

MEDICARE FORM

AVASTIN[™] (bevacizumab) ALYMSYS[™] (bevacizumab-maly) MVASI[™] (bevacizumab-awwb) VEGZELMA[®] (bevacizumab-adcd) ZIRABEV[™] (bevacizumab-bvzr) Medication Precertification Request FAX: 1-855-734-9389 PHONE: 1-855-364-0974 For other lines of business:

Please use other form

For Ohio MMP:

Note: Alymsys, Vegzelma, and Zirabev are non-preferred. The preferred products are Avastin and Mvasi.

Page 1 of 3

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

Please indicate: Sta	rt of treatmer	t: Start date _	/ /	☐ Continuation	on of therap	oy, Date c	of last treatmen	t//	
Precertification Request							Fax: _		
A. PATIENT INFORMATION									
First Name:			Last Name:				DOB:		
Address:			.I	City:			State:	ZIP:	
Home Phone:	1			Cell Phone:			Email:		
Patient Current Weight:	lbs or	kgs Patie	nt Height:inch	es orcr	ms Allergie	es:			
B. INSURANCE INFORM									
			Does patient have ot	her coverage?	☐ Yes	□No			
Group #:				If yes, provide ID#: Carrier Name: _					
Insured:			Insured:						
Medicare: ☐ Yes ☐ No		de ID #:	M	edicaid: 🗌 Ye	es 🗌 No I	If yes, prov	vide ID #:		
C. PRESCRIBER INFORM	MATION								
First Name:			Last Name:		(Check O	(Check One): M.D. D.O. N.P. F			
Address:				City:			State:	ZIP:	
Phone:	Fax:		St Lic #:	NPI #:		DEA #:		UPIN:	
Provider Email:			Office Contact Name	:			Phone:		
Specialty (Check one):	Oncologist	☐ Ophthalmo	ologist 🗌 Other: _						
☐ Administration code(s) (CPT): Address: E. PRODUCT INFORMATION Request is for: ☐ AVASTIN (bevacizumab) ☐ ALY ☐ VEGZELMA (bevacizumab-adcd) Dose:			∕MSYS™ (bevacizum □ ZIRABEV (bevac	Special Specia	Specialty Pharmacy Name: Address: Phone: TIN: maly)		Fax: PIN:		
F. DIAGNOSIS INFORMA							100.0		
Primary ICD Code: G. CLINICAL INFORMAT			_ Secondary ICD Co	<u> </u>					
For Initiation Requests (cli Ophthalmic disorders: ☐ Yes ☐ No Is this reque	est for Avastin No Has the p (e.g., rast) No Was the a is:	ntation required treatment? patient tried and f h, nausea, vomiti adverse event un cluding myopic o	d for all requests): failed treatment with Aving)? nexpected and not attributhoroidal neovascularize	astin due to a do outed to the activ ation (mCNV), a	ocumented ir re ingredient ngioid streak	ntolerable a as describ	adverse event ed in the prescri tis [including cho	proiditis secondary	
Diabetic macular edema Macular edema following Neovascular (wet) Age-F Neovascular glaucoma Polypoidal choroidal vas Proliferative diabetic retir	g retinal vein oc Related Macula culopathy nopathy	cclusion (RVO)		rubeosis mais, ρ	seuuoxanthi	oma elastic	oun, and trauma	J	



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Page 2 of 3

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

For Ohio MMP:

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Patient First Name	Patient Last Name	Patient Phone	Patient DOB			
G. CLINICAL INFORMATION (co	ntinued) – Required clinical in	formation must be completed in	its entirety for all precertification			
Oncology indications:						
Note: Alymsys, Vegzelma, and Zirab						
Yes No Has the patient had pr		-				
Yes No Has the patient had a		, ,	ect all that apply)			
	rumab)	·				
Please explain if there are any other m patient's diagnosis? (select all that app		not use any of the following preferred p	products when indicated for the			
1	יטיי: :umab)	anarb)				
	umab) 🔲 iwvasi (bevacizumab-av	wwb)				
☐ Yes ☐ No Is this request for Mva	si treatment?					
		with Mvasi due to a documented intoler	able adverse event			
` •	rash, nausea, vomiting)?	at attributed to the active ingredient co	described in the prescribing information?			
Please select the diagnosis:	ne adverse event unexpected and n	of altributed to the active ingredient as	described in the prescribing information?			
Ampullary Adenocarcinoma						
1 1	ıllary adenocarcinoma which applies	to the patient's disease: Intestinal-	type			
☐ Yes ☐ No Does the patien	t have progressive, unresectable, or	metastatic disease?				
	progressive disease unresect	able disease	none of the above			
☐ Anaplastic glioma						
Angiosarcoma						
☐ Yes ☐ No Will the requested medication be given as a single agent therapy? ☐ Breast cancer						
I T	t have recurrent or metastatic diseas	e?				
	☐ recurrent disease ☐ metastatic d					
☐ Cervical cancer						
	t have persistent, recurrent, or metas					
		isease	ne of the above			
Colorectal cancer, including append	diceal adenocarcinoma and anal ade	nocarcinoma				
☐ Glioblastoma ☐ Endometrial carcinoma						
Yes No Does the patient have progressive, advanced, recurrent, or metastatic disease?						
			netastatic disease			
Epithelial ovarian cancer (including carcinosarcoma [malignant mixed Müllerian tumors], clear cell carcinoma, mucinous carcinoma, endometrioid						
carcinoma, serous carcinoma, and malignant sex cord-stromal tumors)						
Fallopian tube cancer						
Hepatocellular carcinoma	t have upresentable or materialis di	22222				
 Yes						
Yes No Will the requested drug be used as initial treatment?						
☐ Yes ☐ No Will the requested medication be given in combination with atezolizumab (Tecentriq)?						
☐ Intracranial and spinal ependymoma (excludes subependymoma)						
☐ Limited and extensive brain metastases						
Low-grade (WHO Grade 1 or 2) Glioma						
☐ Medulloblastoma						
☐ Meningiomas						
Metastatic spine tumors Non squampus pon small cell lung cancer (NSCLC)						
□ 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, metastatic, or unresectable disease unresectable disease none of the above						
Fiease select. [unciasialic disease U un	I HOLIE OF THE WOOLE			

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

Patient First Name	Patient Last Name	Patient Phone	Patient DOB					
G. CLINICAL INFORMATION (cont	inued) – Required clinical information m	nust be completed in its entiret	y for all precertification requests					
☐ Mesothelioma	·	·						
— — .	othelioma which applies to the patient's dis	ease:						
· · · · · · · · · · · · · · · · · · ·	ma malignant peritoneal mesotheliom		☐ tunica vaginalis testis mesothelioma					
other .	_		_ 。					
Please indicate the place in therapy in which the requested drug will be used:								
☐ First-line treatment								
· — —	quested medication be given in combination	. ,	either cisplatin (Platinol) or carboplatin					
` .	n), followed by single-agent maintenance b	evacizumab?						
	patient have unresectable disease?							
Subsequent treatment	d acceptance							
Please select the requeste		in all an and an latin (Danan latin)						
	netrexed (Alimta) and either cisplatin (Plati as the patient received immunotherapy as							
☐ In combination with ate	, , , , , , , , , , , , , , , , , , , ,	mst-me treatment?						
Other	conzumab (recenting)							
Primary central nervous system lym	inhoma							
☐ Primary peritoneal cancer								
Renal cell carcinoma								
T	t have relapsed or stage IV disease? 🔲 re	elanced disease	sease none of the above					
_ · ·	Thave relapsed of stage in disease:	siapsed disease	sease I none of the above					
☐ Small bowel adenocarcinoma☐ Solitary fibrous tumor or hemangiop	oorioytama							
1 .	ed medication be given in combination with	temozolomide (Temodar)?						
☐ Vaginal cancer	su medication be given in combination with	rtemozolomide (remodar):						
	t have persistent recurrent or metastatic	disease?						
→ ☐ Yes ☐ No Does the patient have persistent, recurrent, or metastatic disease? → Please select: ☐ persistent disease ☐ recurrent disease ☐ metastatic disease ☐ none of the above								
Uterine neoplasms								
Yes No Does the patient have progressive, advanced, recurrent, or metastatic disease?								
Please select: progressive disease advanced disease recurrent disease metastatic disease none of the above								
☐ Vulvar squamous cell carcinoma								
Yes No Does the patient have unresectable locally advanced, recurrent, or metastatic disease?								
Please select: unresectable locally advanced disease recurrent disease metastatic disease none of the above								
For Continuation Requests (clinical of	documentation required for all requests	<u>):</u>						
Ophthalmic disorders:								
	nstrated a positive clinical response to ther uction in the rate of vision decline or the ris		enance in best corrected visual acuity [BCVA]					
Oncology indications:								
Yes No Has the patient experienced an unacceptable toxicity or disease progression while on the current regimen?								
H. ACKNOWLEDGEMENT								
Request Completed By (Signature	Required):		Date: //					
any insurance company by providing		ls material information for the	e with the intent to injure, defraud or deceive purpose of misleading, commits a fraudulent					

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