



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:
 - O Genetic test showing a PHEX gene mutation (phosphate regulating gene with homology to endopeptidase on the X chromosome)
 - O 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:
 - o Baseline recumbent length/standing height z score
 - o Baseline serum alkaline phosphatase activity
 - o Baseline Thacher Rickets Severity Score (RSS)
- For members 18 years and older documentation of one of the following:
 - o An attestation from the provider that the member is experiencing skeletal pain
 - o Total healing fracture amount
 - o 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.
 - o For members 18 years and older
 - 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)
 - O 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 Highmark Wholecare Pharmacy Services. **FAX:** (888) 245-2049

If needed, you may call to speak to a Pharmacy Services Representative. **PHONE**: (800) 392-1147 Mon – Fri 8:30am to 5:00pm

3:00	лрт					
PROVIDER IN	IFORMA	TI	ON			
Requesting Provider:		Pro	ovider NPI:			
Provider Specialty:		Of	fice Contact:			
State license #:		Of	fice NPI:			
Office Address:		Of	fice Phone:			
		Of	fice Fax:			
MEMBER INFORMATION						
Member Name:	DOB:					
Member ID:		er weight:		Height:		
REQUESTED DRU	G INFO	RM	ATION			
Medication:	Streng	th:				
Directions:	Quantity:			Refills:		
Is the member currently receiving requested medication?	Yes		Date Medication	Initiated:		
No						
Billing In						
	medically					
	lember's					
Place of Service	e Inform					
Name:		NP	PI:			
Address:		Pho	one:			
MEDICAL HISTORY (Co	omplete f	or A	ALL requests)			
Diagnosis:						
X-linked hypophosphatemia (XLH)	ICD Code:					
Tumor- Induced Osteomalacia						
Other:						
For X-Linked Hypophosphatemia:	0 D1		1			
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						
Serum norodiast growth factor 23 (FGF23) level > 30pg/	IIIL					
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						
with metabolic bolic disease.						
Baseline fasting serum phosphorus concentration:			Referer	nce range:		
				0		
						



CRYSVITA (BUROSUMAB-TWZA) PRIOR AUTHORIZATION FORM (CONTINUED)— PAGE 2 of 3

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	MEMBER 1	INFORMATION			
Member Name:		DOB:			
Member ID:		Member weight:	Height:		
M	EDICAL HISTORY	(Complete for ALL re	equests)		
Will Crysvita be used in together	with oral phosphate and	active vitamin D analog	gs? Yes No		
For Members under 18 years of Please provide one of the follows Baseline recumbent leng Baseline serum alkaline Baseline Thacker Ricket For Members 18 years of age at Please provide one of the follows Attestation from the provide the provide one of the follows are provided one of the follows	ing: th/standing height z score phosphatase activity: s Severity Score (RSS): nd older: ing: vider that the member is o	experiencing skeletal pai	n:		
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					
☐ Yes ☐ No					
CURRENT or PREVIOUS THERAPY					
Medication Name	Strength/ Frequency	Dates of Therapy	Status (Discontinued & Why/Current)		
REAUTHORIZATION					
Has the member experienced a significant improvement with treatment? Yes No Please describe:					

CRYSVITA (BUROSUMAB-TWZA)



PRIOR AUTHORIZATION FORM (CONTINUED)- PAGE 3 of 3

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MEMBER INFORMATION				
Member Name:	DOB:			
Member ID:	Member weight: Height:			
REAUTHO	RIZATION			
Has the member's fasting serum phosphorus concentration in				
Fasting serum phosphorus concentration:	Date collected:			
For X-linked Hypophosphatemia: For Members under 18 years of age: Please provide documentation of one of the following: • An increase in height z score from baseline: • A decrease in serum alkaline phosphatase activity from	om baseline: ive Radiographic Global Impression of Change (RGI-C)			
 Total fractures healing after starting therapy: A decrease in osteoid volume/bone volume from bas 	· 			
For Tumor Induces Osteomalacia:				
please provide at least one of the following:				
Total healing fracture amount before starting therapy:	date taken aseline?			
SUFFORTING INFORMATIO	NOT GENIGAL RATIONALE			
Prescribing Provider Signature	Date			