Continuous monitoring and visualization of optimum spinal cord perfusion pressure in patients with acute cord injury

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**Running Head:** Optimum spinal cord perfusion in acute cord injury

**Key words:** Autoregulation, Blood pressure, Critical care, Spinal cord injuries, Vasopressor agents
ABBREVIATIONS

ASIA, American Spinal Injuries Association

cCPP\textsubscript{opt}, continuous optimum cerebral perfusion pressure. Using a sliding 4-hour window that is updated every minute, PRx is plotted against cerebral perfusion pressure. cCPP\textsubscript{opt} is the cerebral perfusion pressure at which PRx is minimum, i.e. the cerebral perfusion pressure that optimizes brain vascular pressure reactivity. cCPP\textsubscript{opt} applies to individual patients and is updated every minute.

cSCPP\textsubscript{opt}, continuous optimum spinal cord perfusion pressure. Using a sliding 4-hour window, the SCPP of an individual patient is divided into 5 mmHg bins between 40 – 120 mmHg and the mean sPRx for each bin is computed. sPRx is plotted against SCPP and cSCPP\textsubscript{opt} is the SCPP at which sPRx is minimum, i.e. the SCPP that optimizes spinal cord vascular pressure reactivity. cSCPP\textsubscript{opt} applies to an individual patient and is updated every minute.

CSF, cerebrospinal fluid

igSCPP\textsubscript{opt}, individual global optimum spinal cord perfusion pressure. Using the entire SCPP signal of an individual patient, SCPP is divided into 5 mmHg bins between 40 – 120 mmHg and the mean sPRx for each bin is computed. sPRx is plotted against SCPP. igSCPP\textsubscript{opt} is the SCPP at which sPRx is minimum, i.e. the SCPP that optimizes spinal cord vascular pressure reactivity. The igSCPP\textsubscript{opt} value applies to the entire signal of an individual patient.

ISP, intraspinal pressure. Pressure of the swollen spinal cord at the injury site, as it is compressed against dura.

MAP, mean arterial pressure

MD, microdialysis
pSCPP_{opt}, pooled optimum spinal cord perfusion pressure. Using the SCPP signals of several patients, SCPP is divided into 5 mmHg bins between 40 – 120 mmHg and the mean sPRx for each bin is computed. sPRx is plotted against SCPP. pSCPP_{opt} is the SCPP at which sPRx is minimum, i.e. the SCPP that optimizes spinal cord vascular pressure reactivity. The pSCPP_{opt} value applies to the entire patient group.

SCPP, spinal cord perfusion pressure. Mean arterial pressure minus intraspinal pressure.

sPRx, spinal vascular pressure reactivity index. This is the running correlation coefficient between ISP and MAP. sPRx \leq 0 indicates intact pressure reactivity, whereas sPRx > 0 indicates impaired pressure reactivity.

TSCI, traumatic spinal cord injury
ABSTRACT

The optimum spinal cord perfusion pressure (SCPP) after traumatic spinal cord injury (TSCI) is unknown. Here, we describe techniques to compute and display the optimum SCPP in real-time. We recruited adults within 72 hours of severe TSCI (ASIA grades A–C). A pressure probe and a microdialysis catheter were placed on the injured cord. SCPP was computed as mean arterial pressure (MAP) minus intraspinal pressure (ISP), spinal pressure reactivity index (sPRx) as the running ISP/MAP correlation coefficient and continuous optimum SCPP (cSCPP\textsubscript{opt}) as the SCPP that minimizes sPRx in a moving four-hour window. In 45 patients, we monitored ISP and blood pressure. In 14 patients, we also monitored injury site metabolism. cSCPP\textsubscript{opt} could be computed 45 % of the time. Mean cSCPP\textsubscript{opt} varied by up to 60 mmHg between patients. Each patient’s cSCPP\textsubscript{opt} varied with time (standard deviation 10–20 mmHg). Color-coded maps showing the sPRx/SCPP curve evolution enhanced visualization of cSCPP\textsubscript{opt}. Periods when SCPP \( \approx \) cSCPP\textsubscript{opt} were associated with low injury site glucose, high pyruvate and high lactate. Mean SCPP deviation from cSCPP\textsubscript{opt} inversely correlated with neurological outcome at 9–12 months: ASIA grade improved in 30 % patients with <5 mmHg deviation, 10 % with 5–15 mmHg deviation and no-one with >15 mmHg deviation. We conclude that real-time computation and visualization of cSCPP\textsubscript{opt} after TSCI are feasible. cSCPP\textsubscript{opt} appears to enhance glucose utilization at the injury site and varies widely between and within patients. Our data suggest that targeting cSCPP\textsubscript{opt} after TSCI might improve neurological outcome.
INTRODUCTION

