

20 October 2022 EMA/CAT/841175/2022 Human Medicines Division

CAT quarterly highlights and approved ATMPs

October 2022

This report provides information on ATMP approvals and extension of indications of authorised ATMPs, as well as statistical data on product-related activities.

The period covered by this report is: August - October 2022

Advanced therapy medicinal products approvals

During its plenary meeting of October 2022, CAT adopted a positive draft opinion for **Ebvallo** (tabelecleucel) for the following indication: treatment of Epstein-Barr virus positive post-transplant lymphoproliferative disease (EBV+ PTLD). Based on the assessment of the CAT, the Committee for Medicinal Products for Human Use (CHMP) adopted a positive opinion, recommending the granting of a marketing authorisation under exceptional circumstances for the medicinal product Ebvallo. More information on Ebvallo can be found in the <u>Summary of opinion</u>.

Extension of indication of authorised ATMPs

During its plenary meeting of September 2022, CAT adopted an extension of indication for **Yescarta** to include the treatment of adult patients with diffuse large B-cell lymphoma (DLBCL) and high-grade B-cell lymphoma (HGBL) that relapses within 12 months from completion of, or is refractory to, first-line chemoimmunotherapy.

Overview of product-related activities

The Committee discussed ongoing evaluation procedures for ATMPs and other related procedures as summarised in the following tables:

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Initial Evaluation of Marketing Authorisation Applications (MAA) for ATMP						
	2009-2020	2021	2022*	Total		
Submitted MAAs	32	3	1	36		
Positive draft Opinion	18 ⁱ	2	5	25*		
Negative draft opinions	4 ^{i,ii,iii}	0	0	4		
Withdrawals	8 ^{ii, iv}	0	۱v	9		
Ongoing MAAs				2		

Corresponding to 24 ATMPs (see List of authorised ATMPs)

One negative draft opinion and two positive draft opinions for the Glybera

"Negative draft opinion and withdrawal for the Cerepro

ⁱⁱⁱ Two negative draft opinions for Heparesc ^{iv} Luxceptar, Roctavian, Artobend

* Sitoiganap

Variations (Type II) for authorised ATMP							
	2009-2020 2021 2022* Total						
Positive opinion	78	32	37	147			

Scientific recommendation on advanced therapy classification ¹							
2009-2020 2021 2022* Total							
Submitted	489	66	36	591			
Adopted	483	61	39	583			

Scientific advice procedure for ATMPs							
2009-2020 2021 2022* Total							
Number of procedures	442	64	40	546			

PRIME ² Eligibility for ATMPs								
2016-2020 2021 2022* Total								
Discussed	91	14	7	112				
Granted	39	7	3	49				

* Period: January - October 2022

¹ More information on the scientific recommendation on advanced therapy classification and the summaries of ATMP classification can be found on the ATMP classification webpage.

It is stressed that the scientific recommendation on advanced therapy classification does not amount to any endorsement of the plausibility of the product, including the mode of action or therapeutic indication(s) claimed by the applicant. ² PRIority MEdicines (PRIME) scheme. More information can be found at the PRIME webpage.

List of authorised ATMPs

NAME	Type of ATMP	Authorisation Date	Orphan	PRIME ³	Comment
Chondrocelect	ТЕР	5/10/2009	No	No	MA withdrawn July 2016
Glybera	GTMP	25/10/2012	Yes	No	MA not renewed (MA ended Oct. 2017)
MACI	TEP, combined ATMP	27/06/2013	No	No	MA not renewed (MA ended June 2018)
Provenge	СТМР	6/09/2013	No	No	MA withdrawn May 2015
Holoclar	ТЕР	17/02/2015	Yes	No	
Imlygic	GTMP	16/12/2015	No	No	
Strimvelis	GTMP	26/05/2016	Yes	No	
Zalmoxis	СТМР	18/08/2016	Yes	No	MA withdrawn Oct. 2019
Spherox	TEP	10/07/2017	No	No	
Alofisel	СТМР	23/03/2018	Yes	No	
Yescarta	GTMP	23/08/2018	Yes	Yes	
Kymriah	GTMP	23/08/2018	Yes	Yes	
Luxturna	GTMP	22/11/2018	Yes	No	
Zynteglo	GTMP	29/05/2019	Yes	Yes	MA withdrawn March 2022
Zolgensma	GTMP	18/05/2020	Yes	Yes	
Libmeldy	GTMP	17/12/2020	Yes	No	
Tecartus	GTMP	14/12/2020	Yes	Yes	
Skysona	GTMP	16/07/2021	Yes	Yes	MA withdrawn Nov. 2021
Abecma	GTMP	18/08/2021	Yes	Yes	
Breyanzi	GTMP	4/04/2022	No	Yes	
Carvykti	GTMP	25/05/2022	Yes	Yes	

³ PRIME (PRIority MEdicines scheme) was set up in March 2016 to provide early and enhanced scientific and regulatory support to medicines that have the potential to significantly address patients' unmet medical needs.

NAME	Type of ATMP	Authorisation Date	Orphan	PRIME ³	Comment
Upstaza	GTMP	18/07/2022	Yes	No	
Roctavian	GTMP	24/08/2022	Yes	No	
Ebvallo	СТМР	Positive opinion October 2022	Yes	Yes	Commission decision pending

More information on authorised products can be found on: <u>www.ema.europa.eu</u> (type in the product name in the search box)

<u>Abbreviations</u>: ATMP: advanced therapy medicinal product; GTMP: gene therapy medicinal product; CTMP: cell therapy medicinal product; TEP: tissue engineered product; MA: Marketing authorisation

