

Post Authorisation Assessments

AquaVac Vibrio Immersion and Injection Vm 01708/4569

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•	07 June 2023	Change in name and address details of a manufacturer of the finished product.
		Change in name and address details of a manufacturer
		of the active substance. (NI)
•	22 December 2022	Change in name and address details of a manufacturer of the finished product.
		Change in name and address details of a manufacturer
		of the active substance.
•	08 October 2021	Addition to a test procedure for the finished product.
		Addition of a site where batch control/testing takes place.
•	01 October 2020	Change in the name of a manufacturer of the finished
		product, also responsible for batch release.
•	14 August 2020	Change in the name of the marketing authorisation holder from Intervet UK Limited to MSD Animal Health
		UK Limited.
•	20 December 2018	Replacement of a manufacturer responsible for batch
		release of the finished product.
		Addition of a batch release site of the finished product.
		Replacement of a secondary packaging site of the
		finished product.
•	25 April 2014	Change of Manufacturing Authorisation Holder.
•	18 January 2012	Change in immediate packaging of the finished product.
•	02 September 2011	To change the MAH in Portugal only.
•	13 July 2011	Renewal Marketing Authorisation – France as RMS.
•	09 February 2011	To change the MAH from Schering-Plough Ltd to Intervet UK Ltd.