

Member Name: {{MEMFIRST}} {{MEMLAST}} DOB: {{MEMBERDOB}} PA Number: {{PANUMBER}}



{{PANUMCODE}}

Ravicti

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 1-866-249-6155.** If you have questions regarding the prior authorization, please contact CVS Caremark at 1-866-814-5506. 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: {{MEMFIRST}} {{MEMLAST}} **Date:** {{TODAY}}
Patient's ID: {{MEMBERID}} **Patient's Date of Birth:** {{MEMBERDOB}}
Physician's Name: {{PHYFIRST}} {{PHYLAST}}
Specialty: _____, **NPI#:** _____
Physician Office Telephone: {{PHYSICIANPHONE}} **Physician Office Fax:** {{PHYSICIANFAX}}
Request Initiated For: {{DRUGNAME}}

1. What is the patient's diagnosis?
 Urea cycle disorder
 Other _____
2. What is the ICD-10 code? _____
3. Is the product being requested for the treatment of urea cycle disorders? Yes No *If No, skip to #8*
4. The preferred product for your patient's health plan is sodium phenylbutyrate. Can the patient's treatment be switched to the preferred product? *If Yes, please call 1-866-814-5506 to have the updated form faxed to your office OR you may complete the PA electronically (ePA). You may sign up online via CoverMyMeds at: www.covermymeds.com/epa/caremark/ or call 1-866-452-5017.* Yes No
5. Does the patient have documented uncontrolled congestive heart failure, uncontrolled hypertension, or severe renal impairment (i.e., creatinine clearance less than 30 mL/min) and is on a documented sodium-restricted diet? **ACTION REQUIRED: If Yes, attach supporting chart note(s) and skip to #8.** Yes No
6. Does the patient have a documented inability to ingest a sufficient amount of the preferred product as prescribed due to an aversion to the taste or smell? **ACTION REQUIRED: If Yes, attach supporting chart note(s) and skip to #8.** Yes No
7. Does the patient have a documented inability to tolerate the necessary pill burden with the preferred product? **ACTION REQUIRED: If Yes, attach supporting chart note(s).** Yes No *If No, complete this form in its entirety and State Step Therapy section.*
8. Will Ravicti be used for chronic management of urea cycle disorder? Yes No
9. Was the diagnosis confirmed by enzymatic, biochemical, or genetic testing? **ACTION REQUIRED: If Yes, attach supporting chart note(s).** Yes No
10. Is this request for continuation of treatment with Ravicti? Yes No *If No, no further questions.*
11. Is the patient experiencing a reduction in plasma ammonia levels from baseline? Yes No

Send completed form to: Case Review Unit, CVS Caremark Prior Authorization. Fax: 1-866-249-6155

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. Ravicti State Step, ACSF SGM - 1/2021.

CVS Caremark Prior Authorization • 1300 E. Campbell Road • Richardson, TX 75081

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Member Name: {{MEMFIRST}} {{MEMLAST}} **DOB:** {{MEMBERDOB}} **PA Number:** {{PANUMBER}}

State Step Therapy

1. Is the requested drug being used for an FDA-approved indication or an indication supported in the compendia of current literature (examples: AHFS, Lexicomp, Clinical Pharmacology, Micromedex, current accepted guidelines)? Yes No
2. Does the prescribed quantity fall within the manufacturer's published dosing guidelines or within dosing guidelines found in the compendia of current literature (examples: package insert, AHFS, Lexicomp, Clinical Pharmacology, Micromedex, current accepted guidelines)? Yes No
3. Does the patient reside in Maryland? Yes No *If No, skip to #7*
4. Is the alternate drug (sodium phenylbutyrate) FDA-approved for the medical condition being treated?
 Yes No *If No, no further questions.*
5. Has the prescriber provided proof, documented in the patient's chart notes, indicating that the requested drug was ordered for the patient in the past 180 days? Yes No *If No, skip to #7*
6. Has the prescriber provided proof, documented in the patient chart notes, that in their opinion the requested drug is effective for the patient's condition? Yes No *No further questions*
7. Are any of the following conditions met for the alternate drug (sodium phenylbutyrate)?
If Yes, indicate below and no further questions.
 - The alternate drug is contraindicated
 - The alternate drug is likely to cause an adverse reaction, physical or mental harm
 - The alternate drug is expected to be ineffective
 - The alternate drug was previously tried or a drug in the same class or with the same action was previously tried and was stopped due to ineffectiveness or an adverse event
 - The alternate drug is not in the patient's best interest
 - The alternate drug was tried while covered by the current or the previous health benefit plan
 - None of the above, *continue to #8*
8. Is the patient stable or currently receiving a positive therapeutic outcome with the requested drug and a change in the prescription drug is expected to be ineffective or cause harm to the patient? Yes No

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.

X _____

Prescriber or Authorized Signature

Date (mm/dd/yy)

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