Evidence-Based Analysis of Adjacent Segment Degeneration and Disease After LIF: A Narrative Review

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Abstract

Study Design: Narrative review.

Objectives: The etiology of adjacent segment degeneration (ASDeg) and adjacent segment disease (ASDz) after lumbar interbody fusion (LIF) remains controversial. The aim of this narrative review was to provide an evidence-based analysis of the peer-reviewed literature on clinical studies of ASDeg and ASDz after LIF.

Methods: A review was performed utilizing Medline, Embase, and Cochrane databases. Two reviewers independently extracted relevant data from each included study. Statistical comparisons were made when appropriate.

Results: Nine articles that matched the inclusion and exclusion criteria were included. All the studies were Level III and retrospective. MINORS scores ranged from 9.5 to 13. Clinical outcomes were assessed in all 9 studies, but only 6 studies used validated outcomes measures. Only 6 studies reported values for both ASDeg and ASDz. ASDeg alone was reported in 3 studies. Due to the variability in the criteria for designation as ASDz (different radiographic modalities) and ASDeg (different outcomes measures), we were unable to calculate frequency-weighted mean values or compare the various surgical techniques.

Conclusions: This review highlights the various limitations of the current literature on ASDeg and ASDz after lumbar fusion, specifically the absence of a rigorous definition and classification system and an extraordinary heterogeneity in methodology. There needs to be a fundamental shift in the current ASDeg and ASDz research landscape, toward a consensus, so that the high-level clinical research that is essential for treatment of spinal pathology may become available.

Keywords
LIF, adjacent segment degeneration, adjacent segment disease, fusion, lumbar

Introduction

Briggs and Milligan first described lumbar interbody fusion (LIF) for the surgical treatment of chronic low back pain 70 years ago. Although there is considerable debate over the indications for lumbar fusion, it remains the gold standard surgical treatment for patients who fail nonoperative methods in many clinical scenarios.1-4

Adjacent segment degeneration (ASDeg) is defined as radiographic degenerative changes at a spinal level immediately cranial or caudal to the site of a previous fusion procedure. ASDeg can progress to adjacent segment disease (ASDz), a clinical phenomenon characterized by the presentation of new symptoms referable to the adjacent level, presumably related to the degenerative changes.5 Although ASDeg and ASDz are sometimes used as outcome measures of LIF, their etiology remains controversial. One theory holds that adjacent segment pathologies, like ASDeg and ASDz, are simply reflections of the natural history of lumbar degenerative disease, which is often characterized by desiccation of all lumbar segments, and are therefore inevitable. Others cite biomechanical findings to argue that LIF results in increased motion, intradiscal pressure, and strain adjacent to the fusion, leading to an increased risk of ASDeg and ASDz.5-7
Resolving this debate has proven exceedingly difficult due to the challenges of studying LIF in the clinical setting. Although spinal fusion is an established and exceedingly common procedure, the steady evolution of surgical techniques and the availability of a myriad of graft materials, cage designs, and plate fixation systems have given rise to considerable variability between treatment methods. In addition, the current literature on adjacent segment pathology suffers from the absence of a universally accepted radiographic modality for diagnosis of ASDeg or validated outcome instrument for diagnosis of ASDz. Because of this heterogeneity, estimates of ASDz incidence after lumbar fusion range from 1.9% to 30.3%,. One systematic review of ASDeg reported a range of incidence from 8% to 100%. As a result of these barriers to clinical research, there is virtually no high-level evidence on ASDeg and ASDz after LIF. The purpose of this narrative review was to analyze the existing literature to determine if there are sufficient similarities among studies to support a meta-analysis.

**Materials and Methods**

**Literature Search**

A systematic search of PubMed, Embase, and Cochrane Reviews for literature published through June 2013 was done on July 31, 2013. The search terms for PubMed and Embase are presented in Tables 1 and Table 2. The PRISMA flow diagram is presented in Figure 1. Search results were limited to full text articles with a minimum follow-up of 1 year written in the English language. All abstracts were independently reviewed by 2 clinical research fellows and a fellowship-trained orthopedic spine surgeon. The following study designs were excluded from this analysis: meta-analyses, systematic reviews, and review articles. Articles focused on the following topics were also excluded: biomechanics analyses (cadaveric, in vitro), animal studies, trauma, and tumor studies.

GalTable 1. Medline Literature Search.

| Search | Query Input |
|--------|-------------|
| #1     | Spinal Fusion [MeSH] or “Spinal Fusion” |
| #2     | Lumbar [MeSH] or Lumbar |
| #3     | 1 AND 2 |
| #4     | ALIF or “Anterior Lumbar Interbody Fusion” |
| #5     | PLIF or “Posterior Lumbar Interbody Fusion” |
| #6     | PLF or “Posterior Lumbar Fusion” |
| #7     | TLIF or “Transforaminal Lumbar Interbody Fusion” |
| #8     | XLIF or “Extreme Lateral Interbody Fusion” |
| #9     | 4 OR 5 OR 6 OR 7 OR 8 |
| #10    | 3 OR 9 |
| #11    | “adjacent segment” or “adjacent level” |
| #12    | 10 OR 11 |

Table 2. Embase literature Search.

| Search | Query Input |
|--------|-------------|
| #1     | spine fusion.mp. or spine fusion/ |
| #2     | lumbar.mp. or exp lumbar spine/ or exp lumbar disk/ or exp lumbar spinal cord/ |
| #3     | (ALIF or “Anterior Lumbar Interbody Fusion”).mp. [mp=title, abstract, subject headings, heading word, drug trade name, original title, device manufacturer, drug manufacturer, device trade name, keyword] |
| #4     | (PLIF or “Posterior Lumbar Interbody Fusion”).mp. [mp=title, abstract, subject headings, heading word, drug trade name, original title, device manufacturer, drug manufacturer, device trade name, keyword] |
| #5     | (PLF or “Posterior Lumbar Fusion”).mp. [mp=title, abstract, subject headings, heading word, drug trade name, original title, device manufacturer, drug manufacturer, device trade name, keyword] |
| #6     | (TLIF or Transforaminal Lumbar Interbody Fusion).mp. [mp=title, abstract, subject headings, heading word, drug trade name, original title, device manufacturer, drug manufacturer, device trade name, keyword] |
| #7     | (XLIF or “Extreme Lateral Interbody Fusion”).mp. [mp=title, abstract, subject headings, heading word, drug trade name, original title, device manufacturer, drug manufacturer, device trade name, keyword] |
| #8     | (“Adjacent Segment” or “Adjacent Level”).mp. [mp=title, abstract, subject headings, heading word, drug trade name, original title, device manufacturer, drug manufacturer, device trade name, keyword] |
| #9     | 1 AND 2 |
| #10    | 3 OR 4 OR 5 OR 6 OR 7 OR 8 |
| #11    | 8 OR 9 |
| #12    | 10 AND 7 |

Articles (n = 284) from the primary literature search were then retrieved and further examined by 3 reviewers (NNT, LAP, and DRL). After critical examination of the full text of the remaining 19 articles, 10 additional articles were excluded, leaving 9 articles to be included in this review. Relevant data was identified and extracted from the full text of all remaining articles (Table 3).

**Level of Evidence**

The criteria outlined by Wright et al were used to evaluate the relative merit of the studied included in this review. In addition, the MINORS criteria outlined by Slim et al were used to evaluate the studies. Each article was independently graded by 2 authors. Any interreviewer variation, defined as difference in study level or difference in MINORS score >2, would be resolved by a third reviewer.

**Results**

Nine articles that matched the inclusion and exclusion criteria were included (Table 3). All of the studies were determined to be Level 3 evidence and retrospective, and the average MINORS scores ranged from 9.5 to 13 (Table 4). There were a multitude of clinical and radiographic initial
presentations that were reported (isthmic spondylolisthesis, degenerative spondylolisthesis, degenerative disc pain, etc).

**Demographics**

In total, 625 patients were included in the 9 studies with an average of 69 patients per study (range = 9-133). All 9 studies provided demographic data on sex, leading to a total of 254 males and 371 females. Seven studies provided age. The frequency-weighted mean age for those 7 studies was 43.4 (range 45-70.5). The frequency-weighted average follow-up was 60.1 months with a range from 13.8 to 246.

**Surgical Technique**

Six studies specifically analyzed 1- or 2-level lumbar fusion, and 3 studies pooled data on several fusion lengths, ranging from 1 to 4 levels. Plate fixation was used in one of the articles, accounting for 13% of surgeries. Pedicle screw was used in 7 of the articles, accounting for 87% of surgeries.

