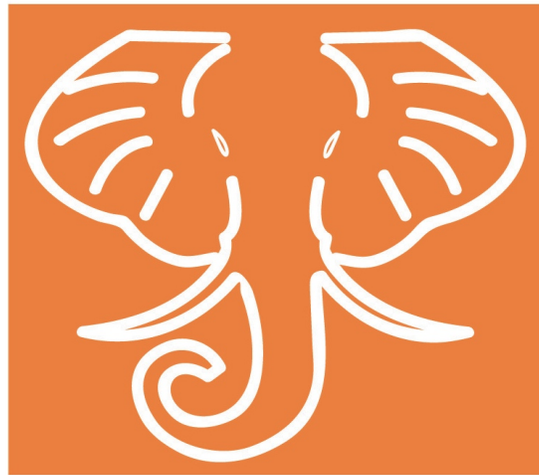


## Approved prescription drug products with therapeutic equivalence evaluations.

[Washington, D.C.?] : U.S. Dept. of Health and Human Services, Public Health Service, Food and Drug Administration, Bureau of Drugs : 1980-

<http://hdl.handle.net/2027/mdp.39015072931606>

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HE 20,4270 : 984 / supp. 6

Pharm  
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Aug 1984 -  
Feb 1985  
Cum. Suppl. 6

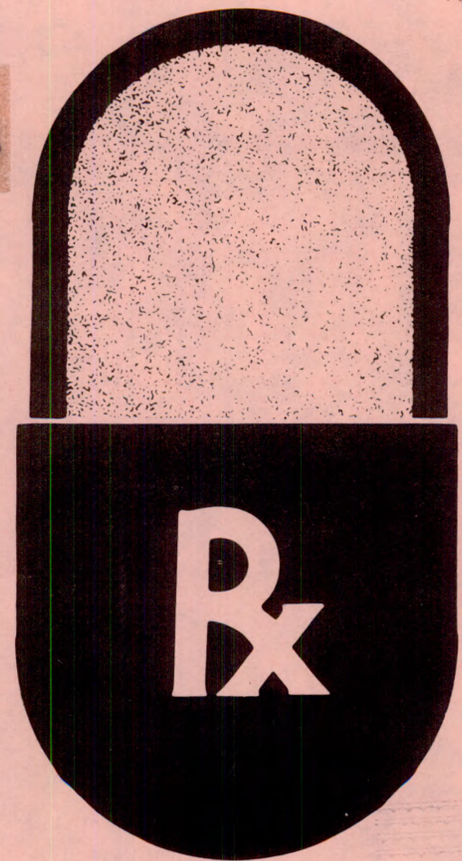
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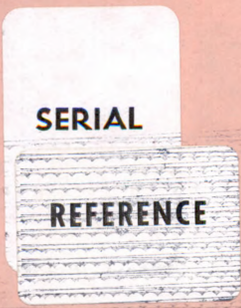
**CUMULATIVE  
SUPPLEMENT 6  
AUG'84 - FEB'85**



# APPROVED PRESCRIPTION DRUG PRODUCTS

WITH  
THERAPEUTIC EQUIVALENCE EVALUATIONS

**5<sup>TH</sup> EDITION**



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FOOD AND DRUG ADMINISTRATION  
APPROVED PRESCRIPTION DRUG PRODUCTS  
WITH  
THERAPEUTIC EQUIVALENCE EVALUATIONS  
CUMULATIVE SUPPLEMENT

I. PREFACE

This cumulative supplement is one of a series of monthly updates to the Approved Prescription Drug Products with Therapeutic Equivalence Evaluations, 5th Edition (the List), to cover interim revisions to the annual publication of the List in its entirety. The List is comprised of several parts and some by their nature, are identified by the term "list." The cumulative supplements routinely provide updates to two of these lists: The Drug Product List and the DESI Addendum.

The List cannot be used effectively without the current cumulative supplement. Users may wish to place an asterisk (\*) in the List to the left of the ingredient(s) in the Drug Product List and the product name in the Addendum to indicate that changes to that entry appear in the cumulative supplement. It is also suggested that earlier cumulative supplements be discarded to avoid possible confusion. In this way, only the List and current cumulative supplement need be referenced.

A. DRUG PRODUCT LIST

The Drug Product List cumulative supplements include the changes made since August 1, 1984. Each subsequent cumulative supplement replaces the previous month's cumulative supplement.

Information in this cumulative supplement follows the format of the Drug Product List. 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.)

Context information on drug products is provided in each cumulative supplement for completeness to assist in locating the proper place in the Drug Product List for the revision. (Strength(s) which already exist in the publication will not be repeated for context.) A page number in parentheses referring to the Drug Product List is located to the right of the ingredient(s).

Additions to the Drug Product List 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.

Deletions from the Drug Product List 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 >DLT> 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.

B. ADDENDUM: DESI Pending List

Information in this cumulative supplement follows the format of the Addendum. Additions and deletions are indicated in the same manner as in the cumulative supplement to the Drug Product List. A change in Current Status of a DESI product is also indicated by an addition and a deletion.

II. SPECIAL NOTES

A. REPORT OF COUNTS FOR THE DRUG PRODUCT LIST

Categories of counts derived from product information in the Drug Product List and from this cumulative supplement are presented. The report includes counts of new molecular entities approved by the agency during the current month.

B. 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 Drug Product List.

| <u>Products</u>           | <u>Federal Register Reference</u> |
|---------------------------|-----------------------------------|
| dicyclomine hydrochloride | JUN 22, 1984 (49 FR 25681)        |
| isosorbide dinitrate      | AUG 3, 1984 (49 FR 31151)         |
| nandrolone decanoate      | JUL 15, 1983 (48 FR 32395)        |

(continued)

Products

Federal Register Reference

(continued)

|   |                            |
|---|----------------------------|
| neomycin sulfate with either:<br>dexamethasone sodium phosphate,<br>fluocinolone acetonide,<br>flurandrenolide,<br>hydrocortisone, or<br>methylprednisolone acetate.<br>[topical anti-infectives for<br>dermatologic use] | MAR 26, 1984 (49 FR 11888) |
| neomycin sulfate, polymyxin B sulfate,<br>bacitracin zinc, and hydrocortisone<br>[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)  |
| parenteral multivitamin products  | SEP 17, 1984 (49 FR 36446) |
| phenazopyridine hydrochloride and<br>sulfamethoxazole   | JUL 29, 1983 (48 FR 34516) |
| sulfanilamide and aminacrine  | AUG 22, 1983 (48 FR 38097) |
| tranlycypromine sulfate   | MAR 22, 1984 (49 FR 10708) |

C. 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> |
|--------------------------------|-----------------------------|-----------------------------|
| OHIO MEDICAL ANESTHETICS       | ANAQUEST                    | ANAQUEST                    |

D. ADDENDUM: DRUG PRICE COMPETITION AND PATENT TERM RESTORATION

The addendum of this supplement provides information required of the Agency by the "Drug Price Competition and Patent Term Restoration Act of 1984."

### III. REPORT OF COUNTS FOR THE DRUG PRODUCT LIST

#### DESCRIPTION OF REPORT

The following report provides summary counts derived from product information in the Drug Product List and the current cumulative supplement. The counts appear in two sections. Section A. refers to the products in the List and Section B. to products in the current cumulative supplement. A new column of data will appear in Section A. each three-month period following July '84. Section A. therefore will provide baseline and quarterly data while Section B. provides monthly activity.

#### USE OF REPORT

From the data presented under Section B., users should be able to observe such things as (1) newly approved, DESI effective 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, changes from prescription to over-the-counter status and discontinued marketing of products; 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.

#### Drug Product Definition

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

#### New Molecular Entity

The active moiety 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.

#### Drug Product Count

This report provides counts in several categories from the list composed of domestically marketed drug products approved for both safety and effectiveness under sections 505 and 507 of the Federal Food, Drug, and Cosmetic Act. Counts of products still pending in the DESI review are not provided. Excluded also are those approved drug products marketed by distributors; those marketed solely abroad; and products now regarded as medical devices, biologics or foods.

REPORT OF COUNTS FOR THE DRUG PRODUCT LIST

A. COUNTS CUMULATIVE BY QUARTERS

| <u>CATEGORIES COUNTED</u>       | <u>JULY '84 (BASELINE)</u> | <u>OCT '84</u> | <u>JAN '85</u> |
|---------------------------------|----------------------------|----------------|----------------|
| DRUG PRODUCTS LISTED            | 7415                       | 7609           | 7746           |
| SINGLE SOURCE                   | 2005 (27.0%)               | 2045 (26.9%)   | 2077 (26.8%)   |
| MULTISOURCE (1)                 | 5410 (72.9%)               | 5564 (73.1%)   | 5669 (73.2%)   |
| THERAPEUTICALLY EQUIVALENT      | 4393 (59.2%)               | 4497 (59.1%)   | 4598 (59.4%)   |
| NOT THERAPEUTICALLY EQUIVALENT  | 999 (13.4%)                | 1032 (13.5%)   | 1038 (13.4%)   |
| EXCEPTIONS (2)                  | 18 (0.3%)                  | 26 (0.3%)      | 23 (0.3%)      |
| NEW MOLECULAR ENTITIES APPROVED | -                          | 4              | 9              |
| NUMBER OF APPLICANTS            | 295                        | 300            | 304            |

B. ACTIVITY FOR SUPPLEMENT NUMBER 6

|                                    | <u>FEB '85</u> | <u>CUMULATIVE</u> |
|------------------------------------|----------------|-------------------|
| DRUG PRODUCTS ADDED:               | 44             | 44                |
| NEWLY APPROVED                     | 43             | 43                |
| DESI EFFECTIVE                     | 1              | 1                 |
| REMARKETED                         | 0              | 0                 |
| DRUG PRODUCTS REMOVED:             | 1              | 1                 |
| WITHDRAWN APPROVAL                 | 0              | 0                 |
| RX TO OTC SWITCH                   | 1              | 1                 |
| DISCONTINUED MARKETING             | 0              | 0                 |
| NET GAIN IN DRUG PRODUCTS          | 43             | 43                |
| SINGLE SOURCE PRODUCTS APPROVED    | 6              | 6                 |
| MULTISOURCE DRUG PRODUCTS APPROVED | 38             | 38                |
| NEW MOLECULAR ENTITIES APPROVED:   | 0              | 0                 |
| AS THE ENTITY                      | 0              | 0                 |
| AS A SALT, ESTER OR DERIVATIVE     | 0              | 0                 |
| OF THE ENTITY                      | 0              | 0                 |

- (1) THERAPEUTIC EQUIVALENCE EVALUATIONS PROVIDED ONLY FOR MULTISOURCE PRODUCTS (I.e., AVAILABLE FROM MORE THAN ONE APPLICANT)  
 (2) AMINO ACID-CONTAINING PRODUCTS OF VARYING COMPOSITION (SEE PAGE 1-5 OF THE LIST)



APPROVED PRESCRIPTION DRUG PRODUCTS  
 DRUG PRODUCT LIST

CUMULATIVE SUPPLEMENT NUMBER 6 / AUGUST '84 - FEBRUARY '85

|  |                   |         |         |
|--|-------------------|---------|---------|
| <u>ACETAMINOPHEN; OXYCODONE HYDROCHLORIDE (PAGE 3-2)</u> |                   |         |         |
| CAPSULE; ORAL  |                   |         |         |
| SECTRAL  |                   |         |         |
| IVES LABS/ANHO   | EQ 200MG BASEX    | N 18917 |         |
|  | EQ 400MG BASEX    | N 18917 | N 87463 |
|  |                   |         |         |
| <u>ACETAMINOPHEN; BUTALBITAL (PAGE 3-1)</u>              |                   |         |         |
| CAPSULE; ORAL  |                   |         |         |
| ESSIC  |                   |         |         |
| GILBERT LABORATORIES                                     | 325MG;50MG;40MGX  |         |         |
|  |                   |         |         |
| <u>ACETAMINOPHEN; BUTALBITAL; CAFFEINE (PAGE 3-1)</u>    |                   |         |         |
| CAPSULE; ORAL  |                   |         |         |
| ESSIC  |                   |         |         |
| GILBERT LABORATORIES                                     | 325MG;50MG;40MGX  | N 88825 | N 18828 |
|  |                   |         |         |
| <u>ACETAMINOPHEN; CODEINE PHOSPHATE (PAGE 3-1)</u>       |                   |         |         |
| TABLET; ORAL   |                   |         |         |
| ESSIC  |                   |         |         |
| GILBERT LABORATORIES                                     | 325MG;50MG;40MGX  | N 87629 |         |
| AB   |                   |         |         |
| FICRIDET   |                   |         |         |
| SANDOZ PHARMS/SANDOZ                                     | 325MG;50MG;40MGX  | N 88616 |         |
| REFAN  |                   |         |         |
| DM GRAHAM LABS   | 325MG;50MG;40MGX  | N 87804 |         |
|  |                   |         |         |
| <u>ACETAMINOPHEN; CODEINE PHOSPHATE (PAGE 3-1)</u>       |                   |         |         |
| TABLET; ORAL   |                   |         |         |
| AM   |                   |         |         |
| ACETAMINOPHEN W/ CODEINE PHOSPHATE #4/                   |                   |         |         |
| AM   |                   |         |         |
| ZENITH LABORATORIES                                      | 300MG;60MG        | N 87083 | N 18241 |
| AA   |                   |         | N 18241 |
| ZENITH LABORATORIES                                      | 300MG;60MG        |         | N 18785 |
|  |                   |         | N 18785 |
|  |                   |         | N 18832 |
|  |                   |         | N 18877 |
|  |                   |         |         |
| <u>ACETAMINOPHEN; HYDROCODONE BITARTRATE (PAGE 3-2)</u>  |                   |         |         |
| TABLET; ORAL   |                   |         |         |
| HYDROCODONE BITARTRATE W/ ACETAMINOPHEN                  |                   |         |         |
| AA   |                   |         |         |
| BARR LABORATORIES  | 500MG;5MGX        |         |         |
|  |                   |         |         |
| <u>ACETAMINOPHEN; OXYCODONE HYDROCHLORIDE (PAGE 3-2)</u> |                   |         |         |
| CAPSULE; ORAL  |                   |         |         |
| TYLOX  |                   |         |         |
| MCNEIL PHARM   | 500MG;5MGX        | N 88790 |         |
| TYLOX-325  |                   |         |         |
| MCNEIL PHARM   | 325MG;5MGX        | N 88246 |         |
|  |                   |         |         |
| <u>ACETIC ACID, GLACIAL (PAGE 3-3)</u>                   |                   |         |         |
| SOLUTION/DROPS; OTIC                                     |                   |         |         |
| ACETIC ACID  |                   |         |         |
| AT   |                   |         |         |
| THAMES PHARMACAL   | 22M               | N 87550 | N 88638 |
|  |                   |         |         |
| <u>ACYCLOVIR (PAGE 3-4)</u>                              |                   |         |         |
| CAPSULE; ORAL  |                   |         |         |
| ZOVIRAX  |                   |         |         |
| BURROUGHS WELLCOME                                       | 200MGX            |         |         |
|  |                   |         |         |
| <u>ALLOPURINOL (PAGE 3-5)</u>                            |                   |         |         |
| TABLET; ORAL   |                   |         |         |
| ALLOPURINOL  |                   |         |         |
| AB   |                   |         |         |
| BOLAR PHARMACEUTICAL                                     | 100MGX            |         |         |
| AB   |                   |         |         |
| BOLAR PHARMACEUTICAL                                     | 300MGX            |         |         |
| AB   |                   |         |         |
| CHELSEA LABORATORIES                                     | 100MGX            |         |         |
| AB   |                   |         |         |
| DANBURY PHARMACAL  | 100MGX            |         |         |
| AB   |                   |         |         |
| DANBURY PHARMACAL  | 300MGX            |         |         |
| AB   |                   |         |         |
|  |                   |         |         |
| <u>AMIDINOCILLIN (PAGE 3-6)</u>                          |                   |         |         |
| INJECTABLE; INJECTION                                    |                   |         |         |
| COACTIN  |                   |         |         |
| HOFFMANN-LA ROCHE  | 250MG/VIALX       |         |         |
|  | 500MG/VIALX       |         |         |
|  | 1GM/VIALX         |         |         |
|  |                   |         |         |
| <u>AMIKACIN SULFATE (PAGE 3-6)</u>                       |                   |         |         |
| INJECTABLE; INJECTION                                    |                   |         |         |
| AMIKIN   |                   |         |         |
| BRISTOL LABS/B-M   | EQ 50MG BASE/MLX  |         |         |
|  | EQ 250MG BASE/MLX |         |         |

AMINO ACIDS (PAGE 3-6)

INJECTABLE; INJECTION  
 BRANCHAMIN 4%  
 TRAVENOL LABS 4%  
 BRANCHAMIN 4% IN PLASTIC CONTAINER  
 TRAVENOL LABS 4%  
 TRAVASOL 10% W/O ELECTROLYTES IN PLASTIC CONTAINER  
 TRAVENOL LABS 10%  
 TRAVASOL 5.5% W/O ELECTROLYTES IN PLASTIC CONTAINER  
 TRAVENOL LABS 5.5%  
 TRAVASOL 8.5% W/O ELECTROLYTES IN PLASTIC CONTAINER  
 TRAVENOL LABS 8.5%

AMINO ACIDS; DEXTROSE (PAGE 3-7)

INJECTABLE; INJECTION  
 AMINOSYN 3.5% W/ DEXTROSE 5% IN PLASTIC CONTAINER  
 ABBOTT LABORATORIES 3.5%;56M/100ML  
 AMINOSYN 3.5% W/ DEXTROSE 25% IN PLASTIC CONTAINER  
 ABBOTT LABORATORIES 3.5%;256M/100ML  
 AMINOSYN 4.25% W/ DEXTROSE 25% IN PLASTIC CONTAINER  
 ABBOTT LABORATORIES 4.25%;256M/100ML

AMINOPHYLLINE (PAGE 3-8)

TABLET; ORAL  
 AMINOPHYLLINE  
 > DLT > /BP/ /CORD LABORATORIES/ /200MG/  
 > ADD > AB /CORD LABORATORIES 200MG

AMINOPHYLLINE; SODIUM CHLORIDE (PAGE 3-9)

INJECTABLE; INJECTION  
 AMINOPHYLLINE IN SODIUM CHLORIDE 0.45%  
 ABBOTT LABORATORIES 100MG/100ML;450MG/100ML  
 200ML/100ML;450MG/100ML  
 AMINOPHYLLINE IN SODIUM CHLORIDE 0.45% IN PLASTIC CONTAINER  
 ABBOTT LABORATORIES 100MG/100ML;450MG/100ML  
 200MG/100ML;450MG/100ML  
 400MG/100ML;450MG/100ML  
 500MG/100ML;450MG/100ML  
 /AMINOPHYLLINE 0.45% IN SODIUM CHLORIDE 0.45%  
 /ABBOTT LABORATORIES//50MG/100ML;450MG/100ML/  
 /AMINOPHYLLINE 0.45% IN SODIUM CHLORIDE 0.45%  
 /ABBOTT LABORATORIES//100MG/100ML;450MG/100ML/  
 /AMINOPHYLLINE 0.45% IN SODIUM CHLORIDE 0.45%  
 /ABBOTT LABORATORIES//200MG/100ML;450MG/100ML/

AMITRIPTYLINE HYDROCHLORIDE (PAGE 3-10)

TABLET; ORAL  
 AMITRIPTYLINE HCL  
 AMERICAN THERAPEUTIC 25MG  
 50MG  
 75MG  
 100MG  
 PAR PHARMACEUTICAL  
 10MG  
 25MG  
 50MG  
 75MG  
 100MG  
 150MG  
 SIDMAK LABORATORIES  
 10MG  
 25MG  
 50MG  
 75MG  
 100MG  
 150MG  
 SUPERPHARM  
 10MG  
 25MG  
 50MG  
 75MG  
 100MG

AMOXICILLIN; POTASSIUM CLAVULANATE (PAGE 3-13)

POWDER FOR RECONSTITUTION; ORAL  
 AUGMENTIN '125'  
 BEECHAM LABS/BEECHAM 125MG/5ML;  
 EQ 31.25MG ACID/5ML  
 N 50575  
 AUGMENTIN '250'  
 BEECHAM LABS/BEECHAM 250MG/5ML;EQ 62.5MG ACID/5ML  
 N 50575

TABLET; ORAL  
 AUGMENTIN '250'  
 BEECHAM LABS/BEECHAM 250MG;EQ 125MG ACID  
 N 50564  
 AUGMENTIN '500'  
 BEECHAM LABS/BEECHAM 500MG;EQ 125MG ACID  
 N 50564

AMPHETAMINE SULFATE (PAGE 3-13)

TABLET; ORAL  
 AMPHETAMINE SULFATE  
 LANNETT  
 5MG  
 10MG  
 N 83901  
 N 83901

ASPIRIN; BUTALBITAL; CAFFEINE (PAGE 3-16)

TABLET; ORAL  
 BUTALBITAL COMPOUND  
 ZENITH LABORATORIES 325MG;50MG;40MG  
 AB N 85441

ASPIRIN; CAFFEINE; PROPOXYPHENE HYDROCHLORIDE (PAGE 3-16)

CAPSULE; ORAL  
PROPOXYPHENE COMPOUND 65  
 AA ZENITH LABORATORIES 389MG; 32.4MG; 65MG  
 PROPOXYPHENE HCL W/ ASPIRIN AND CAFFEINE  
 AA CHELSEA LABORATORIES 389MG; 32.4MG; 65MG

N 83077  
 N 85732

BITOLTEROL MESYLATE (PAGE 3-24)

AEROSOL; INHALATION  
 TORNALATE  
 WINTHROP-BREON/STERL 0.37MG/INH

N 18770

ASPIRIN; METHOCARBANOL (PAGE 3-17)

TABLET; ORAL  
METHOCARBANOL W/ ASPIRIN/  
 METHOCARBANOL AND ASPIRIN

AA AMBAY BAY LABORATORIES 12.5MG/5ML; 10MG/5ML  
 AA AMRENYL MARION LABORATORIES 12.5MG/5ML; 10MG/5ML  
 AA BROVARYL NATL PHARM MFG/BARRE 12.5MG/5ML; 10MG/5ML

N 88626  
 N 09319  
 N 88343

BENZOYL PEROXIDE; ERYTHROMYCIN (PAGE 3-21)

GEL; TOPICAL  
 BENZAMYCIN  
 DERMIK/RORER-AMCHEM 5%; 3%  
BENZATHAZINE; RESERPINE (PAGE 3-21)

N 50557

TABLET; ORAL  
 EXNA-R/  
 AH ROBINS/  
 50MG; 0.125MG/

/N 14861/

BETAMETHASONE DIPROPIONATE (PAGE 3-22)

OINTMENT; TOPICAL  
 ALPHATREX  
 SAVAGE LABS/BYK-GLDN EQ 0.05% BASE  
BETAMETHASONE DIPROPIONATE

AB E FOUGERA/BYK-GLDN EQ 0.05% BASE  
 AB PHARMADERM/BYK-GLDN EQ 0.05% BASE  
 DIPROLENE EQ 0.05%  
 BX SCHERING EQ 0.05%  
 AB DIPROSONE EQ 0.05%  
 AB SCHERING EQ 0.05%

N 19143  
 N 19141  
 N 19140  
 N 18741  
 N 17691

BETAMETHASONE VALERATE (PAGE 3-22)

CREAM; TOPICAL  
 BETATREX  
 SAVAGE LABS/BYK-GLDN EQ 0.1% BASE/  
 AB SAVAGE LABS/BYK-GLDN EQ 0.1% BASE  
 VALNAG  
 AB NMC LABORATORIES EQ 0.1% BASE

/N 18862/  
 N 18862  
 N 70050

OINTMENT; TOPICAL  
 VALNAG  
 AB NMC LABORATORIES EQ 0.1% BASE

N 70051

BROMPHENIRAMINE MALEATE; CODEINE PHOSPHATE;  
 PHENYLPROPANOLAMINE HYDROCHLORIDE (PAGE 3-25)

SYRUP; ORAL

BIFETANE DC 2MG/5ML; 10MG/5ML;  
 BAY LABORATORIES 12.5MG/5ML  
 BROMANATE DC 2MG/5ML; 10MG/5ML;  
 NATL PHARM MFG/BARRE 12.5MG/5ML  
 DIMETANE-DC 2MG/5ML; 10MG/5ML  
 AH ROBINS 12.5MG/5ML

N 88904  
 N 88723  
 N 11694

BROMPHENIRAMINE MALEATE; PHENYLPROPANOLAMINE HYDROCHLORIDE  
 (PAGE 3-25)

ELIXIR; ORAL

BIFHETAP 4MG/5ML; 25MG/5ML  
 BAY LABORATORIES 4MG/5ML; 25MG/5ML  
 BROMANATE 4MG/5ML; 25MG/5ML  
 NATL PHARM MFG/BARRE 4MG/5ML; 25MG/5ML  
 ELIXIR, DIMETAPP/  
 AH ROBINS/  
 4MG/5ML; 25MG/5ML

N 88687  
 N 88688  
 /N 13087/

TABLET; CONTROLLED RELEASE; ORAL/  
 DIMETAPP 12MG; 25MG/  
 AH ROBINS/  
 12MG; 25MG/  
 /N 14436/  
 /N 14436/

BUPRENORPHINE HYDROCHLORIDE (PAGE 3-26)

INJECTABLE; INJECTION  
 BUPRENEX/  
 NORRICH, EASTON/P&S/  
 EQ 0.5MG; BASE/ML

/N 14401/

DRUG PRODUCT LIST / CUMULATIVE SUPPLEMENT NUMBER 6 / AUGUST '84 - FEBRUARY '85

BUTABARBITAL SODIUM (PAGE 3-26)

ELIXIR; ORAL  
 /SODIUM BUTABARBITAL/  
 BUTABARBITAL SODIUM

TABLET; ORAL  
 CAPOTEN  
 ER SQUIBB AND SONS 12.5MGX

N18343

CALCIOTININ (PAGE 3-27)

INJECTABLE; INJECTION  
 CALCINAR  
 /ARMOUR PHARM/  
 /ARMOUR PHARM/  
 ARMOUR PHARM

/300 MCG UNITS/ML/  
 /400 MCG UNITS/VIAL/  
 200 IU/ML  
 400 IU/VIAL

/N:17769/  
 /N:17497/  
 N 17769  
 N 17497

TABLET; ORAL  
 CAPOZIDE 25/15  
 ER SQUIBB AND SONS 25MG;15MGX  
 CAPOZIDE 25/25  
 ER SQUIBB AND SONS 25MG;25MGX  
 CAPOZIDE 50/15  
 ER SQUIBB AND SONS 50MG;15MGX  
 CAPOZIDE 50/25  
 ER SQUIBB AND SONS 50MG;25MGX

N 18709  
 N 18709  
 N 18709  
 N 18709

CAPTOPRIL; HYDROCHLOROTHIAZIDE (PAGE 3-31)

CALCIUM CHLORIDE; DEXTROSE; MAGNESIUM CHLORIDE; SODIUM CHLORIDE; SODIUM LACTATE (PAGE 3-28)

SOLUTION; INTRAPERITONEAL

DELPLEX W/ DEXTROSE 1.5% IN PLASTIC CONTAINER

AI DELMED 25.7MG/100ML;1.5GM/100ML;  
 15.2MG/100ML;56.7MG/100ML;  
 392MG/100MLX N 18883

DELPLEX W/ DEXTROSE 2.5% IN PLASTIC CONTAINER

AI DELMED 25.7MG/100ML;2.5GM/100ML;  
 15.2MG/100ML;56.7MG/100ML;  
 392MG/100MLX N 18883

DELPLEX W/ DEXTROSE 4.25% IN PLASTIC CONTAINER

AI DELMED 25.7MG/100ML;4.25GM/100ML;  
 15.2MG/100ML;56.7MG/100ML;  
 392MG/100MLX N 18883

DELPLEX W/ DEXTROSE 1.5% LOW MAGNESIUM IN PLASTIC CONTAINER

AI DELMED 25.7MG/100ML;1.5GM/100ML;  
 5.08MG/100ML;53.8MG/100ML;  
 448MG/100MLX N 18883

DELPLEX W/ DEXTROSE 2.5% LOW MAGNESIUM IN PLASTIC CONTAINER

AI DELMED 25.7MG/100ML;2.5GM/100ML;  
 5.08MG/100ML;53.8MG/100ML;  
 448MG/100MLX N 18883

DELPLEX W/ DEXTROSE 4.25% LOW MAGNESIUM IN PLASTIC CONTAINER

AI DELMED 25.7MG/100ML;4.25GM/100ML;  
 5.08MG/100ML;53.8MG/100ML;  
 448MG/100MLX N 18883

CALCIUM GLUCEPTATE (PAGE 3-30)

INJECTABLE; INJECTION  
 CALCIUM GLUCEPTATE  
 /AP/ /ANTI-PEDICATION SYS//EQ 30MG CALCIUM/5ML/

/N:87455/

INJECTABLE; INJECTION  
 MEFOXIN IN SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER  
 EQ 20MG BASE/ML;9MG/MLX N 50581  
 MSD/MERCK EQ 40MG BASE/ML;9MG/MLX N 50581

CARBACHOL (PAGE 3-31)

/SOLUTION/DROPS; OPHTHALMIC/  
 INJECTABLE; INJECTION

CEFORANIDE (PAGE 3-33)

INJECTABLE; INJECTION  
 PRECEF

BRISTOL LABS/B-M 500MG/VIALM  
 1GM/VIALM  
 2GM/VIALM  
 10GM/VIALM  
 20GM/VIALM

N 62579  
 N 62579  
 N 62579  
 N 62579

CEFOXITIN SODIUM; DEXTROSE (PAGE 3-33)

INJECTABLE; INJECTION  
 MEFOXIN IN DEXTROSE 5% IN PLASTIC CONTAINER  
 EQ 20MG BASE/ML;50MG/MLX N 50581  
 MSD/MERCK EQ 40MG BASE/ML;50MG/MLX N 50581

CEFOXITIN SODIUM; SODIUM CHLORIDE (PAGE 3-33)

INJECTABLE; INJECTION  
 MEFOXIN IN SODIUM CHLORIDE 0.9% IN PLASTIC CONTAINER  
 EQ 20MG BASE/ML;9MG/MLX N 50581  
 MSD/MERCK EQ 40MG BASE/ML;9MG/MLX N 50581

CEFTIZOXIME SODIUM; DEXTROSE (PAGE 3-33)

INJECTABLE; INJECTION  
 CEFIZOX IN DEXTROSE 5% IN PLASTIC CONTAINER  
 SK&F LABORATORIES EQ 20MG BASE/ML;50MG/ML N 50589  
 EQ 40MG BASE/ML;50MG/ML N 50589

INJECTABLE; INJECTION  
 CHYMOTRYPSIN  
 SMITH LABORATORIES 4,000 UNITS/VIAL N 18663

CEFTRIAXONE SODIUM (PAGE 3-33)

INJECTABLE; INJECTION  
 ROCEPHIN  
 HOFFMANN-LA ROCHE EQ 250MG BASE/VIAL N 50585  
 EQ 500MG BASE/VIAL N 50585  
 EQ 1GM BASE/VIAL N 50585  
 EQ 2GM BASE/VIAL N 50585  
 EQ 10GM BASE/VIAL N 50585

CISPLATIN (PAGE 3-44)

INJECTABLE; INJECTION  
 PLATINOL/  
 /PRISTOL LABS/B-N /10MG/ML/  
 /50MG/VIAL/ N 18057  
 PLATINOL-AQ BRISTOL LABS/B-N 0.5MG/ML N 18057

CELLULOSE SODIUM PHOSPHATE (PAGE 3-34)

POWDER; ORAL  
 CALCIBIND  
 MISSION PHARMACAL 300GM/BOTM N 18757

CLOMIPHENE CITRATE (PAGE 3-45)

TABLET; ORAL  
 CLOMID  
 > DLT > /BP/ /MERRILL, DON/DON CHEN/50MG/ N 16131  
 > ADD > AB MERRELL DON/DON CHEN 50MG N 16131  
 > DLT > /BP/ /CLOMIPHENE CITRATE /50MG/ N 18361  
 > ADD > AB /PLANTEK/IKAPHARM /50MG/ N 18361  
 PLANTEX/IKAPHARM 50MG

CHLORDIAZEPOXIDE HYDROCHLORIDE (PAGE 3-37)

CAPSULE; ORAL  
 CHLORDIAZEPOXIDE HCL  
 LEMMON 5MG N 88705  
 10MG N 88706  
 25MG N 88707

CLONIDINE (PAGE 3-45)

FILM, CONTROLLED RELEASE; PERCUTANEOUS  
 CATAPRES-TTS-1 BOEHRINGER INGELHEIM 2.5MG N 18891  
 CATAPRES-TTS-2 BOEHRINGER INGELHEIM 5MG N 18891  
 CATAPRES-TTS-3 BOEHRINGER INGELHEIM 7.5MG N 18891

CHLORPROPANIDE (PAGE 3-42)

TABLET; ORAL  
 CHLORPROPANIDE  
 BARR LABORATORIES 100MG N 88812  
 250MG N 88813  
 CHELSEA LABORATORIES 100MG N 86865  
 COLMED LABORATORIES 100MG N 88708  
 250MG N 88709  
 CORD LABORATORIES 100MG N 88725  
 250MG N 88726  
 DANBURY PHARMACAL 100MG N 88852  
 250MG N 89826  
 DURAMED PHARMS 100MG N 88918  
 250MG N 88919  
 LEMMON 100MG N 88768  
 SUPERPHARM 100MG N 88694  
 250MG N 88695  
 ZENITH LABORATORIES 100MG N 88840  
 GLUCANIDE 250MG N 88641  
 LEMMON

CODEINE PHOSPHATE; PHENYLEPHRINE HYDROCHLORIDE; PROMETHAZINE HYDROCHLORIDE (PAGE 3-46)

SYRUP; ORAL  
 PHEREGAN VC W/ CODEINE 10MG/5ML; 5MG/5ML; 6.25MG/5ML N 08306  
 AA WYETH LABS/AMHO  
 PROMETH VC W/ CODEINE NATL PHARM MFG/BARRE 10MG/5ML; 5MG/5ML; 6.25MG/5ML N 88764  
 AA PROMETHAZINE VC W/ CODEINE BAY LABORATORIES 10MG/5ML; 5MG/5ML; 6.25MG/5ML N 88896

CODEINE PHOSPHATE; PROMETHAZINE HYDROCHLORIDE (PAGE 3-46)

SYRUP; ORAL  
 PHEREGAN W/ CODEINE 10MG/5ML; 6.25MG/5ML N 08306  
 AA WYETH LABS/AMHO

CODEINE PHOSPHATE; PROMETHAZINE HYDROCHLORIDE (PAGE 3-46)

DESONIDE (PAGE 3-53)

SYRUP; ORAL  
PROMETH W/ CODEINE  
 AA MATL PHARM MFG/BARRE 10MG/5ML;6.25MG/5ML N 88763  
PRMETHAZINE W/ CODEINE  
 AA BAY LABORATORIES 10MG/5ML;6.25MG/5ML N 88875

CREAM; TOPICAL

DESONEN  
 AB OMEN LABS/DERM PRODS 0.05% N 19048  
TRIDESILON  
 AB MILES PHARMS/MILES 0.05% N 17010

CODEINE PHOSPHATE; PSEUDOEPHEDRINE HYDROCHLORIDE; TRIPROLIDINE HYDROCHLORIDE (PAGE 3-46)

DESOXIMETASONE (PAGE 3-53)

SYRUP; ORAL  
ACTIFED W/ CODEINE  
 AA BURROUGHS WELLCOME 10MG/5ML;30MG/5ML;1.25MG/5ML N 12575  
PSEUDOEPHEDRINE O  
 AA BAY LABORATORIES 10MG/5ML;30MG/5ML;1.25MG/5ML N 88833

OINTMENT; TOPICAL  
 TOPICORT  
 HOECHST-ROUSSEL 0.05% N 18594

DEXAMETHASONE; NEOMYCIN SULFATE; POLYMYXIN B SULFATE (PAGE 3-55)

CORTICOTROPIN (PAGE 3-47)

OINTMENT; OPHTHALMIC

> ADD >  
 > ADD > AI  
 > ADD >

DEXACIDIN  
 COOPERVISION PHARMS 0.1%;EQ 3.5MG BASE/GM;  
 10,000 UNITS/GM N 62566

AP CARTER-GLOGAU LABS 40 UNITS/VIAL N 88772

SUSPENSION/DROPS; OPHTHALMIC

AI  
 COOPERVISION PHARMS 0.1%;EQ 3.5MG BASE/ML;  
 10,000 UNITS/ML N 62544

CHROMOLYN SODIUM (PAGE 3-48)

DEXAMETHASONE SODIUM PHOSPHATE (PAGE 3-55)

SOLUTION/DROPS; OPHTHALMIC  
 OPTICROM 4% N 18155  
 FISOXS

SOLUTION/DROPS; OPHTHALMIC  
 DEXAMETHASONE SODIUM PHOSPHATE  
 AI CARTER-GLOGAU LABS EQ 0.1% PHOSPHATE N 88771

CYPROHEPTADINE HYDROCHLORIDE (PAGE 3-51)

DEXAMETHASONE SODIUM PHOSPHATE; NEOMYCIN SULFATE (PAGE 3-56)

TABLET; ORAL  
 CYPROHEPTADINE HCL  
 AM THERAPEUTICS 4MG N 88798

SOLUTION/DROPS; OPHTHALMIC  
 NEDECADRON  
 MS&D/MERCK EQ 0.1% PHOSPHATE;  
 EQ 3.5MG BASE/ML N 50322  
 NEOMYCIN SULFATE-DEXAMETHASONE SODIUM PHOSPHATE  
 EQ 0.1% PHOSPHATE;  
 EQ 3.5MG BASE/ML N 62539  
 PHARMAFAIR

DESERPIDINE; METHYLOTHIAZIDE (PAGE 3-52)

DEXBROMPHENIRAMINE MALEATE; PSEUDOEPHEDRINE SULFATE (PAGE 3-56)

TABLET; ORAL  
 ENDURONYL  
 BP ABBOTT LABORATORIES 0.25MG;5MG N 12775  
 ENDURONYL FORTE  
 BP ABBOTT LABORATORIES 0.5MG;5MG N 12775  
 METHYLOTHIAZIDE AND DESERPIDINE  
 BP BOLAR PHARMACEUTICAL 0.25MG;5MG N 88486  
 BP 0.5MG;5MG N 88452

> DLT > /DEXBROMPHENIRAMINE MALEATE; PSEUDOEPHEDRINE SULFATE/ (PAGE 3-56)  
 > DLT > /TABLET; ORAL/  
 > DLT > /BIPHENOL/  
 > DLT > /SCHEFFINS/

/N 12344/

/116:6045/

DEXTROMETHORPHAN HYDROBROMIDE; PROMETHAZINE HYDROCHLORIDE  
 (PAGE 3-57)

SYRUP; ORAL  
PHENERGAN W/ DEXTROMETHORPHAN  
 AA WYETH LABS./AMHO 15MG/5ML; 6.25MG/5ML N 11265  
PROMETH W/ DEXTROMETHORPHAN  
 AA NATL PHARM MFG/BARRE 15MG/5ML; 6.25MG/5ML N 88762  
PROMETHAZINE DM  
 AA BAY LABORATORIES 15MG/5ML; 6.25MG/5ML N 88864

DEXTROSE (PAGE 3-57)

INJECTABLE; INJECTION  
DEXTROSE 30% IN PLASTIC CONTAINER  
 AP ABBOTT LABORATORIES 30GM/100ML N 19345  
 AP TRAVENOL LABS 30GM/100ML N 17521  
DEXTROSE 38.5% IN PLASTIC CONTAINER  
 ABBOTT LABORATORIES 38.5GM/100ML N 18923  
DEXTROSE 60% IN PLASTIC CONTAINER  
 AP ABBOTT LABORATORIES 60GM/100ML N 19346

DEXTROSE; HEPARIN SODIUM (PAGE 3-58)

INJECTABLE; INJECTION  
HEPARIN SODIUM 10,000 UNITS IN DEXTROSE 5%  
 AP ABBOTT LABORATORIES 5GM/100ML; 10,000 UNITS/100ML N 18911  
HEPARIN SODIUM 1000 UNITS AND DEXTROSE 5% IN PLASTIC CONTAINER  
 AP AM MCGAW/AM HOSP 5GM/100ML; 200 UNITS/100ML N 19130  
 HEPARIN SODIUM 12,500 UNITS IN DEXTROSE 5%  
 ABBOTT LABORATORIES 5GM/100ML; 5,000 UNITS/100ML N 18911  
HEPARIN SODIUM 2000 UNITS AND DEXTROSE 5% IN PLASTIC CONTAINER  
 AP AM MCGAW/AM HOSP 5GM/100ML; 200 UNITS/100ML N 19130  
HEPARIN SODIUM 25,000 UNITS IN DEXTROSE 5%  
 AP ABBOTT LABORATORIES 5GM/100ML; 10,000 UNITS/100ML N 18911  
 HEPARIN SODIUM 5000 UNITS AND DEXTROSE 5% IN PLASTIC CONTAINER  
 AM MCGAW/AM HOSP 5GM/100ML; 1,000 UNITS/100ML N 19130

DEXTROSE; MAGNESIUM CHLORIDE; POTASSIUM CHLORIDE; POTASSIUM PHOSPHATE, DIBASIC; SODIUM ACETATE (PAGE 3-59)

INJECTABLE; INJECTION  
 ISOLYTE P W/ DEXTROSE 5% IN PLASTIC CONTAINER  
 AM MCGAW/AM HOSP 5GM/100ML; 31MG/100ML; 130MG/100ML; 26MG/100ML; 320MG/100ML N 19025

DEXTROSE; THEOPHYLLINE (PAGE 3-62)

INJECTABLE; INJECTION  
THEOPHYLLINE AND DEXTROSE 5% IN PLASTIC CONTAINER  
 AP TRAVENOL LABS 5GM/100ML; 40MG/100ML N 18649  
 AP 5GM/100ML; 80MG/100ML N 18649  
 AP 5GM/100ML; 160MG/100ML N 18649  
 AP 5GM/100ML; 200MG/100ML N 18649  
 AP 5GM/100ML; 400MG/100ML N 18649  
THEOPHYLLINE 0.06% AND DEXTROSE 5% IN PLASTIC CONTAINER  
 AP AM MCGAW/AM HOSP 5GM/100ML; 40MG/100ML N 19083  
THEOPHYLLINE 0.08% AND DEXTROSE 5% IN PLASTIC CONTAINER  
 AP AM MCGAW/AM HOSP 5GM/100ML; 80MG/100ML N 19083  
THEOPHYLLINE 0.16% AND DEXTROSE 5% IN PLASTIC CONTAINER  
 AP AM MCGAW/AM HOSP 5GM/100ML; 160MG/100ML N 19083  
THEOPHYLLINE 0.2% AND DEXTROSE 5% IN PLASTIC CONTAINER  
 AP AM MCGAW/AM HOSP 5GM/100ML; 200MG/100ML N 19212  
THEOPHYLLINE 0.4% AND DEXTROSE 5% IN PLASTIC CONTAINER  
 AP AM MCGAW/AM HOSP 5GM/100ML; 400MG/100ML N 19212  
THEOPHYLLINE IN DEXTROSE 5% IN PLASTIC CONTAINER  
 AP ABBOTT LABORATORIES 5GM/100ML; 40MG/100ML N 19211  
 AP 5GM/100ML; 80MG/100ML N 19211  
 AP 5GM/100ML; 160MG/100ML N 19211  
 AP 5GM/100ML; 200MG/100ML N 19211  
 AP 5GM/100ML; 400MG/100ML N 19211

DICYCLOLINE HYDROCHLORIDE (PAGE 3-64)

CAPSULE; ORAL  
 BENTYL  
 MERRELL DOM/DOM CHEM 10MG N 07409  
 INJECTABLE; INJECTION  
 BENTYL  
 MERRELL DOM/DOM CHEM 10MG/ML N 08370

SYRUP; ORAL  
 BENTYL  
 MERRELL DOM/DOM CHEM 10MG/5ML N 07961

TABLET; ORAL  
 BENTYL  
 MERRELL DOM/DOM CHEM 20MG N 07409

DIETHYLPROPION HYDROCHLORIDE (PAGE 3-65)

TABLET; ORAL  
 DIETHYLPROPION HCL  
 LEMMON 25MG N 88642

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DIHYDROERGOTAMINE MESYLATE; HEPARIN SODIUM; LIDOCAINE HYDROCHLORIDE (PAGE 3-66)

INJECTABLE; INJECTION

ENBOLEX  
 SANDOZ PHARMS/SANDOZ 0.5MG/0.5ML;2,500 UNITS/0.5ML; 5.33MG/0.5MLx N 18885  
 0.5MG/0.7ML;5,000 UNITS/0.7ML; 7.46MG/0.7MLx N 18885

DISOPYRAMIDE PHOSPHATE (PAGE 3-68)

CAPSULE; ORAL

DISOPYRAMIDE PHOSPHATE  
 BIOCRAFT LABS EQ 100MG BASEx N 70101  
 EQ 150MG BASEx N 70102

MORPAC

SEARLE PHARMS EQ 100MG BASE N 17447  
 EQ 150MG BASE N 17447

DISULFIRAM (PAGE 3-68)

TABLET; ORAL

DISULFIRAM  
 PAR PHARMACEUTICAL 250MGx N 88792  
 500MGx N 88793

DIVALPROEX SODIUM (PAGE 3-69)

TABLET, ENTERIC COATED; ORAL

DEPAKOTE  
 ABBOTT LABORATORIES EQ 125MG BASEx N 18723

DOXYCYCLINE HYCLATE (PAGE 3-70)

CAPSULE; ORAL

DOXY-LEMON  
 LETHION EQ 50MG BASEx N 62497  
DOXYCYCLINE HYCLATE  
 PAR PHARMACEUTICAL EQ 50MG BASEx N 62434  
 SUPERPHARM EQ 50MG BASEx N 62469  
 EQ 100MG BASEx N 62469  
 WEST-WARD EQ 50MG BASEx N 62396  
 ZENITH LABORATORIES EQ 50MG BASEx N 62500  
 EQ 100MG BASEx N 62500

TABLET; ORAL

DOXYCYCLINE HYCLATE  
 SUPERPHARM EQ 100MG BASEx N 62494  
 ZENITH LABORATORIES EQ 100MG BASEx N 62505

DOXYLAMINE SUCCINATE (PAGE 3-70)

TABLET; ORAL

DECAPRYN  
 AA MERRELL DOW/DOW CHEM 25MG N 06412  
DOXYLAMINE SUCCINATE  
 AA QUANTUM PHARMICS 25MGx N 88603

EPINEPHRINE BITARTRATE; LIDOCAINE HYDROCHLORIDE (PAGE 3-72)

INJECTABLE; INJECTION

LIGNOSPAN FORTE EQ 0.02MG BASE/ML;2% N 88389  
 DEPROCO  
 LIGNOSPAN STANDARD EQ 0.01MG BASE/ML;2% N 88390  
 DEPROCO

ERYTHROMYCIN (PAGE 3-73)

OINTMENT; TOPICAL

AKNE-MYCIN  
 HERMAL PHARM LABS 2% N 50584

SOLUTION; TOPICAL

SANSAC  
 AI CHEN LABS/DERM PRODS 2% N 62522  
 > ADD > SNAB; TOPICAL  
 > ADD > ERYCETTE  
 > ADD > ORTHO PHARMACEUTICAL 2% N 50594

ESTROGENS, CONJUGATED (PAGE 3-76)

TABLET; ORAL

CONJUGATED ESTROGENS  
 BS ZENITH LABORATORIES 0.3MGx N 88569

ETHINYL ESTRADIOL; ETHYNDIOL DIACETATE (PAGE 3-78)

TABLET; ORAL-21

/DEBULEN/  
 DEMULEN 1/50-21

TABLET; ORAL-28

/DEBULEN-28/  
 DEMULEN 1/50-28

ETHINYL ESTRADIOL; LEVONORGESTREL (PAGE 3-78)

TABLET; ORAL-21

TRIPHASIL-21  
 MYETH LABS/AMHO 0.03MG,0.04MG,0.03MG;  
 0.05MG,0.075MG,0.125MGx N 19192

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ETHINYL ESTRADIOL; LEVONORGESTREL (PAGE 3-78)

TABLET; ORAL-28  
 TRIPHASIL-28  
 WYETH LABS/AMHO 0.03MG, 0.04MG, 0.03MG;  
 0.05MG, 0.075MG, 0.125MG N 19190

FLUOROURACIL (PAGE 3-83)

INJECTABLE; INJECTION  
 FLUOROURACIL  
 SOLOPAK LABORATORIES 50MG/ML N 88766  
 AP 50MG/ML N 88767

ETHINYL ESTRADIOL; NORETHINDRONE ACETATE (PAGE 3-79)

TABLET; ORAL-21  
 /LOESTRIN 21 1.5/30/  
 LOESTRIN 21 1.5/30

FUROSEMIDE (PAGE 3-86)

TABLET; ORAL  
 FUROSEMIDE  
 CORD LABORATORIES 80MG N 18569  
 AB LEDERLE LABS/AM CYAN 80MG N 18415  
 AB PARKE-DAVIS/M-L 80MG N 18419  
 AB LASIX  
 AB HOECHST-ROUSSEL 80MG N 16273

FENTANYL CITRATE (PAGE 3-81)

INJECTABLE; INJECTION  
 FENTANYL CITRATE  
 ABBOTT LABORATORIES EQ 0.05MG BASE/ML N 19115

SENTANICIN SULFATE (PAGE 3-86)

ointment; TOPICAL  
 SENTANICIN SULFATE  
 E FOUGERA/BYK-GLDN EQ IMG BASE/GM N 62533  
 AT PHARMADERM/BYK-GLDN EQ IMG BASE/GM N 62534

FLUNISOLIDE (PAGE 3-82)

AEROSOL; INHALATION  
 BRONALIDE  
 SYNTAX LABS/SYNTAX 0.025MG/INH N 18340

SOLUTION/DROPS; OPHTHALMIC  
 GEMOPTIC

AT ALLERGAN PHARMS EQ 3MG BASE/ML N 62452

FLUOCINOLONE ACETONIDE (PAGE 3-82)

CREAM; TOPICAL  
 FLUOCINOLONE ACETONIDE  
 BAY LABORATORIES 0.01% N 88757  
 AT PHARMAFAIR 0.01% N 88499  
 AT 0.025% N 88506  
 FLUCINID  
 HERBERT LABS/ALLERGN 0.025% N 87156  
 AT /MARION LABORATORIES//0.012% /N. 80434/  
 AT /0.025% /N. 80434/  
 AT

SLUTETHIMIDE (PAGE 3-88)

TABLET; ORAL  
 SLUTETHIMIDE  
 /AA/ /ZENITH LABORATORIES//500MG/ /N. 83683/

ointment; TOPICAL

FLUOCINOLONE ACETONIDE  
 BAY LABORATORIES 0.025% N 88742  
 FLUCINID  
 HERBERT LABS/ALLERGN 0.025% N 87157  
 AT /MARION LABORATORIES//0.025% /N. 80434/  
 AT

SONADOTROPIN, CHORIONIC (PAGE 3-89)

INJECTABLE; INJECTION  
 CHORIONIC GONADOTROPIN  
 CARTER-GLOGAU LABS 2,000 UNITS/VIAL N 17016

HALCINONIDE (PAGE 3-90)

CREAM; TOPICAL  
 /HALCINON/ /HALOS-E

/AT/ /MARION LABORATORIES//0.012% /N. 80434/

FOLICLE STIMULATING HORMONE; LUTEINIZING HORMONE (PAGE 3-85)

LUTEINIZING HORMONE; MENOIOPINS (PAGE 3-118)



HYDROCORTISONE (PAGE 3-99)

POWDER; FOR RX COMPOUNDING

H-CORT  
 /AA/ /PARAFLEX LABORATORIES/100Z/  
 AA TORCH LABORATORIES 100Z

/N' 87834/  
 N 87834

/AB/ /SK&F LABORATORIES/  
 /AB/ /25MG/  
 /BA/ /50MG/

/N' 18083/  
 /N' 18083/  
 /N' 18083/  
 N 83827  
 N 83827  
 N 83827

HYDROCORTISONE ACETATE (PAGE 3-102)

/AEROSOL; TOPICAL/  
 /EPIFOAM/  
 /REED&CARNRICK PHARMS/1Z/

/N' 86457/  
 N 86457

INDOMETHACIN (PAGE 3-108)

HYDROCORTISONE ACETATE; PRAOXINE HYDROCHLORIDE (PAGE 3-103)

AEROSOL; TOPICAL  
 EPIFOAM  
 REED&CARNRICK PHARMS 1Z;1Z

AB 25MG  
 AB 50MG  
 AB 25MG  
 AB 50MG

N 18829  
 N 18829  
 N 18806  
 N 18806

HYDROFLUMETHIAZIDE (PAGE 3-104)

TABLET; ORAL  
 HYDROFLUMETHIAZIDE  
 CHELSEA LABORATORIES 50MG

N 88528

INDOMETHACIN SODIUM TRIHYDRATE (PAGE 3-108)

HYDROFLUMETHIAZIDE; RESERPINE (PAGE 3-104)

TABLET; ORAL  
 RESERPINE AND HYDROFLUMETHIAZIDE  
 BP ZENITH LABORATORIES 50MG;0.125MG

N 88932

INJECTABLE; INJECTION  
 INDOCIN I.V.  
 MS&D/MERCK EQ 1MG BASE/VIALX  
 N 18878

HYDROXYZINE HYDROCHLORIDE (PAGE 3-105)

TABLET; ORAL  
 HYDROXYZINE HCL  
 PUREPAC/KALIPHARMA 10MG  
 25MG  
 50MG  
 SUPERPHARM 10MG  
 25MG  
 50MG

N 88120  
 N 88121  
 N 88122  
 N 88794  
 N 88795  
 N 88796

ISOETHARINE MESYLATE (PAGE 3-110)  
 AEROSOL; INHALATION  
 BRONKONETER  
 /BREON LABS/STERLING/0.61Z/  
 BN BREON LABS/STERLING 0.34MG/INH  
 ISOETHARINE MESYLATE  
 BN NATL PHARM MFG/BARRE 0.34MG/INH

/N' 12339/  
 N 12339  
 N 87558

IBUPROFEN (PAGE 3-106)

TABLET; ORAL  
 RUFEN  
 BOOTS PHARMACEUTICAL 400MG  
 600MG

N 70083  
 N 70088

KANAMYCIN SULFATE (PAGE 3-112)

INJECTABLE; INJECTION  
 KANTREX  
 BRISTOL LABS/B-M  
 AP EQ 75MG BASE/2MLX  
 AP EQ 500MG BASE/2MLX  
 AP EQ 1GM BASE/3MLX

N 62564  
 N 62564  
 N 62564

DRUG PRODUCT LIST / CUMULATIVE SUPPLEMENT NUMBER 6 / AUGUST '84 - FEBRUARY '85

LABELALOL HYDROCHLORIDE (PAGE 3-113)

INJECTABLE; INJECTION  
 NORHODYNE

5MG/MLM N 18687

TABLET; ORAL

NORHODYNE  
 SCHERING

AB 200MG N 18686  
 AB 300MG N 18686  
 AB 500MG N 18686

IRANDATE  
 GLAXO

AB 200MG N 18716  
 AB 300MG N 18716  
 AB 500MG N 18716

LEVONORDEFRIN; MEPIVACAINE HYDROCHLORIDE (PAGE 3-114)

INJECTABLE; INJECTION  
 SCANDINAVIAN L

0.05MG/ML; 22M N 88388

DEPROCO

LIDOCAINE (PAGE 3-114)

AEROSOL; ORAL  
 XYLOCAINE

ASTRA PHARM PRODS 102M N 14394

LINDANE (PAGE 3-116)

LOTION; TOPICAL

LINDANE

AI BAY LABORATORIES 12M N 88190

SHAMPOO; TOPICAL

LINDANE

AI BAY LABORATORIES 12M N 88191

MAGNESIUM CHLORIDE; POTASSIUM CHLORIDE; SODIUM ACETATE; SODIUM CHLORIDE; SODIUM GLUCONATE (PAGE 3-119)

SOLUTION; IRRIGATION

PHYSIOLYTE IN PLASTIC CONTAINER

AP ABBOTT LABORATORIES 30MG/100ML; 37MG/100ML; 370MG/100ML; 530MG/100ML; 500MG/100ML N 19024

SYNOVALTE IN PLASTIC CONTAINER

AP TRAVENOL LABS 30MG/100ML; 37MG/100ML; 368MG/100ML; 526MG/100ML; 502MG/100ML N 19326

MEPERIDINE HYDROCHLORIDE (PAGE 3-122)

INJECTABLE; INJECTION  
 MEPERIDINE HCL

AP ABBOTT LABORATORIES 10MG/ML N 88432  
 AP INTL MEDICATION SYS 10MG/ML N 86332

SYRUP; ORAL

DEMEROL

AA WINTHROP LABS/STERL 50MG/5ML N 05010

MEPERIDINE HCL

AA ROXANE LABORATORIES 50MG/5ML N 88744

TABLET; ORAL

MEPERIDINE HCL

AA BARR LABORATORIES 100MG N 88640

MEPHENTERMINE SULFATE (PAGE 3-123)

INJECTABLE; INJECTION  
 NYAMINE SULFATE

/MYETH. LABS/AMHO/ 15MG/ML; 30MG/ML; 15MG/ML; 30MG/ML N 88248  
 N 88248  
 EQ 15MG BASE/ML  
 EQ 30MG BASE/ML

MYETH LABS/AMHO

MEPIVACAINE HYDROCHLORIDE (PAGE 3-123)

INJECTABLE; INJECTION  
 CARBSCAINE

AP BREON LABS/STERLING 22M N 12250

MEPIVACAINE HCL

AP CARTER-GLOGAU LABS 12M N 88769

POLCCAINE

AP 32M N 88770

SCANDINAVIAN PLAIN

AP 32M N 88653

DEPROCO

AP 32M N 88387

MEPROBAMATE (PAGE 3-123)

TABLET; ORAL

MEPROBAMATE

/AA/ 200MG/ N 86226  
 /AB/ 300MG/ N 86229  
 /MM MAST/

METHICILLIN SODIUM (PAGE 3-127)

INJECTABLE; INJECTION  
CELBIENTIN  
 /AP / BECHAM LABS/BEECHAM / EQ 500MG BASE/VIAL  
 /AP / BECHAM LABS/BEECHAM / EQ 3.66M BASE/VIAL  
 /AP / BECHAM LABS/BEECHAM / EQ 5.46M BASE/VIAL  
 /AP / BECHAM LABS/BEECHAM / EQ 1.86M BASE/VIAL  
 /AP / BECHAM LABS/BEECHAM / EQ 96M BASE/VIAL

N 51493/  
 N 51493/  
 N 51493/  
 N 51493/  
 N 51493/  
 N 51493/

MORPHINE SULFATE (PAGE 3-135)

INJECTABLE; INJECTION  
 DURAMORPH PF  
 ELKINS-SINN/AHROBINS 0.5MG/MLM  
 1MG/MLM

N 18565  
 N 18565

NAFCILLIN SODIUM (PAGE 3-135)

INJECTABLE; INJECTION  
NAFCIL  
 BRISTOL LABS/B-M EQ 10GM BASE/VIALM  
MALLPEN  
 BEECHAM LABS/BEECHAM EQ 10GM BASE/VIAL

N 62527  
 N 61999

NALBUPHINE HYDROCHLORIDE (PAGE 3-136)

INJECTABLE; INJECTION  
 NUBAIN  
 DUPONT PHARMS/DUPONT 20MG/MLM

N 18024

> ADD >

NALTREXONE HYDROCHLORIDE (PAGE 3-136)

TABLETS, ORAL  
TREXAN  
 DUPONT PHARMS/DUPONT 50MG

N 18932

NEOMYCIN SULFATE; POLYMYXIN B SULFATE (PAGE 3-137)

SOLUTION/DROPS; OPHTHALMIC  
STATROL  
 ALCON LABORATORIES EQ 3.5MG BASE/ML;  
 16,250 UNITS/MLM

N 62339

NOMIFENSINE MALEATE (PAGE 3-140)

CAPSULE; ORAL  
MERITAL  
 HOECHST-ROUSSEL 25MG  
 50MG

N 18224  
 N 18224

NOREPINEPHRINE BITARTRATE (PAGE 3-140)

INJECTABLE; INJECTION  
 LEVOPHED  
 /AP / BECHAM LABS/STERLING / EQ 1MG BASE/ML  
 /AP / WINTHROP-BREON/STERL / EQ 1MG BASE/ML

N 07513/  
 N 07513

METHOTREXATE SODIUM (PAGE 3-128)

INJECTABLE; INJECTION  
MEXATE  
 BRISTOL LABS/B-M EQ 250MG BASE/VIALM  
MEXATE-83  
 BRISTOL CARIB/B-M/PR EQ 25MG BASE/MLM

N 86358  
 N 88760

> ADD >  
 > ADD >

METHYLOTHIAZIDE (PAGE 3-129)

TABLET; ORAL  
METHYLOTHIAZIDE  
 CHELSEA LABORATORIES 2.5MG  
 5MG

N 88750  
 N 88724

METRONIDAZOLE (PAGE 3-133)

INJECTABLE; INJECTION  
METRONIDAZOLE  
LYPHONED 500MG/100MLM  
METRYL IV 500MG/100MLM  
LEMNON

N 70071  
 N 70042

TABLET; ORAL

METRONIDAZOLE  
 PAR PHARMACEUTICAL 250MG  
 500MG  
 SIDMAK LABORATORIES 250MG  
 500MG  
 SUPERPHARM 250MG  
 500MG  
METRYL  
 LEMNON 250MG  
METRYL 500  
 LEMNON 500MG

N 70040  
 N 70039  
 N 70027  
 N 70033  
 N 70008  
 N 70009  
 N 70035  
 N 70044

> ADD >

MICONAZOLE NITRATE (PAGE 3-134)

SUPPOSITORY; VAGINAL  
 MONISTAT 3  
 ORTHO PHARMACEUTICAL 200MG

N 18888

DRUG PRODUCT LIST / CUMULATIVE SUPPLEMENT NUMBER 6 / AUGUST '84 - FEBRUARY '85 14

|   |                           |  |         |
|---|---------------------------|--|---------|
| <u>NYSTAIN (PAGE 3-141)</u>   |                           | <u>PHENYLEPHRINE HYDROCHLORIDE; PROMETHAZINE HYDROCHLORIDE</u><br>(PAGE 3-153) |         |
| SUSPENSION; ORAL  |                           |  |         |
| <u>NYSTAIN</u>  |                           |  |         |
| <u>AA</u>   | BAY LABORATORIES          | 100,000 UNITS/MLM  | N 62512 |
| <u>AA</u>   | PHARMAFAIR                | 100,000 UNITS/MLM  | N 62541 |
| TABLET; ORAL  |                           |  |         |
| <u>NYSTAIN</u>  |                           |  |         |
| <u>AA</u>   | QUANTUM PHARMICS          | 500,000 UNITSM   | N 62525 |
| <u>OXTRIPHYLLINE (PAGE 3-143)</u>   |                           |  |         |
| ELIXIR; ORAL  |                           |  |         |
| <u>CHOLEFYL</u>   |                           |  |         |
| <u>AA</u>   | PARKE-DAVIS/M-L           | 100MG/5MLM   | N 09268 |
| <u>AA</u>   | BAY LABORATORIES          | 100MG/5ML  | N 88243 |
| <u>OXYPHENBUTAZONE (PAGE 3-143)</u>   |                           |  |         |
| TABLET; ORAL  |                           |  |         |
| <u>OXYPHENBUTAZONE</u>  |                           |  |         |
| <u>AB</u>   | BOLAR PHARMACEUTICAL      | 100MGM   | N 88399 |
| <u>AB</u>   | TANDEMIL                  |  | N 12542 |
|   | GEIGY/CIBA-GEIGY          | 100MG  |         |
| <u>PENTAMIDINE ISETHIONATE (PAGE 3-148)</u>   |                           |  |         |
| INJECTABLE; INJECTION   |                           |  |         |
| PENTAM 300  |                           |  |         |
|   | LYPHORED                  | 300MG/VIALM  | N 19264 |
| <u>PENTOXIFYLLINE (PAGE 3-149)</u>  |                           |  |         |
| TABLET, CONTROLLED RELEASE; ORAL  |                           |  |         |
| TRENAL  |                           |  |         |
|   | HOECHST-ROUSSEL           | 400MGM   | N 18631 |
| <u>PHENTERMINE HYDROCHLORIDE (PAGE 3-151)</u>   |                           |  |         |
| CAPSULE; ORAL   |                           |  |         |
| PHENTERMINE HCL   |                           |  |         |
| <u>AA</u>   | PHARM BASICS              | 30MGM  | N 88797 |
| <u>PHENYTOIN SODIUM, EXTENDED (PAGE 3-153)</u>  |                           |  |         |
| INJECTABLE; INJECTION   |                           |  |         |
| <u>PHENYTOIN SODIUM</u>   |                           |  |         |
|   | SOLOPAK LABORATORIES      | 50MG/MLM   | N 88519 |
|   |                           | 50MG/MLM   | N 88520 |
|   |                           | 50MG/MLM   | N 88521 |
| <u>PHENYTOIN SODIUM, EXTENDED (PAGE 3-153)</u>  |                           |  |         |
| CAPSULE; ORAL   |                           |  |         |
| <u>DILANTIN</u>   |                           |  |         |
|   | PARKE-DAVIS/M-L           | 100MG  | N 84349 |
|   | EXTENDED PHENYTOIN SODIUM |  | N 88711 |
|   | BOLAR PHARMACEUTICAL      | 100MGM   |         |
| <u>PILLOCARPINE HYDROCHLORIDE (PAGE 3-154)</u>  |                           |  |         |
| GEL; OPHTHALMIC   |                           |  |         |
| PILOPINE HS   |                           |  |         |
|   | ALCON LABORATORIES        | 4%   | N 18796 |
| <u>POLYETHYLENE GLYCOL 3350; POTASSIUM CHLORIDE; SODIUM BICARBONATE; SODIUM CHLORIDE; SODIUM SULFATE (PAGE 3-155)</u> |                           |  |         |
| POWDER FOR RECONSTITUTION; ORAL   |                           |  |         |
| COLYTE  |                           |  |         |
|   | EDLAW PREPARATIONS        | 120GM/PACKET;1.49GM/PACKET;  |         |
|   |                           | 3.36GM/PACKET;2.92GM/PACKET;   | N 18983 |
|   |                           | 11.36GM/PACKETM  |         |
|   |                           | 227.1GM/PACKET;2.82GM/PACKET;  |         |
|   |                           | 6.36GM/PACKET;5.53GM/PACKET;   | N 18983 |
|   |                           | 21.5GM/PACKET;M  |         |
|   |                           | 360GM/PACKET;4.47GM/PACKET;  |         |
|   |                           | 10.08GM/PACKET;8.76GM/PACKET;  | N 18983 |
|   |                           | 34.08GM/PACKETM  |         |

POTASSIUM CHLORIDE (PAGE 3-156)

INJECTABLE; INJECTION  
POTASSIUM CHLORIDE IN PLASTIC CONTAINER  
 AP INVENEX LABS/LIFE 2MEG/MLM  
 AP 2MEG/MLM

N 88901

N 88908

PREDNISOLONE (PAGE 3-159)

TABLET; ORAL  
 PREDNISOLONE 5MGH  
 SUPERPHARM

N 88892

PREDNISOLONE ACETATE; SULFACETAMIDE SODIUM (PAGE 3-160)

OINTMENT; OPHTHALMIC  
 PREDSULFAIR 0.5%;10%  
 PHARMAFAIR  
 VASOCIDIN  
 AT COOPERVISION PHARMS 0.5%;10%  
 AT

N 88032

N 88791

PREDNISONE (PAGE 3-161)

SOLUTION; ORAL  
 PREDNISONE 5MG/5MLM  
 ROXANE LABORATORIES  
 PREDNISONE INTENSOL  
 ROXANE LABORATORIES 5MG/MLM

N 88703

N 88810

PROCAINAMIDE HYDROCHLORIDE (PAGE 3-163)

TABLET; ORAL  
 PREDNISONE 5MGH  
 SUPERPHARM 10MGH  
 BX 20MGH  
 BX  
 BX  
PROCAINAMIDE HCL  
 BULAR PHARMACEUTICAL 250MGH  
 500MGH  
 750MGH  
 AB  
 AB  
 AB  
PROCAN SR  
 PARKE-DAVIS/N-L /500MG/  
 500HG  
 250HG  
 750HG  
 1GMH  
 AB  
 AB  
 AB

N 88533

N 88534

N 88535

N 86468

N 86068

N 86465

N 87510

N 88489

PROCHLORPERAZINE EDISYLATE (PAGE 3-164)

CONCENTRATE; ORAL  
PROCHLORPERAZINE EDISYLATE  
 AA BAY LABORATORIES EQ 10MG BASE/MLM  
 SYRUP; ORAL  
PROCHLORPERAZINE EDISYLATE  
 AA BAY LABORATORIES EQ 5MG BASE/5MLM

N 88598

N 88597

PROMETHAZINE HYDROCHLORIDE (PAGE 3-165)

SYRUP; ORAL  
PROMETHAZINE HYDROCHLORIDE  
 AA BAY LABORATORIES EQ 5MG BASE/5MLM

PROPOXYPHENE HYDROCHLORIDE (PAGE 3-167)

CAPSULE; ORAL  
PROPOXYPHENE HCL  
 AA LENNON 65MGH

N 88615

PROTAMINE SULFATE (PAGE 3-168)

INJECTABLE; INJECTION  
 PROTAMINE SULFATE 250MG/VIALM  
 UPJOHN

N 07413

PSEUDOEPHEDRINE HYDROCHLORIDE; TRIPROLDINE HYDROCHLORIDE (PAGE 3-169)

SYRUP; ORAL  
TRILITRON  
 AA NENTRON PHARMS 30MG/5ML; 1.25MG/5MLM  
 TRIPROLDINE HCL AND PSEUDOEPHEDRINE HCL  
 PHARMAFAIR 30MG/5ML; 1.25MG/5MLM

N 88474

N 88541

QUINIDINE SULFATE (PAGE 3-170)

TABLET; ORAL  
QUIN-SULFN  
 AB/ ROHELL LABORATORIES /200MG/

N 88755/







ADDENDUM  
DESI PENDING LIST - 'EXEMPT' (COURT ORDER) CATEGORY  
CUMULATIVE SUPPLEMENT NUMBER 6 / AUGUST '84 - FEBRUARY '85

ASCORBIC ACID; BIOTIN; CYANOCOBALAMIN; DEXRANTHENOL;  
ERGOCALCIFEROL; FOLIC ACID; NIACINAMIDE; PHYTIADIONE;  
PYRIDOXINE HYDROCHLORIDE; RIBOFLAVIN PHOSPHATE SODIUM;  
THIAMINE HYDROCHLORIDE; VITAMIN A; VITAMIN E (PAGE AD2)

/INJECTABLE; INJECTION/  
/M.V.C. PEDIATRIC/  
/USV. PHARMACEUTICAL/  
/60MG VIAL; 0.02MG VIAL; 0.001MG VIAL; /  
/5MG VIAL; 0.01MG VIAL; 0.14MG VIAL; /  
/1MG VIAL; 0.2MG VIAL; /  
/EQ. 1MG BASE VIAL; 4MG VIAL; /  
/EQ. 1.2MG BASE VIAL; 0.7MG VIAL; /  
/7MG VIAL / N 18928/

ASCORBIC ACID; BIOTIN; CYANOCOBALAMIN; DEXRANTHENOL;  
ERGOCALCIFEROL; FOLIC ACID; NIACINAMIDE; PYRIDOXINE;  
HYDROCHLORIDE; RIBOFLAVIN PHOSPHATE SODIUM; THIAMINE;  
HYDROCHLORIDE; VITAMIN A; VITAMIN E (PAGE AD2)

(SEE SPECIAL NOTE B.)  
/INJECTABLE; INJECTION/  
/M.V.C. /  
/USV. PHARMACEUTICAL/  
/100MG VIAL; 0.06MG VIAL; 0.005MG VIAL; /  
/15MG VIAL; 0.005MG VIAL; 0.4MG VIAL; /  
/40MG VIAL; 4MG VIAL; 3.6MG VIAL; /  
/3MG VIAL; 1MG VIAL; /  
/10. IU VIAL / N 18933/

ASCORBIC ACID; BIOTIN; CYANOCOBALAMIN; DEXRANTHENOL;  
ERGOCALCIFEROL; FOLIC ACID; NIACINAMIDE; PYRIDOXINE;  
HYDROCHLORIDE; RIBOFLAVIN PHOSPHATE SODIUM; THIAMINE; VITAMIN  
E (PAGE AD2)

(SEE SPECIAL NOTE B.)  
/INJECTABLE; INJECTION/  
/MVC PLUS/  
/ASCOT. HOSP. PHARMS/  
/10MG/ML; 0.006MG/ML; 0.5 UGH/ML; /  
/1.5MG/ML; 20. IU/ML; 0.04MG/ML; 4MG/ML; /  
/0.4MG/ML; 0.36MG/ML; 0.3MG/ML; /  
/330. IU/ML / N 18439/

ASCORBIC ACID; BIOTIN; CYANOCOBALAMIN; DEXRANTHENOL; FOLIC  
ACID; NIACINAMIDE; PYRIDOXINE HYDROCHLORIDE; RIBOFLAVIN;  
THIAMINE HYDROCHLORIDE; VITAMIN A; VITAMIN E;  
(PAGE AD2)

(SEE SPECIAL NOTE B.)  
/INJECTABLE; INJECTION/  
/M.V.C. 9+3/  
/LIPPHED/  
/20MG/ML; 0.012MG/ML; 0.001MG/ML; /  
/5MG/ML; 0.08MG/ML; 8MG/ML; 0.8MG/ML; /  
/0.7MG/ML; 0.6MG/ML; 0.660. IU/ML; /  
/40. IU/ML; 2. IU/ML / N 18440/

ASCORBIC ACID; BIOTIN; CYANOCOBALAMIN; FOLIC ACID;  
NIACINAMIDE; PANTOTHENIC ACID; PYRIDOXINE HYDROCHLORIDE; RIBOFLAVIN;  
THIAMINE HYDROCHLORIDE; VITAMIN A; VITAMIN D; VITAMIN E  
(PAGE AD2)

(SEE SPECIAL NOTE B.)  
/INJECTABLE; INJECTION/  
/MULTIVITAMIN ADDITIVE/  
/ABBOTT. LABORATORIES/  
/100MG/5ML; 0.06MG/5ML; 0.005MG/5ML; /  
/0.4MG/5ML; 80MG/5ML; 15MG/5ML; /  
/4.86MG/5ML; 9.5MG/5ML; 3.55MG/5ML; /  
/330. IU/5ML; 200. IU/5ML; /  
/10. IU/5ML / N 18223/

ASCORBIC ACID; BIOTIN; DEXRANTHENOL; NIACINAMIDE; PYRIDOXINE;  
HYDROCHLORIDE; RIBOFLAVIN; THIAMINE HYDROCHLORIDE (PAGE AD2)

(SEE SPECIAL NOTE B.)  
/INJECTABLE; INJECTION/  
/BEROCCA C/  
/HOFFMAN-LA. ROCHE/  
/50MG/ML; 0.1MG/ML; 1.0MG/ML; 4.0MG/ML; /  
/10MG/ML; 5MG/ML; 5MG/ML / N 06071/  
/BEROCCA C. 500/  
/HOFFMAN-LA. ROCHE/  
/125MG/ML; 1.0MG/ML; 1.0MG/ML; 4.0MG/ML; /  
/10MG/ML; 5MG/ML; 5MG/ML / N 06071/

ASCORBIC ACID; DEXRANTHENOL; NIACINAMIDE; PYRIDOXINE;  
HYDROCHLORIDE; RIBOFLAVIN; THIAMINE HYDROCHLORIDE; VITAMIN A;  
VITAMIN D; VITAMIN E (PAGE AD3)

(SEE SPECIAL NOTE B.)  
/INJECTABLE; INJECTION/  
/M.V.C. /  
/USV. PHARMACEUTICAL/  
/50MG/ML; 2.5MG/ML; 1.0MG/ML; 1.5MG/ML; /  
/10MG/ML; 5MG/ML; 1.000. IU/ML; 100. IU/ML; /  
/0.5MG/ML /  
/1.001MG/ML; 5MG/ML; 2.0MG/ML; 3MG/ML; /  
/2MG/ML; 1.01MG/ML; 0.00. IU/ML; /  
/200. IU/ML; 1MG/ML / N 08809/

DIPYRIDAMOLE (PAGE AD4)

TABLET; ORAL  
DIPYRIDAMOLE  
SIDMAK LABORATORIES  
25MG  
50MG  
75MG

> ADD >  
> ADD >  
> ADD >

N 88683  
N 88684  
N 88685

DESI PENDING LIST - 'EXEMPT' (COURT ORDER) CATEGORY  
CUMULATIVE SUPPLEMENT NUMBER 6 / AUGUST '84 - FEBRUARY '85

/ISOSORBIDE DINITRATE/ (PAGE AD5)  
(ALL PRODUCTS - SEE SPECIAL NOTE B.)

/TABLET; ORAL/  
/ISOSORBIDE DINITRATE/  
/BARR. LABORATORIES/ /10MG/

/N. 87566/

/ISOSORBIDE DINITRATE/ (PAGE AD5)  
(ALL PRODUCTS - SEE SPECIAL NOTE B.)

/TABLET; SUBLINGUAL/  
/ISOSORBIDE DINITRATE/  
/BARR. LABORATORIES/ /10MG/

/N. 87565/

/TABLET; CONTROLLED RELEASE; ORAL/  
/ISOSORON/  
/FOREST LABORATORIES/ /10MG/

/N. 88426/

NITROGLYCERIN (PAGE AD7)

/CAPSULE; CONTROLLED RELEASE; ORAL/  
(ALL PRODUCTS - SEE SPECIAL NOTE B.)

/TABLET; CONTROLLED RELEASE; ORAL/  
(ALL PRODUCTS - SEE SPECIAL NOTE B.)

DESI PENDING LIST - OTHER THAN 'EXEMPT' (COURT ORDER) CATEGORY  
CUMULATIVE SUPPLEMENT NUMBER 6 / AUGUST '84 - FEBRUARY '85

CURRENT STATUS - INEFFECTIVE

CURRENT STATUS - INEFFECTIVE

/BENTLY, W. PHARMACEUTICAL/ /MERRILL, DONALD, CHEN/  
/DICYCLONINE HYDROCHLORIDE/; PHENOBARBITAL/

TUSS-ORNADE SK&F LABORATORIES  
CARAMIPHEN EDISYLATE; CHLORPHENIRAMINE MALEATE;  
ISOPROPAMIDE IODIDE; PHENYLPROPANOLAMINE HYDROCHLORIDE

BEROCCA C HOFFMANN-LA ROCHE

ASCORBIC ACID; BIOTIN; DEXPANTHENOL; NIACINAMIDE; PYRIDOXINE  
HYDROCHLORIDE; RIBOFLAVIN; THIAMINE HYDROCHLORIDE

BEROCCA C 500 HOFFMANN-LA ROCHE

ASCORBIC ACID; BIOTIN; DEXPANTHENOL; NIACINAMIDE; PYRIDOXINE  
HYDROCHLORIDE; RIBOFLAVIN; THIAMINE HYDROCHLORIDE

DIMETAPP AH ROBINS

BROMPHENIRAMINE MALEATE; PHENYLEPHRINE HYDROCHLORIDE;  
PHENYLPROPANOLAMINE HYDROCHLORIDE

CURRENT STATUS - EFFECTIVENESS TO BE DETERMINED

M. V. I. PEDIATRIC USV PHARMACEUTICAL  
ASCORBIC ACID; BIOTIN; CYANOCOBALAMIN; DEXPANTHENOL;  
ERGOCALCIFEROL; FOLIC ACID; NIACINAMIDE; PHYTONADIONE;  
PYRIDOXINE HYDROCHLORIDE; RIBOFLAVIN PHOSPHATE SODIUM;  
THIAMINE HYDROCHLORIDE; VITAMIN A; VITAMIN E

> DLT > /ELECTROLYT/ /OWEN, LABS/BERN, PRODS/  
> DLT > /HYDROCORTISONE/

ELIXIR DIMETAPP AH ROBINS

BROMPHENIRAMINE MALEATE; PHENYLEPHRINE HYDROCHLORIDE;  
PHENYLPROPANOLAMINE HYDROCHLORIDE

> DLT > /HC (HYDROCORTISONE)/ /G. AND. N. PHARMACAL/  
> DLT > /HYDROCORTISONE/

> DLT > /HYDROCORTISONE/ /TOSNE, PAULSEN/  
> DLT > /HYDROCORTISONE/

> DLT > /ALCO, CHS/ /ELI LILLY/  
> DLT > /ERYTHRONICIN/

> DLT > /NEOSPORIN B/ /BURROUGHS, WELLCOME/  
> DLT > /GRANICIDIN; NEOMYCIN SULFATE; POLYMYXIN B; SULFATE/

/TERRA-CORTIL/ /PFIZER, LABS/PFIZER/  
/HYDROCORTISONE; OXYTETRACYCLINE, HCL/

> DLT > /NITRACORT/ /OWEN, LABS/BERN, PRODS/  
> DLT > /HYDROCORTISONE/

> DLT > /PRISCOLINE/ /CIBA, CIBA-GEIGY/  
> DLT > /TOLAZOLINE, HYDROCHLORIDE/



## ADDENDUM D: DRUG PRICE COMPETITION AND PATENT TERM RESTORATION

On September 24, 1984, the President signed into law the Drug Price Competition and Patent Term Restoration Act of 1984. The Act amends section 505 of the Federal Food, Drug and Cosmetic Act, authorizing the Agency to accept abbreviated new drug applications for most previously approved drug products. This new legislation also provides for extending the term of a patent which claims a product, use, or method of manufacture that was subject to a regulatory review period in accordance with the Act.

The statute requires that FDA make publicly available a list of approved drug products containing the following information:

- 1) an alphabetical list of all drugs by official and proprietary name approved for safety and effectiveness, with monthly updates;
- 2) the application number and approval date for each drug product approved from January 1, 1982; and
- 3) whether in vitro and/or in vivo bioequivalence studies are required for ANDA approval.

The Approved Prescription Drug Products with Therapeutic Equivalence Evaluations, 5th Edition, (APDP) and its monthly supplements will be used to satisfy this new requirement.

In addition, the APDP will identify drugs which qualify under the new statute for periods of exclusivity (during which ANDAs and paper NDAs for those drugs may not be submitted or made effective as identified below) and will provide information on the current patent status of the listed drugs. Exclusivity prevents the filing and/or approval of ANDAs or paper NDAs. It does not prevent the filing or approval of a second full NDA. Applications qualifying for periods of exclusivity are:

- (1) A new drug application approved between January 1, 1982, and September 24, 1984, for a drug product all active ingredients (including any ester or salt of the active ingredient) of which had never been approved in any other application. Approval of an ANDA or paper NDA for the same drug may not be made effective for a period of ten years from the date of the approval of the original application.

- (2) A new drug application approved after September 24, 1984, for a drug product all active ingredients (including any ester or salt of the active ingredient) of which had never been approved in any other new drug application. Generally, no subsequent ANDA or paper NDA for the same drug may be submitted for a period of five years from the date of approval of the original application, except that such an application may be submitted after four years if it contains a certification that a patent claiming the drug is invalid or will not be infringed by the product for which approval is sought.
- (3) A new drug application approved after September 24, 1984, for a drug product involving an active ingredient (or any ester or salt of that active ingredient) that has been approved in an earlier new drug application and which includes reports of new clinical investigations (other than bioavailability studies). Such investigations must have been conducted or sponsored by the applicant or for which the applicant had a right of reference, and the investigations must have been essential to approval of the application. If these requirements are met, the approval of a subsequent ANDA or paper NDA may not be made effective for the same drug before the expiration of three years from the date of approval of the original application.
- (4) A supplement to a new drug application approved after September 24, 1984, which contains reports of new clinical investigations (other than bioavailability studies) essential to the approval of the supplement and conducted or sponsored by the applicant or to which the applicant had a right of reference. The approval of a subsequent application for a change approved in the supplement may not be made effective for three years from the date of approval of the original supplement.
- (5) A new drug application (or supplement to a new drug application) approved during the period from January 1, 1982, to September 24, 1984, which includes an active ingredient (including any ester or salt of the active ingredient) that has been approved in another application. The approval of a subsequent application for the drug or a significant change made in a supplement may not be made effective for two years from September 24, 1984.

The Act required approved new drug applications to be supplemented with the required patent information by October 24, 1984. Patent information must now be filed with all newly submitted drug applications, and no NDA may be approved after September 24, 1984, without the pertinent patent information. The patent numbers and the expiration dates of any appropriate product or use patent on a marketed drug that is the subject of an approved NDA will be published in the APDP. Patent information on unapproved applications or on patents beyond the scope (i.e., process or manufacturing) of the Act will not be published.

The following explains how the APDP implements this.

### Antibiotics, Insulin and Biologicals

Title I of the Act has been interpreted by the Agency not to include products approved under sections 506 or 507 of the Federal Food, Drug and Cosmetic Act (antibiotic and insulin products). Because of this, (1) antibiotic and insulin products are not considered eligible for exclusivity protection, (2) holders of approved applications for insulin and antibiotic products need not submit the patent information as required of NDA application holders, and (3) Antibiotic Form 6 sponsors are not required to provide the patent certification statement which must be included in ANDAs.

However, Title II, the patent term restoration portion of the Act, specifically addresses antibiotic, non-antibiotic, and human biological products (as those terms are used in the Federal Food, Drug and Cosmetic and Public Health Service Acts) in its provisions.

### Bioavailability/Bioequivalence Requirements

The therapeutic equivalence evaluation codes in Appendix D of the APDP will enable firms to determine whether in vitro and/or in vivo bioavailability/bioequivalence study data must be included with their ANDA submissions.

Currently, drugs approved prior to 1962 fall into three major biopharmaceutical classes: (1) those which pose an actual or potential bioequivalence problem, and for which demonstration of bioequivalence through in vivo testing and acceptable dissolution performance is necessary; (2) those which pose an actual or potential bioequivalence problem but for which an in vivo study may be waived if acceptable dissolution performance is demonstrated (the list of such drugs is provided under TABLE I); and (3) those which pose no actual or potential bioequivalence problem and for which the only biopharmaceutical requirement is demonstration of acceptable dissolution for solid oral dosage forms.

All firms submitting an abbreviated new drug application for a single source drug product or a drug product which was first approved after 1962 will be required to demonstrate in vivo bioequivalence or else submit information sufficient to permit the Agency to waive demonstration of in vivo bioequivalence. Manufacturers of drug products formulated in dosage forms which do not present bioequivalence problems, such as an intravenous solution, may request that the in vivo bioequivalence requirement be waived.

Before the passage of the Drug Price Competition and Patent Term Restoration Act, the Agency approved various drugs with bioavailability/bioequivalence problems and deferred the in vivo testing requirement for a number of reasons. The new law requires information to show that the proposed ANDA drug product is bioequivalent to the listed drug. Therefore, new applications for drugs such as amitriptyline hydrochloride which formerly may have been approved without an in vivo study now require an in vivo study as a condition for approval under the new Act.

### Topicals

In the absence of contrary data, FDA regarded all pharmaceutically equivalent topical products of pre-1962 (DESI) drugs to be therapeutically equivalent. However, the Agency required that applicants for topical drug products initially approved after 1962, including "paper NDAs," either demonstrate the safety and efficacy of their products through clinical trials or through a bioequivalence study in order to be approved and evaluated as therapeutically equivalent.

The new Act requires applicants to demonstrate the bioequivalence of their topical drug product to the listed drug as one of the requirements for ANDA approval. This is the same policy that is presently being used in the "paper NDA" approval process. The Agency is now reviewing the therapeutic equivalence evaluation policy that has been made on the pre-1962 topical products to determine whether a change in this policy is warranted. In the meantime, an in vivo demonstration of bioequivalence will be required for approval of all topical products unless a waiver or in vitro alternatives can be justified by the applicant.

### OTC Drug Products Eligible for Abbreviated New Drug Applications

Previous editions of the APDP excluded OTC drug products, because the main purpose of that publication was to provide information to states regarding FDA's recommendation as to which generic prescription drug products were acceptable candidates for drug product selection. With the passage of the Drug Price Competition and Patent Term Restoration Act of 1984, the Agency now has the responsibility to publish an up-to-date list of all marketed drug products, OTC as well as prescription, that have been approved for safety and efficacy and for which new drug applications are required. There are some drugs for which there are both approved and unapproved OTC drug products in the market place. This situation occurs as a result of the Agency's current OTC compliance policy which allows the marketing of various unapproved OTC drug products pending the effective date of the applicable final OTC monograph. The OTC products included in APDP cumulative supplement TABLE II are limited to those for which approved applications are currently required as a condition of marketing. Appropriate patent numbers, exclusivity information, and expiration dates are also included.

## NDA's Approved by the Office of Biological Research and Review Not Previously Published in the APDP

All products accepted and approved under Section 505 of the Act as NDAs by the Office of Biological Research and Review (OBRR) will now be published in the APDP (see TABLE III). The application holder should have submitted relevant patent and exclusivity information as for other NDA drug products. These products will be listed drugs and ANDA applications may be submitted for marketing of drugs from this group. Appropriate patent numbers, exclusivity information, and expiration dates are also included.

### Patent and Exclusivity Information

It was originally planned that TABLE IV of Cumulative Supplement 2 to the APDP would contain patent and exclusivity information. Because some firms submitted patent information in excess of that covered by the statute, FDA has reviewed all of the patent information to assure that only appropriate patents are listed. The patents that FDA regards as covered by the statutory provisions for submission of patent information are those on the active ingredient or ingredients, or use patents for a particular indication or method of using the product. The Agency will not publish patents relating to chemical intermediates, methods of manufacturing, excipients or formulations. TABLE IV contains patent numbers and expiration dates and, for drug products approved after 1982, the date of approval and application number as required by the Act.

Firms submitting ANDAs after September 24, 1984, that certified that no patent information had been filed should now amend their applications with the appropriate patent certification statement.

TABLES II-IV now identify all drugs which qualify under the new statute for periods of exclusivity. (See pages A-1 & A-2 of the Addendum for an explanation of exclusivity).

FDA has finished reviewing all patent and exclusivity information received initially from interested parties. The Agency believes TABLES II-IV now contain all appropriate patent and exclusivity information that the Agency regards as being covered by the new statute. This table will be updated monthly to include appropriate patent and exclusivity information for new approvals. The exclusivity information column in TABLES II-IV designates the date on which the exclusivity ends and the basis for the exclusivity through the use of codes as explained on pages A-6 and A-7.

FDA invites comments from all interested parties on whether it has excluded any patent or exclusivity information that should have been included, or included patent or exclusivity information that should have been excluded. Any revisions to the list will be published in subsequent supplements.

DO TO SPACE LIMITATIONS IN THE EXCLUSIVITY COLUMNS OF TABLES I-IV THE FOLLOWING ABBREVIATIONS HAVE BEEN DEVELOPED. PLEASE REFER BACK TO THIS PAGE FOR AN EXPLANATION OF THE EXCLUSIVITY ABBREVIATIONS FOUND IN THE TABLES.

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)      |

REFERENCES

NEW DOSING SCHEDULE

|     |  |
|-----|--|
| 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/ELEVEN DAYS DOSING SCHEDULE             |
| D-6 | SEVEN DAYS/NINE DAYS/FIVE DAYS DOSING SCHEDULE   |
| D-7 | BID DOSING                                       |
| D-8 | INTRAVENOUS, EPIDURAL AND INTRATHECAL DOSING     |

INDICATIONS

- 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 POST-MYOCARDIAL INFARCTION
- I-16 ACROMEGALY
- 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 OTITIS MEDIA
- I-22 EXERCISE INDUCED BRONCHOSPASMS
- I-23 MI OR STROKE
- I-24 COMBINED USE WITH NICOTINIC ACID TO LOWER CHOLESTEROL LEVEL
- I-25 BLASTOMYCOSES DERMATITIDES
- I-26 PEDIATRIC SUBARACHNOID VASCULAR
- I-27 PETRIELLIDIUM BOYDII INFECTION
- I-28 HEREDITARY ANGIOEDEMA
- I-29 INTRACORONARY USE
- I-30 PEDIATRIC USE
- I-31 DIRECT ISOTOPIC CYSTOGRAPHY



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

|   |  |  |
|---|--|--|
| ACETAMINOPHEN; ASPIRIN;<br>BUTALBITAL;<br>CAPSULE OR TABLET; ORAL<br>160-165MG; 160-165MG; 50MG             | AMINOPHYLLINE<br>TABLET; ORAL<br>100MG<br>200MG  | ASPIRIN; MEPROBAMATE<br>TABLET; ORAL<br>325MG; 200MG                               |
| ACETAMINOPHEN; ASPIRIN;<br>CAPSULE OR TABLET; ORAL<br>325MG; 325MG; 50MG                                    | ASPIRIN; BUTALBITAL;<br>CAPSULE OR TABLET; ORAL<br>325; 50MG<br>650; 50MG                            | ASPIRIN; METHOCARBAMOL<br>TABLET; ORAL<br>325MG; 200MG                             |
| ACETAMINOPHEN; ASPIRIN;<br>BUTALBITAL;<br>CAPSULE OR TABLET; ORAL<br>160-165MG; 160-165MG; 50MG; 40MG       | ASPIRIN; BUTALBITAL, CAFFEINE<br>CAPSULE OR TABLET; ORAL<br>325MG; 50MG; 40MG;<br>650MG; 50MG; 40MG; | CHLOROTHIAZIDE<br>TABLET; ORAL<br>250MG  |
| ACETAMINOPHEN; ASPIRIN;<br>BUTALBITAL; CAFFEINE<br>CAPSULE OR TABLET; ORAL<br>325MG; 325MG; 50MG; 40MG      | ASPIRIN; CAFFEINE; CARISOPRODOL<br>TABLET; ORAL<br>160MG; 32MG; 200MG                                | ESTROGENS, CONJUGATED; MEPROBAMATE<br>TABLET; ORAL<br>0.4MG; 200MG<br>0.4MG; 400MG |
| ACETAMINOPHEN; BUTALBITAL<br>CAPSULE OR TABLET; ORAL<br>325; 50MG<br>650; 50MG                              | ASPIRIN; CAFFEINE; CARISOPRODOL;<br>CODEINE PHOSPHATE<br>TABLET; ORAL<br>160MG; 32MG; 200MG; 16MG    | HYDROXYZINE HYDROCHLORIDE<br>TABLET; ORAL<br>10MG<br>25MG<br>50MG<br>100MG         |
| ACETAMINOPHEN; BUTALBITAL;<br>CAFFEINE<br>CAPSULE OR TABLET; ORAL<br>325MG; 50MG; 40MG<br>650MG; 50MG; 40MG | ASPIRIN; CARISOPRODOL<br>TABLET; ORAL<br>325MG; 200MG  |  |



**TABLE II. OTC DRUG PRODUCTS WHICH CURRENTLY REQUIRE APPROVED APPLICATIONS AS A CONDITION OF MARKETING**

| <u>ACTIVE INGREDIENT(S)<br/>STRENGTH(S)</u>                 | <u>TRADE NAME<br/>(DOSAGE FORM; ROUTE)</u>        | <u>APPLICANT NAME</u> | <u>NDA NO.<br/>APPROVAL DATE</u> | <u>PATENT NO.<br/>EXP. DATE</u> | <u>EXCLUSIVITY<br/>EXP. DATE</u> |
|---|---|-----------------------|----------------------------------|---------------------------------|----------------------------------|
| ACETAMINOPHEN<br>120MG                                      | NEOPAP<br>(SUPPOSITORY; RECTAL)                   | WEBCON PHARMS/ALCON   | 16-401<br>11-07-68               |                                 |                                  |
| ACETAMINOPHEN<br>650MG                                      | TYLENOL<br>(SUPPOSITORY; RECTAL)                  | MCNEIL LABORATORIES   | 17-756<br>05-26-76               |                                 |                                  |
| ACETAMINOPHEN<br>120MG                                      | TYLENOL<br>(SUPPOSITORY; RECTAL)                  | MCNEIL LABORATORIES   | 17-756<br>05-26-76               |                                 |                                  |
| ACETAMINOPHEN<br>120MG                                      | ACEPHEN<br>(SUPPOSITORY; RECTAL)                  | G AND W LABORATORIES  | 18-060<br>02-09-78               |                                 |                                  |
| ACETAMINOPHEN<br>650MG                                      | ACEPHEN<br>(SUPPOSITORY; RECTAL)                  | G AND W LABORATORIES  | 18-060<br>02-09-78               |                                 |                                  |
| ACETAMINOPHEN<br>650MG                                      | ACETAMINOPHEN<br>(SUPPOSITORY; RECTAL)            | UPSHER-SMITH LABS     | 18-337<br>04-22-80               |                                 |                                  |
| ACETAMINOPHEN<br>120MG                                      | ACETAMINOPHEN<br>(SUPPOSITORY; RECTAL)            | UPSHER-SMITH LABS     | 18-337<br>09-12-83               |                                 |                                  |
| ALUMINUM HYDROXIDE; MAGNESIUM<br>TRISILICATE<br>80MG; 20MG  | GAVISCON<br>(TABLET, CHEWABLE; ORAL)              | MARION LABORATORIES   | 18-685<br>12-09-83               |                                 | NP<br>09-24-86                   |
| ALUMINUM HYDROXIDE; MAGNESIUM<br>TRISILICATE<br>160MG; 40MG | GAVISCON-2<br>(TABLET, CHEWABLE; ORAL)            | MARION LABORATORIES   | 18-685<br>12-09-83               |                                 | NP<br>09-24-86                   |
| BROMPHENIRAMINE MALEATE<br>8MG                              | DIMETANE<br>(TABLET, CONTROLLED<br>RELEASE; ORAL) | AH ROBINS             | 10-799<br>06-10-83               |                                 | RTO<br>09-24-86                  |
| BROMPHENIRAMINE MALEATE<br>12MG                             | DIMETANE<br>(TABLET, CONTROLLED<br>RELEASE; ORAL) | AH ROBINS             | 10-799<br>06-10-83               |                                 | RTO<br>09-24-86                  |
| CHLORHEXIDINE GLUCONATE<br>0.5%                             | HIBITANE<br>(TINCTURE; TOPICAL)                   | ICI AMERICAS          | 18-049<br>12-18-78               |                                 |                                  |
| CHLORHEXIDINE GLUCONATE<br>0.5%                             | HIBISTAT<br>(SOLUTION; TOPICAL)                   | ICI AMERICAS          | 18-300<br>05-23-80               |                                 |                                  |

**TABLE II. OTC DRUG PRODUCTS WHICH CURRENTLY REQUIRE APPROVED APPLICATIONS AS A CONDITION OF MARKETING**

| <u>ACTIVE INGREDIENT(S)<br/>STRENGTH(S)</u>                                     | <u>TRADE NAME<br/>(DOSAGE FORM; ROUTE)</u>              | <u>APPLICANT NAME</u> | <u>NDA NO.<br/>APPROVAL DATE</u> | <u>PATENT NO.<br/>EXP. DATE</u> | <u>EXCLUSIVITY<br/>EXP. DATE</u> |
|---|---|-----------------------|----------------------------------|---------------------------------|----------------------------------|
| CHLORHEXIDINE GLUCONATE<br>4%   | EXIDINE<br>(SOLUTION; TOPICAL)                          | XTTRIUM LABS          | 19-125<br>12-24-84               |                                 |                                  |
| CHLORHEXIDINE GLUCONATE<br>4%   | EXIDINE<br>(AEROSOL; TOPICAL)                           | XTTRIUM LABS          | 19-127<br>12-24-84               |                                 | NDF<br>12-24-87                  |
| CHLORHEXIDINE GLUCONATE<br>4%   | HIBICLENS<br>(SOLUTION; TOPICAL)                        | ICI AMERICAS          | 17-768<br>09-17-76               |                                 |                                  |
| CHLORHEXIDINE GLUCONATE<br>4%   | HIBICLENS<br>(SPONGE; TOPICAL)                          | ICI AMERICAS          | 18-423<br>08-27-81               |                                 |                                  |
| CHLORPHENIRAMINE MALEATE<br>8MG   | TELDRI<br>(CAPSULE, CONTROLLED<br>RELEASE; ORAL)        | MENLEY & JAMES/SKF    | 17-369<br>05-11-78               |                                 |                                  |
| CHLORPHENIRAMINE MALEATE<br>12MG  | TELDRI<br>(CAPSULE, CONTROLLED<br>RELEASE; ORAL)        | MENLEY & JAMES/SKF    | 17-369<br>05-11-78               |                                 |                                  |
| CHLORPHENIRAMINE MALEATE<br>8MG   | CHLOR-TRIMETON<br>(TABLET, CONTROLLED<br>RELEASE; ORAL) | SCHERING              | 07-638<br>10-18-78               |                                 |                                  |
| CHLORPHENIRAMINE MALEATE<br>12MG  | CHLOR-TRIMETON<br>(TABLET, CONTROLLED<br>RELEASE; ORAL) | SCHERING              | 07-638<br>10-18-78               |                                 |                                  |
| CHLORPHENIRAMINE MALEATE;<br>PHENYLPROPANOLAMINE<br>HYDROCHLORIDE<br>8MG; 75MG  | CONTAC<br>(CAPSULE, CONTROLLED<br>RELEASE; ORAL)        | MENLEY & JAMES/SKF    | 18-099<br>02-04-80               |                                 |                                  |
| CHLORPHENIRAMINE MALEATE;<br>PHENYLPROPANOLAMINE<br>HYDROCHLORIDE<br>12MG; 75MG | TRIAMINIC-12<br>(TABLET, CONTROLLED<br>RELEASE; ORAL)   | DORSEY LABS/SANDOZ    | 18-115<br>07-23-81               |                                 |                                  |
| CHLORPHENIRAMINE MALEATE;<br>PHENYLPROPANOLAMINE<br>HYDROCHLORIDE<br>4MG; 25MG  | DEMAZIN<br>(TABLET, CONTROLLED<br>RELEASE; ORAL)        | SCHERING              | 18-556<br>05-14-84               |                                 | NS<br>09-24-86                   |

**TABLE II. OTC DRUG PRODUCTS WHICH CURRENTLY REQUIRE APPROVED APPLICATIONS AS A CONDITION OF MARKETING**

| <u>ACTIVE INGREDIENT(S)</u><br><u>STRENGTH(S)</u>  | <u>TRADE NAME</u><br><u>(DOSAGE FORM; ROUTE)</u>  | <u>APPLICANT NAME</u> | <u>NDA NO.</u><br><u>APPROVAL DATE</u> | <u>PATENT NO.</u><br><u>EXP. DATE</u> | <u>EXCLUSIVITY</u><br><u>EXP. DATE</u> |
|--|---|-----------------------|--|---------------------------------------|--|
| CHLORPHENIRAMINE MALEATE;<br>PHENYLPROPANOLAMINE<br>HYDROCHLORIDE<br>8MG; 75MG                             | PHENYLPROPANOLAMINE HCL<br>W/ CHLORPHENIRAMINE<br>MALEATE<br>(CAPSULE, CONTROLLED<br>RELEASE; ORAL) | CENTRAL PHARMS        | 18-809<br>05-07-84                     |                                       |  |
| CHLORPHENIRAMINE MALEATE;<br>PSEUDOEPHEDRINE SULFATE<br>8MG; 120MG   | CHLOR-TRIMETON<br>(TABLET, CONTROLLED<br>RELEASE; ORAL)   | SCHERING              | 18-397<br>03-31-81                     |                                       |  |
| CHLORPHENIRAMINE POLISTIREX;<br>PHENYLPROPANOLAMINE POLISTIREX<br>EQ 4MG MALEATE/5ML;<br>EQ 37.5MG HCL/5ML | CORSYM<br>(SYRUP; ORAL)   | PENNWALT PHARM        | 18-050<br>01-04-84                     |                                       | NDF<br>09-24-86                        |
| DEXBROMPHENIRAMINE MALEATE;<br>PSEUDOEPHEDRINE SULFATE<br>2MG; 60MG  | DISOPHROL<br>(TABLET; ORAL)   | SCHERING              | 12-394<br>06-03-60                     |                                       | RTO<br>09-24-86                        |
| DEXBROMPHENIRAMINE MALEATE;<br>PSEUDOEPHEDRINE SULFATE<br>6MG; 120MG                                       | DRIXORAL<br>(TABLET, CONTROLLED<br>RELEASE; ORAL)   | SCHERING              | 13-483<br>09-13-82                     |                                       | RTO<br>09-24-86                        |
| DEXBROMPHENIRAMINE MALEATE;<br>PSEUDOEPHEDRINE SULFATE<br>6MG; 120MG                                       | DISOPHROL<br>(TABLET, CONTROLLED<br>RELEASE; ORAL)  | SCHERING              | 13-483<br>09-13-82                     |                                       | RTO<br>09-24-86                        |
| DEXTROMETHORPHAN RESIN COMPLEX<br>EQ 30MG HBR/5ML  | DELSYM<br>(SUSPENSION, CONTROLLED<br>RELEASE; ORAL)   | PENNWALT PHARM        | 18-658<br>10-08-82                     |                                       | NDF<br>09-24-86                        |
| DIPHENHYDRAMINE HYDROCHLORIDE<br>12.5MG/5ML  | BENLYN<br>(SYRUP; ORAL)   | PARKE-DAVIS/W-L       | 06-514<br>08-07-81                     |                                       |  |
| DOXYLAMINE SUCCINATE<br>25MG   | UNISOM<br>(TABLET; ORAL)  | PFIZER                | 18-066<br>10-06-78                     |                                       |  |
| IBUPROFEN<br>200MG   | ADVIL<br>(TABLET; ORAL)   | WHITEHALL LABS/AMHO   | 18-989<br>05-18-84                     | 3385886<br>05-28-85                   | NS<br>09-24-86                         |
| IBUPROFEN<br>200MG   | NUPRIN<br>(TABLET; ORAL)  | UPJOHN MANUFACTURING  | 19-012<br>05-18-84                     | 3385886<br>05-28-85                   | NS<br>09-24-86                         |

TABLE II. OTC DRUG PRODUCTS WHICH CURRENTLY REQUIRE APPROVED APPLICATIONS AS A CONDITION OF MARKETING

| <u>ACTIVE INGREDIENT(S)<br/>STRENGTH(S)</u>   | <u>TRADE NAME<br/>(DOSAGE FORM; ROUTE)</u>                   | <u>APPLICANT NAME</u> | <u>NDA NO.<br/>APPROVAL DATE</u> | <u>PATENT NO.<br/>EXP. DATE</u> | <u>EXCLUSIVITY<br/>EXP. DATE</u> |
|---|--|-----------------------|----------------------------------|---------------------------------|----------------------------------|
| INSULIN SUSPENSION, ISOPHANE,<br>BEEF<br>40 UNITS/ML                                      | SEMILENTE INSULIN<br>(INJECTABLE; INJECTION)                 | SQUIBB-NOVO           | 17-929<br>02-08-77               |                                 |                                  |
| INSULIN SUSPENSION, ISOPHANE,<br>BEEF<br>100 UNITS/ML                                     | SEMILENTE INSULIN<br>(INJECTABLE; INJECTION)                 | SQUIBB-NOVO           | 17-929<br>02-08-77               |                                 |                                  |
| INSULIN SUSPENSION, ISOPHANE,<br>BIOSYNTHETIC HUMAN<br>100 UNITS/ML                       | HUMULIN N<br>(INJECTABLE; INJECTION)                         | ELI LILLY             | 18-781<br>10-28-82               |                                 |                                  |
| INSULIN SUSPENSION, ISOPHANE,<br>MIXED BEEF AND PORK<br>40 UNITS/ML                       | NPH ILETIN (BEEF-PORK)<br>(INJECTABLE; INJECTION)            | LILLY RES LABS DIV    | 17-936<br>02-08-77               |                                 |                                  |
| INSULIN SUSPENSION, ISOPHANE,<br>MIXED BEEF AND PORK<br>100 UNITS/ML                      | NPH ILETIN (BEEF-PORK)<br>(INJECTABLE; INJECTION)            | LILLY RES LABS DIV    | 17-936<br>02-08-77               |                                 |                                  |
| INSULIN SUSPENSION, ISOPHANE,<br>PURIFIED BEEF<br>100 UNITS/ML                            | NPH ILETIN II<br>(INJECTABLE; INJECTION)                     | ELI LILLY             | 18-479<br>06-12-80               |                                 |                                  |
| INSULIN SUSPENSION, ISOPHANE,<br>PURIFIED PORK<br>100 UNITS/ML                            | INSULIN INSULATARD NPH<br>NORDISK<br>(INJECTABLE; INJECTION) | NORDISK               | 18-194<br>01-16-80               |                                 |                                  |
| INSULIN SUSPENSION, ISOPHANE,<br>PURIFIED PORK<br>100 UNITS/ML                            | NPH ILETIN II (PORK)<br>(INJECTABLE; INJECTION)              | ELI LILLY             | 18-345<br>12-05-79               |                                 |                                  |
| INSULIN SUSPENSION, ISOPHANE,<br>PURIFIED PORK<br>100 UNITS/ML                            | PROTAPHANE<br>(INJECTABLE; INJECTION)                        | SQUIBB-NOVO           | 18-623<br>07-30-81               |                                 |                                  |
| INSULIN SUSPENSION, ISOPHANE,<br>PURIFIED PORK; INSULIN,<br>PURIFIED PORK<br>100 UNITS/ML | INSULIN NORDISK MIXTARD<br>(PORK)<br>(INJECTABLE; INJECTION) | NORDISK               | 18-195<br>01-16-80               |                                 |                                  |

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| <u>ACTIVE INGREDIENT(S)</u><br><u>STRENGTH(S)</u>   | <u>TRADE NAME</u><br><u>(DOSAGE FORM; ROUTE)</u>             | <u>APPLICANT NAME</u> | <u>NDA NO.</u><br><u>APPROVAL DATE</u> | <u>PATENT NO.</u><br><u>EXP. DATE</u> | <u>EXCLUSIVITY</u><br><u>EXP. DATE</u> |
|---|--|-----------------------|--|---------------------------------------|--|
| INSULIN SUSPENSION, PROTAMINE ZINC, MIXED BEEF AND PORK<br>100 UNITS/ML                               | PROTAMINE, ZINC & ILETIN (BEEF-PORK) (INJECTABLE; INJECTION) | ELI LILLY             | 17-932<br>02-08-77                     |                                       |  |
| INSULIN SUSPENSION, PROTAMINE ZINC, MIXED BEEF AND PORK; INSULIN, MIXED BEEF AND PORK<br>100 UNITS/ML | PROTAMINE, ZINC & ILETIN (BEEF-PORK) (INJECTABLE; INJECTION) | ELI LILLY             | 17-932<br>02-08-77                     |                                       |  |
| INSULIN SUSPENSION, PROTAMINE ZINC, PURIFIED BEEF<br>40 UNITS/ML                                      | PROTAMINE ZINC INSULIN (INJECTABLE; INJECTION)               | ER SQUIBB AND SONS    | 17-928<br>02-08-77                     |                                       |  |
| INSULIN SUSPENSION, PROTAMINE ZINC, PURIFIED BEEF<br>100 UNITS/ML                                     | PROTAMINE ZINC INSULIN (INJECTABLE; INJECTION)               | ER SQUIBB AND SONS    | 17-928<br>02-08-77                     |                                       |  |
| INSULIN SUSPENSION, PROTAMINE ZINC, PURIFIED BEEF; INSULIN, PURIFIED BEEF<br>100 UNITS/ML             | PROTAMINE ZINC AND ILETIN II (INJECTABLE; INJECTION)         | ELI LILLY             | 18-476<br>06-12-80                     |                                       |  |
| INSULIN SUSPENSION, PROTAMINE ZINC, PURIFIED PORK; INSULIN, PURIFIED PORK<br>100 UNITS/ML             | PROTAMINE ZINC AND ILETIN II(PORK) (INJECTABLE; INJECTION)   | ELI LILLY             | 18-346<br>12-05-79                     |                                       |  |
| INSULIN ZINC SUSPENSION, BEEF<br>40 UNITS/ML  | LENTE INSULIN (INJECTABLE; INJECTION)                        | SQUIBB-NOVO           | 17-998<br>02-08-77                     |                                       |  |
| INSULIN ZINC SUSPENSION, BEEF<br>100 UNITS/ML   | LENTE INSULIN (INJECTABLE; INJECTION)                        | SQUIBB-NOVO           | 17-998<br>02-08-77                     |                                       |  |
| INSULIN ZINC SUSPENSION, BIOSYNTHETIC HUMAN<br>100 UNITS/ML   | MONOTARD HUMAN (INJECTABLE; INJECTION)                       | SQUIBB-NOVO           | 18-777<br>08-30-83                     |                                       |  |
| INSULIN ZINC SUSPENSION, EXTENDED, PURIFIED BEEF<br>100 UNITS/ML                                      | ULTRATARD (INJECTABLE; INJECTION)                            | SQUIBB-NOVO           | 18-385<br>03-17-80                     |                                       |  |

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| <u>ACTIVE INGREDIENT(S)<br/>STRENGTH(S)</u>                         | <u>TRADE NAME<br/>(DOSAGE FORM; ROUTE)</u>        | <u>APPLICANT NAME</u> | <u>NDA NO.<br/>APPROVAL DATE</u> | <u>PATENT NO.<br/>EXP. DATE</u> | <u>EXCLUSIVITY<br/>EXP. DATE</u> |
|---|---|-----------------------|----------------------------------|---------------------------------|----------------------------------|
| INSULIN ZINC SUSPENSION,<br>EXTENDED, PURIFIED BEEF<br>100 UNITS/ML | ULTRALENTE INSULIN<br>(INJECTABLE; INJECTION)     | SQUIBB-NOVO           | 17-997<br>02-08-77               |                                 |                                  |
| INSULIN ZINC SUSPENSION, PROMPT,<br>BEEF<br>100 UNITS/ML            | SEMILENTE INSULIN<br>(INJECTABLE; INJECTION)      | SQUIBB-NOVO           | 17-996<br>02-08-77               |                                 |                                  |
| INSULIN ZINC SUSPENSION, PROMPT,<br>PURIFIED PORK<br>100 UNITS/ML   | SEMITARD<br>(INJECTABLE; INJECTION)               | SQUIBB-NOVO           | 18-382<br>03-17-80               |                                 |                                  |
| INSULIN ZINC SUSPENSION,<br>PURIFIED BEEF<br>100 UNITS/ML           | LENTE ILETIN II<br>(INJECTABLE; INJECTION)        | ELI LILLY             | 18-477<br>06-12-80               |                                 |                                  |
| INSULIN ZINC SUSPENSION,<br>PURIFIED BEEF AND PORK<br>100 UNITS/ML  | LENTARD<br>(INJECTABLE; INJECTION)                | SQUIBB-NOVO           | 18-384<br>03-17-80               |                                 |                                  |
| INSULIN ZINC SUSPENSION,<br>PURIFIED PORK<br>100 UNITS/ML           | LENTE ILETIN II (PORK)<br>(INJECTABLE; INJECTION) | ELI LILLY             | 18-347<br>12-05-79               |                                 |                                  |
| INSULIN ZINC SUSPENSION,<br>PURIFIED PORK<br>100 UNITS/ML           | MONOTARD<br>(INJECTABLE; INJECTION)               | SQUIBB-NOVO           | 18-383<br>03-17-80               |                                 |                                  |
| INSULIN, BIOSYNTHETIC HUMAN<br>100 UNITS/ML                         | ACTRAPID HUMAN<br>(INJECTABLE; INJECTION)         | SQUIBB-NOVO           | 18-778<br>08-30-83               |                                 |                                  |
| INSULIN, BIOSYNTHETIC HUMAN<br>100 UNITS/ML                         | HUMULIN R<br>(INJECTABLE; INJECTION)              | ELI LILLY             | 18-780<br>10-28-82               |                                 |                                  |
| INSULIN, PORK<br>40 UNITS/ML  | INSULIN<br>(INJECTABLE; INJECTION)                | SQUIBB-NOVO           | 17-926<br>02-08-77               |                                 |                                  |
| INSULIN, PORK<br>100 UNITS/ML                                       | INSULIN<br>(INJECTABLE; INJECTION)                | SQUIBB-NOVO           | 17-926<br>02-08-77               |                                 |                                  |

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| <u>ACTIVE INGREDIENT(S)<br/>STRENGTH(S)</u>  | <u>TRADE NAME<br/>(DOSAGE FORM; ROUTE)</u>                 | <u>APPLICANT NAME</u> | <u>NDA NO.<br/>APPROVAL DATE</u> | <u>PATENT NO.<br/>EXP. DATE</u> | <u>EXCLUSIVITY<br/>EXP. DATE</u> |
|--|--|-----------------------|----------------------------------|---------------------------------|----------------------------------|
| INSULIN, PURIFIED BEEF<br>100 UNITS/ML   | REGULAR ILETIN II<br>(INJECTABLE; INJECTION)               | ELI LILLY             | 18-478<br>06-12-80               |                                 |                                  |
| INSULIN, PURIFIED PORK<br>100 UNITS/ML   | INSULIN NORDISK QUICK<br>(PORK)<br>(INJECTABLE; INJECTION) | NORDISK INSULIN LABS  | 18-193<br>01-16-80               |                                 |                                  |
| INSULIN, PURIFIED PORK<br>100 UNITS/ML   | REGULAR ILETIN II (PORK)<br>(INJECTABLE; INJECTION)        | ELI LILLY             | 18-344<br>12-05-79               |                                 |                                  |
| INSULIN, PURIFIED PORK<br>100 UNITS/ML   | ACTRAPID<br>(INJECTABLE; INJECTION)                        | SQUIBB-NOVO           | 18-381<br>03-17-80               |                                 |                                  |
| INSULIN SUSPENSION,<br>ISOPHANE, PURIFIED HUMAN<br>100 UNITS/ML                      | NOVOLIN N<br>(INJECTABLE; INJECTION)                       | NOVO INDUSTRI A/S     | 19-065<br>01-23-85               |                                 |                                  |
| NONOXYNOL-9<br>1GM   | TODAY<br>(SPONGE; VAGINAL)                                 | VLI CORPORATION       | 18-683<br>04-01-83               |                                 | NDF<br>09-24-86                  |
| POTASSIUM IODIDE<br>130MG  | THYRO-BLOCK<br>(TABLET; ORAL)                              | WALLACE LABS/C-W      | 18-307<br>11-09-79               |                                 |                                  |
| POTASSIUM IODIDE<br>1GM/ML   | POTASSIUM IODIDE<br>(SOLUTION; ORAL)                       | ROXANE LABORATORIES   | 18-551<br>02-19-82               |                                 | NDF<br>09-24-86                  |
| POTASSIUM IODIDE<br>130MG  | IOSAT<br>(TABLET; ORAL)                                    | ANBEX                 | 18-664<br>10-14-82               |                                 |                                  |
| PSEUDOEPHEDRINE HYDROCHLORIDE<br>120MG   | SUDAFED S.A.<br>(CAPSULE, CONTROLLED<br>RELEASE; ORAL)     | BURROUGHS WELLCOME    | 17-941<br>01-15-79               |                                 |                                  |
| PSEUDOEPHEDRINE HYDROCHLORIDE;<br>TRIPROLIDINE HYDROCHLORIDE<br>30MG/5ML; 1.25MG/5ML | ACTIFED<br>(SYRUP; ORAL)                                   | BURROUGHS WELLCOME    | 11-935<br>11-26-82               |                                 | RTO<br>09-24-86                  |
| PSEUDOEPHEDRINE HYDROCHLORIDE;<br>TRIPROLIDINE HYDROCHLORIDE<br>60MG; 2.5MG          | ACTIFED<br>(TABLET; ORAL)                                  | BURROUGHS WELLCOME    | 11-936<br>11-26-82               |                                 | RTO<br>09-24-86                  |

