Clinical Policy: Opioid Analgesics
Reference Number: HIM.PA.139
Effective Date: 08.01.18
Last Review Date: 02.19
Line of Business: HIM*

See Important Reminder at the end of this policy for important regulatory and legal information.

*Requests for transmucosal immediate-release fentanyl products (TIRFs) should be evaluated using the Fentanyl IR (Abstral, Actiq, Fentora, Lazanda, Subsys) policy – CP.PMN.127.

Description
Opioid analgesics exert their analgesic effect through opiate receptors distributed in tissues throughout the body.

This policy applies to all formulary long and short acting opioids requiring prior authorization or any non-formulary drug request that has met the formulary exception criteria – HIM.PA.103.

FDA Approved Indication(s)
Opioid analgesics are indicated for the management and treatment of moderate to severe pain.

Policy/Criteria
Provider must submit documentation (including such as office chart notes, lab results or other clinical information) supporting that member has met all approval criteria.

I. Initial Approval Criteria
   A. Cancer or Palliative Care (must meet all):
      1. Prescribed for pain associated with cancer or for palliative care (hospice or any terminal condition);
      2. If request is for a formulary long-acting or short-acting agent requiring prior authorization: member has failed an adequate trial of two other short-acting opioids analgesics dosed around the clock, unless contraindicated or clinically significant adverse effects are experienced;
      3. For OHIO request ONLY: If total dose of opioid exceeds 80 MME per day, member is stable (history of > 7 days of therapy) on current dose or documentation supports gradual upward titration of dose.

   Approval duration: 12 months

   B. Nucynta ER for Diabetic Peripheral Neuropathy (must meet all):
      1. Request is for Nucynta ER;
      2. Diagnosis of diabetic peripheral neuropathy;
      3. Age ≥ 18 years;
      4. Failure of gabapentin at ≥ 1800 mg/day unless contraindicated or clinically significant adverse effects are experienced;
5. Failure of a formulary tricyclic antidepressant (TCA) (e.g., amitriptyline, nortriptyline, imipramine) at up to maximally indicated doses, unless contraindicated or clinically significant adverse effects are experienced;
6. Failure of a formulary serotonin-norepinephrine reuptake inhibitor (SNRI) (e.g., duloxetine, venlafaxine) at up to maximally indicated doses, unless contraindicated or clinically significant adverse effects are experienced;
7. Dose does not exceed 500 mg per day.

Approval duration: Duration of request or 180 days (whichever is less)

C. Short-Acting Agents Requiring Formulary Prior Authorization (PA) ≤ a 14 day Supply (must meet all):
   1. Prescribed for the treatment of pain unrelated to cancer or palliative care;
   2. Failure of two formulary short-acting opioids analgesics dosed around the clock, unless contraindicated or clinically significant adverse effects are experienced;
   3. For OHIO requests ONLY: Total opioid dose does not exceed 80 MME per day or for members who are stable (history of > 7 days of therapy) on doses higher than 80 MME per day, one of the following is met (a or b):
      a. Provider will initiate a dose taper;
         *Future approval will require decrease from current dose.
      b. Documentation that a dose taper has been attempted within the past 6 months, with the reasons for taper failure;
   4. For OHIO requests ONLY: If opioid is being prescribed concomitantly with a benzodiazepine, member meets both of the following (a and b):
      a. Documentation that the provider has acknowledged combined use of opioid medication and benzodiazepine;
      b. Prescribed for short term (< 3 months) use or provider will discontinue concurrent use of benzodiazepine and opioid therapy within a 3-month period.
      *Re-authorization request for concurrent use of opioid and benzodiazepine will not be approved.

Approval Duration: 7 days

D. Long-Acting Agents OR Requests Exceeding a 14-day Supply Within 28 Days OR Requests Exceeding a 28-day Supply Within 90 Days (must meet all):
   *If member is new to Centene benefit and has received 90 days of the opioid in the last 120 days, approve request for 6 months and advise provider to attempt opioid taper.
   1. Prescribed for the treatment of pain unrelated to cancer or palliative care;
   2. Member meets one of the following (a or b):
      a. Prescribed agent is a short-acting agent that is covered without formulary PA requirement or member has failed an adequate trial of two other short-acting opioids analgesics on the formulary, dosed around the clock, unless contraindicated or clinically significant adverse effects are experienced;
      b. Failure of an adequate trial of two short-acting opioids analgesics dosed around the clock; additionally if formulary PA is required, failure of an adequate trial of 2 formulary long-acting agents, unless contraindicated or clinically significant adverse effects are experienced;
3. Failure of at least 2 non-opioid ancillary treatments (such as non-steroidal anti-inflammatory drugs (NSAIDs), acetaminophen, anticonvulsants, antidepressants, etc.) unless contraindicated or clinically significant adverse effect are experienced;

