Division of Health Care Access and Accountability

PHS 107.10(2), Wis. Admin. Code
F-00694 (12/13)

FORWARDHEALTH

PRIOR AUTHORIZATION / PREFERRED DRUG LIST (PA/PDL) FOR CYTOKINE AND CELL ADHESION MOLECULE (CAM) ANTAGONIST DRUGS FOR 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 Ulcerative Colitis Instructions, F-00694A. Providers may refer to the Forms page of the ForwardHealth Portal at https://www.forwardhealth.wi.gov/WIPortal/Content/provider/forms/index.htm.spage for the completion instructions.

Pharmacy providers are required to have a completed Prior Authorization/Preferred Drug List (PA/PDL) for Cytokine and Cell Adhesion Molecule (CAM) Antagonist Drugs for Ulcerative Colitis form signed by the prescriber before calling the Specialized Transmission Approval Technology-Prior Authorization (STAT-PA) system or submitting a prior authorization (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)						
	T = 5 : 45:					
Member Identification Number	3. Date of Bi	rth — Member				
OF OTION III DEFOODINTION INFORMATION						
SECTION II — PRESCRIPTION INFORMATION	_					
4. Drug Name	5. Drug Strength					
6. Date Prescription Written	7. Directions for Use					
8. Name — Prescriber	1	9. National Provider I	dentif	ier (NPI) — P	rescriber
10. Address — Prescriber (Street, City, State, ZIP+4 Code)						
11. Telephone Number — Prescriber						
SECTION III — CLINICAL INFORMATION FOR ULCERATIVE C	OLITIS					
12. Diagnosis Code and Description						
13. Does the member have a diagnosis of ulcerative colitis?				Yes		No
14. Does the member have moderate to severe symptoms of ulcerative colitis?				Yes		No
15. Is the prescription written by a gastroenterologist or through a gastroenterology consultation?			No			



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SECTION III — CLINICAL INFORMATION	FOR ULCERATIVE C	OLITIS (Contir	nued)					
16. Has the member received two or more	of the drugs listed belo	w and taken ea	ch drug	for				
at least three consecutive months and e		factory therape	utic res	ponse				
or experienced a clinically significant ad	verse drug reaction?					Yes		No
If yes, check the boxes next to the drugs the member received. Indicate the dose of the drugs, specific details about the unsatisfactory therapeutic responses or clinically significant adverse drug reactions, and the approximate dates the drugs were taken in the space provided.								
1. oral aminosalicylates (balsalazide	, mesalamine, olsalazi	ne, or sulfasala	zine) _					
2. ☐ 6-mercaptopurine (6MP)								
3. ☐ azathioprine								
4. ☐ oral corticosteroids								
17. Is the member currently using Humira®	for ulcerative colitis?					Yes		No
If yes, complete Section IIIA of this form								
SECTION IIIA — CLINICAL INFORMATIO	N FOR MEMBERS CU	IRRENTLY US	NG HU	MIRA® FOR UL	CER	ATIVE (COLI	ΓIS
18. Has the member been using Humira® for				nths?		Yes		No
19. Has the member shown evidence of clir						Yes		No
SECTION IIIB — ADDITIONAL CLINICAL REQUESTS (Prior authorization requepaper.)								
20. Has the member taken a preferred cytol	kine and CAM antagon	ist drug for at le	east two)				
consecutive months and experienced a		eutic response	or expe	rienced				
a clinically significant adverse drug read	tion?					Yes		No
If yes, indicate the preferred cytokine ar therapeutic response or clinically signific antagonist drug was taken in the space	cant adverse drug read							
SECTION IV — AUTHORIZED SIGNATUR	E							
21. SIGNATURE — Prescriber			22. Da	ate Signed				
SECTION V — FOR PHARMACY PROVID	ERS USING STAT-PA							
23. National Drug Code (11 digits)		24. Days' Sup	ply Red	quested (Up to 3	65 Da	ays)		
25. NPI								
 Date of Service (MM/DD/CCYY) (For S⁻¹ in the past. 	ΓΑΤ-PA requests, the α	date of service r	nay be	up to 31 days in	the fo	uture or	up to	14 days
27. Place of Service								
28. Assigned PA Number								
29. Grant Date	30. Expiration Date			31. Number of	Days	Approv	/ed	
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SECTION VI —	ADDITIONAL	INFORMATIO	N
SECTION VI —	ADDITIONAL	. IINFUNIVIA I IU	IV

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,	additional information in the space below. Additional diagnostic and clinical information explaining the need for the uested may be included here.