Levorphanol versus methadone use: safety considerations

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Abstract: Methadone has unique characteristics that make it an attractive agent for the treatment of chronic pain and opioid drug dependence. However, methadone prescription requires more clinical experience and close monitoring of patients to avoid its undesirable side effects. Recently, levorphanol has emerged as "a forgotten opioid" with a similar profile as methadone. Levorphanol has no impact on QTc prolongation and considerably less drug-drug interactions as compared to methadone. Lack of commercial availability, providers' unfamiliarity, and limited clinical data on its effectiveness remain practical issues. The objective of this article is to review and compare the safety considerations for methadone and levorphanol use.

Keywords: Levorphanol; methadone; N-methyl-D-aspartate receptors (NMDA receptors); opioids; safety

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Introduction

Levorphanol

In recent literature, levorphanol is referred to as "a forgotten opioid". Levorphanol which belongs to the phenanthrene class of opioids, was first approved and marketed under the name Levo-Dromoran in 1953 (1-3). It has high affinities for all three opioid receptors (mu, delta, kappa 1, and kappa 3), where it acts as an agonist and modulates its primary nociceptive action (4,5). Levorphanol is a strong N-methyl-D-aspartate (NMDA) receptor antagonist, and also blocks the reuptake of serotonin (5-HT) and norepinephrine (NE) therefore, can be helpful to treat neuropathic pain (5). Levorphanol has a rapid onset of action achieving peak plasma concentration by 1 hour after oral administration (6). Due to its longer half-life (11–16 hours), and duration of action (6–15 hours), levorphanol can be used as a long-acting opioid (2,6).

Levorphanol undergoes phase II metabolism-glucuronidation through UDP-glucuronosyltransferase to an inactive compound levorphanol-3-glucuronide, which is renally excreted (6). Unlike methadone, cytochrome CYP450 enzyme is not required for its metabolism, and it does not bind to P-glycoprotein in the gut (2). Levorphanol like buprenorphine, has shown a ceiling effect on respiratory depression in animal models; however, its clinical relevance needs further exploration (7). Moreover, levorphanol has no known effects on QTc prolongation (8). In short, the pharmacokinetics and metabolic profile of levorphanol indicate that, it is relatively well absorbed when taken as an oral preparation, can be used as a long-acting opioid, has fewer drug interactions and risk of respiratory depression (2,7).

Levorphanol has been studied mostly in nonmalignant neuropathic pain syndromes, and limited data is available for cancer-related pain (1,9,10). However, with its unique profile, levorphanol can be considered a safe alternative to other opioids especially methadone in conditions where chronic opioid therapy is warranted (9). In addition to its role as a first-line opioid analgesic, levorphanol may be considered in opioid rotation especially in situations where opioid-induced hyperalgesia is a concern (2).
Further research is needed to investigate its role in a palliative care setting and cancer pain management (10). Due to the underutilization of levorphanol, limited data exist on the safety and mortality risk as compared to well-studied opioids, including methadone (9). Summary of pharmacology and pharmacokinetics of levorphanol is outlined in Table 1.

**Methadone**

Methadone 6-((dimethylamino)-4,4-diphenyl-3-heptanone hydrochloride is a synthetic mu-receptor agonist, NMDA receptor antagonist, and blocks the reuptake of 5-HT and NE (11,12). Methadone was first discovered in Germany during World War II; however, its effectiveness in the treatment of opioid dependence was not recognized until the 1950s (13). As compared to levorphanol, it has a weaker NMDA receptor antagonist effect (9). Methadone has a more complex and unpredictable metabolic profile with considerable patient variability due to gene polymorphism, P-glycoprotein dependent oral absorption, and transfer across the blood-brain barrier and gastric mucosa (11,12). In contrast to levorphanol, methadone requires the CYP450 enzyme pathway for its metabolism, which can lead to a higher risk for drug-drug and drug-food interactions (11,14). Methadone has been associated with prolongation of QTc interval in several randomized and cohort studies (15-17).

Clinically, methadone has several routes available, including rectal preparation (18). Characteristics such as high potency, low cost, and excellent oral bioavailability make it an attractive opioid in the management of chronic malignant and nonmalignant pain syndromes (18). Methadone does not have any known active metabolites; therefore, it can be given in patients with compromised renal function. Unfortunately, the use of methadone has been associated with a fivefold risk of overdose deaths as compared to other opioids, which is mainly attributed to its QTc prolongation effect (19). Methadone should be prescribed cautiously in a certain high-risk population such as females, patients with congenital cardiac channel abnormalities, and patients with low potassium and low magnesium (9,20). Summary of pharmacology and pharmacokinetics of methadone is outlined in Table 1.

**Cardiovascular safety considerations**

Methadone is known to cause QTc prolongation, which can predispose patients to develop cardiac arrhythmias, torsades de pointes, and sudden death (15-17,21-23). The S-isomer of methadone is a potent inhibitor of delayed-rectifier potassium current, responsible for the arrhythmogenic activity (9). There is a dose-response relationship between the degree of QTc prolongation and methadone serum concentrations (18). Current literature does not support that methadone has any direct adverse effects on the myocardium per se (18). In one study, comorbid conditions

<table>
<thead>
<tr>
<th>Properties</th>
<th>Levorphanol</th>
<th>Methadone</th>
</tr>
</thead>
<tbody>
<tr>
<td>Chemical name</td>
<td>Levo-3-hydroxy-N-methylmorphinan</td>
<td>6-(dimethylamino)-4,4-diphenyl-3-heptanone hydrochloride</td>
</tr>
<tr>
<td>Chemical class</td>
<td>Phenanthrene</td>
<td>Diphenylheptane</td>
</tr>
<tr>
<td>Opioid receptor agonist activity</td>
<td>Mu, delta, Kappa 1, Kappa 3</td>
<td>Mu, delta, Kappa 1, Kappa 3</td>
</tr>
<tr>
<td>NMDA antagonist activity</td>
<td>Moderate</td>
<td>Weak</td>
</tr>
<tr>
<td>Serotonin and norepinephrine reuptake inhibition activity</td>
<td>Weak</td>
<td>Weak</td>
</tr>
<tr>
<td>Oral bioavailability</td>
<td>50% IM to PO dose ratio is 2:1</td>
<td>Approximately 80%</td>
</tr>
<tr>
<td>Half life</td>
<td>11–30 hours</td>
<td>15–60 hours</td>
</tr>
<tr>
<td>Duration of action</td>
<td>6–15 hours</td>
<td>4–12 hours</td>
</tr>
<tr>
<td>Metabolism</td>
<td>Phase II metabolism-glucuronidation through UDP-glucuronosyltransferase to an inactive compound levorphanol-3-glucuronide</td>
<td>Cytochrome P450 enzymes. CYP2B6, CYP2C19, CYP3A4, and CYP2D6</td>
</tr>
</tbody>
</table>

NMDA, N-methyl-D-aspartate.
such as uncontrolled blood glucose and baseline congestive heart failure were associated with higher mortality among patients who were on methadone maintenance program (MMT) (24). The Federal Drug Administration (FDA) issued a black box warning cautioning clinicians against the fatal QTc prolongation effect of methadone (9). Cochrane review on the effectiveness of electrocardiogram (EKG) screening to prevent morbidity and mortality in patients on methadone could not draw sufficient evidence to support its use. There is a consensus in obtaining a baseline EKG in certain high-risk patients, such as those with structural heart disease, arrhythmias, family or personal history of prolonged QTc, unexplained syncope, and presence of other medications that can prolong QTc (18,25,26). It is recommended to discuss benefit versus risk of methadone therapy if QTc interval is >450 but <500 ms followed by close EKG monitoring (26). In patients with QTc >500 ms, it is recommended to either discontinue or reduce the daily methadone dose (18,26). It is recommended to obtain a follow-up EKG within 2–4 weeks of methadone initiation in patients who were previously considered high risk, those with baseline EKG >450 ms, and those with a history of syncope (18). Also, obtaining an additional EKG, when daily methadone dose reaches 30–40 and 100 mg marks, and anytime when new risk factors or clinical features of arrhythmias appear is recommended (18). It is imperative to discuss the risk and benefits of EKG monitoring when methadone is considered as a first-line agent with comfort-based goals of care or as a second-line agent with curative goals of care (18). In such scenarios, patients and their families may decide not to undergo close monitoring (18).

Levorphanol has no reported effects on QTc prolongation (9). It is considered safe in patients with preexisting risk factors as described above (9). In an observational study of patients with chronic noncancer pain treated with methadone or levorphanol the response rates were 75% and 70% respectively, and levorphanol patients did not require adjuvant analgesics and had no effect on QTc prolongation (8). A comparison of cardiac safety considerations between levorphanol and methadone is outlined in Table 2.

**Hepatobiliary safety considerations**

Methadone is not known to cause an elevation in serum liver enzymes or cause acute liver injury (27). It is highly bound to alpha-1 acid glycoprotein, which is an acute-phase protein secreted by the hepatocytes; therefore, methadone distribution and serum concentration can be affected in patients with liver disease (28). Methadone metabolism is dependent on the phase 1 enzymes (CYP450), which can be impaired in liver diseases (18). Therefore, the dose of methadone should be lowered in patients with advanced liver disease such as cirrhosis (Child-Pugh Class C) (29). Moreover, time interval should be increased during dose titration (18). Methadone use should be avoided in patients with acute fulminant hepatic failure (1,18).

Levorphanol, unlike methadone, is metabolized and excreted as levorphanol-3-glucuronide, which is an inactive metabolite (9). This metabolite is excreted renally (1). There is insufficient data on the hepatic extraction and clearances, but it is generally recommended that dose interval should be increased in patients with hepatic insufficiency (1). Levorphanol can increase pressure in the common bile duct and should be avoided in biliary surgeries (1). A comparison of hepatobiliary safety considerations between levorphanol and methadone is outlined in Table 2.

