



Post Authorisation Assessments

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

•	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.