

## Lucentis, Byooviz Cimerli

## **CareFirst Prior Authorization Request**

CVS Caremark administers the prescription benefit plan for the patient identified. This patient's benefit plan requires prior authorization for certain medications in order for the drug to be covered. To make an appropriate determination, providing the most accurate diagnosis for the use of the prescribed medication is necessary. **Please respond below and fax this form to CVS Caremark toll-free at 1-855-330-1720**. If you have questions regarding the prior authorization, please contact CVS Caremark at **1-888-877-0518**. For inquiries or questions related to the patient's eligibility, drug copay or medication delivery; please contact the Specialty Customer Care Team: CaremarkConnect® 1-800-237-2767.

The recipient of this fax may make a request to opt-out of receiving telemarketing fax transmissions from CVS Caremark. There are numerous ways you may opt-out: The recipient may call the toll-free number at 877-265-2711, at any time, 24 hours a day/7 days a week. The recipient may also send an opt-out request via email to do not call@cvscaremark.com. An opt out request is only valid if it (1) identifies the number to which the request relates, and (2) if the person/entity making the request does not, subsequent to the request, provide express invitation or permission to CVS Caremark to send facsimile advertisements to such person/entity at that particular number. CVS Caremark is required by law to honor an opt-out request within thirty days of receipt.

Patient's Name:		Date:
Patient's ID:		Patient's Date of Birth:
Physician's Name:		
Specialty:		NPI#:
Physician Office Telephone:		Physician Office Fax:
Referring Provider Info: 🗖 Same as Ro	equesting Provid	er
Name:		NPI#:
Fax:		Phone:
Rendering Provider Info: □ Same as Ro Name:	_	
Fax:		Phone:
Required Demographic Information:		
Patient Weight:	kg	
Patient Height:		
Please indicate the place of service for the	requested drug:	
		☐ Off Campus Outpatient Hospital
On Campus Outpatient Hospital	<b>□</b> Office	☐ Pharmacy
What product is being requested?   Luc	entis	ooviz

Exc	ception Criteria Questions: Skip to Criteria questions if the member is MD Risk or VA Risk
A.	Is the product being requested for the treatment of an ocular disorder?  ☐ Yes, Continue to B ☐ No, Skip to Clinical Criteria Questions
B.	Which product is being requested?  ☐ Lucentis, <i>Skip to F</i> ☐ Byooviz, <i>Continue to C</i> ☐ Cimerli, <i>Continue to C</i>
C.	The preferred product for your patient's health plan is Avastin.  Can the patient's treatment be switched to Avastin?  ☐ Yes, Please obtain Form for preferred product and submit for corresponding PA.  ☐ No, Continue to D
D.	Is the patient in an active treatment plan with the requested product?  ☐ Yes, <i>Skip to Clinical Criteria Questions</i> ☐ No, <i>Continue to E</i>
E.	Did the patient have a documented inadequate response, intolerable adverse event, or contraindication to the preferred product Avastin? <i>Action Required: If Yes, attach supporting chart note(s).</i> Yes, <i>Skip to Clinical Criteria Questions</i> No, <i>Skip to Clinical Criteria Questions</i>
F.	The preferred product for your patient's health plan is Avastin.  Can the patient's treatment be switched to the Avastin?  ☐ Yes, <i>Please obtain Form for preferred product and submit for corresponding PA</i> .  ☐ No, <i>Continue to G</i>
G.	Is the patient in an active treatment plan with the requested product?  ☐ Yes, <i>Skip to Clinical Criteria Questions</i> ☐ No, <i>Continue to H</i>
H.	Did the patient have a documented inadequate response, intolerable adverse event, or contraindication to the preferred product Avastin? <i>ACTION REQUIRED: If 'Yes', attach supporting chart note(s)</i> .  Yes, <i>Continue to I</i> No, <i>Continue to I</i>
I.	Did the patient have a documented inadequate response, intolerable adverse event, or contraindication to Byooviz? <i>ACTION REQUIRED: If 'Yes', attach supporting chart note(s)</i> .  Yes, inadequate response, <i>Skip to K</i> Yes, intolerable adverse event, <i>Continue to J</i> Yes, contraindication, <i>Skip to K</i> No, <i>Continue to J</i>
J.	Was the documented intolerable adverse event an expected adverse event attributed to the active ingredient as described in the prescribing information (i.e., known adverse reaction for both the reference product and biosimilar products)? <i>ACTION REQUIRED: If no, attach supporting chart note(s)</i> . $\square$ Yes, <i>Continue to K</i> $\square$ No, <i>Continue to K</i>

