

Prior Authorization Request Form

Adalimumab (Abrilada®, Amgevita®, Hadlima®, Hulio®, Humira®, Hyrimoz®, Idacio®, Simlandi®, Yuflima®), infliximab (Avsola®, Inflectra®, Remicade®, Renflexis®), risankizumab (Skyrizi®), ustekinumab (Stelara®), vedolizumab (Entyvio®) / Moderate to severe Crohn's disease

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Section 1: Information about the plan member and the p	patient			
Name of also mombar	Policy Certifica	Name o	familiar	
Name of plan member	Policy Certifica	ate iname o	f employer	
Name of patient	Date of birth	Telephone		
Name of paroni	Date of birti	Tolophono		
Address (number and street name)	Town/City		Province	Postal code
Section 2: Other prescription drug insurance policies				
Do you have other prescription drug insurance?		□Yes	□No	
If so, please answer the following:				
What type of plan is it?		☐ Private	☐ Public	
Have you ever submitted a claim FOR THIS DRUG to the other insure	r?	□Yes	□No	
What is the status of the claim?		☐ Accepted	Refused	☐ Under review
Did this insurer ask you to complete a prior authorization request?		□Yes	□No	
If so, what is the status of the prior authorization request?		☐ Accepted	Refused	☐ Under review
Please enclose acceptance or refusal documents, if applicable		·		
Section 3: Authorization to disclose personal information	on			
I certify that the information in this prior authorization request is comple				
I authorize physicians and other health care professionals, medical, p Quebec only) and any public or parapublic organization, including Régie personal information including and without limitation, any medical in confidentiality obligation and authorize them to disclose the requested in personal information including and without limitation, any medical information.	paramedical or clinical institutions de la l'assurance maladie du Quinformation and medical evalunformation to SSQ. In addition,	uébec, to disclose to SSQ, Li lations in connection with the lauthorize SSQ to disclose t	ife Insurance Company he processing of this r to the previously named	Inc. (SSQ) any of my relevant equest. I hereby waive their
Photocopies of this document have the same value as the original.				
Signature of patient (parent/legal guardian)		 Date		
IMPORTANT: All correspondence concerning this form will be sent to the	address indicated in the r	particinant's file		
		varticipant 3 nie.		
Send us this duly completed form by mail or by fax to: 1-855-453	3-3942.			
Telephone: 418-651-2588/1-800-380-2588 – Fax: 1-855-453-3942				

DECLARATION OF THE PR	LOCKIDEN						
Section 4: Information al	bout the pres	criber					
Name of prescriber	ame of prescriber Specialty			License no.			
Telephone		Fax					
I hereby certify that the informa	tion in this reque	est is complete, true a	nd accurate.				
Signature of prescriber Date							
Section 5: Drug covered by the authorization							
Drug name Pharma		Pharmace	ceutical form Strength			Dosage	
						Dose: Frequency of administration:	
Type of request	☐ First reques	n 6 Comple	tinuation of treatment te section 7 to proplete section 6 if this is	the first authorization requested fror	n SSQ		
Also, complete section 6 if this is the first authorization requested from SSQ Injection – administered at: Home Outpatient clinic CHSLD Doctor's office Hospital (patient is admitted) Other. Specify:							
Exact location's name and a	ddress						
IMPORTANT: To ensure sound managem to certain conditions.	ent of its group	o insurance plans, S	SQ gives preference	to the use of biosimilar drugs	s. Eligibility fo	or reference biologic products is subject	
IMPORTANT: Please do not provide gene	tic test results.						
Section 6: Clinical inform Diagnosis Moderate or severe Cro Other, specify: Clinical information Patient's weight:	nation (first	request)					
Summary of previous trials or contraindications							
	medical treatm	nent	Reason	or discontinuation		Duration of treatment	
Corticosteroids Name:			Intolerance Contraindication				
Azathioprine Dose:			Intolerance Contraindication				
Mercaptopurine Dose:			Intolerance Contraindication				

Section 6: Clinical information (first request) (con	tinue)	
Summary of previous trials or contraindications		
Drug or other medical treatment	Reason for discontinuation	Duration of treatment
Methotrexate	☐ Ineffectiveness	From
Dose:	☐ Intolerance ☐ Contraindication	From To
	Other, specify :	
Anti-TNF ⁽¹⁾	☐ Absence of clinical benefit despite an induction	
Name:	treatment Loss of clinical benefit	From
Dose:	Intolerance, specify: Contraindication, specify:	То
Anti-TNF ⁽²⁾	Absence of clinical benefit despite an induction treatment	
Name:	Loss of clinical benefit	From
Dose:	Intolerance, specify: Contraindication, specify:	То
Other	☐ Ineffectiveness	
Name:	☐ Intolerance ☐ Contraindication	From
Dose:	Other, specify:	To
Information necessary to evaluate the response to treatm The drug covered by the present authorization request was fir Patient's weight: kg Positive effects observed:		
0.45.40.41.85.41.45.44.45.4		
Section 8: Additional information		