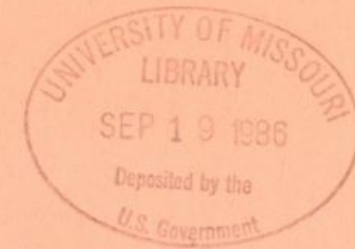


MED
HE 20.4210
985/supp.11

CUMULATIVE
SUPPLEMENT 11
AUG'85-JUL'86

APPROVED
DRUG PRODUCTS
WITH
THERAPEUTIC EQUIVALENCE EVALUATIONS
6TH EDITION



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

APPROVED DRUG PRODUCTS

with

THERAPEUTIC EQUIVALENCE EVALUATIONS

6TH EDITION

CUMULATIVE SUPPLEMENT

JULY 1986

CONTENTS

PAGE

A. INTRODUCTION

1. How to Use the Cumulative Supplement	v
2. Applicant (Name) Changes	vi
3. Prednisone Bioequivalence	vii
4. OTC Drug Products	viii
5. Products Requiring Revised Labeling for Full Approval	ix
6. Injectable Product Package Size Designation	x
7. Report of Counts for the Prescription Drug Product List	xi

B. DRUG PRODUCT LISTS

1. Prescription Drug Product List	1
2. OTC Drug Product List	40
3. Drug Products Approved Under Section 505 of the Act by the Division of Blood and Blood Products List	42

C APPENDICES

1. Orphan Drug Products with Exclusive Approval	44
2. List of Drug Products Which Must Demonstrate <i>in vivo</i> Bioavailability Only if Product Fails to Achieve Adequate Dissolution	48
3. Biopharmaceutic Guidance Availability List	49
4. ANDA Suitability Petitions	51
5. Exclusivity Terms	75
6. Prescription and OTC Drug Product Patent and Exclusivity Data	78

A. INTRODUCTION

1. How to Use the Cumulative Supplement
2. Applicant (Name) Changes
3. Prednisone Bioequivalence
4. OTC Drug Products
5. Products Requiring Revised Labeling for Full Approval
6. Injectable Product Package Size Designation
7. Report of Counts for the Prescription Drug Product List

APPROVED DRUG PRODUCTS

with

THERAPEUTIC EQUIVALENCE EVALUATIONS

6th EDITION

CUMULATIVE SUPPLEMENT

JULY 1986

A. INTRODUCTION

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, 6th Edition (the List). The List is comprised of three drug product lists: The Prescription Drug Product list, the OTC Drug Product list, and the Drug Products Approved Under Section 505 of the Act by the Division of Blood and Blood Products list. 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.

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.

Information in the Cumulative Supplement follows the format of the drug product lists. 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.)

Drug product information is provided in each Cumulative Supplement for completeness to assist in locating the proper place in the drug product lists for the revision. [Strength(s) which already exist in the publication will not be repeated for context.] A page number in parentheses, located to the right of the ingredient(s), refers to the related page in the drug product lists. The effective (marketing) date (the date a product may be marketed), when appropriate, will appear to the left of the approval date.

Additions to the drug product lists and the Appendices are indicated by new information in the Cumulative Supplement. Additions new to the current Cumulative Supplement are indicated by the symbol >ADD> to the left of the line on which new information exists. The >ADD> symbol is dropped in subsequent Cumulative Supplements for that item.

A newly approved product is identified by the lozenge (**) to the right of its strength. This identifier remains throughout all Cumulative Supplements for this edition.

Deletions from the drug product lists and the Appendices are indicated by overstruck print in the Cumulative Supplement. Deletions new to the current Cumulative Supplement are indicated by the symbol >~~DLT~~< (DELETE) to the left of the line containing the overstruck print. The symbol is dropped in subsequent Cumulative Supplements for that item.

Products discontinued from marketing will be flagged in this Cumulative Supplement with the "e" symbol to designate their non-marketed status until such time that the Agency is notified that they are being marketed.

The Appendices of the Cumulative Supplement provide, among other things, updated information required of the Agency by the "Drug Price Competition and Patent Term Restoration Act of 1984."

2. 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 Special Notes 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. The current list of applicant holder changes follows.

APPLICANT (NAME) CHANGES

<u>Former Applicant (Name)</u>	<u>New Applicant (Name)</u>	<u>New Abbreviated Name</u>
VITARINE/PHOENIX	VITARINE PHARMACEUTICALS, INC	VITARINE PHARMS
DRUMMER/PHOENIX	VITARINE PHARMACEUTICALS, INC	VITARINE PHARMS
INVENEX LABS/LIFE	LYPHOMED, INC	LYPHOMED
ONEAL JONES&FELDMAN	FOREST PHARMACEUTICALS, INC SUBSIDIARY OF FOREST LABORATORIES, INC	FOREST PHARMS/FOREST

(continued)

APPLICANT (NAME) CHANGES

(continued)

<u>Former Applicant (Name)</u>	<u>New Applicant (Name)</u>	<u>New Abbreviated Name</u>
AM MCGAW/AM HOSP	KENDALL MCGAW LABORATORIES, INC	KENDALL MCGAW LABS
IVES LABS/AMHO	WYETH LABORATORIES, INC DIVISION OF AMERICAN HOME PRODUCTS CORP	WYETH LABS/AMHO
REID PROVIDENT LABS AND ROWELL LABORATORIES	REID-ROWELL	REID-ROWELL
BAY LABORATORIES	MY-K LABORATORIES, INC	MY-K LABS

3. 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, Cmax, Tmax) 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 Appendix 3 of this Supplement for available guidance from the Division of Bioequivalence.)

4. 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).

Dexbrompheniramine Maleate Pseudoephedrine Sulfate Tablet; Oral	2mg 60mg
Pseudoephedrine HCl Triprolidine HCl Tablet or Capsule; Oral	60mg 2.5mg
Pseudoephedrine HCl Triprolidine HCl Syrup; Oral	30mg/5ml 1.25mg/5ml
Triprolidine HCl Syrup; Oral	1.25mg/5ml
Triprolidine HCl Tablet; Oral	2.5mg

5. 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>
isosorbide dinitrate	AUG 3, 1984 (49 FR 31151)
nandrolone decanoate	JUL 15, 1983 (48 FR 32395)
neomycin sulfate with either: dexamethasone sodium phosphate, fluocinolone acetonide, flurandrenolide,	MAR 26, 1984 (49 FR 11888)
hydrocortisone, or methylprednisolone acetate	
[topical anti-infectives for dermatologic use]	
neomycin sulfate, polymyxin B sulfate, bacitracin zinc, and hydrocortisone [topical ointment]	MAY 4, 1984 (49 FR 19147)
nitroglycerin (capsule, controlled release; oral)	SEP 7, 1984 (49 FR 35428)
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)
sulfamilamide and aminacrine	AUG 22, 1983 (48 FR 38097)
tranylcypromine sulfate	MAR 22, 1984 (49 FR 10708)

6. INJECTABLE PRODUCT PACKAGE SIZE DESIGNATION

When a new drug product (usually injectable) is approved for the same concentration but a different package size than the listed drug, and it has received a period of exclusivity by the Agency, the product will appear as single source (no therapeutic equivalence code displayed) and the potency will reflect the unique package size. Once the period of exclusivity has ended, the product will then conform to the standard ADP format of reflecting "collapsed" package sizes. The current standard is to display package sizes of all small volume parenterals within an NDA as a per ml concentration and large volume parenterals as per 100ml.

7. 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 July '85, 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 LIST

A. COUNTS CUMULATIVE BY QUARTERS

<u>CATEGORIES COUNTED</u>	<u>JULY '85</u>	<u>(BASELINE)</u>	<u>OCT '85</u>	<u>JAN '86</u>	<u>APR '86</u>
DRUG PRODUCTS LISTED	8048	8230	8515	8683	
SINGLE SOURCE	2096 (26.0%)	2100 (25.5%)	2144 (25.1%)	2138 (24.6%)	
MULTISOURCE (1)	5952 (74.0%)	6130 (74.5%)	6371 (74.9%)	6545 (75.4%)	
THERAPEUTICALLY EQUIVALENT	4864 (60.5%)	5034 (61.2%)	5263 (61.8%)	5422 (62.5%)	
NOT THERAPEUTICALLY EQUIVALENT	1054 (13.1%)	1058 (12.9%)	1070 (12.6%)	1068 (12.3%)	
EXCEPTIONS (2)	34 (0.4%)	38 (0.4%)	38 (0.5%)	55 (0.6%)	
NEW MOLECULAR ENTITIES APPROVED	-	5	21	1	
NUMBER OF APPLICANTS	306	313	322	324	

B. ACTIVITY FOR SUPPLEMENT NUMBER 11

	<u>MAY '86</u>	<u>JUN '86</u>	<u>JUL '86</u>	<u>CUMULATIVE</u>
DRUG PRODUCTS ADDED:				
NEWLY APPROVED	60	57	59	84
DESI EFFECTIVE	0	0	5	84
REMARKETED	3	3	0	84
DRUG PRODUCTS REMOVED:				
WITHDRAWN APPROVAL	0	0	0	0
RX TO OTC SWITCH	0	0	0	0
NET GAIN IN DRUG PRODUCTS				
SINGLE SOURCE PRODUCTS APPROVED	60	59	59	84
MULTISOURCE DRUG PRODUCTS APPROVED	5	9	5	84
NEW MOLECULAR ENTITIES APPROVED:	55	50	79	164
AS THE ENTITY	0	0	0	0
AS A SALT, ESTER OR DERIVATIVE	0	0	0	0
OF THE ENTITY	0	0	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 PAGE I-8 OF THE LIST)

B. DRUG PRODUCT LISTS

1. Prescription Drug Product List
2. OTC Drug Product List
3. Drug Products Approved Under Section 505 of the Act
by the Division of Blood and Blood Products List

PRESRIPTIO^N: DRUG PRODUCT LIST
6TH EDITION
CUMULATIVE SUPPLEMENT NUMBER 11 / AUG '85 - JUL '86

1

ACETAMINOPHEN (PAGE 3-1)

INJECTABLE; INJECTION
INJECTAPAP

MCNEIL PHARM

100MG/MLX

N17785 001

MAR 07, 1986

ACETAMINOPHEN; BUTALBITAL (PAGE 3-1)

CAPSULE; ORAL
BANCAP

FOREST PHARM/FOREST

325MG;50MGX

N88889 001

JAN 16, 1986

CAPSULE; ORAL
SEDAPAP-10

MAYRAND

650MG;50MGX

N88944 001

OCT 17, 1985

ACETAMINOPHEN; BUTALBITAL; CAFFFEINE (PAGE 3-1)

CAPSULE; ORAL
ACETAMINOPHEN, BUTALBITAL, AND CAFFEINE

AB MIKART

325MG;50MG;40MGX

N89007 001

MAR 17, 1986

CAPSULE; ORAL
COMPALE

REID-ROWE LL

356.4MG;20MG;16MGX

N88584 001

MAR 04, 1986

SYNALGOS-DC-A

HYETH LABS/AMHO

356.4MG;20MG;16MGX

N89166 001

MAY 14, 1986

ACETAMINOPHEN; CODEINE PHOSPHATE (PAGE 3-1)

TABLET; ORAL
ACETAMINOPHEN AND CODEINE

VITARINE

300MG;15MG

200MG;20MG

300MG;20MG

650MG;20MG

MAR 03, 1986

AA

CAPSULE; ORAL
ACETAMINOPHEN AND HYDROCODONE BITARTRATE

DM GRAHAM LABS

500MG;5MG

AA

BAHCAP HC

FOREST PHARM/FOREST

500MG;5MG

AA

/DR. JONES/FECHAN//500MG;5MG/

/MAR 11, 1983/

N89006 001

AUG 09, 1985

N87961 001

MAR 17, 1983

/JUN 16, 1983/

/JUN 16, 1983/

/JUN 16, 1983/

ACETAMINOPHEN; HYDROCODONE BITARTRATE (PAGE 3-3)

ACYCLOVIR (PAGE 3-5)

CAPSULE; ORAL
HYDROCODEONE
BITARTRATE AND ACETAMINOPHEN

CAPSULE; ORAL
HYDROCODEONE BITARTRATE AND ACETAMINOPHEN
MIKART
500MG/5MG

**TABLET; ORAL
DURADYNE DH**

AA	FOREST PHARM/FOREST	<u>500MG; 5MG</u>	N87809 001 MAR 17, 1983
> <u>ADD</u> >	HYDROCODONE BITARTRATE AND ACETAMINOPHEN	<u>500MG; 5MG</u>	N89271 001 JUL 16, 1986
> <u>ADD</u> >	MIKART	<u>500MG; 5MG</u>	N88871 001 MAY 15, 1986
> <u>ADD</u> >	HORSETT	<u>500MG; 5MG</u>	N88871 001 MAY 15, 1986
AA	HOLLOWAY PHARMS	<u>500MG; 5MG</u>	

ACETAMINOPHEN; PROPOXYPHENE NAPSYLATE (PAGE 3-3)
TABLET; ORAL
PROPOXYPHENE NAPSYLATE AND ACETAMINOPHEN

BARR LABORATORIES	<u>650MG;100MG</u>	N70615 001 MAR 21, 1986
	<u>650MG;100MG</u>	N70771 001 MAR 21, 1986
	<u>650MG;100MG</u>	N70775 001 MAR 21, 1986
CORD LABORATORIES	<u>650MG;100MG</u>	N70443 001 JAN 23, 1986
	<u>650MG;100MG</u>	N70732 001 JAN 03, 1986
LEMMON		N70146 001 AUG 02, 1985
ZENITH LABORATORIES	<u>650MG;100MG</u>	

CETAZOLAMIDE (PAGE 3-4)

TABLET; ORAL
ACETAZOLAMIDE
DANBURY PHARMACEUTICALS
250MG

CETIC ACTIO - FILE ACTAI (PAGE 3-4)

**ACETIC ACID GLACIAL
SOLUTION/DROPS; OTIC**

ACYCLOVIR (PAGE 3-5)

CAPSULE; ORAL
ZOVIRAX

N18828 001
JAN 28, 1985
N 25-1985

ALBUTEROL SULFATE (PAGE 3-6)

N17853 001	N19112 001
MAY 07, 1982	JUL 10, 1986
N17853 002	N19112 002
MAY 07, 1982	JUL 10, 1986

ANSWER (PAGE 3-6)

N70466 001
DEC 24, 1985
N70467 001
DEC 24, 1985
N70268 001
DEC 31, 1985
N70269 001
DEC 31, 1985
N70150 001
DEC 10, 1985
N70147 001
DEC 10, 1985
N70579 001
APR 14, 1986
N70580 001
APR 14, 1986

AMILORIDE HYDROCHLORIDE (PAGE 3-7)

TABLET; ORAL
ANTILORIDE HCL
PAR PHARMACEUTICAL
5MGR
MEDAMOR
MC 2127 MEDICK
AB AB

ACYCLOVIR (PAGE 3-5)

CAPSULE; ORAL
ZOVIRAX
BURROUGHS WELLCOME
200MG
N18828 001
1 JAN 88, 1985

<u>ALBUTEROL SULFATE</u>	(PAGE 3-6)
<u>TABLET; ORAL</u>	
<u>PROVENTIL</u>	
SCHERING	
<u>AB</u>	<u>EQ 2MG BASE</u>
<u>AB</u>	<u>EQ 4MG BASE</u>

<u>VENTOLIN</u>	<u>EQ 2MG BASE</u>	N19112 001
<u>GLAXO</u>		JUL 10, 1986
<u>AB</u>	<u>EQ 4MG BASE</u>	N19112 002
		JUL 10, 1986
<u>AB</u>		

NIOBIRINDI (PAGE 3-6)

TABLET; ORAL <u>ALLOPURINOL</u>		<u>100MG</u>	<u>100MG</u>	<u>100MG</u>	<u>100MG</u>	<u>100MG</u>	<u>100MG</u>	<u>100MG</u>	<u>100MG</u>
AB	BARR LABORATORIES								
AB	CORD LABORATORIES								
AB	PAR PHARMACEUTICAL								
AB	PUREPAC/KALIPHARMA								
AB		NOV 30, 1988 : DEC 24, 1985	NOV 30, 1988 : DEC 24, 1985	/NOV '86; '1986 ./. DEC 31, 1985					
AB		N70466 001	N70467 001	N70268 001	N70269 001	N70150 001	N70147 001	N70579 001	N70580 001
AB		APR 14, 1986	APR 14, 1986	APR 10, 1985	APR 14, 1986				

AMILORIDE HYDROCHLORIDE (PAGE 3-7)

TABLET; ORAL
ANTILORIDE HCL
 PAR PHARMACEUTICAL
51134
MEDAMOR
 MC 2127 MEDICK
 AB AB

AMINO ACIDS (PAGE 3-7)

INJECTABLE; INJECTION

AMINOSYN-HBC 7% IN PLASTIC CONTAINER
ABBOTT LABORATORIES 7/24N19400 001
JUL 23, 1985AMINOSYN-PF 7%
ABBOTT LABORATORIES 7/24N19398 001
SEP 06, 1985AMINOSYN II 3.5%
ABBOTT LABORATORIES 3.5/24N19438 001
APR 03, 1986AMINOSYN II 5%
ABBOTT LABORATORIES 5/24N19438 002
APR 03, 1986AMINOSYN II 7%
ABBOTT LABORATORIES 7/24N19438 003
APR 03, 1986AMINOSYN II 8.5%
ABBOTT LABORATORIES 8.5/24N19438 004
APR 03, 1986AMINOSYN II 10%
ABBOTT LABORATORIES 10/24N19438 005
APR 03, 1986AMINO ACIDS; CALCIUM ACETATE; GLYCERIN; MAGNESIUM ACETATE;
PHOSPHORIC ACID; POTASSIUM CHLORIDE; SODIUM ACETATE; SODIUM
CHLORIDE (PAGE 3-8)

INJECTABLE; INJECTION

/PERIPHRAMINE/
PROCALAMINE
KENDALL MCGRAW LABS3%;20MG/100ML;3GM/100ML;54MG/100ML;
41MG/100ML;150MG/100ML;200MG/100ML;
120MG/100ML
N18582 001
MAY 08, 1982INJECTABLE; INJECTION
AMINOSYN II 3.5% M
ABBOTT LABORATORIES3.5%;32MG/100ML;128MG/100ML;
222MG/100ML;49MG/100ML
N19437 007
APR 03, 1986AMINO ACIDS; MAGNESIUM ACETATE; POTASSIUM ACETATE; SODIUM
CHLORIDE; SODIUM PHOSPHATE, DIBASIC (PAGE 3-9)

INJECTABLE; INJECTION

AMINOSYN II 7% W ELECTROLYTES
ABBOTT LABORATORIES7%;102MG/100ML;45MG/100ML;
522MG/100ML;410MG/100ML
N19437 006
APR 03, 1986AMINO ACIDS; MAGNESIUM CHLORIDE; POTASSIUM CHLORIDE;
POTASSIUM PHOSPHATE, DIBASIC; SODIUM CHLORIDE (PAGE 3-9)INJECTABLE; INJECTION
AMINOSYN II 7% W ELECTROLYTES
ABBOTT LABORATORIES7%;102MG/100ML;45MG/100ML;
522MG/100ML;410MG/100ML
N19437 005
APR 03, 1986AMINO ACIDS; MAGNESIUM CHLORIDE; POTASSIUM CHLORIDE;
POTASSIUM PHOSPHATE, DIBASIC; SODIUM CHLORIDE (PAGE 3-9)INJECTABLE; INJECTION
AMINOSYN II 8.5% W ELECTROLYTES
ABBOTT LABORATORIESN19437 001
APR 03, 1986AMINOSYN II 10% W ELECTROLYTES
ABBOTT LABORATORIESN19437 004
APR 03, 1986AMINOSYN II 10% W ELECTROLYTES
ABBOTT LABORATORIESN19437 004
APR 03, 1986

AMINO ACIDS; MAGNESIUM CHLORIDE; POTASSIUM PHOSPHATE, DIBASIC;

SODIUM ACETATE; SODIUM CHLORIDE (PAGE 3-9)

INJECTABLE; INJECTION
/TRAVASOL, H, J, P, W, ELEC/N19438 001
APR 03, 1986TRAVASOL 3.5% W ELECTROLYTES
TRAVENOL LABS3.5%;51MG/100ML;131MG/100ML;
218MG/100ML;35MG/100ML
N17493 003

AMINOACAPROIC ACID (PAGE 3-9)

INJECTABLE; INJECTION
AMINOACAPROIC ACID
LYPHOMEDN170522 001
APR 03, 1986AMINOACAPROIC ACID
QUAD PHARMSN170694 001
APR 03, 1986

AMINOPHYLLINE (PAGE 3-10)

TABLET; ORAL
AMINOPHYLLINE
CORD LABORATORIESN185262 002
APR 03, 1986AMINOPHYLLINE
/COPD LABORATORIES/N18972 001
APR 03, 1986

AMITRIPTYLINE HYDROCHLORIDE (PAGE 3-11)

TABLET; ORAL
CORDARONE
IVES LABS/AMHON18972 001
APR 03, 1986

AMITRIPTYLINE HYDROCHLORIDE (PAGE 3-14)

TABLET; ORAL
TRIAVIL 4-10
/BP/ /MS&P/MERCK/
MS&D/MERCK/101G/44G/
10MG;4MG
BP/14715'661/
N14715 003

RX DRUG PRODUCT LIST / CUMULATIVE SUPPLEMENT NUMBER 11 / AUG '85 - JUL '86

4

AMOXICILLIN (PAGE 3-15)

CAPSULE; ORAL
AMOXICILLIN
AB LABORATORIOS ATRAI 250MG
AB 500MG
AB UTI-MON
AB 1/3 PARKE-DAVIS/M-L 250MG
AB 500MG
AB POWDER FOR RECONSTITUTION; ORAL
UTI-MON
AB PARK-E-DAVIS/M-L 125MG/5ML
AB 250MG/5ML

AMPICILLIN SODIUM (PAGE 3-17)

INJECTABLE; INJECTION
AMPICILLIN SODIUM
AP ELI LILLY EQ 2GM BASE/VIAL
AP ELKINS-SINN/AHROBINS EQ 125MG BASE/VIAL
AP EQ 250MG BASE/VIAL
AP EQ 500MG BASE/VIAL
AP EQ 1GM BASE/VIAL
AP EQ 2GM BASE/VIAL
AP EQ 10GM BASE/VIAL
AP TOTACELLIN-H
AP BEECHAM LABS./BEECHAM EQ 10GM BASE/VIAL

ASCORBIC ACID; BIOTIN; CYANOCOBALAMIN; DEXPANTHENOL;
ERGOCALCIFEROL; FOLIC ACID; NIACINAMIDE; PYRIDOXINE;
RIBOFLAVIN PHOSPHATE SODIUM; THIAMINE;
VITAMIN A; VITAMIN E (PAGE 3-19)

INJECTABLE; INJECTION
M.V.I.-12 LYOPHILIZED
USV PHARMACEUTICAL 100MG/VIAL; 0.06MG/VIAL; 0.05MG/VIAL;
15MG/VIAL; 200 IU/VIAL; 0.4MG/VIAL;
40MG/VIAL; 4MG/VIAL; 3,200 IU/VIAL; 3.6MG/VIAL;
3MG/VIAL; 3,200 IU/VIAL; 10 IU/VIAL
N18933 002
AUG 08, 1985

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

AB LABORATORIOS ATRAI 250MG
AB 500MG
AB UTI-MON
AB 1/3 PARKE-DAVIS/M-L 250MG
AB 500MG
AB POWDER FOR RECONSTITUTION; ORAL
UTI-MON
AB PARK-E-DAVIS/M-L 125MG/5ML
AB 250MG/5ML

AMPICILLIN SODIUM (PAGE 3-17)

INJECTABLE; INJECTION
AMPICILLIN SODIUM
AP ELI LILLY EQ 2GM BASE/VIAL
AP ELKINS-SINN/AHROBINS EQ 125MG BASE/VIAL
AP EQ 250MG BASE/VIAL
AP EQ 500MG BASE/VIAL
AP EQ 1GM BASE/VIAL
AP EQ 2GM BASE/VIAL
AP EQ 10GM BASE/VIAL
AP TOTACELLIN-H
AP BEECHAM LABS./BEECHAM EQ 10GM BASE/VIAL

INJECTABLE; INJECTION
AMPICILLIN SODIUM
AP ELI LILLY EQ 2GM BASE/VIAL
AP ELKINS-SINN/AHROBINS EQ 125MG BASE/VIAL
AP EQ 250MG BASE/VIAL
AP EQ 500MG BASE/VIAL
AP EQ 1GM BASE/VIAL
AP EQ 2GM BASE/VIAL
AP EQ 10GM BASE/VIAL
AP TOTACELLIN-H
AP BEECHAM LABS./BEECHAM EQ 10GM BASE/VIAL

INJECTABLE; INJECTION
AMPICILLIN SODIUM
AP ELI LILLY EQ 2GM BASE/VIAL
AP ELKINS-SINN/AHROBINS EQ 125MG BASE/VIAL
AP EQ 250MG BASE/VIAL
AP EQ 500MG BASE/VIAL
AP EQ 1GM BASE/VIAL
AP EQ 2GM BASE/VIAL
AP EQ 10GM BASE/VIAL
AP TOTACELLIN-H
AP BEECHAM LABS./BEECHAM EQ 10GM BASE/VIAL

INJECTABLE; INJECTION
BEROCCA PN
HOFFMANN-LA ROCHE 50MG/ML; 0.03MG/ML; 0.0025MG/ML;
7.5MG/ML; 100 IU/ML; 0.2MG/ML; 20MG/ML;
2MG/ML; 1.8MG/ML; 1.5MG/ML; 1,650 IU/ML;
5 IU/ML
N06071 003
OCT 10, 1985

ASPIRIN; BUTALBITAL; CAFFEINE (PAGE 3-19)