About 180,000 people suffer a traumatic spinal cord injury (TSCI) each year\(^1\). The consequences are catastrophic; most patients remain dependent or wheelchair-bound. Currently, the early management of TSCI patients in the neurointensive care unit is variable: In the U.K. there are no clear blood pressure targets\(^2\) whereas the American Association of Neurological Surgeons’ guideline of 85 – 95 mmHg mean arterial pressure (MAP) has little supporting evidence\(^3\). To aid patient management, we developed techniques for monitoring intraspinal pressure (ISP), spinal cord perfusion pressure (SCPP), spinal pressure reactivity (sPRx)\(^4,5\) and metabolism\(^6\) from the injury site. These techniques are safe based on a recent review of 42 patients\(^7\). In general, the optimum spinal cord perfusion pressure (SCPP\(_{opt}\)), defined as the perfusion pressure that minimizes sPRx\(^5\), is around 90 mmHg. In our original analysis, SCPP\(_{opt}\) was computed by pooling data from 18 patients\(^4,5\). Our initial concept of a pooled SCPP\(_{opt}\) requires revision for several reasons: First, there is now evidence that SCPP\(_{opt}\) varies between patients\(^8\) and that the injury site metabolic response is individualistic\(^9\) thus targeting 90 mmHg may not benefit everyone. Second, the SCPP\(_{opt}\) of each patient may change with time. Third, the SCPP\(_{opt}\) can only be computed after the monitoring has been completed and is, therefore, not clinically useful. Here, we show how to compute a continuous SCPP\(_{opt}\) (cSCPP\(_{opt}\)) in real-time and provide evidence that achieving this cSCPP\(_{opt}\) may be beneficial after TSCI.
MATERIALS AND METHODS

Patient recruitment. Approval for the Injured Spinal Cord Pressure Evaluation (ISCoPE) study, including the patient information sheet and consent form, was obtained from the St George’s Joint Research Office and the National Research Ethics Service London – St Giles Committee (No. 10/H0807/23). The study is registered with https://clinicaltrials.gov as NCT02721615. Inclusion criteria are: 1. Severe TSCI defined as American Spinal Injuries Association (ASIA) grade A, B or C; 2. Age 18 – 70 years; 3. Timing between TSCI and surgery ≤72 hours. Exclusion criteria are: 1. Patient unable to consent; 2. Other major injuries or co-morbidities; 3 Penetrating TSCI. Surgery and early management took place at the neurosurgery and neurointensive care units of St. George’s Hospital. Recruitment into the study was discussed with all eligible patients and their families on admission and a patient information sheet was given. Each patient signed a consent form.

Probe placement. Following bony realignment and posterior fixation, a pressure probe (Codman Microsensor Transducer®, Depuy Synthes, Leeds, UK) and a microdialysis (MD) catheter (CMA microdialysis AB, Sweden) were tunnelled through skin into the wound cavity. Under the operating microscope, the dura was opened one level below the injury. The pressure probe and MD catheter were inserted through the durotomy and placed on the spinal cord surface. We previously showed that the surface MD catheter samples from the injured spinal cord, not the cerebrospinal fluid (CSF) 6. The dural opening was sutured and supplemented with fibrin glue (Tisseel®, Baxter, UK). The ISP probe and MD catheter were secured to the skin with silk sutures. A wound drain was set at gravity drainage for a week. Patients were given 5,000 units
dalteparin daily starting on the day after the surgery. Dalteparin was omitted 24 hours before removing the probes and wound drain.

