Amino Acids and Diet in Chronic Pain Management

Post-traumatic Headaches, Migraines, and Sleep Disorders

Comorbidity of Musculoskeletal Injury Pain and PTSD

Tension Headache

OPIOID ANTAGONISTS IN PAIN MANAGEMENT
Achieving effective, durable, and safe pain relief, especially in patients with chronic and/or severe pain conditions, can be a clinical challenge. For many types of pain, prescription opioids are among the most effective analgesics. However, there could be concerns about the development of opioid tolerance or adverse effects, and in some cases opioids seem to worsen pain (eg, hyperalgesia). For certain difficult conditions, such as fibromyalgia or neuropathies, opioids alone are sometimes considered of limited effectiveness.

Healthcare providers interested in pain management must be alert to new or novel approaches that help to overcome deficiencies of opioids, such as treatment-limiting side effects, and as aids in relieving difficult-to-treat pain conditions. In this regard, there is a growing body of evidence suggesting potential benefits of opioid antagonists.

Opioid antagonists—in particular, naloxone and naltrexone—have been available and studied for decades as agents that displace opioid molecules from their neuroreceptors, and block opioids from attaching to and activating those receptors. Such qualities can be of important benefit, as short-acting antagonists like naloxone are used effectively to quickly reverse toxic effects of opioid overmedication or overdose.

Laboratory research and clinical trials have demonstrated the unexpected, paradoxical effects of opioid antagonists as adjuvants for enhancing rather than attenuating analgesic effects of opioids like morphine, oxycodone, and others. Further benefits of opioid antagonists, as monotherapy, for better managing certain chronic pain conditions also have been discovered.

Available evidence suggests that the opioid antagonists naloxone and naltrexone offer potential benefits for enhancing opioid analgesia as well as monotherapy for managing certain challenging pain conditions.

Editor's note: This article is reprinted with permission and minor edits from the author’s peer-reviewed research report, “Opioid Antagonists, Naloxone & Naltrexone—Aids for Pain Management: An Overview of Clinical Evidence,” from Pain Treatment Topics; www.pain-topics.org.

This paper provides an overview of naloxone and naltrexone pharmacology, and briefly examines some of the theoretical foundations of opioid antagonists for pain management. Available evidence from the literature describing opioid-antagonist therapy in adult humans, as portrayed in case examples or clinical trials, is reviewed and summarized. It must be understood, however, that opioid antagonists are not yet FDA-approved as adjuvant analgesics or for other pain management purposes, so their uses described in this report are off-label. Further investigations to assess the safety and efficacy of these applications would be appropriate.

Opioid Antagonist Pharmacology

Opioid antagonists have been available for many decades and are well known for their applications in addiction treatment (naltrexone) and as an antidote for opioid overdose (naloxone). The first report of agents having opioid antagonist-like properties was in 1915, when N-allylnorcodeine was observed to block the respiratory-depressant effects of morphine and heroin. The opioid antagonist nalorphine was synthesized in the 1940s; however, it also had partial agonist activity and its unsettling effects on mood (causing dysphoria) discouraged widespread use for treating opioid intoxication or overdose.

In 1960, naloxone was synthesized as a more potent and less toxic antagonist than nalorphine. Naloxone produces no dysphoria but it has a short duration of action and poor oral bioavailability due to high first-pass hepatic metabolism; it is 15 times more potent by injection than by mouth. The need for a long-acting opioid antagonist as a treatment for addiction, by
blocks the euphoric effects of illicit opioids for an extended period of time, motivated the development of naltrexone in 1963. This antagonist has reasonably good oral bioavailability, a long duration of action, and twice the potency of naloxone (see Table 1).5,8,9

The development of these agents was facilitated by the interesting fact that relatively minor structural changes could convert an opioid agonist drug into one with antagonist actions; that is, blocking various opioid receptors and preventing their activation. Thus, morphine was transformed into nalorphine, and oxymorphone into either naloxone or naltrexone. A more recent development, nalmedine, is a pure mu-receptor antagonist that is at least equipotent with naloxone but longer acting, with a duration of action between naloxone and naltrexone. Additional antagonists that are selectively active at other opioid receptors have been developed largely for experimental purposes.7,8,9

Naloxone and naltrexone are the two opioid antagonists that have been most extensively studied and are commercially available today. They are FDA-approved for the treatment of alcoholism or opioid addiction (naltrexone; eg, Trexan®, Revia®, Depade®) or opioid overdose (naloxone; eg, Narcan®). A long-acting, high-dose depot form of naltrexone (Vivitrol®) for intramuscular injection also is approved for addiction therapy. In the interest of abuse deterrence rather than therapeutic enhancement, these antagonists also are being used or tested as ingredients in specially formulated opioid analgesics to deter their misuse or abuse. These agonist-antagonist combinations contain a sustained-release opioid with a sequestered antagonist, usually naltrexone. The concept is that the antagonist remains latent and is only released in the event of tampering, such as by crushing or dissolving the product, in an attempt to extract the abusable opioid ingredient. The activated antagonist would then impede desired opioid effects making the product less prone to abuse.13

As noted above, two important features distinguishing naloxone from naltrexone are the very limited oral bioavailability of naloxone and the extended action of naltrexone. Both, naltrexone and naloxone may interact with all classes of opioid receptors but they do so most strongly at mu-opioid receptors.3 However, the typical naltrexone doses used for addiction therapy (50–100 mg/day orally) and naloxone dosing for reversing opioid overdose (0.4–1.0 mg/dose IV or IM) are, in most cases, different from the dosing protocols used in pain management applications (discussed below). In general, the literature refers to doses of antagonists, particularly naltrexone, in the 1 to 5 mg range as “low dose”; whereas, those less than 1 mg, in microgram amounts, are usually designated as “ultralow dose.”

