Epidural steroid injections (ESIs) are commonly used for management of lumbosacral radicular pain. Midline interlaminar (MIL) or transforaminal (TF) routes are commonly used. The TF route, although associated with higher delivery of drug to the ventral epidural space, has serious complications including spinal cord injury and permanent paralysis reported in literature. Therefore, there is a search for a technically better route with fewer complications and greater drug delivery into the ventral epidural space. Recently, a parasagittal interlaminar (PIL) approach has been defined.

Objectives: We conducted this study to compare therapeutic effectiveness of 3 techniques of ESIs in patients having unilateral lumbar radiculopathy. Further, effect of ESI on bone mineral density (BMD) and serum osteocalcin levels were studied.

Study Design: Randomized double-blind trial.

Setting: Pain clinic of a tertiary care hospital.

Methods: Sixty-five patients were randomly allocated into group MIL, group PIL, and group TF to receive epidural injection with 80 mg of methylprednisolone and 2 mL of 2% lidocaine. Effective pain relief and improvement in disability were assessed using Visual Analog Scale (VAS) and Modified Oswestry Disability Questionnaire (MODQ) scores at 2 weeks, 4 weeks, 3 months, and 6 months, respectively. Patients with < 50% relief received additional injection. Primary outcome of study was effective pain relief at 6 months. Mean change in VAS and MODQ scores, BMD, and serum osteocalcin levels were secondary outcome assessed.

Results: Patients having effective pain relief were significantly higher in group PIL (16 of 20 [80%]) and group TF (15 of 20 [75%]) compared with group MIL. Effective pain relief and improvement in disability were assessed using Visual Analog Scale (VAS) and Modified Oswestry Disability Questionnaire (MODQ) scores at 2 weeks, 4 weeks, 3 months, and 6 months, respectively. Patients with < 50% relief received additional injection. Primary outcome of study was effective pain relief at 6 months. Mean change in VAS and MODQ scores, BMD, and serum osteocalcin levels were secondary outcome assessed.

Limitations: The absence of a placebo control group, small sample size, and relatively short follow-up of 6 months were limitations.

Conclusions: PIL approach is equivalent to TF and superior to MIL approach in terms of effective pain relief and decrease in disability in patients with unilateral lumbar radiculopathy. This study showed no deleterious effect on BMD.

Key words: Epidural steroid, technique, efficacy, bone marrow density, serum osteocalcin
Lumbar disc herniation is a common cause of low back pain (LBP) with radicular leg pain (1-3). LBP can be treated using various modalities such as conservative management, epidural steroid injections (ESIs), and surgical interventions. Conservative options begin with neuropathic medications, topical modalities, or physical therapies. ESIs are offered when there is no improvement in pain relief after conservative treatments (4,5). Many studies have shown that large amounts of phospholipase A2 produced due to disc herniation enhances production of prostaglandins leading to inflammation and pain (6,7). This inflammatory response is reduced by ESIs, either by inhibiting the synthesis or release of proinflammatory substances (8).

Epidural space is usually approached using the interlaminar (IL), caudal, or transforaminal (TF) routes (9). Effectiveness of ESIs for pain relief depends on delivery of drug close to the site of pathology (9-13). It is probably because of this reason that the TF technique is associated with greater effectiveness. However, major complications such as spinal cord injury and paraplegia owing to embolization of the artery of Adamkiewicz has been reported with the technique (14-15). Infarction of the spinal cord with TF injection continues to be reported despite negative aspiration of blood and the use of continuous digital subtraction fluoroscopy (16).

Using the IL route, posterior epidural space can be approached between the adjacent spinous process, that is midline interlaminar (MIL), or the lateral most part of the lamina, that is parasagittal interlaminar (PIL) (11,17). Limited effectiveness of MIL ESI has been reported in various studies (9,11). This may be owing to a limited spread of drug to the ventral epidural space (9,11). Current data suggests that the PIL route has a greater effectiveness than the MIL route in terms of ventral spread of the drug, and thereby better pain relief (11).

