Hypothermic treatment after computer-controlled compression in minipig: A preliminary report on the effect of epidural vs. direct spinal cord cooling

MONIKA ZAVODSKA1, JAN GALIK1, MARTIN MARSALA1,2, STEFANIA PAPCUNOVA1, JAROSLAV PAVE1, ENIKO RACEKOVA1, MARCELA MARTONCIKOVÁ1, IGOR SULLA1,3, MIROSLAV GAJDOS1, IMRICH LUKAC1, JOZEF KAFKA1, VALENT LEDECKY3, IGOR SULLA JR.3, PETER REICHEL3, ALEXANDRA TRBOLOVA3, IGOR CAPIK3, KATARINA BIMBOVA1, MARIA BACOVA1, ANDREA STROPKOVSKA1, ALEXANDRA KISUCKA1, DANA MIKLISOVA6 and NADEZDA LUKACOVA1

1Institute of Neurobiology, Biomedical Research Center, Slovak Academy of Sciences, 040 01 Košice, Slovakia; 2Department of Anesthesiology, Neuroregeneration Laboratory, University of California-San Diego, San Diego, CA 92093, USA; 3Hospital of Slovak Railways, 040 01 Košice; 4Department of Neurosurgery, Faculty of Medicine, University of Pavol Jozef Safarik, 040 66 Košice; 5Department of Small Animal Clinic, University of Veterinary Medicine and Pharmacy, 041 81 Košice; 6Department of Vector-borne Diseases, Institute of Parasitology, Slovak Academy of Sciences, 040 01 Košice, Slovakia

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Abstract. The aim of the present study was to investigate the therapeutic efficacy of local hypothermia (beginning 30 min post-injury persisting for 5 h) on tissue preservation along the rostro-caudal axis of the spinal cord (3 cm cranially and caudally from the lesion site), and the prevention of injury-induced functional loss in a newly developed computer-controlled compression model in minipig (force of impact 18N at L3 level), which mimics severe spinal cord injury (SCI). Minipigs underwent SCI with two post-injury modifications (durotomy vs. intact dura mater) followed by hypothermia through a perfusion chamber with cold (epidural t≈15˚C) saline, DMEM/F12 or enriched DMEM/F12 (SCI/durotomy group) and with room temperature (t≈24˚C) saline (SCI-only group). Minipigs treated with post-SCI durotomy demonstrated slower development of spontaneous neurological improvement at the early postinjury time points, although the outcome at 9 weeks of survival did not differ significantly between the two SCI groups. Hypothermia with saline (t≈15˚C) applied after SCI-durotomy improved white matter integrity in the dorsal and lateral columns in almost all rostro-caudal segments, whereas treatment with medium/enriched medium affected white matter integrity only in the rostral segments. Furthermore, regeneration of neurofilaments in the spinal cord after SCI-durotomy and hypothermic treatments indicated an important role of local saline hypothermia in the functional outcome. Although saline hypothermia (24˚C) in the SCI-only group exhibited a profound histological outcome (regarding the gray and white matter integrity and the number of motoneurons) and neurofilament protection in general, none of the tested treatments resulted in significant improvement of neurological status. The findings suggest that clinically-proven medical treatments for SCI combined with early 5 h-long saline hypothermia treatment without opening the dural sac could be more beneficial for tissue preservation and neurological outcome compared with hypothermia applied after durotomy.

Introduction

Spinal cord injury (SCI) is associated with severe and long-lasting neurological dysfunction in humans. Currently, there is no effective treatment to reverse the consequences resulting from traumatic SCI. While in experimental studies, various pharmacological agents and therapeutic approaches have tended to improve outcomes in animal SCI models (1), unfortunately none of these therapies has been successfully transferred into clinical practice.

After SCI the development of the lesion consists of primary and secondary phases (2,3). The primary phase is the direct mechanical injury to the spinal cord tissues itself, which is followed by a secondary phase leading to a marked enlargement of the primary lesion within several h after SCI. This
slow evolution of the lesion has been referred to as progressive hemorrhagic necrosis or ‘autodestruction’ (4,5). It results in secondary loss of vital spinal cord tissue and, in some species including humans, leads to formation of a cystic cavity surrounded by glial scar tissue.

From the clinical point of view, the secondary phase is very important because it represents the target for therapeutic interventions which could minimize the extent of the final lesion (6). To this date the treatments options for SCI are limited to surgical interventions, early steroid administration and physical therapy. These therapeutic interventions are effective only under specific circumstances (7,8), and their effect on the final outcome is modest at best.

One potential treatment which has shown promising results over the past few decades is therapeutic hypothermia. Although the entire range of the effects of hypothermia is not fully understood, a few observations are notable (9-12). Hypothermia seems to work through several mechanisms which start to operate soon after trauma and may have an important role in the functional outcome. Neuroprotective effects of systemic and regional hypothermia have been described (9-11,13,14). It has been demonstrated that hypothermia has a powerful effect on control of the injury-induced immune response (15), apoptosis and inflammation (13,14,16-22). Hypothermia also delays glutathione release, reduces oxidative stress and increases spinal cord tolerance to ischemia (23-26). There are also numerous articles illustrating the neuroprotective effect of hypothermia on neural stem cell differentiation and survival (15,27-35).