### Special Qualities of Opioid Antagonists

In an excellent review of the role opioid antagonists may play in pain management, Sloan and Hamann15 note that interest in this topic dates back to the 1950s. It was observed that low doses of the antagonist nalorphine had pain-relieving properties in humans comparable to low-dose morphine in some cases. It was suspected that a combined opioid agonist-plus-antagonist agent might offer significant pain relief without the undesirable side effects of opioids alone. However, this line of research was not pursued until quite recently.

It was later discovered that naloxone had an analgesic effect at very low doses but no such effect at higher doses. For example, experiments in opioid-naïve human subjects during the 1970s found that “low-dose” naloxone (up to 2 mg) had analgesic effects that were lacking at higher naloxone doses (eg, 7.5 mg and 10 mg).15 The proposed neurobiological mechanisms behind this are complex and still being defined.

Studies in opioid-free animals have shown that, by causing a transient blockade of opioid receptors, low doses of antagonists stimulate increased production, or upregulation, of mu-opioid receptors in regions of the brain that control pain responses.15 Therefore, it seems plausible that after antagonist effects wear off — which may take minutes or hours depending on the agent and dose — greater numbers of opioid receptors are available to bind with pain-relieving opioids in the circulatory system, whether externally administered (eg, morphine) or endogenous peptides (eg, endorphins). At the same time, it has been noted that the body responds to the temporary opioid-receptor blockade by producing substantially increased amounts of beneficial natural endorphins.15,16 However, if the opioid-receptor blockade by antagonist is too extensive or continues too long, these pain relieving benefits are not realized, so the dose and half-life of the particular antagonist are critical factors.

Laboratory experiments further demonstrated that opioid antagonists may help to prevent the potential hyperalgesic (pain enhancing) and other negative effects of opioid pain relievers. At a cellular level, it is believed that exposure to opioid analgesics at certain doses and for prolonged periods of time may lead to aberrant signaling patterns by opioid receptors. These patterns can

### Table 1. Pharmacokinetic Data

<table>
<thead>
<tr>
<th></th>
<th>Naltrexone</th>
<th>Naloxone</th>
</tr>
</thead>
<tbody>
<tr>
<td>Chemical Formula</td>
<td>C20H23NO4</td>
<td>C19H21NO4</td>
</tr>
<tr>
<td>Oral Bioavailability</td>
<td>Up to 40%</td>
<td>2% (high absorption but extensive first-pass metabolism)</td>
</tr>
<tr>
<td>Metabolism</td>
<td>Hepatic</td>
<td>Hepatic</td>
</tr>
<tr>
<td>Peak Concentration</td>
<td>1–2 hours</td>
<td>10 minutes</td>
</tr>
<tr>
<td>Half-life</td>
<td>Up to 14 hrs (oral)</td>
<td>30–81 min. (IV, IM)</td>
</tr>
<tr>
<td>Duration of Action</td>
<td>Up to 24+ hrs</td>
<td>1–4 hrs</td>
</tr>
<tr>
<td>Excretion</td>
<td>Renal, Biliary</td>
<td>Renal, Biliary</td>
</tr>
</tbody>
</table>

**Sources:** Crabtree 19846; Gonzales and Brogden 19888; Reisine and Pasternak 19969

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13
be reversed and restored to normal by the actions of opioid antagonists in very low concentrations.13,17

The ultimate effects of these mechanisms, demonstrated in animal studies and human trials, appears to be an enhancement by low- or ultralow-dose naloxone or naltrexone of opioid-agonist antinociceptive (pain relieving) efficacy. Along with this, tolerance to and physiologic dependency on opioid agonists, such as morphine and others, may be diminished.14 Essentially, appropriately low doses of opioid antagonists have been postulated to "reset" the opioid-receptor system for a period of time,15 which seems analogous to how rebooting a malfunctioning computer clears memory, refreshes the software, and often restores normal function. The best type of antagonist, the optimal dose of it for achieving these effects, and the duration of effects have been explored by various investigators, as described below. Note that in most of the literature, ultralow-dose applications of antagonists are usually expressed in microgram amounts. However, since the more well-known applications of naloxone and naltrexone are in milligram increments, doses in this paper are converted from micrograms to milligrams (1 mcg = .001mg).

Research on Naloxone in Pain Management

The traditional role of naloxone has been the complete reversal of opioid effects, particularly in cases of opioid overmedication or life-threatening overdose. However, the potentials of this agent in pain management have been described in a number of studies.

