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



## {{PANUMCODE}}

## Stelara

## **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.

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 Name: {{MEMFIRST}} {{MEMLAST}} Date: {{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 a) Loading dose:  ☐ Stelara SQ 45 mg ☐ Stelara SQ 90 mg ☐ Stelara IV ☐ Other	e and frequency?  Quantity and Frequency: Quantity and Frequency: Quantity and Frequency:			
	b) Maintenance dose:  Stelara SQ 45 mg  Stelara SQ 90 mg  Stelara IV  Other	Quantity and Frequency: Quantity and Frequency: Quantity and Frequency:			
2.	☐ Moderately to severely a☐ Active psoriatic arthritis	o-existent plaque psoriasis active Crohn's disease (CD) active ulcerative colitis (UC) s WITHOUT co-existent plaq			
3.		Patient's		kg / lbs (circle one)	
<u>Sec</u> 4.	a) Psoriatic arthritis: Coser b) Crohn's disease: Humir c) Ulcerative colitis: Humi Can the patient's treatment Yes - Please specify: your office OR you may co www.covermymeds.com/ep No Not applicable -	ntyx, Enbrel, Humira, Otezla, Remicade ira, Remicade be switched to a preferred property of Yes, plea omplete the PA electronically on/caremark/ or call 1-866-4. Requested for condition not	roduct? se call 1-866-814-550 o (ePA). You may sign 52-5017. listed above, skip to S	06 to have the updated form faxed to up online via CoverMyMeds at:	

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5.	Is this request for continuation of therapy with the requested product?   Yes   No If No, skip to #7					
6.	Is the patient currently receiving the requested product through samples or a manufacturer's patient assistance program? If unknown, answer Yes. $\square$ Yes $\square$ No If No, skip to Section B: All Requests.					
7.	Does the patient have a documented inadequate response or intolerable adverse event with any of the following preferred products? ACTION REQUIRED: If Yes, attach supporting chart note(s). Indicate ALL that apply.  Cosentyx: Inadequate response Intolerable adverse event					
8.	Does the patient have one of the following documented clinical reasons to avoid the preferred products that are TNF inhibitors (Enbrel and/or Humira)? <i>ACTION REQUIRED: If Yes, attach supporting chart note(s)</i> .  Yes - History of demyelinating disorder - <i>Indicate drug(s)</i> :  Yes - History of congestive heart failure- <i>Indicate drug(s)</i> :  Yes - History of hepatitis B virus infection- <i>Indicate drug(s)</i> :  Yes - Autoantibody formation/lupus-like syndrome (attributed to TNF inhibitor)  Indicate drug(s):  Yes - Risk of lymphoma- Indicate drug(s):  No - none of the above  Not applicable - requested medication is a TNF inhibitor  If No - none of the above OR Not applicable - requested medication is not a TNF inhibitor, complete this form in its entirety and State Step Therapy section.					
	ection B: All Requests  Will the requested drug be used in combination with any other biologic (e.g., Humira) or targeted synthetic disease modifying antirheumatic drug (DMARD) (e.g., Olumiant, Otezla, Xeljanz)?   Yes  No					
10.	). Has the patient ever received (including current utilizers) a biologic (e.g., Humira) or targeted synthetic DMARI (e.g., Olumiant, Xeljanz) associated with an increased risk of tuberculosis? <i>If Yes, skip to #12</i> • Yes • No					
11.	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? If Yes, skip to #14  \ \square\$ Yes \ \square\$ No					
12.	Does the patient have risk factors for tuberculosis (TB) (e.g., persons with close contact to people with infectiou 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])? $\square$ Yes $\square$ No If No, skip to #17					
13.	Has the patient been tested for tuberculosis (TB) within the previous 12 months? □ Yes □ No					
14.	4. What were the results of the TB test? □ Positive for TB □ Negative for TB, skip to #17 □ Unknown					
15.	Does the patient have latent or active tuberculosis (TB)? $\square$ Latent $\square$ Active $\square$ Unknown					
16.	. Has treatment for latent tuberculosis (TB) infection been initiated or completed?  ☐ Yes - treatment initiated ☐ Yes - treatment completed ☐ No					
17.	. Is the patient currently receiving Stelara?					
18.	Is this request for continuation of therapy with the requested drug?  ☐ Yes ☐ No If No, skip to diagnosis section.					

