



## Orencia

### Prior Authorization Request

CVS Caremark administers the prescription benefit plan for the patient identified. This patient's benefit plan requires prior authorization for certain medications in order for the drug to be covered. To make an appropriate determination, providing the most accurate diagnosis for the use of the prescribed medication is necessary. **Please respond below and fax this form to CVS Caremark toll-free at 1-855-330-1720.** If you have questions regarding the prior authorization, please contact CVS Caremark at **1-888-877-0518**. For inquiries or questions related to the patient's eligibility, drug copay or medication delivery; please contact the Specialty Customer Care Team: CaremarkConnect® 1-800-237-2767.

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**Patient's Name:** \_\_\_\_\_ **Date:** \_\_\_\_\_  
**Patient's ID:** \_\_\_\_\_ **Patient's Date of Birth:** \_\_\_\_\_  
**Physician's Name:** \_\_\_\_\_  
**Specialty:** \_\_\_\_\_ **NPI#:** \_\_\_\_\_  
**Physician Office Telephone:** \_\_\_\_\_ **Physician Office Fax:** \_\_\_\_\_

**Referring Provider Info:**  Same as Requesting Provider

**Name:** \_\_\_\_\_ **NPI#:** \_\_\_\_\_  
**Fax:** \_\_\_\_\_ **Phone:** \_\_\_\_\_

**Rendering Provider Info:**  Same as Referring Provider  Same as Requesting Provider

**Name:** \_\_\_\_\_ **NPI#:** \_\_\_\_\_  
**Fax:** \_\_\_\_\_ **Phone:** \_\_\_\_\_

*Approvals may be subject to dosing limits in accordance with FDA-approved labeling, accepted compendia, and/or evidence-based practice guidelines.*

**Required Demographic Information:**

*Patient Weight:* \_\_\_\_\_ kg

*Patient Height:* \_\_\_\_\_ cm

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**Exception Criteria Questions:**

A. Is the product being requested for the treatment of an ADULT patient (18 years of age or older) with one of the following indications?

- Ankylosing spondylitis
- Crohn's disease
- Plaque psoriasis
- Psoriatic arthritis
- Rheumatoid arthritis
- Ulcerative colitis

Yes  No *If No, skip to Site of Service Questions*

B. These are the preferred products for which coverage is provided for treatment of the following indications:

- Ankylosing spondylitis, psoriatic arthritis, rheumatoid arthritis: **Remicade and Simponi Aria**
- Plaque psoriasis: **Ilumya and Remicade**
- Crohn's disease, ulcerative colitis: **Entyvio and Remicade**
- **Stelara IV** is indicated for a one time induction dose for Crohn's disease and ulcerative colitis.

Can the patient's treatment be switched to a preferred product?

Yes, *Please obtain Form for preferred product and submit for corresponding PA.*

No

*If diagnosis is Plaque psoriasis, skip to Question K*

C. Is this request for continuation of therapy with the requested product?  Yes  No, *If No, skip to Question E*

D. Is the patient currently receiving the requested product through samples or a manufacturer's patient assistance program? If unknown, answer Yes.  Yes  No, *skip to Site of Service Questions*

E. What is the diagnosis?

Ankylosing spondylitis

Crohn's disease, *skip to Question H*

Psoriatic arthritis

Rheumatoid arthritis

Ulcerative colitis, *skip to Question H*

Other, *skip to Site of Service Questions*

F. Does the patient have a documented inadequate response or intolerable adverse event to both of the preferred products (Remicade, Simponi Aria)? **Action Required: If 'Yes', attach supporting chart note(s).**

Yes, *skip to Site of Service Questions*  No

G. Does the patient have one of the following documented clinical reasons to avoid the preferred products that are TNF inhibitors (Remicade, Simponi Aria)? **Action Required: If 'Yes', attach supporting chart note(s).**

Not applicable – Requested medical is a TNF inhibitor *skip to Site of Service Questions*

