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Biomechanical Determination of Distal Level for Fusions across the Cervicothoracic Junction

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Introduction

Adjacent segment pathology is known to occur in ~25% of patients who have undergone anterior cervical diskectomy and fusion within 10 years of the operation. This postoperative degeneration occurs at a steady rate of ~2.9% of patients per year. The cause of this early adjacent disk degeneration has not been fully elucidated. Currently, two differing ideas exist as to what causes this degeneration.

Some groups hypothesize that eliminating motion of spinal segments leads to increased forces on the adjacent segments. Other groups suggest that the disk degeneration is simply part of a natural process.

Eliminating motion by performing a fusion has been shown to increase the intradiscal pressure at the adjacent levels in biomechanical models. Eck et al simulated a fusion at C5–C6 and reported increases in the intradiscal pressures at both adjacent levels.
levels during flexion when compared with the prefusion values. Using a model reproducing a two-level C5–C7 fusion, Park et al showed increases in the intradiscal pressure at C4–C5 after fusion. Increases in the cervical intradiscal pressure after adjacent-level fusion, however, are not universally reproducible.

Patients who had long cervical fusions down to C7 may experience degeneration at the level adjacent to the most caudally fused segment. The purpose of our study is to determine whether long cervical and cervicothoracic fusions increase the intradiscal pressure at the adjacent caudal disk. Furthermore, we hoped to determine which thoracic end vertebra causes the least amount of increase in the adjacent-level intradiscal pressure. This may provide information about which level is the most ideal to end a fusion crossing the cervicothoracic junction. We hypothesized that extending past T1 and ending the fusion in the upper thoracic spine would provide a more biomechanically sound construct because of the added stability from the rib cage. Our hope is that this knowledge will enable surgeons to perform selective posterior fusions to levels that are better able to accommodate the stiff lever arm of a cervical fusion.

**Methods**

Six fresh frozen cadaveric cervicothoracic spine specimens with intact rib cages and sternums were dissected. The extraneous soft tissue was removed and care was taken to preserve the costovertebral joints, costosternal joints, facet joints, capsular ligaments, interspinous ligaments, supraspinous ligament, ligamentum flavum, anterior longitudinal ligament, and posterior longitudinal ligament in each specimen. The atlas and occiput were removed, and the specimens were stabilized caudally by the vertebral column and pelvis were not included in these specimens to focus on the thoracic vertebrae. The amputated rib cages and sternums were dissected. The extraneous ligaments, interspinous ligaments, supraspinous ligament, ligamentum flavum, anterior longitudinal ligament, and posterior longitudinal ligament in each specimen. The atlas and occiput were removed, and the specimens were stabilized caudally by the vertebral column and pelvis were not included in these specimens to focus on the thoracic vertebrae. The amputated rib cages and sternums were dissected. The extraneous

The cervical spine was then plated anteriorly from C3 to C7 using an anterior cervical plate (Stryker Spine, Kalamazoo, Michigan, United States). We simulated a common clinical scenario with previous multilevel anterior fusions ending at C7 resulting in degeneration at the cervicothoracic junction. The C7–T1 disk pressure was recorded again in the neutral position and after application of bending forces in four positions: flexion, extension, and left and right bending. Lateral mass screws were then placed in C5 and C6 and attached to pedicle screws in T1. Four points of fixation posteriorly above the cervicothoracic junction were deemed sufficient fixation after a previous anterior fusion. All the pedicle screws were placed in a straight-ahead trajectory using a freehand technique. The T1–T2 disk pressures were recorded again in all five positions. Sequential extension of the posterior construct was performed using polyaxial pedicle screws one level at a time (Stryker Spine; Fig. 1). The intradiscal pressures were measured at the subjacent disk from C7–T1 to T9–T10.

The changes in the intradiscal pressure for each level were calculated using the formula: (post bending value – pre neutral value) – (pre bending value – pre neutral value), where post is postinstrumentation and pre is preinstrumentation. The changes at each level for flexion, extension, and lateral bending were combined and a post hoc analysis was performed.

