Long-term safety and efficacy of oxycodone/naloxone prolonged-release tablets in patients with moderate-to-severe chronic cancer pain

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Support Care Cancer (2015) 23:823–830
DOI 10.1007/s00520-014-2435-5

Abstract

Aim To evaluate the long-term safety and efficacy of prolonged-release oxycodone/naloxone (OXN PR) and its impact on quality of life (QoL), in patients with moderate-to-severe cancer pain.

Methods This was an open-label extension (OLE) of a 4 week, randomized, double-blind (DB) study in which patients with moderate-to-severe cancer pain had been randomized to OXN PR or oxycodone PR (OxyPR). During the OLE phase, patients were treated with OXN PR capsules (≤20/60 mg/day) for ≤24 weeks. Outcome measures included safety, efficacy and QoL.

Results One hundred and twenty-eight patients entered the OLE, average pain scores based on the modified Brief Pain Inventory—Short Form were low and stable over the 24-week period. The improvement in bowel function and constipation symptoms as measured by the Bowel Function Index and patient assessment of constipation in patients treated with OXN PR during the 4-week DB study was maintained. In patients treated with OxyPR during the DB phase, bowel function and constipation symptoms were improved during the OLE. In the DB and in the OLE, health status and QoL were similar for patients treated with OXN PR and OxyPR. There were no unexpected safety or tolerability issues.

Conclusions In patients with moderate-to-severe cancer pain, long-term use of OXN PR is well tolerated and effective, resulting in sustained analgesia, improved bowel function and improved symptoms of constipation.

Keywords Oxycodone · Naloxone · Opioid-induced constipation · Analgesia · Bowel function

Introduction

Pain is a common and debilitating symptom of cancer and can significantly impact the lives of patients through detrimental effects on physical functioning, psychological well-being and social interactions [1]. Pain affects a large proportion of cancer patients, with prevalence rates ranging from 33 % in patients after curative treatment to 64 % in patients with advanced, metastatic or terminal phase disease [2, 3]. In a 2009 European survey, 56 % of cancer patients were demonstrated to suffer pain several times per month, with over 90 % of these reporting their pain as moderate to severe in intensity [4].

Opioid analgesics are the established treatment for the relief of moderate-to-severe cancer pain and several different opioids are now available [5, 6]. For years, morphine has been the accepted gold standard for the treatment of cancer pain and is endorsed by the European Association of Palliative Care [6]. Oxycodone (a semi-synthetic opioid analgesic) has proven efficacy in the management of severe pain of different...
aetiologies [7–10]. A series of systematic reviews on the use of opioids in cancer patients found that oxycodone and hydromorphone have efficacy and tolerability profiles comparable with that of morphine [11, 12]. Although opioids have proven analgesic efficacy, their use is frequently complicated by a range of side effects, which include nausea, sedation, euphoria/dysphoria, constipation and itching [13, 14]. The most common and often most debilitating side effect is opioid-induced bowel dysfunction (OIBD), which comprises a constellation of gastrointestinal (GI) adverse events (AEs) including constipation, hard dry stools, straining, incomplete evacuation, bloating, abdominal cramping, abdominal distension and increased gastric reflux [13, 15]. The primary symptom of OIBD is constipation, which occurs in approximately 90 % of cancer patients receiving opioid therapy and persists over time [14, 16, 17]. Current strategies to manage OIBD are non-specific and frequently ineffective [18].

Increasing dietary fibre, fluid intake and physical mobility are usually insufficient to prevent or treat OIBD and the majority of patients receiving long-term opioid therapy require laxatives [13]. Laxatives do not address the underlying opioid receptor-mediated cause of constipation and evidence for their effectiveness is weak; therefore, OIC persists in many patients [13, 19]. A novel therapeutic approach to bypass the GI effects of opioids involves co-administration of an opioid antagonist (naloxone). When administered orally in a prolonged-release (PR) formulation, naloxone has a low systemic bioavailability (<3 %) and antagonises peripheral opioid receptors in the GI tract with little impact on centrally-acting opioid analgesia [20–22]. Oral naloxone can, therefore, improve OIC [23, 24]. In patients with moderate-to-severe non-cancer pain, randomized phase III trials have demonstrated that co-administration of prolonged-release (PR) oxycodone with naloxone (OXN PR) provides effective analgesia whilst significantly reducing the impact of OIBD and OIC compared with oxycodone PR alone (OxyPR) [25–27] and that this effect is maintained over the long term (≤52 weeks) [28]. Similar safety and efficacy has been demonstrated in a randomized phase II study of OxyPR vs OXN PR in patients with moderate-to-severe cancer pain [29]; although, the long-term effects on this patient population has not yet been demonstrated. Here, we report results of a long-term open-label extension (OLE) of the phase II study in which patients were maintained on or switched to OXN PR.

