

24 July 2025
EMA/CHMP/319030/2025

Committee for medicinal products for human use (CHMP)

Minutes for the meeting on 21-24 July 2025

Chair: Bruno Sepedes – Vice-Chair: Outi Mäki-Ikola

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 set of minutes 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, these minutes 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 minutes 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

The Chairperson opened the meeting by welcoming all participants. The meeting was held in person.

In accordance with the Agency's policy on handling of declarations of interests of scientific Committees' members and experts, based on the declarations of interest submitted by the Committee members, alternates and experts and based on the topics in the agenda of the meeting, the Committee Secretariat announced the restricted involvement of some Committee members, alternates and experts for concerned agenda topics.

Participants were asked to declare any changes, omissions or errors to their declared interests and/or additional restrictions concerning the matters for discussion. No new or additional interests or restrictions were declared. Restrictions applicable to this meeting are captured in the list of participants included in the minutes.

Discussions, deliberations and voting took place in full respect of the restricted involvement of Committee members and experts in line with the relevant provisions of the Rules of Procedure. All decisions, recommendations and advice were agreed by consensus, unless otherwise specified.

1.2. Adoption of agenda

CHMP agenda for 21-24 July 2025.

The CHMP adopted the agenda.

1.3. Adoption of the minutes

CHMP minutes for the 22-25 April 2025 and the 19-22 May 2025 meetings.

The CHMP adopted the minutes for the 22-25 April 2025 and the 19-22 May 2025 plenary meetings.

Minutes from Preparatory and Organisational Matters (PROM) meeting held on 14 July 2025.

The CHMP adopted the minutes from the PROM meeting held on 14 July 2025.

2. Oral Explanations

2.1. Pre-authorisation procedure oral explanations

2.1.1. Pridopidine - Orphan - EMEA/H/C/006261

Prilenia Therapeutics B.V.; treatment of Huntington's disease

Scope: Oral explanation

Action: Oral explanation to be held on 22 July 2025 at 11:00

List of Outstanding Issues adopted on 22.05.2025. List of Questions adopted on 12.12.2024.

Patient representatives

An oral explanation was held on 22 July 2025. The presentation by the applicant focused on the clinical data in support of the application.

3.1

2.2. Re-examination procedure oral explanations

2.2.1. Kisunla - Donanemab - EMEA/H/C/006024

Eli Lilly Nederland B.V.; to slow disease progression in adult patients with Alzheimer's disease (AD).

Scope: Oral explanation

Action: Oral explanation to be held on 21 July 2025 at 14:00

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

Opinion adopted on 27.03.2025. List of Outstanding Issues adopted on 12.12.2024, 25.04.2024. List of Questions adopted on 14.12.2023.

An oral explanation was held on 21 July 2025. The presentation by the applicant focused on the clinical data in support of the application.

See 3.5

2.2.2. Aplidin - plitidepsin - Orphan - EMEA/H/C/004354

Pharma Mar, S.A.; treatment of multiple myeloma

Scope: Oral explanation

Action: Oral explanation to be held on 23 July 2025 at 11:00

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

An oral explanation was held on 23 July 2025. The presentation by the applicant focused on the clinical data in support of the application.

See 3.7

2.3. Post-authorisation procedure oral explanations

No items

2.4. Referral procedure oral explanations

No items

3. Initial applications

3.1. Initial applications; Opinions

3.1.1. Aqneursa - L-Acetylleucine - Orphan - EMEA/H/C/006327

Intrabio Ireland Limited; chronic treatment of Niemann-Pick Type C (NPC) in adults and children from birth

Scope: Opinion

Action: For adoption

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

List of Outstanding Issues adopted on 25.04.2025, 27.02.2025. List of Questions adopted on 17.10.2024.

The Committee confirmed that all issues previously identified in this application had been addressed.

The Committee adopted a positive opinion recommending the granting of a marketing authorisation by consensus together with the CHMP assessment report and translation timetable.

The legal status was agreed as medicinal product subject to medical prescription.

The CHMP noted the letter of recommendation dated 23 July 2025.

The summary of opinion was circulated for information.

The CHMP adopted the similarity assessment report

The EMA public health communication was circulated for information.

3.1.2. BILDYOS - Denosumab - EMEA/H/C/006434

Henlius Europe GmbH; treatment of osteoporosis and bone loss

Scope: Opinion

Action: For adoption

Similar biological application (Article 10(4) of Directive No 2001/83/EC)

List of Outstanding Issues adopted on 27.02.2025. List of Questions adopted on 19.09.2024.

The Committee confirmed that all issues previously identified in this application had been addressed.

The Committee adopted a positive opinion recommending the granting of a marketing authorisation by consensus together with the CHMP assessment report and translation timetable.

The legal status was agreed as medicinal product subject to medical prescription.

The summary of opinion was circulated for information.

3.1.3. BILPREVDA - Denosumab - EMEA/H/C/006435

Henlius Europe GmbH; prevention of skeletal related events in adults with advanced malignancies involving bone

Scope: Opinion

Action: For adoption

Similar biological application (Article 10(4) of Directive No 2001/83/EC)

List of Outstanding Issues adopted on 27.02.2025. List of Questions adopted on 19.09.2024.

The Committee confirmed that all issues previously identified in this application had been addressed.

The Committee adopted a positive opinion recommending the granting of a marketing authorisation by consensus together with the CHMP assessment report and translation timetable.

The legal status was agreed as medicinal product subject to medical prescription.

The summary of opinion was circulated for information.

3.1.4. Ektetyl - Sebetalstat - Orphan - EMEA/H/C/006211

Kalvista Pharmaceuticals (Ireland) Limited; treatment of hereditary angioedema (HAE) attacks in adult and adolescents aged 12 years and older

Scope: Opinion

Action: For adoption

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

List of Outstanding Issues adopted on 22.05.2025. List of Questions adopted on 12.12.2024.

The Committee confirmed that all issues previously identified in this application had been addressed.

The Committee adopted a positive opinion recommending the granting of a marketing authorisation by consensus together with the CHMP assessment report and translation timetable.

Furthermore, the CHMP considered that sebetalstat is a new active substance, as claimed by the applicant.

The legal status was agreed as medicinal product subject to medical prescription.

The summary of opinion was circulated for information.

The CHMP adopted the similarity assessment report

3.1.5. Elevidys - Delandistrogene moxeparvovec - Orphan - ATMP - EMEA/H/C/005293

Roche Registration GmbH; treatment of ambulatory patients aged 3 to 7 years old with Duchenne muscular dystrophy

Scope: Opinion

Action: For adoption

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

List of Outstanding Issues adopted on 16.04.2025. List of Questions adopted on 11.10.2024.

Based on the draft opinion prepared by the CAT, the Committee adopted a negative opinion by consensus recommending the refusal of the granting of the marketing authorisation. The CHMP assessment report was adopted.

The refusal question-and-answer document was circulated for information.

3.1.6. Eylexvi - Aflibercept - EMEA/H/C/006282

biolitec pharma Limited Zweigniederlassung Jena; treatment of age-related macular degeneration (AMD) and visual impairment

Scope: Opinion

Action: For adoption

Similar biological application (Article 10(4) of Directive No 2001/83/EC)

List of Outstanding Issues adopted on 25.04.2025. List of Questions adopted on 14.11.2024.

The Committee confirmed that all issues previously identified in this application had been addressed.

The Committee adopted a positive opinion recommending the granting of a marketing authorisation by consensus together with the CHMP assessment report and translation timetable.

The legal status was agreed as medicinal product subject to restricted medical prescription.

The summary of opinion was circulated for information.

3.1.7. JELRIX - Autologous cartilage-derived articular chondrocytes, in-vitro expanded - ATMP - EMEA/H/C/004594

TETEC Tissue Engineering Technologies AG; repair of symptomatic, localised, full-thickness

cartilage defects of the knee joint grade III or IV

Scope: Opinion

Action: For adoption

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

List of Outstanding Issues adopted on 13.06.2025, 21.03.2025. List of Questions adopted on 19.04.2024.

Based on the draft opinion prepared by the CAT, the Committee adopted a negative opinion by majority recommending the refusal of the granting of the marketing authorisation. The CHMP assessment report was adopted.

The divergent position was appended to the opinion.

The refusal question-and-answer document was circulated for information.

The CHMP noted the re-examination request by the applicant.

3.1.8. Yeytuo - Lenacapavir - EMEA/H/C/006658

Accelerated assessment

Gilead Sciences Ireland Unlimited Company; pre-exposure prophylaxis to prevent HIV-1

Scope: Opinion

Action: For adoption

Known active substance (Article 8(3) of Directive No 2001/83/EC)

List of Questions adopted on 20.05.2025.

The Committee confirmed that all issues previously identified in this application had been addressed.

The Committee adopted a positive opinion recommending the granting of a marketing authorisation by consensus together with the CHMP assessment report and translation timetable.

The legal status was agreed as medicinal product subject to restricted medical prescription.

The summary of opinion was circulated for information.

The EMA public health communication was circulated for information.

3.1.9. Lenacapavir Gilead - Lenacapavir - Article 58 - EMEA/H/W/006659

Accelerated assessment

Gilead Sciences Ireland Unlimited Company; pre-exposure prophylaxis to prevent HIV-1

Scope: Opinion

Action: For adoption

Known active substance (Article 8(3) of Directive No 2001/83/EC)

The Committee confirmed that all issues previously identified in this application had been addressed.

The Committee adopted the scientific opinion for lenacapavir in accordance with Article 58 of Regulation (EC) No. 726/2004.

The legal status was agreed as medicinal product subject to restricted medical prescription.

The summary of opinion was circulated for information.

3.1.10. Macitentan Accord - Macitentan - EMEA/H/C/006524

Accord Healthcare; treatment of pulmonary arterial hypertension (PAH)

Scope: Opinion

Action: For adoption

Generic application (Article 10(1) of Directive No 2001/83/EC), Generic of Opsumit

List of Outstanding Issues adopted on 25.04.2025. List of Questions adopted on 19.09.2024.

The Committee confirmed that all issues previously identified in this application had been addressed.

The Committee adopted a positive opinion recommending the granting of a marketing authorisation by consensus together with the CHMP assessment report and translation timetable.

The legal status was agreed as medicinal product subject to restricted medical prescription.

The summary of opinion was circulated for information.

The CHMP adopted the similarity assessment report

3.1.11. Macitentan AccordPharma - Macitentan - EMEA/H/C/006523

Accord Healthcare; treatment of pulmonary arterial hypertension (PAH)

Scope: Opinion

Action: For adoption

Generic application (Article 10(1) of Directive No 2001/83/EC), Generic of Opsumit

List of Outstanding Issues adopted on 25.04.2025. List of Questions adopted on 19.09.2024.

The Committee confirmed that all issues previously identified in this application had been addressed.

The Committee adopted a positive opinion recommending the granting of a marketing authorisation by consensus together with the CHMP assessment report and translation timetable.

The legal status was agreed as medicinal product subject to restricted medical prescription.

The summary of opinion was circulated for information.

The CHMP adopted the similarity assessment report

3.1.12. Nurzigma - Pridopidine - Orphan - EMEA/H/C/006261

Prilenia Therapeutics B.V.; treatment of Huntington's disease

Scope: Opinion

Action: For adoption

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

List of Outstanding Issues adopted on 22.05.2025. List of Questions adopted on 12.12.2024.

Patient representatives

The Committee adopted a negative opinion by consensus recommending the refusal of the granting of the conditional marketing authorisation. The CHMP assessment report was adopted.

The question-and-answer document was circulated for information.

See 2.1

3.1.13. ROMVIMZA - Vimsetinib - Orphan - EMEA/H/C/006363

Deciphera Pharmaceuticals (Netherlands) B.V.; Treatment of adult patients with tenosynovial giant cell tumour (TGCT) who are not amenable to surgery

Scope: Opinion

Action: For adoption

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

List of Outstanding Issues adopted on 25.04.2025. List of Questions adopted on 14.11.2024.

The Committee confirmed that all issues previously identified in this application had been addressed.

The Committee adopted a positive opinion recommending the granting of a marketing authorisation by consensus together with the CHMP assessment report and translation timetable.

Furthermore, the CHMP considered that Vimsetinib is a new active substance, as claimed by the applicant.

The legal status was agreed as medicinal product subject to restricted medical prescription.

The summary of opinion was circulated for information.

3.1.14. Tryngolza - Olezarsen - Orphan - EMEA/H/C/006477

Ionis Ireland Limited; treatment of familial chylomicronemia syndrome

Scope: Opinion

Action: For adoption

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

List of Outstanding Issues adopted on 22.05.2025. List of Questions adopted on 12.12.2024.

The Committee confirmed that all issues previously identified in this application had been addressed.

The Committee adopted a positive opinion recommending the granting of a marketing authorisation by consensus together with the CHMP assessment report and translation timetable.

Furthermore, the CHMP considered that Olezarsen is a new active substance, as claimed by the applicant.

The legal status was agreed as medicinal product subject to medical prescription.

The CHMP noted the letter of recommendation dated 23 July 2025.

The summary of opinion was circulated for information.

The CHMP adopted the similarity assessment report.

3.1.15. Usrenty - Ustekinumab - EMEA/H/C/006794

Biosimilar Collaborations Ireland Limited; treatment of Crohn's Disease, treatment of Plaque psoriasis, Psoriatic arthritis (PsA)

Scope: Opinion

Action: For adoption

Similar biological application (Article 10(4) of Directive No 2001/83/EC)

The Committee confirmed that all issues previously identified in this application had been addressed.

The Committee adopted a positive opinion recommending the granting of a marketing authorisation by consensus together with the CHMP assessment report and translation timetable.

The legal status was agreed as medicinal product subject to restricted medical prescription.

The summary of opinion was circulated for information.

3.1.16. Voranigo - Vorasidenib - Orphan - EMEA/H/C/006284

Les Laboratoires Servier; treatment of predominantly non-enhancing astrocytoma or oligodendrolioma

Scope: Opinion

Action: For adoption

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

List of Outstanding Issues adopted on 30.01.2025, 19.09.2024, 25.07.2024. List of Questions adopted on 23.04.2024.

