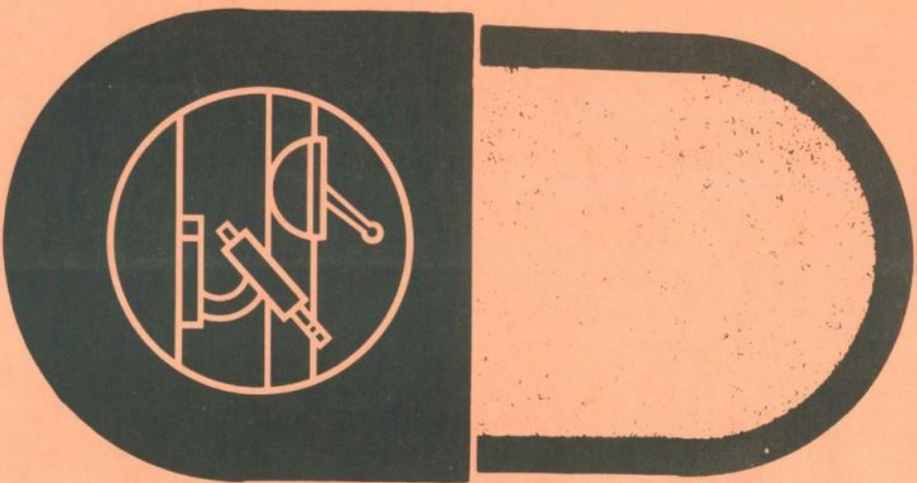
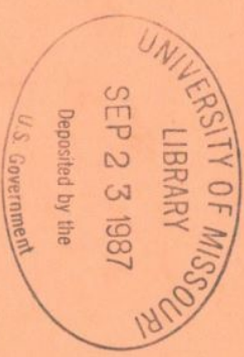


HE 20.4210  
987/Supp. 7



**CUMULATIVE  
SUPPLEMENT 7  
JAN'87-JUL'87**



# **APPROVED DRUG PRODUCTS**

**WITH  
THERAPEUTIC EQUIVALENCE EVALUATIONS  
7<sup>TH</sup> EDITION**

**U.S. DEPARTMENT OF HEALTH AND HUMAN SERVICES  
PUBLIC HEALTH SERVICE  
FOOD AND DRUG ADMINISTRATION  
CENTER FOR DRUGS AND BIOLOGICS**

APPROVED DRUG PRODUCTS  
with  
THERAPEUTIC EQUIVALENCE EVALUATIONS  
7TH EDITION

CUMULATIVE SUPPLEMENT 7

JULY 1987

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APPROVED DRUG PRODUCTS  
with  
THERAPEUTIC EQUIVALENCE EVALUATIONS  
7th EDITION  
CUMULATIVE SUPPLEMENT 7  
JULY 1987

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, 7th Edition (the List). The List is composed of three parts: approved prescription drug products with therapeutic equivalence evaluations, over-the-counter (OTC) drug products that require approved applications as a condition of marketing, and drug products approved by the Division of Blood and Blood Products under Section 505 of the Act.

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 Approved Under Section 505 of the Act by the Division of Blood and Blood Products 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 left of the active ingredient name for each product, along with the application number and product number (FDA's internal file number). 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 for an explanation of the use codes and exclusivity 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 effective (marketing) date (the date a product may be marketed), when appropriate, will appear to the left of the approval date.

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. A newly approved product is also identified by a lozenge (⌘) to the right of its strength which remains throughout all Cumulative Supplements for this edition.

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 overstruck print. The >\_DLT\_> symbol is dropped in subsequent Cumulative Supplements for that item. The overstruck print will remain in the Prescription and OTC Drug Product Lists in all Cumulative Supplements for this edition. However, the overstruck print in the Patent and Exclusivity Data will be dropped in subsequent Cumulative Supplements.

Products discontinued from marketing or that have had their application withdrawn, for other than safety or effectiveness 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 7th Edition List will then be added to the "Discontinued Drug Product List" appearing in the 8th Edition.

## 1.2 PREDNISONE BIOEQUIVALENCE

The Agency has determined that in vitro data are sufficient to demonstrate bioequivalence of prednisone products. This decision is based on past bioavailability studies on a variety of prednisone products sponsored under FDA contract which established an in vitro and in vivo correlation with a variety of in vitro apparatus and media. The studies demonstrated that the dissolution rate using apparatus such as the spin filter, USP basket, and paddle correlated with the rate of drug absorption. The initial paddle used in the above studies was a tilting blade paddle. When the USP adopted a fixed blade paddle method, it raised the issue of whether

the same correlation existed for the tilting blade paddle. Following the October 15, 1977, effective date of the new USP prednisone tablet dissolution specification, the Agency initiated an extensive voluntary dissolution certification program for all marketed prednisone tablet products. This program continued until each firm demonstrated that every prednisone product could consistently meet the new USP standard. Firms failing to meet the new standard were required to remove their product from the market or reformulate to an acceptable product. As a result of this program, when marketed prednisone tablet products were resurveyed in 1980, all met the USP standard.

A selected sample of the products in an Agency bioavailability study conducted in 1982 on marketed prednisone tablets revealed no statistically significant differences in the key bioavailability parameters (AUC, C<sub>max</sub>, T<sub>max</sub>) for prednisone tablets.

Therefore, FDA will change the therapeutic equivalence code from BX to AB on any approved prednisone tablets if the application is supplemented with an acceptable comparative in vitro dissolution study. (See Section 3.7 of the 7th Edition List for available guidance from the Division of Bioequivalence.)

### 1.3 OTC DRUG PRODUCTS

The following drug products identified in the "OTC Drug Product List" of this publication as requiring approved applications may be marketed on the firm's own responsibility without an application under the Agency's existing OTC drug marketing policies so long as applicable proposed or tentative final monographs are followed (see 21 CFR 330.13).

Pseudoephedrine Hydrochloride	60mg
Triprolidine Hydrochloride Tablet or Capsule; Oral	2.5mg
Pseudoephedrine Hydrochloride	30mg/5ml
Triprolidine Hydrochloride Syrup; Oral	1.25mg/5ml
Triprolidine Hydrochloride Syrup; Oral	1.25mg/5ml
Triprolidine Hydrochloride Tablet; Oral	2.5mg

#### 1.4 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 7, 1984 (49 FR 35428)
Nitroglycerin (ointment;topical)	SEP 3, 1986 (51 FR 31371)
Nitroglycerin (tablet, controlled release;oral)	SEP 7, 1984 (49 FR 35428)
Nitroglycerin (tablet, controlled release;buccal)	JUL 5, 1985 (50 FR 27688)
Phenazopyridine Hydrochloride and Sulfamethoxazole	JUL 29, 1983 (48 FR 34516)
Tranlylcypromine Sulfate	MAR 22, 1984 (49 FR 10708)

#### 1.5 GAVICON

Gaviscon is an over-the-counter (OTC) product which has been marketed since September 1970. The active ingredients, aluminum hydroxide and magnesium trisilicate, for this product were reviewed by the OTC's Antacid Panel and were considered to be safe and effective ingredients (Category I) by that panel. However, the tablet failed to pass the antacid test which is required of all antacid products. It was, therefore, placed in Category III for lack of effectiveness. A full NDA with clinical studies was submitted by Marion Laboratories, Inc., and approved by FDA, December 9, 1983. Gaviscon's activity in treating reflux acidity is made possible by the physical-chemical properties of the inactive ingredients, sodium bicarbonate and alginic acid. Therefore, all ANDAs which cite Gaviscon tablets as the listed drug must contain the inactive ingredients, sodium bicarbonate and alginic acid. A full NDA will be required to support the effectiveness of the drug product if different inactive ingredients are substituted for sodium bicarbonate or alginic acid or if different proportions of these ingredients are used.

#### 1.6 APPLICANT (NAME) CHANGES

Because 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, the cumulation of these transfers and name changes will be identified in this section only. Where only partial approved product lines are transferred between applicants, each approved product involved will appear as an applicant name change in the Cumulative Supplement.

APPLICANT (NAME) CHANGES

<u>FORMER APPLICANT (NAME)</u>	<u>NEW APPLICANT (NAME)</u>	<u>NEW ABBREVIATED NAME</u>
COOPERVISION PHARMS	IOLAB PHARMACEUTICALS	IOLAB
CARTER-GLOGAU LABORATORIES	STERIS LABORATORIES	STERIS LABS
ASCOT HOSPITAL PHARMACEUTICALS	ASCOT DIVISION OF TRAVENOL LABORATORIES	ASCOT
WILLIAM H RORER INC	RORER PHARMACEUTICAL CORP SUB RORER GROUP	RORER PHARM
USV (PR) DEVELOPMENT CORPORATION	RORER PHARMACEUTICAL CORP SUB RORER GROUP	RORER PHARM
USV LABORATORIES INC	RORER PHARMACEUTICAL CORP SUB RORER GROUP	RORER PHARM
USV PHARMACEUTICAL CORP	RORER PHARMACEUTICAL CORP SUB RORER GROUP	RORER PHARM

1.7 CONJUGATED ESTROGEN TABLETS

Conjugated estrogen tablets are presently coded BS (not therapeutically equivalent) based on in vivo data indicating differences produced by different conjugated estrogen tablets in urinary excretion levels of the active ingredients. These differences were believed to be directly related to the differences in composition permitted by the official standards for the estrogenic steroids in conjugated estrogen products. The USP monograph was recently revised to narrow the range of differences permitted.

Nevertheless, FDA's Biopharmaceutics Research Branch recently demonstrated problems with dissolution of conjugated estrogen tablets, apparently because of the products' coating. The coating on at least some conjugated estrogen products behaves like an enteric coating. Therefore, the Agency has decided to require in vivo bioequivalence studies for all new applications for conjugated estrogen tablets and for any such product to be coded AB (therapeutically equivalent). Thus, all new or pending applications for conjugated estrogen tablets must contain in vivo studies and previously approved conjugated estrogen tablets will be coded as BP (not therapeutically equivalent) unless an acceptable in vivo bioequivalence study is submitted by the applicant holder. Requests for guidance on conducting bioavailability/bioequivalence studies should be addressed to the Division of Bioequivalence, HFN-250, 5600 Fishers Lane, Rockville, MD 20857.

## 1.8 CORRECTIONS TO THE 7TH EDITION

- a. The locator tab for the "OTC Drug Product List" is placed incorrectly within the List.
- b. There is no locator tab on the back cover for the "Discontinued Drug Product List."
- c. A recent approval has shown that the language in the "BC" code definition did not accurately reflect the use of the BC code for controlled-release products which may meet bioequivalence criteria for approval, but differ in rate such that they would not be considered therapeutically equivalent.

Therefore, please note that on pages 1-5 and 1-6 of the Introduction to the Approved Drug Products with Therapeutic Equivalence Evaluations, 7th Edition, the language defining the AB and BC codes has been revised.

### AB

#### Products meeting necessary bioequivalence requirements

The AB evaluation generally denotes products that: (1) contain an active ingredient in a dosage form for which the submission of bioavailability or clinical data is required for approval or to permit therapeutic equivalence evaluations, and (2) for which the applicant has provided adequate studies to establish the bioavailability and bioequivalence of its product. Products generally will be coded AB if a study is submitted demonstrating bioequivalence, even if the study currently is not required for approval. This category also includes those few drugs with more than one approved application but only one manufacturer. It should be noted that if only one product under a drug ingredient heading is coded AB, it signifies that only that product is supported by bioavailability data. It does not signify that this product is therapeutically equivalent to the other drugs under the same heading. Thus, one product under a drug ingredient heading, coded AB is not therapeutically equivalent to a drug product under the same heading that is coded BD, BP, or BT. Drugs coded AB under an ingredient heading are considered therapeutically equivalent only to other drugs coded AB under that heading.

### BC

#### Controlled-release tablets, controlled-release capsules, and controlled-release injectables

Although bioavailability studies have been conducted on these dosage forms, they are subject to bioavailability differences, primarily because firms developing controlled-release products for the same active ingredient rarely employ the same formulation approach. FDA, therefore, does not evaluate different controlled-release dosage forms

containing the same active ingredient in equal strength as therapeutically equivalent unless equivalence between individual products for both rate and extent has been specifically demonstrated through appropriate bioequivalence studies. Controlled-release products for which such bioequivalence data are available have been coded AB.

- d. In the following products dextrose and sodium chloride are considered vehicles and not active ingredients, therefore, they will no longer appear as part of the active ingredient heading. These ingredients may continue to appear in the trade name for those products which contain them. The active ingredient headings in the 7th Edition affected are:

Alcohol; Dextrose  
Aminophylline; Sodium Chloride  
Ammonium Chloride; Sodium Chloride  
Bretylum Tosylate; Dextrose  
Cefazolin Sodium; Dextrose  
Cefoperazone Sodium; Dextrose  
Cefotaxime Sodium; Dextrose  
Cefotaxime Sodium; Sodium Chloride  
Cefoxitin Sodium; Dextrose  
Cefoxitin Sodium; Sodium Chloride  
Ceftizoxime Sodium; Dextrose  
Cephalothin Sodium; Dextrose  
Cephalothin Sodium; Sodium Chloride  
Cimetidine Hydrochloride; Sodium Chloride  
Dextrose; Dopamine Hydrochloride  
Dextrose; Gentamicin Sulfate  
Dextrose; Lidocaine Hydrochloride  
Dextrose; Heparin Sodium  
Dextrose; Mannitol  
Dextrose; Oxytocin  
Dextrose; Theophylline  
Gentamicin Sulfate; Sodium Chloride  
Heparin Sodium; Sodium Chloride  
Ranitidine Hydrochloride; Sodium Chloride

- e. The following products are corrections to a printing error that appeared on page 3-204. Please record the correct NDA Numbers in the List.

PROCAINAMIDE HYDROCHLORIDE

CAPSULE; ORAL;  
PROCAINAMIDE HCL  
LEDERLE LABS/AM CYAN

	<u>375MG</u>	N86952 001
	<u>500MG</u>	N86943 001
VANGARD LABS/MWM	<u>250MG</u>	N87643 001

## 1.9 CHANGE OF A THERAPEUTIC EQUIVALENCE CODE FOR A DRUG ENTITY

This section explains the procedures the Agency will use when, in response to a petition or on its own initiative, it is considering a change in the therapeutic equivalence code for approved multisource drug products. Such changes will generally occur when the Agency becomes aware of new scientific information affecting therapeutic equivalence. These procedures will be used when all drug products found in the "Drug Product List" under a specific drug entity and dosage form are being considered for a change. The change may be from the code signifying that the drug does not present a bioequivalence problem drug (e.g., AA) to a code signifying a bioequivalence problem (e.g., BP), or vice versa. A change of a single product code from BP to AB as a result of a bioequivalence study is not applicable in this section.

This section lists those drug entities that are actively being considered by the Agency for reclassification. Before making a change in the code, the Agency will announce in this section of the Cumulative Supplement that it is considering the change and will invite comment. Comments, along with scientific data, may be sent to the Division of Bioequivalence, HFN-250, Room 17B06, 5600 Fishers Lane, Rockville, MD 20857. The comment period will generally be 60 days in length, and the closing date for comments will be listed in the description of the proposed change for each drug entity.

The most useful type of scientific data is an in vivo bioavailability/bioequivalence study conducted on batches of the subject drug. These submissions should present a full description of the analytical procedures and equipment used, a validation of the analytical methodology, including the standard curve, a description of the method of calculating results, and a description of the pharmacokinetic and statistical models used in analyzing the data. Anecdotal or testimonial information is the least useful to the Agency, and such submissions are discouraged. However, copies of supporting reports published in the scientific literature or unpublished material are welcome.

The Agency is currently considering a change in therapeutic equivalence evaluation for the following drug(s):

### Benztropine mesylate:

The Agency initially did not classify benztropine mesylate as having an actual or potential bioequivalence problem. (42 FR 1624, January 7, 1977). Benztropine mesylate tablets (Cogentin) is a DESI drug product that was raised to the effective status on November 7, 1970 (35 FR 211). It remained single source until January 1984. At that time, the Agency reviewed its status regarding a potential bioequivalence problem. Based principally on a published article, Tune, L., and Coyle, J.T., "Acute Extrapyrarnidal Side Effects: Serum Levels of Neuroleptics and Anticholinergics," *Psychopharmacology*, 1981;75:9-15, the Agency decided that benztropine mesylate did present a potential bioequivalence problem because of the possibility of nonlinear kinetics. As a result, an in vivo bioequivalence study was required to demonstrate bioequivalence and to gain approval of an ANDA.

Recently, two pharmaceutical firms have asked the Agency to change the therapeutic equivalence code for benztropine mesylate oral tablets from BP to AA. Although the Agency disagrees with the arguments on the basis that the requests were primarily legal and regulatory, the Agency used the opportunity to reassess the merits of its earlier decision. Upon a careful re-review of the article in question and another search of the literature, the Agency now believes that there is an insufficient basis upon which to evaluate benztropine mesylate as having a potential bioequivalence problem. In addition, one of the authors of the article has advised the Agency that he does not believe the data in the article provide a basis for concluding that benztropine mesylate displays nonlinear kinetics. In addition, the drug is freely soluble in water and does not generally meet the criteria, described in 21 CFR 320.52, for a drug posing a bioequivalence problem.

