STIMULANTS AND RELATED AGENTS PRIOR AUTHORIZATION FORM







(form effective 1/8/2024)

Fax to PerformRx $^{\text{SM}}$ at **1-855-851-4058**, or to speak to a representative, call **1-888-674-8720**.

I PRIOR AUTHORI	ZATION REQUEST	INFORMATION								
	enewal request	Total # of pages:								
Name of office contact:		Contact's phone number:			LTC	LTC facility contact/phone:				
PATIENT INFORM	1ATION									
Patient name:				Patient ID #:			DOB:			
Street address:			1							
Apt #:	City/state/zip:				Phone:					
PRESCRIBER INF	ORMATION									
Prescriber name:										
Specialty:			NPI:				State license #:			
Street address:				'			'			
Suite #:	City/state/zip:									
Phone:	Fax:									
CLINICAL INFOR	MATION		·				·			
Drug requested:										
Dosage form (tablet, ODT, suspension, etc.):						Quantity:		# months requested:		
Diagnosis (submit documentation):						Diagnosis	Diagnosis code (required):			
INITIAL REQUES	TS									
		lication within the past 90 da	ıys?			☐ Yes	Submit (Submit documentation		
					□ No	□ No				
For a non-preferred drug: Does the beneficiary have a history of trial and failure of or contraindication or intolerance to						☐ Yes	List preferred medications tried:			
the preferred drugs in this class that are approved or medically accepted for treatment of the beneficiary's condition? Refer to https://papdl.com/preferred-drug-list for a list of preferred and non-preferred drugs in this class.						□No	□ No			
Complete the sections below that are applicable to the beneficiary and this request and SUBMIT DOCUMENTATION for each item. For an analeptic Stimulants and Related Agents (e.g., Provigil, Nuvigil, Sunosi, Wakix) Is not receiving concurrent treatment with sedative/hypnotic medications Is receiving concurrent treatment with sedative/hypnotic medications reason: For the treatment of narcolepsy: Has a diagnosis of narcolepsy that is consistent with current International Classification of Sleep Disorders criteria (e.g., MSLT, overnight PSG, hypocretin-1 concentration, clinical assessment, etc.) For the treatment of shift work sleep disorder: Has a diagnosis of shift work sleep disorder that is consistent with current International Classification of Sleep Disorders criteria (e.g., shift work schedule, sleep log and actigraphy monitoring, other causes ruled out, clinical assessment, etc.) For the treatment of obstructive sleep apnea/hypopnea syndrome (OSAHS): Has a diagnosis of OSAHS that is consistent with current International Classification of Sleep Disorders criteria (e.g., overnight PSG, out-of-center sleep testing, associated medical or psychiatric disorders, clinical assessment, etc.) Tried and failed continuous positive airway pressure (CPAP) while adherent to treatment to resolve daytime sleepiness demonstrated by: Epworth Sleepiness Scale >10 Multiple sleep latency test (MSLT) <8 minutes Crannot use CPAP — reason: Tried and failed an oral appliance for OSAHS to resolve daytime sleepiness										
☐ For the treatment of ☐ Is currently receiv ☐ Is not receiving tr ☐ For a child <4 years ☐ Is prescribed the ☐ pediatric neu ☐ child/adolesc	f fatigue related to multipring treatment for MS eatment for MS — reason of age: requested medication ANE rologist				ith one of the followi	ng specialis	ts:			



INITIAL REQUESTS (continued)							
☐ For a beneficiary ≥18 years of age: ☐ For the treatment of ADHD: ☐ Has a diagnosis of ADHD that is consistent with current DSM criteria							
 For the treatment of narcolepsy: Has a diagnosis of narcolepsy consistent with current International Classification of Sleep Disorders criteria (e.g., MSLT, overnight PSG, CSF hypocretin-1 concentration, clinical assessment) For the treatment of binge eating disorder: Has a diagnosis of moderate to severe binge eating disorder that is consistent with the current DSM criteria Tried and failed (or cannot try) SSRIs (unless beneficiary has comorbid ADD or ADHD) 							
Tried and failed (or cannot try) topiramate (unless beneficiary has comorbid ADD or ADHD) Was referred for cognitive behavioral therapy or other psychotherapy For a stimulant agent: Was assessed for potential risk of misuse, abuse, and/or addiction based on family and social history Was educated regarding the potential adverse effects of stimulants, including the risk of misuse, abuse, and addiction Has documentation that the provider checked the PDMP for the beneficiary's controlled substance prescription history For a beneficiary with a history of substance dependency, abuse, or diversion: Has results of a recent UDS for licit and illicit drugs with the potential for abuse (including specific testing for oxycodone, fentanyl, and tramadol) that is consistent with prescribed controlled substances							
RENEWAL REQUESTS							
Has the beneficiary experienced a positive clinical response since starting the requested medication?	□ Yes	Submit documentation					
PLEASE FAX COMPLETED FORM WITH REQUIRED CLINICAL DOCUMENTATION							
Prescriber signature:		Date:					

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