CAPSULE; ORAL
BUTALBITAL W/ ASPIRIN AND CAFFEINE
AB CHELSEA LABORATORIES 325MG; 50MG; 40MG
N86231 002
FEB 12, 1985

CAPSULE; ORAL
BUTALBITAL W/ ASPIRIN AND CAFFEINE
AB CHELSEA LABORATORIES 325MG; 50MG; 40MG
N17534 005
APR 16, 1986

CAPSULE; ORAL
BUTALBITAL W/ ASPIRIN AND CAFFEINE
AB CHELSEA LABORATORIES 325MG; 50MG; 40MG
N86231 002
FEB 12, 1985

CAPSULE; ORAL
BUTALBITAL W/ ASPIRIN AND CAFFEINE
AB CHELSEA LABORATORIES 325MG; 50MG; 40MG
N17534 005
APR 16, 1986

CAPSULE; ORAL
BUTALBITAL W/ ASPIRIN AND CAFFEINE
AB CHELSEA LABORATORIES 325MG; 50MG; 40MG
N86231 002
FEB 12, 1985

CAPSULE; ORAL
BUTALBITAL W/ ASPIRIN AND CAFFEINE
AB CHELSEA LABORATORIES 325MG; 50MG; 40MG
N17534 005
APR 16, 1986

CAPSULE; ORAL
BUTALBITAL W/ ASPIRIN AND CAFFEINE
AB CHELSEA LABORATORIES 325MG; 50MG; 40MG
N17534 005
APR 16, 1986

RX DRUG PRODUCT LIST / CUMULATIVE SUPPLEMENT NUMBER 11 / AUG '85 - JUL '86

5

ASPIRIN; BUTALBITAL; CAFFEINE (PAGE 3-19)

TABLET; ORAL <u>BUTALBITAL N/ ASPIRIN AND CAFFEINE</u>	<u>NEST-HARD</u>	<u>325MG; 50MG; 40MG</u>	NB6162 002 FEB 16, 1985	
<u>FLOORTHAL</u>	<u>SANDOZ PHARMS./SANDOZ</u>	<u>325MG; 50MG; 40MG</u>	N17534 003 APR 16, 1986	
<u>LANORTHAL</u>	<u>LANNETT</u>	<u>325MG; 50MG; 40MG</u>	N86986 002 OCT 18, 1985	

ASPIRIN; CARISOPRODOL (PAGE 3-20)

TABLET; ORAL <u>CARISOPRODOL COMPOUND</u>	<u>BOLAR PHARMACEUTICAL</u>	<u>325MG; 200MG</u>	N88809 001 OCT 03, 1985	
<u>SOMA COMPOUND</u>	<u>WALLACE PHARMS./C-W</u>	<u>325MG; 200MG</u>	N12365 005 JUL 11, 1983	

ASPIRIN; METHOCARBAMOL (PAGE 3-20)

TABLET; ORAL <u>METHOCARBAMOL AID ASPIRIN</u>	<u>MCNEIL CONSUMER PROD</u>	<u>325MG; 400MG</u>	N89193 901 FEB 12, 1985	
--	-----------------------------	---------------------	----------------------------	--

BACITRACIN ZINC; HYDROCORTISONE; NEOMYCIN SULFATE; POLYMYXIN B SULFATE (PAGE 3-23)

OINTMENT; TOPICAL <u>CORTISPORIN</u>	<u>BURROUGHS WELLCOME</u>	<u>400 UNITS/GM; 1%; EQ 3.5MG BASE/GM; 5,000 UNITS/GM</u>	N50168 002 MAY 04, 1985	
<u>HEOMYCIN & POLYMYXIN B SULFATES & BACITRACIN ZINC ®</u>				
<u>HYDROCORTISONE</u>	<u>PHARMAFAIR</u>	<u>400 UNITS/GM; 1%; EQ 3.5MG BASE/GM; 5,000 UNITS/GM</u>	N62381 001 SEP 06, 1985	

BENZYL PENICILLOYL-POLYLYSINE (PAGE 3-25)

INJECTABLE; INJECTION <u>KREMER'S®-SHRAN/</u>	<u>SCHWARZ PHARMS</u>	<u>/6.0MOL/L/</u> <u>60 UMOL/L</u>	/N50114 001/ N50114 001	
--	-----------------------	---------------------------------------	----------------------------	--

BETAMETHASONE BENZOATE (PAGE 3-25)

CREAM; TOPICAL <u>/BÉTAMÉTHASONE/</u>	<u>PARKER-DAVIS/N-L/</u>	<u>/6.25%/</u> <u>PARKER-DAVIS/N-L</u>	/N16998 001/ N16998 002
GEL; TOPICAL <u>/BÉTAMÉTHASONE/</u>	<u>UTICORT</u>	<u>0.25%</u>	
OINTMENT; TOPICAL <u>/BÉTAMÉTHASONE/</u>	<u>UTICORT</u>		
BETAMETHASONE DIPROPOXIMATE (PAGE 3-25)			
CREAM; TOPICAL <u>DIPROPOXIMATE</u>	<u>BX</u>	<u>EQ 0.05% BASEM</u>	N19408 001 JAN 31, 1986
LOTION; TOPICAL <u>ALPHATREX</u>	<u>AB</u>	<u>EQ 0.05% BASEM</u>	N70273 001 AUG 12, 1985
<u>BETAMETHASONE DIPROPIONATE</u>	<u>AB</u>	<u>EQ 0.05% BASEM</u>	N70275 001 AUG 12, 1985
OINTMENT; TOPICAL <u>BETA-VAL</u>	<u>AB</u>	<u>EQ 0.05% BASEM</u>	N70274 001 AUG 12, 1985
BETAMETHASONE VALERATE (PAGE 3-26)			
CREAM; TOPICAL <u>BETAMETHASONE VALERATE</u>	<u>CLAY-PARK LABS</u>	<u>EQ 0.1% BASEM</u>	N70053 001 JUN 10, 1986
OINTMENT; TOPICAL <u>BETA-VAL</u>	<u>LEMMON</u>	<u>EQ 0.1% BASEM</u>	N70069 001 DEC 19, 1985
BETAXOLOL HYDROCHLORIDE (PAGE 3-27)			
SOLUTION/DROPS; OPHTHALMIC <u>BETOPTIC</u>	<u>ALCON LABORATORIES</u>	<u>EQ 0.5% BASEM</u>	N19270 001 AUG 30, 1985

BETHANECHOL CHLORIDE (PAGE 3-27)

TABLET; ORAL
BETHANECHOL CHLORIDE 5MG
AA SIDMAK LABORATORIES 5MG
AA 50MG

N89095 001
DEC 19, 1985
N890% 001
DEC 19, 1985

BROMPHENIRAMINE MALEATE; PHENYLPROPANOLAMINE HYDROCHLORIDE (PAGE 3-29)

/TABLET; CONTROLLED RELEASE; ORAL
/DTM APP/
/AH ROBINS/ /12MG/75MG/
/AP. 62, 1984/

BRETYLIUM TOSYLATE (PAGE 3-28)

INJECTABLE; INJECTION <u>BRETYLIUM TOSYLATE</u>		50MG/ML	AP	N19033 001 APR 29, 1986 : APR 16, 1986	INJECTABLE; INJECTION <u>SENSORCATH</u>	0.25'M	AP	N70552 001 MAY 21, 1986
AP	ABBOTT LABORATORIES	50MG/ML	AP	N70545 001 MAY 14, 1986	ASTRA PHARM PRODS	0.52'M	AP	N70553 001 MAY 21, 1986
AP	ELKINS-SINN/AHROBINS	50MG/ML	AP	N70546 001 MAY 14, 1986		0.75'M	AP	N70554 001 MAY 21, 1986
AP		50MG/ML						
AP	INT'L MEDICATION SYS	50MG/ML	AP	N70119 001 APR 29, 1986 : MAR 06, 1986	<u>BUPIVACAINE HYDROCHLORIDE; DEXTROSE</u> (PAGE 3-29)			
AP	LYPHOMED	50MG/ML	AP	N70134 001 APR 29, 1986 : FEB 12, 1986	INJECTABLE; INJECTION MARCAINE SPINAL			
AP	ABBOTT LABORATORIES	50MG/ML	AP	N19030 001 APR 16, 1986	AP			
AP	<u>BRETYLOL</u> AM CRITICAL CARE/AHS	50MG/ML	AP	N17954 001 APR 29, 1986 : APR 16, 1986	<u>BUPROPION HYDROCHLORIDE</u> (PAGE 3-30)			
INJECTABLE; INJECTION <u>BRETYLIUM TOSYLATE</u>		50MG/ML	AP	N18644 001 APR 29, 1986	TABLET; ORAL WELLBUTRIN			
AP	ABBOTT LABORATORIES	200MG/100ML; 5GM/100ML	AP	N19005 002 APR 29, 1986 : APR 16, 1986	3 BURROUGHS WELLCOME	50MG	AP	N18644 002 DEC 30, 1985
AP		400MG/100ML; 5GM/100ML	AP	N19005 003 APR 29, 1986 : APR 16, 1986	3	75MG	AP	N18644 003 DEC 30, 1985
AP		800MG/100MG; 5GM/100ML	AP	N19005 001 APR 29, 1986 : APR 16, 1986	100MG		AP	N18644 003 DEC 30, 1985
AP	ABBOTT LABORATORIES	200MG/100ML; 5GM/100ML	AP	N19008 002 APR 29, 1986 : APR 16, 1986	<u>BUTOCONAZOLE NITRATE</u> (PAGE 3-31)			
AP		400MG/100ML; 5GM/100ML	AP	N19008 003 APR 29, 1986 : APR 16, 1986	CREAM; VAGINAL FEMSTAT			
AP		800MG/100MG; 5GM/100ML	AP	N19008 001 APR 29, 1986 : APR 16, 1986	SYNTEX LABS/SYNTEX	2'M	AP	N19215 001 NOV 25, 1985
AP	KENDALL MCGAN LABS	100MG/100ML; 5GM/100ML	AP	N19121 001 APR 29, 1986	SUPPOSITORY; VAGINAL FEMSTAT			
AP		200MG/100ML; 5GM/100ML	AP	N19121 002 APR 29, 1986	SYNTEX LABS/SYNTEX	100MG	AP	N19359 001 NOV 25, 1985
AP		400MG/100ML; 5GM/100ML	AP	N19121 003 APR 29, 1986				

/CALCIUM CHLORIDE; ANHYDROUS (PAGE 3-31)
CALCIUM CHLORIDE, ANHYDROUS (PAGE 3-31)

CALCITONIN, SALMON (PAGE 3-31)

INJECTABLE; INJECTION

MIACALCIN

SANDOZ PHARMS/SANDOZ 100 IU/ML

N17808 001
JUL 03, 1986

CALCIUM CHLORIDE; DEXTOSE; MAGNESIUM CHLORIDE; SODIUM ACETATE; SODIUM CHLORIDE (PAGE 3-32)

SOLUTION; INTRAPERITONEAL

DIALYTE W/ DEXTROSE 2.5% IN PLASTIC CONTAINER

KENDALL MCGRAW LABS

2.9MG/100ML; 2.5GM/100ML;

15MG/100ML; 610MG/100ML;

N18460 006
JAN 29, 1986

CALCIUM CHLORIDE; DEXTOSE; MAGNESIUM CHLORIDE; SODIUM CHLORIDE; SODIUM LACTATE (PAGE 3-32)

SOLUTION; INTRAPERITONEAL

DIALYTE W/ DEXTROSE 1.5% IN PLASTIC CONTAINER

KENDALL MCGRAW LABS

2.6MG/100ML; 1.5GM/100ML;

5MG/100ML; 530MG/100ML;

N18460 007
JAN 29, 1986

CALCIUM CHLORIDE; POTASSIUM CHLORIDE; SODIUM SODIUM CHLORIDE; SODIUM LACTATE (PAGE 3-34)

INJECTABLE; INJECTION

TPH ELECTROLYTES IN PLASTIC CONTAINER

AP Abbott Laboratories

16.5MG/ML; 25.4MG/ML; 74.6MG/ML;
121MG/ML; 16.1MG/ML

N19399 001
JUN 16, 1986

AP Abbott Laboratories

16.5MG/ML; 25.4MG/ML; 74.6MG/ML;
121MG/ML; 16.1MG/ML

N18895 001
JUL 20, 1984

CALCIUM CHLORIDE; POTASSIUM CHLORIDE; SODIUM SODIUM CHLORIDE; SODIUM LACTATE (PAGE 3-35)

INJECTABLE; INJECTION

LACTATED RINGER'S IN PLASTIC CONTAINER

AP Abbott Laboratories

20MG/100ML; 30MG/100ML; 600MG/100ML;
310MG/100ML

N19485 001
OCT 24, 1985

SOLUTION; IRRIGATION

LACTATED RINGER'S IN PLASTIC CONTAINER

AT Abbott Laboratories

20MG/100ML; 30MG/100ML; 600MG/100ML;
310MG/100ML

N19416 001
JAN 17, 1986

CARBACHOL (PAGE 3-36)

INJECTABLE; INJECTION

CARBACHOL

AP PHARMAFAIR

0.012%

MOSTAT ALCON LABORATORIES 0.01%

N16968 001

CARBAMAZEPINE (PAGE 3-36)

TABLET; ORAL
CARBAMAZEPINE
 AB COLMED LABORATORIES 200MG
TEGRETOL
 AB GEIGY/CIBA-GEIGY 200MG

CEFOTETAN DISODIUM (PAGE 3-36)

INJECTABLE; INJECTION
CEFOTAN
 STUART PHARMS./ICI EQ 1GM BASE/VIAL
 N70300 001 MAY 15, 1985
 N16608 001

CEFTAZIDIME (PAGE 3-36)

INJECTABLE; INJECTION
FORTAZ
 GLAXO 500MG/VIAL
VITACARN AP 1GM/VIAL
 KENDALL MCGRAW LABS APR 10, 1986
L-Carnitine/
CARNITOR AP 2GM/VIAL
SIGMA-TAU AP 6GM/VIAL
 N18948 001 DEC 27, 1985

CARNITINE, L- (PAGE 3-37)

SOLUTION; ORAL
VITACARN 1GM/10ML
 KENDALL MCGRAW LABS APR 10, 1986
L-Carnitine/
CARNITOR 330MG
SIGMA-TAU

CETAMOL NAFATE (PAGE 3-37)

INJECTABLE; INJECTION
MANDOL
 ELI LILLY EQ 1GM BASE/VIAL
 N62560 001 SEP 10, 1985
 EQ 2GM BASE/VIAL
 N62560 002 SEP 10, 1985
TAZZIDIME
 ELI LILLY AP 500MG/VIAL
 N62640 001 NOV 20, 1985
CEFAZOL
 ELI LILLY EQ 500MG BASE/VIAL
 N62557 001 SEP 10, 1985
 EQ 1GM BASE/VIAL
 N62557 002 SEP 10, 1985
TAZZIDIME IN PLASTIC CONTAINER
 ELI LILLY 1GM/VIAL
CEFOPERAZONE SODIUM (PAGE 3-38)

INJECTABLE; INJECTION
CEFOBID IN PLASTIC CONTAINER
 ROERIG/PFIZER EQ 40MG BASE/ML
 N50613 001 JUL 23, 1986
 >ADD>
 >ADD>
 >ADD>

CEFUROXIME SODIUM (PAGE 3-40)

INJECTABLE; INJECTION

KEFLUROX

ELI LILLY

EQ 750MG BASE/VIAL

N62591 001

JAN 10, 1986

CHLORPHENIRAMINE MALEATE; PHENYLPROPANOLOLAMINE HYDROCHLORIDE (PAGE 3-46)

AP	EQ 750MG BASE/VIAL	N62592 001	JAN 10, 1986	DRIVE	CAPSULE, CONTROLLED RELEASE ; ORAL
AP	EQ 1.5GM BASE/VIAL	N62591 002	JAN 10, 1986	BC	BF ASCHER
AP	EQ 1.5GM BASE/VIAL	N62592 002	JAN 10, 1986	ORNADE	12MG; 75MG
AP	EQ 1.5GM BASE/VIAL	N62590 001	JAN 10, 1986	BC	SK&F LABORATORIES
AP	EQ 1.5GM BASE/VIAL	N62590 002	JAN 10, 1986		12MG; 75MG

KEFUROX IN PLASTIC CONTAINER

ELI LILLY

EQ 750MG BASE/VIAL

N62590 001

JAN 10, 1986

CHLORPROPAMIDE (PAGE 3-48)

AP	EQ 1.5GM BASE/VIAL	N62590 002	JAN 10, 1986	TABLET; ORAL	CHLORPROPAMIDE
AP	EQ 1.5GM BASE/VIAL	N62590 002	JAN 10, 1986	AB	HALSEY DRUG
AP	EQ 1.5 GM BASE/VIAL	N50558 002	OCT 19, 1983		100MG
AP	EQ 1.5 GM BASE/VIAL	N50558 003	OCT 19, 1983		250MG

CHLORTHALIDONE (PAGE 3-49)

AP	EQ 750MG BASE/VIAL	N62590 001	JAN 10, 1986	TABLET; ORAL	CHLORTHALIDONE
AP	EQ 1.5GM BASE/VIAL	N62590 002	JAN 10, 1986	AB	MUTUAL PHARM
AP	EQ 1.5 GM BASE/VIAL	N50558 002	OCT 19, 1983		25MG
AP	EQ 1.5 GM BASE/VIAL	N50558 003	OCT 19, 1983		50MG

CEPHALOTHIN SODIUM (PAGE 3-40)

INJECTABLE; INJECTION

CEPHALOTHIN SODIUM

ABBOTT LABORATORIES

EQ 1GM BASE/VIAL

N62547 001

SEP 11, 1985

AP	EQ 1GM BASE/VIAL	N62548 001	SEP 11, 1985	> ADD >	AB
AP	EQ 2GM BASE/VIAL	N62547 002	SEP 11, 1985	> ADD >	AB
AP	EQ 2GM BASE/VIAL	N62548 002	SEP 11, 1985	> ADD >	AB

KEFLITH IN PLASTIC CONTAINER

ELI LILLY

EQ 1GM BASE/VIAL

N62549 001

SEP 10, 1985

CHLORAMPHENICOL (PAGE 3-42)

AP	EQ 2GM BASE/VIAL	N62549 002	SEP 10, 1985	INJECTABLE; INJECTION	CHROMIC CHLORIDE IN PLASTIC CONTAINER
AP	EQ 2GM BASE/VIAL	N62549 002	SEP 10, 1985	AB	ABBOTT LABORATORIES EQ 0.004MG CHROMIUM/MHL

AP	EQ 1GM BASE/VIAL	N62549 001	SEP 10, 1985	CHROMIC CHLORIDE (PAGE 3-50)	N18861 001
AP	EQ 2GM BASE/VIAL	N62549 002	SEP 10, 1985	INJECTABLE; INJECTION	JUN 26, 1986
AP	EQ 2GM BASE/VIAL	N62548 002	SEP 11, 1985	CHROMIC CHLORIDE IN PLASTIC CONTAINER	
AP	EQ 2GM BASE/VIAL	N62548 002	SEP 11, 1985	ABBOTT LABORATORIES EQ 0.004MG CHROMIUM/MHL	

CILASTATIN SODIUM; IMIPENEM (PAGE 3-50)

AT	SOLUTION/DROPS; OPHTHALMIC	N62628 001	INJECTABLE; INJECTION	CILASTATIN SODIUM; IMIPENEM (PAGE 3-50)
AT	CHLORAMPHENICOL	SEP 25, 1985	PRIMAXIN	
AT	CARTER-GLOGAU LABS	0.5/24	MS&D RES LABS/MERCK	

AT	AT	EQ 250MG BASE/VIAL;	EQ 250MG BASE/VIAL;	N50587 001
AT	AT	500MG/VIAL	500MG/VIAL	NOV 26, 1985
AT	AT	EQ 500MG BASE/VIAL;	EQ 500MG BASE/VIAL;	N50587 002
AT	AT	500MG/VIAL	500MG/VIAL	NOV 26, 1985

CODEINE PHOSPHATE; PSEUDOEPHEDRINE HYDROCHLORIDE; TRIPROLIDINE HYDROCHLORIDE (PAGE 3-53)

SYRUP; ORAL
HESITAFED®
LIFE LABORATORIES
1.0MG/5ML; 2.0MG/5ML;
1.25MG/5ML

**INTRAUTERINE DEVICE; INTRAUTERINE
CL-7** 89MG
3 SEARLE PHARMS
TATUM-T 120MG
3 SEARLE PHARMS

CROMOLYN SODIUM (PAGE 3-55)

CYCLOPHOSPHAMIDE (PAGE 3-57)

<u>LYOPHILIZED CITROXAN</u>	<u>100MG/VIAL</u>	N12142 006
BRISTOL LABS/B-H		DEC 05, 1985
<u>AP</u>	<u>200MG/VIAL</u>	N12142 007
		DEC 10, 1985
<u>AP</u>	<u>500MG/VIAL</u>	N12142 008
		JAN 04, 1984
<u>AP</u>	<u>1GM/VIAL</u>	N12142 010
		SEP 24, 1985
<u>AP</u>	<u>2GM/VIAL</u>	N12142 009

CYPROHEPTADINE HYDROCHLORIDE (PAGE 3-58)

<u>SYRUP; ORAL</u> <u>CYPROMEPTADINE HCL</u> <u>HALSEY DRUG</u>	<u>2Mg/5mL</u>	<u>N89199 001</u> <u>JUL 03, 1986</u>
<u>> ADD > AA</u> <u>> ADD ></u>	<u>TABLET; ORAL</u> <u>CYPROMEPTADINE HCL</u> <u>HALSEY DRUG</u>	<u>4Mg</u> <u>N89057 001</u> <u>JUL 03, 1986</u>

DEXAMETHASONE SODIUM PHOSPHATE (PAGE 3-62)

PAGE 3-63)

<u>AT</u>	<u>DEXAMETHASONE SODIUM PHOSPHATE; NEOMYCIN SULFATE</u>	<u>SOLUTION/DROPS; OPHTHALMIC</u>
	<u>NEOMYCIN SULFATE-Dexamethasone Sodium Phosphate</u>	<u>HEMISERINE SULFATE-Dexamethasone Sodium Phosphate</u>
	<u>CARTER-GLOGAU LABS</u>	<u>EQ 0.1% PHOSPHATE;</u>
		<u>EQ 3.5MG BASE /ML H</u>
		<u>N62714 0</u>
		<u>JUL 21, 19%</u>

<u>200MG/VIAL</u>	<u>1GM/VIAL</u>	<u>100MG/VIAL</u>	<u>200MG/VIAL</u>
1000 J NOV 1986	1000 J NOV 1986	1000 J NOV 1986	1000 J NOV 1986
JUL 03, 1986	N88374 001	N89194 001	N89195 001
<u>1GM/VIAL</u>	<u>SEP 24,</u>	<u>AUG 27,</u>	<u>AUG 27,</u>
1986 :	JUL 03, 1986	JUL 07, 1986	JUL 07, 1986
<u>CYCLOPHOSPHAMIDE</u>	<u>LYOPHIELIZED</u>	<u>POLARANTHE</u>	<u>SCHERING</u>
<u>LYPHOMED</u>		<u>AB</u>	<u>AB</u>
		<u>DECHLORPHENANTHE MALEATE</u>	<u>DECHLORPHENANTHE MALEATE</u>
		<u>SIDMAK LABORATORIES</u>	<u>SIDMAK LABORATORIES</u>
		<u>2MCG</u>	<u>2MCG</u>
		<u>TABLET; ORAL</u>	<u>TABLET; ORAL</u>

PAGE 3-633

N12142 005 JG 30, 1982	> ADD > AI > ADD > > ADD > > ADD >	CARTER-GLOGAU LABS EQ 0.1% PHOSPHATE; EQ 3.5MG BASE/MIL	N62714 0 JUL 21, 1986
---	---	--	--

DEXAMETHASONE SODIUM PHOSPHATE; NEOMYCIN SULFATE

卷之三

卷之三

DEXTOSE (PAGE 3-64)

INJECTABLE; INJECTION
DEXTOSE 5% IN PLASTIC CONTAINER

AP ABBOTT LABORATORIES 5GM/100ML
SEP 17, 1985 N19479 001
N16673 003
OCT 30, 1985

AP TRAVENOL LABS 50MG/ML
JUN 03, 1986

DEXTOSE 5% IN PLASTIC CONTAINER

AP ABBOTT LABORATORIES 500MG/ML
JUN 03, 1986

DEXTOSE; LIDOCAINE HYDROCHLORIDE (PAGE 3-66)

INJECTABLE; INJECTION
LIDOCAINE HCL 0.2% IN DEXTROSE 5% IN PLASTIC CONTAINER
ABBOTT LABORATORIES 5GM/100ML; 200MG/100ML N18954 001
JUL 09, 1985

DEXTOSE; MAGNESIUM CHLORIDE; POTASSIUM CHLORIDE; POTASSIUM PHOSPHATE, DIBASIC; SODIUM LACTATE; SODIUM PHOSPHATE, MONOBASIC (PAGE 3-66)

INJECTABLE; INJECTION
IONOSOL B AND DEXTROSE 5% IN PLASTIC CONTAINER
ABBOTT LABORATORIES 5GM/100ML; 53MG/100ML; 100MG/100ML;
100MG/100ML; 180MG/100ML;
280MG/100ML; 16MG/100ML N19515 001
MAY 08, 1986

DEXTOSE; MAGNESIUM CHLORIDE; POTASSIUM CHLORIDE; POTASSIUM PHOSPHATE, MONOBASIC; SODIUM LACTATE; SODIUM PHOSPHATE, MONOBASIC (PAGE 3-66)

INJECTABLE; INJECTION
IONOSOL MB AND DEXTROSE 5% IN PLASTIC CONTAINER
ABBOTT LABORATORIES 5GM/100ML; 30MG/100ML; 141MG/100ML;
15MG/100ML; 260MG/100ML;
25MG/100ML N19513 001
MAY 08, 1986

