

Division: Pharmacy Policy	Subject: Prior Authorization Criteria
Original Development Date: Original Effective Date:	February 26, 2021
Revision Date:	March 17, 2021, February 1, 2022, April 22, 2022, April 14, 2023

# VILTEPSO<sup>TM</sup> (viltolarsen)

## LENGTH OF AUTHORIZATION: SIX MONTHS

#### **REVIEW CRITERIA:**

- Patient must have the diagnosis of Duchenne Muscular Dystrophy (DMD).
- Submission of medical records (e.g., chart notes, laboratory values) as genetic test is required to confirm that a patient's mutation of the DMD gene is amenable to exon 53 skipping.
- Medication is prescribed by or in consultation with a neurologist or a physician who specializes in treatment of DMD (e.g., pediatric neurologist, cardiologist, or pulmonary specialist).
- Patient has been on stable dose of oral corticosteroids for at least 24 weeks prior to starting therapy unless contraindicated or intolerant.
- Patient is not concurrently treated with other DMD antisense oligonucleotides (e.g. golodirsen, casimersen, or eteplirsen).
- If the patient is ambulatory, functional level determination of baseline assessment of ambulatory function (six-minute walk test (6MWT), time to run/walk 10-meter test (TTRW), time to climb 4-stair test (TTCLIMB), time to stand (TTSTAND) or North Star Ambulatory Assessment (NSAA)) is required.
- If not ambulatory, patient must have a Brooke Upper Extremity Function Scale of five or less documented OR a Forced Vital Capacity of 30% or more.

#### **CONTINUATION OF THERAPY:**

- Patient met initial review criteria.
- Documentation of improvement, maintenance or slowing of disease progression:
  - o For ambulatory patients submission of 6MWT, TTRW, TTCLIMB, TTSTAND or NSAA.
  - For non-ambulatory patients submission of Brooke Upper Extremity Function Scale (five or less) documented OR a Forced Vital Capacity document (30% or more)
- Patient is not concurrently treated with other DMD antisense oligonucleotides (e.g. golodirsen, casimersen, or eteplirsen).

### DOSING AND ADMINISTRATION:

- Refer to product labeling at <a href="https://www.accessdata.fda.gov/scripts/cder/daf/">https://www.accessdata.fda.gov/scripts/cder/daf/</a>
- Available as 250 mg/5 mL (50 mg/mL) single-dose vial

