Common Drug Review ¹					
CADTU		ubmission Status			
CADTH Product:	Eylea				
Generic Name: aflibercept					
Manufacturer/Applicant: Bayer Inc.					
Indication(s): Macular edema secondary to central retinal vein occlusion					
Submission Type:	New Indication	Date NOC Issued: 2014-Oct-16			
Date Submission Received:	2014-Oct-17	Application Fee Schedule ¹ :		Schedule B	
Orginal Targeted CDEC Meeting:	2015-Mar-18	Priority Review Status: Not Red		Not Requested	
Phase	Target Time (Business Days)	Target Date ²	Actual Date	Comments	
Submission/resubmission accepted for review	10	2014-Oct-31	2014-Oct-31	 Submission placed in queue in accordance with CDR procedures. Review to be initiated pending availability of resources and target dates will be updated. Review has been initiated 2014-Nov-28 	
Patient group input submission received ³		2014-Oct-08	2014-Oct-08	 Call for patient input posted on 2014-Aug-19 Patient group input deadline: 2014-Oct-08 Patient input submission received 	
Patient group input summary comments received	5	2014-Dec-11	2014-Dec-11	 Patient input summary sent for review on 2014-Dec-04 Patient input summary feedback deadline: 2014-Dec-11 Patient input summary feedback received 	
CDR review reports sent to manufacturer ⁴	45	2015-Jan-22	2015-Feb-24	- New target date: 2015-Feb-19 - New target date: 2015-Feb-24	
Comments from manufacturer on CDR review reports received by CADTH	7	2015-Feb-02	2015-Mar-05	- New target date: 2015-Mar-02 - New target date: 2015-Mar-05	
Redaction response from manufacturer on CDR review reports received by CADTH	5	2015-Feb-09	2015-Mar-12	- New target date: 2015-Mar-09 - New target date: 2015-Mar-12	
CDEC meeting		2015-Mar-18	2015-Apr-08	 New target date: 2015-Apr-08 The April 2015 Canadian Drug Expert Committee (CDEC) meeting will be held on April 8, to accommodate for the 2015 CADTH Symposium from April 12 to 14, 2015, in Saskatoon, SK. 	
CDEC recommendation & redacted CDR review reports sent to drug plans and manufacturer	5 to 7	2015-Mar-25	2015-Apr-16	- New target date: 2015-Apr-15 - New target date: 2015-Apr-16	
Embargo period and validation of redacted CDR review reports					
Manufacturers may make a request for reconsideration and drug plans may make a request for clarification of the recommendation	10	2015-Apr-09	2015-Apr-30	- New target date: 2015-Apr-29 - New target date: 2015-Apr-30	
Final recommendation sent to drug plans and manufacturer (No requests for clarification are made AND no request for reconsideration is made or request for reconsideration is resolved)	5	2015-Apr-16	2015-May-07	- New target date: 2015-May-07 - Notice of final recommendation issued	
CDEC final recommendation posted ⁶	Variable	2015-May-11	2015-May-11		
Final CDR review reports and patient input posted ⁷	Variable	2016-Jun-16	2016-Jun-16		

¹ Refer to appendix 1, section 2.2.1 of the Procedure for the CADTH Common Drug Review (August 2014), in the Common Drug Review section of www.cadth.ca for more details.

² The target dates for this report are based on the targeted CDEC meeting schedule, which is posted on www.cadth.ca.

³ The deadline for patient group input is 15 business days after CADTH receives the submission or up to 35 business days if advance notice (20 business days maximum) of a submission is received from the manufacturer.

⁴ Target time is calculated, based on the date the reviewers receive copies of the manufacturer's submission. Target time does not include the time allocated for receipt of manufacturer's additional electronic copies (5 business days) and time allocated for distribution of electronic copies to reviewers (3 business days).

⁵ The 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.

⁶ The target date for posting the CDEC Final Recommendation depends on several factors including the need for consultation with the manufacturer regarding redaction issues.

⁷ The timing of the posting of the CDR review report(s) depends on several factors, including the need for consultation with the manufacturer in case of disagreement with regard to redactions made.

Refer to the Procedure for the CADTH Common Drug Review in the Common Drug Review section of www.cadth.ca for more details about the CDR process.

This CDR Project Status Report is posted every other week, reflecting status as of the end of day Wednesday (4:00 pm ET) of that week.