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IMPORTANT NOTICE

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GENERAL NOTICE ALGEMENE KENNISGEWING

NOTICE 848 OF 2010

MEDICINES CONTROL COUNCIL

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

1. The applicant shall ensure that the medicine is manufactured and controlled in terms of the current Good Manufacturing Practices as determined by Council
2. The manufacture of this medicine is subject to regular investigation and inspections by the inspectors appointed in terms of Section 26 of the Act, to assess compliance with current Good Manufacturing Practices.
3. The information in the package insert shall be updated on a regular basis to conform to the package insert recently approved by Council.
4. The applicant must comply with all the legal requirements of the Medicines and Related Substances Act, 1965 (Act 101 of 1965).
5. The registration of this medicine shall be subject to review at intervals as determined by Council regarding its quality, safety and efficacy, and the registration of this medicine may be varied subject to issues Council may deem fit.
6. The first two production batches must be fully validated in terms of the detailed process validation protocol submitted at the time of application for registration, and the validation report must be submitted within a month after completion of the validation.
7. The product may be advertised to the professions only.
8. The provisional shelf-life allocated must be confirmed with stability data over the full shelf-life period on the first two production batches (well-known API) or first three production batches (NCE) in accordance with the Stability Guideline. Stability testing submitted on any pilot batches must also be completed and reported on. The stability report must be submitted within six months after completion of the stability trial. However, Council has to be informed immediately if any parameter shows a significant change or out-of-specification result during the stability trial.
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 848 VAN 2010**MEDISYNEBEHEERRAAD****VOORWAARDES VIR DIE REGISTRASIE VAN 'N MEDISYNE IN TERME VAN DIE
BEPALINGS VAN ARTIKEL 15(7) VAN DIE WET OP BEHEER VAN MEDISYNE EN
VERWANTE STOWWE, 1965 (WET 101 VAN 1965)**

1. Die applikant sal verseker dat die medisyne vervaardig en beheer word ooreenkomstig huidige Goeie Vervaardigingspraktyke soos bepaal deur die Medisynebeheerraad.
2. Die vervaardiging van hierdie medisyne is onderhewig aan gereelde ondersoeke en inspeksies deur inspekteurs, aangestel ingevolge Artikel 25 van die Wet, om die nakoming van Goeie Vervaardigingspraktyke te bepaal.
3. Die inligting soos vervat in the voubiljet moet op 'n gereelde grondslag opgedateer word in ooreenstemming met 'n voubiljet wat onlangs deur die Raad goedgekeur is.
4. Die applikant moet voldoen aan alle wetlike vereistes van die Wet op Medisyne en Verwante Stowwe, 1965 (Wet 101 van 1965).
5. Die registrasie van hierdie medisyne is onderhewig aan gereelde hersiening met tussenpose soos deur die Raad bepaal, rakende kwaliteit, veiligheid en effektiwiteit, en die registrasie van hierdie medisyne kan gewysig word onderhewig aan kwessies na goëddunke van die Raad.
6. Die eerste twee produksielotte moet ten volle gevalideer word ooreenkomstig die breedvoerige prosesvalidasie protokol wat ingedien is ten tye van die aansoek om registrasie, en die validasieverlag moet binne die bestek van een maand na die voltooiing van die validasie ingedien word.
7. Die produk mag slegs aan die beroepe geadverteer word.
8. Die toegekende voorlopige rakkleeftyd moet bevestig word met stabiliteitsdata volgens die Stabiliteitsriglyn op die eerste twee produksielotte (bekende aktiewe bestanddele), of eerste drie produksielotte (nuwe chemiese entiteite), wat die volle rakkleeftydperiode dek. Stabiliteitstoetse wat op proeflotte ingedien is, moet ook voltooi word en die verslae ingedien word. Die stabiliteitsverslag moet binne ses maande na voltooiing van die stabiliteitsproef ingedien word. Die Raad moet egter onmiddellik in kennis gestel word indien enige parameter 'n betekenisvolle verandering of buite-spesifikasieresultaat tydens die stabiliteitsproef lewer.
9. 'n Na-registrasie-inspeksie moet op die eerste produksielot van elke plaaslike vervaardiger uitgevoer word.
10. 'n Na-registrasie-inspeksie moet op die eerste produksielot van die ingevoerde produk uitgevoer word.
11. Bemaking van die produk mag slegs 'n aanvang neem nadat 'n bevredigende na-registrasie-inspeksieverlag gedien het.
12. Een monster van elke lot moet tesame met vier kopieë van die protokolle vir die toets van die massalot en die vullot sowel as ses kopieë van die vrystellingsertifikaat wat uitgereik is deur die verantwoordelike beheerliggaam in die land waar die produk vervaardig word, ingedien word by die Raad vir lotvrystellingsdoeleindes.
13. Die vervaldatum toegeken, sal gewysig word deur 'n verklaring by te voeg dat die virusstamme tans vir Suid-Afrikaanse gebruik gedurende die gespesifiseerde jaar aanbeveel word.
14. Die stamme van die oorspronklike saadvirusse moet elke jaar deur die Departement van Gesondheid goedgekeur word.

MRF 15

Registration number: 36/2.7/0223

Name of medicine: GRAND-PA PARACETAMOL TABLETS

Dosage form: TABLET

Active ingredients: EACH TABLET CONTAINS:
PARACETAMOL 500,0 mg
SODIUM BICARBONATE 630,0 mg

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

Applicant: GROUP LABORATORIES SA (PTY) LTD

Manufacturer: GLAXOSMITHKLINE,
DUNGARVAN, IRELAND
GLAXOSMITHKLINE,
EPPING, CAPE TOWN

Packer: GLAXOSMITHKLINE,
DUNGARVAN, IRELAND
GLAXOSMITHKLINE,
EPPING, CAPE TOWN

Laboratory: FPRC: GLAXOSMITHKLINE,
DUNGARVAN, IRELAND
GLAXOSMITHKLINE,
EPPING, CAPE TOWN

FPRR: GROUP LABORATORIES,
EPPING, CAPE TOWN

Shelf-life: 24 months

Date of registration: 23 JULY 2010

MRF15

Registration number: A40/16.4/0358

Name of medicine: STREPSILS STRAWBERRY SUGAR-FREE

Dosage form: LOZENGES

Active ingredients: EACH LOZENGE
CONTAINS:
AMYL METACRESOL 0,6 mg
DICHLOOROBENZYL-
ALCOHOL 1,2 mg

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

Applicant: RECKITT BENCKISER
PHARMACEUTICALS (PTY)
LTD

Manufacturer: BOOTS MANUFACTURING,
NOTTINGHAM, UK

Packer: BOOTS MANUFACTURING,
NOTTINGHAM, UK
NATUR PRODUKT
PHARMA LTD, OSTROW,
MAZOWIECKA, POLAND
PHARMACEUTICAL
CONTRACTORS, ISANDO,
KEMPTON PARK

Laboratory: FPRC: BOOTS MANUFACTURING,
NOTTINGHAM, UK
NATUR PRODUKT
PHARMA LTD, OSTROW,
MAZOWIECKA, POLAND
PHARMACEUTICAL
CONTRACTORS, ISANDO,
KEMPTON PARK
PHARMA-Q, INDUSTRIA-
WEST, JOHANNESBURG

FPRR: RECKITT BENCKISER
PHARMACEUTICALS,
ELANDSFONTEIN

Shelf-life: 24 months (Provisional)

Date of registration: 23 JULY 2010

MRF 15

Registration number: 41/20.2.8/0747

Name of medicine: PREZISTA

Dosage form: TABLET

Active ingredients: EACH TABLET CONTAINS:
DARUNAVIR ETHANOLATE
EQUIVALENT TO
DARUNAVIR 300,0 mg

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

Applicant: PHARMACARE LIMITED

Manufacturer: JANSSEN ORTHO LLC,
GURABO, PUERTO RICO

Packer: JANSSEN-CILAG SpA, SAN
MICHELE, LATINA, ITALY
PHARMACARE LTD, KORSTEN,
PORT ELIZABETH

Laboratory: FPRC: JANSSEN-CILAG SpA, SAN
MICHELE, LATINA, ITALY

FPRC/FPRR: PHARMACARE LTD, KORSTEN,
PORT ELIZABETH

FPRR: PHARMACARE LTD,
WOODMEAD, SANDTON

Shelf-life: 24 months (Provisional)

Date of registration: 23 JULY 2010

MRF 15

Registration number: 41/26/0933
Name of medicine: GEMTAZ 1 g
Dosage form: INJECTION
Active ingredients: EACH VIAL CONTAINS:
 GEMCITABINE
 HYDROCHLORIDE
 EQUIVALENT TO
 GEMCITABINE 1,0 g
Conditions of registration: 1, 2, 3, 4, 5, 6, 7, 8
Applicant: PHARMAPLAN (PTY) LTD
Manufacturer: SUN PHARMACEUTICALS
 INDUSTRIES LTD, HALOL,
 DISTRICT PANCHMAHAL,
 GUJARAT, INDIA
Packer: SUN PHARMACEUTICALS
 INDUSTRIES LTD, HALOL,
 DISTRICT PANCHMAHAL,
 GUJARAT, INDIA
Laboratory: FPRC: SUN PHARMACEUTICALS
 INDUSTRIES LTD, HALOL,
 DISTRICT PANCHMAHAL,
 GUJARAT, INDIA
 CONSULTING CHEMICAL
 LABORATORIES,
 ATLASVILLE, BOKSBURG
FPRR: PHARMAPLAN, MIDRAND,
 RSA
Shelf-life: 24 months (Provisional)
Date of registration: 23 JULY 2010

