

COMMONWEALTH OF VIRGINIA DEPARTMENT OF MEDICAL ASSISTANCE SERVICES Service Authorization (SA) Form

Xolair® (omalizumab)

If the following information is not complete, correct, or legible, the SA process can be delayed.

Please use one form per member.

MEMBER INFORMATION		
Last Name:	First Name:	
Medicaid ID Number:	Date of Birth:	
	Weight in Kilograms:	
PRESCRIBER INFORMATION		
Last Name:	First Name:	
NPI Number:		
Phone Number:	Fax Number:	
DRUG INFORMATION		
Drug Name/Form:		
Chronoth.		
Dosing Frequency:		
Length of Therapy:		
Quantity per Day:		
	ssistance Services considers the use of concomitant therapy with	

The Virginia Department of Medical Assistance Services considers the use of concomitant therapy with Cinqair®, Dupixent®, Fasenra®, Nucala®, Tezspire™ and Xolair® to be experimental and investigational. Safety and efficacy of theses combinations have **NOT** been established and will **NOT** be permitted.

(Form continued on next page.)

M	ember's Last Name: Member's First Name:				
DI	DIAGNOSIS AND MEDICAL INFORMATION				
Fo	r severe* asthma initial approval, complete the following questions to receive a 6-month approval:				
1.	Is the member 6 years of age or older? AND Yes No				
2.	Does the member have a diagnosis of severe *asthma? AND Yes No				
3.	Does the member have a positive skin test or in vitro reactivity to a perennial aero-allergen; AND Yes No				
4.	Does the member weigh between 20 kg (44 lbs.) and 150 kg (330 lbs.); AND Yes No				
5.	Does the member have serum total IgE level, measured before the start of treatment, of either: • ≥ 30 IU/mL and ≤ 700 IU/mL in patients age ≥ 12 years; OR • ≥ 30 IU/mL and ≤ 1300 IU/mL in patients age 6 to < 12 years; AND ☐ Yes ☐ No				
6.	Will coadministration with another monoclonal antibody be avoided (e.g., mepolizumab, reslizumab, benralizumab, dupilumab, tezepelumab-ekko)? AND Yes No				
7.	 Will this be used for add-on maintenance treatment in members regularly receiving both (unless otherwise contraindicated) of the following: Medium-to high-dose inhaled corticosteroids; AND An additional controller medication (e.g., long-acting beta agonist, leukotriene modifiers)? Yes No 				
8.	corticosteroid treatment (in addition to the regular maintenance therapy defined above) or one exacerbation resulting in a hospitalization? AND Yes No				
(Fo	orm continued on next page.)				

Member's First Name:

Member's Last Name:

 9. Does the member have at least one of the following for assessment of clinical status: Use of systemic corticosteroids Use of inhaled corticosteroids Number of hospitalizations, ER visits, or unscheduled visits to healthcare provider due to Forced expiratory volume in 1 second (FEV₁)? Yes No 	condition
For severe* asthma renewal, complete the following questions to receive a 12-month approval:	
10. Has the member been assessed for toxicity? AND Yes No	
 11. Does the member have improvement in asthma symptoms or asthma exacerbations as evidence decrease in one or more of the following: Use of systemic corticosteroids Hospitalizations ER visits Unscheduled visits to healthcare provider Improvement from baseline in forced expiratory volume in 1 second (FEV₁)? Yes No 	ed by
For chronic idiopathic urticartia/chronic spontaneous urticaria initial approval, complete the folloquestions to receive a 6-month approval:	owing
12. Is the member 12 years of age or older? AND	
☐ Yes ☐ No	
 13. Is the underlying cause of the patient's condition is NOT considered to be any other allergic corother form(s) of urticaria? AND Yes No 	idition(s) o
14. Is the member avoiding triggers (e.g., NSAIDs, etc.)? AND Yes No	
15. Documented baseline score from an objective clinical evaluation tool, such as: urticaria activity (UAS7), angioedema activity score (AAS), Dermatology Life Quality Index (DLQI), Angioedema C Life (AE-QoL), urticaria control test (UCT), angioedema control test (AECT), or Chronic Urticaria Life Questionnaire (CU-Q2oL)? AND Yes No (Form continued on next page.)	uality of
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Member's Last Name:	Member's First Name:	
16. Has the member had an inadequate response to a scheduled dosing of a second-generation H1-antihi	•	
17. Has the member had an inadequate response to a scheduled dosing of at least one of the following:	e to a one or more-month trial on previous therapy with ving:	
 Up-dosing/dose advancement (up to 4-fold) of 	of a second generation H1-antihistamine	
 Add-on therapy with a leukotriene antagonist 	: (e.g., montelukast, zafirlukast, etc.)	
 Add-on therapy with another H1-antihistamir 	1e**	
 Add-on therapy with a H2-antagonist (e.g. rar Yes No 	nitidine, famotidine, etc.)	
For chronic idiopathic urticartia/chronic spontaneous receive a 12-month approval:	urticaria renewal, complete the following questions t	
18. Has the member been assessed for toxicity? AND		
Yes No		
19. Does the member have a clinical improvement as of UAS7, AAS, DLQI, AE-QoL, UCT, AECT, CU-Q2oL, etc	documented an objective clinical evaluation tool? (e.g.,	
For chronic rhinosinusitis with nasal polyps (CRSwNP) receive a 6-month approval:	initial approval, complete the following questions to	
20. Is the member 18 years of age or older? AND Yes No		
21. Has the member failed on at least 8 weeks of intra-	nasal corticosteroid therapy? AND	
22. Does the member have at least 3 of the following in history of sino-nasal surgery are only required to have	ndicators for biologic treatment (note: members with a ave at least 3 of the indicators):	
 Patient has evidence of type 2 inflammation (150 cells/μL, or total IgE ≥ 100 IU/mL) 	e.g., tissue eosinophils ≥ 10/hpf, blood eosinophils ≥	
 Patient has required ≥ 2 courses of systemic of corticosteroids, unless contraindicated 	corticosteroids per year or >3 months of low dose	
 Disease significantly impairs the patient's qua 	•	
Patient has experienced significant loss of sm		
 Patient has a comorbid diagnosis of asthma; Yes No 	AND	

