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## GENERAL NOTICE ALGEMENE KENNISGEWING

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### NOTICE 77 OF 2007

#### 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 No. 101 OF 1965)

1. The applicant shall ensure that the medicine is manufactured and controlled in terms of 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 No. 101 of 1965).
5. The registration of this medicine shall be subject to regular review 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 registration dossier is subject to review at intervals as determined by Council.
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 77 VAN 2007****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 NO. 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 26 van die Wet, om die nakoming van Goeie Vervaardigingspraktyke te bepaal.
3. Die inligting soos vervat in die 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 No. 101 van 1965).
5. Die registrasie van hierdie medisyne is onderhewig aan gereelde hersiening rakende kwaliteit, veiligheid en effektiwiteit, en die registrasie van hierdie medisyne kan gewysig word onderhewig aan kwessies soos goedgevind deur 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 registrasie-aansoek is onderhewig aan hersiening met tussenposes soos deur die Raad bepaal.
8. 'n Na-registrasie-inspeksie moet op die eerste produksielot van die plaaslike 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 in 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: 29/34/0761  
Name of medicine: AIR LIQUIDE MEDICAL NITROUS OXIDE  
Dosage form: GAS  
Active ingredients: EACH CYLINDER CONTAINS:  
NITROUS OXIDE 100,0 %  
Conditions of registration: 1, 2, 3, 4, 5, 6  
Applicant: AIR LIQUIDE (PTY) LTD  
Manufacturer: AIR LIQUIDE, ALRODE, GERMISTON  
Packer: AIR LIQUIDE, ALRODE, GERMISTON  
Laboratory: FPRC/FPRR: AIR LIQUIDE, ALRODE, GERMISTON  
Shelf-life: 24 months  
Date of registration: 1 DECEMBER 2006

**MRF 15**

Registration number: 32/7.1/0428  
Name of medicine: DILTIAZEM HEXAL 90  
Dosage form: TABLETS  
Active ingredients: EACH TABLET CONTAINS:  
DILTIAZEM HYDROCHLORIDE 90,0 mg  
Conditions of registration: 1, 2, 3, 4, 5, 6  
Applicant: HEXAL PHARMA (SA) (PTY) LTD  
Manufacturer: HEXAL AG, HOLZKIRCHEN, GERMANY  
Packer: HEXAL AG, HOLZKIRCHEN, GERMANY  
Laboratory: FPRC: HEXAL AG, HOLZKIRCHEN, GERMANY  
CONSULTING CHEMICAL  
LABORATORIES, ATLASVILLE, BOKSBURG, RSA  
FPRR: HEXAL PHARMA, PINETOWN, RSA  
Shelf-life: 24 months (provisional)  
Date of registration: 1 DECEMBER 2006

MRF 15

Registration number: 34/16.1/0366  
Name of medicine: ILIADIN 0,01 %  
Dosage form: DROPS  
Active ingredients: EACH 1,0 ml SOLUTION CONTAINS:  
OXYMETAZOLINE HYDROCHLORIDE 0,1 mg

Conditions of registration: 1, 2, 3, 4, 5, 6  
Applicant: MERCK (PTY) LTD  
Manufacturer: MERCK PHARMACEUTICAL MANUFACTURING,  
WADEVILLE, GERMISTON  
Packer: MERCK PHARMACEUTICAL MANUFACTURING,  
WADEVILLE, GERMISTON  
Laboratory: FPRC: MERCK PHARMACEUTICAL MANUFACTURING,  
WADEVILLE, GERMISTON  
FPRR: MERCK, MODDERFONTEIN, RSA  
Shelf-life: 36 months  
Date of registration 1 DECEMBER 2006

MRF 15

Registration number: 36/2.7/0183  
Name of medicine: EXCEDRIN  
Dosage form: TABLET  
Active ingredients: EACH TABLET CONTAINS:  
PARACETAMOL 250,0 mg  
ASPIRIN 250,0 mg  
CAFFEINE 65,0 mg

Conditions of registration: 1, 2, 3, 4, 5, 6  
Applicant: BRISTOL-MYERS SQUIBB (PTY) LTD  
Manufacturer: BRISTOL-MYERS SQUIBB, MOUNT VERNON, INDIANA,  
USA  
Packer: BRISTOL-MYERS SQUIBB, MOUNT VERNON, INDIANA,  
USA  
MERCK PHARMACEUTICAL MANUFACTURING,  
WADEVILLE, GERMISTON  
DIVPHARM MANUFACTURING & PACKAGING,  
LONGDALE, INDUSTRIA  
Laboratory: FPRC: BRISTOL-MYERS SQUIBB, MOUNT VERNON, INDIANA,  
USA  
MERCK PHARMACEUTICAL MANUFACTURING,  
WADEVILLE, GERMISTON  
CONSULTING CHEMICAL LABORATORIES, ATLASVILLE  
BOKSBURG, RSA  
FPRR: BRISTOL-MYERS SQUIBB, BEDFORDVIEW, RSA  
Shelf-life: 24 months  
Date of registration: 1 DECEMBER 2006

## MRF 15

Registration number: 37/30.1/0221  
 Name of medicine: TRITANRIX-HB + HIBERIX COMBO PACK  
 Dosage form: INJECTION  
 Active ingredients: EACH COMBO PACK CONTAINS:  
 TRITANRIX-HB INJECTION CONTAINING PER  
 0,5 ml DOSE:  
 DIPHTHERIA TOXOID 30,0 I.U.  
 TETANUS TOXOID 60,0 I.U.  
 INACTIVATED BORDETELLA PERTUSSIS  
 4,0 I.U.  
 HEPATITIS B VIRUS SURFACE ANTIGEN  
 10,0 ug  
 HIBERIX VACCINE CONTAINING PER 0,5 ml DOSE:  
 HAEMOPHILUS INFLUENZA TYPE b 10,0 ug

Conditions of registration: 1, 2, 3, 4, 5, 6  
 Applicant: GLAXOSMITHKLINE SOUTH AFRICA (PTY) LTD  
 Manufacturer: GLAXOSMITHKLINE BIOLOGICALS s.a, RIXENSART,  
 BELGIUM  
 SACHSISCHES SERUMWERK DRESDEN,  
 DRESDEN, GERMANY  
 Packer: GLAXOSMITHKLINE BIOLOGICALS  
 MANUFACTURING s.a, RIXENSART, BELGIUM  
 SACHSISCHES SERUMWERK DRESDEN,  
 DRESDEN, GERMANY  
 GLAXOSMITHKLINE BIOLOGICALS  
 MANUFACTURING s.a, WAVRE, BELGIUM  
 GLAXOSMITHKLINE S.A., EPPING INDUSTRIA  
 Laboratory: FPRC: GLAXOSMITHKLINE BIOLOGICALS s.a,  
 RIXENSART, BELGIUM  
 FPRC/FPRR GLAXOSMITHKLINE S.A., EPPING INDUSTRIA  
 Shelf-life: 36 months  
 Date of registration: 1 DECEMBER 2006

## MRF 15

Registration number: 37/26/0538  
 Name of medicine: EPIRUBICIN-LEMERY 10 mg  
 Dosage form: INJECTION  
 Active ingredients: EACH VIAL CONTAINS:  
 EPIRUBICIN HYDROCHLORIDE 10,0 mg  
 Conditions of registration: 1, 2, 3, 4, 5, 6  
 Applicant: KEY ONCOLOGICS (PTY) LTD  
 Manufacturer: LEMERY, S.A. de C.V., HUICHAPAN XOCH, MEXICO  
 Packer: LEMERY, S.A. de C.V., HUICHAPAN XOCH, MEXICO  
 Laboratory: FPRC: LEMERY, S.A. de C.V., HUICHAPAN XOCH, MEXICO  
 INSTITUTE FOR PHARMACEUTICAL SERVICES,  
 SILVERTONDALE, RSA  
 FPRR: KEY ONCOLOGICS, SANDTON, RSA  
 Shelf-life: 24 months  
 Date of registration: 1 DECEMBER 2006

MRF 15

Registration number: 37/26/0539  
Name of medicine: EPIRUBICIN-LEMERY 50 mg  
Dosage form: INJECTION  
Active ingredients: EACH VIAL CONTAINS:  
EPIRUBICIN HYDROCHLORIDE 50,0 mg  
Conditions of registration: 1, 2, 3, 4, 5, 6  
Applicant: KEY ONCOLOGICS (PTY) LTD  
Manufacturer: LEMERY, S.A. de C.V., HUICHAPAN XOCH, MEXICO  
Packer: LEMERY, S.A. de C.V., HUICHAPAN XOCH, MEXICO  
Laboratory: FPRC: LEMERY, S.A. de C.V., HUICHAPAN XOCH, MEXICO  
INSTITUTE FOR PHARMACEUTICAL SERVICES,  
SILVERTONDALE, RSA  
FPRR: KEY ONCOLOGICS, SANDTON, RSA  
Shelf-life: 24 months  
Date of registration: 1 DECEMBER 2006

