

Siliq

Prior Authorization Request

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.

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Patient's Name: {{MEMFIRST}} {{MEMLAST}}	Bate: {{TODAY}}				
Patient's ID: {{MEMBERID}}	Patient's Date of Birth: {{MEMBERDOB}}				
Physician's Name: {{PHYFIRST}} {{PHYLAST}	}				
Specialty:,	NPI#:				
Physician Office Telephone: {{PHYSICIANPHONE}} Physician Office Fax: {{PHYSICIANFAX}}					
Request Initiated For: {{DRUGNAME}}					

1.	What is the prescribed dose and frequency?		
	a) Loading dose:		
	Siliq 210mg	Quantity and Frequency:	
	Other:		
	b) Maintenance dose:		
	□ Siliq 210mg	Quantity and Frequency:	
	□ Other:		

- 2. What is the diagnosis? \Box Moderate to severe plaque psoriasis \Box Other
- 3. What is the ICD-10 code?
- 4. These are the preferred products for which coverage is provided for the treatment of the following indication: Plaque psoriasis: Humira, Otezla, Remicade, Skyrizi, Stelara (SC), Taltz, Tremfya. Can the patient's treatment be switched to a preferred product?

□ Yes - Please specify: If Yes, please call 1-866-814-5506 to have the updated form faxed to your office OR you may complete the PA electronically (ePA). You may sign up online via CoverMyMeds at: www.covermymeds.com/epa/caremark/ or call 1-866-452-5017. 🗆 No

□ Not applicable - Requested for condition other than plaque psoriasis, *skip to* #7

5. Does the patient have a documented inadequate response or intolerable adverse event with any of the following preferred products indicated for plaque psoriasis? ACTION REQUIRED: If Yes, attach supporting chart *note(s). Indicate ALL that apply.*

Cimzia syringe:	Inadequate response	□ Intolerable adverse event
Humira:	Inadequate response	□ Intolerable adverse event
Otezla:	Inadequate response	□ Intolerable adverse event
Skyrizi:	Inadequate response	□ Intolerable adverse event
□ Stelara SC:	Inadequate response	□ Intolerable adverse event
Taltz:	Inadequate response	□ Intolerable adverse event
Tremfya:	Inadequate response	□ Intolerable adverse event
□ No - none of the above		

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

Note: This fax may contain medical information that is privileged and confidential and is solely for the use of individuals named above. If you are not the intended recipient you hereby are advised that any dissemination, distribution, or copying of this communication is prohibited. If you have received the fax in error, please immediately notify the sender by telephone and destroy the original fax message. Siliq VF, ACSF SGM - 1/2022.

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Member Name: {{MEMFIRST}} {{MEMLAST}} DOB: {{MEMBERDOB}} PA Number: {{PANUMBER}}

