MONOCLONAL ANTIBODIES (MABs) — ANTI-IL, ANTI-IgE, ANTI-TSLP PRIOR AUTHORIZATION FORM





(form effective 1/6/2025)

Fax to PerformRx[™] at **1-866-497-1387**, or to speak to a representative call **1-800-588-6767**.

PRIOR AUTHOR	ZATION REQU	EST INFORMATION						
□ New request □ Renewal request Total # of pages:								
Name of office contact:			Contact's phone number:		LTC fac	LTC facility contact/phone:		
PATIENT INFORM	MATION							
Patient name:				Patient ID #:			DOB:	
Street address:								
Apt #: City/state/zip:				Phone:				
PRESCRIBER INF	ORMATION							
Prescriber name:								
Specialty:				NPI:			State license #:	
Street address:								
Suite #:	#: City/state/zip:							
Phone:	ione:			Fax:				
CLINICAL INFOR	MATION		I					
Medication requested: Strength:								
Preferred Medications:			Non-Preferred Medications:			losage form (pen, vial, etc):		
□ Fasenra Pen	Tezspire Pen	🗆 Cinqair Vial						
Hasenra Syringe Nucala Autoinjector	□ Fasenra Syringe □ Tezspire Syringe □ Nucala Autoinjector □ Xolair Autoinjector							
Nucala Vial Nucala Vial Xolair Syringe Xolair Vial								
Dose and directions:				Quantity:		D	Ouration: months	
Diagnosis:				Dx code <u>(required)</u> :		N	Veight: Ibs/kg	
Has the beneficiary used the requested medication in the past 90 days? Submit documentation.						C	□ Yes – date of last dose:	
							□ No	
Is the requested medication being prescribed by or in consultation with a specialist?						C	Submit documentation of	
							□ No consultation, if applicable.	
PHARMACY INFORMATION (Prescriber to identify the pharmacy that is to dispense the medication):								
Deliver to: 🗆 Patient's Home 🔹 Physician's Office 🔅 Patient's Preferred Pharmacy Name:								
NPI#:								
Pharmacy Phone #: Pharmacy Fax #: I acknowledge that the patient agrees with the pharmacy chosen for delivery of this medication.								
		the pharmady broben for derivery e						
INITIAL REQUES	15	Complete all sections t	that apply to	the benefic	iary and this request.			
Check all that apply and submit documentation for each item.								
For a non-preferred drug in this class: Does the beneficiary have a history of trial and failure of preferred agents in this class that are approved or medically accepted for treatment of the be Refer to https://papdl.com/preferred-drug-list for a list of preferred and non-preferred agents are approved or medically accepted for treatment of the berefit to https://papdl.com/preferred-drug-list for a list of preferred and non-preferred agents are approved or medically accepted for treatment of the berefit to https://papdl.com/preferred-drug-list for a list of preferred and non-preferred agents are approved or medically accepted for treatment of the berefit to https://papdl.com/preferred-drug-list for a list of preferred and non-preferred agents are approved or the berefit to https://papdl.com/preferred-drug-list for a list of preferred and non-preferred agents are approved or the berefit to https://papdl.com/preferred-drug-list for a list of preferred and non-preferred agents are approved or the berefit to https://papdl.com/preferred-drug-list for a list of preferred and non-preferred agents are approved or the berefit to https://papdl.com/preferred-drug-list for a list of preferred and non-preferred agents are approved or the berefit to https://papdl.com/preferred-drug-list for a list of preferred agents are approved or the berefit to https://papdl.com/preferred-drug-list for a list of preferred agents are approved or the berefit to https://papdl.com/preferred.com/preferred-drug-list for a list of preferred agents are approved or the berefit to https://papdl.com/preferred.com/preferred-drug-list for a list of preferred agents are approved or the berefit to https://papdl.com/preferred-drug-list for a list of preferred-drug-list for a list of preferred agents are approved or the berefit to https://papdl.com/preferred-drug-list for a list of preferred-drug-list for a list of prefer				eneficiary's condition?		the [\Box Yes \Box List medications tried:	
						0	□ No	
1. For treatment of AS	THMA:							
		nt with the FDA-approved indication f	for the		n anti-IgE MAB (e.g., XOLA			
requested medication despite use of maximal therapeutic doses of or has contraindication or an intolerance to the following (check all that apply):							ma induced by an unavoidable perennial	
□ inhaled glucocorticoid					 Diagnosis confirmed by positive skin test or radioallergosorbent test (RAST Has a pretreatment serum total IgE measurement of: 			
□ long-acting beta-agonist (LABA) □ leukotriene modifier				, , , , , , , , , , , , , , , , , , ,				
 other (e.g., tiotropium, theophylline): Will continue to use maximal standard asthma controller medications in addition to the requested medication 					□ For an anti-IL MAB (e.g., CINQAIR □ Has asthma of an eosinophilic ph		NRA, NUCALA): De – Absolute blood eosinophil count:	
					/mL Date obtain			
			□ Has severe asthma			E).		
					□ For an anti-TSLP (e.g., TEZSPIRE): □ Has severe asthma			