The Committee confirmed that all issues previously identified in this application had been addressed.

The Committee adopted a positive opinion recommending the granting of a marketing authorisation by consensus together with the CHMP assessment report and translation timetable.

Furthermore, the CHMP considered that Vorasidenib is a new active substance, as claimed by the applicant.

The legal status was agreed as medicinal product subject to restricted medical prescription.

The CHMP noted the letter of recommendation dated 21 July 2025.

The summary of opinion was circulated for information.

The CHMP adopted the similarity assessment report.

3.1.17. [Zurzuvae - Zuranolone - EMEA/H/C/006488](#)

Biogen Netherlands B.V.; treatment of postpartum depression (PPD) in adults

Scope: Opinion

Action: For adoption

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

List of Outstanding Issues adopted on 22.05.2025. List of Questions adopted on 12.12.2024.

The Committee confirmed that all issues previously identified in this application had been addressed.

The Committee adopted a positive opinion recommending the granting of a marketing authorisation by consensus together with the CHMP assessment report and translation timetable.

Furthermore, the CHMP considered that Zuranolone is a new active substance, as claimed by the applicant.

The legal status was agreed as medicinal product subject to medical prescription.

The CHMP noted the letter of recommendation dated 18 July 2025.

The summary of opinion was circulated for information.

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

3.2.1. [Clesrovimab - EMEA/H/C/006497](#)

prevention of infections with respiratory syncytial virus (RSV) and lower respiratory tract disease (LRTD)

Scope: List of outstanding issues

Action: For adoption

List of Questions adopted on 27.03.2025.

The Committee was reminded of the status of this application and its remaining outstanding

issues.

The Committee adopted a list of outstanding issues with a specific timetable.

3.2.2. Denosumab - EMEA/H/C/006239

prevention of skeletal related events in adults with advanced malignancies involving bone

Scope: List of outstanding issues

Action: For adoption

List of Questions adopted on 27.03.2025.

The Committee was reminded of the status of this application and its remaining outstanding issues.

The Committee adopted a list of outstanding issues with a specific timetable.

3.2.3. Enzalutamide - EMEA/H/C/006612

treatment of prostate cancer

Scope: List of outstanding issues

Action: For adoption

List of Questions adopted on 27.02.2025.

The Committee was reminded of the status of this application and its remaining outstanding issues.

The Committee adopted a list of outstanding issues with a specific timetable.

3.2.4. Insulin icodex / Semaglutide - EMEA/H/C/006279

treatment of adults with type 2 diabetes mellitus insufficiently controlled on basal insulin or glucagon-like peptide 1 (GLP-1) receptor agonists

Scope: List of outstanding issues; Request by the applicant for an extension to the clock-stop to respond to the list of questions adopted in February 2025.

Action: For adoption

List of Questions adopted on 27.02.2025.

The Committee was reminded of the status of this application and its remaining outstanding issues.

The Committee adopted a list of outstanding issues with a specific timetable.

The Committee denied the request by the applicant for an extension to the clock-stop to respond to the list of questions adopted in February 2025.

3.2.5. Elinzanetant - EMEA/H/C/006298

for the treatment of moderate to severe vasomotor symptoms (VMS)

Scope: List of outstanding issues

Action: For adoption

List of Questions adopted on 27.02.2025.

The Committee was reminded of the status of this application and its remaining outstanding issues.

The Committee adopted a list of outstanding issues with a specific timetable.

3.2.6. ACELLULAR PERTUSSIS VACCINE - EMEA/H/C/006304

indicated as active booster immunization against *pertussis* of persons aged 11 years onwards and passive protection against *pertussis* in early infancy following maternal immunization during pregnancy

Scope: List of outstanding issues

Action: For adoption

List of Questions adopted on 14.11.2024.

The Committee was reminded of the status of this application and its remaining outstanding issues.

The Committee adopted a list of outstanding issues with a specific timetable.

3.2.7. Rivaroxaban - EMEA/H/C/006643

prevention of atherothrombotic events

Scope: List of outstanding issues; Request by the applicant for an extension to the clock-stop to respond to the list of questions adopted in February 2025.

Action: For adoption

List of Questions adopted on 27.02.2025.

The Committee was reminded of the status of this application and its remaining outstanding issues.

The Committee adopted a list of outstanding issues with a specific timetable.

The Committee denied the request by the applicant for an extension to the clock-stop to respond to the list of questions adopted in February 2025.

3.2.8. Teduglutide - EMEA/H/C/006564

treatment of Short Bowel Syndrome

Scope: List of outstanding issues

Action: For adoption

List of Questions adopted on 27.02.2025.

The Committee was reminded of the status of this application and its remaining outstanding issues.

The Committee adopted a list of outstanding issues with a specific timetable.

3.2.9. Rilzabrutinib - Orphan - EMEA/H/C/006425

Sanofi B.V.; for the treatment of persistent or chronic immune thrombocytopenia (ITP)

Scope: List of outstanding issues

Action: For adoption

List of Questions adopted on 27.02.2025.

The Committee was reminded of the status of this application and its remaining outstanding issues.

The Committee adopted a list of outstanding issues with a specific timetable.

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

3.3.1. Copper (64Cu) oxodotreotide - Orphan - EMEA/H/C/006608

Cis Bio International; positron emission tomography (PET) for localization of somatostatin receptor positive neuroendocrine neoplasms (NENs).

Scope: List of questions

Action: For adoption

The Committee discussed the issues identified in this application.

The Committee adopted the CHMP recommendation and scientific discussion together with the list of questions.

3.3.2. Liraglutide - EMEA/H/C/006620

treatment of diabetes and weight management

Scope: List of questions; Request by the applicant for an extension to the clock-stop to respond to the list of questions to be adopted in July 2025.

Action: For adoption

The Committee discussed the issues identified in this application.

The Committee adopted the CHMP recommendation and scientific discussion together with the list of questions.

The Committee denied the request by the applicant for an extension to the clock-stop to respond to the list of questions adopted in July 2025.

3.3.3. Furosemide - PUMA - EMEA/H/C/006617

treatment of all conditions requiring diuresis due to mechanical obstruction or venous insufficiency.

Scope: List of questions

Action: For adoption

The Committee discussed the issues identified in this application.

The Committee adopted the CHMP recommendation and scientific discussion together with the list of questions.

3.3.4. Brensocatib - PRIME - EMEA/H/C/005820

Accelerated assessment

treatment of non-cystic fibrosis bronchiectasis

Scope: List of questions

Action: For adoption

The Committee discussed the issues identified in this application.

The Committee adopted the CHMP recommendation and scientific discussion together with the list of questions.

3.3.5. Apitegromab - PRIME - Orphan - EMEA/H/C/005909

Scholar Rock Netherlands B.V.; treatment of 5q spinal muscular atrophy (SMA)

Scope: List of questions; Request by the applicant for an extension to the clock-stop to respond to the list of questions to be adopted in July 2025.

Action: For adoption

The Committee discussed the issues identified in this application.

The Committee adopted the CHMP recommendation and scientific discussion together with the list of questions

The Committee denied the request by the applicant for an extension to the clock-stop to respond to the list of questions adopted in July 2025.

3.3.6. Semaglutide - EMEA/H/C/006426

treatment of non-cirrhotic metabolic dysfunction-associated steatohepatitis with liver fibrosis

Scope: List of questions

Action: For adoption

The Committee discussed the issues identified in this application.

The Committee adopted the CHMP recommendation and scientific discussion together with the list of questions.

3.3.7. Liraglutide - EMEA/H/C/006615

treatment of adults, adolescents and children aged 10 years and above with insufficiently

controlled type 2 diabetes mellitus as an adjunct to diet and exercise

Scope: List of questions; Request by the applicant for an extension to the clock-stop to respond to the list of questions to be adopted in July 2025.

Action: For adoption

The Committee discussed the issues identified in this application

The Committee adopted the CHMP recommendation and scientific discussion together with the list of questions.

The Committee denied the request by the applicant for an extension to the clock-stop to respond to the list of questions adopted in July 2025.

3.3.8. Tovorafenib - Orphan - EMEA/H/C/006140

Ipsen Pharma; treatment of paediatric low-grade glioma (LGG)

Scope: List of questions

Action: For adoption

The Committee discussed the issues identified in this application.

The Committee adopted the CHMP recommendation and scientific discussion together with the list of questions.

3.3.9. Paltusotine - Orphan - EMEA/H/C/006636

Voisin Consulting Life Sciences; Maintenance treatment in adult patients with acromegaly

Scope: List of questions

Action: For adoption

The Committee discussed the issues identified in this application

The Committee adopted the CHMP recommendation and scientific discussion together with the list of questions.

3.3.10. Pertuzumab - EMEA/H/C/006583

treatment of breast cancer

Scope: List of questions

Action: For adoption

The Committee discussed the issues identified in this application.

The Committee adopted the CHMP recommendation and scientific discussion together with the list of questions.

3.3.11. Remibrutinib - EMEA/H/C/006313

treatment of chronic spontaneous urticaria in patients with inadequate response to H1 antihistamine

Scope: List of questions

Action: For adoption

The Committee discussed the issues identified in this application.

The Committee adopted the CHMP recommendation and scientific discussion together with the list of questions.

[3.3.12. INFLUENZA VACCINE - EMEA/H/C/006674](#)

immunisation for the prevention of influenza disease

Scope: List of questions

Action: For adoption

The Committee discussed the issues identified in this application.

The Committee adopted the CHMP recommendation and scientific discussion together with the list of questions.

[3.3.13. Autologous melanoma-derived tumour infiltrating lymphocytes, ex vivo-expanded - ATMP - EMEA/H/C/006563](#)

treatment of melanoma

Scope: List of questions; Request by the applicant for an extension to the clock-stop to respond to the list of questions to be adopted in July 2025.

Action: For information

The CHMP was updated on discussions at the CAT. The Committee was reminded of the status of this application and its remaining outstanding issues.

The Committee endorsed a list of questions with a specific timetable, as adopted by CAT.

The Committee denied the request by the applicant for an extension to the clock-stop to respond to the list of questions adopted in July 2025.

[3.3.14. Tocilizumab - EMEA/H/C/006416](#)

treatment of rheumatoid arthritis and other immunological conditions

Scope: List of questions

Action: For adoption

The Committee discussed the issues identified in this application.

The Committee adopted the CHMP recommendation and scientific discussion together with the list of questions.

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

3.4.1. Belumosudil - Orphan - EMEA/H/C/006421

Sanofi Winthrop Industrie; Treatment of chronic graft-versus host disease (cGVHD) after failure of at least two prior lines of systemic therapy.

Scope: Update on the procedure

Request by the applicant for an extension to the clock-stop to respond to the list of outstanding issues adopted in June 2025.

Action: For information

List of Outstanding Issues adopted on 19.06.2025. List of Questions adopted on 30.01.2025.

The CHMP noted the update on the procedure and denied the request by the applicant for an extension to the clock-stop to respond to the list of outstanding issues adopted in June 2025.

3.4.2. Golimumab - EMEA/H/C/006621

treatment of rheumatoid arthritis, psoriatic arthritis and ankylosing spondylitis

Scope: Request by the applicant for an extension to the clock-stop to respond to the list of questions adopted in May 2025.

Action: For adoption

List of questions adopted on 22.05.2025

The Committee denied the request by the applicant for an extension to the clock-stop to respond to the list of questions adopted in May 2025.

3.4.3. Mavorixafor - Orphan - EMEA/H/C/006496

Treatment of WHIM syndrome

Scope: Request by the applicant for an extension to the clock-stop to respond to the list of questions adopted in May 2025.

Action: For adoption

List of questions adopted on 22.05.2025

The Committee agreed to the request by the applicant for an extension to the clock-stop to respond to the list of questions adopted in May 2025.

3.4.4. Plozasiran - Orphan - EMEA/H/C/006579

Accelerated assessment

Arrowhead Pharmaceuticals Ireland Limited; treatment of familial chylomicronaemia syndrome (FCS).

Scope: Request by the applicant for an extension to the clock-stop to respond to the list of

questions adopted in June 2025.

Action: For adoption

List of questions 19.06.2025

The Committee agreed to the request by the applicant for an extension to the clock-stop to respond to the list of questions adopted in June 2025.

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

3.5.1. Atropine sulfate FGK - Atropine - PUMA - EMEA/H/C/006385

FGK Representative Service GmbH; treatment of myopia in children aged 3 years and older

Scope: Adoption of timetable, questions to the AHEG

Action: For adoption

Known active substance (Article 8(3) of Directive No 2001/83/EC)

Opinion adopted on 22.05.2025. List of Outstanding Issues adopted on 27.02.2025. List of Questions adopted on 19.09.2024.

The CHMP adopted the timetable and the questions to the AHEG.

3.5.2. Austedo - Deutetrabenazine - EMEA/H/C/006371

Teva GmbH; treatment of tardive dyskinesia

Scope: Request for re-examination, appointment of re-examination rapporteurs

Action: For adoption

Opinion adopted on 19.06.2025. List of Outstanding Issues adopted on 27.02.2025. List of Questions adopted on 25.07.2024.

The CHMP noted the request for re-examination and appointed re-examination rapporteur and re-examination co-rapporteur.

3.5.3. Kisunla - Donanemab - EMEA/H/C/006024

Eli Lilly Nederland B.V.; to slow disease progression in adult patients with Alzheimer's disease (AD).

Scope: Opinion

Action: For adoption

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

Third party intervention

Opinion adopted on 27.03.2025. List of Outstanding Issues adopted on 12.12.2024, 25.04.2024. List of Questions adopted on 14.12.2023.

An oral explanation was held on 21 July 2025. The presentation by the applicant focused on

the clinical data in support of the application.

The Committee adopted a positive opinion recommending the granting of a marketing authorisation by majority (17 out of 32) together with the CHMP assessment report and translation timetable.

