## **AvMed**

## PHARMACY PRIOR AUTHORIZATION/STEP-EDIT REQUEST\*

<u>Directions</u>: <u>The prescribing physician must sign and clearly print name (preprinted stamps not valid)</u> on this request. All other information may be filled in by office staff; <u>fax to 1-305-671-0200</u>. No additional phone calls will be necessary if all information (<u>including phone and fax #s</u>) on this form is correct. <u>If the information provided is not complete, correct, or legible, the authorization process can be delayed.</u>

<u>Drug Requested</u>: Firdapse<sup>®</sup> (amifampridine phosphate)

MEMBER & PRESCRIBER INFORMATION	: Authorization may be delayed if incomplete.		
Member Name:			
Member Sentara #:	Date of Birth:		
Prescriber Name:			
Prescriber Signature:			
Office Contact Name:			
Phone Number:			
NPI #:			
DRUG INFORMATION: Authorization may be del	ayed if incomplete.		
Drug Name/Form/Strength:			
Dosing Schedule:	Length of Therapy:		
Diagnosis:	ICD Code, if applicable:		
ight (if applicable): Date weight obtained:			

## **Recommended Dosage:**

Age and body weight	Initial daily dosage	Titration regimen	Maximum single dose	Maximum total daily maintenance dosage
<ul> <li>Adults (any weight)</li> <li>Pediatric patients weighing 45 kg or more</li> </ul>	15 mg to 30 mg daily, in 3 to 5 divided doses	Increase total daily dosage by 5 mg every 3 or 4 days	20 mg	100 mg given in divided doses
<ul> <li>Pediatric patients weighing less than 45 kg</li> </ul>	5 mg to 15 mg daily, in 3 to 5 divided doses	Increase total daily dosage by 2.5 mg every 3 or 4 days	10 mg	50 mg given in divided doses

**Quantity Limit:** 300 tablets per 30 days

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**CLINICAL CRITERIA**: Check below all that apply. All criteria must be met for approval. To support each line checked, all documentation, including lab results, diagnostics, and/or chart notes, must be provided or request may be denied.

Initia	al Authorization: 6 months
	Medication must be prescribed by or in consultation with a neurologist
	Member must be 6 years of age or older
	Member must have a diagnosis of Lambert-Eaton myasthenic syndrome
	Lambert-Eaton myasthenic syndrome diagnosis has been confirmed by <b>ONE</b> of the following ( <b>must submit labs for documentation</b> ):
	<ul> <li>□ Presence of anti-P/Q-type voltage-gated calcium channel (VGCC) antibodies</li> <li>□ A confirmatory electrodiagnostic study [e.g., repetitive nerve stimulation (RNS), needle electromyography (EMG), single-fiber electromyography (SFEMG)]</li> </ul>
	Provider must submit chart notes documenting moderate to severe muscle weakness that interferes with function
	Provider attests other differential diagnoses such as Myasthenia gravis have been ruled out
	Provider attests the member does <u>NOT</u> have a history of seizures or take medications that lower the seizure threshold (e.g., bupropion, tramadol, amphetamines, theophylline)
	Provider attests the member is <b>NOT</b> using alcohol
	Member is <u>NOT</u> receiving Firdapse <sup>®</sup> in combination with similar potassium channel blockers, such as Ampyra <sup>®</sup> (dalfampridine), or used in combination with compounded formulation of 3,4 diaminopyridin
	Provider must submit a baseline assessment or chart notes documenting at least <b>ONE</b> of the following measures <b>(check all that apply)</b> :
	□ Dynamometry
	☐ Timed 25 Foot Walk test ☐ Timed Up and Go (TUG) test
	☐ Timed Up and Go (TUG) test
suppo	<b>athorization:</b> 12 months. Check below all that apply. All criteria must be met for approval. To ort each line checked, all documentation, including lab results, diagnostics, and/or chart notes, must be ded or request may be denied.
	Provider must submit chart notes of a positive clinical symptomatic response to Firdapse <sup>®</sup> therapy with improvement from the initial submitted baseline assessment in at least <u>ONE</u> of the following (check althat apply; current assessment must be submitted):

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☐ Timed 25 Foot Walk test (a quantitative mobility and leg function performance test based on a timed 25-foot walk; an average increase of more than 20% in the timed 25-foot walk may indicate a

□ Dynamometry

significant change in gait)

- ☐ Timed Up and Go (TUG) test (assesses patient's function, weakness and mobility. The test measures the time it takes for patients to rise from a chair, walk a short distance, return to their chair and climb stairs approximately three times; >30% time increase from baseline indicates deterioration)
  - 11–20 seconds is within normal limits for frail elderly and disabled patients
  - Greater than 20 seconds suggests the person needs assistance and indicates further examination and intervention may be required
  - 30 seconds or more suggests that the person may be prone to falls

## Medication being provided by Specialty Pharmacy - Proprium Rx

Not all drugs may be covered under every Plan.

If a drug is non-formulary on a Plan, documentation of medical necessity will be required.

\*\*Use of samples to initiate therapy does not meet step edit/ preauthorization criteria. \*\*

\*Previous therapies will be verified through pharmacy paid claims or submitted chart notes. \*