Loitman16 reported an interesting case series of three patients with intractable chronic pain—despite receiving large doses of strong opioids, other medications, and complementary procedures—who underwent a brief and temporary inpatient opioid-receptor reversal (that is, detoxification) procedure with intravenous naloxone (0.6–1.2mg). This induced rapid withdrawal as the opioids were displaced from their receptors by the antagonist. Distressing withdrawal symptoms were ameliorated by sedation and, after a brief period of time (about 20 minutes), opioid pain relievers were readministered.

All patients responded very positively to the procedure, achieving much greater pain relief than before the brief detoxification and at significantly lower doses of opioid analgesics. Overall, Loitman noted that vastly improved pain relief was achieved in less than one hour on about one-third of the original opioid dose. Although exact mechanisms were unclear, the opioid receptors were thought to have been “reset” (as discussed above), overcoming prior opioid tolerance and/or hyperalgesia.

A number of clinical investigations have suggested potential benefits of ultralow-dose oral naloxone administration for postoperative pain:

• In patients treated with buprenorphine (a mixed agonist-antagonist opioid) postoperatively and experiencing breakthrough pain, the addition of oral ultralow-dose naloxone (0.08–0.4mg) provided effective and long-lasting analgesia (median 22 hrs).17 A more recently reported investigation, by La Vincente and colleagues,18 also confirmed the enhancement of buprenorphine analgesia by ultralow-dose oral naloxone (exact dose unspecified).

• Oral naloxone alone, either 0.4 or 1.0 mg, produced significant analgesic effects compared with placebo for postoperative dental pain in a trial of 90 patients. In a second trial, oral naloxone (0.4mg) plus pentazocine (60 mg, a mixed agonist-antagonist opioid) produced greater dental-pain relief than monotherapy with pentazocine (60mg) or morphine (15mg). However, naloxone (0.4mg) plus morphine (8mg) produced less analgesia than morphine (8mg) alone, suggesting that analgesia augmentation by naloxone is influenced by the type and dose of opioid.19

• In a study involving postoperative IV PCA (patient-controlled analgesia), 60 patients received morphine under patient control, plus either placebo or naloxone—0.00025 mg/kg/hr or 0.001mg/kg/hr—via steady IV infusion. Good pain relief was experienced in all groups; however, side effects (nausea, vomiting, pruritus) were reduced by naloxone at either dose and consumption of PCA morphine was significantly reduced by the lowest naloxone dose, suggesting a morphine-enhancement effect of the antagonist.20

• Other postoperative trials compared IV PCA morphine alone versus IV PCA morphine plus an IV naloxone bolus (0.8mg) administered with each morphine dose (which was equivalent to 0.38 mg/kg/hr of naloxone on average). In the 92 patients studied, investigators found no differences between groups in pain relief, side effects, or morphine consumption.21 Comparing this study with the Gan et al.22 trial above suggests that a continuous naloxone infusion rather than bolus, and at doses of naloxone that are less than in this study, may be required to accrue beneficial effects during naloxone/morphone PCA.

• A more recently reported case series, incorporating intravenous nalbuphine (5mg) plus naloxone (0.2mg IV), used postoperatively in 4 women undergoing outpatient gynecological procedures, found improved analgesia with this mixture.23

• In a case report, a patient with chronic refractory pain following laminectomy was treated with an intrathecal combination of morphine (2mg) and ultralow-dose naloxone (.000020 mg; that is, 20 nanograms). Dramatic pain relief began within 20 minutes, peaked at 1 hour, and persisted with repeated infusions. Up to 80% improvement was reported by the patient while receiving an intrathecal infusion of 5mg morphine plus .000050mg naloxone daily during a 3-year followup period.24

The trials in postoperative patients suggest that combining low-dose nalox-
one with opioid agonists can be safe and enhance effectiveness of the analgesic. PCA-based studies further suggest that the augmentation of opioid-agonist effects during PCA is best with sustained ultralow-level antagonism of opioid receptors versus the more complete and intermittent blockade of those receptors by a somewhat higher dose of naloxone.13

Naloxone for Opioid-induced Constipation
Another proposed application of low-dose opioid antagonists has been to help prevent or relieve opioid-induced constipation that can be a problematic side effect of long-term opioid analgesics. Meissner et al.17 recently reported a clinical trial in 202 patients receiving oxycodone for chronic noncancer pain. Subjects administered 40, 60, or 80mg/day of extended-release oxycodone were randomized to also receive either 10, 20, or 40mg/day of extended-release oral naloxone or placebo.