The divergent opinion (Daniela Philadelphy, Thalia Marie Blicher, Alexandre Moreau, Antonio Gómez Outes, Sol Ruiz, Emilia Mavrokordatou, Frantisek Drafi, Beata Maria Jakline Ullrich, Aris Angelis, Blanka Hirschlerová, Tomáš Radiměřský, Bruno Delafont, Peter Mol, Jayne Crowe, Outi Mäki-Ikola) was appended to the opinion.

The legal status was agreed as medicinal product subject to restricted medical prescription.

The summary of opinion was circulated for information.

The EMA public health communication was circulated for information.

See 2.2

3.6. Initial applications in the decision-making phase

No items

3.7. Withdrawals of initial marketing authorisation application

3.7.1. Lifileucel - ATMP - EMEA/H/C/004741

treatment of unresectable or metastatic melanoma

Scope: Withdrawal of initial marketing authorisation application

Action: For information

List of Outstanding Issues adopted on 16.05.2025. List of Questions adopted on 06.12.2024.

The CHMP noted the withdrawal of the initial marketing authorisation application.

3.7.2. Aplidin - plitidepsin - Orphan - EMEA/H/C/004354

Pharma Mar, S.A.; treatment of multiple myeloma

Scope: Opinion

Restart the 2018 re-examination procedure relating to the initial marketing authorisation application for Aplidin following the adoption of Commission Implementing Decision C(2024) 4469 final of 28 June 2024 which revoked Commission Implementing Decision C(2018) 4831 final of 17 July 2018 refusing marketing authorisation for 'Aplidin – plitidepsin'. That decision was revoked following the judgment of 14 March 2024 in D & A Pharma v Commission and EMA, C 291/22 P.

Action: For adoption

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

An oral explanation was held on 23 July 2025. The presentation by the applicant focused on

the clinical data in support of the application.

The MAH withdrew the re-examination application.

See 2.2

3.7.3. Bifikafusp alfa / Onfekafusp alfa - EMEA/H/C/005651

neoadjuvant treatment of adult patients with locally advanced fully resectable melanoma

Scope: Withdrawal of initial marketing authorisation application

Action: For information

The CHMP noted the withdrawal of the initial marketing authorisation application.

3.7.4. Eflornithine – EMEA/H/C/006067

treatment of high-risk neuroblastoma responsive to prior multiagent, multimodality therapy

Scope: Withdrawal of initial marketing authorisation application

Action: For information

The CHMP noted the withdrawal of the initial marketing authorisation application.

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. Azacitidine Accord - Azacitidine - EMEA/H/C/005147/X/0021

Accord Healthcare S.L.U.;

Rapporteur: Hrefna Gudmundsdottir, PRAC Rapporteur: Bianca Mulder

Scope: "Extension application to introduce a new pharmaceutical form (film-coated tablet) associated with new strengths (200 and 300 mg) and new route of administration (oral use). The RMP (version 2.0) is updated in accordance."

Action: For adoption

List of Outstanding Issues adopted on 22.05.2025. List of Questions adopted on 12.12.2024.

The Committee confirmed that all issues previously identified in this application had been addressed.

The Committee adopted a positive opinion by consensus together with the CHMP Assessment Report and translation timetable.

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. Enzalutamide Viatris - Enzalutamide - EMEA/H/C/006299/X/0003

Viatris Limited;

Rapporteur: Tomas Radimersky, PRAC Rapporteur: Maria del Pilar Rayon

Scope: "Extension application to add a new strength of 160 mg for solution for film-coated tablets.

The RMP (version 1.0) is updated in accordance."

Action: For adoption

List of Questions adopted on 27.03.2025.

The Committee discussed the issues identified in this application, relating to quality, clinical and non-clinical aspects.

The Committee adopted the CHMP recommendation and scientific discussion together with the list of outstanding issues and a specific timetable.

4.2.2. Hetlioz - Tasimelteon - Orphan - EMEA/H/C/003870/X/0039

Vanda Pharmaceuticals Netherlands B.V.;

Rapporteur: Jayne Crowe, PRAC Rapporteur: Adam Przybylkowski

Scope: "Extension application to introduce a new pharmaceutical form associated with new strength (4 mg/ml oral solution). The new formulation is indicated for the treatment of nighttime sleep disturbances in Smith-Magenis Syndrome (SMS) in paediatric patients 3 to 15 years of age. The RMP (version 5.0) is updated in accordance."

Action: For adoption

List of Questions adopted on 27.02.2025.

The Committee discussed the issues identified in this application, relating to clinical and orphan similarity aspects.

The Committee adopted the CHMP recommendation and scientific discussion together with the list of outstanding issues and a specific timetable.

4.2.3. Livmarli - Maralixibat - Orphan - EMEA/H/C/005857/X/0015

Mirum Pharmaceuticals International B.V.;

Rapporteur: Janet Koenig, PRAC Rapporteur: Adam Przybylkowski

Scope: "Extension application to introduce a new pharmaceutical form (tablet) associated with new strengths 10 mg, 15mg, 20 mg and 30 mg. The RMP (version 5.0) is updated in accordance."

Request by the applicant for an extension to the clock-stop to respond to the list of outstanding issues to be adopted in July 2025.

Action: For adoption

List of Questions adopted on 27.02.2025.

The Committee discussed the issues identified in this application, relating to quality and clinical aspects.

The Committee adopted the CHMP recommendation and scientific discussion together with the list of outstanding issues and a specific timetable.

The Committee denied the request by the applicant for an extension to the clock-stop to respond to the list of outstanding issues adopted in July 2025.

4.2.4. Pyrukynd - Mitapivat - Orphan - EMEA/H/C/005540/X/0010/G

Agios Netherlands B.V.;

Rapporteur: Alexandre Moreau, PRAC Rapporteur: Adam Przybylkowski

Scope: "Extension application to introduce a new strength (100 mg film-coated tablet) associated with a new orphan indication for the "treatment of adult patients with non-transfusion-dependent and transfusion-dependent alpha- or beta-thalassaemia". The extension application is grouped with a type II quality variation (C.I.4) to update of sections 4.2 and 5.2 of the SmPC in order to update pharmacokinetic information based on final results from study AG348-C-024 listed as a category 3 study in the RMP; this is a Phase 1, Open-label, Single-dose, Pharmacokinetic Study of Mitapivat in Subjects with Moderate Hepatic Impairment Compared to Matched Healthy Control Subjects with Normal Hepatic Function. The RMP (version 1.1) is updated in accordance."

Action: For adoption

List of Questions adopted on 27.03.2025.

The Committee discussed the issues identified in this application, relating to clinical aspects.

The Committee adopted the CHMP recommendation and scientific discussion together with the list of outstanding issues and a specific timetable.

4.2.5. Remsima - Infliximab - EMEA/H/C/002576/X/0149

Celltrion Healthcare Hungary Kft.;

Rapporteur: Outi Mäki-Ikola, PRAC Rapporteur: Kimmo Jaakkola

Scope: "Extension application to introduce a new pharmaceutical form (concentrate for solution for infusion) associated with a new strength (40 mg/ml)."

Action: For adoption

List of Questions adopted on 25.04.2025.

The Committee discussed the issues identified in this application, relating to quality aspects.

The Committee adopted the CHMP recommendation and scientific discussion together with the list of outstanding issues and a specific timetable.

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. Abrysvo - Respiratory syncytial virus vaccine (bivalent, recombinant) - EMA/X/0000258051

Pfizer Europe MA EEIG

Rapporteur: Jayne Crowe

Scope: Extension application to introduce a new pharmaceutical form Powder and solvent for solution for injection in multidose container.

Action: For adoption

The Committee discussed the issues identified in this application, relating to quality and clinical aspects.

The Committee adopted the CHMP recommendation and scientific discussion together with the list of questions and a specific timetable.

4.3.2. Akynzeo - Fosnetupitant / Netupitant / Palonosetron - EMA/X/0000258060

Helsinn Birex Pharmaceuticals Limited

Rapporteur: Finbarr Leacy, PRAC Rapporteur: Amelia Cupelli

Scope: Extension application to introduce a new pharmaceutical form (300 mg / 0.5 ml oral suspension).

Action: For adoption

The Committee discussed the issues identified in this application, relating to quality aspects.

The Committee adopted the CHMP recommendation and scientific discussion together with the list of questions and a specific timetable.

4.3.3. Camcevi – Leuprorelin - EMA/X/0000258054

Accord Healthcare S.L.U.

Rapporteur: Johanna Lähteenvuo, PRAC Rapporteur: Amelia Cupelli

Scope: Extension application to add a new strength of 21 mg for Leuprorelin prolonged-release suspension for injection pre-filled syringe, for subcutaneous (SC) administration.

Action: For adoption

The Committee discussed the issues identified in this application, relating to quality and clinical aspects

The Committee adopted the CHMP recommendation and scientific discussion together with the list of questions and a specific timetable.

4.3.4. Jorveza – Budesonide - EMA/X/0000257468

Dr. Falk Pharma GmbH

Rapporteur: Janet Koenig, PRAC Rapporteur: Zane Neikena

Scope: Extension application to introduce a new pharmaceutical form associated with new strength (0.2 mg/ml oral suspension). The new presentation is indicated for paediatric patients 2 to 17 years of age.

Action: For adoption

The Committee discussed the issues identified in this application, relating to quality aspects.

The Committee adopted the CHMP recommendation and scientific discussion together with the list of questions and a specific timetable.

4.3.5. Lojuxta – Lomitapide - EMA/X/0000258068

Chiesi Farmaceutici S.p.A.

Rapporteur: Patrick Vrijlandt, PRAC Rapporteur: Bianca Mulder

Scope: Extension application to add a new strength of 2 mg hard capsules.

This application is grouped with

- type II variation (C.I.6.a): an Extension of Indication to include treatment of paediatric patients aged 5 years and older with homozygous familial hypercholesterolaemia (HoFH) for LOJUXTA, based on final results from the pivotal paediatric study APH-19; this is a phase 3, single-arm, open-label, international, multi-centre study to evaluate the efficacy and safety of lomitapide in paediatric patients with homozygous familial hypercholesterolaemia (HOFH) on stable lipid-lowering therapy. As a consequence, sections 4.1, 4.2, 4.4, 4.6, 4.8, 5.1, 5.2 and 5.3 of the SmPC are updated. The Annex II and Package Leaflet are updated accordingly. The RMP version 7.1 has also been submitted. In addition, the MAH took the opportunity to bring the PI in line with the latest QRD template version 10.4.

Action: For adoption

The Committee discussed the issues identified in this application, relating to quality aspects.

The Committee adopted the CHMP recommendation and scientific discussion together with the list of questions and a specific timetable.

4.3.6. Nexviadyme - Avalglucosidase alfa - EMA/X/0000258013

Sanofi B.V.

Rapporteur: Christian Gartner

Scope: Quality

Action: For adoption

The Committee discussed the issues identified in this application, relating to quality aspects.

The Committee adopted the CHMP recommendation and scientific discussion together with the list of questions and a specific timetable.

4.3.7. Olumiant – Baricitinib - EMA/X/0000257923

Eli Lilly Nederland B.V.

Rapporteur: Peter Mol, PRAC Rapporteur: Adam Przybylkowski

Scope: Extension application to introduce a new pharmaceutical form (oral suspension) associated with a new strength (2 mg/ml).

Action: For adoption

The Committee discussed the issues identified in this application EMA/X/0000257923.

The Committee adopted the CHMP recommendation and scientific discussion together with the list of questions and a specific timetable.

4.3.8. Scemblix – Asciminib - EMA/X/0000256688

Novartis Europharm Limited

Rapporteur: Jante Koenig, Co-Rapporteur: Peter Mol, PRAC Rapporteur: Eva Jirsová

Scope: Extension application to introduce a new strength (100 mg film-coated tablets) grouped with a type II variation (C.I.6.a) to add a new indication (treatment of adult patients with Philadelphia chromosome-positive chronic myeloid leukaemia in chronic phase (Ph+ CML-CP) harbouring the T315I mutation), based on final results from study CABL001X2101 and study CABL001A2004. Study CABL001X2101 is a Phase I, multicentre, open-label, dose escalation FIH study to define the MTD/RDEs, to characterize safety and tolerability, and to assess the PK profile and preliminary evidence of efficacy of asciminib given as single agent or in combination with either nilotinib or imatinib or dasatinib in patients with Ph+ CML or Ph+ ALL.

Study CABL001A2004 assessed the real-world effectiveness of asciminib and treatment patterns in patients with Chronic Myeloid Leukaemia with T315I mutation. As a consequence, sections 1, 2, 3, 4, 5, 6 and 8 of the SmPC are updated. The Package Leaflet and Labelling are updated in accordance. Version 3.0 of the RMP has also been submitted. As part of the application the MAH is requesting a 1-year extension of the market protection.

Action: For adoption

The Committee discussed the issues identified in this application, relating to clinical aspects.

The Committee adopted the CHMP recommendation and scientific discussion together with the list of questions and a specific timetable.

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

No items

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

No items

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. Alhemo – Concizumab - EMA/VR/0000244862

Novo Nordisk A/S

Rapporteur: Patrick Vrijlandt, Co-Rapporteur: Daniela Philadelphia, PRAC Rapporteur: Marie Louise Schougaard Christiansen

Scope: Extension of indication to include treatment of haemophilia A without inhibitors and haemophilia B without inhibitors for ALHEMO based on final results from study NN7415-4307; this is an interventional study to investigate efficacy and safety of concizumab prophylaxis in patients with haemophilia A or B without inhibitors. As a consequence, sections 4.1, 4.2, 4.8, 5.1 and 5.2 of the SmPC are updated. The Package Leaflet is updated in accordance. Version 2.0 of the RMP has also been submitted. In addition, the MAH took the opportunity to introduce minor changes to the PI.

Action: For adoption

The Committee confirmed that all issues previously identified in this application had been addressed.

The Committee adopted a positive opinion by consensus together with the CHMP Assessment Report and translation timetable.

The summary of opinion was circulated for information.

The CHMP adopted the similarity assessment report

5.1.2. BAQSIMI – Glucagon - EMA/VR/0000244909

Amphastar France Pharmaceuticals

Rapporteur: Karin Janssen van Doorn, PRAC Rapporteur: Eamon O Murchu

Scope: Extension of indication to include treatment of severe hypoglycaemia in paediatric patients aged 1 and over with diabetes mellitus for BAQSIMI, based on final results from study I8R-MC-IGBO; this is an Open-Label, Multi-Center, Single-Dose Study to Assess the Safety, Tolerability, Pharmacodynamics, and Pharmacokinetics of Nasal Glucagon in Paediatric Patients with Type 1 Diabetes Aged 1 to <4 years; As a consequence, sections 4.1, 4.2, 4.8, 5.1 and 5.2 of the SmPC are updated. The Package Leaflet is updated in accordance. Version 1.0 of the RMP has also been submitted. In addition, the Marketing authorisation holder (MAH) took the opportunity to introduce a correction in the Package Leaflet.

Action: For adoption

The Committee confirmed that all issues previously identified in this application had been addressed.

The Committee adopted a positive opinion by consensus together with the CHMP Assessment Report and translation timetable.

The summary of opinion was circulated for information.

5.1.3. BESONSA - Inotuzumab ozogamicin - EMA/VR/0000257310

Pfizer Europe MA EEIG

Rapporteur: Filip Josephson, PRAC Rapporteur: Gabriele Maurer

Scope: Extension of indication to include treatment of paediatric patients 1 year and older with relapsed or refractory CD22-positive B-cell precursor acute lymphoblastic leukaemia (ALL) for BESONSA, based on final results from studies ITCC-059 (WI203581) and INO-Ped-ALL-1 (WI235086).

Study WI203581 is a Phase 1/2, multicentre, European, multi-cohort, open-label study in paediatric patients (≥ 1 and < 18 years of age) with R/R CD22-positive ALL; Study WI235086 is an open-label, Phase 1 study to assess safety and tolerability of InO in Japanese paediatric patients with R/R CD22-positive AL.

As a consequence, sections 4.1, 4.2, 4.8, 5.1 and 6.6 of the SmPC are updated. The Package Leaflet is updated in accordance. Version 4.0 of the RMP has also been submitted. In addition, the Marketing authorisation holder (MAH) took the opportunity to update the list of local representatives in the Package Leaflet.

Action: For adoption

The Committee discussed the issues identified in this application, relating to clinical aspects.

The Committee adopted a request for supplementary information with a specific timetable.

5.1.4. Breyanzi - Lisocabtagene maraleucel – ATMP – EMA/VR/0000265024

Bristol-Myers Squibb Pharma EEIG

CAT Rapporteur: Concetta Quintarelli, CHMP Coordinator: Paolo Gasparini

Scope: A grouped application comprised of two Type II variations, as follows:

Type II (C.I.6): Extension of indication to include the treatment of adult patients with relapsed or refractory mantle cell lymphoma (MCL) after at least two lines of systemic therapy, including a Bruton's tyrosine kinase (BTK) inhibitor for BREYANZI, based on results from the pivotal Study 017001 MCL Cohort (TRANSCEND-NHL-001); this is a Phase 1, Multicentre, Open-Label Study of JCAR017, CD19-targeted Chimeric Antigen Receptor (CAR) T Cells, for Relapsed and Refractory (R/R) B-cell Non-Hodgkin Lymphoma (NHL). As a consequence, sections 4.1, 4.4, 4.8, 5.1 and 5.2 of the SmPC are updated. The Package leaflet is updated in accordance. Version 7.0 of the RMP has also been submitted. In addition, the MAH took the opportunity to introduce minor editorial changes to the PI and to update the list of local representatives in the Package Leaflet.

Quality variation

Action: For adoption

The CHMP was updated on discussions at the CAT. The Committee discussed the issues identified in this application.

The Committee discussed the issues identified in this application, relating to quality aspects .

The Committee endorsed a request for supplementary information with a specific timetable, as adopted by CAT.

5.1.5. Cejemy – Sugemalimab - EMA/VR/0000261157

Cstone Pharmaceuticals Ireland Limited

Rapporteur: Filip Josephson, PRAC Rapporteur: Petar Mas

Scope: Extension of indication to include the treatment of unresectable stage III non-small-cell lung cancer (NSCLC) with no sensitising EGFR mutations, or ALK, ROS1 genomic tumour aberrations in adults whose disease has not progressed following concurrent or sequential platinum-based chemoradiotherapy for CEJEMLY, based on final results from study CS1001-301; this is a Phase III, multicentre, randomised, double-blind, placebo-controlled study assessing the efficacy and safety of sugemalimab as consolidation therapy versus placebo in participants with locally advanced or unresectable stage III NSCLC who have not progressed after concurrent or sequential chemoradiotherapy. 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. Version 2.0 of the RMP has also been submitted.

Action: For adoption

The Committee discussed the issues identified in this application, relating to clinical aspects.

The Committee adopted a request for supplementary information with a specific timetable.

5.1.6. Clopidogrel Zentiva - Clopidogrel - EMEA/H/C/000975/II/0092

Zentiva k.s.;

Rapporteur: Fátima Ventura, PRAC Rapporteur: Carla Torre

Scope: "Extension of indication to include, in combination with acetylsalicylic acid (ASA), patients with ST segment elevation acute myocardial infarction (STEMI) who are undergoing percutaneous coronary intervention (PCI) for CLOPIDOGREL ZENTIVA. As a consequence, sections 4.1, 4.2, 4.4 and 5.1 of the SmPC are updated. Version 0.1 of the RMP has also been submitted. In addition, the Marketing authorisation holder (MAH) took the opportunity to update the list of local representatives in the Package Leaflet, introduce minor editorial changes to the PI and bring it in line with the latest QRD template version 10.4."

Action: For adoption

Request for Supplementary Information adopted on 25.04.2025.

The Committee confirmed that all issues previously identified in this application had been addressed.

The Committee adopted a positive opinion by consensus together with the CHMP Assessment Report and translation timetable.

The summary of opinion was circulated for information.

5.1.7. CRYSVITA – Burosumab - EMA/VR/0000263400

Kyowa Kirin Holdings B.V.

Rapporteur: Kristina Dunder

Scope: Extension of indication to include treatment of X-linked hypophosphataemia (XLH) in paediatric patients from birth to less than 1 year of age for CRYSVITA, based on final results from study BUR-CL207; this is a phase 1/2 Open-label, Multicentre, Non-randomized Study to Evaluate Safety, Pharmacodynamics, Pharmacokinetics and Effect of Burosumab in Paediatric Patients from Birth to Less than 1 Year of Age with XLH; 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.

Action: For adoption

The Committee discussed the issues identified in this application, relating to clinical aspects.

The Committee adopted a request for supplementary information with a specific timetable.

5.1.8. Elucirem/ Vueway – Gadopiclenol - EMA/VR/0000249008

Guerbet, Bracco Imaging S.p.A

Rapporteur: Patrick Vrijlandt, PRAC Rapporteur: Martin Huber

Scope: Extension of indication to include treatment of new population (0 to 2 years of age patients) for ELUCIREM / VUEWAY, based on final results from study GDX-44-015; this is a phase ii clinical study concerning gadopiclenol pharmacokinetics, safety and efficacy in paediatric patients < 2 years of age undergoing contrast-enhanced MRI; extension of indication is also supported with the non-clinical data. 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. Version 0.4 of the RMP has also been submitted. In addition, the Marketing authorisation holder (MAH) took the opportunity to remove Annex IV from the PI. As part of the application, the MAH is requesting a 1-year extension of the market protection.

Action: For adoption

The Committee discussed the issues identified in this application, relating to clinical aspects.

The Committee adopted a request for supplementary information with a specific timetable.

5.1.9. Eylea – Aflibercept - EMA/VR/0000264981

Bayer AG

Rapporteur: Jean-Michel Race, PRAC Rapporteur: Zoubida Amimour

Scope: A grouped application comprised of two Type II Variations, as follows:

C.I.6: Extension of indication to include the treatment of visual impairment due to macular oedema secondary to retinal vein occlusion (branch, central and hemiretinal RVO) for EYLEA, based on results from study 22153 (QUASAR); this is a randomized, double-masked, active-controlled Phase 3 study of the efficacy and safety of aflibercept 8 mg in macular oedema secondary to retinal vein occlusion. 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

accordingly. The RMP version 36.1 has also been submitted.

C.I.4: Update of section 4.2 of the SmPC in order to change posology recommendations of the approved indications nAMD and DME based on the results from study 22153 (QUASAR) and post-hoc analysis of the pivotal studies 20968 (PULSAR), 21091 (PHOTON) and Phase II study 21086 (CANDELA).

Action: For adoption

The Committee discussed the issues identified in this application, relating to clinical aspects.

The Committee adopted a request for supplementary information with a specific timetable.

5.1.10. Gazyvaro – Obinutuzumab - EMA/VR/0000244907

Roche Registration GmbH

Rapporteur: Boje Kvorning Pires Ehmsen, PRAC Rapporteur: Mari Thorn

Scope: Extension of indication to include treatment of adult patients with active lupus nephritis who are receiving standard therapy for GAZYVARO, based on results from study Regency (CA41705). This is an ongoing, Phase III, randomized, double-blind, placebo-controlled, multicentre study evaluating the efficacy and safety of obinutuzumab administered at standard infusion rates in patients with ISN/RPS 2003 Class III or IV lupus nephritis treated with standard-of-care therapy.

As a consequence, sections 4.1, 4.2, 4.4, 4.8, 5.1, 5.2 and 6.6 of the SmPC are updated. The Package Leaflet is updated in accordance. Version 11 of the RMP has also been submitted. In addition, the Marketing authorisation holder (MAH) took the opportunity to update the list of local representatives in the Package Leaflet.

Action: For adoption

The Committee discussed the issues identified in this application, relating to clinical aspects.

The Committee adopted a request for supplementary information with a specific timetable.

5.1.11. Iclusig – Ponatinib - EMA/VR/0000263550

Incyte Biosciences Distribution B.V.

Rapporteur: Filip Josephson, Co-Rapporteur: Ewa Balkowiec Iskra, PRAC Rapporteur: Mari Thorn

Scope: Extension of indication to include treatment of adult patients with newly-diagnosed Ph+ ALL for ICLUSIG, based on interim results from study Ponatinib-3001 (PhALLCON); this is a phase 3, randomized, open-label, multicentre study comparing ponatinib versus imatinib, administered in combination with reduced intensity chemotherapy, in patients with newly diagnosed Ph+ ALL; supportive data were derived from two single-arm, open-label clinical studies (AP24534 11 001 in combination with chemotherapy and INCB 84344-201 as monotherapy). 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. Version 23.2 of the RMP has also been submitted. In addition, earlier approved updates were incorporated to the PI.

Action: For adoption

The Committee discussed the issues identified in this application, relating to clinical and

pharmacovigilance aspects.

The Committee adopted a request for supplementary information with a specific timetable.

5.1.12. Invokana - Canagliflozin - EMEA/H/C/002649/II/0069

Janssen-Cilag International N.V.;

Rapporteur: Janet Koenig, PRAC Rapporteur: Martin Huber

Scope: "Extension of indication to include treatment of paediatric patients with type 2 diabetes mellitus aged 10 years old and older for INVOKANA, based on final results from study JNJ-28431754DIA3018 as well as study JNJ-28431754DIA1055. Study JNJ-28431754DIA3018 is a double-blind, placebo-controlled, 2-arm, parallel-group, multicentre Phase 3 study in participants with T2DM >10 and <18 years of age who had inadequate glycemic control (i.e., HbA1c of >6.5% to <11.0%). As a consequence, sections 4.1, 4.2, 4.4, 4.5, 4.8, 5.1 and 5.2 of the SmPC are updated. The Package Leaflet is updated in accordance. Version 13.1 of the RMP has also been submitted. In addition, the Marketing authorisation holder (MAH) took the opportunity to introduce minor changes to the PI and update the list of local representatives in the Package Leaflet."

Action: For adoption

Request for Supplementary Information adopted on 27.03.2025, 14.11.2024.

The Committee confirmed that all issues previously identified in this application had been addressed.

The Committee adopted a positive opinion by consensus together with the CHMP Assessment Report and translation timetable.

The summary of opinion was circulated for information.

5.1.13. Keytruda – Pembrolizumab - EMA/VR/0000245108

Merck Sharp & Dohme B.V.

Rapporteur: Paolo Gasparini, PRAC Rapporteur: Bianca Mulder

Scope: Extension of indication to include, KEYTRUDA as monotherapy, for the treatment of resectable locally advanced head and neck squamous cell carcinoma (HNSCC) as neoadjuvant treatment, continued as adjuvant treatment in combination with radiation therapy with or without platinum-containing chemotherapy and then as monotherapy in adults, based on the results of study P689V01MK3475 (KEYNOTE-689); this is a Phase 3, randomised, open-label study evaluating pembrolizumab as neoadjuvant therapy and in combination with standard of care as adjuvant therapy for stage III or IVA, resectable, locoregionally advanced head and neck squamous cell carcinoma. Consequently, sections 4.1, 4.2, 4.8 and 5.1 of the SmPC are updated. The Package Leaflet is updated in accordance. The RMP version 48.1 has also been submitted. In addition, the MAH took the opportunity to introduce some minor editorial changes to the PI.

Action: For adoption

The Committee discussed the issues identified in this application, relating to clinical aspects.

The Committee adopted a request for supplementary information with a specific timetable.

5.1.14. LIBTAYO – Cemiplimab - EMA/VR/0000264999

Regeneron Ireland Designated Activity Company

Rapporteur: Boje Kvorning Pires Ehmsen, PRAC Rapporteur: Bianca Mulder

Scope: Extension of indication to include adjuvant treatment of adult patients with Cutaneous Squamous Cell Carcinoma (CSCC) at high risk of recurrence after surgery and radiation for LIBTAYO, based on interim results from study R2810-ONC-1788; this is a phase 3, randomized, placebo-controlled, double-blind study of adjuvant cemiplimab versus placebo after surgery and radiation therapy in patients with high risk CSCC; As a consequence, sections 4.1, 4.2, 4.8, 5.1, and 6.6 of the SmPC are updated. The Package Leaflet is updated in accordance. Version 4.2 of the RMP has also been submitted. In addition, the Marketing authorisation holder (MAH) took the opportunity to update the warnings for the excipients proline and polysorbate to reflect EU guidance (Section 4.4), and also updated Annex IID of the PI in line with the updates made to the RMPv4.2 to consolidate the aRMMs.

