AvMed

PHARMACY PRIOR AUTHORIZATION/STEP-EDIT REQUEST*

Directions: The prescribing physician must sign and clearly print name (preprinted stamps not valid) on this request. All other information may be filled in by office staff; fax to 1-305-671-0200. No additional phone calls will be necessary if all information (including phone and fax #s) on this form is correct. If the information provided is not complete, correct, or legible, the authorization process can be delayed.

Drug Requested: Dupixent[®] (dupilumab)

MEMBER & PRESCRIBER INFORMATION: Authorization may be delayed if incomplete.

Member Name:		
	Date of Birth:	
Prescriber Name:		
	Date:	
Office Contact Name:		
	Fax Number:	
NPI #:		
	DN: Authorization may be delayed if incomplete.	
Dosing Schedule:	Length of Therapy:	
Diagnosis:	ICD Code, if applicable:	
Weight (if applicable):	Date weight obtained:	
Diagnosis	Recommended Dose	
Atopic Dermatitis	Adult:	

Atopic Dermatus	 Initial: 600 mg (given as two 300 mg injections) Maintenance: 300 mg once every other week
	 <u>Children ≥ 6 years and Adolescents ≤ 17 years:</u> 15 to < 30 kg - Initial: 600 mg once (administered as two 300 mg injections). Maintenance: 300 mg every 4 weeks 30 to < 60 kg - Initial: 400 mg once (administered as two 200 mg injections). Maintenance: 200 mg every other week 60 kg - Initial: 600 mg once (administered as two 300 mg injections). Maintenance: 300 mg every other week
	 <u>Children ≥ 6 months to 5 years of age:</u> 5 to <15 kg – Initial and maintenance: 200mg (one 200 mg injection) every 4 weeks 15 to < 30 kg – Initial and maintenance: 300mg (one 300 mg injection) every 4 weeks

Diagnosis	Recommended Dose
Asthma, moderate to severe	 Children ≥ 12 years, Adolescents and Adults: Initial: 400 mg (given as two 200 mg injections) or 600 mg (given as two 300 mg injections) Maintenance: 200 mg (following 400 mg initial dose) or 300 mg (following 600 mg initial dose) once every other week
	 <u>Children ≥ 6 years and Adolescents < 12 years:</u> 15 to <30 kg: 100 mg every other week or 300 mg every 4 weeks. ≥ 30 kg: 200 mg every other week
Asthma, oral corticosteroid dependent or with comorbid moderate to severe atopic dermatitis	 Initial: 600 mg (given as two 300 mg injections) Maintenance: 300 mg once every other week
Chronic rhinosinusitis with nasal polyposis	 300 mg once every other week 200 mg syringes are <u>NOT</u> approved for chronic rhinosinusitis with nasal polyposis
Eosinophilic Esophagitis	• Initial and maintenance: 300 mg once every week
Prurigo Nodularis	 Initial: 600 mg (given as two 300 mg injections) Maintenance: 300 mg once every other week

Quantity Limits:

- 100 mg/0.67 mL prefilled syringe: 2 prefilled syringes per 28 days
- 200 mg/1.14 mL pen-injector: 2 pens per 28 days
- 200 mg/1.14 mL prefilled syringe: 2 prefilled syringes per 28 days
- 300 mg/2 mL pen-injector: 2 pens per 28 days
- 300 mg/2 mL prefilled syringe: 2 prefilled syringes per 28 days

*The Health Plan considers the use of concomitant therapy with Adbry[™], Cinqair[®], Dupixent[®], Fasenra[®], Nucala[®], Tezspire[™] and Xolair[®] to be experimental and investigational. Safety and efficacy of these combinations have <u>NOT</u> been established and will <u>NOT</u> be permitted. In the event a member has an active Adbry[™], Cinqair[®], Fasenra[®], Nucala[®], Tezspire[™] or Xolair[®] authorization on file, all subsequent requests for Dupixent[®] will <u>NOT</u> be approved.

CLINICAL CRITERIA: Check below all that apply. <u>All criteria must be met for approval</u>. To support each line checked, all documentation, including lab results, diagnostics, and/or chart notes, must be provided or request may be denied.

