

2024 PRIOR AUTHORIZATION REQUEST FORM

Individual and Family Plans

Kevzara

Fax back to: (833) 605-4407 Phone: (215) 991-4300

Jefferson Health Plans manages the pharmacy drug benefit for your patient. Certain requests for coverage require review with the prescribing physician. Please answer the following questions and fax this form to the number listed above.

PLEASE NOTE: Any infor	mation (patient, prescribe	er, drug, labs) left blank, illegible,	or not attached WILL delay the review process.
Patient Name:		Prescriber Name:	
Member Number:		Fax: Phone:	
Date of Birth:		Office Contact:	
Line of Business: Exchange	- PA	NPI:	State Lic ID:
Address:		Address:	
City, State ZIP:		City, State ZIP:	
Primary Phone:		Specialty/facility na	ame (if applicable):
REQUEST FOR EXPEDITED REVIEW: the enrollee or the enrollee's ability to re		ing below, I certify that the standard re	view timeframe may seriously jeopardize the life or health of
Drug Name:			
Strength:			
Directions / SIG:			
Please attach any pertine	-	uding labs and information fo er the following questions and	r this member that may support approval. d sign.
Q1. Request type:			
☐ Initial		☐ Continuat	ion
Q2. Is the medication b	eing prescribed b	y or in consultation with	a rheumatologist?
□ Yes		□No	
tuberculosis skin test [l	PPD], an interferor rapy for persons w	n-release assay [IGRA], vho are naïve to biologic	(TB) test (which can include a or a chest x-ray) within 6 drugs or targeted synthetic
☐ Yes		□ No	
Q4. Is the requested manageted synthetic drug		sed concomitantly with a	ny other biologic drug or
☐ Yes		□ No	
Q5. What is the patien	's diagnosis?		

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Patient Name:	Prescriber Name:			
☐ Rheumatoid arthritis (RA) - Go to 6	☐ Polymyalgia rheumatica (PMR) - Go to 10			
Q6. For moderately to severely active RA, has the patient previously received or is unable to receive a biologic or targeted synthetic drug (e.g., Rinvoq, Xeljanz) indicated for moderately to severely active rheumatoid arthritis? Please attach documentation				
☐ Yes	□ No			
Q7. For moderately to severely active RA, has the patient had an inadequate response to at least a 3-month trial of methotrexate despite adequate dosing (i.e., titrated to at least 15 mg/week) OR is unable to take methotrexate?				
☐ Yes	□ No			
Q8. For RA, has the patient tested positive for Rheumatoid factor (RF) and Anti-cyclic citrullinated peptide (anti-CCP)? Please attach documentation.				
☐ Yes	□ No			
Q9. Has the patient been tested for RF, Anti-CCP and C-reactive protein (CRP) and/or erythrocyte sedimentation rate (ESR)? Please attach documentation.				
☐ Yes	□ No			
Q10. For treatment of polymyalgia rheumatica (PMR), does the member meet all of the following criteria: A) 18 years of age or older; B) Documented diagnosis of polymyalgia rheumatica (PMR); C) Documented inadequate response, contraindication, or intolerance to systemic corticosteroids or steroid tapers?				
☐ Yes	□ No			
Q11. For continuation, has the patient achieved or maintained a positive clinical response as evidenced by disease activity improvement of at least 20% from baseline in tender joint count, swollen joint count, pain, or disability?				
☐ Yes	□ No			

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Patient Name:	Prescriber Name:
Q12. Additional Information:	
Prescriber Signature	Date
ŭ	2024 Prior Authorization Reques

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