

MEDICARE FORM Stelara® (ustekinumab) Specialty

Stelara® (ustekinumab) Specialty Medication Precertification Request

age 1 of 3

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

For Ohio MMP:

FAX: 1-855-734-9389 PHONE: 1-855-364-0974 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 of treatment: Continuation of the								
∟ Precertification Req		лару. Баце от	iasi irealineni	Phone:		Fax:			
A. PATIENT INFORM				I Hone		1 ax			
	MATION		Lost Name			DOB:			
First Name:			Last Name:	T ₀		DOB:	Taip		
Address:				City:		State:	ZIP:		
Home Phone:		Work Phone:		Cell Phone:		Email:			
Current Weight:		s Height: _	inches or	_ cms Allergies:					
B. INSURANCE INFO									
Aetna Member ID #			Does patient have other coverage?						
Group #:			If yes, provide ID#: Carrier Name:						
Insured:			Insured:						
C. PRESCRIBER IN	FORMATION								
First Name:			Last Name:	1	(Check One):	D.O. N.P. P.A.		
Address:				City:		State:	ZIP:		
Phone:	Fax:		St Lic #:	NPI #:	DEA #:		UPIN:		
Provider Email:		Offic	ce Contact Name:		Phone:				
D. DISPENSING PRO	OVIDER/ADMINISTR	ATION INFOR	MATION						
Place of Administra	ation:			Dispensing Provide	er/Pharmacy:				
☐ Self-administered	☐ Self-administered ☐ Physician's Office ☐ Home				☐ Physician's Office ☐ Retail Pharmacy				
Outpatient Infusion Center Phone:				☐ Specialty Pharmacy ☐ Mail Order ☐ Other:					
):			Name:	Name:				
☐ Home Infusion Cen				Address:					
	e:			City: State: ZIP: Phone: Fax:					
☐ Administration code Address:	TIN: PIN:								
City:	Sta	ate: ZIF	 P:	NPI:					
· ·				E. PRODUCT INFO	RMATION				
NPI:				☐ 45mg ☐ 90mg					
Please explain if there are any medical reason(s) why the patient cannot self-				Frequency:					
inject the requested	arug:			HCPCS Code: IV SC					
E DIAGNOSIS INFO	PMATION - Please i	ndicate primary	ICD Code and specify	any other any other w	here applicable	a /*\			
Primary ICD Code:			ary ICD Code:		Other ICD Co				
			tion must be completed			uc			
For Initiation Request	<u> </u>		<u> </u>	d for ALL precentification	ni requests.				
				are preferred for MA pl	ans For MAPI) nlans Entyvio	Inflectra and		
Note: Stelara is non-preferred. Entyvio, Inflectra, Remicade, and Simponi Aria are preferred for MA plans. For MAPD plans, Entyvio, Inflectra, and Remicade are preferred for ulcerative colitis and Enbrel, Humira, Otezla, Rinvoq, Skyrizi, and Xeljanz/Xeljanz XR are preferred for other indications.									
	Preferred products vary based on indication. ☐ Yes ☐ 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)									
☐ Enbrel (etanercept)☐ Humira (adalimumab)☐ Otezla (apremilast)☐ Rinvoq (upadacitinib)☐ Skyrizi (risankizumab-rzaa)☐ Xeljanz/Xeljanz XR (tofacitinib)									
Please explain if there	are any other medical	,	ne patient cannot use an	y of the following preferr	red products wh	nen 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)									
☐ Enbrel (etanercept) ☐ Humira (adalimumab) ☐ Otezla (apremilast) ☐ Rinvoq (upadacitinib) ☐ Skyrizi (risankizumab-rzaa) ☐ Xeljanz/Xeljanz XR (tofacitinib)									
☐ ⊼eijanz/⊼eijanz ⊼K (tolacitinib)									



