AvMed

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 1-305-671-0200</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 the information provided is not complete, correct, or legible, the authorization process can be delayed.</u>

Drug Requested: Fasenra® SQ (benralizumab) (Pharmacy)

Quantity Limits: 1 syringe per 56 days (both strengths)

MEMBER & PRESCRIBER IN	FORMATION: Authorization may be delayed if incomplete.	
Member Name:		
Member AvMed #:	Date of Birth:	
Prescriber Name:		
Prescriber Signature:	Date:	
Office Contact Name:		
	Fax Number:	
NPI #:		
DRUG INFORMATION: Authori	zation may be delayed if incomplete.	
Drug Form/Strength:		
	Length of Therapy:	
Diagnosis:	ICD Code, if applicable:	
Weight (if applicable):	Date weight obtained:	
Recommended Dosing:		
□ Asthma, severe eosinophilic:		
 Adult and Adolescent Patients 1 	2 Years of Age and Older:	
o 30 mg every 4 weeks for the first 3 doses followed by once every 8 weeks thereafter		
 Pediatric Patients 6 Years to 11 Y 	ears of Age:	
	 Weighing Less Than 35 kg: the recommended dosage is 10 mg every 4 weeks for the first 3 dose followed by once every 8 weeks thereafter 	
 Weighing 35 kg or More: the refollowed by once every 8 wee 	recommended dosage is 30 mg every 4 weeks for the first 3 doses ks thereafter	
Eosinophilic granulomatosis with polyangiitis (EGPA): 30 mg every 4 weeks		

Med	Medication will be (select <u>ONE</u> of the following):				
Ţ	□ Self-Administered (pharmacy benefit)				
(☐ Administered by Provider (medical benefit)				
*The Health Plan 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 \underline{NOT} been established and will \underline{NOT} be permitted. In the event a member has an active Cinqair®, Dupixent®, Nucala®, Tezspire™ or Xolair® authorization on file, all subsequent requests for Fasenra® will \underline{NOT} be approved.					
• 7	Will the member be discontinuing a previously prescribed biologic if approved for requested medication? ☐ Yes OR ☐ No				
I	ledication to be discontinued: Effective date:				
I	ledication to be initiated: Effective date:				
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.					
	Piagrania. Agabana garana again ambilia				
	Diagnosis: Asthma, severe eosinophilic ial Authorization: 12 months				
Ini	ial Authorization: 12 months Prescribed by or in consultation with an allergist, immunologist or pulmonologist				
Ini	ial Authorization: 12 months Prescribed by or in consultation with an allergist, immunologist or pulmonologist Member is 6 years of age or older				
<u>Ini</u>	ial Authorization: 12 months Prescribed by or in consultation with an allergist, immunologist or pulmonologist Member is 6 years of age or older Has the member been approved for Fasenra® previously through the AvMed medical department?				
<u>Ini</u>	Prescribed by or in consultation with an allergist, immunologist or pulmonologist Member is 6 years of age or older Has the member been approved for Fasenra® previously through the AvMed medical department? Yes □ No Member has been diagnosed with severe eosinophilic phenotype defined by a baseline (pre-Fasenra®) peripheral blood eosinophil level ≥ 150 cells/microliter at the initiation of treatment Member is currently being treated with ONE of the following unless there is a contraindication or intolerance to these medications and must be compliant on therapy for at least 90 consecutive days within a year of request:				
<u>Ini</u>	Prescribed by or in consultation with an allergist, immunologist or pulmonologist Member is 6 years of age or older Has the member been approved for Fasenra® previously through the AvMed medical department? ☐ Yes ☐ No Member has been diagnosed with severe eosinophilic phenotype defined by a baseline (pre-Fasenra®) peripheral blood eosinophil level ≥ 150 cells/microliter at the initiation of treatment Member is currently being treated with ONE of the following unless there is a contraindication or intolerance to these medications and must be compliant on therapy for at least 90 consecutive days				

