

# Ilaris

## **Prior Authorization Request**

Your patient's benefit plan requires prior authorization for certain medications. In order to make appropriate medical necessity determinations, your patient's diagnosis and other clinical information is required. Please complete the information requested on the form below and fax this form to Priority Partners, toll-free at 1-866-212-4756 to initiate the review process. If you have questions regarding the prior authorization please contact Priority Partners at 888-819-1043 Option 4.

Date:	
Patient's Date of Birth:	
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Criteria Questions:
What is the ICD-10 code?
<ol> <li>Will the requested drug be used in combination with any other biologic (e.g., Humira) or targeted synthetic drug (e.g., Olumiant, Otezla, Xeljanz)?</li> <li>Yes, Continue to 2</li> <li>No, Continue to 2</li> </ol>
<ul> <li>2. Has the patient ever received (including current utilizers) a biologic (e.g., Humira) or targeted synthetic drug (e.g., Olumiant, Xeljanz) associated with an increased risk of tuberculosis?</li> <li>☐ Yes, Continue to 6</li> <li>☐ No, Continue to 3</li> </ul>
3. Has the patient had a tuberculosis (TB) test (e.g., tuberculosis skin test [PPD], interferon-release assay [IGRA], chest x-ray) within 6 months of initiating therapy?  ☐ Yes, Continue to 4  ☐ No, Continue to 4
4. What were the results of the tuberculosis (TB) test?  Positive for TB (If checked, go to 5)  Negative for TB (If checked, go to 6)  Unknown (If checked, no further questions)
5. Which of the following applies to the patient?  Patient has latent TB and treatment for latent TB has been initiated (If checked, go to 6)  Patient has latent TB and treatment for latent TB has been completed (If checked, go to 6)  Patient has latent TB and treatment for latent TB has not been initiated (If checked, go to 6)  Patient has active TB (If checked, go to 6)
6. What is the patient's diagnosis?  Cryopyrin-Associated Periodic Syndromes (CAPS), including Familial Cold Autoinflammatory Syndrome (FCAS) and Muckle-Wells Syndrome (MWS) (If checked, go to 7)
Tumor Necrosis Factor Receptor Associated Periodic Syndrome (TRAPS) (If checked, go to 16)
☐ Hyperimmunoglobulin D Syndrome (HIDS)/Mevalonate Kinase Deficiency (MKD) (If checked, go to 24) ☐ Familial Mediterranean Fever (FMF) (If checked, go to 31)
Systemic juvenile idiopathic arthritis (sJIA) (If checked, go to 39)
Polyarticular juvenile idiopathic arthritis (pJIA) (If checked, <i>no further questions</i> )
Gout flares (If checked, go to 49)
☐Pseudogout (also known as calcium pyrophosphate deposition disease) flares (If checked, go to 49)
☐Adult-onset Still's disease (AOSD) (If checked, go to 58)
□Other, please specify. (If checked, no further questions)
7. Is the patient 4 years of age or older?  Tyes, Continue to 8

 $\square$  No, Continue to 8

8. Is the requested drug being prescribed by or in consultation with a rheumatologist or immunologist?  ☐ Yes, Continue to 9  ☐ No, Continue to 9
<ul> <li>9. Is this request for continuation of therapy with the requested drug?</li> <li>☐ Yes, Continue to 10</li> <li>☐ No, Continue to 12</li> </ul>
10. Is the patient currently receiving the requested drug through samples or a manufacturer's patient assistance program?  Tyes (If checked, go to 12)
□ No (If checked, go to 12) □ Unknown (If checked, go to 12)
11. 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?  Test, No Further Questions No, No Further Questions
12. What is the patient's diagnosis?
□Familial cold autoinflammatory syndrome (FCAS) (If checked, go to 13)
☐Muckle-Wells syndrome (MWS) (If checked, go to 14) ☐Other (If checked, no further questions)
13. Does the patient have classic signs and symptoms of familial cold autoinflammatory syndrome (FCAS) (i.e., recurrent, intermittent fever and rash that were often exacerbated by exposure to generalized cool ambient temperature)?  Temperature to 15 No, Continue to 15
14. Does the patient have classic signs and symptoms of Muckle-Wells syndrome (MWS) (i.e., chronic fever and rash of waxing and waning intensity, sometimes exacerbated by exposure to generalized cool ambient temperature)?  Yes, Continue to 15 No, Continue to 15
<ul> <li>15. Does the patient have functional impairment limiting the activities of daily living?</li> <li>☐ Yes, No Further Questions</li> <li>☐ No, No Further Questions</li> </ul>
<ul> <li>16. Is the requested drug being prescribed by or in consultation with a rheumatologist or immunologist?</li> <li>□ Yes, Continue to 17</li> <li>□ No, Continue to 17</li> </ul>
17. Is this request for continuation of therapy with the requested drug?  ☐ Yes, Continue to 18 ☐ No, Continue to 20