The published studies have defined the most effective target temperature range for systemic hypothermia (36-39) and they suggest that the most effective cooling is such that i) is initiated early after injury, ii) lasts for long periods (up to 48-h) and iii) is followed by slow rewarming (13,38).

Relatively little experimental data are available for local hypothermia. In general, the results indicate that longer durations of local hypothermia (at least 48 h) yield better results than shorter durations (14,40-43). However, there are studies indicating that duration of hypothermia <48 h may be sufficient. In the literature, some animal studies found that local hypothermia improved outcomes after SCI (37,38,42,44-50), whereas others concluded that it has no effect (51-54). Interestingly, in a clinical study by Hansbout and Hansebout (55), positive results were observed with local hypothermia lasting <4 h. In this study, the cooling was initiated 7.1 h after injury, the mean duration of the treatment was 3.7 h and there was an uncontrolled rewarming rate.

In the present study, we wanted to test the protective effect of local hypothermia after precisely-controlled SCI model in minipigs (56,57) representing a preclinical model which mimics human patients more closely than the widely-used rodent models. In a series of experiments, we modified i) the cooling approach (durotomy vs. intact dura mater), ii) the kind of cooling solutions (saline, medium or enriched medium), and iii) the degree of hypothermia (15 or 24°C just above the SCI). The outcome parameters involved long-term behavioral testing for the recovery of neurological functions, and detailed histological analyses of the spinal cord lesions involving axonal quantification using a selective marker of mature axons (SMI-312).

Materials and methods

Animals. A total of 24 adult female Göttingen-Minnesota-Liběchov minipigs each weighing 25-35 kg were used in this study. The animals were maintained in standard conditions with ad libitum access to food and water. The experimental protocols were prepared in accordance with the EC Council Directive (2010/63/EU) regarding the use of animals in research and approved by the State Veterinary and Food Administration of the Slovak Republic (decision no. 1319/13-221) as well as by the Ethical Commission of the University of Veterinary Medicine and Pharmacy (Kosice, Slovak Republic). All efforts were made to minimize the size of experimental groups and animal suffering.

Experimental groups. Two sets of experiments with modifications in post-injury approach (durotomy vs. intact dura mater) and temperature of perfusion solutions (15 or 24°C) were carried out.

The animals in Experiment 1 were divided into the following groups: i) naïve control-without anesthesia or surgical intervention (n=3); ii) SCI at L3 level + durotomy (5 h) and survival for 9 weeks (n=3); iii) SCI at L3 level + durotomy followed by 4°C saline treatment (target temperature: 15°C for 5 h) and survival for 9 weeks (n=3); iv) SCI at L3 level + durotomy followed by 4°C medium treatment (target temperature, 15°C for 5 h) and survival for 9 weeks (n=3); v) SCI at L3 level + durotomy followed by enriched 4°C medium treatment (target temperature: 15°C for 5 h) and survival for 9 weeks (n=3).

In order to relieve the edema and provide direct contact of cooling solutions with the spinal cord, the minipigs underwent durotomy immediately after SCI. Medium (DMEM/F12) was composed of rh Transferrin 100 mg/l, rh Insulin 25 mg/l, Glucose 1.56 g/l, Progesterone 6.3 µg/l, Putrescine 16.1 mg/l, Selenium 5.2 µg/l. Enriched medium was composed of DMEM/F12 + rh bFGF 10 µg/l, rh BDNF 10 µg/l, rh GDNF 10 µg/l, rh VEGF 10 µg/l, Creatine 20 mM. The treatments were initiated 30 min after SCI and durotomy, and maintained for 5 h (see details of hypothermic treatment below).

The animals in Experiment 2 were divided into the following groups: i) naïve control-without anesthesia or surgical intervention (n=3); ii) SCI at L3 level (intact dura mater) and survival for 9 weeks (n=3); iii) SCI at L3 level (intact dura mater) followed by room temperature saline treatment (target temperature: 24°C for 5 h) and survival for 9 weeks (n=3). The treatment was initiated 30 min after SCI and maintained for 5 h (see details of hypothermic treatment below).

Pre-contusion procedures. Three days before the surgery, the animals were pre-treated with a combination of penicillin and streptomycin, administered intramuscularly in a dose of 0.5 ml/30 kg (Norbrook Laboratories, Newry, Northern Ireland). The minipigs were premedicated 30 min before surgical procedure with an intramuscular injection of Stresnil (2 mg/kg; Janssen Pharmaceutica, Beerse, Belgium) and Atropin (0.5 mg/kg; Biotika, Slovenská Ľupča, Slovakia). Anesthesia was induced by intravenous administration of Thiopental (10 mg/kg; Czech Pharma, Czech Republic)
followed by identification of the precise location of the L3 vertebra using plain X-rays in a lateral projection (58). Afterwards, the animals were endotracheally intubated (diameter, 5.0–6.0 mm) and anesthesia was maintained by inhalation of 1.5% Sevoflurane (Baxter, Prague, Czech Republic) mixed with oxygen. Analgesia was supported by administration of Butomidor (0.4 mg/kg, i.v.; Richter Pharma, Austria). Catheters for the administration of infusions and medications were inserted bilaterally into the cephalic and auricular veins. Standard monitoring of life-function indicators (mainly heart rate, respiratory rate, blood pressure and oxygen saturation) was performed during the whole surgical procedure and hypothermic treatment. Hydration was maintained with lactated Ringer’s solution.