**Renal safety considerations**

Methadone has been considered safe in patients with renal failure (30). Methadone has inactive metabolites, which are primarily excreted in the gut and are not dialyzable in patients undergoing hemodialysis (31). Generally, no dose adjustments have been recommended in patients with renal failure (31). Metabolites of the levorphanol, on the other hand, are renally excreted and are not dialyzable due to the high volume of distribution and increased protein binding (1). It is recommended to increase dose interval in patients with compromised renal function (1). Like methadone, levorphanol can cause urinary retention due to its anticholinergic side effects (1). A comparison of renal safety considerations between levorphanol and methadone is outlined in Table 2.

**Drug interactions**

Methadone metabolism is dependent on cytochrome P450 enzymes. The most common enzymes are CYP2B6, CYP2C19, CYP3A4, and CYP2D6 (32,33). S-methadone, which is associated with QTc prolongation, is metabolized by CYP2B6; therefore drugs, which are inducers and inhibitors of this enzyme, can affect methadone plasma concentration, metabolism, and clearance (9). Also, CYP2B6 polymorphism can result in up to 16 different allelic variants with minimal to no expression of CYP2B6,
leading to high interpatient variability (32,33). Initiating a drug that acts as an inducer or discontinuing an inhibitor of CYP3A4, can decrease methadone levels (18,26). In such cases it is recommended to monitor symptoms of increased pain or opioid withdrawal (18). Patients should be instructed to use breakthrough opioids for pain or withdrawal symptoms (18). Likewise, discontinuing a drug that acts as an inducer or initiating an inhibitor of CYP3A4, will increase methadone levels, in such cases, it is recommended to empirically reduce methadone dose by 25–50%, and monitor patients for any overdose symptoms (18). A selected list of common inducers and inhibitors of CYP3A4 is outlined in Table 3.

Contrary to methadone, levorphanol has fewer drug-drug interactions since it is not dependent on CYP450 enzymes (1). Generally, drugs that inhibit glucuronidation such as tricyclic antidepressants, phenothiazines, and ranitidine can potentiate the effects of levorphanol (1). In contrast, drugs that induce glucuronidation such as carbamazepine, phenobarbital, phenytoin, and rifampin can decrease its effects (1). Concurrent use of monoamine oxidase inhibitors and levorphanol is not recommended (1). A comparison of drug interactions between levorphanol and methadone is outlined in Table 2.
Safety considerations in sleep disorders

Methadone worsens central and obstructive sleep apnea in a dose-dependent fashion (34-36). Methadone use has been associated with diminished respiratory response to PCO₂, widens the alveolar-arterial oxygen gradient through hypoventilation, and directly increases apnea-hypopnea in a dose-dependent manner (34). There are no guidelines available on screening and monitoring of sleep disorders while on methadone (18). Clinicians should consider alternative options in patients with preexisting sleep apnea (18). There is no data available regarding the interactions of levorphanol and sleep-disordered breathing. A comparison of safety considerations in sleep disorders between levorphanol and methadone is outlined in Table 2.

Mortality risk

Between 1999–2010, methadone-associated deaths have disproportionately increased by 600% as compared to an increase in deaths due to other opioids of 138% (19). When used as a first-line opioid in the treatment of chronic pain among hospitalized patients, the out of hospital mortality increased to 46% during the follow-up period (37). In the majority of the published data on methadone-associated death, it is difficult to differentiate the cause of death from respiratory depression and fatal arrhythmias (38). Nevertheless; methadone safety concerns are widely accepted due to its unique pharmacology (18, 26, 39). Recently, various organizations have published consensus guidelines to promote safer use of methadone (18, 26, 39). In recent years, methadone-associated deaths have declined, which might be associated with increased awareness and safe prescription patterns (40).

Drug availability and cost consideration

In the United States, methadone is manufactured by Ascent Pharmaceuticals INC (41). It is available in 5 and 10 mg oral preparations (42). Methadone is widely available and is covered by Medicare and most of the insurance plans (42). The average retail price of 5 mg oral tablet (#90) is $31.28 (42).

Levorphanol is manufactured by Sentynl Therapeutics INC, and Virtus Pharmaceuticals (41). It is available in 1, 2 and 3 mg oral preparations (43). In our experience, levorphanol is not readily available in the majority of the pharmacies and may have regional differences (10). The average retail price of 2 mg oral tablet (#90) is $4,490.42 (43).

Conclusions

Levorphanol has a more predictable pharmacokinetic profile, with a shorter half-life and prolonged duration of action as compared to methadone. Unlike methadone, it undergoes glucuronidation to an inactive metabolite, which is renally excreted. Levorphanol has no QTc prolongation risk, fewer drug-drug interactions, and like buprenorphine may have a ceiling effect on respiratory depression. Despite the safer profile of levorphanol, the limited knowledge regarding its use in the palliative care setting, lack of easy availability along with the high cost of the drug may
make prescribing levorphanol a challenge as compared to methadone. Further research is needed to investigate the role of levorphanol in the setting of palliative care and cancer pain which may subsequently make the drug more accessible and affordable for our patients.

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Footnote

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Ethical Statement: The authors are accountable for all aspects of the work in ensuring that questions related to the accuracy or integrity of any part of the work are appropriately investigated and resolved.

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