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| <u>ACTIVE INGREDIENT(S)<br/>STRENGTH(S)</u>   | <u>TRADE NAME<br/>(DOSAGE FORM; ROUTE)</u>                   | <u>APPLICANT NAME</u> | <u>NDA NO.<br/>APPROVAL DATE</u> | <u>PATENT NO.<br/>EXP. DATE</u> | <u>EXCLUSIVITY<br/>EXP. DATE</u> |
|---|--|-----------------------|----------------------------------|---------------------------------|----------------------------------|
| PSEUDOEPHEDRINE HYDROCHLORIDE;<br>TRIPROLDINE HYDROCHLORIDE<br>60MG; 2.5MG          | ACTIFED<br>(CAPSULE; ORAL)                                   | BURROUGHS WELLCOME    | 19-208<br>01-15-85               |                                 | RTO<br>09-24-86                  |
| PSEUDOEPHEDRINE HYDROCHLORIDE;<br>TRIPROLDINE HYDROCHLORIDE<br>30MG/5ML; 1.25MG/5ML | ALLERBAN PLUS<br>(SYRUP; ORAL)                               | BAY LABORATORIES      | 88-116<br>03-04-83               |                                 | RTO<br>09-24-86                  |
| PSEUDOEPHEDRINE HYDROCHLORIDE;<br>TRIPROLDINE HYDROCHLORIDE<br>60MG; 2.5MG          | TRI-SUDO<br>(TABLET; ORAL)                                   | MD PHARMACEUTICAL     | 85-024<br>01-10-84               |                                 | RTO<br>09-24-86                  |
| PSEUDOEPHEDRINE HYDROCHLORIDE;<br>TRIPROLDINE HYDROCHLORIDE<br>60MG; 2.5MG          | TRIPODRINE<br>(TABLET; ORAL)                                 | DANBURY PHARMACAL     | 88-112<br>01-20-83               |                                 | RTO<br>09-24-86                  |
| PSEUDOEPHEDRINE HYDROCHLORIDE;<br>TRIPROLDINE HYDROCHLORIDE<br>30MG/5ML; 1.25MG/5ML | TRIOFED<br>(SYRUP; ORAL)                                     | NATL PHARM MFG/BARRE  | 88-115<br>03-04-83               |                                 | RTO<br>09-24-86                  |
| PSEUDOEPHEDRINE HYDROCHLORIDE;<br>TRIPROLDINE HYDROCHLORIDE<br>30MG/5ML; 1.25MG/5ML | TRIOPOSED<br>(SYRUP; ORAL)                                   | HALSEY DRUG           | 88-213<br>03-30-84               |                                 | RTO<br>09-24-86                  |
| PSEUDOEPHEDRINE HYDROCHLORIDE;<br>TRIPROLDINE HYDROCHLORIDE<br>60MG; 2.5MG          | TRIPROLDINE HCL<br>AND PSEUDOEPHEDRINE HCL<br>(TABLET; ORAL) | CHELSEA LABORATORIES  | 88-118<br>01-26-84               |                                 | RTO<br>09-24-86                  |
| PSEUDOEPHEDRINE HYDROCHLORIDE;<br>TRIPROLDINE HYDROCHLORIDE<br>60MG; 2.5MG          | TRIOPOSED<br>(TABLET; ORAL)                                  | HALSEY DRUG           | 88-192<br>05-01-84               |                                 | RTO<br>09-24-86                  |
| PSEUDOEPHEDRINE HYDROCHLORIDE;<br>TRIPROLDINE HYDROCHLORIDE<br>60MG; 2.5MG          | TRIPROLDINE<br>AND PSEUDOEPHEDRINE<br>(TABLET; ORAL)         | SOLAR PHARMACEUTICAL  | 88-318<br>01-13-84               |                                 | RTO<br>09-24-86                  |
| PSEUDOEPHEDRINE HYDROCHLORIDE;<br>TRIPROLDINE HYDROCHLORIDE<br>30MG/5ML; 1.25MG/5ML | TRIOPOSED<br>(SYRUP; ORAL)                                   | HALSEY DRUG           | 88-213<br>05-01-84               |                                 | RTO<br>09-24-86                  |

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| <u>ACTIVE INGREDIENT(S)<br/>STRENGTH(S)</u> | <u>TRADE NAME<br/>(DOSAGE FORM; ROUTE)</u>       | <u>APPLICANT NAME</u> | <u>NDA NO.<br/>APPROVAL DATE</u> | <u>PATENT NO.<br/>EXP. DATE</u> | <u>EXCLUSIVITY<br/>EXP. DATE</u> |
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| PSEUDOEPHEDRINE SULFATE<br>120MG            | AFRINOL<br>(TABLET, CONTROLLED<br>RELEASE; ORAL) | SCHERING              | 18-191<br>10-30-80               |                                 |                                  |
| TIOCONAZOLE<br>1%                           | TROSYD<br>(CREAM; TOPICAL)                       | PFIZER CEN RES/PFIZR  | 18-682<br>02-18-83               | 4062966<br>12-13-94             | NCE<br>02-18-93                  |
| TRIPROLIDINE HYDROCHLORIDE<br>2.5MG         | ACTIDIL<br>(TABLET; ORAL)                        | BURROUGHS WELLCOME    | 11-110<br>04-14-58               |                                 | RTO<br>09-24-86                  |
| TRIPROLIDINE HYDROCHLORIDE<br>2.5MG         | TRIPROLIDINE HCL<br>(TABLET; ORAL)               | BOLAR PHARMACEUTICAL  | 84-453<br>02-06-76               |                                 | RTO<br>09-24-86                  |
| TRIPROLIDINE HYDROCHLORIDE<br>2.5MG         | TRIPROLIDINE HCL<br>(TABLET; ORAL)               | DANBURY PHARMACAL     | 85-094<br>02-07-77               |                                 | RTO<br>09-24-86                  |
| TRIPROLIDINE HYDROCHLORIDE<br>2.5MG         | TRIPROLIDINE HCL<br>(TABLET; ORAL)               | DRUMMER/PHOENIX       | 85-610<br>03-21-78               |                                 | RTO<br>09-24-86                  |
| TRIPROLIDINE HYDROCHLORIDE<br>1.25MG/5ML    | ACTIDIL<br>(SYRUP; ORAL)                         | BURROUGHS WELLCOME    | 11-496<br>07-24-58               |                                 | RTO<br>09-24-86                  |
| TRIPROLIDINE HYDROCHLORIDE<br>1.25MG/5ML    | BAYDYL<br>(SYRUP; ORAL)                          | BAY LABORATORIES      | 87-963<br>01-18-83               |                                 | RTO<br>09-24-86                  |
| TRIPROLIDINE HYDROCHLORIDE<br>1.25MG/5ML    | TRIPROLIDINE HCL<br>(SYRUP; ORAL)                | NATL PHARM MFG/BARRE  | 85-940<br>07-13-79               |                                 | RTO<br>09-24-86                  |
| TRIPROLIDINE HYDROCHLORIDE<br>1.25MG/5ML    | TRIPROLIDINE HCL<br>(SYRUP; ORAL)                | PHARMS ASSOC/BEACH    | 87-514<br>02-10-82               |                                 | RTO<br>09-24-86                  |



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|---|--|-----------------------|----------------------------------|---------------------------------|----------------------------------|
| ANTICOAGULANT CITRATE DEXTROSE SOLUTION USP                 | NONE (INJECTABLE; INJECTION)               | CUTTER BIOL/MILES     | 10-102<br>12-14-61               |                                 |                                  |
| ANTICOAGULANT CITRATE DEXTROSE SOLUTION USP                 | NONE (INJECTABLE; INJECTION)               | DELMED                | 11-912<br>9-2-59                 |                                 |                                  |
| ANTICOAGULANT CITRATE DEXTROSE SOLUTION USP                 | NONE (INJECTABLE; INJECTION)               | TRAVENOL LABS         | 10-855<br>06-11-59               |                                 |                                  |
| ANTICOAGULANT CITRATE DEXTROSE SOLUTION USP                 | NONE (INJECTABLE; INJECTION)               | TRAVENOL LABS         | 16-918<br>3-17-78                |                                 |                                  |
| ANTICOAGULANT CITRATE PHOSPHATE DEXTROSE ADENINE-1 SOLUTION | NONE (INJECTABLE; INJECTION)               | CUTTER BIOL/MILES     | 80-77<br>11-6-80                 |                                 |                                  |
| ANTICOAGULANT CITRATE PHOSPHATE DEXTROSE ADENINE SOLUTION   | NONE (INJECTABLE; INJECTION)               | DELMED                | 78-519<br>4-23-80                |                                 |                                  |
| ANTICOAGULANT CITRATE PHOSPHATE DEXTROSE ADENINE SOLUTION   | NONE (INJECTABLE; INJECTION)               | TERUMO AMERICA        | 82-528<br>11-3-82                |                                 |                                  |
| ANTICOAGULANT CITRATE PHOSPHATE DEXTROSE ADENINE SOLUTION   | NONE (INJECTABLE; INJECTION)               | TRAVENOL LABS         | 77-420<br>5-12-78                |                                 |                                  |
| ANTICOAGULANT CITRATE PHOSPHATE DEXTROSE SOLUTION USP       | NONE (INJECTABLE; INJECTION)               | CUTTER BIOL/MILES     | 16-527<br>6-22-70                |                                 |                                  |
| ANTICOAGULANT CITRATE PHOSPHATE DEXTROSE SOLUTION USP       | NONE (INJECTABLE; INJECTION)               | CUTTER BIOL/MILES     | 80-222<br>8-23-82                |                                 |                                  |
| ANTICOAGULANT CITRATE PHOSPHATE DEXTROSE SOLUTION USP       | NONE (INJECTABLE; INJECTION)               | DELMED                | 16-907<br>5-15-73                |                                 |                                  |
| ANTICOAGULANT CITRATE PHOSPHATE DEXTROSE SOLUTION USP       | NONE (INJECTABLE; INJECTION)               | TERUMO AMERICA        | 78-1211<br>6-10-81               |                                 |                                  |
| ANTICOAGULANT CITRATE PHOSPHATE DEXTROSE SOLUTION USP       | NONE (INJECTABLE; INJECTION)               | TRAVENOL LABS         | 17-401<br>12-6-77                |                                 |                                  |
| ANTICOAGULANT CITRATE PHOSPHATE DEXTROSE SOLUTION USP       | NONE (INJECTABLE; INJECTION)               | TRAVENOL LABS         | 81-1012<br>6-28-83               |                                 |                                  |

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|---|---|-----------------------|----------------------------------|---------------------------------|----------------------------------|
| ANTICOAGULANT CITRATE PHOSPHATE<br>DEXTROSE SOLUTION USP WITH:<br>AS-1: DEXTROSE USP 2.2GM/100ML,<br>SODIUM CHLORIDE USP 0.9GM/100ML,<br>MANNITOL USP 0.75GM/100ML,<br>ADENINE 0.27GM/100ML   | ADSOR <sup>R</sup> RED CELL<br>PRESERVATION SOLUTION<br>(INJECTABLE; INJECTION) | TRAVENOL LABS         | 81-1104<br>5-16-83               |                                 |                                  |
| ANTICOAGULANT CITRATE PHOSPHATE<br>DOUBLE DEXTROSE SOLUTION WITH:<br>AS-2: CITRIC ACID USP<br>0.42GM/100ML, DIBASIC SODIUM<br>PHOSPHATE USP 0.285GM/100ML,<br>SODIUM CHLORIDE USP 0.718<br>GM/100ML, ADENINE 0.017GM/100ML,<br>DEXTROSE USP 0.396GM/100ML,<br>SODIUM CITRATE USP 0.588GM/100ML        | AS-2 NUTRICEL ADITIVE<br>SYSTEM<br>(INJECTABLE; INJECTION)                      | CUTTER BIOL/MILES     | 82-915<br>9-22-83                |                                 |                                  |
| ANTICOAGULANT CITRATE PHOSPHATE<br>DOUBLE DEXTROSE SOLUTION WITH:<br>AS-3: CITRIC ACID USP 0.042<br>GM/100ML, MONOBASIC SODIUM<br>PHOSPHATE USP 0.276GM/100ML,<br>SODIUM CHLORIDE USP 0.410<br>GM/100ML, ADENINE 0.30<br>GM/100ML, DEXTROSE USP 1.10<br>GM/100ML, SODIUM CITRATE USP<br>0.588GM/100ML | AS-3 NUTRICEL ADDITIVE<br>SYSTEM<br>(INJECTABLE; INJECTION)                     | CUTTER BIOL/MILES     | 82-915<br>10-19-84               |                                 |                                  |
| ANTICOAGULANT HEPARIN SOLUTION<br>USP   | NONE<br>(INJECTABLE; INJECTION)   | DELMED                | 77-822<br>5-17-78                |                                 |                                  |
| ANTICOAGULANT HEPARIN SOLUTION<br>USP   | NONE<br>(INJECTABLE; INJECTION)   | TRAVENOL LABS         | 81-1217<br>5-16-83               |                                 |                                  |
| ANTICOAGULANT SODIUM CITRATE<br>SOLUTION USP  | NONE<br>(INJECTABLE; INJECTION)   | ALPHA THERAPEUTIC     | 81-416<br>10-12-83               |                                 |                                  |
| ANTICOAGULANT SODIUM CITRATE<br>SOLUTION USP  | NONE<br>(INJECTABLE; INJECTION)   | CUTTER BIOL/MILES     | 76-305<br>6-30-78                |                                 |                                  |
| ANTICOAGULANT SODIUM CITRATE<br>SOLUTION USP  | NONE<br>(INJECTABLE; INJECTION)   | DELMED                | 16-702<br>12-28-70               |                                 |                                  |

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| <u>ACTIVE INGREDIENT(S)<br/>STRENGTH(S)</u>                             | <u>TRADE NAME<br/>(DOSAGE FORM; ROUTE)</u> | <u>APPLICANT NAME</u> | <u>NDA NO.<br/>APPROVAL DATE</u> | <u>PATENT NO.<br/>EXP. DATE</u> | <u>EXCLUSIVITY<br/>EXP. DATE</u> |
|---|--|-----------------------|----------------------------------|---------------------------------|----------------------------------|
| ANTICOAGULANT SODIUM CITRATE<br>SOLUTION USP                            | NONE<br>(INJECTABLE; INJECTION)            | TERUMO AMERICA        | 78-1214<br>2-8-80                |                                 |                                  |
| ANTICOAGULANT SODIUM CITRATE<br>SOLUTION USP                            | NONE<br>(INJECTABLE; INJECTION)            | TRAVENOL LABS         | 77-923<br>1-20-78                |                                 |                                  |
| DEXTRAN 40, 10%<br>10GM/100ML IN<br>DEXTROSE 5%<br>5GM/100ML            | NONE<br>(INJECTABLE; INJECTION)            | ABBOTT LABORATORIES   | 16-375<br>7-25-67                |                                 |                                  |
| DEXTRAN 40, 10%<br>10GM/100ML IN<br>SODIUM CHLORIDE 0.9%<br>0.9GM/100ML | NONE<br>(INJECTABLE; INJECTION)            | ABBOTT LABORATORIES   | 16-375<br>7-25-67                |                                 |                                  |
| DEXTRAN 75, 6%<br>6GM/100ML IN<br>SODIUM CHLORIDE 0.9%<br>0.9GM/100ML   | NONE<br>(INJECTABLE; INJECTION)            | ABBOTT LABORATORIES   | 8-819<br>3-31-53                 |                                 |                                  |
| DEXTRAN 75, 6%<br>6GM/100ML IN<br>SODIUM CHLORIDE 0.9%<br>0.9GM/100ML   | NONE<br>(INJECTABLE; INJECTION)            | ABBOTT LABORATORIES   | 8-819<br>3-31-53                 |                                 |                                  |
| DEXTRAN 75, 6%<br>6GM/100ML IN<br>SODIUM CHLORIDE 0.9%<br>0.9GM/100ML   | NONE<br>(INJECTABLE; INJECTION)            | ABBOTT LABORATORIES   | 18-253<br>2-4-83                 |                                 |                                  |
| DEXTRAN 40, 10%<br>10GM/100ML IN<br>DEXTROSE 5%<br>5GM/100ML            | NONE<br>(INJECTABLE; INJECTION)            | AMERICAN MCGAW        | 16-767<br>4-6-70                 |                                 |                                  |
| DEXTRAN 40, 10%<br>10GM/100ML IN<br>SODIUM CHLORIDE 0.9%<br>0.9GM/100ML | NONE<br>(INJECTABLE; INJECTION)            | AMERICAN MCGAW        | 16-767<br>4-6-70                 |                                 |                                  |

**TABLE III. NDA'S APPROVED BY THE OFFICE OF BIOLOGICAL RESEARCH AND REVIEW NOT PREVIOUSLY PUBLISHED**

| <u>ACTIVE INGREDIENT(S)<br/>STRENGTH(S)</u>                             | <u>TRADE NAME<br/>(DOSAGE FORM; ROUTE)</u> | <u>APPLICANT NAME</u> | <u>NDA NO.<br/>APPROVAL DATE</u> | <u>PATENT NO.<br/>EXP. DATE</u> | <u>EXCLUSIVITY<br/>EXP. DATE</u> |
|---|--|-----------------------|----------------------------------|---------------------------------|----------------------------------|
| DEXTRAN 70, 6%<br>6GM/100ML IN<br>SODIUM CHLORIDE 0.9%<br>0.9GM/100ML   | NONE<br>(INJECTABLE; INJECTION)            | AMERICAN MCGAW        | 9-024<br>8-18-69                 |                                 |                                  |
| DEXTRAN 40, 10%<br>10GM/100ML IN<br>DEXTROSE 5%<br>5GM/100ML            | NONE<br>(INJECTABLE; INJECTION)            | CUTTER BIOL/MILES     | 16-653<br>9-23-69                |                                 |                                  |
| DEXTRAN 40, 10%<br>10GM/100ML IN<br>SODIUM CHLORIDE 0.9%<br>0.9GM/100ML | NONE<br>(INJECTABLE; INJECTION)            | CUTTER BIOL/MILES     | 16-653<br>9-23-69                |                                 |                                  |
| DEXTRAN 70, 6%<br>6GM/100ML IN<br>SODIUM CHLORIDE 0.9%<br>0.9GM/100ML   | NONE<br>(INJECTABLE; INJECTION)            | CUTTER BIOL/MILES     | 8-716<br>8-11-69                 |                                 |                                  |
| DEXTRAN 40, 10%<br>10GM/100ML IN<br>DEXTROSE 5%<br>5GM/100ML            | NONE<br>(INJECTABLE; INJECTION)            | PHARMACHEM            | 16-836<br>11-14-70               |                                 |                                  |
| DEXTRAN 40, 10%<br>10GM/100ML IN<br>SODIUM CHLORIDE 0.9%<br>0.9GM/100ML | NONE<br>(INJECTABLE; INJECTION)            | PHARMACHEM            | 16-836<br>11-14-70               |                                 |                                  |
| DEXTRAN 75, 6%<br>6GM/100ML IN<br>SODIUM CHLORIDE 0.9%<br>0.9GM/100ML   | NONE<br>(INJECTABLE; INJECTION)            | PHARMACHEM            | 8-564<br>9-19-52                 |                                 |                                  |
| DEXTRAN 75, 6%<br>6GM/100ML IN<br>SODIUM CHLORIDE 0.9%<br>0.9GM/100ML   | NONE<br>(INJECTABLE; INJECTION)            | PHARMACHEM            | 16-759<br>8-19-70                |                                 |                                  |
| DEXTRAN 1<br>150MG/ML IN<br>SODIUM CHLORIDE 0.6%<br>6MG/ML              | PROMIT<br>(INJECTABLE; INJECTION)          | PHARMACIA LABS        | 83-715<br>10-30-84               |                                 | NCE<br>10-30-89                  |

TABLE III. NDAs APPROVED BY THE OFFICE OF BIOLOGICAL RESEARCH AND REVIEW NOT PREVIOUSLY PUBLISHED

| <u>ACTIVE INGREDIENT(S)<br/>STRENGTH(S)</u>                             | <u>TRADE NAME<br/>(DOSAGE FORM; ROUTE)</u>           | <u>APPLICANT NAME</u> | <u>NDA NO.<br/>APPROVAL DATE</u> | <u>PATENT NO.<br/>EXP. DATE</u> | <u>EXCLUSIVITY<br/>EXP. DATE</u> |
|---|--|-----------------------|----------------------------------|---------------------------------|----------------------------------|
| DEXTRAN 40, 10%<br>10GM/100ML IN<br>DEXTROSE 5%<br>5GM/100ML            | RHEOMACRODEX <sup>R</sup><br>(INJECTABLE; INJECTION) | PHARMACIA LABS        | 14-716<br>1-18-67                |                                 |                                  |
| DEXTRAN 40, 10%<br>10GM/100ML IN<br>SODIUM CHLORIDE 0.9%<br>0.9GM/100ML | RHEOMACRODEX <sup>R</sup><br>(INJECTABLE; INJECTION) | PHARMACIA LABS        | 14-716<br>1-18-67                |                                 |                                  |
| DEXTRAN 70, 6%<br>6GM/100ML IN<br>DEXTROSE 5%<br>5GM/100ML              | MACRODEX <sup>R</sup><br>(INJECTABLE; INJECTION)     | PHARMACIA LABS        | 6-826<br>6-8-54                  |                                 |                                  |
| DEXTRAN 70, 6%<br>6GM/100ML IN<br>SODIUM CHLORIDE 0.9%<br>0.9GM/100ML   | MACRODEX <sup>R</sup><br>(INJECTABLE; INJECTION)     | PHARMACIA LABS        | 6-826<br>6-8-54                  |                                 |                                  |
| DEXTRAN 40, 10%<br>10GM/100ML IN<br>DEXTROSE 5%<br>5GM/100ML            | GENTRAN <sup>R</sup> 40<br>(INJECTABLE; INJECTION)   | TRAVENOL LABS         | 16-628<br>11-4-68                |                                 |                                  |
| DEXTRAN 40, 10%<br>10GM/100ML IN<br>SODIUM CHLORIDE 0.9%<br>0.9GM/100ML | GENTRAN <sup>R</sup> 40<br>(INJECTABLE; INJECTION)   | TRAVENOL LABS         | 16-628<br>11-4-68                |                                 |                                  |
| DEXTRAN 40, 10%<br>10GM/100ML<br>DEXTROSE 5%<br>5GM/100ML               | GENTRAN <sup>R</sup> 40<br>(INJECTABLE; INJECTION)   | TRAVENOL LABS         | 84-619<br>2-22-85                |                                 |                                  |
| DEXTRAN 40, 10%<br>10GM/100ML IN<br>SODIUM CHLORIDE 0.9%<br>0.9GM/100ML | GENTRAN <sup>R</sup> 40<br>(INJECTABLE; INJECTION)   | TRAVENOL LABS         | 84-620<br>2-22-85                |                                 |                                  |
| DEXTRAN 75, 6%<br>6GM/100ML IN<br>SODIUM CHLORIDE 0.9%<br>0.9GM/100ML   | GENTRAN <sup>R</sup> 75<br>(INJECTABLE; INJECTION)   | TRAVENOL LABS         | 16-607<br>1-26-70                |                                 |                                  |

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| <u>ACTIVE INGREDIENT(S)</u><br><u>STRENGTH(S)</u>   | <u>TRADE NAME</u><br><u>(DOSAGE FORM; ROUTE)</u>                                      | <u>APPLICANT NAME</u> | <u>NDA NO.</u><br><u>APPROVAL DATE</u> | <u>PATENT NO.</u><br><u>EXP. DATE</u> | <u>EXCLUSIVITY</u><br><u>EXP. DATE</u> |
|---|---|-----------------------|--|---------------------------------------|--|
| DEXTRAN 75, 6%<br>INVERTED SUGAR 10%<br>6GM/100ML; 10GM/100ML<br>IN SODIUM CHLORIDE 0.9%<br>0.9GM/100ML | 6% GENTRAN <sup>R</sup> 75 AND<br>10% TRAVERT <sup>R</sup><br>(INJECTABLE; INJECTION) | TRAVENOL LABS         | 8-788<br>2-9-53                        |                                       |  |
| HETASTARCH, 6%<br>6GM/100ML IN<br>SODIUM CHLORIDE 0.9%<br>0.9GM/100ML                                   | HESPAR <sup>N</sup><br>(INJECTABLE; INJECTION)  | AM CRITICAL CARE      | 16-889<br>7-17-72                      | 3523938<br>8-11-87                    |  |
| PROPIOLACTONE 99%<br>99GM/100ML   | BETAPRONE<br>(SOLUTION; CHEMICAL<br>STERILIZING AGENT)                                | ONEAL JONES&FELDMAN   | 11-657<br>9-11-59                      |                                       |  |
| UROKINASE<br>5000 IU/VIAL   | ABBOKINASE OPEN-CATHETER<br>(INJECTABLE; INJECTION)                                   | ABBOTT LABORATORIES   | 76-1021<br>12-15-83                    |                                       | NS<br>09-24-86                         |
| UROKINASE<br>250,000 IU/VIAL  | ABBOKINASE<br>(INJECTABLE; INJECTION)   | ABBOTT LABORATORIES   | 76-1021<br>7-31-78                     |                                       | I-29<br>09-24-86                       |
| UROKINASE<br>250,000 IU/VIAL  | BREOKINASE<br>(INJECTABLE; INJECTION)   | STERLING DRUG         | 17-873<br>8-28-79                      |                                       |  |

**TABLE IV. NDA'S APPROVED FROM 1-1-82 TO 2-28-85 AND NDA'S WITH APPROPRIATE PATENT AND EXCLUSIVITY INFORMATION**

| <u>ACTIVE INGREDIENT(S)</u><br><u>STRENGTH(S)</u>               | <u>TRADE NAME</u><br><u>(DOSAGE FORM; ROUTE)</u>                  | <u>APPLICANT NAME</u> | <u>NDA NO.</u><br><u>APPROVAL DATE</u> | <u>PATENT NO.</u><br><u>EXP. DATE</u>      | <u>EXCLUSIVITY</u><br><u>EXP. DATE</u> |
|---|---|-----------------------|--|--|--|
| ACEBUTOLOL HYDROCHLORIDE<br>EQ 200MG BASE                       | SECTRAL<br>(CAPSULE; ORAL)  | IVES LABS/AMHO        | 18-917<br>12-28-84                     | 3726919<br>04-10-90<br>3857952<br>12-31-91 | NCE<br>12-28-89                        |
| ACEBUTOLOL HYDROCHLORIDE<br>EQ 300MG BASE                       | SECTRAL<br>(CAPSULE; ORAL)  | IVES LABS/AMHO        | 18-917<br>12-28-84                     | 3726919<br>04-10-90<br>3857952<br>12-31-91 | NCE<br>12-28-89                        |
| ACEBUTOLOL HYDROCHLORIDE<br>EQ 400MG BASE                       | SECTRAL<br>(CAPSULE; ORAL)  | IVES LABS/AMHO        | 18-917<br>12-28-84                     | 3726919<br>04-10-90<br>3857952<br>12-31-91 | NCE<br>12-28-89                        |
| ACETAMINOPHEN; PENTAZOCINE HYDROCHLORIDE<br>625MG; EQ 25MG BASE | TALACEN<br>(TABLET; ORAL)   | STERLING DRUG         | 18-458<br>09-23-82                     | 4105659<br>08-08-95                        | NC<br>09-24-86                         |
| ACETIC ACID, GLACIAL<br>250MG/100ML                             | ACETIC ACID 0.25%<br>IN PLASTIC CONTAINER<br>(SOLUTION; URETHRAL) | TRAVENOL LABS         | 18-523<br>02-19-82                     |  |  |
| ACETOHYDROXAMIC ACID<br>250MG                                   | LITHOSTAT<br>(TABLET; ORAL)                                       | URO-RESEARCH          | 18-749<br>05-31-83                     |  | NCE<br>05-31-93                        |
| ACYCLOVIR<br>5%   | ZOVIRAX<br>(OINTMENT; TOPICAL)                                    | BURROUGHS WELLCOME    | 18-604<br>03-29-82                     | 4199574<br>04-22-97                        | NCE<br>03-29-92                        |
| ACYCLOVIR<br>200MG  | ZOVIRAX<br>(CAPSULE; ORAL)  | BURROUGHS WELLCOME    | 18-828<br>01-25-85                     | 4199574<br>04-22-97                        | NCE<br>03-29-92                        |
| ACYCLOVIR SODIUM<br>EQ 500MG BASE/VIAL                          | ZOVIRAX<br>(INJECTABLE; INJECTION)                                | BURROUGHS WELLCOME    | 18-603<br>10-22-82                     | 4199574<br>04-22-97                        | NCE<br>03-29-92                        |
| ALBUTEROL<br>0.09MG/INH   | PROVENTIL<br>(AEROSOL; INHALATION)                                | SCHERING              | 17-559<br>05-01-81                     | 3644353<br>02-22-89<br>3705233<br>12-05-89 | 1-22<br>09-24-86                       |
| ALBUTEROL<br>0.09MG/INH   | VENTOLIN<br>(AEROSOL; INHALATION)                                 | GLAXO                 | 18-473<br>05-01-81                     | 3644353<br>02-22-89<br>3705233<br>12-05-89 |  |

TABLE IV. NDA'S APPROVED FROM 1-1-82 TO 2-28-85 AND NDA'S WITH APPROPRIATE PATENT AND EXCLUSIVITY INFORMATION

| <u>ACTIVE INGREDIENT(S)<br/>STRENGTH(S)</u> | <u>TRADE NAME<br/>(DOSAGE FORM; ROUTE)</u> | <u>APPLICANT NAME</u> | <u>NDA NO.<br/>APPROVAL DATE</u> | <u>PATENT NO.<br/>EXP. DATE</u>            | <u>EXCLUSIVITY<br/>EXP. DATE</u> |
|---|--|-----------------------|----------------------------------|--|----------------------------------|
| ALBUTEROL SULFATE<br>EQ 2MG BASE            | PROVENTIL<br>(TABLET; ORAL)                | SCHERING              | 17-853<br>05-07-82               | 3644353<br>02-22-89<br>3705233<br>12-05-89 | NE<br>09-24-86                   |
| ALBUTEROL SULFATE<br>EQ 4MG BASE            | PROVENTIL<br>(TABLET; ORAL)                | SCHERING              | 17-853<br>05-07-82               | 3644353<br>02-22-89<br>3705233<br>12-05-89 | NE<br>09-24-86                   |
| ALCLOMETASONE DIPROPIONATE<br>0.05%         | VADERM<br>(OINTMENT; TOPICAL)              | SCHERING              | 18-702<br>12-14-82               | 4124707<br>11-07-95                        | NCE<br>12-14-92                  |
| ALCLOMETASONE DIPROPIONATE<br>0.05%         | VADERM<br>(CREAM; TOPICAL)                 | SCHERING              | 18-707<br>12-14-82               | 4124707<br>11-07-95                        | NCE<br>12-14-92                  |
| ALLOPURINOL<br>100MG                        | ALLOPURINOL<br>(TABLET; ORAL)              | BOLAR PHARMACEUTICAL  | 18-241<br>11-16-84               |  |                                  |
| ALLOPURINOL<br>300MG                        | ALLOPURINOL<br>(TABLET; ORAL)              | BOLAR PHARMACEUTICAL  | 18-241<br>11-16-84               |  |                                  |
| ALLOPURINOL<br>100MG                        | ALLOPURINOL<br>(TABLET; ORAL)              | CHELSEA LABORATORIES  | 18-785<br>09-28-84               |  |                                  |
| ALLOPURINOL<br>300MG                        | ALLOPURINOL<br>(TABLET; ORAL)              | CHELSEA LABORATORIES  | 18-785<br>09-28-84               |  |                                  |
| ALLOPURINOL<br>100MG                        | ALLOPURINOL<br>(TABLET; ORAL)              | DANBURY PHARMACAL     | 18-832<br>09-28-84               |  |                                  |
| ALLOPURINOL<br>300MG                        | ALLOPURINOL<br>(TABLET; ORAL)              | DANBURY PHARMACAL     | 18-877<br>09-28-84               |  |                                  |
| ALLOPURINOL<br>100MG                        | ZYLOPRIM<br>(TABLET; ORAL)                 | BURROUGHS WELLCOME    | 16-084<br>08-19-66               | 3624205<br>11-30-88                        |                                  |
| ALLOPURINOL<br>300MG                        | ZYLOPRIM<br>(TABLET; ORAL)                 | BURROUGHS WELLCOME    | 16-084<br>01-14-74               | 3624205<br>11-30-88                        |                                  |
| ALLOPURINOL<br>100MG                        | LOPURIN<br>(TABLET; ORAL)                  | BOOTS PHARMACEUTICAL  | 18-297<br>06-10-80               | 3624205<br>11-30-88                        |                                  |

**TABLE IV. NDA'S APPROVED FROM 1-1-82 TO 2-28-85 AND NDA'S WITH APPROPRIATE PATENT AND EXCLUSIVITY INFORMATION**

| <u>ACTIVE INGREDIENT(S)</u><br><u>STRENGTH(S)</u>            | <u>TRADE NAME</u><br><u>(DOSAGE FORM; ROUTE)</u>    | <u>APPLICANT NAME</u> | <u>NDA NO.</u><br><u>APPROVAL DATE</u> | <u>PATENT NO.</u><br><u>EXP. DATE</u>      | <u>EXCLUSIVITY</u><br><u>EXP. DATE</u> |
|--|---|-----------------------|--|--|--|
| ALLOPURINOL<br>300MG   | LOPURIN<br>(TABLET; ORAL)                           | BOOTS PHARMACEUTICAL  | 18-297<br>06-10-80                     | 3624205<br>11-30-88                        |  |
| ALPRAZOLAM<br>0.25MG   | XANAX<br>(TABLET; ORAL)                             | UPJOHN                | 18-276<br>10-16-81                     | 3987052<br>10-19-93<br>3980789<br>09-14-93 |  |
| ALPRAZOLAM<br>0.5MG  | XANAX<br>(TABLET; ORAL)                             | UPJOHN                | 18-276<br>10-16-81                     | 3987052<br>10-19-93<br>3980789             |  |
| ALPRAZOLAM<br>1MG  | XANAX<br>(TABLET; ORAL)                             | UPJOHN                | 18-276<br>10-16-81                     | 3987052<br>10-19-93<br>3980789<br>09-14-93 |  |
| AMCINONIDE<br>0.1%   | CYCLOCORT<br>(CREAM; TOPICAL)                       | LEDERLE LABS/AM CYAN  | 18-116<br>10-18-71                     | 4158055<br>06-12-96                        |  |
| AMCINONIDE<br>0.1%   | CYCLOCORT<br>(OINTMENT; TOPICAL)                    | LEDERLE LABS/AM CYAN  | 18-498<br>11-13-81                     | 4158055<br>06-12-96                        |  |
| AMILORIDE HYDROCHLORIDE;<br>HYDROCHLOROTHIAZIDE<br>5MG; 50MG | MODURETIC 5/50<br>(TABLET; ORAL)                    | MS&D/MERCK            | 18-201<br>10-05-81                     | 3781430<br>12-25-90                        |  |
| AMINO ACIDS<br>6.9%  | FREAMINE HBC 6.9%<br>(INJECTABLE; INJECTION)        | AM MCGAW/AM HOSP      | 16-822<br>05-17-83                     |  | NS<br>09-24-86                         |
| AMINO ACIDS<br>6.5%  | RENAMIN W/O ELECTROLYTES<br>(INJECTABLE; INJECTION) | TRAVENOL LABS         | 17-493<br>10-15-82                     |  | NS<br>09-24-86                         |
| AMINO ACIDS<br>8.5%  | NOVAMINE 8.5%<br>(INJECTABLE; INJECTION)            | CUTTER LABS/MILES     | 17-957<br>08-09-82                     |  |  |
| AMINO ACIDS<br>11.4%   | NOVAMINE 11.4%<br>(INJECTABLE; INJECTION)           | CUTTER LABS/MILES     | 17-957<br>08-09-82                     |  |  |
| AMINO ACIDS<br>8%  | HEPATAMINE 8%<br>(INJECTABLE; INJECTION)            | AM MCGAW/AM HOSP      | 18-676<br>08-03-82                     | 3950529<br>04-13-93                        | NS<br>09-24-86                         |

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| <u>ACTIVE INGREDIENT(S)<br/>STRENGTH(S)</u> | <u>TRADE NAME<br/>(DOSAGE FORM; ROUTE)</u>   | <u>APPLICANT NAME</u> | <u>NDA NO.<br/>APPROVAL DATE</u> | <u>PATENT NO.<br/>EXP. DATE</u> | <u>EXCLUSIVITY<br/>EXP. DATE</u> |
|---|--|-----------------------|----------------------------------|---------------------------------|----------------------------------|
| AMINO ACIDS<br>4%                           | BRANCHAMIN 4%<br>(INJECTABLE; INJECTION)   | TRAVENOL LABS         | 18-678<br>09-28-84               | 4438144<br>03-20-01             | NS<br>09-24-86                   |
| AMINO ACIDS<br>4%                           | BRANCHAMIN 4%<br>IN PLASTIC CONTAINER<br>(INJECTABLE; INJECTION)                     | TRAVENOL LABS         | 18-684<br>09-28-84               | 4438144<br>03-20-01             | NS<br>09-24-86                   |
| AMINO ACIDS<br>6.5%                         | NEOPHAM 6.5%<br>(INJECTABLE; INJECTION)  | CUTTER-VITRUM         | 18-792<br>01-17-84               |                                 | NS<br>09-24-86                   |
| AMINO ACIDS<br>3.5%                         | AMINOSYN 3.5%<br>IN PLASTIC CONTAINER<br>(INJECTABLE; INJECTION)                     | ABBOTT LABORATORIES   | 18-804<br>05-15-84               |                                 | NS<br>09-24-86                   |
| AMINO ACIDS<br>3.5%                         | AMINOSYN 3.5%<br>IN PLASTIC CONTAINER<br>(INJECTABLE; INJECTION)                     | ABBOTT LABORATORIES   | 18-875<br>08-08-84               |                                 | NS<br>09-24-86                   |
| AMINO ACIDS<br>5.2%                         | AMINESS 5.2% ESSENTIAL<br>AMINO ACIDS W/ HISTADINE<br>(INJECTABLE; INJECTION)        | CUTTER-VITRUM         | 18-901<br>04-06-84               |                                 |                                  |
| AMINO ACIDS<br>5.5%                         | TRAVASOL 5.5%<br>W/O ELECTROLYTES<br>IN PLASTIC CONTAINER<br>(INJECTABLE; INJECTION) | TRAVENOL LABS         | 18-931<br>08-23-84               |                                 | NS<br>09-24-86                   |
| AMINO ACIDS<br>8.5%                         | TRAVASOL 8.5%<br>W/O ELECTROLYTES<br>IN PLASTIC CONTAINER<br>(INJECTABLE; INJECTION) | TRAVENOL LABS         | 18-931<br>08-23-84               |                                 |                                  |
| AMINO ACIDS<br>10%                          | TRAVASOL 10%<br>W/O ELECTROLYTES<br>IN PLASTIC CONTAINER<br>(INJECTABLE; INJECTION)  | TRAVENOL LABS         | 18-931<br>08-23-84               |                                 |                                  |
| AMINO ACIDS<br>6%                           | TROPHAMINE 6%<br>(INJECTABLE; INJECTION)   | AM MCGAW/AM HOSP      | 19-018<br>07-20-84               |                                 | NS<br>09-24-86                   |

**TABLE IV. NDA'S APPROVED FROM 1-1-82 TO 2-28-85 AND NDA'S WITH APPROPRIATE PATENT AND EXCLUSIVITY INFORMATION**

| <u>ACTIVE INGREDIENT(S)<br/>STRENGTH(S)</u>   | <u>TRADE NAME<br/>(DOSAGE FORM; ROUTE)</u>   | <u>APPLICANT NAME</u> | <u>NDA NO.<br/>APPROVAL DATE</u> | <u>PATENT NO.<br/>EXP. DATE</u> | <u>EXCLUSIVITY<br/>EXP. DATE</u> |
|---|--|-----------------------|----------------------------------|---------------------------------|----------------------------------|
| AMINO ACIDS; CALCIUM ACETATE;<br>GLYCERIN; MAGNESIUM ACETATE;<br>PHOSPHORIC ACID; POTASSIUM CHLORIDE;<br>SODIUM ACETATE; SODIUM CHLORIDE<br>3%; 26MG/100ML; 3GM/100ML;<br>54MG/100ML; 41MG/100ML;<br>149MG/100ML; 204MG/100ML;<br>117MG/100ML | PERIPHERAMINE<br>(INJECTABLE; INJECTION)   | AM MCGAM/AM HOSP      | 18-582<br>05-08-82               |                                 | NC<br>09-24-86                   |
| AMINO ACIDS; DEXTROSE<br>3.5%; 5%   | AMINOSYN 3.5%<br>W/ DEXTROSE 5%<br>IN PLASTIC CONTAINER<br>(INJECTABLE; INJECTION)   | ABBOTT LABORATORIES   | 19-120<br>10-11-84               |                                 |                                  |
| AMINO ACIDS; DEXTROSE<br>3.5%; 25%  | AMINOSYN 3.5%<br>W/ DEXTROSE 25%<br>IN PLASTIC CONTAINER<br>(INJECTABLE; INJECTION)  | ABBOTT LABORATORIES   | 19-118<br>10-11-84               |                                 |                                  |
| AMINO ACIDS; DEXTROSE<br>4.25%; 25%   | AMINOSYN 4.25%<br>W/ DEXTROSE 25%<br>IN PLASTIC CONTAINER<br>(INJECTABLE; INJECTION) | ABBOTT LABORATORIES   | 19-119<br>10-11-84               |                                 |                                  |
| AMINO ACIDS; MAGNESIUM ACETATE;<br>PHOSPHORIC ACID; POTASSIUM ACETATE;<br>SODIUM CHLORIDE<br>3.5%; 21MG/100ML; 40MG/100ML;<br>128MG/100ML; 234MG/100ML  | AMINOSYN 3.5% M<br>IN PLASTIC CONTAINER<br>(INJECTABLE; INJECTION)                   | ABBOTT LABORATORIES   | 18-804<br>05-15-84               |                                 | NC<br>09-24-86                   |
| AMINO ACIDS; MAGNESIUM ACETATE;<br>PHOSPHORIC ACID; POTASSIUM ACETATE;<br>SODIUM CHLORIDE<br>3.5%; 21MG/100ML; 40MG/100ML;<br>128MG/100ML; 234MG/100ML  | AMINOSYN 3.5% M<br>IN PLASTIC CONTAINER<br>(INJECTABLE; INJECTION)                   | ABBOTT LABORATORIES   | 18-875<br>08-08-84               |                                 | NC<br>09-24-86                   |
| AMINOACETIC ACID<br>1.5GM/100ML   | AMINOACETIC ACID 1.5%<br>IN PLASTIC CONTAINER<br>(SOLUTION; IRRIGATION)              | TRAVENOL LABS         | 18-522<br>02-19-82               |                                 |                                  |

**TABLE IV. NDA'S APPROVED FROM 1-1-82 TO 2-28-85 AND NDA'S WITH APPROPRIATE PATENT AND EXCLUSIVITY INFORMATION**

| <u>ACTIVE INGREDIENT(S)<br/>STRENGTH(S)</u>                | <u>TRADE NAME<br/>(DOSAGE FORM; ROUTE)</u>   | <u>APPLICANT NAME</u> | <u>NDA NO.<br/>APPROVAL DATE</u> | <u>PATENT NO.<br/>EXP. DATE</u>            | <u>EXCLUSIVITY<br/>EXP. DATE</u> |
|--|--|-----------------------|----------------------------------|--|----------------------------------|
| AMINOCAPROIC ACID<br>250MG/ML                              | AMINOCAPROIC ACID<br>(INJECTABLE; INJECTION)   | ELKINS-SINN/AHROBINS  | 18-590<br>10-29-82               |  |                                  |
| AMINOGLUTETHIMIDE<br>250MG                                 | CYTADREN<br>(TABLET; ORAL)   | CIBA/CIBA-GEIGY       | 18-202<br>10-29-80               | 3595960<br>07-27-88<br>3944671<br>03-16-93 |                                  |
| AMINOPHYLLINE<br>300MG/5ML                                 | SOMOPHYLLIN<br>(ENEMA; RECTAL)   | FISONS                | 18-232<br>04-02-82               |  | NR<br>09-24-86                   |
| AMINOPHYLLINE; SODIUM CHLORIDE<br>100MG/100ML; 450MG/100ML | AMINOPHYLLINE W/<br>SODIUM CHLORIDE 0.45%<br>IN PLASTIC CONTAINER<br>(INJECTABLE; INJECTION) | ABBOTT LABORATORIES   | 18-924<br>12-12-84               |  |                                  |
| AMINOPHYLLINE; SODIUM CHLORIDE<br>200MG/100ML; 450MG/100ML | AMINOPHYLLINE W/<br>SODIUM CHLORIDE 0.45%<br>IN PLASTIC CONTAINER<br>(INJECTABLE; INJECTION) | ABBOTT LABORATORIES   | 18-924<br>12-12-84               |  |                                  |
| AMINOPHYLLINE; SODIUM CHLORIDE<br>400MG/100ML; 450MG/100ML | AMINOPHYLLINE W/<br>SODIUM CHLORIDE 0.45%<br>IN PLASTIC CONTAINER<br>(INJECTABLE; INJECTION) | ABBOTT LABORATORIES   | 18-924<br>12-12-84               |  |                                  |
| AMINOPHYLLINE; SODIUM CHLORIDE<br>500MG/100ML; 450MG/100ML | AMINOPHYLLINE W/<br>SODIUM CHLORIDE 0.45%<br>IN PLASTIC CONTAINER<br>(INJECTABLE; INJECTION) | ABBOTT LABORATORIES   | 18-924<br>12-12-84               |  |                                  |
| AMITRIPTYLINE HYDROCHLORIDE<br>10MG                        | ELAVIL<br>(TABLET; ORAL)   | MS&D/MERCK            | 12-703<br>04-07-61               | 3384663<br>05-21-85<br>3428735<br>02-18-86 |                                  |
| AMITRIPTYLINE HYDROCHLORIDE<br>25MG                        | ELAVIL<br>(TABLET; ORAL)   | MS&D/MERCK            | 12-703<br>07-05-74               | 3384663<br>05-21-85<br>3428735<br>02-18-86 |                                  |

TABLE IV. NDA'S APPROVED FROM 1-1-82 TO 2-28-85 AND NDA'S WITH APPROPRIATE PATENT AND EXCLUSIVITY INFORMATION

| <u>ACTIVE INGREDIENT(S)<br/>STRENGTH(S)</u>                     | <u>TRADE NAME<br/>(DOSAGE FORM; ROUTE)</u> | <u>APPLICANT NAME</u> | <u>NDA NO.<br/>APPROVAL DATE</u> | <u>PATENT NO.<br/>EXP. DATE</u>            | <u>EXCLUSIVITY<br/>EXP. DATE</u> |
|---|--|-----------------------|----------------------------------|--|----------------------------------|
| AMITRIPTYLINE HYDROCHLORIDE<br>50MG                             | ELAVIL<br>(TABLET; ORAL)                   | MS&D/MERCK            | 12-703<br>04-07-61               | 3384663<br>05-21-85<br>3428735<br>02-18-86 |                                  |
| AMITRIPTYLINE HYDROCHLORIDE<br>75MG                             | ELAVIL<br>(TABLET; ORAL)                   | MS&D/MERCK            | 12-703<br>10-28-76               | 3384663<br>05-21-85<br>3428735<br>02-18-86 |                                  |
| AMITRIPTYLINE HYDROCHLORIDE<br>100MG                            | ELAVIL<br>(TABLET; ORAL)                   | MS&D/MERCK            | 12-703<br>10-28-76               | 3384663<br>05-21-85<br>3428735<br>02-18-86 |                                  |
| AMITRIPTYLINE HYDROCHLORIDE<br>150MG                            | ELAVIL<br>(TABLET; ORAL)                   | MS&D/MERCK            | 12-703<br>09-17-76               | 3384663<br>05-21-85<br>3428735<br>02-18-86 |                                  |
| AMITRIPTYLINE HYDROCHLORIDE<br>10MG/ML                          | ELAVIL<br>(INJECTABLE; INJECTION)          | MS&D/MERCK            | 12-704<br>04-11-61               | 3384663<br>05-21-85<br>3428735<br>02-18-86 |                                  |
| AMITRIPTYLINE HYDROCHLORIDE;<br>CHLORDIAZEPOXIDE<br>12.5MG; 5MG | LIMBITROL<br>(TABLET; ORAL)                | HOFFMANN-LA ROCHE     | 16-949<br>12-23-77               | 3384663<br>05-21-85<br>4316897<br>02-23-99 |                                  |
| AMITRIPTYLINE HYDROCHLORIDE;<br>CHLORDIAZEPOXIDE<br>25MG; 10MG  | LIMBITROL<br>(TABLET; ORAL)                | HOFFMANN-LA ROCHE     | 16-949<br>12-23-77               | 3384663<br>05-21-85<br>4316897<br>02-23-99 |                                  |
| AMITRIPTYLINE HYDROCHLORIDE;<br>PERPHENAZINE<br>10MG; 4MG       | ETRAFON A<br>(TABLET; ORAL)                | SCHERING              | 14-713<br>12-30-65               | 3384663<br>05-21-85                        |                                  |
| AMITRIPTYLINE HYDROCHLORIDE;<br>PERPHENAZINE<br>25MG; 2MG       | ETRAFON 2-25<br>(TABLET; ORAL)             | SCHERING              | 14-713<br>12-30-65               | 3384663<br>05-21-85                        |                                  |

**TABLE IV. NDA'S APPROVED FROM 1-1-82 TO 2-28-85 AND NDA'S WITH APPROPRIATE PATENT AND EXCLUSIVITY INFORMATION**

| <u>ACTIVE INGREDIENT(S)</u><br><u>STRENGTH(S)</u>         | <u>TRADE NAME</u><br><u>(DOSAGE FORM; ROUTE)</u> | <u>APPLICANT NAME</u> | <u>NDA NO.</u><br><u>APPROVAL DATE</u> | <u>PATENT NO.</u><br><u>EXP. DATE</u>                             | <u>EXCLUSIVITY</u><br><u>EXP. DATE</u> |
|---|--|-----------------------|--|---|--|
| AMITRIPTYLINE HYDROCHLORIDE;<br>PERPHENAZINE<br>25MG; 4MG | ETRAFON-FORTE<br>(TABLET; ORAL)                  | SCHERING              | 14-713<br>12-30-65                     | 3384663<br>05-21-85   |  |
| AMITRIPTYLINE HYDROCHLORIDE;<br>PERPHENAZINE<br>10MG; 2MG | ETRAFON 2-10<br>(TABLET; ORAL)                   | SCHERING              | 14-713<br>12-30-65                     | 3384663<br>05-21-85   |  |
| AMITRIPTYLINE HYDROCHLORIDE;<br>PERPHENAZINE<br>10MG; 4MG | TRIAVIL 4-10<br>(TABLET; ORAL)                   | MS&D/MERCK            | 14-715<br>12-30-65                     | 3384663<br>05-21-85<br>3428735<br>02-18-86                        |  |
| AMITRIPTYLINE HYDROCHLORIDE;<br>PERPHENAZINE<br>25MG; 2MG | TRIAVIL 2-25<br>(TABLET; ORAL)                   | MS&D/MERCK            | 14-715<br>08-23-65                     | 3384663<br>05-21-85<br>3428735<br>02-18-86                        |  |
| AMITRIPTYLINE HYDROCHLORIDE;<br>PERPHENAZINE<br>10MG; 2MG | TRIAVIL 2-10<br>(TABLET; ORAL)                   | MS&D/MERCK            | 14-715<br>04-04-67                     | 3384663<br>05-21-85<br>3428735<br>02-18-86                        |  |
| AMITRIPTYLINE HYDROCHLORIDE;<br>PERPHENAZINE<br>25MG; 4MG | TRIAVIL 4-25<br>(TABLET; ORAL)                   | MS&D/MERCK            | 14-715<br>08-25-65                     | 3384663<br>05-21-85<br>3428735<br>02-18-86                        |  |
| AMITRIPTYLINE HYDROCHLORIDE;<br>PERPHENAZINE<br>50MG; 4MG | TRIAVIL 4-50<br>(TABLET; ORAL)                   | MS&D/MERCK            | 14-715<br>03-15-78                     | 3384663<br>05-21-85<br>3428735<br>02-18-86                        |  |
| AMOXAPINE<br>25MG   | ASENDIN<br>(TABLET; ORAL)                        | LEDERLE LABS/AM CYAN  | 18-021<br>09-22-80                     | 3546226<br>12-08-87<br>3663696<br>05-16-89<br>3681357<br>08-01-89 |  |

**TABLE IV. NDA'S APPROVED FROM 1-1-82 TO 2-28-85 AND NDA'S WITH APPROPRIATE PATENT AND EXCLUSIVITY INFORMATION**

| <u>ACTIVE INGREDIENT(S)<br/>STRENGTH(S)</u>                             | <u>TRADE NAME<br/>(DOSAGE FORM; ROUTE)</u> | <u>APPLICANT NAME</u> | <u>NDA NO.<br/>APPROVAL DATE</u> | <u>PATENT NO.<br/>EXP. DATE</u>                                   | <u>EXCLUSIVITY<br/>EXP. DATE</u> |
|---|--|-----------------------|----------------------------------|---|----------------------------------|
| AMOXAPINE<br>50MG   | ASENDIN<br>(TABLET; ORAL)                  | LEDERLE LABS/AM CYAN  | 18-021<br>09-22-80               | 3546226<br>12-08-87<br>3663696<br>05-16-89<br>3681357<br>08-01-89 |                                  |
| AMOXAPINE<br>100MG  | ASENDIN<br>(TABLET; ORAL)                  | LEDERLE LABS/AM CYAN  | 18-021<br>09-22-80               | 3546226<br>12-08-87<br>3663696<br>05-16-89<br>3681357<br>08-01-89 |                                  |
| AMOXAPINE<br>150MG  | ASENDIN<br>(TABLET; ORAL)                  | LEDERLE LABS/AM CYAN  | 18-021<br>09-22-80               | 3546226<br>12-08-87<br>3663696<br>05-16-89<br>3681357<br>08-01-89 |                                  |
| AMRINONE LACTATE<br>EQ 5MG BASE/ML                                      | INOCOR<br>(INJECTABLE; INJECTION)          | WINTHROP LABS/STERL   | 18-700<br>07-31-84               | 4072746<br>02-07-95   | NCE<br>07-31-94                  |
| ASPIRIN; CAFFEINE;<br>DIHYDROCODEINE BITARTRATE<br>356.4MG; 30MG; 16MG  | SYNALGOS-DC<br>(CAPSULE; ORAL)             | IVES LABS/AMHO        | 11-483<br>09-06-83               |   |                                  |
| ASPIRIN; CAFFEINE;<br>ORPHENADRINE CITRATE<br>385MG; 30MG; 25MG         | NORGESIC<br>(TABLET; ORAL)                 | RIKER LABS/3M         | 13-416<br>10-27-82               |   |                                  |
| ASPIRIN; CAFFEINE;<br>ORPHENADRINE CITRATE<br>770MG; 60MG; 50MG         | NORGESIC FORTE<br>(TABLET; ORAL)           | RIKER LABS/3M         | 13-416<br>10-27-82               |   |                                  |
| ASPIRIN; CAFFEINE;<br>PROPOXYPHENE HYDROCHLORIDE<br>389MG; 32.4MG; 32MG | DARVON COMPOUND<br>(CAPSULE; ORAL)         | ELI LILLY INDSTRS/PR  | 10-996<br>03-08-83               |   |                                  |

**TABLE IV. NDA'S APPROVED FROM 1-1-82 TO 2-28-85 AND NDA'S WITH APPROPRIATE PATENT AND EXCLUSIVITY INFORMATION**

| <u>ACTIVE INGREDIENT(S)<br/>STRENGTH(S)</u>                             | <u>TRADE NAME<br/>(DOSAGE FORM; ROUTE)</u> | <u>APPLICANT NAME</u> | <u>NDA NO.<br/>APPROVAL DATE</u> | <u>PATENT NO.<br/>EXP. DATE</u>                                   | <u>EXCLUSIVITY<br/>EXP. DATE</u> |
|---|--|-----------------------|----------------------------------|---|----------------------------------|
| ASPIRIN; CAFFEINE;<br>PROPOXYPHENE HYDROCHLORIDE<br>389MG; 32.4MG; 65MG | DARVON COMPOUND-65<br>(CAPSULE; ORAL)      | ELI LILLY INDSTRS/PR  | 10-996<br>03-08-83               |   |                                  |
| ASPIRIN; CARISOPRODOL<br>325MG; 200MG                                   | SOMA COMPOUND<br>(TABLET; ORAL)            | WALLACE PHARMS/C-W    | 12-365<br>07-11-83               |   |                                  |
| ASPIRIN; CARISOPRODOL;<br>CODEINE PHOSPHATE<br>325MG; 200MG; 16MG       | SOMA COMPOUND W/ CODEINE<br>(TABLET; ORAL) | WALLACE PHARMS/C-W    | 12-366<br>07-11-83               |   |                                  |
| ASPIRIN; MEPROBAMATE<br>325MG; 200MG                                    | EQUAGESIC<br>(TABLET; ORAL)                | WYETH LABS/AMHO       | 11-702<br>12-29-83               |   |                                  |
| ASPIRIN; PENTAZOCINE HYDROCHLORIDE<br>325MG; EQ 12.5MG BASE             | TALWIN COMPOUND<br>(TABLET; ORAL)          | WINTHROP LABS/STERL   | 16-891<br>11-12-75               | 4105659<br>08-08-95   |                                  |
| ATENOLOL<br>50MG  | TENORMIN<br>(TABLET; ORAL)                 | STUART PHARMS/ICI AM  | 18-240<br>08-19-81               | 3663607<br>05-16-89<br>3934032<br>01-20-93<br>3836671<br>09-17-91 |                                  |
| ATENOLOL<br>100MG   | TENORMIN<br>(TABLET; ORAL)                 | STUART PHARMS/ICI AM  | 18-240<br>08-19-81               | 3663607<br>05-16-89<br>3934032<br>01-20-93<br>3836671<br>09-17-91 |                                  |
| ATENOLOL; CHLORTHALIDONE<br>100MG; 25MG                                 | TENORETIC 100<br>(TABLET; ORAL)            | STUART PHARMS/ICI AM  | 18-760<br>06-08-84               | 3663607<br>05-16-89<br>3934032<br>01-20-93<br>3836671<br>09-17-91 | NC<br>09-24-86                   |