Yes – History of demyelinating disorder, *skip to Site of Service Questions*

Yes – History of congestive heart failure *skip to Site of Service Questions*

Yes – History of hepatitis B virus infection *skip to Site of Service Questions*

Yes – Autoantibody formation/lupus-like syndrome (attributed to TNF inhibitor) *skip to Site of Service Questions*

Yes – Risk of lymphoma *skip to Site of Service Questions*

No – None of the above *skip to Site of Service Questions*

H. Does the patient have a documented inadequate response or intolerable adverse event to both of the preferred products (Entyvio, Remicade)? **Action Required: If 'Yes', attach supporting chart note(s).** *If Yes, skip to Site of Service Questions*  Yes  No

I. Does the patient have one of the following documented clinical reasons to avoid the preferred product that is a TNF inhibitor (Remicade)? **Action Required: If 'Yes', attach supporting chart note(s).**

Not applicable – requested medication is a TNF inhibitor

Yes – History of demyelinating disorder

Yes – History of congestive heart failure

Yes – History of hepatitis B virus infection

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- Yes – Autoantibody formation/lupus-like syndrome (attributed to TNF inhibitor)
  - Yes – Risk of lymphoma
  - No- None of the above
- J. Does the patient have a documented inadequate response or intolerable adverse event to the preferred product that is not a TNF inhibitor (Entyvio)? **Action Required: If 'Yes', attach supporting chart note(s).**  
 Yes  No *For yes or no, skip to Site of Service Questions*
- K. Does the patient have a documented inadequate response or intolerable adverse event to the preferred products indicated for plaque psoriasis (Ilumya, Remicade)? **Action Required: If 'Yes', attach supporting chart note(s).**  
*If Yes, skip to Site of Service Questions*  Yes  No
- L. Does the patient have one of the following documented clinical reasons to avoid the preferred product that is a TNF inhibitor (Remicade)? **Action Required: If 'Yes', attach supporting chart note(s).**  
 Not applicable – requested medication is a TNF inhibitor  
 Yes – History of demyelinating disorder  
 Yes – History of congestive heart failure  
 Yes – History of hepatitis B virus infection  
 Yes – Autoantibody formation/lupus-like syndrome (attributed to TNF inhibitor)  
 Yes – Risk of lymphoma  
 No- None of the above
- M. Does the patient have a documented inadequate response or intolerable adverse event to the preferred product that is not a TNF inhibitor (Ilumya)? **Action Required: If 'Yes', attach supporting chart note(s).**  Yes  No

**Site of Service Questions (SOS):**

- A. Where will this drug be administered?
- |   |   |
|---|---|
| <input type="checkbox"/> Ambulatory surgical, <i>skip to Clinical Questions</i> | <input type="checkbox"/> Home infusion, <i>skip to Clinical Questions</i> |
| <input type="checkbox"/> Off-campus Outpatient Hospital                         | <input type="checkbox"/> On-campus Outpatient Hospital                    |
| <input type="checkbox"/> Physician office, <i>skip to Clinical Questions</i>    | <input type="checkbox"/> Pharmacy, <i>skip to Clinical Questions</i>      |
- B. Is the patient less than 21 years of age or 65 years of age or older?  
 Yes – less than 21 years old  
 Yes – age 65 years or older, *skip to Clinical Criteria Questions*  
 No, *Skip to Question D.*
- C. Is this request to continue previously established treatment with the requested medication?  
 Yes – This is a continuation of an existing treatment  
 No – This is a new therapy request (patient has not received requested medication in the last 6 months) *skip to Clinical Criteria Questions*
- D. Has the patient experienced an adverse event with the requested product that has not responded to conventional interventions (eg acetaminophen, steroids, diphenhydramine, fluids, or other pre- medications) or a severe adverse event (anaphylaxis, anaphylactoid reactions, myocardial infarction, thromboembolism, or seizures) during or immediately after an infusion? **ACTION REQUIRED: If Yes, please attach supporting clinical documentation.**  
 Yes, *skip to Clinical Criteria Questions*  No
- E. Is the patient medically unstable which may include respiratory, cardiovascular, or renal conditions that may limit the member's ability to tolerate a large volume or load or predispose the member to a severe adverse event that cannot be managed in an alternate setting without appropriate medical personnel and equipment?  
**ACTION REQUIRED: If Yes, please attach supporting clinical documentation.**  
 Yes, *skip to Clinical Criteria Questions*  No
- F. Does the patient have severe venous access issues that require the use of special interventions only available in the outpatient hospital setting? **ACTION REQUIRED: If Yes, please attach supporting clinical documentation.**  
 Yes, *skip to Clinical Criteria Questions*  No

