Comparison of the Therapeutic Effect between a Transforaminal along with a Caudal Epidural Injection, as Well as Two-Level Transforaminal Epidural Injections in a Radiculopathy Patient
요천추병증에 대한 미골부 경막 외 스테로이드 주사의 기존 주간공 경막 외 스테로이드 주사와의 치료 효과 비교

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Purpose: The aim of this study was to evaluate the therapeutic effect of a transforaminal epidural steroid injection (TFESI) along with a caudal epidural steroid injection (ESI), compared to two-level TFESIs in a multi-level radiculopathy patient.

Materials and Methods: A total of 895 lumbar ESIs were performed in 492 patients with multi-level radiculopathy from January 2012 to January 2015. Before injections were performed, the initial Numeric Rating Scale (NRS) score was assessed in all patients, categorized into no pain (excellent), mild (good, NRS: 1–3), moderate (fair, NRS: 4–6), and severe pain (poor, NRS: 7–10). Therapeutic effects were examined for two groups: one-level TFESI along with caudal and ESI two-level TFESIs. Patient outcomes were assessed by NRS in a serial follow-up at one, three, and six months.

Results: One TFESI along with caudal ESI was performed in 274 patients and two TFESIs for 218. For the former group with one TFESI along with caudal ESI, excellent results were shown: 219 (79.9%) patients after one month, 200 (72.9%) after three, and 193 (70.4%) after six months. In the patient group with two TFESIs (n = 218) the outcomes were also very good: 152 (69.7%) after one month, 131 (60.0%) after three months, and 123 (56.4%) patients after six months. The therapeutic effect of one TFESI along with caudal ESI was better than two TFESIs in for one, three, and six months (p < 0.01).

Conclusion: Transforaminal epidural steroid with caudal epidural injection is a more effective tool for lumbosacral radiculopathy than two level transforaminal injections in multi-level radiculopathy patients.

Index terms
Spine
Low Back Pain
Epidural Injection

INTRODUCTION

Lumbosacral radiculopathy is a common medical and socio-economic problem with a lifetime prevalence estimated to be around 40–60% (1-3). Intervertebral disc herniation and degenerative lumbar spinal stenosis are the two most common causes of lumbosacral radiculopathy (4). Degenerative lumbar spinal stenosis is a common disease in the elderly population, which is rapidly increasing (5). However, in most cases, the initial clinical symptoms and imaging findings are nonspecific, discordant or insufficient for diagnosis (6-9). In addition, some patients complain about multi-level radiculopathy.

Currently, epidural steroid injection (ESI) is used for the treatment of both acute and chronic lower back pain. Its use relieves...
symptoms attributed to lumbosacral radiculopathy. In multi-level radiculopathy patients, therapeutic effect of one level epidural steroid injection has been limited. Thus, we focused on additional epidural injections for treating multi-level herniated disc patients. The objectives of our study were comparing the therapeutic effect and complication rates between two level injections and a one level injection along with caudal injection.

MATERIALS AND METHODS

Patients

Lumbar ESIs were performed from January 2012 to January 2015. Among the patients, 895 consecutive lumbar ESIs in 492 were reviewed. We received approval from Institutional Review Board. Total 895 lumbar ESIs (285 men, 207 women; mean age, 63.9 years) were used for analysis. There is no significant difference in the two groups’ age range. The initial Numeric Rating Scale (NRS) score was assessed in every patient (Table 1), dividing them into no, mild (NRS: 1–3), moderate (NRS: 4–6), and severe pain (NRS: 7–10) categories. We classified the patients into two groups based on the NRS and defined marked improvement with no pain as a satisfactory result for therapeutic effect. Retrospectively, the therapeutic effect difference in the groups was evaluated: one level transforaminal epidural steroid injection (TFESI) along with caudal epidural injection and two level TFESIs for lumbosacral radiculopathy. The inclusion criteria were presence of low back or leg pain, clear radiculopathy, such as evidenced by computed tomography (CT) or MRI, which was identified by radiologic reports, and the presence of follow-up medical records after caudal ESI. Exclusion criteria were unclear descriptions of symptoms, the absence of evidence of radiculopathy on cross-sectional images or of follow-up data. Injections for two groups were performed for patients with multi-level radiculopathy and contralateral mild radiculopathy or chronic back pain. There was no significant statistical difference in two patient groups’ symptoms, who were injected the same steroid dose at each time. Therapeutic effects were analyzed at one, three, and six months after injection in the two comparison groups and assessed by using the NRS.

