

Please indicate: Start of treatment: Start date _

MEDICARE FORM

1

Actemra® (tocilizumab) Injectable **Medication Precertification Request**

Page 1 of 3

(All fields must be completed and legible for precertification review.)

1

Virginia (HMO D-SNP) FAX: 1-833-280-5224 **PHONE:** 1-855-463-0933

For other lines of business: Please use other form.

Note: Actemra is non-preferred. Preferred products may vary based on indication. See section G below.

	Continuation of therapy: Da	ate of last treatment/	/					
Precertification	Requested By:		Phone:		Fax:			
A. PATIENT IN								
First Name:		Last Name:			DOB:			
Address:			City:		State:	ZIP:		
Home Phone:	Work F	Phone:	Cell Phone:		Email:			
Current Weight:	lbs_orkgs	Height: inches or	cms	Allergies:				
		····g····						
		Doos patient have at	hor oovorago?					
	ID #:		Does patient have other coverage? Yes No If yes, provide ID#: Carrier Name:					
Insured:			If yes, provide ID#: Carrier Name: Insured:					
		insured.						
First Name:		Last Name:		(Check One):	□ M.D. □ D.	0. 🗌 N.P. 🗌 P.A.		
Address:			City:	(0.000.010)	State:	ZIP:		
Phone:	Fax:	St Lic #:	NPI #:	DEA #:	- L	JPIN:		
Provider Email:		Office Contact Name:		Phone:				
	G PROVIDER/ADMINISTRATIO							
Place of Admin			Dispensing Pro	vider/Pharmacy:				
☐ Self-administe		ce		Dispensing Provider/Pharmacy:				
<u> </u>	— ,		Specialty Pharmacy Mail Order					
	Name:							
	n Center Phone:							
	N							
Administration	n code(s) (CPT):		Address:					
Address:			City:	5	State:	ZIP:		
City:	State:	ZIP:	Phone:		Fax:			
Phone:	Fax:		TIN:	PIN:				
TIN:	TIN: PIN:			NPI:				
NPI:			E. PRODUCT I					
Please explain i inject the reque	f there are any medical reason(s) why the patient cannot self-		Actemra (toci	lizumab) IV			
inject the reque	sted drug.			Actemra (tocilizumab) SC				
			HCPCS Code:	D	ose:			
			Frequency:					
	INFORMATION - Please indicat			applicable (*).				
Primary ICD Co			er ICD Code:					
G. CLINICAL IN	FORMATION - Required clinica	l information must be complete	ed in its <u>entirety</u> for	all precertification	i requests.			
For Initiation rec	uests (clinical documentation re	equired):						
🗌 Yes 🗌 No	Will Actemra (tocilizumab) be use	ed concomitantly with apremilast,	tofacitinib, or other	biologic DMARDs	(e.g., adalimuma	ıb, infliximab)?		
Yes No Has the patient been tested for TB with a PPD test, interferon-release assay (IGRA) or chest x-ray within 6 months of initiating a biologic therapy?								
	(check all that apply): PPD test	st 🔲 interferon-gamma assay (I	IGRA) 🗌 chest x-ra	ау				
	Please enter results of the TB test results: Positive Negative Unknown							
	If positive, Does the patient have	e latent or active TB? Latent	Active					
	If latent TB, 🗌 Yes 🗌 No Will	TB treatment be started before i	nitiation of therapy v	vith Actemra (tociliz	zumab)?			
	is non-preferred. Inflectra, Re	-		-	lumira, Kevzar	a, Rinvoq, and		
	Xeljanz/Xeljanz XR are preferred for MAPD plans. Preferred products may vary based on indication. Yes No Has the patient had prior therapy with Actemra (tocilizumab) within the last 365 days?							
		. ,	•	•				
	Has the patient had a trial and failure, intolerance, or contraindication to any of the following? (select all that apply)							
🗌 Yes 🔲 No				n to any of the following? (select all that apply)				
	Enbrel (etanercept)					nz XR (tofacitinib)		



MEDICARE FORM Actemra[®] (tocilizumab) Injectable Medication Precertification Request

(All fields must be completed and legible for precertification review.)

Page 2 of 3

Virginia (HMO D-SNP) FAX: 1-833-280-5224 PHONE: 1-855-463-0933

For other lines of business: Please use other form.

Note: Actemra is non-preferred. Preferred products may vary based on indication. See section G.

