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

Drug Requested: Nucala® SQ (mepolizumab) (Pharmacy)

{Eosinophilic Granulomatosis Polyangiitis (EGPA)}

MEMBER & PRESCRIBER INFO	DRMATION: Authorization may be delayed if incomplete.
Member Name:	
Member Sentara #:	Date of Birth:
Prescriber Name:	_
Prescriber Signature:	
Office Contact Name:	
Phone Number:	Fax Number:
DEA OR NPI #:	
DRUG INFORMATION: Authorizat	tion may be delayed if incomplete.
Drug Form/Strength:	
	Length of Therapy:
Diagnosis:	ICD Code, if applicable:
Weight:	Date:
Recommended Dosage: 300 mg/mL SubQ single-dose prefilled auto-injector/single-dos	once every 4 weeks administered as 3 separate 100-mg injections; e prefilled syringe
Nucala®, Tezspire® and Xolair® to be expecombinations have NOT been established	comitant therapy with Cinqair®, Dupixent®, Fasenra®, erimental and investigational. Safety and efficacy of these and will NOT be permitted. In the event a member has an expire® or Xolair® authorization on file, all subsequent requests
	v all that apply. All criteria must be met for approval. To a, including lab results, diagnostics, and/or chart notes, must be
Initial Authorization: 12 months	
☐ Medication must be prescribed by an	allergist, immunologist, or pulmonologist
☐ Member must be 18 years of age or o	lder

			ember must have diagnosis of Eosinophilic Granulomatosis with Polyangiitis (EGPA) (Churg-Strauss adrome) > 6 months based on the history or presence of asthma						
	La	b do	cur	nentation must show an eosinophil count of ≥150 cells/microliter at baseline					
		All Mo Pu Sir Ma Glo All Pa	hber must have documentation of <u>TWO</u> of the following: A biopsy showing evidence of EGPA Mono-or polyneuropathy Pulmonary infiltrates, non-fixed on chest x-rays Sino-nasal abnormality Magnetic Resonance Imaging or Echocardiography of cardiomyopathy Glomerulonephritis Alveolar hemorrhage (by bronchoalveloar lavage) Palpable purpura Anti-neutrophil cytoplasmic anti-body (ANCA) positive (Myeloperoxidase or proteinase 3)						
□ Relapsing disease:									
			Mo	ember must have a history of at least <u>ONE</u> confirmed EGPA relapse requiring: An increase in oral corticosteroids (OCS) dose Initiation or increased dose of immunosuppressive therapy (e.g., azathioprine, cyclophosphamide, methotrexate, or mycophenolate mofetil) Hospitalization ast have occurred > 12 weeks but < 2 years prior to initiation while receiving a dose of ednisone (or equivalent) of >7.5 milligram per day (mg/day) for <u>at least 90 consecutive days</u>					
		Re	efra	actory disease:					
			Re	Failure to attain remission (Birmingham Vasculitis Activity Score (BVAS) =0) and OCS dose <7.5 mg/day prednisone or equivalent) for at least 90 consecutive days within the last 6 months following induction treatment with a standard regimen (e.g., azathioprine cyclophosphamide, methotrexate, mycophenolate mofetil, or high-dose corticosteroids (> 15 mg/day prednisone), administered for at least 3 months Within 6 months prior to initiation, recurrence of symptoms of EGPA while tapering oral corticosteroids (OCS), occurring at any dose level ≥7.5 mg/day prednisone or equivalent taken for at least 90 consecutive days					

(Continued on next page)

2

Exclusion Criteria – Therapy will \underline{NOT} be approved if member has history of any of the following:

- Organ/life threatening EGPA within 3 months prior to initiation
- Malignancy: current malignancy or previous history of cancer in remission for < 12 months
- Unstable cardiovascular disease: Ejection fraction < 20%, New York Heart Association Class III/IV failure, acute myocardial infarction diagnosed less than 3 months
- Unstable liver disease: Presence of ascites, encephalopathy, coagulopathy, hypoalbuminemia, esophageal or gastric varices, cirrhosis, and known biliary abnormalities (with the exception of Gilbert's syndrome or asymptomatic gallstones
- Rituximab within the past year; IVIg within the past 6 months; omalizumab within the past 4 months
- Pregnancy, breast-feeding, absence of contraception if female of child-bearing age

Reauthorization: 12 months. All criteria must be checked for approval. To support each line checked, all documentation (lab results, diagnostics, and/or chart notes) must be provided or request may be denied.

Member must meet ONE of the following:
\square Documentation of remission or improvement in the Birmingham Vasculitis Activity Score (BVAS) or prednisone/prednisolone daily dose of \leq 7.5mg
☐ Documentation of decrease in maintenance dose of systemic corticosteroids, improvement in asthmatistic symptoms or asthmatic exacerbations
☐ Documentation of disease flares with tapering of corticosteroid therapy or immunotherapy

Medication	being p	rovided by	y a S	pecialty	y Pharmac	y – Proj	prium	Rx

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.