

APPROVED DRUG PRODUCTS

With Therapeutic Equivalence Evaluations



The "Orange Book"

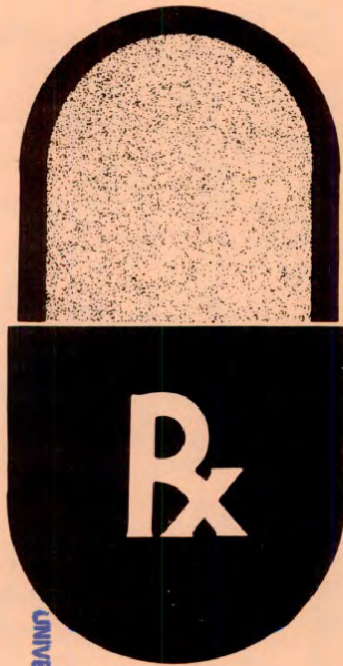
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4th ed.
Suppl. 1

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**CUMULATIVE
SUPPLEMENT 1
AUG '83 - SEP '83**

SERIAL



APPROVED PRESCRIPTION DRUG PRODUCTS

WITH
THERAPEUTIC EQUIVALENCE EVALUATIONS

4TH 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, 4th 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, 1983. 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.

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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 >DELETED (DELETE) to the left of the line containing the overstruck print. The >DELETED 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. SUBSCRIPTION FORM

A subscription form for the publication has been provided at the end of this supplement for use in obtaining additional subscriptions.

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 '83. 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. Also not included in the counts are those duplicate products of a given applicant whose only distinguishing characteristics are items such as package size, inactive ingredient(s), color and alternate manufacturing sites. These various counts are excluded because the Drug Product List itself excludes products from these categories.

REPORT OF COUNTS FOR THE DRUG PRODUCT LIST

A. COUNTS CUMULATIVE BY QUARTERS

JULY '83 (BASELINE)

6679

1908 (28.6%)

4771 (71.4%)

3804 (57.0%)

957 (14.3%)

10 (0.1%)

DRUG PRODUCTS LISTED

SINGLE SOURCE

MULTISOURCE(1)

THERAPEUTICALLY EQUIVALENT

NOT THERAPEUTICALLY EQUIVALENT

EXCEPTIONS(2)

NEW MOLECULAR ENTITIES APPROVED

304

NUMBER OF APPLICANTS

B. ACTIVITY FOR SUPPLEMENT NUMBER 1

AUG '83

SEPT '83

CUMULATIVE

DRUG PRODUCTS ADDED:

NEWLY APPROVED

DESI EFFECTIVE

REMARKETED

REMARKETED

DRUG PRODUCTS REMOVED:

WITHDRAWN APPROVAL

RX TO OTC SWITCH

DISCONTINUED MARKETING

NET GAIN IN DRUG PRODUCTS

SINGLE SOURCE PRODUCTS APPROVED

MULTISOURCE DRUG PRODUCTS APPROVED

NEW MOLECULAR ENTITIES APPROVED:

AS THE ENTITY

AS A SALT, ESTER OR DERIVATIVE

OF THE ENTITY

(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 1 / AUGUST '83 & SEPTEMBER '83

1

AMINOPHYLLINE (PAGE 3-8)

> DLT > /140101; ORAL/
 > ADD > SOLUTION; ORAL
 > ADD > AMINOPHYLLINE
 > ADD > AA ROXANE LABORATORIES 105MG/5ML

TABLET; ORAL

> ADD > AMINOPHYLLINE
 > ADD > BD BARR LABORATORIES 100MG
 > ADD > BD 200MG

> ADD > ASPIRIN; BUTALBITAL; CAFFEINE (PAGE 3-15)

> ADD > TABLET; ORAL
 > ADD > ASPIRIN AND CAFFEINE W/ BUTALBITAL
 > ADD > PUREPAC/KALIPHARMA 325MG;50MG;40MG

ASPIRIN; BUTALBITAL; CAFFEINE; PHENACETIN (PAGE 3-15)

TABLET; ORAL

> DLT > /66/ A.P.C. W/ BUTALBITAL
 > DLT > /PUREPAC/KALIPHARMA/ /200MG;50MG;40MG;150MG/

> ADD > ASPIRIN; CAFFEINE; DIHYDROCODEINE BITARTRATE (PAGE 3-15)

