Radiooperative and Patient-Reported Outcomes of Lordotic Versus Non-lordotic Static Interbody Devices in Minimally Invasive Transforaminal Lumbar Interbody Fusion: A Longitudinal Comparative Cohort Study

Michael H. Lawless 1, Chad F. Claus 1, Doris Tong 1, Noah Jordan 2, Amarpal Dosanjh 3, Connor T. Hanson 1, Daniel A. Carr 1, Clifford M. Houseman 1

1. Division of Neurosurgery, Ascension Providence Hospital, Michigan State University, College of Human Medicine, Southfield, USA
2. Surgery, University of Kentucky College of Medicine, Lexington, USA
3. College of Osteopathic Medicine, Michigan State University, East Lansing, USA

Corresponding author: Michael H. Lawless, mike7@vt.edu

Abstract

Introduction

Minimally invasive transforaminal lumbar interbody fusion (MI-TLIF) is increasingly used to treat lumbar degenerative pathology. Its effect on sagittal parameters remains controversial. Static and expandable lordotic interbody devices (cages) were developed to improve segmental and overall lumbar lordosis. This study aimed to compare the radiographic and patient-reported outcomes (PROs) between static lordotic and non-lordotic titanium cages in patients undergoing 1-2 level MI-TLIF for degenerative conditions.

Methods

We reviewed consecutive eligible patients who underwent 1-2 level MI-TLIF (7/2017-11/2019) at a single institution by multiple surgeons. Standing X-rays and PROs were collected at preoperative, 1-month, and 6-month postoperative intervals. Using univariate analyses, we compared the two cohorts regarding confounders, radiographic parameters, and proportions of patients reaching minimal clinically important difference (MCID) for PROs.

Results

One-hundred-twenty-five patients were reviewed. Forty-seven had lordotic and seventy-eight non-lordotic cages. The lordotic cohort was significantly younger than the non-lordotic (55.9 years vs. 60.7 years, p=0.042). The baseline radiographic parameters were not significantly different between cohorts. At the preoperative-6-month interval, the lordotic cohort had significant improvement in lumbar lordosis versus non-lordotic cohort (2.95° ± 7.2° vs. -0.3° ± 7.1°, p=0.024). Both cohorts showed improvement in segmental lordosis, anterior and posterior interspace height, and low subsidence grade with no significant difference between cohorts at all intervals. Overall, 69.1-83.8% of patients achieved MCID in all PROs with no significant difference between cohorts.

Conclusions

The use of a static lordotic titanium cage in 1-2 level MI-TLIF did not result in significantly different radiographic improvements or PROs compared with a non-lordotic cage.

Categories: Neurosurgery
Keywords: tlif, transforaminal lumbar interbody fusion, lordotic interbody device, lordotic cage, minimally invasive, lumbar osteoarthritis

Introduction

Minimally invasive transforaminal lumbar interbody fusion (MI-TLIF) has become a popular and effective technique for the surgical management of focal lumbar degenerative pathology due to shorter hospital stays, quicker return to work, and increased cost-effectiveness compared to a traditional open posterior lumbar interbody fusion [1-3]. Literature supports the maintenance of spinopelvic harmony (pelvic incidence-lumbar lordosis mismatch within 9°) for improved clinical outcomes [4]. There is controversy regarding the effect of MI-TLIF on sagittal parameters due to its indeterminate effect on the thinning of the anterior spinal column [5-7].

Multiple biomedical device companies have developed static and expandable lordotic interbody devices.
(cages) to improve segmental and regional lordosis to combat sagittal imbalance. However, the actual effect of devices remains unproven. This study compared the radiographic and clinical outcomes in patients undergoing 1-2 level MI-TLIF with static lordotic to non-lordotic titanium cages.

We hypothesize that a lordotic cage will lead to a more significant increase in segmental and lumbar lordosis compared to a non-lordotic cage.

**Materials And Methods**

Institutional Review Board (IRB) approval was obtained for the acquisition and analysis of this data. Due to the retrospective nature of this study, consent was not required by IRB. Our study was prepared per the Strengthening the reporting of observational studies in epidemiology (STROBE) guidelines, circa 2014 [8].

