

## Request for Prior Authorization for Gaucher Disease: Enzyme Replacement and Substrate Reduction Therapy Website Form – <a href="https://www.highmarkhealthoptions.com">www.highmarkhealthoptions.com</a>

**Submit request via: Fax - 1-855-476-4158** 

All requests for Gaucher Disease: Enzyme Replacement and Substrate Reduction Therapy require a Prior Authorization and will be screened for medical necessity and appropriateness using the criteria listed below.

## Gaucher Disease: Enzyme Replacement and Substrate Reduction Therapy Prior Authorization Criteria:

- Confirmation of Type 1 Gaucher Disease diagnosis by one of the following:
  - o A beta-glucosidase leukocyte (BGL) test
  - o A genetic test
- Is age appropriate according to FDA approved package labeling, nationally recognized compendia, or peer-reviewed medical literature.
- The requested dose and frequency is in accordance with FDA-approved labeling, nationally recognized compendia, and/or evidence-based practice guidelines
- Member must also have one or more of the following (supported by the corresponding documentation):
  - o Anemia
    - Documentation of a baseline low hemoglobin level (laboratory reference range must be submitted)
  - o Thrombocytopenia
    - Documentation of a baseline low platelet level (laboratory reference range must be submitted)
  - o A provider attestation that the member has one of the following:
    - Bone Disease
    - Hepatomegaly or splenomegaly supported by documentation of a current baseline liver or spleen size
- For Cerdelga (eliglustat):
  - Documentation the member is a CYP2D6 extensive metabolizer (EM), intermediate metabolizer (IM), or poor metabolizer (PM) as detected by an FDA-cleared test
  - o Coverage is not provided for CYP2D6 ultra-rapid metabolizers.
- For Zavesca (miglustat):
  - o Documentation the member is unable to use enzyme replacement therapy.
- **Initial Duration of Approval:** 6 months

## • Reauthorization criteria:

O Documentation the member had a positive clinical response or stabilization as demonstrated by one of the following:



- An increase in hemoglobin level from baseline (taken within last 6 months)
- An increase in platelet levels from baseline (taken within last 6 months)
- An attestation from the provider of an improvement in bone disease
- A reduction in liver or spleen size from baseline (taken yearly)
- Reauthorization Duration of Approval: 12 months



## GAUCHER DISEASE ERT AND SRT THERAPY 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 INFORMATION Requesting Physician: NPI: Physician Specialty: Office Contact: Office Address: Office Phone: Office Fax: MEMBER INFORMATION Member Name: DOB: Member weight: Member ID: Height: DRUG INFORMATION Medication: Strength: Directions: Ouantity: Refills: Is the member currently receiving requested medication? 

Yes No Date Initiated: Is this medication being used for a chronic or long-term condition for which the medication may be necessary for the life of the patient? Yes No **BILLING INFORMATION** This medication will be billed: 
at a pharmacy OR medically (if medically please provide a JCODE: Place of Service: Hospital Provider's office Member's home Other PLACE OF SERVICE INFORMATION Name: NPI: Address: Phone: **MEDICAL HISTORY** 1. **Member's Diagnosis:** Type 1 Gaucher Disease Other If requesting for Type 1 Gaucher Disease how was the diagnosis confirmed? *Please provide supporting documentation* Beta-glucosidase leukocyte (BGL) test 
Genetic Testing 3. Does the member have any of the following: please select all that apply ☐ Anemia Thrombocytopenia Bone Disease Hepatomegaly or splenomegaly 4. Please provide the following and laboratory reference range: Baseline hemoglobin level: Date taken:\_\_\_\_ Baseline platelet level:

Baseline spleen size: Date taken:\_\_\_\_\_ Date taken:\_\_\_\_\_ Baseline liver size: Date taken: For Cerdelga(eliglustat) Only: What type of CYP2D6 metabolizer is the member? (please provide documentation of an FDA-cleared test) Extensive Metabolizer

Intermediate Metabolizer



ember ID:			
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ease provide doc	umentation of why the mem	ber is not a candidate for o	enzyme replacement therapy:
	C	URRENT OR PREVIOUS	STHERAPV
Medication Name	1	Dates of Therapy	Status (Discontinued & Why or Current)
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