MRF 15

Registration number: 41/26/0934
Name of medicine: GEMTAZ 200 mg
Dosage form: INJECTION
Active ingredients: EACH VIAL CONTAINS:
 GEMCITABINE
 HYDROCHLORIDE
 EQUIVALENT TO
 GEMCITABINE 200,0 mg
Conditions of registration: 1, 2, 3, 4, 5, 6, 7, 8
Applicant: PHARMAPLAN (PTY) LTD
Manufacturer: SUN PHARMACEUTICALS
 INDUSTRIES LTD, HALOL,
 DISTRICT PANCHMAHAL,
 GUJARAT, INDIA
Packer: SUN PHARMACEUTICALS
 INDUSTRIES LTD, HALOL,
 DISTRICT PANCHMAHAL,
 GUJARAT, INDIA
Laboratory: FPRC: SUN PHARMACEUTICALS
 INDUSTRIES LTD, HALOL,
 DISTRICT PANCHMAHAL,
 GUJARAT, INDIA
 CONSULTING CHEMICAL
 LABORATORIES,
 ATLASVILLE, BOKSBURG
FPRR: PHARMAPLAN, MIDRAND,
 RSA
Shelf-life: 24 months (Provisional)
Date of registration: 23 JULY 2010

MRF 15

Registration number: 42/20.2.8/0202
Name of medicine: NYSIVIR ORAL SOLUTION 2 g
Dosage form: SOLUTION
Active ingredients: EACH 100,0 ml SOLUTION
 CONTAINS:
 DIDANOSINE 2,0 g
Conditions of registration: 1, 2, 3, 4, 5, 6, 7, 8
Applicant: AUROBINDO PHARMA (PTY)
 LTD
Manufacturer: AUROBINDO PHARMA LTD,
 UNIT III, QUTHUBULLAPUR
 MANDAL, RANGA REDDY
 DISTRICT, ANDHRA PRADESH,
 INDIA
Packer: AUROBINDO PHARMA LTD,
 UNIT III, QUTHUBULLAPUR
 MANDAL, RANGA REDDY
 DISTRICT, ANDHRA PRADESH,
 INDIA
Laboratory: FPRC: AUROBINDO PHARMA LTD,
 UNIT III, QUTHUBULLAPUR
 MANDAL, RANGA REDDY
 DISTRICT, ANDHRA PRADESH,
 INDIA
FPRR: AUROBINDO PHARMA,
 MEYERSDAL, JOHANNESBURG
Shelf-life: 24 months (Provisional)
Date of registration: 23 JULY 2010

MRF 15

Registration number: 42/20.2.8/0203
Name of medicine: NYSIVIR ORAL SOLUTION
 4 g
Dosage form: SOLUTION
Active ingredients: EACH 100,0 ml CONTAINS:
 DIDANOSINE 4,0 g

Conditions of registration: 1, 2, 3, 4, 5, 6, 7, 8
Applicant: AUROBINDO PHARMA (PTY) LTD
Manufacturer: AUROBINDO PHARMA LTD,
 UNIT III, QUTHUBULLAPUR
 MANDAL, RANGA REDDY
 DISTRICT, ANDHRA
 PRADESH, INDIA
Packer: AUROBINDO PHARMA LTD,
 UNIT III, QUTHUBULLAPUR
 MANDAL, RANGA REDDY
 DISTRICT, ANDHRA
 PRADESH, INDIA
Laboratory: FPRC: AUROBINDO PHARMA LTD,
 UNIT III, QUTHUBULLAPUR
 MANDAL, RANGA REDDY
 DISTRICT, ANDHRA
 PRADESH, INDIA
FPRR: AUROBINDO PHARMA,
 MEYERSDAL,
 JOHANNESBURG
Shelf-life: 24 months (Provisional)
Date of registration: 23 JULY 2010

MRF15

Registration number: 42/21.5.4/0218
Name of medicine: SEREFLO 25/50 HFA
Dosage form: INHALER
Active ingredients: EACH ACTUATION
 DELIVERS:
 SALMETEROL XINAFOATE
 EQUIVALENT TO
 SALMETEROL 25,0 µg
 FLUTICASONE
 PROPIONATE 50,0 µg

Conditions of registration: 1, 2, 3, 4, 5, 6, 7
Applicant: CIPLA MEDPRO (PTY) LTD
Manufacturer: CIPLA LTD, UNIT II,
 VERNA, SALCETTE, GOA,
 INDIA
Packer: CIPLA LTD, UNIT II,
 VERNA, SALCETTE, GOA,
 INDIA
Laboratory: FPRC: CIPLA LTD, UNIT II,
 VERNA, SALCETTE, GOA,
 INDIA
FPRR: CIPLA MEDPRO, ROSEN
 PARK, BELLVILLE
Shelf-life: 24 months
Date of registration: 23 JULY 2010

MRF 15

Registration number: 42/21.5.4/0219
Name of medicine: SEREFLO 25/125 HFA
Dosage form: INHALER
Active ingredients: EACH ACTUATION DELIVERS:
 SALMETEROL XINAFOATE
 EQUIVALENT TO
 SALMETEROL 25,0 µg
 FLUTICASONE PROPIONATE
 125,0 µg

Conditions of registration: 1, 2, 3, 4, 5, 6, 7
Applicant: CIPLA MEDPRO (PTY) LTD
Manufacturer: CIPLA LTD, UNIT II, VERNA,
 SALCETTE, GOA, INDIA
Packer: CIPLA LTD, UNIT II, VERNA,
 SALCETTE, GOA, INDIA
Laboratory: FPRC: CIPLA LTD, UNIT II, VERNA,
 SALCETTE, GOA, INDIA
FPRR: CIPLA MEDPRO, ROSEN PARK,
 BELLVILLE
Shelf-life: 24 months
Date of registration: 23 JULY 2010

MRF 15

Registration number:	42/21.5.4/0220
Name of medicine:	SEREFLO 25/250 HFA
Dosage form:	INHALER
Active ingredients:	EACH ACTUATION DELIVERS: SALMETEROL XINAFOATE EQUIVALENT TO SALMETEROL 25,0 µg FLUTICASON PROPIONATE 250,0 µg
Conditions of registration:	1, 2, 3, 4, 5, 6, 7
Applicant:	CIPLA MEDPRO(PTY) LTD
Manufacturer:	CIPLA LTD, UNIT II, VERNA, SALCETTE, GOA, INDIA
Packer:	CIPLA LTD, UNIT II, VERNA, SALCETTE, GOA, INDIA
Laboratory:FPRC:	CIPLA LTD, UNIT II, VERNA, SALCETTE, GOA, INDIA
FPRR:	CIPLA MEDPRO, ROSENPAK, BELLVILLE
Shelf-life:	12 months
Date of registration:	23 JULY 2010

MRF15

Registration number:	42/1.2/0423
Name of medicine:	EFEGEN XR 75
Dosage form:	CAPSULE
Active ingredients:	EACH CAPSULE CONTAINS: VENLAFAXINE HYDROCHLORIDE EQUIVALENT TO VENLAFAXINE 75,0 mg
Conditions of registration:	1, 2, 3, 4, 5, 6, 7, 8
Applicant:	RANBAXY (S.A.) (PTY) LTD
Manufacturer:	RANBAXY LABORATORIES LTD, PAONTA SAHIB, SIRMOUR, HIMACHAL PRADESH, INDIA
Packer:	RANBAXY LABORATORIES LTD, PAONTA SAHIB, SIRMOUR, HIMACHAL PRADESH, INDIA
Laboratory: FPRC:	RANBAXY LABORATORIES LTD, PAONTA SAHIB, SIRMOUR, HIMACHAL PRADESH, INDIA KHULULEKANI LABORATORY SERVICES, COVENTRY PARK, MIDRAND CONSULTING CHEMICAL LABORATORIES, ATLASVILLE, BOKSBURG
FPRR:	RANBAXY (SA), CENTURION, RSA
Shelf-life:	24 months (Provisional)
Date of registration:	23 JULY 2010

MRF 15

Registration number:	42/1.2/0424
Name of medicine:	EFEGEN XR 150
Dosage form:	CAPSULE
Active ingredients:	EACH CAPSULE CONTAINS: VENLAFAXINE HYDROCHLORIDE EQUIVALENT TO VENLAFAXINE 75,0 mg
Conditions of registration:	1, 2, 3, 4, 5, 6, 7, 8
Applicant:	RANBAXY (S.A.) (PTY) LTD
Manufacturer:	RANBAXY LABORATORIES LTD, PAONTA SAHIB, SIRMOUR, HIMACHAL PRADESH, INDIA
Packer:	RANBAXY LABORATORIES LTD, PAONTA SAHIB, SIRMOUR, HIMACHAL PRADESH, INDIA
Laboratory: FPRC:	RANBAXY LABORATORIES LTD, PAONTA SAHIB, SIRMOUR, HIMACHAL PRADESH, INDIA KHULULEKANI LABORATORY SERVICES, COVENTRY PARK, MIDRAND CONSULTING CHEMICAL LABORATORIES, ATLASVILLE, BOKSBURG
FPRR:	RANBAXY (SA), CENTURION, RSA
Shelf-life:	24 months (Provisional)
Date of registration:	23 JULY 2010

MRF 15

Registration number: 42/1.2/0425
 Name of medicine: RAN VENLAFAXINE XR 75
 Dosage form: CAPSULE
 Active ingredients: EACH CAPSULE CONTAINS:
 VENLAFAZINE
 HYDROCHLORIDE
 EQUIVALENT TO
 VENLAFAXINE 75,0 mg
 Conditions of registration: 1, 2, 3, 4, 5, 6, 7, 8
 Applicant: RANBAXY (S.A.) (PTY) LTD
 Manufacturer: RANBAXY LABORATORIES
 LTD, PAONTA SAHIB,
 SIRMOUR, HIMACHAL
 PRADESH, INDIA
 Packer: RANBAXY LABORATORIES
 LTD, PAONTA SAHIB,
 SIRMOUR, HIMACHAL
 PRADESH, INDIA
 Laboratory:
 FPRC: RANBAXY LABORATORIES
 LTD, PAONTA SAHIB,
 SIRMOUR, HIMACHAL
 PRADESH, INDIA
 KHULULEKANI
 LABORATORY SERVICES,
 COVENTRY PARK,
 MIDRAND
 CONSULTING CHEMICAL
 LABORATORIES,
 ATLASVILLE, BOKSBURG
 FPRR: RANBAXY (SA),
 CENTURION, RSA
 Shelf-life: 24 months (Provisional)
 Date of registration: 23 JULY 2010