There was no loss of oxycodone analgesic efficacy in patients administered naloxone at any of these higher doses, probably due to naloxone’s limited oral bioavailability. However, bowel function significantly improved with increasing naloxone dose, and there were no unexpected adverse events. A 2:1 dose ratio of oxycodone/naloxone was judged as most efficacious. Other large, randomized, placebo-controlled trials had found the same favorable results in patients with chronic noncancer pain and back pain.29,29

A combination product—using extended-release oxycodone and extended-release naloxone; 2:1 ratio—was recently approved for marketing in Europe under the name Targin®, with an emphasis on its approved for marketing in Europe under novel indication of opioid-induced constipation.30

Two other relatively new and specialized opioid antagonists, methylnaltrexone (Relistor®) and alvimopan (Entereg®), have been introduced for managing opioid-induced constipation. These agents block only peripheral opioid receptors in the gut. And, unlike naloxone and naltrexone, these antagonists do not cross the blood-brain barrier, so their actions potentially reverse opioid-induced constipation without precipitating withdrawal symptoms or altering analgesic qualities of opioids.31

The FDA approved oral alvimopan in 2008 to accelerate the restoration of normal bowel function in hospitalized adult patients who have undergone partial large or small bowel resection surgery.32 At about the same time, methylnaltrexone bromide was FDA-approved as an injectable medication to help restore bowel function in adult patients with advanced illness who are receiving analgesic opioids on a continuous basis.33

Naloxone Potential Side-Effects
On rare occasions, nausea, vomiting, hypertension, pulmonary edema, tachycardia, arrhythmia, and cardiac arrest have been implicated with naloxone doses as small as 0.1-0.4 mg.34,35

Applications of Naltrexone
As a longer-acting, orally-bioavailable antagonist, naltrexone is easy to administer and may offer advantages over naloxone. In opioid-tolerant animals, naltrexone was shown to produce increased response to morphine, possibly by upregulating the opioid-receptor system. Also of interest, animal experiments suggest that very low doses of naltrexone appear to enhance opioid analgesia to a greater extent when the amount of opioid drug is actually reduced,36,37 which proposes an opioid-sparing effect in humans. A number of investigations in human subjects have explored the potential of low- or ultralow-dose naltrexone as an opioid adjuvant and as monotherapy for the management of several pain-related conditions.

Crohn’s Disease. In 2007, successful results of the first clinical trial of low-dose naltrexone for active Crohn’s disease were published by investigators at the Pennsylvania State University College of Medicine.16 In an open-label study, 17 patients were administered low-dose oral naltrexone (4.5 mg/day at bed-time) for 12 weeks. Statistically significant improvements were found in CDAI (Crohn’s Disease Activity Index) scores and quality of life indicators, increased rates of remission, and decreases in inflammatory markers in serum. Overall, 89% of patients exhibited a favorable response to naltrexone therapy, 67% achieved remission, and 2 subjects with open fistulas had closure. Side effects of therapy were reportedly minimal, with sleep disturbances in 7 patients most common. Reported, the National Institutes of Health has granted funding for this same group of researchers to conduct a larger placebo-controlled trial.

Irritable Bowel Syndrome. Kariv et al.38 reported an open-label pilot study of ultralow-dose naltrexone (0.5 mg/day) in 42 patients with confirmed irritable bowel syndrome (IBS). During 4 weeks of treatment and 4 weeks of followup, global assessments of symptom relief improved in about three-quarters of the subjects (76%) and mean numbers of pain-free days per week were modestly but significantly increased from baseline. Naltrexone therapy was well tolerated, without any significant adverse reactions.

Despite this initial success, a large, Phase III multi-center, randomized, placebo-controlled trial of oral naltrexone for treating IBS in 600 women was discontinued early due to unacceptable outcomes.39 Naltrexone demonstrated a favorable safety profile, and patients reported statistically meaningful relief of painful IBS symptoms during the second month of treatment compared with placebo. However, these beneficial effects did not persist into the third month, which was the primary endpoint of interest. Possibly, a critical factor was that this trial (and the one by Kariv et al.38 above) used 0.5 mg/day of naltrexone, which is well below the 4.5mg every night that others have recommended as monotherapy [described at LDNinfo.org40; Smith et al41; also see, fibromyalgia, below].

Fibromyalgia. Low-dose naltrexone (LDN) has been purportedly used off-label for the treatment of fibromyalgia for some time, and a small clinical study (n=10) funded by the American Fibromyalgia Syndrome Association (AFSA) was conducted in 2007 at the Stanford University Systems Neuroscience and Pain Lab.41 Details and results of this single-blind trial of LDN for the treatment of fibromyalgia have not been published (as of February 2009), but according to some reports the outcomes were successful and the majority of study participants asked to continue on drug therapy after the conclusion of the study [4.5 mg/day oral naltrexone, per correspondence with researcher S. Mackey42]. Side-effects were essentially nonexistent, except for 3 reports of mild sleep disturbance [also see, LDN for Fibromyalgia43].

Enrollment is ongoing, as of January 2009, for a large double-blind, placebo-controlled trial of LDN for fibromyalgia.44 While the exact mechanisms of naltrex-
one benefits for fibromyalgia are unknown, the researchers suggest it is possible that low doses of the drug block receptors on glial cells that are responsible for fibromyalgia symptoms. Prior to being experimentally administered, it has not been reported in the literature, and the role of the nervous system in this syndrome and its appropriateness as a target for treatment require further definition. The potential for ultralow-dose naltrexone to benefit neuropathic pain conditions needs further exploration.

Neuropathic Pain. Recently reported laboratory experiments, and a case example, suggest benefits of opioid antagonists in ameliorating neuropathic pain. Prior investigations demonstrated that activated spinal cord glial cells (microglia and astrocytes) contribute importantly to neuropathic pain; although, how nerve irritation or injury activate glia is still undefined. In rodents, Hutchinson et al. described a patient with chronic, painful diabetic neuropathy in humans. Cruciani et al described a patient with chronic painful diabetic neuropathy being treated unsuccessfully with methadone. The addition of ultralow-dose naltrexone (0.002 mg/day) dramatically improved pain relief and, at the same time, allowed a 16% reduction in methadone dose.

Based on laboratory experiments and the very limited clinical evidence, low- or ultralow-dose naltrexone might potentially benefit reflex sympathetic dystrophy (RSD, also known as complex regional pain syndrome or CRPS). However, this has not been reported in the literature, and the role of the nervous system in this syndrome and its appropriateness as a target for treatment require further definition. The potential for ultralow-dose naltrexone to benefit neuropathic pain conditions needs further exploration.

Naltrexone Potential Side-Effects

Administering larger doses of naltrexone to patients maintained on pure opioid agonists for pain can result in withdrawal signs and symptoms. These may be more sustained than those encountered with naltrexone due to the longer half-life of naltrexone and its significant concentrations in the central nervous system.

Naltrexone Combined with Opioid Agonists

Successful Phase II trials and Phase III trials have been completed on a combination of naltrexone plus intrathecal morphine infusion. In the first trial, 243 patients with chronic, painful osteoarthritis were administered placebo, oxycodone QID (4x/day), or oxycodone + oxycodone + naltrexone (Oxytrex) either QID or BID (2x/day). The total daily dose of oxycodone (titrated up to 80 mg/day or less) was the same in all treatment groups on average, but the Oxytrex groups additionally received either 0.002 or 0.004 mg naltrexone per day. Oxytrex BID (ie, oxycodone + 0.002 mg/day naltrexone) provided significantly superior pain relief compared with oxycodone alone QID, Oxytrex QID, or placebo.

The potential benefits of such agonist/antagonist combinations seem promising; however, this may depend on the type of opioid and/or the naltrexone dose. In an experimental trial in healthy humans, using a thermal discomfort test, the addition of 0.25, 0.5, or 1.0 mg of oral naltrexone neither aided nor deterred analgesic effects of oral hydrocodone. These findings are very preliminary, and the naltrexone doses in this study were many-fold larger than in the Oxytrex formulation, so further research on combination products seems warranted.

Hamann and Sloan reported a clinical pilot study to assess the analgesic efficacy of ultralow-dose oral naltrexone combined with intrathecal morphine infusions in patients with chronic, refractory nonmalignant pain. Subjects being administered continuous intrathecal morphine were randomized in double-blind fashion to receive either oral placebo (n=5) or oral naltrexone—0.1 mg (n=3) or 0.01 mg (n=7)—every 12 hours for 7 days. The group receiving 0.1 mg BID (0.2 mg/day total) of adjuvant naltrexone exhibited significant reductions in pain intensity and greater pain relief compared with placebo; whereas, the 0.01 mg BID naltrexone dose (0.02 mg/day total) was ineffective. There were no serious adverse events or evidence of opioid withdrawal, and side effects were minor and similar across all treatment groups (eg, headache, nausea, sedation, dry mouth).

Aid for Opioid Tapering

Another interesting application has been the use of very low doses of naltrexone to diminish withdrawal signs/symptoms in
## Table 2. Summary - Opioid Antagonists for Pain in Human Trials/Cases

(See text for further discussion.)