Spinal cord contusion. The animals were placed into a special immobilization apparatus (Fig. 1A). After disinfection of the lumbar area, a midline skin and subcutaneous fat incision was performed from L2 to L4 segments. Muscles were dissected from the spinous processes and vertebral arches and retracted, then a dorsal laminectomy at the L3 segment was performed. Afterwards the minipigs received the muscle relaxant suxamethonium (ViDiTo, Kosice, Slovakia) and a digital force gauge connected to a 5-mm diameter circular aluminum bar (Fig. 1B). Parameters for SCI (velocity: 10 mm/sec; force: 18N) were preset by the computer software (56). To control the uniformity of the impact force, the contusion curve was recorded in each animal (Fig. 1C). The force impact was uniform without significant differences between animals or groups. The real peak force in Experiment 1 was 21.08 N ± 0.79 and in Experiment 2 it was 20.72 N ± 0.97. After SCI, the paraspinal musculature and skin were sutured with several absorbable stitches or durotomy and subsequent hypothermic treatments were performed depending on the experimental group. The animals were housed in thresomes to recover. Post-operative care included bladder expression twice a day, antibiotic administration Penstrepten (0.5 ml/30 kg/day; Biotika) for 10 days, cleaning the hind limbs and monitoring of skin irritation and development of decubitus ulcers.

Implantation of perfusion chamber. In the case of animals undergoing experimental treatment, the perfusion chamber was implanted above the exposed spinal dural sac containing the spinal cord immediately after performing the SCI (Fig. 1D and E). A cone-shaped chamber with both ends opened was made from ABS plastic using a uPrint SE 3D printer (Stratasys, Eden Prairie, MN, USA). It had ellipse-shaped openings, the lower opening measuring 20x14 mm and the upper one 40x30 mm; the height of the chamber was 50 mm. The chamber was fitted with two tubings (inflow and outflow). The smaller opening of the chamber was about the size of the laminae, so it could be freely placed directly on the dura mater without any fixation to the spine or paravertebral muscles. Two superficially-placed stitches were used to partially close the wound, making the skin edges slightly embracing the chamber.

Hypothermia. Hypothermia with cold solutions was carried out locally with the help of the perfusion chamber. The inflow tubing was connected to the tank with sterile cold perfusion solution. The cooling of the injury site was initiated 30 min after SCI or SCI + durotomy. At the beginning, the solution drip rate was manually adapted to ensure that the temperature of fluid in the perfusion chamber (just above the SCI) was maintained at 15°C in Experiment 1 or at 24°C in Experiment 2. Given that cooling was applied locally, the temperature of the solution in the chamber decreased to 15°C or 24°C very rapidly, max. within 5-6 min and was measured at 30 min intervals using a needle-tip thermometer (Omega Bio-Tek, Inc., Norcross, GA, USA) during the whole treatment procedure. Hypothermic treatment of the injured spinal cord lasted for 5 h. Continuous inflow of the perfusion solution into the chamber was regulated by a peristaltic pump (bubble rise distance, flow 2 ml/min; Heidolph, Schwabach, Germany). The outflow tubing was used to drain excess perfusion solution. The average temperature of cooling solutions in the chamber (just above the site of injury) was 15.43°C ± 3.75 in Experiment 1 and 24.06°C ± 1.95 in Experiment 2. There were no significant differences in epidural temperature between groups during the whole treatment period. After the therapy, the perfusion chamber was extracted, the dura mater in Experiment 1 was closed with absorbable sutures, and the wound was sutured in layers. These animals were housed in separated cages to recover.

To minimize variability in body temperature and to prevent systemic hypothermia during treatment the minipigs were covered with an isothermal foil. The rectal temperature of each animal was measured at 30 min intervals. When the body temperature dropped below 36°C, the animals were heated with a stream of warm air blown below the blanket covering the animal. The average rectal temperature was 36.47°C ± 0.70.
Spinal cord tissue preparation. At the end of the experiment, the animals were anesthetized by intravenous administration of Thiopental (10 mg/kg; Czech Pharma) and transcardially perfused with heparinized saline (5l) followed by 4% paraformaldehyde in 0.1 M phosphate-buffered saline (PBS; pH 7.4, 5l). The lumbar spinal cords encompassing the injury site were carefully dissected out, cleaned of the dura mater remnants and post-fixed in 4% paraformaldehyde overnight. Next day, the spinal cords were cut into seven 1 cm segments: cranial (+3, +2, +1), central (0) and caudal (-1, -2, -3) segments. The spinal cord segments were cryoprotected in a solution of 30% sucrose in PBS at 4˚C for 2 days, cut into transverse serial sections (30 µm) on a cryostat (Leica Microsystems, Wetzlar, Germany) and used for histological and immunohistochemical analyses.

