Comparison of Transforaminal and Parasagittal Epidural Steroid Injections in Patients With Radicular Low Back Pain

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1. Background

Epidural steroid injection (ESI), including transforaminal (TF) epidural injections and interlaminar (IL) epidural steroid injections are commonly performed procedures for the management of lumbosacral radicular pain (1). Although both procedures are minimally invasive and effective (2), there is, however, lack of substantial evidence about the superior efficacy of one of these interventional approaches.

The IL epidural injection can be delivered through midline, paramedian, or parasagittal approaches (3). The TF entry can be directed more closely to the assumed site of pathology in the anterolateral (AL) epidural space, requiring less volume of injection than the IL route. Therefore, TF approach is considered more efficacious than the midline IL approach, probably due to better ventral epidural spread (4). In the meantime, the concerns regarding the safety of the TF approach lead to the search for a technically better route, with lesser complications in the IL approach.

The parasagittal interlaminar (PIL) route could induce higher ventral epidural spread, with fewer complications than TF (5). Although there is limited evidence in the literature showing the effectiveness of PIL in redirecting epidural injection to ventral space, however, it has not been thoroughly studied compared to the TF approach.

2. Objectives

This study aims to compare the effectiveness of PIL and TF ESI in relieving the pain and disability of patients with lumbosacral pain.

3. Patients and Methods

The study was reviewed and approved by the University Review Board of Shahid Beheshti University of Medi-
cal Sciences, Tehran, Iran, and the Ethics Committee of Akhtar Hospital, Shahid Beheshti University of Medical Sciences, Tehran, Iran. Information about the trial was given comprehensively, both orally and in written form, to the parents. All patients gave their written informed consents prior to their inclusion in the study, accordingly.

3.1. Patient Selection
This double blind clinical trial enrolled patients aged between 18 to 75 years, with a diagnosis of low back pain and unilateral lumbosacral radicular pain, due to a contained herniated degenerated disc, with a minimum of 6 months duration, not responding to medications and physical therapies. Magnetic resonance imaging (MRI) was performed to correlate the symptoms and exact disc level protrusion, or contained herniation.

Patients were excluded if they had any clinically significant or unstable medical or psychiatric illness, previous surgery on the lumbar spine, facet joint arthropathy, spinal canal stenosis, unstable neurological deficits, or cauda equina syndrome. Those who had received lumbar ESI in the past, were allergic to corticosteroids, taking anticoagulants or had bleeding diathesis, were taking systemic corticosteroids, pregnant women, or those being ESI injected within 30 days of trial, were also excluded.

Randomization was performed based on accidental numbers assigned by the computer to one of the groups of the study. Study was blinded to both sides, in which neither the patients, nor the physician evaluating, knew the group of study. Patients were given a code by the computer and the code was only known to the physician, who delivered pocket containing group of patient. Then, the code was registered as anonymous to the physician, who performed one of the procedures, based on the label inside the pocket accompanying the patients. After performing the procedure, the patient was evaluated by another physician, who was also blind to the code and group of patients.

3.2. Parasagittal Interlaminar Epidural Steroid Injection Procedure
An 18-gauge, 3.5 inches, Tuohy needle was introduced at the level of disc lesion and advanced in a posterior to anterior direction, vertical to the body surface. After determination of the most lateral place for needle entrance, in fluoroscopy anteroposterior (AP) view, the needle was introduced into the epidural space of the affected side, using the loss-of-resistance and this parasagittal orientation of the needle was maintained throughout the procedure. Bevel direction was positioned toward lateral.

3.3. Transforaminal Procedure
In the TF group, a 22-gauge, 3.5 inch Quincke’s needle was introduced at the level of disc lesion, using first an AP and, subsequently, an oblique orientation (15 - 30°) of the fluoroscopy C-arm to achieve the “Scotty Dog” appearance of the lumbar spine and then directed until the needle tip was in the posterior and superior aspect of the intervertebral foramen, as checked in the lateral imaging, and in line with the pedicle on AP view.

