

Paetna Erbitux® (cetuximab) Injectable **Medication Precertification Request**

Page 1 of 2

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

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

FAX: 1-888-267-3277

For Medicare Advantage Part B: Please Use Medicare Request Form

Please indicate:											
	☐ Continua	tion of therapy	: Date	of last treatment	/						
Precertification R		-				Phon	e:		Fa	x:	
A. PATIENT INFOR	RMATION										
First Name:				Last Name:					ı	1	
Address:						City:			State:	ZIP:	
Home Phone:			Work	Phone:				Cell P	hone:		
DOB:	Alle	ergies:							Emai	l:	
Current Weight:	lbs	or	kgs	Height: _		inches	or	cms	;		
B. INSURANCE INF	FORMATION										
Aetna Member ID	#:			Does patient have of		_					
Group #:				If yes, provide ID#: _			_ Carrier	Name:			
Insured:				Insured:							
Medicare: Tes		, provide ID #:		N	ledi	caid: Yes	☐ No	If yes, pro	ovide ID #:		
C. PRESCRIBER IN	NFORMATION			Look Names			(0	haala Onal	. \square M D		
First Name:				Last Name:			(C	песк Опе)] N.P. □ P.A.
Address:				Τ	_	City:	- 1		State:	ZIP:	
Phone:	Fax	K:		St Lic #:		NPI #:		DEA #:	1	UPIN:	
Provider Email:				Office Contact Name	e:				Phor	ne:	
				ologist 🗌 Other: _							
D. DISPENSING PR		NISTRATION IN	NFORM.	ATION							
Place of Administration: Dispensing Provider/Pharmacy: Patient Selected choice									oice		
☐ Self-administered ☐ Physician's Office					-	Physician's Office Retail Pharmacy					
Outpatient Infusion Center Phone: Center Name:				Specialty I			narmac	y L	Other:		
☐ Home Infusion (Phone:			_	Name:					
Home Infusion Center Phone: Agency Name:				Address							
☐ Administration c						Phone:			Fax	«:	
Address:					_	TIN:			PIN	l:	
E. PRODUCT INFO	RMATION										
Request is for Erb	itux: Dose: _			Frequency:							, [
F. DIAGNOSIS INF	ORMATION - F	Please indicate p	rimary I	CD Code and specify a	any o	other where app	licable.				
Primary ICD Code:			_ Seco	ondary ICD Code:				Other I	CD Code:		
				on must be completed i	n its	entirety for all p	precertific	ation reque	sts.		
For All Requests (
				arcinoma, anal ader							
Please indicate the clinical setting in which the requested drug will be used: Unresectable/inoperable disease Advanced disease Metastatic disease Other											
		• •		nical failure on panitum			_				
Please select which of the following applies to the patient:											
 ☐ RAS (KRAS and NRAS) mutation status is negative (wild-type): What is the place in therapy in which the requested drug will be used? ☐ First-line treatment ☐ Subsequent treatment 											
☐ Yes ☐ No Is this request for treatment of colon cancer?											
☐ Yes ☐ No Is the tumor left-sided? ☐ Yes ☐ No Is the tumor positive for BRAF V600E mutation?											
☐ Yes ☐ No Is the tumor positive for BRAF V600E mutation? ☐ Yes ☐ No Will the requested drug be used in combination with encorafenib (Braftovi)?											
☐ KRAS G12C mutation positive:											
What is the requested regimen? ☐ In combination with sotorasib (Lumakras) ☐ In combination with adagrasib (Krazati) ☐ Other ☐ Yes ☐ No Has the patient previously received treatment with chemotherapy?											
Other or unkn		oreviousiy recell	eu neal	ment with chemothera	μy!						
☐ Non-small cell lung cancer											
	☐ Yes ☐ No Will the requested drug be used in combination with afatinib (Gilotrif)? ☐ Yes ☐ No ☐ Unknown Does the patient have a known sensitizing epidermal growth factor receptor (EGFR) mutation?										
				_	-	-					inih Ilrac 1\0
∟ res ∟ No	nas the patie	Yes No Has the patient progressed on EGFR tyrosine kinase inhibitor therapy (e.g., afatinib [Gilotrif], erlotinib [Tarceva], gefitinib [Iressa])?									



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Page 2 of 2

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Patient First Name	Patient Last Name	Patient Phone	Patient DOB					
G. CLINICAL INFORMATION (continued) - F	Required clinical information must be comp	leted in its <u>entirety</u> for all precert	fication requests.					
What is the place in therapy in which the	requested drug will be used? ☐ Initial to	reatment 🔲 Subsequent trea	tment					
Please indicate the clinical setting in which	the requested drug will be used: Recur	rent disease 🔲 Advanced dis	sease					
☐ Other								
☐ Occult primary head and neck cancer								
☐ Yes ☐ No Will the requested drug I	be used as a single agent?							
☐ Yes ☐ No Will the requested drug I	be used for chemoradiation?							
☐ Penile cancer								
☐ Yes ☐ No Will the requested drug I	be used as a single agent?							
What is the place in therapy in which the requested drug will be used? Initial treatment Subsequent treatment								
Please indicate the clinical setting in which the requested drug will be used: Metastatic disease Other								
☐ Squamous cell carcinoma of the head	and neck							
☐ Yes ☐ No Is the patient unfit for surgery?								
☐ Yes ☐ No Will the requested drug be used in combination with radiation?								
Please indicate the clinical setting in which the requested drug will be used: Locally or regionally advanced disease Unresectable disease								
☐ Recurrent disease ☐ Persistent dise	ease							
☐ Squamous cell skin cancer								
☐ Yes ☐ No Will the requested drug I	8 8							
Please indicate the clinical setting in which the requested drug will be used: Unresectable/Inoperable/incompletely resected disease								
☐ Locally advanced disease ☐ Region		Distant metastatic disease 🗌	Other					
For continuation of therapy (clinical docu								
Yes No Is there evidence of disease	progression or unacceptable toxicity wh	nile on the current regimen?						
H. ACKNOWLEDGEMENT								
Request Completed By (Signature Requir	red):		Date: / /					
Any person who knowingly files a request for insurance company by providing materially insurance act, which is a crime and subjects	false information or conceals material	information for the purpose of						

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