

20 March 2017
EMA/CHMP/184758/2017
Inspections, Human Medicines Pharmacovigilance and Committees Division

Committee for medicinal products for human use (CHMP)

Draft agenda for the meeting on 20-23 March 2017

Chair: Tomas Salmonson – Vice-Chair: Harald Enzmann

20 March 2017, 13:00 – 19:30, room 2A

21 March 2017, 08:30 – 19:30, room 2A

22 March 2017, 08:30 – 19:30, room 2A

23 March 2017, 08:30 – 15:00, room 2A

Health and safety information

In accordance with the Agency's health and safety policy, delegates are to be briefed on health, safety and emergency information and procedures prior to the start of the meeting.

Disclaimers

Some of the information contained in this agenda is considered commercially confidential or sensitive and therefore not disclosed. With regard to intended therapeutic indications or procedure scopes listed against products, it must be noted that these may not reflect the full wording proposed by applicants and may also vary during the course of the review. Additional details on some of these procedures will be published in the [CHMP meeting highlights](#) once the procedures are finalised and start of referrals will also be available.

Of note, this agenda is a working document primarily designed for CHMP members and the work the Committee undertakes.

Note on access to documents

Some documents mentioned in the agenda cannot be released at present following a request for access to documents within the framework of Regulation (EC) No 1049/2001 as they are subject to on-going procedures for which a final decision has not yet been adopted. They will become public when adopted or considered public according to the principles stated in the Agency policy on access to documents (EMA/127362/2006).

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1. Introduction

1.1. Welcome and declarations of interest of members, alternates and experts

Pre-meeting list of participants and restrictions in relation to declarations of interests applicable to the items of the agenda for the CHMP plenary session to be held 20-23 March 2017. See March 2017 CHMP minutes (to be published post April 2017 CHMP meeting).

1.2. Adoption of agenda

CHMP agenda for 20-23 March 2017

1.3. Adoption of the minutes

CHMP minutes for 20-23 February 2017.

2. Oral Explanations

2.1. Pre-authorisation procedure oral explanations

2.1.1. - dinutuximab beta - Orphan - EMEA/H/C/003918

APEIRON Biologics AG; treatment of neuroblastoma

Scope: Oral explanation

Action: Possible oral explanation to be held on 21 March 2017 at time 09:00

List of Outstanding Issues adopted on 23.02.2017, 13.10.2016, 26.05.2016. List of Questions adopted on 24.09.2015.

2.1.2. - nusinersen - Orphan - EMEA/H/C/004312

Accelerated assessment

Biogen Idec Ltd; for the treatment of Spinal Muscular Atrophy (SMA)

Scope: Oral explanation/Day 180 list of outstanding issue

Action: Possible oral explanation to be held on 22 March 2017 at time 11:00

List of Questions adopted on 24.01.2017.

2.2. Re-examination procedure oral explanations

2.3. Post-authorisation procedure oral explanations

2.4. Referral procedure oral explanations

3. Initial applications

3.1. Initial applications; Opinions

3.1.1. - fluciclovine (18F) - EMEA/H/C/004197

diagnostic agent for PET of adult men with suspected recurrence of prostate cancer

Scope: Opinion

Action: For adoption

List of Outstanding Issues adopted on 15.12.2016. List of Questions adopted on 28.04.2016.

3.1.2. - ivabradine - EMEA/H/C/004241

treatment of angina pectoris

Scope: Opinion

Action: For adoption

List of Outstanding Issues adopted on 15.12.2016. List of Questions adopted on 01.04.2016.

3.1.3. - pentosan polysulfate sodium - Orphan - EMEA/H/C/004246

bene-Arzneimittel GmbH; treatment of Interstitial Cystitis

Scope: Opinion

Action: For adoption

List of Outstanding Issues adopted on 10.11.2016. List of Questions adopted on 23.06.2016.

3.1.4. - nonacog beta pegol - Orphan - EMEA/H/C/004178

Novo Nordisk A/S; treatment of haemophilia B

Scope: Opinion

Action: For adoption

Oral explanation held on 15.09.2016. List of Outstanding Issues adopted on 15.09.2016.
List of Questions adopted on 26.05.2016.

3.1.5. - meningococcal group B vaccine (recombinant, component, adsorbed) -
EMEA/H/C/004051

prevent invasive meningococcal disease caused by *Neisseria meningitidis* serogroup B

Scope: Opinion

Action: For adoption

List of Outstanding Issues adopted on 26.01.2017. List of Questions adopted on
15.09.2016.

**3.2. Initial applications; List of outstanding issues (Day 180; Day 120 for
procedures with accelerated assessment timetable)**

3.2.1. - dengue tetravalent vaccine (live, attenuated) - EMEA/H/C/004171

indicated for the prevention of dengue disease caused by dengue virus serotypes 1, 2, 3
and 4

Scope: Day 180 list of outstanding issue

Action: For adoption

List of Questions adopted on 21.07.2016.

3.2.2. - efavirenz / emtricitabine / tenofovir disoproxil - EMEA/H/C/004250

treatment of HIV-1 infection

Scope: Day 180 list of outstanding issue

Action: For adoption

List of Questions adopted on 10.11.2016.

3.2.3. - alpha-1-antitrypsin - Orphan - EMEA/H/C/003934

Kamada BioPharma Limited at Fieldfisher LLP; treatment and maintenance therapy of adult
patients with congenital deficiency of alpha-1 antitrypsin and lung disease with clinical
evidence of emphysema and airway obstruction (FEV1/SVC<70%)

Scope: Day 180 list of outstanding issue

Action: For adoption

List of Questions adopted on 21.07.2016.

3.2.4. - nitisinone - EMEA/H/C/004281

treatment of hepatorenal tyrosinemia type 1

Scope: Day 180 list of outstanding issue

Action: For adoption

List of Questions adopted on 21.07.2016.

3.2.5. - ocrelizumab - EMEA/H/C/004043

treatment of multiple sclerosis

Scope: Day 180 list of outstanding issue

Action: For adoption

List of Questions adopted on 15.09.2016.

3.2.6. - etirinotecan pegol - EMEA/H/C/003874

treatment of breast cancer with brain metastases

Scope: Day 180 list of outstanding issue

Action: For adoption

List of Questions adopted on 10.11.2016.

3.3. Initial applications; List of questions (Day 120; Day 90 for procedures with accelerated assessment timetable)

3.3.1. - adalimumab - EMEA/H/C/004319

treatment of rheumatoid arthritis, axial spondyloarthritis, psoriasis, hidradenitis suppurativa (HS), Crohn's disease, ulcerative colitis and uveitis.

Scope: Day 120 list of questions

Action: For adoption

3.3.2. - darunavir - EMEA/H/C/004273

treatment of HIV-1 infection

Scope: Day 120 list of questions

Action: For adoption

3.3.3. - dupilumab - EMEA/H/C/004390

treatment of moderate-to-severe atopic dermatitis

Scope: Day 120 list of questions

Action: For adoption

3.3.4. - naloxone - EMEA/H/C/004325

for emergency use for known or suspected opioid overdose as manifested by respiratory and/or central nervous system depression

Scope: Day 120 list of questions

Action: For adoption

3.3.5. - ciclosporin - EMEA/H/C/004229

for the treatment of moderate dry eye disease in adults

Scope: Day 120 list of questions

Action: For adoption

3.3.6. - rucaparib - Orphan - EMEA/H/C/004272

Clovis Oncology UK Ltd; treatment of ovarian cancer

Scope: Day 120 list of questions

Action: For adoption

3.3.7. - pegfilgrastim - EMEA/H/C/004413

treatment of neutropenia

Scope: Day 120 list of questions

Action: For adoption

3.4. Update on on-going initial applications for Centralised procedure

3.4.1. - entecavir - EMEA/H/C/004458

treatment of chronic hepatitis B virus infection

Scope: Request by the applicant for an extension to the clock stop to respond to the day 120 list of questions adopted on 15 December 2016. Adopted by written procedure on 9 March 2017.

Action: For information

List of questions adopted on 15.12.2016.

3.4.2. - iloperidone - EMEA/H/C/004149

treatment of schizophrenia

Scope: Letter from the applicant dated 8 March 2017 requesting an extension of clock stop to respond to the List of Outstanding Issues adopted on 23 February 2017.

Action: For adoption

List of outstanding issue adopted on 23.02.2017, List of Questions adopted on 28.04.2016.

3.4.3. - brodalumab - EMEA/H/C/003959

moderate to severe plaque psoriasis

Scope: PRAC advice

Action: For discussion

List of Outstanding Issues adopted on 26.01.2017, 10.11.2016, 15.09.2016. List of Questions adopted on 01.04.2016.

3.4.4. - masitinib - Orphan - EMEA/H/C/004398

AB Science; treatment of amyotrophic lateral sclerosis

Scope: Letter from the applicant dated 16 March 2017 requesting an extension of clock-stop to respond to the List of Questions adopted on 26 January 2017

Action: For adoption

List of Questions adopted on 26.01.2017

3.4.5. - d-biotin - EMEA/H/C/004153

treatment of progressive multiple sclerosis (primary or secondary)

Scope: Letter from the applicant dated 28 February 2017 requesting an extension of clock stop to respond to the List of Questions adopted on 15 December 2016

Action: For adoption

List of Questions adopted on 15.12.2016.

3.4.6. - cariprazine - EMEA/H/C/002770

treatment of schizophrenia

Scope: Letter from the applicant dated 14 March 2017 requesting an extension of clock stop to respond to the List of Outstanding Issues adopted on 23 February 2017.

Action: For adoption

List of Outstanding Issues adopted on 23.02.2017. List of Questions adopted on 21.07.2016.

3.4.7. - abaloparatide - EMEA/H/C/004157

treatment of osteoporosis

Scope: extension of the clock stop to respond to the List of Outstanding Issues adopted on 15 December 2016

Action: For adoption

List of Outstanding Issues adopted on 15.12.2016. List of Questions adopted on 01.04.2016.

3.5. Re-examination of initial application procedures under Article 9(2) of Regulation no 726/2004

3.6. Initial applications in the decision-making phase

3.7. Withdrawals of initial marketing authorisation application

3.7.1. Blectifor - caffeine citrate - Orphan - EMEA/H/C/004100

Viridian Pharma Ltd; indicated in preterm neonates for the prevention of bronchopulmonary dysplasia

Rapporteur: Milena Stain, Co-Rapporteur: Agnes Gyurasics, PRAC Rapporteur: Ana Sofia Diniz Martins

Scope: Withdrawal of initial marketing authorisation application

Action: For information

3.7.2. Enpaxiq - pacritinib - Orphan - EMEA/H/C/004193

CTI Life Sciences Limited; treatment of myelofibrosis

Scope: Withdrawal of initial marketing authorisation application

Action: For information

List of Questions adopted on 15.09.2016.

4. Extension of marketing authorisation according to Annex I of Commission Regulation (EC) No 1234/2008

4.1. Extension of marketing authorisation according to Annex I of Commission Regulation (EC) No 1234/2008; Opinion

4.1.1. Benepali - etanercept - EMEA/H/C/004007/X/0016

Samsung Bioepis UK Limited (SBUK)

Rapporteur: Andrea Laslop, PRAC Rapporteur: Patrick Batty

Scope: "To add a new strength of 25 mg solution for injection in pre-filled syringe."

Action: For adoption

List of Questions adopted on 15.12.2016.

4.1.2. Brilique - ticagrelor - EMEA/H/C/001241/X/0034

AstraZeneca AB

Rapporteur: Johann Lodewijk Hillege

Scope: "To add new pharmaceutical form (orodispersible tablets 90 mg) to the currently approved presentations for Brilique."

Action: For adoption

List of Outstanding Issues adopted on 26.01.2017. List of Questions adopted on 15.09.2016.

4.2. Extension of marketing authorisation according to Annex I of Commission Regulation (EC) No 1234/2008; Day 180 list of outstanding issues

4.2.1. Mimpara - cinacalcet - EMEA/H/C/000570/X/0055/G

Amgen Europe B.V.

Rapporteur: Kristina Dunder, Co-Rapporteur: Andrea Laslop, PRAC Rapporteur: Ulla Wändel Liminga

Scope: "Extension application to introduce a new pharmaceutical form associated with new strengths (1 mg, 2.5 mg and 5 mg hard capsules) grouped with a type II variation (C.1.6.a) to include paediatric use in the approved indication.

As a consequence, sections 4.2 and 4.4 of the SmPC are updated to detail posology in paediatric patients and to update the safety information, respectively.

The Package Leaflet and Labelling are updated in accordance.

In addition, the Marketing authorisation holder (MAH) took the opportunity to update the list of local representatives in the Package Leaflet.

Furthermore, the PI is brought in line with the latest QRD template version 10."

Action: For adoption

List of Questions adopted on 13.10.2016.

4.3. Extension of marketing authorisation according to Annex I of Commission Regulation (EC) No 1234/2008; Day 120 List of question

4.3.1. Tasigna - nilotinib - Orphan - EMEA/H/C/000798/X/0088/G

Novartis Europarm Ltd

Rapporteur: Sinan B. Sarac, Co-Rapporteur: Harald Enzmann, PRAC Rapporteur: Doris Stenver

Scope: "Extension of Indication to include treatment of paediatric patients with newly diagnosed Philadelphia chromosome-positive chronic myelogenous leukemia in chronic phase (Ph+ CML-CP), or with Ph+ CML-CP resistant or intolerant to prior therapy including imatinib, based on results from two clinical studies in paediatric patients conducted in accordance with the approved Tasigna Paediatric Investigation Plan (PIP), the Phase I PK study CAMN107A2120 and the Phase II safety and efficacy study CAMN107A2203. An updated RMP version 18.0 was provided as part of the application.

Extension application to add a new strength of 50mg hard capsules.

In addition, the applicant proposes to merge the SmPCs for the 50 mg and 200 mg strengths."

Action: For adoption

4.4. Update on on-going extension application according to Annex I of Commission Regulation (EC) No 1234/2008

4.5. Re-examination procedure of extension of marketing authorisation according to Annex I of Commission Regulation (EC) No 1234/2008

5. Type II variations - variation of therapeutic indication procedure according to Annex I of Commission Regulation (EC) No 1234/2008

5.1. Type II variations - variation of therapeutic indication procedure according to Commission Regulation (EC) No 1234/2008; Opinions or Requests for supplementary information

5.1.1. Bydureon - exenatide - EMEA/H/C/002020/II/0041

AstraZeneca AB

Rapporteur: Kristina Dunder, PRAC Rapporteur: Qun-Ying Yue

Scope: "Update of section 4.1 of the SmPC in order to align with more recently approved glucose-lowering agents and with "Reflection paper on the wording of indication for medicinal products for treatment of type 2 diabetes" and update of section 5.1 based on the study D5553C00003 (Duration 8 study) which evaluated concomitant add-on treatment with the combination of exenatide once weekly 2 mg and dapagliflozin 10 mg once daily in patients with type 2 diabetes mellitus who have inadequate glycaemic control on metformin. The Package Leaflet is updated accordingly. In addition, the Marketing authorisation holder (MAH) took the opportunity to make minor editorial changes in the SmPC and Package Leaflet. Furthermore, the updated RMP version 24 has been submitted."

Action: For adoption

5.1.2. [Genvoya - elvitegravir / cobicistat / emtricitabine / tenofovir alafenamide - EMEA/H/C/004042/II/0026](#)

Gilead Sciences International Ltd

Rapporteur: Robert James Hemmings, PRAC Rapporteur: Amelia Cupelli

Scope: "Extension of Indication to include paediatric patients from 6 years of age to less than 12 years of age, with body weight of at least 25kg, infected with human immunodeficiency virus-1 (HIV-1) without any known mutations associated with resistance to the integrase inhibitor class, emtricitabine or tenofovir, for Genvoya.

As a consequence, sections 4.2, 4.8, 5.1 and 5.2 of the SmPC are updated based on the analysis of the paediatric study GS-US-292-0106 (Cohort 2) "A Phase 2/3, Open-Label Study of the Pharmacokinetics, Safety, and Antiviral Activity of the Elvitegravir/Cobicistat/Emtricitabine/Tenofovir Alafenamide (E/C/F/TAF) Single Tablet Regimen (STR) in HIV-1 Infected Antiretroviral Treatment Naïve Adolescents and Virologically Suppressed Children".

The Package Leaflet and the Risk Management Plan (v. 3) are updated in accordance."

Action: For adoption

5.1.3. [Humira - adalimumab - EMEA/H/C/000481/II/0163](#)

AbbVie Ltd.

Rapporteur: Kristina Dunder, PRAC Rapporteur: Ulla Wändel Liminga

Scope: "Extension of Indication to include new indication for treatment of chronic non-infectious uveitis in paediatric patients for Humira. As a consequence, sections 4.1, 4.2, 4.8 and 5.1 of the SmPC are updated. The Package Leaflet and the RMP are updated in accordance.

In addition, the Marketing authorisation holder (MAH) took the opportunity to implement an alternative format statement for blind/partially sighted patients into the Package Leaflet as it was introduced with procedure EMEA/H/C/000481/N/0155.

Furthermore, the MAH has made some editorial changes to the Package leaflet."

Action: For adoption

5.1.4. Izba - travoprost - EMEA/H/C/002738/II/0005

Alcon Laboratories (UK) Ltd

Rapporteur: Concepcion Prieto Yerro, Co-Rapporteur: Greg Markey, PRAC Rapporteur: Almath Spooner

Scope: "Extension of Indication to include treatment of paediatric patients aged 2 months to < 18 years with ocular hypertension or paediatric glaucoma in order to decrease of elevated intraocular pressure. As a consequence, sections 4.1, 4.2, 4.4, 4.8, 5.1 and 5.2 of the SmPC are updated. The Package leaflet has been updated accordingly. In addition, the marketing authorisation holder took the opportunity to introduce minor corrections in the SmPC and to update the list of local representatives in the PL. The RMP has updated to version 9.0"

Action: For adoption

Request for Supplementary Information adopted on 15.12.2016.

5.1.5. Keytruda - pembrolizumab - EMEA/H/C/003820/II/0014

Merck Sharp & Dohme Limited

Rapporteur: Daniela Melchiorri, Co-Rapporteur: Jan Mueller-Berghaus, PRAC Rapporteur: Sabine Straus

Scope: "Extension of Indication to include the treatment of classical Hodgkin Lymphoma (cHL) in adults who have refractory disease, or who have relapsed after greater than 3 prior lines of therapy, based on the results from study KEYNOTE-087, an open-label Phase II trial of pembrolizumab in subjects with relapsed or refractory cHL and study KEYNOTE-013, a Phase Ib multi-cohort trial of pembrolizumab in subjects with hematologic malignancies. As a consequence, sections 4.1, 4.2, 4.8, 5.1 and 5.2 of the SmPC are updated and the Package Leaflet is updated accordingly. An updated RMP version 5.0 was provided as part of the application."

Action: For adoption

Request for Supplementary Information adopted on 26.01.2017.

5.1.6. Nplate - romiplostim - Orphan - EMEA/H/C/000942/II/0060/G

Amgen Europe B.V.

Rapporteur: Aranzazu Sancho-Lopez, Co-Rapporteur: Paula Boudewina van Hennik, PRAC Rapporteur: Eva A. Segovia

Scope: "C.I.6.a - Extension of Indication to include paediatric population for Nplate: to register Nplate for the use in the paediatric chronic immune (idiopathic) thrombocytopenic purpura (ITP) patients: 1 year of age and older.

As a consequence Product information has been updated accordingly.

The RMP version 18 has also been submitted.

Furthermore, the PI is brought in line with the latest QRD template version 10.

B.II.e.5.c – To add a low-dose romiplostim 125 microgram vial presentation for powder for solution for injection (4 vials pack).

B.II.e.5.a.1 – To add a 1 vial pack size of a low-dose romiplostim 125 microgram presentation."

Action: For adoption

5.1.7. Opdivo - nivolumab - EMEA/H/C/003985/II/0017

Bristol-Myers Squibb Pharma EEIG

Rapporteur: Aranzazu Sancho-Lopez, Co-Rapporteur: Paula Boudewina van Hennik, PRAC
Rapporteur: Brigitte Keller-Stanislawska

Scope: "Extension of Indication to include treatment of recurrent or metastatic squamous cell cancer of the head and neck (SCCHN) after platinum-based therapy in adults for OPDIVO.

As a consequence, sections 4.1, 4.2, 4.4, 4.8, 5.1, of the SmPC are updated in order to add the proposed new indication, add a warning that patients with a baseline performance score ≥ 2 , untreated brain metastasis, active autoimmune disease, or medical conditions requiring systemic immunosuppression were excluded from the SCCHN clinical trial and update the undesirable effect and safety information. Labelling is updated in accordance. Moreover, the updated RMP version 6.0 has been submitted"

Action: For adoption

Request for Supplementary Information adopted on 26.01.2017, 13.10.2016.

5.1.8. Opdivo - nivolumab - EMEA/H/C/003985/II/0019

Bristol-Myers Squibb Pharma EEIG

Rapporteur: Aranzazu Sancho-Lopez, Co-Rapporteur: Paula Boudewina van Hennik, PRAC
Rapporteur: Brigitte Keller-Stanislawska

Scope: "Extension of Indication to include the treatment of locally advanced unresectable or metastatic urothelial carcinoma in adults after failure of prior platinum-containing therapy for OPDIVO.

As a consequence, sections 4.1, 4.4, 4.8, 5.1 and 5.2 of the SmPC are updated in order to add the proposed indication, add a warning about the patient populations excluded from the clinical trial, and update the safety information. The Package Leaflet is updated in accordance.

Moreover, the updated RMP version 7.0 has been submitted"

Action: For adoption

Request for Supplementary Information adopted on 15.12.2016.

5.1.9. Opdivo - nivolumab - EMEA/H/C/003985/II/0029

Bristol-Myers Squibb Pharma EEIG

Rapporteur: Aranzazu Sancho-Lopez, Co-Rapporteur: Paula Boudewina van Hennik, PRAC
Rapporteur: Brigitte Keller-Stanislawska

Scope: "Extension of Indication to include the treatment of hepatocellular carcinoma after prior sorafenib therapy in adults for OPDIVO.

As a consequence, sections 4.1, 4.2, 4.4, 4.8 and 5.1 of the SmPC are updated. The Package Leaflet is updated in accordance.
Moreover, the updated RMP version 8.0 has been submitted."

Action: For adoption

5.1.10. Opdivo - nivolumab - EMEA/H/C/003985/II/0030

Bristol-Myers Squibb Pharma EEIG

Rapporteur: Aranzazu Sancho-Lopez, Co-Rapporteur: Paula Boudewina van Hennik, PRAC
Rapporteur: Brigitte Keller-Stanislawska

Scope: "Extension of indication to include treatment of adults with mismatch repair deficient (dMMR) or microsatellite instability high (MSI-H) metastatic colorectal cancer after prior fluoropyrimidine based therapy for OPDIVO.

As a consequence, sections 4.1, 4.4, 4.8 and 5.1 of the SmPC are updated in order to add the new indication and update the safety information. The Package Leaflet is updated in accordance.

RMP version 9.0 is submitted with this application"

Action: For adoption

5.1.11. RoActemra - tocilizumab - EMEA/H/C/000955/II/0066

Roche Registration Limited

Rapporteur: Jan Mueller-Berghaus, Co-Rapporteur: Agnes Gyurasics, PRAC Rapporteur:
Brigitte Keller-Stanislawska

Scope: "Extension of indication to include an indication in adult patients for the treatment of giant cell arteritis for the subcutaneous formulation of RoActemra. As a consequence, sections 4.1, 4.2, 4.4, 4.5, 4.8, 5.1 and 5.2 of the SmPC are updated to reflect information relevant to this indication. The Package Leaflet is updated in accordance."

Action: For adoption

5.1.12. Soliris - eculizumab - Orphan - EMEA/H/C/000791/II/0090

Alexion Europe SAS

Rapporteur: Aranzazu Sancho-Lopez, PRAC Rapporteur: Eva A. Segovia

Scope: "Extension of Indication of Soliris to include the 'treatment of Refractory generalized Myasthenia Gravis (gMG) patients who are antiacetylcholine receptor (AChR) antibody-positive'.

As a consequence, sections 4.1, 4.2, 4.4, 4.8, 5.1 and 5.2 of the SmPC are updated to include information on the new indication and to include the new methodology to calculate the Adverse Drug Reaction frequencies (section 4.8).

The RMP is updated accordingly (version 14.0)."

Action: For adoption

5.1.13. Zebinix - eslicarbazepine acetate - EMEA/H/C/000988/II/0053

Bial - Portela & C^a, S.A.

Rapporteur: Martina Weise, Co-Rapporteur: Ondřej Slanař, PRAC Rapporteur: Martin Huber

Scope: "Extension of indication to include the use of Zebinix as monotherapy in adults, in addition to the previously authorised indication as adjunctive therapy. As a consequence, Sections 4.1, 4.2, 4.4, 4.8, 5.1 and 5.2 of the SmPC are updated. The Package Leaflet is updated in accordance. This submission includes an updated RMP (version 15.0). In addition, the MAH is claiming an additional 1-year period of market protection under Article 14(11) of Regulation (EC) No 726/2004.", Request for 1 year of market protection for a new indication (Article 14(11) of Regulation (EC) 726/2004)

Action: For adoption

Request for Supplementary Information adopted on 15.12.2016, 21.07.2016.

5.1.14. Zykadia - ceritinib - EMEA/H/C/003819/II/0012

Novartis Europharm Ltd

Rapporteur: Aranzazu Sancho-Lopez, Co-Rapporteur: Bjorg Bolstad, PRAC Rapporteur: Ulla Wändel Liminga

Scope: "Extension of Indication to include new indication/population for Zykadia as first-line treatment of adult patients with anaplastic lymphoma kinase (ALK)-positive advanced non-small cell lung cancer (NSCLC).

As a consequence, sections 4.1, 4.2, 4.4, 4.5, 4.8, 5.1, 5.2 of the SmPC are updated to update the information based primarily on the supporting study, CLDK378A2301 (ASCEND-4). The Package Leaflet is updated in accordance.

An updated Risk Management Plan (Version 6) is also included in the application."

Action: For adoption

- 5.2. **Update on on-going Type II variation; variation of therapeutic indication procedure according to Commission Regulation (EC) No 1234/2008**
- 5.3. **Re-examination of Type II variation; variation of therapeutic indication procedure according to Commission Regulation (EC) No 1234/2008**

6. Ancillary medicinal substances in medical devices

- 6.1. **Ancillary medicinal substances in medical devices; Opinions/ Day 180 list of outstanding issues / Day 120 list of questions**

- 6.2. **Update of Ancillary medicinal substances in medical devices**

7. Procedure under Article 83(1) of Regulation (EC) 726/2004 (Compassionate Use)

- 7.1. **Procedure under Article 83(1) of Regulation (EC) 726/2004 (Compassionate Use)**

8. Pre-submission issues

8.1. Pre-submission issue

8.1.1. – Iterminovir – Orphan - H0004536

Merck Sharp & Dohme Limited, Indicated for prophylaxis of cytomegalovirus (CMV) infection or disease in adult CMV-seropositive recipients [R+] of an allogeneic hematopoietic stem cell transplant (HSCT)

Scope: Briefing note and Rapporteurs' recommendation on the request for accelerated assessment.

Action: For adoption

8.1.2. – erenumab - H0004447

Prophylaxis of migraine in adults

Scope: Briefing note and Rapporteurs' recommendation on the request for accelerated assessment.

Action: For adoption

8.2. Priority Medicines (PRIME)

Disclosure of information related to priority medicines cannot be released at present time as these contain commercially confidential information

8.2.1. List of applications received

Action: For information

8.2.2. Recommendation for PRIME eligibility

Action: For adoption

9. Post-authorisation issues

9.1. Post-authorisation issues

9.1.1. Mozobil - plerixafor – Orphan - EMEA/H/C/001030/II/0030/G

Genzyme Europe BV

Rapporteur: Paula Boudewina van Hennik, PRAC Rapporteur: Sabine Straus,

Scope: "Submission of the final report from study ARD12858 (MOZ23510) "A pilot, exploratory, randomized, phase 2 safety study evaluating tumor cell (plasma cell) mobilization and apheresis product contamination in plerixafor plus non-pegylated G-CSF mobilized patients and in non pegylated G-CSF alone mobilized patients" listed as a category 3 study in the RMP.

Submission of the final report from study OBS13611 (MOZ18009), a multicenter, noninterventional registry designed to evaluate the long-term outcomes for patients who received plerixafor for stem cell mobilization and completed hematopoietic stem cell transplantation (HSCT) compared with patients who received other mobilization methods and completed HSCT, listed as a category 3 study in the RMP.

Submission of the final report from study OBS13612 (MOZ19310), monitoring the plerixafor off-label transplant use, in patients and donors in EBMT centers performing autologous transplants and/or allogeneic transplants, listed as a category 3 study in the RMP."

Action: For discussion

9.1.2. Fampyra - fampridine - EMEA/H/C/002097/II/0036/G

MAH: Biogen Idec Ltd

Rapporteur: Johann Lodewijk Hillege, PRAC Rapporteur: Sabine Straus

Scope: "This is a grouped variation proposing updates:

- to the SmPC sections 4.2, 5.1, Annex II and Package Leaflet based on the clinical study Enhance,
- to the SmPC section 4.6 based on the data from pregnancy registry.
- Further changes to the PI, section 4.2 and 5.2 of the SmPC have been introduced based on the Core Data Sheet (CDS) and PRAC review of the Fampyra PSUR 03.

The RMP (version 11) has been updated accordingly. In addition, the Marketing authorisation holder (MAH) took the opportunity to bring the PI in line with the latest QRD template version 10.0.

With this application the MAH requests to switch the marketing authorisation from conditional to standard."

Action: For discussion

Request for Supplementary Information adopted on 26.01.2017.

9.1.3. Epclusa - sofosbuvir/velpatasvir - EMEA/H/C/004210/WS1075/0006

Harvoni - ledipasvir/sofosbuvir - EMEA/H/C/003850/WS1075/0043

Sovaldi – sofosbuvir - EMEA/H/C/002798/WS1075/0037 - WS1075

MAH: Gilead Sciences International Ltd

Lead Rapporteur: Filip Josephson, Lead PRAC Rapporteur: Ana Sofia Diniz Martins

Scope: "Submission of the final non-clinical study report PC-334-2035 assessing the potential for a pharmacokinetic interaction via transporter or enzyme based inhibition when sofosbuvir and other Direct Acting Antivirals (DAAs) are used concomitantly with amiodarone. The RMPs (Epclusa – RMP version 1.0, Harvoni – RMP version 2.0, Sovaldi – RMP version 5.0) have been updated accordingly."

Action: For adoption

Request for Supplementary Information adopted on 26.01.2017

9.1.4. Hemoprostol - misoprostol - Article 58 - EMEA/H/W/002652

Linepharma International Limited; treatment and prevention of Post Partum Haemorrhage

Rapporteur: Paula Boudewina van Hennik, Co-Rapporteur: Nithyanandan Nagercoil, PRAC

Rapporteur: Alexandre Moreau

Scope: Withdrawal of marketing authorisation

Action: For information

Article 58 of Regulation (EC) No 726/2004

The medicinal product Hemoprostol (200 µg, tablet) was exclusively intended for markets outside the European Union.

10. Referral procedures

- 10.1. **Procedure for Centrally Authorised products under Article 20 Council Regulation (EC) No 726/2004**
- 10.2. **Requests for CHMP Opinion under Article 5(3) of Regulation (EC) No 726/2004**
- 10.3. **Procedure under Articles 5(2) and 10 of the Regulation (EC) No 726/2004**
- 10.4. **Disagreement between Member States on application for medicinal product (potential serious risk to public health) –under Article 29(4) of Directive 2001/83/EC**
- 10.5. **Harmonisation - Referral procedure under Article 30 of Directive 2001/83/EC**
- 10.6. **Community Interests - Referral under Article 31 of Directive 2001/83/EC**

10.6.1. Micro Therapeutics Research Labs, India - EMEA/H/A-31/1450

Rapporteur: Johann Lodewijk Hillege, Co-Rapporteur: Milena Stain

Scope: Opinion

Action: For adoption

Reliability of the data of bioequivalence studies

10.6.2. Symbioflor 2, Escherichia Coli bacteria (cells and autolysate) - EMEA/H/A-31/1441

Symbiopharm GmbH,

Rapporteur: Harald Enzmann, Co-rapporteur: Milena Stain;

Scope: Opinion

Article 31 triggered by the BfArM in Germany in March 2016 requesting the review of the benefit-risk balance for Symbioflor 2 and associated names following concerns that the effectiveness of the medicine(s) has not been adequately demonstrated.

Action: For adoption

10.7. Re-examination Procedure under Article 32(4) of Directive 2001/83/EC

10.8. Procedure under Article 107(2) of Directive 2001/83/EC

10.9. Disagreement between Member States on Type II variation– Arbitration procedure initiated by MAH under Article 6(13) (EC) No 1084/2003

10.10. Procedure under Article 29 Regulation (EC) 1901/2006

10.11. Referral under Article 13 Disagreement between Member States on Type II variation– Arbitration procedure initiated by Member State under Article 13 (EC) No 1234/2008)

10.11.1. Cardioxane - Dexrazoxane - EMEA/H/A-13/1453

Clinigen Group

Rapporteur: Alexandre Moreau, Co-Rapporteur: Greg Markey

RMS: FR, CMS: CZ, DE, ES, IT, NL, PL & UK

Decentralised Procedure numbers: FR/H/283/01/II/27G

Scope: List of Questions/Opinion

Article 13 triggered by the ANSM in France in January 2017 requesting the CHMP's opinion whether the proposed lifting of the contraindication for a subset of anthracycline treated children is justified.

Action: For adoption

11. Pharmacovigilance issue

11.1. Early Notification System

March 2017 Early Notification System on envisaged CHMP/CMDh outcome accompanied by communication to the general public.

Action: For information

12. Inspections

12.1. GMP inspections

Disclosure of information related to GMP inspections will not be published as it undermines the purpose of such inspections

12.2. GCP inspections

Disclosure of information related to GCP inspections will not be published as it undermines the purpose of such inspections

12.3. Pharmacovigilance inspections

Disclosure of information related to Pharmacovigilance inspections will not be published as it undermines the purpose of such inspections

12.4. GLP inspections

Disclosure of information related to GLP inspections will not be published as it undermines the purpose of such inspections

13. Innovation Task Force

13.1. Minutes of Innovation Task Force

Action: For information

13.2. Innovation Task Force briefing meetings

Disclosure of information related to briefing meetings taking place with applicants cannot be released at present time as deemed to contain commercially confidential information

13.2.1. Briefing meeting

Meeting date: 22 March 2017

Action: For information

13.3. Requests for CHMP Opinion under Article 57(1)J and (1)P of Regulation (EC) No 726/2004

13.4. Nanomedicines activities

14. Organisational, regulatory and methodological matters

14.1. Mandate and organisation of the CHMP

14.1.1. Review of experience with the revised RMP assessment process for new marketing authorisations

Action: For information

14.1.2. ATMP guideline on safety and efficacy follow-up and risk management (EMA/CHMP/65416/2016)

Scope: Comments received by CAT and PRAC before the consultation of GCG.

