Long-term efficacy and safety of oxycodone–naloxone prolonged release in geriatric patients with moderate-to-severe chronic noncancer pain: a 52-week open-label extension phase study

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Background: Two-thirds of older people suffer from chronic pain and finding valid treatment options is essential. In this 1-yearlong investigation, we evaluated the efficacy and safety of prolonged-release oxycodone–naloxone (OXN-PR) in patients aged ≥70 (mean 81.7) years.

Methods: In this open-label prospective study, patients with moderate-to-severe noncancer chronic pain were prescribed OXN-PR for 1 year. The primary endpoint was the proportion of patients who achieved ≥30% reduction in pain intensity after 52 weeks of treatment, without worsening bowel function. The scheduled visits were at baseline (T0), after 4 weeks (T4), and after 52 weeks (T52).

Results: Fifty patients completed the study. The primary endpoint was achieved in 78% of patients at T4 and 96% at T52 (P<0.0001). Pain intensity, measured on a 0–10 numerical rating scale, decreased from 6.0 at T0 to 2.8 at T4 and to 1.7 at T52 (P<0.0001). Mean daily dose of oxycodone increased from 10 to 14.4 mg (T4) and finally to 17.4 mg (T52). Bowel Function Index from 35.1 to 28.7 at T52. No changes were observed in cognitive functions (Mini-Mental State Examination evaluation), while daily functioning improved (Barthel Index from 53.1 to 61.0, P<0.0001). The Screener and Opioid Assessment for Patients with Pain-Revised score at 52 weeks was 2.6 (standard deviation 1.6), indicating a low risk of aberrant medication-related behavior. In general, OXN-PR was well tolerated.

Conclusion: This study of the long-term treatment of chronic pain in a geriatric population with OXN-PR shows satisfying analgesic effects achieved with a stable low daily dose, coupled with a good safety profile and, in particular, with a reduction of constipation, often present during opioid therapy. Our findings support the indications of the American Geriatrics Society, suggesting the use of opioids to treat pain in older people not responsive to acetaminophen or nonsteroidal anti-inflammatory drugs.

Keywords: oxycodone/naloxone, geriatric, chronic noncancer pain, constipation

Introduction
Managing chronic pain is, in itself, a complex task due to the wide range of painful clinical pictures and the limits of therapeutic tools. The World Health Organization (WHO) first introduced the three-step analgesic ladder in 1986 with the aim of creating a framework for the rational use of analgesics. The ladder consists of the use of nonopioid analgesics for mild pain (step 1), “weak” opioids, like codeine and tramadol, associated generally with nonopioids for moderate pain (step 2), and “strong” opioids with or without nonopioids for strong pain (step 3).1 On one hand,
adherence to the analgesic ladder and other pain guidelines is relatively infrequent, and on the other hand, the clinical usefulness of step 2 medications (weak opioids such as tramadol, codeine, hydrocodone) has been challenged in the literature, and the routine use of step 2 medications may be associated with disadvantages. In a 2010 Cochrane review, opioid therapy for chronic noncancer pain (CNCP) was considered controversial due to concerns regarding long-term effectiveness and safety, particularly the risk of side effects, tolerance, dependence, or abuse. However, there are data, supported by weak evidence, showing that patients who were able to continue opioids for a long term experienced clinically significant pain relief. When CNCP affects older people, critical issues such as comorbidities, coprescription of medication, frailty, cognitive dysfunction, and chronic illness can further increase, and providing effective pain management requires particular care in the choice of drugs and doses. On this basis, in 1998, the American Geriatrics Society published a clinical practice guideline for the management of chronic pain in older populations, recommending a cautious approach to the use of opioids. Furthermore, current British guidelines for the treatment of pain in older people acknowledge the validity of skipping step 2 of the WHO’s pain ladder, according to which the use of strong opioids at low doses is a valid option for the treatment of moderate-to-severe pain not adequately controlled with acetaminophen (paracetamol) and/or nonsteroidal anti-inflammatory drugs (NSAIDs). During therapy with opioids, the most common adverse events are related to bowel dysfunction, and include constipation; straining; hard, dry stools; incomplete evacuation; bloating; and increased gastric reflux. In older patients, constipation naturally increases with age and a concomitant opioid treatment can worsen the problem. Given that chronic pain affects ~60% of people aged more than 65 years, the problem is imperative and requires a combination of three key aspects to be addressed: to relieve pain, to limit adverse events, and to pay attention to the advanced age.

