

Fax completed form to: (855) 840-1678 If this is an URGENT request, please call (800) 882-4462 (800.88.CIGNA)

Alymsys (bevacizumab-maly) Avastin (bevacizumab)

PHYSICIAN INFORMATION			PATIENT INFORMATION				
* Physician Name:			*Due to privacy regulations we will not be able to respond via fax with the outcome of our review unless all asterisked (*) items on this				
Specialty:	* DEA, NPI or	TIN:		form are completed.*			
Office Contact Person:		* Patient Name:					
Office Phone:			* Cigna ID:		* Date of I	Birth:	
Office Fax:			* Patient Street Address:				
Office Street Address:			City:		State:		Zip:
City:	State:	Zip:	Patient	Phone:			
Urgency: ☐ Standard ☐ Urgent (In checking this box, I attest to the fact that applying the standard review time frame may seriously jeopardize the customer's life, health, or ability to regain maximum function)							
Medication Requested: ☐ Avastin ☐ Alymsys Is this a new start? ☐ Yes ☐ No Start date: Dose: Frequency of therapy: Duration of therapy: Will this medication be given concurrently with other agents? ☐ Yes ☐ No If yes, please specify: What is your patient's current weight? ICD10:							
Where will this medication be obtained? Accredo Specialty Pharmacy** Prescriber's office stock (billing on a medical claim form) Other (please specify): **Medication orders can be placed with Accredo via E-prescribe - Accredo (1620 Century Center Pkwy, Memphis, TN 38134-8822 NCPDP 4436920), Fax 888.302.1028, or Verbal 866.759.1557						ecialty pharmacy	
Facility and/or doctor dis Facility Name: Address (City, State, Zip Cod	St	Iministering (ate:	medicat	tion: Tax ID#:			
NOTE: Per some	Cigna plans, infusio	on of medication	n MUST	occur in the lowest cost	, medically	appropria	te setting
Is this infusion occurring in a facility affiliated with hospital outpatient setting?				☐ Yes ☐ No			
If yes- Is this patient a candid Option Case Manager?			• •	such as AIS, MDO, hon edical necessity rationa	,	sistance of	f a Specialty Care
Is your patient a candidate for home infusion? Does the physician have an in-office infusion site?						Yes No No Yes No No	
Is the requested medication f the patient?	or a chronic or lonç	g-term conditior	n for whic	th the prescription medi	cation may	be necess	sary for the life of ☐ Yes ☐ No
Diagnosis: AIDS-related Kaposi sarce Ampullary adenocarcinom angiosarcoma cervical cancer (carcinom colon or rectal cancer (col non-small cell lung cancer endometrial cancer epithelial ovarian cancer (hepatocellular carcinoma small bowel adenocarcino other (please specify):	a of the cervix) orectal cancer, CR (NSCLC) including serous, n (HCC)	,	metrioid,	☐ fallopian tube cance ☐ granulosa cell ovari ☐ CNS/brain tumor ☐ pleural mesothelion ☐ primary peritoneal cell radiation necrosis aell cancer (RCclear-cell, Brenner or tree solitary fibrous tume vulvar squamous cell	an cancer cancer ind uncontr CC) ansitional or	cell) iopericytor	