**Patient assessments.** Neurological examinations were done on admission and at two weeks postoperatively, i.e. before discharge to the rehabilitation facility. A CT and MRI of the whole spine were done on admission. Another CT was done within 48 hours of surgery to check screw and probe positions. MRI of the injured spinal cord was done at about two weeks and at 6–12 months after surgery.

**Intraspinal pressure and spinal cord perfusion pressure.** The pressure probe was connected to a Codman Intracranial Pressure box linked via a ML221 amplifier to a PowerLab running LabChart v.7.3.5 (AD Instruments, Oxford, UK). Blood pressure was recorded from a radial artery catheter, kept at the same level as the ISP probe, connected to the Philips Intellivue MX800 bedside monitoring system (Philips, Guildford, UK) in turn connected to the PowerLab system. The ISP and arterial blood pressure signals were sampled at 1 kHz for up to a week. LabChart was used to compute SCPP, defined as mean arterial pressure (MAP) minus ISP.

**Spinal pressure reactivity.** The spinal pressure reactivity index sPRx, a measure of spinal cord vascular reactivity, is the running correlation coefficient between ISP and MAP calculated over a five-minute period as described \(^4,^5\). If the spinal cord vasculature reacts normally, sPRx ≤ 0. If autoregulation is impaired, sPRx > 0.
**Microdialysis.** MD monitoring was started postoperatively in the neurointensive care unit. Central nervous system perfusion fluid (CMA Microdialysis AB, Sweden) was perfused at a rate of 0.3 \( \mu \text{L/min} \) using the CMA106 infusion pump (CMA Microdialysis AB, Sweden). MD vials were changed hourly, stored at 4 °C, and batched analysed up to 24 hours later using ISCUS Flex (CMA Microdialysis AB, Sweden). The first two samples from each patient were discarded to allow priming of the MD catheter and stabilization of the metabolite concentrations. All samples were analyzed for glucose, lactate, pyruvate, glycerol and glutamate. 100-fold changes in metabolite concentration, compared with the preceding hour, were excluded from analysis. For details see our earlier papers 6, 10.

**Optimum spinal cord perfusion pressure.** We used the software ICM+ (www.neurosurg.cam.ac.uk/icmplus). SCPP was divided into 5 mmHg bins between 40 – 120 mmHg and the mean sPRx for each bin was computed. To obtain the pooled SCPP\(_{\text{opt}}\) (pSCPP\(_{\text{opt}}\)) we used the entire signal of each patient. We averaged the sPRx for all patients in each SCPP bin and plotted these values *versus* SCPP. pSCPP\(_{\text{opt}}\) is the minimum of this curve. The same method was used to plot sPRx (averaged for all patients) *versus* ISP. To obtain the individual global spinal cord perfusion pressure (igSCPP\(_{\text{opt}}\)), we used the entire signal of each patient and plotted sPRx *versus* SCPP curve for each patient (rather than averaging over all patients). The igSCPP\(_{\text{opt}}\) is the minimum of each curve. The same method was used to obtain the individual continuous optimum spinal cord perfusion pressure (cSCPP\(_{\text{opt}}\)), but by using a four-hour window (unless otherwise stated) rather than the entire signal, moving forward each minute.
**Enhanced visualization.** sPRx *versus* SCPP curves were produced each minute, using a sliding four-hour window. The horizontal axis is time and the vertical axis is SCPP. At each time point, sPRx values are color-coded green (intact reactivity), yellow (borderline reactivity) or red (impaired reactivity). Time points, for which no cSCPP\textsubscript{opt} could be determined, are represented by vertical bands filled with monotonic green. A non-causal, exponentially weighted moving average filter was applied to the image along the time axis. This filter has a smoothing effect in time and fills in some of the gaps with appropriately weighted average of the preceding and following data values. The measured cSCPP\textsubscript{opt} values were smoothed with the same filter and added to the image. Details are given elsewhere \textsuperscript{11}. The images shown here were generated in ICM+ with the current implementation of the MATLAB tool \textsuperscript{12}.

