Program Number | 2018 P 2014-16
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Program | Prior Authorization/Medical Necessity - Long-Acting Opioid Pain Medications
Medication | Includes both brand and generic versions of the listed products unless otherwise noted: Arymo ER^ (morphine sulfate extended-release), Avinza^ (morphine sulfate controlled-release capsules), Dolophine (methadone), Duragesic (fentanyl transdermal)^ 12, 25, 50, 75, 100 mcg/hr, Embeda^ (morphine sulfate and naltrexone), Exalgo^ (hydromorphone extended-release), fentanyl transdermal (37.5, 62.5, 87.5 mcg/hr), Hysingla ER^ (hydrocodone extended-release), Kadian^ (morphine sulfate sustained-release capsules^), Morphabond ER^ (morphine sulfate extended-release), morphine sulfate controlled-release (generic MS Contin), MS Contin, Nucynta ER (tapentadol extended-release), Opana ER (oxymorphone extended-release), OxyConti^ (oxycodone controlled-release^), Xtampza ER (oxycodone extended-release), Zohydro ER (hydrocodone extended-release)
Effective Date | 8/1/2018; Oxford only: 9/1/2018

1. **Background:**
   Long-acting opioid analgesics, Arymo ER, Avinza, Embeda, Exalgo, fentanyl transdermal, hydromorphone ER, Hysingla ER, Kadian, Morphabond ER, MS Contin, Nucynta ER, Opana ER, oxycodone hcl ER, OxyContin, oxymorphone ER, Xtampza ER and Zohydro ER are indicated for the management of moderate to severe pain when a continuous, around-the-clock opioid is needed for an extended period of time and for which alternative treatment options are not appropriate. They are not intended for use as an as needed analgesic.
   Long-acting opioids are not indicated for pain in the immediate postoperative period (the first 12-24 hours following surgery), or if the pain is mild, or not expected to persist for an extended period of time. They are only indicated for postoperative use if the patient is already receiving the drug prior to surgery or if the postoperative pain is expected to be moderate to severe and persist for an extended period of time. Physicians should individualize treatment, moving from parenteral to oral analgesics as appropriate.
Long-acting opioids should not be used in treatment naïve patients. Physicians should individualize treatment in every case, initiating therapy at the appropriate point along a progression from non-opioid analgesics, such as non-steroidal anti-inflammatory drugs and acetaminophen to opioids in a plan of pain management such as those outlined by the World Health Organization, the Agency for Healthcare Research and Quality, the Federation of State Medical Boards Model Guidelines, or the American Pain Society.

The CDC and the American Academy of Neurology recommends the following best practices in the prescription of long-acting opioids:

- Nonpharmacologic therapy and nonopioid pharmacologic therapy are preferred for chronic pain.
- Before starting opioid therapy, treatment goals should be established with patients that include realistic goals for pain and function and should consider how therapy will be discontinued if benefits do not outweigh risks. Track pain and function at every visit (at least every 3 months) using a brief, validated instrument. Continue opioid therapy only if there is clinically meaningful improvement in pain and function that outweighs risks to patient safety.
- When starting opioid therapy for chronic pain, clinicians should prescribe immediate-release opioids instead of extended release/long-acting opioids.
- Document the daily morphine equivalent dose (MED) in mg/day from all sources of opioids. Access the state prescription drug monitoring program (PDMP) data at treatment initiation and periodically during treatment. Currently all states except for Missouri have a PDMP.
- To avoid increased risk of respiratory depression, long-acting opioids should not be prescribed concurrently with benzodiazepines. Screen for past and current substance abuse and for severe depression, anxiety, and PTSD prior to initiation.
- Use random urine drug screening prior to initiation and periodically during treatment with a frequency according to risk.
- Use a patient treatment agreement, signed by both the patient and prescriber that address risks of use and responsibilities of the patient. Avoid escalating doses above 50-90 mg/day MED unless sustained meaningful improvement in pain and function is attained, and not without consultation with a pain management specialist.
- Clinicians should evaluate benefits and harms of continued therapy at least every 3 months. If benefits do not outweigh harms, opioids should be tapered and discontinued. Evaluation should include assessment of substance use disorder/opioid dependence. Validated scales (such as the DAST-10) are available at www.drugabuse.gov.

Section Overview
Section 2. Medical Necessity Coverage Criteria for Book of Business
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## Section 7: Medical Necessity Coverage Criteria for State of West Virginia

### 2. Coverage Criteria

(Refer to section overview for state specific criteria and supply limit coverage criteria)

<table>
<thead>
<tr>
<th>A. Cancer or End of Life (defined as a &lt; 2 year life expectancy) related pain</th>
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<tr>
<td>1. <strong>Dolophine, fentanyl transdermal patch (generic Duragesic)</strong> 12, 25, 50, 75, 100 mcg/hr, <strong>Nucynta ER</strong>, <strong>methadone</strong>, <strong>morphine sulfate controlled-release tablets (generic MS Contin)</strong>, and <strong>Xtampza ER</strong> will be approved for cancer related pain based on the following criteria:</td>
</tr>
</tbody>
</table>
| a. Patient is being treated for cancer or end of life related pain  

-AND-  

b. Submission of medical records documenting active cancer diagnosis or life expectancy of < 2 years. |
| 2. **Arymo ER**, **Avinza**, **Duragesic**, **Embeda**, **Exalgo**, **fentanyl transdermal patch (37.5, 62.5, 87.5 mcg/hr)**, **Hysingla ER**, **Kadian**, **Morphabond ER**, **MS Contin**, **oxymorphone extended release and Zohydro ER** [Applies to all brand and generic versions of listed products except generic morphine sulfate controlled-release tablets (generic MS Contin) and fentanyl transdermal patch (generic Duragesic strengths) ] will be approved based on **ALL** of the following criteria: |
| a. Patient is being treated for cancer or end of life related pain  

-AND-  

b. Submission of medical records documenting active cancer diagnosis or life expectancy of < 2 years.  

-AND-  

c. The patient has a history of failure, contraindication or intolerance to a trial of **ALL** of the following (Document date of trial):  

(1) Nucynta ER  
(2) morphine sulfate controlled-release tablets (specifically generic MS Contin)  
(3) Xtampza ER |
| 3. **OxyContin** and oxycodone controlled-release (Authorized Generic for |
OxyContin™ will be approved based on ALL of the following criteria:

a. Patient is being treated for cancer or end of life related pain

-AND-

b. Submission of medical records documenting active cancer diagnosis or life expectancy of < 2 years.

-AND-

c. **ONE** of the following:

   (1) The patient has a history of failure, contraindication or intolerance to ALL of the following (Document date of trial):

      (a) Xtampza ER
      (b) Nucynta ER
      (c) morphine sulfate controlled-release tablets (specifically generic MS Contin)

   --OR--

   (2) **Both** of the following:

      (a) The patient requires more than 320 mg/day of controlled-release oxycodone.

   --AND--

      (b) The patient has a history of failure, contraindication or intolerance to BOTH of the following (Document dates of trial):

      i. Nucynta ER
      ii. morphine sulfate controlled-release tablets (specifically generic MS Contin)

Authorization will be issued for 24 months up to the dose allowed by supply limit review (please refer supply limit criteria). If the patient is currently taking the requested long-acting opioid OR was recently switched from another long-acting opioid and does not meet the medical necessity initial authorization criteria requirements for long-acting opioids, a denial should be issued and a maximum 90-day authorization may be authorized one time for the requested drug/strength combination up to the requested quantity.
for transition to an alternative treatment.