<table>
<thead>
<tr>
<th>Application</th>
<th>Antagonist Dose</th>
<th>Clinical Outcome</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Naloxone (IV, Intrathecal, or Oral – as specified)</strong></td>
<td></td>
<td></td>
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</tr>
<tr>
<td>Brief opioid detoxification for intractable pain</td>
<td>0.6–1.2mg IV (ca. 20 minutes)</td>
<td>Significant pain relief at lower opioid doses. Requires close monitoring.</td>
<td>Loitman 2006&lt;sup&gt;19&lt;/sup&gt;</td>
</tr>
<tr>
<td>+ Buprenorphine for postop breakthrough pain</td>
<td>0.08–0.4mg oral</td>
<td>Effective, long-lasting analgesia</td>
<td>Schmidt et al. 1985&lt;sup&gt;20&lt;/sup&gt;; La Vincente et al. 2008&lt;sup&gt;21&lt;/sup&gt;</td>
</tr>
<tr>
<td>Monotherapy for acute postop dental pain</td>
<td>0.4 or 1.0mg oral</td>
<td>Significant analgesia compared with placebo.</td>
<td>Levine et al. 1988&lt;sup&gt;22&lt;/sup&gt;</td>
</tr>
<tr>
<td>+ Pentazocine (60mg) for dental pain</td>
<td>0.4mg oral</td>
<td>Greater analgesia than monotherapy with pentazocine or 15mg morphine.</td>
<td>Levine et al. 1988&lt;sup&gt;22&lt;/sup&gt;</td>
</tr>
<tr>
<td>+ PCA morphine postop</td>
<td>0.00025 or 0.001 mg/kg/h steady infusion IV</td>
<td>Reduced side effects and lower morphine consumption versus placebo</td>
<td>Gan et al. 1997&lt;sup&gt;23&lt;/sup&gt;</td>
</tr>
<tr>
<td>+ PCA morphine postop</td>
<td>0.8mg dose/IV bolus (= 0.38 mg/kg/h on average)</td>
<td>No benefit.</td>
<td>Sartain et al. 2003&lt;sup&gt;24&lt;/sup&gt;</td>
</tr>
<tr>
<td>+ Nalbuphine (5mg IV) postop</td>
<td>0.2mg IV</td>
<td>Improved analgesia.</td>
<td>Gordon et al. 2007&lt;sup&gt;25&lt;/sup&gt;</td>
</tr>
<tr>
<td>+ Intrathecal morphine (5mg) for intractable chronic pain</td>
<td>0.000050 mg/day intrathecal during followup</td>
<td>Significantly improved ongoing pain relief.</td>
<td>Hamann et al. 2008&lt;sup&gt;26&lt;/sup&gt;</td>
</tr>
<tr>
<td>+ Oxycodone (chronic noncancer pain; chronic back pain)</td>
<td>10, 20, 40 mg/d oral</td>
<td>No loss of analgesia, improved bowel function at 2:1 oxycodone/naloxone ratio.</td>
<td>Meissner et al. 2009&lt;sup&gt;27&lt;/sup&gt;; Simpson et al. 2008&lt;sup&gt;28&lt;/sup&gt;; Vondrackova et al. 2008&lt;sup&gt;29&lt;/sup&gt;</td>
</tr>
<tr>
<td><strong>Naltrexone (Oral)</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Crohn’s Disease</td>
<td>4.5 mg/day</td>
<td>Favorable response in 89% of patients.</td>
<td>Smith et al. 2007&lt;sup&gt;30&lt;/sup&gt;</td>
</tr>
<tr>
<td>Irritable Bowel Syndrome (IBS)</td>
<td>0.5 mg/day</td>
<td>Pain/symptom relief during first 2 months.</td>
<td>Kariv et al. 2006&lt;sup&gt;31&lt;/sup&gt;</td>
</tr>
<tr>
<td>Fibromyalgia</td>
<td>4.5 mg/day</td>
<td>Unknown, but anecdotal reports favorable.</td>
<td>SNAPL 2008&lt;sup&gt;32&lt;/sup&gt;</td>
</tr>
<tr>
<td>Neuropathic Pain (combined with methadone)</td>
<td>0.002 mg/day</td>
<td>Improved analgesia at lower methadone dose.</td>
<td>Cruciani et al. 2003&lt;sup&gt;33&lt;/sup&gt;</td>
</tr>
<tr>
<td>+ Oxycodone (Oxytrex®) for osteoarthritis or chronic back pain</td>
<td>0.002 mg/day</td>
<td>Reduced opioid consumption and side effects, including resolution of opioid-induced constipation.</td>
<td>Chindalore et al. 2005&lt;sup&gt;34&lt;/sup&gt;; Webster et al. 2006&lt;sup&gt;35&lt;/sup&gt;</td>
</tr>
<tr>
<td>+ Hydrocodone (experimental)</td>
<td>0.25, 0.5, or 1.0 mg</td>
<td>No beneficial effect.</td>
<td>Cantilena et al. 2004&lt;sup&gt;36&lt;/sup&gt;</td>
</tr>
<tr>
<td>+ Intrathecal morphine for chronic noncancer pain</td>
<td>0.2 mg/day</td>
<td>Reduced pain intensity, improved pain relief.</td>
<td>Hamann and Sloan 2007&lt;sup&gt;37&lt;/sup&gt;</td>
</tr>
<tr>
<td>Opioid (methadone) tapering</td>
<td>0.125 or 0.250mg in multiple doses/day</td>
<td>Reduced withdrawal symptoms and opioid craving compared with placebo.</td>
<td>Mannelli et al. 2006&lt;sup&gt;38&lt;/sup&gt;, 2008&lt;sup&gt;39&lt;/sup&gt;</td>
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</tbody>
</table>
patients being tapered from long-term opioid therapy. Mannelli et al 55 describe a study in methadone-maintained patients administered ultralow-dose oral naltrexone (0.125-0.250 mg/dose) during inpatient opioid detoxification. Overall signs of withdrawal were repressed, even as naltrexone doses were increased daily, and the opioid-tapering process was completed without adverse incidents or patient discomfort.

A larger (n = 174), randomized, placebo-controlled study by this group 54 found similar benefits of ultralow-dose oral naltrexone (0.125mg or 0.250 mg/dose) combined with tapered methadone during a 6-day outpatient detoxification protocol. Naltrexone-treated patients experienced milder withdrawal symptoms and reduced opioid craving compared with placebo-group subjects. There were no treatment-related adverse events reported.

