DEPARTMENT OF HEALTH SERVICES

Division of Medicaid Services F-01430 (01/2019)

STATE OF WISCONSIN

Wis. Admin. Code § DHS 107.10(2)

FORWARDHEALTH PRIOR AUTHORIZATION DRUG ATTACHMENT FOR XYREM

INSTRUCTIONS: Type or print clearly. Before completing this form, read the Prior Authorization Drug Attachment for Xyrem Instructions, F-01430A. Providers may refer to the Forms page of the ForwardHealth Portal at www.forwardhealth.wi.gov/WIPortal/Content/provider/forms/index.htm.spage for the completion instructions.

Pharmacy providers are required to have a completed Prior Authorization Drug Attachment for Xyrem form signed by the prescriber before submitting a prior authorization (PA) request on the Portal, by fax, or by mail. Providers may call Provider Services at 800-947-9627 with questions.

SECTION I – MEMBER INFORMATION						
Name – Member (Last, First, Middle Initial)						
1. Name – Wember (Last, First, Widdle Hillar)						
2. Member ID Number	3. Date of Birth – Member					
SECTION II – PRESCRIPTION INFORMATION	<u> </u>					
4. Drug Name	5. Drug Strength					
6. Date Prescription Written	7. Directions for Use					
8. Refills						
9. Name – Prescriber	10. National Provider Identifier – Prescriber					
11. Address – Prescriber (Street, City, State, Zip+4 Code)						
12. Phone Number – Prescriber						
12. Filone Number – Frescriber						
SECTION III – CLINICAL INFORMATION						
Note: A copy of the member's current medical records that support	rt a clinical correlation between the member's test results and the					
member's condition of narcolepsy with cataplexy or narcolepsy with	hout cataplexy needs to be submitted with the PA request,					
including the following:Test results and provider interpretation for the polysomnogram	(PSG) and multiple sleep latency test (MSLT)					
 For members with excessive daytime sleepiness (EDS), a copy of the Epworth sleepiness scale (ESS) questionnaire, 						
maintenance of wakefulness test (MWT), or MSLT	manatrata aliniaal improvament ingluding a deergage in actanlaw.					
or a decrease in the member's daytime sleepiness, supported	monstrate clinical improvement, including a decrease in cataplexy by an ESS, MWT, or MSLT					
13. Diagnosis Code and Description						
14. Does the member have narcolepsy with cataplexy?	☐ Yes ☐ No					
If you indicate in the appear helpy the estapley asymptoms asy	navianced by the member and how frequently they come					
If yes, indicate in the space below the cataplexy symptoms exp	beneficed by the member and now frequently they occur.					
15. Does the member have narcolepsy without cataplexy?	☐ Yes ☐ No					
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SECTION III – CLINICAL INFORMATION (Continued)				
16. Is the member 7 years of age or older?		Yes		No
17. Does the member have a succinic semialdehyde dehydrogenase deficiency?		Yes		No
 18. As required by the Xyrem Risk Evaluation and Mitigation Strategy (REMS) Program: Has the prescriber counseled the member on the contraindication between Xyrem and alcohol? 		Yes		No
Has the member agreed to be abstinent from alcohol while being treated with Xyrem?		Yes		No
19. Indicate which symptom(s) of narcolepsy Xyrem is being used to treat.				
☐ Cataplexy				
☐ Excessive Daytime Sleepiness				
Other				
20. Does the member have a history of substance abuse, addiction, or diversion?		Yes		No
21. Is the member taking any sedative hypnotics?		Yes		No
22. Is the member taking central nervous system (CNS) depressants (for example, anxiolytics, barbiturates, opioids)?		Yes		No
If yes, indicate the CNS depressants and daily doses.				
1				
2				
3				
Are any of the above listed CNS depressants contributing to the member's daytime sleepiness?		Yes		No
If no, indicate how the prescriber evaluated the CNS depressants and determined they are no daytime sleepiness.	t cont	ributing	to the	member's
23. Has the member had an overnight PSG sleep study followed by an MSLT?		Yes		No
24. Does the member have EDS that interferes with normal activities on a daily basis?		Yes		No
25. Has the member completed an ESS questionnaire, MWT, or MSLT?		Yes		No
 26. Has the prescriber ruled out or treated the member for each of the following potential causes of EDS? Other sleep disorders including sleep apnea 		Yes		No
 Chronic pain or illness that disrupts normal sleep patterns Mood disorders such as depression Caffeine or nicotine use causing poor quality of nighttime sleep 				
27. Has the member experienced an unsatisfactory therapeutic response or experienced a				
clinically significant adverse drug reaction with a stimulant?		Yes		No
If yes, list the stimulant and dose, specific details about the unsatisfactory therapeutic response drug reaction, and the approximate dates the stimulant was taken in the space provided.	se or o	clinically	/ signif	icant adverse

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SECTION III – CLINICAL INFORMATION (Continued)					
28. Does the member have a medical condition(s) preventing the use of a stimulant?		Yes		No	
If yes, list the medical condition(s) that prevents the use of a stimulant in the space provided.					
29. Is there a clinically significant drug interaction between another medication the member is					
taking and stimulants?		Yes		No	
If yes, list the medication(s) and interaction(s) in the space provided.					
30. Has the member experienced an unsatisfactory therapeutic response after the medication has been titrated to a maximum recommended daily dose or experienced a clinically					
significant adverse drug reaction with armodafinil or modafinil?		Yes		No	
If yes, list the drug and dose, specific details about the unsatisfactory therapeutic response or reaction, and the approximate dates armodafinil or modafinil or were taken in the space provid		ally signi	fican	t adverse drug	
reaction, and the approximate dates annough in or modelling of were taken in the space provide	cu.				
31. Does the member have a medical condition(s) preventing the use of armodafinil					
or modafinil?		Yes		No	
If yes, list the medical condition(s) that prevents the use of armodafinil or modafinil in the space	e pro	vided.			
32. Is there a clinically significant drug interaction between another medication the member is taking and armodafinil or modafinil?		Yes		No	
If yes, list the medication(s) and interaction(s) in the space provided.					
in yes, list the medication(s) and interaction(s) in the space provided.					
33. Has the member experienced an unsatisfactory therapeutic response or experienced a					
clinically significant adverse drug reaction with tricyclic antidepressant (TCA), selective		Yes		No	
serotonin reuptake inhibitor (SSRI), or serotonin norepinephrine reuptake inhibitor (SNRI)?				No	
If yes, list the TCA, SSRI, or SNRI and dose, specific details about the unsatisfactory therapeu adverse drug reaction, and the approximate dates the TCA, SSRI, or SNRI was taken in the specific details about the unsatisfactory therapeu				nically significant	
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SECTION IV – AUTHORIZED SIGNATURE		
34. SIGNATURE – Prescriber	35. Date Signed	
SECTION V – ADDITIONAL INFORMATION	1	

36. Include any additional information in the space below. Additional diagnostic and clinical information explaining the need for the drug requested may also be included here.