K.	Did the patient have a documented inadequate response Action Required: If yes, attach supporting chart notes  ☐ Yes, inadequate response, Skip to M ☐ Yes, intolerable adverse event, Continue to L ☐ Yes, contraindication, Skip to M ☐ No, Continue to L	se, intolerable adverse event, or contraindication to Cimerli? $e(s)$ .
L.		ected adverse event attributed to the active ingredient as adverse reaction for both the reference product and biosimilatorting chart note(s).
M.	Did the patient have a documented inadequate response Vabysmo)? <i>Action Required: If yes, attach supporting</i> ☐ Yes, <i>Continue to Clinical Criteria Questions</i> ☐ No, <i>Continue to Clinical Criteria Questions</i>	
<u>Cri</u>	teria Questions:	
1.	What is the diagnosis?	
	Diabetic Macular Edema, Continue to 2	
	Neovascular (Wet) Age-Related Macular Degeneration	n, Continue to 2
	Macular Edema Following Retinal Vein Occlusion, Co	ontinue to 2
	Diabetic Retinopathy, Continue to 2	
	Myopic Choroidal Neovascularization, Continue to 2	
	Other, please specify,	Continue to 2
	Is this a request for continuation of therapy? Yes, <i>Continue to 3</i> No, <i>No Further Questions</i>	
se	Has the patient demonstrated a positive clinical response rected visual acuity [BCVA] or visual field, or a reductive vision loss)?  Yes, No Further Questions No, No Further Questions	se to therapy (e.g., improvement or maintenance in best ction in the rate of vision decline or the risk of more

Step Therapy Override: Complete if Applicable for the state of Maryland.		Please Circle	
Is the requested drug being used to treat stage four advanced metastatic cancer?	Yes	No	
Is the requested drug's use consistent with the FDA-approved indication or the National Comprehensive Cancer Network Drugs & Biologics Compendium indication for the	Yes	No	
treatment of stage four advanced metastatic cancer and is supported by peer-reviewed medical literature?			
Is the requested drug being used for an FDA-approved indication OR an indication supported in the compendia of current literature (examples: AHFS, Lexicomp, Clinical Pharmacology, Micromedex, current accepted guidelines)?	Yes	No	
Does the prescribed quantity fall within the manufacturer's published dosing guidelines or within dosing guidelines found in the compendia of current literature (examples: package insert, AHFS, Lexicomp, Clinical Pharmacology, Micromedex, current accepted guidelines)?	Yes	No	
Do patient chart notes document the requested drug was ordered with a paid claim at the pharmacy, the pharmacy filled the prescription and delivered to the patient or other documentation that the requested drug was prescribed for the patient in the last 180 days?	Yes	No	
Has the prescriber provided proof documented in the patient chart notes that in their opinion the requested drug is effective for the patient's condition?	Yes	No	

Step Therapy Override: Complete if Applicable for the state of Virginia.		Please Circle	
Is the requested drug being used for an FDA-approved indication or an indication supported in the compendia of current literature (examples: AHFS, Micromedex, current accepted guidelines)?	Yes	No	
Does the prescribed dose and quantity fall within the FDA-approved labeling or within dosing guidelines found in the compendia of current literature?	Yes	No	
Is the request for a brand drug that has an AB-rated generic equivalent or interchangeable biological product available?	Yes	No	
Has the patient had a trial and failure of the AB-rated generic equivalent or interchangeable biological product due to an adverse event (examples: rash, nausea, vomiting, anaphylaxis) that is thought to be due to an inactive ingredient?			
Is the preferred drug contraindicated?	Yes	No	
Is the preferred drug expected to be ineffective based on the known clinical characteristics of the patient and the prescription drug regimen?		No	
Has the patient tried the preferred drug while on their current or previous health benefit plan and it was discontinued due to lack of efficacy or effectiveness, diminished effect, or an adverse event?		No	
Is the patient currently receiving a positive therapeutic outcome with the requested drug for their medical condition?	Yes	No	

I attest that this information is accurate and true, and that documentation supporting this information is available for review if requested by CVS Caremark or the benefit plan sponsor.

X	
Prescriber or Authorized Signature	Date (mm/dd/yy)