

Post Authorisation Assessments

Clavucill 40 mg/10 mg, Tablets for Dogs and Cats Vm 19968/4003

•	15 July 2022	Updated certificate of suitability for an already approved manufacture.
		Updated certificate of suitability for an already approved
		manufacture.
		New certificate of suitability from a new manufacturer for
		an active substance.
		New certificate of suitability from a new manufacturer for
		an active substance.
•	27 December 2018	Changes to the labelling and package leaflet.
•	13 November 2018	Change in distributor details from: Chenelle Animal
		Heath Ltd., 7 Rodney Street, Liverpool, L1 9HZ, UK, to
		V.M.D. n.v., Hoge Mauw 900, 2370 Arendonk, Belgium.
•	02 October 2018	Change in shape or dimensions of the container or
	47.51 1 0040	closure (immediate packaging).
•	17 November 2016	Change in the safety database of an existing
	20 1010 2015	pharmacovigilance system as described in the DDPS.
•	30 July 2015	Introduction of a new pharmacovigilance system, which
•	03 June 2015	has not been assessed by the relevant NCA/EMA. Change in the name of the medicinal product from
•		Clavucill Tablets 50 mg to Clavucill 40 mg/10 mg,
		tablets for dogs and cats.
•	25 March 2015	Change to the MAH.
•	23 December 2014	Reduction of shelf-life of the finished product in the UK,
_		from 3 years to 2 years.
•	04 December 2014	Change in the specification limits of the finished product
		at both release and end of shelf-life.
•	05 August 2014	Harmonisation of SPC and product literature with
	-	Ireland
•	16 June 2014	Change in the shelf life of the finished product.
•	28 January 2014	Submission of updated Ph. Eur. Certificates of
		Suitability.
•	04 July 2013	Renewal procedure.
•	19 May 2009	To update the reference number of the certificate of
		suitability for Potassium Clavulanate.