Comparison between transforaminal lumbar interbody fusion and posterior lumbar interbody fusion in treatment of lumbar spondylolisthesis

Shao-Yu Han1*, Quan Xiao2*, Guo-Tai Zhu3, Jian Dai3, Xiao-Ming Tang3, Hai-Lang Sun3

1 Third Department of Orthopedics, Huaiyin Hospital, Huai’an 223300, Jiangsu, China; 2 Department of Orthopedics, Lianshui People’s Hospital, Huai’an 223400, Jiangsu, China; 3 Department of Orthopedics, Huai’an First People’s Hospital, Huai’an 223001, Jiangsu, China. *Co-first authors.

Received September 11, 2015; Accepted January 30, 2016; Epub February 15, 2016; Published February 29, 2016

Abstract: The aim of this study was to compare the clinical efficacy between transforaminal lumbar interbody fusion (TLIF) and posterior lumbar interbody fusion (PLIF) in treatment of lumbar spondylolisthesis. A total of 62 patients with lumbar spondylolisthesis were treated at our hospital from 2010 to 2013. These patients were divided into a TLIF group that included 36 patients (mean age: 60 years) and a PLIF group that included 26 patients (mean age: 57 years) according to different surgical methods. For example, the pedicle screws were fixed first and then transforaminal lumbar interbody fusion or posterior lumbar interbody fusion was performed, and the pedicle screws or intralaminar spreading device were used to distract. Next, the surgical durations, the volume of bleeding during surgery, postoperative drainage, and complications were compared between the two groups. All 62 patients were well operated, 2 patients developed a dural sac tear, and 1 patient had injury to the nerve roots in the PLIF group. The average follow-up duration was 20 months and 18 months in the TLIF group and PLIF group respectively. Both TLIF and PLIF technologies could effectively decompress the interbody fusion and fix posterior endplates steadied centrum, which alleviated clinical symptoms, but TLIF had some advantages over PLIF, such as smaller trauma, low incidence rate of nervous injury or dural sac injury, and better protection of end structures.

Keywords: Lumbar spondylolisthesis, pedicle screws fixation, transforaminal lumbar interbody fusion, posterior lumbar interbody fusion

Introduction

Lumbar spondylolisthesis (LSL) is a common disease and causes lumbosacral pain [1-5]. The pathological mechanisms of LSL are very complicated; furthermore, the duration is long and the efficacy of treatment of LSL is uncertain. Conservative treatment is not effective and patients develop progressive neuropathic dysfunction. Spondylolisthesis fusion is the gold standard treatment to be performed [2, 3]. Pedicle screw fixation combined with grating fusion has been acknowledged widely by many experts, but the type of grating fusion used is still controversial. This study was a retrospective analysis to compare the clinical efficacy of treatment of LSL through transforaminal lumbar interbody fusion (TLIF) and posterior lumbar interbody fusion (PLIF).

Materials and methods

General data

A total of 62 patients with LSL, including 36 men and 26 women, aged 45 to 76 years, with an average age of 50 years, were enrolled in this study from March 2010 to March 2013. The duration ranged from 9 months to 20 years, with an average of 48 months. All patients were divided into the TLIF group, which included 20 men and 16 women, aged 44 to 74 years with an average of 60 years; and the PLIF group that included 16 men and 10 women, aged 45 to 72 years with an average of 57 years.

The TLIF group had 16 patients with lumbar degenerative spondylolisthesis and 20 with isthmic spondylolisthesis. The PLIF group had...
Treatment of lumbar spondylolisthesis