B. Non-cancer and Non-End of Life pain

1. Initial Authorization

   a. Dolophine, fentanyl transdermal patch (generic Duragesic) 12, 25, 50, 75, 100 mcg/hr, Nucynta ER, methadone, morphine sulfate controlled-release tablets (generic MS Contin), and Xtampza ER will be approved based on the following criteria:

   (1) Prescriber attests to **ALL** of the following:
   - The information provided is true and accurate to the best of their knowledge and they understand that UnitedHealthcare may perform a routine audit and request the medical information necessary to verify the accuracy of the information provided.
   - Treatment goals are defined, including estimated duration of treatment.
   - Treatment plan includes the use of a non-opioid analgesic and/or non-pharmacologic intervention
   - Patient has been screened for substance abuse/opioid dependence
   - If used in patients with medical comorbidities or if used concurrently with a benzodiazepine or other drugs that could potentially cause drug-drug interactions, the prescriber has acknowledged that they have completed an assessment of increased risk for respiratory depression.
   - Pain is moderate to severe and expected to persist for an extended period of time
   - Pain is chronic
   - Pain is not postoperative (unless the patient is already receiving chronic opioid therapy prior to surgery, or if the postoperative pain is expected to be moderate to severe and persist for an extended period of time)
   - Pain management is required around the clock with a long acting opioid

   -AND-

   (2) **ONE** of the following:
(a) The patient is new to the plan (as evidenced by coverage effective date of less than or equal to 120 days) and is currently established on the requested long-acting opioid.

-OR-

(b) ONE of the following:

i. **ALL** of the following:

   a) The patient is being treated for moderate to severe chronic pain that is **non-neuropathic** (examples of neuropathic pain include neuralgias, neuropathies, fibromyalgia)

   -AND-

   b) Prior to the start of therapy with the long-acting opioid, the patient has failed an adequate (minimum of 4 week) trial of a short-acting opioid. (Document drug(s), duration and date of trial).

-OR-

ii. **ALL** of the following:

   a) The patient is being treated for moderate to severe **neuropathic pain or fibromyalgia**

   -AND-

   b) Unless it is contraindicated, the patient has not exhibited an adequate response to 8 weeks of treatment with gabapentin titrated to a therapeutic dose. (Document duration, dose and date of trial)

   -AND-

   c) Unless it is contraindicated, the patient has not exhibited an adequate response to at least 6 weeks of treatment with a tricyclic antidepressant titrated to the maximum tolerated dose. (Document drug, dose, date, and duration of trial)

b. **Arymo ER**, **Avinza**, **Duragesic**, **Embeda**, **Exalgo**, fentanyl transdermal (37.5, 62.5, 87.5 mcg/hr), **Hysingla ER**, **Kadian**,
Morphabond ER\textsuperscript{\textregistered}, MS Contin, OxyContin\textsuperscript{\textregistered}, oxycodone controlled-release (Authorized Generic for OxyContin)\textsuperscript{\textregistered} oxymorphone extended release\textsuperscript{\textregistered} and Zohydro ER [Applies to all brand and generic versions of listed products except generic morphine sulfate controlled-release tablets (generic MS Contin) and fentanyl transdermal patch (generic Duragesic strengths)] will be approved for non-cancer and non-end of life related pain based on ALL of the following criteria:

(1) The prescriber attests to **ALL** of the following:

- The information provided is true and accurate to the best of their knowledge and they understand that UnitedHealthcare may perform a routine audit and request the medical information necessary to verify the accuracy of the information provided.
- Treatment goals are defined, including estimated duration of treatment.
- Treatment plan includes the use of a non-opioid analgesic and/or non-pharmacologic intervention
- Patient has been screened for substance abuse/opioid dependence
- If used in patients with medical comorbidities or if used concurrently with a benzodiazepine or other drugs that could potentially cause drug-drug interactions, the prescriber has acknowledged that they have completed an assessment of increased risk for respiratory depression.
- Pain is moderate to severe and expected to persist for an extended period of time
- Pain is chronic
- Pain is not postoperative (unless the patient is already receiving chronic opioid therapy prior to surgery, or if the postoperative pain is expected to be moderate to severe and persist for an extended period of time)
- Pain management is required around the clock with a long acting opioid

-AND-

(2) **ONE** of the following:

(a) **ALL** of the following:

i. The patient is being treated for moderate to severe chronic
pain that **is non-neuropathic** (examples of neuropathic pain include neuralgias, neuropathies, fibromyalgia).

-AND-

ii. Prior to the start of therapy with the long-acting opioid, the patient has failed an adequate (minimum of 4 week) trial of a short-acting opioid. (Document drug(s), duration and date of trial).b

-AND-

iii. The patient has a history of failure, contraindication or intolerance to a trial of **ALL** of the following (Document dates of trial):
   a) Nucynta ER
   b) morphine sulfate controlled-release tablets (specifically generic MS Contin)
   c) Xtampza ER

-OR-

(b) **ALL** of the following:

i. The patient is being treated for moderate to severe **neuropathic pain or fibromyalgia**

-AND-

ii. Unless it is contraindicated, the patient has not exhibited an adequate response to 8 weeks of treatment with gabapentin titrated to a therapeutic dose.\(^b\) (Document dose, duration and date of trial)

-AND-

iii. Unless it is contraindicated, the patient has not exhibited an adequate response to at least 6 weeks of treatment with a tricyclic antidepressant titrated to the maximum tolerated dose.\(^b\) (Document drug, dose, date and duration of trial).
iv. The patient has a history of failure, contraindication or intolerance to a trial of **ALL** of the following (Document dates of trial):

1) Nucynta ER
2) morphine sulfate controlled-release tablets (specifically generic MS Contin)
3) Xtampza ER

Authorization will be issued for 6 months for non-cancer and non-end of life pain up to the dose allowed by supply limit review (please refer to supply limit criteria). If the member is **currently** taking the requested long-acting opioid OR was recently switched from another long-acting opioid and does not meet the medical necessity initial authorization criteria requirements for long-acting opioids, a denial should be issued and a maximum 90-day authorization may be authorized one time for the requested drug/strength combination up to the requested quantity for transition to an alternative treatment.

2. Reauthorization

a. **Arymo ER**, **Avinza**, **Embeda**, **Exalgo**, **Dolophine**, **Duragesic**, fentanyl transdermal patch (37.5, 62.5, 87.5 mcg/hr), **Hysingla ER**, **Kadian**, methadone, **Morphabond ER**, **MS Contin, Nucynta ER, , OxyContin**, oxycodone controlled-release (Authorized Generic for OxyContin), oxymorphone ER **Xtampza ER** and **Zohydro ER** [Applies to all brand and generic versions of listed products)] will be reauthorized based on the following criteria:

1) Patient demonstrates meaningful improvement in pain and function (Document improvement in function or pain score improvement)

   -**AND**-

2) Identify rationale for not tapering and discontinuing opioid. (Document rationale).

