

MEDICARE FORM Stelara[®] (ustekinumab) Specialty Medication Precertification Request

Page 1 of 3

(Please return **Pages 1 to 3** for precertification of medications.)

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

Please indicate: Start								
		erapy: Date of	last treatment	/ / Dhanai		E ave		
Precertification Requested	-			Phone:		Fax:		
A. PATIENT INFORMATION	N					5.05		
First Name:			Last Name:			DOB:		
Address:				City:		State:	ZIP:	
Home Phone:		Work Phone:		Cell Phone:		Email:		
Current Weight: Ibs		ls Height: _	inches or	_ cms Allergies:				
B. INSURANCE INFORMAT					_			
Aetna Member ID #:			Does patient have other coverage? Yes No					
Group #:			If yes, provide ID#: Carrier Name:					
			Insured:					
C. PRESCRIBER INFORMA	ATION		1 ()]					
First Name:			Last Name:		(Check One		D.O. 🗌 N.P. 🗌 P.A.	
Address:				City:	1	State:	ZIP:	
Phone:	Fax:	<u>.</u>	St Lic #:	NPI #:	DEA #:		UPIN:	
Provider Email:		Offic	ce Contact Name:		Phone:			
D. DISPENSING PROVIDE	R/ADMINIST	RATION INFOR	MATION					
Place of Administration:				Dispensing Provider/	-			
			Home	Physician's Office Retail Pharmacy Specialty Pharmacy Mail Order Other:			41 ···	
Outpatient Infusion Center Center Name:								
Home Infusion Center				Name: Address:				
Agency Name:				City:	S	tate:	ZIP:	
Administration code(s) (CP				Phone:		Fax:		
Address:				TIN:		PIN:		
City:				NPI:				
Phone:				E. PRODUCT INFORM				
NPI:	TIN: PIN:			Request is for Stelara (ustekinumab) (Check One):				
Please explain if there are a	ny medical re	ason(s) why the	patient cannot self-	45mg 90mg Route: Frequency:				
inject the requested drug:		., .		HCPCS Code:				
F. DIAGNOSIS INFORMAT								
Primary ICD Code:						de:		
G. CLINICAL INFORMATIO	-			d for ALL precertification	equests.			
For Initiation Requests (clini								
Note: Stelara is non-preferre Remicade are preferred for								
Preferred products vary bas	ed on indicat	ion.				•		
\Box Yes \Box No Has the patient had prior therapy with Stelara (ustekinumab) 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) □ Entyvio (vedolizumab) □ Inflectra (infliximab-dyyb) □ Remicade (infliximab) □ Simponi Aria (golimumab) 								
Yes No Has the patient had a trial and failure, intolerance, or contraindication to any of the following? (select all that apply)								
	• • •		umab) 🔲 Otezla (apre	emilast) 🔲 Rinvoq (upada	citinib) 🗌 S	Skyrizi (risankizı	umab-rzaa)	
	Xeljanz XR (tof / other medical		ne patient cannot use ar	av of the following preferred	products w	nen indicated fo	r 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)								
🗌 Entyvio (vedolizumab) 🔲 Inflectra (infliximab-dyyb) 🔲 Remicade (infliximab) 🔲 Simponi Aria (golimumab)								
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			calliot 400 u	,	1 20000 1			
🗌 Enbrel (etanercept) 🔲 Humira (adalimumab) 🗍 Otezla (apremilast) 🗍 Rinvoq (upadacitinib) 🗍 Skyrizi (risankizumab-rzaa) 🗍 Xeljanz/Xeljanz XR (tofacitinib)								