To date, no randomized clinical trials (RCTs) have compared TF, PIL, and MIL techniques for assessing the effectiveness of ESI in controlling pain in patients with chronic unilateral radicular pain. Therefore, the present study was undertaken to compare the therapeutic effectiveness of these 3 routes for administering ESI in patients with unilateral radicular LBP that has not responded to conservative treatment. Because use of a neuraxial steroid may induce osteoporosis secondary to increased bone resorption and decreased bone formation, effect of steroid given epidurally on bone mineral density (BMD) was also evaluated using dual energy x-ray absorptiometry (DEXA) scan (18,19). We hypothesized that lateral PIL approach would be associated with better clinical outcome as compared with the other 2 approaches. The primary aim of the study was to define the incidence of effective pain relief at 6 months following ESI. Other secondary aims measured were mean change in Visual Analog Scale (VAS) and Modified Oswestry Disability Questionnaire (MODQ) scores over a period of 6 months, contrast medium spread pattern on fluoroscopy, number of ESIs required to achieve VAS score < 50% of baseline, and BMD using DEXA scan at baseline and at 3 months after ESI. Other markers of BMD such as serum osteocalcin levels, serum calcium levels, and vitamin D3 levels at baseline and at 3 months after first ESI were also measured.

**METHODS**

**Study Design**

The study was a prospective, single-center, randomized, double-blind, active-controlled clinical trial. It was conducted in accordance with the Consolidated Standards of Reporting Trials (CONSORT) guidelines (20) and followed the principles of the Declaration of Helsinki. The institutional review board of the Post Graduate Institute of Medical Education and Research, Chandigarh, India approved study protocol and all patients provided written informed consent. Registration with the Clinical trial registry-India occurred on January 12, 2016 (registration number 2016/01/006514).

**Patients**

Adult patients of either gender between the age group of 20 and 50 years, with a diagnosis of chronic lumbar back pain and a unilateral radicular component for at least a 3-month duration not responding to conservative therapies, and having a pain score of at least 50 as assessed on 0 to 100 VAS at baseline were eligible for study recruitment. All patients underwent BMD evaluation, and only those with a T score +1.0 or more of total hip and spine was enrolled. Magnetic resonance imaging was performed to correlate level of herniation with symptoms of patient.

Patients who had surgery on the lumbar spine in the past, lumbar canal stenosis, spondylolisthesis, facet joint arthropathy, allergy to contrast medium or steroid, and bleeding diathesis were excluded. Patients with a history of systemic steroid use, use of
lumbar ESI in the past 6 months, or a history of any
disorder known to affect the bone turnover were also
excluded.

**Randomization and Blinding**

Randomization was performed by an independent
pharmacist using a computer generated randomization
schedule (Software Research Randomizer (Urbaniak, GC
& Plous S 2013, version 4.0), blocks of 6). Random numbers
were kept in opaque, sealed envelopes and opened by
an independent anesthesiologist at the time of injection.
None of the study investigators, including the outcome
assessor, had access to the randomization sequence. ESIs
were performed using one of the 3 approaches: group
MIL ESI using midline IL approach, group PIL ESI using PIL
approach, group TF ESI using TF approach.

All procedures were performed by a single investi-
gator (J.K.M.) and followed by other investigators (B.G.
and K.P.G.). Study cases were kept in between clinical
nonstudy cases during the procedure and for follow-up.
This was done to enhance the blinding and allocation
concealment. Both patients and the investigator assessing
the patient were unaware of the group allocation.

**Study Intervention and Procedure**

After application of standard monitoring, patients
were placed in the prone position. Level of intervention
was determined by clinical presentation and confirmed
using magnetic resonance imaging findings. An initial
anteroposterior fluoroscopic image was obtained, and
level of intervention identified.

In the MIL group, skin at target site was infiltrated
with 2% lidocaine. An 18-gauge, 3.5-inch Tuohy needle
was introduced at the midpoint between 2 spinous
processes and epidural space identified using the loss-
of-resistance to saline solution technique. Final position
of needle in-between the 2 spinous processes was con-
firmed with fluoroscopy before giving contrast medium.

In the PIL group, the needle was introduced into
the most lateral part of the epidural space at target lev-
el, and advanced from posterior to anterior direction.
Parasagittal orientation of the needle was maintained
throughout the procedure.

In the TF group, fluoroscope was rotated through
15 to 20 degrees to get an ipsilateral oblique view until
the superior articular process of the infrasegmental
level was seen at the 6 o’clock position of the target
pedicle. The overlying soft tissue was anesthetized with
1% lidocaine. A 22-gauge spinal needle was then ad-
vanced into the “safe triangle,” inferior to the pedicle

and superolateral to the exiting spinal nerve. Final
needle tip was verified by fluoroscopy as caudad to the
pedicle shadow in anteroposterior and mid or ventral
aspect of foramen in lateral views.

