
GENERAL NOTICE ALGEMENE KENNISGEWING

NOTICE 351 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.
 - a. 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.

Registration number:	12/3, 1/15	Registration number:	A05/21, 1/02
Name of medicine:	NOROCARP INJECTION FOR DOGS	Name of medicine:	DRAXXIN 100 mg/ml
Dosage form:	INJECTION	Dosage form:	INJECTION
Active ingredients:	EACH 1,0 ml SOLUTION CONTAINS: CARPROFEN 50,0 mg	Active ingredients:	EACH 1,0 ml SOLUTION CONTAINS: TULATHROMYCIN 100,0 mg
Conditions of registration:	1, 2, 3, 4, 5, 6	Conditions of registration:	1, 2, 3, 4, 5, 6
Applicant:	NORBROOK LABORATORIES S.A. (PTY) LTD	Applicant:	PFIZER LABORATORIES (PTY) LTD
Manufacturer:	NORBROOK LABORATORIES LTD, NEWRY, NORTHERN IRELAND	Manufacturer:	PFIZER GLOBAL MANUFACTURING, AMBOISE CEDEX, FRANCE
Packer:	NORBROOK LABORATORIES LTD, NEWRY, NORTHERN IRELAND	Packer:	PFIZER GLOBAL MANUFACTURING, AMBOISE CEDEX, FRANCE
Laboratory: FPRC:	NORBROOK LABORATORIES LTD, NEWRY, NORTHERN IRELAND	Laboratory: FPRC:	PFIZER GLOBAL MANUFACTURING, AMBOISE CEDEX, FRANCE
FPRR:	NORBROOK LABORATORIES, CENTURION, RSA		SOUTH AFRICAN BUREAU OF STANDARDS, GROENKLOOF, PRETORIA
Shelf-life:	24 months	FPRR:	PFIZER LABORATORIES, SANDTON, RSA
Date of registration:	2 FEBRUARY 2007	Shelf-life:	24 months
		Date of registration:	1 DECEMBER 2006

Registration number:	38/16.4/0021	Registration number:	A39/20.2.8/0343
Name of medicine:	TEETH AND GUM DEFENCE LISTERINE	Name of medicine:	APEX-ACYCLOVIR 200 mg
Dosage form:	SOLUTION	Dosage form:	TABLET
Active ingredients:	EACH 20,0 ml SOLUTION CONTAINS: SODIUM FLUORIDE 4,420 mg THYMOL 12,780 mg ALCOHOL (95 %) 4,540 ml	Active ingredients:	EACH TABLET CONTAINS: ACYCLOVIR 200,0 mg
Conditions of registration:	1, 2, 3, 4, 5, 6	Conditions of registration:	1, 2, 3, 4, 5, 6
Applicant:	PFIZER LABORATORIES (PTY) LTD	Applicant:	CAMOX PHARMACEUTICALS (PTY) LTD
Manufacturer:	PFIZER GLOBAL MANUFACTURING, RETREAT CAPE TOWN	Manufacturer:	MEDREICH STERILAB LTD, VIRGONAR BANGALORE, INDIA
Packer:	PFIZER GLOBAL MANUFACTURING, RETREAT CAPE TOWN	Packer:	MEDREICH STERILAB LTD, VIRGONAR BANGALORE, INDIA
Laboratory: FPRC/FPRR:	PFIZER GLOBAL MANUFACTURING, RETREAT CAPE TOWN	Laboratory: FPRC:	MEDREICH STERILAB LTD, VIRGONAR BANGALORE, INDIA SOUTH AFRICAN BUREAU OF STANDARDS, GROENKLOOF, PRETORIA INSTITUTE FOR PHARMACEUTICAL SERVICES, SILVERTONDALE, RSA INSPECTORATE M&L, ORMONDE, JOHANNESBURG
Shelf-life:	24 months (provisional)	FPRR	CAMOX PHARMACEUTICALS, AMALGAM, JOHANNESBURG, RSA
Date of registration	1 DECEMBER 2006	Shelf-life:	24 months (provisional)
		Date of registration:	1 DECEMBER 2006

Registration number:	A39/20.2.8/0344	Registration number:	A40/20.2.8/0261
Name of medicine:	APEX-ACYCLOVIR 400 mg	Name of medicine:	CIPLA DUOVIR AND NEVIRAPINE CO-PACK
Dosage form:	TABLET	Dosage form:	TABLET
Active ingredients:	EACH TABLET CONTAINS: ACYCLOVIR S: 400,0 mg	Active ingredients:	EACH CARTON CONTAINS: DUOVIR TABLETS CONTAINING: LAMIVUDINE 150,0 mg ZIDOVUDINE 300,0 mg NEVIRAPINE TABLETS CONTAINING: NEVIRAPINE 200,0 mg
Conditions of registration:	1, 2, 3, 4, 5, 6	Conditions of registration:	1, 2, 3, 4, 5, 6
Applicant:	CAMOX PHARMACEUTICALS (PTY) LTD	Applicant:	CIPLA LIFE SCIENCES (PTY) LTD
Manufacturer:	MEDREICH STERILAB LTD, VIRGONAR BANGALORE, INDIA	Manufacturer:	CIPLA LTD, VIKHROLI, MUMBAI, INDIA
Packer:	MEDREICH STERILAB LTD, VIRGONAR BANGALORE, INDIA	Packer:	CIPLA LTD, VIKHROLI, MUMBAI, INDIA
Laboratory: FPRC:	MEDREICH STERILAB LTD, VIRGONAR BANGALORE, INDIA SOUTH AFRICAN BUREAU OF STANDARDS, GROENKLOOF, PRETORIA INSTITUTE FOR PHARMACEUTICAL SERVICES, SILVERTONDALE, RSA INSPECTORATE M&L, ORMONDE, JOHANNESBURG	Laboratory: FPRC:	CIPLA LTD, VIKHROLI, MUMBAI, INDIA
FPRR:	CAMOX PHARMACEUTICALS, AMALGAM, JOHANNESBURG, RSA	FPRR:	CIPLA LIFE SCIENCES, ROSEN PARK, BELLVILLE, RSA
Shelf-life:	24 months (provisional)	Shelf-life:	24 months
Date of registration:	1 DECEMBER 2006	Date of registration:	2 FEBRUARY 2007

MRF 15		MRF 15	
Registration number:	A40/7.1/0284	Registration number:	A40/20.1.2/0329
Name of medicine:	INDO AMLODIPINE-5	Name of medicine:	INDO AMOXICYCLIN 250
Dosage form:	TABLET	Dosage form:	CAPSULE
Active ingredients:	EACH TABLET CONTAINS: AMLODIPINE BESYLATE EQUIVALENT TO AMLODIPINE 5.0 mg	Active ingredients:	EACH CAPSULE CONTAINS: AMOXICYCLIN TRIHYDRATE EQUIVALENT TO AMOXICYCLIN 250.0 mg
Conditions of registration:	1 2, 3, 4, 5, 6	Conditions of registration:	1 2, 3, 4, 5, 6
Applicant:	DEZZO TRADING (392) (PTY) LTD t/a INDO PHARMA	Applicant:	DEZZO TRADING (392) (PTY) LTD t/a INDO PHARMA
Manufacturer:	KOPRAN LTD, KHALAPUR, RAIGAD, INDIA	Manufacturer:	KOPRAN LTD, KHALAPUR, RAIGAD, INDIA
Patent:	KOPRAN LTD, KHALAPUR, RAIGAD, INDIA	Patent:	KOPRAN LTD, KHALAPUR, RAIGAD, INDIA
Laboratory: FPRC:	KOPRAN LTD, KHALAPUR, RAIGAD, INDIA CONSULTING CHEMICAL LABORATORIES, STAR STREET, BOKSBURG, RSA	Laboratory: FPRC:	KOPRAN LTD, KHALAPUR, RAIGAD, INDIA
FPRR:	DEZZO TRADING (392) t/a INDO PHARMA, ANCHORVILLE, LENASIA, JOHANNESBURG, RSA	FPRR:	DEZZO TRADING (392) t/a INDO PHARMA, ANCHORVILLE, LENASIA, JOHANNESBURG, RSA
Shelf-life:	24 months (provisional)	Shelf-life:	36 months
Date of registration:	2 FEBRUARY 2007	Date of registration:	2 FEBRUARY 2007

MRF 15

Registration number:	A40/20.1.2/0330
Name of medicine	INDO AMOXYCILLIN 500
Dosage form	CAPSULE
Active ingredients:	EACH CAPSULE CONTAINS: AMOXYCILLIN TRIHYDRATE EQUIVALENT TO AMOXYCILLIN 500,0 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
FPRR:	DEZZO TRADING (392) t/a INDO PHARMA, ANCHORVILLE, LENASIA, JOHANNESBURG, RSA
Shelf-life:	36 months
Date of registration:	2 FEBRUARY 2007

MDE 15

Registration number:	A40/7.1/0435
Name of medicine:	AMLOBLOC 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
Applicant:	SANDOZ (PTY) LTD
Manufacturer:	LEK PHARMACEUTICALS d.d., VEROVSKOVA, SLOVENIA
Packer:	LEK PHARMACEUTICALS d.d., VEROVSKOVA, SLOVENIA
Laboratory:	FPRC: NOVARTIS, SPARTAN, KEMPTON PARK LEK PHARMACEUTICALS d.d., VEROVSKOVA, SLOVENIA
FPRR:	NOVARTIS, SPARTAN, KEMPTON PARK SOUTH AFRICAN BUREAU OF STANDARDS, GROENKLOOF, PRETORIA
Shelf-life:	FPRR: ANALYTICON, TERENURE, KEMPTON PARK SANDOZ, SPARTAN, KEMPTON PARK 24 months (provisional)
Date of registration	2 FEBRUARY 2007

MRF 15	MDE 15
Registration number: A40.7.10036	Registration number: A40/20.2.8/0460
Name of medicine: AMLOBLOC 10	TRIOMUNE 30
Dosage form: TABLET	TABLET
Active ingredients: EACH TABLET CONTAINS: AMLODIPINE BESYLATE EQUIVALENT TO AMLODIPINE 10,0 mg	EACH TABLET CONTAINS: STAVUDINE 30,0 mg LAMIVUDINE 150,0 mg NEVIRAPINE 200,0 mg
Conditions of registration: 1, 2, 3, 4, 5, 6	1, 2, 3, 4, 5, 6
Applicant: SANDOZ (PTY) LTD	CIPLA MEDPRO (PTY)
Manufacturer: LEK PHARMACEUTICALS d.d., VEROVSKOVA, SLOVENIA	CIPLA LTD, PATALGANGA, MAHARASHTRA, INDIA
Packer: LEK PHARMACEUTICALS d.d., VEROVSKOVA, SLOVENIA	CIPLA LTD, PATALGANGA, MAHARASHTRA, INDIA
La berritory: FPRC NOVARTIS, SPARTAN, KEMPTON PARK	CIPLA LTD, PATALGANGA, MAHARASHTRA, INDIA ELI LILLY & CO, BASINGSTOKE, HAMPSHIRE UK
FPRC: LEK PHARMACEUTICALS d.d., VEROVSKOVA, SLOVENIA	CIPLA LTD, PATALGANGA, MAHARASHTRA, INDIA ELI LILLY & CO, BASINGSTOKE, HAMPSHIRE UK
Shelf-life: NOVARTIS, SPARTAN, KEMPTON PARK	CIPLA LTD, PATALGANGA, MAHARASHTRA, INDIA ELI LILLY & CO, BASINGSTOKE, HAMPSHIRE UK
Date of registration: SOUTH AFRICAN BUREAU OF STANDARDS, GROENKLOOF, PRETORIA	CIPLA LTD, PATALGANGA, MAHARASHTRA, INDIA ELI LILLY & CO, BASINGSTOKE, HAMPSHIRE UK
ANALYTICON, TERENURE, KEMPTON PARK	CIPLA LTD, PATALGANGA, MAHARASHTRA, INDIA ELI LILLY & CO, BASINGSTOKE, HAMPSHIRE UK
SANDOZ, SPARTAN, KEMPTON PARK	CIPLA LTD, PATALGANGA, MAHARASHTRA, INDIA ELI LILLY & CO, BASINGSTOKE, HAMPSHIRE UK
24 months (provisional)	CIPLA LTD, PATALGANGA, MAHARASHTRA, INDIA ELI LILLY & CO, BASINGSTOKE, HAMPSHIRE UK
2 FEBRUARY 2007	CIPLA LTD, PATALGANGA, MAHARASHTRA, INDIA ELI LILLY & CO, BASINGSTOKE, HAMPSHIRE UK

MRF 15	A40/21.2/0466	Registration number:	A40/21.2/0467
	INDO METFORMIN-500	Name of medicine:	INDO METFORMIN-850
	TABLET	Dosage form:	TABLET
	EACH TABLET CONTAINS: METFORMIN HYDROCHLORIDE 500,0 mg	Active ingredients:	EACH TABLET CONTAINS: METFORMIN HYDROCHLORIDE 850,0 mg
	1, 2, 3, 4, 5, 6	Conditions of registration:	1, 2, 3, 4, 5, 6
	DEZZO TRADING (392) (PTY) LTD t/a INDO PHARMA	Applicant:	DEZZO TRADING (392) (PTY) LTD t/a INDO PHARMA
	RUSAN PHARMA LTD, GANDHIDHAM-KUTCH, INDIA	Manufacturer:	RUSAN PHARMA LTD, GANDHIDHAM-KUTCH, INDIA
	RUSAN PHARMA LTD, GANDHIDHAM-KUTCH, INDIA	Packer:	RUSAN PHARMA LTD, GANDHIDHAM-KUTCH, INDIA
	RUSAN PHARMA LTD, GANDHIDHAM-KUTCH, INDIA	Laboratory: FPRC	RUSAN PHARMA LTD, GANDHIDHAM-KUTCH, INDIA
	CONSULTING CHEMICAL LABORATORIES, ATLASVILLE, BOKSBURG, RSA		CONSULTING CHEMICAL LABORATORIES, ATLASVILLE, BOKSBURG, RSA
	DEZZO TRADING (392) t/a INDO PHARMA, ANCHORVILLE, LENASIA, JOHANNESBURG, RSA	FPRR:	DEZZO TRADING (392) t/a INDO PHARMA, ANCHORVILLE, LENASIA, JOHANNESBURG, RSA
	24 months (provisional)	Shelf-life:	24 months (provisional)
	2 FEBRUARY 2007	Date of registration:	2 FEBRUARY 2007

MRF 15

Registration number:	A40/7.1.3/0722
Name of medicine:	AURO-LISINOPRIL 5 mg
Dosage form:	TABLET
Active ingredients:	EACH TABLET CONTAINS: LISINOPRIL DIHYDRATE EQUIVALENT TO LISINOPRIL 5.0 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
Shelf-life:	FPRC: AUROBINDO PHARMA, ROSEBANK, JOHANNESBURG
Date of registration:	24 months (provisional) 2 FEBRUARY 2007