Table 1. The general data of two groups

<table>
<thead>
<tr>
<th>Group</th>
<th>Sex</th>
<th>No. Male</th>
<th>Female</th>
<th>Age (years, x±s)</th>
<th>Spondylolisthesis types</th>
<th>Spondylolysis Grade</th>
<th>Spondylolisthesis segment</th>
<th>Follow-up time</th>
</tr>
</thead>
<tbody>
<tr>
<td>TLIF</td>
<td></td>
<td>36</td>
<td>20</td>
<td>16</td>
<td>Degeneration</td>
<td>I</td>
<td>L4</td>
<td>20.81±8.43</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>59.69±8</td>
<td>Isthmus</td>
<td>II</td>
<td>L5</td>
<td></td>
</tr>
<tr>
<td>PLIF</td>
<td></td>
<td>26</td>
<td>16</td>
<td>10</td>
<td>I</td>
<td>I</td>
<td>L4</td>
<td>18.73±8.15</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>57.31±9.04</td>
<td>I</td>
<td>II</td>
<td>L5</td>
<td></td>
</tr>
</tbody>
</table>

T (c²) - 0.222 0.098 0.178 0.066 0.124 0.970
P value - 0.638 0.277 0.894 0.798 0.725 0.336

According to spondylolisthesis segment classification, the TLIF group had 26 patients with grade I and 10 with grade II injuries, whereas the PLIF group had 18 patients with grade I and 8 with grade II injuries.

According to spondylolisthesis classification, the TLIF group had 15 patients with L4 and L5 spondylolisthesis and 21 patients with L5-S1 spondylolisthesis, whereas the PLIF group had 12 patients with L4 and L5 spondylolisthesis and 14 patients with L5-S1 spondylolisthesis. All patients manifested single lumbar spondylolisthesis and had differing degrees of lumbar pain plus low limb symptoms; moreover, all patients were treated for at least six months by conservative therapy that had resulted in a poor outcome. Furthermore, routine examinations such as radiography, computed tomography (CT), and magnetic resonance imaging (MRI) were performed before the surgery. Patients with a history of a lumbar operation, spondylolisthesis over grade II, tumor or malformation were excluded from this study. This study was conducted in accordance with the declaration of Helsinki. This study was conducted with approval from the Ethics Committee of the Huai’an First People’s Hospital. Written informed consent was obtained from all participants.

Operation

Patients underwent general anesthesia and were placed prone on a surgical frame when the surgeon began by making a vertical incision to expose the facet joints and the lordotic position of the lumbar spine. Spondylolisthetic lumbar loosening was seen in patients with isthmic spondylolisthesis and distinguished by rising spinous processes; additionally, zygapophyseal hypertrophy was obviously seen in patients with lumbar degenerative spondylolisthesis. Under the assistance of fluoroscopy, the positions of upper and lower pedicle screws were confirmed and tracing pins were placed. Pedicle screws were then placed in the standard fashion.

In the TLIF group, we adhered to the following procedure: The zygapophysis was removed with the use of an osteotome, and the skin, muscles, and soft tissues were gently retracted to expose the lateral aspect of the spinous process, the lamina, and the facet joint. The thecal sac and traversing nerve roots were mobilized and retracted to the midline, with care taken to protect the dural sac and neural contents with a retractor. Then the interbody disc was exposed.

In the PLIF group, we adhered to the following procedure: The corresponding lumbar segments and ligamentum flavum were removed to expose the thecal sac and traversing nerve roots, which were slightly pulled with a nerve hook to expose the interbody disc. Next, the interbody disc was incised in the two groups, and the vertebral endplates were removed with annular debris. Proper interbody fusion devices

Table 2. Comparisons of postoperation parameters between two groups

<table>
<thead>
<tr>
<th>Group</th>
<th>n</th>
<th>Surgical Duration (min)</th>
<th>Amount of bleeding during operation (ml)</th>
<th>Volume of postoperative drainage (ml)</th>
<th>Complications (n)</th>
</tr>
</thead>
<tbody>
<tr>
<td>TLIF</td>
<td>36</td>
<td>134.17±27.40</td>
<td>246.94±48.33</td>
<td>151.67±46.93</td>
<td>1</td>
</tr>
<tr>
<td>PLIF</td>
<td>26</td>
<td>130.38±30</td>
<td>271.92±42.43</td>
<td>181.15±60.29</td>
<td>3</td>
</tr>
</tbody>
</table>

T (c²) - 0.515 2.111 2.165 1.919
P value - 0.608 0.039 0.034 0.166
were confirmed and broken bones were added into the interbody space as trial bone grating. Finally, screws were fixed and a titanium rod was placed. After they were identified to be without active bleeding, drainage was performed, followed by closing of the incisions.

