

Fax completed form to: (855) 840-1678 If this is an URGENT request, please call (800) 882-4462 (800.88.CIGNA)

## Fasenra (benralizumab)

PHYSICIA	PATIENT INFORMATION							
* Physician Name:			*Due to privacy regulations we will not be able to respond via fax with the outcome of our review unless all asterisked (*) items on this form are completed.*					
Specialty: * DEA, NPI or TIN:								
Office Contact Person:			* Patient Name:					
Office Phone:			* Cigna ID: * Date of Birth:					
Office Fax:			* Patient Street Address:					
Office Street Address:			City:	Sta	te: Zip:			
City:	State:	Zip:	Patient Phone:					
Urgency:         Standard         Urgent (In checking this box, I attest to the fact that applying the standard review time frame may seriously jeopardize the customer's life, health, or ability to regain maximum function)								
Medication Requested: Fasenra 10 mg/0.5 mL s Fasenra 30mg/ml syring Fasenra 30mg/ml Pen Other (please specify):								
Directions for use:	Directions for use:		Dose: Quantity:					
Duration of therapy:			ICD10:					
Where will this medication be obtained?  Accredo Specialty Pharmacy** Hospital Outpatient Retail pharmacy Other (please specify):			<ul> <li>Home Health / Home Infusion vendor</li> <li>Physician's office stock (billing on a medical claim form)</li> <li>**Cigna's nationally preferred specialty pharmacy</li> </ul>					
**Medication orders can be placed with Accredo via E-prescribe - Accredo (1620 Century Center Pkwy, Memphis, TN 38134-8822   NCPDP 4436920), Fax 888.302.1028, or Verbal 866.759.1557								
Facility and/or doctor dispensing and administering nFacility Name:State:Address (City, State, Zip Code):			nedication: Tax ID#:					
Where will this drug be administered? Patient's Home Hospital Outpatient			<ul><li>Physician's Office</li><li>Other (please specify):</li></ul>					
<b>NOTE:</b> Per some C	igna plans, infu	sion of medication N	IUST occur in the leas	t intensiv	ve, medically app	propriate setting.		
Is this patient a candidate for assistance of a Specialty Ca		•			e, physician's off ide medical nece			
Is the requested medication the patient?	for a chronic or	long-term condition	for which the prescrip	otion med	lication may be r	necessary for the life of ☐ Yes ☐ No		
What is your patient's d	liagnosis?							
<ul> <li>☐ Asthma</li> <li>☐ Chronic Obstructive Puln</li> <li>☐ Hypereosinophilic Syndra</li> <li>☐ other (please specify):</li> </ul>		(COPD)						

Clinical Information:		
Is this initial therapy, is the patient restarting therapy, or is the patient currently receiving Fasenra for at I Initial therapy Currently receiving Fasenra for at least 6 months Restarting therapy with Fasenra None of the above	east 6 mo	onths?
**(if Currently receiving Fasenra and for asthma) Has the patient responded to therapy? Note: Examples of a respon therapy are decreased asthma exacerbations; decreased asthma symptoms; decreased hospitalizations, emergence urgent care, or medical clinic visits due to asthma; and decreased requirement for oral corticosteroid therapy.		
(if no) Please provide support for continued use.		
(if Currently receiving Fasenra and for asthma) Has the patient continued to receive therapy with one inhaled cortico inhaled corticosteroid-containing combination?	osteroid Ol Ves	
(if asthma) Is your patient currently being treated with another monoclonal antibody therapy (for example, Adbry, Cir Nucala, Tezspire or Xolair)?	nqair, Dupi Yes	xent, □ No
(if yes or unknown) Please provide the rationale for concurrent use.	🗌 Yes	🗌 No
(if initial, if 12 yo or older for asthma) Does the patient have a forced expiratory volume in 1 second (FEV1) less that that is NOT due to smoking-related chronic obstructive pulmonary disease?	n 80% pred Ves	dicted
(if yes) Does the patient have an FEV1/forced vital capacity (FVC) less than 0.80?	🗌 Yes	🗌 No
(if no) Does the patient have an increase of over 12% AND greater than 200ml in FEV1 following standard dose of a short-acting bronchodilator? Note: The above lung function criteria may be me to or during asthma treatment.		
(if no) Does the patient have an increase of over 12% AND greater than 200ml in FEV1 I visits? Note: The above lung function criteria may be met at any time prior to or during as		tment.
(if no) Does the patient have an increase of over 12% AND greater than 200ml baseline to after at least 4 weeks of asthma treatment? Note: The above lung fu be met at any time prior to or during asthma treatment.	in FEV1 fr	om
(if no) Did the patient have a positive exercise challenge test? Note: The function criteria may be met at any time prior to or during asthma treat		•
(if no) Did the patient have a positive bronchial challenge test lung function criteria may be met at any time prior to or during treatment.	? Note: Th	
(if initial, if less than 12 yo and for asthma) Does the patient have a forced expiratory volume in 1 second (FEV1) les predicted that is NOT due to smoking-related chronic obstructive pulmonary disease?	s than 80⁰ ☐ Yes	%
(if yes) Does the patient have an FEV1/forced vital capacity (FVC) less than 0.80?	🗌 Yes	🗌 No
(if no) Does the patient have an increase of over 12% in FEV1 following administration of a standa short-acting bronchodilator? Note: The above lung function criteria may be met at any time prior to treatment.	o or during	
(if no) Does the patient have an increase of over 12% in FEV1 between prescriber visits? lung function criteria may be met at any time prior to or during asthma treatment.	? Note: The Yes	
(if no) Does the patient have an increase of over 12% in FEV1 from baseline to weeks of asthma treatment? Note: The above lung function criteria may be met to or during asthma treatment.		e prior
(if no) Did the patient have a positive exercise challenge test? Note: Th function criteria may be met at any time prior to or during asthma treatr		
(if no) Did the patient have a positive bronchial challenge test lung function criteria may be met at any time prior to or during treatment.	? Note: Th asthma	

