Guidance for the Sample Order Form

The sample order form on page 2 is for the sole purpose of providing a template that can be used to order NEXVIAZYME (avalglucosidase alfa-ngpt). If you choose to use this sample order form, please copy and paste the information on page 2.

IMPORTANT SAFETY INFORMATION AND INDICATION

INDICATION

NEXVIAZYME (avalglucosidase alfa-ngpt) is used for the treatment of patients 1 year of age and older with late-onset Pompe disease [lysosomal acid alpha-glucosidase (GAA) deficiency].

IMPORTANT SAFETY INFORMATION

WARNING: SEVERE HYPERSENSITIVITY REACTIONS, INFUSION-ASSOCIATED REACTIONS, and RISK OF ACUTE CARDIORESPIRATORY FAILURE IN SUSCEPTIBLE PATIENTS

Hypersensitivity Reactions Including Anaphylaxis: Patients treated with NEXVIAZYME have experienced lifethreatening hypersensitivity reactions, including anaphylaxis. Appropriate medical support measures, including cardiopulmonary resuscitation equipment, should be readily available during NEXVIAZYME administration. If a severe hypersensitivity reaction (e.g., anaphylaxis) occurs, NEXVIAZYME should be discontinued immediately and appropriate medical treatment should be initiated. In patients with severe hypersensitivity reaction, a desensitization procedure to NEXVIAZYME may be considered.

Infusion-Associated Reactions (IARs): Patients treated with NEXVIAZYME have experienced severe IARs. If severe IARs occur, consider immediate discontinuation of NEXVIAZYME, initiation of appropriate medical treatment, and the benefits and risks of readministering NEXVIAZYME following severe IARs. Patients with an acute underlying illness at the time of NEXVIAZYME infusion may be at greater risk for IARs. Patients with advanced Pompe disease may have compromised cardiac and respiratory function, which may predispose them to a higher risk of severe complications from IARs.

Risk of Acute Cardiorespiratory Failure in Susceptible
Patients: Patients susceptible to fluid volume overload, or
those with acute underlying respiratory illness or
compromised cardiac or respiratory function for whom
fluid restriction is indicated may be at risk of serious
exacerbation of their cardiac or respiratory status during
NEXVIAZYME infusion. More frequent monitoring of vitals
should be performed during NEXVIAZYME infusion.

WARNINGS AND PRECAUTIONS

Hypersensitivity Reactions Including Anaphylaxis: See Boxed WARNING. Prior to NEXVIAZYME administration, consider pretreating with antihistamines, antipyretics, and/or corticosteroids. The risks and benefits of readministering NEXVIAZYME following severe hypersensitivity reaction (including anaphylaxis) should be considered. If a mild or moderate hypersensitivity reaction occurs, the infusion rate may be slowed or temporarily stopped.

Infusion-Associated Reactions: See Boxed WARNING. IARs may still occur in patients after receiving pretreatment. If mild or moderate IARs occur regardless of pretreatment, decreasing the infusion rate or temporarily stopping the infusion may ameliorate the symptoms.

Risk of Acute Cardiorespiratory Failure in Susceptible Patients: See Boxed WARNING.

ADVERSE REACTIONS

The most common adverse reactions (>5%) were headache, fatigue, diarrhea, nausea, arthralgia, dizziness, myalgia, pruritus, vomiting, dyspnea, erythema, paresthesia and urticaria.

Please see full <u>Prescribing Information</u> for complete details, including **Boxed WARNING**.

Sample Order Form

It is recommended that the following information, at a minimum, be included in the physician order to the pharmacy and infusion staff.

| Patient Nar | me: | Diagnosis: | |
|---------------------------------|--------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|----------------------------------------------------------------------------------------------------------------------------------------------------------------------------|------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| | | Patient Weight (kg | |
| | te: | | |
| Notify Dr | | _at phone/pager | when patient arrives. |
| 2) Obtain 3) Prepare | 20 mg/kg (for patients we total volume of mL. 40 mg/kg (for patients we total volume of mL. onitor vital signs during NE commended that patients se | eighing ≥30 kg) avalglucosidase eighing <30 kg) avalglucosidase XVIAZYME infusion as prescri susceptible to fluid volume ov mised cardiac or respiratory f | e alfa-ngpt in 5% dextrose injection, to a e alfa-ngpt in 5% dextrose injection, to a bed by the healthcare provider. It is verload, or those with acute underlying function for whom fluid restriction is |
| 4) Pre-tre | eatment medication (if pres | cribed by the treating physicia | an): |
| proteir a) b) c) d) | Begin the infusion at rate If no signs of IARs Increase the infusion rate If no signs of IARs Increase the infusion rate If no signs of IARs Increase the infusion rate If no signs of IARs Increase the infusion rate remainder of the infusion for at least 30 minutes. If If 5-step process, increase this rate for the remainder | of 1 mg/kg/hr (mL/hr) and tomg/kg/hr (mL/hr) tomg/kg/hr (mL/hr) tomg/kg/hr (mL/hr) if 4-step process. If 5-step process of IARs the infusion rate tomg/er of the infusion. | manner. Use a 0.2 µm in-line, low and administer for at least 30 minutes. and administer for at least 30 minutes and administer for at least 30 minutes and administer at this rate for the ocess, this rate should be administered kg/hr (mL/hr) and administer at |
| - | nfusion line with 5% dextro stered. Do not IV push the | | n rate to ensure the entire dose is |
| | vital signs h ged observation times. | nours after completion of the i | infusion. Some patients may require |
| Contact Dr. associated | | immediately in the event of | f a hypersensitivity reaction or infusion- |
| | uspected adverse reaction or www.fda.gov/medwatch | | -1610, option 1 or the FDA at 1-800- |
| Physician's Phone/Pag | Signature: | Date: | |

Please see full <u>Prescribing Information</u> for complete details, including **Boxed WARNING.**