



Request for Prior Authorization IL-5 ANTAGONISTS

1.877.386.4695 **Provider Help Desk** 1.866.399.0928

FAX Completed Form To

(PLEASE PRINT - ACCURACY IS IMPORTANT)

IA Me	dicaid	Me	mbe	r ID	#			Pa	atient name				DC	В					
Patien	t add	lress																	
Provider NPI							Prescriber name			Phone									
Prescriber address							Fax												
Pharmacy name Ad								Ad	ddress			Phone							
Prescriber must complete all information above. It must be legible, correct, and complete or form will be returned.																			
Pharmacy NPI									Pharmacy fax	NE	C 								

Prior authorization is required for IL-5 antagonists. Requests will not be considered with concurrent use with another monoclonal antibody. Payment will be considered under the following conditions:

- 1) Patient meets the FDA approved age for submitted diagnosis; and
- 2) Is dosed within FDA approved dosing for submitted diagnosis and age; and
- 3) Patient has a diagnosis of severe asthma with an eosinophilic phenotype; and
 - a) Patient has a pretreatment blood eosinophil count of ≥150 cells per mcL within the previous 6 weeks or blood eosinophils of ≥300 cells per mcL within 12 months prior to initiation of therapy; and
 - b) Symptoms are inadequately controlled with documentation of current treatment with a high-dose inhaled corticosteroid (ICS) given in combination with a controller medication (long-acting beta2-agonist [LABA] and leukotriene receptor antagonist [LTRA]) for a minimum of 3 consecutive months, with or without oral corticosteroids. Patient must be compliant with therapy, based on pharmacy claims; and
 - c) Patient has a history of two (2) or more exacerbations in the previous year despite regular use of high-dose ICS plus a LABA and LTRA; and
 - d) A pretreatment forced expiratory volume in 1 second (FEV₁) <80% predicted in adults and < 90% in adolescents; or
- 4) Patient has a diagnosis of eosinophilic granulomatosis with polyangiitis; and
 - a) Patient has documentation of an adequate trial and therapy failure with systemic glucocorticoids; and
 - b) One of the following:
 - i. Eosinophil count greater than 1000 cells/mcL: or
 - ii. Eosinophil count greater than 10% of the total leukocyte count; and
- 5) Prescribed by or in consultation with an allergist, immunologist, pulmonologist, or rheumatologist.

If the criteria for coverage are met, an initial authorization will be given for 3 months to assess the need for continued therapy. Requests for continuation of therapy will be based on continued medical necessity and will be considered when the following criteria are met:

Severe Asthma with an Eosinophilic Phenotype:

- 1) Patient continues to receive therapy with an ICS, LABA and LTRA; and
- 2) Patient has experienced a reduction in asthma signs and symptoms including wheezing, chest tightness, coughing, shortness of breath; or
- 3) Patient has experienced a decrease in administration of rescue medication (albuterol); or
- 4) Patient has experienced a decrease in exacerbation frequency; or
- 5) Patient has experienced an increase in predicted FEV₁ from the pretreatment baseline.

Eosinophilic Granulomatosis with Polyangiitis:

1) Patient has demonstrated a positive clinical response to therapy (increase in remission time).

The required trials may be overridden when documented evidence is provided that use of these agents would be medically contraindicated.

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Non-Preferred								
☐ Fasenra	☐ Nucala Auto-Injector ☐ Nucala Pref	illed Syringe						
Strength	Dosage Instructions	Quantity	Days Supply					
Diagnosis:								
Is prescriber and	allergist, immunologist, pulmonologist, or rheuma	tologist?						
☐ Yes, documen	t specialty:							
■ No If no, note of	consultation with specialist:							
Consultation Date:	Physician Name, Specialty & Phone: _							
Will the patient be	e taking requested medication in combination with	another monoclonal antiboo	ly? ☐ No ☐ Yes					
☐ Severe Asthr	na with an Eosinophilic Phenotype:							
Pretreatment bloc	od eosinophil count (attach lab):	Date Obtained:	Date Obtained:					
OR								
Blood eosinophil	count obtained within 12 months prior to initiation	of treatment (attach lab):						
Date Obtained:								
Pretreatment Bas	eline ppFEV ₁ :	Date Obtained:						
Document curren	t use of:							
High-dose inhale	d corticosteroid: Drug Name:	Strength:						
	s:							
Long-Acting Beta	ı2-Agonist: Drug Name:	Strength:						
Dosing Instructions	s:	Trial start dat	e:					
Leukotriene Rece	eptor Antagonist: Drug Name:	Strength:						
Dosing Instructions	3:	Trial start dat	e:					
Does patient have ICS plus a LABA	e a history of two (2) or more exacerbations in the pand LTRA?	previous year despite regula	r use of high-dose					
☐ Eosinophilic	Granulomatosis with Polyangiitis:							
Document trial of	systemic glucocorticoid: Drug Name:	Strength:						
	3:							
	od eosinophil count (attach lab):	Date Obtained:						





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(PLEASE PRINT – ACCURACY IS IMPORTANT) Eosinophil count greater than 10% of the total leukocyte count (attach lab): Date Obtained:_____ For Renewals Only: Severe Asthma with an Eosinophilic Phenotype: Does patient continue to receive therapy with an ICS, LABA and LTRA? □ No ☐ Yes Please indicate if the patient has experienced any of the following (check all that apply): Reduction in asthma signs and symptoms including: wheezing chest tightness 0 coughing shortness of breath ☐ Decrease in administration of rescue medications (albuterol) ☐ Decrease in exacerbation frequency ☐ Increase in ppFEV₁ from the pretreatment baseline Current ppFEV₁: Date Obtained: Please describe: **Eosinophilic Granulomatosis with Polyangiitis:** Has patient demonstrated a positive clinical response to therapy (increase in remission time)? ☐ No Yes, please describe: Medical or contraindication reason to override trial requirements: Attach lab results and other documentation as necessary. Prescriber signature (Must match prescriber listed above.) Date of submission

IMPORTANT NOTE: In evaluating requests for prior authorization the consultant will consider the treatment from the standpoint of medical necessity only. If approval of this request is granted, this does not indicate that the member continues to be eligible for Medicaid. It is the responsibility of the provider who initiates the request for prior authorization to establish by inspection of the member's Medicaid eligibility card and, if necessary by contact with the county Department of Human Services, that the member continues to be eligible for Medicaid.

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