# BlueCross BlueShield

# BEVACIZUMAB PRIOR APPROVAL REQUEST

Send completed form to: Service Benefit Plan Prior Approval P.O. Box 52080 MC 139 Phoenix, AZ 85072-2080 Attn. Clinical Services Fax: 1-877-378-4727

Federal Employee Program. **PRIOR APPROVAL REQUEST** Additional information is required to process your claim for prescription drugs. Please complete the patient portion, and have the prescribing physician complete the physician portion and submit this completed form.

Patient Information (required)			<b>Provider Information</b> (required)			
Date:			Provider Name:			
Patient Name:			Specialty:	N	IPI:	
Date of Birth:	Sex: Male	□Female	Office Phone:	0	Office Fax:	
Street Address:			Office Street Address:			
City:	State:	Zip:	City:	State:	Zip:	
Patient ID:			Physician Signature:			
	Pl	HYSICIAN (	COMPLETES			
NOTE: Form must be completed in its entirety for processing						

#### Please select medication:

Alymsys (bevacizumab-maly)	□Avastin (bevacizumab)	☐Mvasi (bevacizumab-awwb)		
□Vegzelma (bevacizumab-adcd)	□Zirabev (bevacizumab-bvzr)			

\*\*Check www.fepblue.org/formulary to confirm which medication is part of the patient's benefit

Has the patient been on this medication continuously for the last 6 months <u>excluding samples</u>? *Please select answer below:* **YES** – this is a PA renewal for **CONTINUATION** of therapy, please answer the questions on <u>PAGE 3</u>
**NO** - this is **INITIATION** of therapy, please answer the questions below:

2. Is this request for brand or generic? Brand Generic

3. Requests for Alymsys (bevacizumab-maly), Avastin (bevacizumab), or Vegzelma (bevacizumab-adcd): Does the patient have an intolerance or contraindication or have they had an inadequate treatment response to ONE of the following medications: Zirabev or Mvasi?  $\Box$ Yes  $\Box$ No

## 4. What is the patient's diagnosis?

Glioblastoma multiforme (GBM)

a. Will this medication be used as a single-agent therapy? **D**Yes **D**No

b. Has there been progression of the disease following prior therapy? □Yes □No

□ Metastatic cervical cancer <u>OR</u> □ Persistent cervical cancer <u>OR</u> □ Recurrent cervical cancer

- a. Will the patient be treated with paclitaxel (Taxol)? **U**Yes **U**No
- b. Will the patient be treated with cisplatin?  $\Box$ Yes  $\Box$ No\*
  - \**If NO*, will the patient be treated with topotecan (Hycamtin)?  $\Box$ Yes  $\Box$ No

Detastatic colorectal cancer

a. Is this medication being used as first-line treatment? Please select answer below:

□Yes, first-line treatment: Is the patient receiving concurrent IV chemotherapy with 5-Fluorouracil (5-FU)? □Yes □No

□No: Will the patient be receiving concurrent therapy with ONE of the following regimens: fluoropyrimidine and irinotecan-based chemotherapy, fluoropyrimidine and oxaliplatin-based chemotherapy, 5-fluorouracil-based chemotherapy, or trifluridine and tipiracil (Lonsurf)? □Yes\* □No

\*If YES, select answer: D5-Fluorouracil-based chemotherapy DFluoropyrimidine-irinotecan based chemotherapy

□Fluoropyrimidine-oxaliplatin based chemotherapy □Trifluridine and tipiracil (Lonsurf)

 $\Box Metastatic hepatocellular carcinoma (HCC) \underline{OR} \Box Unresectable hepatocellular carcinoma (HCC)$ 

a. Has the patient received prior systemic therapy? □Yes □No

b. Will this medication be given in combination with atezolizumab (Tecentriq)? **\Box** Yes **\Box** No

Detastatic renal cell carcinoma

a. Will the patient be receiving concurrent therapy with interferon-alfa? Yes No

## PLEASE PROCEED TO PAGE 2 FOR ADDITIONAL DIAGNOSES

PAGE 1 of 4



Other (*please specify*): \_

# BEVACIZUMAB

Federal Employee Program. PRIOR APPROVAL REQUEST

Additional information is required to process your claim for prescription drugs. Please complete the patient portion, and have the prescribing physician complete the physician portion and submit this completed form.

