Emerging therapies for patients with symptoms of opioid-induced bowel dysfunction

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Abstract: Opioid-induced bowel dysfunction (OIBD) comprises gastrointestinal (GI) symptoms, including dry mouth, nausea, vomiting, gastric stasis, bloating, abdominal pain, and opioid-induced constipation, which significantly impair patients’ quality of life and may lead to undertreatment of pain. Traditional laxatives are often prescribed for OIBD symptoms, although they display limited efficacy and exert adverse effects. Other strategies include prokinetics and change of opioids or their administration route. However, these approaches do not address underlying causes of OIBD associated with opioid effects on mostly peripheral opioid receptors located in the GI tract. Targeted management of OIBD comprises purely peripherally acting opioid receptor antagonists and a combination of opioid receptor agonist and antagonist. Methylnaltrexone induces laxation in 50%–60% of patients with advanced diseases and OIBD who do not respond to traditional oral laxatives without inducing opioid withdrawal symptoms with similar response (45%–50%) after an oral administration of naloxegol. A combination of prolonged-release oxycodone with prolonged-release naloxone (OXN) in one tablet (a ratio of 2:1) provides analgesia with limited negative effect on the bowel function, as oxycodone displays high oral bioavailability and naloxone demonstrates local antagonist effect on opioid receptors in the GI tract and is totally inactivated in the liver. OXN in daily doses of up to 80 mg/40 mg provides equally effective analgesia with improved bowel function compared to oxycodone administered alone in patients with chronic non-malignant and cancer-related pain. OIBD is a common complication of long-term opioid therapy and may lead to quality of life deterioration and undertreatment of pain. Thus, a complex assessment and management that addresses underlying causes and patomechanisms of OIBD is recommended. Newer strategies comprise methylnaltrexone or OXN administration in the management of OIBD, and OXN may be also considered as a preventive measure of OIBD development in patients who require opioid administration.

Keywords: methylnaltrexone, naloxegol, opioid-induced constipation, oxycodone/naloxone, quality of life, opioid-induced bowel dysfunction

Introduction

Pain is a worldwide problem, and all efforts should be made to allow its effective management in each suffering patient. It is of paramount importance to assess pain precisely in its physical but also psychological, social, and spiritual dimensions, especially in patients suffering from chronic pain syndromes. Chronic pain management rules are based on the analgesic ladder established in 1986 by the World Health Organization (WHO). In most patients, pain is successfully relieved through the use of pharmacotherapy including opioids alone, or in combination with adjuvant analgesics in accordance with the WHO analgesic ladder. Pain management guidelines for cancer patients have been recently updated by the EAPC (European Association for Palliative Care) and ESMO (European Society for Medical Oncology).
Morphine, along with oxycodone and hydromorphone administered orally, are recommended as the first choice opioids at the third step of the WHO analgesic ladder, which also comprises additional opioids (transdermal formulations of fentanyl and buprenorphine, methadone and tapentadol) for the treatment of cancer patients with moderate-to-severe pain intensity. Currently, instead of weak opioids (opioids for mild-to-moderate pain), it is possible to use low doses of strong opioids (opioids for moderate-to-severe pain): morphine up to 30 mg, oxycodone up to 20 mg, and hydromorphone up to 4 mg per day, administered by the oral route on the second step of the WHO analgesic ladder.\textsuperscript{10}

Opioids are often successfully used for pain management, but they may also induce numerous and potentially serious adverse effects (AE). Although tolerance develops only for some opioid AE, such as sedation, there may be little or no tolerance development to opioid-induced gastrointestinal (GI) AE. Therefore, patients should be closely monitored by the staff to avoid or decrease the intensity of opioid-induced AE that may significantly affect patients’ quality of life (QoL) and lead to non-compliance with opioid regimens resulting in undertreatment of chronic pain.\textsuperscript{11} One common opioid adverse effect is a group of symptoms associated with the GI tract, the so-called opioid-induced bowel dysfunction (OIBD).\textsuperscript{12}

**Epidemiology of OIBD**

OIBD is a frequent phenomenon. Among 40 cancer patients with different primary tumor locations admitted to a palliative medicine inpatient unit, 84% suffered from dry mouth, 71% complained of early satiety, 58% reported constipation, 56% anorexia, 50% bloating, 48% nausea, 42% abdominal pain, and 34% vomiting.\textsuperscript{13} In an Internet study conducted in Europe and in the US, of 322 patients taking oral opioids for chronic cancer-related and noncancer pain, and laxatives, 45% reported less than three bowel movements (BM) per week. Constipation was reported by 81% of patients and straining during defecation by 58% of the patients surveyed. The most bothersome symptoms were, in order of rank: constipation, straining, fatigue, small and hard stools, and insomnia. A third of patients had missed, decreased, or stopped using opioids in order to make it easier to have a BM.\textsuperscript{14} In a study conducted in Spain among 317 outpatients taking different opioids for chronic cancer-related and non-cancer pain, 94.6% of patients reported at least one symptom of OIBD, with constipation reported most frequently (91.6%). Nearly half of the patients reported three or more symptoms with severity equal or above 4 on the numerical rating scale (11-point scale from 0 to 10). No significant differences were found in the prevalence of symptoms between patients treated with different opioids (morphine, oxycodone, fentanyl, or buprenorphine). A decrease related to moderate-to-severe GI symptom severity in the well-being of patients was detected.\textsuperscript{15}

In patients with non-malignant pain taking a median morphine dose of 127.5 mg, the most common symptoms comprised chronic abdominal pain (58.2% of patients), constipation (46.9%), gastro-esophageal reflux (33%), nausea (27%), and vomiting (10%).\textsuperscript{16} In another multicenter, observational study, among 2,324 patients surveyed, morphine (89.6%) and transdermal fentanyl (74.1%) more often induced OIBD than oxycodone and transdermal buprenorphine (59.3% each). Age over 70 years, cancer-related pain, and transdermal fentanyl were the risk factors for the development of OIBD symptoms.\textsuperscript{17} In another study conducted among 4,040 patients, similar constipation intensity was found in those treated with controlled-release (CR) morphine, CR oxycodone, and transdermal fentanyl with no difference between cancer and noncancer patients.\textsuperscript{18} In France, among 520 cancer patients with pain treated with opioids, 61.7% showed the degree of constipation that is problematic for the patient; 85.7% of patients were considered constipated according to physicians’ assessment despite laxatives used by 84.7% of patients. QoL was significantly reduced in constipated versus (vs) non-constipated patients ($P<0.0001$). Hospitalization, pain, and changes in laxative and opioid treatment are all consequences of OIBD development.\textsuperscript{19}

