Analysis of the Relationship between Ligamentum Flavum Thickening and Lumbar Segmental Instability, Disc Degeneration, and Facet Joint Osteoarthritis in Lumbar Spinal Stenosis

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**Study Design:** Cross-sectional study.

**Purpose:** To investigate the relationship between ligamentum flavum (LF) thickening and lumbar segmental instability and disc degeneration and facet joint osteoarthritis.

**Overview of Literature:** Posterior spinal structures, including LF thickness, play a major role in lumbar spinal canal stenosis (LSCS) pathogenesis [1-3]. The LF covers a considerable part of the posterior and lateral spinal canal (LF) thickness, play a major role in lumbar spinal canal stenosis (LSCS) pathogenesis [1-3]. The LF covers a considerable part of the posterior and lateral spinal canal

**Methods:** We examined 419 patients who underwent computed tomography (CT) myelography and magnetic resonance imaging after complaints of clinical symptoms. To investigate LF hypertrophy, 57 patients whose lumbar vertebra had normal disc heights at L4–5 were selected to exclude LF buckling as a hypertrophy component. LF thickness, disc space widening angulation in flexion, segmental angulation, presence of a vacuum phenomenon, and lumbar lordosis at T12–S1 were investigated. Disc and facet degeneration were also evaluated. Facet joint orientation was measured via an axial CT scan.

**Results:** The mean LF thickness in all patients was 4.4±1.0 mm at L4–5. There was a significant correlation between LF thickness and disc degeneration; LF thickness significantly increased with severe disc degeneration and facet joint osteoarthritis. There was a tendency toward increased LF thickness in more sagittalized facet joints than in coronalized facet joints. Logistic regression analysis showed that LF thickening was influenced by segmental angulation and facet joint osteoarthritis. Patient age was associated with LF thickening.

**Conclusions:** LF hypertrophy development was associated with segmental instability and severe disc degeneration, severe facet joint osteoarthritis, and a sagittalized facet joint orientation.

**Keywords:** Disc degeneration; Ligamentum flavum thickness; Lumbar spine; Facet orientation
walls [4]. As LF thickening can reduce spinal canal diameter, LF thickness has been considered as a cause of LSCS [5]. Canal narrowing mechanically compresses the nerve root or cauda equina [6-9] and contributes to lower back pain and sciatica, even in the absence of a bulging annulus fibrosus, herniated nucleus pulposus, or osseous spurs [10].

LF thickening pathogenesis is unclear, and whether the process is attributable to tissue hypertrophy or buckling remains controversial. Canal narrowing, in part, results from LF hypertrophy. In a normal ligament, elastic fibers are predominant and comprise 60% to 70% of the extracellular matrix [11,12]. In contrast, a hypertrophied LF shows an increased amount of collagen fibers, calcification, ossification, and chondrometaplasia [13-18].

The cause of LF hypertrophy is multifactorial and includes activity level, age, and mechanical stress. LF thickness in LSCS without disc space narrowing may sometimes be detected by radiographic findings. In such cases, LF thickness is not caused by LF buckling, but by LF hypertrophy. Mechanical stress-induced tissue damage might be the triggering event for an inflammatory reaction and subsequent development of tissue scarring [16,19]. The accumulation of such scarring may result in LF hypertrophy [20]. This study aimed to clarify LF hypertrophy pathogenesis associated with mechanical stress by investigating the relationship between segmental instability, disc degeneration, and facet joint osteoarthritis.

Materials and Methods

1. Participants

Prospective participants included 419 patients who underwent computed tomography (CT) myelography and magnetic resonance imaging (MRI) after complaining of clinical symptoms related to the spine, spinal cord, or cauda equina. Patients were evaluated between January 2007 and September 2013. Patients with evidence of bone metastasis, congenital anomalies, scoliosis, discitis, osteomyelitis, spondylolysis, spondylolisthesis, ossification, or fracture were excluded. Furthermore, with the intended purpose of investigating pure LF hypertrophy, 57 patients (13.6%) with a normal L4–L5 disc height were selected. A previous study reported that the disc height for healthy males aged 20 to 30 years was 11.8 mm [21]. Therefore, we selected patients whose disc height was more than two-thirds (8 mm) the normal measurement. The study group comprised 41 men and 16 women with an average age of 63.2 years (range, 22–89 years).