3.4. Confirmation of Epidural Steroid Injection
In both groups, once the needle was in position, and after negative aspiration for cerebrospinal fluid and blood, 1 mL contrast dye (OMNIPAQUE ™, GE Healthcare, Hatfield, UK) was injected to confirm the epidural space distribution in the AP view. This was followed by further injection of 3 mL of contrast under fluoroscopy, to confirm the spread of the contrast, as well as to verify that no contrast medium attained the intravascular, subarachnoid, subdural, or intra-discal spread. Lateral images were taken to evaluate the ventral epidural space. Ventral spread was defined as present if contrast travelled along the posterior longitudinal ligament or adjacent to the posterior aspect of the contiguous vertebral body at the level of needle insertion. Perineural spread and segmental spread were also noted on AP view. After epidural space confirmation, 2 mL of triamcinolone (1 mL = 40 mg), plus 2 mL of bupivacaine (2 mL = 10 mg) and 6 mL sterile normal saline were injected. All the patients were monitored for at least 30 minutes after procedure.

3.5. Data Recording
Sixty four patients were randomized to receive fluoroscopically guided epidural injection, through either the PIL or TF approach. Patients were evaluated for effective pain relief [numeric rating scale (NRS) < 3] by 0 - 10 NRS and functional improvement by Oswestry Disability Index (6) (ODI < 20%) at 2 and 4 weeks. Any complications or side effects were recorded.

4. Results
In this prospective cohort of 64 patients with chronic low back pain, 32 patients received TF epidural injection and 32 received epidural through PIL. Demographic characteristics are listed in Table 1. There were no significant differences in age, sex, body mass index (BMI), duration of pain, pre-procedure pain score (NRS), and functional status (ODI), prior to procedure between the two groups (P > 0.05) (Table 1).

4.1. Effective Pain Relief
Effective pain relief (NRS < 3) was observed in 77.3% (95% CI: 67 – 90.5%) of patients in the parasagittal group and 74.2% (95% CI: 62.4 – 89.4%) of patients in the TF group (P = 0.34), at 4 weeks (Figure 1). Besides, mean NRS pain score was also compared between the two groups. Mean NRS was not significantly different in the two groups, prior to epidural injection (P > 0.05). Mean NRS score was not significantly different in PIL group, compared to TF, at 4 weeks (P = 0.19) (Figure 1).
Table 1. Demographic Characteristics of the Patients

| Demographic Characteristic       | Parasagittal b | TF b | P Value |
|---------------------------------|----------------|------|---------|
| Age, y                          | 49.2 ± 15.5    | 50.5 ± 16.6 | 0.32    |
| Gender (Male/Female)            | 19/13          | 20/12 | 0.52    |
| BMI, kg/m²                      | 22.1 ± 3.4     | 21.7 ± 3.1 | 0.17    |
| Duration of pain, mo            | 8.7 ± 6.5      | 8.7 ± 7.9 | 0.67    |
| Pre-procedure NRS               | 7.1 ± 2.5      | 7.5 ± 2.4 | 0.13    |
| Pre-procedure ODI, %            | 55.2 ± 12.4%   | 53.6 ± 16.1% | 0.082  |

Abbreviations: BMI, Body Mass Index; NRS, Numeric Rating Scale; ODI, Oswestry Disability Index.

Figure 1. Comparison of the Proportion of Patients Achieving Effective Pain Relief in Parasagittal and Midline Epidural Injection

The NRS < 3 at 4 weeks after PIL or TF epidural injection (left figure); Mean pain score (NRS) during follow-up time between two groups of PIL and TF epidural injection (right figure) and Abbreviations: NRS, numeric rating scale; PIL, parasagittal interlaminar; TF, transforminal.

Figure 2. Comparison of the Proportion of Patients Improving Disability Index

The ODI was < 20%, at 2 weeks between the two groups of parasagittal and transforminal epidural injection (left figure); Mean ODI score during the follow-up time between the two groups of parasagittal and midline epidural injection (right figure).