Histology. The selected sections were placed in PBS for 10 min, in 70% alcohol for 2 h and overnight in 0.1% Luxol fast blue (LFB) at room temperature. Next day, the sections were rinsed in distilled water and PBS for 3 min, put into 0.05% aqueous lithium carbonate followed by 40% alcohol, PBS and distilled water. Afterwards, the slices were counterstained with 0.2% cresyl violet for 4 min and rinsed in distilled water. Finally, the sections were dehydrated through a graded series of ethanol and xylene and covered with coverslips. Sections were visualized using an Olympus BX51/BX52 light microscope (magnification x5, fitted with an Olympus DP50 digital camera coupled with a computer equipped with Olympus DP Image software, version 3.1; Olympus Corporation, Tokyo, Japan). Slices from each segment were evaluated. The areas of spared white and grey matter were measured using ImageJ 1.44 (National Institutes of Health, Bethesda, MD, USA), expressed as a percentage and compared to control (control values were expressed as 100%). The cresyl violet counterstain enabled motor neurons to be counted. Motor neurons located in the ventral horn (VH) with clearly visible nuclei on both sides of the spinal cord were counted.

Immunohistochemistry. Selected slides taken from each rostral and caudal segment (except for the epicenter of the injury) were pre-treated in PBS for 3x10 min and incubated in blocking solution consisting of 5% normal goat serum in 0.1 M PBS with 0.3% Triton-X 100 (normal goat serum; Vector Laboratories, Burlingame, CA, USA) for 2 h at room temperature. Afterwards, the slices were incubated overnight at 4˚C with mouse monoclonal anti-neurofilament primary antibody (Pan-Axonal Neurofilament Marker SMI-312; dilution, 1:1,000; Abcam, Cambridge, UK) for 24 h at 4˚C. In the next step, the sections were rinsed 3 times in PBS,
followed by incubation in secondary anti-mouse antibody (RhodamineRed; dilution, 1:200; Invitrogen; Thermo Fisher Scientific, Inc., Waltham, MA, USA) for 90 min at room temperature. Sections were rinsed in PBS 3×10 min, mounted with Fluoromount (Serva, Heideberg, Germany) and covered with coverslips. SMI-312 positive axons were counted by an observer in a blinded procedure using a x20 lens over an area of 800×600 µm in three specific regions of the white matter: dorsal, lateral and ventral columns (DC, LC, VC) (Fig. 2), using Image J graphics software (ImageJ 1.44; National Institutes of Health). The images were converted into a binary image using a set threshold. NF positive axons were counted automatically using the ‘analyze particles’ tool (57, 59). The number of neurofilaments (expressed as percentage of NF/mm²) was evaluated bilaterally from ten randomly-selected transverse slices of each spinal cord section. In the control group the number of neurofilaments was expressed as 100%.

Statistical analysis. GraphPad Prism 6 software (GraphPad Software, Inc., La Jolla, CA, USA) was used for the statistical analyses. The differences in behavioral outcomes between individual groups were assessed using ordinary one-way ANOVA followed by Sidak’s multiple comparisons test with statistical significance set at P<0.05 for each week separately. The data collected from the white and grey matter sparing, number of motoneurons and number of neurofilaments were also subjected to ordinary one-way ANOVA followed by Sidak’s multiple comparison test; each spinal cord segment was assessed separately. The unpaired Student t-test was used to determine the effect of durotomy. The comparison between SCI group from Experiment 1 and SCI group from Experiment 2 was performed and P<0.05 was considered to indicate a statistically significant difference. Spearman’s correlation analysis was used to establish the relationship between spinal cord tissue integrity and the neurological outcome of the animals. The data from all 24 animals from both experiments were utilized. All data are presented as mean ± standard deviation.

Results

Experiment 1: Spinal cord exposed by durotomy

Behavioral assessment. Neurological outcomes of minipigs recorded during a 9-weeks survival period are summarized in Fig. 3. All animals displayed complete paraplegia immediately after SCI. In the SCI group, one week after SCI, two animals suffered from complete paraplegia and one was capable of slight movement in one hip joint (ANOVA summary for 1st week: F (4, 10)=575.8; P<0.0001; Control vs. SCI P<0.0001). During the whole survival period, the animals slowly recovered some of their hind-limb functions characterized by slight movement in all three joints. The outcome improved gradually and at the end of the survival period their motor score reached 7±0.0 (ANOVA summary for 9th week: F (4, 10)=59.05; P<0.0001; Control vs. SCI P<0.0001). At the 9th week, these animals were not able to support their body weight, and the motor activity of their hindlimbs consisted of extensive movements in all three joints and ‘sweeping’, which was usually restricted to one pelvic extremity.

None of the treatments improved the final motor scores compared to the SCI group. However, there were significantly lower scores in the 3rd, 4th, 7th and 8th weeks (SCI vs. en. medium P<0.05) in the neurological outcome of the group where the enriched medium was used. Final outcome was 3.0±2.65 (SCI vs. en. medium, P=0.0545). In the group treated with 4°C saline, there was transient improvement of motor functions between the 3rd and 7th week of the follow up period.
Macroscopic changes at the lesion site. After 9 weeks, the spinal cord was markedly narrowed at the injury site (Fig. 4Ab). When removing the dura mater, we noted extensive fibrotic adhesions between the dura and underlying spinal cord tissue. On transversely-cut 1 cm segments, the central part contained a distorted cavity spreading in both cranial and caudal directions from the epicenter. The size of the cavity seemed to be slightly reduced in hypothermia-treated groups. No differences could be seen in the animals treated with medium or enriched medium.

Histological outcome. The cross-sectional areas of spared white and grey matter tissue on histological sections, taken from seven consecutive spinal cord segments, were measured. Nine weeks after SCI, the lesion formed a distorted spindle-shaped cavity. The data from morphometric analyses form a V-shaped curve which corresponds to the amount of preserved tissue in the damaged spinal cord (Figs. 4 and 5).