Methods

Study design

This was a 24 week, uncontrolled, open-label extension of a 4 week, randomized, double-blind study (OXN2001S; NCT00513656) [29] conducted to evaluate the long-term safety and efficacy of OXN PR vs OxyPR in patients with moderate-to-severe cancer pain. The study was performed in compliance with good clinical practice and in accordance with the Declaration of Helsinki.

Inclusion and exclusion criteria

Patients completing the double-blind period and those who had discontinued prematurely due to constipation were eligible to enter the OLE phase. Patients aged ≥18 years were enrolled into the double-blind phase of the study if they had moderate-to-severe chronic cancer pain that required ‘round-the-clock’ opioid therapy and constipation secondary to opioid therapy [29].

Study treatment

During the double-blind phase, participants were randomized to OXN PR or OxyPR for 4 weeks (≤120 mg/day). During the OLE, all patients received OXN PR every 12 h for ≤27 weeks. Study Visit 9a (Day 1) was the first visit of the OLE phase of the study and typically occurred on the same day as Visit 9 (end of the double-blind phase) (Figure 1A).

The initial starting dose was the effective analgesic dose based on oxycodone or OXN PR that the patient was on at the end of the double-blind phase. Any patient on a dose of ≤80 mg/day OxyPR was switched directly to OXN PR. For example, all patients on 80 mg/day OxyPR started with a daily dose of OXN PR 80/40 mg. Those patients on >80 mg/day OxyPR were switched to OXN PR in a stepwise manner. All patients receiving 90, 100, 110, or 120 mg/day OxyPR were started on OXN PR 40/20 mg 12 hourly with the remainder of their oxycodone dose given as OxyPR. Dose titration was permitted at the discretion of the investigator to ≤120 mg/day. A total of 19 patients received OXN PR at a daily dose of >80 mg/day; safety findings for these patients were no different from the whole population.

Rescue medications (Oxy IR 5 mg and laxative bisacodyl 5 mg) were supplied for the first 7 days of the study. After 7 days, laxative use was documented as concomitant medication.

Study assessments

Efficacy was evaluated based on the following measurements: Brief Pain Inventory—Short Form (BPI-SF), a patient-rated scale based on ‘least’, ‘average’ and ‘worst’ pain felt over the previous week [30]; Bowel Function Index (BFI) a comprising three components (ease of defecation, feeling of incomplete bowel evacuations and personal judgement of constipation) [31, 32]; patient assessment of constipation (PAC-SYM) [33]; rescue medications (Oxy IR and laxative); number of bowel movements; health status (EuroQol EQ-5D) and quality of life (QoL) (European Organisation for Research and Treatment of
Cancer Quality of Life Questionnaire-Core 30; EORTC QLQ-C30) [34, 35]. Safety assessments included adverse events (AEs) and serious AEs, physical and laboratory evaluations, 12-lead electrocardiograph, and evaluation of the symptoms of opiate withdrawal as measured by the Subjective Opiate Withdrawal Scale (SOWS) collected 1 day and 1 week following the start of the OLE [36].

Statistical analysis

All analyses were performed on the OLE safety population (patients who received at least one dose of study medication during the OLE. All efficacy and safety variables were summarized descriptively. All continuous variables were summarized using the following descriptive statistics: n, mean, standard deviation (SD), median, minimum and maximum. The frequency and percentage of observed levels were reported for all categorical measures.

Results

Patient characteristics

A total of 128 patients entered the OLE phase of which 126 (98.4 %) had previously completed the double-blind phase and 2 (1.6 %) had discontinued due to constipation. All 128 patients in the OLE phase received study medication and 68 patients (53.1 %) completed the study. Fifty-three per cent and 56.9 % of patients were male and female, respectively, and the mean age was 62.5 years. The most common cancers were breast (19.5 %), lung (8.6 %) and prostate (11.7 %) (Table 1).

