

Prior Authorization Criteria
Duchenne Muscular Dystrophy Antisense Oligonucleotides

All requests for Duchenne Muscular Dystrophy (DMD) Antisense Oligonucleotides require a prior authorization and will be screened for medical necessity and appropriateness using the criteria listed below.

DMD antisense oligonucleotides include Exondys 51 (eteplirsen), Vyondys 53 (golodirsen), Viltepso (viltolarsen) and Amondys 45 (casimersen). New products with this classification will require the same documentation.

Coverage may be provided with a diagnosis of Duchenne Muscular Dystrophy (DMD) and all of the following criteria is met:

- A confirmed diagnosis of DMD by submission of lab testing demonstrating mutation of the dystrophin gene amenable to exon skipping of the applicable target exon
- The member will receive concurrent corticosteroids unless contraindicated or intolerant
- Must be prescribed by or in consultation with a neurologist who has experience in the treatment and management of DMD
- There is documentation of a baseline evaluation, including a standardized assessment of motor function, by a neurologist with experience treating DMD;
- The requested dose and frequency is in accordance with FDA-approved labeling, nationally recognized compendia, and/or evidence-based practice guidelines
- Is age-appropriate according to FDA-approved package labeling, nationally recognized compendia, or peer-reviewed medical literature
- Vyondys 53 (golodirsen) will not be used concomitantly with Viltepso (vitolarsen).
- **Initial Duration of Approval:** 6 months
- **Reauthorization criteria**
 - The member has documentation of an annual evaluation, including an assessment of motor function ability, by a neurologist who has experience in the treatment and management of DMD;
 - Based on the prescriber's assessment, the member continues to benefit from therapy;
 - Vyondys 53 (golodirsen) will not be used concomitantly with Viltepso (vitolarsen).
 - The member is receiving concurrent corticosteroids unless contraindicated or intolerant
- **Reauthorization Duration of Approval:** 6 months

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.

Attachments:

Table 1. DMD Medications and Target Exon

<u>Generic Name</u>	<u>Brand Name</u>	<u>Target Exon</u>
Eteplirsén	Exondys 51	51
Golodirsén	Vyondys 53	53
Viltolarsén	Viltepso	53
Casimersén	Amondys 45	45

Table 2. Brooke Upper Extremity Scale

Score	Description
1	Starting with arms at the sides, the patient can abduct the arms in a full circle until they touch above the head
2	Can raise arms above head only by flexing the elbow (shortening the circumference of the movement) or using accessory muscles
3	Cannot raise hands above head, but can raise an 8-oz glass of water to the mouth
4	Can raise hands to the mouth, but cannot raise an 8-oz glass of water to the mouth
5	Cannot raise hands to the mouth, but can use hands to hold a pen or pick up pennies from the table
6	Cannot raise hands to the mouth and has no useful function of hands

Duchenne Muscular Dystrophy Antisense Oligonucleotides 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 weight: Height:

REQUESTED DRUG INFORMATION

Medication:	Strength:
Directions:	Quantity: Refills:
Is the member currently receiving requested medication? <input type="checkbox"/> Yes <input type="checkbox"/> No Date Medication Initiated:	

Billing Information

This medication will be billed: <input type="checkbox"/> at a pharmacy OR <input type="checkbox"/> medically, JCODE:
Place of Service: <input type="checkbox"/> Hospital <input type="checkbox"/> Provider's office <input type="checkbox"/> Member's home <input type="checkbox"/> Other

Place of Service Information

Name:	NPI:
Address:	Phone:

MEDICAL HISTORY (Complete for ALL requests)

Diagnosis:	ICD-10:
Is there lab testing demonstrating the member has a mutation of the dystrophin gene amenable to exon skipping? <input type="checkbox"/> Yes <input type="checkbox"/> No If Yes, Which Exon is amenable?	
Will the member be using concurrent corticosteroids? <input type="checkbox"/> Yes <input type="checkbox"/> No	
If no, please explain:	
Is a baseline evaluation including baseline motor function testing included with the request? <input type="checkbox"/> Yes <input type="checkbox"/> No	
Is Vyondys 53 being used concomitantly with Viltepso? <input type="checkbox"/> Yes <input type="checkbox"/> No	

CURRENT or PREVIOUS THERAPY

Medication Name	Strength/ Frequency	Dates of Therapy	Status (Discontinued & Why / Current)

REAUTHORIZATION

Has the member experienced a clinical benefit with treatment? <input type="checkbox"/> Yes <input type="checkbox"/> No
Is an annual evaluation including motor function testing included with the request? <input type="checkbox"/> Yes (documentation attached) <input type="checkbox"/> No
Is Vyondys 53 being used concomitantly with Viltepso? <input type="checkbox"/> Yes <input type="checkbox"/> No
Is the member receiving concurrent corticosteroids unless contraindicated or intolerant? <input type="checkbox"/> Yes <input type="checkbox"/> No

SUPPORTING INFORMATION or CLINICAL RATIONALE

Prescribing Provider Signature

Date

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