We performed a single-center retrospective review on consecutive patients who underwent one or two-level MI-TLIF between July 2017 and November 2019 using lordotic and non-lordotic titanium cages. The lordotic cages' lordosis was 6° or 12°. The degree of lordosis was based on surgeon preference. Both lordotic and non-lordotic cages were non-articulating and from the same manufacturer (Stryker, Kalamazoo, MI). Exclusion criteria included patients undergoing revision surgery, history of prior lumbar fusion, non-degenerative indication, worker's compensation payer status, or automobile accident-related.

**Operative technique**

Patients were placed under general anesthesia, intubated, and rolled prone onto a radiolucent Jackson table (Mizuho, Union City, CA). Patients were prepped and draped in a standard sterile fashion. Bilateral pedicles at operative levels were cannulated with a Jamshidi needle, and Kirshner wires were placed under biplanar fluoroscopic guidance. Fixed tubular dilators were then sequentially dilated on the ipsilateral facet to the patient’s radicular pain and secured to a rigid arm attachment (Medtronic, Minneapolis, MN). A high-speed burr followed by a combination of micro-curettes and Kerrison rongeur were then used to perform a complete facetectomy and partial laminectomy to expose Kambin’s triangle [9]. A thorough discectomy and endplate preparation was completed using a combination of curettes, disc shavers, and pituitary rongeur. Blunt trial distractors were then used to help select an appropriately sized cage. Cancellous allograft was then packed anteriorly into the disc space followed by insertion of the static titanium cage (Stryker, Kalamazoo, MI) anteriorly to engage the apophyseal ring under fluoroscopic guidance. A minimally effective dose of bone morphogenetic protein (BMP), at 1.28 mg, was then placed posterior to the cage, followed by morselized autograft obtained during facetectomy [10,11]. After confirming hemostasis, the tubular dilator was removed, and screws were inserted over guidewires and secured with rods and locking caps. The wound was closed in anatomical layers.

**Clinical and radiographic measurements**

Patient demographics collected included age, sex, and body mass index. For all patients, standing lateral radiographs were obtained preoperatively, at 4 weeks, and 6 months postoperatively. Patient-reported outcomes (PROs) consisted of visual analog scale (VAS) scores, Oswestry Disability Index (ODI), and the 12-item short-form health survey physical and mental component summary (SF-12 PCS, SF-12 MCS) [12-14].

PROs were collected preoperatively, 6 months, and 1 year postoperatively. Primary outcomes included clinical and radiographic outcome measures.

Standing films were evaluated, and sagittal parameters were measured by neurosurgical resident physicians using validated image analysis software (Surgimap, NYC, NY) (Figure 1). Primary radiographic outcome measures included pelvic incidence (PI), pelvic tilt (PT), sacral slope (SS), lumbar lordosis (LL), PI-LL mismatch, segmental lordosis, anterior and posterior interspace height, and cage subsidence. Cage subsidence into the vertebral endplates was graded using a validated scale, with grade 0: 0%-24%, grade I: 25%-49%, grade II: 50%-74%, and grade III: 75%-100% collapse of the level [5].
FIGURE 1: Illustrative Post-Operative Standing Radiograph
Example of 6-month postoperative standing lateral radiograph of one-level lordotic cage demonstrating anterior cage placement and measured sagittal parameters using validated imaging software (Surgimap, NYC, NY).

Statistical analysis
Using previously reported differences in segmental lordosis between static and expandable lordotic cages at 17.3°± 5.4 vs. 20.3° for a power of 80%, a sample size of 102 was calculated [15]. Baseline and demographic characteristics were summarized using descriptive statistics. Paired t-tests were used to compare pre- and postoperative radiographic measures and PROs in the lordotic and non-lordotic cohorts. Minimum clinically important difference (MCID) was calculated for improvement in PRO from preoperative to 6-month and 12-month postoperative intervals between lordotic and non-lordotic cohorts [16]. For patients with two-level procedures, the interspace with the largest pre- to postoperative change was used in analyses. Sensitivity analysis was not used. Statistical significance was set at p < 0.05. Sampling bias was mitigated by using consecutive patients. Relevant confounding variables for PROs were used, including BMI, age, operative levels, sex, index PRO score, and preoperative PRO scores were used in multivariable analyses of the effect of radiographic parameters on PROs to avoid omitted variable bias.

