

**Request for Prior Authorization for Gaucher Disease: Enzyme Replacement and Substrate Reduction Therapy**  
**Website Form – [www.highmarkhealthoptions.com](http://www.highmarkhealthoptions.com)**  
**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:
  - A beta-glucosidase leukocyte (BGL) test
  - 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):
  - Anemia
    - Documentation of a baseline low hemoglobin level (laboratory reference range must be submitted)
  - Thrombocytopenia
    - Documentation of a baseline low platelet level (laboratory reference range must be submitted)
  - 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
  - Coverage is not provided for CYP2D6 ultra-rapid metabolizers.
- For Zavesca (miglustat):
  - Documentation the member is unable to use enzyme replacement therapy.
- **Initial Duration of Approval: 6 months**
- **Reauthorization criteria:**
  - 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)
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- **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 ID:	Member weight:	Height:

**DRUG INFORMATION**

Medication:	Strength:	
Directions:	Quantity:	Refills:
Is the member currently receiving requested medication? <input type="checkbox"/> Yes <input type="checkbox"/> 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? <input type="checkbox"/> Yes <input type="checkbox"/> 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**

- 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
- Does the member have any of the following: please select all that apply**  
 Anemia  
 Thrombocytopenia  
 Bone Disease  
 Hepatomegaly or splenomegaly
- Please provide the following and laboratory reference range:**  
 Baseline hemoglobin level: \_\_\_\_\_ Date taken: \_\_\_\_\_  
 Baseline platelet level: \_\_\_\_\_ Date taken: \_\_\_\_\_  
 Baseline spleen size: \_\_\_\_\_ 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

- Poor Metabolizer  
 Ultra-Rapid Metabolizer

**Member Name:** \_\_\_\_\_ **DOB:** \_\_\_\_\_  
**Member ID:** \_\_\_\_\_

**For Zavesca (miglustat) only:**  
**Please provide documentation of why the member is not a candidate for enzyme replacement therapy:**  
\_\_\_\_\_  
\_\_\_\_\_  
\_\_\_\_\_

**CURRENT OR PREVIOUS THERAPY**

Medication Name	Strength/Frequency	Dates of Therapy	Status (Discontinued & Why or Current)

**REAUTHORIZATION**

- Has the member tolerated therapy and shown improvement  Yes  No
- If the member has bone disease has the member shown improvement  Yes  No  N/A
- Please provide the following:
 

Current hemoglobin level: _____	Date taken: _____
Current platelet level: _____	Date taken: _____
Current spleen size: _____	Date taken: _____
Current liver size: _____	Date taken: _____

**SUPPORTING INFORMATION or CLINICAL RATIONALE**

\_\_\_\_\_  
\_\_\_\_\_  
\_\_\_\_\_  
\_\_\_\_\_

Prescribing Physician Signature	Date