   -**AND**-

3) Prescriber attests to **ALL** of the following:
   - Treatment goals are defined, including estimated duration of
• Treatment plan includes the use of a non-opioid analgesic and/or non-pharmacologic intervention
• Patient has been screened for substance abuse/opioid dependence
• If used in patients with medical comorbidities or if used concurrently with a benzodiazepine or other drugs that could potentially cause drug-drug interactions, the prescriber has acknowledged that they have completed an assessment of increased risk for respiratory depression.
• The information provided is true and accurate to the best of their knowledge and they understand that UnitedHealthcare may perform a routine audit and request the medical information necessary to verify the accuracy of the information provided.
• Pain is moderate to severe and expected to persist for an extended period of time
• Pain is chronic
• Pain is not postoperative (unless the patient is already receiving chronic opioid therapy prior to surgery, or if the postoperative pain is expected to be moderate to severe and persist for an extended period of time)
• Pain management is required around the clock with a long acting opioid

Authorization will be issued for 6 months for non-cancer and non-end of life pain up to the dose allowed by supply limit review (please refer to supply limit criteria). If the patient is currently taking the requested long-acting opioid OR was recently switched from another long-acting opioid and does not meet the medical necessity reauthorization criteria requirements for long-acting opioids, a denial should be issued and a maximum 90-day authorization may be authorized one time for the requested drug/strength combination up to the requested quantity for transition to an alternative treatment.

a. State mandates may apply. Any federal regulatory requirements and the member specific benefit plan coverage may also impact coverage criteria. Other policies and utilization management programs may apply.
b. For Kentucky business, only a 30-day trial will be required.
A. Cancer or End of Life (defined as a < 2 year life expectancy) related pain

1. Dolophine, fentanyl transdermal (generic Duragesic) 12, 25, 50, 75, 100 mcg/hr, methadone, morphine sulfate controlled-release (generic MS Contin), Nucynta ER and Xampza ER will be approved for cancer related pain based on the following criteria:

   a. Patient is being treated for cancer or end of life related pain

   -AND-

   b. Submission of medical records documenting active cancer diagnosis or life expectancy of < 2 years.

2. Arymo ER^, Avinza^, Duragesic^, Embeda^, Exalgo^, fentanyl transdermal (37.5, 62.5, 87.5 mcg/hr)^, Hysingla ER^, Kadian^, Morphabond ER^, MS Contin, OxyContin^, oxycodone controlled-release (Authorized Generic for OxyContin)^ oxymorphone extended release and Zohydro ER [Applies to all brand and generic versions of listed products except fentanyl transdermal patch (generic Duragesic) and morphine sulfate controlled-release tablets (generic MS Contin)] will be approved based on ALL of the following criteria:

   a. Patient is being treated for cancer or end of life related pain

   -AND-

   b. Submission of medical records documenting active cancer diagnosis or life expectancy of <2 years.

   -AND-

   c. The patient has a history of failure, contraindication or intolerance to a trial of the following (Document date of trial):

      (1) morphine sulfate controlled-release tablets (specifically generic MS Contin)

Authorization will be issued for 24 months up to the dose allowed by supply limit review (please refer to supply limit criteria). If the patient is currently taking the requested long-acting opioid OR was recently switched from
another long-acting opioid and does not meet the medical necessity initial
authorization criteria requirements for long-acting opioids, a denial should
be issued and a maximum 90-day authorization may be authorized one time
for the requested drug/strength combination up to the requested quantity
for transition to an alternative treatment.

B. Non-cancer and Non-End of Life pain

1. Dolophine, fentanyl transdermal (generic Duragesic) 12, 25, 50, 75, 100
mcg/hr, methadone, morphine sulfate controlled-release (generic MS
Contin), Nucynta ER and Xtampza ER will be approved based on the
following criteria:

a. Prescriber attests to ALL of the following:
   • Treatment goals are defined, including estimated duration of
treatment.
   • Treatment plan includes the use of a non-opioid analgesic and/or non-
pharmacologic intervention
   • Patient has been screened for substance abuse/opioid dependence
   • If used in patients with medical comorbidities or if used concurrently
with a benzodiazepine or other drugs that could potentially cause
drug-drug interactions, the prescriber has acknowledged that they
have completed an assessment of increased risk for respiratory
depression.
   • The information provided is true and accurate to the best of their
knowledge and they understand that UnitedHealthcare may perform a
routine audit and request the medical information necessary to verify
the accuracy of the information provided.
   • Pain is moderate to severe and expected to persist for an extended
period of time
   • Pain is chronic
   • Pain is not postoperative (unless the patient is already receiving
chronic opioid therapy prior to surgery, or if the postoperative pain is
expected to be moderate to severe and persist for an extended period
of time)
   • Pain management is required around the clock with a long acting
opioid

-AND-
b. **ONE** of the following:

(1) The patient is new to the plan (as evidenced by coverage effective date of less than or equal to 120 days) and is currently established on the requested long-acting opioid.

-OR-

(2) **ONE** of the following:

   i. **ALL** of the following:

      (a) The patient is being treated for moderate to severe chronic pain that is **non-neuropathic** (examples of neuropathic pain include neuralgias, neuropathies, fibromyalgia)

      -AND-

      (b) Prior to the start of therapy with the long-acting opioid, the patient has failed a four week trial of a generic short-acting opioid. (Document drug(s) and date of trial)

   -OR-

   ii. **ALL** of the following:

      (a) The patient is being treated for moderate to severe neuropathic pain or fibromyalgia

      -AND-

      (b) Unless it is contraindicated, the patient has not exhibited an adequate response to 4 weeks of treatment with gabapentin titrated to a therapeutic dose. (Document dose, duration, and date of trial)

      -AND-

      (c) Unless it is contraindicated, the patient has not exhibited an adequate response to 4 weeks of treatment with a generic tricyclic antidepressant titrated to the maximum tolerated dose. (Document drug, dose, date of trial)

b. **Arymo ER^, Avinza^, Duragesic^, Embeda^, Exalgo^, fentanyl**
transdermal (37.5, 62.5, 87.5 mcg/hr)^, Hysingla ER^, Kadian^, Morphabond ER^, MS Contin, OxyContin^, oxycodone controlled-release (Authorized Generic for OxyContin)^ oxymorphone extended release and Zohydro ER [Applies to all brand and generic versions of listed products except fentanyl transdermal patch (generic Duragesic) and morphine sulfate controlled-release tablets (generic MS Contin)] will be approved for non-cancer and non-end of life related pain based on the following criteria:

(1) Prescriber attests to ALL of the following:
  - Treatment goals are defined, including estimated duration of treatment.
  - Treatment plan includes the use of a non-opioid analgesic and/or non-pharmacologic intervention
  - Patient has been screened for substance abuse/opioid dependence
  - If used in patients with medical comorbidities or if used concurrently with a benzodiazepine or other drugs that could potentially cause drug-drug interactions, the prescriber has acknowledged that they have completed an assessment of increased risk for respiratory depression.
  - The information provided is true and accurate to the best of their knowledge and they understand that UnitedHealthcare may perform a routine audit and request the medical information necessary to verify the accuracy of the information provided.
  - Pain is moderate to severe and expected to persist for an extended period of time
  - Pain is chronic
  - Pain is not postoperative (unless the patient is already receiving chronic opioid therapy prior to surgery, or if the postoperative pain is expected to be moderate to severe and persist for an extended period of time)
  - Pain management is required around the clock with a long acting opioid

-AND-

(2) ONE of the following:

(a) ALL of the following:
i. The patient is being treated for moderate to severe chronic pain that is **non-neuropathic** (examples of neuropathic pain include neuralgias, neuropathies, fibromyalgia).

-AND-

ii. Prior to the start of therapy with the long-acting opioid, the patient has failed a four week trial of a generic short-acting opioid. (Document drug(s) and date of trial)<sup>a</sup>.

-AND-

iii. The patient has a history of failure, contraindication or intolerance to a trial of the following (Document date of trial)<sup>a</sup>:
   a) morphine sulfate controlled-release tablets (specifically generic MS Contin)

-OR-

(b) **ALL** of the following:

i. The patient is being treated for moderate to severe neuropathic pain or fibromyalgia

-AND-

ii. Unless it is contraindicated, the patient has not exhibited an adequate response to 4 weeks of treatment with gabapentin titrated to a therapeutic dose. (Document duration, dose and date of trial)<sup>a</sup>

-AND-

iii. Unless it is contraindicated, the patient has not exhibited an adequate response to 4 weeks of treatment with a generic tricyclic antidepressant titrated to the maximum tolerated dose. (Document drug, dose, and date of trial)<sup>a</sup>.