MRF15

Registration number: 42/1.2/0426
 Name of medicine: RAN VENLAFAXINE XR
 150 mg
 Dosage form: CAPSULE
 Active ingredients: EACH CAPSULE
 CONTAINS:
 VENLAFAZINE
 HYDROCHLORIDE
 EQUIVALENT TO
 VENLAFAXINE 150,0 mg
 Conditions of registration: 1,2,3,4,5,6,7,8
 Applicant: RANBAXY (S.A.) (PTY) LTD
 Manufacturer: RANBAXY LABORATORIES
 LTD, PAONTA SAHIB,
 SIRMOUR, HIMACHAL
 PRADESH, INDIA
 Packer: RANBAXY LABORATORIES
 LTD, PAONTA SAHIB,
 SIRMOUR, HIMACHAL
 PRADESH, INDIA
 Laboratory: FPRC: RANBAXY LABORATORIES
 LTD, PAONTA SAHIB,
 SIRMOUR, HIMACHAL
 PRADESH, INDIA
 KHULULEKANI
 LABORATORY SERVICES,
 COVENTRY PARK,
 MIDRAND
 CONSULTING CHEMICAL
 LABORATORIES,
 ATLASVILLE, BOKSBURG
 FPRR: RANBAXY (SA),
 CENTURION, RSA
 Shelf-life: 24 months (Provisional)
 Date of registration: 23 JULY 2010

MRF 15

Registration number: 42/3.1/0467
 Name of medicine: AURO MELOXICAM 7,5 mg
 Dosage form: TABLET
 Active ingredients: EACH TABLET CONTAINS:
 MELOXICAM 7,5 mg
 Conditions of registration: 1, 2, 3, 4, 5, 6, 7, 8
 Applicant: AUROBINDO PHARMA (PTY)
 LTD
 Manufacturer: AUROBINDO PHARMA LTD,
 UNIT III, QUTHUBULLAPUR
 MANDAL, RANGA REDDY
 DISTRICT, ANDHRA PRADESH,
 INDIA
 Packer: AUROBINDO PHARMA LTD,
 UNIT III, QUTHUBULLAPUR
 MANDAL, RANGA REDDY
 DISTRICT, ANDHRA PRADESH,
 INDIA
 Laboratory:
 FPRC: AUROBINDO PHARMA LTD,
 UNIT III, QUTHUBULLAPUR
 MANDAL, RANGA REDDY
 DISTRICT, ANDHRA PRADESH,
 INDIA
 FPRR: AUROBINDO PHARMA,
 MEYERSDAL, JOHANNESBURG
 Shelf-life: 24 months (Provisional)
 Date of registration: 23 JULY 2010

MRF 15

Registration number: 42/3.1/0468
Name of medicine: AURO MELOXICAM 15 mg
Dosage form: TABLET
Active ingredients: EACH TABLET CONTAINS:
 MELOXICAM 15.0 mg
Conditions of registration: 1, 2, 3, 4, 5, 6, 7, 8
Applicant: AUROBINDO PHARMA (PTY) LTD
Manufacturer: AUROBINDO PHARMA LTD,
 UNIT III, QUTHUBULLAPUR
 MANDAL, RANGA REDDY
 DISTRICT, ANDHRA PRADESH,
 INDIA
Packer: AUROBINDO PHARMA LTD,
 UNIT III, QUTHUBULLAPUR
 MANDAL, RANGA REDDY
 DISTRICT, ANDHRA PRADESH,
 INDIA
Laboratory: FPRC: AUROBINDO PHARMA LTD,
 UNIT III, QUTHUBULLAPUR
 MANDAL, RANGA REDDY
 DISTRICT, ANDHRA PRADESH,
 INDIA
FPRR: AUROBINDO PHARMA,
 MEYERSDAL, JOHANNESBURG
Shelf-life: 24 months (Provisional)
Date of registration: 23 JULY 2010

MRF15

Registration number: 41/3.1/0469
Name of medicine: FLAMARYX 7,5 mg
Dosage form: TABLET
Active ingredients: EACH TABLET CONTAINS:
 MELOXICAM 7,5 mg
Conditions of registration: 1, 2, 3, 4, 5, 6, 7, 8
Applicant: AUROBINDO PHARMA (PTY) LTD
Manufacturer: AUROBINDO PHARMA LTD,
 UNIT III, QUTHUBULLAPUR
 MANDAL, RANGA REDDY
 DISTRICT, ANDHRA PRADESH,
 INDIA
Packer: AUROBINDO PHARMA LTD,
 UNIT III, QUTHUBULLAPUR
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 DISTRICT, ANDHRA PRADESH,
 INDIA
Laboratory: FPRC: AUROBINDO PHARMA LTD,
 UNIT III, QUTHUBULLAPUR
 MANDAL, RANGA REDDY
 DISTRICT, ANDHRA PRADESH,
 INDIA
FPRR: AUROBINDO PHARMA,
 MEYERSDAL,
 JOHANNESBURG
Shelf-life: 24 months (Provisional)
Date of registration: 23 JULY 2010

MRF 15

Registration number: 41/3.1/0470
Name of medicine: FLAMARYX 15 mg
Dosage form: TABLET
Active ingredients: EACH TABLET CONTAINS:
 MELOXICAM 15,0 mg
Conditions of registration: 1, 2, 3, 4, 5, 6, 7, 8
Applicant: AUROBINDO PHARMA (PTY) LTD
Manufacturer: AUROBINDO PHARMA LTD, UNIT III,
 QUTHUBULLAPUR MANDAL, RANGA
 REDDY DISTRICT, ANDHRA
 PRADESH, INDIA
Packer: AUROBINDO PHARMA LTD, UNIT III,
 QUTHUBULLAPUR MANDAL, RANGA
 REDDY DISTRICT, ANDHRA
 PRADESH, INDIA
Laboratory: FPRC: AUROBINDO PHARMA LTD, UNIT III,
 QUTHUBULLAPUR MANDAL, RANGA
 REDDY DISTRICT, ANDHRA
 PRADESH, INDIA
FPRR: AUROBINDO PHARMA,
 MEYERSDAL, JOHANNESBURG
Shelf-life: 24 months (Provisional)
Date of registration: 23 JULY 2010

MRF 15

Registration number: 42/20.2.8/0936

Name of medicine: SONKE LAMIVUDINE + ZIDOVUDINE 150/300 AND SONKE EFAVIRENZ 600 CO-PACK

Dosage form: TABLETS

Active ingredients: EACH BLISTER CONTAINS:
2 x SONKE LAMIVUDINE + ZIDOVUDINE TABLETS CONTAINING
LAMIVUDINE 150,0 mg
ZIDOVUDINE 300,0 mg

1 x SONKE EFAVIRENZ 600 TABLET CONTAINING
EFAVIRENZ 600,0 mg

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

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

Manufacturer: RANBAXY LABORATORIES LTD, PAONTA SAHIB, SIRMOUR, HIMACHAL PRADESH, INDIA

Packer: RANBAXY LABORATORIES LTD, PAONTA SAHIB, SIRMOUR, HIMACHAL PRADESH, INDIA

Laboratory: FPRC: RANBAXY LABORATORIES LTD, PAONTA SAHIB, SIRMOUR, HIMACHAL PRADESH, INDIA
KHULULEKANI LABORATORY SERVICES, COVENTRY PARK, MIDRAND
CONSULTING CHEMICAL LABORATORIES, ATLASVILLE, BOKSBURG
BE-TABS PHARMACEUTICALS, ROODEPOORT, RSA

FPRR: RANBAXY (S.A.), CENTURION, RSA

Shelf-life: 24 months

Date of registration: 23 JULY 2010

MRF15

Registration number: 42/20.2.8/0973

Name of medicine: RENZIR 50 mg

Dosage form: CAPSULE

Active ingredients: EACH CAPSULE CONTAINS:
EFAVIRENZ 50,0 mg

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

Applicant: ADCOCK INGRAM LIMITED

Manufacturer: ADCOCK INGRAM HEALTHCARE, WADEVILLE, GERMISTON

Packer: ADCOCK INGRAM HEALTHCARE, WADEVILLE, GERMISTON

Laboratory: FPRC/FPRR ADCOCK INGRAM HEALTHCARE, WADEVILLE, GERMISTON
ADCOCK INGRAM LTD, AEROTON, JOHANNESBURG

FPRR: ADCOCK INGRAM LTD, ERAND GARDENS, MIDRAND

Shelf-life: 24 months (Provisional)

Date of registration: 23 JULY 2010

MRF 15

Registration number: 42/20.2.8/0974

Name of medicine: RENZIR 100 mg

Dosage form: CAPSULE

Active ingredients: EACH CAPSULE CONTAINS:
EFAVIRENZ 100,0 mg

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

Applicant: ADCOCK INGRAM LIMITED

Manufacturer: ADCOCK INGRAM HEALTHCARE, WADEVILLE, GERMISTON

Packer: ADCOCK INGRAM HEALTHCARE, WADEVILLE, GERMISTON

Laboratory: FPRC/FPRR ADCOCK INGRAM HEALTHCARE, WADEVILLE, GERMISTON
ADCOCK INGRAM LTD, AEROTON, JOHANNESBURG

FPRR: ADCOCK INGRAM LTD, ERAND GARDENS, MIDRAND

Shelf-life: 24 months (Provisional)

