

MEDICARE FORM

Lucentis® (ranibizumab), Byooviz™ (ranibizumab-nuna), Cimerli™ (ranibizumab-eqrn) Ínjectable **Medication Precertification Request**

Page 1 of 2

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

For Michigan MMP: 1-844-241-2495

PHONE: 1-855-676-5772 (TTY: 711)

For other lines of business: Please use other form.

Note: Lucentis and Cimerli are nonpreferred. The preferred products are bevacizumab (Avastin) first followed by Byooviz. Avastin (C9257) and bevacizumab biosimilars do not require

Please indicate:	☐ Start of treatment: Sta				precertification	for ophthalmic use.
Precertification Re	equested By:			e:	Fax:	
A. PATIENT INFOR	MATION					
First Name:		Last Name:			DOB:	
Address:		-	City:		State:	ZIP:
Home Phone:	Work Ph	one:	Cell Phone:		E-mail:	·
Current Weight:	lbs orkgs He	ight: inches or	cms Allergies:			
B. INSURANCE INF	ORMATION					
Aetna Member ID #	# :	Does patient ha	ave other coverage?	☐ Yes ☐ No		
			ID#:	Carrier Name:		
Medicare: Yes [☐ No If yes, provide ID #:	<u> </u>	Medicaid: ☐ Yes ☐			
C. PRESCRIBER IN	FORMATION					
First Name:		Last Name:		(Check Or	e):	D.O. 🗌 N.P. 🗌 P.A
Address:		<u>.</u>	City:		State:	ZIP:
Phone:	Fax:	St Lic #:	NPI #:	DEA #:	UI	PIN:
Provider Email:	•	Office Contact	Name:	Phone:	•	
D. DISPENSING PR	OVIDER/ADMINISTRATION	INFORMATION				
Center Nar Home Infusion C Agency Na Administration C Address: City: Phone:	ion Center Phone: _ me: Center Phone: _ ime: code(s) (CPT): State: Fax: PIN:	ZIP:	Other:	_	State: Fax: PIN:	ZIP:
	RWATION Lucentis (ranibizumab) [Ryooviz (ranihizumak	nuna) 🗆 Cimorli (ra	nibizumah oarn)		
Dose:	Lucentis (rambizumab)		Frequency:	ilibizulliab-eqili)	HCPCS co	ode:
	DRMATION – Please indicate	primary ICD Code and spe		olicable.		
					Code:	
	RMATION – Required clinical					
Note: Lucentis and and bevacizumab Yes No Ha Yes No Ha Yes No Ha Please explain if the	merli Requests: (clinical description of Cimerli are non-preferred biosimilars do not require as the patient had a trial and as the patient had a trial and as the patient had a trial and are are are any other medical refere	d. The preferred products precertification for optopy with Lucentis (ranibiz failure, intolerance, or contained in the patient of the patient	ets are bevacizumab (Anthalmic use. umab) or Cimerli (ranibizontraindication to bevaciontraindication to Byooverannot use bevacizumab	zumab-eqrn) within izumab (Avastin)? iz (ranibizumab-nur o (Avastin).	the last 365 day	, ,

Continued on next page



MEDICARE FORM

Lucentis® (ranibizumab), Byooviz™ (ranibizumab-nuna), Cimerli™ (ranibizumab-eqrn) Injectable **Medication Precertification Request**

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

For Michigan MMP: FAX: 1-844-241-2495

PHONE: 1-855-676-5772 (TTY: 711)

For other lines of business:

Please use other form.

Note: Lucentis and Cimerli are nonpreferred. The preferred products are bevacizumab (Avastin) first followed by Byooviz. Avastin (C9257) and bevacizumab biosimilars do not require precertification for ophthalmic use.