Diagnosis: Moderate-to-Severe Atopic Dermatitis

Initial Authorization: 4 months

- □ Prescribed by or in consultation with an allergist, dermatologist or immunologist
- □ Member is 6 months of age or older
- Member has a diagnosis of <u>moderate to severe atopic dermatitis</u> with disease severity confirmed by <u>ONE</u> of the following (chart notes documenting disease severity and BSA involvement must be included):
 - □ Body Surface Area (BSA) involvement >10%
 - □ Eczema Area and Severity Index (EASI) score ≥ 16
 - □ Investigator's Global Assessment (IGA) score ≥ 3
 - □ Scoring Atopic Dermatitis (SCORAD) score ≥ 25
- Member has tried and failed, has a contraindication, or intolerance to <u>ALL</u> four of the following therapies (chart notes documenting contraindication(s) or intolerance must be attached; trials will be verified using pharmacy claims and/or submitted chart notes):
 - □ 30 days (14 days for very high potency) of therapy with <u>ONE</u> medium to very-high potency topical corticosteroid in the past 180 days
 - □ 30 days of therapy with <u>ONE</u> of the following topical calcineurin inhibitors in the past 180 days:
 - □ tacrolimus 0.03 % or 0.1% ointment
 - □ pimecrolimus 1% cream (generic Elidel) [requires prior authorization]
 - □ 90 days of phototherapy (e.g., NB UV-B, PUVA) unless the member is not a candidate and/or has an intolerance or contraindication to therapy
 - □ 90 days of therapy with <u>ONE</u> of the following oral immunosuppressants in the past 180 days:
 - □ azathioprine
 - □ cyclosporine
 - □ methotrexate
 - □ mycophenolate

Diagnosis: Moderate-to-Severe Atopic Dermatitis

Reauthorization: 12 months

□ Member has experienced a positive clinical response to Dupixent[®] therapy (e.g., reduced BSA involvement, decrease in severity based on physician assessment) (chart notes must be submitted)

Diagnosis: Moderate-to-Severe Asthma

Initial Authorization: 12 months

- □ Prescribed by or in consultation with an allergist, immunologist or pulmonologist
- □ Member is 6 years of age or older

- □ Member has been diagnosed with <u>ONE</u> of the following (check the diagnoses below that applies):
 - 1.) Eosinophilic phenotype asthma defined by a baseline (pre-Dupixent[®] treatment) peripheral blood eosinophil level greater than or equal to 150 cells per microliter and meets <u>ALL</u> the following clinical criteria:
 - Member is currently being treated with <u>ONE</u> of the following unless there is a contraindication or intolerance to these medications and must be compliant on therapy <u>for at least 90 consecutive</u> <u>days</u> within a year of request:
 - □ High-dose inhaled corticosteroid (ICS) (e.g., greater than 500 mcg fluticasone propionate equivalent/day) <u>AND</u> an additional asthma controller medication (e.g., leukotriene receptor antagonist, long-acting beta-2 agonist (LABA), theophylline)
 - One maximally dosed combination ICS/LABA product (e.g., Advair[®] (fluticasone propionate/salmeterol), Dulera[®] (mometasone/formoterol), Symbicort[®] (budesonide/formoterol))
 - □ Member has experienced <u>ONE</u> of the following (check box that applies):
 - □ More than 2 exacerbations requiring additional medical treatment (e.g., oral corticosteroids, emergency department, urgent care visits or hospitalizations within the past 12 months)
 - □ Any prior intubation for an asthma exacerbation
 - □ Member has a baseline forced expiratory volume (FEV1) < 80% predicted normal (< 90% for members 6-17 years old) submitted within year of request
 - Provider must submit member blood eosinophil count after a trial and failure of at least 90 days of therapy with high dose inhaled corticosteroids <u>AND</u> long-acting inhaled beta-2 agonist. A failure of these medications is defined as a blood count > 150 cells/microliter (submit labs collected within the past 12 months)

Eosinophil count: _____ Date: _____

2.) Oral corticosteroid dependent asthma and meets <u>ALL</u> the following clinical criteria:

- Member is currently being treated with <u>ONE</u> of the following unless there is a contraindication or intolerance to these medications and must be compliant on therapy <u>for at least 90 consecutive</u> <u>days</u> within a year of request:
 - □ High-dose inhaled corticosteroid (ICS) (e.g., greater than 500 mcg fluticasone propionate equivalent/day) <u>AND</u> an additional asthma controller medication (e.g., leukotriene receptor antagonist, long-acting beta-2 agonist (LABA), theophylline)
 - One maximally dosed combination ICS/LABA product (e.g., Advair[®] (fluticasone propionate/salmeterol), Dulera[®] (mometasone/formoterol), Symbicort[®] (budesonide/formoterol))
- □ Member has experienced <u>ONE</u> of the following (check box that applies):
 - □ More than 2 exacerbations requiring additional medical treatment (e.g., oral corticosteroids, emergency department, urgent care visits or hospitalizations within the past 12 months)
 - □ Any prior intubation for an asthma exacerbation
- □ Member has a baseline forced expiratory volume (FEV1) < 80% predicted normal (< 90% for members 6-17 years old) submitted within year of request