The Agency requests that interested parties submit comments with respect to the Agency's proposal to change the therapeutic equivalence code for listed benztropine mesylate oral tablets from BP to AA. We request that such comments be received no later than September 30, 1987.

Nortriptyline hydrochloride:

Presently, Eli Lilly and Sandoz Pharmaceuticals have received approval to market nortriptyline hydrochloride capsules, Aventyl and Pamelor, respectively. A recent article, Dubovsky, S.L., "Single Case Study: Severe Nortriptyline Intoxication due to Change from Generic to a Trade Preparation," Journal of Nervous and Mental Disease, 1987;175:115-17. indicates that it would be appropriate to change the therapeutic equivalence code for Aventyl and Pamelor from BP to BD.

The Agency will change the therapeutic equivalence code of nortriptyline hydrochloride capsules from BP to BD unless scientific data are submitted that adequately controvert the evidence presented in the cited article. The Agency is soliciting comments from interested parties who desire to submit scientific data in support of, or in disagreement with, this proposal. We request that such comments be received no later than October 30, 1987.

## 1.10 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. Thus, 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 those approved drug products marketed by distributors; those marketed solely abroad; and products now regarded as medical devices, biologics or foods.

The counts appear in two sections. Section A. provides baseline and quarterly data. The baseline column refers to the products in the List. For each three-month period following December '86, a column of quarterly data is added which incorporates counts of product activity from the previous quarter(s) with those in the baseline count. Section B. refers to products in the Cumulative Supplements and provides monthly activity with a cumulative count for the current quarter.

### 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 part of a combination.

### USE OF REPORT

From the data presented under Section B., users should be able to observe such things as (1) newly approved and remarketed drug products which are added to the List; (2) products that are being removed from the List as the result of withdrawal of approval and changes from prescription to over-the-counter status; and, (3) trends in approval of products as either multisource or single source during each month within the quarter. The report does not reflect category changes from multisource to single source and vice versa. However, the net gain that results from all additions, deletions and category changes is reflected in the quarterly counts for multisource and single source products.

REPORT OF COUNTS FOR THE DRUG PRODUCT LISTA. COUNTS CUMULATIVE BY QUARTERS

<u>CATEGORIES COUNTED</u>	<u>DEC '86 (BASELINE)</u>	<u>MAR '87</u>
DRUG PRODUCTS LISTED	8957	9183
SINGLE SOURCE	2103 (23.5%)	2095 (22.8%)
MULTISOURCE (1)	6854 (76.5%)	7088 (77.2%)
THERAPEUTICALLY EQUIVALENT	5838 (65.2%)	6093 (66.4%)
NOT THERAPEUTICALLY EQUIVALENT	967 (10.8%)	950 (10.3%)
EXCEPTIONS (2)	49 ( 0.5%)	45 ( 0.5%)
NEW MOLECULAR ENTITIES APPROVED	-	2
NUMBER OF APPLICANTS	333	334

B. ACTIVITY FOR SUPPLEMENT NUMBER 7

	<u>JUL '87</u>	<u>CUMULATIVE</u>
DRUG PRODUCTS ADDED:		
NEWLY APPROVED	76	76
DESI EFFECTIVE	0	0
REMARKETED	0	0
DRUG PRODUCTS REMOVED:	0	0
WITHDRAWN APPROVAL	0	0
RX TO OTC SWITCH	0	0
NET GAIN IN DRUG PRODUCTS	76	76
SINGLE SOURCE PRODUCTS APPROVED	7	7
MULTISOURCE DRUG PRODUCTS APPROVED	69	69
NEW MOLECULAR ENTITIES APPROVED:	0	0
AS THE ENTITY	0	0
AS A SALT, ESTER OR DERIVATIVE	0	0
OF THE ENTITY	0	0

(1) THERAPEUTIC EQUIVALENCE EVALUATIONS PROVIDED ONLY FOR MULTISOURCE PRODUCTS (I.E., AVAILABLE FROM MORE THAN ONE APPLICANT)

(2) AMINO ACID-CONTAINING PRODUCTS OF VARYING COMPOSITION (SEE INTRODUCTION, PAGE 1-8 OF THE LIST)



ALBUTEROL SULFATE

SOLUTION; INHALATION

VENTOLIN  
GLAXO

N19269 002  
JAN 16, 1987

EQ 0.5% BASEM

>\_ADD\_>  
>\_ADD\_>  
>\_ADD\_>  
>\_ADD\_>  
>\_ADD\_>  
>\_ADD\_>  
>\_ADD\_>

INJECTABLE; INJECTION  
AMINOSYN 10% (PH6)  
ABBOTT LABS

10%

N17673 008  
NOV 18, 1985

SYRUP; ORAL  
PROVENTIL  
SCHERING

N18062 001  
JAN 19, 1983

EQ 2MG BASE/5ML

7%

N17673 006  
NOV 18, 1985

VENTOLIN  
GLAXO

N19621 001  
JUN 10, 1987

EQ 2MG BASE/5MLM

8.5%

N17673 007  
NOV 18, 1985

TABLET, CONTROLLED RELEASE; ORAL

PROVENTIL  
SCHERING

N19383 001  
JUL 13, 1987

EQ 4MG BASEM

INJECTABLE; INJECTION  
AMINOCAPROIC ACID IN PLASTIC CONTAINER  
ABBOTT LABS

250MG/MLM

N70010 001  
MAR 09, 1987

ALLOPURINOL

TABLET; ORAL

ALLOPURINOL  
MUTUAL PHARM

N71449 001  
JAN 09, 1987

100MGH

TABLET; ORAL  
AMITRIPTYLINE HCL  
BARR LABS

150MGH

N89423 001  
FEB 17, 1987

AB

AB

N71450 001  
JAN 09, 1987

300MGH

10MG/  
25MG/  
50MG/  
75MG/  
100MG/  
150MG/  
10MG  
25MG  
50MG  
75MG  
100MG  
150MG

N86616/001/  
N86859/001/  
N86857/001/  
N86860/001/  
N86854/001/  
N86853/001/  
N86610 001  
N86859 001  
N86857 001  
N86860 001  
N86854 001  
N86853 001

LOPURIN

BOOTS PHARMS

N71586 001  
APR 02, 1987

100MGH

EMMON

N71587 001  
APR 02, 1987

300MGH

AMANTADINE HYDROCHLORIDE

CAPSULE; ORAL

AMANTADINE HCL  
BOLAR PHARM

N71382 001  
JAN 21, 1987

100MGH

EMMON

N71293 001  
FEB 18, 1987

100MGH

AMILORIDE HYDROCHLORIDE; HYDROCHLOROTHIAZIDE

TABLET; ORAL

AMILORIDE HCL AND HYDROCHLOROTHIAZIDE  
BIOCRAFT LABS

N70795 001  
APR 17, 1988

5MG;50MGH

EMMON

N70795 001  
APR 17, 1988

5MG;50MGH

EMMON

AMITRIPTYLINE HYDROCHLORIDE

TABLET; ORAL  
AMITRIPTYLINE HCL  
MUTUAL PHARM

> ADD > AB N89398 001 10MGx  
> ADD > AB JUL 14, 1987  
> ADD > AB N89399 001 25MGx  
> ADD > AB JUL 14, 1987  
> ADD > AB N89400 001 50MGx  
> ADD > AB JUL 14, 1987  
> ADD > AB N89401 001 75MGx  
> ADD > AB JUL 14, 1987  
> ADD > AB N89402 001 100MGx  
> ADD > AB JUL 14, 1987  
> ADD > AB N89403 001 150MGx  
> ADD > AB JUL 14, 1987

AMITRIPTYLINE HYDROCHLORIDE; PERPHENAZINE

TABLET; ORAL  
PERPHENAZINE AND AMITRIPTYLINE HCL  
CHELSEA LABS

AB N71558 001 50MG;4MGx  
MAR 02, 1987

AMPHOTERICIN B

INJECTABLE; INJECTION  
AMPHOTERICIN B

AP LYPHOMED 50MG/VIALx

FUNGIZONE  
SQUIBB

AP 50MG/VIAL N60517 001

AMPICILLIN SODIUM

INJECTABLE; INJECTION  
AMPICILLIN SODIUM

AP IBI SPA EQ 250MG BASE/VIALx

AP EQ 500MG BASE/VIALx

AP EQ 1GM BASE/VIALx

AP EQ 1GM BASE/VIALx

AP EQ 2GM BASE/VIALx

N62719 001  
MAY 12, 1987

N62719 003  
MAY 12, 1987

N62719 002  
MAY 12, 1987

N62634 002  
JAN 09, 1987

N62634 003  
JAN 09, 1987

AMPICILLIN SODIUM

INJECTABLE; INJECTION  
POLYICILLIN-N

AP BRISTOL LABS EQ 1GM BASE/VIALx N62738 001  
FEB 19, 1987  
AP EQ 2GM BASE/VIALx N62738 002  
FEB 19, 1987

ASPIRIN; CAFFEINE; ORPHENADRINE CITRATE

TABLET; ORAL  
MORGESIC  
RIKER LABS 385MG;30MG;25MG N13416 003  
OCT 27, 1982  
MORGESIC FORTE  
RIKER LABS 770MG;60MG;50MG N13416 004  
OCT 27, 1982

AB MORGESIC  
PAR PHARM 385MG;30MG;25MG N71642 001  
JUN 23, 1987  
AB ORPHENGESIC FORTE  
PAR PHARM 770MG;60MG;50MG N71643 001  
JUN 23, 1987

ASPIRIN; MEPROBAMATE

TABLET; ORAL  
MEPROGESIC  
VITARINE

AB 325MG;200MGx N89127 001  
MAR 02, 1987

/AB/ MEPROGESIC /  
/QUANTUM PHARMCS/

/325MG;200MG/ N66746/661/  
/JUN/81;/1984/

Q-GESIC

AB QUANTUM PHARMCS 325MG;200MG N88740 001  
JUN 01, 1984

ATROPINE

INJECTABLE; INJECTION

ATROPEN

AP SURVIVAL TECH EQ 2MG SULFATE/0.7ML N17106 001

ATROPINE

AP KALI DUPHAR EQ 2MG SULFATE/0.7MLx N71295 001  
JAN 30, 1987

BACITRACIN

INJECTABLE; INJECTION

BACITRACIN

QUAD PHARMS

AP

10,000 UNITS/VIAL

N62696 001

APR 17, 1987

50,000 UNITS/VIAL

N62696 002

APR 17, 1987

AP UPJOHN

10,000 UNITS/VIAL

N60733 001

BECLOMETHASONE DIPROPIONATE

SPRAY; INHALATION/NASAL

BECONASE AQ

GLAXO

> ADD >

> ADD >

> ADD >

0.042MG/INH

N19389 001

JUL 27, 1987

BETAMETHASONE

CREAM; TOPICAL

CELESTONE

SCHERING

0.2%

N14762 001

BETAMETHASONE DIPROPIONATE

CREAM; TOPICAL

BETAMETHASONE DIPROPIONATE

NYC LABS

AB

EQ 0.05% BASE

N70885 001

FEB 03, 1987

THAMES PHARMA

EQ 0.05% BASE

N71143 001

JUN 17, 1987

DIPROLENE AF

EQ 0.05% BASE

N19555 001

APR 27, 1987

LOTION; TOPICAL

BETAMETHASONE DIPROPIONATE

NYC LABS

AB

EQ 0.05% BASE

N71085 001

FEB 03, 1987

ointment; TOPICAL

BETAMETHASONE DIPROPIONATE

NYC LABS

AB

EQ 0.05% BASE

N71012 001

FEB 03, 1987

BETAMETHASONE VALERATE

CREAM; TOPICAL

BETAMETHASONE VALERATE

PHARMAFAIR

AB

EQ 0.1% BASE

N70485 001

MAY 29, 1987

LOTION; TOPICAL

BETAMETHASONE VALERATE

PHARMAFAIR

AB

EQ 0.1% BASE

N70484 001

MAY 29, 1987

ointment; TOPICAL

BETAMETHASONE VALERATE

PHARMAFAIR

AB

EQ 0.1% BASE

N70486 001

MAY 29, 1987

BLEOMYCIN SULFATE

INJECTABLE; INJECTION

BLENOXANE

BRISTOL LABS

/NIPPON/KAYAKU/

N14762 001

N50443 001

/EQ/15 UNITS BASE/VIAL

/EQ/15 UNITS/BASE/VIAL/

/N61847/001/

BRETYLIUM IOSYLATE

INJECTABLE; INJECTION

BRETYLIUM IOSYLATE

LYPHOMED

100MG/ML

N71298 001

FEB 13, 1987

BUPIVACAINE HYDROCHLORIDE

INJECTABLE; INJECTION

BUPIVACAINE HCL

ABBOTT LABS

AP		0.25/M	N70583 001	FEB 17, 1987
AP		0.25/M	N70586 001	MAR 03, 1987
AP		0.25/M	N70590 001	FEB 17, 1987
AP		0.5/M	N70584 001	FEB 17, 1986
AP		0.5/M	N70597 001	MAR 03, 1987
AP		0.5/M	N70609 001	MAR 03, 1987
AP		0.75/M	N70585 001	MAR 03, 1987
AP		0.75/M	N70587 001	MAR 03, 1987
AP		0.75/M	N71202 001	APR 15, 1987

SENSORCAINE

ASTRA PHARM PRODS

CALCIUM GLUCEPTATE

INJECTABLE; INJECTION

CALCIUM GLUCEPTATE

LYPHOMED

AP		EQ 90MG CALCIUM/5MLM	N89373 001	APR 30, 1987
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CARBAMAZEPINE

TABLET; ORAL

CARBAMAZEPINE

PARKE DAVIS

AB		200MG	N70429 001	JAN 02, 1987
AB		200MG	N71479 001	JUL 24, 1987

> ADD >  
> ADD >

CEFADROXIL

CAPSULE; ORAL

CEFADROXIL

ZENITH LABS

AB		EQ 500MG BASEM	N62766 001	MAR 03, 1987
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CEFADROXIL

TABLET; ORAL

CEFADROXIL

ZENITH LABS

AB		EQ 1GM BASEM	N62774 001	APR 08, 1987
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CEFOPERAZONE SODIUM

INJECTABLE; INJECTION

CEFOBID IN PLASTIC CONTAINER

ROERIG

> ADD >		EQ 20MG BASE/MLM	N50613 002	JUL 31, 1987
> ADD >				

CEFOTAXIME SODIUM

INJECTABLE; INJECTION

CLAFORAN

HOECHST

		EQ 1GM BASE/VIALM	N62659 001	JAN 13, 1987
		EQ 2GM BASE/VIALM	N62659 002	JAN 13, 1987

CEFOXITIN SODIUM

INJECTABLE; INJECTION

MEFOXIN

MS&D

		EQ 1GM BASE/VIALM	N62757 001	JAN 08, 1987
		EQ 2GM BASE/VIALM	N62757 002	JAN 08, 1987

CEFTRIAZONE SODIUM

INJECTABLE; INJECTION

ROCEPHIN

ROCHE

		EQ 500MG BASE/VIALM	N62654 001	APR 30, 1987
		EQ 1GM BASE/VIALM	N62654 002	APR 30, 1987
		EQ 2GM BASE/VIALM	N62654 003	APR 30, 1987

ROCEPHIN W/ DEXTROSE IN PLASTIC CONTAINER

ROCHE

		EQ 10MG BASE/MLM	N50624 001	FEB 11, 1987
		EQ 20MG BASE/MLM	N50624 002	FEB 11, 1987
		EQ 40MG BASE/MLM	N50624 003	FEB 11, 1987



