**Review**

**Synthèse**

**Principles of opioid use in chronic noncancer pain**

Jacqueline Gardner-Nix

**Case 1**

A 33-year-old truck driver presented with left shoulder pain dating from a motor vehicle accident 4 years before. His pain was described as “constant” and “constricting,” with “stabbing electric, shock-like” characteristics. It had increased over the past 2 years to intensities between 7 and 10 out of 10 (0 = no pain, 10 = excruciating pain). He had undergone excision of the lateral end of the left clavicle and rotator cuff decompression 3 years earlier, receiving physiotherapy before and after surgery. He had also used heat and a machine providing transcutaneous electric nerve stimulation and had received intra-articular injections of anaesthetic agents and steroids. When first seen in the pain clinic he was taking, on average, one tablet of acetaminophen/codeine/caffeine (300 mg/30 mg/15 mg) and one tablet of acetaminophen/oxycodone (325 mg/5 mg) 5 times daily. He also drank 10 cups of coffee daily, had been unable to sleep more than 3 hours in 24 for the last 2 years and was working night shifts.

Apart from smoking a pack of cigarettes per day, the patient had no other risk factors for addiction, and he described having a supportive wife and 2 daughters. What modifications to his medication regimen and lifestyle might provide him with better pain management?

**Case 2**

A 40-year-old man came to the pain clinic with a 15-year history of ankylosing spondylitis and a 5-year history of rheumatoid arthritis. His medications for the last 2 years included prednisone (20 mg/d), methotrexate (22.5 mg/wk intravenously) and amitriptyline (50 mg at bedtime). In addition, over the last year he had been taking sustained-release morphine, the dose having been gradually increased from 30 to 115 mg every 12 hours. He had never had good pain control since the onset of the rheumatoid arthritis and had been extremely constipated for several months despite taking two 5-mg tablets of bisacodyl and two 100-mg tablets of docusate sodium 4 times daily, as well as using a Fleet enema intermittently. There was evidence of inflammatory activity in his joints, particularly his right elbow and left knee, and he attended appointments in a wheelchair.

The patient was receiving active management from his rheumatologist (methotrexate and intra-articular steroid injections), had no family or personal history of abusing alcohol or recreational or prescription drugs, and had a supportive wife and 2 young children. What changes to his analgesic regimen might improve his pain management?

**The last decade has brought acceptance that opioids can be used as a component of the management of noncancer pain that is chronic (lasting more than 6 months) when other approaches have failed and the quality of life is poor because of the pain. In recent years the annual consumption of morphine, including for noncancer pain, has risen substantially in developed countries.**

Recent guidelines and evidence-based recommendations for the use of opioids in conditions other than acute and cancer pain have advised the use of sustained-release or long-acting opioids, with avoidance of repeated injection, for chronic pain.

Several randomized controlled studies have demonstrated the efficacy of opioids in a variety of noncancer conditions, including postherpetic neuralgia and low back pain. The World Health Organization (WHO) analgesic ladder (Fig. 1) has opioids on the 2nd and 3rd steps, to be used when other strategies, such as physiotherapy, massage and therapy with non-opioid medications (acetaminophen, nonsteroidal anti-inflammatory agents, tricyclic antidepressants, anticonvulsants and topical preparations), have failed to provide sufficient pain control and the quality of life is poor because of the pain.

Perceived barriers to opioid prescribing (Box 1) are still contributing to inadequate pain management. It is now acknowledged that opioids may be appropriate in a subset of the population with a variety of conditions that cause chronic pain, including those that are impossible to diagnose exactly. For many nonmalignant conditions, opioids may allow lowered pain intensity (or more acceptable pain characteristics), an acceptable side effect profile, and enhanced function and quality of life. They may also allow other pain management strategies to become more accessible. This paper discusses approaches to the use of opioids in chronic noncancer pain.

**Fig. 1: World Health Organization analgesic ladder.** Least invasive routes of administration: transdermal and oral.
Diagnosing pain

Every effort should be made to diagnose and appropriately treat the condition underlying the pain, but pain still requires management even when diagnosis remains elusive. Most pain clinics prefer that patients have had an appropriate diagnostic workup (blood work, imaging, nerve conduction studies) before referral. Pain sites, qualities (throbbing, aching, shooting, burning, paresthesias, stabbing) and intensities should be recorded. Frequency and duration of pain exacerbations, triggers, aggravating and relieving factors, and effect on sleep and mood should be elicited. Several methods of assessing pain intensity exist: numeric scales (0 to 5 or 0 to 10, from no pain to excruciating pain), verbal scales (none, mild, moderate, severe), visual analogue scales and facial series (sad to happy, most often used for children). Scales are only useful in assessing progress within the same person, and they likely incorporate psychosocial influences. The best and worst pain scores should be recorded at each visit, along with an estimate from the patient of the proportion of time at different intensities, including the proportions at the lowest and the average intensities. Functional impairment and changes in impairment caused by pain and pain management should be recorded, with information from family members (with the patient’s permission) to corroborate function and status.