**TABLE IV. NDA'S APPROVED FROM 1-1-82 TO 2-28-85 AND NDA'S WITH APPROPRIATE PATENT AND EXCLUSIVITY INFORMATION**

| <u>ACTIVE INGREDIENT(S)<br/>STRENGTH(S)</u>                 | <u>TRADE NAME<br/>(DOSAGE FORM; ROUTE)</u>        | <u>APPLICANT NAME</u> | <u>NDA NO.<br/>APPROVAL DATE</u> | <u>PATENT NO.<br/>EXP. DATE</u>                                   | <u>EXCLUSIVITY<br/>EXP. DATE</u> |
|---|---|-----------------------|----------------------------------|---|----------------------------------|
| ATENOLOL; CHLORTHALIDONE<br>50MG; 25MG                      | TENORETIC 50<br>(TABLET; ORAL)                    | STUART PHARMS/ICI AM  | 18-760<br>06-08-84               | 3663607<br>05-16-89<br>3934032<br>01-20-93<br>3836671<br>09-17-91 | NC<br>09-24-86                   |
| ATRACURIUM BESYLATE<br>10MG/ML                              | TRACRIUM<br>(INJECTABLE; INJECTION)               | BURROUGHS WELLCOME    | 18-831<br>11-23-83               | 4179507<br>12-18-96   | NCE<br>11-23-93                  |
| ATROPINE SULFATE; DIFENOXIN HYDROCHLORIDE<br>0.025MG; 0.5MG | MOTOFEN HALF-STRENGTH<br>(TABLET; ORAL)           | MCNEIL LABORATORIES   | 17-744<br>07-14-78               | 3646207<br>02-28-89   |                                  |
| ATROPINE SULFATE; DIFENOXIN HYDROCHLORIDE<br>0.025MG; 1MG   | MOTOFEN<br>(TABLET; ORAL)                         | MCNEIL LABORATORIES   | 17-744<br>07-14-78               | 3646207<br>02-28-89   |                                  |
| AZATADINE MALEATE<br>1MG                                    | OPTIMINE<br>(TABLET; ORAL)                        | SCHERING              | 17-601<br>03-29-77               | 3419565<br>12-31-85<br>3717647<br>02-20-90                        | NC<br>09-24-86                   |
| AZATADINE MALEATE;<br>PSEUDOEPHEDRINE SULFATE<br>1MG; 120MG | TRINALIN<br>(TABLET, CONTROLLED<br>RELEASE; ORAL) | SCHERING              | 18-506<br>03-23-82               | 3419565<br>12-31-85<br>3717647                                    | NC<br>09-24-86                   |
| BACLOFEN<br>10MG  | LIORESAL<br>(TABLET; ORAL)                        | GEIGY/CIBA-GEIGY      | 17-851<br>11-22-77               | 02-20-90<br>3471548<br>10-07-86                                   |                                  |
| BACLOFEN<br>20MG  | LIORESAL DS<br>(TABLET; ORAL)                     | GEIGY/CIBA-GEIGY      | 17-851<br>01-20-82               | 3471548<br>10-07-86   | NS<br>09-24-86                   |
| BENDROFLUMETHIAZIDE<br>2.5MG                                | NATURETIN-2.5<br>(TABLET; ORAL)                   | ER SQUIBB AND SONS    | 12-164<br>12-07-59               | 3392168<br>07-09-85   |                                  |
| BENDROFLUMETHIAZIDE<br>5MG                                  | NATURETIN-5<br>(TABLET; ORAL)                     | ER SQUIBB AND SONS    | 12-164<br>12-07-59               | 3392168<br>07-09-85   |                                  |
| BENDROFLUMETHIAZIDE<br>10MG                                 | NATURETIN-10<br>(TABLET; ORAL)                    | ER SQUIBB AND SONS    | 12-164<br>03-29-77               | 3392168<br>07-09-85   |                                  |

TABLE IV. NDA'S APPROVED FROM 1-1-82 TO 2-28-85 AND NDA'S WITH APPROPRIATE PATENT AND EXCLUSIVITY INFORMATION

| <u>ACTIVE INGREDIENT(S)<br/>STRENGTH(S)</u>  | <u>TRADE NAME<br/>(DOSAGE FORM; ROUTE)</u>     | <u>APPLICANT NAME</u> | <u>NDA NO.<br/>APPROVAL DATE</u> | <u>PATENT NO.<br/>EXP. DATE</u>            | <u>EXCLUSIVITY<br/>EXP. DATE</u> |
|--|--|-----------------------|----------------------------------|--|----------------------------------|
| BENDROFLUMETHIAZIDE; NADOLOL<br>5MG; 40MG  | CORZIDE<br>(TABLET; ORAL)                      | ER SQUIBB AND SONS    | 18-647<br>05-25-83               | 398201<br>09-21-93<br>3935267<br>01-27-93  | NC<br>09-24-86                   |
| BENDROFLUMETHIAZIDE; NADOLOL<br>5MG; 80MG  | CORZIDE<br>(TABLET; ORAL)                      | ER SQUIBB AND SONS    | 18-647<br>05-25-83               | 398201<br>09-21-93<br>3935267<br>01-27-93  | NC<br>09-24-86                   |
| BENTIROMIDE<br>500MG/7.5ML   | CHYMEX<br>(SOLUTION; ORAL)                     | ADRIA LABORATORIES    | 18-366<br>12-29-83               | 3801562<br>04-02-91<br>3745212<br>07-10-90 | NCE<br>12-29-93                  |
| BETAMETHASONE<br>0.6MG   | CELESTONE<br>(TABLET; ORAL)                    | SCHERING              | 12-657<br>04-17-61               | 3485854<br>12-23-86                        |                                  |
| BETAMETHASONE<br>0.6MG/5ML   | CELESTONE<br>(SYRUP; ORAL)                     | SCHERING              | 14-215<br>04-18-64               | 3485854<br>12-23-86                        |                                  |
| BETAMETHASONE<br>0.2%  | CELESTONE<br>(CREAM; TOPICAL)                  | SCHERING              | 14-762<br>04-10-64               | 3485854<br>12-23-86                        |                                  |
| BETAMETHASONE ACETATE;<br>BETAMETHASONE SODIUM PHOSPHATE<br>3MG/ML; EQ 3MG BASE/ML | CELESTONE SOLUSPAN<br>(INJECTABLE; INJECTION)  | SCHERING              | 14-602<br>03-03-65               | 3485854<br>12-23-86                        |                                  |
| BETAMETHASONE DIPROPIONATE<br>EQ 0.05% BASE  | DIPROLENE<br>(OINTMENT; TOPICAL)               | SCHERING              | 18-741<br>07-27-83               |  |                                  |
| BETAMETHASONE DIPROPIONATE<br>EQ 0.05% BASE  | BETAMETHASONE DIPROPIONATE<br>(CREAM; TOPICAL) | PHARMADERM/BYK-GLDN   | 19-136<br>06-26-84               |  |                                  |
| BETAMETHASONE DIPROPIONATE<br>EQ 0.05% BASE  | BETAMETHASONE DIPROPIONATE<br>(CREAM; TOPICAL) | E FOUGERA/BYK-GLDN    | 19-137<br>06-26-84               |  |                                  |
| BETAMETHASONE DIPROPIONATE<br>EQ 0.05% BASE  | ALPHATREX<br>(CREAM; TOPICAL)                  | SAVAGE LABS/BYK-GLDN  | 19-138<br>06-26-84               |  |                                  |

**TABLE IV. NDA'S APPROVED FROM 1-1-82 TO 2-28-85 AND NDA'S WITH APPROPRIATE PATENT AND EXCLUSIVITY INFORMATION**

| <u>ACTIVE INGREDIENT(S)</u><br><u>STRENGTH(S)</u>             | <u>TRADE NAME</u><br><u>(DOSAGE FORM; ROUTE)</u>  | <u>APPLICANT NAME</u> | <u>NDA NO.</u><br><u>APPROVAL DATE</u> | <u>PATENT NO.</u><br><u>EXP. DATE</u>  | <u>EXCLUSIVITY</u><br><u>EXP. DATE</u> |
|---|---|-----------------------|--|--|--|
| BETAMETHASONE DIPROPIONATE<br>EQ 0.05% BASE                   | BETAMETHASONE DIPROPIONATE<br>(OINTMENT; TOPICAL) | PHARMADERM/BYK-GLDN   | 19-140<br>09-04-84                     |  |  |
| BETAMETHASONE DIPROPIONATE<br>EQ 0.05% BASE                   | BETAMETHASONE DIPROPIONATE<br>(OINTMENT; TOPICAL) | E FOUGERA/BYK-GLDN    | 19-141<br>09-04-84                     |  |  |
| BETAMETHASONE DIPROPIONATE<br>EQ 0.05% BASE                   | ALPHATREX<br>(OINTMENT; TOPICAL)                  | SAVAGE LABS/BYK-GLDN  | 19-143<br>09-04-84                     |  |  |
| BETAMETHASONE DIPROPIONATE<br>EQ 0.05% BASE                   | DIPROSONE<br>(CREAM; TOPICAL)                     | SCHERING              | 17-536<br>01-29-75                     |  | D-1<br>09-24-86                        |
| BETAMETHASONE DIPROPIONATE<br>EQ 0.05% BASE                   | DIPROSONE<br>(OINTMENT; TOPICAL)                  | SCHERING              | 17-691<br>04-15-76                     |  | D-1<br>09-24-86                        |
| BETAMETHASONE DIPROPIONATE<br>EQ 0.05% BASE                   | DIPROSONE<br>(LOTION; TOPICAL)                    | SCHERING              | 17-781<br>02-01-77                     |  | D-1<br>09-24-86                        |
| BETAMETHASONE DIPROPIONATE<br>EQ 0.1% BASE                    | DIPROSONE<br>(AEROSOL; TOPICAL)                   | SCHERING              | 17-829<br>05-24-77                     |  | D-1<br>09-24-86                        |
| BETAMETHASONE DIPROPIONATE; CLOTRIMAZOLE<br>EQ 0.05% BASE; 1% | LOTTRISONE<br>(CREAM; TOPICAL)                    | SCHERING              | 18-827<br>07-10-84                     | 3660577<br>05-02-89<br>3705172<br>12-05-89<br>4298604<br>11-03-98<br>3839573<br>10-01-91 | NC<br>09-24-86                         |
| BETAMETHASONE VALERATE<br>EQ 0.1% BASE                        | BETA-VAL<br>(CREAM; TOPICAL)                      | LEMMON                | 18-642<br>03-24-83                     |  |  |
| BETAMETHASONE VALERATE<br>EQ 0.1% BASE                        | BETADERM<br>(CREAM; TOPICAL)                      | TJ ROACO              | 18-839<br>06-30-83                     |  |  |
| BETAMETHASONE VALERATE<br>EQ 0.1% BASE                        | BETAMETHASONE VALERATE<br>(CREAM; TOPICAL)        | PHARMADERM/BYK-GLDN   | 18-860<br>08-31-83                     |  |  |
| BETAMETHASONE VALERATE<br>EQ 0.1% BASE                        | BETAMETHASONE VALERATE<br>(CREAM; TOPICAL)        | E FOUGERA/BYK-GLDN    | 18-861<br>08-31-83                     |  |  |

**TABLE IV. NDA'S APPROVED FROM 1-1-82 TO 2-28-85 AND NDA'S WITH APPROPRIATE PATENT AND EXCLUSIVITY INFORMATION**

| <u>ACTIVE INGREDIENT(S)<br/>STRENGTH(S)</u> | <u>TRADE NAME<br/>(DOSAGE FORM; ROUTE)</u>    | <u>APPLICANT NAME</u> | <u>NDA NO.<br/>APPROVAL DATE</u> | <u>PATENT NO.<br/>EXP. DATE</u>            | <u>EXCLUSIVITY<br/>EXP. DATE</u> |
|---|---|-----------------------|----------------------------------|--|----------------------------------|
| BETAMETHASONE VALERATE<br>EQ 0.1% BASE      | BETATREX<br>(CREAM; TOPICAL)                  | SAVAGE LABS/BYK-GLDN  | 18-862<br>08-31-83               |  |                                  |
| BETAMETHASONE VALERATE<br>EQ 0.1% BASE      | BETATREX<br>(OINTMENT; TOPICAL)               | SAVAGE LABS/BYK-GLDN  | 18-863<br>08-31-83               |  |                                  |
| BETAMETHASONE VALERATE<br>EQ 0.1% BASE      | BETAMETHASONE VALERATE<br>(OINTMENT; TOPICAL) | PHARMADERM/BYK-GLDN   | 18-864<br>08-31-83               |  |                                  |
| BETAMETHASONE VALERATE<br>EQ 0.1% BASE      | BETAMETHASONE VALERATE<br>(OINTMENT; TOPICAL) | E FOUGERA/BYK-GLDN    | 18-865<br>08-31-83               |  |                                  |
| BETAMETHASONE VALERATE<br>EQ 0.1% BASE      | BETAMETHASONE VALERATE<br>(LOTION; TOPICAL)   | E FOUGERA/BYK-GLDN    | 18-866<br>08-31-83               |  |                                  |
| BETAMETHASONE VALERATE<br>EQ 0.1% BASE      | BETATREX<br>(LOTION; TOPICAL)                 | SAVAGE LABS/BYK-GLDN  | 18-867<br>08-31-83               |  |                                  |
| BETAMETHASONE VALERATE<br>EQ 0.1% BASE      | BETAMETHASONE VALERATE<br>(LOTION; TOPICAL)   | PHARMADERM/BYK-GLDN   | 18-870<br>08-31-83               |  |                                  |
| BETHANIDINE SULFATE<br>10MG                 | TENATHAN<br>(TABLET; ORAL)                    | AH ROBINS             | 17-675<br>05-29-81               | 3495013<br>02-10-87                        |                                  |
| BETHANIDINE SULFATE<br>25MG                 | TENATHAN<br>(TABLET; ORAL)                    | AH ROBINS             | 17-675<br>05-29-81               | 3495013<br>02-10-87                        |                                  |
| BITOLTEROL MESYLATE<br>0.8%                 | TORNALATE<br>(AEROSOL; INHALATION)            | WINTHROP-BREON/STERL  | 18-770<br>12-28-84               | 4138581<br>02-06-96                        | NCE<br>12-28-89                  |
| BRETYLIUM TOSYLATE<br>50MG/ML               | BRETYLLOL<br>(INJECTABLE; INJECTION)          | AM CRITICAL CARE/AHS  | 17-954<br>07-18-78               | RE29618<br>04-29-86                        |                                  |
| BROMOCRIPTINE MESYLATE<br>EQ 2.5MG BASE     | PARLODEL<br>(TABLET; ORAL)                    | SANDOZ PHARMS/SANDOZ  | 17-962<br>06-28-78               | 3752888<br>08-14-90<br>3752814<br>08-14-90 | I-16<br>12-14-87                 |
| BROMOCRIPTINE MESYLATE<br>EQ 5MG BASE       | PARLODEL<br>(CAPSULE; ORAL)                   | SANDOZ PHARMS/SANDOZ  | 17-962<br>03-01-82               | 3752888<br>08-14-90<br>3752814<br>08-14-90 | I-16<br>12-14-87                 |

TABLE IV. NDA'S APPROVED FROM 1-1-82 TO 2-28-85 AND NDA'S WITH APPROPRIATE PATENT AND EXCLUSIVITY INFORMATION

| <u>ACTIVE INGREDIENT(S)</u><br><u>STRENGTH(S)</u>  | <u>TRADE NAME</u><br><u>(DOSAGE FORM; ROUTE)</u>  | <u>APPLICANT NAME</u> | <u>NDA NO.</u><br><u>APPROVAL DATE</u> | <u>PATENT NO.</u><br><u>EXP. DATE</u>      | <u>EXCLUSIVITY</u><br><u>EXP. DATE</u> |
|--|---|-----------------------|--|--|--|
| BROMODIPHENHYDRAMINE HYDROCHLORIDE;<br>CODEINE PHOSPHATE<br>12.5MG/5ML; 10MG/5ML   | AMBENYL<br>(SYRUP; ORAL)                          | MARION LABORATORIES   | 09-319<br>01-10-84                     |  |  |
| BROMPHENIRAMINE MALEATE;<br>CODEINE PHOSPHATE;<br>PHENYLPROPANOLAMINE HYDROCHLORIDE<br>2MG/5ML; 10MG/5ML; 12.5MG/5ML       | DIMETANE-DC<br>(SYRUP; ORAL)                      | AH ROBINS             | 11-694<br>03-29-84                     |  |  |
| BROMPHENIRAMINE MALEATE;<br>DEXTROMETHORPHAN HYDROBROMIDE;<br>PSEUDOEPHEDRINE HYDROCHLORIDE<br>2MG/5ML; 10MG/5ML; 30MG/5ML | DIMETANE-DX<br>(SYRUP; ORAL)                      | AH ROBINS             | 11-694<br>03-29-84                     |  |  |
| BROMPHENIRAMINE MALEATE;<br>DEXTROMETHORPHAN HYDROBROMIDE;<br>PSEUDOEPHEDRINE HYDROCHLORIDE<br>2MG/5ML; 10MG/5ML; 30MG/5ML | DIMETANE-DX<br>(SYRUP; ORAL)                      | AH ROBINS             | 19-279<br>08-24-84                     |  |  |
| BROMPHENIRAMINE MALEATE;<br>PHENYLPROPANOLAMINE HYDROCHLORIDE<br>12MG; 75MG  | DIMETAPP<br>(TABLET, CONTROLLED<br>RELEASE; ORAL) | AH ROBINS             | 12-436<br>04-02-84                     |  |  |
| BROMPHENIRAMINE MALEATE;<br>PHENYLPROPANOLAMINE HYDROCHLORIDE<br>4MG/5ML; 25MG/5ML   | ELIXIR DIMETAPP<br>(ELIXIR; ORAL)                 | AH ROBINS             | 13-087<br>03-29-84                     |  |  |
| BUMETANIDE<br>1MG  | BUMEX<br>(TABLET; ORAL)                           | HOFFMANN-LA ROCHE     | 18-225<br>02-28-83                     | 3634583<br>01-11-89<br>3806534<br>04-23-91 | NCE<br>02-28-93                        |
| BUMETANIDE<br>0.5MG  | BUMEX<br>(TABLET; ORAL)                           | HOFFMANN-LA ROCHE     | 18-225<br>02-28-83                     | 3634583<br>01-11-89<br>3806534             | NCE<br>02-28-93                        |
| BUMETANIDE<br>0.25MG/ML  | BUMEX<br>(INJECTABLE; INJECTION)                  | HOFFMANN-LA ROCHE     | 18-226<br>02-28-83                     | 3634583<br>01-11-89<br>3806534<br>04-23-91 | NCE<br>02-28-93                        |

**TABLE IV. NDA'S APPROVED FROM 1-1-82 TO 2-28-85 AND NDA'S WITH APPROPRIATE PATENT AND EXCLUSIVITY INFORMATION**

| <u>ACTIVE INGREDIENT(S)<br/>STRENGTH(S)</u>                                | <u>TRADE NAME<br/>(DOSAGE FORM; ROUTE)</u> | <u>APPLICANT NAME</u> | <u>NDA NO.<br/>APPROVAL DATE</u> | <u>PATENT NO.<br/>EXP. DATE</u>  | <u>EXCLUSIVITY<br/>EXP. DATE</u> |
|--|--|-----------------------|----------------------------------|--|----------------------------------|
| BUPIVACAINE HYDROCHLORIDE; DEXTROSE<br>0.75%; 8.25%                        | MARCAINE SPINAL<br>(INJECTABLE; INJECTION) | BREON LABS/STERLING   | 18-692<br>05-04-84               |  | NC<br>09-24-86                   |
| BUPIVACAINE HYDROCHLORIDE;<br>EPINEPHRINE BITARTRATE<br>0.5%; 0.0091MG/ML  | SENSORCAINE<br>(INJECTABLE; INJECTION)     | ASTRA PHARM PRODS     | 18-304<br>09-02-83               |  |                                  |
| BUPIVACAINE HYDROCHLORIDE;<br>EPINEPHRINE BITARTRATE<br>0.75%; 0.0091MG/ML | SENSORCAINE<br>(INJECTABLE; INJECTION)     | ASTRA PHARM PRODS     | 18-304<br>09-02-83               |  |                                  |
| BUTORPHANOL TARTRATE<br>1MG/ML   | STADOL<br>(INJECTABLE; INJECTION)          | BRISTOL LABS/B-M      | 17-857<br>08-22-78               | 3819635<br>06-25-91  |                                  |
| BUTORPHANOL TARTRATE<br>2MG/ML   | STADOL<br>(INJECTABLE; INJECTION)          | BRISTOL LABS/B-M      | 17-857<br>08-22-78               | 3819635<br>06-25-91  |                                  |
| CALCEFEDIOL, ANHYDROUS<br>0.02MG   | CALDEROL<br>(CAPSULE; ORAL)                | UPJOHN                | 18-312<br>08-05-80               | 3833622<br>09-03-91<br>3565924<br>03-23-86   |                                  |
| CALCEFEDIOL, ANHYDROUS<br>0.05MG   | CALDEROL<br>(CAPSULE; ORAL)                | UPJOHN                | 18-312<br>08-05-80               | 3833622<br>09-03-91<br>3565924<br>03-23-86   |                                  |
| CALCITONIN<br>200 IU/VIAL  | CALCIMAR<br>(INJECTABLE; INJECTION)        | ARMOUR PHARM          | 17-769<br>12-21-84               |  | I-18<br>12-21-87                 |
| CALCITONIN<br>400 IU/VIAL  | CALCIMAR<br>(INJECTABLE; INJECTION)        | ARMOUR PHARM          | 17-497<br>12-21-84               |  | I-18<br>12-21-87                 |
| CALCITRIOL<br>0.25 UGM   | ROCALTROL<br>(CAPSULE; ORAL)               | HOFFMANN-LA ROCHE     | 18-044<br>08-17-78               | 3697559<br>10-10-89<br>4391802<br>07-05-00<br>4341774<br>07-27-99<br>4225596<br>09-30-97 |                                  |

TABLE IV. NDA'S APPROVED FROM 1-1-82 TO 2-28-85 AND NDA'S WITH APPROPRIATE PATENT AND EXCLUSIVITY INFORMATION

| <u>ACTIVE INGREDIENT(S)<br/>STRENGTH(S)</u>  | <u>TRADE NAME<br/>(DOSAGE FORM; ROUTE)</u>  | <u>APPLICANT NAME</u> | <u>NDA NO.<br/>APPROVAL DATE</u> | <u>PATENT NO.<br/>EXP. DATE</u>  | <u>EXCLUSIVITY<br/>EXP. DATE</u> |
|--|---|-----------------------|----------------------------------|--|----------------------------------|
| <b>CALCITRIOL<br/>0.5 UGM</b>  | <b>ROCALTROL<br/>(CAPSULE; ORAL)</b>  | HOFFMANN-LA ROCHE     | 18-044<br>08-17-78               | 3697559<br>10-10-89<br>4391802<br>07-05-00<br>4341774<br>07-27-99<br>4225596<br>09-30-97 |                                  |
| <b>CALCIUM CHLORIDE; DEXTROSE;<br/>MAGNESIUM CHLORIDE; POTASSIUM CHLORIDE;<br/>SODIUM ACETATE;<br/>SODIUM CHLORIDE; SODIUM CITRATE<br/>34MG/100ML; 5GM/100ML; 30MG/100ML;<br/>74MG/100ML; 640MG/100ML; 500MG/100ML;<br/>74MG/100ML</b> | <b>ISOLYTE E W/ DEXTROSE 5%<br/>IN PLASTIC CONTAINER<br/>(INJECTABLE; INJECTION)</b>                    | AM MCGAW/AM HOSP      | 18-269<br>01-17-83               |  |                                  |
| <b>CALCIUM CHLORIDE; DEXTROSE;<br/>MAGNESIUM CHLORIDE;<br/>SODIUM ACETATE; SODIUM CHLORIDE<br/>510MG/100ML; 30GM/100ML; 200MG/100ML;<br/>9.2GM/100ML; 9.6GM/100ML</b>  | <b>DIALYTE CONCENTRATE<br/>W/ DEXTROSE 30%<br/>IN PLASTIC CONTAINER<br/>(SOLUTION; INTRAPERITONEAL)</b> | AM MCGAW/AM HOSP      | 18-807<br>08-26-83               |  |                                  |
| <b>CALCIUM CHLORIDE; DEXTROSE;<br/>MAGNESIUM CHLORIDE; SODIUM ACETATE;<br/>SODIUM CHLORIDE<br/>510MG/100ML; 50GM/100ML; 200MG/100ML;<br/>9.2GM/100ML; 9.6GM/100ML</b>  | <b>DIALYTE CONCENTRATE<br/>W/ DEXTROSE 50%<br/>IN PLASTIC CONTAINER<br/>(SOLUTION; INTRAPERITONEAL)</b> | AM MCGAW/AM HOSP      | 18-807<br>08-26-83               |  |                                  |
| <b>CALCIUM CHLORIDE; DEXTROSE;<br/>MAGNESIUM CHLORIDE; SODIUM ACETATE;<br/>SODIUM CHLORIDE<br/>510MG/100ML; 30GM/100ML; 200MG/100ML;<br/>9.4GM/100ML; 11GM/100ML</b>   | <b>DIALYTE CONCENTRATE<br/>W/ DEXTROSE 30%<br/>IN PLASTIC CONTAINER<br/>(SOLUTION; INTRAPERITONEAL)</b> | AM MCGAW/AM HOSP      | 18-807<br>08-26-83               |  |                                  |
| <b>CALCIUM CHLORIDE; DEXTROSE;<br/>MAGNESIUM CHLORIDE; SODIUM ACETATE;<br/>SODIUM CHLORIDE<br/>510MG/100ML; 50GM/100ML; 200MG/100ML;<br/>9.4GM/100ML; 11GM/100ML</b>   | <b>DIALYTE CONCENTRATE<br/>W/ DEXTROSE 50%<br/>IN PLASTIC CONTAINER<br/>(SOLUTION; INTRAPERITONEAL)</b> | AM MCGAW/AM HOSP      | 18-807<br>08-26-83               |  |                                  |

TABLE IV. NDA'S APPROVED FROM 1-1-82 TO 2-28-85 AND NDA'S WITH APPROPRIATE PATENT AND EXCLUSIVITY INFORMATION

| <u>ACTIVE INGREDIENT(S)<br/>STRENGTH(S)</u>  | <u>TRADE NAME<br/>(DOSAGE FORM; ROUTE)</u>  | <u>APPLICANT NAME</u> | <u>NDA NO.<br/>APPROVAL DATE</u> | <u>PATENT NO.<br/>EXP. DATE</u> | <u>EXCLUSIVITY<br/>EXP. DATE</u> |
|--|---|-----------------------|----------------------------------|---------------------------------|----------------------------------|
| CALCIUM CHLORIDE; DEXTROSE;<br>MAGNESIUM CHLORIDE;<br>SODIUM CHLORIDE; SODIUM LACTATE<br>25.7MG/100ML; 1.5GM/100ML;<br>15.2MG/100ML; 567MG/100ML; <b>392MG/100ML</b> | DELFLX<br>W/ DEXTROSE 1.5%<br>IN PLASTIC CONTAINER<br>(SOLUTION; INTRAPERITONEAL)                   | DELMED                | 18-883<br>11-30-84               |                                 |                                  |
| CALCIUM CHLORIDE; DEXTROSE;<br>MAGNESIUM CHLORIDE;<br>SODIUM CHLORIDE; SODIUM LACTATE<br>25.7MG/100ML; 2.5GM/100ML;<br>15.2MG/100ML; 567MG/100ML; <b>392MG/100ML</b> | DELFLX<br>W/ DEXTROSE 2.5%<br>IN PLASTIC CONTAINER<br>(SOLUTION; INTRAPERITONEAL)                   | DELMED                | 18-883<br>11-30-84               |                                 |                                  |
| CALCIUM CHLORIDE; DEXTROSE;<br>MAGNESIUM CHLORIDE;<br>SODIUM CHLORIDE; SODIUM LACTATE<br>25.7MG/100ML; 4.25GM/100ML;<br>15.2MG/100ML; 567MG/100ML; 392MG/100ML       | DELFLX<br>W/ DEXTROSE 4.25%<br>IN PLASTIC CONTAINER<br>(SOLUTION; INTRAPERITONEAL)                  | DELMED                | 18-883<br>11-30-84               |                                 |                                  |
| CALCIUM CHLORIDE; DEXTROSE;<br>MAGNESIUM CHLORIDE; SODIUM CHLORIDE;<br>SODIUM LACTATE<br>25.7MG/100ML; 1.5GM/100ML;<br>5.08MG/100ML; 538MG/100ML; 448MG/100ML        | DELFLX<br>W/ DEXTROSE 1.5%<br>LOW MAGNESIUM<br>IN PLASTIC CONTAINER<br>(SOLUTION; INTRAPERITONEAL)  | DELMED                | 18-883<br>11-30-84               |                                 |                                  |
| CALCIUM CHLORIDE; DEXTROSE;<br>MAGNESIUM CHLORIDE; SODIUM CHLORIDE;<br>SODIUM LACTATE<br>25.7MG/100ML; 2.5GM/100ML;<br>5.08MG/100ML; 538MG/100ML; 448MG/100ML        | DELFLX<br>W/ DEXTROSE 2.5%<br>LOW MAGNESIUM<br>IN PLASTIC CONTAINER<br>(SOLUTION; INTRAPERITONEAL)  | DELMED                | 18-883<br>11-30-84               |                                 |                                  |
| CALCIUM CHLORIDE; DEXTROSE;<br>MAGNESIUM CHLORIDE; SODIUM CHLORIDE;<br>SODIUM LACTATE<br>25.7MG/100ML; 4.25GM/100ML;<br>15.2MG/100ML; 567MG/100ML; 392MG/100ML       | DELFLX<br>W/ DEXTROSE 4.25%<br>LOW MAGNESIUM<br>IN PLASTIC CONTAINER<br>(SOLUTION; INTRAPERITONEAL) | DELMED                | 18-883<br>11-30-84               |                                 |                                  |

**TABLE IV. NDA'S APPROVED FROM 1-1-82 TO 2-28-85 AND NDA'S WITH APPROPRIATE PATENT AND EXCLUSIVITY INFORMATION**

| <u>ACTIVE INGREDIENT(S)<br/>STRENGTH(S)</u>  | <u>TRADE NAME<br/>(DOSAGE FORM; ROUTE)</u>  | <u>APPLICANT NAME</u> | <u>NDA NO.<br/>APPROVAL DATE</u> | <u>PATENT NO.<br/>EXP. DATE</u> | <u>EXCLUSIVITY<br/>EXP. DATE</u> |
|--|---|-----------------------|----------------------------------|---------------------------------|----------------------------------|
| CALCIUM CHLORIDE; DEXTROSE;<br>MAGNESIUM CHLORIDE; SODIUM CHLORIDE;<br>SODIUM LACTATE<br>25.7MG/100ML; 1.5GM/100ML;<br>5.08MG/100ML; 538MG/100ML; 448MG/100ML                                  | INPERSOL-LM<br>W/ DEXTROSE 1.5%<br>IN PLASTIC CONTAINER<br>(SOLUTION; INTRAPERITONEAL)  | ABBOTT LABORATORIES   | 18-379<br>07-07-82               |                                 |                                  |
| CALCIUM CHLORIDE; DEXTROSE;<br>MAGNESIUM CHLORIDE; SODIUM CHLORIDE;<br>SODIUM LACTATE<br>25.7MG/100ML; 2.5GM/100ML;<br>5.08MG/100ML; 538MG/100ML; 448MG/100ML                                  | INPERSOL-LM<br>W/ DEXTROSE 2.5%<br>IN PLASTIC CONTAINER<br>(SOLUTION; INTRAPERITONEAL)  | ABBOTT LABORATORIES   | 18-379<br>07-07-82               |                                 |                                  |
| CALCIUM CHLORIDE; DEXTROSE;<br>MAGNESIUM CHLORIDE; SODIUM CHLORIDE;<br>SODIUM LACTATE<br>25.7MG/100ML; 4.25GM/100ML; 5.08MG/100ML;<br>538MG/100ML; 448MG/100ML                                 | INPERSOL-LM<br>W/ DEXTROSE 4.25%<br>IN PLASTIC CONTAINER<br>(SOLUTION; INTRAPERITONEAL) | ABBOTT LABORATORIES   | 18-379<br>07-07-82               |                                 |                                  |
| CALCIUM CHLORIDE; DEXTROSE;<br>MAGNESIUM CHLORIDE; SODIUM CHLORIDE;<br>SODIUM LACTATE<br>26MG/100ML; 2.5GM/100ML; 15MG/100ML;<br>560MG/100ML; 390MG/100ML                                      | DIALYTE<br>W/ DEXTROSE 2.5%<br>IN PLASTIC CONTAINER<br>(SOLUTION; INTRAPERITONEAL)      | AM MCGAW/AM HOSP      | 18-460<br>11-02-83               |                                 |                                  |
| CALCIUM CHLORIDE; DEXTROSE;<br>POTASSIUM CHLORIDE; SODIUM CHLORIDE<br>33MG/100ML; 5GM/100ML;<br>30MG/100ML; 860MG/100ML  | DEXTROSE 5% AND RINGER'S<br>IN PLASTIC CONTAINER<br>(INJECTABLE; INJECTION)             | TRAVENOL LABS         | 18-635<br>02-07-83               |                                 |                                  |
| CALCIUM CHLORIDE; MAGNESIUM CHLORIDE;<br>POTASSIUM CHLORIDE; SODIUM ACETATE;<br>SODIUM CHLORIDE<br>16.5MG/ML; 25.4MG/ML; 74.6MG/ML;<br>121MG/ML; 16.1MG/ML                                     | TPN ELECTROLYTES<br>IN PLASTIC CONTAINER<br>(INJECTABLE; INJECTION)                     | ABBOTT LABORATORIES   | 18-895<br>07-20-84               |                                 | NC<br>09-24-86                   |
| CALCIUM CHLORIDE; MAGNESIUM CHLORIDE;<br>POTASSIUM CHLORIDE; SODIUM ACETATE;<br>SODIUM CHLORIDE; SODIUM CITRATE<br>35MG/100ML; 30MG/100ML; 74MG/100ML;<br>640MG/100ML; 500MG/100ML; 74MG/100ML | ISOLYTE E<br>IN PLASTIC CONTAINER<br>(INJECTABLE; INJECTION)                            | AM MCGAW/AM HOSP      | 18-899<br>10-31-83               |                                 |                                  |

**TABLE IV. NDA'S APPROVED FROM 1-1-82 TO 2-28-85 AND NDA'S WITH APPROPRIATE PATENT AND EXCLUSIVITY INFORMATION**

| <u>ACTIVE INGREDIENT(S)<br/>STRENGTH(S)</u>   | <u>TRADE NAME<br/>(DOSAGE FORM; ROUTE)</u>                            | <u>APPLICANT NAME</u> | <u>NDA NO.<br/>APPROVAL DATE</u> | <u>PATENT NO.<br/>EXP. DATE</u> | <u>EXCLUSIVITY<br/>EXP. DATE</u> |
|---|---|-----------------------|----------------------------------|---------------------------------|----------------------------------|
| CALCIUM CHLORIDE; MAGNESIUM CHLORIDE;<br>POTASSIUM CHLORIDE;<br>SODIUM CHLORIDE<br>17.6MG/100ML; 325.3MG/100ML;<br>119.3MG/100ML; 643MG/100ML | PLEGISOL<br>IN PLASTIC CONTAINER<br>(SOLUTION;<br>PERFUSION, CARDIAC) | ABBOTT LABORATORIES   | 18-608<br>02-26-82               |                                 | NC<br>09-24-86                   |
| CALCIUM CHLORIDE; POTASSIUM CHLORIDE;<br>SODIUM ACETATE; SODIUM CHLORIDE<br>20MG/100ML; 30MG/100ML; 380MG/100ML;<br>600MG/100ML               | ACETATED RINGER'S<br>IN PLASTIC CONTAINER<br>(INJECTABLE; INJECTION)  | AM MCGAW/AM HOSP      | 18-725<br>11-29-82               |                                 |                                  |
| CALCIUM CHLORIDE; POTASSIUM CHLORIDE;<br>SODIUM CHLORIDE<br>33MG/100ML; 30MG/100ML; 860MG/100ML   | RINGER'S<br>IN PLASTIC CONTAINER<br>(SOLUTION; IRRIGATION)            | TRAVENOL LABS         | 18-495<br>02-19-82               |                                 |                                  |
| CALCIUM CHLORIDE; POTASSIUM CHLORIDE;<br>SODIUM CHLORIDE<br>33MG/100ML; 30MG/100ML; 860MG/100ML   | RINGERS INJECTION<br>IN PLASTIC CONTAINER<br>(INJECTABLE; INJECTION)  | TRAVENOL LABS         | 18-648<br>02-07-83               |                                 |                                  |
| CALCIUM CHLORIDE; POTASSIUM CHLORIDE;<br>SODIUM CHLORIDE<br>33MG/100ML; 30MG/100ML; 860MG/100ML   | RINGER'S<br>IN PLASTIC CONTAINER<br>(INJECTABLE; INJECTION)           | AM MCGAW/AM HOSP      | 18-721<br>11-09-82               |                                 |                                  |
| CALCIUM CHLORIDE; POTASSIUM CHLORIDE;<br>SODIUM CHLORIDE; SODIUM LACTATE<br>20MG/100ML; 30MG/100ML;<br>600MG/100ML; 310MG/100ML               | LACTATED RINGER'S<br>IN PLASTIC CONTAINER<br>(SOLUTION; IRRIGATION)   | TRAVENOL LABS         | 18-494<br>02-19-82               |                                 |                                  |
| CALCIUM CHLORIDE; POTASSIUM CHLORIDE;<br>SODIUM CHLORIDE; SODIUM LACTATE<br>20MG/100ML; 30MG/100ML;<br>600MG/100ML; 310MG/100ML               | LACTATED RINGER'S<br>IN PLASTIC CONTAINER<br>(SOLUTION; IRRIGATION)   | AM MCGAW/AM HOSP      | 18-681<br>12-27-82               |                                 |                                  |
| CALCIUM CHLORIDE; POTASSIUM CHLORIDE;<br>SODIUM CHLORIDE; SODIUM LACTATE<br>20MG/100ML; 30MG/100ML;<br>600MG/100ML; 310MG/100ML               | LACTATED RINGER'S<br>IN PLASTIC CONTAINER<br>(SOLUTION; IRRIGATION)   | TRAVENOL LABS         | 18-921<br>04-03-84               |                                 |                                  |

**TABLE IV. NDA'S APPROVED FROM 1-1-82 TO 2-28-85 AND NDA'S WITH APPROPRIATE PATENT AND EXCLUSIVITY INFORMATION**

| <u>ACTIVE INGREDIENT(S)<br/>STRENGTH(S)</u>  | <u>TRADE NAME<br/>(DOSAGE FORM; ROUTE)</u> | <u>APPLICANT NAME</u> | <u>NDA NO.<br/>APPROVAL DATE</u> | <u>PATENT NO.<br/>EXP. DATE</u>            | <u>EXCLUSIVITY<br/>EXP. DATE</u>    |
|--|--|-----------------------|----------------------------------|--|-------------------------------------|
| CALCIUM METRIZOATE; MAGNESIUM METRIZOATE;<br>MEGLUMINE METRIZOATE; METRIZOATE SODIUM<br>0.78MG/ML; 0.15MG/ML; 75.9MG/ML; 16.6MG/ML | ISOPAQUE 440<br>(INJECTABLE; INJECTION)    | WINTHROP LABS/STERL   | 16-847<br>11-17-73               | 3476802<br>11-04-86                        |                                     |
| CALCIUM; MEGLUMINE; METRIZOIC ACID<br>0.35MG/ML; 140.1MG/ML; 461.8MG/ML  | ISOPAQUE 280<br>(INJECTABLE; INJECTION)    | WINTHROP LABS/STERL   | 17-506<br>04-30-74               | 3476802<br>11-04-86                        |                                     |
| CAPTOPRIL<br>12.5MG  | CAPOTEN<br>(TABLET; ORAL)                  | ER SQUIBB AND SONS    | 18-343<br>01-17-85               | 4105776<br>08-08-95                        | I-20<br>09-24-86<br>D-7<br>10-12-87 |
| CAPTOPRIL<br>25MG  | CAPOTEN<br>(TABLET; ORAL)                  | ER SQUIBB AND SONS    | 18-343<br>04-06-81               | 4105776<br>08-08-95                        | I-20<br>09-24-86<br>D-7<br>10-12-87 |
| CAPTOPRIL<br>50MG  | CAPOTEN<br>(TABLET; ORAL)                  | ER SQUIBB AND SONS    | 18-343<br>04-06-81               | 4105776<br>08-08-95                        | I-20<br>09-24-86<br>D-7<br>10-12-87 |
| CAPTOPRIL<br>100MG   | CAPOTEN<br>(TABLET; ORAL)                  | ER SQUIBB AND SONS    | 18-343<br>04-06-81               | 4105776<br>08-08-95                        | I-20<br>09-24-86<br>D-7<br>10-12-87 |
| CAPTOPRIL; HYDROCHLOROTHIAZIDE<br>25MG; 15MG   | CAPOZIDE 25/15<br>(TABLET; ORAL)           | ER SQUIBB AND SONS    | 18-709<br>10-12-84               | 4105776<br>08-08-95<br>4217347<br>08-12-97 | NC<br>10-12-87                      |
| CAPTOPRIL; HYDROCHLOROTHIAZIDE<br>50MG; 15MG   | CAPOZIDE 50/15<br>(TABLET; ORAL)           | ER SQUIBB AND SONS    | 18-709<br>10-12-84               | 4105776<br>08-08-95<br>4217347<br>08-12-97 | NC<br>10-12-87                      |
| CAPTOPRIL; HYDROCHLOROTHIAZIDE<br>50MG; 25MG   | CAPOZIDE 50/25<br>(TABLET; ORAL)           | ER SQUIBB AND SONS    | 18-709<br>10-12-84               | 4105776<br>08-08-95<br>4217347<br>08-12-97 | NC<br>10-12-87                      |

TABLE IV. NDA'S APPROVED FROM 1-1-82 TO 2-28-85 AND NDA'S WITH APPROPRIATE PATENT AND EXCLUSIVITY INFORMATION

| <u>ACTIVE INGREDIENT(S)<br/>STRENGTH(S)</u> | <u>TRADE NAME<br/>(DOSAGE FORM; ROUTE)</u> | <u>APPLICANT NAME</u> | <u>NDA NO.<br/>APPROVAL DATE</u> | <u>PATENT NO.<br/>EXP. DATE</u>   | <u>EXCLUSIVITY<br/>EXP. DATE</u> |
|---|--|-----------------------|----------------------------------|---|----------------------------------|
| CARBAMAZEPINE<br>200MG                      | TEGRETOL<br>(TABLET; ORAL)                 | GEIGY/CIBA-GEIGY      | 16-608<br>03-11-68               | 4409212<br>10-11-00   |                                  |
| CARBAMAZEPINE<br>100MG                      | TEGRETOL<br>(TABLET, CHEWABLE; ORAL)       | GEIGY/CIBA-GEIGY      | 18-281<br>12-14-81               | 4409212<br>10-11-00   |                                  |
| CARBIDOPA<br>25MG                           | LODOSYN<br>(TABLET; ORAL)                  | MS&D/MERCK            | 17-830<br>04-25-77               | 3462536<br>08-19-86<br>3830827<br>08-20-91<br>3781415<br>12-25-90   |                                  |
| CARBIDOPA; LEVODOPA<br>10MG; 100MG          | SINEMET<br>(TABLET; ORAL)                  | MS&D/MERCK            | 17-555<br>05-02-75               | 3462536<br>08-19-86<br>3769424<br>10-30-90<br>3781415<br>12-25-90<br>3830827<br>08-20-91<br>RE29892<br>10-30-90 |                                  |
| CARBIDOPA; LEVODOPA<br>25MG; 250MG          | SINEMET<br>(TABLET; ORAL)                  | MS&D/MERCK            | 17-555<br>05-02-75               | 3462536<br>08-19-86<br>3769424<br>10-30-90<br>3781415<br>12-25-90<br>3830827<br>08-20-91<br>RE29892<br>10-30-90 |                                  |

TABLE IV. NDA'S APPROVED FROM 1-1-82 TO 2-28-85 AND NDA'S WITH APPROPRIATE PATENT AND EXCLUSIVITY INFORMATION

| <u>ACTIVE INGREDIENT(S)</u><br><u>STRENGTH(S)</u> | <u>TRADE NAME</u><br><u>(DOSAGE FORM; ROUTE)</u>    | <u>APPLICANT NAME</u> | <u>NDA NO.</u><br><u>APPROVAL DATE</u> | <u>PATENT NO.</u><br><u>EXP. DATE</u>   | <u>EXCLUSIVITY</u><br><u>EXP. DATE</u> |
|---|---|-----------------------|--|---|--|
| CARBIDOPA; LEVODOPA<br>25MG; 100MG                | SINEMET<br>(TABLET; ORAL)                           | MS&D/MERCK            | 17-555<br>05-02-75                     | 3462536<br>08-19-86<br>3769424<br>10-30-90<br>3781415<br>12-25-90<br>3830827<br>08-20-91<br>RE29892<br>10-30-90 |  |
| CARBOPROST TROMETHAMINE<br>EQ 0.25MG BASE/ML      | PROSTIN/15M<br>(INJECTABLE; INJECTION)              | UPJOHN                | 17-989<br>01-09-79                     | 3728382<br>04-17-90   |  |
| CELLULOSE SODIUM PHOSPHATE<br>2.5GM/PACKET        | CALCIBIND<br>(POWDER; ORAL)                         | MISSION PHARMACAL     | 18-757<br>12-28-82                     |   | NCE<br>12-28-92                        |
| CERULETIDE DIETHYLAMINE<br>0.02MG/ML              | TYMTRAN<br>(INJECTABLE; INJECTION)                  | ADRIA LABORATORIES    | 18-296<br>12-24-81                     | 3472832<br>10-14-86   |  |
| CHENODIOL<br>250MG                                | CHENIX<br>(TABLET; ORAL)                            | ROMELL LABORATORIES   | 18-513<br>07-28-83                     |   | NCE<br>07-28-93                        |
| CHLORDIAZEPOXIDE<br>25MG                          | LIBRITABS<br>(TABLET; ORAL)                         | ROCHE PRODUCTS        | 13-071<br>10-31-66                     | 4316897<br>02-23-99   |  |
| CHLORDIAZEPOXIDE<br>5MG                           | LIBRITABS<br>(TABLET; ORAL)                         | ROCHE PRODUCTS        | 13-071<br>10-31-66                     | 4316897<br>02-23-99   |  |
| CHLORDIAZEPOXIDE<br>10MG                          | LIBRITABS<br>(TABLET; ORAL)                         | ROCHE PRODUCTS        | 13-071<br>10-31-66                     | 4316897<br>02-23-99   |  |
| CHLORDIAZEPOXIDE<br>30MG                          | LIBRELAZE<br>(CAPSULE, CONTROLLED<br>RELEASE; ORAL) | HOFFMANN-LA ROCHE     | 17-813<br>09-12-83                     | 4316897<br>02-23-99   | NDF<br>09-24-86                        |
| CHLORDIAZEPOXIDE HYDROCHLORIDE<br>5MG             | LIBRIUM<br>(CAPSULE; ORAL)                          | ROCHE PRODUCTS        | 12-249<br>02-24-60                     | 4316897<br>02-23-99   |  |
| CHLORDIAZEPOXIDE HYDROCHLORIDE<br>10MG            | LIBRIUM<br>(CAPSULE; ORAL)                          | ROCHE PRODUCTS        | 12-249<br>02-24-60                     | 4316897<br>02-23-99   |  |

TABLE IV. NDA'S APPROVED FROM 1-1-82 TO 2-28-85 AND NDA'S WITH APPROPRIATE PATENT AND EXCLUSIVITY INFORMATION

| <u>ACTIVE INGREDIENT(S)<br/>STRENGTH(S)</u>                        | <u>TRADE NAME<br/>(DOSAGE FORM; ROUTE)</u> | <u>APPLICANT NAME</u> | <u>NDA NO.<br/>APPROVAL DATE</u> | <u>PATENT NO.<br/>EXP. DATE</u> | <u>EXCLUSIVITY<br/>EXP. DATE</u> |
|--|--|-----------------------|----------------------------------|---------------------------------|----------------------------------|
| CHLORDIAZEPOXIDE HYDROCHLORIDE<br>25MG                             | LIBRIUM<br>(CAPSULE; ORAL)                 | ROCHE PRODUCTS        | 12-249<br>02-24-60               | 4316897<br>02-23-99             |                                  |
| CHLORDIAZEPOXIDE HYDROCHLORIDE<br>100MG/AMP                        | LIBRIUM<br>(INJECTABLE; INJECTION)         | HOFFMANN-LA ROCHE     | 12-301<br>07-21-61               | 4316897<br>02-23-99             |                                  |
| CHLORDIAZEPOXIDE HYDROCHLORIDE;<br>CLIDINIUM BROMIDE<br>5MG; 2.5MG | LIBRAX<br>(CAPSULE; ORAL)                  | HOFFMANN-LA ROCHE     | 12-750<br>05-02-61               | 4316897<br>02-23-99             |                                  |
| CHLORDIAZEPOXIDE; ESTROGENS, CONJUGATED<br>5MG; 0.2MG              | MENRIUM 5-2<br>(TABLET; ORAL)              | HOFFMANN-LA ROCHE     | 14-740<br>10-27-69               | 4316897<br>02-23-99             |                                  |
| CHLORDIAZEPOXIDE; ESTROGENS, CONJUGATED<br>5MG; 0.4MG              | MENRIUM 5-4<br>(TABLET; ORAL)              | HOFFMANN-LA ROCHE     | 14-740<br>10-27-69               | 4316897<br>02-23-99             |                                  |
| CHLORDIAZEPOXIDE; ESTROGENS, CONJUGATED<br>10MG; 0.4MG             | MENRIUM 10-4<br>(TABLET; ORAL)             | HOFFMANN-LA ROCHE     | 14-740<br>10-27-69               | 4316897<br>02-23-99             |                                  |
| CHLOROXINE<br>2%   | CAPITROL<br>(SHAMPOO; TOPICAL)             | WESTWOOD PHARMS       | 17-594<br>10-19-76               | 3886277<br>05-27-92             |                                  |
| CHLORTHALIDONE; CLONIDINE HYDROCHLORIDE<br>15MG; 0.1MG             | COMBIPRES<br>(TABLET; ORAL)                | BOEHRINGER INGELHEIM  | 17-503<br>08-22-74               | 3454701<br>07-08-86             |                                  |
| CHLORTHALIDONE; CLONIDINE HYDROCHLORIDE<br>15MG; 0.2MG             | COMBIPRES<br>(TABLET; ORAL)                | BOEHRINGER INGELHEIM  | 17-503<br>08-22-74               | 3454701<br>07-08-86             |                                  |
| CHLORTHALIDONE; CLONIDINE HYDROCHLORIDE<br>15MG; 0.3MG             | COMBIPRES<br>(TABLET; ORAL)                | BOEHRINGER INGELHEIM  | 17-503<br>04-10-84               | 3454701<br>07-08-86             |                                  |
| CHOLESTYRAMINE<br>EQ 4GM RESIN/PACKET                              | QUESTRAN<br>(POWDER; ORAL)                 | MEAD JOHNSON/B-M      | 16-019<br>12-06-66               | 3383281<br>05-18-85             | I-23<br>09-24-86                 |
| CHOLESTYRAMINE<br>EQ 4GM RESIN/PACKET                              | QUESTRAN<br>(POWDER; ORAL)                 | MEAD JOHNSON/B-M      | 16-640<br>08-03-73               | 3383281<br>05-18-85             | I-23<br>09-24-86                 |
| CHYMOPAPAIN<br>12,500 UNITS/VIAL                                   | DISCASC<br>(INJECTABLE; INJECTION)         | TRAVENOL LABS         | 18-625<br>01-18-84               |                                 | NCE<br>11-10-92                  |

TABLE IV. NDA'S APPROVED FROM 1-1-82 TO 2-28-85 AND NDA'S WITH APPROPRIATE PATENT AND EXCLUSIVITY INFORMATION

| <u>ACTIVE INGREDIENT(S)<br/>STRENGTH(S)</u>   | <u>TRADE NAME<br/>(DOSAGE FORM; ROUTE)</u> | <u>APPLICANT NAME</u> | <u>NDA NO.<br/>APPROVAL DATE</u> | <u>PATENT NO.<br/>EXP. DATE</u>            | <u>EXCLUSIVITY<br/>EXP. DATE</u> |
|---|--|-----------------------|----------------------------------|--|----------------------------------|
| CHYMOPAPAIN<br>10,000 UNITS/VIAL              | CHYMODIACTIN<br>(INJECTABLE; INJECTION)    | SMITH LABORATORIES    | 18-663<br>11-10-82               | 4439423<br>03-26-01                        | NCE<br>11-10-92                  |
| CHYMOPAPAIN<br>4,000 UNITS/VIAL               | CHYMODIACTIN<br>(INJECTABLE; INJECTION)    | SMITH LABORATORIES    | 18-663<br>08-21-84               | 4439423<br>03-26-01                        | NCE<br>11-10-92                  |
| CICLOPIROX OLAMINE<br>1%                      | LOPROX<br>(CREAM; TOPICAL)                 | HOECHST-ROUSSEL       | 18-748<br>12-30-82               | 3883545<br>05-13-92                        | NCE<br>12-30-92                  |
| CIMETIDINE<br>200MG                           | TAGAMET<br>(TABLET; ORAL)                  | SK&F LAB              | 17-920<br>08-16-77               | 3950333<br>04-13-93<br>4024271<br>05-17-94 |                                  |
| CIMETIDINE<br>300MG                           | TAGAMET<br>(TABLET; ORAL)                  | SK&F LAB              | 17-920<br>08-16-77               | 3950333<br>04-13-93<br>4024271<br>05-17-94 |                                  |
| CIMETIDINE<br>400MG                           | TAGAMET<br>(TABLET; ORAL)                  | SK&F LAB              | 17-920<br>12-14-83               | 3950333<br>04-13-93<br>4024271<br>05-17-94 | NS<br>09-24-86                   |
| CIMETIDINE HYDROCHLORIDE<br>EQ 300MG BASE/5ML | TAGAMET<br>(SOLUTION; ORAL)                | SK&F LAB              | 17-924<br>08-16-77               | 3950333<br>04-13-93<br>4024271<br>05-17-94 |                                  |
| CIMETIDINE HYDROCHLORIDE<br>EQ 150MG BASE/ML  | TAGAMET<br>(INJECTABLE; INJECTION)         | SK&F LAB              | 17-939<br>08-16-77               | 3950333<br>04-13-93<br>4024271<br>05-17-94 |                                  |
| CINOXACIN<br>250MG                            | CINOBAC<br>(CAPSULE; ORAL)                 | ELI LILLY             | 18-067<br>06-13-80               | 3669965<br>06-13-89                        |                                  |
| CINOXACIN<br>500MG                            | CINOBAC<br>(CAPSULE; ORAL)                 | ELI LILLY             | 18-067<br>06-13-80               | 3669965<br>06-13-89                        |                                  |

**TABLE IV. NDA'S APPROVED FROM 1-1-82 TO 2-28-85 AND NDA'S WITH APPROPRIATE PATENT AND EXCLUSIVITY INFORMATION**

| <u>ACTIVE INGREDIENT(S)<br/>STRENGTH(S)</u>  | <u>TRADE NAME<br/>(DOSAGE FORM; ROUTE)</u>                              | <u>APPLICANT NAME</u> | <u>NDA NO.<br/>APPROVAL DATE</u> | <u>PATENT NO.<br/>EXP. DATE</u> | <u>EXCLUSIVITY<br/>EXP. DATE</u> |
|--|---|-----------------------|----------------------------------|---------------------------------|----------------------------------|
| CISPLATIN<br>0.5MG/ML  | PLATINOL-AQ   | BRISTOL LABS/B-M      | 18-507<br>07-18-84               | 4177263<br>12-04-96             | NDF<br>09-24-86                  |
| CITRIC ACID; MAGNESIUM OXIDE;<br>SODIUM CARBONATE<br>3.24G/100ML; 380MG/100ML; 430MG/100ML | IRRIGATING SOLUTION G<br>IN PLASTIC CONTAINER<br>(SOLUTION; IRRIGATION) | TRAVENOL LABS         | 18-519<br>06-22-82               |                                 | NC<br>09-24-86                   |
| CITRIC ACID; MAGNESIUM OXIDE;<br>SODIUM CARBONATE<br>3.24G/100ML; 380MG/100ML; 430MG/100ML | UROLOGIC G<br>IN PLASTIC CONTAINER<br>(SOLUTION; IRRIGATION)            | ABBOTT LABORATORIES   | 18-904<br>05-27-83               |                                 | NC<br>09-24-86                   |
| CLEMASTINE FUMARATE;<br>PHENYLPROPANOLAMINE HYDROCHLORIDE<br>EQ 1MG BASE; 75MG             | TAVIST D<br>(TABLET, CONTROLLED<br>RELEASE; ORAL)                       | DORSEY LABS/SANDOZ    | 18-298<br>12-15-82               | 3933999<br>01-20-93             | NDF<br>09-24-86                  |
| CLOMIPHENE CITRATE<br>50MG   | CLOMIPHENE CITRATE<br>(TABLET; ORAL)                                    | PLANTEK/IKAPHARM      | 18-361<br>03-22-82               |                                 |                                  |
| CLONAZEPAM<br>0.5MG  | CLONOPIN<br>(TABLET; ORAL)  | HOFFMANN-LA ROCHE     | 17-533<br>06-04-75               | 4316897<br>02-23-99             |                                  |
| CLONAZEPAM<br>1MG  | CLONOPIN<br>(TABLET; ORAL)  | HOFFMANN-LA ROCHE     | 17-533<br>06-04-75               | 4316897<br>02-23-99             |                                  |
| CLONAZEPAM<br>2MG  | CLONOPIN<br>(TABLET; ORAL)  | HOFFMANN-LA ROCHE     | 17-533<br>06-04-75               | 4316897<br>02-23-99             |                                  |
| CLONIDINE<br>2.5MG   | CATAPRES-TTS-1<br>(FILM, CONTROLLED RELEASE;<br>PERCUTANEOUS)           | BOEHRINGER INGELHEIM  | 18-891<br>10-10-84               | 3454701<br>07-08-86             | NR<br>10-10-87                   |
| CLONIDINE<br>5MG   | CATAPRES-TTS-2<br>(FILM, CONTROLLED RELEASE;<br>PERCUTANEOUS)           | BOEHRINGER INGELHEIM  | 18-891<br>10-10-84               | 3454701<br>07-08-86             | NR<br>10-10-87                   |
| CLONIDINE<br>7.5MG   | CATAPRES-TTS-3<br>(FILM, CONTROLLED RELEASE;<br>PERCUTANEOUS)           | BOEHRINGER INGELHEIM  | 18-891<br>10-10-84               | 3454701<br>07-08-86             | NR<br>10-10-87                   |
| CLONIDINE HYDROCHLORIDE<br>0.1MG   | CATAPRES<br>(TABLET; ORAL)  | BOEHRINGER INGELHEIM  | 17-407<br>09-03-74               | 3454701<br>07-08-86             |                                  |

