Iatrogenic Spinal Cord Injury Resulting From Cervical Spine Surgery.

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Iatrogenic Spinal Cord Injury Resulting From Cervical Spine Surgery

Abstract

Study Design: Retrospective cohort study of prospectively collected data.

Objective: To examine the incidence of iatrogenic spinal cord injury following elective cervical spine surgery.

Methods: A retrospective multicenter case series study involving 21 high-volume surgical centers from the AOSpine North America Clinical Research Network was conducted. Medical records for 17,625 patients who received cervical spine surgery (levels from C2 to C7) between January 1, 2005, and December 31, 2011, were reviewed to identify occurrence of iatrogenic spinal cord injury.

Results: In total, 3 cases of iatrogenic spinal cord injury following cervical spine surgery were identified. Institutional incidence rates ranged from 0.0% to 0.24%. Of the 3 patients with quadriplegia, one underwent anterior-only surgery with 2-level cervical corpectomy, one underwent anterior surgery with corpectomy in addition to posterior surgery, and one underwent posterior decompression and fusion surgery alone. One patient had complete neurologic recovery, one partially recovered, and one did not recover motor function.

Conclusion: Iatrogenic spinal cord injury following cervical spine surgery is a rare and devastating adverse event. No standard protocol exists that can guarantee prevention of this complication, and there is a lack of consensus regarding evaluation and treatment when it does occur. Emergent imaging with magnetic resonance imaging or computed tomography myelography to evaluate for compressive etiology or malpositioned instrumentation and avoidance of hypotension should be performed in cases of intraoperative and postoperative spinal cord injury.
Study Rationale and Context
Iatrogenic spinal cord injury resulting from elective cervical spine surgery is a rare and devastating adverse event. The incidence of iatrogenic spinal cord injury following cervical spine operations is challenging to determine. Flynn reviewed 82,114 anterior cervical spine operations and documented a postoperative neurological injury rate of 0.3%. Lee et al examined 1445 anterior cervical spine surgery patients and reported a rate of 0.1% spinal cord injury with neurological deficit.

Some instances of postoperative neurological deficit following cervical spine surgery can be predicted by neuromonitoring changes during the procedure, or due to an obvious intraoperative event leading to injury of the spinal cord, while others may only be recognized when the patient emerges from anesthesia. Each of these scenarios requires a unique response from the surgeon.

Given the low incidence, it is not surprising that clearly defined protocols to manage interoperative spinal cord injury during elective cervical spine surgery have not been developed. Most spine surgeons will encounter this adverse event once or twice, if at all, in an entire career. Given these small numbers, appropriate practices must be determined more based on consensus rather than data. A review of cases of iatrogenic spinal cord injury might serve to inform development of a plan for response to such events.

Although the incidence is rare, the impact of iatrogenic spinal cord injury resulting from cervical spine surgery is substantial and has potential for serious patient, surgeon, institutional, and medicolegal ramifications. The purpose of this investigation is to examine the rate of iatrogenic spinal cord injury associated with cervical spine surgery and to report patient and surgical factors associated with these injuries.

Objective or Clinical Question
This study aimed to evaluate the incidence and factors associated with iatrogenic spinal cord injury during elective cervical spine surgery.

Methods
We conducted a retrospective multicenter case series study involving 21 high-volume spine surgical centers from the AOSpine North America Clinical Research Network, selected for their clinical research infrastructure and experience. Medical records for 17,625 patients who received cervical spine surgery (levels from C2 to C7) between January 1, 2005, and December 31, 2011, were reviewed to identify occurrence of 21 predefined treatment adverse events.

Adverse events examined included reintubation requiring evacuation, esophageal perforation, epidural hematoma, C5 palsy, recurrent laryngeal nerve palsy, superior laryngeal nerve palsy, hypoglossal or glossopharyngeal nerve palsy, dural tear, brachial plexopathy, blindness, graft extrusion, misplaced screws requiring reoperation, anterior cervical infection, carotid artery injury or cerebrovascular accident, vertebral artery injuries, Horner’s syndrome, thoracic duct injury, quadriplegia, intraoperative death, revision of arthroplasty, and pseudomeingocele. This investigation examined only patients with quadriplegia following surgery.