MRF 15

Registration number:	A40/7.1.3/0723
Name of medicine:	AURO-LISINOPRIL 10 mg
Dosage form:	TABLET
Active ingredients:	EACH TABLET CONTAINS: LISINOPRIL DIHYDRATE EQUIVALENT TO LISINOPRIL 10.0 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
Shelf-life:	FPRC: AUROBINDO PHARMA, ROSEBANK, JOHANNESBURG
Date of registration:	24 months (provisional) 2 FEBRUARY 2007

MRF 15	MDE 15
Registration number:	41/20.2.8/0235
Name of medicine:	ADCO-NEVIRAPINE TABLETS
Dosage form:	TABLET
Active ingredients:	EACH TABLET CONTAINS: NEVIRAPINE 200,0 mg
Conditions of registration:	1, 2, 3, 4, 5, 6
Applicant:	ADCOCK INGRAM LIMITED
Manufacturer:	ADCOCK INGRAM HEALTHCARE, WADEVILLE, GERMISTON
Packer:	ADCOCK INGRAM HEALTHCARE, WADEVILLE, GERMISTON ADCOCK INGRAM LTD, BRYANSTON, JOHANNESBURG
Laboratory:	FPRC: FPRC/FPRR ADCOCK INGRAM HEALTHCARE, WADEVILLE, GERMISTON ADCOCK INGRAM LTD, BRYANSTON, JOHANNESBURG
Shelf-life:	24 months (provisional)
Date of registration:	2 FEBRUARY 2007

Registration number:	A39/20.1.1/0303
Name of medicine:	APEX-CEFTRIAXONE 1 g
Dosage form:	INJECTION
Active ingredients:	EACH VIAL CONTAINS: CEFTRIAXONE SODIUM EQUIVALENT TO CEFTRIAXONE 1,0 g
Conditions of registration:	1, 2, 3, 4, 5, 6
Applicant:	CAMOX PHARMACEUTICALS (PTY) LTD
Manufacturer:	MJ BIOPHARM, RAIGAD, MAHARASHTRA, INDIA
Packer:	MJ BIOPHARM, RAIGAD, MAHARASHTRA, INDIA
Laboratory:	FPRC: MJ BIOPHARM, RAIGAD, MAHARASHTRA, INDIA IPCA LABORATORIES, ATHAL, DADRA & NAGAR HAVELI, INDIA SOUTH AFRICAN BUREAU OF STANDARDS, GROENKLOOF, PRETORIA INSPECTORATE M&L, ORMONDE, JOHANNESBURG INSTITUTE FOR PHARMACEUTICAL SERVICES, SILVERTONDALE
FPRR:	CAMOX PHARMACEUTICALS, AMALGAM, JOHANNESBURG
Shelf-life:	24 months
Date of registration:	1 DECEMBER 2006

MRF 15

Registration number:	41/20.2.8/0236
Name of medicine:	ADCO-LAMIVUDINE TABLETS
Dosage form:	TABLET
Active ingredients:	EACH TABLET CONTAINS: LAMIVUDINE 150,0 mg
Conditions of registration:	1, 2, 3, 4, 5, 6
Applicant:	ADCOCK INGRAM LIMITED
Manufacturer:	ADCOCK INGRAM HEALTHCARE, WADEVILLE, GERMISTON
Packer:	ADCOCK INGRAM HEALTHCARE, WADEVILLE, GERMISTON ADCOCK INGRAM LTD, BRYANSTON, JOHANNESBURG
Laboratory:	FPRC/FPRR ADCOCK INGRAM HEALTHCARE, WADEVILLE, GERMISTON ADCOCK INGRAM LTD, BRYANSTON, JOHANNESBURG
Shelf-life:	24 months (provisional)
Date of registration:	2 FEBRUARY 2007

Registration number:	41/20.2.8/0237
Name of medicine:	ADCO-ZIDOVUDINE TABLETS
Dosage form:	TABLET
Active ingredients:	EACH TABLET CONTAINS: ZIDOVUDINE 300,0 mg
Conditions of registration:	1, 2, 3, 4, 5, 6
Applicant:	ADCOCK INGRAM LIMITED
Manufacturer:	ADCOCK INGRAM HEALTHCARE, WADEVILLE, GERMISTON
Packer:	ADCOCK INGRAM HEALTHCARE, WADEVILLE, GERMISTON ADCOCK INGRAM LTD, BRYANSTON, JOHANNESBURG
Laboratory:	FPRC/FPRR ADCOCK INGRAM HEALTHCARE, WADEVILLE, GERMISTON ADCOCK INGRAM LTD, BRYANSTON, JOHANNESBURG
Shelf-life:	24 months (provisional)
Date of registration:	2 FEBRUARY 2007

MDE 15	
Registration number:	41/7.3/0092
Name of medicine:	CIPLA-SUMATRIPTAN 100
Dosage form:	TABLET
Active ingredients:	EACH TABLET CONTAINS: SUMATRIPTAN SUCCINATE EQUIVALENT TO SUMATRIPTAN 100,0 mg
Conditions of registration:	1, 2, 3, 4, 5, 6
Applicant:	CIPLA LIFE SCIENCES (PTY) LTD
Manufacturer:	CIPLA LTD, UNIT III, VERNA, GOA, INDIA
Packer:	CIPLA LTD, UNIT III, VERNA, GOA, INDIA
Library:	FPRC
FPRR:	CIPLA LIFE SCIENCES, ROSENPARK, BELLVILLE
Shelf-life:	24 months
Date of registration:	2 MARCH 2007

MDE 15	
Registration number:	A40/7.1.3/0724
Name of medicine:	AURO-LISINOPRIL 20 mg
Dosage form:	TABLET
Active ingredients:	EACH TABLET CONTAINS: LISINOPRIL DIHYDRATE EQUIVALENT TO LISINOPRIL 20,0 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
Library:	FPRC
FPRR:	AUROBINDO PHARMA, ROSEBANK, JOHANNESBURG
Shelf-life:	24 months (provisional)
Date of registration:	2 FEBRUARY 2007

MRF 15

Registration number: 37/34/0528

Name of medicine: NORBROOK WATER FOR INJECTION

Dosage form: INJECTION

Active ingredients: EACH VIAL CONTAINS:
WATER FOR INJECTIONS 10 ml

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

Applicant: NORBROOK LABORATORIES S.A. (PTY) LTD

Manufacturer: NORBROOK LABORATORIES LTD, NEWRY,
NORTHERN IRELAND

Packer: NORBROOK LABORATORIES LTD, NEWRY,
NORTHERN IRELAND

Laboratory: FPRC NORBROOK LABORATORIES LTD, NEWRY,
NORTHERN IRELAND

FPRC NORBROOK LABORATORIES, CENTURION, RSA

Shelf-life: 36 months

Date of registration: 2 MARCH 2007

MRF 15

Registration number: 38/13.1/0195

Name of medicine: MDI POVIDONE GEL

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 PHARMACEUTICAL SERVICES,
SILVERTONDALE, RSA

FPRC: MDI cc, MENLO PARK, PRETORIA

Shelf-life: 24 months (provisional)

