



Kansas Medical Assistance Program PA Phone 800-933-6593 PA Fax 800-913-2229 **Aetna Better Health of KS** PA Pharmacy Phone 855-221-5656 PA Pharmacy Fax 844-807-8453



Sunflower PA Pharmacy Phone 877-397-9526 PA Pharmacy Fax 866-399-0929



UnitedHealthcare
PA Pharmacy Phone 800-310-6826
PA Pharmacy Fax 866-940-7328

Prior Authorization for Opioid Products Indicated for Pain Management

Long Acting*	Short Acting*		
Buprenorphine (Butrans, Belbuca)	Benzhydrocodone		
Fentanyl transdermal (Duragesic)	Butorphanol		
Hydrocodone extended-release (Zohydro ER, Hysingla ER)	Codeine		
Hydromorphone extended-release (Exalgo)	Dihydrocodeine		
Methadone	Fentanyl		
Morphine controlled-release/extended-release (Kadian ER, Avinza, MS Contin, Oramorph, Arymo ER)	Hydrocodone		
Morphine/Naltrexone (Embeda)	Hydromorphone		
Oxycodone extended-release (OxyContin)	Levorphanol Tartrate		
Oxycodone extended-release (Xtampza ER)	Meperidine		
Oxycodone/Naloxone (Targiniq ER)	Morphine		
Oxycodone/Naltrexone (Troxyca ER)	Opium		
Oxymorphone extended-release (generic non-crush resistant)	Oxycodone		
Oxymorphone extended-release (Opana ER-crush resistant)	Oxymorphone		
Tapentadol extended-release (Nucynta ER)	Pentazocine/Naloxone		
Tramadol extended-release (Ultram ER, Ryzolt)	Tapentadol		
	Tramadol		

^{*}Includes brand and generic versions of the listed products unless otherwise noted (all salt forms, single and combination products, and all brand and generic formulations).

Beneficiary Information					
Name:					
Medicaid ID #:	Date of Birth:	Gender:			
Billing Provider Information (Pharmacy, Physician or Facility)					
Name:	Medicaid ID #:_		-		
NPI #:	Phone #:	Fax #:			
Prescriber Information					
Name:		Medicaid ID #:			
NPI #:	Phone #:	Fax #:			

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PATIE	ENT NAME:			MEDICAID ID:			
REQU	ESTED DR	UG					
Drug N	Name:		Dosage Strength:	Quantity:	Day Supply:		
Direct	ions:				<u> </u>		
Length	n of Therap	y:	Diagnosis:		_ ICD 10 Code:		
	•		ne, patient must have a diagnosis of terminal				
• If	, , , , , , , , , , , , , , , , ,						
Please	e complete			-			
	•	•	F THE FOLLOWING SECTIONS BASED ON THE PA CRITERI	A FOR THE MEDICATION BEIN	NG REQUESTED.)		
1)	•	care, or d	ated for pain related to active cancer diagonous the patient reside in an assisted or cu				
			Please Indicate: ☐ Cancer ☐ Sickle Cell	Disease ☐ Hospice/I	Palliative Care Custodial Care		
			**If YES , complete section H If NO proceed to question 2	,			
2)			·				
2)	-	being trea] YES	ated for pain related to a hospital discharg If YES, complete section A	ge, post-surgery or acu	ite trauma?		
		NO	If NO, complete questions 3 through 7				
3)	Has patie	nt received	d an opioid prescription for < 90 days in a	look back period of 4	months?		
] YES	If YES				
			 → For initial request, complete section → For renewal of a previous approval 				
] NO		,			
4)	Has patie	nt received	d an opioid prescription for ≥ 90 days in a	look back period of 4	months?		
] YES					
	 → For initial request, complete section B & D → For renewal of a previous approval, complete Section D & E 						
] NO	For renewar or a previous approvar	, complete section D	OX E		
5)	Does dos	e exceed 9	0 MME/day?*				
] YES] NO	If YES, complete Section F				
			vw.agencymeddirectors.wa.gov/calculator/dosecalculator.htm	riptionDrugCovContra/Downloads/Opio	oid-Morphine-EQ-Conversion-Factors-Aug-2017.pdf		
6)	CMS MME Conversion Guide https://www.cms.gov/Medicare/Prescription-Drug-Coverage/PrescriptionDrugCovContra/Downloads/Opioid-Morphine-EQ-Conversion-Factors-Aug-2017.pdf 6) Is the request for a long-acting opioid?*						
-		YES	If YES, complete Section G				
] NO					
	s exceeding 90 a look back p	-	and long acting opioid requests will only be apprononths.	ved for patients who have	received opioid prescriptions for ≥90		
7)	Is the req	uest for a	non-preferred medication?				
		YES	If YES, complete Section H				
] NO edicaid Pref	erred Drug List (PDL): http://www.kdheks.gov/hcf/	/pharmacy/download/PDLList.	<u>pdf</u>		

