





Fax Completed Form To 1.833.404.2392 **Prescriber Help Desk** 1.833.587.2012

Online covermymeds.com/main/ prior-authorization-forms/

Request for Prior Authorization High Dose Opioids

(PLEASE PRINT - ACCURACY IS IMPORTANT)

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IA Me	dicaid	Men	ber	ID #				Pat	tient name			DC	ЭВ						
Patien	t addı	ress																	
Provid	er NI	PI							Prescriber name			Pho	one						
Prescr	iber a	addre	ss									Fax	(
Pharmacy name Ad						Ad	Address			Phone									
Presc	riber	mus	t cor	mple	te al	l info	rma	ation	n above. It must be legible, correct, and co	mplet	e or f	orm	will I	be re	eturi	ned.			
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Prior authorization is required for use of high dose opioids ≥ 90 morphine milligram equivalents (MME) per day. (See CDC Guideline for Prescribing Opioids for Chronic Pain at https://www.cdc.gov/opioids/healthcare-professionals/prescribing/guideline/index.html) Patients undergoing active cancer treatment or end-of-life care will not be subject to the criteria below. Payment will be considered when the following is met:

- 1. Requests for non-preferred opioids meet criteria for coverage (see criteria for Long-Acting Opioids and/or Short-Acting Opioids); and
- 2. Patient has a diagnosis of severe, chronic pain with a supporting ICD-10 code. Requests for a diagnosis of fibromyalgia or migraine will not be considered; and
- 3. Patient has tried and failed at least two nonpharmacologic therapies (physical therapy; weight loss; alternative therapies such as manipulation, massage, and acupuncture; or psychological therapies such as cognitive behavior therapy (CBT); and
- 4. Patient has tried and failed at least two nonopioid pharmacologic therapies (acetaminophen, NSAIDs, or selected antidepressants and anticonvulsants); and
- 5. There is documentation demonstrating an appropriate upward titration or an appropriate conversion from other opioid medications;
- 6. Pain was inadequately controlled at the maximum allowed dose without prior authorization for the requested opioid(s); and
- 7. Pain was inadequately controlled by two other chemically distinct preferred long-acting opioids at the maximum allowed dose without prior authorization; and
- 8. Chart notes from a recent office visit or telehealth visit for pain management are included documenting the following: a) Treatment plan, including all therapies to be used concurrently (pharmacologic and nonpharmacologic); and b) Treatment goals; and
- 9. Patient has been informed of the risks of high-dose opioid therapy; and
- 10. The prescriber has reviewed the patient's use of controlled substances on the Iowa Prescription Monitoring Program website and determined that use of high-dose opioid therapy is appropriate for this patient; and
- 11. The patient's risk for opioid addiction, abuse and misuse has been reviewed and prescriber has determined the patient is a candidate for high-dose opioid therapy; and
- 12. A signed chronic opioid therapy management plan between the prescriber and patient dated within 12 months of this request is included; and
- 13. The requested dosing interval is no more frequent than the maximum FDA-approved dosing interval; and
- 14. Patient has documentation of receipt of an opioid reversal agent (e.g. as seen in pharmacy claims or documentation from the Iowa PMP of dispensation [attach documentation]) within the prior 24 months of high dose opioid request for the emergency treatment of an opioid overdose; and
- 15. Patient has been educated on opioid overdose prevention; and
- 16. Patient's household members have been educated on the signs of opioid overdose and how to administer an opioid reversal agent; and
- 17. Patient will not be using opioids and benzodiazepines concurrently or a taper plan to discontinue the benzodiazepine must be submitted with initial and subsequent requests; and
- 18. A documented dose reduction is attempted at least annually.

If criteria for coverage are met, initial requests will be given for three months. Requests for continuation of high-dose opioid therapy will be considered every six months with the following:

- 1. High-dose opioid therapy continues to meet treatment goals, including sustained improvement in pain and function; and
- 2. Patient has not experienced an overdose or other serious adverse event; and
- 3. Patient is not exhibiting warning signs of opioid use disorder; and
- 4. The benefits of opioids continue to outweigh the risks; and
- 5. A documented dose reduction has been attempted at least annually, and the prescriber has determined the dose cannot be reduced at this time; and

