



7. Has the patient had a documented inadequate response or intolerable adverse event with any of the following preferred products? Please indicate ALL that apply. **ACTION REQUIRED: If Yes, attach supporting chart note(s).**
- |                                                  |                                              |                                                    |
|--------------------------------------------------|----------------------------------------------|----------------------------------------------------|
| <input type="checkbox"/> Enbrel:                 | <input type="checkbox"/> Inadequate response | <input type="checkbox"/> Intolerable adverse event |
| <input type="checkbox"/> Humira:                 | <input type="checkbox"/> Inadequate response | <input type="checkbox"/> Intolerable adverse event |
| <input type="checkbox"/> Kevzara:                | <input type="checkbox"/> Inadequate response | <input type="checkbox"/> Intolerable adverse event |
| <input type="checkbox"/> Orenzia (SC/ClickJect): | <input type="checkbox"/> Inadequate response | <input type="checkbox"/> Intolerable adverse event |
| <input type="checkbox"/> No - none of the above  |                                              |                                                    |
8. Does the patient have one of the following documented clinical reasons to avoid Enbrel and Humira?  
**ACTION REQUIRED: If Yes, attach supporting chart note(s).**
- |                                                                                                                       |                                                                           |
|-----------------------------------------------------------------------------------------------------------------------|---------------------------------------------------------------------------|
| <input type="checkbox"/> Yes - History of demyelinating disorder                                                      | <input type="checkbox"/> Yes - Autoantibody formation/lupus-like syndrome |
| <input type="checkbox"/> Yes - History of congestive heart failure                                                    | <input type="checkbox"/> Yes - Risk of lymphoma                           |
| <input type="checkbox"/> Yes - History of hepatitis B virus infection <input type="checkbox"/> No - none of the above |                                                                           |

**Section B: All Requests**

9. Is this request for continuation of therapy?  Yes  No *If No, skip to #13*
10. Is the patient currently receiving Actemra through samples or a manufacturer's patient assistance program?  Yes  No  Unknown *If Yes or Unknown, skip to #13*
11. How long has the patient been receiving the requested medication? \_\_\_\_\_ months  
*If less than 3 months, no further questions.*
12. Has the patient achieved or maintained positive clinical response to treatment as evidenced by low disease activity or improvement in signs and symptoms? *If Yes, no further questions*  Yes  No
13. Has the patient received any of the following medications?  
*If Yes, please indicate the most recent medication and skip to diagnosis section.*
- |                                                                                                                                   |                                    |                                  |                                 |                                    |                                       |                                  |                                  |
|-----------------------------------------------------------------------------------------------------------------------------------|------------------------------------|----------------------------------|---------------------------------|------------------------------------|---------------------------------------|----------------------------------|----------------------------------|
| <input type="checkbox"/> Cimzia                                                                                                   | <input type="checkbox"/> Cosentyx  | <input type="checkbox"/> Enbrel  | <input type="checkbox"/> Humira | <input type="checkbox"/> Inflectra | <input type="checkbox"/> Kevzara      | <input type="checkbox"/> Kineret | <input type="checkbox"/> Orenzia |
| <input type="checkbox"/> Remicade                                                                                                 | <input type="checkbox"/> Renflexis | <input type="checkbox"/> Rituxan | <input type="checkbox"/> Siliq  | <input type="checkbox"/> Simponi   | <input type="checkbox"/> Simponi Aria | <input type="checkbox"/> Stelara | <input type="checkbox"/> Taltz   |
| <input type="checkbox"/> Tremfya <input type="checkbox"/> Xeljanz <input type="checkbox"/> Xeljanz XR <input type="checkbox"/> No |                                    |                                  |                                 |                                    |                                       |                                  |                                  |
14. Has the patient undergone pretreatment screening for latent tuberculosis (TB) infection with either a TB skin test or an interferon gamma release assay (e.g., QFT-GIT, T-SPOT.TB)?  Yes  No

**Complete the following section based on the patient's diagnosis, if applicable.**

**Section C: Rheumatoid Arthritis**

15. Has the patient experienced an inadequate response after at least 3 months of treatment with the methotrexate dose greater than or equal to 20 mg per week? *If Yes, no further questions*  Yes  No
16. Has the patient experienced intolerance to methotrexate? *If Yes, no further questions*  Yes  No
17. Does the patient have a contraindication to methotrexate?  Yes  No  
***If Yes, indicate the contraindication:*** \_\_\_\_\_

**Section D: Polyarticular Juvenile Idiopathic Arthritis**

18. Has the patient experienced an inadequate response to a tumor necrosis factor (TNF) inhibitor (e.g., Enbrel, Humira, or Remicade) after at least 3 months of treatment?  
*If Yes, no further questions*  Yes  No
19. Has the patient experienced an intolerable adverse event to a tumor necrosis factor (TNF) inhibitor (e.g., Enbrel, Humira, or Remicade)? *If Yes, no further questions*  Yes  No
20. Does the patient have contraindication to a tumor necrosis factor (TNF) inhibitors (e.g., Enbrel, Humira, or Remicade)?  Yes  No

**Section E: Systemic Juvenile Idiopathic Arthritis**

21. Has the patient experienced an inadequate response to ANY of the following?
- |                                                                                                                   |
|-------------------------------------------------------------------------------------------------------------------|
| <input type="checkbox"/> At least 2 weeks of treatment with corticosteroids (e.g. prednisone, methylprednisolone) |
| <input type="checkbox"/> At least 3 months of treatment with methotrexate                                         |

- At least 3 months of treatment with leflunomide
- No – No history of an inadequate response to any of the above

*I attest that this information is accurate and true, and that documentation supporting this information is available for review if requested by CVS Caremark or the benefit plan sponsor.*

X \_\_\_\_\_  
**Prescriber or Authorized Signature** **Date (mm/dd/yy)**