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>(Pharmacy) 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: (select drug below)

□ tasimelteon (Hetlioz®) Capsules	☐ Hetlioz® (tasimelteon) Liquid
MEMBER & PRESCRIBER INFOR	MATION: Authorization may be delayed if incomplete.
Member Name:	<u> </u>
Member Sentara #:	
Prescriber Name:	
Prescriber Signature:	
Office Contact Name:	
	Fax Number:
DEA OR NPI #:	
DRUG INFORMATION: Authorization	n may be delayed if incomplete.
Drug Form/Strength:	
	Length of Therapy:
Diagnosis:	ICD Code, if applicable:
Weight:	Date:
Ouantity Limit: 30 capsules/30 days, or if <	≤28 kg: 0.7 mg/kg/dose once daily
	all that apply. All criteria must be met for approval. To including lab results, diagnostics, and/or chart notes, must be
☐ For diagnosis of non-24-hour-sleep	o-wake disorder
Draggrihad by or in congultation with a s	enecialist in clean disorders
□ Prescribed by or in consultation with a s□ Member is ≥ 18 years of age	specialist in sleep disorders
- > 6 1 1 1 1 1 2 2 2 4 1	-cleen-wake disorder
C	disorder such as sleep apnea or insomnia
•	t perception in both eyes (nonfunctioning retinas)

	Member has a history of contraindication or intolerance to melatonin or ramelteon (Rozerem®) therapy (please submit chart notes)
	OR
	Member has history of failure of at least 6 months of uninterrupted daily treatment with melatonin or ramelteon (Rozerem®). Failure is defined as inability to achieve entrainment, clinically meaningful or significant increases in nighttime sleep or decreases in daytime sleep.
	Dates of melatonin or ramelteon therapy:
	(Therapy with melatonin or ramelteon (Rozerem®) will be verified through pharmacy paid claims or submitted chart notes.)
suppo	NICAL CRITERIA: 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.
\Box F	or diagnosis of Smith-Magenis syndrome
	The provider is a specialist experienced in treating patients diagnosed with Smith-Magenis Syndrome [i.e., sleep specialist, developmental/behavioral provider]
	The provider confirms a diagnosis of Smith-Magenis Syndrome (SMS) by all of the following:
	□ Submission of documentation detailing symptomology confirming SMS, and not due to another medical diagnosis (i.e., trisomy 21, Williams syndrome, brachydactyly-intellectual deficit syndrome (del 2q37), Prader-Willi syndrome)
	□ Submission of the results from a genetic panel confirming a deletion at chromosome 17p11.2 OR variant involving RAI1
	□ Submission of detailed history, progress notes, and/or actigraphy focusing on pattern of sleep disturbances affecting the patient (quality, average sleep time)
	For Hetlioz LQ [™] , the patient is between 3 and 15 years of age and documentation of current weight and requested dose must be submitted and follow FDA-approved dosing guidelines
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.