

Request for Prior Authorization BIOLOGICALS FOR HIDRADENITIS SUPPURATIVA

FAX Completed Form To 1 (877) 733-3195

Provider Help Desk 1 (844) 236-1464

Solowa Health Link Hawki

(PLEASE PRINT – ACCURACY IS IMPORTANT) IA Medicaid Member ID # DOB Patient name Patient address Provider NPI Prescriber name Phone Fax Prescriber address Pharmacy name Address Phone Prescriber must complete all information above. It must be legible, correct, and complete or form will be returned. Pharmacy NPI Pharmacy fax

Prior authorization (PA) is required for biologicals FDA approved or compendia indicated for the treatment of Hidradenitis Suppurativa (HS). Payment for non-preferred biologic agents will be considered only for cases in which there is documentation of a previous trial and therapy failure with a preferred biologic agent.

Payment will be considered under the following conditions: 1) Request adheres to all FDA approved labeling for requested drug and indication, including age, dosing, contraindications, warnings and precautions, drug interactions, and use in special populations. 2) Patient has a diagnosis of moderate to severe HS with Hurley Stage II or III disease; and 3) Patient has at least three (3) abscesses or inflammatory nodules; and 4) Patient has documentation of adequate trials and therapy failures with the following: a) Daily treatment with topical clindamycin; b) Oral clindamycin plus rifampin; c) Maintenance therapy with a preferred tetracycline. If criteria for coverage are met, initial requests will be given for 4 months. Additional authorizations will be considered upon documentation of clinical response to therapy. Clinical response is defined as at least a 50% reduction in total abscess and inflammatory nodule count with no increase in abscess count and no increase in draining fistula count from initiation of therapy. The required trials may be overridden when documented evidence is provided that use of these agents would be medically contraindicated.

Non-Preferred

☐ Bimzelx

☐ Cosentyx

Other Humira Biosimilar:

☐ Humira☐ Simlandi☐ Yusimry	,,					
	Strength	Dosage Instructions	Quantity	Days Supply		
Diagnosis:						
☐ Hidradenitis Suppurativa: Hurley Stage ☐ I ☐ II ☐ III						
☐ Other:_						
Does patien	t have at least thre	ee (3) absesses or inflammatory nod	lules?			
П № П	Yes: Abscess/Nod	ule count:	Date ob	Date obtained:		

Preferred

☐ Adalimumab-aacf

☐ Adalimumab-adbm

☐ Adalimumab-fkip

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Topical Clindamycin Trial Name/Dose: Reason for failure:						
Oral Clindamycin Plus Rifampin Trial:						
Clindamycin: Dose:	Trial dates:					
	Trial dates:					
Maintenance Preferred Tetracycline Trial:						
Name/Dose: Reason for failure:	Trial dates:					
<u>Renewals</u>						
Document response to therapy:						
Abscess/Nodule Count: Increase Decrease (provide count): Date obtained:						
Has patient had an increase in draining fistula count since initiation of therapy? □ No □ Yes						
Other medical conditions to consider:						
Possible drug interactions/conflicting drug therapies:						
ttach lab results and other documentation as necessary.						
Prescriber signature (Must match prescriber listed above.) Date of submission	ın					

IMPORTANT NOTE: In evaluating requests for prior authorization the consultant will consider the treatment from the standpoint of medical necessity only. If approval of this request is granted, this does not indicate that the member continues to be eligible for Medicaid. It is the responsibility of the provider who initiates the request for prior authorization to establish by inspection of the member's Medicaid eligibility card and, if necessary by contact with the county Department of Health and Human Services, that the member continues to be eligible for Medicaid.

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