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

Post Authorisation Assessments

Rimadyl Palatable Tablets for Dogs 50 mg

•	August 2023	Deletion of a manufacturing site for an active substance.
•	29 March 2022	Change in shape or dimensions of the container or
		closure (immediate packaging).
•	24 March 2022	Changes in the qualitative and quantitative composition
		of the immediate packaging of the finished product for
	24 March 2022	solid pharmaceutical forms.
•	24 March 2022	Changes in the qualitative and quantitative composition of the immediate packaging of the finished product for
		solid pharmaceutical forms.
•	09 April 2020	Change in the address of the marketing authorisation
		holder from Zoetis UK Limited, 5th Floor, 6 St. Andrew
		Street, London, EC4A 3AE to Zoetis UK Limited, First
		Floor, Birchwood Building, Springfield Drive,
		Leatherhead, Surrey, KT22 7LP. Change(s) in the SPC, Labelling or Package Leaflet of
		veterinary medicinal product intended to implement the
		outcome of a procedure concerning PSUR.
•	June 2018	Change in the specification limits of the finished product.
•	06 April 2016	A change to the restricted part of the ASMF.
•	30 March 2016	Deletion of a manufacturing site (for an active substance,
		intermediate or finished product, packaging site,
		manufacturer responsible for batch release, site where
		batch control takes place, or supplier of a starting material, reagent or excipient (when mentioned in the
		dossier).
•	20 October 2015	Change in name of manufacturer.
•	09 June 15	Change in dimensions of the immediate packaging
		container.
•	06 December 2013	Addition of an alternative API manufacturer.
•	14 February 2013	Variation to delete an active substance manufacturer.
•	15 April 2009	Variation to align the product specifications with the new
	14 Fobruary 2009	European Pharmacopoeia Monograph. Variation to bring the SPC/Labelling in line with the
•	14 February 2008	Veterinary Regulations, 2005. Transfer of legal category
		from POM to POM-V.
•	09 January 2008	Renewal.
•	27 March 2006	Change in the specification of an excipient.
•	23 February 2006	Batch Control.
•	07 February 2006	Change in the storage conditions of the finished product.
•	02 February 2006	Addition of a site for secondary assembly.
•	11 January 2006	Variation concerning the 'Indications'.

•	04 November 2005	Batch Control.
•	27 October 2005	Batch Control.
•	27 October 2005	Batch Control.
•	27 September 2005	Change to the packaging shape and dimensions.
•	23 June 2005	Addition of a distributor.
•	17 March 2005	Changes to the finished product.
•	23 February 2004	Change in the specifications of an excipient to comply with the European Pharmacopoeia.
•	30 September 2003	Variation to change the name of the active substance manufacturer.