Date of registration: 23 JULY 2010

MRF 15

Registration number:	42/20.2.8/0976
Name of medicine:	REFAVIN 50 mg
Dosage form:	CAPSULE
Active ingredients:	EACH CAPSULE CONTAINS: EFAVIRENZ 50,0 mg
Conditions of registration:	1, 2, 3, 4, 5, 6, 7, 8
Applicant:	ADCOCK INGRAM LIMITED
Manufacturer:	ADCOCK INGRAM HEALTHCARE, WADEVILLE, GERMISTON
Packer:	ADCOCK INGRAM HEALTHCARE, WADEVILLE, GERMISTON
Laboratory: FPRC/FPRR:	ADCOCK INGRAM HEALTHCARE, WADEVILLE, GERMISTON ADCOCK INGRAM LTD, AEROTON, JOHANNESBURG
FPRR:	ADCOCK INGRAM LTD, ERAND GARDENS, MIDRAND
Shelf-life:	24 months (Provisional)
Date of registration:	23 JULY 2010

MRF15

Registration number:	42/20.2.8/0977
Name of medicine:	REFAVIN 100 mg
Dosage form:	CAPSULE
Active ingredients:	EACH CAPSULE CONTAINS: EFAVIRENZ 100,0 mg
Conditions of registration:	1, 2, 3, 4, 5, 6, 7, 8
Applicant:	ADCOCK INGRAM LIMITED
Manufacturer:	ADCOCK INGRAM HEALTHCARE, WADEVILLE, GERMISTON
Packer:	ADCOCK INGRAM HEALTHCARE, WADEVILLE, GERMISTON
Laboratory: FPRC/FPRR:	ADCOCK INGRAM HEALTHCARE, WADEVILLE, GERMISTON ADCOCK INGRAM LTD, AEROTON, JOHANNESBURG
FPRR:	ADCOCK INGRAM LTD, ERAND GARDENS, MIDRAND
Shelf-life:	24 months (Provisional)
Date of registration:	23 JULY 2010

MRF 15

Registration number:	42/20.2.8/0979
Name of medicine:	ADCO EFAVIRENZ 50 mg
Dosage form:	CAPSULE
Active ingredients:	EACH CAPSULE CONTAINS: EFAVIRENZ 50,0 mg
Conditions of registration:	1, 2, 3, 4, 5, 6, 7, 8
Applicant:	ADCOCK INGRAM LIMITED
Manufacturer:	ADCOCK INGRAM HEALTHCARE, WADEVILLE, GERMISTON
Packer:	ADCOCK INGRAM HEALTHCARE, WADEVILLE, GERMISTON
Laboratory: FPRC/FPRR:	ADCOCK INGRAM HEALTHCARE, WADEVILLE, GERMISTON ADCOCK INGRAM LTD, AEROTON, JOHANNESBURG
FPRR:	ADCOCK INGRAM LTD, ERAND GARDENS, MIDRAND
Shelf-life:	24 months (Provisional)
Date of registration:	23 JULY 2010

MRF 15

Registration number:	42/20.2.8/0980
Name of medicine:	ADCO EFAVIRENZ 100 mg
Dosage form:	CAPSULE
Active ingredients:	EACH CAPSULE CONTAINS: EFAVIRENZ 100,0 mg
Conditions of registration:	1, 2, 3, 4, 5, 6, 7, 8
Applicant:	ADCOCK INGRAM LIMITED
Manufacturer:	ADCOCK INGRAM HEALTHCARE, WADEVILLE, GERMISTON
Packer:	ADCOCK INGRAM HEALTHCARE, WADEVILLE, GERMISTON
Laboratory: FPRC/FPRR:	ADCOCK INGRAM HEALTHCARE, WADEVILLE, GERMISTON ADCOCK INGRAM LTD, AEROTON, JOHANNESBURG
FPRR:	ADCOCK INGRAM LTD, ERAND GARDENS, MIDRAND
Shelf-life:	24 months (Provisional)
Date of registration:	23 JULY 2010

MRF15

Registration number:	42/21.2/1033
Name of medicine:	GLIMY 1
Dosage form:	TABLET
Active ingredients:	EACH TABLET CONTAINS: GLIMEPIRIDE 1,0 mg
Conditions of registration:	1, 2, 3, 4, 5, 6, 7, 8
Applicant:	DR REDDY'S LABORATORIES (PTY) LTD
Manufacturer:	DR REDDY'S LABORATORIES LTD, QUTHUBULLAPUR MANDAL, RANGA REDDY DISTRICT, ANDHRA PRADESH, INDIA
Packer:	DR REDDY'S LABORATORIES LTD, QUTHUBULLAPUR MANDAL, RANGA REDDY DISTRICT, ANDHRA PRADESH, INDIA
Laboratory: FPRC:	DR REDDY'S LABORATORIES LTD, QUTHUBULLAPUR MANDAL, RANGA REDDY DISTRICT, ANDHRA PRADESH, INDIA RESEARCH INSTITUTE FOR INDUSTRIAL PHARMACY, NORTH-WEST UNIVERSITY, POTCHEFSTROOM
FPRR:	DR REDDY'S LABORATORIES, MURRAYFIELD, PRETORIA
Shelf-life:	24 months (Provisional)
Date of registration:	23 JULY 2010

MRF 15

Registration number:	42/21.2/1034
Name of medicine:	GLIMY 2
Dosage form:	TABLET
Active ingredients:	EACH TABLET CONTAINS: GLIMEPIRIDE 2,0 mg
Conditions of registration:	1, 2, 3, 4, 5, 6, 7, 8
Applicant:	DR REDDY'S LABORATORIES (PTY) LTD
Manufacturer:	DR REDDY'S LABORATORIES LTD, QUTHUBULLAPUR MANDAL, RANGA REDDY DISTRICT, ANDHRA PRADESH, INDIA
Packer:	DR REDDY'S LABORATORIES LTD, QUTHUBULLAPUR MANDAL, RANGA REDDY DISTRICT, ANDHRA PRADESH, INDIA
Laboratory: FPRC:	DR REDDY'S LABORATORIES LTD, QUTHUBULLAPUR MANDAL, RANGA REDDY DISTRICT, ANDHRA PRADESH, INDIA RESEARCH INSTITUTE FOR INDUSTRIAL PHARMACY, NORTH-WEST UNIVERSITY, POTCHEFSTROOM
FPRR:	DR REDDY'S LABORATORIES, MURRAYFIELD, PRETORIA
Shelf-life:	24 months (Provisional)
Date of registration:	23 JULY 2010

MRF 15

Registration number:	42/21.2/1035
Name of medicine:	GLIMY 4
Dosage form:	TABLET
Active ingredients:	EACH TABLET CONTAINS: GLIMEPIRIDE 4,0 mg
Conditions of registration:	1, 2, 3, 4, 5, 6, 7, 8
Applicant:	DR REDDY'S LABORATORIES (PTY) LTD
Manufacturer:	DR REDDY'S LABORATORIES LTD, QUTHUBULLAPUR MANDAL, RANGA REDDY DISTRICT, ANDHRA PRADESH, INDIA
Packer:	DR REDDY'S LABORATORIES LTD, QUTHUBULLAPUR MANDAL, RANGA REDDY DISTRICT, ANDHRA PRADESH, INDIA
Laboratory: FPRC:	DR REDDY'S LABORATORIES LTD, QUTHUBULLAPUR MANDAL, RANGA REDDY DISTRICT, ANDHRA PRADESH, INDIA RESEARCH INSTITUTE FOR INDUSTRIAL PHARMACY, NORTH-WEST UNIVERSITY, POTCHEFSTROOM
FPRR:	DR REDDY'S LABORATORIES, MURRAYFIELD, PRETORIA
Shelf-life:	24 months (Provisional)
Date of registration:	23 JULY 2010

MRF15

Registration number:	43/7.1.3/0032
Name of medicine:	MIGROBEN 80 TABLET
Dosage form:	TABLET
Active ingredients:	EACH TABLET CONTAINS: VALSARTAN 80,0 mg
Conditions of registration:	1, 2, 3, 4, 5, 6, 7
Applicant:	NOVARTIS SOUTH AFRICA (PTY) LTD
Manufacturer:	NOVARTIS PHARMA STEIN AG, STEIN, SWITZERLAND NOVARTIS PHARMACEUTICA S.A., BARBERA DEL VALLES, BARCELONA, SPAIN
Packer:	NOVARTIS PHARMA STEIN AG, STEIN, SWITZERLAND NOVARTIS PHARMACEUTICA S.A., BARBERA DEL VALLES, BARCELONA, SPAIN ALLPACK AG, REINACH, SWITZERLAND KONAPHARMA AG, PRATTELN, SWITZERLAND NOVARTIS PHARMA GmbH, WEHR/BADEN, GERMANY IVERS-LEE AG, BURGDORF, SWITZERLAND NOVARTIS SA, SPARTAN, KEMPTON PARK
Laboratory: FPRC:	NOVARTIS PHARMA STEIN AG, STEIN, SWITZERLAND NOVARTIS PHARMACEUTICA S.A., BARBERA DEL VALLES, BARCELONA, SPAIN IVERS-LEE AG, BURGDORF, SWITZERLAND M&L LABORATORY SERVICES, ORMONDE, JOHANNESBURG
FPRC/FPRR:	NOVARTIS SA, SPARTAN, KEMPTON PARK
Shelf-life:	36 months
Date of registration:	23 JULY 2010

