SENTARA COMMUNITY PLAN (MEDICAID)

MEDICAL PRIOR AUTHORIZATION/STEP-EDIT REQUEST*

<u>Directions:</u> The prescribing physician must sign and clearly print name (preprinted stamps not valid) on this request. All other information may be filled in by office staff; <u>fax to 1-844-305-2331</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 can be delayed</u>.

<u>Drug Requested</u>: Empaveli® (pegcetacoplan) SQ (J3490) (Medical)

Paroxysmal Nocturnal Hemoglobinuria (PNH)

Member Name:		
Member Sentara #:		
Prescriber Name:		
	Date:	
Office Contact Name:		
Phone Number:		
DEA OR NPI #:		
DRUG INFORMATION: Autho	rization may be delayed if incomplete.	
Drug Form/Strength:		
	Length of Therapy:	
Diagnosis:	ICD Code, if applicable:	
	Date:	

Max quantity limits:

- 8 (eight) SQ infusions every 28 days
- Empaveli® 1080 mg/20 mL solution in single-use vials for injection supplied in 8-count cartons

Recommended Dosage:

- Maintenance 1080 mg twice weekly
- Dosage Adjustment: For lactate dehydrogenase (LDH) levels > 2 levels ULN, adjust pegcetacoplan dosing regimen to 1080 mg every 3 days. Monitor LDH twice weekly for at least 4 weeks after a dose increase.

(Continued on next page)

CLINICAL CRITERIA: 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.

Initi	al A	uthorization: 6 months			
	Med	ication must be prescribed by or in consultation with a hematologist or nephrologist			
	Prescriber must be enrolled in the Empaveli® Risk Evaluation and Mitigation Strategy (REMS) program				
	Member must meet ONE of the following:				
	□ E	Empaveli [®] will be used as switch therapy <u>AND</u> member meets <u>ALL</u> the following:			
		Member failed Soliris® or Ultomiris® and must meet renewal criteria			
		Member does <u>NOT</u> have a systemic infection			
		Member must be vaccinated against encapsulated bacteria (Streptococcus pneumoniae, Neisseria meningitidis, and Haemophilus influenzae type B) at least two weeks prior to initiation of Empaveli® therapy and revaccinated according to current medical guidelines for vaccine use			
	L	Empaveli [®] will <u>NOT</u> be used in combination with other complement inhibitor therapies (e.g., Ultomiris [®] , Soliris [®] or Fabhalta [®])			
		OR			
	□ N	Member is treatment-naive AND member meets ALL the following:			
		Member must have a diagnosis of Paroxysmal Nocturnal Hemoglobinuria (PNH) confirmed by detection of PNH clones of at least 10% by flow cytometry testing (must submit labs)			
		Flow cytometry pathology report must demonstrate at least two (2) different glycosylphosphatidylinositol (GPI) protein deficiencies (e.g., CD55, CD59, etc.) within two (2) different cell lines from granulocytes, monocytes, erythrocytes (must submit labs)			
		Member has laboratory evidence of significant intravascular hemolysis (i.e. LDH \geq 1.5 x ULN) AND has experienced ONE of the following additional indications for therapy (must submit chart notes and labs):			
		☐ Member is transfusion dependent (defined by having a transfusion within the last 12 months) and symptomatic anemia			
		☐ Presence of a thrombotic event (e.g., DVT, PE)			
		☐ Presence of organ damage secondary to chronic hemolysis (i.e. renal insufficiency, pulmonary insufficiency, or hypertension)			
		☐ Member is pregnant and potential benefit outweighs potential fetal risk			
		 Member has abdominal pain requiring admission to hospital 			
		Member does <u>NOT</u> have a systemic infection			
		Member must be administered a meningococcal vaccine at least two weeks prior to initiation of Empaveli® therapy and revaccinated according to current medical guidelines for vaccine use			
		Empaveli [®] will <u>NOT</u> be used in combination with other complement inhibitor therapies (e.g., Ultomiris [®] , Soliris [®] or Fabhalta [®])			

provi	ded or request may be det	nied.	luding lab results, diagnostics, and/or chart notes, must be
	Provider attests to an absinfections [septicemia ar	*	ole toxicity from the drug (e.g. serious meningococcal nfusion reactions)
	-	-	esponse indicated by at least <u>ONE</u> of the following (check all document improvement):
	☐ Decrease in serum L	DH	
	☐ Stabilization/increase	e in hemoglobin lev	/el
	☐ Decrease in packed I	RBC transfusion red	quirement
	☐ Reduction in thromb	oembolic events	
Med	ication being provid	ed by (check bo	x below that applies):
Med	ication being provid Physician's office	ed by (check bo	x below that applies): □ Specialty Pharmacy – Proprium Rx

**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. *

Reauthorization: 6 months. Check below all that apply. All criteria must be met for approval. To