FORWARDHEALTH PRIOR AUTHORIZATION / PREFERRED DRUG LIST (PA/PDL) FOR IMMUNOMODULATORS, ATOPIC DERMATITIS – TOPICAL

INSTRUCTIONS: Type or print clearly. Before completing this form, read the Prior Authorization/Preferred Drug List (PA/PDL) for Immunomodulators, Atopic Dermatitis–Topical Instructions, F-02572A. Prescribers may refer to the Forms page of the ForwardHealth Portal at https://www.forwardhealth.wi.gov/WIPortal/Subsystem/Publications/ForwardHealthCommunications.aspx?panel=Forms for the completion instructions.

Pharmacy providers are required to have a completed Prior Authorization/Preferred Drug List (PA/PDL) for Immunomodulators, Atopic Dermatitis–Topical form signed and dated 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. Prescribers and pharmacy 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			
9. Address – Prescriber (Street, City, State, Zip+4 Code)			
10. Phone Number – Prescriber	11. National Provider Identifier (NPI) – Prescriber		
SECTION III – CLINICAL INFORMATION			
12. Diagnosis Code and Description			
13. Does the member have atopic dermatitis?	🗅 Yes 🛛 No		



14. Has the member used a topical steroid for at least two c experienced an unsatisfactory therapeutic response?	onsecutive months and Yes No			
If yes, list the name and strength of the topical steroid, specific details about the unsatisfactory therapeutic response, and the approximate dates that the topical steroid was used in the space provided.				
15. Has the member used a topical steroid and experienced adverse drug reaction?	a clinically significant			
If yes, list the name and strength of the topical steroid, s reaction, and the approximate dates that the topical ster	specific details about the clinically significant adverse drug oid was used in the space provided.			
16. Has the member used a topical calcineurin inhibitor for a months and experienced an unsatisfactory therapeutic r				
If yes, list the name and strength of the topical calcineur therapeutic response, and the approximate dates that th provided.	in inhibitor used, specific details about the unsatisfactory ne topical calcineurin inhibitor was used in the space			
17. Has the member used a topical calcineurin inhibitor and significant adverse drug reaction?If yes, list the name and strength of the topical calcineur significant adverse drug reaction, and the approximate of space provided.	🗅 Yes 🔲 No			
SECTION IV – AUTHORIZED SIGNATURE				
18. SIGNATURE – Prescriber	19. Date Signed			
SECTION V - FOR PHARMACY PROVIDERS USING STA	T-PA			
20. National Drug Code (11 Digits)	21. Days' Supply Requested (Up to 365 Days)			
22. NPI	1			
23. Date of Service (DOS) (mm/dd/ccyy) (For STAT-PA req 14 days in the past.)	uests, the DOS may be up to 31 days in the future or up to			

24.	Place	of	Service
24.	Flace	UI.	Service

25. Assigned PA Number

26. Grant Date	27. Expiration Date	28. Number of Days Approved

SECTION VI – ADDITIONAL INFORMATION

29. Include any additional information in the space below. Additional diagnostic and clinical information explaining the need for the drug requested may also be included here.