

MEDICARE FORM

Orencia® (abatacept) Injectable **Medication Precertification Request**

Page 1 of 2

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

For Ohio MMP:

FAX: 1-855-734-9389 **PHONE**: 1-855-364-0974

For other lines of business:

Please use other form.

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

Please indicate: Start of treatmer	t, Start Date: / /	Continuation of thera	y, date of last treatment:	1 1	
Precertification Requested By:		Phone:	Fax:		
A. PATIENT INFORMATION					
First Name:	Last Name	:	DOB:		
Address:		City:		ZIP:	
Home Phone:	Work Phone:	Cell Phone:	Email:	-11 .	
		1			
Patient Current Weight: lbs B. INSURANCE INFORMATION	or kgs Pallent Heigh	:: inches or cms	Allergies:		
	Deec notic	ant have other coveres 2	-		
Aetna Member ID #:		nt have other coverage?			
Group #: Insured:		ide ID#: Carri	er Name:		
	Insured:				
C. PRESCRIBER INFORMATION	Lost Name	. (6)	sock anal: □MD □DO		
First Name:	Last Name	•	neck one): M.D. D.O		
Address:		City:		IP:	
Phone: Fax:	St Lic #:	NPI #:		JPIN:	
Provider Email:	Office Contact	Name:	Phone:		
D. DISPENSING PROVIDER/ADMINISTRATION INFORMATION					
Place of Administration:		Dispensing Provide			
	sician's Office	Physician's Offic		СУ	
	Phone:				
Center Name: Other:					
☐ Home Infusion Center					
			Address:		
Address:			State:		
Address:	State: 7ID:		Fax:		
Phone:		1 IIV	PIN:		
TIN:					
NPI:		E. PRODUCT INFO	RMATION		
Please explain if there are any medical reason(s) why the patient cannot self-			Request is for: Orencia (abatacept):		
inject the requested drug:			Frequency:		
		HCPCS Code:			
F. DIAGNOSIS INFORMATION - Ple					
Primary ICD Code:				_	
G. CLINICAL INFORMATION - Requ	uired clinical information must b	e completed for ALL precertificatio	n requests.		
For Initiation requests (clinical docur	nentation required):				
☐ Yes ☐ No Will Orencia (abatacep	t) be used concomitantly with apr	emilast, tofacitinib, or other biologic [DMARDs (e.g., adalimumab, infl	liximab)?	
Yes No Has the patient been to	ested for TB with a PPD test, inte	feron-release assay (IGRA) or chest	x-ray within 6 months of initiatir	ng a	
biologic therapy?					
└────────────────────────────────────					
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, and Simponi Aria are preferred for MA plans. Enbrel, Humira, Kevzara, Otezla, Rinvoq,					
Skyrizi, 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)					
☐ 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) ☐ 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)					



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Please use other form.

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Patient First Name	Patient Last Name	Patient Phone	Patient DOB			
G CLINICAL INFORMATION (continued)	Required clinical information must be a	completed in its entirety for all i	precertification requests			
	G. CLINICAL INFORMATION (continued) – Required clinical information must be completed in its entirety for all precertification 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)	sacon(e) anal and panem cannot acc any cr	and remarking presented products	The second of the patients			
☐ Enbrel (etanercept) ☐ H	☐ Enbrel (etanercept) ☐ Humira (adalimumab) ☐ Kevzara (sarilumab) ☐ Otezla (apremilast) ☐ Rinvoq (upadacitinib)					
☐ Skyrizi (risankizumab-rzaa) Xeljanz/Xeljanz XR (tofacitinib)					
Juvenile idiopathic arthritis (juvenile rheumatoid arthritis)						
Please indicate the severity of the patient's disease: Mild Moderate Severe Ves No Is there evidence that the disease is active?						
Yes No Has the patient had an ineffective						
	t with Enbrel (etanercept) not tolerated or con ct: ☐ not tolerated ☐ contraindicated	traindicated?				
Psoriatic Arthritis	t. Hot tolerated Contraindicated					
Yes No Is there evidence that the dise	ease is active?					
Yes No Does the patient have axial ps						
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-ax ☐ Yes ☐ No Was treatme	ial psoriatic arthritis?					
· — —	ent with methotrexate ineffective? No Was treatment with methotrexate not	tolerated or contraindicated?				
	→ Please select: ☐ not tolerated ☐	contraindicated				
	Yes No Was a trial with a co					
	•	cyclophosphamide				
		Other: Please explain:				
Rheumatoid Arthritis						
Please indicate the severity of the patient's rhe		Severe				
☐ Yes ☐ No Is there evidence that the disease is active?						
☐ Yes ☐ No Was treatment with methotrexate ineffective? ☐ Yes ☐ No Was treatment with methotrexate not tolerated or contraindicated?						
> Please sele	ect: not tolerated contraindicated					
☐ Yes ☐	☐ Yes ☐ No Was treatment with another conventional DMARD (other than methotrexate) ineffective? → 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 ☐ No Is there clinical documentation supporting disease stability? ☐ Yes ☐ No Is there clinical documentation supporting disease improvement?						
☐ Yes ☐ No Does the patient have any risk factors for TB?						
Yes No Has the patient had a TB test within the past year?						
	nat apply): ☐ PPD test ☐ interferon-gamn results of the TB test: ☐ Positive ☐ Nega		′			
Yes No Is this continuation request a r						
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 infusion?						
· · · · · · · · · · · · · · · · · · ·	☐ No Could the adverse reaction be mana	aged through pre-medication in t	he home or office setting?			
H. ACKNOWLEDGEMENT						
Request Completed By (Signature Req	juired):		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	s such person to criminal and civil penalti	es.				

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