

1 October 2012 EMA/615951/2012 Committee for Medicinal Products for Veterinary Use (CVMP)

## CVMP Monthly report of application procedures, guidelines and related documents

September 2012

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								
	95-09	2010	2011	2012	Total			
Submitted	80	21	26	21	148			
Advice given	73	18	24	20	135			

Initial evaluation									
	95-09	2010	2011	2012	Total				
Full (Submitted)	124	16	8	5	153				
Abridged/ generics (Submitted)	11	2	3	0	16				
Withdrawals	12	1	0	1	14				
Positive opinions	104	14	19	5	142				
Negative opinions	1	0	0	0	1				

Marketing authorisations								
	95-09	2010	2011	2012	Total			
Granted	100	9	22	7	138			
Withdrawals	2	4	1	0	7			
Not renewed	2	0	0	0	2			

Extensions										
	95-09	2010	2011	2012	Total					
Submitted	72	3	7	6	88					
Withdrawals	3	1	0	0	4					
Positive	47	8	4	9	68					
opinions										
Negative	0	0	0	0	0					
opinions										

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Variations – applications submitted									
	95-09	2010	2011	2012	Total				
Type IA	412	76	125	77	905				
Type IB	412	63	87	65	703				
Type II	250	26	45	36	357				
Transfers	14	8	3	2	27				

Renewals									
	95-09	2010	2011	2012	Total				
Submitted	68	7	14	8	97				
Positive	65	8	12	7	92				
opinions									
Negative	0	0	0	0	0				
opinions									

Arbitrations and Community referrals									
	95-09	2010	2011	2012	Total				
Referrals	47	12	12	2	79				
submitted									
Opinions	35	11	10	9	65				
reached <sup>1</sup>	(5)	(1)		(1)	(7)				

<sup>1</sup> Re-examination of opinions in brackets

## Substances considered as not falling within the scope of Regulation (EC) No 470/2009

	2011	2012	Total
Submitted	7	2	7
Agreed	9	3	12
Scientific advice	0	0	0
recommended			

MUMS/ Limited market classification							
2011 2012 Tota							
Positive with financial	8	11	19				
incentives							
Positive without financial	12	2	14				
incentives							
Negative	1	1	2				

# Establishment of MRLs for new substances95-09201020112012TotalSubmitted7031074

Submitted	70	3	1	0	74
Withdrawals	5	0	0	0	5
Positive	56	2	4	1	63
opinions <sup>2</sup>					
Negative	7	0	0	0	7
opinions <sup>3</sup>					

Extensions / modifications/extrapolations of MRLs

	95-09	2010	2011	2012	Total
Submitted	100	10	13	3	126
Withdrawals	4	0	2	0	6
Positive	116	3	12	8 (2)	139
opinions <sup>2</sup>					
Negative	6	0	0	0	6
opinions					

 <sup>2</sup> 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.
 <sup>3</sup> Including one opinion concluding that final MRL

<sup>3</sup> Including one opinion concluding that final MRL could not be established for a substance with provisional maximum residue limits previously established

#### CVMP opinions in 2012 on medicinal products for veterinary use

Positive opinions

Pr	oduct	•	Marketing	Th	erapeutic area	ΕN		Eu	Iropean
			authorisation	•	Target species	•	Validation		ommission
•	Invented name INN		holder	•	Summary of indication	•	Opinion Active time Clock stop	•	Opinion received Date of decision Notification Official Journal
•	Zulvac 1+8 Bovis Inactivated Bluetongue virus, serotype 1 and 8, strain BTV-1	•	Pfizer Limited	•	Cattle Vaccine for the active immunisation of cattle for the prevention of viraemia caused by Bluetongue Virus, serotype 1 and 8.	•	04/02/2011 12/01/2012 152 191	•	12/01/2012 08/03/2012 12/03/2012 27/04/2012
•	Poulvac E. Coli	•	Pfizer Limited	•	Chickens Vaccine for the active immunisation to reduce mortality and lesions associated with E. Coli serotype 078	•	09/02/2011 11/04/2012 210 219	•	13/04/2012 15/06/2012 20/06/2012 27/07/2012
•	Porcilis ColiClos	•	Intervet Internatinal B.V.	•	Piglets Vaccine for the passive immunisation against E. Coli and C. perfringens	•	12/10/2010 11/04/2012 210 339	•	16/04/2012 14/06/2012 17/06/2012 27/07/2012
•	Cardalis tablets Benazepril and spironolactone	•	Ceva Santé Animale	•	Dogs Indicated for the treatment of congestive heart failure caused by chronic degenerative valvular disease	•	13/07/2011 16/05/2012 208 99	•	16/05/2012 23/07/2012 25/07/2012 31/08/2012
•	Nobivac L4	•	Intervet Internatinal B.V.	•	Dogs Vaccine containing inactivated Leptospira strains and indicated for the active immunisation of dogs to reduce infection and/or urinary excretion caused by Leptospira strains.	•	04/01/2012 16/05/2012 201 256	•	16/05/2012 16/07/2012 18/07/2012 31/08/2012