Date of registration: 2 MARCH 2007

MRF 15

Registration number:	A38/30.1/0525
Name of medicine:	PENTAXIM
Dosage form:	INJECTION
Active ingredients:	EACH 0,5 ml DOSE CONTAINS: PURIFIED DIPHTHERIA TOXOID 30,0 iu PURIFIED TETANUS TOXOID 40,0 iu PURIFIED PERTUSSIS TOXOID 25,0 ug PURIFIED PERTUSSIS FILAMENTOUS HAEMAGGLUTININ 25,0 ug POLIOVIRUS D ANTIGEN TYPE 1 40,0 D units POLIOVIRUS D ANTIGEN TYPE 2 8,0 D units POLIOVIRUS D ANTIGEN TYPE 3 32,0 D units HAEMOPHILUS INFLUENZA TYPE b POLYSACCHARIDE 100 ug
Conditions of registration:	1, 2, 3, 4, 5, 6
Applicant:	AVENTIS PHARMA (PTY) LTD
Manufacturer:	AVENTIS PASTEUR, L'ETOILE, FRANCE
Packer:	AVENTIS PASTEUR, L'ETOILE, FRANCE
Laboratory:	FPRC AVENTIS PASTEUR, L'ETOILE, FRANCE
Shelf-life:	FPRR: AVENTIS PHARMA, WALTLOO, PRETORIA 36 months
Date of registration:	2 MARCH 2007

MRF 15

Registration number:	A38/30.1/0526
Name of medicine:	TETRAXIM
Dosage form:	INJECTION
Active ingredients:	EACH 0,5 ml DOSE CONTAINS: PURIFIED DIPHTHERIA TOXOID 30,0 iu PURIFIED TETANUS TOXOID 40,0 iu PURIFIED PERTUSSIS TOXOID 25,0 ug PURIFIED PERTUSSIS FILAMENTOUS HAEMAGGLUTININ 25,0 ug POLIOVIRUS D ANTIGEN TYPE 1 40,0 D units POLIOVIRUS D ANTIGEN TYPE 2 8,0 D units POLIOVIRUS D ANTIGEN TYPE 3 32,0 D units
Conditions of registration:	1, 2, 3, 4, 5, 6
Applicant:	AVENTIS PHARMA (PTY) LTD
Manufacturer:	AVENTIS PASTEUR, L'ETOILE, FRANCE
Packer:	AVENTIS PASTEUR, L'ETOILE, FRANCE
Laboratory:	FPRC AVENTIS PASTEUR, L'ETOILE, FRANCE
Shelf-life:	FPRR: AVENTIS PHARMA, WALTLOO, PRETORIA 36 months
Date of registration:	2 MARCH 2007

MRF 15		MRF 15	
Registration number:	A39/20.1.7/0383	Registration number:	A39/10.2.1/0482
Name of medicine:	MERCK-ITRACONAZOLE 100 mg	Name of medicine:	ASTHAVENT RESPULES
Dosage form:	CAPSULE	Dosage form:	SOLUTION
Active ingredients:	EACH CAPSULE CONTAINS: ITRACONAZOLE 100.0 mg	Active ingredients:	EACH 2.5 ml SOLUTION CONTAINS: SALBUTAMOL SULPHATE EQUIVALENT TO SALBUTAMOL 2.5 mg
Conditions of registration:	1, 2, 3, 4, 5, 6	Conditions of registration:	1, 2, 3, 4, 5, 6
Applicant:	MERCK GENERICS RSA (PTY) LTD	Applicant:	CIPLA-MEDPRO (PTY) LTD
Manufacturer:	MARTEC PHARMACEUTICAL INC, TOPPING, KANSAS CITY, USA	Manufacturer:	CIPLA LTD, UNIT I, VERNA, GOA, INDIA
Packer:	MARTEC PHARMACEUTICAL INC, TOPPING, KANSAS CITY, USA GENERICS (UK) LTD, POTTERS BAR, HERTFORDSHIRE, UK GERARD LABORATORIES, DUBLIN, IRELAND MERCK PHARMACEUTICAL MANUFACTURING, WADEVILLE, GERMISTON	Packer:	CIPLA LTD, UNIT I, VERNA, GOA, INDIA
Laboratory:	MARTEC PHARMACEUTICAL INC, TOPPING, KANSAS CITY, USA GENERICS (UK) LTD, POTTERS BAR, HERTFORDSHIRE, UK GERARD LABORATORIES, DUBLIN, IRELAND MERCK PHARMACEUTICAL MANUFACTURING, WADEVILLE, GERMISTON RESEARCH INSTITUTE FOR INDUSTRIAL PHARMACY, NORTH-WEST UNIVERSITY, POTCHEFSTROOM SOUTH AFRICAN BUREAU OF STANDARDS	Laboratory:	CIPLA LTD, UNIT I, VERNA, GOA, INDIA
FPRC:		FPRC:	
Shelf-life:	24 months	Shelf-life:	24 months
Date of registration:	2 MARCH 2007	Date of registration:	2 MARCH 2007

MRF 15	
Registration number:	A39/10.2.1/0483
Name of medicine:	CIPLA-SALBUTAMOL RESPULES
Dosage form:	SOLUTION
Active ingredients:	EACH 2.5 ml SOLUTION CONTAINS: SALBUTAMOL 2.5 mg
Conditions of registration:	1, 2, 3, 4, 5, 6
Applicant:	CIPLA LIFE SCIENCES (PTY) LTD
Manufacturer:	CIPLA LTD, UNIT I, VERNA, GOA, INDIA
Packer:	CIPLA LTD, UNIT I, VERNA, GOA, INDIA
Laboratory:	FPRC: CIPLA LTD, UNIT I, VERNA, GOA, INDIA
	FPRR: CIPLA LIFE SCIENCES, ROSENPARK, BELLVILLE PHARMA DYNAMICS, SILVERWOOD, WESTLAKE, RSA
Shelf-life:	24 months
Date of registration:	2 MARCH 2007

MRF 15	
Registration number:	A39/8.4/0518
Name of medicine:	VENOFUNDIN
Dosage form:	SOLUTION
Active ingredients:	EACH 1000.0 ml SOLUTION CONTAINS: HETASTARCH 60.0 g SODIUM CHLORIDE 9.0 g
Conditions of registration:	1, 2, 3, 4, 5, 6
Applicant:	B BRAUN MEDICAL (PTY) LTD
Manufacturer:	B BRAUN MEDICAL AG, CRISSIER, SWITZERLAND
Packer:	B BRAUN MEDICAL AG, CRISSIER, SWITZERLAND
Laboratory:	FPRC: B BRAUN MEDICAL AG, CRISSIER, SWITZERLAND CONSULTING CHEMICAL LABORATORIES, ATLASVILLE, BOKSBURG
	FPRR: B BRAUN MEDICAL, HONEYDEW, GAUTENG
Shelf-life:	24 months
Date of registration:	2 MARCH 2007

MRF 15		MRF 15	
Registration number:	A397.1/0546	Registration number:	A397.1/0545
Name of medicine:	AUSTELL-AMLODIPINE 10 mg	Name of medicine:	AUSTELL-AMLODIPINE 5 mg
Dosage form:	TABLET	Dosage form:	TABLET
Active ingredients:	EACH TABLET CONTAINS: AMLODIPINE BESYLATE EQUIVALENT TO AMLODIPINE 10,0 mg	Active ingredients:	EACH TABLET CONTAINS: AMLODIPINE BESYLATE EQUIVALENT TO AMLODIPINE 5,0 mg
Conditions of registration:	1, 2, 3, 4, 5, 6	Conditions of registration:	1, 2, 3, 4, 5, 6
Applicant:	AUSTELL LABORATORIES (PTY) LTD	Applicant:	AUSTELL LABORATORIES (PTY) LTD
Manufacturer:	IPCA LABORATORIES LIMITED, SILVASSA, DADRA & NAGAR HAVELI, INDIA	Manufacturer:	IPCA LABORATORIES LIMITED, SILVASSA, & NAGAR HAVELI, INDIA
Packer:	IPCA LABORATORIES LIMITED, SILVASSA, DADRA & NAGAR HAVELI, INDIA	Packer:	IPCA LABORATORIES LIMITED, SILVASSA, & NAGAR HAVELI, INDIA
Laboratory:	IPCA LABORATORIES LIMITED, SILVASSA, DADRA & NAGAR HAVELI, INDIA SOUTH AFRICAN BUREAU OF STANDARDS, GROENKLOOF, PRETORIA INSPECTORATE M&L, ORMONDE, JOHANNESBURG INSTITUTE FOR PHARMACEUTICAL SERVICES, SILVERTONDALE, RSA	Laboratory:	IPCA LABORATORIES LIMITED, SILVASSA, DADRA & NAGAR HAVELI, INDIA SOUTH AFRICAN BUREAU OF STANDARDS, GROENKLOOF, PRETORIA INSPECTORATE M&L, ORMONDE, JOHANNESBURG INSTITUTE FOR PHARMACEUTICAL SERVICES, SILVERTONDALE, RSA
For:	AUSTELL LABORATORIES, SPRINGFIELD, JOHANNESBURG	For:	AUSTELL LABORATORIES, SPRINGFIELD, JOHANNESBURG
Shelf-life:	24 months (provisional)	Shelf-life:	24 months (provisional)
Date of registration:	2 MARCH 2007	Date of registration:	2 MARCH 2007