After negative aspiration, 0.5 mL iohexol (300 mg/
ml) (Omnipaque, GE Healthcare, London, United King-
dom) was injected to confirm placement of the needle
in the epidural space. Contrast medium (3.5 mL) was
then further injected to record the pattern of spread of
the contrast medium, and to exclude any intravascular,
subarachnoid, subdural, or intradiscal spread. Contrast
medium was injected under continuous fluoroscopy in
the TF group. Nerve root delineation was confirmed in
the TF group.

Lateral images were used to evaluate ventral epi-
dural spread. Ventral spread was present if contrast
medium was seen hugging the posterior aspect of the
contiguous vertebral body at or above the level of
needle insertion in the lateral image. The perineural
spread was defined as nerve root infiltration of con-
trast medium. Extent of spread of contrast medium
was defined in terms of caudal and cephalic segments
travelled by the contrast medium.

A solution of 80 mg of methylprednisolone acetate
(Depo-Medrol injection, Pfizer Products India Pvt Ltd,
Mumbai, India) with 2 mL of 1% lidocaine (total vol-
ume 4 mL) was then injected.

**Postintervention**

Patients were assessed for neurologic complication
including postural headache, motor weakness, newly
developed pain, paraplegia, and paresthesia.

**Follow-Up**

All patients were followed for a period of 6
months in the pain clinic by a blinded investigator
(B.G. and K.P.G.) who was not aware of the technique
used for performing ESI. Pain relief was recorded using
VAS ranging from 0 to 10 and disability and impair-
ment using MODQ at an interval of 2 weeks, 4 weeks,
3 months, and 6 months. Patients who reported 50%
or less pain relief from baseline after ESI received ad-
ditional injection with the same approach at least 15
days apart with a maximum of 3 injections. Those who
reported > 50% pain relief received further ESIs only
if pain increased to 50% of baseline again. In patients
who developed > 50% pain relief compared with the
baseline, the intervention was defined to be successful.

All patients underwent DEXA scan for BMD at time
of enrollment and at 3 months.
Serum calcium, vitamin D3, and serum osteocalcin were evaluated at baseline and at 3 months.

**Cointerventions**

All patients continued to receive conservative management (pregabalin/gabapentin, nonsteroidal antiinflammatory drugs, and therapeutic exercise program before joining the study). No calcium supplementation was given during the study period.

**Statistical Analyses and Sample Size for the Study**

Statistical analyses were performed using SPSS Version 23 (IBM Corporation, Armonk, NY). As per the previous studies, we calculated that to estimate an effect size of 0.56, we needed at least 20 patients in each group at beta of 90% and alpha error of 0.05 (10,12,14). Mean and medians were calculated for all quantitative variables and measures of dispersion, standard deviation, or standard error were calculated. Normality of data were reviewed using the Kolmogorov–Smirnov test. For normally distributed data, means of 2 groups were compared using the t test. For skewed data, the Mann–Whitney U test was applied. Qualitative or categorical variables were described as frequencies and proportions. Proportions were compared using the chi-square test. For time-dependent repeated variables such as VAS and MODQ scores, repeated measures analysis of variance (ANOVA) was applied. To estimate mean change in VAS and MODQ scores, one-way ANOVA was used followed by Bonferroni correction. Serial values of serum calcium, serum vitamin D3, serum osteocalcin, and DEXA scan were compared using the paired t test.

**Results**

Figure 1 shows the CONSORT of patients. Demographic data, baseline VAS and MODQ scores were comparable between the 3 groups (Table 1). Three hundred and fifty-one patients were screened, and 65 patients were included in the study. There was a loss to follow-up in 4 patients. Data were analyzed for 61 patients: 38 patients had single disc herniation, and 23 patients had disc herniation at 2 levels. The number of patients with one or 2 disc herniation were equally distributed in the 3 groups.

**Primary Outcome**

**Effectiveness of Intervention**

Effectiveness of intervention defined as 50% reduction in VAS scores at 6 months showed a statistically significant difference among the 3 groups ($P = 0.024$; chi-square test). On subgroup analysis, group PIL (16 of 20) and group TF (15 of 20) were significantly more effective than group MIL (12 of 21) ($P = 0.016$ and $P = 0.038$, respectively; chi-square test). However, groups PIL and TF were comparable in effectiveness of intervention ($P = 0.50$; chi-square test) (Fig. 2).