Patient First Name	Patient Last Name	Patient Phone	Patient DOB						
C. CLINICAL INFORMATION (continued)	Described alinical information must be soon	lated in its antiraty for all proporti	Spatian regulate						
G. CLINICAL INFORMATION (continued) – Required clinical information must be completed in its entirety for all precertification requests.									
For Byooviz Requests: (clinical documentation required for all requests)									
Note: Bevacizumab (Avastin) is preferred first prior to Byooviz. Avastin (C9257) and bevacizumab biosimilars do not require precertification for ophthalmic use.									
Yes No Has the patient had prior therapy with Byooviz (ranibizumab-nuna) within the last 365 days?									
☐ Yes ☐ No Has the patient had a trial and failure, intolerance, or contraindication to bevacizumab (Avastin)?									
Please explain if there are any other medical reason(s) that the patient cannot use bevacizumab (Avastin).									
What is the patient's BCVA (best corrected visual acuity) prior to initiating treatment:/(e.g., 20/320)									
☐ Yes ☐ No Is this request for intravitrea									
→ Please indicate which eye: ☐ OD (right eye) ☐ OS (left eye) ☐ OU (both eyes)									
Yes No Will Lucentis (ranibizumab)			inhibitor?						
Yes No Will the medication be given in the same eye as Lucentis (ranibizumab)?									
Yes No Does the patient have any of the following contraindications to Lucentis (ranibizumab)? (check all that apply)									
Endophthalmitis Ocular infection Hypersensitivity									
Please identify which documented diagnosis the patient is being treated for: Diabetic retinopathy Diabetic macular edema Macular edema following retinal vein occlusion (RVO) Polypoidal choroidal vasculopathy									
☐ Myopic Choroidal Neovascularization (mCNV) ☐ Neovascular (wet) (age related macular degeneration) AMD ☐ Neovascular glaucoma									
☐ Pseudoxanthoma elasticum	()(3	3 ,							
Yes No Is this a reque	est for re-treatment?								
Rare causes of choroidal neovascularization									
Please identify the cause of choroidal neovascularization:									
Angioid streaks Choroiditis (including choroiditis secondary to ocular histoplasmosis) Idiopathic degenerative myopia									
Retinal dystrophies Rubeosis iridis Trauma Other: Please identify:									
Yes No Is this a request for re-treatment?									
☐ ☐ ☐ ☐ ☐ ☐ ☐ ☐ ☐ ☐ ☐ ☐ ☐ ☐ ☐ ☐ ☐ ☐ ☐									
Please indicate the stage of disease: Stage 1 Stage 2 Stage 3 Stage 4 Stage 5									
For Continuation Requests:									
Please indicate length of time on Lucentis (ranibizumab), Byooviz (ranibizumab-nuna), or Cimerli (ranibizumab-eqrn):									
Please indicate the patient's current BCVA:/(e.g., 20/320)									
Please choose the patient response: BC	·								
☐ Small vision loss (defined as maximum of 3 lines or 15 letters lost on visual acuity exam)									
□ None of the above									
	Yes No Has the patient had improvement in field vision?								
Yes No Has the patient experienced a hypersensitivity reaction to Lucentis (ranibizumab), Byooviz (ranibizumab-nuna), or Cimerli (ranibizumab-egrn)?									
Please indicate which of the following hypersensitivity reactions the patient experienced:									
☐ anaphylactoid reactions ☐ pruritus ☐ rash ☐ severe anaphylactic reactions ☐ severe intraocular inflammation									
urticaria Other: Please explain:									
Yes No Is this continuation request		of Lucentis (ranibizumab), Byo	oviz (ranibizumab-nuna), or						
Cimerli (ranibizumab-eqrn)?									
H. ACKNOWLEDGEMENT									
Request Completed By (Signature Requi	red):		Date: /						
Any person who knowingly files a request for authorization of coverage of a medical procedure or service with the intent to injure, defraud or deceive any insurance company by providing materially false information or conceals material information for the purpose of misleading, commits a fraudulent insurance act, which is a crime and subjects such person to criminal and civil penalties.									

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