**Pathophysiology of OIBD**

OIBD symptoms reflect a complex impact of opioids on the GI tract (Table 1). The most typical and usually most burdensome symptoms of OIBD are called opioid-induced constipation (OIC). However, OIBD reflects the impact of opioids on the whole GI tract, which comprises such symptoms as: dry mouth; gastro-esophageal reflux-related symptoms (heartburn); nausea; vomiting; chronic abdominal pain; bloating; constipation-related symptoms, such as straining, hard stools, painful, infrequent, and incomplete BM; and diarrhea-related symptoms, such as urgency, loose BM, and frequent BM.\textsuperscript{12}

**Opioid impact on gut motility**

The pathomechanism of OIBD is associated mainly with the peripheral opioid effect on $\mu$-opioid receptors in the gut wall, with probably less central effects.\textsuperscript{20} The $\mu$-opioid receptors are present in neurons of myenteric and submucosal plexus and immune cells in the lamina propria.\textsuperscript{21} Opioid receptors (predominantly $\mu$, but also $\kappa$ and $\delta$) are located in the
myenteric plexus and in the submucosal plexus of the gut wall. The myenteric plexus lies between the longitudinal and circular layers and extends the length of the intestine, and it is mainly responsible for the control of motor activity of the gut. The submucosal plexus controls secretion and absorption; µ-opioid receptors are activated in the wall of the stomach, and small and large intestine by endogenous (enkephalins, endorphins, and dynorphins) and exogenous (eg, morphine, oxycodone, and methadone) opioids, and modify GI function. Activation of µ-opioid receptors inhibits excitatory and inhibitory neural pathways within the enteric nervous system, which coordinates motility. Inhibition of excitatory neural pathways decreases peristaltic contractions. In turn, the blockade of inhibitory neural pathways increases GI muscle activity and elevates resting muscle tone, spasm, and non-propulsive motility patterns. These mechanisms induce delayed gastric emptying and slow intestinal transit.22

Several neurotransmitters are involved in the regulation process of gut motility. Acetylcholine release activates the cholinergic excitatory motoneurons in the longitudinal smooth muscles. Nitric oxide and vasoactive intestinal peptide control the inhibition of the non-cholinergic inhibitory neurons in the circular smooth muscles. The coordination of the contractile and propulsive gut motility is determined by the balance between acetylcholine and nitrous oxide plus vasoactive intestinal peptide release. Opioids inhibit the release of these neurotransmitters and disrupt the coordination of gut motility.12

In the esophagus, opioids induce non-peristaltic contractions with an incomplete relaxation of the lower esophageal sphincter. This may manifest as dysphagia. In the small and large intestine, opioids increase resting contractile tone in the circular muscle and decrease tonic inhibition of the muscle tone, which together leads to an increased tone in the circular muscle layer. Increased tone is accompanied by enhanced rhythmic contractions and occasional, high-amplitude, non-propulsive phasic contractions and associated changes in smooth muscle electrical activity. All these abnormalities result in an increased segmental contraction along with decreased propulsive forward peristalsis. Clinically, these changes manifest as constipation, gut spasm, and abdominal cramps. A consequence of this peristaltic disruption is stasis of luminal contents, which leads to an increased passive absorption of fluids, dryer and harder stools.12

### Opioid impact on gut secretion

Activation of opioid receptors in the submucosa inhibits water and electrolyte secretion into the gut lumen, and increases fluid absorption from the intestine and blood flow in the gut wall.23 Opioids increase activity in the sympathetic nervous system and decrease the secretion. Endocrine cells located in the epithelium play a role in regulating motor activity and secretion in the gut. Studies conducted in mice indicate that peripheral µ-opioid receptors inhibit the transit independently of central µ-receptors.24 Opioids increase ileocecal and anal sphincter tones and impair defecation reflex through reduced sensitivity to distension, and increased internal anal sphincter

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Table 1 Pharmacological mechanisms and clinical symptoms of opioid-induced bowel dysfunction

<table>
<thead>
<tr>
<th>Pharmacological mechanisms</th>
<th>Clinical symptoms</th>
</tr>
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<tbody>
<tr>
<td>Decreased saliva production</td>
<td>Xerostomia</td>
</tr>
<tr>
<td>Dystotility of the lower esophageal sphincter</td>
<td>Gastro-esophageal reflux (or, rarely, dysphagia)</td>
</tr>
<tr>
<td>Decreased gastric secretion, emptying and motility</td>
<td>Delayed absorption of medication, upper abdominal discomfort</td>
</tr>
<tr>
<td>Disturbed fluid secretion and absorption</td>
<td>Constipation</td>
</tr>
<tr>
<td>Abnormal bowel motility, increased resting contractile tone in the small and large intestines, and sphincter dysfunction</td>
<td>Straining, incomplete bowel evacuation, bloating, abdominal distension, constipation</td>
</tr>
<tr>
<td>Increased amplitudes of non-propulsive segmental bowel contractions</td>
<td>Spasm, abdominal cramps and pain, stasis of luminal contents, and hard dry stool</td>
</tr>
<tr>
<td>Constriction of sphincter of Oddi</td>
<td>Biliary colic, epigastric discomfort, and pain</td>
</tr>
<tr>
<td>Increased anal sphincter tone and impaired reflex relaxation during rectal distension</td>
<td>Evacuation disorders</td>
</tr>
<tr>
<td>Diminished intestinal, pancreatic, and biliary secretion</td>
<td>Hard, dry stools</td>
</tr>
<tr>
<td>Abnormal bowel motility, increased fermentation and meteorism, opioid-induced hyperalgesia</td>
<td>Chronic visceral pain</td>
</tr>
<tr>
<td>Central effects of opioids</td>
<td>Nausea and vomiting, anorexia</td>
</tr>
</tbody>
</table>

Notes: Adapted from: Springer International Publishing AG; Drugs; Opioid-induced bowel dysfunction: pathophysiology and management; 72(14); 2012; 1847–1865; Brock C, Olesen SS, Olesen AE, Frøkjaer JB, Andresen T, Drewes AM; Copyright © Springer International Publishing AG 2012; with kind permission of Springer Science+Business Media.11
tive. Morphine causes sphincter contraction and decreases emptying of pancreatic juice and bile, which may delay digestion. The anal sphincter dysfunction is an important factor in the sensation of anal blockage.

Peripheral and central effects of opioids on the gut
Central mechanisms of opioid effects on GI tract were demonstrated in an experimental study in which intracerebroventricular administration of morphine in rats, inhibited GI propulsion. This effect was reversed by intracerebroventricular administration of naloxone and vagotomy. Intrathecal administration of morphine reduced gastroduodenal motility, and morphine administered intramuscularly, yielded additional effects. Thus, both central and peripheral opioid action may play a role in opioid GI effects. The indirect evidence of both central and peripheral components of opioid effects on bowel function can be observed in the 50%–60% response rate to the treatment of OIBD with methylnaltrexone (MNTX), which displays purely peripheral μ-opioid receptor antagonist effects. However, in humans, it seems that the peripheral opioid effects dominate GI transit. OIBD is the consequence of reduced GI motility, increased absorption of fluids from the gut, and decreased epithelial secretion. The stool remains in the gut lumen for a longer time; therefore, more fluid is reabsorbed and the stool becomes hard and dry. These effects are also associated with opioids’ inhibition of secretomotor neurons in the epithelium of the gut.

Clinical assessment of patients with OIBD
The complex assessment of patients with symptoms of OIBD is necessary for the application of effective treatment. Several subjective scales were developed for that purpose, e.g., the Bowel Function Index (BFI). The BFI is a three-item questionnaire that assesses the severity of ease of defecation, feeling of incomplete bowel evacuation, and patients’ personal judgment of constipation using a 0 to 100 numerical rating (0, no symptoms, and 100, the most severe symptoms) (Table 2). Patients rate these variables according to their experience of the preceding 7 days. The BFI score is calculated as the mean of the three component scores. Reference results of non-constipated chronic pain patients’ BFI scores lie within the range of 0–28.8. The BFI was validated in clinical studies conducted in cancer and noncancer patients. Changes in BFI scores equal or above 12 are likely to be clinically meaningful, and those lower than 7.5 unlikely to be clinically relevant, whilst changes between 7.5 and 12 are uncertain and not clear, and require further evaluation.

The Patient Assessment of Constipation comprises two instruments: the symptom questionnaire (PAC–SYM), and the quality of life questionnaire (PAC–QOL). The PAC–SYM contains 12 items assigned to three subscales: stool symptoms, rectal symptoms, and abdominal symptoms (Table 3). In addition, the PAC–SYM also contains a question about the frequency of BM during the last 7 days. The PAC–QOL is composed of 28 items grouped into four subscales related to dissatisfaction (five items), physical discomfort (four items), psychosocial discomfort (eight items), and worries and concerns (eleven items) (Table 4). A

### Table 2 The Bowel Function Index (BFI)

<table>
<thead>
<tr>
<th>Item</th>
<th>Intensity</th>
</tr>
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<tbody>
<tr>
<td>Ease of defecation during the last 7 days according to patient assessment</td>
<td>0 1 2 3 4 5 6 7 8 9 10</td>
</tr>
<tr>
<td>Feeling of incomplete bowel evacuation during the last 7 days according to patient assessment</td>
<td>0 1 2 3 4 5 6 7 8 9 10</td>
</tr>
<tr>
<td>Personal judgment of patient regarding constipation during the last 7 days</td>
<td>0 1 2 3 4 5 6 7 8 9 10</td>
</tr>
</tbody>
</table>

**Note:** Adapted from: Validation of the Bowel Function Index to detect clinically meaningful changes in opioid-induced constipation; Rentz AM, Yu Y, Müller-Lissner S, Leyendecker P; Journal of Medical Economics; 2009; Copyright © 2009 Informa Healthcare; adapted by permission of the publisher Informa Healthcare.
validation study has confirmed that PAC–QOL is internally consistent, reproducible, valid, and responsive to improvements over time. The objective assessment comprises the Bristol Stool Chart—a simple and easy-to-use method of objective assessment detecting stool frequency and stool consistency. However, only a moderate correlation between stool form and whole-gut or colonic transit time can be demonstrated.

Clinical evaluation is composed of a detailed history that should take into account GI symptoms with comprehensive assessment of other physical problems along with psychological and spiritual concerns. Careful physical examination includes palpation of abdomen, auscultation of peristalsis, and rectal examination. Specifically, acute abdominal diseases should be excluded such as bowel obstruction or gut perforation. Investigations comprise plain radiography, transit time studies, manometry, and anorectal dysfunction tests. However, apart from plain radiography, these investigations are rarely used in patients with advanced diseases.

Management of OIBD
Several drug groups may be used for the management of OIBD. However, most of these do not address the underlying cause of this condition. The currently available interventions for patients with OIBD symptoms are presented in Table 5.

Non-targeted treatment of OIBD
Prokinetic agents
Metoclopramide is commonly prescribed for patients with symptoms of gastroparesis. It works through both peripheral (in the upper GI tract) and central mechanisms by inducing antidiopaminergic effects. The typical dose of metoclopramide is usually 10 mg three times a day. The most common AE comprise restlessness, drowsiness, and fatigue. Metoclopramide may induce extra-pyramidal effects, especially in younger patients and in children, and therefore should be avoided in these patients. Nonetheless, it is recommended to be used for a short period of time, ie, a few days in other patients. Metoclopramide is an inhibitor
of CYP2D6 enzyme. Concurrent use of antidepressants such as tricyclics, selective serotonin reuptake inhibitors, and antidepressants acting as serotonin–noradrenaline reuptake inhibitors (venlafaxine, or duloxetine) may enhance AE of metoclopramide. Domperidone may be an alternative drug, which in contrast to metoclopramide, does not cross the blood–brain barrier and displays less risk of extra-pyramidal effects. Cisapride, a 5-HT₃ receptor agonist, works in the whole GI tract, but it is not recommended due to the high risk of cardiotoxicity.