Action: For adoption

The Committee discussed the issues identified in this application, relating to clinical aspects.

The Committee adopted a request for supplementary information with a specific timetable.

5.1.15. mRESVIA - Respiratory syncytial virus mRNA vaccine (nucleoside modified) - EMA/VR/0000248175

Moderna Biotech Spain S.L.

Rapporteur: Jan Mueller-Berghaus, PRAC Rapporteur: Jean-Michel Dogné

Scope: To modify the approved therapeutic indication to include active immunisation for the prevention of lower respiratory tract disease (LRTD) caused by Respiratory Syncytial Virus in individuals 18 through 59 years of age who are at increased risk for LRTD caused by RSV for mRESVIA, based on results from Study mRNA-1345-P303 (Part A) - A Phase 3 Study to Evaluate the Immunogenicity and Safety of mRNA-1345, an mRNA Vaccine Targeting Respiratory Syncytial Virus, in High-risk Adults. As a consequence, sections 4.1, 4.6, 4.8 and 5.1 of the SmPC and the corresponding sections of the Package Leaflet are updated accordingly. The updated RMP Version 1.0 has also been submitted. In addition, the MAH took the opportunity to implement editorial changes to the PI.

Action: For adoption

The Committee confirmed that all issues previously identified in this application had been addressed.

The Committee adopted a positive opinion by consensus together with the CHMP Assessment Report and translation timetable.

The summary of opinion was circulated for information.

5.1.16. Neuraceq - Florbetaben (18F) - EMA/VR/0000227744

Life Molecular Imaging GmbH

Rapporteur: Antonio Gomez-Outes, PRAC Rapporteur: Martin Huber

Scope: Update of section 4.5 of the SmPC to reflect new preclinical data and editorial update of section 5.1. The Package Leaflet is also updated with discontinuation of the paper copy SmPC. In addition, Annex IID is updated with deletion of additional risk minimisation measures. The RMP version 7.0 is agreed.

Action: For adoption

The Committee confirmed that all issues previously identified in this application had been addressed.

The Committee adopted a positive opinion by consensus together with the CHMP Assessment Report and translation timetable.

The CHMP noted the Q&A document.

The summary of opinion was circulated for information.

5.1.17. Noxafil – Posaconazole - EMA/VR/0000263360

Merck Sharp & Dohme B.V.

Rapporteur: Alexandre Moreau, PRAC Rapporteur: Zoubida Amimour

Scope: Extension of indication for NOXAFIL to include treatment of patients two years of age and older for invasive aspergillosis (IA) based on final results from study MK-5592-104 (P104); this is a Phase 2, open-label, noncomparative clinical study that evaluated the safety, efficacy, and PK of POS in paediatric participants aged 2 to <18 years with IA. As a consequence, sections 4.1, 4.2, 4.8, 5.1 and 5.2 of the SmPC are updated. The Package Leaflet and Labelling are updated in accordance. Version 18.1 of the RMP has also been submitted. In addition, the Marketing authorisation holder (MAH) took the opportunity to implement editorial changes to the PI.

Action: For adoption

The Committee discussed the issues identified in this application, relating to clinical aspects.

The Committee adopted a request for supplementary information with a specific timetable.

5.1.18. Recarbrio - Imipenem / Cilastatin / Relebactam - EMA/VR/0000265089

Merck Sharp & Dohme B.V.

Rapporteur: Filip Josephson, Co-Rapporteur: Alar Irs, PRAC Rapporteur: Adam Przybylkowski

Scope: Extension of indication to extend the approved adult indications for RECARBRI to include treatment of paediatric population from birth to <18 years of age, based on final results from two paediatric studies (MK-7655A-021 and MK-7655A-020); phase 2/3 study MK-7655A-021 addressed safety, tolerability, efficacy and PK, and phase 1b study MK-7655A-020 addressed PK, safety, and tolerability of MK-7655A in paediatric subjects from birth to less than 18 years of age with confirmed or suspected gram-negative infections. As a consequence, sections 4.1, 4.2, 4.8, 5.1, 5.2, and 6.6 of the SmPC are updated. The Package Leaflet is updated in accordance. Version 2.1 of the RMP has also been submitted. In addition, the Marketing authorisation holder (MAH) took the opportunity to update the list of local representatives in the Package Leaflet and implement minor editorial corrections.

Action: For adoption

The Committee discussed the issues identified in this application, relating to clinical and non-clinical aspects.

The Committee adopted a request for supplementary information with a specific timetable.

5.1.19. Scemblix – Asciminib - EMA/VR/0000265010

Novartis Europharm Limited

Rapporteur: Janet Koenig, PRAC Rapporteur: Eva Jirsová

Scope: A grouped application consisting of:

C.I.6.a: Extension of indication to include treatment of adult patients with newly diagnosed or previously treated Philadelphia chromosome-positive chronic myeloid leukaemia (Ph+ CML) in chronic phase (CP) for SCEMBLIX, based on primary and key secondary analysis results from study CABL001J12301 (ASC4FIRST, J12301); this is an ongoing Phase III, multi-centre, open-label, randomized study of oral asciminib (80 mg once daily, q.d.) versus Investigator selected tyrosine kinase inhibitor (TKI) in patients with newly diagnosed Ph+ CML-CP, with the primary and key secondary objectives to compare the major molecular response (MMR) rates at Week 48 and Week 96, respectively. As a consequence, sections 4.1, 4.8 and 5.1 of the SmPC are updated. The Package Leaflet is updated accordingly. RMP version 4.0 has also been submitted. As part of the application the MAH is requesting a 1-year extension of the market protection.

C.I.4: Update of sections 4.2, 4.5, 5.1, 5.2 and 5.3 of the SmPC in order to introduction of a new posology regimen based on results from studies CABL001J12301 and CABL001A2302 (ASC4OPT, A2302). CABL001A2302 is an ongoing Phase IIIb, multi-centre, open-label, treatment optimization study of oral asciminib (80 mg daily, randomized to 40 mg b.i.d. or 80 mg q.d.) in patients with Ph+ CML-CP previously treated with two or more TKIs, with the primary objective to estimate the MMR rate at Week 48 of all the patients (40 mg b.i.d. and 80 mg q.d.) with no evidence of MMR at baseline. The Package Leaflet is updated accordingly. RMP version 4.0 has also been submitted.

Action: For adoption

The Committee discussed the issues identified in this application, relating to clinical and non-clinical aspects .

The Committee adopted a request for supplementary information with a specific timetable.

5.1.20. SIRTURO – Bedaquiline - EMA/VR/0000249065

Janssen Cilag International

Rapporteur: Filip Josephson, PRAC Rapporteur: Karin Bolin

Scope: Extension of indication to include treatment of paediatric patients (2 years to less than 5 years of age and weighing at least 7 kg) with pulmonary tuberculosis (TB) due to *Mycobacterium tuberculosis* resistant to at least rifampicin and isoniazid, for SIRTURO, based on the Week 24 primary analysis from Cohort 3 (≥ 2 to < 5 years of age) of Study TMC207-C211; this is an open-label, multicentre, single-arm study to evaluate pharmacokinetics, safety/tolerability, antimycobacterial activity and dose selection of

bedaquiline in children (birth to <18 years) with multidrug-resistant-TB (MDR-TB). Long-term follow-up to Week 120 in participants of Cohort 1 (≥ 12 to <18 years of age) and Cohort 2 (≥ 5 to <12 years of age) have also been submitted. As a consequence, sections 4.1, 4.2, 4.8, 5.1 and 5.2 of the SmPC are updated. The Package Leaflet is updated in accordance. Version 11.1 of the RMP has also been approved. In addition, the Marketing authorisation holder (MAH) took the opportunity to update the list of local representatives in the Package Leaflet and introduce minor changes to the PI.

Action: For adoption

The Committee confirmed that all issues previously identified in this application had been addressed.

The Committee adopted a positive opinion by consensus together with the CHMP Assessment Report and translation timetable.

The summary of opinion was circulated for information.

5.1.21. Sogroya – Somapacitan - EMA/VR/0000264734

Novo Nordisk A/S

Rapporteur: Patrick Vrijlandt, PRAC Rapporteur: Martin Huber

Scope: Grouped extension of indication application to include treatment of children born small for gestational age (SGA), Noonan syndrome (NS) and idiopathic short stature (ISS) for SOGROYA, based on interim results from the pivotal, confirmatory phase 3 study NN8640-4467 supported by the phase 3 study NN8640-4469 and the phase 2 study NN8640-4245. Study 4467 is a study comparing the effect and safety of once weekly dosing of somapacitan with daily Norditropin as well as evaluating long-term safety of somapacitan in a basket study design in children with short stature either born small for gestational age or with Turner syndrome, Noonan syndrome, or idiopathic short stature. Study 4469 is a study evaluating the safety and efficacy of once-weekly dosing of somapacitan in a basket study design in paediatric participants with short stature either born small for gestational age or with Turner syndrome, Noonan syndrome, or idiopathic short stature. Study 4245 is a dose-finding trial evaluating the effect and safety of once-weekly treatment of somapacitan compared to daily Norditropin in children with short stature born small for gestational age with no catch-up growth by 2 years of age or older. As a consequence, sections 4.1, 4.2, 4.8, 5.1 and 5.2 of the SmPC are updated. The Package Leaflet is updated in accordance. Version 4.0 of the RMP has also been submitted. Furthermore, the PI is brought in line with the latest QRD template version 10.4. As part of the application, the MAH is requesting a 1-year extension of the market protection.

Action: For adoption

The Committee discussed the issues identified in this application, relating to clinical aspects.

The Committee adopted a request for supplementary information with a specific timetable.

5.1.22. Taltz - Ixekizumab - EMEA/H/C/003943/II/0053

Eli Lilly and Co (Ireland) Limited;

Rapporteur: Kristina Dunder, PRAC Rapporteur: Gabriele Maurer

Scope: "Extension of indication for TALTZ to include treatment alone or in combination with methotrexate, of active JPsA and ERA in patients 6 years of age and older and with a body weight of at least 25 kg, who have had an inadequate response to, or who are intolerant of, conventional therapy, based on week 16 results from study I1F-MC-RHCG. As a consequence, sections 4.1, 4.2, 4.8, 5.1 and 5.2 of the SmPC are updated. The Package Leaflet is updated in accordance. Version 8.2 of the RMP has also been updated. Furthermore, the PI is in line with the latest QRD template version 10.4."

Action: For adoption

Request for Supplementary Information adopted on 27.03.2025, 14.11.2024.

The Committee confirmed that all issues previously identified in this application had been addressed.

The Committee adopted a positive opinion by consensus together with the CHMP Assessment Report and translation timetable.

The summary of opinion was circulated for information.

5.1.23. Tevibra - Tislelizumab - EMEA/H/C/005919/II/0018

Beone Medicines Ireland Limited;

Rapporteur: Jan Mueller-Berghaus, PRAC Rapporteur: Bianca Mulder

Scope: "Extension of indication for Tevibra in combination with platinum-containing chemotherapy as neoadjuvant treatment and then continued as monotherapy as adjuvant treatment, for the treatment of adult patients with resectable NSCLC based on interim results from study BGB-A317-315. Study BGB-A317-315 is a phase 3 randomized, placebo-controlled, double-blind study to compare the efficacy and safety of neoadjuvant treatment with tislelizumab plus platinum-based doublet chemotherapy followed by adjuvant tislelizumab versus neoadjuvant treatment with placebo plus platinum-based doublet chemotherapy followed by adjuvant placebo in patients with resectable Stage II or IIIA NSCLC. As a consequence, sections 4.1, 4.2, 5.1, and 6.6 of the SmPC are updated. The Package Leaflet is updated in accordance. Version 6.0 of the RMP has also been agreed."

Action: For adoption

Request for Supplementary Information adopted on 27.03.2025.

The Committee confirmed that all issues previously identified in this application had been addressed.

The Committee adopted a positive opinion by consensus together with the CHMP Assessment Report and translation timetable.

The summary of opinion was circulated for information.

5.1.24. TEZSPIRE – Tezepelumab - EMA/VR/0000245013

AstraZeneca AB

Rapporteur: Finbarr Leacy, Co-rapporteur: Ewa Balkowiec Iskra, PRAC Rapporteur: Eva Jirsová

Scope: Extension of indication to include treatment of Chronic Rhinosinusitis with Nasal

Polyps (CRSwNP) for Tezspire, based on results from study WAYPOINT (D5242C00001); this is a global, multicentre, randomised, double-blind, parallel-group, placebo-controlled study that evaluated the efficacy and safety of tezepelumab compared with placebo in the treatment of CRSwNP. As a consequence, sections 4.1, 4.2, 4.8, 5.1 and 5.2 of the SmPC are updated. The Package Leaflet and Labelling are updated in accordance. Version 4.1 of the RMP has also been submitted. In addition, the Marketing authorisation holder (MAH) took the opportunity to implement editorial changes and to update the PI and the Package Leaflet in accordance with the latest EMA excipients guideline.

Action: For adoption

The Committee discussed the issues identified in this application, relating to clinical aspects
The Committee adopted a request for supplementary information with a specific timetable.

5.1.25. VEYVONDI - Vonicog alfa - EMA/VR/0000264863

BAXALTA INNOVATIONS GmbH

Rapporteur: Jan Mueller-Berghaus, Co-Rapporteur: Daniela Philadelphia, PRAC Rapporteur: Mari Thorn

Scope: Extension of indication to include treatment of haemorrhage in children aged less than 18 years for VEYVONDI, based on results from studies 071102 and SHP677-304. Study 071102 is a phase 3, prospective, multicentre, uncontrolled, open-label clinical study to determine the efficacy, safety, and tolerability of rVWF with or without ADVATE in the treatment and control of bleeding episodes, the efficacy and safety of rVWF in elective and emergency surgeries, and the pharmacokinetics (PK) of rVWF in children diagnosed with severe VWD; study SHP677-304 is a phase 3B, prospective, open-label, uncontrolled, multicentre study on long term safety and efficacy of vonicog alfa in paediatric and adult subjects with severe VWD.