MRF 15

Registration number:	43/7.1.3/0033
Name of medicine:	MIGROBEN 160 TABLET
Dosage form:	TABLET
Active ingredients:	EACH TABLET CONTAINS: VALSARTAN 160,0 mg
Conditions of registration:	1, 2, 3, 4, 5, 6, 7
Applicant:	NOVARTIS SOUTH AFRICA (PTY) LTD
Manufacturer:	NOVARTIS PHARMA STEIN AG, STEIN, SWITZERLAND NOVARTIS PHARMACEUTICA S.A., BARBERA DEL VALLES, BARCELONA, SPAIN
Packer:	NOVARTIS PHARMA STEIN AG, STEIN, SWITZERLAND NOVARTIS PHARMACEUTICA S.A., BARBERA DEL VALLES, BARCELONA, SPAIN ALLPACK AG, REINACH, SWITZERLAND KONAPHARMA AG, PRATTELN, SWITZERLAND NOVARTIS PHARMA GmbH, WEHR/BADEN, GERMANY IVERS-LEE AG, BURGDORF, SWITZERLAND NOVARTIS SA, SPARTAN, KEMPTON PARK
Laboratory: FPRC:	NOVARTIS PHARMA STEIN AG, STEIN, SWITZERLAND NOVARTIS PHARMACEUTICA S.A., BARBERA DEL VALLES, BARCELONA, SPAIN IVERS-LEE AG, BURGDORF, SWITZERLAND M&L LABORATORY SERVICES, ORMONDE, JOHANNESBURG
FPRC/FPRR:	NOVARTIS SA, SPARTAN, KEMPTON PARK
Shelf-life:	36 months
Date of registration:	23 JULY 2010

MRF 15

Registration number:	43/7.1.3/0034
Name of medicine:	RINTEZEX 80 TABLET
Dosage form:	TABLET
Active ingredients:	EACH TABLET CONTAINS: VALSARTAN 80,0 mg
Conditions of registration:	1, 2, 3, 4, 5, 6, 7
Applicant:	NOVARTIS SOUTH AFRICA (PTY) LTD
Manufacturer:	NOVARTIS PHARMA STEIN AG, STEIN, SWITZERLAND NOVARTIS PHARMACEUTICA S.A., BARERA DEL VALLES, BARCELONA, SPAIN
Packer:	NOVARTIS PHARMA STEIN AG, STEIN, SWITZERLAND NOVARTIS PHARMACEUTICA S.A., BARERA DEL VALLES, BARCELONA, SPAIN ALLPACK AG, REINACH, SWITZERLAND KONAPHARMA AG, PRATTELN, SWITZERLAND NOVARTIS PHARMA GmbH, WEHR/BADEN, GERMANY IVERS-LEE AG, BURGDORF, SWITZERLAND NOVARTIS SA, SPARTAN, KEMPTON PARK
Laboratory:	FPRC: NOVARTIS PHARMA STEIN AG, STEIN, SWITZERLAND NOVARTIS PHARMACEUTICA S.A., BARERA DEL VALLES, BARCELONA, SPAIN IVERS-LEE AG, BURGDORF, SWITZERLAND M&L LABORATORY SERVICES, ORMONDE, JOHANNESBURG
	FPRC/FPRR: NOVARTIS SA, SPARTAN, KEMPTON PARK
Shelf-life:	36 months
Date of registration:	23 JULY 2010

MRF15

Registration number:	43/7.1.3/0035
Name of medicine:	RINTEZEX 160 TABLET
Dosage form:	TABLET
Active ingredients:	EACH TABLET CONTAINS: VALSARTAN 160,0 mg
Conditions of registration:	1, 2, 3, 4, 5, 6, 7
Applicant:	NOVARTIS SOUTH AFRICA (PTY) LTD
Manufacturer:	NOVARTIS PHARMA STEIN AG, STEIN, SWITZERLAND NOVARTIS PHARMACEUTICA S.A., BARERA DEL VALLES, BARCELONA, SPAIN
Packer:	NOVARTIS PHARMA STEIN AG, STEIN, SWITZERLAND NOVARTIS PHARMACEUTICA S.A., BARERA DEL VALLES, BARCELONA, SPAIN ALLPACK AG, REINACH, SWITZERLAND KONAPHARMA AG, PRATTELN, SWITZERLAND NOVARTIS PHARMA GmbH, WEHR/BADEN, GERMANY IVERS-LEE AG, BURGDORF, SWITZERLAND NOVARTIS SA, SPARTAN,
Laboratory:	FPRC: NOVARTIS PHARMA STEIN AG, STEIN, SWITZERLAND NOVARTIS PHARMACEUTICA S.A., BARERA DEL VALLES, BARCELONA, SPAIN IVERS-LEE AG, BURGDORF, SWITZERLAND M&L LABORATORY SERVICES, ORMONDE, JOHANNESBURG
	FPRC/FPRR: NOVARTIS SA, SPARTAN, KEMPTON PARK
Shelf-life:	36 months
Date of registration:	23 JULY 2010

MRF 15

Registration number:	43/7.1.3/0036
Name of medicine:	ZOMEVEK 80 TABLET
Dosage form:	TABLET
Active ingredients:	EACH TABLET CONTAINS: VALSARTAN 80,0 mg
Conditions of registration:	1, 2, 3, 4, 5, 6, 7
Applicant:	NOVARTIS SOUTH AFRICA (PTY) LTD
Manufacturer:	NOVARTIS PHARMA STEIN AG, STEIN, SWITZERLAND NOVARTIS PHARMACEUTICA S.A., BARERA DEL VALLES, BARCELONA, SPAIN
Packer:	NOVARTIS PHARMA STEIN AG, STEIN, SWITZERLAND NOVARTIS PHARMACEUTICA S.A., BARERA DEL VALLES, BARCELONA, SPAIN ALLPACK AG, REINACH, SWITZERLAND KONAPHARMA AG, PRATTELN, SWITZERLAND NOVARTIS PHARMA GmbH, WEHR/BADEN, GERMANY IVERS-LEE AG, BURGDORF, SWITZERLAND NOVARTIS SA, SPARTAN
Laboratory:	FPRC: NOVARTIS PHARMA STEIN AG, STEIN, SWITZERLAND NOVARTIS PHARMACEUTICA S.A., BARERA DEL VALLES, BARCELONA, SPAIN IVERS-LEE AG, BURGDORF, SWITZERLAND M&L LABORATORY SERVICES, ORMONDE, JOHANNESBURG
	FPRC/FPRR: NOVARTIS SA, SPARTAN, KEMPTON PARK
Shelf-life:	36 months
Date of registration:	23 JULY 2010

MRF 15

Registration number:	43/7.1.3/0037
Name of medicine:	ZOMEVEK 160 TABLET
Dosage form:	TABLET
Active ingredients:	EACH TABLET CONTAINS: VALSARTAN 160,0 mg
Conditions of registration:	1, 2, 3, 4, 5, 6, 7
Applicant:	NOVARTIS SOUTH AFRICA (PTY) LTD
Manufacturer:	NOVARTIS PHARMA STEIN AG, STEIN, SWITZERLAND NOVARTIS PHARMACEUTICA S.A., BARBERA DEL VALLES, BARCELONA, SPAIN
Packer:	NOVARTIS PHARMA STEIN AG, STEIN, SWITZERLAND NOVARTIS PHARMACEUTICA S.A., BARBERA DEL VALLES, BARCELONA, SPAIN ALLPACK AG, REINACH, SWITZERLAND KONAPHARMA AG, PRATTELN, SWITZERLAND NOVARTIS PHARMA GmbH, WEHR/BADEN, GERMANY IVERS-LEE AG, BURGDORF, SWITZERLAND NOVARTIS SA, SPARTAN, KEMPTON PARK
Laboratory: FPRC:	NOVARTIS PHARMA STEIN AG, STEIN, SWITZERLAND NOVARTIS PHARMACEUTICA S.A., BARBERA DEL VALLES, BARCELONA, SPAIN IVERS-LEE AG, BURGDORF, SWITZERLAND M&L LABORATORY SERVICES, ORMONDE, JOHANNESBURG
FPRC/FPRR:	NOVARTIS SA, SPARTAN, KEMPTON PARK
Shelf-life:	36 months
Date of registration:	23 JULY 2010

MRF15

Registration number:	43/7.1.3/0080
Name of medicine:	CO-ZOMEVEK 80/12,5 TABLET
Dosage form:	TABLET
Active ingredients:	EACH TABLET CONTAINS: VALSARTAN 80,0 mg HYDROCHLOROTHIAZIDE 12,5 mg
Conditions of registration:	1, 2, 3, 4, 5, 6, 7
Applicant:	NOVARTIS SOUTH AFRICA (PTY) LTD
Manufacturer:	NOVARTIS PHARMA STEIN AG, STEIN, SWITZERLAND NOVARTIS FARMA S.p.A., TORRE ANNUNZIATA, ITALY
Packer:	NOVARTIS PHARMA STEIN AG, STEIN, SWITZERLAND NOVARTIS FARMA S.p.A., TORRE ANNUNZIATA, ITALY ALLPACK AG, REINACH, SWITZERLAND KONAPHARMA AG, PRATTELN, SWITZERLAND NOVARTIS PHARMA GmbH, WEHR/BADEN, GERMANY IVERS-LEE AG, BURGDORF, SWITZERLAND LAMP S. PROSPERO SpA, SAN PROSPERO, MODENA, ITALY MIPHARM SpA, MILAN, ITALY NOVARTIS SA, SPARTAN, KEMPTON PARK
Laboratory: FPRC:	NOVARTIS PHARMA STEIN AG, STEIN, SWITZERLAND NOVARTIS FARMA S.p.A., TORRE ANNUNZIATA, ITALY NOVARTIS PHARMA GmbH, WEHR/BADEN, GERMANY M&L LABORATORY SERVICES, ORMONDE, JOHANNESBURG
FPRC/FPRR:	NOVARTIS SA, SPARTAN, KEMPTON PARK
Shelf-life:	24 months
Date of registration:	23 JULY 2010