> ADD > CAPSULE; ORAL
 > ADD > SYNALGOS-DC
 > ADD > IVES LABS/AMMO 356.4MG;30MG;16MG

ASPIRIN; CAFFEINE; PHENACETIN; PROPOXYPHENE HYDROCHLORIDE (PAGE 3-15)

CAPSULE; ORAL

> DLT > /66/ PROPOXYPHENE COMPOUND 65
 > DLT > /TONE PAULSEN/ /227MG;32.4MG;162MG;65MG/
 > DLT > /REPRO COMPOUND 65/
 > DLT > /66/ /REID-PROVIDENT LABS/ /227MG;32.4MG;162MG;65MG/
 > DLT > /SK-65 COMPOUND/
 > DLT > /66/ /SK&F LABORATORIES/ /227MG;32.4MG;162MG;65MG/

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

CAPSULE; ORAL

> ADD > DARVON COMPOUND-65
 > ADD > AA ELI LILLY INDUSTRIES/PR 389MG;32.4MG;65MG
 > ADD > SK-65 COMPOUND
 > ADD > AA SK&F LABORATORIES 389MG;32.4MG;65MG

> ADD > ASPIRIN; CARISOPRODOL (PAGE 3-16)

> ADD > TABLET; ORAL
 > ADD > SOMA COMPOUND
 > ADD > WALLACE PHARMS/C-W 325MG;200MG

> ADD > ASPIRIN; CARISOPRODOL; CODEINE PHOSPHATE (PAGE 3-16)

> ADD > TABLET; ORAL
 > ADD > SOMA COMPOUND W/ CODEINE
 > ADD > WALLACE PHARMS/C-W 325MG;200MG;16MG

> ADD > ASPIRIN; HYDROCODONE BITARTRATE (PAGE 3-16)

> ADD > TABLET; ORAL
 > ADD > VICOPRIN
 > ADD > KNOLL PHARMACEUTICAL 500MG;5MG

BACAMPICILLIN HYDROCHLORIDE (PAGE 3-18)

TABLET; ORAL

> ADD > SPECTROBID
 > ADD > PFIZER LABS/PFIZER 800MG

BETAMETHASONE VALERATE (PAGE 3-21)

CREAM; TOPICAL

> ADD > BETAMETHASONE VALERATE
 > ADD > AB E FOUGERA/BYK-GLDN EQ 0.1% BASE
 > ADD > AB PHARMADERM/BYK-GLDN EQ 0.1% BASE
 > ADD > BETATREX
 > ADD > AB SAVAGE LABS/BYK-GLDN EQ 0.1% BASE

LOTION; TOPICAL

> ADD > BETAMETHASONE VALERATE
 > ADD > AB E FOUGERA/BYK-GLDN EQ 0.1% BASE
 > ADD > AB PHARMADERM/BYK-GLDN EQ 0.1% BASE
 > ADD > BETATREX
 > ADD > AB SAVAGE LABS/BYK-GLDN EQ 0.1% BASE
 > ADD > AB VALTSONE
 > ADD > AB SCHERING EQ 0.1% BASE

OINTMENT; TOPICAL

> ADD > BETAMETHASONE VALERATE
 > ADD > AB E FOUGERA/BYK-GLDN EQ 0.1% BASE
 > ADD > AB PHARMADERM/BYK-GLDN EQ 0.1% BASE
 > ADD > BETATREX
 > ADD > AB SAVAGE LABS/BYK-GLDN EQ 0.1% BASE
 > ADD > AB VALTSONE
 > ADD > AB SCHERING EQ 0.1% BASE

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DEXAMETHASONE (PAGE 3-49)

> ADD > SOLUTION; ORAL
> ADD > DEXAMETHASONE
> ADD > ROXANE LABORATORIES 0.5MG/5MLX
> ADD > DEXAMETHASONE INTENSOL
> ADD > ROXANE LABORATORIES 0.5MG/0.5MLX

TABLET; ORAL
> ADD > BP DECAIDRON 6MG
> ADD > BP MS&D/MERCK 6MG
> ADD > BP DEXAMETHASONE 6MG
> ADD > BP ROXANE LABORATORIES 1MGX

