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Post Authorisation Assessments

Hyogen Emulsion for Injection for Pigs Vm 15052/5046

•	28 April 2024	Change in name of a quality control testing site.
•	28 March 2024	A new final product control test, testing the extractable
		volume of the finished product in addition to currently
		applied filling volume test performed during the filling
		process.
		To extend the shelf life of the vaccine from 15 months to
		24 months.
		An additional new test measuring the paraffin oil content
		in the vaccine. A sandwich ELISA assay for antigen quantification using
		the same methodology, like the proposed in-vitro
		sandwich ELISA potency test. The proposed test is
		intended as in-process control test on the inactivated,
		concentrated antigen suspension.
		A new in-vitro method (sandwich ELISA assay) for
		control the potency of final product, replacing the current
		one, which is an in-vivo/iv-vitro test system measuring
		rabbit serology.
		Follow-up updates based on earlier variations.
		One-off alignment of the product information with version
	07.1	9.0* of the QRD templates.
•	27 January 2023	Increase in the batch size of the finished product.
•	13 October 2022	Change in the address of the MAH from Unit 3 Anglo
		Office Park, White Lion Road Amersham, Buckinghamshire HP7 9FB to Explorer House, Mercury
		Park, Wycombe Lane, Wooburn Green, High Wycombe,
		Buckinghamshire, HP10 0HH, United Kingdom.
•	16 March 2022	Change in the SPC, labelling or package leaflet due to
	_	new data.
•	06 September 2021	Change in the fill volume of the finished product.
•	06 July 2021	Change in the manufacturer of a starting material used in
		the manufacturing process of the active substance.
•	23 June 2021	Replacement to a test procedure for the finished product.
•	10 June 2020	Replacement of a site where batch control/testing takes
	0	place.
•	27 March 2020	Renewal - UK as CMS.
•	21 November 2019	Changes in the SPC, Labelling or Package Leaflet of
		human medicinal products intended to implement the
		outcome of a procedure concerning PSUR or PASS, or
		the outcome of the assessment done by the competent
		authority under Articles 45 or 46 of Regulation

		1901/2006.
•	21 November 2019	Increase in batch size of active substance or
		intermediate used in the manufacturing process of the
		active substance.
•	01 October 2019	Change to an approved stability protocol.
•	22 May 2019	Change in the fill volume of the finished product.
•	08 April 2019	Change in test procedure for an excipient.
•	31 January 2019	Change in the invented name of the veterinary medicinal product from Hyobloc (SE) and Mhyogen (DK) to Mhyogen vet in SE and DK.
•	19 September 2017	Change in the QPPV of an existing pharmacovigilance system as described in the DDPS. Change of the back-up procedure of the QPPV of an existing pharmacovigilance system as described in the DDPS.
•	08 August 2017	Replacement to a test procedure for the finished product.
•	06 October 2016	Change in the specification limits of the finished product. Change in the manufacturing process of the active substance.
•	20 September 2016	Addition of an alternative site for the animal testing phase of the potency test.
•	06 January 2016	Submission of an updated DDPS.