(Form continued on next page.)

IVI	ember's Last Name: Member's First Name:
23	 The member does not have any of the following: Antrochoanal polyps Nasal septal deviation that would occlude at least one nostril Disease with lack of signs of type 2 inflammation Cystic fibrosis Mucoceles; AND Yes No
24	. Have other causes of nasal congestion/obstruction been ruled out (e.g., acute sinusitis, nasal infection or upper respiratory infection, rhinitis medicamentosa, tumors, infections, granulomatosis)? AND Yes No
25	. Has the physician assessed baseline disease severity utilizing an objective measure/tool? AND Yes No
26	Will therapy be used in combination with intranasal corticosteroids unless unable to tolerate or is contraindicated?YesNo
Fo	r CRSwNP renewal, complete the following questions to receive a 12-month approval:
27	. Has the member been assessed for toxicity? AND Yes No
28	b. Does the member have disease response as indicated by improvement in signs and symptoms compared to baseline in one or more of the following: nasal/obstruction symptoms, improvement of sinus opacifications as assessed by CT-scans and/or an improvement on a disease activity scoring tool [e.g., nasa polyposis score (NPS), nasal congestion (NC) symptom severity score, sinonasal outcome test-22 (SNOT-22), etc.]? OR
29	 Did the member have improvement in at least one of the following response criteria: Reduction in nasal polyp size Reduction in need for systemic corticosteroids Improvement in quality of life Improvement in sense of smell Reduction of impact of comorbidities? Yes
	(Form continued on next page.)

Me	ember's Last Name: Member's First Name:		
	For IgE-Mediated Food Allergy initial approval, complete the following questions to receive a 6-month approval:		
1.	Is the member 1 year of age or older? AND		
	Yes No		
2.	Is the prescribing physician an allergist or immunologist or has an allergist or immunologist been consulted? AND		
	Yes No		
3.	Does the member have a diagnosed food allergy as confirmed by:		
	a. A positive skin prick test under a drop of allergen extract; ORb. A positive IgE screening to identified foods? AND		
	Yes No		
4.	Will the member continue to practice allergen avoidance?		
	☐ Yes ☐ No		
	r IgE-Mediated Food Allergy initial renewal, complete the following questions to receive a 12-month proval:		
1.	Has the member has been assessed for toxicity? AND		
	☐ Yes ☐ No		
2.	Is the member experiencing a clinical response and improvement as attested by the prescriber?		
	☐ Yes ☐ No		
	* Components of severity for classifying asthma as severe may include any of the following (not all-inclusive):		
•	Symptoms throughout the day Nighttime awakenings, often 7 times/week SABA use for symptom control occurs several times per day Extremely limited normal activities Lung function (percent predicted FEV ₁) < 60% Exacerbations requiring oral systemic corticosteroids are generally more frequent and intense relative to moderate asthma		
Ву	escriber Signature (Required) r signature, the physician confirms the above information is accurate and verifiable by member records.		
	ease include ALL requested information; Incomplete forms will delay the SA process. bmission of documentation does NOT guarantee coverage by the Department of Medical Assistance Services.		
Pri Ati P.(e completed form may be: FAXED TO 800-932-6651 , phoned to 800-932-6648, or mailed to: me Therapeutics Management LLC tn: GV – 4201 D. Box 64811 Paul, MN 55164-0811		