Action: For information

14.2. Coordination with EMA Scientific Committees

14.2.1. Pharmacovigilance Risk Assessment Committee (PRAC)

Summary of recommendations and advice of PRAC meeting held on 6-9 March 2017

Action: For information

List of Union Reference Dates and frequency of submission of Periodic Safety Update Reports (EURD list) for March 2017

Action: For adoption

14.2.2. Committee for Advanced Therapies (CAT)

CAT draft minutes of meeting held on 15-17 March 2017

Action: For information

14.2.3. Committee for Herbal Medicinal Products (HMPC)

Report from the HMPC meeting

Action: For information

14.2.4. Paediatric Committee (PDCO)

PIPs reaching D30 at March 2017 PDCO

Action: For information

Report from the PDCO meeting held on 21-24 March 2017

Action: For information

14.2.5. Committee for Orphan Medicinal Products (COMP)

Report from the COMP meeting held on 14-15 March 2017

Action: For information

14.2.6. Coordination Group for Mutual Recognition and Decentralised Procedures – Human (CMDh)

Report from the Coordination Group for Mutual Recognition and Decentralised Procedures – Human (CMDh) on the meeting held on 20-22 March 2017

Action: For information

14.3. Coordination with EMA Working Parties/Working Groups/Drafting Groups

14.3.1. Scientific Advice Working Party (SAWP)

Report from the SAWP meeting held on 6-9 March 2017. Table of conclusions

Action: For information

Scientific advice letters: Disclosure of information related to scientific advice letters cannot be released at present time as these contain commercially confidential information.

14.3.2. Scientific Advice Working Party (SAWP)

Appointment of a replacement SAWP member and alternate following resignation of Thomas Lang. The required area of expertise is statistics.

Action: For adoption

14.3.3. Blood Products Working Party (BPWP)

Vice-Chair: Karri Penttilä,

Scope: Election of a new Chairperson of the Blood Products Working Party (BPWP).

Action: For adoption

Nomination of new observer Marie Louise Schougaard Christiansen (DK) to the BPWP

Action: For adoption

14.3.4. Ad-hoc Influenza Working Group

Scope: EU Strain selection for the Influenza Vaccines for the Season 2017/2018

Action: For adoption

Scope: EU Recommendation for the Seasonal Influenza Vaccine Composition for the Season 2017/2018

Action: For adoption

Report from the Ad Hoc Influenza working group to the BWP

14.3.5. CHMP ad-hoc drafting group

Scope: Revision of the Guideline on clinical development of fixed combination medicinal products

Action: For adoption

14.3.6. Biostatistics Working Party (BSWP)

Chair: Anja Schiel/Thomas Lang

Scope: Reflection paper on statistical methodology for the comparative assessment of quality attributes in drug development

Action: For adoption for 6-months public consultation

14.3.7. Pharmacokinetics Working Party (PKWP)

Chair: Jan Welink/Alfredo Garcia-Arieta

Scope: Guideline on equivalence studies for the demonstration of therapeutic equivalence for products that are locally applied locally acting in the gastrointestinal tract as addendum to the guideline on the clinical requirements for locally applied, locally acting products containing known constituents (CPMP/EWP/239/95 Rev. 1)

Action: For adoption for 6 months public consultation

14.3.8. Pharmacogenomics Working Party (PGWP)

Chair: Krishna Prasad/Markus Paulmichl,

Draft agenda for the F2F meeting on 27-28 March 2017 (Doc. Ref. EMA/75533/2017)

Action: For information

14.3.9. Oncology Working Party (ONCWP)

Chair: Pierre Demolis

Nomination of new observer Doris J. Hovgaard (DK) to the ONCWP

Action: For adoption

14.3.10. Cardiovascular Working Party (CVSWP)

Chair: Pieter de Graeff/Kristina Dunder

Call for nomination of CVSWP core member following resignation of Karsten Bruins Slot

Action: For information

14.3.11. Biologics Working Party (BWP)

Chair: Sol Ruiz,

Call for nomination of BWP Vice-Chair, the term of current Vice Chair ending in April 2017.

Nominations should be sent by 6 April 2017

Action: For information

14.3.12. Central Nervous System Working Party (CNSWP)

Chair: Karl Broich,

Nomination of new observer Eskild Colding-Jørgensen (DK) to the CNSWP

Action: For adoption

14.4. Cooperation within the EU regulatory network

14.5. Cooperation with International Regulators

14.6. Contacts of the CHMP with external parties and interaction with the Interested Parties to the Committee

14.7. CHMP work plan

14.8. Planning and reporting

14.9. Others

15. Any other business

15.1. AOB topic

15.1.1. Preparedness of the system and capacity increase

Action: For discussion

15.1.2. Revision of the 'Guideline on strategies to identify and mitigate risks for first-in-human clinical trials with investigational medicinal products'

Scope: update on the revision
workshop to be held 28 March 2017 (draft programme)

Action: For information

16. Explanatory notes

The notes below give a brief explanation of the main sections and headings in the CHMP agenda and should be read in conjunction with the agenda or the minutes.

Oral explanations (section 2)

The items listed in this section are those for which marketing authorisation holders (MAHs) or applicants have been invited to the CHMP plenary meeting to address questions raised by the Committee. Oral explanations normally relate to on-going applications (section 3, 4 and 5) or referral procedures (section 10) but can relate to any other issue for which the CHMP would like to discuss with company representatives in person.

Initial applications (section 3)

This section lists applications for marketing authorisations of new medicines that are to be discussed by the Committee.

Section 3.1 is for medicinal products nearing the end of the evaluation and for which the CHMP is expected to adopt an **opinion** at this meeting on whether marketing authorisation should be granted. Once adopted, the CHMP opinion will be forwarded to the European Commission for a final legally binding decision valid throughout the EU.

The other items in the section are listed depending on the stage of the evaluation, which is shown graphically below:



The assessment of an application for a new medicine takes up to 210 'active' days. This active evaluation time is interrupted by at least one 'clock-stop' during which time the applicant prepares the answers to questions from the CHMP. The clock stop happens after day 120 and may also happen after day 180, when the CHMP has adopted a list of questions or outstanding issues to be addressed by the company. Related discussions are listed in the agenda under sections 3.2 (**Day 180 List of outstanding issues**) and 3.3 (**Day 120 list of questions**).

CHMP discussions may also occur at any other stage of the evaluation, and these are listed under section 3.4, **update on ongoing new applications for centralised procedures**.

The assessment leads to an opinion from the CHMP by day 210. Following a CHMP opinion the European Commission takes usually 67 days to issue a legally binding decision (i.e. by day 277 of the procedure). CHMP discussions on products that have received a CHMP opinion and are awaiting a decision are listed under section 3.6, **products in the decision making phase**.

Extension of marketing authorisations according to Annex I of Reg. 1234/2008 (section 4)

Extensions of marketing authorisations are applications for the change or addition of new strengths,

formulations or routes of administration to existing marketing authorisations. Extension applications follow a 210-day evaluation process, similarly to applications for new medicines (see figure above).

Type II variations - Extension of indication procedures (*section 5*)

Type II variations are applications for a change to the marketing authorisation which requires an update of the product information and which is not covered in section 4. Type II variations include applications for a new use of the medicine (extension of indication), for which the assessment takes up to 90 days. For the applications listed in this section, the CHMP may adopt an opinion or request supplementary information from the applicant.

Ancillary medicinal substances in medical devices (*section 6*)

Although the EMA does not regulate medical devices it can be asked by the relevant authorities (the so-called Notified Bodies) that are responsible for regulating these devices to give a scientific opinion on a medicinal substance contained in a medical device.

Re-examination procedures (new applications) under article 9(2) of regulation no 726/2004 (*section 3.5*)

This section lists applications for new marketing authorisation for which the applicant has requested a re-examination of the opinion previously issued by the CHMP.

Re-examination procedures (*section 5.3*)

This section lists applications for type II variations (including extension of indication applications) for which the applicant has requested re-examination of the opinion previously issued by the CHMP.

Withdrawal of application (*section 3.7*)

Applicants may decide to withdraw applications at any stage during the assessment and a CHMP opinion will therefore not be issued. Withdrawals are included in the agenda for information or discussion, as necessary.

Procedure under article 83(1) of regulation (EC) 726/2004 (compassionate use) (*section 7*)

Compassionate use is a way of making available to patients with an unmet medical need a promising medicine which has not yet been authorised (licensed) for their condition. Upon request, the CHMP provides recommendations to all EU Member States on how to administer, distribute and use certain medicines for compassionate use.

Pre-submission issues (*section 8*)

In some cases the CHMP may discuss a medicine before a formal application for marketing authorisation is submitted. These cases generally refer to requests for an accelerated assessment for medicines that are of major interest for public health or can be considered a therapeutic innovation. In case of an accelerated assessment the assessment timetable is reduced from 210 to 150 days.

Post-authorisation issues (*section 9*)

This section lists other issues concerning authorised medicines that are not covered elsewhere in the agenda. Issues include supply shortages, quality defects, some annual reassessments or renewals or type II variations to marketing authorisations that would require specific discussion at the plenary.

Referral procedures (*section 10*)

This section lists referrals that are ongoing or due to be started at the plenary meeting. A referral is a procedure used to resolve issues such as concerns over the safety or benefit-risk balance of a medicine or a class of medicines. In a referral, the EMA is requested to conduct a scientific assessment of a

particular medicine or class of medicines on behalf of the EU. Further information on such procedures can be found [here](#).

Pharmacovigilance issues (section 11)

This section lists issues that have been discussed at the previous meeting of the PRAC, the EMA's committee responsible for evaluating and monitoring safety issues for medicines. Feedback is provided by the PRAC. This section also refers to the early notification system, a system used to notify the European regulatory network on proposed EMA communication on safety of medicines.

Inspections Issues (section 12)

This section lists inspections that are undertaken for some medicinal products. Inspections are carried out by regulatory agencies to ensure that marketing authorisation holders comply with their obligations. Inspection can relate to good manufacturing practice (GMP), good clinical practice (GCP), good laboratory practice (GLP) or good pharmacovigilance practice (GVP).

Innovation task force (section 13)

The Innovation Task Force (ITF) is a body set up to encourage early dialogue with applicants developing innovative medicines. Minutes from the last ITF meeting as well as any related issue that requires discussion with the CHMP are listed in this section of the agenda. Further information on the ITF can be found [here](#).

Scientific advice working party (SAWP) (section 14.3.1)

This section refers to the monthly report from the CHMP's Scientific Advice Working Party (SAWP) on scientific advice given to companies during the development of medicines. Further general information on SAWP can be found [here](#).

Satellite groups / other committees (section 14.2)

This section refers to the reports from groups and committees making decisions relating to human medicines: the Coordination Group for Mutual Recognition and Decentralised Procedures – Human (CMDh), the Committee for Orphan Medicinal Products (COMP), the Committee for Herbal Medicinal Products (HMPC), Paediatric Committee (PDCO), the Committee for Advanced Therapies (CAT) and the Pharmacovigilance Risk Assessment Committee (PRAC).

Invented name issues (section 14.3)

This section lists issues related to invented names proposed by applicants for new medicines. The CHMP has established the Name Review Group (NRG) to perform reviews of the invented names. The group's main role is to consider whether the proposed names could create a public-health concern or potential safety risk. Further information can be found [here](#).

More detailed information on the above terms can be found on the EMA website: www.ema.europa.eu/

20 March 2017
EMA/CHMP/184754/2017

Annex to March 2017 CHMP Agenda

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A. PRE SUBMISSION ISSUES

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Report on Eligibility to Centralised Procedure for
March 2017: **For adoption**

A.2. APPOINTMENT OF RAPPORTEUR / CO-RAPPORTEUR FULL APPLICATIONS

Final Outcome of Rapporteurship allocation for
March 2017: **For adoption**

A.3. PRE-SUBMISSION ISSUES FOR INFORMATION

Disclosure of information related to pre-submission of initial applications cannot be released at present time as these contain commercially confidential information.

B. POST-AUTHORISATION PROCEDURES OUTCOMES

B.1. Annual re-assessment outcomes

B.1.1. Annual reassessment for products authorised under exceptional circumstances

Ceplene - histamine dihydrochloride -
EMEA/H/C/000796/S/0030, Orphan
MAH: Meda AB, Rapporteur: David Lyons, PRAC
Rapporteur: Almath Spooner

Kolbam - cholic acid -
EMEA/H/C/002081/S/0020, Orphan
MAH: Retrophin Europe Ltd, Rapporteur: Robert James Hemmings, PRAC Rapporteur: Patrick Batty

Lojuxta - lomitapide -
EMEA/H/C/002578/S/0023
MAH: Aegerion Pharmaceuticals Limited,
Rapporteur: Johann Lodewijk Hillege, PRAC
Rapporteur: Menno van der Elst
Request for Supplementary Information adopted
on 23.02.2017, 10.11.2016.

Raxone - idebenone -
EMEA/H/C/003834/S/0005, Orphan
MAH: Santhera Pharmaceuticals (Deutschland)

GmbH, Rapporteur: John Joseph Borg, PRAC
Rapporteur: Carmela Macchiarulo
Request for Supplementary Information adopted
on 26.01.2017.

Vedrop - tocofersolan -
EMEA/H/C/000920/S/0019
MAH: Orphan Europe S.A.R.L., Rapporteur:
Greg Markey, PRAC Rapporteur: Julie Williams
Request for Supplementary Information adopted
on 26.01.2017.

Vyndaqel - tafamidis -
EMEA/H/C/002294/S/0036, Orphan
MAH: Pfizer Limited, Rapporteur: Joseph
Emmerich, PRAC Rapporteur: Caroline Laborde

Xagrid - anagrelide -
EMEA/H/C/000480/S/0077
MAH: Shire Pharmaceutical Contracts Ltd.,
Rapporteur: Alexandre Moreau, Co-Rapporteur:
Koenraad Norga, PRAC Rapporteur: Caroline
Laborde

B.2. RENEWALS OF MARKETING AUTHORISATIONS OUTCOMES

B.2.1. Renewals of Marketing Authorisations requiring 2nd Renewal

NovoThirteen - catridecacog -
EMEA/H/C/002284/R/0020
MAH: Novo Nordisk A/S, Rapporteur: Joseph
Emmerich, Co-Rapporteur: Jan Mueller-
Berghaus, PRAC Rapporteur: Caroline Laborde

Rasilez - aliskiren -
EMEA/H/C/000780/R/0112
MAH: Novartis Europharm Ltd, Rapporteur:
Daniela Melchiorri, Co-Rapporteur: Melinda
Sobor, PRAC Rapporteur: Carmela Macchiarulo
Request for Supplementary Information adopted
on 23.02.2017.

B.2.2. Renewals of Marketing Authorisations for unlimited validity

Cuprymina - copper (64Cu) chloride -
EMEA/H/C/002136/R/0014
MAH: Sparkle S.r.l., Rapporteur: Greg Markey,
Co-Rapporteur: Daniela Melchiorri, PRAC
Rapporteur: Patrick Batty

Dacogen - decitabine -
EMEA/H/C/002221/R/0030, Orphan

MAH: Janssen-Cilag International NV,
Rapporteur: Alexandre Moreau, Co-Rapporteur:
Greg Markey, PRAC Rapporteur: Caroline
Laborde

Enurev Breezhaler - glycopyrronium bromide - EMEA/H/C/002691/R/0020
MAH: Novartis Europharm Ltd, Duplicate,
Duplicate of Seebri Breezhaler, Rapporteur:
Hanne Lomholt Larsen, Co-Rapporteur: David
Lyons, PRAC Rapporteur: Torbjorn Callreus

Inlyta - axitinib -
EMEA/H/C/002406/R/0021
MAH: Pfizer Limited, Rapporteur: Bjorg Bolstad,
Co-Rapporteur: Sinan B. Sarac, PRAC
Rapporteur: Helga Haugom Olsen

Revestive - teduglutide -
EMEA/H/C/002345/R/0038, Orphan
MAH: Shire Pharmaceuticals Ireland Ltd,
Rapporteur: Sinan B. Sarac, Co-Rapporteur:
Harald Enzmann, PRAC Rapporteur: Torbjorn
Callreus

Seebri Breezhaler - glycopyrronium -
EMEA/H/C/002430/R/0020
MAH: Novartis Europharm Ltd, Rapporteur:
Hanne Lomholt Larsen, Co-Rapporteur: David
Lyons, PRAC Rapporteur: Torbjorn Callreus

Torisel - temsirolimus -
EMEA/H/C/000799/R/0065, Orphan
MAH: Pfizer Limited, Rapporteur: Harald
Enzmann, Co-Rapporteur: Paula Boudeolina van
Hennik, PRAC Rapporteur: Martin Huber

Tovanor Breezhaler - glycopyrronium -
EMEA/H/C/002690/R/0022
MAH: Novartis Europharm Ltd, Duplicate,
Duplicate of Seebri Breezhaler, Rapporteur:
Hanne Lomholt Larsen, Co-Rapporteur: David
Lyons, PRAC Rapporteur: Torbjorn Callreus

Zoledronic acid Mylan - zoledronic acid -
EMEA/H/C/002482/R/0013
MAH: MYLAN S.A.S, Generic, Generic of
Zometa, Rapporteur: Milena Stain, PRAC
Rapporteur: Doris Stenver

Zoledronic acid Teva - zoledronic acid -
EMEA/H/C/002439/R/0018
MAH: Teva B.V., Generic, Generic of Zometa,
Rapporteur: Filip Josephson, PRAC Rapporteur:

Ulla Wändel Liminga

Zoledronic acid Teva Pharma - zoledronic acid - EMEA/H/C/002437/R/0014

MAH: Teva B.V., Generic, Generic of Aclasta,
Rapporteur: Filip Josephson, PRAC Rapporteur:
Doris Stenver

B.2.3. Renewals of Conditional Marketing Authorisations

**Fampyra - fampridine -
EMEA/H/C/002097/R/0037**

MAH: Biogen Idec Ltd, Rapporteur: Johann
Lodewijk Hillege, Co-Rapporteur: Martina Weise,
PRAC Rapporteur: Sabine Straus

B.3. POST-AUTHORISATION PHARMACOVIGILANCE OUTCOMES

Signal detection

PRAC recommendations on signals adopted at
the PRAC meeting held on 6-9 March 2017
PRAC:

PSUR procedures for which PRAC adopted a
recommendation for variation of the terms of
the MA at its 6-9 March 2017 meeting:

EMEA/H/C/PSUSA/00000940/201608

(deferiprone)

CAPS:

Ferriprox (EMEA/H/C/000236) (deferiprone),
MAH: Apotex Europe BV, Rapporteur: Alexandre
Moreau, PRAC Rapporteur: Caroline Laborde,
"01/09/2015 - 31/08/2016"

EMEA/H/C/PSUSA/00000962/201607

(desloratadine)

CAPS:

Aerius (EMEA/H/C/000313) (desloratadine),
MAH: Merck Sharp & Dohme Limited,
Rapporteur: Koenraad Norga

Azomyr (EMEA/H/C/000310) (desloratadine),
MAH: Merck Sharp & Dohme Limited,
Rapporteur: Koenraad Norga

Dasselta (EMEA/H/C/002310) (desloratadine),
MAH: KRKA, d.d., Novo mesto, Rapporteur:
Melinda Sobor

Desloratadine Actavis (EMEA/H/C/002435)
(desloratadine), MAH: Actavis Group PTC ehf,
Rapporteur: Melinda Sobor

Desloratadine ratiopharm
(EMEA/H/C/002404) (desloratadine), MAH:

ratiopharm GmbH, Rapporteur: Koenraad Norga

Desloratadine Teva (EMEA/H/C/002419)

(desloratadine), MAH: Teva B.V., Rapporteur:

Melinda Sobor

Neoclarityn (EMEA/H/C/000314)

(desloratadine), MAH: Merck Sharp & Dohme

Limited, Rapporteur: Koenraad Norga

NAPS:

Alerdes, 0,5 mg/ml, roztwór doustny 20454

PL - SYMPHAR SP. Z O.O.

Alerdes, 5 mg, tabletki powlekane 20067 PL

- SYMPHAR SP. Z O.O.

ALERDIN 0,5 mg/ml oralna otopina UP/I-

530-09/11-01/368 HR - BELUPO D.D.

ALERDIN 2,5 mg raspadljive tablete za usta

UP/I-530-09/11-01/369 HR - BELUPO D.D.

ALERDIN 5 mg raspadljive tablete za usta

UP/I-530-09/11-01/370 HR - BELUPO D.D.

Aleric Deslo Active, 2,5 mg, tabletki

ulegające rozpadowi w jamie ustnej 20562

PL - US PHARMACIA SP. Z O.O.

Aleric Deslo Active, 5 mg, tabletki ulegające

rozpadowi w jamie ustnej 20564 PL - US

PHARMACIA SP. Z O.O.

Aleric Deslo, 0,5 mg/ml, roztwór doustny

20561 PL - US PHARMACIA SP. Z O.O.

Aleric Deslo, 5 mg, tabletki powlekane

20563 PL - US PHARMACIA SP. Z O.O.

Alvotadin 5 mg filmtabлетта OGYI-T-

22014/25 HU - ALVOGEN IPCO S.AR.L

Clarderin, 0,5 mg/ml, roztwór doustny

21102 PL - PHARMASWISS ČESKÁ REPUBLIKA

S.R.O.

Clarderin, 5 mg, tabletki ulegające

rozpadowi w jamie ustnej 21037 PL -

PHARMASWISS ČESKÁ REPUBLIKA S.R.O.

Clarus 0,5 mg/ml solução oral 5634019 PT

- LABORATÓRIOS BASI – INDÚSTRIA

FARMACÊUTICA, S.A.

Dareq 0,5 mg/ml Oral solution 21528 CY -

DELOBIS PHARMACEUTICALS LTD

Dareq 5 mg Film-coated tablets 21529 CY -

DELOBIS PHARMACEUTICALS LTD

Dehistar, 0,5 mg/ml, roztwór doustny

19985 PL - FARMACEUTYCZNA SPOLDZIELNIA

PRACY GALENA

Dehistar, 5 mg, tabletki powlekane 19992

PL - FARMACEUTYCZNA SPOLDZIELNIA PRACY

GALENA

DELESIT 5 mg filmom obalené tablety

24/0417/12-S SK - CIPLA (UK) LIMITED

DELESIT 5 mg potahované tablety
24/682/12-C CZ - CIPLA (UK) LIMITED
Delortan Allergy, 5 mg, tabletki powlekane
22669 PL - WARSZAWSKIE ZAKLADY
FARMACEUTYCZNE POLFA S.A.
Delortan, 0,5 mg/ml, roztwór doustny
20061 PL - WARSZAWSKIE ZAKLADY
FARMACEUTYCZNE POLFA S.A.
Delortan, 5 mg, tabletki powlekane 20059
PL - WARSZAWSKIE ZAKLADY
FARMACEUTYCZNE POLFA S.A.
Deslix, 0,5 mg/ml, roztwór doustny 20415
PL - MEDANA PHARMA SPOLKA AKCYJNA
Deslix, 5 mg, tabletki powlekane 20414 PL -
MEDANA PHARMA SPOLKA AKCYJNA
DESLODYNA pro, 5 mg, tabletki ulegajace
rozpadowi w jamie ustnej 19963 PL -
HASCO-LEK
Deslodyna pro, smeltetabletter 5 mg 48300
DK - HASCO-LEK
Deslodyna, 0,5 mg/ml, roztwór doustny
19960 PL - HASCO-LEK
Deslodyna, 2,5 mg, tabletki ulegajace
rozpadowi w jamie ustnej 19962 PL -
HASCO-LEK
Deslodyna, 5 mg, tabletki powlekane 19961
PL - HASCO-LEK
Deslodyna, filmovertrukne tabletter 48298
DK - HASCO-LEK
Deslodyna, oral opløsning 48314 DK -
HASCO-LEK
Deslodyna, smeltetabletter 2,5 mg 48299
DK - HASCO-LEK
Deslomed 5 mg filmom obalené tablety
24/0748/11-S SK - CANDE S.R.O.
Deslor 0.5 mg/ml Oral Solution
PA0711/202/002 IE - ROWEX LTD
Deslor 48335 DK - ROWEX LTD
Deslor 48336 DK - ROWEX LTD
Deslor 5 mg Film-Coated Tablets
PA0711/202/001 IE - ROWEX LTD
Deslora-Denk 5 mg Filmtabletten
90185.00.00 DE - DENK PHARMA GMBH & CO.
KG
Desloraderm 0,5 mg/ml Lösung zum
Einnehmen 1-31334 AT - DERMAPHARM
GMBH
Desloraderm 0,5 mg/ml Lösung zum
Einnehmen 84588.00.00 DE - DERMAPHARM
AG
Desloraderm 5 mg Filmtabletten 1-31333

AT - DERMAPHARM GMBH
Desloraderm 5 mg Filmtab
84587.00.00 DE - DERMAPHARM AG
Desloradia 5 mg Filmtab 2192095 DE
- M.R. PHARMA, DE
Desloratadin +pharma 5 mg Filmtab
1-31178 AT - +PHARMA ARZNEIMITTEL GMBH
Desloratadin +pharma 5 mg potahované
tablety 24/331/12-C CZ - +PHARMA
ARZNEIMITTEL GMBH
Desloratadin - 1 A Pharma 5 mg
Filmtab 84359.00.00 DE - 1 A PHARMA
GMBH
DESLORATADIN ACTAVIS 0,5 MG/ML
BELSŐLEGES OLDAT OGYI-T-22197/04 HU -
ACTAVIS GROUP PTC EHF.
Desloratadin AL 5 mg Filmtab
84536.00.00 DE - ALIUD PHARMA GMBH
Desloratadin AL, filmovertrukne tabletter
48325 DK - ALIUD PHARMA GMBH
Desloratadin Apofri 5 mg filmdragerade
tabletter 52183 SE - APOFRI AB
Desloratadin Apotex 0,5 mg/ml perorální
roztok 24/406/12 C CZ - APOTEX EUROPE
B.V.
Desloratadin Apotex 5 mg potahované
tablety 24/209/13-C CZ - APOTEX EUROPE
BV
Desloratadin Aristo 0,5 mg/ml Lösung zum
Einnehmen 1-31628 AT - ARISTO PHARMA
GMBH (ART 57)
Desloratadin Aristo 0,5 mg/ml Lösung zum
Einnehmen 84989.00.00 DE - ARISTO
PHARMA GMBH (ART 57)
Desloratadin Aristo® 5 mg Filmtab 1-
31618 AT - ARISTO PHARMA GMBH (ART 57)
Desloratadin Aristo® 5 mg Filmtab
84988.00.00 DE - ARISTO PHARMA GMBH
(ART 57)
Desloratadin axcount 0,5 mg/ml Lösung
zum Einnehmen 94143.00.00 DE - AXCOUNT
GENERIKA GMBH
Desloratadin axcount 2,5 mg
Schmelztabletten 93355.00.00 DE -
AXCOUNT GENERIKA GMBH
Desloratadin axcount 5 mg Filmtab
93357.00.00 DE - AXCOUNT GENERIKA GMBH
Desloratadin axcount 5 mg
Schmelztabletten 93356.00.00 DE -
AXCOUNT GENERIKA GMBH
Desloratadin Cipla 5 mg filmom obložene

tablete HR-H-302001390 HR - CIPLA EUROPE

NV

Desloratadin Cipla 5 mg Filmtabletten

94826.00.00 DE - CIPLA EUROPE NV

Desloratadin Cipla, filmovertrukne tabletter

55618 DK - CIPLA EUROPE NV

DESLORATADIN DR.MAX 5 MG

POTAHOVANÉ TABLETY 24/489/12-C CZ -

DR. MAX

Desloratadin Genericon 5 mg Filmtabletten

1-31147 AT - GENERICON PHARMA

GESELLSCHAFT M.B.H.

Desloratadin Glenmark 5 mg Tabletten

84037.00.00 DE - GLENMARK

PHARMACEUTICALS EUROPE LIMITED

Desloratadin Glenmark 5mg tabletter

48522 DK - GLENMARK PHARMACEUTICALS

EUROPE LIMITED

Desloratadin HCS® 5 mg Filmtabletten

90865.00.00 DE - TAD PHARMA GMBH

Desloratadin HEXAL 0,5 mg/ml Lösung zum

Einnehmen 84366.00.00 DE - HEXAL AG

Desloratadin HEXAL 5 mg Filmtabletten

84365.00.00 DE - HEXAL AG

Desloratadin IBERMEDGEN 0.5 mg/ml

peroralen raztvor II-21146 BG -

IBERMEDGEN, S.A.

Desloratadin IBERMEDGEN 5 mg filmirani

tabletki II-21147 BG - IBERMEDGEN, S.A.

Desloratadin Krka 5 mg potahované tablety

24/213/14-C CZ - KRKA, D.D., NOVO MESTO

Desloratadin M.R. Pharma 0,5 mg/ml

Lösung zum Einnehmen 2192096 DE - M.R.

PHARMA, DE

Desloratadin M.R. Pharma 5 mg

Filmtabletten 2192094 DE - M.R. PHARMA,

DE

Desloratadin Mylan 5 mg filmdragerade

tablett 46075 SE - MYLAN AB

Desloratadin Mylan 5 mg kalvopäällysteiset

tabletit 29741 FI - MYLAN AB

Desloratadin Mylan 5 mg potahované

tablety 24/683/12-C CZ - GENERICS [UK]

LIMITED

Desloratadin Mylan, oral opløsning 48308

DK - MYLAN AB

Desloratadin Ranbaxy, filmovertrukne

tablett 48284 DK - RANBAXY PHARMACIE

GENERIQUES

Desloratadin ratiopharm 2,5 mg

Schmelztabletten 1-31313 AT - RATIOPHARM

ARZNEIMITTEL VERTRIEBS-GMBH

Desloratadin ratiopharm 5 mg

Schmelztabletten 1-31314 AT - RATIOPHARM

ARZNEIMITTEL VERTRIEBS-GMBH

Desloratadin Sandoz 0,5 mg/ml drank

BE420034 BE - SANDOZ N.V.

Desloratadin Sandoz 5 mg - Filmtabletten

1-31302 AT - SANDOZ GMBH

Desloratadin Sandoz 5 mg comprimés

pelliculés 2013010011 LU - SANDOZ N.V.

Desloratadin Sandoz 5 mg filmomhulde

tabletten BE420016 BE - SANDOZ N.V.

Desloratadin Sandoz 5 mg filmomhulde

tabletten BE420025 BE - SANDOZ N.V.

Desloratadin Saneca 5 mg tablety

24/0318/12-S SK - SANECA

PHARMACEUTICALS

Desloratadin Saneca 5 mg tablety

24/628/12-C CZ - SANECA

PHARMACEUTICALS

Desloratadin Sipla 5 mg filmirani tabletki

20130159 BG - CIPLA (UK) LIMITED

Desloratadin Sofarma 0,5 mg/ml peroralen

raztvorp 20150207 BG - SOPHARMA AD

Desloratadin Sofarma 5 mg filmirani

tabletki 20160045 BG - SOPHARMA AD

Desloratadin Specifar 2.5mg

smeltabletter 48293 DK - SPECIFAR S.A.

Desloratadin Specifar 5mg filmovertrukne

tabletter 48292 DK - SPECIFAR S.A.

Desloratadin STADA 0,5 mg/ml Lösung zum

Einnehmen 84537.00.00 DE - STADAPHARM

GMBH

Desloratadin Stada 0,5 mg/ml oraaliliuos

29571 FI - STADA ARZNEIMITTEL AG

Desloratadin Stada 5 mg 24/335/15-C CZ -

STADA ARZNEIMITTEL AG

Desloratadin STADA 5 mg filmdragerade

tabletter 45771 SE - STADA ARZNEIMITTEL

AG

Desloratadin STADA 5 mg filmdragerade

tabletter 45775 SE - STADA ARZNEIMITTEL

AG

Desloratadin STADA 5 mg Filmtabletten 1-

31291 AT - STADA ARZNEIMITTEL GMBH

Desloratadin Stada 5 mg kalvopäällysteinen

tabletti 29570 FI - STADA ARZNEIMITTEL AG

Desloratadin Stada, filmovertrukne

tabletter 48321 DK - STADA ARZNEIMITTEL

AG

Desloratadin STADA® 5 mg Filmtabletten

84535.00.00 DE - STADAPHARM GMBH
Desloratadin Stadfa 0,5 mg/ml peroralen
raztvorp 20120244 BG - STADA
ARZNEIMITTEL AG
Desloratadin Teva 0,5 mg/ml belsőleges
oldat OGYI-T-22096/05 HU - TEVA
GYÓGYSZERGYÁR ZRT
Desloratadin Teva 0,5 mg/ml belsőleges
oldat OGYI-T-22096/06 HU - TEVA
GYÓGYSZERGYÁR ZRT
Desloratadin Teva 0,5 mg/ml belsőleges
oldat OGYI-T-22096/07 HU - TEVA
GYÓGYSZERGYÁR ZRT
Desloratadin Teva 0,5 mg/ml belsőleges
oldat OGYI-T-22096/08 HU - TEVA
GYÓGYSZERGYÁR ZRT
Desloratadin Teva 0,5 mg/ml belsőleges
oldat OGYI-T-22096/09 HU - TEVA
GYÓGYSZERGYÁR ZRT
Desloratadin Teva 2,5 mg szájban
diszpergálódó tabletta OGYI-T-22096/01
HU - TEVA GYÓGYSZERGYÁR ZRT
Desloratadin Teva 2,5 mg szájban
diszpergálódó tabletta OGYI-T-22096/02
HU - TEVA GYÓGYSZERGYÁR ZRT
Desloratadin Teva 5 mg szájban
diszpergálódó tabletta OGYI-T-22096/03
HU - TEVA GYÓGYSZERGYÁR ZRT
Desloratadin Teva 5 mg szájban
diszpergálódó tabletta OGYI-T-22096/04
HU - TEVA GYÓGYSZERGYÁR ZRT
Desloratadin Zentiva 0,5 mg/ml perorální
roztok 24/326/12-C CZ - ZENTIVA, K.S.
DESLORATADIN ZENTIVA 0,5 mg/ml
perorálny roztok 24/0010/12-S SK -
ZENTIVA, K.S.
Desloratadin Zentiva 5 mg filmirani tabletki
20120167 BG - ZENTIVA, K.S.
DESLORATADIN ZENTIVA 5 mg filmom
obalené tablety 24/0008/12-S SK -
ZENTIVA, K.S.
Desloratadin Zentiva 5 mg potahované
tablety 24/324/12-C CZ - ZENTIVA, K.S.
Desloratadin Алвоген 0,5 mg/ml peroralen
raztvorp II-18062 BG - ALVOGEN IPCO S.AR.L
Desloratadin Алвоген 5 mg filmirani
tabletki II-18059 BG - ALVOGEN IPCO S.AR.L
Desloratadin Софарма 5 mg filmirani
tabletki 20160045 BG - SOPHARMA AD
Desloratadin "Actavis PTC", oral opløsning
49155 DK - ACTAVIS GROUP PTC EHF.