Table 1. Baseline Characteristic of Enrolled Patients

<table>
<thead>
<tr>
<th>Patients</th>
<th>Two Level Injection</th>
<th>One level + Caudal</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gender</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Male</td>
<td>135</td>
<td>150</td>
</tr>
<tr>
<td>Female</td>
<td>83</td>
<td>124</td>
</tr>
<tr>
<td>Age, yr</td>
<td></td>
<td></td>
</tr>
<tr>
<td>≤ 60</td>
<td>88</td>
<td>124</td>
</tr>
<tr>
<td>&gt; 60</td>
<td>130</td>
<td>150</td>
</tr>
<tr>
<td>Primary NRS</td>
<td></td>
<td></td>
</tr>
<tr>
<td>1–3</td>
<td>18</td>
<td>24</td>
</tr>
<tr>
<td>4–6</td>
<td>40</td>
<td>50</td>
</tr>
<tr>
<td>7–10</td>
<td>160</td>
<td>200</td>
</tr>
</tbody>
</table>

NRS = Numeric Rating Scale

Technique

All therapeutic TFESI were conducted under fluoroscopic guidance by a musculoskeletal radiologist experienced in performing spinal interventions. All the injections were performed as outpatient procedures, after obtaining informed consent. With a patient in susceptible position, the tube was rotated obliquely to ensure the facet joint injection pars interarticularis and a clearly visible for it. Next, the skin and subcutaneous tissues overlying the desired pedicle and nerve root were anesthetized with lidocaine. The needle was then advanced under intermittent fluoroscopy into the neural foramen. If it is not possible to inject in a safe triangle, we had a procedure employing the postero-lateral approach. Serial lateral views were obtained to ensure that the antero-posterior position of the needle tip was appropriate. The needle position was checked through fluoroscopy using injections of contrast materials’ small amounts. Postero-anterior and lateral spot radiographs were obtained to document contrast material distribution in epidural space. The steroid was slowly and gently injected (Fig. 1).

Caudal injection was performed in same manner as above, palpating the sacral cornua and localizing the entry point of the needle on A–P view. Sacral cornua were used as landmarks to identify the sacral hiatus, as the sacral hiatus is bounded bilaterally by them. Then, the needle position was confirmed on the oblique, lateral view at the level S2–3. Nonionic contrast materials were injected to the sacral epidural space; then, a mixture of steroid and normal saline was slowly injected (Fig. 2).

Clinical Assessment

Before the injection was performed, the radiologist recorded the duration of symptoms, level, and cause of nerve root impingement on the CT or MR images and the affected dermato-
Comparison of the Therapeutic Effect between a TFESI along with a Caudal ESI and Two Level TFESIs

Mal distribution on the medical charts. Patient outcomes were assessed at one, three, and six month serial follow-up using a NRS that ranged from 0 to 10, subjectively expressed by the patients for the degree of improvement: excellent, good, fair, and poor. An excellent degree represented marked improvement, much improved (> 3-point NRS decreases), good was improvement but with a lesser effect (1–2 point NRS decreases). A fair degree indicated no interval change since the first injection and a poor one meant worsening of pain during the study.

