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GENERAL NOTICE ALGEMENE KENNISGEWING

NOTICE 1269 OF 2008

MEDICINES CONTROL COUNCIL

CONDITIONS OF REGISTRATION OF A MEDICINE IN TERMS OF THE PROVISIONS OF SECTION 15(7) OF THE MEDICINES AND RELATED SUBSTANCES ACT, 1965 (ACT 101 OF 1965)

1. The applicant shall ensure that the medicine is manufactured and controlled in terms of the current Good Manufacturing Practices as determined by Council.
2. The manufacture of this medicine is subject to regular investigation and inspections by the inspectors appointed in terms of Section 26 of the Act, to assess compliance with current Good Manufacturing Practices.
3. The information in the package insert shall be updated on a regular basis to conform to the package insert recently approved by Council.
4. The applicant must comply with all the legal requirements of the Medicines and Related Substances Act, 1965 (Act 101 of 1965).
5. The registration of this medicine shall be subject to review at intervals as determined by Council regarding its quality, safety and efficacy, and the registration of this medicine may be varied subject to issues Council may deem fit.
6. The first two production batches must be fully validated in terms of the detailed process validation protocol submitted at the time of application for registration, and the validation report must be submitted within a month after completion of the validation.
7. The product may be advertised to the professions only.
8. A post-registration inspection must be conducted in the first production batch of the locally manufactured product.
9. A post-registration inspection must be conducted on the first production batch manufactured by each local manufacturer.
10. A post-registration inspection must be conducted on the first production batch of the imported product.
11. Marketing of the product may only commence following a satisfactory post-registration inspection report.
12. One sample of every batch, together with four copies of the protocol for testing of the bulk lot and filling lot, and six copies of the certificate of release issued by a competent authority in the country in which the product was manufactured, must be submitted to the Council for lot release purposes.
13. The expiry date allocated shall be modified by adding a statement that the virus strains are currently recommended for South African usage in the specific year.
14. The strains of the master seed viruses must be approved by the Department of Health for each year.

KENNISGEWING 1269 VAN 2008**MEDISYNEBEHEERRAAD****VOORWAARDES VIR DIE REGISTRASIE VAN 'N MEDISYNE IN TERME VAN DIE
BEPALINGS VAN ARTIKEL 15(7) VAN DIE WET OP BEHEER VAN MEDISYNE EN
VERWANTE STOWWE, 1965 (WET 101 VAN 1965)**

1. Die applikant sal verseker dat die medisyne vervaardig en beheer word ooreenkomstig huidige Goeie Vervaardigingspraktyke soos bepaal deur die Medisynebeheerraad.
2. Die vervaardiging van hierdie medisyne is onderhewig aan gereelde ondersoeke en inspeksies deur inspekteurs, aangestel ingevolge Artikel 25 van die Wet, om die nakoming van Goeie Vervaardigingspraktyke te bepaal.
3. Die inligting soos vervat in the voubiljet moet op 'n gereelde grondslag opgedateer word in ooreenstemming met 'n voubiljet wat onlangs deur die Raad goedgekeur is.
4. Die applikant moet voldoen aan alle wetlike vereistes van die Wet op Medisyne en Verwante Stowwe, 1965 (Wet 101 van 1965).
5. Die registrasie van hierdie medisyne is onderhewig aan gereelde hersiening met tussenpose soos deur die Raad bepaal, rakende kwaliteit, veiligheid en effektiwiteit, en die registrasie van hierdie medisyne kan gewysig word onderhewig aan kwessies na goeddunke van die Raad.
6. Die eerste twee produksielotte moet ten volle gevalideer word ooreenkomstig die breedvoerige prosesvalidasie protokol wat ingedien is ten tye van die aansoek om registrasie, en die validasieverslag moet binne die bestek van een maand na die voltooiing van die validasie ingedien word.
7. Die produk mag slegs aan die beroepe geadverteer word.
8. 'n Na-registrasie-inspeksie moet op die eerste produksielot van die plaaslik vervaardigde produk uitgevoer word.
9. 'n Na-registrasie-inspeksie moet op die eerste produksielot van elke plaaslike vervaardiger uitgevoer word.
10. 'n Na-registrasie-inspeksie moet op die eerste produksielot van die ingevoerde produk uitgevoer word.
11. Bemaking van die produk mag slegs 'n aanvang neem nadat 'n bevredigende na-registrasie-inspeksieverslag gedien het.
12. Een monster van elke lot moet tesame met vier kopieë van die protokolle vir die toets van die massalot en die vullot sowel as ses kopieë van die vrystellingsertifikaat wat uitgereik is deur die verantwoordelike beheerliggaam in die land waar die produk vervaardig word, ingedien word by die Raad vir lotvrystellingsdoeleindes.
13. Die vervaldatum toegeken, sal gewysig word deur 'n verklaring by te voeg dat die virusstamme tans vir Suid-Afrikaanse gebruik gedurende die gespesifiseerde jaar aanbeveel word.
14. Die stamme van die oorspronklike saadvirusse moet elke jaar deur die Departement van Gesondheid goedgekeur word.

MRF 15

Registration number:	36/2.2/0235
Name of medicine:	DOXYNITE DROPS
Dosage form:	DROPS
Active ingredients:	EACH 0,5 ml DROPS CONTAIN: DOXYLAMINE SUCCINATE 25,0 mg
Conditions of registration:	1, 2, 3, 4, 5, 6, 7
Applicant:	EQUITY PHARMACEUTICALS (PTY) LTD
Manufacturer:	PHARMA-Q, INDUSTRIA, JOHANNESBURG
Packer:	PHARMA-Q, INDUSTRIA, JOHANNESBURG
Laboratory:	FPRC: PHARMA-Q, INDUSTRIA, JOHANNESBURG
	FPRR: EQUITY PHARMACEUTICALS, HAZELWOOD, PRETORIA
Shelf-life:	24 months
Date of registration:	15 AUGUST 2008

MRF 15

Registration number:	37/20.2.2/0100
Name of medicine:	TIBOZOLE
Dosage form:	TABLET
Active ingredients:	EACH TABLET CONTAINS: MICONAZOLE NITRATE 10,0 mg
Conditions of registration:	1, 2, 3, 4, 5, 6, 7
Applicant:	JANSSEN PHARMACEUTICA (PTY) LTD
Manufacturer:	CLONMEL HEALTHCARE LTD, CLONMEL, IRELAND
Packer:	SANICO NV, TURNHOUT, BELGIUM JANSSEN PHARMACEUTICA, WOODMEAD, JOHANNESBURG
Laboratory:	FPRC: CLONMEL HEALTHCARE LTD, CLONMEL, IRELAND SANICO NV, TURNHOUT, BELGIUM
	FPRC/FPRR: JANSSEN PHARMACEUTICA, WOODMEAD, JOHANNESBURG
Shelf-life:	30 months
Date of registration:	15 AUGUST 2008

MRF 15

Registration number:	37/7.1/0302
Name of medicine:	FEDALOC 30 mg SR
Dosage form:	TABLET
Active ingredients:	EACH TABLET CONTAINS: NIFEDIPINE 30,0 mg
Conditions of registration:	1, 2, 3, 4, 5, 6, 7, 8
Applicant:	COMPUPHARM (PTY) LTD
Manufacturer:	VALPHARMA SA, SERRAVALLA, REPUBLIC OF SAN MARINO EUDERMA S.p.A., RIMINI, ITALY
Packer:	VALPHARMA SA, SERRAVALLA, REPUBLIC OF SAN MARINO EUDERMA S.p.A., RIMINI, ITALY
Laboratory:	FPRC: VALPHARMA SA, SERRAVALLA, REPUBLIC OF SAN MARINO EUDERMA S.p.A., RIMINI, ITALY SEDEK AGRIKEM, KAMEELDRIFT, PRETORIA INSTITUTE FOR PHARMACEUTICAL SERVICES, SILVERTONDALE, PRETORIA
	FPRR: COMPUPHARM, LYNNWOOD, PRETORIA
Shelf-life:	24 months (Provisional)
Date of registration:	15 AUGUST 2008

MRF 15

Registration number:	37/7.1/0303
Name of medicine:	FEDALOC 60 mg SR
Dosage form:	TABLET
Active ingredients:	EACH TABLET CONTAINS: NIFEDIPINE 60,0 mg
Conditions of registration:	1, 2, 3, 4, 5, 6, 7, 8
Applicant:	COMPUPHARM (PTY) LTD
Manufacturer:	VALPHARMA SA, SERRAVALLA, REPUBLIC OF SAN MARINO EUDERMA S.p.A., RIMINI, ITALY
Packer:	VALPHARMA SA, SERRAVALLA, REPUBLIC OF SAN MARINO EUDERMA S.p.A., RIMINI, ITALY
Laboratory:	FPRC: VALPHARMA SA, SERRAVALLA, REPUBLIC OF SAN MARINO EUDERMA S.p.A., RIMINI, ITALY SEDEK AGRIKEM, KAMEELDRIFT, PRETORIA INSTITUTE FOR PHARMACEUTICAL SERVICES, SILVERTONDALE, PRETORIA
	FPRR: COMPUPHARM, LYNNWOOD, PRETORIA
Shelf-life:	24 months (Provisional)
Date of registration:	15 AUGUST 2008

MRF 15

Registration number:	37/8.3/0434
Name of medicine:	COSMOFER
Dosage form:	SOLUTION
Active ingredients:	EACH 1,0 ml SOLUTION CONTAINS: IRON DEXTRAN EQUIVALENT TO ELEMENTARY IRON 50,0 mg
Conditions of registration:	1, 2, 3, 4, 5, 6, 7
Applicant:	PHARMAPLAN (PTY) LTD
Manufacturer:	SOLUPHARM GmbH, MELSUNGEN, GERMANY WEIMAR PHARMA GmbH, RASTATT, GERMANY
Packer:	SOLUPHARM GmbH, MELSUNGEN, GERMANY WEIMAR PHARMA GmbH, RASTATT, GERMANY
Laboratory:	FPRC: SOLUPHARM GmbH, MELSUNGEN, GERMANY WEIMAR PHARMA GmbH, RASTATT, GERMANY PHARMACOSMOS, SOENDERGADE, VIBY, DENMARK CONSULTING CHEMICAL LABORATORIES, ATLASVILLE, BOKSBURG
	FPRR: PHARMAPLAN, MIDRAND, RSA
Shelf-life:	36 months
Date of registration:	15 AUGUST 2008

MRF 15

Registration number:	37/20.2.2/0630
Name of medicine:	MICMAT
Dosage form:	TABLET
Active ingredients:	EACH TABLET CONTAINS: MICONAZOLE NITRATE 10,0 mg
Conditions of registration:	1, 2, 3, 4, 5, 6, 7
Applicant:	JANSSEN PHARMACEUTICA (PTY) LTD
Manufacturer:	CLONMEL HEALTHCARE LTD, TIPPERARY, CLONMEL, IRELAND
Packer:	SANICO NV, TURNHOUT, BELGIUM JANSSEN PHARMACEUTICA, WOODMEAD, SANDTON
Laboratory:	FPRC: CLONMEL HEALTHCARE LTD, TIPPERARY, CLONMEL, IRELAND SANICO NV, TURNHOUT, BELGIUM
	FPRC/FPRR: JANSSEN PHARMACEUTICA, WOODMEAD, SANDTON
Shelf-life:	24 months
Date of registration:	15 AUGUST 2008

MRF 15

Registration number: 38/20.2.2/0157
 Name of medicine: CANDIZOLE VAGINAL TABLET
 Dosage form: TABLET
 Active ingredients: EACH TABLET CONTAINS:
 CLOTRIMAZOLE 500,0 mg
 Conditions of registration: 1, 2, 3, 4, 5, 6, 7
 Applicant: PHARMACARE LIMITED
 Manufacturer: TECHNIKON LABORATORIES, ROBERTVILLE,
 FLORIDA
 COLUMBIA PHARMACEUTICALS, BARDENE,
 BOKSBURG
 Packer: TECHNIKON LABORATORIES, ROBERTVILLE,
 FLORIDA
 COLUMBIA PHARMACEUTICALS, BARDENE,
 BOKSBURG
 Laboratory: FPRC: TECHNIKON LABORATORIES, ROBERTVILLE,
 FLORIDA
 COLUMBIA PHARMACEUTICALS, BARDENE,
 BOKSBURG
 SOUTH AFRICAN BUREAU OF STANDARDS,
 GROENKLOOF, PRETORIA
 FPRC/FPRR: PHARMACARE LTD, WOODMEAD, SANDTON
 Shelf-life: 24 months
 Date of registration: 15 AUGUST 2008

MRF 15

Registration number: A05/3.1.1/03
 Name of medicine: PETCAM
 Dosage form: SUSPENSION
 Active ingredients: EACH 1,0 ml SUSPENSION CONTAINS:
 MELOXICAM 1,5 mg
 Conditions of registration: 1, 2, 3, 4, 5, 6, 7
 Applicant: CIPLA MEDPRO (PTY) LTD
 Manufacturer: CIPLA LTD (UNIT I), SALCETTE, GOA, INDIA
 Packer: CIPLA LTD (UNIT I), SALCETTE, GOA, INDIA
 Laboratory: FPRC: CIPLA LTD (UNIT I), SALCETTE, GOA, INDIA
 FPRR: CIPLA MEDPRO, ROSENPARK, BELLVILLE
 Shelf-life: 36 months
 Date of registration: 15 AUGUST 2008

