

Request for Prior Authorization



Provider Help Desk I (844) 236-1464

CNS STIMULANTS AND ATOMOXETINE

FAX Completed Form To I (877) 733-3195

(PLEASE PRINT – ACCURACY IS IMPORTANT)

IA Medicaid Member ID #	Patient name	DOB			
Patient address					
Provider NPI	Prescriber name	Phone			
Prescriber address Fax					
Pharmacy name Address		Phone			
Prescriber must complete all information above. It must be legible, correct, and complete or form will be returned.					
Pharmacy NPI Pharmacy fax NDC		2			

Prior Authorization (PA) is required for CNS stimulants and atomoxetine for patients 21 years of age or older. Prior to requesting PA for any covered diagnosis, the prescriber must review the patient's use of controlled substances on the Iowa Prescription Monitoring Program (PMP) website. Request must adhere to all FDA approved labeling for requested drug and indication, including age, dosing, contraindications, warnings and precautions, drug interactions, and use in specific populations. Payment for CNS stimulants and atomoxetine will be considered when patient has an FDA approved or compendia indication for requested drug under the following conditions: I) Attention Deficit Hyperactivity Disorder (ADHD) meeting the DSM-5 criteria and confirmed by a standardized rating scale (such as Conners, Vanderbilt, Brown, Snap-IV). Symptoms must have been present before twelve (12) years of age and there must be clear evidence of clinically significant impairment in two or more current environments (social, academic, or occupational). Documentation of a recent clinical visit that confirms improvement in symptoms from baseline will be required for renewals or patients newly eligible that are established on medication to treat ADHD. Adults (\geq 21 years of age) are limited to the use of long-acting agents only. If a supplemental dose with a short-acting agent is needed for an adult in the mid to late afternoon, requests will be considered under the following circumstances: the dose of the long-acting agent has been optimized, documentation is provided a short-acting agent of the same chemical entity is medically necessary (e.g. employed during the day with school in the evening), and will be limited to one unit dose per day. Children (< 21 years of age) are limited to the use of longacting agents with one unit of a short acting agent per day. Use of an amphetamine agent plus a methylphenidate agent will not be considered for a diagnosis of ADHD. 2) Narcolepsy with diagnosis confirmed with a recent sleep study (ESS, MSLT, PSG). 3) Excessive sleepiness from obstructive sleep apnea/hypopnea syndrome (OSAHS) with documentation of non-pharmacological therapies tried (weight loss, position therapy, CPAP at maximum titration, BiPAP at maximum titration or surgery) and results from a recent sleep study (ESS, MSLT, PSG) with the diagnosis confirmed by a sleep specialist.

Payment for a non-preferred agent will be authorized only for cases in which there is documentation of previous trial and therapy failure with a preferred agent. * If a non-preferred long-acting medication is requested, a trial with the preferred extended release product of the same chemical entity (methylphenidate class) or chemically related agent (amphetamine class) is required. The required trials may be overridden when documented evidence is provided that the use of these agents would be medically contraindicated.

Requests for Vyvanse for Binge Eating Disorder must be submitted on the Binge Eating Disorder Agents PA form.

Preferred Non-Preferred Adderall Amphetamine Salt Combo Amphetamine ER Caps Adderall XR Armodafinil Adhansia XR* Atomoxetine Adzenys ER Susp Concerta Adzenys XR ODT Dexmethylphenidate ER Caps Amphetamine ER Suspension Dexmethylphenidate Tabs Amphetamine Sulfate Tabs Dextroamphetamine EE Caps Aptensio XR* Dextroamphetamine Tabs Azstarys Dyanavel XR Suspension Cotempla* Methylphenidate CD Caps Daytrana Methylphenidate IR Tabs Desoxyn Methylphenidate ER Tabs Dexedrine Methylphenidate LA Caps Dyanavel XR Chew Tab Methylphenidate Solution Evekeo Modafini Focalin **Ouillichew ER** Focalin XR Sunosi (step through armodafinil or modafinil) Iornay PM Methylin Solution

Methylphenidate Chew Methylphenidate TD Patch Methylphenidate ER 45mg, 63mg, 72mg Tabs Methylphenidate ER Caps* Methylphenidate XR Caps* Mydayis* 🗌 Nuvigil Procentra Provigil Quillivant XR Relexxii þ Ritalin Ritalin LA* Strattera Vyvanse ☐ Xelstrym

Request for Prior Authorization

CNS STIMULANTS AND ATOMOXETINE

(PLEASE PRINT – ACCURACY IS IMPORTANT)

Strength	Dosage Instructions	Quantity	Days Supply
Diagnosis:			
Attent	ion Deficit Hyperactivity Disorder (ADH	D)	
Did patient	have inattentive or hyperactive/impulsive symp	toms present prior to age	12? 🗌 Yes 🔲 No
Date of mos	st recent clinical visit confirming improvement	in symptoms from baseline	2:
Rating scale	used to determine diagnosis:		
Documenta	tion of clinically significant impairment in two c	or more current environr	nents (social, academic, or occupational).
Current Env	vironment & description:		
Current Env	vironment 2 & description:		
	or short-acting agents:		
Has dose of	long-acting agent been optimized? 🔲 Yes	🗌 No	
Adults: Prov	ride medical necessity for the addition of a sho	rt-acting agent:	
Children: Pr	ovide medical necessity for the need of more	than one unit of a short-ac	ting agent:
UNDERSTONE	Veight Loss CPAP Date: biPAP Date: urgery Date: ifics: psis confirmed by a sleep specialist? Yes r (specify)	sition therapy Maximum titration? Maximum titration?	
	view of patient's controlled substances u	se on the Iowa PMP we	ebsite:
🗌 No 🗌 Yes	Date Reviewed:		
Please documer reasons:	nt prior psychostimulant trial(s) and failures(s)	including drug name(s) str	ength, dose, exact date ranges and failure
Other - Please ranges:	e provide all pertinent medication trial(s) relati	ng to the diagnosis includi	ng drug name(s) strength, dose and exact date
Reason for use	of Non-Preferred drug requiring approval:		
Prescriber sig	nature (Must match prescriber listed above.)		Date of submission

IMPORTANT NOTE: In evaluating requests for prior authorization the consultant will consider the treatment from the standpoint of medical necessity only. If approval of this request is granted, this does not indicate that the member continues to be eligible for Medicaid. It is the responsibility of the provider who initiates the request for prior authorization to establish by inspection of the member's Medicaid eligibility card and, if necessary by contact with the county Department of Human Services, that the member continues to be eligible for Medicaid.