

4 July 2011 EMA/CVMP/37837/2011 Veterinary Medicines and Product Data Management

Committee for Medicinal Products for Veterinary Use

Monthly report of application procedures, guidelines and related documents June 2011

The CVMP monthly report includes statistical data for the current and previous two years on scientific advice, initial evaluations, variations, line extensions, renewals, MRLs initial evaluations and MRLs extensions/modifications and arbitration and referral procedures.

In addition, the report includes a summary table of the opinions issued by the CVMP in the current year and a list of adopted guidelines and other public documents.

Applications for medicinal products for veterinary use and maximum residue limits (MRLs)

Scientific advice requests							
	95-08 2009 2010 2011 Total						
Submitted	69	11	21	13	114		
Advice given 65 8 18 10 101							

Initial evaluation					
	95-08	2009	2010	2011	Total
Full	110	14	16	2	142
(Submitted)					
Abridged/	10	1	2	1	14
generics					
(Submitted)					
Withdrawals	12	0	1	0	13
Positive	91	13	14	12	130
opinions					
Negative	1	0	0	0	1
opinions					

Marketing authorisations						
	95-08 2009 2010 2011 Total					
Granted	88	12	9	14	123	
Withdrawals	2	0	4	0	6	
Not renewed	2	0	0	0	2	

Extensions					
	95-08	2009	2010	2011	Total
Submitted	60	12	3	2	77
Withdrawals	2	1	1	0	4
Positive	40	7	8	2	57
opinions					
Negative	0	0	0	0	0
opinions					



Variations – applications submitted							
	95-08	95-08 2009 2010 2011 Total					
Type IA	339	32	76	47	631		
Type IB	339	41	63	33	031		
Type II	210	40	26	9	285		
Transfers	11	3	8	3	25		

Renewals						
	95-08	2009	2010	2011	Total	
Submitted	50	18	7	9	84	
Positive	48	17	8	5	78	
opinions						
Negative	0	0	0	0	0	
opinions						

Arbitrations and Community referrals						
	95-08 2009 2010 2011 Total					
Referrals submitted	38	9	12	9	68	
Opinions reached ¹	20	15 (5)	11 (1)	7	53 (6)	

¹ Re-examination of opinions in brackets

Substances considered as not falling within the				
scope of Regulation (EC) No 470/2009				
2011 Total				
Submitted	2	2		
Agreed	6	6		
Scientific advice recommended 0 0				

MUMS/ Limited market classification					
	2011	Total			
Positive with financial incentives	4	4			
Positive without financial	8	8			
incentives					
Negative	1	1			

Establishment of MRLs for new substances						
	95-08	2009	2010	2011	Total	
Submitted	66	4	3	1	74	
Withdrawals	5	0	0	0	5	
Positive	54	2	2	3	61	
opinions ²						
Negative	7	0	0	0	7	
opinions ³						

Extensions / modifications/extrapolations of MRLs							
	95-08	2009	2010	2011	Total		
Submitted	98	2	10	3	113		
Withdrawals	4	0	0	0	4		
Positive	113	3	3	4	122		
opinions ²							
Negative	6	0	0	0	6		
opinions							
Extrapolations	50	0	0	0	50		

² Including opinions recommending the extension of the expiry date for provisional MRLS or definitive MRLs for substances with previously provisional maximum residue limits
³ Including one opinion concluding that final MRL could not be established for a substance with

