



Cimzia (for Maryland Only) Prior Authorization Request

Send completed form to: Case Review Unit, CVS Caremark Prior Authorization Fax: 1-866-249-6155

CVS Caremark administers the prescription benefit plan for the patient identified. This patient's benefit plan requires prior authorization for certain medications in order for the drug to be covered. To make an appropriate determination, providing the most accurate diagnosis for the use of the prescribed medication is necessary. **Please respond below and fax this form to CVS Caremark toll-free at 1-866-249-6155**. If you have questions regarding the prior authorization, please contact CVS Caremark at **1-866-814-5506**. For inquiries or questions related to the patient's eligibility, drug copay or medication delivery; please contact the Specialty Customer Care Team: CaremarkConnect[®] 1-800-237-2767.

The recipient of this fax may make a request to opt-out of receiving telemarketing fax transmissions from CVS Caremark. There are numerous ways you may opt-out: The recipient may call the toll-free number at 877-265-2711, at any time, 24 hours a day/7 days a week. The recipient may also send an opt-out request via email to do_not_call@cvscaremark.com. An opt out request is only valid if it (1) identifies the number to which the request relates, and (2) if the person/entity making the request does not, subsequent to the request, provide express invitation or permission to CVS Caremark to send facsimile advertisements to such person/entity at that particular number. CVS Caremark is required by law to honor an opt-out request within thirty days of receipt.

Patient's ID:		Patient's Date of Birth:	
Specialty:		NPI#:Physician Office Fax:	
Ph	ysician Office Telephone:	Physician Office Fax:	
Re	quest Initiated For:		
1.	Has the patient been diagnosed with any of the followard Moderately to severely active rheumatoid arthritural Active psoriatic arthritis (PsA) Active ankylosing spondylitis (AS) Active axial spondyloarthritis Moderately to severely active Crohn's disease (Composite Other	is (RA)	
2.	What is the ICD-10 code?		
Sec	ction A: Preferred Product		
3.	a) Rheumatoid arthritis: Enbrel, Humira, Kevzara b) Psoriatic arthritis: Cosentyx, Enbrel, Humira, C c) Ankylosing spondylitis: Cosentyx, Enbrel, Hum d) Crohn's disease (CD): Humira (primary); Cim *Note: Secondary preferred product for CD is Cimzia. Thad a documented inadequate response or intolerable at Can the patient's treatment be switched to a preferred	Otezla nira zia (secondary)* This preferred product option only applies to members who have dverse event with Humira. ed product? es, please call 1-866-814-5506 to have the updated form electronically (ePA). You may sign up online via remark/ or call 1-866-452-5017.	
4.	Is this request for continuation of therapy with the r	requested product? \(\begin{aligned} \text{Yes} \\ \begin{aligned} \text{No} & \text{No, skip to \$\pi6\$} \end{aligned} \)	
5.	Is the patient currently receiving the requested prod program? If unknown, answer Yes. ☐ Yes ☐ No	luct through samples or a manufacturer's patient assistance of If No, skip to Section B: All Requests	
recip		tial and is solely for the use of individuals named above. If you are not the intended g of this communication is prohibited. If you have received the fax in error, please e. Cimzia MD Step, PDPD SGM - 3/2018.	

CVS Caremark is an independent company that provides pharmacy benefit management services to CareFirst BlueCross BlueShield and CareFirst BlueChoice, Inc. members.