DEXTOSE; POTASSIUM CHLORIDE; POTASSIUM LACTATE; SODIUM CHLORIDE; SODIUM PHOSPHATE, MONOBASIC (PAGE 3-66)

INJECTABLE; INJECTION
IONOSOL T AND DEXTROSE 5% IN PLASTIC CONTAINER
ABBOTT LABORATORIES 5GM/100ML; 111MG/100ML; 256MG/100ML;
146MG/100ML; 207MG/100ML N19514 001
MAY 08, 1986

DEXTOSE; POTASSIUM CHLORIDE; SODIUM CHLORIDE (PAGE 3-68)

INJECTABLE; INJECTION
DEXTROSE 5% AND SODIUM CHLORIDE 0.3% W/ POTASSIUM CHLORIDE 0.075% IN PLASTIC CONTAINER
ABBOTT LABORATORIES 5GM/100ML; 74.5MG/100ML;
300MG/100ML N18876 001
JAN 17, 1986

DEXTROSE 5% AND SODIUM CHLORIDE 0.3% W/ POTASSIUM CHLORIDE 0.15% IN PLASTIC CONTAINER
ABBOTT LABORATORIES 5GM/100ML; 149MG/100ML;
300MG/100ML N18876 002
JAN 17, 1986

DEXTROSE 5% AND SODIUM CHLORIDE 0.3% W/ POTASSIUM CHLORIDE 0.224% IN PLASTIC CONTAINER
ABBOTT LABORATORIES 5GM/100ML; 224MG/100ML;
300MG/100ML N18876 003
JAN 17, 1986

DEXTROSE 5% AND SODIUM CHLORIDE 0.33% AND POTASSIUM CHLORIDE 0.075% IN PLASTIC CONTAINER
KENDALL MCGRAW LABS 5GM/100ML; 75MG/100ML;
330MG/100ML N18268 011
JAN 18, 1986

DEXTROSE 5% AND SODIUM CHLORIDE 0.33% AND POTASSIUM CHLORIDE 0.15% IN PLASTIC CONTAINER
KENDALL MCGRAW LABS 5GM/100ML; 150MG/100ML;
330MG/100ML N18268 012
JAN 18, 1986

DEXTROSE 5%, SODIUM CHLORIDE 0.33% AND POTASSIUM CHLORIDE 0.22% IN PLASTIC CONTAINER
KENDALL MCGRAW LABS 5GM/100ML; 220MG/100ML;
330MG/100ML N18268 013
JAN 18, 1986

DEXTROSE 5%, SODIUM CHLORIDE 0.33% AND POTASSIUM CHLORIDE 0.30% IN PLASTIC CONTAINER
KENDALL MCGRAW LABS 5GM/100ML; 300MG/100ML;
330MG/100ML N18268 014
JAN 18, 1986

DEXTROSE; SODIUM CHLORIDE (PAGE 3-70)

INJECTABLE; INJECTION
DEXTROSE 5% AND SODIUM CHLORIDE 0.3% IN PLASTIC CONTAINER
ABBOTT LABORATORIES 5GM/100ML; 225MG/100ML;
5GM/100ML; 225MG/100ML N17606 001
N19482 001
OCT 04, 1985

INJECTABLE; INJECTION
DEXTROSE 5% AND SODIUM CHLORIDE 0.45% IN PLASTIC CONTAINER
ABBOTT LABORATORIES 5GM/100ML; 450MG/100ML;
5GM/100ML; 450MG/100ML N19484 001
OCT 04, 1985

INJECTABLE; INJECTION
DEXTROSE 5% AND SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER
ABBOTT LABORATORIES 5GM/100ML; 900MG/100ML;
5GM/100ML; 900MG/100ML N19483 001
OCT 04, 1985

DEXTROSE, THEOPHYLLINE (PAGE 3-70)

INJECTABLE; INJECTION
THEOPHYLLINE AND DEXTROSE 5% IN PLASTIC CONTAINER
 TRAVENOL LABS 5GM/100ML; 320MG/100ML NOV 13, 1985

DIAZEPAM (PAGE 3-72)

INJECTABLE; INJECTION
DIAZEPAM
 CARTER-GLOGAU LABS 5MG/ML
 AP
 ELKINS-SINN/AHROBINS 5MG/ML
 AP
 5MG/ML
 AP
 5MG/ML
 AP
 LYPHOMED 5MG/ML
 AP
VALTUM
 HOFFMANN-LA ROCHE 5MG/ML

TABLET; ORAL
DIAZEPAM

BARR LABORATORIES

2MG
 AB
 5MG
 AB
 10MG
 AB
 CHELSEA LABORATORIES 2MG
 AB
 5MG
 AB
 10MG
 AB
 CORD LABORATORIES 2MG
 AB
 5MG
 AB
 10MG
 AB
 LEDERLE LABS/AM CYAN 2MG

DIAZEPAM (PAGE 3-72)

TABLET; ORAL
DIAZEPAM

MYLAN PHARMS 2MG
 AB
 5MG
 AB
 10MG
 AB
 PAR PHARMACEUTICAL 2MG
 AB
 5MG
 AB
 10MG
 AB
 PARKE-DAVIS/W-L 2MG
 AB
 5MG
 AB
 10MG
 AB
 PUREPAC/KALIPHARMA 2MG
 AB
 5MG
 AB
 10MG
 AB
 ROXANE LABORATORIES 2MG
 AB
 5MG
 AB
 10MG
 AB
 SUPERPHARM 2MG
 AB
 5MG
 AB
 10MG
 AB
 ZENITH LABORATORIES 2MG
 AB
 5MG
 AB
 10MG
 AB
Q-PAM QUANTUM PHARMICS 2MG
 AB
 5MG
 AB
 10MG

N70323 001 SEP 04, 1985
 N70324 001 SEP 04, 1985
 N70325 001 SEP 04, 1985
 N70462 001 FEB 25, 1986
 N70463 001 FEB 25, 1986
 N70464 001 FEB 25, 1986
 N70209 001 SEP 04, 1985
 N70210 001 SEP 04, 1985
 N70222 001 SEP 04, 1985
 N70781 001 MAR 19, 1986
 N70706 001 MAR 19, 1986
 N70707 001 MAR 19, 1986
 N70356 001 JUN 17, 1986
 N70357 001 JUN 17, 1986
 N70358 001 JUN 17, 1986
 N70642 001 DEC 11, 1985
 N70643 001 DEC 11, 1985
 N70360 001 SEP 04, 1985
 N70361 001 SEP 04, 1985
 N70362 001 SEP 04, 1985
 N70423 001 DEC 12, 1985
 N70424 001 DEC 12, 1985
 N70425 001 DEC 12, 1985

DIAZEPAM (PAGE 3-72)

TABLET; ORAL
VALIUM
 AB HOFFMANN-LA ROCHE
 AB AB AB

N13263 002
 N13263 004
 N13263 006

2MG
 5MG
 10MG

DIPHENHYDRAMINE HYDROCHLORIDE (PAGE 3-76)

CAPSULE; ORAL
DIPHENHYDRAMINE HCL
 AA PIONEER PHARMS
 AA

2.5MG
 5.0MG

DICYCLOMINE HYDROCHLORIDE (PAGE 3-73)

CAPSULE; ORAL
BENTYL
 AB MERRELL DOW/DOW CHEM 10MG

DICYCLOMINE HCL
 AB BOLAR PHARMACEUTICAL 10MG

CHELSEA LABORATORIES 10MG
 AB CORD LABORATORIES

 TABLET; ORAL
BENTYL
 AB MERRELL DOW/DOW CHEM 2.0MG

DICYCLOMINE HCL
 AB BARR LABORATORIES 20MG

 AB BOLAR PHARMACEUTICAL 20MG

N07409 001
 OCT 15, 1984

 N83179 001
 FEB 12, 1986
 N85082 001
 JUN 19, 1986

 N07409 001
 OCT 15, 1984

 N84600 001
 JUL 29, 1985
 N84361 001
 FEB 06, 1986

AUG 28, 1985
 N17741 001

 AP
 AP
 AP
 AP
 AP

DISOPYRAMIDE PHOSPHATE (PAGE 3-77)

CAPSULE; ORAL
DISOPYRAMIDE PHOSPHATE
 AB BARR LABORATORIES EQ 100MG BASEN

 AB BOLAR PHARMACEUTICAL EQ 150MG BASEN

 AB CORD LABORATORIES EQ 100MG BASEN

 AB ZENITH LABORATORIES EQ 150MG BASEN

 AB

N70351 001
 DEC 17, 1985
 N70352 001
 DEC 17, 1985
 N70240 001
 FEB 02, 1986
 N70241 001
 FEB 02, 1986
 N70470 001
 DEC 10, 1985
 N70471 001
 DEC 10, 1985
 N70186 001
 NOV 18, 1985
 N70187 001
 NOV 18, 1985

DIFLORASONE DIACETATE (PAGE 3-74)

CREAM; TOPICAL
DIFLORASONE DIACETATE 0.05%
 BX UP JOHN

FLORONE 0.05%
 BX UP JOHN

OINTMENT; TOPICAL
DIFLORASONE DIACETATE 0.05%
 BX UP JOHN

FLORONE 0.05%
 BX UP JOHN

N19259 001
 AUG 28, 1985

 N19260 001
 AUG 28, 1985

 N17994 001

 AP
 AP
 AP
 AP
 AP

DOPAMINE HYDROCHLORIDE (PAGE 3-78)

INJECTABLE; INJECTION
DOPAMINE HCL
 AP ASTRA PHARM PRODS 4.0MG/MLN

 AP 8.0MG/MLN

 AP 8.0MG/MLN

 AP 8.0MG/MLN

 AP 16.0MG/MLN

 AP 16.0MG/MLN

 AP 16.0MG/MLN

N70087 001
 OCT 23, 1985
 N70089 001
 OCT 23, 1985
 N70090 001
 OCT 23, 1985
 N70091 001
 OCT 23, 1985
 N70092 001
 OCT 23, 1985
 N70093 001
 OCT 23, 1985
 N70094 001
 OCT 23, 1985

DOPAMINE HYDROCHLORIDE (PAGE 3-78)INJECTABLE; INJECTIONDOPAMINE HCLLYPHODED160MG/ML

N70364 001

DEC 04, 1985

AB

DOXY FAULDING

EQ 100MG BASE

N50582 001

JUL 22, 1985

N62653 001

OCT 30, 1985

AP

SOLOPAK LABORATORIES

40MG/ML

N70011 001

AUG 29, 1985

AB

PARKE-DAVIS/W-L

EQ 100MG BASE

N70046 001

AUG 29, 1985

AP

40MG/ML

N70047 001

AUG 29, 1985

AP

80MG/ML

N70558 001

SEP 20, 1985

AP

80MG/ML

N70559 001

SEP 20, 1985

AP

DOXASTAT

PARKE-DAVIS/W-L

40MG/ML

N70047 001

AUG 29, 1985

AP

80MG/ML

N70047 001

AUG 29, 1985

AP

INTROZEN

AM CRITICAL CARE/AHS

160MG/ML

N17395 003

AP

DOXEPIPIN HYDROCHLORIDE (PAGE 3-78)

AB

CAPSULE; ORAL

DOXEPIPIN HCL

CHELSEA LABORATORIES

EQ 25MG BASE

AB

EQ 50MG BASE

AB

EQ 100MG BASE

AB

EQ 25MG BASE

AB

EQ 50MG BASE

AB

EQ 75MG BASE

AB

EQ 100MG BASE

AB

EQ 25MG BASE

AB

EQ 50MG BASE

AB

EQ 75MG BASE

AB

EQ 100MG BASE

AB

MYLAN PHARMS

AB

EQ 25MG BASE

AB

EQ 50MG BASE

AB

EQ 75MG BASE

AB

EQ 100MG BASE

AB

EQ 100MG BASE

N70790 001

MAY 13, 1986

AB

N70791 001

MAY 13, 1986

AB

N70792 001

MAY 13, 1986

AB

N70793 001

MAY 13, 1986

AB

N70794 001

MAY 13, 1986

AB

N70795 001

MAY 15, 1986

AB

N70827 001

MAY 15, 1986

AB

N70828 001

MAY 15, 1986

AB

N70825 001

MAY 15, 1986

AB

N70789 001

MAY 13, 1986

AB

N70790 001

MAY 13, 1986

AB

N70791 001

MAY 13, 1986

AB

N70792 001

MAY 13, 1986

AB

N70793 001

MAY 13, 1986

AB

N70794 001

MAY 13, 1986

AB

N70795 001

MAY 13, 1986

AB

N70796 001

MAY 13, 1986

AB

N70797 001

MAY 13, 1986

AB

N70798 001

MAY 13, 1986

AB

N70799 001

MAY 13, 1986

AB

N70800 001

MAY 13, 1986

AB

N70801 001

MAY 13, 1986

AB

N70802 001

MAY 13, 1986

AB

N70803 001

MAY 13, 1986

AB

N70804 001

MAY 13, 1986

AB

N70805 001

MAY 13, 1986

AB

N70806 001

MAY 13, 1986

AB

N70807 001

MAY 13, 1986

AB

N70808 001

MAY 13, 1986

AB

N70809 001

MAY 13, 1986

AB

N70810 001

MAY 13, 1986

AB

N70811 001

MAY 13, 1986

AB

N70812 001

MAY 13, 1986

AB

N70813 001

MAY 13, 1986

AB

N70814 001

MAY 13, 1986

AB

N70815 001

MAY 13, 1986

AB

N70816 001

MAY 13, 1986

AB

N70817 001

MAY 13, 1986

AB

N70818 001

MAY 13, 1986

AB

N70819 001

MAY 13, 1986

AB

N70820 001

MAY 13, 1986

AB

N70821 001

MAY 13, 1986

AB

N70822 001

MAY 13, 1986

AB

N70823 001

MAY 13, 1986

AB

N70824 001

MAY 13, 1986

AB

N70825 001

MAY 13, 1986

AB

N70826 001

MAY 13, 1986

AB

N70827 001

MAY 13, 1986

AB

N70828 001

MAY 13, 1986

AB

N70829 001

MAY 13, 1986

AB

N70830 001

MAY 13, 1986

AB

N70831 001

MAY 13, 1986

AB

N70832 001

MAY 13, 1986

AB

N70833 001

MAY 13, 1986

AB

N70834 001

MAY 13, 1986

AB

N70835 001

MAY 13, 1986

AB

N70836 001

MAY 13, 1986

AB

N70837 001

MAY 13, 1986

AB

N70838 001

MAY 13, 1986

AB

N70839 001

MAY 13, 1986

AB

N70840 001

MAY 13, 1986

AB

N70841 001

MAY 13, 1986

AB

N70842 001

MAY 13, 1986

AB

N70843 001

MAY 13, 1986

AB

N70844 001

MAY 13, 1986

AB

N70845 001

MAY 13, 1986

AB

N70846 001

MAY 13, 1986

AB

N70847 001

MAY 13, 1986

AB

N70848 001

MAY 13, 1986

AB

N70849 001

MAY 13, 1986

AB

N70850 001

MAY 13, 1986

AB

N70851 001

MAY 13, 1986

AB

N70852 001

MAY 13, 1986

AB

N70853 001

MAY 13, 1986

AB

>_ADD_>	ETHINYL ESTRADOL; NORETHINDRONE; FERROUS FUMARATE (PAGE 3-89)	FLUROMETHOLONE (PAGE 3-93)
>_ADD_>	TABLET; ORAL-28 NORQUEST FE SYNTEX (FP) 0.035MG; 1MG; 7.5MG	N18926 001 AB FLUOR-DOP SUSPENSION/DROPS; OPHTHALMIC FEB 27, 1986
>_ADD_>		JUL 18, 1986 COOPERVISION PHARMS 0.12M
>_ADD_>		AB FML ALLERGAN PHARMS 0.12M N16851 002 JUL 28, 1982
		FML FORTE ALLERGAN PHARMS 0.25M N19216 001 APR 23, 1986
		N16144 001
		FLUROMETHOLONE ACETATE (PAGE 3-93)
		SUSPENSION/DROPS; OPHTHALMIC
		OMNITROL ALCON LABORATORIES 0.12M
		N19079 001 FEB 11, 1986
		OCT 31, 1985 N18830 001
		N18830 002 200MG/ML
		OCT 31, 1985
		FLUOROURACIL (PAGE 3-93)
		INJECTABLE; INJECTION
		FLUOROURACIL AP INT'L PHARM PROD 50MG/ML
		AP LYPHOMED 50MG/ML
		N88929 001 MAR 04, 1986
		N89152 001 MAR 21, 1986
		/N16446.0001/ /AUG 17, 1984/
		FLUNISOLIDE (PAGE 3-92)
		AEROSOL; INHALATION
		/BIRONALIDE/ /SYNTEX/LABES/SYNTEX/ 6.025MG/144H/
		AEROBID KEY PHARMACEUTICALS 0.025MG/TINH
		N18340 001 AUG 17, 1984
		FLUPHENAZINE DECANATE (PAGE 3-94)
		INJECTABLE; INJECTION
		FLUPHENAZINE AQ QUAD PHARMS 25MG/ML
		N70762 001 FEB 20, 1986
		PROLODION DECANATE
		AI ER SQUIBB AND SONS 25MG/ML
		N16727 001 SEP 11, 1985
		FLUPHENAZINE HYDROCHLORIDE (PAGE 3-94)
		CONCENTRATE; ORAL
		PERMITTIL AA SCHERING 5MG/ML
		N16008 001 NOV 07, 1985
		FLUOROMETHOLONE (PAGE 3-93)
		OINTMENT; OPHTHALMIC
		FML ALLERGAN PHARMS 0.12M
		N17760 001 SEP 04, 1985

RX DRUG PRODUCT LIST / CUMULATIVE SUPPLEMENT NUMBER 11 / JUL '86 - AUG '85

ELIUIAZEPAM HYDROCHLORIDE (PAGE 3-95)

THE VENETIAN

LEADERLINE, MARCH, 1935

FOLVITE LEADERLE LABS./AM CYAN SMG/ML

<u>FULL ACCT</u>	BARR LABORATORIES	1MGR
AA	PIONEER PHARMS	1MGR

FUR USE/EHIE (PAGE 3-98)

<u>10MG/ML</u>	N70014 001 SEP 09, 1985
<u>10MG/ML</u>	N70095 001 SEP 09, 1985
<u>10MG/ML</u>	N70096 001 SEP 09, 1985
<u>10MG/ML</u>	N70023 001 FEB 05, 1986
<u>10MG/ML</u>	N70078 001 FEB 05, 1986

FUROSEMIDE (PAGE 3-96)

TABLET; ORAL <u>FUROSEPTIDE</u>		<u>20MG#</u>
		<u>BARR LABORATORIES</u>
		<u>DANBURY PHARMACAL</u>
N16721	001	<u>AB</u>
N16721	002	<u>AB</u>
		<u>N70344 001</u>
		<u>NOV 27, 1985</u>
		<u>N70345 001</u>
		<u>NOV 27, 1985</u>
		<u>N70444 001</u>
		<u>JAR 20, 1986</u>
		<u>N70445 001</u>
		<u>JAR 20, 1986</u>
		<u>AB</u>
		<u>AB</u>
		<u>AB</u>

/N65847.6661

N89202 001
FEB 18, 1986

N8894.9 001
SEP 13, 1985

SEP 09, 1985
N70095 001

N70023 001
FEB 05, 1986
N70078 001
FEB 05, 1986

TABLET; ORAL

<u>PROJECT NUMBER</u>	<u>TESTER</u>	<u>TEST</u>	<u>DATE</u>	<u>RESULT</u>
AB	BARR LABORATORIES	20MG <small>HR</small>	N70043 SEP 26, 1985	001
AB	DANBURY PHARMACAL	20MG <small>HR</small>	N700412 FEB 26, 1986	001
AB		40MG <small>HR</small>	N700413 FEB 26, 1986	001
AB	ROXANE LABORATORIES	50MG <small>HR</small>	N70086 JAN 24, 1986	001
AB	WATSON LABORATORIES	20MG <small>HR</small>	N70449 NOV 22, 1985	001
AB		40MG <small>HR</small>	N70450 NOV 22, 1985	001
AB		80MG <small>HR</small>	N70528 NOV 22, 1985	001

GENTAMICIN SULFATE (PAGE 3-97)

INJECTABLE, INJECTION CENTRAFATR

<u>GENTAMICIN SULFATE</u>	<u>ABOTT LABORATORIES</u>	<u>EQ 10MG BASE/ML</u>	N62612 004 FEB 20, 1986
SOLUTION/DROPS; OPHTHALMIC			
<u>GENTAMICIN SULFATE</u>	<u>CARTER-GLOGAU LABS</u>	<u>EQ 3MG BASE/ML</u>	N62523 001 NOV 25, 1985
<u>AP</u>			<u>AT</u>

THE INVESTIGATIVE INTERVIEW

N62588 006
JAN 06, 1986

N62588 007
JAN 06, 1986

N62588 008
JAN 06, 1986

N62588 009
JAN 06, 1986

N62588 010
JAN 06, 1986

GENTAMICIN SULFATE; SODIUM CHLORIDE (PAGE 3-98)

HAL OPERATOR (PAGE 3-102)

INJECTABLE; INJECTION

<u>GENTAMICIN SULFATE IN PLASTIC CONTAINER</u>			
AP	ABBOTT LABORATORIES	EQ 1.2MG BASE/ML; 9MG/ML	N62588 001 JAN 06, 1986
AP		EQ 1.4MG BASE/ML; 9MG/ML	N62588 002 JAN 06, 1986
AP		EQ 1.6MG BASE/ML; 9MG/ML	N62588 003 JAN 06, 1986
AP		EQ 1.8MG BASE/ML; 9MG/ML	N62588 004 JAN 06, 1986
AP		EQ 2MG BASE/ML; 9MG/ML	N62588 005 JAN 06, 1986

GLYCINE (PAGE 3-100)

**SOLUTION; IRRIGATION
AMINOCETIC ACID 1.5**

AT / TRAVENOL LABS / **AT** TRAVENOL LABS **AT** **GLYCENE 1.5% IN PLASTIC CONTAINER** / 1.5G/1.00ML

GLYCOPYRROLATE (PAGE 3-1)

ADD >

TABU ET AL.