**Desloratadin "Stada Arzneimittel",
filmovertukne tabletter 54011 DK - STADA
ARZNEIMITTEL AG**

**Desloratadin "Stada Arzneimittel", oral
opløsning 48324 DK - STADA ARZNEIMITTEL
AG**

**Desloratadin "Stada", oral opløsning 48323
DK - STADA ARZNEIMITTEL AG**

**Desloratadin-Hormosan 5 mg Filmtabletten
84742.00.00 DE - HORMOSAN PHARMA GMBH**

**Desloratadina Actavis 2.5 mg
orodispersible tablets 5462502 PT -
ACTAVIS GROUP PTC EHF.**

**Desloratadina Actavis 5 mg orodispersible
tablets 5462825, 2510 PT - ACTAVIS GROUP
PTC EHF.**

**Desloratadina Almus 5 mg comprimidos
recubiertos con película EFG 76305 ES -
ALMUS FARMACEUTICA S.A**

**Desloratadina Alter 5 mg comprimidos
bucodispersables EFG 77984 ES -
LABORATORIOS ALTER, S.A.**

**Desloratadina ALTER 5 mg comprimidos
orodispersíveis 5580436 PT - ALTER, S.A.**

**Desloratadina ALTER 5 mg comprimidos
orodispersíveis 5580444 PT - ALTER, S.A.**

**Desloratadina ALTER 5 mg comprimidos
orodispersíveis 5580451 PT - ALTER, S.A.**

**Desloratadina Alter 5 mg comprimidos
recubiertos con película 77557 ES -
LABORATORIOS ALTER, S.A.**

**Desloratadina ALTER 5 mg comprimidos
revestidos por película 5580469 PT - ALTER,
S.A.**

**Desloratadina ALTER 5 mg comprimidos
revestidos por película 5580477 PT - ALTER,
S.A.**

**Desloratadina ALTER 5 mg comprimidos
revestidos por película 5580501 PT - ALTER,
S.A.**

**Desloratadina Apotex 5 mg comprimidos
recubiertos con película EFG 75985 ES -
APOTEX EUROPE BV**

**Desloratadina Apotex AG 5 mg comprimidos
recubiertos con película EFG 77669 ES -
APOTEX EUROPE BV**

**Desloratadina Aristo 0,5 mg/ml solución
oral EFG 76599 ES - ARISTO PHARMA IBERIA,
S.L.**

**Desloratadina Aristo 0,5mg/ml solução oral
5568936 PT - ARISTO PHARMA IBERIA, S.L.**

Desloratadina Aristo 0,5mg/ml solução oral

5568944 PT - ARISTO PHARMA IBERIA, S.L.

Desloratadina Aristo 0,5mg/ml solução oral

5568951 PT - ARISTO PHARMA IBERIA, S.L.

Desloratadina Aristo 0,5mg/ml solução oral

5568969 PT - ARISTO PHARMA IBERIA, S.L.

Desloratadina Aristo 0,5mg/ml solução oral

5568977 PT - ARISTO PHARMA IBERIA, S.L.

Desloratadina Aristo 0,5mg/ml solução oral

5569009 PT - ARISTO PHARMA IBERIA, S.L.

Desloratadina Aristo 5 mg comprimidos

**bucodispersables EFG 77277 ES - ARISTO
PHARMA IBERIA, S.L.**

Desloratadina Aristo 5mg comprimidos

revestidos por película 5568779 PT -

ARISTO PHARMA IBERIA, S.L.

Desloratadina Aristo 5mg comprimidos

revestidos por película 5568803 PT -

ARISTO PHARMA IBERIA, S.L.

Desloratadina Aristo 5mg comprimidos

revestidos por película 5568811 PT -

ARISTO PHARMA IBERIA, S.L.

Desloratadina Aristo 5mg comprimidos

revestidos por película 5568829 PT -

ARISTO PHARMA IBERIA, S.L.

Desloratadina Aristo 5mg comprimidos

revestidos por película 5568837 PT -

ARISTO PHARMA IBERIA, S.L.

Desloratadina Aristo 5mg comprimidos

revestidos por película 5568845 PT -

ARISTO PHARMA IBERIA, S.L.

DESLORETADINA AUROBINDO 0,5 mg/ml

soluzione orale 041570019 IT - AUROBINDO

PHARMA (ITALIA) S.R.L.

DESLORETADINA AUROBINDO 0,5 mg/ml

soluzione orale 041570021 IT - AUROBINDO

PHARMA (ITALIA) S.R.L.

DESLORETADINA AUROBINDO 0,5 mg/ml

soluzione orale 041570033 IT - AUROBINDO

PHARMA (ITALIA) S.R.L.

DESLORETADINA AUROBINDO 0,5 mg/ml

soluzione orale 041570045 IT - AUROBINDO

PHARMA (ITALIA) S.R.L.

DESLORETADINA AUROBINDO 0,5 mg/ml

soluzione orale 041570058 IT - AUROBINDO

PHARMA (ITALIA) S.R.L.

DESLORETADINA AUROBINDO 0,5 mg/ml

soluzione orale 041570060 IT - AUROBINDO

PHARMA (ITALIA) S.R.L.

DESLORETADINA AUROBINDO 0,5 mg/ml

soluzione orale 041570072 IT - AUROBINDO

PHARMA (ITALIA) S.R.L.

DESLORATADINA AUROBINDO 0,5 mg/ml

soluzione orale 041570084 IT - AUROBINDO

PHARMA (ITALIA) S.R.L.

Desloratadina Azevedos 0,5 mg/ml solução

oral 5634274 PT - LABORATÓRIOS AZEVEDOS

- INDÚSTRIA FARMACÊUTICA, S.A.

Desloratadina Azevedos 0,5 mg/ml solução

oral 5634308 PT - LABORATÓRIOS AZEVEDOS

- INDÚSTRIA FARMACÊUTICA, S.A.

Desloratadina Azevedos 0,5 mg/ml solução

oral 5634316 PT - LABORATÓRIOS AZEVEDOS

- INDÚSTRIA FARMACÊUTICA, S.A.

Desloratadina Azevedos 0,5 mg/ml solução

oral 5634324 PT - LABORATÓRIOS AZEVEDOS

- INDÚSTRIA FARMACÊUTICA, S.A.

Desloratadina Azevedos 0,5 mg/ml solução

oral 5634332 PT - LABORATÓRIOS AZEVEDOS

- INDÚSTRIA FARMACÊUTICA, S.A.

Desloratadina Azevedos 0,5 mg/ml solução

oral 5634340 PT - LABORATÓRIOS AZEVEDOS

- INDÚSTRIA FARMACÊUTICA, S.A.

Desloratadina Basi 5 mg comprimidos

revestidos por película 5459433 PT -

LABORATÓRIOS BASI – INDÚSTRIA

FARMACÊUTICA, S.A.

Desloratadina Basi 5 mg comprimidos

revestidos por película 5459441 PT -

LABORATÓRIOS BASI – INDÚSTRIA

FARMACÊUTICA, S.A.

Desloratadina Basi 5 mg comprimidos

revestidos por película 5459458 PT -

LABORATÓRIOS BASI – INDÚSTRIA

FARMACÊUTICA, S.A.

Desloratadina Basi 5 mg comprimidos

revestidos por película 5459466 PT -

LABORATÓRIOS BASI – INDÚSTRIA

FARMACÊUTICA, S.A.

Desloratadina Bluelife 0,5 mg Solução oral

14/H/0084/004 PT - BLUELIFE, SOCIEDADE

UNIPESSOAL LDA.

Desloratadina Bluelife 2.5 mg Comprimido

orodispersível 14/H/0084/002 PT -

BLUELIFE, SOCIEDADE UNIPESSOAL LDA.

Desloratadina Bluelife 5 mg Comprimido

orodispersível 14/H/0084/003 PT -

BLUELIFE, SOCIEDADE UNIPESSOAL LDA.

Desloratadina Bluelife 5 mg Comprimido

revestido por película 14/H/0084/001 PT -

BLUELIFE, SOCIEDADE UNIPESSOAL LDA.

Desloratadina Bluepharma 5 mg

comprimidos revestidos por película

5428578 PT - BLUEPHARMA GENÉRICOS -
COMÉRCIO DE MEDICAMENTOS, S.A.

Desloratadina Bluepharma 5 mg

comprimidos revestidos por película

5428602 PT - BLUEPHARMA GENÉRICOS -
COMÉRCIO DE MEDICAMENTOS, S.A.

Desloratadina Bluepharma 5 mg

comprimidos revestidos por película

5428610 PT - BLUEPHARMA GENÉRICOS -
COMÉRCIO DE MEDICAMENTOS, S.A.

Desloratadina Bluepharma 5 mg

comprimidos revestidos por película

5428628 PT - BLUEPHARMA GENÉRICOS -
COMÉRCIO DE MEDICAMENTOS, S.A.

Desloratadina Bluepharma 5 mg

comprimidos revestidos por película

5428636 PT - BLUEPHARMA GENÉRICOS -
COMÉRCIO DE MEDICAMENTOS, S.A.

Desloratadina Bluepharma 5 mg

comprimidos revestidos por película

5428644 PT - BLUEPHARMA GENÉRICOS -
COMÉRCIO DE MEDICAMENTOS, S.A.

Desloratadina Bluepharma 5 mg

comprimidos revestidos por película

5428651 PT - BLUEPHARMA GENÉRICOS -
COMÉRCIO DE MEDICAMENTOS, S.A.

Desloratadina Ciclum 0,5 mg/ml solução

oral 5477740 PT - CICLUM FARMA

UNIPESSOAL, LDA

Desloratadina Ciclum 0,5 mg/ml solução

oral 5477757 PT - CICLUM FARMA

UNIPESSOAL, LDA

Desloratadina Ciclum 0,5 mg/ml solução

oral DK/H/2065/001 PT - CICLUM FARMA

UNIPESSOAL, LDA

Desloratadina Ciclum 5 mg comprimidos

revestidos por película 5477732 PT -

CICLUM FARMA UNIPESSOAL, LDA

Desloratadina Ciclum 5 mg comprimidos

revestidos por película DK/H/2050/001 PT

- CICLUM FARMA UNIPESSOAL, LDA

desloratadina cinfa 5 mg comprimidos

bucodispersables EFG. 78370 ES -

LABORATORIOS CINFA, S.A.

desloratadina cinfa 5 mg comprimidos

recubiertos con película EFG 75335 ES -

LABORATORIOS CINFA, S.A.

Desloratadina Cinfa 5 mg comprimidos

revestidos por película 5458906 PT - CINFA

PORTUGAL, LDA.

Desloratadina Cipla 5 mg compresse

rivestite con film 044367011 IT - CIPLA

EUROPE NV

Desloratadina Cipla 5 mg compresse

rivestite con film 044367023 IT - CIPLA

EUROPE NV

Desloratadina Cipla 5 mg compresse

rivestite con film 044367035 IT - CIPLA

EUROPE NV

Desloratadina Cipla 5 mg compresse

rivestite con film 044367047 IT - CIPLA

EUROPE NV

Desloratadina Cipla 5 mg compresse

rivestite con film 044367050 IT - CIPLA

EUROPE NV

Desloratadina Cipla 5 mg compresse

rivestite con film 044367062 IT - CIPLA

EUROPE NV

Desloratadina Cipla 5 mg comprimate

filmate 8741/2016/06 RO - CIPLA EUROPE

NV

Desloratadina Cipla 5 mg comprimidos

recubiertos con película EFG 80568 ES -

CIPLA EUROPE NV

Desloratadina Combix 5 mg comprimidos

recubiertos con película EFG 76.043 ES -

LABORATORIOS COMBIX, S.L.U.

DESLORATADINA DOC 5 mg compresse

rivestite con film 040718013 IT - DOC

GENERICI S.R.L.

DESLORATADINA DOC 5 mg compresse

rivestite con film 040718025 IT - DOC

GENERICI S.R.L.

DESLORATADINA DOC 5 mg compresse

rivestite con film 040718037 IT - DOC

GENERICI S.R.L.

DESLORATADINA DOC 5 mg compresse

rivestite con film 040718049 IT - DOC

GENERICI S.R.L.

DESLORATADINA DOC 5 mg compresse

rivestite con film 040718052 IT - DOC

GENERICI S.R.L.

DESLORATADINA DOC Generici 2.5 mg

compresse orodispersibili 040810057 IT -

DOC GENERICI S.R.L.

DESLORATADINA DOC Generici 5 mg

compresse orodispersibili 040810 IT - DOC

GENERICI S.R.L.

DESLORATADINA DOC Generici 5 mg

compresse rivestite con film 040810 IT -

DOC GENERICI S.R.L.

DESLORATADINA EG 5 mg compresse
rivestite con film 040733014 IT - EG SPA
DESLORATADINA EG 5 mg compresse
rivestite con film 040733026 IT - EG SPA
DESLORATADINA EG 5 mg compresse
rivestite con film 040733038 IT - EG SPA
DESLORATADINA EG 5 mg compresse
rivestite con film 040733040 IT - EG SPA
DESLORATADINA EG 5 mg compresse
rivestite con film 040733053 IT - EG SPA
DESLORATADINA EG 5 mg compresse
rivestite con film 040733065 IT - EG SPA
DESLORATADINA EG 5 mg compresse
rivestite con film 040733077 IT - EG SPA
DESLORATADINA EG 5 mg compresse
rivestite con film 040733089 IT - EG SPA
DESLORATADINA EG 5 mg compresse
rivestite con film 040733091 IT - EG SPA
DESLORATADINA EG 5 mg compresse
rivestite con film 040733103 IT - EG SPA
DESLORATADINA EG 5 mg compresse
rivestite con film 040733115 IT - EG SPA
DESLORATADINA EG 5 mg compresse
rivestite con film 040733127 IT - EG SPA
Desloratadina Farmoz 0,5 mg/ml solução oral 5451059 PT - FARMOZ - SOCIEDADE TÉCNICO MEDICINAL, S.A.
Desloratadina Farmoz 5 mg comprimidos revestidos por película 5451067 PT - FARMOZ - SOCIEDADE TÉCNICO MEDICINAL, S.A.
Desloratadina Farmoz 5 mg comprimidos revestidos por película 5451075 PT - FARMOZ - SOCIEDADE TÉCNICO MEDICINAL, S.A.
Desloratadina Flas Combix 5 mg comprimidos bucodispersables EFG 76.037 ES - LABORATORIOS COMBIX, S.L.U.
Desloratadina Genepharm 0,5 mg/ml solução oral 5657028 PT - GENEPHARM S.A.
Desloratadina Genepharm 0,5 mg/ml solução oral 5657036 PT - GENEPHARM S.A.
Desloratadina Genepharm 0,5 mg/ml solução oral 5657044 PT - GENEPHARM S.A.
Desloratadina Genepharm 0,5 mg/ml solução oral 5657051 PT - GENEPHARM S.A.
Desloratadina Genepharm 0,5 mg/ml solução oral 5657077 PT - GENEPHARM S.A.
Desloratadina Genepharm 0,5 mg/ml solução oral 5657101 PT - GENEPHARM S.A.
Desloratadina Genepharm 0,5 mg/ml

solução oral 5657119 PT - GENEPHARM S.A.
Desloratadina Genepharm 0,5 mg/ml
solução oral 5657127 PT - GENEPHARM S.A.
Desloratadina Genepharm 0,5 mg/ml
solução oral 5657135 PT - GENEPHARM S.A.
Desloratadina Genepharm 0,5 mg/ml
solução oral 5657143 PT - GENEPHARM S.A.
Desloratadina Genepharm 0,5 mg/ml
solução oral 5657150 PT - GENEPHARM S.A.
Desloratadina Genepharm 0,5 mg/ml
solução oral 5657168 PT - GENEPHARM S.A.
Desloratadina Genepharm 0,5 mg/ml
solução oral 5657169 PT - GENEPHARM S.A.
Desloratadina Genepharm 0,5 mg/ml
solução oral 5657176 PT - GENEPHARM S.A.
Desloratadina Genepharm 2,5 mg
comprimidos orodispersíveis 5582077 PT -
GENEPHARM S.A.
Desloratadina Genepharm 5 mg
comprimidos orodispersíveis 5582101 PT -
GENEPHARM S.A.
Desloratadina Genepharm 5 mg
comprimidos revestidos por película
5582069 PT - GENEPHARM S.A.
Desloratadina Generis 0,5 mg/ml solucao
oral 5415351 PT - GENERIS FARMACÉUTICA,
S.A.
Desloratadina Generis 0,5 mg/ml solucao
oral 5415369 PT - GENERIS FARMACÉUTICA,
S.A.
Desloratadina Generis 0,5 mg/ml solucao
oral 5415377 PT - GENERIS FARMACÉUTICA,
S.A.
Desloratadina Generis 0,5 mg/ml solucao
oral 5415401 PT - GENERIS FARMACÉUTICA,
S.A.
Desloratadina Generis 0,5 mg/ml solucao
oral 5415419 PT - GENERIS FARMACÉUTICA,
S.A.
Desloratadina Generis 0,5 mg/ml solucao
oral 5417522 PT - GENERIS FARMACÉUTICA,
S.A.
Desloratadina Generis 0,5 mg/ml solucao
oral 5417530 PT - GENERIS FARMACÉUTICA,
S.A.
Desloratadina Generis 0,5 mg/ml solucao
oral 5417548 PT - GENERIS FARMACÉUTICA,
S.A.
Desloratadina Generis 0,5 mg/ml solucao
oral 5417555 PT - GENERIS FARMACÉUTICA,
S.A.

Desloratadina Generis 0,5 mg/ml solucao

**oral 5418215 PT - GENERIS FARMACÉUTICA,
S.A.**

**Desloratadina Generis 2,5 mg comprimidos
orodispersíveis 5413315 PT - GENERIS
FARMACÉUTICA, S.A.**

**Desloratadina Generis 2,5 mg comprimidos
orodispersíveis 5442504 PT - GENERIS
FARMACÉUTICA, S.A.**

**Desloratadina Generis 5 mg comprimidos
orodispersíveis 5413323 PT - GENERIS
FARMACÉUTICA, S.A.**

**Desloratadina Generis 5 mg comprimidos
revestidos por película 5413331 PT -
GENERIS FARMACÉUTICA, S.A.**

**Desloratadina Germed 0,5 mg/ml solución
oral EFG 75521 ES - ARISTO PHARMA IBERIA,
S.L.**

**Desloratadina Germed 0,5 mg/ml solución
oral EFG 75521 ES - ARISTO PHARMA IBERIA,
S.L.**

**Desloratadina Germed 0,5 mg/ml solução
oral 5469606 PT - GERMED FARMACÉUTICA,
LDA.**

**Desloratadina Germed 0,5 mg/ml solução
oral 5469614 PT - GERMED FARMACÉUTICA,
LDA.**

**Desloratadina Germed 0,5 mg/ml solução
oral 5469622 PT - GERMED FARMACÉUTICA,
LDA.**

**Desloratadina Germed 0,5 mg/ml soluzione
orale 040983088 IT - GERMED PHARMA S.R.L.**

**Desloratadina Germed 0,5 mg/ml soluzione
orale 040983090 IT - GERMED PHARMA S.R.L.**

**Desloratadina Germed 0,5 mg/ml soluzione
orale 040983102 IT - GERMED PHARMA S.R.L.**

**Desloratadina Germed 2,5 mg compresse
orodispersibili 040983049 IT - GERMED
PHARMA S.R.L.**

**Desloratadina Germed 2,5 mg compresse
orodispersibili 040983052 IT - GERMED
PHARMA S.R.L.**

**Desloratadina Germed 2,5 mg compresse
orodispersibili 040983064 IT - GERMED
PHARMA S.R.L.**

**Desloratadina Germed 2,5 mg comprimidos
orodispersíveis 5469549 PT - GERMED
FARMACÉUTICA, LDA.**

**Desloratadina Germed 2,5 mg comprimidos
orodispersíveis 5469556 PT - GERMED
FARMACÉUTICA, LDA.**

Desloratadina Germed 2,5 mg comprimidos

orodispersíveis 5469564 PT - GERMED

FARMACÊUTICA, LDA.

Desloratadina Germed 5 mg compresse

orodispersibili 040983076 IT - GERMED

PHARMA S.R.L.

Desloratadina Germed 5 mg compresse

rivestite con film 040983013 IT - GERMED

PHARMA S.R.L.

Desloratadina Germed 5 mg compresse

rivestite con film 040983025 IT - GERMED

PHARMA S.R.L.

Desloratadina Germed 5 mg compresse

rivestite con film 040983037 IT - GERMED

PHARMA S.R.L.

Desloratadina Germed 5 mg comprimidos

bucodispersables EFG 75523 ES - ARISTO

PHARMA IBERIA, S.L.

Desloratadina Germed 5 mg comprimidos

bucodispersables EFG 75523 ES - ARISTO

PHARMA IBERIA, S.L.

Desloratadina Germed 5 mg comprimidos

orodispersíveis 5469572 PT - GERMED

FARMACÊUTICA, LDA.

Desloratadina Germed 5 mg comprimidos

recubiertos con película EFG 75522 ES -

ARISTO PHARMA IBERIA, S.L.

Desloratadina Germed 5 mg comprimidos

recubiertos con película EFG 75522 ES -

ARISTO PHARMA IBERIA, S.L.

Desloratadina Germed 5 mg comprimidos

revestidos por película 5469515 PT -

GERMED FARMACÊUTICA, LDA.

Desloratadina Germed 5 mg comprimidos

revestidos por película 5469523 PT -

GERMED FARMACÊUTICA, LDA.

Desloratadina Germed 5 mg comprimidos

revestidos por película 5469531 PT -

GERMED FARMACÊUTICA, LDA.

Desloratadina Glenmark 5 mg Comprimidos

5643200 PT - GLENMARK PHARMACEUTICALS

EUROPE LIMITED

Desloratadina Kern Pharma 0,5 mg/ml

solución oral EFG 77592 ES - KERN PHARMA,

S.L.

Desloratadina KERN PHARMA 5 mg

comprimidos recubiertos con película EFG

75.304 ES - KERN PHARMA, S.L.

Desloratadina Krka 5 mg Comprimido

revestido por película 5398953 PT - KRKA

FARMACÊUTICA, UNIPESSOAL LDA.

Desloratadina Krka 5 mg Comprimido
revestido por película 5398961 PT - KRKA
FARMACÉUTICA, UNIPESSOAL LDA.

Desloratadina Krka 5 mg Comprimido
revestido por película 5399233 PT - KRKA
FARMACÉUTICA, UNIPESSOAL LDA.

Desloratadina Labesfal 0,5 mg/ml Solução
oral 5472857 PT - GENERIS FARMACÉUTICA,
S.A.

Desloratadina Labesfal 0,5 mg/ml Solução
oral 5472865 PT - GENERIS FARMACÉUTICA,
S.A.

Desloratadina Labesfal 5 mg comprimidos
revestidos por película 5428412 PT -
GENERIS FARMACÉUTICA, S.A.

Desloratadina Labesfal 5 mg comprimidos
revestidos por película 5428420 PT -
GENERIS FARMACÉUTICA, S.A.

Desloratadina Labesfal 5 mg comprimidos
revestidos por película 5428438 PT -
GENERIS FARMACÉUTICA, S.A.

Desloratadina Labesfal 5 mg comprimidos
revestidos por película 5428446 PT -
GENERIS FARMACÉUTICA, S.A.

Desloratadina Labesfal 5 mg comprimidos
revestidos por película 5428453 PT -
GENERIS FARMACÉUTICA, S.A.

Desloratadina Labesfal 5 mg comprimidos
revestidos por película 5428461 PT -
GENERIS FARMACÉUTICA, S.A.

Desloratadina Labesfal 5 mg comprimidos
revestidos por película 5428479 PT -
GENERIS FARMACÉUTICA, S.A.

Desloratadina Labesfal OD 2,5 mg
Comprimido orodispersível 5472873 PT -
GENERIS FARMACÉUTICA, S.A.

Desloratadina Labesfal OD 2,5 mg
Comprimido orodispersível 5472907 PT -
GENERIS FARMACÉUTICA, S.A.

Desloratadina Labesfal OD 5 mg
Comprimidos orodispersíveis 5472915 PT -
GENERIS FARMACÉUTICA, S.A.

Desloratadina Lesvi 5 mg comprimidos
recubiertos con película EFG 75529 ES -
LABORATORIOS LESVI, S.L.

Desloratadina Lupin 5 mg compresse
rivestite con film 040827014 IT - LUPIN
(EUROPE) LIMITED

Desloratadina Lupin 5 mg compresse
rivestite con film 040827026 IT - LUPIN
(EUROPE) LIMITED

**Desloratadina Lupin 5 mg compresse
rivestite con film 040827038 IT - LUPIN
(EUROPE) LIMITED**

**Desloratadina Lupin 5 mg compresse
rivestite con film 040827040 IT - LUPIN
(EUROPE) LIMITED**

**Desloratadina Lupin 5 mg compresse
rivestite con film 040827053 IT - LUPIN
(EUROPE) LIMITED**

**Desloratadina Lupin 5 mg compresse
rivestite con film 040827065 IT - LUPIN
(EUROPE) LIMITED**

**Desloratadina Lupin 5 mg compresse
rivestite con film 040827077 IT - LUPIN
(EUROPE) LIMITED**

**Desloratadina Lupin 5 mg compresse
rivestite con film 040827089 IT - LUPIN
(EUROPE) LIMITED**

**Desloratadina Lupin 5 mg compresse
rivestite con film 040827091 IT - LUPIN
(EUROPE) LIMITED**

**Desloratadina Lupin 5 mg compresse
rivestite con film 040827103 IT - LUPIN
(EUROPE) LIMITED**

**Desloratadina Lupin 5 mg compresse
rivestite con film 040827115 IT - LUPIN
(EUROPE) LIMITED**

**Desloratadina Lupin 5 mg compresse
rivestite con film 040827127 IT - LUPIN
(EUROPE) LIMITED**

**Desloratadina Lupin 5 mg compresse
rivestite con film 040827139 IT - LUPIN
(EUROPE) LIMITED**

**Desloratadina Lupin 5 mg compresse
rivestite con film 040827141 IT - LUPIN
(EUROPE) LIMITED**

**Desloratadina Lupin 5 mg compresse
rivestite con film 040827154 IT - LUPIN
(EUROPE) LIMITED**

**Desloratadina Lupin 5 mg compresse
rivestite con film 040827166 IT - LUPIN
(EUROPE) LIMITED**

**Desloratadina Lupin 5 mg compresse
rivestite con film 040827178 IT - LUPIN
(EUROPE) LIMITED**

**Desloratadina Lupin 5 mg compresse
rivestite con film 040827180 IT - LUPIN
(EUROPE) LIMITED**

**Desloratadina Mepha, 0,5 mg/ml, solução
oral 5415427 PT - MEPA-INVESTIGACAO
DESENVOLVIMENTO E FABRICACAO**

FARMACEUTICA, LDA.

Desloratadina Mepha, 0,5 mg/ml, solução

oral 5415435 PT - MEPHA-INVESTIGACAO

DESENVOLVIMENTO E FABRICACAO

FARMACEUTICA, LDA.

Desloratadina Mepha, 0,5 mg/ml, solução

oral 5415443 PT - MEPHA-INVESTIGACAO

DESENVOLVIMENTO E FABRICACAO

FARMACEUTICA, LDA.

Desloratadina Mepha, 0,5 mg/ml, solução

oral 5415450 PT - MEPHA-INVESTIGACAO

DESENVOLVIMENTO E FABRICACAO

FARMACEUTICA, LDA.

Desloratadina Mepha, 0,5 mg/ml, solução

oral 5415468 PT - MEPHA-INVESTIGACAO

DESENVOLVIMENTO E FABRICACAO

FARMACEUTICA, LDA.

Desloratadina Mepha, 0,5 mg/ml, solução

oral 5418223 PT - MEPHA-INVESTIGACAO

DESENVOLVIMENTO E FABRICACAO

FARMACEUTICA, LDA.

Desloratadina Mepha, 0,5 mg/ml, solução

oral 5418231 PT - MEPHA-INVESTIGACAO

DESENVOLVIMENTO E FABRICACAO

FARMACEUTICA, LDA.

Desloratadina Mepha, 0,5 mg/ml, solução

oral 5418249 PT - MEPHA-INVESTIGACAO

DESENVOLVIMENTO E FABRICACAO

FARMACEUTICA, LDA.

Desloratadina Mepha, 0,5 mg/ml, solução

oral 5418256 PT - MEPHA-INVESTIGACAO

DESENVOLVIMENTO E FABRICACAO

FARMACEUTICA, LDA.

Desloratadina Mepha, 0,5 mg/ml, solução

oral 5418264 PT - MEPHA-INVESTIGACAO

DESENVOLVIMENTO E FABRICACAO

FARMACEUTICA, LDA.

Desloratadina Mepha, 5 mg, comprimidos

revestidos por película 5413307 PT -

MEPHA-INVESTIGACAO DESENVOLVIMENTO E

FABRICACAO FARMACEUTICA, LDA.

Desloratadina MYLAN 0,5 mg/ml solución

oral EFG 76343 ES - MYLAN

PHARMACEUTICALS S.L.

Desloratadina Mylan 0,5 mg/ml solução

oral 5439872 PT - MYLAN, LDA

Desloratadina MYLAN 5 mg comprimidos

recubiertos con película EFG 76625 ES -

MYLAN PHARMACEUTICALS S.L.

Desloratadina Mylan 5 mg comprimidos

revestidos por película 5415476 PT - MYLAN,

LDA

**Desloratadina Mylan 5 mg comprimidos
revestidos por película 5415500 PT - MYLAN,**
LDA

**Desloratadina Mylan 5 mg comprimidos
revestidos por película 5415518 PT - MYLAN,**
LDA

**Desloratadina Mylan Generics 5 mg
compresse rivestite con film 041081011/M**

IT - MYLAN S.P.A.

**Desloratadina Mylan Generics 5 mg
compresse rivestite con film 041081023/M**

IT - MYLAN S.P.A.

**Desloratadina Mylan Generics 5 mg
compresse rivestite con film 041081035/M**

IT - MYLAN S.P.A.

**Desloratadina Mylan Generics 5 mg
compresse rivestite con film 041081047/M**

IT - MYLAN S.P.A.

**Desloratadina Mylan Generics 5 mg
compresse rivestite con film 041081050/M**

IT - MYLAN S.P.A.

**Desloratadina Mylan Generics 5 mg
compresse rivestite con film 041081062/M**

IT - MYLAN S.P.A.

**Desloratadina Mylan Generics 5 mg
compresse rivestite con film 041081074/M**

IT - MYLAN S.P.A.

**Desloratadina Mylan Generics 5 mg
compresse rivestite con film 041081086/M**

IT - MYLAN S.P.A.

**Desloratadina Mylan Generics 5 mg
compresse rivestite con film 041081098/M**

IT - MYLAN S.P.A.

**Desloratadina Mylan Generics 5 mg
compresse rivestite con film 041081100/M**

IT - MYLAN S.P.A.

**Desloratadina Mylan Generics 5 mg
compresse rivestite con film 041081112/M**

IT - MYLAN S.P.A.

**Desloratadina Mylan Generics 5 mg
compresse rivestite con film 041081124/M**

IT - MYLAN S.P.A.

**Desloratadina NORMON 0,5 mg/ml solución
oral EFG 75842 ES - LABORATORIOS**
NORMON, S.A.

**Desloratadina Normon 0,5 mg/ml solução
oral 5664933 PT - LABORATÓRIOS NORMON,
S.A.**

**Desloratadina NORMON 5 mg comprimidos
recubiertos con película EFG 75840 ES -**

LABORATORIOS NORMON, S.A.

Desloratadina Normon 5 mg comprimidos

revestidos por película 5664925 PT -

LABORATÓRIOS NORMON, S.A.

Desloratadina Pharmakern 0,5 mg/ml

solução oral 5558150 PT - PHARMAKERN

PORTUGAL – PRODUTOS FARMACÊUTICOS,

SOCIEDADE UNIPESSOAL, LDA.

Desloratadina Pharmakern 0,5 mg/ml

solução oral 5558168 PT - PHARMAKERN

PORTUGAL – PRODUTOS FARMACÊUTICOS,

SOCIEDADE UNIPESSOAL, LDA.

Desloratadina Pharmakern 5 mg

comprimidos revestidos por película

5450721 PT - PHARMAKERN PORTUGAL –

PRODUTOS FARMACÊUTICOS, SOCIEDADE

UNIPESSOAL, LDA.

Desloratadina Pharmakern 5 mg

comprimidos revestidos por película

5450739 PT - PHARMAKERN PORTUGAL –

PRODUTOS FARMACÊUTICOS, SOCIEDADE

UNIPESSOAL, LDA.

Desloratadina Qualigen 0.5 mg/ml solución

oral EFG 76.761 ES - QUALIGEN, S.L.

Desloratadina Qualigen 5 mg comprimidos

recubiertos con película EFG 75542 ES -

QUALIGEN, S.L.

Desloratadina ratiopharm 0,5 mg/ml

solución oral EFG 77664 ES - RATIOFARM

ESPAÑA SA

Desloratadina ratiopharm 0,5 mg/ml

Solução Oral 5580253 PT - RATIOFARM-

COMERCIO E INDUSTRIA DE PRODUTOS

FARMACEUTICOS LDA

Desloratadina ratiopharm 2,5 mg

comprimidos orodispersíveis 5492137 PT -

RATIOFARM-COMERCIO E INDUSTRIA DE

PRODUTOS FARMACEUTICOS LDA

Desloratadina ratiopharm 5 mg

comprimidos orodispersíveis 5492145 PT -

RATIOFARM-COMERCIO E INDUSTRIA DE

PRODUTOS FARMACEUTICOS LDA

Desloratadina Sandoz 0,5mg/ml solución

oral EFG 76283 ES - SANDOZ FARMACÉUTICA,

S.A.

DESLORATADINA SANDOZ 040722011 IT -

SANDOZ S.P.A.

DESLORATADINA SANDOZ 040722023 IT -

SANDOZ S.P.A.

DESLORATADINA SANDOZ 040722035 IT -

SANDOZ S.P.A.

DESLORATADINA SANDOZ 040722047 IT -

SANDOZ S.P.A.

DESLORATADINA SANDOZ 040722050 IT -

SANDOZ S.P.A.

DESLORATADINA SANDOZ 040722062 IT -

SANDOZ S.P.A.

DESLORATADINA SANDOZ 040722074 IT -

SANDOZ S.P.A.

DESLORATADINA SANDOZ 040722086 IT -

SANDOZ S.P.A.

DESLORATADINA SANDOZ 040722098 IT -

SANDOZ S.P.A.

DESLORATADINA SANDOZ 040722100 IT -

SANDOZ S.P.A.

DESLORATADINA SANDOZ 040722112 IT -

SANDOZ S.P.A.

DESLORATADINA SANDOZ 040722124 IT -

SANDOZ S.P.A.

DESLORATADINA SANDOZ 040722136 IT -

SANDOZ S.P.A.

DESLORATADINA SANDOZ 040722148 IT -

SANDOZ S.P.A.

DESLORATADINA SANDOZ 040722151 IT -

SANDOZ S.P.A.

DESLORATADINA SANDOZ 040722163 IT -

SANDOZ S.P.A.

DESLORATADINA SANDOZ 040722175 IT -

SANDOZ S.P.A.

DESLORATADINA SANDOZ 040722187 IT -

SANDOZ S.P.A.

DESLORATADINA SANDOZ 040722199 IT -

SANDOZ S.P.A.

DESLORATADINA SANDOZ 040722201 IT -

SANDOZ S.P.A.

DESLORATADINA SANDOZ 040722213 IT -

SANDOZ S.P.A.

DESLORATADINA SANDOZ 040722225 IT -

SANDOZ S.P.A.

DESLORATADINA SANDOZ 040722237 IT -

SANDOZ S.P.A.

