Is a single low dose of intrathecal morphine a useful adjunct to patient-controlled analgesia for postoperative pain control following lumbar spine surgery? A preliminary report

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BACKGROUND: Several studies addressing intrathecal morphine (ITM) use following spine surgery have been published either involving the pediatric population, using mid- to high-dose ITM, or not in conjunction with morphine patient-controlled analgesia (PCA).

OBJECTIVES: To determine whether low-dose ITM is a useful adjunct to PCA for postoperative pain control following elective lumbar spine surgery in adults.

METHODS: Thirty-two patients were enrolled in a double-blinded randomized controlled trial, and received either ITM or intrathecal placebo. Postoperatively, all patients were given a PCA pump and observed for the first 24 h in a step-down unit. Measurements of: total PCA morphine consumed in the first 24 h; intensity of pain; pruritus; nausea at 4 h, 8 h and 24 h; time to first ambulation; length of hospital stay; and occurrences of respiratory depression were recorded.

RESULTS: The total PCA use was significantly lower in the ITM group. There were lower average pain scores in the ITM group, which increased to that of the intrathecal placebo group over 24 h; however, this failed to attain statistical significance. There were no differences in nausea, pruritus, time to first ambulation or hospital length stay. There were no cases of respiratory depression in either group.

CONCLUSIONS: ITM may be a useful adjunct to PCA, but did not decrease time to ambulation or length of stay.

Key Words: Intrathecal morphine; Lumbar spine surgery

Une seule faible dose de morphine par voie intrathécale est-elle un ajout utile à l’analgésie contrôlée par le patient pour contrôler la douleur après une opération de la colonne lombaire? Un rapport préliminaire

HISTORIQUE : Plusieurs études portent sur l’utilisation de morphine par voie intrathécale (MIT) après une opération de la colonne vertébrale auprès de la population pédiatrique, à l’aide de doses modérées à élevées de MIT ou auprès de la population adulte conjointement à une analgésie contrôlée par le patient (ACP) au moyen de morphine.

OBJECTIFS : Déterminer si une faible dose de MIT est un ajout utile à l’ACP pour contrôler la douleur postopératoire après une opération non urgente de la colonne lombaire chez des adultes.

MÉTHODOLOGIE : Trente-deux patients ont participé à un essai aléatoire dans un centre de soins courants. Les chercheurs ont collecté les mesures totales d’ACP au moyen de morphine consommées pendant les 24 premières heures, l’intensité de la douleur, le prurit, les nausées au bout de quatre, huit et 24 heures, le délai jusqu’à l’ambulation, la durée d’hospitalisation et les occurrences de dépression respiratoire.

RÉSULTATS : L’utilisation totale d’ACP était considérablement plus faible dans le groupe prenant de la MIT. Les scores de douleur moyens étaient moins élevés dans le groupe prenant de la MIT, mais sur 24 heures, ils correspondaient à ceux du groupe prenant un placebo par voie intrathécale. Ce résultat n’avait toutefois pas de signification statistique. Il n’y avait pas de différence sur le plan des nausées, du prurit, du délai jusqu’à l’ambulation ou de la durée d’hospitalisation. Aucun des groupes n’incluait de cas de dépression respiratoire.

CONCLUSIONS : La MIT peut être un ajout utile à l’ACP, mais elle ne réduit pas le délai avant l’ambulation ni la durée d’hospitalisation.

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The primary aim of the present study was to determine whether administration of low-dose ITM is a useful adjunct to PCA for providing postoperative pain control following elective lumbar spine surgery in the adult population. The secondary goals were to determine whether ITM affected hospital length of stay, time to ambulation and occurrences of respiratory depression.

METHODS

Patients undergoing elective lumbar laminectomy performed by a single surgeon, with or without fusion, and with or without instrumentation, were considered for entry into the study. Inclusion criteria were signed informed consent and age ≥16 years and <70 years. Exclusion criteria were pregnancy, allergy/intolerance to any of the study medications, chronic morphine use, history of sleep apnea and inability to use PCA.

Intraoperatively, patients all received intravenous fentanyl 1 µg/kg to 3 µg/kg, intravenous sodium thiopental 3 mg/kg to 5 mg/kg and neuromuscular relaxant, at the discretion of the anesthesiologist; per rectum acetylmorphine 1300 mg postinduction; maintenance of anesthesia with 60% to 70% nitrous oxide and isoflurane at 0.02% to 1.0%; neuromuscular blockade at the discretion of the attending anesthesiologist; intravenous ondansetron 4 mg before emergence and reversal of neuromuscular blockade at the discretion of the management anesthesiologist, who was blinded to patient allocation, with the proviso that background infusions were not to be used.

All patients were monitored for the first 18 h to 24 h postoperatively in a step-down unit with continuous oxygen saturation monitoring. The postoperative outcome measures that were evaluated and recorded were: pain intensity, graded by the patient from 1 to 10; total PCA morphine used in the first 24 h postoperatively, measured in milligrams; time to first ambulation, measured in minutes; length of hospital stay, measured in days; nausea intensity, graded by the patient from 1 to 4; pruritis intensity, graded by the patient from 1 to 4; and episodes of respiratory depression (rate < 9 breaths/min).

Statistical analysis was performed using a series of independent-samples t tests to compare the two groups with regard to all of the various outcomes. With this sample size, P<0.05 was considered to be statistically significant. While there are three temporal measures for each patient, repeated-measures ANOVA was not performed because within-person change over time was not of interest; rather, the between-group differences at each point in time were the focus of the analysis. No a priori adjustments were made for multiple comparisons; rather, all P values and sample sizes are presented to enable the reader to interpret the data in the context of the comparisons.

RESULTS

The ITM group included 18 patients and the ITP group included 14 patients. The groups were not significantly different according to patient age, sex, American Society of Anesthesiologists classification and whether a fusion was performed in addition to the lumbar decompression (Table 1).

The average total PCA use over the first postoperative 24 h was significantly less in the ITM group (32.7 mg versus 59.4 mg; P=0.04) (Table 2). There were lower average pain scores in the ITM group at 4 h (P=0.11) and at 8 h (P=0.07) postoperatively; however, this did not attain statistical significance. There was no difference at 24 h postoperatively (P=0.90) (Figure 1). There were no episodes of respiratory depression experienced by patients in either group.

There was no difference between the groups with regard to nausea at 4 h (P=0.14), 8 h (P=0.23) and 24 h (P=0.56) postoperatively; pruritis at 4 h (P=0.15), 8 h (P=0.32) and 24 h (P=0.91) postoperatively; time to first ambulation (P=0.81) and hospital length of stay (P=0.14) (Table 2).

Sample size calculations were derived on the basis of the cumulative patient-controlled morphine delivered in the first 24 h, as reported by France et al (19). SDs were not provided but are estimated on the basis of the magnitude in the current study. Using a mean ± SD of

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**Table 1** Demographic characteristics of participants

| Characteristic | ITM | ITP | P* |
|---------------|-----|-----|----|
| Sex, female/male, n/n | 7/11 | 3/11 | 0.29 |
| Age, years, mean | 54.6 | 54.8 | 0.96 |
| American Society of Anesthesiologists classification, mean† | 2.1 | 2.3 | 0.69 |
| Fusion, n | 6 | 3 | 0.46 |

*Calculated using χ² tests or two-tailed t tests as appropriate; †Range 1–3. ITM Intrathecal morphine; ITP Intrathecal placebo

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**Table 2** Independent-samples tests for group statistics

| Group | Time postoperatively, h | Intrathecal morphine | Intrathecal placebo | P* |
|-------|-------------------------|----------------------|---------------------|----|
| Total 24 h patient-controlled analgesia, mg | | | | |
| Nausea intensity† | 4 | 1.4 | 1.1 | 0.14 |
| 8 | 1.3 | 1.1 | 0.23 |
| 24 | 1.4 | 1.2 | 0.56 |
| Pruritus intensity† | 4 | 1.5 | 1.1 | 0.15 |
| 8 | 1.7 | 1.4 | 0.32 |
| 24 | 1.5 | 1.5 | 0.91 |
| Time to ambulation, min | | | | |
| Hospital length of stay, days | 1853 | 1705 | 0.81 |

Data presented as mean unless otherwise indicated. †On a scale of 1 to 4
placebo groups.

Figure 1) Pain scores in the intrathecal morphine versus intrathecal placebo groups.

7±5.2 and a mean of 25.5±19.1, an alpha of 0.05 and a two-tailed test, the study would have a power of 80% with only 10 patients in each group; this suggests that the current number of 18 and 14 patients would provide adequate power to detect difference in morphine delivery.

The study by France et al (19) noted few differences in the complication rates. Itchiness was noted in eight experimental and two control patients; need for nalaxone due to a respiratory rate <8 breaths/min was demonstrated in three experimental patients and none of the control patients. Neither attained statistical significance. Using an alpha of 0.05 and a power of 80%, 130 patients per group would be required for this to attain statistical significance, suggesting that the current study, as well as the study by France et al (19), were both underpowered to detect these complications. However, sample size calculations are not as relevant in the case of relatively rare events and, given that the use of morphine was the primary outcome, the present study had sufficient power for this outcome.

DISCUSSION

Similar to France et al (19), we found that the average total PCA use over the first postoperative 24 h was significantly lower in the ITM group, although we used a low dose rather than the mid-range dose that France et al (19) used. Therefore, low-dose ITM is an effective early adjunct to PCA in decreasing the amount of self-administered systemic morphine in patients undergoing lumbar spine surgery. The average pain scores in the ITM group were lower for the first 8 h postoperatively; however, this did not attain statistical significance.

There were no episodes of respiratory depression experienced by patients in the ITM with PCA group. Therefore, although the present study had a small sample size, our data support clinicians who believe that respiratory depression is dose dependent and advocate using low-dose ITM.

There were no differences in measured nausea or pruritus between the ITM and ITP groups. Interestingly, therefore, decreasing the amount of systemic morphine using ITM did not result in less nausea or pruritis in our patients.

We found no difference in time to ambulation or length of stay with the use of ITM. We believe this may be because time to ambulation and length of stay after spine surgery is determined not by postoperative pain but by multiple factors such as preoperative deconditioning, postoperative fatigability and the patients' home support system. Therefore, in the present study, ITM was not a useful adjunct for decreasing hospital resource utilization in lumbar spine surgery.

The strength of our study was in its design methodology. The weakness is the small sample size, which may have reduced our ability to find a statistically significant difference in pain scores in favour of ITM.

CONCLUSION

ITM was an effective adjunct to PCA for decreasing the amount of self-administered systemic morphine required by patients undergoing lumbar spine surgery. Further studies, including larger sample sizes, are needed to show that a low dose provides superior early postoperative pain relief compared with PCA alone, with less respiratory depression than mid-range doses of ITM.

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