Choice of opioid

Opioids act at receptors on the pre- and postsynaptic membranes of neurons in the central and peripheral nervous system. Three major families of opioid receptors have been described: µ, κ and δ. Most of the opioids used clinically act on the μ receptors. Work in animal models and cell cultures has demonstrated multiple μ-receptor subtypes, which may support clinical observations that the response to a given opioid, with respect to both efficacy and side effects, can vary widely between individuals, probably owing to genetic polymorphism. About 7% to 10% of white people respond poorly or not at all to codeine; they lack the cytochrome P450 enzyme CYP2D6, which converts codeine to its active metabolites, including morphine. No trend has been reported in the literature regarding greater effectiveness of specific opioids for specific pain conditions. A patient cannot be considered unresponsive to opioids until all appropriate opioids have been tried.

Codeine and oxycodone, plain or in combination with acetaminophen, are usually perceived to be on the 2nd step of the WHO analgesic ladder, but any of the strong opioids (fentanyl, hydromorphone, methadone, morphine and oxycodone) can be considered to be on the 2nd or 3rd step because in lower doses they are equivalent to combinations of codeine or oxycodone with acetaminophen.

Titration is usually done with long-acting sustained-release opioids (Table 1) to promote smoother pain relief and improved compliance, though some physicians prefer to titrate short-acting opioids and convert to a sustained-release form when a maintenance dose is achieved. Selection of the initial opioid is based on the criteria shown in Table 2. For example, if the efficacy and side effect profile of an acetaminophen/oxycodone combination had been acceptable, sustained-release oxycodone could be used initially, but if prolonged sedation, nausea or intractable constipation had been present, transdermal fentanyl might be a wiser choice. And if the patient’s answers to the CAGE-AID questionnaire (Box 2) suggest a history of drug abuse, methadone might be selected and short-acting opioids for “rescue” analgesia not be offered. Once an opioid is chosen, the dose is increased to balance improved pain control and function against side effects of the opioid.

Decisions to switch to alternatives after titration with the initial opioid may be based on the approaches outlined in Table 3. Approximately equianalgesic dosages of other opioids (Table 1) may vary with the individual, and a 10% to 20% decrease in the equianalgesic dose is recommended when switching because different types of opioid receptors are affected by different opioids. Caution should be exercised when switching to methadone because it is about 7 to 10 times more potent than morphine when given long term. Very low doses of methadone (2.5 to 5 mg every 12 hours or, in patients over 65 years of age, 1 mg every 12 hours) are advisable while the dosage of the initial opioid is being tapered. Relatively small doses of methadone can provide excellent analgesia in certain individuals in whom larger doses of alternative opioids have
been unsuccessful, but this approach requires slow titration over several weeks, as a reservoir of methadone builds in the patient’s fatty and other tissues. Sometimes pain is controlled only after several weeks of a stable dose of methadone.

The prescription of opioids for chronic pain in some patients, such as the elderly and those with hepatic or renal disease, deserves special caution because of the potential accumulation of toxic metabolites in these people. Accumulation of active metabolites such as morphine-3-glucuronide and morphine-6-glucuronide may cause reduced cognitive function, increased anxiety, allodynia and clonus. The accumulation varies with the individual and the mode of drug delivery. Often for patients over 65 the initial dose is lower and the titration slower than for younger patients. Impairment of cognitive function due to the accumulation of active metabolites may be alleviated by a switch to transdermal fentanyl, which is metabolized in the liver to inactive metabolites.

Patients may find their function and activity levels enhanced by judicious use of “rescue” or “breakthrough” analgesia. For exacerbations of pain that are spontaneous or related to activity, stress or other factors, rescue analgesia can be provided by a short-acting opioid, taken from a few times a month to several times a day. The dose should be tailored to the maintenance analgesic dose (one-third to one-sixth of the 12-hourly dose), and the frequency should be tailored to the patient’s needs and response. Some patients find the effectiveness reduced if the drug is used too frequently. Patients with addictive tendencies or rebound headaches may be best managed with only sustained-release long-acting opioids.