4. For OHIO requests ONLY: Total opioid dose does not exceed 80 MME per day or for members who are stable (history of > 7 days of therapy) on doses higher than 80 MME per day, one of the following is met (a or b):
   c. Provider will initiate a dose taper;
      \*Future approval will require decrease from current dose.
   d. Documentation that a dose taper has been attempted within the past 6 months, with the reasons for taper failure;

5. For OHIO requests ONLY: If opioid is being prescribed concomitantly with a benzodiazepine, member meets both of the following (a and b):
   a. Documentation that the provider has acknowledged combined use of opioid medication and benzodiazepine;
   b. Prescribed for short term (< 3 months) use or provider will discontinue concurrent use of benzodiazepine and opioid therapy within a 3-month period.

Approval duration: Duration of request or 30 days (whichever is less)

E. Other diagnoses/indications – Not applicable

II. Continued Therapy
A. Cancer or Palliative Care (must meet all):
   1. Currently receiving prescribed agent via Centene benefit for cancer and palliative care or have previously met initial approval criteria;
   2. For OHIO requests ONLY: If total dose of opioid exceeds 80 MME/day, member is stable (history of > 7 days of therapy) on current dose or documentation supports gradual upward titration of dose.

Approval duration: 12 months

B. Nucynta ER for Diabetic Peripheral Neuropathy (must meet all):
   1. Currently receiving Nucynta ER for the diagnosis of diabetic peripheral neuropathy or member has met initial approval criteria;
   2. Provider submits medical justification supporting continued need of Nucynta ER;
   3. If request is for a dose increase, new dose does not exceed 500 mg per day.

Approval duration: Duration of request or 30 days (whichever is less)

C. Short Acting Agents Requiring Formulary Prior Authorization (PA) ≤ a 14 day Supply (must meet all):
   1. Previously received medication via Centene benefit or has previously met the initial approval criteria;
   2. For OHIO requests ONLY: Total opioid dose should NOT exceed 80 MME/day or if the current dose is higher than 80 MME/day, one of the following is met (a, b or c):
      a. Dose reduction has occurred since previous approval;
      b. A dose taper has been attempted within the past 6 months and was not successful;
      \*Reason(s) for taper failure must be provided
c. Prescribed by or consultation with a pain management specialist;
3. For OHIO requests ONLY: If the requested dose is for > 80 MME/day, an increase in dose has not occurred since previous approval;
4. For OHIO requests ONLY: If opioid is being prescribed concomitantly with a benzodiazepine, member meets all of the following (a, b, and c):
   a. Currently receiving concurrent opioid and benzodiazepine therapy via Centene benefit or member has previously met the initial approval criteria;
   b. Documentation supports that discontinuation of combination opioid and benzodiazepine therapy has been attempted in the last 3 months without success;
   c. Prescribed by or in consultation with a pain management specialist.

Approval Duration: 7 days

D. Long-Acting Agents OR Requests Exceeding a 14-day Supply Within 28 Days OR 28-day Supply Within 90 Days (must meet all):
   *If member is new to Centene benefit and has received 90 days of the opioid in the last 120 days, approve request for 6 months and advise provider to attempt opioid taper.
   1. Previously received medication via Centene benefit or member has previously met initial approval criteria;
   2. Has received more than a 7-day supply of opioid in the last 90 days; 
      If member does not meet this requirement, please use the initial approval criteria to review this request
   3. Provider submits medical justification supporting continued need of opioid analgesics;
   4. For OHIO requests ONLY: Total opioid dose should NOT exceed 80 MME/day or if the current dose is higher than 80 MME/day, one of the following is met (a, b or c):
      a. Dose reduction has occurred since previous approval;
      b. A dose taper has been attempted within the past 6 months and was not successful; 
         *Reason(s) for taper failure must be provided
      c. Prescribed by or consultation with a pain management specialist;
   5. For OHIO requests ONLY: If the requested dose is for > 80 MME/day, an increase in dose has not occurred since previous approval;
   6. For OHIO requests ONLY: If opioid is being prescribed concomitantly with a benzodiazepine, member meets all of the following (a, b, and c):
      a. Currently receiving concurrent opioid and benzodiazepine therapy via Centene benefit or member has previously met the initial approval criteria;
      b. Documentation supports that discontinuation of combination opioid and benzodiazepine therapy has been attempted in the last 3 months without success;
      c. Prescribed by or in consultation with a pain management specialist.

