

MEDICARE FORM

Fasenra® (benralizumab) Injectable Medication Precertification Request

Page 1 of 2

(All fields must be completed and legible for precertification review.)

For Illinois MMP: FAX: 1-855-320-8445 PHONE: 1-866-600-2139 For other lines of business: Please use other form

Note: Fasenra is non-preferred. The preferred products are Nucala

and Xolair.

Please indicate:	☐ Start of treatment: Sta☐ Continuation of therap						
Precertification Requested By:				Phone: Fax:			
A. PATIENT INFOR	MATION						
First Name:			Last Name:				
Address:			City:		State:	ZIP:	
Home Phone:		Work Phone:		Cell Phone:			
DOB:	Allergies:			E-mail:			
Current Weight:	lbs or	_ kgs Heigl	nt: inches	or cms			
B. INSURANCE INF	ORMATION						
Aetna Member ID #:		If yes, provide ID	Does patient have other coverage?				
Medicare: Tes	☐ No If yes, provide ID #		Medicaid: Yes	☐ No If yes, pro	vide ID #:		
C. PRESCRIBER IN	IFORMATION						
First Name:		Last Name:		(Check One	<i>϶):</i> ☐ M.D. ☐	D.O. 🗌 N.P. 🗌 P.A	
Address:			City:		State:	ZIP:	
Phone:	Fax:	St Lic #:	NPI #:	DEA #:	UF	PIN:	
Provider E-mail:		Office Contact N	ame:		Phone:		
Specialty (Check of	ne):	Allergist					
D. DISPENSING PROVIDER/ADMINISTRATION INFORMATION Place of Administration: Self-administered Physician's Office Outpatient Infusion Center Phone: Center Name: Home Infusion Center Phone: Agency Name: Administration code(s) (CPT): Address:			☐ Physician ☐ Specialty Name: ☐ Phone: ☐ Address:	Dispensing Provider/Pharmacy: Patient Selected choice Physician's Office Retail Pharmacy Specialty Pharmacy Other: Name: Phone: Address: TIN: PIN:			
E. PRODUCT INFO			F				
<u> </u>	senra (benralizumab) Dose DRMATION – Please indicate			aliaahla			
					ode:		
Primary ICD Code: Secondary ICD Code: G. CLINICAL INFORMATION – Required clinical information must be completed.							
For All Requests (cl Note: Fasenra is no Yes No Has Yes No Has Please explain if the diagnosis? (select all	Inical documentation requirements on preferred. The preferred parties the patient had prior therapy the patient had a trial and fail Nucala (mepolizumab)	ed): products are Nucala, and > with Fasenra within the last ure, intolerance, or contraine Xolair (omalizumab) on(s) that the patient cannot	(olair. 365 days? dication to any of the fol	llowing? (select all tha	at apply)	or the patient's	

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Patient First Name Patient Last Name Patient Phone Patient DOB G. CLINICAL INFORMATION (continued) - Required clinical information must be completed in its entirety for all precertification requests. ☐ Yes ☐ No Is this infusion request in an outpatient hospital setting? → 🔲 Yes 🔲 No Has the patient experienced an adverse event with the requested product that has not responded to conventional interventions (e.g., acetaminophen, steroids, diphenhydramine, fluids, other pre-medications or slowing of infusion rate) or a severe adverse event (anaphylaxis, anaphylactoid reactions, myocardial infarction, thromboembolism, or seizures) during or immediately after an infusion? Yes No Does the patient have significant behavioral issues and/or physical or cognitive impairment that would impact the safety of the infusion therapy AND the patient does not have access to a caregiver? → Please provide a description of the behavioral issue or impairment: _ Yes No Is the patient medically unstable which may include respiratory, cardiovascular, or renal conditions that may limit the member's ability to tolerate a large volume or load or predispose the member to a severe adverse event that cannot be managed in an alternate setting without appropriate medical personnel and equipment? → Please provide a description of the condition: ☐ Cardiovascular: _ Respiratory: Renal: Other: ☐ Yes ☐ No Is the medication prescribed by or in consultation with an allergist, immunologist, or pulmonologist? ☐ Yes ☐ No Does the patient have a documented diagnosis of asthma? ☐ Yes ☐ No Will the patient continue to use maintenance asthma treatments (i.e., inhaled corticosteroids, additional controller) in combination with the requested medication? Yes No Will the patient receive the requested medication concomitantly with other biologics indicated for asthma (e.g., Cinqair, Dupixent, Nucala, Tezspire, Xolair)? For Initiation Requests (clinical documentation required): Please indicate the patient's baseline (e.g., before significant oral steroid use) blood eosinophil count in cells per microliter: ☐ Yes ☐ No Does the patient have uncontrolled asthma as demonstrated by experiencing two or more asthma exacerbations requiring oral or injectable corticosteroid treatment within the past year? > 🗌 Yes 🔲 No Does the patient have uncontrolled asthma as demonstrated by experiencing one or more asthma exacerbations resulting in hospitalization or emergency medical care visit within the past year? ⇒ ☐ Yes ☐ No Does the patient have uncontrolled asthma as demonstrated by experiencing poor symptom control (frequent). symptoms or reliever use, activity limited by asthma, night walking due to asthma) within the past year? ☐ Yes ☐ No Does the patient have inadequate asthma control despite current treatment with an inhaled corticosteroid and additional controller (long acting beta2-agonist, long-acting muscarinic antagonist, leukotriene modifier, or sustained release theophylline) at optimized doses? ☐ Yes ☐ No Is the patient dependent on systemic corticosteroids? For Continuation Requests (clinical documentation required): ☐ Yes ☐ No Is this continuation request a result of the patient receiving samples or a manufacturer's patient assistance program? Yes No Has asthma control improved on the requested medication treatment as demonstrated by a reduction in the frequency and/or severity of symptoms and exacerbations? > 🗌 Yes 🔲 No Has asthma control improved on the requested medication treatment as demonstrated by a reduction in the daily maintenance of oral corticosteroid dose? H. ACKNOWLEDGEMENT Request Completed By (Signature Required): Any person who knowingly files a request for authorization of coverage of a medical procedure or service with the intent to injure, defraud or deceive any

insurance company by providing materially false information or conceals material information for the purpose of misleading, commits a fraudulent

insurance act, which is a crime and subjects such person to criminal and civil penalties.

The plan may request additional information or clarification, if needed, to evaluate requests.