MRF 15

Registration number: A05/3.1.1/04
 Name of medicine: RIMADYL AQUEOUS INJECTION
 Dosage form: INJECTION
 Active ingredients: EACH 1,0 ml SOLUTION CONTAINS:
 CARPROFEN 50,0 mg

 Conditions of registration: 1, 2, 3, 4, 5, 6, 7
 Applicant: PFIZER LABORATORIES (PTY) LTD
 Manufacturer: VERICORE LTD, DUNDEE, SCOTLAND
 Packer: VERICORE LTD, DUNDEE, SCOTLAND
 Laboratory: FPRC: VERICORE LTD, DUNDEE, SCOTLAND

 FPRR: PFIZER LABORATORIES, SANDTON,
 JOHANNESBURG
 Shelf-life: 36 months
 Date of registration: 15 AUGUST 2008

MRF 15

Registration number: A38/4/0412
 Name of medicine: SEPTANEST WITH ADRENALINE 1/100 000
 Dosage form: INJECTION
 Active ingredients: EACH 2,2 ml CARTRIDGE CONTAINS:
 ARTICAIN HYDROCHLORIDE 88,0 mg
 ADRENALINE TARTRATE EQUIVALENT TO
 ADRENALINE 22,0 ug

 Conditions of registration: 1, 2, 3, 4, 5, 6, 7
 Applicant: THE DENTAL WAREHOUSE (PTY) LTD
 Manufacturer: SEPTODONT, SAINT MAUR DES FOSSES,
 FRANCE
 Packer: SEPTODONT, SAINT MAUR DES FOSSES,
 FRANCE
 Laboratory: FPRC: SEPTODONT, SAINT MAUR DES FOSSES,
 FRANCE
 SOUTH AFRICAN BUREAU OF STANDARDS,
 GROENKLOOF, PRETORIA
 CONSULTING CHEMICAL LABORATORIES,
 ATLASVILLE, BOKSBURG, RSA

 FPRR: THE DENTAL WAREHOUSE, SANDTON,
 RSA
 Shelf-life: 24 months
 Date of registration: 15 AUGUST 2008

MRF 15

Registration number: A38/7.1.3/0604
 Name of medicine: ASPEN ENALAPRIL 10 mg
 Dosage form: TABLET
 Active ingredients: EACH TABLET CONTAINS:
 ENALAPRIL MALEATE 10,0 mg
 Conditions of registration: 1, 2, 3, 4, 5, 6, 7, 8
 Applicant: PHARMACARE LIMITED
 Manufacturer: GENPHARM INC, ETOBICOKE, ONTARIO,
 CANADA
 MERCK FARMA y QUIMICA S.A, MOLLET DEL
 VALLES, BARCELONA, SPAIN
 Packer: GENPHARM INC, ETOBICOKE, ONTARIO,
 CANADA
 MERCK FARMA y QUIMICA S.A, MOLLET DEL
 VALLES, BARCELONA, SPAIN
 GERARD LABORATORIES, DUBLIN, IRELAND
 GENERICS UK LTD, POTTERS BAR,
 HERTFORDSHIRE, UK
 Laboratory: FPRC: GENPHARM INC, ETOBICOKE, ONTARIO,
 CANADA
 MERCK FARMA y QUIMICA S.A, MOLLET DEL
 VALLES, BARCELONA, SPAIN
 SOUTH AFRICAN BUREAU OF STANDARDS,
 GROENKLOOF, PRETORIA
 RESEARCH INSTITUTE FOR INDUSTRIAL
 PHARMACY, NORTH-WEST UNIVERSITY,
 POTCHEFSTROOM
 FPRR PHARMACARE LTD, WOODMEAD, SANDTON
 Shelf-life: 24 months (Provisional)
 Date of registration: 15 AUGUST 2008

MRF 15

Registration number: A38/7.1.3/0605
 Name of medicine: ASPEN ENALAPRIL 20 mg
 Dosage form: TABLET
 Active ingredients: EACH TABLET CONTAINS:
 ENALAPRIL MALEATE 20,0 mg
 Conditions of registration: 1, 2, 3, 4, 5, 6, 7, 8
 Applicant: PHARMACARE LIMITED
 Manufacturer: GENPHARM INC, ETOBICOKE, ONTARIO,
 CANADA
 MERCK FARMA y QUIMICA S.A, MOLLET DEL
 VALLES, BARCELONA, SPAIN
 Packer: GENPHARM INC, ETOBICOKE, ONTARIO,
 CANADA
 MERCK FARMA y QUIMICA S.A, MOLLET DEL
 VALLES, BARCELONA, SPAIN
 GERARD LABORATORIES, DUBLIN, IRELAND
 GENERICS UK LTD, POTTERS BAR,
 HERTFORDSHIRE, UK
 Laboratory: FPRC: GENPHARM INC, ETOBICOKE, ONTARIO,
 CANADA
 MERCK FARMA y QUIMICA S.A, MOLLET DEL
 VALLES, BARCELONA, SPAIN
 SOUTH AFRICAN BUREAU OF STANDARDS,
 GROENKLOOF, PRETORIA
 RESEARCH INSTITUTE FOR INDUSTRIAL
 PHARMACY, NORTH-WEST UNIVERSITY,
 POTCHEFSTROOM
 FPRR PHARMACARE LTD, WOODMEAD, SANDTON
 Shelf-life: 24 months (Provisional)
 Date of registration: 15 AUGUST 2008

MRF 15

Registration number: A40/2.5/0004
 Name of medicine: ASPEN PHENYTOIN SODIUM 250 mg/5 ml
 Dosage form: INJECTION
 Active ingredients: EACH 5,0 ml SOLUTION CONTAINS:
 PHENYTOIN SODIUM 250,0 mg

 Conditions of registration: 1, 2, 3, 4, 5, 6, 7, 8
 Applicant: PHARMACARE LIMITED
 Manufacturer: LABORATORIO REIG JOFRE S.A, BARCELONA, SPAIN
 STRIDES ARCOLAB LTD, BANGALORE, INDIA
 Packer: LABORATORIO REIG JOFRE S.A, BARCELONA, SPAIN
 STRIDES ARCOLAB LTD, BANGALORE, INDIA
 Laboratory: FPRC: LABORATORIO REIG JOFRE S.A, BARCELONA, SPAIN
 STRIDES ARCOLAB LTD, BANGALORE, INDIA
 SOUTH AFRICAN BUREAU OF STANDARDS,
 GROENKLOOF, PRETORIA
 RESEARCH INSTITUTE FOR PHARMACEUTICAL
 SERVICES, NORTH-WEST UNIVERSITY,
 POTCHEFSTROOM
 M&L LABORATORY SERVICES, ORMONDE,
 JOHANNESBURG
 FPRR: PHARMACARE LTD, WOODMEAD, SANDTON
 Shelf-life: 24 months (provisional)
 Date of registration: 15 AUGUST 2008

MRF 15

Registration number: A40/20.1.1/0114
 Name of medicine: CIPROCINA
 Dosage form: INFUSION
 Active ingredients: EACH 1,0 ml SOLUTION CONTAINS:
 CIPROFLOXACIN LACTATE EQUIVALENT
 TO
 CIPROFLOXACIN 2,0 mg

 Conditions of registration: 1, 2, 3, 4, 5, 6, 7, 8
 Applicant: PHARMAPLAN (PTY) LTD
 Manufacturer: CLARIS LIFESCIENCES LTD, ELLISBRIDGE,
 AHMEDABAD, INDIA
 Packer: CLARIS LIFESCIENCES LTD, ELLISBRIDGE,
 AHMEDABAD, INDIA
 Laboratory: FPRC: CLARIS LIFESCIENCES LTD, ELLISBRIDGE,
 AHMEDABAD, INDIA
 CONSULTING CHEMICAL LABORATORIES,
 ATLASVILLE, BOKSBURG, RSA
 FPRR: PHARMAPLAN, MIDRAND, RSA
 Shelf-life: 24 months (Provisional)
 Date of registration: 15 AUGUST 2008

MRF 15

Registration number:	A40/3.1/0174
Name of medicine:	PEDEA
Dosage form:	INJECTION
Active ingredients:	EACH AMPOULE CONTAINS: IBUPROFEN 10,0 mg
Conditions of registration:	1, 2, 3, 4, 5, 6, 7, 8
Applicant:	TEMA MEDICAL (PTY) LTD
Manufacturer:	MERCKLE GmbH, BLAUBEUREN, GERMANY
Packer:	MERCKLE GmbH, BLAUBEUREN, GERMANY
Laboratory:	FPRC: MERCKLE GmbH, BLAUBEUREN, GERMANY TECHNIKON LABORATORIES, ROBERTVILLE, FLORIDA
	FPRR: TEMA MEDICAL, SANDTON, JOHANNESBURG
Shelf-life:	24 months (Provisional)
Date of registration:	15 AUGUST 2008

MRF 15

Registration number:	A40/2.9/0203
Name of medicine:	DUROGESIC 12 µg/h
Dosage form:	TRANSDERMAL PATCH
Active ingredients:	EACH PATCH CONTAINS: FENTANYL 2,1 mg
Conditions of registration:	1, 2, 3, 4, 5, 6, 7, 8
Applicant:	JANSSEN PHARMACEUTICA (PTY) LTD
Manufacturer:	ALZA IRELAND LTD, CASHEL, COUNTY TIPPERARY, IRELAND
Packer:	ALZA IRELAND LTD, CASHEL, COUNTY TIPPERARY, IRELAND JANSSEN PHARMACEUTICA NV, BEERSE, BELGIUM
Laboratory:	FPRC: ALZA IRELAND LTD, CASHEL, COUNTY TIPPERARY, IRELAND JANSSEN PHARMACEUTICA NV, BEERSE, BELGIUM MICROCHEM LABORATORIES, DUNGARVAN COUNTY, WATERFORD, IRELAND
	FPRC/FPRR: JANSSEN PHARMACEUTICA, WOODMEAD, JOHANNESBURG, RSA
Shelf-life:	24 months (Provisional)
Date of registration:	15 AUGUST 2008

MRF 15

Registration number: A40/7.1.3/0292
 Name of medicine: CO-SARBEN 150/12,5
 Dosage form: TABLET
 Active ingredients: EACH TABLET CONTAINS:
 IRBESARTAN 150,0 mg
 HYDROCHLOROTHIAZIDE 12,5 mg
 Conditions of registration: 1, 2, 3, 4, 5, 6, 7
 Applicant: SANOFI-SYNTHELABO (PTY) LTD
 Manufacturer: BRISTOL-MYERS SQUIBB CO., EVANSVILLE
 INDIANA, USA
 SANOFI WINTHROP INDUSTRIE, AMBARES, FRANCE
 Packer: SANOFI WINTHROP INDUSTRIE, AMBARES, FRANCE
 AVENTIS PHARMA, WALTLOO, PRETORIA
 PHARMACEUTICAL CONTRACTORS, ISANDO
 Laboratory: FPRC: SANOFI WINTHROP INDUSTRIE, AMBARES, FRANCE
 M&L LABORATORY SERVICES, ORMONDE,
 JOHANNESBURG
 FPRC/FPRR: AVENTIS PHARMA, WALTLOO, PRETORIA
 FPRR: SANOFI-SYNTHELABO, MIDRAND, RSA
 Shelf-life: 36 months
 Date of registration: 15 AUGUST 2008

MRF 15

Registration number: A40/7.1.3/0293
 Name of medicine: CO-SARBEN 300/12,5
 Dosage form: TABLET
 Active ingredients: EACH TABLET CONTAINS:
 IRBESARTAN 300,0 mg
 HYDROCHLOROTHIAZIDE 12,5 mg
 Conditions of registration: 1, 2, 3, 4, 5, 6, 7
 Applicant: SANOFI-SYNTHELABO (PTY) LTD
 Manufacturer: BRISTOL-MYERS SQUIBB CO., EVANSVILLE
 INDIANA, USA
 SANOFI WINTHROP INDUSTRIE, AMBARES,
 FRANCE
 Packer: SANOFI WINTHROP INDUSTRIE, AMBARES,
 FRANCE
 AVENTIS PHARMA, WALTLOO, PRETORIA
 PHARMACEUTICAL CONTRACTORS, ISANDO
 Laboratory: FPRC: SANOFI WINTHROP INDUSTRIE, AMBARES,
 FRANCE
 M&L LABORATORY SERVICES, ORMONDE,
 JOHANNESBURG
 FPRC/FPRR: AVENTIS PHARMA, WALTLOO, PRETORIA
 FPRR: SANOFI-SYNTHELABO, MIDRAND, RSA
 Shelf-life: 36 months
 Date of registration: 15 AUGUST 2008

