## Louisiana Medicaid Delandistrogene Moxeparvovec-rokl (Elevidys Medicaid Polys Med

The *Louisiana Uniform Prescription Drug Prior Authorization Form* should be utilized to request clinical authorization for delandistrogene moxeparvovec-rokl (Elevidys M.).

Additional Point-of-Sale edits may apply.

By submitting the authorization request, the prescriber attests to the conditions available HERE.

This The indication for non-ambulatory patients is approved under accelerated approval based on expression of Elevidys  $\mathbb{R}^{TM}$  micro-dystrophin in skeletal muscle-observed in patients treated with Elevidys  $\mathbb{R}^{TM}$ . Continued approval for this indication may be contingent upon verification and description of clinical benefit in a confirmatory trial(s).

## **Approval Criteria**

- The recipient is 4 or 5 years of age or older on the date of the request; AND
- The recipient has a diagnosis of Duchenne muscular dystrophy (DMD) confirmed by genetic testing; **AND**
- The recipient had baseline laboratory tests demonstrating rAAVrh74 antibody titers < 1:400 as determined by ELISA binding immunoassay; **AND**
- This medication is prescribed by a neurologist; AND
- The following are true and stated on the request:
  - o Elevidys RTM is not prescribed concurrently with exon skipping therapies; AND
  - o The recipient has been receiving oral corticosteroid therapy for DMD; AND
  - The recipient will continue to receive oral corticosteroid therapy, unless contraindicated or clinically significant adverse effects are experienced; AND
  - o The recipient has never received a dose of Elevidys@; AND
  - The recipient does **NOT** have any deletion in exon 8 and/or exon 9 in the DMD gene;
     .AND
  - The recipient has ambulatory function.

## Duration of approval: 6 months – allow 1 dose per lifetime

## References

ClinicalTrials.gov. A Gene Transfer Therapy Study to Evaluate the Safety and Efficacy of Delandistrogene Moxeparvovec (SRP-9001) in Participants With Duchenne Muscular Dystrophy (DMD) (EMBARK). <a href="https://classic.clinicaltrials.gov/ct2/show/NCT05096221">https://classic.clinicaltrials.gov/ct2/show/NCT05096221</a>

ClinicalTrials.gov. A Gene Transfer Therapy Study to Evaluate the Safety of and Expression From Delandistrogene Moxeparvovec (SRP-9001) in Participants With Duchenne Muscular Dystrophy (DMD) (ENDEAVOR). https://classic.clinicaltrials.gov/ct2/show/NCT04626674

Elevidys (delandistrogene moxeparvovec-rokl) [package insert]. Cambridge, MA: Sarepta Therapeutics, Inc; October 2023 June 2024. <a href="https://www.elevidyshcp.com/pihttps://www.elevidys.com/PI">https://www.elevidyshcp.com/pihttps://www.elevidys.com/PI</a>

Revision / Date	<b>Implementation Date</b>
Policy created / July 2023	January 2024
Formatting changes, updated references / March 2024	July 2024
Expanded age to 4 years and older, removed ambulatory requirement, updated references / June 2024	January 2025