

5th August 2010 EMA/502619/2010

### Medicinal products for human use

Monthly figures — July 2010

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.

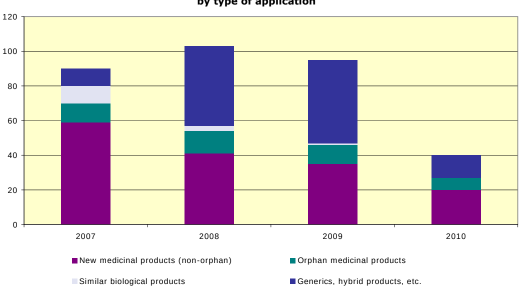


Monthly figures — July 2010

	2007		2008		2009		2010	
	Started	Finalised	Started	Finalised	Started	Finalised	Started	Finalised
Non-orphan medicinal products								
New products	59	40	41	40	35	49	20	11
Advanced therapy medicinal products	na	na	na	na	0	1	0	0
Advanced therapy Art. 29 transition products	na	na	na	na	0	0	0	0
Paediatric-use (PUMA) products	0	0	0	0	0	0	0	0
Well-established use, abridged, hybrid & non- prescription switch products	4	4	16	11	10	14	4	3
Generic products	6	5	30	4	38	51	9	12
Similar biological products	10	5	3	6	1	0	0	1
Total products	79	54	90	61	84	114	33	27
Orphan medicinal products								
New products	11	11	13	11	11	11	7	3
Advanced therapy medicinal products	na	na	na	na	0	0	1	0
Total product applications	90	65	103	72	95	125	40	30

<sup>\*</sup> Finalised applications exclude applications withdrawn prior to opinion

# Marketing authorisation application evaluations started by type of application

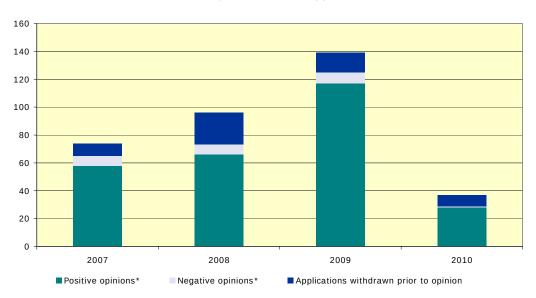


Monthly figures — July 2010

Pre-authorisation: Outcome of the evaluation of marketing authorisation applications								
	2007	2008	2009	2010				
Positive opinions*	58	66	117	28				
Opinions recommending conditional marketing**	5	2	1	4				
Negative opinions*	7	7	8	1				
Applications withdrawn prior to opinion	9	23	14	8				
Re-examinations requested	5	9	7	2				
Opinions after accelerated assessment	5	2	0	1				

<sup>\*</sup> Only the final outcome in case of the re-examination of an opinion under Art. 9(2) of Regulation (EC) No. 726/2004 is reported

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



<sup>\*</sup> Only the final outcome in case of the re-examination of an opinion under Art. 9(2) of Regulation (EC) No. 726/2004 is reported

<sup>\*\*</sup> Included in the figures for positive opinions

Monthly figures — July 2010

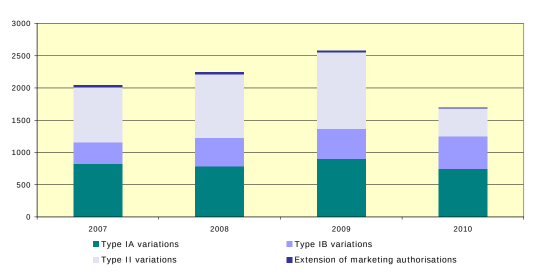
Scientific services									
	2007		2008		2009		2010		
	Received	Finalised	Received	Finalised	Received	Finalised	Received	Finalised	
Compassionate-use opinions	0	0	0	0	1	0	1	2	
Art. 58 (WHO) scientific opinions	1	0	0	0	0	0	1	0	
Opinions on ancillary medicinal substances in medical devices*	3	0	1	0	0	1	2	0	
Plasma master file (includes initial certification, variations and annual re-certification)	16	17	19	23	23	23	6	12	
Vaccine antigen master file	0	0	0	0	0	0	0	0	

<sup>\*</sup> Consultation in accordance with Council Directive 93/42/EEC concerning medical devices as amended by Directive 00/70/EC as regards medical devices incorporating stable derivates of human blood or plasma and Directive 2001/14/EC

Monthly figures — July 2010

	2007		2008		2009		2010	
	Received	Finalised	Received	Finalised	Received	Finalised	Received	Finalised
Type IA variations	822	820	783	783	897	842	748	552
Type IB variations	338	292	445	462	470	412	502	339
Type II variations	853	777	981	877	1186	1142	432	571
Extension of marketing authorisations	32	28	37	35	24	31	12	18
Percentage of variations submitted in grouped notifications/applications <sup>†</sup>	N/A	N/A	N/A	N/A	N/A	N/A	47%	29%
Multi-product Type IA groups	N/A	N/A	N/A	N/A	N/A	N/A	17	9
Worksharing variation applications	N/A	N/A	N/A	N/A	N/A	N/A	50	17
Annual reassessments	24	25	24	24	21	17	8	14
Renewals*	46	44	65	59	46	54	34	18

#### Post-authorisation: Variations received



<sup>\*</sup> Includes renewals of conditional marketing authorisations
† Excluding groups in worksharing or in multi-product Type IA groups