



An FDA-approved treatment for retinal conditions¹

Available in 0.3 mg & 0.5 mg dosing strengths



Who is CIMERLI® for?

CIMERLI® is a prescription medicine used to treat patients with:

Wet age-related macular degeneration (wAMD)

Diabetic macular edema and diabetic retinopathy (DME and DR) Myopic choroidal neovascularization (mCNV)

Macular edema following retinal vein occlusion (RVO)

IMPORTANT SAFETY INFORMATION

What Important Safety Information should I know about CIMERLI® (ranibizumab-eqrn)?

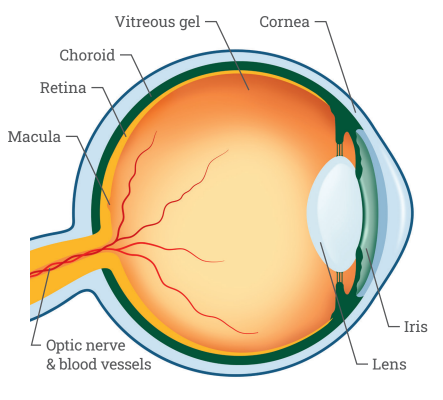
• You should not receive CIMERLI® if you have an infection in or around the eye or are allergic to CIMERLI® or any of its ingredients. CIMERLI® is a prescription medication given by injection into the eye, and it has side effects. CIMERLI® is not for everyone. Some patients using ranibizumab products have had detached retinas and serious eye infections. If your eye becomes red, sensitive to light, or painful, or if you have a change in vision, call or visit your eye doctor right away

Understandingyour eye condition

If you've been diagnosed with

- Wet age-related macular degeneration (wAMD)
- Diabetic retinopathy and/or diabetic macular edema (DR, DME)
- Myopic choroidal neovascularization (mCNV)
- Macular edema following retinal vein occlusion (RVO)

you may be experiencing issues with your vision.



Healthy eye

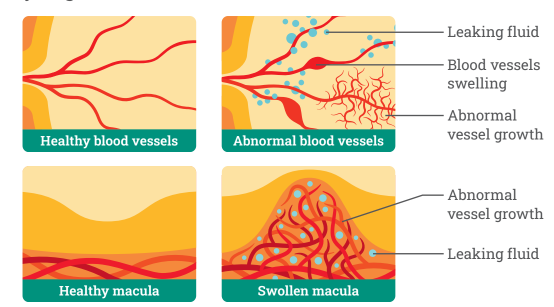
IMPORTANT SAFETY INFORMATION (CONTINUED)

What Important Safety Information should I know about CIMERLI® (ranibizumab-eqrn)?

- Some patients have had increased eye pressure before and within 1 hour of an injection. Your eye doctor should check your eye pressure and eye health before and after your CIMERLI® injection
- Uncommonly, patients using ranibizumab products have had serious, sometimes fatal, problems
 related to blood clots, such as heart attacks or strokes. Fatal events were seen more often in patients
 with DME and DR with ranibizumab products compared with patients who did not receive ranibizumab
 products. Although there were only few fatal events which included causes of death typical of patients
 with advanced diabetic complications, these events may be caused by ranibizumab products

In these conditions, problems with eyesight can occur when:

- Abnormal blood vessels grow in a part of the eye known as the retina
- Or, when these blood vessels leak fluid into an area of the retina called the macula



One of the contributing factors that leads to the overgrowth of these blood vessels is the presence of a protein called vascular endothelial growth factor (VEGF). Medicine to help treat these conditions, called anti-VEGF treatment, works by blocking the VEGF protein.

CIMERLI® is an anti-VEGF treatment that slows the growth of abnormal blood vessels in the eye.¹

IMPORTANT SAFETY INFORMATION (CONTINUED)

What Important Safety Information should I know about CIMERLI® (ranibizumab-eqrn)?

- Severe inflammation of the veins in the retina, with or without blockage, usually with preexisting
 inflammation in the eye or after other treatments in the eye, have been reported with the use of
 ranibizumab products. You should tell your eye doctor if you have any changes in vision immediately.
 Your eye doctor may stop your treatment
- Some patients using ranibizumab products have serious side effects related to the injection. These include serious infections inside the eye, detached retinas, and cataracts. The most common eye-related side effects are increased redness in the white of the eye, eye pain, small specks in vision, and increased eye pressure. The most common non-eye-related side effects are nose and throat infections, anemia, nausea, and cough



What is CIMERLI®?