Itopride is a peripheral antidopaminergic agent that also increases acetylcholine levels due to the inhibition of acetylcholinesterase, and the prokinetic effect observed in the whole GI tract. It does not activate 5-HT₄ or 5-HT³ receptors. Itopride is metabolized through a mono-oxidase system, and there is little risk of significant pharmacokinetic interactions with other medications. Itopride does not permeate the blood–brain barrier and poses no risk of extra-pyramidal effects. The recommended dose is normally 50 mg three times a day.

Newer prokinetics, such as prucalopride, lubiprostone, and linaclotide have recently been introduced. Prucalopride is a selective 5-HT₃ receptor agonist which stimulates propulsive gut motility in vitro and in vivo. Prucalopride in the daily doses of 2–4 mg accelerates the whole gut, gastric, small bowel, and colonic transit in patients with chronic constipation. The recommended dose is usually 2 mg once daily. However, prucalopride was introduced for the treatment of chronic constipation, predominantly in women. The most common AE comprise headache (25%–30% of treated patients), nausea (12%–24%), abdominal cramps (16%–23%), and diarrhea (12%–19%). A phase-2 controlled study conducted during 4 weeks in 196 patients with noncancer pain and OIC found an increase in the number of complete spontaneous BM in patients treated with prucalopride at a daily dose of 2–4 mg compared to placebo, but the difference did not reach statistical significance. Both itopride and prucalopride do not interfere with cardiac functioning.

Lubiprostone activates type-2 chloride channels and increases intestinal fluid secretion, thereby facilitating...
intestinal transit and increasing the passage of stool. It was demonstrated to be effective in patients with OIC suffering from non-malignant pain. Lubiprostone is registered by the US Food and Drug Administration (FDA) for the management of OIBD in patients with chronic non-malignant pain based on the results of three phase-3 trials (NCT00595946, NCT01298219, and NCT00597428) that demonstrated superiority of the drug over placebo. The most common AE comprised nausea (11% of treated patients), diarrhea (8%), and abdominal pain (4%). However, in contrast to morphine, lubiprostone may be ineffective in patients treated with methadone.\(^5\)

Linaclotide is an agent that regulates chloride secretion on the intestinal epithelial cells through activation of guanylate cyclase C. Linaclotide stimulates the GI secretion and transit, and also modulates visceral sensitivity. It increases stool frequency, stool weight, and improves the ease of stool passage. The main AE of linaclotide is diarrhea.\(^6\) Linaclotide may be potentially considered for the management of OIBD, although this drug was only used in patients with chronic constipation and irritable bowel syndrome.\(^7\)

### Laxatives

Numerous laxatives are prescribed for the prevention and treatment of OIBD, particularly for constipation-related symptoms. The most commonly used group of laxatives for OIBD comprise osmotic agents and stimulants. Nonetheless, they possess limited efficacy and display their own AE.\(^8\) Moreover, treatment with laxatives should be limited to a short period of time. Patients with chronic non-malignant and cancer-related pain usually have to take laxatives for the long term, as tolerance to the constipating effect of opioids develops to little extent or not at all. OIBD is often unsuccessfully managed due to not only the lack of efficacy and AE of traditional laxatives, but also due to the lack of appropriate clinical evaluation and knowledge on OIBD management and expertise of the medical staff.\(^9\)

### Change of opioid route of administration and opioid rotation (switch)

Another approach is a change from oral to parenteral or to transdermal route of opioid administration. However, all opioids display OIBD including transdermal formulations.\(^10\) From the point of view of physicochemical properties, it may be beneficial to switch from hydrophilic opioids such as morphine, oxycodone, or hydromorphone to lipophilic opioids such as fentanyl, buprenorphine, or methadone due to more involvement of GI receptors with hydrophilic formulations and less with lipophilic formulations.\(^11\) However, only few studies demonstrated improvement in constipation, predominantly after the switch from oral to transdermal opioids.\(^12,13\)

### Targeted treatment of OIBD

Newer strategies for the management of OIBD include administration of purely peripherally acting opioid receptor antagonists, predominantly MNTX and a combination of prolonged-release (PR) oxycodone with PR naloxone (OXN) in one tablet. MNTX is administered in the form of subcutaneous injections when treatment of OIBD with traditional oral laxatives has failed to provide effective bowel movements.\(^14\) OXN renders analgesia with limited negative effect of oxycodone on bowel function.\(^15\)

#### Combining PR oxycodone with PR naloxone

**Pharmacodynamics and pharmacokinetics**

Oxycodone is an opioid agonist that predominantly binds to \(\mu\) and \(\kappa\) opioid receptors.\(^15\) Naloxone is an opioid antagonist that acts at \(\mu, \kappa,\) and \(\delta\) opioid receptors. Naloxone displaces opioid agonists from the receptors due to higher affinity, and when administered by the oral route it improves bowel function in patients with OIBD.\(^16\) The chemical structure of oxycodone and naloxone is shown on Figure 1. OXN tablets contain PR oxycodone and PR naloxone in different strengths: 5 mg/2.5 mg, 10 mg/5 mg, 20 mg/10 mg, and 40 mg/20 mg. The 2:1 ratio of oxycodone/naloxone was deemed optimal in a phase-2 study conducted in patients with severe chronic pain as effective analgesia and improvement in bowel function;\(^17\) the observed AE were acceptable.\(^18\) OXN is registered for patients with severe pain that may only be successfully treated with opioid analgesics.\(^19,20\)

OXN analgesia provided by oxycodone is not reversed by naloxone; opioid withdrawal is not observed (Figure 2). An important difference between the administration of immediate-release (IR) and PR formulations of naloxone is that IR naloxone may attenuate analgesia or induce opioid

![](https://www.dovepress.com/)

**Figure 1** Chemical structures of oxycodone and naloxone.
withdrawal. PR naloxone prevents the saturation of hepatic enzymes responsible for naloxone metabolism, and reduces the risk of passing naloxone on to systemic circulation and opioid antagonism in the central nervous system. OXN is contraindicated in patients with bowel obstruction, acute abdominal conditions, diarrhea, allergy to the drug, and hepatic failure. OXN should not be administered before and 12–24 hours after the surgery. As no data exist for the administration of OXN in children and patients under 18 years of age, the drug should not be administered in these patient populations.