**Statistical analysis.** Two-tailed t-test was used to compare metabolite level within 5 mmHg *versus* more than 5 mmHg from the cSCPP\textsubscript{opt} with significance at \( P < 0.01^{**} \) and \( 0.005^{#} \). We used the Spearman coefficient to quantify the correlation between neurological outcome *versus* deviation from cSCPP\textsubscript{opt} or MAP guideline.
RESULTS

Demographics. Table 1 summarizes details of the 45 patients. Most are young, with 84% <60 years old. Males outnumber females by 3.5:1. Cervical TSCIs are more common than thoracic at 56 vs. 44%. Most had neurologically complete TSCI on admission; two thirds were ASIA grade A. Overall, 13 % (6/45) patients improved by one grade and 16 % (7/45) by at least two grades at 9 – 12 months. Of those who were grade A 17 % improved versus 44 % of grade B and 67 % of grade C.

Pooled SCPP\textsubscript{opt}. Using data from all 45 patients, we plotted sPRx \textit{versus} ISP and sPRx \textit{versus} SCPP. Figure 1A shows that as ISP increases, sPRx increases. This suggests that higher ISP correlates with more marked loss of injury site autoregulation. Figure 1B shows a U-shaped relationship between sPRx and SCPP, which suggests that the optimum SCPP is around 80 – 90 mmHg. As SCPP falls below 80 mmHg or rises above 90 mmHg, injury site autoregulation progressively worsens. Since the optimum SCPP of 80 – 90 mmHg was computed by pooling data from all patients, we term this pooled SCPP\textsubscript{opt} (pSCPP\textsubscript{opt}). A major drawback of pSCPP\textsubscript{opt} is that it does not provide information about the variability of SCPP\textsubscript{opt} between patients.

Individual global SCPP\textsubscript{opt}. To test the hypothesis that SCPP\textsubscript{opt} varies between patients, we computed SCPP\textsubscript{opt} for individual patients by plotting the sPRx \textit{versus} SCPP for each individual patient. We termed this the individual global (igSCPP\textsubscript{opt}) because it only applies to an individual patient and the entire monitoring period is used. Figure 1C shows sPRx \textit{versus} SCPP curve for three patients; two had a U-shaped relationship with approximately 20 mmHg difference in their igSCPP\textsubscript{opt}, whereas in one there was no correlation between sPRx and SCPP. igSCPP\textsubscript{opt} values
ranged from 50 – 60 mmHg in some patients up to 100 – 110 mmHg in others (Figure 1D). In 13 % (6/45) patients, igSCPP\textsubscript{opt} could not be computed because the relation between sPRx \textit{versus} SCPP is not U-shaped. Unlike pSCPP\textsubscript{opt}, which is averaged over many patients, igSCPP\textsubscript{opt} provides individual patient information. Because igSCPP\textsubscript{opt} requires the entire ISP and MAP signals, igSCPP\textsubscript{opt} does not indicate how a patient’s SCPP\textsubscript{opt} changes with time and can only be computed after the monitoring has ended. Thus, igSCPP\textsubscript{opt} cannot be used to guide patient management in real time in the neurointensive care unit.

