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)</u> on this request. All other information may be filled in by office staff; <u>fax to 1-800-750-9692</u>. No additional phone calls will be necessary if all information <u>(including phone and fax #s)</u> on this form is correct. <u>If the information provided is not complete, correct, or legible, the authorization process may be delayed.</u>

Sodium Phenylbutyrate Products

<u>Drug Requested</u> : (select one from below)				
□ sodium phenylbutyrate (Buphenyl®) □ Powder □ Tablets	 Pheburane[®] (sodium phenylbutyrate) oral pellets 	□ Olpruva [™] (sodium phenylbutyrate) oral suspension		
MEMBER & PRESCRIBE	R INFORMATION: Authoriza	tion may be delayed if incomplete.		
Member Name:				
		Date of Birth:		
Prescriber Name:				
Prescriber Signature:		Date:		
Office Contact Name:				
Phone Number:	ne Number: Fax Number:			
DEA OR NPI #:				
DRUG INFORMATION: A	authorization may be delayed if incor	nplete.		
Drug Form/Strength:				
	Length of Therapy:			
Diagnosis:	ICD Code, if applicable:			
Weight:	Date:			
Quantity Limits: Maximum dail	y dose of 20 grams per day for all form	alations		
	neck below all that apply. All criteria amentation, including lab results, dia			
Initial Authorization: 12 m	onths			
☐ Prescriber is a specialist in the	he management of urea cycle disorde	ers		
☐ Member's current weight:	and height:	must be noted		

PA Buphenyl_1	Pheburane (Phar	macy)(CORE
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	Member has a confirmed diagnosis of chronic hyperammonemia due to a urea cycle disorder (UCD) amenable to treatment with sodium phenylbutyrate as verified by genetic, enzymatic or biochemical testing (submit labs confirming diagnosis)	
	Member does NOT have a diagnosis of UCD with N-acetylglutamate synthase (NAGS) deficiency	
	Sodium phenylbutyrate will NOT be used in treatment of acute hyperammonemia	
	Member will be maintained on a protein restricted diet while using sodium phenylbutyrate therapy	
	Member's blood ammonia levels, CBC with differential, hepatic and renal function will be monitored regularly while using this medication	
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 on a protein restricted diet while using sodium phenylbutyrate therapy	
	Member's current weight: and height: must be noted	
	Member has a documented positive clinical response to therapy and fasting ammonia levels have normalized since last approval of requested medication (chart notes and/or labs must be submitted)	
Medication being provided by Specialty Pharmacy - PropriumRx		
**	Use of samples to initiate therapy does not meet step edit/ preauthorization criteria.**	

*Previous therapies will be verified through pharmacy paid claims or submitted chart notes. *