

Nevada Medicaid – Molina Healthcare Monoclonal Antibody Agents Prior Authorization Request Form

Please provide the information below. Please print your answer, attach supporting documentation, sign, date, and return to our office as soon as possible to expedite this request. FAX responses to: (844) 259-1689. Phone: (833) 685-2103.

Member Information (required)			Provider Information (required)				
Member Name:			Provider Name:				
Molina ID#:			NPI#:		Specialty:		
Date of Birth:			Office Phone:				
Street Address:			Office Fax:				
City:	State:	Zip:	Office Street Add	dress:			
Phone:			City:		State:	Zip:	
		Medication Ir	nformation (requ	uired)			
Medication Name:			Strength:	Dosage Form:			
Check if requesting brand			Directions for Use:				
Check if request is for conti	nuation of the	erapy					
Drug-Specific Information (required)							
Cinqair® (reslizumab)							
Will the recipient use the requested antiasthmatic monoclonal antibody in combination with other antiasthmatic monoclonal antibodies? □ Yes □ No							
What is the recipient's diag	nosis? 🗆 🛚 S	evere eosinoph	nilic-phenotype a	sthma			
		Other:		ICD-10 Code	(s):		
Is the recipient 18 years of age or older? Yes No							
Is the medication prescribed by or in consultation with a pulmonologist or an allergist/immunologist?							
Is the recipient uncontrolled on current therapy that includes a high dose corticosteroid? U Yes D No							
Is the recipient on a secondary asthma inhaler?							
Is the requested dose to be 3mg/kg via intravenous infusion of 20 to 50 minutes every four weeks?							
If no , please provide the requested dose:							
Will the prescriber submit documentation of the recipient's vaccination status along with this request? Yes No							
Dupixent® (dupilumab)							
Please select the recipient's	s diagnosis b	elow and answe	r the following diag	gnosis-related qu	uestions:		
Atopic Dermatitis							
Does the recipient have a diagnosis of moderate to severe atopic dermatitis? Yes No							
Does the recipient have a trial and failure, contraindication, or intolerance to one medium to high potency topical							
corticosteroid (e.g., betamethasone, triamcinolone)?							
Yes, drug/response: No Does the recipient have a trial and failure or intolerance to any of the following? Tacrolimus topical ointment							
□ Elidel® (pimecrolumus) topical cream □ Recipient is not a candidate for therapy (e.g., immunocompromised)							
Is the medication prescribed by or in consultation with a dermatologist or an allergist/immunologist? \Box Yes \Box No							
Is the request for recertification of Dupixent®?							
If yes , is there documentation of positive clinical response to Dupixent®?							
(Dupixent® (dupilumab) criteria continued on next page)							

Dupixent® (dupilumab) continued

□ Moderate to Severe Asthma

Is the recipient 12 years of age or older? **Yes No**

Is the recipient currently dependent on oral corticosteroids for the treatment of asthma? Is the recipient's asthma of the eosinophilic phenotype as defined by a baseline (pre-treatment) peripheral blood eosinophil level greater than or equal to 150 cells per microliter? **Q Yes O** No Select any of the following that apply to the recipient:

One or more asthma exacerbations requiring systemic corticosteroids within the past 12 months

□ Any prior intubation for an asthma exacerbation

□ Prior asthma-related hospitalization within the past 12 months

Is the recipient currently utilizing one maximally dosed combination ICS/LABA product (e.g., Advair® [fluticasone propionate/salmeterol], Dulera® [mometasone/formoterol], Symbicort® [budesonide/formoterol])?

□ Yes □ No □ Recipient has contraindication/intolerance

Is the recipient currently utilizing both a 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)?
Yes Vo Recipient has contraindication/intolerance Is the medication prescribed by or in consultation with a pulmonologist or an allergist/immunologist?
Q Yes Q No Is the request for recertification of Dupixent®? **U** Yes **U** No

If yes, is there documentation of a positive clinical response to Dupixent® therapy (e.g., reduction in exacerbations,

Chronic Rhinosinusitis with Nasal Polyposis (CRSwNP)

Has the recipient had an inadequate response to two months of treatment with an intranasal corticosteroid (e.g.,

fluticasone, mometasone)?
Q Yes
No
Recipient has contraindication/intolerance

If yes, please document drug(s), dose, duration, and date of trial: _

Will the medication be used in combination with another agent for CRSwNP? **Yes No** Is the medication prescribed by or in consultation with an allergist/immunologist?
Q Yes
Q No Is the request for recertification of Dupixent®? **U** Yes **D** No

If yes, is there documentation of a positive clinical response to Dupixent® therapy? □ Yes □ No

Other diagnosis:

ICD-10 Code(s): ___

Fasenra® (benralizumab)

Will the recipient use the requested antiasthmatic monoclonal antibody in combination with other antiasthmatic monoclonal antibodies?
Q Yes
Q No

What is the recipient's diagnosis? **Devere eosinophilic-phenotype asthma**

Other:_____ICD-10 Code(s): _____

Is the recipient 12 years of age or older? **U** Yes **U** No

Select any of the following that apply to the recipient:

Two or more asthma exacerbations requiring systemic corticosteroids within the past 12 months

□ Any prior intubation for an asthma exacerbation

□ Prior asthma-related hospitalization within the past 12 months

Is the recipient currently utilizing one maximally dosed combination ICS/LABA product (e.g., Advair® [fluticasone propionate/salmeterol], Dulera® [mometasone/formoterol], Symbicort® [budesonide/formoterol])?

