

Nemluvio

Patient Ir	nformation:			
Name:				
Member	ID:			
Address:				
City, Stat	e, Zip:			
Date of B				
	•			
Prescribe	er Information:			
Name:				
NPI:				
Phone No	umber:			
Fax Num	ber			
Address:				
City, Stat	e, Zip:			
Requeste	ed Medication			
Rx Name				
Rx Streng	gth			
Rx Quan				
Rx Frequ	•			
Rx Route	-			
Administr	ration:			
Diagnosis	s and ICD Code:			
prescribed quantities of Upon rece SECTIO requests medicat	a medication for your can be provided. Plea ipt of the completed NA: Please not s. Pharmacy prinches that are not provided in the provided in	efit requires that we review certain requests for coverage with the proposition patient that requires Prior Authorization before benefit coverage or coverage complete the following questions then fax this form to the toll-free not form, prescription benefit coverage will be determined based or set that supporting clinical documentation is required or authorization reviews can be subject to trial with a set listed within the criteria. The policies are subject to this, MDH transmittals and updates to treatment guidents.	rerage of a umber liste the plan for AL addition change	additional ed below. i's rules. LPA al
1	Is the request an II	NITIAL or CONTINUATION of therapy?		
•	-	• •		
	[] Initial (If checked	I, go to 7)		
	[] Continuation (If o	checked, go to 2)		
2	Is the patient curre	ently receiving the requested medication?	Yes	No

	[If no, skip to question 7.]		
3	Has the patient been receiving medication samples of Nemluvio? [If yes, skip to question 7.]	Yes	No
4	Does the patient have a previously approved prior authorization (PA) on file with the current plan?	Yes	No
	[Note: If the patient does NOT have a previously approved PA on file for the requested medication with the current plan, the renewal request will be considered under initial therapy.]		
	[If no, skip to question 7.]		
5	Will the patient be concurrently receiving the requested medication in combination with any monoclonal antibody, anti-IL4, anti-IL5, or TSLP inhibitor such as Adbry, Cinqair, Dupixent, Fasenra, Nucala, Tezspire, or Xolair?	Yes	No
	[If yes, no further questions.]		
6	Has the patient been taking the requested medication for at least 4 months and has experienced a clinically significant benefit from the medication, as documented by the prescriber? ACTION REQUIRED: Submit supporting documentation.	Yes	No
	[Note: Atopic dermatitis Examples of a response to the requested medication are marked improvements in erythema, induration/papulation/edema, excoriations, and lichenification; reduced pruritus; decreased requirement for other topical or systemic therapies; reduced body surface area affected with atopic dermatitis; or other responses observed. Prurigo Nodularis Prurigo nodularis Examples of response include: A) Reduced nodular lesion count, B) Decreased pruritus, OR C) Reduced nodular lesion size]		
	[No further questions.]		
7	What is the indication or diagnosis?		
	[] Atopic dermatitis (If checked, go to 8)		
	[] Prurigo Nodularis (If checked, go to 24)		
	[] Other (If checked, no further questions)		

8	Is the patient greater than or equal to 12 years of age?	Yes	No
	[If no, no further questions.]		
9	Will the patient be concurrently receiving the requested medication in combination with any monoclonal antibody, anti-IL4, anti-IL5, or TSLP inhibitor such Adbry, Cinqair, Dupixent, Fasenra, Nucala, Tezspire, or Xolair?	Yes	No
	[If yes, no further questions.]		
10	Does the patient have a documented diagnosis of moderate to serve atopic dermatitis?	Yes	No
	[If no, no further questions.]		
11	Does the prescribed dosing exceed FDA approved indication?	Yes	No
	[If yes, no further questions.]		
12	Has documentation been provided to confirm that the patient has an IGA score of greater than or equal to 3 and an EASI score of greater than or equal to 16? ACTION REQUIRED: Submit supporting documentation.	Yes	No
	[If no, no further questions.]		
13	Does the patient have atopic dermatitis involvement estimated to be greater than or equal to 10% of the body surface area (BSA) according to the prescriber? ACTION REQUIRED: Submit supporting documentation.	Yes	No
	[If yes, skip to question 15.]		
14	Does the patient have atopic dermatitis affecting the following areas: hands, face, feet, eyes/eyelids, neck, scalp, skin folds, and/or genitalia?	Yes	No
	[If no, no further questions.]		
15	Has the patient tried at least TWO medium-, medium-high, high-, and/or super-high-potency prescription topical corticosteroids? ACTION REQUIRED: Submit supporting documentation.	Yes	No
	[If no, no further questions.]		
16	Has the patient tried tacrolimus ointment? ACTION REQUIRED: Submit supporting documentation.	Yes	No
	[If no, no further questions.]		

17	Were the topical corticosteroids and tacrolimus ointment EACH applied daily for at least 28 consecutive days?	Yes	No
	[If no, no further questions.]		
18	Was inadequate efficacy demonstrated with the topical therapies, according to the prescriber?	Yes	No
	[If no, no further questions.]		
19	Has the patient tried Zoryve? ACTION REQUIRED: Submit supporting documentation.	Yes	No
	[If no, no further questions.]		
20	Was Zoryve applied daily for at least 56 consecutive days?	Yes	No
	[If no, no further questions.]		
21	Was inadequate efficacy demonstrated with Zoryve, according to the prescriber?	Yes	No
	[If no, no further questions.]		
22	Has the patient tried Dupixent for at least 4 months? ACTION REQUIRED: Submit supporting documentation.	Yes	No
	[If no, no further questions.]		
23	Is the requested medication prescribed by or in consultation with an allergist, immunologist, or dermatologist?	Yes	No
	[No further questions.]		
24	Is the patient greater than or equal to 18 years of age?	Yes	No
	[If no, no further questions.]		
25	Is the requested medication prescribed by or in consultation with an allergist, immunologist, or dermatologist?	Yes	No
	[If no, no further questions.]		
26	Does the patient have greater than equal to 20 identifiable nodular lesions in total on both arms, and/or both legs, and/or trunk? ACTION REQUIRED: Submit	Yes	No

	supporting documentation.		
	[If no, no further questions.]		
27	Has documentation been provided to confirm that the patient has an IGA score of greater than or equal to 3? ACTION REQUIRED: Submit supporting documentation. [If no, no further questions.]	Yes	No
28	Has the patient experienced pruritus for 6 weeks or longer? [f yes, skip to question 31.]	Yes	No
29	Does the patient have a secondary cause of prurigo nodularis that has been identified (such as, medication induced, non- dermatologic condition such as neuropathy or psychiatric disease)? [If no, skip to question 31.]	Yes	No
30	Has the secondary cause been adequately managed according to the prescriber? [If no, no further questions.]	Yes	No
31	Has the patient tried at least TWO high- or super-high-potency prescription topical corticosteroid? [If no, no further questions.]	Yes	No
32	Was each topical corticosteroid applied daily for at least 14 consecutive days? [If no, no further questions.]	Yes	No
33	Was inadequate efficacy demonstrated with the topical corticosteroid therapy, according to the prescriber? [If no, no further questions.]	Yes	No
34	Has the patient tried and failed at least TWO traditional systemic agents for prurigo nodularis for at least 3 months? ACTION REQUIRED: Submit supporting documentation. [Note: Examples include systemic immunosuppressants, topical calcineurin inhibitors, or psoralen plus ultraviolet A light (PUVA).] [If no, no further questions.]	Yes	No



35	Does the prescribed dosing exceed FDA approved indication?	Yes	No
	[If yes, no further questions.]		
36	Will the patient be concurrently receiving the requested medication in combination with any monoclonal antibody, anti-IL4, anti-IL5, or TSLP inhibitor such as Cinqair, Fasenra, Tezspire, or Xolair?	Yes	No

Please document the diagnoses, symptoms, and/or any other information important to this review.

SECTION B: Physician Signature

PHYSICIAN SIGNATURE

DATE

FAX COMPLETED FORM TO: 1-833-896-0656

Disclaimer: An authorization is not a guarantee of payment. Member must be eligible at the time services are rendered. Services must be a covered Health Plan Benefit and medically necessary with prior authorization as per Plan policy and procedures.

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If you have any questions, call: 1-888-258-8250

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