



EUROPEAN MEDICINES AGENCY
SCIENCE MEDICINES HEALTH

17 July 2019
EMA/CAT/499958/2019
Inspections, Human Medicines Pharmacovigilance and Committees Division

Committee for Advanced Therapies (CAT)

Minutes of the meeting on 19-21 June 2019

Chair: Martina Schübler-Lenz; Vice-Chair: Ilona Reischl

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 these 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 CAT meeting reports once the procedures are finalised.

Of note, the minutes are a working document primarily designed for CAT 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

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 current meeting, the Committee Secretariat announced that no restriction in the involvement of meeting participants in upcoming discussions was identified.

Participants in this meeting 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.

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 and as included in the list of participants. All decisions taken at this meeting were made in the presence of a quorum of members (i.e. 22 or more members were present in the room). M Turner declared a potential conflict of interest for the product under agenda point 2.7.1 and 6.3.3 . No further new or additional interests or restrictions were declared.

The CAT chair informed the CAT that this will be the last meeting of the current civil society members, and that new members and alternates representing clinicians and patients' organisations have been appointed by the European Commission for a mandate of 3 years from 1 July 2019 (see 7.1.2). The CAT chaired thanked the current members and alternates for their contributions to the work of the CAT over the last 3 years.

The CAT chair welcomed the new member from Cyprus, Rafaella Pontou, who attended the CAT for the first time.

1.2. Adoption of agenda

The CAT agenda for 19-21 June 2019 meeting was adopted with two additions (2.12.3 Yescarta and 7.5.1 ICH – Non-clinical biodistribution studies for gene therapy products)

1.3. Adoption of the minutes

The CAT minutes for 22-24 May 2019 meeting with the incorporation of comments received from CAT members were adopted.

2. Evaluation of ATMPs

2.1. Opinions

No items

2.2. Oral explanations

No items

2.3. Day 180 list of outstanding issues

2.3.1. Viable T-cells - Orphan - EMEA/H/C/002397

Kiadis Pharma Netherlands B.V.; adjunctive treatment in haematopoietic stem cell transplantation (HSCT) for a malignant disease

Scope: List of Outstanding Issues (LoOIs)

Action: for adoption

List of Outstanding Issues adopted on 14.09.2018, 25.05.2018. List of Questions adopted on 08.09.2017.

The Rapporteurs presented their assessment of the responses to the second List of Outstanding Issues. CAT subsequently adopted the third List of Outstanding Issues.

CAT agreed with the proposal to organise an Ad-Hoc Expert Group to provide input to the CAT. CAT adopted the list of questions to the Ad-hoc expert group. A request for nomination of experts will be sent to CAT and CHMP.

CAT adopted the review timetable.

Post-meeting note: CHMP proposed amendments to the list of questions to the Ad-hoc expert group.

2.3.2. Onasemnogene abeparvovec - Orphan - EMEA/H/C/004750

Accelerated assessment

AveXis Netherlands B.V.; treatment of spinal muscular atrophy (SMA)

Scope: List of Outstanding Issues (LoOIs)

Action: for adoption

List of Questions adopted on 22.02.2019.

The Rapporteurs presented their assessment of the responses to the List of questions. CAT adopted the list of outstanding issues.

CAT agreed with the proposal to organise a Scientific Advisory Group (SAG, Neurology) to provide input to the CAT. The CAT adopted a list of questions for the experts in the SAG Neurology to address. A request for nomination of additional experts will be sent to CAT and CHMP.

The accelerated timetable was reverted to a normal timetable. CAT adopted the review timetable.

2.4. Day 120 list of questions

No items

2.5. Day 80 assessment reports

No items

2.6. Update on ongoing initial applications

No items

2.7. New applications

2.8. Withdrawal of initial marking authorisation application

No items

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

No items

2.10. GMP and GCP inspections requests

No items

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

2.11.1. Spherox - spheroids of human autologous matrix-associated chondrocytes - EMEA/H/C/002736/II/0005/G

CO.DON AG

Rapporteur: Lisbeth Barkholt, CHMP Coordinator: Kristina Dunder

Scope: safety: update of the product information to reflect the study results of the 36-month follow up data for trial cod 16 HS 13¹ and the final study report with 60-month follow-up data for trial cod 16 HS 14². Request for supplementary information (RSI).

Action: for adoption

Request for supplementary information adopted on 24.05.2019.