CEPHRADINE

POWDER FOR RECONSTITUTION; ORAL

CEPHRADINE

BIOCRAFT LABS

125MG/5MLM

N62693 001  
JAN 09, 1987

INJECTABLE; INJECTION  
CHROMIC CHLORIDE

AP LYPHOMED

EQ 0.004MG CHROMIUM/MLM N19271 001  
MAY 05, 1987

AB

BIOCRAFT LABS

250MG/5MLM

N62693 002  
JAN 09, 1987

CHROMIC CHLORIDE IN PLASTIC CONTAINER

AP ABBOTT LABS

EQ 0.004MG CHROMIUM/ML N18961 001  
JUN 26, 1986

CHLORPHENIRAMINE MALEATE

INJECTABLE; INJECTION

CHLOR-TRIMETON

AP @ SCHERING

100MG/ML

N08794 001

CILASTATIN SODIUM; IMIPENEM

INJECTABLE; INJECTION

PRIMAXIN  
MS&D

EQ 250MG BASE/VIAL;  
250MG/VIALM N62756 001  
JAN 08, 1987

CHLORTHALIDONE

TABLET; ORAL

CHLORTHALIDONE

AB COLMED LABS

25MG

N89051 001  
JUN 01, 1987

AB

COLMED LABS

50MG

N89052 001  
JUN 01, 1987

CLINDAMYCIN PHOSPHATE

GEL; TOPICAL

CLEOCIN T

UPJOHN

EQ 1% BASEM N50615 001  
JAN 07, 1987

CHLORTHALIDONE; CLONIDINE HYDROCHLORIDE

TABLET; ORAL

CLONIDINE HCL AND CHLORTHALIDONE

AB MYLAN PHARMS

15MG;0.1MG

N71323 001  
FEB 09, 1987

AB

MYLAN PHARMS

15MG;0.2MG

N71324 001  
FEB 09, 1987

AB

MYLAN PHARMS

15MG;0.3MG

N71325 001  
FEB 09, 1987

INJECTABLE; INJECTION

CLEOCIN

UPJOHN MFG

CLINDAMYCIN PHOSPHATE

AP ABBOTT LABS

EQ 150MG BASE/ML N61839 001

EQ 150MG BASE/MLM N62800 001  
JUL 24, 1987

EQ 150MG BASE/MLM N62801 001  
JUL 24, 1987

COMBIPRES

BOEHR INGEL

15MG;0.1MG

N17503 001

AB

BOEHR INGEL

15MG;0.2MG

N17503 002

AB

BOEHR INGEL

15MG;0.3MG

N17503 003  
APR 10, 1984

CLONIDINE HYDROCHLORIDE

TABLET; ORAL

CLONIDINE HCL

AB BOLAR PHARM

0.1MG# N70395 001  
MAR 23, 1987

AA

PARAFON FORTE DSC  
MCNEIL PHARM

250MG

N88928 001  
MAY 08, 1987

PARAFON FORTE DSC

500MG

N11529 002  
JUN 15, 1987

0.2MG# N70396 001  
MAR 23, 1987

0.3MG# N70397 001  
MAR 23, 1987

>\_ADD\_>  
>\_ADD\_>  
>\_ADD\_>

CLONIDINE HYDROCHLORIDE

TABLET; ORAL

CLONIDINE HCL

MYLAN PHARMS

AB 0.1MGx  
 > ADD > AB N70315 001  
 > ADD > JUN 09, 1987  
 AB 0.2MGx  
 > ADD > AB N70316 001  
 > ADD > JUN 09, 1987  
 AB 0.3MGx  
 > ADD > AB N70317 001  
 > ADD > JUN 09, 1987

CLORAZEPATE DIPOTASSIUM

CAPSULE; ORAL

CLORAZEPATE DIPOTASSIUM

ABLE LABS

> ADD > AB 3.75MGx  
 > ADD > AB N71777 001  
 > ADD > AB JUL 14, 1987  
 > ADD > AB 7.5MGx  
 > ADD > AB N71778 001  
 > ADD > AB JUL 14, 1987  
 > ADD > AB 15MGx  
 > ADD > AB N71779 001  
 > ADD > AB JUL 14, 1987  
 AB AM THERPTCS 3.75MGx  
 JUN 23, 1987 : JAN 08, 1987  
 AB 7.5MGx  
 JUN 23, 1987 : JAN 08, 1987  
 AB 15MGx  
 JUN 23, 1987 : JAN 08, 1987  
 AB 3.75MGx  
 JUN 23, 1987 : MAY 20, 1987  
 AB 7.5MGx  
 JUN 23, 1987 : MAY 20, 1987  
 AB 15MGx  
 JUN 23, 1987 : MAY 20, 1987

TRANXENE

ABBOTT LABS

3.75MG  
 7.5MG  
 15MG

N17105 001  
 N17105 002  
 N17105 003

TABLET; ORAL

CLORAZEPATE DIPOTASSIUM

ABLE LABS

AB 3.75MGx  
 AB 7.5MGx  
 AB 15MGx  
 AB 3.75MGx  
 AB 7.5MGx  
 AB 15MGx

N71780 001  
 JUN 26, 1987  
 N71781 001  
 JUN 26, 1987  
 N71782 001  
 JUN 26, 1987  
 N71747 001  
 JUN 23, 1987 : JUN 09, 1987  
 N71748 001  
 JUN 23, 1987 : JUN 09, 1987  
 N71749 001  
 JUN 23, 1987 : JUN 09, 1987

AM THERPTCS

CLORAZEPATE DIPOTASSIUM

TABLET; ORAL

CLORAZEPATE DIPOTASSIUM

MYLAN PHARMS

> ADD > AB 3.75MGx  
 > ADD > AB N71856 001  
 > ADD > AB JUL 17, 1987  
 > ADD > AB 7.5MGx  
 > ADD > AB N71857 001  
 > ADD > AB JUL 17, 1987  
 > ADD > AB 15MGx  
 > ADD > AB N71858 001  
 > ADD > AB JUL 17, 1987

TRAMXENE

ABBOTT LABS

3.75MG  
 7.5MG  
 15MG  
 N17105 006  
 N17105 007  
 N17105 008

CODEINE PHOSPHATE; PHENYLEPHRINE HYDROCHLORIDE; PROMETHAZINE HYDROCHLORIDE

SYRUP; ORAL

PHERAZINE VC W/ CODEINE

HALSEY DRUG

AA 10MG/5ML; 5MG/5ML;  
 6.25MG/5MLx  
 N88870 001  
 MAR 02, 1987

CUPRIC SULFATE

INJECTABLE; INJECTION

CUPRIC SULFATE

LYPHOMED

EQ 0.4MG COPPER/MLx  
 N19350 001  
 MAY 05, 1987

CYCLOPENTOLATE HYDROCHLORIDE

SOLUTION/DROPS; OPHTHALMIC

CYCLOGYL

ALCON LABS

0.5%

PENTOLAIR

PHARMAFAIR

0.5%  
 N84109 001

DESIPRAMINE HYDROCHLORIDE

TABLET; ORAL

DESIPRAMINE HCL

VITARINE

AB 25MGx  
 AB 50MGx  
 N71601 001  
 JUN 05, 1987  
 N71588 001  
 JUN 05, 1987

DESIPRAMINE HYDROCHLORIDE

TABLET; ORAL

MORPRAMIN  
MERRILL DOW

AB 25MG  
AB 50MG

N14399 001  
N14399 003

DEXAMETHASONE SODIUM PHOSPHATE

INJECTABLE; INJECTION

DEXAMETHASONE SODIUM PHOSPHATE

AP EQ 4MG PHOSPHATE/ML

AP EQ 10MG PHOSPHATE/ML

AP EQ 20MG PHOSPHATE/ML

AP EQ 24MG PHOSPHATE/ML

N89280 001  
MAR 18, 1987  
N89281 001  
MAR 18, 1987  
N89282 001  
MAR 18, 1987  
N89372 001  
MAR 18, 1987

DEXTROMETHORPHAN HYDROBROMIDE; PROMETHAZINE HYDROCHLORIDE

SYRUP; ORAL

PHERAZINE DM  
HALSEY DRUG

AA 15MG/5ML ; 6.25MG/5ML

N88913 001  
MAR 02, 1987

DIAZEPAM

CONCENTRATE; ORAL  
DIAZEPAM INTENSOL  
ROXANE LABS

5MG/ML

N71415 001  
APR 03, 1987

INJECTABLE; INJECTION

DIAZEPAM

LEDERLE LABS

5MG/ML

N71308 001  
JUL 17, 1987

5MG/ML

N71309 001  
JUL 17, 1987

5MG/ML

N71310 001  
JUL 17, 1987

SOLUTION; ORAL

DIAZEPAM  
ROXANE LABS

5MG/5ML

N70928 001  
APR 03, 1987

DIAZEPAM

TABLET; ORAL

DIAZEPAM  
COLMED LABS

AB 2MG

N70903 001  
APR 01, 1987

AB 5MG

N70904 001  
APR 01, 1987

AB 10MG

N70905 001  
APR 01, 1987

AB 2MG

N71134 001  
FEB 03, 1987

AB 5MG

N71135 001  
FEB 03, 1987

AB 10MG

N71136 001  
FEB 03, 1987

DICYCLOMINE HYDROCHLORIDE

CAPSULE; ORAL

DICYCLOMINE HCL

AB 10MG

N84505 001  
OCT 21, 1986

DIPHENHYDRAMINE HYDROCHLORIDE

CAPSULE; ORAL

DIPHENHYDRAMINE HCL

AA 25MG

N89488 001  
JAN 02, 1987

AA 50MG

N89489 001  
JAN 02, 1987

DIPYRIDAMOLE

TABLET; ORAL

PERSANTINE  
BOEHR INGEL

50MG

N12836 004  
FEB 06, 1987

75MG

N12836 005  
FEB 06, 1987

DISOPYRAMIDE PHOSPHATE

CAPSULE; ORAL  
DISOPYRAMIDE PHOSPHATE  
 INTERPHARM

AB EQ 100MG BASEM N71190 001  
 JAN 15, 1987  
 AB EQ 150MG BASEM N71191 001  
 JAN 15, 1987  
 AB EQ 100MG BASEM N70940 001  
 FEB 09, 1987  
 AB EQ 150MG BASEM N70941 001  
 FEB 09, 1987

CAPSULE; ORAL  
DOXEPTIN HCL  
 DANBURY PHARMA

AB EQ 10MG BASEM N71485 001  
 APR 30, 1987  
 AB EQ 25MG BASEM N71486 001  
 APR 30, 1987  
 AB EQ 50MG BASEM N71238 001  
 APR 30, 1987  
 AB EQ 75MG BASEM N71326 001  
 APR 30, 1987  
 AB EQ 100MG BASEM N71239 001  
 APR 30, 1987

DOPAMINE HYDROCHLORIDE

INJECTABLE; INJECTION  
DOPAMINE HCL

AP LUITPOLD PHARMS 40MG/MLM N70799 001  
 FEB 11, 1987  
 AP 80MG/MLM N70820 001  
 FEB 11, 1987  
 AP 160MG/MLM N70826 001  
 FEB 11, 1987

DOXORUBICIN HYDROCHLORIDE

INJECTABLE; INJECTION  
ADRIAMYCIN  
FARMITALIA

AB 10MG/VIAL /16MG/VIAL/ N50467 001  
 20MG/VIAL /20MG/VIAL/ N50467 003  
 MAY 20, 1985  
 AB 50MG/VIAL N50467 002  
 N50467 004  
 JUL 22, 1987  
 AB 150MG/VIALM N50467 004  
 JUL 22, 1987

DOPAMINE HCL IN DEXTROSE 5% IN PLASTIC CONTAINER

AP TRAVENOL LABS 80MG/100MLM N19615 001  
 MAR 27, 1987  
 AP 160MG/100MLM N19615 002  
 MAR 27, 1987  
 AP 320MG/100MLM N19615 003  
 MAR 27, 1987  
 AP 640MG/100MLM N19615 004  
 MAR 27, 1987

DOXEPTIN HYDROCHLORIDE

CAPSULE; ORAL  
DOXEPTIN HCL

AB CHELSEA LABS EQ 10MG BASEM N70952 001  
 MAR 04, 1987  
 AB CORD LABS EQ 10MG BASEM N71487 001  
 MAR 02, 1987  
 AB EQ 100MG BASEM N71562 001  
 MAR 02, 1987

ENFLURANE

LIQUID; INHALATION  
ENFLURANE

>\_ADD\_> AN ABBOTT LABS 99.9% SEP 08, 1987 : JUL 27, 1987 N70803 001  
 >\_ADD\_> AN ABBOTT LABS 99.9%  
 >\_ADD\_> AN ANAQUEST 99.9% N17087 001

EPINEPHRINE; LIDOCAINE HYDROCHLORIDE

INJECTABLE; INJECTION  
XYLOCAINE W/ EPINEPHRINE  
 ASTRA PHARM PRODS

>\_ADD\_> 0.005MG/ML;1% N06488 018  
 >\_ADD\_> 0.005MG/ML;2% N06488 019  
 NOV 13, 1986  
 NOV 13, 1986

RX DRUG PRODUCT LIST / CUMULATIVE SUPPLEMENT NUMBER 7 / JAN'87 - JUL'87

ERYTHROMYCIN

SWAB; TOPICAL

ERYCETTE

ORTHO PHARM

2/2

N50594 001  
FEB 15, 1985

0.035MG; 0.5MG

N70686 001  
JAN 29, 1987

T-STAT

WESTWOOD PHARMS

2/2

N62748 001  
JUL 23, 1987

0.035MG; 1MG

N70687 001  
JAN 29, 1987

ERYTHROMYCIN ETHYLSUCCINATE

SUSPENSION; ORAL

ERYTHROMYCIN ETHYLSUCCINATE

NASKA PHARMA

EQ 400MG BASE/5ML

N62674 001  
MAR 10, 1987

N19545 001  
APR 20, 1987

ESTRADIOL CYPIONATE

INJECTABLE; INJECTION

ESTRADIOL CYPIONATE

QUAD PHARMS

5MG/ML

N89310 001  
FEB 09, 1987

POWDER FOR RECONSTITUTION; ORAL

PEPCID

MS&D RES LABS

40MG/5ML

N19527 001  
FEB 02, 1987

ESTROGENS, CONJUGATED

TABLET; ORAL

CONJUGATED ESTROGENS

CHELSEA LABS

0.625MG

1.25MG

2.5MG

0.625MG

1.25MG

2.5MG

0.625MG

1.25MG

2.5MG

N85800 001  
N85801 001  
N85826 001  
N83356 001  
N83360 001  
N84650 001  
N83354 003  
N83592 001  
N85908 001

200MG

N18830 002  
OCT 31, 1985

ETHINYL ESTRADIOL; NORETHINDRONE

TABLET; ORAL-21

GYNEX 0.5/35E-21

GYNEX LABS

0.035MG; 0.5MG

N70684 001  
JAN 29, 1987

GYNEX 1/35E-21

GYNEX LABS

0.035MG; 1MG

N70685 001  
JAN 29, 1987

ETHINYL ESTRADIOL; NORETHINDRONE

TABLET; ORAL-28

GYNEX 0.5/35E-28

GYNEX LABS

0.035MG; 0.5MG

N70686 001  
JAN 29, 1987

GYNEX 1/35E-28

GYNEX LABS

0.035MG; 1MG

N70687 001  
JAN 29, 1987

ETIDRONATE DISODIUM

INJECTABLE; INJECTION

DIDRONEL

NORWICH EATON

50MG/ML

N19545 001  
APR 20, 1987

FAMOTIDINE

POWDER FOR RECONSTITUTION; ORAL

PEPCID

MS&D RES LABS

40MG/5ML

N19527 001  
FEB 02, 1987

FLECAINIDE ACETATE

TABLET; ORAL

TAMBOCOR

RIKER LABS

200MG

N18830 002  
OCT 31, 1985

FLUNISOLIDE

AEROSOL, METERED; INHALATION

AEROBID

KEY PHARMS

0.25MG/INH

N18340 001  
AUG 17, 1984

FLUOCINONIDE

CREAM; TOPICAL

FLUOCINONIDE

THAMES PHARMA

0.05%

N71500 001  
JUN 10, 1987

FLUOROMETHOLONE ACETATE

SUSPENSION/DROPS; OPHTHALMIC  
FLAREX

ALCON LABS 0.1%  
/MINTHROP/  
/ALCON/LABS/  
/6.1%/  
/N19079/001/  
/FEB/11, 1986/  
/N19079/001/  
/FEB/11, 1986/

N19079 001  
FEB 11, 1986

AP  
INJECTABLE; INJECTION  
FUROSEMIDE  
CARTER GLOGAU  
WINTHROP BREON

10MG/MLM  
10MG/MLM

N70604 001  
JAN 02, 1987  
N70578 001  
JUL 08, 1987

FLUOROURACIL

INJECTABLE; INJECTION  
FLUOROURACIL  
LYPHOMED

AP 50MG/MLM  
AP 50MG/MLM  
AP 50MG/MLM  
AP 50MG/MLM  
AP 50MG/MLM

N89428 001  
JAN 12, 1987  
N89519 001  
MAR 12, 1987  
N89368 001  
FEB 03, 1987  
N89455 001  
FEB 03, 1987  
N89434 001  
MAR 26, 1987

AA  
SOLUTION; ORAL  
FUROSEMIDE  
ROXANE LABS  
LASIX  
HOECHST  
TABLET; ORAL  
FUROSEMIDE  
WATSON LABS

10MG/MLM  
40MG/5MLM  
10MG/ML

N70434 001  
APR 22, 1987  
N70433 001  
APR 22, 1987  
N17688 001

FLUPHENAZINE DECANOATE

INJECTABLE; INJECTION  
FLUPHENAZINE DECANOATE  
LYPHOMED

> ADD >  
> ADD > A0  
> ADD >

N71413 001  
JUL 14, 1987

AT  
SOLUTION/DROPS; OPHTHALMIC  
GENTAMICIN SULFATE  
MAURRY BIO

EQ 3MG BASE/MLM

N62635 001  
JAN 08, 1987

FLUPHENAZINE HYDROCHLORIDE

INJECTABLE; INJECTION  
FLUPHENAZINE HCL  
LYPHOMED

AP 2.5MG/MLM  
AP 2.5MG/ML

N89556 001  
APR 16, 1987

AP  
INJECTABLE; INJECTION  
GLUCAGON  
LILLY  
QUAD PHARMS

EQ 1MG BASE/VIAL  
EQ 10MG BASE/VIAL  
EQ 1MG BASE/VIALM

N12122 001  
N12122 002  
N71022 001  
MAR 04, 1987  
N71023 001  
MAR 04, 1987

FLURAZEPAM HYDROCHLORIDE

CAPSULE; ORAL  
FLURAZEPAM HCL  
COLMED LABS

> ADD > AB  
> ADD >  
> ADD > AB  
> ADD >

N70562 001  
JUL 09, 1987  
N70563 001  
JUL 09, 1987

AT  
GRAMICIDIN; NEOMYCIN SULFATE; POLYMYXIN B SULFATE  
SOLUTION/DROPS; OPHTHALMIC  
NEOMYCIN AND POLYMYXIN B SULFATES AND GRAMICIDIN  
STERIS LABS