MRF 15

Registration number:	43/7.1.3/0081
Name of medicine:	CO-ZOMEVEK 160/12,5 TABLET
Dosage form:	TABLET
Active ingredients:	EACH TABLET CONTAINS: VALSARTAN 160,0 mg HYDROCHLOROTHIAZIDE 12,5 mg
Conditions of registration:	1, 2, 3, 4, 5, 6, 7
Applicant:	NOVARTIS SOUTH AFRICA (PTY) LTD
Manufacturer:	NOVARTIS PHARMA STEIN AG, STEIN, SWITZERLAND NOVARTIS FARMA S.p.A., TORRE ANNUNZIATA, ITALY
Packer:	NOVARTIS PHARMA STEIN AG, STEIN, SWITZERLAND NOVARTIS FARMA S.p.A., TORRE ANNUNZIATA, ITALY ALLPACK AG, REINACH, SWITZERLAND KONAPHARMA AG, PRATTELN, SWITZERLAND NOVARTIS PHARMA GmbH, WEHR/BADEN, GERMANY IVERS-LEE AG, BURGDORF, SWITZERLAND LAMP S. PROSPERO SpA, SAN PROSPERO, MODENA, ITALY MIPHARM SpA, MILAN, ITALY NOVARTIS SA, SPARTAN, KEMPTON PARK
Laboratory: FPRC:	NOVARTIS PHARMA STEIN AG, STEIN, SWITZERLAND NOVARTIS FARMA S.p.A., TORRE ANNUNZIATA, ITALY NOVARTIS PHARMA GmbH, WEHR/BADEN, GERMANY M&L LABORATORY SERVICES, ORMONDE, JOHANNESBURG
FPRC/FPRR:	NOVARTIS SA, SPARTAN, KEMPTON PARK
Shelf-life:	24 months
Date of registration:	23 JULY 2010

MRF 15

Registration number: 43/7.1.3/0082

Name of medicine: CO-ZOMEVEK 160/25 TABLET

Dosage form: TABLET

Active ingredients: EACH TABLET CONTAINS:
VALSARTAN 160,0 mg
HYDROCHLOROTHIAZIDE
25,0 mg

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

Applicant: NOVARTIS SOUTH AFRICA (PTY) LTD

Manufacturer: NOVARTIS PHARMA STEIN AG, STEIN, SWITZERLAND
NOVARTIS FARMA S.p.A., TORRE ANNUNZIATA, ITALY

Packer: NOVARTIS PHARMA STEIN AG, STEIN, SWITZERLAND
NOVARTIS FARMA S.p.A., TORRE ANNUNZIATA, ITALY
ALLPACK AG, REINACH, SWITZERLAND
KONAPHARMA AG, PRATTELN, SWITZERLAND
NOVARTIS PHARMA GmbH, WEHR/BADEN, GERMANY
IVERS-LEE AG, BURGDORF, SWITZERLAND
LAMP S. PROSPERO SpA, SAN PROSPERO, MODENA, ITALY
MIPHARM SpA, MILAN, ITALY
NOVARTIS SA, SPARTAN, KEMPTON PARK

Laboratory: FPRC: NOVARTIS PHARMA STEIN AG, STEIN, SWITZERLAND
NOVARTIS FARMA S.p.A., TORRE ANNUNZIATA, ITALY
NOVARTIS PHARMA GmbH, WEHR/BADEN, GERMANY
M&L LABORATORY SERVICES, ORMONDE, JOHANNESBURG

FPRC/FPRR: NOVARTIS SA, SPARTAN, KEMPTON PARK

Shelf-life: 24 months

Date of registration: 23 JULY 2010

MRF15

Registration number: 43/7.1.3/0083

Name of medicine: CO-MIGROBEN 80/12,5 TABLET

Dosage form: TABLET

Active ingredients: EACH TABLET CONTAINS:
VALSARTAN 80,0 mg
HYDROCHLOROTHIAZIDE
12,5 mg

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

Applicant: NOVARTIS SOUTH AFRICA (PTY) LTD

Manufacturer: NOVARTIS PHARMA STEIN AG, STEIN, SWITZERLAND
NOVARTIS FARMA S.p.A., TORRE ANNUNZIATA, ITALY

Packer: NOVARTIS PHARMA STEIN AG, STEIN, SWITZERLAND
NOVARTIS FARMA S.p.A., TORRE ANNUNZIATA, ITALY
ALLPACK AG, REINACH, SWITZERLAND
KONAPHARMA AG, PRATTELN, SWITZERLAND
NOVARTIS PHARMA GmbH, WEHR/BADEN, GERMANY
IVERS-LEE AG, BURGDORF, SWITZERLAND
LAMP S. PROSPERO SpA, SAN PROSPERO, MODENA, ITALY
MIPHARM SpA, MILAN, ITALY
NOVARTIS SA, SPARTAN, KEMPTON PARK

Laboratory: FPRC: NOVARTIS PHARMA STEIN AG, STEIN, SWITZERLAND
NOVARTIS FARMA S.p.A., TORRE ANNUNZIATA, ITALY
NOVARTIS PHARMA GmbH, WEHR/BADEN, GERMANY
M&L LABORATORY SERVICES, ORMONDE, JOHANNESBURG

FPRC/FPRR: NOVARTIS SA, SPARTAN, KEMPTON PARK

Shelf-life: 24 months

Date of registration: 23 JULY 2010

MRF 15

Registration number: 43/7.1.3/0084

Name of medicine: CO-MIGROBEN 160/12,5 TABLET

Dosage form: TABLET

Active ingredients: EACH TABLET CONTAINS:
VALSARTAN 160,0 mg
HYDROCHLOROTHIAZIDE
12,5 mg

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

Applicant: NOVARTIS SOUTH AFRICA (PTY) LTD

Manufacturer: NOVARTIS PHARMA STEIN AG, STEIN, SWITZERLAND
NOVARTIS FARMA S.p.A., TORRE ANNUNZIATA, ITALY

Packer: NOVARTIS PHARMA STEIN AG, STEIN, SWITZERLAND
NOVARTIS FARMA S.p.A., TORRE ANNUNZIATA, ITALY
ALLPACK AG, REINACH, SWITZERLAND
KONAPHARMA AG, PRATTELN, SWITZERLAND
NOVARTIS PHARMA GmbH, WEHR/BADEN, GERMANY
IVERS-LEE AG, BURGDORF, SWITZERLAND
LAMP S. PROSPERO SpA, SAN PROSPERO, MODENA, ITALY
MIPHARM SpA, MILAN, ITALY
NOVARTIS SA, SPARTAN, KEMPTON PARK

Laboratory: FPRC: NOVARTIS PHARMA STEIN AG, STEIN, SWITZERLAND
NOVARTIS FARMA S.p.A., TORRE ANNUNZIATA, ITALY
NOVARTIS PHARMA GmbH, WEHR/BADEN, GERMANY
M&L LABORATORY SERVICES, ORMONDE, JOHANNESBURG

FPRC/FPRR: NOVARTIS SA, SPARTAN, KEMPTON PARK

Shelf-life: 24 months

Date of registration: 23 JULY 2010

MRF 15

Registration number:	43/7.1.3/0085
Name of medicine:	CO-MIGROBEN 160/25 TABLET
Dosage form:	TABLET
Active ingredients:	EACH TABLET CONTAINS: VALSARTAN 160,0 mg HYDROCHLOROTHIAZIDE 25,0 mg
Conditions of registration:	1, 2, 3, 4, 5, 6, 7
Applicant:	NOVARTIS SOUTH AFRICA (PTY) LTD
Manufacturer:	NOVARTIS PHARMA STEIN AG, STEIN, SWITZERLAND NOVARTIS FARMA S.p.A., TORRE ANNUNZIATA, ITALY
Packer:	NOVARTIS PHARMA STEIN AG, STEIN, SWITZERLAND NOVARTIS FARMA S.p.A., TORRE ANNUNZIATA, ITALY ALLPACK AG, REINACH, SWITZERLAND KONAPHARMA AG, PRATTELN, SWITZERLAND NOVARTIS PHARMA GmbH, WEHR/BADEN, GERMANY IVERS-LEE AG, BURGDORF, SWITZERLAND LAMP S. PROSPERO SpA, SAN PROSPERO, MODENA, ITALY MIPHARM SpA, MILAN, ITALY NOVARTIS SA, SPARTAN, KEMPTON PARK
Laboratory: FPRC:	NOVARTIS PHARMA STEIN AG, STEIN, SWITZERLAND NOVARTIS FARMA S.p.A., TORRE ANNUNZIATA, ITALY NOVARTIS PHARMA GmbH, WEHR/BADEN, GERMANY M&L LABORATORY SERVICES, ORMONDE, JOHANNESBURG
FPRC/FPRR:	NOVARTIS SA, SPARTAN, KEMPTON PARK
Shelf-life:	24 months
Date of registration:	23 JULY 2010