Efficacy

OXN PR dose

Mean duration of exposure to OXN PR was 122.2 days and the mean daily dose was 56.2 mg. The majority of subjects (69.5 %) remained on their starting dose for the duration of the study. Thirty-four subjects progressed to a slightly higher dose...
Table 1 Patient demographics and clinical characteristics (safety population)

| Characteristic | OXN PR (N=128) |
|----------------|----------------|
| Mean age, years (SD) | 62.5 (10.3) |
| Age ≤65 years, n (%) | 74 (57.8) |
| Male, n (%) | 68 (53.1) |
| Race, n (%) | 127 (99.2) |
| Black | 1 (0.8) |
| BMI, kg/m² (n=123) | 25.3 (5.1) |
| Median (range) | 25.1 (16–40) |

BPI-SF average pain scores (LOCF) by study visit (OLE safety population).

Pain (BPI-SF)

At the end of the double-blind phase of the study in which patients were treated with OxyPR or OXN PR, there was no significant difference in pain control (as assessed by the BPI-SF) between the two treatments [29]. At the beginning of the OLE, BPI-SF pain scores for ‘average pain’ were low in both groups: mean (SD) 3.42 (2.03) and 3.63 (1.76) for OXN PR and OxyPR, respectively. BPI-SF scores remained consistently low and stable in both groups over the OLE period with scores of 3.71 (2.12) for OxyPR and 3.56 (2.27) for OXN PR at Week 24 (Figure 2).

Bowel function index

At the end of the double-blind phase, OXN PR is both statistically significantly superior in respect to BFI and has also demonstrated a change that is clinically relevant compared with OxyPR [29]. For patients that received OXN PR during the DB phase (OLE baseline score [SD]: 38.08 [26.94]), the improvement in BFI was maintained during the OLE (mean BFI: 37.90 [28.06] at Week 12 and 39.02 [27.70] at Week 24). For patients treated with OxyPR during the DB phase (OLE baseline score: 46.61 [26.84]), mean BFI scores were reduced to 38.52 (27.69) at Week 12 and 38.78 (27.45) at Week 24, comparable to those who had previously received OXN PR (Figure 3).

Constipation symptoms (PAC-SYM)

At the end of the double-blind phase, a statistically significantly greater improvement in constipation symptoms was observed for OXN PR compared with OxyPR [29]. At the beginning of the OLE phase, mean (SD) PAC-SYM total symptoms scores were 10.55 (8.60) and 15.53 (10.20) for OXN PR and OxyPR, respectively. The improvement in constipation symptoms seen in the OXN PR group was maintained during the OLE, with a mean PAC-SYM (SD) of 12.14 (9.65) at Week 24. For patients who had received OxyPR during the double-blind period, mean PAC-SYM scores were reduced to 12.81 (10.10) at Week 24 (Table 2).

Rescue medication

During the first 7 days of the OLE, the need for rescue medication was low. A total of 37.5 % of patients used laxatives. The mean bisacodyl dose in 7 days was 6.09 mg, which translates to approximately one tablet used in 7 days. A mean of 1.1 (range: 0–3.9) capsules of Oxy IR 5 mg were required for rescue analgesia per day.

QoL

In the double-blind phase of the study, health status and QoL were similar between OXN PR and OxyPR groups [29].
These results were maintained during the OLE. In the total OLE population, the mean (SD) EQ-5D index score was 0.49 (0.31) at day 1, 0.59 (0.25) at week 12 and 0.46 (0.38) at week 24. The EORTC QLQ-C30 global health status scores were 45.83 (20.36) at day 1, 53.08 (20.07) at week 12 and 43.06 (23.90) at week 24. For the specific EORTC constipation scale, the score was at least maintained and suggested an improvement over the OLE with mean (SD) scores in the total OLE population of 45.31 (35.18) at day 1, 38.49 (35.66) at week 12 and 42.71 (34.44) at week 24.