0.025MG/ML; EQ 1.75MG BASE/ML;  
10,000 UNITS/MLM

N62788 001  
JUN 11, 1987

HALOPERIDOL

TABLET; ORAL  
HALOPERIDOL  
BARR LABS

AB	> ADD >	0.5MG	N71156 001	AP	EQ 5MG BASE/MLM	N71187 001
AB	> ADD >	1MG	JAN 02, 1987	LYPHOMED		JAN 20, 1987
AB	> ADD >	2MG	N71157 001	QUAD PHARMS	EQ 5MG BASE/MLM	N71082 001
AB	> ADD >	2MG	JAN 02, 1987			JAN 02, 1987
AB	> ADD >	0.5MG	N71172 001			
AB	> ADD >	1MG	JAN 02, 1987			
AB	> ADD >	2MG	N70981 001			
AB	> ADD >	5MG	MAR 06, 1987			
AB	> ADD >	10MG	N70982 001			
AB	> ADD >	20MG	MAR 06, 1987			
AB	> ADD >	5MG	N70983 001			
AB	> ADD >	10MG	MAR 06, 1987			
AB	> ADD >	20MG	N70984 001			
AB	> ADD >	0.5MG	MAR 06, 1987			
AB	> ADD >	1MG	N71220 001			
AB	> ADD >	2MG	JUL 07, 1987			
AB	> ADD >	5MG	N71221 001			
AB	> ADD >	10MG	JUL 07, 1987			
AB	> ADD >	20MG	N71237 001			
AB	> ADD >	0.5MG	JUL 20, 1987			
AB	> ADD >	1MG	N71255 001			
AB	> ADD >	2MG	FEB 17, 1987			
AB	> ADD >	5MG	N71269 001			
AB	> ADD >	0.5MG	FEB 17, 1987			
AB	> ADD >	1MG	N71256 001			
AB	> ADD >	2MG	FEB 17, 1987			
AB	> ADD >	5MG	N71257 001			
AB	> ADD >	0.5MG	FEB 17, 1987			
AB	> ADD >	1MG	N71128 001			
AB	> ADD >	2MG	FEB 17, 1987			
AB	> ADD >	5MG	N71129 001			
AB	> ADD >	0.5MG	FEB 17, 1987			
AB	> ADD >	1MG	N71130 001			
AB	> ADD >	2MG	FEB 17, 1987			
AB	> ADD >	5MG	N71131 001			
AB	> ADD >	10MG	FEB 17, 1987			
AB	> ADD >	20MG	N71132 001			
AB	> ADD >	0.5MG	MAY 12, 1987			
AB	> ADD >	1MG	N71133 001			
AB	> ADD >	2MG	MAY 12, 1987			

HALOPERIDOL LACTATE

INJECTABLE; INJECTION

HALOPERIDOL

AP	> ADD >	EQ 5MG BASE/MLM				N71187 001
AP	> ADD >	EQ 5MG BASE/MLM				JAN 20, 1987
AP	> ADD >	EQ 5MG BASE/MLM				N71082 001
AP	> ADD >	EQ 5MG BASE/MLM				JAN 02, 1987
AP	> ADD >	10,000 UNITS/MLM				
AP	> ADD >	10,000 UNITS/MLM				
AP	> ADD >	2,000 UNITS/100ML				
AP	> ADD >	5,000 UNITS/100ML				
AP	> ADD >	10,000 UNITS/100MLM				

HEPARIN SODIUM

INJECTABLE; INJECTION

HEPARIN SODIUM PRESERVATIVE FREE

WINTHROP BREON

AP	> ADD >	10,000 UNITS/MLM				N89522 001
AP	> ADD >	10,000 UNITS/MLM				MAY 04, 1987

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

TRAVENOL LABS

AP	> ADD >	2,000 UNITS/100ML				N18814 002
AP	> ADD >	2,000 UNITS/100ML				JUL 09, 1985

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

TRAVENOL LABS

AP	> ADD >	5,000 UNITS/100ML				N18814 003
AP	> ADD >	5,000 UNITS/100ML				JUL 09, 1985
AP	> ADD >	10,000 UNITS/100MLM				N18814 004
AP	> ADD >	10,000 UNITS/100MLM				JUL 02, 1987

HEXACHLOROPHENE

EMULSION; TOPICAL

SOY-DOME

3 MILES PHARMS

AT	> ADD >	3%				N17405 001
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HYDRALAZINE HYDROCHLORIDE; HYDROCHLOROTHIAZIDE

CAPSULE; ORAL

HYDRALAZINE HCL AND HYDROCHLOROTHIAZIDE

SUPERPHARM

AB	> ADD >	25MG; 25MG				N89200 001
AB	> ADD >	50MG; 50MG				FEB 09, 1987
AB	> ADD >	50MG; 50MG				N89201 001
AB	> ADD >	50MG; 50MG				FEB 09, 1987

HALOPERIDOL LACTATE

INJECTABLE; INJECTION

HALDOL

MCNEIL LABS

AP	> ADD >	EQ 5MG BASE/ML				N15923 001
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HYDROCHLOROTHIAZIDE; LABETALOL HYDROCHLORIDE

TABLET; ORAL  
NORMOZIDE  
SCHERING

AB 2.5MG; 100MG  
AB 2.5MG; 200MG  
AB 2.5MG; 300MG  
AB 2.5MG; 400MG  
AB 2.5MG; 100MG  
AB 2.5MG; 200MG  
AB 2.5MG; 300MG  
AB 2.5MG; 400MG

N19046 001  
APR 06, 1987  
N19046 002  
APR 06, 1987  
N19046 003  
APR 06, 1987  
N19046 004  
APR 06, 1987  
N19174 001  
APR 10, 1987  
N19174 002  
APR 10, 1987  
N19174 003  
APR 10, 1987  
N19174 004  
APR 10, 1987

TABLET; ORAL

PROPRANOLOL HCL & HYDROCHLOROTHIAZIDE  
DURAMED PHARMS

AB 2.5MG; 40MG  
AB 2.5MG; 80MG

N71126 001  
MAR 02, 1987  
N71127 001  
MAR 02, 1987

PROPRANOLOL HCL AND HYDROCHLOROTHIAZIDE

AB 2.5MG; 40MG  
AB 2.5MG; 80MG

N70946 001  
MAR 04, 1987  
N70947 001  
APR 01, 1987

TRANDATE-HCT  
GLAXO

AB 2.5MG; 100MG  
AB 2.5MG; 200MG  
AB 2.5MG; 300MG  
AB 2.5MG; 400MG

N19174 001  
APR 10, 1987  
N19174 002  
APR 10, 1987  
N19174 003  
APR 10, 1987  
N19174 004  
APR 10, 1987

HYDROCHLOROTHIAZIDE; SPIRONOLACTONE

TABLET; ORAL

SPIRONOLACTONE AND HYDROCHLOROTHIAZIDE  
MUTUAL PHARM

> ADD > 2.5MG; 2.5MG  
> ADD > 2.5MG; 2.5MG

N89534 001  
JUL 02, 1987

HYDROCHLOROTHIAZIDE; METHYLDOPA

TABLET; ORAL  
METHYLDOPA AND HYDROCHLOROTHIAZIDE  
INVAMED

AB 1.5MG; 2.5MG  
AB 2.5MG; 2.5MG  
AB 1.5MG; 2.5MG  
AB 2.5MG; 2.5MG  
AB 30MG; 500MG  
AB 50MG; 500MG

N70829 001  
MAR 09, 1987  
N70830 001  
MAR 09, 1987  
N70616 001  
FEB 02, 1987  
N70612 001  
FEB 02, 1987  
N70613 001  
FEB 02, 1987  
N70614 001  
FEB 02, 1987

AB 1.5MG  
OINTMENT; TOPICAL  
HYDROCORTISONE  
PHARMADERM

N88842 001  
FEB 09, 1987

HYDROCORTISONE BUTYRATE

SOLUTION; TOPICAL  
LOCOID  
GIST BROCADES

0.1%  
0.1%

N19116 001  
FEB 25, 1987

HYDROCORTISONE SODIUM PHOSPHATE

INJECTABLE; INJECTION  
HYDROCORTISONE SODIUM PHOSPHATE  
QUAD PHARMS

AP EQ 50MG BASE/ML

N89581 001  
MAY 28, 1987

HYDROCORTONE

AP MS&D  
EQ 50MG BASE/ML

N12052 001

> ADD > HYDROCHLOROTHIAZIDE; PINDOLOL

TABLET; ORAL  
VISKAZIDE  
SANDOZ PHARMS

25MG; 5MG  
25MG; 10MG

N18872 001  
JUL 22, 1987  
N18872 002  
JUL 22, 1987

> ADD >  
> ADD >



IOPAMIDOL

INJECTABLE; INJECTION

ISOVUE-200  
 SQUIBB DIAGS  
 41%  
 N18735 001  
 DEC 31, 1985  
 > ADD > AB  
 > ADD >  
 /ISOVUE-N/ADD/  
 SQUIBB/DIAGS/  
 /41%/  
 N18735/ADD/  
 DEC/31/1985/  
 26%  
 N18735 005  
 OCT 21, 1986  
 > ADD > AB  
 > ADD >

ISOVUE-128  
 SQUIBB DIAGS

N18754 001  
 JUL 31, 1987  
 25MG

IRON DEXTRAN COMPLEX

INJECTABLE; INJECTION  
 INFERON  
 FISON  
 /MERRELL/DOM/  
 EQ 50MG IRON/ML  
 /EQ 50MG IRON/ML/  
 N10787 002  
 N10787/ADD/  
 EQ 50MG BASE/VIAL  
 EQ 50MG BASE/VIAL  
 EQ 50MG BASE/VIAL

N70480 001  
 JAN 02, 1987  
 N89496 001  
 MAR 05, 1987

INJECTABLE; INJECTION

POWDER FOR RECONSTITUTION; ORAL  
 LEUCOVORIN CALCIUM  
 LEDERLE LABS  
 N10787 002  
 N10787/ADD/  
 EQ 60MG BASE/VIAL

N08107 003  
 JAN 30, 1987

ISOSORBIDE DINITRATE

TABLET; ORAL  
 BARR LABS  
 5MG  
 N86166 002  
 SEP 19, 1986  
 10MG  
 N86169 001  
 SEP 19, 1986  
 20MG  
 N86167 001  
 SEP 19, 1986  
 5MG  
 N89190 001  
 FEB 17, 1987  
 10MG  
 N89191 001  
 FEB 17, 1987  
 20MG  
 N89192 001  
 FEB 17, 1987

N71104 001  
 MAR 04, 1987

TABLET; ORAL

LEUCOVORIN CALCIUM  
 LEDERLE LABS  
 EQ 15MG BASE  
 LITHIUM CARBONATE  
 CAPSULE; ORAL  
 LITHIUM CARBONATE  
 BOLAR PHARM  
 300MG  
 150MG  
 600MG  
 ROXANE LABS

N70407 001  
 MAR 19, 1987  
 N17812 002  
 JAN 28, 1987  
 N17812 003  
 JAN 28, 1987

KANAMYCIN SULFATE

CAPSULE; ORAL  
 KANTREX  
 BRISTOL LABS  
 EQ 500MG BASE  
 N62726 001  
 MAR 06, 1987  
 INJECTABLE; INJECTION  
 KANAMYCIN SULFATE  
 PHARMAFAIR  
 AP  
 EQ 75MG BASE/2ML  
 N62668 001  
 MAY 07, 1987  
 AP  
 EQ 500MG BASE/2ML  
 N62672 001  
 MAY 07, 1987  
 AP  
 EQ 1GM BASE/3ML  
 N62669 001  
 MAY 07, 1987

N71403 001  
 APR 21, 1987  
 N71404 001  
 APR 21, 1987  
 N71141 001  
 APR 21, 1987

TABLET; ORAL

LORAZEPAM  
 PUREPAC PHARM  
 0.5MG  
 1MG  
 2MG

RX DRUG PRODUCT LIST / CUMULATIVE SUPPLEMENT NUMBER 7 / JAN '87 - JUL '87

LORAZEPAM

TABLET; ORAL  
LORAZEPAM  
SUPERPHARM

AB 0.5MG~~M~~  
AB 1MG~~M~~  
AB 2MG~~M~~  
AB 0.5MG~~M~~  
AB 1MG~~M~~  
AB 2MG~~M~~

> ADD >  
> ADD > AB  
> ADD >  
> ADD > AB  
> ADD >  
N71245 001  
FEB 09, 1987  
N71246 001  
FEB 09, 1987  
N71247 001  
FEB 09, 1987  
N71086 001  
MAR 23, 1987  
N71087 001  
MAR 23, 1987  
N71088 001  
MAR 23, 1987

CAPSULE; ORAL  
MECLODITUM

EQ 50MG BASE~~M~~  
EQ 100MG BASE~~M~~  
EQ 50MG BASE~~M~~  
EQ 100MG BASE~~M~~  
EQ 50MG BASE~~M~~  
EQ 100MG BASE~~M~~

N71380 001  
JUL 14, 1987  
N71381 001  
JUL 14, 1987  
N71362 001  
FEB 10, 1987  
N71363 001  
FEB 10, 1987  
N71468 001  
APR 15, 1987  
N71469 001  
APR 15, 1987

MECLOFENAMATE SODIUM  
AM THERPTCS

AB  
AB  
AB  
AB

DANBURY PHARMA

MANGANESE SULFATE

INJECTABLE; INJECTION  
MANGANESE SULFATE  
LYPHOMED

EQ 0.1MG MANGANESE/ML~~M~~

N19228 001  
MAY 05, 1987

METHOCARBAMOL

TABLET; ORAL  
METHOCARBAMOL  
AM THERPTCS

AA 500MG~~M~~  
AA 750MG~~M~~

N89417 001  
FEB 11, 1987  
N89418 001  
FEB 11, 1987

MANNITOL

INJECTABLE; INJECTION  
MANNITOL 10% IN PLASTIC CONTAINER  
ABBOTT LABS

AP 10GM/100ML~~M~~

AP 12.5GM/50ML~~M~~  
AP 12.5GM/50ML~~M~~

AP 5GM/100ML~~M~~

N19603 002  
JAN 08, 1987  
N89239 001  
MAY 06, 1987  
N89240 001  
MAY 06, 1987  
N19603 001  
JAN 08, 1987

METHOTREXATE SODIUM

INJECTABLE; INJECTION  
ABITREXATE  
INTL PHARM

AP  
> ADD > AP  
> ADD >  
> ADD > AP  
> ADD >  
> ADD > AP  
> ADD >  
EQ 25MG BASE/ML~~M~~  
EQ 50MG BASE/VIAL~~M~~  
EQ 100MG BASE/VIAL~~M~~  
EQ 250MG BASE/VIAL~~M~~

N89161 001  
MAR 10, 1987  
N89354 001  
JUL 17, 1987  
N89355 001  
JUL 17, 1987  
N89356 001  
JUL 17, 1987

MECLIZINE HYDROCHLORIDE

TABLET; ORAL  
ANTIVERT  
ROERIG

50MG

N10721 001  
JAN 20, 1982

METHOXSALEN

CAPSULE; ORAL  
METHOXSALEN  
BP @ CORD LABS

10MG

N87781 001  
JUN 08, 1982

METHYLDOPA

TABLET; ORAL  
METHYLDOPA  
PAR PHARM

AB 125MG~~M~~ N70535 001  
JAN 02, 1987  
AB 250MG~~M~~ N70536 001  
JAN 02, 1987  
AB 500MG~~M~~ N70537 001  
JAN 02, 1987

METHYLDOPATE HYDROCHLORIDE

INJECTABLE; INJECTION  
METHYLDOPATE HCL  
ABBOTT LABS

AP 50MG/ML~~M~~ N70698 001  
JUN 15, 1987  
AP 50MG/ML~~M~~ N70699 001  
JUN 15, 1987  
AP 50MG/ML~~M~~ N70691 001  
JUN 19, 1987  
AP 50MG/ML~~M~~ N70849 001  
JUN 19, 1987  
AP 50MG/ML~~M~~ N70841 001  
JAN 02, 1987

METOCLOPRAMIDE HYDROCHLORIDE

INJECTABLE; INJECTION  
METOCLOPRAMIDE HCL  
SOLOPAK LABS

AP EQ 10MG BASE/2ML~~M~~ N70622 001  
MAR 02, 1987  
AP EQ 10MG BASE/2ML~~M~~ N70623 001  
MAR 02, 1987  
REGLAN  
ROBINS  
EQ 10MG BASE/ML~~M~~ N17862 004  
MAY 28, 1987

SYRUP; ORAL  
METOCLOPRAMIDE HCL  
BIOCRAFT LABS

>\_ADD\_> AA EQ 5MG BASE/5ML~~M~~ N70819 001  
>\_ADD\_> AA EQ 5MG BASE/5ML~~M~~ N70949 001  
MAR 06, 1987  
REGLAN  
ROBINS  
EQ 5MG BASE/5ML~~M~~ N18821 001  
MAR 25, 1983