Challenges to opioid prescribing

Between 3% and 17% of people are potential addicts; these individuals usually have a family history or a personal history of addiction. Addicts crave a substance, often initially seeking its euphoric effects but later taking the drug just to

Table 1: Long-acting sustained-release opioids available in Canada

<table>
<thead>
<tr>
<th>Opioid</th>
<th>Sustained-release brand names</th>
<th>Dosing frequency (h)</th>
<th>Approximately equianalgesic dosages of short-acting opioids (opioid: morphine)</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Codeine</td>
<td>Codeine Contin</td>
<td>8–12</td>
<td>10 mg codeine: 1 mg morphine (10:1)</td>
<td></td>
</tr>
<tr>
<td>Morphine*</td>
<td>M S Contin, M S Contin Suppositories, M-Eslon, Kadian, Oramorph</td>
<td>8–12 except for Kadian, which is given once a day</td>
<td>1 mg oxycodone: 2 mg morphine (1:2)</td>
<td></td>
</tr>
<tr>
<td>Oxycodone</td>
<td>Oxycontin</td>
<td>8–12</td>
<td>1 mg oxycodone: 2 mg morphine (1:2)</td>
<td></td>
</tr>
<tr>
<td>Hydromorphone*</td>
<td>Hydromorph Contin</td>
<td>8–12</td>
<td>1 mg hydromorphone: 5 mg morphine (1:5)</td>
<td></td>
</tr>
<tr>
<td>Transdermal fentanyl</td>
<td>Duragesic</td>
<td>48–72</td>
<td>25-µg/h patch, changed every 48–72 hours: 45–60 mg daily morphine</td>
<td></td>
</tr>
<tr>
<td>Methadone</td>
<td>Metadol</td>
<td>8–12</td>
<td>Not established. About 10 mg morphine: 1 mg methadone</td>
<td></td>
</tr>
</tbody>
</table>

Note: NMDA = N-methyl-D-aspartate

*When converting from oral to parenteral route, reduce the dose by half.
feel normal, and will do anything to acquire it, despite causing themselves harm. However, people who use opioids for chronic pain usually experience physical dependence, which is different from addiction in that those with physical dependence usually experience withdrawal symptoms if they abruptly stop taking the drug instead of gradually tapering the dosage. The CAGE-AID\textsuperscript{15} and SISAP\textsuperscript{22} questionnaires (Boxes 2 and 3) are useful in the initial history-taking among patients with chronic pain to identify those at higher risk of addiction. Those at higher risk may be treated with opioids as long as controls such as stricter boundaries are defined in a signed written contract (Table 4)\textsuperscript{1} and steps such as using methadone as the opioid of choice or dispensing smaller quantities at defined intervals (e.g., weekly or biweekly at the local pharmacy) are in place.

Opioids are often perceived as imperfectly controlling chronic nonmalignant pain. Pain intensities often are not reduced below 4 or 5 on a scale of 0 to 10, even when doses are increased to the point at which side effects are unacceptable.

### Box 2: CAGE-AID questionnaire\textsuperscript{16}

In the past have you ever
- Felt that you wanted or needed to cut down on your drinking or drug use?
- Been annoyed or angered by others complaining about your drug use
- Felt guilty about the consequences of your drinking or drug use?
- Had a drink or taken a drink in the morning (eye opener) to decrease hangover or withdrawal symptoms?

AID = Adapted to Include Drugs

One positive response suggests caution in prescribing opioids; 2 or more positive responses suggest increased vigilance by the physician prescribing opioids.

