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Post Authorisation Assessments

Nisamox 50 mg Tablets for Dogs and Cats

	45 Cohmison, 2022	Deletion of a new significant energification negrounder of
•	15 February 2022	Deletion of a non-significant specification parameter of an excipient.
•	10 December 2019	Addition of a secondary packaging site of the finished
		product.
		Submission of an updated Ph. Eur. certificate of
		suitability for an active substance from an already
		approved manufacturer. Submission of an updated Ph. Eur. certificate of
		suitability for an active substance from an already
		approved manufacturer.
	29 November 2019	Minor change in the manufacturing process of an
	20 110 101111101 20 10	immediate release solid oral dosage form.
		Qualitative / quantitative changes to the excipients.
•	03 June 2019	Addition of a manufacturer responsible for batch release
		of the finished product.
•	14 January 2019	Change in RMS from UK to ES.
•	04 January 2019	Update of the test procedure to comply with the updated
		general Ph. Eur monograph.
		Changes to a test procedure for the finished product.
•	28 December 2017	Change in distributor details from Bayer plc, Animal
		Health, Strawberry Hill, Newbury, RG14 1JA to Bayer
		plc, 400 South Oak Way, Green Park, Reading, RG2 6AD.
	23 March 2016	Submission of a new or updated Ph. Eur. certificate of
	25 Maion 2010	suitability
		Submission of a new or updated Ph. Eur. certificate of
		suitability
		Submission of a new or updated Ph. Eur. certificate of
		suitability
		Deletion of a Ph. Eur. certificate of suitability
		Deletion of a Ph. Eur. certificate of suitability
		Submission of a new or updated Ph. Eur. certificate of
	28 November 2014	suitability Update to the DDPS.
•	03 January 2014	Submission of updated Ph. Eur. Certificates of Suitability
•	US January 2014	for an already approved manufacturer.
•	24 November 2011	Changes to SPC section 2 and appropriate sections of
		the Product Literature.
•	03 June 2011	Change of product name in Italy only.
•	27 April 2011	Removal of a manufacturing site of the active substance.
•	10 November 2010	Change of distributor.
•	05 August 2009	Renewal.

•	25 April 2008	Change in shelf life from 18 months to 24 months.
•	09 April 2008	Change to composition of immediate packaging.
•	17 April 2007	Addition of a manufacturer of the active substance.
•	24 January 2007	Addition of a target species – Cats.
•	28 October 2005	Change of shelf life from 1 year to 18 months.
•	15 November 2004	Mutual Recognition Procedure, UK as RMS.
•	16 May 2003	Additional presentation.