4.2. Functional Improvement During Follow up Time

Number of patients with improved disability (measured by ODI < 20%) was not significantly different in parasagittal group (78% of cases), compared to the TF group (76% of cases), at 4 weeks (P = 0.21) (Figure 2). Besides, mean ODI score was also compared between the two groups. Mean ODI was not significantly different in the PIL compared to TF group, at 4 weeks follow up (P = 0.15) (Figure 2). There was no adverse effect observed in any of our patients.
5. Discussion

Intervertebral disc herniation, spinal stenosis, intervertebral disc degeneration, and failed back syndrome are the most common diagnosis of chronic radicular low back pain. Epidural injections are one of the most commonly performed interventions in managing chronic low back pain.

Among various approaches for ESI, TF is considered as target specific and more effective, compared to midline IL ESI (7). This may be due to blind administration of IL or needle placement in the dorsal space, under fluoroscopic guidance, leading to distribution of the drug to the dorsal space, rather than ventral space (8). Botwin et al. (9) evaluated lumbar IL epidural injections in epidurography pattern. They showed that dorsal contrast of flow occurred in 100% of injections; however, ventral spread of the contrast was seen only in 36% of the patients. In another study, Choi and Barbella (10), in an evaluation of contrast patterns of IL epidural injections, showed excellent spread of contrast into the nerve root and the ventral epidural space in all patients, utilizing a paramedian approach.

The advantage of TF over midline IL injections is attributed to the enhanced deposition of medication in ventral epidural space, close to the source of pain, with a smaller dose of medication (11). There is evidence suggesting that TF allows for greater ventral epidural spread of corticosteroid (12), and ventral epidural spread of corticosteroid has been associated with higher pain and functional improvements.

In our study, there was no significant difference in pain score and functional disability, after 4 weeks of follow up, between PIL and TF epidural injections. This indicates that, in both approaches, the drug is able to reach the ventral space. The existing data suggests long-term efficacy benefits are greater for TF, compared to IL (13-15). However, it remains unclear if TF ESI result in clinically or statistically significant improvements in pain and functional outcomes, compared to IL (16).

Although we did not observe any complications of TF or PIL ESI, however, other literature reviews have indicated that TF are more often implicated in complications, compared to IL ESI, including intravascular injection in up to 23% of lumbar epidural injection cases (17), which can lead to spinal cord infarction and paralysis. A meta-analysis (18) showed that TF resulted in better short-term pain improvement and fewer long-term surgical interventions than midline IL ESI. However, TF injection complications risk must be taken into consideration (19). There have been reports of pneumocephalus during TF. The complication of dural puncture is documented in the context of a lumbar TF (20). The incidence of vascular penetration, during contrast confirmed fluoroscopically guided TF epidural injections have been reported to 8.9-21.3%, depending on the level of injection (21). Previous study demonstrates a high incidence of intravascular injections in TF lumbosacral epidural injections (22). Even in severe cases, studies have presented a case of quadripareisis and brainstem herniation after selective cervical TF (23). The TF, compared to IL ESI, are associated with a 12-fold increased risk of intradiscal injection (24), which can potentially weaken the disc or lead to discitis (25). Other methods, such as intradiscal ozone injection, have been shown to have a positive effect on disk herniation (26).

In conclusion, PIL epidural injection is as effective as TF epidural injection in improving pain and functional status in patients with chronic lumbosacral low back pain, due to disc degeneration. Parasagittal approach holds the advantage of avoiding the risk of complications associated with the TF approach.

Authors' Contributions

Masoud Hashemi suggested the study design. Mohamad Reza Aryani and Gholamreza Mohseni performed data collection, data analysis and interpretation, statistics, Sirus Momenzadeh and Sajad Razavi performed critical revision and manuscript approval, Seyed Amir Mohajerani performed manuscript writing.

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