Safety profile and probe placement accuracy of intraspinal pressure monitoring for traumatic spinal cord injury: Injured Spinal Cord Pressure Evaluation study

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OBJECTIVE A novel technique for monitoring intraspinal pressure and spinal cord perfusion pressure in patients with traumatic spinal cord injury was recently described. This is analogous to monitoring intracranial pressure and cerebral perfusion pressure in patients with traumatic brain injury. Because intraspinal pressure monitoring is a new technique, its safety profile and impact on early patient care and long-term outcome after traumatic spinal cord injury are unknown. The object of this study is to review all patients who had intraspinal pressure monitoring to date at the authors’ institution in order to define the accuracy of intraspinal pressure probe placement and the safety of the technique.

METHODS At the end of surgery to fix spinal fractures, a pressure probe was inserted intradurally to monitor intraspinal pressure at the injury site. Postoperatively, CT scanning was performed within 48 hours and MRI at 2 weeks and 6 months. Neurointensive care management and complications were reviewed. The American Spinal Injury Association Impairment Scale (AIS) grade was determined on admission and at 2 to 4 weeks and 12 to 18 months postoperation.

RESULTS To date, 42 patients with severe traumatic spinal cord injuries (AIS Grades A–C) had undergone intraspinal pressure monitoring. Monitoring started within 72 hours of injury and continued for up to a week. Based on postoperative CT and MRI, the probe position was acceptable in all patients, i.e., the probe was located at the site of maximum spinal cord swelling. Complications were probe displacement in 1 of 42 patients (2.4%), CSF leakage that required wound resuturing in 3 of 42 patients (7.1%), and asymptomatic pseudomeningocele that was diagnosed in 8 of 42 patients (19.0%). Pseudomeningocele was diagnosed on MRI and resolved within 6 months in all patients. Based on the MRI and neurological examination results, there were no serious probe-related complications such as meningitis, wound infection, hematoma, wound breakdown, or neurological deterioration. Within 2 weeks postoperatively, 75% of patients were extubated and 25% underwent tracheostomy. Norepinephrine was used to support blood pressure without complications. Overall, the mean intraspinal pressure was around 20 mm Hg, and the mean spinal cord perfusion pressure was around 70 mm Hg. In laminectomized patients, the intraspinal pressure was significantly higher in the supine than lateral position by up to 18 mm Hg after thoracic laminectomy and 8 mm Hg after cervical laminectomy. At 12 to 18 months, 11.4% of patients had improved by 1 AIS grade and 14.3% by at least 2 AIS grades.

CONCLUSIONS These data suggest that after traumatic spinal cord injury intradural placement of the pressure probe is accurate and intraspinal pressure monitoring is safe for up to a week. In patients with spinal cord injury who had laminectomy, the supine position should be avoided in order to prevent rises in intraspinal pressure.

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KEY WORDS critical care; monitoring; paralysis; perfusion pressure; spinal cord injury; trauma

ABBREVIATIONS AIS = American Spinal Injury Association Impairment Scale; ASIA = American Spinal Injury Association; CPP = cerebral perfusion pressure; CPPopt = optimum cerebral perfusion pressure; ICP = intracranial pressure; ISP = intraspinal pressure; MAP = mean arterial pressure; NBD = Neurogenic Bowel Dysfunction; NICU = neurointensive care unit; SCIM III = Spinal Cord Independence Measure III; SCPP = spinal cord perfusion pressure; SCPPopt = optimum spinal cord perfusion pressure; TBI = traumatic brain injury; TSCI = traumatic spinal cord injury; WISCI II = Walking Index for Spinal Cord Injury.
Intraspinal pressure monitoring