Trained research staff at each site abstracted the data from medical records, surgical charts, radiologic imaging, narratives, and other source documents for the patients who experienced one or more of the adverse events from the list. Data were transcribed into study-specific paper case report forms. Copies of case report forms were transferred to the AOSpine North America Clinical Research Network Methodological Core for processing, cleaning, and data entry.

Results
Three cases of quadriplegia were reported from 12,903 patients. Incidence rates of the participating centers ranged from 0.0% to 0.24%.

Of the 3 patients suffering iatrogenic spinal cord injury, 2 were male and 1 was female. The mean age was 57.3 years, with an average hospital length of stay of 12 days. One injury occurred in 2007 and 2 occurred in 2011. All 3 were nonsmokers. The diagnosis and reason for surgery was myelopathy for 2 patients and degenerative disk disease for 1 patient.

One patient underwent anterior surgery only with 2-level cervical corpectomy (C5, C6), one underwent posterior surgery only, and one underwent circumferential surgery (anterior and posterior) including cervical corpectomy. Two patients underwent surgery from C3 to C7, while one patient underwent surgery from C4 to C7. All 3 patients had interoperative neuromonitoring (IONM) utilized during the procedure. Poor baseline neuromonitoring signals were noted in one patient, no baseline motor response was noted in another, with data unknown from the third patient.

In patient 1, a 67-year-old patient who underwent 2-level anterior corpectomy of C5 and C6, a dural defect was identified during resection of ossification of the posterior longitudinal ligament (PLL) with subsequent neuromonitoring change following its removal. The dural defect was covered then with a Duragen patch, followed by graft placement. The patient underwent the remaining portion of the surgical procedure prior to closure. The patient had a partial recovery but had residual upper and lower extremity weakness at follow-up.

Patient 2 was a 36-year-old patient who underwent both transcranial motor evoked potential and somatosensory evoked potential monitoring. At the outset of the case, the patient had no motor response on monitoring. The patient underwent
Response to Interoperative Neuromonitoring Alerts

Neuromonitoring utilizing somatosensory evoked potentials and transcranial motor evoked potentials (tcMEPs) is frequently used in cervical spine surgery and may help surgeons intervene to reverse the immediate cause of intraoperative spinal cord injury.9,10 For procedures performed in the prone position, obtaining potentials with the neck in a neutral posture (prior to prone positioning) may be beneficial in some cases to provide baseline neurophysiologic data. Potentials can then be repeated in the prone surgical position to help identify cervical positioning related neuromonitoring alterations.

Some instances of intraoperative neuromonitoring changes occur due to spinal cord hypoperfusion.2,3,8,11,12 Spinal cord oxygenation and perfusion are known to correlate with neuromonitoring alerts. Direct correlation between cerebrospinal fluid (CSF) oxygenation and TcMEPs has been shown in a pig model with clamping of spinal radicular arteries, with reversal of these neuromonitoring changes following unclamping of the vessels.13 In a canine study, multiple bilateral spinal radicular vessel ligation was required to create irreversible neurological deficit.14 In human studies examining neuromonitoring changes during scoliosis surgery, neuromonitoring changes associated with hypotension are often reversible with mean arterial pressure (MAP) elevation, and do not lead to permanent postoperative neurological deficit in the majority of situations.15 In cases of neuromonitoring changes without an obvious reversible surgical explanation, evaluation of blood pressure and correction of hypotension if possible should be undertaken.

Literature regarding the utility of neuromonitoring during cervical spine surgery is relatively limited. An investigation by Clark et al16 retrospectively reviewed 140 patients with cervicothoracic spondylotic myelopathy undergoing spine surgery, of which 16 (11%) had intraoperative decreases in tcMEPs. In total, there were 8 patients from this group who awoke with neurological deficits: 5 with C5 palsy and 2 with paraparesis. A significant correlation (P < .001) was found between persistent tcMEP changes and postoperative neurological deficits, with a sensitivity of 75%, specificity of 98%, positive predictive value of 75%, and a negative predictive value of 98%. In patients with vascular disease, the sensitivity of tcMEPs decreased to 60%.

