

MEDICARE FORM

Tremfya® (guselkumab) Medication Precertification Request

Page 1 of 2

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

For New Jersey HMO D-SNP: FAX: 1-833-322-0034 PHONE: 1-844-362-0934

For other lines of business: Please use other form.

Note: Tremfya is non-preferred. Preferred products vary based on indication. See section G below

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Please indicate	<u> </u>		_	, ,				
Drocortification	Continuation of therapy:	Date of last treatment				For		
	Requested By:			Phone: _		Fax:		
A. PATIENT INFO	JRMATION	Last Name:				DOB:		
Address:		Last Name.		ty:		State:	ZIP:	
Home Phone:	Work Phor			ell Phone:		_	ZIF.	
				1		E-mail:		
	lbs or kgs Heigh	t: inches or	cms	Allergies:				
B. INSURANCE I		Deep notions has	ra athau		Yes □ No			
			pes patient have other coverage?					
			Insured:				_	
Medicare □ Ye	s No If yes, provide ID #:			edicaid: Yes	□ No If yes n	rovide ID #·		
C. PRESCRIBER				culculu. 163	□ No II yes, p	10 VIGC 1D #		
First Name:	THE SKIIIATION	Last Name:			(Check One	e):] D.O. [] N.P. [] P.A.	
Address:				City:	(00	State:	ZIP:	
Phone:	Fax:	St Lic #:		NPI #:	DEA #:		UPIN:	
Provider E-mail:	Į. d.v.	Office Contact N			<i>52, (n</i> .	Phone:		
	k one): Dermatologist Ga			atalogist		1 110110.		
	PROVIDER/ADMINISTRATION IN	=	Kileuilla	atologist 🗀 Oth	er			
		FORMATION		Disponsing Pro	idor/Pharmacy:	Patient Sele	ected choice	
Place of Administration: Self-administered Physician's Office				Dispensing Provider/Pharmacy: Patient Selected choice ☐ Physician's Office ☐ Retail Pharmacy				
_								
	fusion Center			Other:	=	="		
☐ Home Infusion				-			_	
	N.L							
	n code(s) (CPT):			Address:				
Address:	Stata	ZID.		-			ZIP:	
	State: Fax:							
	PIN:							
NPI:				INI II.				
E. PRODUCT INI								
	guselkumab (Tremfya) Dose: _			Frequency:				
	NFORMATION – Please indicate pri	•	city any	other where applica				
Primary ICD Cod	e:S FORMATION – Required clinical info	Secondary ICD Code: _	tod in its	a antiraty for all proc	Other ICD Co			
	quests (clinical documentation	· · · · · · · · · · · · · · · · · · ·	tea in its	s <u>entirety</u> for all pred	erillication reques	ilS.		
		• '	apremila	ast, tofacitinib, or o	ther biologic DMA	ARDs		
☐ Yes ☐ No Will guselkumab (Tremfya) be used concomitantly with apremilast, tofacitinib, or other biologic DMARDs (e.g., adalimumab,infliximab)?								
Yes No Has the patient been tested for TB with a PPD test, interferon-release assay (IGRA) or chest x-ray within 6 months of initiating a								
$ \longrightarrow $	biologic therapy? (check all that apply): ☐ PPD te	st	na assav	/(IGRA) □ ches	x-rav			
If positive, Does the patient have latent or active TB? Latent Active								
	If latent TB, ☐ Yes ☐ No Wil	I TB treatment be starte	ed befor	e initiation of thera	py with guselkum	nab (Tremfya))?	
	s non-preferred. Inflectra, Remi					umira, Otezla	ı, Rinvoq, Skyrizi, and	
	XR are preferred for MAPD plants the nations had prior therapy with		_		on.			
☐ Yes ☐ No Has the patient had prior therapy with Tremfya (guselkumab) within the last 365 days? ☐ Yes ☐ No Has the patient had a trial and failure, intolerance, or contraindication to any of the following? (select all that apply)								
☐ Inflectra (infliximab-dyyb) ☐ Remicade (infliximab) ☐ Simponi Aria (golimumab)								
☐ Yes ☐ No F	 Yes ☐ No Has the patient had a trial and failure, intolerance, or contraindication to any of the following? (select all that apply) ☐ Enbrel (etanercept) ☐ Humira (adalimumab) ☐ Otezla (apremilast) ☐ Rinvoq (upadacitinib) ☐ Skyrizi (Risankizumab-rzaa) 							
		(adalimumab) 🔲 Otezla	a (apren	nilast) ∐ Rinvoq (ι	ıpadacitinib) 🔲 🕄	Skyrizi (Risank	(izumab-rzaa)	
	☐ Xeljanz/Xeljanz XR (tofacitinib)							