MED 15

Registration number:	A40/21.2/0151
Name of medicine:	ACTAZON 15
Dosage form:	TABLET
Active ingredients:	EACH TABLET CONTAINS: PIOGLITAZONE HYDROCHLORIDE EQUIVALENT TO PIOGLITAZONE 15,0 mg
Conditions of registration:	1, 2, 3, 4, 5, 6
Applicant:	CIPLA MEDPRO (PTY) LTD
Manufacturer:	CIPLA LTD, SALCETTE, GOA, INDIA
Packer:	CIPLA LTD, SALCETTE, GOA, INDIA
Laboratory:	FPRC: CIPLA LTD, SALCETTE, GOA, INDIA FPRR: CIPLA MEDPRO, ROSENPARK, BELLVILLE
Shelf-life:	24 months
Date of registration:	2 MARCH 2007

MED 4 E

Registration number:	A40/21.2/0152
Name of medicine:	ACTAZON 30
Dosage form:	TABLET
Active ingredients:	EACH TABLET CONTAINS: PIOGLITAZONE HYDROCHLORIDE EQUIVALENT TO PIOGLITAZONE 30,0 mg
Conditions of registration:	1, 2, 3, 4, 5, 6
Applicant:	CIPLA MEDPRO (PTY) LTD
Manufacturer:	CIPLA LTD, SALCETTE, GOA, INDIA
Packer:	CIPLA LTD, SALCETTE, GOA, INDIA
Laboratory:	FPRC: CIPLA LTD, SALCETTE, GOA, INDIA FPRR: CIPLA MEDPRO, ROSENPARK, BELLVILLE
Shelf-life:	24 months
Date of registration:	2 MARCH 2007

MRF 15	
Registration number:	A40/2.5/0168
Name of medicine:	PHARMA DYNAMICS LAMOTRIGINE 200 mg
Dosage form:	TABLET
Active ingredients:	EACH TABLET CONTAINS: LAMOTRIGINE 200,0 mg
Conditions of registration:	1, 2, 3, 4, 5, 6
Applicant:	PHARMA DYNAMICS (PTY) LTD
Manufacturer:	MEDOCHEMIE LTD, LIMASSOL, CYPRUS
Packer:	MEDOCHEMIE LTD, LIMASSOL, CYPRUS DIVPHARM MANUFACTURING & PACKAGING, LONGDALE, JOHANNESBURG TECHNIKON LABORATORIES, FLORIDA, RSA PHARMACEUTICAL ENTERPRISES, N'DABENI, KZN IMPILO DRUGS, ISITHEBE, KZN
Laboratory:	FPRC: MEDOCHEMIE LTD, LIMASSOL, CYPRUS CONSULTING CHEMICAL LABORATORIES, ATLASVILLE, BOKSBURG, RSA TECHNIKON LABORATORIES, FLORIDA, RSA IMPILO DRUGS, ISITHEBE, KZN
Shelf-life:	FPRR: PHARMA DYNAMICS, SILVERWOOD, WESTLAKE, RSA 24 months
Date of registration:	2 MARCH 2007

MRF 15	
Registration number:	A40/2.5/0170
Name of medicine:	PHARMA DYNAMICS LAMOTRIGINE 00 mg
Dosage form:	TABLET
Active ingredients:	EACH TABLET CONTAINS: LAMOTRIGINE 100,0 mg
Conditions of registration:	1, 2, 3, 4, 5, 6
Applicant:	PHARMA DYNAMICS (PTY) LTD
Manufacturer:	MEDOCHEMIE LTD, LIMASSOL, CYPRUS
Packer:	MEDOCHEMIE LTD, LIMASSOL, CYPRUS DIVPHARM MANUFACTURING & PACKAGING, LONGDALE, JOHANNESBURG TECHNIKON LABORATORIES, FLORIDA, RSA PHARMACEUTICAL ENTERPRISES, N'DABENI, KZN IMPILO DRUGS, ISITHEBE, KZN
Laboratory:	FPRC: MEDOCHEMIE LTD, LIMASSOL, CYPRUS CONSULTING CHEMICAL LABORATORIES, ATLASVILLE, BOKSBURG, RSA TECHNIKON LABORATORIES, FLORIDA, RSA IMPILO DRUGS, ISITHEBE, KZN
Shelf-life:	FPRR: PHARMA DYNAMICS, SILVERWOOD, WESTLAKE, RSA 24 months
Date of registration:	2 MARCH 2007

MRF 15		MRF 15	
Registration number:	A40/2.5/0171	Registration number:	A40/2.5/0172
Name of medicine:	PHARMA DYNAMICS LAMOTRIGINE 25 mg	Name of medicine:	PHARMA DYNAMICS LAMOTRIGINE 50 mg
Dosage form:	TABLET	Dosage form:	TABLET
Active ingredients:	EACH TABLET CONTAINS: LAMOTRIGINE 25,0 mg	Active ingredients:	EACH TABLET CONTAINS: LAMOTRIGINE 50,0 mg
Conditions of registration:	1, 2, 3, 4, 5, 6	Conditions of registration:	1, 2, 3, 4, 5, 6
Applicant:	PHARMA DYNAMICS (PTY) LTD	Applicant:	PHARMA DYNAMICS (PTY) LTD
Manufacturer:	MEDOCHÉMIE LTD, LIMASSOL, CYPRUS	Manufacturer:	MEDOCHÉMIE LTD, LIMASSOL, CYPRUS
Packer:	MEDOCHÉMIE LTD, LIMASSOL, CYPRUS DIVPHARM MANUFACTURING & PACKAGING, LONGDALE, JOHANNESBURG TECHNIKON LABORATORIES, FLORIDA, RSA PHARMACEUTICAL ENTERPRISES, N'DABENI, KZN IMPILO DRUGS, ISITHEBE, KZN	Manufacturer:	MEDOCHÉMIE LTD, LIMASSOL, CYPRUS DIVPHARM MANUFACTURING & PACKAGING, LONGDALE, JOHANNESBURG TECHNIKON LABORATORIES, FLORIDA, RSA PHARMACEUTICAL ENTERPRISES, N'DABENI, KZN IMPILO DRUGS, ISITHEBE, KZN
Laboratory:	FPRC: MEDOCHÉMIE LTD, LIMASSOL, CYPRUS CONSULTING CHEMICAL LABORATORIES, ATLASVILLE, BOKSBURG, RSA TECHNIKON LABORATORIES, FLORIDA, RSA IMPILO DRUGS, ISITHEBE, KZN	Laboratory:	FARC: MEDOCHÉMIE LTD, LIMASSOL, CYPRUS CONSULTING CHEMICAL LABORATORIES, ATLASVILLE, BOKSBURG, RSA TECHNIKON LABORATORIES, FLORIDA, RSA IMPILO DRUGS, ISITHEBE, KZN
Shelf-life:	24 months	Shelf-life:	24 months
Date of registration:	2 MARCH 2007	Date of registration:	2 MARCH 2007