TABLET; ORAL  
METOCLOPRAMIDE HCL  
BARR LABS

AB EQ 10MG BASE~~M~~ N70660 001  
FEB 10, 1987

METOCLOPRAMIDE HYDROCHLORIDE

TABLET; ORAL  
METOCLOPRAMIDE HCL  
BOLAR PHARM

AB EQ 10MG BASE~~M~~ N70363 001  
MAR 02, 1987  
AB EQ 10MG BASE~~M~~ N70850 001  
FEB 03, 1987  
AB EQ 10MG BASE~~M~~ N70598 001  
FEB 02, 1987  
AB EQ 10MG BASE~~M~~ N70926 001  
JUN 26, 1987  
AB EQ 10MG BASE~~M~~ N70645 001  
MAY 11, 1987  
REGLAN  
ROBINS  
EQ 5MG BASE~~M~~ N17854 002  
MAY 05, 1987

METRIZAMIDE

INJECTABLE; INJECTION  
AMIPAQUE  
MINTHROP BREON

EQ 2.5GM/VIAL N17982 003  
SEP 12, 1983  
EQ 13.5GM/VIAL N17982 004  
SEP 12, 1983

METRONIDAZOLE

TABLET; ORAL  
SATRIG  
SAVAGE LABS

AB 500MG~~M~~ N70731 001  
JUN 08, 1987

MEZLOCILLIN SODIUM MONOHYDRATE

INJECTABLE; INJECTION  
MEZLIN  
MILES PHARMS

EQ 3GM BASE/VIAL~~M~~ N62697 001  
JAN 22, 1987  
EQ 4GM BASE/VIAL~~M~~ N62697 002  
JAN 22, 1987

MIDAZOLAM HYDROCHLORIDE

INJECTABLE; INJECTION  
VERSED  
ROCHE

EQ 1MG BASE/ML~~M~~ N18654 002  
MAY 26, 1987

MINOXIDIL

TABLET; ORAL  
LOXTEH  
 UPJOHN  
 AB N18154 001 2.5MG 25MG/ML MAR 23, 1987  
 AB N18154 003 10MG

MEHODYL  
 QUANTUM PHARMCS  
 AB N71534 001 10MG MAR 19, 1987

MEHOXIDIL  
 DANBURY PHARMA  
 AB N71344 001 2.5MG MAR 03, 1987  
 AB N71345 001 10MG MAR 03, 1987

MOMETASONE FUROATE

CREAM; TOPICAL  
 ELOCON  
 SCHERING  
 AB N19625 001 0.1% MAY 06, 1987

OINTMENT; TOPICAL  
 ELOCON  
 SCHERING  
 AB N19543 001 0.1% APR 30, 1987

MORPHINE SULFATE

TABLET, CONTROLLED RELEASE; ORAL  
 MS CONTIN  
 PURDUE FRDRK  
 AB N19516 001 30MG MAY 29, 1987

NALOXONE HYDROCHLORIDE

INJECTABLE; INJECTION  
NALOXONE HCL  
 ABBOTT LABS  
 AB N70252 001 0.02MG/ML JAN 16, 1987  
 AB N70253 001 0.02MG/ML JAN 16, 1987  
 AB N70254 001 0.4MG/ML JAN 07, 1987  
 AB N70255 001 0.4MG/ML JAN 07, 1987  
 AB N70256 001 0.4MG/ML JAN 07, 1987  
 AB N70257 001 0.4MG/ML JAN 07, 1987

NAPROXEN

SUSPENSION; ORAL  
 NAPROSYN  
 SYNTEX LABS  
 AB N18965 001 25MG/ML MAR 23, 1987

NITROGLYCERIN

INJECTABLE; INJECTION  
NITROGLYCERIN  
 LYPHOMED  
 AP N71203 001 5MG/ML MAY 08, 1987  
 AP N71094 001 5MG/ML JUL 31, 1987  
 AP N71095 001 10MG/ML JUL 31, 1987

NITROSTAT

PARKE DAVIS  
 AP N70863 001 5MG/ML JAN 08, 1987  
 AP N70871 001 10MG/ML JAN 08, 1987  
 AP N70872 001 10MG/ML JAN 08, 1987

NYSTATIN

PASTILLE; ORAL  
 MYCOSTATIN  
 SQUIBB  
 AB N50619 001 200,000 UNITS APR 09, 1987

SUSPENSION; ORAL  
NYSTATIN  
 AA BIOCRAFT LABS  
 AB N62670 001 100,000 UNITS/ML JUN 18, 1987

NYSTATIN; TRIAMCINOLONE ACETONIDE

CREAM; TOPICAL  
NYSTATIN-TRIAMCINOLONE ACETONIDE  
 AT THAMES PHARMA  
 AB N62347 001 100,000 UNITS/GH; 0.1% MAR 30, 1987

OINTMENT; TOPICAL  
MYKACET  
 AT NMC LABS  
 AB N62733 001 100,000 UNITS/GH; 0.1% MAR 09, 1987



PROCHLORPERAZINE MALEATE

TABLET; ORAL

PROCHLORPERAZINE MALEATE

AB DURAMED PHARMS EQ 5MG BASEM N89484 001  
 > ADD > AB JAN 20, 1987  
 > ADD > AB N89485 001  
 > ADD > AB JAN 20, 1987  
 > ADD > AB N89486 001  
 > ADD > AB JAN 20, 1987

PROPRANOLOL HYDROCHLORIDE

TABLET; ORAL

PROPRANOLOL HCL

60MGM N71791 001  
 JUL 15, 1987  
 90MGM N71792 001  
 JUL 15, 1987

PROPRANOLOL HYDROCHLORIDE

CAPSULE, CONTROLLED RELEASE; ORAL  
 INDERAL LA  
 AYERST LABS 60MGM N18553 004  
 MAR 18, 1987

CONCENTRATE; ORAL  
 PROPRANOLOL HCL INTENSOL  
 ROXANE LABS 80MG/MLM N71388 001  
 MAY 15, 1987

SOLUTION; ORAL  
 PROPRANOLOL HCL  
 ROXANE LABS 20MG/5MLM N70979 001  
 MAY 15, 1987  
 40MG/5MLM N70690 001  
 MAY 15, 1987

TABLET; ORAL  
PROPRANOLOL HCL  
 BOLAR PHARM 10MGM N70378 001  
 MAR 19, 1997  
 20MGM N70379 001  
 MAR 19, 1987  
 40MGM N70380 001  
 MAR 19, 1987  
 60MGM N70381 001  
 MAR 19, 1987  
 80MGM N70382 001  
 MAR 19, 1987  
 60MGM N70143 001  
 JAN 15, 1987  
 10MGM N71368 001  
 MAY 05, 1987  
 20MGM N71369 001  
 MAY 05, 1987  
 40MGM N71370 001  
 MAY 05, 1987  
 80MGM N71371 001  
 MAY 05, 1987

INJECTABLE; INJECTION  
PROTAMINE SULFATE

AP LYPHOMED 10MG/MLM N89454 001  
 APR 07, 1987

QUAZEPAM

TABLET; ORAL  
 DORMALIN  
 SCHERING 7.5MGM N18708 003  
 FEB 26, 1987

QUINIDINE GLUCONATE

TABLET, CONTROLLED RELEASE; ORAL  
QUINIDINE GLUCONATE

AB HALSEY DRUG 324MGM N89476 001  
 APR 10, 1987  
 AB MUTUAL PHARM 324MGM N89338 001  
 FEB 11, 1987

RITODRINE HYDROCHLORIDE

INJECTABLE; INJECTION  
RITODRINE HCL

AP LYPHOMED 10MG/MLM N71188 001  
 JUL 23, 1987  
 AP LYPHOMED 15MG/MLM N71189 001  
 JUL 23, 1987

SODIUM CHLORIDE

INJECTABLE; INJECTION  
 SODIUM CHLORIDE 23.4% IN PLASTIC CONTAINER  
 LYPHOMED 234MG/MLM N19329 001  
 APR 22, 1987

SOMATROPIN, BIOSYNTHETIC

INJECTABLE; INJECTION  
HUMATROPE  
LILLY

5MG/VIAL  
N19640 004  
MAR 08, 1986

AI  
CREAM; VAGINAL  
AVC  
MERRELL DOM

15%  
15%

N06530 003  
JAN 27, 1987

SPIRONOLACTONE

TABLET; ORAL

AA/  
SPIRONOLACTONE  
/SUPERPHARM/

25MG  
N89364 001  
NOV 07, 1986

AI  
VAGITROL  
LEMMON

N88718 001  
SEP 19, 1985

SULFACETAMIDE SODIUM

SOLUTION/DROPS; OPHTHALMIC  
SULFACETAMIDE SODIUM

AI  
STERIS LABS  
30%  
N89068 001  
MAY 05, 1987

AI  
SUPPOSITORY; VAGINAL  
AVC  
MERRELL DOM

1.05GM  
165MG

N06044 003

SULFAMETHOXAZOLE; TRIMETHOPRIM

INJECTABLE; INJECTION

AP  
SULFAMETHOXAZOLE AND TRIMETHOPRIM  
ELKINS SINN

80MG/ML; 16MG/ML  
DEC 29, 1987 : APR 30, 1987  
N70627 001  
80MG/ML; 16MG/ML  
DEC 29, 1987 : APR 30, 1987  
N70628 001  
80MG/ML; 16MG/ML  
DEC 29, 1987 : JAN 16, 1987  
N70223 001

AP  
LYPHOMED

N18217 001  
DEC 24, 1985

TABLET; ORAL

AB/  
SULFAMETHOXAZOLE AND TRIMETHOPRIM DOUBLE STRENGTH  
/PLANTEX/

800MG; 160MG  
SEP 19, 1985  
N70037 001

AB  
STUART PHARMS  
EQ 10MG BASE  
BARR LABS

N17970 001  
N70929 001  
AUG 20, 2002 : APR 01, 1987

AB/  
SULFAMETHOXAZOLE AND TRIMETHOPRIM SINGLE STRENGTH  
/PLANTEX/

400MG; 80MG  
SEP 19, 1985  
N70030 001

AB  
TECHNETIUM TC-99M MEBROFENIN KIT

N18963 001  
JAN 21, 1987

INJECTABLE; INJECTION  
CHOLETEC  
SQUIBB DIAGS

N/A

TECHNETIUM Tc-99m PYROPHOSPHATE KIT

INJECTABLE; INJECTION  
AN-PYROTEC

AP CIS US N/AM N19039 001  
JUN 30, 1987

TEMZEPAM

CAPSULE; ORAL  
TEMZEPAM

AB BOLAR PHARM 15MG# N70383 001  
MAR 23, 1987  
AB 30MG# N70384 001  
MAR 23, 1987  
AB PAR PHARM 15MG# N71456 001  
APR 21, 1987  
AB 30MG# N71457 001  
APR 21, 1987

THEOPHYLLINE

TABLET, CONTROLLED RELEASE; ORAL

AB DURAPHYL 300MG# N88505 001  
APR 03, 1985  
BC FOREST LABS 100MG N88503 001  
APR 03, 1985  
BC 200MG N88504 001  
APR 03, 1985  
BC LABID 250MG N87225 001  
NORWICH EATON  
BC THEOLAIR-SR 200MG# N88369 001  
RIKER LABS 250MG# N86363 001  
JUL 16, 1987  
BC 300MG# N88364 001  
JUL 16, 1987  
BC 500MG# N89132 001  
JUL 16, 1987

/AB/ /THEOPHYLLINE/  
/BC/ /FOREST/LABS/  
/BC/ /

/N88505/001/  
/APR/03,1985/  
/N88503/001/  
/APR/03,1985/  
/N88504/001/  
/APR/03,1985/

THIOTHIXENE

CAPSULE; ORAL

AB NAVANE ROERIG 1MG N16584 001  
AB 2MG N16584 002  
AB 5MG N16584 003  
AB 10MG N16584 004  
AB CHELSEA LABS 2MG# N71626 001  
JUN 25, 1987  
AB 5MG# N71627 001  
JUN 25, 1987  
AB 10MG# N71628 001  
JUN 25, 1987  
AB CORD LABS 1MG# N71610 001  
JUN 24, 1987  
AB 2MG# N71570 001  
JUN 24, 1987  
AB 5MG# N71529 001  
JUN 24, 1987  
AB 10MG# N71530 001  
JUN 24, 1987  
AB DANBURY PHARMA 1MG# N70600 001  
JUN 05, 1987  
AB 2MG# N70601 001  
JUN 05, 1987  
AB 5MG# N70602 001  
JUN 05, 1987  
AB 10MG# N70603 001  
JUN 05, 1987  
AB MYLAN PHARMS 1MG# N71090 001  
JUN 23, 1987  
AB 2MG# N71091 001  
JUN 23, 1987  
AB 5MG# N71092 001  
JUN 23, 1987  
AB 10MG# N71093 001  
JUN 23, 1987

