

PRIOR AUTHORIZATION REQUEST FORM

Asciminib (Scemblix®) / Chronic myeloid leukemia (CML) in chronic phase for adult

DECLARATION OF THE INSURED PERSON

Section 1: Information about the p	lan member and the	patient						
Name of plan member	Insurance policy / certificate		Name of employer					
Name of patient	Date of birth (YYYY/MM/DD)		Telephone					
Address (house number and street name)	City/Town		Province	Postal code				
Section 2: Other prescription drug insurance policies								
Do you have other prescription drug insura			☐ Yes	□ No				
If so, please answer the following:								
What type of plan is it?		☐ Private	☐ Public					
Have you ever submitted a claim for this d	rug to the other insurer?		☐ Yes	□ No				
What is the status of the claim?		☐ Accepted	l □ Refused	☐ Under review				
Did this insurer ask you to complete a prio	r authorization request?		☐ Yes	□ No				
If so, what is the status of the prior au	status of the prior authorization request?			☐ Under review				
Please enclose acceptance or refu	sal documents, if app	licable						
Section 3: Authorization to disclose	<u> </u>							
I certify that the information in this prior authorization request is complete, accurate and true.								
I authorize physicians and other health care professionals, medical, paramedical or clinical institutions, care coordinators, members of SSQ's Preferred Pharmacy Network (outside Quebec only) and any public or parapublic organization, including Régie de l'assurance maladie du Québec, to disclose to SSQ, Life Insurance Company Inc. (SSQ) any of my relevant information including and without limitation, any medical information, and medical evaluations in connection with the processing of this request. I hereby waive their confidentiality obligation and authorize them to disclose the requested information to SSQ. In addition, I authorize SSQ to disclose to the previously named third parties any of my relevant personal information including and without limitation any medical information and medical evaluations in connection with the processing of this request. Photocopies of this document have the same value as the original.								
Signature of patient (parent/legal	guardian)		Dat	e				
INADORTANT .								
IMPORTANT:								
All correspondence concerning this form will be sent to the address indicated in the plan member's file.								
Send us this duly completed form by mail or by fax to: 1-855-453-3942.								
Telephone: 418-651-2588/1-800-380-2588 – Fax: 1-855-453-3942 Address: 2525 Laurier Blvd, P.O. Box 10500, Quebec City, QC G1V 4H6								
ssq.ca								



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DECLARATION OF THE PRESCRIBER

Section 4: Information about the prescriber								
Name of prescriber	Name of prescriber		Specialty		Licence No.:			
				1				
Telephone				Fax				
I hereby certify that the information in this request is complete, true, and accurate:								
Signature of prescriber				Date				
Section 5 : Drug covered								
Drug name	Pharmaceutical form	Stren	gth	Dosage				
Scemblix					 of administration:			
				riequency o	i dullillisti ation.			
Type of request	☐ First request			☐ Continua	ation of treatment			
	Complete section 6			Complete section 7				
				•	e section 6 if this is the first requested from SSQ			
IMPORTANT:								
Please do not provide a	Please do not provide any genetic test results							
Costing C. Clinical information (final manual)								
Section 6 : Clinical information (first request)								
Diagnosis								
Chronic myeloid leukemia in chronic phase as per Health Canada approved indication								
For information purposes: According to Heath Canada, Asciminib is indicated for the treatment of adult patients with Philadelphia chromosome-positive chronic myeloid chronic (CML) in chronic phase (PC) previously treated with two or more tyrosine kinase inhibitors.								
☐ Other. Specify:								



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Section 6 : Clinical information (first request) (cont'd)								
Summary of previous trials or contraindications								
Medication of other medical treatment	Results	Treatment period (if applicable)						
Category Name: Dosage:	☐ Ineffectiveness☐ Intolerance☐ Contraindication☐ Other. Specify	from						
Category Name :	☐ Ineffectiveness ☐ Intolerance	from						
Dosage :	Contraindication Other. Specify	to						
Category Name :	☐ Ineffectiveness ☐ Intolerance	from						
Dosage :	Contraindication Other. Specify	to						
Hematologic response Yes Elements of observed hematological response: No Expected clinical benefit with continuation of treatment:								
Section 8 : Additional information								