Mean number of injections administered to achieve 50% reduction in pain scores in the 3 groups were similar (group MIL $2.33 \pm 0.65$, group PIL $2.10 \pm 0.64$, group TF $2.05 \pm 0.68$, $P = 0.347$; one-way ANOVA). Percentage of patients who received only one ESI for effective pain relief were 2 of 21 (9.5%) in group MIL, 3 of 20 (15%) in group PIL, and 5 of 20 (25%) in group TF. Additionally, 10 of 21 (47.6%) patients in group MIL, 12 of 20 (60%) in group PIL, and 10 of 20 (50%) in group TF required 2 ESIs for effective pain relief. Percentages of patients receiving 3 ESIs for effective pain relief were 9 of 21 (42.9%) in group MIL, 5 of 20 (25%) in group PIL, and 5 of 20 (25%) in group TF (Fig. 3).

**Secondary Outcome**

**VAS Score**

Repeated measures ANOVA showed significant difference in serial VAS scores between the 3 groups over the follow-up time ($P = 0.00$). We observed that all modalities were effective in each of the groups, as most of the patients reported a significant decrease in VAS score from baseline at all time intervals, that is at 15 days, 1-month, 3-month, and 6-month follow-up (VAS time interaction). VAS scores at 3- and 6-month follow-up showed statistically significant difference in pain scores among the 3 groups (group MIL $3.81 \pm 1.25$, group PIL $2.7 \pm 0.86$, group TF $3.05 \pm 0.75$, $P = 0.002$ at 3 months, and group MIL $3.86 \pm 1.01$, group PIL $2.35 \pm 0.74$, group TF $2.45 \pm 0.51$, $P = 0.000$ at 6 months; one-way ANOVA). Further, patients receiving ESI in group PIL and group TF showed significantly lower VAS scores than group MIL ($P = 0.02$, $P = 0.50$ at 3 months, and $P = 0.00$, $P = 0.02$ at 6 months, respectively; Bonferroni correction). However, group PIL and group TF did not significantly differ in VAS score ($P = 0.799$ at 3 months, and $P = 0.972$ at 6 months; Bonferroni correction) (Fig. 4).

**MODQ**

Repeated measures ANOVA revealed significant MODQ time interaction within the 3 groups, and MODQ group interaction after epidural injection. Statistically
Fig. 1. CONSORT statement.
There was significant difference between percentage of patients demonstrating unilateral or bilateral spread among the 3 groups (P < 0.00; chi-square test). On individual group comparison, TF had a greater number of unilateral contrast medium spread (41 of 41, 100%), followed by group PIL (28 of 42, 66.7%). A significant difference between percentages of patients having unilateral or bilateral spread was observed between groups (MIL and PIL, P < 0.000; groups PIL and TF, P < 0.000; and groups MIL and TF, P < 0.000; chi-square test). Anterior spread was seen in only 51% of ESIs performed in the MIL group as compared with 85.7% in the PIL group (P < 0.000; chi-square test). Further anterior spread was seen in 82.9% of ESIs performed in the TF group (P = 0.002; chi-square test) when compared with the MIL group. However, there was no significant difference observed between percentages of anterior spread of contrast medium between group PIL and TF (P = 0.727; chi-square test).

Perineural spread was significantly different among the 3 groups (P = 0.000; chi-square test). Group TF (85.4%) showed a statistically significant better spread in comparison to groups PIL (57.1%; P = 0.005) and MIL (16.3%; P = 0.000). Further, better perineural spread was observed in group PIL than group MIL (P = 0.000).

The average number of vertebral segments of cephalic spread of contrast medium in group MIL was 2.88 ± 1.092, group PIL was 3.21 ± 1.001, and in group TF was 3.15 ± 0.937. There was no statistically significant difference between means values of cephalic spread between the 3 groups (P = 0.247, one-way ANOVA).

Fluoroscopy exposure time was significantly different in the 3 study groups (P = 0.00; one-way ANOVA). On between-group comparisons, ESIs in group TF took more fluoroscopy time (13.34 ± 1.353 seconds), followed by group PIL (9.10 ± 0.878 seconds). There was significant difference in fluoroscopy time between group MIL and PIL (P = 0.000).

Table 1. Demographic data, baseline VAS and MODQ scores.