Oxycodone displays high oral bioavailability (60%–87%) and is metabolized primarily in the liver and in the intestine wall, predominantly through CYP3A to noroxycodone, and to less extent through CYP2D6 to oxymorphone. As oxycodone is metabolized through CYP3A and CYP2D6, drugs inhibiting these enzymes, especially CYP3A, should be avoided due to the risk of an increase in oxycodone plasma levels that may lead to toxicity. The highest risk is associated with concurrent administration of CYP2D6 and CYP3A4 inhibitors. The risk of drug interactions in the case of naloxone is lower due to the type of metabolism (glucuronidation) without P-450 system involvement.

Oxycodone and its metabolites are excreted with urine and with feces. The volume of distribution of oxycodone equals 2–3 L/kg. The maximum plasma concentration is reached within 25 minutes after intravenous injection, 1.3 hours after IR administration, and 2.6 hours after the administration of a CR formulation. The $T_{1/2}$ (half-life) is approximately 2–3 hours after intravenous administration, 3 hours after administration with an IR, and approximately 5 hours after treatment with PR oxycodone tablets. Oxycodone predominantly binds to albumin (45%).

Naloxone exhibits low oral bioavailability (<3%) and extensive first-pass metabolism in the liver, with the formation of naloxone-3-glucuronide (NAL-3-G). Another naloxone metabolic pathway is N-dealkylation and the reduction of the 6-keto group to 6-alpha-naloxol and 6-beta-naloxol, although only the latter has been detected in humans. NAL-3-G is measured in plasma instead of the parent compound, which is undetectable after oral administration. Naloxone and its metabolites are excreted with urine. The effect of orally administered naloxone depends on normal liver function; thus, any hepatic impairment should be carefully considered. In patients suffering from liver failure, PR oxycodone/PR naloxone administration is not recommended.

Highly fatty meals may insignificantly increase oxycodone bioavailability. Food consumption does not affect NAL-3-G levels, and OXN may be administered in fasting condition or with food. OXN was not studied in patients during pregnancy and delivery. However, oxycodone and naloxone penetrate to placenta, and their long-term administration in pregnant women may evoke opioid withdrawal syndrome in neonates. Oxycodone permeates to the milk; thus, OXN should not be administered in lactating women.

**Figure 2** Mechanisms of action of opioid agonist/antagonist combinations to counteract opioid-induced bowel dysfunction development. **Abbreviation:** CNS, central nervous system.
Dosing guidelines
The starting OXN doses in opioid-naïve patients are normally in the range of 5 mg/2.5 mg–10 mg/5 mg administered twice daily (every 12 hours). In patients not responding to opioids, for mild-to-moderate pain (tramadol, codeine, dihydrocodeine, or hydrocodone), the initial dose usually recommended equals 10 mg/5 mg or 20 mg/10 mg twice daily. The dose may be titrated to achieve satisfactory analgesia and tolerable AE. The recommended maximal daily dose of OXN is 40 mg/20 mg administered twice daily. However, higher daily doses of up to 120 mg/60 mg were safely administered in clinical studies.77 When OXN doses exceed the recommended dose range, a rotation to oxycodone administered alone, or other opioids for moderate-to-severe pain intensity should be instituted.

When rotating from other opioids for moderate-to-severe pain to OXN, the starting dose is established individually, depending on the amount of the opioid administered, analgesia, AE, and complex clinical evaluation. Patients treated with OXN should have access to IR formulations of opioids for breakthrough pain management. In the case of IR oxycodone, a single dose usually equals approximately a sixth of the daily oxycodone dose used in OXN. However, when using IR opioids for breakthrough pain management, especially rapid onset fentanyl formulations, the treatment should always be started with the lowest dose and then be titrated to achieve good analgesia and acceptable toxicity.78 OXN should be administered with caution in patients with renal impairment; in patients with renal failure, a switch to fentanyl or buprenorphine should be considered.8 As OXN is contained in PR formulations, the whole tablets should be swallowed without crushing, gnawing, or chewing.

Adverse effects
AE of OXN and PR oxycodone are similar. The frequency of diarrhea is higher in patients treated with OXN compared to those treated with PR oxycodone alone (5.2% vs 2.6%),89 with less frequent nausea (6.3% vs 10.5%), vomiting (1.3% vs 4.3%), abdominal pain (1.3% vs 4.3%) and dyspepsia (0.6% vs 2.5%), compared to PR oxycodone administered alone.80 These effects are associated with naloxone antagonist effect on opioid receptors in the GI tract,81 and prokinetic properties.82 In another study, most GI AE were more frequent in patients treated with OXN (23.8%) compared with PR oxycodone (16.3%): abdominal pain (7.7% vs 1.5%), constipation (0.8% vs 1.5%), diarrhea (4.6% vs 3.0%), nausea (10.0% vs 6.7%), and vomiting (3.1% vs 0.7%), respectively.83

Long-term treatment with OXN over a period of up to 52 weeks in patients with chronic pain who completed 12 weeks of treatment in two phase-3 studies,79,80 was safe.84 A total of 6.3% of treated patients discontinued therapy with OXN due to AE. The most commonly reported AE were constipation (9.2%), nausea (7.7%), back pain and depression (6.3% each), and diarrhea (3.2%). Withdrawal symptoms were experienced by 0.53% of all treated patients.84 Similar incidences of all AE from OXN (61.0%) and PR oxycodone (57.3%), GI AE (20.9% and 21.7%, respectively; diarrhea, 5.1% and 3.7%, respectively), and discontinuation rate due to AE (4.8% and 6.4%, respectively) were found in an analysis of two controlled studies83,85 conducted in patients with chronic non-malignant pain.85 No increase in the incidence of opioid withdrawal syndrome during treatment with OXN in daily doses up to 120 mg/60 mg compared to oxycodone was found.83 Long-term study of cancer (24 weeks) and non-cancer patients (1 year) confirmed good analgesic efficacy and safety without withdrawal symptoms during treatment with OXN.86,87

