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SCIENCE MEDICINES HEALTH

15 July 2020
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Human Medicines Division

Committee for Advanced Therapies (CAT)

Minutes of the meeting on 17-19 June 2020

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

Disclaimers

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

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

Note on access to documents

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



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1. Introduction

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

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. 21 or more members were present in the room). All decisions, recommendations and advice were agreed by consensus, unless otherwise specified.

The CAT chair welcomed the new alternate for Sweden, Maria Lüttgen and thanked the previous Swedish alternate, Björn Carlsson, for his contributions to the CAT over the last 9 years.

1.2. Adoption of agenda

The CAT agenda for 17-19 June 2020 meeting was adopted.

1.3. Adoption of the minutes

The CAT minutes for 18-20 May 2020 meeting 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

No items

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

2.7.1. Idecabtagene vicleucel - Orphan - EMEA/H/C/004662

Accelerated assessment

Celgene Europe BV; treatment of multiple myeloma

Scope: Timetable for assessment

Action: for adoption

The timetable was adopted.

2.8. Withdrawal of initial marketing 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. Kymriah - tisagenlecleucel - Orphan - EMEA/H/C/004090/II/0022/G

Novartis Europharm Limited

Rapporteur: Rune Kjeken

Scope: quality: Opinion

Action: for adoption

CAT noted the outcome of the BWP discussion. The opinion was adopted.

2.11.2. Yescarta - axicabtagene ciloleucel - Orphan - EMEA/H/C/004480/II/0015

Kite Pharma EU B.V.

Rapporteur: Jan Mueller-Berghaus

Scope: quality: Opinion.

Action: for adoption

Request for Supplementary Information adopted on 24.04.2020, 21.02.2020.

CAT noted the outcome of the BWP discussion. The opinion was adopted.

2.12. Extension applications

No items

2.13. Other Post-Authorisation Activities

2.13.1. Imlygic - talimogene laherparepvec - EMEA/H/C/002771/ANX/001.1

Amgen Europe B.V.

Rapporteur: Olli Tenhunen, CHMP Coordinator: Tuomo Lapveteläinen

Scope: clinical

From initial MAA:

The MAH should submit the preliminary results of Study 20120325 (a phase 2, multicenter, open-label, single-arm trial to evaluate the correlation between objective response rate and baseline intratumoral CD8+T-lymphocyte density in subjects with unresected stage IIIB to IVM1c melanoma treated with talimogene laherparepvec). Interim report

Action: for adoption

CAT agreed with the conclusions in the Rapporteur's assessment report

2.13.2. Yescarta - axicabtagene ciloleucel - Orphan - EMEA/H/C/004480/REC/009

Kite Pharma EU B.V.

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

Scope: quality

Action: for adoption

CAT agreed with the conclusions in the Rapporteur's assessment report. The post-authorisation measure is considered adopted.

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

Timetable:

-Start of the procedure:	19.06.2020
-Draft CAT co-ordinator's report:	03.07.2020
-ITF peer-review comments:	08.07.2020
-Revised scientific recommendation:	10.07.2020
-Adoption of scientific recommendation by CAT:	17.07.2020

4.1. New requests – Appointment of CAT Coordinator

4.1.1. Recombinant adeno-associated viral vector (serotype 8) carrying an optimised gene for human cyclic nucleotide gated channel subunit alpha 3 (CNGA3) protein – H0005726

Intended for the treatment of achromatopsia caused by mutations in the CNGA3 gene

Scope: appointment of CAT Coordinator and adoption of timetable

Action: for adoption

The CAT coordinator was appointed.

4.1.2. Irradiated allogeneic induced-pluripotent stem cells expressing pluripotent genes and cancer-specific embryonic neo-antigens – H0005108

Intended for the treatment malignant solid tumours including all epithelial cancers in sub-group type harbouring a stemness mesenchymal-like signature and haematopoietic malignancies

Scope: appointment of CAT Coordinator and adoption of timetable

Action: for adoption

The CAT coordinator was appointed.

4.1.3. Autologous naïve regulatory T cells transduced with a lentiviral vector encoding for a Chimeric Antigen Receptor (CAR) to recognize the HLA-A*02 antigen – H0005713

Intended for the prevention of immune-mediated graft rejection in HLA-A*02 mismatched renal transplantation

Scope: appointment of CAT Coordinator and adoption of timetable

Action: for adoption

The CAT coordinator was appointed.