The patients were not informed about the injection method to ensure a blinded protocol for the study. We defined satisfactory results as patients with a reduction of pain as reflected in NRS and no residual pain. After epidural steroid injection, the above-described one, three, and six month serial follow-ups were conducted at a clinical department, as proposed in the literature and based on the duration of therapeutically effective corticosteroids. Follow-up after ESI was scheduled for one month. At follow-up, the outcome was measured on a four-point patient satisfaction scale and recorded on the medical chart. If there was no pain, observations without repeated ESI were considered. If we decided this, the next follow-up was scheduled for three and six months later. We advised patients to return to our hospital immediately, even before the routine follow up date, if symptoms recurred.

Statistical Analysis

The Student t-test was used to evaluate age differences between two groups. Statistical analysis was also performed using Fisher’s exact test and multiple logistic regression analysis to evaluate differences between the two level TFESIs and one level TFESI subsamples along with caudal epidural injection.

The Fisher’s exact test was used to determine differences in sex, therapeutic effects, and the complication rate between the two patient groups. Multiple logistic regression analysis was also performed to assess therapeutic effects. Analyses were performed for both short- and mid-term therapeutic effects. Analyses took place by using the software program SPSS, version 10.0 (SPSS Inc., Chicago, IL, USA). A p value of less than 0.05 was considered to indicate statistical significance.

RESULTS

274 (55.6%) of 492 patients were treated by one TFESI along with caudal epidural injections; 218 (44.4%) were subjected to two level TFESIs. 219 (79.9%) and they achieved a satisfactory result after one TFESI along with caudal epidural injections (group 1) after one month, and 152 (69.7%) of the 218 patients after two level TFESIs (group 2). After a three month follow-up, 200 (72.9%) of the 274 patients obtained a good outcome after one TFESI along with caudal epidural injections and 131 (60.0%) of the 218 patients achieved a satisfactory result after two level TFESIs. After the six month follow-up, 193 (70.4%) of the 274 subjects as well, after one TFESI along with caudal epidural injections, and the findings were likewise for 123 (56.4%) of the
DISCUSSION

Nonsurgical treatment for spinal stenosis varies, but it includes bed rest, nonsteroidal anti-inflammatory drugs, analgesics, oral administration of corticosteroids, physical therapy, and ESI. According to a cohort study by Simotas et al. (11) of non-surgically treated patients with lumbar spinal stenosis, three years after treatment, nine of the 49 patients had undergone surgical interventions. Twelve of the 40 patients not operated also reported no or only mild pain. The authors concluded that aggressive non-operative treatment for spinal stenosis remains a reasonable option (11).

The indication and efficacy of ESI still remains controversial. This technique is currently used as an intermediate treatment for back pain of various causes and duration. It is not considered curative, but a number of patients have reported long and short-term pain relief (12-14). Many reports have been published about the use of TFESI in patients with chronic lower back pain (15); however, to the best of our knowledge, few reports are available about the outcome predictors of TFESI treatment, especially in terms of the injection level numbers. The therapeutic effect of conventional TFESI is limited to multi-level herniated disc patients, because it is a target-specific treatment for leveled originated radiculopathy. Caudal injections are performed for multi-level disc patients and—previously—two level injections did not achieve a sufficient therapeutic effect.

Caudal injection is designed for maximizing the therapeutic effect and to cover broad disc pain. Lee et al. (16) reported that approximately 85% of patients showed improvements after an initial caudal ESI and some 55% displayed excellent amelioration after a series of caudal ESI. According to a study by Delport et al. (17), of the 140 participants who underwent TFESI or caudal injection, 32% reported more than three months of pain relief, 39% less than that, while 29% stated that there was no relief. Manchikanti et al. (18) stated that significant pain relief (≥ 50%) was demonstrated in 55–65% of the patients with spinal stenosis after the use of caudal injection. While Botwin et al. (19) had published that 65% of patients after six weeks, 62% after six months, and 54% after 12 months had achieved a successful outcome, reporting an at least over-50% reduction in visual analog pain scores after caudal injection, but our results appear more optimistic. Fukusaki et al. (20) reported that ESI had no beneficial effect on the pseudoclaudication associated with spinal canal stenosis. On the other hand, Kim et al. (21) could demonstrate that fluoroscopy-guided intra-articular facet joint injection exhibited excellent immediate effectiveness and good prolonged pain relief (> two months) in patients with chronic low back pain. According to our study, one TFESI along with caudal epidural injection was more effective than two level TFESIs. This finding supports the idea that the former can more directly to reduce inflammation and relieve pain.