As a consequence, sections 4.1, 4.2, 4.8, 5.1 and 5.2 of the SmPC are updated. The Package Leaflet is updated accordingly. Version 6.0 of the RMP 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 10.4, to update the PI in accordance with the latest EMA excipients guideline, and to implement editorial changes to the PI.

Action: For adoption

The Committee discussed the issues identified in this application, relating to clinical aspects.
The Committee adopted a request for supplementary information with a specific timetable.

5.1.26. Xerava – Eravacycline - EMA/VR/0000265697

Paion Pharma GmbH

Rapporteur: Filip Josephson

Scope: Extension of indication to include treatment of complicated intra-abdominal infections (cIAI) from the age of 8 years and older for XERAVA, based on final results from study TP-434-028; this is a phase 1, open-label, multicentre study to determine the pharmacokinetics and safety of intravenous eravacycline in children with suspected or confirmed bacterial infection; As a consequence, sections 4.1, 4.2, 5.1, 5.2, and 6.6 of the

SmPC are updated. The Package Leaflet is 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 10.4.

Action: For adoption

The Committee discussed the issues identified in this application, relating to clinical aspects.

The Committee adopted a request for supplementary information with a specific timetable.

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

No items

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

No items

6. Medical devices

6.1. Ancillary medicinal substances - initial consultation

6.2. Ancillary medicinal substances – post-consultation update

No items

6.3. Companion diagnostics - initial consultation

6.3.1. In vitro diagnostic medical device - EMEA/H/D/006768

qualitative determination of antibodies to adeno-associated virus serotype 74 (AAVrh74) in human serum and/or plasma

Scope: Opinion

Action: For adoption

The CHMP was updated on discussions at the CAT.

The Committee endorsed the list of questions with a specific timetable as adopted by the CAT.

6.3.2. VENTANA HER2 Dual ISH DNA Probe Cocktail RxRx - In vitro diagnostic medical device - EMEA/H/D/006723

Roche Diagnostics GmbH; to determine HER2 gene status by enumeration of the ratio of the HER2 gene to Chromosome 17 by light microscopy

Scope: Opinion

Action: For adoption

List of questions adopted on 19.06.2025.

The Committee confirmed that all issues previously identified in this application had been addressed.

The Committee adopted a positive opinion by consensus together with the CHMP Assessment Report and translation timetable.

6.3.3. VENTANA HER2 (4B5) Rabbit Monoclonal Primary Antibody RxRx - In vitro diagnostic medical device - EMEA/H/D/006724

Roche Diagnostics GmbH; semi-quantitative detection of HER2 antigen by immunohistochemistry (IHC) in sections of formalin-fixed, paraffin-embedded breast carcinoma, gastric carcinoma, and biliary tract cancer

Scope: Opinion

Action: For adoption

List of questions adopted on 19.06.2025.

The Committee confirmed that all issues previously identified in this application had been addressed.

The Committee adopted a positive opinion by consensus together with the CHMP Assessment Report and translation timetable.

6.4. Companion diagnostics – follow-up consultation

No items

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)

No items

8. Pre-submission issues

8.1. Pre-submission issue

8.1.1. Acoziborole - Article 58 – H0006686

Treatment of Human African Trypanosomiasis (HAT or sleeping sickness) caused by *T.b. gambiense*

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

Action: For adoption

The CHMP agreed to the request for accelerated assessment and adopted the briefing note and Rapporteurs' recommendation on the Request for Accelerated Assessment.

8.2. Priority Medicines (PRIME)

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

The CHMP adopted the recommendations for PRIME eligibility.

The individual outcomes are listed in the PRIME Monthly Report on the EMA website, in the PRIME homepage, under Outcome of eligibility section.

9. Post-authorisation issues

9.1. Post-authorisation issues

9.1.1. Amvuttra - Vutrisiran - Orphan - EMEA/H/C/005852/II/0015

Alnylam Netherlands B.V.;

Rapporteur: Janet Koenig, Co-Rapporteur: Fátima Ventura, PRAC Rapporteur: Liana Martirosyan

Scope: Revised assessment report

Action: For adoption

Opinion adopted on 25.04.2025. Request for Supplementary Information adopted on 30.01.2025.

The CHMP adopted the revised assessment report.

9.1.2. Tecovirimat SIGA – Tecovirimat - EMA/S/0000248804

Siga Technologies Netherlands B.V.

Rapporteur: Jayne Crowe, PRAC Rapporteur: Martin Huber

Scope: Annual reassessment

Action: For adoption

In the evaluation of the annual re-assessment new efficacy data have emerged from recent clinical trials suggesting a lack of effectiveness in the treatment of mpox.

The findings from these emerging and forthcoming data need to be reviewed, taking into account all available data, to determine whether there is an impact on the benefit-risk balance in the authorised indications.

In view of the above, the EC initiated an Article 20 referral (EC) No 726/2004 and requests the CHMP to assess the above concerns and their impact on the benefit risk balance for the centrally authorised medicinal product Tecovirimat SIGA.

See 10.1.3

9.1.3. WS2780

Riltrava Aerosphere-EMEA/H/C/005311/WS2780/0017

Trixeo Aerosphere-EMEA/H/C/004983/WS2780/0024

AstraZeneca AB

Lead Rapporteur: Finbarr Leacy, Lead PRAC Rapporteur: Jan Neuhauser

Scope: Quality

Action: For adoption

Request for Supplementary Information adopted on 22.05.2025, 30.01.2025.

The Committee confirmed that all issues previously identified in this application had been addressed.

The Committee adopted a positive opinion by consensus together with the CHMP Assessment Report and translation timetable.

The summary of opinion was circulated for information.

The EMA public health communication was circulated for information.

9.1.4. Fluenz Tetra - Influenza vaccine (live attenuated, nasal) – EMEA/H/C/002617

AstraZeneca AB; Prophylaxis of influenza in individuals 24 months to less than 18 years.

Rapporteur: Christophe Focke, Co-Rapporteur: Ingrid Wang

Scope: Withdrawal of marketing authorisation

Action: For information

The CHMP noted the withdrawal of the marketing authorisation.

9.1.5. Volibris – Ambrisentan - EMA/VR/0000266441

Glaxosmithkline (Ireland) Limited

Rapporteur: Antonio Gomez-Outes

Scope: Update of sections 4.2 and 5.3 of the SmPC in order to update the recommendations for the paediatric population, based on non-clinical data. In addition, the MAH took the opportunity to introduce minor editorial and formatting changes to the PI.

Action: For adoption

The Committee discussed the issues identified in this application, relating to non-clinical and clinical aspects.

The Committee adopted a request for supplementary information with a specific timetable.

9.1.6. COMIRNATY - COVID-19 mRNA vaccine - EMA/VR/0000275515

BioNTech Manufacturing GmbH

Rapporteur: Filip Josephson

Scope: Quality

Action: For adoption

The Committee confirmed that all issues previously identified in this application had been addressed.

The Committee adopted a positive opinion by consensus together with the CHMP Assessment Report and translation timetable.

The summary of opinion was circulated for information.

9.1.7. BIMERVAX - COVID-19 vaccine (recombinant, adjuvanted) - EMA/VR/0000279224

Hipra Human Health S.L.

Rapporteur: Beata Maria Jakline Ullrich

Scope: Quality

Action: For adoption

The Committee discussed the issues identified in this application.

The Committee adopted a request for supplementary information with a specific timetable.

9.1.8. Spikevax - COVID-19 mRNA vaccine - EMA/VR/0000278795

Moderna Biotech Spain S.L.

Rapporteur: Jan Mueller-Berghaus

Scope: Quality

Action: For adoption

The Committee confirmed that all issues previously identified in this application had been addressed.

The Committee adopted a positive opinion by consensus together with the CHMP Assessment Report and translation timetable.

The summary of opinion was circulated for information.

9.1.9. Holoclar - ex vivo expanded autologous human corneal epithelial cells containing stem cells – ATMP - EMEA/H/C/002450/R/0058

Holostem S.r.l.

Rapporteur: Egbert Flory, CHMP coordinators: Jan Mueller-Berghaus and Paolo Gasparini

Scope: Revised opinion adopted via written procedure on 01.07.2025.

Action: For information

The CHMP noted the written procedure.

9.1.10. Evrysdi – Risdiplam – EMEA/H/C/005145

Roche Registration GmbH; treatment of spinal muscular atrophy (SMA)

Rapporteur: Fatima Ventura, Co-Rapporteur: Paolo Gasparini

Scope: DHPC letter and Communication plan for QD2025-240

Action: For adoption

The CHMP adopted the DHPC letter and the communication plan for QD2025-240.

9.1.11. Inrebic - Fedratinib - EMEA/H/C/005026/II/0027, Orphan

Bristol-Myers Squibb Pharma EEIG

Rapporteur: Peter Mol

Scope: Withdrawal of Type II variation

Action: For information

Request for Supplementary Information adopted on 27.02.2025.

The Committee noted the withdrawal of the Type II variation.

10. Referral procedures

10.1. Procedure for Centrally Authorised products under Article 20 of Regulation (EC) No 726/2004

10.1.1. Oxbryta - Voxelotor - EMEA/H/A-20/1538/C/004869/0014

Pfizer Europe MA EEIG

Referral Rapporteur: Patrick Vrijlandt, Referral Co- Rapporteur: Alexandre Moreau

Scope: Revised timetable, draft list of experts for AHEG

Action: For adoption

The EC initiated a procedure under Article 20 of Regulation (EC) No 726/2004 to assess the benefit-risk balance of Oxbryta in its authorised indication. The initiation of the review

follows an imbalance of deaths between voxelotor and placebo observed in clinical trials. The findings from these emerging safety data need to be further reviewed, taking into account all available data, to determine whether there is an impact on the benefit-risk balance of Oxbryta in its authorised indication.

List of outstanding issues adopted 22.05.2025, 12.12.2024. List of questions adopted on 29.07.2024

The CHMP adopted the revised timetable and draft list of experts for AHEG.

10.1.2. IXCHIQ - Chikungunya vaccine (live) - EMA/REF/0000269473

Valneva Austria GmbH

Referral PRAC Rapporteur: Gabriele Maurer

Scope: Opinion

Action: For adoption

Review of the benefit-risk balance following procedure triggered by the European Commission (EC) under Article 20 of Regulation (EC) No 726/2004 resulting from pharmacovigilance data; PRAC recommendation.

Based on the recommendation prepared by the PRAC, the Committee adopted a positive opinion by consensus.

10.1.3. Tecovirimat SIGA – Tecovirimat - EMA/REF/0000287477

Siga Technologies Netherlands B.V.

Referral Rapporteur: Jayne Crowe, Referral Co-Rapporteur: Vilma Petrikaite

Scope: Start of procedure, appointment of rapporteurs, list of questions, timetable

Action: For adoption

The European Commission (EC) initiated a procedure under Article 20 of Regulation (EC) No 726/2004 and requested the Agency/CHMP to assess the benefit-risk balance of Tecovirimat SIGA. The review was prompted by emerging data from clinical trials, which raised concerns about a potential lack of efficacy. These findings need to be reviewed in the context of all available data and their potential impact on the benefit-risk of Tecovirimat SIGA in its authorised indications. In addition, the EC requests the Agency/CHMP to give its opinion, as soon as possible, as to whether temporary measures are necessary to ensure the safe and effective use of this medicinal product.

The CHMP appointed as referral rapporteur Jayne Crowe (IE) and Vilma Petrikaite (LT) as referral Co-Rapporteur.

The CHMP agreed that no temporary measures were necessary at this stage. The CHMP considered at this point that a detailed assessment of further data from the trials raising concern of a possible lack of efficacy in the context of all available data on Tecovirimat SIGA was necessary, before making recommendations on its authorised use.

The CHMP adopted a list of questions with a procedural timetable.

10.2. Requests for CHMP Opinion under Article 5(3) of Regulation (EC) No 726/2004

10.2.1. Colistimethate sodium (CMS) – EMEA/H/A-5(3)/1524

Various MAHs

Referral Rapporteur: Janet Koenig, Referral Co-Rapporteur: Ewa Balkowiec Iskra

Scope: Revised list of outstanding issues

Action: For adoption

Review of the ratio of polymyxins E1 and E2 in colistin starting material and of the (sulfomethylation) composition profile of CMS finished product.

List of outstanding issues adopted on 21.03.2024, 22.02.2024, 22.05.2025. List of questions adopted on 22.06.2023.

The CHMP adopted a revised 2nd list of outstanding issues.

10.3. Procedure under Articles 5(2) and 10 of Regulation (EC) No 726/2004

No items

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

No items

10.5. Harmonisation - Referral procedure under Article 30 of Directive 2001/83/EC

No items

10.6. Community Interests - Referral under Article 31 of Directive 2001/83/EC

No items

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

No items

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

No items

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

No items

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

No items

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

No items

11. Pharmacovigilance issue

11.1. Early Notification System

July 2025 Early Notification System on envisaged CHMP/CMDh outcome accompanied by communication to the general public.

Action: For information

12. Inspections

12.1. GMP inspections

Information related to GMP inspections will not be published as it undermines the purpose of such inspections.

12.2. GCP inspections

Information related to GCP inspections will not be published as it undermines the purpose of such inspections.

12.3. Pharmacovigilance inspections

Information related to Pharmacovigilance inspections will not be published as it undermines the purpose of such inspections.

12.4. GLP inspections

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

No items

13.2. Innovation Task Force briefing meetings

Information related to briefing meetings taking place with applicants cannot be released at the present time as it is deemed to contain commercially confidential information.

No items

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

No items

13.4. Nanomedicines activities

No items

14. Organisational, regulatory and methodological matters

14.1. Mandate and organisation of the CHMP

14.1.1. Vote by Proxy

Jan Mueller-Berghaus (Co-opted member) gave proxy to Janet Koenig (DE) for the entire duration of the meeting.

