

Instructions:

This form is used by Kaiser Permanente and/or participating providers for coverage of **Xyrem (sodium oxybate)**. <u>Please complete all sections, incomplete forms will delay processing</u>. <u>Fax this form back to Kaiser Permanente within 24</u> <u>hours (fax: 1-866-331-2104)</u>. If you have any questions or concerns, please call 1-866-331-2103. **Requests will not be considered unless all sections are complete**.

KP-MAS Formulary can be found at: Pharmacy | Community Provider Portal | Kaiser Permanente

	1 – Patient Information	
Patient Name:	Kaiser Medical ID#:	Date of Birth:
-	2 – Prescriber Information	
Prescriber Name:	Specialty:	NPI:
Prescriber Address:		
Prescriber Phone #:	Prescriber Fax #:	
, , , ,	r referral number from Kaiser Permanente? der referral number here:	
3 – Pharmacy Information		
Pharmacy Name:	Pharmacy NPI:	
Pharmacy Phone #	Pharmacy Fax #:	
	4 – Drug Therapy Requested	
Drug 1: Name/Strength/Formulati	on:	
Drug 2: Name/Strength/Formulati	on:	
Sig:		
	5– Diagnosis/Clinical Criteria	
1. Is this request for initial o	r continuing therapy?	

Initial therapy
Continuing therapy, State date: ______

2. Indicate the patient's diagnosis for the requested medication:

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Clinical Criteria:

- Is the prescriber a Pulmonologist (Sleep Specialist) or Neurologist?
 □ No □ Yes
- Is the prescriber enrolled in the Xyrem Patient Success Program?
 □ No □ Yes
- Is the patient between 7 years to 65 years of age?
 □ No □ Yes
- Is the patient on any sedative-hypnotic agents, opioids, benzodiazepines, or alcohol?
 □ No □ Yes

Treatment of excessive daytime sleepiness in narcolepsy:

- 5. Has the patient had an adequate trial (≥2 months) of a preferred stimulant (methylphenidate, amphetamine salt combination, dextroamphetamine) AND modafinil/armodafinil, unless contraindicated?
 □ No □ Yes
- Has the patient had an adequate trial of Sunosi (≥2 months) AND Wakix (≥2 months), unless contraindicated?
 □ No □ Yes
- Has the patient had an adequate trial (≥2 months) of Xywav?
 □ No □ Yes

Treatment of cataplexy due to narcolepsy:

- 8. Has the patient had an adequate trial (≥2 months) of at least 2 of the following: TCAs, SSRI, or SNRI or there is a contraindication?
 □ No □ Yes
- 9. Has the patient had an adequate trial (≥2 months) of Wakix AND Xywav, unless contraindicated? □ No □ Yes

For continuation of therapy, please respond to <u>additional questions</u> below:

- Does the patient have documentation of positive clinical response to therapy?
 □ No □ Yes
- Does the patient continue to be under the care of a specialist?
 □ No □ Yes

7 – Prescriber Sign-Off

Additional Information –

- 1. Please submit chart notes/medical records for the patient that are applicable to this request.
- 2. If member has not tried preferred agent(s) please provide rationale/explanation and any additional supporting information that should be taken into consideration for the requested medication:

I certify that the information provided is accurate	. Supporting documentation is available for State audits.
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Prescriber Signature:

Date:

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