DEXAMETHASONE SODIUM PHOSPHATE (PAGE 3-50)

> DLT > INJECTABLE; INJECTION
> ADD > ~~HEXADROL PHOSPHATE~~
> ADD > AP HEXADROL
> ADD > AP ORGANON/AKZONA EQ 4MG PHOSPHATE/ML
> ADD > EQ 20MG PHOSPHATE/ML

DEXTROSE; DOPAMINE HYDROCHLORIDE (PAGE 3-53)

> DLT > INJECTABLE; INJECTION
> ADD > ~~DOPAMINE HCL IN DEXTROSE 5%~~
> ADD > AP DOPAMINE HCL
> ADD > AP ABBOTT LABORATORIES 5GM/100ML; 80MG/100ML
> ADD > AP 5GM/100ML; 160MG/100ML
> ADD > AP DOPAMINE HCL IN PLASTIC CONTAINER
> ADD > AP ABBOTT LABORATORIES 5GM/100ML; 80MG/500MLX
> ADD > AP 5GM/100ML; 160MG/100MLX
> ADD > AP 5GM/100ML; 320MG/100MLX

DIETHYLPROPION HYDROCHLORIDE (PAGE 3-59)

TABLET; ORAL
> ADD > AA DIETHYLPROPION HCL
> ADD > AA CAMALL 25MGX

DIPHENHYDRAMINE HYDROCHLORIDE (PAGE 3-61)

CAPSULE; ORAL
> DLT > ~~SK-DIPHENHYDRAMINE~~
> ADD > SK&F LABORATORIES /12MG/
ELIXIR; ORAL
> DLT > ~~SK-DIPHENHYDRAMINE~~
> DLT > ~~SK&F LABORATORIES~~ /12.5MG/5ML/

DISULFIRAM (PAGE 3-63)

TABLET; ORAL
> ADD > BX DISULFIRAM
> ADD > BX CHELSEA LABORATORIES 250MGX
> ADD > BX 500MGX

DOXYCYCLINE HYCLATE (PAGE 3-64)

CAPSULE; ORAL
> ADD > AB DOXYCYCLINE HYCLATE
> ADD > AB CHELSEA LABORATORIES EQ 50MG BASE

EPINEPHRINE; LIDOCAINE HYDROCHLORIDE (PAGE 3-66)

INJECTABLE; INJECTION
> ADD > LIDOCATON
> ADD > AP PHARMATON/SZ 0.01MG/ML; 22X
> ADD > AP 0.02MG/ML; 22X

ERYTHROMYCIN (PAGE 3-67)

OINTMENT; OPHTHALMIC
> ADD > ERYTHROMYCIN
> ADD > AT E FOUGERA/BYK-GLDN 5MG/GMX
> ADD > AT PHARMADERM/BYK-GLDN 5MG/GMX
> ADD > AT TLOTYCIN
> ADD > AT DISTA PRODS/LILLY 5MG/GM

GENTAMICIN SULFATE (PAGE 3-79)

CREAM; TOPICAL
> ADD > GENTAFAIR
> ADD > AT PHARMAFAIR EQ 1MG BASE/GMX
> ADD > GENTAMICIN SULFATE
> ADD > AT NMC LABORATORIES EQ 1MG BASE/GMX

INJECTABLE; INJECTION
> ADD > GENTAMICIN SULFATE
> ADD > AP ABBOTT LABORATORIES EQ 60MG BASE/100MLX
> ADD > AP EQ 70MG BASE/100MLX
> ADD > AP EQ 80MG BASE/100MLX
> ADD > AP EQ 90MG BASE/100MLX
> ADD > AP EQ 100MG BASE/100MLX
> ADD > AP EQ 1.2MG BASE/MLX
> ADD > AP EQ 1.4MG BASE/MLX
> ADD > AP EQ 1.6MG BASE/MLX
> ADD > AP EQ 1.8MG BASE/MLX
> ADD > AP EQ 2MG BASE/MLX
> ADD > AP EQ 10MG BASE/MLX
> ADD > AP EQ 40MG BASE/MLX
> ADD > AP EQ 40MG BASE/MLX