18. Is the patient currently receiving the requested drug through samples or a manufacturer's patient assistance program?
☐Yes (If checked, go to 20)
□No (If checked, go to 19) □Unknown (If checked, go to 20)
19. 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, <i>No Further Questions</i> ☐ No, <i>No Further Questions</i>
20. Does the patient have chronic or recurrent disease activity?  ☐ Yes, Continue to 21  ☐ No, Continue to 21
21. Has the patient had active flares within the last 6 months? <i>ACTION REQUIRED</i> : If Yes, please attach chart notes or medical record documentation indicating number of active flares within the last 6 months. <i>ACTION REQUIRED</i> : Submit supporting documentation ☐ Yes, <i>Continue to 22</i> ☐ No, <i>Continue to 22</i>
22. What is the patient's Physician's Global Assessment (PGA) score? Indicate score. <i>ACTION REQUIRED</i> : Please attach chart notes or medical record documentation indicating Physician's Global Assessment score.
□Less than 2 <i>ACTION REQUIRED</i> : Submit supporting documentation (If checked, go to 23)
□2 or greater <i>ACTION REQUIRED</i> : Submit supporting documentation (If checked, <i>no further questions</i> ) □Unknown (If checked, go to 23)
23. What is the patient's C-reactive protein (CRP) level in mg/L? Indicate in mg/L. <i>ACTION REQUIRED</i> : Please attach laboratory result indicating patient's C-reactive protein (CRP) level.
mg/L <i>ACTION REQUIRED</i> : Submit supporting documentation ( <i>No further questions</i> )
24. Is the requested drug being prescribed by or in consultation with a rheumatologist or immunologist? ☐ Yes, <i>Continue to 25</i> ☐ No, <i>Continue to 25</i>
25. Is this request for continuation of therapy with the requested drug?  ☐ Yes, Continue to 26  ☐ No, Continue to 28
26. Is the patient currently receiving the requested drug through samples or a manufacturer's patient assistance program?
☐Yes (If checked, go to 28)
□No (If checked, go to 27) □Unknown (If checked, go to 28)

27. 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, <i>No Further Questions</i> ☐ No, <i>No Further Questions</i>
28. Has the patient had active flares within the last 6 months? <i>ACTION REQUIRED</i> : If Yes, please attach char notes or medical record documentation indicating number of active flares within the last 6 months. <i>ACTION REQUIRED</i> : Submit supporting documentation  Yes, <i>Continue to 29</i> No, <i>Continue to 29</i>
29. What is the patient's Physician's Global Assessment (PGA) score? Indicate score. <i>ACTION REQUIRED</i> : Please attach chart notes or medical record documentation indicating Physician's Global Assessment score.
□Less than 2 <i>ACTION REQUIRED</i> : Submit supporting documentation (If checked, go to 30)
□ 2 or greater <i>ACTION REQUIRED</i> : Submit supporting documentation (If checked, <i>no further questions</i> ) □ Unknown (If checked, go to 30)
30. What is the patient's C-reactive protein (CRP) level in mg/L? Indicate in mg/L. <i>ACTION REQUIRED</i> : Please attach laboratory result indicating patient's C-reactive protein (CRP) level.
mg/L ACTION REQUIRED: Submit supporting documentation (No further questions)
31. Is the requested drug being prescribed by or in consultation with a rheumatologist or immunologist?  ☐ Yes, Continue to 32  ☐ No, Continue to 32
32. Is this request for continuation of therapy with the requested drug?  ☐ Yes, Continue to 33  ☐ No, Continue to 35
33. Is the patient currently receiving the requested drug through samples or a manufacturer's patient assistance program?
☐Yes (If checked, go to 35) ☐No (If checked, go to 34) ☐Unknown (If checked, go to 35)
34. 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 Further Questions ☐ No, No Further Questions
35. Does the patient have active disease with flares within the last 6 months? <i>ACTION REQUIRED</i> : If Yes, please attach chart notes or medical record documentation indicating number of active flares within the last 6 months. <i>ACTION REQUIRED</i> : Submit supporting documentation ☐ Yes, <i>Continue to 36</i> ☐ No, <i>Continue to 36</i>
36. What is the patient's C-reactive protein (CRP) level in mg/L? Indicate in mg/L. <i>ACTION REQUIRED</i> : Please attach laboratory result indicating patient's C-reactive protein (CRP) level.  mg/L <i>ACTION REQUIRED</i> : Submit supporting documentation (go to 37)