In 2008, we showed that intraspinal pressure (ISP) is elevated after TSCI at the injury site in mice. We subsequently introduced a technique to monitor ISP in patients with TSCI and thus compute spinal cord perfusion pressure (SCPP) as MAP minus ISP. Our analysis shows that the ISP and SCPP waveforms after TSCI have similar characteristics as the ISP and CPP waveforms after TBI, respectively. We, therefore, proposed that the concepts of pressure reactivity (quantified using PRx), optimum cerebral perfusion pressure (CPPopt), and compensatory reserve (quantified using RAP), which had been developed for TBI, could also be applied to the damaged spinal cord to yield spinal pressure reactivity (sPRx), optimum spinal cord perfusion pressure (SCPPopt), and spinal compensatory reserve (sRAP). After TSCI, bony decompression (i.e., realignment of the fracture and laminectomy) does not effectively reduce the elevated ISP at the injury site. Our data showed that in addition to bony decompression, expansion duraplasty was also required to effectively reduce ISP, increase SCPP, and improve the pressure reactivity at the injury site. Laminectomy with duraplasty after TSCI is analogous to decompressive craniectomy after TBI, which involves not only bony but also dural decompression. The dura is a major, but unappreciated, cause of spinal cord compression after TSCI and may explain why the benefits of early bony decompression (without opening the dura) after TSCI remain controversial.

To monitor ISP, we place a pressure probe intradurally at the injury site using a microscope at the end of posterior spinal fixation. Our data suggest that, at the injury site, subdural ISP is the same as intraparenchymal ISP. It is important to note that ISP at the injury site, where the spinal cord is swollen and compressed against the dura, is different from subdural pressure above or below the injury site and extradural pressure. The positioning of the ISP probe is, therefore, crucial for accurate measurement of ISP. Since ISP monitoring is invasive there are potential risks, including CSF leakage, pseudomeningocele, CSF or wound infection, spinal cord damage, and hematoma at the injury site. It is also possible that ISP monitoring compromises the care of TSCI patients indirectly, e.g., by prolonging intubation or increasing complications from inotrope use. To date, we have monitored ISP from 42 patients with severe TSCI. Here, we report the accuracy of ISP probe placement and the safety of ISP monitoring.

Patient Recruitment

Inclusion criteria are: 1) severe TSCI defined as American Spinal Injury Association (ASIA) Impairment Scale (AIS) Grade A, B, or C; 2) age 18 to 70 years; and 3) timing between TSCI and surgery not more than 72 hours. All types of spine trauma were included, including compression, distraction, and rotation. Exclusion criteria are: 1) patients unable to provide consent; 2) other major injuries or comorbidities; and 3) penetrating spinal cord injury. Surgery and early management took place at the neurosurgical unit of St. George’s Hospital, a tertiary referral center that covers a population of about 2.5 million in southwest London. Recruitment into the study was discussed with eligible patients and their families on admission, and the patient information sheet was given.

Surgical Technique

All patients underwent spinal realignment and fixation via a posterior approach. Spinal fixation involved lateral mass screws for cervical injuries and pedicle screws for thoracolumbar injuries. Four of 42 patients underwent additional anterior cervical fixation with plate and screws. For anterior cord compression in the thoracolumbar spine, the surgeon would push the bone fragments from the spinal canal back into the vertebral body via a posterior approach. Following posterior spinal fixation, a Codman Microsensor Transducer (DePuy Synthes) was tunneled through the skin with a 14-gauge introducer. The Codman probe is not labeled for ISP monitoring, as used here. The dura was perforated 1 level below the injury with a 21-gauge needle bent at 90° to prevent damage to the underlying spinal cord. After calibration, the Codman probe was advanced through the dural perforation in the subdural space under the operating microscope until the probe tip was at the level of the injury based on the measurements on the MR scan and ISP recordings. Fibrin glue (Tisseel) was placed on the dura. A wound drain was inserted superficial to the lumbar fascia and allowed to drain under gravity. The Codman probe was secured to the skin with multiple silk sutures, including a tightening stitch around the exit site to prevent CSF leakage.

