Member Name: {{MEMFIRST}} {{MEMLAST}} DOB: {{MEMBERDOB}} PA Number: {{PANUMBER}}									
{{PANUMCODE}}									
{{DISPLAY_PAGNAME}}} {{PACDESCRIPTION}}									
forr {{C	This fax machine is located in a secure location as required by HIPAA regulations. Fax complete signed and dated forms to {{COMPANY_NAME}} at {{CLIENT_PAG_FAX}}. Please contact {{COMPANY_NAME}} at {{CLIENT_PAG_PHONE}} with questions regarding the prior authorization process. When conditions are met, we will authorize the coverage of {{DRUGNAME}}.								
Patient's Name: {{MEMFIRST}} {{MEMLAST}} Date: {{TODAY}} Patient's ID: {{MEMBERID}} Patient's Date of Birth: {{MEMBERDOB}} Physician's Name: {{PHYFIRST}} {{PHYLAST}} Patient Phone: << MEMPHONE>> Specialty: NPI#: Physician Office Telephone: {{PHYSICIANPHONE}} Physician Office Fax: {{PHYSICIANFAX}} Physician Office Address: << PHYADDRESS1>> << PHYADDRESS2>> << PHYCITY>>, << PHYSTATE>> Orug Name: {{DRUGNAME}}									
Qu	anti	ty: Frequen	ncy: Strength: Expected Length of Therapy:						
Rou	ite (of Administration: sis: < <diagnosis>> ICD Code</diagnosis>	Expected Length of Therapy:						
	Wł	nat is the prescribed quantity and free Loading dose: Orencia IV 250 mg Orencia SQ 125 mg Orencia SQ 87.5 mg Orencia SQ 50 mg Other Maintenance dose: Orencia IV 250 mg Orencia SQ 125 mg Orencia SQ 125 mg Orencia SQ 125 mg Orencia SQ 87.5 mg Orencia SQ 87.5 mg Orencia SQ 50 mg							
2.	□ Other								
3.	Wł	nat is the ICD-10 code?							
4.	Wł	nat is the patient's weight?	kg						
	ection A: All Requests Will the requested drug be used in combination with any other biologic (e.g., Humira) or targeted synthetic drug (e.g., Olumiant, Otezla, Xeljanz) for the same indication? Yes No								
2.			g current utilizers) a biologic (e.g., Humira) or targeted synthetic drug th an increased risk of tuberculosis? <i>If Yes, skip to #6</i> □ Yes □ No						

Mo	ember Name: {{MEMFIRST}} {{MEMLAST}} DOB: {{MEMBERDOB}} PA Number: {{PANUMBER}}				
3.	Has the patient had a tuberculosis (TB) test (e.g., tuberculosis skin test [TST], interferon-release assay [IGRA]) within 6 months of initiating therapy? ☐ Yes ☐ No				
4.	What were the results of the tuberculosis (TB) test? ☐ Positive for TB ☐ Negative for TB, skip to #6 ☐ Unknown				
5.	Which of the following applies to the patient? ☐ Patient has latent TB and treatment for latent TB has been initiated ☐ Patient has latent TB and treatment for latent TB has been completed ☐ Patient has latent TB and treatment for latent TB has not been initiated ☐ Patient has active TB				
6.	Is the requested drug being prescribed by or in consultation with a: ☐ Rheumatologist ☐ Dermatologist ☐ Oncologist ☐ Hematologist ☐ Other:				
7.	Is the patient currently receiving Orencia?				
8.	Is this request for continuation of therapy with the requested drug? ☐ Yes ☐ No If No, skip to diagnosis initiation section.				
9.	Is the patient currently receiving the requested drug through samples or a manufacturer's patient assistance program? <i>If Yes or Unknown, skip to diagnosis initiation section.</i> \square Yes \square Unknown \square No				
10.	Has the patient achieved or maintained a positive clinical response as evidenced by low disease activity or improvement in signs and symptoms since starting treatment with the requested drug? Yes No				
Con	nplete the following section based on the patient's diagnosis, if applicable.				
Sec	tion B: Rheumatoid Arthritis				
Con	ntinuation				
1.	Has the patient experienced substantial disease activity improvement (e.g., at least 20% from baseline) in tender joint count, swollen joint count, pain, or disability? <i>ACTION REQUIRED: If Yes, please attach chart notes or medical record documentation supporting positive clinical response and substantial disease activity improvement.</i> \square Yes \square No				
Init	iation				
2.	Has the patient ever received or is currently receiving a biologic (e.g., Humira) or targeted synthetic drug (e.g., Rinvoq, Xeljanz) indicated for the treatment of moderately to severely active rheumatoid arthritis (excluding receiving the drug via samples or a manufacturer's patient assistance program)? **ACTION REQUIRED: If Yes, please attach chart notes, medical record documentation, or claims history supporting previous medications tried and no further questions. **Description** Description**				
3.	Does the patient meet either of the following: a) the patient was tested for the rheumatoid factor (RF) biomarker and the RF biomarker test was positive, or b) the patient was tested for the anti-cyclic citrullinated peptide (anti-CCP) biomarker and the anti-CCP biomarker test was positive? ACTION REQUIRED: If Yes please attach laboratory results, chart notes, or medical record documentation of biomarker testing and skip to #4. \(\sigma\) Yes \(\sigma\) No				
4.	Has the patient been tested for all of the following biomarkers: a) rheumatoid factor (RF), b) anti-cyclic citrullinated peptide (anti-CCP), and c) C-reactive protein (CRP) and/or erythrocyte sedimentation rate (ESR)? <i>ACTION REQUIRED: If Yes, please attach laboratory results, chart notes, or medical record documentation of biomarker testing.</i> \square Yes \square No <i>If No, no further questions</i>				
4.	Has the patient experienced an inadequate response after at least 3 months of treatment with methotrexate at dose greater than or equal to 15 mg per week? ACTION REQUIRED: If Yes, please attach chart notes, medical record documentation, or claims history supporting previous medications tried, including response to therapy and no further questions. Yes No				
5.	Has the patient experienced an intolerance to methotrexate? ACTION REQUIRED: If Yes, please attach chart notes, medical record documentation, or claims history supporting previous medications tried, including response to therapy and no further questions. \square Yes \square No				