### Table 3: Problems often leading to a switch in opioid

<table>
<thead>
<tr>
<th>Problem</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Persistent nausea</td>
<td>Often worse with morphine.</td>
</tr>
<tr>
<td>with or without clonus</td>
<td>Reducing dose may give poor analgesia; switching may improve analgesia and alertness.</td>
</tr>
<tr>
<td>Excessive sweating</td>
<td>Benztrapine (2 mg/d) may help if tolerated. Switching may also help or may be a “class effect.”</td>
</tr>
<tr>
<td>Intractable constipation</td>
<td>Transdermal fentanyl may reduce constipation when the constipation due to codeine and morphine is not relieved by high doses of laxatives.</td>
</tr>
<tr>
<td>Intractable itching</td>
<td>Opioids can cause histamine release that is only partly reduced by antihistamines. Switching from morphine to transdermal fentanyl may be successful. If the itching is a class effect, try 0.5 mL of 1 mg/mL naltrexone solution daily.</td>
</tr>
<tr>
<td>No pain relief</td>
<td>Increase dose; may be unresponsive and require switching.</td>
</tr>
<tr>
<td>Weight change</td>
<td>Common with methadone; consider switching.</td>
</tr>
<tr>
<td>Peripheral edema</td>
<td>Can be temporary and dose-dependent. Switching may eliminate edema. If it occurs with any opioid, consider using diuretics.</td>
</tr>
<tr>
<td>Sexual dysfunction,</td>
<td>Usually a class effect. Testosterone levels can decrease with opioid therapy; impotence and low libido are sometimes reported with all opioids but are sometimes resolved with switching.\textsuperscript{3} Oligomenorrhea and amenorrhea are dose-dependent and possibly due to hyperprolactinemia; refer to an endocrinologist or gynecologist for possible hormone replacement therapy.</td>
</tr>
<tr>
<td>oligomenorrhea and amenorrhea</td>
<td></td>
</tr>
<tr>
<td>Dyspnea</td>
<td>Dyspnea that is not respiratory depression can be a class effect and can be alleviated by switching or reducing the opioid dose. It is not alleviated much by asthma medications.</td>
</tr>
<tr>
<td>Dysphoria and mood changes</td>
<td>Depression, panic attacks and other mood changes can occur with all opioids and sometimes disappear on switching.</td>
</tr>
<tr>
<td>Sleep disturbances</td>
<td>Opioids can interfere with restorative sleep, but so can pain. If insomnia is due to the opioid, try switching. Try a sedative tricyclic antidepressant.</td>
</tr>
<tr>
<td>No drug plan</td>
<td>Sustained-release opioids are expensive. Methadone is an inexpensive alternative that should be tried before returning to inexpensive short-acting opioids, which need to be taken more often.</td>
</tr>
</tbody>
</table>
In contrast, pain intensities in acute and cancer pain can often be lowered to 0 to 3 out of 10. This refractoriness of chronic noncancer pain often leads physicians to conclude that psychologic factors are at play. However, experimental work in animals and cell cultures has demonstrated that depleted opioid receptors on damaged nerves, changes in the receptor-binding sites and down-regulation of the receptors may be the cause of the observed reduction in opioid efficacy. It is also probable that genetic polymorphism in humans influences pain perception and response to medications.

Although many patients report somewhat reduced efficacy after the first month of opioid administration, a small percentage appear to become markedly tolerant (the drugs appear to become almost entirely ineffective over relatively short periods, such as weeks). Opioid rotation will likely be effective in these patients because a different subtype of receptor will respond to the new opioid while the receptors that had previously responded recover. Anecdotal evidence suggests that combining opioids to lower pain intensity, change pain characteristics and retard the development of tolerance may be more effective than using a single drug by targeting a broader range of opioid receptors in those refractory to therapy with a single opioid.

There is no evidence in the literature to suggest that driving be denied to patients receiving opioid therapy for chronic pain except during titration to establish the maintenance dose and when there is a higher likelihood of cognitive dysfunction, as with older patients. A decision to ban driving depends on the judgement of the treating physician, in collaboration with the patient and family. A medical driving test may be considered.

Long-term effects of opioids

To date, there are few reports of toxic effects in organs from long-term opioid use in nonaddicts. When opioids are used responsibly at stable doses, a balance between pain control and side effects can be established, and these drugs can be considered safer than nonsteroidal anti-inflammatory agents or large quantities (> 2 g/d) of acetaminophen. There is one report of the potential for cardiac arrhythmias in female patients on high-dose methadone therapy for chronic pain. It is therefore advisable to do electrocardiography and measure the serum levels of electrolytes and magnesium before and after methadone therapy is started in patients who are likely to be receiving a high dose (> 200 mg/d).

Cases revisited

Case 1

The patient likely had neuropathic pain resulting from damaged nerves and ischemic pain resulting from constriction by scar tissue of the blood supply of nerves around the site of trauma and surgery. Sleep deprivation likely was contributing to his reduced ability to cope with pain. He was therefore asked to taper his coffee intake to 2 cups per day and change his night shifts to day shifts. His medication was switched from acetaminophen/codeine/caffeine and acetaminophen/oxycodone preparations to sustained-release oxycodone, titrated to 30 mg every 8 hours, with up to 4 acetaminophen/oxycodone tablets (325 mg/5 mg) daily for rescue analgesia. For the neuropathic component of his pain, gabapentin therapy (300 mg at bedtime) was started, but he was unable to tolerate it because of somnolence that persisted even with a low dose. Therefore, gabapentin was switched to nortriptyline, titrated from 10 to 75 mg, before bedtime. Currently his pain intensity varies between 3 and 6 out of 10. He now sleeps 8 of every 24 hours, is alert during waking hours and exercises at a