THIOTHIXENE HYDROCHLORIDE

CONCENTRATE; ORAL

AA NAVANE EQ 5MG BASE/ML N16758 001  
AB THIOTHIXENE HCL  
LEMMON EQ 5MG BASE/ML# N71184 001  
JUN 22, 1987

TOBRAMYCIN SULFATE

INJECTABLE; INJECTION  
NEBCIN  
LILLY

EQ 10MG BASE/MLM

N62707 001  
APR 29, 1987

INJECTABLE; INJECTION  
TRIMETHOBENZAMIDE HCL

AP  
WINTHROP BREON  
100MG/MLM

N88804 001  
APR 03, 1987

TOLAZAMIDE

TABLET; ORAL  
TOLAZAMIDE  
MUTUAL PHARM

>\_ADD\_>  
>\_ADD\_>  
>\_ADD\_>  
>\_ADD\_>  
>\_ADD\_>  
>\_ADD\_>

100MGH  
250MGH  
500MGH

N71357 001  
JUL 16, 1987  
N71358 001  
JUL 16, 1987  
N71359 001  
JUL 16, 1987

TABLET; ORAL  
TRIMETHOPRIM  
BIOCRAFT LABS

AB  
200MGH

N71259 001  
JUN 18, 1987

TOLBUTAMIDE

TABLET; ORAL  
TOLBUTAMIDE  
BOLAR PHARM

AB  
AB

250MGH  
500MGH

N89110 001  
MAY 29, 1987  
N89111 001  
MAY 29, 1987

CAPSULE; ORAL  
VALPROIC ACID  
FORMUTEC

AB  
>\_ADD\_>  
>\_ADD\_>  
250MGH  
250MGH

N70631 001  
JUN 11, 1987  
N70195 001  
JUL 02, 1987

TRAZODONE HYDROCHLORIDE

TABLET; ORAL  
TRAZODONE HCL  
BARR LABS

AB  
AB  
AB  
AB

50MGH  
100MGH  
50MGH  
100MGH

N71258 001  
MAR 25, 1987  
N71196 001  
MAR 25, 1987  
N70491 001  
APR 29, 1987  
N70492 001  
APR 29, 1987

INJECTABLE; INJECTION  
LYPHOCIN  
LYPHOMED

AP  
EQ 500MG BASE/VIALM

N62663 001  
MAR 17, 1987

VANCOCCIN HCL  
LILLY

AP  
EQ 500MG BASE/VIALM  
EQ 1GM BASE/VIALM

N62716 001  
MAR 13, 1987  
N62716 002  
MAR 13, 1987

TRIAMCINOLONE ACETONIDE

PASTE; DENTAL  
ORALONE  
THAMES PHARMA

>\_ADD\_>  
>\_ADD\_>  
>\_ADD\_>

0.12M

N71383 001  
JUL 06, 1987

INJECTABLE; INJECTION  
VERAPAMIL HCL  
ABBOTT LABS

AP  
2.5MG/MLM  
2.5MG/MLM  
2.5MG/MLM  
2.5MG/MLM

N70737 001  
MAY 06, 1987  
N70738 001  
MAY 06, 1987  
N70739 001  
MAY 06, 1987  
N70740 001  
MAY 06, 1987

RX DRUG PRODUCT LIST / CUMULATIVE SUPPLEMENT NUMBER 7 / JAN '87 - JUL '87

VERAPAMIL HYDROCHLORIDE

INJECTABLE; INJECTION  
VERAPAMIL HCL

SOLOPAK LABS

> ADD > AP  
> ADD >  
> ADD > AP  
> ADD >  
> ADD > AP  
> ADD >  
> ADD >

2.5MG/ML  
2.5MG/ML  
2.5MG/ML  
2.5MG/ML

N70695 001  
JUL 31, 1987  
N70696 001  
JUL 31, 1987  
N70697 001  
JUL 31, 1987  
N70577 001  
FEB 02, 1987

6.3MCI/ML

N17283 001

XENON, XE-133

INJECTABLE; INJECTION

XENON XE 133  
@ DUPONT DIAG

> ADD >

XYLOSE

POWDER; ORAL

XYLO-PFAN  
ADRIA LABS

AA

N17605 001

AA

25GM/BOI  
25GM/BOI

N18856 001  
MAR 26, 1987

VINBLASTINE SULFATE

INJECTABLE; INJECTION

VINBLASTINE SULFATE

BEN VENUE LABS

AP

10MG/VIAL

N89395 001  
APR 09, 1987  
N89515 001  
APR 29, 1987  
N89311 001  
MAR 23, 1987

ZIDOVUDINE

CAPSULE; ORAL

RETROVIR  
BURROUGHS WELLC

100MG

N19655 001  
MAR 19, 1987

VINCRIStINE SULFATE

INJECTABLE; INJECTION

VINCASAR PFS

ADRIA LABS

AP

1MG/ML

N71426 001  
JUL 17, 1987

INJECTABLE; INJECTION

ZINC SULFATE  
LYPHOMED

EQ 1MG ZINC/ML

N19229 002  
MAY 05, 1987

VINCRIStINE SULFATE

INTL PHARM

AP

1MG/ML

N70873 001  
FEB 19, 1987

WARFARIN POTASSIUM

TABLET; ORAL

ATHROMBIN-K

@ PURDUE FRDRK

@

@

2MG  
10MG  
25MG

N11771 007  
N11771 005  
N11771 006

WARFARIN SODIUM

TABLET; ORAL

ATHROMBIN

BX @ PURDUE FRDRK

BX @

@

5MG  
10MG  
25MG

N11771 003  
N11771 002  
N11771 001

ACETAMINOPHEN

SUPPOSITORY; RECTAL  
ACETAMINOPHEN  
ROXANE LABS

120MG#

N71010 001  
MAY 12, 1987

N71011 001  
MAY 12, 1987

SUPPOSITORIA

120MG#

N70607 001  
APR 06, 1987

UPSHER SMITH

325MG#

N18337 002

DEXBROMPHENIRAMINE MALEATE; PSEUDOEPHEDRINE SULFATE

TABLET, CONTROLLED RELEASE; ORAL  
BROMPHERIL

6MG;120MG#

N89116 001  
JAN 22, 1987

DIPHENHYDRAMINE HYDROCHLORIDE

SYRUP; ORAL  
ANTITUSSIVE  
PERRIGO

12.5MG/5ML#

N71292 001  
APR 10, 1987

VICKS FORMULA 44

12.5MG/5ML#

N70524 001  
JAN 14, 1987

DOXYLAMINE SUCCINATE

TABLET; ORAL  
DOXY-SLEEP-AID  
PAR PHARM

25MG#

N70156 001  
JUL 02, 1987

ASPIRIN

TABLET, CONTROLLED RELEASE; ORAL  
MEASURIN

650MG#

N16030 002

WINTHROP BREON

8-HOUR BAYER

650MG#

N16030 001

BACITRACIN

OINTMENT; TOPICAL  
BACITRACIN  
COMBE

500 UNITS/GM#

N62799 001  
MAY 14, 1987

TABLET; ORAL  
ACHES-N-PAIN  
LEDERLE LABS

200MG#

N71065 001  
MAY 28, 1987

IBUPRIN  
SIDMAK LABS

200MG#

N71773 001  
JUL 16, 1987

IBUPROFEN  
INTERPHARM

200MG#

N71333 001  
FEB 17, 1987

MUTUAL PHARM

200MG#

N71229 001  
APR 01, 1987

PAR PHARM

200MG#

N71575 001  
MAY 08, 1987

PUREPAC PHARM

200MG#

N71664 001  
FEB 03, 1987

NEUVIL

200MG#

N71144 001  
JAN 20, 1987

TRENDAR  
WHITEHALL LABS

200MG

N16989 002  
JUL 10, 1986

CHLORHEXIDINE GLUCONATE

SPONGE; TOPICAL  
CHLORHEXIDINE GLUCONATE  
KENDALL

4/M

N19490 001  
MAR 27, 1987

> ADD >  
> ADD >  
> ADD >

INSULIN ZINC SUSP EXTENDED BIOSYNTHETIC HUMAN

INJECTABLE; INJECTION			
HUMULIN U			
LILLY			
	40 UNITS/ML		N19571 001
			JUN 10, 1987
	100 UNITS/ML		N19571 002
			JUN 10, 1987

POVIDONE-IODINE

Sponge; Topical			
E-Z SCRUB 241			
DESERET MED			
	10%		N19476 001
			JAN 07, 1987

PSEUDOEPHEDRINE POLISTIREX

Suspension, Controlled Release; Oral			
PSEUDO-12			
PENNMALT			
	EQ 60MG HCL/5ML		N19401 001
			JUN 19, 1987

SODIUM MONOFLUOROPHOSPHATE

Paste; Dental			
EXTRA-STRENGTH AIM			
LEVER BROTHERS			
	1.2%		N19518 001
			JUN 03, 1987

PENTASTARCH 10% IN SODIUM CHLORIDE 0.9%

INJECTABLE; INJECTION  
PENTASPAR ( )  
DUPONT CRI CARE

10G/100ML; 0.9G/100ML

N 841207  
MAY 19, 1987

## ORPHAN DRUG PRODUCTS WITH EXCLUSIVE APPROVAL

SECTION 526 OF THE FEDERAL FOOD, DRUG, AND COSMETIC ACT CONTAINS PROVISIONS WHEREBY FDA MAY DESIGNATE A SPONSOR'S DRUG, ANTIBIOTIC, OR BIOLOGICAL PRODUCT AS A "DESIGNATED ORPHAN DRUG". SECTION 527 OF THE ACT ESTABLISHES A PROCESS WHEREBY A SPONSOR MAY RECEIVE SEVEN YEARS OF EXCLUSIVE APPROVAL STATUS IF THAT SPONSOR IS THE FIRST TO ACHIEVE NEW DRUG, ANTIBIOTIC, OR BIOLOGICAL PRODUCT APPROVAL FOR A DESIGNATED ORPHAN DRUG FOR THE DESIGNATED INDICATION(S). THE EXCLUSIVE APPROVAL MAY BE REVOKED BY WRITTEN CONSENT OF THE SPONSOR OR BY FDA ACTION AFTER FINDING THAT THE SPONSOR HOLDING EXCLUSIVE APPROVAL CANNOT ASSURE THE AVAILABILITY OF SUFFICIENT QUANTITIES OF THE DRUG TO MEET THE NEEDS OF PATIENTS WITH THE DESIGNATED ORPHAN INDICATION(S).

ORPHAN DRUG EXCLUSIVE APPROVAL STATUS (CODED ODE) APPLIES ONLY TO THE APPROVED OR LICENSED INDICATION(S) FOR WHICH ORPHAN DRUG DESIGNATION HAS BEEN GRANTED PURSUANT TO SECTION 526 OF THE ACT.

FOR THE FOLLOWING DRUG PRODUCTS WITH ORPHAN DRUG EXCLUSIVE APPROVAL STATUS, THE SPONSOR HAS SEVEN YEARS OF EXCLUSIVE APPROVAL FOR THE APPROVED INDICATION BEGINNING ON THE DATE OF NDA, ANTIBIOTIC APPLICATION, OR BIOLOGICAL LICENSE APPROVAL FOR THE DRUG. NO SUBSEQUENT SPONSOR MAY RECEIVE APPROVAL OF AN NDA, BIOLOGICAL LICENSE, PAPER NDA, ANTIBIOTIC APPLICATION, ANDA, OR ABBREVIATED ANTIBIOTIC APPLICATION DURING THE SEVEN YEAR PERIOD FOR THE DRUG AND INDICATION(S) FOR WHICH A PERSON MAINTAINS ODE STATUS UNLESS THE EXCLUSIVE APPROVAL HAS BEEN REVOKED AS DESCRIBED ABOVE OR THE SUBSEQUENT SPONSOR HAS OBTAINED WRITTEN CONSENT FROM THE SPONSOR WHO HAS RECEIVED EXCLUSIVE APPROVAL.

BIOLOGICAL PRODUCTS, ANTIBIOTICS, AND DRUGS THAT HAVE BEEN APPROVED UNDER SECTION 505 OR 507 OF THE ACT OR UNDER SECTION 351 OF THE PUBLIC HEALTH SERVICE ACT FOR MARKETING AND HAVE BEEN GIVEN ORPHAN DRUG EXCLUSIVE APPROVAL WILL BE NOTED BY THE ABBREVIATION ODE IN THE PATENT AND EXCLUSIVITY DATA APPENDIX. DRUG PRODUCTS THAT HAVE RECEIVED THE WRITTEN PERMISSION OF THE SPONSOR THAT HAS ORPHAN DRUG EXCLUSIVE APPROVAL TO BE APPROVED UNDER SECTION 527(B)(2) OF THE ACT ARE ALSO NOTED BY THE ABBREVIATION ODE IN THE PATENT AND EXCLUSIVITY DATA APPENDIX. THESE DRUG PRODUCTS DO NOT HAVE ANY EXCLUSIVE APPROVAL RIGHTS OF THEIR OWN, BUT CAN BE MARKETED BECAUSE OF THE CONSENT GIVEN BY THE SPONSOR THAT HAS EXCLUSIVE APPROVAL. THESE PRODUCTS ARE MARKED BY AN (\*) NEXT TO THE APPLICANT'S NAME.

REFER BACK TO THE APPROVED DRUG PRODUCTS WITH THERAPEUTIC EQUIVALENCE EVALUATIONS, 7TH EDITION FOR A FULL LISTING OF ORPHAN DRUG PRODUCTS WITH EXCLUSIVE APPROVAL DATA. ONLY NEW DATA WILL BE ADDED TO THE CUMULATIVE SUPPLEMENT.

## ORPHAN DRUG PRODUCTS WITH EXCLUSIVE APPROVAL

## DRUG PRODUCTS

ACTIVE INGREDIENT(S) STRENGTH(S)	TRADE NAME DOSAGE FORM; ROUTE	APPLICANT	APPLICATION NUMBER APPROVAL DATE	EXCLUSIVITY EXP. DATE
CALCITONIN, HUMAN 0.5MG/VIAL	CIBACALCIN INJECTABLE; INJECTION	CIBA PHARM	18470 001 OCT 31, 1986	ODE OCT 31, 1993
ETIDRONATE DISODIUM 50MG/ML	DIDRONEL I.V. INJECTABLE; INJECTION	NORWICH EATON	19545 001 APR 24, 1987	ODE APR 24, 1994
PENTASTARCH 10% IN SODIUM CHLORIDE 0.9% 10GM/100ML; 0.9GM/100ML	PENTASPERN INJECTABLE; INJECTION	DUPONT CRI CARE	841207 001 MAY 19, 1987	ODE MAY 19, 1994
SOMATROPIN, BIOSYNTHETIC 5MG/VIAL	HUMATROPE INJECTABLE; INJECTION	LILLY	19640 004 MAR 08, 1987	ODE MAR 08, 1994
ZIDOVUDINE 100MG	RETROVIR CAPSULE; ORAL	BURROUGHS WELLC	19655 001 MAR 19, 1987	ODE MAR 19, 1994

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

NO JULY 1987 ACTIONS

## BIOPHARMACEUTIC GUIDANCE AVAILABILITY

THE FOLLOWING IS A LIST OF GUIDANCES AVAILABLE FOR IN VIVO BIOEQUIVALENCE STUDIES AND IN VITRO DISSOLUTION TESTING AVAILABLE FROM THE DIVISION OF BIOEQUIVALENCE, HFN-250, ROOM 17B-06, 5600 FISHERS LANE, ROCKVILLE, MD 20857. COMMENTS AND SUGGESTIONS CONCERNING THESE GUIDANCES ARE ENCOURAGED AND SHOULD BE SENT TO THE DIVISION OF BIOEQUIVALENCE.

REFER BACK TO THE APPROVED DRUG PRODUCTS WITH THERAPEUTIC EQUIVALENCE EVALUATIONS, 7TH EDITION FOR A FULL LISTING OF BIOPHARMACEUTIC GUIDANCE AVAILABILITY DATA. ONLY NEW DATA WILL BE ADDED TO THE CUMULATIVE SUPPLEMENT.

NAME OF DRUG	DATE	REVISED DATE
ALBUTEROL (TABLET)	MAY 05, 1987	
CEPHALEXIN (TABLET AND CAPSULE)	AUG 13, 1986	MAR 19, 1987
CLORAZEPATE DIPOTASSIUM	MAR 10, 1986	FEB 17, 1987
DESIPRAMINE HYDROCHLORIDE (TABLET)	APR 28, 1987	
DISSOLUTION TESTING (GENERAL)	APR 01, 1978*	
HALOPERIDOL (TABLET)	APR 30, 1987	
LEUCOVORIN CALCIUM (TABLET)	APR 28, 1987	
POTASSIUM CHLORIDE (TABLET AND CAPSULE, SLOW RELEASE)	JAN 17, 1987	

\* THIS DATE WAS INCORRECTLY LISTED IN THE 7TH EDITION AS APR 19, 1985.

## ANDA SUITABILITY PETITIONS

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) AND (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, 7TH EDITION FOR A FULL LISTING OF ANDA SUITABILITY PETITIONS DATA. ONLY NEW DATA WILL BE ADDED TO THE CUMULATIVE SUPPLEMENT.

## PETITIONS APPROVED

DRUG NAME DOSAGE FORM; ROUTE	STRENGTH (CONTAINER SIZE)	DOCKET NUMBER	PETITIONER	REASON FOR PETITION	STATUS
ACETAMINOPHEN; BUTALBITAL; CAFFEINE TABLET; ORAL	500MG 50MG 40MG	86 P-0514/CP	FOREST LABS	NEW STRENGTH	APPROVED JUL 15, 1987
ACETAMINOPHEN; HYDROCODONE BITARTRATE TABLET; ORAL	325MG 2.5MG	87 P-0129/CP	MIKART	NEW STRENGTH	APPROVED JUN 08, 1987

ANDA SUITABILITY PETITIONS

PETITIONS APPROVED

DRUG NAME DOSAGE FORM; ROUTE	STRENGTH (CONTAINER SIZE)	DOCKET NUMBER	PETITIONER	REASON FOR PETITION	STATUS
ACETAMINOPHEN; HYDROCODONE BITARTRATE TABLET; ORAL	325MG 5MG	87 P-0129/CP	MIKART	NEW STRENGTH	APPROVED JUN 08, 1987
ACETAMINOPHEN; HYDROCODONE BITARTRATE TABLET; ORAL	325MG 7.5MG	87 P-0129/CP	MIKART	NEW STRENGTH	APPROVED JUN 08, 1987
ACETAMINOPHEN; HYDROCODONE BITARTRATE TABLET; ORAL	325MG 10MG	87 P-0129/CP	MIKART	NEW STRENGTH	APPROVED JUN 08, 1987
ACETAMINOPHEN; HYDROCODONE BITARTRATE SOLUTION; ORAL	325MG/15ML 2.5MG/15ML	87 P-0129/ CP02	MIKART	NEW STRENGTH	APPROVED JUN 08, 1987

## ANDA SUITABILITY PETITIONS

## PETITIONS APPROVED

DRUG NAME DOSAGE FORM; ROUTE	STRENGTH (CONTAINER SIZE)	DOCKET NUMBER	PETITIONER	REASON FOR PETITION	STATUS
ACETAMINOPHEN; HYDROCODONE BITARTRATE SOLUTION; ORAL	325MG/15ML 5MG/15ML	87 P-0129/ CP02	MIKART	NEW STRENGTH	APPROVED JUN 08, 1987
ACETAMINOPHEN; HYDROCODONE BITARTRATE SOLUTION; ORAL	325MG/15ML 7.5MG/15ML	87 P-0129/ CP02	MIKART	NEW STRENGTH	APPROVED JUN 08, 1987
ACETAMINOPHEN; HYDROCODONE BITARTRATE SOLUTION; ORAL	325MG/15ML 10MG/15ML	87 P-0129/ CP02	MIKART	NEW STRENGTH	APPROVED JUN 08, 1987
ACETAMINOPHEN; HYDROCODONE BITARTRATE LIQUID; ORAL	500MG/15ML 7.5MG/15ML	85 P-0439/ CP0003	RUSS PHARMS	NEW STRENGTH	APPROVED APR 01, 1987

## ANDA SUITABILITY PETITIONS

## PETITIONS APPROVED

DRUG NAME DOSAGE FORM; ROUTE	STRENGTH (CONTAINER SIZE)	DOCKET NUMBER	PETITIONER	REASON FOR PETITION	STATUS
ACETAMINOPHEN; HYDROCODONE BITARTRATE TABLET; ORAL	500MG 2.5MG	85 P-0439/ CP002	KING AND SPAULDING	NEW STRENGTH	APPROVED MAR 18, 1987
ACETAMINOPHEN; HYDROCODONE BITARTRATE TABLET; ORAL	500MG 7.5MG	85 P-0439/CP	KING AND SPAULDING	NEW STRENGTH	APPROVED MAR 17, 1987
ACETAMINOPHEN; HYDROCODONE BITARTRATE TABLET; ORAL	500MG 10MG	87 P-0170/CP	LUCHEM PHARM	NEW STRENGTH	APPROVED JUL 07, 1987
ACETAMINOPHEN; HYDROCODONE BITARTRATE CAPSULE; ORAL	650MG 7.5MG	85 P-0390/CP	UAD LABS	NEW STRENGTH NEW DOSAGE FORM	APPROVED MAR 17, 1987

## ANDA SUITABILITY PETITIONS

## PETITIONS APPROVED

DRUG NAME DOSAGE FORM; ROUTE	STRENGTH (CONTAINER SIZE)	DOCKET NUMBER	PETITIONER	REASON FOR PETITION	STATUS
ACETAMINOPHEN; HYDROCODONE BITARTRATE TABLET; ORAL	650MG 7.5MG	85 P-0390/CP	UAD LABS	NEW STRENGTH NEW DOSAGE FORM	APPROVED MAR 17, 1987
ACETAMINOPHEN; HYDROCODONE BITARTRATE TABLET; ORAL	750MG 7.5MG	85 P-0169/PRC*	KNOLL PHARM	NEW STRENGTH	APPROVED MAR 13, 1987
AMINOPHYLLINE INJECTABLE; INJECTION	10MG/ML (10ML/VIAL)	87 P-0103/CP	LYPHOMED	NEW STRENGTH	APPROVED JUL 07, 1987
ASPIRIN; HYDROCODONE BITARTRATE TABLET; ORAL	500MG 7.5MG	87 P-0100/CP	KING AND SPAULDING	NEW STRENGTH	APPROVED APR 24, 1987
BRETYLIUM TOSYLATE INJECTABLE; INJECTION	200MG/ML (10ML/CONTAINER)	85 P-0546/CP	INTL MEDTN SYS	NEW STRENGTH	APPROVED JAN 20, 1987

\*ORIGINAL PETITION DENIED NOV 07, 1985; PETITION FOR RECONSIDERATION APPROVED MAR 13, 1987.