Safety

Overall, 93.8% of patients experienced at least one AE in the OLE phase of which 28.1% were classed as having AEs related to study treatment according to investigators. Fifty-nine patients (46.1%) experienced serious adverse events (SAEs). A total of four SAEs in four patients were related to study treatment: constipation, dyspnoea, paralytic ileus and worsening of cancer-related pain. Two additional SAEs in one patient were assessed as treatment-related by the sponsor. A

### Table 2: Patient assessment of constipation using PAC-SYM (OLE safety population)

|                      | Treatment during the DB phase | All OLE patients (N=128) |
|----------------------|-------------------------------|--------------------------|
|                      | OxyPR (n=62)                  | OXN PR (n=66)            |
| **Total symptom score** |                               |                          |
| Day 1                |                               |                          |
| n                    | 62                            | 66                       | 128                      |
| Mean (SD)            | 15.53 (10.20)                 | 10.55 (8.60)             | 12.96 (9.70)             |
| Week 12              |                               |                          |
| n                    | 41                            | 43                       | 84                       |
| Mean (SD)            | 13.22 (11.83)                 | 9.67 (7.43)              | 11.40 (9.93)             |
| Week 24              |                               |                          |
| n                    | 52                            | 44                       | 96                       |
| Mean (SD)            | 12.81 (10.10)                 | 11.34 (9.14)             | 12.14 (9.65)             |
| **Frequency of symptoms** |                               |                          |
| Day 1                |                               |                          |
| n                    | 61                            | 66                       | 127                      |
| Mean (SD)            | 1.93 (1.31)                   | 1.44 (1.07)              | 1.68 (1.21)              |
| Week 12              |                               |                          |
| n                    | 41                            | 43                       | 84                       |
| Mean (SD)            | 1.63 (1.39)                   | 1.30 (1.06)              | 1.46 (1.24)              |
| Week 24              |                               |                          |
| n                    | 51                            | 43                       | 94                       |
| Mean (SD)            | 1.67 (1.34)                   | 1.77 (1.27)              | 1.71 (1.30)              |
total of 38 patients discontinued during the OLE. In total, 28 patients (21.9 %) died during the study. Twenty-four of these deaths were due to disease progression or events related to disease progression; the other deaths were due to sepsis, worsening dyspnoea, lung embolism and cardiac arrest. The death from worsening of dyspnoea was recorded as unlikely to be related to study medication. All other deaths were recorded as unrelated to study medication. There were no clinically important changes in vital signs, laboratory values and electrocardiographs. Mean (SD) SOWS scores in the OLE safety population were comparable with those collected in the respective double-blind phase and decreased from 7.90 (7.39) at day 1 to 7.36 (6.24) at week 1 of the OLE. SOWS scores were comparable in both patient groups of the double-blind phase (Table 3).

Discussion

Pain is a common symptom of cancer, and if poorly treated, can adversely affect patients’ physical functioning, psychological well-being and social interactions [4]. Whilst opioids are widely recommended to relieve pain in cancer patients [6, 37], successful management requires that the benefits of these agents outweigh the impact of treatment-related side effects [38]. Opioid-induced bowel dysfunction, particularly OIC, is the most frequently reported AE experienced by patients receiving long-term opioid therapy and can be sufficiently severe to undermine the effectiveness of pain management [13].

To our knowledge, this OLE study is the largest trial to date evaluating OIC in cancer pain and provides a sound basis for evaluation of long-term safety and efficacy data. Results from this study demonstrate that long-term treatment (of at least 6 months) with OXN PR provides sustained analgesic efficacy for patients with moderate-to-severe cancer pain whilst maintaining improved bowel function. Increases in doses were small and there were no withdrawals due to lack of therapeutic effect indicating that pain control was maintained for the duration of the study for most subjects. The average pain scores based on the modified BPI-SF were low and stable over the 24-week OLE, indicating good analgesic efficacy during long-term treatment with OXN PR. The use of rescue medication was low and did not lead to any concern over the switching from OxyPR to OXN PR. The improvement in bowel function demonstrated with OXN PR during the double-blind phase of the study was maintained during the 24-week OLE phase, and patients who had received OxyPR during the double-blind period experienced improvement in bowel function following switch to OXN PR. Health status and QoL appeared to be maintained over the course of the OLE, with some possible improvements during the earlier part of the study.

It should be noted that data for QoL at week 24 (visit 13) included patients who had discontinued earlier in the study. Week 12 (visit 12), however, included only those data recorded at the visit. This may have contributed to any observed differences, particularly since the majority of discontinuations were due to disease progression/worsening of the condition and associated AEs.