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| <u>ACTIVE INGREDIENT(S)<br/>STRENGTH(S)</u> | <u>TRADE NAME<br/>(DOSAGE FORM; ROUTE)</u> | <u>APPLICANT NAME</u> | <u>NDA NO.<br/>APPROVAL DATE</u> | <u>PATENT NO.<br/>EXP. DATE</u>                                   | <u>EXCLUSIVITY<br/>EXP. DATE</u> |
|---|--|-----------------------|----------------------------------|---|----------------------------------|
| CLONIDINE HYDROCHLORIDE<br>0.2MG            | CATAPRES<br>(TABLET; ORAL)                 | BOEHRINGER INGELHEIM  | 17-407<br>09-03-74               | 3454701<br>07-08-86   |                                  |
| CLONIDINE HYDROCHLORIDE<br>0.3MG            | CATAPRES<br>(TABLET; ORAL)                 | BOEHRINGER INGELHEIM  | 17-407<br>09-20-79               | 3454701<br>07-08-86   |                                  |
| CLORAZEPATE DIPOTASSIUM<br>3.75MG           | TRANXENE<br>(CAPSULE; ORAL)                | ABBOTT LABORATORIES   | 17-105<br>06-23-72               | RE28315<br>06-23-87   |                                  |
| CLORAZEPATE DIPOTASSIUM<br>7.5MG            | TRANXENE<br>(CAPSULE; ORAL)                | ABBOTT LABORATORIES   | 17-105<br>06-23-72               | RE28315<br>06-23-87   |                                  |
| CLORAZEPATE DIPOTASSIUM<br>15MG             | TRANXENE<br>(CAPSULE; ORAL)                | ABBOTT LABORATORIES   | 17-105<br>06-23-72               | RE28315<br>06-23-87   |                                  |
| CLORAZEPATE DIPOTASSIUM<br>22.5MG           | TRANXENE SD<br>(TABLET; ORAL)              | ABBOTT LABORATORIES   | 17-105<br>03-31-75               | RE28315<br>06-23-87   |                                  |
| CLORAZEPATE DIPOTASSIUM<br>11.25MG          | TRANXENE SD<br>(TABLET; ORAL)              | ABBOTT LABORATORIES   | 17-105<br>08-04-76               | RE28315<br>06-23-87   |                                  |
| CLORAZEPATE DIPOTASSIUM<br>3.75MG           | TRANXENE<br>(TABLET; ORAL)                 | ABBOTT LABORATORIES   | 17-105<br>03-10-80               | RE28315<br>06-23-87   |                                  |
| CLORAZEPATE DIPOTASSIUM<br>7.5MG            | TRANXENE<br>(TABLET; ORAL)                 | ABBOTT LABORATORIES   | 17-105<br>03-10-80               | RE28315<br>06-23-87   |                                  |
| CLORAZEPATE DIPOTASSIUM<br>15MG             | TRANXENE<br>(TABLET; ORAL)                 | ABBOTT LABORATORIES   | 17-105<br>03-10-80               | RE28315<br>06-23-87   |                                  |
| CLOTTRIMAZOLE<br>1%                         | LOTRIMIN<br>(SOLUTION; TOPICAL)            | SCHERING              | 17-613<br>02-03-75               | 3660577<br>05-02-89<br>3705172<br>12-05-89<br>3839573<br>10-01-91 |                                  |

**TABLE IV. NDA'S APPROVED FROM 1-1-82 TO 2-28-85 AND NDA'S WITH APPROPRIATE PATENT AND EXCLUSIVITY INFORMATION**

| <u>ACTIVE INGREDIENT(S)<br/>STRENGTH(S)</u> | <u>TRADE NAME<br/>(DOSAGE FORM; ROUTE)</u> | <u>APPLICANT NAME</u> | <u>NDA NO.<br/>APPROVAL DATE</u> | <u>PATENT NO.<br/>EXP. DATE</u>                                   | <u>EXCLUSIVITY<br/>EXP. DATE</u> |
|---|--|-----------------------|----------------------------------|---|----------------------------------|
| CLOTRIMAZOLE<br>1%                          | LOTTRIMIN<br>(CREAM; TOPICAL)              | SCHERING              | 17-619<br>03-18-75               | 3660577<br>05-02-89<br>3705172<br>12-05-89<br>3839573<br>10-01-91 |                                  |
| CLOTRIMAZOLE<br>1%                          | GYNE-LOTTRIMIN<br>(CREAM; VAGINAL)         | SCHERING              | 18-052<br>11-08-78               | 3839573<br>10-01-91<br>3705172<br>12-05-89<br>3660577<br>05-02-89 |                                  |
| CLOTRIMAZOLE<br>100MG                       | GYNE-LOTTRIMIN<br>(TABLET; VAGINAL)        | SCHERING              | 17-717<br>03-24-76               | 3839573<br>10-01-91<br>3705172<br>12-05-89<br>3660577<br>05-02-89 |                                  |
| CLOTRIMAZOLE<br>1%                          | MYCELEX<br>(SOLUTION; TOPICAL)             | MILES PHARMS/MILES    | 18-181<br>01-15-79               | 3839573<br>10-01-91<br>3705172<br>12-05-89<br>3660577<br>05-02-89 |                                  |
| CLOTRIMAZOLE<br>100MG                       | MYCELEX-G<br>(TABLET; VAGINAL)             | MILES PHARMS/MILES    | 18-182<br>02-27-79               | 3839573<br>10-01-91<br>3705172<br>12-05-89<br>3660577<br>05-02-89 |                                  |
| CLOTRIMAZOLE<br>1%                          | MYCELEX<br>(CREAM; TOPICAL)                | MILES PHARMS/MILES    | 18-183<br>01-15-79               | 3839573<br>10-01-91<br>3705172<br>12-05-89<br>3660577<br>05-02-89 |                                  |

**TABLE IV. NDA'S APPROVED FROM 1-1-82 TO 2-28-85 AND NDA'S WITH APPROPRIATE PATENT AND EXCLUSIVITY INFORMATION**

| <u>ACTIVE INGREDIENT(S)<br/>STRENGTH(S)</u>  | <u>TRADE NAME<br/>(DOSAGE FORM; ROUTE)</u> | <u>APPLICANT NAME</u> | <u>NDA NO.<br/>APPROVAL DATE</u> | <u>PATENT NO.<br/>EXP. DATE</u>                                   | <u>EXCLUSIVITY<br/>EXP. DATE</u> |
|--|--|-----------------------|----------------------------------|---|----------------------------------|
| CLOTTRIMAZOLE<br>1%  | MYCELEX-G<br>(CREAM; VAGINAL)              | MILES PHARMS/MILES    | 18-230<br>02-16-79               | 3839573<br>10-01-91<br>3705172<br>12-05-89<br>3660577<br>05-02-89 |                                  |
| CLOTTRIMAZOLE<br>10MG  | MYCELEX<br>(TROCHE/LOZENGE; ORAL)          | MILES PHARMS/MILES    | 18-713<br>06-17-83               | 3839573<br>10-01-91<br>3705172<br>12-05-89<br>3660577<br>05-02-89 | NDF<br>09-24-86                  |
| CLOTTRIMAZOLE<br>1%  | LOTRIMIN<br>(LOTION; TOPICAL)              | SCHERING              | 18-813<br>02-17-84               | 3839573<br>10-01-91<br>3705172<br>12-05-89<br>3660577<br>05-02-89 |                                  |
| CODEINE PHOSPHATE;<br>PHENYLEPHRINE HYDROCHLORIDE;<br>PROMETHAZINE HYDROCHLORIDE<br>10MG/5ML; 5MG/5ML; 6.25MG/5ML    | PHENERGAN VC W/<br>(SYRUP; ORAL) CODEINE   | WYETH LABS/AMHO       | 08-306<br>04-02-84               |   |                                  |
| CODEINE PHOSPHATE;<br>PROMETHAZINE HYDROCHLORIDE<br>10MG/5ML; 6.25MG/5ML   | PHENERGAN W/<br>(SYRUP; ORAL) CODEINE      | WYETH LABS/AMHO       | 08-306<br>04-02-84               |   |                                  |
| CODEINE PHOSPHATE;<br>PSEUDOEPHEDRINE HYDROCHLORIDE;<br>TRIPROLIDINE HYDROCHLORIDE<br>10MG/5ML; 30MG/5ML; 1.25MG/5ML | ACTIFED W/<br>(SYRUP; ORAL) CODEINE        | BURROUGHS WELLCOME    | 12-575<br>04-04-84               |   |                                  |
| COLESTIPOL HYDROCHLORIDE<br>5GM/PACKET   | COLESTID<br>(GRANULE; ORAL)                | UPJOHN                | 17-563<br>04-04-77               | 3692895<br>09-19-89   | I-24<br>09-24-86                 |
| COLESTIPOL HYDROCHLORIDE<br>500GM/BOT  | COLESTID<br>(GRANULE; ORAL)                | UPJOHN                | 17-563<br>04-04-77               | 3692895<br>09-19-89   | I-24<br>09-24-86                 |

**TABLE IV. NDA'S APPROVED FROM 1-1-82 TO 2-28-85 AND NDA'S WITH APPROPRIATE PATENT AND EXCLUSIVITY INFORMATION**

| <u>ACTIVE INGREDIENT(S)</u><br><u>STRENGTH(S)</u> | <u>TRADE NAME</u><br><u>(DOSAGE FORM; ROUTE)</u>  | <u>APPLICANT NAME</u> | <u>NDA NO.</u><br><u>APPROVAL DATE</u> | <u>PATENT NO.</u><br><u>EXP. DATE</u>   | <u>EXCLUSIVITY</u><br><u>EXP. DATE</u> |
|---|---|-----------------------|--|---|--|
| COPPER<br>89MG                                    | CU-7<br>(INTRAUTERINE DEVICE;<br>INTRAUTERINE)    | SEARLE PHARMS         | 17-408<br>02-25-74                     | 3563235<br>02-16-88<br>4040417<br>08-09-94<br>3783861<br>01-08-91<br>3803308<br>12-01-87<br>RE28399<br>04-29-92 |  |
| COPPER<br>120MG                                   | TATUM-T<br>(INTRAUTERINE DEVICE;<br>INTRAUTERINE) | SEARLE PHARMS         | 18-205<br>08-16-79                     | 3563235<br>02-16-88<br>4040417<br>08-09-94<br>3783861<br>01-08-91<br>3803308<br>12-01-87<br>RE28399<br>04-29-92 |  |
| CROMOLYN SODIUM<br>20MG                           | INTAL<br>(CAPSULE; INHALATION)                    | FISONS                | 16-990<br>06-20-73                     | 3686412<br>08-22-89<br>3777033<br>08-22-89<br>3419578<br>12-31-85   | I-22<br>09-24-86                       |
| CROMOLYN SODIUM<br>4%                             | NASALCROM<br>(SOLUTION; NASAL)                    | FISONS                | 18-306<br>03-18-83                     | 3686412<br>08-22-89<br>3777033<br>08-22-89<br>3419578<br>12-31-85<br>3975536<br>08-17-93<br>4053628<br>10-11-94 | NDF<br>09-24-86                        |

**TABLE IV. NDA'S APPROVED FROM 1-1-82 TO 2-28-85 AND NDA'S WITH APPROPRIATE PATENT AND EXCLUSIVITY INFORMATION**

| <u>ACTIVE INGREDIENT(S)</u><br><u>STRENGTH(S)</u> | <u>TRADE NAME</u><br><u>(DOSAGE FORM; ROUTE)</u> | <u>APPLICANT NAME</u> | <u>NDA NO.</u><br><u>APPROVAL DATE</u> | <u>PATENT NO.</u><br><u>EXP. DATE</u>   | <u>EXCLUSIVITY</u><br><u>EXP. DATE</u> |
|---|--|-----------------------|--|---|--|
| CROMOLYN SODIUM<br>4%                             | OPTICROM<br>(SOLUTION; OPHTHALMIC)               | FISONS                | 18-155<br>10-03-84                     | 3686412<br>08-22-89<br>3777033<br>08-22-89<br>3419578<br>12-31-85<br>3975536<br>08-17-93<br>4053628<br>10-11-94 | NDF<br>10-03-87                        |
| CROMOLYN SODIUM<br>10MG/ML                        | INTAL<br>(SOLUTION; INHALATION)                  | FISONS                | 18-596<br>05-28-82                     | 3686412<br>08-22-89<br>3777033<br>08-22-89<br>3419578<br>12-31-85<br>3975536<br>08-17-93                        | I-22<br>01-19-88                       |
| CYCLOBENZAPRINE HYDROCHLORIDE<br>5MG              | FLEXERIL<br>(TABLET; ORAL)                       | MS&D/MERCK            | 17-821<br>08-26-77                     | 3454643<br>07-08-86<br>3882246<br>05-06-92  |  |
| CYCLOBENZAPRINE HYDROCHLORIDE<br>10MG             | FLEXERIL<br>(TABLET; ORAL)                       | MS&D/MERCK            | 17-821<br>08-26-77                     | 3454643<br>07-08-86<br>3882246<br>05-06-92  |  |
| CYCLOPHOSPHAMIDE<br>1GM/VIAL                      | CYTOXAN<br>(INJECTABLE; INJECTION)               | MEAD JOHNSON/B-M      | 12-142<br>08-30-82                     |   | NS<br>09-24-86                         |
| CYCLOPHOSPHAMIDE<br>1GM/VIAL                      | NEOSAR<br>(INJECTABLE; INJECTION)                | ADRIA LABORATORIES    | 87-442<br>07-08-83                     |   | NS<br>09-24-86                         |
| CYCLOPHOSPHAMIDE<br>2GM/VIAL                      | CYTOXAN<br>(INJECTABLE; INJECTION)               | MEAD JOHNSON/B-M      | 12-142<br>08-30-82                     |   | NS<br>09-24-86                         |

**TABLE IV. NDA'S APPROVED FROM 1-1-82 TO 2-28-85 AND NDA'S WITH APPROPRIATE PATENT AND EXCLUSIVITY INFORMATION**

| <u>ACTIVE INGREDIENT(S)<br/>STRENGTH(S)</u> | <u>TRADE NAME<br/>(DOSAGE FORM; ROUTE)</u>  | <u>APPLICANT NAME</u> | <u>NDA NO.<br/>APPROVAL DATE</u> | <u>PATENT NO.<br/>EXP. DATE</u>            | <u>EXCLUSIVITY<br/>EXP. DATE</u> |
|---|---|-----------------------|----------------------------------|--|----------------------------------|
| DANTROLENE SODIUM<br>25MG                   | DANTRIUM<br>(CAPSULE; ORAL)                 | NORWICH EATON/P&G     | 17-443<br>01-15-74               | 3415821<br>12-10-85                        |                                  |
| DANTROLENE SODIUM<br>100MG                  | DANTRIUM<br>(CAPSULE; ORAL)                 | NORWICH EATON/P&G     | 17-443<br>01-15-74               | 3415821<br>12-10-85                        |                                  |
| DANTROLENE SODIUM<br>50MG                   | DANTRIUM<br>(CAPSULE; ORAL)                 | NORWICH EATON/P&G     | 17-443<br>10-10-75               | 3415821<br>12-10-85                        |                                  |
| DANTROLENE SODIUM<br>20MG/VIAL              | DANTRIUM<br>(INJECTABLE; INJECTION)         | NORWICH EATON/P&G     | 18-264<br>09-18-79               | 3415821<br>12-10-85                        |                                  |
| DEFEROXAMINE MESYLATE<br>500MG/VIAL         | DEFERAL MESYLATE<br>(INJECTABLE; INJECTION) | CIBA/CIBA-GEIGY       | 16-267<br>04-01-68               | 3471476<br>10-07-86                        |                                  |
| DESIPRAMINE HYDROCHLORIDE<br>25MG           | PERTOFRANE<br>(CAPSULE; ORAL)               | USV LABORATORIES      | 13-621<br>12-18-64               | 3454698<br>07-08-86<br>3454554<br>07-08-86 |                                  |
| DESIPRAMINE HYDROCHLORIDE<br>50MG           | PERTOFRANE<br>(CAPSULE; ORAL)               | USV LABORATORIES      | 13-621<br>04-10-68               | 3454698<br>07-08-86<br>3454554<br>07-08-86 |                                  |
| DESIPRAMINE HYDROCHLORIDE<br>25MG           | NORPRAMIN<br>(TABLET; ORAL)                 | MERRELL DOW/DOW CHEM  | 14-399<br>11-20-64               | 3454698<br>07-08-86<br>3454554<br>07-08-86 |                                  |
| DESIPRAMINE HYDROCHLORIDE<br>50MG           | NORPRAMIN<br>(TABLET; ORAL)                 | MERRELL DOW/DOW CHEM  | 14-399<br>01-09-67               | 3454698<br>07-08-86<br>3454554<br>07-08-86 |                                  |
| DESIPRAMINE HYDROCHLORIDE<br>75MG           | NORPRAMIN<br>(TABLET; ORAL)                 | MERRELL DOW/DOW CHEM  | 14-399<br>03-01-77               | 3454698<br>07-08-86<br>3454554<br>07-08-86 |                                  |

**TABLE IV. NDA'S APPROVED FROM 1-1-82 TO 2-28-85 AND NDA'S WITH APPROPRIATE PATENT AND EXCLUSIVITY INFORMATION**

| <u>ACTIVE INGREDIENT(S)</u><br><u>STRENGTH(S)</u> | <u>TRADE NAME</u><br><u>(DOSAGE FORM; ROUTE)</u> | <u>APPLICANT NAME</u> | <u>NDA NO.</u><br><u>APPROVAL DATE</u> | <u>PATENT NO.</u><br><u>EXP. DATE</u>      | <u>EXCLUSIVITY</u><br><u>EXP. DATE</u> |
|---|--|-----------------------|--|--|--|
| DESIPRAMINE HYDROCHLORIDE<br>100MG                | NORPRAMIN<br>(TABLET; ORAL)                      | MERRELL DOW/DOW CHEM  | 14-399<br>03-01-77                     | 3454698<br>07-08-86<br>3454554<br>07-08-86 |  |
| DESIPRAMINE HYDROCHLORIDE<br>150MG                | NORPRAMIN<br>(TABLET; ORAL)                      | MERRELL DOW/DOW CHEM  | 14-399<br>03-01-77                     | 3454698<br>07-08-86<br>3454554<br>07-08-86 |  |
| DESIPRAMINE HYDROCHLORIDE<br>10MG                 | NORPRAMIN<br>(TABLET; ORAL)                      | MERRELL DOW/DOW CHEM  | 14-399<br>02-11-82                     | 3454698<br>07-08-86<br>3454554<br>07-08-86 | NS<br>09-24-86                         |
| DESMOPRESSIN ACETATE<br>0.01%                     | DDAVP<br>(SOLUTION; NASAL)                       | ARMOUR PHARM          | 17-922<br>02-21-78                     | 3497491<br>02-24-87                        |  |
| DESMOPRESSIN ACETATE<br>0.004MG/ML                | DDAVP<br>(INJECTABLE; INJECTION)                 | ARMOUR PHARM          | 18-938<br>03-30-84                     | 3497491<br>02-24-87                        | NDF<br>09-24-86                        |
| DESONIDE<br>0.05%                                 | DESONEN<br>(CREAM; TOPICAL)                      | OWEN LABS/DERM PRODS  | 19-048<br>12-14-84                     |  |  |
| DESOXIMETASONE<br>0.05%                           | TOPICORT<br>(GEL; TOPICAL)                       | HOECHST-ROUSSEL       | 18-586<br>03-29-82                     |  | NDF<br>09-24-86                        |
| DESOXIMETASONE<br>0.05%                           | TOPICORT<br>(OINTMENT; TOPICAL)                  | HOECHST-ROUSSEL       | 18-594<br>01-17-85                     |  | NDF<br>09-24-86                        |
| DESOXIMETASONE<br>0.25%                           | TOPICORT<br>(OINTMENT; TOPICAL)                  | HOECHST-ROUSSEL       | 18-763<br>09-30-83                     |  | NDF<br>09-24-86                        |
| DEXAMETHASONE<br>0.5MG                            | DECADRON<br>(TABLET; ORAL)                       | MS&D/MERCK            | 11-664<br>10-30-58                     | 3375261<br>03-26-85<br>RE28369<br>03-26-85 |  |
| DEXAMETHASONE<br>0.75MG                           | DECADRON<br>(TABLET; ORAL)                       | MS&D/MERCK            | 11-664<br>10-30-58                     | 3375261<br>03-26-85<br>RE28369<br>03-26-85 |  |

**TABLE IV. NDA'S APPROVED FROM 1-1-82 TO 2-28-85 AND NDA'S WITH APPROPRIATE PATENT AND EXCLUSIVITY INFORMATION**

| <u>ACTIVE INGREDIENT(S)</u><br><u>STRENGTH(S)</u> | <u>TRADE NAME</u><br><u>(DOSAGE FORM; ROUTE)</u> | <u>APPLICANT NAME</u> | <u>NDA NO.</u><br><u>APPROVAL DATE</u> | <u>PATENT NO.</u><br><u>EXP. DATE</u>      | <u>EXCLUSIVITY</u><br><u>EXP. DATE</u> |
|---|--|-----------------------|--|--|--|
| DEXAMETHASONE<br>1.5MG                            | DECADRON<br>(TABLET; ORAL)                       | MS&D/MERCK            | 11-664<br>10-30-58                     | 3375261<br>03-26-85<br>RE28369<br>03-26-85 |  |
| DEXAMETHASONE<br>0.25MG                           | DECADRON<br>(TABLET; ORAL)                       | MS&D/MERCK            | 11-664<br>07-26-79                     | 3375261<br>03-26-85<br>RE28369<br>03-26-85 |  |
| DEXAMETHASONE<br>4MG                              | DECADRON<br>(TABLET; ORAL)                       | MS&D/MERCK            | 11-664<br>07-26-79                     | 3375261<br>03-26-85<br>RE28369<br>03-26-85 |  |
| DEXAMETHASONE<br>6MG                              | DECADRON<br>(TABLET; ORAL)                       | MS&D/MERCK            | 11-664<br>07-30-82                     | 3375261<br>03-26-85<br>RE28369<br>03-26-85 | NS<br>09-24-86                         |
| DEXAMETHASONE<br>6MG                              | DEXAMETHASONE<br>(TABLET; ORAL)                  | PAR PHARMACEUTICAL    | 88-481<br>11-28-83                     |  | NS<br>09-24-86                         |
| DEXAMETHASONE<br>6MG                              | DEXAMETHASONE<br>(TABLET; ORAL)                  | ROXANE LABORATORIES   | 88-316<br>09-15-83                     |  | NS<br>09-24-86                         |
| DEXAMETHASONE<br>0.5MG/5ML                        | DECADRON<br>(ELIXIR; ORAL)                       | MS&D/MERCK            | 12-376<br>09-02-60                     | 3375261<br>03-26-85<br>RE28369<br>03-26-85 |  |
| DEXAMETHASONE<br>0.5MG/5ML                        | HEXADROL<br>(ELIXIR; ORAL)                       | ORGANON/AKZONA        | 12-674<br>04-23-64                     | RE28369<br>03-26-85                        |  |
| DEXAMETHASONE<br>0.5MG                            | HEXADROL<br>(TABLET; ORAL)                       | ORGANON/AKZONA        | 12-675<br>07-01-78                     | RE28369<br>03-26-85                        |  |
| DEXAMETHASONE<br>0.75MG                           | HEXADROL<br>(TABLET; ORAL)                       | ORGANON/AKZONA        | 12-675<br>07-01-78                     | RE28369<br>03-26-85                        |  |

TABLE IV. NDA'S APPROVED FROM 1-1-82 TO 2-28-85 AND NDA'S WITH APPROPRIATE PATENT AND EXCLUSIVITY INFORMATION

| <u>ACTIVE INGREDIENT(S)<br/>STRENGTH(S)</u>          | <u>TRADE NAME<br/>(DOSAGE FORM; ROUTE)</u>  | <u>APPLICANT NAME</u> | <u>NDA NO.<br/>APPROVAL DATE</u> | <u>PATENT NO.<br/>EXP. DATE</u>            | <u>EXCLUSIVITY<br/>EXP. DATE</u> |
|--|---|-----------------------|----------------------------------|--|----------------------------------|
| DEXAMETHASONE<br>1.5MG                               | HEXADROL<br>(TABLET; ORAL)                  | ORGANON/AKZONA        | 12-675<br>09-24-65               | RE28369<br>03-26-85                        |                                  |
| DEXAMETHASONE<br>4MG                                 | HEXADROL<br>(TABLET; ORAL)                  | ORGANON/AKZONA        | 12-675<br>07-01-74               | RE28369<br>03-26-85                        |                                  |
| DEXAMETHASONE<br>10MG/25GH                           | DECASPRAY<br>(AEROSOL; TOPICAL)             | MS&D/MERCK            | 12-731<br>03-29-61               | 3375261<br>03-26-85<br>RE28369<br>03-26-85 |                                  |
| DEXAMETHASONE<br>0.04%                               | HEXADROL<br>(CREAM; TOPICAL)                | ORGANON/AKZONA        | 13-304<br>01-09-67               | RE28369<br>03-26-85                        |                                  |
| DEXAMETHASONE<br>0.1%                                | DECADERM<br>(GEL; TOPICAL)                  | MS&D/MERCK            | 13-538<br>05-03-65               | 3375261<br>03-26-85<br>RE28369<br>03-26-85 |                                  |
| DEXAMETHASONE ACETATE<br>EQ 8MG BASE/ML              | DECADRON-LA<br>(INJECTABLE; INJECTION)      | MS&D/MERCK            | 16-675<br>09-06-73               | 3375261<br>03-26-85<br>RE28369<br>03-26-85 |                                  |
| DEXAMETHASONE SODIUM PHOSPHATE<br>EQ 0.05% PHOSPHATE | DECADRON<br>(OINTMENT; OPHTHALMIC)          | MS&D/MERCK            | 11-977<br>09-02-59               | 3375261<br>03-26-85<br>RE28369<br>03-26-85 |                                  |
| DEXAMETHASONE SODIUM PHOSPHATE<br>EQ 0.1% PHOSPHATE  | DECADRON<br>(CREAM; TOPICAL)                | MS&D/MERCK            | 11-983<br>08-26-59               | 3375261<br>03-26-85<br>RE28369<br>03-26-85 |                                  |
| DEXAMETHASONE SODIUM PHOSPHATE<br>EQ 0.1% PHOSPHATE  | DECADRON<br>(SOLUTION;<br>OPHTHALMIC, OTIC) | MS&D/MERCK            | 11-984<br>09-23-59               | 3375261<br>03-26-85<br>RE28369<br>03-26-85 |                                  |

**TABLE IV. NDA'S APPROVED FROM 1-1-82 TO 2-28-85 AND NDA'S WITH APPROPRIATE PATENT AND EXCLUSIVITY INFORMATION**

| <u>ACTIVE INGREDIENT(S)</u><br><u>STRENGTH(S)</u>  | <u>TRADE NAME</u><br><u>(DOSAGE FORM; ROUTE)</u>                | <u>APPLICANT NAME</u> | <u>NDA NO.</u><br><u>APPROVAL DATE</u> | <u>PATENT NO.</u><br><u>EXP. DATE</u>      | <u>EXCLUSIVITY</u><br><u>EXP. DATE</u> |
|--|---|-----------------------|--|--|--|
| DEXAMETHASONE SODIUM PHOSPHATE<br>EQ 4MG PHOSPHATE/ML                                      | DECADRON<br>(INJECTABLE; INJECTION)                             | MS&D/MERCK            | 12-071<br>05-12-61                     | 3375261<br>03-26-85<br>RE28369<br>03-26-85 |  |
| DEXAMETHASONE SODIUM PHOSPHATE<br>EQ 24MG PHOSPHATE/ML                                     | DECADRON<br>(INJECTABLE; INJECTION)                             | MS&D/MERCK            | 12-071<br>03-01-77                     | 3375261<br>03-26-85<br>RE28369<br>03-26-85 |  |
| DEXAMETHASONE SODIUM PHOSPHATE<br>EQ 0.1MG PHOSPHATE/INH                                   | DECADRON<br>(AEROSOL; INHALATION)                               | MS&D/MERCK            | 13-413<br>09-17-62                     | 3375261<br>03-26-85<br>RE28369<br>03-26-85 |  |
| DEXAMETHASONE SODIUM PHOSPHATE<br>EQ 0.1MG PHOSPHATE/INH                                   | DECADRON<br>(AEROSOL; NASAL)                                    | MS&D/MERCK            | 14-242<br>12-17-65                     | 3375261<br>03-26-85<br>RE28369<br>03-26-85 |  |
| DEXAMETHASONE SODIUM PHOSPHATE<br>EQ 4MG PHOSPHATE/ML                                      | HEXADROL<br>(INJECTABLE; INJECTION)                             | ORGANON/AKZONA        | 14-694<br>03-14-75                     | RE28369<br>03-26-85                        |  |
| DEXAMETHASONE SODIUM PHOSPHATE<br>EQ 10MG PHOSPHATE/ML                                     | HEXADROL<br>(INJECTABLE; INJECTION)                             | ORGANON/AKZONA        | 14-694<br>03-14-75                     | RE28369<br>03-26-85                        |  |
| DEXAMETHASONE SODIUM PHOSPHATE<br>EQ 20MG PHOSPHATE/ML                                     | HEXADROL<br>(INJECTABLE; INJECTION)                             | ORGANON/AKZONA        | 14-694<br>04-27-81                     | RE28369<br>03-26-85                        |  |
| DEXAMETHASONE SODIUM PHOSPHATE;<br>LIDOCAINE HYDROCHLORIDE<br>EQ 4MG PHOSPHATE/ML; 10MG/ML | DECADRON W/ XYLOCAINE<br>(INJECTABLE; INJECTION)                | MS&D/MERCK            | 13-334<br>07-11-62                     | 3375261<br>03-26-85<br>RE28369<br>03-26-85 |  |
| DEXTROMETHORPHAN HYDROBROMIDE;<br>PROMETHAZINE HYDROCHLORIDE<br>15MG/5ML; 6.25MG/5ML       | PHENERGAN W/ DEXTROMETHORPHAN<br>(SYRUP; ORAL)                  | WYETH LABS/AMHO       | 11-265<br>04-02-84                     |  |  |
| DEXTROSE<br>60GM/100ML   | DEXTROSE 60% IN PLASTIC<br>CONTAINER<br>(INJECTABLE; INJECTION) | TRAVENOL LABS         | 17-521<br>03-26-82                     |  |  |

**TABLE IV. NDA'S APPROVED FROM 1-1-82 TO 2-28-85 AND NDA'S WITH APPROPRIATE PATENT AND EXCLUSIVITY INFORMATION**

| <u>ACTIVE INGREDIENT(S)</u><br><u>STRENGTH(S)</u> | <u>TRADE NAME</u><br><u>(DOSAGE FORM; ROUTE)</u>                  | <u>APPLICANT NAME</u> | <u>NDA NO.</u><br><u>APPROVAL DATE</u> | <u>PATENT NO.</u><br><u>EXP. DATE</u> | <u>EXCLUSIVITY</u><br><u>EXP. DATE</u> |
|---|---|-----------------------|--|---------------------------------------|--|
| DEXTROSE<br>70GM/100ML                            | DEXTROSE 70% IN PLASTIC<br>CONTAINER<br>(INJECTABLE; INJECTION)   | TRAVENOL LABS         | 17-521<br>03-26-82                     |                                       |  |
| DEXTROSE<br>60GM/100ML                            | DEXTROSE 60%<br>IN PLASTIC CONTAINER<br>(INJECTABLE; INJECTION)   | ABBOTT LABORATORIES   | 19-346<br>01-25-85                     |                                       |  |
| DEXTROSE<br>30GM/100ML                            | DEXTROSE 30%<br>IN PLASTIC CONTAINER<br>(INJECTABLE; INJECTION)   | ABBOTT LABORATORIES   | 19-345<br>01-26-85                     |                                       |  |
| DEXTROSE<br>60GM/100ML                            | DEXTROSE 60% IN PLASTIC<br>CONTAINER<br>(INJECTABLE; INJECTION)   | AM MCGAW/AM HOSP      | 17-995<br>04-27-78                     | 3729568<br>04-24-90                   |  |
| DEXTROSE<br>60GM/100ML                            | DEXTROSE 60%<br>(INJECTABLE; INJECTION)                           | AM MCGAW/AM HOSP      | 17-995<br>09-22-82                     | 3729568<br>04-24-90                   |  |
| DEXTROSE<br>70GM/100ML                            | DEXTROSE 70% IN PLASTIC<br>CONTAINER<br>(INJECTABLE; INJECTION)   | ABBOTT LABORATORIES   | 18-561<br>03-23-82                     |                                       |  |
| DEXTROSE<br>40GM/100ML                            | DEXTROSE 40% IN PLASTIC<br>CONTAINER<br>(INJECTABLE; INJECTION)   | ABBOTT LABORATORIES   | 18-562<br>03-23-82                     |                                       |  |
| DEXTROSE<br>50GM/100ML                            | DEXTROSE 50% IN PLASTIC<br>CONTAINER<br>(INJECTABLE; INJECTION)   | ABBOTT LABORATORIES   | 18-563<br>03-23-82                     |                                       |  |
| DEXTROSE<br>20GM/100ML                            | DEXTROSE 20% IN PLASTIC<br>CONTAINER<br>(INJECTABLE; INJECTION)   | ABBOTT LABORATORIES   | 18-564<br>03-23-82                     |                                       |  |
| DEXTROSE<br>38.5GM/100ML                          | DEXTROSE 38.5% IN PLASTIC<br>CONTAINER<br>(INJECTABLE; INJECTION) | ABBOTT LABORATORIES   | 18-923<br>09-19-84                     |                                       |  |

**TABLE IV. NDA'S APPROVED FROM 1-1-82 TO 2-28-85 AND NDA'S WITH APPROPRIATE PATENT AND EXCLUSIVITY INFORMATION**

| <u>ACTIVE INGREDIENT(S)<br/>STRENGTH(S)</u>                | <u>TRADE NAME<br/>(DOSAGE FORM; ROUTE)</u>   | <u>APPLICANT NAME</u> | <u>NDA NO.<br/>APPROVAL DATE</u> | <u>PATENT NO.<br/>EXP. DATE</u> | <u>EXCLUSIVITY<br/>EXP. DATE</u> |
|--|--|-----------------------|----------------------------------|---------------------------------|----------------------------------|
| DEXTROSE<br>50MG/ML  | DEXTROSE 5% IN PLASTIC<br>CONTAINER<br>(INJECTABLE; INJECTION)                                   | ABBOTT LABORATORIES   | 19-222<br>07-13-84               |                                 |                                  |
| DEXTROSE; DOPAMINE HYDROCHLORIDE<br>5GM/100ML; 80MG/100ML  | DOPAMINE HCL<br>(INJECTABLE; INJECTION)  | ABBOTT LABORATORIES   | 18-132<br>02-04-82               |                                 | NC<br>09-24-86                   |
| DEXTROSE; DOPAMINE HYDROCHLORIDE<br>5GM/100ML; 160MG/100ML | DOPAMINE HCL<br>(INJECTABLE; INJECTION)  | ABBOTT LABORATORIES   | 18-132<br>02-04-82               |                                 | NC<br>09-24-86                   |
| DEXTROSE; DOPAMINE HYDROCHLORIDE<br>5GM/100ML; 80MG/100ML  | DOPAMINE HCL IN PLASTIC<br>CONTAINER<br>(INJECTABLE; INJECTION)                                  | ABBOTT LABORATORIES   | 18-826<br>09-30-83               |                                 | NC<br>09-24-86                   |
| DEXTROSE; DOPAMINE HYDROCHLORIDE<br>5GM/100ML; 160MG/100ML | DOPAMINE HCL IN PLASTIC<br>CONTAINER<br>(INJECTABLE; INJECTION)                                  | ABBOTT LABORATORIES   | 18-826<br>09-30-83               |                                 | NC<br>09-24-86                   |
| DEXTROSE; DOPAMINE HYDROCHLORIDE<br>5GM/100ML; 320MG/100ML | DOPAMINE HCL IN PLASTIC<br>CONTAINER<br>(INJECTABLE; INJECTION)                                  | ABBOTT LABORATORIES   | 18-826<br>09-30-83               |                                 | NC<br>09-24-86                   |
| DEXTROSE; HEPARIN SODIUM<br>5GM/100ML; 200 UNITS/100ML     | HEPARIN SODIUM 1,000 UNITS<br>AND DEXTROSE 5% IN<br>PLASTIC CONTAINER<br>(INJECTABLE; INJECTION) | AM MCGAM/AM HOSP      | 19-130<br>12-31-83               |                                 | NC<br>09-24-86                   |
| DEXTROSE; HEPARIN SODIUM<br>5GM/100ML; 200 UNITS/100ML     | HEPARIN SODIUM 2,000 UNITS<br>AND DEXTROSE 5% IN<br>PLASTIC CONTAINER<br>(INJECTABLE; INJECTION) | AM MCGAM/AM HOSP      | 19-130<br>12-31-83               |                                 | NC<br>09-24-86                   |
| DEXTROSE; HEPARIN SODIUM<br>5GM/100ML; 1,000 UNITS/100ML   | HEPARIN SODIUM 5,000 UNITS<br>AND DEXTROSE 5% IN<br>PLASTIC CONTAINER<br>(INJECTABLE; INJECTION) | AM MCGAM/AM HOSP      | 19-130<br>12-31-83               |                                 | NC<br>09-24-86                   |

TABLE IV. NDA'S APPROVED FROM 1-1-82 TO 2-28-85 AND NDA'S WITH APPROPRIATE PATENT AND EXCLUSIVITY INFORMATION

| <u>ACTIVE INGREDIENT(S)</u><br><u>STRENGTH(S)</u>           | <u>TRADE NAME</u><br><u>(DOSAGE FORM; ROUTE)</u>  | <u>APPLICANT NAME</u> | <u>NDA NO.</u><br><u>APPROVAL DATE</u> | <u>PATENT NO.</u><br><u>EXP. DATE</u> | <u>EXCLUSIVITY</u><br><u>EXP. DATE</u> |
|---|---|-----------------------|--|---------------------------------------|--|
| DEXTROSE; HEPARIN SODIUM<br>5GM/100ML; 4,000 UNITS/100ML    | HEPARIN SODIUM 20,000 UNITS<br>AND DEXTROSE 5% IN<br>PLASTIC CONTAINER<br>(INJECTABLE; INJECTION) | TRAVENOL LABS         | 18-814<br>10-31-83                     |                                       | NC<br>09-24-86                         |
| DEXTROSE; HEPARIN SODIUM<br>5GM/100ML; 5,000 UNITS/100ML    | HEPARIN SODIUM<br>12,500 UNITS<br>IN DEXTROSE 5%<br>(INJECTABLE; INJECTION)                       | ABBOTT LABORATORIES   | 18-911<br>01-30-85                     |                                       |  |
| DEXTROSE; HEPARIN SODIUM<br>5GM/100ML; 10,000 UNITS/100ML   | HEPARIN SODIUM<br>10,000 UNITS<br>IN DEXTROSE 5%<br>(INJECTABLE; INJECTION)                       | ABBOTT LABORATORIES   | 18-911<br>01-30-85                     |                                       |  |
| DEXTROSE; HEPARIN SODIUM<br>5GM/100ML; 10,000 UNITS/100ML   | HEPARIN SODIUM<br>25,000 UNITS<br>IN DEXTROSE 5%<br>(INJECTABLE; INJECTION)                       | ABBOTT LABORATORIES   | 18-911<br>01-30-85                     |                                       |  |
| DEXTROSE; LIDOCAINE HYDROCHLORIDE<br>5GM/100ML; 800MG/100ML | LIDOCAINE HCL 0.8%<br>IN DEXTROSE 5%<br>IN PLASTIC CONTAINER<br>(INJECTABLE; INJECTION)           | ABBOTT LABORATORIES   | 18-388<br>11-05-82                     |                                       | NS<br>09-24-86                         |
| DEXTROSE; LIDOCAINE HYDROCHLORIDE<br>5GM/100ML; 800MG/100ML | LIDOCAINE HCL 0.8%<br>AND DEXTROSE 5%<br>IN PLASTIC CONTAINER<br>(INJECTABLE; INJECTION)          | TRAVENOL LABS         | 18-461<br>02-22-82                     |                                       | NS<br>09-24-86                         |
| DEXTROSE; LIDOCAINE HYDROCHLORIDE<br>5GM/100ML; 200MG/100ML | LIDOCAINE HCL 0.2%<br>AND DEXTROSE 5%<br>IN PLASTIC CONTAINER<br>(INJECTABLE; INJECTION)          | AM MCGAW/AM HOSP      | 18-967<br>03-30-84                     |                                       | NS<br>09-24-86                         |
| DEXTROSE; LIDOCAINE HYDROCHLORIDE<br>5GM/100ML; 400MG/100ML | LIDOCAINE HCL 0.4%<br>AND DEXTROSE 5%<br>IN PLASTIC CONTAINER<br>(INJECTABLE; INJECTION)          | AM MCGAW/AM HOSP      | 18-967<br>03-30-84                     |                                       | NS<br>09-24-86                         |

TABLE IV. NDA'S APPROVED FROM 1-1-82 TO 2-28-85 AND NDA'S WITH APPROPRIATE PATENT AND EXCLUSIVITY INFORMATION

| <u>ACTIVE INGREDIENT(S)<br/>STRENGTH(S)</u>   | <u>TRADE NAME<br/>(DOSAGE FORM; ROUTE)</u>  | <u>APPLICANT NAME</u> | <u>NDA NO.<br/>APPROVAL DATE</u> | <u>PATENT NO.<br/>EXP. DATE</u> | <u>EXCLUSIVITY<br/>EXP. DATE</u> |
|---|---|-----------------------|----------------------------------|---------------------------------|----------------------------------|
| DEXTROSE; LIDOCAINE HYDROCHLORIDE<br>5GM/100ML; 800MG/100ML   | LIDOCAINE HCL 0.8%<br>AND DEXTROSE 5%<br>IN PLASTIC CONTAINER<br>(INJECTABLE; INJECTION)                              | AM MCGAW/AM HOSP      | 18-967<br>03-30-84               |                                 | NS<br>09-24-86                   |
| DEXTROSE; MAGNESIUM CHLORIDE;<br>POTASSIUM CHLORIDE;<br>POTASSIUM PHOSPHATE DIBASIC;<br>SODIUM ACETATE<br>5GM/100ML; 31MG/100ML;<br>130MG/100ML; 26MG/100ML;<br>320MG/100ML | ISOLYTE P W/<br>DEXTROSE 5% IN<br>PLASTIC CONTAINER<br>(INJECTABLE; INJECTION)  | AM MCGAW/AM HOSP      | 19-025<br>12-27-84               |                                 |                                  |
| DEXTROSE; POTASSIUM CHLORIDE<br>5GM/100ML; 75MG/100ML   | DEXTROSE 5% AND<br>POTASSIUM CHLORIDE 0.075%<br>IN PLASTIC CONTAINER<br>(INJECTABLE; INJECTION)                       | AM MCGAW/AM HOSP      | 18-744<br>11-09-82               |                                 |                                  |
| DEXTROSE; POTASSIUM CHLORIDE<br>5GM/100ML; 150MG/100ML  | DEXTROSE 5% AND<br>POTASSIUM CHLORIDE 0.15%<br>IN PLASTIC CONTAINER<br>(INJECTABLE; INJECTION)                        | AM MCGAW/AM HOSP      | 18-744<br>11-09-82               |                                 |                                  |
| DEXTROSE; POTASSIUM CHLORIDE<br>5GM/100ML; 220MG/100ML  | DEXTROSE 5% AND<br>POTASSIUM CHLORIDE 0.22%<br>IN PLASTIC CONTAINER<br>(INJECTABLE; INJECTION)                        | AM MCGAW/AM HOSP      | 18-744<br>11-09-82               |                                 |                                  |
| DEXTROSE; POTASSIUM CHLORIDE<br>5GM/100ML; 300MG/100ML  | DEXTROSE 5% AND<br>POTASSIUM CHLORIDE 0.3%<br>IN PLASTIC CONTAINER<br>(INJECTABLE; INJECTION)                         | AM MCGAW/AM HOSP      | 18-744<br>11-09-82               |                                 |                                  |
| DEXTROSE; POTASSIUM CHLORIDE; POTASSIUM<br>PHOSPHATE, MONOBASIC; SODIUM CHLORIDE;<br>SODIUM LACTATE<br>5GM/100ML; 205MG/100ML; 100MG/100ML;<br>120MG/100ML; 220MG/100ML     | DEXTROSE 5% AND ELECTROLYTE<br>NO 75 IN PLASTIC CONTAINER<br>(INJECTABLE; INJECTION)                                  | TRAVENOL LABS         | 18-840<br>06-29-83               |                                 |                                  |
| DEXTROSE; POTASSIUM CHLORIDE; SODIUM<br>CHLORIDE<br>5GM/100ML; 150MG/100ML; 450MG/100ML   | DEXTROSE 5%, SODIUM CHLORIDE<br>0.45% AND POTASSIUM CHLORIDE<br>10MEQ IN PLASTIC CONTAINER<br>(INJECTABLE; INJECTION) | TRAVENOL LABS         | 18-566<br>02-10-83               |                                 |                                  |

TABLE IV. NDA'S APPROVED FROM 1-1-82 TO 2-28-85 AND NDA'S WITH APPROPRIATE PATENT AND EXCLUSIVITY INFORMATION

| <u>ACTIVE INGREDIENT(S)<br/>STRENGTH(S)</u>  | <u>TRADE NAME<br/>(DOSAGE FORM; ROUTE)</u>   | <u>APPLICANT NAME</u> | <u>NDA NO.<br/>APPROVAL DATE</u> | <u>PATENT NO.<br/>EXP. DATE</u> | <u>EXCLUSIVITY<br/>EXP. DATE</u> |
|--|--|-----------------------|----------------------------------|---------------------------------|----------------------------------|
| DEXTROSE; POTASSIUM CHLORIDE; SODIUM CHLORIDE<br>5GM/100ML; 224MG/100ML; 450MG/100ML | DEXTROSE 5%, SODIUM CHLORIDE 0.45% AND POTASSIUM CHLORIDE 15MEQ IN PLASTIC CONTAINER (INJECTABLE; INJECTION) | TRAVENOL LABS         | 18-566<br>02-10-83               |                                 |                                  |
| DEXTROSE; POTASSIUM CHLORIDE; SODIUM CHLORIDE<br>5GM/100ML; 300MG/100ML; 450MG/100ML | DEXTROSE 5%, SODIUM CHLORIDE 0.45% AND POTASSIUM CHLORIDE 20MEQ IN PLASTIC CONTAINER (INJECTABLE; INJECTION) | TRAVENOL LABS         | 18-566<br>02-10-83               |                                 |                                  |
| DEXTROSE; POTASSIUM CHLORIDE; SODIUM CHLORIDE<br>5GM/100ML; 150MG/100ML; 450MG/100ML | DEXTROSE 5%, SODIUM CHLORIDE 0.45% AND POTASSIUM CHLORIDE 20MEQ IN PLASTIC CONTAINER (INJECTABLE; INJECTION) | TRAVENOL LABS         | 18-566<br>02-10-83               |                                 |                                  |
| DEXTROSE; POTASSIUM CHLORIDE; SODIUM CHLORIDE<br>5GM/100ML; 224MG/100ML; 450MG/100ML | DEXTROSE 5%, SODIUM CHLORIDE 0.45% AND POTASSIUM CHLORIDE 30MEQ IN PLASTIC CONTAINER (INJECTABLE; INJECTION) | TRAVENOL LABS         | 18-566<br>02-10-83               |                                 |                                  |
| DEXTROSE; POTASSIUM CHLORIDE; SODIUM CHLORIDE<br>5GM/100ML; 300MG/100ML; 450MG/100ML | DEXTROSE 5%, SODIUM CHLORIDE 0.45% AND POTASSIUM CHLORIDE 40MEQ IN PLASTIC CONTAINER (INJECTABLE; INJECTION) | TRAVENOL LABS         | 18-566<br>02-10-83               |                                 |                                  |
| DEXTROSE; POTASSIUM CHLORIDE; SODIUM CHLORIDE<br>5GM/100ML; 150MG/100ML; 200MG/100ML | DEXTROSE 5%, SODIUM CHLORIDE 0.2% AND POTASSIUM CHLORIDE 10MEQ IN PLASTIC CONTAINER (INJECTABLE; INJECTION)  | TRAVENOL LABS         | 18-567<br>02-16-83               |                                 |                                  |
| DEXTROSE; POTASSIUM CHLORIDE; SODIUM CHLORIDE<br>5GM/100ML; 224MG/100ML; 200MG/100ML | DEXTROSE 5%, SODIUM CHLORIDE 0.2% AND POTASSIUM CHLORIDE 15MEQ IN PLASTIC CONTAINER (INJECTABLE; INJECTION)  | TRAVENOL LABS         | 18-567<br>02-16-83               |                                 |                                  |
| DEXTROSE; POTASSIUM CHLORIDE; SODIUM CHLORIDE<br>5GM/100ML; 150MG/100ML; 200MG/100ML | DEXTROSE 5%, SODIUM CHLORIDE 0.2% AND POTASSIUM CHLORIDE 20MEQ IN PLASTIC CONTAINER (INJECTABLE; INJECTION)  | TRAVENOL LABS         | 18-567<br>02-16-83               |                                 |                                  |

**TABLE IV. NDA'S APPROVED FROM 1-1-82 TO 2-28-85 AND NDA'S WITH APPROPRIATE PATENT AND EXCLUSIVITY INFORMATION**

| <u>ACTIVE INGREDIENT(S)<br/>STRENGTH(S)</u>  | <u>TRADE NAME<br/>(DOSAGE FORM; ROUTE)</u>   | <u>APPLICANT NAME</u> | <u>NDA NO.<br/>APPROVAL DATE</u> | <u>PATENT NO.<br/>EXP. DATE</u> | <u>EXCLUSIVITY<br/>EXP. DATE</u> |
|--|--|-----------------------|----------------------------------|---------------------------------|----------------------------------|
| DEXTROSE; POTASSIUM CHLORIDE; SODIUM CHLORIDE<br>5GM/100ML; 224MG/100ML; 200MG/100ML | DEXTROSE 5%, SODIUM CHLORIDE 0.2% AND POTASSIUM CHLORIDE 30MEQ IN PLASTIC CONTAINER (INJECTABLE; INJECTION)  | TRAVENOL LABS         | 18-567<br>02-16-83               |                                 |                                  |
| DEXTROSE; POTASSIUM CHLORIDE; SODIUM CHLORIDE<br>5GM/100ML; 75MG/100ML; 330MG/100ML  | DEXTROSE 5%, SODIUM CHLORIDE 0.33% AND POTASSIUM CHLORIDE 5MEQ IN PLASTIC CONTAINER (INJECTABLE; INJECTION)  | TRAVENOL LABS         | 18-629<br>03-23-82               |                                 |                                  |
| DEXTROSE; POTASSIUM CHLORIDE; SODIUM CHLORIDE<br>5GM/100ML; 150MG/100ML; 330MG/100ML | DEXTROSE 5%, SODIUM CHLORIDE 0.33% AND POTASSIUM CHLORIDE 10MEQ IN PLASTIC CONTAINER (INJECTABLE; INJECTION) | TRAVENOL LABS         | 18-629<br>03-23-82               |                                 |                                  |
| DEXTROSE; POTASSIUM CHLORIDE; SODIUM CHLORIDE<br>5GM/100ML; 224MG/100ML; 330MG/100ML | DEXTROSE 5%, SODIUM CHLORIDE 0.33% AND POTASSIUM CHLORIDE 15MEQ IN PLASTIC CONTAINER (INJECTABLE; INJECTION) | TRAVENOL LABS         | 18-629<br>03-23-82               |                                 |                                  |
| DEXTROSE; POTASSIUM CHLORIDE; SODIUM CHLORIDE<br>5GM/100ML; 150MG/100ML; 330MG/100ML | DEXTROSE 5%, SODIUM CHLORIDE 0.33% AND POTASSIUM CHLORIDE 20MEQ IN PLASTIC CONTAINER (INJECTABLE; INJECTION) | TRAVENOL LABS         | 18-629<br>03-23-82               |                                 |                                  |
| DEXTROSE; POTASSIUM CHLORIDE; SODIUM CHLORIDE<br>5GM/100ML; 75MG/100ML; 330MG/100ML  | DEXTROSE 5%, SODIUM CHLORIDE 0.33% AND POTASSIUM CHLORIDE 10MEQ IN PLASTIC CONTAINER (INJECTABLE; INJECTION) | TRAVENOL LABS         | 18-629<br>03-23-82               |                                 |                                  |
| DEXTROSE; POTASSIUM CHLORIDE; SODIUM CHLORIDE<br>5GM/100ML; 300MG/100ML; 330MG/100ML | DEXTROSE 5%, SODIUM CHLORIDE 0.33% AND POTASSIUM CHLORIDE 20MEQ IN PLASTIC CONTAINER (INJECTABLE; INJECTION) | TRAVENOL LABS         | 18-629<br>03-23-82               |                                 |                                  |
| DEXTROSE; POTASSIUM CHLORIDE; SODIUM CHLORIDE<br>5GM/100ML; 224MG/100ML; 330MG/100ML | DEXTROSE 5%, SODIUM CHLORIDE 0.33% AND POTASSIUM CHLORIDE 30MEQ IN PLASTIC CONTAINER (INJECTABLE; INJECTION) | TRAVENOL LABS         | 18-629<br>03-23-82               |                                 |                                  |

TABLE IV. NDA'S APPROVED FROM 1-1-82 TO 2-28-85 AND NDA'S WITH APPROPRIATE PATENT AND EXCLUSIVITY INFORMATION

| <u>ACTIVE INGREDIENT(S)<br/>STRENGTH(S)</u>  | <u>TRADE NAME<br/>(DOSAGE FORM; ROUTE)</u>   | <u>APPLICANT NAME</u> | <u>NDA NO.<br/>APPROVAL DATE</u> | <u>PATENT NO.<br/>EXP. DATE</u> | <u>EXCLUSIVITY<br/>EXP. DATE</u> |
|--|--|-----------------------|----------------------------------|---------------------------------|----------------------------------|
| DEXTROSE; POTASSIUM CHLORIDE; SODIUM CHLORIDE<br>5GM/100ML; 300MG/100ML; 330MG/100ML | DEXTROSE 5%, SODIUM CHLORIDE 0.33% AND POTASSIUM CHLORIDE 40MEQ IN PLASTIC CONTAINER (INJECTABLE; INJECTION) | TRAVENOL LABS         | 18-629<br>03-23-82               |                                 |                                  |
| DEXTROSE; THEOPHYLLINE<br>5GM/100ML; 40MG/100ML                                      | THEOPHYLLINE IN DEXTROSE 5% IN PLASTIC CONTAINER (INJECTABLE; INJECTION)                                     | ABBOTT LABORATORIES   | 19-211<br>12-14-84               |                                 |                                  |
| DEXTROSE; THEOPHYLLINE<br>5GM/100ML; 40MG/100ML                                      | THEOPHYLLINE 0.04% AND DEXTROSE 5% IN PLASTIC CONTAINER (INJECTABLE; INJECTION)                              | AM MCGAW/AM HOSP      | 19-083<br>11-07-84               |                                 |                                  |
| DEXTROSE; THEOPHYLLINE<br>5GM/100ML; 80MG/100ML                                      | THEOPHYLLINE IN DEXTROSE 5% IN PLASTIC CONTAINER (INJECTABLE; INJECTION)                                     | ABBOTT LABORATORIES   | 19-211<br>12-14-84               |                                 |                                  |
| DEXTROSE; THEOPHYLLINE<br>5GM/100ML; 80MG/100ML                                      | THEOPHYLLINE 0.08% AND DEXTROSE 5% IN PLASTIC CONTAINER (INJECTABLE; INJECTION)                              | AM MCGAW/AM HOSP      | 19-083<br>11-07-84               |                                 |                                  |
| DEXTROSE; THEOPHYLLINE<br>5GM/100ML; 160MG/100ML                                     | THEOPHYLLINE IN DEXTROSE 5% IN PLASTIC CONTAINER (INJECTABLE; INJECTION)                                     | ABBOTT LABORATORIES   | 19-211<br>12-14-84               |                                 |                                  |
| DEXTROSE; THEOPHYLLINE<br>5GM/100ML; 160MG/100ML                                     | THEOPHYLLINE 0.16% AND DEXTROSE 5% IN PLASTIC CONTAINER (INJECTABLE; INJECTION)                              | AM MCGAW/AM HOSP      | 19-083<br>11-07-84               |                                 |                                  |
| DEXTROSE; THEOPHYLLINE<br>5GM/100ML; 200MG/100ML                                     | THEOPHYLLINE IN DEXTROSE 5% IN PLASTIC CONTAINER (INJECTABLE; INJECTION)                                     | ABBOTT LABORATORIES   | 19-211<br>12-14-84               |                                 |                                  |

TABLE IV. NDA'S APPROVED FROM 1-1-82 TO 2-28-85 AND NDA'S WITH APPROPRIATE PATENT AND EXCLUSIVITY INFORMATION

| <u>ACTIVE INGREDIENT(S)<br/>STRENGTH(S)</u>      | <u>TRADE NAME<br/>(DOSAGE FORM; ROUTE)</u>  | <u>APPLICANT NAME</u> | <u>NDA NO.<br/>APPROVAL DATE</u> | <u>PATENT NO.<br/>EXP. DATE</u> | <u>EXCLUSIVITY<br/>EXP. DATE</u> |
|--|---|-----------------------|----------------------------------|---------------------------------|----------------------------------|
| DEXTROSE; THEOPHYLLINE<br>5GM/100ML; 200MG/100ML | THEOPHYLLINE 0.2%<br>AND DEXTROSE 5%<br>IN PLASTIC CONTAINER<br>(INJECTABLE; INJECTION) | AM MCGAW/AM HOSP      | 19-212<br>11-07-84               |                                 |                                  |
| DEXTROSE; THEOPHYLLINE<br>5GM/100ML; 400MG/100ML | THEOPHYLLINE IN<br>DEXTROSE 5%<br>IN PLASTIC CONTAINER<br>(INJECTABLE; INJECTION)       | ABBOTT LABORATORIES   | 19-211<br>12-14-84               |                                 |                                  |
| DEXTROSE; THEOPHYLLINE<br>5GM/100ML; 400MG/100ML | THEOPHYLLINE 0.4%<br>AND DEXTROSE 5%<br>IN PLASTIC CONTAINER<br>(INJECTABLE; INJECTION) | AM MCGAW/AM HOSP      | 19-212<br>11-07-84               |                                 |                                  |
| DEXTROSE; THEOPHYLLINE<br>5GM/100ML; 400MG/100ML | THEOPHYLLINE AND DEXTROSE 5%<br>IN PLASTIC CONTAINER<br>(INJECTABLE; INJECTION)         | TRAVENOL LABS         | 18-649<br>07-26-82               |                                 |                                  |
| DEXTROSE; THEOPHYLLINE<br>5GM/100ML; 80MG/100ML  | THEOPHYLLINE AND DEXTROSE 5%<br>IN PLASTIC CONTAINER<br>(INJECTABLE; INJECTION)         | TRAVENOL LABS         | 18-649<br>07-26-82               |                                 |                                  |
| DEXTROSE; THEOPHYLLINE<br>5GM/100ML; 160MG/100ML | THEOPHYLLINE AND DEXTROSE 5%<br>IN PLASTIC CONTAINER<br>(INJECTABLE; INJECTION)         | TRAVENOL LABS         | 18-649<br>07-26-82               |                                 |                                  |
| DEXTROSE; THEOPHYLLINE<br>5GM/100ML; 200MG/100ML | THEOPHYLLINE AND DEXTROSE 5%<br>IN PLASTIC CONTAINER<br>(INJECTABLE; INJECTION)         | TRAVENOL LABS         | 18-649<br>07-26-82               |                                 |                                  |
| DEXTROSE; THEOPHYLLINE<br>5GM/100ML; 400MG/100ML | THEOPHYLLINE AND DEXTROSE 5%<br>IN PLASTIC CONTAINER<br>(INJECTABLE; INJECTION)         | TRAVENOL LABS         | 18-649<br>07-26-82               |                                 |                                  |