TABLET; ORAL

HALOPERIDOL (PAGE 3-102)

TABLET; ORAL
HALDOL MCNEIL PHAR

1MG 2MG 5MG 10MG 20MG

HEPARIN SODIUM (PAGE 3-103)

INJECTABLE: INJECTION

卷之三

166 / ERIK SINSIN

卷之三

TABLE I: OBAL

HALOPERIDOL DECANOATE (PAGE 3-102)

**INJECTABLE; INJECTION
HALDOL DECANOATE
MCNEIL PHARM**

CONCENTRATE; ORAL			
<u>HALDOL</u>			
MCKEIL LABORATORIES	EQ 2MG BASE/ML		N15922 001
<u>HALOPERIDOL</u>			
BAY LABORATORIES	EQ 2MG BASE/ML	APR 15, 1986	N70710 001
NATL PHARM MFG/BARRE	EQ 2MG BASE/ML	APR 15, 1986	MAR 07, 1986
SEARLE PHARNS	EQ 2MG BASE/ML	APR 15, 1986	N70318 001
			APR 11, 1986
			N70726 001
			IN 10 1986

NOV 10, 1964

HYDROCHLORTIAZIDE; METHYLDOPA (PAGE 3-110)

TABLET; ORAL
METHYLDOPA AND HYDROCHLORTIAZIDE
BOLAR PHARMACEUTICAL 25MG; 250MG

AB N70365 001
MAR 19, 1986
N70366 001
APR 16, 1986
N70367 001
MAR 19, 1986
N70368 001
APR 16, 1986
N70182 001
JAN 15, 1986
N70183 001
JAN 15, 1986
N70543 001
JAN 15, 1986
N70544 001
JAN 15, 1986
N70264 001
JAN 23, 1986
N70265 001
JAN 23, 1986
N70688 001
APR 24, 1986
N70689 001
APR 24, 1986

AB CORD LABORATORIES
15MG; 250MG
AB CORD LABORATORIES
25MG; 250MG
AB CORD LABORATORIES
30MG; 500MG
AB CORD LABORATORIES
50MG; 500MG
AB CORD LABORATORIES
15MG; 250MG
AB CORD LABORATORIES
25MG; 250MG
AB CORD LABORATORIES
30MG; 500MG
AB CORD LABORATORIES
50MG; 500MG

AB MYLAN PHARMS
15MG; 250MG

AB PUREPAC/KALIPHARMA
25MG; 250MG

AB PUREPAC/KALIPHARMA
50MG; 500MG

AB N70368 001
APR 16, 1986
N70182 001
JAN 15, 1986
N70183 001
JAN 15, 1986
N70543 001
JAN 15, 1986
N70544 001
JAN 15, 1986
N70264 001
JAN 23, 1986
N70265 001
JAN 23, 1986
N70688 001
APR 24, 1986
N70689 001
APR 24, 1986

AB N70367 001
APR 16, 1986
N70182 001
JAN 15, 1986
N70183 001
JAN 15, 1986
N70543 001
JAN 15, 1986
N70544 001
JAN 15, 1986
N70264 001
JAN 23, 1986
N70265 001
JAN 23, 1986
N70688 001
APR 24, 1986
N70689 001
APR 24, 1986

AB N70368 001
APR 16, 1986
N70182 001
JAN 15, 1986
N70183 001
JAN 15, 1986
N70543 001
JAN 15, 1986
N70544 001
JAN 15, 1986
N70264 001
JAN 23, 1986
N70265 001
JAN 23, 1986
N70688 001
APR 24, 1986
N70689 001
APR 24, 1986

AB N70369 001
APR 16, 1986
N70182 001
JAN 15, 1986
N70183 001
JAN 15, 1986
N70543 001
JAN 15, 1986
N70544 001
JAN 15, 1986
N70264 001
JAN 23, 1986
N70265 001
JAN 23, 1986
N70688 001
APR 24, 1986
N70689 001
APR 24, 1986

AB N70370 001
APR 16, 1986
N70182 001
JAN 15, 1986
N70183 001
JAN 15, 1986
N70543 001
JAN 15, 1986
N70544 001
JAN 15, 1986
N70264 001
JAN 23, 1986
N70265 001
JAN 23, 1986
N70688 001
APR 24, 1986
N70689 001
APR 24, 1986

AB N70371 001
APR 16, 1986
N70182 001
JAN 15, 1986
N70183 001
JAN 15, 1986
N70543 001
JAN 15, 1986
N70544 001
JAN 15, 1986
N70264 001
JAN 23, 1986
N70265 001
JAN 23, 1986
N70688 001
APR 24, 1986
N70689 001
APR 24, 1986

AB N70372 001
APR 16, 1986
N70182 001
JAN 15, 1986
N70183 001
JAN 15, 1986
N70543 001
JAN 15, 1986
N70544 001
JAN 15, 1986
N70264 001
JAN 23, 1986
N70265 001
JAN 23, 1986
N70688 001
APR 24, 1986
N70689 001
APR 24, 1986

AB N70373 001
APR 16, 1986
N70182 001
JAN 15, 1986
N70183 001
JAN 15, 1986
N70543 001
JAN 15, 1986
N70544 001
JAN 15, 1986
N70264 001
JAN 23, 1986
N70265 001
JAN 23, 1986
N70688 001
APR 24, 1986
N70689 001
APR 24, 1986

AB N70374 001
APR 16, 1986
N70182 001
JAN 15, 1986
N70183 001
JAN 15, 1986
N70543 001
JAN 15, 1986
N70544 001
JAN 15, 1986
N70264 001
JAN 23, 1986
N70265 001
JAN 23, 1986
N70688 001
APR 24, 1986
N70689 001
APR 24, 1986

AB N70375 001
APR 16, 1986
N70182 001
JAN 15, 1986
N70183 001
JAN 15, 1986
N70543 001
JAN 15, 1986
N70544 001
JAN 15, 1986
N70264 001
JAN 23, 1986
N70265 001
JAN 23, 1986
N70688 001
APR 24, 1986
N70689 001
APR 24, 1986

AB N70376 001
APR 16, 1986
N70182 001
JAN 15, 1986
N70183 001
JAN 15, 1986
N70543 001
JAN 15, 1986
N70544 001
JAN 15, 1986
N70264 001
JAN 23, 1986
N70265 001
JAN 23, 1986
N70688 001
APR 24, 1986
N70689 001
APR 24, 1986

AB N70377 001
APR 16, 1986
N70182 001
JAN 15, 1986
N70183 001
JAN 15, 1986
N70543 001
JAN 15, 1986
N70544 001
JAN 15, 1986
N70264 001
JAN 23, 1986
N70265 001
JAN 23, 1986
N70688 001
APR 24, 1986
N70689 001
APR 24, 1986

AB N70378 001
APR 16, 1986
N70182 001
JAN 15, 1986
N70183 001
JAN 15, 1986
N70543 001
JAN 15, 1986
N70544 001
JAN 15, 1986
N70264 001
JAN 23, 1986
N70265 001
JAN 23, 1986
N70688 001
APR 24, 1986
N70689 001
APR 24, 1986

AB N70379 001
APR 16, 1986
N70182 001
JAN 15, 1986
N70183 001
JAN 15, 1986
N70543 001
JAN 15, 1986
N70544 001
JAN 15, 1986
N70264 001
JAN 23, 1986
N70265 001
JAN 23, 1986
N70688 001
APR 24, 1986
N70689 001
APR 24, 1986

AB N70380 001
APR 16, 1986
N70182 001
JAN 15, 1986
N70183 001
JAN 15, 1986
N70543 001
JAN 15, 1986
N70544 001
JAN 15, 1986
N70264 001
JAN 23, 1986
N70265 001
JAN 23, 1986
N70688 001
APR 24, 1986
N70689 001
APR 24, 1986

HYDROCHLORTIAZIDE; METYLDOPA (PAGE 3-110)

/Hydrochlorothiazide; Metyldopa (PAGE 3-110)
TABLET; ORAL
METHYLDOPA AND HYDROCHLORTIAZIDE
/SUSPENSION; ORAL/
/TUSSCONEX/
/PENWALT 'PHARM/'
/Eq. 5mg. BASE/5ml/

/NI 6768.666/

HYDROCORTISONE (PAGE 3-112)

CREAM; TOPICAL
ALA-CORT
DE-L-RAY LABORATORIES 1/2
HYDROCORTISONE
PHARMADERM/ALTANA 1/2
Lotion; TOPICAL
ALA-CORT
DEL-RAY LABORATORIES 1/2
ALA-SCALP
DEL-RAY LABORATORIES 2%
HYDROCORTISONE
THAMES PHARMACAL 1/2

OINTMENT; TOPICAL
HYDROCORTISONE IN ABSORBASE

AT CAROLINA MED PRODS 1/2
HYDROCORTISONE; NEOMYCIN SULFATE; POLYMYXIN B SULFATE
(PAGE 3-115)

SUSPENSION; OTIC
NEOTECN AND POLYMYXIN B SULFATES AND HYDROCORTISONE
CARTER-GLOGAU LABS 1/2; Eq. 3.5MG BASE/ML;
10,000 UNITS/ML
NOV 06, 1985
AT PHARMAFAIR 1/2; Eq. 3.5MG BASE/ML;
10,000 UNITS/ML
NOV 26, 1985

SUSPENSION/DROPS; OPHTHALMIC
CORTESPORIN
BURROUGHS WELLCOME 1/2; Eq. 3.5MG BASE/ML;
10,000 UNITS/ML
NOV 06, 1985
AT HECTOCHEM SULFATE-POLYMYXIN B SULFATE-HYDROCORTISONE
PHARMAFAIR 1/2; Eq. 3.5MG BASE/ML;
10,000 UNITS/ML
NOV 26, 1985

TABLET; ORAL
SPIRONOLACTONE
PUREPAC/KALIPHARMA 25MG; 250MG
SUPERPAC 25MG; 250MG
/Eq. 5mg. BASE/5ml;
10,000 UNITS/ML
NOV 06, 1985
AT PHARMAFAIR 1/2; Eq. 3.5MG BASE/ML;
10,000 UNITS/ML
NOV 24, 1985

HYDROCORTISONE ACETATE; NEOMYCIN SULFATE; POLYMYXIN B SULFATE
(PAGE 3-116)

CREAM; TOPICAL CORTISPORIN	BURROUGHS WELLCOME	0.5%;EQ 3.5MG BASE/GM; 10,000 UNITS/GM	N50218 001 AUG 09, 1985	> ADD > AB > ADD > AB > ADD > AB > ADD > AB > ADD > AB	TABLET; ORAL <u>HYDROXYZINE HCL</u> AMIDE PHARMACEUTICAL <u>10MG</u>	N89071 001 JUL 22, 1986
<u>HYDROCORTISONE BUTYRATE (PAGE 3-116)</u>						N89072 001 JUL 22, 1986
CREAM; TOPICAL HYDROCORTISONE BUTYRATE	BX 3 GIST-BROCADES	0.1%	N18514 001 MAY 31, 1982	AB	TABLET; ORAL <u>HYDROXYZINE HCL</u> AMIDE PHARMACEUTICAL <u>25MG</u>	N89073 001 JUL 22, 1986
LOCOID	OWEN LABS/DERM PRODS	0.1%	N18795 001 JAN 07, 1983	AB	COLMED LABORATORIES <u>HYDROXYZINE HCL</u> AMIDE PHARMACEUTICAL <u>50MG</u>	N89121 001 MAR 20, 1986
OINTMENT; TOPICAL HYDROCORTISONE BUTYRATE	BX 3 GIST-BROCADES	0.1%	N18652 001 OCT 29, 1982	AB	COLMED LABORATORIES <u>HYDROXYZINE HCL</u> AMIDE PHARMACEUTICAL <u>50MG</u>	N89122 001 MAR 20, 1986
LOCOID	OWEN LABS/DERM PRODS	0.1%	N19106 001 JUL 03, 1984	AB	COLMED LABORATORIES <u>HYDROXYZINE HCL</u> AMIDE PHARMACEUTICAL <u>50MG</u>	N89123 001 MAR 20, 1986
<u>HYDROFLUMETHIAZIDE; RESERPINE (PAGE 3-117)</u>						N89381 001 MAY 19, 1986
TABLET; ORAL HYDROFLUMETHIAZIDE AND RESERPINE PAR PHARMACEUTICAL 50MG;0.125MG						N89382 001 MAY 19, 1986
						N89383 001 MAY 19, 1986
						N88540 001 OCT 22, 1985
						N88551 001 OCT 22, 1985
						N88529 001 OCT 22, 1985
						N88617 001 JAN 10, 1986
						N88618 001 JAN 10, 1986
						N88619 001 JAN 10, 1986

HYDROXYZINE HYDROCHLORIDE (PAGE 3-118)

TABLET; ORAL <u>HYDROXYZINE HCL</u>	AB	TABLET; ORAL <u>HYDROXYZINE HCL</u> PAR PHARMACEUTICAL	AB	CAPSULE; ORAL <u>HYDROXYZINE PAMOTE</u> PAR PHARMACEUTICAL	AB	CAPSULE; ORAL <u>HYDROXYZINE PAMOTE</u> PAR PHARMACEUTICAL	AB
ELKINS-SINN/AHROBINS	N85551 002	/654441/662/ N88862 001	AB	EQ 25MG HCL	N89145 001	EQ 25MG HCL	N71264 001
HYDROXYZINE HCL /ELKINS-SINN/AHROBINS/50MG/ML	FEB 14, 1986	FEB 14, 1986	AB	EQ 50MG HCL	MAR 17, 1986	EQ 50MG HCL	JUL 25, 1986
PHARMAFAIR	N89106 001	N89106 001	AB	IBUPROFEN (PAGE 3-120)	N89146 001	IBUPROFEN (PAGE 3-120)	N70038 001
25MG/ML	FEB 14, 1986	FEB 14, 1986	AB	TABLET; ORAL <u>IBUPROFEN</u>	MAR 17, 1986	TABLET; ORAL <u>IBUPROFEN</u>	SEP 06, 1985
50MG/ML	N88881 001	N88881 001	AB	BOOTS PHARMACEUTICAL 800MG	N70041 001	BOOTS PHARMACEUTICAL 800MG	N70041 001
	FEB 14, 1986	FEB 14, 1986	> ADD > AB	CHELSEA LABORATORIES 400MG	SEP 06, 1985	CHELSEA LABORATORIES 400MG	SEP 06, 1985
	N89107 001	N89107 001	> ADD > AB	600MG		600MG	
	FEB 14, 1986	FEB 14, 1986	AB				

HYDROXYZINE PAMOTE (PAGE 3-120)

TABLET; ORAL <u>IBUPROFEN</u>	AB	TABLET; ORAL <u>IBUPROFEN</u>	AB
BOOTS PHARMACEUTICAL 800MG	AB	CHELSEA LABORATORIES 400MG	AB

TOPAMIDOL (PAGE 3-123)INJECTABLE; INJECTIONISOVUE-370
ER SQUIBB AND SONS
76/4N18735 003
DEC 31, 1985
ISOVUE-M 200
ER SQUIBB AND SONS
41/4N18735 001
DEC 31, 1985
ISOVUE-M 300
ER SQUIBB AND SONS
61/4N18735 004
DEC 31, 1985ISONIAZID (PAGE 3-125)SYRUP; ORAL
LANNAZIDAA
LANNETT
50MG/5MLNB9243 001
FEB 03, 1986KANAMYCIN SULFATE (PAGE 3-126)INJECTABLE; INJECTIONKANAMYCIN SULFATE
QUAD PHARMS
APEQ 75MG BASE/2ML
EQ 500MG BASE/2MLN62642 001
FEB 03, 1986
N62642 002EQ 1GM BASE/3ML
EQ 75MG BASE/2MLN62642 003
FEB 03, 1986
N62605 003EQ 500MG BASE/2ML
EQ 1GM BASE/3MLN62605 001
FEB 26, 1986
N62605 002EQ 300MG CARBONATE/5ML
FEB 26, 1986KETOCONAZOLE (PAGE 3-127)CREAM; TOPICAL
NIZORAL
JANSSEN PHARMA
2/4N19084 001
DEC 31, 1985KETOPROFEN (PAGE 3-127)
CAPSULE; ORAL
ORUDIS
WYETH : ABS./AHC
50MG
75MGLABETALOL HYDROCHLORIDE (PAGE 3-127)INJECTABLE; INJECTIONNORMODYNE
SCHERING
AP
TRANDATE
GLAXO
5MG/MLN18686 001
AUG 01, 1984
N19425 001
DEC 31, 1985N18735 004
DEC 31, 1985
LACTULOSE (PAGE 3-127)SYRUP; ORAL
LACTULOSE
AA
ROXANE LABORATORIES
10GM/15MLN17906 001
LEUCOVORIN CALCIUM (PAGE 3-127)TABLET; ORAL
LEUCOVORIN CALCIUM
BX
LEDERLE LABS/AM CYAN EQ 5MG BASE
N18459 001
JAN 30, 1986N18342 001
JUL 08, 1983
LEVOBUNOLOL HYDROCHLORIDE (PAGE 3-128)
BX
BURROUGHS WELLCOME
EQ 5MG BASE
SOLUTION/DROPS; OPHTHALMIC
BETAGAN
ALLERGAN PHARMS
0.5%
N19219 002
DEC 19, 1985

LITHIUM CITRATE (PAGE 3-132)

SYRUP; ORAL
LITHIUM CITRATE
AA
MY-K LABS
EQ 300MG CARBONATE/5MLN17755 001
MAY 21, 1986TABLET; ORAL
ATIVAN
AB
WYETH LABS/AMHO
AB
AB
N17794 001
N17794 002
N17794 003N18754 002
JAN 09, 1986
N18754 003
JAN 09, 1986

LORAZEPAM (PAGE 3-132)

TABLET; ORAL
LORAZEPAM
AB AM THERAPEUTIC 0.5MG
AB 1MG
AB 2MG
AB BARR LABORATORIES 0.5MG
AB 1MG
AB 2MG
AB DANBURY PHARMACAL 0.5MG
> ADD > AB QUANTUM PHARMICS 0.5MG
AB 1MG
AB 2MG
AB LOXAPINE SUCCINATE (PAGE 3-132)

TABLET; ORAL
LOXITANE
3 LEDERLE LABS/AM CYAN EQ 10MG BASE
3 EQ 25MG BASE
3 EQ 50MG BASE
MANGANESE CHLORIDE (PAGE 3-134)

INJECTABLE; INJECTION
MANGANESE CHLORIDE IN PLASTIC CONTAINER
ABBOTT LABORATORIES EQ 0.1MG MANGANESE/ML JUN 26, 1986

MANNITOL (PAGE 3-134)
SOLUTION; IRRIGATION
RESECTISOL /AM MCNA/ AM HOSP /AM MCNA/ AM HOSP RESECTISOL IN PLASTIC CONTAINER AM MCNA/AM HOSP 5GM/100ML

MECLIZINE HYDROCHLORIDE (PAGE 3-135)

TABLET; ORAL
MECLIZINE HCl
AA SIDMAK LABORATORIES 12.5MG
N70727 001 DEC 11, 1985
MAR 07, 1986 N88732 001
N70728 001 DEC 11, 1985
MAR 07, 1986 N88733 001
N70729 001 DEC 11, 1985
MAR 07, 1986 N88734 001
N70472 001 DEC 11, 1985
DEC 10, 1985 N89113 001
N70473 001 AUG 20, 1985
DEC 10, 1985 N89114 001
N70474 001 AUG 20, 1985
DEC 10, 1985
N71117 001
JUL 24, 1986 N11839 003
N71118 001
JUL 24, 1986
N71110 001
JUL 24, 1986
N70200 001
AUG 09, 1985
N70201 001
AUG 09, 1985
N70202 001
AUG 09, 1985
SERONO LABS
MEDROXYPROGESTERONE ACETATE (PAGE 3-136)

TABLET; ORAL
PROVERA
UPJOHN
5MG
NENOTROPINS (PAGE 1-137)
INJECTABLE; INJECTION
PERGONAL /SERONO LABS/
75 IU/AMP
150 IU/AMP
MAY 20, 1985
/N17646.661/
/N17646.662/
N17646 001
N17646 002
METHOCARBAMOL (PAGE 3-142)
TABLET; ORAL
METHOCARBAMOL
AA PIONEER PHARNS 500MG
AA 750MG
METHOTREXATE SODIUM (PAGE 3-143)
INJECTABLE; INJECTION
FOLEY
ADRIA LABS./ERBAMONT EQ 250MG BASE/VIAL
AP ADRIA LABS./ERBAMONT EQ 25MG BASE/ML
N88731 001 DEC 13, 1985
N89082 001 DEC 13, 1985
N88954 001 OCT 24, 1985
N89180 001 JAN 03, 1986
N89181 001 JAN 03, 1986
N89182 001 JAN 03, 1986
N89183 001 JAN 03, 1986

METHOTREXATE SODIUM (PAGE 3-143)INJECTABLE; INJECTIONMETHOTREXATE LPFLEDERLE LABS./AM CYAN EQ 25MG BASE/ML

AP	<u>METHOTREXATE SODIUM</u>	N11719 007	MAR 31, 1982	AB	TABLET; ORAL <u>METHYLDOPA</u>
AP	<u>INTL PHARM PRODS</u>	N88648 001	MAY 09, 1986	AB	<u>250MG</u>
AP	<u>LYPHOMED</u>	N89323 001	JUN 13, 1986	AB	<u>500MG</u>
AP	<u>EQ 2.5MG BASE/ML</u>	N88935 001	OCT 11, 1985	AB	<u>250MG</u>
AP	<u>EQ 20MG BASE/VIAL</u>	N89322 001	JUN 13, 1986	AB	<u>500MG</u>
AP	<u>EQ 25MG BASE/ML</u>	N89263 001	JUN 13, 1986	AB	<u>125MG</u>
AP	<u>EQ 50MG BASE/VIAL</u>	N88936 001	OCT 11, 1985	AB	<u>250MG</u>
AP	<u>EQ 100MG BASE/VIAL</u>	N89937 001	OCT 11, 1985	AB	<u>500MG</u>
AP	<u>QUAD PHARMS</u>	N89308 001	JUL 10, 1986	AB	<u>125MG</u>
AP	<u>EQ 25MG BASE/ML</u>	N89309 001	JUL 10, 1986	AB	<u>250MG</u>
AP	<u>EQ 20MG BASE/VIAL</u>	N89293 001	JUL 10, 1986	AB	<u>500MG</u>
AP	<u>EQ 50MG BASE/VIAL</u>	N89294 001	JUL 10, 1986	AB	<u>125MG</u>
AP	<u>EQ 100MG BASE/VIAL</u>	N89295 001	JUL 10, 1986	AB	<u>250MG</u>
AP	<u>EQ 250MG BASE/VIAL</u>	N89296 001	JUL 10, 1986	AB	<u>500MG</u>
AP	<u>METHOTREXATE</u>			AB	<u>ROXANE LABORATORIES</u>
AP	<u>LEDERLE LABS./AM CYAN EQ 2.5MG BASE/ML</u>	N11719 004	AB	<u>250MG</u>	
AP	<u>HEMAT</u>	N86358 004	AB	<u>500MG</u>	
AP	<u>BRISTOL LABS./B-M</u>		AB	<u>250MG</u>	

METHYLCLOTHIAZIDE (PAGE 3-143)

AB	<u>TABLET; ORAL METHYLCLOTHIAZIDE</u>	2.5MG	N89135 001	AP	<u>INJECTABLE; INJECTION ALDOMET</u>
AB			FEB 12, 1986	AP	<u>MS&D/MERCK METHYLDOPATE HCL</u>

METHYLDOPATE HYDROCHLORIDE (PAGE 3-144)

>ADD>	<u>AP</u>	<u>MS&D/MERCK METHYLDOPATE HCL</u>	<u>50MG/ML</u>
>ADD>	<u>AP</u>	<u>ELKINS-SINN/AHROBINS</u>	<u>50MG/ML</u>
>ADD>	<u>AP</u>	<u>LYPHOMED</u>	<u>50MG/ML</u>

METHYLDOPA (PAGE 3-144)

AP	<u>METHYLDOPA</u>	<u>BOLAR PHARMACEUTICAL</u>	<u>125MG</u>	N70245 001
AB				FEB 25, 1986
AB				N70246 001
AB				FEB 25, 1986
AB				N70247 001
AB				FEB 25, 1986
AB				N70703 001
AB				JUN 06, 1986
AB				N70625 001
AB				JUN 06, 1986
AB				N7070 003
AB				OCT 15, 1985
AB				N7084 001
AB				OCT 15, 1985
AB				N7085 001
AB				OCT 15, 1985
AB				N70331 001
AB				APR 15, 1986
AB				N70332 001
AB				APR 15, 1986
AB				N70333 001
AB				APR 15, 1986
AB				N70749 001
AB				FEB 07, 1986
AB				N70750 001
AB				FEB 07, 1986
AB				N70452 001
AB				FEB 07, 1986
AB				N70192 001
AB				APR 25, 1986
AB				N70193 001
AB				APR 25, 1986
AB				N70194 001
AB				APR 25, 1986
AB				N70098 001
AB				FEB 20, 1986
AB				N70343 001
AB				FEB 20, 1986

N13401 001
N70291 001
JUL 01, 1986
N70652 001
JUN 03, 1986

METHYLPREDNISOLONE SODIUM SUCCINATE (PAGE 3-145)INJECTABLE; INJECTION METRONIDAZOLE (PAGE 3-148)

<u>METHYLPREDNISOLONE SODIUM SUCCINATE</u>		<u>INJECTABLE; INJECTION</u>	<u>METRONIDAZOLE (PAGE 3-148)</u>
AP	<u>QUAD PHARMS</u>	<u>EQ 40MG BASE/VIAL</u>	N89264 001 JAN 22, 1986
AP		<u>EQ 125MG BASE/VIAL</u>	N89265 001 JAN 22, 1986
AP		<u>EQ 500MG BASE/VIAL</u>	N89266 001 JAN 22, 1986
AP		<u>EQ 1GM BASE/VIAL</u>	N89267 001 JAN 22, 1986
AP	<u>LYPHOMED</u>	<u>EQ 40MG BASE/VIAL</u>	N89143 001 MAR 28, 1986
AP		<u>EQ 125MG BASE/VIAL</u>	N89144 001 MAR 28, 1986
AP		<u>EQ 500MG BASE/VIAL</u>	N89186 001 MAR 28, 1986
AP		<u>EQ 1GM BASE/VIAL</u>	N89187 001 MAR 28, 1986
AP		<u>EQ 1GM BASE/VIAL</u>	N89188 001 MAR 28, 1986
AP		<u>EQ 1GM BASE/VIAL</u>	N89189 001 MAR 28, 1986
AP			

METOCLOPRAMIDE HYDROCHLORIDE (PAGE 3-147)INJECTABLE; INJECTION

<u>METOCLOPRAMIDE HCl</u>		<u>INJECTABLE; INJECTION</u>	<u>METRONIDAZOLE (PAGE 3-148)</u>
AP	<u>LYPHOMED</u>	<u>EQ 10MG BASE/2ML</u>	N70293 001 JAN 24, 1986
AP	<u>QUAD PHARMS</u>	<u>EQ 10MG BASE/2ML</u>	N70671 001 MAY 27, 1986
AP	<u>REGLAN</u>	<u>EQ 10MG BASE/2ML</u>	N17862 001 N17862 003
AP	<u>AH ROBINS</u>	<u>EQ 50MG BASE/10ML</u>	AUG 03, 1984
AP		<u>EQ 150MG BASE/30ML</u>	N17862 002 AUG 03, 1984
AB	<u>CHELSEA LABORATORIES</u>	<u>EQ 10MG BASE</u>	N70632 001 OCT 28, 1985
AB	<u>MAXOLON</u>		N70106 001 MAR 04, 1986
AB	<u>BEECHAM LABS/BEECHAM</u>	<u>EQ 10MG BASE</u>	
AB			

METOCLOPRAMIDE HYDROCHLORIDE (PAGE 3-149)

<u>METOCLOPRAMIDE HCl</u>		<u>INJECTABLE; INJECTION</u>	<u>METRONIDAZOLE (PAGE 3-148)</u>
AP	<u>LYPHOMED</u>	<u>FLAGYL I.V.</u>	
AP	<u>QUAD PHARMS</u>	<u>SEARLE PHARMS</u>	
AP	<u>REGLAN</u>	<u>METRONIDAZOLE HCl</u>	
AP	<u>AH ROBINS</u>	<u>LYPHOMED</u>	
AB	<u>CHELSEA LABORATORIES</u>	<u>EQ 10MG BASE</u>	
AB	<u>DANBURY PHARMACY</u>		
AB	<u>PAR PHARMACEUTICAL</u>		
AB	<u>PUREPAC/KALIPHARMA</u>		

<u>MIDAZOLAM HYDROCHLORIDE (PAGE 3-149)</u>		<u>INJECTABLE; INJECTION</u>	<u>METRONIDAZOLE (PAGE 3-148)</u>
AB	<u>CHESLEA LABORATORIES</u>	<u>EQ 10MG BASE</u>	N70453 001 JUN 06, 1985
AB	<u>DANBURY PHARMACY</u>	<u>EQ 10MG BASE</u>	N70511 001 JAN 22, 1986
AB	<u>PAR PHARMACEUTICAL</u>	<u>EQ 10MG BASE</u>	N70342 001 MAR 25, 1986
AB	<u>PUREPAC/KALIPHARMA</u>	<u>EQ 10MG BASE</u>	N70581 001 OCT 17, 1985
AB			