The most affected area was the epicenter of the injury (segment 0) where the white matter sparing corresponded to 26.20±6.99. A detailed analysis of spared white matter was carried out by dividing it into dorsal, lateral and ventral columns (Fig. 4). We assessed the volume of preserved white matter columns and found differences indicating the largest loss of tissue in the dorsal columns. The volumes of preserved white matter ranked in descending order: Ventral columns (74.21%±6.73) > lateral columns (69.14%±6.05) > dorsal columns (61.69%±5.45).

Nine weeks after SCI, the gray matter preservation was quite low at the lesion site (20.99±6.59%) and in segments near the injury epicenter (48.26±7.53% at +1 and 65.00±7.56% at -1 segment) (Fig. 5A). In addition, we counted the number of motoneurons in the VHs of control animals was 16.58±1.54. SCI + durotomy significantly reduced the cumulative number of motoneurons to 6.17±1.73. A large
necrotic focus could be seen in the epicenter of the injury, where the number of motoneurons dropped to 0.39±1.04. The number of motoneurons increased gradually along with the distance from the epicenter (Fig. 5B). Occasionally occurring motoneurons were observed up to 1 cm in rostro-caudal direction; there were 3.11±2.70 cells rostrally and 5.72±0.75 cells caudally. The motoneurons were clearly detectable 2 cm from the epicenter, although their number was still quite low (5.94±1.80 rostrally and 8.06±1.00 caudally) in comparison with controls. There was only a slight increase in occurrence of motoneurons in segment +3 (9.5±1.79) and in segment -3 (10.44±3.05) (Fig. 5B).

Morphometric analyses indicated that individual treatments had positive effect on tissue sparing after SCI. The most prominent improvement was detected in the white matter, where 4˚C saline positively affected tissue sparing especially in the dorsal and lateral columns in almost all rostro-caudal segments. Hypothermia with medium and/or enriched medium positively affected the white matter integrity in the rostral segments of the dorso and lateral columns (Fig. 4B and C). Significant sparing (P<0.05) was also found in the caudal segment distant from the epicenter of the injury in the group treated with enriched medium (Fig. 4C). No significant differences were noted in the ventral columns. The hypothermia treatment was less pronounced in the grey matter. The largest effect could be seen after 4˚C saline treatment, especially in the segments distant from the epicenter. The most prominent improvement in number of motoneurons was noticed after hypothermia in the caudal segments (Fig. 5B). Significant improvement in the rostral segments was visible only in the saline or medium treated groups (Fig. 5B).

Immunohistochemistry. SMI-312 immunolabeling was performed to visualize individual axons in the white matter and to count their number in individual areas of interest (Figs. 2, 6 and 7). These data characterize the density of axons in selected areas of the dorsal, lateral and ventral columns, and their changes after SCI (Fig. 7). Nine weeks after SCI, loss of SMI-312-positive axons was significant in all white matter

Figure 6. Experiment 1. Representative images of neurofilament immunohistochemistry. Sections were taken from +1 segment of transverse spinal cord sections (dorsal, lateral and ventral columns) from (A-C) control, (D-F) after SCI and subsequent hypothermic treatment with (G-I) cold saline, (J-L) en. medium or (M-O) medium. Scale bar, 100 µm. SCI, spinal cord injury; en. medium, enriched medium.
columns along the rostro-caudal axis. The most affected were the lateral columns. The white matter disintegrity after SCI did not allow the number of neurofilaments to be quantified at the lesion site (segment 0).

Hypothermic treatment with 4°C saline increased the number of NF in all columns in comparison to the SCI. However, this improvement was significant (SCI vs. saline P<0.05) only in the lateral (+1 segment) and ventral columns (+1 and -3 segments). Hypothermia with medium negatively influenced the number of NF in the caudal segments (Fig. 7A-C).

**Experiment 2: Intact dura mater**

**Behavioral assessment.** The outcomes of minipigs in this experiment were very similar to those in Experiment 1 (Fig. 8). One week after SCI, the animals suffered complete paraplegia with almost no movement ability in all joints of the hind limbs (ANOVA summary for 1st week: F (2, 6)=495.5; P<0.0001; Control vs. SCI P<0.0001). At the end of the survival period, the score was 8.7±3.35 points (ANOVA summary for 9th week: F (2, 6)=13.41; P=0.0028; Control vs. SCI P=0.0051).

We did not notice any significant differences between the SCI group and the saline treated group (24°C) at any time point during the whole survival period. The final neurological score was 7.67±4.73 (SCI vs. Saline P=0.9683).

**Histological outcome.** The pattern of spinal cord tissue disintegrity after SCI was quite similar to that described in Experiment 1 (Figs. 9 and 10). At the lesion epicenter, the cumulative white matter sparing was 31.48%±7.45 and the spared gray matter was 23.7%±4.35. The volumes of preserved white matter ranked in descending order: ventral columns (77.91%±7.48) > dorsal columns (75.54%±5.89) > lateral columns (72.23%±6.35).

