

#### Request for Prior Authorization IL-5 ANTAGONISTS

**FAX Completed Form To**1 (800) 574-2515

**Provider Help Desk** 1 (877) 776-1567

(PLEASE PRINT – ACCURACY IS IMPORTANT)

	•	,		
IA Medicaid Member ID #	Patient name	DOB		
Patient address				
Provider NPI	Prescriber name	Phone		
Prescriber address		Fax		
Pharmacy name	Address	Phone		
Prescriber must complete all information above. It must be legible, correct, and complete or form will be returned.				
Pharmacy NPI	Pharmacy fax	NDC 		

Prior authorization is required for IL-5 antagonists. Requests will not be considered with concurrent use with another monoclonal antibody. Payment for a non-preferred agent will be authorized only for cases in which there is documentation of a previous trial and therapy failure with a preferred agent. Payment will be considered under when patient has an FDA approved or compendia indication for the requested drug under the following conditions:

- I) Request adheres to all FDA approved labeling for requested drug and indication, including age, dosing, contraindications, warnings and precautions, drug interactions, and use in specific populations: and
- 2) Patient has a diagnosis of severe asthma with an eosinophilic phenotype; and
  - a) Patient has a pretreatment blood eosinophil count of ≥150 cells/mcL within the previous 6 weeks or blood eosinophils of ≥300 cells/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
- 3) 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 > 1000 cells/mcL; or
  - ii. Eosinophil count > 10% of the total leukocyte count; or
- 4) Patient has a diagnosis of hypereosinophilic syndrome (HES); and
  - a) Patient has been diagnosed with HES for ≥ 6 months prior to starting treatment; and
  - b) Documentation that non-hematologic secondary causes of HES have been ruled out; and
  - c) Documentation patient does not have FIP1L1-PDGFR $\alpha$  kinase-positive HES; and
  - d) Documentation of ≥ 2 HES flares within the previous 12 months while on stable HES therapy (e.g., chronic or episodic oral corticosteroids, immunosuppressive, or cytotoxic therapy); and
  - e) Patient has a blood eosinophil count ≥ 1,000 cells/mcL; and
  - f) Medication will be used in combination with stable doses of at least one other HES therapy; or
- 5) Patient has a diagnosis of chronic rhinosinusitis with nasal polyps (CRSwNP); and
  - a) Documentation mepolizumab will be used as an add-on maintenance treatment with a nasal corticosteroid spray; and
  - b) Documentation of an adequate trial and therapy failure with at least one preferred medication from each of the following categories:
  - i. Nasal corticosteroid spray; and
  - ii. Oral corticosteroid; and
- 6) Prescribed by or in consultation with an allergist, hematologist, immunologist, otolaryngologist, pulmonologist, or rheumatologist.

If the criteria for coverage are met, an initial authorization will be given for 3 months for a diagnosis of severe asthma with an eosinophilic phenotype and eosinophilic granulomatosis with polyangiitis or 6 months for a diagnosis of hypereosinophilic syndrome or CRSwNP 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:

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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 FEV<sub>1</sub> from the pretreatment baseline.

Eosinophilic Granulomatosis with Polyangiitis:

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

Hypereosinophilic Syndrome:

- 1) Patient has demonstrated a positive clinical response to therapy (improvement of symptoms and/or reduction in the number of flares): and
- 2) Medication continues to be used in combination with stable doses of at least one other HES therapy.

Chronic Rhinosinusitis with Nasal Polyps (CRSwNP):

- 1) Patient has demonstrated positive clinical response to therapy (improvement in symptoms); and
- 2) Continues to receive medication as add-on maintenance therapy with a nasal corticosteroid spray.

