

Cimzia® (certolizumab pegol) Injectable **Medication Precertification Request**

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

For Medicare Advantage Part B: For other lines of business: Please use commercial form.

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

Would you like to use electronic prior authorization? Consider using **Availity**, our electronic prior authorization portal. Learn more about Availity from the links in the table below.

For phone or fax requests, refer to the table below for routing information. To determine which box to use, refer to the patient's Aetna ID card. State specific special needs and Medicare-Medicaid Plans may be designated on the front of the ID card or in the website URL on the back of the card. If you don't see your specific plan listed, call the number on the back of the member's ID card to confirm routing information.

For Aetna Medicare Advantage and Allina Health Aetna Medicare members send request to:

Phone: 1-866-503-0857 (TTY: 711)

Fax: 1-844-268-7263

Availity: https://www.aetna.com/health-care-professionals/resource-center/availity.html

For Aetna Medicare Advantage Virginia Dual Eligible Special Needs Plans (HMO D-SNP)

send request to:

Phone: <u>1-855-463-0933</u> Fax: 1-833-280-5224

Availity: https://www.aetnabetterhealth.com/virginia-hmosnp/providers/portal

For Aetna Assure Premier Plus Medicare Advantage New Jersey Dual Eligible Special Needs Plans

(HMO D-SNP) send request to:

Phone: 1-844-362-0934 Fax: 1-833-322-0034

Availity: https://www.aetnabetterhealth.com/new-jersey-hmosnp/providers/portal.html

For Aetna Better Health of Illinois Premier Medicare Medicaid Plan (MMP) send request to:

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

Availity: https://www.aetnabetterhealth.com/illinois/providers/portal

For Aetna Better Health of Ohio Premier Medicare Medicaid Plan (MMP) send request to:

Phone: <u>1-855-364-0974</u> Fax: 1-855-734-9389

Availity: https://www.aetnabetterhealth.com/ohio/providers/portal

For Aetna Better Health of Michigan Premier Medicare Medicaid Plan (MMP) send request to:

Phone: 1-855-676-5772 Fax: 1-844-241-2495

Availity: https://www.aetnabetterhealth.com/michigan/providers/portal.html



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→ Please describe the nature of the failure of the preferred drug

→ When was the member's adverse reaction to the preferred drug?

☐ No Has the patient had an adverse reaction to any of the following? (if yes, select all that apply below)

→ Please describe the nature of the adverse reaction to the preferred drug

- ☐ Entyvio (vedolizumab) ☐ Inflectra (infliximab-dyyb) ☐ Renflexis (infliximab-abda) ☐ Simponi Aria (golimumab)

Preferred products vary based on indication. See section G below. (All fields must be completed and legible for precertification review.) Please indicate: Start of treatment: Start date ____/ Continuation of therapy: Date of last treatment / / Precertification Requested By: A. PATIENT INFORMATION First Name: Last Name: DOB: Address: City: State: ZIP: Home Phone: Work Phone: Cell Phone: Email: Patient Current Weight: cms Allergies: lbs or kgs Patient Height: inches or **B. INSURANCE INFORMATION** Aetna Member ID #: 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, provide ID #: C. PRESCRIBER INFORMATION First Name: Last Name: (Check One): M.D. D.O. N.P. P.A. Address: City: State: ZIP: UPIN: Phone: Fax: St Lic #: NPI#: DEA #: Phone: Provider Email: Office Contact Name: Specialty (Check one): Gastroenterologist Rheumatologist Dermatologist Other: D. DISPENSING PROVIDER/ADMINISTRATION INFORMATION Dispensing Provider/Pharmacy: Patient Selected choice Place of Administration: ☐ Self-administered ☐ Physician's Office ☐ Home ☐ Retail Pharmacy ☐ Physician's Office Outpatient Infusion Center Phone: ☐ Specialty Pharmacy ☐ Mail Order Center Name: ___ Other: Home Infusion Center Phone: Name: _____ Agency Name: Agency Name. ______ Address: Address: City: State: ZIP: City: _____ State: ____ ZIP: ____ Phone: _____ Fax: _____ Phone: _____ Fax: _____ TIN: _____ PIN: ____ TIN: _____ PIN: ____ NPI: NPI: E. PRODUCT INFORMATION Request is for Cimzia (certolizumab pegol) Frequency: _ HCPCS Code: F. DIAGNOSIS INFORMATION - Please indicate primary ICD code and specify any other any other where applicable (*). Primary ICD Code: Secondary ICD Code: Other ICD Code: G. CLINICAL INFORMATION - Required clinical information must be completed for ALL precertification requests. For Initiation Requests (clinical documentation required for all requests): Note: Cimzia is non-preferred. Entyvio, Inflectra, Renflexis and Simponi Aria are preferred for MA plans. For MAPD plans, Cosentyx SC, Enbrel, Humira, Idacio, Rinvoq, Skyrizi, Sotyktu, Stelara, Tremfya, Tyenne SC and Xeljanz/Xeljanz XR are preferred. Preferred products vary based on indication. ☐ Yes ☐ No Has the patient had prior therapy with Cimzia (certolizumab pegol) within the last 365 days? ☐ No Has the patient had a trial and failure of any of the following? (if yes, select all that apply below) ─ ☐ Entyvio (vedolizumab) ☐ Inflectra (infliximab-dyyb) ☐ Renflexis (infliximab-abda) ☐ Simponi Aria (golimumab) → When was the member's trial and failure of the preferred drug?