The assessment of the responses to the RSI was presented. CAT adopted a second RSI.

2.11.2. Spherox - spheroids of human autologous matrix-associated chondrocytes - EMEA/H/C/002736/II/0008

CO.DON AG

Rapporteur: Lisbeth Barkholt, CHMP Coordinator: Kristina Dunder

Scope: quality Request for supplementary information (RSI).

Action: for adoption

The assessment of the responses to the RSI was presented. The second RSI, as discussed and agreed at the BWP, was adopted.

2.11.3. YESCARTA - axicabtagene ciloleucel - Orphan - EMEA/H/C/004480/II/0006

Kite Pharma EU B.V.

Rapporteur: Jan Mueller-Berghaus, CHMP Coordinator: Jan Mueller-Berghaus

Scope: quality Opinion

¹ Study cod 16 HS 13, is a prospective, randomised, open label, multicentre Phase-III clinical trial to compare the efficacy and safety of the treatment with the autologous chondrocyte transplantation product co.don chondrosphere (ACT3D-CS) with microfracture in subjects with cartilage defects of the knee with a defect size between 1 and 4 cm²

² Study cod 16 HS 14, is a prospective, randomised, open label, multicentre Phase-II clinical trial to investigate the efficacy and safety of the treatment of large defects (4-10 cm²) with three different doses of the autologous chondrocyte transplantation product co.don chondrosphere (ACT3D-CS) in subjects with cartilage defects of the knee.

Action: for adoption

Request for Supplementary Information adopted on 17.04.2019.

The assessment of the responses to the RSI was presented. BWP discussed and agreed with the proposed change. The opinion was adopted.

2.11.4. YESCARTA - axicabtagene ciloleucel - Orphan - EMEA/H/C/004480/II/0008

Kite Pharma EU B.V.

Rapporteur: Jan Mueller-Berghaus, CHMP Coordinator: Jan Mueller-Berghaus

Scope: quality Request for supplementary information (RSI).

Action: for adoption

The assessment of the responses to the RSI was presented. The second RSI, as discussed and agreed at the BWP, was adopted.

2.12. Other Post-Authorisation Activities

2.12.1. Strimvelis - autologous CD34+ enriched cell fraction that contains CD34+ cells transduced with retroviral vector that encodes for the human ADA cDNA sequence - Orphan - EMEA/H/C/003854/ANX/004.1

Orchard Therapeutics (Netherlands) BV

Rapporteur: Sol Ruiz; PRAC Rapporteur: Menno van der Elst

Scope: clinical and PhV. From initial MAA: non-interventional PASS: In order to investigate the long term safety and efficacy of Strimvelis gene therapy, the MAH should conduct and submit the results of a long term prospective, non-interventional follow up study using data from a registry of patients with adenosine deaminase severe combined immunodeficiency (ADA-SCID) treated with Strimvelis. The MAH will follow up on the risk of immunogenicity, insertional mutagenesis and oncogenesis as well as hepatic toxicity. The MAH will review the occurrence of angioedema, anaphylactic reactions, systemic allergic events and severe cutaneous adverse reactions during the FU period, particularly in those patients who had unsuccessful response and received ERT or SCT. The MAH will also evaluate intervention-free survival.

The MAH shall plan to include regular progress reports of the registry in the PSUR and provide interim study reports every 2 years until the registry finishes. Interim registry reports to be submitted every 2 years. Final CSR - to be submitted after the 50th patient has 15 year follow-up visit; Q4 2037.

Action: for adoption

The outcome of the assessment by the PRAC Rapporteur of the first interim report on the non-interventional PASS was presented. The report was adopted by CAT.

2.12.2. Zalmoxis - nalotimagene carmaleucel - Orphan - EMEA/H/C/002801/R/0015

MolMed S.p.A

Rapporteur: Carla Herberts, Co-Rapporteur: Sol Ruiz, CHMP Coordinator: Paula Boudewina van Hennik, PRAC Rapporteur: Brigitte Keller-Stanislawski

Scope: 1 year renewal of Marketing Authorisation

Action: for adoption

Request for Supplementary Information adopted on 24.05.2019.

The assessment of the responses to the RSI was presented. The opinion on the renewal of the marketing authorisation was adopted.

2.12.3. YESCARTA - axicabtagene ciloleucel - Orphan - EMEA/H/C/004480

Kite Pharma EU B.V.

