

MEDICARE FORM

Cimzia[®] (certolizumab pegol) Injectable Medication Precertification Request

Page 1 of 3

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

For Ohio MMP: FAX: 1-855-734-9389 PHONE: 1-855-364-0974

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						
🗌 Continu	uation of therapy: Date o	of last treatment				
Precertification Requested	Ву:		Phon	e:	Fax: _	
A. PATIENT INFORMATION						
First Name:		Last Name:			DOB:	
Address:		City:			State:	ZIP:
Home Phone:	Work Phone:		Cell Phone:		Email:	
Patient Current Weight:	_lbs orkgs Pati	ient Height: inch	es or cms	s Allergies:		
B. INSURANCE INFORMATIC	DN					
Aetna Member ID #:		Does patient have ot				
Group #:		_ If yes, provide ID#:		_ Carrier Name: _		
Insured:		Insured:				
Medicare: Yes No If		M	edicaid: 🗌 Yes	□ No If yes, prov	vide ID #:	
C. PRESCRIBER INFORMATI	ON					
First Name:		Last Name:		(Check One):		D. 🗌 N.P. 🗌 P.A.
Address:	T	City:			State:	ZIP:
Phone:	Fax:	St Lic #:	NPI #:	DEA #:	T	UPIN:
Provider Email:		Office Contact Name	:		Phone:	
Specialty (Check one): Gastroenterologist Rheumatologist Dermatologist Other:						
D. DISPENSING PROVIDER/	ADMINISTRATION INFORM	MATION				
Administration code(s) (CP Address: City: Phone: TIN: NP1:	Phone: T): State: Fax: PIN:	ZIP:		/ Pharmacy	_ State: Fax:	
E. PRODUCT INFORMATION						
Request is for Cimzia (certo			-	ncy:		
F. DIAGNOSIS INFORMATION	N - Please indicate primary					
Primary ICD Code:		Secondary ICD Cod			er ICD Code:	
G. CLINICAL INFORMATION	•		or ALL precertificat	ion requests.		
For All Requests (clinical doc			a ana muafannad fa	* MA plana Fax MA	DD plana Enh	al Humina Kaynana
☐ Entyvio (v ☐ Yes ☐ No Has the patie ☐ Enbrel (et ☐ Skyrizi (ri: Please explain if there are any diagnosis (select all that apply)	Xeljanz/Xeljanz XR are pre ent had prior therapy with C ent had a trial and failure, in vedolizumab) ☐ Inflectra (ent had a trial and failure, in tanercept) ☐ Humira (ada sankizumab-rzaa) ☐ Xelja other medical reason(s) tha	eferred. Preferred production imzia (certolizumab pegutolerance, or contraindic (infliximab-dyyb) tolerance, or contraindic (infliximab-dyyb) tolerance, or contraindic (infliximab) limumab) Kevzara (anz/Xeljanz XR (tofacitin at the patient cannot use	ucts vary based o ol) within the last 3 ation to any of the micade (infliximab) ation to any of the sarilumab)	on indication. 65 days? following? (select al ○	l that apply) golimumab) I that apply)] Rinvoq (upada s when indicated	citinib)