The combination of oral prolonged-release oxycodone and naloxone (OXN-PR) in a fixed 2:1 ratio was developed to mitigate the gastrointestinal adverse effects of opioids, while preserving the aim to reduce pain. The pharmacological basis of this agonist–antagonist association has been widely described elsewhere. The clinical effects of the oral OXN-PR combination were shown in a number of controlled randomized trials and in observational studies including patients with cancer and noncancer-related chronic pain. These studies have provided evidence of effective analgesia and a significant reduction in the incidence of opioid-induced constipation, with an improvement in compliance and quality of life. Two 52-week extension phase open-label trials assessing the long-term efficacy and safety of OXN-PR in managing CNCP demonstrated stable relief of pain with tolerable side effects.

There are scarce data from the literature addressing pain treatment with OXN-PR in older patients. This study presents the results of the 52-week extension phase of a 4-week prospective observational study, evaluating the efficacy and tolerability of OXN-PR in older patients naïve to strong opioids, who were recruited by an Italian geriatric unit specialized in physical rehabilitation. Here, the long-term analgesic effects and safety of OXN-PR, together with the functional and cognitive status and the level of opioid dependence are analyzed.

Materials and methods

This was a single-center, noncontrolled, prospective observational study performed at Santa Margherita Institute of Pavia, Italy, to assess the long-term analgesic efficacy and safety of OXN-PR in older people with moderate-to-severe CNCP. The 4-week findings of the study have been previously reported. Briefly, this was an open-label prospective study assessing older patients naïve to strong opioids with moderate-to-severe chronic pain. Patients were prescribed OXN-PR at an initial dose of 10/5 mg daily for 4 weeks, and in case of insufficient analgesia, the initial daily dose could be increased gradually. Changes in cognitive state, daily functioning, quality of life, constipation, and other adverse events were assessed. Low-dose OXN-PR was found effective and well tolerated in the short term, without affecting the cognitive status and bowel function. The study was conducted in accordance with the principles of the Declaration of Helsinki and International Conference on Harmonization Guidelines for Good Clinical Practice. Policlinico San Matteo of Pavia Ethics Committee approved the study protocol and all patients provided written informed consent.

Patient population

Patients who completed the 4 weeks’ observation achieving at least partial pain control (no severe pain, i.e., numerical rating scale [NRS] <7) in the absence of major side effects entered the extension phase and received OXN-PR for up to 52 weeks. Furthermore, consecutive outpatients referred to the center from January to March 2014 and receiving a new prescription for OXN-PR were included in this 52-week observation.
Patients were eligible for inclusion in the study if they met the following criteria: age ≥70 years; presence of chronic pain lasting ≥3 months, average pain intensity (PI) score ≥4 measured on a 0–10 NRS; requiring around-the-clock WHO step 3 opioids (strong opioids) despite analgesic treatment (NSAIDs, acetaminophen alone or in combination with low doses of codeine); no previous treatment with around-the-clock strong opioids; and absence of contraindications to oxycodone use.

Patients with the following were excluded: inability to take oral medication; symptomatic cerebral metastases; known psychiatric disease; cognitive impairment or dementia not allowing appropriate pain assessment (scores <18 at Mini-Mental State Examination [MMSE; range: 0–30], indicating moderate-to-severe cognitive impairment); severe chronic obstructive pulmonary disease or other respiratory disease with hypoxemia and/or hypercapnia; paralytic ileus; moderate or severe liver insufficiency (total bilirubin ≥1.5 mg/dL); moderate or severe renal insufficiency (creatinine clearance <30 mL/min according to Cockcroft-Gault formula); or hypersensitivity to the active substances or to any of the excipients of the study medication.

Patients were evaluated at baseline (T0), 4 weeks (T4), and after 52 weeks (T52; final observation).

Assessment and endpoints
Demographic information and selected clinical data were recorded at baseline: age, sex, and previous treatments for pain. The PIs, assessed on a verbally administered 0–10 point NRS referred to the last 7 days, were recorded as pain at rest, pain during movement, pain during the day, and pain at nighttime, at baseline (T0), after 4 weeks (T4), and after 52 weeks (T52). A mean of the four PIs was reported as mean of each single day assessment for the four daily items.