CIMERLI® is an FDA-approved biosimilar that is interchangeable with Lucentis® (ranibizumab injection)²

CIMERLI® is a biosimilar—a type of biologic medicine that is highly similar to the FDA-approved biologic Lucentis®.3

CIMERLI® can be used to treat **the same eye conditions as Lucentis®**, and as an interchangeable biosimilar it is expected to have **the same effectiveness and safety**, which makes it possible to be directly substituted for Lucentis®.^{1,3}

With no clinically meaningful differences between the two medicines, your doctor can choose to start or transition you to CIMERLI® without impacting your treatment.

IMPORTANT SAFETY INFORMATION (CONTINUED)

What Important Safety Information should I know about CIMERLI® (ranibizumab-eqrn)?

- Reports of a tear in the outermost layer of the retina in patients with neovascular AMD (age-related macular degeneration) have been reported after treatment with ranibizumab products
- CIMERLI® is for prescription use only

Why are biosimilars important?^{4,5}

Biosimilars were established to

- Help reduce out-of-pocket& medication costs for patients
- Make important treatments accessible to patients who need them
- Potentially help reduce healthcare spending

all while maintaining the same quality, safety, and effectiveness of treatment.



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What you can expect from CIMERLI^{®1,3}



Your doctor has chosen to start or transition you to CIMERLI®.

Here are some things you may find helpful to know:



The same treatment approach: Both treatments are given as injections to the eye and are dosed the same way. Treatment is typically recommended monthly, but your doctor will determine a dosing schedule that is appropriate for you.



The same way of working: Both treatments work the same way in the eye and are expected to produce the same clinical results.



The same effectiveness and safety: CIMERLI® can be substituted for Lucentis® without compromising treatment impact or safety. CIMERLI® was carefully reviewed and thoroughly evaluated by a rigorous FDA-approval process, and it is required to meet high FDA standards for consistent manufacturing to ensure medication quality and safety.

IMPORTANT SAFETY INFORMATION (CONTINUED)

What Important Safety Information should I know about CIMERLI® (ranibizumab-eqrn)?

• Some patients have had increased eye pressure before and within 1 hour of an injection. Your eye doctor should check your eye pressure and eye health before and after your CIMERLI® injection





Financial support is available

Sandoz One Source® for CIMERLI® offers a range of financial assistance programs for patients who are prescribed CIMERLI®.

Please see following pages for more information.

IMPORTANT SAFETY INFORMATION (CONTINUED)

What Important Safety Information should I know about CIMERLI® (ranibizumab-eqrn)?

 Uncommonly, patients using ranibizumab products have had serious, sometimes fatal, problems related to blood clots, such as heart attacks or strokes. Fatal events were seen more often in patients with DME and DR with ranibizumab products compared with patients who did not receive ranibizumab products. Although there were only few fatal events which included causes of death typical of patients with advanced diabetic complications, these events may be caused by ranibizumab products



Financial support is available

Co-Pay Savings Program

The Co-Pay Savings Program may cover out-of-pocket expenses associated with CIMERLI® (ranibizumab-eqrn) and the injection procedure for eligible patients with commercial insurance*:

Please see the financial support eligibility criteria on page 10.



per dose of CIMERLI® including injection with a maximum annual benefit of \$16,000 per calendar year*

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Your eye doctor may stop your treatment

Financial support is available

Sandoz Patient Assistance (SPA)

CIMERLI® (ranibizumab-eqrn) may be available to you at no cost if you are uninsured or functionally underinsured.

Independent Foundation Support

Sandoz One Source® may be able to help you find financial support through charitable foundations, or you can choose to contact these independent assistance foundations directly.

Please see the financial support eligibility criteria on page 10.

Ask your doctor's office to help you apply to CIMERLI® financial support programs

To learn more, visit CIMERLI.com/patients or contact 1-844-4SANDOZ (1-844-472-6369).

IMPORTANT SAFETY INFORMATION (CONTINUED)

What Important Safety Information should I know about CIMERLI® (ranibizumab-eqrn)?

