Blue Cross Blue Shield/Blue Care Network of Michigan Medication Authorization Request Form



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This form is to be used by participating physicians to obtain coverage for **drugs covered under the medical benefit**. For <u>commercial members only</u>, please complete this form and submit via fax to 1-877-325-5979. If you have any questions regarding this process, please contact BCBSM Provider Relations and Servicing or the Medical Drug Helpdesk at 1-800-437-3803 for assistance.

PATIENT INFORMATION	PHYSICIAN INFORMATION
Name	Name
ID Number	Specialty
D.O.B. $\frac{///}{\Box}$ MM/DD/YYYY	Address
Diagnosis	City /State/Zip
Drug Name Simponi Aria	Phone: Fax:
Dose and Quantity	NPI
Directions	Contact Person
Date of Service(s)	Contact Person
	Phone / Ext.

STEP 1: DISEASE STATE INFORMATION

Required Demographic Information:

 Patient Weight:
 kg

 Patient Height:
 ft

Will the provider be administering the medication to the FEP member within the health plan's geographic service area? □ Yes □ No If No, a prior authorization is not required through this process.

Prior authorizations are required for FEP members that will be serviced by a provider within the health plan's geographic service area. If you are not a provider in the geographic service area, please contact the health plan for questions regarding the FEP member's benefit requirements.

Is this member's FEP coverage primary or secondary coverage?

□ If primary, continue with question set.

□ If secondary, an authorization is not needed through this process. Please contact the member's primary coverage for determination of benefit and additional information.

Site of Care:

A. At what location will the member be receiving the requested medication?

Depresent Physician's office, home infusion, non-hospital affiliated ambulatory infusion center.

Outpatient hospital infusion center. Please provide the name of the infusion center and rationale why the patient must receive this medication in a hospital outpatient setting.

Other. Please specify.

Criteria Questions:

- 1. Has the patient been on Simponi ARIA therapy continuously for the last 6 months, excluding samples? Please select answer below:
 - **YES** this is a PA renewal for **CONTINUATION** of therapy, please answer the questions on **Continuation Section**.:
 - **NO** this is **INITIATION** of therapy, please answer the following questions:
- 2. Has the patient been tested for latent tuberculosis (TB)? □Yes* □No
 If YES, was the result of the test positive or negative for TB infection? □Negative □Positive
 *If POSITIVE, has the patient completed treatment or is the patient currently receiving treatment for latent TB? □Yes □No
- 3. Is the patient at risk for Hepatitis B Virus (HBV) infection? □Yes* □No
 *If YES, has HBV infection been ruled out or has the patient already started treatment for HBV infection? □Yes □No
- 4. Does the patient have any active infections including tuberculosis (TB) or hepatitis B virus (HBV)? **U**Yes **U**No
- 5. Will the patient be given live vaccines while on Simponi Aria therapy? **U**Yes **D**No
- 6. Will Simponi Aria be used in combination with another biologic DMARD or targeted synthetic DMARD? □Yes* □No **If YES*, please specify medication: _____
- 7. What is the patient's diagnosis?
 - □ Ankylosing Spondylitis (AS) (axial spondyloarthritis)
 - a. Does the patient have active ankylosing spondylitis? Yes No
 - b. Has the patient had either an inadequate response or intolerance to at least two different NSAIDs over a four-week period in total at maximum recommended or tolerated dose? \Box Yes \Box No*
 - *If NO, does the patient have a contraindication to NSAIDs? \Box Yes \Box No
 - c. Does the prescriber agree not to exceed the FDA labeled maintenance dose of 2 mg/kg IV every 8 weeks? Yes No
 - Delyarticular Juvenile Idiopathic Arthritis (pJIA)
 - a. Does the patient have an intolerance or contraindication or have they had an inadequate treatment response to a 3-month trial of at least one conventional disease-modifying antirheumatic drug (DMARD)? □Yes □No
 - b. Does the prescriber agree to administer Simponi Aria within the FDA labeled maintenance dose of 80 mg/m² (based on body surface area) every eight weeks? □Yes □No
 - D Psoriatic Arthritis (PsA)
 - a. Does the patient have active psoriatic arthritis (PsA)? □Yes □No
 - b. Does the patient have an intolerance or contraindication or have they had an inadequate treatment response to a 3-month trial of at least one conventional disease-modifying antirheumatic drug (DMARD)? \Box Yes \Box No
 - c. Does the prescriber agree not to exceed the FDA labeled maintenance dose of 80 mg/m2 (based on body surface area) every 8 weeks? □Yes □No
 - **C** Rheumatoid Arthritis (RA)
 - a. Does the patient have moderate to severely active rheumatoid arthritis? \Box Yes \Box No
 - b. Does the patient have an intolerance or contraindication or have they had an inadequate treatment response to a three-month trial of at least one conventional DMARD? \Box Yes \Box No
 - c. Does the patient have an intolerance or contraindication to methotrexate (MTX)? \Box Yes **If NO*, will Simponi Aria be used in combination with methotrexate? \Box Yes \Box No
 - d. Does the prescriber agree not to exceed the FDA labeled maintenance dose of 2 mg/kg IV every 8 weeks? Yes INo
 - Other diagnosis (*please specify*):

