Prior Authorization Criteria



Elevidys® (delandistrogene moxeparvovec-rokl) PA Criteria

Evrysdi is indicated for the treatment of ambulatory pediatric patients aged 4 years of age and older with Duchenne muscular dystrophy (DMD) with a confirmed mutation in the DMD gene.

Provider must submit documentation (such as office chart notes, lab results or other clinical information) supporting that member has met all approval criteria.

Limitation: One single dose per lifetime. One kit based on patient weight.
\square Yes \square No Patient is 4 years of age or older;
AND
\square Yes \square No Patient has been diagnosed with Duchenne Muscular Dystrophy (DMD)
AND
\square Yes \square No $$ Patient has a confirmed mutation of the DMD gene between exons 18 to
exon 58;
\square Yes \square No The mutation of the DMD gene is NOT a deletion in exon 8 or exon 9;
AND
\square Yes \square No Patient must have a baseline anti-AArh74 total binding antibody titer of <
1:400 as measured by ELISA;
AND
\square Yes \square No $$ Patient is ambulatory, and provider will attest.
AND
\square Yes \square No Patient is receiving physical and/or occupational therapy;
AND
\square Yes \square No $$ Patient is \textbf{NOT} on concomitant the rapy with DMD-directed antisense
oligonucleotides (e.g. golodirsen, casimersen, viltolarsen, eteplirsen, etc.);
AND

\square Yes \square No Patient has NOT received a DMD-directed antisense oligonucleotides
within the past 30 days;
AND
\square Yes \square No Patient does NOT have an active infection, including clinically important
localized infections;
AND
\square Yes \square No $$ Patient has been on a stable dose of a corticosteroid, unless
contraindicated or intolerance, prior to the start of therapy and will be used
concomitantly with a corticosteroid regimen pre and post-infusion (refer to the
package insert for recommended corticosteroid dosing during therapy);
AND
\square Yes \square No $$ Patient's troponin-1 levels will be monitored at baseline, weekly for the
first month, and subsequently as clinically indicated;
AND
\square Yes \square No Patient's platelet counts will be monitored at baseline, weekly for the first
two weeks, and subsequently as clinically indicated;
AND
\square Yes \square No $$ Patient will have laboratory liver assessments performed weekly prior to
and following therapy for the first 3 months, and as indicated.
See Package Insert for specific details on Contraindications/Warnings/Precautions
Dosing The recommended dose is $1.33\times10^{14}\mathrm{vector}$ genomes per kilogram (vg/kg) of body weight (or 10 mL/kg body weight).
Calculate the dose as follows: Elevidys dose (in mL) = patient body weight (in kilogram) \times 10
The multiplication factor 10 represents the per kilogram dose (1.33 \times 1014 vg/kg) divided by the amount of vector genome copies per mL of the Elevidys suspension (1.33 \times 1013 vg/mL).
Number of vials needed = Elevidys dose (in mL) divided by 10 (round to the nearest

number of vials).