2. LF thickness measurement

Lumbar images were obtained using a 1.5 Tesla magnet (EXCELART Vantage, Toshiba Medical, Tokyo, Japan). LF thickness was measured at L4–5 for 57 patients on axial T2-weighted MRI images at the facet joint level (Fig. 1). Thickness at the middle LF portion was measured. Images were projected using a DICOM viewer. LF thickness was measured using electronic calipers at a DICOM workstation. Measurements were repeated twice by the same orthopedic surgeon; mean values were recorded. In asymmetrical thickness cases, the larger LF thickness value was used.

3. Assessment of lumbar segmental spinal motion

A radiologic assessment of lumbar segmental spinal motion was measured on dynamic film at flexion-extension. Disc space widening angulation in flexion and segmental angulation (angle of extension–angle of flexion) on lateral radiographs of extension and flexion were used to assess radiologic instability at the L4–5 segment. The presence of a vacuum phenomenon in the L4–5 disc and lumbar lordosis at T12–S1 was investigated by lateral radiograph.
4. Disc degeneration assessment

To evaluate vertebral discs, a major component for segmental stability, disc degeneration was classified into five grades according to Pfirrmann’s criteria using T2-weighted sagittal images [22]. The grade was determined by the degenerative status of the nucleus, annulus, and endplate (grade I, normal; grade V, most advanced degeneration).

5. Lumbar facet joint assessment

Facet joint orientation was measured on an axial CT scan. A facet line was drawn between the anteromedial and posterolateral points of the L4–5 facet. A mid-sagittal line, drawn through the disc, was considered the sagittal line. The angle formed by the left or right facet line and mid-sagittal line was measured and recorded in degrees (Fig. 2). Two angles (left and right) were averaged and recorded as facet joint orientation.

Facet joint morphology was classified into four groups according to Imajo et al. [23]: sagittalized facet joints indicated a smaller facet angle, coronalized facet joints were nearly 90°, C-shaped facet joints were angled at about 45°, and J-shaped facet joints were angled at about 30° (Fig. 3). To evaluate facet joint osteoarthritis, osteophyte size at the L4–L5 facet joints was determined and classified into four grades according to Grogan et al. [24]: Grade 1 indicates no osteophyte; grade 2, a mild or possible osteophyte; grade 3, a moderate osteophyte; and grade 4, a large osteophyte. When different grades were assigned to the right and left facet joints in the same segment, the grade for the most severe osteoarthritis was used.

6. Statistical analysis

Statistical analyses were performed with SPSS ver. 18.0 (SPSS Inc., Chicago, IL, USA). Relationships between LF thickness and age, disc widening angulation, segmental angulation, a vacuum phenomenon, lumbar lordosis, facet angle, disc degeneration, and facet joint osteoarthritis were analyzed by Spearman’s correlation. Bonferroni correction was used to examine the relationship between LF thickness and disc degeneration, facet joint osteoarthritis, and facet joint orientation. A logistic regression analysis was performed to test whether each parameter was independently associated with LF thickness. Studies have proposed to set an upper limit for normal LF thickness of <4 mm [25,26]. Therefore, LF thickening was identified when LF thickness was ≥4 mm. Logistic regression was applied.
to determine which variables could predict the risk of LF thickness, with the end point being ≥4 mm. Odds ratios (ORs) and 95% confidence intervals were determined for each predictive variable. Demographics and radiological parameters according to an LF of <4 mm or ≥4 mm were statistically analyzed by the Mann-Whitney U test. The grade system of disc degeneration and facet joint osteoarthritis was compared at <4 mm and ≥4 mm by the chi-square test. Statistical significance was defined as \( p < 0.05 \).

Intraobserver and interobserver agreement were assessed to be good-to-excellent with a kappa statistic of >0.70 for each parameter.

## Results

### 1. Lumbar segmental spinal motion, disc degeneration, and facet joint orientation

The mean disc space widening angulation at L4–L5 was 4.2°±5.2° (range, -10° to +17°). The mean segmental angulation at L4–L5 was 6.8°±3.4° (range, 0° to +15°). The average lumbar lordosis was 45.2°±12.1° (range, +15° to +76°). Fourteen patients (24.6%) had a vacuum phenomenon (Table 1). Disc degeneration was identified as grade II in 4 patients (7.0%), grade III in 20 (35.1%), and grade IV in 33 (57.9%). No patients had grade I or V.

