Comparative Evaluation of Intrathecal Use of two Different Doses of Dexmedetomidine Along with Bupivacaine in Lower Abdominal Surgeries

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Running Title- Intrathecal Use Of Two Different Doses Of Dexmedetomidine

Abstract:
Introduction: The addition of adjuvants to local anaesthetics in spinal anaesthesia avoids intraoperative somatic and visceral pain and provides prolonged post operative analgesia. Aim –The present study was designed to determine the dose related effects of intrathecal dexmedetomidine added as adjuvant to 0.5% hyperbaric bupivacaine on block characteristics, haemodynamics and analgesia potentiating effects. Patients and method –This prospective randomised double blind study included 90 patients undergoing lower abdominal surgeries, who were randomly allocated into three groups of 30 patients each. Group B received 12.5 mg of 0.5% hyperbaric bupivacaine intrathecally, group D5 received 12.5 mg of 0.5% hyperbaric bupivacaine and 5 µg of dexmedetomidine, group D10 was given 12.5 mg of 0.5% hyperbaric bupivacaine and 10 µg of dexmedetomidine, all three made up to a total volume of 3 ml with NS. The three groups were compared with respect to haemodynamic parameters, onset and regression of motor and sensory block, duration of analgesia, doses of rescue analgesia required and 24 hours complications. The mean time of onset of sensory block to T10 was (in minutes) group B 11.6±1.12, group D5 5.84±2.02, group D10- 4.92±1.23 (p <0.001). Total duration of sensory block was (in minutes) group B 172.61±24.68, group D5 268.34±28.42, group D10 346.28±44.8 (p <0.001). Total duration of motor block (in minutes) group B 150.66±18.64, group D5 268.44±24.85, group D10 322.9±49.68 (p <0.001). Duration of analgesia (in minutes) - group B 124.01±8.552, group D5 194.68±18.44, group D10 290.48±20.64 (p <0.001). Conclusion: dexmedetomidine added to hyperbaric bupivacaine intrathecally has a dose dependent favourable effect on the onset and regression of motor and sensory block.

Keywords: spinal, dexmedetomidine, bupivacaine

I. Introduction

Effective management of perioperative and postoperative pain after lower abdominal surgeries represents an important component of postoperative recovery as it serves to blunt the autonomic, somatic and endocrine reflexes with a resultant potential of decreasing perioperative morbidity.¹ Multimodal drug analgesia as well as regional techniques have been the most common practice to treat perioperative pain but no method has been identified as yet to specifically block nociception without associated side effects.² Intrathecal block is preferred technique because it is simple and easy to administer, quite economical, has a rapid onset of action with less failure rates and superior level of blockade.³ Bupivacaine is the most commonly used amide local anaesthetics that has a prolonged duration of action and lower incidence of toxic reactions.⁴ It acts by blocking release of norepinephrine at locus ceruleus and exerts its analgesic action both at spinal and sensory levels.⁵ Various adjuncts such as opioids, epinephrine, neostigmine, magnesium, midazolam, ketamine and clonidine have been added to intrathecal local anaesthetics to prolong analgesia and accelerate functional recovery.⁶ Dexmedetomidine is an s enantiomer of medetomidine, a highly selective α2 adrenoceptor agonist with hypnotic, sedative, anxiolytic, opioid sparing and analgesic properties without producing significant respiratory depression.⁷,⁸ It acts by blocking release of norepinephrine at locus ceruleus and exerts its analgesic action both at spinal and sensory levels.⁹

Addition of dexmedetomidine to spinal bupivacaine produces shorter onset of motor block and a prolongation in the duration of motor and sensory block with preserved haemodynamics and minimal side effects.⁴ Previous studies have described the intrathecal use of dexmedetomidine in a wide range (2-15 µg).¹⁰,¹¹ To compare the subarachnoid block characteristics (onset and duration of motor and sensory block) 5 µg and 10 µg of dexmedetomidine was added to 12.5 mg of 0.5% heavy bupivacaine intrathecally. Our study sought to investigate the dose dependent effects of dexmedetomidine on the duration of motor and sensory block, duration