After 24°C saline treatment, there was an increase in white matter preservation (Fig. 9B-D). The area of preserved white matter was significant (SCI vs. saline P<0.05) at several levels of the evaluated segments in the dorsal and lateral columns. The largest differences were noted in the lateral columns where hypothermia increased white matter preservation by 13.71%±4.75 (Fig. 9C). This improvement was significant along the rostro-caudal axis from +2 to -2 segment. The gray matter integrity was influenced as well (Fig. 10). Quantitative analysis revealed overall improvement by 16.68±5.76% in comparison to the SCI group. The improvement was significant (SCI vs. saline P<0.05) in all segments except for the epicenter itself (Fig. 10A). The number of motoneurons was also improved by saline treatment (Fig. 10B). As shown in Fig. 10B, the number of motoneurons dropped from the initial value of 16.58±1.54 cells in the control group to cumulative 5.7±1.78 motoneurons in the SCI group. The number of motoneurons was significantly higher in all segments caudal to the injury epicenter (Fig. 10B). The cumulative number of motoneurons in the VHs in the saline-treated group increased to 6.9±1.57 cells.
Immunohistochemistry. Nine weeks after SCI, the density of neurofilaments strongly decreased in all white matter columns (Figs. 11 and 12). Again, the lowest density of SMI-312 positive axons (59.16±5.57% of spared neurofilaments) was identified in the lateral columns.

Although saline hypothermia had beneficial influence on neurofilaments' preservation in all white matter columns (Figs. 11 and 12), the effect of treatment was more prominent in the dorsal and the lateral columns than in the ventral columns (Fig. 12).

Durotomy vs. intact dura mater. To determine the effect of durotomy, the data from both SCI groups from Experiment 1 (durotomy) and Experiment 2 (intact dura mater) are summarized in Fig. 13. We detected significantly better neurological outcome in animals with intact dura mater at the 4th-6th weeks (P<0.05), but the differences were non-significant at the end of the survival period (Fig. 13A).

The animals with intact dura mater also showed improvement in white matter preservation (segments +3, +2, -2, -3) and in density of SMI-312 positive axons (from +3 to +1 segments) (Fig. 13B and D). In contrast, both the grey matter sparing and preservation of motoneurons deteriorated in segments distant from the lesion site (Fig. 13C and D).

Correlation. To determine the sensitivity of the individual outcome parameters, we utilized the data from all groups from experiments 1 and 2 (n=24) and performed a Spearman's correlation analysis between individual outcome parameters and final behavioral scores (Fig. 14). We found moderately positive correlation between the neurological score and the number of neurofilaments at 9 weeks (r=0.6746; P=0.0003) followed by weakly positive correlation between neurological score and number of motoneurons in the VHs (r=0.4660; P=0.0217), cumulative white matter sparing (r=0.4708; P=0.0202) and cumulative grey matter sparing (r=0.4760; P=0.0187).

Discussion

In the present study, we tested the efficacy of local hypothermia on the outcome parameters in a minipig SCI model utilizing a computer-controlled impact device. Two different approaches to local hypothermia treatment were compared: i) Post-SCI durotomy allowing direct access of perfusion solutions to the spinal cord tissue, and ii) post-SCI treatment with intact dura mater.

Comparison of the outcomes from both SCI groups (in Experiments 1 and 2) demonstrates that the durotomy itself had no impact on the final neurological outcome. However, the recovery of the animals treated with durotomy was noticeably slower and the neurological score of these minipigs was significantly worse from the 4th to 6th week. This finding was supported by our histological observations where the white matter outcome parameters showed significantly worsened tissue and axonal preservation in the group treated with durotomy. Based on data from the literature, durotomy itself performed soon after SCI could be beneficial. The positive effects are generally attributed to alleviated intraspinal pressure, subsequent improvement in blood perfusion of the injured tissues, and also to reduction of...
lesion volume due to attenuation of post-traumatic cystic cavitation (60-63). Beneficial influence achieved through durotomy applied after acute SCI was described as early as in 1988 by Perkins and Deane (64). Based on later studies, spinal cord decompression with durotomy has substantial effect on neurological outcome (60,62,63). It has been described, in terms of its application, not worsening the neurological state of the patients in clinical studies (65,66). Iannotti et al (67) reported beneficial effects of durotomy itself and combined with an allograft patch performed 4 h after injury. These researchers found that cases with durotomy showed improved recovery and significant reduction in lesion volume.

On the other hand, durotomy disrupts the natural circulation of cerebrospinal fluid (CSF), which can be associated with adverse effects. Smith et al (63) showed that durotomy itself had detrimental effect and increased scar and cavity formation. There are evidences that even more severe complications (e.g., low-CSF-pressure headache with meningitis or transient quadriplegia) could be associated with durotomy (68). Jones et al (69,70) performed two separate studies utilizing the porcine model of weight drop and compression SCI followed by decompression surgery four h after primary injury. Both studies describe in detail the eventual consequences of the performed durotomy. The first study showed that decompression following SCI resulted in varying degrees of spinal cord swelling, occlusion of the subarachnoid space and blockage of CSF flow. According to them, an important factor in this process is the initial severity of the injury. These observations may partly explain the lack of benefit of decompression in some patients. They suggest a need to reduce spinal cord swelling in order to optimize the clinical outcome after acute SCI (69). The second study posits that intradural swelling may induce secondary pathology due to interruption of normal CSF flow and constraints on swelling by the surrounding meninges, leading to inconclusive results of decompression and neurological outcome following SCI (70). In our study we observed spinal cord swelling in all groups which underwent durotomy. In spite of the treatment procedures, the swelling persisted after five h, when the dura mater was sutured in order to stop CSF leakage and development of meningitis. We assume that

