Adjacent Segment Pathology after Anterior Cervical Fusion

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Anterior cervical fusion has become a standard of care for numerous pathologic conditions of the cervical spine. However, subsequent development of clinically significant disc disease at levels adjacent to fused discs is a serious long-term complication of this procedure. As more patients live longer after surgery, it is foreseeable that adjacent segment pathology (ASP) will develop in increasing numbers of patients. Also, ASP has been studied more intensively with the recent popularity of motion preservation technologies like total disc arthroplasty. The true nature and scope of ASP remains poorly understood. The etiology of ASP is most likely multifactorial. Various factors including altered biomechanical stresses, surgical disruption of soft tissue and the natural history of cervical disc disease contribute to the development of ASP. General factors associated with disc degeneration including gender, age, smoking and sports may play a role in the development of ASP. Postoperative sagittal alignment and type of surgery are also considered potential causes of ASP. Therefore, a spine surgeon must be particularly careful to avoid unnecessary disruption of the musculoligamentous structures, reduced risk of direct injury to the disc during dissection and maintain a safe margin between the plate edge and adjacent vertebrae during anterior cervical fusion.

Keywords: Cervical vertebrae/surgery; Spinal fusion; Complications; Adjacent segment pathology; Risk factors; Reoperation

Introduction

Anterior cervical discectomy and fusion (ACDF) was first reported in 1955 [1]. This technique provides relatively easy access to surgical sites, a lower bleeding risk, and a good view of the operative field [2,3]. Over the past 60 years, ACDF has become a standard method for many pathologic conditions of the cervical spine. Anterior cervical fusion in conjunction with decompression provides relief of radicular complaints and improvement of myelopathic findings in over 90% of cases [1,4]. However, subsequent development of clinically significant disc disease at levels adjacent to fused discs is a serious long-term complication of this procedure (Fig. 1) [5,6]. Also, as more patients live longer after surgery, it is foreseeable that adjacent segment pathology (ASP) will develop in increasing numbers of patients. ASP has become a common topic in spine surgery because of the development of motion preservation technologies like total disc arthroplasty that theoretically should lead to a decrease in this pathology. With the recent popularity of this new motion preservation technology, the study of ASP has increased enormously.

However, controversy remains as to whether these
Adjacent-segment pathology after anterior cervical fusion conditions are related to altered biomechanics [5,7-11] or represent the natural history of cervical spondylosis at the adjacent segment [12-14]. Clearly, patients who have undergone cervical fusions are at an increased risk of developing ASP. What remains unknown is how much of this risk is related to the surgical procedure and the natural history of their disease.

This review presents data on the incidence, pathogenesis and clinical impact of ASP after cervical fusion, and summarizes the analyses of predictive factors and prevention of ASP. Many comparative studies associated with total disc arthroplasty have been reported. However, a comprehensive summary is beyond the scope of the present review.

**Terminology**

The true incidence and clinical impact of degenerative changes at the adjacent segment is unclear because of the lack of a universally accepted classification system that rigorously addresses clinical and radiological issues.

In the scientific literature, “adjacent segment degeneration”, “adjacent segment disease”, “ASD”, “junctional disease”, “junctional problem” and “junctional stenosis” have been used to describe spinal degenerative pathology, which coexists in the adjacent spine after a previous spinal fusion. “Degeneration” has also been poorly defined, and may separately refer to osteophyte formation, intervertebral disc degeneration, spinal stenosis, segmental instability, facet arthrosis or significant structural deformity including kyphosis and scoliosis. Thus, although “ASD” is a commonly used term, its definition is imprecise and it is created varied and ambiguous literature regarding this pathology [15]. Recently, the term “adjacent segment degeneration” has been used to describe radiographic changes seen at levels adjacent to a previous spinal fusion that do not necessarily correlate with any clinical findings. On the other hand, the term “adjacent segment disease” has been used to refer to the development of new clinical symptoms that correspond to radiographic changes adjacent to the level of a previous spinal fusion. Despite the clear distinction between these two terms, they are also often used interchangeably in the literature.