## ANDA SUITABILITY PETITIONS

## PETITIONS APPROVED

DRUG NAME DOSAGE FORM; ROUTE	STRENGTH (CONTAINER SIZE)	DOCKET NUMBER	PETITIONER	REASON FOR PETITION	STATUS
BRETYLIUM TOSYLATE IN DEXTROSE 5% INJECTABLE; INJECTION	10MG/ML (50ML/CONTAINER)	87 P-0065/CP	LYPHOMED	NEW STRENGTH	APPROVED APR 27, 1987
BRETYLIUM TOSYLATE IN DEXTROSE 5% INJECTABLE; INJECTION	10MG/ML (100ML/CONTAINER)	87 P-0128/CP	LYPHOMED	NEW STRENGTH	APPROVED JUL 22, 1987
CHLORPHENIRAMINE MALEATE; PSEUDOEPHEDRINE HYDROCHLORIDE TABLET, CONTROLLED RELEASE; ORAL	12MG 120MG	87 P-0165/CP	SANDOZ CONSUMER	NEW DOSAGE FORM	APPROVED MAY 19, 1987
CHOLESTYRAMINE CAPSULE; ORAL	EQ 500MG RESIN	86 P-0474/CP	BRISTOL MYERS	NEW DOSAGE FORM NEW STRENGTH	APPROVED JAN 30, 1987
CHOLESTYRAMINE TABLET; ORAL	EQ 800MG RESIN	86 P-0475/CP	BRISTOL MYERS	NEW DOSAGE FORM NEW STRENGTH	APPROVED JAN 30, 1987

ANDA SUITABILITY PETITIONS

PETITIONS APPROVED

DRUG NAME DOSAGE FORM; ROUTE	STRENGTH (CONTAINER SIZE)	DOCKET NUMBER	PETITIONER	REASON FOR PETITION	STATUS
CYTARABINE INJECTABLE; INJECTION	1000MG/VIAL	86 P-0313/CP	QUAD PHARMS	NEW STRENGTH	APPROVED MAY 07, 1987
CYTARABINE INJECTABLE; INJECTION	20MG/ML (50ML CONTAINER)	86 P-0428/ CP0002	ADRIA LABS	NEW STRENGTH	APPROVED MAY 07, 1987
DEXTROMETHORPHAN POLISTIREX SUSPENSION, CONTROLLED RELEASE; ORAL	EQ 15MG HBR/5ML	87 P-0088/CP	KING AND SPAULDING	NEW STRENGTH	APPROVED APR 27, 1987
DIAZOXIDE INJECTABLE; INJECTION	15MG/ML (10ML/CONTAINER)	87 P-0061/CP	LYPHOMED	NEW STRENGTH	APPROVED APR 30, 1987
FENOPROFEN CALCIUM TABLET; ORAL	EQ 200MG BASE EQ 300MG BASE	87 P-0133/CP	BARR LABS	NEW STRENGTH	APPROVED AUG 04, 1987
FLUOROURACIL INJECTABLE; INJECTION	50MG/ML (50ML/VIAL)	86 P-0490/CP	ADRIA LABS	NEW STRENGTH	APPROVED JAN 09, 1987
LEUCOVORIN CALCIUM INJECTABLE; INJECTION	EQ 25MG BASE/VIAL	86 P-0240/CP	BURROUGHS WELLC	NEW STRENGTH	APPROVED JAN 29, 1987
LEUCOVORIN CALCIUM INJECTABLE; INJECTION	EQ 5MG BASE/ML (10ML AND 20ML/VIALS)	86 P-0241/CP	QUAD PHARMS	NEW STRENGTH	APPROVED JUL 28, 1987

## ANDA SUITABILITY PETITIONS

## PETITIONS APPROVED

DRUG NAME DOSAGE FORM; ROUTE	STRENGTH (CONTAINER SIZE)	DOCKET NUMBER	PETITIONER	REASON FOR PETITION	STATUS
LEUCOVORIN CALCIUM INJECTABLE; INJECTION	EQ 100MG BASE/VIAL	86 P-0152/CP	BEN VENUE LABS	NEW STRENGTH	APPROVED JAN 20, 1987
LEUCOVORIN CALCIUM TABLET; ORAL	EQ 10MG BASE	86 P-0258/CP	LEDERLE LABS	NEW STRENGTH	APPROVED JAN 16, 1987
LORAZEPAM SOFT GELATIN CAPSULE; ORAL	0.5MG 1MG 2MG	87 P-0037/CP	APPLIED LABS	NEW DOSAGE FORM	APPROVED MAR 10, 1987
LORAZEPAM TABLET; ORAL	0.5MG 1MG 2MG	85 P-0515/CP	WYETH INC	NEW DOSAGE FORM	APPROVED FEB 25, 1986
METHYLDOPATE HYDROCHLORIDE IN 5% DEXTROSE INJECTABLE; INJECTION	2.5MG/ML (100ML/CONTAINER)	86 P-0410/ CP0002	KING AND SPAULDING	NEW STRENGTH	APPROVED MAR 10, 1987
METHYLDOPATE HYDROCHLORIDE IN 5% DEXTROSE INJECTABLE; INJECTION	5MG/ML (100ML/CONTAINER)	86 P-0410/ CP0003	KING AND SPAULDING	NEW STRENGTH	APPROVED MAR 10, 1987

## ANDA SUITABILITY PETITIONS

## PETITIONS APPROVED

DRUG NAME DOSAGE FORM; ROUTE	STRENGTH (CONTAINER SIZE)	DOCKET NUMBER	PETITIONER	REASON FOR PETITION	STATUS
MORPHINE SULFATE INJECTABLE; INJECTION	0.5MG/ML (2ML/AMP)	87 P-0106/CP	ASTRA PHARM PRODS	NEW STRENGTH	APPROVED JUL 15, 1987
MORPHINE SULFATE INJECTABLE; INJECTION	1MG/ML (2ML/AMP)	87 P-0106/CP	ASTRA PHARM PRODS	NEW STRENGTH	APPROVED JUL 15, 1987
OXAZEPAM CAPSULE; ORAL	10MG 15MG 30MG	87 P-0157/CP	BARR LABS	NEW DOSAGE FORM NEW STRENGTH	APPROVED JUL 17, 1987
OXAZEPAM TABLET; ORAL	15MG 30MG	85 P-0516/CP	WYETH INC	NEW DOSAGE FORM	APPROVED FEB 25, 1986
NITROGLYCERIN IN DEXTROSE 5% INJECTABLE; INJECTION	0.5MG/ML (100ML/CONTAINER)	86 P-0099/ CP0004	ABBOTT LABS	NEW STRENGTH	APPROVED FEB 02, 1987
PROMETHAZINE HYDROCHLORIDE INJECTABLE; INJECTION	25MG/ML (2ML/VIAL)	87 P-0087/ CP00002	LYPHOMED	NEW STRENGTH	APPROVED MAY 01, 1987
PROMETHAZINE HYDROCHLORIDE INJECTABLE; INJECTION	50MG/ML (2ML/VIAL)	87 P-0087/CP	LYPHOMED	NEW STRENGTH	APPROVED MAY 01, 1987

## ANDA SUITABILITY PETITIONS

## PETITIONS APPROVED

DRUG NAME DOSAGE FORM; ROUTE	STRENGTH (CONTAINER SIZE)	DOCKET NUMBER	PETITIONER	REASON FOR PETITION	STATUS
SODIUM NITROPRUSSIDE INJECTABLE; INJECTION	25MG/ML (2ML/VIAL)	87 P-0039/CP	ABBOTT LABS	NEW DOSAGE FORM	APPROVED MAR 10, 1987
THEOPHYLLINE CAPSULE, CONTROLLED RELEASE; ORAL	400MG	86 P-0471/ CP0002	SEARLE RESEARCH AND DEVELOPMENT	NEW STRENGTH	APPROVED MAR 10, 1987
VINBLASTINE SULFATE INJECTABLE; INJECTION	1MG/ML (25ML/VIAL)	87 P-0112/CP	QUAD PHARMS	NEW DOSAGE FORM NEW STRENGTH	APPROVED JUN 08, 1987
VINBLASTINE SULFATE INJECTABLE; INJECTION	1MG/ML (30ML/VIAL)	87 P-0211/CP	LYPHOMED	NEW STRENGTH	APPROVED JUL 28, 1987

## ANDA SUITABILITY PETITIONS

## PETITIONS DENIED

DRUG NAME DOSAGE FORM; ROUTE	STRENGTH (CONTAINER SIZE)	DOCKET NUMBER	PETITIONER	REASON FOR PETITION	STATUS
ACETAMINOPHEN; DIHYDROCODEINE BITARTRATE CAPSULE; ORAL	356.4MG 20MG	86 P-0040/CP	DUNHALL PHARMACEUTICALS	NEW STRENGTH NEW COMBINATION	DENIED FEB 12, 1987
ASPIRIN; CAFFEINE; HYDROCODONE BITARTRATE TABLET; ORAL	224MG 32MG 5MG	86 P-0243/CP	MASON PHARMS INC	NEW COMBINATION NEW DOSAGE FORM NEW STRENGTH	DENIED JUN 12, 1987
ASPIRIN; CAFFEINE; HYDROCODONE BITARTRATE TABLET; ORAL	325MG 30MG 5MG	85 P-0455/CP	CENTRAL PHARM	NEW COMBINATION NEW DOSAGE FORM NEW STRENGTH	DENIED JUN 08, 1987
ASPIRIN; CAFFEINE; HYDROCODONE BITARTRATE TABLET; ORAL	356.4MG 30MG 5MG	86 P-0243/ CP002	MASON PHARMS INC	NEW COMBINATION NEW DOSAGE FORM	DENIED JUN 16, 1987
HYDROCORTISONE; SALICYLIC ACID; SULFUR CREAM; TOPICAL	0.25% 2.35% 4%	86 P-0439/CP	C&M PHARMA	NEW COMBINATION NEW INGREDIENT	DENIED MAY 06, 1987

## ANDA SUITABILITY PETITIONS

## PETITIONS DENIED

DRUG NAME DOSAGE FORM; ROUTE	STRENGTH (CONTAINER SIZE)	DOCKET NUMBER	PETITIONER	REASON FOR PETITION	STATUS
PROCAINAMIDE HYDROCHLORIDE TABLET; ORAL	500MG 750MG 1000MG	85 P-0181/CP	FOREST LABS	NEW DOSAGE FORM	DENIED APR 21, 1987
PROCAINAMIDE HYDROCHLORIDE TABLET, CONTROLLED RELEASE; ORAL	500MG 750MG 1000MG	86 P-0328/CP	KV PHARM	NEW DOSAGE FORM	DENIED APR 21, 1987

## EXCLUSIVITY TERMS

DUE TO SPACE LIMITATIONS IN THE EXCLUSIVITY COLUMN, ABBREVIATIONS AND REFERENCES HAVE BEEN DEVELOPED. REFER BACK TO THE APPROVED DRUG PRODUCTS WITH THERAPEUTIC EQUIVALENCE EVALUATIONS, 7TH 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-13 INCREASED MAXIMUM DAILY DOSAGE RECOMMENDATION

## NEW INDICATION

I-54 CONTRAST ENHANCEMENT OF COMPUTED TOMOGRAPHIC BODY IMAGING  
 I-55 PEDIATRIC ANGIOCARDIOGRAPHY  
 I-56 INTRAVENOUS DIGITAL SUBTRACTION ANGIOGRAPHY  
 I-57 PERIPHERAL VENOGRAPHY (PHLEBOGRAPHY)  
 I-58 EXCRETORY UROGRAPHY  
 I-59 ARTHROGRAPHY  
 I-60 HYSTEROSALPINGOGRAPHY  
 I-61 AORTOGRAPHY  
 I-62 TREATMENT OF JUVENILE ARTHRITIS  
 I-63 BIOPSY PROVEN MINIMAL CHANGE NEPHROTIC SYNDROME IN CHILDREN  
 I-64 LONG-TERM TREATMENT OF ANGINA PECTORIS  
 I-65 ADULT INTRAVENOUS CONTRAST ENHANCED COMPUTED TOMOGRAPHY OF THE HEAD AND BODY  
 I-66 PREVENTION OF POSTOPERATIVE NAUSEA AND VOMITING  
 I-67 PREVENTION OF POSTOPERATIVE DEEP VEIN THROMBOSIS AND PULMONARY EMBOLISM IN TOTAL HIP REPLACEMENT SURGERY

## EXCLUSIVITY TERMS

## PATENT USE CODE

U-1 PREVENTION OF PREGNANCY  
 U-2 CYCLIC CONTROL  
 U-3 TREATMENT OF AMENORRHEA, DYSMENORRHEA, AND FUNCTIONAL UTERINE BLEEDING  
 U-4 TREATMENT OR PROPHYLAXIS OF ANGINA PECTORIS AND ARRHYTHMIA  
 U-5 TREATMENT OF HYPERTENSION  
 U-6 TREATING MAMMALS SUFFERING [FROM] ANXIETY  
 U-7 PROVIDING PREVENTION AND TREATMENT OF EMESIS AND NAUSEA IN MAMMALS  
 U-8 REDUCING INTRAVASCULAR PRESSURE IN MAMMALS  
 U-9 METHOD OF PRODUCING BRONCHODILATION  
 U-10 METHOD OF PRODUCING SYMPATHOMIMETIC EFFECTS  
 U-11 INCREASING CARDIAC CONTRACTILITY  
 U-12 TREATMENT OF BURNS  
 U-13 CONTROL OF EMESIS ASSOCIATED WITH ANY CANCER CHEMOTHERAPY AGENT  
 U-14 TREATMENT OF STRESS-INDUCED DEPRESSION  
 U-15 DIAGNOSTIC METHOD FOR DISTINGUISHING BETWEEN HYPOTHALMIC MALFUNCTIONS OR LESIONS IN HUMANS  
 U-16 TREATMENT OR PROPHYLAXIS OF CARDIAC DISORDERS  
 U-17 METHOD FOR TREATMENT OF HERPETIC INFECTIONS

