

Dispense as written/Brand medically necessary

Referral Status:	MRN:	
New referral	Order change	Order Renewal
Patient preferred clinic:		

Substitution permitted

1111 001011			Pat	tient preferred clinic:					
	one: 1-800-809-1265 Fax: 1-866-872-89								
	ıcala® (mepolizumab) Standarı	d Plan of Trea	tm	ent for Asthm	a				
	TIENT DEMOGRAPHICS:								
	Date of Referral:			Patient's Phone:					
	ient Name:		Address:						
	e of Birth:		_	ty, State, Zip:	1	1 10		Lunica	
Hei	ght in inches: Weight: LB	or KG	G G	ender:	Allergies:	Se	e list	NDKA	
DIA	AGNOSIS: (PLEASE COMPLETE 2 <sup>ND</sup> AND	3 <sup>RD</sup> DIGITS TO CO	MP	LETE ICD 10 FOR B	ILLING )				
	J45.50 - Severe persistent asthma, uncomplicate				,				
	J45.52 - Severe persistent asthma with status as								
	J45.51 - Severe persistent asthma with (acute) e	exacerbation							
	Other:								
REC	QUESTED DOCUMENTATION:	PREVIOUS ADMIN	IISTI	RATION: HAS THIS PA	ATIENT TAKEN THIS	S MEDICATIO	N BEF	ORE?	
1	Insurance information	IF NO:	IF YES:						
2	Most recent History & Physical	PLEASE STATE		LAST INJECTION DATE:					
3	Full medication list	REQUIRED WASHOUT FROM PREVIOUS	NE	EXT INJECTION DATE:					
4	Tried and failed therapies	THERAPY:	IF	ORDER CHANGE:					
5	Blood eosinophil level (pre-treatment baseline count greater than or equal to 150 cells/mcL)		Continue current order until insurance approved						
		Provider Attestation	on fo	r HCP administration:					
	Provider attestation that the patient or caregiver are not competent or are physically unable to administer the Nucala product FDA labeled for self-			Patient has experienced severe hypersensitivity reactions (e.g., anaphylaxis, angioedema, bronchospasm, or hypotension) to Nucala within the past 6 months and					
	_administration	labeled for self-		•	and direct monitoring by		•		
	Patient has a history of uncontrolled disease and orderin their clinical opinion, it is not advisable to try the self-adn			<u></u>					
	requested drug	illilistered formulation of			<ul> <li>ordering provider attests</li> <li>f-administered formulation</li> </ul>			on, it is not	
	The location and circumstances for self-administration a potential treatment of anaphylaxis should that arise.	re not adequate for the		authouble to all and som			-9		
*Spe	cific reactions:								
ME	DICATION ORDERS:								
	E: Patient may be ineligible to receive Nucala® (mepoli	zumah) if natient has sig	ns/s	vmntoms of narasitic infe	ction is currently being	treated for a na	rasitic in	fection or is	
	ng acute bronchospasm and/or asthma attack.	zamas) ii patient nas sig	5113/3	ymptoms or parasitic line	ction, is currently being	treated for a pa	i asitic iii	rection, or is	
D0	SE/FREQUENCY:								
_	Nucala <sup>®</sup> (mepolizumab) 100 mg every	four (4) wooks vis		houtanoous inicatio	an.				
_	Indicata (mepolizumab) 100 mg every	iour (4) weeks via	Sui	ocularieous injectio	Of 1				
		ubcutaneous inject	tion	to the upper arm, t	high, or abdomen	-			
SPI	ECIAL ORDERS:								
Ext	tended post treatment monitoring: monitor					ter second i	njectio	on, and 15	
		minutes after eac	n sı	ubsequent injection					
			<b>\</b>	Refills x 12 month	ns unless noted oth	erwise here:			
AD	VERSE REACTION & ANAPHYLAXIS ORI	DERS:							
Adm	ninister acute infusion and anaphylaxis medication	s per Palmetto Infusio	n sta	anding adverse reaction	orders which can be	found at our	æ:	anassas a	
	site or scan here.	o per i annotto initiolo	11 010	anding daverse redelion	orders, willow can be	Tourid at our			
							ا، <u>س</u>	TEARES INSTITUTE	
PRI	ESCRIBER INFORMATION:								
	OVIDER NAME:			PHONE:					
ADDRESS: CITY, STATE, ZIP:				FAX:					
				NPI:		D 4 ===			
PKI	ESCRIBER SIGNATURE: (No stamp signa	tures)				DATE			