AvMed

PHARMACY PRIOR AUTHORIZATION/STEP-EDIT REQUEST*

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

<u>Drug Requested</u>: Xolair[®] (omalizumab) (self-administered) (Pharmacy)

MEMBER & PRESCRIBER INFORM	MATION: Authorization may be delayed if incomplete.
Member Name:	
Member AvMed #:	
Prescriber Name:	
Prescriber Signature:	Date:
Office Contact Name:	
Phone Number:	Fax Number:
DEA OR NPI #:	
DRUG INFORMATION: Authorization	may be delayed if incomplete.
Drug Form/Strength:	
Dosing Schedule:	Length of Therapy:
Diagnosis:	ICD Code, if applicable:
Weight:	Date:
Quantity Limits: 1 syringe/auto-injector/vial	per 28 days
□ 75 mg/0.5 mL auto-injector	•
□ 75 mg/0.5 mL prefilled syringe	
☐ 150 mg/1 mL auto-injector	
□ 150 mg/1 mL prefilled syringe	
□ 150 mg/1.2 mL powder vial	
□ 300 mg/2 mL auto-injector	
□ 300 mg/2 mL prefilled syringe	

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

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

□ **DIAGNOSIS:** <u>Moderate to Severe Persistent Asthma</u> — with a positive skin test or in vitro reactivity to a perennial aeroallergen and symptoms are inadequately controlled with inhaled corticosteroids.

Initial Authorization: 12 months

Recommended Dosage: Maximum dosages will be based on a member weight of 150 kg.

Subcutaneous XOLAIR Doses Every 2 or 4 Weeks* for Patients 12 Years of Age and Older with Asthma

Pretreatment Serum IgE (IU/mL)	Dosing Freq.	Body Weight									
		30-60 kg	>60-70 kg	>70-90 kg	>90-150 kg						
			Dose	(mg)							
≥30-100	Every	150	150	150	300						
>100-200	4	300	300	300	225						
>200-300	weeks	300	225	225	300						
>300-400	Every	225	225	300							
>400-500	2	300	300	375							
>500-600	weeks	300 Insufficient Data									
>600-700		375	375 to Recommend a Dose								

Subcutaneous XOLAIR Doses Every 2 or 4 Weeks* for Pediatric Patients with Asthma Who Begin XOLAIR

Between the Ages of 6 to < 12 years

Pre-treatment	Dosing	Body Weight										
Serum IgE (IU/mL)	Freq.	20-25	>25-30	>30-40	>40-50	>50-60	>60-70	>70-80	>80-90	>90-125	>125-150	
(IO/IIIL)		kg	kg	kg	kg	kg	kg	kg	kg	kg	kg	
		Dose (mg)										
30-100		75	75	75	150	150	150	150	150	300	300	
>100-200		150	150	150	300	300	300	300	300	225	300	
>200-300	Every	150	150	225	300	300	225	225	225	300	375	
>300-400	4	225	225	300	225	225	225	300	300			
>400-500	weeks	225	300	225	225	300	300	375	375			
>500-600		300	300	225	300	300	375					
>600-700		300	225	225	300	375						
>700-800		225	225	300	375							
>800-900	_	225	225	300	375							
>900-1000	Every	225	300	375		Insufficient Data to Recommend a Dose						
>1000-1100	weeks	225	300	375		IIISUIII	cient Da	ita to Ke	comme	iu a Dose		
>1100-1200		300	300									
>1200-1300		300	375									

	Pre	escribed by or in consultation with an allergist or pulmonologist						
		s the member been approved for Xolair® previously through AvMed medical department? Yes □ No						
Member is currently being treated with ONE of the following unless there is a contraindication of intolerance to these medications and must be compliant on therapy for at least 90 consecutive day within a year of request:								
		Medium to 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))						
	Me	ember must meet ONE of the following:						
		Member is ≥ 6 and < 12 years of age with a pre-treatment IgE level of 30-1300 Member is ≥ 12 years of age with a pre-treatment IgE level of 30-700						
		IgE level: Test Date:						
	Me	ember has experienced ONE of the following (check box that applies):						
		More than > 2 exacerbations requiring additional medical treatment (e.g., an increase in oral corticosteroid dose, emergency department, urgent care visits or hospitalizations) within the past 12 months						
		Any prior intubation for an asthma exacerbation						
D	iag	nosis: Moderate-to-Severe Persistent Asthma						
lea [*]	uth	orization: 12 months						
		ember has experienced a sustained positive clinical response to Xolair® therapy as demonstrated by at st ONE 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						
		ember is currently being treated with <u>ONE</u> of the following unless there is a contraindication or olerance to these medications:						
		Medium to 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)						