A recent Spine Focus issue recommended uniform terminology comprising radiographic adjacent segment pathology without symptoms (RASP) and clinical adjacent segment pathology with symptoms (CASP). The term adjacent segment pathology (ASP) was proposed as an umbrella term to refer to the breadth of clinical and/or radiographical changes at adjacent motion segments that developed subsequent to a previous spinal fusion. Under this umbrella, RASP and CASP are then used to categorize radiographical features and clinical manifestations, respectively. The intent is to have a uniform and unambiguous terminology.
Pathogenesis

Development of ASP following anterior cervical fusion has been amply described in the literature [5,12,16-23]. The cause of this degenerative process has been extensively studied in biomechanical studies using animal and cadaver models [24-27]. Still, the true nature and scope of ASP remains poorly understood. Whether ASP following ACDF represents a true iatrogenic postoperative complication or a progression of the natural history of cervical spondylosis remains unclear. The etiology of ASP is most likely multifactorial. No study has proven that a single risk factor directly correlates with this pathology. The natural history of degeneration, changes in intradiscal pressure, anatomy disruption and sagittal malalignment have been proposed as etiologic factors [28-30].

Many studies have focused on the altered biomechanics at the adjacent levels after fusion that result in increased mobility [5,7,9,10], increased loading [8] or increased intradiscal pressure [11] that ultimately accelerate disc degeneration [25,31-33]. Increased mechanical demands adversely affect the disc by interfering with its normal nutritional supply. Impaired disc nutrition is the most significant cause of disc degeneration [31]. Increased pressure within the disc is believed to inhibit diffusion of nutrients, which leads to the accumulation of waste products [34]. Cadaveric studies of the spine have demonstrated that changes in pressure, force, and motion occur at adjacent levels as a result of arthrodesis [24,35]. Although these studies do not account for the true in vivo properties of living tissues and the contribution of the spinal musculature, they provide some insight into the altered forces at the adjacent segment after a fusion.

In a study that involved humans and animals, the authors reported that the mobile segments adjacent to the fusion segment showed an increased range of motion, and this effect is increased with multilevel fusions [36]. Another study reported that regardless of the fusion method, cervical fusion increased the mobility of the adjacent segments and facilitated degenerative change [37]. Also, at 7.2 years of follow-up monitoring, 32% of patients had recurrent pain and 16% required additional surgery.

Reported rates of ASP in patients with fusion seem to be higher than the rates of de novo degeneration in patients without fusion, suggesting that the fusion itself may have a contributory effect in the development of ASP.

Whereas some authors advocate the hypothesis of altered biomechanics, others focus on natural progression at adjacent levels after fusion. In addition, the aforementioned disc changes occurred during the normal aging process. As the disc ages, its nutrition is impaired because the presence of fewer peripheral arteries and of calcification of the cartilage end plates reduces the vascular supply. Furthermore, loss of viable cells, modification of matrix proteins and fatigue failure of the matrix occur [34]. Physiologic aging of the cervical spine has been observed in several cross-sectional and longitudinal studies in healthy volunteers. Magnetic resonance imaging (MRI) in 63 asymptomatic volunteers revealed that the intervertebral disc was narrowed or had degenerated at one or more cervical levels in 25% of those under 40 years of age and in 60% of those older than 40 years of age [38]. MRI of the cervical intervertebral discs of 497 asymptomatic subjects demonstrated a linear increase in degenerative findings with age, from 17% of men and 12% of women aged 20 to 29 years compared with 86% of men and 89% of women aged >60 years [39].

Fusion and nonfusion procedures have been compared to determine whether the fusion itself might be causative in the development of ASP. If the biomechanical consequences of cervical fusion were the only factor in the development of ASP, nonfusion procedures like anterior discectomy and posterior foraminotomy would not increase the risk. In a relatively short follow-up of >1 years involving 253 patients who underwent anterior cervical discectomy with fusion or discectomy alone, the authors did not find any difference in the rate of CASP between patients who underwent discectomy with fusion and those who underwent discectomy alone [40]. A retrospective review of a cohort of 846 patients after posterior foraminotomy without fusion with an average follow-up of 2.8 years reported 79 patients required additional procedures for CASP (9%; approximate annual incidence, 3%) [41]. A prospective study reported that after anterior cervical fusion, 41% of the patients had radiographic evidence of ASP at an average 4.5-year follow-up, compared with 50% of those who underwent posterior cervical laminoforaminotomy [42]. These clinical observations suggest that anterior decompression with fusion and posterior decompression without fusion may lead to similar rates of ASP.

The relationship between the number of the fused segments and ASP is contentious. Some studies have demonstrated an increased risk of degeneration with increasing length of fusion [43-46], while other studies reported
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that the increase in the number of fused segments did not result in a higher rate of ASP [47,48]. In one study, the risk of CASP after multilevel fusion was significantly less than after single-level fusion [12]. These findings are contrary with the results expected if the biomechanical consequences of fusion were the only cause of ASP. In recent years, total disc arthroplasty has become an alternate surgical procedure that may replace spinal fusion in patients with degenerative disc disease. The goals of disc arthroplasty are to preserve motion and restore the intervertebral disc and foraminal height to prevent recurrence of nerve root compression. But, after a 2- to 4-year follow-up, ASP was also found in total disc arthroplasty patients [49]. Therefore, whether ASP is caused by the natural progression of aging or by increased motion stress related to biomechanical factors secondary to the surgical fusion itself remains debatable.

Moreover, general factors linked with disc degeneration including gender, smoking and sporting activities may play a role in the development of ASP. Soft tissue disruption is also considered a potential cause of ASP, although this is much more difficult to experimentally elucidate. Surgical damage to the anterior longitudinal ligament or to the longus colli muscles can contribute to ASP. Incorrect needle placement during intraoperative radiographic-level confirmation was reported to increase the relative risk of developing ASP by three-fold [50]. The authors attributed the increased ASP to both the dissection over the adjacent level and the puncture of the annulus.

Considering the collective data, we think that various factors including altered biomechanical stresses, surgical disruption of soft tissue and the natural history of cervical disc disease are causal for the development of ASP (Fig. 2).

Clinical ASP and Reoperation

ASP has been reported in multiple studies after both anterior and posterior fusions, and at single and multilevel. However, the reported incidence has varied. At an average 100.6 month follow-up, 92% of the 180 patients treated by ACDF demonstrated additional radiographic degeneration at the adjacent disc levels [51]. Another study reported that 92% of 177 patients treated by ACDF showed some degree of radiographic change [52]. Still another study reported RASP in 67% of participants after a mean follow-up of 27 months after anterior cervical fusion [7].

In contrast with previous studies about radiographic changes of adjacent level after cervical fusion, present
studies focused on CASP. Hilibrand et al. [12] reported that new CASP occurred at a rate of 2.9% per year and that 25.6% would have new disease at an adjacent level within 10 years after ACDF. A review of the outcomes of 177 patients after ACDF found that 19.2% developed CASP during a minimum 10 years of follow-up, of which 6.8% required reoperation [52]. Others observed new spondylosis in 25% of 121 patients and progression of preexisting spondylosis in another 25% of patients who had previously undergone anterior cervical fusion with an average follow-up of 5 years [53].

The incidence of cervical RASP has ranged widely from 18.33% [54] to 96% [37], as has CASP, with values from 11.1% [55] to 38.1% [7]. RASP occurs at a significantly higher rate than CASP, a finding clearly supported by the studies reviewed [20,22,51,54,56-61]. The relationship between RASP and CASP has been considered, specifically whether RASP can predict the development of CASP. No correlation between RASP and the development of clinical symptoms referable to radiographic changes has been apparent [7,51,60]. Consequently, radiographic evidence of degeneration of disc spaces adjacent to the site of anterior cervical fusion may be meaningful only when it is associated with clinical symptoms of radiculopathy or myelopathy referable to that level.

Varying rates of surgery to repair ASP have been reported. Several studies reported similar rates of 6% [51], 6.3% [18], 6.7% [55] and 6.8% [52], while higher rates of 9.1% [13], 16% [37] and 17% [22,62] have been documented on other studies. When the number of years of follow-up is divided into the overall prevalence, the annual incidence of CASP requiring additional surgery is between 0.8% and 4% (Table 1).

Risk factors for follow-up surgery include type of surgery, smoking and female sex. In one study, the posterior arthrodesis group had a 7.5-times greater risk of ASP requiring follow-up surgery than those receiving posterior decompression, and a 3.0-times greater risk than the anterior arthrodesis group [63]. Possible causes included soft-tissue injury of ligaments, facet capsule and muscle during posterior arthrodesis, or the rigidity of posterior instrumentation.

### Plate-To-Disc Distance and Adjacent Level Ossification

Anterior osteophyte formation is more frequently observed in those with fusion than in those without fusion. Fusion with cervical plates has been used widely in settings including trauma, disc herniation, spondylosis and tumors because of advantages that include initial stability, higher fusion rate, recovery of normal lordosis and the lack of necessity for external support. However, in patients with an anterior cervical plate, the distance between the plate and adjacent segments may influence the osteophyte at adjacent segments [64]. The association between adjacent level ossification and disease has not been proven, but the placement of the plate closer to the adjacent level could potentially disrupt the adjacent level anatomy, such as the anterior longitudinal ligament.

Some study reported that a plate-to-disc distance of less than 5 mm results in moderate or severe ossification of the adjacent segments and therefore a surgeon performing ACDF should try to achieve a distance of 5 mm or more [64,65]. Another study reported that most CASP occurred when the plate-to-disc distance was less than 5 mm. Goffin et al. [51] also recommended using the shortest plate possible to avoid intrusion of the plate into the adjacent segments. A cutoff of 3 mm has been suggested to avoid adjacent level ossification development (ALOD) [66]. ALOD results in heterotopic bone formation along the anterior longitudinal ligament at the adjacent levels. In theory, ALOD may be caused by an inflammatory reaction between the plate and anterior longitudinal ligament leading to ossification. This is evidenced by higher rates of ALOD seen in plated ACDF compared with noninstrumented ACDF [64,67,68].

ALOD is a sequela of a close plate and may also result from dissection at the disc. The closer the plate lies to the disc, the greater the risk of soft tissue dissection at the anus, such as a soft tissue injury by a periosteal elevator or other surgical tool used when placing the cervical plate. This can lead to subsequent ossification. But, the precise mechanism of ALOD is still unknown.

Recently, other authors [69,70] reported that although both ALOD and ASP can result in bone formation, the pathological mechanism of ALOD differs from that of osteophytic growth in ASP. They reported that patients with ALOD typically have better preservation of disc height compared with patients with ASP. Because ALOD in its severest stages causes ankylosis and a decrease in motion, the result is a “relative protection” of the adjacent level from degenerative changes. Patients with ASP are more likely to have disc protrusions, facet arthrosis,
Table 1. Literature review regarding adjacent-segment pathology

| Study                          | No. of cases in study | Duration of follow-up in years (range) | No. of patients with RASP (%) | No. of patients with CASP (%) | No. of patients undergoing follow-up surgery (%) | Clinical characteristics |
|-------------------------------|-----------------------|----------------------------------------|------------------------------|------------------------------|-----------------------------------------------|--------------------------|
| Baba et al. [5]               | 106                   | 8.5 (5–16.3)                           | 26 (24.5)                    | N/A                          | 17 (16)                                       | Dynamic spinal canal stenosis at the adjacent level showed unfavorable neurologic results. |
| Dohler et al. [7]             | 21                    | 2.3±1.3                                | 14 (66.7)                    | 8 (38.1)                     | N/A                                           | No relationship between the degree of slippage and either neck or arm pain |
| Hilibrand et al. [12]        | 374                   | Max 21                                 | N/A                          | 55 (14.7)                    | 27 (7.2)                                      | CASP was the result of progressive spondylosis |
| Hilibrand et al. [13]        | 417                   | 5.6 (2–15.3)                           | N/A                          | N/A                          | 38 (9.1)                                      | Anterior cervical corpectomy and strut grafting resulted in a significantly higher rate of arthrodesis than interbody grafting in patients surgically treated for adjacent segment disease. |
| Ishihara et al. [18]         | 112                   | 9.4 (2–19)                             | N/A                          | 19 (17)                      | 7 (6.3)                                       | Incidence of CASP was higher when there was preoperative disc degeneration |
| Yue et al. [22]              | 71                    | 7.2 (5.4–11.1)                         | 52 (73.2)                    | N/A                          | 12 (16.9)                                     | No relationship between the incidence of adjacent-segment pathology and follow-up surgery |
| Gore and Sepic [37]          | 50                    | 21.1 (19.5–24.7)                       | 48 (96)                      | 16 (32)                      | 8 (16)                                        | No relationship among the incidence of RASP and CASP |
| Goffin et al. [51]           | 180                   | 8.4±2.6                                | 166 (92.2)                   | N/A                          | 11 (6.1)                                      | No relationship between the incidence of adjacent-segment pathology and follow-up surgery |
| Chung et al. [52]            | 177                   | 16.2 (10.0–25.2)                       | 163 (92.1)                   | 34 (19.2)                    | 12 (6.8)                                      | No relationship among the incidence of RASP, CASP, and follow-up surgery |
| Kawakami et al. [54]         | 60                    | 4.5 (2–9)                              | 11 (18.3)                    | N/A                          | N/A                                           | No significant difference in frequency of disturbance of adjacent segments between patients with and without axial symptoms |
| Teramoto et al. [55]         | 45                    | 10 (4–18)                              | 23 (51.1)                    | 5 (11.1)                     | 3 (6.7)                                       | CASP was the result of progressive spondylosis |
| Komura et al. [56]           | 102                   | 5.1±2.6 (2–10.8)                       | 35 (34.3)                    | 12 (11.8)                    | 3 (2.9)                                       | Significant higher incidence of CASP in the short level ACDF than long-level ACDF and Significant lesser incidence of RASP, CASP in the patients in whom C5–6 and C6–7 are fused than among those in whom C5–6 or C6–7 is left |
| Wu et al. [61]               | 19385                 | 10                                     | N/A                          | N/A                          | 7.6 per 1000 person-years (0.8)               | Kaplan-Meier analysis demonstrated a considerable portion of patients (5.6%) underwent reoperation for ASD at the end of the 10-year cohort |
| Williams et al. [62]         | 60                    | 4.6 (2–9)                              | 50 (83.3)                    | N/A                          | 10 (16.6)                                     | Patient who underwent cervical discectomy and interbody fusion with radicular symptoms had a higher rate of improvement than those whose symptoms were non-radicular. |

All values are expressed as mean±standard deviation. 
RASP, radiographic adjacent-segment pathology; CASP, clinical adjacent-segment pathology; N/A, not applicable; Max, maximum.
spondylolisthesis, endplate sclerosis and/or osteophyte formation. Also, the base of ALOD is situated anterior to the vertebral body, covering about half of it. However, the osteophytes seen in ASP are located anteroinferiorly in the vertebral body, covering no more than one-third of the body. It is not clear whether ASP is due to surgery or the natural history of degenerative disease. ALOD, on the contrary, seems to be an iatrogenic problem that has a high likelihood of developing if the plate is placed too close to the adjacent segment. More importantly, ALOD matures within the first 2 years after surgery, unlike in ASP, in which osteophytic bone and disc degeneration gradually increases over time [12,65,68].

**Prevention**

Although most cases of ASP may be unavoidable, some of these postoperative changes might be technique dependent. Incorrect needle placement during intraoperative radiographic-level confirmation was reported to increase the relative risk of developing ASP by three-fold [50]. The plate-to-disc distance is important, with separation by <5 mm driving the development of moderate or severe ossification of the adjacent segments [64,65]. Both of these studies highlight the importance of avoiding the soft tissue injury at the adjacent level. The disc has a poor capacity for regeneration and any significant injury will likely result in accelerated degeneration. Therefore, a spine surgeon must be particularly careful to discreet decision of the operative range through careful preoperative assessments, minimal exposure of the operation field, use of a short anterior cervical plate, decrease of surrounding soft tissue damage like anterior longitudinal ligament, needle placement in vertebral body during intraoperative radiographic-level confirmation and preservation of lordosis due to plate bending and delicate wedging of bone grafts during anterior cervical fusion.

**Conclusions**

Anterior cervical fusion is the preferred surgical procedure for symptomatic cervical spinal disease and cervical spondylosis. However, ASP after fusion is a common effect of this procedure and highly morbid condition, but remains a poorly understood. Surgeons should avoid unnecessary disruption of the musculoligamentous structures along the anterior vertebral border, reduce the risk of direct injury to the disc during dissection and maintain a safe margin between the plate edge and adjacent vertebrae. Various and continuous researches will improve the quality of clinical data on ASP and elucidate the true etiology and incidence of this condition.

**Conflict of Interest**

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

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