-AND-

iv. The patient has a history of failure, contraindication or intolerance to a trial of the following (Document date of trial)<sup>a</sup>: 
1) morphine sulfate controlled-release tablets (specifically generic MS Contin)

Authorization will be issued for 6 months for non-cancer and non-end of life pain up to the dose allowed by supply limit review (please refer to supply limit criteria). If the patient is currently taking the requested long-acting opioid OR was recently switched from another long-acting opioid and does not meet the medical necessity initial authorization criteria requirements for long-acting opioids, a denial should be issued and a maximum 90-day authorization may be authorized one time for the requested drug/strength combination up to the requested quantity for transition to an alternative treatment.

2. Reauthorization

a. Arymo ER^, Avinza^, Embeda^, Exalgo^, Dolophine, Duragesic^, fentanyl transdermal patch (37.5, 62.5, 87.5 mcg/hr)^, Hysingla ER^, Kadian^, methadone, Morphabond ER^, MS Contin, Nucynta ER, OxyContin^, oxycodone controlled-release (Authorized Generic for OxyContin)^, oxymorphone extended release, Xtampza ER and Zohydro ER [Applies to all brand and generic versions of listed products] will be reauthorized based on the following criteria:

(1) Patient demonstrates meaningful improvement in pain and function (Document improvement in function or pain score improvement)

-AND-

(2) Identify rationale for not tapering and discontinuing opioid. (Document rationale).

-AND-

(3) Prescriber attests to ALL of the following:
  - Treatment goals are defined, including estimated duration of treatment.
  - Treatment plan includes the use of a non-opioid analgesic and/or non-pharmacologic intervention
  - Patient has been screened for substance abuse/opioid dependence
  - If used in patients with medical comorbidities or if used
concurrently with a benzodiazepine or other drugs that could potentially cause drug-drug interactions, the prescriber has acknowledged that they have completed an assessment of increased risk for respiratory depression.

- The information provided is true and accurate to the best of their knowledge and they understand that UnitedHealthcare may perform a routine audit and request the medical information necessary to verify the accuracy of the information provided.
- Pain is moderate to severe and expected to persist for an extended period of time
- Pain is chronic
- Pain is not postoperative (unless the patient is already receiving chronic opioid therapy prior to surgery, or if the postoperative pain is expected to be moderate to severe and persist for an extended period of time)
- Pain management is required around the clock with a long acting opioid

Authorization will be issued for 6 months for non-cancer and non-end of life pain up to the dose allowed by supply limit review (please refer to supply limit criteria). If the patient is currently taking the requested long-acting opioid OR was recently switched from another long-acting opioid and does not meet the medical necessity reauthorization criteria requirements for long-acting opioids, a denial should be issued and a maximum 90-day authorization may be authorized one time for the requested drug/strength combination up to the requested quantity for transition to an alternative treatment.

a. For Connecticut only a trial/failure of generic medications is required. Trial may not be more than 30 days.

4. Coverage Criteria for the State of Florida:

   A. Cancer or End of Life (defined as a < 2 year life expectancy) related pain

   1. Dolophine, fentanyl transdermal (generic Duragesic) 12, 25, 50, 75, 100 mcg/hr, methadone, morphine sulfate controlled-release tablets(generic MS Contin), Nucynta ER and Xstampza ER will be approved for cancer related pain based on ALL of the following criteria:

      a. Patient is being treated for cancer or end of life related pain
b. Submission of medical records documenting active cancer diagnosis or life expectancy of < 2 years.

2. Arymo ER^, Avinza^, Duragesic^, Embeda^, Exalgo^, fentanyl transdermal (37.5, 62.5, 87.5 mcg/hr)^, Hysingla ER^, Kadian^, Morphabond ER^, MS Contin, Oxycontin^, oxycodone controlled-release^ (Authorized Generic for Oxycontin), oxymorphone extended release^ and Zohydro ER [Applies to all brand and generic versions of listed products except generic morphine sulfate controlled-release tablets (generic MS Contin) and fentanyl transdermal patch (generic Duragesic)] will be approved for non-cancer and non-end of life related pain based on ALL the following criteria:

a. Patient is being treated for cancer or end of life related pain -AND-

b. Submission of medical records documenting active cancer diagnosis or life expectancy of <2 years.

-AND-

c. ONE of the following:

(1) The patient has a history of failure, contraindication or intolerance to a trial of Xtampza ER. (Document date of trial).

-OR-

(2) If request is for Oxycontin or oxycodone controlled-release (Authorized generic for OxyContin) the patient requires more than 320 mg/day of controlled-release oxycodone.

Authorization will be issued for 24 months up to the dose allowed by supply limit review (please refer to supply limit criteria). If the patient is currently taking the requested long-acting opioid OR was recently switched from another long-acting opioid and does not meet the medical necessity initial authorization criteria requirements for long-acting opioids, a denial should be issued and a maximum 90-day authorization may be authorized one time for the requested drug/strength combination up to the requested quantity.
B. Non-cancer and Non-End of Life pain

1. Initial Authorization

a. Dolophine, fentanyl transdermal 12, 25, 50, 75, 100 mcg/hr, methadone, morphine sulfate controlled-release tablets (specifically generic MS Contin), Nucynta ER and Xtampza ER will be approved based on the following criteria:

(1) Prescriber attests to ALL of the following:
- Treatment goals are defined, including estimated duration of treatment.
- Treatment plan includes the use of a non-opioid analgesic and/or non-pharmacologic intervention
- Patient has been screened for substance abuse/opioid dependence
- If used in patients with medical comorbidities or if used concurrently with a benzodiazepine or other drugs that could potentially cause drug-drug interactions, the prescriber has acknowledged that they have completed an assessment of increased risk for respiratory depression.
- The information provided is true and accurate to the best of their knowledge and they understand that UnitedHealthcare may perform a routine audit and request the medical information necessary to verify the accuracy of the information provided.
- Pain is moderate to severe and expected to persist for an extended period of time
- Pain is chronic
- Pain is not postoperative (unless the patient is already receiving chronic opioid therapy prior to surgery, or if the postoperative pain is expected to be moderate to severe and persist for an extended period of time)
- Pain management is required around the clock with a long acting opioid

-AND-

(2) ONE of the following:
(a) The patient is new to the plan (as evidenced by coverage effective date of less than or equal to 120 days) and is currently established on the requested long-acting opioid.

-OR-

(b) **ONE** of the following:

i. **ALL** of the following:

   a) The patient is being treated for moderate to severe chronic pain that is **non-neuropathic** (examples of neuropathic pain include neuralgias, neuropathies, fibromyalgia).

   -AND-

   ii. Prior to the start of therapy with the long-acting opioid, the patient has failed an adequate (minimum of 4 week) trial of a short-acting opioid. (Document drug(s), duration and date of trial).

