tecentriq [®] (atezolizumab) Medication Precertification Request						Phone: 1-8	certification Notification 666-752-7021 (TTY: 711) 888-267-3277	
	Page 1 of (All fields n		ompleted and legible	for pred	certification review	.)		are Advantage Part B: Medicare Request Form
	Start of treatment: Star			-	1			
	Continuation of therap						F	
	uested By:				Phone:		Fax:	
A. PATIENT INFORMA	ATION			Last N	amai			
				City:	ane.		State:	ZIP:
Address: Home Phone: Work Ph						State.	ZIF.	
DOB:	Allergies:	Work Phone:				Email:		
		kao	Haight		inches or			
B. INSURANCE INFO	Ibs or	<u>kgs</u>	Height	·	inches or	Cm	S	
			Does patient have	othor o				
			If yes, provide ID#:		-			
			Insured:					
] No If yes, provide ID #:				aid: 🗌 Yes 🔲		ovide ID #·	
C. PRESCRIBER INFO				mouro		110 il 300, pi	ondo 18 #: _	
First Name:			Last Name:			(Check O	ne): 🗌 M.D	. 🗌 D.O. 🗌 N.P. 🗌 P.A.
Address:			I	Cit	y:	· ·	State:	ZIP:
Phone:	Fax:		St Lic #:	NF	'l #:	DEA #:		UPIN:
Provider Email:			Office Contact Nan	ne:			Phor	ie:
Specialty (Check one): 🗌 Oncologist 🔲	Hemato	ologist 🗌 Other:					
Place of Administrat Self-administered Outpatient Infusion Center Name Home Infusion Ce	Physician's Of n Center Phone: e:	fice				ffice armacy	☐ Retail Ph ☐ Other:	armacy
• •	e:							
Address:					TIN:		PIN:	· · · · · · · · · · · · · · · · · · ·
Administration cod					·		1 111.	
	ntrig (atezolizumab) Dos	e:			Frequency:			
	MATION – Please indicate		CD Code and specify			ole.		
Primary ICD Code:			dary ICD Code:			Other ICD	Code:	
	IATION – Required clinical i		,					
Yes No Has the (e.g., C) Alveolar Soft Part Please indicate the	e clinical setting in which the ill the requested medication the requested medication b ill the requested medication e clinical setting in which the	e progres a (pembro requeste be used eing used be used	ssion while on program lizumab), Bavencio (a ed medication will be as a single agent? d to treat small cell ne in combination with e	aveluma used: [euroend etoposid used: [b), or Imfinzi (durv] Unresectable di ocrine carcinoma e and either cispla	alumab))? sease	istatic diseas ECC)? in?	e 🗌 Other
☐ Other ☐ Yes	clinical setting: Unresec	be used i	in combination with b			ease 🗌 Diseas	se with exten	sive liver tumor burden
🗌 Yes 🗌 No 🛛 Wi	Il the requested medication	be used f	for initial treatment?					

Tecentriq[®] (atezolizumab) Medication Precertification Request

Page 2 of 3

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 Aetna Precertification Notification

 Phone:
 <u>1-866-752-7021</u> (TTY: <u>711</u>)

 FAX:
 <u>1-888-267-3277</u>

(All fields must be completed and legible for precertification review.)

