FORWARDHEALTH PRIOR AUTHORIZATION DRUG ATTACHMENT FOR MODAFINIL AND NUVIGIL

INSTRUCTIONS: Type or print clearly. Before completing this form, read the Prior Authorization Drug Attachment for Modafinil and Nuvigil Instructions, F-00079A. 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 Modafinil and Nuvigil form signed by the prescriber before calling the Specialized Transmission Approval Technology-Prior Authorization (STAT-PA) system or submitting a 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

1. Name – Member (Last, First, Middle Initial)

2. Member ID Number	3. Date of Birth – Member						
SECTION II – PRESCRIPTION INFORMATION							
4. Drug Name	5. Drug Strength						
6. Date Prescription Written	7. Directions for Use						
8. Refills	1						
9. Name – Prescriber		10. National Provider Identifier (NPI) – Prescriber					
				,	,		
11. Address – Prescriber (Street, City, State, Zip+4 Code)							
12. Phone Number – Prescriber							
SECTION III – CLINICAL INFORMATION							
13. Diagnosis Code and Description							
13. Diagnosis Code and Description							
SECTION III A – CLINICAL INFORMATION FOR OBSTRUCTIVE				1084			
	E SLEEP APNE						
14. Does the member have OSAHS?			Yes		No		
If yes, complete the remainder of Section III A.							
If no, proceed to Section III B, Element 18.							
15. Has the member had an overnight polysomnogram (PSG) slee							
Apnea-Hypopnea Index (AHI) greater than or equal to five events per hour?			Yes		No		
Indicate the member's AHI: events per hour.							
					Co	ntinuea	



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SECTION III A – CLINICAL INFORMATION FOR OSAHS (Continued)						
16. Is the member taking any other stimulants or related agents?		Yes		No		
17. Has the member tried continuous positive airway pressure (CPAP)?		Yes		No		
				-		
If requested by ForwardHealth, the provider is required to submit the test results and provider interpretation for the PSG. SECTION III B – CLINICAL INFORMATION FOR NARCOLEPSY						
18. Does the member have narcolepsy?		Yes		No		
If yes, complete the remainder of Section III B. If no, proceed to Section III C, Element 22.						
19. Has the member had an overnight PSG sleep study followed by a multiple sleep						
latency test (MSLT) that confirm the member has narcolepsy?		Yes		No		
If yes, provide responses to the following questions regarding the PSG and MSLT sleep studi	ies:					
PSG						
A. Was the member's total sleep time less than 360 minutes?		Yes		No		
B. Did the member experience significant sleep interruptions (for example, respiratory						
events, periodic leg movements)?		Yes		No		
C. Did the provider interpretation indicate the member had an adequate night's sleep?		Yes		No		
MSLT						
D. Was the MSLT conducted the morning after the overnight PSG?		Yes		No		
E. Was the average sleep latency for all naps greater than eight minutes?		Yes		No		
F. Indicate the number of sleep onset rapid eye movement periods (SOREMPs) the member achieved during the MSLT.			SC	DREMPs		
0 = No SOREMPs 1 = One SOREMP 2 = Two or more SOREMPs						
If requested by ForwardHealth, the provider is required to submit the test results and provider interpretation for the PSG and MSLT, along with medical record documentation supporting a clinical correlation between the test results and a diagnosis of narcolepsy.						
20. Is the member taking any sedative hypnotics?		Yes		No		
21. Is the member taking central nervous system (CNS) depressants (for example, anxiolytics,		Ň				
barbiturates, or 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 not contributing to the member's daytime sleepiness.						

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SECTION III C – CLINICAL INFORMATION FOR SHIFT WORK SLEEP DISORDER				
2. Does the member have shift work sleep disorder?		Yes		No
If yes, complete the remainder of Section III C. If no, proceed to Section III D, Element 27.				
3. Is the member a night-shift worker?		Yes		No
If yes, indicate the member's current employer and weekly work schedule.				
4. Is the member taking any sedative hypnotics?		Yes		No
5. Is the member taking CNS depressants (for example, anxiolytics, barbiturates, or opioids)?		Yes		No
If yes, indicate the CNS depressants and daily doses. 1				
2				
3				
3Are any of the above listed CNS depressants contributing to the member's daytime sleepiness?		Yes		No
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SECTION III D – CLINICAL INFORMATION FOR ADHD (Complete this section only for PA requests for modafinil, if applicable.) (Continued)							
30. Has the member previously taken atomoxetine and experienced an unsatisfactory therapeutic response or experienced a clinically significant adverse drug reaction?				Yes		No	
31. Does the member have a medical history of substance abuse disorder?			Yes		No		
32. Does the member have a serious risk of drug diversion?			Yes		No		
SECTION IV – AUTHORIZED SIGNATURE							
33. SIGNATURE – Prescriber		34. Date Signed					
SECTION V – FOR PHARMACY PROVIDERS USING STAT-PA							
		equested (Up to 365 Days)					
37. NPI							
38. Date of Service (mm/dd/ccyy) (For STAT-PA requests, the date of service may be up to 31 days in the future or up to 14 days in the past.)							
39. Place of Service							
40. Assigned PA Number							
41. Grant Date	42. Expiration Date		43. Number of Days Approved				
SECTION VI – ADDITIONAL INFORMATION							
44. 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.