Bowel function, evaluated by the Bowel Function Index (BFI) according to Rentz et al, was recorded at the same time points. The BFI, a measure of general bowel function recently validated as a reproducible tool that detects clinically meaningful changes in opioid-induced constipation, comprises scores ranging from 0 (free from symptoms) to 100 (most severe symptoms). In patients with chronic pain, normal bowel function is defined as a BFI of ≤29, and a ≥12-point change in BFI score represents a clinically meaningful change in constipation severity.

Cognitive state was assessed using the MMSE (normal value ≥25). Daily functioning, evaluated by the Barthel Index (the maximum score achievable, equal to 100, indicates that the patient can perform all activities independently) and the measure of interference of pain in the patient’s life, was evaluated at T0, T4, and T52 by seven items from the validated Italian version of the Brief Pain Inventory-Short Form (general activity, walking ability, normal work, moods, enjoyment of life, sleeping, and relations with other people) with an 11-point NRS for each item, ranging from 0= no impairment to 10= most severe impairment.

Safety evaluations were performed at T4 and T52 by recording the presence and severity (on a verbally administered 0–10 point NRS, where 0= absence and 10= extreme severity) of symptoms commonly related to opioid treatment (nausea/vomiting, dry mouth, dizziness, drowsiness, and pruritus) and any other treatment-related adverse events that occurred or worsened in intensity and/or frequency after the first intake of OXN-PR.

The patient’s risk for opioid abuse was assessed at T52 using the Screening and Opioid Assessment for Patients with Pain-Revised (SOAPP-R), a 24-item, self-administered questionnaire designed to predict aberrant medication-related behaviors among persons with chronic pain. Items are rated from 0= never to 4= very often, and summed to generate the total SOAPP-R score, which ranges from 0 to 96.

The starting dose of OXN-PR at T0 was 5/2.5 mg of naloxone twice a day. During the 4-week observation (T4), the daily dose could be modified up to a maximum of 10/5 mg twice a day in case of insufficient analgesia. To control pain, all patients were instructed to take a rescue dose of acetaminophen 500 mg when they did not achieve satisfactory pain control. In case of rescue medication dosing more than twice in 1 day, patients were instructed to contact our center. Any other medications needed for the treatment of any other concomitant medical condition, including laxatives, were continued. During the 52-week extension phase, all patients received open-label OXN-PR and the starting dose of OXN-PR was the effective analgesic dose of OXN that the patient received at the end of the previous 4-week observation. Dose titration was permitted to a maximum of 40 mg/day at the discretion of the investigator. Concomitant physical and rehabilitative therapy was allowed throughout the protocol.

The primary endpoint was the proportion of responders, namely, patients who achieved a ≥30% reduction in PI score (evaluated as mean of each single day assessment for the four daily items of the four pain measures) from baseline to 52 weeks of treatment with OXN-PR without a worsening of bowel function.

Secondary endpoints were: the values of BFI score, MMSE, Barthel Index, Brief Pain Inventory (BPI),
and SOAPP-R at T52, the proportion of patients reporting adverse events at T52, their severity, and the absolute daily dose of OXN-PR at T52.

Statistical analysis

In the descriptive analysis, absolute frequency was used for categorical variables and central trend and dispersion measurements (mean or median, standard deviations) were used for quantitative continuous variables. Normality of data distribution was verified by the Shapiro–Wilk test.

The significance of difference between pairs of continuous variables was evaluated by the Student’s t-test or the Wilcoxon test, as appropriate. Changes over time in categorical variables were evaluated with the Cochran’s Q test, the Friedman test, or repeated measures analysis of variance, when appropriate. A P-value of <0.05 was considered statistically significant. Analysis was performed with the Statistical Analysis Software version 9.2 (SAS Institute Inc., Cary, NC, USA).

Results

Among the 60 enrolled patients who started the 52-week observation, 50 patients (83.3%) completed the follow-up study. One patient experienced a severe adverse event (drowsiness) leading to OXN-PR discontinuation within the first week. Nine other patients discontinued treatment prematurely: patient disposition is reported in Figure 1. The demographic and clinical characteristics of the 60 patients included 36 subjects who were aged ≥80 years. The PI at rest was documented. In a similar way, the PI as a mean ± standard deviation, if not otherwise reported. The sum of percentages may not be equal to 100 due to rounding. *Expressed as mean of each single day assessment for the four daily items; †Pain interference composite score comprising the seven Brief Pain Inventory measures of interference of pain in the patient’s daily life.

