Veterinary Medicines Directorate

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

Post Authorisation Assessments

Vasotop 2.5 mg Tablets Vm 01708/4404

•	09 September 2022	Submission of a new certificate of suitability for an
		already approved manufacturer.
•	11 February 2022	Change in storage conditions of the finished product from
		Do not store above 30°C. to Do not store above
		25°C.
		Change in the specification parameters and/or limits of
	09 June 2021	the finished product. Change in the name of the marketing authorisation
•		holder from Intervet UK Limited to MSD Animal Health
		UK Limited.
•	23 August 2018	Reduction of the shelf life of the finished product as
		packaged for sale from 24 months to 18 months.
•	04 January 2013	Changes to the SPC and product literature doe to
	47 March 0040	deletion of presentations.
•	17 March 2010	Changes to the SPC and product literature in order to maintain harmonisation across the product range.
•	08 July 2009	Change to shelf-life specifications.
•	20 June 2008	Renewal.
	28 December 2006	Changes to the SPC and product literature to bring them
		into line with new legislation.
•	09 August 2006	Change to the qualitative/quantitative composition of the
		immediate packaging material.
•	09 August 2006	Change in formulation.
•	09 August 2006	Change in the product name.
•	23 June 2006	Change to test procedure of an excipient.
•	25 May 2006	Change in the test procedure of the finished product.
•	20 July 2005	Amendments to the package leaflet.
•	23 June 2005	Change of distributor.
•	17 March 2005	Change to packaging design.
•	29 September 2004	Renewal.
•	11 April 2003	Additional manufacturer/assembler of dosage form.
•	22 June 2001	Change of distributor.
•	17 April 2000	Change in dosage schedule for standard dose.
•	29 February 2000	Change in name and address od MAH including name
		change of manufacturer of the active substance and
	20 Juno 1000	assembler of dosage from.
•	30 June 1999	Change to manufacturer of dosage form (assembly).
•	30 June 1999	Change to manufacturer of dosage form.

VMD/L4/GAT/018/C