**Results**

The changes in the intradiscal pressure for each level are shown in graph format in Figs. 2 and 3 for flexion-extension and lateral bending, respectively. There were no significant differences (p = 0.13 to 0.89) when comparing the changes of the juxtafusion intradiscal pressure at each level for the separate forces of flexion, extension, and left and right bending. In flexion, the largest mean increase in the intradiscal pressure occurred at the T6–T7 level, and in extension the largest increase occurred at T8–T9. In left and right bending together, the largest increase occurred at T9–T10.

Post hoc analysis revealed decreases in the intradiscal pressure when combining all forces at T2–T3 and T7–T8. In an exploratory analysis, a t test comparing the combined changes in the intradiscal pressure for flexion, extension, and lateral bending at T7–T8 to other levels did not show a significant difference (−30.8 kPa versus −16.1 kPa p = 0.76). A t test comparing combined the intradiscal pressure changes at T2–T3 to other levels did show a significant difference (−141.0 kPa versus −3.3 kPa, p = 0.005).

**Discussion**

Adjacent segment pathology is a general term applicable to both the radiographic and clinical (CASP) changes at a segment adjacent to a previously operated spinal motion segment. Recently, Lawrence et al performed a systematic review to determine the risk of adjacent segment pathology in the
They found the prevalence of CASP to range from 11 to 12% at 5 years, 16 to 38% at 10 years, and 33% at 17 years. They also found factors that may contribute to the development of CASP including age less than 60 years, fusing adjacent to C5–C6 and/or C6–C7 levels, a pre-existing disk herniation, and/or dural compression secondary to spinal stenosis. Multiple biomechanical studies have focused on the effects of multilevel fusions on the adjacent levels. Prasarn et al demonstrated an increase in the sagittal range of motion of 31.3% above and 33.9% below a fused motion segment after comparing a two-level with a one-level anterior cervical disectomy and fusion. Another biomechanical study by Dmitriev et al also showed an increased range of motion at the distal level following C3–C6 arthrodesis.

Quantifying mechanical demands on intervertebral disks is important because these increased demands are known to have adverse effects on the disk. The stress on the disk can interfere with its nutritional requirements and cause early degeneration. The intervertebral disk relies on the diffusion of nutrients from the peripheral blood vessels and vertebral end plates through the extracellular matrix because it lacks a true blood supply. Therefore, an increase in pressure may interfere with the entrance of nutrients to the intervertebral disk as well as the escape of waste products. The accumulation of waste products leads to increased levels of lactate, which results in increased pH and impaired metabolism. Normal aging processes such as an increase in collagen type 1 and a decrease in collagen type 2, proteoglycans, and chondroitin sulfate are also accelerated with increased pressure on the intervertebral disk, and these changes lead to disk degeneration.

Theoretically, fusing caudally into the thoracic spine may result in less change in the adjacent-level intradiscal pressure due to the inherent stability of the ribs. A fusion into the thoracic spine would also avoid ending a construct at the cervicothoracic junction, an area of potential instability where the lordotic and mobile cervical spine meets the rigid and kyphotic lumbar spine. Longer constructs, however, increase the lever arm and may overcome the stability of the ribcage, increasing the adjacent-level pressure. The compromise between these two concepts may have a balancing effect on the intradiscal pressure changes. Our hypothesis was that these opposing biomechanical

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Fig. 1 Anterior, posterior, and lateral views of an instrumented specimen.

Fig. 2 Graph showing the absolute change between preinstrumented and postinstrumented adjacent-level intradiscal pressure at each level between C7–T1 and T9–T10 for flexion and extension.

Fig. 3 Graph showing the absolute change between preinstrumented and postinstrumented adjacent-level intradiscal pressure at each level between C7–T1 and T9–T10 for lateral bending.
concepts would create an equilibrium that would allow for lower-pressure changes in the upper thoracic spine.

