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Post Authorisation Assessments

Nuflor 40 mg/g Premix for Medicated Feeding Stuff for Swine Vm 01708/3042

•	22 March 2024	Change in the specification parameters of an active substance.
•	February 2024	Change in the specification parameters of an active substance.
•	09 February 2023	Tightening of specification limits for the active substance.
•	27 January 2022	Addition of a manufacturer of the active substance or addition of a site of manufacture.
•	14 August 2020	Change in the name of the marketing authorisation holder from Intervet UK Limited to MSD Animal Health UK Limited.
•	07 June 2018	Updates to the SPC and QRD texts.
•	29 June 2016	Change in the manufacturer of a starting material used in the manufacturing process.
•	06 March 2015	Change in the batch size of the active substance. Tightening of specification limits of the active substance.
•	20 November 2014	Change in test procedure for the active substance. Change in the re-test period of the active substance. To introduce a second manufacturing process.
•	31 July 2014	Deletion of a manufacturing site for the active substance.
•	17 February 2012	To change the specification parameter and/or limits of the finished product.
•	25 January 2012	Variation to change the MAH and distributor from Schering Plough Ltd to Intervet UK Ltd.
•	23 January 2012	Deletion of an active substance manufacturer.
•	20 December 2011	Renewal procedure – France as RMS.
•	16 December 2011	To add an additional manufacturing site. To add an alternative site for primary and secondary packaging operations and batch control/release testing of the finished product. Minor changes to the manufacturing process.
•	02 September 2011	To update the specification limits of the active substance.
•	14 March 2011	To change the name of the MAH in Portugal only.
•	09 February 2011	To add a new manufacturer of the active substance.
•	18 October 2007	Change in part of primary packaging material not in contact with finished product. Change in composition of immediate packaging: all other pharm forms.