4.1.4. [Live-attenuated, genetically modified Mycobacterium bovis expressing the gene coding for listeriolysin from Listeria monocytogenes – H0005714](#)

Intended for treatment of non-muscle invasive bladder cancer

Scope: appointment of CAT Coordinator and adoption of timetable

Action: for adoption

The CAT coordinator was appointed.

4.2. **Day 30 ATMP scientific recommendation**

4.2.1. [Allogeneic CD34+-enhanced cell suspension derived from umbilical cord blood – H0005712](#)

Intended for the treatment of patients with inherited metabolic disorders [cerebral adrenoleukodystrophy, Hurler syndrome] where haematopoietic stem cell transplant is indicated

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 6 July 2020.

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. [Aggregates of defined size of human embryonic stem cell derived insulin secreting pancreatic beta cells, encapsulated within an encapsulation device – H0005721](#)

Intended for the treatment of type I diabetes mellitus

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 6 July 2020.

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. [Homogenate of antlerogenic stem cells – H0005710](#)

Intended for the treatment of chronic obstructive pulmonary disease, bronchial asthma

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 6 July 2020.

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.

4.2.4. Autologous adipose-derived mesenchymal stem cells, cartilage lesions – H0005700

Intended for the treatment of cartilage lesions

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 6 July 2020.

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. Wharton's jelly derived mesenchymal cells, myelitis – H0005701

Intended for the treatment of myelitis

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 6 July 2020.

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.

4.2.6. Wharton's jelly derived mesenchymal cells, meningitis – H0005693

intended for the treatment of meningitis

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 6 July 2020.

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.

4.2.7. Wharton's jelly derived mesenchymal cells, meningomyelocele – H0005704

Intended for the treatment of meningomyelocele, myelomeningocele, spina bifida

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 6 July 2020.

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.

4.2.8. Wharton's jelly derived mesenchymal cells, cerebellum syndrome – H0005705

Intended for the treatment of cerebellum syndrome

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 6 July 2020.

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.

4.2.9. [Wharton's jelly derived mesenchymal cells, encephalitis - H0005706](#)

Intended for the treatment of encephalitis

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 6 July 2020.

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.

4.2.10. [Wharton's jelly derived mesenchymal cells, Krabbe disease - H0005707](#)

Intended for the treatment of Globoid cell leukodystrophy (Krabbe disease)

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 6 July 2020.

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.

4.2.11. [Wharton's jelly derived mesenchymal cells, osteoarthritis - H0005708](#)

Intended for the treatment of osteoarthritis

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 6 July 2020.

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.

4.2.12. [Wharton's jelly derived mesenchymal cells, spinal and bulbar muscular atrophy – H0005709](#)

Intended for the treatment of spinal and bulbar muscular atrophy

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 6 July 2020.

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)

4.3.1. Autologous adipose-derived mesenchymal stem cell, diabetic foot syndrome - H0005699

Intended for the treatment of diabetic foot syndrome

Scope: awaiting responses from the applicant to the LoQs. Revised ATMP scientific recommendation

Action: for adoption

Postponed, awaiting responses from the applicant

4.4. Finalisation of procedure

4.4.1. Recombinant adeno-associated viral vector (serotype 8) carrying an optimised gene for human cyclic nucleotide gated channel subunit beta 3 (CNGB3) protein - H0005013

Intended for the treatment of achromatopsia caused by mutations in the CNGB3 gene

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

Action: for information

The information was noted.

4.4.2. Genetically modified *Lactococcus lactis* strain , engineered to secrete human pro-insulin and human IL-10 - H0005671

Intended for the treatment of clinical recent-onset Type 1 diabetes mellitus

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

Action: for information

The information was noted.

4.4.3. Autologous CD34+ cells transduced with a lentiviral vector encoding a modified γ -globin gene - H0005672

Intended for the treatment of sickle cell disease (SCD) and β -thalassemia

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

Action: for information

The information was noted.

4.4.4. Human autologous hematopoietic stem cells transduced with a lentiviral vector containing codon-optimized cDNA encoding for functional human alpha galactosidase - H0005696

Intended for the treatment of Fabry disease

Scope: comments received from the European Commission. Final ATMP scientific recommendation

Action: for discussion

CAT discussion the comments received from the European Commission. It was agreed not to change the classification report.

4.4.5. [Human autologous hematopoietic stem cells transduced with a lentiviral vector containing codon-optimized cDNA encoding for functional human glucocerebrosidase - H0005698](#)

Intended for the treatment of Gaucher disease

Scope: comments received by the European Commission. Final ATMP scientific recommendation

Action: for discussion

See also under 4.4.4

4.4.6. [Wharton's jelly derived mesenchymal cells – H0005673](#)

Intended for the treatment of patients with COVID-19 infections

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

Action: for information

The information was noted.

4.4.7. [Wharton's jelly derived mesenchymal stem cell, COVID-19 - H0005674](#)

Intended for the treatment of patients with COVID-19 infections

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

Action: for information

The information was noted.

4.4.8. [Wharton's jelly derived mesenchymal stem cell, Optic atrophy - H0005688](#)

Intended for the treatment of optic atrophy

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

Action: for information

The information was noted.

4.4.9. [Wharton's jelly derived mesenchymal stem cell, IFAP syndrome - H0005689](#)

Intended for the treatment of patients with Ichthyosis follicularis with alopecia and photophobia (IFAP) syndrome

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

Action: for information

The information was noted.

4.4.10. Wharton's jelly derived mesenchymal stem cell, Bone marrow transplant rejection - H0005690

Intended for the treatment of bone marrow transplant rejection

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

Action: for information

The information was noted.

4.4.11. Wharton's jelly derived mesenchymal stem cell, Secondary graft failure - H0005691

Intended for the treatment of secondary bone marrow transplant failure/secondary graft failure

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

Action: for information

The information was noted.

4.4.12. Wharton's jelly derived mesenchymal stem cell, Progressive supranuclear palsy - H0005692

Intended for the treatment of progressive supranuclear palsy

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

Action: for information

The information was noted.

4.4.13. Wharton's jelly derived mesenchymal stem cell, Multiple system atrophy - H0005711

intended for the treatment of multiple system atrophy

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

Action: for information

The information was noted.

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

Timetable:

-Final Briefing Package:	28.06.2020
-Start of the procedure at SAWP:	03.07.2020
-CAT report due by:	08.07.2020
-CAT recommendation:	17.07.2020

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

No items

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

Timetable for assessment:

Procedure start:	11.06.2020
SAWP recommendation:	09.07.2020
CAT recommendation:	17.07.2020
CHMP adoption of report and final recommendation:	23.07.2020

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

Sweden: Maria Lüttgen – membership mandate to start on 15 June 2020

Sweden: Björn Carlsson – membership mandate to end on 14 June 2020

Action: for information

The information was noted. See 1.1.

7.2. Coordination with EMA Scientific Committees

No items

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

7.3.1. Guideline on registry-based studies – consultation of committees

Scope: introduction of the guideline to committees' members in advance of the written consultation planned in June-July 2020 (deadline: 31 July 2020)

Action: For information

CAT noted the presentation on the guideline on registry-based studies. CAT members were informed that the final draft of the guideline will be circulated for comments shortly and were asked to provide written comments on the final draft by end of July 2020.

Post-meeting comment: the EMA confirmed that the CTFG and the CTEG will be consulted on the draft guideline.

7.3.2. Scientific advice for ATMPs

Scope: new procedure for providing CAT input to SAWP on scientific advices for ATMPs

Action: for discussion

EMA presented a proposal for a new procedure to provide CAT input to SAWP on scientific advice for ATMPs. CAT agreed with a 3-months pilot to try out the new procedure, starting from September 2020.

7.3.3. Working Party with Patients' and Consumers' Organisations (PCWP) and Working Party with Healthcare Professionals' Organisations (HCPWP)

Scope: meeting Summary from the PCWP/HCPWP joint meeting, held on 3-4 March 2020

Action: for information

The information was noted.

7.4. Cooperation within the EU regulatory network

7.4.1. European Union Network-Pharmacovigilance Oversight Group (EU-POG)

CAT: John-Joseph Borg

Scope: call for nomination of a new CAT member to join the EU-POG. Mandate of the EU-POG

Action: for nomination of a CAT member

Note: former CAT member was Corina Spreitzer who resigned in March 2020

An overview of the mandate and activities of the EU-POG was provided. CAT members interested to join this group should inform CAT secretariat.