<u>N18873 002</u>	<u>DEC 30, 1985</u>
<u>N18873 003</u>	<u>DEC 30, 1985</u>
<u>N18873 004</u>	<u>DEC 30, 1985</u>

<u>N18654 001</u>	<u>DEC 20, 1985</u>

MONOOCTANOIN (PAGE 3-150)

LIQUID; PERFUSION, BILARY

MOCTANIN
/Ascor. Hops./Pharm/ / 100%/
ETHITEK PHARMS100%
IMNALOXONE HYDROCHLORIDE (PAGE 3-151)

INJECTABLE; INJECTION

HALOXONE
ELKINS-SINN/AHROBINS 0.4MG/ML
SEP 24, 1986 : OCT 22, 1985
AP
0.4MG/ML
SEP 24, 1986 : OCT 22, 1985
AP
0.4MG/ML
SEP 24, 1986 : OCT 22, 1985
AP
INTL MEDICATION SYS 0.4MG/ML
SEP 24, 1986 : NOV 06, 1985
AP
0.4MG/ML
SEP 24, 1986 : JAN 17, 1986
AP
MYETH LABS/AMHO 0.02MG/ML
SEP 24, 1986 : OCT 02, 1985
AP
0.02MG/ML
SEP 24, 1986 : OCT 02, 1985
AP
0.4MG/ML
SEP 24, 1986 : OCT 02, 1985
AP
0.4MG/ML
SEP 24, 1986 : OCT 02, 1985
AP
HALOXONE HCL
WINTHROP-BREON/STERL 0.02MG/ML
SEP 24, 1986 : APR 18, 1986
AP
0.4MG/ML
SEP 24, 1986 : APR 18, 1986
AP
MARGAN DUPONT PHARMS/DUPONT 0.02MG/ML
AP
/\$/
1MG/ML
JUN 14, 1982

NALBUPHINE HYDROCHLORIDE (PAGE 3-151)NALIDIXIC ACID (PAGE 3-151)

TABLET; ORAL
TALWIN NX
/WINTHROP-BREON/STERL/ 0.5MG BASE/
/DEC. 16., 1986/
WINTHROP-BREON/STERL EQ 0.5MG BASE;
EQ 50MG BASE
DEC 16, 1982

TABLET; ORAL
NANDROLONE DECANOATE (PAGE 3-151)
DECA-DURABOLIN
ORGANON/AZKONA 50MG/ML
N13132 001
JUN 12, 1986
N13132 002
JUN 12, 1986
N13132 003
JUN 12, 1986

TABLET; ORAL
N14214 002
N14214 004
N14214 005
> ADD > AO
/\$/
50MG/ML
100MG/ML
200MG/ML

TABLET; ORAL
N14214 002
N14214 004
N14214 005
> ADD > AO
> ADD > AO
> ADD > AO
> ADD > AO
/\$/
50MG/ML
100MG/ML
200MG/ML

TABLET; ORAL
N18733 001
DEC 16, 1982

TABLET; ORAL
N18024 001
N18024 001
MAY 27, 1982

TABLET; ORAL
N16636 002
N16636 001
N16636 003
JUN 14, 1982

TABLET; ORAL
N14214 002
N14214 004
N14214 005
> ADD > AO
> ADD > AO
> ADD > AO
> ADD > AO
/\$/
50MG/ML
100MG/ML
200MG/ML

RX DRUG PRODUCT LIST / CUMULATIVE SUPPLEMENT NUMBER 11 / AUG '85 - JUL '86

29

NANDROLONE DECANOATE (PAGE 3-151)

INJECTABLE; INJECTION

NANDROLONE DECANOATE

AO LEMMON 50MG/ML

AO 50MG/ML

AO QUAD PHARMS 50MG/ML

AO 100MG/ML

AO 200MG/ML

NEOMYCIN SULFATE; POLYMYXIN B SULFATE (PAGE 3-153)

SOLUTION; IRRIGATION

AT NEOSPORIN G.U. IRRIGANTBURROUGHS WELLCOME EQ 40MG BASE/ML;
200,000 UNITS/MLAT NEOMYCIN AND POLYMYXIN B SULFATESCARTER-GLOGAU LABS EQ 40MG BASE/ML;
200,000 UNITS/ML

NEOMYCIN SULFATE; TRIAMCINOLONE ACETONIDE (3-153)

CREAM; TOPICAL

> ADD > AT NEOMYCIN SULFATE-TRIAMCINOLONE ACETONIDE

> ADD > AT E FOUGERA/ALTANA EQ 3.5MG BASE/GM; 0.1/4 JUL 21, 1986 N62600 001

> ADD > AT NEOMYCIN SULFATE-TRIAMCINOLONE ACETONIDE

> ADD > AT PHARMADERM/ALTANA EQ 3.5MG BASE/GM; 0.1/4 JUL 21, 1986 N62595 001

> ADD > AT MYTREX A

> ADD > AT SAVAGE LABS/ALTANA EQ 3.5MG BASE/GM; 0.1/4 JUL 21, 1986 N62598 001

> ADD > AT

NIFEDIPINE (PAGE 3-154)

CAPSULE; ORAL

ADALAT

AO NB8554 001 FEB 10, 1986 AB MILES PHARM/VILES 1.0MG/ML

AO NB7598 001 OCT 06, 1983 AB PROCARDIA PFIZER LABS./PFIZER 1.0MG/ML

AO NB9248 001 JUN 25, 1986 > ADD > AB 2.0MG/ML

AO NB9249 001 JUN 25, 1986 > ADD > AB 2.0MG/ML

AO NB9250 001 JUN 25, 1986 NITROGLYCERIN (PAGE 3-154)

AEROSOL; ORAL
NITROLINGUAL
G POHL-BOSKAMP 0.4MG/SPRAY/N

NITROFENESTINE MALEATE (PAGE 3-155)

INJECTABLE; INJECTION

AT HETROGLYCERIN INTL MEDICATION SYS SMG/ML

AT LYPHOMED SMG/ML

AT SOLOPAK LABORATORIES SMG/ML

AT SMG/ML

NYSTATIN (PAGE 3-156)

POWDER; ORAL

NYELSTAT

AO AA LEDERLE LABS/AM CYAN 100%

AT NYSTATIN PADDACK LABORATORIES 100%

AO AA MAY 26, 1985

NYSTATIN (PAGE 3-156)

<u>SUSPENSION; ORAL</u>	<u>NYSTATIN</u>	<u>100,000 UNITS/ML</u>	N62571 001 OCT 29, 1985	<u>OXYPHENBUTAZONE (PAGE 3-159)</u>			
<u>AA</u>	<u>NASKA PHARMACAL</u>			<u>TABLET; ORAL</u>	<u>OXYPHENBUTAZONE</u>	<u>AB</u>	<u>3 BOLAR PHARMACEUTICAL 100MG</u>
							NR83399 001 SEP 17, 1984
<u>TABLET; ORAL</u>	<u>NYSTATIN</u>	<u>500,000 UNITS</u>	N62506 001 JAN 16, 1984	<u>PARGYLINE HYDROCHLORIDE (PAGE 3-160)</u>			
<u>AA</u>	<u>LEMMON</u>		N62524 001 NOV 26, 1985	<u>TABLET; ORAL</u>	<u>EUTONYL</u>	<u>AB</u>	<u>3 ABBOTT LABORATORIES 50MG</u>
							N13448 004
<u>TABLET; VAGINAL</u>	<u>NYSTATIN</u>	<u>100,000 UNITS</u>	N62615 001 OCT 17, 1985	<u>PENICILLIN G POTASSIUM (PAGE 3-161)</u>			
<u>AI</u>	<u>SIDMAX LABORATORIES</u>			<u>POWDER FOR RECONSTITUTION; ORAL</u>	<u>PENICILLIN G POTASSIUM</u>	<u>AA</u>	<u>200,000 UNITS/5ML</u>
						<u>AA</u>	<u>250,000 UNITS/5ML</u>
						<u>AA</u>	<u>400,000 UNITS/5ML</u>
<u>NYSTATIN; TRIAMCINOLONE ACETONIDE (PAGE 3-157)</u>							NR60752 003 NR60752 002 NR60752 001
<u>CREAM; TOPICAL</u>	<u>NYCO-TRIZASET III</u>	<u>100,000 UNITS/GM; 0.125</u>	N61954 002 SEP 20, 1985	<u>PERMETHRIN (PAGE 3-164)</u>			
<u>AI</u>	<u>LEMMON</u>			<u>LOTION; TOPICAL</u>	<u>NIX</u>	<u>AA</u>	<u>MAR 31, 1986</u>
							N13435 001
<u>NYTREX F</u>	<u>SAVAGE LABS/ALTANA</u>	<u>100,000 UNITS/GM; 0.125</u>	N62597 001 OCT 08, 1985		<u>BURROUGHS WELLCOME</u>	<u>AA</u>	
<u>AI</u>							
<u>NYSTATIN-TRIAMCINOLONE ACETONIDE</u>							
<u>AI</u>	<u>E FOUGERA/ALTANA</u>	<u>100,000 UNITS/GM; 0.125</u>	N62599 001 OCT 08, 1985	<u>PHENTERMINE HYDROCHLORIDE (PAGE 3-167)</u>			
<u>AI</u>	<u>PHARMADERM/ALTANA</u>	<u>100,000 UNITS/GM; 0.125</u>	N62596 001 OCT 08, 1985	<u>CAPSULE; ORAL</u>	<u>/ADT/</u>	<u>AA/</u>	
<u>NYSTATIN AND TRIAMCINOLONE ACETONIDE</u>					<u>/ADT/</u>	<u>/ADT/</u>	
<u>> ADD ></u>	<u>PHARMAFAIR</u>	<u>100,000 UNITS/GM; 0.125</u>	N62657 001 JUL 30, 1986				
<u>> ADD ></u>	<u>AI</u>						NR87126 001
<u>> ADD ></u>							
<u>OINTMENT; TOPICAL</u>	<u>NYCO-TRIZASET III</u>	<u>100,000 UNITS/GM; 0.125</u>	N62045 002 NOV 26, 1985	<u>PHENTERMINE HCL</u>	<u>30MG</u>	<u>AA</u>	<u>APR 25, 1986</u>
<u>AI</u>	<u>LEMMON</u>						NR87777 001
<u>NYCOLOG-III</u>							NOV 01, 1985
<u>AI</u>	<u>ER SQUIBB AND SONS</u>	<u>100,000 UNITS/GM; 0.125</u>	N60572 001 JUN 28, 1985		<u>30MG</u>	<u>AA</u>	
							NR87126 001
<u>NYTREX F</u>	<u>SAVAGE LABS/ALTANA</u>	<u>100,000 UNITS/GM; 0.125</u>	N62601 001 OCT 09, 1985	<u>PHENYL BUTAZONE (PAGE 3-168)</u>			
<u>AI</u>							
<u>NYSTATIN-TRIAMCINOLONE ACETONIDE</u>				<u>CAPSULE; ORAL</u>	<u>PHENYL BUTAZONE</u>	<u>AB</u>	<u>100MG</u>
<u>AI</u>	<u>E FOUGERA/ALTANA</u>	<u>100,000 UNITS/GM; 0.125</u>	N62602 001 OCT 09, 1985				
							NR88994 001
<u>AI</u>	<u>PHARMADERM/ALTANA</u>	<u>100,000 UNITS/GM; 0.125</u>	N62603 001 OCT 09, 1985	<u>TABLET; ORAL</u>	<u>PHENYL BUTAZONE</u>	<u>AB</u>	<u>100MG</u>
							DEC 04, 1985
<u>NYSTATIN AND TRIAMCINOLONE ACETONIDE</u>							
<u>AI</u>	<u>CLAY-PARK LABS</u>	<u>100,000 UNITS/GM; 0.125</u>	N62280 002 OCT 10, 1985				NR8863 001
<u>> ADD ></u>	<u>AI</u>	<u>PHARMAFAIR</u>	<u>100,000 UNITS/GM; 0.125</u>				DEC 04, 1985
<u>> ADD ></u>							

RX DRUG PRODUCT LIST / CUMULATIVE SUPPLEMENT NUMBER 11 / AUG '85 - JUL '86

31

PHENYLEPHRINE HYDROCHLORIDE; PROMETHAZINE HYDROCHLORIDE
(PAGE 3-168)POTASSIUM CHLORIDE (PAGE 3-171)SYRUP; ORAL
PROMETHAZINE VC PLATH

AA HR CENCI LABS 5MG/5ML; 6.25MG/5ML NOV 22, 1985 N88815 001

PHENYTOIN SODIUM, EXTENDED (PAGE 3-169)CAPSULE; ORAL
/EXTENDED PHENYTOIN SODIUM/
BOLAR PHARMACEUTICAL 100MGAB /SETRED/
PIENTEX BOLAR PHARMACEUTICAL 100MG DEC 21, 1984 N88711 001PHENYTOIN SODIUM, PROMPT (PAGE 3-169)CAPSULE; ORAL
/BX/
/BX/ DANBURY, ZENITH LABORATORIES /100MG/
100MG/ PROMPT PHENYTOIN SODIUM
BX DANBURY PHARMACAL 100MG
BX ZENITH LABORATORIES 100MGPIPERAZINE CITRATE (PAGE 3-170)TABLET; ORAL
ANTEPAR
3 BURROUGHS WELLCOME EQ 500MG BASE NO9102 003POLYETHYLENE GLYCOL 3350; POTASSIUM CHLORIDE; SODIUM BICARBONATE; SODIUM CHLORIDE; SODIUM SULFATE (PAGE 3-170)SOLUTION; ORAL
OCL
a ABBOTT LABORATORIES 6GM/100ML; 7.5MG/100ML; 16.9MG/100ML;
14.6MG/100ML;
1.29GM/100ML APR 30, 1986 N19284 001PRALIDOXIME CHLORIDE (PAGE 3-174)INJECTABLE; INJECTION
PRALIDOXIME CHLORIDE
AP SURVIVAL TECHNOLOGY 300MG/ML
/AP/ /PROTOPAM/
/SYRUP/ 'TECHNIPROS//300MG/ML/
/AP/ /300MG/ML/ /300MG/ML/PREDNISOLONE (PAGE 3-174)SYRUP; ORAL
PRELONE MURO PHARMACEUTICAL 15MG/5MLNB 90081 001
FEB 04, 1986POTASSIUM CHLORIDE (PAGE 3-171)INJECTABLE; INJECTION
POTASSIUM CHLORIDE
AP MAURY BIOLOGICAL 2MEQ/ML NO88286 001
SEP 05, 1985TABLET, CONTROLLED RELEASE; ORAL
K-DUR 10
BC KEY PHARMACEUTICALS 10MEQ
K-DUR 20
KEY PHARMACEUTICALS 20MEQ
/AP/ /A/S BENZON/
KALINORM /AP/ /AP/ /
/AP/ /AP/ /AP/ /
BC CIBA/CIBA-GEIGY 10MEQ
KLOR-CON BC UPSHER-SMITH LABS 8MEQ
SLOW-K BC CIBA-GEIGY 8MEQ
/TABLET; ORAL/
TABLET, CONTROLLED RELEASE; ORAL
POTASSIUM CITRATE (PAGE 3-173)
/POTASSIUM CITRATE/
UROCIT-K UNIV TX HLTH SCI CTR 5MEQ
N19071 001 AUG 30, 1985N19439 002
JUN 13, 1986N19439 001
JUN 13, 1986
/AP/ /AP/ /
N19381 001
APR 16, 1986N19439 001
JUN 13, 1986
/AP/ /AP/ /
N19381 001
APR 16, 1986

PREDNISONOLONE ACETATE; SULFACETAMIDE SODIUM (PAGE 3-175)

PREUDOMINE (PAGE 3-176)

N12813	002	> ADD >	<u>AB</u>	TONNE PAULSEN	<u>10MG</u>	NB9028	001
		> ADD >		/WEST-HARD/ <u>AB</u>	/WEST-HARD/ 50MG	JUL 24, 1986 NB8465	001
N88837	001					JUN 01, 1984 NB8832	001
		DEC 24, 1985		AB	WEST-HARD		

SOLUTION; ORAL PEDIAPRED FISONS		EQ 5MG BASE/5MLX	N19157 001 MAY 28, 1986	> ADD > AB > ADD > > ADD > > ADD > > ADD >	CAPSULE; ORAL <u>PROCAINAMIDE HCL</u> CORD LABORATORIES <u>250MG</u>	NB89219 001 JUL 01, 1986
PREDNISONE (PAGE 3-176)						
TABLET; ORAL DELTASONE UP-JOHN	AB AB AB	5MG 10MG 20MG	N09986 002 N09986 006 N09986 007	> ADD > AB > ADD > > ADD > > ADD >	INJECTABLE; INJECTION <u>PROCAINAMIDE HCL</u> ABBOTT LABORATORIES <u>100MG/MLX</u>	NB89069 001 FEB 12, 1986
ORASONE REID-RONELL LABS / REID-RONELL LABS	AB AB AB	50MG 50MG 50MG	/N85999 001 /N85999 001 /N85999 001	> ADD > AB > ADD > > ADD >	500MG/MLX	NB89070 001 FEB 12, 1986
PREDNISONE BARR LABORATORIES	AB AB AB	5MG 10MG 20MG	/N86959 001 /N86959 001 /N86959 001	> ADD > AB > ADD > > ADD >	100MG/MLX	NB89029 001 APR 17, 1986
PREDNISONE PHARMACAL / PHARMACAL /	AB AB AB	5MG 10MG 20MG	/N85162 001 /N85162 001 /N85162 001	> ADD > AB > ADD > > ADD >	500MG/MLX	NB89030 001 APR 17, 1986
DANBURY PHARMACAL	AB AB AB	5MG 10MG 20MG	/N85161 001 /N85161 001 /N85161 001	> ADD > AB > ADD > > ADD >	100MG/MLX	NB88824 001 NOV 19, 1985
DURAMED PHARMS	AB AB AB	5MG 10MG 20MG	/N88395 001 /N88395 001 /N88395 001	> ADD > AB > ADD > > ADD >	500MG/MLX	NB88830 001 NOV 20, 1985
	AB AB AB	10MG 20MG 5MG	N88395 001 N88396 001 N89245 001	OCT 04, 1983 OCT 04, 1983 DEC 04, 1985	250MG	NB89257 001 MAY 30, 1986
	AB AB AB	5MG 750MG 500MG	N88394 001 N88394 001 N89247 001	OCT 04, 1983 OCT 04, 1983 DEC 04, 1985	500MG	NB89026 001 OCT 22, 1985
	AB AB AB	10MG 20MG 20MG	N88396 001 N89246 001 N89247 001	OCT 04, 1983 DEC 04, 1985 DEC 04, 1985	750MG	NB89027 001 OCT 22, 1985
	AB AB	INVAMED			500MG	NB89284 001 JUN 23, 1986

PROPRANOLOL HYDROCHLORIDE (PAGE 3-183)

RANITIDINE HYDROCHLORIDE (PAGE 3-187)

PROTAMINE SULFATE (PAGE 3-184)

<u>INJECTABLE; INJECTION</u>	<u>SECRETIN-KABI</u>	<u>/15CU/VIAL/</u>	<u>N18290 001</u>
<u>PROTAMINE SULFATE</u>	<u>/KABIVITRUM/</u>	<u>75CU/VIAL</u>	
ELI LILLY	PHARMACIA/PHARMACIA		
QUAD PHARMS			
<u>50MG/VIAL</u>			
<u>UP JOHN</u>			
<u>50MG/VIAL</u>			
<u>QUAZEPAM (PAGE 3-186)</u>			
<u>TABLET; ORAL</u>			
DORMALIN			
SCHERING			
<u>15MG</u>			
<u>QUINIDINE GLUCONATE (PAGE 3-186)</u>			
<u>TABLET, CONTROLLED RELEASE; ORAL</u>			
QUINALAN			
LANNETT			
<u>324MGR</u>			
<u>SODIUM BICARBONATE (PAGE 3-191)</u>			
<u>INJECTABLE; INJECTION</u>			
SODIUM BICARBONATE IN PLASTIC CONTAINER			
ABBOTT LABORATORIES	0.9MEQ/ML		
<u>1MEQ/ML</u>			
<u>N18810 001</u>			
<u>DEC 23, 1985</u>			
<u>N18708 001</u>			
<u>DEC 27, 1985</u>			
<u>AB</u>	<u>TRAIVENOL LABS</u>	<u>12/</u>	
<u>ULTRA DERM</u>			
CHESEBROUGH-PONDS			
<u>12/</u>			
<u>N88081 001</u>			
<u>FEB 10, 1986</u>			
<u>N89164 001</u>			
<u>NOV 21, 1985</u>			
<u>BC</u>	<u>QUINIDINE GLUCONATE</u>	<u>324MGR</u>	
	SUPERPHARM		
<u>AB</u>	<u>QUINIDINE GLUCONATE</u>	<u>324MGR</u>	
	SUPERPHARM		
<u>N19443 001</u>			
<u>JUN 03, 1986</u>			
<u>N19443 002</u>			
<u>JUN 03, 1986</u>			

RX DRUG PRODUCT LIST / CUMULATIVE SUPPLEMENT NUMBER 11 / AUG'85 - JUL '86

35

SODIUM BICARBONATE; TARTARIC ACID (PAGE 3-191)

GRANULE, EFFERVESCENT; ORAL BAROS MALLINCKRODT 4.60MG/GM;420MG/GM
AP ABBOTT LABORATORIES 900MG/100ML

SODIUM CHLORIDE (PAGE 3-191)

INJECTABLE; INJECTION SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER
AP ABBOTT LABORATORIES 900MG/100ML
AP TRAVENOL LABS 2MG/ML

SODIUM IODIDE, I-123 (PAGE 3-193)

CAPSULE; ORAL SODIUM IODIDE I-123
③ BENEDICT NUCR PHARM 400 UCI

SODIUM NITROPRUSSIDE (PAGE 3-194)

INJECTABLE; INJECTION METROPRESS
AP ABBOTT LABORATORIES 50MG/VIAL

SOMATREM (PAGE 3-195)

INJECTABLE; INJECTION PROTROPIN GENENTECH

SOMATROPIN (PAGE 3-195)

INJECTABLE; INJECTION ASELLACRIN 10
③ SERONO LABS 10 IU/VIAL
ASELLACRIN 2
③ SERONO LABS 2 IU/VIAL
CRESCORMON
③ KABIVITRUM 4 IU/VIAL

SPIRONOLACTONE (PAGE 3-196)

TABLET; ORAL SPIRONOLACTONE
N18509 001 > ADD > AB
AUG 07, 1985 > ADD >
25MG

SODIUM CHLORIDE (PAGE 3-196)

> ADD > STANOZOLOL (PAGE 3-196)
> ADD >
> ADD >
N19480 001 > ADD >
SEP 17, 1985 WINSTROL
N16677 004 > ADD >
OCT 30, 1985 MINTHROP-BREON/STERL 2MG

SULCONAZOLE NITRATE (PAGE 3-197)

SOLUTION; TOPICAL SULCOSYN
SYNTEX LABS/SYNTEX 1/24
N18671 003 > ADD >
MAY 27, 1982 G AND H LABORATORIES 3.7%;2.86%;3.42%;0.64%
N70566 001 > ADD >
JUN 09, 1986 JUN 09, 1986 N88607 001
SULFABENZAMIDE; SULFACETAMIDE; SULFATHIAZOLE; UREA (PAGE 3-197)

SULFAMETHOXAZOLE; TRIMETHOPRIM (PAGE 3-198)

CREAM; VAGINAL GYNE-SULF
AI > ADD >
SUSPENSION; ORAL SEPTRA GRAPE
AB BURROUGHS WELLCOME 200MG/5ML;400MG/5ML
N19107 001 > ADD >
OCT 17, 1985 SULFAMETHOXAZOLE AND TRIMETHOPRIM
PLANTEK/IKAPHARM 200MG/5ML;400MG/5ML
N70028 001 JUN 02, 1987 : OCT 29, 1985 N17598 002
SULFAMETHOXAZOLE; TRIMETHOPRIM (PAGE 3-198)

SULFAMETHOXAZOLE AND TRIMETHOPRIM

N70203 001 > DLT > AB
400MG;800MG /JUN '82; '83; '84; '85; '86; '87; '88; '89; '90; '91; '92; '93; '94; '95
800MG;1600MG /JUN '82; '83; '84; '85; '86; '87; '88; '89; '90; '91; '92; '93; '94; '95
N70204 001 > DLT > AB
SIDMAK LABORATORIES 400MG;800MG /JUN '82; '83; '84; '85; '86; '87; '88; '89; '90; '91; '92; '93; '94; '95
800MG;1600MG /JUN '82; '83; '84; '85; '86; '87; '88; '89; '90; '91; '92; '93; '94; '95
N70215 001 > DLT > AB
SULFAMETHOXAZOLE AND TRIMETHOPRIM DOUBLE STRENGTH
PLANTEK/IKAPHARM 800MG;1600MG /JUN 02, 1987 : SEP 19, 1985 N70037 001
SULFAMETHOXAZOLE AND TRIMETHOPRIM SINGLE STRENGTH
PLANTEK/IKAPHARM 400MG;800MG /JUN 02, 1987 : SEP 19, 1985 N70030 001

SULFANILAMIDE (PAGE 3-199)

CREAM; VAGINAL
VAGITROL
LEMON
15GM

N88718 001
SEP 19, 1985
AP
TECHNECOLL
MALLINCKRODT
N/A
TESULOID
ER SQUIBB AND SONS
N/A
N17059 001
N16923 001

SULFINPYRAZONE (PAGE 3-200)

CAPSULE; ORAL
SULFINPYRAZONE

AB PAR PHARMACEUTICAL 200MG

N88934 001
SEP 06, 1985
AP
RESTOROL
SANDOZ PHARMS/SANDOZ
15MG
30MG
AB
/Sdhl4z/
TEMAZ
QUANTUM PHARMICS
15MG
30MG

SULFISOXAZOLE DIOLAMINE (PAGE 3-200)