MRF15

Registration number:	43/7.1.3/0087
Name of medicine:	RINTEZEX CO 80/12,5 TABLET
Dosage form:	TABLET
Active ingredients:	EACH TABLET CONTAINS: VALSARTAN 80,0 mg HYDROCHLOROTHIAZIDE 12,5 mg
Conditions of registration:	1, 2, 3, 4, 5, 6, 7
Applicant:	NOVARTIS SOUTH AFRICA (PTY) LTD
Manufacturer:	NOVARTIS PHARMA STEIN AG, STEIN, SWITZERLAND NOVARTIS FARMA S.p.A., TORRE ANNUNZIATA, ITALY
Packer:	NOVARTIS PHARMA STEIN AG, STEIN, SWITZERLAND NOVARTIS FARMA S.p.A., TORRE ANNUNZIATA, ITALY ALLPACK AG, REINACH, SWITZERLAND KONAPHARMA AG, PRATTELN, SWITZERLAND NOVARTIS PHARMA GmbH, WEHR/BADEN, GERMANY IVERS-LEE AG, BURGDORF, SWITZERLAND LAMP S. PROSPERO SpA, SAN PROSPERO, MODENA, ITALY MIPHARM SpA, MILAN, ITALY NOVARTIS SA, SPARTAN, KEMPTON PARK
Laboratory: FPRC:	NOVARTIS PHARMA STEIN AG, STEIN, SWITZERLAND NOVARTIS FARMA S.p.A., TORRE ANNUNZIATA, ITALY NOVARTIS PHARMA GmbH, WEHR/BADEN, GERMANY M&L LABORATORY SERVICES, ORMONDE, JOHANNESBURG
FPRC/FPRR:	NOVARTIS SA, SPARTAN, KEMPTON PARK
Shelf-life:	24 months
Date of registration:	23 JULY 2010

MRF 15

Registration number:	43/7.1.3/0088
Name of medicine:	RINTEZEX CO 160/12,5 TABLET
Dosage form:	TABLET
Active ingredients:	EACH TABLET CONTAINS: VALSARTAN 160,0 mg HYDROCHLOROTHIAZIDE 12,5 mg
Conditions of registration:	1, 2, 3, 4, 5, 6, 7
Applicant:	NOVARTIS SOUTH AFRICA (PTY) LTD
Manufacturer:	NOVARTIS PHARMA STEIN AG, STEIN, SWITZERLAND NOVARTIS FARMA S.p.A., TORRE ANNUNZIATA, ITALY
Packer:	NOVARTIS PHARMA STEIN AG, STEIN, SWITZERLAND NOVARTIS FARMA S.p.A., TORRE ANNUNZIATA, ITALY ALLPACK AG, REINACH, SWITZERLAND KONAPHARMA AG, PRATTELN, SWITZERLAND NOVARTIS PHARMA GmbH, WEHR/BADEN, GERMANY IVERS-LEE AG, BURGDORF, SWITZERLAND LAMP S. PROSPERO SpA, SAN PROSPERO, MODENA, ITALY MIPHARM SpA, MILAN, ITALY NOVARTIS SA, SPARTAN, KEMPTON PARK
Laboratory: FPRC:	NOVARTIS PHARMA STEIN AG, STEIN, SWITZERLAND NOVARTIS FARMA S.p.A., TORRE ANNUNZIATA, ITALY NOVARTIS PHARMA GmbH, WEHR/BADEN, GERMANY M&L LABORATORY SERVICES, ORMONDE, JOHANNESBURG
FPRC/FPRR:	NOVARTIS SA, SPARTAN, KEMPTON PARK
Shelf-life:	24 months
Date of registration:	23 JULY 2010

MRF 15

Registration number: 43/7.1.3/0089
Name of medicine: RINTEZEX CO 160/25 TABLET
Dosage form: TABLET
Active ingredients: EACH TABLET CONTAINS:
VALSARTAN 160,0 mg
HYDROCHLOROTHIAZIDE
25,0 mg

Conditions of registration: 1, 2, 3, 4, 5, 6, 7
Applicant: NOVARTIS SOUTH AFRICA
(PTY) LTD
Manufacturer: NOVARTIS PHARMA STEIN AG,
STEIN, SWITZERLAND
NOVARTIS FARMA S.p.A.,
TORRE ANNUNZIATA, ITALY

Packer: NOVARTIS PHARMA STEIN AG,
STEIN, SWITZERLAND
NOVARTIS FARMA S.p.A.,
TORRE ANNUNZIATA, ITALY
ALLPACK AG, REINACH,
SWITZERLAND
KONAPHARMA AG, PRATTELN,
SWITZERLAND
NOVARTIS PHARMA GmbH,
WEHR/BADEN, GERMANY
IVERS-LEE AG, BURGDORF,
SWITZERLAND
LAMP S. PROSPERO SpA, SAN
PROSPERO, MODENA, ITALY
MIPHARM SpA, MILAN, ITALY
NOVARTIS SA, SPARTAN,
KEMPTON PARK

Laboratory: FPRC: NOVARTIS PHARMA STEIN AG,
STEIN, SWITZERLAND
NOVARTIS FARMA S.p.A.,
TORRE ANNUNZIATA, ITALY
NOVARTIS PHARMA GmbH,
WEHR/BADEN, GERMANY
M&L LABORATORY SERVICES,
ORMONDE, JOHANNESBURG

FPRC/FPRR: NOVARTIS SA, SPARTAN,
KEMPTON PARK

Shelf-life: 24 months
Date of registration: 23 JULY 2010

MRF15

Registration number: 43/21.2/0605
Name of medicine: DRL GLIMEPIRIDE 1
Dosage form: TABLET
Active ingredients: EACH TABLET CONTAINS:
GLIMEPIRIDE 1,0 mg

Conditions of registration: 1, 2, 3, 4, 5, 6, 7, 8
Applicant: DR REDDY'S LABORATORIES
(PTY) LTD
Manufacturer: DR REDDY'S LABORATORIES
LTD, QUTHUBULLAPUR
MANDAL, RANGA REDDY
DISTRICT, ANDHRA PRADESH,
INDIA

Packer: DR REDDY'S LABORATORIES
LTD, QUTHUBULLAPUR
MANDAL, RANGA REDDY
DISTRICT, ANDHRA PRADESH,
INDIA
DRA PHARMACEUTICALS,
IRENE, CENTURION
TECHNIKON LABORATORIES,
ROBERTVILLE, FLORIDA
DIVPHARM MANUFACTURING
& PACKAGING, LONGDALE,
JOHANNESBURG

Laboratory: FPRC: DR REDDY'S LABORATORIES
LTD, QUTHUBULLAPUR
MANDAL, RANGA REDDY
DISTRICT, ANDHRA PRADESH,
INDIA
RESEARCH INSTITUTE FOR
INDUSTRIAL PHARMACY,
NORTH-WEST UNIVERSITY,
POTCHEFSTROOM

FPRR: DR REDDY'S LABORATORIES,
MURRAYFIELD, PRETORIA

Shelf-life: 24 months (Provisional)
Date of registration: 23 JULY 2010

MRF 15

Registration number: 43/21.2/0606
Name of medicine: DRL GLIMEPIRIDE 2
Dosage form: TABLET
Active ingredients: EACH TABLET CONTAINS:
GLIMEPIRIDE 2,0 mg

Conditions of registration: 1, 2, 3, 4, 5, 6, 7, 8
Applicant: DR REDDY'S LABORATORIES (PTY)
LTD
Manufacturer: DR REDDY'S LABORATORIES LTD,
QUTHUBULLAPUR MANDAL, RANGA
REDDY DISTRICT, ANDHRA
PRADESH, INDIA

Packer: DR REDDY'S LABORATORIES LTD,
QUTHUBULLAPUR MANDAL, RANGA
REDDY DISTRICT, ANDHRA
PRADESH, INDIA
DRA PHARMACEUTICALS, IRENE,
CENTURION
TECHNIKON LABORATORIES,
ROBERTVILLE, FLORIDA
DIVPHARM MANUFACTURING &
PACKAGING, LONGDALE,
JOHANNESBURG

Laboratory: FPRC:: DR REDDY'S LABORATORIES LTD,
QUTHUBULLAPUR MANDAL, RANGA
REDDY DISTRICT, ANDHRA
PRADESH, INDIA
RESEARCH INSTITUTE FOR
INDUSTRIAL PHARMACY, NORTH-
WEST UNIVERSITY,
POTCHEFSTROOM

FPRR: DR REDDY'S LABORATORIES,
MURRAYFIELD, PRETORIA

Shelf-life: 24 months (Provisional)
Date of registration: 23 JULY 2010

MRF 15

Registration number: 43/21.2/0607
 Name of medicine: DRL GLIMEPIRIDE 4
 Dosage form: TABLET
 Active ingredients: EACH TABLET CONTAINS:
 GLIMEPIRIDE 4,0 mg
 Conditions of registration: 1, 2, 3, 4, 5, 6, 7, 8
 Applicant: DR REDDY'S LABORATORIES (PTY) LTD
 Manufacturer: DR REDDY'S LABORATORIES LTD, QUTHUBULLAPUR MANDAL, RANGA REDDY DISTRICT, ANDHRA PRADESH, INDIA
 Packer: DR REDDY'S LABORATORIES LTD, QUTHUBULLAPUR MANDAL, RANGA REDDY DISTRICT, ANDHRA PRADESH, INDIA
 DRA PHARMACEUTICALS, IRENE, CENTURION TECHNIKON LABORATORIES, ROBERTVILLE, FLORIDA DIVPHARM MANUFACTURING & PACKAGING, LONGDALE, JOHANNESBURG
 Laboratory: FPRC: DR REDDY'S LABORATORIES LTD, QUTHUBULLAPUR MANDAL, RANGA REDDY DISTRICT, ANDHRA PRADESH, INDIA
 RESEARCH INSTITUTE FOR INDUSTRIAL PHARMACY, NORTH-WEST UNIVERSITY, POTCHEFSTROOM
 FPRR: DR REDDY'S LABORATORIES, MURRAYFIELD, PRETORIA
 Shelf-life: 24 months (Provisional)
 Date of registration: 23 JULY 2010

