## 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 can be delayed.</u>

**Drug Requested:** Fabhalta<sup>®</sup> (iptacopan)

MEMBER & PRESCRIBER INF	<b>FORMATION:</b> Authorization 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: Authoriz	zation may be delayed if incomplete.
Drug Name/Form/Strength:	
Dosing Schedule:	Length of Therapy:
Diagnosis:	ICD Code, if applicable:
Weight (if applicable):	Date weight obtained:

## **Recommended Dosage:**

- Paroxysmal nocturnal hemoglobinuria: 200 mg orally twice daily
   Conversion from C5 inhibitors:
  - o Conversion from Soliris® (eculizumab): When converting from eculizumab to iptacopan, initiate iptacopan no later than 1 week following the last eculizumab dose.
  - o Conversion from Ultomiris® (ravulizumab): When converting from ravulizumab to iptacopan, initiate iptacopan no later than 6 weeks following the last ravulizumab dose.
- Primary immunoglobulin A nephropathy: 200 mg orally twice daily

**Quantity Limit:** 2 capsules per day (for **BOTH** indications)

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**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.

D	iag	nos	sis:	Paroxysmal Nocturnal Hemoglobinuria (PNH)
iti	al A	Aut	hoı	rization: 6 months
	Me	dic	atio	n must be prescribed by or in consultation with a hematologist or nephrologist
	Pre	escri	iber	must be enrolled in the Fabhalta® Risk Evaluation and Mitigation Strategy (REMS) program
	Me	emb	er n	nust be 18 years of age or older
	Me	emb	er n	nust meet <b>ONE</b> of the following:
		Fal	bhal	ta® will be used as switch therapy AND member meets ALL the following:
				ember failed Soliris® or Ultomiris® and must meet renewal criteria
			Me	ember does NOT have a systemic infection
			Ne ini	ember must be vaccinated against encapsulated bacteria (Streptococcus pneumoniae, isseria meningitidis, and Haemophilus influenzae type B) at least two weeks prior to tiation of Fabhalta <sup>®</sup> therapy and revaccinated according to current medical guidelines for ecine use
				bhalta <sup>®</sup> will <u>NOT</u> be used in combination with other complement inhibitor therapies (e.g., npaveli <sup>®</sup> , Soliris <sup>®</sup> , Ultomiris <sup>®</sup> or Voydeya <sup>™</sup> )
				<u>OR</u>
		Me	emb	er is treatment-naive AND member meets ALL the following:
				ember must have a diagnosis of Paroxysmal Nocturnal Hemoglobinuria (PNH) confirmed by tection of PNH clones of at least 10% by flow cytometry testing (must submit labs)
			gly	ow cytometry pathology report must demonstrate at least two (2) different vcosylphosphatidylinositol (GPI) protein deficiencies (e.g., CD55, CD59, etc.) within two (2) ferent cell lines from granulocytes, monocytes, erythrocytes (must submit labs)
			exp	ember has laboratory evidence of significant hemolysis (i.e. LDH $\geq$ 1.5 x ULN) <u>AND</u> has perienced <u>ONE</u> of the following additional indications for therapy (must submit chart notes d labs):
				Member is transfusion dependent (defined by having a transfusion within the last 12 months) and has 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
		Me	emb	er does NOT have a systemic infection
				er must be administered a meningococcal vaccine <b>at least two weeks prior</b> to initiation of ta <sup>®</sup> therapy and revaccinated according to current medical guidelines for vaccine use

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	Fabhalta <sup>®</sup> will <u>NOT</u> be prescribed concurrently with another FDA approved product prescribed for treatment of PNH (e.g., Bkemv <sup>™</sup> , Epysqli <sup>™</sup> , PiaSky <sup>®</sup> , Ultomiris <sup>®</sup> , Soliris <sup>®</sup> or Empaveli <sup>®</sup> )
□ D	Diagnosis: Paroxysmal Nocturnal Hemoglobinuria (PNH)
Rea	uthorization: 12 months
	Provider attests to an absence of unacceptable toxicity from the drug (e.g., serious meningococcal infections [septicemia and/or meningitis])
	Member has experienced positive disease response indicated by at least <u>ONE</u> of the following (check all that apply; results must be submitted to document improvement):
	□ Decrease in serum LDH
	☐ Stabilization/increase in hemoglobin level
	☐ Decrease in packed RBC transfusion requirement
	□ Reduction in thromboembolic events
supp	<b>NICAL CRITERIA:</b> Check below all that apply. All criteria must be met for approval. To ort each line checked, all documentation, including lab results, diagnostics, and/or chart notes, must be ided or request may be denied.
□ P	Primary Immunoglobulin A Nephropathy (IgAN)
Initi	ial Authorization: 6 months
	Member is 18 years of age or older
	Provider is a nephrologist
	Member has a diagnosis of biopsy-proven, primary immunoglobulin A nephropathy (IgAN) and is at risk of rapid disease progression
	Member has been on a stable, maximized dose of a renin-angiotensin system (RAS) inhibitor (≥ 50% of maximum labeled dose), including either an angiotensin-converting enzyme (ACE) inhibitor or angiotensin receptor blocker (ARB), for at least 90 days (verified chart notes and/or pharmacy paid claims)
	Members' lab test results taken within the last 30 days must be submitted to document <u>ALL</u> the following:
	□ Total urine protein $\ge 1$ g/day
	☐ Urine protein-to-creatinine ratio is $\geq 1.5 \text{ g/g}$
	$\Box  eGFR \ge 30 \text{ mL/min/1.73 m}^2$
	Member will avoid concomitant therapy with major interacting drugs, including <u>ALL</u> the following:
	Strong CYP2C8 inhibitors (e.g., gemfibrozil)
	• CYP2C8 inducers (e.g., rifampin)

	Mem	ber must meet ONE of the following:
		lember has had an unsuccessful 3-month trial of oral generic budesonide EC capsules (must submit nart notes or lab test results confirming therapy failure)
	la	Tember has an intolerance or hypersensitivity to oral generic budesonide EC capsules, or an FDA beled contraindication to oral generic budesonide EC capsules that is not expected to occur with the quested agent (documentation of intolerance or hypersensitivity must be submitted)
		ber has had unsuccessful 3-month trials of Filspari <sup>®</sup> <u>AND</u> Tarpeyo <sup>®</sup> (must submit chart notes or est results confirming therapy failure)
		ber is <u>NOT</u> using concomitant therapy with any of the following: Tarpeyo <sup>®</sup> , Filspari <sup>®</sup> , Fabhalta <sup>®</sup> or complement inhibitor therapies (e.g., Empaveli <sup>®</sup> , Soliris <sup>®</sup> , Ultomiris <sup>®</sup> or Voydeya <sup>™</sup> )
	Diag	nosis: Primary Immunoglobulin A Nephropathy (IgAN)
		nosis: Primary Immunoglobulin A Nephropathy (IgAN)  orization: 12 months
	autho	
	Mem'stabil	orization: 12 months
Re	Memi Memi stabil	per continues to meet all initial authorization criteria  ther must have reduction in proteinuria from baseline after initial approval, and reduction or dization in proteinuria after subsequent approvals (current lab test results must be submitted for mentation)  there has NOT experienced any treatment-restricting adverse effects (e.g., serious and life-threatening)

 $\label{eq:medication} \textbf{Medication being provided by Specialty Pharmacy} - \textbf{Proprium Rx}$ 

\*\*Use of samples to initiate therapy does not meet step edit/ preauthorization criteria.\*\*

\*Previous therapies will be verified through pha rmacy paid claims or submitted chart notes.\*