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**Figure 10. Experiment 2.** Gray matter sparing and preservation of motoneurons 9 weeks after SCI and hypothermic treatment with saline performed on transverse sections dyed with Luxol fast blue and cresyl violet staining (A-B). Transverse spinal cord sections taken from caudal segments (-3) of (C) control, (D) after SCI and (E) treatment with saline. (F-H) High-power microphotographs of the boxed areas in C-E showing motoneurons. Data are the mean values ± standard deviation. Ordinary one-way analysis of variance followed by Sidak's multiple comparison test was used for the assessment of differences between experimental groups in each segment of spinal cord separately. (A) Gray matter sparing; †P<0.05 vs. Control; *P<0.05 vs. SCI. (B) Number of motor neurons in VHs; †P<0.0001, vs. Control; *P<0.05 vs. SCI. Scale bar in C-E, 2 mm; in F-H, 500 µm. SCI, spinal cord injury; VH, ventral horn.
delayed closure of the dura encircling the swollen segments of the spinal cord could be one of the reasons for the worsened outcomes noted in the animals that underwent durotomy.

Schumacher \textit{et al} \cite{71} reported an association between neurofilament loss and impaired hind-limb motor function after SCI. Significantly lower white matter integrity and reduction of neurofilaments in durotomy-treated animals could reflect multiple factors causing variable outcomes following the interruption of long-projection pyramidal (lateral cortico-spinal) and extrapyramidal (rubrospinal and reticulospinal) tracts. Neurofilaments are particularly abundant in large myelinated axons and are essential for axon radial growth and axon caliber maintenance during development \cite{72, 73}.

Wang \textit{et al} \cite{74} reported that NF gene transcriptional regulation is crucial for NF expression, predominantly in axonal regeneration and degenerative diseases.

The opposite effect was seen in gray matter. Quantitative analysis confirmed the loss of gray matter and motoneurons at the lesion site and in rostro-caudal segments up to 2 cm in both SCI models. These changes were less dramatic in the SCI-durotomy group but only in segments (+3 and -3), i.e., distant from the lesion site. The response of motoneurons to SCI depends on their ability to handle calcium. Previous results have indicated an unusually high influx of Ca\textsuperscript{2+} upon their stimulation \cite{75-77}. We have reported increased expression of parvalbumin, a Ca\textsuperscript{2+}-binding protein, in motoneurons ten days after Th9 transection \cite{78}. The results suggest that this Ca\textsuperscript{2+}-binding protein may rescue trauma-affected neuronal circuits involved in motor control.

\textit{Effects of hypothermia in Experiment 1}. When planning to carry out therapeutic local hypothermia in the acute stage of spinal cord compression in minipigs, we took into consideration two critical aspects, i.e., the cooling conditions and the penetration of perfusion solutions, both essential for the success of treatment. Under certain circumstances, radical hypothermia...
Figure 13. Comparison of outcome parameters in SCI groups (SCI opened and SCI intact dura mater). The animals survived for 9 weeks. Data are the mean values ± standard deviation. Unpaired Student t test was used for statistical evaluation of data. *P<0.05 SCI opened vs. SCI intact. (A) Behavioral recovery, (B) cumulative white matter sparing, (C) cumulative grey matter sparing, (D) number of neurofilaments and (E) number of motoneurons in VHs were assessed. SCI, spinal cord injury; VH, ventral horn.

Figure 14. Scatterplots illustrating relationship between final neurological score of animals from all experimental groups and (A) cumulative white matter sparing, (B) cumulative grey matter sparing, (C) number of neurofilaments and (D) number of motoneurons in spinal cord VHs. Indicated are the Spearman rank correlation coefficients (r) and the P-values. The number of XY pairs is 24. VH, ventral horn.
appears more promising than modest hypothermia (79-81). Vanický et al (82) reported that deep spinal cord hypothermia (t<15°C) via epidural cooling with 5°C saline provided effective protection against long-lasting spinal cord ischemia in rabbits. We have previously shown the benefit of five hours of local hypothermia (t=19°C) in an 8N and 15N computer-controlled compression model of SCI in minipig (57). The results indicated that modulation of the microcircuits leading to better functional outcome may depend on the preservation of axons in the lateral columns of cranial and caudal segments immediately adjacent to the lesion, and the protection of neurofilaments in the same segments of lateral funiculi (57).

In the present study, durotomy allowed direct penetration of solutions (saline, medium or enriched medium) through the outer surface of the uncovered spinal cord within 3 cm in cranio-caudal direction. However, none of these treatments improved the final neurological outcome, and quite surprisingly, the application of the cold enriched medium caused significant worsening of the scores which became apparent at the 4th week and persisted almost until the end of the survival period. Morphometric analyses of the lesion size indicated that individual treatment procedures did improve some tissue preservation in both the white and grey matter. Saline hypothermia activated the processes that led to better regeneration of the white matter within the dorsal and lateral columns in almost all cranio-caudal segments. We also found improvements in neurofilament expression and the sparing of grey matter/motoneurons. Earlier in vitro study showed that a particular neuronal population in monolayer cultures of dissociated murine spinal cord was especially sensitive to hypothermic stress (83). When the spinal cord cultures were exposed to the temperatures below 17°C, the neuronal perikarya and dendrites swelled, with the majority of the swollen neurons dying during the phase of rewarming to 37°C, while glial and other nonneuronal cell types were unaffected. The authors claimed that prolonged exposure (2-6 h) of monolayer cultures to the temperatures below 17°C caused significant neuronal death mediated primarily by the NMDA receptor-ion channel complexes in the dendro-somatic membranes. In the present study we report that the temperature of the saline (at 15 or 24°C) significantly reduced motoneuron death. Surface cooling of the spinal cord creates a substantial temperature gradient within the cord tissue (84) which may differently influence the viability of spinal motoneurons in in vivo experiments.