Some patients using ranibizumab products have serious side effects related to the injection.
These include serious infections inside the eye, detached retinas, and cataracts. The most common
eye-related side effects are increased redness in the white of the eye, eye pain, small specks in vision,
and increased eye pressure. The most common non-eye-related side effects are nose and throat
infections, anemia, nausea, and cough

Financial support eligibility criteria

*Patient Drug and Injection Co-Pay Eligibility Criteria:

- Be prescribed CIMERLI® (ranibizumab-eqrn) for a medically appropriate purpose consistent with its FDA-approved labeling within 180 days of program enrollment
- Have commercial (private or non-governmental)
 health insurance that covers the medication costs
 of CIMFRI I®
- For Administration: Not a resident or get treatments in Minnesota, Rhode Island
 - If a resident of Massachusetts, injection administration may only be paid directly to the patient. Additional information may be required
- Not covered by any federal, state, or governmentfunded healthcare program, such as Medicare, Medicare Advantage, Medicare Part D, Veterans Affairs, Department of Defense, or TRICARE
- Not seek reimbursement from any third party, including payers, charitable foundations, or flexible spending account (FSAs) or healthcare savings accounts (HSAs) for all or any part of the benefit received by Coherus through this program
- Other restrictions apply, see <u>Terms & Conditions</u> at CIMERLI.com
- It is not valid for cash-paying patients or where prohibited by law
- Co-Pay Savings Program subject to change or discontinuation without notice. This is not health insurance

To be eligible for SPA assistance you must:

- Reside in the United States or a U.S. Territory
- Have limited or no prescription insurance coverage
- Meet income guidelines adjusted for household size, for the medication for which the patient is seeking assistance
- Have a valid prescription for the Sandoz medication
- Be treated by a licensed U.S. healthcare provider
- Complete and sign consent form and, when applicable, provide income documentation

IMPORTANT SAFETY INFORMATION (CONTINUED)

What Important Safety Information should I know about CIMERLI® (ranibizumab-eqrn)?

- Reports of a tear in the outermost layer of the retina in patients with neovascular AMD (age-related macular degeneration) have been reported after treatment with ranibizumab products
- CIMERLI® is for prescription use only

Call your doctor for medical advice about side effects. To report SUSPECTED ADVERSE REACTIONS, contact Sandoz, Inc at 1-800-525-8747 or FDA at 1-800-FDA-1088 or www.fda.gov/medwatch. For additional Safety Information, please talk to your doctor and see Important Safety Information throughout. Click for <u>full Prescribing Information</u>.

References: 1. CIMERLI® (ranibizumab-eqrn) prescribing information. Princeton, NJ: Sandoz, Inc. **2.** Purple Book Database of Licensed Biological Products. U.S. Food and Drug Administration. https://purplebooksearch.fda.gov/faqs. Updated 2024. Accessed on June 4, 2024. **3.** Biosimilar and Interchangeable Products. https://www.fda.gov/drugs/biosimilars/biosimilar-and-interchangeable-products#biological. Published October 23, 2017. Accessed on May 19, 2022. **4.** The U.S. Generic & Biosimilar Medicines Savings Report. Accessiblemeds.org. https://accessiblemeds.org/sites/default/files/2021-10/AAM-2021-US-Generic-Biosimilar-Medicines-Savings-Report-web.pdf. Published 2021. Accessed on February 22, 2022. **5.** Makurvet F. Biologics vs. small molecules: Drug costs and patient access. *Med Drug Discov*. 2021;9:100075. doi:10.1016/j.medidd.2020.100075



CIMERLI® is given as an injection in the eye to treat certain retinal conditions¹:

- Wet age-related macular degeneration (wAMD)
- Diabetic retinopathy and/or diabetic macular edema (DR, DME)
- Myopic choroidal neovascularization (mCNV)
- Macular edema following retinal vein occlusion (RVO)



IMPORTANT SAFETY INFORMATION (CONTINUED)

What Important Safety Information should I know about CIMERLI® (ranibizumab-eqrn) injection?

• Some patients have had increased eye pressure before and within 1 hour of an injection. Your eye doctor should check your eye pressure and eye health before and after your CIMERLI® injection

For additional Safety Information, please talk to your doctor and click for the CIMERLI® <u>full Prescribing Information</u>.

Financial support is available with Sandoz One Source® for CIMERLI®

To learn more, visit **CIMERLI.com/patients** or contact 1-844-4SANDOZ (1-844-472-6369).



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