

Page 1 of 2

New Patient Start: Prescriber/Physician Order Form



PATIENT INFORMATION		
Address City _	□ Male □ Female Date of Birth/ (mm/dd/yyyy)	
Language: English Other		
	Relationship to Patient	
Primary Contact Email	Dimensional Content Coll	
Primary Contact Phone	_ Primary Contact Cell	
Please attach copies of patient insurance and prescription cards—front and back.		
MEDICAL INFORMATION		
Diagnosis: Alpha-mannosidosis (ICD-10-CM E77.1 Defects in glycoprotein Allergies: None Specify Methods of Diagnosis (check all that apply): Methods of Diagnosis (check all that apply): Enzyme Assay Genetic Testing Tissue Biopsy Prior treatment and dose:	ast date of prior treatment and dose:	
Please attach copies of medical history/physical summary, urine oligosaccharides, acid α-mannosidase activity in leukocytes, current medications, genetic testing results, and allergies.		
LAMZEDE [®] (VELMANASE ALFA-TYCV) PRESCRIPTION		
Frequency Number of Refills *The recommended dosage is 1 mg/kg (actual body weight) administered once weekly as an intravenous infusion. Please list any additional treatment information, including follow-up evaluations: PRESCRIBER/OFFICE INFORMATION		
	_ Practice/Group Name	
	Suite Suite	
Office Contact Person		
	_ Office Fax	
List of Facilities Where Physician Has Privileges		
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Please continue on page 2. Please see Important Safety Information for Lamzede, including Boxed Warning for Severe Hypersensitivity Reactions, on <u>page 2</u> and <u>Full Prescribing Information</u>.



Page 2 of 2

New Patient Start: Prescriber/Physician Order Form (cont'd)



SITE OF SERVICE

Preferred Acquisitions Channe	əl:	
Buy and Bill (If site does not allow white bagging, send the form directly to EVERSANA [®] .)	Specialty Pharmacy to Bill Please send the prescription directly to:	□ Other
	EVERSANA Life Science Services PH: 1-855-282-4883 FAX: 1-855-929-2828	
Preferred Site of Infusion:		
Site of Infusion		
Contact Person		

Indication

Lamzede[®] (velmanase alfa-tycv) is indicated for the treatment of non-central nervous system manifestations of alpha-mannosidosis in adult and pediatric patients.

Important Safety Information

WARNING: SEVERE HYPERSENSITIVITY REACTIONS

Hypersensitivity Reactions Including Anaphylaxis

Patients treated with Lamzede have experienced hypersensitivity reactions, including anaphylaxis. Appropriate medical support measures, including cardiopulmonary resuscitation equipment, should be readily available during Lamzede administration. If a severe hypersensitivity reaction (e.g., anaphylaxis) occurs, discontinue Lamzede immediately and initiate appropriate medical treatment. In patients with severe hypersensitivity reaction, a desensitization procedure to Lamzede may be considered.

Considerations Due to Hypersensitivity Reactions and/or Infusion-Associated Reactions (IARs)

Prior to Lamzede administration, consider pretreating with antihistamines, antipyretics, and/or corticosteroids. Inform patients and caregivers of the signs and symptoms of hypersensitivity reactions and IARs and instruct them to seek medical care immediately if such symptoms occur.

- If a severe hypersensitivity reaction (including anaphylaxis) or severe IAR occurs, immediately discontinue Lamzede administration and initiate appropriate medical treatment.
- In the event of a mild to moderate hypersensitivity reaction or a mild to moderate IAR, consider temporarily holding the infusion for 15 to 30 minutes, slowing the infusion rate to 25% to 50% of the recommended rate, and initiating appropriate medical treatment.

Hypersensitivity Reactions Including Anaphylaxis

Anaphylaxis and severe hypersensitivity signs and symptoms included cyanosis, hypotension, emesis, urticaria, erythema, facial swelling, pyrexia, and tremor.

Infusion-Associated Reactions (IARs)

The most frequent symptoms of IARs that occurred in >10% of the population were pyrexia, chills, erythema, vomiting, cough, urticaria, rash, and conjunctivitis.

Females of Reproductive Potential

Advise females of reproductive potential to use effective contraception during treatment and for 14 days after the last dose if Lamzede is discontinued. For females of reproductive potential, verify that the patient is not pregnant prior to initiating treatment with Lamzede.

Embryo-Fetal Toxicity

Based on findings from animal reproduction studies, Lamzede may cause embryo-fetal harm when administered to a pregnant female.

Common Adverse Reactions

The most common adverse reactions (incidence >20%) are hypersensitivity reactions including anaphylaxis, nasopharyngitis, pyrexia, headache, and arthralgia.

Please see Full Prescribing Information for Lamzede.