SENTARA HEALTH PLANS

PHARMACY PRIOR AUTHORIZATION/STEP-EDIT REQUEST*

<u>Directions</u>: <u>The prescribing physician must sign and clearly print name (preprinted stamps not valid) on this request</u>. All other information may be filled in by office staff; **fax to 1-800-750-9692**. No additional phone calls will be necessary if all information (including phone and fax #s) on this form is correct. <u>If information provided is not complete, correct, or legible, authorization may be delayed.</u>

<u>Drug Requested</u>: **Ravicti**[®] (glycerol phenylbutyrate)

MEMBER & PRESCRIBER INFORMATION: Authorization may be delayed if incomplete.			
Meml	ber Name:		
Member Sentara #:			
Presci	riber Name:		
Prescriber Signature:			
Office	e Contact Name:		
Phone Number:		Fax Number:	
DEA	OR NPI #:		
DRU	UG INFORMATION: Authoriz	zation may be delayed if incomplete.	
Drug	Form/Strength:		
		Length of Therapy:	
Diagnosis:		ICD Code:	
Weight:		Date:	
<u>Quan</u>	tity Limits: 17.5 mL (19 grams) per	day	
suppo		low all that apply. All criteria must be met for approval. To tion, including lab results, diagnostics, and/or chart notes, must be	
Initi	ial Authorization: 12 months		
	Prescriber is a specialist in the man	agement of urea cycle disorders	
	Member is 2 months of age or older has been noted by provider	r and current weight: and height:	
		of chronic hyperammonemia due to a urea cycle disorder (UCD) as iochemical testing (submit labs confirming diagnosis)	
	Member does NOT have a diagnost	is of UCD with N-acetylglutamate synthase (NAGS) deficiency	
	Ravicti will NOT be used in treatm	nent of acute hyperammonemia	

	Member has had a 30-day trial and failure of a sodium phenylbutyrate product (generic Buphenyl [®] , Pheburane [®] , Olpruva [™]) as documented by ONE of the following:			
	□ Fasting ammonia level > 0.5 times the upper limit of normal while compliantly taking a sodium phenylbutyrate product (generic Buphenyl [®] , Pheburane [®] , Olpruva [™]) (submit labs for documentation)			
	☐ Member has a history of intolerance to a sodium phenylbutyrate product (generic Buphenyl [®] , Pheburane [®] , Olpruva [™]) (submit chart notes documenting clinically significant medication intolerance and completed Med Watch form)			
	Member will be maintained on a protein restricted diet while using Ravicti® therapy			
	Members with moderate to severe hepatic impairment (Child-Pugh score B or C) will be initiated on 4.5 mL/m²/day (submit current labs including albumin, PT/INR and total bilirubin)			
	Does the member have some residual enzyme activity?			
	• If yes, member must be initiated on 4.5 mL/m²/day and titrated according to guidelines			
Reauthorization: 12 months. Check below all that apply. All criteria must be met for approval. To support each line checked, all documentation, including lab results, diagnostics, and/or chart notes, must be provided or request may be denied.				
	Member has been maintained a protein restricted diet while using Ravicti® therapy			
	Member's current weight: and height: must be noted			
	Member has a documented positive clinical response to Ravicti® therapy and fasting ammonia levels have normalized since last approval of Ravicti® (submit chart notes and labs to support positive)			
Medication being provided by Specialty Pharmacy – Proprium Rx				
	Use of samples to initiate therapy does not meet step edit/ preauthorization criteria.**			
" <u>Pre</u>	*Previous therapies will be verified through pharmacy paid claims or submitted chart notes. *			

^{*}Approved by Pharmacy and Therapeutics Committee: 3/21/2019; 11/18/2022; 9/21/2023 REVISED/UPDATED/REFORMATTED: 5/15/2019; 11/30/2022; 10/17/2023