A COMPARATIVE STUDY OF ISOBARIC 0.5% LEVOBUPIVACAINE COMBINED WITH 50 MCG FENTANYL VERSUS ISOBARIC 0.5% ROPIVACAINE COMBINED WITH 50 MCG FENTANYL IN LUMBAR EPIDURAL ANAESTHESIA FOR ELECTIVE INFRATEMBILICAL SURGERY

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ABSTRACT

BACKGROUND
This prospective, randomised, double-blinded study compared the onset and duration of epidural anaesthesia produced by levobupivacaine and ropivacaine.

MATERIALS AND METHODS
Seventy adult patients of ASA physical status I and II were divided into two groups of 35 each by pre-decided randomisation schedule, Group R to receive epidural isobaric ropivacaine 0.5% 15 mL combined with 50 mcg fentanyl and Group L to receive epidural isobaric levobupivacaine 0.5% 15 mL combined with 50 mcg fentanyl. A blinded observer evaluated onset and regression of motor and sensory block, and requirement of rescue analgesia.

RESULTS
Onset of sensory block was comparable in both groups. Onset of motor block was longer in group R (18.4+/- 1.77 min. in Group R vs. 15.69 +/- 0.76 min. in group L). In both groups, maximum sensory level reached was T5. Duration of motor block was found to be significantly shorter in group R as compared to group L (175.9 +/- 8.53 min. in Group L vs. 150.9 +/- 7.12 min. in Group R). Duration of analgesia was comparable in both groups (231.0 +/- 9.2195 min. in Group L and 233.54 +/- 8.4064 min. in Group R). Haemodynamic variables were comparable in both groups.

CONCLUSIONS
Levobupivacaine 0.5% produced a sensory block of similar onset, quality, and duration as ropivacaine 0.5% but a longer duration of motor block.

KEYWORDS
Levobupivacaine, Ropivacaine, Lumbar Epidural Anaesthesia, Infraumbilical Surgery.

blockade of 0.75% ropivacaine was comparable to 0.5% bupivacaine.9

Levobupivacaine hydrochloride is the levo stereoisomer form of the racemic form of bupivacaine hydrochloride. Levobupivacaine has demonstrated anaesthetic potency similar to bupivacaine, superior pharmacokinetic profile, lesser cardiovascular and central nervous system toxicity than bupivacaine and better perioperative haemodynamic stability.10 Some studies have shown that, depending on the dose used, levobupivacaine may produce a significantly longer duration of sensory block than bupivacaine.11 On the other hand, some studies showed that levobupivacaine produced same duration of sensory block but less amount of motor block than bupivacaine.12

Levobupivacaine and ropivacaine cause less residual motor blockade compared to bupivacaine. Hence, considering the advent of new drug levobupivacaine in Indian market we proposed to compare levobupivacaine with ropivacaine.

The purpose of this study was to evaluate the duration of analgesia and motor blockade, perioperative haemodynamic effects and side effect profiles of equal dose of isobaric levobupivacaine 0.5% and fentanyl combination versus isobaric ropivacaine 0.5% and fentanyl combination in adult patients undergoing elective infraumbilical surgery under epidural anaesthesia.

MATERIALS AND METHODS
A randomised prospective clinical study of patients undergoing elective infraumbilical abdominal surgeries receiving either epidural ropivacaine or levobupivacaine was undertaken after obtaining written informed consent from the patients and institutional ethics committee approval. For the purpose of sample size calculation, the difference in duration of analgesia was taken as the primary outcome measure. It was calculated that 35 subjects would be required per group in order to detect a difference of 30 min. with 80% study power and 5% probability of Type-I error. The calculation assumed a standard deviation of 45 min. for the duration of sensory block. Sample size calculation was done by n.master-2.0 (Department of Biostatistics, Christian Medical College, Vellore; 2012) software.

Seventy patients were divided into two groups of 35 each by pre-decided randomisation schedule, Group R to receive epidural isobaric ropivacaine 0.5% 15 mL combined with 50 mcg fentanyl and Group L to receive epidural isobaric levobupivacaine 0.5% 15 mL combined with 50 mcg fentanyl. Adult patients aged between 19 and 60 years age and BMI 18.5 to 29.9 kg/meter 2 were included in the study.

Exclusion criteria included patient refusal, infection at the site of injection, coagulopathy, neurological disorders and psychiatric disorder, heart diseases, haemodynamically compromised patients, sepsis, gross anatomical abnormality of vertebral column and known allergy to the study drugs. After preanaesthetic checkup, patients were given tablet 10 mg oral diazepam and tablet ranitidine 150 mg night before operation and kept 8 hours fasting before surgery.

After receiving the patients in operation theatre, an intravenous line was established with an 18G cannula in a large vein of hand and 10 to 15 mL/kg body weight lactated Ringer’s solution administered over 1 hr.

The patients were monitored by NIBP, continuous ECG & pulse oximetry. Under strict aseptic conditions, with the patient in sitting or lateral position skin and subcutaneous tissue was infiltrated with 1% lignocaine (2 mL) at L2-L3 or L3-L4 disc space. Epidural space was identified in the midline with an 18G Tuohy needle using loss of resistance technique with air. Epidural catheter was carefully introduced through the needle. After removal of needle 3 mL of lignocaine 2% with adrenaline was administered through the catheter as a test dose. After 5 min. if there is no evidence of intravascular/intrathecal placement of catheter, 15 mL of test drug was injected in an incremental manner (5 mL drug over 15 secs & interval between each injection 2 min.) such that total injection time will be 4 min. 45 seconds. After completion of injection (time 0), patient was placed supine. Surgery was commenced when the sensory block reached the dermatome level T6.

Sensory blockade was assessed by pinprick using a blunt tipped needle and onset of sensory block (time from epidural injection to the time T 10 blockade was achieved), maximum height reached were noted. It was tested every 1 min. interval till maximum height of block reached and thereafter at 15 min. interval intraoperatively & postoperatively until rescue analgesia required.

Motor block was assessed using modified Bromage scale7 and graded as 0: No motor paralysis; 1: Inability to raise extended leg; 2: Inability to flex knee; 3: Inability to flex ankle. Time for onset of motor block (time from epidural injection to the time Bromage Grade 0 changed to Grade 1), maximum motor block and complete motor recovery noted. It was assessed every 5 min. till onset and every 15 min. thereafter unless restricted access during surgery prevented.

Heart rate, mean blood pressure was recorded every 5 minutes for first 30 min. and then at 15 min. interval till the end of surgery and then at 30 min. interval till rescue analgesia was required. Bradycardia, defined as heart rate (HR) <50/min, was managed by Inj. Atropine 0.6 mg IV bolus (may be repeated). Hypotension, defined by decrease in mean arterial blood pressure more than 30% below baseline or in normotensive patients, fall in systolic pressure below 90 mmHg was managed by Inj. Mephentermine 3-6 mg IV bolus (titrated to patient response) along with fluid bolus (both crystalloids and colloids). We used crystalloids like Ringer’s lactate, 0.9% Normal Saline as intravenous fluid. Obvious side effects like nausea and vomiting, hypotension, pruritus, retention of urine, respiratory depression were monitored.

Analgesia was assessed by VAS pain score which is a linear pain scoring tool ranging from 0 to 10 cm where 0 means no pain and 10 means worst possible pain. Rescue Analgesia was administered postoperatively when VAS score >4 or when patient requested for analgesia. Eight mL of 0.125% racemic bupivacaine was then administered through the epidural catheter. The duration of analgesia was assessed by time to rescue analgesia.

All raw data of study parameters were entered into a Microsoft Excel spread sheet and statistical assessment of the data was carried out using the statistical software Statistica 6.0 [Tulsa, Oklahoma: Stat Soft Inc.,2001] and GraphPad Prism version 5 [San Diego, California: GraphPad Software Inc., 2007]. Kolmogorov-Smirnov goodness-of-fit test was applied to see whether the data distribution is normal. Results were
summarised by descriptive statistics such as mean and standard deviation for numerical variables that are normally distributed and median and interquartile range for those that are skewed. For the variables which showed a normal distribution of intergroup comparisons, independent Two-Sample T-Test was used. For the variables, which were skewed, the Mann-Whitney U test was used. Within group comparisons repeated measurements were performed with the Variance Analysis. Comparisons between two qualitative variables were performed with Pearson’s Chi-Square Test and Fisher’s Exact Test. p value <0.05 was considered to be statistically significant.