INITIAL REQUESTS (continued)

2. For treatment of CHRONIC SPONTANEOUS (IDIOPATHIC) URTICARIA:

- \Box Has a history of urticaria for a period of ≥ 6 weeks
- Requires use of systemic steroids to control urticarial symptoms
- □ Tried and failed the maximally tolerated dose of an H1 antihistamine (e.g., cetirizine/levocetirizine, fexofenadine, loratadine/desloratadine) taken for at least two weeks or has a contraindication or an intolerance to H1 antihistamines

3. For treatment of EGPA:

- \Box Has a history of asthma
- □ Has an absolute blood eosinophil count ≥1000/microliter
- \Box Has a blood eosinophil level >10% of leukocytes
- □ Has evidence of the following *(check all that apply)*:
 - □ histopathological evidence of:
 - eosinophilic vasculitis
 - perivascular eosinophilic infiltration
 - $\hfill\square$ eosinophil-rich granulomatous inflammation
 - □ neuropathy (nerve deficit or conduction abnormality)
 - pulmonary infiltrates, non-fixed
 - □ sino-nasal abnormality
 - □ cardiomyopathy
 - □ glomerulonephritis
 - alveolar hemorrhage
 - □ palpable purpura
 - □ positive test for ANCA
- Requires systemic glucocorticoids to maintain remission
- □ Has a contraindication or an intolerance to systemic glucocorticoids
- □ Has severe EGPA as defined by national treatment guidelines
- □ Tried and failed or has a contraindication or an intolerance to rituximab or cyclophosphamide

4. For treatment of HYPEREOSINOPHILIC SYNDROME (HES):

- □ Has documented FIP1L1-PDGFRA-negative HES
- □ Has organ damage or dysfunction
- □ Has a blood eosinophil count ≥1000/microliter
- □ Requires or has required systemic glucocorticoids to maintain remission
- Has a contraindication or an intolerance to systemic glucocorticoids

5. For treatment of NASAL POLYPS:

- □ Has a history of trial and failure of or contraindication or intolerance to nasal corticosteroids
- □ For an anti-IgE MAB (e.g., XOLAIR):
 - □ Has a pretreatment serum total IgE measurement of:
- 6. For treatment of ALL OTHER DIAGNOSES:
 - List other treatments tried (including start/stop dates, dose, outcomes):

RENEWAL REQUESTS

1. For treatment of ASTHMA:

- Experienced measurable evidence of improvement in the severity of the asthma condition
- □ Will continue to use optimally titrated doses of or has a contraindication or an intolerance to the following (check all that apply):
 - □ inhaled glucocorticoid
 - □ leukotriene modifier
 - □ long-acting beta-agonist (LABA)
 - □ other (e.g., tiotropium, theophylline):

2. For treatment of CHRONIC SPONTANEOUS (IDIOPATHIC) URTICARIA:

- □ Experienced an improvement in symptoms
- Document rationale for continued use:

3. For treatment of EGPA:

□ Experienced measurable evidence of improvement in disease activity □ Reduction in use of systemic glucocorticoids for the treatment of EGPA

4. For treatment of HYPEREOSINOPHILIC SYNDROME (HES):

- Experienced measurable improvement in disease activity
 - Reduction in use of systemic glucocorticoids for the treatment of HES

PLEASE FAX COMPLETED FORM WITH REQUIRED CLINICAL DOCUMENTATION

Prescriber signature:

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Date