Two studies reported using only posterior lumbar fusion (PLF) for fusion, and 3 studies reported using only anterior lumbar interbody fusion (ALIF). One study reported using only posterior lumbar interbody fusion (PLIF), and 3 studies were either nonspecific or reported multiple surgical techniques. Because of the variability in measuring ASDeg and ASDz, we were unable to compare incidence for the various surgical techniques.

**Outcome Assessment**

Clinical outcomes were assessed in all 9 studies. Six used validated outcomes measures, including the visual analog scale, Oswestry Disability Index, Low Back Outcome Scale, and Japanese Orthopedic Association score. The remaining 3 studies reported tolerance of walking, pain scale, and reoperation.

Radiographic evaluation was reported in all 9 of the studies. Of the modalities employed, plain radiographs were used in 5 studies, with computed tomography in 2 studies and magnetic resonance imaging in 4 studies. One study did not specify radiographic modality.

**Outcomes: ASDeg and ASDz**

Only 6 articles reported incidence for both ASDeg and ASDz. Adjacent segment degeneration alone was reported in 3 studies. However, due to the variability in criteria for designation as ASDz or ASDeg, we were unable to calculate frequency-weighted mean values.
Discussion

Long-term follow-up studies on LIF have described radiographic deterioration of discs adjacent to the level of index surgery.\textsuperscript{5,26,27} Adjacent segment pathology is differentiated in the literature into radiographic degeneration (ASDeg) and the associated clinical entity (ASDz).\textsuperscript{28} Unfortunately, there is no consensus on the definitions or classification systems for ASDeg or ASDz. Nevertheless, this distinction is useful for surgical decision making and patient counseling regarding the likelihood of the need for reoperation.

The relationship between ASDeg and ASDz after LIF remains an important topic of research interest. There is some data indicating that the severity of ASDeg might predict clinical outcome, specifically ASDz, but the evidence is limited.\textsuperscript{29} Other studies have found no association between the degree of ASDeg and clinical or functional outcomes, and some suggest that routine radiographic follow-up has limited utility.\textsuperscript{14,23,30-33} Rigorous definitions and classification methods for ASDeg and ASDz are required before

Table 3. Summary of Literature.

| Study          | Number of Patients (Males/Females) | Age (Average) | Surgeries Performed | Follow-up: Months, Average (Range) | Clinical Assessment Tools | Radiographic Modality | ASDDegeneration (% Patients) | ASDDisease (% Patients) |
|----------------|-----------------------------------|---------------|---------------------|----------------------------------|--------------------------|-----------------------|--------------------------|--------------------------|
| Bae et al\textsuperscript{12} | 103 (39/94)                       | 48.5          | Mini-ALIF; mini-TLIF | 51.9 (36-94)                     | Visual Analog Scale, ODI, patient’s return to work status | CT                     | 10.7                     | 1.9                      |
| Chou et al\textsuperscript{25} | 32 (20/12)                        | 70.5          | PLF                 | 56 (48-66)                       | Tolerance of walking claudication and the severity of back pain or radicular pain | X-ray                  | 18.8                     | 12.5                     |
| Kim et al\textsuperscript{17}  | 94 (23/71)                        | 50.2          | ALIF and PLIF       | 70 (62-86)                       | VAS and ODI scores       | MRI                   | 75.7                     | 18.1                     |
| Axelsson et al\textsuperscript{19} | 9 (3/6)                          | 45            | PLF with/without instrumentation and ALIF | 60                      | Pain Scale (3 options) | X-ray                  | 22.2                     | 11.1                     |
| Rahm et al\textsuperscript{18} | 102 (69/33)                       | 54            | Anterior fusion with a pararectal approach; AP fusion with a ventral pararectal approach | 13.8                   | ODI/Patient Satisfaction Scores | X-ray                  | 26.5                     | 27.5                     |
| Wai et al\textsuperscript{22}  | 39 (19/20)                        | 58.3          | ALIF                | 246 (240-306)                    | Low Back Outcome Scale | MRI                   | 74.3                     | NA                       |
| Okuda et al\textsuperscript{23} | 87 (38/49)                        | 64            | PLIF                | 43 (24-78)                       | Japanese Orthopedic Association score | MRI and CT            | 67                       | NA                       |
| Min et al\textsuperscript{21}  | 48 (14/34)                        | NA            | ALIF, PLIF, TLIF   | 44.6 (24-68)                     | Japanese Orthopedic Association score | X-ray                  | 62.5                     | NA                       |
| Gillet et al\textsuperscript{24} | 106 (35/71)                      | NA            | PLF                 | 60                               | Did not specify          | Did not specify       | 41                       | 20                       |

