

## **Post Authorisation Assessments**

## Prevensa 250 mg + 62.5 mg Spot-on Solution for Large Dogs Vm 00879/4148

•	19 December 2022	Extension of the re-test period of the active substance
	03 March 2022	Removal of all references to Local Representative.
•		Change in the SPC, labelling or package leaflet following
		assessment of the same change for the reference
		product.
•	25 June 2021	Minor change to an approved test procedure for the
		active substance used in the manufacturing process of
		the active substance.
		Introduction of a re-test period of the active substance.
		Submission of a new Ph. Eur. certificate of suitability for
		an active substance from a new manufacturer.
•	23 October 2020	Change of MAH from Bayer plc, 400 South Oak Way,
		Green Park, Reading, Berkshire, RG2 6AD to Elanco
		Europe Ltd., Form 2, Bartley Way, Bartley Wood
		Business Park, Hook, RG27 9XA, United Kingdom.
•	24 April 2020	Renewal – National.
•	18 September 2018	Change in distributor details from Bayer plc, Animal
		Health Division, Strawberry Hill, Newbury, Berkshire, RG14 1JA to Bayer plc, 400 South Oak Way, Green
		park, Reading, Berkshire, RG2 6AD.
	29 August 2018	Change in the invented name of the veterinary medicinal
	207/090012010	product from Multi-parasite 250 mg + 62.5 mg Spot-on
		Solution for Large Dogs to Prevensa 250 mg + 62.5 mg
		Spot-on Solution for Large Dogs.
•	20 June 2018	Change in the SPC, labelling or package leaflet following
		assessment of the same change for the reference
		product.
•	27 July 2017	Change in the name of a manufacturer used in the
		manufacture of the active substance.
•	05 May 2017	Change in the address of the marketing authorisation
		holder from Bayer plc, Animal Health Division, Bayer
		House, Strawberry Hill, Newbury, Berkshire, RG14 1JA
		to Bayer plc, 400 South Oak Way, Green Park, Reading,
-	01 September 2016	Berkshire, RG2 6AD. Changes to SPC and product literature following the
•		assessment of the same change for the reference
		product.
•	24 May 2016	Changes to a DDPS following the assessment of the
		same DDPS in relation to another medicinal product of
		the same MAH
•	17 May 2016	Addition of a new therapeutic indication for the treatment
	-	of cutaneous dirofilariosis (adult stages of Dirofilaria

		repens.
•	8 December 2015	Addition of a secondary packaging site
•	25 August 2015	Change in test procedure for the finished product.
•	30 June 2015	Submission of an updated certificate of suitability.