Abbreviations: BFI, Bowel Function index; MMSE, Mini-Mental state examination; NRS, Numerical Rating Scale; NSAIDs, nonsteroidal anti-inflammatory drugs; WHO, World Health Organization.

score (25.7±2.7), Barthe Index score (53.9±13.9), and BPI mean score within the items recorded (6.0±1.6) were also reported.

Changes in PI at the three scheduled time points for patients who completed the follow-up are reported in Table 2. A substantial, statistically significant (P<0.0001), decrease of pain was documented. In a similar way, the PI as a mean of the four measures was significantly reduced.

Table 1 Demographic and baseline characteristics of the study population

<table>
<thead>
<tr>
<th>Demographic and baseline characteristics</th>
<th>N=60</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age, years (median; range)</td>
<td>81.7±15.7 (83; 70–94)</td>
</tr>
<tr>
<td>Patients &gt;80 years, n (%)</td>
<td>36 (60)</td>
</tr>
<tr>
<td>Females/males, n (%)</td>
<td>48/12 (80.0/20.0)</td>
</tr>
<tr>
<td>Pain intensity, mean NRS score†</td>
<td>6.7±1.2</td>
</tr>
<tr>
<td>Pain intensity at rest, NRS score</td>
<td>6.0±1.9</td>
</tr>
<tr>
<td>Pain intensity on movement, NRS score</td>
<td>8.1±0.9</td>
</tr>
<tr>
<td>Daily pain intensity, NRS score</td>
<td>7.6±1.0</td>
</tr>
<tr>
<td>Nocturnal pain intensity, NRS score</td>
<td>5.2±2.2</td>
</tr>
<tr>
<td>Previous analgesic treatment, n (%)</td>
<td></td>
</tr>
<tr>
<td>Step 1 WHO drugs</td>
<td></td>
</tr>
<tr>
<td>NSAIDs</td>
<td>7 (11.7)</td>
</tr>
<tr>
<td>Acetaminophen</td>
<td>39 (65.0)</td>
</tr>
<tr>
<td>WHO step 2 drugs</td>
<td></td>
</tr>
<tr>
<td>Codeine</td>
<td>5 (8.3)</td>
</tr>
<tr>
<td>Tramadol</td>
<td>17 (28.3)</td>
</tr>
<tr>
<td>Rescue analgesics, n (%)</td>
<td>51 (85.0)</td>
</tr>
<tr>
<td>Cognitive state, MMSE score (median)</td>
<td>25.7±2.7 (26)</td>
</tr>
<tr>
<td>Daily functioning, Barthel Index score</td>
<td>53.9±13.9 (55.0)</td>
</tr>
<tr>
<td>Brief Pain Inventory, composite score‡</td>
<td>6.0±1.6 (6.4)</td>
</tr>
<tr>
<td>Bowel function, BFI score (median)</td>
<td>35.9±19.1 (40.0)</td>
</tr>
</tbody>
</table>

Notes: All values are expressed as numbers (percent) or mean ± standard deviation, if not otherwise reported. The sum of percentages may not be equal to 100 due to rounding. *Expressed as mean of each single day assessment for the four daily items; †Pain interference composite score comprising the seven Brief Pain Inventory measures of interference of pain in the patient’s daily life.

Table 2 Pain intensity (NRS score) during the 52 weeks of oxycodone/naloxone prolonged-release treatment

<table>
<thead>
<tr>
<th>Parameter</th>
<th>T0</th>
<th>T4</th>
<th>T52</th>
<th>P-value*</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pain at rest, mean ± SD</td>
<td>6.0±1.9</td>
<td>2.8±1.7*</td>
<td>1.7±1.4**</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>Pain on movement, mean ± SD</td>
<td>8.0±1.0</td>
<td>4.4±2.0*</td>
<td>3.5±1.2***</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>Pain at daytime, mean ± SD</td>
<td>7.5±1.1</td>
<td>3.9±2.3*</td>
<td>3.4±1.3*</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>Pain at nighttime, mean ± SD</td>
<td>5.3±1.2</td>
<td>2.1±1.9*</td>
<td>1.1±1.3*</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>Pain intensity, mean ± SD‡</td>
<td>6.7±1.2</td>
<td>3.3±1.8*</td>
<td>2.4±1.1*</td>
<td>&lt;0.0001</td>
</tr>
</tbody>
</table>

Notes: Included in this analysis were all patients completing the 52-weeks treatment (n=50). *Expressed as mean of each single day assessment for the four daily items; †P52 weeks versus baseline; ‡P=0.0001 versus previous observation; ***P=0.0001 versus previous observation; ****P=0.00024 versus previous observation; ††P=0.0670 versus previous observation.