Abbreviations: ALIF, anterior lumbar interbody fusion; TLIF, transforaminal lumbar interbody fusion; VAS, Visual Analog Scale; ODI, Oswestry Disability Index; CT, computed tomography; PLF, posterior lumbar fusion; PLIF, posterior lumbar interbody fusion; MRI, magnetic resonance imaging.

Table 4. Summary of Level of Evidence and MINORS Scores.

| Study          | Level of Evidence | MINORS Score |
|----------------|-------------------|--------------|
| Bae et al\textsuperscript{12} | Level III (Retrospective) | 12.5         |
| Chou et al\textsuperscript{25} | Level III (Retrospective) | 9.5          |
| Kim et al\textsuperscript{17}  | Level III (Retrospective) | 13           |
| Axelsson et al\textsuperscript{19} | Level III (Retrospective) | 11           |
| Rahm et al\textsuperscript{18} | Level III (Retrospective) | 10           |
| Wai et al\textsuperscript{22}  | Level III (Retrospective) | 11           |
| Okuda et al\textsuperscript{23} | Level III (Retrospective) | 10           |
| Min et al\textsuperscript{21}  | Level III (Retrospective) | 11.5         |
| Gillet et al\textsuperscript{24} | Level III (Retrospective) | 11           |
high-quality studies can address this question with any certainty.

The central debate revolves around the etiology of adjacent segment pathology, specifically whether ASDeg and ASDz are part of the natural history of the disease or induced by biomechanical adjustments postsurgery. Proponents of the natural history theory argue that lumbar degenerative disease is characterized by gradual desiccation of all lumbar discs, and therefore deterioration would occur even without surgery. The natural history argument is supported indirectly by evidence that many asymptomatic patients exhibit progressive abnormal radiographic findings. One study found that disc degeneration progresses in 41% of asymptomatic patients and another demonstrated that 57% of asymptomatic patients over 60 years of age have abnormal findings on magnetic resonance imaging scans of the lumbar spine. In addition, Wai et al determined in a series of 39 patients that the majority of degenerative changes occurred homogeneously across lumbar levels, suggesting that deterioration is determined by characteristics specific to the patient. Similarly, Pisellé et al found that in patients with degenerated adjacent disc, 50% of other levels also exhibited degeneration. In patients with normal adjacent discs, only 8.6% of the remaining discs had degeneration. Evidence of a genetic predisposition for lumbar degenerative disease offers further support for this argument. Twin studies attribute 26% to 76% of the variability in disease incidence to genetic factors. Although many proponents of the natural history theory do not fully dismiss the biomechanical explanation of ASDeg and ASDz, they argue that it is negligible.

This argument is complicated by evidence, albeit limited and low level, that lumbar fusion patients have an elevated rate of ASDeg and ASDz over nonsurgical controls. The biomechanical argument supplies an attractive explanation for this possible elevation of ASDeg and ASDz incidence. Numerous biomechanical studies using animal and human models demonstrate that lumbar fusion can result in pressure, force, and motion adjustments at adjacent levels. However, these studies only demonstrate changes in biomechanical forces, and although various animal models have demonstrated ASDeg and ASDz after fusion, there is no quality clinical evidence that biomechanical changes directly cause radiographic (ASDeg) and clinical (ASDz) deterioration.

Furthermore, both cadaveric and animal studies are severely limited in their ability to reproduce the in vivo properties of the human spine, and some authors argue that this results in an overestimation of mechanical alterations. In one biomechanical study, Rohlmann et al concluded that interspecimen differences and the complex loading conditions that do not reproduce the in vivo environment best explain the observed changes in pressures. Clinical studies have also challenged the biomechanical argument. In a radiographic study of hypermobility after lumbar spine fusion, Axelsson et al found no increase in mean mobility 5 years after surgery and no correlation between mobility and clinical outcomes. However, although this study is clinical, it suffers from a small sample size (n = 9) and other limitations. Because of the absence of high-quality clinical research, biomechanical studies will remain central to the debate, despite their obvious limitations.