These protective effects can be attributed to transient hypothermia. Previously, strong protective effects of local spinal cord hypothermia had been demonstrated in spinal cord ischemia experiments (24,82,85,86). Local hypothermia was utilized in experimental studies after traumatic SCI as well. Cooling was generally initiated no later than 30 min after acute SCI, its duration varied between 20 min and 48 h and the temperature varied between 19°C up to 35°C. In general, these studies showed that hypothermia is more effective when initiated early after injury and lasts for a longer time. Shorter duration may require more profound cooling to achieve similar effect (35). In general, these data suggest that long periods of local hypothermia are tolerated well and may improve long-term tissue sparing and functional outcome (14,35,40-43,47,57,87).

Treatment with cold medium and/or enriched medium significantly preserved the white matter in dorsal (+3 segment) and lateral (+3 and +1 segment) columns located cranially to the lesion site. The medium negatively affected neurofilaments predominantly in the caudal segments and led to non-significantly lower behavioral outcome than in the SCI + durotomy group from the 7th to 9th week. In the case of the enriched medium-treated minipigs, the neurological outcome was very low during the whole survival period. It appears that low temperature of medium and/or enriched medium influenced the blood flow and slowed down the cell metabolism resulting in cell inability to profit from the ingredients contained in the cooling solutions. The mechanisms underlying protection by hypothermia are believed to be slowing down the enzymatic activity of reactions that require adenosine triphosphate (88). The results of our study confirm that exposure to low temperatures may be stressful for the cells treated with medium and/or enriched medium. It is still questionable whether increasing the target temperature during enriched medium/medium hypothermia could be able to improve histological/neurological outcomes.

Effect of hypothermia in Experiment 2. In Experiment 2 we tested a less invasive saline (24°C) treatment through the intact dura mater, minimizing the risk of spinal parenchymal infection. This moderate hypothermia treatment had significant effect on tissue preservation, but this effect was not associated with improved motor recovery. Several studies described similar results, where spinal cord hypothermia protected tissue from excessive loss without significant improvement in neurological function (41,57). Similarly, Teh et al (87) observed that hypothermia improved tissue preservation and neurological outcome during survival, but not the final neurological outcome. On the other hand, Casas et al (40) showed that regional cooling applied 30 min after moderate contusive SCI was not beneficial or detrimental in terms of tissue sparing, neuronal preservation, or locomotor outcome. They suggested, that this method of cooling may reduce blood flow in the injured spinal cord and exacerbate secondary injury.

The results of this pilot study extend the use of our reproducible minipig model for preclinical SCI experiments. The data indicate that further therapeutic interventions can modify the outcomes we have followed. The degree of injury we have used (18N SCI) is perhaps too severe for experimental therapy, and for observing any changes in neurological outcome, and less severe degrees of injury would be preferable. Our observations indicate that posttraumatic edema is a serious pathophysiological mechanism contributing to injury. When planning interventions with durotomy, prolonged duration of post-traumatic swelling should be taken into consideration. The optimal degree of hypothermia and its timing should be studied systematically. Further experimental/clinical studies are required to determine the optimal cooling parameters (therapeutic window, optimal temperature and duration) and to test the effectiveness of hypothermic therapy in various trauma-induced SCI models using large experimental animals.

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Availability of data and materials

The datasets used and/or analyzed during the current study are available from the corresponding authors on reasonable request.

Authors’ contributions

MMars, NL and JG were involved in creating the study concept and designing the experiments. VL, AT, IC and PR were involved in anesthetizing animals and post-surgical animal care. MMars, IS, MG, IL, JK, IS Jr., JG and JP were involved in spinal cord compression and implantation of the perfusion chamber. JG, SP and MZ prepared the perfusion chamber. JG, MZ, SP and JP assisted in hypothermic treatment administration. JP, MZ, SP, JG, KB and MB were involved in transcatheter perfusion and spinal cord tissue preparation. MZ, SP, ER, MMart, AS and AK performed the immunohistochemical staining and histological procedures. PR, MZ, SP and JG performed the behavioral assessments. MZ, NL and DM performed data analysis. MZ and NL wrote the paper.

Ethics approval and consent to participate

The experimental protocols were prepared in accordance with the EC Council Directive (2010/63/EU) regarding the use of animals in research and approved by the State Veterinary and Food Administration of the Slovak Republic (decision no. 1319/13-221) as well as by the Ethical Commission of the University of Veterinary Medicine and Pharmacy in Kosice, Slovak Republic. All efforts were made to minimize the size of experimental groups and animal suffering.

Patient consent for publication

Not applicable.

Competing interests

The authors declare no conflicts of interest.

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