## Woodham Lane, New Haw, Addlestone, Surrey KT15 3LS Telephone +44 (0)1932 336911

## **Post Authorisation Assessments**

## Vitamin B1 10% w/v Solution for Injection Vm 50146/4029

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• 01 August 2023	Change in the manufacturing process of the finished product.
	Change in immediate packaging of the finished product: -
	Change in type of container or addition of a new
	container.
• 18 March 2021	Replacement of a secondary packaging site of the
	finished product.
• 26 January 2021	Change in the QPPV of an existing pharmacovigilance
	system as described in the DDPS.
• 23 July 2019	Change in the name of a manufacturer used in the
	manufacture of the active substance.
	Change in the name and address of a manufacturer of
05.0-1-1-0040	the finished product, also responsible for batch release.
• 25 October 2018	Change of the back-up procedure of the QPPV of an
	existing pharmacovigilance system as described in the DDPS.
	Change of MAH, from Bimeda Chemicals Ltd., Broomhill Road, Tallaght, Dublin 24, Ireland to Bimeda Animal
	Health Limited, 2 / 3 / 4 Airton Close, Tallaght, Dublin 24,
	Ireland.
• 15 October 2018	Change in distributor details. From Bimeda Chemicals
10 0010201 2010	Ltd, Broomhill Road, Tallaght, Dublin 24, Ireland to
	Bimeda ®, Cross Vetpharm Group UK Ltd., Unit 2, Bryn
	Cefni, Llangefni, Anglesey, LL77 7XA, United Kingdom
• 03 July 2013	Submission of an updated European Pharmacopoeia
	Certificate of Suitability from a currently approved active
	substance manufacturer.
• 17 May 2012	Grouped variation to change the finished product
	specification.
• 28 July 2010	Grouped variation to change the name of an active
	substance manufacturer via the submission of an
	updated European Pharmacopoeia Certificate of
	Suitability from a currently approved active substance
	manufacturer.
• 08 January 2009	Variation to bring the SPC/Labelling in line with the
	Veterinary Regulations, 2005. Transfer of the legal
10 Contomber 2000	category from POM to POM-V.
• 10 September 2008	Variation to the formulation of the finished product.
• 17 January 2008	Renewal.
• 12 December 2003	Renewal.
<ul> <li>22 May 2003</li> </ul>	Addition of an assembler of dosage form.

ſ	•	16 November 2001	Addition of a dosage form manufacturer.
ſ	•	26 July 1999	Renewal.