

## **Pharmacy Prior Authorization Form**

For Prior Authorization, please fax to: 877 974-4411 toll free, or 616 942-8206 Commercial (Traditional) Commercial (Individual/Optimized) This form applies to: Medicaid This request is: ☐ **Urgent** (life threatening) ☐ **Non-Urgent** (standard review) Urgent means the standard review time may seriously jeopardize the life or health of the patient or the patient's ability Fentanyl citrate lozenge (generic Actiq®) Member Last Name: First Name: DOB: Gender: ID #: Primary Care Physician: Prov. Phone: \_\_\_\_\_ Prov. Fax: \_\_\_\_\_ Requesting Provider: Provider Address: Provider NPI: Contact Name: Provider Signature: **Product Information** ☐ New request ☐ Continuation request Drug product: ☐ Fentanyl citrate 200 mcg lozenge Start date (or date of next dose): ☐ Fentanyl citrate 400 mcg lozenge Date of last dose (if applicable): ☐ Fentanyl citrate 600 mcg lozenge Dosing frequency: \_\_ ☐ Fentanyl citrate 800 mcg lozenge ☐ Fentanyl citrate 1,200 mcg lozenge ☐ Fentanyl citrate 1,600 mcg lozenge **Precertification Requirements** Before this drug is covered, the patient must meet all of the following requirements: 1. This drug is being used to manage breakthrough pain in cancer patients already receiving and tolerant to around-the-clock opioid therapy for persistent cancer pain

## **Dosage Limit**

dose of another opioid for 1 week or longer.

2. Age 16 or older

When criteria are met, the quantity of fentanyl citrate lozenge will be limited to #120 lozenges per 30 days.

**Note:** Authorization for indications, dosing, or a route of administration not approved by the Food and Drug Administration (FDA) or recognized in CMS-accepted compendia (e.g. DrugDex, AHFS, U.S. Pharmacopeia, and also Clinical Pharmacology for oncology indications only) require supporting evidence for coverage. Please provide two published peer-reviewed literature articles supporting the appropriateness of the drug, the dosing of the drug, or the route of administration to be used for the identified indication.

Note: Patients are considered opioid tolerant when taking oral morphine 60 mg/day or more, transdermal fentanyl 25 mcg/hr, oral oxycodone 30 mg/day, oral hydromorphone 8 mg/day, oral oxymorphone 25 mg/day, oral hydrocodone 60mg/day, or an equianalgesic



Pı	riority Health Precertification Documer	ntation			
1.	What condition is this drug being reques  Breakthrough cancer pain  Cancer diagnosis:			_	
	Other – the patient's condition is:				
2.	What is the patient's current opioid treatr	ment (must be aroun	d-the-clock opioids	s for persistent cancer p	oain):
	Drug	Dose	•	Dates	•
					:
3.	What other breakthrough pain medication				•
	Drug 	Dose		Dates	
					•
4.	Please provide other rationale for use, if	necessary:			