<table>
<thead>
<tr>
<th></th>
<th>Group MIL (n = 21)</th>
<th>Group PIL (n = 20)</th>
<th>Group TF (n = 20)</th>
<th>P Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age (years)*</td>
<td>42.71 (± 7.47)</td>
<td>41.15 (± 7.38)</td>
<td>37.65 (± 6.72)</td>
<td>0.08</td>
</tr>
<tr>
<td>Gender</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Male</td>
<td>12 (57.1%)</td>
<td>7 (33.3%)</td>
<td>11 (52.2%)</td>
<td>0.158</td>
</tr>
<tr>
<td>Female</td>
<td>9 (42.9%)</td>
<td>14 (66.7%)</td>
<td>12 (47.8%)</td>
<td></td>
</tr>
<tr>
<td>Height (cm)*</td>
<td>162.62 (± 7.76)</td>
<td>159.05 (± 7.59)</td>
<td>164.45 (± 9.66)</td>
<td>0.126</td>
</tr>
<tr>
<td>Weight (kg)*</td>
<td>67.24 (± 8.78)</td>
<td>63.35 (± 8.00)</td>
<td>66.95 (± 11.16)</td>
<td>0.349</td>
</tr>
<tr>
<td>Body mass index</td>
<td>25.46 (± 3.262)</td>
<td>25.42 (± 2.08)</td>
<td>24.75 (± 3.60)</td>
<td>0.708</td>
</tr>
<tr>
<td>(kg/m²)*</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Baseline VAS*</td>
<td>7.90 (± 0.625)</td>
<td>7.88 (± 0.972)</td>
<td>7.80 (± 0.951)</td>
<td>0.923</td>
</tr>
<tr>
<td>Baseline MODQ*</td>
<td>60.33 (± 2.921)</td>
<td>61.20 (± 4.275)</td>
<td>60.75 (± 3.582)</td>
<td>0.747</td>
</tr>
<tr>
<td>Number of disc</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>herniations</td>
<td>1</td>
<td>13 (61.9%)</td>
<td>14 (70%)</td>
<td>11 (55%)</td>
</tr>
<tr>
<td></td>
<td>2</td>
<td>8 (38.1%)</td>
<td>6 (30%)</td>
<td>9 (45%)</td>
</tr>
</tbody>
</table>
| *- data expressed as mean ± SD

Fig. 2. Bar diagram showing effective pain relief at various time intervals. Compared using the chi-square test; P < 0.05 is significant.

significant difference in MODQ scores were noted among the 3 groups at 3 months (group MIL 41.76 ± 8.64, group PIL 30.65 ± 5.76, group TF 30.75 ± 5.76; P = 0.000) and 6 months (group MIL 37.81 ± 7.21, group PIL 23.70 ± 5.13, group TF 24.15 ± 3.18; P = 0.000) using one-way ANOVA (Fig. 5).

**Contrast Medium Spread Pattern**

A total of 132 ESIs were administered during various time intervals described in the protocol in our study (group MIL 49, group PIL 42, and group TF 41).
and PIL, group MIL and TF, and in group PIL and TF ($P = 0.029$, $P = 0.000$, $P = 0.00$, respectively; post hoc analysis using the Bonferroni test).

There was no statistically significant difference in mean serum osteocalcin levels, serum vitamin D3 levels, and mean DEXA hip and spine t and z scores among the 3 groups at baseline and at 3 months (Table 2).

**DISCUSSION**

The present study was designed to compare the therapeutic effectiveness of 3 routes for administering ESI in patients with unilateral radicular LBP not responding to conservative treatment. Because the use of neuraxial steroid may induce osteoporosis secondary to increased bone resorption and decreased
bone formation, effect of epidural steroid on BMD using DEXA scan was evaluated (21,22). We found that 80% of patients reported effective pain relief in the PIL group, followed by 75% in the TF group, and only 43% in the MIL group. There was a significant reduction in VAS and MODQ scores in all 3 groups as compared with baseline. However, decrease in VAS and MODQ scores were greater in group TF and PIL as compared to group MIL at 3 and 6 months. We did not find any difference in number of injections required in the 3 groups for effective pain relief. The spread of contrast medium on fluoroscopy showed better anterior spread in patients receiving ESI using PIL and TF approach as compared with MIL. Mean serum osteocalcin levels, serum vitamin D3 levels, and mean DEXA hip and spine t and z scores did not differ significantly from the baseline.

