## **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	
<ul> <li>c. SABA use for symptom control occurring so d. Extremely limited normal activities</li> </ul>	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) <b>O</b>	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		
<ol> <li>Will the beneficiary use Tezspire for the relie</li> <li>Will the beneficiary use Tezspire in combinat</li> </ol>		
omalizumab, mepolizumab, reslizumab, dupilu	mab)? 🗆 <b>Yes 🗆 No</b>	
9. Does the beneficiary have hypersensitivity to	o tezepelumab-ekko (Tezspire) or any of its	s excipients? 🗆 <b>Yes</b> 🗆 <b>No</b>
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)	)
<ul> <li>a. Use of systemic corticosteroids</li> <li>b. Two-fold or greater decrease in inhaled co</li> </ul>	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.