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Patient First Name	Patient Last Name	Patient Phone	Patient DOB					
G. CLINICAL INFORMATION (continue	d) – Required clinical informatic	n must be completed in its entire	ty for all precertification requests					
Please explain if there are any other med				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 t drug (DMARD) (e.g. Oli	be used in combination with any umiant, Otezla, Xeljanz)?	other biologic (e.g., Humira) or t	argeted synthetic disease-modifying ant	i-rheumatic				
Yes No Has the patient ever rece	eived (including current utilizers)	a biologic (e.g., Humira) or targ	eted synthetic DMARD (e.g., Olumiant, 2	Xeljanz)				
	ased risk of tuberculosis? patient had a tuberculosis (TB)	test (e.g., tuberculosis skin test	[PPD], interferon-release assay [IGRA],	chest x-ray)				
	months of initiating therapy?	interferen comme accov (ICDA)						
		interferon-gamma assay (IGRA) osis (TB) test:						
If posit	<i>ive,</i> Does the patient have laten	t or active tuberculosis TB? 🗌 la	itent 🗌 active 🔲 unknown					
If latent		Has treatment for latent tubercul Please select:	osis (TB) infection been initiated or comp ted treatment completed	pleted?				
For Initiation Requests (clinical docum								
Ankylosing spondylitis and axial spon								
Please indicate loading dose at weeks 0								
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?								
		uate response with at least TWC ⁻ contraindication to at least two		√SAIDs), or				
Crohn's disease								
Please indicate loading dose at weeks 0, 2, and 4: Please indicate maintenance dose: frequency: weeks								
Yes No Has the patient been diagnosed with moderately to severely active Crohn's disease (CD)? Yes No Has the patient ever received (including current utilizers) a biologic (e.g., Humira) indicated for moderately to severely active Crohn's disease?								
	e patient have fistulizing Crohn'							
			o at least one conventional therapy option or intolerance to at least one convention					
	option (e.g., azathioprine [Azasan, Imur	an], budesonide [Entocort EC], ciproflox	acin				
			thylprednisolone [Solu-Medrol], methotr asalazine [Azulfidine, Sulfazine], rifaximi					
	tacrolin			n [/ thatan],				
		alazine (Azulfidine, Sulfazine)	☐ Metronidazole (Flagyl) e (Entocort EC)	an Imuran)				
			SC Methylprednisolone (Solu-Medro					
	Rifaximin (Xifaxan)	Tacrolimus						
Plaque psoriasis Please indicate loading dose at weeks 0), 2 and 4: Please indic	ate maintenance dose:	frequency: weeks					
Yes No Has the patient been dia	agnosed with moderate to sever	e plaque psoriasis?						
Yes No Has the patient ever rec plaque psoriasis?	ceived (including current utilizers	s) Otezla or a biologic (e.g., Hum	ira) indicated for the treatment of moder	ate to severe				
Yes I No Are cru			in, intertriginous areas) affected?					
Please → Please If less than 10% of BSA		/ surface area (BSA) affected (p	ior to starting the requested medication)):%				
		uate response, or has an intoler	ance to phototherapy (e.g., UVB, PUVA)) or				
	acologic treatment with methotre							
	and acitretin?	a clinical reason to avoid pharm	acologic treatment with methotrexate, cy	yclosporine				
		l reason to avoid pharmacologic						
Clinical diagnosis of alcohol use disorder, alcoholic liver disease or other chronic liver disease Breastfeeding Cannot be used due to risk of treatment-related toxicity Drug interaction								
	Pregnancy or curre	ently planning pregnancy	,					
	-		ents (e.g., liver or kidney disease, blood	dyscrasias,				
	uncontrolled hyper							



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Patient First Name	Patient Last Name	Patient Phone	Patient DOB					
G CLINICAL INFORMATION (continu	und) - Required clinical information mus	t be completed in its entirety for all precer	tification requests					
G. CLINICAL INFORMATION (continued) – Required clinical information must be completed in its <u>entirety</u> for all precertification requests. Psoriatic arthritis								
Please indicate loading dose at weeks		aintenance dose: frequency:	weeks					
·	diagnosed with active psoriatic arthritis (,						
Rheumatoid arthritis	psoriatic arthritis with co-existent plaqu	e psonasis :						
Please indicate loading dose at weeks	0, 2 and 4: Please indicate ma		weeks					
	diagnosed with moderately to severely a	()						
) indicated for moderately to severely ac	logic (e.g., Humira) or targeted synthetic tive rheumatoid arthritis?	disease modifying drug (DMARD)					
	he patient been tested for the rheumatoi							
	e indicate test result: 🗌 positive 📋 ne							
	he patient been tested for the anti-cyclic se indicate test result:	citrullinated peptide (anti-CCP) biomarke	?					
	he patient been tested for the C-reactive							
	e indicate test result: positive ne							
	he patient been tested for the erythrocyt e indicate test result: positive ne							
🗌 Yes 🗌 No Hast	he patient experienced an inadequate re	sponse after at least 3 months of treatme	nt with methotrexate at a dose greater					
	or equal to 20mg per week?	n intelerance to methotroveto?						
	es							
[> Please indicate the contrained	lication:						
		adverse event 🔲 Renal impairment 🔲						
		nol use disorder, alcoholic liver disease or ses 🔲 Significant drug interaction 🔲 N						
		clinically significant pulmonary fibrosis						
	Pregnancy or currently pl							
		rombocytopenia, leukopenia, significant a	-					
For Continuation Requests (clinical o	locumentation required for all reques							
Please indicate maintenance dose:								
		mples or a manufacturer's patient assista						
	ed or maintained positive clinical respon it with the requested drug?	se as evidenced by low disease activity o	r improvement in signs and symptoms					
Ankylosing spondylitis and axial spo								
Please indicate which of the following								
Functional status Total spinal pain Inflammation (e.g., morning stiffness) None of the above								
Crohn's disease ☐ Yes ☐ No Has the patient achieved	ved or maintained remission?							
☐ 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	ie mucosa 🔲 Hematocrit					
Plaque psoriasis	scoring tool (e.g., Cronn's Disease Activ	ity Index [CDAI] score)	bove					
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,								
flaking Psoriatic arthritis only	, scaling, burning, cracking, pain)?							
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								
Request Completed By (Signature Required): 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 act, which is a crime and subjects such person to criminal and civil penalties.								