Purpose: The efficacy of low dose intrathecal lidocaine-sufentanil was compared with intrathecal sufentanil for short duration outpatient gynecological laparoscopy.

Methods: Thirteen ASA I and II patients undergoing gynecological laparoscopy were studied in a randomized double-blind trial. Patients received either intrathecal 10 mg lidocaine plus 10 µg sufentanil (Group LS) or intrathecal 20 µg sufentanil (Group S), each diluted to 3 mL with sterile water through a 27g Whitacre needle in the sitting position. Sensory and motor recovery were assessed with pinprick and a modified Bromage scale.

Results: One of seven Group LS patients and two of five Group S patients required conversion to general anesthesia for failed skin test with forceps. Two of the remaining three Group S patients felt sharpness with skin incision. The study was terminated early because of inadequate anesthesia in Group S. The small sample size (n=9) made statistical analysis uninformative.

Conclusion: Intrathecal 20 µg sufentanil is unsuitable as a sole agent for gynecological laparoscopy.

Objectif: Comparer l’efficacité de l’injection intrathécale d’une faible dose de lidocaïne-sufentanil au sufentanil seul dans le contexte d’une laparoscopie gynécologique ambulatoire de courte durée.

Méthode: L’essai randomisé et à double insu a porté sur 13 patientes, ASA I et II, qui devaient subir une laparoscopie gynécologique. On a procédé à l’injection intrathécale, soit 10 mg de lidocaïne plus 10 µg de sufentanil (groupe LS), soit 20 µg de sufentanil (groupe S), complétés dans chaque cas au volume de 3 mL avec de l’eau stérile et administrés, en position assise, à l’aide d’une aiguille Whitacre de calibre 27. La récupération sensitive et motrice a été évaluée par la réaction à la piqûre et par une échelle de Bromage modifiée.


Conclusion: L’administration intrathécale de 20 µg de sufentanil seul est inadéquate pour la laparoscopie gynécologique.

Selective spinal anesthesia for outpatient laparoscopy.
III: Sufentanil vs lidocaine-sufentanil

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ALTHOUGH short duration outpatient gynecological laparoscopy is usually performed on patients under general anesthesia, there are recent publications describing effective anesthesia for laparoscopy using small-dose hypobaric lidocaine-fentanyl spinal anesthesia. While this technique results in less motor block, faster sensory recovery and earlier discharge compared with conventional dose spinal anesthesia, doses of lidocaine as low as 10 mg still produce mild motor weakness of the lower extremities in some patients. In addition, complete return of sensation to pinprick remains the rate-limiting step to early discharge. Intrathecal sufentanil (ITS) as a sole agent has been used successfully for analgesia during labour and delivery, and extracorporeal shock wave lithotripsy, while preserving motor function of the lower extremities.

We hypothesized that ITS alone would provide adequate surgical conditions for short duration outpatient gynecological laparoscopy, while achieving the goal of early discharge with fully intact sensory and motor function. In the present study, intrathecal sufentanil was compared with small-dose hypobaric lidocaine-sufentanil spinal anesthesia for laparoscopy with respect to operating conditions, recovery of function, and side effects.

Methods

The study was conducted as a single-centre, randomized, prospective double-blind trial. Institutional and university ethics approvals were obtained along with written informed consent from each patient recruited. Exclusion criteria were neurological or neuromuscular disease, infection at the intended site of needle insertion for spinal anesthesia, coagulopathy, and allergy to lidocaine or sufentanil.

Thirteen ASA I and II women scheduled to undergo outpatient gynecological laparoscopic procedures of less than 60 min duration were enrolled in the study and randomized to two groups. Group LS received a hypobaric solution of 10 mg lidocaine 1% with 10 µg sufentanil diluted with sterile water to a total volume of 3 ml. Group S received a hypobaric solution of 20 µg sufentanil, mixed in sterile water to a final volume of 3 mL. Solutions were prepared aseptically by an anesthesiologist not performing the anesthetic and both solutions appeared identical.

An intravenous infusion of sodium chloride 0.9% was established preoperatively. In the operating room, patients were monitored with an ECG, pulse oximeter, and automatic blood pressure cuff. Spinal anesthesia was administered in the sitting position with a midline approach at the L3-4 level using a 27g Whitacre spinal needle through an introducer. The solution was injected, after aspiration of cerebrospinal fluid, as rapidly as possible in a single shot with the bevel cephalad. After sitting for one minute, the patient was placed in 20 to 30° reverse Trendelenberg while lithotomy positioning and skin preparation occurred.

The extent of motor blockade and sensory loss to pinprick were measured at three and five minutes post spinal injection, prior to the surgical start. Testing the skin with toothed forceps prior to surgery was instituted after the third subject (Group S). She was the first patient to note sharpness during trocar insertion, in spite of blunted sensation to pinprick testing at three and five minutes. Patients were placed in Trendelenberg during carbon dioxide (CO₂) insufflation into the peritoneum to minimize diaphragmatic irritation from CO₂ and consequent shoulder tip pain.

Intraoperative patient comfort was continually evaluated by verbal communication, and discomfort was graded by the patient as mild, moderate, or severe. In the presence of an obviously inadequate block (defined as sharp sensation when testing the abdomen with toothed forceps) a GA was induced. Anxiety and abdominal or shoulder discomfort were treated with increments of 1 mg midazolam and 250 µg alfentanil iv, respectively. Surgical conditions were graded by the surgeon as poor, fair, good, or excellent.

Assessment in the recovery room consisted of sensory and motor block measurements and evaluation of pruritus on arrival and every 30 min. Sensation to pinprick was assessed with a 20g needle until the S₁ dermatome at the lateral aspect of the foot normalized. Motor block was assessed with a modified Bromage scale until the patient was able to perform a deep knee bend independently. Pruritus was evaluated using a sliding Visual Analogue Scale anchored at 0 (no pruritus) and 10 (maximum pruritus), and evaluations were discontinued when pruritus became absent or the patient was discharged. Patients reporting pruritus were offered treatment with 40 µg naloxone iv and reassessed after 15 min when the dose was repeated if necessary. Nausea and abdominal pain were graded according to the most severe manifestation over the duration of recovery, using a modified Prince Henry Pain Scale. A measurement of satisfaction was obtained just before discharge by asking the patient: “If you had to have the surgery again would you choose the same spinal anesthetic?”

Results

Thirteen women age 32–72 yr were enrolled. One was excluded after she developed abrupt onset of paresthesia coincident with the beginning of subarachnoid
TABLE I Intraoperative characteristics of small-dose spinal lidocaine-sufentanil (Group LS) vs sufentanil (Group S)

<table>
<thead>
<tr>
<th>Sensory level at 3'</th>
<th>Motor block†</th>
<th>Surgical conditions‡</th>
<th>Supplements</th>
<th>Complications§</th>
<th>Abdominal discomfort</th>
</tr>
</thead>
<tbody>
<tr>
<td>Patient 1</td>
<td>T10</td>
<td>T5</td>
<td>4</td>
<td>excellent</td>
<td>0</td>
</tr>
<tr>
<td>Patient 2</td>
<td>T11</td>
<td>T6</td>
<td>4</td>
<td>good</td>
<td>0</td>
</tr>
<tr>
<td>Patient 3</td>
<td>T6</td>
<td>T4</td>
<td>4</td>
<td>fair</td>
<td>1</td>
</tr>
<tr>
<td>Patient 4</td>
<td>T10</td>
<td>T9</td>
<td>4</td>
<td>good</td>
<td>0</td>
</tr>
<tr>
<td>Patient 5</td>
<td>T9</td>
<td>T7</td>
<td>4</td>
<td>good</td>
<td>0</td>
</tr>
<tr>
<td>Patient 6</td>
<td>T6</td>
<td>T6</td>
<td>4</td>
<td>good</td>
<td>0</td>
</tr>
<tr>
<td>Mean</td>
<td>T8</td>
<td>T5</td>
<td>4</td>
<td>GA</td>
<td>GA</td>
</tr>
</tbody>
</table>

Group S

| Patient 1 | T6 | T4 | 4 | good | 1 | 500 | 0 | 0 | mild | mild |
| Patient 2 | T9 | T8 | 4 | GA | GA | GA | GA | GA | GA | GA |
| Patient 3 | T9 | T8 | 4 | GA | GA | GA | GA | GA | GA | GA |
| Patient 4 | T6 | T4 | 4 | good | 0 | 0 | 0 | 0 | 0 | 0 |
| Patient 5 | T5 | T4 | 4 | good | 2 | 750 | mod | 0 | mild | mild |
| Mean | T6.4 | T4.8 | 4 | 1 | 416.7 |

* Graded by the surgeon
† Modified Bromage scale: 1=unable to move feet or knees, 2=moves foot/feet, 3=moves knee/knees, 4=straight leg raise
‡ Graded on a scale: 0=not present, 1=mild, 2=mod, 3=severe
GA = general anesthetic conversion; data ineligible for consideration

TABLE II Recovery characteristics of small-dose spinal lidocaine-sufentanil (Group LS) vs sufentanil (Group S)

<table>
<thead>
<tr>
<th>Time to S1 Sensation* (min)</th>
<th>Time to Motor Recovery* (min)</th>
<th>Time to ambulation‡ (min)</th>
<th>Pruritis (VAS /10)</th>
<th>Nausea†</th>
</tr>
</thead>
</table>

Group LS

| Patient 1 | 120 | 0 | 37 | 0.25 | 0.25 |
| Patient 2 | 30 | 60 | 93 | 8.00 | 0 |
| Patient 3 | 30 | 0 | 21 | 3.50 | 0 |
| Patient 4 | 30 | 0 | 35 | 5.00 | 0 |
| Patient 5 | 60 | 0 | 34 | 3.25 | mild |
| Mean | 75 | 40.8 | 4.04 |

Group S

| Patient 1 | 90 | 30 | 61 | 2.00 | severe |
| Patient 2 | GA | GA | GA | GA | GA |
| Patient 3 | GA | GA | GA | 4.50 | 0 |
| Patient 4 | GA | GA | GA | GA | GA |
| Patient 5 | 30 | 0 | 29 | 5.00 | severe |
| Mean | 60 | 40 | 3.83 |

* From time of arrival in recovery room to time of sensory or motor recovery (return of S1 sensation to pinprick & ability to deep knee bend, respectively)
† From time of spinal injection to time of first successful independent ambulation
‡ Maximal grade of nausea experienced: absent, mild, moderate-no vomiting, severe vomiting
GA = general anesthetic conversion; data ineligible for consideration
In the study, it became evident that the usual depth of anesthesia provided by small dose spinal anesthesia was not being met in all subjects. Only one of five (20%) Group S patients had adequate anesthesia during trocar insertion at the beginning of laparoscopy. This success rate was felt to be sufficiently low to warrant early closure of the study on ethical grounds, even though the goal of 20 patients (10 per group) was not achieved and the small sample size did not allow for meaningful statistical analysis.

Interestingly, a “level” of dullness to pinprick was always established in Group S patients, reflecting the profound segmental antinociception of intrathecal opioids. Sufentanil has no local anesthetic effect on nerve conduction in isolated dorsal root axons when it is administered in clinically relevant concentrations. In labouring women receiving ITS, the lack of sympathectomy, as measured by differences between calf and toe temperatures, and normal motor strength also support the fact that sufentanil does not act significantly as a local anesthetic. The decrease in blood pressure after ITS for labour analgesia is not caused by sympathectomy, but rather due to pain relief. In male volunteers, ITS does not affect blood pressure, yet other effects of sufentanil including pruritus and sensory changes to pinprick and cold are similar to those in women during labour.

Achievement of a differential level of sensation to pinprick in Group S patients did not reliably translate into adequate anesthesia for laparoscopy. Local anesthetics and opioids are frequently used together during spinal and epidural anesthesia for their dose-sparing effects to minimize toxicity and side effects. A study of intrathecal sufentanil and epidural bupivacaine alone and in combination in women during labour showed that markedly reduced doses of local anesthetics and opioids can be used in combination to provide adequate analgesia compared to either drug alone.

Isobolographic evaluation concluded that ITS and epidural bupivacaine had a potentially synergistic interaction, although a purely additive interaction could not be excluded due to the study design. It is likely that the equivalent dose of intrathecal 10 mg lidocaine with 10 µg sufentanil is much greater than 20 µg sufentanil since the combined sufentanil and bupivacaine fractional dose ED₅₀ was found to be approximately one-third for sufentanil and one-tenth for bupivacaine (of the single-drug ED₅₀).

Although there have been several case reports of respiratory arrest associated with intrathecal sufentanil, 20 µg ITS is felt to be safe for outpatient anesthesia. Factors which may increase the risk of respiratory arrest following intrathecal opioids include: high doses of opioid, repeated doses of opioid, intravenous sedatives or opioids, advanced age, co-existing disease, lack of opioid tolerance, and general anesthesia. Respiratory depression after ITS has been reported as early as four minutes after administration and as late as 30 min after administration. SO ur procedure duration ranged from 5 to 33 min. Patients were then

General anesthesia (GA) was induced and the patient recovered uneventfully, with no neurological deficit. The study was terminated early on ethical grounds because a large number of unacceptable blocks occurred compared to our usual experience with low dose local anesthetic and opioid mixtures. Two of five (40%) Group S patients had inadequate sensory block of the abdomen, as determined by the toothed forceps test, and one of seven (14%) Group LS patients had sharp perineal sensation without abdominal testing, and were converted to GA. A further two of the remaining three (67%) Group S patients, who passed the forceps test, felt mild transient sharpness during skin incision. Patients who received GA were excluded from the intraoperative (Table I) and recovery data (Table II).

Abdominal cramping and shoulder tip pain associated with CO₂ insufflation were common (three of six in Group LS and two of three in Group S). Only one of three Group S patients was satisfactorily comfortable with one dose or less of alfentanil, whereas five of six Group LS patients were comfortable with this dose. Higher doses of alfentanil may be related to increased postoperative nausea and vomiting (PONV) but this was difficult to interpret from the small amount of data. One of three Group LS patients compared with two of three Group S patients experienced some degree of PONV. Frequency of pruritus was pared with two of three Group S patients experienced some degree of PONV.

Achievement of a differential level of sensation to pinprick in Group S patients did not reliably translate into adequate anesthesia for laparoscopy.
monitored in the recovery room for more than 30 min. The mean duration of anesthesia in outpatients who received 20 µg ITS for extracorporeal shock wave lithotripsy was 64 ± 13 min, which clearly outlasted the time period at risk.

The abdominal toothed forceps skin test was not performed on the final study patient from Group LS, who had pressure discomfort with the vaginal speculum examination prior to laparoscopy and sharp perineal sensation on testing with a tenaculum. Her discomfort did not ease with 500 µg alfentanil iv and, consequently, GA was induced. This was the first "failure" of a lidocaine-sufentanil spinal solution in three series of patients at our institution. Retrospectively, we should have performed the skin test of the abdomen, as per protocol. This patient may have been adequately blocked for the laparoscopic procedure, without sacral anesthesia, but we were so sensitized to the possibility of block failure that we prematurely converted to GA.

Intrathecal 10 mg lidocaine with 10 µg sufentanil has been shown to be effective for laparoscopy, with the benefit of immediate postoperative ambulation, and reduced time to recovery of sensation to touch and pinprick compared to other low dose spinal anesthetic solutions. Intrathecal sufentanil is used effectively for labour and delivery analgesia with an ED₅₀ and ED₉₀ of 2.6 and 8.9 µg, respectively, in labouring women. In a study comparing 20 µg ITS with conventional dose hyperbaric lidocaine for extracorporeal shock wave lithotripsy, there was no significant difference in supplementation required or in intraoperative and postoperative pain between groups. Patients receiving intrathecal sufentanil could ambulate, tolerate oral intake, and void earlier than the lidocaine group and were discharged home significantly earlier.

The dose of 20 µg ITS was felt by the authors to be a safe and effective dose that was not associated with excessive sedation. Intrathecal 20 µg sufentanil has also been used successfully at our institution prior to this study for three short duration outpatient laparoscopic procedures.

In summary, 20 µg sufentanil intrathecally is inferred by this study to be inferior to 10 mg lidocaine with 10 µg sufentanil intrathecally for providing adequate anesthesia reliably for short duration gynecological laparoscopic procedures. Moreover, there are no advantages with respect to side effects or rapidity of recovery of sensory and motor function.

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**References**

13. Camann W, Aboulade A, Eisenach J, McLeod D, Datta S. Intrathecal sufentanil and epidural bupivacaine for labor analgesia: dose-response of individual agents and in...