MRF 15

Registration number: 37/11.1/0629  
Name of medicine: ASACOL 800  
Dosage form: TABLET  
Active ingredients: EACH TABLET CONTAINS:  
MESALAZINE 800,0 mg  
Conditions of registration: 1, 2, 3, 4, 5, 6  
Applicant: AVENTIS PHARMA (PTY) LTD  
Manufacturer: WULFING PHARMA (GmbH), GRONAU/LEINE,  
GERMANY  
Packer: AVENTIS PHARMA, WALTLOO, PRETORIA  
Laboratory: FPRC: WULFING PHARMA (GmbH), GRONAU/LEINE, GERMANY  
TILLOTS PHARMA AG, ZIEFEN, SWITZERLAND  
FPRC/FPRR: AVENTIS PHARMA, WALTLOO, PRETORIA  
Shelf-life: 24 months  
Date of registration: 1 DECEMBER 2006



## MRF 15

Registration number:	38/2.6.5/0030
Name of medicine	ZYPREXA VELOTAB 5 mg
Dosage form	TABLET
Active ingredients:	EACH TABLET CONTAINS: OLANZAPINE 5,0 mg
Conditions of registration:	1, 2, 3, 4, 5, 6
Applicant:	ELI LILLY (S.A.) (PTY) LTD
Manufacturer:	CARDINAL HEALTH, SWINDON, WILTSHIRE, UK
Packer:	ELI LILLY & CO, BASINGSTOKE, HAMPSHIRE UK PCI SERVICES, SHOTGATE, ESSEX, UK PCI SERVICES, WESTHOUGHTON, BOLTON, UK PCI SERVICES, CORBY, NORTHAMPTONSHIRE, UK
Laboratory:	FPRC: ELI LILLY & CO, BASINGSTOKE, HAMPSHIRE UK SOUTH AFRICAN BUREAU OF STANDARDS, GROENKLOOF, PRETORIA
	FPRR: ELI LILLY, BRYANSTON, RSA
Shelf-life:	24 months
Date of registration:	1 DECEMBER 2006

## MRF 15

Registration number:	38/11.10/0070
Name of medicine:	MOASON 500 SUPPOSITORIES
Dosage form:	SUPPOSITORY
Active ingredients:	EACH SUPPOSITORY CONTAINS: MESALAZINE 500,0 mg
Conditions of registration:	1, 2, 3, 4, 5, 6
Applicant:	ASTELLAS PHARMA (PTY) LTD
Manufacturer:	AMCAPHARM GmbH, ROSBACH, GERMANY ASTELLAS PHARMA SpA, MILAN, ITALY
Packer:	AMCAPHARM GmbH, ROSBACH, GERMANY ASTELLAS PHARMA SpA, MILAN, ITALY
Laboratory:	FPRC: AMCAPHARM GmbH, ROSBACH, GERMANY ASTELLAS PHARMA SpA, MILAN, ITALY CONSULTING CHEMICAL LABORATORIES, ATLASVILLE, BOKSBURG
	FPRR: ASTELLAS PHARMA, BEDFORDVIEW, RSA
Shelf-life:	24 months (provisional)
Date of registration	1 DECEMBER 2006

**MRF 15**

Registration number: 38/11.10/0071  
Name of medicine: MOASON 250 SUPPOSITORIES  
Dosage form: SUPPOSITORY  
Active ingredients: EACH SUPPOSITORY CONTAINS:  
MESALAZINE 250,0 mg  
Conditions of registration: 1, 2, 3, 4, 5, 6  
Applicant: ASTELLAS PHARMA (PTY) LTD  
Manufacturer: AMCAPHARM GmbH, ROSBACH, GERMANY  
ASTELLAS PHARMA SpA, MILAN, ITALY  
Packer: AMCAPHARM GmbH, ROSBACH, GERMANY  
ASTELLAS PHARMA SpA, MILAN, ITALY  
Laboratory: FPRC: AMCAPHARM GmbH, ROSBACH, GERMANY  
ASTELLAS PHARMA SpA, MILAN, ITALY  
CONSULTING CHEMICAL LABORATORIES,  
ATLASVILLE, BOKSBURG  
FPRR: ASTELLAS PHARMA, BEDFORDVIEW, RSA  
Shelf-life: 24 months (provisional)  
Date of registration: 1 DECEMBER 2006

**MRF 15**

Registration number: 38/2.6.5/0073  
Name of medicine: ZYPREXA VELOTAB 10 mg  
Dosage form: TABLET  
Active ingredients: EACH TABLET CONTAINS:  
OLANZAPINE 10,0 mg  
Conditions of registration: 1, 2, 3, 4, 5, 6  
Applicant: ELI LILLY (S.A.) (PTY) LTD  
Manufacturer: CARDINAL HEALTH, SWINDON, WILTSHIRE, UK  
Packer: ELI LILLY & CO, BASINGSTOKE, HAMPSHIRE UK  
PCI SERVICES, SHOTGATE, ESSEX, UK  
PCI SERVICES, WESTHOUGHTON, BOLTON, UK  
PCI SERVICES, CORBY, NORTHAMPTONSHIRE, UK  
Laboratory: FPRC: ELI LILLY & CO, BASINGSTOKE, HAMPSHIRE UK  
SOUTH AFRICAN BUREAU OF STANDARDS,  
GROENKLOOF, PRETORIA  
FPRR: ELI LILLY, BRYANSTON, RSA  
Shelf-life: 24 months  
Date of registration: 1 DECEMBER 2006

## MRF 15

Registration number: 38/13.1/0194  
 Name of medicine: POVIGEL  
 Dosage form: GEL  
 Active ingredients: EACH 1,0 g GEL CONTAINS:  
 POVIDONE IODINE 100,0 mg  
  
 Conditions of registration 1, 2, 3, 4, 5, 6  
 Applicant: MEDICINE DEVELOPERS INTERNATIONAL cc  
 Manufacturer: IMPILO DRUGS (1966), ISITHEBE, KZN, RSA  
 Packer: IMPILO DRUGS (1966), ISITHEBE, KZN, RSA  
 Laboratory: FPRC  
 IMPILO DRUGS (1966), ISITHEBE, KZN, RSA  
 CONSULTING CHEMICAL LABORATORIES,  
 ATLASVILLE, BOKSBURG  
 CONSULTING MICROBIOLOGICAL LABORATORIES,  
 MOREHILL, BENONI  
 SOUTH AFRICAN BUREAU OF STANDARDS,  
 GROENKLOOF, PRETORIA  
 INSTITUTE FOR PHARMACEUTICALS SERVICES,  
 SILVERTONDALE, RSA  
  
 FPRR: MDI cc, MENLO PARK, PRETORIA  
 Shelf-life: 24 months (provisional)  
 Date of registration: 1 DECEMBER 2006

## MRF 15

Registration number: 38/30.3/0281  
 Name of medicine: IMMUNINE 200 I.U  
 Dosage form: INJECTION  
 Active ingredients: EACH VIAL CONTAINS:  
 HUMAN COAGULATION FACTOR IX 200,0 I.U  
  
 Conditions of registration: 1, 2, 3, 4, 5, 6  
 Applicant: ADCOCK INGRAM CRITICAL CARE LTD  
 Manufacturer: BAXTER AG, INDUSTRIESTRASSE, VIENNA, AUSTRIA  
 BAXTER AG, BENATZKYGASSE,  
 VIENNA, AUSTRIA  
 BAXTER HEALTHCARE CORP., ROCHESTER,  
 MI, USA  
 BAXTER S.p.A., RIETI, RUFINA, ITALY  
 BAXTER AG, LANGE ALLEE 24-B, VIENNA,  
 AUSTRIA  
  
 Packer: ADCOCK INGRAM CRITICAL CARE, AEROTON,  
 JOHANNESBURG  
  
 Laboratory: FPRC: BAXTER AG, LANGE ALLEE 24, A-1220, VIENNA,  
 AUSTRIA  
 BAXTER AG, LANG ALLEE 24, A-1220, VIENNA,  
 AUSTRIA  
 BAXTER AG, INDUSTRIESTRASSE, VIENNA,  
 AUSTRIA  
 BAXTER AG, BENATZKYGASSE, VIENNA, AUSTRIA  
 BAXTER AG, ORTH/DONAU, AUSTRIA  
  
 FPRC/FPRR: ADCOCK INGRAM CRITICAL CARE, AEROTON,  
 JOHANNESBURG  
  
 Shelf-life: 36 months  
 Date of registration: 1 DECEMBER 2006

## MRF 15

Registration number: 38/30.3/0282  
 Name of medicine: IMMUNINE 600 I.U.  
 Dosage form: INJECTION  
 Active ingredients: EACH VIAL CONTAINS:  
 HUMAN COAGULATION FACTOR IX 600,0 I.U.  
 Conditions of registration: 1, 2, 3, 4, 5, 6  
 Applicant: ADCOCK INGRAM CRITICAL CARE LTD  
 Manufacturer: BAXTER AG, INDUSTRIESTRASSE, VIENNA, AUSTRIA  
 BAXTER AG, BENATZKYGASSE, VIENNA, AUSTRIA  
 BAXTER HEALTHCARE CORP., ROCHESTER, MI, US  
 BAXTER S.p.A., RIETI, RUFINA, ITALY  
 BAXTER AG, LANGE ALLEE 24-B, VIENNA,  
 AUSTRIA  
 Packer: ADCOCK INGRAM CRITICAL CARE, AEROTON,  
 JOHANNESBURG  
 BAXTER AG, LANGE ALLEE 24, VIENNA, AUSTRIA  
 Laboratory : FPRC: BAXTER AG, LANG ALLEE 24, A-1220, VIENNA,  
 AUSTRIA  
 BAXTER AG, INDUSTRIESTRASSE, VIENNA,  
 AUSTRIA  
 BAXTER AG, BENATZKYGASSE, VIENNA, AUSTRIA  
 BAXTER AG, ORTH/DONAU, AUSTRIA  
 FPRC/FPRR: ADCOCK INGRAM CRITICAL CARE, AEROTON,  
 JOHANNESBURG  
 Shelf-life: 36 months  
 Date of registration: 1 DECEMBER 2006