MRF 15

Registration number: A40/34/0502
 Name of medicine: AVONEX SOLUTION FOR INJECTION
 Dosage form: INJECTION
 Active ingredients: EACH PRE-FILLED SYRINGE CONTAINS:
 INTERFERON BETA 1a 30,0 µg
 Conditions of registration: 1, 2, 3, 4, 5, 6, 7
 Applicant: PHARMAPLAN (PTY) LTD
 Manufacturer: VETTER PHARMA-FERTIGUNG GmbH & CO,
 SCHUTZENSTRASSE, RAVENBURG, GERMANY
 Packer: VETTER PHARMA-FERTIGUNG GmbH & CO,
 SCHUTZENSTRASSE, RAVENBURG, GERMANY
 BIOGEN IDEC B.V, HOOFDORP, THE NETHERLANDS
 VETTER PHARMA-FERTIGUNG GmbH & CO,
 LAGENARGEN, GERMANY
 VETTER PHARMA-FERTIGUNG GmbH & CO,
 HOLBEINSTRASSE, RAVENSBURG, GERMANY
 Laboratory: FPRC: VETTER PHARMA-FERTIGUNG GmbH & CO,
 SCHUTZENSTRASSE, RAVENBURG, GERMANY
 BIOGEN IDEC B.V, HOOFDORP, THE NETHERLANDS
 BIOGEN IDEC B.V, AMSTERDAM, THE NETHERLANDS
 BIORELIANCE LTD, STIRLING, SCOTLAND, UK
 COVANCE LABORATORIES LTD, HARROGATE, NORTH
 YORKSHIRE, UK
 CONSULTING CHEMICAL LABORATORIES,
 ATLASVILLE, BOKSBURG
 FPRR: PHARMAPLAN, MIDRAND
 Shelf-life: 18 months
 Date of registration: 15 AUGUST 2008

MRF 15

Registration number: A40/21.12/0656
 Name of medicine: DRL-FINASTERIDE 1 mg
 Dosage form: TABLET
 Active ingredients: EACH TABLET CONTAINS:
 FINASTERIDE 1,0 mg
 Conditions of registration: 1, 2, 3, 4, 5, 6, 7, 8
 Applicant: DR REDDY'S LABORATORIES (PTY) LTD
 Manufacturer: DR REDDY'S LABORATORIES LTD, RANGA
 REDDY DISTRICT, ANDHRA PRADESH, INDIA
 Packer: DR REDDY'S LABORATORIES LTD, RANGA
 REDDY DISTRICT, ANDHRA PRADESH, INDIA
 DRA PHARMACEUTICALS, IRENE, CENTURION
 Laboratory: FPRC: DR REDDY'S LABORATORIES LTD, RANGA
 REDDY DISTRICT, ANDHRA PRADESH, INDIA
 INSTITUTE FOR PHARMACEUTICALS SERVICES,
 SILVERTONDALE, PRETORIA
 RESEARCH INSTITUTE FOR INDUSTRIAL
 PHARMACY, NORTH-WEST UNIVERSITY,
 POTCHEFSTROOM
 FPRR: DR REDDY'S LABORATORIES, MURRAYFIELD
 PRETORIA
 Shelf-life: 24 months (Provisional)
 Date of registration: 15 AUGUST 2008

MRF 15

Registration number: A40/7.5/0711
 Name of medicine: GULF PRAVASTATIN 20
 Dosage form: TABLET
 Active ingredients: EACH TABLET CONTAINS:
 PRAVASTATIN SODIUM 20,0 mg

 Conditions of registration: 1, 2, 3, 4, 5, 6, 7
 Applicant: GULF DRUG COMPANY (PTY) LTD
 Manufacturer: PT NOVELL PHARMACEUTICAL LABORATORIES,
 GUNUNG PUTRI, BOGOR, INDONESIA
 ALEMBIC LTD, PANCHMAHALS, GUJARAT, INDIA

 Packer: PT NOVELL PHARMACEUTICAL LABORATORIES,
 GUNUNG PUTRI, BOGOR, INDONESIA
 ALEMBIC LTD, PANCHMAHALS, GUJARAT, INDIA

 Laboratory: FPRC: PT NOVELL PHARMACEUTICAL LABORATORIES,
 GUNUNG PUTRI, BOGOR, INDONESIA
 ALEMBIC LTD, PANCHMAHALS, GUJARAT, INDIA
 SOUTH AFRICAN BUREAU OF STANDARDS,
 GROENKLOOF, PRETORIA
 M & L LABORATORY SERVICES, ORMONDE,
 JOHANNESBURG
 CONSULTING CHEMICAL LABORATORIES,
 ATLASVILLE, BOKSBURG
 PHARMA-Q, INDUSTRIA WEST, JOHANNESBURG
 INSTITUTE FOR PHARMACEUTICAL SERVICES,
 SILVERTONDALE, PRETORIA, RSA
 CONSULTING MICROBIOLOGICAL LABORATORY,
 MOREWILL, BEYERSPARK, BOKSBURG

 FPRR: GULF DRUG COMPANY, MOUNT EDGEcombe

 Shelf-life: 36 months

 Date of registration: 15 AUGUST 2008

MRF 15

Registration number: A40/20.1.1/0734
 Name of medicine: OREOXAL 40 mg/5 ml
 Dosage form: SUSPENSION
 Active ingredients: EACH 5,0 ml SUSPENSION CONTAINS:
 CEFPODOXIME PROXETIL EQUIVALENT TO
 CEFPODOXIME 40,0 mg

 Conditions of registration: 1, 2, 3, 4, 5, 6, 7
 Applicant: SANDOZ S.A. (PTY) LTD
 Manufacturer: SANDOZ PVT LTD, TURBHE, NAVI MUMBAI,
 INDIA

 Packer: SANDOZ PVT LTD, TURBHE, NAVI MUMBAI,
 INDIA

 Laboratory: FPRC: SANDOZ PVT LTD, TURBHE, NAVI MUMBAI,
 INDIA
 NOVARTIS, SPARTAN, KEMPTON PARK
 ANALYTICON, TERENURE, KEMPTON PARK
 SOUTH AFRICAN BUREAU OF STANDARDS,
 GROENKLOOF, PRETORIA

 FPRR: SANDOZ S.A., SPARTAN, KEMPTON PARK

 Shelf-life: 24 months

 Date of registration: 15 AUGUST 2008

MRF 15

Registration number: A40/20.2.8/0736
Name of medicine: STOCRIN 50 mg
Dosage form: TABLET
Active ingredients: EACH TABLET CONTAINS:
EFAVIRENZ 50,0 mg
Conditions of registration: 1, 2, 3, 4, 5, 6, 7
Applicant: MSD (PTY) LTD
Manufacturer: MERCK SHARP & DOHME, SOUTH GRANVILLE,
NEW SOUTH WALES, AUSTRALIA
Packer: MERCK SHARP & DOHME, SOUTH GRANVILLE,
NEW SOUTH WALES, AUSTRALIA
MERCK SHARP & DOHME BV, HAARLEM, THE
NETHERLANDS
MSD, HAFWAY HOUSE, RSA
Laboratory: FPRC: MERCK SHARP & DOHME, SOUTH GRANVILLE,
NEW SOUTH WALES, AUSTRALIA
FPRC/FPRR: MSD, HAFWAY HOUSE, RSA
Shelf-life: 24 months
Date of registration: 15 AUGUST 2008

MRF 15

Registration number: A40/20.2.8/0737
Name of medicine: STOCRIN 200 mg
Dosage form: TABLET
Active ingredients: EACH TABLET CONTAINS:
EFAVIRENZ 200,0 mg
Conditions of registration: 1, 2, 3, 4, 5, 6, 7
Applicant: MSD (PTY) LTD
Manufacturer: MERCK SHARP & DOHME, SOUTH
GRANVILLE, NEW SOUTH WALES,
AUSTRALIA
Packer: MERCK SHARP & DOHME, SOUTH
GRANVILLE, NEW SOUTH WALES,
AUSTRALIA
MERCK SHARP & DOHME BV, HAARLEM,
THE NETHERLANDS
MSD, HAFWAY HOUSE, RSA
Laboratory: FPRC: MERCK SHARP & DOHME, SOUTH
GRANVILLE, NEW SOUTH WALES,
AUSTRALIA
FPRC/FPRR: MSD, HAFWAY HOUSE, RSA
Shelf-life: 24 months
Date of registration: 15 AUGUST 2008

MRF 15

Registration number:	41/7.1.3/0042
Name of medicine:	DIACE CO 20 TABLETS
Dosage form:	TABLET
Active ingredients:	EACH TABLET CONTAINS: LISINOPRIL DIHYDRATE EQUIVALENT TO LISINOPRIL 20,0 mg HYDROCHLOROTHIAZIDE 12,5 mg
Conditions of registration:	1, 2, 3, 4, 5, 6, 7, 8
Applicant:	ZYDUS HEALTHCARE S.A. (PTY) LTD
Manufacturer:	ZYDUS CADILA HEALTHCARE LTD, SANAND, AHMEDABAD, INDIA
Packer:	ZYDUS CADILA HEALTHCARE LTD, SANAND, AHMEDABAD, INDIA
Laboratory:	FPRC: ZYDUS CADILA HEALTHCARE LTD, SANAND, AHMEDABAD, INDIA INSTITUTE FOR PHARMACEUTICAL SERVICES, SILVERTONDALE, PRETORIA RESEARCH INSTITUTE FOR INDUSTRIAL PHARMACY, NORTH-WEST UNIVERSITY, POTCHEFSTROOM
	FPRR: ZYDUS HEALTHCARE, VAN DER HOFF PARK, POTCHEFSTROOM
Shelf-life:	24 months (Provisional)
Date of registration:	15 AUGUST 2008

MRF 15

Registration number:	41/5.10/0075
Name of medicine:	DANTRON 4 VIAL
Dosage form:	INJECTION
Active ingredients:	EACH 2,0 ml SOLUTION CONTAINS: ONDANSETRON HYDROCHLORIDE DIHYDRATE EQUIVALENT TO ONDANSETRON 4,0 mg
Conditions of registration:	1, 2, 3, 4, 5, 6, 7, 8
Applicant:	SANDOZ S.A. (PTY) LTD
Manufacturer:	SANDOZ CANADA Inc, BOUCHERVILLE, QUEBEC, CANADA
Packer:	SANDOZ CANADA Inc, BOUCHERVILLE, QUEBEC, CANADA NOVARTIS S.A., SPARTAN, KEMPTON PARK
Laboratory:	FPRC: SANDOZ CANADA Inc, BOUCHERVILLE, QUEBEC, CANADA NOVARTIS S.A., SPARTAN, KEMPTON PARK SOUTH AFRICAN BUREAU OF STANDARDS, GROENKLOOF, PRETORIA
	FPRR: SANDOZ S.A., SPARTAN, KEMPTON PARK
Shelf-life:	24 months (Provisional)
Date of registration:	15 AUGUST 2008

MRF 15

Registration number: 41/5.10/0076
 Name of medicine: DANTRON 8 VIAL
 Dosage form: INJECTION
 Active ingredients: EACH 4,0 ml SOLUTION CONTAINS:
 ONDANSETRON HYDROCHLORIDE DIHYDRATE
 EQUIVALENT TO ONDANSETRON 8,0 mg

 Conditions of registration: 1, 2, 3, 4, 5, 6, 7, 8
 Applicant: SANDOZ S.A. (PTY) LTD
 Manufacturer: SANDOZ CANADA Inc, BOUCHERVILLE, QUEBEC,
 CANADA
 Packer: SANDOZ CANADA Inc, BOUCHERVILLE, QUEBEC,
 CANADA
 NOVARTIS S.A., SPARTAN, KEMPTON PARK

 Laboratory: FPRC: SANDOZ CANADA Inc, BOUCHERVILLE, QUEBEC,
 CANADA
 NOVARTIS S.A., SPARTAN, KEMPTON PARK
 SOUTH AFRICAN BUREAU OF STANDARDS,
 GROENKLOOF, PRETORIA

 FPRR: SANDOZ S.A., SPARTAN, KEMPTON PARK
 Shelf-life: 24 months (Provisional)
 Date of registration: 15 AUGUST 2008

MRF 15

Registration number: 41/5.10/0077
 Name of medicine: NAUSETRON 4 VIAL
 Dosage form: INJECTION
 Active ingredients: EACH 2,0 ml SOLUTION CONTAINS:
 ONDANSETRON HYDROCHLORIDE
 DIHYDRATE EQUIVALENT TO
 ONDANSETRON 4,0 mg

 Conditions of registration: 1, 2, 3, 4, 5, 6, 7, 8
 Applicant: SANDOZ S.A. (PTY) LTD
 Manufacturer: SANDOZ CANADA Inc, BOUCHERVILLE,
 QUEBEC, CANADA
 Packer: SANDOZ CANADA Inc, BOUCHERVILLE,
 QUEBEC, CANADA
 NOVARTIS S.A., SPARTAN, KEMPTON
 PARK

 Laboratory: FPRC: SANDOZ CANADA Inc, BOUCHERVILLE,
 QUEBEC, CANADA
 NOVARTIS S.A., SPARTAN, KEMPTON
 PARK
 SOUTH AFRICAN BUREAU OF STANDARDS,
 GROENKLOOF, PRETORIA

 FPRR: SANDOZ S.A., SPARTAN, KEMPTON PARK
 Shelf-life: 24 months (Provisional)
 Date of registration: 15 AUGUST 2008

MRF 15

Registration number: 41/5.10/0078
 Name of medicine: NAUSETRON 8 VIAL
 Dosage form: INJECTION
 Active ingredients: EACH 4,0 ml SOLUTION CONTAINS:
 ONDANSETRON HYDROCHLORIDE DIHYDRATE
 EQUIVALENT TO ONDANSETRON 8,0 mg

 Conditions of registration: 1, 2, 3, 4, 5, 6, 7, 8
 Applicant: SANDOZ S.A. (PTY) LTD
 Manufacturer: SANDOZ CANADA Inc, BOUCHERVILLE, QUEBEC,
 CANADA

 Packer: SANDOZ CANADA Inc, BOUCHERVILLE, QUEBEC,
 CANADA
 NOVARTIS S.A., SPARTAN, KEMPTON PARK

 Laboratory: FPRC: SANDOZ CANADA Inc, BOUCHERVILLE, QUEBEC,
 CANADA
 NOVARTIS S.A., SPARTAN, KEMPTON PARK
 SOUTH AFRICAN BUREAU OF STANDARDS,
 GROENKLOOF, PRETORIA