Rapporteur: Jan Mueller-Berghaus, CHMP Coordinator: Jan Mueller-Berghaus

Scope: feedback, initiated by the Rapporteur, on a question from the MAH .

Action: for discussion

The Rapporteur informed the CAT of a question from Kite Pharma . This is outside of a formal procedure.

3. Certification of ATMPs

Information related to this section cannot be released at the present time as it is deemed to contain commercially confidential information.

3.1. Opinion

No items

3.2. Day 60 Evaluation Reports

No items

3.3. New Applications

No items

4. Scientific Recommendation on Classification of ATMPs

A new procedure for appointment of CAT coordinators for ATMP classification started at the June 2019 CAT meeting. The new procedures will ensure a better sharing of workload amongst all CAT members and will increase the experience of the CAT members.

4.1. New requests – Appointment of CAT Coordinator

4.1.1. Autologous, *ex vivo* expanded, clonal neoantigen specific tumour infiltrating lymphocytes – H0005417

Intended for the treatment of solid tumours

Scope: appointment of CAT Coordinator and adoption of timetable

Action: for adoption

Following CAT member was appointed as CAT coordinator.

4.1.2. Autologous CD34+ cells transduced with lentiviral vector encoding human γ -globinG16D and short-hairpin RNA734 – H0005415

Intended for the treatment of moderate to severe Sickle Cell

Scope: appointment of CAT Coordinator and adoption of timetable

Action: for adoption

Following CAT member was appointed as CAT coordinator

4.1.3. Autologous tumour-infiltrating lymphocytes (TIL) – H0005414

Intended for the treatment of solid tumours

Scope: appointment of CAT Coordinator and adoption of timetable

Action: for adoption

Following CAT member was appointed as CAT coordinator

4.1.4. CD34+ haematopoietic stem/progenitor cells enriched with normal mitochondria derived from white blood cells from a related donor - H0005416

Intended for the treatment of non-inherited mtDNA deletion syndromes

Scope: appointment of CAT Coordinator and adoption of timetable

Action: for adoption

Following CAT member was appointed as CAT coordinator .

4.1.5. Purified recombinant adeno-associated viral vector serotype 2 (AAV2) encoding the complementary DNA (cDNA) of human Rab escort protein type 1 (REP1) – H0005418

Intended for the treatment of choroideremia (CHM)

Scope: appointment of CAT Coordinator and adoption of timetable

Action: for adoption

Following CAT member was appointed as CAT coordinator

4.2. Day 30 ATMP scientific recommendation

4.2.1. Modified Vaccinia Ankara-Bavarian Nordic- Brachyury (MVA-BN-Brachyury) and recombinant fowlpox virus (FPV-Brachyury) encoding the human brachyury gene and three human costimulatory molecules known as TRICOM (triad of costimulatory molecules): B7.1, intercellular adhesion molecule-1 (ICAM-1), and leukocyte function-associated antigen-3 (LFA-3) – H0005394

Intended for the treatment of chordoma

Scope: ATMP scientific recommendation

Action: for adoption

CAT discussed the ATMP classification report. CAT secretariat to send the draft scientific recommendation to the European Commission for comments by 5 July 2019.

The CAT recommendation will be considered final if no comments are received from the European Commission. The final report will be sent thereafter to the applicant.

4.2.2. Autologous CD34+ cells – H0005399

Intended for the treatment of no-option critical limb ischemia

Scope: ATMP scientific recommendation

Action: for adoption

CAT discussed the ATMP classification report. CAT secretariat to send the draft scientific recommendation to the European Commission for comments by 5 July 2019.

The CAT recommendation will be considered final if no comments are received from the European Commission. The final report will be sent thereafter to the applicant.

4.2.3. [Uncapped, non-coding ribonucleic acid – H0005400](#)

Intended for the treatment of adenoid cystic carcinoma, squamous cell carcinoma of the head and neck, melanoma and squamous cell carcinoma of the skin

Scope: ATMP scientific recommendation

Action: for adoption

CAT discussed the ATMP classification report.

CAT proposed to put the following question to the applicant.

The classification of this product will be further discussed at the July 2019 CAT, taking into account the feedback from the applicant.