DESLORATADINA SANDOZ 040722249 IT -

SANDOZ S.P.A.

DESLORATADINA SANDOZ 040722252 IT -

SANDOZ S.P.A.

DESLORATADINA SANDOZ 040722264 IT -

SANDOZ S.P.A.

DESLORATADINA SANDOZ 040722276 IT -

SANDOZ S.P.A.

DESLORATADINA SANDOZ 040722288 IT -

SANDOZ S.P.A.

DESLORATADINA SANDOZ 040722290/M-

040722340 IT - SANDOZ S.P.A.
DESLORATADINA SANDOZ 040722326/M-
040722377 IT - SANDOZ S.P.A.
Desloratadina Sandoz 5 mg comprimidos
recubiertos EFG 76284 ES - SANDOZ
FARMACÉUTICA, S.A.
Desloratadina Sandoz 5461348 PT -
SANDOZ FARMACÉUTICA LDA.
Desloratadina Sandoz 5461355 PT -
SANDOZ FARMACÉUTICA LDA.
Desloratadina Sandoz 5461363 PT -
SANDOZ FARMACÉUTICA LDA.
Desloratadina Sandoz 5461371 PT -
SANDOZ FARMACÉUTICA LDA.
Desloratadina Sandoz 5461405 PT -
SANDOZ FARMACÉUTICA LDA.
Desloratadina Sandoz 5461413 PT -
SANDOZ FARMACÉUTICA LDA.
Desloratadina Sandoz 5461421 PT -
SANDOZ FARMACÉUTICA LDA.
Desloratadina Specifar 5 mg comprimidos
recubiertos con película EFG 76.344 ES -
SPECIFAR S.A.
Desloratadina STADA 0,5 mg/ml solución
oral EFG 75.387 ES - LABORATORIO STADA,
S.L.
Desloratadina STADA 5 mg comprimidos
recubiertos con película EFG 75.528 ES -
LABORATORIO STADA, S.L.
Desloratadina Tarbis 0,5 mg/ml solución
oral EFG 76340 ES - TARBIS FARMA, S.L.
Desloratadina Tarbis 5 mg comprimidos
recubiertos con película EFG 76174 ES -
TARBIS FARMA, S.L.
Desloratadina Tecnigen 0,5 mg/ml solución
oral EFG 76.331 ES - TECNIMEDE ESPAÑA,
INDUSTRIA FARMACÉUTICA, S.A.
Desloratadina TecniGen 5 mg comprimidos
recubiertos con película EFG 76.332 ES -
TECNIMEDE ESPAÑA IND. FCA., S.A.
Desloratadina TecniGen 5 mg comprimidos
recubiertos con película EFG 76.332 ES -
TECNIMEDE ESPAÑA, INDUSTRIA
FARMACÉUTICA, S.A.
Desloratadina Teva 0,5 mg/ml solución oral
EFG 77666 ES - TEVA PHARMA S.L.U
Desloratadina Teva 5 mg comprimidos
bucodispersables EFG 76005 ES - TEVA
PHARMA S.L.U
Desloratadina Teva 5 mg comprimidos
orodispersíveis 5458609 PT - TEVA PHARMA

– PRODUTOS FARMACÊUTICOS LDA

Desloratadina toLife 2,5 mg comprimidos

orodispersíveis 5413265 PT - TOLIFE -

PRODUTOS FARMACÊUTICOS, S.A.

Desloratadina toLife 2,5 mg comprimidos

orodispersíveis 5442512 PT - TOLIFE -

PRODUTOS FARMACÊUTICOS, S.A.

Desloratadina toLife 5 mg comprimidos

orodispersíveis 5413273 PT - TOLIFE -

PRODUTOS FARMACÊUTICOS, S.A.

Desloratadina toLife 5 mg comprimidos

revestidos por película 5443320 PT - TOLIFE

- PRODUTOS FARMACÊUTICOS, S.A.

Desloratadina Viso Farmacéutica 5 mg

comprimidos EFG 76148 ES - GLENMARK

PHARMACEUTICALS EUROPE LIMITED

Desloratadina Worldrugs 0,5 mg/ml

solução oral 5646328 PT - WORLDRUGS,

LDA.

Desloratadina Zentiva 5 mg compresse

rivestite con film 040872018 IT - ZENTIVA

ITALIA SRL

Desloratadina Zentiva 5 mg compresse

rivestite con film 040872020 IT - ZENTIVA

ITALIA SRL

Desloratadina Zentiva 5 mg compresse

rivestite con film 040872032 IT - ZENTIVA

ITALIA SRL

Desloratadina Zentiva 5 mg compresse

rivestite con film 040872044 IT - ZENTIVA

ITALIA SRL

Desloratadina Zentiva 5 mg compresse

rivestite con film 040872057 IT - ZENTIVA

ITALIA SRL

Desloratadina Zentiva 5 mg compresse

rivestite con film 040872069 IT - ZENTIVA

ITALIA SRL

Desloratadina Zentiva 5 mg compresse

rivestite con film 040872071 IT - ZENTIVA

ITALIA SRL

Desloratadina Zentiva 5 mg compresse

rivestite con film 040872083 IT - ZENTIVA

ITALIA SRL

Desloratadina Zentiva 5 mg compresse

rivestite con film 040872095 IT - ZENTIVA

ITALIA SRL

Desloratadina Zentiva 5 mg compresse

rivestite con film 040872107 IT - ZENTIVA

ITALIA SRL

Desloratadină Alvogen 0,5 mg/ ml soluție

orală 4535/2012/01-08 RO - ALVOGEN IPCO

S.AR.L

Desloratadină Alvogen 5 mg comprimate

filmate 4534/2012/01-28 RO - ALVOGEN

IPCO S.AR.L

DESLORATADINĂ SANDOZ 0.5 mg soluție

orală 4660/2012/01 RO - S.C. SANDOZ

S.R.L.

DESLORATADINĂ SANDOZ 0.5 mg soluție

orală 4660/2012/02 RO - S.C. SANDOZ

S.R.L.

DESLORATADINĂ SANDOZ 0.5 mg soluție

orală 4660/2012/03 RO - S.C. SANDOZ

S.R.L.

DESLORATADINĂ SANDOZ 0.5 mg soluție

orală 4660/2012/04 RO - S.C. SANDOZ

S.R.L.

DESLORATADINĂ SANDOZ 0.5 mg soluție

orală 4660/2012/05 RO - S.C. SANDOZ

S.R.L.

DESLORATADINĂ SANDOZ 5 mg

comprimate filmate 4659/2012/01 RO -

S.C. SANDOZ S.R.L.

DESLORATADINĂ SANDOZ 5 mg

comprimate filmate 4659/2012/02 RO -

S.C. SANDOZ S.R.L.

DESLORATADINĂ SANDOZ 5 mg

comprimate filmate 4659/2012/03 RO -

S.C. SANDOZ S.R.L.

DESLORATADINĂ SANDOZ 5 mg

comprimate filmate 4659/2012/04 RO -

S.C. SANDOZ S.R.L.

DESLORATADINĂ SANDOZ 5 mg

comprimate filmate 4659/2012/05 RO -

S.C. SANDOZ S.R.L.

DESLORATADINĂ SANDOZ 5 mg

comprimate filmate 4659/2012/06 RO -

S.C. SANDOZ S.R.L.

DESLORATADINĂ SANDOZ 5 mg

comprimate filmate 4659/2012/07 RO -

S.C. SANDOZ S.R.L.

DESLORATADINĂ SANDOZ 5 mg

comprimate filmate 4659/2012/08 RO -

S.C. SANDOZ S.R.L.

DESLORATADINĂ SANDOZ 5 mg

comprimate filmate 4659/2012/09 RO -

S.C. SANDOZ S.R.L.

DESLORATADINĂ SANDOZ 5 mg

comprimate filmate 4659/2012/10 RO -

S.C. SANDOZ S.R.L.

DESLORATADINĂ SANDOZ 5 mg

comprimate filmate 4659/2012/11 RO -

S.C. SANDOZ S.R.L.

DESLORATADINĂ SANDOZ 5 mg

comprimate filmate 4659/2012/12 RO -

S.C. SANDOZ S.R.L.

DESLORATADINĂ SANDOZ 5 mg

comprimate filmate 4659/2012/13 RO -

S.C. SANDOZ S.R.L.

DESLORATADINĂ SANDOZ 5 mg

comprimate filmate 4659/2012/14 RO -

S.C. SANDOZ S.R.L.

DESLORATADINĂ SANDOZ 5 mg

comprimate filmate 4659/2012/15 RO -

S.C. SANDOZ S.R.L.

DESLORATADINĂ SANDOZ 5 mg

comprimate filmate 4659/2012/16 RO -

S.C. SANDOZ S.R.L.

DESLORATADINĂ SANDOZ 5 mg

comprimate filmate 4659/2012/17 RO -

S.C. SANDOZ S.R.L.

DESLORATADINĂ SANDOZ 5 mg

comprimate filmate 4659/2012/18 RO -

S.C. SANDOZ S.R.L.

DESLORATADINĂ SANDOZ 5 mg

comprimate filmate 4659/2012/19 RO -

S.C. SANDOZ S.R.L.

DESLORATADINĂ SANDOZ 5 mg

comprimate filmate 4659/2012/20 RO -

S.C. SANDOZ S.R.L.

DESLORATADINĂ SANDOZ 5 mg

comprimate filmate 4659/2012/21 RO -

S.C. SANDOZ S.R.L.

DESLORATADINĂ SANDOZ 5 mg

comprimate filmate 4659/2012/22 RO -

S.C. SANDOZ S.R.L.

DESLORATADINĂ SANDOZ 5 mg

comprimate filmate 4659/2012/23 RO -

S.C. SANDOZ S.R.L.

DESLORATADINĂ SANDOZ 5 mg

comprimate filmate 4659/2012/24 RO -

S.C. SANDOZ S.R.L.

DESLORATADINĂ SANDOZ 5 mg

comprimate filmate 4659/2012/25 RO -

S.C. SANDOZ S.R.L.

DESLORATADINĂ SANDOZ 5 mg

comprimate filmate 4659/2012/26 RO -

S.C. SANDOZ S.R.L.

DESLORATADINĂ SANDOZ 5 mg

comprimate filmate 4659/2012/27 RO -

S.C. SANDOZ S.R.L.

DESLORATADINĂ SANDOZ 5 mg

comprimate filmate 4659/2012/28 RO -

S.C. SANDOZ S.R.L.

DESLORATADINĂ SANDOZ 5 mg

comprimate filmate 4659/2012/29 RO -

S.C. SANDOZ S.R.L.

DESLORATADINĂ SANDOZ 5 mg

comprimate filmate 4659/2012/30 RO -

S.C. SANDOZ S.R.L.

DESLORATADINĂ SANDOZ 5 mg

comprimate filmate 4659/2012/31 RO -

S.C. SANDOZ S.R.L.

DESLORATADINĂ SANDOZ 5 mg

comprimate filmate 4659/2012/32 RO -

S.C. SANDOZ S.R.L.

DESLORATADINĂ SANDOZ 5 mg

comprimate filmate 4659/2012/33 RO -

S.C. SANDOZ S.R.L.

DESLORATADINĂ SANDOZ 5 mg

comprimate filmate 4659/2012/34 RO -

S.C. SANDOZ S.R.L.

DESLORATADINĂ SANDOZ 5 mg

comprimate filmate 4659/2012/35 RO -

S.C. SANDOZ S.R.L.

DESLORATADINĂ SANDOZ 5 mg

comprimate filmate 4659/2012/36 RO -

S.C. SANDOZ S.R.L.

DESLORATADINĂ SANDOZ 5 mg

comprimate filmate 4659/2012/37 RO -

S.C. SANDOZ S.R.L.

DESLORATADINĂ SANDOZ 5 mg

comprimate filmate 4659/2012/38 RO -

S.C. SANDOZ S.R.L.

DESLORATADINĂ SANDOZ 5 mg

comprimate filmate 4659/2012/39 RO -

S.C. SANDOZ S.R.L.

DESLORATADINĂ SANDOZ 5 mg

comprimate filmate 4659/2012/40 RO -

S.C. SANDOZ S.R.L.

DESLORATADINĂ SANDOZ 5 mg

comprimate filmate 4659/2012/41 RO -

S.C. SANDOZ S.R.L.

DESLORATADINĂ SANDOZ 5 mg

comprimate filmate 4659/2012/42 RO -

S.C. SANDOZ S.R.L.

DESLORATADINĂ SANDOZ 5 mg

comprimate filmate 4659/2012/43 RO -

S.C. SANDOZ S.R.L.

DESLORATADINĂ SANDOZ 5 mg

comprimate filmate 4659/2012/44 RO -

S.C. SANDOZ S.R.L.

DESLORATADINĂ SANDOZ 5 mg

comprimate filmate 4659/2012/45 RO -

S.C. SANDOZ S.R.L.

DESLORATADINĂ SANDOZ 5 mg

comprimate filmate 4659/2012/46 RO -

S.C. SANDOZ S.R.L.

DESLORATADINĂ SANDOZ 5 mg

comprimate filmate 4659/2012/47 RO -

S.C. SANDOZ S.R.L.

DESLORATADINĂ SANDOZ 5 mg

comprimate filmate 4659/2012/48 RO -

S.C. SANDOZ S.R.L.

DESLORATADINĂ SANDOZ 5 mg

comprimate filmate 4659/2012/49 RO -

S.C. SANDOZ S.R.L.

DESLORATADINĂ SANDOZ 5 mg

comprimate filmate 4659/2012/50 RO -

S.C. SANDOZ S.R.L.

DESLORATADINĂ SANDOZ 5 mg

comprimate filmate 4659/2012/51 RO -

S.C. SANDOZ S.R.L.

DESLORATADINĂ SANDOZ 5 mg

comprimate filmate 4659/2012/52 RO -

S.C. SANDOZ S.R.L.

DESLORATADINĂ SANDOZ 5 mg

comprimate filmate 4659/2012/53 RO -

S.C. SANDOZ S.R.L.

DESLORATADINĂ SANDOZ 5 mg

comprimate filmate 4659/2012/54 RO -

S.C. SANDOZ S.R.L.

DESLORATADINĂ SANDOZ 5 mg

comprimate filmate 4659/2012/55 RO -

S.C. SANDOZ S.R.L.

DESLORATADINĂ SANDOZ 5 mg

comprimate filmate 4659/2012/56 RO -

S.C. SANDOZ S.R.L.

Desloratadină Terapia 5 mg comprimate
filmate 4481/2012/01 RO - TERAPIA S.A.

Desloratadină Terapia 5 mg comprimate
filmate 4481/2012/02 RO - TERAPIA S.A.

Desloratadină Terapia 5 mg comprimate
filmate 4481/2012/03 RO - TERAPIA S.A.

Desloratadină Terapia 5 mg comprimate
filmate 4481/2012/04 RO - TERAPIA S.A.

Desloratadină Terapia 5 mg comprimate
filmate 4481/2012/05 RO - TERAPIA S.A.

Desloratadină Terapia 5 mg comprimate
filmate 4481/2012/06 RO - TERAPIA S.A.

Desloratadină Terapia 5 mg comprimate
filmate 4481/2012/07 RO - TERAPIA S.A.

Desloratadină Terapia 5 mg comprimate
filmate 4481/2012/08 RO - TERAPIA S.A.

Desloratadină Terapia 5 mg comprimate

filmate 4481/2012/09 RO - TERAPIA S.A.

Desloratadină Terapia 5 mg comprimate

filmate 4481/2012/10 RO - TERAPIA S.A.

Desloratadină Teva 5 mg comprimate

orodispersabile 4575/2012/01 RO - TEVA

PHARMACEUTICALS S.R.L

Desloratadină Teva 5 mg comprimate

orodispersabile 4575/2012/02 RO - TEVA

PHARMACEUTICALS S.R.L

Desloratadină Teva 5 mg comprimate

orodispersabile 4575/2012/03 RO - TEVA

PHARMACEUTICALS S.R.L

Desloratadină Teva 5 mg comprimate

orodispersabile 4575/2012/04 RO - TEVA

PHARMACEUTICALS S.R.L

Desloratadină Teva 5 mg comprimate

orodispersabile 4575/2012/05 RO - TEVA

PHARMACEUTICALS S.R.L

Desloratadină Teva 5 mg comprimate

orodispersabile 4575/2012/06 RO - TEVA

PHARMACEUTICALS S.R.L

Desloratadină Teva 5 mg comprimate

orodispersabile 4575/2012/07 RO - TEVA

PHARMACEUTICALS S.R.L

Desloratadină Teva 5 mg comprimate

orodispersabile 4575/2012/08 RO - TEVA

PHARMACEUTICALS S.R.L

Desloratadină Teva 5 mg comprimate

orodispersabile 4575/2012/09 RO - TEVA

PHARMACEUTICALS S.R.L

Desloratadine +pharma, 5 mg, tabletki

powlekane 20082 PL - +PHARMA

ARZNEIMITTEL GMBH

Desloratadine 0.5 mg/ml oral solution PL

17907/0502 UK - BRISTOL LABORATORIES

LTD (BERKHAMSTED)

Desloratadine 0.5 mg/ml Oral Solution PL

20416/0242 UK - CRESCENT PHARMA LIMITED

Desloratadine 0.5mg/ml Oral Solution

PL24668/0162 UK - CADUCEUS PHARMA

LIMITED

Desloratadine 1A Farma 48333 DK - 1A

FARMA A/S

Desloratadine 1A Farma 48334 DK - 1A

FARMA A/S

Desloratadine 2.5 mg orodispersible tablets

PL 17907/0499 UK - BRISTOL LABORATORIES

LTD (BERKHAMSTED)

Desloratadine 5 mg film-coated tablets PL

17907/0501 UK - BRISTOL LABORATORIES

LTD (BERKHAMSTED)

Desloratadine 5 mg Film-Coated Tablets PL
20416/0241 UK - CRESCENT PHARMA LIMITED
Desloratadine 5 mg orodispersible tablets
PL 17907/0500 UK - BRISTOL LABORATORIES
LTD (BERKHAMSTED)
Desloratadine 5 mg tablets PL 25258/0052
UK - GLENMARK PHARMACEUTICALS EUROPE
LIMITED
Desloratadine 5mg Film-coated Tablets PL
35507/0035 UK - LUPIN (EUROPE) LIMITED
DESLORATADINE ACTAVIS 0,5 MG/ML
ORAALILIUOS 30005 FI - ACTAVIS GROUP
PTC EHF.
DESLORATADINE ACTAVIS, 0,5 MG/ML,
ROZTWÓR DOUSTNY 20518 PL - ACTAVIS
GROUP PTC EHF.
DESLORATADINE ALMUS 5 mg, comprimé
pelliculé 269 410-7 FR - ALMUS FRANCE
DESLORATADINE ALMUS 5 mg, comprimé
pelliculé 269 411-3 FR - ALMUS FRANCE
DESLORATADINE ALMUS 5 mg, comprimé
pelliculé 269 413-6 FR - ALMUS FRANCE
DESLORATADINE ALMUS 5 mg, comprimé
pelliculé 269 414-2 FR - ALMUS FRANCE
DESLORATADINE ALMUS 5 mg, comprimé
pelliculé 584 394-4 FR - ALMUS FRANCE
DESLORATADINE ALMUS 5 mg, comprimé
pelliculé 584 395-0 FR - ALMUS FRANCE
DESLORATADINE ALMUS 5 mg, comprimé
pelliculé 584 396-7 FR - ALMUS FRANCE
Desloratadine Apotex 0.5 mg/ml oral
solution 48309 DK - APOTEX EUROPE B.V.
Desloratadine Apotex 5 mg comprimidos
revestidos por película NOT YET KNOWN PT
- APOTEX EUROPE B.V.
Desloratadine Apotex 5 mg filmomhulde
tabletten RVG 110966 NL - APOTEX EUROPE
BV
DESLORATADINE ARROW 0,5 mg/ml,
solution buvable 50638 FR - ARROW
GENERIQUES
DESLORATADINE ARROW 5mg, comprimé
pelliculé 50454 FR - ARROW GENERIQUES
DESLORATADINE BIOGARAN 0,5 mg/ml,
solution buvable 3400941963070 FR -
BIOGARAN
DESLORATADINE BIOGARAN 0,5 mg/ml,
solution buvable 3400941963131 FR -
BIOGARAN
DESLORATADINE BIOGARAN 0,5 mg/ml,
solution buvable 3400941963360 FR -

BIOGARAN

**DESLORATADINE BIOGARAN 0,5 mg/ml,
solution buvable 3400941963421 FR -**

BIOGARAN

**DESLORATADINE BIOGARAN 0,5 mg/ml,
solution buvable 3400941963599 FR -**

BIOGARAN

**DESLORATADINE BIOGARAN 0,5 mg/ml,
solution buvable 3400941963650 FR -**

BIOGARAN

**DESLORATADINE BIOGARAN 0,5 mg/ml,
solution buvable 3400941963711 FR -**

BIOGARAN

**DESLORATADINE BIOGARAN 0,5 mg/ml,
solution buvable 3400941963889 FR -**

BIOGARAN

**DESLORATADINE BIOGARAN 0,5 mg/ml,
solution buvable 3400941963940 FR -**

BIOGARAN

**DESLORATADINE BIOGARAN® 5 mg,
comprimé pelliculé 3400930021149 FR -**

BIOGARAN

**DESLORATADINE BIOGARAN® 5 mg,
comprimé pelliculé 3400941999031 FR -**

BIOGARAN

**DESLORATADINE BIOGARAN® 5 mg,
comprimé pelliculé 3400941999499 FR -**

BIOGARAN

**Desloratadine CF 0,5 mg/ml, drank RVG
109205 NL - CENTRAFARM B.V.**

**Desloratadine CF 5 mg, filmomhulde
tabletten RVG 109204 NL - CENTRAFARM B.V.
Desloratadine Cipla 5 mg film-coated
tablets MA1059/00401 MT - CIPLA EUROPE
NV**

**Desloratadine Cipla 5 mg film-coated
tablets PA1963/006/001 IE - CIPLA EUROPE
NV**

**Desloratadine Cipla 5 mg film-coated
tablets PL 43362/0015 UK - CIPLA EUROPE
NV**

**Desloratadine Cipla 5 mg filmdragerade
tabletter 45973 SE - CIPLA (UK) LIMITED**

**Desloratadine Cipla 5 mg filmomhulde
tabletten BE489253 BE - CIPLA EUROPE NV**

**Desloratadine Cipla 5 mg filmtabletta OGYI-
T-22321/01 HU - CIPLA (UK) LIMITED**

Desloratadine Cipla 5 mg tabletter,

filmdrasjerte 15-10584 NO - CIPLA EUROPE

NV

**Desloratadine Clomel 0.5 mg/ml oral
solution PA0126/241/001 IE - CLONMEL**

HEALTHCARE LTD.

**DESLORATADINE CRISTERS 5 mg,
comprimé pelliculé 34009 419 658 9 0 FR -
CRISTERS**

**DESLORATADINE CRISTERS 5 mg,
comprimé pelliculé 34009 419 659 5 1 FR -
CRISTERS**

**DESLORATADINE CRISTERS 5 mg,
comprimé pelliculé 34009 419 662 6 2 FR -
CRISTERS**

**DESLORATADINE CRISTERS 5 mg,
comprimé pelliculé 34009 419 663 2 3 FR -
CRISTERS**

**DESLORATADINE CRISTERS 5 mg,
comprimé pelliculé 34009 419 665 5 2 FR -
CRISTERS**

**DESLORATADINE CRISTERS 5 mg,
comprimé pelliculé 34009 419 666 1 3 FR -
CRISTERS**

**DESLORATADINE CRISTERS 5 mg,
comprimé pelliculé 34009 419 668 4 2 FR -
CRISTERS**

**DESLORATADINE CRISTERS 5 mg,
comprimé pelliculé 34009 419 669 0 3 FR -
CRISTERS**

**DESLORATADINE CRISTERS 5 mg,
comprimé pelliculé 34009 419 671 5 3 FR -
CRISTERS**

**DESLORATADINE CRISTERS 5 mg,
comprimé pelliculé 34009 419 672 1 4 FR -
CRISTERS**

**DESLORATADINE CRISTERS 5 mg,
comprimé pelliculé 34009 419 674 4 3 FR -
CRISTERS**

**DESLORATADINE CRISTERS 5 mg,
comprimé pelliculé 34009 419 680 4 4 FR -
CRISTERS**

**DESLORATADINE CRISTERS 5 mg,
comprimé pelliculé 34009 419 681 0 5 FR -
CRISTERS**

**DESLORATADINE CRISTERS 5 mg,
comprimé pelliculé 34009 419 682 7 3 FR -
CRISTERS**

**DESLORATADINE CRISTERS 5 mg,
comprimé pelliculé 34009 419 683 3 4 FR -
CRISTERS**

DESLORATADINE CRISTERS 5 mg,

comprimé pelliculé 34009 419 685 6 3 FR -

CRISTERS

DESLORATADINE CRISTERS 5 mg,

comprimé pelliculé 34009 419 686 2 4 FR -

CRISTERS

DESLORATADINE CRISTERS 5 mg,

comprimé pelliculé 34009 419 687 9 2 FR -

CRISTERS

DESLORATADINE CRISTERS 5 mg,

comprimé pelliculé 34009 419 688 5 3 FR -

CRISTERS

DESLORATADINE CRISTERS 5 mg,

comprimé pelliculé 34009 419 689 1 4 FR -

CRISTERS

DESLORATADINE CRISTERS 5 mg,

comprimé pelliculé 34009 419 691 6 4 FR -

CRISTERS

DESLORATADINE CRISTERS 5 mg,

comprimé pelliculé 34009 419 692 2 5 FR -

CRISTERS

DESLORATADINE CRISTERS 5 mg,

comprimé pelliculé 34009 580 449 9 1 FR -

CRISTERS

DESLORATADINE CRISTERS 5 mg,

comprimé pelliculé 34009 580 451 3 4 FR -

CRISTERS

DESLORATADINE CRISTERS 5 mg,

comprimé pelliculé 34009 580 453 6 3 FR -

CRISTERS

DESLORATADINE CRISTERS 5 mg,

comprimé pelliculé 34009 580 457 1 4 FR -

CRISTERS

DESLORATADINE CRISTERS 5 mg,

comprimé pelliculé 34009 580 458 8 2 FR -

CRISTERS

DESLORATADINE CRISTERS 5 mg,

comprimé pelliculé 34009 580 459 4 3 FR -

CRISTERS

Desloratadine Dr. Max, 5 mg, tabletki

powlekane 20897 PL - DR. MAX

Desloratadine EG 0,5 mg/ml drank

BE409735 BE - EUROGENERICSA

Desloratadine EG 0,5 mg/ml Lösung zum

Einnehmen BE409735 BE - EUROGENERICSA

SA

Desloratadine EG 0,5 mg/ml solution

buvable 2013120609 LU - EUROGENERICSA

Desloratadine EG 0,5 mg/ml solution

buvable BE409735 BE - EUROGENERICSA

DESLORATADINE EG 0,5 mg/ml, solution

buvable NL50636 FR - EG LABO -

LABORATOIRES EUROGENERICs

Desloratadine EG 5 mg comprimés

pelliculés 2013120608 LU - EUROGENERICs

SA

Desloratadine EG 5 mg comprimés

pelliculés BE412921 BE - EUROGENERICs SA

Desloratadine EG 5 mg filmomhulde

tabletten BE412921 BE - EUROGENERICs SA

Desloratadine EG 5 mg FilmtabLetten

BE412921 BE - EUROGENERICs SA

DESLORATADINE EG 5 mg, comprimé

pelliculé NL40899 FR - EG LABO -

LABORATOIRES EUROGENERICs

Desloratadine ESP Pharma 0,5 mg/ml oral

solution 20130245 BG - ACTAVIS GROUP PTC

EHF.

DESLORATADINE EVOLUGEN 5 mg,

comprimé pelliculé 34009 279 101 7 3 FR -

EVOLUPHARM

DESLORATADINE EVOLUGEN 5 mg,

comprimé pelliculé 34009 279 102 3 4 FR -

EVOLUPHARM

DESLORATADINE EVOLUGEN 5 mg,

comprimé pelliculé 34009 279 104 6 3 FR -

EVOLUPHARM

DESLORATADINE EVOLUGEN 5 mg,

comprimé pelliculé 34009 279 105 2 4 FR -

EVOLUPHARM

DESLORATADINE EVOLUGEN 5 mg,

comprimé pelliculé 34009 279 106 9 2 FR -

EVOLUPHARM

DESLORATADINE EVOLUGEN 5 mg,

comprimé pelliculé 34009 279 107 5 3 FR -

EVOLUPHARM

DESLORATADINE EVOLUGEN 5 mg,

comprimé pelliculé 34009 279 108 1 4 FR -

EVOLUPHARM

DESLORATADINE EVOLUGEN 5 mg,

comprimé pelliculé 34009 586 864 8 1 FR -

EVOLUPHARM

DESLORATADINE EVOLUGEN 5 mg,

comprimé pelliculé 34009 586 865 4 2 FR -

EVOLUPHARM

DESLORATADINE EVOLUGEN 5 mg,

comprimé pelliculé 34009 586 866 0 3 FR -

EVOLUPHARM

DESLORATADINE EVOLUGEN 5 mg,

comprimé pelliculé 34009 586 867 7 1 FR -

EVOLUPHARM

DESLORATADINE EVOLUGEN 5 mg,

comprimé pelliculé 34009 586 868 3 2 FR -

EVOLUPHARM

DESLORATADINE EVOLUGEN 5 mg,
comprimé pelliculé 34009 586 870 8 2 FR -

EVOLUPHARM

DESLORATADINE EVOLUGEN 5 mg,
comprimé pelliculé 34009279 109 8 2 FR -

EVOLUPHARM

Desloratadine Gedeon Richter 5 mg film-
coated tablets PL 04854-0125 UK - GEDEON
RICHTER PLC.

Desloratadine Gedeon Richter 5 mg
filmtabletta OGYI-T-22039/01 HU - GEDEON
RICHTER PLC.

Desloratadine Gedeon Richter 5 mg
filmtabletta OGYI-T-22039/02 HU - GEDEON
RICHTER PLC.

Desloratadine Gedeon Richter 5 mg
filmtabletta OGYI-T-22039/03 HU - GEDEON
RICHTER PLC.

Desloratadine Gedeon Richter 5 mg
filmtabletta OGYI-T-22039/04 HU - GEDEON
RICHTER PLC.

Desloratadine Gedeon Richter 5 mg
filmtabletta OGYI-T-22039/05 HU - GEDEON
RICHTER PLC.

Desloratadine Gedeon Richter 5 mg
filmtabletta OGYI-T-22039/06 HU - GEDEON
RICHTER PLC.

Desloratadine Gedeon Richter 5 mg
filmtabletta OGYI-T-22039/07 HU - GEDEON
RICHTER PLC.

Desloratadine Genoptim, 0,5 mg/ml,
roztwór doustny 20574 PL - SYOPTIS
PHARMA SP Z O O

Desloratadine Genoptim, 5 mg, tabletki
powlekane 20573 PL - SYOPTIS PHARMA SP
Z O O

Desloratadine Glenmark 5 mg tabletit
29566 FI - GLENMARK PHARMACEUTICALS
EUROPE LIMITED

Desloratadine Glenmark 5 mg tablets
PA1462/007/001 IE - GLENMARK
PHARMACEUTICALS EUROPE LIMITED

Desloratadine Glenmark 5 mg tabletter
29566 FI - GLENMARK PHARMACEUTICALS
EUROPE LIMITED

Desloratadine Glenmark 5mg tabletten RVG
109194 NL - GLENMARK PHARMACEUTICALS
EUROPE LIMITED

Desloratadine Glenmark 5mg tabletter
45749 SE - GLENMARK PHARMACEUTICALS

EUROPE LIMITED

**DESLORATADINE MYLAN 0,5 mg/ml,
solution buvable NL 50409 FR - MYLAN S.A.S**
**Desloratadine Mylan 5 mg film-coated
tablets PL 04569/1302 UK - GENERICS [UK]**

LIMITED

DESLORATADINE MYLAN 5 mg, comprimé

pelliculé NL 41140 FR - MYLAN S.A.S

**Desloratadine Mylan 5 mg, filmomhulde
tabletten RVG 109515 NL - MYLAN B.V.**

DESLORATADINE MYLAN GENERIQUES 5

mg, comprimé pelliculé NL 50494 FR -

MYLAN S.A.S

**DESLORATADINE MYLAN PHARMA 5 mg,
comprimé pelliculé NL50396 FR - MYLAN**

S.A.S

**Desloratadine Mylan, 0,5 mg/ml, roztwór
doustny 20785 PL - MYLAN S.A.S**

**Desloratadine Mylan, 5 mg, tabletki
powlekane 20846 PL - MYLAN S.A.S**

**Desloratadine Peseri, 0,5 mg/ml, roztwór
doustny 20479 PL - BIO PROFIL POLSKA SP Z
OO**

**Desloratadine Peseri, 2,5 mg, tabletki
u1egajce rozpadowi wjamie ustnej 20491
PL - PESERI TRADING LIMITED**

**Desloratadine Peseri, 5 mg, tabletki
ulegajace rozpadowi wjamie ustnej 20492
PL - PESERI TRADING LIMITED**

**DESLORATADINE RANBAXY 5 mg,
comprimé pelliculé NL 40918 FR - RANBAXY
PHARMACIE GENERIQUES**

**Desloratadine ratiopharm 0,5 mg/ml
oraaliliuos 30284 FI - RATIOPHARM GMBH**

**Desloratadine ratiopharm 0,5 mg/ml oral
lösning 30284 FI - RATIOPHARM GMBH**

**Desloratadine ratiopharm 2,5 mg
munsönderfallande tabletter 29581 FI -
RATIOPHARM GMBH**

**Desloratadine ratiopharm 2,5 mg
smeltetabletter 11-8180 NO - RATIOPHARM
GMBH**

**Desloratadine ratiopharm 2,5 mg tabletter,
suussa hajoava 29581 FI - RATIOPHARM
GMBH**

**Desloratadine ratiopharm 5 mg
munndreifitöflur IS/1/13/079/01 IS -
RATIOPHARM GMBH**

**Desloratadine ratiopharm 5 mg
munsönderfallande tabletter 29582 FI -
RATIOPHARM GMBH**

Desloratadine ratiopharm 5 mg
smeltetabletter 11-8181 NO - RATIOPHARM
GMBH

Desloratadine ratiopharm 5 mg tabletti,
suussa hajoava 29582 FI - RATIOPHARM
GMBH

Desloratadine ratiopharm, 0,5 mg/ml,
roztwór doustny 21410 PL - RATIOPHARM
GMBH

Desloratadine Sandoz 0,5 mg/ml, drank
RVG 109202 NL - SANDOZ B.V.