 FPRR: SANDOZ S.A., SPARTAN, KEMPTON PARK

 Shelf-life: 24 months (Provisional)
 Date of registration: 15 AUGUST 2008

MRF 15

Registration number: 41/7.1.3/0187
 Name of medicine: LYSIN CO 10
 Dosage form: TABLET
 Active ingredients: EACH TABLET CONTAINS:
 LISINOPRIL DIHYDRATE EQUIVALENT TO
 LISINOPRIL 10,0 mg
 HYDROCHLOROTHIAZIDE 12,5 mg

 Conditions of registration: 1, 2, 3, 4, 5, 6, 7
 Applicant: BE-TABS PHARMACEUTICALS (PTY) LTD
 Manufacturer: AUROBINDO PHARMA LTD, UNIT III, RANGA
 REDDY DISTRICT, ANDHRA PRADESH,
 INDIA

 Packer: AUROBINDO PHARMA LTD, UNIT III, RANGA
 REDDY DISTRICT, ANDHRA PRADESH,
 INDIA

 Laboratory: FPRC: AUROBINDO PHARMA LTD, UNIT III, RANGA
 REDDY DISTRICT, ANDHRA PRADESH,
 INDIA

 FPRR: BE-TABS PHARMACEUTICALS,
 ROODEPOORT, RSA

 Shelf-life: 24 months
 Date of registration: 15 AUGUST 2008

MRF 15		
Registration number:	41/7.1.3/0188	
Name of medicine:	LYSIN CO 20	
Dosage form:	TABLET	
Active ingredients:	EACH TABLET CONTAINS: LISINOPRIL DIHYDRATE EQUIVALENT TO LISINOPRIL 20,0 mg HYDROCHLOROTHIAZIDE 12,5 mg	
Conditions of registration:	1, 2, 3, 4, 5, 6, 7	
Applicant:	BE-TABS PHARMACEUTICALS (PTY) LTD	
Manufacturer:	AUROBINDO PHARMA LTD, UNIT III, RANGA REDDY DISTRICT, ANDHRA PRADESH, INDIA	
Packer:	AUROBINDO PHARMA LTD, UNIT III, RANGA REDDY DISTRICT, ANDHRA PRADESH, INDIA	
Laboratory:	FPRC:	AUROBINDO PHARMA LTD, UNIT III, RANGA REDDY DISTRICT, ANDHRA PRADESH, INDIA
	FPRR:	BE-TABS PHARMACEUTICALS, ROODEPOORT, RSA
Shelf-life:	24 months	
Date of registration:	15 AUGUST 2008	

MRF 15		
Registration number:	41/2.5/271	
Name of medicine:	LAMOD 5	
Dosage form:	TABLET	
Active ingredients:	EACH TABLET CONTAINS: LAMOTRIGINE 5,0 mg	
Conditions of registration:	1, 2, 3, 4, 5, 6, 7	
Applicant:	ARROW PHARMA SOUTH AFRICA (PTY) LTD	
Manufacturer:	ARROW PHARMA (MALTA), HAL FAR, MALTA	
Packer:	ARROW PHARMA (MALTA), HAL FAR, MALTA DIVPHARM MANUFACTURING & PACKAGING, LONGDALE, JOHANNESBURG TECHNIKON LABORATORIES, ROBERTVILLE, FLORIDA	
Laboratory:	FPRC:	ARROW PHARMA (MALTA), HAL FAR, MALTA SELAMINE t/a ARROW GENERICS, DUBLIN, IRELAND TECHNIKON LABORATORIES, ROBERTVILLE, FLORIDA M&L LABORATORY SERVICES, ORMONDE, JOHANNESBURG
	FPRR:	ARROW PHARMA SA, WOODMEAD, SANDTON 24 months
Shelf-life	24 months	
Date of registration:	15 AUGUST 2008	

MRF 15

Registration number: 41/2.5/272
 Name of medicine: LAMOD 25
 Dosage form: TABLET
 Active ingredients: EACH TABLET CONTAINS:
 LAMOTRIGINE 25,0 mg
 Conditions of registration: 1, 2, 3, 4, 5, 6, 7
 Applicant: ARROW PHARMA SOUTH AFRICA (PTY) LTD
 Manufacturer: ARROW PHARMA (MALTA), HAL FAR, MALTA
 Packer: ARROW PHARMA (MALTA), HAL FAR, MALTA
 DIVPHARM MANUFACTURING & PACKAGING,
 LONGDALE, JOHANNESBURG
 TECHNIKON LABORATORIES, ROBERTVILLE,
 FLORIDA
 Laboratory: FPRC: ARROW PHARMA (MALTA), HAL FAR, MALTA
 SELAMINE t/a ARROW GENERICS, DUBLIN,
 IRELAND
 TECHNIKON LABORATORIES, ROBERTVILLE,
 FLORIDA
 M&L LABORATORY SERVICES, ORMONDE,
 JOHANNESBURG
 FPRR: ARROW PHARMA SA, WOODMEAD,
 SANDTON
 Shelf-life: 24 months
 Date of registration: 15 AUGUST 2008

MRF 15

Registration number: 41/2.5/273
 Name of medicine: LAMOD 50
 Dosage form: TABLET
 Active ingredients: EACH TABLET CONTAINS:
 LAMOTRIGINE 50,0 mg
 Conditions of registration: 1, 2, 3, 4, 5, 6, 7
 Applicant: ARROW PHARMA SOUTH AFRICA (PTY) LTD
 Manufacturer: ARROW PHARMA (MALTA), HAL FAR, MALTA
 Packer: ARROW PHARMA (MALTA), HAL FAR, MALTA
 DIVPHARM MANUFACTURING & PACKAGING, LONGDALE,
 JOHANNESBURG
 TECHNIKON LABORATORIES, ROBERTVILLE, FLORIDA
 Laboratory: FPRC: ARROW PHARMA (MALTA), HAL FAR, MALTA
 SELAMINE t/a ARROW GENERICS, DUBLIN, IRELAND
 TECHNIKON LABORATORIES, ROBERTVILLE, FLORIDA
 M&L LABORATORY SERVICES, ORMONDE, JOHANNESBURG
 FPRR: ARROW PHARMA SA, WOODMEAD, SANDTON
 Shelf-life: 24 months
 Date of registration: 15 AUGUST 2008

MRF 15

Registration number: 41/2.5/274
 Name of medicine: LAMOD 100
 Dosage form: TABLET
 Active ingredients: EACH TABLET CONTAINS:
 LAMOTRIGINE 100,0 mg
 Conditions of registration: 1, 2, 3, 4, 5, 6, 7
 Applicant: ARROW PHARMA SOUTH AFRICA (PTY) LTD
 Manufacturer: ARROW PHARMA (MALTA), HAL FAR, MALTA
 Packer: ARROW PHARMA (MALTA), HAL FAR, MALTA
 DIVPHARM MANUFACTURING & PACKAGING,
 LONGDALE, JOHANNESBURG
 TECHNIKON LABORATORIES, ROBERTVILLE,
 FLORIDA
 Laboratory: FPRC: ARROW PHARMA (MALTA), HAL FAR, MALTA
 SELAMINE t/a ARROW GENERICS, DUBLIN, IRELAND
 TECHNIKON LABORATORIES, ROBERTVILLE,
 FLORIDA
 M&L LABORATORY SERVICES, ORMONDE,
 JOHANNESBURG
 FPRR: ARROW PHARMA SA, WOODMEAD, SANDTON
 Shelf-life: 24 months
 Date of registration: 15 AUGUST 2008

MRF 15

Registration number: 41/2.5/275
 Name of medicine: LAMOD 200
 Dosage form: TABLET
 Active ingredients: EACH TABLET CONTAINS:
 LAMOTRIGINE 200,0 mg
 Conditions of registration: 1, 2, 3, 4, 5, 6, 7
 Applicant: ARROW PHARMA SOUTH AFRICA (PTY) LTD
 Manufacturer: ARROW PHARMA (MALTA), HAL FAR, MALTA
 Packer: ARROW PHARMA (MALTA), HAL FAR, MALTA
 DIVPHARM MANUFACTURING & PACKAGING,
 LONGDALE, JOHANNESBURG
 TECHNIKON LABORATORIES,
 ROBERTVILLE, FLORIDA
 Laboratory: FPRC: ARROW PHARMA (MALTA), HAL FAR, MALTA
 SELAMINE t/a ARROW GENERICS, DUBLIN, IRELAND
 TECHNIKON LABORATORIES,
 ROBERTVILLE, FLORIDA
 M&L LABORATORY SERVICES, ORMONDE,
 JOHANNESBURG
 FPRR: ARROW PHARMA SA, WOODMEAD, SANDTON
 Shelf-life: 24 months
 Date of registration: 15 AUGUST 2008

MRF 15

Registration number: 41/20.2.8/0296
 Name of medicine: VARI-ZIDOVUDINE 300 mg
 Dosage form: TABLET
 Active ingredients: EACH TABLET CONTAINS:
 ZIDOVUDINE 300,0 mg

 Conditions of registration: 1, 2, 3, 4, 5, 6, 7
 Applicant: LEBASI PHARMACEUTICALS CC

 Manufacturer: VARICHEM, PHARMACEUTICALS, HARARE,
 ZIMBABWE
 Packer: VARICHEM, PHARMACEUTICALS, HARARE,
 ZIMBABWE
 Laboratory: FPRC: VARICHEM, PHARMACEUTICALS, HARARE,
 ZIMBABWE
 RESEARCH INSTITUTE FOR INDUSTRIAL
 PHARMACY, NORTH-WEST UNIVERSITY,
 POTCHEFSTROOM
 INSTITUTE FOR PHARMACEUTICAL SERVICES,
 SILVERTONDALE, PRETORIA
 CONSULTING CHEMICAL LABORATORIES,
 ATLASVILLE, BOKSBURG

 FPRR: LEBASI PHARMACEUTICALS, NOORDBRUG,
 POTCHEFSTROOM

 Shelf-life: 24 months
 Date of registration: 15 AUGUST 2008

MRF 15

Registration number: 41/18/0299
 Name of medicine: FOSRENOL 250 mg
 Dosage form: TABLET
 Active ingredients: EACH TABLET CONTAINS:
 LANTHANUM CARBONATE HYDRATE
 EQUIVALENT TO LANTHANUM 250,0 mg

 Conditions of registration: 1, 2, 3, 4, 5, 6, 7, 8
 Applicant: ADCOCK INGRAM CRITICAL CARE (PTY)
 LTD

 Manufacturer: BCM LTD, NOTTINGHAM,
 NOTTINGHAMSHIRE, UK
 Packer: BCM LTD, NOTTINGHAM,
 NOTTINGHAMSHIRE, UK
 Laboratory: FPRC: BCM LTD, NOTTINGHAM,
 NOTTINGHAMSHIRE, UK
 READING SCIENTIFIC SERVICES LTD,
 READING, BERKSHIRE, UK
 SOUTHERN TESTING & RESEARCH,
 WILSON, NORTH CAROLINA, USA
 SSCI Inc, WEST LAFAYETTE,
 INDIANAPOLIS, USA
 TEPNEL SCIENTIFIC SERVICES,
 EDINBURGH, YORKSHIRE, UK
 M&L LABORATORY SERVICES, ORMONDE,
 JOHANNESBURG

 FPRR: ADCOCK INGRAM CRITICAL CARE,
 AEROTON, JOHANNESBURG

 Shelf-life: 24 months (Provisional)
 Date of registration: 15 AUGUST 2008

MRF 15

Registration number: 41/18/0300
Name of medicine: FOSRENOL 500 mg
Dosage form: TABLET
Active ingredients: EACH TABLET CONTAINS:
LANTHANUM CARBONATE HYDRATE
EQUIVALENT TO LANTHANUM 500,0 mg
Conditions of registration: 1, 2, 3, 4, 5, 6, 7, 8
Applicant: ADCOCK INGRAM CRITICAL CARE (PTY) LTD
Manufacturer: BCM LTD, NOTTINGHAM, NOTTINGHAMSHIRE,
UK
Packer: BCM LTD, NOTTINGHAM, NOTTINGHAMSHIRE,
UK
Laboratory: FPRC: BCM LTD, NOTTINGHAM, NOTTINGHAMSHIRE,
UK
READING SCIENTIFIC SERVICES LTD, READING,
BERKSHIRE, UK
SOUTHERN TESTING & RESEARCH, WILSON,
NORTH CAROLINA, USA
SSCI Inc, WEST LAFAYETTE, INDIANAPOLIS, USA
TEPNEL SCIENTIFIC SERVICES, EDINBURGH,
YORKSHIRE, UK
M&L LABORATORY SERVICES, ORMONDE,
JOHANNESBURG
FPRR: ADCOCK INGRAM CRITICAL CARE, AEROTON,
JOHANNESBURG
Shelf-life: 24 months (Provisional)
Date of registration: 15 AUGUST 2008