## MRF 15

Registration number: 38/30.3/0283  
 Name of medicine: IMMUNINE I 200 I.U.  
 Dosage form: INJECTION  
 Active ingredients: EACH VIAL CONTAINS:  
 HUMAN COAGULATION FACTOR IX 1 200,0 I.U.  
 Conditions of registration: 1, 2, 3, 4, 5, 6  
 Applicant: ADCOCK INGRAM CRITICAL CARE LTD  
 Manufacturer: BAXTER AG, INDUSTRIESTRASSE, VIENNA,  
 AUSTRIA  
 BAXTER AG, BENATZKYGASSE, VIENNA, AUSTRIA  
 BAXTER HEALTHCARE CORP., ROCHESTER, MI  
 BAXTER S.p.A., RIETI, RUFINA, ITALY  
 BAXTER AG, LANGE ALLEE 24-B, VIENNA,  
 AUSTRIA  
 Packer: ADCOCK INGRAM CRITICAL CARE, AEROTON,  
 JOHANNESBURG  
 BAXTER AG, LANGE ALLEE 24, A-1220, VIENNA,  
 AUSTRIA  
 Laboratory: FPRC: BAXTER AG, LANG ALLEE 24, A-1220, VIENNA, AUSTRIA  
 BAXTER AG, INDUSTRIESTRASSE, VIENNA,  
 AUSTRIA  
 BAXTER AG, BENATZKYGASSE, VIENNA, AUSTRIA  
 BAXTER AG, ORTH/DONAU, AUSTRIA  
 FPRC/FPRR: ADCOCK INGRAM CRITICAL CARE, AEROTON,  
 JOHANNESBURG  
 Shelf-life: 36 months  
 Date of registration: 1 DECEMBER 2006

## MRF 15

Registration number: A38/21.2/0681  
 Name of medicine: FORMINAL 50  
 Dosage form: TABLET  
 Active ingredients: EACH TABLET CONTAINS:  
 METFORMIN HYDROCHLORIDE 500,0 mg  
 Conditions of registration: 1, 2, 3, 4, 5, 6  
 Applicant: GULF DRUG COMPANY (PTY) LTD  
 Manufacturer: ALEMBIC LTD, GUJARAT, INDIA  
 Packer: ALEMBIC LTD, GUJARAT, INDIA  
 Laboratory: FPRC: ALEMBIC LTD, GUJARAT, INDIA  
 SOUTH AFRICAN BUREAU OF STANDARDS,  
 GROENKLOOF, PRETORIA, RSA  
 INSPECTORATE M&L, ORMONDE, JOHANNESBURG  
 CONSULTING CHEMICAL LABORATORIES,  
 ATLASVILLE, BOKSBURG, RSA  
 PHARMA-Q, INDUSTRIA WEST, RSA  
 INSTITUTE FOR PHARMACEUTICAL SERVICES,  
 SILVERTONDALE, RSA  
 CONSULTING MICROBIOLOGICAL LABORATORY,  
 MOREHILL, BOKSBURG, RSA  
 FPRR: GULF DRUG CO., MOUNT EDGECOMBE, RSA  
 Shelf-life: 24 months (provisional)  
 Date of registration: 1 DECEMBER 2006

## MRF 15

Registration number: A38/21.2/0682  
 Name of medicine: FORMINAL 850  
 Dosage form: TABLET  
 Active ingredients: EACH TABLET CONTAINS:  
 METFORMIN HYDROCHLORIDE 850,0 mg  
 Conditions of registration: 1, 2, 3, 4, 5, 6  
 Applicant: GULF DRUG COMPANY (PTY) LTD  
 Manufacturer: ALEMBIC LTD, GUJARAT, INDIA  
 Packer: ALEMBIC LTD, GUJARAT, INDIA  
 Laboratory: FPRC: ALEMBIC LTD, GUJARAT, INDIA  
 SOUTH AFRICAN BUREAU OF STANDARDS,  
 GROENKLOOF, PRETORIA, RSA  
 INSPECTORATE M&L, ORMONDE, JOHANNESBURG  
 CONSULTING CHEMICAL LABORATORIES,  
 ATLASVILLE, BOKSBURG, RSA  
 PHARMA-Q, INDUSTRIA WEST, RSA  
 INSTITUTE FOR PHARMACEUTICAL SERVICES,  
 SILVERTONDALE, RSA  
 CONSULTING MICROBIOLOGICAL LABORATORY  
 MOREHILL, BOKSBURG, RSA  
 FPRR: GULF DRUG CO., MOUNT EDGECOMBE, RSA  
 Shelf-life: 24 months (provisional)  
 Date of registration: 1 DECEMBER 2006

MRF 15

Registration number: A38/7.5/0710  
Name of medicine: BIOVAC SIMVASTATIN 10  
Dosage form: TABLET  
Active ingredients: EACH TABLET CONTAINS:  
SIMVASTATIN 10,0 mg  
Conditions of registration: 1, 2, 3, 4, 5, 6  
Applicant: ARROW PHARMA SOUTH AFRICA (PTY) LTD  
Manufacturer: RX MANUFACTURING INC, ONTARIO, CANADA  
Packer: RX MANUFACTURING INC, ONTARIO, CANADA  
CONTRACT PHARMACEUTICALS, ONTARIO,  
CANADA  
QUALITY LIMITED, BRIERCLIFFE, BURNLEY, UK  
Laboratory: FPRC: RX MANUFACTURING INC, ONTARIO, CANADA  
QUALITY LIMITED, BRIERCLIFFE, BURNLEY, UK  
RESEARCH INSTITUTE FOR INDUSTRIAL  
PHARMACY, UNIVERSITY, POTCHEFSTROOM  
SEDEK AGRIKEM, KAMEELDRIFT  
FPRR: ARROW PHARMA SA, WOODMEAD, RSA  
Shelf-life: 24 months  
Date of registration: 1 DECEMBER 2006

MRF 15

Registration number: A38/7.5/0711  
Name of medicine: BIOVAC SIMVASTATIN 20  
Dosage form: TABLET  
Active ingredients: EACH TABLET CONTAINS  
SIMVASTATIN 20,0 mg  
Conditions of registration: 1, 2, 3, 4, 5, 6  
Applicant: ARROW PHARMA SOUTH AFRICA (PTY) LTD  
Manufacturer: RX MANUFACTURING INC, ONTARIO, CANADA  
Packer: RX MANUFACTURING INC, ONTARIO, CANADA  
CONTRACT PHARMACEUTICALS, ONTARIO,  
CANADA  
QUALITY LIMITED, BRIERCLIFFE, BURNLEY, UK  
Laboratory: FPRC: RX MANUFACTURING INC, ONTARIO, CANADA  
QUALITY LIMITED, BRIERCLIFFE, BURNLEY, UK  
RESEARCH INSTITUTE FOR INDUSTRIAL  
PHARMACY, UNIVERSITY, POTCHEFSTROOM  
SEDEK AGRIKEM, KAMEELDRIFT  
FPRR: ARROW PHARMA SA, WOODMEAD, RSA  
Shelf-life: 24 months  
Date of registration: 1 DECEMBER 2006

**MRF 15**

Registration number: A38/7.5/0712  
Name of medicine: BIOVAC SIMVASTATIN 40  
Dosage form: TABLET  
Active ingredients: EACH TABLET CONTAINS:  
SIMVASTATIN 40,0 mg  
Conditions of registration: 1, 2, 3, 4, 5, 6  
Applicant: ARROW PHARMA SOUTH AFRICA (PTY) LTD  
Manufacturer: RX MANUFACTURING INC, ONTARIO, CANADA  
Packer: RX MANUFACTURING INC, ONTARIO, CANADA  
CONTRACT PHARMACEUTICALS, ONTARIO,  
CANADA  
QUALITY LIMITED, BRIERCLIFFE, BURNLEY, UK  
Laboratory: FPRC RX MANUFACTURING INC, ONTARIO, CANADA  
QUALITY LIMITED, BRIERCLIFFE BURNLEY, UK  
RESEARCH INSTITUTE FOR INDUSTRIAL  
PHARMACY, UNIVERSITY, POTCHEFSTROOM  
SEDEK AGRIKEM, KAMEELDRIFT  
FPRR: ARROW PHARMA SA, WOODMEAD, RSA  
Shelf-life: 24 months  
Date of registration: 1 DECEMBER 2006

**MRF 15**

Registration number: A39/7.5/0017  
Name of medicine: ASPAVOR 10  
Dosage form: TABLET  
Active ingredients: EACH TABLET CONTAINS  
ATORVASTATIN CALCIUM EQUIVALENT TO  
ATORVASTATIN 10,0 mg  
Conditions of registration: 1, 2, 3, 4, 5, 6  
Applicant: PFIZER LABORATORIES (PTY) LTD  
Manufacturer: GODECKE GmbH, FREIBURG, GERMANY  
PFIZER PHARMACEUTICALS LTD PDPL, VEGA BEJA,  
PUERTO RICO  
PFIZER IRELAND PHARMACEUTICALS, CO CORK,  
IRELAND  
Packer: GODECKE GmbH, FREIBURG, GERMANY  
Laboratory: FPRC: GODECKE GmbH, FREIBURG, GERMANY  
PFIZER GLOBAL MANUFACTURING, RETREAT,  
CAPE TOWN  
FPRR: PFIZER LABORATORIES, SANDTON, RSA  
Shelf-life: 36 months  
Date of registration: 1 DECEMBER 2006

