Switching from Methadone to a Different Opioid: What Is the Equianalgesic Dose Ratio?


Abstract

Introduction: Methadone (ME) is a highly effective opioid agonist used for difficult pain syndromes. However, in the management of cancer pain with strong opioids, rotation to a different opioid (opioid rotation) may be required because of side effects or poor pain control. Rotation from methadone to another opioid has received limited study and therefore may be difficult because of the absence of a uniformly accepted dose conversion ratio.

Methods: Retrospectively reviewed consecutive medical records of patients undergoing an opioid rotation from methadone to an alternative opioid were evaluated. For inclusion, patients were required to have received methadone for at least 3 days and have reached stable dose of the alternative opioid(s) during the 7 days following. Stable dose was defined as a 30% or less change in opioid dose from one day to the next.

Results: Records of 39 patients met inclusion criteria. Excluded from analysis were 5 patients who were restarted on methadone within 7 days, 2 with irregular opioid use resulting in negligible regular opioid doses post-switch, and 3 due to concerns about reliability of multiple routes used for fentanyl. Data from 29 patients, 10 female, mean age 48±14.4 years, were evaluable. The mean dose ratio for oral methadone to oral morphine equivalent daily dose (MEDD) was 1:4.7 (95% confidence interval [CI], 3.0–6.5; n = 16), and for intravenous (IV) methadone to MEDD was 1:13.5 (95% CI, 6.6–20.5; n = 13), p = 0.06. Methadone dose was significantly correlated to stable MEDD after switching opioids for both methadone IV and oral (Spearman = 0.86, p = 0.0001 and Spearman = 0.72, p = 0.0024), respectively. Mean day of achieving stable dose was day 2.5±0.2 for IV methadone and day 2.6±0.3 for oral methadone.

Conclusion: These dose ratios are new findings that may assist in switching patients more safely to alternative opioids when side effects or pain problems occur when patients are receiving methadone. An important difference in analgesic potency appears to exist between IV and oral ME. Future research with prospective studies is required.

Introduction

Methadone is a synthetic opioid agonist that is used increasingly in the management of cancer pain.1,2 The strategy of opioid rotation, that is switching from one opioid drug to another, is substantiated by the medical literature and is commonly used in clinical practice to manage opioid side effects or inadequate analgesia.3–5 The use of equianalgesic dose ratios provides a method to determine equivalent and safe analgesic doses for opioids that differ in potency.2,6 Many studies have addressed the equianalgesic dose ratios for switching between morphine and hydromorphone, morphine and oxycodone, and when switching from morphine to methadone.7–10 Only two small studies report on rotations involving methadone in the opposite direction, that is when switching from methadone to an alternate opioid.1,8 The direction of opioid rotation is important, as the conversion factors that result are not necessarily equivalent when switching the opioid in the opposite direction. One study by Lawlor et al.8 reports on six patients switched from methadone to morphine. The study by Moryl et al.1 reports that 12 of 13 patients were unable to complete a rotation from methadone to another opioid due to pain and adverse side effects. This is contrary to the experience of our group as we enjoyed successful pain control in switching patients from methadone to an alternative opioid.
Methods

Institutional Review Board (IRB) approval was obtained to perform a retrospective analysis of consecutive opioid rotations from methadone to an alternative opioid performed by the Department of Symptom Control and Palliative Care (eight board-certified palliative medicine specialists with significant expertise in the use of methadone and opioid analgesia supported by fellows and advance practice nurses that evaluate patients daily). Using the computerized pharmacy database, medical records were screened for inclusions and exclusion criteria. Opioid rotation was performed empirically based on the experience of most physicians using a methadone: morphine conversion dose ratio of 1:5 or 1:10 as a benchmark. Standard practice had doses adjusted up or down by a minimum of 25%–30% until pain was controlled or side effects resolved. Breakthrough pain doses of 10% of the total daily opioid dose (or equivalent) were typically pre-

\[
\text{METHODS - MEDD}_1 \quad \text{MEDD}_2 \quad \text{MEDD}_3
\]

\[
\text{MEDD}_3 = \text{Conversion dose ratio} \\
\text{Methadone}
\]

**FIG. 1.** Calculation of conversion ratio.

**FIG. 2.** Study outline.
SWITCHING FROM METHADONE TO A DIFFERENT OPIOID

Inclusion and exclusion criteria were determined by utilizing criteria from previous studies in order to obtain reasonable estimates of stable methadone dose prior to switch and to eliminate possible confounding factors such as delirium, discharge before stable dose, or death.

