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Post Authorisation Assessments

Tricaine Pharmaq 1000 mg/g Powder for Solution for Fish Treatment Vm 11003/4014

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30 June 2022	Replacement to a test procedure for the active substance.
	Changes in the specification parameters of the active
	substance.
09 January 2020	Change in the specification parameters and/or limits of
	an active substance.
	Change in the specification limits of the finished product.
11 December 2019	Addition of a site where batch control/testing takes place.
26 November 2018	Change in RMS from UK to NO.
21 November 2018	Changes to the labelling and package leaflet.
06 April 2018	Renewal UK as RMS
28 July 2017	Change in type of container for the finished product.
20 July 2016	Minor change to an approved test procedure.
	Change in name of manufacturer of the active
	substance.
	Additional site of batch control/testing.
02 June 2015	Update to the DDPS.
15 August 2014	Grouped variation: To replace the HPLC test procedure
	for assay and related substances, API and finished
	product.
	To remove secondary test procedures for the API and/or
	the finished product.
	To implement other updates to the ASMF.
09 January 2014	To add an alternative batch release site.
	11 December 2019 26 November 2018 21 November 2018 06 April 2018 28 July 2017 20 July 2016 02 June 2015 15 August 2014