

AVASTIN[®] (bevacizumab)
ALYMSYS[®] (bevacizumab-maly)
MVASI[®] (bevacizumab-awwb)
VEGZELMA[®] (bevacizumab-adcd)
ZIRABEV[™] (bevacizumab-bvzr)
Medication Precertification Request

Aetna Precertification Notification Phone: 1-866-752-7021 (TTY: 711)

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

For Medicare Advantage Part B: Please Use Medicare Request Form

Page 1 of 3 (All fields must be completed and legible for precertification review.)

Please indicate:		ent: Start date _	e completed and legible		•	review.)			
☐ Continuation of therapy, Date of						_			
Precertification Requested By:				Phone:		Fax:			
A. PATIENT INFO	RMATION								
First Name:			Last Name:					DOB:	
Address:				Cit	y:			State:	ZIP:
Home Phone:		Work Phone:		Се	Il Phone:			Email:	
Patient Current We	eight: lbs or _	kgs Patie	ent Height: inch	es o	or cms	Allergie	es:		
B. INSURANCE II	NFORMATION								
Aetna Member ID #:			Does patient have other coverage?			i □ No			
Group #:		If yes, provide ID#:		Carrier Name: _					
Insured:			Insured:						
	s ☐ No If yes, prov	ride ID #:	M	ledic	aid: 🗌 Yes [☐ No	If yes, prov	ride ID #:	
C. PRESCRIBER	INFORMATION								
First Name:			Last Name:	-			(Check O	ne):	☐ D.O. ☐ N.P. ☐ P.A
Address:					City:			State:	ZIP:
Phone:	Fax:		St Lic #:		NPI#:		DEA #:		UPIN:
Provider Email:			Office Contact Name	e:				Phone:	
Specialty (Check of	one): Oncologis	t 🗌 Ophthalm	ologist 🗌 Other: _					•	
D. DISPENSING F	PROVIDER/ADMINI	STRATION INFO	DRMATION						
Place of Administ					Dispensing P	Provide	r/Pharmac	y: Patient Se	lected choice
☐ Self-administered ☐ Physician's Office				□ Physician's Office			☐ Retail Pharmacy		
Outpatient Infu	sion Center F	hone:	Specialty Pharmacy			-			
Center Na	ame:				Name:		-		
☐ Home Infusion	Center F	hone:							
	ame:				Phone:				
Administration code(s) (CPT):			TIN:			PIN:			
E. PRODUCT INF									
Request is for:] AVASTIN™ (beva		☐ ALYMSYS™ (bev						
	ີ MVASI™ (bevacia	zumab-awwb)	☐ VEGZELMA® (bev	/aciz	umab-adcd)	☐ ZIR	ABEV™ (k	evacizumab-	-bvzr)
Dose:			Frequen						
		se indicate prima	ary ICD code and spec			e applic			
Primary ICD Code	e:	1 1: : 1: 6	Secondary ICD Co	ode:				ICD Code:	
			nation must be comple	etea	in its <u>entirety</u> to	or all pre	ecertificatio	n requests.	
Ophthalmic disord	<u>ests (clinical docum</u> lers:	entation require	d for all requests):						
	this request for Avasti	n treatment?							
I — — —	<u>.</u>		failed treatment with Av	/astin	due to a docun	nented i	ntolerable a	dverse event	
	_ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \	sh, nausea, vomit	0,						
_	_		nexpected and not attrib			•		•	ribing information?
Please select the o		patient had an ine	effective response, conf	traino	lication or intole	erance to	Byooviz O	R Cimerii?	
☐ Choroidal neova	ascularization (CNV) (0 , 1	choroidal neovasculariz pia, retinal dystrophies,		` ''				•
☐ Diabetic macula		_ , , ,			,			•	
	due to retinal vein oc	• •							
	et) Age-Related Macu	lar Degeneration	(AMD)						
☐ Neovascular gla									
☐ Polypoidal chord									
☐ Proliferative diabetic retinopathy ☐ Retinopathy of prematurity									