**TABLE IV. NDA'S APPROVED FROM 1-1-82 TO 2-28-85 AND NDA'S WITH APPROPRIATE PATENT AND EXCLUSIVITY INFORMATION**

| <u>ACTIVE INGREDIENT(S)<br/>STRENGTH(S)</u>              | <u>TRADE NAME<br/>(DOSAGE FORM; ROUTE)</u>            | <u>APPLICANT NAME</u> | <u>NDA NO.<br/>APPROVAL DATE</u> | <u>PATENT NO.<br/>EXP. DATE</u> | <u>EXCLUSIVITY<br/>EXP. DATE</u> |
|--|---|-----------------------|----------------------------------|---------------------------------|----------------------------------|
| DIATRIZOATE<br>MEGLUMINE<br>30%                          | RENO-M-DIP<br>(INJECTABLE; INJECTION)                 | ER SQUIBB AND SONS    | 10-040<br>01-08-60               |                                 | I-7; I-8<br>09-24-86             |
| DIATRIZOATE MEGLUMINE;<br>DIATRIZOATE SODIUM<br>52%; 8%  | RENOGRAFIN-60<br>(INJECTABLE; INJECTION)              | ER SQUIBB AND SONS    | 10-040<br>08-29-74               |                                 | I-8<br>09-24-86                  |
| DIATRIZOATE MEGLUMINE;<br>DIATRIZOATE SODIUM<br>66%; 10% | RENOGRAFIN-76<br>(INJECTABLE; INJECTION)              | ER SQUIBB AND SONS    | 10-040<br>10-27-72               |                                 | I-5<br>09-24-86                  |
| DIAZEPAM<br>2MG  | VALIUM<br>(TABLET; ORAL)                              | HOFFMANN-LA ROCHE     | 13-263<br>11-15-63               | 4316897<br>02-23-99             |                                  |
| DIAZEPAM<br>5MG  | VALIUM<br>(TABLET; ORAL)                              | HOFFMANN-LA ROCHE     | 13-263<br>11-15-63               | 4316897<br>02-23-99             |                                  |
| DIAZEPAM<br>10MG   | VALIUM<br>(TABLET; ORAL)                              | HOFFMANN-LA ROCHE     | 13-263<br>11-15-63               | 4316897<br>02-23-99             |                                  |
| DIAZEPAM<br>5MG/ML                                       | VALIUM<br>(INJECTABLE; INJECTION)                     | HOFFMANN-LA ROCHE     | 16-087<br>08-24-66               | 4316897<br>02-23-99             |                                  |
| DIAZEPAM<br>15MG   | VAL RELEASE<br>(CAPSULE; CONTROLLED<br>RELEASE; ORAL) | HOFFMANN-LA ROCHE     | 18-179<br>03-12-81               | 4316897<br>02-23-99             |                                  |
| DIAZOXIDE<br>15MG/ML                                     | HYPERSTAT<br>(INJECTABLE; INJECTION)                  | SCHERING              | 16-996<br>01-22-73               |                                 | I-1<br>09-24-86                  |
| DICYCLOMINE HYDROCHLORIDE<br>10MG                        | BENTYL<br>(CAPSULE; ORAL)                             | MERRELL DOW/DOW CHEM  | 07-409<br>10-15-84               |                                 |                                  |
| DICYCLOMINE HYDROCHLORIDE<br>20MG                        | BENTYL<br>(CAPSULE; ORAL)                             | MERRELL DOW/DOW CHEM  | 07-409<br>10-15-84               |                                 |                                  |
| DICYCLOMINE HYDROCHLORIDE<br>10MG/ML                     | BENTYL<br>(INJECTABLE; INJECTION)                     | MERRELL DOW/DOW CHEM  | 08-370<br>10-15-84               |                                 |                                  |

**TABLE IV. NDA'S APPROVED FROM 1-1-82 TO 2-28-85 AND NDA'S WITH APPROPRIATE PATENT AND EXCLUSIVITY INFORMATION**

| <u>ACTIVE INGREDIENT(S)<br/>STRENGTH(S)</u>   | <u>TRADE NAME<br/>(DOSAGE FORM; ROUTE)</u> | <u>APPLICANT NAME</u> | <u>NDA NO.<br/>APPROVAL DATE</u> | <u>PATENT NO.<br/>EXP. DATE</u>            | <u>EXCLUSIVITY<br/>EXP. DATE</u> |
|---|--|-----------------------|----------------------------------|--|----------------------------------|
| DICYCLOMINE HYDROCHLORIDE<br>10MG/5ML   | BENTYL<br>(SYRUP; ORAL)                    | MERRELL DOM/DOW CHEM  | 07-961<br>10-15-84               |  |                                  |
| DIFLORASONE DIACETATE<br>0.05%  | FLORONE<br>(CREAM; TOPICAL)                | UPJOHN                | 17-741<br>09-14-77               | 3980778<br>09-14-93                        |                                  |
| DIFLORASONE DIACETATE<br>0.05%  | FLORONE<br>(OINTMENT; TOPICAL)             | UPJOHN                | 17-994<br>03-01-78               | 3980778<br>09-14-93                        |                                  |
| DIFLUNISAL<br>250MG   | DOLOBID<br>(TABLET; ORAL)                  | MS&D/MERCK            | 18-445<br>04-19-82               | 3714226<br>08-01-89<br>3674870<br>07-04-89 | NCE<br>04-19-92                  |
| DIFLUNISAL<br>500MG   | DOLOBID<br>(TABLET; ORAL)                  | MS&D/MERCK            | 18-445<br>04-19-82               | 3714226<br>08-01-89<br>3674870<br>07-04-89 | NCE<br>04-19-92                  |
| DIGOXIN<br>0.2MG  | LANOXICAPS<br>(CAPSULE; ORAL)              | BURROUGHS WELLCOME    | 18-118<br>07-26-82               |  | NDF<br>09-24-86                  |
| DIGOXIN<br>0.05MG   | LANOXICAPS<br>(CAPSULE; ORAL)              | BURROUGHS WELLCOME    | 18-118<br>07-26-82               |  | NDF<br>09-24-86                  |
| DIGOXIN<br>0.1MG  | LANOXICAPS<br>(CAPSULE; ORAL)              | BURROUGHS WELLCOME    | 18-118<br>07-26-82               |  | NDF<br>09-24-86                  |
| DHYDROERGOTAMINE MESYLATE;<br>HEPARIN SODIUM; LIDOCAINE HYDROCHLORIDE<br>0.5MG/0.5ML; 2500 UNITS/0.5ML;<br>5.33MG/0.5ML | EMBOLEX<br>(INJECTABLE; INJECTION)         | SANDOZ PHARMS/SANDOZ  | 18-885<br>11-30-84               | 4451458<br>05-29-01                        | NC<br>11-30-87                   |
| DHYDROERGOTAMINE MESYLATE;<br>HEPARIN SODIUM; LIDOCAINE HYDROCHLORIDE<br>0.5MG/0.7ML; 5000 UNITS/0.7ML;<br>7.46MG/0.7ML | EMBOLEX<br>(INJECTABLE; INJECTION)         | SANDOZ PHARMS/SANDOZ  | 18-885<br>11-30-84               | 4451458<br>05-29-01                        | NC<br>11-30-87                   |
| DILTIAZEM HYDROCHLORIDE<br>30MG   | CARDIZEM<br>(TABLET; ORAL)                 | MARION LABORATORIES   | 18-602<br>11-05-82               | 3562257<br>02-09-88                        | NCE<br>11-05-92                  |

**TABLE IV. NDA'S APPROVED FROM 1-1-82 TO 2-28-85 AND NDA'S WITH APPROPRIATE PATENT AND EXCLUSIVITY INFORMATION**

| <u>ACTIVE INGREDIENT(S)<br/>STRENGTH(S)</u>    | <u>TRADE NAME<br/>(DOSAGE FORM; ROUTE)</u>           | <u>APPLICANT NAME</u> | <u>NDA NO.<br/>APPROVAL DATE</u> | <u>PATENT NO.<br/>EXP. DATE</u>            | <u>EXCLUSIVITY<br/>EXP. DATE</u> |
|--|--|-----------------------|----------------------------------|--|----------------------------------|
| DILTIAZEM HYDROCHLORIDE<br>60MG                | CARDIZEM<br>(TABLET; ORAL)                           | MARION LABORATORIES   | 18-602<br>11-05-82               | 3562257<br>02-09-88                        | NCE<br>11-05-92                  |
| DINOPROST TROMETHAMINE<br>EQ 5MG BASE/ML       | PROSTIN F2 ALPHA<br>(INJECTABLE; INJECTION)          | UPJOHN                | 17-434<br>11-26-73               | 3706789<br>12-19-89<br>3778506<br>12-11-90 |                                  |
| DINOPROSTONE<br>20MG                           | PROSTIN E2<br>(SUPPOSITORY; VAGINAL)                 | UPJOHN                | 17-810<br>08-23-77               | 3899587<br>08-12-92<br>3598858<br>08-10-88 |                                  |
| DIPIVEFRIN HYDROCHLORIDE<br>0.1%               | PROPINE<br>(SOLUTION; OPHTHALMIC)                    | ALLERGAN PHARMS       | 18-239<br>05-02-80               | 3839584<br>10-01-91<br>3809714<br>05-07-91 |                                  |
| DISOPYRAMIDE PHOSPHATE<br>EQ 100MG BASE        | NORPACE CR<br>(CAPSULE, CONTROLLED<br>RELEASE; ORAL) | SEARLE/SEARLE PHARMS  | 18-655<br>07-20-82               |  | NDF<br>09-24-86                  |
| DISOPYRAMIDE PHOSPHATE<br>EQ 150MG BASE        | NORPACE CR<br>(CAPSULE, CONTROLLED<br>RELEASE; ORAL) | SEARLE/SEARLE PHARMS  | 18-655<br>07-20-82               |  | NDF<br>09-24-86                  |
| DIVALPROEX SODIUM<br>EQ 250MG BASE             | DEPAKOTE<br>(TABLET, ENTERIC COATED;<br>ORAL)        | ABBOTT LABORATORIES   | 18-723<br>03-10-83               |  | NE<br>09-24-86                   |
| DIVALPROEX SODIUM<br>EQ 500MG BASE             | DEPAKOTE<br>(TABLET, ENTERIC COATED;<br>ORAL)        | ABBOTT LABORATORIES   | 18-723<br>03-10-83               |  | NE<br>09-24-86                   |
| DOBUTAMINE HYDROCHLORIDE<br>EQ 250MG BASE/VIAL | DOBUTREX<br>(INJECTABLE; INJECTION)                  | ELI LILLY             | 17-820<br>07-18-78               | 3987200<br>10-19-93                        |                                  |
| DOPAMINE HYDROCHLORIDE<br>80MG/ML              | DOPAMINE HCL<br>(INJECTABLE; INJECTION)              | ABBOTT LABORATORIES   | 18-132<br>07-09-82               |  |                                  |

**TABLE IV. NDA'S APPROVED FROM 1-1-82 TO 2-28-85 AND NDA'S WITH APPROPRIATE PATENT AND EXCLUSIVITY INFORMATION**

| <u>ACTIVE INGREDIENT(S)<br/>STRENGTH(S)</u> | <u>TRADE NAME<br/>(DOSAGE FORM; ROUTE)</u> | <u>APPLICANT NAME</u> | <u>NDA NO.<br/>APPROVAL DATE</u> | <u>PATENT NO.<br/>EXP. DATE</u> | <u>EXCLUSIVITY<br/>EXP. DATE</u> |
|---|--|-----------------------|----------------------------------|---------------------------------|----------------------------------|
| DOPAMINE HYDROCHLORIDE<br>80MG/ML           | DOPAMINE<br>(INJECTABLE; INJECTION)        | ELKINS-SINN/AHROBINS  | 18-398<br>03-22-82               |                                 |                                  |
| DOPAMINE HYDROCHLORIDE<br>40MG/ML           | DOPAMINE HCL<br>(INJECTABLE; INJECTION)    | BRISTOL LABS/B-M      | 18-549<br>03-11-83               |                                 |                                  |
| DOPAMINE HYDROCHLORIDE<br>40MG/ML           | DOPAMINE<br>(INJECTABLE; INJECTION)        | ASTRA PHARM PRODS     | 18-656<br>06-28-83               |                                 |                                  |
| DOXEPIN HYDROCHLORIDE<br>EQ 25MG BASE       | SINEQUAN<br>(CAPSULE; ORAL)                | PFIZER LABS/PFIZER    | 16-798<br>09-23-69               | 3420851<br>01-07-86             |                                  |
| DOXEPIN HYDROCHLORIDE<br>EQ 50MG BASE       | SINEQUAN<br>(CAPSULE; ORAL)                | PFIZER LABS/PFIZER    | 16-798<br>09-23-69               | 3420851<br>01-07-86             |                                  |
| DOXEPIN HYDROCHLORIDE<br>EQ 10MG BASE       | SINEQUAN<br>(CAPSULE; ORAL)                | PFIZER LABS/PFIZER    | 16-798<br>03-31-75               | 3420851<br>01-07-86             |                                  |
| DOXEPIN HYDROCHLORIDE<br>EQ 100MG BASE      | SINEQUAN<br>(CAPSULE; ORAL)                | PFIZER LABS/PFIZER    | 16-798<br>03-31-75               | 3420851<br>01-07-86             |                                  |
| DOXEPIN HYDROCHLORIDE<br>EQ 75MG BASE       | SINEQUAN<br>(CAPSULE; ORAL)                | PFIZER LABS/PFIZER    | 16-798<br>06-04-76               | 3420851<br>01-07-86             |                                  |
| DOXEPIN HYDROCHLORIDE<br>EQ 150MG BASE      | SINEQUAN<br>(CAPSULE; ORAL)                | PFIZER LABS/PFIZER    | 16-798<br>03-15-78               | 3420851<br>01-07-86             |                                  |
| DOXEPIN HYDROCHLORIDE<br>EQ 10MG BASE       | ADAPIN<br>(CAPSULE; ORAL)                  | PENNMALT PHARM        | 16-987<br>01-31-72               | 3420851<br>01-07-86             |                                  |
| DOXEPIN HYDROCHLORIDE<br>EQ 25MG BASE       | ADAPIN<br>(CAPSULE; ORAL)                  | PENNMALT PHARM        | 16-987<br>01-31-72               | 3420851<br>01-07-86             |                                  |
| DOXEPIN HYDROCHLORIDE<br>EQ 50MG BASE       | ADAPIN<br>(CAPSULE; ORAL)                  | PENNMALT PHARM        | 16-987<br>01-31-72               | 3420851<br>01-07-86             |                                  |
| DOXEPIN HYDROCHLORIDE<br>EQ 100MG BASE      | ADAPIN<br>(CAPSULE; ORAL)                  | PENNMALT PHARM        | 16-987<br>12-12-77               | 3420851<br>01-07-86             |                                  |
| DOXEPIN HYDROCHLORIDE<br>EQ 75MG BASE       | ADAPIN<br>(CAPSULE; ORAL)                  | PENNMALT PHARM        | 16-987<br>04-15-80               | 3420851<br>01-07-86             |                                  |

TABLE IV. NDA'S APPROVED FROM 1-1-82 TO 2-28-85 AND NDA'S WITH APPROPRIATE PATENT AND EXCLUSIVITY INFORMATION

| <u>ACTIVE INGREDIENT(S)<br/>STRENGTH(S)</u>               | <u>TRADE NAME<br/>(DOSAGE FORM; ROUTE)</u> | <u>APPLICANT NAME</u> | <u>NDA NO.<br/>APPROVAL DATE</u> | <u>PATENT NO.<br/>EXP. DATE</u>                                   | <u>EXCLUSIVITY<br/>EXP. DATE</u> |
|---|--|-----------------------|----------------------------------|---|----------------------------------|
| DOXEPTIN HYDROCHLORIDE<br>EQ 10MG BASE/ML                 | SINEQUAN<br>(CONCENTRATE; ORAL)            | PFIZER LABS/PFIZER    | 17-516<br>03-11-74               | 3420851<br>01-07-86   |                                  |
| ECONAZOLE NITRATE<br>1%                                   | SPECTAZOLE<br>(CREAM; TOPICAL)             | ORTHO PHARMACEUTICAL  | 18-751<br>12-23-82               | 3717655<br>02-20-90<br>3839574<br>10-01-91                        | NCE<br>12-23-92                  |
| ENFLURANE<br>99.9%  | ETHRANE<br>(LIQUID; INHALATION)            | ANAQUEST/BOC          | 17-087<br>08-28-72               | 3469011<br>09-23-86<br>3527813<br>09-08-87                        |                                  |
| EPINEPHRINE; ETIDOCAINE HYDROCHLORIDE<br>0.005MG/ML; 0.5% | DURANEST<br>(INJECTABLE; INJECTION)        | ASTRA PHARM PRODS     | 17-751<br>08-30-76               | 3862321<br>01-21-92<br>3812147<br>05-21-91                        |                                  |
| EPINEPHRINE; ETIDOCAINE HYDROCHLORIDE<br>0.005MG/ML; 1%   | DURANEST<br>(INJECTABLE; INJECTION)        | ASTRA PHARM PRODS     | 17-751<br>08-30-76               | 3862321<br>01-21-92<br>3812147<br>05-21-91                        |                                  |
| EPINEPHRINE; ETIDOCAINE HYDROCHLORIDE<br>0.005MG/ML; 1.5% | DURANEST<br>(INJECTABLE; INJECTION)        | ASTRA PHARM PRODS     | 17-751<br>08-30-76               | 3862321<br>01-21-92<br>3812147<br>05-21-91                        |                                  |
| ERGOLOID MESYLATES<br>1MG                                 | HYDERGINE LC<br>(CAPSULE; ORAL)            | SANDOZ PHARMS/SANDOZ  | 18-706<br>01-18-83               |   | NDF<br>09-24-86                  |
| ESTROGENS, CONJUGATED<br>0.9MG                            | PREMARIN<br>(TABLET; ORAL)                 | AYERST LABS/AMHO      | 04-782<br>01-26-84               |   | NS<br>09-24-86                   |
| ETHINYL ESTRADIOL; LEVONORGESTREL<br>0.03MG; 0.15MG       | NORDETTE-21<br>(TABLET; ORAL-21)           | WYETH LABS/AMHO       | 18-668<br>05-10-82               | 3666858<br>05-30-89<br>3850911<br>11-26-91<br>3959322<br>11-26-91 | NC<br>09-24-86                   |

TABLE IV. NDA'S APPROVED FROM 1-1-82 TO 2-28-85 AND NDA'S WITH APPROPRIATE PATENT AND EXCLUSIVITY INFORMATION

| <u>ACTIVE INGREDIENT(S)<br/>STRENGTH(S)</u>   | <u>TRADE NAME<br/>(DOSAGE FORM; ROUTE)</u> | <u>APPLICANT NAME</u> | <u>NDA NO.<br/>APPROVAL DATE</u> | <u>PATENT NO.<br/>EXP. DATE</u>  | <u>EXCLUSIVITY<br/>EXP. DATE</u> |
|---|--|-----------------------|----------------------------------|--|----------------------------------|
| ETHINYL ESTRADIOL; LEVONORGESTREL<br>0.03MG; 0.15MG                                       | NORDETTE-28<br>(TABLET; ORAL-28)           | WYETH LABS/AMHO       | 18-782<br>07-21-82               | 3666858<br>05-30-89<br>3850911<br>11-26-91<br>3959322<br>11-26-91                        | NC<br>09-24-86                   |
| ETHINYL ESTRADIOL; LEVONORGESTREL<br>0.03MG; 0.05MG<br>0.04MG; 0.075MG<br>0.03MG; 0.125MG | TRIPHASIL-28<br>(TABLET; ORAL-28)          | WYETH LABS/AMHO       | 19-190<br>11-01-84               | 3666858<br>05-30-89<br>3850911<br>11-26-91<br>3959322<br>11-26-91<br>3957982<br>05-18-93 | NS<br>11-01-87                   |
| ETHINYL ESTRADIOL; LEVONORGESTREL<br>0.03MG; 0.05MG<br>0.04MG; 0.075MG<br>0.03MG; 0.125MG | TRIPHASIL-21<br>(TABLET; ORAL-21)          | WYETH LABS/AMHO       | 19-192<br>11-01-84               | 3666858<br>05-30-89<br>3850911<br>11-26-91<br>3959322<br>11-26-91<br>3957982<br>05-18-93 | NS<br>11-01-87                   |
| ETHINYL ESTRADIOL; NORETHINDRONE<br>0.035MG; 0.5MG AND 1MG                                | ORTHO-NOVUM 10/11-21<br>(TABLET; ORAL-21)  | ORTHO PHARMACEUTICAL  | 18-354<br>01-11-82               |  | D-5<br>09-24-86                  |
| ETHINYL ESTRADIOL; NORETHINDRONE<br>0.035MG; 0.5MG AND 1MG                                | ORTHO-NOVUM 10/11-28<br>(TABLET; ORAL-28)  | ORTHO PHARMACEUTICAL  | 18-354<br>01-11-82               |  | D-5<br>09-24-86                  |
| ETHINYL ESTRADIOL; NORETHINDRONE<br>0.035MG; 0.5MG AND 1MG                                | TRI-NORINYL 21-DAY<br>(TABLET; ORAL-21)    | SYNTEX (FP)           | 18-977<br>04-13-84               | 4390531<br>06-28-00  | D-6<br>09-24-86                  |
| ETHINYL ESTRADIOL; NORETHINDRONE<br>0.035MG; 0.5MG AND 1MG                                | TRI-NORINYL 28-DAY<br>(TABLET; ORAL-28)    | SYNTEX (FP)           | 18-977<br>04-13-84               | 4390531<br>06-28-00  | D-6<br>09-24-86                  |
| ETHINYL ESTRADIOL; NORETHINDRONE<br>0.035MG; 0.5MG, 0.75MG AND 1MG                        | ORTHO-NOVUM 7/7/7-21<br>(TABLET; ORAL-21)  | ORTHO PHARMACEUTICAL  | 18-985<br>04-04-84               |  | D-3<br>09-24-86                  |

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| <u>ACTIVE INGREDIENT(S)<br/>STRENGTH(S)</u>                        | <u>TRADE NAME<br/>(DOSAGE FORM; ROUTE)</u> | <u>APPLICANT NAME</u> | <u>NDA NO.<br/>APPROVAL DATE</u> | <u>PATENT NO.<br/>EXP. DATE</u>                                   | <u>EXCLUSIVITY<br/>EXP. DATE</u> |
|--|--|-----------------------|----------------------------------|---|----------------------------------|
| ETHINYL ESTRADIOL; NORETHINDRONE<br>0.035MG; 0.5MG, 0.75MG AND 1MG | ORTHO-NOVUM 7/7/7-28<br>(TABLET; ORAL-28)  | ORTHO PHARMACEUTICAL  | 18-985<br>04-04-84               |   | D-3<br>09-24-86                  |
| ETHINYL ESTRADIOL; NORETHINDRONE<br>0.035MG; 0.5MG AND 1MG         | ORTHO-NOVUM 7/14-21<br>(TABLET; ORAL-21)   | ORTHO PHARMACEUTICAL  | 19-004<br>04-04-84               |   | D-4<br>09-24-86                  |
| ETHINYL ESTRADIOL; NORETHINDRONE<br>0.035MG; 0.5MG AND 1MG         | ORTHO-NOVUM 7/14-28<br>(TABLET; ORAL-28)   | ORTHO PHARMACEUTICAL  | 19-004<br>04-04-84               |   | D-4<br>09-24-86                  |
| ETHINYL ESTRADIOL; NORGESTREL<br>0.05MG; 0.5MG                     | OVRAL<br>(TABLET; ORAL-21)                 | WYETH LABS/AMHO       | 16-672<br>04-16-68               | 3666858<br>05-30-89<br>3850911<br>11-26-91<br>3959322<br>11-26-91 |                                  |
| ETHINYL ESTRADIOL; NORGESTREL<br>0.05MG; 0.5MG                     | OVRAL-28<br>(TABLET; ORAL-28)              | WYETH LABS/AMHO       | 16-806<br>11-26-68               | 3666858<br>05-30-89<br>3850911<br>11-26-91<br>3959322<br>11-26-91 |                                  |
| ETHINYL ESTRADIOL; NORGESTREL<br>0.03MG; 0.3MG                     | LO/OVRAL<br>(TABLET; ORAL-21)              | WYETH LABS/AMHO       | 17-612<br>03-17-75               | 3666858<br>05-30-89<br>3850911<br>11-26-91<br>3959322<br>11-26-91 |                                  |
| ETHINYL ESTRADIOL; NORGESTREL<br>0.03MG; 0.3MG                     | LO/OVRAL-28<br>(TABLET; ORAL-28)           | WYETH LABS/AMHO       | 17-802<br>03-16-76               | 3666858<br>05-30-89<br>3850911<br>11-26-91<br>3959322<br>11-26-91 |                                  |
| ETIDOCaine HYDROCHLORIDE<br>0.5%                                   | DURANEST<br>(INJECTABLE; INJECTION)        | ASTRA PHARM PRODS     | 17-751<br>08-30-76               | 3862321<br>01-21-92<br>3812147<br>05-21-91                        |                                  |

**TABLE IV. NDA'S APPROVED FROM 1-1-82 TO 2-28-85 AND NDA'S WITH APPROPRIATE PATENT AND EXCLUSIVITY INFORMATION**

| <u>ACTIVE INGREDIENT(S)<br/>STRENGTH(S)</u> | <u>TRADE NAME<br/>(DOSAGE FORM; ROUTE)</u>        | <u>APPLICANT NAME</u> | <u>NDA NO.<br/>APPROVAL DATE</u> | <u>PATENT NO.<br/>EXP. DATE</u>  | <u>EXCLUSIVITY<br/>EXP. DATE</u> |
|---|---|-----------------------|----------------------------------|--|----------------------------------|
| ETIDOCAINE HYDROCHLORIDE<br>1%              | DURANEST<br>(INJECTABLE; INJECTION)               | ASTRA PHARM PRODS     | 17-751<br>08-30-76               | 3862321<br>01-21-92<br>3812147<br>05-21-91   |                                  |
| ETIDRONATE DISODIUM<br>200MG                | DIDRONEL<br>(TABLET; ORAL)                        | NORMICH EATON/P&G     | 17-831<br>09-01-77               | 4254114<br>03-03-98<br>4216211<br>08-05-97<br>4137309<br>01-30-96<br>3683080<br>08-08-89<br>4254114<br>03-03-98<br>4216211<br>08-05-97<br>4137309<br>01-30-96<br>3683080<br>08-08-89 | NS<br>09-24-86                   |
| ETIDRONATE DISODIUM<br>400MG                | DIDRONEL<br>(TABLET; ORAL)                        | NORMICH EATON/P&G     | 17-831<br>07-06-84               |  |                                  |
| ETOMIDATE<br>2MG/ML                         | AMIDATE<br>(INJECTABLE; INJECTION)                | ABBOTT LABORATORIES   | 18-227<br>09-07-82               |  | NCE<br>09-07-92                  |
| ETOPOSIDE<br>20MG/ML                        | VEPSID<br>(INJECTABLE; INJECTION)                 | BRISTOL LABS/B-M      | 18-768<br>11-10-83               | 3524844<br>08-18-87  | NCE<br>11-10-93                  |
| FENFLURAMINE HYDROCHLORIDE<br>60MG          | PONDIMIN<br>(TABLET, CONTROLLED<br>RELEASE; ORAL) | AH ROBINS             | 16-618<br>07-27-82               |  | NDF<br>09-24-86                  |
| FENOPROFEN CALCIUM<br>EQ 300MG BASE         | MALFON<br>(CAPSULE; ORAL)                         | DISTA PRODS/LILLY     | 17-604<br>03-16-76               | 3600437<br>08-17-88  |                                  |
| FENOPROFEN CALCIUM<br>EQ 200MG BASE         | MALFON 200<br>(CAPSULE; ORAL)                     | DISTA PRODS/LILLY     | 17-604<br>10-15-80               | 3600437<br>08-17-88  |                                  |
| FENOPROFEN CALCIUM<br>EQ 600MG BASE         | MALFON<br>(TABLET; ORAL)                          | DISTA PRODS/LILLY     | 17-710<br>03-16-76               | 3600437<br>08-17-88  |                                  |

**TABLE IV. NDA'S APPROVED FROM 1-1-82 TO 2-28-85 AND NDA'S WITH APPROPRIATE PATENT AND EXCLUSIVITY INFORMATION**

| <u>ACTIVE INGREDIENT(S)<br/>STRENGTH(S)</u> | <u>TRADE NAME<br/>(DOSAGE FORM; ROUTE)</u>    | <u>APPLICANT NAME</u> | <u>NDA NO.<br/>APPROVAL DATE</u> | <u>PATENT NO.<br/>EXP. DATE</u> | <u>EXCLUSIVITY<br/>EXP. DATE</u> |
|---|---|-----------------------|----------------------------------|---------------------------------|----------------------------------|
| FENTANYL CITRATE<br>EQ 0.05MG BASE/ML       | FENTANYL CITRATE<br>(INJECTABLE; INJECTION)   | ABBOTT LABORATORIES   | 19-115<br>01-12-85               |                                 |                                  |
| FENTANYL CITRATE<br>EQ 0.05MG BASE/ML       | FENTANYL<br>(INJECTABLE; INJECTION)           | ELKINS-SINN/AHROBINS  | 19-101<br>07-11-84               |                                 |                                  |
| FLUNISOLIDE<br>0.025MG/INH                  | BRONALIDE<br>(AEROSOL; INHALATION)            | SYNTEX LABS/SYNTEX    | 18-340<br>08-17-84               |                                 | NDF<br>09-24-86                  |
| FLUOCINONIDE<br>0.05%                       | LIDEX<br>(SOLUTION; TOPICAL)                  | SYNTEX LABS/SYNTEX    | 18-849<br>04-06-84               |                                 | NDF<br>09-24-86                  |
| FLUOCINONIDE<br>0.05%                       | VASODERM<br>(CREAM; TOPICAL)                  | K-LINE PHARMS         | 19-117<br>06-26-84               |                                 |                                  |
| FLUPHENAZINE DECANOATE<br>25MG/ML           | PROLIXIN DECANOATE<br>(INJECTABLE; INJECTION) | ER SQUIBB AND SONS    | 16-727<br>06-20-72               | 3394131<br>07-23-85             |                                  |
| FLUPHENAZINE ENANTHATE<br>25MG/ML           | PROLIXIN ENANTHATE<br>(INJECTABLE; INJECTION) | ER SQUIBB AND SONS    | 16-110<br>03-15-67               | 3394131<br>07-23-85             |                                  |
| FLURANDRENOLIDE<br>0.004MG/SQ CM            | CORDRAN<br>(TAPE; TOPICAL)                    | DISTA PRODS/LILLY     | 16-455<br>07-29-69               | 3632740<br>01-04-89             |                                  |
| FLURAZEPAM HYDROCHLORIDE<br>15MG            | DALMANE<br>(CAPSULE; ORAL)                    | ROCHE PRODUCTS        | 16-721<br>04-07-70               | 4316897<br>02-23-99             |                                  |
| FLURAZEPAM HYDROCHLORIDE<br>30MG            | DALMANE<br>(CAPSULE; ORAL)                    | ROCHE PRODUCTS        | 16-721<br>04-07-70               | 4316897<br>02-23-99             |                                  |
| FUROSEMIDE<br>20MG                          | FUROSEMIDE<br>(TABLET; ORAL)                  | CHELSEA LABORATORIES  | 18-369<br>05-14-82               |                                 |                                  |
| FUROSEMIDE<br>40MG                          | FUROSEMIDE<br>(TABLET; ORAL)                  | CHELSEA LABORATORIES  | 18-369<br>05-14-82               |                                 |                                  |
| FUROSEMIDE<br>40MG                          | FUROSEMIDE<br>(TABLET; ORAL)                  | SUPERPHARM            | 18-370<br>02-10-83               |                                 |                                  |
| FUROSEMIDE<br>20MG                          | FUROSEMIDE<br>(TABLET; ORAL)                  | SUPERPHARM            | 18-370<br>06-26-84               |                                 |                                  |

TABLE IV. NDA'S APPROVED FROM 1-1-82 TO 2-28-85 AND NDA'S WITH APPROPRIATE PATENT AND EXCLUSIVITY INFORMATION

| <u>ACTIVE INGREDIENT(S)</u><br><u>STRENGTH(S)</u> | <u>TRADE NAME</u><br><u>(DOSAGE FORM; ROUTE)</u> | <u>APPLICANT NAME</u> | <u>NDA NO.</u><br><u>APPROVAL DATE</u> | <u>PATENT NO.</u><br><u>EXP. DATE</u> | <u>EXCLUSIVITY</u><br><u>EXP. DATE</u> |
|---|--|-----------------------|--|---------------------------------------|--|
| FUROSEMIDE<br>20MG                                | FUROSEMIDE<br>(TABLET; ORAL)                     | ZENITH LABORATORIES   | 18-413<br>11-30-83                     |                                       |  |
| FUROSEMIDE<br>40MG                                | FUROSEMIDE<br>(TABLET; ORAL)                     | ZENITH LABORATORIES   | 18-413<br>11-30-83                     |                                       |  |
| FUROSEMIDE<br>20MG                                | FUROSEMIDE<br>(TABLET; ORAL)                     | LEDERLE LABS/AM CYAN  | 18-415<br>07-27-82                     |                                       |  |
| FUROSEMIDE<br>40MG                                | FUROSEMIDE<br>(TABLET; ORAL)                     | LEDERLE LABS/AM CYAN  | 18-415<br>07-27-82                     |                                       |  |
| FUROSEMIDE<br>80MG                                | FUROSEMIDE<br>(TABLET; ORAL)                     | LEDERLE LABS/AM CYAN  | 18-415<br>11-26-84                     |                                       |  |
| FUROSEMIDE<br>20MG                                | FUROSEMIDE<br>(TABLET; ORAL)                     | PARKE-DAVIS/M-L       | 18-419<br>01-31-83                     |                                       |  |
| FUROSEMIDE<br>40MG                                | FUROSEMIDE<br>(TABLET; ORAL)                     | PARKE-DAVIS/M-L       | 18-419<br>01-31-83                     |                                       |  |
| FUROSEMIDE<br>80MG                                | FUROSEMIDE<br>(TABLET; ORAL)                     | PARKE-DAVIS/M-L       | 18-419<br>11-13-84                     |                                       |  |
| FUROSEMIDE<br>10MG/ML                             | FUROSEMIDE<br>(INJECTABLE; INJECTION)            | PARKE-DAVIS/M-L       | 18-420<br>02-26-82                     |                                       |  |
| FUROSEMIDE<br>10MG/ML                             | FUROSEMIDE<br>(INJECTABLE; INJECTION)            | LYPHOMED              | 18-507<br>07-30-82                     |                                       |  |
| FUROSEMIDE<br>80MG                                | FUROSEMIDE<br>(TABLET; ORAL)                     | CORD LABORATORIES     | 18-569<br>08-14-84                     |                                       |  |
| FUROSEMIDE<br>10MG/ML                             | FUROSEMIDE<br>(INJECTABLE; INJECTION)            | NATCON                | 18-579<br>11-30-83                     |                                       |  |
| FUROSEMIDE<br>10MG/ML                             | FUROSEMIDE<br>(INJECTABLE; INJECTION)            | ABBOTT LABORATORIES   | 18-667<br>05-28-82                     |                                       |  |
| FUROSEMIDE<br>10MG/ML                             | FUROSEMIDE<br>(INJECTABLE; INJECTION)            | WYETH LABS/AMHO       | 18-670<br>07-20-82                     |                                       |  |

TABLE IV. NDA'S APPROVED FROM 1-1-82 TO 2-28-85 AND NDA'S WITH APPROPRIATE PATENT AND EXCLUSIVITY INFORMATION

| <u>ACTIVE INGREDIENT(S)</u><br><u>STRENGTH(S)</u> | <u>TRADE NAME</u><br><u>(DOSAGE FORM; ROUTE)</u> | <u>APPLICANT NAME</u> | <u>NDA NO.</u><br><u>APPROVAL DATE</u> | <u>PATENT NO.</u><br><u>EXP. DATE</u> | <u>EXCLUSIVITY</u><br><u>EXP. DATE</u> |
|---|--|-----------------------|--|---------------------------------------|--|
| FUROSEMIDE<br>40MG                                | FUROSEMIDE<br>(TABLET; ORAL)                     | DRUMMER/PHOENIX       | 18-750<br>07-30-84                     |                                       |  |
| FUROSEMIDE<br>20MG                                | FUROSEMIDE<br>(TABLET; ORAL)                     | INTL MEDICATION SYS   | 18-753<br>02-28-84                     |                                       |  |
| FUROSEMIDE<br>40MG                                | FUROSEMIDE<br>(TABLET; ORAL)                     | INTL MEDICATION SYS   | 18-753<br>02-28-84                     |                                       |  |
| FUROSEMIDE<br>40MG                                | FUROSEMIDE<br>(TABLET; ORAL)                     | BARR LABORATORIES     | 18-790<br>11-29-83                     |                                       |  |
| FUROSEMIDE<br>20MG                                | FUROSEMIDE<br>(TABLET; ORAL)                     | ROXANE LABORATORIES   | 18-823<br>11-10-83                     |                                       |  |
| FUROSEMIDE<br>40MG                                | FUROSEMIDE<br>(TABLET; ORAL)                     | ROXANE LABORATORIES   | 18-823<br>11-10-83                     |                                       |  |
| FUROSEMIDE<br>20MG                                | FUROSEMIDE<br>(TABLET; ORAL)                     | KALAPHARM             | 18-868<br>06-28-83                     |                                       |  |
| FUROSEMIDE<br>40MG                                | FUROSEMIDE<br>(TABLET; ORAL)                     | KALAPHARM             | 18-868<br>06-28-83                     |                                       |  |
| FUROSEMIDE<br>10MG/ML                             | FUROSEMIDE<br>(INJECTABLE; INJECTION)            | INVENEX LABS/LIFE     | 18-902<br>05-22-84                     |                                       |  |
| FUROSEMIDE<br>10MG/ML                             | FUROSEMIDE<br>(INJECTABLE; INJECTION)            | INVENEX LABS/LIFE     | 19-036<br>08-13-84                     |                                       |  |
| GEMFIBROZIL<br>200MG                              | LOPID<br>(CAPSULE; ORAL)                         | PARKE-DAVIS/M-L       | 18-422<br>12-21-81                     | 3674836<br>07-04-89                   |  |
| GEMFIBROZIL<br>300MG                              | LOPID<br>(CAPSULE; ORAL)                         | PARKE-DAVIS/M-L       | 18-422<br>12-21-81                     | 3674836<br>07-04-89                   |  |
| GLIPIZIDE<br>5MG                                  | GLUCOTROL<br>(TABLET; ORAL)                      | ROERIG/PFIZER         | 17-783<br>05-08-84                     | 3669966<br>04-21-92                   | NCE<br>05-08-94                        |
| GLIPIZIDE<br>10MG                                 | GLUCOTROL<br>(TABLET; ORAL)                      | ROERIG/PFIZER         | 17-783<br>05-08-84                     | 3669966<br>04-21-92                   | NCE<br>05-08-94                        |

**TABLE IV. NDA'S APPROVED FROM 1-1-82 TO 2-28-85 AND NDA'S WITH APPROPRIATE PATENT AND EXCLUSIVITY INFORMATION**

| <u>ACTIVE INGREDIENT(S)<br/>STRENGTH(S)</u> | <u>TRADE NAME<br/>(DOSAGE FORM; ROUTE)</u> | <u>APPLICANT NAME</u> | <u>NDA NO.<br/>APPROVAL DATE</u> | <u>PATENT NO.<br/>EXP. DATE</u> | <u>EXCLUSIVITY<br/>EXP. DATE</u> |
|---|--|-----------------------|----------------------------------|---------------------------------|----------------------------------|
| GLYBURIDE<br>1.25MG                         | MICRONASE<br>(TABLET; ORAL)                | UPJOHN                | 17-498<br>05-01-84               | 3426067                         | NCE                              |
|   |  |                       |                                  | 04-21-92                        | 05-01-94                         |
|   |  |                       |                                  | 3454635                         |                                  |
|   |  |                       |                                  | 04-21-92                        |                                  |
|   |  |                       |                                  | 3507954                         |                                  |
| GLYBURIDE<br>2.5MG                          | MICRONASE<br>(TABLET; ORAL)                | UPJOHN                | 17-498<br>05-01-84               | 04-21-92                        | NCE                              |
|   |  |                       |                                  | 04-21-92                        | 05-01-94                         |
|   |  |                       |                                  | 3454635                         |                                  |
|   |  |                       |                                  | 04-21-92                        |                                  |
|   |  |                       |                                  | 3507954                         |                                  |
| GLYBURIDE<br>5MG                            | MICRONASE<br>(TABLET; ORAL)                | UPJOHN                | 17-498<br>05-01-84               | 3426067                         | NCE                              |
|   |  |                       |                                  | 04-21-92                        | 05-01-94                         |
|   |  |                       |                                  | 3454635                         |                                  |
|   |  |                       |                                  | 04-21-92                        |                                  |
|   |  |                       |                                  | 3507954                         |                                  |
| GLYBURIDE<br>1.25MG                         | DIABETA<br>(TABLET; ORAL)                  | HOECHST-ROUSSEL       | 17-532<br>05-01-84               | 3426067                         | NCE                              |
|   |  |                       |                                  | 04-21-92                        | 05-01-94                         |
|   |  |                       |                                  | 3454635                         |                                  |
|   |  |                       |                                  | 04-21-92                        |                                  |
|   |  |                       |                                  | 3507961                         |                                  |

**TABLE IV. NDA'S APPROVED FROM 1-1-82 TO 2-28-85 AND NDA'S WITH APPROPRIATE PATENT AND EXCLUSIVITY INFORMATION**

| <u>ACTIVE INGREDIENT(S)</u><br><u>STRENGTH(S)</u> | <u>TRADE NAME</u><br><u>(DOSAGE FORM; ROUTE)</u>  | <u>APPLICANT NAME</u> | <u>NDA NO.</u><br><u>APPROVAL DATE</u> | <u>PATENT NO.</u><br><u>EXP. DATE</u> | <u>EXCLUSIVITY</u><br><u>EXP. DATE</u> |
|---|---|-----------------------|--|---------------------------------------|--|
| GLYBURIDE<br>2.5MG                                | DIABETA<br>(TABLET; ORAL)                         | HOECHST-ROUSSEL       | 17-532                                 | 3426067                               | NCE                                    |
|   |   |                       | 05-01-84                               | 04-21-92                              | 05-01-94                               |
|   |   |                       |  | 3454635                               |  |
|   |   |                       |  | 04-21-92                              |  |
|   |   |                       |  | 3507961                               |  |
|   |   |                       |  | 04-21-92                              |  |
| GLYBURIDE<br>5MG                                  | DIABETA<br>(TABLET; ORAL)                         | HOECHST-ROUSSEL       | 17-532                                 | 3426067                               | NCE                                    |
|   |   |                       | 05-01-84                               | 04-21-92                              | 05-01-94                               |
|   |   |                       |  | 3454635                               |  |
|   |   |                       |  | 04-21-92                              |  |
|   |   |                       |  | 3507961                               |  |
|   |   |                       |  | 04-21-92                              |  |
| GONADORELIN HYDROCHLORIDE<br>EQ 0.1MG BASE/VIAL   | FACTREL<br>(INJECTABLE; INJECTION)                | AYERST LABS/AMHO      | 18-123                                 | 3947569                               | NCE                                    |
|   |   |                       | 09-30-82                               | 03-30-93                              | 09-30-92                               |
|   |   |                       |  | 4110438                               |  |
|   |   |                       |  | 08-29-95                              |  |
|   |   |                       |  | 3947569                               |  |
|   |   |                       |  | 03-30-93                              |  |
| GONADORELIN HYDROCHLORIDE<br>EQ 0.5MG BASE/VIAL   | FACTREL<br>(INJECTABLE; INJECTION)                | AYERST LABS/AMHO      | 18-123                                 | 3947569                               | NCE                                    |
|   |   |                       | 09-30-82                               | 03-30-93                              | 09-30-92                               |
|   |   |                       |  | 4110438                               |  |
|   |   |                       |  | 08-29-95                              |  |
|   |   |                       |  | 3947569                               |  |
|   |   |                       |  | 03-30-93                              |  |
| GONADOTROPIN, CHORIONIC<br>2,000 UNITS/VIAL       | CHORIONIC GONADOTROPIN<br>(INJECTABLE; INJECTION) | CARTER-GLOGAU LABS    | 17-016                                 |                                       |  |
|   |   |                       | 12-27-84                               |                                       |  |
| GUANABENZ ACETATE<br>EQ 4MG BASE                  | WYTENSIN<br>(TABLET; ORAL)                        | WYETH LABS/AMHO       | 18-587                                 | 3658993                               | NCE                                    |
|   |   |                       | 09-07-82                               | 04-25-89                              | 09-07-92                               |
| GUANABENZ ACETATE<br>EQ 8MG BASE                  | WYTENSIN<br>(TABLET; ORAL)                        | WYETH LABS/AMHO       | 18-587                                 | 3658993                               | NCE                                    |
|   |   |                       | 09-07-82                               | 04-25-89                              | 09-07-92                               |

**TABLE IV. NDA'S APPROVED FROM 1-1-82 TO 2-28-85 AND NDA'S WITH APPROPRIATE PATENT AND EXCLUSIVITY INFORMATION**

| <u>ACTIVE INGREDIENT(S)<br/>STRENGTH(S)</u> | <u>TRADE NAME<br/>(DOSAGE FORM; ROUTE)</u>    | <u>APPLICANT NAME</u> | <u>NDA NO.<br/>APPROVAL DATE</u> | <u>PATENT NO.<br/>EXP. DATE</u> | <u>EXCLUSIVITY<br/>EXP. DATE</u> |
|---|---|-----------------------|----------------------------------|---------------------------------|----------------------------------|
| GUANADREL SULFATE<br>10MG                   | HYLOREL<br>(TABLET; ORAL)                     | UPJOHN                | 18-104<br>12-29-82               | 3547951<br>12-15-87             | NCE<br>12-29-92                  |
| GUANADREL SULFATE<br>25MG                   | HYLOREL<br>(TABLET; ORAL)                     | UPJOHN                | 18-104<br>12-29-82               | 3547951<br>12-15-87             | NCE<br>12-29-92                  |
| HALAZEPAM<br>20MG                           | PAXIPAM<br>(TABLET; ORAL)                     | SCHERING              | 17-736<br>09-24-81               | 3429874<br>02-25-86             |                                  |
| HALAZEPAM<br>40MG                           | PAXIPAM<br>(TABLET; ORAL)                     | SCHERING              | 17-736<br>09-24-81               | 3429874<br>02-25-86             |                                  |
| HALOPERIDOL<br>0.5MG                        | HALDOL<br>(TABLET; ORAL)                      | MCNEIL PHARM          | 15-921<br>04-12-67               | 3438991<br>04-15-86             | NS<br>09-24-86                   |
| HALOPERIDOL<br>1MG                          | HALDOL<br>(TABLET; ORAL)                      | MCNEIL PHARM          | 15-921<br>04-12-67               | 3438991<br>04-15-86             |                                  |
| HALOPERIDOL<br>2MG                          | HALDOL<br>(TABLET; ORAL)                      | MCNEIL PHARM          | 15-921<br>04-12-67               | 3438991<br>04-15-86             |                                  |
| HALOPERIDOL<br>5MG                          | HALDOL<br>(TABLET; ORAL)                      | MCNEIL PHARM          | 15-921<br>04-16-74               | 3438991<br>04-15-86             |                                  |
| HALOPERIDOL<br>10MG                         | HALDOL<br>(TABLET; ORAL)                      | MCNEIL PHARM          | 15-921<br>04-16-74               | 3438991<br>04-15-86             |                                  |
| HALOPERIDOL<br>20MG                         | HALDOL<br>(TABLET; ORAL)                      | MCNEIL PHARM          | 15-921<br>02-02-82               | 3438991<br>04-15-86             | NS<br>09-24-86                   |
| HALOPERIDOL LACTATE<br>EQ 2MG BASE/ML       | HALDOL<br>(CONCENTRATE; ORAL)                 | MCNEIL LABORATORIES   | 15-922<br>04-12-67               | 3438991<br>04-15-86             |                                  |
| HALOPERIDOL LACTATE<br>EQ 5MG BASE/ML       | HALDOL<br>(INJECTABLE; INJECTION)             | MCNEIL LABORATORIES   | 15-923<br>05-18-71               | 3438991<br>04-15-86             |                                  |
| HEPARIN SODIUM<br>10 UNITS/ML               | HEPARIN LOCK FLUSH<br>(INJECTABLE; INJECTION) | INVENEX LABS/LIFE     | 17-029<br>05-06-82               |                                 |                                  |

**TABLE IV. NDA'S APPROVED FROM 1-1-82 TO 2-28-85 AND NDA'S WITH APPROPRIATE PATENT AND EXCLUSIVITY INFORMATION**

| <u>ACTIVE INGREDIENT(S)<br/>STRENGTH(S)</u>                        | <u>TRADE NAME<br/>(DOSAGE FORM; ROUTE)</u>  | <u>APPLICANT NAME</u> | <u>NDA NO.<br/>APPROVAL DATE</u> | <u>PATENT NO.<br/>EXP. DATE</u> | <u>EXCLUSIVITY<br/>EXP. DATE</u> |
|--|---|-----------------------|----------------------------------|---------------------------------|----------------------------------|
| HEPARIN SODIUM; SODIUM CHLORIDE<br>200 UNITS/100ML; 900MG/100ML    | HEPARIN SODIUM 1000 UNITS<br>AND SODIUM CHLORIDE 0.9%<br>IN PLASTIC CONTAINER<br>(INJECTABLE; INJECTION)  | TRAVENOL LABS         | 18-609<br>04-28-82               |                                 |                                  |
| HEPARIN SODIUM; SODIUM CHLORIDE<br>200 UNITS/100ML; 900MG/100ML    | HEPARIN SODIUM 2000 UNITS<br>AND SODIUM CHLORIDE 0.9%<br>IN PLASTIC CONTAINER<br>(INJECTABLE; INJECTION)  | TRAVENOL LABS         | 18-609<br>04-28-82               |                                 |                                  |
| HEPARIN SODIUM; SODIUM CHLORIDE<br>500 UNITS/100ML; 900MG/100ML    | HEPARIN SODIUM 5000 UNITS<br>AND SODIUM CHLORIDE 0.9%<br>IN PLASTIC CONTAINER<br>(INJECTABLE; INJECTION)  | TRAVENOL LABS         | 18-609<br>04-28-82               |                                 |                                  |
| HEPARIN SODIUM; SODIUM CHLORIDE<br>1,000 UNITS/100ML; 900MG/100ML  | HEPARIN SODIUM 5000 UNITS<br>IN SODIUM CHLORIDE 0.9%<br>IN PLASTIC CONTAINER<br>(INJECTABLE; INJECTION)   | ABBOTT LABORATORIES   | 18-916<br>01-31-84               |                                 |                                  |
| HEPARIN SODIUM; SODIUM CHLORIDE<br>10,000 UNITS/100ML; 900MG/100ML | HEPARIN SODIUM 10,000 UNITS<br>IN SODIUM CHLORIDE 0.9%<br>IN PLASTIC CONTAINER<br>(INJECTABLE; INJECTION) | ABBOTT LABORATORIES   | 18-916<br>01-31-84               |                                 |                                  |
| HEPARIN SODIUM; SODIUM CHLORIDE<br>5,000 UNITS/100ML; 900MG/100ML  | HEPARIN SODIUM 12,500 UNITS<br>IN SODIUM CHLORIDE 0.9%<br>IN PLASTIC CONTAINER<br>(INJECTABLE; INJECTION) | ABBOTT LABORATORIES   | 18-916<br>01-31-84               |                                 |                                  |
| HEPARIN SODIUM; SODIUM CHLORIDE<br>100 UNITS/ML; 4.5MG/ML          | HEPARIN SODIUM 5,000 UNITS<br>IN SODIUM CHLORIDE 0.45%<br>IN PLASTIC CONTAINER<br>(INJECTABLE; INJECTION) | ABBOTT LABORATORIES   | 18-916<br>01-31-84               |                                 |                                  |
| HEPARIN SODIUM; SODIUM CHLORIDE<br>100 UNITS/ML; 4.5MG/ML          | HEPARIN SODIUM 5,000 UNITS<br>IN SODIUM CHLORIDE 0.45%<br>IN PLASTIC CONTAINER<br>(INJECTABLE; INJECTION) | ABBOTT LABORATORIES   | 18-911<br>01-30-85               |                                 |                                  |

**TABLE IV. NDA'S APPROVED FROM 1-1-82 TO 2-28-85 AND NDA'S WITH APPROPRIATE PATENT AND EXCLUSIVITY INFORMATION**

| <u>ACTIVE INGREDIENT(S)<br/>STRENGTH(S)</u>                        | <u>TRADE NAME<br/>(DOSAGE FORM; ROUTE)</u>   | <u>APPLICANT NAME</u> | <u>NDA NO.<br/>APPROVAL DATE</u> | <u>PATENT NO.<br/>EXP. DATE</u> | <u>EXCLUSIVITY<br/>EXP. DATE</u> |
|--|--|-----------------------|----------------------------------|---------------------------------|----------------------------------|
| HEPARIN SODIUM; SODIUM CHLORIDE<br>10,000 UNITS/100ML; 450MG/100ML | HEPARIN SODIUM 10,000 UNITS<br>IN SODIUM CHLORIDE 0.45%<br>IN PLASTIC CONTAINER<br>(INJECTABLE; INJECTION) | ABBOTT LABORATORIES   | 18-916<br>01-31-84               |                                 |                                  |
| HEPARIN SODIUM; SODIUM CHLORIDE<br>10,000 UNITS/100ML; 450MG/100ML | HEPARIN SODIUM 10,000 UNITS<br>IN SODIUM CHLORIDE 0.45%<br>(INJECTABLE; INJECTION)                         | ABBOTT LABORATORIES   | 18-911<br>01-30-85               |                                 |                                  |
| HEPARIN SODIUM; SODIUM CHLORIDE<br>10,000 UNITS/100ML; 900MG/100ML | HEPARIN SODIUM 10,000 UNITS<br>IN SODIUM CHLORIDE 0.9%<br>(INJECTABLE; INJECTION)                          | ABBOTT LABORATORIES   | 18-911<br>01-30-85               |                                 |                                  |
| HEPARIN SODIUM; SODIUM CHLORIDE<br>5,000 UNITS/100ML; 450MG/100ML  | HEPARIN SODIUM 12,500 UNITS<br>IN SODIUM CHLORIDE 0.45%<br>IN PLASTIC CONTAINER<br>(INJECTABLE; INJECTION) | ABBOTT LABORATORIES   | 18-916<br>01-31-84               |                                 |                                  |
| HEPARIN SODIUM; SODIUM CHLORIDE<br>5,000 UNITS/100ML; 900MG/100ML  | HEPARIN SODIUM 12,500 UNITS<br>IN SODIUM CHLORIDE 0.9%<br>(INJECTABLE; INJECTION)                          | ABBOTT LABORATORIES   | 18-911<br>01-30-85               |                                 |                                  |
| HEPARIN SODIUM; SODIUM CHLORIDE<br>5,000 UNITS/100ML; 900MG/100ML  | HEPARIN SODIUM 25,000 UNITS<br>IN SODIUM CHLORIDE 0.9%<br>(INJECTABLE; INJECTION)                          | ABBOTT LABORATORIES   | 18-911<br>01-30-85               |                                 |                                  |
| HEPARIN SODIUM; SODIUM CHLORIDE<br>5,000 UNITS/100ML; 450MG/100ML  | HEPARIN SODIUM 25,000 UNITS<br>IN SODIUM CHLORIDE 0.45%<br>IN PLASTIC CONTAINER<br>(INJECTABLE; INJECTION) | ABBOTT LABORATORIES   | 18-916<br>01-31-84               |                                 |                                  |
| HEPARIN SODIUM; SODIUM CHLORIDE<br>10,000 UNITS/100ML; 450MG/100ML | HEPARIN SODIUM 25,000 UNITS<br>IN SODIUM CHLORIDE 0.45%<br>IN PLASTIC CONTAINER<br>(INJECTABLE; INJECTION) | ABBOTT LABORATORIES   | 18-916<br>01-31-84               |                                 |                                  |
| HEPARIN SODIUM; SODIUM CHLORIDE<br>5,000 UNITS/100ML; 900MG/100ML  | HEPARIN SODIUM 25,000 UNITS<br>IN SODIUM CHLORIDE 0.9%<br>IN PLASTIC CONTAINER<br>(INJECTABLE; INJECTION)  | ABBOTT LABORATORIES   | 18-916<br>01-31-84               |                                 |                                  |

**TABLE IV. NDA'S APPROVED FROM 1-1-82 TO 2-28-85 AND NDA'S WITH APPROPRIATE PATENT AND EXCLUSIVITY INFORMATION**

| <u>ACTIVE INGREDIENT(S)<br/>STRENGTH(S)</u>                | <u>TRADE NAME<br/>(DOSAGE FORM; ROUTE)</u> | <u>APPLICANT NAME</u> | <u>NDA NO.<br/>APPROVAL DATE</u> | <u>PATENT NO.<br/>EXP. DATE</u>            | <u>EXCLUSIVITY<br/>EXP. DATE</u> |
|--|--|-----------------------|----------------------------------|--|----------------------------------|
| HEXACHLOROPHENE<br>3%                                      | TURGEX<br>(SOLUTION; TOPICAL)              | XTRTRIUM LABS         | 19-055<br>11-30-84               |  |                                  |
| HYDROCHLOROTHIAZIDE;<br>METOPROLOL TARTRATE<br>25MG; 50MG  | LOPRESSOR HCT 50/25<br>(TABLET; ORAL)      | GEIGY/CIBA-GEIGY      | 18-303<br>12-31-84               | 3876802<br>04-08-92<br>3998790<br>12-21-93 | NC<br>12-31-87                   |
| HYDROCHLOROTHIAZIDE;<br>METOPROLOL TARTRATE<br>25MG; 100MG | LOPRESSOR HCT 100/25<br>(TABLET; ORAL)     | GEIGY/CIBA-GEIGY      | 18-303<br>12-31-84               | 3876802<br>04-08-92<br>3998790             | NC<br>12-31-87                   |
| HYDROCHLOROTHIAZIDE;<br>METOPROLOL TARTRATE<br>50MG; 100MG | LOPRESSOR HCT 100/50<br>(TABLET; ORAL)     | GEIGY/CIBA-GEIGY      | 18-303<br>12-31-84               | 3876802<br>04-08-92<br>3998790<br>12-21-93 | NC<br>12-31-87                   |
| HYDROCHLOROTHIAZIDE; TIMOLOL MALEATE<br>25MG; 10MG         | TIMOLIDE<br>(TABLET; ORAL)                 | MS&D/MERCK            | 18-061<br>12-11-81               | 3655663<br>04-11-89<br>4238485<br>12-09-97 |                                  |
| HYDROCHLOROTHIAZIDE; TRIAMTERENE<br>50MG; 75MG             | MAXZIDE<br>(TABLET; ORAL)                  | MYLAN PHARMS          | 19-129<br>10-22-84               | 4444769<br>04-24-01                        | NS<br>10-22-87                   |
| HYDROCORTISONE ACETATE<br>10%                              | CORTIFOAM<br>(AEROSOL; RECTAL)             | REED&CARNRICK PHARMS  | 17-351<br>02-10-82               |  | NDF<br>09-24-86                  |
| HYDROCORTISONE BUTYRATE<br>0.1%                            | LOCOID<br>(CREAM; TOPICAL)                 | OWEN LABS/DERM PRODS  | 18-795<br>01-07-83               |  | NP<br>09-24-86                   |
| HYDROCORTISONE BUTYRATE<br>0.1%                            | LOCOID<br>(OINTMENT; TOPICAL)              | OWEN LABS/DERM PRODS  | 19-106<br>07-03-84               |  | NP<br>09-24-86                   |
| HYDROCORTISONE VALERATE<br>0.2%                            | WESTCORT<br>(OINTMENT; TOPICAL)            | WESTWOOD PHARMS       | 18-726<br>08-08-83               |  | NDF<br>09-24-86                  |
| HYDROMORPHONE HYDROCHLORIDE<br>10MG/ML                     | DILAUIDID-HP<br>(INJECTABLE; INJECTION)    | KNOLL PHARMACEUTICAL  | 19-034<br>01-11-84               |  | NCE<br>01-11-94                  |