(if CNS/brain tumor) What is your patient's diagnosis? ☐ anaplastic glioma (including anaplastic astrocytoma, anaplastic oligodendroglioma and anaplastic oligoastrocytom ☐ central nervous system (CNS) brain metastases ☐ central nervous system (CNS) meningioma ☐ ependymoma ☐ glioblastoma (including glioblastoma multiforme) ☐ leptomeningeal metastases ☐ medulloblastoma	na)
primary central nervous system (CNS) lymphoma subependymoma spine tumor other (please specify): (if other to either question above) Is this use related to chemotherapy or oncology (cancer)?	Yes ☐ No ☐
Clinical Information (if NSCLC) Does your patient have non-squamous cell NSCLC? (if NSCLC) Does your patient have unresectable, locally advanced, recurrent, or metastatic disease? (if NSCLC) Is the drug requested being given as first-line therapy? Yes No, patient has tried other drugs before for this diagnosis	Yes No Yes No No
☐ Unknown (if first-line) Will the drug requested be given in combination with carboplatin and paclitaxel?	Yes ☐ No ☐
(if pleural mesothelioma) Will the drug requested be used in combination with pemetrexed (Alimta, Pemfexy) and EIT Paraplatin (carboplatin)? (if pleural mesothelioma) What is your patient's stage? ☐ stage 1 (I)-stage 3a (IIIa) ☐ stage 3b (IIIb)-stage 4 (IV)	HER cisplatin or Yes
 ☐ unknown (if stage 1-3a) Does your patient have unresectable disease? (if not unresectable OR unknown stage) Does your patient have medically inoperable tumors? (if inoperable tumors) What is your patient performance status? ☐ PS 0-2 ☐ PS 3-4 ☐ unknown 	Yes No Yes No No
(if cervical) Does your patient have persistent, recurrent, or metastatic disease? (if cervical) Will the drug requested be used in combination with paclitaxel and either cisplatin or carboplatin OR paclitopotecan (Hycamtin)?	Yes ☐ No ☐ taxel and Yes ☐ No ☐
(if CRC or spine tumor) Does your patient have metastatic disease? (if CRC) How is the drug requested being used in your patient's treatment? in combination with a fluorouracil (Adrucil, 5-FU) based chemotherapy regimen in combination with fluoropyrimidine-irinotecan (Camptosar)- OR fluoropyrimidine-oxaliplatin-based chemotherapy In combination with trifluridine and tipiracil (Lonsurf) other	Yes 🗌 No 🗍
(if in combo with Lonsurf chemo) Is this medication being used as second-line treatment in patients who have treated with fluoropyrimidine-, oxaliplatin- and irinotecan-based chemotherapy, an anti-VEGF biological ther wild-type, an anti-EGFR therapy? (if in combo with 5-FU chemo) Is this medication being used as a first or second-line therapy? (if in combo with fluoropyrimidine-irinotecan- or fluoropyrimidine-oxaliplatin-based chemo) Did your patient progression while on a first-line bevacizumab (Alymsys, Avastin, Mvasi, Vegzelma, and Zirabev)-containing	apy, AND if RAS Yes ☐ No ☐ Yes ☐ No ☐ have disease
(if endometrial) Which of the following best describes the requested drug's role in your patient's therapy? ☐ for disease progression after failure of first-line therapy ☐ for the treatment of advanced or recurrent disease ☐ other	
other (if advanced or recurrent) Will the drug requested be used in combination with carboplatin and paclitaxel?	Yes 🗌 No 🗌
(if HCC) Does your patient have unresectable or metastatic disease? (if HCC) Will the requested drug be used in combination with Tecentriq (atezolizumab)? (if HCC) Has the patient received prior systemic therapy for this diagnosis in the past?	Yes
(if RCC) Does your patient have relapsed or metastatic disease? (if RCC) What is the histology of the disease? □ non-clear cell □ predominantly clear cell □ other	Yes 🗌 No 🗌