PRESCRIPTION AND OTC DRUG PRODUCT  
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APPL/PROD	TRADE NAME; INGREDIENT NAME	PATENT NUMBER	PATENT EXPIRES	USE CODE	EXCLUS CODE	EXCLUS EXPIRES
18917 001	SECTRAL; ACEBUTOLOL HYDROCHLORIDE	3857952	DEC 31, 1993	U-4		
18917 003	SECTRAL; ACEBUTOLOL HYDROCHLORIDE	3857952	DEC 31, 1993	U-4		
19243 001	PROVENTIL; ALBUTEROL SULFATE	3705233	DEC 05, 1989		NDF	JAN 14, 1990
19243 002	PROVENTIL; ALBUTEROL SULFATE	3644353	FEB 22, 1989		NDF	JAN 14, 1990
>ADD>		3705233	DEC 05, 1989		NDF	JUL 13, 1990
>ADD>		3644353	FEB 22, 1989			
19383 001	PROVENTIL; ALBUTEROL SULFATE	3705233	DEC 05, 1989			
19621 001	VENTOLIN; ALBUTEROL SULFATE	3644353	FEB 22, 1989			
19353 001	ALFENTA; ALFENTANIL HYDROCHLORIDE	3705233	DEC 05, 1989			
18700 001	INOCOR; AMRINONE LACTATE	4167574	SEP 11, 1996	U-11	NCE	DEC 29, 1991
19389 001	BECONASE AQ; BECLMETHASONE DIPROPIONATE	4072746	FEB 07, 1995		NCE	JUL 31, 1994
19270 001	BETOPTIC; BETAXOLOL HYDROCHLORIDE	4252984	JUL 31, 1999		NP	JUL 27, 1990
18770 001	TORNALATE; BITOLTEROL MESYLATE	4336400	JUN 22, 1999	U-10	NCE	AUG 30, 1990
		4336400	JUN 22, 1999	U-9		
		4336400	JUN 22, 1999	U-10		
18644 001	WELLBUTRIN; BUPROPION HYDROCHLORIDE	3885046	MAY 20, 1994			
18644 002	WELLBUTRIN; BUPROPION HYDROCHLORIDE	3885046	MAY 20, 1994			
18644 003	WELLBUTRIN; BUPROPION HYDROCHLORIDE	3885046	MAY 20, 1994			
19215 001	FEMSTAT; BUTOCONAZOLE NITRATE	4078071	MAR 07, 1997			
18470 001	CIBACALCIN; CALCITONIN, HUMAN	RE32347	JUN 30, 1998		NCE	NOV 25, 1990
18057 001	PLATINOL; CISPLATIN	4177263	DEC 04, 1996		NCE	OCT 31, 1991
18057 002	PLATINOL; CISPLATIN	4177263	DEC 04, 1996		ODE	OCT 31, 1993
18057 003	PLATINOL-AQ; CISPLATIN	4177263	DEC 04, 1996			
19322 001	TEMOVATE; CLOBETASOL PROPIONATE	3721687	MAR 20, 1992			
19323 001	TEMOVATE; CLOBETASOL PROPIONATE	3721687	MAR 20, 1992			
12141 001	CYTOXAN; CYCLOPHOSPHAMIDE				NCE	DEC 27, 1990
12141 002	CYTOXAN; CYCLOPHOSPHAMIDE				NCE	DEC 27, 1990
12142 001	CYTOXAN; CYCLOPHOSPHAMIDE				I-63	APR 29, 1990
12142 002	CYTOXAN; CYCLOPHOSPHAMIDE				I-63	APR 29, 1990
12142 003	CYTOXAN; CYCLOPHOSPHAMIDE				I-63	APR 29, 1990
12142 004	CYTOXAN; CYCLOPHOSPHAMIDE				I-63	APR 29, 1990
12142 005	CYTOXAN; CYCLOPHOSPHAMIDE				I-63	APR 29, 1990
12142 006	LYOPHILIZED CYTOXAN; CYCLOPHOSPHAMIDE				I-63	APR 29, 1990
12142 007	LYOPHILIZED CYTOXAN; CYCLOPHOSPHAMIDE				I-63	APR 29, 1990
12142 008	LYOPHILIZED CYTOXAN; CYCLOPHOSPHAMIDE				I-63	APR 29, 1990
12142 009	LYOPHILIZED CYTOXAN; CYCLOPHOSPHAMIDE				I-63	APR 29, 1990
12142 010	LYOPHILIZED CYTOXAN; CYCLOPHOSPHAMIDE				I-63	APR 29, 1990
18885 002	EMBOLEX; DIHYDROERGOTAMINE MESYLATE	4402949	SEP 06, 2000			
12836 004	PERSANTINE; DIPYRIDAMOLE					
12836 005	PERSANTINE; DIPYRIDAMOLE					
17820 002	DOBUTREX; DOBUTAMINE HYDROCHLORIDE	3987200	OCT 19, 1993	U-11		

APPL/PROD	TRADE NAME; INGREDIENT NAME	PATENT NUMBER	PATENT EXPIRES	USE CODE	EXCLUS CODE	EXCLUS EXPIRES
19386 002	BREVIBLOC; ESMOLOL HYDROCHLORIDE	4593119	JUN 03, 2003		NCE	DEC 31, 1991
16672 001	OVRAL; ETHINYL ESTRADIOL	4387103	JUN 07, 2000	U-16		
		3666858	MAY 30, 1989	U-1		
		3666858	MAY 30, 1989	U-2		
		3666858	MAY 30, 1989	U-3		
16805 001	OVRAL-28; ETHINYL ESTRADIOL	3666858	MAY 30, 1989	U-1		
		3666858	MAY 30, 1989	U-2		
		3666858	MAY 30, 1989	U-3		
17612 001	LO/OVRAL; ETHINYL ESTRADIOL	3666858	MAY 30, 1989	U-1		
		3666858	MAY 30, 1989	U-2		
		3666858	MAY 30, 1989	U-3		
17802 001	LO/OVRAL-28; ETHINYL ESTRADIOL	3666858	MAY 30, 1989	U-1		
		3666858	MAY 30, 1989	U-2		
		3666858	MAY 30, 1989	U-3		
18668 001	NORDETTE-21; ETHINYL ESTRADIOL	3666858	MAY 30, 1989	U-1		
		3666858	MAY 30, 1989	U-2		
		3666858	MAY 30, 1989	U-3		
18782 001	NORDETTE-28; ETHINYL ESTRADIOL	3666858	MAY 30, 1989	U-1		
		3666858	MAY 30, 1989	U-2		
		3666858	MAY 30, 1989	U-3		
19190 001	TRIPHASIL-28; ETHINYL ESTRADIOL	3666858	MAY 30, 1989	U-1		
		3957982	MAY 18, 1993	U-1		
		3666858	MAY 30, 1989	U-2		
		3666858	MAY 30, 1989	U-3		
19192 001	TRIPHASIL-21; ETHINYL ESTRADIOL	3666858	MAY 30, 1989	U-1		
		3666858	MAY 30, 1989	U-2		
		3666858	MAY 30, 1989	U-3		
19545 001	DIDRONEL; ETIDRONATE DISODIUM	4254114	MAR 03, 1998			
		4216211	AUG 05, 1997			
		4137309	JAN 30, 1996			
		3683080	AUG 08, 1989			
		4283408	AUG 11, 1998			
		4005209	JAN 25, 1996			
		4005209	JAN 25, 1996			
19527 001	PEPCID; FAMOTIDINE	3793457	FEB 19, 1991			
18830 001	TAMBOCOR; FLECAINIDE ACETATE	3755427	AUG 28, 1990			
18830 002	TAMBOCOR; FLECAINIDE ACETATE	4110438	AUG 29, 1995			
19415 002	METRODIN; FLUMAZENIL	3947569	MAR 30, 1993			
19404 001	OCUFEN; FLURBIPROFEN SODIUM	4110438	AUG 29, 1995			
18123 001	FACTREL; GONADORELIN HYDROCHLORIDE	3947569	MAR 30, 1993	U-14		
18123 002	FACTREL; GONADORELIN HYDROCHLORIDE	3947569	MAR 30, 1993	U-15		
18123 003	FACTREL; GONADORELIN HYDROCHLORIDE	4110438	AUG 29, 1995	U-14		
		4110438	AUG 29, 1995	U-15		
		3947569	MAR 30, 1993	U-14		
		3947569	MAR 30, 1993	U-15		

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APPL/PROD	TRADE NAME; INGREDIENT NAME	PATENT NUMBER	PATENT EXPIRES	USE CODE	EXCLUS CODE	EXCLUS EXPIRES
18587 001	WYTENSIN; GUANABENZ ACETATE	3658993	APR 25, 1989	U-5	NCE	SEP 07, 1992
18587 002	WYTENSIN; GUANABENZ ACETATE	3658993	APR 25, 1989	U-5	NCE	SEP 07, 1992
18587 003	WYTENSIN; GUANABENZ ACETATE	3658993	APR 25, 1989	U-5	NCE	SEP 07, 1992
>ADD>						
>ADD>						
18872 001	VISKAZIDE; HYDROCHLOROTHIAZIDE	4066755	JAN 03, 1995		NCE	SEP 03, 1992
18872 002	VISKAZIDE; HYDROCHLOROTHIAZIDE	4012444	MAR 15, 1994		NCE	SEP 03, 1992
19046 001	NORMOZIDE; HYDROCHLOROTHIAZIDE	4066755	JAN 03, 1995		NCE	SEP 03, 1992
19046 002	NORMOZIDE; HYDROCHLOROTHIAZIDE	4012444	MAR 15, 1994		NC	APR 06, 1990
19046 003	NORMOZIDE; HYDROCHLOROTHIAZIDE	4066755	JAN 03, 1995		NC	APR 06, 1990
19046 004	NORMOZIDE; HYDROCHLOROTHIAZIDE	4012444	MAR 15, 1994		NC	APR 06, 1990
19174 001	TRANDATE-HCT; HYDROCHLOROTHIAZIDE	4066755	JAN 03, 1995		NC	APR 06, 1990
19174 002	TRANDATE-HCT; HYDROCHLOROTHIAZIDE	4012444	MAR 15, 1994		NC	APR 10, 1990
19174 003	TRANDATE-HCT; HYDROCHLOROTHIAZIDE	4066755	JAN 03, 1995		NC	APR 10, 1990
19174 004	TRANDATE-HCT; HYDROCHLOROTHIAZIDE	4012444	MAR 15, 1994		NC	APR 10, 1990
19571 001	HUMULIN U; INSULIN ZINC SUSP EXTENDED BIOSYNTHETIC HUMAN				NC	APR 10, 1990
19571 002	HUMULIN U; INSULIN ZINC SUSP EXTENDED BIOSYNTHETIC HUMAN				NP	JUN 10, 1990
18956 001	OMNIPAQUE 180; IOHEXOL	4396597	JUL 14, 1998		NP	JUN 10, 1990
18956 002	OMNIPAQUE 240; IOHEXOL	4250113	DEC 26, 1999		I-65	MAY 12, 1990
18956 003	OMNIPAQUE 300; IOHEXOL	4396597	JUL 14, 1998		NCE	DEC 26, 1990
18956 004	OMNIPAQUE 350; IOHEXOL	4250113	DEC 26, 1999		I-65	MAY 12, 1990
18735 001	ISOVUE 200; IOPAMIDOL	4396597	JUL 14, 1998		NCE	DEC 26, 1990
		4250113	DEC 26, 1999		I-65	MAY 12, 1990
		4001323	JAN 04, 1996		NCE	DEC 31, 1990
					NR	JUL 07, 1990
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>ADD>						
18735 002	ISOVUE-300; IOPAMIDOL	4001323	JAN 04, 1996		I-57	JUL 07, 1990
18735 003	ISOVUE-370; IOPAMIDOL	4001323	JAN 04, 1996		NCE	DEC 31, 1990
18735 004	ISOVUE-M 300; IOPAMIDOL	4001323	JAN 04, 1996		NCE	DEC 31, 1990
13295 002	CONRAY-43; IOTHALAMATE MEGLUMINE				NCE	DEC 31, 1990
					I-54	DEC 18, 1989

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APPL/PROD	TRADE NAME; INGREDIENT NAME	PATENT NUMBER	PATENT EXPIRES	USE CODE	EXCLUS CODE	EXCLUS EXPIRES
18905 002	HEXABRIX; IOXAGLATE MEGLUMINE	4094966	JUN 13, 1995		I-54	OCT 22, 1989
		4065554	DEC 27, 1994		I-36	OCT 22, 1989
		4065553	DEC 27, 1994		I-6	OCT 22, 1989
		4014986	MAR 29, 1996		NCE	JUL 26, 1990
					I-55	OCT 22, 1989
					I-56	OCT 22, 1989
					I-57	OCT 22, 1989
					I-58	OCT 22, 1989
					I-59	OCT 22, 1989
					I-60	OCT 22, 1989
					I-61	OCT 22, 1989
		3641127	FEB 08, 1991		NCE	JAN 09, 1991
		3641127	FEB 08, 1991		NCE	JAN 09, 1991
		3641127	FEB 08, 1991		NCE	JAN 09, 1991
		4005063	JAN 25, 1996		NCE	JAN 09, 1991
		3497599	JAN 26, 1988	U-12		APR 09, 1990
		4137300	JAN 30, 1996			
		4536386	AUG 20, 2002	U-13		APR 30, 1992
		4536386	AUG 20, 2002	U-13		
		3998790	DEC 21, 1993		I-66	MAY 28, 1990
		3998790	DEC 21, 1993		NS	MAY 28, 1990
		3954872	MAY 04, 1995		I-64	JUN 27, 1989
		3954872	MAY 04, 1995		I-64	JUN 27, 1989
		3954872	MAY 04, 1995		NCE	DEC 30, 1990
		3954872	MAY 04, 1995		NCE	DEC 30, 1990
		4280957	MAY 04, 1995		NCE	DEC 30, 1990
		4472393	SEP 18, 2001		NCE	DEC 20, 1990
		4472393	SEP 18, 2001		NCE	APR 30, 1992
		4472393	SEP 18, 2001		NCE	APR 30, 1992
		4087547	MAY 02, 1995	U-8	NDF	MAY 29, 1990
		4087545	MAY 02, 1995	U-7		
		3928598	DEC 23, 1992	U-6		
		3928009	NOV 18, 1992			
		3998966	DEC 21, 1993		NCE	DEC 26, 1990
		3904682	SEP 09, 1992		I-62	MAR 23, 1990
		3998966	DEC 21, 1993		D-13	MAR 23, 1990
		3904682	SEP 09, 1992		I-62	MAR 23, 1990
		3998966	DEC 21, 1993		D-13	MAR 23, 1990
		3904682	SEP 09, 1992		I-62	MAR 23, 1990
		4009197	SEP 09, 1992		D-13	MAR 23, 1990
		4001301	SEP 09, 1992			
		3998966	DEC 21, 1993			
		3904682	SEP 09, 1992		NDF	MAR 23, 1990
		3904682	SEP 09, 1992			
17581 002	NAPROSYN; NAPROXEN					
17581 003	NAPROSYN; NAPROXEN					
17581 004	NAPROSYN; NAPROXEN					
18965 001	NAPROSYN; NAPROXEN					

>ADD>

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APPL/PROD	TRADE NAME; INGREDIENT NAME	PATENT NUMBER	PATENT EXPIRES	USE CODE	EXCLUS CODE	EXCLUS EXPIRES
19384 002	NOROXIN; NORFLOXACIN	4639458	JAN 27, 2004			
17031 001	OVRETTE; NORGESTREL	3666858	MAY 30, 1989	U-1		
		3666858	MAY 30, 1989	U-2		
		3666858	MAY 30, 1989	U-3		
18553 004	INDERAL LA; PROPRANOLOL HYDROCHLORIDE	4138475	FEB 06, 1996			
19536 001	INDERAL; PROPRANOLOL HYDROCHLORIDE	4600708	JUL 15, 2003		D-7	OCT 31, 1989
18708 003	DORMALIN; QUAZEPAM	3920818	NOV 18, 1992			
		3845039	OCT 29, 1991			
		4211771	JUL 08, 1999			
18859 001	VIRAZOLE; RIBAVIRIN				NCE	DEC 27, 1990
19518 001	EXTRA-STRENGTH AIM; SODIUM MONOFLUOROPHOSPHATE				NCE	DEC 31, 1990
19518 002	EXTRA-STRENGTH AIM; SODIUM MONOFLUOROPHOSPHATE				NS	AUG 06, 1989
19107 001	PROTROPIN; SOMATREM				NS	AUG 06, 1989
19640 004	HUMATROPE; SOMATROPIN, BIOSYNTHETIC				NCE	OCT 17, 1990
18217 001	SUPROL; SUPROFEN	4658021	APR 14, 2004		ODE	MAR 08, 1994
18963 001	CHOLETEC; TECHNITIUM TC-99M MEBROFENIN KIT				NCE	DEC 24, 1990
18682 001	TROSYD; TIOCONAZOLE	4035376	JUL 12, 1996		NCE	JAN 21, 1992
19355 001	VAGISLAT; TIOCONAZOLE	4418208	NOV 29, 2000			
14103 003	ONCOVIN; VINCRISTINE SULFATE	4661493	APR 28, 2004	U-17		
19655 001	RETROVIR; ZIDOVUDINE	4661493	APR 28, 2004	U-17		
		4619935	OCT 28, 2003		ODE	MAR 19, 1994
					NCE	MAR 19, 1992

DRUG PRODUCTS APPROVED UNDER SECTION 505 OF THE ACT  
 BY THE DIVISION OF BLOOD AND BLOOD PRODUCTS LIST  
 PATENT AND EXCLUSIVITY DATA

APPL/PROD	TRADE NAME; INGREDIENT NAME	PATENT NUMBER	PATENT EXPIRES	USE CODE	EXCLUS CODE	EXCLUS EXPIRES
83715 001	PROMIT; DEXTRAN 1 IN SODIUM CHLORIDE 0.6%	4201772	AUG 17, 1998		NCE	OCT 30, 1989
841207 001	PENTASpan; PENTASTARCH 10% IN SODIUM CHLORIDE 0.9%				ODE	MAY 19, 1994



**SUBSCRIPTION FORM**  
**APPROVED DRUG PRODUCTS**  
**WITH**  
**THERAPEUTIC EQUIVALENCE EVALUATIONS**  
**7TH EDITION (1987)**

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Check/money order enclosed for \$ \_\_\_\_\_

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AUTHORIZING  
SIGNATURE:

DATE:

DESCRIPTION	QUANTITY	UNIT PRICE	TOTAL PRICE
The 7th Edition is published in March 1987. Subscription includes the Approved Drug Products publication and monthly Cumulative Supplements.  DOMESTIC (Stock No. 917-001-00000-6)		@ \$86.00	\$
FOREIGN (Stock No. 917-001-00000-6)		@ \$107.50	\$
ENTER TOTAL			\$