

## **Post Authorisation Assessments**

## Atopica 10 mg Soft Capsules for Dogs Vm 00879/3011

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•	25 March 2024	Tightening of specification limits of an intermediate.
		Tightening of specification limits of an intermediate.
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•	23 June 2023	Pharmacovigilance sections updated to align product.
•	23 June 2023	One-off alignment of the product information with version 9.0* of the QRD template.
•	28 February 2023	Tightening of in-process limits of the active substance. Minor changes to a test method of the active substance. Addition of a new Ph.Eur from a new manufacturer for a non-sterile active substance.
•	05 December 2022	Introduction of a re-test period for a new active substance manufacturer source.
•	08 March 2022	Change in the name of a manufacturer used in the manufacture of the active substance.
•	08 July 2021	Change in the name and address of a manufacturer of the active substance.
•	09 June 2021	Change in the specification parameters and/or limits of
		an excipient.
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		an excipient.
•	25 September 2020	Change in the address of the MAH from Elanco Europe
		Ltd, Lilly House, Priestley Road, Basingstoke,
		Hampshire, RG24 9NL, United Kingdom to Elanco
		Europe Ltd, Form 2, Bartley Way, Bartley Wood,
	25 March 2020	Business Park, Hook, RG27 9XA, United Kingdom.
•	25 March 2020	Change in the specification parameters and/or limits of
		an active substance, used in the manufacturing process of the active substance.
		Removal of non-significant test parameter from the raw
		material specifications.
		Minor change to the restricted part of an Active
		Substance Master File.
•	30 August 2019	Addition of a new specification parameter with its
		corresponding test method of an active substance used
		in the manufacturing process of the active substance.
•	17 June 2019	Change in source of an excipient or reagent with TSE
		risk
•	05 June 2019	Change in the safety database of an existing
		pharmacovigilance system as described in the DDPS.
•	10 May 2019	Tightening of specification limits of an excipient
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•	07 March 2017	Introduction of a new pharmacovigilance system.
•	30 September 2016	Change in the name and address of the Marketing
		Authorisation Holder.
		Change of distributor details.
•	15 August 2016	Change in the name of a manufacturer of the finished
		product including manufacturer responsible for batch
	00 1.1. 0040	release.
•	06 July 2016	Change in the name and address of the Marketing
	25 November 2015	Authorisation Holder in Spain and Italy only. Widening of specification limits.
•		Changes in the manufacturing process of the active
		substance.
		Deletion of a manufacturer of the active substance.
		Tightening of specification limits.
		Addition of an identification test.
		Deletion of tests from raw material specifications.
•	31 July 2015	Grouped variation to change the name of the bulk
		manufacturer, replacement of sites for both batch release and Microbial tests and the deletion of both a
		manufacturing site and a packaging site.
•	07 October 2014	Update to the text in Section 4.5 and 4.9 on the SPC and
	• • • • • • • • • • • • • •	the relevant sections of the product literature.
•	07 February 2014	Change to the address of the MAH in Portugal only.
•	15 January 2014	Changes to test procedures for the active substance and
		intermediate used in the manufacturing process of the
		active substance.
		Changes to the specification limits of the active
		substance and of the intermediate used in the
	15 October 2013	manufacturing process of the active substance. Changes to the specification limits of the finished
•		product.
		Minor changes to the test procedures for the finished
		product.
•	05 July 2013	To change the address of the MAH in France only.
•	28 March 2013	To change the address of the MAH in Denmark, Finland,
		Norway and Sweden and corrections to the address of
	07 February 0010	the MAH in Cyprus.
•	27 February 2013	To update the SPC and product literature text and to change the ATCVet code.
•	11 July 2008	Change to the markings of the finished product.
•	29 May 2008	Renewal
	09 October 2007	To change the address of the marketing authorisation
		holder (MAH) and distributor.
•	20 July 2006	Addition of new packaging sites, to change the pack size
	-	of the finished product and to change the dimensions of
		the immediate packaging.
•	12 April 2006	Change in the composition of the product.
•	07 March 2006	To include an alternative quality control site, replacement
		of the site responsible for batch release and replacement
	26 Sontombor 2005	of a manufacturing site.
•	26 September 2005	Addition of a new test for the finished product.
•	24 May 2005	Changes to the finished product specification.
•	29 April 2005	To change the name of a manufacturer of the finished

		product.
•	06 April 2005	Changes to the manufacturing process, addition of a manufacturer and a change in the name of a manufacturer.
•	07 December 2004	Change in the name of a supplier.