

State of Oklahoma

WICHIBC	er Name:	Date of Birth:	Member ID#:				
		Drug Information					
Pharm	acy billing (NDC:) Fill Date:					
Dose:_		Regimen:					
Billing Provider Information							
Pharmacy NPI:		Pharmacy Name:					
Pharmacy Phone:		Pharmacy Fax:					
Prescriber Information							
Prescri	iber NPI:	Prescriber Name:					
Prescri	iber Phone:	Prescriber Fax:	Specialty:				
Clinical Information							
Prescriber Phone: Prescriber Fax: Specialty:							
	dose ICS comp	iled at least 1 other asthma controller medication obliantly for at least the past 3 months.	_				

Page 1 of 3

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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Pharm - 115 12/22/2023



State of Oklahoma SoonerCare

Date of Birth:



Member ID#:

Health Care Authority Dupixent® (Dupilumab) Prior Authorization Form

Me	ember Name:	Date of Birth:	Member ID#:			
		Clinical Information				
Pa	age 2 of 3—Please complete and retur	n <u>all</u> pages. <i>Failure to comple</i>	te all pages will result in processing delays.			
3.	duration of 16 weeks): A. Is member inadequately controlled. B. Has the member failed 1 medium. YesNo i. If yes, please provide the ma. Drug: b. Was the trial at least 2 ii. If no, is there a contraindic Tier-1 topical corticosteroid a. If yes, please described. C. Has the member failed 1 topical of YesNo i. If yes, please provide the ma. Drug: b. Was the trial at least 2	ed with topical prescription the n potency to very-high potency medication and duration of treation or documented intolerant ds? Yes No e: calcineurin inhibitor [e.g., Elide medication and duration of treation or documented intolerant decident and duration of treation or documented intolerant decident and duration?	Tier-1 topical corticosteroid? atment: Date of trial: No ce to medium potency to very-high potency el® (pimecrolimus), Protopic® (tacrolimus)]?			
4.	If diagnosis is Chronic Rhinosinusit approvals will be for the duration of 6. A. Will Dupixent® be used as add-oryesNo B. Does the member have a trial withor documented intolerance)? Yes i. If yes, please provide the r. C. Has the member required prior since the member been treated with contraindication or documented in the	is with Nasal Polyposis (CR: months): In maintenance treatment for in the intranasal corticosteroid that is No nedication used and dates of ino-nasal surgery? Yes No_ ith systemic corticosteroids for intolerance)? Yes No_ ns of chronic rhinosinusitis (e.g. n/congestion, nasal discharge) _ No_ e of nasal polyposis by direct eleive intranasal corticosteroid the	t resulted in failure (or have a contraindication use: O CRSwNP in the past 2 years (or have a I., facial pain/pressure, reduction or loss of for 12 weeks or longer despite attempts at xamination, sinus CT scan, or endoscopy? Iderapy? Yes No Basal corticosteroid therapy? Yes No			
5.	If diagnosis is Eosinophilic Esophag duration of 6 months): A. Does the member have 2 or more B. Does the member have ≥ 15 intra	re episodes of dysphagia per w	following (Initial approvals will be for the reek? Yes Non-power field (eol/hpf)? Yes No			
	(continued on next page)					

Page 2 of 3

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State of Oklahoma SoonerCare





Health Care Authority Dupixent® (Dupilumab) Prior Authorization Form

Member Name:	Date of Birth:	_ Member ID#:				
	Clinical Information					
*Page 3 of 3—Please complete and return <u>all</u> pages. <i>Failure to complete all pages will result in processing delays.</i> * C. Has the member failed 1 high-dose proton pump inhibitor?						
	medication and duration of treatment:					
a. Drug:	Dat 8 weeks in duration? Yes No	te of trial:				
ii. If no, is there a contraindid Yes No	cation or documented intolerance to h	igh-dose proton pump inhibitors?				
a. If yes, please describ	e: wed inhaled respiratory corticosteroid	(a.g. hudaaanida\2				
Yes No	medication and duration of treatment:					
a. Drug:		te of trial:				
	8 weeks in duration? Yes No					
the state of the s	cation or documented intolerance to s	wallowed inhaled respiratory				
	No					
 a. If yes, please describ 6. If diagnosis is Prurigo Nodularis (Pl months): 	e:	l approvals will be for the duration of 6				
	s of PN for at least 3 months? Yes	No				
B. Does the member have a Worst	-Itch Numeric Rating Scale (WI-NRS)					
C. Does the member have ≥ 20 PN	lesions? Yes No					
 D. Has the prescriber ruled out all of 						
Yes No	m potency to very-high potency Tier-1					
a. Drug:	medication and duration of treatment: Dat	te of trial:				
b. Was the trial at least i	2 weeks in duration? Yes No					
Tier-1 topical corticosteroi						
a. If yes, please describe F. Has the member failed 1 tonical	e: calcineurin inhibitor [e.g., Elidel [®] (pim	pecrolimus) Protonic® (tacrolimus)12				
Yes No						
i. If yes, please provide the a. Drug:	medication and duration of treatment: Dat	te of trial:				
b. Was the trial at least :	2 weeks in duration? Yes No					
ii. If no, is there a contraindic	cation or documented intolerance to to	opical calcineurin inhibitors?				
Yes No						
a. If yes, please describ	e:					
For Continued Authorization: 1. Is member compliant with therap	v2 Ves No					
 Is member compliant with therap Is member responding well to the 	erapy? Yes No					
Compliance with all of the prior authoriz information must be provided and Soon	cation criteria is a condition for paym erCare may verify through further re					
drug history will be reviewed prior to ap	•					
Prescriber Signature:	Dat					
By signature, the physician confirms the criteria information above is accurate and verifiable in patient records.						
Please do not send in chart notes. Specific information/documentation will be requested if necessary.						

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