Continued on next page

For Medicare Advantage Part B:

For other lines of business:

Please use commercial form. Note: Cimzia is non-preferred.



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Patient First Name	Patient Last Name	Patient Phone	Patient DOB		
G. CLINICAL INFORMATION (continue	l e d) – Required clinical information must be co	moleted in its entirety for all prece	rtification requests		
-					
For Initiation Requests continued (clinical documentation required for all requests): No Has the patient had a trial and failure of any of the following? (if yes, select all that apply below) Cosentyx SC (secukinumab) Enbrel (etanercept) Humira (adalimumab) Idacio (adalimumab-aacf) Rinvoq (upadacitinib) Skyrizi (risankizumab-raza) Sotyktu (deucravacitinib) Stelara (ustekinumab) Tremfya (guselkumab) Tyenne SC (tocilizumab-aazg) Xeljanz/Xeljanz XR (tofacitinib) When was the member's trial and failure of the preferred drug? Please describe the nature of the failure of the preferred drug? Humira (adalimumab) Idacio (adalimumab-aacf) Rinvoq (upadacitinib) Skyrizi (risankizumab-razaa) Enbrel (etanercept) Humira (adalimumab) Idacio (adalimumab-aacf) Rinvoq (upadacitinib) Skyrizi (risankizumab-aazg) Xeljanz/Xeljanz XR (tofacitinib) Stelara (ustekinumab) Tremfya (guselkumab) Tyenne SC (tocilizumab-aazg) Xeljanz/Xeljanz XR (tofacitinib) When was the member's adverse reaction to the preferred drug? Please describe the nature of the adverse reaction to the preferred drug? Please explain if there are any contraindications or 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) Renflexis (infliximab-abda) Simponi Aria (golimumab)					
the patient's diagnosis (select all that ap	nbrel (etanercept)	mab) 🔲 Idacio (adalimumab-aa	acf) Rinvoq (upadacitinib)		
For All Requests (clinical documentati	on required for all requests):				
drug (DMARD) (e.g., Ollowson Andrewson Andrews		g., Humira) or targeted synthetic lerculosis skin test [PPD], interferonmma assay (IGRA) chest x-racccccccccccccccccccccccccccccccccccc	DMARD (e.g., Olumiant, Xeljanz) on-release assay [IGRA], chest x-ray) ay known e unknown cion been initiated or completed?		
Ankylosing spondylitis and axial spor					
Please indicate loading dose at weeks (Please select which of the following app Yes No Has the patient ever reconspondyloarthritis? Yes No Has the		e.g., Humira) indicated for active a e with at least TWO nonsteroidal a	yloarthritis nkylosing spondylitis or active axial		
Please indicate loading dose at weeks (☐ Yes ☐ No Has the patient been di ☐ Yes ☐ No Has the patient ever red ☐ Yes ☐ No Does the	o, 2, and 4: Please indicate maintener agnosed with moderately to severely active Coeived (including current utilizers) a biologic (energy patient have fistulizing Crohn's Disease? No Has the patient tried and had an inactive patient have continuous for a continuous continuous for a cont	rohn's disease (CD)? e.g., Humira) indicated for modera dequate response to at least one of a contraindication or intolerance	tely to severely active Crohn's disease? conventional therapy option? to at least one conventional therapy		
	[Cipro], mercaptopur	ri], prednisone, sulfasalazine [Azul fidine, Sulfazine) ☐ Metronidazo ne ☐ Budesonide (Entocort EC Methotrexate IM or SC ☐ Methyl	ne [Solu-Medrol], methotrexate IM or SC, (fidine, Sulfazine], rifaximin [Xifaxan], ole (Flagyl))		