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HYDROXYZINE PAMOATE (PAGE 3-96)CAPSULE; ORAL
HYDROXYZINE PAMOATE

> ADD > AB VANGARD LABS/HNM EQ 25MG HCLM
> ADD > AB EQ 50MG HCLM

IMIPRAMINE HYDROCHLORIDE (PAGE 3-97)

TABLET; ORAL

> DLT > /AP/ /ANTIPRESS/ /25MG/
> DLT > /BP/ /LEMMON/ /25MG/
> DLT > /AP/ /IMAVATE/ /25MG/
> DLT > /AB/ /AM ROBIN/ /25MG/
> DLT > /AB/ /50MG/

ISOETHARINE HYDROCHLORIDE (PAGE 3-100)

SOLUTION; INHALATION

ISOETHARINE HCL

> ADD > AM INTL MEDICATION SYS 0.167%
> ADD > AM ROXANE LABORATORIES 0.167%
> ADD > AM TRAVENOL LABS 0.25%

KANAMYCIN SULFATE (PAGE 3-102)

INJECTABLE; INJECTION

KANAMYCIN SULFATE

> ADD > AP INTL MEDICATION SYS EQ 500MG BASE/2MLM
> ADD > AP EQ 1GM BASE/2MLM
> DLT > /AP/ /KANTREX/ /75MG/2ML/
> DLT > /AP/ /BRISTOL LABS/B-M/ /500MG/2ML/
> DLT > /AP/ /1GM/2ML/
> ADD > AP BRISTOL LABS/B-M EQ 75MG BASE/2ML
> ADD > AP EQ 500MG BASE/2ML
> ADD > AP EQ 1GM BASE/2ML

LIDOCAINE HYDROCHLORIDE (PAGE 3-104)

INJECTABLE; INJECTION

LIDOCATON

> ADD > AP PHARMATON/SZ 22%

METHYLTESTOSTERONE (PAGE 3-121)

TABLET; BUCCAL/SUBLINGUAL

METHYLTESTOSTERONE

> DLT > /BP/ /JOHNE PAULSEN/ /10MG/

METRONIDAZOLE (PAGE 3-122)

INJECTABLE; INJECTION

METRO T.V. IN PLASTIC CONTAINER

> ADD > AP AM MCGAW/AM HOSP 500MG/100MLM

TABLET; ORAL

METRONIDAZOLE

> ADD > AB PAR PHARMACEUTICAL 250MG
> ADD > AB 500MG

NANDROLONE DECANOATE (PAGE 3-125)

INJECTABLE; INJECTION

NANDROLONE DECANOATE

> ADD > MAURRY BIOLOGICAL 100MG/MLM

NITROGLYCERIN (PAGE 3-128)

INJECTABLE; INJECTION

NITROHAL

> ADD > AP G POHL-BOSKAMP 5MG/MLM
> ADD > 1MG/MLM

TRIDIL

> ADD > AM CRITICAL CARE/AHS 0.5MG/MLM

OXTRIPHYLLINE (PAGE 3-131)

TABLET, ENTERIC COATED; ORAL

CHOLEDYL

> ADD > AB PARKE-DAVIS/W-L 100MG
> ADD > AB 200MG

OXTRIPHYLLINE

> ADD > AB BOLAR PHARMACEUTICAL 100MG
> ADD > AB 200MG

OXYTETRACYCLINE HYDROCHLORIDE (PAGE 3-132)

CAPSULE; ORAL

OXYLOXAS

> DLT > /AB/ /PARKE-DAVIS/W-L/ /EQ 250MG BASE/

PENICILLIN V POTASSIUM (PAGE 3-135)

POWDER FOR RECONSTITUTION; ORAL

PENICILLIN VK

> DLT > /AP/ /J.P. H.V. /EQ 125MG BASE/2ML/
> DLT > /AA/ /EQ 250MG BASE/2ML/

PSEUDOEPHEDRINE HYDROCHLORIDE; TRIPROlidine HYDROCHLORIDE (PAGE 3-155)

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DRUG PRODUCT LIST / CUMULATIVE SUPPLEMENT NUMBER 1 / AUGUST '83 & SEPTEMBER '83

