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Post Authorisation Assessments

AquaVac ERM Concentrate for Dip Suspension for Rainbow Trout Vm 01708/5124

•	27 June 2023	Change in the name and address of the
		manufacturer of the finished product. (GB)
		Change in the name and address of the
	07.0 1 1 0001	manufacturer of the active substance. (GB)
•	27 September 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
•	01 October 2020	finished product, also responsible for batch
		release.
	14 August 2020	Change in the name of the marketing
	, tagast 2020	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.
		Replacement of a manufacturer responsible for
		batch release of the finished product.
		Deletion of a packaging site.
		Replacement of a secondary packaging site of the
		finished product.
•	09 February 2018	Change in the RMS from UK to IE.
•	08 July 2015	Change to the design and technical specifications
	45.14	of the product labels.
•	15 May 2014	Change in the name and address of the
	07.1	manufacturing authorisation holder.
•	27 January 2012	Change in the immediate packaging for the
	04 August 2011	finished product. To change the name of the MAH in Portugal only,
•	04 August 2011	from Schering-Plough II to Intervet Portugal.
•	15 October 2010	Renewal.
	08 September 2009	Change of Marketing Authorisation Holder.
	14 August 2009	Change of Marketing Authorisation Holder.
•		Variation to increase the size of maximum
•	10 January 2007	
	11 December 2006	blending volume of final product. Variation to change Marketing Authorisation
•	TI December 2000	Holder in Denmark only.
_	21 March 2006	MRP procedure – UK as RMS.
•	2 1 Water 2000	IVII \(\text{PIOCECUTE} = OI\ as I\\ IVIO.