MRF 15

Registration number:	A40/2.5/0183
Name of medicine:	ADCO-MICTRIN 200
Dosage form:	TABLET
Active ingredients:	EACH TABLET CONTAINS: LAMOTRIGINE 200.0 mg
Conditions of registration:	1, 2, 3, 4, 5, 6
Applicant:	ADCOCK INGRAM LIMITED
Manufacturer:	DELTA LIMITED, HAFNARFJORDUR, ICELAND
Packer:	DELTA LIMITED, HAFNARFJORDUR, ICELAND ADCOCK INGRAM HEALTHCARE, WADEVILLE, GERMISTON
Laboratory: FPRC:	DELTA LIMITED, HAFNARFJORDUR, ICELAND
FPRC/FPRR:	ADCOCK INGRAM HEALTHCARE, WADEVILLE, GERMISTON
Shelf-life:	24 months
Date of registration:	2 MARCH 2007

MRF 16

Registration number:	A40/2.5/0185
Name of medicine:	ADCO-MICTRIN 50
Dosage form:	TABLET
Active ingredients:	EACH TABLET CONTAINS: LAMOTRIGINE 50.0 mg
Conditions of registration:	1, 2, 3, 4, 5, 6
Applicant:	ADCOCK INGRAM LIMITED
Manufacturer:	DELTA LIMITED, HAFNARFJORDUR, ICELAND
Packer:	DELTA LIMITED, HAFNARFJORDUR, ICELAND ADCOCK INGRAM HEALTHCARE, WADEVILLE, GERMISTON
Laboratory: FPRC:	DELTA LIMITED, HAFNARFJORDUR, ICELAND
FPRC/FPRR:	ADCOCK INGRAM HEALTHCARE, WADEVILLE, GERMISTON
Shelf-life:	24 months
Date of registration:	2 MARCH 2007

MRF 15	
A40/2.5/0186	Registration number: A40/1.4.3/0247
ADCO-MICTRIN 100	Name of medicine: LANCAP 15 mg
TABLET	Dosage form: CAPSULE
EACH TABLET CONTAINS: LAMOTRIGINE 100,0 mg	Active ingredients: EACH CAPSULE CONTAINS: LANSOPRAZOLE 15,0 mg
1, 2, 3, 4, 5, 6	Conditions of registration: 1, 2, 3, 4, 5, 6
ADCOCK INGRAM LIMITED	Applicant: PHARMA DYNAMICS (PTY) LTD
DELTA LIMITED, HAFNARFJORDUR, ICELAND	Manufacturer: KRKA DD, NOVO MESTO, SLOVENIA
ADCOCK INGRAM LIMITED, HAFNARFJORDUR, ICELAND	Packer: KRKA DD, NOVO MESTO, SLOVENIA
ADCOCK INGRAM HEALTHCARE, WADEVILLE, TON	DIVPHARM MANUFACTURING & PACKAGING, LONGDALE, JOHANNESBURG
	TECHNIKON LABORATORIES, FLORIDA, RSA
	PHARMACEUTICAL ENTERPRISES, N'DABENI, KZN
	IMPILO DRUGS, ISITHEBE, KZN
DELTA LIMITED, HAFNARFJORDUR, ICELAND	Laboratory: KRKA DD, NOVO MESTO, SLOVENIA
	FPRC: CONSULTING CHEMICAL LABORATORIES, ATLASSVILLE, BOKSBURG, RSA
ADCOCK INGRAM HEALTHCARE, WADEVILLE, GERMISTON	FPRC/FPRR: TECHNIKON LABORATORIES, FLORIDA, RSA
24 months	IMPILO DRUGS, ISITHEBE, KZN
2 MARCH 2007	
	Shelf-life: PHARMA DYNAMICS, SILVERWOOD, WESTLAKE, RSA
	24 months
	Date of registration: 2 MARCH 2007

MRF 15		MRF 15	
Registration number:	A40/11.4.3/0248	Registration number:	A40/27/0266
Name of medicine:	LANCAP 30 mg	Name of medicine:	EXJADE 125 mg
Dosage form:	CAPSULE	Dosage form:	TABLET
Active ingredients:	EACH CAPSULE CONTAINS: LANSOPRAZOLE 30,0 mg	Active ingredients:	EACH TABLET CONTAINS: DEFERASIROX 125,0 mg
Conditions of registration:	1, 2, 3, 4, 5, 6	Conditions of registration:	1, 2, 3, 4, 5, 6
Applicant:	PHARMA DYNAMICS (PTY) LTD	Applicant:	NOVARTIS SOUTH AFRICA (PTY) LTD
Manufacturer:	KRKA DD, NOVO MESTO, SLOVENIA	Manufacturer:	NOVARTIS PHARMA STEIN AG, STEIN, SWITZERLAND
Packer:	KRKA DD, NOVO MESTO, SLOVENIA DIVPHARM MANUFACTURING & PACKAGING, LONGDALE, JOHANNESBURG TECHNIKON LABORATORIES, FLORIDA, RSA PHARMACEUTICAL ENTERPRISES, N'DABENI, KZN	Importer:	NOVARTIS PHARMA STEIN AG, STEIN, SWITZERLAND
Laboratory:	FPRC: KRKA DD, NOVO MESTO, SLOVENIA CONSULTING CHEMICAL LABORATORIES, ATLASVILLE, BOKSBURG RSA TECHNIKON LABORATORIES, FLORIDA, RSA IMPILO DRUGS, ISITHEBE, KZN	Laboratory:	FPRC: NOVARTIS PHARMA STEIN AG, STEIN, SWITZERLAND INSPECTORATE M&L, ORMONDE, JOHANNESBURG
FPRR:	PHARMA DYNAMICS, SILVERWOOD, WESTLAKE, RSA	FPRC/FPRR:	NOVARTIS, SPARTAN, KEMPTON PARK
Shelf-life:	24 months	Shelf-life:	24 months (provisional)
Date of registration:	2 MARCH 2007	Date of registration:	2 MARCH 2007

AIDE 1E

Registration number: A40/27/0267
 Name of medicine: EXJADE 250 mg
 Dosage form: TABLET
 Active ingredients: EACH TABLET CONTAINS: DEFERASIROX
 Conditions of registration: 1, 2, 3, 4, 5, 6
 Applicant: NOVARTIS SOUTH AFRICA (PTY) LTD
 Manufacturer: NOVARTIS PHARMA STEIN AG, STEIN, SWITZERLAND
 Packer: NOVARTIS PHARMA STEIN AG, STEIN, SWITZERLAND
 Laboratory: FERC: NOVARTIS PHARMA STEIN AG, STEIN, SWITZERLAND
 INSPECTORATE M&L, ORMONDE, JOHANNESBURG
 FPRC/FPRR: NOVARTIS, SPARTAN, KEMPTON PARK
 Shelf-life: 24 months (provisional)
 Date of registration: 2 MARCH 2007