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**MRF 15**

Registration number: A39/7.5/0018  
Name of medicine: ASPAVOR 20  
Dosage form: TABLET  
Active ingredients: EACH TABLET CONTAINS:  
ATORVASTATIN CALCIUM EQUIVALENT TO  
ATORVASTATIN 20,0 mg  
Conditions of registration: 1, 2, 3, 4, 5, 6  
Applicant: PFIZER LABORATORIES (PTY) LTD  
Manufacturer: GODECKE GmbH, FREIBURG, GERMANY  
PFIZER PHARMACEUTICALS LTD PDPL, VEGA  
BEJA, PUERTO RICO  
PFIZER IRELAND PHARMACEUTICALS, CO CORK,  
IRELAND  
Packer: GODECKE GmbH, FREIBURG, GERMANY  
Laboratory: FPRC: GODECKE GmbH, FREIBURG, GERMANY  
PFIZER GLOBAL MANUFACTURING, RETREAT,  
CAPE TOWN  
FPRR: PFIZER LABORATORIES, SANDTON, RSA  
Shelf-life: 36 months  
Date of registration 1 DECEMBER 2006

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**MRF 15**

Registration number: A39/7.5/0019  
Name of medicine: ASPAVOR 40  
Dosage form: TABLET  
Active ingredients: EACH TABLET CONTAINS:  
ATORVASTATIN CALCIUM EQUIVALENT TO  
ATORVASTATIN 40,0 mg  
Conditions of registration: 1, 2, 3, 4, 5, 6  
Applicant: PFIZER LABORATORIES (PTY) LTD  
Manufacturer: GODECKE GmbH, FREIBURG, GERMANY  
PFIZER PHARMACEUTICALS LTD PDPL, VEGA BEJA,  
PUERTO RICO  
PFIZER IRELAND PHARMACEUTICALS, CO CORK,  
IRELAND  
Packer: GODECKE GmbH, FREIBURG, GERMANY  
Laboratory: FPRC: GODECKE GmbH, FREIBURG, GERMANY  
PFIZER GLOBAL MANUFACTURING, RETREAT, CAPE  
TOWN  
FPRR: PFIZER LABORATORIES, SANDTON, RSA  
Shelf-life: 36 months  
Date of registration: 1 DECEMBER 2006

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**MRF 15**

Registration number: A39/20.2.8/0112  
Name of medicine: VARI-NEVIRAPINE 200 mg  
Dosage form: TABLET  
Active ingredients: EACH TABLET CONTAINS:  
NEVIRAPINE 200,0 mg  
Conditions of registration: 1, 2, 3, 4, 5, 6  
Applicant: LEBASI PHARMACEUTICALS cc  
Manufacturer: VARICHEM PHARMACEUTICALS (PVT) LTD, HARARE,  
ZIMBABWE  
Packer: VARICHEM PHARMACEUTICALS (PVT) LTD, HARARE,  
ZIMBABWE  
Laboratory: FPRC: VARICHEM PHARMACEUTICALS (PVT) LTD, HARARE,  
ZIMBABWE  
RESEARCH INSTITUTE FOR INDUSTRIAL  
PHARMACY, UNIVERSITY, POTCHEFSTROOM  
INSTITUTE FOR PHARMACEUTICAL SERVICES,  
SILVERTONDALE, RSA  
CONSULTING CHEMICAL LABORATORIES,  
ATLASVILLE, BOKSBURG  
FPRR: LEBASI PHARMACEUTICALS, POTCHEFSTROOM  
Shelf-life: 24 months  
Date of registration: 1 DECEMBER 2006

**MRF 15**

Registration number: A39/11.5/0115  
Name of medicine: ISPAGEL  
Dosage form: POWDER  
Active ingredients: EACH SACHET CONTAINS:  
ISPAGHULA HUSK 3,5 g  
Conditions of registration: 1, 2, 3, 4, 5, 6  
Applicant: PHARMA DYNAMICS (PTY) LTD  
Manufacturer: LABORATORIOS BELMAC S.A., ZARAGOZA, SPAIN  
Packer: LABORATORIOS BELMAC S.A., ZARAGOZA, SPAIN  
DIVPHARM MANUFACTURING & PACKAGING,  
LONGDALE, JOHANNESBURG  
TECHNIKON LABORATORIES, FLORIDA, RSA  
PHARMACEUTICAL ENTERPRISES, N'DABENI, KZN  
IMPILO DRUGS, ISITHEBE, KZN  
Laboratory: FPRC: LABORATORIOS BELMAC S.A., ZARAGOZA, SPAIN  
CONSULTING CHEMICAL LABORATORIES,  
ATLASVILLE, BOKSBURG, RSA  
TECHNIKON LABORATORIES, FLORIDA, RSA  
IMPILO DRUGS, ISITHEBE, KZN  
FPRR: PHARMA DYNAMICS, SILVERWOOD, WESTLAKE, RSA  
Shelf-life: 48 months  
Date of registration: 1 DECEMBER 2006

**MRF 15**

Registration number: A39/11.5/0135  
Name of medicine: PHARMA DYNAMICS ISPAGHULA HUSK  
Dosage form: POWDER  
Active ingredients: EACH SACHET CONTAINS:  
ISPAGHULA HUSK 3,5 g  
Conditions of registration: 1, 2, 3, 4, 5, 6  
Applicant: PHARMA DYNAMICS (PTY) LTD  
Manufacturer: LABORATORIOS BELMAC S.A., ZARAGOZA, SPAIN  
Packer: LABORATORIOS BELMAC S.A., ZARAGOZA, SPAIN  
DIVPHARM MANUFACTURING & PACKAGING,  
LONGDALE, JOHANNESBURG  
TECHNIKON LABORATORIES, FLORIDA, RSA  
PHARMACEUTICAL ENTERPRISES, N'DABENI, KZN  
IMPILO DRUGS, ISITHEBE, KZN  
Laboratory: FPRC: LABORATORIOS BELMAC S.A., ZARAGOZA, SPAIN  
CONSULTING CHEMICAL LABORATORIES,  
ATLASVILLE, BOKSBURG, RSA  
TECHNIKON LABORATORIES, FLORIDA, RSA  
IMPILO DRUGS, ISITHEBE, KZN  
FPRR PHARMA DYNAMICS, SILVERWOOD, WESTLAKE,  
RSA  
Shelf-life: 48 months  
Date of registration: 1 DECEMBER 2006

**MRF 15**

Registration number: A39/20.2.8/0279  
Name of medicine: DAS – STAVUDINE 40 mg  
Dosage form: CAPSULES  
Active ingredients: EACH CAPSULE CONTAINS:  
STAVUDINE 40,0 mg  
Conditions of registration: 1, 2, 3, 4, 5, 6  
Applicant: STRIDES S.A. (PTY) LTD  
Manufacturer: STRIDES ARCOLAB LTD, ANEKAL TALUK, BANGALORE,  
INDIA  
Packer: STRIDES ARCOLAB LTD, ANEKAL TALUK, BANGALORE,  
INDIA  
Laboratory: FPRC: STRIDES ARCOLAB LTD, ANEKAL TALUK, BANGALORE,  
INDIA  
COLUMBIA PHARMACEUTICALS, BARDENE, BOKSBURG  
INSTITUTE FOR PHARMACEUTICAL SERVICES,  
SILVERTONDALE, RSA  
NOVARTIS S.A., SPARTAN, KEMPTON PARK  
FPRR: STRIDES S.A., ARCADIA, PRETORIA  
Shelf-life: 24 months  
Date of registration: 1 DECEMBER 2006

**MRF 15**

Registration number: A39/23/0182  
Name of medicine: AMINOVEN INFANT 10 %  
Dosage form: SOLUTION  
Active ingredients: EACH 1000,0 ml SOLUTION CONTAINS:  
TOTAL AMINO ACIDS 100,0 g  
SEE ADDENDUM  
Conditions of registration: 1, 2, 3, 4, 5, 6  
Applicant: FRESENIUS KABI SOUTH AFRICA (PTY) LTD  
Manufacturer: FRESENIUS KABI AUSTRIA GmbH, GRAZ, AUSTRIA  
Packer: FRESENIUS KABI AUSTRIA GmbH, GRAZ, AUSTRIA  
Laboratory: FPRC: FRESENIUS KABI AUSTRIA GmbH, GRAZ, AUSTRIA  
KHULULEKANI LABORATORIES, MIDRAND, RSA  
BODENE t/a INTRAMED, KORSTEN, PORT ELIZABETH  
FPRR: FRESENIUS KABI, MIDRAND, RSA  
Shelf-life: 24 months  
Date of registration: 1 DECEMBER 2006

**ADDENDUM:**

L-isoleucine	8,00 g
L-leucine	13,00 g
L-methionine	3,12 g
L-lysine-acetate	12,00g
L-phenylalanine	3,75 g
L-threonine	4,40 g
L-tryptophan	2,01 g
L-valine	9,00 g
L-arginine	7,50 g
L-histidine	4,76 g
L-alanine	9,30 g
Glycine	4,15 g
L-proline	9,71 g
L-serine	7,67 g
N-acetyl- L-cysteine	0,77 g
Taurine	0,40 g
N-acetyl-L-tyrosine	5,176 g
L-malic acid	2,62 g

