

Long-Acting Opioid Request Form

Healthy Connections	%
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s this	request	for medication prescribed for treatment of pain related to cancer, palliative care, or end-of-life care?
∃Yes	□ No	(If yes, approve for six months. If no, approve for a one-month duration; must submit each month for
		three months, then subsequent approvals will be in three-month intervals.)

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Member name:	Member ID #:	Member date of birth:			
Medication allergies:	Member weight (kg):	Member height (ft/in):			
Prescriber name:	Prescriber specialty:	Medicaid provider ID # or NPI#:			
Prescriber address:					
Prescriber phone number:	Office fax number:	Office contact:			
This request is for: A long-acting opioid	d and patient has had previous history of sho	ort-acting opioids.			
Please explain medical necessity in detail:					
Drug information (one drug per requ	est form)				
Drug name/dosage form		Strength			
Directions		Quantity requested			
Request is for: 🗆 Initiation of therapy 🗆	Continuation of therapy				
For continuation of therapy, is the dose cu	rrently being tapered? ☐ Yes ☐ No				
If no, please explain:					
Treatment information					
This medication is being used for: acut	e condition □ chronic condition (check o	ne only)			
Is this medication being used for postoperative pain? \Box Yes \Box No \Box If yes, date of surgery:					

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Diagnoses for which the (ICD code and description	opioid is prescribed (includ n)	de primary and secondary	diagnoses applicable to	this request):	
Diagnosis		Date of diagnosis			
Diagnosis			Date of diagnosis		
	atments that have been tr			onpharmacological:	
Pharmacological treat	tments (including prefe	erred and nonpreferred	d medications)		
Drug / Strength	Long-acting or short acting (if applicable)	Directions	Start date/end date	Reason for discontinuation (if applicable)	
Nonpharmacological t	treatments				
	Treatment		Start da	ate/end date	

Prescriber attestation

Please indicate **Yes/True** or **No/False** for each of the following attestations. Explanation is required for each **No/False** answer in order for the request to be considered for approval.

	Yes (True)	No (False)	The prescriber attests to the following:				
			1. The scripts program will be accessed each time a controlled prescription is written for this patient.				
SOIC			2. Diagnosis is chronic pain and patient previously utilized short-acting opioids for this condition for at least three months without adequate pain relief and now requires around-the-clock pain management.				
			3. A Pain Treatment Plan and goals (updated upon reauthorization) consistent with the SC LLR Joint Revised Pain Management Guidelines is provided.				
			4. Patient is currently using or has been prescribed a separate nonopioid medication for baseline pain relief or a medical reason is given why nonopioid therapy cannot be used. Nonopioid medication can include, but is not limited to:				
			i. Antidepressants: Amitriptyline, Nortriptyline, Duloxetine, Venlafaxine, Savella				
			ii. Anticonvulsants: Gabapentin capsules, Carbamazepine				
			iii. Muscle relaxants: Baclofen, Cyclobenzaprine, Methocarbamol, Tizanidine tablets				
LONG - ACTING OPIOIDS			iv. NSAIDs: Aspirin, Celebrex (ST required), Diclofenac, Etodolac, Ibuprofen, Indomethacin, Meloxicam, Nabumetone, Naproxen, Salsalate, Sulindac				
Ę			v. Nonopioid analgesics: Acetaminophen				
G - AC			5. Dose prescribed is based on use of a functional pain scale with comparison to baseline and/or previous functional pain scale. Functional pain scale must be included in documentation.				
ő			6. Prescriber has attested to the following:				
_			a. An Opioid Treatment Agreement signed by both the patient and prescriber is on file.				
			b. Checking the SCRIPTS monitoring program for each request.				
			c. Benefits and potential harms of opioid use have been discussed with this patient. In addition, risks of combining opioids with other central nervous system depressants such as benzodiazepines, alcohol, other sedatives, skeletal muscle relaxants, illicit drugs such as heroin, or other opioids has been discussed with this patient.				
			d. If patient has a high-risk condition stated in the Centers for Disease Control and Prevention Guidelines (e.g., sleep apnea or other causes of sleep-disordered breathing, patients with renal or hepatic insufficiency, older adults, pregnant women, patients with depression or other mental health conditions, and patients with alcohol or other substance use disorders) prescriber attests to discussing heightened risks of opioid use and has educated patient on naloxone use and has considered prescribing naloxone.				
			7. For nonformulary opioid products, patient must meet criteria 1 – 6 and try and fail two formulary longacting opioid drugs.				
			8. For reauthorizations only : A treatment plan that includes current and previous goals of therapy for both pain and function has been developed for this patient.				
			No/False to any of the above attestations, please explain in detail:				
Pres	criber si	gnature	Date				