(if non-clear) Does your patient have advanced papillary renal cell carcinoma [RCC] (including hereditary leiomyom cell cancer [HLRCC])?	natosis and renal Yes
(if yes) Will the drug requested be used in combination with Afinitor (everolimus) or Tarceva (erlotinib)?	Yes 🗌 No 🗌
(if predominant clear cell) Which best describes how the drug requested will be used? ☐ as first-line therapy ☐ following disease progression while on previous therapy ☐ neither of the above	
(if non-clear or after disease progression with clear cell) Will the drug requested be used as single-agent therapy? (if predominant clear cell and first-line) Will the drug requested be used in combination with Intron-A?	Yes No Yes No
(if granulosa cell ovarian) Does your patient have relapsed disease?	Yes 🗌 No 🗌
(if angiosarcoma, CNS brain mets, endometrial, ependymoma, granulosa cell ovarian, lep mets, medulloblastoma, pr lymphoma, spine tumor, radiation necrosis and uncontrolled cerebral edema) Will the drug requested be used as sing	
(if anaplastic glioma or glioblastoma) Does your patient have recurrent disease?	Yes 🗌 No 🗌
(if ependymoma) Does the patient have progressive disease?	Yes 🗌 No 🗌
(if CNS meningioma) Does your patient have recurrent or progressive disease? (if CNS meningioma) Is the lesion surgically inaccessible (meaning that standard surgical techniques can't reach it)? (if CNS meningioma) Is radiation a possible option?	Yes No Yes No Yes No
(if CNS brain mets, lep mets, or spine tumor) Is the drug requested being given to control symptoms? (if solitary fibrous tumor/hemangiopericytoma) Will the drug requested be used in combination with Temodar (temozo	Yes ☐ No ☐ olomide)? Yes ☐ No ☐
(if epithelial ovarian, fallopian tube, peritoneal) Is your patient's cancer associated with homologous recombination depositive status? (if HRD positive) Did the patient have gene testing showing genomic instability AND/OR a deleterious or suspected mutation? (if yes) Has your patient had a complete or partial response to first-line platinum-based chemotherapy (carb cisplatin)? (if complete or partial response) Will the requested drug be used for first-line maintenance treatment? (if first-line maintenance) Does your patient have advanced disease? (if advanced disease) Will the requested drug be used in combination with Lynparza (olaparib)?	eficiency (HRD) Yes ☐ No ☐ I deleterious BRCA Yes ☐ No ☐
(if epithelial ovarian, fallopian tube, or primary peritoneal and not to ANY of the previous 6 questions) Does your patie or IV disease? (if stage III or IV) Has your patient had surgical resection? (if resection) Will/Was the drug requested used in combination with carboplatin and paclitaxel, followed by s therapy with bevacizumab (Alymsys, Avastin, Mvasi, Zirabev)?	Yes No Yes No No
(if no to any of the previous 3 questions) Does your patient have persistent or recurrent disease? (if persistent or recurrent) Has your patient been treated with bevacizumab (Alymsys, Avastin, Mvasi, Zirabev) before	
(if treated with bevacizumab before) Is your patient currently on bevacizumab (Alymsys, Avastin, Mvasi, or Zirabev diagnosis?	Yes ☐ No ☐) for this Yes ☐ No ☐
(if no bevacizumab before OR currently on) Will the drug requested be used as single-agent therapy?	Yes 🗌 No 🗌
(if not single agent) Was your patient previously treated with carboplatin or cisplatin (platinum therapy)?	
 Yes, and patient was platinum-refractory (no response with progression during treatment) Yes, and patient was platinum-resistant (showed initial response to chemotherapy but relapsed within 6 round of chemotherapy) Yes, and patient was platinum-sensitive No, patient was not treated with platinum therapy Unknown 	months of last
(if platinum-sensitive) Will the drug requested be used in combination with EITHER paclitaxel and carboplati (Gemzar) and carboplatin)? (if platinum-resistant) Will the drug requested be used in combination with liposomal doxorubicin (Doxil or Li OR topotecan (Hycamtin)?	Yes 🔲 No 🗌