Results
Patient demographics
A total of 176 patients met the inclusion criteria. Ten patients were excluded due to worker’s compensation status or injury secondary to an automobile accident and 41 patients were excluded due to prior lumbar fusion. A total of 125 patients were reviewed. All patients had a degenerative indication for surgery (e.g. spondylolisthesis, recurrent disc herniation). A total of 78 patients had a non-lordotic cage, of which 56 were one-level, and 22 were two-level. Forty-seven patients had a lordotic cage, 44 had a 6° cage, and three had a 12° cage. Thirty-nine of the lordotic cages were one-level, and eight were two-level. The lordotic cohort was younger than the non-lordotic (55.9 years vs. 60.7 years, p = 0.042) (Table 1). There was no significant difference in gender or weight distribution between cohorts (Table 1). In both the lordotic and non-lordotic cohorts, the most common operative levels were L4-5 for one-level surgery and L3-5 for two-level surgery, with no significant difference in the distribution of operative levels between both cohorts (Table 1).
One patient in the lordotic cohort developed an epidural hematoma requiring evacuation on postoperative day 2. Another patient in the lordotic cohort had a durotomy that required repair with dural graft, fibrin glue, and several days of lumbar catheter drainage with no subsequent cerebrospinal fluid (CSF) leak.

Of the included patients, one patient in the non-lordotic cohort did not have a 4-week follow-up standing X-ray. Fifteen patients did not have a 6-month follow-up standing X-ray (11 - non-lordotic; 4 - lordotic).

**Radiographic parameters**

Non-lordotic and lordotic cohorts had similar preoperative radiographic parameter measurements, with no significantly different parameters (Table 2). The average preoperative LL was 52.1° and 50.5°, segmental lordosis 13.7° and 13.2°, PI - LL mismatch 5.3° and 1.6°, PT 20.8° and 18.7°, anterior disc height 10.2 mm and 11.1 mm, in non-lordotic and lordotic cohorts, respectively (Table 2).
|                  | 4 Week       | 6 Month      | ∆ Pre - 6 Month | PI – LL Mismatch | Pelvic Tilt | Sacral Slope | Segmental Lordosis | Anterior Disc Height | Posterior Disc Height | Cage Subsidence |
|------------------|--------------|--------------|----------------|------------------|-------------|--------------|-------------------|----------------------|----------------------|-----------------|
|                  | 50.4 ± 13.7  | 51.2 ± 12.9  | -0.74          | -5.64 - 4.17     | 5.25 ± 12.4 | 3.45         | -1.27 ± 6.26      | 10.2 ± 4.7           | 5.62 ± 2.46         | ---             |
|                  | 51.7 ± 12.8  | 53.7 ± 11.7  | -1.99          | -6.77 - 2.80     | 10.6 ± 8.1  | 1.34         | 0.15              | 15.0 ± 3.4           | 8.62 ± 2.68         | 38 (56.7)       |
|                  |              |              |                |                  | 8.94 ± 7.38 | -0.02        | -0.04             | 14.7 ± 8.0           | 7.75 ± 2.46         | 25 (59.5)       |
|                  |              |              |                |                  | 5.78 ± 4.79 | -1.03        | -3.00             | 3.94 ± 4.28         | 2.12 ± 2.68         |              |
|                  |              |              |                |                  | 19.2 ± 7.7  | 1.06         | -2.10             | 2.27 ± 7.47         | 2.15 ± 6.10         | 4 (6.0)         |
|                  |              |              |                |                  | 6.81 ± 5.13 | -1.35        | -3.93             | 0.05 ± 7.87         | 0.12                 |                |
|                  |              |              |                |                  |              |              |                   |                     | 0                   |                |
|                  |              |              |                |                  |              |              |                   |                     | 1                   |                |
|                  |              |              |                |                  |              |              |                   |                     | 2                   |                |

**TABLE 2: Radiographic Parameters**

P-value < 0.05 considered significant; PI, pelvic incidence; LL, lumbar lordosis; 95% CI, 95% confidence interval. Continuous data are reported as mean ± SD and written in degrees. PI-LL mismatch presented as the absolute value.
At the 4-week postoperative interval averages were as follows in non-lordotic and lordotic cohorts, respectively: LL decreased -1.4° and increased 0.7°, segmental lordosis increased 1.7° and increased 2.3°, PT increased 0.3° and increased 1.4°, PI - LL mismatch were 5.2° and 5.2°, and anterior disc height increased 5.1 mm and 4.5 mm, with no significant differences (Table 2).

