

17 October 2013 EMA/587321/2013 Committee for Medicinal Products for Veterinary Use (CVMP)

CVMP Monthly report of application procedures, guidelines and related documents

September 2013

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											
95-10 2011 2012 2013 Total											
Submitted	Submitted 101 26 28 28 183										
Advice given 91 24 29 25 169											

Initial evaluation										
	95-10	2011	2012	2013	Total					
Full	140	8	12	18	178					
(Submitted)										
Abridged/	13	3	0	0	16					
generics										
(Submitted)										
Withdrawals	13	0	1	0	14					
Positive	118	19	9	8	154					
opinions										
Negative	1	0	0	0	1					
opinions										

Marketing authorisations											
95-10 2011 2012 2013 Total											
Granted	111	24	8	11	154						
Withdrawals 6 1 3 3 13											
Not renewed	Not renewed 2 0 0 0 2										

Extensions					
	95-10	2011	2012	2013	Total
Submitted	75	7	8	4	94
Withdrawals	4	0	1	0	5
Positive	55	4	10	7	76
opinions					
Negative	0	0	0	0	0
opinions					



Variations – applications submitted										
95-10 2011 2012 2013 Total										
Type IA	551	120	104	137	1174					
Type IB	331	101	96	65	1174					
Type II	276	45	52	22	395					
Transfers	<u> </u>									

Renewals										
	95-10	2011	2012	2013	Total					
Submitted	75	14	10	12	111					
Positive opinions	73	12	10	11	106					
Negative opinions	0	0	0	0	0					

Arbitrations and Community referrals										
95-10 2011 2012 2013 Total										
Referrals	59	12	12	9	92					
submitted										
Opinions	46	10	11	11	78					
reached.1	(6)		(1)	(3)	(10)					

¹ Re-examination of opinions in brackets

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the scope of Regulation (EC) No 470/2009										
2010 2011 2012 2013 Total										
Submitted	5	5	9	12	31					
Agreed	0	10	6	7	23					
Scientific	0	0	0	3	3					
advice	advice									
recommend										

Substances considered as not falling within

r									
MUMS/ Limited market classification									
2011 2012 2013 Total									
Positive with	8	16	10	34					
financial incentives									
Positive without	12	5	9	26					
financial incentives									
Negative	1	1	2	4					

Establishment of MRLs for new substances										
95-10 2011 2012 2013 Total										
Submitted	73	1	1	3	78					
Withdrawals	5	0	0	2	7					
Positive	58	4	1	2	65					
opinions. ² .										
Negative	7	0	0	0	7					
opinions.3										

Extensions / modifications/extrapolations of MRLs									
	95-10	2011	2012	2013	Total				
Submitted	110	13	5	5	133				
Withdrawals	4	2	0	0	6				
Positive	119	12	8 (2)	2	141				
opinions.2									
Negative	6	0	0	0	6				
opinions									

² Including opinions recommending the extension of the expiry date for provisional MRLs or definitive MRLs for substances with previously provisional maximum residue limits. Re-examination of opinions are indicated in brackets.

are indicated in brackets.

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 2013 on medicinal products for veterinary use

Positive opinions

Pre	oduct	•	Marketing	Th	erapeutic area	EN	IA/CVMP	Eu	ropean
	Invented		authorisation	•	Target species	•	Validation	Co	mmission
	name		holder	•	Summary of	•	Opinion	•	Opinion
•	INN				indication	•	Active time		received
	TIVIV					•	Clock stop	•	Decision
								•	Notification
								•	Official Journal
•	Meloxidolor	•	Le Vet Beheer	•	Dogs, cats, cattle,	•	15/12/2012	•	07/02/2013
•	Meloxicam		B.V.		pigs and horses	•	07/02/2013	•	22/04/2013
				•	Anti-inflammatory	•	210	•	24/04/2013
					and anti-rheumatic	•	212	•	C 156 of
									31/05/2013
•	ECOPORC	•	IDT Biologika	•	Piglets	•	15/12/2012	•	08/02/2013
	Shiga		GmbH	•	Vaccine for the	•	07/02/2013	•	10/04/2013
					active immunisation	•	210	•	12/04/2013
					to reduce the	•	212	•	C 156 of
					mortality and clinical				31/05/2013
					sings of oedema				
					disease				
•	Oncept IL-2	•	MERIAL	•	Cats	•	09/11/2012	•	07/03/2013
				•	Immunotherapy	•	07/03/2013	•	03/05/2013
					product to be used in	•	205	•	07/05/2013
					addition to surgery	•	280	•	C 184 of
					and radiotherapy				28/06/2013
					with fibrosarcoma				
					without metastasis				
					or lymph node				
					involvement				
•	Equilis West	•	Intervet	•	Horses	•	17/01/2012	•	11/04/2013
	Nile		International BV	•	For the active	•	11/04/2013	•	06/06/2013
					immunisation of	•	208	•	16/07/2013
					horses against West	•	240	•	C 250 of
					Nile virus (WNV) to				30/08/2013
					prevent virus				
					viraemia and to				
					reduce clinical				
					symptoms of disease				
					and lesions in the				
					brain				