Although neuromonitoring may be able to predict some cases of postoperative neurological deficit, the appropriate response by the surgeon, anesthesia staff, and neurophysiologist is not clear in many cases. Ziewacz et al designed a checklist for responding to neuromonitoring changes during spinal myelopathy and deformity spine surgery in 2012.17 They utilized expert consensus and aviation and surgical literature to create their algorithm (Figure 1), which highlights initial logical responses to MEP changes as well as additional considerations if the MEPs do not respond to initial interventions. Surgeon responses recommended include stopping the current manipulation, assessing the field for structural spinal cord compression, and consideration for further spinal cord decompression and stenosis is present.

Although there is relatively little literature specific to neuromonitoring changes during cervical spine surgery, there is a large body of work regarding thoracolumbar spinal deformity surgery, which may be informative to cervical spine surgery.10 The incidence of spinal cord injury has been reported to occur in 0.26% to 1.75% of thoracolumbar deformity operations.11,18 The surgeon and surgical team response to neuromonitoring changes in spinal deformity surgery have been thoroughly evaluated in a Delphi Consensus Report by Vitale et al.10 In this investigation, they separated the “mechanically stable spine” from the unstable spine following spinal osteotomy as appropriate response in these 2 situations differs substantially. This
delineation may similarly be useful for cervical spine surgery, in which cervical spine trauma or spinal osteotomy may require a specific surgeon response to neuromonitoring changes. Although designed for thoracolumbar deformity surgery, the results of this Delphi Report provide a valuable guide for response to neuromonitoring alerts during cervical spine surgery. Recommended responses to neuromonitoring changes include an intraoperative pause, summoning the attending

**Figure 1.** Checklist for neuromonitoring (MEP) alerts in patients with myelopathy or spinal deformity. From Ziewacz et al.17

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**Checklist for Neuromonitoring (MEP) Alert in Patients with Myelopathy or Deformity**

**Spine Surgeon:**
- Stop current manipulation
- Assess field for structural cord compression (misplaced hardware or bone graft, osteophytes, or hematoma)
- Perform further decompression if stenosis is present
- Consider reversing correction of a spinal deformity

**Neurophysiologist:**
- Repeat trials of MEPs and SSEPs to rule out potential false positive
- Check all leads to make sure no pull-out, may add leads in proximal muscle groups if possible
- Assess the pattern of changes
  - Asymmetric changes (associated with cord or nerve root injury)
  - Symmetric changes (associated with anesthetic or hypotension issues)
- Quantify improvement and communicate to the surgical team

**Anesthesiologist:**
- Check if neuromuscular blockade (muscle relaxant) given
- If yes, check train of four (TOF)
- Verify that no change in anesthetic administration occurred
- Assess anesthetic depth
  - BP
  - RR
  - HR
  - BIS monitor (if available)
- Restore or maintain blood pressure (goal mean arterial pressure of 90-100)
- Check Hemoglobin/Hematocrit (goal hemoglobin >9-10)
- Check temperature and I/O’s for adequate resuscitation
- Check extremity position in case of plexus palsy
- Lighten depth of anesthesia
  - Reduce to 1/3 MAC or temporarily eliminate inhaled agents (i.e. desflurane)
  - Reduce intravenous anesthetics such as propofol (which may accumulate systemically during the case and blunt MEPs)
  - Add adjuvant agents such as Ketamine to permit reduction of MEP suppressive agents (i.e. propofol and inhalational anesthetics)

**IF No Change:**
- Increase MAP >100
- Consider Steroid Administration
- Consider Wake-up test
- Consider Aborting surgery
- Consider Calcium Channel Blocker (topical to cord or iv)

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*The checklist assumes baseline anesthetic regimen is 1/3-1/2 MAC of halogenated anesthetic (desflurane) and TIVA (total intravenous anesthesia) with propofol +/- ketamine.*

**Fig. 2.** Checklist for the response to an intraoperative neuromonitoring alert. BIS = bispectral index; BP = blood pressure; HR = heart rate; I/O = input/output; MAC = minimum alveolar concentration; MAP = mean arterial pressure; MEP = motor evoked potential; RR = respiration rate; SSEP = somatosensory evoked potential.
anesthesiologist and senior neurophysiologist, determination of the need for intraoperative imaging, optimization of patient MAPs, hematocrit, pCO₂, and temperature, consideration of a wakeup test, checking technical neuromonitoring factors, and evaluation and correction of surgical factors (ie, remove traction, remove instrumentation or bone graft, etc; Figure 2).