Abbreviations: NRS, Numerical Rating Scale; SD, standard deviation.
All patients started OXN-PR with a dose of 5/2.5 mg twice a day. During the follow-up, the dose was increased to 20/10 mg in 42% of patients at 4 weeks and to 40/20 mg at 52 weeks in only 6% of patients (Figure 2). Overall, the mean daily dose of oxycodone increased from 10 mg (T0) to 14.4±4.9 mg (T4) to the final 17.4±7.7 mg (T52) (P<0.0001).

The findings of the secondary endpoints are shown in Figure 3. The MMSE score remained stable during the follow-up; the Barthel Index increased slightly between T0 and T4 (P<0.0001) without further changes up to T52; BPI items improved during the first month of treatment (P<0.0001), remaining at the same level thereafter. The mean SOAPP-R score at 52 weeks was 2.6 (standard deviation 1.6), thus suggesting a low risk of aberrant medication-related behavior in our population.

The improvement in pain control was paralleled by a slight improvement in bowel function: mean BFI values decreased from 35.1 at baseline to 28.7 at T52 (P=0.0002) (Figure 4). An opposite trend was observed with regard to the proportion of patients receiving supplementary laxatives: 38% at T0, 53% at T4, and 29% at T52 (data not shown).

The proportion of the responders according to the primary endpoint, namely, patients who achieved a ≥30% reduction in PI in the absence of a bowel function worsening, was 78% at T4 and 96% at T52 (P<0.0001) (data not shown).

Figure 5 shows the prevalence and severity of commonly observed opioid-related side effects and tremor at different points of the observation. Nausea/vomiting and tremor gradually decreased during the follow-up. Dizziness, drowsiness, and pruritus only slightly decreased during the first 4 weeks and then increased a little up to 52 weeks. Dry mouth worsened slightly over the 52-week observation.

**Discussion**

Older patients exhibit a high incidence of chronic pain conditions, partially due to age-related alterations of the peripheral somatosensory system and increased activity of the central glial cells. Microglia are the primary modulators of pain neurons and their sensitization promotes a persistent condition of neuroinflammation. Aging is associated with impaired endogenous inhibitory systems with dysfunctional changes in pain modulatory capacity.

Due to these changes and the higher frailty of older people, chronic painful and disabling conditions are very common. Moreover, pharmacokinetic and pharmacodynamic variability is a major determinant of the dose–effect relationship in older patients and should also be considered when choosing the most appropriate analgesic regimen. Available analgesics include nonopioids such as acetaminophen, NSAIDs, cyclooxygenase-2 inhibitors, and opioids.

Acetaminophen has been recommended by the American Geriatrics Society clinical practice guideline and the...
American College of Rheumatology15–37 as the first-line agent in the treatment of mild-to-moderate chronic pain, given its moderate toxicity. Nevertheless, it has poor analgesic efficacy in severe pain, which limits its utilization. The toxicity, both renal and gastrointestinal, of NSAIDs is increased in older patients.3,39 Their prolonged use may compromise the renal function and increase the risk of cerebrovascular attacks, as well as congestive heart failure in patients with cardiovascular disease by tenfold. Cyclooxygenase-2 selective inhibitors offer improved gastrointestinal tolerance, but pose similar risk to renal and cardiovascular function.40 Among the opioids, tramadol can be indicated for moderate pain unrelieved by acetaminophen,41 with some limitations due to its low pharmacological potency, the presence of a ceiling effect, and the risk of a serotonergic syndrome. Finally, “strong opioids”, commonly used in cancer pain patients and the third step of WHO guidelines, are now increasingly employed for all types of pain including the chronic pain of musculoskeletal disease. Opioids also present some risks of negative effects, specifically possible poor responsiveness, side effects, tolerance, abuse, and addiction. While that does not prevent their use in CNCP, it suggests careful selection of the patients, and a monitoring of some clinical parameters during the treatment is necessary.

A strength and originality of our study consists in the combination of CNCP, geriatric patients (82 years of age on average), and long-term treatment (1 year). The objectives of therapy in this case were relief of pain, improvement of physical daily activity and quality of life in the absence of important side effects, and drug consumption not clinically targeted.