OPHTHALMIC; SOLUTION

SULFISOXAZOLE DIOLAMINE
AT 2 BARNES-HIND PHARNS EQ 4% BASE

AT GANTREZIN EQ 4% BASE
HOFFMAN-LAROCHE

N84148 001
NO7757 002
AB
>ADD > AB
BARR LABORATORIES
15MG

AB
>ADD > AB
30MG
AB
>ADD > AB
COLMED LABORATORIES
15MG

AB
>ADD > AB
30MG
AB
>ADD > AB
MYLAN PHARMS
15MG

AB
>ADD > AB
30MG
AB
>ADD > AB
N70490 001
JUL 07, 1986
N70919 001
JUL 07, 1986
N70920 001
JUL 07, 1986

N18217 001
DEC 24, 1985
AB
>ADD > AB
30MG
AB
>ADD > AB
30MG

TECHNETIUM, TC-99M, SULFUR COLLOID (PAGE 3-203)

INJECTABLE; INJECTION
TECHNETIUM TC-99M SULFUR COLLOID
/GAMMA DIA-3 LEP/
SOLUTION; INJECTION, ORAL
TECHNETIUM TC-99M SULFUR COLLOID
GAMMA DIA-3 LEP,
/SULFUR COLLOID KIT/
/AN-SULFUR COLLOID /N/A/
AP CIS-US N/A
AP /TECHNECOLL
/MALLINCKRODT/ /N/A/
AP /TESULOID
/ER. SQUIBB. AND SONS/ /N/A/
N17724 001
/N/A/
N17858 001
N16923 001
THEOPHYLLINE (PAGE 3-206)
CAPSULE, CONTROLLED RELEASE; ORAL
THEO-DUR SPRINKLE
KEY PHARMACEUTICALS 50MG

N62486 001
JUL 24, 1986
N62486 002
JUL 24, 1986
N88022 001
SEP 10, 1985
N88016 001
SEP 10, 1985
N88015 001
SEP 10, 1985
N88015 001
SEP 10, 1985

TEMAZEPM (PAGE 3-203)

CAPSULE; ORAL
TEMAZEPM

N70564 001
OCT 15, 1985
N70547 001
OCT 15, 1985
AB
>ADD > AB
BARR LABORATORIES
15MG
AB
>ADD > AB
30MG
AB
>ADD > AB
COLMED LABORATORIES
15MG
AB
>ADD > AB
30MG
AB
>ADD > AB
MYLAN PHARMS
15MG
AB
>ADD > AB
N70490 001
JUL 07, 1986
N70919 001
JUL 07, 1986
N70920 001
JUL 07, 1986

TETRACYCLINE HYDROCHLORIDE (PAGE 3-205)

CAPSULE; ORAL
TETRACYCLINE HCl

N71174 001
JUL 10, 1986
N71175 001
JUL 10, 1986
N70489 001
JUL 07, 1986
N70490 001
JUL 07, 1986
N70919 001
JUL 07, 1986
N70920 001
JUL 07, 1986
AB
>ADD > AB
15MG
AB
>ADD > AB
30MG
AB
>ADD > AB
COLMED LABORATORIES
15MG
AB
>ADD > AB
30MG
AB
>ADD > AB
MYLAN PHARMS
15MG
AB
>ADD > AB
N70490 001
JUL 07, 1986
N70919 001
JUL 07, 1986
N70920 001
JUL 07, 1986

RX DRUG PRODUCT LIST / CUMULATIVE SUPPLEMENT NUMBER 11 / AUG '85 - JUL '86

THEOPHYLLINE (PAGE 3-206)

CAPSULE, CONTROLLED RELEASE; ORAL

THEOPHYLLINE-SR

BC RP SCHERER 300MG

Elixir; Oral
THEOPHYL 225
/KNOELL PHARMACEUTICAL/ 112.5MG/115ML/
MCNEIL PHARM 112.5MG/15ML

SYRUP; ORAL

ACURBROH
MERRELL DOW/DOW CHEM 150MG/15MLTABLET; ORAL
THEOPHYLLINE
NATL PHARM MFG/BARRE 150MG/15MLTABLET; ORAL
QUIBRON-T
MEAD JOHNSON/B-M 300MGSLO-PHYLLIN
/BD/
/BD/
WILLIAM H RORER/
AB AB
THEOPHYL-225
/KNOELL PHARMACEUTICAL/ 225MG/
MCNEIL PHARM 225MGTABLET, CHEWABLE; ORAL
THEOPHYL
MCNEIL PHARM 100MGTABLET, CONTROLLED RELEASE; ORAL
THEO-DUR
KEY PHARMACEUTICALS 450MGTABLET, CONTROLLED RELEASE; ORAL
THEO-DUR
ROXANE LABORATORIES 30MG/MLTHIORDAZINE HYDROCHLORIDE (PAGE 3-209)

CONCENTRATE; ORAL

THIORDAZINE HCL INTENSOL

AA ROXANE LABORATORIES 30MG/ML
AA 100MG/MLTHEOPHYLLINE (PAGE 3-206)TOLAZAMIDE (PAGE 3-212)

TABLET; ORAL

TOLAZAMIDE
BARR LABORATORIESN88255 001
JUN 12, 1986
AB
AB
AB/N86445 661/
N86485 001
AB
AB
AB

CHELSEA LABORATORIES 100MG

250MG
500MG
100MG
250MG
500MGN88746 001
NOV 22, 1985
AB
AB
AB

COLMED LABORATORIES

250MG
500MG
250MG
500MG
250MG
500MGN86545 001
AB
AB
AB
AB

CORD LABORATORIES

250MG
500MG
250MG
500MG
250MG
500MGN88656 001
AUG 22, 1985
AB
AB
AB
AB

DANBURY PHARMACAL

100MG
250MG
500MG
100MG
250MG
100MGN85202 001
N85204 001
N84726 001
N84726 001
AB
AB
AB

DURAMED PHARMS

100MG
250MG
500MG
100MG
100MG
250MG
500MG
100MGN86506 001
SEP 12, 1985
AB
AB
AB

MYLAN PHARMS

250MG
500MG
100MG
250MG
500MG
100MG
250MG
500MG
100MGN89131 001
JUN 25, 1986
AB
AB
AB

PAR PHARMACEUTICAL

JAN 02, 1986
JAN 06, 1986
JUN 16, 1986
JUN 16, 1986
DEC 16, 1985
N88942 001
DEC 16, 1985

TRIENTINE HYDROCHLORIDE (PAGE 3-216)

CAPSULE; ORAL
CUPRID MS&D RES LABS/MERCK 250M

N19194 001
NOV 08, 1985

YANCOMYCIN HYDROCHI OBIDE (BAGE 7-220)

CAPSULE; ORAL
VANCOCIN HCL

N50606 001
APR 15, 1986
N50606 002
APR 15, 1986

TRIMETHOBENZAMIDE HYDROCHLORIDE (PAGE 3-217)

**INJECTABLE; INJECTION
TRIMETHOBENZAMIDE HC
SOLOPAK LABORATORIES**

N88960 001
APR 04, 1986
N89043 001
APR 04, 1986
N89094 001
APR 04, 1986

INJECTABLES; INJECTION

> ADD >	<u>ELI LILLY</u>	<u>EQ 500MG BASE/VIAL</u>	<u>NE0180 001</u>
> ADD >	<u>ELI LILLY</u>	<u>EQ 500MG BASE/VIAL</u>	<u>NE2476 001</u>
> ADD >	<u>ELI LILLY</u>	<u>EQ 1GM BASE/VIAL</u>	<u>MAR 15, 1986</u>
> ADD >	<u>ELI LILLY</u>	<u>EQ 1GM BASE/VIAL</u>	<u>NE2476 002</u>
> ADD >	<u>ELI LILLY</u>	<u>EQ 1GM BASE/VIAL</u>	<u>MAR 21, 1986</u>
> ADD >	<u>VANCOLE</u> <u>LEDERLE</u>	<u>PARENTERALS</u>	<u>EQ 500MG BASE/VIAL</u>
> ADD >			<u>NE2682 001</u>
> ADD >			" " 22, 1986

卷之三

BARR LABORATORIES 100M⁶

100MEN N70494 001
JAN 22, 1986
200MEN N70495 001
SEP 26, 1986 : MAR 14, 1986

**SOLUTION/DROPS; OPHTHALMIC
TOPICAL**

VALPROIC ACID (PAGE 3-220)

CAPSULE; ORAL	
<u>DEPAKEINE</u>	
ABBOTT LABORATORIES	250MG
<u>VALPROIC ACID</u>	250MG
PAR PHARMACEUTICAL	

VALPROATE SODIUM (PAGE 3-220)

SYRUP; ORAL
DEPAKENE
ABOTT LABORATORIES
EQ 250MG BASE / 5ML
MYPROTC ACID
HY-K LABS
EQ 250MG BASE / 5ML

VANCOUVER HYDROCHI OBTDE (PAGE 3-220)

מִנְיָמִינָה וְעַלְיוֹן

EQ 125MG BASEM

N60180 001
N62476 001
MAR 15, 1986
N62476 002
MAR 21, 1986
N60180 002
MAR 21, 1986
N62682 001
MAR 22, 1986

卷之三

N70451 001
DEC 16, 1985
N70225 001
NOV 12, 1985
N70617 001
NOV 12, 1985
N70348 001
MAY 01, 1986
N70672 001
MAR 07, 1986

VINBLASTINE SULFATE (PAGE 3-221)

卷之三

VINCRISTINE SULFATE (PAGE 3-221)

INJECTABLE; INJECTION

ONCOGEN[®]
ELI LILLY

1MG/ML

N14103 003

MAR 07, 1984

AP VINCERISTINE SULFATE

1MG/ML

N70777 001

APR 29, 1986

AP QUAD PHARMS

1MG/ML

N70778 001

MAY 01, 1986

AP

MARFARIN SODIUM (PAGE 3-221)

TABLET; ORAL

COLMED[®]
/BX/ DUPONT[®] PHARMS/DUPONT 1.5MG/AB DUPONT[®] PHARMS/DUPONT 2.5MG

AB MARFARIN SODIUM

COLMED LABORATORIES 2.5MG

N88720 001

AUG 06, 1985

ZINC CHLORIDE (PAGE 3-223)

INJECTABLE; INJECTION

ZINC CHLORIDE IN PLASTIC CONTAINER

ABBOTT LABORATORIES EQ 1MG ZINC/ML

N19559 001

JUN 26, 1986

CHLORHEXIDINE GLUCONATE (PAGE 3-224)

SOLUTION; TOPICAL CIDA-STAT	HUNTINGTON LABS	2/24	N19258 001 JUL 22, 1986	SYRUP; ORAL BELDIN HALSEY DRUG	12.5MG/5MLX	N89179 001 JUN 05, 1986
CHG SCRUB	HUNTINGTON LABS	4/24	N19258 002 JUL 22, 1986	DIPHEN BAY LABORATORIES	12.5MG/5MLX	N70118 001 OCT 01, 1985
EXIDINE	XTTRIUM LABS	2/24	N19422 001 DEC 17, 1985	HYDRAMINE NATL PHARM MFG/BARRE	12.5MG/5MLX	N70205 001 JAN 28, 1986
		2.5/24	N19421 001 DEC 17, 1985			
>ADD>	STERI-STAT	N70104 001 JUL 24, 1986				
>ADD>	MEDICAL SYS RES	4/24				
>ADD>						

CHLORPHENIRAMINE MALEATE; PHENYLPROPANOLAMINE HYDROCHLORIDE
(DAGE 3-225.)

<u>TABLET, CONTROLLED RELEASE; ORAL PHENYLPROANALPINE HCL W/ CHLORPHENIRAMINE MALEATE</u>		<u>DORSEY LABS/SAICCOZ 12MG;75MG</u>	<u>N19613 0.1 JUN 16, 1986</u>	<u>TABLET; ORAL IBUPROFEN</u>	<u>BARR LABORATORIES</u>	<u>200MG#</u>	<u>SEP 24, 1986 : DEC 24, 1985</u>	<u>N70493 001</u>
<u>CHLORPHENIRAMINE MALEATE; PSEUDOEPHEDRINE HYDROCHLORIDE</u>				<u>CHELSEA LABORATORIES</u>	<u>200MG#</u>	<u>SEP 24, 1986 : MAY 07, 1986</u>	<u>N70605 001</u>	
<u>(PAGE 3-225)</u>				<u>DANBURY PHARMACAL</u>	<u>200MG#</u>	<u>SEP 24, 1986 : MAR 05, 1986</u>	<u>N70435 001</u>	
<u>CAPSULE, CONTROLLED RELEASE; ORAL ISOCLOL</u>		<u>AM CRITICAL CARE/AHS</u>	<u>8MG;120MG#</u>	<u>OHM LABORATORIES</u>	<u>200MG#</u>	<u>SEP 24, 1986 : JUL 15, 1986</u>	<u>N71163 001</u>	
<u>MAR 06, 1986</u>				<u>PAR PHARMACEUTICAL</u>	<u>200MG#</u>	<u>SEP 24, 1986 : OCT 18, 1985</u>	<u>N70481 001</u>	
<u>CHLORPHENIRAMINE POLISTIREX; CODEINE POLISTIREX</u>				<u>MEDIPREP MCNEIL CONSUMER PROD</u>	<u>200MG#</u>	<u>SEP 24, 1986 : NOV 06, 1986</u>	<u>N70475 001</u>	
<u>(PAGE 3-225)</u>					<u>200MG#</u>	<u>SEP 24, 1986 : JUN 26, 1986</u>	<u>N71215 001</u>	
<u>SYRUP; ORAL PENNUTSUSS</u>		<u>PENNWALT PHARM</u>	<u>EQ 4MG MALEATE/5ML; EQ 10MG BASE/5ML</u>	<u>N18928 001 AUG 14, 1985</u>	<u>>ADD > >ADD ></u>	<u>INSULIN, PURIFIED, HUMAN, SEMISYNTHETIC; INSULIN SUSPENSION, ISOPHANE, PURIFIED HUMAN (SEMISYNTHETIC)</u>	<u>(PAGE 3-226)</u>	
<u>(PAGE 3-225)</u>					<u>>ADD > >ADD > >ADD > >ADD ></u>	<u>INJECTABLE; INJECTION NOVOLIN 70/30 SQUIBB/NOVO</u>		
						<u>30 UNITS/ML;70 UNITS/ML N19441 001 JUL 11, 1986</u>		

INSULIN SUSPENSION, BIOSYNTHETIC HUMAN (PAGE 3-226)

INJECTABLE; INJECTION
HUMULIN BR
ELI LILLY

100 UNITS/ML
APR 28, 1986

INSULIN SUSPENSION, ISOPHANE, BIOSYNTHETIC HUMAN (PAGE 3-226)

INJECTABLE; INJECTION
HUMULIN N
ELI LILLY

100 UNITS/ML
OCT 28, 1985

INSULIN SUSPENSION, ISOPHANE, PURIFIED HUMAN (SEMSYNTETIC)
(PAGE 3-226)

INJECTABLE; INJECTION
INSULATARD NPH HUMAN
NORDISK USA

100 UNITS/ML
MAY 30, 1986

INSULIN ZINC SUSPENSION, BIOSYNTHETIC HUMAN (PAGE 3-226)

INJECTABLE; INJECTION
HUMULIN L
ELI LILLY

100 UNITS/ML
SEP 30, 1985

INSULIN, PURIFIED, HUMAN, SEMSYNTETIC (PAGE 3-227)

INJECTABLE; INJECTION
VELOSULIN HUMAN
NORDISK USA

100 UNITS/ML
MAY 30, 1986

OXYMETAZOLINE HYDROCHLORIDE (PAGE 3-228)

SOLUTION/DROPS; OPHTHALMIC
OCUCLEAR
SCHERING

0.025%
MAY 30, 1986

POVIDONE-IODINE (PAGE 3-228)

SPONGE; TOPICAL
POVIDONE-IODINE
PARKE-DAVIS/DESERET

20%
NOV 29, 1985

PSEUDOEPHEDRINE HYDROCHLORIDE (PAGE 3-228)

CAPSULE; CONTROLLED RELEASE; ORAL
/SUDAFED®/S.A./
SUDAFED 12 HOUR

PYRITHIONE ZINC (PAGE 3-228)

LOTION; TOPICAL HEAD & SHOULDERS CONDITIONER PROCTER AND GAMBLE	0.3% N19412 001 MAR 10, 1986
	0.3% N19412 002 MAR 10, 1986
	0.3% N19412 003 MAR 10, 1986
	0.3% N19412 004 MAR 10, 1986

DRUG PRODUCTS APPROVED UNDER SECTION 505 OF THE ACT / CUMULATIVE SUPPLEMENT NUMBER 11 / AUG '85 - JUL '86
BY THE DIVISION OF BLOOD AND BLOOD PRODUCTS LIST

NO SEPTEMBER - JULY APPROVALS

C. APPENDICES

1. Orphan Drug Products with Exclusive Approval
2. List of Drug Products Which Must Demonstrate in vivo Bioavailability Only if Product Fails to Achieve Adequate Dissolution
3. Biopharmaceutic Guidance Availability List
4. ANDA Suitability Petitions
5. Exclusivity Terms
6. Prescription and OTC Drug Product Patent and Exclusivity Data

APPENDIX 1**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.

APPENDIX 1

BIOLOGICAL PRODUCTS

<u>Active Ingred.(s)</u> <u>Strength</u>	<u>Trade Name</u> <u>Dosage Form; Route</u>	<u>Applicant</u>	<u>License Number</u> <u>Approval Date</u>	<u>Exclusivity</u> <u>Exp.Date</u>
Hemin 313mg/amp	Panhematin Injectable; Injection	Abbott Laboratories	43 Jul 20, 1983	ODE Jul 20, 1990
Digoxin Immune Fab (OVINE)	Digibind Injectable; Injection	Burroughs Wellcome	129 Apr 22, 1986	ODE Apr 22, 1993

APPENDIX 1

DRUG PRODUCTS

<u>Active Ingred.(s)</u> <u>Strength(s)</u>	<u>Trade Name</u> <u>Dosage Form; Route</u>	<u>Applicant</u>	<u>Appl. Prod.</u> <u>Approval Date</u>	<u>Exclusivity</u> <u>Exp. Date</u>
Chenodiol 250mg	Chenix Tablet; Oral	Rowell Laboratories	18513 002 Jul 28, 1983	ODE Jul 28, 1990
Cromolyn Sodium 4%	Opticrom Solution/Drops; Ophthalmic	Fisons	18155 001 Oct 3, 1984	ODE Oct 3, 1991
Carnitine, L- 330mg	L-Carnitine Tablet; Oral	Sigma-Tau	18948 001 Dec 27, 1985	ODE Dec 27, 1992
Carnitine, L- 1gm/10ml	Vitacarn Solution; Oral	Kendall McGaw Labs*	19257 001 Apr 10, 1986	
Naltrexone Hydrochloride 50mg	Trexan Tablet; Oral	Dupont Pharm's	18932 001 Nov 20, 1984	ODE Nov 20, 1991
Monooctanoic 100%	Moctanin Liquid; Perfusion Biliary	Ascot Hosp Pharms	19368 001 Oct 29, 1985	ODE Oct 29, 1992
Pentamidine Isethionate 300mg/ml	Pentam 300 Injectable; Injection	LyphoMed	19264 001 Oct 16, 1984	ODE Oct 16, 1991

(continued)

*Refer to Appendix I narrative

APPENDIX 1

DRUG PRODUCTS

(continued)

<u>Active Ingred. (s) Strength(s)</u>	<u>Trade Name Dosage Form; Route</u>	<u>Applicant</u>	<u>Appl. Prod. Approval Date</u>	<u>Exclusivity Exp. Date</u>
Potassium Citrate 5meq	Uroxit-K Tablet; Oral	Univ of Tx Hlth Sci Ctr	19071 001 Aug 30, 1985	ODE Aug 30, 1992
Somatrem 5mg/vial	Protropin Injectable; Injection	Genentech	19107 001 Oct 17, 1985	ODE Oct 17, 1992
Trientine Hydrochloride 250mg	Cupid Capsule; Oral	Merck Sharp and Do Home Res Labs	19194 001 Nov 8, 1985	ODE Nov 08, 1992

APPENDIX 2

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

Acetaminophen; Aspirin; Butalbital Capsule or Tablet; Oral 160-165mg; 160-165mg; 50mg	Aminophylline Tablet; Oral 100mg 200mg	Aspirin; Carisoprodol; Codeine Phosphate 325mg; 200mg; 10mg
Acetaminophen; Aspirin; Butalbital Capsule or Tablet; Oral 325mg; 325mg; 50mg	Aspirin; Butalbital; Capsule or Tablet; Oral 325mg; 50mg 650mg; 50mg	Aspirin; Meprobamate Tablet; Oral 325mg; 200mg
Acetaminophen; Aspirin; Butalbital; Caffeine Capsule or Tablet; Oral 160-165mg; 160-165mg; 50mg; 40mg	Aspirin; Butalbital; Caffeine Capsule or Tablet; Oral 325mg; 50mg; 40mg; 650mg; 50mg; 40mg;	Aspirin; Methocarbamol Tablet; Oral 325mg; 200mg
Acetaminophen; Aspirin; Butalbital; Caffeine Capsule or Tablet; Oral 325mg; 325mg; 50mg; 40mg	Aspirin; Caffeine; Carisoprodol Tablet; Oral 160mg; 32mg; 200mg	Chlorothiazide Tablet; Oral 250mg
Acetaminophen; Butalbital Capsule or Tablet; Oral 325mg; 50mg 650mg; 50mg	Aspirin; Caffeine; Carisoprodol; Codeine Phosphate Tablet; Oral 160mg; 32mg; 200mg; 16mg	Estrogens, Conjugated; Meprobamate Tablet; Oral 0.4mg; 200mg 0.4mg; 400mg
Acetaminophen; Butalbital; Caffeine Capsule or Tablet; Oral 325mg; 50mg; 40mg	Aspirin; Carisoprodol Tablet; Oral 325mg; 200mg	Hydroxyzine Hydrochloride Tablet; Oral 10mg 25mg 50mg. 100mg

APPENDIX 3

BIOPHARMACEUTIC GUIDANCE AVAILABILITY LIST

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 T7B-06, 5600 Fishers Lane, Rockville, MD 20857. Comments and suggestions concerning these guidances are encouraged and should be sent to the Division of Bioequivalence.

<u>Name of Drug</u>	<u>Date</u>	<u>Revised Date</u>
Acetohexamide	Nov 15, 1985	
Allopurinol	Jul 15, 1985	
Amiloride Hydrochloride	Mar 29, 1985	
Aminophylline Suppositories	Jul 05, 1983	
Amitriptyline Hydrochloride	Jul 05, 1983	
Anticholinergic Drugs (Controlled Release)	Nov 07, 1980	
Baclofen	May 05, 1986	
Carbamazepine	Dec 05, 1984	Aug 04, 1986
Chlordiazepoxide Hydrochloride	Jul 05, 1983	
Chlorpropamide	Jul 05, 1983	
Chlorthalidone	Jul 05, 1983	
Clofibrate	Apr 07, 1986	
Clonidine Hydrochloride	Dec 05, 1984	
Clorazepate Dipotassium	Mar 10, 1986	
Diazepam (revised)	Jul 08, 1985	
Dicyclomine Hydrochloride	Aug 10, 1984	
Dipyridamole	Jul 05, 1983	
Disopyramide Phosphate	Jul 09, 1985	
Dissolution Testing (General)	Apr 19, 1985	
Doxepin Hydrochloride	Apr 02, 1985	
Erythromycin	Apr 05, 1977	
Flurazepam	Oct 15, 1985	

(continued)

APPENDIX 3

(continued)

Name of Drug	Date	Revised Date
Hydrochlorothiazide	Jul 25, 1983	
Hydroxyzine Hydrochloride (Dissolution Only)	Jan 27, 1981	
Hydroxyzine Pamoate	Jul 26, 1983	
Indometacin	Apr 06, 1985	
Isosorbide Dinitrate	Jun 04, 1985	
Isosorbide Dinitrate (Controlled Release Products)	Sep 19, 1985	
Lorazepam	Dec 03, 1984	
Methylprednisolone	Jun 12, 1986	
Methyltestosterone	Nov 16, 1979	
Metoclopramide	Dec 27, 1984	
Minoxidil	Apr 02, 1986	
Nitrofurantoin (Macrocrystalline)	Oct 29, 1985	
Phentermine Hydrochloride (Dissolution)	Nov 21, 1980	
Phentermine Hydrochloride (Slow Dissolving; Dissolution)	Nov 21, 1980	
Phenylbutazone & Oxyphenbutazone	Jul 26, 1983	
Prednisone (Dissolution Only)	Jul 10, 1985	
Probenecid	Jul 26, 1983	
Procainamide	Jul 25, 1983	
Propranolol	May 19, 1984	
Quinidine Gluconate (Controlled Release)	Jun 15, 1981	
Spironolactone	Jul 25, 1983	
Sulfinpyrazone	Jul 15, 1983	
Temazepam	Aug 1985	
Theophylline (Controlled Release)	Apr 1984	
Theophylline (Immediate Release)	Nov 02, 1983	
Tolazamide	Aug 22, 1984	
Tolbutamide	Jan 1982	
Trazodone	Nov 15, 1985	
Verapamil	Jul 1985	
		Apr 30, 1986

APPENDIX 4

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 (List I., Petitions Approved) and (2) is not suitable for submission as an ANDA (List II., 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.