While it is not clear that a separate checklist for IONM alerts during cervical spine surgery is needed, the work done by Ziewacz et al¹⁷ and Vitale et al¹⁰ would serve as an excellent starting point. The review of cases here suggests that a standardized approach to use of IONM, as well as response to alerts, is not part of current practice among cervical spine surgeons as the monitoring strategy differed in each of the 3 cases presented in this investigation.

**Response to Postoperative Motor Deficit**

A separate but related issue is how the operating surgeon should respond when a patient awakens from anesthesia with a new motor deficit of clinical significance. The optimal postoperative management following iatrogenic cervical spinal cord injury should generally include emergent MRI or computed tomography (CT) myelogram to evaluate for spinal cord compression from hematoma, bone graft, vertebral displacement, or malpositioned instrumentation. If a compressive etiology is discovered, return to the operating room for alleviation of the cause of neural compression is indicated at the earliest possible opportunity that the patient can safely tolerate.

Additionally, avoidance of hypotension with induced hypertension is recommended in cases of acute spinal cord injury of any etiology. Keeping MAPs >85 mm Hg has been shown to improve motor function and bowel/bladder recovery following traumatic spinal cord injury,¹⁹ and may be performed for up to 7 days, although some centers perform only 48 to 72 hours of MAP elevation. Optimizing spinal cord oxygenation and avoiding hypotension are important interventions in optimizing outcomes following iatrogenic spinal cord injury.

The neurological sequelae of traumatic spinal cord injury occurs due to an initial traumatic mechanical injury followed by secondary insult stemming from ischemia, reperfusion, ionic dysregulation, cellular excitotoxicity, swelling, and free-radical–mediated peroxidation.²⁰ Numerous prospective human studies have been performed to investigate pharmacologic interventions to reverse the deleterious inflammatory response and neurological deficits from traumatic spinal cord injury, although unfortunately none have proven dramatically successful thus far. Therefore at this time, no strong recommendations regarding steroids or other investigational medications can be made to provide to patients who suffer iatrogenic spinal cord injury resulting from cervical spine surgery.²⁰
Other strategies to mitigate spinal cord injury may exist. Placing a CSF drain is commonly performed to decrease CSF pressure in an attempt to prevent spinal cord injury during thoracoabdominal aorta surgery\(^1\); no data currently exist to examine whether this may be beneficial in cases of iatrogenic spinal cord injury during cervical spine surgery.

This review demonstrates a similar lack of a protocol-based approach to discovery of a new postoperative neurological deficit. As case numbers will be too small to develop such a protocol based on data, a consensus-based approach appears appropriate. Postoperative institutional safety improvement review of protocols and procedures are imperative following serious adverse events such as iatrogenic spinal cord injury and were likely performed in each of the cases presented in this investigation. Unfortunately, details of individual institution safety improvement initiatives were not included in our data set.

**Conclusion**

Iatrogenic spinal cord injury following elective cervical spine surgery is a rare and devastating adverse event occurring in up to 0.24% of cases in this multicenter cohort. This study was limited in its ability thoroughly assess risk factors and outcomes of this adverse event due to the rarity of the event and the small number of cases encountered. No standard protocol exists that can guarantee prevention of this complication, and there is a lack of consensus regarding evaluation and treatment when it does occur. Utilization of IONM and response to intraoperative alerts should be standardized based on surgeon consensus. Similarly, response to postoperative motor deficits is not yet protocolized. Emergent imaging with MRI or CT myelography to evaluate for compressive etiology or malpositioned instrumentation, appropriate surgical correction when appropriate, and maintenance of adequate mean arterial blood pressure should generally be performed in cases of postoperative spinal cord injury.

**Authors’ Note**

This study was ethically approved by the institutional ethics committees at all participating sites.

**Declaration of Conflicting Interests**

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**References**


