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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 40 mg Chewable Tablets for Dogs >10-20 kg

Vm 42058/5051

| | 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. |
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| | | Minor changes to an approved test procedure for the finished |
| | | product. |
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| | | product. |
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| | | product. |
| • | 18 October 2023 | Extension of the re-test period of the active substance where |
| | | no Ph. Eur. Certificate of Suitability covering the retest period |
| | 10.0 | is part of the approved dossier. |
| • | 18 September 2023 | Minor changes in the manufacturing process of the drug |
| | | product intermediate. |
| | | Addition of a site for the manufacturing process of the drug |
| | 44.0 | 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. |
| | 0. 34., 2020 | Change in batch size of finished product. |
| | | 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. |
| • | 17 February 2023 | Additional indication: For reduction of the risk of infection with |
| | ,==== | 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. |

| • | 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. |