<u>I. Petitions Approved</u>	<u>Drug Name Dosage Form; Route</u>	<u>Strength (Container Size)</u>	<u>Docket Number</u>	<u>Petitioner</u>	<u>Reason for Petition</u>	<u>Status</u>
	Acetaminophen; Codeine Phosphate Capsule; Oral	500mg 30mg	84 P-0228/CP	McNeil Pharm	New Dosage Form New Strength	Approved Jun 02, 1986
	Acetaminophen; Codeine Phosphate Capsule; Oral	500mg 60mg	84 P-0228/CP	McNeil Pharm	New Dosage Form New Strength	Approved Jun 02, 1986
	Acetaminophen; Codeine Phosphate Soft Gelatin Capsule; Oral	300mg 30mg	85 P-0543/CP	Softan	New Dosage Form	Approved Mar 18, 1986
	Acetaminophen; Codeine Phosphate Soft Gelatin Capsule; Oral	500mg 7.5mg	85 P-0543/ CP0002	Softan	New Dosage Form New Strength	Approved Mar 19, 1986

(continued)

APPENDIX 4

I. Petitions Approved

(continued)

<u>Drug Name Dosage Form; Route</u>	<u>Strength (Container Size)</u>	<u>Docket Number</u>	<u>Petitioner</u>	<u>Reason for Petition</u>	<u>Status</u>
Acetaminophen; Codeine Phosphate Soft Gelatin Capsule; Oral	500mg 15mg	85 P-0543/ CP0002	Softan	New Dosage Form New Strength	Approved Mar 19, 1986
Acetaminophen; Codeine Phosphate Elixir; Oral	160mg/5ml 6mg/5ml	86 P-0133/CP	Kleinfield, Kaplan and Becker	New Strength	Approved May 21, 1986
Acetaminophen; Codeine Phosphate Tablet; Oral	500mg 15mg	86 P-0161/CP	Roxane Laboratories	New Strength	Approved May 8, 1986
Acetaminophen; Codeine Phosphate Tablet; Oral	500mg 30mg	86 P-0161/CP	Roxane Laboratories	New Strength	Approved May 8, 1986
Acetaminophen; Codeine Phosphate Tablet; Oral	500mg 60mg	86 P-0161/CP	Roxane Laboratories	New Strength	Approved May 8, 1986
Acetaminophen; Hydrocodone Bitartrate Solution; Oral	500mg/15ml 5mg/15ml	84 P-0391/CP	UAD Laboratories	New Dosage Form	Approved Jul 2, 1985

(continued)

APPENDIX 4

I. Petitions Approved

(continued)

<u>Drug Name Dosage Form; Route</u>	<u>Strength (Container Size)</u>	<u>Docket Number</u>	<u>Petitioner</u>	<u>Reason for Petition</u>	<u>Status</u>
Acetaminophen; Oxycodone Hydrochloride Solution; Oral	325mg/5ml 5mg/5ml	85 P-0085/CP	Roxane Laboratories	New Dosage Form	Approved Aug 23, 1985
Acetaminophen; Oxycodone Hydrochloride Soft Gelatin capsule; Oral	500mg 5mg	85 P-0543/ CP0003	Softan	New Dosage Form	Approved Mar 18, 1986
Acetaminophen; Propoxyphene Hydrochloride Soft Gelatin capsule; Oral	500mg 32mg	85 P-0581/CP	Softan	New Dosage Form New Strength	Approved Mar 18, 1986
Acetaminophen Suppository; Rectal	80mg	85 P-0403/CP	Upsher-Smith Labs	New Dosage Form (Pediatric)	Approved Oct 16, 1985
Aminocaproic Acid Injectable; Injection	500mg/ml 10ml/vial	85 P-0308/CP	Abbott Laboratories	New Strength	Approved Feb 12, 1986
Aminophylline Injectable; Injection	10mg/ml 10ml/vial	85 P-0459/CP	Abbott Laboratories	New Strength	Approved Feb 12, 1986

(continued)

APPENDIX 4

I. Petitions Approved

(continued)

<u>Drug Name Dosage Form; Route</u>	<u>Strength (Container Size)</u>	<u>Docket Number</u>	<u>Petitioner</u>	<u>Reason for Petition</u>	<u>Status</u>
Aminophylline Injectable; Injection	50mg/ml 20ml/vial	85 P-0459/CP	Abbott Laboratories	New Strength	Approved Feb 12, 1986
Azatadine Maleate; Phenylpropanolamine Hydrochloride Sustained Release Capsule; Oral	1mg 75mg	85 P-0492/CP	SK&F Laboratories	New Combination New Dosage Form	Approved Jan 28, 1986
Benztropine Mesylate Syrup; Oral	0.5mg/5ml	85 P-0423/CP	RIM Consulting	New Dosage Form	Approved Oct 16, 1985
Bretlyium Tosylate Injectable; Injection	100mg/ml	86 P-0157/CP	Lyphomed	New Strength	Approved May 8, 1986
Brompheniramine Maleate; Pseudoephedrine Hydrochloride Sustained Release Capsule; Oral	12mg 120mg	85 P-0095/CP	UAD Laboratories	New Combination New Dosage Form	Approved Dec 13, 1985
Cholestyramine Tablet, Chewable; Oral	Eq 4gm Resin	86 P-0123/CP	Parke-Davis Labs/W-L	New Dosage Form	Approved Jun 20, 1986

(continued)

APPENDIX 4

I. Petitions Approved
 (continued)

<u>Drug Name Dosage Form; Route</u>	<u>Strength (Container Size)</u>	<u>Docket Number</u>	<u>Petitioner</u>	<u>Reason for Petition</u>	<u>Status</u>
Chlorpheniramine Maleate; Phenylpropanolamine Hydrochloride Controlled-release Capsule; Oral	10mg 75mg	85 P-0149/CP	Dura Pharmaceuticals	New Strength	Approved Dec 13, 1985
Chlorhexidine Gluconate Solution; Topical	1.5%	84 P-0417/CP	Parke Davis	New Strength	Approved Sep 18, 1985
Codeine Phosphate; Dexbrompheniramine Maleate; Phenylpropanolamine Hydrochloride Syrup; Oral	10mg/5ml 1mg/5ml 12.5mg/5ml	85 P-0269/CP	Bock Pharmacal	New Combination	Approved Dec 6, 1985
Dexbrompheniramine Maleate; Phenylpropanolamine Hydrochloride Time Release Capsule; Oral	6mg 75mg	85 P-0238/ CP0002	Bock Pharmacal	New Combination	Approved Dec 13, 1985
Dexbrompheniramine Maleate; Pseudoephedrine Hydrochloride Sustained Release Capsule; Oral	6mg 120mg	85 P-0140/CP	Central Pharms	New Combination New Dosage Form	Approved Dec 13, 1985

(continued)

APPENDIX 4

I. Petitions Approved

(continued)

<u>Drug Name Dosage Form; Route</u>	<u>Strength (Container Size)</u>	<u>Docket Number</u>	<u>Petitioner</u>	<u>Reason for Petition</u>	<u>Status</u>
Dexbrompheniramine Maleate; Pseudoephedrine Sulfate Sustained Release Capsule; Oral	6mg 120mg	85 P-0140/ CP0002	Central Pharms	New Dosage Form	Approved Jan 22, 1986
Diazepam Solution; Oral	5mg/5ml	85 P-0090/CP	Roxane Laboratories	New Dosage Form	Approved Sep 19, 1985
Diazepam Syrup; Oral	2mg/5ml	85 P-0499/CP	Carolina Med Prods	New Dosage Form	Approved Feb 28, 1986
Diazepam Intenso Solution (Concentrate); Oral	5mg/ml	85 P-0566/CP	Roxane Laboratories	New Dosage Form	Approved Mar 18, 1986
Diphenhydramine Hydrochloride Concentrate; Oral	50mg/ml	84 P-0174/CP	Roxane Laboratories	New Strength	Approved Sep 11, 1985
Disopyramide Phosphate Tablet, Controlled Release; Oral	200mg 300mg	84 N-0116/CP	Biocraft Labs	New Dosage Form New Strength	Approved Jun 03, 1986
Disulfiram Suspension; Oral	500mg/30ml	85 P-0215/CP	Paddock Laboratories	New Dosage Form	Approved Oct 8, 1985

(continued)

APPENDIX 4

I. Petitions Approved

<u>Drug Name Dosage Form; Route</u>	<u>Strength (Container Size)</u>	<u>Docket Number</u>	<u>Petitioner</u>	<u>Reason for Petition</u>	<u>Status</u>
Estradiol Tablet; Oral	0.5mg	84 P-0308/CP	Key Pharmaceuticals	New Strength	Approved Mar 24, 1986
Fluorouracil Injectable; Injection	25mg/ml	85 P-0208/CP	Intl Pharm Prods	New Strength	Approved Oct 8, 1985
Fluorouracil Injectable; Injection	50mg/20ml	86 P-0080/CP	Ben Venue Labs	New Strength	Approved Apr 2, 1986
Flurazepam Concentrate; Oral	30mg/ml	85 P-0081/CP	Roxane Laboratories	New Dosage Form	Approved Jul 10, 1985
Flurazepam Hydrochloride Solution; Oral	15mg/5ml	85 P-0091/CP	Roxane Laboratories	New Dosage Form	Approved Oct 25, 1985
Furosemide Solution; Oral	40mg/5ml	85 P-0106/ CP0002	Roxane Laboratories	New Strength	Approved Sep 19, 1985
Furosemide Concentrate; Oral	80mg/ml	85 P-0106/CP	Roxane Laboratories	New Strength	Approved Sep 19, 1985
Haloperidol Solution; Oral	2mg/5ml	85 P-0076/ CP0002	Roxane Laboratories	New Strength	Approved Mar 26, 1986

(continued)

APPENDIX 4

I. Petitions Approved

(continued)

<u>Drug Name Dosage Form; Route</u>	<u>Strength (Container Size)</u>	<u>Docket Number</u>	<u>Petitioner</u>	<u>Reason for Petition</u>	<u>Status</u>
Haloperidol Solution; Oral	5mg/5ml	85 P-0080/CP	Roxane Laboratories	New Strength	Approved Sep 19, 1985
Hydralazine Hydrochloride Solution; Oral	25mg/5ml	85 P-0074/CP	Roxane Laboratories	New Dosage Form	Approved Jul 3, 1985
Ibuprofen Capsule; Oral	200mg	84 P-0383/CP	Sterling Drug	New Dosage Form	Approved Jun 25, 1985
Ibuprofen Soft Gelatin Capsule; Oral	300mg 400mg 600mg	85 P-0563/CP	Softan	New Dosage Form	Approved Mar 19, 1986
Indomethacin Suspension; Oral	25mg/5ml	85 P-0077/ CP0002	Roxane Laboratories	New Dosage Form	Approved Jul 19, 1985
Isoniazid Concentrate; Oral	50mg/ml	85 P-0468/CP	Carolina Med Prods	New Strength	Approved Dec 13, 1985
Ketoconazole Suspension; Oral	20mg/ml	85 P-0147/CP	Janssen Pharma	New Dosage Form	Approved Sep 27, 1985

(continued)

APPENDIX 4

I. Petitions Approved

<u>Drug Name Dosage Form; Route</u>	<u>Strength (Container Size)</u>	<u>Docket Number</u>	<u>Petitioner</u>	<u>Reason for Petition</u>	<u>Status</u>
Leucovorin Calcium Tablet; Oral	15mg	85 P-0487/CP	Lederle Labs/AM Cyan	New Strength	Approved Jan 28, 1986
Meperidine Hydrochloride Concentrate; Oral	100mg/ml	84 P-0175/CP	Roxane Laboratories	New Strength	Approved Jun 7, 1985
Metaproterenol Sulfate Solution; Inhalation	10mg/2.5ml	85 P-0509/CP	Armour Pharm	New Strength	Approved Feb 28, 1986
Metaproterenol Sulfate Solution; Inhalation	10mg/3ml	85 P-0429/ CP0002	Dey Laboratories	New Strength	Approved Feb 28, 1986
Metaproterenol Sulfate Solution; Inhalation	15mg/3ml	85 P-0429/CP	Dey Laboratories	New Strength	Approved Feb 28, 1986
Metoclopramide Hydrochloride Injectable; Injection	5mg/ml 20ml/vial	86 P-0036/CP	AM Critical Care/AHS	New Strength	Approved Mar 18, 1986
Metoclopramide Hydrochloride Injectable; Injection	5mg/ml 50ml/vial	85 P-0545/CP	Lyphomed	New Strength	Approved Feb 28, 1986
Metoclopramide Hydrochloride Injectable; Injection	5mg/ml 50ml/vial	85 P-0540/CP	Quad Pharms	New Strength	Approved Feb 28, 1986

(continued)

APPENDIX 4

I. Petitions Approved

(continued)

<u>Drug Name Dosage Form; Route</u>	<u>Strength (Container Size)</u>	<u>Docket Number</u>	<u>Petitioner</u>	<u>Reason for Petition</u>	<u>Status</u>
Metoclopramide Hydrochloride 5mg/ml Injectable; Injection	5mg/ml 100ml/vial	85 P-0540/CP	Quad Pharm	New Strength	Approved Feb 28, 1986
Methyldopate Hydrochloride Injectable; Injection	50mg/ml 10ml/vial	85 P-0404/CP	AM Critical Care/AHS	New Strength	Approved Oct 25, 1985
Methyltestosterone Capsule; Oral	25mg	85 P-0067/CP	Star Pharmaceuticals	New Dosage Form	Approved Aug 23, 1985
Miconazole Nitrate Cream; Vaginal	4%	84 P-0398/CP	Ortho Pharmaceutical	New Strength	Approved Mar 31, 1986
Naloxone Hydrochloride Injectable; Injection	1mg/ml 5ml/vial	86 P-0079/CP	AM Critical Care/AHS	New Strength	Approved May 7, 1986
Naloxone Hydrochloride Injectable; Injection	1mg/ml 10ml/vial	86 P-0079/CP	AM Critical Care/AHS	New Strength	Approved May 7, 1986
Nitroglycerin Injectable; Injection	5mg/ml	86 P-0025/CP	Lyphomed	New Strength	Approved Apr 1, 1986
Nitroglycerin Injectable; Injection	10mg/ml	85 P-0134/CP	Marion Laboratories	New Strength	Approved Sep 19, 1985
Nitroglycerin in 5% Dextrose Injectable; Injection	10mg/100ml (500ml Container)	86 P-0099/CP	Abbott Laboratories	New Strength	Approved Apr 1, 1986

(continued)

APPENDIX 4

I. Petitions Approved

(continued)

<u>Drug Name Dosage Form; Route</u>	<u>Strength (Container Size)</u>	<u>Docket Number</u>	<u>Petitioner</u>	<u>Reason for Petition</u>	<u>Status</u>
Nitroglycerin in 5% Dextrose Injectable; Injection	2.0mg/100ml (250ml Container)	86 P-0099/ CP0002	Abbott Laboratories	New Strength	Approved Apr 1, 1986
Nitroglycerin in 5% Dextrose Injectable; Injection	4.0mg/100ml (250ml and 500ml Containers)	86 P-0099/ CP0003	Abbott Laboratories	New Strength	Approved Apr 1, 1986
Probucol Tablet; Oral	500mg	85 P-0337/CP	Merrell Dow/Dow Chem	New Strength	Approved Oct 25, 1985
Procainamide Hydrochloride Tablet; Oral	375mg	85 P-0125/CP	Key Pharmaceuticals	New Strength	Approved Sep 19, 1985
Propranolol Hydrochloride Capsule; Oral	10mg 20mg 40mg 60mg 80mg 90mg	86 P-0045/CP	Nutripharm Labs	New Dosage Form	Approved Mar 19, 1986

(continued)

APPENDIX 4

I. Petitions Approved

(continued)

<u>Drug Name Dosage Form; Route</u>	<u>Strength (Container Size)</u>	<u>Docket Number</u>	<u>Petitioner</u>	<u>Reason for Petition</u>	<u>Status</u>
Propranolol Hydrochloride Solution; Oral	40mg/5ml	85 P-0073/CP	Roxane Laboratories	New Dosage Form	Approved Jul 8, 1985
Propranolol Hydrochloride Concentrate; Oral	80mg/ml	85 P-0073/CP0002	Roxane Laboratories	New Dosage Form	Approved Jul 19, 1985
Propranolol Hydrochloride Solution; Oral	20mg/5ml	85 P-0073/CP0003	Roxane Laboratories	New Dosage Form	Approved Sep 24, 1985
Propranolol Hydrochloride Tablet, Constant-Release; Oral	160mg	85 P-0129/CP	Verex Laboratories	New Dosage Form	Approved Sep 25, 1985
Propranolol Hydrochloride Tablet, Controlled Release; Oral	80mg 120mg 160mg	85 P-0197/CP	Forest Laboratories	New Dosage Form	Approved Sep 27, 1985
Pyridostigmine Bromide Tablet; Oral	30mg	85 P-0412/CP	Kali-Duphar Labs	New Strength	Approved Jan 22, 1986

(continued)

APPENDIX 4

I. Petitions Approved

(continued)

<u>Drug Name Dosage Form; Route</u>	<u>Strength (Container Size)</u>	<u>Docket Number</u>	<u>Petitioner</u>	<u>Reason for Petition</u>	<u>Status</u>
Ritodrine Hydrochloride in Dextrose 5% Injectable; Injection	30mg/100ml 500ml Container	86 P-0100/CP	Abbott Laboratories	New Strength	Approved May 7, 1986
Scopolamine Transdermal System/24 Hour Film, Controlled Release; Percutaneous	1mg	85 P-0168/CP	Ciba Consumer Pharmas	New Strength (Dosing Interval)	Approved Sep 27, 1985
Spironolactone Syrup; Oral	25mg/5ml	85 P-0510/CP	Carolina Med Prods	New Dosage Form	Approved Jan 22, 1986
Spironolactone Oral; Injection	25mg/5ml	86 P-0055/CP	Carolina Med Prods	New Dosage Form	Approved Mar 28, 1986
Theophylline Capsule; Oral	150mg 300mg	85 P-0175/CP	Mead Johnson/B-M	New Strength	Approved Oct 8, 1985
Thiothixene Hydrochloride Solution; Oral	5mg/5ml	86 P-0178/CP	Eliis Pharmaceutical	New Strength	Approved Jun 04, 1986

(continued)

APPENDIX 4

I. Petitions Approved

(continued)

<u>Drug Name Dosage Form; Route</u>	<u>Strength (Container Size)</u>	<u>Docket Number</u>	<u>Petitioner</u>	<u>Reason for Petition</u>	<u>Status</u>
Vinblastine Sulfate Injectable; Injection	1mg/ml	86 P-0056/CP	Quad Pharm	New Dosage Form	Approved Mar 28, 1986
Vincristine Sulfate Injectable; Injection	2mg/vial	85 P-0016/CP	Bristol Labs/B-M	New Dosage Form	Approved Nov 8, 1985

APPENDIX 4

II. Petitions Denied

<u>Drug Name Dosage Form; Route</u>	<u>Strength (Container Size)</u>	<u>Docket Number</u>	<u>Petitioner</u>	<u>Reason for Petition</u>	<u>Status</u>
Acetaminophen; Hydrocodone Bitartrate Tablet; Oral	650mg 10mg	85 P-0015/CP	Applied Labs	New Strength	Denied Nov 7, 1985
Acetaminophen; Hydrocodone Bitartrate Tablet; Oral	750mg 7.5mg	85 P-0169/CP	Knoll Pharmaceutical	New Strength	Denied Nov 7, 1985
Aminocaproic Acid Injectable; Injection	500mg/ml	85 P-0064/CP	Abbott Laboratories	New Strength	Denied May 29, 1985
Aminophylline Injectable; Injection	10mg/ml	85 P-0066/CP	Abbott Laboratories	New Strength	Denied May 3, 1985
Aminophylline Injectable; Injection	50mg/ml	85 P-0066/CP	Abbott Laboratories	New Strength	Denied May 3, 1985
5-Aminosalicylic Acid Suppository; Rectal	500mg	84 P-0425/CP	Reid-Rowell	New Ingredient	Denied Jun 05, 1986

(continued)

APPENDIX 4

II. Petitions Denied

(continued)

<u>Drug Name Dosage Form; Route</u>	<u>Strength (Container Size)</u>	<u>Docket Number</u>	<u>Petitioner</u>	<u>Reason for Petition</u>	<u>Status</u>
Aspirin; Chloroxazone Tablets; Oral	325mg 250mg	85 P-0071/CP	McNeil Pharm	New Combination	Denied Sep 3, 1985
Aspirin; Butalbital; Caffeine; Codeine Phosphate Capsule; Oral	325mg 50mg 40mg 7.5mg	85 P-0101/CP	Sandoz Pharms/Sandoz	New Combination	Denied Sep 11, 1985
Aspirin; Butalbital; Caffeine; Codeine Phosphate Capsule; Oral	325mg 50mg 40mg 15mg	85 P-0101/CP	Sandoz Pharms/Sandoz	New Combination	Denied Sep 11, 1985

(continued)

APPENDIX 4

II. Petitions Denied

(continued)

<u>Drug Name Dosage Form; Route</u>	<u>Strength (Container Size)</u>	<u>Docket Number</u>	<u>Petitioner</u>	<u>Reason for Petition</u>	<u>Status</u>
Aspirin; Butalbital; Caffeine; Codeine Phosphate Capsule; Oral	325mg 50mg 40mg 30mg	85 P-0101/CP CP0002	Sandoz Pharms/Sandoz	New Combination	Denied Sep 11, 1985
Aspirin; Butalbital; Caffeine; Codeine Phosphate Capsule; Oral	325mg 50mg 40mg 60mg	85 P-0101/ CP0002	Sandoz Pharms/Sandoz	New Combination	Denied Sep 11, 1985
Benzoyl Metronidazole Suspension; Injection	200mg/5ml	85 P-0258/CP	Apkon Laboratories	New Ester New Ingredient	Denied Mar 19, 1986
Betamethasone Dipropionate Miconazole Nitrate Cream; Topical	0.05% 2%	85 P-0271/CP	Ortho Pharmaceutical	New Combination	Denied Apr 18, 1986

(continued)

APPENDIX 4

II. Petitions Denied

(continued)

<u>Drug Name Dosage Form; Route</u>	<u>Strength (Container Size)</u>	<u>Docket Number</u>	<u>Petitioner</u>	<u>Reason for Petition</u>	<u>Status</u>
Bretylium Tosylate Injectable; Injection	2mg/ml	85 P-0063/CP	Abbott Laboratories	New Strength	Denied May 29, 1985
Bretylium Tosylate Injectable; Injection	4mg/ml	85 P-0063/ CP0002	Abbott Laboratories	New Strength	Denied May 29, 1985
Bretylium Tosylate Injectable; Injection	8mg/ml	85 P-0063/ CP0003	Abbott Laboratories	New Strength	Denied May 29, 1985
Bretylium Tosylate Injectable; Injection	10mg/ml	85 P-0063/ CP0004	Abbott Laboratories	New Strength	Denied May 29, 1985
Caffeine; Ergotamine Tartrate; Pentobarbital Tablet; Oral	100mg 1mg 30mg	85 P-0433/CP	Sandoz Pharms/Sandoz	New Combination	Denied Nov 8, 1985
Caffeine; Ergotamine Tartrate; Pentobarbital Sodium Suppository; Rectal	200mg 2mg 60mg	85 P-0433/ CP0002	Sandoz Pharms/Sandoz	New Combination	Denied Nov 8, 1985

(continued)

APPENDIX 4

II. Petitions Denied

(continued)

<u>Drug Name Dosage Form; Route</u>	<u>Strength (Container Size)</u>	<u>Docket Number</u>	<u>Petitioner</u>	<u>Reason for Petition</u>	<u>Status</u>
Cholecalciferol Capsule; Oral	1.25mg	84 P-0161/CP	Pharmacaps	New Active Ingredient	Denied Feb 13, 1986
Codeine Phosphate; Ibuprofen Capsule; Oral	30mg 200mg	84 P-0388/CP	McNeil Pharm	New Combination	Denied Sep 16, 1985
Codeine Phosphate; Ibuprofen Capsule; Oral	60mg 200mg	84 P-0388/CP	McNeil Pharm	New Combination	Denied Sep 16, 1985
Codeine Phosphate; Ibuprofen Tablet; Oral	30mg 200mg	84 P-0388/CP	McNeil Pharm	New Combination	Denied Sep 16, 1985
Codeine Phosphate; Ibuprofen Tablet; Oral	60mg 200mg	84 P-0388/CP	McNeil Pharm	New Combination	Denied Sep 16, 1985
Choline Magnesium Trisalicylate Codeine Phosphate Tablet; Oral	500mg 30mg	85 P-0142/CP	Purdue Frederick	New Combination	Denied Jul 21, 1986

(continued)

APPENDIX 4

<u>Drug Name Dosage Form; Route</u>	<u>Strength (Container Size)</u>	<u>Docket Number</u>	<u>Petitioner</u>	<u>Reason for Petition</u>	<u>Status</u>
Choline Magnesium Trisalicylate; Codeine Phosphate Tablet; Oral	500mg 60mg	85 P-0142/CP	Purdue Frederick	New Combination	Denied Jul 21, 1986
Dextromethorphan Hydrobromide Tablet, Controlled Release; Oral	60mg	85 P-0135/CP	Ciba Consumer Pharm	New Salt New Ingredient	Denied Jul 17, 1986
Diatrizoate Meglumine; Lidocaine Hydrochloride Injectable; Injection	60% 1.5mg/ml	84 P-0325/CP	Cook Imaging	New Combination	Denied Sep 3, 1985
Diazepam Intensol Concentrate; Oral	10mg/ml	85 P-0075/CP	Roxane Laboratories	New Dosage Form	Denied Sep 24, 1985
Tri-Phasic Contraceptive Tablet; Oral(21 and 28 days) Ethinyl Estradiol Norethindrone	0.05mg 0.5mg	84 P-0443/CP	Ortho Pharmaceutical	New Strength (Dose Schedule)	Denied Sep 3, 1985
Ethinyl Estradiol Norethindrone	0.05mg 0.75mg				
Ethinyl Estradiol Norethindrone	0.05mg 1.0mg				

(continued)

APPENDIX 4

II. Petitions Denied

(continued)