Send completed form to: Service Benefit Plan Prior Approval P.O. Box 52080 MC 139 Phoenix, AZ 85072-2080 Attn. Clinical Services Fax: 1-877-378-4727

PAGE 2 - PHYSICIAN COMPLETES				
Patient Name:	DOB:	Patient ID: R		
Non squemous non small call lu	a concor			
□Non-squamous non-small cell lui a. Is this medication being use	-	□No		
-				
b. Is the cancer unresectable, le	•			
c. Will the patient be receiving				
Ocular disease resulting from intr a. Please select one of the follo		cluding.		
Angioid streaks	Ocular histoplasmosis	□Macular edema secondary to retinal vascular occlusion		
<ul><li>Diabetic macular edema</li><li>Neovascular glaucoma</li></ul>	<ul><li>Progressive high myopia</li><li>Retinopathy of prematurity</li></ul>	□Neovascular (Wet) Age-related Macular Degeneration (AMD) □Proliferative diabetic retinopathy		
b. Will this medication be used indications? □Yes* □No		scular Endothelial Growth Factor (VEGF) inhibitors for ocular		
*If YES, please specify th	e medication:			
*VEGF Inhibitors: Beovu Vabysmo (faricimab-svoa)		a HD (aflibercept), Lucentis (ranibizumab), Susvimo (ranibizumab),		
$\Box$ Epithelial ovarian cancer <u>OR</u>		<b>DR D</b> Primary peritoneal cancer		
-	-	Yes* (*If YES, answer the following questions)		
	or stage IV disease? <b>\Box</b> Yes			
ii. Will this medication be	-	boplatin (Paraplatin) and paclitaxel (Taxol) for up to 6 cycles $\Box$ No		
b. Is the cancer recurrent plati	•••			
*If YES, please select one of	of the following:			
		tion be given concurrently with paclitaxel (Taxol/Onxal), or topotecan (Hycamtin)?		
* <i>If YES</i> , please s	elect one of the following belo	)W:		
□paclitaxel (Taxe	ol/Onxal) Degylated liposon	nal doxorubicin (Doxil/Caelyx)		
		tion be given in combination with carboplatin (Paraplatin) and a single agent? □Yes □No*		
* <i>If NO</i> , will this	•	nation with carboplatin (Paraplatin) and gemcitabine (Gemzar)		
-		* (*If YES, answer the following questions)		
i. Will this medication be	given in combination with ola	parib (Lynparza)? Tyes INo		
ii. Has the patient had a c	omplete or partial response to	platinum-based chemotherapy?		
*If YES, please select of	one of the following below:			
Complete resp	oonse to platinum-based chem-	otherapy Partial response to platinum-based chemotherapy		
d. Is the cancer associated wit	h homologous recombination	deficiency (HRD) positive status?  Yes*  No		
		Distitive status defined by deleterious or suspected deleterious BRCA <i>If YES, select one of the following below)</i> □No		
Deleterious or	suspected deleterious BRCA r	nutation <u>OR</u> Genomic instability		

PAGE 2 of 4

# BlueCross BlueShield

# BEVACIZUMAB PRIOR APPROVAL REQUEST

Send completed form to: Service Benefit Plan Prior Approval P.O. Box 52080 MC 139 Phoenix, AZ 85072-2080 Attn. Clinical Services Fax: 1-877-378-4727

Federal Employee Program. PRIOR

Additional information is required to process your claim for prescription drugs. Please complete the patient portion, and have the prescribing physician complete the physician portion and submit this completed form.

Patient Information (required)			<b>Provider Information</b> (required)			
Date:			Provider Name:			
Patient Name:			Specialty:		NPI:	
Date of Birth:Sex: <b>M</b> ale <b>F</b> emale		□Female	Office Phone:		Office Fax:	
Street Address:		Office Street Address:				
City:	State:	Zip:	City:	Stat	te:	Zip:
Patient ID:			Physician Signature:			
	Р	HYSICIAN (	COMPLETES			
<b>CONTINUATION OF THERAPY (PA RENEWAL)</b>						
<b>NOTE:</b> Form must be completed in its <b>entirety</b> for processing						

Please	select	medication:
--------	--------	-------------

□Alymsys (bevacizumab-maly)	□Avastin (bevacizumab)	□Mvasi (bevacizumab-awwb)
□Vegzelma (bevacizumab-adcd)	□Zirabev (bevacizumab-bvzr)	

\*\*Check www.fepblue.org/formulary to confirm which medication is part of the patient's benefit

Has the patient been on this medication continuously for the last 6 months excluding samples? *Please select answer below:* □ NO - this is INITIATION of therapy, please answer the questions on <u>PAGE 1</u>