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Patient First Name	Patient Last Name	Patient Phone	Patient DOB			
G. CLINICAL INFORMATION (continu	led) – Required clinical information must be com	pleted in its entirety for all precertifica	ation requests.			
Immune checkpoint inhibitor-related	G. CLINICAL INFORMATION (continued) – Required clinical information must be completed in its entirety for all precertification requests. Immune checkpoint inhibitor-related toxicity Yes No Has the patient been diagnosed with severe immunotherapy-related inflammatory arthritis?					
☐ Yes ☐ No Has the patient been of Plaque psoriasis Please indicate loading dose at weeks ☐ Yes ☐ No Has the patient been of plaque psoriasis? ☐ Yes ☐ No Are of Please ☐ If less than 10% of BS☐ Yes ☐ No Has the pharm	iagnosed with severe immunotherapy-related inf 0, 2 and 4: Please indicate maintenance diagnosed with moderate to severe plaque psorial eceived (including current utilizers) Otezla or a burucial body areas (e.g., hands, feet, face, neck, see indicate the percentage of body surface area (65A: The patient experienced an inadequate response, macologic treatment with methotrexate, cyclospotes Does the patient have a clinical reason and acitretin? Please indicate clinical reason to avoid the company of	ce dose: frequency: asis? iologic (e.g., Humira) indicated for the scalp, genitals/groin, intertriginous are (BSA) affected (prior to starting the re- , or has an intolerance to phototherap rine or acitretin? on to avoid pharmacologic treatment void pharmacologic treatment: disorder, alcoholic liver disease or other ad due to risk of treatment-related toxic	e treatment of moderate to severe eas) affected? quested medication):% y (e.g., UVB, PUVA) or with methotrexate, cyclosporine er chronic liver disease city			
	uncontrolled hypertension)					
☐ Yes ☐ No Has the patient been	Other, please explain: 0, 2 and 4: Please indicate maintenance diagnosed with active psoriatic arthritis (PsA)? psoriatic arthritis with co-existent plaque psoriase	ce dose: frequency:	weeks			
Please indicate loading dose at weeks Yes No Has the patient been of the patient ever record (e.g., Rinvoq, Xeljanz)	0, 2 and 4: Please indicate maintenance diagnosed with moderately to severely active rhe eceived (including current utilizers) a biologic (e.g. indicated for moderately to severely active rheu	eumatoid arthritis (RA)? g., Humira) or targeted synthetic disea matoid arthritis?				
Pleas □ Yes □ No Has ti Pleas	ne patient been tested for the rheumatoid factor of eindicate test result: positive negative for the anti-cyclic citrullinate indicate test result: positive negative negative spatient been tested for the C-reactive protein	□ not completed ated peptide (anti-CCP) biomarker? □ not completed				
Pleas ☐ Yes ☐ No Has t	e indicate test result: positive negative negative ne patient been tested for the erythrocyte sedime	☐ not completed entation rate (ESR) biomarker?				
☐ Yes ☐ No Hast	e indicate test result: ☐ positive ☐ negative ☐ negati		ith methotrexate at a dose greater			
☐ Y€	Please indicate the contraindication:					
	☐ History of intolerance or adverse of ☐ Clinical diagnosis of alcohol use d		er chronic liver disease			
		topenia, leukopenia, significant anem	nia)			
Other, please explain:						
Please indicate maintenance dose:						
☐ Yes ☐ No Is the patient currently receiving the requested drug through samples or a manufacturer's patient assistance program? ☐ Yes ☐ No Has the patient achieved or maintained positive clinical response as evidenced by low disease activity or improvement in signs and symptoms since starting treatment 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						



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Patient First Name	Patient Last Name	Patient Phone	Patient DOB			
G. CLINICAL INFORMATION (continu	red) – Required clinical information must be	e completed in its <u>entirety</u> for all precert	fication requests.			
Crohn's disease						
☐ Yes ☐ No Has the patient achiev	ed 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 ☐ Improvement on a disease activity scoring tool (e.g., Crohn's Disease Activity Index [CDAI] score) ☐ 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, flaking, scaling, burning, cracking, pain)?						
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 tend	ler joint count, swollen joint count, pain,	or disability:%			
H. ACKNOWLEDGEMENT						
Request Completed By (Signature	Required):		Date: //			
insurance company by providing ma	uest for authorization of coverage of a materially false information or conceals nubjects such person to criminal and civil	material information for the purpose				

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