The mean facet angle was 40.9°±8.4° (range, 23.5° to 55.5°), and facet joint morphology was sagittalized in 2 patients (3.5%), coronalized in 3 (5.2%), C-shaped in 36 (63.2%), and J-shaped in 16 (28.1%). Facet joint osteoarthritis was grade 1 in 1 patient (1.8%), grade 2 in 19 (33.3%), grade 3 in 28 (49.1%), and grade 4 in 9 (15.8%). The Bonferroni correction was not valid because the grade 1 group included only one patient.

### 2. Relationships between LF thickness and parameters

The mean LF thickness was 4.4±1.0 mm for L4–L5. LF thickness significantly correlated with age (\( r = 0.55, p < 0.01 \)) and the vacuum phenomenon (\( r = 0.32, p < 0.05 \)) and was independent of disc space widening angulation, segmental angulation, lumbar lordosis, and facet angle (\( p > 0.05 \)) (Table 1).

The grade distribution of disc degeneration was as follows: grade II, was 3.2±0.1 mm; grade III, was 3.9±0.7 mm; and grade IV, was 4.8±1.0 mm. LF thickness was greater in the grade IV group compared with the grade II (\( p < 0.01 \)) and grade III (\( p < 0.01 \)) groups, with a significant positive association between LF thickness and grade IV.

![Fig. 4. Mean ligamentum flavum (LF) thickness of each grade of disc degeneration. The average thickness of grade II was 3.2±0.1 mm, grade III was 3.9±0.7 mm, and grade IV was 4.8±1.0 mm. LF thickness was greater in grade IV than in grade II (p<0.01) and grade III (p<0.01).](image-url)

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**Table 1. Relationships between thickness of LF and parameters**

| Parameter                        | Value   | \( p \)-value |
|----------------------------------|---------|---------------|
| Age (yr)                         | 63.2    | \( S \, (p<0.01, r=0.55) \) |
| Angulation of disc space widening (°) | 4.2±5.2 | NS (\( p=0.82, r=0.31 \)) |
| Segmental angulation (°)         | 6.8±3.4 | NS (\( p=0.60, r=0.07 \)) |
| Lumbar lordosis (°)              | 45.2±12.1 | NS (\( p=0.90, r=0.17 \)) |
| Vacuum phenomenon                | 14/57 (24.6%) | \( S \, (p<0.05, r=0.32) \) |
| Facet angle (°)                  | 40.9±8.4 | NS (\( p=0.35, r=0.13 \)) |

The values are given as the mean and the standard deviation. LF, ligamentum flavum; \( S \), significant; \( r \), correlation coefficient; NS, not significant.
increasing disc degeneration severity (Fig. 4). In addition, LF thickness was significantly correlated with age ($r=0.55$, $p<0.01$). Furthermore, age was significantly correlated with disc degeneration ($r=0.67$, $p<0.01$) and facet joint osteoarthritis ($r=0.56$, $p<0.01$) (Fig. 5). This association between LF thickness and age was mirrored by a significant positive correlation between age and grade of disc degeneration and facet joint osteoarthritis.

Regarding facet joint morphology, the average sagittalized facet thickness was $5.0\pm1.8$ mm; coronalized facet thickness was $3.9\pm0.8$ mm, C-shaped facet thickness was $4.4\pm1.0$ mm, and J-shaped facet thickness was $4.3\pm0.8$ mm (Fig. 6). No statistical significance was noted, but LF thickness was greater for more sagittalized facet joints than more coronalized facet joints.

Thirty-one patients (54.4%) had an LF thickness of $\geq4$ mm and 26 (45.6%) had an LF thickness of $<4$ mm. Table 2 shows demographics and radiological parameters according to an LF thickness of $<4$ mm or $\geq4$ mm. Logistic regression analysis showed that LF thickening was influenced by disc space widening (OR, 0.86; $p=0.033$) and facet joint osteoarthritis (OR, 3.57; $p=0.032$). Patient age was also associated with LF thickening (OR, 1.08; $p=0.041$) (Table 3). Conversely, segmental angulation, the vacuum phenomenon, and lumbar lordosis were not associated with LF thickening. Disc degeneration and facet joint osteoarthritis were the result of aging, and thus lumbar segmental instability occurred because of lumbar disc degeneration.
Ligamentum flavum thickening and lumbar instability

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space widening and sagittalized facet joints.

