

Post Authorisation Assessments

Enrox Max 100 mg/ml Solution for Injection for Cattle and Pigs Vm 01656/5066

•	21 March 2024	One-off alignment of the product information with version 9.0 of the QRD templates.
•	24 July 2023	Submission of a new Ph. Eur. CEP from a new
		manufacturer for a non-sterile: – active substance.
•	24 May 2023	Change to comply with an update of the relevant monograph of the Ph. Eur.
		Change to comply with Ph. Eur. by removing reference to
		the internal test method and test method number.
		Changes to the quality part of the dossier: Deletion of - a
		Ph. Eur. CEP for an active substance.
•	20 January 2023	Change to comply with an update of the relevant
		monograph of the Ph. Eur.
		Change to comply with Ph. Eur. by removing reference to
		the internal test method and test method number.
		Deletion of Ph.Eur CEP for an active substance.
•	02 March 2020	Addition of a site where batch control/testing takes place.
		Submission of an updated Ph. Eur. certificate of
		suitability for an active substance from an already
		approved manufacturer.
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		suitability for an active substance from an already
		approved manufacturer.
•	26 July 2019	Update of the test procedure to comply with the updated
		general Ph. Eur monograph.
•	24 December 2018	Changes to an existing pharmacovigilance system as
		described in the DDPS.
•	17 July 2018	Renewal – UK CMS.
•	02 December 2015	Minor changes to an approved test procedure
•	29 April 2015	To add UK local representative information to package
		leaflet.
•	01 April 2015	Submission of an updated and new certificate of
		suitability.
•	09 December 2013	Introduction of a new pharmacovigilance system.
•	09 December 2013	Change of invented name of the product from 'Enrox
		Max 100 mg/ml Solution for Injection' to 'Enroxal Max
		100 mg/ml Solution for Injection' in Germany only.
•	28 November 2013	Change of MA holder from Billev Pharma aps to Krka Dd.
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