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>.

IV Eculizumab Products - Paroxysmal Nocturnal Hemoglobinuria (PNH) (Medical)

Drug Requested: select one drug b	pelow	
□ Bkemv [®] (eculizumab-aeeb) Q5152	□ Epysqli® (eculizumab-aagh) Q5151	□ Soliris [®] (eculizumab) J1299
MEMBER & PRESCRIBER	INFORMATION: Authorization	n may be delayed if incomplete.
Member Name:		
Member Sentara #:	Date of Birth:	
Prescriber Name:		
Prescriber Signature:		Date:
Office Contact Name:		
Phone Number:	Fax Number:	
NPI #:		
DRUG INFORMATION: Au	thorization may be delayed if incomp	lete.
Drug Name/Form/Strength:		
Dosing Schedule:	Length of Therapy:	
Diagnosis:	gnosis: ICD Code, if applicable:	
Weight (if applicable):	Date w	eight obtained:
	is box, the timeframe does not jeopard maximum function and would not sub	

(Continued on next page)

Recommended Dosage:

- Maximum Quantity Limit 4 vials every 14 days; one 300 mg vial (30 mL) = 150 billable units [1 billable unit per 2 mg]
 - IV Induction 600 mg weekly for 4 doses
 - o Maintenance 900 mg at week 5, then 900 mg every 2 weeks thereafter

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.

Initial Authorization: 6 months			
	Medication must be prescribed by or in consultation with a hematologist or nephrologist		
	Prescriber must be enrolled in the Soliris® Risk Evaluation and Mitigation Strategy (REMS) program		
	Member must be 18 years of age or older		
	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 2 different glycosylphosphatidylinositol (GPI) protein deficiencies (e.g., CD55, CD59, etc.) within 2 different cell lines from granulocytes, monocytes, erythrocytes (must submit labs)		
	Member must have <u>ONE</u> of the following indications for therapy (must submit chart notes and labs): ☐ Member is transfusion dependent as defined by having a transfusion within the last 12 months and <u>ONE</u> of the following:		
	☐ Member's hemoglobin is less than or equal to 7 g/dL		
	☐ Member has symptoms of anemia and the hemoglobin is less than or equal to 9 g/dL		
	□ Member has high lactate dehydrogenase (LDH) level (defined as ≥ 1.5 times the upper limit of the normal range with clinical symptoms)		
	☐ Presence of a thrombotic event (e.g., DVT, PE)		
	☐ Presence of organ damage secondary to chronic hemolysis		
	☐ Member is pregnant and potential benefit outweighs potential fetal risk		
	For Bkemv [®] and Soliris [®] requests: Member must have documentation of an inadequate response, contraindication or intolerance to <u>BOTH</u> of the following:		
	☐ Ultomiris [™] (ravulizumab) (*requires prior authorization)		
	☐ Epysqli® (eculizumab-aagh) (*requires prior authorization)		
	Member does NOT have a systemic infection		
	Member must meet ONE of the following:		
	Member must be administered a meningococcal vaccine at least two weeks prior to initiation of Soliris® therapy and revaccinated according to current medical guidelines for vaccine use		
	☐ Member has NOT received a meningococcal vaccination at least two weeks prior to the initiation		

(Continued on next page)

risks of developing a meningococcal infection

of therapy with eculizumab and documented the risks of delaying eculizumab therapy outweigh the

(Continued from previous page)

Medication will NOT be prescribed concurrently with another FDA approved product prescribed for
treatment of PNH (e.g., Bkemv [™] , Epysqli [™] , PiaSky [®] , Ultomiris [®] , Empaveli [®] , or Fabhalta [®])

Reauthorization: 6 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.

 Provider attests to an absence of unacceptable toxicity from the drug (e.g., serious meningococcal infections (septicemia and/or meningitis), infusion reactions, serious infections) Member has experienced positive disease response indicated by at least <u>ONE</u> of the following (check

- □ Documentation of a recent (within 3 months) LDH level that shows a reduction from baseline
 □ Documentation that the member has stabilized hemoglobin levels as supported by ONE of the
- □ Documentation that the member has stabilized hemoglobin levels as supported by **ONE** of the following:
 - ☐ Member had a reduction in number of transfusions **OR** units of packed red cells transfused from baseline
 - ☐ Member maintained a hemoglobin concentration above 7 g/dL **OR** maintained a hemoglobin concentration above 9 g/dL if member had a baseline hemoglobin level above 7 g/dL but below 9 g/dL
- ☐ Member had a reduction in thrombotic events (e.g., DVT, PE)

☐ Member continues to meet all initial authorization criteria

EXCLUSIONS. Therapy will **NOT** be approved if member has history of any of the following:

- Unresolved meningococcal disease
- Any systemic bacterial or significant infections that have not been treated with appropriate antibiotics

Medication being provided by: Please check applicable box below.		
□ Location/site of drug administration:		
NPI or DEA # of administering location:		
<u>OR</u>		

□ Specialty Pharmacy

For urgent reviews: Practitioner should call Sentara Health Plans Pre-Authorization Department if they believe a standard review would subject the member to adverse health consequences. Sentara Health Plan's definition of urgent is a lack of treatment that could seriously jeopardize the life or health of the member or the member's ability to regain maximum function.

**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. *