

Ilumya® (tildrakizumab-asmn) Medication Precertification Request

Page 1 of 2

(All fields must be completed and legible for precertification review.)

Aetna Precertification Notification Phone: <u>1-866-752-7021</u> (TTY: <u>711</u>)

FAX: <u>1-888-267-3277</u>

For Medicare Advantage Part B: Please Use Medicare Request Form

Please indicate:				, ,				
			last treatment	<u> </u>		Ган		
Precertification Re	•			Phone	:	Fax:		
A. PATIENT INFOR	RMATION		Loot Name			DOD:		
First Name:			Last Name:	l a		DOB:	1	
Address:		Т		City:		State:	ZIP:	
Home Phone:		Work Phone:		Cell Phone:	T	Email:		
Patient Current Wei	ght: lbs or	kgs Patien	t Height:inches	orcms	Allergies:			
B. INSURANCE IN								
Aetna Member ID #:			Does patient have other coverage?					
Group #:			If yes, provide ID#: Carrier Name: Insured:					
Insured:								
	☐ No If yes, provid	e ID #:	M	edicaid: U Yes	☐ No If yes, prov	/ide ID #:		
C. PRESCRIBER II First Name:	NFORMATION		Last Name:		(Check On	A). 🗆 M D	☐ D.O. ☐ N.P. ☐ P.A.	
Address:			Last Name.	City:	(Oneck On	State:	ZIP:	
			Ct 1 :- #:		DEA #	State.		
Phone:	Fax:		St Lic #:	NPI #:	DEA #:	T	UPIN:	
Provider Email:			Office Contact Name:		Phone:			
	ne): 🔲 Dermatologi							
D. DISPENSING P	ROVIDER/ADMINIST	RATION INFOR	RMATION		Provider/Pharmac			
Self-administered Physician's Office Outpatient Infusion Center Phone: Center Name: Home Infusion Center Phone: Agency Name: Administration code(s) (CPT): Address:				Specialty Name: Address: Phone:	Name:Address:		☐ Other	
E. PRODUCT INFO								
-								
			y ICD code and specif					
			Secondary ICD Co					
G. CLINICAL INFORMATION - Required clinical information must be completed in its entirety for all precertification requests.								
For All Requests (clinical documentation required): Yes								

Continued on next page



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Page 2 of 2

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Aetna Precertification Notification

Phone: <u>1-866-752-7021 (TTY: 711)</u>

FAX: 1-888-267-3277

For Medicare Advantage Part B: Please Use Medicare Request Form

Patient First Name	Patient Last Name	Patient Phone	Patient DOB						
G. CLINICAL INFORMATION (continued	N - Required clinical information must be	completed in its entirety for a	Il precertification requests						
For Initiation Requests (clinical document		e completed in its <u>entirety</u> for a	iii precertification requests.						
Plaque psoriasis	ation required).								
Please indicate loading dose at weeks 0, an	nd 4: Please indicate maintenance	dose: frequency: _	weeks						
☐ Yes ☐ No Has the patient been diagno									
Yes No Is the requested drug being	• •	_							
Yes No Has the patient ever receive the treatment of moderate to program)?	ed or is currently receiving a biologic (e.g., to severe plaque psoriasis (excluding received)	Humira) or targeted synthetic dro ving the drug via samples or a m	ug (e.g., Sotyktu, Otezla) indicated for anufacturer's patient assistance						
☐ Yes ☐ No Are crucial	body areas (e.g., hands, feet, face, neck, s	scalp, genitals/groin, intertriginou	us areas) affected?						
-	Please indicate the percentage of body surface area (BSA) affected (prior to starting the requested medication):%								
If less than 10% of BSA:			(10/0 5/0/4)						
☐ Yes ☐ No Has the patient experienced an inadequate response, or has an intolerance to phototherapy (e.g., UVB, PUVA) or pharmacologic treatment with methotrexate, cyclosporine or acitretin?									
I · · ·	No Does the patient have a clinical reaso		ent with methotrexate, cyclosporine						
T	and acitretin?	to avoia pilaaooiogio aoaa.							
	→ Please indicate clinical reason to avo	id pharmacologic treatment with	methotrexate, cyclosporine						
			ver disease or other chronic liver disease						
	☐ Breastfeeding ☐ Drug interaction		Iverse event ☐ Hypersensitivity related toxicity ☐ Significant comorbidity						
			dyscrasias, uncontrolled hypertension)						
	Other, please explain:	•	• • • • • • • • • • • • • • • • • • • •						
For Continuation Requests (clinical docum									
Please indicate maintenance dose: frequency:weeks									
☐ Yes ☐ No Is the patient currently receiving the requested drug through samples or a manufacturer's patient assistance program?									
☐ Yes ☐ No Is the requested drug being prescribed by or in consultation with a dermatologist?									
Yes No Has the patient achieved or maintained positive clinical response as evidenced by low disease activity or improvement in signs and symptoms of the condition since starting treatment with the requested drug?									
Yes No Has the patient experienced		ffected from baseline?							
	tient experienced an improvement in signs		from baseline (e.g., itching, redness,						
	aling, burning, cracking, pain)?	, ,							
H. ACKNOWLEDGEMENT									
Request Completed By (Signature Requ	uired):		Date:/ /						
Any person who knowingly files a request any insurance company by providing matinsurance act, which is a crime and subjection.	erially false information or conceals mat	erial information for the purpos							

The plan may request additional information or clarification, if needed, to evaluate requests.