Pro	oduct	•	Marketing	Th	erapeutic area	ΕN	IA/CVMP	Eu	ropean
	Invented		authorisation	•	Target species	•	Validation		mmission
	name		holder	•	Summary of	•	Opinion	•	Opinion
•	INN				indication	•	Active time		received
						•	Clock stop	•	Decision
								•	Notification
	Due 7ime		Doobringer		Coto		15/03/2012	•	Official Journal
•	ProZinc Insulin	•	Boehringer Ingelheim	•	Cats For the treatment of	•	16/05/2013	•	16/05/2013 12/07/2013
•	(human)		Vetmedica	•	diabetes mellitus to		210	•	16/07/2013
	(Harriari)		GmbH		achieve reduction of	•	218	•	C 250 of
					hyperglycaemia and		2.10		30/08/2013
					improvement of				
					associated clinical				
					signs	L			
•	AFTOVAXPUR	•	MERIAL	•	Cattle, sheep, pigs	•	12/10/2012	•	16/05/2013
	DOE			•	Vaccine containing a	•	16/05/2013	•	15/07/2013
					maximum of three	•	210	•	17/07/2013
					inactivated, purified	•	737	•	C 250 of
					foot-and-mouth-				30/08/2013
					disease (FMD) virus				
					strains out of seven				
	APOQUEL	•	Zoetis Belgium	•	authorised strains Dogs	•	15/08/2012	•	18/07/2013
	Oclacitinib		SA	•	Treatment of clinical		18/07/2013		10/07/2013
	maleate		<i>57</i> .		manifestations of	•	210		
	mareate				pruritus associated	•	127		
					with allergic				
					dermatitis in dogs				
					and treatment of				
					clinical				
					manifestations of				
					atopic dermatitis in				
					dogs.				4.0.40
•	Trifexis	•	Eli Lilly & Co Ltd	•	Dogs	•	15/02/2012	•	18/07/2013
•	Spinosad /			•	Treatment and	•	17/07/2013		
	milbemycin oxime				prevention of flea	•	210		
	UXIIIIE				infestations in dogs when the concurrent	•	308		
					prevention of				
					heartworm disease				
					and/or treatment of				
					specified				
					gastrointestinal				
					nematode infections				
					is indicated.				

CVMP opinions in 2013 on establishment of MRLs

Positive opinions

Substance	Target species	EMA/CVMP	European Commission
		 Validation 	Opinion received
		 Opinion 	 Regulation
		 Active time 	Official Journal
		 Clock stop 	omeiai sournai
• Diclazuril	 Rabbits 	• 12/09/2012	• 18/02/2013
		• 07/02/2013	
		• 148	
		• 0	
 Butafosfan 	All mammalian food	• 16/01/2013	• 26/06/2013
	producing species	• 13/06/2013	
		• 148	
		• 0	
 Chloroform 	All mammalian food	• 11/10/2013	• 26/06/2013
	producing species	• 13/06/2013	
		• 175	
		• 71	
Triptorelin aceta	Porcine species	• 13/02/2013	• 18/07/2013
	·	• 18/07/2013	
		• 155	
		• 0	

Arbitrations and Community referrals in 2013

Type of referral	Date of clock start	Product name
	CVMP opinion	• INN
Referral under Article 35 of Directive 2001/82/EC	 15/09/2011 11/04/2013 18/07/2013 (re-examination) 	 All long acting formulations for injection containing barium selenate for all food producing species Barium selenate
Referral under Article 33(4) of Directive 2001/82/EC	 12/10/2011 13/06/2012 07/02/2013 (re-examination) 	Nuflor Swine Once 450 mg/mlFlorfenicol
Referral under Article 35 of Directive 2001/82/EC	12/04/201212/06/2013	 All injectable and pour-on veterinary medicinal products containing doramectin that are intended for use in mammalian food-producing species Doramectin
Referral under Art. 34 of Directive 2001/82/EC	15/05/201218/07/2013	Micotil 300 Injectie and associated namesTilmicosin