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DIAGNOSIS: Chronic Idiopathic Urticaria									
niti	al Authorization: 12 months								
ecommended Dosage: 150 mg or 300 mg by subcutaneous injection every 4 weeks									
	Prescribed by or in consultation with an allergist or pulmonologist								
	Member is > 12 years of age								
	Member has failed ONE (1) of the foweeks:	ollowing H1 antihistamines at 4 tim	nes the initial dose for at least 4						
	□ levocetirizine 10 mg – 20 mg QD	☐ desloratadine 10 – 20 mg QD	☐ fexofenadine 120 mg − 240 mg BID						
	□ cetirizine 20 mg – 40 mg QD	□ loratadine 20 mg – 40 mg QD							
	Member has remained symptomatic of pharmacy paid claims):	despite treatment with <u>ALL</u> the following	lowing therapies (verified by						
	☐ Hydroxyzine 10 mg – 25 mg take	·							
	-	st 4 weeks (e.g., montelukast, zafir	•						
	H2 antihistamine, for treatment of cimetidine)	f acute exacerbations, for at least 5	days (e.g., famotidine,						
ı D	iagnosis: Chronic Idiopathic U	J rticaria							
Rea	uthorization: 12 months								
	Members disease status has been re-e condition warrants continued treatme								
	Provider has submitted chart notes do in the number of hives, a decrease in	• • • • • • • • • • • • • • • • • • • •	• • •						
	Symptoms returned when the Xolair® (chart notes must be submitted for of therapy beyond the next dosing it	documentation supporting taper	ing of dose and/or withholding						

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□ DIAGNOSIS: Chronic Rhinosinusitis with Nasal Polyps (CRSwNP)

Initial Authorization: 12 months

Recommended Dosage:

Pretreatment Serum IgE (IU/mL)	Dosing	Bodyweight								
271.11	Freq.	>30-40 kg	>40-50 kg	>50-60 kg	>60-70 kg	>70-80 kg	>80-90 kg	>90-125 kg	> 125-150 kg	
		17		*** ***	Dose	(mg)	•	10		
30 - 100		75	150	150	150	150	150	300	300	
>100 - 200		150	300	300	300	300	300	450	600	
>200 - 300	_	225	300	300	450	450	450	600	375	
>300 - 400	Every 4	300	450	450	450	600	600	450	525	
>400 - 500	Weeks	450	450	600	600	375	375	525	600	
>500 - 600		450	600	600	375	450	450	600		
>600 - 700		450	600	375	450	450	525			
>700 - 800		300	375	450	450	525	600			
>800 - 900		300	375	450	525	600				
>900 - 1000	Euros	375	450	525	600					
>1000 - 1100	Every 2	375	450	600						
>1100 - 1200	Weeks	450	525	600	Insu	ıfficient Da	nta to Reco	ommend a	Dose	
>1200 - 1300		450	525							
>1300 - 1500		525	600							

Prescribed by or in consultation with an allergist, immunologist, or	otolaryngologist
Pre-treatment IgE level of 30-1500:	Test Date:
Member is 18 years of age or older	
Member has a <u>diagnosis of CRSwNP</u> confirmed by the American and Neck Surgery Clinical Practice Guideline (Update): Adult Sinu Academy of Allergy Asthma & Immunology (AAAAI) with <u>ONE</u>	usitis (AAO-HNSF 2015)/American
☐ Anterior rhinoscopy	
□ Nasal endoscopy	
☐ Computed tomography (CT)	

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Ц		ocumented diagnosis of chronic rhinosinusitis defined by at least 12 weeks of the following (chart note ust be submitted):
		Mucosal inflammation AND at least two of the following:
		□ Decreased sense of smell
		☐ Facial pressure, pain, fullness
		☐ Mucopurulent drainage
		□ Nasal obstruction
	is a	ember is currently being treated with medications in at least <u>two</u> of the following categories unless ther a contraindication or intolerance to these medications and <u>must</u> be compliant on therapy <u>for at least 90 insecutive days</u> within a year of request (chart notes documenting contraindication(s) or intolerance ust 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)
	Me	ember is refractory, ineligible, or intolerant to ONE of the following:
		Systemic corticosteroids
		Sino-nasal surgery
	Me	ember is requesting Xolair® (omalizumab) as add-on therapy to maintenance intranasal corticosteroids
		ember has had an unsuccessful 6-month trial of Dupixent® (dupilumab) OR Nucala® (mepolizumab) erified by pharmacy paid claims)
C	hro	onic Rhinosinusitis with Nasal Polyps (CRSwNP)
<u>Rea</u>	<u>uth</u>	orization: 12 months.
	im	ember has experienced a positive clinical response to Xolair® therapy (e.g., reduced nasal polyp size, proved nasal congestion, reduced sinus opacification, decreased sino-nasal symptoms, improved sense smell) (please submit chart notes)
	Me	ember has decreased utilization of oral corticosteroids (verified by pharmacy paid claims)
		ember has been compliant on Xolair® therapy and continues to receive therapy with an intranasal rticosteroid (verified by pharmacy paid claims)
		(Continued on next page)