DESLORATADINE SANDOZ 0,5 mg/ml,
solution buvable 273 167-6 FR - SANDOZ

DESLORATADINE SANDOZ 0,5 mg/ml,
solution buvable 273 168-2 FR - SANDOZ

DESLORATADINE SANDOZ 0,5 mg/ml,
solution buvable 273 169-9 FR - SANDOZ

DESLORATADINE SANDOZ 0,5 mg/ml,
solution buvable 273 170-7 FR - SANDOZ

DESLORATADINE SANDOZ 0,5 mg/ml,
solution buvable 273 171-3 FR - SANDOZ

DESLORATADINE SANDOZ 0,5 mg/ml,
solution buvable 273 173-6 FR - SANDOZ

DESLORATADINE SANDOZ 0,5 mg/ml,
solution buvable 273 174-2 FR - SANDOZ

DESLORATADINE SANDOZ 0,5 mg/ml,
solution buvable 273 175-9 FR - SANDOZ

DESLORATADINE SANDOZ 0,5 mg/ml,
solution buvable 273 176-5 FR - SANDOZ

DESLORATADINE SANDOZ 0,5 mg/ml,
solution buvable 273 177-1 FR - SANDOZ

Desloratadine Sandoz 022141 CY - SANDOZ
GMBH

Desloratadine Sandoz 022142 CY - SANDOZ
GMBH

Desloratadine Sandoz 48329 DK - SANDOZ
A/S

Desloratadine Sandoz 48330 DK - SANDOZ
A/S

DESLORATADINE SANDOZ 5 mg comprimé
pelliculé 275 897-1 FR - SANDOZ

DESLORATADINE SANDOZ 5 mg comprimé
pelliculé 275 898-8 FR - SANDOZ

DESLORATADINE SANDOZ 5 mg comprimé
pelliculé 275 899-4 FR - SANDOZ

DESLORATADINE SANDOZ 5 mg comprimé
pelliculé 275 900-2 FR - SANDOZ

Desloratadine Sandoz 5 mg filmdragerade
tablettter 45779 SE - SANDOZ A/S

Desloratadine Sandoz 5 mg filmuhúðaðar
töflur IS/1/12/053/01 IS - SANDOZ A/S

Desloratadine Sandoz 5 mg
kalvopäällysteinen tabletti 29573 FI -
SANDOZ A/S
Desloratadine Sandoz 5 mg, filmomhulde
tabletten RVG 109201 NL - SANDOZ B.V.
Desloratadine Smeltablet 2,5 mg Teva,
orodispergeerbare tabletten RVG 109206
NL - TEVA NEDERLAND B.V.
Desloratadine Smeltablet 5 mg Teva,
orodispergeerbare tabletten RVG 109207
BE - TEVA NEDERLAND B.V.
Desloratadine Sopharma 0,5 mg/ml
geriamasis tirpalas LT/1/15/3736/001 LT -
SOPHARMA AD
Desloratadine Sopharma 0,5 mg/ml
oraaliliuos 32430 FI - SOPHARMA AD
Desloratadine Sopharma 0,5 mg/ml oral
lösning 51396 SE - SOPHARMA AD
Desloratadine Sopharma 0,5 mg/ml
roztwór doustny 22747 PL - SOPHARMA AD
Desloratadine Sopharma 0,5 mg/ml
suukaudne lahus 874615 EE - SOPHARMA AD
Desloratadine Sopharma 0,5 mg/ml
šķidums iekšķīgai lietošanai 15-0132 LV -
SOPHARMA AD
Desloratadine Sopharma 5 mg apvalkotās
tabletes 16-0008 LV - SOPHARMA AD
Desloratadine Sopharma 5 mg apvalkotās
tabletes 16-0008 LV - SOPHARMA AD
Desloratadine Sopharma 5 mg film-coated
tablets 33349 FI - SOPHARMA AD
Desloratadine Sopharma 5 mg film-coated
tablets 33349 FI - SOPHARMA AD
Desloratadine Sopharma 5 mg
filmdragerade tabletter 52877 SE -
SOPHARMA AD
Desloratadine Sopharma 5 mg
filmdragerade tabletter 52877 SE -
SOPHARMA AD
Desloratadine Sopharma 5 mg öhukese
polümeerikattega tabletid 901016 EE -
SOPHARMA AD
Desloratadine Sopharma 5 mg öhukese
polümeerikattega tabletid 901016 EE -
SOPHARMA AD
Desloratadine Sopharma 5 mg plevele
dengtos tablettes LT/1/16/3873/002 LT -
SOPHARMA AD
Desloratadine Sopharma 5 mg plevele
dengtos tablettes LT/1/16/3873/002 LT -
SOPHARMA AD

Desloratadine Sopharma 5 mg plevele
dengtos tabletės LT/1/16/3873/003 LT -
SOPHARMA AD

Desloratadine Sopharma 5 mg plevele
dengtos tabletės LT/1/16/3873/003 LT -
SOPHARMA AD

Desloratadine Sopharma 5 mg plevele
dengtos tabletės LT/1/16/3873/004 LT -
SOPHARMA AD

Desloratadine Sopharma 5 mg plevele
dengtos tabletės LT/1/16/3873/004 LT -
SOPHARMA AD

Desloratadine Sopharma 5 mg plevele
dengtos tabletės LT/1/16/3873/005 LT -
SOPHARMA AD

Desloratadine Sopharma 5 mg plevele
dengtos tabletės LT/1/16/3873/005 LT -
SOPHARMA AD

Desloratadine Sopharma 5 mg plevele
dengtos tabletės LT/1/16/3873/006 LT -
SOPHARMA AD

Desloratadine Sopharma 5 mg plevele
dengtos tabletės LT/1/16/3873/006 LT -
SOPHARMA AD

Desloratadine Sopharma 5 mg plevele
dengtos tabletės LT/1/16/3873/007 LT -
SOPHARMA AD

Desloratadine Sopharma 5 mg plėvele
dengtos tabletės LT/1/16/3873/001 LT -
SOPHARMA AD

Desloratadine Sopharma 5 mg plėvele
dengtos tabletės LT/1/16/3873/001 LT -
SOPHARMA AD

Desloratadine Sopharma, 5 mg, tabletki
powlekane 23119 PL - SOPHARMA AD

Desloratadine Sopharma, 5 mg, tabletki
powlekane 23119 PL - SOPHARMA AD

Desloratadine Stada 0,5 mg/ml belsőleges
oldat OGYI-T-22041/13 HU - STADA
ARZNEIMITTEL AG

Desloratadine Stada 0,5 mg/ml belsőleges
oldat OGYI-T-22041/14 HU - STADA
ARZNEIMITTEL AG

Desloratadine Stada 0,5 mg/ml belsőleges
oldat OGYI-T-22041/15 HU - STADA
ARZNEIMITTEL AG

Desloratadine Stada 0,5 mg/ml belsőleges
oldat OGYI-T-22041/16 HU - STADA

ARZNEIMITTEL AG

Desloratadine Stada 0,5 mg/ml belsőleges

oldat OGYI-T-22041/17 HU - STADA

ARZNEIMITTEL AG

Desloratadine Stada 0,5 mg/ml belsőleges

oldat OGYI-T-22041/18 HU - STADA

ARZNEIMITTEL AG

Desloratadine Stada 5 mg filmtabletta

OGYI-T-22041/01 HU - STADA ARZNEIMITTEL

AG

Desloratadine Stada 5 mg filmtabletta

OGYI-T-22041/02 HU - STADA ARZNEIMITTEL

AG

Desloratadine Stada 5 mg filmtabletta

OGYI-T-22041/03 HU - STADA ARZNEIMITTEL

AG

Desloratadine Stada 5 mg filmtabletta

OGYI-T-22041/04 HU - STADA ARZNEIMITTEL

AG

Desloratadine Stada 5 mg filmtabletta

OGYI-T-22041/05 HU - STADA ARZNEIMITTEL

AG

Desloratadine Stada 5 mg filmtabletta

OGYI-T-22041/06 HU - STADA ARZNEIMITTEL

AG

Desloratadine Stada 5 mg filmtabletta

OGYI-T-22041/07 HU - STADA ARZNEIMITTEL

AG

Desloratadine Stada 5 mg filmtabletta

OGYI-T-22041/08 HU - STADA ARZNEIMITTEL

AG

Desloratadine Stada 5 mg filmtabletta

OGYI-T-22041/09 HU - STADA ARZNEIMITTEL

AG

Desloratadine Stada 5 mg filmtabletta

OGYI-T-22041/10 HU - STADA ARZNEIMITTEL

AG

Desloratadine Stada 5 mg filmtabletta

OGYI-T-22041/11 HU - STADA ARZNEIMITTEL

AG

Desloratadine Stada 5 mg filmtabletta

OGYI-T-22041/12 HU - STADA ARZNEIMITTEL

AG

Desloratadine Teva 0,5 mg/ml drank

BE437342 BE - TEVA PHARMA BELGIUM

N.V./S.A

Desloratadine Teva 0,5 mg/ml perorální

roztok 24/250/13-C CZ - TEVA

PHARMACEUTICALS CR, S.R.O.

Desloratadine Teva 0,5 mg/ml, drank RVG

110902 NL - TEVA NEDERLAND B.V.

Desloratadine Teva 0,5 mg/ml, drank RVG

110903 NL - TEVA NEDERLAND B.V.

Desloratadine Teva 0,5 mg/ml, solution

buvable BE437342 BE - TEVA PHARMA

BELGIUM N.V./S.A

DESLORATADINE TEVA 0,5mg/ml LÖSUNG

ZUM EINNEHMEN BE437342 BE - TEVA

PHARMA BELGIUM N.V./S.A

Desloratadine Teva 0,5 mg/ml perorale

raztvor 20130221 BG - TEVA

PHARMACEUTICALS BULGARIA EOOD

Desloratadine Teva 2,5 mg comprimés

orodispersibles BE419596 BE - TEVA

PHARMA BELGIUM N.V./S.A

Desloratadine Teva 2,5 mg

Schmelztabletten BE419596 BE - TEVA

PHARMA BELGIUM N.V./S.A

Desloratadine Teva 5 mg orodispergeerbare

tabletten BE419605 BE - TEVA PHARMA

BELGIUM N.V./S.A

Desloratadine Teva 5 mg Schmelztabletten

BE419605 BE - TEVA PHARMA BELGIUM

N.V./S.A

Desloratadine Teva Oral Solution 0.5

mg/ml PL 00289/1699 UK - TEVA UK

LIMITED

Desloratadine Teva Pharma B.V. 0,5 mg/ml

posimo dialyma 21782 CY - TEVA PHARMA

B.V.

Desloratadine Teva Pharma B.V. 0,5 mg/ml

posimo dialyma 49652/13-6-2014 GR -

TEVA PHARMA B.V.

DESLORATADINE TEVA SANTE 5 mg,

comprimé pelliculé NL50684 FR - TEVA

SANTÉ

DESLORATADINE URGO 5 mg, comprimé

pelliculé 34009 273 263 5 6 FR -

LABORATOIRES URGO

DESLORATADINE URGO 5 mg, comprimé

pelliculé 34009 273 264 1 7 FR -

LABORATOIRES URGO

DESLORATADINE URGO 5 mg, comprimé

pelliculé 34009 273 265 8 5 FR -

LABORATOIRES URGO

DESLORATADINE URGO 5 mg, comprimé

pelliculé 34009 273 266 4 6 FR -

LABORATOIRES URGO

DESLORATADINE URGO 5 mg, comprimé

pelliculé 34009 273 267 0 7 FR -

LABORATOIRES URGO

DESLORATADINE URGO 5 mg, comprimé

pelliculé 34009 273 268 7 5 FR -
LABORATOIRES URG
DESLORATADINE URG 5 mg, comprimé
pelliculé 34009 273 269 3 6 FR -
LABORATOIRES URG
DESLORATADINE URG 5 mg, comprimé
pelliculé 34009 273 270 1 8 FR -
LABORATOIRES URG
DESLORATADINE URG 5 mg, comprimé
pelliculé 34009 273 271 8 6 FR -
LABORATOIRES URG
DESLORATADINE URG 5 mg, comprimé
pelliculé 34009 273 272 4 7 FR -
LABORATOIRES URG
DESLORATADINE URG 5 mg, comprimé
pelliculé 34009 273 273 0 8 FR -
LABORATOIRES URG
DESLORATADINE URG 5 mg, comprimé
pelliculé 34009 584 801 9 5 FR -
LABORATOIRES URG
DESLORATADINE URG 5 mg, comprimé
pelliculé 34009 584 802 5 6 FR -
LABORATOIRES URG
DESLORATADINE URG 5 mg, comprimé
pelliculé 34009 584 803 1 7 FR -
LABORATOIRES URG
Desloratadine Zentiva 0,5 mg/ml
geriamasis tirpalas LT/1/12/2847/001 LT -
ZENTIVA, K.S.
Desloratadine Zentiva 0,5 mg/ml
geriamasis tirpalas LT/1/12/2847/002 LT -
ZENTIVA, K.S.
Desloratadine Zentiva 0,5 mg/ml
geriamasis tirpalas LT/1/12/2847/003 LT -
ZENTIVA, K.S.
Desloratadine Zentiva 0,5 mg/ml
geriamasis tirpalas LT/1/12/2847/004 LT -
ZENTIVA, K.S.
Desloratadine Zentiva 0,5 mg/ml
geriamasis tirpalas LT/1/12/2847/005 LT -
ZENTIVA, K.S.
DESLORATADINE ZENTIVA 0,5 mg/ml,
solution buvable 34009 217 616 3 4 FR -
SANOFI-AVENTIS FRANCE
DESLORATADINE ZENTIVA 0,5 mg/ml,
solution buvable 34009 217 618 6 3 FR -
SANOFI-AVENTIS FRANCE
DESLORATADINE ZENTIVA 0,5 mg/ml,
solution buvable 34009 217 619 2 4 FR -
SANOFI-AVENTIS FRANCE
DESLORATADINE ZENTIVA 0,5 mg/ml,

solution buvable 34009 217 620 0 6 FR -
SANOFI-AVENTIS FRANCE
DESLORATADINE ZENTIVA 0,5 mg/ml,
solution buvable 34009 217 621 7 4 FR -
SANOFI-AVENTIS FRANCE
DESLORATADINE ZENTIVA 0,5 mg/ml,
solution buvable 34009 217 622 3 5 FR -
SANOFI-AVENTIS FRANCE
DESLORATADINE ZENTIVA 0,5 mg/ml,
solution buvable 34009 217 624 6 4 FR -
SANOFI-AVENTIS FRANCE
DESLORATADINE ZENTIVA 0,5 mg/ml,
solution buvable 34009 217 625 2 5 FR -
SANOFI-AVENTIS FRANCE
DESLORATADINE ZENTIVA 0,5 mg/ml,
solution buvable 34009 217 626 9 3 FR -
SANOFI-AVENTIS FRANCE
Desloratadine Zentiva 0,5 mg/ml,
suukaudne lahus 774512 EE - ZENTIVA, K.S.
Desloratadine Zentiva 5 mg plèvèle dengtos
tabletès LT/1/12/2847/006 LT - ZENTIVA,
K.S.
Desloratadine Zentiva 5 mg plèvèle dengtos
tabletès LT/1/12/2847/007 LT - ZENTIVA,
K.S.
Desloratadine Zentiva 5 mg plèvèle dengtos
tabletès LT/1/12/2847/008 LT - ZENTIVA,
K.S.
Desloratadine Zentiva 5 mg plèvèle dengtos
tabletès LT/1/12/2847/009 LT - ZENTIVA,
K.S.
Desloratadine Zentiva 5 mg plèvèle dengtos
tabletès LT/1/12/2847/010 LT - ZENTIVA,
K.S.
Desloratadine Zentiva 5 mg plèvèle dengtos
tabletès LT/1/12/2847/011 LT - ZENTIVA,
K.S.
Desloratadine Zentiva 5 mg plèvèle dengtos
tabletès LT/1/12/2847/012 LT - ZENTIVA,
K.S.
Desloratadine Zentiva 5 mg plèvèle dengtos
tabletès LT/1/12/2847/013 LT - ZENTIVA,
K.S.
Desloratadine Zentiva 5 mg plèvèle dengtos
tabletès LT/1/12/2847/014 LT - ZENTIVA,
K.S.
Desloratadine Zentiva 5 mg plèvèle dengtos
tabletès LT/1/12/2847/015 LT - ZENTIVA,

K.S.

**DESLORATADINE ZENTIVA 5 MG,
COMPRIME PELLICULE 34009 217 480 4 8**
FR - SANOFI-AVENTIS FRANCE
**DESLORATADINE ZENTIVA 5 MG,
COMPRIME PELLICULE 34009 217 481 0 9**
FR - SANOFI-AVENTIS FRANCE
**DESLORATADINE ZENTIVA 5 MG,
COMPRIME PELLICULE 34009 217 482 7 7**
FR - SANOFI-AVENTIS FRANCE
**DESLORATADINE ZENTIVA 5 MG,
COMPRIME PELLICULE 34009 217 486 2 8**
FR - SANOFI-AVENTIS FRANCE
**DESLORATADINE ZENTIVA 5 MG,
COMPRIME PELLICULE 34009 217 487 9 6**
FR - SANOFI-AVENTIS FRANCE
**DESLORATADINE ZENTIVA 5 MG,
COMPRIME PELLICULE 34009 217 488 5 7**
FR - SANOFI-AVENTIS FRANCE
**DESLORATADINE ZENTIVA 5 MG,
COMPRIME PELLICULE 34009 580 922 6 8**
FR - SANOFI-AVENTIS FRANCE
**DESLORATADINE ZENTIVA 5 MG,
COMPRIME PELLICULE 34009 580 923 2 9**
FR - SANOFI-AVENTIS FRANCE
**DESLORATADINE ZENTIVA 5 MG,
COMPRIME PELLICULE 34009 580 924 9 7**
FR - SANOFI-AVENTIS FRANCE
**DESLORATADINE ZENTIVA 5 MG,
COMPRIME PELLICULE 340092 17 48338 FR**
- SANOFI-AVENTIS FRANCE
**DESLORATADINE ZENTIVA 5 MG,
COMPRIME PELLICULE 340092 17 489 1 8**
FR - SANOFI-AVENTIS FRANCE
**DESLORATADINE ZENTIVA 5 MG,
COMPRIME PELLICULE 340092 17 4922 9**
FR - SANOFI-AVENTIS FRANCE
**DESLORATADINE ZENTIVA 5 MG,
COMPRIME PELLICULE 340092 174 8567 FR**
- SANOFI-AVENTIS FRANCE
**DESLORATADINE ZENTIVA 5 MG,
COMPRIME PELLICULE 340092 174 916 8**
FR - SANOFI-AVENTIS FRANCE
**Desloratadine Zentiva 5 mg, õhukese
polümeerikattega tabletid 774612 EE -
ZENTIVA, K.S.**
**DESLORATADINE ZYDUS 5 mg, comprimé
pelliculé 34009 268 610 2 5 FR - ZYDUS
FRANCE**
**DESLORATADINE ZYDUS 5 mg, comprimé
pelliculé 34009 268 611 9 3 FR - ZYDUS**

FRANCE

DESLORATADINE ZYDUS 5 mg, comprimé
pelliculé 34009 268 612 5 4 FR - ZYDUS

FRANCE

DESLORATADINE ZYDUS 5 mg, comprimé
pelliculé 34009 268 613 1 5 FR - ZYDUS

FRANCE

Desloratadine "Hexal" 48331 DK - HEXAL

A/S

Desloratadine "Hexal" 48332 DK - HEXAL

A/S

Desloratadine/Genepharm 0.5 mg/ml

posimo dialyma 31215 GR - GENEPHARM S.A.

Desloratadine/Genepharm 2.5 mg diskia

diaspeirovema sto stoma 31212 GR -

GENEPHARM S.A.

Desloratadine/Genepharm 5 mg diskia

diaspeirovema sto stoma 31214 GR -

GENEPHARM S.A.

Desloratadine/Genepharm 5 mg diskia

epikalimena me lepto ymenio 31213 GR -

GENEPHARM S.A.

Desloratadyna Apotex, 5 mg, tabletki

powlekane 21023 PL - APOTEX EUROPE BV

Desloratatine Teva 2,5 mg

orodispergeerbare tabletten BE419596 BE -

TEVA PHARMA BELGIUM N.V./S.A

Desloratatine Teva 5 mg orodispergeerbare
tabletten BE419605 BE - TEVA PHARMA

BELGIUM N.V./S.A

Deslordinis 5 mg filmom obložene tablete HR-
H-465680296 HR - PLIVA HRVATSKA D.O.O.

Deslorius 0,5 mg/ml posimo dialyma
67125/11-9-13 GR - ALET PHARMACEUTICALS

SA

Deslorius 2,5 mg diskia diaspeirovema sto
stoma 67123/11-9-13 GR - ALET

PHARMACEUTICALS SA

Deslorius 5 mg diskia diaspeirovema sto
stoma 67124/11-9-13 GR - ALET

PHARMACEUTICALS SA

Deslorius 5 mg diskia epikalimena me lepto
ymenio 67122/11-9-13 GR - ALET

PHARMACEUTICALS SA

Dynid 0,5 mg/ml soluție orală
4507/2012/01 RO - GLENMARK

PHARMACEUTICALS S.R.O.

Dynid 0,5 mg/ml soluție orală
4507/2012/02 RO - GLENMARK

PHARMACEUTICALS S.R.O.

Dynid 0,5 mg/ml soluție orală

4507/2012/03 RO - GLENMARK

PHARMACEUTICALS S.R.O.

Dynid 0,5 mg/ml soluție orală

4507/2012/04 RO - GLENMARK

PHARMACEUTICALS S.R.O.

Dynid 0,5 mg/ml soluție orală

4507/2012/05 RO - GLENMARK

PHARMACEUTICALS S.R.O.

Dynid 0,5 mg/ml soluție orală

4507/2012/06 RO - GLENMARK

PHARMACEUTICALS S.R.O.

Dynid 0,5 mg/ml soluție orală

4507/2012/07 RO - GLENMARK

PHARMACEUTICALS S.R.O.

Dynid 0,5 mg/ml soluție orală

4507/2012/08 RO - GLENMARK

PHARMACEUTICALS S.R.O.

Dynid 5 mg comprimate 4684/2012/01 RO

- GLENMARK PHARMACEUTICALS S.R.O.

Dynid 5 mg comprimate 4684/2012/02 RO

- GLENMARK PHARMACEUTICALS S.R.O.

Dynid 5 mg comprimate 4684/2012/03 RO

- GLENMARK PHARMACEUTICALS S.R.O.

Dynid 5 mg comprimate 4684/2012/04 RO

- GLENMARK PHARMACEUTICALS S.R.O.

Dynid 5 mg comprimate 4684/2012/05 RO

- GLENMARK PHARMACEUTICALS S.R.O.

Dynid 5 mg comprimate 4684/2012/06 RO

- GLENMARK PHARMACEUTICALS S.R.O.

Dynid 5 mg comprimate 4684/2012/07 RO

- GLENMARK PHARMACEUTICALS S.R.O.

Dynid 5 mg comprimate 4684/2012/08 RO

- GLENMARK PHARMACEUTICALS S.R.O.

Dynid 5 mg comprimate 4684/2012/09 RO

- GLENMARK PHARMACEUTICALS S.R.O.

Dynid 5 mg comprimate 4684/2012/10 RO

- GLENMARK PHARMACEUTICALS S.R.O.

Dynid 5 mg Tablets PL 33882/0021 UK -

GLENMARK PHARMACEUTICALS S.R.O.

Dynid, 0,5 mg/ml, roztwór doustny 20075

PL - GLENMARK PHARMACEUTICALS S.R.O.

Dynid, 5 mg, tabletki 20191 PL - GLENMARK

PHARMACEUTICALS S.R.O.

EFESTAD 5 mg compresse rivestite con film

040855013 IT - CRINOS S.P.A.

EFESTAD 5 mg compresse rivestite con film

040855025 IT - CRINOS S.P.A.

EFESTAD 5 mg compresse rivestite con film

040855037 IT - CRINOS S.P.A.

EFESTAD 5 mg compresse rivestite con film

040855049 IT - CRINOS S.P.A.

Efestad 5 mg film-coated tablets
PA0126/223/001 IE - CLONMEL HEALTHCARE LTD.

Efestad 5 mg filmirani tabletki 20120172

BG - STADA ARZNEIMITTEL AG

Esradin 5 mg filmom obalené tablety

24/0111/15-S SK - KRKA, D.D., NOVO MESTO

Esradin 5 mg filmsko obložene tablete

5363-I-488/13 SI - KRKA, D.D., NOVO MESTO

Esradin 5 mg filmsko obložene tablete

5363-I-489/13 SI - KRKA, D.D., NOVO MESTO

Esradin 5 mg filmtabletta OGYI-T-

21941/01 HU - KRKA, D.D., NOVO MESTO

Esradin 5 mg filmtabletta OGYI-T-

21941/02 HU - KRKA, D.D., NOVO MESTO

Esradin 5 mg filmtabletta OGYI-T-

21941/03 HU - KRKA, D.D., NOVO MESTO

Esradin 5 mg filmtabletta OGYI-T-

21941/04 HU - KRKA, D.D., NOVO MESTO

Esradin 5 mg filmtabletta OGYI-T-

21941/05 HU - KRKA, D.D., NOVO MESTO

Esradin 5 mg filmtabletta OGYI-T-

21941/06 HU - KRKA, D.D., NOVO MESTO

Flynise 2,5 mg burnoje disperguojamos tabletės LT/1/15/3700/001 LT - ACTAVIS

GROUP PTC EHF.

Flynise 2,5 mg burnoje disperguojamos tabletės LT/1/15/3700/002 LT - ACTAVIS

GROUP PTC EHF.

Flynise 2,5 mg burnoje disperguojamos tabletės LT/1/15/3700/003 LT - ACTAVIS

GROUP PTC EHF.

Flynise 2,5 mg burnoje disperguojamos tabletės LT/1/15/3700/004 LT - ACTAVIS

GROUP PTC EHF.

Flynise 2,5 mg burnoje disperguojamos tabletės LT/1/15/3700/005 LT - ACTAVIS

GROUP PTC EHF.

Flynise 2,5 mg burnoje disperguojamos tabletės LT/1/15/3700/006 LT - ACTAVIS

GROUP PTC EHF.

Flynise 2,5 mg burnoje disperguojamos tabletės LT/1/15/3700/007 LT - ACTAVIS

GROUP PTC EHF.

Flynise 2,5 mg burnoje disperguojamos tabletės LT/1/15/3700/008 LT - ACTAVIS

GROUP PTC EHF.

Flynise 2,5 mg burnoje disperguojamos tabletės LT/1/15/3700/009 LT - ACTAVIS

GROUP PTC EHF.

Flynise 2,5 mg burnoje disperguojamos

Flynise 2,5 mg burnoje disperguojamos

tabletēs LT/1/15/3700/010 LT - ACTAVIS

GROUP PTC EHF.

Flynise 2,5 mg burnoje disperguojamos

tabletēs LT/1/15/3700/011 LT - ACTAVIS

GROUP PTC EHF.

Flynise 2,5 mg burnoje disperguojamos

tabletēs LT/1/15/3700/012 LT - ACTAVIS

GROUP PTC EHF.

Flynise 2,5 mg burnoje disperguojamos

tabletēs LT/1/15/3700/013 LT - ACTAVIS

GROUP PTC EHF.

Flynise 2,5 mg mutē dispergējamās

tabletes 15-0072 LV - ACTAVIS GROUP PTC

EHF.

Flynise 2,5 mg mutē dispergējamās

tabletēs 15-0071 LV - ACTAVIS GROUP PTC

EHF.

Flynise 2,5 mg suus dispergeeruvad

tabletid 867115 EE - ACTAVIS GROUP PTC

EHF.

Flynise 2,5 mg szájban diszpergálódó

tabletta OGYI-T-22796/01-09 HU - ACTAVIS

GROUP PTC EHF.

Flynise 5 mg burnoje disperguojamos

tabletēs LT/1/15/3700/014 LT - ACTAVIS

GROUP PTC EHF.

Flynise 5 mg burnoje disperguojamos

tabletēs LT/1/15/3700/015 LT - ACTAVIS

GROUP PTC EHF.

Flynise 5 mg burnoje disperguojamos

tabletēs LT/1/15/3700/016 LT - ACTAVIS

GROUP PTC EHF.

Flynise 5 mg burnoje disperguojamos

tabletēs LT/1/15/3700/017 LT - ACTAVIS

GROUP PTC EHF.

Flynise 5 mg burnoje disperguojamos

tabletēs LT/1/15/3700/018 LT - ACTAVIS

GROUP PTC EHF.

Flynise 5 mg burnoje disperguojamos

tabletēs LT/1/15/3700/019 LT - ACTAVIS

GROUP PTC EHF.

Flynise 5 mg burnoje disperguojamos

tabletēs LT/1/15/3700/020 LT - ACTAVIS

GROUP PTC EHF.

Flynise 5 mg burnoje disperguojamos

tabletēs LT/1/15/3700/021 LT - ACTAVIS

GROUP PTC EHF.

Flynise 5 mg burnoje disperguojamos

tabletēs LT/1/15/3700/022 LT - ACTAVIS

GROUP PTC EHF.

Flynise 5 mg burnoje disperguojamos

tabletēs LT/1/15/3700/023 LT - ACTAVIS

GROUP PTC EHF.

Flynise 5 mg burnoje disperguojamos

tabletēs LT/1/15/3700/024 LT - ACTAVIS

GROUP PTC EHF.

Flynise 5 mg burnoje disperguojamos

tabletēs LT/1/15/3700/025 LT - ACTAVIS

GROUP PTC EHF.

Flynise 5 mg burnoje disperguojamos

tabletēs LT/1/15/3700/026 LT - ACTAVIS

GROUP PTC EHF.

Flynise 5 mg film-coated tablets

201500363 BG - ACTAVIS GROUP PTC EHF.

Flynise 5 mg filmdragerade tabletter 50926

SE - ACTAVIS GROUP PTC EHF.

Flynise 5 mg filmdrasjerte tabletter 14-

10010 NO - ACTAVIS GROUP PTC EHF.

Flynise 5 mg filmtabletta OGYI-T-22794/01

HU - ACTAVIS GROUP PTC EHF.

Flynise 5 mg filmtabletta OGYI-T-22794/02

HU - ACTAVIS GROUP PTC EHF.

Flynise 5 mg filmtabletta OGYI-T-22794/03

HU - ACTAVIS GROUP PTC EHF.

Flynise 5 mg filmtabletta OGYI-T-22794/04

HU - ACTAVIS GROUP PTC EHF.

Flynise 5 mg filmtabletta OGYI-T-22794/05

HU - ACTAVIS GROUP PTC EHF.

Flynise 5 mg filmtabletta OGYI-T-22794/06

HU - ACTAVIS GROUP PTC EHF.

Flynise 5 mg filmtabletta OGYI-T-22794/07

HU - ACTAVIS GROUP PTC EHF.

Flynise 5 mg filmtabletta OGYI-T-22794/08

HU - ACTAVIS GROUP PTC EHF.

Flynise 5 mg filmtabletta OGYI-T-22794/09

HU - ACTAVIS GROUP PTC EHF.

Flynise 5 mg filmtabletta OGYI-T-22794/10

HU - ACTAVIS GROUP PTC EHF.

Flynise 5 mg filmuhúðaðar töflur

IS/1/15/022/01 IS - ACTAVIS GROUP PTC

EHF.

Flynise 5 mg kalvopäälysteiset tabletit

32142 FI - ACTAVIS GROUP PTC EHF.

Flynise 5 mg muté dispergéjamás tabletes

15-0074 LV - ACTAVIS GROUP PTC EHF.

Flynise 5 mg mutē dispergējamās tabletes

15-0073 LV - ACTAVIS GROUP PTC EHF.

Flynise 5 mg Orodispersible Tablets PA

1380.162.1 IE - ACTAVIS GROUP PTC EHF.

Flynise 5 mg suus dispergeeruvad tabletid

867215 EE - ACTAVIS GROUP PTC EHF.

Flynise 5 mg szájban diszpergálódó tabletta

OGYI-T-22796/10-18 HU - ACTAVIS GROUP

PTC EHF.

Flynise 53724 DK - ACTAVIS GROUP PTC EHF.

Flynise 53725 DK - ACTAVIS GROUP PTC EHF.

Flynise 53752 DK - ACTAVIS GROUP PTC EHF.

Flynise 5mg, tabletki pwolekane 22440 PL -
ACTAVIS GROUP PTC EHF.

Flynise MA 628/12401 MT - ACTAVIS GROUP
PTC EHF.

Flynise MA 628/12402 MT - ACTAVIS GROUP
PTC EHF.

GOLDESIN 20338 PL - TACTICA

PHARMACEUTICAL SP. Z O.O.

GOLDESIN, oral solution, 0,5mg/ml 20339

PL - TACTICA PHARMACEUTICAL SP. Z O.O.

Hitaxa fast, 2,5 mg, tabletki ulegajace

rozpadowi w jamie ustnej 20223 PL -

ADAMED

Hitaxa fast, 5 mg, tabletki ulegajace

rozpadowi w jamie ustnej 20224 PL -

ADAMED

Hitaxa, 0,5 mg/ml, roztwór doustny 20360

PL - ADAMED

Hitaxa, 2,5 mg, tabletki ulegajace

rozpadowi w jamie ustnej 20256 PL -

ADAMED

Hitaxa, 5 mg, tabletki ulegajace rozpadowi

w jamie ustnej 20257 PL - ADAMED

Jovesto 0,5 MG/ML peroralen raztvor

20120370 BG - SANDOZ PHARMACEUTICALS

D.D.

JOVESTO 0,5 MG/ML PERORÁLNÍ ROZTOK

24/411/12-C CZ - SANDOZ S.R.O.

JOVESTO 0,5 mg/ml perorálny roztok

24/0203/12-S SK - SANDOZ

PHARMACEUTICALS D.D.

Jovesto 20358 PL - SANDOZ GMBH

Jovesto 48337 DK - SANDOZ A/S

Jovesto 48338 DK - SANDOZ A/S

Jovesto 5 mg film-coated tablets 20120371

BG - SANDOZ PHARMACEUTICALS D.D.

JOVESTO 5 mg filmom obalené tablety

24/0202/12-S SK - SANDOZ

PHARMACEUTICALS D.D.

JOVESTO 5 MG POTAHOVANÉ TABLETY

24/410/12-C CZ - SANDOZ S.R.O.

JOVESTO, 0,5 MG/ML, ROZTWÓR DOUSTNY

20359 PL - SANDOZ GMBH

Laboratoria PolfaŁódź ALERGO MAX 20478

PL - BIO PROFIL POLSKA SP Z OO

Lentrica 48341 DK - SANDOZ A/S

Lentrica 5 mg filmsko obložene tablete

5363-I-939/12 SI - LEK PHARMACEUTICALS

D.D. LJUBLJANA

Lorabel 0,5 mg/ml peroralna raztopina

5363-I-74/14 SI - BELUPO D.O.O.

Lordestin 0,5 mg/ml belsöleges oldat

OGYI-T-22038/06 HU - GEDEON RICHTER

PLC.

Lordestin 0,5 mg/ml belsöleges oldat

OGYI-T-22038/07 HU - GEDEON RICHTER

PLC.

Lordestin 0,5 mg/ml belsöleges oldat

OGYI-T-22038/08 HU - GEDEON RICHTER

PLC.

Lordestin 0,5 mg/ml soluție orală

6327/2014/01 RO - GEDEON RICHTER

ROMÂNIA S.A.

Lordestin 0,5 mg/ml soluție orală

6327/2014/02 RO - GEDEON RICHTER

ROMÂNIA S.A.

Lordestin 0,5 mg/ml soluție orală

6327/2014/03 RO - GEDEON RICHTER

ROMÂNIA S.A.