**YES** – this is a PA renewal for **CONTINUATION** of therapy, please answer the questions below:

2. Is this request for brand or generic? Brand Generic

3. What is the patient's diagnosis?

Glioblastoma multiforme (GBM)

a. Will this medication be used as a single-agent therapy?  $\Box$ Yes  $\Box$ No

□ Metastatic cervical cancer <u>OR</u> □ Persistent cervical cancer <u>OR</u> □ Recurrent cervical cancer

a. Will the patient be treated with paclitaxel (Taxol)? **U**Yes **U**No

b. Will the patient be treated with cisplatin? Yes No\*

\**If NO*, will the patient be treated with topotecan (Hycamtin)?  $\Box$ Yes  $\Box$ No

□ Metastatic colorectal cancer

a. Is this medication being used as first-line treatment? Please select answer below:

 $\Box$  Yes, first-line treatment: Is the patient receiving concurrent IV chemotherapy with 5-Fluorouracil (5-FU)?  $\Box$  Yes  $\Box$  No

❑No: Will the patient be receiving concurrent therapy with ONE of the following regimens: fluoropyrimidine and irinotecan-based chemotherapy, fluoropyrimidine and oxaliplatin-based chemotherapy, 5-fluorouracil-based chemotherapy, or trifluridine and tipiracil (Lonsurf)? □Yes\* □No

\**If YES*, select answer: D5-Fluorouracil-based chemotherapy DFluoropyrimidine-irinotecan based chemotherapy

Fluoropyrimidine-oxaliplatin based chemotherapy Trifluridine and tipiracil (Lonsurf)

 $\Box Metastatic hepatocellular carcinoma (HCC) \underline{OR} \Box Unresectable hepatocellular carcinoma (HCC)$ 

a. Will this medication be given in combination with atezolizumab (Tecentriq)?  $\Box$ Yes  $\Box$ No

Detastatic renal cell carcinoma

a. Will the patient be receiving concurrent therapy with interferon-alfa?  $\Box$ Yes  $\Box$ No

□Non-squamous non-small cell lung cancer

a. Will the patient be receiving concurrent therapy with carboplatin and paclitaxel? **D**Yes **D**No

## PLEASE PROCEED TO PAGE 4 FOR ADDITIONAL DIAGNOSES

PAGE 3 of 4



# BEVACIZUMAB

Federal Employee Program. PRIOR APPROVAL REQUEST

Additional information is required to process your claim for prescription drugs. Please complete the patient portion, and have the prescribing physician complete the patient portion and submit this completed form.

Send completed form to: Service Benefit Plan Prior Approval P.O. Box 52080 MC 139 Phoenix, AZ 85072-2080 Attn. Clinical Services Fax: 1-877-378-4727

#### **PAGE 2 - PHYSICIAN COMPLETES**

DOB: Patient ID: R Patient Name: \_\_\_\_ Ocular disease resulting from intravitreal neovascularization including: a. Please select one of the following below: Angioid streaks Ocular histoplasmosis Macular edema secondary to retinal vascular occlusion Diabetic macular edema □Progressive high myopia Neovascular (Wet) Age-related Macular Degeneration (AMD) □Neovascular glaucoma □Retinopathy of prematurity □Proliferative diabetic retinopathy b. Will this medication be used in combination with other Vascular Endothelial Growth Factor (VEGF) inhibitors for ocular indications? **D**Yes\* **No** \**If YES*, please specify the medication: \*VEGF Inhibitors: Beovu (brolucizumab-dbll), Eylea/Eylea HD (aflibercept), Lucentis (ranibizumab), Susvimo (ranibizumab), Vabysmo (faricimab-svoa) OR □Fallopian tube cancer **OR** Primary peritoneal cancer Epithelial ovarian cancer a. Will this medication be used as single agent therapy post initial surgical resection?  $\Box$ Yes  $\Box$ No b. Is the cancer recurrent platinum resistant or recurrent platinum sensitive? □Yes\* □Cancer is not recurrent \*If YES, please select one of the following: **Recurrent Platinum Resistant:** Will this medication be given concurrently with paclitaxel (Taxol/Onxal), pegylated liposomal doxorubicin (Doxil/Caelyx), or topotecan (Hycamtin)? **U**Yes\* \*If YES, please select one of the following below: □paclitaxel (Taxol/Onxal) □pegylated liposomal doxorubicin (Doxil/Caelyx) □topotecan (Hycamtin) **Recurrent Platinum Sensitive:** Will this medication be used as single agent therapy? □No c. Is the patient's cancer considered to be advanced? **D**Yes\* **D**No \**If YES*, will this medication be given in combination with olaparib (Lynparza)? **U**Yes Other (*please specify*): \_

PAGE 4 of 4