# 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 (<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>

**Drug Requested:** Wakix<sup>®</sup> (pitolisant)

MEMBER & PRESCRIBER INFORM	MATION: Authorization may be delayed if incomplete.
Member Name:	
Member Sentara #:	Date of Birth:
Prescriber Name:	
Prescriber Signature:	
Office Contact Name:	
Phone Number:	Fax Number:
DEA OR NPI #:	
DRUG INFORMATION: Authorization	may be delayed if incomplete.
Drug Form/Strength:	
Dosing Schedule:	Length of Therapy:
Diagnosis:	ICD Code, if applicable:
Weight:	Date:

<u>Maximum Approved Daily Dosing</u>: Narcolepsy (excessive daytime sleepiness/cataplexy): Initial: 8.9 mg once daily for 1 week, then increase to 17.8 mg once daily for 1 week; may further increase dose based on response and tolerability during week 3 to a maximum dose of 35.6 mg once daily

• Wakix for narcolepsy with or without cataplexy will not be approved in conjunction with Sunosi or Xyrem/Xywav. Sentara considers the use of concomitant therapy with Wakix and Sunosi or Xyrem/Xywav to be experimental and investigational. Safety and efficacy of these combinations has not been established and will not be permitted. In the event a member has an active Sunosi or Xyrem/Xywav authorization on file, all subsequent requests for Wakix will not be approved.

**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.

☐ Member is at least 18 years old

<u>AND</u>

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Member does not have a history of any of the following: substance abuse, serious CV disorder, history
of QT prolongation, current use of medications that cause QT prolongation, hepatic or renal
abnormalities or a psychiatric disorder

### **AND**

☐ Member does not have a sleep-related breathing disorder or periodic limb movement disorder (polysomnography results must be submitted for documentation)

### **DIAGNOSIS:** Please check one of the applicable diagnoses below

- □ Excessive Daytime Sleepiness associated with Narcolepsy. 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.
  - ☐ Member has a diagnosis of excessive daytime sleepiness (EDS) associated with narcolepsy (MSLT confirming diagnosis of narcolepsy must be submitted)

#### **AND**

☐ Provider must submit the member's baseline Epworth Sleepiness Scale score (rating scale must be attached)

### <u>AND</u>

☐ Member has failed a 30-day trial of modafinil or armodafinil (verified by paid pharmacy claims)

#### AND

- ☐ Member has failed a 30-day trial of Sunosi\* (\*Sunosi requires prior authorization; trial will be verified by paid pharmacy claims)
- □ Narcolepsy with Cataplexy. 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.
  - ☐ Member has a diagnosis of narcolepsy with cataplexy (MSLT confirming diagnosis of narcolepsy and chart notes documenting the occurrence of more than one episode of cataplexy at baseline prior to treatment with Wakix must be submitted. If polysomnography required, please submit with MSLT).

#### **AND**

□ Provider must submit the member's baseline Epworth Sleepiness Scale score (rating scale must be attached)

#### AND

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PA Wakix (CORE) (Continued from previous page)

Member must have a <b>2-month trial</b> and failure of <b>ONE</b> of the following anti-cataplectic therapies
(verified by pharmacy paid claims; documentation of intolerance or treatment failure must be
submitted, unless use is contraindicated. Please attach clinical documentation citing
contraindication):
□ SSRI (i.e. fluoxetine)
☐ TCA (i.e. clomipramine, imipramine, desipramine or protriptyline)
□ SNRI (i.e. venlafaxine or duloxetine)

# 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. \*