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PHENIDMETRAZINE TARTRATE (PAGE 3-138)

TABLET, ORAL
PHENMETRAZINE TARTRATE
FERNDAL LABS
55MG

PHENTERMINE HYDROCHLORIDE (PAGE 3-139)

CAPSULE; ORAL
ADIPEX-P
LEMON
> ADD >
37.5MGx

POTASSIUM CHLORIDE (PAGE 3-143)

INJECTABLE; INJECTION

POTASSIUM CHLORIDE

TRAYENOL LABS/

INTEG/NL

INTEG/NL

INTEG/NL

INTEG/NL

< DLT >/AP

< DLT >/AP

< DLT >/AP

< DLT >/AP

PREDNISOLONE SODIUM PHOSPHATE (PAGE 3-147)

SOLUTION/DROPS; OPHTHALMIC	< DLT >	< DLT >/4T	< DLT >/4T
/HYDROLYSOL			
/MSD/NECK			
METREON			
SCHERING			
EO 0.5% PHOSPHATE			
/EO 0.5% PHOSPHATE			

PROCAINAMIDE HYDROCHLORIDE (PAGE 3-149)

CAPSOLE, ORAL
 PROGRAM
 PARKE-DAVIS/M-L
 /50MG/
 /50MG/

PROPOXYPHENE HYDROCHLORIDE (PAGE 3-153)

CAPSULE; ORAL
PROPXYPHENE HCL
RICHLYN LABORATORIES 65MG
> ADD > AA

PROPYLTHIOURACIL (PAGE 3-154)

PROPLITHIURACIL /HVLN.PHAMS/ /50MS/

PSEUDOEPHEDRINE HYDROCHLORIDE; TRIPROLINE HYDROCHLORIDE (PAGE 3-155)

SYRUP; ORAL
PSEUDODINE
BAY LABORATORIES
30MB/5ML; 1.25MB/5ML
/TILAGIN/
/NATL PHARM N°6/BAYNE/30MB/5ML; 1.25MB/5ML
> 0.17 >/AA/
> 0.16 >
> 0.17 >/AA/

TABLET, ORAL
/TID/ /DAILY/ /60mg/ /AND/
/AA/

QUINIDINE SULFATE (PAGE 3-157)
TABLET; ORAL
QUINIDINE SULFATE
VITARINE/WEST CHEM 300MGX
> ADD > AB

SELENIUM SULFIDE (PAGE 3-161)

LOTION/SHAMPOO; TOPICAL
SELENIUM SULFIDE
BAY LABORATORIES
2.52%

CREAM; TOPICAL
/STIVER, S.I./EAT/4474#

> 550 < />
> 011 < /ST05TEROL; ALPHA/ (PAGE 3-161)

[illegible]

SOMATROPIN (PAGE 3-165)

ASCELLACRIN 2

< ADD >
ASCELLACRIN 10

SPIRONOLACTONE (PAGE 3-166)

TABLET; ORAL
SPIRONOLACTONE

>_ADD_> AB PUREPAC/KALIPHARMA 25MG

TESTOSTERONE (PAGE 3-173)

INJECTABLE; INJECTION
TESTOSTERONE

>_ADD_> CARTER-GLOGAU LABS 50MG/ML

TETRACYCLINE HYDROCHLORIDE (PAGE 3-174)

TABLET; ORAL
PANMYCIN

>_DLT_>/AB/ UPJOHN /250MG/

>_DLT_>/AB/ SUNMYCIN
ER SQUIBB AND SONS 250MG

THEOPHYLLINE (PAGE 3-176)

CAPSULE, CONTROLLED RELEASE; ORAL
SOMOPHYLLIN-CRT

>_ADD_> BC FISONS 100MG

>_ADD_> THEO-24

>_ADD_> BC SEARLE/SEARLE PHARMS 100MG

>_ADD_> 200MG

>_ADD_> 300MG

SOLUTION; ORAL

THEOLAIR

>_ADD_> AA RIKER LABS/3M 80MG/15ML

>_ADD_> THEOPHYLLINE

>_ADD_> AA ROXANE LABORATORIES 80MG/15ML

TABLET, CONTROLLED RELEASE; ORAL
THEOCONTIN

>_ADD_> BC PURDUE FREDERICK 200MG

THEO-DUR

>_ADD_> BC KEY PHARMACEUTICALS 200MG

THIORIDAZINE HYDROCHLORIDE (PAGE 3-178)

