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</u>, <u>correct</u>, <u>or legible</u>, <u>authorization may be delayed</u>.

Prophylaxis Hereditary Angioedema (HAE) (Pharmacy)

<u>Drug Requested</u> (select applicable drug below):

	Cinryze® (C1 Esterase Inhibitor Human)		Haegarda® (C1 Esterase Inhibitor Human)			
	Orladeyo® (berotralstat)		Takhzyro® (lanadelumab)			
3.6		ONT.				
NI.	EMBER & PRESCRIBER INFORMATION	UN:	Authorization may be delayed if incomplete.			
Mer	mber Name:					
	mber Sentara #:					
Pres	scriber Name:					
	scriber Signature:					
Offi	ice Contact Name:					
Pho	ne Number:		Fax Number:			
DE A	A OR NPI #:					
	RUG INFORMATION: Authorization may be					
	g Form/Strength:					
			Length of Therapy:			
Diagnosis:			ICD Code:			
	ight:		Oate:			
	sing Limit: (see below)					
	Haegarda: Administer 60 International Units (IU) p	er k	g body weight twice weekly (every 3 or 4 days)			
	☐ Quantity Limit:					
	☐ Haegarda 2000 IU SDV kit: 16 kits per 28 da	ays				
	☐ Haegarda 3000 IU SDV kit: 8 kits per 28 day	-				
Cinryze: Administer 1,000 units intravenous every 3 or 4 days						
	☐ Quantity Limit:		- -			
	☐ Cinryze 500-unit vial: 16 vials per 30 day	ys				

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PA Prophylaxis Hereditary Angiodema (Pharmacy) (CORE)

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 □ Orladeyo: Adults and children ≥ 12 years of age - One capsule taken orally once daily with □ Quantity Limit: 						
	_	□ 150 mg capsules: 1 capsule per day				
		110 mg capsules: 1 capsule per day (For members with Moderate to severe impairment (Child-Pugh class B and C)				
		Quantity Limit:				
		□ 300 mg vial/syringe per 14 days				
		□ 150 mg syringe per 28 days				
ach	line	CAL CRITERIA: Check below all that apply. All criteria must be met for approval. To support checked, all documentation, including lab results, diagnostics, and/or chart notes, must be provided on any be denied.				
niti	al A	Authorization: 12 months				
		ent of acute attacks of Hereditary Angioedema Criteria:				
		escribed by or in consultation with a specialist in allergy, immunology, hematology, pulmonology or edical genetics				
		AND				
		ovider attests the requested medication will <u>NOT</u> be used in combination with other prophylactic erapies targeting C1 inhibitors or kallikrein (e.g., Haegarda or Takhzyro)				
		<u>AND</u>				
	Pro	ovider attests the patient is avoiding BOTH of the following possible triggers for HAE attacks:				
		Estrogen-containing oral contraceptive agents AND hormone replacement therapy				
		Antihypertensive agents containing ACE inhibitors				
		AND				
	Me	ember must meet medication specific age requirement:				
		Haegarda: Member must be at least 6 years of age				
		Cinryze: Member must be at least 6 years of age				
		Takhzyro: Member must be at least 2 years of age				
		Orladeyo: Member must be at least 12 years of age				
		AND				
	Me	ember has a history of ONE of the following:				
		Three (3) or more severe HAE attacks per month (i.e., airway swelling, debilitating cutaneous or				

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gastrointestinal episodes)

or

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- □ Disablement for more than 5 days per month by HAE
- □ Recurrent laryngeal attacks caused by HAE

AND

□ Treatment of member with "on-demand" therapy (i.e., Kalbitor®, Firazyr®, Ruconest®, or Berinert®) did NOT provide satisfactory control or access to "on-demand therapy" is limited (failure is defined as more than 5 attacks/month for 4 months consecutively within the same year)

AND

☐ Member failed, is intolerant, or has a contraindication to attenuated (17 alpha-alkylated) androgens (e.g., Danazol®) for HAE prophylaxis

AND

- □ Orladeyo (berotralstat) Criteria Only:
 - ☐ Member has tried and failed the preferred prophylaxis HAE medication Takhzyro[®] (lanadelumab)

Member has ONE of the following clinical presentations that is consistent with a HAE subtype, confirmed by repeat blood testing (please submit chart notes for symptoms and lab values to confirm the HAE subtype):

II.A. HAE I: (all bullet points must apply)

- □ Low C1 inhibitor (C1-INH) antigenic level (C1-INH antigenic level below the lower limit of normal as defined by the laboratory performing the test)
- □ Low C4 level (C4 below the lower limit of normal as defined by the laboratory performing the test)
- □ Low C1-INH functional level (C1-INH functional level below the lower limit of normal as defined by the laboratory performing the test) **in addition to ONE** of the following:
 - ☐ Member has a family history of HAE
 - □ Acquired angioedema has been ruled out (i.e., patient onset of symptoms occur prior to 30 years old, normal C1q levels, patient does not have underlying disease such as lymphoma or benign monoclonal gammopathy [MGUS])

OR

II.B. HAE II (C1-Inhibitor dysfunction): (all bullet points must apply)

- □ Normal to elevated C1-INH antigenic level
- □ Low C4 level (C4 below the lower limit of normal as defined by the laboratory performing the test)
- □ Low C1-INH functional level (C1-INH functional level below the lower limit of normal as defined by the laboratory performing the test) **in addition to ONE** of the following:
 - ☐ Member has a family history of HAE
 - □ Acquired angioedema has been ruled out (i.e., onset of symptoms occurred prior to 30 years old, normal C1q levels, patient does not have underlying disease such as lymphoma or benign monoclonal gammopathy [MGUS])

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No- approve dosing every 2 weeks

		<u>OR</u>
II.C. 🗆	H	AE III with normal C1-INH: (all bullet points must apply)
		Normal C1-INH antigenic level
		Normal C4 level
		Normal C1-INH functional level
		Repeat blood testing during an attack has confirmed the member does NOT have abnormal lab values indicative of HAE I or HAE II
		Member had an inadequate response or intolerance to an adequate trial of prophylactic therapy with ONE of following:
		□ antifibrinolytic agent: (□ tranexamic acid (TXA) OR □ aminocaproic acid)
		□ 17α- alkylated androgen: danazol
		□ progestins (female members only)
		AND
		ONE of the following:
		 Member has a known HAE-causing mutation (e.g., mutation of coagulation factor XII gene [F12 mutation], mutation in the angiopoietin-1 gene, mutation in the plasminogen gene or kininogen-1) Member has a family history of HAE and documented evidence of lack of efficacy of chronic high-dose antihistamine therapy (e.g., cetirizine standard dosing at up to four times daily or an alternative equivalent, given for at least one month or an interval long enough to expect three or more angioedema attacks) AND corticosteroids
check	æd,	orization: Check below all that apply. All criteria must be met for approval. To support each line all documentation, including lab results, diagnostics, and/or chart notes, must be provided or request enied.
	Me	ember must continue to meet initial criteria
		<u>AND</u>
	Sig	gnificant improvement in severity and duration of attacks has been achieved and sustained
		<u>AND</u>
		ember has experienced an absence of unacceptable toxicity from the drug (e.g., hypersensitivity actions)
		<u>AND</u>
	Fo	r Takhzyro Renewal Only: Has the member been attack free for greater than > 6 months?
		Yes- approve dosing every 4 weeks

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Medication	being provid	led by a S	pecialty	Pharmac	v - Pro	priumRx

**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. *