

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

## **Post Authorisation Assessments**

## Advantage 80 mg Spot-on Solution for Cats (>8 kg) Vm 00879/4106

Update to antiparasitic warnings as per EMA guideline. Alignment of the product information with version 9.0* of the QRD templates.  Or April 2025 Change(s) in the name or address or contact details of a qualified person for pharmacovigilance. (GB)  19 January 2025 Deletion of a non-significant specification parameter in the shelf-life specification of the finished product. (NI).  11 January 2025 Change(s) in the name or address or contact details of a qualified person for pharmacovigilance (QPPV). Introduction of a summary of the PSMF or changes to the summary of the active substance.  14 Postrary 2023 Change in the re-test period of the active substance.  14 October 2020 Amendments to the Summary of Product Characteristics and to the product literature with regard to safe use of the product, where a collar is worn by the target species.  14 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.  14 November 2018 Change in the safety database of an existing pharmacovigilance system as described in the DDPS.  18 September 2018 Change in RMS from UK to AT.  19 Change in RMS from UK to AT.  10 Change in RMS from UK to AT.  10 Change in RMS from UK to AT.  11 Change in the address of the marketing authorisation holder from Bayer plc, Animal Health Division, Bayer House, Strawberry Hill, Newbury, Berkshire, RG2 6AD.  21 August 2018 Change in the address of the marketing authorisation holder from Bayer plc, Animal Healt		T.,
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02 February 2017 Change in pack size of the finished product.	31 March 2017	
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27 May 2016	Doloto Unidrug Distribution Croup Limited as a distributer
27 May 2016	Delete Unidrug Distribution Group Limited as a distributor.
18 May 2016	Changes to a DDPS following the assessment of the same DDPS in relation to another medicinal product of the same MAH
17 September 2015	Change in product name in Austria only.
16 April 2014	Change in MAH
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13 April 2012	Five changes to the test procedure for the active substance or starting material, reagent or intermediate used in the
	manufacturing process of the active substance.
	Two changes to the specification parameters/limits of an active
	substance, starting material, intermediate or reagent used in the
	manufacture of the active substance.
	Five changes to in process tests/limits applied during the
	manufacture of the active substance.
	Two changes in the manufacturing process of the active
45 Danamahan 0044	substance.
15 December 2011	Six changes in the specification parameters/limits of an active substance, starting material, intermediate or reagent used in the
	manufacture of the active substance.
	Change to in process test/limits applied during the manufacture
	of the active substance.
12 August 2011	Change to specification parameters/limits of an excipient.
02 March 2011	Change of distributor.
05 January 2011	Change in legal category from POM-V to NFA-VPS.
	Change to therapeutic indications regarding flea allergy
	dermatitis.
30 July 2010	Change to in process tests/limits applied during the manufacture
04 Marr 0040	of the finished product.
21 May 2010	Change in name/address of MAH.
02 April 2009	Update specification of an excipient to comply with Ph. Eur.
04 September 2008	Change of MAH.
03 September 2008	Change of MAH name/address.
02 April 2008	Change of MAH name/address.
09 January 2008	Renewal.
20 June 2006	Change in name of manufacturer of active substance.
23 November 2005	Variation to update the packaging to improve legibility and clarity
44.4 ".0005	on the product labelling and package leaflet.
14 April 2005	Change of shelf life from 3 to 5 years.
18 February 2005	Corrections/text changes to the SPC and Labels.
09 October 2003	Change in distributor.
14 March 2003	Renewal.
15 August 2002	Addition of an 'Indication'.
09 July 2002	Change in manufacturing process of the finished product.
19 February 2002	Repeat use procedure (UK as RMS).
22 May 2001	Addition of an FAD claim.
23 October 2000	Change of manufacturing process.
28 January 2000	Change of packaging.
22 March 1999	Change of shelf life from 2 to 3 years.
19 February 1999	Change of safety warnings.
08 August 1997	Mutual recognition procedure (UK as RMS).
19 March 1997	Update of licence particulars.
10 Maiori 1001	production of motion particulars.