Our outcomes contrast with some previously published results. In the cervical spine, Eck et al showed an increase in mobility and intradiscal pressure in spinal segments adjacent to fusions during the simulation of physiologic movements. Specifically, the segmental motion and intradiscal pressure during flexion and extension were measured at C4–C5 and C6–C7 before and after the fusion of C5–C6 by anterior plating. The results show significant increases in pressure during flexion at both the C4–C5 level (73.2%) and the C6–C7 level (45.3%). In our model, we did not observe significant changes in the pressure after adjacent-level fusion. Our study differed from the Eck model in several important ways. Notably, our bending was force-controlled and the Eck team used a displacement-controlled model. The authors did not report the prefusion and postfusion bending moments. We also started our pressure measurements at the cervicothoracic junction and proceeded caudally, and the analyzed intradiscal pressures in the Eck study were centered at the midcervical spine.

In concordance with our study, increases in the cervical juxtafusion intradiscal pressures have not been universally reproducible in biomechanical models. Rao et al simulated C5–C6 grafted fusion and C5–C6 anterior plating. Like in our study, they observed high interspecimen variability in the intradiscal pressure measurements. They reported a significant reduction in the intradiscal pressure of the superior adjacent disk after both grafted fusion and anterior plating. In extension, there were no significant changes in the intradiscal pressure at the inferior adjacent disk. In our model, we did not see significant increases in the intradiscal pressure at C7–T1 with any bending motion after simulating a fusion from C3–C7. That being said, the largest increases in intradiscal pressure occurred with the constructs ending from T6 to T9. Surgeons may consider ending constructs that cross the cervicothoracic junction in the upper thoracic spine rather than the mid to lower thoracic regions. In fact, the lowest combined pressures were measured at T2–T3. This may be due to a balance between the increasing stress of a longer lever arm with the increased stability of the thoracic ribs. In situations where there is lack of focal kyphosis or significant instability, it may be feasible to end constructs at T2.

This study had several weaknesses. There are inherent limitations that come with cadaveric biomechanical studies. Our model could not account for the changes that the surgical dissection and instrumentation have on living tissue nor could it incorporate the effects of active muscular control. In addition, we did not measure the changes in the motion or facet pressures at the levels adjacent to our fusion. The adjacent-level motion changes and facet loading may contribute to early disk degeneration. Treating physicians must also take into account individual patient anatomy that may influence the choice of caudal fusion levels, such as a kyphotic deformity, listhetic instability, or pre-existing disk/facet arthropathy. In fact, we found that some of the specimens had significant degeneration at multiple levels of the thoracic spine leading to asymmetric collapse, osteophyte formation, and facet hypertrophy on one side more than the other without significant deformity. This may explain some of the variation in values between left and right bending at specific levels. Also, simulating a multilevel anterior fusion ending at C7 may not be biomechanically equivalent to a posterior fusion construct ending at C7, and future biomechanical studies should be performed for this clinical scenario. Finally, we did not load our specimens to failure, therefore we could not comment on other causes of failure of long cervicothoracic fusions such as caudal-level instrumentation pullout or breakage.

Conclusion

We did not find a significant difference in intradiscal pressure changes when analyzing flexion, extension, and lateral bending separately. However, in an exploratory analysis, the combined change in the juxtafusion intradiscal pressure for flexion, extension, and lateral bending at T2–T3 was significantly less than the combined pressure at other levels. Although this change needs to be confirmed with future studies also analyzing facet pressures and three-dimensional motion, it may reflect a balance between the rigidity afforded by the rib cage in the thoracic spine and the length of the lever arm of the construct. When crossing the cervicothoracic junction with long cervical fusions, surgeons may want to consider stopping at the T2–T3 level if there are no other mitigating factors.

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Note
Of note, lateral mass screws were used in this study and are not U.S. Food and Drug Administration–approved for this indication.

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References