

Post Authorisation Assessments

Engemycin 10% DD Solution for Injection Vm 01708/4623

•	23 January 2024	Addition of a test procedure for the finished product.
•	16 August 2023	Deletion of a manufacturing site of the finished product.
•	31 July 2023	Change in immediate packaging of the finished product.
•	10 August 2022	Change in excipient specification to comply with the Ph.Eur.
•	23 December 2021	Addition of a site where batch control/testing takes place. Addition of a site where batch control/testing takes place. Addition of a manufacturer responsible for batch release where batch control/testing takes place. Addition of a manufacturing site of the finished product. Addition of secondary packaging site of the finished product.
•	13 July 2021	Change in distributor details. From various addresses to Same as MAH, Alternative distributor in Northern Ireland, Intervet Ireland Limited, Magna Drive, Magna Business Park, Citywest Road, Dublin 24.
•	11 January 2021	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. 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. Submission of an updated Ph. Eur. certificate of suitability for an active substance from an already approved manufacturer.
•	03 November 2020	Submission of a new Ph. Eur. certificate of suitability for an active substance from an already approved manufacturer.
•	17 July 2020	Change of MAH from Intervet International BV represented by: Intervet UK Ltd. Walton Manor, Walton, Milton Keynes, Buckinghamshire, MK7 7AJ to MSD Animal Health UK Limited, Walton Manor, Walton, Milton Keynes, Buckinghamshire, MK7 7AJ.
•	23 June 2020	Deletion of manufacturing site for an active substance
•	24 May 2016	Deletion of a manufacturing sites (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). Submission of a new or updated Ph. Eur. certificate of suitability
•	20 August 2013	Submission of an updated Ph. Eur. Certificate of Suitability for an active substance
•	16 February 2012	Submission of an updated Ph. Eur. Certificate of Suitability for an active substance
•	09 February 2011	Submission of an updated Ph. Eur. Certificate of Suitability for an active substance

 30 December 2008 Updates to section 4.11 of the SPC 12 September 2007 Change of batch size 03 April 2007 Addition of a manufacturer of an active substance 09 November 2006 Change of legal category from POM to POM-V Changes to the SPC and Product Literature to bring in line with new legislation 02 May 2006 Renewal 23 June 2005 Change of distributor 21 April 2005 Renewal Change of withdrawal period for: Meat from Pigs from 8 days to 14 days Meat from Cattle – from 8 days to 21 days (prolonged dosing regimen) 28 January 2005 Change of withdrawal period for: Milk from Sheep – not for use in sheep producing milk for human consumption Milk from Cattle – from 48 hours (24hour dosing regimen)/72 hours (prolonged action dosing regimen) to 6 days for all dosing regimens Meat from Sheep – from 8 days to 14 days 23 April 2003 Addition of a manufacturing site for batch release 22 August 2001 Change of distributor 23 April 2003 Change of distributor 24 August 1998 Deletion of a target species 27 October 1997 Change of type of sterile container 			
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