## MRF 15

Registration number: A39/25.2/0275  
 Name of medicine: OLICLINOMEL N8-800  
 Dosage form: INFUSION  
 Active ingredients: EACH BAG CONTAINS  
 a) 12 % AMINO ACIDS SOLUTION compartment  
 b) 31,25 % GLUCOSE SOLUTION compartment  
 c) 15 % LIPID EMULSION compartment  
 SEE ADDENDUM  
 Conditions of registration: 1, 2, 3, 4, 5, 6  
 Applicant: ADCOCK INGRAM CRITICAL CARE (PTY) LTD  
 Manufacturer: CLINTEC PARENTERAL S.A., MONTARGIS, FRANCE  
 BAXTER S.A., LESSINES, BELGIUM  
 Packer: CLINTEC PARENTERAL S.A., MONTARGIS, FRANCE  
 BAXTER S.A., LESSINES, BELGIUM  
 ADCOCK INGRAM CRITICAL CARE, AEROTON,  
 JOHANNESBURG, RSA  
 Laboratory: FPRC: CLINTEC PARENTERAL S.A., MONTARGIS, FRANCE  
 BAXTER S.A., LESSINES, BELGIUM  
 FPRC/FPRR: ADCOCK INGRAM CRITICAL CARE, AEROTON,  
 JOHANNESBURG, RSA  
 Shelf-life: 24 months  
 Date of registration: 1 DECEMBER 2006

## ADDENDUM

- a) 12 % AMINO ACIDS SOLUTION compartment containing per 1,0 litre:
- |                 |         |         |
|-----------------|---------|---------|
| L-alanine       | 25,88 g |         |
| L-arginine      |         | 14,38 g |
| Glycine         | 12,88 g |         |
| L-histidine     |         | 6,00 g  |
| L-isoleucine    | 7,50 g  |         |
| L-leucine       | 9,13 g  |         |
| L-lysine        | 7,25 g  |         |
| L-methionine    | 5,00 g  |         |
| L-phenylalanine | 7,00 g  |         |
| L-proline       | 8,50 g  |         |
| L-serine        | 6,25 g  |         |
| L-threonine     | 5,25 g  |         |
| L-tryptophan    | 2,25 g  |         |
| L-tyrosine      | 0,50 g  |         |
| L-valine        | 7,25 g  |         |
- b) 31,25 % GLUCOSE SOLUTION compartment containing per 1,0 litre:
- |  |          |
|--|----------|
| Glucose monohydrate equivalent to<br>Glucose | 312,50 g |
|--|----------|
- c) 15 % LIPID EMULSION compartment containing per 1,0 litre:
- |  |          |
|--|----------|
| Refined soya-bean oil +<br>Refined olive oil | 150,00 g |
|--|----------|

**MRF 15**

Registration number: A39/28/0614

Name of medicine: OPTIRAY 300-75 ml

Dosage form: SOLUTION

Active ingredients: EACH 1,0 ml SOLUTION CONTAINS:  
IOVERSOL 300,0 mg

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

Applicant: TYCO HEALTHCARE (PTY) LTD

Manufacturer: TYCO HEALTHCARE INC, RALEIGH, NORTH CAROLINA, USA  
TYCO HEALTHCARE INC, POINTE CLAIRE, QUEBEC, CANADA

Packer: TYCO HEALTHCARE INC, RALEIGH, NORTH CAROLINA, USA  
TYCO HEALTHCARE INC, POINTE CLAIRE, QUEBEC, CANADA

Laboratory: FPRC: TYCO HEALTHCARE INC, RALEIGH, NORTH CAROLINA, USA  
TYCO HEALTHCARE INC, POINTE CLAIRE, QUEBEC, CANADA  
FPRR: BIOCHEMICAL & SCIENTIFIC cc, HILTON, KZN  
TYCO HEALTHCARE, MIDRAND, RSA

Shelf-life: 36 months

Date of registration: 1 DECEMBER 2006

**MRF 15**

Registration number: A39/20.1.1/618

Name of medicine: RAN-CLARITHROMYCIN MR 500

Dosage form: TABLET

Active ingredients: EACH TABLET CONTAINS:  
CLARITHROMYCIN 500,0 mg

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

Applicant: RANBAXY (S.A.) (PTY) LTD

Manufacturer: RANBAXY LABORATORIES LTD, DEWAS, INDIA

Packer: RANBAXY LABORATORIES LTD, DEWAS, INDIA

Laboratory: FPRC: RANBAXY LABORATORIES LTD, DEWAS, INDIA  
KHULULEKANI LABORATORY SERVICES, MIDRAND, RSA  
CENTRE FOR QUALITY ASSURANCE OF MEDICINES,  
UNIVERSITY, POTCHEFSTROOM  
FPRR: RANBAXY, CENTURION, RSA

Shelf-life: 24 months (provisional)

Date of registration: 1 DECEMBER 2006

**MRF 15**

Registration number: A39/20.1.1/619  
Name of medicine: KLARITHRAN MR 500  
Dosage form: TABLET  
Active ingredients: EACH TABLET CONTAINS:  
CLARITHROMYCIN 500,0 mg  
Conditions of registration: 1, 2, 3, 4, 5, 6  
Applicant: RANBAXY (S.A.) (PTY) LTD  
Manufacturer: RANBAXY LABORATORIES LTD, DEWAS, INDIA  
Packer: RANBAXY LABORATORIES LTD, DEWAS, INDIA  
Laboratory: FPRC: RANBAXY LABORATORIES LTD, DEWAS, INDIA  
KHULULEKANI LABORATORY SERVICES, MIDRAND,  
RSA  
CENTRE FOR QUALITY ASSURANCE OF MEDICINES,  
UNIVERSITY, POTCHEFSTROOM  
FPRR: RANBAXY, CENTURION, RSA  
Shelf-life: 24 months (provisional)  
Date of registration: 1 DECEMBER 2006

**MRF 15**

Registration number: A39/24/0620  
Name of medicine: POTASSIUM CHLORIDE B BRAUN 7,45 %  
Dosage form: SOLUTION  
Active ingredients: EACH 100,0 ml SOLUTION CONTAINS:  
POTASSIUM CHLORIDE 7,456 g  
Conditions of registration: 1, 2, 3, 4, 5, 6  
Applicant: B BRAUN MEDICAL (PTY) LTD  
Manufacturer: B BRAUN MELSUNGEN AG, BERLIN, GERMANY  
Packer: B BRAUN MELSUNGEN AG, BERLIN, GERMANY  
Laboratory: FPRC: B BRAUN MELSUNGEN AG, BERLIN, GERMANY  
CONSULTING CHEMICAL LABORATORIES, ATLASVILLE,  
BOKSBURG  
FPRR: B BRAUN MEDICAL, HONEYDEW, RSA  
Shelf-life: 36 months  
Date of registration: 1 DECEMBER 2006

**MRF 15**

Registration number: A39/24/0621  
Name of medicine: POTASSIUM CHLORIDE B BRAUN 14,9 %  
Dosage form: SOLUTION  
Active ingredients: EACH 100,0 ml SOLUTION CONTAINS:  
POTASSIUM CHLORIDE 14,9 g  
Conditions of registration: 1, 2, 3, 4, 5, 6  
Applicant: B BRAUN MEDICAL (PTY) LTD  
Manufacturer: B BRAUN MELSUNGEN AG, BERLIN, GERMANY  
Packer: B BRAUN MELSUNGEN AG, BERLIN, GERMANY  
Laboratory: FPRC: B BRAUN MELSUNGEN AG, BERLIN, GERMANY  
CONSULTING CHEMICAL LABORATORIES,  
ATLASVILLE, BOKSBURG  
FPRR: B BRAUN MEDICAL, HONEYDEW, RSA  
Shelf-life: 36 months  
Date of registration: 1 DECEMBER 2006

**MRF 15**

Registration number: A40/20.1.1/0019  
Name of medicine: SPEC-CEFAXONE 250 mg  
Dosage form: INJECTION  
Active ingredients: EACH VIAL CONTAINS:  
CEFTRIAZONE SODIUM EQUIVALENT TO  
CEFTRIAZONE 250,0 mg  
Conditions of registration: 1, 2, 3, 4, 5, 6  
Applicant: SPECPHARM (PTY) LTD  
Manufacturer: LUPIN LTD, MANDIDEEP, MADHYA PRADESH, INDIA  
Packer: LUPIN LTD, MANDIDEEP, MADHYA PRADESH, INDIA  
Laboratory: FPRC: LUPIN LTD, MANDIDEEP, MADHYA PRADESH, INDIA  
ANALYTICON, TERENURE, KEMPTON PARK  
FPRR: SPECPHARM, SANDTON, RSA  
Shelf-life: 24 months  
Date of registration: 1 DECEMBER 2006