**TABLE IV. NDA'S APPROVED FROM 1-1-82 TO 2-28-85 AND NDA'S WITH APPROPRIATE PATENT AND EXCLUSIVITY INFORMATION**

| <u>ACTIVE INGREDIENT(S)</u><br><u>STRENGTH(S)</u> | <u>TRADE NAME</u><br><u>(DOSAGE FORM; ROUTE)</u>     | <u>APPLICANT NAME</u> | <u>NDA NO.</u><br><u>APPROVAL DATE</u> | <u>PATENT NO.</u><br><u>EXP. DATE</u> | <u>EXCLUSIVITY</u><br><u>EXP. DATE</u> |
|---|--|-----------------------|--|---------------------------------------|--|
| HYDROXYUREA<br>500MG                              | HYDREA<br>(CAPSULE; ORAL)                            | ER SQUIBB AND SONS    | 16-295<br>12-07-67                     | 3968249<br>07-06-93                   |  |
| IBUPROFEN<br>400MG                                | MOTRIN<br>(TABLET; ORAL)                             | UPJOHN MANUFACTURING  | 17-463<br>09-19-74                     | 3385886<br>05-28-85                   | I-2<br>09-24-86                        |
| IBUPROFEN<br>300MG                                | MOTRIN<br>(TABLET; ORAL)                             | UPJOHN MANUFACTURING  | 17-463<br>09-19-74                     | 3385886<br>05-28-85                   | I-2<br>09-24-86                        |
| IBUPROFEN<br>600MG                                | MOTRIN<br>(TABLET; ORAL)                             | UPJOHN MANUFACTURING  | 17-463<br>03-09-79                     | 3385886<br>05-28-85                   | I-2<br>09-24-86                        |
| IBUPROFEN<br>400MG                                | RUFEN<br>(TABLET; ORAL)                              | BOOTS PHARMACEUTICAL  | 18-197<br>05-19-81                     | 3385886<br>05-28-85                   | I-2<br>09-24-86                        |
| IBUPROFEN<br>600MG                                | RUFEN<br>(TABLET; ORAL)                              | BOOTS PHARMACEUTICAL  | 18-197<br>03-05-84                     | 3385886<br>05-28-85                   | I-2<br>09-24-86                        |
| INDAPAMIDE<br>2.5MG                               | LOZOL<br>(TABLET; ORAL)                              | USV PHARMACEUTICAL    | 18-538<br>07-06-83                     | 3565911<br>02-23-88                   | NCE<br>07-06-93                        |
| INDOMETHACIN<br>50MG                              | INDOCIN<br>(SUPPOSITORY; RECTAL)                     | MS&D RES LABS/MERCK   | 17-814<br>08-13-84                     |                                       | NDF<br>09-24-86                        |
| INDOMETHACIN<br>75MG                              | INDOCIN SR<br>(CAPSULE, CONTROLLED<br>RELEASE; ORAL) | MS&D/MERCK            | 18-185<br>02-23-82                     |                                       | NDF<br>09-24-86                        |
| INDOMETHACIN<br>25MG                              | INDOMETHACIN<br>(CAPSULE; ORAL)                      | CHELSEA LABORATORIES  | 18-690<br>07-31-84                     |                                       |  |
| INDOMETHACIN<br>50MG                              | INDOMETHACIN<br>(CAPSULE; ORAL)                      | CHELSEA LABORATORIES  | 18-690<br>07-31-84                     |                                       |  |
| INDOMETHACIN<br>25MG                              | INDOMETHACIN<br>(CAPSULE; ORAL)                      | ZENITH LABORATORIES   | 18-730<br>05-04-84                     |                                       |  |
| INDOMETHACIN<br>50MG                              | INDOMETHACIN<br>(CAPSULE; ORAL)                      | ZENITH LABORATORIES   | 18-730<br>05-04-84                     |                                       |  |

TABLE IV. NDA'S APPROVED FROM 1-1-82 TO 2-28-85 AND NDA'S WITH APPROPRIATE PATENT AND EXCLUSIVITY INFORMATION

| <u>ACTIVE INGREDIENT(S)<br/>STRENGTH(S)</u>        | <u>TRADE NAME<br/>(DOSAGE FORM; ROUTE)</u> | <u>APPLICANT NAME</u> | <u>NDA NO.<br/>APPROVAL DATE</u> | <u>PATENT NO.<br/>EXP. DATE</u> | <u>EXCLUSIVITY<br/>EXP. DATE</u> |
|--|--|-----------------------|----------------------------------|---------------------------------|----------------------------------|
| INDOMETHACIN<br>50MG                               | INDOMETHACIN<br>(CAPSULE; ORAL)            | PAR PHARMACEUTICAL    | 18-829<br>08-06-84               |                                 |                                  |
| INDOMETHACIN<br>25MG                               | INDOMETHACIN<br>(CAPSULE; ORAL)            | PAR PHARMACEUTICAL    | 18-829<br>08-06-84               |                                 |                                  |
| INDOMETHACIN<br>25MG                               | INDOMETHACIN<br>(CAPSULE; ORAL)            | LEDERLE LABS/AM CYAN  | 18-851<br>05-18-84               |                                 |                                  |
| INDOMETHACIN<br>50MG                               | INDOMETHACIN<br>(CAPSULE; ORAL)            | LEDERLE LABS/AM CYAN  | 18-851<br>05-18-84               |                                 |                                  |
| INDOMETHACIN<br>25MG                               | INDOMETHACIN<br>(CAPSULE; ORAL)            | MYLAN PHARMS          | 18-858<br>04-20-84               |                                 |                                  |
| INDOMETHACIN<br>50MG                               | INDOMETHACIN<br>(CAPSULE; ORAL)            | MYLAN PHARMS          | 18-858<br>04-20-84               |                                 |                                  |
| INDOMETHACIN<br>25MG                               | INDOMETHACIN<br>(CAPSULE; ORAL)            | PARKE-DAVIS/M-L       | 18-806<br>11-23-84               |                                 |                                  |
| INDOMETHACIN<br>50MG                               | INDOMETHACIN<br>(CAPSULE; ORAL)            | PARKE-DAVIS/M-L       | 18-806<br>11-23-84               |                                 |                                  |
| INDOMETHACIN SODIUM TRIHYDRATE<br>EQ 1MG BASE/VIAL | INDOCIN I. V.<br>(INJECTABLE; INJECTION)   | MS&D/MERCK            | 18-878<br>01-30-85               |                                 | I-6<br>09-24-86                  |
| IODAMIDE MEGLUMINE<br>24%                          | RENOVUE-DIP<br>(INJECTABLE; INJECTION)     | ER SQUIBB AND SONS    | 17-903<br>07-10-78               |                                 |                                  |
| IODAMIDE MEGLUMINE<br>65%                          | RENOVUE-65<br>(INJECTABLE; INJECTION)      | ER SQUIBB AND SONS    | 17-902<br>07-24-78               |                                 | I-6<br>09-24-86                  |
| IODOHIPPURATE SODIUM,<br>I-123<br>1MCI/ML          | NEPHROFLOW<br>(INJECTABLE; INJECTION)      | MEDI-PHYSICS          | 18-289<br>12-28-84               |                                 | NCE<br>12-28-89                  |
| IODOXAMATE MEGLUMINE<br>9.9%                       | CHOLOVUE<br>(INJECTABLE; INJECTION)        | ER SQUIBB AND SONS    | 18-076<br>08-14-81               | 3654272<br>04-04-89             |                                  |

**TABLE IV. NDA'S APPROVED FROM 1-1-82 TO 2-28-85 AND NDA'S WITH APPROPRIATE PATENT AND EXCLUSIVITY INFORMATION**

| <u>ACTIVE INGREDIENT(S)<br/>STRENGTH(S)</u> | <u>TRADE NAME<br/>(DOSAGE FORM; ROUTE)</u> | <u>APPLICANT NAME</u> | <u>NDA NO.<br/>APPROVAL DATE</u> | <u>PATENT NO.<br/>EXP. DATE</u>                                   | <u>EXCLUSIVITY<br/>EXP. DATE</u> |
|---|--|-----------------------|----------------------------------|---|----------------------------------|
| IDOXYAMATE MEGLUMINE<br>40.3%               | CHOLOVUE<br>(INJECTABLE; INJECTION)        | ER SQUIBB AND SONS    | 18-077<br>08-14-81               | 3654272<br>04-04-89   |                                  |
| ISOFLURANE<br>99.9%                         | FORANE<br>(GAS; INHALATION)                | ANAQUEST/BOC          | 17-624<br>12-18-79               | 3535425<br>01-24-93<br>3535388<br>01-24-93                        |                                  |
| ISOTRETINOIN<br>10MG                        | ACCUTANE<br>(CAPSULE; ORAL)                | HOFFMANN-LA ROCHE     | 18-662<br>05-07-82               | 4200647<br>04-29-97<br>4322438<br>03-30-99<br>4464394<br>08-07-01 | NCE<br>05-07-92                  |
| ISOTRETINOIN<br>20MG                        | ACCUTANE<br>(CAPSULE; ORAL)                | HOFFMANN-LA ROCHE     | 18-662<br>03-28-83               | 4200647<br>04-29-97<br>4322438<br>03-30-99<br>4464394<br>08-07-01 | NCE<br>05-07-92                  |
| ISOTRETINOIN<br>40MG                        | ACCUTANE<br>(CAPSULE; ORAL)                | HOFFMANN-LA ROCHE     | 18-662<br>05-07-82               | 4200647<br>04-29-97<br>4322438<br>03-30-99<br>4464394<br>08-07-01 | NCE<br>05-07-92                  |
| KETOCONAZOLE<br>200MG                       | NIZORAL<br>(TABLET; ORAL)                  | JANSSEN PHARMA        | 18-533<br>06-12-81               | 4335125<br>06-15-99   | I-25<br>09-24-86                 |
| LABELALOL HYDROCHLORIDE<br>200MG            | NORMODYNE<br>(TABLET; ORAL)                | SCHERING              | 18-686<br>08-01-84               | 4012444<br>03-15-94<br>4006755<br>01-03-95                        | NCE<br>08-01-94                  |
| LABELALOL HYDROCHLORIDE<br>300MG            | NORMODYNE<br>(TABLET; ORAL)                | SCHERING              | 18-686<br>08-01-84               | 4012444<br>03-15-94<br>4006755<br>01-03-95                        | NCE<br>08-01-94                  |

TABLE IV. NDA'S APPROVED FROM 1-1-82 TO 2-28-85 AND NDA'S WITH APPROPRIATE PATENT AND EXCLUSIVITY INFORMATION

| <u>ACTIVE INGREDIENT(S)<br/>STRENGTH(S)</u> | <u>TRADE NAME<br/>(DOSAGE FORM; ROUTE)</u> | <u>APPLICANT NAME</u> | <u>NDA NO.<br/>APPROVAL DATE</u> | <u>PATENT NO.<br/>EXP. DATE</u>  | <u>EXCLUSIVITY<br/>EXP. DATE</u> |
|---|--|-----------------------|----------------------------------|--|----------------------------------|
| LABETALOL HYDROCHLORIDE<br>400MG            | NORMODYNE<br>(TABLET; ORAL)                | SCHERING              | 18-686<br>08-01-84               | 4012444<br>03-15-94<br>4006755<br>01-03-95   | NCE<br>08-01-94                  |
| LABETALOL HYDROCHLORIDE<br>5MG/ML           | NORMODYNE<br>(INJECTABLE; INJECTION)       | SCHERING              | 18-687<br>08-01-84               | 4012444<br>03-15-94<br>4006755<br>01-03-95<br>4328213<br>05-04-99  | NCE<br>08-01-94                  |
| LABETALOL HYDROCHLORIDE<br>200MG            | TRANDATE<br>(TABLET; ORAL)                 | GLAXO                 | 18-716<br>08-01-84               | 4012444<br>03-15-94<br>4006755<br>01-03-95   | NCE<br>08-01-94                  |
| LABETALOL HYDROCHLORIDE<br>300MG            | TRANDATE<br>(TABLET; ORAL)                 | GLAXO                 | 18-716<br>08-01-84               | 4012444<br>03-15-94<br>4006755<br>01-03-95   | NCE<br>08-01-94                  |
| LABETALOL HYDROCHLORIDE<br>400MG            | TRANDATE<br>(TABLET; ORAL)                 | GLAXO                 | 18-716<br>08-01-84               | 4012444<br>03-15-94<br>4006755<br>01-03-95   | NCE<br>08-01-94                  |
| LACTULOSE<br>10GM/15ML                      | CEPHULAC<br>(SYRUP; ORAL)                  | MERRELL DOW/DOW CHEM  | 17-657<br>03-25-76               | 3461204<br>08-12-86<br>3867524<br>02-18-92<br>3860708<br>01-14-92<br>3860707<br>01-14-92<br>3562388<br>02-09-88<br>3558774<br>01-26-88 |                                  |

**TABLE IV. NDA'S APPROVED FROM 1-1-82 TO 2-28-85 AND NDA'S WITH APPROPRIATE PATENT AND EXCLUSIVITY INFORMATION**

| <u>ACTIVE INGREDIENT(S)<br/>STRENGTH(S)</u> | <u>TRADE NAME<br/>(DOSAGE FORM; ROUTE)</u>           | <u>APPLICANT NAME</u> | <u>NDA NO.<br/>APPROVAL DATE</u> | <u>PATENT NO.<br/>EXP. DATE</u>            | <u>EXCLUSIVITY<br/>EXP. DATE</u> |
|---|--|-----------------------|----------------------------------|--|----------------------------------|
| LEUCOVORIN CALCIUM<br>EQ 5MG BASE           | WELLCOVORIN<br>(TABLET; ORAL)                        | BURROUGHS WELLCOME    | 18-342<br>07-08-83               |  | NDF<br>09-24-86                  |
| LEUCOVORIN CALCIUM<br>EQ 25MG BASE          | WELLCOVORIN<br>(TABLET; ORAL)                        | BURROUGHS WELLCOME    | 18-342<br>07-08-83               |  | NDF<br>09-24-86                  |
| LITHIUM CARBONATE<br>450MG                  | ESKALITH CR<br>(TABLET, CONTROLLED<br>RELEASE; ORAL) | SK&F LABORATORIES     | 18-152<br>03-29-82               |  | NS<br>09-24-86                   |
| LITHIUM CARBONATE<br>300MG                  | LITHIUM CARBONATE<br>(TABLET; ORAL)                  | ROXANE LABORATORIES   | 18-558<br>01-29-82               |  |                                  |
| LOPERAMIDE HYDROCHLORIDE<br>2MG             | IMODIUM<br>(CAPSULE; ORAL)                           | JANSSEN PHARMA        | 17-694<br>12-28-76               | 3714159<br>01-30-90                        | I-30<br>09-24-86                 |
| LOPERAMIDE HYDROCHLORIDE<br>1MG/5ML         | IMODIUM<br>(SOLUTION; ORAL)                          | JANSSEN PHARMA        | 19-037<br>07-31-84               | 3714159<br>01-30-90                        | NDF<br>09-24-86                  |
| LOXAPINE HYDROCHLORIDE<br>EQ 50MG BASE/ML   | LOXITANE<br>(INJECTABLE; INJECTION)                  | LEDERLE LABS/AM CYAN  | 18-039<br>10-26-79               | 3546226<br>12-08-87                        |                                  |
| LOXAPINE HYDROCHLORIDE<br>EQ 25MG BASE/ML   | LOXITANE<br>(CONCENTRATE; ORAL)                      | LEDERLE LABS/AM CYAN  | 17-658<br>05-04-76               | 3546226<br>12-08-87<br>4049809<br>09-20-94 |                                  |
| LOXAPINE SUCCINATE<br>EQ 5MG BASE           | LOXITANE<br>(CAPSULE; ORAL)                          | LEDERLE LABS/AM CYAN  | 17-525<br>10-25-77               | 3546226<br>12-08-87                        |                                  |
| LOXAPINE SUCCINATE<br>EQ 10MG BASE          | LOXITANE<br>(CAPSULE; ORAL)                          | LEDERLE LABS/AM CYAN  | 17-525<br>02-25-75               | 3546226<br>12-08-87                        |                                  |
| LOXAPINE SUCCINATE<br>EQ 25MG BASE          | LOXITANE<br>(CAPSULE; ORAL)                          | LEDERLE LABS/AM CYAN  | 17-525<br>02-25-75               | 3546226<br>12-08-87                        |                                  |
| LOXAPINE SUCCINATE<br>EQ 50MG BASE          | LOXITANE<br>(CAPSULE; ORAL)                          | LEDERLE LABS/AM CYAN  | 17-525<br>02-25-75               | 3546226<br>12-08-87                        |                                  |

TABLE IV. NDA'S APPROVED FROM 1-1-82 TO 2-28-85 AND NDA'S WITH APPROPRIATE PATENT AND EXCLUSIVITY INFORMATION

| <u>ACTIVE INGREDIENT(S)<br/>STRENGTH(S)</u>   | <u>TRADE NAME<br/>(DOSAGE FORM; ROUTE)</u>                         | <u>APPLICANT NAME</u> | <u>NDA NO.<br/>APPROVAL DATE</u> | <u>PATENT NO.<br/>EXP. DATE</u> | <u>EXCLUSIVITY<br/>EXP. DATE</u> |
|---|--|-----------------------|----------------------------------|---------------------------------|----------------------------------|
| MAFENIDE ACETATE<br>EQ 85MG BASE/GM   | SULFAMYLON<br>(CREAM; TOPICAL)                                     | WINTHROP LABS/STERL   | 16-763<br>01-24-69               | 3497599<br>01-26-88             |                                  |
| MAGNESIUM ACETATE TETRAHYDRATE; POTASSIUM<br>ACETATE; SODIUM CHLORIDE<br>32MG/100ML; 128MG/100ML; 234MG/100ML   | PLASMA-LYTE 56 IN PLASTIC<br>CONTAINER<br>(INJECTABLE; INJECTION)  | TRAVENOL LABS         | 19-047<br>06-15-84               |                                 | NC<br>09-24-86                   |
| MAGNESIUM CHLORIDE; POTASSIUM CHLORIDE;<br>POTASSIUM PHOSPHATE, MONOBASIC; SODIUM<br>ACETATE; SODIUM CHLORIDE; SODIUM GLUCONATE;<br>SODIUM PHOSPHATE<br>30MG/100ML; 37MG/100ML; 0.82MG/100ML;<br>370MG/100ML; 530MG/100ML; 500MG/100ML;<br>12MG/100ML | ISOLYTES PH 7.4 IN PLASTIC<br>CONTAINER<br>(INJECTABLE; INJECTION) | AM MCGAW/AM HOSP      | 19-006<br>04-04-84               |                                 | NC<br>09-24-86                   |
| MAGNESIUM CHLORIDE; POTASSIUM CHLORIDE;<br>SODIUM ACETATE; SODIUM CHLORIDE; SODIUM<br>GLUCONATE<br>30MG/100ML; 37MG/100ML; 222MG/100ML;<br>526MG/100ML; 502MG/100ML   | PHYSIOSOL IN PLASTIC<br>CONTAINER<br>(SOLUTION; IRRIGATION)        | ABBOTT LABORATORIES   | 17-637<br>07-08-82               |                                 | NC<br>09-24-86                   |
| MAGNESIUM CHLORIDE; POTASSIUM CHLORIDE;<br>SODIUM ACETATE; SODIUM CHLORIDE; SODIUM<br>GLUCONATE<br>30MG/100ML; 37MG/100ML; 222MG/100ML;<br>526MG/100ML; 502MG/100ML   | PHYSIOSOL IN PLASTIC<br>CONTAINER<br>(SOLUTION; IRRIGATION)        | ABBOTT LABORATORIES   | 18-406<br>07-08-82               |                                 | NC<br>09-24-86                   |
| MAGNESIUM CHLORIDE; POTASSIUM CHLORIDE;<br>SODIUM ACETATE; SODIUM CHLORIDE; SODIUM<br>GLUCONATE<br>30MG/100ML; 37MG/100ML; 370MG/100ML;<br>530MG/100ML; 500MG/100ML   | PHYSIOLYTE IN PLASTIC<br>CONTAINER<br>(SOLUTION; IRRIGATION)       | AM MCGAW/AM HOSP      | 19-024<br>06-08-84               |                                 | NC<br>09-24-86                   |
| MAGNESIUM CHLORIDE; POTASSIUM CHLORIDE;<br>SODIUM ACETATE; SODIUM CHLORIDE;<br>SODIUM GLUCONATE<br>30MG/100ML; 37MG/100ML; 368MG/100ML;<br>526MG/100ML; 502MG/100ML   | SYNOVALYTE<br>IN PLASTIC CONTAINER<br>(SOLUTION; IRRIGATION)       | TRAVENOL LABS         | 19-326<br>01-25-85               |                                 |                                  |

TABLE IV. NDA'S APPROVED FROM 1-1-82 TO 2-28-85 AND NDA'S WITH APPROPRIATE PATENT AND EXCLUSIVITY INFORMATION

| <u>ACTIVE INGREDIENT(S)</u><br><u>STRENGTH(S)</u>  | <u>TRADE NAME</u><br><u>(DOSAGE FORM; ROUTE)</u> | <u>APPLICANT NAME</u> | <u>NDA NO.</u><br><u>APPROVAL DATE</u> | <u>PATENT NO.</u><br><u>EXP. DATE</u> | <u>EXCLUSIVITY</u><br><u>EXP. DATE</u> |
|--|--|-----------------------|--|---------------------------------------|--|
| MAGNESIUM SULFATE; POTASSIUM CHLORIDE;<br>POTASSIUM PHOSPHATE, MONOBASIC; SODIUM<br>CHLORIDE; SODIUM PHOSPHATE<br>20MG/100ML; 40MG/100ML; 6.25MG/100ML;<br>800MG/100ML; 8.75MG/100ML | TIS-U-SOL<br>(SOLUTION; IRRIGATION)              | TRAVENOL LABS         | 18-508<br>02-19-82                     |                                       | NC<br>09-24-86                         |
| MALATHION<br>0.5%  | PRIODERM<br>(LOTION; TOPICAL)                    | PURDUE FREDERICK      | 18-613<br>08-02-82                     |                                       | NCE<br>08-02-92                        |
| MAPROTILINE HYDROCHLORIDE<br>25MG  | LUDIOMIL<br>(TABLET; ORAL)                       | CIBA/CIBA-GEIGY       | 17-543<br>12-01-80                     | 3399201<br>08-27-85                   |  |
| MAPROTILINE HYDROCHLORIDE<br>50MG  | LUDIOMIL<br>(TABLET; ORAL)                       | CIBA/CIBA-GEIGY       | 17-543<br>12-01-80                     | 3399201<br>08-27-85                   |  |
| MAPROTILINE HYDROCHLORIDE<br>75MG  | LUDIOMIL<br>(TABLET; ORAL)                       | CIBA/CIBA-GEIGY       | 17-543<br>09-30-82                     | 3399201<br>08-27-85                   | NS<br>09-24-86                         |
| MAZINDOL<br>1MG  | SANOREX<br>(TABLET; ORAL)                        | SANDOZ PHARMS/SANDOZ  | 17-247<br>06-14-73                     | 3763178<br>10-02-90                   |  |
| MAZINDOL<br>2MG  | SANOREX<br>(TABLET; ORAL)                        | SANDOZ PHARMS/SANDOZ  | 17-247<br>06-14-73                     | 3763178<br>10-02-90                   |  |
| MAZINDOL<br>2MG  | MAZANOR<br>(TABLET; ORAL)                        | WYETH LABS/AMHO       | 17-980<br>08-28-80                     | 3763178<br>10-02-90                   |  |
| MAZINDOL<br>1MG  | MAZANOR<br>(TABLET; ORAL)                        | WYETH LABS/AMHO       | 17-980<br>02-02-82                     | 3763178<br>10-02-90                   |  |
| MEBENDAZOLE<br>100MG   | VERMOX<br>(TABLET, CHEWABLE; ORAL)               | JANSSEN PHARMA        | 17-481<br>06-28-74                     | 3657267<br>04-18-89                   |  |
| MEDROXYPROGESTERONE ACETATE<br>100MG/ML  | DEPO-PROVERA<br>(INJECTABLE; INJECTION)          | UPJOHN                | 12-541<br>01-16-76                     | 3377364<br>04-09-85                   |  |
| MEDROXYPROGESTERONE ACETATE<br>400MG/ML  | DEPO-PROVERA<br>(INJECTABLE; INJECTION)          | UPJOHN                | 12-541<br>01-16-76                     | 3377364<br>04-09-85                   |  |

**TABLE IV. NDA'S APPROVED FROM 1-1-82 TO 2-28-85 AND NDA'S WITH APPROPRIATE PATENT AND EXCLUSIVITY INFORMATION**

| <u>ACTIVE INGREDIENT(S)<br/>STRENGTH(S)</u>         | <u>TRADE NAME<br/>(DOSAGE FORM; ROUTE)</u>          | <u>APPLICANT NAME</u> | <u>NDA NO.<br/>APPROVAL DATE</u> | <u>PATENT NO.<br/>EXP. DATE</u> | <u>EXCLUSIVITY<br/>EXP. DATE</u> |
|---|---|-----------------------|----------------------------------|---------------------------------|----------------------------------|
| MEGLUMINE; METRIZOIC ACID<br>140.1MG/ML; 461.8MG/ML | ISOPAQUE-280<br>(INJECTABLE; INJECTION)             | WINTHROP LABS/STERL   | 17-506<br>04-30-74               | 3476802<br>11-04-86             |                                  |
| METAPROTERENOL SULFATE<br>20MG                      | ALUPENT<br>(TABLET; ORAL)                           | BOEHRINGER INGELHEIM  | 15-874<br>05-13-74               | 3422196<br>01-14-86             |                                  |
| METAPROTERENOL SULFATE<br>10MG                      | ALUPENT<br>(TABLET; ORAL)                           | BOEHRINGER INGELHEIM  | 15-874<br>08-08-77               | 3422196<br>01-14-86             |                                  |
| METAPROTERENOL SULFATE<br>0.65MG/INH                | ALUPENT<br>(AEROSOL; INHALATION)                    | BOEHRINGER INGELHEIM  | 16-402<br>07-31-73               | 3422196<br>01-14-86             |                                  |
| METAPROTERENOL SULFATE<br>10MG/5ML                  | ALUPENT<br>(SYRUP; ORAL)                            | BOEHRINGER INGELHEIM  | 17-571<br>05-23-75               | 3422196<br>01-14-86             |                                  |
| METAPROTERENOL SULFATE<br>5%                        | ALUPENT<br>(SOLUTION; INHALATION)                   | BOEHRINGER INGELHEIM  | 17-659<br>09-18-80               | 3422196<br>01-14-86             |                                  |
| METAPROTERENOL SULFATE<br>0.6%                      | ALUPENT<br>(SOLUTION; INHALATION)                   | BOEHRINGER INGELHEIM  | 18-761<br>06-30-83               | 3422196<br>01-14-86             |                                  |
| METHYLDOPA<br>250MG                                 | METHYLDOPA<br>(TABLET; ORAL)                        | CORD LABORATORIES     | 18-934<br>06-29-84               |                                 |                                  |
| METHYLDOPA<br>500MG                                 | METHYLDOPA<br>(TABLET; ORAL)                        | CORD LABORATORIES     | 18-934<br>06-29-84               |                                 |                                  |
| METHYLPHENIDATE HYDROCHLORIDE<br>20MG               | RITALIN-SR<br>(TABLET, CONTROLLED<br>RELEASE; ORAL) | CIBA/CIBA-GEIGY       | 18-029<br>03-30-82               |                                 | NDF<br>09-24-86                  |
| METOCLOPRAMIDE<br>EQ 5MG BASE/5ML                   | REGLAN<br>(SYRUP; ORAL)                             | AH ROBINS             | 18-821<br>3-25-83                |                                 | NDF<br>09-24-86                  |
| METOCLOPRAMIDE HYDROCHLORIDE<br>EQ 5MG BASE/ML      | REGLAN<br>(INJECTABLE; INJECTION)                   | AH ROBINS             | 17-862<br>02-07-79               |                                 | I-12; I-13;<br>I-14<br>09-24-87  |
| METOCLOPRAMIDE HYDROCHLORIDE<br>EQ 10MG BASE        | REGLAN<br>(TABLET; ORAL)                            | AH ROBINS             | 17-854<br>12-30-80               |                                 | I-4<br>09-24-86                  |

**TABLE IV. NDA'S APPROVED FROM 1-1-82 TO 2-28-85 AND NDA'S WITH APPROPRIATE PATENT AND EXCLUSIVITY INFORMATION**

| <u>ACTIVE INGREDIENT(S)</u><br><u>STRENGTH(S)</u> | <u>TRADE NAME</u><br><u>(DOSAGE FORM; ROUTE)</u> | <u>APPLICANT NAME</u> | <u>NDA NO.</u><br><u>APPROVAL DATE</u> | <u>PATENT NO.</u><br><u>EXP. DATE</u> | <u>EXCLUSIVITY</u><br><u>EXP. DATE</u> |
|---|--|-----------------------|--|---------------------------------------|--|
| METOPROLOL TARTRATE<br>50MG                       | LOPRESSOR<br>(TABLET; ORAL)                      | GEIGY/CIBA-GEIGY      | 17-963<br>08-07-78                     | 3998790<br>12-21-93                   |  |
| METOPROLOL TARTRATE<br>100MG                      | LOPRESSOR<br>(TABLET; ORAL)                      | GEIGY/CIBA-GEIGY      | 17-963<br>08-07-78                     | 3998790<br>12-21-93                   |  |
| METOPROLOL TARTRATE<br>1MG/ML                     | LOPRESSOR<br>(INJECTABLE; INJECTION)             | GEIGY/CIBA-GEIGY      | 18-704<br>03-30-84                     | 3998790<br>12-21-93                   | NDF<br>09-24-86                        |
| METRIZAMIDE<br>3.75GM/VIAL                        | AMIPAQUE<br>(INJECTABLE; INJECTION)              | WINTHROP LABS/STERL   | 17-982<br>08-23-78                     | 3701771<br>10-31-89                   | I-26<br>09-24-86                       |
| METRIZAMIDE<br>6.75GM/VIAL                        | AMIPAQUE<br>(INJECTABLE; INJECTION)              | WINTHROP LABS/STERL   | 17-982<br>08-23-78                     | 3701771<br>10-31-89                   | I-26<br>09-24-86                       |
| METRONIDAZOLE<br>500MG                            | METRONIDAZOLE<br>(TABLET; ORAL)                  | ZENITH LABORATORIES   | 18-517<br>05-05-82                     |                                       |  |
| METRONIDAZOLE<br>250MG                            | METRONIDAZOLE<br>(TABLET; ORAL)                  | CHELSEA LABORATORIES  | 18-599<br>09-17-82                     |                                       |  |
| METRONIDAZOLE<br>500MG                            | METRONIDAZOLE<br>(TABLET; ORAL)                  | CHELSEA LABORATORIES  | 18-599<br>02-13-84                     |                                       |  |
| METRONIDAZOLE<br>250MG                            | METRYL<br>(TABLET; ORAL)                         | DRUMMER/PHOENIX       | 18-620<br>03-04-82                     |                                       |  |
| METRONIDAZOLE<br>500MG                            | METRYL 500<br>(TABLET; ORAL)                     | DRUMMER/PHOENIX       | 18-620<br>06-02-83                     |                                       |  |
| METRONIDAZOLE<br>500MG/100ML                      | METRO I.V.<br>(INJECTABLE; INJECTION)            | AM MCGAW/AM HOSP      | 18-674<br>08-31-82                     |                                       |  |
| METRONIDAZOLE<br>250MG                            | METRONIDAZOLE<br>(TABLET; ORAL)                  | CORD LABORATORIES     | 18-740<br>10-22-82                     |                                       |  |
| METRONIDAZOLE<br>500MG                            | METRONIDAZOLE<br>(TABLET; ORAL)                  | CORD LABORATORIES     | 18-740<br>10-22-82                     |                                       |  |
| METRONIDAZOLE<br>250MG                            | METRONIDAZOLE<br>(TABLET; ORAL)                  | DANBURY PHARMACAL     | 18-764<br>09-17-82                     |                                       |  |

TABLE IV. NDA'S APPROVED FROM 1-1-82 TO 2-28-85 AND NDA'S WITH APPROPRIATE PATENT AND EXCLUSIVITY INFORMATION

| <u>ACTIVE INGREDIENT(S)<br/>STRENGTH(S)</u> | <u>TRADE NAME<br/>(DOSAGE FORM; ROUTE)</u>                         | <u>APPLICANT NAME</u> | <u>NDA NO.<br/>APPROVAL DATE</u> | <u>PATENT NO.<br/>EXP. DATE</u> | <u>EXCLUSIVITY<br/>EXP. DATE</u> |
|---|--|-----------------------|----------------------------------|---------------------------------|----------------------------------|
| METRONIDAZOLE<br>500MG                      | METRONIDAZOLE<br>(TABLET; ORAL)                                    | DANBURY PHARMACAL     | 18-764<br>12-20-82               |                                 |                                  |
| METRONIDAZOLE<br>250MG                      | METRONIDAZOLE<br>(TABLET; ORAL)                                    | BARR LABORATORIES     | 18-818<br>02-16-83               |                                 |                                  |
| METRONIDAZOLE<br>500MG                      | METRONIDAZOLE<br>(TABLET; ORAL)                                    | BARR LABORATORIES     | 18-818<br>02-16-83               |                                 |                                  |
| METRONIDAZOLE<br>250MG                      | METRONIDAZOLE<br>(TABLET; ORAL)                                    | PAR PHARMACEUTICAL    | 18-845<br>08-18-83               |                                 |                                  |
| METRONIDAZOLE<br>250MG                      | PROTOSTAT<br>(TABLET; ORAL)  | ORTHO PHARMACEUTICAL  | 18-871<br>03-02-83               |                                 |                                  |
| METRONIDAZOLE<br>500MG                      | PROTOSTAT<br>(TABLET; ORAL)  | ORTHO PHARMACEUTICAL  | 18-871<br>03-02-83               |                                 |                                  |
| METRONIDAZOLE<br>500MG/100ML                | METRONIDAZOLE<br>(INJECTABLE; INJECTION)                           | ABBOTT LABORATORIES   | 18-889<br>11-18-83               |                                 |                                  |
| METRONIDAZOLE<br>500MG/100ML                | METRONIDAZOLE IN PLASTIC<br>CONTAINER<br>(INJECTABLE; INJECTION)   | ABBOTT LABORATORIES   | 18-890<br>11-18-83               |                                 |                                  |
| METRONIDAZOLE<br>500MG/100ML                | METRO I.V. IN PLASTIC<br>CONTAINER<br>(INJECTABLE; INJECTION)      | AM MCGAM/AM HOSP      | 18-900<br>09-29-83               |                                 |                                  |
| METRONIDAZOLE<br>500MG/100ML                | METRONIDAZOLE<br>(INJECTABLE; INJECTION)                           | ELKINS-SINN/AHROBINS  | 18-907<br>03-30-84               |                                 |                                  |
| METRONIDAZOLE<br>500MG/100ML                | FLAGYL I.V. RTU<br>(INJECTABLE; INJECTION)                         | SEARLE PHARMS         | 18-353<br>05-29-81               |                                 | I-11<br>12-20-87                 |
| METRONIDAZOLE<br>500MG/100ML                | FLAGYL I.V. RTU<br>IN PLASTIC CONTAINER<br>(INJECTABLE; INJECTION) | SEARLE PHARMS         | 18-657<br>12-24-81               |                                 | I-11<br>12-20-87                 |
| METRONIDAZOLE<br>500MG                      | METRONIDAZOLE<br>(TABLET; ORAL)                                    | PAR PHARMACEUTICAL    | 18-930<br>08-18-83               |                                 |                                  |

**TABLE IV. NDA'S APPROVED FROM 1-1-82 TO 2-28-85 AND NDA'S WITH APPROPRIATE PATENT AND EXCLUSIVITY INFORMATION**

| <u>ACTIVE INGREDIENT(S)<br/>STRENGTH(S)</u>          | <u>TRADE NAME<br/>(DOSAGE FORM; ROUTE)</u> | <u>APPLICANT NAME</u> | <u>NDA NO.<br/>APPROVAL DATE</u> | <u>PATENT NO.<br/>EXP. DATE</u>            | <u>EXCLUSIVITY<br/>EXP. DATE</u> |
|--|--|-----------------------|----------------------------------|--|----------------------------------|
| METRONIDAZOLE<br>250MG                               | METRONIDAZOLE<br>(TABLET; ORAL)            | LNK INTERNATIONAL     | 19-029<br>04-10-84               |  |                                  |
| METRONIDAZOLE<br>HYDROCHLORIDE<br>EQ 500MG BASE/VIAL | FLAGYL I.V.<br>(INJECTABLE; INJECTION)     | SEARLE PHARMS         | 18-353<br>11-28-80               |  | I-11<br>12-20-87                 |
| MICONAZOLE<br>10MG/ML                                | MONISTAT<br>(INJECTABLE; INJECTION)        | JANSSEN PHARMA        | 18-040<br>10-04-78               | 3717655<br>02-20-90<br>3839574<br>10-01-91 | I-27<br>09-24-86                 |
| MICONAZOLE NITRATE<br>2%                             | MONISTAT 7<br>(CREAM; VAGINAL)             | ORTHO PHARMACEUTICAL  | 17-450<br>01-30-74               | 3717655<br>02-20-90<br>3839574<br>10-01-91 |                                  |
| MICONAZOLE NITRATE<br>2%                             | MONISTAT-DERM<br>(CREAM; TOPICAL)          | ORTHO PHARMACEUTICAL  | 17-494<br>01-30-74               | 3717655<br>02-20-90<br>3839574<br>10-01-91 |                                  |
| MICONAZOLE NITRATE<br>2%                             | MONISTAT-DERM<br>(LOTION; TOPICAL)         | ORTHO PHARMACEUTICAL  | 17-739<br>12-16-75               | 3717655<br>02-20-90<br>3839574<br>10-01-91 |                                  |
| MICONAZOLE NITRATE<br>100MG                          | MONISTAT 7<br>(SUPPOSITORY; VAGINAL)       | ORTHO PHARMACEUTICAL  | 18-520<br>03-15-82               | 3717655<br>02-20-90<br>3839574<br>10-01-91 |                                  |
| MICONAZOLE NITRATE<br>200MG                          | MONISTAT 3<br>(SUPPOSITORY; VAGINAL)       | ORTHO PHARMACEUTICAL  | 18-888<br>08-15-84               | 3717655<br>02-20-90<br>3839574<br>10-01-91 | NS<br>09-24-86                   |
| MINOXIDIL<br>2.5MG                                   | LONITEN<br>(TABLET; ORAL)                  | UPJOHN                | 18-154<br>10-18-79               | 3461461<br>08-12-86                        |                                  |

TABLE IV. NDA'S APPROVED FROM 1-1-82 TO 2-28-85 AND NDA'S WITH APPROPRIATE PATENT AND EXCLUSIVITY INFORMATION

| <u>ACTIVE INGREDIENT(S)</u><br><u>STRENGTH(S)</u> | <u>TRADE NAME</u><br><u>(DOSAGE FORM; ROUTE)</u> | <u>APPLICANT NAME</u> | <u>NDA NO.</u><br><u>APPROVAL DATE</u> | <u>PATENT NO.</u><br><u>EXP. DATE</u>      | <u>EXCLUSIVITY</u><br><u>EXP. DATE</u> |
|---|--|-----------------------|--|--|--|
| MINOXIDIL<br>10MG                                 | LONITEN<br>(TABLET; ORAL)                        | UPJOHN                | 18-154<br>10-18-79                     | 3461461<br>08-12-86                        |  |
| MOLINDONE HYDROCHLORIDE<br>5MG                    | MOBAN<br>(TABLET; ORAL)                          | DUPONT PHARMS/DUPONT  | 17-111<br>07-03-74                     | 3491093<br>01-20-87                        |  |
| MOLINDONE HYDROCHLORIDE<br>10MG                   | MOBAN<br>(TABLET; ORAL)                          | DUPONT PHARMS/DUPONT  | 17-111<br>07-03-74                     | 3491093<br>01-20-87                        |  |
| MOLINDONE HYDROCHLORIDE<br>25MG                   | MOBAN<br>(TABLET; ORAL)                          | DUPONT PHARMS/DUPONT  | 17-111<br>07-03-74                     | 3491093<br>01-20-87                        |  |
| MOLINDONE HYDROCHLORIDE<br>50MG                   | MOBAN<br>(TABLET; ORAL)                          | DUPONT PHARMS/DUPONT  | 17-111<br>01-05-81                     | 3491093<br>01-20-87                        |  |
| MOLINDONE HYDROCHLORIDE<br>100MG                  | MOBAN<br>(TABLET; ORAL)                          | DUPONT PHARMS/DUPONT  | 17-111<br>01-05-81                     | 3491093<br>01-20-87                        |  |
| MOLINDONE HYDROCHLORIDE<br>20MG/ML                | MOBAN<br>(CONCENTRATE; ORAL)                     | DUPONT PHARMS/DUPONT  | 17-938<br>12-28-79                     | 3491093<br>01-20-87                        |  |
| MORPHINE SULFATE<br>0.5MG/ML                      | DURAMORPH PF<br>(INJECTABLE; INJECTION)          | ELKINS-SINN/AHROBINS  | 18-565<br>09-18-84                     |  | NR; D-8<br>09-24-86                    |
| MORPHINE SULFATE<br>1MG/ML                        | DURAMORPH PF<br>(INJECTABLE; INJECTION)          | ELKINS-SINN/AHROBINS  | 18-565<br>09-18-84                     |  | NR; D-8<br>09-24-86                    |
| NADOLOL<br>40MG                                   | CORGARD<br>(TABLET; ORAL)                        | ER SQUIBB AND SONS    | 18-063<br>12-10-79                     | 3982021<br>09-21-93<br>3935267<br>01-27-93 |  |
| NADOLOL<br>80MG                                   | CORGARD<br>(TABLET; ORAL)                        | ER SQUIBB AND SONS    | 18-063<br>12-10-79                     | 3982021<br>09-21-93<br>3935267<br>01-27-93 |  |
| NADOLOL<br>120MG                                  | CORGARD<br>(TABLET; ORAL)                        | ER SQUIBB AND SONS    | 18-063<br>12-10-79                     | 3982021<br>09-21-93<br>3935267<br>01-27-93 |  |

**TABLE IV. NDA'S APPROVED FROM 1-1-82 TO 2-28-85 AND NDA'S WITH APPROPRIATE PATENT AND EXCLUSIVITY INFORMATION**

| <u>ACTIVE INGREDIENT(S)<br/>STRENGTH(S)</u> | <u>TRADE NAME<br/>(DOSAGE FORM; ROUTE)</u> | <u>APPLICANT NAME</u> | <u>NDA NO.<br/>APPROVAL DATE</u> | <u>PATENT NO.<br/>EXP. DATE</u>            | <u>EXCLUSIVITY<br/>EXP. DATE</u> |
|---|--|-----------------------|----------------------------------|--|----------------------------------|
| NADOLOL<br>160MG                            | CORGARD<br>(TABLET; ORAL)                  | ER SQUIBB AND SONS    | 18-063<br>12-10-79               | 3982021<br>09-21-93<br>3935267<br>01-27-93 |                                  |
| NADOLOL<br>40MG                             | CORGARD<br>(TABLET; ORAL)                  | ER SQUIBB AND SONS    | 18-064<br>12-10-79               | 3982021<br>09-21-93<br>3935267<br>01-27-93 |                                  |
| NADOLOL<br>80MG                             | CORGARD<br>(TABLET; ORAL)                  | ER SQUIBB AND SONS    | 18-064<br>12-10-79               | 3982021<br>09-21-93<br>3935267<br>01-27-93 |                                  |
| NADOLOL<br>120MG                            | CORGARD<br>(TABLET; ORAL)                  | ER SQUIBB AND SONS    | 18-064<br>12-10-79               | 3982021<br>09-21-93<br>3935267<br>01-27-93 |                                  |
| NADOLOL<br>160MG                            | CORGARD<br>(TABLET; ORAL)                  | ER SQUIBB AND SONS    | 18-064<br>12-10-79               | 3982021<br>09-21-93<br>3935267<br>01-27-93 |                                  |
| NALBUPHINE HYDROCHLORIDE<br>10MG/ML         | NUBAIN<br>(INJECTABLE; INJECTION)          | DUPONT PHARMS/DUPONT  | 18-024<br>05-15-79               | 3393197<br>07-16-85                        | NS<br>09-24-86                   |
| NALBUPHINE HYDROCHLORIDE<br>20MG/ML         | NUBAIN<br>(INJECTABLE; INJECTION)          | DUPONT PHARMS/DUPONT  | 18-024<br>05-27-82               |  |                                  |
| NALIDIXIC ACID<br>250MG                     | NEGRAM<br>(TABLET; ORAL)                   | WINTHROP LABS/STERL   | 14-214<br>12-27-67               | 3590036<br>06-29-88                        |                                  |
| NALIDIXIC ACID<br>500MG                     | NEGRAM<br>(TABLET; ORAL)                   | WINTHROP LABS/STERL   | 14-214<br>03-06-64               | 3590036<br>06-29-88                        |                                  |
| NALIDIXIC ACID<br>1GM                       | NEGRAM<br>(TABLET; ORAL)                   | WINTHROP LABS/STERL   | 14-214<br>03-06-64               | 3590036<br>06-29-88                        |                                  |

**TABLE IV. NDA'S APPROVED FROM 1-1-82 TO 2-28-85 AND NDA'S WITH APPROPRIATE PATENT AND EXCLUSIVITY INFORMATION**

| <u>ACTIVE INGREDIENT(S)<br/>STRENGTH(S)</u>                                 | <u>TRADE NAME<br/>(DOSAGE FORM; ROUTE)</u> | <u>APPLICANT NAME</u> | <u>NDA NO.<br/>APPROVAL DATE</u> | <u>PATENT NO.<br/>EXP. DATE</u>  | <u>EXCLUSIVITY<br/>EXP. DATE</u> |
|---|--|-----------------------|----------------------------------|--|----------------------------------|
| NALIDIXIC ACID<br>250MG/5ML   | NEGRAM<br>(SUSPENSION; ORAL)               | WINTHROP LABS/STERL   | 17-430<br>04-17-73               | 3590036<br>06-29-88  |                                  |
| NALOXONE HYDROCHLORIDE<br>1MG/ML  | NARCAN<br>(INJECTABLE; INJECTION)          | DUPONT PHARMS/DUPONT  | 16-636<br>06-14-82               |  | NS<br>09-24-86                   |
| NALOXONE HYDROCHLORIDE; PENTAZOCINE<br>HYDROCHLORIDE<br>0.5MG; EQ 50MG BASE | TALWIN NX<br>(TABLET; ORAL)                | WINTHROP LABS/STERL   | 18-733<br>12-16-82               | 4105659<br>08-08-95  | NC<br>09-24-86                   |
| NALTREXONE HYDROCHLORIDE<br>50MG  | TREXAN<br>(TABLET; ORAL)                   | DUPONT PHARMS/DUPONT  | 18-932<br>11-20-84               |  | NCE<br>11-20-89                  |
| NAPROXEN<br>125MG   | NAPROSYN<br>(TABLET; ORAL)                 | SYNTEX PR             | 17-581<br>03-11-76               | 3998966<br>12-21-93<br>4009197<br>09-09-92<br>4001301<br>09-09-92<br>3904682<br>09-09-92 | NS<br>09-24-86                   |
| NAPROXEN<br>250MG   | NAPROSYN<br>(TABLET; ORAL)                 | SYNTEX PR             | 17-581<br>03-11-76               |  |                                  |
| NAPROXEN<br>375MG   | NAPROSYN<br>(TABLET; ORAL)                 | SYNTEX PR             | 17-581<br>07-18-80               |  |                                  |

**TABLE IV. NDA'S APPROVED FROM 1-1-82 TO 2-28-85 AND NDA'S WITH APPROPRIATE PATENT AND EXCLUSIVITY INFORMATION**

| <u>ACTIVE INGREDIENT(S)<br/>STRENGTH(S)</u> | <u>TRADE NAME<br/>(DOSAGE FORM; ROUTE)</u> | <u>APPLICANT NAME</u> | <u>NDA NO.<br/>APPROVAL DATE</u> | <u>PATENT NO.<br/>EXP. DATE</u>  | <u>EXCLUSIVITY<br/>EXP. DATE</u> |
|---|--|-----------------------|----------------------------------|--|----------------------------------|
| NAPROXEN<br>500MG                           | NAPROSYN<br>(TABLET; ORAL)                 | SYNTEX PR             | 17-581<br>04-15-82               | 3998966<br>12-21-93<br>4009197<br>09-09-92<br>4001301<br>09-09-92<br>3904682<br>09-09-92 | NS<br>09-24-86                   |
| NAPROXEN SODIUM<br>275MG                    | ANAPROX<br>(TABLET; ORAL)                  | SYNTEX PR             | 18-164<br>09-04-80               | 3998966<br>12-21-93<br>4001301<br>09-09-92<br>4009197<br>09-09-92                        |                                  |
| NICLOSAMIDE<br>500MG                        | NICLOCIDE<br>(TABLET, CHEWABLE; ORAL)      | MILES PHARMS/MILES    | 18-669<br>05-14-82               |  | NCE<br>05-14-92                  |
| NICOTINE RESIN COMPLEX<br>EQ 2MG BASE       | NICORETTE<br>(GUM, CHEWING; ORAL)          | MERRELL DOM/DOM CHEM  | 18-612<br>01-13-84               |  | NCE<br>01-13-94                  |
| NIFEDIPINE<br>10MG                          | PROCARDIA<br>(CAPSULE; ORAL)               | PFIZER LABS/PFIZER    | 18-482<br>12-31-81               | 3644627<br>02-22-89  |                                  |
| NITROGLYCERIN<br>0.5MG/ML                   | TRIDIL<br>(INJECTABLE; INJECTION)          | AM CRITICAL CARE/AHS  | 18-537<br>06-16-83               |  | NDF<br>09-24-86                  |
| NITROGLYCERIN<br>5MG/ML                     | NITROSTAT<br>(INJECTABLE; INJECTION)       | PARKE-DAVIS/M-L       | 18-588<br>12-23-83               |  | NDF<br>09-24-86                  |
| NITROGLYCERIN<br>5MG/ML                     | NITRO-BID<br>(INJECTABLE; INJECTION)       | MARION LABORATORIES   | 18-621<br>01-05-82               |  | NDF<br>09-24-86                  |
| NITROGLYCERIN<br>1MG/ML                     | NITRONAL<br>(INJECTABLE; INJECTION)        | G POHL-BOSKAMP        | 18-672<br>08-30-83               |  | NDF<br>09-24-86                  |
| NITROGLYCERIN<br>5MG/ML                     | NITRONAL<br>(INJECTABLE; INJECTION)        | G POHL-BOSKAMP        | 18-672<br>08-30-83               |  | NDF<br>09-24-86                  |

TABLE IV. NDA'S APPROVED FROM 1-1-82 TO 2-28-85 AND NDA'S WITH APPROPRIATE PATENT AND EXCLUSIVITY INFORMATION

| <u>ACTIVE INGREDIENT(S)<br/>STRENGTH(S)</u>     | <u>TRADE NAME<br/>(DOSAGE FORM; ROUTE)</u> | <u>APPLICANT NAME</u> | <u>NDA NO.<br/>APPROVAL DATE</u> | <u>PATENT NO.<br/>EXP. DATE</u>                                   | <u>EXCLUSIVITY<br/>EXP. DATE</u> |
|---|--|-----------------------|----------------------------------|---|----------------------------------|
| NITROGLYCERIN<br>0.8MG/ML                       | NITROL<br>(INJECTABLE; INJECTION)          | KREMERS-URBAN         | 18-774<br>01-19-83               |   | NDF<br>09-24-86                  |
| NOMIFENSINE MALEATE<br>25MG                     | MERITAL<br>(CAPSULE; ORAL)                 | HOECHST-ROUSSEL       | 18-224<br>12-31-84               |   | NCE<br>12-31-89                  |
| NOMIFENSINE MALEATE<br>50MG                     | MERITAL<br>(CAPSULE; ORAL)                 | HOECHST-ROUSSEL       | 18-224<br>12-31-84               |   | NCE<br>12-31-89                  |
| NORETHINDRONE ACETATE<br>5MG                    | AYGESTIN<br>(TABLET; ORAL)                 | AYERST LABS/AMHO      | 18-405<br>04-21-82               |   |                                  |
| NORGESTREL<br>0.075MG                           | OVRETTE<br>(TABLET; ORAL)                  | WYETH LABS/AMHO       | 17-031<br>10-23-73               | 3666858<br>05-30-89<br>3850911<br>11-26-91<br>3959322<br>11-26-91 |                                  |
| NORTRIPTYLINE HYDROCHLORIDE<br>EQ 10MG BASE     | AVENTYL HCL<br>(CAPSULE; ORAL)             | ELI LILLY             | 14-684<br>11-06-64               | 3922305<br>11-25-92   |                                  |
| NORTRIPTYLINE HYDROCHLORIDE<br>EQ 25MG BASE     | AVENTYL HCL<br>(CAPSULE; ORAL)             | ELI LILLY             | 14-684<br>11-06-64               | 3922305<br>11-25-92   |                                  |
| NORTRIPTYLINE HYDROCHLORIDE<br>EQ 10MG BASE/5ML | AVENTYL HCL<br>(SOLUTION; ORAL)            | ELI LILLY             | 14-685<br>11-06-64               | 3922305<br>11-25-92   |                                  |
| NORTRIPTYLINE HYDROCHLORIDE<br>EQ 10MG BASE/5ML | PAMELOR<br>(SOLUTION; ORAL)                | SANDOZ PHARMS/SANDOZ  | 18-012<br>08-01-77               | 3922305<br>11-25-92   |                                  |
| NORTRIPTYLINE HYDROCHLORIDE<br>EQ 10MG BASE     | PAMELOR<br>(CAPSULE; ORAL)                 | SANDOZ PHARMS/SANDOZ  | 18-013<br>08-01-77               | 3922305<br>11-25-92   |                                  |
| NORTRIPTYLINE HYDROCHLORIDE<br>EQ 25MG BASE     | PAMELOR<br>(CAPSULE; ORAL)                 | SANDOZ PHARMS/SANDOZ  | 18-013<br>08-01-77               | 3922305<br>11-25-92   |                                  |
| NORTRIPTYLINE HYDROCHLORIDE<br>EQ 75MG BASE     | PAMELOR<br>(CAPSULE; ORAL)                 | SANDOZ PHARMS/SANDOZ  | 18-013<br>06-14-79               | 3922305<br>11-25-92   |                                  |

**TABLE IV. NDA'S APPROVED FROM 1-1-82 TO 2-28-85 AND NDA'S WITH APPROPRIATE PATENT AND EXCLUSIVITY INFORMATION**

| <u>ACTIVE INGREDIENT(S)<br/>STRENGTH(S)</u> | <u>TRADE NAME<br/>(DOSAGE FORM; ROUTE)</u> | <u>APPLICANT NAME</u> | <u>NDA NO.<br/>APPROVAL DATE</u> | <u>PATENT NO.<br/>EXP. DATE</u>                                   | <u>EXCLUSIVITY<br/>EXP. DATE</u> |
|---|--|-----------------------|----------------------------------|---|----------------------------------|
| NORTRIPTYLINE HYDROCHLORIDE<br>EQ 50MG BASE | PAMELOR<br>(CAPSULE; ORAL)                 | SANDOZ PHARMS/SANDOZ  | 18-013<br>06-14-79               | 3922305<br>11-25-92   |                                  |
| OXAMNIQUINE<br>250MG                        | VANSTIL<br>(CAPSULE; ORAL)                 | PFIZER LABS/PFIZER    | 18-069<br>07-23-80               | 3903283<br>09-02-92<br>3821228<br>06-28-91<br>3925391<br>12-09-92 |                                  |
| OXPRENOLOL HYDROCHLORIDE<br>20MG            | TRASICOR<br>(CAPSULE; ORAL)                | CIBA/CIBA-GEIGY       | 18-166<br>12-28-83               | 3483221<br>12-09-86   | NCE<br>12-28-93                  |
| OXPRENOLOL HYDROCHLORIDE<br>40MG            | TRASICOR<br>(CAPSULE; ORAL)                | CIBA/CIBA-GEIGY       | 18-166<br>12-28-83               | 3483221<br>12-09-86   | NCE<br>12-28-93                  |
| OXPRENOLOL HYDROCHLORIDE<br>80MG            | TRASICOR<br>(CAPSULE; ORAL)                | CIBA/CIBA-GEIGY       | 18-166<br>12-28-83               | 3483221<br>12-09-86   | NCE<br>12-28-93                  |
| OXPRENOLOL HYDROCHLORIDE<br>160MG           | TRASICOR<br>(CAPSULE; ORAL)                | CIBA/CIBA-GEIGY       | 18-166<br>12-28-83               | 3483221<br>12-09-86   | NCE<br>12-28-93                  |
| PANCURONIUM BROMIDE<br>2MG/ML               | PAVULON<br>(INJECTABLE; INJECTION)         | ORGANON/AKZONA        | 17-015<br>10-24-72               | 3553212<br>01-05-88   |                                  |
| PANCURONIUM BROMIDE<br>1MG/ML               | PAVULON<br>(INJECTABLE; INJECTION)         | ORGANON/AKZONA        | 17-015<br>09-14-73               | 3553212<br>01-05-88   |                                  |
| PARAMETHASONE ACETATE<br>1MG                | HALDRONE<br>(TABLET; ORAL)                 | ELI LILLY             | 12-772<br>04-17-61               | 3499016<br>03-03-87   |                                  |
| PARAMETHASONE ACETATE<br>2MG                | HALDRONE<br>(TABLET; ORAL)                 | ELI LILLY             | 12-772<br>04-17-61               | 3499016<br>03-03-87   |                                  |
| PENTAGASTRIN<br>0.25MG/ML                   | PEPTAVLON<br>(INJECTABLE; INJECTION)       | AYERST LABS/AMHO      | 17-048<br>07-26-74               | 3896103<br>07-22-92   |                                  |
| PENTAMIDINE ISETHIONATE<br>300MG/VIAL       | PENTAM 300<br>(INJECTABLE; INJECTION)      | LYPHOMED              | 19-264<br>10-16-84               |   |                                  |