Has the patient had a documented inadequate response or intolerable adverse event with any of the following preferred products? Please indicate ALL that apply. <i>ACTION REQUIRED: If Yes, attach supporting chart note(s)</i> .						
	☐ Inadequate response	☐ Intole	erable adverse event			
☐ Enbrel:	☐ Inadequate response		erable adverse event			
☐ Humira:			erable adverse event			
			erable adverse event			
			erable adverse event			
			erable adverse event			
None of the above, <i>complete</i>	ione of the above, complete this form in its entirety and also complete Texas state step Therapy Section					
Is this request for continuation of therapy?						
Is the patient currently receiving Cimzia through samples or a manufacturer's patient assistance program? ☐ Yes ☐ No ☐ Unknown If Yes or Unknown, skip to #11						
How long has the patient been receiving the requested medication? months If less than 3 months, no further questions.						
D. Has the patient achieved or maintained positive clinical response to treatment as evidenced by low disease activity or improvement in signs and symptoms? <i>If Yes, no further questions</i> □ Yes □ No						
I. Has the patient received any of the following medications? If Yes, please indicate the most recent medication and skip to diagnosis section. □ Actemra □ Cosentyx □ Enbrel □ Entyvio □ Humira □ Inflectra □ Kevzara □ Kineret □ Orencia □ Remicade □ Renflexis □ Rituxan □ Siliq □ Simponi □ Simponi Aria □ Stelara □ Taltz □ Tremfya □ Tysabri □ Xeljanz □ Xeljanz XR □ No						
2. Has the patient undergone pretreatment screening for latent tuberculosis (TB) infection with either a TB skin test or an interferon gamma release assay (e.g., QFT-GIT, T-SPOT.TB)? Yes No						
nplete the following section bas	sed on the patient's diagno	sis, if app	plicable.			
tion C: Phoumatoid Arthritis						
B. Has the patient experienced an inadequate response after at least 3 months of treatment with the methotrexate dose greater than or equal to 20 mg per week? \(\begin{array}{c}\D \text{Yes} \D \text{No}\)						
4. Has the patient experienced intolerance to methotrexate? If Yes, no further questions ☐ Yes ☐ No						
i. Does the patient have a contraindication to methotrexate? \(\subseteq \text{ Yes} \) No \(\text{If Yes, indicate the contraindication:} \)						
ection D: Ankylosing Spondylitis or Axial Spondyloarthritis 6. Has the patient experienced an inadequate response with at least TWO nonsteroidal anti-inflammatory drugs (NSAIDs), or has an intolerance or contraindication to at least two NSAIDs? Yes No						
Has the patient tried and had ar		least one	conventional therapy option?			
☐ Yes - Sulfasalazine (Azulfid☐ Yes - Mesalamine, oral (Asa☐ Yes - Metronidazole (Flagyl☐ Yes - Ciprofloxacin (Cipro)☐ Yes - Prednisone	line, Sulfazine) acol, Pentasa, Delzicol, Lia ()	alda)	☐ Yes - Azathioprine (Azasan, Imuran) ☐ Yes - Mercaptopurine (Purinethol) ☐ Yes - Methotrexate ☐ Yes - Methylprednisolone (Solu-Medrol) ☐ Yes - Rifaximin (Xifaxan) ☐ No			
	preferred products? Please ind note(s). Cosentyx: Enbrel: Humira: Kevzara: Orencia (SC/ClickJect): Otezla: None of the above, complete tion B: All Requests Is this request for continuation Is the patient currently receiving Yes No Unknown How long has the patient been If less than 3 months, no furthe Has the patient achieved or ma activity or improvement in sign Has the patient received any of If Yes, please indicate the most Actemra Cosentyx If Orencia Remicade Formulate Traltz Tremfya Tysa Has the patient undergone pretitest or an interferon gamma relimplete the following section base tion C: Rheumatoid Arthritis Has the patient experienced and dose greater than or equal to 20. Has the patient experienced into Does the patient have a contrait of Yes, indicate the contraindiction D: Ankylosing Spondylitis. Has the patient experienced and (NSAIDs), or has an intolerance tion E: Crohn's Disease Has the patient tried and had an If Yes, indicate below and no fited Yes - Mesalamine, oral (Asa Yes - Metronidazole (Flagyl) Yes - Ciprofloxacin (Cipro) Yes - Prednisone	preferred products? Please indicate ALL that apply. AC note(s). Cosentyx: Inadequate response Enbrel: Inadequate response Humira: Inadequate response Humira: Inadequate response Humira: Inadequate response Orencia (SC/ClickJect): Inadequate response Orencia (SC/ClickJect): Inadequate response Orencia (SC/ClickJect): Inadequate response None of the above, complete this form in its entirety an tion B: All Requests Is this request for continuation of therapy? Yes No Is the patient currently receiving Cimzia through samples Yes No Unknown If Yes or Unknown, skip to How long has the patient been receiving the requested me If less than 3 months, no further questions. Has the patient achieved or maintained positive clinical re activity or improvement in signs and symptoms? If Yes, please indicate the most recent medication and skif Actemra Cosentyx Enbrel Entyvio Hu Orencia Remicade Renflexis Rituxan Taltz Tremfya Tysabri Xeljanz Xeljanz Has the patient undergone pretreatment screening for late test or an interferon gamma release assay (e.g., QFT-GIT nuplete the following section based on the patient's diagnor tion C: Rheumatoid Arthritis Has the patient experienced an inadequate response after dose greater than or equal to 20 mg per week? Yes [Has the patient experienced intolerance to methotrexate? Does the patient have a contraindication to methotrexate? Does the patient have a contraindication to methotrexate? The patient experienced an inadequate response with (NSAIDs), or has an intolerance or contraindication to at tion E: Crohn's Disease Has the patient tried and had an inadequate response to at If Yes, indicate below and no further questions. Yes - Mesalamine, oral (Asacol, Pentasa, Delzicol, Lia Yes - Mesalamine, oral (Asacol, Pentasa, Delzicol, Lia Yes - Metronidazole (Flagyl) Yes - Ciprofloxacin (Cipro)	preferred products? Please indicate ALL that apply. *ACTION RI note(s). Cosentyx:			

18. Does the patient have a contraindication or intolerance to at least one conventional therapy option (e.g.,

azathioprine [Azasan, Imuran], budesonide [Entocort EC], ciprofloxacin [Cipro], mesalamine [Asacol, Delzicol,

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	XPrescriber or Authorized Signature	Date (mm/dd/yy)
inf	I attest that this information is accurate and true, and that document information is available for review if requested by CVS Caremark o	
La	I attact that this information is accounts and two and that documen	station supporting this
6.	6. Has the prescriber provided proof documented in the patient chardrug is effective for the patient's condition? ☐ Yes ☐ No	rt notes that in their opinion the requested
5.	5. Do patient chart notes document the requested drug was ordered pharmacy filled the prescription and delivered to the patient or of prescribed for the patient in the last 180 days? ☐ Yes ☐ No	1 1
4.	4. Does the prescribed quantity fall within the manufacturer's publi guidelines found in the compendia of current literature (examples Pharmacology, Micromedex, current accepted guidelines)?	s: package insert, AHFS, Lexicomp, Clinical
3.	3. Is the requested drug being used for an FDA-approved indication of current literature (examples: AHFS, Lexicomp, Clinical Pharn guidelines)? ☐ Yes ☐ No	
2.	2. Is the requested drug's use consistent with the FDA-approved inc Network Drugs & Biologics Compendium indication for the trea and is supported by peer-reviewed medical literature? ☐ Yes ☐	tment of stage four advanced metastatic cancer
	Maryland State Step Therapy 1. Is the requested drug being used to treat stage four advanced met ☐ Yes ☐ No If No, skip to #3	astatic cancer?
	Pentasa, Liaida], mercaptopurine [Purinethol], methylprednisoloi [Flagyl], prednisone, sulfasalazine [Azulfidine, Sulfazine], rifaxi ☐ Yes ☐ No	2, ,