

10 July 2017 EMA/412075/2017 Veterinary Medicines Division

Monthly report on application procedures, guidelines and related documents for veterinary medicines

June 2017

This report, which is updated every month, provides current information related to the volume and evaluation of pre- and post-authorisation applications for medicinal products for veterinary use received by the European Medicines Agency (EMA) for the current and previous three years on:

- scientific advice requests;
- applications for initial evaluations, extensions, variations and renewals concerning marketing authorisations (MAs);
- applications for initial evaluations, extensions, modifications and extrapolations for maximum residue limits (MRLs);
- arbitration and referral procedures;
- requests for classification and re-classifications of products as Minor Use Minor Species (MUMS)/limited market.

In addition, the report includes a summary table of the opinions issued by the Committee for Medicinal Products for Veterinary Use (CVMP) in the current year, as well as a list of adopted guidelines and other public guidance documents.

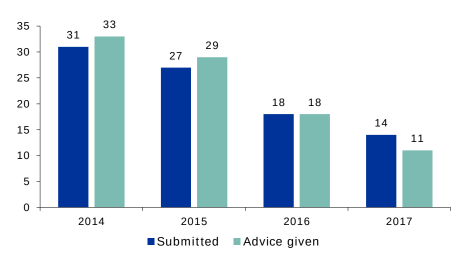
The purpose is only to provide ongoing factual information. Commentaries and analysis are provided in the Agency's annual reports.



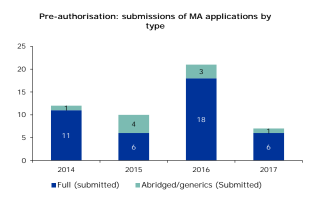
Statistics on pre- and post-authorisation applications for medicinal products for veterinary use

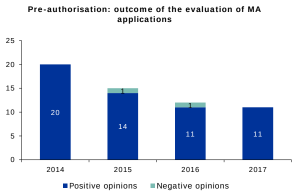
Scientific advice requests					
	2014	2015	2016	2017	
Submitted and validated	31	27	18	14	
Advice given	33	29	18	11	

Scientific advice requests submitted and advice given



Initial evaluation of marketing authorisation applications							
	2014 2015 2016 2017						
Full (submitted)	11	6	18	6			
Abridged/generics (submitted)	1	4	3	1			
Withdrawals	3	0	1	1			
Positive opinions	20	14	11	11			
Negative opinions	0	1	1	0			



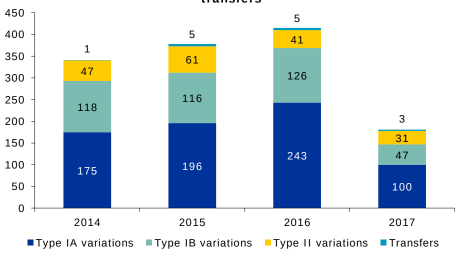


Marketing authorisations					
	2014	2015	2016	2017	
Granted	19	17	7	11	
Withdrawals	1	3	1	0	
Refusal	0	1	0	0	
Not renewed	0	0	1	0	

Extensions — applications					
	2014	2015	2016	2017	
Submitted	6	3	3	4	
Withdrawals	1	0	0	0	
Positive opinions	2	6	5	2	
Negative opinions	0	1	0	0	

Variations — applications submitted					
	2014	2015	2016		
Type-IA variations	175	196	243	100	
Type-IB variations	118	116	126	47	
Type-II variations	47	61	41	31	
Transfers	1	5	5	3	





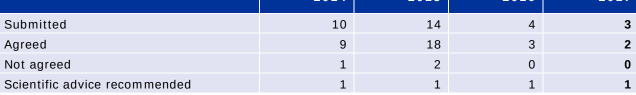
Renewals — applications					
	2014	2015	2016	2017	
Submitted	10	24	13	2	
Positive opinions	15	19	14	5	
Negative opinions	0	0	0	0	

Establishment of MRLs for new substances ¹ — applications						
2014 2015 2016 20						
Submitted	4	4	6	2		
Withdrawals	0	1	0	2		
Positive opinions ^{2,3}	4	3 (1)	2	2		
Negative opinions	0	0	0	0		

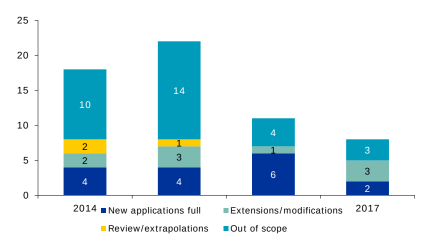
Extensions/modifications of MRLs ⁴ — applications					
	2014	2015	2016	2017	
Submitted	2	3	1	3	
Withdrawals	0	0	1	0	
Positive opinions ²	8	2	3	0	
Negative opinions	0	0	0	0	

Review of opinions/extrapolations of MRLs° – requests from Commission or Member States							
2014 2015 2016 2							
Submitted	2	1	0	0			
Opinion ²	2	3	0	0			

Substances considered as not falling within the scope of Regulation (EC) No 470/2009 requests 2014 2015 2016 2017 Submitted 14 10 4



MRL-related submissions



¹ Establishment of MRLs for new substances under article 3 of Regulation (EC) No 470/2009.