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**Extensions of Low-Dose Naltrexone (LDN) Theory & Application**

Besides the potential advantages of opioid antagonists for pain management described above, off-label low-dose naltrexone (LDN) protocols have been reported as benefiting a number of important immunological and neurological disorders. This is believed due at least in part to the observation that many body tissues, including virtually every cell of the body’s immune system, have receptors that are routinely stimulated by endogenous opioid peptides—including endorphins and enkephalins. 55 Therefore, by acting at those receptors, LDN has been demonstrated as possibly having important capabilities to modulate the immune system and promote healing. 15,40

LDN has been shown to halt disease progression in certain cancers and to reduce symptoms of autism. 40 Some authors have recently suggested that, since LDN can upregulate endogenous opioid activity, it may have a role in promoting stress resilience and emotional well-being, as well as the amelioration of depression. 40

The typical therapeutic dosage range for LDN in these applications is from 1.75mg to 4.5mg taken at bedtime. It has been suggested that doses below this range are likely to have little effect for neurodegenerative and other disorders, while doses above this range could block endogenous opioid peptides for too long a period of time and interfere with naltrexone’s effectiveness. 40

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**Prescribing Opioid Antagonists**

Opioid antagonists are non-scheduled drugs that may be prescribed by any appropriately licensed health-care provider. However, low- or ultralow-dose formulations of naloxone and naltrexone are not commercially available and must be specially compounded by pharmacies equipped for this purpose.

The website, LDNinfo.org, under the editorial direction of David Gluck, MD, provides recommendations regarding the ordering of low-dose naltrexone. 41 Apparent ly, low-dose naltrexone prescriptions in 1mg to 5mg doses are being filled by many local and mail order pharmacies, and Gluck notes that a 30-day supply of 4.5mg naltrexone capsules may cost from $15 to $38 (USD). He specifies that pharmacies should be instructed to not provide naltrexone in slow-release (SR) or timed-release form. Furthermore, calcium carbonate fillers should not be employed in compounding capsules because they may interfere with naltrexone absorption.

The website also cautions that it is important that the compounding pharmacy has a reputation for consistent reliability in the quality and accuracy of opioid antagonist preparations that it delivers. (See ‘Compounding Considerations’ below.) In one investigation, the FDA found significant error rates in compounded prescriptions produced at randomly selected pharmacies. More than a third (34%) of sampled products failed one or more standard quality tests performed by the FDA, including inadequate potency in 90% of the failing products. 56

Recommendations are lacking regarding where to order naloxone or naltrexone in ultralow, microgram doses (<1.0mg). However, contacts with several compounding pharmacies confirm that both naltrexone and naloxone powders are readily available and most of those pharmacies are well-equipped to prepare doses in any amounts specified. 56 Presumably, the same cautions as above would apply to ultralow-dose formulations; ie, avoidance of slow-release formulations and calcium carbonate fillers.

**Compounding Considerations**

Compounding pharmacies must use an analytical balance with a very high degree of precision and without fluctuations or loss of product. For extremely low doses (below the sensitivity of a scale) an aliquot may be used. This involves adding active ingredient measured to the limits of the scale plus a proportional amount of diluent to make a “stock” supply of known strength. From this, the exact amount containing the desired dose of active ingredient is withdrawn and packaged.

**Summary And Conclusions**

Select applications of naloxone and naltrexone may offer novel and potentially effective approaches for resolving difficult pain management challenges. Table 2 briefly summarizes the applications, doses, and outcomes in research on adult human subjects that were described in more detail above.

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6. Bernhard Bihari, MD, in New York discovered its beneficial effects in the newly emerging disease of HIV/AIDS. 15

7. Mannelli et al 55 describe a study in methadone-maintained patients administered ultralow-dose oral naltrexone (0.125-0.250 mg/dose) during inpatient opioid detoxification. Overall signs of withdrawal were repressed, even as naltrexone doses were increased daily, and the opioid-tapering process was completed without adverse incidents or patient discomfort.

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12. Select applications of naloxone and naltrexone may offer novel and potentially effective approaches for resolving difficult pain management challenges. Table 2 briefly summarizes the applications, doses, and outcomes in research on adult human subjects that were described in more detail above.
Both of these agents have passed animal and clinical toxicity studies, and are FDA-approved for applications other than those described in this paper. Therefore, it is not surprising that they have exhibited favorable safety profiles when applied at low- and ultralow-dose levels, with few if any reports of adverse events or side effects at these doses when used individually as monotherapy or in combination with opioid analgesics.

Explanatory mechanisms of action behind the benefits of opioid antagonists in pain management are still under investigation. A significant body of laboratory research, in animals or tissue cultures, suggests that the opioid-receptor system is very complex, affecting a variety of tissue and cell types, and with functions extending beyond antinociception (ie, pain-relief). It appears that blockade of opioid receptors exerted by very small concentrations of naloxone or naltrexone may stimulate a cascade of actions at the cellular and molecular levels that may contribute to their multiple beneficial effects. Some of those proposed actions were briefly described in this paper.