Mercadante et al88 in a case report depicted a cancer patient with severe pain who required high OXN daily doses (240 mg/120 mg), which turned out to be ineffective. A switch to PR oxycodone administered alone at a daily dose of 240 mg provided satisfactory analgesia. This case report suggests that at higher doses (240 mg/120 mg daily), OXN provides inferior analgesia compared to PR oxycodone administered alone at a respective daily dose (240 mg).88

Kang et al89 depicted a female patient diagnosed with gastric cancer with peritoneal seeding, extensive thrombosis in the portal vein, and abdominal pain. PR oxycodone at a dose of 10 mg administered twice daily provided satisfactory analgesia but caused OIC not amenable to laxative therapy. For this reason, the patient was rotated to OXN at a dose of 10 mg/5 mg administered twice daily. After taking OXN, the patient experienced severe pain and symptoms such as chills, diarrhea, general weakness, and sweating suggesting opioid withdrawal. The patient returned to the former treatment with PR oxycodone 10 mg administered twice daily, and her pain was again effectively relieved, and other symptoms disappeared. This result suggests development of a collateral circulation due to portal vein thrombosis, and naloxone directly reaching the central nervous system and evoking withdrawal symptoms.89 Numerous other studies have demonstrated clinical effectiveness of OXN in patients with chronic non-malignant pain,67,88,79,83,84,90–95 cancer-related pain,96–99 and postoperative pain.100,101
Purely peripherally acting opioid receptor antagonists

Treatment with opioid receptor antagonists that act purely peripherally may be used when traditional oral laxatives fail.\textsuperscript{102}

MNTX

Pharmacodynamics and pharmacokinetics

MNTX is a quaternary methyl derivative of naltrexone, preferential, non-selective, peripheral \(\mu\)-opioid receptor antagonist.\textsuperscript{102} The addition of the methyl group to the

Table 6 Mechanisms of action of purely peripherally acting \(\mu\)-opioid receptor antagonists (PAMORA)

<table>
<thead>
<tr>
<th>Place of action</th>
<th>Opioid agonists’ effects</th>
<th>Opioid antagonists’ effects</th>
</tr>
</thead>
<tbody>
<tr>
<td>Central nervous system</td>
<td>Analgesia</td>
<td>No effect on analgesia</td>
</tr>
<tr>
<td></td>
<td>Nausea</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Vomiting</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Sedation</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Decreased sensation of dyspnea</td>
<td></td>
</tr>
<tr>
<td>Blood–brain barrier</td>
<td>Decreased gastric secretion, emptying and motility</td>
<td>PAMORA eg, methylxaltrexone, naloexogol</td>
</tr>
<tr>
<td></td>
<td>Inhibition of propulsive intestinal contractions</td>
<td>Inhibition of opioid-induced gastrointestinal adverse effects</td>
</tr>
<tr>
<td></td>
<td>Diminished intestinal, pancreatic and biliary secretion</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Increased absorption of fluids and electrolytes</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Increased anal sphincter tone</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Constipation</td>
<td></td>
</tr>
</tbody>
</table>

Notes: Adapted from Springer International Publishing AG; Journal of Gastroenterology; Physiology, signaling, and pharmacology of opioid receptors and their ligands in the gastrointestinal tract: current concepts and future perspectives; 49(1): 2014; 24–45; Sobczak M, Salaga M, Storr MA, Fichna J. Copyright © The Authors 2013; with kind permission of Springer Science+Business Media.\textsuperscript{104}
nitrogen ring increases polarity and reduces lipophilicity of MNTX. As a consequence, MNTX does not cross the blood–brain barrier and displays limited absorption from the gut. The chemical structures of naloxone, MNTX, naloxegol, and alvimopan are shown in Figure 3. When administered systemically (orally, subcutaneously, or intravenously), MNTX does not reverse opioid analgesia, which is the case for MNTX intrathecal administration.103 The principal mechanisms of action of MNTX are shown in Table 6.104

MNTX binds to µ-opioid receptors and with less potency to κ-opioid receptors, and it does not bind to δ-opioid receptors. The inhibition of propulsive motor activity in the gut by opioids is primarily mediated by the activation of local µ-opioid receptors in the plexus myentericus of the GI tract. MNTX administration reverses the inhibiting effect of opioids on the gut during opioid therapy in patients and reduces fecal transit time in volunteers, the latter suggesting endogenous opioids’ role in the regulation of gut motility. MNTX induces bowel movement in approximately 50%–60% treated patients within 4 hours of its administration.33,34 MNTX is registered for the treatment of OIC in patients with advanced illness who require palliative care. MNTX is used in the treatment of OIC in advanced diseases in adult patients when conventional oral laxatives are ineffective.64

MNTX is suggested to have little oral bioavailability. However, a study conducted in healthy volunteers demonstrated the efficacy of oral MNTX in the prevention of the delay in oro-cecal transit time after intravenous morphine administration.105 The levels of MNTX in plasma after ingestion of enteric-coated oral formulation (to prevent the absorption of the drug in the stomach and to release MNTX in the intestine) were substantially lower in comparison to those observed after administration of uncoated formulations, which may suggest a more efficient local action on opioid receptors in the gut of enteric-coated MNTX oral formulations.106 MNTX is usually administered subcutaneously and is rapidly absorbed, with the peak plasma concentration attained within 30 minutes, and plasma $T_{1/2}$ is approximately 8 hours.107

