



EUROPEAN MEDICINES AGENCY
SCIENCE MEDICINES HEALTH

19 January 2026
EMA/5869/2026
Human Medicines Division

Monthly statistics report: December 2025

Medicinal products for human use (cumulative figures for the year to date)

This document provides current information related to the volume and evaluation of marketing authorisation and post-authorisation applications for medicinal products for human use received by the European Medicines Agency.

The purpose is only to provide ongoing factual information. Commentaries and analysis are provided in the Agency's annual reports.

Official address Domenico Scarlattilaan 6 • 1083 HS Amsterdam • The Netherlands

Address for visits and deliveries Refer to www.ema.europa.eu/how-to-find-us

Send us a question Go to www.ema.europa.eu/contact **Telephone** +31 (0)88 781 6000

An agency of the European Union



Table 1. Pre-authorisation: Marketing-authorisation applications*

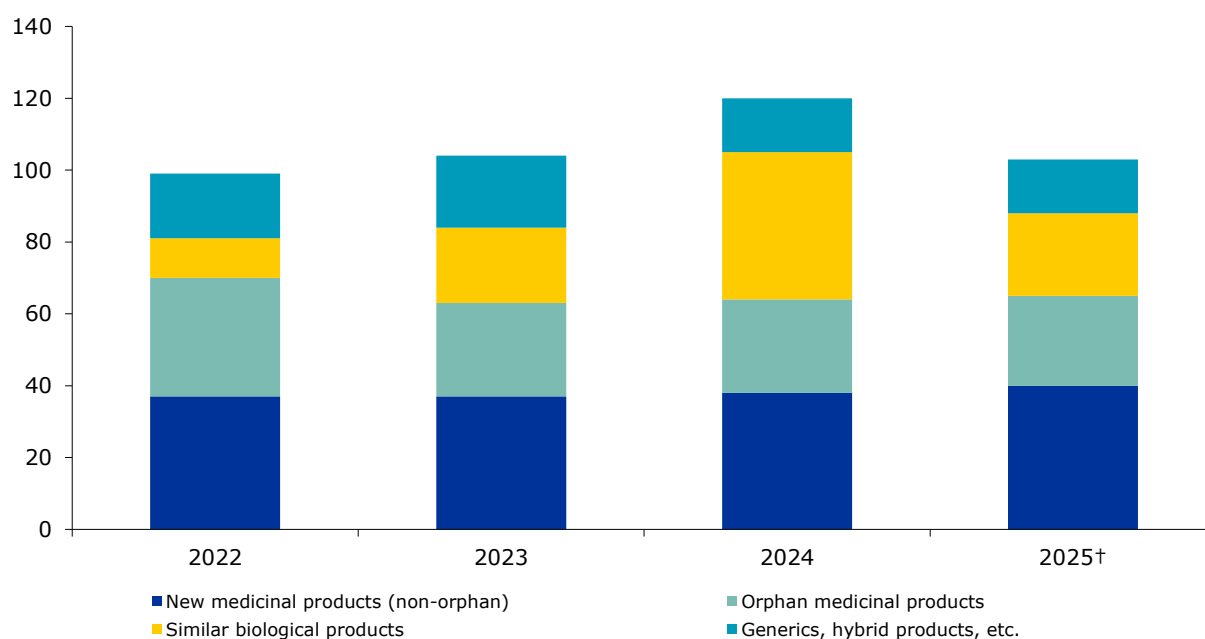
	2022		2023		2024		2025 [†]	
	Started	Finalised	Started	Finalised	Started	Finalised	Started	Finalised
Non-orphan medicinal products								
New products	35	27	35	26	33	44	37	30
Advanced-therapy medicinal products	0	0	1	0	2	1	1	1
Paediatric-use (PUMA) products	2	0	1	2	3	2	2	1
Well-established use, abridged, hybrid and informed consent products	3	7	4	4	3	2	6	3
Generic products	15	23	16	14	12	17	9	10
Similar biological products	11	10	21	8	41	28	23	41
Sub-total product applications	66	67	78	54	94	94	78	86
Orphan medicinal products[◇]								
New products	32	19	23	25	21	25	23	20
Advanced-therapy medicinal products	1	6	3	1	5	0	2	5
Total product applications	99	92	104	80	120	119	103	111

* Finalised applications exclude applications withdrawn prior to opinion.

[†] Figures for the current year are cumulative, year to date. Figures for preceding years are totals for the year.