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| <u>ACTIVE INGREDIENT(S)<br/>STRENGTH(S)</u>                                       | <u>TRADE NAME<br/>(DOSAGE FORM; ROUTE)</u>                     | <u>APPLICANT NAME</u> | <u>NDA NO.<br/>APPROVAL DATE</u> | <u>PATENT NO.<br/>EXP. DATE</u> | <u>EXCLUSIVITY<br/>EXP. DATE</u> |
|---|--|-----------------------|----------------------------------|---------------------------------|----------------------------------|
| PENTAZOCINE LACTATE<br>EQ 30MG BASE/ML  | TALWIN<br>(INJECTABLE; INJECTION)                              | WINTHROP LABS/STERL   | 16-194<br>07-24-67               | 4105659<br>08-08-95             |                                  |
| PENTETATE INDIUM DISODIUM, IN-111<br>TICI/ML                                      | MPI INDIUM DTPA IN 111<br>(INJECTABLE; INJECTION)              | MEDI-PHYSICS          | 17-707<br>02-18-82               |                                 | NCE<br>02-18-92                  |
| PENTOXIFYLLINE<br>400MG   | TRENTAL<br>(TABLET, CONTROLLED<br>RELEASE; ORAL)               | HOECHST-ROUSSEL       | 18-631<br>08-30-84               | 3737433<br>06-05-90             | NCE<br>08-30-94                  |
| PHENYLEPHRINE HYDROCHLORIDE; PROMETHAZINE<br>HYDROCHLORIDE<br>5MG/5ML; 6.25MG/5ML | PHENERGAN VC<br>(SYRUP; ORAL)                                  | WYETH LABS/AMHO       | 08-604<br>04-02-84               |                                 |                                  |
| PILOCARPINE<br>5MG  | OCUSERT PILO-20<br>(INSERT, CONTROLLED<br>RELEASE; OPHTHALMIC) | ALZA                  | 17-431<br>07-29-74               | 391628<br>06-08-93              |                                  |
| PILOCARPINE<br>11MG   | OCUSERT PILO-40<br>(INSERT, CONTROLLED<br>RELEASE; OPHTHALMIC) | ALZA                  | 17-548<br>07-29-72               | 391628<br>06-08-93              |                                  |
| PILOCARPINE HYDROCHLORIDE<br>4%   | PILOPINE HS<br>(GEL; OPHTHALMIC)                               | ALCON LABORATORIES    | 18-796<br>10-01-84               |                                 | NDF<br>10-01-87                  |
| PIMOZIDE<br>2MG   | ORAP<br>(TABLET; ORAL)   | MCNEIL PHARM          | 17-473<br>07-31-84               |                                 | NCE<br>07-31-94                  |
| PINDOLOL<br>5MG   | VISKEN<br>(TABLET; ORAL)                                       | SANDOZ PHARMS/SANDOZ  | 18-285<br>09-03-82               | 3471515<br>10-07-86             | NCE<br>09-03-92                  |
| PINDOLOL<br>10MG  | VISKEN<br>(TABLET; ORAL)                                       | SANDOZ PHARMS/SANDOZ  | 18-285<br>09-03-82               | 3471515<br>10-07-86             | NCE<br>09-03-92                  |
| PINDOLOL<br>15MG  | VISKEN<br>(TABLET; ORAL)                                       | SANDOZ PHARMS/SANDOZ  | 18-285<br>09-03-82               | 3471515<br>10-07-86             | NCE<br>09-03-92                  |

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| <u>ACTIVE INGREDIENT(S)<br/>STRENGTH(S)</u>   | <u>TRADE NAME<br/>(DOSAGE FORM; ROUTE)</u>       | <u>APPLICANT NAME</u> | <u>NDA NO.<br/>APPROVAL DATE</u> | <u>PATENT NO.<br/>EXP. DATE</u>  | <u>EXCLUSIVITY<br/>EXP. DATE</u> |
|---|--|-----------------------|----------------------------------|--|----------------------------------|
| PIROXICAM<br>10MG   | FELDENE<br>(CAPSULE; ORAL)                       | PFIZER LABS/PFIZER    | 18-147<br>04-06-82               | 3591584<br>07-06-88<br>3674876<br>07-04-89<br>3862319<br>01-21-92<br>4100347<br>07-11-95<br>3927002<br>12-16-92<br>RE29668<br>12-10-91 | NCE<br>04-06-92                  |
| PIROXICAM<br>20MG   | FELDENE<br>(CAPSULE; ORAL)                       | PFIZER LABS/PFIZER    | 18-147<br>04-06-82               | 3591584<br>07-06-88<br>3674876<br>07-04-89<br>3862319<br>01-21-92<br>4100347<br>07-11-95<br>3927002<br>12-16-92<br>RE29668<br>12-10-91 | NCE<br>04-06-92                  |
| POLYETHYLENE GLYCOL 3350;<br>POTASSIUM CHLORIDE;<br>SODIUM BICARBONATE;<br>SODIUM CHLORIDE;<br>SODIUM SULFATE<br>236GM/BOT;<br>2.97GM/BOT;<br>6.74GM/BOT;<br>5.86GM/BOT;<br>22.74GM/BOT | GOLYTELY<br>(POWDER FOR<br>RECONSTITUTION; ORAL) | BRAINTREE LABS        | 19-011<br>07-13-84               |  |                                  |

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| <u>ACTIVE INGREDIENT(S)<br/>STRENGTH(S)</u>   | <u>TRADE NAME<br/>(DOSAGE FORM; ROUTE)</u>     | <u>APPLICANT NAME</u> | <u>NDA NO.<br/>APPROVAL DATE</u> | <u>PATENT NO.<br/>EXP. DATE</u>                                   | <u>EXCLUSIVITY<br/>EXP. DATE</u> |
|---|--|-----------------------|----------------------------------|---|----------------------------------|
| POLYETHYLENE GLYCOL 3350;<br>POTASSIUM CHLORIDE;<br>SODIUM BICARBONATE;<br>SODIUM CHLORIDE;<br>SODIUM SULFATE<br>120GM/PACKET;<br>1.49GM/PACKET;<br>3.36GM/PACKET;<br>2.92GM/PACKET;<br>11.36GM/PACKET  | COLYTE<br>(POWDER FOR<br>RECONSTITUTION; ORAL) | EDLAW PREPARATIONS    | 18-983<br>10-26-84               |   |                                  |
| POLYETHYLENE GLYCOL 3350;<br>POTASSIUM CHLORIDE;<br>SODIUM BICARBONATE;<br>SODIUM CHLORIDE;<br>SODIUM SULFATE<br>227.1GM/PACKET;<br>2.82GM/PACKET;<br>6.36GM/PACKET;<br>5.53GM/PACKET;<br>21.5GM/PACKET | COLYTE<br>(POWDER FOR<br>RECONSTITUTION; ORAL) | EDLAW PREPARATIONS    | 18-983<br>10-26-84               |   |                                  |
| POLYETHYLENE GLYCOL 3350;<br>POTASSIUM CHLORIDE;<br>SODIUM BICARBONATE;<br>SODIUM CHLORIDE;<br>SODIUM SULFATE<br>360GM/PACKET;<br>4.47GM/PACKET;<br>10.08GM/PACKET;<br>8.76GM/PACKET;<br>34.08GM/PACKET | COLYTE<br>(POWDER FOR<br>RECONSTITUTION; ORAL) | EDLAW PREPARATIONS    | 18-983<br>10-26-84               |   |                                  |
| POLYTHIAZIDE; PRAZOSIN HYDROCHLORIDE<br>0.5MG; 1MG  | MINIZIDE<br>(CAPSULE; ORAL)                    | PFIZER LABS/PFIZER    | 17-986<br>06-13-80               | 3511836<br>05-12-87<br>3663706<br>05-16-89<br>4130647<br>12-19-95 |                                  |

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| <u>ACTIVE INGREDIENT(S)<br/>STRENGTH(S)</u>                     | <u>TRADE NAME<br/>(DOSAGE FORM; ROUTE)</u>  | <u>APPLICANT NAME</u> | <u>NDA NO.<br/>APPROVAL DATE</u> | <u>PATENT NO.<br/>EXP. DATE</u>                                   | <u>EXCLUSIVITY<br/>EXP. DATE</u> |
|---|---|-----------------------|----------------------------------|---|----------------------------------|
| POLYTHIAZIDE; PRAZOSIN HYDROCHLORIDE<br>0.5MG; 2MG              | MINIZIDE<br>(CAPSULE; ORAL)   | PFIZER LABS/PFIZER    | 17-986<br>06-13-80               | 3511836<br>05-12-87<br>3663706<br>05-16-89<br>4130647<br>12-19-95 |                                  |
| POLYTHIAZIDE; PRAZOSIN HYDROCHLORIDE<br>0.5MG; 5MG              | MINIZIDE<br>(CAPSULE; ORAL)   | PFIZER LABS/PFIZER    | 17-986<br>06-13-80               | 3511836<br>05-12-87<br>3663706<br>05-16-89<br>4130647<br>12-19-95 |                                  |
| POTASSIUM ACETATE<br>2MEQ/ML                                    | POTASSIUM ACETATE IN<br>PLASTIC CONTAINER<br>(INJECTABLE; INJECTION)                                    | ABBOTT LABORATORIES   | 18-896<br>07-20-84               |   | NDF<br>09-24-86                  |
| POTASSIUM CHLORIDE<br>10MEQ                                     | KLOTRIX<br>(TABLET, CONTROLLED<br>RELEASE; ORAL)  | MEAD JOHNSON/B-M      | 17-850<br>05-22-80               | 4140756<br>02-20-96   |                                  |
| POTASSIUM CHLORIDE; SODIUM CHLORIDE<br>150MG/100ML; 900MG/100ML | SODIUM CHLORIDE 0.9% AND<br>POTASSIUM CHLORIDE 10MEQ<br>IN PLASTIC CONTAINER<br>(INJECTABLE; INJECTION) | TRAVENOL LABS         | 18-630<br>02-17-83               |   |                                  |
| POTASSIUM CHLORIDE; SODIUM CHLORIDE<br>300MG/100ML; 900MG/100ML | SODIUM CHLORIDE 0.9% AND<br>POTASSIUM CHLORIDE 20MEQ<br>IN PLASTIC CONTAINER<br>(INJECTABLE; INJECTION) | TRAVENOL LABS         | 18-630<br>02-17-83               |   |                                  |
| POTASSIUM CHLORIDE; SODIUM CHLORIDE<br>150MG/100ML; 900MG/100ML | SODIUM CHLORIDE 0.9% AND<br>POTASSIUM CHLORIDE 20MEQ<br>IN PLASTIC CONTAINER<br>(INJECTABLE; INJECTION) | TRAVENOL LABS         | 18-630<br>02-17-83               |   |                                  |
| POTASSIUM CHLORIDE; SODIUM CHLORIDE<br>300MG/100ML; 900MG/100ML | SODIUM CHLORIDE 0.9% AND<br>POTASSIUM CHLORIDE 40MEQ<br>IN PLASTIC CONTAINER<br>(INJECTABLE; INJECTION) | TRAVENOL LABS         | 18-630<br>02-17-83               |   |                                  |

TABLE IV. NDA'S APPROVED FROM 1-1-82 TO 2-28-85 AND NDA'S WITH APPROPRIATE PATENT AND EXCLUSIVITY INFORMATION

| <u>ACTIVE INGREDIENT(S)<br/>STRENGTH(S)</u>                     | <u>TRADE NAME<br/>(DOSAGE FORM; ROUTE)</u>   | <u>APPLICANT NAME</u> | <u>NDA NO.<br/>APPROVAL DATE</u> | <u>PATENT NO.<br/>EXP. DATE</u>  | <u>EXCLUSIVITY<br/>EXP. DATE</u> |
|---|--|-----------------------|----------------------------------|--|----------------------------------|
| POTASSIUM CHLORIDE; SODIUM CHLORIDE<br>75MG/100ML; 900MG/100ML  | SODIUM CHLORIDE 0.9% AND<br>POTASSIUM CHLORIDE 0.075%<br>IN PLASTIC CONTAINER<br>(INJECTABLE; INJECTION) | AM MCGAW/AM HOSP      | 18-722<br>11-09-82               |  |                                  |
| POTASSIUM CHLORIDE; SODIUM CHLORIDE<br>150MG/100ML; 900MG/100ML | SODIUM CHLORIDE 0.9% AND<br>POTASSIUM CHLORIDE 0.15%<br>IN PLASTIC CONTAINER<br>(INJECTABLE; INJECTION)  | AM MCGAW/AM HOSP      | 18-722<br>11-09-82               |  |                                  |
| POTASSIUM CHLORIDE; SODIUM CHLORIDE<br>220MG/100ML; 900MG/100ML | SODIUM CHLORIDE 0.9% AND<br>POTASSIUM CHLORIDE 0.22%<br>IN PLASTIC CONTAINER<br>(INJECTABLE; INJECTION)  | AM MCGAW/AM HOSP      | 18-722<br>11-09-82               |  |                                  |
| POTASSIUM CHLORIDE; SODIUM CHLORIDE<br>300MG/100ML; 900MG/100ML | SODIUM CHLORIDE 0.9% AND<br>POTASSIUM CHLORIDE 0.3%<br>IN PLASTIC CONTAINER<br>(INJECTABLE; INJECTION)   | AM MCGAW/AM HOSP      | 18-722<br>11-09-82               |  |                                  |
| PRALIDOXIME CHLORIDE<br>300MG/ML                                | PROTOPAM CHLORIDE<br>(INJECTABLE; INJECTION)   | AYERST LABS/AMHO      | 18-799<br>12-13-82               |  | NDF<br>09-24-86                  |
| PRALIDOXIME CHLORIDE<br>300MG/ML                                | PRALIDOXIME CHLORIDE<br>(INJECTABLE; INJECTION)  | SURVIVAL TECHNOLOGY   | 18-986<br>12-13-82               |  | NDF<br>09-24-86                  |
| PRAZEPAM<br>20MG  | CENTRAX<br>(CAPSULE; ORAL)   | PARKE-DAVIS/M-L       | 18-144<br>05-10-82               |  | NS<br>09-24-86                   |
| PRAZIQUNTEL<br>600MG  | BILTRICIDE<br>(TABLET; ORAL)   | MILES PHARMS/MILES    | 18-714<br>12-29-82               | 4001411<br>01-04-94  | NCE<br>12-29-92                  |
| PRAZOSIN HYDROCHLORIDE<br>5MG                                   | MINIPRESS<br>(CAPSULE; ORAL)   | PFIZER LABS/PFIZER    | 17-442<br>06-23-76               | 3511836<br>05-12-87<br>3663706<br>05-16-89<br>4092315<br>05-30-95<br>4130647<br>12-19-95 |                                  |

**TABLE IV. NDA'S APPROVED FROM 1-1-82 TO 2-28-85 AND NDA'S WITH APPROPRIATE PATENT AND EXCLUSIVITY INFORMATION**

| <u>ACTIVE INGREDIENT(S)<br/>STRENGTH(S)</u> | <u>TRADE NAME<br/>(DOSAGE FORM; ROUTE)</u> | <u>APPLICANT NAME</u> | <u>NDA NO.<br/>APPROVAL DATE</u> | <u>PATENT NO.<br/>EXP. DATE</u>  | <u>EXCLUSIVITY<br/>EXP. DATE</u> |
|---|--|-----------------------|----------------------------------|--|----------------------------------|
| PRAZOSIN HYDROCHLORIDE<br>1MG               | MINIPRESS<br>(CAPSULE; ORAL)               | PFIZER LABS/PFIZER    | 17-442<br>06-23-76               | 3511836<br>05-12-87<br>3663706<br>05-16-89<br>4092315<br>05-30-95<br>4130647<br>12-19-95 |                                  |
| PRAZOSIN HYDROCHLORIDE<br>2MG               | MINIPRESS<br>(CAPSULE; ORAL)               | PFIZER LABS/PFIZER    | 17-442<br>06-23-76               | 3511836<br>05-12-87<br>3663706<br>05-16-89<br>4092315<br>05-30-95<br>4130647<br>12-19-95 |                                  |
| PROBUCOL<br>250MG                           | LORELCO<br>(TABLET; ORAL)                  | MERRELL DOW/DOW CHEM  | 17-535<br>02-01-77               | 3576883<br>04-27-88<br>3862332<br>01-21-92   |                                  |
| PROCARBAZINE HYDROCHLORIDE<br>EQ 50MG BASE  | MATULANE<br>(CAPSULE; ORAL)                | HOFFMANN-LA ROCHE     | 16-785<br>07-22-69               | 3520926<br>07-21-87  |                                  |
| PROPRANOLOL HYDROCHLORIDE<br>10MG           | INDERAL<br>(TABLET; ORAL)                  | AYERST LABS/AMHO      | 16-418<br>11-13-67               |  | I-15<br>09-24-86                 |
| PROPRANOLOL HYDROCHLORIDE<br>20MG           | INDERAL<br>(TABLET; ORAL)                  | AYERST LABS/AMHO      | 16-418<br>10-16-74               |  | I-15<br>09-24-86                 |
| PROPRANOLOL HYDROCHLORIDE<br>40MG           | INDERAL<br>(TABLET; ORAL)                  | AYERST LABS/AMHO      | 16-418<br>11-13-67               |  | I-15<br>09-24-86                 |
| PROPRANOLOL HYDROCHLORIDE<br>60MG           | INDERAL<br>(TABLET; ORAL)                  | AYERST LABS/AMHO      | 16-418<br>10-18-82               |  | NS<br>09-24-86                   |
| PROPRANOLOL HYDROCHLORIDE<br>80MG           | INDERAL<br>(TABLET; ORAL)                  | AYERST LABS/AMHO      | 16-418<br>10-16-74               |  | I-15<br>09-24-86                 |

**TABLE IV. NDA'S APPROVED FROM 1-1-82 TO 2-28-85 AND NDA'S WITH APPROPRIATE PATENT AND EXCLUSIVITY INFORMATION**

| <u>ACTIVE INGREDIENT(S)<br/>STRENGTH(S)</u> | <u>TRADE NAME<br/>(DOSAGE FORM; ROUTE)</u>           | <u>APPLICANT NAME</u> | <u>NDA NO.<br/>APPROVAL DATE</u> | <u>PATENT NO.<br/>EXP. DATE</u>            | <u>EXCLUSIVITY<br/>EXP. DATE</u> |
|---|--|-----------------------|----------------------------------|--|----------------------------------|
| PROPRANOLOL HYDROCHLORIDE<br>80MG           | INDERAL LA<br>(CAPSULE, CONTROLLED<br>RELEASE; ORAL) | AYERST LABS/AMHO      | 18-553<br>04-19-83               |  | NDF<br>09-24-86                  |
| PROPRANOLOL HYDROCHLORIDE<br>90MG           | INDERAL<br>(TABLET; ORAL)                            | AYERST LABS/AMHO      | 16-418<br>10-18-82               |  | NS<br>09-24-86                   |
| PROPRANOLOL HYDROCHLORIDE<br>120MG          | INDERAL LA<br>(CAPSULE, CONTROLLED<br>RELEASE; ORAL) | AYERST LABS/AMHO      | 18-553<br>04-19-83               |  | NDF<br>09-24-86                  |
| PROPRANOLOL HYDROCHLORIDE<br>160MG          | INDERAL LA<br>(CAPSULE, CONTROLLED<br>RELEASE; ORAL) | AYERST LABS/AMHO      | 18-553<br>04-19-83               |  | NDF<br>09-24-86                  |
| PROTAMINE SULFATE<br>250MG/VIAL             | PROTAMINE SULFATE<br>(INJECTABLE; INJECTION)         | UPJOHN                | 07-413<br>08-02-84               |  | NS<br>09-24-86                   |
| PROTIRELIN<br>0.5MG/ML                      | THYPINONE<br>(INJECTABLE; INJECTION)                 | ABBOTT LABORATORIES   | 17-638<br>11-05-76               | 3746697<br>07-17-90                        |                                  |
| PROTIRELIN<br>0.5MG/ML                      | RELEFACT TRH<br>(INJECTABLE; INJECTION)              | HOECHST-ROUSSEL       | 18-087<br>07-18-78               | 3746697<br>07-17-90                        |                                  |
| PROTRIPTYLINE HYDROCHLORIDE<br>5MG          | VIVACTIL<br>(TABLET; ORAL)                           | MS&D/MERCK            | 16-012<br>09-27-67               | 3372196<br>03-05-85                        |                                  |
| PROTRIPTYLINE HYDROCHLORIDE<br>10MG         | VIVACTIL<br>(TABLET; ORAL)                           | MS&D/MERCK            | 16-012<br>09-27-67               | 3372196<br>03-05-85                        |                                  |
| PYRANTEL PAMOATE<br>EQ 250MG BASE/5ML       | ANTIMINTH<br>(SUSPENSION; ORAL)                      | ROERIG/PFIZER         | 16-883<br>12-30-71               | 3644624<br>02-22-89<br>3549624<br>12-22-87 |                                  |
| RANITIDINE HYDROCHLORIDE<br>EQ 150MG BASE   | ZANTAC<br>(TABLET; ORAL)                             | GLAXO                 | 18-703<br>06-09-83               | 4128658<br>12-05-95                        | NCE<br>06-09-93                  |
| RANITIDINE HYDROCHLORIDE<br>EQ 25MG BASE/ML | ZANTAC<br>(INJECTABLE; INJECTION)                    | GLAXO                 | 19-090<br>10-19-84               | 4128658<br>12-05-95                        | NCE<br>06-09-93                  |

**TABLE IV. NDA'S APPROVED FROM 1-1-82 TO 2-28-85 AND NDA'S WITH APPROPRIATE PATENT AND EXCLUSIVITY INFORMATION**

| <u>ACTIVE INGREDIENT(S)</u><br><u>STRENGTH(S)</u> | <u>TRADE NAME</u><br><u>(DOSAGE FORM; ROUTE)</u>                  | <u>APPLICANT NAME</u> | <u>NDA NO.</u><br><u>APPROVAL DATE</u> | <u>PATENT NO.</u><br><u>EXP. DATE</u>      | <u>EXCLUSIVITY</u><br><u>EXP. DATE</u> |
|---|---|-----------------------|--|--|--|
| RITODRINE HYDROCHLORIDE<br>10MG                   | YUTOPAR<br>(TABLET; ORAL)   | ASTRA PHARM PRODS     | 18-555<br>12-12-80                     | 3410944<br>11-12-85                        |  |
| RITODRINE HYDROCHLORIDE<br>10MG/ML                | YUTOPAR<br>(INJECTABLE; INJECTION)                                | ASTRA PHARM PRODS     | 18-580<br>12-12-80                     | 3410944<br>11-12-85                        |  |
| RITODRINE HYDROCHLORIDE<br>15MG/ML                | YUTOPAR<br>(INJECTABLE; INJECTION)                                | ASTRA PHARM PRODS     | 18-580<br>09-27-84                     | 3410944<br>11-12-85                        |  |
| SAFFLOWER OIL; SOYBEAN OIL<br>10%; 10%            | LIPOSYN II 20%<br>(INJECTABLE; INJECTION)                         | ABBOTT LABORATORIES   | 18-991<br>08-27-84                     |  | NP<br>09-24-86                         |
| SAFFLOWER OIL; SOYBEAN OIL<br>5%; 5%              | LIPOSYN II 10%<br>(INJECTABLE; INJECTION)                         | ABBOTT LABORATORIES   | 18-997<br>08-27-84                     |  | NP<br>09-24-86                         |
| SARALASIN ACETATE<br>EQ 0.6MG BASE/ML             | SARENIN<br>(INJECTABLE; INJECTION)                                | NORMICH EATON/P&G     | 18-009<br>05-29-81                     | 3932624<br>01-13-93<br>3886134<br>05-27-92 |  |
| SCOPOLAMINE<br>1.5MG                              | TRANSERM-SCOP<br>(FILM, CONTROLLED<br>RELEASE; PERCUTANEOUS)      | CIBA/CIBA-GEIGY       | 17-874<br>12-31-79                     | 4031894<br>06-28-94<br>4262003<br>04-14-98 |  |
| SELENIUM SULFIDE<br>2.5%                          | SELSUN<br>(SHAMPOO/LOTION; TOPICAL)                               | ABBOTT LABS           | 07-936<br>05-17-51                     |  | I-3<br>09-24-86                        |
| SILVER SULFADIAZINE<br>1%                         | SILVADENE<br>(CREAM; TOPICAL)                                     | MARION LABORATORIES   | 17-381<br>11-26-73                     | 3761590<br>09-24-90                        |  |
| SILVER SULFADIAZINE<br>1%                         | SSD<br>(CREAM; TOPICAL)   | TRAVENOL LABS         | 18-578<br>02-25-82                     |  |  |
| SINCALIDE<br>0.005MG/VIAL                         | KINEVAC<br>(INJECTABLE; INJECTION)                                | ER SQUIBB AND SONS    | 17-697<br>07-21-76                     | 3839315<br>10-01-91                        |  |
| SODIUM ACETATE, ANHYDROUS<br>2MEQ/ML              | SODIUM ACETATE IN<br>PLASTIC CONTAINER<br>(INJECTABLE; INJECTION) | ABBOTT LABORATORIES   | 18-893<br>05-04-83                     |  | PP<br>09-24-86                         |

**TABLE IV. NDA'S APPROVED FROM 1-1-82 TO 2-28-85 AND NDA'S WITH APPROPRIATE PATENT AND EXCLUSIVITY INFORMATION**

| <u>ACTIVE INGREDIENT(S)<br/>STRENGTH(S)</u> | <u>TRADE NAME<br/>(DOSAGE FORM; ROUTE)</u>  | <u>APPLICANT NAME</u> | <u>NDA NO.<br/>APPROVAL DATE</u> | <u>PATENT NO.<br/>EXP. DATE</u> | <u>EXCLUSIVITY<br/>EXP. DATE</u> |
|---|---|-----------------------|----------------------------------|---------------------------------|----------------------------------|
| SODIUM CHLORIDE<br>450MG/100ML              | SODIUM CHLORIDE 0.45%<br>IN PLASTIC CONTAINER<br>(SOLUTION; IRRIGATION)                   | TRAVENOL LABS         | 18-497<br>02-19-82               |                                 |                                  |
| SODIUM CHLORIDE<br>9MG/ML                   | BACTERIOSTATIC SODIUM<br>CHLORIDE 0.9% IN PLASTIC<br>CONTAINER<br>(INJECTABLE; INJECTION) | ABBOTT LABORATORIES   | 18-800<br>10-29-82               |                                 |                                  |
| SODIUM CHLORIDE<br>9MG/ML                   | SODIUM CHLORIDE 0.9%<br>IN PLASTIC CONTAINER<br>(INJECTABLE; INJECTION)                   | ABBOTT LABORATORIES   | 18-803<br>10-29-82               |                                 |                                  |
| SODIUM CHLORIDE<br>2.5MEQ/ML                | SODIUM CHLORIDE IN<br>PLASTIC CONTAINER<br>(INJECTABLE; INJECTION)                        | ABBOTT LABORATORIES   | 18-897<br>07-20-84               |                                 |                                  |
| SODIUM CHLORIDE<br>3GM/100ML                | SODIUM CHLORIDE 3% IN<br>PLASTIC CONTAINER<br>(INJECTABLE; INJECTION)                     | TRAVENOL LABS         | 19-022<br>11-01-83               |                                 |                                  |
| SODIUM CHLORIDE<br>5GM/100ML                | SODIUM CHLORIDE 5% IN<br>PLASTIC CONTAINER<br>(INJECTABLE; INJECTION)                     | TRAVENOL LABS         | 19-022<br>11-01-83               |                                 |                                  |
| SODIUM CHLORIDE<br>9MG/ML                   | SODIUM CHLORIDE 0.9% IN<br>PLASTIC CONTAINER<br>(INJECTABLE; INJECTION)                   | ABBOTT LABORATORIES   | 19-217<br>07-13-84               |                                 |                                  |
| SODIUM CHLORIDE<br>9MG/ML                   | SODIUM CHLORIDE 0.9%<br>IN PLASTIC CONTAINER<br>(INJECTABLE; INJECTION)                   | ABBOTT LABORATORIES   | 19-218<br>07-13-84               |                                 |                                  |
| SODIUM IODIDE, I-123<br>100 UCI             | SODIUM IODIDE I 123<br>(CAPSULE; ORAL)  | BENEDICT NUCLR PHARM  | 18-671<br>05-27-82               |                                 |                                  |
| SODIUM IODIDE, I-123<br>200 UCI             | SODIUM IODIDE I 123<br>(CAPSULE; ORAL)  | BENEDICT NUCLR PHARM  | 18-671<br>05-27-82               |                                 |                                  |

**TABLE IV. NDA'S APPROVED FROM 1-1-82 TO 2-28-85 AND NDA'S WITH APPROPRIATE PATENT AND EXCLUSIVITY INFORMATION**

| <u>ACTIVE INGREDIENT(S)<br/>STRENGTH(S)</u>                                     | <u>TRADE NAME<br/>(DOSAGE FORM; ROUTE)</u>                           | <u>APPLICANT NAME</u> | <u>NDA NO.<br/>APPROVAL DATE</u> | <u>PATENT NO.<br/>EXP. DATE</u> | <u>EXCLUSIVITY<br/>EXP. DATE</u> |
|---|--|-----------------------|----------------------------------|---------------------------------|----------------------------------|
| SODIUM IODIDE, I-123<br>400 UCI   | SODIUM IODIDE I 123<br>(CAPSULE; ORAL)                               | BENEDICT NUCLR PHARM  | 18-671<br>05-27-82               |                                 |                                  |
| SODIUM LACTATE<br>5MEQ/ML   | SODIUM LACTATE IN<br>PLASTIC CONTAINER<br>(INJECTABLE; INJECTION)    | ABBOTT LABORATORIES   | 18-947<br>09-05-84               |                                 | NS<br>09-24-86                   |
| SODIUM NITROPRUSSIDE<br>50MG/VIAL   | SODIUM NITROPRUSSIDE<br>(INJECTABLE; INJECTION)                      | ELKINS-SINN/AHROBINS  | 18-581<br>07-28-82               |                                 |                                  |
| SODIUM PHOSPHATE, DIBASIC; SODIUM<br>PHOSPHATE, MONOBASIC<br>142MG/ML; 276MG/ML | SODIUM PHOSPHATES<br>IN PLASTIC CONTAINER<br>(INJECTABLE; INJECTION) | ABBOTT LABORATORIES   | 18-892<br>05-10-83               |                                 | NP<br>09-24-86                   |
| SOMATROPIN<br>2 IU/VIAL   | ASELLACRIN 2<br>(INJECTABLE; INJECTION)                              | SERONO LABS           | 17-726<br>07-21-83               |                                 | NS<br>09-24-86                   |
| SORBITOL<br>3GM/100ML   | SORBITOL 3% IN PLASTIC<br>CONTAINER<br>(SOLUTION; IRRIGATION)        | TRAVENOL LABS         | 18-512<br>05-27-82               |                                 |                                  |
| SOYBEAN OIL<br>10%  | SOYACAL 10%<br>(INJECTABLE; INJECTION)                               | ALPHA THERAPEUTIC     | 18-465<br>06-29-83               |                                 |                                  |
| SOYBEAN OIL<br>10%  | TRAVAMULSION 10%<br>(INJECTABLE; INJECTION)                          | TRAVENOL LABS         | 18-660<br>02-26-82               |                                 |                                  |
| SOYBEAN OIL<br>20%  | TRAVAMULSION 20%<br>(INJECTABLE; INJECTION)                          | TRAVENOL LABS         | 18-758<br>02-15-83               |                                 |                                  |
| SOYBEAN OIL<br>20%  | SOYACAL 20%<br>(INJECTABLE; INJECTION)                               | ALPHA THERAPEUTIC     | 18-786<br>06-29-83               |                                 |                                  |
| SOYBEAN OIL<br>10%  | LIPOSYN III 10%<br>(INJECTABLE; INJECTION)                           | ABBOTT LABORATORIES   | 18-969<br>09-24-84               |                                 |                                  |
| SOYBEAN OIL<br>20%  | LIPOSYN III 20%<br>(INJECTABLE; INJECTION)                           | ABBOTT LABORATORIES   | 18-970<br>09-25-84               |                                 |                                  |
| STANZOLOL<br>2MG  | WINSTROL<br>(TABLET; ORAL)   | WINTHROP LABS/STERL   | 12-885<br>11-30-61               | 3704295<br>11-28-89             | I-28<br>09-24-86                 |

TABLE IV. NDA'S APPROVED FROM 1-1-82 TO 2-28-85 AND NDA'S WITH APPROPRIATE PATENT AND EXCLUSIVITY INFORMATION

| <u>ACTIVE INGREDIENT(S)<br/>STRENGTH(S)</u>           | <u>TRADE NAME<br/>(DOSAGE FORM; ROUTE)</u>                             | <u>APPLICANT NAME</u> | <u>NDA NO.<br/>APPROVAL DATE</u> | <u>PATENT NO.<br/>EXP. DATE</u>            | <u>EXCLUSIVITY<br/>EXP. DATE</u> |
|---|--|-----------------------|----------------------------------|--|----------------------------------|
| STREPTOZOCIN<br>1GM/VIAL                              | ZANOSAR<br>(INJECTABLE; INJECTION)                                     | UPJOHN                | 17-961<br>05-07-82               |  | NCE<br>05-07-92                  |
| SUCRALFATE<br>1GM                                     | CARAFATE<br>(TABLET; ORAL)   | MARION LABORATORIES   | 18-333<br>10-30-81               | 3432489<br>03-11-86                        |                                  |
| SUFENTANIL CITRATE<br>EQ 0.05MG BASE/ML               | SUFENTA<br>(INJECTABLE; INJECTION)                                     | JANSSEN PHARMA        | 19-050<br>05-04-84               | 3998834<br>12-21-93                        | NCE<br>05-04-94                  |
| SULFAMETHOXAZOLE; TRIMETHOPRIM<br>400MG; 80MG         | BACTRIM<br>(TABLET; ORAL)  | HOFFMANN-LA ROCHE     | 17-377<br>07-30-73               | RE28636<br>06-02-87                        |                                  |
| SULFAMETHOXAZOLE; TRIMETHOPRIM<br>800MG; 160MG        | BACTRIM DS<br>(TABLET; ORAL)   | HOFFMANN-LA ROCHE     | 17-377<br>03-01-78               | RE28636<br>06-02-87                        |                                  |
| SULFAMETHOXAZOLE; TRIMETHOPRIM<br>200MG/5ML; 40MG/5ML | BACTRIM<br>(SUSPENSION; ORAL)  | HOFFMANN-LA ROCHE     | 17-560<br>04-16-75               | RE28636<br>06-02-87                        | I-21<br>09-24-86                 |
| SULFAMETHOXAZOLE; TRIMETHOPRIM<br>200MG/5ML; 40MG/5ML | BACTRIM PEDIATRIC<br>(SUSPENSION; ORAL)                                | HOFFMANN-LA ROCHE     | 17-560<br>12-10-79               | RE28636<br>06-02-87                        | I-21<br>09-24-86                 |
| SULFAMETHOXAZOLE; TRIMETHOPRIM<br>80MG/ML; 16MG/ML    | BACTRIM<br>(INJECTABLE; INJECTION)                                     | HOFFMANN-LA ROCHE     | 18-374<br>06-23-81               | 3551564<br>12-29-87<br>RE28636<br>06-02-87 |                                  |
| SULFAMETHOXAZOLE; TRIMETHOPRIM<br>400MG; 80MG         | SULFAMETHOXAZOLE AND<br>TRIMETHOPRIM<br>(TABLET; ORAL)                 | DRUMMER/PHOENIX       | 18-598<br>05-19-82               |  |                                  |
| SULFAMETHOXAZOLE; TRIMETHOPRIM<br>800MG; 160MG        | SULFAMETHOXAZOLE AND<br>TRIMETHOPRIM DOUBLE STRENGTH<br>(TABLET; ORAL) | DRUMMER/PHOENIX       | 18-598<br>05-19-82               |  |                                  |
| SULFAMETHOXAZOLE; TRIMETHOPRIM<br>200MG/5ML; 40MG/5ML | SULFATRIM PEDIATRIC<br>(SUSPENSION; ORAL)                              | NATL PHARM MFG/BARRE  | 18-615<br>01-07-83               |  |                                  |
| SULFAMETHOXAZOLE; TRIMETHOPRIM<br>200MG/5ML; 40MG/5ML | SULFATRIM<br>(SUSPENSION; ORAL)  | NATL PHARM MFG/BARRE  | 18-615<br>01-07-83               |  |                                  |

**TABLE IV. NDA'S APPROVED FROM 1-1-82 TO 2-28-85 AND NDA'S WITH APPROPRIATE PATENT AND EXCLUSIVITY INFORMATION**

| <u>ACTIVE INGREDIENT(S)<br/>STRENGTH(S)</u>           | <u>TRADE NAME<br/>(DOSAGE FORM; ROUTE)</u>                             | <u>APPLICANT NAME</u> | <u>NDA NO.<br/>APPROVAL DATE</u> | <u>PATENT NO.<br/>EXP. DATE</u>            | <u>EXCLUSIVITY<br/>EXP. DATE</u> |
|---|--|-----------------------|----------------------------------|--|----------------------------------|
| SULFAMETHOXAZOLE; TRIMETHOPRIM<br>200MG/5ML; 40MG/5ML | SMZ-TMP<br>(SUSPENSION; ORAL)  | BIOCRAFT LABS         | 18-812<br>01-28-83               |  |                                  |
| SULFAMETHOXAZOLE; TRIMETHOPRIM<br>200MG/5ML; 40MG/5ML | SMZ-TMP PEDIATRIC<br>(SUSPENSION; ORAL)                                | BIOCRAFT LABS         | 18-812<br>06-10-83               |  |                                  |
| SULFAMETHOXAZOLE; TRIMETHOPRIM<br>400MG; 80MG         | SULFAMETHOXAZOLE AND<br>TRIMETHOPRIM<br>(TABLET; ORAL)                 | DANBURY PHARMACAL     | 18-852<br>05-09-83               |  |                                  |
| SULFAMETHOXAZOLE; TRIMETHOPRIM<br>800MG; 160MG        | SULFAMETHOXAZOLE AND<br>TRIMETHOPRIM DOUBLE STRENGTH<br>(TABLET; ORAL) | DANBURY PHARMACAL     | 18-854<br>05-09-83               |  |                                  |
| SULFAMETHOXAZOLE; TRIMETHOPRIM<br>400MG; 80MG         | SULFAMETHOXAZOLE &<br>TRIMETHOPRIM<br>(TABLET; ORAL)                   | HEATHER DRUG          | 18-946<br>08-10-84               |  |                                  |
| SULFAMETHOXAZOLE; TRIMETHOPRIM<br>800MG; 160MG        | SULFAMETHOXAZOLE &<br>TRIMETHOPRIM<br>(TABLET; ORAL)                   | HEATHER DRUG          | 18-946<br>08-10-84               |  |                                  |
| SULFASALAZINE<br>500MG                                | AZULFIDINE<br>(TABLET, ENTERIC COATED;<br>ORAL)                        | PHARMACIA/PHARMACIA   | 07-073<br>04-06-83               |  | NDF<br>09-24-86                  |
| SULFASALAZINE<br>500MG                                | SULFASALAZINE<br>(TABLET, ENTERIC COATED;<br>ORAL)                     | BOLAR PHARMACEUTICAL  | 88-052<br>05-24-83               |  | NDF<br>09-24-86                  |
| SULINDAC<br>150MG                                     | CLINORIL<br>(TABLET; ORAL)   | MS&D/MERCK            | 17-911<br>09-27-78               | 3654349<br>04-04-89<br>3725548<br>04-03-90 |                                  |
| SULINDAC<br>200MG                                     | CLINORIL<br>(TABLET; ORAL)   | MS&D/MERCK            | 17-911<br>09-27-78               | 3725548<br>04-03-90<br>3654349<br>04-04-89 |                                  |

TABLE IV. NDA'S APPROVED FROM 1-1-82 TO 2-28-85 AND NDA'S WITH APPROPRIATE PATENT AND EXCLUSIVITY INFORMATION

| <u>ACTIVE INGREDIENT(S)<br/>STRENGTH(S)</u>                                   | <u>TRADE NAME<br/>(DOSAGE FORM; ROUTE)</u>         | <u>APPLICANT NAME</u> | <u>NDA NO.<br/>APPROVAL DATE</u> | <u>PATENT NO.<br/>EXP. DATE</u>            | <u>EXCLUSIVITY<br/>EXP. DATE</u> |
|---|--|-----------------------|----------------------------------|--|----------------------------------|
| SUTILAINS<br>82,000 UNITS/GM  | TRAVASE<br>(OINTMENT; TOPICAL)                     | TRAVENOL LABS         | 12-828<br>06-12-69               | 3409719<br>11-05-85                        |                                  |
| TECHNETIUM, TC-99M SODIUM PERTECHNETATE<br>GENERATOR<br>0.22-2.22CI/GENERATOR | MINITEC<br>(SOLUTION; INTRAVENOUS,<br>ORAL)        | ER SQUIBB AND SONS    | 17-339<br>06-03-74               |  | I-31<br>09-24-86                 |
| TECHNETIUM, TC-99M, ALBUMIN COLLOID<br>KIT<br>N/A                             | MICROLITE<br>(INJECTABLE; INJECTION)               | MED DIAG/NE NUCLEAR   | 18-263<br>03-25-83               |  |                                  |
| TECHNETIUM, TC-99M, DISOFENIN KIT<br>N/A                                      | HEPATOLITE<br>(INJECTABLE; INJECTION)              | MED DIAG/NE NUCLEAR   | 18-467<br>03-16-82               |  | NP<br>09-24-86                   |
| TECHNETIUM, TC 99M,<br>PYROPHOSPHATE KIT<br>N/A                               | PHOSPHOTEC<br>(INJECTABLE; INJECTION)              | ER SQUIBB AND SONS    | 17-680<br>10-20-76               |  | I-9<br>09-24-86                  |
| TECHNETIUM, TC-99M, GLUCEPTATE KIT<br>N/A                                     | TECHNISCAN GLUCEPTATE<br>(INJECTABLE; INJECTION)   | MS&D/MERCK            | 18-272<br>01-27-82               |  |                                  |
| TECHNETIUM, TC-99M, MEDRONATE<br>N/A  | OSTEOLITE<br>(INJECTABLE; INJECTION)               | MED DIAG/NE NUCLEAR   | 17-972<br>12-16-77               |  |                                  |
| TECHNETIUM, TC-99M, MEDRONATE<br>N/A  | AMERSCAN<br>(INJECTABLE; INJECTION)                | AMERSHAM/RADIOCHEM    | 18-335<br>08-05-82               |  |                                  |
| TECHNETIUM, TC-99M, SUCCIMER KIT<br>N/A                                       | MPI DMSA KIDNEY REAGENT<br>(INJECTABLE; INJECTION) | MEDI-PHYSICS          | 17-944<br>05-18-82               | 4208398<br>06-17-97<br>4233285<br>11-11-97 | NP<br>09-24-86                   |
| TERBUTALINE SULFATE<br>1MG/ML   | BRICANYL<br>(INJECTABLE; INJECTION)                | MERRELL DOW/DOW CHEM  | 17-466<br>03-25-74               | 3937838<br>02-10-93<br>4011258<br>03-08-94 |                                  |
| TERBUTALINE SULFATE<br>2.5MG  | BRICANYL<br>(TABLET; ORAL)                         | MERRELL DOW/DOW CHEM  | 17-618<br>04-22-75               | 3937838<br>02-10-93<br>4011258<br>03-08-94 |                                  |

TABLE IV. NDA'S APPROVED FROM 1-1-82 TO 2-28-85 AND NDA'S WITH APPROPRIATE PATENT AND EXCLUSIVITY INFORMATION

| <u>ACTIVE INGREDIENT(S)<br/>STRENGTH(S)</u> | <u>TRADE NAME<br/>(DOSAGE FORM; ROUTE)</u>          | <u>APPLICANT NAME</u> | <u>NDA NO.<br/>APPROVAL DATE</u> | <u>PATENT NO.<br/>EXP. DATE</u>            | <u>EXCLUSIVITY<br/>EXP. DATE</u> |
|---|---|-----------------------|----------------------------------|--|----------------------------------|
| TERBUTALINE SULFATE<br>5MG                  | BRICANYL<br>(TABLET; ORAL)                          | MERRELL DOM/DOM CHEM  | 17-618<br>04-22-75               | 3937838<br>02-10-93<br>4011258<br>03-08-94 |                                  |
| TERBUTALINE SULFATE<br>2.5MG                | BRETHINE<br>(TABLET; ORAL)                          | GEIGY/CIBA-GEIGY      | 17-849<br>05-17-76               | 3937838<br>02-10-93<br>4011258<br>03-08-94 |                                  |
| TERBUTALINE SULFATE<br>5MG                  | BRETHINE<br>(TABLET; ORAL)                          | GEIGY/CIBA-GEIGY      | 17-849<br>05-17-76               | 3937838<br>02-10-93<br>4011258<br>03-08-94 |                                  |
| TERBUTALINE SULFATE<br>1MG/ML               | BRETHINE<br>(INJECTABLE; INJECTION)                 | GEIGY/CIBA-GEIGY      | 18-571<br>11-30-81               | 3937838<br>02-10-93<br>4011258<br>03-08-94 |                                  |
| TERBUTALINE SULFATE<br>0.2MG/INH            | BRETHAIRE<br>(AEROSOL; INHALATION)                  | GEIGY/CIBA-GEIGY      | 18-762<br>08-17-84               | 3937838<br>02-10-93<br>4011258<br>03-08-94 | NDF<br>09-24-86                  |
| THALLOUS CHLORIDE, TL-201<br>2MC/ML         | THALLOUS CHLORIDE TL 201<br>(INJECTABLE; INJECTION) | MEDI-PHYSICS          | 18-110<br>02-01-82               |  | NS<br>09-24-86                   |
| THALLOUS CHLORIDE, TL-201<br>1MC/ML         | THALLOUS CHLORIDE TL 201<br>(INJECTABLE; INJECTION) | AMERSHAM/RADIOCHEM    | 18-548<br>12-30-82               |  |                                  |
| TIMOLOL MALEATE<br>5MG                      | BLOCADREN<br>(TABLET; ORAL)                         | MS&D/MERCK            | 18-017<br>11-25-81               | 3655663<br>04-11-89                        |                                  |
| TIMOLOL MALEATE<br>10MG                     | BLOCADREN<br>(TABLET; ORAL)                         | MS&D/MERCK            | 18-017<br>11-25-81               | 3655663<br>04-11-89                        |                                  |
| TIMOLOL MALEATE<br>20MG                     | BLOCADREN<br>(TABLET; ORAL)                         | MS&D/MERCK            | 18-017<br>11-25-81               | 3655663<br>04-11-89                        |                                  |

TABLE IV. NDA'S APPROVED FROM 1-1-82 TO 2-28-85 AND NDA'S WITH APPROPRIATE PATENT AND EXCLUSIVITY INFORMATION

| <u>ACTIVE INGREDIENT(S)<br/>STRENGTH(S)</u> | <u>TRADE NAME<br/>(DOSAGE FORM; ROUTE)</u> | <u>APPLICANT NAME</u> | <u>NDA NO.<br/>APPROVAL DATE</u> | <u>PATENT NO.<br/>EXP. DATE</u>            | <u>EXCLUSIVITY<br/>EXP. DATE</u> |
|---|--|-----------------------|----------------------------------|--|----------------------------------|
| TIMOLOL MALEATE<br>EQ 0.25% BASE            | TIMOPTIC<br>(SOLUTION; OPHTHALMIC)         | MS&D/MERCK            | 18-086<br>08-17-78               | 4195085<br>03-25-97<br>3655663<br>04-11-89 |                                  |
| TIMOLOL MALEATE<br>EQ 0.5% BASE             | TIMOPTIC<br>(SOLUTION; OPHTHALMIC)         | MS&D/MERCK            | 18-086<br>08-17-78               | 4195085<br>03-25-97<br>3655663<br>04-11-89 |                                  |
| TOCAINIDE HYDROCHLORIDE<br>400MG            | TONOCARD<br>(TABLET; ORAL)                 | MS&D/MERCK            | 18-257<br>11-09-84               | 4218477<br>08-19-97<br>4237068<br>12-02-97 | NCE<br>11-09-89                  |
| TOCAINIDE HYDROCHLORIDE<br>600MG            | TONOCARD<br>(TABLET; ORAL)                 | MS&D/MERCK            | 18-257<br>11-09-84               | 4218477<br>08-19-97<br>4237068<br>12-02-97 | NCE<br>11-09-89                  |
| TOLAZAMIDE<br>100MG                         | TOLAZAMIDE<br>(TABLET; ORAL)               | ZENITH LABORATORIES   | 18-894<br>11-02-84               |  |                                  |
| TOLAZAMIDE<br>250MG                         | TOLAZAMIDE<br>(TABLET; ORAL)               | ZENITH LABORATORIES   | 18-894<br>11-02-84               |  |                                  |
| TOLAZAMIDE<br>500MG                         | TOLAZAMIDE<br>(TABLET; ORAL)               | ZENITH LABORATORIES   | 18-894<br>11-02-84               |  |                                  |
| TOLAZOLINE HYDROCHLORIDE<br>25MG/ML         | PRISCOLINE<br>(INJECTABLE; INJECTION)      | CIBA/CIBA-GEIGY       | 06-403<br>02-22-85               |  |                                  |
| TOLMETIN SODIUM<br>EQ 200MG BASE            | TOLECTIN<br>(TABLET; ORAL)                 | MCNEIL LABORATORIES   | 17-628<br>03-24-76               | 3752826<br>08-14-90                        |                                  |
| TOLMETIN SODIUM<br>EQ 400MG BASE            | TOLECTIN DS<br>(CAPSULE; ORAL)             | MCNEIL LABORATORIES   | 18-084<br>10-30-79               | 3752826<br>08-14-90                        |                                  |
| TRAZODONE HYDROCHLORIDE<br>50MG             | DESYREL<br>(TABLET; ORAL)                  | MEAD JOHNSON/B-M      | 18-207<br>12-24-81               | 3381009<br>04-30-85                        |                                  |

TABLE IV. NDA'S APPROVED FROM 1-1-82 TO 2-28-85 AND NDA'S WITH APPROPRIATE PATENT AND EXCLUSIVITY INFORMATION

| <u>ACTIVE INGREDIENT(S)</u><br><u>STRENGTH(S)</u> | <u>TRADE NAME</u><br><u>(DOSAGE FORM; ROUTE)</u> | <u>APPLICANT NAME</u> | <u>NDA NO.</u><br><u>APPROVAL DATE</u> | <u>PATENT NO.</u><br><u>EXP. DATE</u>      | <u>EXCLUSIVITY</u><br><u>EXP. DATE</u> |
|---|--|-----------------------|--|--|--|
| TRAZODONE HYDROCHLORIDE<br>100MG                  | DESYREL<br>(TABLET; ORAL)                        | MEAD JOHNSON/B-M      | 18-207<br>12-24-81                     | 3381009<br>04-30-85                        |  |
| TRETINOIN<br>0.05%                                | RETIN-A<br>(SOLUTION; TOPICAL)                   | ORTHO PHARMACEUTICAL  | 16-921<br>10-20-71                     | 3729568<br>04-24-90                        |  |
| TRETINOIN<br>0.1%                                 | RETIN-A<br>(CREAM; TOPICAL)                      | ORTHO PHARMACEUTICAL  | 17-340<br>01-26-73                     | 3729568<br>04-24-90                        |  |
| TRETINOIN<br>0.05%                                | RETIN-A<br>(CREAM; TOPICAL)                      | ORTHO PHARMACEUTICAL  | 17-522<br>07-19-74                     | 3729568<br>04-24-90                        |  |
| TRETINOIN<br>0.01%                                | RETIN-A<br>(GEL; TOPICAL)                        | ORTHO PHARMACEUTICAL  | 17-955<br>10-05-78                     | 3729568<br>04-24-90                        |  |
| TRETINOIN<br>0.025%                               | RETIN-A<br>(GEL; TOPICAL)                        | ORTHO PHARMACEUTICAL  | 17-579<br>04-18-75                     | 3729568<br>04-24-90                        |  |
| TRIAMCINOLONE ACETONIDE<br>0.25MG/INH             | AZMACORT<br>(AEROSOL; INHALATION)                | WILLIAM H RORER       | 18-117<br>04-23-83                     | 3897779<br>08-05-92<br>3927806<br>12-23-92 | NDF<br>09-24-86                        |
| TRIAZOLAM<br>0.25MG                               | HALCION<br>(TABLET; ORAL)                        | UPJOHN                | 17-892<br>11-15-82                     | 3980790<br>09-14-93<br>3987052<br>10-19-93 | NCE<br>11-15-92                        |
| TRIAZOLAM<br>0.5MG                                | HALCION<br>(TABLET; ORAL)                        | UPJOHN                | 17-892<br>11-15-82                     | 3980790<br>09-14-93<br>3987052<br>10-19-93 | NCE<br>11-15-92                        |
| TRILOSTANE<br>30MG                                | MODRASTANE<br>(CAPSULE; ORAL)                    | WINTHROP LABS/STERL   | 18-719<br>12-21-84                     |  | NCE<br>12-21-89                        |
| TRILOSTANE<br>60MG                                | MODRASTANE<br>(CAPSULE; ORAL)                    | WINTHROP LABS/STERL   | 18-719<br>12-21-84                     |  | NCE<br>12-21-89                        |
| TRIMETHOPRIM<br>200MG                             | PROLOPRIM<br>(TABLET; ORAL)                      | BURROUGHS WELLCOME    | 17-943<br>07-14-82                     |  | NS<br>09-24-86                         |

TABLE IV. NDA'S APPROVED FROM 1-1-82 TO 2-28-85 AND NDA'S WITH APPROPRIATE PATENT AND EXCLUSIVITY INFORMATION

| <u>ACTIVE INGREDIENT(S)<br/>STRENGTH(S)</u> | <u>TRADE NAME<br/>(DOSAGE FORM; ROUTE)</u>                           | <u>APPLICANT NAME</u> | <u>NDA NO.<br/>APPROVAL DATE</u> | <u>PATENT NO.<br/>EXP. DATE</u>                                   | <u>EXCLUSIVITY<br/>EXP. DATE</u> |
|---|--|-----------------------|----------------------------------|---|----------------------------------|
| TRIMETHOPRIM<br>200MG                       | TRIMPEX 200<br>(TABLET; ORAL)  | HOFFMANN-LA ROCHE     | 17-952<br>11-09-82               |   | NS<br>09-24-86                   |
| TRIMETHOPRIM<br>100MG                       | TRIMETHOPRIM<br>(TABLET; ORAL)                                       | BIOCRAFT LABS         | 18-679<br>07-30-82               |   |                                  |
| TRIMPRAMINE MALEATE<br>EQ 100MG BASE        | SURMONTIL<br>(CAPSULE; ORAL)   | IVES LABS/AMHO        | 16-792<br>09-15-82               |   | NS<br>09-24-86                   |
| VECURONIUM BROMIDE<br>10MG/VIAL             | NORCURON (NC-45)<br>(INJECTABLE; INJECTION)                          | ORGANON/AKZONA        | 18-776<br>04-30-84               | 3553212<br>01-05-88<br>4237126<br>12-02-97<br>4297351<br>10-27-98 | NCE<br>04-30-94                  |
| VERAPAMIL HYDROCHLORIDE<br>80MG             | ISOPTIN<br>(TABLET; ORAL)  | KNOLL PHARMACEUTICAL  | 18-593<br>03-08-82               |   | NR<br>09-24-86                   |
| VERAPAMIL HYDROCHLORIDE<br>120MG            | ISOPTIN<br>(TABLET; ORAL)  | KNOLL PHARMACEUTICAL  | 18-593<br>03-08-82               |   | NR<br>09-24-86                   |
| VERAPAMIL HYDROCHLORIDE<br>80MG             | CALAN<br>(TABLET; ORAL)  | SEARLE/SEARLE PHARMS  | 18-817<br>09-10-84               |   | NR<br>09-24-86                   |
| VERAPAMIL HYDROCHLORIDE<br>120MG            | CALAN<br>(TABLET; ORAL)  | SEARLE/SEARLE PHARMS  | 18-817<br>09-10-84               |   | NR<br>09-24-86                   |
| VERAPAMIL HYDROCHLORIDE<br>2.5MG/ML         | CALAN<br>(INJECTABLE; INJECTION)                                     | SEARLE PHARMS         | 18-925<br>03-30-84               |   |                                  |
| VERAPAMIL HYDROCHLORIDE<br>2.5MG/ML         | CALAN<br>(INJECTABLE; INJECTION)                                     | SEARLE PHARMS         | 19-038<br>03-30-84               |   |                                  |
| WATER FOR INJECTION, STERILE<br>100%        | STERILE WATER FOR INJECTION<br>IN PLASTIC CONTAINER<br>(LIQUID; N/A) | TRAVENOL LABS         | 18-595<br>01-17-83               |   |                                  |
| WATER FOR INJECTION, STERILE<br>100%        | STERILE WATER IN PLASTIC<br>CONTAINER<br>(LIQUID; N/A)               | TRAVENOL LABS         | 18-632<br>06-30-82               |   |                                  |

TABLE IV. NDA'S APPROVED FROM 1-1-82 TO 2-28-85 AND NDA'S WITH APPROPRIATE PATENT AND EXCLUSIVITY INFORMATION

| <u>ACTIVE INGREDIENT(S)<br/>STRENGTH(S)</u> | <u>TRADE NAME<br/>(DOSAGE FORM; ROUTE)</u>                           | <u>APPLICANT NAME</u> | <u>NDA NO.<br/>APPROVAL DATE</u> | <u>PATENT NO.<br/>EXP. DATE</u> | <u>EXCLUSIVITY<br/>EXP. DATE</u> |
|---|--|-----------------------|----------------------------------|---------------------------------|----------------------------------|
| WATER FOR INJECTION, STERILE<br>100%        | STERILE WATER IN PLASTIC<br>CONTAINER<br>(LIQUID; N/A)               | ABBOTT LABORATORIES   | 18-801<br>10-27-82               |                                 |                                  |
| WATER FOR INJECTION, STERILE<br>100%        | BACTERIOSTATIC WATER IN<br>PLASTIC CONTAINER<br>(LIQUID; N/A)        | ABBOTT LABORATORIES   | 18-802<br>10-27-82               |                                 |                                  |
| WATER FOR INJECTION, STERILE<br>100%        | STERILE WATER FOR INJECTION<br>IN PLASTIC CONTAINER<br>(LIQUID; N/A) | AM MCGAW/AM HOSP      | 19-077<br>03-02-84               |                                 |                                  |
| XENON, XE-127<br>5MCI/VIAL                  | XENON XE 127<br>(GAS; INHALATION)                                    | MALLINCKRODT          | 18-536<br>10-01-82               |                                 | NCE<br>10-01-92                  |
| XENON, XE-127<br>10MCI/VIAL                 | XENON XE 127<br>(GAS; INHALATION)                                    | MALLINCKRODT          | 18-536<br>10-01-82               |                                 | NCE<br>10-01-92                  |
| XENON, XE-133<br>10MCI/VIAL                 | XENON XE 133<br>(GAS; INHALATION)                                    | MALLINCKRODT          | 18-327<br>03-09-82               |                                 |                                  |
| XENON, XE-133<br>20MCI/VIAL                 | XENON XE 133<br>(GAS; INHALATION)                                    | MALLINCKRODT          | 18-327<br>03-09-82               |                                 |                                  |

