

INFUSION* Phone: 1-800-809-1265 Fax: 1-866-872-8920

Referral Status:	MRN:	
New referral	Order change	Order Renewal
Patient preferred clinic:		

Substitution permitted

Krystexxa [®] (pegloticase) Standard Plan of Treatmen	Krvstexxa[®]	(pegloticase)	Standard Plan	of Treatmen
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_	ENT DEMOGRAPH											
	of Referral:	nco	•			Patie	nt's Phone:					
	nt Name:					Addre						
	of Birth:						State, Zip:					
	nt in inches:	We	eight:	LE	or KG	Gend		Allergies	:	See lis	st	NKDA
				- ND				1 2		1 1		
DIAG	GNOSIS: (PLEASE C				3 ND DIGITS TO CO	MPLE	TE ICD 10 FOR BIL	LING)				
	M1A Chronic g M1A Chronic g											
	M10 Idiopathic	-	-	111								
	- Other:	y got										
REO	UESTED DOCUMEN	NTA	TION:		PREVIOUS ADMI	NISTR	ATION: HAS THIS P	ATIENT TAK	EN THIS N	VEDICATIC	DN BI	EFORE?
1	Insurance information				IF NO:	IF YE						
2	Most recent History &	Phys	sical		PLEASE STATE	LAST	INFUSION DATE:					
3	Full medication list				REQUIRED WASHOUT	NEXT	INFUSION DATE:					
4	Tried and failed therap	ies			THERAPY:	IF OR	DER CHANGE:					
5	Baseline serum uric ad	cid le	evel				Continuo ou	www.mt.ordo	r until in a			wed
6	G6PD serum level						Continue cu	irrent ordei	runui ins	surance a	opro	vea
7	Specify if patient is pre	escril	bed prop	hylaxis for g	jout flare:	8	Specify if patient is p therapy:	rescribed metl	hotrexate or	other immur	iomod	dulation
MFD	ICATION ORDERS:											
	: Patient may be ineligi		o receive	e Krystexxa	[®] if patient has a diagn	osis of	G6PD or has new or w	orsenina symr	otoms of CH	F If appropri	iate i	tis
	mended that Krystexxa [®]										,	
	EDICATION TO BE ADMIN											
Manu	facturing guidelines sug	gest			of IV corticosteroids an	nd antihi				1		
	Diphenhydramine	_	25mg	50mg		_	Acetaminophen	325mg	500mg	650mg		1000mg
IV	Methylprednisolone		40mg	125mg	Other:	_	Famotidine	20mg	40mg			
	Famotidine Other:	-	20mg	40 mg		РО	Diphenhydramine Fexofenadine	25mg	50mg 180mg			
MED	DICATION/DOSE:						Cetirizine	60mg 10mg	roomy			
	Krystexxa [®] (peglo	tica	aa) 0 m	a in 250.	n NC N/ to infuso		Loratadine	10mg				
	over 2 hours	uca	se) o n	ig in 250i			Other:	Tomy				
	Monitor patient	for	one (1) hour po	ost infusion		PARAMETERS: (Ph	armacist to n	orform clini	cal lab moni	toring	a)
	monitor putient		mpleti			LAD	Serum uric acid level					
							infusion if 2 consecut	•				
FREC	QUENCY:					Ple	ase ensure all lab v	work is faxe	d to Palm	etto Infusio	on Se	<u>ervices</u>
	Dosing every 2 we	eks	6			SPEC	CIAL/LAB ORDERS:					
	Other:						7					
lf 2 d	loses (4 weeks) of the	rapv	are mis	sed. then i	eferring provider mu	st aive	written clearance to	resume thera	pv or treatn	nent will be	disco	
							Refills x 12 months					
LINE	USE/CARE ORDER	s.					ADVERSE REACT	ΊΟΝ & ΔΝΔ	ΦΗΥΙΔΧΙ			
	Start PIV/Access CVC				ADVERSE REACTION & ANAPHYLAXIS ORDERS: Administer acute infusion and anaphylaxis							
	Flush device per fac		standar	rd flushing	procedure		medications per Paln).(1) 2 1		
\checkmark		Jinty	Standar	a nashing	procedure		adverse reaction orde				hie	
							our website or scan h	nere.			36	
											0.4783	
	SCRIBER INFORMA	TIO	N:				PLIONE					
	VIDER NAME:						PHONE:					
	RESS:						FAX:					
-	, STATE, ZIP:		/		· · · · · · · · · · · · · · · · · · ·		NPI:					
PRES	SCRIBER SIGNATU	RE:	(No sta	imp signa	tures)					DATE:		
1												

Dispense as written/Brand medically necessary

Patient Enrollment Form Once complete, submit by fax 1-877-633-9522 or email GoutHBYS@horizontherapeutics.com

Complete all required fields, including prescriber's signature and date, to initiate patient enrollment process.

For patient support and/or assistance obtaining patient signature, call Horizon By Your Side at 1-877-633-9521.

