Mechanisms underlying cell death in ischemia-like damage to the rat spinal cord in vitro

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New spinal cord injury (SCI) cases are frequently due to non-traumatic causes, including vascular disorders. To develop mechanism-based neuroprotective strategies for acute SCI requires full understanding of the early pathophysiological changes to prevent disability and paralysis. The aim of our study was to identify the molecular and cellular mechanisms of cell death triggered by a pathological medium (PM) mimicking ischemia in the rat spinal cord in vitro. We previously showed that extracellular Mg2+ (1 mM) worsened PM-induced damage and inhibited locomotor function. The present study indicated that 1 h of PM + Mg2+ application induced delayed pyknosis chiefly in the spinal white matter via overactivation of poly (ADP-ribose) polymerase 1 (PARP1), suggesting cell death mediated by the process of parthanatos that was largely suppressed by pharmacological block of PARP-1. Gray matter damage was less intense and concentrated in dorsal horn neurons and motoneurons that became immunoreactive for the mitochondrial apoptosis-inducing factor (the intracellular effector of parthanatos) translocated into the nucleus to induce chromatin condensation and DNA fragmentation. Immunoreactivity to TRPM ion channels believed to be involved in ischemic brain damage was also investigated. TRPM2 channel expression was enhanced 24 h later in dorsal horn and motoneurons, whereas TRPM7 channel expression concomitantly decreased. Conversely, TRPM7 expression was found earlier (3 h) in white matter cells, whereas TRPM2 remained undetectable. Simulating acute ischemic-like damage in vitro in the presence of Mg2+ showed how, during the first 24 h, this divalent cation unveiled differential vulnerability of white matter cells and motoneurons, with distinct changes in their TRPM expression.

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The clinical scenario of new spinal cord injury (SCI) cases is changing because trauma is no more the leading cause.1 Non-traumatic SCI originates from vertebral stenosis, tumors, or cardiovascular disorders that comprise a cluster of post-surgical paralysis after abdominal aorta surgery.1 Although non-traumatic lesions are usually incomplete, the clinical symptoms are severe, and include paralysis and sensory dysfunction. A typical feature of SCI is the delayed onset of a secondary lesion, with cell death affecting the initially spared neighboring tissue through complex mechanisms. This is an important issue, as a prerequisite to the development of mechanism-based neuroprotective strategies for acute SCI is a detailed understanding of the early pathophysiological changes. On the assumption that brain lesions are transiently associated with decreased extracellular Mg2+ concentration,2 even if this observation has not been systematically confirmed in humans,3 Mg2+ solution infusion was tested and was found to worsen the outcome in patients with traumatic brain lesions.4,5 This result was also observed in stroke patients.6 In accordance with this view are reports on the lack of effectiveness of exogenous Mg2+ on experimental SCI or neonatal brain injury.7,8

To explore the cell death mechanisms, our laboratory has developed an in vitro neonatal rat spinal cord model to mimic the metabolic perturbation (hypoxia, aglycemia, oxidative stress, acidosis, and toxic free radicals with low extracellular Mg2+ occurring in vivo after ischemic SCI.9 This model is based on the transient application of a toxic medium (termed ‘pathological medium’, PM) that recapitulates the chief elements of metabolic dysfunction occurring in vivo and depresses locomotor network function with cell death predominantly in the white matter via an apoptotic pathway. We have observed that more extensive pyknosis with block of locomotor network activity is evoked when PM is applied with a standard concentration (1 mM) of extracellular Mg2+ when compared with preparations treated with Mg-free PM.

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Abbreviations: AIF, apoptosis-inducing factor; AUs, arbitrary units; Chat, choline acetyltransferase; DAPI, 4,6-diamidino-2-phenylindole; DTT, dithiothreitol; EDTA, ethylenediaminetetraacetic acid; ELISA, enzyme-linked immunosorbent assay; H2AX, phospho-histone2A.X(Ser139); HEPES, 4-(2-hydroxyethyl)-1-piperazineethanesulfonic acid; L3, lumbar spinal cord segment 3; NeuN, neuronal nuclei; PM, pathological medium; PBS, phosphate-buffered solution; PJ34, N-(5,6-dihydro-6-oxo-2-phenanthridinyl)-2-acetamide; PMSF, phenylmethylsulfonyl fluoride; SNP, sodium nitroprusside; SCI, spinal cord injury; PAR, poly-ADP-ribose; PARP-1, poly (ADP-ribose) polymerase 1; ROI, region of interest; SMI 32, neurofilament H non-phosphorylated monoclonal antibody; T13, thoracic spinal cord segment 13; TRIS, tromethamine; TRPM2, transient receptor potential cation channel, subfamily M, member 2; TRPM7, transient receptor potential cation channel, subfamily M, member 7

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Although Mg²⁺ is essential for numerous enzymatic reactions,¹¹ novel targets of Mg²⁺ on the central nervous system have emerged. In particular, Mg²⁺ gates the activity of TRPM7, a bifunctional protein containing a protein kinase fused to an ion channel¹² and having a key role in neuronal death caused by oxidative stress and excitotoxicity.¹³ Further, TRPM2 (a member of the same channel family and modulated by Mg²⁺) is involved in the delayed death of brain neurons after experimental ischemia.¹³–¹⁵ The aim of the present report was to investigate whether the increased severity of spinal damage evoked by PM + Mg²⁺ was due to recruitment of additional cell death pathways, as well as study their identity and the potential involvement of TRPM2 and TRPM7.