MRF 15

Registration number: 41/18/0301
Name of medicine: FOSRENOL 750 mg
Dosage form: TABLET
Active ingredients: EACH TABLET CONTAINS:
LANTHANUM CARBONATE HYDRATE
EQUIVALENT TO LANTHANUM 750,0 mg
Conditions of registration: 1, 2, 3, 4, 5, 6, 7, 8
Applicant: ADCOCK INGRAM CRITICAL CARE (PTY)
LTD
Manufacturer: BCM LTD, NOTTINGHAM,
NOTTINGHAMSHIRE, UK
Packer: BCM LTD, NOTTINGHAM,
NOTTINGHAMSHIRE, UK
Laboratory: FPRC: BCM LTD, NOTTINGHAM,
NOTTINGHAMSHIRE, UK
READING SCIENTIFIC SERVICES LTD,
READING, BERKSHIRE, UK
SOUTHERN TESTING & RESEARCH,
WILSON, NORTH CAROLINA, USA
SSCI Inc, WEST LAFAYETTE,
INDIANAPOLIS, USA
TEPNEL SCIENTIFIC SERVICES,
EDINBURGH, YORKSHIRE, UK
M&L LABORATORY SERVICES, ORMONDE,
JOHANNESBURG
FPRR: ADCOCK INGRAM CRITICAL CARE,
AEROTON, JOHANNESBURG
Shelf-life: 24 months (Provisional)
Date of registration: 15 AUGUST 2008

MRF 15

Registration number:	41/18/0302
Name of medicine:	FOSRENOL 1 000 mg
Dosage form:	TABLET
Active ingredients:	EACH TABLET CONTAINS: LANTHANUM CARBONATE HYDRATE EQUIVALENT TO LANTHANUM 1 000,0 mg
Conditions of registration:	1, 2, 3, 4, 5, 6, 7, 8
Applicant:	ADCOCK INGRAM CRITICAL CARE (PTY) LTD
Manufacturer:	BCM LTD, NOTTINGHAM, NOTTINGHAMSHIRE, UK
Packer:	BCM LTD, NOTTINGHAM, NOTTINGHAMSHIRE, UK
Laboratory:	FPRC: BCM LTD, NOTTINGHAM, NOTTINGHAMSHIRE, UK READING SCIENTIFIC SERVICES LTD, READING, BERKSHIRE, UK SOUTHERN TESTING & RESEARCH, WILSON, NORTH CAROLINA, USA SSCI Inc, WEST LAFAYETTE, INDIANAPOLIS, USA TEPNEL SCIENTIFIC SERVICES, EDINBURGH, YORKSHIRE, UK M&L LABORATORY SERVICES, ORMONDE, JOHANNESBURG FPRR: ADCOCK INGRAM CRITICAL CARE, AEROTON, JOHANNESBURG
Shelf-life:	24 months (Provisional)
Date of registration:	15 AUGUST 2008

MRF 15

Registration number:	41/1.2/0371
Name of medicine:	WELLBUTRIN XL 150
Dosage form:	TABLET
Active ingredients:	EACH TABLET CONTAINS: BUPROPION HYDROCHLORIDE 150,0 mg
Conditions of registration:	1, 2, 3, 4, 5, 6, 7
Applicant:	GLAXOSMITHKLINE SOUTH AFRICA (PTY) LTD
Manufacturer:	BIOVAIL CORPORATION, STEINBACH, CANADA
Packer:	BIOVAIL CORPORATION, STEINBACH, CANADA GLAXO WELLCOME GmbH & Co, BAD OLDESLOE, GERMANY GLAXOSMITHKLINE, EPPING, CAPE TOWN
Laboratory:	FPRC: BIOVAIL CORPORATION, STEINBACH, CANADA INOPHARM INC, MARKHAM, CANADA GLAXO WELLCOME GmbH & Co, BAD OLDESLOE, GERMANY FPRC/FPRR: GLAXOSMITHKLINE, EPPING, CAPE TOWN
Shelf-life:	18 months
Date of registration:	15 AUGUST 2008

MRF 15

Registration number: 41/1.2/0372
Name of medicine: WELLBUTRIN XL 300
Dosage form: TABLET
Active ingredients: EACH TABLET CONTAINS:
BUPROPION HYDROCHLORIDE 300,0 mg
Conditions of registration: 1, 2, 3, 4, 5, 6, 7
Applicant: GLAXOSMITHKLINE SOUTH AFRICA (PTY) LTD
Manufacturer: BIOVAIL CORPORATION, STEINBACH, CANADA
Packer: BIOVAIL CORPORATION, STEINBACH, CANADA
GLAXO WELLCOME GmbH & Co, BAD
OLDESLOE, GERMANY
GLAXOSMITHKLINE, EPPING, CAPE TOWN
Laboratory: FPRC: BIOVAIL CORPORATION, STEINBACH, CANADA
INOPHARM INC, MARKHAM, CANADA
GLAXO WELLCOME GmbH & Co, BAD
OLDESLOE, GERMANY
FPRR: GLAXOSMITHKLINE, EPPING, CAPE TOWN
Shelf-life: 18 months
Date of registration: 15 AUGUST 2008

MRF 15

Registration number: 41/20.1.1/0388
Name of medicine: AURO-CEFALEXIN TABLETS 500 mg
Dosage form: TABLET
Active ingredients: EACH TABLET CONTAINS:
CEFALEXIN MONOHYDRATE EQUIVALENT TO
CEFALEXIN 500,0 mg
Conditions of registration: 1, 2, 3, 4, 5, 6, 7, 8
Applicant: AUROBINDO PHARMA (PTY) LTD
Manufacturer: AUROBINDO PHARMA LTD, UNIT VI, MEDAK
DISTRICT, ANDHRA PRADESH, INDIA
Packer: AUROBINDO PHARMA LTD, UNIT VI, MEDAK
DISTRICT, ANDHRA PRADESH, INDIA
Laboratory: FPRC: AUROBINDO PHARMA LTD, UNIT VI, MEDAK
DISTRICT, ANDHRA PRADESH, INDIA
FPRR: AUROBINDO PHARMA, ROSEBANK,
JOHANNESBURG
Shelf-life: 24 months (Provisional)
Date of registration: 15 AUGUST 2008

MRF 15

Registration number: 41/20.1.1/0389
 Name of medicine: AURO-CEFALEXIN TABLETS 1 000 mg
 Dosage form: TABLET
 Active ingredients: EACH TABLET CONTAINS:
 CEFALEXIN MONOHYDRATE EQUIVALENT TO
 CEFALEXIN 1 000,0 mg
 Conditions of registration: 1, 2, 3, 4, 5, 6, 7, 8
 Applicant: AUROBINDO PHARMA (PTY) LTD
 Manufacturer: AUROBINDO PHARMA LTD, UNIT VI, MEDAK
 DISTRICT, ANDHRA PRADESH, INDIA
 Packer: AUROBINDO PHARMA LTD, UNIT VI, MEDAK
 DISTRICT, ANDHRA PRADESH, INDIA
 Laboratory: FPRC: AUROBINDO PHARMA LTD, UNIT VI, MEDAK
 DISTRICT, ANDHRA PRADESH, INDIA
 FPRC/FPRR: AUROBINDO PHARMA, ROSEBANK,
 JOHANNESBURG
 Shelf-life: 24 months (Provisional)
 Date of registration: 15 AUGUST 2008

MRF 15

Registration number: 41/7.5/0409
 Name of medicine: COLITE 10 mg TABLETS
 Dosage form: TABLET
 Active ingredients: EACH TABLET CONTAINS:
 PRAVASTATIN SODIUM 10,0 mg
 Conditions of registration: 1, 2, 3, 4, 5, 6, 7
 Applicant: ZYDUS HEALTHCARE S.A. (PTY) LTD
 Manufacturer: ZYDUS CADILA HEALTHCARE LTD,
 SANAND, AHMEDABAD, INDIA
 Packer: ZYDUS CADILA HEALTHCARE LTD,
 SANAND, AHMEDABAD, INDIA
 Laboratory: FPRC: ZYDUS CADILA HEALTHCARE LTD,
 SANAND, AHMEDABAD, INDIA
 INSTITUTE FOR PHARMACEUTICAL
 SERVICES, SILVERTONDALE, PRETORIA
 RESEARCH INSTITUTE FOR INDUSTRIAL
 PHARMACY, NORTH-WEST UNIVERSITY,
 POTCHEFSTROOM
 FPRR: ZYDUS HEALTHCARE, VAN DER HOFF
 PARK, POTCHEFSTROOM
 Shelf-life: 24 months
 Date of registration: 15 AUGUST 2008

MRF 15

Registration number: 41/7.5/0410
 Name of medicine: COLITE 20 mg TABLETS
 Dosage form: TABLET
 Active ingredients: EACH TABLET CONTAINS:
 PRAVASTATIN SODIUM 20,0 mg
 Conditions of registration: 1, 2, 3, 4, 5, 6, 7
 Applicant: ZYDUS HEALTHCARE S.A. (PTY) LTD
 Manufacturer: ZYDUS CADILA HEALTHCARE LTD, SANAND,
 AHMEDABAD, INDIA
 Packer: ZYDUS CADILA HEALTHCARE LTD, SANAND,
 AHMEDABAD, INDIA
 Laboratory: FPRC: ZYDUS CADILA HEALTHCARE LTD, SANAND,
 AHMEDABAD, INDIA
 INSTITUTE FOR PHARMACEUTICAL SERVICES,
 SILVERTONDALE, PRETORIA
 RESEARCH INSTITUTE FOR INDUSTRIAL
 PHARMACY, NORTH-WEST UNIVERSITY,
 POTCHEFSTROOM
 FPRR: ZYDUS HEALTHCARE, VAN DER HOFF PARK,
 POTCHEFSTROOM
 Shelf-life: 24 months
 Date of registration: 15 AUGUST 2008

MRF 15

Registration number: 41/7.5/0411
 Name of medicine: COLITE 40 mg TABLETS
 Dosage form: TABLET
 Active ingredients: EACH TABLET CONTAINS:
 PRAVASTATIN SODIUM 40,0 mg
 Conditions of registration: 1, 2, 3, 4, 5, 6, 7
 Applicant: ZYDUS HEALTHCARE S.A. (PTY) LTD
 Manufacturer: ZYDUS CADILA HEALTHCARE LTD, SANAND,
 AHMEDABAD, INDIA
 Packer: ZYDUS CADILA HEALTHCARE LTD, SANAND,
 AHMEDABAD, INDIA
 Laboratory: FPRC: ZYDUS CADILA HEALTHCARE LTD, SANAND,
 AHMEDABAD, INDIA
 INSTITUTE FOR PHARMACEUTICAL SERVICES,
 SILVERTONDALE, PRETORIA
 RESEARCH INSTITUTE FOR INDUSTRIAL
 PHARMACY, NORTH-WEST UNIVERSITY,
 POTCHEFSTROOM
 FPRR: ZYDUS HEALTHCARE, VAN DER HOFF PARK,
 POTCHEFSTROOM
 Shelf-life: 24 months
 Date of registration: 15 AUGUST 2008

MRF 15

Registration number:	41/7.5/0412
Name of medicine:	COLITE 80 mg TABLETS
Dosage form:	TABLET
Active ingredients:	EACH TABLET CONTAINS: PRAVASTATIN SODIUM 80,0 mg
Conditions of registration:	1, 2, 3, 4, 5, 6, 7
Applicant:	ZYDUS HEALTHCARE S.A. (PTY) LTD
Manufacturer:	ZYDUS CADILA HEALTHCARE LTD, SANAND, AHMEDABAD, INDIA
Packer:	ZYDUS CADILA HEALTHCARE LTD, SANAND, AHMEDABAD, INDIA
Laboratory:	FPRC: ZYDUS CADILA HEALTHCARE LTD, SANAND, AHMEDABAD, INDIA INSTITUTE FOR PHARMACEUTICAL SERVICES, SILVERTONDALE, PRETORIA RESEARCH INSTITUTE FOR INDUSTRIAL PHARMACY, NORTH-WEST UNIVERSITY, POTCHEFSTROOM
	FPRR: ZYDUS HEALTHCARE, VAN DER HOFF PARK, POTCHEFSTROOM
Shelf-life:	24 months
Date of registration:	15 AUGUST 2008

MRF 15

Registration number:	41/1.2/0440
Name of medicine:	ODIVEN 37,5
Dosage form:	TABLET
Active ingredients:	EACH TABLET CONTAINS: VENLAFAXINE HYDROCHLORIDE EQUIVALENT TO VENLAFAXINE 37,5 mg
Conditions of registration:	1, 2, 3, 4, 5, 6, 7, 8
Applicant:	ZYDUS HEALTHCARE SA (PTY) LTD
Manufacturer:	ZYDUS CADILA HEALTHCARE LTD, SANAND, AHMEDABAD, INDIA
Packer:	ZYDUS CADILA HEALTHCARE LTD, SANAND, AHMEDABAD, INDIA
Laboratory:	FPRC: ZYDUS CADILA HEALTHCARE LTD, SANAND, AHMEDABAD, INDIA INSTITUTE FOR PHARMACEUTICAL SERVICES, SILVERTONDALE, PRETORIA RESEARCH INSTITUTE FOR INDUSTRIAL PHARMACY, NORTH-WEST UNIVERSITY, POTCHEFSTROOM
	FPRR: ZYDUS HEALTHCARE, VAN DER HOFF PARK, POTCHEFSTROOM
Shelf-life:	24 months (Provisional)
Date of registration:	15 AUGUST 2008

