



## Prior Authorization Criteria

MISSISSIPPI DIVISION OF  
**MEDICAID**

### **eteplirsen (Exondys 51™) CRITERIA\***

Diagnosis of Duchenne Muscular Dystrophy (DMD) with mutation amenable to exon 51 skipping; ICD-10 code: \_\_\_\_\_

For both the initial and the reauthorization requests, providers must submit documentation of a recent clinical progress note including patient's current age and weight with the date the weight was obtained; the weight must be dated no more than 30 days before the request date, a copy of the prescription, and the eteplirsen (Exondys 51™) prior authorization form.

INITIAL AUTHORIZATION requests shall also include documentation of:

- the genetic laboratory test result with specific mutation and
- physical function with age appropriate testing tools used to measure physical function.

REAUTHORIZATION requests shall also include documentation of either:

- an increase in the physical function from baseline or
- that baseline physical function has been maintained.

Providers should use the same testing instrument as used in the baseline evaluation for physical function. If re-use of the initial testing instrument is not appropriate, for example, due to change in client status or restricted age range of the testing tool, the provider must explain the reason for the change.

Prior authorization requests for eteplirsen (Exondys 51™) may be approved if the following criteria are met: (*Yes should be checked for each statement*):

**INITIAL AUTHORIZATION: (prior authorization will be issued for, no more than, a six-month duration):**

- Yes  No Males age  $\geq$  3 years,
- Yes  No Patient has a diagnosis of Duchenne muscular dystrophy (DMD) with mutation amenable to exon 51 skipping **confirmed** by genetic testing (attach results of genetic testing),
- Yes  No eteplirsen (Exondys 51™) is prescribed by or in consultation with a neurologist or a physician who specializes in treatment of DMD; (i.e. pediatric neurologist, cardiologist or pulmonary specialist),
- Yes  No Patient is on a stable dose of a corticosteroid or has documented reason not to be on this medication,
- Yes  No Patient is not ventilator dependent,
- Yes  No Patient is not in medically intractable congestive heart failure,

Yes  No Comprehensive progress notes submitted document age appropriate and functional level determination of baseline assessment. Testing tools that can be used to demonstrate physical function include, but are not limited to:

- Baseline 6-Minute Walk Test (6MWT)],
  - **OR**
- Brooke Upper Extremity Scale (some useful hand function present for use of adaptive technology),
  - **OR**
- Forced Vital Capacity assessment,

Yes  No Request is for FDA approved dosing,

Yes  No Copy of prescription is provided,

Yes  No Patient's weight in kilograms and the date weight recorded.

Dates of Weight/Visit: \_\_\_\_\_ Weight (in kg) \_\_\_\_\_

**REAUTHORIZATION: (prior authorization will be issued for, no more than, a six-month duration):**

Yes  No Patient initial authorization criteria for initial treatment are still met,

Yes  No Patient has not had significant decline in neuromotor function or forced vital capacity while on eteplirsen (Exondys 51™),

Yes  No Statement from prescribing physician that the patient has been compliant with treatment,

Yes  No Copy of prescription is provided,

Yes  No Patient's weight in kilograms and the date weight recorded.

Dates of Weight/Visit: \_\_\_\_\_ Weight (in kg) \_\_\_\_\_

Each prescription may be written for 6 months; however the prescription must specify that medication is to be dispensed at no more than four (4) weeks supply at a time. Mississippi Division of Medicaid will only allow a 28 day supply to be dispensed at a time to ensure that the patient tolerates the medication prior to subsequent medication doses being dispensed to the physician clinic/hospital for administration.

*\*NOTE: Mississippi Division of Medicaid's coverage of eteplirsen (Exondys 51™) is contingent upon continued FDA approval.*

Update Effective 2/3/2020 V3