-OR-

(c) **ALL** of the following:

i. The patient is being treated for moderate to severe **neuropathic pain or fibromyalgia**

   -AND-

   ii. Unless it is contraindicated, the patient has not exhibited an adequate response to 8 weeks of treatment with gabapentin titrated to a therapeutic dose. (Document duration, dose and date of trial)

   -AND-

   iii. Unless it is contraindicated, the patient has not exhibited an adequate response to at least 6 weeks of treatment with a tricyclic antidepressant titrated to the maximum tolerated dose. (Document drug, dose, date and duration of trial).

b. **Arymo ER^, Avinza^, Duragesic^, Embeda^, Exalgo^, fentanyl**
transdermal (37.5, 62.5, 87.5 mcg/hr)^, Hysingla ER^, Kadian^, Morphabond ER^, MS Contin, Oxycontin^, oxycodone controlled-release^ (Authorized Generic for Oxycontin), oxymorphone extended release^ and Zohydro ER [Applies to all brand and generic versions of listed products except generic morphine sulfate controlled-release tablets (generic MS Contin) and fentanyl transdermal (generic Duragesic)] will be approved for non-cancer and non-end of life related pain based on the following criteria:

(1) Prescriber attests to ALL of the following:

- Treatment goals are defined, including estimated duration of treatment.
- Treatment plan includes the use of a non-opioid analgesic and/or non-pharmacologic intervention
- Patient has been screened for substance abuse/opioid dependence
- If used in patients with medical comorbidities or if used concurrently with a benzodiazepine or other drugs that could potentially cause drug-drug interactions, the prescriber has acknowledged that they have completed an assessment of increased risk for respiratory depression.
- The information provided is true and accurate to the best of their knowledge and they understand that UnitedHealthcare may perform a routine audit and request the medical information necessary to verify the accuracy of the information provided.
- Pain is moderate to severe and expected to persist for an extended period of time
- Pain is chronic
- Pain is not postoperative (unless the patient is already receiving chronic opioid therapy prior to surgery, or if the postoperative pain is expected to be moderate to severe and persist for an extended period of time)
- Pain management is required around the clock with a long acting opioid

-AND-

(2) ONE of the following:

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21
(a) **ALL** of the following:

i. The patient is being treated for moderate to severe chronic pain that **is non-neuropathic** (examples of neuropathic pain include neuralgias, neuropathies, fibromyalgia)

-**AND**-

ii. Prior to the start of therapy with the long-acting opioid, the patient has failed an adequate (minimum of 4 week) trial of a short-acting opioid. (Document drug(s), duration and date of trial).

-**AND**-

iii. The patient has a history of failure, contraindication or intolerance to a trial of Xtampza ER (Document date of trial).

-**OR**-

(b) **ALL** of the following:

i. The patient is being treated for moderate to severe **neuropathic pain or fibromyalgia**

-**AND**-

ii. Unless it is contraindicated, the patient has not exhibited an adequate response to 8 weeks of treatment with gabapentin titrated to a therapeutic dose. (Document duration, dose and date of trial)

-**AND**-

iii. Unless it is contraindicated, the patient has not exhibited an adequate response to at least 6 weeks of treatment with a tricyclic antidepressant titrated to the maximum tolerated dose. (Document drug, dose, date and duration of trial).

-**AND**-

iv. The patient has a history of failure, contraindication or
intolerance to a trial of Xtampza ER. (Document date of trial).

Authorization will be issued for 6 months for non-cancer and non-end of life pain up to the dose allowed by supply limit review (please refer to supply limit criteria). If the patient is currently taking the requested long-acting opioid OR was recently switched from another long-acting opioid and does not meet the medical necessity initial authorization criteria requirements for long-acting opioids, a denial should be issued and a maximum 90-day authorization may be authorized one time for the requested drug/strength combination up to the requested quantity for transition to an alternative treatment.

2. Reauthorization

a. Arymo ER^, Avinza^, Dolophine, Duragesic, Embeda, Exalgo^, fentanyl transdermal (37.5, 62.5, 87.5 mcg/hr)^, Hysingla ER^, Kadian^, methadone, Morphabond ER^, MS Contin, Nucynta ER, Oxycontin^, oxycodone controlled-release^ (Authorized Generic for Oxycontin), oxymorphone extended release, Xtampza ER and Zohydro ER [Applies to all brand and generic versions of listed products] will be reauthorized based on ALL of the following criteria:

   (1) Patient demonstrates meaningful improvement in pain and function (Document improvement in function or pain score improvement)

   -AND-

   (2) Identify rationale for not tapering and discontinuing opioid. (Document rationale).

   -AND-

   (3) Prescriber attests to ALL of the following:

   • Treatment goals are defined, including estimated duration of treatment.

   • Treatment plan includes the use of a non-opioid analgesic and/or non-pharmacologic intervention

   • Patient has been screened for substance abuse/opioid dependence

   • If used in patients with medical comorbidities or if used
concurrently with a benzodiazepine or other drugs that could potentially cause drug-drug interactions, the prescriber has acknowledged that they have completed an assessment of increased risk for respiratory depression.

- The information provided is true and accurate to the best of their knowledge and they understand that UnitedHealthcare may perform a routine audit and request the medical information necessary to verify the accuracy of the information provided.
- Pain is moderate to severe and expected to persist for an extended period of time
- Pain is chronic
- Pain is not postoperative (unless the patient is already receiving chronic opioid therapy prior to surgery, or if the postoperative pain is expected to be moderate to severe and persist for an extended period of time)
- Pain management is required around the clock with a long-acting opioid

Authorization will be issued for 6 months for non-cancer and non-end of life pain up to the dose allowed by supply limit review (please refer to supply limit criteria). If the patient is currently taking the requested long-acting opioid OR was recently switched from another long-acting opioid and does not meet the medical necessity reauthorization criteria requirements for long-acting opioids, a denial should be issued and a maximum 90-day authorization may be authorized one time for the requested drug/strength combination up to the requested quantity for transition to an alternative treatment.

5. Coverage Criteria for State of Maryland

A. Cancer or End of Life (define as a < 2 year life expectancy) related pain

1. Dolophine, Embeda, fentanyl transdermal (generic Duragesic) 12, 25, 50, 75, 100 mcg/hr, methadone, Xtampza ER, Nucynta ER and, morphine sulfate controlled-release tablet (generic MS Contin) will be approved for cancer related pain based on ALL of the following criteria:

   a. Patient is being treated for cancer or end of life related pain

   -AND-

   b. Submission of medical records documenting active cancer diagnosis or
life expectancy of < 2 years.

2. **Arymo ER^, Avinza^, Duragesic Exalgo^, fentanyl transdermal (37.5, 62.5, 87.5 mcg/hr)^, Hysingla ER^, Kadian^, Morphabond ER^, MS Contin, oxymorphone extended release^ and Zohydro ER [Applies to all brand and generic versions of listed products except generic fentanyl transdermal (generic Duragesic) and generic morphine sulfate controlled-release tablets (generic MS Contin) will be approved based on **ALL** of the following criteria:

a. Patient is being treated for cancer or end of life related pain

-AND-

b. Submission of medical records documenting active cancer diagnosis or life expectancy of < 2 years.

-AND-

c. The patient has a history of failure, contraindication or intolerance to a trial of at least **three** of the following (Document drug name and date of trial):

  (a) Nucynta ER
  (b) morphine sulfate controlled-release tablets (specifically generic MS Contin)
  (c) Embeda
  (d) Xtampza ER

3. **OxyContin^ and oxycodone controlled-release (Authorized Generic for OxyContin)^** will be approved based on **ALL** of the following criteria:

a. Patient is being treated for cancer or end of life related pain

-AND-

b. Submission of medical records documenting active cancer diagnosis or life expectancy of <2 years.

-AND-

c. The patient has a history of failure, contraindication or intolerance to a trial of at least **TWO** of the following (Document drug name date of trial):
a) Nucynta ER  
b) morphine sulfate controlled-release tablets (specifically generic MS Contin)  
c) Embeda

-AND-

d. **ONE** of the following:

(a) The patient has a history of failure, contraindication or intolerance to a trial of **Xtampza ER** (Document date of trial).

-OR-

(b) The patient requires more than 320 mg/day of controlled-release oxycodone.

Authorization will be issued for 24 months up to the dose allowed by supply limit review (please refer to supply limit criteria). If the patient is currently taking the requested long-acting opioid OR was recently switched from another long-acting opioid and does not meet the medical necessity initial authorization criteria requirements for long-acting opioids, a denial should be issued and a maximum 90-day authorization may be authorized one time for the requested drug/strength combination up to the requested quantity for transition to an alternative treatment.

