

OptumRx has partnered with CoverMyMeds to receive prior authorization requests, saving you time and often delivering real-time determinations.

Visit <u>go.covermymeds.com/OptumRx</u> to begin using this free service. Please note: All information below is required to process this request. Mon-Fri: 5am to 10pm Pacific / Sat: 6am to 3pm Pacific

Provigil® (modafinil) Prior Authorization Request Form (Page 1 of 2)

	DO NOT COPY FOR FU	TURE USE. FORMS A	RE UPDATED FREQUENTLY	AND MAY B	<u>E BARCODI</u>	<u> </u>	
Membe	er Information	Provider Information (required)					
Member Name:			Provider Name:				
Insurance ID#:			NPI#:		Specialty:		
Date of Birth:			Office Phone:				
Street Address:			Office Fax:				
City:	State:	Zip:	Office Street Address:	Street Address:			
Phone:			City:	State:		Zip:	
		Medication	Information (require	ed)			
Medication Name:			Strength: Dosage Form:				
☐ Check if requesting brand			Directions for Use:				
☐ Check if request is for continuation of therapy							
Clinical Information (required)							
depressive disorder or bipolar depression □ Ob□ Fatigue due to multiple sclerosis □ Sh□ Idiopathic hypersomnia			questions for that diagnosis: arcolepsy bstructive sleep apnea/hypopnea syndrome (OSAHS) nift work sleep disorder (SWSD) 10 Code(s):				
Does the patient hav		ailure, contraindicatio	order or bipolar depression, or intolerance to at least		ressants fro	om different	
If this is a reauthorization request, answer the following: Is there documentation of positive clinical response to Provigil (modafinil) therapy? Yes No							
Fatigue due to mult	iple sclerosis:	-					
For brand Provigil request, answer the following:							
Does the patient have a history of failure, contraindication, or intolerance to modafinil (generic Provigil)? Yes No							
If this is a reauthorization request, answer the following: Is the patient experiencing relief of fatigue with Provigil (modafinil) therapy? Yes No							
Idiopathic hyperson		been confirmed by a	aloon atudy? D Vac D N	٥			
Has the diagnosis of idiopathic hypersomnia been confirmed by a sleep study? Q Yes Q No If a sleep study has not been completed, please provide justification confirming why a sleep study would not be feasible:							
If this is a reauthorization request, answer the following:							
			odafinil) therapy? 🗖 Yes 🛚	□No			
Narcolepsy: Has the diagnosis of	narcolepsy been confirm	med by a sleep study	? □ Yes □ No				
If a sleep study has r	not been completed, ple	ase provide justificati	on confirming why a sleep	study would	not be feas	sible:	
 □ Amphetamine □ Amphetamine-dex □ Armodafinil □ Dexedrine □ Dextroamphetami □ Dextroamphetami If this is a reauthori 	ktroamphetamine ine ine extended-release (E zation request, answe	☐ Metadate EF☐ Methylin☐ Methylpheni☐ Methylpheni☐ Modafinil R) r the following:	date chewable tablet	□ R □ Si □ W □ Ze	unosi		

This document and others if attached contain information that is privileged, confidential and/or may contain protected health information (PHI). The Provider named above is required to safeguard PHI by applicable law. The information in this document is for the sole use of OptumRx. Proper consent to disclose PHI between these parties has been obtained. If you received this document by mistake, please know that sharing, copying, distributing or using information in this document is against the law. If you are not the intended recipient, please notify the sender immediately. Office use only: Provigil-Modafinil_CMS_2020Apr-W



Provigil® (modafinil) Prior Authorization Request Form (Page 2 of 2) DO NOT COPY FOR FUTURE USE. FORMS ARE UPDATED FREQUENTLY AND MAY BE BARCODED

Obstructive sleep apnea/hypopnea syndrome (OSAHS): Has the diagnosis of OSAHS been confirmed by a sleep study? ☐ Yes ☐ No						
If a sleep study has not been completed, please provide justification confirming why a sleep study would not be feasible:						
Was the diagnosis of OSAHS defined by 15 or more obstructive respiratory events per hour of sleep? Yes No						
Was the diagnosis of OSAHS defined by 5 or more obstructive respiratory events per hour of sleep AND one of these symptoms (unintentional sleep episodes during wakefulness, daytime sleepiness, unrefreshing sleep, fatigue, insomnia, waking up breath holding/gasping/choking, loud snoring or breathing interruptions during sleep)? Yes No						
Select the medications the patient has a failure, contraindication, or intolerance to: Armodafinil Sunosi						
☐ Armodafinil ☐ Modafinil ☐ Sunosi If this is a reauthorization request, answer the following:						
Is there documentation of positive clinical response to Provigil (modafinil) therapy? Yes No						
Shift work sleep disorder (SWSD):						
Has SWSD been confirmed by one of the following (select from the two options below)? Symptoms of excessive sleepiness or insomnia, for at least 3 months, which is associated with a work period (usually night work) that occurs during the normal sleep period.						
□ Sleep study demonstrating loss of a normal sleep-wake pattern (i.e., disturbed chronobiologic rhythmicity).						
Has it been confirmed that no other medical condition or medication accounts for the symptoms? ☐ Yes ☐ No						
Select the medications the patient has a failure, contraindication, or intolerance to:						
□ Armodafinil □ Modafinil						
If this is a reauthorization request, answer the following: Is there documentation of positive clinical response to Provigil (modafinil) therapy? Yes No						
Quantity limit requests: What is the quantity requested per DAY? What is the reason for exceeding the plan limitations? □ Titration or loading-dose purposes						
☐ Patient is on a dose-alternating schedule (e.g., one tablet in the morning and two tablets at night, one to two tablets at bedtime)						
 □ Requested strength/dose is not commercially available □ There is a medically necessary justification why the patient cannot use a higher commercially available strength to achieve the same dosage and remain within the same dosing frequency. Please specify: 						
Other:						
Are there any other comments, diagnoses, symptoms, medications tried or failed, and/or any other information the physician feels is important to this review?						
Please note: This request may be denied unless all required information is received. If the patient is not able to meet the above standard prior authorization requirements, please call 1-800-711-4555. For urgent or expedited requests please call 1-800-711-4555. This form may be used for non-urgent requests and faxed to 1-844-403-1028.						

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