MRF 15

Registration number: 41/1.2/0441
 Name of medicine: ODIVEN 50
 Dosage form: TABLET
 Active ingredients: EACH TABLET CONTAINS:
 VENLAFAXINE HYDROCHLORIDE EQUIVALENT
 TO VENLAFAXINE 50,0 mg
 Conditions of registration: 1, 2, 3, 4, 5, 6, 7, 8
 Applicant: ZYDUS HEALTHCARE SA (PTY) LTD
 Manufacturer: ZYDUS CADILA HEALTHCARE LTD, SANAND,
 AHMEDABAD, INDIA
 Packer: ZYDUS CADILA HEALTHCARE LTD, SANAND,
 AHMEDABAD, INDIA
 Laboratory: FPRC: ZYDUS CADILA HEALTHCARE LTD, SANAND,
 AHMEDABAD, INDIA
 INSTITUTE FOR PHARMACEUTICAL SERVICES,
 SILVERTONDALE, PRETORIA
 RESEARCH INSTITUTE FOR INDUSTRIAL
 PHARMACY, NORTH-WEST UNIVERSITY,
 POTCHEFSTROOM
 FPRR: ZYDUS HEALTHCARE, VAN DER HOFF PARK,
 POTCHEFSTROOM
 Shelf-life: 24 months (Provisional)
 Date of registration: 15 AUGUST 2008

MRF 15

Registration number: 41/1.2/0442
 Name of medicine: ODIVEN 75
 Dosage form: TABLET
 Active ingredients: EACH TABLET CONTAINS:
 VENLAFAXINE HYDROCHLORIDE
 EQUIVALENT TO VENLAFAXINE 75,0 mg
 Conditions of registration: 1, 2, 3, 4, 5, 6, 7, 8
 Applicant: ZYDUS HEALTHCARE SA (PTY) LTD
 Manufacturer: ZYDUS CADILA HEALTHCARE LTD,
 SANAND, AHMEDABAD, INDIA
 Packer: ZYDUS CADILA HEALTHCARE LTD,
 SANAND, AHMEDABAD, INDIA
 Laboratory: FPRC: ZYDUS CADILA HEALTHCARE LTD,
 SANAND, AHMEDABAD, INDIA
 INSTITUTE FOR PHARMACEUTICAL
 SERVICES, SILVERTONDALE, PRETORIA
 RESEARCH INSTITUTE FOR INDUSTRIAL
 PHARMACY, NORTH-WEST UNIVERSITY,
 POTCHEFSTROOM
 FPRR: ZYDUS HEALTHCARE, VAN DER HOFF
 PARK, POTCHEFSTROOM
 Shelf-life: 24 months (Provisional)
 Date of registration: 15 AUGUST 2008

MRF 15

Registration number: 41/20.1.2/0532
 Name of medicine: BINDOCLAV 375 mg
 Dosage form: TABLET
 Active ingredients: EACH TABLET CONTAINS:
 AMOXYCILLIN TRIHYDRATE EQUIVALENT TO
 AMOXYCILLIN 250,0 mg
 POTASSIUM CLAVULANATE EQUIVALENT TO
 CLAVULANIC ACID 125,0 mg
 Conditions of registration: 1, 2, 3, 4, 5, 6, 7, 8
 Applicant: AUROBINDO PHARMA (PTY) LTD
 Manufacturer: AUROBINDO PHARMA, UNIT XII,
 HYDERABAD, ANDHRA PRADESH, INDIA
 Packer: AUROBINDO PHARMA, UNIT XII,
 HYDERABAD, ANDHRA PRADESH, INDIA
 Laboratory: FPRC: AUROBINDO PHARMA, UNIT XII,
 HYDERABAD, ANDHRA PRADESH, INDIA
 FPRR: AUROBINDO PHARMA, ROSEBANK,
 JOHANNESBURG
 Shelf-life: 24 months (Provisional)
 Date of registration: 15 AUGUST 2008

MRF 15

Registration number: 41/20.1.2/0533
 Name of medicine: BINDOCLAV 625 mg
 Dosage form: TABLET
 Active ingredients: EACH TABLET CONTAINS:
 AMOXYCILLIN TRIHYDRATE EQUIVALENT TO
 AMOXYCILLIN 500,0 mg
 POTASSIUM CLAVULANATE EQUIVALENT TO
 CLAVULANIC ACID 125,0 mg
 Conditions of registration: 1, 2, 3, 4, 5, 6, 7, 8
 Applicant: AUROBINDO PHARMA (PTY) LTD
 Manufacturer: AUROBINDO PHARMA, UNIT XII, HYDERABAD,
 ANDHRA PRADESH, INDIA
 Packer: AUROBINDO PHARMA, UNIT XII, HYDERABAD,
 ANDHRA PRADESH, INDIA
 Laboratory: FPRC: AUROBINDO PHARMA, UNIT XII, HYDERABAD,
 ANDHRA PRADESH, INDIA
 FPRR: AUROBINDO PHARMA, ROSEBANK,
 JOHANNESBURG
 Shelf-life: 24 months (Provisional)
 Date of registration: 15 AUGUST 2008

MRF 15

Registration number:	41/20.1.2/0534
Name of medicine:	BINDOCLAV 1 000 mg
Dosage form:	TABLET
Active ingredients:	EACH TABLET CONTAINS: AMOXYCILLIN TRIHYDRATE EQUIVALENT TO AMOXYCILLIN 875,0 mg POTASSIUM CLAVULANATE EQUIVALENT TO CLAVULANIC ACID 125,0 mg
Conditions of registration:	1, 2, 3, 4, 5, 6, 7, 8
Applicant:	AUROBINDO PHARMA (PTY) LTD
Manufacturer:	AUROBINDO PHARMA, UNIT XII, HYDERABAD, ANDHRA PRADESH, INDIA
Packer:	AUROBINDO PHARMA, UNIT XII, HYDERABAD, ANDHRA PRADESH, INDIA
Laboratory:	FPRC: AUROBINDO PHARMA, UNIT XII, HYDERABAD, ANDHRA PRADESH, INDIA
	FPRR: AUROBINDO PHARMA, ROSEBANK, JOHANNESBURG
Shelf-life:	24 months (Provisional)
Date of registration:	15 AUGUST 2008

MRF 15

Registration number:	41/20.1.2/0535
Name of medicine:	AURO-AMOXICLAV 375 mg
Dosage form:	TABLET
Active ingredients:	EACH TABLET CONTAINS: AMOXYCILLIN TRIHYDRATE EQUIVALENT TO AMOXYCILLIN 250,0 mg POTASSIUM CLAVULANATE EQUIVALENT TO CLAVULANIC ACID 125,0 mg
Conditions of registration:	1, 2, 3, 4, 5, 6, 7, 8
Applicant:	AUROBINDO PHARMA (PTY) LTD
Manufacturer:	AUROBINDO PHARMA, UNIT XII, HYDERABAD, ANDHRA PRADESH, INDIA
Packer:	AUROBINDO PHARMA, UNIT XII, HYDERABAD, ANDHRA PRADESH, INDIA
Laboratory:	FPRC: AUROBINDO PHARMA, UNIT XII, HYDERABAD, ANDHRA PRADESH, INDIA
	FPRR: AUROBINDO PHARMA, ROSEBANK, JOHANNESBURG
Shelf-life:	24 months (Provisional)
Date of registration:	15 AUGUST 2008

MRF 15

Registration number: 41/20.1.2/0536
 Name of medicine: AURO-AMOXICLAV 625 mg
 Dosage form: TABLET
 Active ingredients: EACH TABLET CONTAINS:
 AMOXYCILLIN TRIHYDRATE EQUIVALENT TO 500,0 mg
 AMOXYCILLIN
 POTASSIUM CLAVULANATE EQUIVALENT TO 125,0 mg
 CLAVULANIC ACID

 Conditions of registration: 1, 2, 3, 4, 5, 6, 7, 8
 Applicant: AUROBINDO PHARMA (PTY) LTD
 Manufacturer: AUROBINDO PHARMA, UNIT XII, HYDERABAD,
 ANDHRA PRADESH, INDIA
 Packer: AUROBINDO PHARMA, UNIT XII, HYDERABAD,
 ANDHRA PRADESH, INDIA
 Laboratory: FPRC: AUROBINDO PHARMA, UNIT XII, HYDERABAD,
 ANDHRA PRADESH, INDIA

 FPRR: AUROBINDO PHARMA, ROSEBANK,
 JOHANNESBURG
 Shelf-life: 24 months (Provisional)
 Date of registration: 15 AUGUST 2008

MRF 15

Registration number: 41/20.1.2/0537
 Name of medicine: AURO-AMOXICLAV 1 000 mg
 Dosage form: TABLET
 Active ingredients: EACH TABLET CONTAINS:
 AMOXYCILLIN TRIHYDRATE
 EQUIVALENT TO 875,0 mg
 AMOXYCILLIN
 POTASSIUM CLAVULANATE
 EQUIVALENT TO 125,0 mg
 CLAVULANIC ACID

 Conditions of registration: 1, 2, 3, 4, 5, 6, 7, 8
 Applicant: AUROBINDO PHARMA (PTY) LTD
 Manufacturer: AUROBINDO PHARMA, UNIT XII,
 HYDERABAD, ANDHRA PRADESH, INDIA
 Packer: AUROBINDO PHARMA, UNIT XII,
 HYDERABAD, ANDHRA PRADESH, INDIA
 Laboratory: FPRC: AUROBINDO PHARMA, UNIT XII,
 HYDERABAD, ANDHRA PRADESH, INDIA

 FPRC/FPRR: AUROBINDO PHARMA, ROSEBANK,
 JOHANNESBURG
 Shelf-life: 24 months (Provisional)
 Date of registration: 15 AUGUST 2008

MRF 15

Registration number: 41/20.1.2/0546
 Name of medicine: BETACLAV 375
 Dosage form: TABLET
 Active ingredients: EACH TABLET CONTAINS:
 AMOXYCILLIN TRIHYDRATE EQUIVALENT TO
 AMOXYCILLIN 250,0 mg
 POTASSIUM CLAVULANATE EQUIVALENT TO
 CLAVULANIC ACID 125,0 mg
 Conditions of registration: 1, 2, 3, 4, 5, 6, 7
 Applicant: BE-TABS PHARMACEUTICALS (PTY) LTD
 Manufacturer: AUROBINDO PHARMA LTD, UNIT XII,
 HYDERABAD, ANDHRA PRADESH, INDIA
 Packer: AUROBINDO PHARMA LTD, UNIT XII,
 HYDERABAD, ANDHRA PRADESH, INDIA
 Laboratory: FPRC: AUROBINDO PHARMA LTD, UNIT XII,
 HYDERABAD, ANDHRA PRADESH, INDIA
 FPRC/FPRR: BE-TABS PHARMACEUTICALS, ROODEPOORT
 Shelf-life: 24 months
 Date of registration: 15 AUGUST 2008

MRF 15

Registration number: 41/20.1.2/0547
 Name of medicine: BETACLAV 625
 Dosage form: TABLET
 Active ingredients: EACH TABLET CONTAINS:
 AMOXYCILLIN TRIHYDRATE EQUIVALENT TO
 AMOXYCILLIN 500,0 mg
 POTASSIUM CLAVULANATE EQUIVALENT TO
 CLAVULANIC ACID 125,0 mg
 Conditions of registration: 1, 2, 3, 4, 5, 6, 7
 Applicant: BE-TABS PHARMACEUTICALS (PTY) LTD
 Manufacturer: AUROBINDO PHARMA LTD, UNIT XII,
 HYDERABAD, ANDHRA PRADESH, INDIA
 Packer: AUROBINDO PHARMA LTD, UNIT XII,
 HYDERABAD, ANDHRA PRADESH, INDIA
 Laboratory: FPRC: AUROBINDO PHARMA LTD, UNIT XII,
 HYDERABAD, ANDHRA PRADESH, INDIA
 FPRR: BE-TABS PHARMACEUTICALS, ROODEPOORT
 Shelf-life: 24 months
 Date of registration: 15 AUGUST 2008

MRF 15

Registration number: 41/20.1.2/0548
 Name of medicine: BETACLAV 1 000
 Dosage form: TABLET
 Active ingredients: EACH TABLET CONTAINS:
 AMOXYCILLIN TRIHYDRATE EQUIVALENT TO 875,0 mg
 AMOXYCILLIN 875,0 mg
 POTASSIUM CLAVULANATE EQUIVALENT TO 125,0 mg
 CLAVULANIC ACID 125,0 mg
 Conditions of registration: 1, 2, 3, 4, 5, 6, 7
 Applicant: BE-TABS PHARMACEUTICALS (PTY) LTD
 Manufacturer: AUROBINDO PHARMA LTD, UNIT XII,
 HYDERABAD, ANDHRA PRADESH, INDIA
 Packer: AUROBINDO PHARMA LTD, UNIT XII,
 HYDERABAD, ANDHRA PRADESH, INDIA
 Laboratory: FPRC: AUROBINDO PHARMA LTD, UNIT XII,
 HYDERABAD, ANDHRA PRADESH, INDIA
 FPRC/FPRR: BE-TABS PHARMACEUTICALS, ROODEPOORT
 Shelf-life: 24 months
 Date of registration: 15 AUGUST 2008

