DEPARTMENT OF HEALTH SERVICES

Division of Medicaid Services F-01950 (01/2018)

STATE OF WISCONSIN

Wis. Admin. Code § DHS 107.10(2)

FORWARDHEALTH

PRIOR AUTHORIZATION / PREFERRED DRUG LIST (PA/PDL) FOR CYTOKINE AND CELL ADHESION MOLECULE (CAM) ANTAGONIST DRUGS FOR CROHN'S DISEASE AND ULCERATIVE COLITIS

Instructions: Type or print clearly. Before completing this form, read the Prior Authorization/Preferred Drug List (PA/PDL) for Cytokine and Cell Adhesion Molecule (CAM) Antagonist Drugs for Crohn's Disease and Ulcerative Colitis Instructions, F-01950A. Providers may refer to the Forms page of the ForwardHealth Portal at www.forwardhealth.wi.gov/WIPortal/Content/provider/forms/index.htm.spage for the completion instructions.

Pharmacy providers are required to have a completed PA/PDL for Cytokine and CAM Antagonist Drugs for Crohn's Disease and Ulcerative Colitis form signed by the prescriber before calling the Specialized Transmission Approval Technology-Prior Authorization (STAT-PA) system or submitting a PA request on the Portal, by fax, or by mail. Providers may call Provider Services at 800-947-9627 with questions.

SECTION I – MEMBER INFORMATION						
1. Name – Member (Last, First, Middle Initial)						
2. Member ID Number	3. Date of Birth – Member					
SECTION II – PRESCRIPTION INFORMATION						
4. Drug Name	5. Drug Strength					
6. Date Prescription Written	7. Directions for Use					
8. Name – Prescriber	National Provider Identifier (NPI) – Prescriber					
10. Address – Prescriber (Street, City, State, ZIP+4 Code)						
11. Telephone Number – Prescriber						
SECTION III – CLINICAL INFORMATION FOR CROHN'S DISEASE AND ULCERATIVE COLITIS						
12. Diagnosis Code and Description						
13. Does the member have Crohn's disease?	☐ Yes ☐ No					
14. Does the member have ulcerative colitis?	☐ Yes ☐ No					
15. Is the prescription written by a gastroenterologist or through a gastroenterology consultation?						

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SECTION III - CLINICAL INFORMATION FOR CROHN'S DISEASE AND ULCERATIVE COLITIS (Con	tinue	d)		
16. Is the member currently using the requested cytokine and CAM antagonist drug?		Yes		No
If yes, indicate the approximate date therapy was started.				
17. Has the member attempted any of the following drugs for Crohn's disease or ulcerative colitis: 6 mercaptopurine (6MP), azathioprine, oral aminosalicylates				
(balsalazide, mesalamine, olsalazine, sulfasalazine), or methotrexate?		Yes		No
If yes, indicate the drug name(s), dose, specific details about the treatment response, and the approx taken in the space provided. If additional space is needed, continue documentation in Section VI of the			each (drug was
18. Has the member attempted other drugs for Crohn's disease or ulcerative colitis				
(e.g., antibiotics, glucocorticoids, or IV immunomodulators such as infliximab)?		Yes		No
If yes, indicate the drug name(s), dose, specific details about the treatment response, and the approx taken in the space provided. If additional space is needed, continue documentation in Section VI of the			each (drug was
taken in the space provided. If additional space is needed, continue documentation in Section vi or the	15 1011	11.		

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SECTION III A – ADDITIONAL CLINICAL INFORMATION FOR NON-PREFERRED CYTOKINE AND CAM ANTAGONIST DRUG REQUESTS (PA requests for non-preferred cytokine and CAM antagonist drugs must be submitted on paper.)

19. Indicate the cytokine and CAM antagonist drugs the member has taken and provide specific details regarding the treatment response. If additional space is needed, continue documentation in Section VI of this form.

Note: A copy of the member's medical records must be submitted with the PA request to support the condition being treated, details regarding previous medication use, and outline the member's current treatment plan.

1. Drug Name	Dose	Dates Taken	
Reason for Discontinuation			
2. Drug Name	Dose	Dates Taken	
Reason for Discontinuation			
3. Drug Name	Dose	Dates Taken	
Reason for Discontinuation			
20. Indicate the clinical reason(s) why the pr	rescriber is requesting a non-	preferred cytokine and CAM antagonist drug.	
SECTION IV – AUTHORIZED SIGNATURE	<u> </u>		
21. SIGNATURE – Prescriber		22. Date Signed	
SECTION V – FOR PHARMACY PROVIDE	ERS USING STAT-PA		
23. National Drug Code (11 digits)	24. D	Supply Requested (Up to 365 Days)	
25. NPI			
26. Date of Service (MM/DD/CCYY) (For ST in the past.	ΓΑΤ-PA requests, the date of	service may be up to 31 days in the future or up to 14 days	
27. Place of Service			

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SECTION V – FOR PHARMACY PROVIDERS USING STAT-PA (Continued)						
28. Assigned PA Number						
29. Grant Date	30. Expiration Date	31. Number of Days Approved				
SECTION VI - ADDITIONAL INFORMATION						

^{32.} Include any additional information in the space below. Additional diagnostic and clinical information explaining the need for the product requested may be included here.