

Updated: 05/2024 PARP Approved: 05/2024

Prior Authorization Criteria **Luxturna (voretigene neparvovec-rzyl)**

All requests for Luxturna (voretigene neparvovec-rzyl) require a prior authorization and will be screened for medical necessity and appropriateness using the criteria listed below.

Coverage may be provided with a <u>diagnosis</u> of biallelic *RPE65* mutation-associated retinal dystrophy and the following criteria is met:

- Must have a diagnosis of retinal dystrophy with confirmed RPE65 mutation in both alleles confirmed by both of the following:
 - O Clinical documentation confirming diagnosis of Leber congenital amaurosis (LCA) or Retinitis pigmetosa (RP) including clinical features, funduscopic appearance, and results of testing such as dark-adapted thresholds, Ganzfeld-flash electroretinography (ERG), and when appropriate, perimetry.
 - Documentation of positive genetic test result confirming a biallelic pathogenic or likely pathogenic RPE65 mutation (homozygote or compound heterozygote) by a CLIA-approved mutational test.
- Must be prescribed by or in consultation with an ophthalmologist
- Must have viable retinal cells as determined by at least one of the following:
 - o Area of retina within the posterior pole of greater than 100 µm thickness per OCT
 - o At least 3 disc areas of retina without atrophy or pigmentary degeneration within the posterior pole
 - o Remaining visual field within 30 degrees of fixation as measured by a III4e isopter or equivalent
- Is age-appropriate according to FDA-approved package labeling, nationally recognized compendia, or peer-reviewed medical literature
- The requested dose and frequency is in accordance with FDA-approved labeling, nationally recognized compendia, and/or evidence-based practice guidelines
- Has not previously received treatment with voretigene neparvovec-rzyl in the requested treatment eye(s)
- **Initial Duration of Approval:** 1 injection per eye (1 month)
- Reauthorization criteria
 - o None one time use

Coverage may be provided for any non-FDA labeled indication if it is determined that the use is a medically accepted indication supported by nationally recognized pharmacy compendia or peer-reviewed medical literature for treatment of the diagnosis(es) for which it is prescribed. These requests will be reviewed on a case by case basis to determine medical necessity.

When criteria are not met, the request will be forwarded to a Medical Director for review. The physician reviewer must override criteria when, in their professional judgment, the requested medication is medically necessary.



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LUXTURNA PRIOR AUTHORIZATION FORM

Please complete and fax all requested information below including any progress notes, laboratory test results, or chart documentation			
as applicable to Highmark Wholecare Pharmacy Services. FAX: (888) 245-2049			
If needed, you may call to speak to a Pharmacy Services Representative. PHONE : (800) 392-1147 Mon – Fri 8:30am to 5:00pm			
PROVIDER INFORMATION			
Requesting Provider:		Provider NPI:	
Provider Specialty:		Office Contact:	
State license #:		Office NPI:	
Office Address:		Office Phone:	
		Office Fax:	
MEMBER INFORMATION			
Member Name: DOB:			
Member ID: Member v		<u> </u>	
REQUESTED DRUG INFORMATION			
Medication: Strength:			
Directions:	Quantit	•	
Is the member currently receiving requested medication? \(\subseteq \text{Yes} \)	No	Date Medication Initiated:	
Billing Information			
This medication will be billed: at a pharmacy OR medically, JCODE:			
Place of Service: Hospital Provider's office Member'		Other	
Place of Service Information			
Name:		NPI:	
Address:		Phone:	
MEDICAL HISTORY (Complete for ALL requests)			
Diagnosis:	ICD Cod	le:	
Does the member have confirmed RPE65 mutation in both alleles?	Yes	No	
Is there clinical documentation confirming diagnosis of Leber congenital amaurosis (LCA) or Retinitis pigmetosa (RP) including			
clinical features, funduscopic appearance, and results of testing such as dark-adapted thresholds, Ganzfeld-flash electroretinography			
(ERG), and when appropriate, perimetry? Yes No			
Is there documentation of positive genetic test result confirming a biallelic pathogenic or likely pathogenic RPE65 mutation			
(homozygote or compound heterozygote) by a MoIDX-approved mutational test? Yes No			
Does the member have viable retinal cells? Select all that apply to the member:			
Area of retina within the posterior pole of greater than 100 µm thickness per optical coherence tomography (OCT)			
☐ At least 3 disc areas of retina without atrophy or pigmentary degeneration within the posterior pole ☐ Remaining visual field within 30 degrees of fixation as measured by a III4e isopter or equivalent			
Which eye is being treated? Left Right Both			
Has the member previously received treatment with voretigene neparvovec-rzyl in the requested eye(s)? Yes No			
SUPPORTING INFORMATION			
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Prescribing Provider Signature		Date	