4.2.4. [Messenger ribonucleic acid \(mRNA\) coding for coiled-coil domain-containing protein 40 \(CCDC40\) protein – H0005395](#)

Intended for the treatment of primary ciliary dyskinesia (PCD) caused by biallelic mutation in the CCDC40 gene

Scope: ATMP scientific recommendation

Action: for adoption

CAT discussed the ATMP classification report. CAT secretariat to send the draft scientific recommendation to the European Commission for comments by 5 July 2019.

The CAT recommendation will be considered final if no comments are received from the European Commission. The final report will be sent thereafter to the applicant.

4.2.5. [Autologous peripheral blood T cells CD4 and CD8 selected and CD3 and CD28 activated transduced with retroviral vector expressing anti-CD19 CD28/CD3-zeta chimeric antigen receptor – H0005396](#)

Intended for the treatment of various types of cancer

Scope: ATMP scientific recommendation

Action: for adoption

CAT discussed the ATMP classification report. CAT secretariat to send the draft scientific recommendation to the European Commission for comments by 5 July 2019.

The CAT recommendation will be considered final if no comments are received from the European Commission. The final report will be sent thereafter to the applicant.

4.3. [Day 60 revised scientific recommendation \(following list of questions\)](#)

No items

4.4. Finalisation of procedure

4.4.1. Human embryonic stem cell-derived Müller cells – H0005356

Intended for the treatment of primary open angle glaucoma

Scope: the European Commission raised no comments. Final ATMP scientific recommendation

Action: for information

The information was noted.

4.4.2. Allogeneic neonatal human cardiac progenitor cells – H0005357

Intended for the treatment of cardiac failure

Scope: the European Commission raised no comments. Final ATMP scientific recommendation

Action: for information

The information was noted.

4.4.3. Allogeneic human enucleated red cell therapy expressing Anabaena variabilis (Av) phenylalanine ammonia lyase (AvPAL) – H0005355

Intended for the treatment of phenylketonuria (PKU)

Scope: the European Commission raised no comments. Final ATMP scientific recommendation

Action: for information

The information was noted.

4.5. Follow-up and guidance

No items

5. Scientific Advice

Information related to this section cannot be released at the present time as it is deemed to contain commercially confidential information.

5.1. New requests – appointment of CAT Rapporteurs

5.2. CAT reports

5.3. List of Issues

5.4. Finalisation of SA procedures

6. Pre-Authorisation Activities

Information related to this section cannot be released at the present time as it is deemed to contain commercially confidential information.

6.1. Paediatric investigation plans

- 6.2. ITF briefing meetings in the field of ATMPs
- 6.3. Priority Medicines (PRIME) – Eligibility requests
 - 6.3.1. Month 0 - Start of the procedure
 - 6.3.2. Month 1 – Discussion of eligibility
 - 6.3.3. Month 2 – Recommendation of eligibility
 - 6.3.4. Ongoing support

7. Organisational, regulatory and methodological matters

7.1. Mandate and organisation of the CAT

7.1.1. CAT membership

Cyprus: Rafaella Pontou – new member. New membership mandate from 16 May 2019

Cyprus: Isavella Kyriakidou – new alternate. New membership mandate from 16 May 2019

Cyprus: Marina Ieridi – membership ended on 15 May 2019

Cyprus: Maria Vassiliou - membership ended on 15 May 2019

France: Nathalie Morgensztejn – new alternate. New membership started on 03 June 2019

Action: for information

The information was noted. The CAT chair welcomed Rafaella Pontou, who attended CAT for the first time.

7.1.2. Appointed members and alternates to represent clinicians and patients' associations to Committee for Advanced Therapies (CAT), July 2019 to June 2022

Scope: Commission decision dated 28 May 2019 (ref. C(2019) 3919) on the new appointment of representatives of civil societies to CAT for a mandate of three years, from 1 July 2019 to 30 June 2022

Patients' associations:

-Member: Kerstin Sollerbrant affiliated to the Swedish Childhood Cancer Fund;

-Alternate: Lydie Meheus affiliated to The Anticancer Fund.

-Member: Kieran Breen affiliated to the European Parkinson's Disease Association;

-Alternate: Roland Pochet affiliated to the Belgian Brain Council.

Clinicians:

-Member: Bernd Gansbacher from the European Society of Gene and Cell Therapy (ESGCT);

-Alternate: Birgitte Klindt Poulsen from University Hospital of Aarhus and Aalborg.

-Member: Alessandro Aiuti from Vita Salute Raffaele University;

-Alternate: Alessandra Renieri from University of Siena.

Action: for information

Note: the CAT Secretariat will organise an induction meeting for the new cohort of representatives of civil societies.

The information was noted. The CAT chaired thanked the current members and alternates representing the patients' associations and the clinicians for their contributions to the work of the CAT over the last 3 years.

7.1.3. Strategic Review & Learning meeting – joint CAT/Clinical trial facilitation group (CTFG), Bucharest, Romania, 13-14 June 2019

CAT: Simona Badoi

Scope: feedback from the SRLM meeting

Action: for discussion

Note: a half day of this SRLM was held jointly with the CTFG.

Topic postponed until the July CAT meeting.

7.2. Coordination with EMA Scientific Committees

7.2.1. Committee for Medicinal Products for Human Use (CHMP)

Scope: Summary of Outcomes (SoO) for the May 2019 meeting

Action: for information

The information was noted.

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

7.3.1. Meeting report from workshop with stakeholders on support to quality development in early access approaches (i.e. PRIME, breakthrough therapies), 26 November 2018

CAT experts: Matthias Renner, Marcel Hoefnagel

Scope: presentation on the draft revised workshop report following industry comments. Compiled industry comments and responses

Action: for discussion

The report from the workshop was presented. CAT members were asked to provide comments on the report by Friday 28 June 2019. Publication of the final report is expected during summer 2019.

7.4. Cooperation within the EU regulatory network

7.4.1. Environmental risk assessments of medicinal products containing/consisting of genetically modified organisms (GMOs) through the centralised procedure

Scope: presentation of the procedure

Action: for information

EMA presented the current procedure for the review of the Environmental Risk Assessment (ERA) of medicines containing GMOs through the centralised procedure. Feedback was provided on discussions on two issues arising from recent experience .

It was agreed to set-up a CAT drafting group to review the current procedure, which should be ready for adoption at next CAT meeting. Following CAT members will join this group: Margarida Menezes Ferreira, Carla Herberts, Rune Kjekken, Violaine Closson Carella and Jan Mueller Berghaus. The Commission representative will also join. Feedback from the discussions will be provided at the July CAT meeting.

7.4.2. European Commission's pharmaceutical committee on hospital exemption: upcoming discussions

Scope: Update on upcoming discussions in the Pharmaceutical Committee

Action: for information

The Commission representative informed CAT of the upcoming discussion about hospital exemption in the pharmaceutical committee.

The Guideline on GCP for ATMPs will also be presented at the Pharmaceutical Committee for adoption.

7.4.3. Principles for good manufacturing practice/viral vectors for genetically modified cells/advance therapy medicinal products

Scope: presentation of the outcome of the survey for viral vector inspections

Action: for information

EMA presented the outcome of a survey with GMP inspectors on inspections of viral vector manufacturing sites. A drafting group from the GMP inspectors working group will prepare a Questions and Answer document to harmonise the approach.

7.5. Cooperation with international regulators

7.5.1. ICH³ – Non-clinical biodistribution studies for gene therapy products

Scope: Appointment of CAT members / experts to join the ICH drafting group on non-clinical biodistribution studies for gene therapy products.

Action: for agreement

CAT was informed of the ICH proposal to develop the above mentioned Guideline. For EU, two experts have to be appointed. CAT appointed following CAT member as EU expert: Rune Kjekken. A second CAT member/expert will be nominated later.

7.6. CAT work plan

None

7.7. Planning and reporting

7.7.1. Planning estimates of forthcoming ATMP MAAs

Scope: Q2/2019 update of the business pipeline report for the human scientific committees

Action: for information

The information was noted.

7.8. Others

7.8.1. Global consultation on the review and update of the Changsha Communiqué on Xenotransplantation, 12 – 14 December 2018, Changsha, China

CAT: Ralf Tönjes, PEI-DE

Scope: feedback from the meeting

Action: for information

³ ICH: International Council for Harmonisation of Technical Requirements for Pharmaceuticals for Human Use

Ralf Tönjes provided feedback from the meeting which he attended as EMA/CAT representative. The update to the WHO document (Changsha Communique on Xenotransplantation) was presented.

Further to a question from the CAT chair, it was noted that very few member states have authorised clinical trials with xenotransplantation products.

8. Any other business

No items

Date of next CAT meeting:

17-19 July 2019

9. Explanatory notes

The Notes give a brief explanation of relevant agenda items and should be read in conjunction with the agenda.

