

Division: Pharmacy Policy	Subject: State of Florida's Agency for Health Care Administration's
	Prior Authorization Criteria
Original Development Date: August 26, 2014	August 26, 2014
Original Effective Date:	
Revision Date:	August 5, 2021, January 13, 2022

HETLIOZ® (tasimelteon) capsules and HETLIOZ LQ™ (tasimelteon) oral suspension

LENGTH OF AUTHORIZATION: UP TO 6 MONTHS

INITIAL REVIEW CRITERIA (ALL OF THE FOLLOWING MUST BE TRUE):

- If seeking approval for **Hetlioz**[®] capsules
 - Patient must be \ge 18 years old.
 - Patient must have a diagnosis of Non-24-hour sleep-wake disorder ("non-24") documented in clinical notes or health conditions.

OR

- o Patient must be ≥16 years old
- o Patient has a diagnosis of Nighttime sleep disturbances in Smith-Magenis Syndrome (SMS)
- Patient must have confirmed deletion 17p11.2 (cytogenetic analysis or microarray) or RAI1 gene mutation is identified
- If seeking approval for Hetlioz LQTM oral suspension
 - o Patient must be 3 to 15 years old.
 - Patient must have a diagnosis of Nighttime sleep disturbances in Smith-Magenis Syndrome (SMS).
 - Patient must have confirmed deletion 17p11.2 (cytogenetic analysis or microarray) or RAI1 gene mutation is identified.
- Do NOT approve for insomnia

CONTINUATION OF THERAPY:

- Patient met initial review criteria.
- Documentation of positive clinical response.
- Dosing is appropriate as per labeling or is supported by compendia.

DOSING & ADMINISTRATION:

- Refer to product labeling at https://www.accessdata.fda.gov/scripts/cder/daf/
- Dosage Forms:
 - o 20 mg capsule (**Hetlioz**[®])
 - o 4 mg/mL oral suspension (Hetlioz LQTM)