#### CVMP opinions in 2012 on establishment of MRLs

Positive opinions

Substance	Target species	EMA/CVMP	European
• INN		<ul><li>Validation</li><li>Opinion</li><li>Active time</li><li>Clock stop</li></ul>	Commission <ul> <li>Opinion received</li> <li>Date of regulation</li> </ul>
			Official Journal
Sodium salicylate (After provisional MRLs)	• Turkeys	<ul> <li>n/a</li> <li>09/02/2012</li> <li>90</li> <li>0</li> </ul>	• 15/02/2012
Prednisolone	• Horses	<ul> <li>12/10/2011</li> <li>08/03/2012; 14/06/2012 (Re-examination)</li> <li>148</li> <li>0</li> </ul>	• 20/06/2012
Monensin	Bovine species	<ul> <li>15/06/2011</li> <li>08/03/2012</li> <li>205</li> <li>63</li> </ul>	• 21/03/2012
Phoxim	All food producing except fin fish	<ul> <li>04/01/2010</li> <li>08/03/2012</li> <li>210</li> <li>220</li> </ul>	• 21/03/2012
Diclazuril	• Poultry	<ul> <li>09/11/2011</li> <li>13/04/2012</li> <li>156</li> <li>0</li> </ul>	• 20/04/2012
Double stranded ribonucleic acid homologous to viral ribonucleic acid coding for part of the coat protein and part of the intergenic region of Israel Acute Paralysis Virus	• Bees	<ul> <li>09/10/2010</li> <li>13/04/2012</li> <li>210</li> <li>312</li> </ul>	• 20/04/2012
Eprinomectin	Ovine and caprine	<ul> <li>18/05/2010</li> <li>13/04/2012</li> <li>183</li> <li>515</li> </ul>	• 20/04/2012
Monepantel	Ovine and caprine     milk	<ul> <li>13/09/2011</li> <li>16/05/2012</li> <li>210</li> <li>36</li> </ul>	• 25/05/2012

Manganese	All food producing	• 15/02/2012	• 25/07/2012
carbonate	species	<ul><li>12/07/2012</li><li>148</li></ul>	
		• 0	

#### Arbitrations and Community referrals in 2012

Type of referral	<ul><li>Date of clock start</li><li>CVMP opinion</li></ul>	Product name     INN
Referral under Art. 34 of Directive 2001/82/EC	<ul><li>09/11/2010</li><li>13/06/2012</li></ul>	Baytril 10% oral solution and associated names
2001/02/20		Enrofloxacin
Referral under Art. 35 of Directive 2001/82/EC	<ul> <li>09/03/2011</li> <li>08/03/2012</li> <li>13/06/2012 (re-examination)</li> </ul>	Veterinary medicinal products containing active substances belonging to the class of flukicides for which no MRL has been established in milk and which are intended for use in ruminants producing milk for human consumption
Referral under Art.	• 04/05/2011	Prontax 5 mg/ml pour-on solution for cattle
33(4) of Directive 2001/82/EC	• 08/02/2012	Doramectin
Referral under Art. 33(4) of Directive 2001/82/EC	<ul><li>04/05/2011</li><li>08/02/2012</li></ul>	Prontax 10 mg/ml solution for injection for sheep, cattle and pigs
2001/82/EC		Doramectin
Referral under Art. 35 of Directive 2001/82/EC	<ul><li>04/05/2011</li><li>08/03/2012</li></ul>	All pre-mixes for medicated feedingstuffs containing 40, 100 or 200 g tilmicosin per kg pre-mix
		• Tilmicosin
Referral under Art. 34 of Directive 2001/82/EC	<ul><li>14/09/2011</li><li>08/03/2012</li></ul>	Milaxyn Plus, Strantel Plus, Prazical Plus, Voxical Plus, Exitel Plus, Cazitel Plus and Prazitel Plus and associated names
		Praziquantel, pyrantel and febantel
Referral under Art. 35 of Directive 2001/82/EC	• 15/09/2011	All long acting formulations for injection containing barium selenate for all food producing species
		Barium selenate
Procedure under Art.	• 15/09/2011	• N/a
30(3) of Regulation (EC) No 726/2004	• 11/07/2012	• Dapsone
Referral under Article 33(4) of Directive 2001/82/EC	<ul><li>12/10/2011</li><li>13/06/2012</li></ul>	Nuflor 300 mg/ml solution for injection for cattle and sheep