<table>
<thead>
<tr>
<th>Box 3: SISAP (Screening Instrument for Substance Abuse Potential) questionnaire</th>
</tr>
</thead>
<tbody>
<tr>
<td>• If you drink alcohol, how many drinks do you have in a typical day?</td>
</tr>
<tr>
<td>• How many drinks do you have in a typical week?</td>
</tr>
<tr>
<td>• Have you used marijuana or hashish in the past year?</td>
</tr>
<tr>
<td>• Have you ever smoked cigarettes?</td>
</tr>
<tr>
<td>• What is your age?</td>
</tr>
<tr>
<td>Increased vigilance is required by the physician prescribing opioids for men who consume more than 4 drinks a day or 16 drinks a week, women who consume more than 3 drinks a day or 12 drinks a week, patients who admit to marijuana use in the past year and patients under 40 who smoke.</td>
</tr>
</tbody>
</table>

Table 4: Sample opioid contract or agreement

I will not go to other physicians (including in emergency departments) for prescriptions for pain medication.

I will reliably attend my appointments with the doctor.

I will notify the doctor’s office if I have side effects from the prescribed medication (e.g., oversedation), and I will not drive or operate machinery while taking this medication if I feel impaired.

If a specific quantity of medication is prescribed to last me until my next scheduled appointment, I will not request earlier prescription refills.

I will accept the use of long-acting analgesia as maintenance.

I agree to other pain consultations/management strategies as necessary.

I understand that lost/stolen/spilt medications will not be replaced.

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<table>
<thead>
<tr>
<th>Patient’s signature: Date:</th>
</tr>
</thead>
<tbody>
<tr>
<td>Doctor’s signature: Date:</td>
</tr>
</tbody>
</table>

*Some patients prefer to use short-acting analgesics exclusively. They may be looking for some euphoria not available to them with sustained-release opioids.*
gym, with only gentle stretching of the shoulder. His wife reports that he is much more affable with his family and colleagues. His doses of medication have been consistent for 4 years. He is permitted to drive for work in Ontario but not in the United States because of the opioid medication.

**Case 2**

The patient was considered to have mainly nociceptive pain from intractable inflammation. His sustained-release morphine was switched to transdermal fentanyl, titrated from 125 to 200 µg/h. The change resulted in good relief from excessive constipation (such that he needed only 2 bisacodyl and 2 docusate sodium tablets daily) and improved pain control for the next 4 months. Despite the stability of his disease, his analgesia became less effective, and the opioid medication was rotated to sustained-release oxycodone, titrated from 80 to 160 mg every 12 hours. Pain control was restored for the next 2 months. He was assessed by his rheumatologist for a trial of infliximab (a new disease-modifying anti-rheumatic drug), for which the prednisone dosage needed to be titrated down to 10 mg/d; in this process, his pain was not managed even with an increase in the sustained-release oxycodone to 200 mg every 12 hours. He became somnolent and exhibited clonus, and his right elbow and left knee swelled and became more painful. He was admitted to hospital for pain management and joint aspirations, and the opioid was switched to a subcutaneous hydromorphone infusion, 5 to 8 mg/h. He experienced no pain relief, remaining somnolent with clonus. A switch to oral methadone, titrated to 15 mg every 8 hours, led to pain relief, and he regained alertness and stopped having clonus. He was discharged from hospital a few days later.

**Comment**

Opioids can be beneficial and safe as a component of the long-term management of noncancer pain. Restrictions on their use have lifted in Canada in the last decade. Physicians are acquiring increased expertise in their use, and opioids are beginning to be perceived as less invasive than recurrent surgical intervention, such as repeated spinal surgery, if they provide adequate relief at a stable dosage. Avenues to be explored include using opioid combinations to improve efficacy, devising strategies to eliminate analgesic tolerance, and developing new analgesics without undesirable opioid side effects, such as constipation, and new drugs that enhance endogenous opioid levels. As education about opioids increases and outdated prejudices decline, fewer patients with chronic noncancer pain will be denied a trial of opioid therapy when pain control is poor and causing suffering.

This article has been peer reviewed.

Dr. Gardner-Nix is with the Department of Anaesthesia, University of Toronto, the Pain Management Program, Sunnybrook and Women’s College Health Sciences Centre, and the Department of Anaesthesia, St. Michael’s Hospital Pain Clinic, Toronto, Ont.

**Correspondence to:** Dr. Jacqueline Gardner-Nix, Pain Management Program, Sunnybrook and Women’s College Health Sciences Centre, Rm. MG 630, 2075 Bayview Ave., Toronto ON M4N 3M5; fax 416 480-4772; jackie.gardner@sympatico.ca

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**References**