Monoclonal Antibodies: Tezspire



Beneficiary Information

1. Beneficiary Last Name:	2. First Name:	
3. Beneficiary ID #:	4. Beneficiary Date of Birth:	5. Beneficiary Gender:
Prescriber Information		
6. Prescribing Provider NPI #:		
7. Requester Contact Information - Name:	Phone	e #: Ext
Drug Information		
8. Drug Name:	9. Strength:	10. Quantity Per 30 Days:
11. Length of Therapy (in days): 🛛 up to 30	Days 🗌 60 Days 🗌 90 Days 🗌 120 D	Days 🗌 180 Days 🔲 365 Days 🗌 Other
Clinical Information		
Initial Approval:		
1. Is the beneficiary age 12 years of age or olde		
2. Does the beneficiary have a diagnosis of seven		
3. Does the beneficiary have at least 1 of the fo	ollowing? Yes No Please indicate which the set of th	ch one(s)
a. Symptoms throughout the day		
b. Nighttime awakenings, often 7x/week	overal times per dev	
 c. SABA use for symptom control occurring so d. Extremely limited normal activities 	everal times per day	
e. Lung function (percent predicted FEV1) < 6	50%	
f. Exacerbations requiring oral systemic corti		tense relative to moderate asthma
	ce treatment for a beneficiary who regular	rly received BOTH of the following? \Box Yes \Box No
b. An additional controller medication (e.g., I	ong-acting beta agonist, leukotriene modif	fiers
5. Has the beneficiary had, in the previous year	$r_{i} \ge 2$ exacerbations requiring oral or injecta	able corticosteroid treatment (in addition to the
regular maintenance therapy defined above) O	R one exacerbation resulting in a hospitali	ization? 🗆 Yes 🗆 No
6. Is there a baseline measurement of ≥ 1 of th	e following for assessment of clinical status	s? Yes No Please indicate which one(s).
a. Use of systemic corticosteroids	-	
b. Use of inhaled corticosteroids		
c. Number of hospitalizations, ER visits, or u	nscheduled visits to healthcare provider du	ue to condition
d. FEV1		
 Will the beneficiary use Tezspire for the relie Will the beneficiary use Tezspire in combinat 		
omalizumab, mepolizumab, reslizumab, dupilu	mab)? 🗆 Yes 🗆 No	
9. Does the beneficiary have hypersensitivity to	o tezepelumab-ekko (Tezspire) or any of its	s excipients? 🗆 Yes 🗆 No
10. Does the beneficiary have an active or untr	eated helminth infection? \Box Yes \Box No	
11. Will Tezspire be administered concurrently	with live vaccines? \Box Yes \Box No	
Initial approval can be for up to 6 months		
For continuation of therapy, please answer qu		
		ns, asthma exacerbations, or airway function as
evidenced by decrease in \geq 1 of the following?	\Box res \Box No Please indicate which one(s))
 a. Use of systemic corticosteroids b. Two-fold or greater decrease in inhaled co 	prticosteroid use for at least 2 days	
c. Hospitalizations	recoster of a crease of ago	
d. ER visits		
e. Unscheduled visits to healthcare provider		
f. Improvement from baseline in FEV1		

13. Has the beneficiary experienced any serious treatment-related adverse events (e.g., parasitic [helminth] infection, severe hypersensitivi reactions)?
Yes
No Reauthorizations can be for up to 6 months ** Please provide medical records documenting the beneficiary's current Asthma status and response to Tezspire treatment** Signature of Prescriber: _ Date:

(Prescriber Signature Mandatory)

I certify that the information provided is accurate and complete to the best of my knowledge, and I understand that any falsification, omission, or concealment of material fact may subject me to civil or criminal liability.