[◇] These figures reflect the orphan status of the medicinal products at the time of the CHMP opinion. EMA's Committee for Orphan Medicinal Products (COMP) then assesses whether the orphan designation should be maintained.

Marketing authorisation application evaluations started by type of application



[†] Figures for the current year are cumulative, year to date. Figures for preceding years are totals for the year.

Table 2. Pre-authorisation: Outcome of the evaluation of marketing authorisation applications*

	2022	2023	2024	2025 [†]
Positive opinions (total 1 st opinions)	89	76	113	102
- new active substance (NAS)**		40	46	36
- conditional marketing authorisation**	9	7	8	8
- under exceptional circumstances**	5	1	4	2
- after accelerated assessment**	5	3	3	3
Negative opinions	3	4	6	9
Applications withdrawn prior to 1 st opinion ^{††}	11	15	9	14
Applications withdrawn after a 1 st opinion (e.g. during re-examination) ^{††}		3	1	6
Re-examinations requested	2	4	5	9
Re-examination - Positive opinions	0	1	1	2

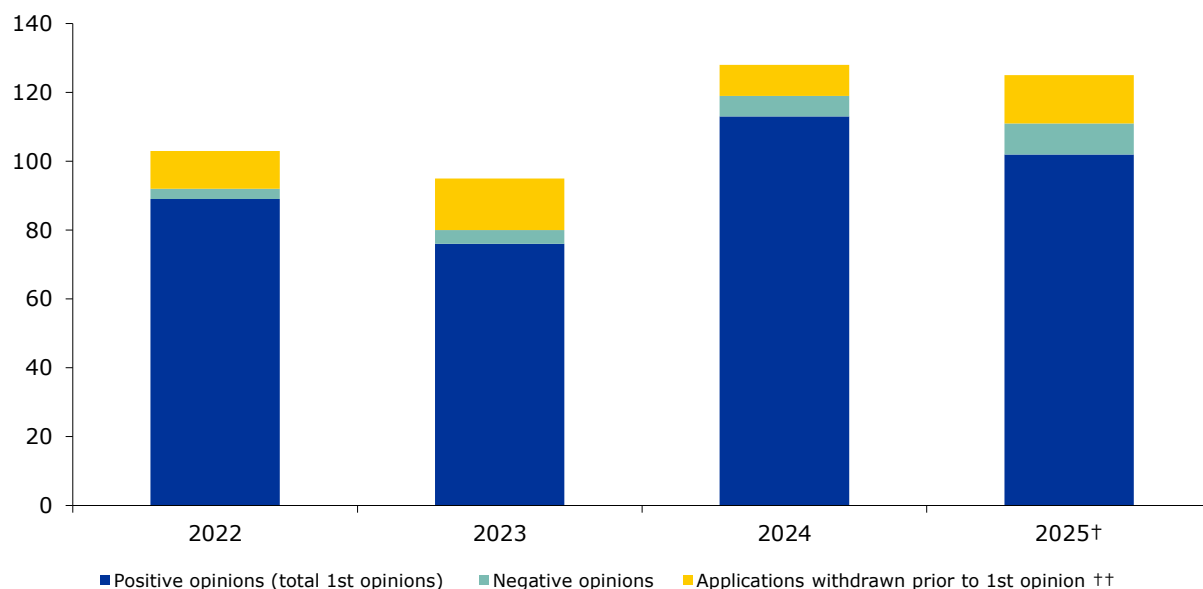
* Applicants can request a re-examination. The first five rows present the outcome of the evaluation before a re-examination (or a re-consideration). The final row shows the number of changes from a negative to a positive opinion following a re-examination or a re-consideration.

** Included in the figures for positive opinions. Duplicate products, if any, are included in the figures.

[†] Figures for the current year are cumulative, year to date. Figures for preceding years are totals for the year.

^{††} Date as per withdrawal letter date. It may first be communicated at the following month's CHMP meeting.

Pre-authorisation: Outcome of the evaluation of marketing authorisation applications



[†] Figures for the current year are cumulative, year to date. Figures for preceding years are totals for the year.

^{††} Date as per withdrawal letter date. It may first be communicated at the following month's CHMP meeting.