**MRF 15**

Registration number: A40/20.1.1/0020  
Name of medicine: SPEC-CEFAXONE 500 mg  
Dosage form: INJECTION  
Active ingredients: EACH VIAL CONTAINS  
CEFTRIAXONE SODIUM EQUIVALENT TO  
CEFTRIAXONE 500,0 mg  
Conditions of registration: 1, 2, 3, 4, 5, 6  
Applicant: SPECPHARM (PTY) LTD  
Manufacturer: LUPIN LTD, MANDIDEEP, MADHYA PRADESH, INDIA  
Packer: LUPIN LTD, MANDIDEEP, MADHYA PRADESH, INDIA  
Laboratory: FPRC: LUPIN LTD, MANDIDEEP, MADHYA PRADESH, INDIA  
ANALYTICON, TERENURE, KEMPTON PARK  
FPRR: SPECPHARM, SANDTON, RSA  
Shelf-life: 24 months  
Date of registration: 1 DECEMBER 2006

**MRF 15**

Registration number: A40/20.1.1/0021  
Name of medicine: SPEC-CEFAXONE 1 g  
Dosage form: INJECTION  
Active ingredients: EACH VIAL CONTAINS:  
CEFTRIAXONE SODIUM EQUIVALENT TO  
CEFTRIAXONE 1000,0 mg  
Conditions of registration: 1, 2, 3, 4, 5, 6  
Applicant: SPECPHARM (PTY) LTD  
Manufacturer: LUPIN LTD, MANDIDEEP, MADHYA PRADESH, INDIA  
Packer: LUPIN LTD, MANDIDEEP, MADHYA PRADESH, INDIA  
Laboratory: FPRC: LUPIN LTD, MANDIDEEP, MADHYA PRADESH, INDIA  
ANALYTICON, TERENURE, KEMPTON PARK  
FPRR: SPECPHARM, SANDTON, RSA  
Shelf-life: 24 months  
Date of registration: 1 DECEMBER 2006



## MRF 15

Registration number: A40/10.2.1/0064  
 Name of medicine: SEREVENT INHALER CFC-FREE  
 Dosage form: INHALER  
 Active ingredients: EACH ACTUATION DELIVERS:  
 SALMETEROL XINAFOATE EQUIVALENT TO  
 SALMETEROL 25,0 ug  
 Conditions of registration: 1, 2, 3, 4, 5, 6  
 Applicant: GLAXOSMITHKLINE SOUTH AFRICA (PTY) LTD  
 Manufacturer: GLAXO WELLCOME PRODUCTION, EVREUX, FRANCE  
 Packer: GLAXO WELLCOME PRODUCTION, EVREUX, FRANCE  
 GLAXOSMITHKLINE S.A., EPPING, CAPE TOWN  
 Laboratory:FPRC: GLAXO WELLCOME PRODUCTION, EVREUX, FRANCE  
 FPRC/FPRR: GLAXOSMITHKLINE S.A., EPPING, CAPE TOWN  
 Shelf-life: 24 months (provisional)  
 Date of registration: 1 DECEMBER 2006

## MRF 15

Registration number: A40/7.1/0106  
 Name of medicine: ZANIDIP 20  
 Dosage form: TABLET  
 Active ingredients: EACH TABLET CONTAINS:  
 LERCANIDIPINE HYDROCHLORIDE 20,0 mg  
 Conditions of registration: 1, 2, 3, 4, 5, 6  
 Applicant: PHARMAPLAN (PTY) LTD  
 Manufacturer: RECORDATI INDUSTRIA CHIMICA E FARMACEUTICA,  
 MILAN, ITALY  
 Packer: RECORDATI INDUSTRIA CHIMICA E FARMACEUTICA,  
 MILAN, ITALY  
 Laboratory: FPRC: RECORDATI INDUSTRIA CHIMICA E FARMACEUTICA,  
 MILAN, ITALY  
 CONSULTING CHEMICAL LABORATORIES, ATLASVILLE,  
 BOKSBURG  
 FPRR: PHARMAPLAN, MIDRAND, RSA  
 Shelf-life: 36 months  
 Date of registration: 1 DECEMBER 2006

**MRF 15**

Registration number: A40/15.1/0164  
Name of medicine: VIGAMOX EYE DROPS  
Dosage form: DROPS  
Active ingredients: EACH 1,0 ml SOLUTION CONTAINS:  
MOXIFLOXACIN HYDROCHLORIDE EQUIVALENT TO  
MOXIFLOXACIN 5,0 mg  
Conditions of registration: 1, 2, 3, 4, 5, 6  
Applicant: ALCON LABORATORIES (S.A.) (PTY) LTD  
Manufacturer: S.A. ALCON-COUVREUR N.V, PUURS, BELGIUM  
Laboratory: FPRC: S.A. ALCON-COUVREUR N.V, PUURS, BELGIUM  
RESEARCH INSTITUTE FOR INDUSTRIAL PHARMACY,  
UNIVERSITY, POTCHEFSTROOM  
FPRR: ALCON LABORATORIES, BRYANSTON,  
JOHANNESBURG, RSA  
Shelf-life: 24 months  
Date of registration: 1 DECEMBER 2006

**MRF 15**

Registration number: A40/21.2/0193  
Name of medicine: AUSTELL – GLIMEPIRIDE 1 mg  
Dosage form: TABLET  
Active ingredients: EACH TABLET CONTAINS:  
GLIMEPIRIDE 1,0 mg  
Conditions of registration: 1, 2, 3, 4, 5, 6  
Applicant: AUSTELL LABORATORIES (PTY) LTD  
Manufacturer: IPCA LABORATORIES LTD, ATHAL, DADRA & NAGAR  
HAVELI, INDIA  
Packer: IPCA LABORATORIES LTD, ATHAL, DADRA & NAGAR  
HAVELI, INDIA  
Laboratory: FPRC: IPCA LABORATORIES LTD, ATHAL, DADRA & NAGAR  
HAVELI, INDIA  
SOUTH AFRICAN BUREAU OF STANDARDS,  
GROENKLOOF, PRETORIA  
INSPECTORATE M&L, ORMONDE, JOHANNESBURG  
INSTITUTE FOR PHARMACEUTICAL SERVICES,  
SILVERTONDALE  
FPRR: AUSTELL LABORATORIES, SPRINGFIELD,  
JOHANNESBURG  
Shelf-life: 24 months (provisional)  
Date of registration: 1 DECEMBER 2006

## MRF 15

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Registration number: A40/21.2/0194

Name of medicine: AUSTELL – GLIMEPIRIDE 2 mg

Dosage form: TABLET

Active ingredients: EACH TABLET CONTAINS:  
GLIMEPIRIDE 2,0 mg

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

Applicant: AUSTELL LABORATORIES (PTY) LTD

Manufacturer: IPCA LABORATORIES LTD, ATHAL, DADRA & NAGAR HAVELI, INDIA

Packer: IPCA LABORATORIES LTD, ATHAL, DADRA & NAGAR HAVELI, INDIA

Laboratory: FPRC: IPCA LABORATORIES LTD, ATHAL, DADRA & NAGAR HAVELI, INDIA  
SOUTH AFRICAN BUREAU OF STANDARDS,  
GROENKLOOF, PRETORIA  
INSPECTORATE M&L, ORMONDE, JOHANNESBURG  
INSTITUTE FOR PHARMACEUTICAL SERVICES,  
SILVERTONDALE

FPRR: AUSTELL LABORATORIES, SPRINGFIELD,  
JOHANNESBURG

Shelf-life: 24 months (provisional)

Date of registration: 1 DECEMBER 2006

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## MRF 15

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Registration number: A40/21.2/0195

Name of medicine: AUSTELL – GLIMEPIRIDE 3 mg

Dosage form: TABLET

Active ingredients: EACH TABLET CONTAINS:  
GLIMEPIRIDE 3,0 mg

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

Applicant: AUSTELL LABORATORIES (PTY) LTD

Manufacturer: IPCA LABORATORIES LTD, ATHAL, DADRA & NAGAR HAVELI, INDIA

Packer: IPCA LABORATORIES LTD, ATHAL, DADRA & NAGAR HAVELI, INDIA

Laboratory: FPRC: IPCA LABORATORIES LTD, ATHAL, DADRA & NAGAR HAVELI, INDIA  
SOUTH AFRICAN BUREAU OF STANDARDS,  
GROENKLOOF, PRETORIA  
INSPECTORATE M&L, ORMONDE, JOHANNESBURG  
INSTITUTE FOR PHARMACEUTICAL SERVICES,  
SILVERTONDALE

FPRR: AUSTELL LABORATORIES, SPRINGFIELD,  
JOHANNESBURG

Shelf-life: 24 months (provisional)

Date of registration: 1 DECEMBER 2006

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## MRF 15

Registration number: A40/21.2/0196  
 Name of medicine: AUSTELL -- GLIMEPIRIDE 4 mg  
 Dosage form: TABLET  
 Active ingredients: EACH TABLET CONTAINS:  
 GLIMEPIRIDE 4,0 mg  
 Conditions of registration: 1, 2, 3, 4, 5, 6  
 Applicant: AUSTELL LABORATORIES (PTY) LTD  
 Manufacturer: IPCA LABORATORIES LTD, ATHAL, DADRA & NAGAR  
 HAVELI, INDIA  
 Packer: IPCA LABORATORIES LTD, ATHAL, DADRA & NAGAR  
 HAVELI, INDIA  
 Laboratory: FPRC: IPCA LABORATORIES LTD, ATHAL, DADRA & NAGAR  
 HAVELI, INDIA  
 SOUTH AFRICAN BUREAU OF STANDARDS,  
 GROENKLOOF, PRETORIA  
 INSPECTORATE M&L, ORMONDE, JOHANNESBURG  
 INSTITUTE FOR PHARMACEUTICAL SERVICES,  
 SILVERTONDALE  
 FPRR: AUSTELL LABORATORIES, SPRINGFIELD,  
 JOHANNESBURG  
 Shelf-life: 24 months (provisional)  
 Date of registration: 1 DECEMBER 2006