AIDE 1F

Registration number: A40/27/0268
 Name of medicine: EXJADE 500 mg
 Dosage form: TABLET
 Active ingredients: EACH TABLET CONTAINS: DEFERASIROX
 Conditions of registration: 1, 2, 3, 4, 5, 6
 Applicant: NOVARTIS SOUTH AFRICA (PTY) LTD
 Manufacturer: NOVARTIS PHARMA STEIN AG, STEIN, SWITZERLAND
 Packer: NOVARTIS PHARMA STEIN AG, STEIN, SWITZERLAND
 Laboratory: FPRC: NOVARTIS PHARMA STEIN AG, STEIN, SWITZERLAND
 INSPECTORATE M&L, ORMONDE, JOHANNESBURG
 FPRC/FPRR: NOVARTIS, SPARTAN, KEMPTON PARK
 Shelf-life: 24 months (provisional)
 Date of registration: 2 MARCH 2007

MRF 15

Registration number: A40/2.5/0348
 Name of medicine: TOPLEP 25
 Dosage form: TABLET
 Active ingredients: EACH TABLET CONTAINS: TOPIRAMATE 25,0 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: RANBAXY LABORATORIES LTD, DEWAS, INDIA
 FPRC: KHULULEKANI LABORATORY SERVICES, MIDRAND, RSA
 FPRR: RANBAXY, CENTURION, RSA

Shelf-life: 24 months
 Date of registration: 2 MARCH 2007

MRF 15

Registration number: A40/2.5/0349
 Name of medicine: TOPLEP 50
 Dosage form: TABLET
 Active ingredients: EACH TABLET CONTAINS: TOPIRAMATE 50,0 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: RANBAXY LABORATORIES LTD, DEWAS, INDIA
 FPRC: KHULULEKANI LABORATORY SERVICES, MIDRAND, RSA
 FPRR: RANBAXY, CENTURION, RSA

Shelf-life: 24 months
 Date of registration: 2 MARCH 2007

MRF 15

A40/2.5/0350

Name of medicine: TOPLEP 100

Dosage form: TABLET

Active ingredients: EACH TABLET CONTAINS: 100,0 mg TOPIRAMATE

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
KHULILEKANI LABORATORY SERVICES, MIDRAND, RSA

FPRR: RANBAXY, CENTURION, RSA

Shelf-life: 24 months

Date of registration: 2 MARCH 2007

MRF 15

Registration number: A40/2.5/0351

Name of medicine: TOPLEP 200

Dosage form: TABLET

Active ingredients: EACH TABLET CONTAINS: 200,0 mg TOPIRAMATE

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
KHULILEKANI LABORATORY SERVICES, MIDRAND, RSA

FPRR: RANBAXY, CENTURION, RSA

Shelf-life: 24 months

Date of registration: 2 MARCH 2007

MRF 15

Registration number: A40/5.10/0484

Name of medicine: ZYDUS-ONDANSETRON 4 mg INJECTION

Dosage form: INJECTION

Active ingredients: EACH 1,0 ml SOLUTION CONTAINS:
ONDANSETRON HYDROCHLORIDE EQUIVALENT
TO
ONDANSETRON 2,0 mg

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

Applicant: ZYDUS HEALTHCARE SA (PTY) LTD

Manufacturer: ZYDUS CADILA HEALTHCARE LIMITED, SANAND,
AHMEDABAD, INDIA

Packer: ZYDUS CADILA HEALTHCARE LIMITED, SANAND,
AHMEDABAD, INDIA

Laboratory: FPRC: ZYDUS CADILA HEALTHCARE LIMITED, SANAND,
AHMEDABAD, INDIA
INSTITUTE FOR PHARMACEUTICAL AND
CHEMICAL SERVICES, SILVERTONDALE, RSA
INSTITUTE FOR INDUSTRIAL PHARMACY,
NORTH-WEST UNIVERSITY, POTCHEFSTROOM

FPRR: ZYDUS HEALTHCARE, VAN DER HOFF PARK,
POTCHEFSTROOM

Shelf-life: 24 months (provisional)

Date of registration: 2 MARCH 2007

MRF 15

Registration number: A40/5.10/0485

Name of medicine: ZYDUS-ONDANSETRON 8 mg INJECTION

Dosage form: INJECTION

Active ingredients: EACH 1,0 ml SOLUTION CONTAINS:
ONDANSETRON HYDROCHLORIDE EQUIVALENT
TO
ONDANSETRON 2,0 mg

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

Applicant: ZYDUS HEALTHCARE SA (PTY) LTD

Manufacturer: ZYDUS CADILA HEALTHCARE LIMITED, SANAND,
AHMEDABAD, INDIA

Importer: ZYDUS CADILA HEALTHCARE LIMITED, SANAND,
AHMEDABAD, INDIA

Laboratory: FPRC: ZYDUS CADILA HEALTHCARE LIMITED, SANAND,
AHMEDABAD, INDIA
INSTITUTE FOR PHARMACEUTICAL AND
CHEMICAL SERVICES, SILVERTONDALE, RSA
INSTITUTE FOR INDUSTRIAL PHARMACY,
NORTH-WEST UNIVERSITY, POTCHEFSTROOM

FPRR: ZYDUS HEALTHCARE, VAN DER HOFF PARK,
POTCHEFSTROOM

Shelf-life: 24 months (provisional)

Date of registration: 2 MARCH 2007

registration number:	A40/20.1.2/0000	Registration number:	A40/7.1.3/0601
Name of medicine:	BIO-AMOKSIKLAV 1 000	Name of medicine:	QUINAGEN 5 mg
Dosage form:	TABLET	Dosage form:	TABLET
Active ingredients:	EACH TABLET CONTAINS: AMOXICILLIN TRIHYDRATE EQUIVALENT TO AMOXICILLIN 875,0 mg POTASSIUM CLAVULANATE EQUIVALENT TO CLAVULANIC ACID 125,0 mg	Active ingredients:	EACH TABLET CONTAINS: QUINAPRIL HYDROCHLORIDE EQUIVALENT TO QUINAPRIL 5,0 mg
Conditions of registration:	1, 2, 3, 4, 5, 6	Conditions of registration:	1, 2, 3, 4, 5, 6
Applicant:	BIOTECH LABORATORIES (PTY) LTD	Applicant:	MERCK GENERICS RSA (PTY) LTD
Manufacturer:	LEK PHARMACEUTICAL & CHEMICAL CO d.d., PREVALJE, SLOVENIA	Manufacturer:	MERCK FARMA Y QUIMICA S.A., BARCELONA, SPAIN MERCK PHARMACEUTICAL MANUFACTURING, WADEVILLE, GERMISTON
Packer:	LEK PHARMACEUTICAL & CHEMICAL CO d.d., PREVALJE, SLOVENIA DIVPHARM MANUFACTURING & PACKAGING, LONGDALE, INDUSTRIA	Packer:	MERCK FARMA Y QUIMICA S.A., BARCELONA, SPAIN MERCK PHARMACEUTICAL MANUFACTURING, WADEVILLE, GERMISTON GERARD LABORATORIES, DUBLIN, IRELAND GENERICS (UK) LTD, POTTERS BAR, HERTFORDSHIRE, UK
Laboratory:	FPRC: LEK PHARMACEUTICAL & CHEMICAL CO d.d., PREVALJE, SLOVENIA LEK PHARMACEUTICAL & CHEMICAL CO d.d., LJUBLJANA, SLOVENIA INSTITUTE FOR PHARMACEUTICAL SERVICES, SILVERTONDALE, RSA	Laboratory:	FPRC: MERCK FARMA Y QUIMICA S.A., BARCELONA, SPAIN MERCK PHARMACEUTICAL MANUFACTURING, WADEVILLE, GERMISTON GERARD LABORATORIES, DUBLIN, IRELAND GENERICS (UK) LTD, POTTERS BAR, HERTFORDSHIRE, UK RESEARCH INSTITUTE FOR INDUSTRIAL PHARMACY, NORTH-WEST UNIVERSITY, POTCHEFSTROOM
Shelf-life:	FPRC: BIOTECH LABORATORIES, MIDRAND, RSA	Shelf-life:	FPRC: MERCK GENERICS RSA, MODDERFONTEIN, RSA
Date of registration:	24 months 2 MARCH 2007	Date of registration:	24 months 2 MARCH 2007