Type of referral	Date of clock start     CVMP opinion	Product name     INN
		Florfenicol
Referral under Article	• 12/10/2011	Hipralona Enro-S and its generics
35 of Directive 2001/82/EC	• 13/04/2012	Enrofloxacin
Referral under Article 33(4) of Directive	<ul><li>10/01/2012</li><li>13/06/2012</li></ul>	Nuflor Swine Once 450 mg/ml solution for injection
2001/82/EC		Florfenicol
Referral under Article 35 of Directive 2001/82/EC	• 12/04/2012	All injectable and pour-on veterinary medicinal products containing doramectin that are intended for use in mammalian foodproducing species
		Doramectin
Referral under Art. 34	• 15/05/2012	Micotil 300 Injectie and associated names
of Directive 2001/82/EC		• Tilmicosin
Referral under Article 33(4) of Directive	• 15/05/2012	Florgane 300 mg/ml suspension for injection for cattle and pigs
2001/82/EC		• Florfenicol
Referral under Article 33(4) of Directive	• 11/07/2012	Melosolute 40 mg/ml solution for injection for cattle, pigs and horses
2001/82/EC		Meloxicam
Referral under Article 33(4) of Directive	• 11/07/2012	<ul> <li>Strenzen 500/125 mg/g powder for use in drinking water for pigs</li> </ul>
2001/82/EC		Amoxicillin/clavulanic acid
Referral under Article 35 of Directive 2001/82/EC	• 12/09/2012	<ul> <li>Suanovil 20 and associated names, Captalin and associated names and generic products thereof, including pending applications</li> </ul>
		• Spiramycin
Referral under Article 35 of Directive 2001/82/EC	• 12/09/2012	Dexadreson 2 mg/ml and associated names, and generic products thereof, including pending applications
		Dexamethasone

#### Guidelines and working documents in 2012

#### **CVMP** Quality

Reference number	Document title	Status
EMEA/CVMP/134/02-	Draft guideline on the Active	Adopted June 2012

Reference number	Document title	Status
Rev.3/CHMP/QWP/277/02-Rev.3	Substance Master File Procedure	
EMEA/CHMP/CVMP/QWP/17760/2 009-Rev.1	Draft guideline on the Use of Near Infrared Spectroscopy by the Pharmaceutical Industry and the Data Requirements for New Submissions and Variations	Adopted for consultation, January 2012 (End of consultation 30 April 2012)
EMA/CHMP/CVMP/QWP/70278/20 12-Rev.1	Draft guideline on process validation	Adopted for consultation, March 2012 (End of consultation September 2012)
EMA/705532/2011	Questions and Answers on Post Approval Change Management Protocols	Adopted March 2012
Not applicable	Questions and Answers on the Uniformity of Dosage Units	Adopted April 2012
EMA/CHMP/CVMP/QWP/199250/2 009	Guideline on setting specifications for related impurities in antibiotics	Adopted June 2012

#### **CVMP Safety**

Reference number	Document title	Status
EMA/CVMP/SWP/355689/2006	Draft guideline on the approach to establish a pharmacological ADI.	Adopted for consultation, January 2012 (End of consultation 31 July 2012)
EMA/CVMP/SWP/878228/2011	Concept paper introducing a review and update of existing EU guidelines on residues studies to bring these into line with the VICH metabolism and residues guidelines VICH 46-49	Adopted for consultation, February 2012 (End of consultation 31 May 2012)

