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Post Authorisation Assessments

Anthelmin Plus Flavour Tablets for Dogs Vm 01656/4015

•	23 February 2024	Update to a Ph. Eur. CEP for an already authorised manufacturer of the active substance. (GB)
•	23 February 2024	Update to a Ph. Eur. CEP for an already authorised manufacturer of the active substance. Update to a Ph. Eur. CEP for an already authorised manufacturer of the active substance. (NI)
•	19 January 2024	Introduction of a summary of the PSMF or changes to the summary of the PSMF not already covered elsewhere in this Annex. (NI)
•	January 2024	Update to a Ph. Eur. CEP for an already authorised manufacturer of the active substance. (GB)
•	04 December 2023	Introduction of a summary of the PSMF. (GB)
•	05 October 2023	Substantial changes in the updated version of the ASMF.
•	07 June 2023	Submission of a new certificate of suitability.
•	16 March 2023	New certificate of suitability from a new manufacturer.
•	06 January 2023	Updated certificate of suitability from an already approved manufacturer.
•	11 October 2022	Updated certificate of suitability from an already approved manufacturer.
•	07 April 2022	Submission of an updated Ph. Eur. certificate of suitability for an active substance from an already approved manufacturer. Submission of an updated Ph. Eur. certificate of suitability for an active substance from an already approved manufacturer.
•	13 April 2021	Minor changes to an approved test procedure of the finished product.
•	16 March 2021	Changes to the labelling and/or package leaflet.
•	26 January 2021	Submission of an updated Ph. Eur. certificate of suitability for an active substance from an already approved manufacturer. Submission of an updated Ph. Eur. certificate of suitability for an active substance from an already approved manufacturer.
•	25 November 2020	Submission of a new Ph. Eur. certificate of suitability for an active substance from a new manufacturer.
•	14 August 2020	Changes to the labelling.
•	09 July 2020	Minor change in the manufacturing process of the finished product. Minor change in the manufacturing process of the

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		finished product.
		Minor change in the manufacturing process of the
		finished product. Minor change in the manufacturing process of the
		finished product.
		Minor change in the manufacturing process of the
		finished product.
		Minor change in the manufacturing process of the
		finished product.
		Addition of a manufacturer responsible for batch
		release of the finished product.
		Change in the shape or dimensions of the
		pharmaceutical form.
	40 A	Addition of a manufacturing site of the finished product.
•	16 August 2019	Addition of a site where batch control/testing takes place.
	27 June 2019	Submission of an updated Ph. Eur. certificate of
•	21 Julio 2019	suitability for an active substance from an already
		approved manufacturer.
		Submission of an updated Ph. Eur. certificate of
		suitability for an active substance from an already
		approved manufacturer.
•	27 June 2019	Addition of a site where batch control/testing takes
		place.
		Deletion of manufacturing site for finished product.
		Replacement of a secondary packaging site of the
		finished product.
		Replacement of a primary packaging site of the finished product.
•	11 June 2019	Change in the contact details of the QPPV of an
	11 04110 2010	existing pharmacovigilance system as described in the
		DDPS.
•	26 October 2018	Change in RMS from UK to IE.
•	23 October 2018	Update to the Local Representative details.
•	25 July 2017	Change to comply with an update of the relevant
		monograph of the Ph. Eur. or national pharmacopoeia
		of a Member State.
•	12 April 2017	Deletion of manufacturing site for an active substance
•	01 December 2016	Mock-ups approved.
•	07 October 2016	Addition of a new test method.
	OF Contombox 2010	Addition of an active substance manufacturer.
•	05 September 2016	Change in the (invented) name of the medicinal product
	05 August 2016	in Spain and Portugal. Increase in the shelf-life of the finished product as
•	00 August 2010	packaged for sale, from 2 to 3 years.
•	02 June 2016	Renewal – UK as RMS
	25 November 2015	To update the administration advice in section 4.9 of
	20 110 10111111111111111111111111111111	the SPC
•	22 April 2015	Addition of UK local representative information to
		package leaflet.
•	23 March 2015	Change in distributor details.
•	09 May 2014	Receipt of an updated Certificate of Suitability for an
		active substance.

•	19 December 2013	To update the pharmacovigilance system.
•	19 December 2013	Deletion of a manufacturing site for primary and secondary packaging.
•	01 November 2013	Submission of an updated Ph. Eur. certificate of suitability.
•	21 March 2013	Changes to the labelling or the package leaflet which are not connected with summary of product characteristics
•	01 March 2013	Deletion of a secondary packaging site and deletion of a manufacturing site responsible for batch release.
•	12 October 2012	Deletion of a manufacturing site for the active substance febantel; deletion of a manufacturing site for the active substance praziquantel; addition of a primary packaging site for the finished product; addition of a manufacturing site (not responsible for batch release or batch control) for the finished product.
•	10 August 2012	Change of name of veterinary medicinal product.
•	06 January 2012	Submission of a new or updated Ph. Eur. certificate of suitability.