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- G. Does the patient have significant behavioral issues and/or physical or cognitive impairment that would impact the safety of the infusion therapy AND the patient does not have access to a caregiver? **ACTION REQUIRED: If Yes, please attach supporting clinical documentation. Indicate and continue to Clinical Criteria Questions**  
 Yes  No

**Clinical Criteria Questions:**

1. What is the prescribed quantity and frequency?
  - a) **Loading dose:**
    - Orencia IV 250 mg    Quantity and frequency: \_\_
    - Orencia SQ 125 mg    Quantity and frequency: \_\_
    - Orencia SQ 87.5 mg    Quantity and frequency: \_\_
    - Orencia SQ 50 mg    Quantity and frequency: \_\_
    - Other \_\_\_\_\_
  - b) **Maintenance dose:**
    - Orencia IV 250 mg    Quantity and frequency: \_\_
    - Orencia SQ 125 mg    Quantity and frequency: \_\_
    - Orencia SQ 87.5 mg    Quantity and frequency: \_\_
    - Orencia SQ 50 mg    Quantity and frequency: \_\_
    - Other \_\_\_\_\_
2. Has the patient been diagnosed with any of the following?
  - Moderately to severely active rheumatoid arthritis (RA)
  - Moderately to severely active **polyarticular** juvenile idiopathic arthritis (pJIA)
  - Moderately to severely active **oligoarticular** juvenile idiopathic arthritis
  - Active psoriatic arthritis (PsA)
  - Chronic graft versus host disease
  - Immune checkpoint inhibitor-related toxicity
  - Systemic juvenile idiopathic arthritis (sJIA)
  - Other \_\_\_\_\_
3. What is the ICD-10 code? \_\_\_\_
4. What is the patient's weight? \_\_Kg

**Section A: All Requests**

5. Will the requested drug be used in combination with any other biologic (e.g. Humira) or targeted synthetic disease-modifying antirheumatic drug (DMARD) (e.g., Olumiant, Otezla, Xeljanz)?  Yes  No
6. Has the patient ever received (including current utilizers) a biologic (e.g., Humira) or targeted synthetic DMARD (e.g., Olumiant, Xeljanz) associated with an increased risk of tuberculosis? *If Yes, skip to #8*  Yes  No
7. 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? *If Yes, skip to #10*  Yes  No
8. Does the patient have risk factors for tuberculosis (TB) (e.g., persons with close contact to people with infectious TB disease; persons who have recently immigrated from areas of the world with high rates of TB [e.g., Africa, Asia, Eastern Europe, Latin America, Russia]; children less than 5 years of age who have a positive TB test; groups with high rates of TB transmission [e.g., homeless persons, injection drug users, persons with HIV infection], or persons who work or reside with people who are at an increased risk for active TB [e.g., hospitals, long-term care facilities, correctional facilities, homeless shelters])?  Yes  No *If No, skip to #13*
9. Has the patient been tested for tuberculosis (TB) within the previous 12 months?  Yes  No
10. What were the results of the tuberculosis (TB) test?
  - Positive for TB
  - Negative for TB, *skip to #13*
  - Unknown
11. Does the patient have latent or active tuberculosis (TB)?  Latent  Active  Unknown

