US Family Health Plan Prior Authorization Request Form for upadacitinib (Rinvog ER)

To be completed and signed by the prescriber. To be used only for prescriptions which are to be filled through the Department of Defense (DoD) US Family Health Plan Pharmacy Program. US Family Health Plan is a TRICARE contractor for DoD.

The completed form may be faxed to 855-273-5735

OR

The patient may attach the completed form to the prescription and **mail** it to:

Attn: Pharmacy, 77 Warren St, Brighton, MA 02135

QUESTIONS? Call 1-877-880-7007

For Atopic Dermatitis, prior authorization expires after 12 months. Renewal PA criteria will be approved indefinitely. For renewal of therapy an initial USFHP prior authorization approval is required. Medical documentation must be attached. Failure to provide could result in denial.

Step	Please complete patient and physician information (please print):				
1	Patient Name: Phy	sician Name:			
	Address:	Address:			
	Sponsor ID #	Phone #:			
	•	Secure Fax #:			
Step	Please complete clinical assessment:				
2	Is the requested medication being used for non- radiographic axial spondyloarthritis, rheumatoid arthritis, psoriatic arthritis, ulcerative colitis, ankylosing spondylitis, or Crohn's disease?	□ Yes	□ No		
		proceed to question 2	proceed to question 6		
	2. Humira is the Department of Defense's preferred targeted biologic agent. Has the patient tried Humira?	□ Yes	□ No		
		proceed to question 3	proceed to question 5		
	3. Has the patient had an inadequate response to Humira?	□ Yes	□ No		
		proceed to question 6	proceed to question 4		
	4. Has the patient experienced an adverse reaction to Humira that is not expected to occur with the requested agent?	□ Yes	□ No		
		proceed to question 6	STOP		
			Coverage not approved		
	5. Does the patient have a contraindication to Humira (adalimumab)?	□ Yes	□ No		
		proceed to question 6	STOP		

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6.	What is the indication or diagnosis?	☐ Moderate to severe active rheumatoid arthritis – proceed to question 7		
		☐ Moderate to severe atopic dermatitis - proceed to question 14		
		☐ Active psoriatic arthritis (PsA) - proceed to question 9		
		☐ Moderately to severely active ulcerative colitis - proceed to question 12		
		☐ Moderately to severely active Crohn's disease - proceed to question 13		
		☐ Ankylosing spondylitis – proceed to question 22		
		☐ Non-radiographic axial spondyloarthritis – proceed		
		to question 21		
		☐ Other indication or diagnosis – STOP: Coverage not approved		
7.	The provider acknowledges that for rheumatoid arthritis a trial of Xeljanz or Olumiant is required before Rinvoq.			
		Proceed to question 8		
8.	Has the patient experienced an inadequate response or adverse reaction to Xeljanz OR Xeljanz XR OR Olumiant?	□ Yes	□ No	
		proceed to question 11	proceed to question 10	
9.	Has the patient experienced an inadequate response or adverse reaction to Xeljanz OR Xeljanz XR?	☐ Yes proceed to question 11	☐ No proceed to question 10	
		p. 2 - 2 - 2 - 2 - 2 - 2 - 2 - 2 - 2 - 2	p	
10.	Does the patient have a contraindication to Xeljanz OR	☐ Yes	□ No	
	Xeljanz XR OR Olumiant?	proceed to question 11	STOP Coverage not approved	
11.	Has the patient had an inadequate response or an	☐ Yes	□ No	
	intolerance to methotrexate or other nonbiologic	proceed to question 13	STOP	
	disease-modifying antirheumatic drugs (DMARDs)?		Coverage not approved	
12.	Has the patient had an inadequate response to non- biologic systemic therapy (for example – methotrexate, aminosalicylates (e.g. sulfasalazine, mesalamine),	☐ Yes	□ No	
		proceed to question 13	STOP Coverage not approved	
	corticosteroids, immunosuppressants (e.g. azathioprine), etc?			
13.	Is the patient 18 years of age or older?	☐ Yes	□ No	
		proceed to question 27	STOP	
			Coverage not approved	
T	Has the patient received this medication under the TRICARE benefit in the last 6 months? Please choose "No" if the patient did not previously have a TRICARE approved PA for Rinvoq ER.	☐ Yes	□ No	
		(subject to verification)	proceed to question 16	
•	approved FA IOI MIIIVOY EK.	proceed to question 15		

