR 15, 2011		
>ADD>			5212176	MAY 18, 2010		
>ADD>			4251532	FEB 17, 2000		
>ADD>			4112097	SEP 05, 1995		
>ADD>	20347 002	TERAZOSIN HYDROCHLORIDE; HYTRIN	5294615	MAR 15, 2011		
>ADD>			5212176	MAY 18, 2010		
>ADD>			4251532	FEB 17, 2000		
>ADD>			4112097	SEP 05, 1995		
>ADD>	20347 003	TERAZOSIN HYDROCHLORIDE; HYTRIN	5294615	MAR 15, 2011		
>ADD>			5212176	MAY 18, 2010		
>ADD>			4251532	FEB 17, 2000		
>ADD>			4112097	SEP 05, 1995		
>ADD>	20347 004	TERAZOSIN HYDROCHLORIDE; HYTRIN	5294615	MAR 15, 2011		
>ADD>			5212176	MAY 18, 2010		
>ADD>			4251532	FEB 17, 2000		
>ADD>			4112097	SEP 05, 1995		
	20192 001	TERBINAFINE HYDROCHLORIDE; LAMISIL	4755534	JUL 05, 2005	U-73	NCE DEC 30, 1999
	19762 001	TESTOSTERONE; TESTODERM	4867982	FEB 16, 2005		
			4725439	FEB 16, 2005		
			4704282	NOV 03, 2004	NDF	OCT 12, 1996
	19762 002	TESTOSTERONE; TESTODERM	4867982	FEB 16, 2005		
			4725439	FEB 16, 2005		
			4704282	NOV 03, 2004	NDF	OCT 12, 1996

PRESCRIPTION AND OTC DRUG PRODUCT  
PATENT AND EXCLUSIVITY DATA

APPL/PROD NUMBER	INGREDIENT NAME; TRADE NAME	PATENT NUMBER	PATENT EXPIRES	USE CODE	EXCLUS CODE	EXCLUS EXPIRES	
20330 001	TIMOLOL MALEATE; TIMOPTIC-XE	4861760	AUG 29, 2006				
		4195085	MAR 25, 1997		NP	NOV 04, 1996	
20330 002	TIMOLOL MALEATE; TIMOPTIC-XE	4861760	AUG 29, 2006				
		4195085	MAR 25, 1997		NP	NOV 04, 1996	
20326 001	TRIMETREXATE GLUCURONATE; NEUTREXIN	4694007	SEP 15, 2004	U-91	ODE	DEC 17, 2000	
>ADD>	20388 001	VINORELBINE TARTRATE; NAVELBINE			NCE	DEC 23, 1999	
	19908 001	ZOLPIDEM TARTRATE; AMBIEN	4382938	MAY 10, 2005	U-74	NCE	DEC 16, 1997
	19908 002	ZOLPIDEM TARTRATE; AMBIEN	4382938	MAY 10, 2005	U-74	NCE	DEC 16, 1997