B. Non-cancer and Non-End of Life pain

1. Initial Authorization

a. **Dolophine, fentanyl transdermal (generic Duragesic)** 12, 25, 50, 75, 100 mcg/hr, **Embeda, Xtampza ER, Nucynta ER, methadone and morphine sulfate controlled-release (generic MS Contin)** will be approved based on the following criteria:

(1) Prescriber attests to **ALL** of the following:

- Treatment goals are defined, including estimated duration of treatment.
- Treatment plan includes the use of a non-opioid analgesic and/or non-pharmacologic intervention
- Patient has been screened for substance abuse/opioid dependence
- If used in patients with medical comorbidities or if used
concurrently with a benzodiazepine or other drugs that could potentially cause drug-drug interactions, the prescriber has acknowledged that they have completed an assessment of increased risk for respiratory depression.

- The information provided is true and accurate to the best of their knowledge and they understand that UnitedHealthcare may perform a routine audit and request the medical information necessary to verify the accuracy of the information provided.

- Pain is moderate to severe and expected to persist for an extended period of time

- Pain is chronic

- Pain is not postoperative (unless the patient is already receiving chronic opioid therapy prior to surgery, or if the postoperative pain is expected to be moderate to severe and persist for an extended period of time)

- Pain management is required around the clock with a long acting opioid

   -AND-

(2) **ONE** of the following:

   (a) The patient is new to the plan (as evidenced by coverage effective date of less than or equal to 120 days) and is currently established on the requested long-acting opioid.

   -OR-

   (b) **ONE** of the following:

      i. **ALL** of the following:

         a) The patient is being treated for moderate to severe chronic pain that is **non-neuropathic** (examples of neuropathic pain include neuralgias, neuropathies, fibromyalgia).

      -AND-

      b) Prior to the start of therapy with the long-acting opioid, the patient has failed an adequate (minimum of 4 week) trial of
a short-acting opioid. (Document drug(s), duration and date of trial).

-OR-

ii. **ALL** of the following:

a) The patient is being treated for moderate to severe *neuropathic pain or fibromyalgia*

-AND-

b) Unless it is contraindicated, the patient has not exhibited an adequate response to 8 weeks of treatment with gabapentin titrated to a therapeutic dose. (Document duration, dose and date of trial)

-AND-

c) Unless it is contraindicated, the patient has not exhibited an adequate response to at least 6 weeks of treatment with a tricyclic antidepressant titrated to the maximum tolerated dose. (Document drug, dose, date and duration of trial).

b. **Arymo ER^, Avinza^, Duragesic, Exalgo^, fentanyl transdermal (37.5, 62.5, 87.5 mcg/hr)^, Hysingla ER^, Kadian^, Morphabond ER^, MS Contin, OxyContin^, oxycodone controlled-release (Authorized Generic for OxyContin)^oxymorphone extended release^ and Zohydro ER [Applies to all brand and generic versions of listed products except generic fentanyl transdermal (generic Duragesic) and generic morphine sulfate controlled-release tablets (generic MS Contin)]** will be approved for non-cancer and non-end of life related pain based on the following criteria:

(1) Prescriber attests to **ALL** of the following:

- Treatment goals are defined, including estimated duration of treatment.
- Treatment plan includes the use of a non-opioid analgesic and/or non-pharmacologic intervention
- Patient has been screened for substance abuse/opioid dependence
- If used in patients with medical comorbidities or if used concurrently with a benzodiazepine or other drugs that could
potentially cause drug-drug interactions, the prescriber has acknowledged that they have completed an assessment of increased risk for respiratory depression.

- The information provided is true and accurate to the best of their knowledge and they understand that UnitedHealthcare may perform a routine audit and request the medical information necessary to verify the accuracy of the information provided.
- Pain is moderate to severe and expected to persist for an extended period of time
- Pain is chronic
- Pain is not postoperative (unless the patient is already receiving chronic opioid therapy prior to surgery, or if the postoperative pain is expected to be moderate to severe and persist for an extended period of time)
- Pain management is required around the clock with a long acting opioid

-AND-

(2) **ONE** of the following:

(a) **ALL** of the following:

i. The patient is being treated for moderate to severe chronic pain that is **non-neuropathic** (examples of neuropathic pain include neuralgias, neuropathies, fibromyalgia).

-AND-

ii. Prior to the start of therapy with the long-acting opioid, the patient has failed an adequate (minimum of 4 week) trial of a short-acting opioid. (Document drug(s), duration and date of trial).

-AND-

iii. The patient has a history of failure, contraindication or intolerance to a trial of at least **THREE** of the following (Document drugs, dates of trial):
   a) Nucynta ER
   b) morphine sulfate controlled-release tablets (specifically
(b) **ALL** of the following:

i. The patient is being treated for moderate to severe **neuropathic pain or fibromyalgia**

-AND-

ii. Unless it is contraindicated, the patient has not exhibited an adequate response to 8 weeks of treatment with gabapentin titrated to a therapeutic dose. (Document duration, dose and date of trial)

-AND-

iii. Unless it is contraindicated, the patient has not exhibited an adequate response to at least 6 weeks of treatment with a tricyclic antidepressant titrated to the maximum tolerated dose. (Document drug, dose, date and duration of trial).

-AND-

iv. The patient has a history of failure, contraindication or intolerance to a trial of at least **three** of the following: (Document drugs and date of trial):

a) Nucynta ER
b) morphine sulfate controlled-release tablets (specifically generic MS Contin)
c) Embeda
d) Xtampza ER

Authorization will be issued for 6 months for non-cancer and non-end of life pain up to the dose allowed by supply limit review (please refer to supply limit criteria). If the patient is currently taking the requested long-acting opioid OR was recently switched from another long-acting opioid and does not meet the medical necessity initial authorization criteria requirements for long-acting opioids, a denial should be issued and a maximum 60-day authorization may be authorized one time for the requested drug/strength.
combination up to the requested quantity for transition to an alternative treatment.

2. Reauthorization
   a. Arymo ER\(^\text{®}\), Avinza\(^\text{®}\), Dolophine, Duragesic\(^\text{®}\), Embeda, Exalgo\(^\text{®}\), fentanyl transdermal (37.5, 62.5, 87.5 mcg/hr)\(^\text{®}\), Hysingla ER\(^\text{®}\), Kadian\(^\text{®}\), methadone, Morphabond ER\(^\text{®}\), MS Contin, Nucynta ER, OxyContin\(^\text{®}\), oxycodone controlled-release (Authorized Generic for OxyContin)\(^\text{®}\), Oxymorphone ER, Xtampza ER and Zohydro ER [Applies to all brand and generic versions of listed products] will be reauthorized based on the following criteria:

   (1) Patient demonstrates meaningful improvement in pain and function
       (Document improvement in function or pain score improvement)

   -AND-

   (2) Identify rationale for not tapering and discontinuing opioid.
       (Document rationale).

