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Post Authorisation Assessments

For details of post authorisation assessments prior to 1st January 2021, please refer to the <u>EMA</u> website.

Simparica 5 mg Chewable Tablets for Dogs 1.3-2.5 kg

Vm 42058/5052

	18 May 2024	Alternate test method for a starting material added.
•	04 May 2024	Addition of a new specification parameter for a starting
		material.
•	23 February 2024	Change in the shelf-life or storage conditions of the finished
		product.
•	22 February 2024	One-off alignment of the product information with version 9.0*
		of the QRD templates.
•	22 December 2023	Editorial changes to Part 2 of the dossier.
		Editorial changes to Part 2 of the dossier.
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		Editorial changes to Part 2 of the dossier.
		Minor changes to an approved test procedure for the finished
		product.
		Minor changes to an approved test procedure for the finished
		product. Minor changes to an approved test procedure for the finished
		product.
•	18 October 2023	Extension of the re-test period of the active substance where
	10 0000001 2020	no Ph. Eur. Certificate of Suitability covering the retest period
		is part of the approved dossier.
•	18 September 2023	Minor changes in the manufacturing process of the drug
	10 00010111201 2020	product intermediate.
		Addition of a site for the manufacturing process of the drug
		product intermediate.
•	14 September 2023	Change in batch size of the drug product intermediate.
		Minor changes to the registered method for the drug product
		intermediate.
		Minor changes to the registered method for the drug product
		intermediate.
		Minor changes to the registered method for the drug product
		intermediate.
•	31 July 2023	Change in batch size of finished product.
		Change in batch size of finished product.
	47 Amril 2022	Change in batch size of finished product.
•	17 April 2023	Addition of an alternative supplier of a starting material.
•	20 February 2023	Deletion of packaging components suppliers. Additional indication: For reduction of the risk of infection with
•	17 February 2023	Babesia canis canis via transmission by Dermacentor
		reticulatus for 28 days after treatment. The effect is indirect
		due to the product's activity against the vector.
		Associated warning in Section 4.4
•	22 December 2022	Addition of a secondary packaging site of a finished product.
•	22 December 2022	Addition of a primary packaging site of a non-sterile finished
		product.
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•	31 October 2022	Change in name and address of a manufacturer of the active substance.
•	19 October 2022	Changes to labelling to include GB details in blue box.
•	23 August 2022	Change in the name of a supplier of the active substance. Change in the name of a supplier of the active substance. Deletion of a supplier of the active substance. Deletion of a supplier of the active substance.
•	31 May 2022	Change in the name of a supplier of starting material.
•	09 March 2022	Changes to a test procedure for the immediate packaging of the active substance. Change in manufacturer of the active substance.