Patients were included if: they had cancer pain and were rotated from methadone to another strong opioid as an inpatient, had received methadone 3 days or more before the rotation, had reached stable dose of the new opioid 2 days or more before discharge or death, reached stable dose 1–7 days post-switch, and methadone was able to be completely stopped at time of rotation. Stable dose was defined as the dose maintained with 30% or less change for more than 24 hours.

Patients were excluded if: they were discharged from the hospital less than 48 hours after the opioid switch, were receiving opioids via intrathecal or epidural pump, if opioid rotation involved more than 3 strong opioids other than methadone, or if they were receiving sedation with midazolam.

Data collection included age, gender, race, primary tumor, pain score, reason for opioid switch, and date of death if available.

The dose of all opioids other than methadone were converted to morphine equivalent daily dose (MEDD) for purposes of analysis, using established conversion ratios for these opioids (these ratios are summarized at www.palliative.org/PC/ClinicalInfo/AssessmentTools, using parenteral MEDD). For patients receiving methadone and a second opioid prior to the switch, the MEDD of the second opioid was subtracted from the MEDD calculated for the day when stable dose was reached. The remainder was used to calculate the equianalgesic dose ratio with the previous methadone dose (Fig. 1).

Summary statistics were generated to describe the demographic and clinical characteristics of the patients. Scatter plots of methadone dose on the day prior to switch versus stable MEDD dose were obtained. Regression modes with no Y-intercept were fit to predict the new opioid dose based on the observed methadone dose. Equianalgesic dose ratios were calculated. Statistical analysis of the relationship be-

<table>
<thead>
<tr>
<th>Variable</th>
<th>IV methadone to oral MEDD n(%), n = 13</th>
<th>Oral methadone to oral MEDD n (%), n = 16</th>
<th>Total n (%), n = 29</th>
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<tr>
<td>Gender</td>
<td>Female 5 (38%)</td>
<td>5 (31%)</td>
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<td>White 8 (62%)</td>
<td>11 (69%)</td>
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<td>Other 5 (38%)</td>
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<td>Age</td>
<td>Mean (SD) 49 (12)</td>
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<td>48 (14)</td>
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<td>3 (19%)</td>
<td>6 (21%)</td>
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<td></td>
<td>Opioid toxicity 3 (23%)</td>
<td>5 (31%)</td>
<td>8 (28%)</td>
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<td>Other 3 (23%)</td>
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<td>Unknown 4 (31%)</td>
<td>1 (6%)</td>
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<td>Methadone Dose</td>
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<td>5 (17%)</td>
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<td>10–19 mg/d 1 (8%)</td>
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<td>6 (21%)</td>
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<tr>
<td></td>
<td>20–34 mg/d 2 (15%)</td>
<td>4 (25%)</td>
<td>6 (21%)</td>
</tr>
<tr>
<td></td>
<td>35–57 mg/d 4 (31%)</td>
<td>3 (19%)</td>
<td>7 (24%)</td>
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<td></td>
<td>&gt;57 mg/d 2 (15%)</td>
<td>3 (19%)</td>
<td>5 (17%)</td>
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<tr>
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<td>4 (25)</td>
<td>8 (28)</td>
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<tr>
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<td>Gynecological 2 (15)</td>
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</tr>
<tr>
<td></td>
<td>Other 1 (8)</td>
<td>5 (31)</td>
<td>6 (21)</td>
</tr>
</tbody>
</table>

IV, intravenous; MEDD, morphine equivalent daily dose; SD, standard deviation.

FIG. 3. Linear regression: intravenous methadone to oral morphine equivalent daily dose (MEDD).
between dose ratio and methadone dose was performed. Linear, quadratic, and cubic models were fit to the data. Spearman’s correlation coefficient was calculated to measure the strength of the relationship between the previous methadone dose and the stable opioid dose.

Results

The computerized pharmacy database determined 265 opioid rotations from methadone to other opioids. These were further screened for inclusions/exclusion criteria (Fig. 2).

Data from the 29 remaining patients were classified into two distinct subgroups: IV methadone switched to oral MEDD and oral methadone switched to oral MEDD (Table 1). Among the IV and oral methadone groups approximately equal numbers were switched to morphine (n = 6, n = 7, respectively) and hydromorphone (n = 5, n = 5). The remaining patients received oxycodone (n = 1, n = 2 respectively), a combination of morphine and hydromorphone (n = 1, n = 1), and a combination of morphine, hydromorphone and oxycodone (n = 1, oral methadone group). The majority of the cases represented the use of methadone plus other opioid(s) prior to the switch, who were then rotated to morphine (n = 13) or hydromorphone (n = 10).

Plots of the linear regression for both IV methadone and oral methadone are displayed in Figures 3 and 4, from these are derived the estimated equianalgesic conversion dose ratios (Table 2). Analysis of the relationship between dose ratio and methadone dose did not yield a statistically significantly relationship for any of the models used.

Methadone dose was significantly correlated to stable MEDD after switching opioids for both: methadone IV (Spearman = 0.86, p = 0.0001) and ME oral (Spearman = 0.72, p = 0.0024). The mean day of achieving stable dose was on day 2.5 ± 0.2 for intravenous ME, and on day 2.6 ± 0.3 for oral methadone.

Prior to the switch a pain score was documented in the records on only 12 of 29 patients (mean 3.9 ± 2.4 on 0–10 verbal rating scale): At the time of stable dose a pain score was available for only 8 of 29 patients (mean 3.9 ± 1.6 on 0–10 verbal rating scale).

Discussion

The term “equianalgesic dose ratio” refers to the ratio of the dose of two opioids required to produce the same analgesic effect.10 Equianalgesic dose ratios and “relative potency ratios” can cause confusion and are dangerous as they are inverse mathematical expressions to reflect the same relationships.10 In this study the terminology to indicate the direction of the opioid rotation (from methadone to other opioids) is used consistently as this is the novel feature of this study. To clarify and avoid dangerous misinterpretation, we have found intravenous methadone and oral methadone to be 13.5 and 4.7 times more potent than an equivalent oral MEDD of a strong opioid. Of interest we find a difference between the intravenous and oral methadone conversion dose ratios that obtained borderline statistical significance by p value, likely due to low numbers, and reached statistical significance based on confidence intervals. This was not expected based on methadone’s high oral bioavailability. This appears to be clinically important and likely due to presystemic (first pass) elimination.

These results run contrary to those of Moryl et al.1 who reported only one successful rotation from intravenous

![FIG. 4. Linear regression: oral methadone to oral morphine equivalent daily dose (MEDD).](image)

Table 2. Estimated Conversion Dose Ratios and Mean Dosages

<table>
<thead>
<tr>
<th>Variable</th>
<th>IV methadone oral MEDD</th>
<th>Oral methadone to oral MEDD</th>
</tr>
</thead>
<tbody>
<tr>
<td>Methadone dose</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mean mg/d (SD)</td>
<td>30 (25)</td>
<td>35 (31)</td>
</tr>
<tr>
<td>Median mg/d (range)</td>
<td>20 (2–75)</td>
<td>26 (5–105)</td>
</tr>
<tr>
<td>Stable MEDD dose</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mean mg/d (SD)</td>
<td>411 (552)</td>
<td>191 (192)</td>
</tr>
<tr>
<td>Median mg/d (range)</td>
<td>275 (43–2138)</td>
<td>110 (15–610)</td>
</tr>
<tr>
<td>Estimated conversion dose ratio</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mean (95% CI)</td>
<td>13.5 (6.6–20.5)</td>
<td>4.7 (3.0–6.5), p = 0.06a</td>
</tr>
<tr>
<td>Median (range)</td>
<td>15.0 (3.7–37.5)</td>
<td>4.7 (0.5–15.3)</td>
</tr>
</tbody>
</table>

*p value reflects the difference between mean IV and oral methadone conversion dose ratios.

IV, intravenous; MEDD, morphine equivalent daily dose; SD, standard deviation; CI, confidence interval.
methadone to intravenous hydromorphone and 12 patients who were unsuccessful in switching to intravenous morphine, hydromorphone, fentanyl, and levorphanol from methadone. These patients were restarted on methadone. In contrast we have found that of 39 rotations from methadone that met inclusion criteria, only 5 (13%) needed to be restarted on methadone. Also, of our successful rotations the mean time to achieving stable dose was 2.5–2.6 days, indicating rapid stabilization of opioid requirements. It is unclear what may be responsible for the difference in findings between our report and those of Moryl et al. One important difference may have been the higher doses of IV methadone administered to the patients in the Moryl et al. study (2–80 mg/hr intravenous) compared to lower doses of intravenous and oral methadone administered to our patients prior to opioid rotation. This may explain the significant opioid induced neurotoxicity they report (dysphoria, confusion, sedation, myoclonus) as well as persisting pain which may possibly reflect the presence of hyperalgesia or allodynia.

Lawlor et al. has reported a subanalysis of 6 patients rotated from oral methadone to oral morphine. The reported median dose ratio was 8.25 with an interquartile range of 4.37–11.3. Our findings in 16 patients report a mean ratio of 4.7 (95% CI 3.0–6.5) and median ratio of 4.7 (range, 0.5–15.3; Table 2).

We were interested in determining if the conversion dose ratio would change based on methadone dose, as has been reported by Ripamonti et al. in a study of patients rotated in the opposite manner, i.e. from morphine to oral methadone. Statistical analysis of the relationship between dose ratio and methadone dose was performed. Linear, quadratic, and cubic models were fit to the data. None of the models yielded a statistically significant relationship. Therefore, unlike Ripamonti et al. who were able to conclude that there is an increasing dose ratio based on increasing doses of morphine when switching from morphine to methadone, we are unable to conclude that there exists a mathematical relationship that varies the dose ratio based on the dose of methadone, when rotating from methadone to another opioid.

To our knowledge this is the most extensive study of patients rotated from methadone to other opioids, an area of limited research but of increasing importance. Chronic, not single dose opioid use was investigated among a broad sampling of the cancer population. Other strengths include a rigorous inclusion and exclusion criteria to minimize confounders and inclusion of data for both intravenous and oral methadone use. The use of opioid consumption rather than analgesic reporting has been established as an alternate viable method of establishing equianalgesic dose ratios. Multiple studies support the validity of the MEED conversion ratios used in the methods section this study and in clinical practice.

Limitations of this study include: its retrospective nature, relatively low numbers, predominance of white males, lack of control for hepatic and renal impairment, lack of direct drug to drug dose comparison, no control for radiation, chemotherapy, hormonal therapy or other interventions that may influence pain control, and limited data available related to pain scores. A further limitation relates to methadone’s long half-life. In this regard a design utilizing a longer period of methadone administration before rotation, and reporting of dosages of other opioids at longer follow-up may be beneficial in more accurately calculating the equianalgesic dose ratios.

Although the authors believe this study improves our knowledge of the relationship between analgesic requirements that occurs when switching from methadone to an alternative opioid, we urge caution in the use of these conversion dose ratios due to the limitations listed above. Further studies utilizing prospective design and larger numbers are required to confirm these results. We suggest that until further studies are completed, it would be judicious to rotate patients in clinical practice from methadone to other opioids using conversion dose ratios more conservative than described in this paper. It would be safer to underestimate the analgesic effect of methadone and use breakthrough opioid doses to “catch up,” than risk overdose with the new opioid due to a possible overestimation of methadone’s analgesic potency.

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Author Disclosure Statement

No competing financial interests exist.

References


