

Bavencio® (avelumab) Injectable Medication Precertification Request

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

Aetna Precertification Notification

Phone: 1-866-752-7021 **FAX:** 1-888-267-3277

For Medicare Advantage Part B: Please Use Medicare Request Form

Please indicate:	☐ Start of treatment: Sta							
Precertification Re	equested By:		ie:	Fax:				
A. PATIENT INFOR								
First Name:			Last Name:					
Address:			City:		State:	ZIP:		
Home Phone:		Work Phone:	,	Cell Phone:	I.			
DOB:	Allergies:	111011111111111111111111111111111111111		Email:				
	Ibs or	kas Height	:inches	I				
B. INSURANCE INF		_ kgs		orome	,			
		Doos nationt have	other coverage?	□ Voc. □ No.				
	# :		Does patient have other coverage?					
Group #:Insured:			If yes, provide ID#: Carrier Name:Insured:			-		
Medicare: Yes	☐ No If yes, provide ID #:		Medicaid: Yes	☐ No If yes, pro	ovide ID #:	_		
C. PRESCRIBER IN								
First Name:		Last Name:		(Check Or	ne): 🔲 M.D. 🔲	D.O. 🗌 N.P. 🗌 P.A.		
Address:		<u> </u>	City:	•	State:	ZIP:		
Phone:	Fax:	St Lic #:	NPI #:	DEA #:	UP	IN:		
Provider Email:		Office Contact Na			Phone:			
Specialty (Check o	ne): Oncologist							
	OVIDER/ADMINISTRATION	<u>'</u>						
Center Na Home Infusion (Agency Na	ion Center Phone: me: Center Phone: mme: code(s) (CPT):		Name: Address: Phone:	Pharmacy [Fax:			
E. PRODUCT INFO	RMATION							
Request is for Bay	encio (avelumab): Dose: _		Frequency:					
F. DIAGNOSIS INFO	DRMATION – Please indicate	primary ICD Code and specif	y any other where app	olicable.				
Primary ICD Code:		Secondary ICD Code:		Other ICD (Code:			
G. CLINICAL INFORMATION – Required clinical information must be completed in its entirety for all precertification requests. For All Requests (Clinical documentation required for all requests): Yes No Has the patient experienced disease progression while receiving another PD-1 or PD-L1 inhibitor (e.g., Opdivo (nivolumab), Keytruda (pembrolizumab), Tecentriq (atezolizumab), Bavencio (avelumab), and Imfinzi (durvalumab).)? Bladder Urothelial Cancer Yes No Will the requested drug be used as a single agent? Maintenance therapy request only: Yes No Will the requested drug be used as maintenance therapy? Yes No Did the patient experience disease progression on first-line platinum-containing chemotherapy (e.g., cisplatin, carboplatin)? Subsequent therapy request only: Please indicate how the requested drug will be used: First line treatment Subsequent treatment Please select the clinical setting in which the requested drug will be used: Metastatic disease Locally advanced disease Post-cystectomy Please indicate the clinical setting in which the requested drug will be used following cystectomy: Metastatic disease Local recurrence Other								
☐ Preserved bladder → Please indicate the clinical setting in which the requested drug will be used in a preserved bladder: ☐ Muscle invasive local recurrent ☐ Muscle invasive persistent disease ☐ Other								