(if epithelial ovarian) Which type of epithelial tumor does your patient have? ☐ serous or endometrioid ☐ mucinous ☐ clear cell ☐ unknown or other	
(if serous/endometrioid or mucinous) Will the drug requested be used as adjuvant therapy?	Yes 🗌 No 🗌
(if mucinous and NOT adjuvant) Does your patient have persistent or recurrent disease?	Yes 🗌 No 🗌
(if serous/endometrioid) What is the tumor grade? ☐ grade 1 ☐ grade 2 ☐ grade 3 ☐ unknown	
(if serous/endometrioid, mucinous, or granulosa cell) What is your patient's cancer stage? ☐ Stage 1 (I) ☐ Stage 2 (II) ☐ Stage 3 (III) ☐ Stage 4 (IV) ☐ unknown	
(if serous/endometrioid, adjuvant, and stage II/III/IV) Will the drug requested be used in combination with car paclitaxel?	rboplatin and Yes
(if mucinous, adjuvant and stage II/III/IV) Is the drug requested being used as any of the following? ☐ as combination therapy with carboplatin or paclitaxel ☐ as combination therapy with capecitabine (Xeloda) and oxaliplatin ☐ as combination therapy with fluorouracil (Adrucil, 5-FU), leucovorin, and oxaliplatin ☐ none of the above	
(if mucinous and persistent or recurrent) Is the drug requested being used as any of the following? ☐ as combination therapy with fluorouracil (Adrucil, 5-FU), leucovorin, and oxaliplatin ☐ as combination therapy with capecitabine (Xeloda) and oxaliplatin ☐ neither of the above	
(if small bowel adenocarcinoma) Will this drug be used in combination with either a Xeloda (capecitabine) or a 5-fluor regimen?	ouracil (5-FU) Yes
(if small bowel adenocarcinoma) Does the patient have advanced or metastatic disease?	Yes 🗌 No 🗌
(if small bowel adenocarcinoma) Will the patient be using this medication as initial therapy?	Yes 🗌 No 🗌
(if no) Will the patient be using this medication as subsequent therapy in patients who previously received in Opdivo (nivolumab)?	itial therapy with Yes
(if vulvar squamous cell carcinoma) Will the drug requested be used in combination with paclitaxel and EITHER cisple (carboplatin)?	atin or Paraplatin Yes
(if vulvar squamous cell carcinoma) Which best describes your patient's diagnosis? unresectable locally advanced disease with residual tumor at primary site locally advanced disease with positive margins following resection as primary treatment for metastatic disease beyond the pelvis for isolated groin/pelvic recurrence if prior external beam radiation therapy (EBRT) for clinical nodal or distant recurrence with multiple pelvic nodes, distant metastasis, or prior pelvic EBRT other	
The covered alternatives are: Mvasi (bevacizumab-awwb) and Zirabev (bevacizumab-bvzr). For the alternatives tried medication name and strength, date(s) taken and for how long, and what the documented results were of taking each including any intolerances or adverse reactions your patient experienced.	

For Mvasi (bevacizumab-awwb), which of the following applies to your patient? Patient has not tried this medication. Patient tried this medication, but it didn't work or didn't work well enough. Patient tried this medication, but had an allergic or adverse reaction. Other
(If allergic or adverse reaction) Is there documentation that this reaction was due to a formulation difference in the inactive ingredients between the requested medication and Mvasi (bevacizumab-awwb) (for example, difference in dyes, fillers, preservatives)? Yes ☐ No ☐
(If yes) Please provide details to support
For Zirabev (bevacizumab-bvzr), which of the following applies to your patient? Patient has not tried this medication. Patient tried this medication, but it didn't work or didn't work well enough. Patient tried this medication, but had an allergic or adverse reaction. Other
(If allergic or adverse reaction) Is there documentation that this reaction was due to a formulation difference in the inactive ingredients between the requested medication and Zirabev (bevacizumab-bvzr) (for example, difference in dyes, fillers, preservatives)? Yes ☐ No ☐
(If yes) Please provide details to support.
Additional pertinent information: (including disease stage, prior therapy, performance status, and names/doses/admin schedule of any agents to be used concurrently)
Attestation: I attest the information provided is true and accurate to the best of my knowledge. I understand that the Health Plan or insurer its designees may perform a routine audit and request the medical information necessary to verify the accuracy of the information reported on this form.
Prescriber Signature: Date:
Save Time! Submit Online at: www.covermymeds.com/main/prior-authorization-forms/cigna/ or via SureScripts in your EHR.
Our standard response time for prescription drug coverage requests is 5 business days. If your request is urgent, it is important that you call us to expedite the request. View our Prescription Drug List and Coverage Policies online at cigna.com.

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