For Medicare Advantage Part B: Please use Medicare Request Form

Patient First Name	Patient Last Name	Patient Phone	Patient DOB						
G. CLINICAL INFORMATION (continued) -	- Required clinical information must l	be completed in its entirety for all	precertification requests.						
G. CLINICAL INFORMATION (continued) – Required clinical information must be completed in its <u>entirety</u> for all precertification requests. Melanoma Please indicate the clinical setting in which the requested medication will be used: Unresectable disease Metastatic disease Other Yes No Unknown Is the tumor positive for BRAF V600 mutation? Yes No Will the requested medication be used in combination with cobimetinib (Cotellic) and vemurafenib (Zelboraf)?									
☐ Yes ☐ Yes ☐ Yes ☐ Yes ☐ No Will the requested drug be used in combination with obsidiation with obsidiation with obsidiation with obsidiation (2010011)?									
□ Non-small cell lung cancer (NSCLC)	□ Non-small cell lung cancer (NSCLC)								
What is the clinical setting in which the requested drug will be used? Recurrent disease Advanced disease Metastatic disease Other Yes No Unknown Is the tumor negative for EGFR exon 19 deletions, L858R mutations, and ALK rearrangements? Yes No Is testing for these genomic tumor aberrations not feasible due to insufficient tissue?									
☐ Yes ☐ No Will the requested medication be used as a single agent? What is the place in therapy in which the requested medication will be used? ☐ Initial treatment ☐ Subsequent treatment									
For tumor negative for EGFR exon 19 deletions, L858R mutations, and ALK rearrangements: What is the requested regimen? Single agent In combination with bevacizumab									
Please indicate the place in therapy: ☐ Continued maintenance therapy ☐ Subsequent therapy									
 Yes □ No □ Unknown Is the patient's tumor PD-L1 positive? Yes □ No Will the requested medication be used as a single agent? 									
 ☐ Small cell lung cancer (small cell carcinoma) ☐ Yes ☐ No Does the patient have extensive-stage disease? ☐ Yes ☐ No Will the requested medication be used in combination with etoposide and carboplatin (followed by single agent maintenance)? ☐ Yes ☐ No Will the requested medication be used for initial treatment? 									
For Continuation Requests (clinical docum	entation required for all requests	<u>:</u>							
 Yes □ No Has the patient experienced disease progression while receiving another programmed death receptor-1 (PD-1) or programmed death ligand 1 (PD-L1) inhibitor (e.g., Opdivo (nivolumab), Keytruda (pembrolizumab), Bavencio (avelumab), or Imfinzi (durvalumab))? Yes □ No Is there evidence of disease progression or unacceptable toxicity while on the current regimen? Yes □ No Is this infusion request in an outpatient hospital setting? 									
└────────────────────────────────────	ent continuing on a maintenance req nited to the following?		inistered combination chemotherapy including r non-small cell lung cancer (NSCLC)						
	er combination chemotherapy ase enter the regimen: Other:								
Yes No Is the pati pneumoni transverse	ent experiencing severe toxicity requ itis, Stevens-Johnson syndrome, act e myelitis, myocarditis, pericarditis, a	uiring continuous monitoring (e.g Ite pancreatitis, primary adrenal i	. Grade 2-4 bullous dermatitis, transaminitis, insufficiency aseptic meningitis, encephalitis, function, conduction abnormalities)?						
Yes No Has the p interventi severe ac immediat		diphenhydramine, fluids, other pr ctoid reactions, myocardial infarc	has not responded to conventional re-medications or slowing of infusion rate) or a ction, thromboembolism, or seizures) during or						
☐ Yes ☐ No Does the	· ·		ecial interventions only available in the						
	ease explain: patient have significant behavioral is	sues and/or physical or cognitive	e impairment that would impact the safety						
infusion th	herapy AND the patient does not have ease provide a description of the beh	e access to a caregiver?							

Tecentriq[®] (atezolizumab) Medication Precertification Request

Page 3 of 3

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For Medicare Advantage Part B: Please use Medicare Request Form

Patient First Name	Patient Last Name	Patient Phone	Patient DOB				
G. CLINICAL INFORMATION (continued) -	Required clinical information must be comr	leted in its entirety for all precertit	fication requests				
C. CEINICAE INI ONMATION (Continued) -		leted in its <u>entirety</u> for all precertin					
Yes No Is the patient medically unstable which may include respiratory, cardiovascular, or renal conditions that may limit the patient's ability to tolerate a large volume or load or predispose the patient to a severe adverse event that cannot be managed in an alternate setting without appropriate medical personnel and equipment?							
→ Please provide a description of the condition: □ Cardiovascular:							
Respiratory:							
☐ Renal:							
	Ε	Other:					
Yes No Is the patient within the initial 6 months of starting therapy?							
└────────────────────────────────────							
H. ACKNOWLEDGEMENT							
Request Completed By (Signature Requi	red):		Date: / /				
Any person who knowingly files a request for authorization of coverage of a medical procedure or service with the intent to injure, defraud or deceive any							

Any person who knowingly files a request for authorization of coverage of a medical procedure or service with the intent to injure, defraud or deceive any insurance company by providing materially false information or conceals material information for the purpose of misleading, commits a fraudulent insurance act, which is a crime and subjects such person to criminal and civil penalties.

The plan may request additional information or clarification, if needed, to evaluate requests.