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- 6. The prescriber has reviewed the patient's use of controlled substances on the Iowa Prescription Monitoring Program website and determined that continued use of high-dose opioid therapy is appropriate for this patient; and
- 7. Patient will not be using opioids and benzodiazepines concurrently or a taper plan to discontinue the benzodiazepine must be submitted with subsequent requests; and
- 8. Patient has documentation of receipt of an opioid reversal agent (e.g. as seen in pharmacy claims or documentation from the lowa PMP of dispensation [attach documentation]) within 24 months of high dose opioid request for the emergency treatment of an opioid overdose; and

Drug name:	Strength:				
Dosage instructions:	Quantity:				
Drug name:	Strength:				
Dosage instructions:	Quantity:	Days supply:			
Diagnosis:		ICD-10 code:			
* Proceed to Prescriber Signature for active cancer	treatment or end of life care diagnoses.				
Initial Requests:					
Document non-pharmacologic therapies (suc and acupuncture; or psychological therapies such as		e therapies such as manipulation, massage,			
Non pharmacological treatment trial #1:					
Trial dates:	Failure reason:				
Non pharmacological treatment trial #2:					
Trial dates:	Failure reason:				
Document two nonopioid pharmacologic the	erapies (acetaminophen, NSAIDs, or select	ed antidepressants, and anticonvulsants)			
Nonopioid pharmacologic trial #1: Name/dose:					
Trial dates:	Failure reason:				
Nonopioid pharmacologic trial #2: Name/dose:					
Trial dates:	Failure reason:				
·					
Document upward titration or conversion from	om other opioid medications:				
	dose allowed without prior authorization fo	or the requested opioid(s)?			
Was pain inadequately controlled at the maximum Document dose and trial dates: Was pain inadequately controlled by two other che	dose allowed without prior authorization fo	or the requested opioid(s)?			
Document upward titration or conversion from the second se	dose allowed without prior authorization for semically distinct preferred long-acting opioic below.	or the requested opioid(s)? No Y			
Was pain inadequately controlled at the maximum Document dose and trial dates: Was pain inadequately controlled by two other che prior authorization? No Yes Document	dose allowed without prior authorization for emically distinct preferred long-acting opioic below.	or the requested opioid(s)? No Yos Yos Yos Yos Yos Yos Yos Yos Yos Yo			
Was pain inadequately controlled at the maximum Document dose and trial dates: Was pain inadequately controlled by two other che prior authorization? No Yes Document Preferred long-acting narcotic trial #1: Name/dose	dose allowed without prior authorization for smically distinct preferred long-acting opioic below.	or the requested opioid(s)? No			

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□ No

☐ Yes

Attach notes from a recent office visit for pain management documenting both of the following:

Has patient been informed of the risks of high-dose opioid therapy?

Treatment goals

Treatment plan, including all therapies to be used concurrently (pharmacologic and nonpharmacologic)







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Prescriber review of patient's controlled substance use on the Iowa PMP website: No Yes Date reviewed:
Is long-acting opioid use appropriate for patient based on PMP review and patient's risk for opioid addiction, abuse and misuse?
□ No □ Yes
Attach a signed chronic opioid therapy management plan between the prescriber and patient dated within 12 months of this request.
Has patient received an opioid reversal agent within 24 months of this request for the emergency treatment of an opioid overdose?
No Yes Date Received (attach Iowa PMP record if not in pharmacy claims):
Has patient been educated on opioid overdose prevention?
Has patient's household members been educated on the signs of opioid overdose and how to administer an opioid reversal agent? No Yes Date:
Is patient using opioids and benzodiazepines concurrently? No Yes (provide taper plan to discontinue the benzodiazepine)
Date of patient's most recent documented dose reduction:
Renewals:
Does high-dose opioid therapy continue to meet treatment goals, including sustained improvement in pain and function? No Yes (describe):
Has patient experienced an overdose or other serious adverse event?
Is patient exhibiting warning signs of opioid use disorder?
Do the benefits of opioids continue to outweigh the risks?
Date of patient's most recent documented dose reduction:
Updated prescriber review of patient's controlled substances use on the Iowa PMP website: No Yes Date reviewed:
Is patient using opioids and benzodiazepines concurrently?
Has patient received an opioid reversal agent within 24 months of this request for the emergency treatment of an opioid overdose?
☐ No ☐ Yes Date Received (attach Iowa PMP record if not in pharmacy claims):
Has patient been reeducated on opioid overdose prevention?
Has patient's household members been reeducated on the signs of opioid overdose and how to administer an opioid reversal agent? No Yes Date:
Attach a signed chronic opioid therapy management plan between the prescriber and patient dated within 12 months of this request.
Prescriber signature (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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