MRF 15

Registration number: 41/5.4.1/0604
 Name of medicine: REQUIP XL 2 mg
 Dosage form: TABLET
 Active ingredients: EACH TABLET CONTAINS:
 ROPINIROLE HYDROCHLORIDE
 EQUIVALENT TO 2,0 mg
 ROPINIROLE 2,0 mg
 Conditions of registration: 1, 2, 3, 4, 5, 6, 7
 Applicant: GLAXOSMITHKLINE SOUTH AFRICA
 (PTY) LTD
 Manufacturer: SMITHKLINE BEECHAM PLC, CRAWLEY,
 WEST SUSSEX, UK
 Packer: SMITHKLINE BEECHAM PLC, CRAWLEY,
 WEST SUSSEX, UK
 CARDINAL HEALTH LTD, GREAT
 OAKLEY, CORBY, NORTHAMPTONSHIRE,
 UK
 GLAXOSMITHKLINE, EPPING, CAPE
 TOWN
 Laboratory: FPRC: SMITHKLINE BEECHAM PLC, CRAWLEY,
 WEST SUSSEX, UK
 FPRC/FPRR: GLAXOSMITHKLINE, EPPING, CAPE
 TOWN
 Shelf-life: 36 months
 Date of registration: 15 AUGUST 2008

MRF 15

Registration number: 41/5.4.1/0605
Name of medicine: REQUIP XL 3 mg
Dosage form: TABLET
Active ingredients: EACH TABLET CONTAINS:
ROPINIROLE HYDROCHLORIDE EQUIVALENT
TO
ROPINIROLE 3,0 mg
Conditions of registration: 1, 2, 3, 4, 5, 6, 7
Applicant: GLAXOSMITHKLINE SOUTH AFRICA (PTY) LTD
Manufacturer: SMITHKLINE BEECHAM PLC, CRAWLEY, WEST
SUSSEX, UK
Packer: SMITHKLINE BEECHAM PLC, CRAWLEY, WEST
SUSSEX, UK
CARDINAL HEALTH LTD, GREAT OAKLEY,
CORBY, NORTHAMPTONSHIRE, UK
GLAXOSMITHKLINE, EPPING, CAPE TOWN
Laboratory: FPRC: SMITHKLINE BEECHAM PLC, CRAWLEY, WEST
SUSSEX, UK
FPRC/FPRR: GLAXOSMITHKLINE, EPPING, CAPE TOWN
Shelf-life: 36 months
Date of registration: 15 AUGUST 2008

MRF 15

Registration number: 41/5.4.1/0606
Name of medicine: REQUIP XL 4 mg
Dosage form: TABLET
Active ingredients: EACH TABLET CONTAINS:
ROPINIROLE HYDROCHLORIDE EQUIVALENT
TO
ROPINIROLE 4,0 mg
Conditions of registration: 1, 2, 3, 4, 5, 6, 7
Applicant: GLAXOSMITHKLINE SOUTH AFRICA (PTY)
LTD
Manufacturer: SMITHKLINE BEECHAM PLC, CRAWLEY,
WEST SUSSEX, UK
Packer: SMITHKLINE BEECHAM PLC, CRAWLEY,
WEST SUSSEX, UK
CARDINAL HEALTH LTD, GREAT OAKLEY,
CORBY, NORTHAMPTONSHIRE, UK
GLAXOSMITHKLINE, EPPING, CAPE TOWN
Laboratory: FPRC: SMITHKLINE BEECHAM PLC, CRAWLEY,
WEST SUSSEX, UK
FPRC/FPRR: GLAXOSMITHKLINE, EPPING, CAPE TOWN
Shelf-life: 36 months
Date of registration: 15 AUGUST 2008

MRF 15

Registration number: 41/5.4.1/0607
 Name of medicine: REQUIP XL 8 mg
 Dosage form: TABLET
 Active ingredients: EACH TABLET CONTAINS:
 ROPINIROLE HYDROCHLORIDE
 EQUIVALENT TO
 ROPINIROLE 8,0 mg
 Conditions of registration: 1, 2, 3, 4, 5, 6, 7
 Applicant: GLAXOSMITHKLINE SOUTH AFRICA (PTY) LTD
 Manufacturer: SMITHKLINE BEECHAM PLC, CRAWLEY, WEST SUSSEX, UK
 Packer: SMITHKLINE BEECHAM PLC, CRAWLEY, WEST SUSSEX, UK
 CARDINAL HEALTH LTD, GREAT OAKLEY, CORBY, NORTHAMPTONSHIRE, UK
 GLAXOSMITHKLINE, EPPING, CAPE TOWN
 Laboratory: FPRC: SMITHKLINE BEECHAM PLC, CRAWLEY, WEST SUSSEX, UK
 FPRC/FPRR: GLAXOSMITHKLINE, EPPING, CAPE TOWN
 Shelf-life: 36 months
 Date of registration: 15 AUGUST 2008

MRF 15

Registration number: 41/20.2.8/0737
 Name of medicine: ADCO-ZIDOVUDINE 100 mg
 Dosage form: CAPSULE
 Active ingredients: EACH CAPSULE CONTAINS:
 ZIDOVUDINE 100,0 mg
 Conditions of registration: 1, 2, 3, 4, 5, 6, 7, 8
 Applicant: ADCOCK INGRAM LIMITED
 Manufacturer: ADCOCK INGRAM HEALTHCARE, WADEVILLE, GERMISTON
 Packer: ADCOCK INGRAM HEALTHCARE, WADEVILLE, GERMISTON
 ADCOCK INGRAM LTD, AEROTON, JOHANNESBURG
 Laboratory: FPRC: ADCOCK INGRAM HEALTHCARE, WADEVILLE, GERMISTON
 FPRC/FPRR: ADCOCK INGRAM LTD, AEROTON, JOHANNESBURG
 FPRR: ADCOCK INGRAM LTD, BRYANSTON, JOHANNESBURG
 Shelf-life: 24 months (Provisional)
 Date of registration: 15 AUGUST 2008

MRF 15

Registration number: 41/20.2.8/0738
Name of medicine: ADCO-ZIDOVUDINE 250 mg
Dosage form: CAPSULE
Active ingredients: EACH CAPSULE CONTAINS:
ZIDOVUDINE 250,0 mg

Conditions of registration: 1, 2, 3, 4, 5, 6, 7, 8
Applicant: ADCOCK INGRAM LIMITED
Manufacturer: ADCOCK INGRAM HEALTHCARE, WADEVILLE,
GERMISTON
Packer: ADCOCK INGRAM HEALTHCARE, WADEVILLE,
GERMISTON
ADCOCK INGRAM LTD, AEROTON, JOHANNESBURG
Laboratory: FPRC: ADCOCK INGRAM HEALTHCARE, WADEVILLE,
GERMISTON

FPRC/FPRR: ADCOCK INGRAM LTD, AEROTON, JOHANNESBURG
FPRR: ADCOCK INGRAM LTD, BRYANSTON, JOHANNESBURG

Shelf-life: 24 months (Provisional)
Date of registration: 15 AUGUST 2008

MRF 15

Registration number: 41/8.1/0812
Name of medicine: GULF AMLODIPINE 5
Dosage form: TABLET
Active ingredients: EACH TABLET CONTAINS:
AMLODIPINE BESYLATE EQUIVALENT TO
AMLODIPINE 5,0 mg

Conditions of registration: 1, 2, 3, 4, 5, 6, 7, 8
Applicant: GULF DRUG COMPANY (PTY) LTD
Manufacturer: ALEMBIC LTD, TAL-HALOL, PANCHMAHALS,
GUJARAT, INDIA
Packer: ALEMBIC LTD, TAL-HALOL, PANCHMAHALS,
GUJARAT, INDIA
Laboratory: FPRC: ALEMBIC LTD, TAL-HALOL, PANCHMAHALS,
GUJARAT, INDIA
SOUTH AFRICAN BUREAU OF STANDARDS,
GROENKLOOF, PRETORIA
M&L LABORATORY SERVICES, ORMONDE,
JOHANNESBURG
CONSULTING CHEMICAL LABORATORIES,
ATLASVILLE, BOKSBURG
PHARMA-Q, INDUSTRIA WEST, JOHANNESBURG
INSTITUTE FOR PHARMACEUTICALS SERVICES,
SILVERTONDALE, PRETORIA
CONSULTING MICROBIOLOGICAL
LABORATORIES, MOREWILL, BOKSBURG

FPRR: GULF DRUG CO, MOUNT EDGECOMBE, KZN

Shelf-life: 24 months (Provisional)
Date of registration: 15 AUGUST 2008

MRF 15

Registration number: 41/8.1/0813
Name of medicine: GULF AMLODIPINE 10
Dosage form: TABLET
Active ingredients: EACH TABLET CONTAINS:
 AMLODIPINE BESYLATE EQUIVALENT TO
 AMLODIPINE 10,0 mg
Conditions of registration: 1, 2, 3, 4, 5, 6, 7, 8
Applicant: GULF DRUG COMPANY (PTY) LTD
Manufacturer: ALEMBIC LTD, TAL-HALOL, PANCHMAHALS,
 GUJARAT, INDIA
Packer: ALEMBIC LTD, TAL-HALOL, PANCHMAHALS, GUJARAT,
 INDIA
Laboratory: **FPRC:** ALEMBIC LTD, TAL-HALOL, PANCHMAHALS, GUJARAT,
 INDIA
 SOUTH AFRICAN BUREAU OF STANDARDS,
 GROENKLOOF, PRETORIA
 M&L LABORATORY SERVICES, ORMONDE,
 JOHANNESBURG
 CONSULTING CHEMICAL LABORATORIES,
 ATLASVILLE, BOKSBURG
 PHARMA-Q, INDUSTRIA WEST, JOHANNESBURG
 INSTITUTE FOR PHARMACEUTICALS SERVICES,
 SILVERTONDALE, PRETORIA
 CONSULTING MICROBIOLOGICAL LABORATORIES,
 MOREWILL, BOKSBURG
FPRR: GULF DRUG CO, MOUNT EDGEcombe, KZN
Shelf-life: 24 months (Provisional)
Date of registration: 15 AUGUST 2008

MRF 15

Registration number: 41/7.1.3/0820
Name of medicine: CIPLAZAR 25
Dosage form: TABLET
Active ingredients: EACH TABLET CONTAINS:
 LOSARTAN POTASSIUM EQUIVALENT TO
 LOSARTAN 25,0 mg
Conditions of registration: 1, 2, 3, 4, 5, 6, 7
Applicant: CIPLA MEDPRO (PTY) LTD
Manufacturer: CIPLA LTD, (UNIT III), SALCETTE, GOA,
 INDIA
Packer: CIPLA LTD, (UNIT III), SALCETTE, GOA,
 INDIA
Laboratory: **FPRC:** CIPLA LTD, (UNIT III), SALCETTE, GOA,
 INDIA
FPRR: CIPLA MEDPRO, ROSENPARK, BELLVILLE
Shelf-life: 24 months
Date of registration: 15 AUGUST 2008

MRF 15

Registration number: 41/7.1.3/0821
Name of medicine: CIPLAZAR 50
Dosage form: TABLET
Active ingredients: EACH TABLET CONTAINS:
LOSARTAN POTASSIUM EQUIVALENT TO
LOSARTAN 50,0 mg
Conditions of registration: 1, 2, 3, 4, 5, 6, 7
Applicant: CIPLA MEDPRO (PTY) LTD
Manufacturer: CIPLA LTD, (UNIT III), SALCETTE, GOA, INDIA
Packer: CIPLA LTD, (UNIT III), SALCETTE, GOA, INDIA
Laboratory: FPRC: CIPLA LTD, (UNIT III), SALCETTE, GOA, INDIA

FPRR: CIPLA MEDPRO, ROSENPARK, BELLVILLE
Shelf-life: 24 months
Date of registration: 15 AUGUST 2008

MRF 15

Registration number: 41/7.1.3/0822
Name of medicine: CIPLAZAR 100
Dosage form: TABLET
Active ingredients: EACH TABLET CONTAINS:
LOSARTAN POTASSIUM EQUIVALENT TO
LOSARTAN 100,0 mg
Conditions of registration: 1, 2, 3, 4, 5, 6, 7
Applicant: CIPLA MEDPRO (PTY) LTD
Manufacturer: CIPLA LTD, (UNIT III), SALCETTE, GOA,
INDIA
Packer: CIPLA LTD, (UNIT III), SALCETTE, GOA,
INDIA
Laboratory: FPRC: CIPLA LTD, (UNIT III), SALCETTE, GOA,
INDIA

FPRR: CIPLA MEDPRO, ROSENPARK, BELLVILLE
Shelf-life: 24 months
Date of registration: 15 AUGUST 2008

MRF 15

Registration number: 41/1.2/0843
 Name of medicine: CITRAZ 5
 Dosage form: TABLET
 Active ingredients: EACH TABLET CONTAINS:
 ESCITALOPRAM OXALATE EQUIVALENT TO
 ESCITALOPRAM 5,0 mg
 Conditions of registration: 1, 2, 3, 4, 5, 6, 7, 8
 Applicant: DR REDDY'S LABORATORIES (PTY) LTD
 Manufacturer: DR REDDY'S LABORATORIES LTD, RANGA
 REDDY DISTRICT, ANDHRA PRADESH, INDIA
 Packer: DR REDDY'S LABORATORIES LTD, RANGA
 REDDY DISTRICT, ANDHRA PRADESH, INDIA
 DRA PHARMACEUTICALS, CENTURION, RSA
 Laboratory: FPRC: DR REDDY'S LABORATORIES LTD, RANGA
 REDDY DISTRICT, ANDHRA PRADESH, INDIA
 INSTITUTE FOR PHARMACEUTICAL SERVICES,
 SILVERTONDALE, PRETORIA
 RESEARCH INSTITUTE FOR INDUSTRIAL
 PHARMACY, NORTH-WEST UNIVERSITY,
 POTCHEFSTROOM
 FPRR: DR REDDY'S LABORATORIES, MURRAYFIELD,
 PRETORIA
 Shelf-life: 24 months (Provisional)
 Date of registration: 15 AUGUST 2008