**Discussion**

The LF covers the posterior spinal canal wall. LF hypertrophy may eventually compress the spinal cord, cauda equina, or nerve roots [6,27-29]. Canal narrowing in LSCS results from degenerative changes in posterior lumbar spine structures, with LF thickening and bony proliferation of the facet joints being major factors in its development [19]. The mechanism of LF hypertrophy from the perspectives of anatomy, histology, and biology has been investigated. Causes of LF hypertrophy include age, spinal level, mechanical stress endured, and growth factors. Histological research data have shown that an increased LF thickness is associated with an increase in fibrosis (i.e., an increase in collagen fiber content) and a decrease in elastic fibers [13,16-18]. Epidemiologically, mechanical stress attributable to spinal instability may initiate LF hypertrophy [16,25,30,31]. However, the mechanism inciting lumbar mechanical stress remains unknown. In our study, we used radiographic measures of L4–5 segmental instability to elucidate the mechanism between mechanical stress and LF thickness.

LF thickening pathogenesis is unclear, and whether LF thickening is due to tissue hypertrophy or buckling remains controversial. The exact function of the LF is also unknown, but it is a connective tissue thought to affect intrinsic spine stability, control intervertebral movement, and maintain a smooth posterior dural sac surface [32].

Using a transmission electron microscopy LF assessment, Postacchini et al. [14] concluded that reduced elasticity

| Parameter                        | <4 mm     | ≥4 mm     | p-value   |
|----------------------------------|-----------|-----------|-----------|
| Age (yr)                         | 56.3±2.9  | 68.9±1.7  | S (p<0.01)|
| Angulation of disc space widening (°) | 5.4±0.8   | 3.2±1.0   | NS (p=0.14)|
| Segmental angulation (°)         | 6.4±0.7   | 7.1±0.6   | NS (p=0.38)|
| Lumbar lordosis (°)              | 45.6±2.6  | 44.9±2.0  | NS (p=0.60)|
| Vacuum phenomenon, n (%)         | 2/57 (4.0)| 12/57 (21.1)| S (p<0.01)|

**Table 2.** Measured parameters of <4 mm and ≥4 mm LF thickness

The values are given as the mean and the standard deviation. Statistical analyses were performed by Mann-Whitney U test. The grade system of disc degeneration and facet joint osteoarthritis were by chi-square test. LF, ligamentum flavum; S, significant; NS, not significant; NA, not available.

| Independent variable                        | p-value | OR   | 95% CI   |
|---------------------------------------------|---------|------|----------|
| Age                                         | 0.041   | 1.08 | 1.00–1.15|
| Angulation of disc space widening           | 0.033   | 0.86 | 0.75–0.99|
| Facet joints osteoarthritis                 | 0.032   | 3.57 | 1.11–11.44|

**Table 3.** Logistic regression analysis for the thickness of LF

LF, ligamentum flavum; OR, odds ratio; CI, confidence interval.
may cause the LF to bulge into the spinal canal, even in the standing position with an LF of normal thickness. In addition, a cause of thickening may be LF buckling into the spinal canal after disc collapse. To evaluate this theory, Altinkaya et al. [33] examined the relationship between LF thickness and intervertebral disc degeneration and disc height; they reported a significant increase in LF thickness with increased disc degeneration. Based on these findings, they hypothesized that spinal segment buckling, secondary to disc degeneration, resulted in LF thickening and buckling into the spinal canal. Conversely, Sakamaki et al. [4] did not identify a correlation between LF thickness and disc degeneration. We investigated patients with preserved disc height because each intervertebral disc has a different segmental mobility. During flexion-extension, there is usually a cephalocaudal increase in the range of motion in the lumbar spine; the L4–5 area bears the highest load and tends to experience the most motion. Our study included 57 patients (13.6%) with preserved L4–5 disc heights; 31 (7.4%) patients had an LF thickness of ≥4 mm, LF hypertrophy, and no buckling. Furthermore, disc degeneration and facet joint osteoarthritis significantly correlated with aging; LF thickness also significantly correlated with disc degeneration and facet joints osteoarthritis. In our study, LF thickness increased with age, supporting previous research suggesting that LF thickness is an age-dependent phenomenon [4,16,34]. Thus, our results indicate that thickening is not necessarily LF buckling into the spinal canal associated with disc degeneration.

