



# Simponi Aria Order Form

Patient Name: \_\_\_\_\_ DOB: \_\_\_\_\_  
 Phone: \_\_\_\_\_ Address: \_\_\_\_\_  
 City: \_\_\_\_\_ State: \_\_\_\_\_ Zip: \_\_\_\_\_ Email: \_\_\_\_\_  
 Sex: \_\_\_\_\_ Height: \_\_\_\_\_ Weight: \_\_\_\_\_ Allergies: \_\_\_\_\_

### DIAGNOSIS:

- Rheumatoid Arthritis ICD-10: \_\_\_\_\_
- Rheumatoid Arthritis with Rheumatoid Factor ICD-10: M05.A
- Psoriatic Arthritis ICD-10: \_\_\_\_\_
- Ankylosing Spondylitis ICD-10: \_\_\_\_\_
- Other: \_\_\_\_\_ ICD-10: \_\_\_\_\_

### ORDER FOR SIMPONI ARIA (GOLIMUMAB):

- Initial Dose: 2mg/kg IV at weeks 0, 4, and then every 8 weeks x 1 year**
- 2mg/kg IV every 8 weeks x 1 year**
- Other Dose: \_\_\_\_\_ Frequency: \_\_\_\_\_ x 1 year

### PRE-MEDICATIONS:

- Pre-Medications may be PRN (as needed)
- Acetaminophen 650mg PO
- Diphenhydramine 25mg PO or IV or Zyrtec 10 mg PO
- Hydrocortisone 100mg IV or Methylprednisolone 125mg IV
- Additional Pre-Medications: \_\_\_\_\_

### MAY ADMINISTER IF NEEDED FOR ALLERGIC REACTION:

- Nevada Infusion Hypersensitivity Reaction Order Set**
- Other: \_\_\_\_\_

ACCESS: Peripheral IV, Port, Midline, or PICC line

FLUSHING: 10 mls NS pre/post infusion OR Heparin 5ml for port – 100 units/ml

NURSING: Per Nevada Infusion

LABS ORDERS: \_\_\_\_\_ Fax results to: \_\_\_\_\_

### PROVIDER INFORMATION:

Physician Name: \_\_\_\_\_ NPI: \_\_\_\_\_  
 Physician Signature: \_\_\_\_\_ Date: \_\_\_\_\_  
 Point of Contact: \_\_\_\_\_ Phone: \_\_\_\_\_ Email: \_\_\_\_\_

**Please Fax This Form With - DEMOGRAPHICS, LABS, MEDICATION LIST and H&P: 775-470-8478**

\*\*Insurance verification/authorization is always obtained by Nevada Infusion prior to scheduling patients. \*\*



Patient Name: \_\_\_\_\_ DOB: \_\_\_\_\_

**Please Include Required Documentation for Expedited Order Processing & Insurance Approval:**

- Signed provider orders (page 1)
- Patient demographic and insurance information
- Patient's current medication list
- Supporting recent clinical notes and H&P (to support primary diagnosis)
- Supporting documentation to include past tried and/or failed therapies
- Supporting clinical notes to include any past tried and/or failed therapies, intolerance, benefits, or contraindications to conventional therapy
  - Has the patient had a documented contraindication/intolerance or failed trial of a DMARD, NSAID, or conventional therapy (i.e., MTX, leflunomide)?
    - Yes OR  No
    - If yes, which drug(s)? \_\_\_\_\_
  - Does the patient have a contraindication/intolerance or failed trial to at least one biologic (i.e., Humira, Enbrel, Stelara, Cimzia)?
    - Yes OR  No
    - If yes, which drug(s)? \_\_\_\_\_
  - Include labs and/or test results to support diagnosis (please attach results):
    - Rheumatoid factor
    - Anti-Cyclic citrullinated peptide (anti-CCP)
    - CRP and/or ESR

If applicable - Last known biological therapy: \_\_\_\_\_ and last date received: \_\_\_\_\_.  
If the patient is switching to biologic therapies, please perform a wash out period of \_\_\_\_\_ weeks prior to starting Simponi Aria.

Other medical necessity: \_\_\_\_\_

**Additional REQUIRED Information:**

- TB screening test completed within 12 months - (please attach results)
  - Positive OR  Negative
- Hepatitis B screening test completed. This includes Hepatitis B antigen and Hepatitis B core antibody total (not IgM) - (please attach results)
  - Positive OR  Negative

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