

Request for Prior Authorization SHORT ACTING OPIOIDS



FAX Completed Form To

1 (877) 733-3195

(PLEASE PRINT – ACCURACY IS IMPORTANT)

IA Medicaid Member ID #	Patient name	C	OOB			
Patient address						
Provider NPI	Prescriber name	P	hone			
Prescriber address		F	ax			
Pharmacy name	Address	P	hone			
Prescriber must complete all informa	tion above It must be legih	le correct and complete or form	will be returned			
Pharmacy NPI	Pharmacy fax					
Prior authorization (PA)is requir	ed for all non-preferred	short acting onioids PA is	also required for members			
Prior authorization (PA)is required for all non-preferred short acting opioids. PA is also required for members when the total daily opioid dose (combined across all opioids) exceeds the set morphine milligram equivalent (MME) threshold (include High Dose Opioids PA form with request). Payment will be considered under the following conditions: 1) Patient has pain severe enough to require opioid treatment; and 2) Patient has tried and failed at least two nonpharmacologic therapies; and 3) Patient has tried and failed at least two nonpharmacologic therapies; and 3) Patient has tried and failed at least two nonopioid pharmacologic therapies; and 3) Patient has tried and failed at least two nonopioid pharmacologic therapies; and 4) Patient has documentation of previous trials and therapy failures with three (3) chemically distinct preferred short acting opioids (based on opioid ingredient only) at therapeutic doses; and 5) The prescriber has reviewed the patient's use of controlled substances on the lowa Prescription Monitoring Program (PMP) website and has determined that use of a short-acting opioid is appropriate for this member based on review of PMP and the patient's risk for opioid addiction, abuse and misuse prior to requesting prior authorization; and 6) Patient has been informed of the common adverse effects and serious adverse effects of opioids; and 7) For patients taking concurrent benzodiazepines, the prescriber must document the following: a. The risks of using opioids and benzodiazepines concurrently has been discussed with the patient; and b. Documentation as to why concurrent use is medically necessary is provided; and c. A plan to taper the benzodiazepine is provided, if appropriate. If criteria for coverage are met, an initial authorization will be given for 3 months. Additional approvals will be considered if the following criteria are met: 1) Patient has experienced improvement in pain control and level of functioning; and 2) Prescriber has reviewed the patient's use of controlled substances on						
Preferred(*PleaserefertothePDLfclistofpreferredalternatives)Acetaminophen/CodeineOxycodoHydrocodone/APAP(5/325)Hydromorphone TabTramadoMorphine Sulfate TabOxycodone Cap/Tab	one /APAP	eferred Butalbital/APAP/Caff/Codeine Butalbital/ASA/Caff/Codeine Combunox Hydrocodone/APAP (5/300, 7.5/300, 10/300) Hydrocodone/Ibuprofen	Nucynta Oxymorphone Oxycodone/APAP (7.5/325, 10/325) Primlev Roxicodone			

Other (specify)

Meperidine

Strength	Dosage Instructions	Quantity	Days Supply
Diagnosis:			

Tramadol 100mg

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			nt loss, alternative therapies such as s cognitive behavior therapy [CBT], etc,)
Non-Pharmacological Treatmen	t Trial #1:		
Non-Pharmacological Treatmen	t Trial #2:		
Document 2 nonopioid pharm Nonopioid Pharmacologic Trial #		-	NDs) Trial Dates:
Failure reason:			
			Trial Dates:
Document trials with three pre	eferred chemically dis	tinct short acting op	bioids
Preferred Trial 1: Drug Name		Strength	Dosage Instructions
Trial start date:	Trial end date:		
Failure reason:			
Preferred Trial 2: Drug Name		Strength	Dosage Instructions
Trial start date:	Trial end date:		
Failure reason:			
Preferred Trial 3: Drug Name		Strength	Dosage Instructions
Trial start date:	Trial end date:		
Failure reason:			
Prescriber review of patient's	controlled substance	s use on the Iowa P	MP website: No 🗌 Yes Date Reviewed:
Is short-acting opioid use app and misuse?		ased on PMP review	and patient's risk for opioid addiction, abuse
confusion, tolerance, physical	dependence, and wi	thdrawal symptoms	n, dry mouth, nausea, vomiting, drowsiness, when stopping opioids) and serious adverse ous opioid use disorder) of opioids?
Patients taking concurrent be	nzodiazepines:		
Have the risks of using opioids a	nd benzodiazepines co	oncurrently been disc	ussed with the patient?
Medical necessity for concurrent	use:		

PAA-1103

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Provide plan to taper the benzodiazepine or medical rationale why not appropriate:

Renewals
Has patient experienced improvement in pain control and level of functioning?
□ No □ Yes (describe):
Updated prescriber review of patient's controlled substances use on the lowa PMP website (since initial request
Continued use of a short-acting opioid is appropriate for this member?
□ No □ Yes (describe):
Patients taking concurrent benzodiazepines:
Have the risks of using opioids and benzodiazepines concurrently been discussed with the patient?
Medical necessity for concurrent use:
Provide plan to taper the benzodiazepine or medical rationale why not appropriate:
Other medical conditions to consider:

Attach lab results and other documentation as necessary.		
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.