The Pedicle screw system used was CD Horizon Legacy system (Medtronic), and the interbody fusion instrument used was Capstone lumbar interbody fusion instrument (Medtronic) (Minneapolis, Minnesota, USA).

**Postoperative treatment**

Patients were nursed and drainage tubes were placed for 24-48 h after the surgery. Meanwhile, antibiotics were administrated for 1 day, steroids and dehydrating agents were used for 3 days after the surgery. Sutures were removed 12-14 days after the surgery. Furthermore, patients needed to exercise the lumbar and back muscles. According to the patients’ condition, they tried to walk with a waistband support about 1 week after the surgery. Additionally, postoperative follow-up was performed.

**Assessment**

The operative duration, amount of bleeding during the surgery, and postoperative drainage volume were observed and recorded. The following Imaging evaluations were performed: Interbody fusion was evaluated according to the Simmons Method. There was no bright area around cage on the radiogram, and the cage did not move above six months after the surgery. The angle of the fused segment on the flexion-extension radiograph was less than 5°. Clinical effect was assessed by visual analogue score (VAS) and Oswestry disability index (ODI). The clinical symptoms, physical sign, and sphincter function were assessed before and after the surgery, and during the last follow-up. The improvement rate was calculated according to ODI scores, (preoperative scores-postoperative scores)/preoperative scores × 100%. “Excellent” meant that the improvement rate was more than 75%; “good” meant that the improvement rate ranged from 50% to 74%; “medium” meant that the improvement rate ranged from 25% to 49%, and “poor” meant that the improvement rate was less than 24%.

**Statistical analysis**

Statistical analysis was performed using the SPSS10.0 software. Measured data was presented as means ± SD. Student t test and chi-square test were used to compare data between the two groups. P < 0.05 denoted a significant statistical difference.

**Results**

A total of 62 patients underwent the procedures successfully. Two patients developed dural sac tears, and 1 patient had an injury to the nerve roots in the PLIF group. There was no obvious difference in the general data between the two groups (Table 1). The surgical duration time was of statistical significance between the TLIF group (134.17±27.40 min) and the PLIF group (130.38±30 min) (t = 2.111, P = 0.039). The volume of postoperative drainage was of statistical significance between the TLIF group (151.67±46.93 ml) and the PLIF group (181.15±60.29 ml) (t = 2.165, P = 0.034, Table 2). The follow-up time ranged from 12 to 48 months, with an average of 20 months in the TLIF group and 9-42 months (mean age: 18 months) in the PLIF group. The VAS and ODI scores were not statistically significant at any time between the TLIF group and the PLIF group (P > 0.05, Table 3). The improvement rate and acceptance rate were 75±17% and 88.9%, respectively, in the TLIF group and 70±16% and 84.5%, respectively, in the PLIF group. One fusion device moved without nervous symptoms in the PLIF group, and 1 patient developed fat liquefaction at the site of the incision in the TLIF group. Both were treated according-

<table>
<thead>
<tr>
<th>Group</th>
<th>n</th>
<th>Pre-operation</th>
<th>Post-operation</th>
<th>Last follow-up</th>
<th>Pre-operation</th>
<th>Post-operation</th>
<th>Last follow-up</th>
</tr>
</thead>
<tbody>
<tr>
<td>TLIF</td>
<td>36</td>
<td>7.11±0.95</td>
<td>2.44±1.42*</td>
<td>2.25±1.59*</td>
<td>67.50±13.43</td>
<td>24.50±12.19*</td>
<td>20.56±10.33*</td>
</tr>
<tr>
<td>PLIF</td>
<td>26</td>
<td>7.12±0.91</td>
<td>2.65±1.26*</td>
<td>2.35±1.38*</td>
<td>68.08±13.31</td>
<td>25.15±10.74*</td>
<td>20.69±10.63*</td>
</tr>
</tbody>
</table>

Note: Vs. pre-operation, *P < 0.05.