provisional maximum residue limits previously established

CVMP opinions in 2011 on medicinal products for veterinary use

Positive opinions

Pro	oduct	Marketing	Therapeutic area	EMA/CVMP	European Commission
		authorisation	Target species	Validation	Opinion received
•	Invented name INN	holder	Summary of	• Opinion	Date of decision
	TIVIV		indication	 Active time 	 Notification
				 Clock stop 	Official Journal
•	CaniLeish	Virbac S.A.	• Dogs	• 17/03/2010	• 13/01/2011
			Vaccine against	• 12/01/2011	• 14/03/2011
			Leishmania	• 210	
			infection	• 91	
•	ZULVAC 1 + 8	Pfizer Limited	• Sheep	• 18/03/2010	• 13/01/2011
	Ovis		Vaccine for	• 12/01/2011	• 14/03/2011
			prevention of viraemia caused	180119	
			by Bluetongue	• 119	
			Virus serotypes 1		
			and 8		
•	BLUEVAC BTV8	CZ Veterinaria	Cattle, sheep	• 17/01/2009	• 10/02/2011
		S.A	Vaccine for	• 09/02/2011	• 14/04/2011
			active	• 210	
			immunisation	• 543	
			against		
			bluetongue		
			disease		
•	Procox	Bayer Animal	• Dogs	• 16/02/2010	• 11/02/2011
•	Emodepside and	Health GmbH	Treatment of	• 09/02/2011	• 20/04/2011
	toltrazuril		dogs when	• 210	
			mixed parasitic infections,	• 148	
			caused by		
			caused by certain specific		
			roundworms and		
			coccidia are		
			suspected or		
			demonstrated		
•	Veraflox	Bayer Animal	Dogs, cats	• 19/05/2009	• 11/02/2011
•	Pradofloxacin	Health GmbH	Treatment for	• 14/07/2010	• 12/04/2011
			dogs and cats	• 205	
			with particular	• 217	
			infections caused	• 09/02/2011	
			by certain	(re-consideration)	
			specific and	•	
			susceptible		
			pathogens		

Pro	oduct	Marketing	Therapeutic area EMA/CVMP European Commission
•	Invented name INN	authorisation holder	 Target species Summary of indication Opinion Opinion Date of decision Notification Clock stop Opinion received Date of decision Notification
•	Zuprevo Tildipirosin	Intervet International BV	 Pigs, cattle Treatment of bacterial infections in the respiratory tract in pigs and cattle 16/02/2010 08/03/2011 06/05/2011 06/05/2011
•	CERTIFECT Fipronil, (S)- methoprene, amitraz	MERIAL SAS	 Dogs Treatment and prevention of infestations with ticks, alone or in association with fleas and/or chewing lice 16/03/2010 09/03/2011 06/05/2011 06/05/2011
•	MS-H Vaccine Mycoplasma synoviae strain MS-H	Pharmsure Ltd	 Chickens Vaccine to reduce air sac lesions and reduce the number of eggs with abnormal shell formation caused by Mycoplasma synoviae 15/12/2009 08/04/2011 14/06/2011 14/06/2011
•	Recuvyra Fentanyl	Nexcyon Pharmaceuticals Ltd	 Dogs Control of post-operative pain associated with major orthopaedic and soft tissue surgery 16/12/2009 05/05/2011 05/05/2011 210 294
•	Emdocam Meloxicam	Emdoka bvba	 Cattle, pigs, horses For treatment in respiratory infections, diarrhoea and mastitis in cattle. For treatment in non-infectious locomotor disorders and in puerperal 18/05/2010 09/06/2011 175 211 Total ment in 175 211 Total ment in 175 Total ment in 18/05/2010 09/06/2011 09/06/2011 09/06/2011

Pro	oduct	•	Marketing	Th	erapeutic area	EM	1A/CVMP	Eu	ropean Commission
•	Invented name INN		authorisation holder	•	Target species Summary of indication	•	Validation Opinion Active time Clock stop	•	Opinion received Date of decision Notification Official Journal
					septicaemia and toxaemia in pigs. In horses for treatment in musculo-skeletal disorders as well for the relief of pain in equine colic.				
•	Proteq West Nile West Nile recombinant canarypox virus (vCP2017 virus)	•	MERIAL	•	Horses Vaccine for the active immunisation of horses against West Nile disease	•	18/05/2010 09/06/2011 196 190	•	09/06/2011
•	Zulvac 1 Bovis Inactivated Bluetongue virus, serotype 1, strain BTV-1	•	Pfizer Limited	•	Cattle Active immunisation of cattle for the prevention of viraemia caused by Bluetongue Virus, serotype 1	•	12/08/2010 09/06/2011 180 120	•	
•	Zulvac 1 Ovis Inactivated Bluetongue Virus, serotype 1, strain BTV-1	•	Pfizer Limited	•	Sheep Active immunisation of sheep for the prevention of viraemia caused by Bluetongue Virus, serotype 1	•	15/07/2010 08/06/2011 179 148	•	