For the most part, clinical research to date on low- or ultralow doses of opioid antagonists in humans has been limited. Except for several larger, controlled trials, reports in the literature have included small numbers of patients or represented case examples. Collectively, the research appears to offer some suggestions for the application and dosing of naloxone or naltrexone, but there are no well-established or standardized protocols when it comes to their use in pain management. Still, the available evidence, as summarized in Table 2, does propose a number of general possibilities that may be of interest to pain practitioners and their adult patients:

1. Brief detoxification via IV naloxone for difficult cases of opioid-unresponsive intractable pain, opioid tolerance, or suspected opioid-induced hyperalgesia.
2. Ultralow-dose oral naloxone combined with various opioid agonists for managing acute postoperative pain.
3. Adjuvant ultralow-dose naloxone (continuous IV infusion) combined with patient controlled analgesia (PCA) postoperatively.
4. Ultralow-dose naltrexone (oral) or naloxone (intrathecal) as a component of intrathecal opioid analgesia for difficult cases of intractable pain.
5. Ultralow-dose naltrexone combined with opioid agonists to provide an opioid-sparing effect, offering equivalent pain relief at lower opioid doses.
6. Oral ultra-low dose naloxone or naltrexone combined with oral opioid analgesics to help prevent or reverse opioid-induced constipation and potentially ameliorate other opioid side effects.
7. Ultralow-dose naltrexone to help facilitate more comfortable opioid-agonist tapering.

Certainly, further research, including large-scale clinical trials, are needed to better understand the capabilities and safety of opioid antagonists for treating pain conditions of various types. Meanwhile, healthcare providers interested in using low- or ultralow-dose naloxone or naltrexone would need to cautiously prescribe these agents off-label for compounding at properly equipped pharmacies.

Clearly, the type of antagonist (naloxone or naltrexone), mode of administration (oral, IV, IT), the dose (low or ultralow), the pain disorder being treated, comorbidities, and concurrent opioid analgesia can all interact to make important differences. For example, the first commercial agonist/antagonist combination products — Targin, approved in Europe, and Oxytrex, in development — have used vastly differing doses of either naloxone or naltrexone, respectively, due to the dissimilar oral bioavailabilities of the two antagonists.

Hopefully, other opioid analgesics incorporating naloxone or naltrexone for pain management applications will be formally tested and developed. Whether or not sponsors will come forward to initiate the necessary but costly clinical trials and seek appropriate regulatory agency approvals remains to be seen.

As monotherapy, the development of FDA-approved formulations of low- and ultralow-dose naloxone or naltrexone used alone for pain management purposes may face some formidable hurdles. For example, at a U.S. government-sponsored conference, it was noted that, “[Low-dose naltrexone] has been presented to big pharmaceutical companies, but there has been little support provided, as LDN is commercially available and it is unclear whether or not there could be patent coverage for a new indication at [the very low dosages].”

In sum, the opioid antagonists naloxone and naltrexone appear to offer promise for helping to achieve more effective pain management in some of the most difficult and challenging situations facing healthcare providers and their adult patients. Naloxone and naltrexone are available today and, although they are approved for other applications, it is important that practitioners become aware of the therapeutic options that these agents may provide for patient care.

“Naloxone and naltrexone are available today and, although they are approved for other applications, it is important that practitioners become aware of the therapeutic options that these agents may provide for patient care.”

Competing Interests Disclosure/Disclaimer
The author and reviewers report no competing interests. Pain Treatment Topics is supported in part by educational grants from Purdue Pharma L.P. and, formerly, from Covidien/Mallinckrodt. These companies manufacture certain products mentioned in this report; however, they neither suggested this report topic nor did they have any role in its development, review, or revision.

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Opioid Antagonists in Pain Management

The contents of this report are for educational purposes and are not intended to endorse or promote the off-label prescribing of any drugs. Practitioners are advised to study the available evidence and use professional discretion in their prescribing decisions.

Report Reviewers
Pain Treatment Topics acknowledges and thanks the following professionals for their expert reviews and assistance:

Jeffrey Fudin, BS, PharmD, DAAPM. Clinical Pharmacy Specialist, Pain Management, VAMC Albany; Adjunct Associate Professor of Pharmacy Practice, Albany College of Pharmacy & Health Sciences; and, Diplomat, American Academy of Pain Management.

Edward Hesterlee, PharmD. Associate Professor of Pharmacy Practice, University of Arkansas Medical Sciences (UAMS) College of Pharmacy, Little Rock, Arkansas, and a consultant to the pharmaceutical industry and national pharmacy associations.

Seth I. Kaufman, MD. Pain management and palliative care practitioner, West Clinic, Baptist Hospital, and Methodist Hospital, Memphis, Tenn.

Lee A. Krall, PharmD, BCGPS. Faculty member, Center for Pain Medicine and Regional Anesthesia, University of Iowa Hospitals and Clinics, Iowa City, Iowa.

James D. Toombs, MD. Staff physician, Division of Primary Care/Pain Medicine, Harry S. Truman Memorial Veterans’ Hospital, Columbia, Missouri.

Stewart B. Leavitt, MA, PhD is the founding Publisher/Editor-in-Chief of Pain Treatment Topics and has more than 25 years of experience in healthcare education and medical communications serving numerous government agencies, pharmaceutical companies, and private organizations. He was educated in biomedical communications at the University of Illinois at Chicago College of Medicine and then served as a Commissioned Officer in the U.S. Public Health Service at the National Institutes of Health, Clinical Center, Bethesda. He then went on to earn Masters and Doctorate degrees specializing in health/medical education at Northwestern University, Illinois, where he was an instructor and a Ford Fellow researching urban healthcare delivery systems. He is a member of the American Academy of Pain Management and a founding member of the International Association for Pain & Chemical Dependency.

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