RESULTS
Demographic profiles (age, sex, body wt, BMI) and type and mean duration of surgery was comparable in both the groups. (Tables 1,2)

The mean time for onset of sensory block was comparable in both groups (7.57 +/- 0.58 min in group L vs. 7.4 +/- 0.55 min. in group R). Mean time for onset of motor block was longer in group R (18.4 +/- 1.77 min. in Group R vs. 15.69 +/- 0.76 min. in group L) (Figure 1). Peak sensory block height attained was similar in both the groups (T5 -9, T6- 25 in levobupivacaine group and T5- 9, T6- 25 in ropivacaine group).

Modified Bromage scores in both the groups were comparable (2.9 +/- 0.35 in Group L vs. 2.6 +/- 0.94). Duration of motor block as assessed by Modified Bromage score was found to be significantly shorter in group R as compared to group L (175.9 +/- 8.53 min. in Group L vs. 150.9 +/- 7.12 min. in Group R) (Figure 2).

The duration of analgesia was 231.0 +/- 9.219 min. and 233.54 +/- 8.406 min. in Group L and R respectively which showed no statistical significance (p value 0.2321).

Haemodynamic variables were comparable in both groups [Figure 3] and [Figure 4]. There was no incidence of bradycardia in both the groups. There was incidence of significant hypotension in 3 cases of group L compared to 2 in group R which showed no statistical difference. There were no postoperative sequelae like headache, badcach, nausea and vomiting for next 24 h. Two patients in the ropivacaine group and one patient in the levobupivacaine group were excluded from the study due to technical failure of the block.

<table>
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<tr>
<th>SB</th>
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<tr>
<td>L</td>
<td>R</td>
<td>231.0 +/- 9.219</td>
</tr>
<tr>
<td>R</td>
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<td>233.54 +/- 8.406</td>
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Table 2. Sensory and Motor Block Characteristics

<table>
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<th>AGE (yrs.)</th>
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<th>R</th>
<th>P value</th>
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<tr>
<td>39</td>
<td>40</td>
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Table 1. Demographic Profile, Type and Duration of Surgery

Demographic profiles, type and mean duration of surgery was comparable in both the groups.
was associated with relatively longer duration of postoperative analgesia.\(^\text{18}\)

In 2004, Cline et al\(^\text{19}\) compared 0.5% levobupivacaine and 0.5% ropivacaine in combination with 1:200,000 epinephrine for axillary brachial plexus block, and found that sensory analgesia was significantly longer with levobupivacaine than with ropivacaine, but ropivacaine patients showed a faster recovery of motor function. This was similar to our study findings.

Senard et al\(^\text{20}\) in 2004 compared the efficacy, dose requirements, side effects and motor block with epidural infusion of 0.1% levobupivacaine or 0.1% ropivacaine with added 0.1 mg/hour morphine after major abdominal surgery, and showed no differences in quality of pain relief and hourly consumption of the local anaesthetic mixture between the two groups; however, recovery of unassisted ambulation was quicker with ropivacaine than levobupivacaine (76% of patients were able to ambulate on the second postoperative day with ropivacaine versus 48% with levobupivacaine; P < 0.05) which correlated with our study.

The interpretation of the finding for ropivacaine causing a less intense motor block and a more rapid recovery of the sensory and motor functions has been the subject of some controversy. Some have argued that this is a specific drug effect of ropivacaine demonstrating an increased separation of the sensory and motor blocking effects by virtue of a lower lipid solubility whereas others claim that the observed differences are merely due to reduced potency of ropivacaine compared with bupivacaine. In our study, we found patients in the ropivacaine group having a similar duration of sensory block, but motor block was less intense and shorter duration with ropivacaine. This finding appears to add weight to the argument for an increased motor/sensory difference with ropivacaine. If the differences in that study were due just to differences in potency, we would expect parallel differences in the motor and sensory components.

**REFERENCES**


