

Woodham Lane, New Haw, Addlestone, Surrey KT15 3LS Telephone +44 (0)1932 336911

Post Authorisation Assessments

Trimediazine BMP Premix for Medicated Feeding Stuff Vm 08007/4064

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•	02 May 2018	Change in the address of the marketing authorisation holder from Vetoquinol UK Limited, Vetoquinol House, Great Slade, Buckingham Industrial Park, Buckingham, MK18 1PA to Vetoquinol UK Limited, Steadings Barn, Pury Hill Business park, Nr. Alderton, Towcester, Northamptonshire, NN12 7LS.
•	06 December 2017	Addition of a manufacturer of the active substance or addition of a site of manufacture.
•	29 December 2016	Change to manufacturer responsible for batch release.
•	29 June 2016	Deletion of a manufacturing site for an active ingredient. Submission of an updated certificate of suitability. Submission of an updated certificate of suitability.
•	15 August 2014	Deletion of a manufacturer of the finished product and primary packaging site. Deletion of a site for control of the finished product. Deletion of secondary packaging site.
•	21 September 2012	Grouped variation to submit updated European Pharmacopoeia Certificates of Suitability for already approved active substance manufacturers. Deletion of an active substance manufacturer.
•	08 February 2012	Deletion of a non-significant specification parameter.
•	12 January 2011	Variation to amend the user safety warnings on the SPC and Product Literature.
•	27 October 2010	Submission of an updated European Pharmacopoeia Certificate of Suitability for an already approved active substance manufacturer.
•	16 March 2009	Variation to change the batch release arrangements and quality control testing of the finished product.
•	16 March 2009	Variation to change the batch release arrangements and quality control testing of the finished product.
•	20 January 2009	Submission of an updated European Pharmacopoeia Certificate of Suitability for an already approved active substance manufacturer.
•	14 November 2008	Deletion of a manufacturer.
•	09 July 2008	Variation to increase the shelf life of the finished product.
•	05 September 2007	Variation to bring the SPC/Labelling in line with the Veterinary Regulations, 2005. Transfer of the legal category from POM to POM-V.
•	17 May 2007	Addition of an active substance manufacturer.
•	17 May 2007	Addition of an active substance manufacturer.
•	20 September 2006	Renewal.

•	25 April 2006	Variation to change the batch release arrangements.
•	06 March 2006	Addition of an active substance manufacturer.
•	24 February 2006	Submission of a new European Pharmacopoeia Certificate of Suitability for a new active substance manufacturer.
•	09 March 2005	Renewal.
•	22 December 2004	Addition of a secondary assembler of dosage form.
•	24 August 2004	Variation to remove a manufacturer/assembler of dosage form.
•	19 August 2004	Variation to change the address of the Marketing Authorisation Holder.
•	27 May 2004	Addition of an active substance manufacturer.
•	30 January 2004	Variation to change the site of finished product testing.
•	22 August 2003	Decrease in the meat withdrawal period (pigs).
•	25 June 2003	Addition of an assembler.
•	20 August 1999	Non-sterile container (size).
•	18 August 1997	Change to the specification of the finished product.
•	07 February 1997	Update Licence Particulars.
•	03 September 1996	Change to the shelf life of the finished product.
•	31 July 1996	Change of name and address of the PL/ATC Holder.
•	05 October 1995	Change to the formulation of the finished product.