SENTARA COMMUNITY PLAN (MEDICAID)

MEDICAL PRIOR AUTHORIZATION/STEP-EDIT REQUEST*

Directions: The prescribing physician must sign and clearly print name (preprinted stamps not valid) on this request. All other information may be filled in by office staff; <u>fax to 1-844-305-2331</u>. No additional phone calls will be necessary if all information (including phone and fax #s) on this form is correct. If information provided is not complete, correct, or legible, authorization can be delayed.

Drug Requested: Luxturna[™] (voretigen neparvovec-rzy) Subretinal Injection (Medical) (J-3398)

Luxturna[™] should <u>ONLY</u> be administered in a surgical suite under controlled aseptic conditions by a surgeon experienced in performing intraocular surgery.

MEMBER & PRESCRIBER INFORMATION: Authorization may be delayed if incomplete.

Member Name:	
Member Sentara #:	
Prescriber Name:	
Prescriber Signature:	
Office Contact Name:	
Phone Number:	Fax Number:
DEA OR NPI #:	
DRUG INFORMATION: Authorization may be delayed if incomplete.	
Drug Form/Strength:	
Dosing Schedule:	Length of Therapy:
Diagnosis:	ICD Code, if applicable:
Weight:	Date:

□ Standard Review. In checking this box, the timeframe does not jeopardize the life or health of the member or the member's ability to regain maximum function and would not subject the member to severe pain.

CLINICAL CRITERIA: Check below all that apply. All criteria must be met for approval. To support each line checked, all documentation, including lab results, diagnostics, and/or chart notes, must be provided or request may be denied.

Initial Authorization: One dose per eye/per lifetime. Renewal is NOT authorized.

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□ Has member received LuxturnaTM in the past?

□ Yes □ No

- **Yes**, in the same eye (**Provide details and progress notes with date and time stamp.**)
- □ Yes, but in the other eye only (Provide details and progress notes with date and time stamp.)
- D No
- **Unknown**

MEDICAL INFORMATION. Supportive documentation must be provided for all answers below or request may be delayed or denied.

 1. Does member have vision loss due to inherited retinal dystrophy?
 □ Yes □ No

 AND

2. Did member undergo genetic testing?

 $\Box \quad Yes - include a copy of the test results \qquad \Box \quad No - denial of drug$

If YES:

a) Did genetic testing confirm biallelic mutation of the RPE65 gene (two confirmed pathogenic or likely pathogenic loss of function variants)?

AND

b) Indicate which of the following genetic test was performed to confirm biallelic pathogenic and/or likely pathogenic RPE65:

□ Single-gene test □ None listed

Provide the date of the test:

3. Are the RPE65 gene mutations classifications based on the ACMG standards and guidelines for the interpretation of sequence variants (2015)? □ Yes □ No

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4. Does member have sufficiently viable retinal cells as determined by optical coherence tomography (OCT) and/or ophthalmoscopy?
 □ Yes □ No (denial of drug)

If **YES**, does the optical coherence tomography (OCT) and/or ophthalmoscopy show any of the following? (Check and provide documentation of at least one of the below.)

□ Area of retina within the posterior pole of greater than 100 µm thickness per OCT

OR

□ At least 3 disc areas of retina without atrophy or pigmentary degeneration within the posterior pole

AND

□ Visual field within 20 degrees in any meridian as measured by III4e isopter **OR** equivalent in both eyes

OR

□ Visual acuity worse than 20/60 in **both** eyes

Medication being provided by (check applicable box(es) below):
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Physician's office OR Specialty Pharmacy – PropriumRx

For urgent reviews: Practitioner should call Sentara Health Pre-Authorization Department if they believe a standard review would subject the member to adverse health consequences. Sentara Health's definition of urgent is a lack of treatment that could seriously jeopardize the life or health of the member or the member's ability to regain maximum function.

******Use of samples to initiate therapy does not meet step edit/ preauthorization criteria. ******

Previous therapies will be verified through pharmacy paid claims or submitted chart notes.