



FAX Completed Form To 1.833.404.2392

Pharmacy Help Desk 1.800.460.8988

Prescriber Help Desk

Request for Prior Authorization SODIUM OXYBATE PRODUCTS

(PLEASE PRINT - ACCURACY IS IMPORTANT) 1.833.587.2012 IA Medicaid Member ID # Patient name DOB Patient address Provider NPI Prescriber name Phone Prescriber address Fax Pharmacy name Address Phone Prescriber must complete all information above. It must be legible, correct, and complete or form will be returned. Pharmacy NPI Pharmacy fax NDC Prior authorization is required for sodium oxybate containing products. Payment will be considered under the following conditions: 1) A diagnosis of cataplexy associated with narcolepsy verified by a recent sleep study (including a PSG, MSLT, and ESS) and previous trial and therapy failure at a therapeutic dose with one of the following tricyclic antidepressants: clomipramine, imipramine, or protriptyline; or 2) A diagnosis of excessive daytime sleepiness associated with narcolepsy verified by a recent sleep study (including a PSG, MSLT, and ESS) and previous trials and therapy failures at a therapeutic dose with a preferred amphetamine and nonamphetamine stimulant; and 3) Patient meets the FDA approved age; and 4) is prescribed within the FDA approved dosing; and 5) Patient and provider are enrolled in the Xyrem REMS Program; and 6) Patient has been instructed to not drink alcohol when using Xyrem®; and 7) Patient has been counseled regarding the potential for abuse and dependence and will be closely monitored for signs of abuse and dependence; and 8) Requests for patients with concurrent use of a sedative hypnotic or a semialdehyde dehydrogenase deficiency will not be considered; and 9) The presciber must review the patient's use of controlled substances on the lowa Prescription Monitoring Program website prior to requesting prior authorization. The required trials may be overridden when documented evidence is provided that use of these agents would be medically contraindicated. Non-Preferred ☐ Xyrem □ Xywav **Dosage Instructions** Strength Quantity **Days Supply** ☐ Cataplexy associated with Narcolepsy (Please provide results from a recent ESS, MSLT, and PSG) Trial of preferred tricyclic antidepressant drug: Drug Name & Dose: Trial Dates: Failure Reason: ☐ Excessive Daytime Sleepiness associated with Narcolepsy (Please provide results from a recent ESS, MSLT, and PSG) Trial of preferred amphetamine stimulant: Drug Name & Dose: ______ Trial Dates: _____ Failure Reason: Trial of preferred non-amphetamine stimulant: Drug Name & Dose: Trial dates: Failure Reason: Medical or contraindication reason to override trial requirements: ____ ☐ No Patient is enrolled in the Xyrem[®] REMS Program: Yes Patient has been counseled and will be closely monitored for signs of abuse: \(\text{Yes} \) No Patient has a semialdehyde dehydrogenase deficiency: Yes Patient has been instructed to not drink alcohol when using Xyrem®: Prescriber review of patient's controlled substances use on the lowa PMP website:

Yes Date Reviewed:

No Attach lab results and other documentation as necessary. Prescriber signature (Must match prescriber listed above.) Date of submission

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.