\textbf{Individual continuous SCPP\textsubscript{opt}.} To overcome the problems with igSCPP\textsubscript{opt}, we computed a running SCPP\textsubscript{opt} for each patient, which we term the individual continuous SCPP\textsubscript{opt}, abbreviated as cSCPP\textsubscript{opt}, by analogy to the continuous optimum cerebral perfusion pressure (cCPP\textsubscript{opt}) for brain injury\textsuperscript{11,13}. cSCPP\textsubscript{opt} was calculated using a four-hour window, updated every minute. Fig. 2 shows sPRx \textit{versus} SCPP plots, obtained using four-hour windows. Some plots (plot a) were U-shaped thus allowing cSCPP\textsubscript{opt} to be computed. In other cases, cSCPP\textsubscript{opt} could not be calculated because the injured cord was underperfused (descending curve, plot b) or overperfused (ascending curve, plot c) or there was no correlation between sPRx and SCPP (plot d). We then varied the time window from 2 – 10 hours to compute the cSCPP\textsubscript{opt}. For each time window the SCPP\textsubscript{opt} is calculated, the window is moved forward 60 seconds and the SCPP\textsubscript{opt} result is updated. The process continues for the entire SCPP signal. Therefore, when process is finished, there is a SCPP\textsubscript{opt} vector every 60 seconds. Each vector has four components (a, b, c, d) and can be (1, 0, 0, 0), (0, 1, 0, 0), (0, 0, 1, 0) or (0, 0, 0, 1) as follows: a - optimum value found, b - hypoperfused, c - hyperperfused, d - no relation between sPRx and SCPP. We then computed the frequencies with which a, b, c, d occur in the entire SCPP signal. The process
was repeated for each patient. Fig. 2B shows that time windows greater or equal to four hours maximize the chance of obtaining the cSCPP\textsubscript{opt}. With a four-hour window, cSCPP\textsubscript{opt} could be computed 45% of the time. Fig. 3 shows many signals (ISP, MAP, SCPP, sPRx, and cSCPP\textsubscript{opt}) simultaneously displayed for a single patient. SCPP – cSCPP\textsubscript{opt} is also displayed to allow doctors to see in real-time how their treatment deviates from the optimum.

**Histograms of continuous SCPP\textsubscript{opt}**. We plotted each patient’s cSCPP\textsubscript{opt} histogram (Fig. 4A). Distributions varied widely; some were unimodal-symmetrical with a predominant cSCPP\textsubscript{opt} peak (patient i) or a wide range of cSCPP\textsubscript{opt} values (patient ii) or bi-modal (patient iii). Left- and right-skewed unimodal distributions were also found (not shown). Fig. 4B shows the standard deviations of these distributions; in most patients, the standard deviation was high at 10 – 15 mmHg and in some as high as 15 – 20 mmHg. The high variability in cSCPP\textsubscript{opt} suggests that a patient’s SCPP\textsubscript{opt} is dynamic and, therefore, targeting the same SCPP is inadequate. Fig. 4C shows significant positive correlation between igSCPP\textsubscript{opt} and mean cSCPP\textsubscript{opt}.

**Neurological outcome**. We hypothesized that larger deviation of SCPP from cSCPP\textsubscript{opt} correlates with worse neurological outcome at 9 – 12 months. The data show that as the difference between SCPP and the cSCPP\textsubscript{opt} (averaged over each patient’s entire monitoring period) decreases, the chance of ASIA grade improvement increases (Fig. 5A) and the chance of having a complete neurological deficit at follow-up (ASIA A) decreases (Fig. 5B). There are only two patients with positive mean deviation of SCPP from cSCPP\textsubscript{opt}; thus, we do not know if hyperperfusion correlates with worse neurological outcome. There was no correlation between the American Association of Neurological Surgeons’ guideline of 85 – 95 mmHg MAP and the chance of
ASIA grade improvement (Fig. 5C) or the chance of having complete deficit (Fig. 5D) at 9 – 12 months.

**Enhanced visualization.** Displaying cSCPP\textsubscript{opt} as a line (Fig. 3) is clinically helpful, but has drawbacks. First, it fails to show the range of cSCPP\textsubscript{opt} at any one time that are associated with intact autoregulation, i.e. sPRx $\leq 0$; a cSCPP\textsubscript{opt} line simplistically implies that cSCPP\textsubscript{opt} at any one time is a single value rather than a range. Second, the cSCPP\textsubscript{opt} line has gaps whenever the optimum cannot be calculated. We, therefore, used an enhanced visualization technique, recently developed to overcome these issues when monitoring in severe traumatic brain injury\textsuperscript{11,12}. Fig. 6 shows two monitoring periods from the same TSCI patient. The top trace shows a wide range (60 – 90 mmHg) of cSCPP\textsubscript{opt}. Initially, the actual SCPP was within this range, but between 00:30 and 01:45 hours the injured cord was hypoperfused. After two days (bottom trace) the range of cSCPP\textsubscript{opt} narrowed substantially. Between 13:00 and 14:00 hours, the cSCPP\textsubscript{opt} range was narrow at around 78 mmHg, between 14:00 and 16:00 hours it was 65 – 85 mmHg and, after 18:30, it narrowed again at around 85 mmHg. The enhanced visualization technique not only displays the range of cSCPP\textsubscript{opt} values at any one time, but also fills some of the gaps in the signal.

