

Post Authorisation Assessments

Milbemax 12.5 mg/125 mg chewable tablets for dogs Vm 00879/5031

| • | 13 April 2024 | One-off alignment of the product information with version |
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| | | 9.0* of the QRD templates. |
| • | 19 January 2024 | Editorial changes to SPC, package leaflet or labelling. |
| • | 29 August 2023 | Deletion of Heavy metals from specifications of natural chicken flavour. |
| • | 28 July 2023 | Submission of an updated Ph. Eur. CEP from an already approved manufacturer for a non-sterile: – active substance. |
| • | 14 September 2022 | Change in the specification limits of the finished product. |
| • | 04 February 2022 | Deletion of manufacturing site for an active substance. |
| • | 14 October 2021 | Submission of an updated Ph. Eur. certificate of suitability for an active substance from an already approved manufacturer. |
| • | 11 February 2021 | Addition of a new specification parameter with its corresponding test method of a reagent used in the manufacturing process of the active substance. 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. |
| • | 02 February 2021 | Addition of a secondary packaging site of the finished product. Addition of a primary packaging site of the finished product. |
| • | 28 September 2020 | Minor changes to an approved test procedure of the finished product. Minor changes to an approved test procedure of the finished product. Minor changes to an approved test procedure of the finished product. |
| • | 16 September 2020 | Change in the address of the Marketing Authorisation Holder from Elanco Europe Ltd, Lilly House, Priestley Road, Basingstoke, Hampshire, RG24 9NL, United Kingdom to Elanco Europe Ltd, Form 2, Bartley Way, Bartley Wood Business Park, Hook, RG27 9XA, United Kingdom. |
| • | 10 December 2019 | Minor changes to an approved test procedure of the finished product. Minor changes to an approved test procedure of the finished product. |

| | | Change to comply with an update of the relevant monograph of the Ph. Eur. or national pharmacopoeia of a Member State. |
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| • | 05 June 2019 | Change in the safety database of an existing pharmacovigilance system as described in the DDPS. |
| • | 29 May 2019 | 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. |
| • | 04 April 2019 | Minor change to an approved test procedure for an excipient. Minor change to an approved test procedure for an excipient. Minor change in the manufacturing process of the finished product. |
| | | Change to comply with an update of the relevant monograph of the Ph. Eur. or national pharmacopoeia of a Member State. Addition of an alternative site of an excipient. Addition of a manufacturing site of the finished product. |
| • | 29 January 2019 | Change in the address of a manufacturer used in the manufacture of the active substance. |
| • | 04 October 2018 | Change of specifications of a former non Pharmacopoeial active substance to comply with the Ph. Eur. |
| • | 27 September 2018 | Change in the SPC, Labelling or Package Leaflet of veterinary medicinal products intended to implement the outcome of a procedure concerning PSUR. |
| • | 20 February 2018 | Change in the specification limits of the finished product. Change in the specification limits of the finished product. |
| • | 24 August 2017 | Addition of a supplier of packaging components or devices |
| • | 14 August 2017 | Addition of a new specification parameter with its corresponding test method of an active substance used in the manufacturing process of the active substance Changes to a test procedure (including replacement or addition) for the active substance Introduction of a new site of manufacture Submission of a new Ph. Eur. certificate of suitability for an active substance from a new manufacture |
| • | 21 June 2017 | Change in batch size (including batch size ranges) of active substance used in the manufacturing process of the active substance Change in the specification parameters or limits of an active substance |
| • | 07 March 2017 | Introduction of a new pharmacovigilance system. |
| • | 15 February 2017 | Minor change in the manufacturing process. Addition of a manufacturing site of the finished product. |
| • | 06 September 2016 | Minor changes to sections 4.4 & 4.8 of the SPC. |
| • | 15 August 2016 | Change in the name of a manufacturer of the finished product including manufacturer responsible for batch release |
| • | 06 July 2016 | Change in the name of the Marketing Authorisation Holder from Novartis Santé Animale to Elanco France in France |

| | | and Poland only. |
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| • | 06 July 2016 | Change in the name and address of the Marketing |
| | | Authorisation Holder in Spain and Italy only. |
| • | 16 March 2016 | Change in distributor details |
| | | Change in legal entity |
| • | 15 September 2015 | Change in the immediate packaging of the finished product. |
| • | 24 July 2015 | Submission of an updated TSE Certificate of Suitability. |
| • | 02 May 2015 | To add a new pack size for the finished product. |
| • | 02 March 2015 | Minor changes to sections 4.2 and 4.8 of the SPC. |
| • | 05 November 2014 | Renewal. |
| • | 30 April 2014 | Changes to an existing pharmacovigilance system as described in the DDPS. |
| • | 27 March 2014 | Submission of updated Ph. Eur. Certificates of Suitability. |
| • | 07 February 2014 | Change in MAH address. |
| • | 18 July 2013 | Change in address of manufacturer/supplier of reagent used in the manufacture of the active substance. Change in the manufacturing process of the active substance. |
| | | Change of in process limits applied during manufacture of the active substance. Change to in process tests applied during the manufacture |
| | | of the active substance. Change in specification limits of a reagent involved in the manufacture of the active substance. Minor amendments to part 2 of the dossier. |
| | | Inclusion of restricted part of ASMF in part 2 of the dossier. |
| • | 05 July 2013 | To change the address of the MAH in Denmark, Finland, |
| | | Norway and Sweden and corrections to the address of the MAH in Cyprus. |
| • | 25 June 2013 | Addition of new therapeutic indications. |
| • | 28 March 2013 | To change the address of the marketing authorisation holder (MAH) in France only. |
| • | 16 January 2013 | To reduce the shelf life of the finished product as packaged for sale from 3 years to 2 year. |
| • | 12 November 2011 | Changes to the DDPS that do not impact on the operation of the pharmacovigilance system. |
| • | 30 August 2012 | Variation to change the manufacturer of the tablets. |
| • | 15 February 2012 | Changes to the dimensions of the artwork. |
| • | 07 September 2011 | To change the immediate packaging of the finished product. |
| • | 11 July 2011 | Grouped variation to introduce the batch size range from 100 kg to 500 kg and to make minor changes to the manufacturing process. |
| • | 18 April 2011 | Grouped variation to add an additional manufacturer of the active substance and also to submit an updated Ph. Eur. certificate of suitability for approved manufacturer. |
| • | 06 August 2010 | Grouped variation to add primary and secondary packaging sites. |
| • | 06 August 2010 | Grouped variation to change the name and contact details of the qualified person for pharmacovigilance. |