MRF 15		MRF 15	
Registration number:	A407.1.3/0602	Registration number:	A407.1.3/0603
Name of medicine	QUINAGEN 10 mg	Name of medicine:	QUINAGEN 20 mg
Dosage form:	TABLET	Dosage form:	TABLET
Active ingredients:	EACH TABLET CONTAINS: QUINAPRIL HYDROCHLORIDE EQUIVALENT 10,0 mg	Active ingredients:	EACH TABLET CONTAINS: QUINAPRIL HYDROCHLORIDE EQUIVALENT 20,0 mg
Conditions of registration:	1, 2, 3, 4, 5, 6	Conditions of registration:	1, 2, 3, 4, 5, 6
Applicant:	MERCK GENERICS RSA (PTY) LTD	Applicant:	MERCK GENERICS RSA (PTY) LTD
Manufacturer:	MERCK FARMA Y QUIMICA S.A., BARCELONA, SPAIN MERCK PHARMACEUTICAL MANUFACTURING, WADEVILLE, GERMISTON	Manufacturer:	MERCK FARMA Y QUIMICA S.A., BARCELONA, SPAIN MERCK PHARMACEUTICAL MANUFACTURING, WADEVILLE, GERMISTON
Packer:	MERCK FARMA Y QUIMICA S.A., BARCELONA, SPAIN MERCK PHARMACEUTICAL MANUFACTURING, WADEVILLE, GERMISTON GERARD LABORATORIES, DUBLIN, IRELAND GENERICS (UK) LTD, POTTERS BAR, HERTFORDSHIRE, UK	Packer:	MERCK FARMA Y QUIMICA S.A., BARCELONA, SPAIN MERCK PHARMACEUTICAL MANUFACTURING, WADEVILLE, GERMISTON GERARD LABORATORIES, DUBLIN, IRELAND GENERICS (UK) LTD, POTTERS BAR, HERTFORDSHIRE, UK
Laboratory:	MERCK FARMA Y QUIMICA S.A., BARCELONA, SPAIN MERCK PHARMACEUTICAL MANUFACTURING, WADEVILLE, GERMISTON GERARD LABORATORIES, DUBLIN, IRELAND GENERICS (UK) LTD, POTTERS BAR, HERTFORDSHIRE, UK RESEARCH INSTITUTE FOR INDUSTRIAL PHARMACY, NORTH-WEST UNIVERSITY, POTCHEFSTROOM	Laboratory:	MERCK FARMA Y QUIMICA S.A., BARCELONA, SPAIN MERCK PHARMACEUTICAL MANUFACTURING, WADEVILLE, GERMISTON GERARD LABORATORIES, DUBLIN, IRELAND GENERICS (UK) LTD, POTTERS BAR, HERTFORDSHIRE, UK RESEARCH INSTITUTE FOR INDUSTRIAL PHARMACY, NORTH-WEST UNIVERSITY, POTCHEFSTROOM
FPRR:	MERCK GENERICS RSA, MODDERFONTEIN, RSA	FPRR:	MERCK GENERICS RSA, MODDERFONTEIN, RSA
Shelf-life:	24 months	Shelf-life:	24 months
Date of registration:	2 MARCH 2007	Date of registration:	2 MARCH 2007

MDE 15		MRF 15	
Registration number:	A40/7.1.3/0604	Registration number	41/7.3/0089
Name of medicine:	QUINAGEN 40 mg	Name of medicine:	SUMIG 50
Dosage form:	TABLET	Dosage form:	TABLET
Active ingredients:	EACH TABLET CONTAINS: QUINAPRIL HYDROCHLORIDE EQUIVALENT TO QUINAPRIL 40,0 mg	Active ingredients:	EACH TABLET CONTAINS: SUMATRIPTAN SUCCINATE EQUIVALENT TO SUMATRIPTAN 50,0 mg
Conditions of registration:	1, 2, 3, 4, 5, 6		
Applicant:	MERCK GENERICS RSA (PTY) LTD		
Manufacturer:	MERCK FARMA Y QUIMICA S.A., BARCELONA, SPAIN MERCK PHARMACEUTICAL MANUFACTURING, WADEVILLE, GERMISTON		
Packer:	MERCK FARMA Y QUIMICA S.A., BARCELONA, SPAIN MERCK PHARMACEUTICAL MANUFACTURING, WADEVILLE, GERMISTON GERARD LABORATORIES, DUBLIN, IRELAND GENERICS (UK) LTD, POTTERS BAR, HERTFORDSHIRE, UK	Packer:	CIPLA LTD, UNIT III, VERNA, GOA, INDIA
Laboratory:	FPRC: MERCK FARMA Y QUIMICA S.A., BARCELONA, SPAIN MERCK PHARMACEUTICAL MANUFACTURING, WADEVILLE, GERMISTON GERARD LABORATORIES, DUBLIN, IRELAND GENERICS (UK) LTD, POTTERS BAR, HERTFORDSHIRE, UK RESEARCH INSTITUTE FOR INDUSTRIAL PHARMACY, NORTH-WEST UNIVERSITY, POTCHEFSTROOM	Laboratory: FPRC:	CIPLA LTD, UNIT III, VERNA, GOA, INDIA
Shelf-life:	FPRC: MERCK GENERICS RSA, MODDERFONTEIN, RSA 24 months	FPRR Shelf-life:	CIPLA MEDPRO, ROSENPARK, BELLVILLE 24 months
Date of registration:	2 MARCH 2007	Date of registration:	2 MARCH 2007

MRF 15

Registration number:	41/7.3/0090
Name of medicine:	SUMIG 100
Dosage form:	TABLET
Active ingredients:	EACH TABLET CONTAINS: SUMATRIPTAN SUCCINATE EQUIVALENT TO SUMATRIPTAN 100.0 mg
Conditions of registration:	1, 2, 3, 4, 5, 8
Applicant:	OPD MEDPRO (PTY) LTD
Manufacturer:	CIPLA LTD, UNIT III, VERNA, GOA, INDIA
Packer:	CIPLA LTD, UNIT III, VERNA, GOA, INDIA
Laboratory: FPRC:	CIPLA LTD, UNIT III, VERNA, GOA, INDIA
FPRR:	CIPLA LIFE SCIENCES, ROSEN PARK, BELLVILLE
Shelf-life:	24 months
Date of registration:	2 MARCH 2007

MRF 16

Registration number:	41/7.3/0091
Name of medicine:	CIPLA-SUMATRIPTAN 50
Dosage form:	TABLET
Active ingredients:	EACH TABLET CONTAINS: SUMATRIPTAN SUCCINATE EQUIVALENT TO SUMATRIPTAN 50.0 mg
Conditions of registration:	1, 2, 3, 4, 5, 8
Applicant:	CIPLA LIFE SCIENCES (PTY) LTD
Manufacturer:	CIPLA LTD, UNIT III, VERNA, GOA, INDIA
Packer:	CIPLA LTD, UNIT III, VERNA, GOA, INDIA
Laboratory: FPRC:	CIPLA LTD, UNIT III, VERNA, GOA, INDIA
FPRR:	CIPLA LIFE SCIENCES, ROSEN PARK, BELLVILLE
Shelf-life:	24 months
Date of registration:	2 MARCH 2007