**Injury site metabolism.** After TSCI, injury site metabolism can be monitored using surface MD\textsuperscript{6}. We hypothesized that small deviation of SCPP from cSCPP\textsubscript{opt} correlates with less injury site metabolic derangement. To test this, we quantified the hourly injury site metabolic state in 14 spinal cord injury patients. Figure 7 shows that hourly SCPP close to cSCPP\textsubscript{opt} is associated with low injury site glucose, high lactate and high pyruvate. As SCPP deviates from cSCPP\textsubscript{opt} by
more than 5 mmHg, injury site glucose rises whereas lactate and pyruvate fall. Therefore, cSCPP$_{opt}$ does not minimize injury site metabolic derangement, but appears to maximize glucose utilization at the injury site. There was no correlation between deviation from cSCPP$_{opt}$ and glutamate, glycerol or lactate-to-pyruvate ratio (not shown). cSCPP$_{opt}$ was not associated with a unique metabolic profile (Supplement).
DISCUSSION

In this study, we introduced the novel concept of $\text{cSCPP}_{opt}$, which is a range rather than an individual number. To guide the management of TSCI patients, we showed how $\text{cSCPP}_{opt}$ could be displayed in real-time using an enhanced visualization method.

Our data show that $\text{cSCPP}_{opt}$ varies widely between and within patients. Various reasons may explain such wide inter-patient variability: patients with more extensive microvascular damage, pre-existing hypertension or greater leukocyte infiltration at the injury site will likely require higher SCPP. The large intra-patient variability in $\text{cSCPP}_{opt}$ may be due to the dynamic nature of the pathological processes at the injury site e.g. changes in the numbers of infiltrating leukocytes \(^{14}\) and reactive glia \(^{15}\) as well as changes in injury site edema \(^{16-18}\) and fever \(^{19}\). Most $\text{cSCPP}_{opt}$ distributions are unimodal, but some are bimodal. A bimodal distribution may arise by sudden events that reset the $\text{cSCPP}_{opt}$, e.g. spinal cord hemorrhage. Further studies are required to clarify the significance of the different $\text{cSCPP}_{opt}$ distributions. Based on our data, we suggest that targeting the same SCPP continuously in all patients is inadequate.

$c\text{SCPP}_{opt}$ could only be computed 45% of the time. It would be clinically useful for $\text{cSCPP}_{opt}$ to be computable more often. One way to achieve this may be by using a Dynamic Adaptive Target of Cerebral Autoregulation algorithm, to combine different sPRx values and time windows in a weighted manner to issue a $\text{cSCPP}_{opt}$ recommendation. When applied to cerebral perfusion pressure obtained from TBI patients, such an algorithm produced a $\text{CPP}_{opt}$ recommendation 97% of the entire monitoring period \(^{20,21}\).

Is actively targeting $\text{cSCPP}_{opt}$ safe? Our audit of 42 patients concluded that insertion of the pressure probe and ISP monitoring for up to a week are safe \(^{7}\). However, actively increasing SCPP requires vasopressors, which may produce complications such as cardiac dysrhythmias,
especially in older patients. We previously showed that, after TSCI, the spinal cord swells and is compressed by the dura. Thus, one way to reduce vasopressor requirements is expansion duroplasty, a simple and safe surgical procedure that lowers ISP and sPRx as well as increasing SCPP. Based on our experience thus far, SCPP monitoring is safe and helpful to guide management of TSCI patients.