Approval duration: Duration of request or 30 days (whichever is less)

E. Other diagnoses/indications – Not applicable

III. Diagnoses/Indications for which coverage is NOT authorized – Not applicable

IV. Appendices/General Information
   Appendix A: Abbreviation/Acronym Key
   FDA: Food and Drug Administration
   MME: morphine milligram equivalents
CLINICAL POLICY
Opioids Analgesics

NSAID: non-steroidal anti-inflammatory drug
SNRI: serotonin-norepinephrine reuptake inhibitor
TCA: tricyclic antidepressant

Appendix B: Therapeutic Alternatives
Not applicable

Appendix C: Contraindications/Boxed Warnings
Contraindications: significant respiratory depression; acute or severe bronchial asthma; gastrointestinal obstruction, including paralytic ileus; hypersensitivity to the opioid active ingredient, salts, or any component of the product.
Boxed warnings: potential for addiction, abuse, and misuse; life-threatening respiratory depression; accidental ingestion; neonatal opioid withdrawal syndrome; cytochrome P450 3A4 interactions; risks from concomitant use with benzodiazepines or other CNS depressants.

Appendix D: General Information

<table>
<thead>
<tr>
<th>Opioid Oral Morphine Milligram Equivalent (MME) Conversion Factors</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Type of Opioid (strength units)</strong></td>
</tr>
<tr>
<td>Codeine (mg)</td>
</tr>
<tr>
<td>Dihydrocodeine (mg)</td>
</tr>
<tr>
<td>Fentanyl buccal or SL tablets, or lozenge/troche (mcg)</td>
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<tr>
<td>Fentanyl film or oral spray (mcg)</td>
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<tr>
<td>Fentanyl nasal spray (mcg)</td>
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<tr>
<td>Fentanyl patch (mcg)</td>
</tr>
<tr>
<td>Hydrocodone (mg)</td>
</tr>
<tr>
<td>Hydromorphone (mg)</td>
</tr>
<tr>
<td>Levorphanol tartrate (mg)</td>
</tr>
<tr>
<td>Meperidine hydrochloride (mg)</td>
</tr>
<tr>
<td>Methadone (mg)</td>
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<tr>
<td>&gt; 0, ≤ 20</td>
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<tr>
<td>&gt; 20, ≤ 40</td>
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<tr>
<td>&gt; 40, ≤ 60</td>
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<tr>
<td>&gt; 60</td>
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<tr>
<td>Morphine (mg)</td>
</tr>
<tr>
<td>Opium (mg)</td>
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<tr>
<td>Oxycodone (mg)</td>
</tr>
<tr>
<td>Oxymorphone (mg)</td>
</tr>
<tr>
<td>Pentazocine (mg)</td>
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<tr>
<td>Tapentadol (mg)</td>
</tr>
<tr>
<td>Tramadol (mg)</td>
</tr>
</tbody>
</table>
V. Dosage and Administration

There are numerous narcotic analgesics, please refer to the package insert of your drug of interest for information on appropriate dosage and administration.

VI. Product Availability

There are numerous narcotic analgesics, please refer to the package insert of your drug of interest for product availability information.

VII. References


<table>
<thead>
<tr>
<th>Reviews, Revisions, and Approvals</th>
<th>Date</th>
<th>P&amp;T Approval Date</th>
</tr>
</thead>
<tbody>
<tr>
<td>New policy created</td>
<td>06.09.18</td>
<td>06.18</td>
</tr>
<tr>
<td>No significant changes: modified the day supply requirement for PA override to align with programming; request exceeding 7 day supply/90 days changed to requests exceeding a 14-day supply within 28 Days OR 28-day supply within 90 Days</td>
<td>07.17.18</td>
<td></td>
</tr>
<tr>
<td>Added redirection to TIRF policy; clarified that redirection in section C is to other formulary agents; notated that section C should apply for request for &lt; 14 day supply; added HIM OH state requirements to policy from a previously approved policy.</td>
<td>09.10.18</td>
<td></td>
</tr>
<tr>
<td>1Q 2019 annual review: no significant changes; references reviewed and updated.</td>
<td>11.05.18</td>
<td>02.19</td>
</tr>
</tbody>
</table>

Important Reminder

This clinical policy has been developed by appropriately experienced and licensed health care professionals based on a review and consideration of currently available generally accepted standards of medical practice; peer-reviewed medical literature; government agency/program approval status; evidence-based guidelines and positions of leading national health professional organizations; views of physicians practicing in relevant clinical areas affected by this clinical policy; and other available clinical information. The Health Plan makes no representations and accepts no liability with respect to the content of any external information used or relied upon in developing this clinical policy. This clinical policy is consistent with standards of medical practice current at the time that this clinical policy was approved. “Health Plan” means a health plan that has adopted this clinical policy and that is operated or administered, in whole or in part, by Centene Management Company, LLC, or any of such health plan’s affiliates, as applicable.

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