After intravenous administration, a maximum plasma concentration is observed within 5 minutes, and plasma $T_{1/2}$ equals approximately 2.5–2.9 hours. Approximately 11%–15% of MNTX binds to proteins. MNTX is a weak CYP2D6 inhibitor with no significant drug interactions.108 MNTX undergoes two major metabolic pathways in humans: 1) sulfation of the phenolic group to MNTX-3-sulphate (M2), and 2) reduction of the carbonyl group to two epimeric alcohols, methyl-6α-naltrexol (M4) and methyl-6β-naltrexol (M5). Neither naltrexone, nor its metabolite 6β-naltrexol, was detected in human plasma after administration of MNTX, which confirmed that N-demethylation was not a metabolic pathway in humans.109 Approximately 10% of MNTX is metabolized through the glucuronidation process in the liver. As regards elimination, approximately 40%–50% is excreted mostly unchanged in urine, and approximately 20% through feces, with a total 70% recovery of the administered MNTX dose.110

Systematic reviews found MNTX to be effective in palliative care patients with OIC when conventional laxatives have failed.111,112 However, the safety of the product could not be clearly established; thus, large controlled trials are required.113 A more recent systematic review and meta-analysis of 14 controlled studies including 4,101 patients demonstrated superiority of mu-opioid receptor antagonists over placebo in the treatment of OIC. MNTX, naloxone, and alvimopan all were superior to placebo. AE (diarrhea and abdominal pain) were more common during these drugs’ administration compared to placebo. Reversal of analgesia was not observed during active therapy. Lubiprostone was beneficial in two controlled studies, and prucalopride showed a tendency to elicit a better response in one study.114 However, MNTX failed to improve postoperative nausea and vomiting in women undergoing gynecological surgery.115

**Dosing recommendations**

The drug MNTX is available in ampules containing 12 mg MNTX bromide in the volume of 0.6 mL, and is applied via subcutaneous injections. The single MNTX dose equals 8 mg in patients with body weight of 38–61 kg, or 12 mg if the body mass is 62–114 kg.116 Patients falling outside of this range should receive a dose of 0.15 mg/kg. A bowel movement within 4 hours after MNTX injection is observed in 50%–60% of patients (the median time to bowel movement after the drug administration is 30 minutes). If no therapeutic effect is observed, the injection may be repeated every other day. The efficacy of MNTX in patients who have already responded to the treatment reaches 57%–100% (2–7 doses); when the first, or the first two doses are ineffective, 35% patients respond to the second and 26% to the third dose, respectively.117 The European Medicines Agency (EMA) has approved subcutaneous administration of MNTX for the treatment of OIC in adults with advanced illness.118 The use of MNTX may eliminate the need for unpleasant rectal measures, which are poorly tolerated by patients.119
MNTX dosing is similar in younger and older patients. No dose adjustment is required in patients with mild-to-moderate renal impairment. However, in patients with mild renal failure (creatinine clearance <30 mL/min), the MNTX dose should be reduced by half. No dose adjustment is required for patients with mild-to-moderate hepatic impairment. MNTX is not recommended for patients with end-stage renal impairment requiring dialysis, or for patients with severe liver impairment.

**Adverse Effects**

MNTX may be used in palliative care patients with OIC not amenable to treatment with traditional oral laxatives, as demonstrated in clinical studies. However, long-term efficacy and safety of MNTX has not been clearly established. As MNTX does not cross the blood–brain barrier, the attenuation of analgesia or opioid withdrawal symptoms are not observed. The use of MNTX is contraindicated in patients with bowel obstruction, in acute abdominal conditions, diarrhea, and in the case of allergy to the drug.

MNTX may induce several AE: abdominal pain (28% of treated patients), flatulence (13%), nausea (11%), dizziness (7%), and diarrhea (5%), which usually have mild-to-moderate intensity and are associated with the defecation act. In an extension phase of this study, the most common AE comprised abdominal pain, typically of mild-to-moderate intensity (30.5% of treated patients), malignant tumor progression (24.4%), nausea (20.7%), and vomiting (19.5%). The most frequent AE observed in the analysis of the controlled study were as follows: abdominal pain (19.3%, 15.5%, and 3.7%); diarrhea (6.0%, 11.5%, and 3.7%); and nausea (8.7%, 11.5%, and 6.2%) in patients receiving MNTX once daily, every other day, and placebo, respectively. MNTX may exacerbate vomiting induced by morphine after a single 8 mg dose administration.

In a few adult patients with OIC, advanced illness, and conditions that may induce local or diffuse reduction of structural integrity of the GI tract wall (eg, cancer, peptic ulcer, and Ogilvie’s syndrome), MNTX caused GI perforation. GI perforations have been found in seven patients and involved varying regions of the GI tract, such as stomach, duodenum, and colon; in most cases, GI perforations appear after the first dose of MNTX. The concurrent medical conditions included two cases of duodenal peptic ulcer and single cases of amyotrophic lateral sclerosis, volvulus, bevacizumab use, metastatic colon cancer, and bowel obstruction. Interestingly, in a patient with amyotrophic lateral sclerosis diagnosis and constipation not responsive to traditional laxatives, MNTX was used without opioid administration.

It is recommended that MNTX should be used with caution in patients with present or suspected lesions of the GI tract. Treatment with MNTX should be stopped if severe abdominal symptoms appear or intensify. If severe or persistent diarrhea occurs during the treatment, patients are advised to discontinue the MNTX therapy. In a retrospective chart review conducted on 200 patients who were prescribed MNTX (202 physicians’ orders, 282 doses administered) in a comprehensive cancer center (inpatient and outpatient clinics), the prescription order was often not adherent to the label’s indication. In 18% of orders, MNTX was prescribed within 24 hours of the last bowel movement, 33% of orders were for patients with non-advanced illness, and 25% were for patients who had no OIC; 19% and 3% of orders were for patients who had no history of laxative and opioid use, respectively, and 5% of orders were for patients with documented bowel obstruction.

Clinical studies have demonstrated the effectiveness of MNTX in the treatment of OIC not responding to traditional laxatives in patients with advanced diseases, non-malignant pain, and in other patient populations. However, MNTX was ineffective in shortening the duration of postoperative ileus following segmental colectomy.