CVMP opinions in 2011 on establishment of MRLs for new substances

Positive opinions

Substance INN Methylpredni – solone (after provisional MRLs)	Target speciesBovine	EMA/CVMP Validation Opinion Active time Clock stop n/a 12/01/2011 90 0	European Commission Opinion received Date of regulation Official Journal 27/01/2011
Octenidine dihydrochloride	All mammalian food producing species	11/08/200908/02/2011210246	• 21/02/2011
 Monepantel (after provisional MRLs) 	Caprine	n/a09/03/2011900	• 25/03/2011
Azamethiphos	Fin fish	21/02/201107/04/2011450	• 08/04/2011
Pegylated bovine granulocyte colony stimulating factor	Bovine	16/03/201005/05/2011210205	• 18/05/2011
Lasalocid	Bovine	10/08/201005/05/201121058	• 18/05/2011
Ivermectin	All mammalian food producing species	n/a09/06/20111760	• 20/06/2011

Arbitrations and Community referrals in 2011

Type of referral	Date of clock startCVMP opinion	Product nameINN
Referral under Art. 34 of Directive	• 11/11/2009	Fortekor vet and associated names
2001/82/EC		Benazepril hydrochloride
Referral under Art. 34	• 14/04/2010	Synulox Lactating Cow and associated names
of Directive 2001/82/EC	• 07/06/2011	Amoxicillin, clavulanic acid, prednisolone
Referral under Art.	• 14/07/2010	Combimox Lactating Cow
33(4) of Directive 2001/82/EC	• 07/04/2011	Amoxicillin, clavulanic acid, prednisolone
Referral under Art.	• 14/07/2010	Nisamox Lactating Cow
33(4) of Directive 2001/82/EC	• 07/04/2011	Amoxicillin, clavulanic acid, prednisolone
Referral under Art.	• 14/07/2010	Combisyn Lactating Cow
33(4) of Directive 2001/82/EC	• 07/04/2011	Amoxicillin, clavulanic acid, prednisolone
Referral under Art. 34	• 14/07/2010	Doxycycline 50% WSP and associated names
of Directive 2001/82/EC	• 04/05/2011	Doxycycline hyclate
Referral under Art. 34	• 14/07/2010	Doxyfar 50% WSP and associated names
of Directive	• 04/05/2011	Doxycycline hyclate
2001/82/EC		2 Doxyoyamio Hydiato
Referral under Art. 34 of Directive	• 09/11/2010	Baytril 10% oral solution and associated names
2001/82/EC		Enrofloxacin
Referral under Art.	• 09/02/2011	Clavudale 50 mg tablet for cats and dogs
33(4) of Directive	• 08/06/2011	Amoxicillin and clavulanic acid
2001/82/EC Referral under Art. 35	• 09/03/2011	Votorinary modicinal products containing
of Directive 2001/82/EC	09/03/2011	 Veterinary medicinal products containing active substances belonging to the class of flukicides for which no MRL has been established in milk
Referral under Art. 35 of Directive 2001/82/EC	• 06/04/2011	All veterinary medicinal products containing systemically administered (parenteral and oral) 3rd and 4th generation cephalosporins and intended for use in food producing species
		Cefquinome and ceftiofur
Referral under Art. 33(4) of Directive	• 04/05/2011	 Prontax 10 mg/ml solution for injection for sheep, cattle and pigs
2001/82/EC		• Doramectin

