





M	ember N	Name: Date of Bi	rth:	Member ID#:	
Drug Information					
Physician billing (HCPCS code:) Pharmacy billing* (NDC:) *If Nucala [®] vial for injection is being used and billed by a pharmacy, the medication should be shipped to the health care facility where it will be administered.					
	ose:	Regimen:			
Billing Provider Information					
			Provider Name:		
Provider Phone: Provide					
If Nucala [®] vial for injection will be used, please provide the name of outpatient health care facility where Nucala [®] will be delivered to and administered at:					
		Prescribe	r Information		
			Prescriber Name:		
Sp	pecialty:	: Prescriber Phone:		Prescriber Fax:	
		Clinical	Information		
Page 1 of 3 - Please complete and return <u>all</u> pages. <i>Failure to complete all pages will result in processing delays.</i>					
For Initial Authorization (Initial approval will be for the duration of 6 months):					
1.		u cala[®] vial for injection: Will Nucala [®] vial for injection be administered in a	health care setti	ng hy a health care professional	
	A. Will Nucala [®] vial for injection be administered in a health care setting by a health care professional prepared to manage anaphylaxis? Yes No				
 For Nucala[®] prefilled autoinjector or prefilled syringe: A. Has the member or caregiver been trained by a health care professional on subcutaneous 			sional on subcutaneous		
	administration of Nucala $^{\textcircled{B}}$ prefilled autoinjector or prefilled syringe, monitoring for any				
3	allergic reactions, and storage of Nucala [®] prefilled autoinjector or prefilled syringe? Yes No Please indicate diagnosis and information:				
	Severe Eosinophilic Phenotype Asthma				
	Α.		this medication be used as add-on maintenance treatment for severe eosinophilic		
	phenotype asthma? Yes No i. If yes, please indicate member's daily medications and dose prescribed for treatment of this diagn			se prescribed for treatment of this diagnosis:	
	Drug/Dose: Drug/Dose:				
		Baseline blood eosinophil count: Da		······································	
C. Does member require daily systemic corticosteroids despite compliant use of a medium inhaled corticosteroid (ICS) plus at least 1 additional controller medication? Yes No					
		i. If no, please list number and dates of exa		ing systemic corticosteroids within last 12	
	D.	months: Number: Dates of exa Has the member been evaluated by an allergist, j		pulmonary specialist within the last 12 months	
(or an advanced care practitioner with a supervising physician who is an allergist,					
		pulmonologist, or pulmonary specialist)? Yes If yes, please include name of specialist:			
	E.	Please check all that apply:			
		Member has failed a medium-to-high-dose Dose:	ICS used compli	antly for at least the past 12 months Drug/	
	Member has failed at least 1 other asthma controller medication used in addition to the medium-to-high-				
dose ICS compliantly for at least the past 3 months - Drug/Dose:					
Eosinophilic Granulomatosis with Polyangiitis (EGPA)					
A. Does member have a past history of at least 1 confirmed EGPA relapse [requiring increase in oral					
corticosteroid (OCS) dose, initiation/increased dose of immunosuppressive therapy, or hospitalization] within the past 12 months? YesNo					
Page 1 of 3					
	CONFIDENTIALITY NOTICE				
F		eted prior authorization request form to 888-601-8461 o lectronic Prior Authorization throughCoverMyMeds® or		This document, including any attachments, contains information which is confidential or privileged. If you are not the intended recipient, be aware	
S	ureScripts	s. All requested data must be provided. Incomplete form			
		is without the chart notes will be returned. Pharmacy Coverage Guidelines are available at AetnaBetterHealth com/Oklahoma	please notify the	assisted as a service of the second the seco	



State of Oklahoma SoonerCare

Health Care Authority Nucala[®] (Mepolizumab) Prior Authorization Form

Member Name:

Date of Birth:

Member ID#:

Clinical Information

Page 2 of 3—Please complete and return all pages. Failure to complete all pages will result in processing delays.

3 Please indicate diagnosis and information, continued:

- Eosinophilic Granulomatosis with Polyangiitis (EGPA), continued
 - B. Does member have refractory disease within the last 6 months following induction of standard treatment regimen administered compliantly for at least 3 months? Yes____ No_
 - C. Is diagnosis granulomatosis with polyangiitis (GPA) or microscopic polyangiitis (MPA)? Yes No
 - D. Has member failed to achieve remission despite glucocorticoid therapy (oral prednisone equivalent equal to or greater than 7.5mg/day) for a minimum of 4 weeks duration? Yes____ No_
 - E. Has the member been evaluated by an allergist, pulmonologist, pulmonary specialist, or rheumatologist (or an advanced care practitioner with a supervising physician who is an allergist, pulmonologist, pulmonary specialist, or rheumatologist) within the past 12 months? Yes____ No____ i. If yes, please include name of specialist:

Hypereosinophilic Syndrome (HES)

- A. Has member been diagnosed with HES for ≥6 months without an identifiable non-hematologic secondary cause? Yes No
- B. Does member have a history of at least 2 confirmed HES flares [requiring increase in oral corticosteroid (OCS) dose, initiation/increased dose of cytotoxic or immunosuppressive therapy, or hospitalization] within the past 12 months? Yes No Flare dates:
- C. Please provide member's baseline blood eosinophil count: _____ Date taken: _____
- D. Is HES FIP1L1-PDGFRα kinase-positive? Yes No
- E. Has member failed to achieve remission despite corticosteroid therapy (oral prednisone equivalent ≥10mg/day) for a minimum of 4 weeks duration? Yes No
 - i. If no, is member is unable to tolerate corticosteroid therapy due to significant side effects from glucocorticoid therapy? Yes____ No_
- F. Is the prescriber a hematologist or a specialist with expertise in treatment of HES (or an advanced care practitioner with a supervising physician who is a hematologist or a specialist with expertise in treatment of HES)? Yes No

Chronic Rhinosinusitis with Nasal Polyposis (CRSwNP)

- A. Will Nucala[®] be used as add-on maintenance treatment for inadequately controlled CRSwNP? Yes____ No
- B. Does the member have a trial with intranasal corticosteroid that resulted in failure (or have a contraindication or documented intolerance)? Yes _No
 - i. If yes, please provide the medication used and dates of use:
- C. Has the member required prior sino-nasal surgery? Yes____ No____ D. Has the member been treated with systemic corticosteroids for CRSwNP in the past 2 years (or have a contraindication or documented intolerance)? Yes No
- E. Has the member been evaluated by an otolaryngologist, allergist, immunologist, or pulmonologist (or an advanced care practitioner with a supervising physician who is an allergist, otolaryngologist, allergist, immunologist, or pulmonologist) within the past 12 months? Yes No____
 - i. If yes, please include name of specialist:
- Does the member have symptoms of chronic rhinosinusitis (e.g., facial pain/pressure, reduction or loss of smell, F. nasal blockade/obstruction/congestion, nasal discharge) for 12 weeks or longer despite attempts at medical management? Yes No
- G. Does the member have evidence of nasal polyposis by direct examination, sinus CT scan, or endoscopy? Yes No
- H. Will the member continue to receive intranasal corticosteroid therapy? Yes No
 - i. If yes, does the member have a contraindication to intranasal corticosteroid therapy? Yes No 1. If yes, please provide the member's contraindication:
- Will Nucala[®] be used concurrently with other biologic medications? Yes No Ι.
 - i. If yes, please provide patient-specific information to support the concurrent use of Nucala® with other biologic medications:

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Fax completed prior authorization request form to 888-601-8461 or submit Electronic Prior Authorization throughCoverMyMeds® or SureScripts. All requested data must be provided. Incomplete forms or forms without the chart notes will be returned. Pharmacy Coverage Guidelines are available at AetnaBetterHealth.com/ Oklahoma.

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State of Oklahoma SoonerCare Nucala[®] (Mepolizumab) Prior Authorization Form

Member Name:

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Page 3 of 3—Please complete and return all pages. Failure to complete all pages will result in processing delays.

For Continued Authorization:

- 1. Is the member compliant with therapy? Yes No
- 2. Is the member responding well to therapy? Yes_ No
- 3. If member's diagnosis includes EGPA, please check all that apply:
 - Member has a Birmingham Vasculitis Activity Score (BVAS) of zero
 - Member has fewer EGPA relapses from baseline
 - Member has had a decrease in daily OCS dose regimen from baseline
 - If none of the above, please provide additional information on member's response to therapy:
- 3. If member's diagnosis includes **HES**, please provide the following:
 - A. Is the member responding to Nucala[®] therapy? Yes_ No
 - i. If yes, has member had fewer HES flares from baseline? Yes No
 - a. Please provide number of HES flares: Baseline: Current: ii. If yes, has member had a decrease in daily OCS dosing from baseline? Yes
 - a. Please provide daily OCS dosing: Baseline: Current:

Compliance with all of the prior authorization criteria is a condition for payment for this drug by SoonerCare. All information must be provided and SoonerCare may verify through further requested documentation. The member's drug history will be reviewed prior to approval.

Prescriber Signature:

(By signature, the physician confirms the criteria information above is accurate and verifiable in patient records.)

Pharmacist Signature:

Please do not send in chart notes. Specific information will be requested if necessary. Failure to complete all pages will result in processing delays.

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