7.4.2. Commission initiative related to GMO for medicinal products

Scope: oral feedback from the European Commission

Action: for information

The European Commission representative presented the draft Regulation on the application of certain aspect of the GMO legislation to medicinal product against COVID-19. The explanatory memorandum of the Commission Proposal also includes the Commission's interpretation on the interplay between the GMO and medicinal product legislation in case of medicines used under compassionate use authorisation.

7.5. Cooperation with international regulators

7.5.1. ATMP cluster teleconference with FDA-USA, Health Canada and PMDA-Japan (30 April 2020)

CAT: Martina Schüssler-Lenz

Scope: feedback on the teleconference that took place on 30 April 2020

Action: for information

Following CAT members attended the ATMP cluster teleconference: Martina Schüssler-Lenz, Carla Herberts, Ilona Reischl, Pille Säälük. The EC representative also attended.

Feedback was provided from the discussion during the ATMP cluster of 30 April 2020.

CAT proposed that information from ATMP discussions in other clusters (e.g. blood products cluster, rare diseases cluster) is routinely provided to the CAT.

7.5.2. ATMP cluster teleconference with FDA-USA, Health Canada and PMDA-Japan (25 June 2020)

CAT: Martina Schüssler-Lenz

Scope: agenda for the teleconference to take place on 25 June 2020

Action: for discussion

CAT was informed that the planned ATMP cluster of 25 June 2020 has been postponed.

7.6. CAT work plan

None

7.7. Planning and reporting

7.7.1. Planning estimates of forthcoming ATMP MAAs

Scope: Q2/2020 update of the business pipeline report for the human scientific committees

Action: for information

A short presentation was given on the forthcoming ATMP marketing authorisations.

7.8. Others

7.8.1. European Commission's reflection process on the future of the legislation on medicinal products

Scope: Commission questionnaire (including a question on ATMPs)

Action: for information

The European Commission representative provided a short feedback on a questionnaire that has been launched on 16 June 2020.

<https://ec.europa.eu/info/law/better-regulation/have-your-say/initiatives/12421-Pharmaceutical-Strategy-Timely-patient-access-to-affordable-medicines/public-consultation>

The questionnaire includes one question on ATMPs developed in hospitals.

CAT was also informed that a stakeholders' workshop on the Pharma strategy will be held on 14-15 July 2020.

https://ec.europa.eu/newsroom/sante/newsletter-specific-archive-issue.cfm?archtype=specific&newsletter_service_id=327&newsletter_issue_id=23305&page=1&fullDate=Fri%2019%20Jun%202020&lang=default

8. Any other business

8.1. Participation to external meetings

8.1.1. Participation of CAT members/alternates as speakers or panellist to international conferences

Scope: criteria for participation to international conferences

Action: for discussion

CAT discussed the participation of conferences when CAT members are representing EMA or CAT. CAT members can still join conference on their own behalf / on behalf of their national agencies: members were reminded to include the standard disclaimers in their presentations that the views expressed are their personal opinion and not representing the views of the CAT / EMA.

The aim of the current exercise is to identify criteria that will ensure that CAT is represented at important meetings/conference. Transparency is important, so sometimes several CAT members receive invitations for the same meeting. CAT members were asked to provide a list of the most relevant scientific meetings or conference to attend; any invitation received can also be forwarded to the CAT secretariat for inclusion in the tracking table.

8.1.2. American Society of Gene & Cell Therapy (ASGCT)'s annual meeting, 11th May 2020, Boston MS (USA)

CAT: Jan Mueller-Berghaus

Scope: feedback on the presentation at the ASGCT pre-meeting workshop: 'Commercialization I Workshop'

Action: for information

Feedback was provided on the discussion in the above-mentioned workshop

Date of next CAT meeting:

15-17/07/2020

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

EU NTC: European Union Network Training Centre

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

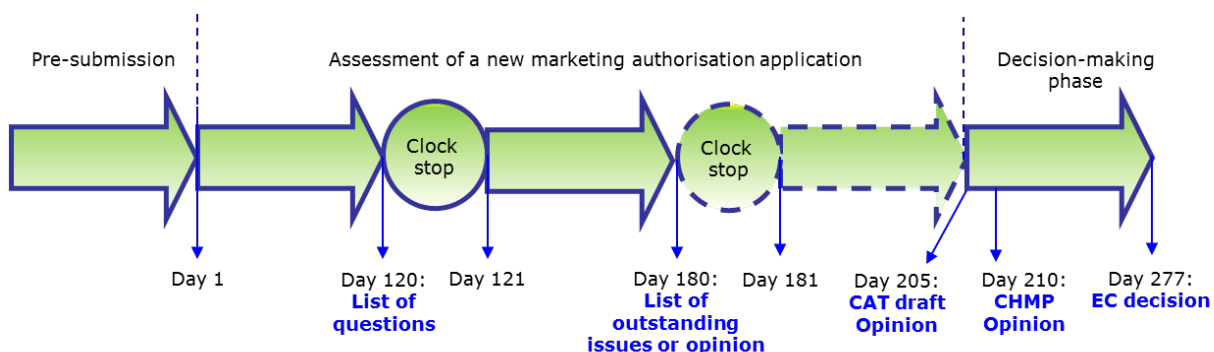
Evaluation of ATMPs (section 2)

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 17-19 June 2020 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	
Silke Dorner	Alternate	Austria	No interests declared	
Claire Beuneu	Member	Belgium	No interests declared	
Belaïd Sekkali	Alternate	Belgium	No interests declared	
Rozalina Kulaksazova	Member	Bulgaria	No interests declared	
Mirna Golemovic	Member	Croatia	No interests declared	
Rafaella Pontou	Member	Cyprus	No interests declared	
Ivana Haunerova	Member	Czech Republic	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	
Violaine Closson	Member	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 Rompoti	Alternate	Greece	No interests declared	
Katalin Lengyel	Member	Hungary	No interests declared	
Maura O'Donovan	Member	Ireland	No interests declared	
Paolo Gasparini	Member	Italy	No interests declared	
Una Riekstina	Member	Latvia	No interests declared	
Romaldas Mačiulaitis	Member (CHMP member)	Lithuania	No interests declared	
Guy Berchem	Member (to CHMP representative)	Luxembourg	No restrictions applicable to this meeting	
John J. Borg	Member (CHMP member)	Malta	No interests declared	
Anthony Samuel	Alternate (to CHMP representative)	Malta	No interests declared	

Name	Role	Member state or affiliation	Outcome restriction following evaluation of e-DoI	Topics on agenda for which restrictions apply
Carla Herberts	Member	Netherlands	No interests declared	
Johannes Hendrikus Ovelgonne	Alternate	Netherlands	No interests declared	
Rune Kjeklen	Member	Norway	No restrictions applicable to this meeting	
Maja Sommerfelt Grønvold	Alternate	Norway	No restrictions applicable to this meeting	
Dariusz Śladowski	Member	Poland	No restrictions applicable to this meeting	
Maria Isabel Borba Vieira	Alternate (to CHMP representative)	Portugal	No interests declared	
Simona Badoi	Member	Romania	No interests declared	
Lukas Slovak	Member	Slovakia	No interests declared	
Alexandra Padova	Alternate	Slovakia	No interests declared	
Metoda Lipnik-Stangelj	Member	Slovenia	No interests declared	
Sol Ruiz	Member (CHMP co-opted member)	Spain	No interests declared	
Marcos Timón	Alternate (to CHMP representative)	Spain	No interests declared	
Lisbeth Barkholt	Member	Sweden	No interests declared	
Maria Luttgen	Alternate	Sweden	No restrictions applicable to this meeting	
Bernd Gänsbacher	Member	Healthcare Professionals' Representative	No interests declared	
Alessandro Aiuti	Member	Healthcare Professionals' Representative	No restrictions applicable to this meeting	
Kerstin Sollerbrant	Member	Patients' Representative	No interests declared	
Lydie Meheus	Alternate	Patients' Representative	No interests declared	
Kieran Breen	Member	Patients' Representative	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
Roland Pochet	Alternate	Patients' Representative	No interests declared	
	Observer/Alternate	European Directorate for the Quality of Medicine & HealthCare(EDQM)	No interests declared	
Barbara Bonamassa	Expert	AIFA-IT		
Gloria María Palomo Carrasco	Expert	AEMPS-ES		
Leire Aguado	Expert	AEMPS-ES		
Celia Cerrato Rivera	Expert	AEMPS-ES		
Blanca García Ochoa	Expert	AEMPS-ES		
Babs Fabriek	Expert	CBG-MEB		

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.