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





(form effective 1/6/2025)

Fax to PerformRx<sup>SM</sup> at **1-855-851-4058**, or to speak to a representative call **1-866-907-7088**.

PRIOR AUTHORIZ	ATION REQUEST	INFORMATION			
☐ New request ☐ Renewal request ☐ Total # of pages:		Total # of pages:			
Name of office contact: Contact			phone number:	LTC facility contact/phone:	
PATIENT INFORMATION					
Patient name:			Patient ID #:	DOB:	
Street address:					
Apt #:	City/state/zip:		Phone:		
PRESCRIBER INFO	ORMATION				
Prescriber name:					
Specialty:			NPI:	State license #:	
Street address:					
Suite #: City/state/zip:					
Phone: Fa			Fax:		
CLINICAL INFORMATION					
Medication requested:				Strength:	
Preferred Medications:			Non-Preferred Medications:	Dosage form (pen, vial, etc):	
☐ Fasenra Pen	□T	ezspire Pen	☐ Cinqair Vial	boodge form (port, viai, oto).	
☐ Fasenra Syringe	□⊺	ezspire Syringe			
☐ Nucala Autoinjector		Kolair Autoinjector			
☐ Nucala Vial		Colair Syringe			
Dose and directions:		(olair Vial	Quantity:	Duration: months	
			Dx code (required):	Weight: lbs/kg	
Diagnosis:	h	in the count of decision of the count of the			
Has the beneficiary used the requested medication in the past 90 days? Submit documentation.					
				□ No	
Is the requested medication being prescribed by or in consultation with a specialist?				☐ Yes Submit documentation of consultation, if applicable.	
PHARMACY INFORMATION (Prescriber to identify the pharmacy that is to dispense the medication):					
Deliver to: ☐ Patient's Hon					
NPI#:					
Pharmacy Phone #:			Pharmacy Fax #:		
☐ I acknowledge that the patient agrees with the pharmacy chosen for delivery of this medication.					
INITIAL REQUEST	S				
Complete all sections that apply to the beneficiary and this request.  Check all that apply and <u>submit documentation</u> 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 ben Refer to https://papdl.com/preferred-drug-list for a list of preferred and non-preferred agen				he ☐ Yes ☐ List medications tried:	
				□ No	
1. For treatment of ASTHMA:					
☐ Has an asthma severity that is consistent with the FDA-approved indication for the ☐ For an anti-IgE MAB (e.g., XOLAIR):					
requested medication despite use of maximal therapeutic doses of or has 🗆 Has moderate-to-severe persistent asthma induced by an unavoidable perennial					
contraindication or an intolerance to the following (check all that apply):  allergen (pollen, mold, dust mites, etc.)  Diagnosis confirmed by positive skin test or radioallergosorbent test (RAST)					
□ Innated glucocordicold □ Diagnosis confirmed by positive skin test or radioaliergosorbent test (RAST) □ Iong-acting beta-agonist (LABA) □ Has a pretreatment serum total IgE measurement of:					
□ leukotriene modifier □ For an anti-II MAR (e.g. CINDAIR FASENRA NUCALA):					
□ other (e.g., tiotropium, theophylline):			☐ Has asthma of an eosinophilic phenotype — Absolute blood eosinophil count:  /mL Date obtained: ☐ Has severe asthma		
☐ Will continue to use maximal standard asthma controller medications in addition to the requested medication					
			☐ For an anti-TSLP (e.g., TEZSPIRE	):	
			☐ Has severe asthma		

IN	IITIAL REQUESTS (continued)
2.	For treatment of CHRONIC SPONTANEOUS (IDIOPATHIC) URTICARIA:  ☐ 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:
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):
R	ENEWAL REQUESTS
	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  I elukotriene modifier  I 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
	For treatment of HYPEREOSINOPHILIC SYNDROME (HES):  Experienced measurable improvement in disease activity Reduction in use of systemic glucocorticoids for the treatment of HES
	LEASE FAY COMPLETED FORM WITH DECLIDED CLINICAL DOCUMENTATION

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Date:

Prescriber signature: