

## **Post Authorisation Assessments**

## Capstar 57 mg Tablets for Large Dogs Vm 00879/3020

<ul> <li>25 January 2024 Deletion of a non-significant specification parameter in the specification parameters of an excipient.</li> <li>29 August 2023 One-off alignment of the product information with version 9.0 of the QRD templates.</li> <li>27 June 2023 Change in test procedure for the finished product: - Other changes to a test procedure.</li> <li>02 November 2021 Deletion of a non-significant parameter of an active substance.</li> <li>24 February 2021 Changes to the labelling and/or package leaflet.</li> <li>28 January 2021 Replacement to a test procedure for the finished product.</li> <li>09 September 2020 Changes to the labelling and/or package leaflet.</li> <li>28 January 2021 Replacement to a test procedure for the finished product.</li> <li>09 September 2020 Change in the address of the Marketing Authorisation Holder from Elanco Europe Ltd, LIIIy House, Priestley Road, Basingstoke, Hampshire, RG24 9NL, United Kingdom.</li> <li>16 March 2020 Minor change in the manufacturing process of the finished product. Addition of new tests and limits applied during the manufacture of the finished product.</li> <li>06 February 2020 Changes to the labelling and/or package leaflet.</li> <li>05 June 2019 Change in shape or dimensions of the container or closure (immediate packaging).</li> <li>19 March 2018 Change in RMS from UK to FR.</li> <li>16 August 2017 Changes to the labelling.</li> <li>07 March 2017 Introduction of a new pharmacovigilance system.</li> <li>15 February 2016 Change of MAH and Distributor, from Novartis Animal Health UK Limited to Elanco Europe Ltd.</li> <li>28 June 2016 Change in the name of a manufacturer of the finished product inter product literature.</li> <li>28 June 2016 Change in the name of the Marketing Authorisation Holder from Novartis Santé Animale to Elanco France in France only.</li> </ul>		05 1 000 1	
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		Authorisation Holder from Novartis Animal Health S.p.A
		to Elanco Italia S.p.A. in Italy only, and from Novartis
		Sanidad Animal, S.L. to Elanco Spain, S.L.U. in Spain
		only.
•	05 April 2016	Deletion of a non-significant in-process test applied
		during the manufacture of the finished product
		Extension of a re-test period of the active substance.
•	06 May 2014	To add/amend identification assays.
•	10 April 2014	Change in test procedures of the finished product and
		change of specification parameters for the finished
		product.
•	27 March 2014	Addition of pack size of 1 blister containing 1 tablet.
•	16 January 2014	Change of MA address in Portugal only.
•	10 October 2013	Change of manufacturer of the active substance.
•	04 July 2013	Change of MA holder address in France only.
•	28 March 2013	Change of MA holder address in Denmark, Finland,
		Norway and Sweden.
•	26 May 2011	Changes in the manufacturing process of the active
		substance
		Addition of a specification parameter of the active
	07.0 1 1 0000	substance
•	07 September 2009	Change to Part II of the dossier
•	25 March 2009	Change in specification of the finished product
•	28 August 2008	Change in test procedure performed on the finished product
•	02 April 2008	Approval of unseen mock ups
•	20 December 2007	Increase of bulk holding time
•	28 September 2007	Renewal
•	03 September 2007	Change of address of the MAH
•	28 March 2007	Change of storage conditions
•	29 March 2006	Change of manufacturer of the active substance
•	12 March 2003	Change of legal category from POM to GSL
•	06 September 2002	Mutual Recognition procedure, UK as RMS