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PATIENT NAME:	MEDICAID ID:
SECTION A:	Request for pain related to a hospital discharge, post-surgery or acute trauma
	ust attest to ALL of the following for PA approval).
Y N	B a property of the second sec
	Prescriber has reviewed controlled substance prescriptions in the Prescription Drug Monitoring Program (PDMP)/K-TRACS.
	Prescriber has counseled the patient on potential respiratory depression.
	Cumulative opioid dose does not exceed 90 MME per day.
	Total day supply for the requested medication does not exceed 21 days (3 weeks).
SECTION B:	INITIAL PA (Prescriber must attest to ALL of the following for PA approval)
Y N	Prescriber has KMAP ID (required for opioid prescription approval)
	Patient has attempted or is contraindicated to treatment with at least 2 non-opioid ancillary treatments (e.g. NSAIDS, antidepressants, acetaminophen) in the last 90 days.
	Trial 1 Drug Name: Date Outcome
	Trial 2 Drug Name: Date Outcome
	List Contraindication or Intolerance (if any):
	Non-pharmacological treatment has been tried and/or is currently being used (e.g. exercise, cognitive behavior therapy, or interventional treatment).
	Prescriber has reviewed prescriptions for controlled substances in the Prescription Drug Monitoring Program (PDMP)/K-TRACS.
	Patient has been screened for substance abuse/opioid dependence.
	If patient is concurrently on a CNS depressant (e.g. benzodiazepines), prescriber has reviewed and will
	address the increased risk of respiratory depression with the patient. Patient has been screened for depression or other mental illnesses.
	If patient is positive for depression, patient is receiving either pharmacological or nonpharmacological treatment.
	Treatment duration and goals are defined with the patient and within the medical record.
	atients with <90 days of opioid prescription in past 4 months - Renewal (must meet ALL following)
Y N	
	Dose/frequency taper has been attempted.
	Reason for not tapering dose/frequency is documented in medical record.
Taper Outcome:	·
* <mark>REQUIRED</mark> -Ra	tionale for not tapering:
SECTION D:	Patients with ≥90 days of opioid prescription in past 4 months (must meet ALL following)
Y N	
	Patient has a pain management/opioid agreement with the prescriber.
	Patient has/will have random urine drug screens as part of their on-going therapy with opioids.
	Rationale for not tapering and discontinuing opioid.*
REQUIRED- Pre	scriber's rationale supporting inability to discontinue opioid therapy:

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PATIEN	T NAME:		MEDIO	CAID ID:			
SECTIO	N E: I	Patients with ≥90 opioid presc	ription in past 4 mo	nths - Ren	ewal	(must meet ALL follo	wing)
Υ	N						
		All narcotic analgesics are written b	<u> </u>				
		Prescriber has reviewed prescriptio (PDMP)/K-TRACS.	ns for controlled substa	nces in the P	Prescrip	tion Drug Monitoring F	'rogram
		Patient will not be maintained on m concurrently.	nore than one long-actin	g and one sl	hort-act	ting opioid analgesic	
		Documentation of treatment durat	ion and treatment goals	.*			
*Treatmo	ent dura	ation and goals:					
SECTIO	N F: [OOSE EXCEEDS 90 MME/DAY (must meet ONE of the	e following	:)		
Υ	N						
		Dose reduction has occurred since p	orevious approval.				
		Previous Dose:	New D	ose:			
		There is documentation of an attem	npted unsuccessful dose	taper withir	n the pa	ast 6 months.	
		Taper Date:					
		Taper Outcome:					
		Provider attests that a dose taper is	not clinically appropria	te for this pa	atient.		
		LONG-ACTING OPIOID (must n	neet ALL of the follow	ing)			
Υ	N	Batter the constitute about 1		20 1 1- 11		20 4	
		Patient has received a short-acting	<u> </u>	•			
		Patient has a documented history of preferred short-acting opioids.	of failure, contraindication	on or intolera	ance to	a trial of at least two	
		Trial 1 Drug Name:	Da	te	Out	tcome	
		Trial 2 Drug Name:	Dat	te	Out	tcome	

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List Contraindication or Intolerance (if any):

PATIEN	T NAME:		MEDICAID ID:
	THE PREI		MEDICATION AT: http://www.kdheks.gov/hcf/pharmacy/download/PDLList.pdf) box and provide the required information to receive the requested non-preferred drug.
Υ	N	INTOLERANCE/ CONTRAINDICATION	
			If there is one preferred agent in the preferred category, has the patient experienced an inadequate response after a trial of the preferred agent at a maximum tolerated dose, or do they have a documented intolerance or contraindication to the preferred agent?
			Trial – Drug Name: Date of Trial:
			List medical intolerance/allergy (if any):
			If there are two or more agents in the preferred category, has the patient experienced an inadequate response after a trial of two or more of the preferred agents at their maximum tolerated dose, or do they have a documented intolerance or contraindication to two or more preferred agents?
			Trial – Drug Name: Date of Trial:
			Trial – Drug Name: Date of Trial:
			List medical intolerance/allergy (if any):
			An appropriate formulation or indication is not available as a preferred drug. Please
			specify which formulation or indication is needed and information supporting the need:

Prescriber's Signature:	Date:	

This form will be returned unprocessed if it is not completed in its entirety.

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