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

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Me	mber Name: {{MEMFIRST}}} {{MEMLAST}} DOB: {{MEMB	ERDOB}} PA Number: {{PANUMBER}}			
19.	Is the patient currently receiving the requested drug through samples or a manufacturer's patient assistance program? <i>If Yes or Unknown, skip to diagnosis section.</i> $\square$ Yes $\square$ No $\square$ Unknown				
20.	Has the patient achieved or maintained positive clinical response to treatment as evidenced by low disease activity or improvement in signs and symptoms since starting treatment with the requested drug?  ☐ Yes ☐ No No further questions				
Cor	nplete the following section based on the patient's diagnosis, if a	pplicable.			
	ction C: Plaque Psoriasis AND/OR Psoriatic Arthritis <b>with</b> Co-Existent Plaque Psoriasis  Has the patient been diagnosed with moderate to severe plaque psoriasis?   Yes  No				
22.	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? <i>If Yes, no further questions.</i> □ Yes □ No				
23.	Are crucial body areas (e.g., hands, feet, face, neck, scalp, genitals/groin, intertriginous areas) affected? <i>If Yes, no further questions.</i> $\square$ Yes $\square$ No				
24.	What is the percentage of body surface area (BSA) affected (prior to starting the requested medication)?				
25.	. Has the patient experienced an inadequate response, or has an intolerance to phototherapy (e.g., UVB, PUVA) or pharmacologic treatment with methotrexate, cyclosporine or acitretin?  If Yes, no further questions   Yes  No				
26.	Does the patient have a clinical reason to avoid pharmacologic treatment with methotrexate, cyclosporine and acitretin?    Yes No  If Yes, indicate clinical reason:				
Sec	tion D: Crohn's Disease				
27.	7. Has the patient ever received (including current utilizers) a biologic (e.g., Humira) indicated for Crohn's disease? <i>If Yes, no further questions.</i> $\square$ Yes $\square$ No				
28.	Has the patient tried and had an inadequate response to at least one conventional therapy option?				
	If Yes, indicate below and no further questions.  Yes - Sulfasalazine (Azulfidine, Sulfazine)	☐ Yes - Metronidazole (Flagyl)			
	☐ Yes - Ciprofloxacin (Cipro) ☐ Yes - Budesonide (Entocort EC)	☐ Yes - Prednisone ☐ Yes - Azathioprine (Azasan, Imuran)			
	☐ Yes - Mercaptopurine (Purinethol)	☐ Yes - Methylprednisolone (Solu-Medrol)			
	☐ Yes - Methotrexate intramuscular (IM) or subcutaneous (SC)☐ Yes - Tacrolimus	☐ Yes - Rifaximin (Xifaxan)☐ No			
29.	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], mercaptopurine [Purinethol], methylprednisolone [Solu-Medrol], methotrexate, metronidazole [Flagyl], prednisone, sulfasalazine [Azulfidine, Sulfazine], rifaximin [Xifaxan], tacrolimus)?				
	tion E: Ulcerative Colitis				
30.	Has the patient ever received (including current utilizers) a biologic (e.g., Humira) or targeted synthetic disease modifying drug (e.g., Xeljanz) indicated for moderately to severely active ulcerative colitis?  If Yes, no further questions.   Yes				
31.	Has the patient tried and had an inadequate response to at least one conventional therapy option?  If Yes, indicate below and no further questions. List continues on following page.  Yes - Azathioprine (Azasan, Imuran)  Yes - Sulfasalazine  Yes - Cyclosporine (Sandimmune)  Yes - Tacrolimus (Prograf)  Yes - Mercaptopurine (Purinethol)				
	☐ Yes - Metronidazole (Flagyl) or Ciprofloxacin (Cipro) (for por	ichitis only)			

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	☐ Yes - Mesalamine (e.g., Asacol, Lialda, Pentasa, Canasa, Rowasa), balsalazide, olsalazine ☐ Yes - Corticosteroid (e.g., budesonide [Entocort, Uceris], hydrocortisone [Cortifoam, Colocort, Solu-Cortef, Cortef], methylprednisolone [Medrol, Solu-Medrol], prednisone) ☐ No			
32.	Does the patient have a contraindication or intolerance to at least one conventional therapy option (e.g., azathioprine [Azasan, Imuran], corticosteroid [e.g., budesonide [Entocort, Uceris], hydrocortisone, methylprednisolone, prednisone, cyclosporine [Sandimmune], mesalamine [Asacol, Lialda, Pentasa, Canasa, Rowasa], balsalazide, olsalazine, mercaptopurine [Purinethol], sulfasalazine, tacrolimus [Prograf], metronidazole/ciprofloxacin [for pouchitis only])?			
1.	State Step Therapy Is the requested drug being used for an FDA-approved indication or an indication supported in the compendia of current literature (examples: AHFS, Lexicomp, Clinical Pharmacology, Micromedex, current accepted guidelines)?   Yes   No			
2.	Does the prescribed quantity fall within the manufacturer's published dosing guidelines or within dosing guidelines found in the compendia of current literature (examples: package insert, AHFS, Lexicomp, Clinical Pharmacology, Micromedex, current accepted guidelines)?   Yes  No			
3.	Does the patient reside in Maryland?			
4.	Is the alternate drug (see below) FDA-approved for the medical condition being treated?  ☐ Yes ☐ No If No, please specify:			
5.	Has the prescriber provided proof, documented in the patient's chart notes, indicating that the requested drug was ordered for the patient in the past 180 days? ☐ Yes ☐ No If No, skip to #7			
6.	Has the prescriber provided proof, documented in the patient chart notes, that in their opinion the requested drug is effective for the patient's condition? $\square$ Yes $\square$ No No further questions			
7.	Are any of the following conditions met for the alternate drug (see below)?  The alternate drug is contraindicated The alternate drug is likely to cause an adverse reaction, physical or mental harm The alternate drug is expected to be ineffective The alternate drug was previously tried or a drug in the same class or with the same action was previously tried and was stopped due to ineffectiveness or an adverse event The alternate drug is not in the patient's best interest The alternate drug was tried while covered by the current or the previous health benefit plan None of the above If Yes, please specify:			
8.	Is the patient stable or currently receiving a positive therapeutic outcome with the requested drug and a change in the prescription drug is expected to be ineffective or cause harm to the patient? $\square$ Yes $\square$ No			
Alte	a) Psoriatic arthritis: Cosentyx, Enbrel, Humira, Otezla, Remicade, Simponi Aria b) Crohn's disease: Humira, Remicade c) Ulcerative colitis: Humira, Remicade			
	ttest that this information is accurate and true, and that documentation supporting this formation is available for review if requested by CVS Caremark or the benefit plan sponsor.			
<b>X</b> _	escriber or Authorized Signature Date (mm/dd/vv)			
Pre	escriber or Authorized Signature Date (mm/dd/vv)			

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