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

Drug Requested: Cinqair® IV (reslizumab) (J2786) (Medical)

MEMBER & PRESCRIBER IN	FORMATION: Authorization may be delayed if incomplete.
Member Name:	
Member Sentara #:	
Prescriber Name:	
	Date:
Office Contact Name:	
Phone Number:	Fax Number:
DEA OR NPI #:	
DRUG INFORMATION: Author	
Drug Form/Strength:	
Dosing Schedule:	Length of Therapy:
Diagnosis:	ICD Code, if applicable:
Weight:	
	x, the timeframe does not jeopardize the life or health of the member or turn function and would not subject the member to severe pain.
Recommended Dosage: Dosage 3m minutes	ng/kg once every 4 weeks by intravenous infusion over 20 - 50

*Sentara considers the use of concomitant therapy with Cinqair®, Dupixent®, Fasenra®, Nucala®, Tezspire™ and Xolair® to be experimental and investigational. Safety and efficacy of these combinations have NOT been established and will NOT be permitted. In the event a member has an active Dupixent®, Fasenra®, Nucala®, Tezspire™ or Xolair® authorization on file, all subsequent requests for Cinqair® will NOT be approved.

(Continued on next page)

each provi	line checked, all documentation, including lab results, diagnostics, and/or chart notes, must be ded or request may be denied. (Trials will be verified using pharmacy claims and/or submitted inotes.)
	al Authorization: 12 months
	Prescribed by or in consultation with an allergist, immunologist or pulmonologist
	Member is 18 years of age or older
	Member has been diagnosed with severe eosinophilic phenotype defined by a baseline (pre-Cinqair®) peripheral blood eosinophil level of ≥ 400 cells/microliter
	Member is currently being treated with <u>ONE</u> of the following unless there is a contraindication or intolerance to these medications and must be compliant on therapy <u>for at least 90 consecutive days</u> within a year of request:
	High-dose inhaled corticosteroid (ICS) (e.g., greater than 500 mcg fluticasone propionate equivalent/day) AND an additional asthma controller medication (e.g., leukotriene receptor antagonist, long-acting beta-2 agonist (LABA), theophylline)
	One maximally dosed combination ICS/LABA product (e.g., Advair® (fluticasone propionate/salmeterol), Dulera® (mometasone/formoterol), Symbicort® (budesonide/formoterol))
	Member has experienced ONE of the following (check box that applies):
	☐ More than 2 exacerbations requiring additional medical treatment (e.g., an increase in oral corticosteroid dose, emergency department, urgent care visits or hospitalizations) within the past 12 months
	☐ Any prior intubation for an asthma exacerbation
	Member has a baseline forced expiratory volume (FEV1) \leq 80% predicted normal submitted within year of request
	Provider must submit member blood eosinophil count collected after a trial and failure of at least 90 consecutive days of therapy with high dose inhaled corticosteroids <u>AND</u> long-acting inhaled beta-2 agonist. A failure of these medications is defined as a blood count > 400 cells/microliter (submit labs collected within the past 12 months)
	Eosinophil count: Date:

(Continued on next page)

CLINICAL CRITERIA: Check below all that apply. All criteria must be met for approval. To support

2

at

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. (Trials will be verified using pharmacy claims and/or submitted chart
notes.)

		ember has experienced a sustained positive clinical response to Cinqair® therapy as demonstrated by ast ONE of the following (check all that apply; chart notes must be submitted) :	
		Increase in percent predicted Forced Expiratory Volume (FEV1) from baseline (pre-treatment)	
		Reduction in the dose of inhaled corticosteroids required to control asthma	
		Reduction in the use of oral corticosteroids to treat/prevent exacerbation	
		Reduction in asthma symptoms such as chest tightness, coughing, shortness of breath or nocturnal awakenings	
		ember is currently being treated with ONE of the following unless there is a ntraindication or intolerance to these medications:	
		High-dose inhaled corticosteroid (ICS) (e.g., greater than 500 mcg fluticasone propionate equivalent/day) <u>AND</u> an additional asthma controller medication (e.g., leukotriene receptor antagonist, long-acting beta-2 agonist (LABA), theophylline)	
		One maximally dosed combination ICS/LABA product (e.g., Advair® (fluticasone propionate/salmeterol), Dulera® (mometasone/formoterol), Symbicort® (budesonide/formoterol))	
Iedication being provided by (check applicable box(es) below):			
	Lo	ocation/site of drug administration:	
	NF	PI or DEA # of administering location:	
		OR	
	Sp	ecialty Pharmacy - PropriumRx	

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