Type of referral	Date of clock start	Product name
	CVMP opinion	• INN
Referral under Article 33(4) of Directive	• 15/05/2012 • 07/03/2013	 Florgane 300 mg/ml suspension for injection for cattle and pigs
2001/82/EC		• Florfenicol
Referral under Article 33(4) of Directive	11/07/201210/04/2013	Strenzen 500/125 mg/g powder for use in drinking water for pigs
2001/82/EC		Amoxicillin/clavulanic acid
Referral under Article 35 of Directive 2001/82/EC	• 12/09/2012	Suanovil 20 and associated names, Captalin and associated names and generic products thereof, including pending applications
		• Spiramycin
Referral under Article 35 of Directive 2001/82/EC	12/09/201218/07/2013	Dexadreson 2 mg/ml and associated names, and generic products thereof, including pending applications
		Dexamethasone
Referral under Article	• 10/10/2012	Linco-Spectin 100 and its associated names
34 of Directive 2001/82/EC		Lincomycin, spectinomycin
Referral under Article 34 of Directive	• 07/11/2012	Baytril 2.5% injectable, Baytril 5% injectable and Baytril 10% injectable and their associated names
2001/82/EC		Enrofloxacin
Referral under Article 35 of Directive 2001/82/EC	• 07/11/2012	All veterinary medicinal products containing enrofloxacin to be administered via the drinking water to chickens and/or turkeys
		Enrofloxacin
Referral under Article 13 of Regulation (EC)	07/11/201207/03/2013	Soludox 500 mg/g powder for use in drinking water for pigs and chickens
No. 1234/2008	• 12/06/2013 (re- examination)	Doxycycline hyclate
Referral under Article	• 10/01/2013	Lidocaine
30(3) of Regulation 726/2004		• Lidocaine
Referral under Article 33(4) of Directive 2001/82/EC	• 07/03/2013 • 17/07/2013	Deltanil 10 mg/ml Pour-on Solution for cattle and sheep and Deltanil 100 mg Spot-on Solution for cattle
		Deltamethrin
Referral under Article	• 07/03/2013	Suifertil 4 mg/ml Oral Solution for Pigs
33(4) of Directive 2001/82/EC	• 18/07/2013	Altrenogest

Type of referral	Date of clock start CVMP opinion	Product name INN
Referral under Article 35 of Directive 2001/82/EC	• 10/04/2013	 All veterinary medicinal products containing altrenogest to be administered orally to pigs and horses Altrenogest
Referral under Article 13 of Regulation (EC) No. 1234/2008	10/04/201316/07/2013	 Cydectin TriclaMox pour-on solution for use in cattle Triclabendazole and moxidectin
Referral under Article 33(4) of Directive 2001/82/EC	• 16/05/2013	 Norbonex 5-mg/ml pour-on solution for beef and dairy cattle Eprinomectin
Referral under Article 33(4) of Directive 2001/82/EC	• 16/05/2013	 Fiprex CAT 52.5 mg spot-on solution for cats, Fiprex S 75 mg spot-on solution for dogs, Fiprex M 150 mg spot-on solution for dogs, Fiprex L 300 mg spot-on solution for dogs and Fiprex XL 412.5 mg spot-on solution for dogs
		Fipronil
Referral under Article 13 of Directive 2001/82/EC	• 16/05/2013	 Baytril 2.5% injectable, Baytril 5% injectable, Baytril 10% injectable and associated names and related veterinary medicinal products Enrofloxacin
Referral under Article 45 of Regulation (EC) No. 726/2004	• 16/05/2013	Suvaxyn PCV (inactivated vaccine)

Guidelines and working documents in 2013

CVMP Quality

Reference number	Document title	Status
EMEA/CVMP/511/03-Rev.1	Annexes to: CPMP/ICH/283/95 Impurities: Guideline for residual	Adopted February 2013
	solvents & CVMP/VICH/509/99	
	Guideline on impurities: residual	
	solvents.	
EMA/CVMP/VICH/858875/2011	VIVH GL 51: Quality: Statistical	Adopted March 2013
	evaluation of stability data	
N/a	Q&A on co-operation between	Adopted May 2013
	assessors and inspectors when real-	
	time release testing is applied.	
N/a	Q&A on setting specifications for	Adopted June 2013
	impurities in veterinary medicinal	
	products	

CVMP Safety

Reference number	Document title	Status
EMA/CVMP/SWP/398880/2012	Concept paper on genotoxic impurities	Adopted for consultation, January 2013 (End of consultation 30 April 2013)
EMA/CVMP/VICH/526/2000	VICH GL 23(R) Safety: Studies to evaluate the safety of residues of veterinary drugs in human food: Genotoxicity testing	Adopted for consultation, January 2013 (End of consultation 31 March 2013)