- 6. Does the patient have one of the following documented clinical reasons to avoid the preferred products that are TNF inhibitors (Humira and Cimzia)? *ACTION REQUIRED: If Yes, attach supporting chart note(s).*
 - □ Yes History of demyelinating disorder
 - □ Yes History of congestive heart failure
 - □ Yes History of hepatitis B virus infection
 - □ Yes Autoantibody formation/lupus-like syndrome (attributed to TNF inhibitor)
 - □ Yes Risk of lymphoma
 - $\hfill\square$ No None of the above
 - □ Not applicable Requested medication is a TNF inhibitor
- 7. Will the requested drug be used in combination with any other biologic (e.g., Humira) or targeted synthetic disease-modifying anti-rheumatic drug (DMARD) (e.g., Olumiant, Otezla, Xeljanz)? Urs No
- 8. Has the patient ever received (including current utilizers) a biologic (e.g., Humira) or targeted synthetic DMARD (e.g., Olumiant, Xeljanz)? If Yes, skip to #8 □ Yes □ No
- 9. Has the patient had a tuberculosis (TB) test (e.g., tuberculosis skin test [PPD], an interferon-release assay [IGRA], or a chest x-ray) within 6 months of initiating therapy? *If Yes, skip to #10* □ Yes □ No
- 10. Does the patient have risk factors for tuberculosis (TB) (e.g., persons with close contact to people with infectious TB disease; persons who have recently immigrated from areas of the world with high rates of TB [e.g., Africa, Asia, Eastern Europe, Latin America, Russia]; children less than 5 years of age who have a positive TB test; groups with high rates of TB transmission [e.g., homeless persons, injection drug users, persons with HIV infection], or persons who work or reside with people who are at an increased risk for active TB [e.g., hospitals, long-term care facilities, correctional facilities, homeless shelters])?
- 11. Has the patient been tested for tuberculosis (TB) within the previous 12 months? \Box Yes \Box No
- 12. What were the results of the tuberculosis (TB) test?
 □ Positive for TB
 □ Negative for TB, *skip to #14*□ Unknown
- 13. Does the patient have latent or active tuberculosis (TB)?
 Latent
 Active
 Unknown
- 14. Has treatment for latent tuberculosis (TB) infection been initiated or completed? □ Yes – treatment initiated □ Yes – treatment completed □ No
- 15. Is this request for continuation of therapy with the requested drug? \Box Yes \Box No If No, skip to #18
- 16. Is the patient currently receiving the requested drug through samples or a manufacturer's patient assistance program? *If Yes or Unknown, skip to #18* □ Yes □ No □ Unknown
- 17. Has the patient achieved or maintained positive clinical response as evidenced by low disease activity or improvement in signs and symptoms of the condition since starting treatment with the requested drug?
 □ Yes □ No
- 18. Has the patient experienced a reduction in body surface area (BSA) affected from baseline?
 ACTION REQUIRED: If Yes, please attach chart notes or medical record documentation of decreased body surface area affected and no further questions.

 Yes
 No
- 19. Has the patient experienced an improvement in signs and symptoms of the condition from baseline (e.g., itching, redness, flaking, scaling, burning, cracking, pain)? *ACTION REQUIRED: If Yes, please attach chart notes or medical record documentation of improvement in signs and symptoms.* □ Yes □ No *No further questions*
- 20. Has the patient ever received (including current utilizers) Otezla or a biologic (e.g., Humira) indicated for the treatment of moderate to severe plaque psoriasis? ACTION REQUIRED: If Yes, please attach chart notes, medical record documentation, or claims history supporting previous medications tried and no further questions. □ Yes □ No

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- 21. Are crucial body areas (e.g., hands, feet, face, neck, scalp, genitals/groin, intertriginous areas) affected? *ACTION REQUIRED: If Yes, please attach chart notes or medical record documentation of affected areas and body surface area affected and no further questions.* □ Yes □ No
- 22. What is the percentage of body surface area (BSA) affected (prior to starting the requested medication)? ACTION REQUIRED: Please attach chart notes or medical record documentation of affected areas and body surface area affected. _______% of BSA If greater than or equal to 10% of BSA, no further questions
- 23. Has the patient experienced an inadequate response, or has an intolerance to phototherapy (e.g., UVB, PUVA) or pharmacologic treatment with methotrexate, cyclosporine and acitretin? *ACTION REQUIRED: If Yes, please attach chart notes, medical record documentation, or claims history supporting previous medications tried, including response to therapy and no further questions.* □ Yes □ No
- 24. Does the patient have a clinical reason to avoid pharmacologic treatment with methotrexate, cyclosporine and acitretin? *ACTION REQUIRED: If Yes, please attach documentation of clinical reason to avoid therapy.*□ Yes □ No *If Yes, indicate clinical reason*

I attest that this information is accurate and true, and that documentation supporting this information is available for review if requested by CVS Caremark or the benefit plan sponsor.

Х

Prescriber or Authorized Signature

Date (mm/dd/yy)

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