Ackerman and Ahmad (9) conducted a randomized study comparing 3 approaches: TF, IL, and caudal ESI in patients with disc herniation. The authors reported...
better pain relief in patients receiving ESI through the TF approach than the IL and caudal approach. This was attributed to placement of drug closer to the site of pathology. In a prospective case-control study, Schaufele et al (17) compared IL and TF approaches with the primary objective of observing improvement in pain. They also reported that TF ESIs were superior to IL in terms of both pain improvement and long-term surgical intervention. In both studies, authors reported better outcomes using the TF approach. However, recent research has concluded that the IL approach is not inferior to the TF approach. Gharibo et al (21) compared the TF and IL approach in patients with LBP due to disc herniations, and reported significant improvement in pain and function with both approaches. The dose of drug used in the 2 groups was different (40 mg triamcinolone in the TF group, and 80 mg of triamcinolone in the IL group).

Second, patients were followed for a short duration of 10 to 16 days only. Rados et al (22) compared the TF and IL approaches in patients with chronic lumbar radiculopathy. The authors demonstrated a significant decrease in VAS scores with both approaches. Again, authors used a higher dose of methylprednisolone in group IL (80 mg) as compared with group TF (40 mg). A total of 53% of ESIs were successful in the IL group, and 63% successful in the TF group. Most of the studies using 80 mg of methylprednisolone in the TF group reported a success rate of 70% to 80% (9,11). It appears that Rados et al (22) probably did not administer equivalent doses in either approaches, and therefore found IL to be as effective as TF.

Although more efficacious, incidence of severe complications in the TF approach are higher. This may be owing to embolization of the artery of Adamkiewicz accompanying the nerve root (15). Some 11.2% of intravascular injections have been reported in the TF approach as compared with 1.9% with IL ESIs (23,24). The PIL approach may provide a suitable alternative to the TF approach in which the drug is deposited close to pathology similar to the TF approach, but at the same time avoids complications associated with TF (10,12-13). Furman et al (25) conducted a pilot study to evaluate the effective pain relief using the PIL approach in patients with lumbar radiculopathy and showed significant improvement in pain scores at 3 months. Because it was a single arm pilot study, further studies addressing this issue are required (25).

Candido et al (10) conducted a randomized study in which IL injection was performed in the lateral most part of the epidural space (i.e., PIL) in one group and TF was performed in the second group. The authors reported comparable VAS scores at all time intervals, that is 2 weeks, 4 weeks, 1 month, 3 months, and 6 months using either approach. Similarly studies conducted by Ghai et al (12) and Hashemi et al (13) also reported comparable pain scores during follow-up using PIL and TF approaches.

Evidence comparing pain relief using MIL with PIL was reported in a single study by Ghai et al (11). The authors found a higher incidence of effective pain relief at 6 months in the PIL group (68.4%) as compared with the MIL group (16.7%). However, the success rate of 16.7% found in the MIL group in the study was lower than that reported in several studies (20,21). We compared the therapeutic effectiveness of the 3 routes for administering ESI in patients with unilateral radicular LBP not responding to conservative treatment and found that the PIL and TF techniques were comparable in effectiveness of intervention.

Lee et al (26) conducted a meta-analysis to investigate efficacy of TF with IL epidural and reported significantly better short-term pain relief. However, quality of evidence for these results were low indicating need of more robust RCTs. Further, authors did not perform a subgroup analysis between MIL and PIL techniques of ESI (26).

Ventral spread of contrast medium on fluoroscopy signifying greater drug deposition was observed in

---

**Table 2. Data comparing biochemical investigations at 3 months with baseline.**

<table>
<thead>
<tr>
<th></th>
<th>Mean ± Standard Deviation</th>
<th>P Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Osteocalcin</td>
<td>Baseline 9.080 ± 3.753</td>
<td>0.077</td>
</tr>
<tr>
<td></td>
<td>3 months 11.877 ± 12.106</td>
<td></td>
</tr>
<tr>
<td>Vitamin D3</td>
<td>Baseline 31.059 ± 0.945</td>
<td>0.437</td>
</tr>
<tr>
<td></td>
<td>3 months 31.5843 ± 0.889</td>
<td></td>
</tr>
<tr>
<td>Serum calcium</td>
<td>Baseline 9.384 ± 0.447</td>
<td>0.279</td>
</tr>
<tr>
<td></td>
<td>3 months 9.430 ± 0.344</td>
<td></td>
</tr>
<tr>
<td>DEXA scan</td>
<td>Baseline –0.266 ± 0.69</td>
<td>0.085</td>
</tr>
<tr>
<td>hip T</td>
<td>3 months –0.310 ± 0.67</td>
<td></td>
</tr>
<tr>
<td>DEXA scan</td>
<td>Baseline –0.185 ± 0.68</td>
<td>0.081</td>
</tr>
<tr>
<td>hip Z</td>
<td>3 months –0.220 ± 0.68</td>
<td></td>
</tr>
<tr>
<td>DEXA scan</td>
<td>Baseline –0.46 ± 0.85</td>
<td>0.086</td>
</tr>
<tr>
<td>spine T</td>
<td>3 months –0.472 ± 0.83</td>
<td></td>
</tr>
<tr>
<td>DEXA scan</td>
<td>Baseline –0.408 ± 0.83</td>
<td>0.601</td>
</tr>
<tr>
<td>spine Z</td>
<td>3 months –0.39 ± 0.82</td>
<td></td>
</tr>
</tbody>
</table>