The 50 patients who completed the study achieved an important analgesic effect throughout 1 year of low-dose OXN-PR. The initial PI was halved at T4 and further reduced at T52. It should not be taken for granted that the efficacy was maintained for such an extended time. Opioid therapy often runs into tolerance, whose appearance is easily recognized by the drop of analgesic effect with a simultaneous compensatory dose escalation. In our study, we found no loss of analgesia and only a moderate increase of the daily dose over time (from 10 mg of oxycodone to 17.4±7.7 mg daily on average). This observation is in agreement with the results of other studies based on OXN therapy.42,43 The first of these, Lazzari et al,42 was a comparative study between prolonged-release oxycodone and OXN in opioid-naive cancer patients with moderate-to-severe pain. Analgesia was achieved and maintained with similarly low and stable dosages over 60 days (from 11.5 to 22.0 mg/day of oxycodone in the OXN group). Trenkwalder et al43 designed a study to evaluate the effects of OXN in the treatment of severe restless legs syndrome; the mean daily dose was 18.1 mg of oxycodone and 9.1 mg of naloxone for a median period of treatment of 281 days.

Stable doses of OXN-PR over time have been repeatedly reported for the treatment of CNCP in opioid-naive patients, after very low starting doses. Furthermore, the long-term steadiness of the opioid dose is associated with persistent analgesia and a limited number of treatment switches and discontinuations. Further studies providing additional contribution to the strategy of treatment of CNCP are required.

The low dose may also explain the good safety profile observed in our geriatric patients. The prevalence of typically expected adverse opioid reactions was at low levels. Only dry mouth tended to be particularly frequent (82%–90% of patients), which is also likely related to the advanced age.
Nausea and vomiting, on the contrary, were infrequent (28% at T4 and 18% at T52), probably due to the action of naloxone at the gastric level by the same mechanism involved at bowel level. Similar observations have also been made in two previous studies.18,42

Special attention has to be paid to the issue of constipation. In this study, the degree of constipation, measured by BFI, consistently decreased from the initial value of 35.1 to 31.6 (T4) and to the final value of 28.7 at T52. This last value, reached at T52, is deemed as a normal value in patients with chronic pain. Considering the age of patients and the prolonged treatment with opioids, this result can be viewed very positively. In parallel, cognitive functions were unequivocally not affected by long-term treatment with OXN, in contrast to the effects observed in studies that used other opioids.44,45 The daily activity tended to improve over time, and similarly, some quality of life parameters, detected by Brief Pain Inventory-Short Form, particularly referring to moods, enjoyment of life, sleeping, and relations with other people, became significantly better.

Of note and of primary relevance, our results confirm the low risk of addiction in older adults without a history of drug abuse, as confirmed by SOAPP-R assessment in the whole population study. An exceedingly low value was found at T52, well below the cutoff score of 18 traditionally used to identify subjects at increased risk of opioid abuse and misuse.

We utilized a composite endpoint as the primary endpoint. Combining efficacy and tolerability parameters (specifically, the proportion of patients with at least 30% reduction in PI, in the absence of a bowel function worsening), we focused on the two main goals of an effective opioid treatment for chronic pain. The proportion of responders was equivalent to 78% at T4 and 96% at T52, which can be considered an excellent outcome.

This study has several limitations related to the methodology used, specifically the observational design and the absence of a control group. Despite this, the 1-year follow-up and generalizability of the findings are its important strengths.

Conclusion
The findings of this study suggest that low doses of OXN-PR are a valid treatment for chronic moderate-to-severe pain in older patients. During 1 year of treatment, they achieved effective analgesia accompanied by few side effects, improvement of bowel function, and a good level of functional state and quality of life.

Data from the present open-label study confirm that in the older patients, a slow titration of opioids is mandatory to avoid adverse events; the rule “start slowly, go slowly” should always be considered when prescribing opioids to this fragile population.

Despite the aforementioned limitations of the present study, our findings suggest that OXN-PR should be considered as a potentially safe approach for managing pain and preventing opioid-induced constipation46 in older adults over the long term because of its particular pharmacological profile.

Further studies in older persons with CNCP, looking at the therapeutic approach and safety of OXN-PR will be of great relevance in a phase of critical evaluation of the long-term treatment of pain.

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Author contributions
All authors contributed in developing the concepts, designing the structure, and writing/revising the manuscript, and approved the final version before submission and agree to be accountable for all aspects of the work.

Disclosure
The authors report no conflicts of interest in this work.

References