Normal LF is composed of 80% elastic fibers and 20% collagen fibers [35]. Upon hypertrophy, LF shows a loss of elastic fibers and an increase in collagen fibers, resulting in fibrosis [17,20]. Previous histological data indicated that LF hypertrophy was characterized by fibrosis, loss of elastic fibers, increased collagen fibers, calcification, degeneration of elastic fibers, and chondrometaplasia [13,16,18-20]. These histological changes are similar to tissue scarring during the post-inflammatory repair process in other organs [36-39]. Sairyo et al. [16,19] reported that damaged LF tissue, induced by mechanical stress, might be the initial trigger for an inflammatory reaction and subsequent development of tissue scarring. Attention has recently focused on angiogenesis, a pivotal component of scarring and the tissue repair process. Moon et al. [40] reported that LF cells interact with macrophage-like cells to produce angiogenesis-related factors, with the exception of transforming growth factor β1. Activated LF cells exposed to macrophages can impact the stimulus of angiogenesis-related factors. This indicates that fibrosis and scarring, triggered by an inflammatory reaction, is the major pathomechanism of LF hypertrophy. Therefore, inflammatory reactions are initiated by trauma such as mechanical stress and are followed by the repair process. Hypertrophied LF, containing more collagen fibers, is vulnerable to repetitive flexion-extension motion resulting in a vicious cycle of scar-repair-scar.

Mechanical stress is one of the most important factors in LF hypertrophy, yet it is unclear precisely what form of mechanical stress induces hypertrophy. Fukuyama et al. [25] reported that LF with radiological degenerative signs such as spondylolisthesis or a vacuum phenomenon exhibits severe hypertrophy and suggested that mechanical instability is a possible cause of LF hypertrophy. In our study, disc space widening was identified as a significant risk factor of LF thickening, although the vacuum phenomenon was not retained as a predictive factor in our logistic regression analysis. Abbas et al. [34] reported that LF thickness in the LSCS group occurred at L3–4 and L4–5 segments because of the relative hypermobility of these two segments compared with the L5–S1 segment, which is stabilized by the iliolumbar ligaments and the large transverse processes of the L5 vertebra. Sairyo et al. [16] investigated LF thickness and used a three-dimensional finite element model to characterize the effects of mechanical stresses on various LF layers. Maximum stress was observed in flexion mode activities such as lifting. More biomechanical stress on the dorsal side of the LF than the ventral side led to LF hypertrophy.

In our study, a logistic regression analysis showed that LF thickening was influenced by disc space widening, facet joint osteoarthritis, and patient age. The possibility of a relationship between LF thickening and facet joint osteoarthritis has rarely been reported. Wang and Yang [41] suggested that a coronal L4–5 facet joint orientation negatively correlated to age, which could explain why aging people are prone to degenerative spondylolisthesis; degenerative spondylolisthesis is considered to be one of the causes of induced mechanical stress. In our study, a greater LF thickness was associated with a more sagittalized facet joint orientation and facet joint osteoarthritis. These findings suggest that the aging process increases disc degeneration and affects the facet joints, possibly leading to segmental instability on flexion-extension motion. Therefore, repetitive flexion-extension motion cycles
may lead to LF hypertrophy.

Our study has several limitations. Foremost, numerous factors contribute to the narrowing of the spinal canal beyond LF thickening, including developmental canal stenosis or spondylolisthesis. To further understand canal narrowing pathogenesis, these additional factors should be included in future analyses. Second, our study group was small and did not include a control group. Therefore, the correlation between LF thickness and LSCS symptoms could not be evaluated within the context of our study. In addition, LF thickening was defined by a thickness of ≥4 mm, representing a narrow range in comparison with previous studies [16,25,26,34,42,43].

Conclusions

LF thickness correlated with age, disc space widening angulation, and facet joint osteoarthritis. LF hypertrophy development was associated with segmental instability and severe disc degeneration, severe facet joints osteoarthritis, and sagittalized facet joint orientation.

Conflict of Interest

No potential conflict of interest relevant to this article was reported.

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