LORDESTIN 5 mg comprimate filmate

8182/2015/01 RO - GEDEON RICHTER

ROMÂNIA S.A.

LORDESTIN 5 mg comprimate filmate

8182/2015/02 RO - GEDEON RICHTER

ROMÂNIA S.A.

LORDESTIN 5 mg comprimate filmate

8182/2015/03 RO - GEDEON RICHTER

ROMÂNIA S.A.

LORDESTIN 5 mg comprimate filmate

8182/2015/04 RO - GEDEON RICHTER

ROMÂNIA S.A.

LORDESTIN 5 mg comprimate filmate

8182/2015/05 RO - GEDEON RICHTER

ROMÂNIA S.A.

Lordestin 5 mg filmtabletta OGYI-T-

22038/01 HU - GEDEON RICHTER PLC.

Lordestin 5 mg filmtabletta OGYI-T-

22038/02 HU - GEDEON RICHTER PLC.

Lordestin 5 mg filmtabletta OGYI-T-

22038/03 HU - GEDEON RICHTER PLC.

Lordestin 5 mg filmtabletta OGYI-T-

22038/04 HU - GEDEON RICHTER PLC.

Lordestin 5 mg filmtabletta OGYI-T-

22038/05 HU - GEDEON RICHTER PLC.

Lordestin, 0,5 mg/ml, roztwór doustny

22110 PL - GEDEON RICHTER POLSKA SP. Z.

O.O.

Lordestin, 5 mg, tabletki powlekane 20074

PL - GEDEON RICHTER POLSKA SP. Z. O.O.

LOTERA 0000 PT - SVUS PHARMA A.S.

LOTERA 5 mg filmom obalené tablety

24/0038/12-S SK - SVUS PHARMA A.S.

LOTERA 5 mg potahované tablety

24/506/12-C CZ - SVUS PHARMA A.S.

RHINOHELP 2,5 mg diskia diaspeironema

sto stoma 24569/03-12-2015 GR - SPECIAL

MEDICINES PHARMACEUTICAL PRODUCTS

SINGLE-MEMBER PRIVATE COMPANY (PC)

RHINOHELP 2,5 mg diskia diaspeironema

sto stoma 24569/03-12-2015 GR - SPECIAL

MEDICINES PHARMACEUTICAL PRODUCTS

SINGLE-MEMBER PRIVATE COMPANY (PC)

RHINOHELP 5 mg diskia diaspeironema sto

stoma 24569/03-12-2015 GR - SPECIAL

MEDICINES PHARMACEUTICAL PRODUCTS

SINGLE-MEMBER PRIVATE COMPANY (PC)

RHINOHELP 5 mg diskia diaspeironema sto

stoma 24569/03-12-2015 GR - SPECIAL

MEDICINES PHARMACEUTICAL PRODUCTS

SINGLE-MEMBER PRIVATE COMPANY (PC)

RHINOHELP 5 mg diskia epikalimena me

lepto ymenio 25654/15/11-03-2016 GR -

SPECIAL MEDICINES PHARMACEUTICAL

PRODUCTS SINGLE-MEMBER PRIVATE COMPANY

(PC)

RHINOHELP 5 mg diskia epikalimena me

lepto ymenio 25654/15/11-03-2016 GR -

SPECIAL MEDICINES PHARMACEUTICAL

PRODUCTS SINGLE-MEMBER PRIVATE COMPANY

(PC)

Rinispes 0,5 mg/ml posimo dialyma

67134/11-09-2013 GR - SPECIFAR S.A.

Rinispes 5 mg diskia epikalimena me lepto

ymentio 69066/11-9-13 GR - SPECIFAR S.A.

Suprodeslon, 0,5 mg/ml, roztwór doustny

20196 PL - S-LAB SP.ZO.O.

Suprodeslon, 5 mg, tabletki powlekane

20193 PL - S-LAB SP.ZO.O.

Teslor, 5 mg, tabletki powlekane 21598 PL

- AFLOFARM FARMACJA POLSKA SP. Z O.O.

Valora, filmoovertrukne tabletter 48322 DK -

STADA ARZNEIMITTEL AG

Yosqiero 48339 DK - SANDOZ A/S

Yosqiero 48340 DK - SANDOZ A/S

Yosqiero 5 mg filmtabletta OGYI-T-

22413/01 HU - SANDOZ HUNGÁRIA KFT

Yosqiero 5 mg filmtabletta OGYI-T-

22413/02 HU - SANDOZ HUNGÁRIA KFT

Yosqiero 5 mg filmtabletta OGYI-T-
22413/03 HU - SANDOZ HUNGÁRIA KFT
Yosqiero 5 mg filmtabletta OGYI-T-
22413/04 HU - SANDOZ HUNGÁRIA KFT
Yosqiero 5 mg õhukese polümeerikattega
tabletid 786212 EE - SANDOZ
PHARMACEUTICALS D.D.
Yosqiero 5 mg õhukese polümeerikattega
tabletid 786212 EE - SANDOZ
PHARMACEUTICALS D.D.
, PRAC Rapporteur: Jean-Michel Dogné, "DLP
15/07/2016; 5 years"

EMEA/H/C/PSUSA/00000963/201607
(desloratadine / pseudoephedrine)
CAPS:
Aerinaze (EMEA/H/C/000772) (desloratadine /
pseudoephedrine sulphate), MAH: Merck Sharp
& Dohme Limited, Rapporteur: Koenraad Norga,
PRAC Rapporteur: Jean-Michel Dogné, "16 July
2012 to 15 July 2016"

EMEA/H/C/PSUSA/00009329/201608
(vemurafenib)
CAPS:
Zelboraf (EMEA/H/C/002409) (vemurafenib),
MAH: Roche Registration Limited, Rapporteur:
Filip Josephson, PRAC Rapporteur: Ulla Wändel
Liminga, "7 August 2015 to 16 August 2016"

EMEA/H/C/PSUSA/00010084/201608
(dabrafenib)
CAPS:
Tafinlar (EMEA/H/C/002604) (dabrafenib),
MAH: Novartis Europharm Ltd, Rapporteur: Filip
Josephson, PRAC Rapporteur: Ulla Wändel
Liminga, "27 August 2015 to 26 August 2016"

EMEA/H/C/PSUSA/00010093/201608
(brimonidine (centrally authorised product only))
CAPS:
Mirvaso (EMEA/H/C/002642) (brimonidine),
MAH: GALDERMA INTERNATIONAL, Rapporteur:
Filip Josephson, PRAC Rapporteur: Patrick Batty,
"22/02/2016 to 21/08/2016"

EMEA/H/C/PSUSA/00010409/201608
(panobinostat)
CAPS:
Farydak (EMEA/H/C/003725) (panobinostat),
MAH: Novartis Europharm Ltd, Rapporteur:
Paula Boudewina van Hennik, PRAC Rapporteur:
Julie Williams, "Changes to SmPC are needed in

section 4.8 as follows: ADRs with a difference in frequency < 1% between active arm and panobinostat arm are not listed (e.g. peripheral neuropathy and constipation known side effects of bortezomib treatment). (period: 23/02/2016 - 22/08/2016)"

EMEA/H/C/PSUSA/00010420/201608

(dinutuximab)

CAPS:

Unituxin (EMEA/H/C/002800) (dinutuximab),

MAH: United Therapeutics Europe Ltd,

Rapporteur: Robert James Hemmings, PRAC

Rapporteur: Sabine Straus, "15th February 2016 to 14th August 2016"

EMEA/H/C/PSUSA/00010450/201608

(cobimetinib)

CAPS:

Cotellic (EMEA/H/C/003960) (cobimetinib),

MAH: Roche Registration Limited, Rapporteur:

Filip Josephson, PRAC Rapporteur: Sabine

Straus, "24 February 2016 to 23 August 2016"

B.4. EPARs / WPARs

chondrosphere - spheroids of human autologous matrix-associated

chondrocytes - EMEA/H/C/002736, ATMP

Applicant: CO.DON AG, treatment of cartilage defects, New active substance (Article 8(3) of Directive No 2001/83/EC)

Diclofenamide SUN - diclofenamide -

EMEA/H/C/004690, Orphan

Applicant: Sun Pharmaceutical Industries Europe B.V., treatment of periodic paralysis, Known active substance (Article 8(3) of Directive No 2001/83/EC)

WPAR

Emtricitabine/tenofovir disoproxil Krka

d.d. - emtricitabine / tenofovir disoproxil -

EMEA/H/C/004686

Applicant: KRKA, d.d., Novo mesto, treatment of HIV-1 infection, Generic, Duplicate, Duplicate of Emtricitabine/Tenofovir disoproxil Krka, Generic application (Article 10(1) of Directive No 2001/83/EC)

Lokelma - sodium zirconium cyclosilicate -

EMEA/H/C/004029

Applicant: AstraZeneca AB, for the treatment of

hyperkalaemia, New active substance (Article 8(3) of Directive No 2001/83/EC)

Natpar - parathyroid hormone -

EMEA/H/C/003861, Orphan

Applicant: Shire Pharmaceuticals Ireland Ltd,
treatment of hypoparathyroidism, Known active
substance (Article 8(3) of Directive No
2001/83/EC)

Pemetrexed Hospira UK Limited -

pemetrexed - EMEA/H/C/004488

Applicant: Hospira UK Limited, treatment of
malignant pleural mesothelioma and non-small
cell lung cancer, Generic, Duplicate, Generic of
Alimta, Duplicate of Pemetrexed
ditromethamine Hospira (WD), Generic
application (Article 10(1) of Directive No
2001/83/EC)

Roteas - edoxaban - EMEA/H/C/004339

Applicant: Daiichi Sankyo Europe GmbH,
prevention of stroke; embolism and treatment
of venous thromboembolism, Informed consent
application (Article 10c of Directive No
2001/83/EC)

Varuby - rolapitant - EMEA/H/C/004196

Applicant: Tesaro UK Limited, prevention of
nausea and vomiting, New active substance
(Article 8(3) of Directive No 2001/83/EC)

B.5. TYPE II VARIATION, WORKSHARING PROCEDURE OUTCOMES

Disclosure of scopes related to Chemistry, Manufacturing, and Controls cannot be released at present time as these contain commercially confidential information.

B.5.1. CHMP assessed procedures scope: Pharmaceutical aspects

Armisarte - pemetrexed -

EMEA/H/C/004109/II/0008/G

MAH: Actavis Group PTC ehf, Rapporteur: Alar
Irs

Request for Supplementary Information adopted
on 09.03.2017.

Weekly start timetable. The Committee

adopted a Request for Supplementary
information together with a specific timetable.

Azarga - brinzolamide / timolol -

EMEA/H/C/000960/II/0035/G

MAH: Alcon Laboratories (UK) Ltd, Rapporteur:
Hanne Lomholt Larsen

Request for Supplementary Information adopted
on 19.01.2017.

Weekly start timetable.

| | | |
|--|---|---|
| Bexsero - meningococcal group B vaccine (rDNA, component, adsorbed) - EMEA/H/C/002333/11/0048 | MAH: GSK Vaccines S.r.l, Rapporteur: Kristina Dunder Request for Supplementary Information adopted on 19.01.2017. | Weekly start timetable. |
| Biopoin - epoetin theta - EMEA/H/C/001036/11/0036/G | MAH: TEVA GmbH, Rapporteur: Alexandre Moreau Request for Supplementary Information adopted on 19.01.2017. | Weekly start timetable. |
| Envarsus - tacrolimus - EMEA/H/C/002655/11/0008/G | MAH: Chiesi Farmaceutici S.p.A., Rapporteur: John Joseph Borg | Weekly start timetable. |
| Eporatio - epoetin theta - EMEA/H/C/001033/11/0035/G | MAH: ratiopharm GmbH, Rapporteur: Alexandre Moreau Request for Supplementary Information adopted on 19.01.2017. | Weekly start timetable. |
| Firazyr - icatibant - EMEA/H/C/000899/11/0036/G, Orphan | MAH: Shire Orphan Therapies GmbH, Rapporteur: Kristina Dunder | Weekly start timetable. |
| Fortacin - lidocaine / prilocaine - EMEA/H/C/002693/11/0015 | MAH: Plethora Solutions Ltd., Rapporteur: Concepcion Prieto Yerro | Weekly start timetable. |
| Hizentra - human normal immunoglobulin - EMEA/H/C/002127/11/0075 | MAH: CSL Behring GmbH, Rapporteur: Jan Mueller-Berghaus | Weekly start timetable. |
| HyQvia - human normal immunoglobulin - EMEA/H/C/002491/11/0033/G | MAH: Baxalta Innovations GmbH, Rapporteur: Jan Mueller-Berghaus Request for Supplementary Information adopted on 26.01.2017. | Weekly start timetable. |
| Inhixa - enoxaparin sodium - EMEA/H/C/004264/11/0004/G | MAH: Techdow Europe AB, Duplicate, Duplicate of Thorinane, Rapporteur: Andrea Laslop Opinion adopted on 02.03.2017. | Positive Opinion adopted by consensus on 02.03.2017. The Icelandic and Norwegian CHMP Members were in agreement with the CHMP recommendation. |

| | |
|---|--|
| Kalydeco - ivacaftor - EMEA/H/C/002494/11/0053/G, Orphan MAH: Vertex Pharmaceuticals (Europe) Ltd., Rapporteur: Concepcion Prieto Yerro Opinion adopted on 09.03.2017. Request for Supplementary Information adopted on 19.01.2017. | Positive Opinion adopted by consensus on 09.03.2017. The Icelandic and Norwegian CHMP Members were in agreement with the CHMP recommendation. |
| Keytruda - pembrolizumab - EMEA/H/C/003820/11/0020/G MAH: Merck Sharp & Dohme Limited, Rapporteur: Daniela Melchiorri Opinion adopted on 09.03.2017. | Positive Opinion adopted by consensus on 09.03.2017. The Icelandic and Norwegian CHMP Members were in agreement with the CHMP recommendation. |
| Lonquex - lipegfilgrastim - EMEA/H/C/002556/11/0030/G MAH: Sicor Biotech UAB, Rapporteur: Greg Markey | Weekly start timetable. |
| MabThera - rituximab - EMEA/H/C/000165/11/0129/G MAH: Roche Registration Limited, Rapporteur: Sinan B. Sarac Request for Supplementary Information adopted on 02.03.2017. | Weekly start timetable. The Committee adopted a Request for Supplementary information together with a specific timetable. |
| MabThera - rituximab - EMEA/H/C/000165/11/0130/G MAH: Roche Registration Limited, Rapporteur: Sinan B. Sarac Request for Supplementary Information adopted on 02.03.2017. | Weekly start timetable. The Committee adopted a Request for Supplementary information together with a specific timetable. |
| Nucala - mepolizumab - EMEA/H/C/003860/11/0007 MAH: GlaxoSmithKline Trading Services, Rapporteur: Nithyanandan Nagercoil | Weekly start timetable. |
| Nuwiq - simoctocog alfa - EMEA/H/C/002813/11/0012/G MAH: Octapharma AB, Rapporteur: Jan Mueller- Berghaus Request for Supplementary Information adopted on 19.01.2017, 13.10.2016, 14.07.2016. | Weekly start timetable. |
| Omnitrope - somatropin - EMEA/H/C/000607/11/0045 MAH: SANDOZ GmbH, Rapporteur: Johann Lodewijk Hillege | Weekly start timetable. |
| Orencia - abatacept - EMEA/H/C/000701/11/0106/G MAH: Bristol-Myers Squibb Pharma EEIG, Rapporteur: Outi Mäki-Ikola | Weekly start timetable. |

Request for Supplementary Information adopted
on 02.02.2017.

Ratiograstim - filgrastim - Weekly start timetable.

EMEA/H/C/000825/II/0053/G

MAH: ratiopharm GmbH, Rapporteur: Outi Mäki-Ikola

ReFacto AF - moroctocog alfa - Weekly start timetable.

EMEA/H/C/000232/II/0139

MAH: Pfizer Limited, Rapporteur: Hanne Lomholt Larsen

Revestive - teduglutide - Weekly start timetable.

EMEA/H/C/002345/II/0035, Orphan

MAH: Shire Pharmaceuticals Ireland Ltd,
Rapporteur: Sinan B. Sarac

Request for Supplementary Information adopted
on 26.01.2017.

Senshio - ospemifene - Positive Opinion adopted by consensus on
EMEA/H/C/002780/II/0010 02.03.2017. The Icelandic and Norwegian CHMP
MAH: Shionogi Limited, Rapporteur: Paula Members were in agreement with the CHMP
Boudewina van Hennik recommendation.

Opinion adopted on 02.03.2017.

Request for Supplementary Information adopted
on 10.11.2016, 04.08.2016.

Siklos - hydroxycarbamide - Weekly start timetable.

EMEA/H/C/000689/II/0031/G, Orphan

MAH: Addmedica, Rapporteur: Koenraad Norga

Tevagrastim - filgrastim - Weekly start timetable.

EMEA/H/C/000827/II/0063/G

MAH: TEVA GmbH, Duplicate, Duplicate of
Biograstim, Rapporteur: Outi Mäki-Ikola

Trisenox - arsenic trioxide - Weekly start timetable.

EMEA/H/C/000388/II/0063/G

MAH: Teva B.V., Rapporteur: Alexandre Moreau

Umbipro (TM) - chlorhexidine - Weekly start timetable.

EMEA/H/W/003799/II/0002/G

MAH: GlaxoSmithKline Trading Services,
Rapporteur: Patrick Salmon

Request for Supplementary Information adopted
on 26.01.2017.

Zostavax - shingles (herpes zoster) vaccine (live) - Weekly start timetable.

EMEA/H/C/000674/II/0109/G

MAH: MSD Vaccins, Rapporteur: Jan Mueller-Berghaus

WS1099/G Weekly start timetable.

Neulasta-

EMEA/H/C/000420/WS1099/0092/G

Ristempa-

EMEA/H/C/003910/WS1099/0009/G

MAH: Amgen Europe B.V., Lead Rapporteur:

Robert James Hemmings

WS1125/G

Weekly start timetable.

Helixate NexGen-

EMEA/H/C/000276/WS1125/0187/G

KOGENATE Bayer-

EMEA/H/C/000275/WS1125/0195/G

MAH: Bayer Pharma AG, Lead Rapporteur: Jan

Mueller-Berghaus

B.5.2. CHMP assessed procedures scope: Non-Clinical and Clinical aspects

Aclasta - zoledronic acid -

Weekly start timetable.

EMEA/H/C/000595/11/0068

MAH: Novartis Europharm Ltd, Rapporteur:

Kristina Dunder, "Update of section 4.8 of the SmPC in order to add the adverse reaction hypophosphataemia with an unknown frequency based on post-marketing spontaneous reports and internal databases. The package leaflet is updated accordingly.

In addition, the Marketing authorisation holder (MAH) took the opportunity to remove the lower level term 'should pain' which is covered by the corresponding preferred term 'musculoskeletal pain', to update the list of local representatives in the Package Leaflet and to bring the product information in line with the latest QRD template version 10."

Afinitor - everolimus -

Weekly start timetable.

EMEA/H/C/001038/11/0051/G

MAH: Novartis Europharm Ltd, Rapporteur:

Harald Enzmann, "C.I.13 Submission of the final clinical study report of study RAD001J2301: A randomized phase-III, double-blind, placebo-controlled multicenter trial of everolimus in combination with trastuzumab and paclitaxel, as first line therapy in women with HER2 positive locally advanced or metastatic breast cancer C.I.13 Submission of the final clinical study report of study RAD001W2301: A randomized Phase III, double-blind, placebo-controlled multicenter trial of everolimus in combination with trastuzumab and vinorelbine, in pretreated women with HER2/neu over-expressing locally advanced or metastatic breast cancer

In addition, the MAH included a report on exposure-response relationship combining data from these two trials."

Cerdelga - eliglustat - Weekly start timetable.

EMEA/H/C/003724/II/0010, Orphan

MAH: Genzyme Europe BV, Rapporteur: Johann Lodewijk Hillegje, "Update of section 5.1 of the SmPC in order to update the safety and efficacy of eliglustat from studies in the GD1 patient population (studies ENGAGE & EDGE)."

In addition, the Marketing authorisation holder (MAH) took the opportunity to update the list of local representatives in the Package Leaflet for Bulgaria and Romania."

Request for Supplementary Information adopted on 26.01.2017.

Cervarix - human papillomavirus vaccine

[types 16, 18] (recombinant, adjuvanted, adsorbed) - EMEA/H/C/000721/II/0081

MAH: GSK Biologicals SA, Rapporteur: Bart Van der Schueren, "To submit the final effectiveness results of clinical study HPV-040, a community randomized study conducted in Finland to evaluate the effectiveness of two vaccination strategies for 12 -15 year old early adolescents using Cervarix, i.e., to vaccinate female adolescents only, or to vaccinate female and male adolescents."

Request for Supplementary Information adopted on 15.12.2016, 15.09.2016.

Effentora - fentanyl - Weekly start timetable.

EMEA/H/C/000833/II/0045

MAH: Teva B.V., Rapporteur: Martina Weise, "Update of sections 4.4 and 4.5 of the SmPC in order to add a warning on increased risk of increased depressant effects with the concomitant use of alcohol and possibility of a fatal outcome with concomitant use of other CNS depressants following a cumulative review on spontaneous reporting and literature review of these risks. The package leaflet has been updated accordingly."

In addition, the marketing authorisation holder took the opportunity to introduce editorial clarifications in Annex I and Annex IIIB and changes in accordance to QRD template 10."

Enbrel - etanercept - Weekly start timetable.

EMEA/H/C/000262/II/0204

MAH: Pfizer Limited, Rapporteur: Robert James

Hemmings, "Update of section 4.8 of the SmPC in order to change the frequency category of the ADR 'elevated liver enzymes' from rare to uncommon and to add some further details on the frequency of elevated liver enzymes reported ADRs with etanercept in double-blind controlled trials with or without concomitant methotrexate use, following the assessment of Enbrel (etanercept) PSUSA/00001295/201602.). The Package Leaflet is updated accordingly. In addition, the Marketing authorisation holder (MAH) took the opportunity to update section 4.4 of the SmPC on traceability of biological medicinal products as requested by the CHMP, to make a small correction in section 6 of the 50 mg solution for injection in a pre-filled pen Package Leaflet and to bring the PI in line with the latest QRD template version 10."

EVOTAZ - atazanavir / cobicistat -
EMEA/H/C/003904/II/0010
MAH: Bristol-Myers Squibb Pharma EEIG,
Rapporteur: Bruno Sepodes, "Proposed changes to the EVOTAZ SmPC to align with the current Company Core Data Sheet (CCDS). During the EVOTAZ MAA procedure, an interim Week 144 CSR for Gilead study GS-US-216-0114 was submitted and the SmPC efficacy and safety data were updated and approved accordingly. However, the resistance data were not updated at that time. As a result, the MAH proposes to update the resistance sub-section in SmPC section 5.1 with study GS-US-216-0114 Week 144 resistance data that were submitted in the context of the MAA. In addition, for clarification purposes, the MAH proposes to use the specific designation of tenofovir disoproxil fumarate throughout the EVOTAZ Product Information (PI) to differentiate this pharmaceutical entity from the tenofovir alafenamide (for which no studies with EVOTAZ have been conducted). Finally, the MAH would like to take this opportunity to implement QRD version 10." Opinion adopted on 16.03.2017.
Request for Supplementary Information adopted on 19.01.2017, 29.09.2016.

Positive Opinion adopted by consensus on 16.03.2017. The Icelandic and Norwegian CHMP Members were in agreement with the CHMP recommendation.

EVRA - ethinylestradiol / norelgestromin - Weekly start timetable.
EMEA/H/C/000410/II/0041
MAH: Janssen-Cilag International NV,
Rapporteur: Paula Boudewina van Hennik,

"Update of sections 4.3 and 4.5 of the SmPC in order to add a contraindication for patients receiving drug combinations with Direct-acting antiviral (DAA) agents that contain paritaprevir/ritonavir, ombitasvir, and/or dasabuvir as these DAs have the potential for a drug-drug interaction with ethinyl estradiol (EE)-containing combined hormonal contraceptives resulting in ALT elevations. The Package Leaflet has been updated accordingly."

Hetlioz - tasimelteon -

Weekly start timetable.

EMEA/H/C/003870/11/0007, Orphan

MAH: Vanda Pharmaceuticals Ltd., Rapporteur: Greg Markey, PRAC Rapporteur: Adam Przybylkowski, "Submission of the final report from Study VEC-162-3T3 NRU-PT, listed as a category 3 study in the RMP. This is a study to assess the phototoxic potential of tasimelteon metabolites M12 and M14 and tasimelteon-phenol (M3 without the glucuronidation) in an in vitro neutral red uptake test using balb/c 3T3, clone 31, fibroblast cells."

HyQvia - human normal immunoglobulin -

EMEA/H/C/002491/11/0032

MAH: Baxalta Innovations GmbH, Rapporteur: Jan Mueller-Berghaus, "Update of section 4.2 and 4.8 of the SmPC in order to add information on infusion site leakage. The Package Leaflet is updated accordingly. In addition, the Marketing authorisation holder (MAH) took the opportunity to update the list of local representatives in the Package Leaflet and to bring the PI in line with the latest QRD template version 10."

Request for Supplementary Information adopted on 26.01.2017.

Invokana - canagliflozin -

EMEA/H/C/002649/11/0026

MAH: Janssen-Cilag International NV, Rapporteur: Martina Weise, "Update of section 4.4 of the SmPC in order to update the safety information: the term 'and fatal' is added when describing the Diabetic Ketoacidosis cases that have been reported. The Package Leaflet is updated accordingly: term 'rare but serious, sometimes life-threatening and fatal' is added when describing Diabetic Ketoacidosis.

In addition, the Marketing authorisation holder (MAH) took the opportunity to update the list of

local representatives in the Package Leaflet."

Request for Supplementary Information adopted
on 26.01.2017.

Kuvan - sapropterin - Weekly start timetable.
EMEA/H/C/000943/II/0046, Orphan

MAH: BioMarin International Limited,
Rapporteur: Patrick Salmon, "Update of section 4.5 to delete the statement that no interaction studies have been performed and section 5.2 to reflect the relevant results of in vitro pharmacokinetic drug interactions studies BMN162-14-021, 022, 023, BMN162-15-036 and 101.
In addition, the MAH took the opportunity of this procedure to improve the wording of section 4.2 and implement minor administrative changes in the SmPC."

Request for Supplementary Information adopted
on 19.01.2017.

Kuvan - sapropterin - Weekly start timetable. The Committee adopted a Request for Supplementary information together with a specific timetable.
EMEA/H/C/000943/II/0048/G, Orphan

MAH: BioMarin International Limited,
Rapporteur: Patrick Salmon, "Update of section 4.9 to add information regarding shortening of QT interval at high doses following review of data of study QTC-001.
Submission of the clinical study report EMR700773-004 (pilot study assessing the effect of sapropterin on cognitive abilities, study prematurely terminated due to enrolment issues)
In addition, the MAH took the opportunity of this procedure to clarify the wording of section 4.2 and section 3 of the PL."

Request for Supplementary Information adopted
on 09.03.2017.

Kyprolis - carfilzomib - Weekly start timetable.
EMEA/H/C/003790/II/0010, Orphan

MAH: Amgen Europe B.V., Rapporteur:
Aranzazu Sancho-Lopez, "Update of section 4.5 of the SmPC in order to inform the prescriber that no Drug Drug Interaction (DDI) studies were conducted at the higher dose (56mg/m2)."
Request for Supplementary Information adopted
on 19.01.2017.

Lonsurf - trifluridine / tipiracil - Weekly start timetable.
EMEA/H/C/003897/II/0003

MAH: Les Laboratoires Servier, Rapporteur:
Paula Boudewina van Hennik, "Submission of

the final report from the pharmacogenomics study (NP35044) of TAS-102 in patients with metastatic colorectal cancer refractory to standard chemotherapy (10040080) in order to fulfil a Recommendation made at the time of the initial MA."

Lumigan - bimatoprost - Weekly start timetable.

EMEA/H/C/000391/11/0052

MAH: Allergan Pharmaceuticals Ireland,
Rapporteur: Hanne Lomholt Larsen, "Update of section 4.8 to add 4 adverse events in the Eye disorders SOC in line with the Company Core Data Sheet. The Package Leaflet has been updated accordingly.

Section 3 of the PL was also amended to improve clarity of instructions.

In addition, the MAH took the opportunity to update the Product Information in line with the QRD template version 10 and implement the unique identifier 2D barcode."

NovoThirteen - catidecacog - Weekly start timetable.

EMEA/H/C/002284/11/0018

MAH: Novo Nordisk A/S, Rapporteur: Joseph Emmerich, "Update of sections 4.4, 4.8, 5.1 and 5.2 of the SmPC in order to consolidate the outcome of the clinical development programme (studies F13CD-3720 and F13CD-3835) submitted in procedures P46/014 and P46/016. Briefly, section 4.4 was updated to reflect that on-demand treatment was used in the extension study F13CD-3720, section 4.8 was updated to reflect the data on number of patients/paediatric patients and exposures, in section 5.1 the bleeding rate was updated, in section 5.2 minor amendments were made to the half-life of NovoThirteen.

In addition, the Marketing authorisation holder (MAH) took the opportunity to update Annex II with minor administrative amendments in line with QRD template 9.1 and Annex III in line with QRD template version 10.0."

Request for Supplementary Information adopted on 26.01.2017.

Prevenar 13 - pneumococcal polysaccharide conjugate vaccine (13-valent, adsorbed) -

EMEA/H/C/001104/11/0149

MAH: Pfizer Limited, Rapporteur: Kristina Dunder, "Submission of the final clinical study

report (CSR) of study B1851018, a Phase 4 study evaluating the impact of 13vPnC in reducing AOM and NP colonisation caused by *S pneumoniae* in healthy children, in accordance with the Pharmacovigilance plan outlined in the EU RMP (version 11.0)."

Request for Supplementary Information adopted on 23.02.2017.

Revestive - teduglutide -

EMEA/H/C/002345/11/0032, Orphan

MAH: Shire Pharmaceuticals Ireland Ltd,
Rapporteur: Sinan B. Sarac, "Update of sections 4.3, 4.4, and 4.8 of the SmPC in order to update the safety information in line with updated CCDS following review of the MAH's safety database. The Package Leaflet is updated accordingly.

In addition, the Marketing authorisation holder (MAH) took the opportunity to make minor editorial changes in section 5.1 of the SmPC."

Request for Supplementary Information adopted on 15.12.2016.

Revadate - eltrombopag / eltrombopag

olamine - EMEA/H/C/001110/11/0042

MAH: Novartis Europharm Ltd, Rapporteur: Aranzazu Sancho-Lopez, "Submission of the ASPIRE (TRC114968) final study report, a Three-Part Study of Eltrombopag in Thrombocytopenic Subjects with Myelodysplastic Syndromes or Acute Myeloid Leukemia (Part 1: Open-Label, Part 2: Randomized, Double-Blind, Part 3: Extension) assessing the potential risk of haematological changes, optimal dose escalation scheme and eltrombopag pharmacokinetics."

Weekly start timetable.

Rotarix - human rotavirus, live attenuated - EMEA/H/C/000639/11/0089

MAH: GlaxoSmithKline Biologicals S.A.,
Rapporteur: Bart Van der Schueren, "Update of section 5.1 to introduce effectiveness data following completion of ecological observational study EPI-ROTA-025 VE AU DB (114910) - An ecological study to assess impact of rotavirus vaccination on hospitalisations for rotavirus gastroenteritis (RV GE) in children <5 years of age in Australia.

In addition, the marketing authorisation holder took the opportunity to introduce clarifications in the SmPC."

Positive Opinion adopted by consensus on 02.03.2017. The Icelandic and Norwegian CHMP Members were in agreement with the CHMP recommendation.

Opinion adopted on 02.03.2017.

Request for Supplementary Information adopted
on 27.10.2016.

Rotarix - human rotavirus, live attenuated Weekly start timetable.
- EMEA/H/C/000639/II/0094

MAH: GlaxoSmithKline Biologicals S.A.,
Rapporteur: Bart Van der Schueren, PRAC
Rapporteur: Jean-Michel Dogné, "Submission of
the final study report for EPI-ROTA-007 VS US
DB (A phase IV, open, observational study of
the safety of Rotarix, administered to a birth
cohort in US States health insurance plans)
which is listed in the section III.4.3 of the Risk
Management Plan (RMP) version 16.
Consequently a revised RMP (version 17) is
submitted in order to update information in
relation to: the EPI-ROTA-007 VS US DB study;
the EPI-ROTA-052 BOD EU SUPP as agreed
during variation EMEA/H/C/0639/II/0086. In
addition, the MAH took this opportunity to
further update the RMP with the new due date
for submission of the final study report for
ROTA-085 PMS."

Request for Supplementary Information adopted
on 09.03.2017.

Teyesuno - tegafur / gimeracil / oteracil - Weekly start timetable.
EMEA/H/C/001242/II/0029

MAH: Nordic Group B.V., Rapporteur: Paula
Boudewina van Hennik, "Submission of the final
clinical study report for Study Salto - A phase
III randomized study of S-1 versus capecitabine
as first line treatment in metastatic colorectal
cancer."

Torisel - temsirolimus - Weekly start timetable.

EMEA/H/C/000799/II/0066, Orphan

MAH: Pfizer Limited, Rapporteur: Harald
Enzmann, "Submission of the further analysis of
a possible association of corticosteroid (pre-
)treatment and frequency and severity of
hypersensitivity/infusion reactions in study
3066K1-4438-WW (B1771007), as requested by
the CHMP during procedures
EMEA/H/C/799/MEA 023.1 and
EMEA/H/C/799/MEA 024.1. No changes to the
PI are proposed."

Torisel - temsirolimus - Weekly start timetable.

EMEA/H/C/000799/II/0067, Orphan

MAH: Pfizer Limited, Rapporteur: Harald
Enzmann, "Submission of the final report from

the Japanese post marketing surveillance (PMS) studies 3066K5-4406 and B1771016 together with the response to the questions raised by the CHMP on the interim report within procedure LEG 031.4."

Translarna - ataluren - Weekly start timetable.

EMEA/H/C/002720/II/0031, Orphan

MAH: PTC Therapeutics International Limited, Rapporteur: Johann Lodewijk Hillege, "Update of section 4.5 of the SmPC in order to update information on interaction with other medicinal products adding adefovir as a medicinal product that is a substrate of OAT1 based on results from study "Safety and PK study of co-administration of ataluren and a sensitive probe substrate of organic anion transporter 1 (OAT1)" (MEA014). The MAH took the occasion to correct a minor typographical error in the SmPC."

Uptravi - selexipag - Weekly start timetable.

EMEA/H/C/003774/II/0007

MAH: Actelion Registration Ltd., Rapporteur: Martina Weise, "Update of sections 4.4 and 4.5 of the SmPC in order to add information on pharmacokinetic interactions with gemfibrozil and rifampicin in healthy subjects, based on the final clinical study report of the completed clinical pharmacology drug-drug interaction study AC-065-113. The Package Leaflet is updated accordingly. In addition, the Marketing authorisation holder (MAH) took the opportunity to update information on the hydrolysis of selexipag based on data from the previously submitted absolute bioavailability study AC-065-110, make minor amendments to sections 5.1 and 5.2 of the SmPC and to bring the PI in line with the latest QRD template version 10."