Table 3. The results of VAS and ODI scores in two groups
Figure 1. Female with the age of 59 years, had lumbar pain for over 10 years, and the lumbar pain worsened and right low limb was numbness and pain six months ago. A-D showed the lumbar anterio-posterior position and flexion-extension X-ray, which indicated the lumbar degeneration and L4 spondylolisthesis forward. E, F showed the lumbar CT, which also noted the lumbar degeneration and L4 spondylolisthesis forward. G, H showed the lumbar MRI, which implied the lumbar degeneration and bulged interbody disc of L1-2, L2-3, and L4 spondylolisthesis forward. I, J postoperative lumbar anterio-posterior position X-ray showed the posterior pedicle screws fixation at L4-5, and L5/L5 right side by interbody rod decompression plus cage grafting, meanwhile, L4 spondylolisthesis reduction was satisfaction and symptoms such as lumbar pain and low limbs numbness were improved. K, L showed the lumbar fixation still existed through X-ray of lumbar anterio-posterior position one year after the surgery and the symptoms were not recurrent.
ly. The pedicle screw system had no loosening or fractures in the two groups, and the interbody fusion rate was 94.4% (34/36) and 92.3% (24/26) in the TLIF group and PLIF group respectively. The average duration of bone grafting fusion was 6.5 months. In addition, Figure 1 showed the typical cases.

Discussion

LSL is a common disease. About 5% patients with lumbosacral pain have LSL, but the cause of LSL remains unclear. Most studies indicate that congenital dysplasia and chronic strain may be the most important factors that cause LSL [4]. Congenital isthmus diastasis or isthmus cracking caused by chronic strain, degenerative intervertebral disc, and unstable intervertebral facet joints cause and increase in lumbar instability and occur in spondylolisthesis between the upper and lower centrum.

Usually, lumbar degenerative spondylolisthesis and isthmic lumbar spondylolisthesis are common cases in clinical practice [3]. Lumbar spondylolisthesis aggravates spinal canal stenosis and compresses nerves so as to present corresponding symptoms. In addition, durative vertebral instability or increase of stress makes corresponding smaller joints wear down and cause hyperplasia. Moreover, many proliferative scars were formed in the area of isthmus diastasis, which aggravates spinal canal and nerve roots stenoses. Furthermore, lumbar spondylolisthesis is generally accompanied by a herniated disk and spinal stenosis, which increases the difficulties of diagnosis and treatment.

Degenerative lumbar vertebrae, discontinuous pedicles, and lumbar spondylolisthesis affect stability of three lumbar spines. Meanwhile, spinal mechanics are changed and the erected body increased stress on the lower lumbar spines, which further causes intervertebral labilization and compensatory hypertrophy that leads to vicious circle [5, 6]. Therefore, better decompression, reduced distraction of lumbar spondylolisthesis and interbody fusions are the basis to treat LSL.

LSL can be treated by surgery or nonsurgical treatment. Generally, surgical treatment is for patients who have found conservative treatment ineffective and have progressive nervous dysfunction [7]. Moreover lumbar instability and spinal canal stenosis cause degenerative spondylolisthesis and isthmic lumbar spondylolisthesis; thus, the treatment principle of LSL presented is that injured spines are reduced, fixated, and a bone graft fusion is performed. Furthermore, the spinal canal and nerve roots should be completely decompressed which makes the bone graft fusion the most important.

