

RHEUMATOLOGY ORDER SET

P: 877.365.5566 | **F:** 855.889.2946

PATIENT IN	NFORMATION: Fax completed form, insurance information, and clinical docu	mentation to 855.889.2946	
Patient Name:	: DOB: Phon	e:	
Patient Status	S: □ New to Therapy □ Continuing Therapy Next Treatment Date:		
MEDICAL IN	NFORMATION		
Patient Weight: lbs. Patient Height: Allergies:			
	Diagnosis:		
□ Rheumatoid Arthritis, Unspecified □ Wegener's granulomatosis □ Unspecified Iridocyclitis □ Ankylosing Spondylitis, Unspecified			
	hropathic Psoriasis, Unspecified Gout		
	□ Rheumatoid Arthritis with Rheumatoid Factor, Unspecified □ Systemic Lupus Erythematosus		
☐ Rheumatoid Arthritis without Rheumatoid Factor, Unspecified ☐ Other:			
THERAPY ORDER			
Drug	Dosing	Refill	
Actemra	☐ 4 mg/kg IV every 4 weeks for doses, then followed by 8mg/kg every 4weeks th ☐ 4 mg/kg IV every 4 weeks ☐ 8 mg/kg IV every 4 weeks ☐ Other dose: mg IV every 4 weeks	/CRS**	
Cimzia	☐ Initial Dose: 400mg subcutaneously at weeks 0, 2, and 4 weeks Maintenance Dose: ☐ 200mg subcutaneously Q 2 weeks OR ☐ 400mg subcutaneously Q	4 weeks	
Krystexxa	☐ 8mg IV every 2 weeks		
Immunoglobulin	□ IV □ SubQ gm/kg x day(s) OR divided over day(s) mg/kg x day(s) OR divided over day(s) Frequency: Every weeks or	f not indicated)	
Orencia	Orencia Dose: mg IV Frequency: □ Every 4 weeks OR □ 0, 2, 4 weeks, and every 4 weeks thereafter		
Simponi Aria	☐ Initial Dose: 2mg/kg at weeks 0, 4, and then every 8 weeks ☐ Maintenance Dose: 2mg/kg every 8 weeks		
Stelara	Initial Dose: 45mg subcutaneously initially, 4 weeks later, followed by 45mg every 12 mg 90mg subcutaneously initially, 4 weeks later, followed by 90mg every 12 mg Maintenance Dose: 45mg subcutaneously every 12 weeks maintenance Dose: 90mg subcutaneously every 12 weeks	weeks weeks	
Infliximab	Dose:mg/kg	equirement	
Rituximab	Dose: \[\begin{array}{ c c c c c c c c c c c c c c c c c c c		
Saphnelo	□ 300mg IV every 4 weeks		
Premedication orders: Tylenol ☐ 1000mg ☐ 500mg PO, please choose one antihistamine: ☐ Diphenhydramine 25-50mg PO/IV ☐ Loratadine 10mg PO ☐ Cetirizine 10mg PO ☐ Cetirizine 10mg IVP Additional premedications: ☐ Solu-Medrol mg IVP ☐ Solu-Cortef mg IVP ☐ Other Lab orders: ☐ Yearly TB QFT (optional) ☐ Baseline HepBcAB total			
PROVIDER	INFORMATION		
By signing this form and utilizing our services, you are authorizing <i>Paragon Healthcare, Inc.</i> and its employees to serve as your prior authorization and specialty pharmacy designated agent in dealing with medical and prescription insurance companies, and to select the preferred site of care for the patient.			
Drovider NDI	c Signature Signature Contact Dorso	Date	
□ Opt out of P	e: Signature: Contact Perso Paragon selecting site of care (if checked, please list site of care):	116	
PREFERRED LOCATION			
City:	State: View our locations here:		

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IMPORTANT NOTICE: This fax is intended to be delivered only to the named address and contains material that is confidential, privileged property, or exempt from disclosure under applicable law. If you are not the named addressee, you should not disseminate, distribute, or copy this fax. Please notify the sender immediately and destroy all copies if you have received this document in error.





COMPREHENSIVE SUPPORT FOR RHEUMATOLOGY THERAPY

PATIENT INFORMATION:
Patient Name: DOB:
REQUIRED DOCUMENTATION FOR REFERRAL PROCESSING & INSURANCE APPROVAL
☐ Include <u>signed</u> and <u>completed</u> order (MD/prescriber to complete page 1)
☐ Include patient demographic information and insurance information
☐ Include patient's medication list
☐ Supporting clinical notes to include any past tried and/or failed therapies, intolerance, benefits, or contraindications to conventional therapy
☐ For biologic orders, has the patient had a documented contraindication/intolerance or failed trial of a conventional therapy (i.e., steroids)? ☐ Yes ☐ No If yes, which drug(s)?
☐ For biologic orders, does the patient have a contraindication/intolerance or failed trial to any other biologic? ☐ Yes ☐ No If yes, which drug(s)?
☐ Include labs and/or test results to support diagnosis
☐ If applicable - Last known biological therapy: and last date received: If patient is switching to biologic therapies, please perform a washout period of weeks prior to starting ordered biologic therapy.
☐ Other medical necessity:
REQUIRED PRE-SCREENING (BASED ON DRUG THERAPY)
 □ TB screening test completed within 12 months - attach results Required for: Actemra, Cimzia, infliximab, Stelara, Simponi Aria, Orencia □ Positive □ Negative
 ☐ Hepatitis B screening (Hepatitis B surface antigen) - ☐ Positive ☐ Negative Required for: Actemra, Cimzia, infliximab, rituximab, Simponi Aria Hepatitis B core antibody total (not IgM) - ☐ Positive ☐ Negative Required for: rituximab
☐ Serum immunoglobulins - attach results Recommended for: rituximab
☐ Baseline creatinine - attach results Required for: IVIG
*If TP or Hanatitic P results are necitive. places provide decumentation of treatment or medical clearance, and a negative CVD (TP+)

"If TB or Hepatitis B results are positive - please provide documentation of treatment or medical clearance, and a negative CXR (TB+)

Paragon Healthcare will complete insurance verification and submit all required documentation for approval to the patient's insurance company for eligibility. Our team will notify you if any additional information is required. We will review financial responsibility with the patient and refer him/her to any available co-pay assistance as needed. Thank you for the referral.

Please fax all information to (855) 889-2946 or call (877) 365-5566 for assistance