Signal Recording

The ISP probe was connected to a Codman ICP box linked via an ML 221 amplifier to a PowerLab running LabChart (version 7.3.5, AD Instruments). Arterial blood pressure was recorded from a radial artery catheter connected to the Philips Intellivue MX800 bedside monitoring system (Philips), which was in turn connected to the PowerLab system. The ISP and arterial blood pressure signals were sampled at 100 Hz or 1 kHz for up to 7 days after surgery. LabChart was used to analyze the signals and compute SCPP, which was defined as MAP minus ISP.

Thromboprophylaxis

Preoperative thromboprophylaxis was achieved by using antithrombosis stockings and intermittent pneumatic foot compression. Postoperatively, in addition to compression stockings, a daily dose of 5000 U of dalteparin sodium was administered subcutaneously starting at 24

Methods

Institutional Research Board Approval

Approvals for the Injured Spinal Cord Pressure Evaluation study, including the patient information sheet and informed consent form, were obtained from St. George’s Joint Research Office and the National Research Ethics Service London—St. Giles Committee. Informed consent was obtained from all patients recruited in the study. Online details are available at http://elv1ml.co.uk/?page_id=73.
hours after surgery. The dalteparin dose was omitted for 24 hours before ISP probe removal.

Inotropes and Target MAP

Neurointensivists were blind to the ISP and SCPP readings and, therefore, no treatment was initiated in response to changes in these pressures. For the first 28 patients, MAP was treated with norepinephrine according to the neurointensivists’ preferences. In the UK, such MAP management is variable. For the next 14 patients, a norepinephrine infusion was administered to support a target MAP of more than 80 mm Hg. Serum lactate and base excess were checked via arterial blood gas analysis at least 3 times per day to assess end-organ ischemia.

Patient Positioning

Patients were turned from supine to lateral and vice versa every 2 to 3 hours to prevent the development of pressure ulcers. Lateral positioning was achieved by placing 2 or more pillows behind the patient’s back to avoid external compression to the wound. The difference in ISP recorded on the observation chart between each turn was averaged for the different injury groups: 1) cervical with laminectomy; 2) cervical without laminectomy; 3) thoracic with laminectomy; and 4) thoracic without laminectomy.

Patient Imaging

CT and MR scanning of the whole spine were done on admission. Another CT scan was obtained within 48 hours of ISP probe insertion to check the screw and probe positions. An MR scan of the injured spinal cord was obtained at about 2 weeks (i.e., after removing the ISP probe) and 6 to 12 months after surgery.

Neurological Assessments

A full neurological examination according to the International Standards for Neurological Classification of Spinal Cord Injury was performed on admission, at 2 weeks postoperatively (i.e., before discharge to the rehabilitation facility), and at the outpatient follow-up at 12 to 18 months. At the outpatient follow-up, additional information was collected using the Walking Index for Spinal Cord Injury score (WISCI II),17 Neurogenic Bowel Dysfunction (NBD) scale,18 and the sphincter management section of the Spinal Cord Independence Measure III (SCIM III) scale.19

Probe Placement Analysis

The ISP probe was visualized on the postoperative CT scan. The 2-week postoperative MR scan was then compared with the postoperative CT scan. On the 2-week postoperative T2-weighted, midline sagittal MR scan, the injury site was divided into 3 zones. Zone 1 has no CSF signal between the swollen, injured spinal cord and the dura. In Zone 2, which lies on either side of Zone 1, the thickness of the CSF space around the cord is < 50% normal. Zone 3 lies next to Zone 2, away from the injury, and has normal CSF space around the cord. We determined the probe tip position to be adequate/good in Zone 1, adequate/fair in Zone 2, and inadequate in Zone 3.

Clinical Data Collection

Neurointensive care unit (NICU) nursing charts, drug administration charts, scans, serum biochemistry, and arterial blood gas data were reviewed. Probe-related and nonrelated complications were charted prospectively and reported to the ethics committee. Patient position (supine, lateral) was recorded hourly by the NICU nurse who was looking after the patient.

