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 (<u>including phone and fax #s</u>) on this form is correct. <u>If information provided is not complete</u>, correct, or legible, authorization can be delayed.

MEMBER & PRESCRIBER INFORMATION: Authorization may be delayed if incomplete.

<u>Drug Requested</u>: Tezspire[™] (tezepelumab) (J2356) (Medical)

provided or request may be denied.

Initial Authorization: 6 months

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Member Name:					
Member Sentara #:	Date of Birth:				
Prescriber Name:					
Prescriber Signature:					
Office Contact Name:					
Phone Number: Fax Number:					
DEA OR NPI #:					
DRUG INFORMATION: Authoriz					
Drug Form/Strength:					
Dosing Schedule:	Length of Therapy:				
Diagnosis:	ICD Code, if applicable:				
Weight:	Date:				
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.					
	adolescents ≥ 12 years: 210 mg/1.9 mL SubQ, single-dose prefilled teks. Tezspire [™] 210mg/1.9ml; 1 syringe or vial=210 billable units.				
Fasenra [®] , Nucala [®] , and Xolair [®] to of these combinations have NOT b	ne use of concomitant therapy with Cinqair®, Dupixent [®] , be experimental and investigational. Safety and efficacy een established and will <u>NOT</u> be permitted. In the event Dupixent [®] , Fasenra [®] , Nucala [®] , and Xolair [®] authorization Tezspire [™] will <u>NOT</u> be approved.				
	ow all that apply. All criteria must be met for approval. To support luding lab results, diagnostics, and/or chart notes, must be				

1.	Has the member been approved for Tezspire TM previously through the Sentara medical department?					
	□ Yes □ No					
2.	Is the member 12 years of age or older?					
	□ Yes □ No					
3.	Does the member have a diagnosis of severe asthma*?					
	□ Yes □ No					
4.	Will coadministration with another monoclonal antibody be avoided (i.e. omalizumab, mepolizumab, reslizumab, benralizumab, and dupilumab)?					
	□ Yes □ No					
5.	Will Tezspire [™] be used for add-on maintenance treatment in members regularly receiving both (unless otherwise contraindicated) of the following:					
	 Medium to high dose inhaled corticosteroids AND 					
	• An additional controller medication (ie. long-acting beta agonist, leukotriene modifiers?					
	□ Yes □ No					
6.	Has the member had two or more exacerbations in the previous year requiring oral or injectable corticosteroid treatment (in addition to the regular maintenance therapy defined above) or one exacerbation resulting in hospitalization?					
	□ Yes □ No					
7.	Does the member have at least one of the following for assessment of clinical status:					
	Use of systemic corticosteroids					
	Use of inhaled corticosteroids					
	• Number of hospitalizations, ER visits, or unscheduled visits to healthcare provider due to condition					
	• Forced expiratory volume in 1 second (FEV ₁)?					
	□ Yes □ No					
8.	Has the member tried and failed an adequate trial of the 2 different preferred products (Fasenra® and Xolair®)?					
	☐ Yes ☐ No ☐ N/A (continued below)					
	If N/A selected for question 8, please answer the following:					
	1. Does the member lack an eosinophilic phenotype with blood eosinophils ≥ 150 cells/ μ L? AND					
	□ Yes □ No					
	2. Does the member lack a serum IgE level > 30 IU/mL? OR					
	□ Yes □ No					
	3. Does the member have another predicted intolerance to the preferred agents?					
	□ Yes □ No					
	Please provide explanation:					

(Continued on next page)

Reauthorization: 12 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.

1.	Has the	member	been	assessed	for	toxicity?	
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□ Yes □ No

- 2. Does the member have improvement in asthma symptoms or asthma exacerbations as evidenced by a decrease in one or more of the following:
 - Use of systemic corticosteroids
 - Hospitalizations
 - ER visits
 - Unscheduled visits to healthcare provider
 - Improvement from baseline in forced expiratory volume in 1 second (FEV₁)?

□ No

*Components of severity for classifying asthma as severe may include any of the following (not all inclusive):

- Symptoms throughout the day
- Nighttime awakenings, often 7 times per week
- SABA use for symptom control occurs several times per day
- Extremely limited normal activities
- Lung function (percent predicted FEV₁) < 60%
- Exacerbations requiring oral systemic corticosteroids are generally more frequent and intense relative to moderate asthma.

Medication being provided by a Specialty Pharmacy - PropriumRx

For urgent reviews: Practitioner should call Sentara Health Pre-Authorization Department if they believe a standard review would subject the member to adverse health consequences. Sentara Health'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. *