

Request for Biologics for Rheumatoid Arthritis (RA) Exceptional Access Program (EAP)

Not for Other Inflammatory Disorders
Not for Paediatric Cases



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

Section 1 - Physician Information			Section 2 - Patient Information		
First Name	Initial	Last Name	First Name	Initial	Last Name
Street #	Street Name		Ontario Health Insurance Number		
City		Postal Code	Gender	<input type="checkbox"/> Male <input type="checkbox"/> Female	Current Weight (kg)
Fax		Telephone (Back Line)	Date of Birth (DD/MM/YYYY)		

Request Type Initial Request (Complete all sections) Renewal Request (Complete sections 3, 4B) EAP #

Section 3 - Drugs, Dose and Regimen Requested *(attach additional sheets if more space is required)*

<input type="checkbox"/> abatacept (Orencia™)	500 mg (<60 kg), 750 mg (60-100 kg), 1000 mg (>100 kg) IV at 0, 2, 4 weeks then every 4 weeks or 125 mg SC once weekly +/- initial IV loading	Dosage
<input type="checkbox"/> adalimumab (Humira®)*	40 mg SC every two weeks	
<input type="checkbox"/> anakinra (Kineret®)*	100 mg SC per day	Dosing Frequency
<input type="checkbox"/> certolizumab (Cimzia™)	400 mg SC at 0, 2, 4 weeks, followed by 200 mg SC every other week (400 mg SC every 4 weeks may be considered for maintenance therapy only)	
<input type="checkbox"/> etanercept (Enbrel®)*	25 mg SC twice weekly or 50 mg SC once weekly.	Route of Administration: <input type="checkbox"/> SC <input type="checkbox"/> IV <input type="checkbox"/> PO
Requests for renewal of Enbrel in patients with RA who initiated therapy prior to July 31, 2017 will be assessed according to established renewal criteria. New requests will not be accepted. Note that Brenzys (LU code 499) and Erelzi (LU code 512) are considered for RA in patients meeting LU criteria.		
<input type="checkbox"/> golimumab (Simponi®)	50 mg SC once monthly	
<input type="checkbox"/> infliximab (Remicade®)*	Maintenance therapy of 3 mg/kg IV every 8 weeks.	
Requests for renewal of Remicade in patients with RA who initiated therapy prior to February 24, 2016 will be assessed according to established renewal criteria. New requests will not be accepted. Note that Inflectra (LU code 468) and Renflexis (LU code 541) are considered for patients with RA meeting LU criteria.		
<input type="checkbox"/> tocilizumab (Actemra®)	4 mg/kg/dose IV every 4 weeks followed by an increase to 8 mg/kg/dose IV based on clinical response; not to exceed 800 mg/dose	
Patients < 100 kg weight, starting dose of 162 mg SC every other week. May increase to weekly dose based on clinical response > 100 kg weight, 162 mg every week		
tofacitinib (Xeljanz™)	Please refer to LU code #480	
rituximab (Rituxan®)	Please use the RA Rituxan EAP form	

*Higher than listed dosages should be accompanied by the rationale for the need for non-standard dosing levels. Please provide additional documentation (i.e. objective evidence) regarding the patient's response, including the swollen joint count, to the standard dosing regimen vs the higher dosing regimen.

Section 4A Indication of Active Disease	Section 4B Response to Treatment					
Diagnosis of active RA: <input type="checkbox"/> ≥ 5 Swollen Joints <input type="checkbox"/> Rheumatoid Factor Positive and/or Anti-CCP Positive AND / OR <input type="checkbox"/> Radiographic Evidence of RA <i>if < 5 swollen joints, provide location of swollen joints</i>	Renewal requests should demonstrate a 20% reduction in swollen joint count and a minimum of improvement in 2 swollen joints over the previous year. For renewals beyond the second year, objective evidence of the preservation of treatment effect must be provided.					
	Clinical Marker	Prior-to Requested Biologic	Renewal 1	Renewal 2	Renewal 3	Renewal 4
	Swollen Joint Count					
	Date (DD/MM/YYYY)					

Section 5 - Previous/Current Disease Modifying Anti-Rheumatic Drug (DMARD) and Biologic Therapies

- Provide details of use and response to separate courses of maximum dose methotrexate (i.e. 20 mg/week) AND maximum dose leflunomide (i.e. 20 mg/day), AND any DMARD combination for at least 3 months each. If maximum doses of methotrexate and leflunomide have not been tried, details of the patient's intolerance or contraindication to these two agents must be given AND details of another DMARD tried in each one's place for at least 3 months should be provided. **OR**
- Provide details of use and response to methotrexate 20 mg/week and leflunomide 20 mg/day in combination for at least 3 months. **OR**
- Provide details of use and response to triple disease modifying anti-rheumatic drug (DMARD) therapy, defined as methotrexate (20 mg/week), sulfasalazine (2 g/day) and hydroxychloroquine (400 mg/day - dose based by weight up to 400 mg per day) for at least 3 months. Note: In cases where the patient could not receive an adequate trial of triple DMARD therapy with methotrexate, sulfasalazine and hydroxychloroquine due to intolerance, then the regular DMARD trial criteria must be met. (i.e. #1 or #2)
- Combination biologics is not funded.

NAME OF DRUG	DOSING REGIMEN	START DATE (DD/MM/YYYY)	END DATE (DD/MM/YYYY)	REASON FOR DISCONTINUATION <small>Details of intolerance, contraindication, or failure at maximum dose must be provided</small>
methotrexate				
leflunomide				

Physician Signature (Mandatory)	CPSO Number	Date (DD/MM/YYYY)
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