#### **CVMP Environmental Risk Assessment**

Reference number	Document title	Status
EMA/CVMP/ERA/409328/2010	Reflection paper on mitigation measures related to the environmental risk assessment of veterinary medicinal products testing	Adopted March 2012
EMA/CVMP/ERA/52740/2012	Draft guidance on the assessment of persistent, bioaccumulative and toxic (PBT) or very persistent and very bioaccumulative (vPvB) substances in veterinary medicine	Adopted for consultation, July 2012 (End of consultation 01 September 2012)
<u>EMA/CVMP/ERA/172074/2008 –</u>	Q&A document on the	Adopted September 2012

Reference number	Document title	Status
Rev.4	implementation of the CVMP	
	Guideline on Environmental Impact	
	Assessment for Veterinary Medicinal	
	Products in support of the VICH	
	Guidelines GL6 (Phase I) and GL38	
	(Phase II)	

#### **CVMP Efficacy**

Reference number	Document title	Status
EMA/CVMP/EWP/81976/2010 replacing EMEA/CVMP/816/00	Guideline on Statistical principles for veterinary clinical trials.	Adopted January 2012
EMA/CVMP/EWP/82829/2009- Rev.2	Revised Questions and Answers on: Testing and evaluation of the efficacy of antiparasitic substances for the treatment and prevention of tick and flea infestations in dogs and cats	Adopted July 2012

#### **CVMP Immunologicals**

Reference number	Document title	Status
EMA/CVMP/IWP/810769/2011 replacing EMEA/CVMP/865/03/final	Guideline on data requirements for removing the target animal batch safety test for immunological veterinary medicinal products in the EU	Adopted January 2012
EMA/CVMP/IWP/4199/2012	Concept paper on the need of revision of the Note for Guidance on the Harmonisation of requirements for equine influenza vaccines	Adopted for consultation, March 2012 (End of consultation 31 May 2012)
EMA/CVMP/IWP/206555/2010	Guideline on requirements for the production and control of immunological veterinary medicinal products	Adopted April 2012 Adoption of the revised version June 2012

#### **CVMP Pharmacovigilance**

Reference number	Document title	Status
EMA/CVMP/126726/2007-Rev.1	Reflection paper on risk management plans for centrally authorised veterinary medicinal products	Adopted February 2012
EMA/CVMP/PhVWP/987984/2011	Public bulletin on veterinary pharmacovigilance for 2011	Adopted February 2012
EMA/SOP/V/4025	Procedure in accordance with Article 78 of Directive 2001/82/EC related	Adopted April 2012

Reference number	Document title	Status
	to pharmacovigilance measures for veterinary medicinal products authorised in the European Union	
EMA/CVMP/10418/2009-Rev.4	CVMP combined VeDDRA list of clinical terms for reporting suspected adverse reactions in animals and humans to veterinary medicinal products	Adopted June 2012
EMA/CVMP/PhVWP/288284/2007- Rev.5	Guidance notes on the use of VeDDRA terminology for reporting suspected adverse reactions in animals and humans	Adopted June 2012
EMA/123352/2004-Rev.6	Call for comments on standard lists for EudraVigilance Veterinary	Adopted June 2012
EMA/CVMP/PhVWP/5507/2011	Concept paper for the revision of the CVMP guideline on harmonising the approach to causality assessment for adverse reactions to veterinary medicinal products	Adopted for consultation, July 2012 (End of consultation 31 October 2012)

#### Application of 3Rs (Replacement, Refinement and Reduction) in testing

Reference number	Document title	Status
EMA/CHMP/CVMP/JEG- 3Rs/252137/2012	Recommendation to marketing authorisation holders, highlighting the need to ensure compliance with 3Rs methods described in the European Pharmacopoeia	Adopted July 2012
EMA/CHMP/CVMP/JEG- 3Rs/169839/2011-Rev.1	Concept paper on the need for revision of the position on the replacement of animal studies by <i>in</i> <i>vitro</i> models	Adopted for consultation, July 2012 (End of consultation 31
		October 2012)

#### General

Reference number	Document title	Status
EMA/899273/2011	Revised list of target species for use in SPCs	Adopted February 2012
EMA/SOP/V/4003	Incident management for medicines for veterinary	Endorsed September 2012