CONTINUATION OF THERAPY (PA RENEWAL) Simponi Aria (golimumab)

1. Has the patient been on Simponi ARIA therapy continuously for the last 6 months, excluding samples? Answer below: **NO** – this is **INITIATION** of therapy, please answer the questions on **Initiation Section**. **YES** – this is a PA renewal for **CONTINUATION** of therapy, please answer the following questions: 2. What is the patient's diagnosis? Ankylosing Spondylitis (AS) (axial spondyloarthritis) a. Does the prescriber agree not to exceed the FDA labeled maintenance dose of 2 mg/kg IV every 8 weeks? \Box Yes \Box No Polyarticular Juvenile Idiopathic Arthritis (pJIA) a. Does the prescriber agree not to exceed the FDA labeled maintenance dose of 80 mg/m2 (based on body surface area) every 8 weeks? **\U**Yes **U**No □ Psoriatic Arthritis (PsA) a. Does the prescriber agree not to exceed the FDA labeled maintenance dose of 2 mg/kg IV every 8 weeks? \Box Yes \Box No Rheumatoid Arthritis (RA) a. Does the patient have an intolerance or contraindication to methotrexate (MTX)? □No* *If NO, will Simponi Aria be used in combination with methotrexate? Yes No b. Does the prescriber agree not to exceed the FDA labeled maintenance dose of 2 mg/kg IV every 8 weeks? □Yes □No □ Other diagnosis (*please specify*): 3. Has the patient's condition improved or stabilized? □Yes □No 4. Does the patient have any active infections including tuberculosis (TB) or Hepatitis B Virus (HBV) infection? □Yes **D**No 5. Will the patient be given live vaccines while on Simponi Aria therapy? **\Box** Yes **\Box** No 6. Will Simponi Aria be used in combination with another biologic disease-modifying antirheumatic drug (DMARD) or targeted synthetic DMARD? **U**Yes* DNo

**If YES*, please specify medication:

*DMARDs: Actemra, Avsola, Cimzia, Cosentyx, Enbrel, Entyvio, Humira, Ilumya, Inflectra, Kevzara, Kineret, Olumiant, Orencia, Otezla, Remicade, Renflexis, Riabni, Rinvoq, Rituxan, Ruxience, Siliq, Simponi, Skyrizi, Stelara, Taltz, Tremfya, Truxima, and Xeljanz/ Xeljanz XR

Chart notes are required for the processing of all requests. Please add any other supporting medical information necessary for our review (required) Coverage will not be provided if the prescribing physician's signature and date are not reflected on this document.

Request for expedited review: I certify that applying the standard review time frame may seriously jeopardize the life or health of the member or the member's ability to regain maximum function

Physician's Nan	ne Physician Signature	Date
Step 2: Checklist	 Form Completely Filled Out Provide chart notes 	Attach test results
Step 3: Submit	By Fax: BCBSM Specialty Pharmacy Mailbox 1-877-325-5979	By Mail: BCBSM Specialty Pharmacy Program P.O. Box 312320, Detroit, MI 48231-2320