The three spines fusion is the key measure that makes up spinal stabilization for spondylolisthesis; meanwhile, spine fusion could prevent breakage of pedicle screws [8, 9]. Spine fusion approaches contained PLF and LIF that included ALIF, PLIF, and TLIF according to bone graft parts [10]. The bone graft fusion parts of PLIF contained the basal part of transverse process and outside of small joints, which is easy to operate and has a big bony mattress with plenty of vessels. However, the exposed wide range of the two sides might cause more bleeding; furthermore, the joint sac and muscles covered the transverse process and zygopophysis so the bone graft fusion bed is not established.

Simple periosteal bone graft fusion might lead to posterior loosening and breakage of pedicle screws [11, 12]. Interbody fusion not only avoids increasing bleeding problems after stripping muscles and establishing a bone graft bed, but also provides plenty of blood support between the maximum distraction and recovery of the intervertebral space height. Moreover, interbody fusion indirectly expands the nerve roots and recovers the spinal physiologic curve. Most of researchers considered that the fusion rate of interbody fusion was higher than posterior fusion [13]. Cheng et al. [14] compared the effect of PLIF and PLF in treatment of LSL and the results indicated that the fusion rate of PLIF was higher than that of PLF, but there was no obvious difference, which was confirmed in a study by Barbanti Brodano et al. [15].

Both TLIF and PLIF are the effective methods to treat lumbar degenerative diseases, including LSL. TLIF and PLIF provide forward support of the spine during the recompression and keep the intervertebral space height intact until the interbody fusion is completed [13, 16]. PLIF is widely used in clinical practice, but the surgical trauma is serious as PLIF destroys most of the posterior structure of the spine. Additionally, it
affects the dural sac and nerve roots. PLIF also acts as a fulcrum for a dural retractor and to preserve a tension band posteriorly. Dural sac formation leads to iatrogenic spinal canal stenosis [13, 16]. In order to reach the aim of distraction, infusion and increase spine stability to preserve more posteriors structures, TLIF technology was termed on the basis of PLIF by Harms in 1982. Harms and Rolinger reported use of a bone graft packed in a titanium mesh that was inserted via a transforaminal route into the disc space, which had similar characteristics, such as small surgical trauma and little bleeding. Additionally, TLIF effectively reduced compression and achieved the better effect of 360 degree fusion for patients with LSL [17].

Xu et al. [18] also concluded that the TLIF procedure was a safe and effective treatment for 60 the Hans with LSL. Furthermore, Kleinsteuck et al. and his team [19] found that decompression with fusion better improved lumbar pain in patients with LDS than with decompression alone, which corroborated with the study of Ha et al. [20]. A study suggested that the TLIF technique had a better treatment for patients with LSL than the PLIF technique [13], in which the results, such as less bleeding and a lower nerve injury rate were also similar with this study. Here, 2 out of 26 patients had dural sac leakage of the cerebrospinal fluid occurred additional treatment. Fortunately, no severe leakage of the cerebrospinal fluid occurred after the surgery, and 1 case occurred of nerve root injury, which was then treated with steroids. After a six month follow-up, the symptoms of this patient improved. Our experience affects the dural sac and nerve roots. PLIF also acts as a fulcrum for a dural retractor and to preserve a tension band posteriorly. Dural sac formation leads to iatrogenic spinal canal stenosis [13, 16]. In order to reach the aim of distraction, infusion and increase spine stability to preserve more posteriors structures, TLIF technology was termed on the basis of PLIF by Harms in 1982. Harms and Rolinger reported use of a bone graft packed in a titanium mesh that was inserted via a transforaminal route into the disc space, which had similar characteristics, such as small surgical trauma and little bleeding. Additionally, TLIF effectively reduced compression and achieved the better effect of 360 degree fusion for patients with LSL [17].

In summary, both the TLIF and PLIF techniques were effective methods to treat LSL and achieve interbody fusion with nerve decompression. Posterior fixation and unfold interbody fusion could achieve distraction, stability, and fusion of the spine. Compared with PLIF, TLIF had some advantages, such as less blood loss, reducing potential nerve injury, and properly preserving the posterior structure of spine.

**Disclosure of conflict of interest**

None.

**References**

Treatment of lumbar spondylolisthesis