Simona Badoi (RO) gave a proxy to Ewa Balkowiec-Iskra (PL) for the entire duration of the meeting.

Alar Irs (EE) gave a proxy to John Joseph Borg (MT) for the Tuesday session of the meeting.

14.1.2. CHMP membership

No items

14.2. Coordination with EMA Scientific Committees

14.2.1. Pharmacovigilance Risk Assessment Committee (PRAC)

List of Union Reference Dates and frequency of submission of Periodic Safety Update

Reports (EURD list) for July 2025

Action: For adoption

The CHMP adopted the EURD list.

14.2.2. Paediatric Committee (PDCO)

PIPs reaching D30 at July 2025 PDCO

Action: For information

The CHMP note the PIPs reaching D30 at the July 2025 PDCO.

Agenda from the PDCO meeting held on 22-25 July 2025

Action: For information

The CHMP noted the agenda.

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

14.3.1. Biologics Working Party (BWP)

Chair: Sean Barry, Vice-Chair: Andreea Barbu

Action: For adoption

The CHMP adopted the BWP reports.

14.3.2. Scientific Advice Working Party (SAWP)

Chair: Paolo Foggi

Report from the SAWP meeting held on 07-10 July 2025. Table of conclusions

Action: For information

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

14.3.3. The CHMP noted the update.Election of Oncology Working Party Vice-Chair

The position of vice-chair is currently being held by Olli Tenhunen (FI).

Action: For election

Nomination(s) received

The CHMP re-elected Olli Tenhunen (FI) as vice-chair of the Oncology Working Party.

14.3.4. Election of 3Rs Working Party Chair and Vice-Chair

Action: For election

Nomination(s) received

The CHMP elected Sonja Beken (BE) as chair, and Sarah Adler-Flindt (DE) as vice-chair of the 3Rs Working Party.

14.3.5. Election of CNSWP Vice-chair

Action: For election

Nomination(s) received

The elections were postponed and the deadline of the call for Central Nervous System Working Party vice-chair was extended.

14.3.6. Election of VWP Vice-chair

Action: For election

Nomination(s) received

The elections were postponed and the deadline of the call for Vaccines Working Party vice-chair was extended.

14.3.7. CVSWP Response to the CHMP request on indication wording

Discussion on the indication wording

Action: For discussion

The CHMP discussed the indication wording.

14.3.8. Nomination of CHMP representatives to the PCWP and HCPWP

Nomination of a CHMP representative (and alternate) for each working party for the mandate June 2025 to May 2028.

Nomination(s) received

Action: For endorsement

The CHMP nominated Ewa Bałkowiec-Iskra (PL) as representative and Fatima Ventura (PT) as alternate to the PCWP.

The CHMP nominated Kristina Nadrah (SI) as representative and Ingrid Wang (NO) as alternate to the HCPWP.

14.3.9. ICH Q3E draft Guideline on Extractables and Leachables - Step 2b

The ICH Q3E Expert Working Group has completed a draft guideline covering the assessment and control of extractables and leachables (E&L). The document is presented

for adoption for a 4-month public consultation.

Action: For information

The CHMP noted the update of the ICH Q3E draft Guideline on Extractables and leachables - Step 2b, which will be adopted at a later stage. [Post-meeting note: The document was adopted via written procedure on the 05th of August.]

14.4. Cooperation within the EU regulatory network

No items

14.5. Cooperation with International Regulators

No items

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

No items

14.7. CHMP work plan

No items

14.8. Planning and reporting

No items

14.9. Others

15. Any other business

15.1. AOB topic

15.1.1. GIREX rules

Analysis of requests for clock-stop extensions and feedback from GIREX.

Action: For discussion

The CHMP discussed the requests for clock-stop extensions and feedback from GIREX.

16. List of participants

List of participants including any restrictions with respect to involvement of members/alternates/experts following evaluation of declared interests for the 21-24 July 2025 CHMP meeting, which was held in-person.

An asterisk (*) after the role, in the second column, signals that the member/alternate attended remotely. Additional experts participated in the meeting, either in person or remotely.

Name	Role	Member State or affiliation	Outcome restriction following evaluation of e-DoI	Topics on agenda for which restrictions apply
Bruno Sepedes	Chair	Portugal	No restrictions applicable to this meeting	
Daniela Philadelphy	Member	Austria	No interests declared	
Christian Gartner	Alternate	Austria	No interests declared	
Christophe Focke	Member	Belgium	No restrictions applicable to this meeting	
Karin Janssen van Doorn	Alternate	Belgium	No interests declared	
Lyubina Racheva Todorova	Member	Bulgaria	No restrictions applicable to this meeting	
Gergana Lazarova	Alternate*	Bulgaria	No restrictions applicable to this meeting	
Margareta Bego	Member	Croatia	No interests declared	
Selma Arapovic Dzakula	Alternate	Croatia	No interests declared	
Emilia Mavrokordatou	Member	Cyprus	No interests declared	
Katerina Savvidou	Alternate*	Cyprus	No interests declared	
Tomas Radimersky	Member	Czechia	No interests declared	
Petr Vrbata	Alternate*	Czechia	No interests declared	
Thalia Marie Estrup Blicher	Member	Denmark	No interests declared	
Boje Kvorning Pires Ehmsen	Alternate	Denmark	No interests declared	
Alar Irs	Member	Estonia	No restrictions applicable to this meeting	
Edward Laane	Alternate	Estonia	No restrictions applicable to this meeting	
Outi Mäki-Ikola	Member (Vice-Chair)	Finland	No restrictions applicable to this meeting	
Johanna Lähteenvu	Alternate	Finland	No interests declared	
Alexandre Moreau	Member	France	No interests declared	
Jean-Michel Race	Alternate	France	No restrictions applicable to this meeting	
Janet Koenig	Member	Germany	No interests declared	
Martin Mengel	Alternate*	Germany	No interests declared	
Aris Angelis	Member	Greece	No participation in discussion, final deliberations and voting on:	4.3.1. Abrysvo - Respiratory syncytial virus vaccine (bivalent,

Name	Role	Member State or affiliation	Outcome restriction following evaluation of e-DoI	Topics on agenda for which restrictions apply
				recombinant) - EMA/X/000025805 1; 5.1.3. BESONSA - Inotuzumab ozogamicin - EMA/VR/00002573 10; 10.1.1. Oxbryta - Voxelotor - EMEA/H/A-20/1538/C/004869 /0014; 5.1.13. Keytruda - Pembrolizumab - EMA/VR/00002451 08; 5.1.17. Noxafil – Posaconazole - EMA/VR/00002633 60; 5.1.18. Recarbrio - Imipenem / Cilastatin / Relebactam - EMA/VR/00002650 89; 3.2.1. – Clesrovimab - EMEA/H/C/00649; 5.1.24. TEZSPIRE – Tezepelumab - EMA/VR/00002450 13; 9.1.3. WS2780 Riltavva Aerosphere- EMEA/H/C/005311 /WS2780/0017 Trixeo Aerosphere- EMEA/H/C/004983 /WS2780/0024; 9.1.4. Fluenz Tetra - Influenza vaccine (live attenuated, nasal) – EMEA/H/C/002617
Anastasia Mountaki	Alternate	Greece	No interests declared	
Robert Porszasz	Member*	Hungary	No restrictions applicable to this meeting	
Beata Maria Jakline Ullrich	Alternate	Hungary	No interests declared	
Hrefna Gudmundsdottir	Member	Iceland	No interests declared	
Hjalti Kristinsson	Alternate	Iceland	No interests declared	
Jayne Crowe	Member	Ireland	No interests declared	
Finbarr Leacy	Alternate	Ireland	No interests declared	
Paolo Gasparini	Member	Italy	No restrictions applicable to this meeting	

Name	Role	Member State or affiliation	Outcome restriction following evaluation of e-DoI	Topics on agenda for which restrictions apply
Maria Grazia Evandri	Alternate*	Italy	No restrictions applicable to this meeting	
Elita Poplavská	Member	Latvia	No restrictions applicable to this meeting	
Vilma Petrikaite	Member	Lithuania	No restrictions applicable to this meeting	
Larisa Gorobets	Alternate	Lithuania	No interests declared	
Martine Trauffler	Member	Luxembourg	No interests declared	
Alexandra Branchu	Alternate*	Luxembourg	No interests declared	
John Joseph Borg	Member	Malta	No restrictions applicable to this meeting	
Peter Mol	Member	Netherlands	No restrictions applicable to this meeting	
Patrick Vrijlandt	Alternate	Netherlands	No restrictions applicable to this meeting	
Ingrid Wang	Member	Norway	No interests declared	
Eva Skovlund	Alternate	Norway	No restrictions applicable to this meeting	
Ewa Balkowiec Iskra	Member	Poland	No restrictions applicable to this meeting	
Fatima Ventura	Member	Portugal	No restrictions applicable to this meeting	
Paulo Paixão	Alternate	Portugal	No restrictions applicable to this meeting	
Simona Badoi	Member*	Romania	No interests declared	
Dana Gabriela Marin	Alternate*	Romania	No interests declared	
Frantisek Drafi	Member	Slovakia	No restrictions applicable to this meeting	
Jana Klimasová	Alternate	Slovakia	No restrictions applicable to this meeting	
Kristina Nadrah	Member	Slovenia	No restrictions applicable to this meeting	
Carolina Prieto Fernandez	Member*	Spain	No interests declared	
Antonio Gomez-Outes	Alternate	Spain	No interests declared	
Kristina Dunder	Member	Sweden	No interests declared	
Filip Josephson	Alternate	Sweden	No interests declared	
Bruno Delafont	Co-opted member	France	No interests declared	
Carla Torre	Co-opted member	Portugal	No restrictions applicable to this meeting	
Jan Mueller-Berghaus	Co-opted member*	Germany	No interests declared	

Name	Role	Member State or affiliation	Outcome restriction following evaluation of e-DoI	Topics on agenda for which restrictions apply
Blanka Hirschlerova	Co-opted member	Czechia	No interests declared	
Sol Ruiz	Co-opted member	Spain	No interests declared	
Jana Schweigertová	Expert	Slovakia	No interests declared	
Nikola Gejgušová	Expert	Slovakia	No interests declared	
Mário Miguel Coelho da Silva Rosa	Expert	Portugal	No restrictions applicable to this meeting	
Ana Catarina Fonseca	Expert	Portugal	No restrictions applicable to this meeting	
Uta Buckpesch-Heberer	Expert	Germany	No interests declared	
Ger van Zandbergen	Expert	Germany	No restrictions applicable to this meeting	
Lukas Malte Aguirre Dávila	Expert	Germany	No restrictions applicable to this meeting	
Julia Katharina Maier	Expert	Germany	No interests declared	
Anja Schmidt	Expert	Germany	No interests declared	
Marion Haberkamp	Expert	Germany	No interests declared	
Christine Greiner	Expert	Germany	No interests declared	
Georgios Aislaitner	Expert	Germany	No interests declared	
Jeanette McCallion	Expert	Ireland	No interests declared	
Liam McDonough	Expert	Ireland	No interests declared	
Sandra Bright	Expert	Ireland	No interests declared	
Dhruva Teja Thurlapati	Expert	Ireland	No interests declared	
Gabriele Maurer	Expert	Germany	No interests declared	
Tihana Slezak	Expert	Croatia	No interests declared	
Hrvoje Rimac	Expert	Croatia	No participation in discussion, final deliberations and voting on:	4.3.5. Lojuxta – Lomitapide – EMA/X/0000258068; 5.1.12. Invokana – Canagliflozin – EMEA/H/C/002649 /II/0069
Lidija Prka	Expert	Croatia	No interests declared	
Višnja Drinovac Vlah	Expert	Croatia	No interests declared	
Melanie Ramberger	Expert	Austria	No interests declared	
Thomas Lang	Expert	Austria	No interests declared	
Bojana Divkovic	Expert	Austria	No interests declared	
Lisa Nika	Expert	Austria	No interests declared	
Sabrina Jenull	Expert	Austria	No interests declared	
Michael Jirout	Expert	Austria	No interests declared	
Silke Dorner	Expert	Austria	No interests declared	
Elina Rantala	Expert	Finland	No interests declared	
Paolo Foggi	Expert	Italy	No interests declared	
Pierre Demolis	Expert	Iceland	No interests declared	
Anna Vikerfors	Expert	Sweden	No interests declared	
Elisabeth Wischnitzki	Expert	Austria	No interests declared	
André Elferink	Expert	Netherlands	No interests declared	
Silvijus Abramavicius	Expert	Lithuania	No interests declared	
Hilke Zander	Expert	Germany	No interests declared	
Cristina Migali	Expert	Italy	No interests declared	

Name	Role	Member State or affiliation	Outcome restriction following evaluation of e-DoI	Topics on agenda for which restrictions apply
Jutta Dedorath	Expert	Germany	No interests declared	
Bruna Dekic	Expert	Germany	No interests declared	
Ulrike Hermes	Expert	Germany	No interests declared	
Susanna Hausmann	Expert	Germany	No interests declared	
Julian Paesler	Expert	Germany	No interests declared	
Peter van de Ven	Expert	Netherlands	No restrictions applicable to this meeting	
Thea Trijntje Klamer	Expert	Netherlands	No interests declared	
Ingrid Schellens	Expert	Netherlands	No interests declared	
Melissa van Tok	Expert	Netherlands	No participation in discussion, final deliberations and voting on:	3.2.1. - Clesrovimab - EMEA/H/C/006497 ; 5.1.13. Keytruda - Pembrolizumab - EMA/VR/00002451 08; 5.1.17. Noxafil - Posaconazole - EMA/VR/00002633 60; 5.1.18. Recarbio - Imipenem / Cilastatin / Relebactam - EMA/VR/00002650 89
Robert Pollmann	Expert	Germany	No interests declared	
Susanne Müller-Egert	Expert	Germany	No interests declared	
Christian Baarlink	Expert	Germany	No interests declared	
Jörg Engelbergs	Expert	Germany	No interests declared	
Simin Oveisi	Expert	France	No restrictions applicable to this meeting	
Céline Jumeau	Expert	France	No interests declared	
Muriel Uzzan	Expert	France	No interests declared	
Ramzi Mraidi	Expert	France	No restrictions applicable to this meeting	
Violette Dirix	Expert	Belgium	No restrictions applicable to this meeting	
Ingrid Bourges	Expert	Belgium	No restrictions applicable to this meeting	
Jean-Michel Dogné	Expert	Belgium	No restrictions applicable to this meeting	
Jean-Noël Talbot	Expert	France	No restrictions applicable to this meeting	
Violaine Closson Carella	Expert	France	No interests declared	
Agnes Mambole Dema	Expert	France	No interests declared	
Ilona G. Reischl	Expert	Austria	No restrictions applicable to this meeting	

Name	Role	Member State or affiliation	Outcome restriction following evaluation of e-DoI	Topics on agenda for which restrictions apply
Anna Mari Lone	Expert	Norway	No restrictions applicable to this meeting	
Ole Henrik Myrdal	Expert	Norway	No interests declared	
Rune Kjeken	Expert	Norway	No interests declared	
Sarah Lang	Expert	Germany	No interests declared	
Sarah Adler-Flint	Expert	Germany	No interests declared	
Ana Rita Lemos	Expert	Portugal	No restrictions applicable to this meeting	
Sabine Mayrhofer	Expert	Germany	No interests declared	
Christoph Furtmann	Expert	Germany	No restrictions applicable to this meeting	
Johanna de Groot	Expert	Netherlands	No interests declared	
Hemme Jacob Hijma	Expert	Netherlands	No restrictions applicable to this meeting	
Susanna Uiterwaal	Expert	Netherlands	No interests declared	
Illiana Meurs	Expert	Netherlands	No interests declared	
Viktoria Starokozhko	Expert	Netherlands	No restrictions applicable to this meeting	
Carlijn Litjens	Expert	Netherlands	No restrictions applicable to this meeting	

A representative from the European Commission attended the meeting

Meeting run with the help of EMA staff.