Abbreviations / Acronyms

AAV: Adeno-Associated Virus

AR: Assessment Report

ATMP: Advanced Therapy Medicinal Product

BWP: Biologics Working Party

CAT: Committee for Advanced Therapies

CHMP: Committee for Medicinal Product for Human Use

COMP: Committee for Orphan Medicinal Products

CTFG: Clinical Trial Facilitation Group

DG: Drafting Group

EC: European Commission

ERA: Environmental Risk Assessment

FDA: Food and Drug Administration

FL: Final Letter

GCG: Guideline Consistency Group

GCP: Good Clinical Practice

GLP: Good Laboratory Practice

GMO: Genetically-modified organism

GMP: Good Manufacturing Practice

GTMP: Gene Therapy Medicinal Product

HTA: Health Technology Assessment Bodies

HSPC: Hematopoietic Stem and Progenitor Cells

ITF: Innovative Task Force

JR: Joint Report

LoOI: List of outstanding issues

LoQ: List of questions

MA: Marketing Authorisation

MAA: Marketing Authorisation Application

MAH: Marketing Authorisation Holder

MSC: Mesenchymal stem cells

PDCO: Paediatric Committee

PMDA: Pharmaceuticals and Medical Devices Agency (Japan)

PIP: Paediatric Investigation Plan

PL: Package leaflet

PRAC: Pharmacovigilance and Risk Assessment Committee #

PRIME: Priority Medicines

RMP: Risk Management Plan

RP: Reflection paper
 RSI: Request for supplementary information
 SAs: Scientific Advices
 SAG-O: Scientific Advisory Group Oncology
 SAWP: Scientific Advice Working Party
 SR: Summary Report
 SWP: Scientific Working Party
 SME: Small and medium size enterprises
 SmPC: Summary of Products Characteristics
 TT: Timetable

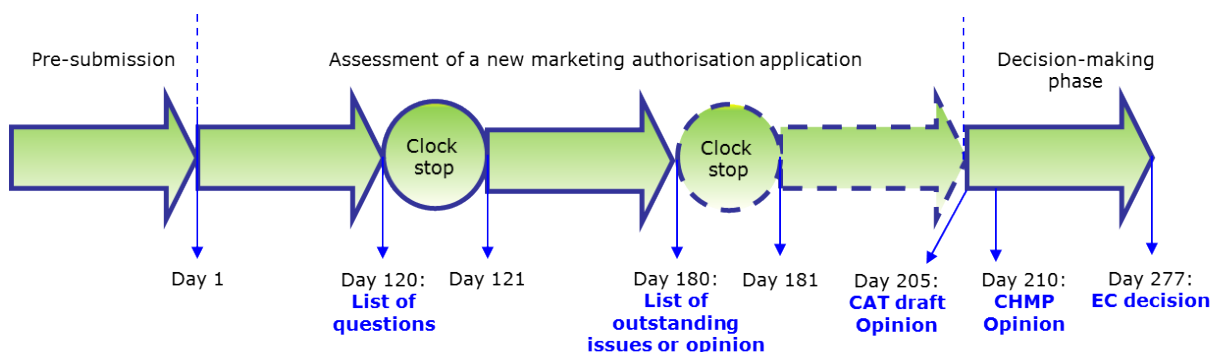
Evaluation of ATMPs (section 2)

This section lists applications for marketing authorisations of new Advanced Therapy Medicinal Products (ATMPs) that are to be discussed by the Committee. It also lists any ATMP related inspection requests (section 2.9) and Post-authorisation activities (section 2.10).

New applications (sections 2.1. to 2.12.)

Section 2.1 is for ATMPs nearing the end of the evaluation and for which the CAT is expected to adopt a draft [opinion](#) at this meeting on whether marketing authorisation should be granted. Once adopted, the CAT opinion is transmitted to the CHMP for final adoption. The CHMP opinion will be forwarded to the European Commission for a final legally binding decision valid throughout the EU. More information on the evaluation of ATMPs can be found [here](#).

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 CAT. The clock stop happens after day 120 and may also happen after day 180, when the CAT has adopted respectively a [Day 120 list of questions](#) (section 2.3) or a List of outstanding issues to be addressed by the company, which is listed in the agenda under sections 2.7 ([Ongoing evaluation procedures](#)). Section 2.7 also includes the CAT discussions at any other timepoint of the evaluation procedure of new applications.