Data expressed as mean ± standard deviation, analyzed using paired t test. P < 0.05 was significant.
most of the patients in the PIL group (85.7%) and in the TF group (82.9%). The previous data studying contrast medium spread has also reported better ventral spread in PIL and TF approaches similar to our study (10,12).

The use of steroid is associated with many adverse effects (27). Glucocorticoid induced osteoporosis is one of the common adverse effects and varies with preparations, duration, dose, and route of administration. Although enough studies have shown decrease in BMD and decrease in bone quality with use of oral or inhaled steroids, limited studies have evaluated the effect of epidural steroids on BMD.

Manchikanti et al (18) conducted a prospective study to evaluate the effect of neuraxial steroid on BMD. The authors demonstrated no significant change in BMD at 3 months, 6 month, and one year in the group receiving neuraxial steroids as compared with baseline (18). This may be because of the very low doses of steroids used in the study. In the study, betamethasone of 3 to 6 mg (equivalent to 20 to 40 mg of methylprednisolone) was used in the caudal approach, and 1.5 to 3 mg betamethasone (equivalent to 10 to 20 mg of methylprednisolone) for the TF approach. As doses routinely used in epidural space are greater than these, results of the study need to be interpreted with caution. Al-Shoha et al (19) evaluated the effect of ESI on BMD in 1,000 patients. The authors reported a significant mean difference in BMD of the hip at 6 months. Study population consisted of postmenopausal women who were most vulnerable to bone loss, a very well-known fact. Another retrospective analysis reported that ESI caused BMD changes in postmenopausal women not taking antioestoprotic medications (28). However, our study found no significant difference in serum calcium, serum osteocalcin, serum vitamin D3, or DEXA hip and spine score values at 3 months from baseline values.

We did not observe any dural puncture or any spread of contrast medium in subarachnoid space, subdural space, and intradiscal space. No patient developed skin lesions or paresthesia. Postdural puncture headache occurred in one patient only, and responded to conservative treatment.

Even with a midline approach, in the majority of the time, the epidural needle ends up being parasagittal or at least paramedian while performing the procedure. To prevent this, different entry points were targeted for the midline and parasagittal approach. The needle was introduced at the midpoint between 2 spinous processes at the effected level in the MIL approach. Final position of needle in-between the 2 spinous processes was confirmed with fluoroscopy before giving contrast medium. However, in the PIL approach, the needle was introduced into the most lateral epidural space of the effected side.

Our decision to use 4 mL was based on the available literature on ESI in the last 10 years in which maximum studies have used a volume ranging from 3 to 10 mL (9,15,25). Also, Makkar et al (29) found no increase in effectiveness of ESI volume of injectate that was increased from 4 to 8 mL.

All TF procedures were performed under real-time fluoroscopy. Vascular puncture was observed in 2 patients in the TF group and required needle relocation. Intravascular injection can also occur in the epidural space secondary to needle placement in the vertebral venous plexus. As posterior internal vertebral venous plexus attenuates in the midline, incidence of intravascular uptake reported in literature during translaminar injection is as low as 2% (24). It is therefore possible to have a higher incidence of intravascular injection with parasagittal epidural injection secondary to placement of the needle in the lateral epidural vein. We did not observe any vascular injection in group PIL.