Our study also has several limitations. First, it was not a prospective study and follow-ups was not long-term designed, although it is difficult to organize a regular schedule for TFESI. We think that it is better for patients to receive TFESI when symptoms appear or worsen. We still believe that the midterm follow-up is sufficient to evaluate the therapeutic effects. Second, initial patient groups had severe back pain. Up to 80% patient’s initial NRS scores were 10 points. Thus, the injection effect appears to be overestimated. We trust that a prospective analysis is the next logical step to further analyze such findings. Furthermore, there was no significant difference in therapeutic effect by symptom duration at a short-term follow-up.

In conclusion, the transforaminal epidural steroid with caudal epidural injection is a more effective tool for multi-level ra-
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diculopathy and contralateral mild radiculopathy or chronic back pain than two level TFESIs.

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요천추병증에 대한 미골부 경막 외 스테로이드 주사의 기존 추간공 경막 외 스테로이드 주사와의 치료 효과 비교

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목적: 여러 층 신경근증 증상을 가진 환자군에서 추간공 경막 및 미골부 경막 외 스테로이드 주사를 함께 시행했을 경우와 두 층 추간공 경막 외 스테로이드 주사를 시행하였을 때의 치료 효과의 차이를 알아보고자 한다.

대상과 방법: 2012년 1월에서 2015년 1월 사이 492명의 여러 층 신경근증 증상이 있는 환자를 대상으로 하였다. 주사 전, 모든 환자에서 초기 통증평가척도(Numerical Rating Scale: 이하 NRS)를 측정하여 경미한(NRS: 1~3), 중등도의(NRS: 4~6), 그리고 심각한(NRS: 7~10) 정도로 나누었다. 환자들은 두 레벨의 경막 외 스테로이드 주사, 그리고 한 레벨의 경막 외 스테로이드 주사와 함께 미골부 경막 외 스테로이드 주사를 병행한 두 가지의 환자군으로 나누어 치료 효과를 비교하였다. 각각의 환자군은 시술 후 1, 3, 6개월에 통증평가척도를 통해 치료 반응을 평가하였다. 평가는 환자의 시술 후 주관적인 통증 해소 정도에 따라 ‘훌륭한, 좋은, 보통의, 악화된’의 4단계로 나누어 평가하였다.

결과: 274명에서 추간공 경막 외 스테로이드 주사와 더불어 미골부 스테로이드 주사를 받았고, 218명에서 두 층 추간공 경막 외 스테로이드 주사를 시험받았다. 추간공 경막 외 스테로이드 주사와 더불어 미골부 스테로이드 주사를 시행받은 274명의 환자 중, 시술 후 1개월 후에 219(79.9%)명이, 3개월 후에 200(72.9%)명이, 그리고 6개월 후에 193(70.4%)명이 ‘훌륭한’ 정도의 치료 반응을 보였다. 두 층 추간공 경막 외 스테로이드 주사를 시행받은 218명의 환자 중, 시술 후 1개월 후에 152(69.7%)명이, 3개월 후에 131(60.0%)명이, 그리고 6개월 후에 123(56.4%)명이 ‘훌륭한’ 정도의 치료 반응을 보였다. 추간공 경막 외 스테로이드 주사와 더불어 미골부 스테로이드 주사를 병행한 경우가 더 나은 치료 효과를 보이는 것이 확인되었다.

결론: 여러 층 신경근증 증상을 가진 환자군에서 추간공 경막 및 미골부 경막 외 스테로이드 주사를 함께 시행하였을 때의 치료 효과가 두 층 추간공 경막 외 스테로이드 주사보다 더 효과적이다.

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