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Patient First Name	Patient Last Name	Patient Phone	Patient DOB						
O CLINICAL INFORMATION D L. III.		('6' - ('							
G. CLINICAL INFORMATION - Required clinic	·	•	delinerum ek indiirine ek\Q						
☐ Yes ☐ No Has the patient been tested for	Will Stelara (ustekinumab) be given concomitantly with apremilast, tofacitinib, or other biologic DMARDs (e.g., adalimumab, infliximab)? 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 therapy? (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?								
Crohn's Disease	If latent TB, ☐ Yes ☐ No Will TB treatment be started before initiation of therapy with Stelara (ustekinumab)?								
	o Does the patient have a diagnosis of fistulizing Crohn's disease?								
Please indicate how long the pa	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 Severe No Does the patient have a documented diagnosis of active Crohn's disease?								
	Please select all signs/symptoms that apply:								
	☐ abdominal pain ☐ arthritis ☐ bleeding ☐ diarrhea ☐ internal fistulae ☐ intestinal obstruction								
_	perianal disease spondylitis weight lo								
Yes No Have the Crohi corticosteroids	o's disease symptoms remained active despite trea	atment with 6-mercaptopurine,	azathioprine, or						
	all medications that apply: 6-mercaptopurine [azathioprine							
	☐ corticosteroids- please identify: ☐ prednisone ☐ hydrocortisone ☐ methylprednisolone ☐ Other:								
	Will the initial (induction) dose of Stelara (ustekinumab) be administered intravenously?								
Yes No Will all doses after the initial dos	e be administered subcutaneously?								
Plaque Psoriasis (Adult and Pediatric) ☐ Yes ☐ No Is there clinical documentation of	of chronic disease?								
Please indicate the severity of t	ne patient's plaque psoriasis: mild modera	te 🗌 severe							
Yes No Is there evidence that the disea									
Yes No Is the patient a candidate for sy	stemic therapy or phototherapy? ☐ systemic therapy ☐ phototherapy and syste	omio thorony							
Please provide the patient's Psoriasis Area and S		эппс шегару							
Please indicate the percentage of body surface ar									
Yes No Does the plaque psoriasis affect	t sensitive areas? <i>If yes</i> , please select: ☐ hands	☐ feet ☐ face ☐ genitals	;						
Adult	and DMADD(a) (a.g. mathetrayeta acetratin ar a	oveleenerine) in effective?							
	Was a trial of systemic conventional DMARD(s) (e.g., methotrexate, acetretin, or cyclosporine) ineffective? Yes No Was the trial with systemic conventional DMARD(s) not tolerated?								
	onventional DMARD(s) contraindicated?	•							
	cyclosporine 🔲 methotrexate 🔲 mycophenolat	te 🔲 Other, please explain: _							
Yes ☐ No Was a trial with phototherapy in ☐ Yes ☐ No Was the trial wi									
Yes No Is phototherapy									
	Psoralens (methoxsalen, trioxsalen) with UVA light	ht (PUVA)							
	UVB with coal tar or dithranol								
	UVB (standard or narrow band) Home UVB								
	None of the above								
Please indicate the length of tria	ıl: ☐ Less than 1 month ☐ 1 month ☐ 2 mont	hs 3 months or greater							
Pediatric									
	effective, not tolerated, or contraindicated?	ht (DUI\(A)							
	Psoralens (methoxsalen, trioxsalen) with UVA light UVB with coal tar or dithranol	III (FUVA)							
	UVB (standard or narrow band)								
	Home UVB								
	None of the above I: ☐ Less than 1 month ☐ 1 month ☐ 2 month	hs							
. idada indicate the length of the		ooo or groutor							

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Patient First Name	Patient Last Name	Patient Phone	Patient DOB			
G. CLINICAL INFORMATION - Required clinical	information must be completed for ALL prec	certification requests.				
Psoriatic Arthritis Yes No Does the patient have co-existent Is there evidence that the disease Does the patient have axial psorial Yes No Was the treatmen Please provide the NSAID #1:	moderate to severe plaque psoriasis? is active? tic arthritis? t with 2 or more non-steroidal anti-inflammatory e names and length of treatment:					
Yes No Does the patient have non-axial properties of the patient have non-axial properties of the patient have non-ax	nave severe disease at presentation, defined as		erosive disease involving			
		exate not tolerated or contraind	I DMARD ineffective? ☐ cyclosporine ☐ leflunomide			
Ulcerative Colitis						
moderately to sev	eviously received a biologic or targeted syntheterely active ulcerative colitis? as the patient tried and had an inadequate responding the patient tried and had an inadequate responding to the patient have a contration option (e.g., azathioprine [Azasa [Entocort, Uceris], methylpred mesalamine [Asacol, Lialda, Psulfasalazine, tacrolimus [Progresses select: Azathioprine [Azasan, Imurar hydrocortisone [Cortifoam, Colocort, Solu-Corte prednisone) Cyclosporine (Sandimmune) Canasa, Rowasa) Mercaptopurine (Purinet Metronidazole (Flagyl) or Ciprofloxacin (Ciptelara (ustekinumab) be administered intravence.	cic disease modifying drug (e.g. conse to at least one convention indication or intolerance to at lessan, Imuran], corticosteroid [e.g. nisolone, prednisone, cyclospo entasa, Canasa, Rowasa], mei graf], metronidazole/ciprofloxacin] Corticosteroid (e.g., budgef, Cortef], methylprednisolone Mesalamine (e.g., Apriso, Ashol) Sulfasalazine Tacrro) (for pouchitis only)	enal therapy option? east one conventional therapy g., budesonide, hydrocortisone rine [Sandimmune], rcaptopurine [Purinethol], in [for pouchitis only])? esonide [Entocort, Uceris], [Medrol, Solu-Medrol], sacol, Lialda, Pentasa,			
For Continuation of Therapy (clinical documentat						
	t of the patient receiving samples of Stelara (us disease stability or improvement? ☐ disease st cors for TB?	tability ☐ improvement (IGRA) ☐ chest x-ray				
For Crohn's Disease, Plaque Psoriasis, Ulcerative Colitis: 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-existent 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.