(if initial and for asthma) Does your patient have a blood eosinophil level at least 150 cells per microliter within the previous 6 weeks or within 6 weeks prior to treatment with Fasenra or another monoclonal antibody therapy that may lower blood eosinophil levels? Note: Examples of monoclonal antibody therapies that may lower blood eosinophil levels include Fasenra, Adbry (tralokinumab-ldrm subcutaneous injection), Cinqair (reslizumab intravenous infusion), Dupixent (dupilumab subcutaneous injection), Nucala (mepolizumab subcutaneous injection), Tezspire (tezepelumab-ekko subcutaneous injection), and Xolair (omalizumab subcutaneous injection).
(if initial and for asthma) Has the patient received at least 3 consecutive months of combination therapy with BOTH: A. An inhaled corticosteroid (medium- or high- dose); AND B. At least one additional asthma controller or asthma maintenance medication? Note: Examples of additional asthma controller or asthma maintenance medications are inhaled long-acting beta2-agonists, inhaled long-acting muscarinic antagonists, and monoclonal antibody therapies (for example, Cinqair, Dupixent, Fasenra, Nucala, Tezspire, Xolair). Use of a combination inhaler containing both an inhaled corticosteroid (medium- or high- dose) and additional asthma controller/maintenance medication(s) would fulfill the requirement for both criteria A and B.
(if initial and for asthma) Does the patient have asthma that is uncontrolled or was uncontrolled at baseline as defined by the patient experiencing two or more asthma exacerbations requiring treatment with systemic corticosteroids in the previous year? Note: "Baseline" is defined as prior to receiving Fasenra or another monoclonal antibody therapy for asthma. Examples of monoclonal antibody therapies for asthma include Cinqair, Dupixent, Fasenra, Nucala, Tezspire, and Xolair.
(if no) Does the patient have asthma that is uncontrolled or was uncontrolled at baseline as defined by the patient experiencing one or more asthma exacerbation(s) requiring hospitalization, an Emergency Department visit, or an urgent care visit in the previous year? Note: "Baseline" is defined as prior to receiving Fasenra or another monoclonal antibody therapy for asthma. Examples of monoclonal antibody therapies for asthma include Cinqair, Dupixent, Fasenra, Nucala, Tezspire, and Xolair.
(if no) Does the patient have asthma that is uncontrolled or was uncontrolled at baseline as defined by asthma that worsens upon tapering of oral (systemic) corticosteroid therapy? Note: "Baseline" is defined as prior to receiving Fasenra or another monoclonal antibody therapy for asthma. Examples of monoclonal antibody therapies for asthma include Cinqair, Dupixent, Fasenra, Nucala, Tezspire, and Xolair.
(if initial and asthma) Is this medication being prescribed by, or in consultation with, an allergist, immunologist, or pulmonologist? ☐ Yes ☐ No
Additional pertinent information: Please provide any additional pertinent clinical information, including: if the patient is currently on the requested drug (with dates of use) and how they have been receiving it (for example: samples, out of pocket).
Attestation: I attest the information provided is true and accurate to the best of my knowledge. I understand that the Health Plan or insurer its designees may perform a routine audit and request the medical information necessary to verify the accuracy of the information reported on this form.
Prescriber Signature: Date:
Save Time! Submit Online at: <u>www.covermymeds.com/main/prior-authorization-forms/cigna/</u> or via SureScripts in your EHR.
Our standard response time for prescription drug coverage requests is 5 business days. If your request is urgent, it is important that you call us to expedite the request. View our Prescription Drug List and Coverage Policies online at cigna.com.
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