SENTARA COMMUNITY PLAN (MEDICAID)

MEDICAL PRIOR AUTHORIZATION/STEP-EDIT REQUEST

<u>Directions:</u> The prescribing physician must sign and clearly print name (preprinted stamps not valid) on this request. All other information may be filled in by office staff; fax to 1-844-305-2331. No additional phone calls will be necessary if all information (including phone and fax #s) on this form is correct. If information provided is not complete, correct, or legible, authorization will be delayed.

Drug Requested: Xenpozyme[™] (olipudase alfa) (J3590/C9399) (Medical)

Member Name:	
	Date of Birth:
Prescriber Name:	
Prescriber Signature:	Date:
	Fax Number:
Phone Number:	
Phone Number: DEA OR NPI #: DRUG INFORMATION: A	Authorization may be delayed if incomplete.
Phone Number: DEA OR NPI #: DRUG INFORMATION: A Drug Form/Strength:	Authorization may be delayed if incomplete.
Phone Number: DEA OR NPI #: DRUG INFORMATION: A Drug Form/Strength: Dosing Schedule:	Authorization may be delayed if incomplete.

Dosing Limits

- A. Quantity Limit (max daily dose) [NDC Unit]:
 - Xenpozyme 20 mg lyophilized powder for reconstitution in a single-dose vial: 58468-0050-xx
 - Xenpozyme 20 mg single-dose vial: 17 vials per 14 days
- B. Max Units (per dose and over time) [HCPCS Unit]:
 - 340 mg every 14 days
 - *Use actual body weight for patients with a BMI \leq 30. For patients with a BMI > 30, calculate adjusted body weight (kg) = (actual height in m)² \times 30

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Xenpozyme (olipudase alfa), a hydrolytic lysosomal sphingomyelin-specific enzyme approved by the Food and Drug Administration (FDA) for treatment of non-central nervous system manifestations of acid sphingomyelinase deficiency (ASMD) in adult and pediatric individuals.

Xenpozyme is an intravenous enzyme replacement therapy and the first FDA-approved treatment for ASMD. Xenpozyme is not expected to cross the blood-brain barrier and is intended to treat the non-central nervous system disease manifestations. Xenpozyme has only been studied in individuals with the type B and A/B phenotypes.

Xenpozyme has a black box warning for severe hypersensitivity reactions including anaphylaxis. Appropriate medical support measures, including cardiopulmonary resuscitation equipment, should be readily available during Xenpozyme administration. If a severe hypersensitivity reaction occurs, Xenpozyme should be discontinued immediately, and appropriate medical treatment should be initiated.

CLINICAL CRITERIA: Check below all that apply. All criteria must be met for approval. To support each line checked, all documentation, including lab results, diagnostics, and/or chart notes, must be provided or request may be denied.

Initial Authorization: 12 months

	ember has a diagnosis of acid sphingomyelinase deficiency (ASMD) confirmed by ONE of the lowing:
	Pathogenic sphingomyelin phosphodiesterase-1 (SMPD1) gene mutation
	Deficiency in acid sphingomyelinase (ASM) enzyme activity <10% of controls as measured in fibroblasts, leukocytes or dried blood spot [Note: NPD-A (infantile neurovisceral ASMD) has not been studied. Genotype-phenotype correlations as well as signs/symptoms may not be conclusive in infants therefore requests will be evaluated on a case-by-case basis]
Me	ember has a clinical presentation consistent with ONE of the following ASMD types:
	ICD Code: E75.241 Niemann-Pick type B
	ICD Code: E75.244 Niemann-Pick type A/B
Re	quested medication will be used for the treatment of non-central nervous system disease manifestation
aw	ember should <u>NOT</u> require invasive ventilatory support OR non-invasive ventilatory support while take and for >12 hours a day (Note: Members requiring ventilatory support will be reviewed on a se-by-case basis)
Pro	ovider must submit documentation for at least ONE of the following:
	Baseline: percent predicted diffusion capacity of the lungs for carbon monoxide (DLco) or age-appropriate pulmonary function testing
	Baseline: spleen and/or liver volumes
	Baseline: plasma lyso-sphingomyelin
	Baseline: Platelet count
	Baseline: height Z-score and skeletal maturation (relevant for pediatric patients)

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PA Xenpozyme (Medical)(Medicaid)

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	Provider has submitted baseline transaminase (alanine aminotransferase [ALT] and aspartate aminotransferase [AST]) levels obtained within 1 month prior to treatment initiation	
	Females of reproductive potential will have pregnancy status verified prior to start of therapy and will use effective contraception during treatment and for 14 days after the last dose if therapy is discontinued	
	Requested dose has been provided:	
suppo	uthorization: 12 months. Check below all that apply. All criteria must be met for approval. To ort each line checked, all documentation, including lab results, diagnostics, and/or chart notes, must be ded or request may be denied.	
	Member has experienced an absence of unacceptable toxicity from the drug (e.g., anaphylaxis and severe hypersensitivity reactions, severe infusion-associated reactions, severely elevated liver transaminases)	
	Member has NOT experienced progressive/irreversible severe cognitive impairment	
	Member has shown disease improvement or stability from pre-treatment baseline as demonstrated by at least ONE of the following:	
	☐ Improvement in or stability in the percent predicted diffusion capacity of the lungs for carbon monoxide (DLco) or other age-appropriate pulmonary function testing	
	☐ Improvement in or stability of spleen and/or liver volumes	
	☐ Reduction in plasma lyso-sphingomyelin	
	☐ Improvement in or stability of platelet count	
	☐ Improvement in linear growth progression as measured by mean height Z-scores (pediatric patients only)	
	Provider has submitted transaminase (alanine aminotransferase [ALT] and aspartate aminotransferase [AST]) levels obtained within the last 30 days	
	Requested dose has been provided:	
Medication being provided by a Specialty Pharmacy - PropriumRx		
	Location/site of drug administration:	
	NPI or DEA # of administering location:	
	OR	
	Specialty Pharmacy - PropriumRx	
	gent reviews: Practitioner should call Sentara Pre-Authorization Department if they believe a standard would subject the member to adverse health consequences. Sentara's definition of urgent is a lack of	

Fo treatment that could seriously jeopardize the life or health of the member or the member's ability to regain maximum function.

**Use of samples to initiate therapy does not meet step edit/preauthorization criteria. ** *Previous therapies will be verified through pharmacy paid claims or submitted chart notes. *

REVISED/UPDATED: 11/29/2022

^{*}Approved by Pharmacy and Therapeutics Committee: 11/18/2022