Request for Biologics for Psoriatic Arthritis (PsA)/Seronegative Arthritis Exceptional Access Program (EAP)



To avoid delays, please ensure that all appropriate information for each section is provided

Not for Other Inflammatory Disorders

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Section 1	– Physician	Informa	tion			Secti	on 2 – P	atient Inf	ormation				
		Initial	Last Name	Last Name		First Na	First Name		Initial	Last Name	Last Name		
Street #	Street Name					Ontario	Ontario Health Insurance Number						
City			Postal Code			Gender Curre Male Female				rent Weight (kg)			
Fax			Telephone			Date of	Date of Birth (DD/MM/YYYY)						
Request Type	New Reques	st (complete al	Il sections)	Rene	ewal Request (d	complete se	ctions 3, 4B)	EAP#					
Section 3	- Products	(attach add	litional shee	ets if mo	re space is	required)							
adalimumab (Humira®)			40 mg SC every two weeks								Dosage		
certolizumab (Cimzia™)			400 mg SC at 0, 2 and 4 weeks followed by 0 OR 400 mg every 4 weeks				maintenance therapy of 200 mg every 2 weeks						
etanercept (Enbrel®)			25 mg SC twice weekly or 50 mg SC once we				reekly				Dosing Frequency		
golimumab (Simponi®)			50 mg SC once monthly										
infliximab (Remicade®)			maintenance therapy¹ of 3-5 mg/kg/dose IV ev				every 8 weeks				D. C. (Advisidadis		
¹Requests for Remicade in patients with PsA who ir established renewal criteria. Note that Inflectra (LU			nitiated Remicade therapy on or prior to February 24, 2 code 470) and Renflexis (LU code 543) are considered								Route of Administrations SC IV PO		
For patients with P with coexistent mo	mab (Taltz [®]) sA and coexistent mild p derate-to severe plaque ng: 80 mg/1.0 ml SC, 16	plaque psoriasis psoriasis (PPs)	. To be used as m , refer to ODB for	onotherapy mulary for a	ccess upon meeti	n with a conve ng the Limited	entional DMARD I Use criteria for	(i.e. MTX). For P PPs; EAP authori					
secukir	numab (Cosent	yx®) 150	mg SC at wee	eks 0, 1, 2	and 3 followed	by monthly	/ maintenance	dosing starting	g at week 4				
patients with coexis	nti-TNF alpha inadequate stent moderate to sever by monthly maintenance	e plaque psorias	sis, use the dosing										
upadac	i tinib (Rinvoq™	⁴) 15	mg PO once da	aily									
Section 4A Indication of Active Disease			Section 4B Response to Treatment										
Diagnosis of active PsA			Renewal requests should demonstrate a 20% reduction in swollen joint count and a minimum of improvement in 2 swollen joints over the										
≥ 5 swollen joints		prev	previous year. For renewals beyond the seco				bjective evide	ence of the pres	servation of trea	itment effect mi	ust be provided	l.	
AND Diagnostic imaging evidence of PsA (x-rays, U/S, MRI)		٠,٠	Clinical Marker	Prior-to Requested Biologic		Rene	Renewal 1 F		2 R	enewal 3	Renew	al 4	
if < 5 swollen joints, provide location of swollen joints If not PsA, please specify diagnosis and enclose copies of relevant diagnostic imaging and bloodwork		Sı	wollen Joint Count		5.09.0								
		odwork	Date D/MM/YYYY)										
Section 5	- Previous/0	Current l	Disease I	Modify	/ing Anti-	Rheum	atic Dru	g (DMAR	D) Thera	ру			
3 months. If patie	f use and response ent has documented ntolerances must be	contraindicat											
NAME OF DMARD			START DA		END DATE (DD/MM/YYYY)				CONTINUATION ure at maximum dose must be provided				
methotrexate													
leflunomide													
sulfasalazine													
Physician Signature (Mandatory)						CPSO N	CPSO Number				Date (DD/MM/YYYY)		
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