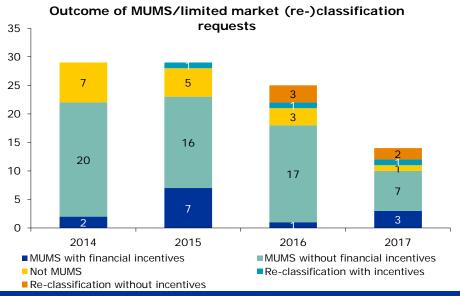
² Including opinions recommending the extension of the expiry date for provisional MRLs or definitive MRLs for substances previously with provisional MRLs.

Re-examinations of opinions are indicated in brackets.

⁴ Extension or modification of MRLs under article 3 of Regulation (EC) No 470/2009.

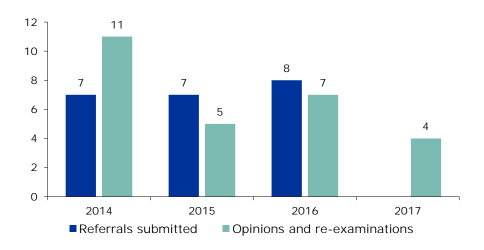
⁵ Review of opinions under article 11 of Regulation (EC) No 470/2009 or requests under article 27 of Regulation (EC) No 470/2009.

MUMS/limited market (re)classification requests — outcome				
	2014	2015	2016	2017
MUMS/limited market with financial incentives	2	6	1	3
MUMS/limited market without financial incentives	20	16	17	7
MUMS/limited market reclassification with financial incentives ⁶	0	1	1	1
MUMS/limited market reclassification without financial incentives ⁶	0	0	3	2
Not MUMS/limited market	7	5	3	1



Arbitrations and referrals				
	2014	2015	2016	2017
Arbitrations and referrals submitted	7	7	8	0
Opinions ⁷	11 (1)	5	7	4(1)

Arbitrations and referrals submitssions and opinions



 $^{^{6}}$ For re-classification the first year available is 2014.

⁷ Re-examinations of opinions are in brackets.

CVMP opinions in 2017 on medicinal products for veterinary use

Positive opinions

Product	Marketing	Target species	Regulatory information
Invented nameINN/Common name	authorisation holder		Procedure numberOpinion date
CredelioLotilaner	Elanco Europe Ltd	• Dog	EMEA/V/C/004247/000016/02/2017
CYTOPOINTLokivetmab	Zoetis Belgium SA	• Dog	EMEA/V/C/003939/000016/02/2017
 Zulvac BTV Ovis Bluetongue vaccine (inactivated) (multistrain: 1-2 strains out of a set of 3) 	Zoetis Belgium SA	• Sheep	EMEA/V/C/004185/000016/02/2017
Ingelvac PCV FLEXPorcine circovirus vaccine (inactivated)	 Boehringer Ingelheim Vetmedica GmbH 	• Pig	EMEA/V/C/004645/000016/03/2017
 RESPIPORC FLUpan H1N1 Swine influenza vaccine (inactivated) 	IDT Biologika GmbH	• Pig	EMEA/V/C/003993/000016/03/2017
 Zeleris Florfenicol/meloxicam	CEVA Santé Animale	• Cattle	EMEA/V/C/004099/000016/03/2017
PrevomaxMaropitant	Le Vet Beheer B.V.	Dogs, Cats	EMEA/V/C/004331/000012/04/2017
ExzoltFluralaner	Intervet International B.V.	• Chickens	EMEA/V/C/004344/000015/06/2017
 Innovax-ND-IBD Newcastle disease, infectious bursal disease and Marek's disease vaccine (live recombinant) 	Intervet International B.V.	• Chickens	EMEA/V/C/004422/000015/06/2017
 Suvaxyn PRRS MLV Porcine respiratory and reproductive syndrome virus vaccine (live) 	Zoetis Belgium SA	 Pigs for fattening, Pigs for reporduction 	EMEA/V/C/004276/000015/06/2017
 VEPURED E. coli verotoxoid vaccine (inactivated recombinant) 	 Laboratorios Hipra, S.A. 	• Pigs	EMEA/V/C/004364/000015/06/2017

CVMP opinions in 2017 on establishment of MRLs

Positive opinions

Product	Target species	Regulatory information
Substance		Procedure numberOpinion date
• Alarelin	All food producing species	EMEA/V/MRL/04706/FULL/000112/04/2017
Bromelain	• Porcine	EMEA/V/MRL/004479/FULL/000111/05/2017