Type of referral	Date of clock startCVMP opinion	Product name INN
Referral under Art. 33(4) of Directive 2001/82/EC	• 04/05/2011	Prontax 5 mg/ml pour-on solution for cattleDoramectin
Referral under Art. 35 of Directive 2001/82/EC	• 04/05/2011	 All pre-mixes for medicated feedingstuffs containing 40, 100 or 200 g tilmicosin per kg pre-mix Tilmicosin
Referral under Art. 78 of Directive 2001/82/EC	• 04/05/2011	 HIPRABOVIS PNEUMOS Emulsion for injection for cattle and associated names Inactivated Mannheimia haemolytica and Histophilus somni

Guidelines and working documents in 2011

CVMP Efficacy

Reference number	Document title	Status
EMA/CVMP/016/00-Rev.2	Guideline on the conduct of bioequivalence studies for veterinary medicinal products	Adopted April 2011
EMA/CVMP/760764/2010	Concept paper on the revision of the CVMP Guideline for the demonstration of efficacy for veterinary medicinal products containing antimicrobial substances	Adopted for consultation, April 2011 (End of consultation 31 July 2011)
EMA/CVMP/EWP/459868/2008	Guideline on demonstration of target animal safety and efficacy of veterinary medicinal products intended for use in farmed finfish	Adopted May 2011

CVMP Environmental Risk Assessment

Reference number	Document title	Status
EMA/CVMP/ERA/147844/2011	Reflection paper on the testing strategy and risk assessment for plants	Adopted for consultation, March 2011 (End of consultation 30 June 2011)
EMA/CVMP/ERA/430327/2009	Guideline on determining the fate of veterinary medicinal products in manure	Adopted March 2011
EMA/CVMP/ERAWP/409328/2010	Reflection paper on risk mitigation measures related to the environmental risk assessment of veterinary medicinal products	Adopted for consultation, May 2011 (End of consultation 31 August 2011)

CVMP Immunologicals

Reference number	Document title	Status
EMA/CVMP/IWP/206555/2010	Guideline on requirements for the production and control of immunological veterinary medicinal products	Adopted for consultation, March 2011 (End of consultation 30 September 2011)
EMA/CVMP/IWP/314550/2010	Guideline on the design of studies to evaluate the safety and efficacy of fish vaccines	Adopted for consultation, March 2011 (End of consultation 30 September 2011)

CVMP Pharmacovigilance

Reference number	Document title	Status
EMA/CVMP/PhVWP/471721/2006	Recommendation on the basic surveillance of EudraVigilance Veterinary (EVVet) data	Adopted February 2011
EMA/CVMP/PhVWP/44873/2011	Public bulletin - Veterinary pharmacovigilance for 2010	Adopted February 2011
EMA/CVMP/10418/2009-Rev.3	CVMP combined VeDDRA list of clinical terms for reporting suspected adverse reactions in animals and humans to veterinary medicinal products	Adopted June 2011
EMA/CVMP/PhVWP/377827/2011	List of species and breeds for electronic reporting of suspected adverse reactions in veterinary pharmacovigilance	Adopted June 2011
EMA/CVMP/PhVWP/288284/2007- Rev.4	Quidance notes on the use of VeDDRA terminology for reporting suspected adverse reactions in animals and humans	Adopted June 2011
SOP/V/4019	Standard operating procedure - Annual review of standard lists to be used in EudraVigilance Veterinary	Adopted June 2011

CVMP Scientific Advisory Group on Antimicrobials

Reference number	Document title	Status
EMA/CVMP/SAGAM/736964/2009	Reflection paper on meticillin- resistant <i>Staphylococcus</i> pseudintermedius (MRSP)	Adopted January 2011

General

Reference number	Document title	Status
EMA/347137/2010	Summary of procedures for consultation by CVMP of Scientific Advisory Groups (SAGs) and ad-hoc expert groups functioning as SAGs in relation to applications for authorisation for veterinary medicinal products	Adopted February 2011