

Iowa Department of Human Services

FAX Completed Form To 1 (800) 574-2515

Provider Help Desk 1 (877) 776-1567

Request for Prior Authorization CNS STIMULANTS AND ATOMOXETINE

	(PLEASE PRINT – ACCURAC	Y 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	ation above. It must be legible, co	orrect, and complete or fo	rm will be returned.	
Pharmacy NPI	Pharmacy fax	NDC		
prescriber must review the patient's use of controlled substances on the lowa Prescription Monitoring Program (PMP) website. Payment for CNS stimulants and atomoxetine will be considered under the following conditions: 1) 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 (≥ 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 long-acting agents with one unit of a short acting agent per day. 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 medicati				
Requests for Vyvanse for Binge Eati			ler Agents PA form.	
Preferred Amphetamine Salt Combo Amphetamine ER Caps Aptensio XR Armodafinil Atomoxetine Concerta Daytrana Dexmethylphenidate Tabs Focalin XR Methylin Solution Methylphenidate IR Tabs Methylphenidate ER 20mg Tabs	Modafinil Quillichew ER Quillivant XR Vyvanse A C D D D D D D D D D D D D	Preferred dderall dderall XR* dzenys ER Susp dzenys XR ODT mphetamine Sulfate Tabs totempla* lesoxyn lexedrine* lexmethylphenidate ER cap lextroamphetamine ER Cap lyanavel XR vekeo ocalin		
Strength Dosage Ir	nstructions	Quantity	_Days Supply	

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Diagnosis:

Attention		
Age of patien	t at onset of symptoms:	_
Date of most	recent clinical visit confirming improvement in sympton	oms from baseline:
Rating scale (used to determine diagnosis:	
Documentation occupational)	on of clinically significant impairment in two or more c	urrent environments (social, academic, or
Current Envir	onment 1 & description:	
Current Envir	onment 2 & description:	
Requests for	short-acting agents for adults:	
Has dose of lo	ong-acting agent been optimized? Yes N	0
Provide medic	cal necessity for the addition of a short-acting agent:	
☐ Narcole	psy (Please provide results from a recent ESS, M	SLT, and PSG)
Have no We CP/ BiP Sur Specifi Diagnos Other (s Prescriber reviev No Yes Da Please document	AP Date: Maximur AP Date: Maximur	Yes If Yes, please indicate below: on therapy on titration? Yes No on titration? No on titration? Yes No on titration? Yes No on titration? Yes No on titration? Yes No
Other - Please p exact date ranges	rovide all pertinent medication trial(s) relating to the os:	liagnosis including drug name(s) strength, dose and
Reason for use of	Non-Preferred drug requiring approval:	
Prescriber signatur	re (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.

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