



**Request for Prior Authorization
TASIMELTEON (HETLIOZ®)**

(PLEASE PRINT – ACCURACY IS IMPORTANT)

Non-24-Hour Sleep-Wake Disorder (Non-24)

Treatment failure with a preferred sedative/hypnotic-non-benzodiazepine agent:

Drug name & dose: _____ Trial dates: _____

Reason for failure: _____

Treatment failure with ramelteon (Rozerem®):

Trial dose: _____ Trial dates: _____

Reason for failure: _____

Possible drug interactions/conflicting drug therapies: _____

Smith-Magenis Syndrome (SMS)

Attach documentation of one of the following:

- Deletion of 17p11.2 (cytogenic analysis or microarray) RAI1 gene mutation

Treatment failure with at least one medication used for sleep disturbances:

Trial drug name & dose: _____ Trial dates: _____

Reason for failure: _____

Requests for continuation therapy:

Has patient's use of tasimelteon been continuous without gaps in treatment? Yes No

Has patient experienced a positive clinical response with tasimelteon therapy? Yes (describe below) No

Patient improvements with tasimelteon (HetlioZ®) therapy (include description):

- Entrainment: _____
- Significant increase in nighttime sleep: _____
- Significant decrease in daytime sleep: _____
- Nighttime sleep quality: _____
- Other: _____

Attach lab results and other documentation as necessary.

Prescriber signature (Must match prescriber listed above.)	Date of submission
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IMPORTANT NOTE: In evaluating requests for prior authorization the consultant will consider the treatment from the standpoint of medical necessity only. If approval of this request is granted, this does not indicate that the member continues to be eligible for Medicaid. It is the responsibility of the provider who initiates the request for prior authorization to establish by inspection of the member's Medicaid eligibility card and, if necessary by contact with the county Department of Human Services, that the member continues to be eligible for Medicaid.