

## **Prior Authorization Request** Form for Monoclonal Antibodies-Anti-IL, Anti-IgE, Anti-TSLP

## FAX this completed form to (844) 205-3386

	OR Mail requests to: PA i	Jepartment	5 KIV	er Park Place	East, Suite 210   Fresho, CA 95720		
I. PROVIDER INFORMATION			II. MEMBER INFORMATION				
Prescriber Name:			Member Name:				
Prescriber Specialty:			Identification #:				
Office Contact Name:			Group #:				
Group Name:			Date of Birth:				
Fax #:			Medication Allergies:				
Phone #:							
III. DRUG INFORMATION (One drug request per form)							
Drug name and strength:  Dosage Inte		erval (sig):		Qty. per Day:			
	D DOCUMENTION (Det e submitted with prior o				ntation demonstrating evidence for each		
Specify diagnosis & diagnosis code relevant to this request:  Dx/Dx Code:							
Requests for all non-preferred medications: Does to member have a history of trial and failure of or contraindication or intolerance to the preferred Mono Antibodies-Anti-IL, Anti-IgE, Anti-TSLP agents? Refer that https://papdl.com/preferred-drug-list for a list of preferred non-preferred medications in this class.			oclonal to	□ Yes	Medications Previously Taken (start and end date and dose):		
☐ If not prescribed by one of the following specialist, pulmonologist, allergist, immunologist, dermatologist, hematologist/oncologist, etc., please indicate a specialist consulted: ☐ If currently using a different Monoclonal Antibodies – Anti-IL, Anti-IgE, Anti-TSLP agent (Fasenra, Nucala, Xolair, Cinqair, Dupixent, Tezspire), will discontinue the other Monoclonal Antibodies – Anti-IL, Anti-IgE, Anti-TSLP agent prior to starting requested ☐ If requesting for daily quantity exceeding daily limit (Refer to <a href="https://www.dhs.pa.gov/providers/Pharmacy-Services/Pages/Quantity-Limits-and-Daily-Dose-Limits.aspx">https://www.dhs.pa.gov/providers/Pharmacy-Services/Pages/Quantity-Limits-and-Daily-Dose-Limits.aspx</a> ), please provide supporting information:							
<b>ASTHMA:</b> ☐ Member's ☐ Member's		sthma contro therapeutic	oller me	edications (ple f or intolerand	ease provide asthma severity):ee or contraindication to asthma controller		
Requested medication will be used with standard asthma controller medications (LABA, LAMA, ICS):							
an unavo	_ ,						
For Cinqair, member has absolute blood eosinophil count 400 cells/microliter or greater:  For Nucala or Fasenra, member has asthma with an eosinophilic phenotype with an absolute blood eosinophil count of a least 150 cells/microl:							
	WAL REQUESTS:						
☐ Documented measurement improvement in severity of asthma evidenced by:							

	Member will continue to use standard asthma controller medications (LABA, LAMA, ICS) (Treatment plan):
CHRO	NIC IDIOPATHIC URTICARIA:
	Documented history of urticarial for at least 6 weeks
	Select all that apply:
	Requires steroids to control urticarial symptoms:
	Documented history of therapeutic failure, contraindication or intolerance to H1 Antihistamine (medication, start
	date and end date):
CHRO	NIC IDIOPATHIC URTICARIA RENEWAL REQUESTS:
	by:
	Prescriber's rationale for continued use:
ESOIN	OPHILIC GRANULOMATOSIS WITH POLYANGIITIS (EGPA):
	For a diagnosis of eosinophilic granulomatosis with polyangiitis (EGPA), both of the following:
	□ Documented history of asthma
	□ Documented history of absolute blood eosinophil count 1000 cells/microL or greater OR blood eosinophil level
	greater than 10% of leukocytes:
	□ Documented history of at least one of the following:
	Histopathological evidence of one of the following:
	☐ Eosinophilic vasculitis
	Perivascular eosinophilic infiltration
	☐ Eosinophil-rich granulomatous inflammation
	☐ Neuropathy, mono or poly (monitor deficit or nerve conduction abnormality)
	Pulmonary infiltrates, non-fixed
	☐ Sino-nasal abnormality
	☐ Cardiomyopathy
	☐ Glomerulonephritis
	☐ Alveolar hemorrhage
	□ Palpable purpura
	Parpable purpura  Positive test for ANCA
Ц	
	Requires systemic glucocorticoids to maintain remission (medication, start date and end date):
_	Has a contraindication or an intolerance to systemic glucocorticoids:
ECOIN	rituximab or cyclophosphamide:
ESUIN	OPHILIC GRANULOMATOSIS WITH POLYANGIITIS (EGPA) RENEWAL REQUESTS:
ь	Documented measurable improvement in eosinophilic with polyangilits disease activity evidenced
	by:
	REOSINOPHILIC SYNDROME (HES):
	For a diagnosis of hypereosinophilic syndrome HES), all of the following:
Ц	
	Has documented FIP1L1-PDGFRA-negative HES with organ damage or dysfunction
	Has documented blood eosinophil count ≥1000 cells/microL
	One of the following:
	Requires or has required systemic glucocorticoids to control symptoms (medication, start date and end date):
	Has documented contraindication or intolerance of systemic glucocorticoids:
HYPEF	REOSINOPHILIC SYNDROME (HES) RENEWAL REQUESTS:
	One of the following:
	☐ Has documented measurable improvement in disease activity evidenced
	by:
	Has documented reduction in use of systemic glucocorticoids for this indication (current dose):
ОТНЕ	R DIAGNOSES:
	consensus treatment guidelines:

IV. ADDITIONAL RATIONALE FOR REQUEST / PERTINENT CLINICAL INFORMATION :							
Appropriate clinical information to support the request on the basis of medical necessity must be submitted.	Provider Signature:	Date:					

Envolve Pharmacy Solutions will respond via fax or phone within 24 hours.

Requests for prior authorization (PA) requests must include member name, ID#, and drug name. Please include lab reports with requests when appropriate (e.g., Culture and Sensitivity; Hemoglobin A1C; Serum Creatinine; CD4; Hematocrit; WBC, etc.)