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12. Has treatment for latent tuberculosis (TB) infection been initiated or completed?  
 Yes - treatment initiated    Yes - treatment completed    No
13. Is the patient currently receiving Orencia?    Yes    No  
*If diagnosis is chronic graft versus host disease, skip to Section F*
14. Is this request for continuation of therapy with the requested drug?  
 Yes    No   *If No, skip to diagnosis section.*
15. Is the patient currently receiving the requested drug through samples or a manufacturer's patient assistance program? *If Yes or Unknown, skip to diagnosis section.*    Yes    No    Unknown
16. Has the patient achieved or maintained positive clinical response as evidenced by low disease activity or improvement in signs and symptoms since starting treatment with the requested drug?  
 Yes    No

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

Section B: Rheumatoid Arthritis

*Continuation*

17. What is the percent of disease activity improvement from baseline in tender joint count, swollen joint count, pain, or disability? ***ACTION REQUIRED: Please attach chart notes or medical record documentation supporting positive clinical response.***    Yes    No   *No further questions*

*Initiation*

18. Has the patient ever received (including current utilizers) a biologic (e.g., Humira) or targeted synthetic DMARD (e.g., Rinvoq, Xeljanz) that is indicated for moderately to severely active rheumatoid arthritis?  
***ACTION REQUIRED: If Yes, please attach chart notes, medical record documentation, or claims history supporting previous medications tried and no further questions.***    Yes    No
19. Does the patient meet BOTH of the following: a) the patient was tested for the rheumatoid factor (RF) biomarker AND b) the RF biomarker test was positive? ***ACTION REQUIRED: If Yes, please attach laboratory results, chart notes, or medical record documentation of biomarker testing and skip to #26.***    Yes    No
20. Does the patient meet BOTH of the following: a) the patient was tested for the anti-cyclic citrullinated peptide (anti-CCP) biomarker AND b) 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 #26.***  
 Yes    No
21. Has the patient been tested for the rheumatoid factor (RF) biomarker? ***ACTION REQUIRED: If Yes, please attach laboratory results, chart notes, or medical record documentation of biomarker testing.***    Yes    No
22. Has the patient been tested for the anti-cyclic citrullinated peptide (anti-CCP) biomarker?  
***ACTION REQUIRED: If Yes, please attach laboratory results, chart notes, or medical record documentation of biomarker testing.***    Yes    No
23. Has the patient been tested for the C-reactive protein (CRP) and/or erythrocyte sedimentation rate (ESR) biomarker(s)? ***ACTION REQUIRED: If Yes, please attach laboratory results, chart notes, or medical record documentation of biomarker testing.***    Yes    No
24. Please indicate if the patient tested positive or negative for the C-reactive protein (CRP) biomarker, or if the test was not completed.    Positive for CRP    Negative for CRP    Test for CRP was not completed
25. Please indicate if the patient tested positive or negative for the erythrocyte sedimentation rate (ESR) biomarker, or if the test was not completed.    Positive for ESR    Negative for ESR    Test for ESR was not completed

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26. Has the patient experienced an inadequate response after at least 3 months of treatment with methotrexate at a dose greater than or equal to 20 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
27. 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.**  Yes  No
28. Does the patient have a contraindication to methotrexate? **ACTION REQUIRED: If Yes, please attach documentation of clinical reason to avoid therapy and indicate the contraindication.**  
 Yes  No **If Yes, please indicate the contraindication: \_\_\_\_\_**

### Section C: Articular Juvenile Idiopathic Arthritis

#### *Continuation*

29. Which of the following has the patient experienced an improvement in from baseline?  
**ACTION REQUIRED: Please attach chart notes or medical record documentation supporting positive clinical response. Indicate ALL that apply and no further questions.**
- Number of joints with active arthritis (e.g., swelling, pain, limitation of motion)  
 Functional ability  
 Number of joints with limitation of movement  
 None of the above

#### *Initiation*

30. Has the patient ever received (including current utilizers) a biologic (e.g., Humira) or targeted synthetic disease-modifying antirheumatic drug (DMARD) indicated for moderately to severely active articular juvenile idiopathic arthritis? **ACTION REQUIRED: If Yes, please attach chart notes, medical record documentation, or claims history supporting previous medications tried and no further questions.**  Yes  No
31. Has the patient had an inadequate response to methotrexate or another non-biologic DMARD 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.**  Yes  No
32. Does the patient have any of the following risk factors?  
 Positive rheumatoid factor  Pre-existing joint damage  
 Positive anti-cyclic citrullinated peptide antibodies  None of the above
33. Does the patient meet any of the following?  
 High-risk joints are involved (e.g., cervical spine, wrist, or hip)  High disease activity  
 High risk for disabling joint disease  None of the above

### Section D: Psoriatic Arthritis

#### *Continuation*

34. Which of the following has the patient experienced an improvement in from baseline?  
**ACTION REQUIRED: Please attach chart notes or medical record documentation supporting positive clinical response. Indicate ALL that apply and no further questions.**
- Number of swollen joints  Enthesitis  Number of tender joints  
 Skin and/or nail involvement  Dactylitis  None of the above

### Section E: Chronic Graft Versus Host Disease

35. 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  No
36. Does the patient have an intolerance or contraindication to corticosteroids? **ACTION REQUIRED: If Yes, please attach documentation of clinical reason to avoid therapy.**  Yes  No

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Section F: Immune Checkpoint Inhibitor-Related Toxicity

37. Does the patient have cardiac toxicity?  Yes  No

<b>Step Therapy Override: Complete if Applicable for the state of Maryland.</b>	Please Circle	
Is the requested drug being used to treat stage four advanced metastatic cancer?	Yes	No
Is the requested drug's use consistent with the FDA-approved indication or the National Comprehensive Cancer Network Drugs & Biologics Compendium indication for the treatment of stage four advanced metastatic cancer and is supported by peer-reviewed medical literature?	Yes	No
Is the requested drug being used for an FDA-approved indication OR an indication supported in the compendia of current literature (examples: AHFS, Lexicomp, Clinical Pharmacology, Micromedex, current accepted guidelines)?	Yes	No
Does the prescribed quantity fall within the manufacturer's published dosing guidelines or within dosing guidelines found in the compendia of current literature (examples: package insert, AHFS, Lexicomp, Clinical Pharmacology, Micromedex, current accepted guidelines)?	Yes	No
Do patient chart notes document the requested drug was ordered with a paid claim at the pharmacy, the pharmacy filled the prescription and delivered to the patient or other documentation that the requested drug was prescribed for the patient in the last 180 days?	Yes	No
Has the prescriber provided proof documented in the patient chart notes that in their opinion the requested drug is effective for the patient's condition?	Yes	No

<b>Step Therapy Override: Complete if Applicable for the state of Virginia.</b>	Please Circle	
Is the requested drug being used for an FDA-approved indication or an indication supported in the compendia of current literature (examples: AHFS, Micromedex, current accepted guidelines)?	Yes	No
Does the prescribed dose and quantity fall within the FDA-approved labeling or within dosing guidelines found in the compendia of current literature?	Yes	No
Is the request for a brand drug that has an AB-rated generic equivalent or interchangeable biological product available?	Yes	No
Has the patient had a trial and failure of the AB-rated generic equivalent or interchangeable biological product due to an adverse event (examples: rash, nausea, vomiting, anaphylaxis) that is thought to be due to an inactive ingredient?	Yes	No
Is the preferred drug contraindicated?	Yes	No
Is the preferred drug expected to be ineffective based on the known clinical characteristics of the patient and the prescription drug regimen?	Yes	No
Has the patient tried the preferred drug while on their current or previous health benefit plan and it was discontinued due to lack of efficacy or effectiveness, diminished effect, or an adverse event?	Yes	No
Is the patient currently receiving a positive therapeutic outcome with the requested drug for their medical condition?	Yes	No

*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)**

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