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Patient First Name	Patient Last Name	Patient Phone	Patient DOB						
G. CLINICAL INFORMATION (continued) –	l Required clinical information must be compl	l leted in its <u>entirety</u> for all precertif	ication requests.						
Please explain if there are any other medical re	eason(s) that the patient cannot use any of	the following preferred products v	hen indicated for the patient's						
diagnosis (select all that apply).									
☐ Inflectra (infliximab-dyyb) ☐ Remicade (infliximab) ☐ Simponi Aria (golimumab)									
Please explain if there are any other medical re	eason(s) that the patient cannot use any of	the following preferred products v	hen indicated for the patient's						
Please explain if there are any other medical reason(s) that the patient cannot use any of the following preferred products when indicated for the patient's diagnosis (select all that apply).									
	umira (adalimumab) 🔲 Otezla (apremilast	t) 🔲 Rinvoq (upadacitinib)							
☐ Skyrizi (Risankizumab-rzaa) ☐ Xeljanz/Xeljanz XR (tofacitinib)									
Planus Pagriagia									
Plaque Psoriasis	2 Mild Moderate D Severe								
What is the severity of the patient's disease									
☐ Yes ☐ No Is there evidence that the disease is active? ☐ Yes ☐ No Is there clinical documentation of chronic disease?									
<u> </u>									
Please provide the patient's Psoriasis Area and Severity Index (PASI) score:									
Yes ☐ No Does the plaque psoriasis involve sensitive areas? <i>If yes</i> , please select: ☐ hands ☐ feet ☐ face ☐ genitals									
☐ Yes ☐ No Is the patient a candidate for systemic treatment with conventional DMARD(s)?									
☐ Yes ☐ No Was the trial with systemic conventional DMARD(s) (e.g., methotrexate, acetretin, or cyclosporine) ineffective?									
Provide the	e name and date range: Name:	Date range:/	to						
☐ Yes ☐ No Was the tri	ial with systemic conventional DMARD(s)) not tolerated?							
☐ Yes ☐ No Are systemic conventional DMARDs contraindicated?									
Yes No Is the patient a candidate for									
	ial with phototherapy ineffective?								
Please che	eck all that apply: Psoralens (methox		t (PUVA)						
	UVB with coal tar o								
	☐ UVB (standard or n ☐ Home UVB	narrow-band)							
Date range	e of phototherapy use://	to /							
	ial with phototherapy not tolerated?								
Yes No Is phototherapy contraindic									
For Continuation of Therapy (clinical documentation required for all requests): Yes No Will guselkumab (Tremfya) be used concomitantly with apremilast, tofacitinib, or other biologic DMARDs (e.g., adalimumab,									
infliximab)?	ar area consernation, that aproximacs,								
Please indicate the length of time on guselku	ımab (Tremfya):								
☐ Yes ☐ No Is there clinical documentation supporting disease stability?									
☐ Yes ☐ No Is there clinical documentation supporting disease improvement?									
Yes No Does the patient have any									
	atient had a TB test within the past year?								
(check all that apply): ☐ PPD test ☐ interferon-gamma assay (IGRA) ☐ chest x-ray									
Please enter the date and results of the TB test: Date: / Results: ☐ Positive ☐ Negative ☐ Unknown									
	R	esuits: Positive Negative	/e Unknown						
H. ACKNOWLEDGEMENT									
Request Completed By (Signature Required): Date:/									
Any person who knowingly files a request for									
any insurance company by providing materi			or misieading, commits a traudulent						

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