37. Has the patient had an inadequate response or intolerance to colchicine? <i>ACTION REQUIRED</i> : If Yes, please attach chart notes, medical record documentation, or claims history supporting previous medications tried, including response to therapy. <i>ACTION REQUIRED</i> : Submit supporting documentation ☐ Yes, <i>No Further Questions</i> ☐ No, <i>Continue to 38</i>			
38. Does the patient have a contraindication to colchicine? <i>ACTION REQUIRED</i> : If Yes, please attach documentation of clinical reason to avoid therapy. <i>ACTION REQUIRED</i> : Submit supporting documentation ☐ Yes, <i>No Further Questions</i> ☐ No, <i>No Further Questions</i>			
39. Is the patient 2 years of age or older?  ☐ Yes, Continue to 40  ☐ No, Continue to 40			
40. Is the requested drug being prescribed by or in consultation with a rheumatologist?  ☐ Yes, Continue to 41  ☐ No, Continue to 41			
41. Is this request for continuation of therapy with the requested drug?  ☐ Yes, Continue to 42  ☐ No, Continue to 45			
42. Is the patient currently receiving the requested drug through samples or a manufacturer's patient assistance program?			
☐ Yes (If checked, go to 45)			
□ No (If checked, go to 43) □ Unknown (If checked, go to 45)			
43. 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, Continue to 44  ☐ No, Continue to 44			
44. Which of the following has the patient experienced an improvement in from baseline? <i>ACTION REQUIRED</i> : Please attach chart notes or medical record documentation supporting a positive clinical response.  ☐ Number of joints with active arthritis (e.g., swelling, pain, limitation of motion) <i>ACTION REQUIRED</i> : Submit supporting documentation (If checked, <i>no further questions</i> )  ☐ Number of joints with limitation of movement <i>ACTION REQUIRED</i> : Submit supporting documentation (If checked, <i>no further questions</i> )  ☐ Functional ability <i>ACTION REQUIRED</i> : Submit supporting documentation (If checked, <i>no further questions</i> )  ☐ Systemic features (e.g., fevers, evanescent rash, lymphadenopathy, hepatomegaly, splenomegaly, or			
serositis) ACTION REQUIRED: Submit supporting documentation (If checked, no further questions)  None of the above (If checked, no further questions)			
<ul> <li>45. Has the patient been diagnosed with active systemic juvenile idiopathic arthritis (sJIA)?</li> <li>□ Yes, Continue to 46</li> <li>□ No, Continue to 46</li> </ul>			

Send completed form to: Priority Partners Fax: 1-866-212-4756

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46. Has the patient ever received or is currently receiving a biologic (e.g., Humira) indicated for the treatment of active systemic juvenile idiopathic arthritis (excluding receiving the drug via samples or a manufacturer's patient assistance program)? <i>ACTION REQUIRED</i> : If Yes, please attach chart notes, medical record documentation, or claims history supporting previous medications tried. <i>ACTION REQUIRED</i> : Submit supporting documentation Yes, <i>No Further Questions</i> □ No, <i>Continue to 47</i>	•
47. Does the patient have active systemic features (e.g., fever, evanescent rash, lymphadenopathy, hepatomegaly splenomegaly, or serositis)?  ☐ Yes, Continue to 48  ☐ No, Continue to 48	,
48. Has the patient had an inadequate response to non-steroidal anti-inflammatory drugs (NSAIDs) or systemic glucocorticoids? <i>ACTION REQUIRED</i> : If Yes, please attach chart notes, medical record documentation, or claims history supporting previous medications tried, including response to therapy. <i>ACTION REQUIRED</i> : Submit supporting documentation  Yes, <i>No Further Questions</i> No, <i>No Further Questions</i>	
<ul> <li>49. Is Ilaris being requested for the management of flares for gout or pseudogout (also known as calcium pyrophosphate deposition disease)?</li> <li>☐ Yes, Continue to 50</li> <li>☐ No, Continue to 50</li> </ul>	
50. Is the requested drug being prescribed by or in consultation with a rheumatologist?  ☐ Yes, Continue to 51  ☐ No, Continue to 51	
<ul> <li>51. Is this request for continuation of therapy with the requested drug?</li> <li>☐ Yes, Continue to 52</li> <li>☐ No, Continue to 54</li> </ul>	
<ul> <li>52. Is the patient currently receiving the requested drug through samples or a manufacturer's patient assistance program?</li> <li>Yes (If checked, go to 54)</li> <li>No (If checked, go to 53)</li> <li>Unknown (If checked, go to 54)</li> </ul>	
<ul> <li>53. 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?</li> <li>Yes, No Further Questions</li> <li>No, No Further Questions</li> </ul>	
54. Has the patient had an inadequate response or intolerance to maximum tolerated doses of non-steroidal anti-inflammatory drugs (NSAIDs) or has a contraindication to NSAIDs? <i>ACTION REQUIRED</i> : If Yes, please attac 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. <i>ACTION REQUIRED</i> : Submit supporting documentation	h

