SENTARA HEALTH PLAN

PHARMACY PRIOR AUTHORIZATION/STEP-EDIT REQUEST

<u>Directions</u>: The prescribing physician must sign and clearly print name (preprinted stamps not valid) on this <u>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 can be delayed.</u>

Drug Requested: Icatibant or Sajazir (Firazyr®) (Pharmacy)

MEMBER & PRESCRIBER INFORMATION: Authorization may be delayed if incomplete.				
Member Name:				
	Date of Birth:			
Prescriber Name:				
	Date:			
Office Contact Name:				
	Fax Number:			
DEA OR NPI #:				
DRUG INFORMATION: Authorization r				
Dosing Schedule:	Length of Therapy:			
Diagnosis:				
Dosing Limit : (see below)				
A. Quantity Limit (max daily dose): Pharmacy Benefit: icatibant or sajazir	(Firazyr) 30mg/3ml vial: 3 subcutaneous pen per 28 days			
B. Max Units (per dose and over time): Medical Benefit: 90 billable units per 2	8 days; 1mg = 1billable			
• J1744 30mg/3mL vial: 1mg=1billable AND	NDC 54092-0702-xx 30mg			
• Coverage is provided for 12 months	and will be eligible for renewal			

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.

Standard Review. In checking this box, the timeframe does not jeopardize the life or health of the member or the member's ability to regain maximum function and would not subject the member to severe pain.

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Initial Approval Criteria - 12 months: The cumulative amount of medication(s) the patient has on-hand, indicated for the acute treatment of HAE, will be taken into account when authorizing. The authorization will provide a sufficient quantity in order for the patient to have a cumulative amount of HAE medication(s) on-hand in order to treat acute attacks for the duration of the authorization (unless otherwise specified).

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	Member must be at least 18 years of age
	<u>AND</u>
	Must be prescribed by or in consultation with a specialist in: allergy, immunology, hematology, pulmonology or medical genetics
	AND
	Provider attests the patient is avoiding <u>ALL</u> of the following possible triggers for HAE attacks: Helicobacter pylori infections (confirmed by lab test)
	 Estrogen-containing oral contraceptive agents AND hormone replacement therapy Antihypertensive agents containing ACE inhibitors
	AND
	Member has a history of one of the following criteria:
	☐ Three (3) or more severe HAE attacks per month (i.e., airway swelling, debilitating cutaneous or gastrointestinal episodes)
	☐ Disablement for more than 5 days per month by HAE
	☐ Recurrent laryngeal attacks caused by HAE
	AND
su	tient has one of the following clinical presentations that is consistent with a HAE btype, confirmed by repeat blood testing (please submit chart notes for symptoms and b 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)

<u>OR</u>

☐ Member has a family history of HAE

benign monoclonal gammopathy [MGUS])

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by the laboratory performing the test) **AND** one of the following:

□ 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

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

		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) AND 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])				
		<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 patient does not have abnormal lab value indicative of HAE I or HAE II				
		Patient had an inadequate response or intolerance to an adequate trial of prophylactic therapy we one of following:				
		□ antifibrinolytic agent: (□ tranexamic acid (TXA) OR □ aminocaproic acid)				
		\Box 17 α - alkylated androgen: danazol				
		□ progestins (female patients only)				
		AND				
		One of the following:				
		□ Patient 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)				
		Patient 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				
		Criteria. All criteria must be checked for approval. To support each line checked, all on (lab results, diagnostics, and/or chart notes) must be provided or request may be denied.				

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

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☐ Member must continue to meet initial criteria

AND

	Significant improvement in severity and duration of attacks have been achieved and sustained						
	-	AND absence of unacceptable toxicity from the drug (e.g. hypersensitivity reactions)					
Medication being provided by (check box below that applies):							
C	☐ Physician's office	OR		Specialty Pharmacy- PropriumRx			
s	standard review would subj	ect the member ld seriously jeop	to adve	na Pre-Authorization Department if they believe a erse health consequences. Optima's definition of urgent the life or health of the member or the member's ability			
	Use of samples to in	itiate therapy	does n	not meet step edit/preauthorization criteria.			

Trevous merupies was severyed among a prairie of para causes of submitted chart notes.