Vokanamet - canagliflozin / metformin -

EMEA/H/C/002656/II/0023

MAH: Janssen-Cilag International NV, Rapporteur: Martina Weise, "Update of section 4.4 of the SmPC in order to update the safety information: the term 'and fatal' is added when describing the Diabetic Ketoacidosis cases that have been reported. The Package Leaflet is updated accordingly: term 'rare but serious, sometimes life-threatening and fatal' is added when describing Diabetic Ketoacidosis.

In addition, the Marketing authorisation holder (MAH) took the opportunity to update the list of local representatives in the Package Leaflet."

Request for Supplementary Information adopted on 26.01.2017.

Zepatier - elbasvir / grazoprevir - Weekly start timetable.
EMEA/H/C/004126/II/0006

MAH: Merck Sharp & Dohme Limited,
Rapporteur: Greg Markey, "Update of section 5.2 of the SmPC in order to update the information on absolute bioavailability of elbasvir following recent Company Core Data Sheet (CCDS) safety information update"

WS1041 Weekly start timetable.

CONTROLOC Control-

EMEA/H/C/001097/WS1041/0025

PANTOLOC Control-

EMEA/H/C/001100/WS1041/0029

PANTOZOL Control-

EMEA/H/C/001013/WS1041/0027

SOMAC Control-

EMEA/H/C/001098/WS1041/0026

MAH: Takeda GmbH, Lead Rapporteur: Greg Markey, "Update of sections 4.3, 4.4, 4.5, 4.6 and 4.8 of the SmPC to reflect that co-administration with HIV protease inhibitors is contraindicated (not only atazanavir), to include a warning about the reduction of the absorption of vitamin B12, and a warning about the increased risk of bone fractures and hypomagnesemia, to include drug interactions with HIV protease inhibitors in section 4.5 of the SmPC, to include that animal studies have shown excretion of pantoprazole in breast milk, and to include fracture of wrist, hip and spine as undesirable effects with unknown frequency. The package leaflet is updated accordingly."

Request for Supplementary Information adopted on 19.01.2017.

WS1066 Weekly start timetable.

Adcirca-EMEA/H/C/001021/WS1066/0026

Cialis-EMEA/H/C/000436/WS1066/0086

MAH: Eli Lilly Nederland B.V., Lead Rapporteur: Concepcion Prieto Yerro, "Update of sections 4.2 and 5.1 of the SmPC in order to reflect the results of study H6D-MC-LVJJ, a randomized, double-blind, placebo-controlled phase 3 trial of tadalafil in the treatment of Duchenne Muscular

Dystrophy (DMD), to fulfil Adcirca P46 019.1
and Cialis P46 045.1."
Request for Supplementary Information adopted
on 19.01.2017.

WS1079 Weekly start timetable.

Exviera-EMEA/H/C/003837/WS1079/0023

Viekirax-

EMEA/H/C/003839/WS1079/0028

MAH: AbbVie Ltd., Lead Rapporteur: Filip Josephson, "Update of section 4.5 to include information on the drug-drug interaction with mTOR inhibitors sirolimus and everolimus. The Package Leaflet is updated accordingly."
Request for Supplementary Information adopted on 19.01.2017.

WS1092

Ebymect-

EMEA/H/C/004162/WS1092/0017

Edistride-

EMEA/H/C/004161/WS1092/0013

Forxiga-

EMEA/H/C/002322/WS1092/0032

Xigduo-EMEA/H/C/002672/WS1092/0028

MAH: AstraZeneca AB, Lead Rapporteur: Kristina Dunder, "Update of sections 4.4 and 5.1 of the SmPC in order to reflect the results of the Phase 3 study D5553C00003: 28-week safety and efficacy, randomised, double-blind comparison of simultaneous administration of exenatide once weekly 2 mg and dapagliflozin once daily 10 mg to exenatide once weekly 2 mg alone and dapagliflozin once daily 10 mg alone in patients with type 2 diabetes with inadequate glycaemic control on metformin. The Package Leaflet is updated accordingly. In addition, the Worksharing applicant (WSA) took the opportunity to update the list of local representatives in the Package Leaflets for Ebymect and Edistride and to introduce minor editorial changes throughout the Product Informations."

WS1106 Weekly start timetable.

Exviera-EMEA/H/C/003837/WS1106/0027

Viekirax-

EMEA/H/C/003839/WS1106/0031

MAH: AbbVie Ltd., Lead Rapporteur: Filip Josephson, "Update of sections 4.4 and 4.5 of the SmPC in order to add a warning stating that concomitant use of tacrolimus with dasabuvir

and ombitasvir/paritaprevir/ritonavir should be avoided unless the benefit outweigh the risks."

WS1113

Weekly start timetable.

Stribild-EMEA/H/C/002574/WS1113/0078**Tybost-EMEA/H/C/002572/WS1113/0035**

MAH: Gilead Sciences International Ltd, Lead

Rapporteur: Robert James Hemmings,

"Submission of the final report from Study GS-

US-236-0128 listed as a category 3 study in
the RMP.

This is a randomized, double-blind phase 3B
study to evaluate the safety and efficacy of
Elvitegravir/Cobicistat/Emtricitabine/Tenofovir
Disoproxil Fumarate versus Ritonavir-boosted
Atazanavir plus Emtricitabine/Tenofovir
Disoproxil Fumarate in HIV-1 infected,
antiretroviral treatment-naive women."

WS1123**Kisplyx-EMEA/H/C/004224/WS1123/0003**

Lenvima-

EMEA/H/C/003727/WS1123/0007

MAH: Eisai Europe Ltd., Lead Rapporteur: Bart
Van der Schueren, "Update of section 4.8 of the
SmPC to add the adverse events "cholecystitis"
with a frequency of common, and the adverse
events "pancreatitis", "amylase

Increased" and "lipase increased" with a
frequency of uncommon, common and common
respectively. The Package Leaflet is updated
accordingly. A correction has been done to
section 5.2. In addition, the Worksharing
applicant (WSA) took the opportunity to
combine the Kisplyx SmPC."

B.5.3. CHMP-PRAC assessed procedures

Benlysta - belimumab -**EMEA/H/C/002015/11/0047**

MAH: Glaxo Group Ltd, Rapporteur: Kristina
Dunder, PRAC Rapporteur: Ulla Wändel Liminga,
"Submission of the final report from study
LBSL99/BEL112626 listed as a category 3 study
in the RMP (MEA010). This is "A Multi-Center,
Open Label, Continuation Trial of Monoclonal
Anti-Blys Antibody in Subjects with SLE who
completed the phase 2 Protocol LBSL02""

Cinqaero - reslizumab -**EMEA/H/C/003912/11/0005/G**

MAH: Teva Pharmaceuticals Limited,

Rapporteur: Johann Lodewijk Hillege, PRAC
Rapporteur: Brigitte Keller-Stanislawska
Request for Supplementary Information adopted
on 26.01.2017.

**Emtricitabine/Tenofovir disoproxil Mylan -
emtricitabine / tenofovir disoproxil -
EMEA/H/C/004050/11/0001**

MAH: MYLAN S.A.S, Generic, Generic of
Truvada, Rapporteur: Romaldas Mačiulaitis,
PRAC Rapporteur: Patrick Batty, "Update of the
SmPC following the assessment of the extension
of indication for the reference product, Truvada,
for pre-exposure prophylaxis. The Package
Leaflet, Annex II and Labelling are updated in
accordance."

Fampyra - fampridine -

EMEA/H/C/002097/11/0036/G

MAH: Biogen Idec Ltd, Rapporteur: Johann
Lodewijk Hillege, PRAC Rapporteur: Sabine
Straus, "This is a grouped variation proposing
updates:

- to the SmPC sections 4.2, 5.1, Annex II and
Package Leaflet based on the clinical study
Enhance,
- to the SmPC section 4.6 based on the data
from pregnancy registry.
- Further changes to the PI, section 4.2 and 5.2
of the SmPC have been introduced based on the
Core Data Sheet (CDS) and PRAC review of the
Fampyra PSUR 03.

The RMP (version 11) has been updated
accordingly. In addition, the Marketing
authorisation holder (MAH) took the opportunity
to bring the PI in line with the latest QRD
template version 10.0.

With this application the MAH requests to switch
the marketing authorisation from conditional to
standard."

Request for Supplementary Information adopted
on 26.01.2017.

Firdapse - amifampridine -

EMEA/H/C/001032/11/0043, Orphan

MAH: BioMarin Europe Ltd, Rapporteur: Greg
Markey, PRAC Rapporteur: Julie Williams,
"Update of sections 4.4 and 5.3 of the SmPC
respectively in order to delete the statements
that amifampridine has not been fully tested in
carcinogenicity models and to provide the
findings from the carcinogenicity reports

required for the completion of SOB 004.

The RMP (v.9) is proposed to be updated accordingly.

In addition, the Marketing authorisation holder (MAH) took the opportunity to request the removal of the requirement to complete carcinogenicity testing in an appropriate model in section E of the Annex II."

Request for Supplementary Information adopted on 26.01.2017, 10.11.2016, 15.09.2016.

Fluenz Tetra - influenza vaccine (live attenuated, nasal) -

Weekly start timetable.

EMEA/H/C/002617/11/0064

MAH: MedImmune LLC, Rapporteur: Bart Van der Schueren, PRAC Rapporteur: Jean-Michel Dogné, "C.I.13: Submission of the final Clinical Study Report for the study number MI-MA194: A Postmarketing Observational Evaluation of the Safety of Fluenz in Children and Adolescents with High-risk Conditions."

Request for Supplementary Information adopted on 09.03.2017.

Imbruvica - ibrutinib -

EMEA/H/C/003791/11/0029, Orphan

MAH: Janssen-Cilag International NV, Rapporteur: Filip Josephson, PRAC Rapporteur: Julie Williams, "Update of sections 4.5 of the SmPC to remove the statement that an interaction between products increasing stomach pH and ibrutinib have not been studied and section 5.2 to include the findings from study CLL1005. The Package Leaflet is not impacted by these changes.

In addition, the RMP is updated to version 6.3 to reflect this new safety information."

Request for Supplementary Information adopted on 15.12.2016.

Lonsurf - trifluridine / tipiracil -

EMEA/H/C/003897/11/0002/G

MAH: Les Laboratoires Servier, Rapporteur:

Paula Boudewina van Hennik, PRAC Rapporteur: Ulla Wändel Liminga, "1) C.I.4 (type II) - Update of sections 4.2, 4.4 and 5.2 of the SmPC following availability of the final clinical study report for the study TO-TAS-102-106, A phase I, open-label study evaluating the safety, tolerability, and pharmacokinetics of TAS-102 in patients with advanced solid tumours and varying degrees of hepatic impairment

(requested in MEA 002). As a consequence of TO-TAS-102-106 study results, the RMP (ver. 5.0) is updated to remove the missing information "Use in patients with moderate to severe hepatic impairment", and to add "Hyperbilirubinaemia in patients with baseline moderate to severe hepatic impairment" as important potential risk.

2) C.I.4 (type II) - Update of sections 4.5 and 5.2 of the SmPC following availability of the results of the in vitro CYP induction study of tipiracil hydrochloride (TPI) using the appropriate concentration of TPI (requested in a recommendation). Section SVII.4 of the RMP is updated accordingly.

3) C.I.4 (type II) - Update of section 4.2 of the SmPC in order to correct inconsistencies in the dose calculation according to body surface area. The package leaflet is updated to add 'interstitial lung disease' in the serious side effects part of section 4.

In addition, the MAH took the opportunity to update Annex IIIA in accordance with the latest QRD template."

Request for Supplementary Information adopted on 26.01.2017.

Mozobil - plerixafor -

EMEA/H/C/001030/11/0030/G, Orphan

MAH: Genzyme Europe BV, Rapporteur: Paula Boudewina van Hennik, PRAC Rapporteur: Sabine Straus, "Submission of the final report from study ARD12858 (MOZ23510) "A pilot, exploratory, randomized, phase 2 safety study evaluating tumor cell (plasma cell) mobilization and apheresis product contamination in plerixafor plus non-pegylated G-CSF mobilized patients and in non pegylated G-CSF alone mobilized patients" listed as a category 3 study in the RMP .

Submission of the final report from study OBS13611 (MOZ18009), a multicenter, noninterventional registry designed to evaluate the long-term outcomes for patients who received plerixafor for stem cell mobilization and completed hematopoietic stem cell transplantation (HSCT) compared with patients who received other mobilization methods and completed HSCT, listed as a category 3 study in the RMP.

Submission of the final report from study

OBS13612 (MOZ19310), monitoring the plerixafor off-label transplant use, in patients and donors in EBMT centers performing autologous transplants and/or allogeneic transplants, listed as a category 3 study in the RMP."

OPDIVO - nivolumab -

EMEA/H/C/003985/II/0024

MAH: Bristol-Myers Squibb Pharma EEIG,
Rapporteur: Aranzazu Sancho-Lopez, PRAC
Rapporteur: Brigitte Keller-Stanislawska,
"Update of section 5.1 of the SmPC in order to reflect the final overall survival and response data, including duration of response with longer follow-up, following completion of PAES CA209037 (Randomized, Open-Label, Phase 3 Trial of nivolumab vs Investigator's Choice in Advanced (Unresectable or Metastatic) Melanoma Patients Progressing Post Anti-CTLA-4 Therapy) and its addendum on predictability of efficacy with biomarkers.

This application fulfils ANX 001 and 003.1.

Annex II has been updated accordingly.

RMP version 5.5 has been submitted within this application."

Request for Supplementary Information adopted on 26.01.2017.

Orencia - abatacept -

Weekly start timetable.

EMEA/H/C/000701/II/0107

MAH: Bristol-Myers Squibb Pharma EEIG,
Rapporteur: Outi Mäki-Ikola, PRAC Rapporteur:
Kirsti Villikka, "Update of sections 4.4 and 4.8 of the SmPC in order to update the safety information following the MAH's initiative to update its clinical trials safety database to include all currently completed Orencia clinical trials for both the IV and SC formulations. The adverse reactions table in section 4.8, as well as the description of selected adverse reactions of special interest is being amended. Section 4.4 is being brought in line with the updated section 4.8.

The package leaflet is being revised accordingly.

An updated Risk Management Plan (Version 22) is also being submitted within this variation."

Request for Supplementary Information adopted on 09.03.2017.

Rekovelle - follitropin delta -

EMEA/H/C/003994/II/0003/G

MAH: Ferring Pharmaceuticals A/S, Rapporteur:
Joseph Emmerich, PRAC Rapporteur: Menno van
der Elst.

Remicade - infliximab -

EMEA/H/C/000240/II/0204

MAH: Janssen Biologics B.V., Rapporteur:
Kristina Dunder, PRAC Rapporteur: Ulla Wändel
Liminga, "Submission of the final registry report
from the C0168T71 study (a review and analysis
of birth outcomes from Swedish, Danish and
Finish medical birth registers) and an evaluation
of pregnancy data from multiple sources.
Section 4.6 of the SmPC, relevant section of the
PL and the RMP version 13.2 has been updated
to reflect the study results.
The MAH has also taken the opportunity to bring
the product in line with the QRD template and
update the local representative section of the
PL."

Soliris - eculizumab -

EMEA/H/C/000791/II/0086/G, Orphan

MAH: Alexion Europe SAS, Rapporteur:
Aranzazu Sancho-Lopez, PRAC Rapporteur: Eva
A. Segovia, "Type II (C.I.4): Update of section
4.8 of the SmPC with the ADR frequencies to
reflect overall exposure to eculizumab in clinical
trials. The Package Leaflet (section 4) is
updated accordingly.
Type II (C.I.3.b): update of section 4.4 of the
SmPC with warning and precautions on
meningococcal vaccination timing as
recommended by PRAC. The Package Leaflet
(sections 2 and 3) Annex II.D and the RMP (ver.
13) are updated accordingly.
In addition, the MAH took the opportunity of this
RMP update to implement the PRAC
recommendation suggesting to remove the off
label use from the missing information, to
provide the exposure data from PSUR 13 and to
update the epidemiology sections with more
complete and recent scientific literature data.
Moreover, the MAH took the opportunity of this
submission to add editorial changes and to bring
the PI in line with the latest QRD template."
Request for Supplementary Information adopted
on 15.12.2016, 15.09.2016.

Tresiba - insulin degludec -

EMEA/H/C/002498/II/0024/G

MAH: Novo Nordisk A/S, Rapporteur: Kristina

Dunder, PRAC Rapporteur: Qun-Ying Yue,
"Grouping of two variations to update sections
4.2 and 5.1 of the SmPC in order to include
updated information on the use of Tresiba in
terms of transfer from other basal insulin
regimens and the effects of Tresiba on
hypoglycaemia.

The Package Leaflet and Labelling are proposed
to be updated accordingly.

An updated RMP (version 7.0) is being
submitted.

The proposed changes reflect the findings from
two studies submitted:

NN1250-3995 (SWITCH 1) and NN1250-3998
(SWITCH 2), comparing the safety and efficacy
of Tresiba and insulin glargine U-100, mainly to
document the hypoglycaemia profile in type 1
diabetes and type 2 diabetes, respectively.

In addition, the Marketing authorisation holder
(MAH) took the opportunity to bring the PI in
line with the latest QRD template version 10.0.
Finally, minor changes have been made to the
SmPC section 4.2 and the corresponding section
of the Package Leaflet to clarify the correct use
of Tresiba."

Request for Supplementary Information adopted
on 26.01.2017.

Tyverb - Ipatinib -

EMEA/H/C/000795/II/0048/G

MAH: Novartis Europharm Ltd, Rapporteur: Filip
Josephson, PRAC Rapporteur: Ulla Wändel
Liminga, "1) C.I.4 (type II): Update of sections
4.4, 4.8, and 5.1 of the SmPC in order to add a
warning on QTc prolongation and update safety
information following the submission of study
report EGF114271 (A Phase IV placebo
controlled single sequence crossover study to
evaluate the effect of repeat oral doses of
lapatinib on cardiac repolarization in patients
with advanced cancer). The Package Leaflet is
updated accordingly.

2) C.I.4 (type II): Update of section 4.8 of the
SmPC in order to further elaborate on the
undesirable effect 'serious cutaneous reactions'
based on the review of the Novartis safety
database. The Package Leaflet is updated
accordingly.

In addition, the Marketing authorisation holder
(MAH) took the opportunity to bring the PI in
line with the latest QRD template version 10.

Moreover, the MAH took the opportunity to update Annex II to delete an Annex II condition which has been fulfilled with procedure ANX.

28.2.

The RMP (version 32) is updated accordingly to the scopes presented above and also to introduce template-related changes, study milestones updates, and to upgrade 'food effect' to an important identified risk (from procedure EMEA/H/C/000795/II/0024)."

Request for Supplementary Information adopted on 10.11.2016.

Voncento - human coagulation factor VIII

/ human von willebrand factor -

EMEA/H/C/002493/II/0017/G

MAH: CSL Behring GmbH, Rapporteur: Paula Boudewina van Hennik, PRAC Rapporteur: Sabine Straus, "C.I.4 (type II): Update of section 4.8 of the SmPC in order to update the frequencies of undesirable effects to reflect the final clinical study data from study CSLCT-BIO-08-53 in haemophilia A paediatric patients. The Package Leaflet is updated accordingly. The submission of the final CSR CSLCT-BIO-08-53 also leads to changes to the RMP (ver. 6.1) in order update the Company Core Safety Information (CCSI).

C.I.11.z (type IB): Submission of a revised RMP in order to remove the commitment to conduct a post-marketing study for haemophilia A patients (CSLCT-BIO-12-78) for Voncento as consequence of new data from study CSLCT-BIO-08-53.

In addition, the Marketing authorisation holder (MAH) took the opportunity to combine different strengths in the SmPC and Package Leaflet."

Request for Supplementary Information adopted on 10.11.2016, 01.04.2016, 19.11.2015.

Wakix - pitolisant -

EMEA/H/C/002616/II/0004/G, Orphan

MAH: BIOPROJET PHARMA, Rapporteur: Joseph Emmerich, PRAC Rapporteur: Kirsti Villikka, "Update of sections 4.4, 4.5, 4.6 and 5.2 of the SmPC based on the final CSR of study P15-02 (to assess the mass balance recovery, metabolite profile and metabolite identification of 14C-pitolisant at steady state conditions, in healthy CYP2D6 phenotyped subjects), P14-07 (to evaluate pharmacokinetic interaction of pitolisant with sodium oxybate and modafinil in

healthy male volunteers) and P15-15 (to evaluate pharmacokinetic interaction of pitolisant with CYP3A4 substrates (midazolam), CYP2B6 substrates (bupropion), UGT2B7 inhibtors (probenecide)) in fulfilment of PAM (MEA 02, 03 and 04). The Package Leaflet is updated accordingly. In addition, the Marketing authorisation holder (MAH) took the opportunity to make minor editorial change in section 4.8 of the SmPC. Moreover, updated RMP version 5.0 has been submitted as part of this application."

Xtandi - enzalutamide - Weekly start timetable.

EMEA/H/C/002639/11/0034

MAH: Astellas Pharma Europe B.V., Rapporteur: Aranzazu Sancho-Lopez, PRAC Rapporteur: Eva A. Segovia, "Update of section 5.1 of the SmPC in order to reflect the final results of the post authorisation efficacy study (PAES) CL-9785-0410 which was a study of enzalutamide in patients with progressive mCRPC previously treated with abiraterone Acetate, listed as a category 3 in the RMP. The RMP version 11.0 has also been submitted."

Xtandi - enzalutamide - Weekly start timetable.

EMEA/H/C/002639/11/0035

MAH: Astellas Pharma Europe B.V., Rapporteur: Aranzazu Sancho-Lopez, PRAC Rapporteur: Eva A. Segovia, "Update of sections 4.4 and 4.8 of the SmPC to reflect the final results of the post authorisation safety study (PASS) CL-9785-0403 which evaluated the risk of seizure among subjects with mCRPC treated with enzalutamide who were at potential increased risk of seizure (UPWARD) and was listed as a category 3 in the RMP. The RMP version 11.0 has also been submitted.

In addition, the Marketing authorisation holder (MAH) took the opportunity to make a correction in section 5.1 of the SmPC."

Request for Supplementary Information adopted on 09.03.2017.

Xtandi - enzalutamide -

EMEA/H/C/002639/11/0036

MAH: Astellas Pharma Europe B.V., Rapporteur: Aranzazu Sancho-Lopez, PRAC Rapporteur: Eva A. Segovia, "Update of sections 4.6 and 5.3 of the SmPC to reflect the final results of study AE-7592-G, "Transfer of Radioactivity into Fetuses and Breast Milk in Rats after a Single Oral

Administration of [14C] MDV3100- ISN: 9785-ME-0046". The Package Leaflet is updated accordingly. The RMP version 11.0 has also been submitted."

Opinion adopted on 09.03.2017.

**Xultophy - insulin degludec / liraglutide -
EMEA/H/C/002647/II/0017**

MAH: Novo Nordisk A/S, Rapporteur: Kristina Dunder, PRAC Rapporteur: Menno van der Elst, "Update of section 4.2 of the SmPC in order to update the information on use of Xultophy in patients with hepatic impairment, based on clinical trial NN2211-1328, the LEAD 1-6 meta-analysis as well as other liraglutide trials.

In addition, 'fatigue' has been added to the tabulated list of adverse reactions in Section 4.8 of the SmPC. The Package Leaflet is updated accordingly.

RMP version 6.0 has also been submitted.

In addition, the Marketing authorisation holder (MAH) took the opportunity to bring the PI in line with the latest QRD template version 10."

WS1075

Epclusa-

EMEA/H/C/004210/WS1075/0006

Harvoni-

EMEA/H/C/003850/WS1075/0043

Sovaldi-EMEA/H/C/002798/WS1075/0037

MAH: Gilead Sciences International Ltd, Lead Rapporteur: Filip Josephson, Lead PRAC Rapporteur: Ana Sofia Diniz Martins, "Submission of the final non-clinical study report PC-334-2035 assessing the potential for a pharmacokinetic interaction via transporter or enzyme based inhibition when sofosbuvir and other Direct Acting Antivirals (DAAs) are used concomitantly with amiodarone

The RMPs (Epclusa – RMP version 1.0, Harvoni – RMP version 2.0, Sovaldi – RMP version 5.0) have been updated accordingly."

Request for Supplementary Information adopted on 26.01.2017.

WS1086

Stribild-EMEA/H/C/002574/WS1086/0077

Tybost-EMEA/H/C/002572/WS1086/0034

MAH: Gilead Sciences International Ltd, Lead Rapporteur: Robert James Hemmings, Lead PRAC Rapporteur: Patrick Batty, "Submission of the final report from Study GS-US-236-0140.

This is a randomized, open-label, phase 4 study evaluating the renal effect of Elvitegravir/Cobicistat/ Emtricitabine/Tenofovir DF or other Tenofovir DF-containing Regimens (Ritonavir-boosted Atazanavir plus Emtricitabine /Tenofovir DF or Efavirenz /Emtricitabine/Tenofovir DF) compared to Ritonavir-boosted Atazanavir plus Abacavir/ Lamivudine in Antiretroviral Treatment-naïve HIV-1 Infected Adults with eGFR ≥70 mL/min."

WS1089/G

Prezista-

EMEA/H/C/000707/WS1089/0086/G

Rezolsta-

EMEA/H/C/002819/WS1089/0018/G

MAH: Janssen-Cilag International NV, Lead

Rapporteur: Johann Lodewijk Hillege, Lead

PRAC Rapporteur: Menno van der Elst,

"Submission of the final report from Study GS-US-236-0140 listed as a category 3 study in the RMP. This is a randomized, open-label, phase 4 study evaluating the renal effect of Elvitegravir/Cobicistat/ Emtricitabine/Tenofovir DF or other Tenofovir DF-containing Regimens (Ritonavir-boosted Atazanavir plus Emtricitabine /Tenofovir DF or Efavirenz /Emtricitabine/Tenofovir DF) compared to Ritonavir-boosted Atazanavir plus Abacavir/ Lamivudine in Antiretroviral Treatment-naïve HIV-1 Infected Adults with eGFR ≥70 mL/min.

The RMP has been updated accordingly and the important potential risks of renal toxicity removed.

Based on cumulative review of the available data, the Prezista and Rezolsta RMPs are updated to remove the important risks of 'pancreatitis', 'convulsions' and 'cardiac conduction abnormalities' and the important risk 'development of drug resistance' in the Rezolsta RMP."

WS1103

Ebymect-

EMEA/H/C/004162/WS1103/0018

Xigduo-EMEA/H/C/002672/WS1103/0029

MAH: AstraZeneca AB, Lead PRAC Rapporteur:

Julie Williams, "The Applicant submitted a Type IB worksharing to update the RMP of Xigduo and its duplicate Ebymect. The proposed changes are in line with the outcome of the article 31 referral on metformin and metformin-containing

medicines regarding the use in patients with moderate renal impairment (EMEA/H/A-31/1432). The Commission Decision for this article 31 referral was adopted on 12th December 2016."

WS1128

Gardasil-

EMEA/H/C/000703/WS1128/0071

Silgard-EMEA/H/C/000732/WS1128/0062

MAH: MSD Vaccins, Lead Rapporteur: Kristina Dunder, Lead PRAC Rapporteur: Qun-Ying Yue, "Update of section 5.1 of the SmPC in order to following/based on the final report for Study P019-21 (Gardasil MEA 060.2 and Silgard MEA 059.2) and fourth interim report for Study P015-21 (Gardasil/Silgard MEA 019.7).

Study P019-21 is a long-term Follow-up Study of Safety, Immunogenicity, and Effectiveness of Gardasil (Human Papillomavirus [Types 6, 11, 16, 18] Recombinant Vaccine) in Mid-Adult Women - The FUTURE III (Females United to Unilaterally Reduce Endo/Ecto Cervical Cancer).

Study P015-21 is a registry-based Study of Protocol V501-015 Subjects, and Recipients of Gardasil recombinant vaccine in Countries with centralized cervical cancer screening infrastructures to evaluate the long-term effectiveness, immunogenicity, and safety of Gardasil.

The RMP version 11 has also been submitted."

B.5.4. PRAC assessed procedures

PRAC Led

Cervarix - human papillomavirus vaccine [types 16, 18] (recombinant, adjuvanted, adsorbed) - EMEA/H/C/000721/II/0086

MAH: GSK Biologicals SA, Rapporteur: Bart Van der Schueren, PRAC Rapporteur: Jean-Michel Dogné, PRAC-CHMP liaison: Bart Van der Schueren, "Submission of the final report from study HPV-039, listed in the RMP as one of the measures to bring additional information on the theoretical risk of acquiring vaccine-induced autoimmune diseases and on pregnancy outcomes after vaccination.

With this submission the MAH fulfils post-authorisation measure MEA 081."

Opinion adopted on 09.03.2017.

PRAC Led

**Corbilta - levodopa / carbidopa /
entacapone - EMEA/H/C/002785/II/0009**

MAH: Orion Corporation, PRAC Rapporteur:
Kirsti Villikka, PRAC-CHMP liaison: Outi Mäki-Ikola, "Submission of the final report of pharmacoepidemiological registry study CCOM998A2001, as requested in PRAC PSUR Assessment report

EMEA/H/C/PSUSA/00000547/201510. The study is listed as category III studies in the Risk Management plan (RMP) of Corbilta and the summary results indicate that treatment with entacapone does not increase the risk of myocardial infarction in patients with Parkinson's disease.

The RMP of Corbilta is updated accordingly from version 1.1 to version 2.0.

MA holder does not propose any changes to the Product Information of Corbilta as a consequence of this Type II variation."

PRAC Led

**Corbilta - levodopa / carbidopa /
entacapone - EMEA/H/C/002785/II/0010**

MAH: Orion Corporation, Rapporteur: Outi Mäki-Ikola, PRAC Rapporteur: Kirsti Villikka, PRAC-CHMP liaison: Outi Mäki-Ikola, "Submission of the final report of pharmacoepidemiological registry study ER11-9411 was requested in PRAC PSUR assessment report

EMEA/H/C/PSUSA/00000547/201510. The study is listed as category III study in the Risk Management Plan (RMP) and the summary results indicate that treatment with entacapone does not increase the risk of prostate cancer in patients with Parkinson's disease.

The RMP of Corbilta is updated accordingly from version 1.1 to version 2.0.

MA holder does not propose any changes to the Product Information of Corbilta as a consequence of this Type II variation."

PRAC Led

Humira - adalimumab -

EMEA/H/C/000481/II/0159

MAH: AbbVie Ltd., Rapporteur: Kristina Dunder, PRAC Rapporteur: Ulla Wändel Liminga, PRAC-CHMP liaison: Kristina Dunder, "Submission of study P06-134: "A Long-Term Non-Interventional Registry to Assess Safety and Effectiveness of Humira in Subjects with

Moderately to Severely Active Crohn's Disease" in fulfilment fo MEA 056.9. The study includes also some paediatric patients and fulfils article 46 paediatric obligations."

Request for Supplementary Information adopted on 10.11.2016.

PRAC Led

Weekly start timetable.

Orencia - abatacept -

EMEA/H/C/000701/II/0108/G

MAH: Bristol-Myers Squibb Pharma EEIG, PRAC Rapporteur: Kirsti Villikka, "This grouping of two type II variations (category C.I.13) covers the submission of the final clinical study reports from epidemiological studies IM101045A & IM101045B, listed as category 3 studies in the RMP.

IM101045A & IM101045B are both observational studies, sharing overlapping safety objectives (e.g.: to assess the risk of infections, infusion-related reactions, autoimmune disorders, injection reactions and combination use)."

Request for Supplementary Information adopted on 09.03.2017.

PRAC Led

Ozurdex - dexamethasone -

EMEA/H/C/001140/II/0025

MAH: Allergan Pharmaceuticals Ireland, Rapporteur: Greg Markey, PRAC Rapporteur: Julie Williams, PRAC-CHMP liaison: Greg Markey, "In line with the RMP commitment, submission of the final report for the Post-Authorisation Safety Study 206207-025 (A Prospective Observational Study to Evaluate Long-Term Safety in Real-World Clinical Practice.)"

Request for Supplementary Information adopted on 26.01.2017.

PRAC Led

Weekly start timetable. The Committee adopted a Request for Supplementary information together with a specific timetable.

Pradaxa - dabigatran etexilate -

EMEA/H/C/000829/II/0100

MAH: Boehringer Ingelheim International GmbH, Rapporteur: Hanne Lomholt Larsen, PRAC Rapporteur: Torbjorn Callreus, PRAC-CHMP liaison: Sinan B. Sarac, "Submission of the final report for study 1160.144, which evaluated the potential off-label use of dabigatran etexilate in Europe: A drug utilisation study in Cegedim France, Denmark, and CPRD

UK."

Request for Supplementary Information adopted
on 09.03.2017.

PRAC Led

**Pradaxa - dabigatran etexilate -
EMEA/H/C/000829/II/0101**

MAH: Boehringer Ingelheim International GmbH, Rapporteur: Hanne Lomholt Larsen, PRAC Rapporteur: Torbjorn Callreus, PRAC-CHMP liaison: Sinan B. Sarac, "Submission of the final report of study 1160.162, an observational study assessing the management of gastrointestinal and urogenital bleeding events in patients with non valvular atrial fibrillation treated with dabigatran etexilate."

Opinion adopted on 09.03.2017.

PRAC Led

**Suboxone - buprenorphine / naloxone -
EMEA/H/C/000697/II/0035**

MAH: Indivior UK Limited, Rapporteur: Martina Weise, PRAC Rapporteur: Martin Huber, PRAC-CHMP liaison: Martina Weise, "Submission of the final study report for PEUS004 , a retrospective observational survey on Suboxone use in France. Consequently , the RMP (RMP 12.1) has been updated."

Opinion adopted on 09.03.2017.

PRAC Led

WS1059

Prezista-

EMEA/H/C/000707/WS1059/0084

Rezolsta-

EMEA/H/C/002819/WS1059/0015

MAH: Janssen-Cilag International NV, Lead Rapporteur: Johann Lodewijk Hillege, Lead PRAC Rapporteur: Menno van der Elst, PRAC-CHMP liaison: Johann Lodewijk Hillege, "Submission of an updated RMP version 3.1 in order to propose the deletion of the cat 3 study TMC114HIV3015 in HIV-1 infected pregnant women and replace the commitment by the assessment of the pharmacokinetics data in HIV-1 pregnant women."

Request for Supplementary Information adopted on 26.01.2017.

PRAC Led

WS1063

Exviera-EMEA/H/C/003837/WS1063/0022

Viekirax-

EMEA/H/C/003839/WS1063/0027

MAH: AbbVie Ltd., Lead Rapporteur: Filip Josephson, Lead PRAC Rapporteur: Dolores Montero Corominas, PRAC-CHMP liaison: Concepcion Prieto Yerro, "To update the RMP for Exviera and Viekirax with the following changes:

1. Addition of information on cases of hepatic decompensation observed in patients with Child-Pugh B hepatic impairment, and the revision of the SmPC to change the dose recommendation of these patients to "not recommended", as well as the addition of statements recommending the monitoring of hepatic function in these patients as approved on 25 January 2016 (Ref: EMEA/H/C/WS/0873).
2. Addition of a reference to nine drug-drug interaction studies as approved on 28 April 2016 (Ref: EMEA/H/C/WS0896/G).
3. Addition to the reference to the completion of rat 2 year carcinogenicity studies on dasabuvir (Exviera) and ombitasvir (Viekirax) as approved on 24 September 2015 (Ref: EMEA/H/C/003837/II/0006 and EMEA/H/C/003839/II/0004).
4. Update of section 4.2 of SmPC for Virkirax to recommend a decrease in treatment duration of 12 weeks in GT4 cirrhotic patients, with a consequential change to sections 4.4 and 5.1 as approved on 18 August 2016 (Ref: EMEA/H/C/003839/II/0022/G).
5. Removal of the nonclinical PAMS 1-3 in the initial RMP, (Ref: EMEA/H/C/03837/MEA/003, EMEA/H/C/038397/MEA/002, EMEA/H/C/03839/MEA/003)."