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of analgesia, need for postoperative analgesia and haemodynamic parameters. Also our aim was to find out the dose dependent increase in dexmedetomidine side effects.

| Table 1: Demographic profile of group B, D5, D10 |
|---------------------------------|----------|----------|---------|----------|
| Demographic profile            | Group B  | Group D5 | Group D10 |
| Mean age in years              | 36.32±10.2 | 40.23±11.20 | 38.11±8.62 |
| Weight in kg                   | 63.42±10.68 | 64.66±8.72 | 64.12±9.0  |
| Sex (%)                        | female 16(53%) | 17(56%) | 18(60%) |
|                                | male 14(47%) | 13(44%) | 12(40%) |
| ASA grade                      | Grade I 22(73%) | 24(80%) | 25(83%) |
|                                | Grade II 8(27%) | 6(20%)  | 5(17%)  |

<table>
<thead>
<tr>
<th>Table 2: Types Of Surgeries (Lower Abdominal Surgeries)</th>
</tr>
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<tbody>
<tr>
<td>surgery</td>
</tr>
<tr>
<td>1 Inguinal hernia</td>
</tr>
<tr>
<td>2 Abdominal hysterectomy</td>
</tr>
<tr>
<td>3 Benign prostatic hypertrophy</td>
</tr>
<tr>
<td>4 Umbilical hernia</td>
</tr>
<tr>
<td>5 Haemorrhoids</td>
</tr>
<tr>
<td>6 Hydrocele</td>
</tr>
<tr>
<td>7 Appendicectomy</td>
</tr>
</tbody>
</table>

II. Materials And Methods

Our study was a randomized, patient and observer blind (double blind), concentration controlled, single centre trial. After obtaining permission from institutional ethics committee written informed consent was taken. A total of 90 patients were randomly allocated into 3 equal groups of 30 each using computer generated random numbers inserted into sealed envelops marked 1-90. Patients having ASA physical status I and II age 20-60 years and of either sex undergoing elective various lower abdominal surgeries under spinal anaesthesia were included in the study.

Exclusion criteria – patient refusal, any known allergy or contraindication to bupivacaine or dexmedetomidine, pregnancy, hepatic, renal or cardiopulmonary abnormalities, alcoholism, diabetes, long term analgesic or anticoagulant therapy, spinal cord deformities, neurological or neuromuscular deficits, paralysis, bleeding diathesis, local skin site infections or patients on α₂ adrenergic receptor antagonists, calcium channel blockers, ACE inhibitors, morbid obesity (Body weight >120 kg) or height <150 cm.

A day before surgery a detailed preanaesthetic checkup was carried out. Patients were asked to restrict fluids and solid by mouth at least six hours before operation. Interpretation of VAS (visual analogue scale) was explained to determine level of analgesia in the postoperative period. This was carried out with 10 cm line. The first end marked ‘0’ means ‘no pain’ and the end marked ‘10’ means ‘worst pain imaginable’.

All patients were clinically examined in the preoperative period, where whole process was explained. On entering the patient in the operating room standard intraoperative monitors like ECG, pulse oximeter, NIBP were attached and baseline parameters (blood pressure, heart rate, respiratory rate) were recorded. The patients were preloaded with lactated Ringer’s solution (RL) 10 ml/kg. Technique used was standardized for all patients. Peripheral IV line was secured with 18 G cannula. After preloading with RL and under aseptic preparation, lumbar puncture was performed at L3 – L4 intervertebral space (L2 – L3, in case difficulty occurs) in median approach with 25G Quinke spinal needle. The patients were randomized into 3 groups B, D5, D10 of 30 patients each using sealed envelope technique. The dose of 0.5% hyperbaric bupivacaine 12.5 mg was identical in all study groups. In group B patients received 12.5 mg of 0.5% bupivacaine which was made upto 3 ml after dilution in .9% saline (NS). Group D5 received 12.5 mg of 0.5% bupivacaine and 5μg of dexmedetomidine, total drug volume made upto 3 ml with .9% saline (NS). Group D10 received 12.5 mg of hyperbaric bupivacaine and 10μg of dexmedetomidine, total volume made upto 3 ml with .9% saline.

The intrathecal drug formula was prepared by a separate anaesthesiologist by a sterile technique who was blinded to the study and block given by a different anaesthetist who also monitored block characteristics and was also blinded to the study groups. PR, RR, BP were monitored every 5 minutes for first 15 minutes then every 10 minutes till the end of surgery and every 30 minutes in the postoperative ward till 1 hour and every 2 hours thereafter.

Incidence of intraoperative hypotension (decrease of systolic BP 20% from baseline or systolic BP<90) was recorded. Hypotension was treated with oxygen, bolus administration of 250 ml lactated Ringer’s solution over 10 minutes, or with intermittent doses of intravenous mephentermine at 6 mg. Bradycardia (HR <50 beats /minute) and tachycardia (HR>100 beats/minute) were also recorded. Episodes of bradycardia were treated with incremental doses of atropine at 0.3 mg administered intravenously. The total duration of surgery was noted.
Sensory block was assessed by loss of sensation to pin prick in midline using a 22G blunt hypodermic needle at 2 minute interval till block reaches T8 at which level surgery was allowed to proceed. Level was assessed at 3 minute interval till no change in level was seen, thereafter every 20 minute interval. Onset of sensory block to T10 dermatome, peak level of sensory block and duration of sensory block (regression to S1 dermatome) was noted.

Degree of motor block was assessed by means of James modified Bromage score (0-no resistance, able to raise a leg straight against resistance, 1-able to raise leg straight but able to flex knee, 2- unable to flex knee but with free movement of feet, 3-unable to move leg or feet). Onset time to reach Bromage 3 and time taken for regression to bromage 0 were noted.

All durations were calculated by taking the time of drug administration intrathecally as time 0. Analgesia was monitored by using VAS score. VAS was recorded 5 minutes before spinal, at start of surgery and then every 15 minute interval till the surgery was over. Postoperatively VAS was recorded half hourly for first hour then 1 hourly for 12 hours, and then 3 hourly for next 12 hours till 24 hours. When patient had VAS> 3 rescue analgesia in the form of diclofenac sodium 75 mg intramuscular (I/M) or when needed inj. Tramadol 50 mg slow IV was given. Time to first dose of rescue analgesia, number of doses of rescue analgesia and the time at which it was repeated was recorded in all groups. The time at which patient demanded first dose was the primary end point of this study because at this point the effect of spinal anaesthesia has weaned off.

In case of failed spinal block patients were given general anaesthesia and these patients were excluded from the study. The quality of surgical analgesia was assessed and graded as: Excellent: if no supplemental drugs were required, Good: only one analgesic required, Fair: if more than one analgesic required, Poor: GA required. If full surgical anaesthesia was not achieved then inj. tramadol 50-100 mg IV was given as supplementary analgesia during surgery. Patients were monitored for sedation every 10 minute interval for first 30 minutes and then every 15 minute interval till completion of surgery. Following sedation score was used

0: no sedation
1: patient somnolent but responding to verbal commands
2: patient somnolent not responding to verbal commands
3: patient somnolent and not responding to verbal commands and manual stimulation.

Our primary outcome after completion of surgery was to compare the duration of motor and sensory block, secondary outcome was to observe the onset of motor and sensory block, level of sensory block achieved, haemodynamic parameters, duration of motor and sensory block, time and dosage of analgesic used, to compare pain scores among the three groups and any side effects noted throughout the study period and 24 hours postoperatively.

Any side effects or complications like hypotension and bradycardia headache, dry mouth, nausea, vomiting, local anaesthetic toxicity, backache and urinary retension were noted in these 24 hours. This was secondary end point of our study.

### III. Statistical Analysis

The statistical analysis of the data was done by using SPSS evaluation version 20 (statistical package for social sciences). The data was expressed as either mean or standard deviation for number and percentages. The demographic data of patients was studied for number and percentages.

The total sample size calculated was 42 (14 patients in each group). Power analysis using the following parameters was carried out \((a=0.05, \beta=0.8)\) total duration of analgesia and time to bromage scale 0. We increased the total number of patients to increase the power of study. \(p\) value <0.01 was considered highly significant. Post hoc power analysis was carried out using a power and sample size calculator. The cut off value for power analysis was taken as at least 80% \((\beta=0.8).\) The effective size/power was calculated for duration of analgesia \((\beta=1)\) and the duration of motor block \((\beta=1)\) determined as >80%. Thus the post hoc assessment of effective size justified the sample size.

### Table 3: Sensory And Motor Block Characteristics

<table>
<thead>
<tr>
<th>characteristic</th>
<th>Group B</th>
<th>Group D5</th>
<th>Group D10</th>
<th>p value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Onset time to T10 (min)</td>
<td>11.6±1.12</td>
<td>5.8±4.5±2.02</td>
<td>4.9±1.23</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Max. level of sensory block</td>
<td>T6</td>
<td>T6</td>
<td>T6</td>
<td>&gt;.5</td>
</tr>
</tbody>
</table>

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There was no statistical difference in the patients demographic variables or duration of surgery as shown in Table 1. Table 2 shows the number of patients in each group undergoing different types of lower abdominal surgeries. The number of patients under each type of surgery performed were similar amongst the groups thereby keeping the comparison unbiased.

The sensory and motor block characteristics are shown in Table 3. The mean time required for onset of sensory block to T10 dermatome in group D10 (4.92±1.23) was more rapid than either group D5 or B. Also onset of sensory block to T10 was more rapid in group D5 than in group B (p<0.001). The maximum upper level of sensory block achieved in group B or D5 was T6-8 dermatome with a median value of T6 and group D10 it was T5-T8 dermatome with a median of T6 dermatome, which was comparable in all three groups. However the maximum level of sensory block was achieved earlier in group D10 (8.02±0.412) than in group D5 or B. Also it was earlier in group D5 than in group B and the difference was highly significant (p<0.001).

The addition of dexmedetomidine has significant effect on sensory block. The mean time taken for regression of sensory block to S1 dermatome was much more prolonged in D10 (346.28±44.8 minutes) than in group D5 (268.44±24.85 minutes) and this was much longer in comparison to group B (150.66±18.64). The prolongation in time to regression in group B vs group D5, group B vs group D10 and group D5 vs group D10 was highly significant statistically by Tuckey’s test (p<0.001).

Complete motor block was achieved earlier in group D10 and group D5 than group B and it was much earlier in group D10 than in group B. It was 9.42±2.36 minutes in group D10 and 12.96±2.64 minutes in group D5 and 18.02±0.96 minutes in group B. Thus the difference was statistically highly significant (p<0.001). Statistical analysis by ANOVA and Tuckey’s showed that the total duration of motor block was significantly prolonged in group D10 and D5 as compared to group B. Similarly it was more prolonged in group D10 as compared to group D5 and the difference was highly significant (p<0.001).

Patients remained pain free for a longer duration in group D10 and D5 than in group B. Pain free duration of rescue analgesia in group D5 was shorter than group B and it was much shorter in group D5 than in group B and the difference was highly significant. Pain free duration of rescue analgesia in group D5 was significant for 4th hour and second dose in group D5 and 10th hour and IV tramadol was given. The second dose of injectable diclofenac was given in the 13th hour.

In group D5 first dose of rescue analgesia (VAS>3) was given in 3rd to 4th hour and second dose in 14th to 15th hour and thus the difference is highly significant in comparison with group B.

In group D10 the first dose of rescue analgesia was given in 5th to 6th hour and second dose at 17th to 18th hour. None of the in groups D5 or D10 required injectable tramadol. When Tuckey’s post hoc analysis was applied the difference in duration of analgesia in between group D5 and group D10 was highly significant which confirmed that the increase in the duration of analgesia after addition of dexmedetomidine is dose dependent.

Table 4: Surgical Characteristics, Adverse Effects And Treatment

<table>
<thead>
<tr>
<th></th>
<th>Group B</th>
<th>Group D5</th>
<th>Group D10</th>
<th>p value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Total IV infusion (in ml)</td>
<td>1100±148.2</td>
<td>921±216.4</td>
<td>1028±284.3</td>
<td>.32</td>
</tr>
<tr>
<td>Duration of surgery(in minutes)</td>
<td>54.66±5.64</td>
<td>43.84±11.68</td>
<td>50.64±3.3</td>
<td>.64</td>
</tr>
<tr>
<td>Billed transfusion (no. of patients)</td>
<td>1</td>
<td>2</td>
<td>1</td>
<td>.54</td>
</tr>
<tr>
<td>Additive analgesia (no. of patients)</td>
<td>2</td>
<td>1</td>
<td>0</td>
<td>.56</td>
</tr>
<tr>
<td>Nausea/vomiting (no. of patients)</td>
<td>1</td>
<td>0</td>
<td>1</td>
<td>.34</td>
</tr>
<tr>
<td>Bradycardia</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>.12</td>
</tr>
<tr>
<td>Hypotension</td>
<td>2</td>
<td>1</td>
<td>3</td>
<td>.64</td>
</tr>
<tr>
<td>Atopine</td>
<td>0</td>
<td>1</td>
<td>1</td>
<td>.36</td>
</tr>
<tr>
<td>Mephentermine</td>
<td>1</td>
<td>2</td>
<td>1</td>
<td>.23</td>
</tr>
<tr>
<td>Shivering</td>
<td>2</td>
<td>1</td>
<td>0</td>
<td>.56</td>
</tr>
</tbody>
</table>

IV. Results

All patients (90) completed the study, there was no statistical difference in the patients demographic variables or duration of surgery as shown in Table 1. Table 2 shows the number of patients in each group undergoing different types of lower abdominal surgeries. The number of patients under each type of surgery performed were similar amongst the groups thereby keeping the comparison unbiased.

The sensory and motor block characteristics are shown in Table 3. The mean time required for onset of sensory block to T10 dermatome in group D10 (4.92±1.23) was more rapid than either group D5 or B. Also onset of sensory block to T10 was more rapid in group D5 than in group B (p<0.001). The maximum upper level of sensory block achieved in group B or D5 was T6-8 dermatome with a median value of T6 and group D10 it was T5-T8 dermatome with a median of T6 dermatome, which was comparable in all three groups. However the maximum level of sensory block was achieved earlier in group D10 (8.02±0.412) than in group D5 or B. Also it was earlier in group D5 than in group B and the difference was highly significant (p<0.001).

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Complete motor block was achieved earlier in group D10 and group D5 than group B and it was much earlier in group D10 than in group B. It was 9.42±2.36 minutes in group D10 and 12.96±2.64 minutes in group D5 and 18.02±0.96 minutes in group B. Thus the difference was statistically highly significant (p<0.001). Statistical analysis by ANOVA and Tuckey’s showed that the total duration of motor block was significantly prolonged in group D10 and D5 as compared to group B. Similarly it was more prolonged in group D10 as compared to group D5 and the difference was highly significant (p<0.001).

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In group D10 the first dose of rescue analgesia was given in 5th to 6th hour and second dose at 17th to 18th hour. None of the in groups D5 or D10 required injectable tramadol. When Tuckey’s post hoc analysis was applied the difference in duration of analgesia in between group D5 and group D10 was highly significant which confirmed that the increase in the duration of analgesia after addition of dexmedetomidine is dose dependent.
The total number of rescue analgesia doses required at 24 hours postoperatively were also significantly less in group D10 as shown in table 3. Haemodynamic parameters remained stable and were comparable in both groups at all measured intervals (fig 1). Incidence of nausea and vomiting was comparable. Only one patient required IV mephentermine.

In our study the onset of sensory block was earlier in dexmedetomidine group compared to other groups. Shukla et al also showed that onset time to peak sensory level was earlier in dexmedetomidine group compared to control group.19

In this prospective, randomized, double blind study in patients scheduled for lower abdominal surgery, we compared the dose dependent effect of 5 µg and 10 µg of dexmedetomidine added to 12.5 mg of intrathecal bupivacaine on the onset time and duration of motor and sensory block as well as on the postoperative rescue analgesia required and associated side effects if any. The demographic profile in all three groups which was statistically insignificant (p>0.05) was quite similar with other research investigations and provided us with a uniform platform to evenly compare the results obtained.

A similar study was conducted by Mustafa et al, in which 66 patients scheduled for urological procedures were randomly divided into 3 groups and given dexmedetomidine 5 µg and 10 µg along with 12.5 mg of bupivacaine.14 Our results are similar and further confirm the fact that when dexmedetomidine is added as adjuvant to bupivacaine in spinal anaesthesia, the prolongation of motor and sensory block occurs in a dose dependent manner that is as the dose of dexmedetomidine is increased more is the duration of motor and sensory block and duration of analgesia. In our study the onset time to highest level was also noted due to level of incision in lower abdominal surgeries. The postoperative analgesia is even more prolonged with 15 µg of dexmedetomidine, which may be beneficial in patients undergoing lengthy and complex surgeries, but this dose leads to higher sedation scores which may be undesirable.12 We chose 5 µg and 10 µg of dexmedetomidine in comparison with plain bupivacaine so that in addition to prolongation of analgesia by dexmedetomidine, its dose dependent effect may be confirmed with minimal side effects.15

Tarbeehet et al and Jamilia RH et al also found that dexmedetomidine has a dose dependent effect on onset and regression of motor and sensory block and time to rescue analgesia with lower VAS scores and minimal side effects when used as an adjuvant to intrathecal bupivacaine. Similar results were found by balder et al.16,17,18

In our study the onset of sensory block was earlier in dexmedetomidine group with much earlier in group D10. Ogan et al showed an earlier significant peak sensory block in dexmedetomidine group compared to other groups. Shukla et al also showed that onset time to peak sensory level was earlier in dexmedetomidine group compared to control group.

A study done by Kanazi et al including 60 patients undergoing transurethral resection of prostrate or bladder tumour under spinal anaesthesia reported shorter onset time of motor block but longer sensory and motor regression times in bupivacaine given with dexmedetomidine (3 µg) as compared with bupivacaine alone. They showed that mean time to sensory regression to S1 was 303 ±75 minutes whereas in our study it was longer, which clearly indicate dose dependent effect of dexmedetomidine.
The regression of motor block to Bromage 0 was 322.94 ± 9.68 minutes with higher dose of dexmedetomidine (10 µg). Similar results were shown by Kanazi et al.20 As regards the first time to require analgesia and total analgesic consumption of diclofenac in 24 hours, group D10 showed a significant increase in time to first analgesic dose (290.48±20.64 minutes) and significant decrease in total analgesic consumption. In agreement with our results, Eid and colleagues, showed a significantly longer time to first analgesic request compared to control group.12 Ashraf and colleagues also showed a significantly longer time to first analgesic request (3.30±0.87 hours) compared to control group 90.23±0.11 hours).

In the present study, no sedative was given during premedication, and thus most of the patients had sedation score in the range of 0 and 1 at all measured intervals in both groups. It has also been observed earlier that addition of low-dose dexmedetomidine to intrathecal bupivacaine does not lead to higher sedation scores. Patients remained hemodynamically stable in both groups at all measured intervals for 24 h. Dexmedetomidine as an intrathecal adjuvant to bupivacaine in a dose dependent manner does not produce any significant hemodynamic changes and vitals remained stable both intraoperatively and postoperatively.

Further studies are required to rule out any short term or long-term adverse effects of intrathecal dexmedetomidine, although in our study 24 hours follow up showed no significant side effects. Patients only in ASA grade I and II were included in our study. Safety of dexmetomidine therefore needs to be evaluated in patients with known cardiovascular or other comorbidities or in pregnancy. Another limitation being that we limited our dose to 10 µg of dexmedetomidine. Further studies are therefore required with higher doses of dexmedetomidine to better know its side effects and limitations. Effect of adding dexmedetomidine to other local anaesthetics like ropivacaine or levobupivacaine in other neuraxial blocks needs further research.

VI. Conclusion

Finally we do conclude that addition of dexmedetomidine in 10 µg dose compared with 5 µg dose provides fast onset and longer duration of motor and sensory blockade and also reduced the requirement of local anaesthetics like ropivacaine or levobupivacaine in other neuraxial blocks needs further research.

Bibliography

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**Figure 1:** Comparison of Preoperative Vitals

![Comparison of Preoperative Vitals](image)