Statistics

The 1-sample t-test was performed using Excel 2010 to compare the changes in ISP in the supine versus lateral position. Statistical significance was taken at $p < 0.05$.

Results

Patient Characteristics

We had 25 (59.5%) cervical, 15 (35.7%) thoracic, and 2 (4.8%) conus medullaris spinal cord injuries (Fig. 1A). The most commonly injured levels were C-5 and C-6. Two-thirds (28 of 42) of the patients were younger than 50 years, and the most frequently injured group was 30- to 40-year-old individuals (Fig. 1B). Overall, the male-to-female ratio was about 3:1: the male-to-female ratio was higher for patients younger than 50 years (7:1), but females predominated with a male-to-female ratio of 2:5 in those older than 50 years. Figure 1C shows the severity of TSCI for each spinal level. Thirteen of 25 (52.0%) cervical TSCIs were classified as AIS Grade A on admission compared with 13 of 15 (76.7%) thoracic TSCIs and 2 of 2 (100%) conus medullaris TSCIs. Overall, 24 of 42 (57.1%) patients had a fall and 18 of 42 (42.9%) had a road traffic accident (Fig. 1D). The most common cause of injury was a fall in cervical TSCI patients (18 of 25 patients; 72.0%) and a road traffic accident in thoracic TSCI patients (10 of 15 patients; 66.7%). Figure 1E shows the hours from injury to surgery, which ranged from 9 to 72 hours with a median of 37 hours.

Probe Position

Figure 2A and B illustrate the method used to determine the probe position and the definition of the 3 zones in relation to the CSF space around the injured spinal cord. Probe position was adequate in all patients, good in 82%, and fair in 18% (Fig. 2C).

NICU Management

On average, ISP was approximately 20 mm Hg, MAP was approximately 90 mm Hg, and SCPP was approximately 70 mm Hg during the 7-day monitoring period, with considerable interpatient variability (Fig. 3A). In patients who underwent laminectomy, ISP was significantly higher in the supine compared with the lateral position (Fig. 3B). The average difference in ISP between supine versus lateral patient position ($\Delta$ISP) was approximately 2 mm Hg for patients with thoracic laminectomies and 1 mm Hg for patients with cervical laminectomies. After thoracic laminectomies, $\Delta$ISP was up to 18 mm Hg, and $\Delta$ISP was up to 8 mm Hg after cervical laminectomies. An increase in ISP while lying supine was observed in all
intraspinal pressure monitoring

Laminectomy patients, regardless of the type of fracture and regardless of anterior decompression. Postoperative MRI ruled out the presence of hematoma compressing the injured spinal cord. Within 2 weeks, all patients were extubated, though about a quarter received a tracheostomy (Fig. 3C). Tracheostomy was required in 28% of patients with cervical injuries compared with 13% patients with thoracic injuries. Two of 9 (22.2%) patients who had a tracheostomy had lung contusions and/or pneumothorax that was evident on the chest CT scan done on admission. Overall, 9.5% (4 of 42) of our patients required a chest drain. Figure 3D–F summarize the use of norepinephrine, as well as the base excess and lactate concentrations as determined from the arterial blood samples.

Complications

Table 1 lists the complications, including those associated with probe insertion as well as other complications. In 3 of 42 patients, we observed CSF leakage from the probe skin site once the probe was removed. In all patients, the CSF leaks settled after suturing the skin. In 19% of patients, we observed a pseudomeningocele on the MR scan obtained at 2 weeks after surgery, which disappeared in all patients on the MR scan obtained at 6 months. The pseudomeningocele was a radiological complication and did not appear to compress the spinal cord in any of the patients. There were no serious probe-related complications such as meningitis, wound infection, or spinal cord hematoma. Table 1 also lists the complications not associated with probe insertion. Such complications were related to the initial trauma, immobility, and intubation/ventilation. The only technical problem associated with data collection was the occasional disconnection of the monitoring equipment.

