



FAX Completed Form To 1.833.404.2392

Pharmacy Help Desk 1.800.460.8988

Prescriber Help Desk 1.833.587.2012

Request for Prior Authorization Dupilumab (Dupixent)

(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 Dupixent (dupilumab). Payment will be considered under the following conditions:

- 1) Patient is within the FDA labeled age for indication; and
- 2) Patient has a diagnosis of moderate-to-severe atopic dermatitis; and
 - a. Is prescribed by or in consultation with a dermatologist, allergist, or immunologist; and
 - b. Patient has failed to respond to good skin care and regular use of emollients; and
 - c. Patient has documentation of an adequate trial and therapy failure with one preferred medium to high potency topical corticosteroid for a minimum of 2 consecutive weeks; and
 - d. Patient has documentation of a previous trial and therapy failure with a topical immunomodulator for a minimum of 4 weeks; and
 - e. Patient has documentation of a previous trial and therapy failure with cyclosporine or azathioprine; and
 - f. Patient will continue with skin care regimen and regular use of emollients; or
- 3) Patient has a diagnosis of moderate to severe asthma with an eosinophilic phenotype (with a pretreatment eosinophil count ≥ 150 cells/mcL within the previous 6 weeks) OR with oral corticosteroid dependent asthma; and
 - a. Is prescribed by or in consultation with an allergist, immunologist, or pulmonologist; and
 - b. Has a pretreatment forced expiratory volume in 1 second (FEV₁) ≤ 80% predicted; and
 - c. Symptoms are inadequately controlled with documentation of current treatment with a high-dose inhaled corticosteroid (ICS) given in combination with a controller medication (e.g. long acting beta₂ agonist [LABA], leukotriene receptor antagonist [LTRA], oral theophylline) for a minimum of 3 consecutive months. Patient must be compliant with therapy, based on pharmacy claims; and
 - d. Patient must have one of the following, in addition to the regular maintenance medications defined above:
 - i. Two (2) or more exacerbations in the previous year, or
 - ii. Require daily oral corticosteroids for at least 3 days; or
- 4) Patient has a diagnosis of inadequately controlled chronic rhinosinusitis with nasal polyposis (CRSwNP); and
 - a. Documentation dupilumab will be used as an add-on maintenance treatment; 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

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- ii. Oral corticosteroid; and
- 5) Dose does not exceed the FDA approved dosing for indication.

If criteria for coverage are met, initial authorizations will be given for 16 weeks to assess the response to treatment. Requests for continuation of therapy will require documentation of a positive response to therapy. The required trials may be overridden when documented evidence is provided that use of these agents would be medically contraindicated.

Non-Preferred			
☐ Dupixent			
Strength	Usage Instructions	Quantity	Day's Supply
Diagnosis:			
	Atopic Dermatitis		
Is prescriber a dermatol	ogist, allergist, or immunologist?	,	
Yes specialty:			
☐ No If no, note consu	ltation with dermatologist, allergist,	or immunologist:	
Consultation date:	Physician name, specia	lty & phone:	
Did patient fail to respon	nd to good skin care and regular	use of emollients?	
☐ Yes ☐ No If yes,	provide documentation below:		
Provide skin care regimen	n, including name and dates of emo	llient use:	
Will patient continue ski	n care regimen and regular use o	of emollients? Yes	☐ No
Preferred medium to hig	h potency topical corticosteroid	trial:	
Drug name & dose:		Trial dates:	
Topical immunomodula	tor trial:		
Drug name & dose:		Trial dates:	
Failure reason:			
Cyclosporine or Azathio	prine trial:		
Drug name & dose:		Trial dates:	

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Failure reason:						
Medical or contraindication reason to override trial requirements: Moderate-to-Severe Asthma with an Eosinophilic Phenotype						
						☐ Yes (attach results) ☐
Is prescriber an allergist, im	munologist, or pulmonologist	?				
Yes specialty:	☐ Yes specialty:					
☐ No If no, note consultation	on with allergist, immunologist, o	or pulmonologist:				
Consultation date:	onsultation date:Physician name, specialty & phone:					
Does patient have a pretreat Yes (attach results)	tment FEV₁ ≤ 80% predicted? No					
Document current treatment	t with a high-dose ICS given in	n combination with a controller medication:				
High-Dose ICS Trial:						
Drug name & dose:		Trial dates:				
Failure reason:						
Controller Medication Trial:						
Drug name & dose:		Trial dates:				
Failure reason:						
Does patient have one of the	e following?					
Two (2) or more exacerbations	s in the previous year? Yes	☐ No				
Require daily oral corticostero	ids for at least 3 days? Yes	☐ No				
☐ Inadequately controlled	chronic rhinosinusitis with na	sal polyposis (CRSwNP)				
Will dupliumab be used as a	ın add-on maintenance treatm	ent?				
Yes (document concomitant maintenance treatment): Drug name & dose:						
☐ No						

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Document adequate trial and therapy failure with at least one preferred medication from each of the following categories:

Nasal Corticosteroid Spray Trial:					
Drug name & dose:	Trial dates:				
Failure reason:					
Oral Corticosteroid Trial:					
Drug name & dose:	Trial dates:				
Failure reason:					
Renewal requests:					
Document positive response to therapy:					
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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