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Post Authorisation Assessments

Vanguard CPV-L Vm 42058/4159

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•	25 March 2024	Change in the source of a starting material used in the
		manufacturing process of the active substance.
•	18 November 2022	Introduction of in vitro antigen capture ELISA potency
		test.
		Use of antigen capture ELISA to quantify antigen.
•	01 May 2020	Change in the address of the marketing authorisation
	-	holder from Zoetis UK Limited, 5th Floor, 6 St. Andrew
		Street, London, EC4A 3AE to Zoetis UK Limited, 1st
		Floor, Birchwood Building, Springfield Drive,
		Leatherhead, Surrey, KT22 7LP.
•	19 October 2018	Change in the specification parameters and/or limits of
		the finished product.
•	09 November 2016	Change in the composition (excipients) of the finished
		product.
•	19 February 2014	Transfer of MA from Pfizer Ltd to Zoetis UK Limited and
	_	change of distributor. Change of name of the active
		substance manufacturer and change of name of the
		finished product manufacturer responsible for batch
		release.
•	16 December 2010	Renewal
•	18 November 2010	Removal of a test.
•	07 July 2010	Reduction of onset of immunity of an active substance.
•	17 February 2010	Addition of suppliers of starting materials.
•	15 July 2009	Changes to Section 4.2 of the SPC.
•	24 May 2006	Changes to the SPC and product literature to bring them
		into line with new legislation.
•	24 May 2006	Change of legal category from POM to POM-V.
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