Neurological Outcome

We assessed neurological outcome using the AIS grade when the patients were discharged from the neurosurgery unit (Table 2) and at 12 to 18 months later (Table 3). Overall, about 10% of our patients improved by the time they were discharged from neurosurgery: 7.1% of patients improved by 1 AIS grade, and 2.4% improved by 2 AIS grades. At 12 to 18 months, about a quarter of our patients improved neurologically: 11.4% of patients improved by 1 AIS grade, and 14.3% improved by 2 or more AIS grades. Improvement by at least 1 AIS grade was inversely correlated with AIS on admission: improvement was seen in 8.3% of AIS Grade A patients compared with 63.6% of AIS Grade B or C patients. The Supplemental Data show additional outcome measures at 12 to 18 months based on the sphincter management section of SCIM III, NBD, and WISCI II.

![Figure 1](https://example.com/figure1.png)

Discussion

Our key finding is that ISP monitoring is safe. Probe-related complications were CSF leakage from the probe’s skin exit site that resolved with suturing and asymptomatic pseudomeningocele that resolved within 6 months. We did not encounter serious complications such as meningitis or spinal cord damage. By comparison, ICP monitoring for brain injury causes intracerebral hematoma in 1.1% to 4.1% of patients and CSF infection in 0.6% to 7.4% of patients. ISP monitoring may have a better profile for serious complications than ICP monitoring because the ISP probe is subdural and inserted under direct vision using a microscope, whereas the ICP probe is intraparenchymal and inserted blindly through a twist-drill craniostomy. After the ISP probe was removed, CSF-related complications that were easily treatable or did not require treatment occurred in 28.8% of patients. In contrast, CSF-related complications are uncommon after the ICP probes are removed. In all patients, the AIS grade at 2 weeks was the same or better than preoperatively. Therefore, spinal fixation surgery, including probe insertion, did not cause neurological deterioration.

We also assessed NICU management and related complications. As expected for intubated patients, chest infections were common. Though most of our patients were extubated within 2 weeks, a quarter required a tracheostomy. In general, tracheostomy was performed in patients with cervical spinal cord injuries and those with lung contusions. These findings are comparable with published series in which the mean time to extubation was 5 days and 81% of complete cervical SCI patients required tracheostomy. We intentionally did not set ISP or SCPP targets in order to study the injured spinal cord over a range of ISPs and SCPPs. Norepinephrine was used to support blood pressure during 75% (3096 of 4108 hours) of monitored hours without associated complications. By comparison, Inoue et al. reported complications with dopamine or phenylephrine usage such as arrhythmias, acidosis, and skin necrosis. In our laminectomized patients,
ISP was significantly higher in the supine than the lateral position by up to 8 mm Hg for cervical injuries and 18 mm Hg for thoracic injuries. This finding has important implications for nursing care: we recommend avoiding wound compression (e.g., by avoiding the supine position or by placing a ring-shaped pillow around the wound). Together, our data suggest that placing an ISP probe may help guide NICU medical and nursing management without major complications.

Even though the ISP probe may not cause detectable spinal cord damage during the monitoring period, the probe may compromise the long-term recovery of patients. We, therefore, determined the AIS grades of the patients who had ISP monitoring at 12 to 18 months following TSCI. Our data show that the improvements in AIS grades in our patients were comparable to the improvements reported in other series. We did not expect improved recovery in our patients, in comparison with published series, because our ISP monitoring study was observational without set ISP or SCPP targets. Though our data suggest that placing an ISP probe may help guide NICU medical and nursing management without major complications.

**TABLE 2. AIS on admission versus at 2 weeks after surgery**

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<th>AIS Grade on Admission</th>
<th>AIS Grade at 2 Wks After Surgery*</th>
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<tr>
<td></td>
<td>A</td>
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<tr>
<td>A</td>
<td>27 (93.1)</td>
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<tr>
<td>B</td>
<td>0 (0.0)</td>
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<tr>
<td>C</td>
<td>0 (0.0)</td>
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* Values are shown as the number of patients (%).

