

MEDICARE FORM

Cimzia® (certolizumab pegol) Injectable Medication Precertification Request

Page 1 of 3

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

FAX: 1-855-320-8445 PHONE: 1-866-600-2139

For other lines of business:

Please use other form.

Note: Cimzia is non-preferred.

Preferred products vary based on indication. See section G below.

Please indicate: Start of treatment: Start date/					indication. See Section & Delow.		
	Continuation of therapy: D	-	1 1				
Precertification Rec	• •			e:	Fax:		
A. PATIENT INFORM							
First Name:		Last Name:			DOB:		
Address:		City:			State:	ZIP:	
Home Phone:	Work Pho	one:	Cell Phone:		Email:		
Patient Current Weigh	nt:kgs	Patient Height:i	nches or cms	s Allergies:			
B. INSURANCE INFO	RMATION						
		Does patient have	Does patient have other coverage? ☐ Yes ☐ No				
Group #:			If yes, provide ID#: Carrier Name:				
Insured:		Insured:					
Medicare: Yes	☐ No If yes, provide ID #:	·	Medicaid: Yes	☐ No If yes, prov	vide ID #:		
C. PRESCRIBER INF	ORMATION						
First Name:		Last Name:		(Check One): [☐ M.D. ☐ D.0	O. 🗌 N.P. 🔲 P.A.	
Address:		City:			State:	ZIP:	
Phone:	Fax:	St Lic #:	NPI#:	DEA #:	•	UPIN:	
Provider Email:	•	Office Contact Na	nme:	<u>.</u>	Phone:		
Specialty (Check one	e): Gastroenterologist	☐ Rheumatologist ☐	☐ Dermatologist ☐	Other:	•		
D. DISPENSING PRO	VIDER/ADMINISTRATION IN	FORMATION		-			
☐ Outpatient Infusion Center Name ☐ Home Infusion Cen Agency Name ☐ Administration code Address:	nter Phone: e: e(s) (CPT):		Specialty Other: _ Name: _ Address: _ City: _	Pharmacy	State:	ZIP:	
	Fax:		Priorie.				
-	PIN:						
NPI:			NPI:				
E. PRODUCT INFORM							
· ·	ia (certolizumab pegol) Do		Frequen	-			
F. DIAGNOSIS INFOR	RMATION - Please indicate pri	mary ICD code and specit	fy any other any other v	where applicable (*).			
Primary ICD Code: _		Secondary ICD	Code:	Othe	er ICD Code:		
G. CLINICAL INFORM	MATION - Required clinical info	ormation must be complete	ed for ALL precertificat	ion requests.			
Note: Cimzia is non-preferred. Entyvio, Inflectra, Remicade, and Simponi Aria are preferred for MA plans. For MAPD plans, Enbrel, Humira, Kevzara, Otezla, Rinvoq, Skyrizi, and Xeljanz/Xeljanz XR are preferred. Preferred products vary based on indication. Yes							
· · · · · · · · · · · · · · · · · · ·							



MEDICARE FORM

Cimzia® (certolizumab pegol) Injectable Medication Precertification Request

Page 2 of 3

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

FAX: 1-855-320-8445 PHONE: 1-866-600-2139

For other lines of business:

Please use other form.

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

Patient First Name	Patient Last Name	Patient Phone	Patient DOB					
G. CLINICAL INFORMATION (continued)	 Required clinical information must 	be completed in its entirety for all pr	recertification 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) Enbrel (etanercept) Humira (adalimumab) Kevzara (sarilumab) Otezla (apremilast) Rinvoq (upadacitinib) Skyrizi (risankizumab-rzaa) Xeljanz/Xeljanz XR (tofacitinib)								
☐ Yes ☐ No Will the requested drug be drug (DMARD) (e.g., Olum	iant, Otezla, Xeljanz)?	, , , , ,						
Yes ☐ No Has the patient ever received (including current utilizers) a biologic (e.g., Humira) or targeted synthetic DMARD (e.g., Olumiant, Xeljanz) associated with an increased risk of tuberculosis? Yes ☐ No Has the patient had a tuberculosis (TB) test (e.g., tuberculosis skin test [PPD], interferon-release assay [IGRA], chest x-ray) within 6 months of initiating therapy? (Check all that apply): ☐ PPD test ☐ interferon-gamma assay (IGRA) ☐ chest x-ray Please enter the results of the tuberculosis (TB) test: ☐ positive ☐ negative ☐ unknown								
If positive If latent to	e, Does the patient have latent or act uberculosis Yes No Has tre Please	ive tuberculosis TB? ☐ latent ☐ a	ctive unknown fection been initiated or completed?					
For Initiation Requests (clinical document								
Ankylosing spondylitis and axial spondy Please indicate loading dose at weeks 0, 2		intenance dose: freque	ncy:weeks					
Please select which of the following applies to the patient: Active ankylosing spondylitis (AS) Active axial spondyloarthritis Yes No Has the patient ever received (including current utilizers) a biologic (e.g., Humira) indicated for active ankylosing spondylitis or active axial spondyloarthritis?								
	atient experienced an inadequate re patient have an intolerance or contra		dal anti-inflammatory drugs (NSAIDs), or					
Crohn's disease	patient have an intolerance of contra	indication to at least two NOAIDs:						
	losed with moderately to severely ac- red (including current utilizers) a biolo patient have fistulizing Crohn's Disea	tive Crohn's disease (CD)? ogic (e.g., Humira) indicated for mod ise?	erately to severely active Crohn's disease?					
☐ Yes ☐	option (e.g., az [Cipro], merca	nt have a contraindication or intolera cathioprine [Azasan, Imuran], budeso otopurine [Purinethol], methylprednis	nce to at least one conventional therapy					
	→ Please select: ☐ Sulfasalazine ☐ Ciprofloxacin (Cipro) ☐ Pr	☐ Methotrexate IM or SC ☐ Me	EC) Azathioprine (Azasan, Imuran)					
Plaque psoriasis	Disease in the term							
Please indicate loading dose at weeks 0, 2 Yes No Has the patient been diag Yes No Has the patient ever receiplaque psoriasis?	nosed with moderate to severe plaqu	ue psoriasis?	-					
Yes No Are crucia	al body areas (e.g., hands, feet, face dicate the percentage of body surfac							
pharmaco	patient experienced an inadequate re ologic treatment with methotrexate, c ☐ No Does the patient have a clinic and acitretin?	yclosporine or acitretin?	ototherapy (e.g., UVB, PUVA) or eatment with methotrexate, cyclosporine					
	Please indicate clinical reaso Clinical diagnosis of alcoh Breastfeeding Cannot Pregnancy or currently pla Significant comorbidity prouncontrolled hypertension	phibits use of systemic agents (e.g.,	ated toxicity Drug interaction					