Diagnosis: Moderate-to-Severe Asthma

Reauthorization: 12 months

- □ Member has experienced a sustained positive clinical response to Dupixent[®] therapy as demonstrated by at least <u>ONE</u> of the following (check all that apply; chart notes must be submitted):
 - □ Increase in percent predicted Forced Expiratory Volume (FEV1) from baseline (pre-treatment)
 - □ Reduction in the dose of inhaled corticosteroids required to control asthma
 - □ Reduction in the use of oral corticosteroids to treat/prevent exacerbation
 - Reduction in asthma symptoms such as chest tightness, coughing, shortness of breath or nocturnal awakenings
- □ Member is currently being treated with <u>ONE</u> of the following unless there is a contraindication or intolerance to these medications:
 - □ High-dose inhaled corticosteroid (ICS) (e.g., greater than 500 mcg fluticasone propionate equivalent/day) <u>AND</u> an additional asthma controller medication (e.g., leukotriene receptor antagonist, long-acting beta-2 agonist (LABA), theophylline)
 - □ One maximally dosed combination ICS/LABA product (e.g., Advair[®] (fluticasone propionate/salmeterol), Dulera[®] (mometasone/formoterol), Symbicort[®] (budesonide/formoterol))

Diagnosis: Chronic rhinosinusitis with nasal polyps (CRSwNP)

Initial Authorization: 12 months

- □ Prescribed by or in consultation with an allergist, immunologist or otolaryngologist
- □ Member is 12 years of age or older
- Member has a <u>diagnosis of CRSwNP</u> confirmed by the American Academy of Otolaryngology-Head and Neck Surgery Clinical Practice Guideline (Update): Adult Sinusitis (AAO-HNSF 2015)/American Academy of Allergy Asthma & Immunology (AAAAI) with <u>ONE</u> of the following clinical procedures:
 - □ Anterior rhinoscopy
 - □ Nasal endoscopy
 - □ Computed tomography (CT)
- □ Member has a documented diagnosis of chronic rhinosinusitis defined by at least 12 weeks of the following (chart notes must be submitted):
 - □ Mucosal inflammation <u>AND</u> at least <u>TWO</u> of the following:
 - Decreased sense of smell
 - □ Facial pressure, pain, fullness
 - □ Mucopurulent drainage
 - Nasal obstruction

- Member is currently being treated with medications in at least <u>TWO</u> of the following categories unless there is a contraindication or intolerance to these medications and must be compliant on therapy <u>for at</u> <u>least 90 consecutive days</u> within a year of request (chart notes documenting contraindication(s) or intolerance must be attached; trials will be verified using pharmacy claims and/or submitted chart notes):
 - □ Nasal saline irrigation
 - □ Intranasal corticosteroids (e.g., fluticasone, budesonide, triamcinolone)
 - Leukotriene receptor antagonists (e.g., montelukast, zafirlukast, zileuton)
- □ Member is refractory, ineligible or intolerant to <u>ONE</u> of the following:
 - □ Systemic corticosteroids
 - □ Sino-nasal surgery
- □ Member is requesting Dupixent[®] (dupilumab) as add-on therapy to maintenance intranasal corticosteroids

Diagnosis: Chronic rhinosinusitis with nasal polyps (CRSwNP)

Reauthorization: 12 months

- Member has experienced a positive clinical response to Dupixent[®] therapy (e.g., reduced nasal polyp size, improved nasal congestion, reduced sinus opacification, decreased sino-nasal symptoms, improved sense of smell) (chart notes must be submitted)
- □ Provider documents a reduction in the use of oral corticosteroids (verified by pharmacy paid claims)
- Member has been compliant with Dupixent[®] therapy and continues to receive therapy with an intranasal corticosteroid (verified by pharmacy paid claims)

Diagnosis: Eosinophilic Esophagitis (EoE)