The 6-month postoperative interval was as follows in non-lordotic and lordotic cohorts, respectively: LL decreased 0.3° and increased 2.9° (p = 0.023), PT decreased 0.8° and increased 0.5°, PI - LL mismatch was 8.9° and 9.0°, segmental lordosis increased 2.3° and 2.2°, and anterior disc height remained increased at 3.9 mm and 3.6 mm (Table 2). The degree of cage subsidence at 6-months postoperatively did not differ significantly between non-lordotic and lordotic cohorts, with the majority in both cohorts being grade 0 (Table 2).

**Patient-reported outcomes**

For PROs, 119 patients (95%) had preoperative baseline data, 81 patients (65%) had 6-month postoperative data, 70 patients (56%) had 12-month postoperative data. For all PROs, there was improvement observed at 6-month and 12-month postoperative intervals compared to preoperative baseline (Table 3).

|                | Non-Lordotic (n=75) | Lordotic (n=43) | % MCID Non-Lordotic | % MCID Lordotic | p-value |
|----------------|---------------------|----------------|--------------------|----------------|---------|
| N=118          |                     |                |                    |                |         |
| VAS Back       |                     |                |                    |                |         |
| Preop          | 8.29 ± 2.24         | 8.37 ± 2.15    | ---                | ---            | ---     |
| 6 Month        | 3.49 ± 2.71         | 4.36 ± 3.43    | 39 (86.7)          | 21 (60.0)      | .006    |
| 12 Month       | 3.73 ± 3.49         | 2.72 ± 3.06    | 29 (74.4)          | 25 (89.3)      | .128    |
| VAS Leg        |                     |                |                    |                |         |
| Preop          | 7.27 ± 3.27         | 7.58 ± 3.13    | ---                | ---            | ---     |
| 6 Month        | 1.98 ± 2.90         | 3.06 ± 4.08    | 35 (76.1)          | 22 (62.9)      | .196    |
| 12 Month       | 2.70 ± 3.54         | 1.97 ± 2.99    | 27 (69.2)          | 24 (85.7)      | .119    |
| ODI            |                     |                |                    |                |         |
| Preop          | 47.5 ± 14.4         | 49.6 ± 15.8    | ---                | ---            | ---     |
| 6 Month        | 28.6 ± 18.3         | 24.2 ± 21.4    | 28 (65.1)          | 24 (70.6)      | .611    |
| 12 Month       | 22.6 ± 20.2         | 26.1 ± 21.5    | 21 (55.3)          | 17 (60.7)      | .658    |
| SF – 12 PCS    |                     |                |                    |                |         |
| Preop          | 27.6 ± 6.8          | 28.7 ± 7.5     | ---                | ---            | ---     |
| 6 Month        | 35.2 ± 11.1         | 38.5 ± 12.0    | 24 (53.3)          | 24 (70.6)      | .120    |
| 12 Month       | 40.3 ± 11.5         | 39.6 ± 11.6    | 26 (68.4)          | 21 (75.0)      | .560    |
| SF – 12 MCS    |                     |                |                    |                |         |
| Preop          | 48.3 ± 11.3         | 45.9 ± 11.9    | ---                | ---            | ---     |
| 6 Month        | 53.2 ± 9.2          | 51.6 ± 10.6    | 22 (48.9)          | 17 (50.0)      | .922    |
| 12 Month       | 54.2 ± 10.3         | 55.3 ± 7.7     | 20 (52.6)          | 18 (64.3)      | .344    |

**TABLE 3: Patient-Reported Outcomes**

Continuous data are shown as mean ± SD. P-values < 0.05 are considered significant. VAS, visual analog scale; ODI, Oswestry disability index; SF-12, Short Form – 12 Questionnaire;

PCS, physical composite score; MCS, mental composite score; MCID, minimal clinically important difference.
There was a significantly greater percentage of patients that met MCID in the non-lordotic versus the lordotic cohort in VAS back pain score at 6-months postoperatively (86.7% vs. 60%, \( p = 0.006 \)) but not at 12-months postoperatively (74.4% vs. 89.3%, \( p = 0.128 \)) (Table 3). There were no other significant differences in percentage meeting MCID in PROs between cohorts (Table 3).

Multivariable analysis of PROs by radiographic parameters revealed pelvic tilt to be significantly associated with VAS back pain score \((-0.09 \beta, -0.16 \text{ to } -0.03 \text{ 95% CI}, p = 0.003)\) (Table 4). There were no other significant associations identified (Table 4).