Is targeting cSCPP$_{opt}$ beneficial? Our MD data suggest that cSCPP$_{opt}$ maximizes glucose utilization by the injured cells. The Kohonen self-organizing maps show that each patient’s injury site metabolic response is unique. Therefore, each patient’s optimal injury site metabolic profile must also be unique, which explains why cSCPP$_{opt}$ is not associated with a single metabolic signature (Supplement). How about neurological outcome? We found that larger deviation from cSCPP$_{opt}$ (hypoperfusion) is associated with worse neurological outcome at 9 – 12 months, though the effect of hyperperfusion on neurological outcome could not be determined. Aries et al. showed that increased deviation from cCPP$_{opt}$ is associated with worse neurological outcome after brain injury. These observations suggest that targeting the continuous optimum perfusion pressure may be beneficial after central nervous system injury.

Are there ways of managing spinal cord perfusion without targeting cSCPP$_{opt}$? One may follow the 85 – 95 mmHg MAP guideline or drain CSF with a lumbar catheter. MAP-oriented management may be inadequate, evident by the lack of a strong correlation between achieving the guideline and neurological outcome as shown in Fig. 6 and by others. A major drawback of the MAP guideline is that it does not consider ISP; thus, the same MAP may hypoperfuse the injury site in one patient and hyperperfuse it in another. Also, the MAP guideline does not consider inter- and intra- patient variability in SCPP$_{opt}$. Draining lumbar CSF is also likely inadequate because, at the injury site, the cord is not surrounded by CSF, but is
compressed against dura\textsuperscript{5,8,24,29,30}. Thus, CSF drainage is unlikely to reduce injury site ISP. Ultimately, a randomized controlled trial is required to definitively determine whether interventions to achieve cSCPP\textsubscript{opt} improve neurological outcome after TSCI.
ACKNOWLEDGEMENTS

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DISCLOSURES

PS and MC receive royalties for ICM+ from the software licensing fee.
REFERENCES


## TABLES

Table 1. Patients’ demographic characteristics.

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FIGURE LEGENDS

Fig. 1. Concepts of pSCPP\textsubscript{opt} and igSCPP\textsubscript{opt}. A. sPRx \textit{versus} ISP and B. sPRx \textit{vs}. SCPP using pooled data from 45 patients. Dotted line shows pSCPP\textsubscript{opt}. C. sPRx \textit{vs}. SCPP for three patients (a, b, c). Dotted line shows igSCPP\textsubscript{opt}. D. igSCPP\textsubscript{opt} values for 45 patients. Nil means igSCPP\textsubscript{opt} not computable. Mean ± standard error.

Fig. 2. Concept of cSCPP\textsubscript{opt}. A. Four sets (a, b, c, d) of corresponding ISP, MAP, SCPP and sPRx signals. cSCPP\textsubscript{opt} can be obtained in ‘a’, but not in ‘b’ (hypoperfused), ‘c’ (hyperperfused) or ‘d’ (no relation between sPRx and SCPP). B. % time \textit{versus} ‘a’, ‘b’, ‘c’ and ‘d’ for time windows between 2 – 10 hours. Mean ± standard error.

Fig. 3. Simultaneous display of multiple signals. Simultaneously displayed ISP, MAP, SCPP and sPRx, cSCPP\textsubscript{opt} and SCPP – cSCPP\textsubscript{opt} signals from one patient.

Fig. 4. Distribution of cSCPP\textsubscript{opt} values. A. cSCPP\textsubscript{opt} distributions in three patients: i – unimodal leptokurtic, ii – unimodal pachykur tic, iii – bimodal. B. Standard deviations of the 45 cSCPP\textsubscript{opt} distributions. C. igSCPP\textsubscript{opt} \textit{versus} mean cSCPP\textsubscript{opt} for 45 patients and best-fit straight line (R\textsuperscript{2} = 0.56).

Fig. 5. Deviation of SCPP from cSCPP\textsubscript{opt} \textit{versus} neurological outcome. A. % patients who improved by at least one ASIA grade \textit{versus} mean SCPP – cSCPP\textsubscript{opt} (R\textsuperscript{2} = 0.95). B. % patients with complete (ASIA A) neurological deficit at follow-up \textit{versus} mean SCPP – cSCPP\textsubscript{opt} (R\textsuperscript{2} = 1.00). C. % patients who improved by at least one ASIA grade \textit{vs}. mean deviation from the 85 –
95 mmHg MAP Guideline \( (R^2 = 0.38) \). \textbf{D.} % patients with complete (ASIA A) neurological deficit at follow-up \textit{versus} mean deviation from MAP Guideline \( (R^2 = 0.67) \).