CONCENTRATE; ORAL
THIORIDAZINE HCL

>_ADD_> AA NATL PHARM MFG/BARRE 100MG/ML

THIORIDAZINE HYDROCHLORIDE (PAGE 3-178)

TABLET; ORAL
MELLARTIL

>_ADD_> AB SANDOZ PHARMS/SANDOZ 100MG

THIORIDAZINE HCL

>_ADD_> AB BOLAR PHARMACEUTICAL 10MG

>_ADD_> AB 100MG

>_ADD_> AB CORD LABORATORIES 10MG

>_ADD_> AB 15MG

>_ADD_> AB 25MG

>_ADD_> AB 50MG

TIMOLOL MALEATE (PAGE 3-179)

TABLET; ORAL
BLOCADREN

>_ADD_> MS&D/MERCK 5MG

TRIAMCINOLONE ACETONIDE (PAGE 3-180)

CREAM; TOPICAL

TRIAMCINOLONE ACETONIDE

>_ADD_> AT BAY LABORATORIES 0.025%

>_ADD_> AT 0.1%

>_ADD_> AT 0.5%

OINTMENT; TOPICAL

TRIAMCINOLONE ACETONIDE

>_ADD_> AT BAY LABORATORIES 0.025%

>_ADD_> AT 0.1%

>_ADD_> AT 0.5%

TRISULFAPYRIMIDINES (PAGE 3-185)

SUSPENSION; ORAL

>_DLT_> /TRISULF/

>_DLT_>/AB/ /BEECHAM LABS/BEECHAM/500MG/5ML/

>_DLT_> /TRISUREID/

>_DLT_>/AB/ /REID-PROVIDENT LABS/500MG/5ML/

TABLET; ORAL

>_DLT_> /TRIPLE SULFA 333/

>_DLT_>/AB/ /ZENITH LABORATORIES/500MG/

TROPICAMIDE (PAGE 3-186)

SOLUTION/DROPS; OPHTHALMIC

HYDRIFAIR

>_ADD_> PHARMAFAIR 0.5%

>_ADD_> AT 1%

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> DLT > /AA/ /JONE, PAULSEN/ /EG 50,000 UNITS BASE/

VITAMIN A PALMITATE (PAGE 3-168)

> DLT > /AA/ /JONE, PAULSEN/ /EG 50,000 UNITS BASE/

> DLT > /TYBATH/ (PAGE 3-166)

DRUG PRODUCT LIST / CUMULATIVE SUPPLEMENT NUMBER 1 / AUGUST '83 & SEPTEMBER '83

8

> DLT > /CARBAMIPHEN EMBUTYLATE/ CHLORPHENTRAMINE MALEATE/ ISOPROPANTINE/
> DLT > /IODIDES/ PHENYLPROPANOLAMINE HYDROCHLORIDE/ (PAGE AD3)
> DLT > /CAPSULE/ CONTROLLED RELEASE/ ORAL/
> DLT > /TUSS-ORNADE/
> DLT > /Saf. LABORATORIES/ /20mg; 40mg; 60mg; 80mg/
> DLT > /SOLUTION/ ORAL/
> DLT > /TUSS-ORNADE/
> DLT > /Saf. LABORATORIES/ /5mg; 10mg; 15mg; 20mg; 25mg; 30mg; 35mg; 40mg; 45mg; 50mg; 55mg; 60mg; 65mg; 70mg; 75mg; 80mg; 85mg; 90mg; 95mg; 100mg;

DIPYRIDAMOLE (PAGE AD4)

TABLET; ORAL
DIPYRIDAMOLE
> ADD > ASCOT HOSP PHARMS 50MG

CURRENT STATUS - INEFFECTIVE

DESIGN PENDING LIST - OTHER THAN 'EXEMPT' (COURT ORDER) CATEGORY
CUMULATIVE SUPPLEMENT NUMBER 1 / AUGUST / '83 & SEPTEMBER / '83

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4TH EDITION (1983)**

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