

Fasenra® (benralizumab) Injectable Medication Precertification Request

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

Aetna Precertification Notification Phone: <u>1-866-752-7021</u> (TTY: <u>711</u>)

FAX: <u>1-888-267-3277</u>

For Medicare Advantage Part B: Please Use Medicare Request Form

	_l Start of treatment: Start dat		1 1		
	Continuation of therapy: Da	ate or last treatment			Fow
A. PATIENT INFORMA	uested By:		Phone		_Fax:
First Name:	Amon	La	ast Name:		
Address:		Ci	ty:	State	e: ZIP:
Home Phone:	W	/ork Phone:	,	Cell Phone:	<u> </u>
DOB:	Allergies:			E-mail:	
Current Weight:	kgs	Height:	inches or _	cms	
B. INSURANCE INFO	RMATION				
Aetna Member ID #:			Does patient have other coverage?		
	No If yes, provide ID #:		edicaid: Yes	No If ves. provide	 ID #:
C. PRESCRIBER INFO				, , , , , , , , , , , , , , , , , ,	
First Name:		Last Name:		(Check One):] M.D. □ D.O. □ N.P. □ P.A
Address:		1	City:	State	ziP:
Phone:	Fax:	St Lic #:	NPI#:	DEA #:	UPIN:
Provider E-mail:	<u>.</u>	Office Contact Name:	:		Phone:
Specialty (Check one	e): Dulmonologist DAlle	rgist Other:		•	
D. DISPENSING PRO	VIDER/ADMINISTRATION INFO	RMATION			
□ Self-administered □ Physician's Office □ Outpatient Infusion Center Phone:			Phone:	rmacy	ail Pharmacy er: Fax:
Address:			_ TIN:		PIN:
E. PRODUCT INFORM					
Request is for: Fase	nra (benralizumab) Dose:		_ Frequency:		
F. DIAGNOSIS INFOR	RMATION – Please indicate prima	ary ICD Code and specify ar	ny other where applicab	le.	
Primary ICD Code: _	Sec.	condary ICD Code:		_ Other ICD Code:	
	IATION – Required clinical inform	nation must be completed in	its entirety for all prece	rtification requests.	
Yes No Is this	severe adverse event immediately after an les No Does the patient have infusion therapy AND	ienced an adverse event wi etaminophen, steroids, diph t (anaphylaxis, anaphylactoi infusion?	nenhydramine, fluids, or id reactions, myocardial es and/or physical or co access to a caregiver? sue or impairment:	other pre-medication infarction, thromboer gnitive impairment tha	s) or a nbolism, or seizures) during or

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Patient First Name	Patient Last Name	Patient Phone	Patient DOB			
G. CLINICAL INFORMATION (continu		on must be completed in its <u>entir</u>	ety for all precertification requests.			
Yes No Does the patient have a d	•					
		biologic (e.g., Adbry, Humira, Dup	oixent), or targeted synthetic drug (e.g., Rinvoq,			
Olumiant, Otezla, Xeljanz) for the same indication?					
☐ Yes ☐ No Is the medication prescrib	s the medication prescribed by or in consultation with an allergist, immunologist, or pulmonologist?					
Yes No Will the patient continue to requested medication?	Will the patient continue to use maintenance asthma treatments (i.e., inhaled corticosteroids, additional controller) in combination with the requested medication?					
For Initiation Requests (clinical docume	ntation required):					
Please indicate the patient's baseline (e.g.	., before significant oral steroid use)	blood eosinophil count in cells pe	r microliter:			
Yes No Has the patient previously						
Yes No Does the patient have uncontrolled asthma as demonstrated by experiencing two or more asthma exacerbations requiring oral o						
	corticosteroid treatment within the p		, averaging and as mare authors average stions			
—→ ☐ Yes ☐		emergency medical care visit with	/ experiencing one or more asthma exacerbations			
	•	0 ,	demonstrated by experiencing poor symptom			
			ctivity limited by asthma, night walking due to			
	· · ·	hin the past year?	, , , , ,			
☐ Yes ☐ No Prior to re	questing the requested medication,	did the patient have inadequate as	sthma control despite current treatment with an			
			ong-acting muscarinic antagonist, leukotriene			
	or sustained release theophylline) at					
☐ Yes ☐ No Is the pati	ent dependent on systemic corticost	eroids?				
For Continuation Requests (clinical doc	umentation required):					
☐ Yes ☐ No Is this continuation reques	st a result of the patient receiving sa	mples or a manufacturer's patient	assistance program?			
Yes No Has asthma control impro symptoms and exacerbati		eatment as demonstrated by a red	uction in the frequency and/or severity of			
	ma control improved on the requestorticosteroid dose?	ed medication treatment as demor	nstrated by a reduction in the daily maintenance			
H. ACKNOWLEDGEMENT						
Request Completed By (Signature Re	equired):		Date: /			
	rially false information or concea	Is material information for the	with the intent to injure, defraud or deceive any purpose of misleading, commits a fraudulent			

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