CADTH

Common Drug Review

Submission Status

Product: Aloxi	
Generic Name: palonosetro	

Manufacturer/Applicant: Eisai Limited

Indication(s): Nausea and vomiting (chemotherapy induced) prevention

Submission Type: Resubmission Date Submission Received: 2015-Feb-06

Date NOC Issued: 2012-Mar-14 Application Fee Schedule¹:

N/A

Submission/resubmission accepted for review 2015-Feb-23 2015-Feb-23 2015-Feb-23 - Review has been initiated 2015-Feb-24 - Call for patient input posted on 2015-Jan-06 - Patient group input submission received 2015-Feb-25 Patient group input submission received 2015-Mer-10 2015-Mer-10 2015-Mer-10 2015-Mer-10 2015-Mer-10 2015-Mer-10 2015-Mer-10 2015-Mer-10 - Patient group input summary sent for review on 2015-Mer-10 - Patient group input summary sent for review on 2015-Mer-10 - Patient group input summary sent for review on 2015-Mer-10 - Patient group input summary sent for review on 2015-Mer-10 - Patient group input summary sent for review on 2015-Mer-10 - Patient group input summary sent for review on 2015-Mer-10 - Patient group summary sent for review on 2015-Mer-10 - Patient group summary sent for review on 2015-Mer-10 - Patient group summary sent for review on 2015-Mer-10 - Patient group summary sent for review on 2015-Mer-10 - Patient group summary sent for review on 2015-Mer-10 - Patient group summary sent for review on 2015-Mer-10 - Patient group summary sent for review on 2015-Mer-10 - Patient group summary sent for review on 2015-Mer-10 - Patient group summary sent for review on 2015-Mer-10 - Patient group summary sent for review on 2015-Mer-10 - Patient group summary sent for review on 2015-Mer-10 - Patient group summary sent for review on 2015-Mer-10 - Patient group summary sent for review on 2015-Mer-10 - Patient group summary sent for review on 2015-Mer-10 - Patient group summary sent for review report(s) - 2015-Mer-20 - New target date: 2015-Jul-24 - New target date: 2015-Aug-10 - Voluntarily withdrawn by the manufacturer on 2015-Jul-30 - New target date: 2015-Aug-10 - Voluntarily withdrawn by the manufacturer on 2015-Jul-30 - New target date: 2015-Aug-10 - New target date: 2015-Aug-10 - New tar	Original Targeted CDEC Meeting:	2015-Jul-15		Priority Review Status: No	t Requested
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Comments from manufacturer on draft CDR review report(s) 2015-May-20 2015-May-20 2015-May-20 2015-May-20 2015-May-20 2015-May-27 2015-May-28 2015-May-	Patient group input summary comments received	2015-Mar-10	2015-Mar-10	- Patient input summary feedback deadline: 2015-	
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¹ Refer to Appendix 1 of the Procedure for the CADTH Common Drug Reviewin the Common Drug Review section of www.cadth.ca for details regarding fee schedules.

² Please refer to the *Procedure for the CADTH Common Drug Review* in the Common Drug Review section of www.cadth.ca for complete details regarding the CDR process and targeted time frames for key milestones,.

³ The call for patient group input is posted 20 business days inadvance of the manufacturer's anticipated date of filing the application. Patient groups have a total of 35 business days for prepating and submitting patient input.

⁴The embargoed CDEC recommendation is held in confidence by all stakeholders and not acted upon until after CADTH has issued the notice of CDEC Final Recommendation.

⁵ The timing for posting the CDEC Final Recommendation and CDR review report(s) depends on several factors including the need for consultation with the manufacturer regarding redaction issues.