



## Post Authorisation Assessments

### Panacur Equine Granules 22.2% w/w Vm 01708/4424

•	13 September 2023	Change in the specification limits of the finished product. Change to the method description of an in-process control test. Change to an in-process control test limit applied during the manufacture of the finished product.
•	07 July 2023	Delete odour testing specification parameter of the active substance.
•	23 November 2022	Extension or introduction of a re-test period/storage period supported by real time data. Minor change to the restricted part of an Active Substance Master File.
•	02 August 2021	Increase in batch size (from 400 - 500 kg to 400 - 600 kg) of the finished product.
•	30 December 2020	Change in the name of the MAH from Intervet UK Limited to MSD Animal Health UK Limited.
•	17 August 2020	Addition of a new specification parameter with its corresponding test method of the active substance. Minor change to the restricted part of an Active Substance Master File.
•	27 April 2020	Addition of a manufacturer of the active substance or addition of a site of manufacture.
•	21 December 2016	Minor change in the manufacturing process of the active substance.
•	08 June 2016	Reduction of the shelf life of the finished product as packaged for sale from 5 years to 36 months.
•	18 January 2012	Change of logo and formatting on the product literature.
•	18 September 2008	Changes to the SPC and product literature to bring them into line with new legislation.
•	18 September 2008	Change of legal category from PML to POM-VPS.
•	03 June 2008	Deletion of a 1kg pack size.
•	03 June 2008	Replacement of a site of manufacturer of finished product and batch release.
•	19 March 2008	Change of equipment used for an in-process control.
•	21 February 2008	Change in batch size of the finished product.
•	24 August 2007	Renewal
•	28 March 2007	Change in the test procedure of the finished product.
•	12 May 2005	Change of distributor for Northern Ireland.
•	04 September 2003	Renewal.
•	03 July 2001	Addition of a distributor for Northern Ireland.
•	30 November 2000	Change in manufacturer and assembler of dosage form.

•	17 March 2000	Change in name and address of the MAH.
•	18 June 1998	Renewal.
•	23 February 1998	Addition of a manufacture of the active substance.
•	19 September 1996	Change of MAH.
•	19 September 1996	Addition of a manufacturer and assembler of the finished product.
•	05 September 1996	Change in therapeutic indications.
•	15 August 1996	Change in dosage and administration.