Request for Supplementary Information adopted on 15.12.2016.

B.5.5. CHMP-CAT assessed procedures**B.5.6. CHMP-PRAC-CAT assessed procedures****B.5.7. PRAC assessed ATMP procedures****B.5.8. Unclassified procedures and worksharing procedures of type I variations**

WS0934/G

Weekly start timetable.

Suboxone-**EMEA/H/C/000697/WS0934/0034/G**

MAH: Indivior UK Limited, Lead Rapporteur:

Martina Weise

Request for Supplementary Information adopted
on 16.02.2017.

WS0972/G**Infanrix hexa-****EMEA/H/C/000296/WS0972/0211/G**

MAH: GSK Biologicals SA, Lead Rapporteur: Bart

Van der Schueren

Opinion adopted on 09.03.2017.

Positive Opinion adopted by consensus on

09.03.2017. The Icelandic and Norwegian CHMP

Members were in agreement with the CHMP
recommendation.

WS1048/G**Infanrix hexa-****EMEA/H/C/000296/WS1048/0212/G**

MAH: GSK Biologicals SA, Lead Rapporteur: Bart

Van der Schueren

Request for Supplementary Information adopted
on 26.01.2017.

WS1111

Weekly start timetable.

Entresto-**EMEA/H/C/004062/WS1111/0011****Neparvis-****EMEA/H/C/004343/WS1111/0009**

MAH: Novartis Europharm Ltd, Lead

Rapporteur: Johann Lodewijk Hillege

WS1121

Weekly start timetable.

Lyrica-EMEA/H/C/000546/WS1121/0086**Pregabalin Pfizer-****EMEA/H/C/003880/WS1121/0016**

MAH: Pfizer Limited, Lead Rapporteur: Johann

Lodewijk Hillege

WS1126

Weekly start timetable.

Gardasil-**EMEA/H/C/000703/WS1126/0070****Silgard-EMEA/H/C/000732/WS1126/0061**

MAH: MSD Vaccins, Lead Rapporteur: Kristina

Dunder

WS1140**Glyxambi-****EMEA/H/C/003833/WS1140/0002****Jentadueto-****EMEA/H/C/002279/WS1140/0037****Trajenta-****EMEA/H/C/002110/WS1140/0027**

MAH: Boehringer Ingelheim International

GmbH, Lead Rapporteur: Johann Lodewijk

B.5.9. Information on withdrawn type II variation / WS procedure

B.5.10. Information on type II variation / WS procedure with revised timetable

Saxenda - liraglutide - Request for clock stop extension.

EMEA/H/C/003780/II/0011

MAH: Novo Nordisk A/S, Rapporteur: Johann Lodewijk Hillege, PRAC Rapporteur: Menno van der Elst, "Based on submission of the LEADER clinical study results (EX2211-3748: liraglutide effect on and action in diabetes, evaluation of cardiovascular outcome results), changes to sections 4.4, and 5.1 of the SmPC are being proposed in order to update the safety information and include a description of the clinical study outcomes. The Package Leaflet and Labelling are updated accordingly. In addition, the Marketing authorisation holder (MAH) took the opportunity to implement minor editorial changes throughout the product information.

The LEADER study was included in the liraglutide RMP as a required pharmacovigilance activity (category 3) to specifically address the important potential risk of cardiovascular disorders in patients with Type 2 Diabetes Mellitus. Updates to the liraglutide RMP based on the study results are also proposed: this variation application fulfils two post-approval commitments in relation to the cardiovascular outcomes trial (MEA 002), as well as to provide additional information on the breast cancer cases found in LEADER (MEA 005). RMP Version 27 was submitted with the application. These liraglutide RMP modifications are in line with the proposed updates to the Saxenda Product Information described above."

Request for Supplementary Information adopted on 23.02.2017.

B.6. START OF THE PROCEDURES TIMETABLES FOR INFORMATION

B.6.1. Start of procedure for New Applications: timetables for information

B.6.2. Start of procedure for Extension application according to Annex I of Reg. 1234/2008): timetables for information

B.6.3. Restart of procedure - responses received to Day 120 List of Questions timetables: for information

- paclitaxel - EMEA/H/C/004154, Orphan

Applicant: Oasmia Pharmaceutical AB,

treatment of ovarian cancer

List of Questions adopted on 23.06.2016.

B.6.4. Annual Re-assessments: timetables for adoption

B.6.5. Renewals of Marketing Authorisations: timetables for adoption provided only if the validation has been completed

B.6.6. VARIATIONS – START OF THE PROCEDURE

Timetables for adoption provided that the validation has been completed.

B.6.7. Type II Variations scope of the Variations: Extension of indication

B.6.8. CHMP assessed procedures scope: Pharmaceutical aspects

- meningococcal group B vaccine (rDNA, component, adsorbed) -

EMEA/H/C/002333/II/0051

- darunavir -

EMEA/H/C/004068/II/0001/G

- etanercept -

EMEA/H/C/000262/II/0207/G

- imatinib - EMEA/H/C/002585/II/0026

, Generic, Generic of Glivec

- infliximab -

EMEA/H/C/002778/II/0050/G

, Duplicate, Duplicate of Remsima

- ivacaftor - EMEA/H/C/002494/II/0057,

Orphan

MAH: Vertex Pharmaceuticals (Europe) Ltd.

- meningococcal group A, C, W135 and Y

conjugate vaccine -

EMEA/H/C/001095/11/0065

- romiplostim -

EMEA/H/C/000942/11/0062/G, Orphan

MAH: Amgen Europe B.V.

- somatropin -

EMEA/H/C/000607/11/0047

- human normal immunoglobulin -

EMEA/H/C/000831/11/0114/G

- infliximab -

EMEA/H/C/002576/11/0042/G

- eculizumab -

EMEA/H/C/000791/11/0093, Orphan

MAH: Alexion Europe SAS

WS1132

Fiasp-EMEA/H/C/004046/WS1132/0002

NovoMix-

EMEA/H/C/000308/WS1132/0089

NovoRapid-

EMEA/H/C/000258/WS1132/0117

Ryzodeg-

EMEA/H/C/002499/WS1132/0022

WS1143

Aflunov-

EMEA/H/C/002094/WS1143/0033

Foclivia-

EMEA/H/C/001208/WS1143/0028

WS1145/G

Aflunov-

EMEA/H/C/002094/WS1145/0034/G

Foclivia-

EMEA/H/C/001208/WS1145/0029/G

WS1155

Abseamed-

EMEA/H/C/000727/WS1155/0063

Binocrit-

EMEA/H/C/000725/WS1155/0063

Epoetin alfa Hexal-

EMEA/H/C/000726/WS1155/0062

B.6.9. CHMP assessed procedures scope: Non-Clinical and Clinical aspects

Cerdelga - eliglustat -

EMEA/H/C/003724/11/0011, Orphan

MAH: Genzyme Europe BV, Rapporteur: Johann Lodewijk Hillege, Update of section 5.1 of the SmPC in order to reflect the final study results from study GZGD00304 ("A Phase 2, Open-Label, Multi-Center Study Evaluating the Efficacy, Safety and Pharmacokinetics of Genz-112638 in Gaucher Type 1 Patients") listed as a category 3 study in the RMP (MEA 007). In addition, the Marketing authorisation holder (MAH) took the opportunity to update the list of local representatives in the Package Leaflet."

Cyanokit - hydroxocobalamin -

EMEA/H/C/000806/11/0031

MAH: SERB SA, Rapporteur: Alexandre Moreau, "Update of sections 4.4 and 4.8 of the SmPC in order to add a warning on renal disorders and to update the safety information on skin and subcutaneous tissue disorders, renal and urinary disorders following a safety signal on renal disorders. The package leaflet is updated accordingly."

Docetaxel Winthrop - docetaxel -

EMEA/H/C/000808/11/0051

MAH: Aventis Pharma S.A., Informed Consent of Taxotere, Rapporteur: Alexandre Moreau, Update of section 4.8 of the SmPC to update the safety information related to electrolyte imbalance. The Package Leaflet is updated accordingly.

In addition, the Marketing authorisation holder (MAH) took the opportunity to update the list of local representatives for Bulgaria and introduce minor corrections in the Package Leaflet."

Edurant - rilpivirine -

EMEA/H/C/002264/11/0025

MAH: Janssen-Cilag International NV, Rapporteur: Johann Lodewijk Hillege "Update of section 4.5 of the SmPC in order to include Pharmacokinetics data of drug-drug interactions between simeprevir and rilpivirine, based on final result from study TMC435-TiDP16-C114; this is a Phase I, 2-panel, open-label, randomized, cross-over study in healthy subjects to investigate the potential drug-drug interaction between simeprevir and RPV."

In addition, the Marketing authorisation holder (MAH) took the opportunity to bring the PI in line with the latest QRD template version 10."

Elonva - corifollitropin alfa -

EMEA/H/C/001106/11/0033

MAH: Merck Sharp & Dohme Limited,
Rapporteur: Paula Boudewina van Hennik,
"Update of section 4.8 of the SmPC to add the new ADR 'hypersensitivity reactions (both local and generalized, including rash)' identified through post-marketing surveillance. The Package Leaflet has been updated accordingly."

Elonva - corifollitropin alfa -

EMEA/H/C/001106/11/0034

MAH: Merck Sharp & Dohme Limited,
Rapporteur: Paula Boudewina van Hennik,
"Update of section 4.5 of the SmPC to add information pertaining to potential hCG cross-reactivity resulting in a false positive pregnancy test.
In addition, the MAH is taking the opportunity to implement changes in the annexes in line with the QRD templates (versions 9.1 and 10) and to propose combined versions of the SmPCs and Package Leaflets for the different strengths."

Eperzan - albiglutide -

EMEA/H/C/002735/11/0032

MAH: GlaxoSmithKline Trading Services,
Rapporteur: Kristina Dunder, "Submission of the final clinical study report of the study 201834: A randomized, double-blind, single-dose, placebo controlled, 2-way cross-over study evaluating effect of albiglutide on cholecystokinin-induced gallbladder emptying in fasting healthy subjects, listed as a category 3 study in the RMP."

Epivir - lamivudine -

EMEA/H/C/000107/11/0104

MAH: ViiV Healthcare UK Limited, Rapporteur: Joseph Emmerich, "Update of section 4.5 of the SmPC of both Epivir tablets and oral solution, and section 4.4 of the SmPC for Epivir Oral solution only, to add information regarding the potential for interaction between lamivudine and sorbitol based on the results of Study 204857. Further, a minor amendment has been implemented throughout the SmPC to update the clinical terminology for 'Pneumocystis carinii pneumonia' to 'Pneumocystis jiroveci pneumonia'. In addition, the MAH has taken the

opportunity to align the product information with the QRD template version 10, to make minor editorial changes in the annexes and to update the contact details of the local representative in Norway in the Package Leaflet."

Galafold - migalastat -

EMEA/H/C/004059/II/0009, Orphan

MAH: Amicus Therapeutics UK Ltd Rapporteur: Johann Lodewijk Hillege, "Update of section 5.1 of the SmPC to add new mutations in Table 2: GalaFold (migalastat) amenability table and to Table 3: Mutations not amenable to GalaFold (migalastat)."

In addition, the Marketing authorisation holder (MAH) took the opportunity to introduce some minor editorial changes to the tables and to update the list of local representatives in the Package Leaflet."

Harvoni - ledipasvir / sofosbuvir -

EMEA/H/C/003850/II/0049

MAH: Gilead Sciences International Ltd, Rapporteur: Filip Josephson, "Update of section 4.8 of the SmPC in order to add angioedema with frequency 'unknown'. The Package Leaflet is updated accordingly."

Kadcyla - trastuzumab emtansine -

EMEA/H/C/002389/II/0031

MAH: Roche Registration Limited, Rapporteur: Sinan B. Sarac, "To update the SmPC section 4.4 Special Warning and Precautions for use and section 4.8 Undesirable Effects to include haemorrhage under its own heading. The package leaflet is amended accordingly."

Lixiana - edoxaban -

EMEA/H/C/002629/II/0012

MAH: Daiichi Sankyo Europe GmbH, Rapporteur: Concepcion Prieto Yerro, "Update of sections 4.2 and 5.1 of the SmPC in order to add information deriving from new clinical data for the use of edoxaban as anticoagulant therapy for patients with non-valvular atrial fibrillation undergoing cardioversion. The Package Leaflet is updated accordingly. In addition, the Marketing authorisation holder (MAH) took the opportunity to update the list of local representatives for Portugal in the Package Leaflet and to bring the PI in line with the latest

Pyramax - pyronaridine / artesunate -

EMEA/H/W/002319/II/0015

MAH: Shin Poong Pharmaceutical Co., Ltd.,
Rapporteur: Joseph Emmerich
Submission of the final report from study SP-C-013-11 listed as a category 3 study in the RMP. This is a phase IIIb/IV comparative, randomised, multi-centre, open label, parallel 3-arm clinical study to assess the safety and efficacy of repeated administration of pyronaridine-artesunate, dihydroartemisinin-piperaquine or artemether-lumefantrine or artesunate-amodiaquine over a 2-year period in children and adult patients with acute uncomplicated Plasmodium sp. malaria."

Taxotere - docetaxel -

EMEA/H/C/000073/II/0125

MAH: Aventis Pharma S.A., Rapporteur: Alexandre Moreau, "Update of section 4.8 of the SmPC to update the safety information related to electrolyte imbalance. The Package Leaflet is updated accordingly.
In addition, the Marketing authorisation holder (MAH) took the opportunity to update the list of local representatives for Bulgaria and introduce minor corrections in the Package Leaflet."

Tybost - cobicistat -

EMEA/H/C/002572/II/0036

MAH: Gilead Sciences International Ltd,
Rapporteur: Robert James Hemmings,
"Submission of the integrated resistance analysis (PC-236-2016) of the genotypic changes in the protease gene for all HIV-1 infected subjects participating in Phase 3 clinical trials of Stribild (GS-US-236-0102, GS-US-236-0103, GS-US-236-0128, GS-US-264-0110, GS-US-236-0121 and GS-US-236-0123) listed as category 3 studies in the RMP."

Wakix - pitolisant -

EMEA/H/C/002616/II/0007, Orphan

MAH: BIOPROJET PHARMA, Rapporteur: Joseph Emmerich, "Submission of the final CSR for Study P11-11; a multi-centre, single dose trial to evaluate the pharmacokinetics of pitolisant in children from 6 to less than 18 years with narcolepsy (Measure 3 of the agreed PIP)."

Zeffix - lamivudine -

EMEA/H/C/000242/II/0069

MAH: Glaxo Group Ltd, Duplicate, Duplicate of Epivir, Rapporteur: Joseph EmmerichUpdate of section 4.5 of the SmPC to add information regarding a potential interaction with sorbitol-containing medicines. The Package Leaflet has been updated accordingly. In addition, the MAH took the opportunity to implement a minor change in the labelling in line with the QRD template version 10."

WS1135

Glyxambi-

EMEA/H/C/003833/WS1135/0003

Jardiance-

EMEA/H/C/002677/WS1135/0030

Synjardy-

EMEA/H/C/003770/WS1135/0026

MAH: Boehringer Ingelheim International

GmbH, Lead Rapporteur: Johann Lodewijk

Hillegge

WS1136

Descovy-

EMEA/H/C/004094/WS1136/0017

Genvoaya-

EMEA/H/C/004042/WS1136/0031

Odefsey-

EMEA/H/C/004156/WS1136/0013

MAH: Gilead Sciences International Ltd, Lead Rapporteur: Robert James Hemmings, "Update of sections 4.4, 4.8. 5.1 and 5.2 of the SmPC in order to provide 48 weeks data from Study GS-US-292-1249; this is a Phase 3b open-label study of the efficacy and safety of elvitegravir/cobicistat/emtricitabine/tenofovir alafenamide single-tablet regimen in HIV-1/Hepatitis B co-infected adults.

The Package Leaflet is updated accordingly.

In addition, the Worksharing applicant (WSA) took the opportunity to make minor administrative changes in the SmPC and the Package Leaflet."

WS1137

Lyrica-EMEA/H/C/000546/WS1137/0087

Pregabalin Pfizer-

EMEA/H/C/003880/WS1137/0017

MAH: Pfizer Limited, Lead Rapporteur: Johann Lodewijk Hillegge"Update of sections 4.8 and 5.1 of the SmPC in order to reflect final results from paediatric study A0081041: "A Double-Blind, Placebo-Controlled, Parallel-Group, Multicenter

Study of the Efficacy and Safety of Pregabalin
as Adjunctive Therapy in Children 4-16 Years of
Age with Partial Onset Seizures"."

WS1144/G

Afinitor-

EMEA/H/C/001038/WS1144/0052/G

Votubia-

EMEA/H/C/002311/WS1144/0042/G

MAH: Novartis Europharm Ltd, Lead

Rapporteur: Harald Enzmann "Update of sections 4.4 and 4.8 of the SmPC in order to include new safety information on stomatitis and its

management based on final results from study CRAD001JUS226: a phase II, single arm study of the use of steroid-based mouthwash to prevent stomatitis in postmenopausal women with advanced or metastatic hormone receptor positive breast cancer being treated with everolimus plus exemestane

Update of section 4.6 of the SmPC in order to add new information on breast-feeding

The Package Leaflets are updated accordingly.

In addition, the Worksharing applicant (WSA) took the opportunity to bring the Afinitor PI in line with the latest QRD template version 10."

WS1152

Descovy-

EMEA/H/C/004094/WS1152/0016

Genvoya-

EMEA/H/C/004042/WS1152/0030

Odefsey-

EMEA/H/C/004156/WS1152/0012

MAH: Gilead Sciences International Ltd, Lead

Rapporteur: Robert James Hemmings, "Update of sections 4.8 and 5.1 of the SmPC in order to amend the information regarding undesirable effects and pharmacodynamic properties of Genvoya, Descovy and Odefsey following Week 144 efficacy and safety data from Study GS-US-292-0112, listed as a category 4 study in the RMP; this is a phase 3 open-label safety study of elvitegravir/cobicistat/emtricitabine/tenofovir alafenamide single-tablet regimen in HIV-1 positive patients with mild to moderate renal impairment.

The Package Leaflet is updated accordingly.

In addition, the Worksharing applicant (WSA) took the opportunity make administrative updated to the Genvoya SmPC."

WS1156**Combivir-****EMEA/H/C/000190/WS1156/0090****Kivexa-EMEA/H/C/000581/WS1156/0072****Triumeq-****EMEA/H/C/002754/WS1156/0042****Trizivir-EMEA/H/C/000338/WS1156/0104**

MAH: ViiV Healthcare UK Limited, Lead

Rapporteur: Joseph Emmerich, "Update of section 4.5 of the SmPC to add information regarding the potential interaction between lamivudine and sorbitol based on the results of Study 204857. The Package Leaflet has been updated accordingly. Further, a minor amendment has been implemented throughout the SmPC in order to update the clinical terminology of *Pneumocystis carinii* pneumonia to *Pneumocystis jiroveci* pneumonia. In addition, the MAH takes the opportunity to make minor editorial changes , to align the annexes with the QRD template version 10 and to update the contact details of the local representative in Norway in the Package Leaflet."

WS1162**Glyxambi-****EMEA/H/C/003833/WS1162/0004****Jentadueto-****EMEA/H/C/002279/WS1162/0038****Trajenta-****EMEA/H/C/002110/WS1162/0028**

MAH: Boehringer Ingelheim International

GmbH, Lead Rapporteur: Johann Lodewijk

Hillegje, Lead PRAC Rapporteur: Menno van der Elst

B.6.10. CHMP-PRAC assessed procedures**Dificlir - fidaxomicin -****EMEA/H/C/002087/11/0028**

MAH: Astellas Pharma Europe B.V., Rapporteur:

Filip Josephson, PRAC Rapporteur: Qun-Ying

Yue, "C.I.11: Submission of an updated RMP

version 7 in order to remove the post-

authorization measure (PAM) MEA003

(concerning clinical study 2819-CL-2001 in

patients with Clostridium difficile Infection who

will receive a second course offidaxomicin) due

to the non-feasibility of the study."

Jinarc - tolvaptan -**EMEA/H/C/002788/11/0006**

MAH: Otsuka Pharmaceutical Europe Ltd,
Rapporteur: Greg Markey, PRAC Rapporteur:
Julie Williams, "Update of section 5.1 of the
SmPC based on final results from study 156-08-
271 (TEMPO 4:4) listed as a PAES in Annex II.
This study is a Multicenter, Open-label,
Extension Study (Extension of Trial 156-04-251)
to Evaluate the Long-term Safety and Efficacy of
Oral Tolvaptan Tablet Regimens in Patients With
Autosomal Dominant Polycystic. It provides data
for Jinarc treatment of autosomal dominant
polycystic kidney disease (ADPKD) over 5 years.
Reference to submission of this study is being
deleted from Annex II.
In addition, the Marketing authorisation holder
(MAH) took the opportunity to add the current
ATC code applicable for tolvaptan as it has been
assigned by WHO.
The RMP version 13.1 has also been submitted
to reflect the completion of the 156-08-271
study."

NINLARO - ixazomib -**EMEA/H/C/003844/11/0002, Orphan**

MAH: Takeda Pharma A/S, Rapporteur: Greg
Markey, PRAC Rapporteur: Ulla Wändel
Liminga"Update of sections 4.8 and 5.1 of the
SmPC to reflect the final overall survival
analysis of C16010 China continuation study, a
phase III study comparing ixazomib plus
lenalidomide and dexamethasone versus
placebo plus lenalidomide in patients with
relapsed and/or refractory multiple myeloma, in
order to fulfil SOB (Specific Obligation) 002.
Annex II.E and the RMP (version 2.0) are
updated accordingly. In addition the Marketing
Authorisation Holder (MAH) took the opportunity
to make a small correction in sections 4.7 and 9
of the SmPC and to the German translations."

OLYSIO - simeprevir -**EMEA/H/C/002777/11/0031**

MAH: Janssen-Cilag International NV,
Rapporteur: Aranzazu Sancho-Lopez, PRAC
Rapporteur: Julie WilliamsUpdate of section 5.1
of the SmPC in order to update the efficacy
information following results from study
HPC3002 A Prospective 3-year Follow-up Study
in Subjects Previously Treated in a Phase IIb or
Phase III Study with a TMC435-containing

Regimen for the Treatment of Hepatitis C Virus (HCV) Infection listed as a category 3 study in the RMP and in fulfilment of MEA005. The RMP version 4.0 has also been submitted which includes updates of changes already agreed in procedures
EMEA/H/C/002777/II/0021, EMEA/H/C/002777/I/0027 and EMEA/H/A-20/1438/C/2777/0019."

Pegasys - peginterferon alfa-2a -
EMEA/H/C/000395/II/0092
MAH: Roche Registration Limited, Rapporteur: Filip Josephson, PRAC Rapporteur: Qun-Ying Yue, "Submission of the final report from a systematic review and individual patient data meta-analysis of PEG-IFN studies to identify optimal stopping rules in order to provide the final outcome related to the assessment of a Response Guided Therapy (RGT) for Pegasys in HBV-infected patients."

TECFIDERA - dimethyl fumarate -
EMEA/H/C/002601/II/0036/G
MAH: Biogen Idec Ltd, Rapporteur: Martina Weise, PRAC Rapporteur: Martin Huber, Submission of a Clinical Study Report for study 109HV321: A Randomized, Double-Blind, Phase 3b Study to Evaluate the Safety and Tolerability of BG00012 when Administered as 240 mg BID (twice daily) Dose Regimen with and without Aspirin Compared to Placebo or Following a Slow Titration (Category 3)
C.I.13: Submission of a Clinical Study Report for study 109MS406 (ASSURE): A Phase 4, Randomized, Double-Blind Study with a Safety Extension Period to Evaluate the Effect of Aspirin on Flushing Events in Subjects with Relapsing-Remitting Multiple Sclerosis Treated with Tecfidera (Dimethyl Fumarate) Delayed-release Capsules (Category 4)"

TECFIDERA - dimethyl fumarate -
EMEA/H/C/002601/II/0037
MAH: Biogen Idec Ltd, Rapporteur: Martina Weise, PRAC Rapporteur: Martin Huber, C.I.4: Submission of a Clinical Study Report for study 109MS307: An Open-Label Study to Assess the Immune Response to Vaccination in Tecfidera-Treated Versus Interferon-Treated Subjects With Relapsing Forms of Multiple Sclerosis (Category 3). Consequently, this variation includes an update to section 4.5 (Interaction

with other medicinal products and other forms of interaction) of the Summary of Product Characteristics (SmPC) and section 2 of the package leaflet."

Yervoy - ipilimumab -

EMEA/H/C/002213/II/0042

MAH: Bristol-Myers Squibb Pharma EEIG,
Rapporteur: Paula Boudewina van Hennik, PRAC
Rapporteur: Sabine StrausUpdate of sections 4.4, 4.8 and 5.1 of the SmPC to reflect the final results of study CA184-169, a randomized double-blind phase III study of ipilimumab administered at 3 mg/kg versus at 10 mg/kg in subjects previously treated or untreated with unresectable or metastatic melanoma, in order to fulfil ANX 014.1. Annex II.D and the RMP (version 14.0) are updated accordingly. In addition the Marketing authorisation holder (MAH) took the opportunity to update the list of local representatives in the Package Leaflet, to include some editorial changes and correct some typos throughout the product information, and to bring the product information in line with the latest QRD template version 10."

Zinbryta - daclizumab -

EMEA/H/C/003862/II/0007

MAH: Biogen Idec Ltd, Rapporteur: Bruno Sepedes, PRAC Rapporteur: Eva A. Segovia"Update of sections 4.4 and 4.8 of the SmPC in order to add autoimmune haemolytic anaemia with frequency 'uncommon' and to include a warning concerning symptoms of this adverse drug reaction.
The Package Leaflet is updated accordingly.
In addition, the Marketing authorisation holder took the opportunity to implement minor editorial amendments throughout the Product Information.
The RMP version 5.0 has also been submitted."

WS1158/G

Humalog-

EMEA/H/C/000088/WS1158/0154/G

Liprolog-

EMEA/H/C/000393/WS1158/0117/G

MAH: Eli Lilly Nederland B.V., Lead Rapporteur: Robert James Hemmings, Lead PRAC
Rapporteur: Julie Williams, Type II (B.IV.1.c):
to add a pre-filled pen: the Humalog and
Liprolog 100 U/ml Junior KwikPen. The Junior

KwikPen can administer insulin in half unit increments and contains the insulin lispro 3ml cartridge that is already approved for use. The pack contains 5 pre-filled pens.

Type IA in (B.II.e.5.a.1): to add a new pack size of 10 (2x5) pre-filled pens (multipack) for the Humalog and Liprolog 100 U/ml Junior KwikPen. This presentation contains the insulin lispro 3ml cartridge that is already approved for use.
Type II (C.I.z): to change the SmPC of the already authorised 100 U/ml Humalog and Liprolog presentations: to change section 4.2 to include the paediatric population; to change section 4.4 to remove text that states that the product should only be used in children in preference to soluble insulin when a fast action of insulin might be beneficial. The PL is updated accordingly."

B.6.11. PRAC assessed procedures

PRAC Led

Avastin - bevacizumab -

EMEA/H/C/000582/11/0095

MAH: Roche Registration Limited, Rapporteur: Sinan B. Sarac, PRAC Rapporteur: Doris Stenver, Submission of an updated RMP version 28.0 in order to remove the post-authorisation measure outlined in section III.4.3 of the RMP consisting of the submission of an extension protocol in order to obtain additional long-term follow-up (LTFU) information from the paediatric population after patients complete the minimum 5.5 year follow-up period as defined in the BO20924 (BERNIE) paediatric study protocol and to amend the date of submission of the final report (addendum CSR) for the BO20924 (BERNIE) study."

PRAC Led

Benlysta - belimumab -

EMEA/H/C/002015/11/0049

MAH: Glaxo Group Ltd, Rapporteur: Kristina Dunder, PRAC Rapporteur: Ulla Wändel Liminga, Submission of an updated RMP version 23 in order to amend the CSR available time line, patient number and the primary and secondary endpoints listed in the EU Risk Management Plan, with regards to study HGS1006-C1121/BEL114054."

PRAC Led

Inovelon - rufinamide -

EMEA/H/C/000660/11/0041, Orphan

MAH: Eisai Ltd, Rapporteur: Alexandre Moreau, PRAC Rapporteur: Claire Ferard "Submission of the final clinical study report for study E2080-E044-401, the European registry of anti-epileptic drug use in patients with Lennox-Gastaut Syndrome (LAG), listed as a category 3 study in the RMP, in order to fulfil MEA 002.1. This is a non-interventional EU registry study entering patients (aged ≥4 years) with LGS who required a modification in anti-epileptic therapy (either the addition of another AED or the change of one drug to another) to evaluate the long-term safety of rufinamide."

PRAC Led

Plenadren - hydrocortisone -

EMEA/H/C/002185/11/0024, Orphan

MAH: Shire Services BVBA, Rapporteur: Kristina Dunder, PRAC Rapporteur: Qun-Ying Yue, PRAC- "Submission of an updated RMP (version 3.1) in order to submit protocol amendments of SHP 617-400 (EU-AIR) study – A European multicentre, multi-country, post-authorisation, observation study (registry) of patients with chronic adrenal insufficiency (category 3). Additionally, the opportunity is being taken to implement a change agreed by the PRAC/CHMP as part of the assessment of MEA 005.3 in July 2016 and remove from the RMP reference to study SHP617-404 (SWE-DUS), a Category 3 study to monitor off-label use of Plenadren to evaluate physician prescribing patterns."

B.6.12. CHMP-CAT assessed procedures

Holoclar - ex vivo expanded autologous human corneal epithelial cells containing stem cells -

**EMEA/H/C/002450/11/0012/G, Orphan,
ATMP**

MAH: Chiesi Farmaceutici S.p.A.

B.6.13. CHMP-PRAC-CAT assessed procedures

B.6.14. PRAC assessed ATMP procedures

B.6.15. Unclassified procedures and worksharing procedures of type I variations

WS1115/G

Ambirix-

EMEA/H/C/000426/WS1115/0084/G

Twinrix Adult-

EMEA/H/C/000112/WS1115/0118/G

Twinrix Paediatric-

EMEA/H/C/000129/WS1115/0119/G

MAH: GSK Biologicals SA, Lead Rapporteur:

Robert James Hemmings

WS1116

Infanrix hexa-

EMEA/H/C/000296/WS1116/0217

MAH: GSK Biologicals SA, Lead Rapporteur: Bart

Van der Schueren

WS1122/G

Hexacima-

EMEA/H/C/002702/WS1122/0060/G

Hexaxim-

EMEA/H/W/002495/WS1122/0066/G

Hexyon-

EMEA/H/C/002796/WS1122/0064/G

MAH: Sanofi Pasteur SA, Lead Rapporteur: Jan

Mueller-Berghaus

WS1129/G

Hexacima-

EMEA/H/C/002702/WS1129/0061/G

Hexaxim-

EMEA/H/W/002495/WS1129/0067/G

Hexyon-

EMEA/H/C/002796/WS1129/0065/G

MAH: Sanofi Pasteur SA, Lead Rapporteur: Jan

Mueller-Berghaus

WS1138

Actos-EMEA/H/C/000285/WS1138/0077

Competact-

EMEA/H/C/000655/WS1138/0065

Glubrava-

EMEA/H/C/000893/WS1138/0051

Glustin-EMEA/H/C/000286/WS1138/0076

Tandemact-

EMEA/H/C/000680/WS1138/0055

MAH: Takeda Pharma A/S, Lead Rapporteur:

Patrick Salmon

WS1139/G**Rivastigmine 1A Pharma-****EMEA/H/C/001181/WS1139/0023/G****Rivastigmine Hexal-****EMEA/H/C/001182/WS1139/0024/G****Rivastigmine Sandoz-****EMEA/H/C/001183/WS1139/0025/G**

MAH: 1 A Pharma GmbH, Informed Consent of

Exelon, Lead Rapporteur: Alexandre Moreau

WS1157**Relvar Ellipta-****EMEA/H/C/002673/WS1157/0030****Revinty Ellipta-****EMEA/H/C/002745/WS1157/0026**

MAH: Glaxo Group Ltd, Lead Rapporteur:

Concepcion Prieto Yerro

B.7. DOCUMENTS TABLED IN MMD AFTER THE CHMP PLENARY**B.7.1. Line listing for Variation Type I and Variation Type II (MMD only) post authorisation procedures from the beginning of the year.****B.7.2. Line listing overview of all applications under the centralised procedure (MMD only). line listing - products - authorised, under evaluation, suspended.xls****B.7.3. Opinion on Marketing Authorisation transfer (MMD only).****B.7.4. Notifications in accordance with Article 61(3) of Council Directive 2001/83/EC (MMD only).****B.7.5. Request for supplementary information relating to Notification of Type I variation (MMD only).****B.7.6. Notifications of Type I Variations (MMD only).****C. Annex C - Post-Authorisation Measures (PAMs), (Line listing of Post authorisation measures with a description of the PAM. Procedures starting in that given month with assessment timetabled)****D. Annex D - Post-Authorisation Measures (PAMs), (Details on PAMs including description and conclusion, for adoption by CHMP in that given month, or finalised ones with PRAC recommendation and no adoption by CHMP needed)****E. Annex E - EMEA CERTIFICATION OF PLASMA MASTER FILES**

Disclosure of information related to plasma master files cannot be released at present time as these contain commercially confidential information.

E.1. PMF Certification Dossiers:

E.1.1. Annual Update

E.1.2. Variations:

E.1.3. Initial PMF Certification:

E.2. Time Tables – starting & ongoing procedures: For information

PMF timetables starting and ongoing procedures Tabled in MMD and sent by post mail (folder E).

F. ANNEX F - Decision of the Granting of a Fee Reduction/Fee Waiver

F.1. Parallel Distribution - Pursuant to Article 9 of Council Regulation (EC) No. 2743/98 of 14 December 1998, as amended

F.2. Request for scientific opinion on justification of exceptional circumstance and for imperative grounds of public health

G. ANNEX G

G.1. Final Scientific Advice (Reports and Scientific Advice letters):

Disclosure of information related to Scientific Advice cannot be released at present time as these contain commercially confidential information.

I Final Scientific Advice (Reports and Scientific Advice letters):

VII Post-Scientific Advice Issues:

Qualification of Biomarkers:

HTA:

G.2. Ongoing procedures

G.3. PRIME

Disclosure of some information related to PRIME cannot be released at present time as these contain commercially confidential information.

G.3.1. List of procedures concluding at 20-23 March 2017 CHMP plenary:

G.3.2. List of procedures starting in March 2017 for April 2017 CHMP adoption of outcomes

H. ANNEX H - Product Shared Mailboxes – e-mail address