### Table 3

| N=105 | B     | 95% CI      | p-value |
|-------|-------|-------------|---------|
| **VAS Back** |       |             |         |
| PI - LL Mismatch | 0.05 | -0.02 - 0.13 | .160 |
| Pelvic Incidence | -0.03 | -0.07 - 0.01 | .096 |
| Pelvic Tilt | -0.09 | -0.16 - -0.03 | .003 |
| Lumbar Lordosis | -0.02 | -0.06 - 0.03 | .406 |
| Sacral Slope | 0.01 | -0.05 - 0.06 | .826 |
| Segmental Lordosis | -0.01 | -0.11 - 0.09 | .834 |
| Anterior Disc Height | 0.03 | -0.20 - 0.26 | .796 |
| Posterior Disc Height | 0.11 | -0.16 - 0.37 | .436 |
| **VAS Leg** |       |             |         |
| PI - LL Mismatch | 0.002 | -0.08 - 0.08 | .960 |
| Pelvic Incidence | -0.02 | -0.07 - 0.02 | .311 |
| Pelvic Tilt | -0.01 | -0.08 - 0.07 | .857 |
| Lumbar Lordosis | -0.01 | -0.06 - 0.04 | .651 |
| Sacral Slope | -0.05 | -0.11 - 0.02 | .130 |
| Segmental Lordosis | -0.05 | -0.14 - 0.05 | .136 |
| Anterior Disc Height | -0.16 | -0.40 - 0.07 | .180 |
| Posterior Disc Height | 0.12 | -0.17 - 0.41 | .422 |
| **ODI** |       |             |         |
| PI - LL Mismatch | 0.07 | -0.31 - 0.45 | .724 |
| Pelvic Incidence | -0.14 | -0.38 - 0.11 | .266 |
| Pelvic Tilt | -0.16 | -0.59 - 0.24 | .405 |
| Lumbar Lordosis | -0.15 | -0.41 - 0.11 | .270 |
| Sacral Slope | -0.12 | -0.42 - 0.18 | .424 |
| Segmental Lordosis | -0.03 | -0.75 - 0.69 | .929 |
| Anterior Disc Height | 0.60 | -0.69 - 1.89 | .365 |
| Posterior Disc Height | 1.69 | 0.18 - 3.20 | .028 |
| **SF - 12 PCS** |       |             |         |
| PI - LL Mismatch | -0.01 | -0.26 - 0.24 | .932 |
| Pelvic Incidence | 0.10 | -0.05 - 0.26 | .200 |
| Pelvic Tilt | 0.29 | 0.01 - 0.56 | .042 |
| Lumbar Lordosis | -0.02 | -0.17 - 0.13 | .767 |
| Sacral Slope | 0.01 | -0.18 - 0.20 | .924 |
### Discussion

This longitudinal cohort study, which included 125 patients undergoing 1- to 2-level MI-TLIF for degenerative indications with a lordotic or non-lordotic static titanium cage, revealed overall no significant differences between the radiographic outcomes or PROs between cohorts.

There are many interbody fusion techniques to treat degenerative disc disease with mechanical instability in the lumbar spine. Transforaminal lumbar interbody fusion has remained a workhorse approach for decades since Harms’s first description in 1998 due to its effectiveness in decompression of the neural elements and stabilization of the lumbar spine [17].

With more recent innovations in minimally invasive approaches, a common criticism of the MI-TLIF approach, particularly in a unilateral approach, is that it is a kyphosing procedure [15]. To combat this, biomechanical device companies have designed a plethora of static, expandable, and articulating interbody devices to improve the segmental and overall lordosis. However, in this study, we found no clinically significant difference between the static lordotic and non-lordotic MI-TLIF titanium cages in one or two-level surgery. This could be related to the limited power of an MI-TLIF on affecting sagittal parameters. This is likely due to the lack of shortening the posterior column via a unilateral facetectomy and limited lengthening of the anterior column without the release of the anterior longitudinal ligament [5].

### Key results

We did find the lumbar lordosis was increased with a static lordotic titanium cage compared to non-lordotic at 6-months postoperatively. We also demonstrated that the lordotic and non-lordotic cage did improve segmental lordosis by, on average, 2.3°. This improvement in segmental lordosis is consistent with previously reported studies [5,15]. However, these findings are discordant as one would expect a consistent effect on the segmental lordosis and lumbar lordosis.