**TABLE 3. AIS on admission versus 12 to 18 months after surgery**

<table>
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<th>AIS Grade at 12–18 Mos After Surgery*</th>
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<tr>
<td>A</td>
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<tr>
<td>22 (91.7)</td>
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<tr>
<td>0 (0.0)</td>
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<td>0 (0.0)</td>
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* Values are shown as the number of patients (%). Seven of 42 patients have not reached the 12- to 18-month follow-up, and therefore only 35 of 42 patients are included.
ing an ISP probe is safe, a randomized controlled trial is necessary to definitively conclude that the probe did not compromise long-term recovery.

In earlier studies,18,19,26 we showed that, after TSCI, 3 intradural compartments form with different pressure profiles: above the injury, at the injury site (where the swollen cord is compressed against dura), and below the injury site. The ISP probe should be placed at the site of injury to monitor the maximum ISP. Technical nuances include measuring the insertion distance on the MR scan and monitoring ISP while inserting the probe. The postoperative CT scans show that ISP probe placement accuracy was adequate in all patients. Intraoperative fluoroscopy could not be used to guide probe insertion because the Codman probe was invisible on fluoroscopy. The chance of probe dislodgment was minimized by suturing the probe to the skin at multiple sites and placing a large, adherent loban drape over the probe. By placing a suture at the probe’s skin exit site, we eliminated the risk of CSF leakage around the probe.

There is controversy about whether aiming for universal CPP targets with ICP monitoring improves outcome after TBI.24 Such controversy led to the idea that CPPopt is not the same in all patients. It has been suggested that, after TBI, NICU treatment should become individualized based on each patient’s CPPopt.29 The CPPopt for a patient can be defined as the CPP that minimizes PRx: i.e., maximizes vascular pressure reactivity. There were no ISP or SCPP targets in our studies because we have not yet determined the optimal ISP and SCPP after TSCI. Currently, in the UK, arterial blood pressure management after TSCI is variable,27 whereas in the US the guidelines of the American Association of Neurological Surgeons of 85–90 mm Hg MAP are usually followed.25 Intervening to elevate the blood pressure without monitoring ISP and SCPP from the injury site is potentially dangerous due to cardiogenic complications.15 From our experience with ISP/SCPP monitoring, and by analogy with the concept of individualized CPPopt in TBI, it seems that SCPPopt varies between patients, thus suggesting individualized SCPP management rather than setting generalized guidelines.20 Currently, ICP monitoring and CPP optimization are the standard of care when managing severe TBI patients in developed countries. Based on our earlier studies19,20,24,26 and the safety profile of ISP monitoring shown here, we hope that our ISP monitoring technique is used by other centers. Ultimately, a randomized controlled trial is warranted to determine if ISP monitoring and SCPP optimization improve outcome after TSCI.

Conclusions

After TSCI, intradural placement of the pressure probe is accurate and ISP monitoring for up to a week is safe. In patients with spinal cord injury who underwent laminectomy, the supine position should be avoided to prevent rises in ISP at the injury site.

Acknowledgments

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References


Disclosures
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Author Contributions
Conception and design: Papadopoulos, Saadoun. Acquisition of data: Papadopoulos, Phang. Analysis and interpretation of data: Papadopoulos, Phang, Saadoun. Drafting the article: Papadopoulos, Phang, Saadoun. Critically revising the article: Papadopoulos, Zoumprouli, Saadoun. Reviewed submitted version of manuscript: all authors. Approved the final version of the manuscript on behalf of all authors: Papadopoulos. Statistical analysis: Phang. Study supervision: Papadopoulos, Saadoun.

Supplemental Information
Online-Only Content
Supplemental material is available with the online version of the article.

Previous Presentations
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