

### Clinical Pharmacy Program Guidelines for Fentanyl IR

Program	Prior Authorization
Medication	Abstral (fentanyl sublingual tablets), Actiq (fentanyl transmucosal lozenge), Fentora (fentanyl buccal tablet), Lazanda (fentanyl nasal spray), Subsys (fentanyl sublingual spray), and fentanyl citrate
Issue Date	6/2012
Pharmacy and Therapeutics Approval Date	8/2017
Effective Date	10/2017

**1. Background:**

Abstral, Actiq, Fentora, Lazanda, Subsys, and fentanyl citrate lozenges (generic Actiq) are opioid analgesics indicated for the management of breakthrough cancer pain in patients who are already receiving and have developed tolerance to around-the-clock opioid therapy for their underlying persistent cancer pain. Patients considered opioid tolerant are those who are taking at least 60 mg of oral morphine daily, at least 25 mcg/hour of transdermal fentanyl, at least 30 mg of oxycodone daily, at least 8 mg of oral hydromorphone daily, at least 25 mg of oral oxymorphone daily or an equianalgesic dose of another opioid for a week or longer. Patients must remain on around-the-clock opioids while taking one of these fentanyl products. Abstral, Actiq, Fentora, Lazanda, fentanyl citrate (generic Actiq) and Subsys must not be used in opioid non-tolerant patients because life-threatening hypoventilation could occur at any dose in patients not on a chronic regimen of opiates.

**2. Coverage Criteria:**

<p>A. Fentanyl citrate lozenges (generic Actiq), will be approved based on the following criteria:</p> <ol style="list-style-type: none"> <li>1. <b>All</b> of the following:             <ol style="list-style-type: none"> <li>a. Submission of medical records demonstrating use is for the management of pain associated with a cancer diagnosis (cancer diagnosis must be documented).</li> </ol> </li> </ol> <p style="text-align: center;"><b>-AND-</b></p> <ol style="list-style-type: none"> <li>b. Use is for the management of breakthrough cancer pain.</li> </ol>
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**-AND-**

- c. Patient must have at least a **one** week history of **one** of the following medications to demonstrate tolerance to opioids:
- 1) Morphine sulfate at a doses of greater than or equal to 60 mg/day
  - 2) Fentanyl transdermal patch at a dose of greater than or equal to 25 mcg/hr
  - 3) Oxycodone at a dose of greater than or equal to 30 mg/day
  - 4) Oral hydromorphone at a dose of greater than or equal to 8 mg/day
  - 5) Oral oxymorphone at a dose of greater than or equal to 25 mg/day
  - 6) An alternative opioid at an equianalgesic dose (e.g., oral methadone greater than or equal to 20 mg/day)

**-AND**

- d. The patient is currently taking a long-acting opioid around the clock for cancer pain.

**-AND-**

- e. **One** of the following:

- 1) The patient is not concurrently receiving an alternative fentanyl transmucosal product.

**-OR-**

- 2) The patient is currently receiving an alternative transmucosal fentanyl product **AND** the prescriber is requesting the termination of all current authorizations for alternative transmucosal fentanyl products in order to begin treatment with the requested medication. Only one transmucosal fentanyl product will be approved at a time. If previous authorizations cannot be terminated, the PA request will be denied.

**Authorization will be approved for 12 months.**

- B. Abstral, Actiq, Fentora, Lazanda, or Subsys will be approved based on the following criteria:

1. **All** of the following:

- a. Submission of medical records demonstrating use is for the management of pain associated with a cancer diagnosis (cancer diagnosis must be documented).

**-AND-**

b. Use is for the management of breakthrough cancer pain.

**-AND-**

c. Patient must have at least a **one** week history of **one** of the following medications to demonstrate tolerance to opioids:

- 1) Morphine sulfate at a doses of greater than or equal to 60 mg/day
- 2) Fentanyl transdermal patch at a dose of greater than or equal to 25 mcg/hr
- 3) Oxycodone at a dose of greater than or equal to 30 mg/day
- 4) Oral hydromorphone at a dose of greater than or equal to 8 mg/day
- 5) Oral oxymorphone at a dose of greater than or equal to 25 mg/day
- 6) An alternative opioid at an equianalgesic dose (e.g., oral methadone greater than or equal to 20 mg/day)

**-AND**

d. The patient is currently taking a long-acting opioid around the clock for cancer pain.

**-AND-**

e. **One** of the following:

- 1) The patient is not concurrently receiving an alternative transmucosal fentanyl product.

**-OR-**

- 2) The patient is currently receiving an alternative transmucosal fentanyl product **AND** the prescriber is requesting the termination of all current authorizations for alternative transmucosal fentanyl products in order to begin treatment with the requested medication. Only one transmucosal fentanyl product will be approved at a time. If previous authorizations cannot be terminated, the PA request will be denied.

**-AND-**

f. History of failure, contraindication, or intolerance to Fentanyl citrate lozenges (generic Actiq)

**Authorization will be approved for 12 months.**

### Quantity Limit Table

DrugName	Strength	Limit
Abstral	100mcg,200mcg, 300mcg,400mcg, 600mcg,800mcg	4 sublingual tablets per day
Actiq	200mcg,400mcg, 600mcg,800mcg, 1200mcg,1600mcg	4 lozenges per day
Fentora	100mcg,200mcg,400mcg,600mcg,800 mcg	4 buccal tablets per day
Lazanda	100 mcg, 400 mcg	5.3 mL (one bottle) per day
Subsys	100mcg,200mcg,400mcg,600mcg,800mcg,1200mcg,1600mcg	12 sprays per day

### 3. References:

1. Abstral package insert. Sentyln Therapeutics, Inc. Solana Beach, CA. December 2016
2. Actiq package insert. Cephalon: Frazer, PA. December 2016.
3. Fentora package insert. Cephalon: Frazer, PA. December 2016.
4. Lazanda package insert. Depomed, Inc. Newark, CA. March 2017.
5. Subsys package insert. Insys Therapeutics: Chandler, AZ. December 2016.

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<b>Change Control</b>	
Date	Change
6/2012	New policy

6/2013	<p>Changed policy name to “Oral Fentanyl Products”</p> <p>Converted policy to new UHC enterprise wide formatting</p> <p>No changes to clinical criteria</p>
3/2015	<p>Removed Onsolis from guideline and notification; Onsolis is no longer on the market. No clinical changes to criteria. Background revisions.</p>
9/2015	<p>Separated initial therapy criteria into preferred with prior authorization and non-preferred sections. Preferred, prior authorization criteria applies to Fentanyl Citrate and the new non-preferred section applies to Abstral, Brand Actiq, Fentora, Lazanda, and Subsys.</p> <p>Added prescriber requirement.</p> <p>Updated quantity limit criteria to simplify review criteria.</p>
9/2016	<p>Updated policy template. Updated clinical criteria to align with Employer &amp; Individual’s policy.</p>
12/2016	<p>Added clarification that prescriber requests the termination of all previous authorizations for transmucosal fentanyl products</p>
8/2017	<p>Added a step through fentanyl citrate lozenges for all branded products. Updated references.</p>