	☐ More than > 2 exacerbations required corticosteroid dose, emergency of	the following (check box that applies): uiring additional medical treatment (e.g., an increase in oral department, urgent care visits or hospitalizations) within the past		
	12 months ☐ Any prior intubation for an asthr	na evacerbation		
	• •	atory volume (FEV1) < 80% predicted normal (< 90% for		
	Provider must submit member blood eosinophil count 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 > 150 cells/microliter (submit labs collected within the past 12 months)			
	Eosinophil count:	Date:		
□ I	Diagnosis: Asthma, severe eosi	nophilic		
supp		ek below all that apply. All criteria must be met for approval. To on, including lab results, diagnostics, and/or chart notes, must be		
	<u> </u>	l positive clinical response to Fasenra® therapy as demonstrated by k all that apply; chart notes must be submitted) :		
	☐ Increase in percent predicted Fo	rced Expiratory Volume (FEV1) from baseline (pre-treatment)		
	☐ Reduction in the dose of inhaled	corticosteroids required to control asthma		
	☐ Reduction in the use of oral cort	icosteroids to treat/prevent exacerbation		
	☐ Reduction in asthma symptoms awakenings	such as chest tightness, coughing, shortness of breath or nocturnal		
	Member is currently being treated wintolerance to these medications:	rith <u>ONE</u> of the following unless there is a contraindication or		
		l (ICS) (e.g., greater than 500 mcg fluticasone propionate nal asthma controller medication (e.g., leukotriene receptor conist (LABA), theophylline)		
	-	on ICS/LABA product (e.g., Advair® (fluticasone		
	propionate/salmeterol), Dulera®	(mometasone/formoterol), Symbicort® (budesonide/formoterol))		
□ Diagnosis: Eosinophilic Granulomatosis Polyangiitis (EGPA)				
<u>Ini</u>	tial Authorization: 12 months			
	Prescribed by or in consultation with	n an allergist, immunologist, pulmonologist, or rheumatologist		
	Member is 18 years of age or older			
	5 7 11 11 55 51 51 46 1			

Has the member been approved for Fasenra [®] previously through the Health Plan medical department? ☐ Yes ☐ No			
Member must have a diagnosis of Eosinophilic Granulomatosis with Polyangiitis (EGPA) (Churg-Strauss Syndrome) based on the history or presence of asthma			
Member must have a blood eosinophil level $> 10\%$ of total white blood cells or an absolute eosinophil count > 1000 cells/mm ³ at baseline			
Eosinophil count: Date:			
Member must have documentation of <u>TWO</u> of the following:			
☐ A biopsy showing histopathologic evidence of eosinophilic vasculitis, perivascular eosinophilic infiltration, or eosinophil rich granulomatous inflammation			
□ Neuropath; 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 or (Myeloperoxidase or proteinase 3)			
Member has active, non-severe disease defined as vasculitis without life-or organ-threatening manifestations. Examples of symptoms in patients with non-severe disease include rhinosinusitis, asthma, mild systemic symptoms, uncomplicated cutaneous disease and mild inflammatory arthritis.			
Member must have a history of ONE of the following:			
□ <u>Relapsing disease</u> :			
☐ Member must have a history of at least ONE 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, rituximab)			
☐ Hospitalization			
☐ Must have occurred within the past 2 years while receiving a dose of prednisone (or equivalent of > 7.5 milligram per day (mg/day) for at least 90 consecutive days	t)		

Refractory	disease:

- □ Refractory disease must meet <u>ONE</u> of the following:
 - □ 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 a standard regimen (e.g., azathioprine cyclophosphamide, methotrexate, mycophenolate mofetil, high-dose corticosteroids or rituximab administered for at least 3 months
 - □ Within the past 6 months, the member has had a recurrence of EGPA symptoms during the tapering of oral corticosteroids (OCS), at any dose level of ≥ 7.5 mg/day of prednisone or equivalent, taken for <u>at least 90 consecutive days</u>
- ☐ Member has been on a stable dose of oral corticosteroid therapy for at least 4 weeks prior to starting treatment (e.g., prednisone or equivalent of ≥ 7.5 mg/day)

□ Diagnosis: Eosinophilic Granulomatosis Polyangiitis (EGPA)

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.

- \square Member must meet **ONE** of the following:
 - □ Documentation of remission or improvement in the Birmingham Vasculitis Activity Score (BVAS)=0 (no active vasculitis) plus prednisone/prednisolone daily dose of ≤ 7.5 mg/day or equivalent
 - □ Documentation of improvement in duration of remission or decrease frequency in the occurrence of relapses
 - Documentation of decrease in maintenance dose of systemic corticosteroids
 - □ Documentation of improvement on a disease activity scoring tool [e.g., Vasculitis Damage Index (VDI), Birmingham Vasculitis Activity Score (BVAS), Forced vital capacity (FVC), Forced Expiratory Volume during first second (FEV1), Asthma Control Questionnaire (6-item version) ACQ-6), etc.]

Medication being provided by a Specialty Pharmacy - Proprium 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.