## 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 <u>1-800-750-9692</u>. 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>

**Drug Requested:** carglumic acid (Carbaglu®)

MEMBER & PRESCRIBER IN	NFORMATION: Authorization may be delayed if incomplete.
Member Name:	
	Date of Birth:
Prescriber Name:	
	Date:
Office Contact Name:	
	Fax Number:
DEA OR NPI #:	
DRUG INFORMATION: Autho	
Drug Form/Strength:	
	Length of Therapy:
Diagnosis:	ICD Code, if applicable:
Weight:	Date:
<b>Recommended Dosage:</b>	
<ul> <li>NAGS deficiency, acute hyperan</li> </ul>	nmonemia: 100 to 250 mg/kg/day given in 2 to 4 divided doses
<ul> <li>NAGS deficiency, chronic hyper</li> </ul>	ammonemia: 10 to 100 mg/kg/day given in 2 to 4 divided doses
<ul> <li>Propionic acidemia or methylma doses (12 hours apart) and for a r</li> </ul>	lonic acidemia, acute hyperammonemia: Oral: 3.3g/m²/day in 2 divided maximum of 7 days
	below all that apply. All criteria must be met for approval. To ation, including lab results, diagnostics, and/or chart notes, must be
□ N-acetylglutamate synthase (I	NAGS) deficiency
<b>Initial Authorization:</b> 6 months	
☐ Provider is or has consulted with cycle disorders	a specialist in medical genetics or other specialist in treatment of urea
☐ Member has diagnosis of NAGS	deficiency as confirmed by genetic testing (submit results)

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	Member is experiencing hyperammonemia despite compliance with standard therapy (submit current plasma ammonia lab test results and chart notes documenting therapies tried)
	For treatment of acute hyperammonemia, carglumic acid will be used in conjunction with standard therapy (i.e. hemodialysis, intravenous sodium benzoate and phenylacetate, protein restriction)
	Prescribed dose will not exceed 250 mg/kg per day initially, followed by a maintenance dose of 100 mg/kg per day
	For approval of brand name Carbaglu: Member has had trial and intolerable life-endangering adverse event with generic carglumic acid tablets (must submit completed MedWatch form and chart notes to document adverse event)
<b>.</b>	N-acetylglutamate synthase (NAGS) deficiency
Rea	authorization: 12 months.
	All initial authorization criteria continues to be met
	Member's plasma ammonia levels have been sustained at or below normal limits for age (submit current lab test results)
	Member is <b>NOT</b> experiencing any symptoms of unacceptable toxicity associated with carglumic acid
	For approval of brand name Carbaglu: Member has had trial and intolerable life-endangering adverse event with generic carglumic acid tablets (must submit completed MedWatch form and chart notes to document adverse event)
	Propionic Acidemia (PA) or Methylmalonic Acidemia (MMA) with acute hyperammonemia
<b>A</b> u	thorization Criteria: 7 day length of authorization. Coverage cannot be renewed.
	Provider is or has consulted with a specialist in medical genetics or other specialist in treatment of urea cycle disorders
	Member has diagnosis of propionic acidemia or methylmalonic acidemia as confirmed by genetic testing (submit results)
	Member's plasma ammonia level is $\geq 70~\mu mol/L$ despite standard of care treatment, such as intravenous hydration and nutritional support (submit current plasma ammonia lab test results and chart notes documenting therapies tried)
	Medication will be used in conjunction with other ammonia-lowering therapies (i.e. intravenous glucose, insulin, L-carnitine, protein restriction, hemodialysis)
	Medication will only be used until the patient's ammonia level is $<$ 50 $\mu$ mol/L and for a maximum duration of 7 days
	For approval of brand name Carbaglu: Member has had trial and intolerable life-endangering adverse event with generic carglumic acid tablets (must submit completed MedWatch form and chart notes to

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document adverse event)

## 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. \*