M	Member Name: {{MEMFIRST}}} {{ME	EMLAST}} DOB:	{{MEMBERDOP	B}} PA Number: {{PANUMBER}}		
6.	Does the patient have a contraindication to methotrexate? <i>ACTION REQUIRED: If Yes, please attach documentation of clinical reason to avoid therapy and indicate the contraindication.</i> \square Yes \square No					
7.	Please indicate the contraindication to History of intolerance or adverse et Risk of treatment-related toxicity Breastfeeding Significant comorbidity prohibits uncontrolled hypertension) Clinical diagnosis of alcohol use di Other	vent use of systemic age	☐ Hypersensitivnts (e.g., liver or l	currently planning pregnancy ity kidney disease, blood dyscrasias,		
	ection C: Polyarticular Juvenile Idiopath	ic Arthritis (pJIA),	Oligoarticular Ju	venile Idiopathic Arthritis		
Coi 1.	 Which of the following has the patien ACTION REQUIRED: Please attach clinical response. Functional ability Functional status 	chart notes or me	dical record docu	imentation supporting positive limitation of movement		
	☐ Number of joints with active arthri					
. .	☐ None of the above					
Init 2.	Has the patient ever received or is currently receiving a biologic (e.g., Humira) or targeted synthetic drug (e.g., Xeljanz) indicated for the treatment of moderately to severely active articular juvenile idiopathic arthritis (excluding receiving the drug via samples or a manufacturer's patient assistance program)? ACTION REQUIRED: If Yes, please attach chart notes, medical record documentation, or claims history supporting previous medications tried and no further questions. Yes					
3.	Has the patient had an inadequate response to methotrexate or another conventional synthetic drug (e.g., leflunomide, sulfasalazine, hydroxychloroquine) administered at an adequate dose and duration? **ACTION REQUIRED: If Yes, please attach chart notes, medical record documentation, or claims history supporting previous medications tried, including response to therapy, and no further questions. **Description** Description** 1. Yes					
4.	Has the patient had an inadequate response to a trial of scheduled non-steroidal anti-inflammatory drugs (NSAIDs) and/or intra-articular glucocorticoids (e.g., triamcinolone hexacetonide)? ACTION REQUIRED: If Yes, please attach chart notes, medical record documentation, or claims history supporting previous medications tried, including response to therapy and skip to #6. Yes No					
5.	Does the patient have one of the following risk factors for poor outcome: a) involvement of ankle, wrist, hip, sacroiliac joint, and/or temporomandibular joint (TMJ), b) presence of erosive disease or enthesitis, c) delay in diagnosis, d) elevated levels of inflammation markers, or e) symmetric disease? If Yes, no further question. Yes					
6.	Does the patient have any of the following risk factors for disease severity and potentially a more refractory disease course: a) positive rheumatoid factor, b) positive anti-cyclic citrullinated peptide antibodies, or c) pre-existing joint damage? Yes No					
7.	1	Does the patient meet any of the following: a) high-risk joints are involved (e.g., cervical spine, wrist, or hip) b) high disease activity, or c) high risk for disabling joint disease? Yes No				
	ection D: Psoriatic Arthritis					
<i>Coi</i> 1.	ontinuation Which of the following has the patien ACTION REQUIRED: Please attach clinical response.					
	 Number of swollen joints □ Enthesitis □ Functional status 	□ Number of ter□ Axial disease□ C-reactive pro	•	□ Dactylitis□ Skin and/or nail involvement□ None of the above		