Finally, this argument is even further complicated by the multitude of surgical techniques for LIF currently in practice, including ALIF, TLIF (transforaminal lumbar interbody fusion), XLIF (extreme lateral interbody fusion), PLF, and PLIF. Some argue that ALIF allows for preservation of the posterior ligaments and soft tissue, which could further prevent the likelihood of ASDeg and ASDz. However, there have been no consistent findings in the literature substantiating this claim or that any of the LIF techniques is superior with respect to ASDeg and ASDz.

There are many proposed risk factors for ASDeg and ASDz, including age, instrumentation, fusion type, fusion length, and degree of lumbar lordosis. The studies in this review addressed some of these risk factors, though pervasive covariability and low sample sizes limit their generalizability and conclusiveness.

There is evidence in the literature, including studies in this review, that the degree of lumbar lordosis is an important risk factor for adjacent segment pathology. In twin studies, that the degree of lumbar lordosis was associated with ASDeg, suggesting that maintaining the preoperative lordosis may protect against the development of ASDeg. It is important to note that many other studies have not reproduced these findings to date.

The number of fused levels is another proposed risk factor for ASDeg and ASDz, and there is evidence that patients with multilevel fusions are at 2 to 3 times greater risk for ASDz. However, in a study of 215 patients, Ghiselli et al reported that single-level fusion patients are up to 3 times more likely to develop ASDz. Six studies in this review included multilevel fusions, and 3 of those evaluated the risk associated with multilevel fusions. Chou et al found a trend toward higher incidence of ASDeg in the multilevel fusion group, but this was not statistically significant (P = .79). Rahm et al detected a similar correlation between the number of levels and ASDeg, but this was also not significant due to a lack of power. Gillet et al observed reoperation rates for single, double, and 3- to 4-level fusions of 11%, 27%, and 33%, respectively. Taken together with the existing literature, it seems likely that there is an increased risk of ASDz with multilevel fusions, but there is need for higher quality studies. The correlation between number of fusion levels and ASDeg remains unclear.

This review is most useful to the extent that it highlights the current limitations in the literature. For one, it illustrates the
low quality of evidence currently available. No Level I or Level II studies met the search criteria: the review was limited to studies with Level III evidence. Furthermore, this review found no studies with MINORS criteria scores greater than 13, the major limiting factor being the retrospective nature of all the studies that met the inclusion criteria.

The review also highlights some of the practical barriers to quality clinical research. The heterogeneity of approaches used to detect ASDeg and ASDz is the most important. For example, while all 9 of the studies in this review analyzed ASDz incidence, they used 10 different clinical assessment tools in 8 combinations. These measures ranged from reoperation to an unvalidated pain scale. Only 2 pairs of studies used the same metric to define ASDz. Similar variation in evaluation systems was evident in the analysis of ASDeg. This inconsistency is common throughout the literature and poses a major barrier to high-quality research.

Indeed, the defining feature of this narrative review was the variability between studies. Surgery type, number and location of fused levels, screw type, cage type, and graft material differed considerably between (and within) studies, when provided at all. For example, average follow-up time varied from 3 to 246 months.20,22 Such inconsistencies introduce a challenging mass of covariables. This reality, considered in the absence of rigorous definition and classification systems, makes it impractical to synthesize the existing literature with any confidence. The extraordinary variability between studies also precludes a more sophisticated statistical approach (ie, meta-analysis).

Conclusion

Despite the high success rate of LIF, follow-up has identified ASDeg and ASDz as potential postoperative complications. The literature is currently divided on whether adjacent segment pathology represents the natural history of the disease or results from surgery-induced biomechanical stresses. This review highlights the various limitations of the current literature on ASDeg and ASDz after lumbar fusion, specifically the absence of a rigorous definition and classification system and an extraordinary heterogeneity in methodology. There needs to be a fundamental shift in the current ASDeg and ASDz research landscape, toward a consensus, so that the high-level clinical research that is essential for treatment of spinal pathology may become available. This shift must begin with the development of a rigorous definition and classification system for ASDeg and ASDz. It will also require more consistent methodology, something that multi-surgeon collaboration could help achieve.

Authors’ Note

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