MRF15

Registration number: 43/20.2.8/0832
 Name of medicine: ASPEN LAMIVUDINE 300 mg
 Dosage form: TABLET
 Active ingredients: EACH TABLET CONTAINS:
 LAMIVUDINE 300,0 mg
 Conditions of registration: 1, 2, 3, 4, 5, 6, 7, 8
 Applicant: PHARMACARE LIMITED
 Manufacturer: STRIDES ARCOLAB LTD, ANEKAL TALUK, BANGALORE, INDIA
 ASPED OSD, GIBAUD ROAD, PORT ELIZABETH
 Packer: STRIDES ARCOLAB LTD, ANEKAL TALUK, BANGALORE, INDIA
 ASPED OSD, GIBAUD ROAD, PORT ELIZABETH
 Laboratory: FPRC: STRIDES ARCOLAB LTD, ANEKAL TALUK, BANGALORE, INDIA
 M&L LABORATORY SERVICES, ORMONDE, JOHANNESBURG RESEARCH INSTITUTE FOR INDUSTRIAL PHARMACY, NORTH-WEST UNIVERSITY, POTCHEFSTROOM
 SABS PHARMACEUTICAL CHEMISTRY LABORATORY, GROENKLOOF, PRETORIA
 FPRC/FPRR: ASPEN OSD, GIBAUD ROAD, PORT ELIZABETH
 FPRR: PHARMACARE LTD, WOODMEAD, SANDTON
 Shelf-life: 24 months (Provisional)
 Date of registration: 23 JULY 2010

MRF 15

Registration number: 44/20.2.8/0294
 Name of medicine: ADCO LAMIVUDINE 300 mg TABLETS
 Dosage form: TABLET
 Active ingredients: EACH TABLET CONTAINS:
 LAMIVUDINE 300,0 mg
 Conditions of registration: 1, 2, 3, 4, 5, 6, 7
 Applicant: ADCOCK INGRAM LIMITED
 Manufacturer: HETERO DRUGS, JEEDIMETLA, HYDERABAD, INDIA
 ADCOCK INGRAM HEALTHCARE, WADEVILLE, GERMISTON
 Packer: HETERO DRUGS, JEEDIMETLA, HYDERABAD, INDIA
 ADCOCK INGRAM HEALTHCARE, WADEVILLE, GERMISTON
 Laboratory: FPRC: HETERO DRUGS, JEEDIMETLA, HYDERABAD, INDIA
 FPRC/FPRR: ADCOCK INGRAM HEALTHCARE, WADEVILLE, GERMISTON
 ADCOCK INGRAM LTD, AEROTON, JOHANNESBURG
 FPRR: ADCOCK INGRAM LTD, ERAND GARDENS, MIDRAND
 Shelf-life: 24 months
 Date of registration: 23 JULY 2010

MRF 15

Registration number: 44/20.2.8/0285
 Name of medicine: RETLAM 300 mg TABLETS
 Dosage form: TABLET
 Active ingredients: EACH TABLET CONTAINS:
 LAMIVUDINE 300,0 mg

Conditions of registration: 1, 2, 3, 4, 5, 6, 7
 Applicant: ADCOCK INGRAM LIMITED
 Manufacturer: HETERO DRUGS, JEEDIMETLA,
 HYDERABAD, INDIA
 ADCOCK INGRAM HEALTHCARE,
 WADEVILLE, GERMISTON

Packer: HETERO DRUGS, JEEDIMETLA,
 HYDERABAD, INDIA
 ADCOCK INGRAM HEALTHCARE,
 WADEVILLE, GERMISTON

Laboratory: FPRC: HETERO DRUGS, JEEDIMETLA,

MRF15

Registration number: 44/20.2.8/0296
 Name of medicine: LAVOS 300 mg TABLETS
 Dosage form: TABLET
 Active ingredients: EACH TABLET CONTAINS:
 LAMIVUDINE 300,0 mg

Conditions of registration: 1, 2, 3, 4, 5, 6, 7
 Applicant: ADCOCK INGRAM LIMITED
 Manufacturer: HETERO DRUGS, JEEDIMETLA,
 HYDERABAD, INDIA
 ADCOCK INGRAM HEALTHCARE,
 WADEVILLE, GERMISTON

Packer: HETERO DRUGS, JEEDIMETLA,
 HYDERABAD, INDIA
 ADCOCK INGRAM HEALTHCARE,
 WADEVILLE, GERMISTON

Laboratory: FPRC: HETERO DRUGS, JEEDIMETLA,

MRF 15

Registration number: 44/30.1/0897
 Name of medicine: AREPANDRIX H1N1
 Dosage form: INJECTION
 Active ingredients: EACH 0,5 ml DOSE CONTAINS:
 SPLIT INFLUENZA VIRUS
 A/CALIFORNIA/7/2009(H1N1)
 V-LIKE VIRUS 3,75 µg

Conditions of registration: 1, 2, 3, 4, 5, 6, 7
 Applicant: GLAXOSMITHKLINE S.A. (PTY) LTD
 Manufacturer: GLAXOSMITHKLINE BIOLOGICALS NORTH
 AMERICA, SAINTE-FOY, QUEBEC,
 CANADA
 GLAXOSMITHKLINE BIOLOGICALS
 MANUFACTURING SA, RIXENSART,
 BELGIUM
 GLAXOSMITHKLINE BIOLOGICALS
 MANUFACTURING SA, WAVRE, BELGIUM
 GLAXOSMITHKLINE BIOLOGICALS, SAINT
 AMAND LES EAUX, FRANCE
 GLAXOSMITHKLINE BIOLOGICALS NORTH
 AMERICA, MARIETTA, PA, USA
 GLAXO WELCOME OPERATIONS UK,
 BARNARD CASTLE, DURHAM, UK
 VETTER PHARMA INTERNATIONAL GmbH,
 MOOSWIESEN, RAVENSBURG, GERMANY
 DSM, GREENVILLE, NC, USA
 BAXTER PHARMACEUTICAL SOLUTIONS,
 BLOOMINGTON, INDIANA, USA
 HOSPIRA INC, MCPHERSON, KANSAS,
 USA

Packer: GLAXOSMITHKLINE BIOLOGICALS NORTH
 AMERICA, SAINTE-FOY, QUEBEC,
 CANADA
 GLAXOSMITHKLINE BIOLOGICALS
 MANUFACTURING SA, RIXENSART,
 BELGIUM
 GLAXOSMITHKLINE BIOLOGICALS
 MANUFACTURING SA, WAVRE, BELGIUM
 GLAXOSMITHKLINE BIOLOGICALS NORTH
 AMERICA, MARIETTA, PA, USA
 GLAXO WELCOME OPERATIONS UK,
 BARNARD CASTLE, DURHAM, UK
 VETTER PHAMRA INTERNATIONAL GmbH,
 MOOSWIESEN, RAVENSBURG, GERMANY
 HOLLISTER-STIER LABORATORIES,
 SPOKANE, WA, USA
 DSM, GREENVILLE, NC, USA
 PIERRE FABRE MEDICAMENT
 PRODUCTION, CHATEAURENARD,
 FRANCE
 PIERRE FABRE MEDICAMENT
 PRODUCTION, CHEMIN DE MAZEROLLES,
 IDRON, FRANCE
 PIERRE FABRE MEDICAMENT
 PRODUCTION, IDRON FRANCE
 BAXTER PHARMACEUTICAL SOLUTIONS,
 BLOOMINGTON, INDIANA, USA
 HOSPIRA INC, MCPHERSON, KANSAS,
 USA
 GLAXOSMITHKLINE S.A., EPPING, CAPE
 TOWN

Laboratory: FPRC: GLAXOSMITHKLINE BIOLOGICALS NORTH

HYDERABAD, INDIA

FPRC/FPRR: ADCOCK INGRAM HEALTHCARE,
WADEVILLE, GERMISTON
ADCOCK INGRAM LTD, AEROTON,
JOHANNESBURG

FPRR: ADCOCK INGRAM LTD, ERAND
GARDENS, MIDRAND

Shelf-life: 24 months

Date of registration: 23 JULY 2010

HYDERABAD, INDIA

FPRC/FPRR: ADCOCK INGRAM HEALTHCARE,
WADEVILLE, GERMISTON
ADCOCK INGRAM LTD, AEROTON,
JOHANNESBURG

FPRR: ADCOCK INGRAM LTD, ERAND
GARDENS, MIDRAND

Shelf-life: 24 months

Date of registration: 23 JULY 2010

AMERICA, SAINTE-FOY, QUEBEC,
CANADA
GLAXOSMITHKLINE BIOLOGICALS
MANUFACTURING SA, RIXENSART,
BELGIUM
GLAXOSMITHKLINE BIOLOGICALS
MANUFACTURING SA, WAVRE, BELGIUM
GLAXOSMITHKLINE BIOLOGICALS, SAINT
AMAND LES EAUX, FRANCE
GLAXOSMITHKLINE BIOLOGICALS NORTH
AMERICA, MARIETTA, PA, USA
GLAXO WELCOME OPERATIONS UK,
BARNARD CASTLE, DURHAM, UK
VETTER PHARMA INTERNATIONAL GmbH,
MOOSWIESEN, RAVENSBURG, GERMANY
HOLLISTER-STIER LABORATORIES,
SPOKANE, WA, USA
DSM, GREENVILLE, NC, USA
PIERRE FABRE MEDICAMENT
PRODUCTION, CHATEAURENARD,
FRANCE
PIERRE FABRE MEDICAMENT
PRODUCTION, CHEMIN DE MAZEROLLES,
IDRON, FRANCE
PIERRE FABRE MEDICAMENT
PRODUCTION, IDRON FRANCE
BAXTER PHARMACEUTICAL SOLUTIONS,
BLOOMINGTON, INDIANA, USA
HOSPIRA INC, MCPHERSON, KANSAS,
USA

FPRC/FPRR: GLAXOSMITHKLINE S.A., EPPING, CAPE
TOWN

Shelf-life: 24 months

Date of registration: 23 JULY 2010