The required trials may be over	ridden when documented evidence is	provided that use of these a	gents would be me	dically contraindicated.	
<u>Preferred</u>		Non-Preferred	Non-Preferred		
Fasenra Auto-Injector	☐ Nucala Auto-Injector	☐ Nucala Prefilled S	Syringe		
Strength	Dosage Instructions		Quantity	Days Supply	
Is prescriber and allergist,	hematologist, immunologist, o	tolaryngologist, pulmon	ologist, or rheu	matologist?	
Yes, document specialty	y:				
■ No If no, note consultation	n with specialist:				
Consultation Date:	Physician Name, Specialty & P	hone:			
Will the patient be taking	requested medication in combi	ination with another mo	onoclonal antibo	dy? 🗌 No 📗 Ye	
Severe Asthma with a	n Eosinophilic Phenotype:				
Pretreatment blood eosing		Date Obtained:			
OR					
Blood eosinophil count obt	ained within 12 months prior t	o initiation of treatmen	t (attach lab):		
Date Obtained:	·		` ,		
Pretreatment Baseline ppFEV:			Date Obtained:		

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**Document current use of:** 

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High-dose inhaled corticosteroid: Drug Name:	Strength:	
Dosing Instructions:	Trial start date:	
Long-Acting Beta2-Agonist: Drug Name:	Strength:	
Dosing Instructions:	•	
Leukotriene Receptor Antagonist: Drug Name:	Strength:	
Dosing Instructions:	Trial start date:	
Does patient have a history of two (2) or more exacerbations in the prev plus a LABA and LTRA?   No Yes (provide dates):	ious year despite regular use of high-dose ICS	
Eosinophilic Granulomatosis with Polyangiitis:		
Document trial of systemic glucocorticoid: Drug Name:	Strength:	
Dosing Instructions:	Trial start & end date:	
Pretreatment blood eosinophil count (attach lab):	Date Obtained:	
OR		
Eosinophil count > 10% of the total leukocyte count (attach lab):	Date Obtained:	
☐ Hypereosinophilic Syndrome:		
Has patient been diagnosed with HES for ≥ 6 months prior to starting tre	eatment?	
□ No □ Yes Date of diagnosis:		
Have non-hematologic secondary causes of HES been ruled out?	o 🔲 Yes	
Does patient have FIP1L1-PDGFRα kinase-positive HES? No Y	'es	
Has patient had $\geq$ 2 HES flares within the previous I2 months while on st	able HES therapy?	
☐ No		
Yes Provide dates of HES flares:		
HES therapy & dates of therapy:		
Does patient have a blood eosinophil count $\geq$ 1,000 cells/mcL? $\square$ No	Yes Date obtained:	
Will medication be used in combination with stable doses of at least one	other HES therapy?	
□ No		
Yes Drug Name & Dosing Instructions:		
CRSwNP:		
Will mepolizumab be used as an add-on maintenance treatment with a r	nasal corticosteroid spray?	



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Document at least one preferred drug trial from each of the following categories:

Nasal corticosteroid spray: Drug Name:	•		
Dosing Instructions:			
Oral corticosteroid: Drug Name:	Strength:		
Dosing Instructions:	-		
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 ap	ply):		
Reduction in asthma signs and symptoms including:  wheezing chest tightness coughing shortness of breath			
Decrease in administration of rescue medications (albuterol)			
Decrease in exacerbation frequency	Data Ohasinada		
Increase in ppFEV <sub>1</sub> from the pretreatment baseline Current ppFEV <sub>1</sub> :	Date Obtained:		
Please describe:			
Eosinophilic Granulomatosis with Polyangiitis:			
Has patient demonstrated a positive clinical response to therapy (increase in remis	sion time)?		
□ No			
Yes, please describe:			
Hypereosinophilic Syndrome:			
Has patient demonstrated a positive clinical response to therapy (improvement in number of flares)?	symptoms and/or reduction in the		
□ No			
Yes, please describe:			
Is medication being used in combination with stable doses of at least one other HES	S therapy?		
□ No □ Yes Drug Name: Dosing Instructions:			

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**CRSwNP:** 

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Has patient demonstrated a positive clinical response to therapy (improvement in symptoms)?					
☐ No					
Yes, please describe:					
Does patient continue to receive medication as add-on therapy with a nasal corticosteroid spray?  No Yes; Provide Drug Name & Dose:  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 Health and Human Services, that the member continues to be eligible for Medicaid.

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