□ Yes □ No □ Recipient has contraindication/intolerance

Is the recipient currently utilizing both a 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)? u Yes u No u Recipient has contraindication/intolerance

(Fasenra® (benralizumab) criteria continued on next page)

Is the medication prescribed by or in consultation with a pulmonologist or an allergist/immunologist? Is the request for recertification of Fasenra®? Is the request for recertificati

If yes, is there documentation of a positive clinical response to Fasenra® therapy? □ Yes □ No

Nucala® (mepolizumab)

Please select the recipient's diagnosis below and answer the following diagnosis-related questions:

Severe Asthma

Is the recipient's asthma of the eosinophilic phenotype as defined by a baseline (pre-treatment) peripheral blood eosinophil level greater than or equal to 150 cells per microliter OR peripheral blood eosinophil levels greater than or equal to 300 cells/microliter from within the past 12 months? **Q Yes Q No**

Is the recipient 6 years of age or older? □ Yes □ No

Select any of the following that apply to the recipient:

• One or more asthma exacerbations requiring systemic corticosteroids within the past 12 months

□ Any prior intubation for an asthma exacerbation

□ Prior asthma-related hospitalization within the past 12 months

Is the recipient currently utilizing one maximally dosed combination ICS/LABA product (e.g., Advair® [fluticasone propionate/salmeterol], Dulera® [mometasone/formoterol], Symbicort® [budesonide/formoterol])?

□ Yes □ No □ Recipient has contraindication/intolerance

Is the recipient currently utilizing both a 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)? Yes No Recipient has contraindication/intolerance Is the medication prescribed by or in consultation with a pulmonologist or an allergist/immunologist? Yes No Is the request for recertification of Nucala®? Yes No

If yes, answer the following:

Is there documentation of a positive clinical response to Nucala® therapy (e.g., reduction in exacerbations, improvement in forced expiratory volume in one second [FEV1], decreased use of rescue medications)?

□ Yes (attach documentation) □ No

Is the recipient currently utilizing a combination ICS/LABA product, or an ICS and an additional asthma controller medication? **Yes No**

D Eosinophilic Granulomatosis with Polyangiitis (EGPA)

Has the recipient's disease relapsed or is it refractory to standard of care therapy (i.e., corticosteroid treatment with or without immunosuppressive therapy)? **U Yes D No**

Is the recipient currently receiving corticosteroid therapy? □ Yes □ No

Is the medication prescribed by or in consultation with a pulmonologist, rheumatologist, or allergist/immunologist?

🗆 Yes 🗅 No

Is the request for recertification of Nucala®? □ Yes □ No

If yes, is there documentation of a positive clinical response to Nucala® therapy (e.g., increase in remission time)? □ Yes □ No

Other diagnosis:

ICD-10 Code(s): ____

Xolair® (omalizumab)

Please select the recipient's diagnosis below and answer the following diagnosis-related questions:

Moderate to Severe Persistent Asthma

Will the recipient use the requested antiasthmatic monoclonal antibody in combination with other antiasthmatic

monoclonal antibodies?
D Yes

Is the recipient 6 years of age or older? □ Yes □ No

(Xolair® (omalizumab) criteria continued on next page)

Xolair® (omalizumab) continued					
	test or Radioallergosorbent (RAST) test to a perennial				
aeroallergen? 🗆 Yes 🗅 No					
Is the medication prescribed by a pulmonologist of	allergist/immunologist? 🗆 Yes 🗅 No				
Has the recipient had an inadequate response, ad	verse reaction or contraindication to inhaled, oral corticosteroids?				
Yes, drug/response:	🗆 No				
Has the recipient had an inadequate response, ad	verse reaction or contraindication to a leukotriene receptor				
	<u> </u>				
Please record the recipient's pretreatment serum t	otal Immunoglobulin E (IgE) level:				
Please record the recipient's current weight:					
Please record the requested dose:	mg everyweeks				
Chronic Idiopathic Urticaria (CIU)					
Will the recipient use the requested antiasthmatic	monoclonal antibody in combination with other antiasthmatic				
monoclonal antibodies? 🗅 Yes 🗅 No					
Is the recipient 12 years of age or older?	□ No				
Has the recipient had an inadequate response, ad	verse reaction or contraindication to two different oral second-				
generation antihistamines? Yes, drug names:	D No				
Has the recipient had an inadequate response, ad	verse reaction or contraindication to an oral second-generation				
antihistamine in combination with a leukotriene red	ceptor antagonist?				
Yes, drug names:	🗅 No				
Is the medication prescribed by a dermatologist, r	neumatologist, or allergist/immunologist? 🛛 Yes 🛛 No				
If no, is there documentation in the recipient's m	edical record that a consultation was done by an				
allergist/immunologist, dermatologist or a rheum	atologist regarding the diagnosis and treatment				
recommendations? Yes (attach document	ation) 🗅 No				
Select the requested dose from the following:					
Initial therapy: 150 mg every four weeks					
Initial therapy: 300 mg every four weeks (P	lease provide clinical rationale for starting therapy at this				
dose:)				
Continuation of therapy: 150 mg every four					
Continuation of therapy: 300 mg every four	weeks				
Other:					
□ Other diagnosis:	ICD-10 Code(s):				
Please attach all supporting documentation to request					
Are there any other comments, diagnoses, symptoms, medications	tried or failed, and/or any other information the physician feels is important to				
his review?					

Please note:

This request may be denied unless all required information is received. For urgent or expedited requests please call (833) 685-2103. This form may be used for non-urgent requests and faxed to (844) 259-1689.

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