

Request for Prior Authorization for Krystexxa (pegloticase)
Website Form – www.highmarkhealthoptions.com
Submit request via: Fax - 1-855-476-4158

All requests for Krystexxa (pegloticase) require a Prior Authorization and will be screened for medical necessity and appropriateness using the criteria listed below.

Krystexxa (pegloticase) Prior Authorization Criteria:

- The member must be 18 years of age or older
- The medication must be prescribed by or in consultation with a rheumatologist
- Documentation the member will be using the medication in combination with methotrexate unless methotrexate is contraindicated or not clinically appropriate
- An attestation from the prescriber that the member has discontinued all oral uratelowering medications prior to starting Krystexxa and will not restart therapy with an oral urate-lowering agent while on Krystexxa
- There must be documentation of a recent (within 1 month) serum uric acid level ≥ 6 mg/dL
- The member must have tried and failed or had an intolerance or contraindication to 3 months of therapy with allopurinol
 - o Failure is considered both of the following:
 - the inability to normalize uric acid to less than 6mg/dL
 - frequent gout flares (≥2 flares/year) and/or nonresolving subcutaneous tophi
- The member must have tried and failed or had an intolerance or contraindication to 3 months of therapy with at least one of the following:
 - o allopurinol in combination with an uricosuric agent (i.e. probenecid.)
 - o another xanthine oxidase inhibitor (XOI) (i.e. febuxostat (non-preferred XOIs require prior authorization))
 - o Failure is considered both of the following:
 - the inability to normalize uric acid to less than 6mg/dL
 - frequent gout flares (≥2 flares/year) and/or nonresolving subcutaneous tophi
- The requested dose and frequency is in accordance with FDA-approved labeling, nationally recognized compendia, and/or evidence-based practice guidelines
- Initial Duration of Approval: 6 months

• Reauthorization criteria:

The member must have documentation from the prescriber indicating improvement or stabilization in condition



- o Documentation of two recent consecutive (within the last 12 months) serum uric acid levels.
 - Rationale from the prescriber for why treatment must be continued if serum uric acid levels increase to above 6 mg/dL, particularly when 2 consecutive levels above 6 mg/dL are observed.
- Reauthorization Duration of Approval: 12 months

Coverage may be provided for any non-FDA labeled indication if it is determined that the use is a medically accepted indication supported by nationally recognized pharmacy compendia or peer-reviewed medical literature for treatment of the diagnosis(es) for which it is prescribed. These requests will be reviewed on a case by case basis to determine medical necessity.



KRYSTEXXA (PEGLOTICASE) PRIOR AUTHORIZATION FORM

Please complete and fax all requested information below including any progress notes, laboratory test results, or chart				
documentation as applicable to Highmark Health Options Pharmacy Services. FAX: (855) 476-4158				
If needed, you may call to speak to a Pharmacy Services Representative.				
PHONE: (844) 325-6251 Monday through Friday 8:00am to 7:00pm				
PROVIDER IN		in to 7.00pm		
Requesting Provider:	NPI:			
Provider Specialty:				
		Office Contact:		
Office Address:	Office Phone:			
	Office Fax:			
MEMBER IN				
Member Name:	OB:			
Member ID:	Member weight:	Height:		
	_			
REQUESTED DRU	G INFORMATION			
Medication:	Strength:			
Directions:	Quantity:	Refills:		
Is the member currently receiving requested medication?	Date Medication In			
Yes No	Date Medication in	macci.		
Is this medication being used for a chronic or long-term condition for which the medication may be necessary for the life				
_ • _	illion for which the m	edication may be necessary for the me		
of the patient? Yes No	c			
	formation			
This medication will be billed: at a pharmacy OR		_		
medically (if medically p	•			
Place of Service: Hospital Provider's office Member's home Other				
Place of Service	ce Information			
Name:	NPI:			
Address:	Phone:			
MEDICAL HISTORY (C	amplete for AII rec	mosts)		
		uests)		
Diagnosis: Chronic Gout Other				
Serum uric acid level: Date taken:				
Has the member discontinued all oral urate lowering medications prior to starting therapy and will the member stay off				
all urate lowering medications while on therapy with Krystexxa? Yes No				
Will the member be using Krystexxa in combination with methotrexate? Yes No				
If no please explain why:				
REAUTHORIZATION				
Has the member shown improvement or stabilization? Yes No				
2 Consecutive serum uric acid levels: Date taken:				
Date taken				



Member Name:	DOB:				
Member ID:					
CURRENT or PREVIOUS THERAPY					
Medication Name	Strength/ Frequency	Dates of Therapy	Status (Discontinued & Why / Current)		
SUPPORTING INFORMATION or CLINICAL RATIONALE					
Prescribing Provider Signature Date			Date		