Arbitrations and referrals in 2017

Ongoing procedures

Type of procedure	Date	Product
	Clock start	Product name
	CVMP opinion	• INN
 Referral under Article 34 of Directive 2001/82/EC 	09/09/201512/04/2017	Denagard 45% and associated namesTiamulin hydrogen fumarate
 Referral under Article 35 of Directive 2001/82/EC 	05/11/201511/05/2017	 All veterinary medicinal products containing moxidectin to be administered to cattle, sheep and horses Moxidectin
 Referral under Article 35 of Directive 2001/82/EC (re-examination) 	17/02/201608/12/201616/03/2017	 All veterinary medicinal products containing zinc oxide to be administered orally to food producing species Zinc oxide
 Referral under Article 35 of Directive 2001/82/EC 	18/05/201616/03/2017	 Veterinary medicinal products containing methylprednisolone hydrogen succinate presented as solutions for injection for intramuscular use in cattle Methylprednisolone hydrogen succinate
 Referral under Article 35 of Directive 2001/82/EC 	13/07/201616/03/2017	 Veterinary medicinal products containing tylosin to be administered parenterally and intended for the treatment of bovine mastitis caused by <i>Mycoplasma spp</i> Tylosin
 Referral under Article 34 of Directive 2001/82/EC 	• 13/07/2016	Girolan and its associated name ApralanApramycin sulfate
 Referral under Article 34 of Directive 2001/82/EC 	• 13/07/2016	Lincocin and associated namesLincomycin
 Referral under Article 35 of Directive 2001/82/EC 	• 07/09/2016	Zanil and associated names, and generic products thereofOxyclozanide

Guidelines and working documents in 2017

CVMP quality

Reference number	Document title	Status
EMA/CHMP/CVMP/QWP/BWP/42 8135/2016	Draft Concept paper on the need for Revision of Note for guidance on quality of water for pharmaceutical use (H+V)	Adopted for consultation January 2017 (End of consultation TBC)
EMA/CHMP/CVMP/QWP/826771/ 2016	Corrigendum to Reflection paper on the Requirements for selection and justification of starting materials for the manufacture of chemical active substances	Adopted January 2017

CVMP safety

Reference number	Document title	Status
EMA/CVMP/SWP/377245/2016	Guideline on assessment and control of DNA reactive (mutagenic) impurities in veterinary medicinal	Adopted for consultation February 2017
	products	(End of consultation 31
		August 2017)

CVMP efficacy

Reference number	Document title	Status
EMA/CVMP/344/1999-Rev.2	Guideline on the conduct of efficacy studies for intramammary products for use in cattle	Adopted January 2017
EMA/CVMP/EWP/573536/2013	Reflection paper on anthelmintic resistance	Adopted April 2017
EMA/CVMP/EWP/016/00-Rev.3	Revised guideline on the conduct of bioequivalence studies for veterinary medicinal products	Adopted for consultation April 2017
		(End of consultation 31 October 2017)

CVMP pharmacovigilance

Reference number	Document title	Status
EMA/CVMP/PhVWP/171122/2016	Revised recommendation for the	Adopted for consultation
	basic surveillance of	February 2017
	Eudravigilance Veterinary (EVVet)	
	data for centrally authorised	(End of consultation 31
	products (CAPs)	August 2017)

Reference number	Document title	Status
EMA/CVMP/PhVWP/303762/2012 - Rev. 1	Revised Questions and answers on serious non-fatal adverse events and reporting rules	Adopted April 2017
EMA/CVMP/PhVWP/357539/2015	Reflection paper on non- spontaneous adverse event reports (literature, internet and social media) for veterinary medicinal products	Adopted May 2017

CVMP antimicrobials

Reference number	Document title	Status

CVMP immunologicals

Reference number	Document title	Status
EMA/CVMP/IWP/592652/2014	CVMP Risk Management Strategy - Managing the risk of the potential presence of replication competent endogenous retrovirus RD114 in starting materials and final products of feline and canine vaccines	Adopted February 2017
EMA/CVMP/IWP/123243/2006- Rev.3	Guideline on data requirements for immunological veterinary medicinal products intended for minor use or minor species (MUMS)/limited market	Adopted April 2017

CVMP environmental risk assessment

Reference number	Document title	Status
EMA/CVMP/ERA/103555/2015	Guideline on assessing the toxicological risk to human health and groundwater communities from veterinary pharmaceuticals in groundwater	Adopted for consultation February 2017 (End of consultation 31 August 2017)
EMA/CVMP/ERA/689041/2015	Guideline on the plant testing strategy for veterinary medicinal products	Adopted March 2017
EMA/CVMP/448211/2015	Authorisation of veterinary medicinal products containing (potential) persistent, bioaccumulative and toxic (PBT) or very persistent and very bioaccumulative (vPvB) substances	Adopted April 2017

CVMP novel therapies

Reference number	Document title	Status
EMA/CVMP/ADVENT/751229/2016	Questions and Answers on allogenic stem cell-based products for veterinary use: specific questions on sterility	Adopted June 2017

Replacement, Reduction, Refinement of animal testing (3Rs)

Reference number	Document title	Status

General

Reference number	Document title	Status
EMA/CVMP/757903/2016	Question and answer on the information contained within section 5.1 of the SPC on pharmacodynamic properties for pharmaceutical products	Adopted February 2017