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 (including phone and fax #s) 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>: DOJOLVI[™] (triheptanoin)

MEMBER & PRESCRIBER IN	NFORMATION: Authorization may be delayed if incomplete.		
Member Name:			
Member AvMed #:	Date of Birth:		
Prescriber Name:			
Prescriber Signature:	Date:		
Office Contact Name:			
Phone Number:	none Number: Fax Number:		
DEA OR NPI #:			
DRUG INFORMATION: Author	orization may be delayed if incomplete.		
Drug Form/Strength:			
Dosing Schedule:	Length of Therapy:		
Diagnosis:	ICD Code, if applicable:		
Weight:	Date:		

RECOMMENDED DOSING:

- Caloic value of DOJOLVI = 8.3 kcal/mL
- Round the total daily dosage to the nearest whole number.
- Divide the total daily dosage into at least four approximately equal individual doses.

$$Total\ Daily\ Dose\ (__mL) = \frac{Patients\ DCI\ (__kcal)\ x\ Target\ __\ \%\ dose\ of\ DCI}{8.3\frac{kcal}{mL}\ of\ DOJOLVI}$$

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•	Initiate DOJOLVI at a total daily dosage of approximately 10% DCI divided into at least four times per
	day and increase to the recommended total daily dosage of up to 35% DCI over a period of 2 to 3 weeks

•	AGE:
•	Total DCI (KCAL):

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.

Initial Authorization Length: 6 months

- □ Patient must have a diagnosis of a long-chain fatty acid oxidation disorder (LC-FAOD) of either: CPTII, VLCAD, LCHAD, OR TFP/MTP AND confirmed by **two of the following** assessments:
 - □ FAOD Deficiency (please document VLCAD, LCHAD, CPTII, or TFP/MTP):

Diagnosis	Age & Date of assessment	FAOD Deficiency: (Please document: VLCAD, LCHAD, CPTII, MTP/TFP)	RESULTS FROM DIAGNOSIS (fill in or send the assessment)	Confirmed Diagnosis
☐ Tandem mass spectrometry (MS/MS)				Acylcarnitine analysis: elevations of acylcarnitines on a newborn blood spot or in plasma https://www.acmg.net/ACMG/Medical-Genetics-Practice-Resources/ACT_Sheets_and_Algorithms.aspx
Genetic Analysis			ACADVL, HADHA, HADHP, CPT2:	Splice variants or nonsense mutations were identified
☐ Enzyme assay (lymphocytes)				Low enzyme activity in cultured fibroblasts
☐ IVP assay				Elevations of long chain acyl CoA

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AND

	Patient must have severe LC-FAOD confirmed by a history of ≥ 1 of the following despite therapy: ($\geq 2X$ upper limit of age/gender-matched normal, or ≥ 500 units/L if age-matched reference not established)					
	Chronic elevated creatine kinase ($[CK] \ge 2$ times the upper limit of normal) with ≥ 2 major clinical events (e.g., hospitalizations, hypoglycemia, cardiomyopathy, and rhabdomyolysis); OR					
	□ Episodic elevated CK with reported muscle dysfunction (e.g., frequent muscle fatigue, exercise intolerance, limitation of exercise); OR					
	□ Highly elevated CK (\geq 4 times the upper limit of normal); OR					
	□ Frequent (\geq 3 within a year or \geq 5 within 2 years) severe major clinical events (e.g., hospitalizations, hypoglycemia, cardiomyopathy, and rhabdomyolysis); OR					
	□ Severe susceptibility to hypoglycemia after short periods of fasting (≥ 2 events within a year that require ongoing prophylactic management or recurrent symptomatic hypoglycemia requiring intervention ≥ 2 times per week); OR					
	☐ Evidence of functional cardiomyopathy (echocardiogram documenting poor ejection fraction);					
	AND					
	Patient is being followed by a clinical specialist knowledgeable in appropriate disease-related dietary management (e.g., medical geneticist, genetic metabolic disorders, or a physician with a board certification in nutrition);					
	AND					
□ Patient is practicing appropriate dietary measures for their age and specific disorder (high callow long-chain fatty acids, avoidance of fasting);						
	AND					
	Patient has tried and failed medium chain triglyceride and continue to have ONE of the following: \square elevated CK] ≥ 2 times the upper limit of normal					
	□ hospitalizations					
	hypoglycemia OP					
	□ cardiomyopathy, OR □ rhabdomyolysis					
	• •					
	AND					
	Patient is NOT taking a pancreatic lipase inhibitor (e.g., orlistat);					
	AND					
	Patient will NOT receive an additional medium chain triglyceride while taking triheptanoin.					

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Reauthorization Approval Length: 12 months. 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.

Patient must continue to meet the above criteria;			
AND			
Patient must demonstrate disease improvement and/or stabilization (e.g., cardiac function, exercise tolerance, reduction in major clinical events, including hospitalization) as evidenced by all of the following:			
☐ Creatinine kinase is within normal limits			
□ Normal glycemic control			
□ No documentation of recent hospitalization			
□ No evidence of muscle fatigue			
AND			
Patient does NOT experience serious treatment-related adverse effects (e.g., gastrointestinal effects).			

Medication being provided by Specialty Pharmacy - PropriumRx

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.