

## Padcev® (enfortumab vedotin-ejfv) 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: Start		_	, ,					
Precertification P	☐ Continuation of therapy:  Requested By:	_			۶.		Fav	:	
A. PATIENT INFOR				FIIONE	·		гах	-	
A. PATIENT INFOR	RWATION		Last	: Name:					
Address:			City				State:	ZIP:	
		Work Phone:	City	•		Call Dhana:	State.	ZIF.	
Home Phone:	All	Work Phone:				Cell Phone:			
DOB:	Allergies:					Email:			
	lbs or k	gs Heigh	t:	inches o	or _	cm	S		
B. INSURANCE IN									
	#:			_					
					_ Car	rrier Name: _			
Medicare:  Yes	☐ No If yes, provide ID #: _		Med	licaid: Yes		No If yes, pr	ovide ID #:		
C. PRESCRIBER II	NFORMATION								
First Name:		Last Name:				(Check O		). D.O. N.	P.
Address:			(	City:			State:	ZIP:	
Phone:	Fax:	St Lic #:	I	NPI #:		DEA #:		UPIN:	
Provider Email:		Office Contact Na	ıme:				Phone	e:	
Specialty (Check of	one):	ner:							
Center Na  Home Infusion Agency N Administration Address:  PRODUCT INFO Request is for Par  Judge 10 10 10 10 10 10 10 10 10 10 10 10 10	sion Center Phone: ame: Center Phone: lame: code(s) (CPT):  DRMATION  dcev (enfortumab vedotin-ejfer FORMATION — Please indicate pr	v) Dose:		Physician's Specialty F Name: Address: Phone: TIN:	s Off Pharr	rice macy	☐ Retail Ph☐ Other:Fax:		
Primary ICD Code:	:	Secondary ICD Code:				_ Other ICD	Code:		
For All Requests (compared to the late of	RMATION — Required clinical infectinical documentation required.  If the requested drug be used as a steep patient ineligible for cisplatin-one in the patient recomposed in the patient recomposed in the patient recomposed in the patient recomposed in the requested drug be used as for the patient reatments in the requested drug be used for a steep patient in the requested drug be used for a clinical setting in which the requested disease ■ Metastatic disease in the patient in the requested drug be used for a clinical setting in which the requested disease ■ Metastatic disease in the patient in the requested drug be used for a clinical setting in which the requested disease ■ Metastatic disease in the patient in the p	I for all requests): a single agent? containing chemotherapy? containing chemotherapy. containin	a platin a progr equent ment nt post rrent o	um-containing ch ammed death red therapy? -cystectomy? r persistent disea	nemo cepto se in	therapy (e.g., r-1 (PD-1) or p preserved bla	cisplatin, cari programmed dder?	death-ligand	nerapy,
Urothelial carcinon	ich clinical setting the requested of ma- upper genitourinary tract tu clinical setting in which the reque	mors or urothelial carcin	oma d	of the prostate					Other



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Patient First Name	Patient Last Name	Patient Phone	Patient DOB										
G. CLINICAL INFORMATION (continued) – Required clinical information must be completed in its entirety for all precertification requests.													
For Continuation Requests (clinical documentation required for all requests):													
Yes No Has the patient experienced disease progression or an unacceptable toxicity while on the current regimen?													
H. ACKNOWLEDGEMENT													
Request Completed By (Signature Require	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.