<u>Drug Name Dosage Form; Route</u>	<u>Strength (Container Size)</u>	<u>Docket Number</u>	<u>Petitioner</u>	<u>Reason for Petition</u>	<u>Status</u>
Fluphenazine Hydrochloride Injectable; Injection	5mg/ml	85 P-0019/CP	ER Squibb and Sons	New Strength	Denied Oct 25, 1985
Heparin Sodium Injectable; Injection	2000 Units/ml	85 P-0065/CP	Abbott Laboratories	New Strength	Denied May 29, 1985
Heparin Sodium Injectable; Injection	4000 Units/ml	85 P-0065/CP	Abbott Laboratories	New Strength	Denied May 29, 1985
Hydrochlorothiazide; Propranolol Hydrochloride; Triamterene Capsule, Controlled Release; Oral	50mg 80mg 75mg	85 P-0571/CP	Ayerst Labs/AMHO	New Combination	Denied May 16, 1986
Hydrochlorothiazide; Propranolol Hydrochloride; Triamterene Capsule, Controlled Release; Oral	50mg 120mg 75mg	85 P-0571/CP	Ayerst Labs/AMHO	New Combination	Denied May 16, 1986
Hydrochlorothiazide; Propranolol Hydrochloride; Triamterene Capsule, Controlled Release;	50mg 160mg 75mg	85 P-0571/CP	Ayerst Labs/AMHO	New Combination	Denied May 16, 1986

(continued)

APPENDIX 4

II. Petitions Denied

(continued)

<u>Drug Name Dosage Form; Route</u>	<u>Strength (Container Size)</u>	<u>Docket Number</u>	<u>Petitioner</u>	<u>Reason for Petition</u>	<u>Status</u>
Ibuprofen; Oxycodone Hydrochloride Capsule; Oral Oral	200mg 5mg	85 P-0141/CP	Dupont Pharms/Dupont	New Combination	Denied Sep 27, 1985
Ibuprofen; Oxycodone Hydrochloride Tablet; Oral	200mg 5mg	85 P-0141/CP	Dupont Pharms/Dupont	New Combination	Denied Sep 27, 1985
Indomethacin Tablet; Oral	25mg	85 P-0025/CP	Verex Laboratories	New Dosage Form	Denied Mar 31, 1986
Indomethacin Tablet; Oral	50mg	85 P-0025/CP	Verex Laboratories	New Dosage Form	Denied Mar 31, 1986
Indomethacin Intenso Solution (Concentrate);	50mg/ml	85 P-0077/CP	Roxane Laboratories	New Dosage Form New Strength	Denied Apr 7, 1986
Indomethacin Tablet, Constant Release; Oral	75mg	85 P-0026/CP	Verex Laboratories	New Dosage Form	Denied Sep 16, 1985
Indomethacin Controlled-release Tablet; Oral	75mg	85 P-0180/CP	Forest Laboratories	New Dosage Form	Denied Apr 7, 1986

(continued)

APPENDIX 4

II. Petitions Denied

(continued)

<u>Drug Name Dosage Form; Route</u>	<u>Strength (Container Size)</u>	<u>Docket Number</u>	<u>Petitioner</u>	<u>Reason for Petition</u>	<u>Status</u>
Methocarbamol Acetaminophen Tablet; Oral	400mg 325mg	85 P-0102/CP	McNeil Pharm	New Combination	Denied Jun 24, 1986
Metoclopramide Hydrochloride 1mg/ml Injectable; Injection	50ml/vial	86 P-0015/CP	Intl Medication Sys	New Strength	Denied Apr 25, 1986
Metoclopramide Hydrochloride 1mg/ml Injectable; Injection	75ml/vial	86 P-0015/CP	Intl Medication Sys	New Strength	Denied Apr 25, 1986
Metoclopramide Hydrochloride 1mg/ml Injectable; Injection	100ml/vial	86 P-0015/CP	Intl Medication Sys	New Strength	Denied Apr 25, 1986
Metoclopramide Hydrochloride 10mg/ml Injectable; Injection		85 P-0062/CP	Abbott Laboratories	New Strength	Denied May 29, 1985
Metoclopramide Hydrochloride 10mg/ml Injectable; Injection		85 P-0457/CP	Abbott Laboratories	New Strength	Denied Apr 18, 1986
Metoclopramide Hydrochloride 20mg/ml Injectable; Injection		85 P-0062/ CP0002	Abbott Laboratories	New Strength	Denied May 29, 1985
Metoclopramide Hydrochloride 20mg/ml Injectable; Injection		85 P-0457/ CP0002	Abbott Laboratories	New Strength	Denied Apr 18, 1986
Metronidazole Sponge; Vaginal	50-125mg/ Sponge	85 P-0117/CP	VLI	New Dosage Form	Denied Oct 8, 1985

(continued)

APPENDIX 4

II. Petitions Denied

(continued)

<u>Drug Name Dosage Form; Route</u>	<u>Strength (Container Size)</u>	<u>Docket Number</u>	<u>Petitioner</u>	<u>Reason for Petition</u>	<u>Status</u>
Nitroglycerin Transdermal System	None Given	84 P-0302/CP	Key Pharmaceuticals	New Dosage Form (New Matrix)	Denied Jul 29, 1985
Phenylephrine Hydrochloride; Sulfathiazole Nasal Suspension; Topical	0.5% 5%	85 P-0205/CP	Tanya W Ross	New Dosage Form New Combination	Denied Nov 14, 1985
Pseudoephedrine Polisterex Controlled Release Capsule; Oral	60mg	85 P-0334/CP	Pennwalt Pharm	New Salt New Ingredient	Denied Mar 19, 1986
Triamcinolone Acetonide Suspension; Injection	2.5mg/ml	85 P-0001/CP	GenDerm	New Strength	Denied Mar 4, 1985
Triamcinolone Acetonide Suspension; Injection	3mg/ml	84 P-0240/CP	Bay Pharmaceuticals	New Strength	Denied Mar 4, 1985

APPENDIX 5
EXCLUSIVITY TERMS

DUE TO SPACE LIMITATIONS IN THE EXCLUSIVITY COLUMN, THE FOLLOWING ABBREVIATIONS HAVE BEEN DEVELOPED. PLEASE REFER BACK TO THIS PAGE FOR AN EXPLANATION OF THE EXCLUSIVITY ABBREVIATIONS FOUND IN THE ADDENDUM.

ABBREVIATIONS

NC	NEW COMBINATION
NCE	NEW CHEMICAL ENTITY
NDF	NEW DOSAGE FORM
NE	NEW ESTER OR SALT OF AN ACTIVE INGREDIENT
NP	NEW PRODUCT
NR	NEW ROUTE
PP	PARENTERAL IN PLASTIC CONTAINER
RTO	PREScription TO OTC STATUS CHANGE
NS	NEW STRENGTH
D	NEW DOSING SCHEDULE (SEE REFERENCE, BELOW)
I	NEW INDICATION (SEE REFERENCE, BELOW)
ODE	ORPHAN DRUG EXCLUSIVITY

REFERENCES

<u>NEW DOSING SCHEDULE</u>	
D-1	ONCE A DAY APPLICATION
D-2	ONCE DAILY DOSING
D-3	SEVEN DAYS/SEVEN DAYS/SEVEN DAYS DOSING SCHEDULE
D-4	SEVEN DAYS/FOURTEEN DAYS DOSING SCHEDULE
D-5	TEN DAYS/ELLEVEN DAYS DOSING SCHEDULE
D-6	SEVEN DAYS/NINE DAYS/FIVE DAYS DOSING SCHEDULE
D-7	BID DOSING

(continued)

APPENDIX 5

(continued)

NEW DOSING SCHEDULE

- D-8 INTRAVENOUS, EPIDURAL AND INTRATHECAL DOSING
- D-9 NARCOTIC OVERDOSE IN ADULTS
- D-10 NARCOTIC OVERDOSE IN CHILDREN
- D-11 POSTOPERATIVE NARCOTIC DEPRESSION IN CHILDREN
- D-12 BEDTIME DOSING OF 800MG FOR TREATMENT

NEW INDICATION

- I-1 SEVERE HYPERTENSION IN PEDIATRICS AND NON-MALIGNANT HYPERTENSION
- I-2 DYSMENORRHEA
- I-3 TREATMENT OF TINEA VERSICOLOR
- I-4 SYMPTOMATIC GASTROESOPHAGEAL REFLUX
- I-5 NEPHROTOMOGRAPHY
- I-6 CONTRAST ENHANCEMENT IN CRANIAL COMPUTED TOMOGRAPHY
- I-7 VENOGRAPHY OF LOWER EXTREMITIES
- I-8 WHOLE-BODY COMPUTED TOMOGRAPHY
- I-9 GATED CARDIAC POOL IMAGING
- I-10 POST-MYOCARDIAL INFARCTION
- I-11 COLORECTAL SURGERY
- I-12 NAUSEA AND VOMITING ASSOCIATED WITH EMETOGENIC CANCER CHEMOTHERAPY
- I-13 CISPLATIN INDUCED EMESIS
- I-14 DIABETIC GASTROPARESIS
- I-15 SHORT TERM TREATMENT OF GASTRIC ULCER DISEASE
- I-16 ACROMEGALY

(continued)

APPENDIX 5

(continued)

NEW INDICATION

- I-17 PITUITARY TUMORS
 I-18 POSTMENOPAUSAL OSTEOPOROSIS
 I-19 ANTIDOTE FOR ACETAMINOPHEN OVERDOSAGE
 I-20 CONGESTIVE HEART FAILURE BID DOSAGE SCHEDULE
I-21 ACUTE/OBTURATE/MEDIA
 I-22 EXERCISE INDUCED BRONCHOSPASMS
 I-23 MYOCARDIAL INFARCTION OR STROKE
 I-24 COMBINED USE WITH NICOTINIC ACID TO LOWER CHOLESTEROL LEVEL
 BLASTOMYCOSES DERMATITIDES
 I-25 PEDIATRIC SUBARACHNOID VASCULAR
 I-26 PETRIELLIUM BOYDII INFECTION
 HEREDITARY ANGIOEDEMA
 I-27 INTRACORONARY USE
 I-28 PEDIATRIC USE
 I-29 DIRECT ISOTOPIC CYSTOGRAPHY
 I-30 POSTPARTUM HEMORRHAGE
 I-31 USE IN METHODONE INDUCED RESPIRATORY DEPRESSION
 I-32 PROLACTIN SECRETING ADENOMAS
 I-33 ANGINA PECTORIS DUE TO CORONARY ATHEROSCLEROSIS
 I-34 ARTERIAL DIGITAL SUBTRACTION ANGIOGRAPHY
 I-35 SPINAL ANESTHESIA
 I-36 PATIENT PREOPERATIVE SKIN PREPARATION
 I-37 ADJUVANT WITH CHEMOTHERAPY FOR TREATMENT OF BREAST CANCER FOLLOWING MASTECTOMY
 I-38 ANTIDOTE FOR ACETAMINOPHEN OVERDOSE
 I-39 MANAGEMENT OF HYPOCALCEMIA AND RESULTANT METABOLIC BONE DISEASE IN RENAL DIALYSIS PATIENTS
 I-40 MAINTENANCE THERAPY AT REDUCED DOSE FOLLOWING HEALING OF ACUTE DUODENAL ULCER
 I-41 TREATMENT OF GASTROESOPHAGEAL REFLUX DISEASE
 I-42 TREATMENT OF SEVERE RECALCITRANT DERMATOPHYTE INFECTIONS
 I-43
 I-44

APPENDIX 6
PREScription AND OTC DRUG PRODUCT
PATENT AND EXCLUSIVITY DATA

78

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

NO SEPTEMBER - JULY ACTIONS

PREScription AND OTC DRUG PRODUCT
PATENT AND EXCLUSIVITY DATA

APPL/PROD	PATENT NUMBER	PATENT EXPIRES	EXCLUSIVITY CODE	EXCLUSIVITY EXPIRES	APPL/PROD	PATENT NUMBER	PATENT EXPIRES	EXCLUSIVITY CODE	EXCLUSIVITY EXPIRES
/16142/001/	/4537883/	AUG/6/	/602/	/16243/003/	/4324779/	/APR/13/1999/	/4324779/	/APR/13/1999/	SEP 24, 1986
/16142/002/	/4537883/	AUG/6/	/602/	/16243/003/	/4324779/	/APR/13/1999/	/4324779/	/APR/13/1999/	SEP 24, 1986
/16142/003/	/4537883/	AUG/6/	/602/	/16243/003/	/4324779/	/APR/13/1999/	/4324779/	/APR/13/1999/	SEP 24, 1986
/16142/004/	/4537883/	AUG/6/	/602/	/16243/003/	/4324779/	/APR/13/1999/	/4324779/	/APR/13/1999/	SEP 24, 1986
/16142/005/	/4537883/	AUG/6/	/602/	/16243/003/	/4324779/	/APR/13/1999/	/4324779/	/APR/13/1999/	SEP 24, 1986
/16142/006/	/4537883/	AUG/6/	/602/	/16243/003/	/4324779/	/APR/13/1999/	/4324779/	/APR/13/1999/	SEP 24, 1986
12142 006	4537883	AUG 27,	2002	16983 001	3634582	JAN 11, 1989			SEP 09, 1988
12142 007	4537883	AUG 27,	2002	16990 001	3860618	JAN 14, 1992			I-36
12142 008	4537883	AUG 27,	2002		RE28636	JUN 02, 1987	/1-21/	/1-21/	/SEP 24, 1986/
12142 009	4537883	AUG 27,	2002		17560 002	RE28636	/1-21/	/1-21/	/SEP 24, 1986/
12142 010	4537883	AUG 27,	2002		17560 001	RE28636	/1-21/	/1-21/	/SEP 24, 1986/
12365 005	4534973	AUG 13,	2002		17581 001	3998966	DEC 21, 1993		
12366 002	4534974	AUG 13,	2002		17601 001	/419565/	/DEC 31, 1995/		
13601 001						/371764/	/FEB 20, 1996/		
13601 002						/383957/	/OCT 01, 1991/		
/14715/601/	/3428735/	/FEB/18/	/1986/		17613 001	/17619 001	/OCT 01, 1991/		
14715 004	3428735	FEB 18,	1986			/383957/	/OCT 01, 1991/		
/16275/601/	/4324779/	/APR/13/	/1999/		/17648/661/	/4324779/	/APR 13, 1999/		
/16275/602/	/4324779/	/APR/13/	/1999/		17717 001	/3839573/	/OCT 01, 1991/		
					17760 001				NDF
									SEP 04, 1988

(continued)

**APPENDIX 6
PRESCRIPTION AND OTC DRUG PRODUCT
PATENT AND EXCLUSIVITY DATA**

APPL/PROD	PATENT NUMBER	PATENT EXPIRES	EXCLUSIVITY CODE	EXCLUSIVITY EXPIRES	APPL/PROD	PATENT NUMBER	PATENT EXPIRES	EXCLUSIVITY CODE	EXCLUSIVITY EXPIRES
17768 001	3855140	DEC 17, 1991	I-38	SEP 24, 1986	18240 001				I-35
17785 001	3960745	DEC 17, 1991	NDF	MAR 07, 1989	18240 002	/46457/001/	/DEC/04/;1991/		SEP 04, 1988
17862 001	4555336	AUG 20, 2002	I-12	SEP 24, 1986	18257 001	/4237068/001/	/DEC/04/;1991/		SEP 04, 1988
17862 002	1536756	AUG 20, 2002	I-13	SEP 24, 1986	18257 002	/4237068/001/	/DEC/04/;1991/		
17862 003	4536386	AUG 20, 2002	I-14	SEP 24, 1986	18257 002	/4237068/001/	/DEC/04/;1991/		
17920 005	3950333	APR 13, 1993	I-12	SEP 24, 1986	18257 002	/4237068/001/	/DEC/04/;1991/		
17970 001	4024271	MAY 17, 1994	D-12	APR 30, 1989	18509 001				
17970 001	4536516	AUG 20, 2002	I-39	DEC 10, 1988	18509 001				
18044 001			I-41	JAN 22, 1989	18513 002				
18044 002			I-41	JAN 22, 1989	18533 001				
18052 001	/4639573/	/DEC/01/;1991/	I-37	SEP 25, 1988	18587 003	3658993	APR 25, 1989	NP	AUG 07, 1988
18053 003					18644 001	3819706	JUN 25, 1991	OEE	JUL 28, 1990
18147 002	/4624668/	/DEC/10/;1991/	I-37	SEP 25, 1988	18644 001	3885046	MAY 20, 1992	I-44	JUN 30, 1989
18154 001	/4100441/	/DEC/11/;1991/	I-37	SEP 25, 1988	4057323	4057323	MAR 26, 2002	NCE	Sep 07, 1992
18154 002	/3927602/	/DEC/16/;1992/	I-37	SEP 25, 1988	4347257	4347257	AUG 31, 1999	NCE	DEC 30, 1990
18154 003	/4F29588/	/DEC/18/;1991/	I-37	SEP 25, 1988	4393078	4393078	JUL 12, 2000		
18155 001	/4103347/	/DEC/11/;1991/	I-37	SEP 25, 1988	4425363	4425363	JAN 10, 2001		
18155 001	/3927602/	/DEC/16/;1992/	I-37	SEP 25, 1988	4435449	4435449	MAR 06, 2001		
18155 001	/46154/661/	/AUG/12/;1986/	I-37	SEP 25, 1988	4438138	4438138	MAR 20, 2001		
18155 001	/3461461/	/AUG/12/;1986/	I-37	SEP 25, 1988	3819706	3819706	JUN 25, 1991		
18155 001	/46154/663/	/AUG/12/;1986/	I-37	SEP 25, 1988	3885046	3885046	MAY 20, 1992		
18155 003	3461461	MAY 07, 1985	I-37	SEP 25, 1988	4057323	4057323	MAR 26, 2002		
18181 001	/4639573/	/DEC/01/;1991/	I-37	SEP 25, 1988	4347257	4347257	AUG 31, 1999		
18182 001	/353573/	/DEC/01/;1991/	I-37	SEP 25, 1988	4393078	4393078	JUL 12, 2000		
18183 001	/363573/	/DEC/01/;1991/	I-37	SEP 25, 1988	4425363	4425363	JAN 10, 2001		
18217 001	4035376	JUL 12, 1994	I-37	SEP 25, 1988	4435449	4435449	MAR 06, 2001		
18230 001	/3639573/	/DEC/01/;1991/	I-37	SEP 25, 1988	4438138	4438138	MAR 20, 2001		

(continued)

APPENDIX 6
PREScription AND OTC DRUG PRODUCT
PATENT AND EXCLUSIVITY DATA

APPL/PROD	PATENT NUMBER	PATENT EXPIRES	EXCLUSIVITY CODE	EXCLUSIVITY EXPIRES	APPL/PROD	PATENT NUMBER	PATENT EXPIRES	EXCLUSIVITY CODE	EXCLUSIVITY EXPIRES
186444 003	3819706	JUN 25, 1991	NCE	DEC 30, 1990	18830	001	3900481	AUG 19, 1992	NCE
	3888046	MAY 20, 1992			18830	001	4005209	JAN 25, 1994	OCT 31, 1990
	4057323	MAR 26, 2002			18830	002	3900481	AUG 19, 1992	NCE
4347257	AUG 31, 1999				18830	002	4005209	JAN 25, 1994	OCT 31, 1990
4393078	JUL 12, 2000				18839	001	4211771	JUL 08, 1997	DEC 31, 1990
4425363	JAN 10, 2001				RE29835		MAR 19, 1991		
4435449	MAR 06, 2001				18873	002	3954872	MAY 04, 1993	NCE
4438138	MAR 20, 2001						4031244	JUN 21, 1994	
4280957	JUL 28, 1998	NCE	DEC 20, 1990		18873	003	3954872	MAY 04, 1993	NCE
4087545	MAY 02, 1995	NCE	DEC 26, 1990				4031244	JUN 21, 1994	DEC 30, 1990
4087547	MAY 02, 1995	/DEC/18, 1993/	/NCE/		18873	004	3954872	MAY 04, 1993	NCE
/18682/661/	/4062966/	DEC 13, 1994	/NCE/		18887	001	4031244	JUN 21, 1994	DEC 30, 1990
18683 001	4062966	JUL 19, 2000					3686412	AUG 22, 1989	NDF
/18682/661/	/3748579/	/JAN/02, 1990/					3777033	AUG 22, 1989	
18689 001	3708579	JAN 02, 1992			18891	001	4559222	DEC 17, 2002	
18701 001	3438991	APR 15, 1986	NE	JAN 14, 1989	18891	002	4559222	DEC 17, 2002	
18703 001			I-42	MAY 30, 1989			4559222	DEC 17, 2002	
18703 001			I-43	MAY 30, 1989	/18691/	/661/	/3857952/	/DEC/31, 1991/	
18703 002	4128658	DEC 05, 1995	NCE	JUN 09, 1993	18917	001	3857952	DEC 31, 1993	
	4521431	JUN 04, 2002		JUN 28, 1988	18917	003	3857952	DEC 31, 1993	
18705 001			I-15	JUN 28, 1988	18928	001	4221778	SEP 09, 1997	
18708 001	3845039	OCT 29, 1991	NDF	OCT 31, 1988	18932	001			ODE
	3920818	NOV 18, 1991			18948	001			NCE
18713 001	/3639573/	/DEC/01, 1991/			/18649/661/	/18649/15/1992/	/DEC/08, 1990	/18649/15/1992/	NCE
18735 001	4001323	JAN 04, 1994	NCE	DEC 31, 1990	18949	001	3878217	APR 15, 1994	NCE
18735 002	4001323	JAN 04, 1994	NCE	DEC 31, 1990			3806546/	APR 25, 1991/	
18735 003	4001323	JAN 04, 1994	NCE	DEC 31, 1990			/1865557/	JUN 24, 1993/	
	18735 004	4001323	JAN 04, 1994	NCE			/3966949/	JUN 29, 1993/	
	18738 001	4055652	OCT 25, 1994	NCE			/4254129/	JUN 29, 1993/	
18754 002	3641127	FEB 08, 1989	NCE	AUG 30, 1990			/4255957/	AUG 25, 1998/	
18754 003	3641127	FEB 08, 1989	NCE	JAN 09, 1991	18956	001	4021481	MAY 03, 1994	DEC 26, 1990
/18677/661/	/4138581/	/FEB/06, 1996/	/NCE/	JAN 09, 1991			4250113	FEB 10, 19980	
18770 001	4138581			/DEC/28, 1989	18956	002	4021481	MAY 03, 1994	DEC 26, 1990
18813 001	/1869574/	/DEC/01, 1991/					4250113	FEB 10, 1998	
18827 001	/3639573/	/DEC/01, 1991/							

(continued)

APPENDIX 6
PRESCRIPTION AND OTC DRUG PRODUCT
PATENT AND EXCLUSIVITY DATA

APPL/PROD	PATENT NUMBER	PATENT EXPIRES	EXCLUSIVITY CODE	EXCLUSIVITY EXPIRES	APPL/PROD	PATENT NUMBER	PATENT EXPIRES	EXCLUSIVITY CODE	EXCLUSIVITY EXPIRES
18956 003	4021481	MAY 03, 1994	NCE	DEC 26, 1990		19219 002	FEB 08, 1989	NCE	DEC 19, 1990
	4250113	FEB 10, 1998				19257 001		NDF	APR 10, 1989
18956 004	4021481	MAY 03, 1994	NCE	DEC 26, 1990		19259 001	SEP 14, 1993	ODE*	DEC 27, 1992
	4250113	FEB 10, 1998				19260 001	SEP 14, 1993		
18972 001						19270 001	4252984	ODE	OCT 16, 1991
18985 001	4544554	JUL 23, 2002				4311708	JAN 19, 1999	NCE	AUG 30, 1990
18985 002	4544554	JUL 23, 2002				4342783	AUG 03, 1999		
18998 001	4374829	FEB 22, 2000	NCE	DEC 24, 1990		19322 001	3721687	NCE	DEC 27, 1990
18998 002	4374829	FEB 22, 2000	NCE	DEC 24, 1990		19323 001	3721687	NCE	DEC 27, 1990
18998 003	4374829	FEB 22, 2000	NCE	DEC 24, 1990		19359 001	4078071	NCE	NOV 25, 1990
19011 001	/4335659/ /4335659/	JUN 15, 1999	NCE/	SEP 24, 1986		19368 001	4205086	NCE	OCT 29, 1990
/19644/661/		JUN 15, 1999	NCE/	/DEC/23//1996/					
19044 001	4335095	JUN 15, 1999	NCE	DEC 23, 1990					
19059 001	4138475	FEB 06, 1996		PETITION FOR EXCLUSIVITY PENDING					
19059 002	4138475	FEB 06, 1996		PETITION FOR EXCLUSIVITY PENDING		19412 001		ODE	OCT 29, 1992
19059 003	4138475	FEB 06, 1996		PETITION FOR EXCLUSIVITY PENDING		19412 002		NS	MAR 10, 1989
19069 001	/4335659/ /4335659/	FEB 06, 1996				19412 003		NS	MAR 10, 1989
19071 001								NS	MAR 10, 1989
19079 001						1942444	MAR 15, 1994	NCE	AUG 01, 1994
19084 001	4335125	JUN 15, 1999	NE	AUG 30, 1988		4066755	JAN 03, 1995		
19107 001			NDF	FEB 11, 1989		19434 001	3950333	APR 13, 1993	
19107 001				DEC 31, 1988			4024271	MAY 17, 1994	
19107 001			NCE	OCT 17, 1990			4024163	MAY 17, 1994	
19194 001			ODE	OCT 17, 1992					
			NCE	NOV 11, 1990					
19215 001	4078071	MAR 07, 1995	ODE	NOV 11, 1992		19478 001	3644627	FEB 22, 1989	
			NCE	NOV 25, 1990			3784684	JAN 08, 1991	

*REFER TO APPENDIX I NARRATIVE