M	ember Name: {{MEMFIRST}} {{MEMLAST}} DOB: {{MEMBERDOB}} PA Number: {{PANUMBER}}		
Inii 2.	Has the patient ever received (including current utilizers) a biologic (e.g., Humira) or targeted synthetic drug (e.g., Rinvoq, Otezla) indicated for the treatment of active psoriatic arthritis (excluding receiving the drug via samples or a manufacturer's patient assistance program)? ACTION REQUIRED: If Yes, please attach chart notes, medical record documentation, or claims history supporting previous medications tried, and no further questions. Yes		
3.	What is the patient's disease severity? \square Mild to moderate \square Severe If Severe, no further questions.		
4.	Does the patient have enthesitis? If Yes, no further questions. \square Yes \square No		
5.	Has the patient had an inadequate response to methotrexate, leflunomide, or another conventional synthetic drug (e.g., sulfasalazine) administered at an adequate dose and duration? <i>ACTION REQUIRED: If Yes, please attach chart notes, medical record documentation, or claims history supporting previous medications tried, including response to therapy and no further questions.</i> \square Yes \square No		
6.	Has the patient had an intolerance to methotrexate, leflunomide, or another conventional synthetic drug (e.g., sulfasalazine)? ACTION REQUIRED: If Yes, please attach chart notes, medical record documentation, or claims history supporting previous medications tried, including response to therapy, and no further questions. Yes No		
7.	Does the patient have a contraindication to methotrexate or leflunomide? ACTION REQUIRED: If Yes, please attach documentation of clinical reason to avoid therapy and indicate the contraindication and skip to #9. \(\sigma\) Yes \(\sigma\) No		
8.	Does the patient have a contraindication to another conventional synthetic drug (e.g., sulfasalazine)? <i>ACTION REQUIRED: If Yes, please attach documentation of clinical reason to avoid therapy and indicate the contraindication.</i> \square Yes \square No		
9.	Please indicate the contraindication to methotrexate or leflunomide. History of intolerance or adverse event Risk of treatment-related toxicity Pregnancy or currently planning pregnancy Hypersensitivity Significant comorbidity prohibits use of systemic agents (e.g., liver or kidney disease, blood dyscrasias, uncontrolled hypertension) Clinical diagnosis of alcohol use disorder, alcoholic liver disease, or other chronic liver disease Other		
<u>Sec</u> 1.	Has the patient experienced an inadequate response to systemic corticosteroids? ACTION REQUIRED: If Yes, please attach chart notes, medical record documentation, or claims history supporting previous medications tried, including response to therapy and no further questions. Yes		
2.	Does the patient have an intolerance or contraindication to corticosteroids? ACTION REQUIRED: If Yes, please attach chart notes, medical record documentation, or claims history supporting previous medications tried (if applicable), including response to therapy. If therapy is not advisable, please attach documentation of clinical reason to avoid therapy. \square Yes \square No		
<u>Sec</u> 1.	tion F: Immune Checkpoint Inhibitor-Related Toxicity Does the patient have myocarditis?		
2.	Has the patient experienced an inadequate response to systemic corticosteroids? ACTION REQUIRED: If Yes, please attach chart notes, medical record documentation, or claims history supporting previous medications tried, including response to therapy and no further questions. Yes		
3.	Does the patient have an intolerance or contraindication to corticosteroids? ACTION REQUIRED: If Yes, please attach chart notes, medical record documentation, or claims history supporting previous medications tried (if applicable), including response to therapy. If therapy is not advisable, please attach documentation		

of clinical reason to avoid therapy. ☐ Yes ☐ No

Mei	mber Name: {{MEMFIRST}} {{MEMLAST}} DOB: {{MEMBERDOB}} PA Number: {{PANUMBER}		
1.]	ion G: Prophylaxis of Acute Graft Versus Host Disease Is the patient undergoing hematopoietic stem cell transplantation (HSCT) from a matched or 1 allelemismatched unrelated donor? Ves No		
2.	Will the requested medication be used in combination with a calcineurin inhibitor (e.g., cyclosporine,		
1	tacrolimus) and methotrexate? \(\begin{align*} \text{Yes} \bigsize \text{No} \\ \t		
_			
prov	est that the medication requested is medically necessary for this patient. I further attest that the information ided is accurate and true, and that the documentation supporting this information is available for review if ested by the claims processor, the health plan sponsor, or, if applicable a state or federal regulatory agency.		
Pres	scriber (Or Authorized) Signature and Date		