**Naloxegol**

Naloxegol (NKTR-118) is a polymer conjugate of the opioid antagonist naloxone administered once daily by the oral route. The polyethylene glycol moiety limits naloxegol capacity to cross the blood–brain barrier. In a phase-2, randomized, double-blind, placebo-controlled, dose-escalation study, the efficacy and safety of three dose levels of oral naloxegol in the treatment of OIC in patients with non-malignant or cancer-related pain was evaluated. The dose of 25 mg naloxegol was chosen to be further explored in phase-3 studies.

Two phase-3, double-blind clinical studies (KODIAC-04 [NCT01309841] and KODIAC-05 [NCT01323790]) have been completed, and demonstrated significant increase in spontaneous BM compared to placebo for the 25 mg naloxegol dose (in one study also for the 12.5 mg dose), with arthralgia as the only AE more common than in the placebo group. In both studies (KODIAC-04, 652 participants; KODIAC-05, 700 participants), outpatients with noncancer pain and OIC were randomly assigned to receive a daily dose of 12.5 or 25 mg of naloxegol or placebo, respectively. The primary endpoint was the 12-week response rate (≥3 spontaneous BM per week, and an increase from baseline of ≥1 spontaneous BM for ≥9 of 12 weeks and for ≥3 of the final 4 weeks) in the intention-to-treat population. The key secondary endpoints were the response rate in the subpopulation of patients with an inadequate response to laxatives before
enrollment, time to first post-dose spontaneous BM, and mean number of days per week with one or more spontaneous BM.

Response rates were significantly higher with 25 mg of naloxegol than with placebo (intention-to-treat population: KODIAC-04, 44.4% vs 29.4%, respectively \[P=0.001\]; KODIAC-05, 39.7% vs 29.3%, respectively \[P=0.02\]; patients with an inadequate response to laxatives: KODIAC-04, 48.7% vs 28.8%, respectively \[P=0.002\]; KODIAC-05, 46.8% vs 31.4%, respectively \[P=0.01\]). In KODIAC-04, response rates were also higher in the group treated with 12.5 mg of naloxegol than placebo (intention-to-treat population, 40.8% vs 29.4%, respectively \[P=0.02\]; in patients with an inadequate response to laxatives, rates for naloxegol vs placebo were 42.6% vs 28.8%, respectively \[P=0.03\]). A shorter time to the first post-dose spontaneous BM and a higher mean number of days per week with one or more spontaneous BM were observed with 25 mg of naloxegol vs placebo in both studies \(P<0.001\), and with 12.5 mg of naloxegol in KODIAC-04 \(P<0.001\). Pain scores and daily opioid dose were similar among the three groups. AE (primarily gastrointestinal) occurred most frequently in the groups treated with 25 mg of naloxegol. Treatment with naloxegol, as compared with placebo, resulted in a significantly higher rate of treatment response, without reducing opioid-mediated analgesia.135,136 The most common AE of naloxegol are as follows: abdominal pain (21% of treated patients), diarrhea (9%), nausea (8%), flatulence (6%), vomiting (5%), headache (4%), and hyperhidrosis (3%).

Naloxegol was approved by the FDA for the treatment of OIC in adult patients with chronic non-malignant pain. Naloxegol is contraindicated in patients with known or suspected GI obstruction and those at increased risk of the potential for GI perforation, patients receiving strong CYP3A4 inhibitors (may increase exposure to naloxegol and precipitate opioid withdrawal symptoms) and patients with a known hypersensitivity reaction to naloxegol or any of its excipients [133 https://www.movantikhcp.com/access 6th December 2014].

**Alvimopan**

Alvimopan is used to restore bowel function after surgery, for no more than 7 days (15 doses) and cannot be prescribed for outpatients. However, it is contraindicated for the treatment of OIBD in patients with advanced diseases due to the risk of myocardial infarction.137–140

**Conclusion and recommendations**

OIBD is a common complication in patients receiving long-term treatment with opioids. Nonetheless, significant progress has been made with targeted therapies for the management of patients with symptoms of OIBD, namely OXN, MNTX, and naloxegol. Any intervention should always be preceded with a meticulous assessment of the OIBD symptoms in the wider context of the underlying disease and comorbidities, and a thorough patient’s assessment including non-medical problems should also be conducted.

The management of OIBD in the first instance may be based on traditional laxatives and prokinetics, especially newer agents such as prucalopride or lubiprostone. The second step may comprise a consideration of an opioid switch, especially from orally administered hydrophilic opioids, such as morphine, oxycodone, and hydromorphone, to more lipophilic transdermal opioids, such as buprenorphine, fentanyl, or drugs with non-opioid components, such as tapentadol or, in the case of mild-to-moderate pain, tramadol. When traditional laxatives and prokinetics are ineffective, the use of peripheral µ-opioid receptor antagonists may be considered, with a preference of oral agents, if available. Furthermore, it needs to be noted that there is uncertainty in the MNTX role, because of the reports of GI perforation. The third step in treating OIBD may constitute rectal measures, including suppositories and enemas. OXN may be considered as a first-step approach in patients who already suffer from or are in high risk of OIBD development. However, limitations of OXN include the currently recommended daily dose range (80 mg/40 mg), and the fact that the patients must have normal hepatic function and portal circulation.

**Future perspectives**

Several purely peripherally acting opioid receptor antagonists, which are currently under investigation, include naldemedine (S-297995; NCT01965652), benvenopran (CB-5945) and axelopran (TD-1211).141 Further progress can be made by a more in-depth exploration of different mechanisms (eg, naloxonazine-sensitive vs naloxonazine-insensitive sites, peripheral or central mechanisms) of OIBD development by different opioids associated with their different impact on the GI tract, and finally the development of more effective, targeted therapies aimed at specific opioid receptor sites.142 Controlled clinical studies are needed to demonstrate long-term efficacy, safety, and cost-effectiveness of new therapies, and to decrease the need for unpleasant and poorly accepted rectal interventions, thereby reducing OIBD costs and improving patients’ compliance to opioid therapy. As a result of a more effective OIBD treatment, lower GI symptom burden and better effectiveness of pain management may be achieved, with a positive influence on patients’ QoL.
Disclosure
The author reports no conflicts of interest in this work.

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