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Patient First Name	Patient Last Name	Patient Phone	Patient DOB					
G. CLINICAL INFORMATION (continued) - F	Required clinical information must be completed	l for ALL precertificati	on requests.					
☐ Stage II or IIIA disease								
Yes No Is tumor present following primary treatment?								
Other								
☐ Endometrial carcinoma								
Please select the clinical setting in which the r	requested drug will be used: Recurrent disease	Metastatic diseas	se 🗌 Other					
	Please indicate how the requested drug will be used: First line treatment Second-line treatment							
☐ Yes ☐ No ☐ Unknown Is the tumor mid	Yes No Unknown Is the tumor microsatellite instability-high (MSI-H) or mismatch repair deficient (dMMR)?							
☐ Yes ☐ No Will the requested drug be us	ed as a single agent?							
☐ Gestational Trophoblastic Neoplasia								
☐ Yes ☐ No Will the requested drug be us	ed as a single agent?							
☐ Yes ☐ No Is the disease resistant to multiagent chemotherapy?								
Please indicate the type of disease the patient	Please indicate the type of disease the patient has:							
☐ High-risk disease								
☐ Intermediate trophoblastic tumor (placental site trophoblastic tumor or epithelioid trophoblastic tumor)								
Please select the clinical setting in which the requested drug will be used:								
☐ Recurrent disease ☐ Progressiv	/e disease ☐ Other							
☐ Yes ☐ No Has the patient prev	iously received treatment with a platinum-based (e	∍.g., cisplatin, carboplat	tin) regimen?					
☐ Other								
Please indicate the patient's disease state:	Metastatic disease							
☐ Primary urothelial carcinoma of the urethra								
☐ Yes ☐ No Will the requested drug be us	ed as a single agent?							
Maintenance therapy request only:								
Yes No Will the requested drug be us	• •							
	sease progression on first-line platinum-containing	, chemotherapy (e.g., c	isplatin, carboplatin)?					
Subsequent therapy request only:								
Please indicate how the requested drug will be used: First line treatment Subsequent treatment								
Please select the clinical setting in which the requested drug will be used: ☐ Recurrent disease ☐ Locally advanced disease ☐ Metastatic disease ☐ Other								
_	ilsease Metastatic disease Other							
Renal Cell Carcinoma	and an analysis of the same dis							
Please indicate the clinical setting in which the								
☐ Advanced disease ☐ Relapsed disease ☐ Stage IV disease ☐ Other Please indicate how the requested drug will be used: ☐ First line treatment ☐ Subsequent treatment								
☐ Yes ☐ No Will the requested drug be us		auneni						
Upper genitourinary tract urothelial carcing								
Yes No Will the requested drug be us								
Maintenance therapy request only:	ed as a siligle agent:							
Yes ☐ No Will the requested drug be used as maintenance therapy?								
Yes No Did the patient experience disease progression on first-line platinum-containing chemotherapy (e.g., cisplatin, carboplatin)?								
Subsequent therapy request only:								
	e used: First line treatment Subsequent tre	eatment						
Please select the clinical setting in which the r	•							
☐ Locally advanced disease ☐ Metastatic d	·							

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Patient First Name	Patient Last Name	Patient Phone	Patient DOB						
		16 011							
	Required clinical information must be completed	for ALL precertification	n requests.						
Urothelial carcinoma of the prostate									
☐ Yes ☐ No Will the requested drug be u Maintenance therapy request only:	sed as a single agent?								
Maintenance therapy request only: ☐ Yes ☐ No Will the requested drug be used as maintenance therapy?									
☐ Yes ☐ No Did the patient experience disease progression on first-line platinum-containing chemotherapy (e.g., cisplatin, carboplatin)?									
Subsequent therapy request only:									
Please indicate how the requested drug will be used: First line therapy Subsequent therapy									
Please select the clinical setting in which the									
☐ Locally advanced disease ☐ Metastatic	disease								
For Continuation Requests (Clinical document	tation required for all requests):								
☐ Yes ☐ No Has the patient experienced disease progression or an unacceptable toxicity while on the current regimen?									
Yes No Is this infusion request in an out									
Yes \(\) No is the patient co	ntinuing on a maintenance regimen that includes p	rovider administered con	nbination chemotherapy?						
	periencing severe toxicity requiring continuous mo	nitoring (e.g., Grade 2-4	bullous dermatitis, transaminitis.						
	evens-Johnson syndrome, acute pancreatitis, prima								
I	litis, myocarditis, pericarditis, arrhythmias, impaired	ventricular function, con	nduction abnormalities)?						
Please explain:									
The state of the s	experienced an adverse event with the requested page. g., acetaminophen, steroids, diphenhydramine, flui								
,	event (anaphylaxis, anaphylactoid reactions, myoc		,						
immediately after		araidi inidiodon, dirombe	Johnsonom, or Johnson Julian G						
Please explain:									
	t have severe venous access issues that require th	e use of special interven	itions only available in the						
outpatient hosp	-								
Please explain:			the Association and the confedence						
	t have significant behavioral issues and/or physical rapy AND the patient does not have access to a ca		that would impact the safety of						
Please explain:		rogivor:							
	edically unstable which may include respiratory, cal								
I	to tolerate a large volume or load or predispose the		dverse event that cannot be						
	alternate setting without appropriate medical perso a description of the condition:	nnei and equipment?							
•	onary:								
☐ Other:									
☐ Yes ☐ No Is the patient wi	thin the initial 6 months of starting therapy?								
> Please indicate	how many continuous months of treatment the pat	ient has received with the	e requested drug:						
H. ACKNOWLEDGEMENT									
Request Completed By (Signature Required): 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 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.