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Post Authorisation Assessments

Nobivac KC Vm 01708/3022

•	03 April 2024	G.I.18 update of product information.
•	26 February 2024	To mention the use of animal derived trypsin in the manufacture of the Byco-C hydrolysed gelatin provided by Croda and to declare a change in the hydrolysis step of the gelatin (excipient) by replacing porcine trypsin by the use of either porcine or bovine trypsin.
•	02 November 2023	Extension of the storage period of antigen.
•	23 December 2021	Deletion of a pack size(s) of the finished product. Change in the SPC, labelling or package leaflet due to new data.
•	28 May 2021	Change in the address of a manufacturer used in the manufacture of the active substance.
•	15 April 2021	Change in the SPC, Labelling or Package Leaflet following a periodic safety update report (PSUR).
•	10 June 2020	Change of MAH from Intervet International BV, Represented by:, Intervet UK Ltd., Walton Manor, Walton, Milton Keynes, Bucks, MK7 7AJ to MSD Animal Health UK Limited, Walton Manor, Walton, Milton Keynes, Buckinghamshire, MK7 7AJ.
•	02 April 2020	Change in the manufacturing process of the active substance.
•	17 April 2019	Changes in the SPC, Labelling or Package Leaflet intended to implement the outcome of a procedure concerning PSUR
•	13 November 2018	Change in the safety database of an existing pharmacovigilance system as described in the DDPS
•	07 March 2018	Change in RMS from UK to IT
•	13 June 2017	Changes in the manufacturing process of the finished product.
•	16 December 2016	Change in the safety database of an existing pharmacovigilance system as described in the DDPS.
•	16 December 2015	An addition of an alternative sterilisation method for stoppers.
•	16 December 2015	An introduction of a real-time PCR test.

implement outcome of PSUR. 29 April 2015 Approval of mock-ups. Removal of the administration device from the packaging. 04 December 2014 Update to the DDPS. 23 July 2013 Significant changes to the SPC/PL. 30 August 2012 Renewal. 17 June 2010 Changes to an existing pharmacovigilance system. 21 October 2009 Change of production site. 01 September 2009 Change of MAH name/address. 17 December 2008 Minor amendments to production process addition of an alternative manufacturer, Repeat Use procedure.
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03 April 2008 Repeat Use procedure.
03 April 2008 Renewal procedure.
30 January 2008 Simple text changes to SPC and product literature.
08 January 2008 To change product literature to comply with CMD\ referral.
07 December 2007 New, updated TSE Ph. Eur CofS for excipier from an approved manufacturer.
14 December 2005 Other.
11 August 2005 Variation to comply with Ph. Eur. or MS national pharmacopoeia.
20 June 2005 Variation to change the distributor.
25 January 2005 Addition of a manufacturer responsible for blending, filling and freeze drying.
09 June 2004 Extension of the duration of immunity.
22 March 2002 Change to the ingredient specification according to TSE compliance.
27 December 2001 Addition of a distributor.
27 December 2001 Change of finished product formulation.

•	19 October 2000	MRP – UK as RMS.
•	30 June 2000	Change of name and address of the Marketing Authorisation Holder.
•	31 March 2000	Change to the therapeutic purpose.