

Kanuma

Prior Authorization Request

CVS Caremark administers the prescription benefit plan for the patient identified. This patient's benefit plan requires prior authorization for certain medications in order for the drug to be covered. To make an appropriate determination, providing the most accurate diagnosis for the use of the prescribed medication is necessary. Please respond below and fax this form to CVS Caremark toll-free at 1-855-330-1720. If you have questions regarding the prior authorization, please contact CVS Caremark at 1-888-877-0518. For inquiries or questions related to the patient's eligibility, drug copay or medication delivery; please contact the Specialty Customer Care Team: CaremarkConnect® 1-800-237-2767.

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Patient's Name:	Date:
Patient's ID:	
Physician's Name:	
Specialty:	
Physician Office Telephone:	
<u>Referring</u> Provider Info: □ Same as Reque Name:	8
Fax:	Phone:
	ring Provider 🛛 Same as Requesting Provider
Name:	NPI#:
Fax:	Phone:

Approvals may be subject to dosing limits in accordance with FDA-approved labeling, accepted compendia, and/or evidence-based practice guidelines.

<u>Required Demographic Information:</u>

Patient Weight: _____kg Patient Height: cm

What is the ICD-10 code?

Site of Service Questions:

A. Where will this drug be administered? Ambulatory surgical, *skip to Clinical Ouestions* □ Off-campus Outpatient Hospital □ Physician office, *skip to Clinical Questions*

□ Home infusion, *skip to Clinical Questions* On-campus Outpatient Hospital □ Pharmacy, *skip to Clinical Questions*

- B. Is this request to continue previously established treatment with the requested medication? \Box Yes \rightarrow This is a continuation of an existing treatment. \Box No \rightarrow This is a new therapy request (patient has not received requested medication in the last 6 months). *skip to* Clinical Criteria Questions
- C. Has the patient experienced an adverse event with the requested product that has not responded to conventional interventions (eg acetaminophen, steroids, diphenhydramine, fluids, other pre-medications or slowing of the infusion rate) or a severe adverse event (anaphylaxis, anaphylactoid reactions, myocardial infarction, thromboembolism, or seizures) during or immediately after an infusion? ACTION REQUIRED: If Yes, Attach supporting clinical *documentation.* \Box Yes, *skip to Clinical Criteria Questions* \Box No

Send completed form to: Case Review Unit CVS Caremark Specialty Programs Fax: 1-855-330-1720 Note: This fax may contain medical information that is privileged and confidential and is solely for the use of individuals named above. If you are not the intended recipient you hereby are advised that any dissemination, distribution, or copying of this communication is prohibited. If you have received the fax in error, please immediately notify the sender by telephone and destroy the original fax message. Kanuma SGM 2094-A - 08/2023.

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- D. Does the patient have laboratory confirmed sebelipase alfa antibodies? ACTION REQUIRED: If Yes, Attach supporting clinical documentation. Use, skip to Clinical Criteria Questions No
- E. Is the patient medically unstable which may include respiratory, cardiovascular, or renal conditions that may limit the member's ability to tolerate a large volume or load or predispose the member to a severe adverse event that cannot be managed in an alternate setting without appropriate medical personnel and equipment?
 ACTION REQUIRED: If Yes, Attach supporting clinical documentation.
 Yes, skip to Clinical Criteria Questions
 No
- F. Does the patient have severe venous access issues that require the use of special interventions only available in the outpatient hospital setting? ACTION REQUIRED: If Yes, Attach supporting clinical documentation.
 □ Yes, skip to Clinical Criteria Questions □ No
- G. Does the patient have significant behavioral issues and/or physical or cognitive impairment that would impact the safety of the infusion therapy AND the patient does not have access to a caregiver? *ACTION REQUIRED: If Yes, Attach supporting clinical documentation.* □ Yes □ No

Clinical Criteria Questions:

1. What is the diagnosis?

Lysosomal acid lipase (LAL) deficiency (*If checked, go to 2*)

- □ Other, please specify. _____ (*If checked, go to 2*)
- 2. Is this a request for continuation of therapy with the requested medication?

□ Yes, Continue to 3

□ No, Continue to 4

3. Is the patient responding to therapy (e.g., improvement, stabilization, or slowing of disease progression for weight-for-age z-score if exhibiting growth failure, low-density lipoprotein [LDL], high-density lipoprotein [HDL], triglycerides, or alanine aminotransferase [ALT])? *ACTION REQUIRED*: If yes, attach lab results or chart notes documenting a positive response to therapy (e.g., improvement, stabilization, or slowing of disease progression for weight-for-age z-score if exhibiting growth failure, LDL, HDL, triglycerides, or ALT). **D** Yes, *No Further Questions*

□ No, No Further Questions

4. Was the diagnosis confirmed by enzyme assay demonstrating a deficiency of lysosomal acid lipase enzyme activity OR by genetic testing? *ACTION REQUIRED*: If yes, attach lysosomal acid lipase enzyme assay or genetic testing results supporting diagnosis.

□ Yes, Continue to 5

□ No, Continue to 5

5. Does the patient have an alanine aminotransferase (ALT) level greater than or equal to 1.5 times the upper limit of normal (based on the age and gender-specific normal ranges) on two consecutive ALT measurements obtained at least one week apart?

□ Yes (*If checked, no further questions*)

□ No [ALT is less than 1.5 times the upper limits of normal (based on the age- and gender-specific normal ranges)] (*If checked, no further questions*)

Unknown (*If checked, no further questions*)

I attest that this information is accurate and true, and that documentation supporting this information is available for review if requested by CVS Caremark or the benefit plan sponsor.

Х

Prescriber or Authorized Signature

Date (mm/dd/yy)

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