## MRF 15

Registration number: A40/34/0258  
 Name of medicine: COPAXONE 20 mg/ml  
 Dosage form: INJECTION  
 Active ingredients: EACH 1,0 ml SOLUTION CONTAINS:  
 GLATIRAMER ACETATE 20,0 mg  
 Conditions of registration: 1, 2, 3, 4, 5, 6  
 Applicant: TEVA PHARMACEUTICALS (PTY) LTD  
 Manufacturer: TEVA PHARMACEUTICAL INDUSTRIES LTD, KFAR-  
 SAVA, ISRAEL  
 Packer: TEVA PHARMACEUTICAL INDUSTRIES LTD, KFAR-  
 SAVA, ISRAEL  
 Laboratory: FPRC: TEVA PHARMACEUTICAL INDUSTRIES LTD, KFAR-  
 SAVA, ISRAEL  
 RESEARCH INSTITUTE FOR INDUSTRIAL PHARMACY,  
 UNIVERSITY, POTCHEFSTROOM  
 SOUTH AFRICAN BUREAU OF STANDARDS,  
 GROENKLOOF, PRETORIA, RSA  
 FPRR: TEVA PHARMACEUTICALS, RUIMSIG, ROODEPOORT,  
 RSA  
 Shelf-life: 24 months  
 Date of registration: 1 DECEMBER 2006

## MRF 15

Registration number: A40/20.1.2/0285  
 Name of medicine: FLUPEN-250  
 Dosage form: CAPSULE  
 Active ingredients: EACH CAPSULE CONTAINS:  
 FLUCLOXACILLIN SODIUM EQUIVALENT TO:  
 FLUCLOXACILLIN 250,0 mg  
 Conditions of registration: 1, 2, 3, 4, 5, 6  
 Applicant: DEZZO TRADING (392) PTY LTD t/a INDO PHARMA  
 Manufacturer: KOPRAN LTD, KHALAPUR, RAIGAD, INDIA  
 Packer: KOPRAN LTD, KHALAPUR, RAIGAD, INDIA  
 Laboratory: FPRC: KOPRAN LTD, KHALAPUR, RAIGAD, INDIA  
 CONSULTING CHEMICAL LABORATORIES,  
 ATLASVILLE, BOKSBURG  
 FPRR: DEZZO TRADING (392) t/a INDO PHARMA, LENASIA,  
 JOHANNESBURG  
 Shelf-life: 24 months (provisional)  
 Date of registration: 1 DECEMBER 2006

## MRF 15

Registration number: A40/5.10/0322  
 Name of medicine: ONICIT  
 Dosage form: INJECTION  
 Active ingredients: EACH 1,0 ml SOLUTION CONTAINS:  
 PALONOSETRON HYDROCHLORIDE EQUIVALENT TO  
 PALONOSETRON 50,0 ug  
 Conditions of registration: 1, 2, 3, 4, 5, 6  
 Applicant: PFIZER LABORATORIES (PTY) LTD  
 Manufacturer: CARDINAL HEALTH, ALBUQUERQUE, NEW MEXICO,  
 USA  
 Packer: CARDINAL HEALTH, ALBUQUERQUE, NEW MEXICO,  
 USA  
 HELSINN BIREX PHARMACEUTICALS, DUBLIN,  
 IRELAND  
 Laboratory: FPRC: CARDINAL HEALTH, ALBUQUERQUE, NEW MEXICO,  
 USA  
 HELSINN BIREX PHARMACEUTICALS, DUBLIN,  
 IRELAND  
 PFIZER GLOBAL MANUFACTURING, RETREAT, CAPE  
 TOWN  
 FPRR: PFIZER LABORATORIES, SANDTON, RSA  
 Shelf-life: 36 months  
 Date of registration: 1 DECEMBER 2006

**MRF 15**

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Registration number: A40/11.4.3/0482

Name of medicine: PANTOCID 40

Dosage form: TABLET

Active ingredients: EACH TABLET CONTAINS:  
PANTOPRAZOLE SODIUM SESQUIHYDRATE  
EQUIVALENT TO  
PANTOPRAZOLE 40,0 mg

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

Applicant: PHARMAPLAN (PTY) LTD

Manufacturer: M J PHARMACEUTICALS LTD, PANCHMAHAL,  
GUJARAT, INDIA

Packer: M J PHARMACEUTICALS LTD, PANCHMAHAL,  
GUJARAT, INDIA

Laboratory: FPRC: M J PHARMACEUTICALS LTD, PANCHMAHAL,  
GUJARAT, INDIA  
CONSULTING CHEMICAL LABORATORIES,  
ATLASVILLE, BOKSBURG

FPRR: PHARMAPLAN, MIDRAND, RSA

Shelf-life: 24 months (provisional)

Date of registration: 1 DECEMBER 2006

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**MRF 15**

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Registration number: A40/3.2/0514

Name of medicine: OSTOMIR 70

Dosage form: TABLET

Active ingredients: EACH TABLET CONTAINS:  
ALENDRONATE SODIUM EQUIVALENT TO  
ALENDRONIC ACID 70,0 mg

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

Applicant: SANDOZ (PTY) LTD

Manufacturer: LEK PHARMACEUTICALS dd, VEROVSKOVA,  
LJUBLJANA, SLOVENIA

Packer: LEK PHARMACEUTICALS dd, VEROVSKOVA,  
LJUBLJANA, SLOVENIA

Laboratory: FPRC: LEK PHARMACEUTICALS dd, VEROVSKOVA,  
LJUBLJANA, SLOVENIA  
NOVARTIS, SPARTAN, KEMPTON PARK  
SOUTH AFRICAN BUREAU OF STANDARDS,  
GROENKLOOF, PRETORIA  
ANALYTICON, TERENURE, KEMPTON PARK

FPRR: SANDOZ, SPARTAN, KEMPTON PARK

Shelf-life: 24 months (provisional)

Date of registration: 1 DECEMBER 2006

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## MRF 15

Registration number: A40/3.2/0515  
 Name of medicine: SANDOZ ALENDRONATE 70  
 Dosage form: TABLET  
 Active ingredients: EACH TABLET CONTAINS:  
 ALENDRONATE SODIUM EQUIVALENT TO  
 ALENDRONIC ACID 70,0 mg  
 Conditions of registration: 1, 2, 3, 4, 5, 6  
 Applicant: SANDOZ (PTY) LTD  
 Manufacturer: LEK PHARMACEUTICALS dd, VEROVSKOVA,  
 LJUBLJANA, SLOVENIA  
 Packer: LEK PHARMACEUTICALS dd, VEROVSKOVA,  
 LJUBLJANA, SLOVENIA  
 Laboratory: FPRC: LEK PHARMACEUTICALS dd, VEROVSKOVA,  
 LJUBLJANA, SLOVENIA  
 NOVARTIS, SPARTAN, KEMPTON PARK  
 SOUTH AFRICAN BUREAU OF STANDARDS,  
 GROENKLOOF, PRETORIA  
 ANALYTICON, TERENCE, KEMPTON PARK  
 FPRR: SANDOZ, SPARTAN, KEMPTON PARK  
 Shelf-life: 24 months (provisional)  
 Date of registration: 1 DECEMBER 2006

## MRF 15

Registration number: A40/20.2.3/0534  
 Name of medicine: RIFINAH 150/75  
 Dosage form: TABLET  
 Active ingredients: EACH TABLET CONTAINS:  
 RIFAMPICIN 150,0 mg  
 ISONIAZID 75,0 mg  
 Conditions of registration: 1, 2, 3, 4, 5, 6  
 Applicant: AVENTIS PHARMA (PTY) LTD  
 Manufacturer: AVENTIS PHARMA, WALTLOO, PRETORIA  
 Packer: AVENTIS PHARMA, WALTLOO, PRETORIA  
 Laboratory: FPRC: SOUTH AFRICAN BUREAU OF STANDARDS,  
 GROENKLOOF, PRETORIA  
 RESEARCH INSTITUTE FOR INDUSTRIAL PHARMACY,  
 UNIVERSITY, POTCHEFSTROOM  
 FPRC/FPRR: AVENTIS PHARMA, WALTLOO, PRETORIA  
 Shelf-life: 24 months (provisional)  
 Date of registration: 1 DECEMBER 2006

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**MRF 15**

Registration number: A40/1.2/0584  
Name of medicine: SANDOZ MIRTAZAPINE 15  
Dosage form: TABLET  
Active ingredients: EACH TABLET CONTAINS:  
MIRTAZAPINE 15,0 mg  
Conditions of registration: 1, 2, 3, 4, 5, 6  
Applicant: SANDOZ (PTY) LTD  
Manufacturer: NOVARTIS LTD, GAZIPUR, BANGLADESH  
Packer: NOVARTIS LTD, GAZIPUR, BANGLADESH  
NOVARTIS S.A., SPARTAN, KEMPTON PARK  
Laboratory: FPRC: NOVARTIS LTD, GAZIPUR, BANGLADESH  
NOVARTIS S.A., SPARTAN, KEMPTON PARK  
SOUTH AFRICAN BUREAU OF STANDARDS,  
GROENKLOOF, PRETORIA  
ANALYTICON, TERENURE, KEMPTON PARK  
FPRR: SANDOZ, SPARTAN, KEMPTON PARK  
Shelf-life: 24 months  
Date of registration: 1 DECEMBER 2006