MEDICARE FORM

Cimzia® (certolizumab pegol) Injectable Medication Precertification Request

Page 3 of 3

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

FAX: 1-855-320-8445 PHONE: 1-866-600-2139

For other lines of business:

Please use other form.

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

Patient First Name	Patient Last Name	Patient Phone	Patient DOB						
C. CLINICAL INFORMATION (confine	und) Paguired clinical information must	t he completed in its entir	oty for all proportification requests						
	ued) – Required clinical information must	t be completed in its <u>entir</u>	ety for all precertification requests.						
Psoriatic arthritis Please indicate loading dose at weeks	s 0, 2 and 4: Please indicate ma	aintenance dose:	frequency: weeks						
	diagnosed with active psoriatic arthritis (l								
☐ Yes ☐ No Does the patient have	psoriatic arthritis with co-existent plaque	e psoriasis?							
Rheumatoid arthritis									
	6 0, 2 and 4: Please indicate madiagnosed with moderately to severely a								
			geted synthetic disease modifying drug (DMARD)						
· ·) indicated for moderately to severely act		g						
	Yes No Has the patient been tested for the rheumatoid factor (RF) biomarker?								
	Please indicate test result: positive negative not completed Yes No Has the patient been tested for the anti-cyclic citrullinated peptide (anti-CCP) biomarker?								
	he patient been tested for the anti-cyclic se indicate test result: positive ne								
	he patient been tested for the C-reactive								
	se indicate test result: positive ne								
	☐ Yes ☐ No Has the patient been tested for the erythrocyte sedimentation rate (ESR) biomarker?								
	se indicate test result: positive ne		onths of treatment with methotrexate at a dose greater						
	or equal to 20mg per week?	sponse after at least 5 m	shirts of treatment with methotrexate at a dose greater						
☐ Y€	es 🗌 No Has the patient experienced ar								
☐ Y∈	es No Does the patient have a contr		ate?						
	> Please indicate the contraind		impairment						
			c liver disease or other chronic liver disease						
			interaction						
	☐ Interstitial pneumonitis or								
	☐ Pregnancy or currently pla								
	☐ Blood dyscrasias (e.g., th								
	Other, please explain:								
Please indicate maintenance dose:	frequency: weeks	<u>is):</u>							
	receiving the requested drug through sa	mples or a manufacturer	s patient assistance program?						
			isease activity or improvement in signs and symptoms						
	nt with the requested drug?								
Ankylosing spondylitis and axial spondyloarthritis									
Please indicate which of the following has the patient experienced: ☐ Functional status ☐ Total spinal pain ☐ Inflammation (e.g., morning stiffness) ☐ None of the above									
Crohn's disease	Jan		70VC						
Yes No Has the patient achieved or maintained remission?									
Please indicate which of the following has the patient experienced:									
Abdominal pain or tenderness Abdominal mass Body weight Diarrhea Endoscopic appearance of the mucosa Hematocrit									
	scoring tool (e.g., Crohn's Disease Activi	ity Index [CDAI] score)	_I None of the above						
Plaque psoriasis									
Yes No Has the patient experienced a reduction in body surface area (BSA) affected from baseline? Yes No Has the patient experienced an improvement in signs and symptoms of the condition from baseline (e.g., itching, redness,									
	, scaling, burning, cracking, pain)?	n dignis and dymptoms of	the condition from passine (e.g., norming, realitiess,						
Psoriatic arthritis only									
Please indicate which of the following has the patient experienced:									
□ Number of swollen joints □ Number of tender joints □ Dactylitis □ Enthesitis □ Skin and/or nail involvement □ None of the above									
Rheumatoid arthritis Please indicate the percent of disease activity improvement from baseline in tender joint count, swollen joint count, pain, or disability: %									
H. ACKNOWLEDGEMENT									
	Do ouring all y		Data						
Request Completed By (Signature		modical procedure or s	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.