

lt's Wholecare.

Prior Authorization Criteria Crysvita (burosumab-twza)

All requests for Crysvita (burosumab-twza) require a prior authorization and will be screened for medical necessity and appropriateness using the criteria listed below.

Coverage may be provided with a <u>diagnosis</u> of X-linked hypophosphatemia (XLH) and the following criteria is met:

- Confirmation of the diagnosis by at least one of the following:
 - Genetic test showing a PHEX gene mutation (phosphate regulating gene with homology to endopeptidase on the X chromosome)
 - Serum fibroblast growth factor 23 (FGF23) level above the upper limit of normal for the reference range for the member's age (reference range must be provided)
- Member must be 6 months or older
- Must be prescribed by or in consultation with a physician who is experienced in the management of patients with metabolic bone disease.
- The requested dose and frequency is in accordance with FDA-approved labeling, nationally recognized compendia, and/or evidence-based practice guidelines.
- An attestation from the provider the Crysvita will not be used together with oral phosphate and active vitamin D analogs
- Baseline fasting serum phosphorus concentration that is below the reference range for the member's age (reference range must be provided)
- For members under 18 years of age documentation of one of the following:
 - $\circ \quad \text{Baseline recumbent length/standing height z score}$
 - Baseline serum alkaline phosphatase activity
 - Baseline Thacher Rickets Severity Score (RSS)
- For members 18 years and older documentation of one of the following:
 - An attestation from the provider that the member is experiencing skeletal pain
 - Total healing fracture amount
 - Baseline osteoid volume/bone volume
- Initial Duration of Approval: 12 months
- Reauthorization criteria
 - For members under 18 years of age
 - An increase in fasting serum phosphorus from baseline taken within last 12 months but not greater than 5.0mg/dL
 - Documentation the member has a positive clinical response or stabilization in their disease demonstrated by one of the following
 - An increase in height z score from baseline
 - A decrease in serum alkaline phosphatase activity from baseline
 - A decrease in the RSS score from baseline or a positive Radiographic Global Impression of Change (RGI-C) score.
 - For members 18 years and older



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- An increase in fasting serum phosphorus from baseline taken within last 12 months (the level must also be below the upper limit of normal for the lab range; reference range must be provided)
- Documentation the member has a positive clinical response or stabilization in their disease demonstrated by one of the following:
 - An attestation there has been improvement in the member's pain
 - Total fractures healing after starting therapy
 - A decrease in osteoid volume/bone volume from baseline
- **Reauthorization Duration of Approval:** 12 months

Coverage may be provided with a <u>diagnosis</u> of FGF23-related hypophosphatemia in Tumor Induced Osteomalacia and the following criteria is met:

- Member must be 2 years of age or older
- Documentation member has a phosphaturic mesenchymal tumor that cannot be resected or localized
- Baseline fasting serum phosphorus concentration that is below the reference range for the member's age (reference range must be provided)
- Must be prescribed by or in consultation with a hematologist, oncologist, or endocrinologist
- Initial Duration of Approval: 12 months
- Reauthorization criteria
 - An increase in fasting serum phosphorus from baseline taken within last 12 months (the level must also be below the upper limit of normal range for the lab range; reference range must be provided)
 - Documentation the member has a positive clinical response or stabilization in their disease demonstrated by one of the following:
 - An attestation there has been improvement in the member's pain
 - Total fractures healing after starting therapy
 - A decrease in osteoid volume/bone volume from baseline
 - Improved growth velocity
- **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.

When criteria are not met, the request will be forwarded to a Medical Director for review. The physician reviewer must override criteria when, in their professional judgment, the requested medication is medically necessary.



CRYSVITA (burosumab-twza) PRIOR AUTHORIZATION FORM			
Please complete and fax all requested information below including any progress notes, laboratory test results, or chart documentation			
as applicable to Gateway Health SM Pharmacy Services. FAX: (888) 245-2049			
	a Pharmacy Services Representative.		
	v through Friday 8:30am to 5:00pm		
PROVIDER INFORMATION Requesting Provider:	NPI:		
Provider Specialty:	Office Contact:		
Office Address:	Office Phone:		
	Office Fax:		
MEMBER INFORMATION			
Member Name:	DOB:		
Gateway ID:	Member weight:pounds orkg		
REQUESTED DRUG INFORMATION			
Medication:	Strength:		
Frequency:	Duration:		
Is the member currently receiving requested medication? Yes	No Date Medication Initiated:		
Billing Information			
This medication will be billed: \Box at a pharmacy OR			
medically (if medically please provide a JCODE:			
	r's home 🗌 Other		
Place of Service Information			
Name:	NPI:		
Address:	Phone:		
MEDICAL HISTORY (Complete for ALL requests)			
Diagnosis: X-linked hypophosphatemia (XLH) ICD-10 Code: Tumor- Induced Osteomalacia ICD-10 Code:			
Other: ICD-10 Code:			
For X-Linked Hypophosphatemia:			
Has the diagnosis been confirmed by at least one of the following? <i>Please select all that are applicable</i> Yes No			
Genetic test showing a PHEX gene mutation (phosphate regulating gene with homology to endopeptidase on the X			
chromosome)			
Serum fibroblast growth factor 23 (FGF23) level > 30pg/mL			
Is the medication prescribed by, or in consultation with, a physician who is experienced in the management of patients with			
metabolic bone disease? Yes No			
Baseline fasting serum phosphorus concentration:	Deference rence		
Basenne fasting serum phosphorus concentration:	Kelerence range:		
Will Crysvita be used in together with oral phosphate and active vitamin D analogs?			
For Members under 18 years of age:			
Please provide one of the following:			
Baseline recumbent length/standing height z score:			
Baseline serum alkaline phosphatase activity:			
Baseline Thacker Rickets Severity Score (RSS):			



For Members 18 years of age and older:

Please provide one of the following:

- Attestation from the provider that the member is experiencing skeletal pain: _____ •
- Total healing fracture amount: ____ •
- Baseline osteoid volume/bone volume:

For Tumor-Induced Osteomalacia

Does the member have a phosphaturic mesenchymal tumor that cannot be resected or localized? 🗌 Yes 🗌 No Baseline fasting serum phosphorus concentration: ______ reference range _____

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CURRENT or PREVIOUS THERAPY				
1				
Strength/ Frequency	Dates of Therapy	Status (Discontinued & Why/Current)		
REAUTHORIZATION Has the member experienced a significant improvement with treatment? Yes No Please describe:				
Has the member's fasting serum phosphorus concentration increased from baseline? Yes No - Fasting serum phosphorus concentration: Date collected:				
e of the following: re from baseline: e phosphatase activity fron	baseline:			
 For Members 18 years of age and older: Attestation from the provider that there has been improvement in the member's pain:				
r starting therapy: tt in skeletal pain from base ne date taken _ e date taken _	date taken _ line?			
UP OLINICAL KATIUNA				
	Date			
	Strength/ Frequency ficant improvement with tree sphorus concentration increation concentration: ge: e of the following: re from baseline: me phosphatase activity from e from baseline or a positive older: er that there has been improvement with the properties of the following: starting therapy: me/bone volume from baseling lowing: e starting therapy: me date taken date taken date taken date taken	Strength/ Frequency Dates of Therapy		



Updated: 10/2021 PARP Approved: 10/2021