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**MRF 15**

Registration number: A40/1.2/0585  
Name of medicine: SANDOZ MIRTAZAPINE 30  
Dosage form: TABLET  
Active ingredients: EACH TABLET CONTAINS:  
MIRTAZAPINE 30,0 mg  
Conditions of registration: 1, 2, 3, 4, 5, 6  
Applicant: SANDOZ (PTY) LTD  
Manufacturer: NOVARTIS LTD, GAZIPUR, BANGLADESH  
Packer: NOVARTIS LTD, GAZIPUR, BANGLADESH  
NOVARTIS S.A., SPARTAN, KEMPTON PARK  
Laboratory: FPRC: NOVARTIS LTD, GAZIPUR, BANGLADESH  
NOVARTIS S.A., SPARTAN, KEMPTON PARK  
SOUTH AFRICAN BUREAU OF STANDARDS,  
GROENKLOOF, PRETORIA  
ANALYTICON, TERENURE, KEMPTON PARK  
FPRR: SANDOZ, SPARTAN, KEMPTON PARK  
Shelf-life: 24 months  
Date of registration: 1 DECEMBER

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**MRF 15**

Registration number: A40/1.2/0586

Name of medicine SANDOZ MIRTAZAPINE 45

Dosage form: TABLET

Active ingredients: EACH TABLET CONTAINS:  
MIRTAZAPINE 45,0 mg

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

Applicant: SANDOZ (PTY) LTD

Manufacturer: NOVARTIS LTD, GAZIPUR, BANGLADESH

Packer: NOVARTIS LTD, GAZIPUR, BANGLADESH  
NOVARTIS S.A., SPARTAN, KEMPTON PARK

Laboratory: FPRC: NOVARTIS LTD, GAZIPUR, BANGLADESH  
NOVARTIS S.A., SPARTAN, KEMPTON PARK  
SOUTH AFRICAN BUREAU OF STANDARDS  
GROENKLOOF, PRETORIA  
ANALYTICON, TERENURE, KEMPTON PARK

FPRR: SANDOZ, SPARTAN, KEMPTON PARK

Shelf-life: 24 months

Date of registration: 1 DECEMBER 2006

**MRF 15**

Registration number: A40/1.2/0654

Name of medicine: AUROLIFT 50 mg

Dosage form: TABLET

Active ingredients: EACH TABLET CONTAINS:  
SERTRALINE HYDROCHLORIDE EQUIVALENT TO  
SERTRALINE 50,0 mg

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

Applicant: AUROBINDO PHARMA (PTY) LTD

Manufacturer: AUROBINDO PHARMA LTD, QUTHUBULLAPUR  
MANDAL, ANDHRA PRADESH, INDIA

Packer: AUROBINDO PHARMA LTD, QUTHUBULLAPUR  
MANDAL, ANDHRA PRADESH, INDIA

Laboratory: FPRC: AUROBINDO PHARMA LTD, QUTHUBULLAPUR  
MANDAL, ANDHRA PRADESH, INDIA

FPRR: AUROBINDO PHARMA, ROSEBANK, JOHANNESBURG

Shelf-life: 24 months

Date of registration: 1 DECEMBER 2006

## MRF 15

Registration number: A40/1.2/0655  
 Name of medicine: AUROLIFT 100 mg  
 Dosage form: TABLET  
 Active ingredients: EACH TABLET CONTAINS:  
 SERTRALINE HYDROCHLORIDE EQUIVALENT TO  
 SERTRALINE 100,0 mg  
 Conditions of registration: 1, 2, 3, 4, 5, 6  
 Applicant: AUROBINDO PHARMA (PTY) LTD  
 Manufacturer: AUROBINDO PHARMA LTD, QUTHUBULLAPUR  
 MANDAL, ANDHRA PRADESH, INDIA  
 Packer: AUROBINDO PHARMA LTD, QUTHUBULLAPUR  
 MANDAL, ANDHRA PRADESH, INDIA  
 Laboratory: FPRC: AUROBINDO PHARMA LTD, QUTHUBULLAPUR  
 MANDAL, ANDHRA PRADESH, INDIA  
 FPRR: AUROBINDO PHARMA, ROSEBANK, JOHANNESBURG  
 Shelf-life: 24 months  
 Date of registration: 1 DECEMBER 2006

## MRF 15

Registration number: A40/20.2.3/0680  
 Name of medicine: CAPASTAT 1 g  
 Dosage form: INJECTION  
 Active ingredients: EACH VIAL CONTAINS:  
 CAPREOMYCIN SULPHATE EQUIVALENT TO  
 CAPREOMYCIN 1,0 g  
 Conditions of registration: 1, 2, 3, 4, 5, 6  
 Applicant: PHARMACARE LIMITED  
 Manufacturer: TEVA PHARMACEUTICAL WORKS CO LTD, TANCSICS  
 M, HUNGARY  
 Packer: TEVA PHARMACEUTICAL WORKS CO LTD, TANCSICS  
 M, HUNGARY  
 PHARMACARE LTD, KORSTEN, PORT ELIZABETH  
 ASPEN PHARMACARE, WILSONIA, EAST LONDON  
 Laboratory: FPRC: ELI LILLY ITALIA S.p.A, SESTO FIORENTINO, ITALY  
 SOUTH AFRICAN BUREAU OF STANDARDS,  
 GROENKLOOF, PRETORIA  
 RESEARCH INSTITUTE FOR INDUSTRIAL PHARMACY,  
 UNIVERSITY, POTCHEFSTROOM  
 INSPECTORATE M&L, ORMONDE, JOHANNESBURG  
 FPRC/FPRR: PHARMACARE LTD, KORSTEN, PORT ELIZABETH  
 ASPEN PHARMACARE, WILSONIA, EAST LONDON  
 Shelf-life: 24 months  
 Date of registration: 17 NOVEMBER

**MRF 15**

Registration number: A40/7.1.3/0682  
Name of medicine: SANDOZ RAMIPRIL 10  
Dosage form: TABLET  
Active ingredients: EACH TABLET CONTAINS:  
RAMIPRIL 10,0 mg  
Conditions of registration: 1, 2, 3, 4, 5, 6  
Applicant: SANDOZ (PTY) LTD  
Manufacturer: NOVARTIS BANGLADESH LTD, GAZIPUR, BANGLADESH  
Packer: NOVARTIS BANGLADESH LTD, GAZIPUR, BANGLADESH  
NOVARTIS S.A., SPARTAN, KEMPTON PARK  
Laboratory: FPRC: NOVARTIS BANGLADESH LTD, GAZIPUR, BANGLADESH  
SOUTH AFRICAN BUREAU OF STANDARDS,  
GROENKLOOF, PRETORIA  
ANALYTICON, TERENURE, KEMPTON PARK  
NOVARTIS S.A., SPARTAN, KEMPTON PARK  
FPRR: SANDOZ, SPARTAN, KEMPTON PARK  
Shelf-life: 24 months (provisional)  
Date of registration: 1 DECEMBER 2006

**MRF 15**

Registration number: A40/7.1.3/0683  
Name of medicine: RAMIPRIL-HEXAL 20  
Dosage form: TABLET  
Active ingredients: EACH TABLET CONTAINS:  
RAMIPRIL 20,0 mg  
Conditions of registration: 1, 2, 3, 4, 5, 6  
Applicant: SANDOZ (PTY) LTD  
Manufacturer: NOVARTIS BANGLADESH LTD, GAZIPUR,  
BANGLADESH  
Packer: NOVARTIS BANGLADESH LTD, GAZIPUR,  
BANGLADESH  
NOVARTIS S.A., SPARTAN, KEMPTON PARK  
Laboratory: FPRC: NOVARTIS BANGLADESH LTD, GAZIPUR,  
BANGLADESH  
SOUTH AFRICAN BUREAU OF STANDARDS,  
GROENKLOOF, PRETORIA  
ANALYTICON, TERENURE, KEMPTON PARK  
NOVARTIS S.A., SPARTAN, KEMPTON PARK  
FPRR: SANDOZ, SPARTAN, KEMPTON PARK  
Shelf-life: 24 months (provisional)  
Date of registration: 1 DECEMBER 2006

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**MRF 15**

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Registration number: A40/7.1.3/0684  
Name of medicine: RETACE 10  
Dosage form: TABLET  
Active ingredients: EACH TABLET CONTAINS:  
RAMIPRIL 10,0 mg  
Conditions of registration: 1, 2, 3, 4, 5, 6  
Applicant: SANDOZ (PTY) LTD  
Manufacturer: NOVARTIS BANGLADESH LTD, GAZIPUR, BANGLADESH  
Packer: NOVARTIS BANGLADESH LTD, GAZIPUR, BANGLADESH  
NOVARTIS S.A., SPARTAN, KEMPTON PARK  
Laboratory: FPRC: NOVARTIS BANGLADESH LTD, GAZIPUR, BANGLADESH  
SOUTH AFRICAN BUREAU OF STANDARDS,  
GROENKLOOF, PRETORIA  
ANALYTICON, TERENURE, KEMPTON PARK  
NOVARTIS S.A., SPARTAN, KEMPTON PARK  
FPRR: SANDOZ, SPARTAN, KEMPTON PARK  
Shelf-life: 24 months (provisional)  
Date of registration: 1 DECEMBER 2006

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