Oral explanation (section 2.2.)

Prior to adoption of the CAT opinion, marketing authorisation applicants are normally invited to the CAT plenary meeting to address questions raised by the Committee.

Oral explanations normally relate to ongoing applications, but they can also relate to any other issue for which the CAT would like to discuss with company representatives in person.

Re-examination procedures (new applications) under article 9(2) of regulation no 726/2004 (section 2.6.)

This section lists applications for new marketing authorisation for ATMPs for which the applicant has requested a re-examination of the opinion previously issued by the CHMP. Similar to the initial evaluation of a marketing authorisation of an ATMP, CAT will adopt a draft re-examination opinion, which is transmitted to the CHMP for final adoption.

Withdrawal of applications (section 2.7.)

This section includes information on marketing authorisation applications that are withdrawn by the applicant. Applicants may decide to withdraw applications at any stage during the assessment and a CAT opinion will therefore not be issued. Withdrawals are included in the agenda for information or discussion, as necessary.

New applications (section 2.9.)

In this section, information is included on upcoming marketing authorisation applications for ATMPs, as well as information on appointment of Rapporteurs for new ATMP applications.

GMP and GCP Inspections Issues (section 2.10.)

This section lists inspections that are undertaken for ATMPs. 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).

Post-authorisation activities (section 2.12.)

This section lists type II variations, extension application according to Annex I of Reg. 1234/2008, re-examination procedures 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 and other issues concerning authorised medicines that are not covered elsewhere in the agenda such as annual reassessments, 5-year renewals, supply shortages, qualify defects. 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, will also be included here.

Certification of ATMPs (section 3)

This section includes the scientific evaluation by the CAT of quality and non-clinical data that small and medium-sized enterprises have generated at any stage of the ATMP development process. More information on the ATMP certification procedure can be found [here](#).

Scientific Recommendation on Classification of ATMPs (Section 4)

This section includes the scientific recommendation by the CAT on whether medicines based on genes, cells or tissues meet the scientific criteria that define ATMPs. More information on the ATMP classification procedure, including the outcomes of finalised classifications, can be found [here](#).

Scientific Advice (section 5)

This section includes all scientific advice given to companies during the development of an ATMP. Information related to the number of ATMP related scientific advices discussed by CAT can be found in the CAT Monthly reports. Further information on SAWP can be found [here](#).

Pre-Authorisation (section 6)

Paediatric Investigation Plan (PIP)

This section includes the discussion of an ATMP before a formal application for marketing authorisation

is submitted. These cases refer for example 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.

CAT contributes to the evaluation of a Paediatric Investigation Plan (PIPs) for ATMPs by the Paediatric Committee. These PIPs are included in this section of the Agenda.

ITF Briefing meeting in the field of ATMPs

This section refers to briefing meetings of the Innovation Task Force and International co-operations activities of the CAT

The Innovation Task Force (ITF) is a body set up to encourage early dialogue with applicants developing innovative medicines. Minutes of meetings with applicants developing ATMPs and of other ITF meetings of interest to the CAT are included in this section of the agenda. Further information on the ITF can be found [here](#).

Priority Medicines (PRIME)

This section includes the new requests for eligibility to PRIME for ATMPs under development, the discussions in CAT of these eligibility requests and the final recommendations for eligibility of ATMPs adopted by CHMP.

CAT will appoint one of its members as the CAT sponsor for each new ATMP eligibility request who will lead the CAT discussion based on the recommendation from the SAWP.

Organisational, regulatory and methodological matters (section 7)

This section includes topics related to regulatory and procedural guidance, CAT workplan, CAT meeting organisation (including CAT membership), planning and reporting, co-ordination with other committees, working parties and scientific advisory groups.

Furthermore, this section refers to the activities of the CAT drafting groups developing scientific guidelines for gene therapy medicinal products and for cell-based medicinal products, cooperation within the EU regulatory network and international regulators as well as direct interaction with interested parties. It also includes topics of scientific interest for the Committee that are not directly related to the work of the CAT drafting groups or CAT associated working parties.

Any other business (section 8)

This section is populated with miscellaneous topics not suitable under the previous headings.

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

10. List of participants

including any restrictions with respect to involvement of members / alternates / experts following evaluation of declared interests for the 19-21 June 2019 meeting.

