Canadian Agency for Drugs and Technologies in Health Product: Olmetec Plus Generic Name: olmesartan medoxomil + hydrochlorothiazide					
Manufacturer: Schering-Plough Canada Inc.					
Submission Type: New					
	Date Submission Received:	2008-Nov-28	Da	te NOC Issued:	2008-Oct-28
	Targeted CEDAC Meeting:	2009-Apr-15	Priority R	eview Granted:	Not Requested
	Phase	Target Time (Business Days)	Target Date**	Actual CDR Date	Comments
	Submission Assessment	5	2008-Dec-05	2008-Dec-05	Category 1 Submission requirements deemed incomplete December 5, 2008.
1	Submission deemed complete			2008-Dec-08	Submission deemed complete.
2	CDR Reviewers' Reports Completed • Reviewers selected and contracted • Literature search and selection completed • Systematic review of clinical data completed • Critical appraisal of pharmacoeconomic (PE) data completed • Clinical and PE reports written • Reports edited and finalized • Reviewers' reports sent to manufacturer	45	2009-Mar-02		Additional information requested December 9, 2008. Additional information received December 10, 2008. Additional information requested January 12, 2009. Additional information requested January 20, 2009. Additional information received January 23, 2009. Additional information received January 28, 2009.
3	Comments from Manufacturer on Reviewers' Reports Received by CDR	7	2009-Mar-11	2009-Mar-12	
4	Reviewers' Reply to Manufacturer's Comments Completed	7	2009-Mar-20	2009-Mar-23	Due date for reviewers' reply March 23, 2009.
5	CEDAC Brief Completed and Sent to CEDAC Members	5	2009-Mar-31	2009-Mar-31	
6	CEDAC Meeting		2009-Apr-15	2009-Apr-15	
7	CEDAC Recommendation and Reasons for Recommendation Sent to Drug Plans, ACP and Manufacturer	5	2009-Apr-22	2009-Apr-22	
8	Embargo Period*** Manufacturers may make a Request for Reconsideration and Drug Plans may make a Request for Clarification of the Recommendation and Reasons for Recommendation	10	2009-May-06	2009-May-06	Request for Clarification received May 6, 2009.
9 (a)	Final Recommendation sent to Drug Plans, ACP, and Manufacturer (No Requests for Clarification are made AND no Request for Reconsideration is made or Request for Reconsideration is Resolved)	5			
	OR				
9 (b)	Clarification and Final Recommendation sent to Drug Plans, ACP, and Manufacturer (Clarification Requested, no Request for Reconsideration made)	5	2009-May-27	2009-May-27	Clarification and Notice of Final Recommendation issued.
	OR				· · · · · · · · · · · · · · · · · · ·
	Placed on CEDAC Agenda For Reconsideration (At Manufacturer's request)	25 Depends on Meeting Dates			
10	Final Recommendation sent to Drug Plans, ACP, and Manufacturer o the Procedure for Common Drug Review on the Common D	5			

* Refer to the Procedure for Common Drug Review on the Common Drug Review section of www.cadth.ca for more details. ** The CDR review process is initiated AFTER a submission is deemed complete. The target dates for this report are based on the CEDAC meeting schedule, which is posted on

www.cadth.ca *** The Recommendation and Reasons for Recommendation are to be held in confidence and not acted upon until after the CDR Directorate has issued the notice of Final Recommendation.

Reflects updates as of Thursday noon.