

Iowa Health Link

Request for Prior Authorization Dupilumab (Dupixent)

FAX Completed Form To 1 (877) 733-3195 Provider Help Desk 1 (844) 236-1464

(PLEASE PRINT - ACCURACY IS IMPORTANT)

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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 for non-preferred agents will be considered when there is documentation of a previous trial and therapy failure with a preferred agent. Payment will be considered when patient has an FDA approved or compendia indication for the requested drug under the following conditions:

- 1) 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's current weight in kilograms (kg) is provided; and

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- 3) 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 will continue with skin care regimen and regular use of emollients; or
- 4) 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 in adults; < 90% predicted in adolescents 12 to 17 years of age; and < 95% predicted in children 6 to 11 years of age; 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. One (1) or more exacerbations in the previous year, or
 - ii. Require daily oral corticosteroids for at least 3 days; or
- 5) 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
 - ii. Oral corticosteroid; or
- 6) Patient has a diagnosis of eosinophilic esophagitis (EoE); and
 - a. Is prescribed by, or in consultation with, and allergist, gastroenterologist, or immunologist; and
 - b. Patient has ≥ 15 intraepithelial eosinophils per high-power field (eos/hpf) as confirmed by endoscopic esophageal biopsy (attach results); and
 - c. Patient has signs and symptoms of esophageal dysfunction (e.g., dysphagia, food impaction, food refusal, abdominal pain. heartburn regurgitation, chest pain and/or, odynophagia); and
 - d. Documentation of previous trials and therapy failures with all of the following:

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- High dose proton pump inhibitor (PPI) for at least 8 weeks; and i.
- ii. Swallowed topical corticosteroid (e.g., fluticasone propionate, oral budesonide suspension); and
- iii. Dietary therapy; or
- 7) Patient has a diagnosis of moderate to severe prurigo nodularis (PN); and
 - Is prescribed by, or in consultation with an allergist, immunologist, or dermatologist; and a.
 - b. Patient has experienced severe to very severe pruritis, as demonstrated by a current Worst Itch-Numeric Rating Scale (WI-NRS) \geq 7; and
 - c. Patient has \geq 20 nodular lesions (attach documentation); and
 - d. Documentation of a previous trial and therapy failure with a high or super high potency topical corticosteroid for at least 14 consecutive days; and
- Dose does not exceed the FDA approved dosing for indication. 8)

If criteria for coverage are met, initial authorizations will be given for 6 months 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.

Preferred

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Strength	Usage Instructions	Quantity	Day's Supply
Diagnosis:			
Patient's current weight in k	g:Date o	obtained:	
Moderate-to-Severe Ato	pic Dermatitis		
Is prescriber a dermatologist	, allergist, or immunologist?		
Yes specialty:			
No If no, note consultation	n with dermatologist, allergist, or immuno	logist:	
Consultation date:	Physician name, specialty & phon	ne:	
Did patient fail to respond to	good skin care and regular use of er	mollients?	
Yes No If yes, provid	le documentation below:		
Provide skin care regimen, includ	ing name and dates of emollient use:		
Will patient continue skin ca	re regimen and regular use of emolli	ients? 🗌 Yes 🗌 No	
Preferred medium to high pe	otency topical corticosteroid trial:		
Drug name & dose:	7	Trial dates:	
Failure reason:			
Topical immunomodulator t	rial:		
Drug name & dose:	ī	Trial dates:	
Failure reason:			
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Moderate-to-Severe A	sthma with an Eosinophilic Phenotype			
Does patient have pretreatment eosinophil count ≥ 150 cells/mcL within the previous 6 weeks? Yes (attach results) No				
Does patient have oral cor	ticosteroid dependent asthma?			
Is prescriber an allergist, in	nmunologist, or pulmonologist?			
Yes specialty:				
No If no, note consultat	ion with allergist, immunologist, or pulmonologist:			
Consultation date:	Physician name, specialty & phone:			
Provide pretreatment FEV	/1 % predicted (attach results):			
Document current treatm	ent with a high-dose ICS given in combination with a controller medication:			
High-Dose ICS Trial:				
Drug name & dose:	Trial dates:			
Failure reason:				
Controller Medication Tria	al:			
Drug name & dose:	Trial dates:			
Failure reason:				
Does patient have one of t	he following?			
One (I) or more exacerbation	is in the previous year? 🗌 Yes 🗌 No			
Require daily oral corticosterc	oids for at least 3 days? 🔲 Yes 🗌 No			
Inadequately controlle	d chronic rhinosinusitis with nasal polyposis (CRSwNP)			
Will dupliumab be used as	an add-on maintenance treatment?			
Yes (document concomita	nt maintenance treatment): Drug name & dose:			
🗌 No				
Document adequate trial a categories:	and therapy failure with at least one preferred medication from each of the following			
Nasal Corticosteroid Spray	y Trial:			
Drug name & dose:	Trial dates:			
Failure reason:				
Oral Corticosteroid Trial:				
Drug name & dose:	Trial dates:			
Failure reason:				

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Eosinophilic Esophag	itis (EoE)	
ls prescriber an allergist,	immunologist, or gastroenterol	logist?
Yes specialty:		
No If no, note consult	ation with allergist, immunologist, or	gastroenterologist:
Consultation date:	Physician name, specialty δ	& phone:
Does patient have ≥ 15 ir esophageal biopsy?	ntraepithelial eosinophils per hig	h-power field (eos/hpf) confirmed by endoscopic
Yes (attach results)] No	
Does patient have signs a	and symptoms of esophageal dys	sfunction?
Yes; provide signs and sy	mptoms:	
🗌 No		
Document previous trial	s and therapy failures with all of	the following:
High Dose PPI :		
Drug name & dose:		Trial dates:
Failure reason:		
Swallowed topical cortic	osteroid:	
Drug name & dose:		Trial dates:
Failure reason:		
Dietary Therapy:		
Dietary Plan:		Trial dates:
Failure reason:		
Moderate to Severe I	Prurigo Nodularis (PN)	
Is prescriber an allergist,	immunologist, or dermatologis	t?
Yes specialty:		
No If no, note consult	ation with allergist, immunologist, or	dermatologist:
Consultation date:	Physician name, specialty &	& phone:
Worst Itch-Numeric Rat	ing Scale (WI-NRS) response:	Date obtained:
Does patient have \ge 20 n	odular lesions? Yes (provid	de documentation) 🛛 No

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Preferred high or super high potency topical corticosteroid trial:

Drug name & dose:Tr	rial 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 Health and Human Services, that the member continues to be eligible for Medicaid.