□ DIAGNOSIS: Immunoglobulin (Ig) E-Mediated Food Allergy

Initial Authorization: 12 months

Recommended Dosage:

Pretreatment Serum IgE (IU/mL)	Dosing						Body	Weight	(kg)					
	Freq.	≥10-12	>12-15	>15-20	>20-25	>25-30	>30-40	>40-50	>50-60	>60-70	>70- 80	>80-90	>90 - 125	>125 - 150
							Do	se (mg)						
≥30 - 100		75	75	75	75	75	75	150	150	150	150	150	300	300
>100 - 200		75	75	75	150	150	150	300	300	300	300	300	450	600
>200 - 300	F	75	75	150	150	150	225	300	300	450	450	450	600	375
>300 - 400	Every 4 Weeks	150	150	150	225	225	300	450	450	450	600	600	450	525
>400 - 500	Weeks	150	150	225	225	300	450	450	600	600	375	375	525	600
>500 - 600		150	150	225	300	300	450	600	600	375	450	450	600	
>600 - 700		150	150	225	300	225	450	600	375	450	450	525		
>700 - 800		150	150	150	225	225	300	375	450	450	525	600		
>800 - 900		150	150	150	225	225	300	375	450	525	600			
>900 - 1000	Every	150	150	225	225	300	375	450	525	600				
>1000 - 1100	2 Weeks	150	150	225	225	300	375	450	600					
>1100 - 1200		150	150	225	300	300	450	525	600	Insuff	icient (lata to R Dose	ecomn	end a
>1200 - 1300		150	225	225	300	375	450	525						
>1300 - 1500		150	225	300	300	375	525	600						
>1500 - 1850			225	300	375	450	600							

- \square Member is ≥ 1 year of age
- ☐ Prescribed by or in consultation with an allergist or immunologist
- □ Member has a baseline immunoglobulin (Ig)E level ≥ 30 IU/mL Note: "Baseline" is defined as prior to receiving any treatment with Xolair® or another monoclonal antibody therapy that may lower IgE levels (e.g., Dupixent® [dupilumab subcutaneous injection], Tezspire™ [tezepelumab-ekko subcutaneous injection]).
- ☐ Member must meet **BOTH** of the following:
 - ☐ Member has a positive skin prick test response to one or more foods
 - ☐ Member has a positive in vitro test (i.e., a blood test) for IgE to one or more foods

	Provider attests member has a history of an allergic reaction to a food that met <u>ALL</u> the following:
	☐ Member demonstrated signs and symptoms of a significant systemic allergic reaction (e.g., hives, swelling, wheezing, hypotension, and gastrointestinal symptoms)
	☐ Reaction occurred within a short period of time following a known ingestion of the food
	☐ Prescriber deemed this reaction significant enough to require a prescription for an epinephrine auto injector (e.g., EpiPen, EpiPen Jr., Auvi-Q, and generic epinephrine auto-injectors)
	Member has been prescribed an epinephrine auto-injector
	Provider attests Xolair® will be used in conjunction with a food allergen-avoidant diet
	Medication will NOT be used in conjunction with Palforzia® or oral immunotherapy (OIT)
□ D	IAGNOSIS: Immunoglobulin (Ig) E-Mediated Food Allergy
To su	uthorization: 12 months. Check below all that apply. All criteria must be met for approval. apport each line checked, all documentation, including lab results, diagnostics, and/or chart notes, be provided or request may be denied.
	Member is compliant with Xolair® therapy
	Provider attests Xolair® will continue to be used in conjunction with a food allergen-avoidant diet
	Member has been prescribed an epinephrine auto-injector

$\label{eq:medication} \textbf{Medication being provided by a Specialty Pharmacy}-\textbf{Proprium }\textbf{Rx}$

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.