The safety profile was as expected for this patient population, in accordance with the safety profile of opioid analgesics in this advanced cancer patient population. Although the overall AE and discontinuation rates were high, this was not unexpected considering the patient population and the duration of the study with respect to the course of cancer. The

### Table 3

| Event Description                                      | Number of Patients with Events, n (%) |
|--------------------------------------------------------|--------------------------------------|
| Total AE                                               | 120 (93.8)                           |
| AEs related to study medication                         | 36 (28.1)                            |
| Discontinuations due to AE                             | 38 (29.7)                            |
| SAE                                                    | 59 (46.1)                            |
| SAEs related to study medication                        | 4 (3.1)                              |
| Blood and lymphatic system disorders                    | 17 (13.3)                            |
| Anaemia                                                | 8 (6.3)                              |
| Gastrointestinal disorders                              | 44 (34.4)                            |
| Abdominal pain                                          | 7 (5.5)                              |
| Constipation                                           | 13 (10.2)                            |
| Diarrhoea                                              | 7 (5.5)                              |
| Nausea                                                 | 18 (14.1)                            |
| Vomiting                                               | 8 (6.3)                              |
| General disorders and administration site conditions    | 46 (35.9)                            |
| Asthenia                                                | 8 (6.3)                              |
| Peripheral oedema                                       | 17 (13.3)                            |
| Pain                                                   | 11 (8.6)                             |
| Pyrexia                                                | 7 (5.5)                              |
| Investigations                                          | 49 (38.3)                            |
| Increased blood uric acid                              | 7 (5.5)                              |
| Increased gamma-glutamyl transferase                   | 7 (5.5)                              |
| Decrease haemoglobin                                   | 15 (11.7)                            |
| Decreased lymphocyte count                             | 10 (7.8)                             |
| Metabolism and nutrition disorders                      | 26 (20.3)                            |
| Anorexia                                               | 11 (8.6)                             |
| Neoplasms benign, malignant and unspecified (including cysts and polyps) | 51 (39.8) |
| Cancer pain                                            | 22 (17.2)                            |
| Malignant neoplasm progression                         | 32 (25.0)                            |

AE adverse event, OLE open-label extension, SAE serious adverse event

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incidence of AEs in the OLE phase was comparable with that in the double-blind phase. The most frequently reported AEs were related to progression of underlying cancer disease, which is reflected in the profile of the frequently reported AEs of malignant neoplasm progression, cancer pain and nausea. The rate of treatment-related AEs (28.1%) in this cancer study was in line with reported AEs in non-malignant pain populations in OLE studies of OXN PR [28]. Although 59 patients (46.1%) experienced SAEs, the proportion considered related to study treatment was low (only 4 patients; 3.1%).

As with any clinical trial, the results of this study should be interpreted with some consideration of its limitations. Open-label extension studies are often uncontrolled and potentially biased due to patient self-selection, and further, randomized trials would better support the long-term use of OXN PR in chronic cancer pain. However, open-label studies, allow for greater patient exposure to medications, which enhances understanding of the efficacy and safety profile of the agent being investigated over a longer time period. As such, these results add to the accumulating evidence for the analgesic efficacy and improvement in bowel function and QoL associated with OXN PR, and suggest that these effects can be maintained during long-term therapy. Overall, the results reported here provide further support for the findings from the 4-week double-blind phase of this trial [29] and are also in line with those seen in patients with non-malignant pain [25–28].

Conclusion

The findings presented here demonstrated that the benefits of OXN PR seen in the double-blind study were maintained in the 24-week OLE. For patients with moderate-to-severe cancer pain, this allows sustained analgesia whilst maintaining improved bowel function and reduced symptoms of constipation. OXN PR was not associated with any unexpected safety or tolerability issues. These findings may reassure clinicians that patients may safely remain on stable drugs for pain as their cancer progresses.

Acknowledgments The authors would like to thank Medicus International for providing editorial assistance.

Funding This study, and the editorial assistance, was funded by Mundipharma Research GmbH & Co. KG.

Disclosures SA and WL have worked as consultants for Mundipharma Research; ML, HD and MH are employees of Mundipharma Research GmbH.

Conflict of interest MJ and HP have no conflict of interest to declare.

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