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Patient First Name	Patient Last Name	Patient Phone	Patient DOB			
G. CLINICAL INFORMATION (co	ntinued) – Required clinical in	nformation must be completed in i	ts <u>entirety</u> for all precertification			
Oncology indications:						
☐ Yes ☐ No Is this request for Mva						
		with Mvasi due to a documented intolera	ble adverse event			
` ` • `	rash, nausea, vomiting)?	and attributed to the active increalisation of	assisted in the prescripting information?			
Please select the diagnosis:	ne adverse event unexpected and r	not attributed to the active ingredient as d	escribed in the prescribing information?			
Ampullary Adenocarcinoma						
	ullarv adenocarcinoma which applie	s to the patient's disease: Intestinal-ty	pe			
	nt have progressive, unresectable, o					
Please select: [☐ progressive disease ☐ unresec	table disease 🔲 metastatic disease 🗀	none of the above			
☐ Angiosarcoma						
Yes No Will the request	ed medication be given as a single	agent therapy?				
☐ Breast cancer						
	t have recurrent or metastatic disea					
'	☐ recurrent disease ☐ metastatic	disease				
Cervical cancer						
Yes No Does the patient have persistent, recurrent, or metastatic disease?						
Please select: persistent disease metastatic disease none of the above						
☐ Colorectal cancer, including appendiceal adenocarcinoma and anal adenocarcinoma ☐ Diffuse high grade gliomas						
☐ Glioblastoma						
Glioma (WHO Grade 1)						
☐ Endometrial carcinoma						
Yes No Does the patien	it have progressive, advanced, recu	rrent, or metastatic disease?				
Please select: [☐ progressive disease ☐ advance	ed disease 🔲 recurrent disease 🔲 me	tastatic disease none of the above			
☐ Epithelial ovarian cancer						
☐ Fallopian tube cancer						
Hepatocellular carcinoma						
Yes No Does the patien						
Please select: ☐ unresectable disease ☐ metastatic disease ☐ none of the above						
 Yes ☐ No Will the requested drug be used as initial treatment? Yes ☐ No Will the requested medication be given in combination with atezolizumab (Tecentriq)? 						
☐ IDH mutant astrocytoma (WHO Grade 2, 3 or 4)						
☐ Intracranial and spinal ependymoma (excludes subependymoma)						
Limited and extensive brain metastases						
☐ Malignant sex cord stromal tumors						
☐ Medulloblastoma						
Meningiomas						
☐ Metastatic spine tumors						
Non-squamous non-small cell lung cancer (NSCLC)						
Yes No Does the patient have recurrent, advanced, metastatic, or unresectable disease?						
Please select: [☐ recurrent disease ☐ advanced	disease 🗌 metastatic disease 🔲 unre	sectable disease none of the above			

Continued on next page.



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Patient First Name	Patient Last Name	Patient Phone	Patient DOB
G. CLINICAL INFORMATION (conti	inued) – Required clinical information must b	e completed in its entirety for all pre	certification requests
☐ Mesothelioma		<u></u>	
└── Please indicate the type of meso	thelioma which applies to the patient's disease:		
	ma 🔲 malignant peritoneal mesothelioma 🔲	pericardial mesothelioma unica v	aginalis testis mesothelioma
other other			ļ
	apy in which the requested drug will be used:		
First-line treatment			
	quested medication be given in combination with		itin (Platinol) or carboplatin
`), followed by single-agent maintenance bevacial atient have unresectable disease?	zumad?	ļ
Subsequent treatment	atient have unlesectable disease:		
Please select the requested	t regimen:		
•	netrexed (Alimta) and either cisplatin (Platinol) o	r carboplatin (Paraplatin)	
	as the patient received immunotherapy as first-li		
☐ In combination with atez	•		
Other	, ,		
Oligodendroglioma (WHO Grade 2 c	or 3)		ļ
☐ Primary central nervous system lym	•		ļ
Primary peritoneal cancer	priorita		
Renal cell carcinoma			ļ
干	have relevand as store IV/ diagonal Tralenas	d disease	and of the above
_	have relapsed or stage IV disease? relapse	d disease stage iv disease n	one of the above
☐ Small bowel adenocarcinoma			
Solitary fibrous tumor or hemangiopo	•	lid- (Td)2	
→ ☐ Yes ☐ No Will the requeste	ed medication be given in combination with temo	ozolomide (Temodar)?	
T 5	have persistent, recurrent, or metastatic diseas	202	
· ·	persistent disease recurrent disease		nove
Uterine neoplasms	persistent disease recuirent disease	metastatic disease	040
	have progressive, advanced, recurrent, or meta	astatic disease?	
→ Please select: Γ	progressive disease advanced disease	☐ recurrent disease ☐ metastatic di	sease none of the above
	ous cell carcinoma and adenocarcinoma		_
Yes No Does the patient	have unresectable locally advanced, recurrent,	or metastatic disease?	
→ Please select:	unresectable locally advanced disease 🔲 re	current disease 🔲 metastatic diseas	e none of the above
For Continuation Requests (clinical d	locumentation required for all requests):		
Ophthalmic disorders:	-		
Yes No Has the patient demon	strated a positive clinical response to therapy (e	e.g., improvement or maintenance in be	est corrected visual acuity [BCVA]
or visual field, or a redu	uction in the rate of vision decline or the risk of r	nore severe vision loss)?	
Oncology indications:			
Yes No Has the patient experie	enced an unacceptable toxicity or disease progr	ession while on the current regimen?	
H. ACKNOWLEDGEMENT			
D	Day 1000		Data / /
Request Completed By (Signature			Date: //
	quest for authorization of coverage of a med		
	materially false information or conceals ma		misleading, commits a fraudulent
insurance act, which is a crime and s	ubjects such person to criminal and civil pen-	aities.	

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