Experts were evaluated against the agenda topics or activities they participated in.

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/



24 July 2025
EMA/CHMP/243581/2025

Annex to 21-24 July 2025 CHMP Minutes

Pre-submission and post-authorisations issues

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A. PRE-SUBMISSION ISSUES

A.1. ELIGIBILITY REQUESTS

Report on Eligibility to Centralised Procedure for Adopted
July 2025: **For adoption**

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

Final Outcome of Rapporteurship allocation for Adopted
July 2025: **For adoption**

B. POST-AUTHORISATION PROCEDURES OUTCOMES

B.1. ANNUAL RE-ASSESSMENT OUTCOMES

B.1.1. ANNUAL REASSESSMENT FOR PRODUCTS AUTHORISED UNDER EXCEPTIONAL CIRCUMSTANCES

Livmarli - Maralixibat -
EMEA/H/C/005857/S/0019, Orphan
Mirum Pharmaceuticals International B.V.,
Rapporteur: Janet Koenig, PRAC Rapporteur:
Adam Przybylkowski
Request for Supplementary Information adopted
on 25.04.2025.

Positive Opinion adopted by consensus together
with the CHMP assessment report and
translation timetable.
The Marketing Authorisation remains under
exceptional circumstances

B.2. RENEWALS OF MARKETING AUTHORISATIONS OUTCOMES

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

B.2.2. Renewals of Marketing Authorisations for unlimited validity

B.2.3. Renewals of Conditional Marketing Authorisations

B.3. POST-AUTHORISATION PHARMACOVIGILANCE OUTCOMES

Signal detection

PRAC recommendations on signals adopted at
the PRAC meeting held on 07-10 July 2025
PRAC:

Signal of Progressive multifocal leukoencephalopathy	The CHMP adopted the PRAC recommendation.
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Varicella Vaccine (live); Measles, Mumps, Rubella and Varicella Vaccine (Live) - PROQUAD (CAP & NAP)

Rapporteur: Jan Mueller-Berghaus, Co-Rapporteur: Paolo Gasparini, PRAC
Rapporteur: Gabriele Maurer

PRAC recommendation on a variation

Action: For adoption

Signal of progressive multifocal leukoencephalopathy	The CHMP adopted the PRAC recommendation.
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Ciltacabtagene autoleucel, idecabtagene vicleucel, tisagenlecleucel – CARVYKTI, Abecma, Kymriah (CAP)

Rapporteur: multiple, Co-Rapporteur: multiple, PRAC Rapporteur: multiple

PRAC recommendation on a variation

Action: For adoption

PSUR procedures for which PRAC adopted a recommendation for variation of the terms of the MA at its July 2025 meeting:

B.5. TYPE II VARIATION, WORKSHARING PROCEDURE OUTCOMES

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

B.5.1. CHMP assessed procedures scope: Pharmaceutical aspects

Idacio - Adalimumab - EMEA/H/C/004475/II/0024/G Fresenius Kabi Deutschland GmbH, Rapporteur: Peter Mol, PRAC Rapporteur: Karin Bolin Request for Supplementary Information adopted on 24.07.2025, 27.03.2025.	Request for supplementary information adopted with a specific timetable.
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Pombiliti - Cipaglucosidase alfa - EMEA/H/C/005703/II/0019 Amicus Therapeutics Europe Limited, Rapporteur: Patrick Vrijlandt Opinion adopted on 03.07.2025. Request for Supplementary Information adopted on 25.04.2025.	Positive Opinion adopted by consensus on 03.07.2025.
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POTELIGEO - Mogamulizumab - EMEA/H/C/004232/II/0028/G, Orphan	Positive Opinion adopted by consensus on
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Kyowa Kirin Holdings B.V., Rapporteur: Peter Mol	26.06.2025.
Opinion adopted on 26.06.2025.	
Request for Supplementary Information adopted on 27.03.2025.	
WS2780	Positive Opinion adopted by consensus on
Riltrava Aerosphere-	24.07.2025.
EMEA/H/C/005311/WS2780/0017	
Trixeo Aerosphere-	See 9.1
EMEA/H/C/004983/WS2780/0024	
AstraZeneca AB, Lead Rapporteur: Finbarr Leacy, Lead PRAC Rapporteur: Jan Neuhauser	
Opinion adopted on 24.07.2025.	
Request for Supplementary Information adopted on 22.05.2025, 30.01.2025.	
B.5.2. CHMP assessed procedures scope: Non-Clinical and Clinical aspects	
Paxlovid - Nirmatrelvir / Ritonavir -	Positive Opinion adopted by consensus on
EMEA/H/C/005973/II/0059/G	10.07.2025.
Pfizer Europe MA EEIG, Rapporteur: Jean-Michel Race, "A grouped application consisting of:	
C.I.4: Update of section 4.5 of the SmPC in order to add drug-drug interaction information with albendazole based on the post-marketing data and literature and to update information on drug-drug interactions with methadone and ethinyl estradiol based on the literature; the Package Leaflet is updated accordingly.	
C.I.4: Update of section 4.5 of the SmPC in order to update information on drug-drug interactions with calcium channel antagonists based on the cumulative safety data and literature."	
Opinion adopted on 10.07.2025.	
Request for Supplementary Information adopted on 15.05.2025, 23.01.2025.	
WS2818	Negative Opinion adopted by consensus on
PecFent-	24.07.2025.
EMEA/H/C/001164/WS2818/0062	
Gruenthal GmbH, Lead Rapporteur: Janet Koenig, "Update of section 4.5 of the SmPC in order to add drug-drug interaction information between opioids and anticholinergics; the Package Leaflet is updated accordingly."	
Request for Supplementary Information adopted on 15.05.2025, 13.03.2025.	
Dengue Tetravalent Vaccine (Live, Attenuated) Takeda-	Positive Opinion adopted by consensus on

Qdenga-**EMEA/H/C/005155/WS2809/0022**

Takeda GmbH, Lead Rapporteur: Sol Ruiz,
"Update of section 4.8 of the SmPC in order to
add eye pain to the list of adverse drug
reactions (ADRs) with frequency uncommon
based on post-marketing data; the Package
Leaflet is updated accordingly. In addition, the
MAH took the opportunity to update the list of
local representatives in the Package Leaflet and
to introduce editorial changes to the PI."

Opinion adopted on 17.07.2025.

Request for Supplementary Information adopted
on 20.03.2025.

B.5.3. CHMP-PRAC assessed procedures**Sunosi - Solriamfetol -****EMEA/H/C/004893/II/0026**

Positive Opinion adopted by consensus on

10.07.2025.

Atnahs Pharma Netherlands B.V., Rapporteur:
Janet Koenig, PRAC Rapporteur: Julia Pallos,
"Update of sections 4.6 and 5.2 of the SmPC in
order to update information on lactation and
breast-feeding based on results from the post-
marketing lactation study JZP110-401 listed as
a category 3 study in the RMP. This was a Phase
4, open-label, single-dose study to evaluate the
PK of solriamfetol in the breast milk and plasma
of healthy postpartum women following oral
administration of a 150 mg solriamfetol tablet.
The Package Leaflet is updated accordingly. The
RMP version 1.3 has also been submitted."

Opinion adopted on 10.07.2025.

Request for Supplementary Information adopted
on 08.05.2025.

ZTALMY - Ganaxolone -**EMEA/H/C/005825/II/0004/G, Orphan**

Request for supplementary information adopted
with a specific timetable.

Immedica Pharma AB, Rapporteur: Peter Mol,
PRAC Rapporteur: Adam Przybylkowski, "A
grouped application comprised of 8 Type II
variations as follows:

1 Type II (C.I.4): Update of section 5.2 of the
SmPC in order to update ganaxolone metabolite
pattern at steady state based on re-analysis of
1042-TQT-1001 listed as a category 3 study in
the RMP to evaluate the ganaxolone steady-
state metabolite.

7 Type II (C.I.13): Submission of the final non-clinical study reports for the in vitro DDI potential and in vivo PK of the metabolite M17 listed as category 3 studies in the RMP.

The RMP version 1.2 has also been submitted. In addition, the MAH took the opportunity to introduce updates to the PI that reflect clarifications and typographical corrections, including to sections 4.2 and 4.4 of the SmPC." Request for Supplementary Information adopted on 24.07.2025, 27.03.2025, 25.07.2024, 11.04.2024.

**ZTALMY - Ganaxolone -
EMEA/H/C/005825/II/0015/G, Orphan**

Immedica Pharma AB, Rapporteur: Peter Mol, PRAC Rapporteur: Adam Przybylkowski, "A grouped application consisting of five Type II variations, as follows:

C.I.13: Submission of the final report from non-clinical study 1022-9241 listed as a category 3 study in the RMP. This is a 26-Week Toxicity Study of Ganaxolone Metabolite, M2, by Oral Gavage in the Sprague-Dawley rat with a 2-Week Recovery Period. The RMP version 3 has also been submitted.

C.I.13: Submission of the final report from non-clinical study 20447815 listed as a category 3 study in the RMP. This is a An Oral (Gavage) Study of the Effects of M2 (Ganaxolone Metabolite) Administration on Embryo/Fetal Development in CD (Sprague Dawley) IGS Rat. The RMP version 3 has also been submitted.

C.I.13: Submission of the final report from Weight of Evidence (WoE) assessment to evaluate the need for a 2-year carcinogenicity study in rats with GNX, listed as a category 3 study in the RMP.

C.I.13: Submission of the final report from WoE assessment to evaluate the need for a 2-year carcinogenicity study in rats with M2, listed as a category 3 study in the RMP.

C.I.13: Submission of the final report from WoE assessment to evaluate the need for a juvenile toxicity study with M2, listed as a category 3

Request for supplementary information adopted with a specific timetable.

study in the RMP. ”

Request for Supplementary Information adopted
on 10.07.2025, 13.03.2025.

B.5.4. PRAC assessed procedures

PRAC Led	Positive Opinion adopted by consensus on
Enbrel - Etanercept -	10.07.2025.

EMEA/H/C/000262/II/0255

Pfizer Europe MA EEIG, PRAC Rapporteur:
Monica Martinez Redondo, PRAC-CHMP liaison:
Antonio Gomez-Outes, “Update of the RMP
version 7.9 to remove the important risks of
“Aplastic Anaemia and Pancytopenia”,
“Congestive Heart Failure in Adult Subjects” and
“Acute Ischaemic Cardiovascular Events in
Adults Subjects” and the missing information
“Immunogenicity Profile and Related Clinical
Outcomes of Etanercept Manufactured using the
revised process in a Real-life Post-marketing
Setting” from the list of SCs. In addition, the
MAH took the opportunity to introduce minor
editorial and formatting changes to the PI as
well as to update the list of local representatives
in the Package Leaflet and align the PI with the
QRD version 10.4.”

Opinion adopted on 10.07.2025.

Request for Supplementary Information adopted
on 10.04.2025, 16.01.2025.

PRAC Led	Positive Opinion adopted by consensus on
Mimpara - Cinacalcet -	10.07.2025.

EMEA/H/C/000570/II/0076

Amgen Europe B.V., PRAC Rapporteur: Mari
Thorn, PRAC-CHMP liaison: Kristina Dunder,
“Submission of the final report from study
20180204 listed as a category 3 study in the
RMP. This is a non-interventional observational
registry study to evaluate the use and safety of
cinacalcet among paediatric patients with
secondary hyperparathyroidism (HPT).”

Opinion adopted on 10.07.2025.

Request for Supplementary Information adopted
on 13.03.2025.

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

B.5.9. Information on withdrawn type II variation / WS procedure

B.5.10. Information on type II variation / WS procedure with revised timetable

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 - EMA CERTIFICATION OF PLASMA MASTER FILES

Information related to plasma master files cannot be released at the present time as these contain commercially confidential information.

E.1. PMF Certification Dossiers

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

G. ANNEX G

G.1. Final Scientific Advice (Reports and Scientific Advice letters):

Information related to Scientific Advice cannot be released at the present time as these contain commercially confidential information.

G.2. PRIME

Some information related to PRIME cannot be released at the present time as these contain commercially confidential information.