Initial Authorization: 12 months

- □ Prescribed by or in consultation with an allergist, immunologist, pulmonologist or gastroenterologist
- □ Member is 12 years of age or older and weighs at least 40 kg
- Member has a documented diagnosis of EoE as evidenced by at least 15 intraepithelial eosinophils per high-powered microscopy field (eos/hpf), or 60 eosinophils/mm² on endoscopic biopsy (chart notes must be submitted)
- □ Member has a history of an average of at least two (2) episodes of dysphagia, with intake of solids, per week or prior history of esophageal dilation
- □ Provider attests to <u>ONE</u> of the following:
 - □ Member does <u>NOT</u> have a diagnosis of gastroesophageal reflux disease (GERD) and/or GERD diagnosis has been ruled out
 - □ Member has a diagnosis of GERD that is being adequately managed by high dose PPI therapy (e.g., omeprazole 40-80 mg daily)

- □ Provider attestation to other causes of esophageal eosinophilia have been ruled out (i.e., active helicobacter pylori infection, hypereosinophilic syndrome and eosinophilic granulomatosis with polyangiitis, Crohn's disease, ulcerative colitis, celiac disease, achalasia)
- □ Member meets <u>ONE</u> of the following:
 - □ Member has tried an elemental diet or an empiric, 6-food elimination diet (i.e., dairy, eggs, wheat, soy, peanuts, fish/shellfish) to treat/manage eosinophilic esophagitis
 - Provider has determined that the individual is <u>NOT</u> an appropriate candidate for dietary modifications (clinical rationale must be documented in submitted chart notes)
- □ Member has tried and failed swallowed topical glucocorticoids (e.g., nebulized or swallowed nasal drops such as budesonide nasal spray or nebulizer solution) for at least 6 -12 weeks

Diagnosis: Eosinophilic Esophagitis (EoE)

Reauthorization: 12 months

- Member has experienced disease response is indicated by improvement in signs and symptoms compared to baseline in one or more of the following: dysphagia/swallowing pain, including chest pain, stomach pain, heartburn, regurgitation, and vomiting (chart notes must be submitted)
- Member is in histologic remission defined as a peak esophageal intraepithelial eosinophil count of at least 6 eos/hpf

Diagnosis: Prurigo Nodularis (PN)

Initial Authorization: 6 months

- □ Prescribed by or in consultation with an allergist, dermatologist or immunologist
- □ Member is 18 years of age or older
- □ Member has a diagnosis of prurigo nodularis (PN) for at least three (3) months (chart notes must be submitted)
- □ Member's disease is <u>NOT</u> secondary to medications or medical conditions (i.e., neuropathy or psychiatric disease)
- Member has an average worst itch score of at least 7 or greater on the Worst Itch Numeric Rating Scale (WI-NRS 0-10) (chart notes must be submitted)
- Member has at least 20 prurigo nodularis lesions, in total, on legs, arms and/or trunk (chart notes must be submitted)

- Member has tried and failed, has a contraindication, or intolerance to <u>ALL</u> four of the following therapies (chart notes documenting contraindication(s) or intolerance must be attached; trials will be verified using pharmacy claims and/or submitted chart notes):
 - □ 30 days (14 days for very high potency) of therapy with <u>ONE</u> medium to very-high potency topical corticosteroid in the past 180 days
 - □ 30 days of therapy with <u>ONE</u> of the following topical calcineurin inhibitors in the past 180 days:
 - □ tacrolimus 0.03 % or 0.1% ointment
 - □ pimecrolimus 1% cream (generic Elidel) [requires prior authorization]
 - □ 90 days of phototherapy (e.g., NB UV-B, PUVA) unless the member is not a candidate and/or has an intolerance or contraindication to therapy
 - □ 90 days of therapy with <u>ONE</u> of the following oral immunosuppressants in the past 180 days:
 - □ azathioprine
 - □ cyclosporine
 - □ methotrexate

Diagnosis: Prurigo Nodularis (PN)

Reauthorization: 12 months

□ Member has experienced disease response as indicated by improvement (reduction) in signs and symptoms compared to baseline in one or more of the following: pruritus severity, number of lesions, and/or WINRS (chart notes must be submitted)

Medication being provided by Specialty Pharmacy – Proprium Rx

Not all drugs may be covered under every Plan.

If a drug is non-formulary on a Plan, documentation of medical necessity will be required. **Use of samples to initiate therapy does not meet step edit/ preauthorization criteria.** *Previous therapies will be verified through pharmacy paid claims or submitted chart notes.*