COMPARISON OF DIFFERENT DOSES OF EPIDURAL BUTORPHANOL 2 MG/4 MG AS ADJUVANT TO 0.125% BUPIVACAINE FOR POSTOPERATIVE ANALGESIA FOR VARIOUS INFRAUMBILICAL SURGERIES

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ABSTRACT

BACKGROUND
Postoperative analgesia is widespread in this era. There are multimodal techniques available for the same. The aim of this study was to compare efficacy, effectiveness, safety, of different doses of butorphanol as adjuvant to Bupivacaine 0.125% for postoperative analgesia. Study design was randomised double blind comparative study.

MATERIALS AND METHODS
This randomized clinical comparative study was conducted in 50 patients of either sex, aged between 18-60 years of ASA grade I and II admitted for elective surgery. Written informed consent was taken and pre-anesthetic evaluation was done. Epidural catheter was inserted, and all patients were given spinal anaesthesia using 0.5% hyperbaric bupivacaine (15 mg). In the postoperative period, when patient complained of pain, or VAS score was 3 or more, study drug was given. In group A, epidural bupivacaine (0.125%) 10 ml and preservative free butorphanol 2 mg diluted to 2 ml with NS, & in Group B, bupivacaine (0.125%) 10 ml and preservative free butorphanol 4 mg diluted to 2 ml with NS was given. Parameters observed were; onset of analgesia; duration of analgesia; cardio- respiratory effects and adverse effects. Data was analysed with SPSS Software 16 (IBM, Armonk, NY, USA). Unpaired t test, was used for qualitative data. chi-square test used for categorical data (n, number, %).

RESULTS
Age, sex, weight, haemodynamics, and ASA grade were comparable in both groups. Mean onset of analgesia was rapid (4.5±1.06 minutes) in group B when compared to group A (6.0±1.32 minutes). This was clinically and statistically significant (p<0.001). Duration of analgesia was longer in group A which was mean of 410±50.29 minutes compared to group B which was a mean of 490±39.32 minutes. This was clinically and statistically significant (p<0.001). 24 hour VAS was 3.0±1.2 in Group A, 2.5±1.0 in Group B. Total Analgesic request in 24 hours was 2.4±0.5 in group A, 1.7±0.2 in group B. There was no significant difference in heart rate, blood pressure, and respiratory rate, as monitored at regular intervals postoperatively between the two study groups. Regarding adverse reactions, sedation was the main side effect in group B (40%) patients, in group A it was (36%). Frequency of pruritis 4% and nausea-vomiting was more in group B (12%), in group A it was 0% & 8% respectively. Hypotension and respiratory depression were in 4% patients in group B which was statistical insignificant. All patients were monitored for 24 hours postoperatively for any untoward effects & for haemodynamic stability as well as for VAS.

CONCLUSION
Butorphanol is a good adjuvant to 0.125% bupivacaine in epidural analgesia. 4 mg butorphanol provides prolonged analgesia than 2 mg.

KEY WORDS
Epidural Butorphanol, Different Doses, Postoperative Analgesia, Infraumbilical Surgeries.

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Butorphanol 2 mg/4 mg have been selected as an adjuvant to Bupivacaine (0.125%) for postoperative epidural analgesia.

Aims & Objectives
A comparison of epidural Bupivacaine and Butorphanol 2 mg versus epidural Bupivacaine and butorphanol 4 mg for post operative analgesia focusing on Onset and duration of analgesia, Cardio respiratory effects, Adverse effects.

MATERIALS AND METHODS
Fifty adult patients of ASA grade I and II, of either sex, belonging to 18-60 years of age, posted for elective infraumbilical surgeries in general surgery, orthopaedics, gynaecology, urology and plastic surgery were selected for the study.

Group Allocation
- Randomisation was done by Randomisation table.(10)
- Randomised group Allocation was done by sealed opaque envelope method.
- Execution of sealed envelope was done at time of combined spinal epidural anaesthesia.

Group A
Bupivacaine (0.125%) 10 ml + Butorphanol (2 mg) in NS 2 ml of total 12 ml volume.

Group B
Bupivacaine (0.125%) 10 ml +Butorphanol (4 mg) in NS 2 ml of total 12 ml volume.

Sample Size
The sample size for the study was taken for convenience.

Exclusion Criteria
1. Patients with cardio-respiratory disorders.
2. Patients with renal and/ or hepatic disorders.
3. Contraindications for epidural anaesthesia.
4. Patients physically dependent on narcotics.
5. Patients with history of drug allergy.
6. Head injury cases.

Pre-Anaesthetic Evaluation
Patients were visited on the previous day of the surgery, a detailed clinical history was taken, and general and systemic examinations were done. Basic laboratory investigations ECG and chest x-ray were carried out routinely on all patients. The patients were explained about the spinal-epidural technique and VAS scale. A written informed consent was taken from each patient.

Premedication
To allay the anxiety and apprehension all patients were given Tablet Alprazolam (0.25 mg) at 10 pm in the night before the surgery. Patients were kept nil orally for 8 hrs before surgery.

Anaesthesia Technique
After taking informed consent with all aseptic antiseptic techniques, double spaced combined spinal epidural anaesthesia (CSEA) technique was used.

- Epidural catheter was inserted for postoperative analgesia in L2-3 space with 18-gauge epidural needle by LOR technique (Loss of Resistance) & epidural catheter of 20 was introduced through it & advanced 3-4 cm in space. It was fixed and later all patients were operated under spinal anaesthesia given in one space below epidural catheter, using hyperbaric bupivacaine (0.5%) 3 ml (15 mg).
- A large bore intravenous line was secured with 18G cannula and Ringer lactate (R.L.) 10 ml/kg/hr infusion was started.
- Routine monitors like ECG, Non-invasive Blood pressure, Pulseoximetry were connected for every case and basal vital signs were recorded before starting the procedure.
- Drugs and equipment necessary for resuscitation and general anaesthesia administration were kept ready. An autoclaved spinal-epidural tray was used.
- Sterile disposable epidural set was used and checked for any manufacturing problems.
- Under all aseptic and antiseptic precautions. The epidural space was identified using 18G Tuohy needle with Loss of Resistance (LOR) technique at L2- L3 interspace. Then 20G epidural catheter was passed through the epidural needle in upward direction till about 4cms of the catheter was in the space. The needle was withdrawn, and the catheter was fixed to the back using adhesive tape. Then spinal anaesthesia was given in the one interspace below the catheter with 23G Quinke needle using hyperbaric Bupivacaine (0.5 %) 3 ml (15 mg)
- Level of sensory blockade was accessed by pinprick method, motor block was accessed by modified Bromage score. After achieving sensory level of T6 & motor block of grade 3 surgery is started. Blood loss, urine output and other routine monitors as described above were observed.
- No narcotics were administered during the intraoperative period.
- Fluid management: To begin with, R.L was infused and maintained with R.L, N.S or D.N.S. Blood was transfused only when indicated.

Post Operative Period
After completion of the surgery, patient was shifted to recovery room and monitoring was continued. When patient recovered from motor blockade, they were shifted to postoperative ward.

In the postoperative period, when the patient first complained of pain, intensity of pain was assessed using VAS scale. When the VAS score was 4 or more, study drug was given through epidural catheter after confirming its proper position as follows:

Group A
Received Bupivacaine (0.125%) 10ml and preservative free Butorphanol 2 mg diluted to 2 ml in NS.

Group B
Received Bupivacaine (0.125%) 10 ml and preservative free butorphanol 4 mg diluted to 2 ml in NS. The intensity of pain and pain relief was assessed using VAS

at 5, 10, 15, 30, 60 minutes 2 hours, and thereafter 2-hourly for 24 hours postoperatively. As and when the patient complains of further pain during the period of observation, intensity of pain was assessed again using VAS to know the effect of the study drug given earlier. If it was 4 or more, rescue analgesia was given in form of Injection Tramadol 50 mg preservative free through epidural catheter slowly as per the institute protocol and the requirements of Epidural Tramadol (Rescue analgesic requests) in 24 hours noted.

Visual Analogue Scale
Visual analogue scale (VAS) consisted of a 10 cm line, marked at 1 cm each on which the patient makes a mark on the line that represents the intensity of pain he/she was experiencing. Mark ‘0’ represents no pain and mark ‘10’ represents worst possible pain. The numbers marked by the patient was taken as units of pain intensity. VAS was explained to each patient in Preoperative period.

Observations
1. Onset of analgesia.
2. Duration of analgesia.
3. Haemodynamic effects: Heart rate, blood pressure and respiratory rate.
4. Adverse effects like sedation, pruritus, nausea, vomiting, respiratory depression and hypotension. Urinary retention could not be studied, as patients in the study had indwelling urinary catheter inserted as part of the surgical management.
5. VAS was monitored for 24 hours.
6. Total Analgesic requests in 24 hours were noted.

Onset of Analgesia
Is the time interval from administration of the study drug (VAS score of 4 or more) to first reduction in pain intensity by at least 10 mm in VAS.

Duration of Analgesia
- Is the time interval between onset of analgesia, till patient complaints of pain (VAS score 3 or more) when rescue medication was given.
- Total analgesic requests in 24 hrs. were noted.

Sedation
Quality of sedation after giving the study drug was based on Ramsay sedation assessment scale.

Hypotension
A fall of 30 % in BP from baseline value.

Respiratory Depression
A respiratory rate of less than 10 breaths/ min.

Bradycardia
- A fall of 20% in pulse rate from base line value.
- Nausea vomiting- 4-point scale.

Management of Adverse Effects

Hypotension
IV fluids and Injection Ephedrine 6 mg kept ready.
Both groups were also comparable with respect of sex distribution, weight, haemodynamics characteristics and ASA Grade (p<0.05) of patients before study drug given.

<table>
<thead>
<tr>
<th>Parameters (Mean+/−SD)</th>
<th>Group A (n=25)</th>
<th>Group B (n=25)</th>
<th>P value</th>
<th>Inference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Onset of analgesia (mins)</td>
<td>6.0+/−1.32</td>
<td>4.5+/−1.06</td>
<td>&lt;0.001</td>
<td>HS</td>
</tr>
<tr>
<td>Duration of analgesia (mins)</td>
<td>410+/−50.29</td>
<td>490+/−39.32</td>
<td>&lt;0.001</td>
<td>HS</td>
</tr>
<tr>
<td>Total analgesic requests in 24 hrs (no.)</td>
<td>2+/−0.5</td>
<td>1+/−0.2</td>
<td>&lt;0.001</td>
<td>HS</td>
</tr>
<tr>
<td>VAS in 24 hrs</td>
<td>8.0+/−1.2</td>
<td>2.5+/−1.0</td>
<td>&lt;0.05</td>
<td>S</td>
</tr>
<tr>
<td>SBP (mm of Hg) in 24 hrs</td>
<td>118+/−12</td>
<td>114+/−14</td>
<td>&gt;0.05</td>
<td>NS</td>
</tr>
<tr>
<td>DBP (mm of Hg) in 24 hrs</td>
<td>68+/−6</td>
<td>66+/−8</td>
<td>&gt;0.05</td>
<td>NS</td>
</tr>
<tr>
<td>RR (per minute) in 24 hrs</td>
<td>14+/−3</td>
<td>12+/−4</td>
<td>&gt;0.05</td>
<td>NS</td>
</tr>
</tbody>
</table>

Table 2. Analgesic & Haemodynamic Characteristics after Study Drug

<table>
<thead>
<tr>
<th>Adverse Effects</th>
<th>Group A</th>
<th>Group B</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sedation</td>
<td>9 36</td>
<td>10 40</td>
</tr>
<tr>
<td>Pruritis</td>
<td>0 0</td>
<td>1 4</td>
</tr>
<tr>
<td>Nausea-Vomiting</td>
<td>2 8</td>
<td>3 12</td>
</tr>
<tr>
<td>Respiratory Depression</td>
<td>0 0</td>
<td>1 4</td>
</tr>
<tr>
<td>Hypotension</td>
<td>0 0</td>
<td>1 4</td>
</tr>
</tbody>
</table>

Table 3. Adverse Effects

**Sedation**

Was observed in 9 patients (45%) of Group A and 3 (12%) patients of Group B. This was statistically significant (p<0.05).

**Pruritis**

Was seen in 1 patient (4%) of group B and in none of the patients of group A which was statistically significant (p<0.05). It was treated with inj. Chlorpheniramine maleate.

**Nausea and Vomiting**

Were observed in 2 patients (8%) in group A and in 3 patients (12%) in group B which was statistically significant (p<0.05).

**Respiratory Depression**

Was seen in only 2 patients (8%).

**Hypotension**

In only 1 patient (4%) of group B and in none of the patients of group A. These were statistically insignificant (p>0.05).

**Sedation Score/Ramsay Sedation Score of 2 or 3 was observed**

In 9 patients (36%) in group A whereas 10 patients (40%) in group B had sedation of RSS score 2/3 which was statistically significant (p<0.001). The quality of sedation was acceptable in the interest of patients’ wellbeing. No airway intervention required in any patient.

Total analgesic requests in 24 hours was 2+/−0.5 in group A, & in it was 1+/−0.2 in group B which states that butorphanol dose dependently provide analgesia. More analgesia & less analgesic requests in group B suggest it.

**DISCUSSION**

The present comparative study done to assess the efficacy and safety of epidural Bupivacaine and Butorphanol versus epidural Bupivacaine and Fentanyl for the management of postoperative pain in lower abdominal and lower limb surgeries. A total of 50 patients belonging to age groups 18-60 years of ASA grade I-II have been taken. Male and female patient ratio was equal. Patients undergoing elective lower abdominal and lower limb surgeries in general surgery, orthopaedics, gynaecology, urology and plastic surgery were selected. During the preoperative assessment patients were explained about the epidural procedure and VAS score.

Pre-medication Tablet Alprazolam 0.25 mg orally was given the night before the surgery. Patients were randomly divided into two groups of 25 each, Group A – Bupivacaine and Butorphanol 2 mg and Group B – Bupivacaine and Butorphanol 4 mg. Epidural catheter was inserted and all patients were given spinal anaesthesia. In the postoperative period, when patient complained of pain, or when VAS score was 4 or more, patients in group A received epidural Bupivacaine 0.125% 10 ml and Butorphanol 2 mg diluted to 2 ml NS and patients in group B received epidural Bupivacaine 0.125% 10 ml & inj. Butorphanol 4 mg diluted to 2 ml in NS. Observations recorded are Onset of analgesia, Duration of postoperative analgesia, Haemodynamic stability and adverse effects.

**Onset of Analgesia**

In our study, the mean time for onset of analgesia in group A was 6 ± 1.32 minutes and in group B was 4.5 ± 1.06 minutes. Majority of patients in group A had onset of analgesia between 4-6 minutes whereas in group B between 2-4 minutes. Statistical analysis showed that onset of analgesia was delayed in group A compared to group B (p<0.001).

Mok et al. in 1986 did a study to evaluate the analgesic efficacy and safety of epidural Butorphanol 4 mg in comparison to that of epidural morphine 5 mg in patients with postoperative pain. Onset of pain relief with epidural Butorphanol appeared at 15 minutes.

Aswini A. et al. in 2009 conducted a comparative study of epidural Butorphanol 4 mg and epidural Fentanyl 100 µg for the relief of postoperative pain in lower abdominal and lower limb surgeries. The onset of analgesia was clinically and statistically significantly late (6 minutes) in Butorphanol group when compared to Fentanyl group (3 minutes).

**Duration of Analgesia**

In the present study, duration of analgesia in group A was 410 ± 50.29 min and in group B was 490 ± 39.32 min. (p<0.001).

Less analgesic requests, Less VAS in 24 hrs, more duration of analgesia & less analgesic requests in group B suggest it.

Mok et al. in 1986 concluded that duration of analgesia with Butorphanol 4 mg averaged 5.4 hrs.

Shivakumar T. C. et al., in 2006 evaluated analgesic efficacy and side effects of 2 doses of epidural Butorphanol in lower abdominal surgeries. Patients were randomly assigned to three groups to receive epidural Bupivacaine 0.5% 16 ml (n=25 control group I), Bupivacaine 0.5% 15 ml + 1 ml 2 mg...
Butorphanol (n=25, group II) and Bupivacaine 0.5% 14 ml + 2 ml of 4 mg Butorphanol (n=25, group III). Maximum patients demanded rescue analgesics in Group I (36%) and Group II (32%) at 7th hour and in group III (40%) at 9th hour.

Neerja Bharti et al, in 2009. The duration of analgesia was prolonged in patients receiving Butorphanol (2 mg, 4 mg) with Bupivacaine (0.125%) combination (8.68 ± 0.82 hrs, 9.82 ± 0.54 hrs) as compared with Butorphanol alone (4.35 ± 0.66 hrs; P < 0.05).

Aswini A et al, in 2009. Duration of analgesia was clinically and statistically longer in Butorphanol group (350 minutes) in comparison to Fentanyl group (230 minutes).

Haemodynamic and Respiratory Effects
In the present study heart rate, blood pressure and respiratory rate remained stable throughout the observatory period. 1 patient in group B had hypotension (Fall in systolic BP <20% of basal reading) and 2 patients in group B had respiratory depression (RR<10/min) which was not anaesthetic practice.

Nausea and Vomiting
In our current study, in group B 2 patients had nausea and vomiting which was significant statistically (p<0.05). In group II 3 patients had nausea and vomiting which was significant statistically (p<0.05). No patients in group A had nausea or vomiting.

Pruritis
In our study none of the patients in group A had pruritis and 4 patients (16%) in group B had pruritis which was statistically significant (p<0.05). Premila Malik, Chhavi Manchanda, Naveen Malhotra in their study showed that there were no significant changes in pulse rate, systolic and diastolic BP, RR and SpO2 in the 2 groups at different time intervals throughout the 24 hours study period (p>0.05).

Aswini A et al, in 2009. opined that there were no significant changes in pulse rate, BP and RR in either group throughout post operative period.

Adverse Effects
Sedation
It was the main side effect group A which constituted 45% compared to B group (12%). Majority of the patients had mild sedation, patient sedated but arousable. This was statistically significant.

Catherine O Hunt in his study has reported a higher incidence of sedation with epidural Butorphanol and is a dose dependent side effect.

Pruritis
In our study none of the patients in group A had pruritis and 4 patients (16%) in group B had pruritis which was statistically significant (p<0.05).

Premila Malik, Chhavi Manchanda, Naveen Malhotra in 2006 showed that pruritis was higher in epidural fentanyl group than butorphanol group. (p<0.05).

Nausea and Vomiting
In our study 2 patient in group A had nausea-vomiting whereas in group B 3 patients had nausea-vomiting which was significant statistically (p<0.05).

No patients on epidural Butorphanol had nausea or vomiting in study conducted By Catheline O Hunt et al. Premila Malik, Chhavi Manchanda, Naveen Malhotra in 2006 showed that the incidence of nausea and vomiting was higher in Fentanyl group than butorphanol group.

Respiratory Depression and Hypotension
In our current study, in group B 2 patients had respiratory depression and 1 patient had hypotension and in none of the patients in group B had hypotension or respiratory depression which were not significant (p>0.05).

No patients had respiratory depression or hypotension with Butorphanol in studies conducted by Catherine O Hunt et al in 1989. CONCLUSION

Epidural butorphanol is safe, effective & dose dependent in providing good pain relief of moderate duration in the postoperative period and is associated with minimal adverse effects. 4 mg is more efficient than 2 mg for postoperative analgesia.

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