Table 3. Scientific services

	2022		2023		2024		2025 [†]	
	Started	Finalised	Started	Finalised	Started	Finalised	Started	Finalised
Compassionate-use opinions	0	0	0	0	0	0	0	0
Art. 58 (WHO) scientific opinions	1	3	0	1	1	0	2	2
Opinions on Companion Diagnostics medical devices (CDx)	4	3	9	8	11	10	12	8
Opinions on ancillary medicinal substances in medical devices*	2	0	0	2	3	1	1	2
Plasma master file (includes initial certification, variations and annual re-certification)	17	23	18	22	20	19	22	20

* Consultation in accordance with Council Directive 93/42/EEC concerning medical devices as amended by Directive 2000/70/EC as regards medical devices incorporating stable derivatives of human blood or plasma and Directive 2001/14/EC.

[†] Figures for the current year are cumulative, year to date. Figures for preceding years are totals for the year.

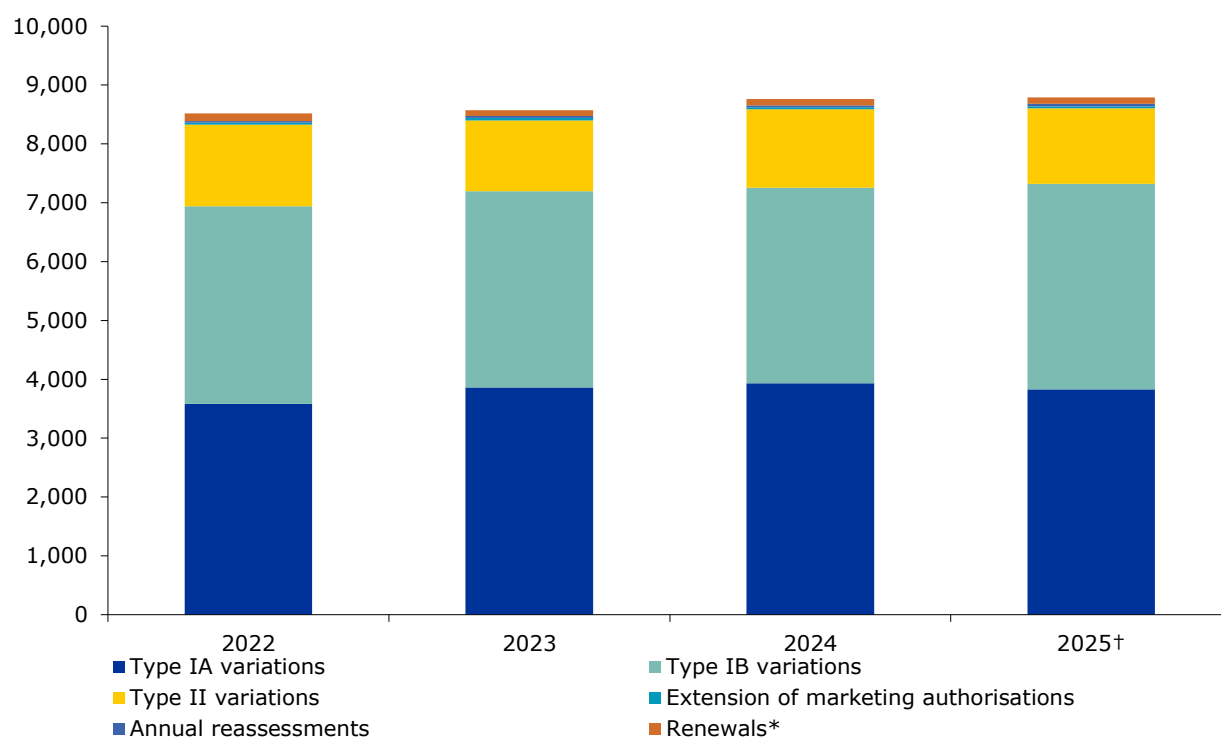
Table 4. Post-authorisation: Variations (scopes), renewals and annual reassessments

	2022		2023		2024		2025 [†]	
	Started	Finalised	Started	Finalised	Started	Finalised	Started	Finalised
Type IA variations	3,586	3,456	3,864	3,883	3,931	3,845	3,831	3,622
Type IB variations	3,354	3,169	3,332	3,303	3,323	3,105	3,490	3,464
Type II variations	1,388	1,373	1,201	1,131	1,333	1,261	1,284	1,175
Extensions of marketing authorisation	31	23	43	32	33	36	35	36
Annual reassessments	27	28	33	29	35	38	39	37
Renewals*	132	129	101	116	107	101	111	117

* Includes renewals of conditional marketing authorisations.

[†] Figures for the current year are cumulative, year to date. Figures for preceding years are totals for the year.

Post-authorisation: Variations, renewals and annual reassessments



[†] Figures for the current year are cumulative, year to date. Figures for preceding years are totals for the year.