# SENTARA HEALTH PLANS

# 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 1-800-750-9692</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</u> complete, correct, or legible, the authorization process can be delayed.

# **Drug Requested:** Myalept<sup>®</sup> (metreleptin)

### **MEMBER & PRESCRIBER INFORMATION:** Authorization may be delayed if incomplete.

Member Name:			
Member Sentara #:			
Prescriber Name:			
Prescriber Signature:			
Office Contact Name:			
Phone Number:			
DEA OR NPI #:			
DRUG INFORMATION: Complete all information	tion below or authorization may be delayed.		
Drug Form/Strength:			
Dosing Schedule:	Length of Therapy:		
Diagnosis:	_ ICD Code, if applicable:		

**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.

# **INITIATION AND CONTINUATION OF TREATMENT – All boxes below must be checked to qualify.**

- □ Member has a leptin deficiency as defined as (a copy of fasting laboratory leptin assay results is required for approval):
  - □ <4.0 ng/mL fasting leptin for females
  - $\Box$  <3.0 ng/mL fasting leptin for males
- □ Member has a diagnosis of (choose indication):
  - □ Acquired generalized lipodystrophy
  - □ Congenital generalized lipodystrophy
- □ Member has a concurrent condition of:

(Continued on next page)

- Diabetes mellitus or insulin resistance and failed 30-day trial of (submit chart notes):
  - □ Metformin, total daily dose of: \_\_\_\_\_

#### AND

- □ High-dose insulin or insulin pump
- □ Hypertriglyceridemia and failed 30-day trial of (submit chart notes):
  - □ Low-fat diet and/or dietary restrictions

#### AND

□ Fenofibrate or fenofibrate derivative

#### OR

□ Niacin or omega-3 fatty acid

#### OR

□ Atorvastatin, simvastatin, pravastatin, rosuvastatin

#### OR

□ Other therapy of (please specify): \_\_\_\_\_

INITIATION OF TREATMENT (submit all labs)		<u>REAUTHORIZATION</u> (submit all labs)	
HbA1c%		HbA1c%	
Fasting glucose	mg/dL	Fasting glucose	mg/dL
Triglyceride	mg/dL	Triglyceride	mg/dL
Patient weight	kg	Patient weight	kg
		Has member experienced clinical improvement or metabolic stabilization while using this medication? (submit chart notes to verify response) • Yes • No	

If approved, response to initial treatment will be <u>assessed after 4 months</u>, then <u>quarterly</u> <u>reassessment</u> will be required for continued approval.

Medication being provided by Specialty Pharmacy - PropriumRx

\*\*<u>Use of samples to initiate therapy does not meet step edit/ preauthorization criteria.</u>\*\* \*<u>Previous therapies will be verified through pharmacy paid claims or submitted chart notes.</u>\*