Patient Information (*Indicates a required field)

First name*	L	ast name*		
Sex*: O Male O Female	D	ate of birth*:	(MM/DD/YYYY)	
Primary language	E	mail address		
Primary telephone*		ave voice message at p ate contact telephone?		O No
O Home O Cell	Consent to se	and text message?	O Yes	O No
Address*				
City*	<u>s</u>	tate*	ZIP code*	
Alternate contact name	A	lternate contact telepl	none	

(*Indicates a required field) (Please include front and back copies of insurance card[s] with this form) Insurance Information

rimary insurance*	Secondary insurance, if applicable
olicy #*	Policy #
olicyholder's first and last name*	Policyholder's first and last name
nsurance company telephone*	Insurance company telephone
Group #*	Group #
olicyholder's DOB*:(MM/DD/YYYY)	Policyholder's DOB:(MM/DD/YYYY)
PA/Medical group name	IPA/Medical group telephone
Reverification request	

Patient is uninsured to my knowledge

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Infusion Facility (*Indicates a required field)

Do you have a preferred infusion facility?* O Yes O No If yes, please fill out the preferred infusion facility information below. If no, Horizon By Your Side will help identify a facility in close proximity to your patient.

The infusion facility is the same as the prescribing office

Facility name* Facility address* City* State* ZIP code* Telephone* Fax* Facility NPI #* Facility tax ID #* Patient Authorization (Required – please see authorization language on page 2) Date: Patient signature (MM/DD/YYYY) Please read page 2

Printed full name

Please see Important Safety Information on page 2 and see Full Prescribing Information, including Boxed Warning, at KRYSTEXXAhcp.com.





ORIZON

First name*			
Address*			
City*		State* ZIP code*	
NPI #*	Tax ID #*	State license #*	
Clinic/hospital affiliation			
Office contact name			
Office contact telephone*		Fax*	
mail address*			
	Telephone 🔘 Er	nail Prescriber specialty*:	
Referring healthcare provider: referred to you by another HCP?	Was this patient	Yes No If yes, please popula	ate:
Name:		Specialty:	
City:		State:	
		Telephone:	
		·	
	or benefits investig a required field)	jation)	
dditional disease manifestation	codes:		
Additional disease manifestation			
Co-administration	Medication	s 🔘 No If yes, please indicate below:	
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I certify that the above therapy is medically necessary, that the information provided is accurate to the best of my knowledge and that my patient is being administered KRYSTEXXA* (pegloticase) injection, 8 mg/mL, for intravenous infusion in accordance with the labeled use of the product. I understand that Horizon Therapeutics USA, Inc. and its affiliates and their respective employees or agents (collectively, "Horizon") will use this information to administer the Horizon By Your Side program (the "Program"), which provides a wide array of patient-focused services, including providing logistical and non-medical treatment support for KRYSTEXXA as prescribed, and educating about the insurance process. By my signature, I also certify that (1) my patient or his/her personal representative has provided a signed HIPAA authorization that allows me to share protected health information with Horizon for purposes of the Program and (2) I have obtained the patient's authorization to release such information as may be required for AllCare Plus Pharmacy (or another party acting on behalf of Horizon) to assess insurance coverage for KRYSTEXXA and assistance in initiating or continuing KRYSTEXXA as prescribed. I further understand and agree that (a) any medication or service provided through the Program as a result of this form is for the named patient only and is not being made in exchange for any express or implied agreement or understanding that I would recommend, prescribe, or use KRYSTEXXA* was based solely on my professional determination of medical necessity; and (c) I will not seek reimbursement for any medication or service provided by or through the Program or the responsibility of the patient and healthcare provider. Horizon makes no representation or guarantee concerning coverage or reimbursement for any item or service. On behalf of the patient, Horizon expects the prescriber's and work with Horizon By Your Side to effectively communicate both in-network and ocumentation are the responsibility of the patient and

State requirements: The prescriber is to comply with his/her state-specific prescription requirements such as e-prescribing, state-specific prescription form, fax language, etc. Noncompliance with state-specific requirements could result in outreach to the prescriber.

By filling out and signing this form, the enrollment process in Horizon By Your Side has initiated; however, your patient must sign a Patient Authorization to complete enrollment in Horizon By Your Side. Please note that your patient will not benefit from the services and support offered by the Program unless your patient signs a Patient Authorization, consenting to receiving such services. If your patient does not sign the Patient Authorization contained within this form, Horizon will contact the patient to determine whether the patient is interested in signing a separate Patient Authorization.

Patient Consent for Patient Information, Enrolling in Services, and Accessing Financial Support (referred to as "Patient Authorization") Please read and provide signature in Patient Authorization section on page 1

I hereby authorize my healthcare providers, my health insurance carriers, and my pharmacies to use and disclose my individually identifiable health information, including my medical records, insurance coverage information, and my name, address, and telephone number to Horizon Therapeutics USA, Inc. and its affiliates and their respective agents and representatives (collectively, "Horizon"), including third parties authorized by Horizon to administer drug support and to dispense drugs (collectively, "Horizon by Your Side") for the following purposes: (1) to establish eligibility for benefits; (2) to communicate with my healthcare providers and me about my treatment or condition and related products; (3) to facilitate the provision of products, supplies, or services by third party including, but not limited to, specialty pharmacies; (4) to register me in any applicable product registration programs required for my treatment; (5) to enroll me in eligible patient support programs offered by Horizon By Your Side and/or Horizon, including nursing or patient access support services (government-reimbursed programs may not be eligible for all support services offered; please contact Horizon By Your Side for determination); and (6) to send me marketing information or offer me products and services related to my treatment or condition (or other products or services in which I might be interested) and to contact me occassionally to obtain my feedback (for market research purposes only) about my treatment, my condition, or my experience with Horizon By Your Side dortherwise as required or permitted by law. Further, I appoint the Program, on my behalf, to proceed with Program send to convey this prescription to the dispensing pharmaccity to the extent permitted under state law. I understand the pharmacies may receive a fee from Horizon health information.

I understand that Horizon, as well as my healthcare providers, cannot require me, as a condition of having access to medications, prescription drugs, treatment, or other care, to sign this Authorization. I understand that I am entitled to a copy of this Authorization.

I understand that information disclosed pursuant to this Authorization in some cases may be redisclosed by the recipient and no longer protected by HIPAA or other privacy laws. But Horizon has agreed to use and disclose my information only for purposes of operating the Program. I understand that I may cancel this Authorization at any time by mailing a signed letter requesting such cancellation to Horizon BY Jour Side, I Horizon Way, Deerfield, IL 60015, but that this cancellation will not apply to any information used or disclosed by my healthcare providers and/or health insurance carriers based on this Authorization is valid for whichever is greater: (a) the duration remaining on this treatment or (b) 10 years from the date signed above. A photocopy of this Authorization will be treated in the same manner as the original.

INDICATION

KRYSTEXXA® (pegloticase) is indicated for the treatment of chronic gout in adult patients who have failed to normalize serum uric acid and whose signs and symptoms are inadequately controlled with xanthine oxidase inhibitors at the maximum medically appropriate dose or for whom these drugs are contraindicated.

Limitations of Use: KRYSTEXXA is not recommended for the treatment of asymptomatic hyperuricemia.

IMPORTANT SAFETY INFORMATION

WARNING: ANAPHYLAXIS AND INFUSION REACTIONS, G6PD DEFICIENCY ASSOCIATED HEMOLYSIS AND METHEMOGLOBINEMIA

Anaphylaxis and infusion reactions have been reported to occur during and after administration of KRYSTEXXA.

- Anaphylaxis may occur with any infusion, including a first infusion, and generally manifests within 2 hours of the infusion. Delayed hypersensitivity reactions have also been reported.
- KRYSTEXXA should be administered in healthcare settings and by healthcare providers prepared to manage anaphylaxis and infusion reactions.
- Premedicate with antihistamines and corticosteroids and closely monitor for anaphylaxis for an appropriate period after administration of KRYSTEXXA.
- Monitor serum uric acid levels prior to each infusion and discontinue treatment if levels increase to above 6 mg/dL, particularly when 2 consecutive levels above 6 mg/dL are observed.
- Screen patients at risk for glucose-6-phosphate dehydrogenase (G6PD) deficiency prior to starting KRYSTEXXA. Hemolysis and methemoglobinemia have been reported with KRYSTEXXA in patients with G6PD deficiency. KRYSTEXXA is contraindicated in patients with G6PD deficiency.

CONTRAINDICATIONS:

- · In patients with G6PD deficiency.
- · In patients with history of serious hypersensitivity reactions, including anaphylaxis, to KRYSTEXXA or any of its components.

WARNINGS AND PRECAUTIONS

Gout Flares: An increase in gout flares is frequently observed upon initiation of anti-hyperuricemic therapy, including KRYSTEXXA. Gout flare prophylaxis with a non-steroidal anti-inflammatory drug (NSAID) or colchicine is recommended starting at least 1 week before initiation of KRYSTEXXA therapy and lasting at least 6 months, unless medically contraindicated or not tolerated.

Congestive Heart Failure: KRYSTEXXA has not been formally studied in patients with congestive heart failure, but some patients in the pre-marketing placebocontrolled clinical trials experienced exacerbation. Exercise caution in patients who have congestive heart failure and monitor patients closely following infusion.

ADVERSE REACTIONS

The most commonly reported adverse reactions (≥5%) are:

KRYSTEXXA co-administration with methotrexate trial:

KRYSTEXXA with methotrexate: gout flares, arthralgia, COVID-19, nausea, and fatigue; KRYSTEXXA alone: gout flares, arthralgia, COVID-19, nausea, fatigue, infusion reaction, pain in extremity, hypertension, and vomiting.

KRYSTEXXA pre-marketing placebo-controlled trials:

gout flares, infusion reactions, nausea, contusion or ecchymosis, nasopharyngitis, constipation, chest pain, anaphylaxis, and vomiting.

For additional information on KRYSTEXXA, please see Full Prescribing Information, including Boxed Warning, at KRYSTEXXAhcp.com.



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