

## Krystexxa Order Form (pegloticase)

FAX TO: 972.499.9210

PATIENT INFORMATION			
		Sex: M F Ht: Wt: lbs kg	
Primary Language:	Allergies:		
Patient Preferred Location:			
<icd 10="" code="" required=""> DIAGNOSIS &amp; CLINICAL INFORMATION</icd>			
ICD 10 Code (PROVIDE COMPLETE CO	DE) <u>Prescribing Information</u>	mation	
M1A 0 Chronic Gout, w/o To M1A 1 Chronic Gout, w/ Top		that the patient discontinue oral urate-lowering medications e week) before starting Krystexxa.	
Other:	Recent data sugge	ests that patients may have improved outcomes when s are taken with Krystexxa.	
		<ul> <li>The recommended dosage is Krystexxa 8 mg every two weeks, co-administered w/weekly methotrexate 15 mg orally.</li> </ul>	
<ul> <li>Krystexxa alone may be used in patients for whom methotrexate is contraindicated or not clinically appropriate.</li> </ul>			
<u>REQUIRED</u> : Demographics and Most Recent: H&P, clinical notes, & medication list. Supporting clinical notes to include any past tried and/or failed therapies, intolerance, outcomes, or contraindications to conventional therapy. <u>LAB RESULTS</u> : G6PD, baseline uric acid > 6.0 mg/dL.			
PRESCRIPTION			
Pre-Medications Required: Acetaminophen: 650 mg PO, may repeat of Diphenhydramine: 25 mg IVP, may repeat Methylprednisolone: 125 mg SIVP Other:	q 6 hours, PRN infusion reaction	Lab Orders+ Required: Uric Acid Level, 24-72 hours prior to infusion If Uric Acid Level > 6 mg/dL upon two consecutive lab draws, hold dose, and contact prescriber  +Medix Infusion will draw maintenance labs unless otherwise directed by Referring Provider	
KRYSTEXXA (pegloticase)			
Loading Dose IV: Infuse 8 mg in 250 mL of 0.9% Sodium Chloride over 2 hours, every 2 weeks for one year  Is the patient on any other disease modifying therapy? Yes No If yes, please note therapy and last dose:  Post Treatment Observations: The patient is observed for 60 minutes following the first and second administrations.			
Adverse Events: In the event of an adverse reaction occurring at a Medix Infusion suite, utilize the Medix Infusion adverse reactions protocol.			
PRESCRIBER INFORMATION			
Prescriber Name: Signature:			
		Ity:	
Supervising Physician:(If Applicable)			
		State: Zip:	
	•	Fax: Email:	