\textbf{Fig. 6. Enhanced visualization of cSCPP\textsubscript{opt}.} Plots of SCPP \textit{versus} time for two periods (4 Jan, 6 Jan). sPRx zones: Red (loss of autoregulation), Green (preserved autoregulation) and Yellow (transition from preserved to impaired autoregulation) as per color scale. Line shows actual SCPP.

\textbf{Fig. 7. Deviation of SCPP from cSCPP\textsubscript{opt} \textit{versus} injury site metabolism.} Glucose, lactate and pyruvate concentration \textit{versus} mean hourly SCPP – cSCPP\textsubscript{opt}. Mean ± standard error. \( P < 0.01^{**}, < 0.005^\# \) for metabolite level at \( 5 – 5 \) mmHg compared with all others together.
FIGURE 2
**Figure 5**

(A) % patients who improved

(B) % AIS grade A patients at follow-up

(C) % patients who improved

(D) % AIS grade A patients at follow-up

Deviation from cSCPP_{opt} (mmHg)

Deviation from MAP guide (mmHg)
SUPPLEMENTARY DATA

**Kohonen self-organizing maps.** We define a vector as a set of five metabolite measurements at a time point. We first removed outliers and interpolated vectors with missing components as we described \(^1\). After filtering and interpolation, the MD dataset comprised 1,611 five-dimensional vectors. The five metabolites were then normalized to a mean of zero and unit standard deviation. Subsequent analysis was done using the self-organizing map (SOM) toolbox for Matlab v.2.0beta at [www.cis.hut.fi/somtoolbox/](http://www.cis.hut.fi/somtoolbox/) \(^2\). The grid length and width were computed from the ratio between the eigenvalues of the training data. The grid was initialized using random weight vectors as we described \(^1\). The SOM algorithm uses competitive learning to project the five-dimensional vectors onto a two-dimensional map where similar vectors are cluster close to each other. The algorithm preserves topological relationships between the five-dimensional vectors as they are mapped onto the two-dimensional grid thus facilitating visualization of clinically important, non-linear relationships between the vectors. We colored the Kohonen grid by metabolite level, patient number and mean hourly deviation from the cSCPP\(_{\text{opt}}\). For details see our earlier paper \(^1\).

We determined whether cSCPP\(_{\text{opt}}\) is associated with a specific injury site metabolic signature. Supplementary Fig. 1A shows Kohonen metabolite SOMs. The upper part of the SOM indicates high glucose, low lactate, pyruvate, glutamate and glycerol and thus represents near-normal metabolism. The bottom left of the SOM indicates low glucose, high lactate and high pyruvate and thus represents hypermetabolism. The bottom right part of the SOM indicates low glucose, high lactate, medium pyruvate as well as high glutamate and glycerol and thus represents ischemia. Patients occupy different regions of the metabolic map, which suggests that they have individualistic injury site metabolic responses to TSCI (Supplementary Fig. 1B, left). Supplementary figure 1B (right) shows that the yellow hexagons are scattered throughout the map, i.e. having SCPP within 5 mmHg of cSCPP\(_{\text{opt}}\) is not associated with a unique metabolic signature. This finding suggests that the optimum injury site metabolic response differs between patients.
Suppl. Fig. 1. Deviation of SCPP from $c\text{SCPP}_{\text{opt}}$ versus injury site metabolism. A. Kohonen SOMs of injury site glucose, lactate, pyruvate, glutamate and glycerol. Heat maps with the average value in each hexagon colour coded. B. (left) Kohonen SOM coloured by patient (P1 – P14). Each hexagon is coloured according to the patient with the most vectors in that hexagon. (right) Kohonen SOM colored according to deviation from $c\text{SCPP}_{\text{opt}}$. Each hexagon is color coded according to its mean SCPP – $c\text{SCPP}_{\text{opt}}$ value.

REFERENCES