

## Fentanyl Transdermal Patch PA Criteria

**\*\*\*Continuation of therapy requests that do not meet criteria may be approved as amended for up to 6 months to allow for PA criteria compliance or dose tapering\*\*\***

1. Is the patient being treated for an OHP funded condition?
  - a. If Yes, move to question #2.
  - b. If No, review documentation for relevant comorbid conditions. If there are relevant comorbid conditions, move to question #2. If there are no relevant comorbid conditions, Cat. 1 denial.
  
2. Is the patient opioid-tolerant?

*Opioid-tolerant is defined as patients who are taking at least 60 mg/day of oral morphine, transdermal fentanyl 25 mcg/hour, oral oxycodone 30 mg/day, oral hydromorphone 8 mg/day, oral oxymorphone 25 mg/day, or equianalgesic dose of another opioid for **at least one week**.*

  - a. If Yes, move to question #3.
  - b. If No, Cat. 3 denial and Cat. 5 denial.

*Fentanyl transdermal patches are not indicated for opioid-naïve patients.*
  
3. Is the medication being used for the treatment of chronic pain severe enough to require daily, around-the-clock, long-term opioid treatment for which alternative treatments are inadequate?
  - a. If Yes, move to question #4.
  - b. If No, Cat. 3 denial and Cat. 5 denial.

*Fentanyl transdermal patches are only indicated for the treatment of chronic pain that meets this criteria.*
  
4. Is the medication being used for the treatment of pain associated with cancer or a terminal illness?
  - a. If Yes, **Approve for 12 months**.
  - b. If No, move to questions #5.
  
5. Does the dosing of the medication exceed the 50 mcg/hr patches?
  - a. If Yes, move to question #6.
  - b. If No, move to question #8.
  
6. Has patient tried and failed medication doses under the 50 mcg/hr patches?
  - a. If Yes, move to question #7.
  - b. If No, forward to Medical Director to assess medical appropriateness (Possible Cat. 5 denial).
  
7. Has patient been evaluated by a pain management specialist?
  - a. If Yes, move to question #8.
  - b. If No, forward to Medical Director to assess medical appropriateness (Possible Cat. 5 denial).
  
8. Is the medication dosing interval within the FDA approved dosing?

*FDA approved dosing interval for fentanyl transdermal patches is every 48 to 72 hours.*

  - a. If Yes, move to question #9.
  - b. If No, Cat. 3 denial and Cat. 5 denial.
  
9. Is there an established pain treatment agreement between patient and provider?

**Pain Treatment Agreements**  
Pain treatment agreements should include plans for random UAs, random pill counts, provider review of the prescription drug monitoring program (PDMP), patient use of a single pharmacy, a material risk notice (MRN), and patient abstinence from illegal drug use, marijuana use, and alcohol abuse.

  - a. If Yes, move to question #10.
  - b. If No, forward to Medical Director to assess medical appropriateness (Possible Cat. 5 denial).

10. Does the request represent a new start or a continuation of therapy?
  - a. If request is a New Start, **Approve for up to 90 days.**
  - b. If request is a continuation of therapy, move to question #11.
  
11. Does the provider submitted documentation indicate that medication use has demonstrated an improvement in patient’s function and pain status?
  - a. If Yes, move to question #12.
  - b. If No, forward to Medical Director to assess medical appropriateness (Possible Cat. 5 denial).
  
12. Has the patient been adherent to their established pain treatment agreement?
  - a. If Yes, **Approve for 6 months.**
  - b. If No, forward to Medical Director to assess medical appropriateness (Possible Cat. 5 denial).

<b>Guide to Denial Categories</b>	<b>Reason for Denial</b>
Category 1	The condition is not on a funded line
Category 3	The use of the medication is considered experimental/ investigational (usually applies to off-label use of a medication)
Category 5	Not medically appropriate
Category 15	Formulary medications have not been exhausted