   -AND-

   (3) Prescriber attests to ALL of the following:
       • Treatment goals are defined, including estimated duration of treatment.
       • Treatment plan includes the use of a non-opioid analgesic and/or non-pharmacologic intervention
       • Patient has been screened for substance abuse/opioid dependence
       • If used in patients with medical comorbidities or if used concurrently with a benzodiazepine or other drugs that could potentially cause drug-drug interactions, the prescriber has acknowledged that they have completed an assessment of increased risk for respiratory depression.
       • The information provided is true and accurate to the best of their knowledge and they understand that UnitedHealthcare may perform a routine audit and request the medical information necessary to verify the accuracy of the information provided.
       • Pain is moderate to severe and expected to persist for an extended period of time
       • Pain is chronic
- Pain is not postoperative (unless the patient is already receiving chronic opioid therapy prior to surgery, or if the postoperative pain is expected to be moderate to severe and persist for an extended period of time)
- Pain management is required around the clock with a long acting opioid

Authorization will be issued for 6 months for non-cancer and non-end of life pain up to the dose allowed by supply limit review (please refer to supply limit criteria). If the patient is currently taking the requested long-acting opioid OR was recently switched from another long-acting opioid and does not meet the medical necessity reauthorization criteria requirements for long-acting opioids, a denial should be issued and a maximum 90-day authorization may be authorized one time for the requested drug/strength combination up to the requested quantity for transition to an alternative treatment.

a. State mandates may apply. Any federal regulatory requirements and the patient specific benefit plan coverage may also impact coverage criteria. Other policies and utilization management programs may apply.

6. Coverage Criteria for the State of Massachusetts (Refer to Section 2)

7. Coverage Criteria for the State of West Virginiaa

A. Cancer or End of Life (defined as a < 2 year life expectancy) related pain

1. Dolophine, Embeda, Hysingla ER, fentanyl transdermal (generic Duragesic) 12, 25, 50, 75, 100 mcg/hr, methadone, morphine sulfate controlled-release tablet (specifically generic MS Contin), X tam pza ER, Nucynta ER will be approved for cancer related pain based on ALL of the following criteria:

   a. Patient is being treated for cancer or end of life related pain

   -AND-

   b. Submission of medical records documenting active cancer diagnosis or life expectancy of < 2 years.

2. Arymo ER^, Avinza^, Duragesic, Exalgo^, fentanyl transdermal (37.5, 62.5, 87.5 mcg/hr)^, Kadian^, Morphabond ER^, MS Contin, OxyContin^, oxycodone controlled-release (Authorized Generic for
OxyContin™, oxymorphone extended-release, and Zohydro ER [Applies to all brand and generic versions of listed products except generic fentanyl transdermal (generic Duragesic) and generic morphine sulfate controlled-release tablets (generic MS Contin)] will be approved based on ALL of the following criteria:

a. Patient is being treated for cancer or end of life related pain

-AND-

b. Submission of medical records documenting active cancer diagnosis or life expectancy of <2 years.

-AND-

c. ONE of the following

(1) The patient has a history of failure, contraindication or intolerance to a trial of Xtampza ER. (Document date of trial).

-OR-

(2) For OxyContin™ and oxycodone controlled-release (Authorized Generic for OxyContin™) the patient requires more than 320 mg/day of controlled-release oxycodone.

Authorization will be issued for 24 months up to the dose allowed by supply limit review (please refer to supply limit criteria). If the patient is currently taking the requested long-acting opioid OR was recently switched from another long-acting opioid and does not meet the medical necessity initial authorization criteria requirements for long-acting opioids, a denial should be issued and a maximum 90-day authorization may be authorized one time for the requested drug/strength combination up to the requested quantity for transition to an alternative treatment.

B. Non-cancer and Non-End of Life pain

1. Initial Authorization

a. Dolophine, Embeda, fentanyl transdermal (generic Duragesic) 12, 25, 50, 75, 100 mcg/hr, Hysingla ER, methadone, morphine sulfate controlled-release tablet (specifically generic MS Contin), Xtampza
ER, and Nucynta ER will be approved based on the following criteria:

(1) Prescriber attests to **ALL** of the following:
- Treatment goals are defined, including estimated duration of treatment.
- Treatment plan includes the use of a non-opioid analgesic and/or non-pharmacologic intervention
- Patient has been screened for substance abuse/opioid dependence
- If used in patients with medical comorbidities or if used concurrently with a benzodiazepine or other drugs that could potentially cause drug-drug interactions, the prescriber has acknowledged that they have completed an assessment of increased risk for respiratory depression.
- The information provided is true and accurate to the best of their knowledge and they understand that UnitedHealthcare may perform a routine audit and request the medical information necessary to verify the accuracy of the information provided.
- Pain is moderate to severe and expected to persist for an extended period of time
- Pain is chronic
- Pain is not postoperative (unless the patient is already receiving chronic opioid therapy prior to surgery, or if the postoperative pain is expected to be moderate to severe and persist for an extended period of time)
- Pain management is required around the clock with a long acting opioid

-AND-

(2) **ONE** of the following:

(a) The patient is new to the plan (as evidenced by coverage effective date of less than or equal to 120 days) and is currently established on the requested long-acting opioid.

-OR-

(b) **ONE** of the following:

i. **ALL** of the following:
a) The patient is being treated for moderate to severe chronic pain that is non-neuropathic (examples of neuropathic pain include neuralgias, neuropathies, fibromyalgia).

-AND-

b) Prior to the start of therapy with the long-acting opioid, the patient has failed an adequate (minimum of 4 week) trial of a short-acting opioid. (Document drug(s), duration and date of trial).

-OR-

ii. ALL of the following:

a) The patient is being treated for moderate to severe neuropathic pain or fibromyalgia

-AND-

b) Unless it is contraindicated, the patient has not exhibited an adequate response to 8 weeks of treatment with gabapentin titrated to a therapeutic dose. (Document duration, dose and date of trial)

-AND-

c) Unless it is contraindicated, the patient has not exhibited an adequate response to at least 6 weeks of treatment with a tricyclic antidepressant titrated to the maximum tolerated dose. (Document drug, dose, date and duration of trial).

b. Arymo ER\textsuperscript{\textregistered}, Avinza\textsuperscript{\textregistered}, Duragesic, Exalgo\textsuperscript{\textregistered}, fentanyl transdermal (37.5, 62.5, 87.5 mcg/hr)\textsuperscript{\textregistered}, Kadian\textsuperscript{\textregistered}, Morphabond ER\textsuperscript{\textregistered}, MS Contin, OxyContin\textsuperscript{\textregistered}, oxycodone controlled-release (Authorized Generic for OxyContin)\textsuperscript{\textregistered}, oxymorphone extended-release and Zohydro ER [Applies to all brand and generic versions of listed products except generic fentanyl transdermal (generic Duragesic) and generic morphine sulfate controlled-release tablets (generic MS Contin)] will be approved based on ALL of the following criteria:

(1) Prescriber attests to ALL of the following:

- Treatment goals are defined, including estimated duration of
- Treatment plan includes the use of a non-opioid analgesic and/or non-pharmacologic intervention
- Patient has been screened for substance abuse/opioid dependence
- If used in patients with medical comorbidities or if used concurrently with a benzodiazepine or other drugs that could potentially cause drug-drug interactions, the prescriber has acknowledged that they have completed an assessment of increased risk for respiratory depression.
- The information provided is true and accurate to the best of their knowledge and they understand that UnitedHealthcare may perform a routine audit and request the medical information necessary to verify the accuracy of the information provided.
- Pain is moderate to severe and expected to persist for an extended period of time
- Pain is chronic
- Pain is not postoperative (unless the patient is already receiving chronic opioid therapy prior to surgery, or if the postoperative pain is expected to be moderate to severe and persist for an extended period of time)
- Pain management is required around the clock with a long acting opioid

-AND-

(2) **ONE** of the following:

(a) **ALL** of the following:

i. The patient is being treated for moderate to severe chronic pain that is **non-neuropathic** (examples of neuropathic pain include neuralgias, neuropathies, fibromyalgia)

-AND-

ii. Prior to the start of therapy with the long-acting opioid, the patient has failed an adequate (minimum of 4 week) trial of a short-acting opioid. (Document drug(s), duration and date of trial).

-AND-
iii. The patient has a history of failure, contraindication or intolerance to a trial of Xtampza ER (Document date of trial).