MRF 15

Registration number: 41/1.2/0844
 Name of medicine: CITRAZ 10
 Dosage form: TABLET
 Active ingredients: EACH TABLET CONTAINS:
 ESCITALOPRAM OXALATE EQUIVALENT TO
 ESCITALOPRAM 10,0 mg
 Conditions of registration: 1, 2, 3, 4, 5, 6, 7, 8
 Applicant: DR REDDY'S LABORATORIES (PTY) LTD
 Manufacturer: DR REDDY'S LABORATORIES LTD, RANGA
 REDDY DISTRICT, ANDHRA PRADESH,
 INDIA
 Packer: DR REDDY'S LABORATORIES LTD, RANGA
 REDDY DISTRICT, ANDHRA PRADESH,
 INDIA
 DRA PHARMACEUTICALS, CENTURION,
 RSA
 Laboratory: FPRC: DR REDDY'S LABORATORIES LTD, RANGA
 REDDY DISTRICT, ANDHRA PRADESH,
 INDIA
 INSTITUTE FOR PHARMACEUTICAL
 SERVICES, SILVERTONDALE, PRETORIA
 RESEARCH INSTITUTE FOR INDUSTRIAL
 PHARMACY, NORTH-WEST UNIVERSITY,
 POTCHEFSTROOM
 FPRR: DR REDDY'S LABORATORIES,
 MURRAYFIELD, PRETORIA
 Shelf-life: 24 months (Provisional)
 Date of registration: 15 AUGUST 2008

MRF 15

Registration number: 41/1.2/0845
Name of medicine: CITRAZ 20
Dosage form: TABLET
Active ingredients: EACH TABLET CONTAINS:
ESCITALOPRAM OXALATE EQUIVALENT TO
ESCITALOPRAM 20,0 mg
Conditions of registration: 1, 2, 3, 4, 5, 6, 7, 8
Applicant: DR REDDY'S LABORATORIES (PTY) LTD
Manufacturer: DR REDDY'S LABORATORIES LTD, RANGA
REDDY DISTRICT, ANDHRA PRADESH, INDIA
Packer: DR REDDY'S LABORATORIES LTD, RANGA
REDDY DISTRICT, ANDHRA PRADESH, INDIA
DRA PHARMACEUTICALS, CENTURION, RSA
Laboratory: FPRC: DR REDDY'S LABORATORIES LTD, RANGA
REDDY DISTRICT, ANDHRA PRADESH, INDIA
INSTITUTE FOR PHARMACEUTICAL SERVICES,
SILVERTONDALE, PRETORIA
RESEARCH INSTITUTE FOR INDUSTRIAL
PHARMACY, NORTH-WEST UNIVERSITY,
POTCHEFSTROOM
FPRR: DR REDDY'S LABORATORIES, MURRAYFIELD,
PRETORIA
Shelf-life: 24 months (Provisional)
Date of registration: 15 AUGUST 2008

MRF 15

Registration number: 41/7.1/0935
Name of medicine: ZANERIL 10/10
Dosage form: TABLET
Active ingredients: EACH TABLET CONTAINS:
LERCANIDIPINE HYDROCHLORIDE 10,0 mg
ENALAPRIL MALEATE 10,0 mg
Conditions of registration: 1, 2, 3, 4, 5, 6, 7
Applicant: PHARMAPLAN (PTY) LTD
Manufacturer: RECORDATI INDUSTRIA CHIMICA &
FARMACEUTICA S.p.A., MILAN, ITALY
Packer: RECORDATI INDUSTRIA CHIMICA &
FARMACEUTICA S.p.A., MILAN, ITALY
Laboratory: FPRC: RECORDATI INDUSTRIA CHIMICA &
FARMACEUTICA S.p.A., MILAN, ITALY
CONSULTING CHEMICAL LABORATORIES,
ATLASVILLE, BOKSBURG
FPRR: PHARMAPLAN, MIDRAND, JOHANNESBURG
Shelf-life: 24 months
Date of registration: 15 AUGUST 2008

MRF 15

Registration number: 41/7.1/0936
 Name of medicine: ZANERIL 10/20
 Dosage form: TABLET
 Active ingredients: EACH TABLET CONTAINS:
 LERCANIDIPINE HYDROCHLORIDE 10,0 mg
 ENALAPRIL MALEATE 20,0 mg
 Conditions of registration: 1, 2, 3, 4, 5, 6, 7
 Applicant: PHARMAPLAN (PTY) LTD
 Manufacturer: RECORDATI INDUSTRIA CHIMICA &
 FARMACEUTICA S.p.A., MILAN, ITALY
 Packer: RECORDATI INDUSTRIA CHIMICA &
 FARMACEUTICA S.p.A., MILAN, ITALY
 Laboratory: FPRC: RECORDATI INDUSTRIA CHIMICA &
 FARMACEUTICA S.p.A., MILAN, ITALY
 CONSULTING CHEMICAL LABORATORIES,
 ATLASVILLE, BOKSBURG
 FPRR: PHARMAPLAN, MIDRAND, JOHANNESBURG
 Shelf-life: 24 months
 Date of registration: 15 AUGUST 2008

MRF 15

Registration number: 41/1.2/1007
 Name of medicine: VENELA 37,5
 Dosage form: TABLET
 Active ingredients: EACH TABLET CONTAINS:
 VENLAFAXINE HYDROCHLORIDE
 EQUIVALENT TO VENLAFAXINE 37,5 mg
 Conditions of registration: 1, 2, 3, 4, 5, 6, 7, 8
 Applicant: DR REDDY'S LABORATORIES (PTY) LTD
 Manufacturer: DR REDDY'S LABORATORIES LTD, RANGA
 REDDY DISTRICT, ANDHRA PRADESH,
 INDIA
 Packer: DR REDDY'S LABORATORIES LTD, RANGA
 REDDY DISTRICT, ANDHRA PRADESH,
 INDIA
 Laboratory: FPRC: DR REDDY'S LABORATORIES LTD, RANGA
 REDDY DISTRICT, ANDHRA PRADESH,
 INDIA
 INSTITUTE FOR PHARMACEUTICALS
 SERVICES, SILVERTONDALE, PRETORIA
 RESEARCH INSTITUTE FOR INDUSTRIAL
 PHARMACY, NORTH-WEST UNIVERSITY,
 POTCHEFSTROOM
 FPRR: DR REDDY'S LABORATORIES,
 MURRAYFIELD, PRETORIA
 Shelf-life: 24 months (Provisional)
 Date of registration: 15 AUGUST 2008

MRF 15

Registration number: 41/1.2/1008
Name of medicine: VENELA 50
Dosage form: TABLET
Active ingredients: EACH TABLET CONTAINS:
VENLAFAXINE HYDROCHLORIDE
EQUIVALENT TO VENLAFAXINE 50 mg
Conditions of registration: 1, 2, 3, 4, 5, 6, 7, 8
Applicant: DR REDDY'S LABORATORIES (PTY) LTD
Manufacturer: DR REDDY'S LABORATORIES LTD, RANGA
PEDDY DISTRICT, ANDHRA PRADESH, INDIA
Packer: DR REDDY'S LABORATORIES LTD, RANGA
REDDY DISTRICT, ANDHRA PRADESH, INDIA
Laboratory: FPRC: DR REDDY'S LABORATORIES LTD, RANGA
REDDY DISTRICT, ANDHRA PRADESH, INDIA
INSTITUTE FOR PHARMACEUTICALS
SERVICES, SILVERTONDALE, PRETORIA
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PHARMACY, NORTH-WEST UNIVERSITY,
POTCHEFSTROOM
FPRR: DR REDDY'S LABORATORIES,
MURRAYFIELD, PRETORIA
Shelf-life: 24 months (Provisional)
Date of registration: 15 AUGUST 2008

MRF 15

Registration number: 41/1.2/1009
Name of medicine: VENELA 75
Dosage form: TABLET
Active ingredients: EACH TABLET CONTAINS:
VENLAFAXINE HYDROCHLORIDE
EQUIVALENT TO VENLAFAXINE 75 mg
Conditions of registration: 1, 2, 3, 4, 5, 6, 7, 8
Applicant: DR REDDY'S LABORATORIES (PTY) LTD
Manufacturer: DR REDDY'S LABORATORIES LTD, RANGA
REDDY DISTRICT, ANDHRA PRADESH,
INDIA
Packer: DR REDDY'S LABORATORIES LTD, RANGA
REDDY DISTRICT, ANDHRA PRADESH,
INDIA
Laboratory: FPRC: DR REDDY'S LABORATORIES LTD, RANGA
REDDY DISTRICT, ANDHRA PRADESH,
INDIA
INSTITUTE FOR PHARMACEUTICALS
SERVICES, SILVERTONDALE, PRETORIA
RESEARCH INSTITUTE FOR INDUSTRIAL
PHARMACY, NORTH-WEST UNIVERSITY,
POTCHEFSTROOM
FPRR: DR REDDY'S LABORATORIES,
MURRAYFIELD, PRETORIA
Shelf-life: 24 months (Provisional)
Date of registration: 15 AUGUST 2008

MRF 15

Registration number: 41/21.2/1089
 Name of medicine: METORED 500
 Dosage form: TABLET
 Active ingredients: EACH TABLET CONTAINS:
 METFORMIN HYDROCHLORIDE 500,0 mg
 Conditions of registration: 1, 2, 3, 4, 5, 6, 7, 8
 Applicant: PHARMAPLAN (PTY) LTD
 Manufacturer: SUN PHARMACEUTICAL INDUSTRIES LTD,
 HALOL, DISTRICT PANCHMAHAL, GUJARAT,
 INDIA
 Packer: SUN PHARMACEUTICAL INDUSTRIES LTD,
 HALOL, DISTRICT PANCHMAHAL, GUJARAT,
 INDIA
 Laboratory: FPRC: SUN PHARMACEUTICAL INDUSTRIES LTD,
 HALOL, DISTRICT PANCHMAHAL, GUJARAT,
 INDIA
 CONSULTING CHEMICAL LABORATORIES,
 ATLASVILLE, BOKSBURG
 FPRR: PHARMAPLAN, MIDRAND, RSA
 Shelf-life: 24 months (Provisional)
 Date of registration: 15 AUGUST 2008

MRF 15

Registration number: 41/21.2/1090
 Name of medicine: METORED 850
 Dosage form: TABLET
 Active ingredients: EACH TABLET CONTAINS:
 METFORMIN HYDROCHLORIDE 850,0 mg
 Conditions of registration: 1, 2, 3, 4, 5, 6, 7, 8
 Applicant: PHARMAPLAN (PTY) LTD
 Manufacturer: SUN PHARMACEUTICAL INDUSTRIES LTD,
 HALOL, DISTRICT PANCHMAHAL,
 GUJARAT, INDIA
 Packer: SUN PHARMACEUTICAL INDUSTRIES LTD,
 HALOL, DISTRICT PANCHMAHAL,
 GUJARAT, INDIA
 Laboratory: FPRC: SUN PHARMACEUTICAL INDUSTRIES LTD,
 HALOL, DISTRICT PANCHMAHAL,
 GUJARAT, INDIA
 CONSULTING CHEMICAL LABORATORIES,
 ATLASVILLE, BOKSBURG
 FPRR: PHARMAPLAN, MIDRAND, RSA
 Shelf-life: 24 months (Provisional)
 Date of registration: 15 AUGUST 2008

MRF 15

Registration number: 41/21.2/1091

Name of medicine: METORED 1 000

Dosage form: TABLET

Active ingredients: EACH TABLET CONTAINS:
METFORMIN HYDROCHLORIDE 1 000,0 mg

Conditions of registration: 1, 2, 3, 4, 5, 6, 7, 8

Applicant: PHARMAPLAN (PTY) LTD

Manufacturer: SUN PHARMACEUTICAL INDUSTRIES LTD,
HALOL, DISTRICT PANCHMAHAL, GUJARAT,
INDIA

Packer: SUN PHARMACEUTICAL INDUSTRIES LTD,
HALOL, DISTRICT PANCHMAHAL, GUJARAT,
INDIA

Laboratory: FPRC: SUN PHARMACEUTICAL INDUSTRIES LTD,
HALOL, DISTRICT PANCHMAHAL, GUJARAT,
INDIA
CONSULTING CHEMICAL LABORATORIES,
ATLASVILLE, BOKSBURG

FPRR: PHARMAPLAN, MIDRAND, RSA

Shelf-life: 24 months (Provisional)

Date of registration: 15 AUGUST 2008

MRF 15

Registration number: 34/30.2/0501

Name of medicine: OCTAGAM

Dosage form: INFUSION

Active ingredients: EACH 1,0 ml SOLUTION CONTAINS:
HUMAN NORMAL IMMUNOGLOBULIN G
50,0 mg

Conditions of registration: 1, 2, 3, 4, 5, 6, 7

Applicant: JE ECKARD

Manufacturer: OCTAPHARMA PHARMAZEUTIKA, VIENNA,
AUSTRIA

Packer: OCTAPHARMA PHARMAZEUTIKA, VIENNA,
AUSTRIA

Laboratory: FPRC: OCTAPHARMA PHARMAZEUTIKA, VIENNA,
AUSTRIA

FPRR: JE ECKARD, MONUMENT PARK, PRETORIA,
RSA

Shelf-life: 24 months

Date of registration: 13 JUNE 2008