

Post Authorisation Assessments

Kesium 50 mg / 12.5 mg Chewable Tablets for Cats and Dogs Vm 15052/4134

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•	12 June 2023	Change in pack size of the finished product: - Change in
		the number of units in a pack outside the range of the
	01 March 0000	currently approved pack sizes.
•	21 March 2023	Deletion of a non-significant in-process tests during the
	01 March 2022	manufacture of the finished product.
•	21 March 2023	Minor change in the manufacturing process of the
	18 October 2022	finished product. Change in the address of the MAH from Unit 3 Anglo
•		Office Park, White Lion Road Amersham,
		Buckinghamshire HP7 9FB to Explorer House, Mercury
		Park, Wycombe Lane, Wooburn Green, High Wycombe,
		Buckinghamshire, HP10 0HH, United Kingdom.
•	07 October 2021	Changes to the labelling and package leaflet.
•	10 September 2021	Minor changes to an approved test procedure of the
		finished product.
		Changes to a test procedure for the finished product.
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•	09 September 2021	Addition of a site where batch control/testing takes place.
		Addition of a primary packaging site of the finished
		product.
		Increase in batch size (from 533.000 tablets (100kg)
		1.000.000 tablets (187.5kg) to 533.000 tablets (100kg) / 1.000.000 tablets (187.5kg) Ceva site: 120/250/285/500
		kg of final blend bulk) of the finished product.
		Minor change in the manufacturing process of an
		immediate release solid oral dosage form or oral
		solutions.
		Tightening of in-process limit applied during the
		manufacture of the finished product.
		Addition of a manufacturing site of the finished product.
•	15 July 2021	Deletion of a non-significant specification parameter of
	-	the immediate packaging of the finished product.
•	06 July 2021	Deletion of a non-significant specification parameter of
		an excipient.
		Deletion of a non-significant specification parameter of

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		an excipient. Minor change to an approved test procedure for an
		excipient.
		Minor change to an approved test procedure for an
		excipient.
•	17 June 2021	Change in the invented name of the veterinary medicinal
		product from Kesium 62.5 mg Chewable Tablets for Cats
		and Dogs to Kesium 50 mg / 12.5 mg Chewable Tablets
		for Cats and Dogs.
•	03 March 2021	Submission of an updated Ph. Eur. certificate of
		suitability for an active substance from an already
	00 A 11 0000	approved manufacturer.
•	03 April 2020	Submission of an updated Ph. Eur. certificate of
		suitability for an active substance from an already
	24 1010 2019	approved manufacturer.
•	24 July 2018	Change in the invented name of the veterinary medicinal
•	12 June 2018	product in DK only. Change in the address of the manufacturer of the
-		finished product.
		Deletion of a manufacturing site responsible for batch
		release.
		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.
•	19 September 2017	Change in the QPPV of an existing pharmacovigilance
		system as described in the DDPS.
		Change of the back-up procedure of the QPPV of an existing pharmacovigilance system as described in the
		DDPS.
•	09 November 2016	Change in the name of a manufacturer of the finished
		product.
		Change in the name of a manufacturer of the finished
		product including manufacturer responsible for batch
		release.
		Change in the name of a manufacturer of the finished
		product including manufacturer responsible for batch
		release.
•	21 October 2016	Mock-ups approved.
		Change in distributor details from Alstoe Ltd to Ceva
	10 August 2016	Animal Health Ltd. Change of Marketing Authorisation Holder from Sogeval
•	19 August 2016	to Ceva Animal Health Ltd.
-	09 August 2016	Submission of an updated certificate of suitability.
	29 June 2016	
•		Introduction of a new pharmacovigilance system which has been assessed by the relevant national competent
		authority/EMA for another product of the same MAH.
-	08 June 2016	Renewal UK as CMS.
	16 February 2016	Submission of an updated Ph. Eur. certificate of
•	101 EDiudiy 2010	suitability.
		Deletion of a Ph. Eur. certificate of suitability
•	4 December 2015	Replacement of a secondary packaging site.
–	1 2000111001 2010	representation a secondary packaging site.

•	31 May 2013	Submission of an updated Ph. Eur. Certificate of Suitability for an active substance from two already approved manufacturers.
•	31 October 2012	To change the shelf-life of the veterinary medicinal product from 18 to 21 months.