We did demonstrate that a lordotic cage was not associated with an increased incidence of subsidence compared with the non-lordotic cage. There was an overall low degree of subsidence in both cohorts. After radical discectomy and endplate preparation, we routinely pack 15 cubic centimeters of cancellous bone allograft into the interspace to buttress the cage to decrease excessive subsidence. We further take advantage of the increased strength of the vertebral body endplate at the midline and anterior aspect of the apophyseal ring and place the cage at this site [18,19]. By placing the cage as far anterior as the anterior perimeter of the apophyseal ring, we aim to increase the anterior interspace height and subsequently segmental and overall lordosis. Our findings demonstrate that the increase in disc height and segmental lordosis was sustained over time, which has been a criticism of static cages in prior studies [7].
Most patients did not have completely collapsed disc spaces or “vacuum phenomenon,” evidenced by the preoperative disc heights. This indicates that there was at least some maintenance of central intradiscal pressure and thus persevered bony endplate cortical structure in keeping with Wolff’s law [20]. This logical preservation of the endplate may further explain our low degree of subsidence in both cohorts.

It is well established that pelvic tilt > 20° indicates the patient is attempting to compensate for sagittal imbalance, which would increase the workload of the paraspinal musculature, which could increase the VAS back score [4]. The average PT for lordotic and non-lordotic cohorts at the 6-month radiograph were 19.2° and 20.2°, respectively.

Generalizability
Our results are generalizable to middle-aged patients with degenerative lumbar spine conditions undergoing one or two-level MI-TLIF with a static titanium cage. To our knowledge, this is the first study to compare the radiographic and clinical outcomes of lordotic and non-lordotic static titanium cages in MI-TLIF.

Limitations
Due to the retrospective design, our study may be subject to some bias, particularly regarding the clinical outcome, that may not be present in a prospective study. We did not assess radiographic coronal imbalance. However, Daubs et al. demonstrated that only sagittal plane correction was a predictor of ODI improvement in the setting of combined coronal and sagittal deformity [21].

Conclusions
This study demonstrates that the use of a static lordotic titanium cage in 1-2 level MI-TLIF did not result in significantly different radiographic or PROs compared with a non-lordotic cage. Lordotic and non-lordotic cage cohorts had low grades of subsidence and improvement in interspace height, segmental lordosis, and PROs postoperatively.

Additional Information
Disclosures

Human subjects: Consent was obtained or waived by all participants in this study. IRBNet issued approval 1478613-1. Institutional Review Board (IRB) approval was obtained for the acquisition and analysis of this data. Animal subjects: All authors have confirmed that this study did not involve animal subjects or tissue. Conflicts of interest: In compliance with the ICMJE uniform disclosure form, all authors declare the following: Payment/services info: All authors have declared that no financial support was received from any organization for the submitted work. Financial relationships: All authors have declared that they have no financial relationships at present or within the previous three years with any organizations that might have an interest in the submitted work. Other relationships: All authors have declared that there are no other relationships or activities that could appear to have influenced the submitted work.