CVMP Environmental Risk Assessment

EMA/CVMP/ERA/718229/2012	Concept paper on assessing the	Adopted for consultation,
	toxicological risk to humans and the	April 2013
	environment of veterinary	
	pharmaceuticals in groundwater	(End of consultation 30 June
		2013)

CVMP Efficacy

Reference number	Document title	Status
EMA/CVMP/EWP/261180/2012	Draft Guideline for the	Adopted for consultation,
	demonstration of efficacy for	May 2013
	veterinary medicinal products	
	containing antimicrobial substances	(End of consultation 30
		November 2013)

CVMP Immunologicals

Reference number	Document title	Status
EMA/CVMP/VICH/463/2002	VICH GL34: Biologicals: Testing for	Adopted March 2013
	the detection of Mycoplasma	
	contamination	
EMA/CVMP/VICH/582610/2009	VICH GL 50: Biologicals:	Adopted March 2013
	Harmonisation of criteria to waive	
	Target Animal Batch Safety Testing	
	(TABST) for inactivated vaccines for	
	veterinary use	
EMA/CVMP/IWP/97961/2013	Draft guideline on the compliance of	Adopted for consultation,
	authorised equine influenza	April 2013
	vaccines with OIE requirements	
		(End of consultation 31
		October 2013)
EMA/CVMP/IWP/594618/2010	Guideline on the requirements for	Adopted July 2013
	combined vaccines and associations	
	of immunological veterinary	
	medicinal products (IVMPs)	

CVMP Pharmacovigilance

Reference number	Document title	Status
EMA/CVMP/PhVWP/536313/2011	Draft Reflection paper on pharmacovigilance communication concerning veterinary medicinal products	Adopted for consultation, February 2013 (End of consultation 31 May 2013)
EMA/CVMP/PhVWP/552/2003- Rev.1	Draft revised Recommendation on harmonising the approach to causality assessment for adverse events to veterinary medicinal products	Adopted for consultation, February 2013 (End of consultation 31 May 2013)
EMA/CVMP/VICH/123940/2006	VICH GL 35 on Pharmacovigilance: Electronic standards for transfer of data	Adopted March 2013
EMA/CVMP/PhVWP/126661/2009- Rev.3	Q&A on Serious non-fatal adverse events and reporting rules	Adopted April 2013
EMA/CVMP/PhVWP/303762/2012	Q&A on PSUR preparation, management and assessment	Adopted April 2013
EMA/CVMP/PhVWP/145186/2013	Q&A on Adverse event reporting	Adopted April 2013
EMA/CVMP/10418/2009-Rev.5	CVMP combined VeDDRA list of clinical terms for reporting suspected adverse reactions in animals and humans to veterinary medicinal products	Adopted June 2013

Reference number	Document title	Status
EMA/CVMP/PhVWP/288284/2007-	Guidance notes on the use of	Adopted June 2013
Rev.6	VeDDRA terminology for reporting	
	suspected adverse reactions in	
	animals and humans	
EMA/123352/2001-Rev.7	Call for comments on standard lists	Adopted June 2013
	for EudraVigilance Veterinary	

CVMP Antimicrobials

Reference number	Document title	Status
EMA/CVMP/680258/2012	Concept paper on the development	Adopted for consultation,
	of a guideline on antimicrobial risk	January 2013
	assessment	
		(End of consultation 30 April
		2013)
EMA/363834/2013	Request for scientific advice on the	Adopted July 2013
	impact on public health and animal	
	health of the use of antibiotics in	
	animals: Answer to the first request	
	from the European Commission	
EMA/755938/2012	Use of colistin products in animals	Adopted July 2013
	within the European Union:	
	development of resistance and	
	possible impact on human and	
	animal health	
EMA/291760/2013	Use of glycylcyclines in animals in	Adopted July 2013
	the European Union: development	
	of resistance and possible impact on	
	human and animal health	

Joint CVMP/ CHMP AHEG on the application of the 3Rs (replacement, refinement and reduction) in regulatory testing of medicinal products $\frac{1}{2}$

Reference number	Document title	Status
EMA/CHMP/CVMP/JEG- 3Rs/746429/2013	Recommendation to marketing authorisation holders for veterinary vaccines, highlighting the need to update marketing authorisations to remove the target animal batch safety test (TABST) following removal of the requirement from the European Pharmacopoeia monographs	Adopted May 2013