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15. For atopic dermatitis, has the patient's disease severity	☐ Yes	□ No
improved and stabilized to warrant continued therapy?	(subject to verification)	STOP
	Sign and date below	Coverage not approved
16. Is the patient greater than or equal to 12 year(s) of age?	☐ Yes	□ No
	proceed to question 17	STOP
		Coverage not approved
17. Is the requested medication prescribed by a	□ Yes	□ No
dermatologist, allergist, or immunologist?	proceed to question 18	STOP
		Coverage not approved
18. Is the patient's disease adequately controlled with	☐ Yes	□ No
other systemic drug products including biologics (for	STOP	proceed to question 19
example, Dupixent)?	Coverage not approved	
19. Does the patient have a contraindication to,	☐ Yes	□ No
intolerability to, or have they failed treatment with ONE	proceed to question 20	STOP
medication in EACH of the following two categories:		Coverage not approved
Topical Corticosteroids AND		
NOTE:		
For patients 18 years of age or older, high potency/class 1 topical corticosteroids (for example, clobetasol propionate 0.05% ointment/cream, fluocinonide 0.05% ointment/cream) is required.		
For patients 12 to 17 years of age, can be any topical corticosteroid.		
 Topical calcineurin inhibitor (for example, pimecrolimus, tacrolimus) 		
20.Does the patient have a contraindication to,	☐ Yes	□ No
intolerability to, inability to access treatment, or has failed treatment with Narrowband UVB phototherapy?	proceed to question 27	STOP
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21. Does the patient have active non-radiographic axial	☐ Yes	□ No
spondyloarthritis?	proceed to question 22	STOP
	' '	Coverage not approved
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22. Is the patient 18 years of age or older?	☐ Yes	□ No
	proceed to question 23	STOP
		Coverage not approved
23. Has the patient experienced an inadequate response to	☐ Yes	□ No
Cosentyx?	proceed to question 26	proceed to question 24
24. Has the patient experienced an adverse reaction to	☐ Yes	□ No
Cosentyx that is not expected to occur with the requested agent?	proceed to question 26	proceed to question 25

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	25. Does the patient have a contraindication to Cosentyx?	☐ Yes	□ No
		proceed to question 26	STOP
			Coverage not approved
	26. Has the patient experienced an inadequate response to	☐ Yes	□ No
	at least TWO NSAIDs (for example: ibuprofen,	proceed to question 27	STOP
	naproxen, diclofenac) over a period of at least two months?		Coverage not approved
	27. Is the provider aware of the FDA safety alerts AND	☐ Yes	□ No
	Boxed Warnings?	proceed to question 28	STOP
			Coverage not approved
	28. Does the patient have a hemoglobin level LESS THAN	☐ Yes	□ No
	8 g/dL?	STOP	proceed to question 29
		Coverage not approved	
	29. Does the patient have an absolute neutrophil count	☐ Yes	□ No
	(ANC) LESS THAN 1,000/mm ³ ?	STOP	proceed to question 30
		Coverage not approved	
	30. Does the patient have an absolute lymphocyte count	☐ Yes	□ No
	(ALC) LESS THAN 500/mm ³ ?	STOP	proceed to question 31
		Coverage not approved	
	31. Will the patient be receiving other targeted	☐ Yes	□ No
	immunomodulatory biologics with Rinvoq ER, except for Otezla, including but not limited to the following:	STOP	proceed to question 32
	Actemra, Cimzia, Cosentyx, Enbrel, Humira, Ilumya,	Coverage not approved	
	Kevzara, Kineret, Olumiant, Orencia, Remicade, Rituxan, Siliq, Stelara, Taltz, Xeljanz or Xeljanz XR or		
	Tremfya and other potent immunosuppressant's (for		
	example: azathioprine, cyclosporine)? 32. Does the patient have a history of venous	☐ Yes	□ No
	thromboembolic (VTE) disease?	STOP	proceed to question 33
		Coverage not approved	proceed to question co
	33. Does the patient have evidence of an active TB	☐ Yes	□ No
	infection within the past 12 months?	STOP	Sign and date below
		Coverage not approved	
)	I certify the above is true to the best of my knowledge. Please sign and date:		
	Prescriber Signature	Date	
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