References
1. Mummaneni PV, Bixson EF, Kereczdus P, et al.: Minimally invasive versus open fusion for Grade I degenerative lumbar spondylolisthesis: analysis of the Quality Outcomes Database. Neurosurg Focus. 2017, 43:E11. DOI 10.3171/2017.5.FOCUS17188
2. Khan NR, Clark AJ, Lee SL, Venable GT, Rosi NB, Foley KT: Surgical outcomes for minimally invasive vs open transforaminal lumbar interbody fusion: An updated systematic review and meta-analysis. Neurosurgery. 2015, 77:847-74. DOI 10.1227/NEU.0000000000000913
3. Parker SL, Mendenhall SK, Shau DN, Zuckerman SL, Godil SS, Cheng JS, McGirt MJ: Minimally invasive versus open transforaminal lumbar interbody fusion for degenerative spondylolisthesis: comparative effectiveness and cost-utility analysis. World Neurosurg. 2014, 82:230-8. DOI 10.1016/j.wneu.2013.01.041
4. Schwab FJ, Blondel B, Bess S, et al.: Radiographical spinopelvic parameters and disability in the setting of adult spinal deformity: a prospective multicenter analysis. Spine (Phila Pa 1976). 2015, 38:E805-12. DOI 10.1097/BRS.0000000000001829
5. Yee TJ, Joseph JR, Terman SW, Park P: Expandable vs static cages in transforaminal lumbar interbody fusion: Radiographic comparison of segmental and lumbar sagittal angles. Neurosurgery. 2017, 81:69-74. DOI 10.1093/neuros/nyw177
6. McMordie JH, Schmidt KP, Gard AP, Gillia CC: Clinical and short-term radiographic outcomes of minimally invasive transforaminal lumbar interbody fusion with expandable lordotic devices. Neurosurgery. 2020, 86:E147-E55. DOI 10.1093/neuros/nyz402
7. Hawasli AH, Khalifeh JM, Chatrath A, Yarbrough CK, Ray WZ: Minimally invasive transforaminal lumbar interbody fusion with expandable versus static interbody devices: radiographic assessment of sagittal segmental and pelvic parameters. Neurosurg Focus. 2017, 43:E10. DOI 10.3171/2017.5.FOCUS17197
8. von Elm E, Altman DG, Egger M, Pocock SJ, Gotzsche PC, Vandekerckhove JP: The Strengthening the Reporting of Observational Studies in Epidemiology (STROBE) Statement: guidelines for reporting observational studies. Int J Surg. 2014, 12:1495-9. DOI 10.1016/j.ijsu.2014.07.013
9. Kamibayashi H, Kasai K, O’Brien E, Zhou L: Transforaminal arthroscopic decompression of lateral recess stenosis. J Neurosurg. 1996, 84:462-7. DOI 10.3171/jn.1996.84.3.0462
10. Lytle EJ, Lawless MH, Paik G, Tong D, Soo TM: The minimally effective dose of bone morphogenetic protein in posterior lumbar interbody fusion: a systematic review and meta-analysis. Spine J. 2020, 20:1286-304. 10.1016/j.spinee.2020.04.012

11. Lytle EJ, Slavnic D, Tong D, et al.: Minimally effective dose of bone morphogenetic protein in minimally invasive lumbar interbody fusions: Six hundred ninety patients in a dose-finding longitudinal cohort study. Spine (Phila Pa 1976). 2019, 44:989-95. 10.1097/BRS.0000000000002995

12. Fairbank JC, Pyenson PB: The Oswestry Disability Index. Spine (Phila Pa 1976). 2000, 25:2940-52. 10.1097/00007632-200011150-00017

13. Ware J Jr, Kosinski M, Keller SD: A 12-Item Short-Form Health Survey: construction of scales and preliminary tests of reliability and validity. Med Care. 1996, 34:220-33. 10.1097/00005650-199603000-00003

14. Heller GZ, Manuguerra M, Chow B: How to analyze the Visual Analogue Scale: Myths, truths and clinical relevance. Scand J Pain. 2016, 15:67-75. 10.1016/j.sjpain.2016.06.012

15. Vaishnav AS, Saville P, McAnany S, et al.: Retrospective review of immediate restoration of lordosis in single-level minimally invasive transforaminal lumbar interbody fusion: A comparison of static and expandable interbody cages. Oper Neurosurg (Hagerstown). 2020, 18:518-23. 10.1093/ons/opz240

16. Parker SL, Adogwa O, Merendenhall SK, et al.: Determination of minimum clinically important difference (MCID) in pain, disability, and quality of life after revision fusion for symptomatic pseudarthrosis. Spine J. 2012, 12:1122-8. 10.1016/j.spinee.2012.10.006

17. Harms J, Jeszenszky D: The unilateral, transforaminal approach for posterior lumbar interbody fusion. Orthop Traumatol. 1998, 6:88-99.

18. Grant JP, Oxlund TR, Dvorak MF: Mapping the structural properties of the lumbosacral vertebral endplates. Spine (Phila Pa 1976). 2001, 26:889-96. 10.1097/00007632-200104150-00012

19. Lowe TG, Hashim S, Wilson LA, O’Brien MF, Smith DA, Diekmann MJ, Trommeter J: A biomechanical study of regional endplate strength and cage morphology as it relates to structural interbody support. Spine (Phila Pa 1976). 2004, 29:2389-94. 10.1097/01.brs.0000143623.18098.e5

20. Benzil EC: Biomechanics of Spine Stabilization. Thieme Medical Publishers, Inc, 2015. 10.1055/b-005-106374

21. Daubs MD, Lenke LG, Bridwell KH, Kim YJ, Hung M, Cheh G, Koester LA: Does correction of preoperative coronal imbalance make a difference in outcomes of adult patients with deformity? Spine (Phila Pa 1976). 2013, 38:476-85. 10.1097/BRS.0b013e5182846e25