	Definit Loot Nows	Detient Dhene	Detient DOD					
Patient First Name	Patient Last Name	Patient Phone	Patient DOB					
G. CLINICAL INFORMATION (continued)	 Required clinical information must be cor 	npleted in its <u>entirety</u> for all pre	ecertification requests.					
Please explain if there are any other medical reason(s) that the patient cannot use any of the following preferred products when indicated for the patient's diagnosis (select all that apply) I Inflectra (infliximab-dyyb) Remicade (infliximab) Simponi Aria (golimumab)								
Please explain if there are any other medical reason(s) that the patient cannot use any of the following preferred products when indicated for the patient's diagnosis (select all that apply)								
Castleman's disease (CD)								
□ Yes □ No Is this request for IV formulation? □ Yes □ No Will Actemra (tocilizumab) be used as a monotherapy? □ Yes □ No Does the patient have unicentric CD? → Please identify if the patient has relapsed or refractory CD: □ Relapsed □ Refractory □ Yes □ No Will Actemra (tocilizumab) be used a second-line therapy? □ Yes □ No Is the patient human immunodeficiency virus (HIV) negative?								
Yes ☐ No Does the patient have docu → ☐ Yes ☐ No Will Actern	nra (tocilizumab) be used as subsequent ther							
Yes No Has the disease progressed	I following treatment of relapsed/refractory or	progressive disease?						
Cytokine release syndrome □ Yes No Is this request for IV formula □ Yes No Does the patient have a door release syndrome?	ation? cumented diagnosis of chimeric antigen recep	otor (CAR) T cell-induced severe	or life threatening cytokine					
☐ Please select which one: ☐ ☐ Yes ☐ No Does the patient have acute	eous formulation? oral artery biopsy or cross-sectional imaging? I temporal artery biopsy ☐ cross-sectional i -phase reactant elevation (i.e., high erythroc serum C-reactive protein [CRP]?	maging						
Juvenile idiopathic arthritis (juvenile rheum	atoid arthritis)							
Is this request for IV formulation or subcutaneous formulation? IV formulation subcutaneous formulation What is the severity of the patient's disease? Mild Moderate Severe Yes No Is there evidence that the disease is active?								
Please s	eumatoid arthritis: 🗌 Mild 📄 Moderate 🔲	l Severe indicated? onal DMARD (other than methot						
Systemic juvenile idiopathic arthritis Is this request for IV formulation or subcutaneous formulation?								
Yes No Is there evidence that the di	sease is active? nptoms include high fevers and painful polyar roidal anti-inflammatory (NSAID) monotherap	thritis?						



MEDICARE FORM Actemra[®] (tocilizumab) Injectable Medication Precertification Request

Page 3 of 3

(All fields must be completed and legible for precertification review.)

For other lines of business: Please use other form.

Note: Actemra is non-preferred. Preferred products may vary based on indication. See section G.

Patient First Name	Patient Last Name	Patient Phone	Patient DOB					
G. CLINICAL INFORMATION (continued)	- Required clinical information must be cor	mpleted in its <u>entirety</u> for all pred	certification requests.					
For ALL continuation of therapy requests (clinical documentation required for all requests):								
☐ Yes ☐ No Is this continuation request	a result of the patient receiving samples of A	ctemra (tocilizumab)?						
	Yes No Will Actemra (tocilizumab) be used concomitantly with apremilast, tofacitinib, or other biologic DMARDs (e.g., adalimumab, infliximab)?							
Yes No Is there clinical documentation supporting disease stability?								
	□ Yes □ No Is there clinical documentation supporting disease improvement?							
☐ Yes ☐ No Does the patient have any risk factors for TB?								
(check all that apply): PPD test interferon-gamma assay (IGRA) chest x-ray								
	er the results of the TB test: Results: Posit							
For IV formulation requests only (continuation of therapy requests only):								
☐ Yes ☐ No Has the patient received A	ctemra (tocilizumab) within the past 6 months	?						
Yes No Does the patient have a documented severe and/or potentially life-threatening adverse event that occurred during or following the previous infusion?								
\square Yes \square	└────────────────────────────────────							
For juvenile idiopathic arthritis (juvenile rh	eumatoid arthritis), rheumatoid arthritis or	systemic juvenile idiopathic ar	thritis only:					
Please indicate the severity of the patient's arthritis at baseline (pretreatment with Actemra (tocilizumab)): 🗌 Mild 🗌 Moderate 🔲 Severe								
H. ACKNOWLEDGEMENT								
Request Completed By (Signature Requ	iired):		Date: / /					
Any person who knowingly files a request for authorization of coverage of a medical procedure or service with the intent to injure, defraud or deceive any insurance company by providing materially false information or conceals material information for the purpose of misleading, commits a fraudulent insurance act, which is a crime and subjects such person to criminal and civil penalties.								

The plan may request additional information or clarification, if needed, to evaluate requests.