Several case reports have documented potential complications such as paraplegia secondary to spinal cord infarction associated with the use of the TF technique (14,23). Incidents have been associated with both safe triangle approach as well as use of particulate steroid. Particulate steroids can cause occlusion of the segmental artery or vertebral artery resulting in cord ischemia (30,31). However, available literature suggests that particulate steroids offer a slightly better VAS score as compared with nonparticulate steroids, and clinicians need to weigh the advantage offered by this difference with the complications reported in the literature with use of particulate steroids (32). To address this issue, a working group constituted under safe use initiative of the US Food and Drug Administration (FDA) issued a warning that injection of corticosteroids into the epidural space of the spine may result in rare, life-threatening complications (33). Manchikanti et al (34) criticized this warning, stating the FDA failed to discuss simple measures to prevent neurologic complications such as initiation of alternate techniques to classic and traditional teachings, avoidance of particulate steroids, use of a blunt needle, and differentiation between different techniques of epidural injection. We used extension tubing to reduce the risk of the needle getting dislodged at the time of change of syringes. All
TF procedures in our study were performed under real-time fluoroscopy.

We used the subpedicular “safe triangle” approach. However, the term “safe” is in reference to the location of the neural but not the vasculature structures. In light of anatomic and radiologic evidence that radicular arteries dwell in the superior part of the foramina, use of alternate techniques have been suggested in literature (35). Many practitioners have proposed adopting the retrodiscal (infraneural) approach with the needle tip positioned into Kambin’s triangle (36,37). Although final position of the needle in this technique might result in less likelihood of radiculomedullary intraarterial cannulation, there may be other risks associated with this needle tip placement. The needle tip may unintentionally be placed too far ventrally and enter the intervertebral disc (38), and risk of subarachnoid or subdural injection may be higher because theoretically, the axillary pouch of the nerve-root sleeve may encroach into Kambin’s triangle. In a retrospective review published by Levi et al (39), the authors reported inadvertent intradiscal injections in 4.7% of patients, and intrathecal injections in 3.1% of patients with use of infraneural/retrodiscal technique. Beyond safety issues in choosing a TF approach, many physicians believe one approach is superior to another for a variety of reasons. At this point in time, it remains unclear if the infraneural or traditional subpedicular technique is superior in effectiveness. Few small trials have investigated this issue without demonstrating any significant clinical difference between the 2 approaches (38,40).

Interventional techniques are one of the commonly used modalities in treating chronic pain, with increasing use and debate to effectiveness. There continues to be a lack of agreement between proponents and opponents on lack of efficacy of certain interventional techniques along with increasing burden of cost (41-45).

RCTs are the gold standard in the evidence-based evaluation of efficacy of intervention modalities/treatment, and are different from other study designs because they are performed under very rigorous conditions. An RCT uses randomization, control group, and double-blind design to minimize bias. There are 2 distinct ways to show efficacy of a particular intervention in an RCT comparison with a (1) placebo control or (2) an active control. RCT with a control assures the investigator that observed treatment effects can be attributed to the intervention, rather than to external factors. No information external to the trial is needed to support the conclusion of effectiveness. However, a study with active control shows that the new therapy is equivalent to or not worse by some defined amount than a known effective treatment (46). Placebo controlled trials are inappropriate in the field of chronic pain, as a delay in treatment increases the disability of the patient and increases the financial burden on society. Active control equivalence trials can be used more appropriately in many systematic reviews and evidence synthesis over single armed trials (47).

There are a few limitations of our study. First, there was no placebo group/control group. This was justifiable because it would be unethical to have a placebo group in which patients are denied pain relief. Second, DEXA scan for BMD was not recorded at 1 and 6 months. Third, we did not study various factors that might have influenced the outcome of our study (48). Our study is limited by a small sample size. For the purpose of the study, only patients with the earlier mentioned fairly strict inclusion/exclusion criteria were enrolled. This eliminates a large proportion of patients typically seen by an Interventional Pain Physician. ESIs are commonly performed for radicular symptoms in patients with multilevel disc herniations, previous blind ESI, and for diagnostic reasons in cases of an unclear diagnosis. It was our goal to limit the study to a well-defined patient population to increase the validity of our results. Finally, we followed our patients for only 6 months. Longer follow-up were desirable to know the long-term effectiveness of the procedure.

Conclusions

The PIL approach is equivalent to the TF approach in terms of effective pain relief and improvement in disability, and superior to the MIL approach in a patient with unilateral lumbar radiculopathy. The PIL approach can be considered as a suitable alternative to the TF approach for administering ESI as both PIL and TF approaches are equally effective, and the PIL approach is associated with fewer complications and technically easy to perform. Low-dose neuraxial steroids are safe in patients with lumbar radiculopathy. This study showed no deleterious effect on BMD.
33. Rathnell JP, Benzon HT, Dreyfuss P, et al. Safeguards to prevent neurologic complications after epidural steroid injections: Consensus opinions from a multidisciplinary working group and