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Patient First Name	Patient Last Name	Patient Phone	Patient DOB					
	ical information must be completed for ALL pred							
	given concomitantly with apremilast, tofacitinib, or							
biologic therapy?	Has the patient been tested for TB with a PPD test, interferon-release assay (IGRAs) or chest x-ray within 6 months of initiation a biologic theorem /2							
	· (check all that apply): PPD test interferon-gamma assay (IGRA) chest x-ray							
	Please enter results of the TB test: positive negative unknown							
	If positive, does the patient have latent or active TB?							
If latent TB, 🗌 Yes 🔲 No Will TB treatment be started before initiation of therapy with Stelara (ustekinumab)?								
Crohn's Disease								
$+$ $ \cdot$ $*$	Does the patient have a diagnosis of fistulizing Crohn's disease? • Please indicate how long the patient has been diagnosed with fistulizing Crohn's disease:							
	Does the patient have a diagnosis of Crohn's disease?							
	→ Please indicate the severity of the patient's disease: ☐ mild ☐ moderate ☐ severe							
	ent have a documented diagnosis of active Crohn's	s disease?						
	 Please select all signs/symptoms that apply: abdominal pain arthritis bleeding diarrhea internal fistulae intestinal obstruction 							
	on perianal disease spondylitis weight l							
☐ Yes ☐ No Have the Cro	hn's disease symptoms remained active despite tre		ine, azathioprine, or					
corticosteroid								
	Please check all medications that apply: 6-mercaptopurine azathioprine corticosteroids- please identify: prednisone hydrocortisone methylprednisolone Other:							
	Yes No Will the initial (induction) dose of Stelara (ustekinumab) be administered intravenously?							
Yes No Will all doses after the initial d	ose be administered subcutaneously?							
Plaque Psoriasis (Adult and Pediatric)								
☐ Yes ☐ No Is there clinical documentation	of chronic disease? the patient's plaque psoriasis: 🗌 mild 🛛 modera	ata. 🗖 sovoro						
☐ Yes ☐ No Is there evidence that the dise								
\square Yes \square No Is the patient a candidate for s								
Please select: D phototherap	y 🔲 systemic therapy 🗌 phototherapy and syst	temic therapy						
Please provide the patient's Psoriasis Area and								
Please indicate the percentage of body surface	area affected by plaque psoriasis:% ect sensitive areas? <i>If yes</i> , please select:	n foot □ fooo □ gooi	tals					
Adult	ici sensitive areas? II yes, piease select. 🗋 nanus							
	itional DMARD(s) (e.g., methotrexate, acetretin, or	cyclosporine) ineffective?						
	with systemic conventional DMARD(s) not tolerated							
	conventional DMARD(s) contraindicated?	_						
] cyclosporine methotrexate mycophenola	ate 📋 Other, please explai	n:					
☐ Yes ☐ No Was a trial with phototherapy ☐ Yes ☐ No Was the trial								
☐ Yes ☐ No Is photothera								
	Soralens (methoxsalen, trioxsalen) with UVA lig	ght (PUVA)						
	UVB with coal tar or dithranol							
]UVB (standard or narrow band)]Home UVB							
	None of the above							
	rial: Less than 1 month 1 month 2 mon	ths 🔲 3 months or greater	-					
Pediatric		-						
	neffective, not tolerated, or contraindicated?							
	Psoralens (methoxsalen, trioxsalen) with UVA lig	ght (PUVA)						
] UVB with coal tar or dithranol] UVB (standard or narrow band)							
] Home UVB							
	None of the above	_						
Please indicate the length of t	ial: 🗌 Less than 1 month 📋 1 month 📋 2 mon	ths 🔲 3 months or greater						

Continued on next page



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Patient First Name	Patient Last Name	Patient Phone	Patient DOB			
G. CLINICAL INFORMATION - Required clinica	l information must be completed for ALL pred	certification requests.				
Psoriatic Arthritis						
	t moderate to severe plaque psoriasis?					
Yes ☐ No Is there evidence that the disease ☐ Yes ☐ No Does the patient have axial psori						
	nt with 2 or more non-steroidal anti-inflammatory	v drugs (NSAIDs) ineffective?				
Please provide t	ne names and length of treatment:					
NSAID #1:						
Yes No Does the patient have non-axial						
multiple joints?	have severe disease at presentation, defined a	s severe disability at onset with	erosive disease involving			
	Was the treatment with methotrexate ineffective	?				
	$ ightarrow$ \Box Yes \Box No Was treatment with methotr		dicated?			
	\rightarrow Please select: \Box not toler					
		ment with another conventiona				
	Please se	elect: Cyclophosphamide				
		sulfasalazine Oth				
Ulcerative Colitis			· · · · · · · · · · · · · · · · · · ·			
Yes No Has the patient been diagnosed v	vith moderately to severely active ulcerative coli	tis (UC)?				
	reviously received a biologic or targeted synthet	tic disease modifying drug (e.g	., Xeljanz) indicated for			
,	verely active ulcerative colitis?	nonce to at least one conventio	and thereasy ention?			
	Has the patient tried and had an inadequate res ☐ Yes ☐ No Does the patient have a contra					
	•		g., budesonide, hydrocortisone			
	[Entocort, Uceris], methylpred					
	mesalamine [Asacol, Lialda, P					
	sulfasalazine, tacrolimus [Prog Please select: 🗌 Azathioprine [Azasan, Imura					
	hydrocortisone [Cortifoam, Colocort, Solu-Cort					
	prednisone) Cyclosporine (Sandimmune)					
	Canasa, Rowasa) 🗌 Mercaptopurine (Purinet		rolimus (Prograf)			
Vec. No. Will the initial (induction) does of	Metronidazole (Flagyl) or Ciprofloxacin (Cip					
Yes No Will the initial (induction) dose of Yes No Will all doses after the initial dose		Susiy ?				
For Continuation of Therapy (clinical documenta						
Please indicate length of time on Stelara (ustekinumab):						
☐ Yes ☐ No Is there clinical documentation of						
Yes No Does the patient have any risk fa		, -				
\square Yes \square No Has the patient h		_				
	pply): PPD test interferon-gamma assay					
For Crohn's Disease, Plaque Psoriasis, Ulcerativ	e results of the TB test: positive negative positive					
Please indicate the severity of the disease at baseline (pretreatment with Stelara (ustekinumab)): mild moderate severe						
For Psoriatic Arthritis:						
Yes No Does the patient have co-existen	t moderate to severe plaque psoriasis?					
H. ACKNOWLEDGEMENT						
Request Completed By (Signature Required)	:		_ Date: / / /			
Any person who knowingly files a request for authorization of coverage of a medical procedure or service with the intent to injure, defraud or deceive any						
insurance company by providing materially false information or conceals material information for the purpose of misleading, commits a fraudulent insurance act, which is a crime and subjects such person to criminal and civil penalties.						
The plan may request additional information or clarification, if needed, to evaluate requests.						