Name	Role	Member state or affiliation	Outcome restriction following evaluation of e-DoI	Topics on agenda for which restrictions apply
Martina Schüssler-Lenz	Chair	Germany	No interests declared	
Ilona Reischl	Member	Austria	No interests declared	
Corina Spreitzer	Alternate	Austria	No restrictions applicable to this meeting	
Claire Beuneu	Member	Belgium	No interests declared	
Belaïd Sekkali	Alternate	Belgium	No interests declared	
Mirna Golemovic	Member	Croatia	No interests declared	
Petra Sokol	Alternate	Croatia	No interests declared	
Rafaella Pontou	Member	Cyprus	No interests declared	
Ondrej Palan	Alternate	Czech Republic	No interests declared	
Anne Pastoft	Member	Denmark	No interests declared	
Toivo Maimets	Member	Estonia	No interests declared	
Pille Saalik	Alternate	Estonia	No interests declared	
Heli Suila	Member	Finland	No interests declared	
Olli Tenhunen	Alternate	Finland	No interests declared	
Violaine Closson	Member	France	No interests declared	
Nathalie Morgensztejn	Alternate	France	No interests declared	
Jan Mueller-Berghaus	Member (CHMP co-opted member)	Germany	No interests declared	
Egbert Flory	Alternate (to CHMP representative)	Germany	No interests declared	
Angeliki Roboti	Alternate	Greece	No interests declared	
Balázs Sarkadi	Alternate	Hungary	No interests declared	
Niamh Curran	Alternate	Ireland	No interests declared	
Paolo Gasparini	Member	Italy	No interests declared	

Name	Role	Member state or affiliation	Outcome restriction following evaluation of e-Dol	Topics on agenda for which restrictions apply
Una Riekstina	Member	Latvia	No interests declared	
Romaldas Mačiulaitis	Member (CHMP member)	Lithuania	No restrictions applicable to this meeting	
Carla Herberts	Member	Netherlands	No interests declared	
Johannes Hendrikus Ovelgonne	Alternate	Netherlands	No interests declared	
Rune Kjekken	Member	Norway	No restrictions applicable to this meeting	
Dariusz Śladowski	Member	Poland	No restrictions applicable to this meeting	
Margarida Menezes-Ferreira	Alternate (to CHMP representative)	Portugal	No interests declared	
Lukas Slovak	Member	Slovakia	No interests declared	
Metoda Lipnik-Stangelj	Member	Slovenia	No interests declared	
Marcos Timón	Alternate (to CHMP representative)	Spain	No interests declared	
Lisbeth Barkholt	Member	Sweden	No interests declared	
John Johnston	Alternate	United Kingdom	No interests declared	
Marc Turner	Member	Healthcare Professionals' Representative	Interest declared	agenda points 2.7.1 and 6.3.3.2.
Bernd Gänsbacher	Member	Healthcare Professionals' Representative	No interests declared	
Willem Fibbe	Alternate	Healthcare Professionals' Representative	No restrictions applicable to this meeting	
Kieran Breen	Member	Patients' Representative	No restrictions applicable to this meeting	
Mariëtte Driessens	Member	Patients' Representative	No restrictions applicable to this meeting	
Erik Briers	Alternate	Patients' Representative	No restrictions applicable to this meeting	
Marcel Hoefnagel	Expert - in person*	CBG-MEB.NL		
Emmely de Vries	Expert - in person*	CBG-MEB.NL		
Marja van de Bovenkamp	Expert - in person*	CBG-MEB.NL		
Janneke van Leeuwen	Expert - in person*	CBG-MEB.NL		
Susanne Poley-Ochmann	Expert - in person*	PEI-DE		
Hans Hillege	Expert - via telephone*	CBG-MEB.NL		

Name	Role	Member state or affiliation	Outcome restriction following evaluation of e-Dol	Topics on agenda for which restrictions apply
Marcel Kwa	Expert - via telephone*	CBG-MEB.NL		
Ria Nibbeling	Expert - via telephone*	CBG-MEB.NL		
Berendina Maria van den Hoorn	Expert - via telephone*	CBG-MEB.NL		
Juliane Rau	Expert - via telephone*	PEI-DE		
Anke Zobywalsi	Expert - via telephone*	PEI-DE		
Maren Hammann	Expert - via telephone*	PEI-DE		
Benjamin Hofner	Expert - via telephone*	PEI-DE		
A representative from the European Commission attended the meeting.				
Meeting run with support from relevant EMA staff.				

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