-OR-

(b) ALL of the following:

i. The patient is being treated for moderate to severe neuropathic pain or fibromyalgia

-AND-

ii. Unless it is contraindicated, the patient has not exhibited an adequate response to 8 weeks of treatment with gabapentin titrated to a therapeutic dose. (Document duration, dose and date of trial)

-AND-

iii. Unless it is contraindicated, the patient has not exhibited an adequate response to at least 6 weeks of treatment with a tricyclic antidepressant titrated to the maximum tolerated dose. (Document drug, dose, date and duration of trial).

-AND-

iv. The patient has a history of failure, contraindication or intolerance to a trial of Xtampza ER. (Document date of trial).

Authorization will be issued for 6 months for non-cancer and non-end of life pain up to the dose allowed by supply limit review (please refer to supply limit criteria). If the patient is currently taking the requested long-acting opioid OR was recently switched from another long-acting opioid and does not meet the medical necessity initial authorization criteria requirements for long-acting opioids, a denial should be issued and a maximum 90-day authorization may be authorized one time for the requested drug/strength combination up to the requested quantity for transition to an alternative treatment.

2. Reauthorization

a. Aynmo ER^, Avinza^, Dolophine, Duragesic, Embeda, Exalgo^,


fentanyl transdermal (37.5, 62.5, 87.5 mcg/hr)^, Hysingla ER, Kadian^, methadone, Morphabond ER^, MS Contin, Nucynta ER, OxyContin^, oxycodone controlled-release (Authorized Generic for OxyContin)^oxymorphone extended-release, Xtampza ER, and Zohydro ER [Applies to all brand and generic versions of listed products] will be approved based on ALL of the following criteria:

1. Patient demonstrates meaningful improvement in pain and function (Document improvement in function or pain score improvement)

   -AND-

2. Identify rationale for not tapering and discontinuing opioid. (Document rationale).

   -AND-

3. Prescriber attests to ALL of the following:
   - Treatment goals are defined, including estimated duration of treatment.
   - Treatment plan includes the use of a non-opioid analgesic and/or non-pharmacologic intervention
   - Patient has been screened for substance abuse/opioid dependence
   - If used in patients with medical comorbidities or if used concurrently with a benzodiazepine or other drugs that could potentially cause drug-drug interactions, the prescriber has acknowledged that they have completed an assessment of increased risk for respiratory depression.
   - The information provided is true and accurate to the best of their knowledge and they understand that UnitedHealthcare may perform a routine audit and request the medical information necessary to verify the accuracy of the information provided.
   - Pain is moderate to severe and expected to persist for an extended period of time
   - Pain is chronic
   - Pain is not postoperative (unless the patient is already receiving chronic opioid therapy prior to surgery, or if the postoperative pain is expected to be moderate to severe and persist for an
Pain management is required around the clock with a long acting opioid

Authorization will be issued for 6 months for non-cancer and non-end of life pain up to the dose allowed by supply limit review (please refer to supply limit criteria). If the patient is currently taking the requested long-acting opioid OR was recently switched from another long-acting opioid and does not meet the medical necessity reauthorization criteria requirements for long-acting opioids, a denial should be issued and a maximum 90-day authorization may be authorized one time for the requested drug/strength combination up to the requested quantity for transition to an alternative treatment.

State mandates may apply. Any federal regulatory requirements and the patient specific benefit plan coverage may also impact coverage criteria. Other policies and utilization management programs may apply.

3. Additional Clinical Rules:
   - Supply limits may be in place.
   - MEDLIMIT (Cumulative Opioid Review) is in place and can be utilized for individual supply limit reviews

^ Arymo ER, Avinza (brand only), Embeda, Exalgo (brand only), Duragesic Hysingla ER, fentanyl 37.5, 62.5 and 87.5 mcg/hr, Morphabond ER, Kadian (brand and generic), oxycodone controlled-release (authorized generic for OxyContin), and OxyContin are typically excluded from coverage. Tried/Failed criteria may be in place. Please refer to plan specifics to determine exclusion status.

4. References:
   2. Embeda Prescribing Information. Pfizer Inc. October 2014
   9. OxyContin Prescribing Information. Purdue Pharma, August 2015.

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<table>
<thead>
<tr>
<th>Program</th>
<th>Prior Authorization/Medical Necessity - Long-Acting Opioid Pain Medications</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Change Control</strong></td>
<td></td>
</tr>
<tr>
<td>Date</td>
<td>Change</td>
</tr>
<tr>
<td>2/2014</td>
<td>New program</td>
</tr>
<tr>
<td>4/2014</td>
<td>Removed step criteria</td>
</tr>
<tr>
<td>1/2015</td>
<td>Added additional products: Hysingla, MS Contin, hydromorphone and Oramorph. Added step criteria for cancer and non-cancer chronic pain with differentiation between neuropathic and non-neuropathic pain. Updated references to include new products’ prescribing information, the AAN position paper, and neuropathic pain treatment guidelines.</td>
</tr>
<tr>
<td>10/2015</td>
<td>Provided clarification regarding which brand and generic versions of listed products are included in the criteria (e.g. which generic morphine sulfate product is preferred and which are non-preferred). Added criteria for patients under the age of 18 years. Added state specific criteria for Maryland and Maine.</td>
</tr>
<tr>
<td>7/2016</td>
<td>Added Xtampza ER as preferred product. Added Indiana and West Virginia step therapy mandate.</td>
</tr>
</tbody>
</table>

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<table>
<thead>
<tr>
<th>Date</th>
<th>Changes</th>
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<tbody>
<tr>
<td>10/2016</td>
<td>Added AR state mandate to supply limit review section. Removed requirement for the Long-Acting Opioid Prior Authorization Fax Form. Revised reauthorization to include the request for all information collected from open-ended questions in lieu of Long-Acting Opioid Prior Authorization Fax Form.</td>
</tr>
<tr>
<td>12/2016</td>
<td>Changed taper allowance from a one-time authorization to a 60-day authorization. Added end of life diagnoses to cancer pain section. Updated supply limits section to allow pre-approval of higher strengths where applicable for dose consolidation. Removed ceiling limit for cancer and end of life diagnoses. Added CT footnote for the trial and failure of short-acting opioids for the book of business criteria.</td>
</tr>
<tr>
<td>1/2017</td>
<td>Added requirement for trial and failure of Xtampza ER prior to approval for OxyContin and oxycodone controlled-release for initial authorization and reauthorization criteria. Clarified that maximum 60-day fill should only be authorized one time.</td>
</tr>
<tr>
<td>3/2017</td>
<td>Added criteria for members new to the plan that should be reviewed as continuation of therapy for preferred products.</td>
</tr>
<tr>
<td>5/2017</td>
<td>Removed Opana ER as a preferred step one product. Added new product Arymo ER to criteria.</td>
</tr>
<tr>
<td>7/2017</td>
<td>Removed fentanyl transdermal as a preferred step one product. Updated reauthorization criteria to review instruments used to assess patients rather than specific scores. Removed requirement for provider attestation for cancer and end of life pain diagnoses. Added Morphabond ER, Troxyca ER, and Vantrela ER.</td>
</tr>
<tr>
<td>8/2017</td>
<td>Updated fentanyl supply limits.</td>
</tr>
<tr>
<td>2/2018</td>
<td>Added morphine sulfate ER (generic MS Contin), Duragesic, and methadone to the program. Added criteria for State of Connecticut. Revised provider attestation and added to initial authorization. Revised reauthorization criteria.</td>
</tr>
<tr>
<td>6/2018</td>
<td>Removed supply limit criteria. Will now utilize MEDLIMIT criteria. Removed Vantrela ER and Troxyca ER- products never brought to market.</td>
</tr>
</tbody>
</table>