	Yes, Continue to 55 No, Continue to 55		
doc ther <b>RE</b>	Has the patient had an inadequate response or intolerance to maximum tolerated doses of colchicine or has a traindication to colchicine? <i>ACTION REQUIRED</i> : If Yes, please attach chart notes, medical record umentation, or claims history supporting previous medications tried (if applicable), including response to rapy. If therapy is not advisable, please attach documentation of clinical reason to avoid therapy. <i>ACTION QUIRED</i> : Submit supporting documentation  Yes, <i>Continue to 56</i> No, <i>Continue to 56</i>		
cort clai Sub	Has the patient had an inadequate response or intolerance to maximum tolerated doses of oral and injectable ticosteroids? <i>ACTION REQUIRED</i> : If Yes, please attach chart notes, medical record documentation, or ms history supporting previous medications tried, including response to therapy. <i>ACTION REQUIRED</i> : mit supporting documentation  Yes, <i>No Further Questions</i> No, <i>Continue to 57</i>		
57. Does the patient have a clinical reason to avoid repeated courses of corticosteroids? <i>ACTION REQUIRED</i> : If Yes, please attach documentation of clinical reason to avoid therapy. <i>ACTION REQUIRED</i> : Submit supporting documentation  ☐ Yes, <i>No Further Questions</i> ☐ No, <i>No Further Questions</i>			
	Is the patient an adult (18 years of age or older)? Yes, Continue to 59 No, Continue to 59		
59.	Is the requested drug being prescribed by or in consultation with a rheumatologist? Yes, <i>Continue to 60</i> No, <i>Continue to 60</i>		
60.	Is this request for continuation of therapy with the requested drug? Yes, Continue to 61 No, Continue to 64		
	Is the patient currently receiving the requested drug through samples or a manufacturer's patient assistance gram?  Yes (If checked, go to 64)  No (If checked, go to 62)  Unknown (If checked, go to 64)		
	Has the patient achieved or maintained a positive clinical response to treatment as evidenced by low disease vity or improvement in signs and symptoms since starting treatment with the requested drug? Yes, <i>Continue to 63</i> No, <i>Continue to 63</i>		
	Which of the following has the patient experienced an improvement in from baseline? <i>ACTION REQUIRED</i> : ase attach chart notes or medical record documentation supporting a positive clinical response.		

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Prescriber or Authorized Signature	Date (mm/dd/yy)
I attest that this information is accurate and true, information is available for review if requested by	
	drug (e.g., methotrexate)? <i>ACTION REQUIRED</i> : If Yes, on, or claims history supporting previous medications tried.
66. Does the patient have active systemic features (e.g lymphadenopathy, hepatomegaly, splenomegaly, or so ☐ Yes, <i>Continue to 67</i> ☐ No, <i>Continue to 67</i>	
64. Has the patient been diagnosed with active adult-o  ☐ Yes, Continue to 65  ☐ No, Continue to 65	nset Still's disease (AOSD)?
Submit supporting documentation (If checked, no furth  Number of joints with limitation of movement AC checked, no further questions)  Functional ability ACTION REQUIRED: Submit questions)	ction REQUIRED: Submit supporting documentation (If supporting documentation (If checked, no further supporting documentation), hepatomegaly, splenomegaly, or ocumentation (If checked, no further questions)
Number of joints with active arthritis (e.g. swelli	ng nain limitation of motion) ACTION PEOUIPED: