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## COMMONWEALTH OF VIRGINIA DEPARTMENT OF MEDICAL ASSISTANCE SERVICES Service Authorization (SA) Form NARCOLEPSY MEDICATIONS

If the following information is not complete, correct, or legible, the SA process can be delayed.

Please use one form per member.

MEMBER INFORMATION				
Last Name:	First Name:			
Medicaid ID Number:	Date of Birth:			
	Weight in Kilograms:			
PRESCRIBER INFORMATION				
Last Name:	First Name:			
NPI Number:				
Phone Number:	Fax Number:			
DRUG INFORMATION				
Minimum age of 18 for the following medication	ons:			
☐ Armodafinil tablet (generic for Nuvigil®) 50 n	ng, 150 mg, 200 mg, 250 mg (QD)			
☐ Modafinil (generic for Provigil®) 100 mg, 200	mg (QD or BID)			
☐ Nuvigil® 50 mg, 150 mg, 200 mg, 250 mg (QD)				
Provigil® 100 mg, 200 mg (QD or BID)				
Sunosi™ (solriamfetol) 75 mg, 150 mg				
Wakix® (pitolisant) 4.45 mg, 17.8 mg				
Drug Name/Form:				
Strength:				
Dosing Frequency:				
Length of Therapy:				
Quantity per Day:				
. ,				

Virginia DMAS SA Form: Narcolepsy Medications

Member's Last Name:	Member's First Name:				
DIAGNOSIS AND MEDICAL INFORMATION					
Please select diagnosis from the following:					
☐ Narcolepsy (sleep study must be attached)					
Excessive daytime sleepiness (EDS) in adult	members with narcolepsy				
Obstructive sleep apnea (sleep study must	be attached)				
Sudden onset of weak or paralyzed muscle	s (cataplexy)				
Shift work sleep disorder:					
Current shift schedule:					
Does not occur during the course of a	another sleep disorder or mental disorder				
Is not due to the direct physiological	effects of a medication or a general medical condition				
Other:					
list about a suitable and and and and					
List pharmaceutical agents attempted and ou	tcome:				
Medical Necessity (Provide clinical evidence the provide clinical rationale for quantity exception	nat the preferred agent(s) will not provide adequate benefit or n requests):				
(Form continued on next page.)					

Virginia DMAS SA Form: Narcolepsy Medications

Me	Member's Last Name: Men	nber's First Name:
No	Non-Preferred Medications	
Fo	For Wakix® (pitolisant):	
1.	Does the member have an International Classification of Statistical Manual of Mental Disorders, Fifth Edition (DS)	· · · · · · · · · · · · · · · · · · ·
	Yes No	
2.	<ol> <li>Does the member have a baseline daytime sleepiness as Sleepiness Scale, Stanford Sleepiness Scale, Karolinska S Questionnaire, or a Visual Analog Scale)? AND</li> </ol>	
	Yes No	
3.	<ol> <li>A mean sleep latency of ≤ 8 minutes AND ≥ 2 sleep onse sleep latency test (MSLT) performed according to stands sleep onset] on the preceding nocturnal polysomnogram AND</li> </ol>	ard techniques (A SOREMP [within 15 minutes of
	Yes No	
4.	4. Either cerebrospinal fluid (CSF) hypocretin-1 concentrate concentration measured by immunoreactivity is either a normal subjects with the same standardized assay; <b>AND</b>	110 pg/mL OR > 1/3 of mean values obtained in
	Yes No	
5.	<ol> <li>The hypersomnolence and/or MSLT findings are not bet sleep, obstructive sleep apnea, delayed sleep phase disc their withdrawal; AND</li> </ol>	•
	Yes No	
6.	<ol> <li>Patient has daily periods of irrepressible need to sleep of months; AND</li> </ol>	r daytime lapses into sleep occurring for ≥ 3
	Yes No	
7.	7. Patient must not be receiving treatment with sedative has zaleplon, benzodiazepines, barbiturates); <b>AND</b>	ypnotic agents (e.g., zolpidem, eszopiclone,
	Yes No	
8.	8. Patient will not use drugs that prolong the QT interval (e amiodarone, sotalol, ziprasidone, chlorpromazine, thior	
/Fr	(Form continued on next page.)	
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Virginia DMAS SA Form: Narcolepsy Medications

Me	mber's Last Name:	Member's First Name:			
9.	Patient will not use histamine-1 (H1) receptor antagonists (e.g., pheniramine maleate, diphenhydramin promethazine, imipramine, clomipramine, mirtazapine) concomitantly; <b>AND</b>				
	Yes No				
10	10. Patient does not have a history of prolonged QTc interval (e.g., QTc interval > 450 milliseconds); ANI				
	Yes No				
11. Therapy will not be used in patients with severe hepatic impairment (Child-Pugh C); AND					
	Yes No				
12	12. Patient does not have end stage renal disease (ESRD) (e.g., eGFR < 15 mL/minute/1.73 m <sup>2</sup> ).				
	Yes No				
For	brand Nuvigil or Provigil:				
1.	Has the member tried and failed the preferred generics for the requested products?				
	Yes No				
For	Renewal:				
1.	Does the member continue to meet initial criteria? AND				
	Yes No				
2.	ore-treatment baseline? AND				
	Yes No				
3.	Has the member not experienced andy treatment	related adverse effects?			
	☐ Yes ☐ No				
Pre	escriber Signature (Required)		Date		
•	signature, the Physician confirms the above inform diverifiable by member records.	ation is accurate			
Suk	ase include ALL requested information; Incompletomission of documentation does NOT guarantee covorices.		•		

Fax this form to 1-866-940-7328

Pharmacy PA call center: 1-800-310-6826