

MEDICARE FORM

Orencia® (abatacept) Injectable Medication Precertification Request

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

For Medicare Advantage Part B: Phone: 1-866-503-0857 (TTY: 711)

FAX: <u>1-844-268-7263</u>

For other lines of business: Please use other form.

se other form.

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

Please indicate: Start of treatment, Start Date:/ Continuation of therapy, date of last treatment:/								
Precertification Requested B		Phone:			Fax:			
A. PATIENT INFORMATION								
First Name:		Last Name:				DOB:		
Address:		•	City	y:		State:	ZIP:	
Home Phone:	Work Phone:		<u> </u>	II Phone:		Email:		
Patient Current Weight:	lbs or kgs Pa	atient Height:i	inche	es or cms Al	llergies	}:		
B. INSURANCE INFORMATION								
Aetna Member ID #:		Does patient have of	other	coverage?	☐ No)		
Group #:		=		Carrier				
Insured:		Insured:						
C. PRESCRIBER INFORMAT	ION							
First Name:		Last Name:		(Che	ck one	<u>∍):</u>	O.O. N.P. P.A.	
Address:			Cit	ity:		State:	ZIP:	
Phone:	Fax:	St Lic #:	NF	PI #:	DEA#	‡ :	UPIN:	
Provider Email:		fice Contact Name:			Phone) :		
D. DISPENSING PROVIDER/	ADMINISTRATION INFO	RMATION						
Place of Administration:			1	Dispensing Provider		-		
☐ Self-administered			☐ Physician's Office ☐ Retail Pharmacy					
Outpatient Infusion Center	· · · · · · · · · · · · · · · · · · ·		_	Specialty Pharmac				
	DI .		-	Other:				
Home Infusion Center			-	Name:				
Agency Name:	١٠		-	Address:				
Address:				City:				
City:		ZIP:		Phone:				
Phone:				TIN:		PIN:		
TIN:				NPI:				
NPI:				E. PRODUCT INFORMATION				
Please explain if there are any	the patient cannot self-			•	• *			
inject the requested drug:				Dose: HCPCS Code:		□ IV □ SC		
F. DIAGNOSIS INFORMATIO	N - Please indicate prima	ary ICD code and speci			re anni			
Primary ICD Code:								
•								
G. CLINICAL INFORMATION - Required clinical information must be completed for ALL precertification requests. For Initiation requests (clinical documentation required):								
Yes No Will Orencia (abatacept) 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 biologic therapy?								
Check all that apply): PPD test interferon-gamma assay (IGRA) chest x-ray								
Please enter the results of the TB test: ☐ Positive ☐ Negative ☐ Unknown If positive, Does the patient have latent or active TB? ☐ Latent ☐ Active								
If latent TB, Yes No Will TB treatment be started before initiation of therapy with Orencia (abatacept)?								
Note: Orencia is non-preferred. Inflectra, Remicade, Simponi Aria, and unbranded infliximab are preferred for MA plans. Enbrel, Humira, Kevzara, Otezla, Rinvoq, Skyrizi, Stelara, and Xeljanz/Xeljanz XR are preferred for MAPD plans. Preferred products vary based on indication.								
☐ Yes ☐ No Has the patient had prior therapy with Orencia (abatacept) 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) ☐ Unbranded infliximab								
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) ☐ Kevzara (sarilumab) ☐ Otezla (apremilast) ☐ Rinvoq (upadacitinib)								
☐ Skyrizi (risankizumab-rzaa) ☐ Stelara (ustekinumab) ☐ Xeljanz/Xeljanz XR (tofacitinib) 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).								
☐ Inflectra (infliximab-dyyb) ☐ Remicade (infliximab) ☐ Simponi Aria (golimumab) ☐ Unbranded infliximab								



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FAX: <u>1-844-268-7263</u>

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Patient First Name	Patient Last Name	Patient Phone	Patient DOB						
G. CLINICAL INFORMATION (continued)	 Required clinical information must be c 	completed in its <u>entirety</u> for all p	ecertification requests.						
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) ☐ Kevzara (sarilumab) ☐ Otezla (apremilast) ☐ Rinvoq (upadacitinib)									
☐ Skyrizi (risankizumab-rzaa) ☐ Stelara (ustekinumab) ☐ Xeljanz/Xeljanz XR (tofacitinib)									
Juvenile idiopathic arthritis (juvenile rheumatoid arthritis)									
Please indicate the severity of the patient's disease: Mild Moderate Severe									
Yes No Is there evidence that the disease is active?									
Psoriatic Arthritis ☐ Yes ☐ No Is there evidence that the disease is active?									
☐ Yes ☐ No Does the patient have axial psoriatic arthritis?									
☐ Yes ☐ No Was the treatment with 2 or more non-steroidal anti-inflammatory drugs (NSAIDs) ineffective?									
Please provide the names of treatment:									
NSAID #1:									
NSAID #2:									
Yes ☐ No Does the patient have non-axi ☐ Yes ☐ No Was treatme									
	No Was treatment with methotrexate not	tolerated or contraindicated?							
	→ Please select: ☐ not tolerated ☐ o								
	☐ Yes ☐ No Was a trial with a co		i-rheumatic drug ineffective?						
		cyclophosphamide cyclospo							
	_	leflunomide							
	Ш	Other: Please explain:							
Rheumatoid Arthritis									
Please indicate the severity of the patient's rheumatoid arthritis: Mild Moderate Severe									
☐ Yes ☐ No Is there evidence that the disease is active? ☐ Yes ☐ No Was treatment with methotrexate ineffective?									
Yes No Was treatment with methotrexate method rective? Yes No Was treatment with methotrexate not tolerated or contraindicated?									
	ect: not tolerated contraindicated								
☐ Yes ☐	No Was treatment with another convention								
Provide select: ☐ azathioprine ☐ hydroxychloroquine ☐ leflunomide ☐ sulfasalazine									
For Continuation requests (clinical documentation required):									
Yes ☐ No Will Orencia (abatacept) be used concomitantly with apremilast, tofacitinib, or other biologic DMARDs (e.g., adalimumab, infliximab)? Please indicate the severity of the patient's disease at baseline (pretreatment with Orencia (abatacept)): ☐ Mild ☐ Moderate ☐ Severe									
Yes									
☐ Yes ☐ No Is there clinical documentation supporting disease improvement?									
☐ Yes ☐ No Does the patient have any risk factors for TB?									
└────────────────────────────────────									
`	at apply): PPD test interferon-gamm	, , <u> </u>							
Please the results of the TB test: ☐ Positive ☐ Negative ☐ Unknown ☐ Yes ☐ No Is this continuation request a result of the patient receiving samples of Orencia (abatacept)?									
For Juvenile idiopathic arthritis (juvenile rheumatoid arthritis) IV formulation only (continuation of therapy requests only):									
☐ Yes ☐ No Has the patient received Orencia (abatacept) within the past 6 months?									
Yes No Does the patient have a documented severe and/or potentially life-threatening adverse event that occurred during or following									
the previous									
,	No Could the adverse reaction be mana	iged through pre-medication in the	e home or office setting?						
H. ACKNOWLEDGEMENT									
Request Completed By (Signature Req	uired):		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									
Insurance act which is a crime and subjects	such person to criminal and civil penaltic	es							

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