Results

Cell death induced by PM + Mg²⁺. Our previous report¹⁰ indicated that a 1-h application of PM containing 1 mM Mg²⁺ induces, 24 h later, significant damage to the in vitro spinal cord. This observation was confirmed by detecting extensive pyknosis in the white matter region at 24 h (see arrow in the example of Figure 1a). Figure 1b indicates the four ROIs used for these experiments, whereas Figure 1c quantifies these data. Significant pyknosis already emerged 3 h after washing out the toxic medium (Figure 1c). It is noteworthy that sham preparations (kept in standard Krebs for up to 24 h) or mock preparations (bathed for 1 h in Mg-free Krebs solution to which 1 mM Mg²⁺ was added to test for any artifact due to solution application and washout procedure) showed no significant pyknosis (Figure 1c). Thus, unlike the outcome with PM without Mg²⁺ (Figure 1c), the presence of this divalent cation worsened an ischemia-like metabolic perturbation, and cell damage developed over 24 h.

Characterization of the white matter damage evoked by PM + Mg²⁺. Using biomarkers previously employed to characterize spinal injury,¹⁶,¹⁷ we next investigated the type of cell damage evoked by PM + Mg²⁺ by studying immunoreactivity to phospho-histone2A.X(Ser139) (H2AX a marker of DNA double-strand breaks)¹⁸ and active-caspase 3 (a marker of apoptosis)¹⁸ in the white matter ROI where extensive pyknosis was found (Figure 1c). Figure 2a shows that the percentage of H2AX-positive cells in the white matter region was higher in treated than in sham preparations at 3 or 24 h. Nevertheless, unlike the number of pyknotic nuclei (Figure 1c), the number of H2AX-positive elements did not increase from 3 to 24 h (Figure 2a), suggesting the presence of early DNA double-strand lesions in keeping with former studies.¹⁸,²⁰ By comparing the 24-h data in Figures 1c and 2a, it seems likely that the PM + Mg²⁺ damage affected the vast majority of white matter elements, as about 75% of them showed pyknosis and 25% had H2AX immunoreactivity. Interestingly, in the same ROI, immunopositivity for active caspase 3 (an important executor of apoptosis), although limited to <5% of the cells, was larger than that in the sham condition (Figure 2b).

The poor expression of active caspase 3 was an unexpected result, as PM (without Mg²⁺) is known to damage the spinal white matter primarily via caspase-dependent apoptosis.¹⁶ When the spinal damage is evoked by excitotoxicity, the main process of cell death is a non-apoptotic mechanism termed parthanatos caused

![Figure 1](image-url)
Figure 2  Characterization of white matter damage evoked by PM + Mg²⁺. (a) Left, examples of immunostaining for H2AX in the ventral quadrant of the spinal cord in which the white matter shows minimal signal positivity in sham conditions. Scale bar: 30 µm; right, percentage of H2AX-positive cells in the ventrolateral white matter after PM + Mg²⁺; n = 9 sections, ***P < 0.001. (b) Left, examples of immunostaining for active caspase 3 in the same ROIs as shown in (a); right, percentage of caspase 3-positive cells in the ventrolateral white matter after PM + Mg²⁺; n = 9 sections, *P = 0.034; ***P < 0.001. (c) Histograms showing pyknotic nuclei after addition (on PM + Mg²⁺ washout) of caspase 3 inhibitor (1 µM), PJ34 (60 µM), or both. n = 9; ***P < 0.001. For data shown in (a), (b) and (c), the Mann–Whitney test was used after performing an ANOVA Tukey test. (d) Histograms show PAR tissue levels in various experimental conditions. For each test, data are from three spinal cords tested in duplicate; one-way ANOVA Tukey test; ***P < 0.001.
by excessive activation of PARP-1 (poly (ADP-ribose) polymerase 1) to synthesize toxic concentrations of PAR (poly-ADP-ribose) that disrupt cell energy stores.\textsuperscript{21} Therefore, in the present report we investigated whether, in the white matter ROI, PM + Mg\textsuperscript{2+} could stimulate excessive production of PAR, which is known to induce pyknosis.\textsuperscript{21,22} Thus, we tested whether application of a selective inhibitor of either PARP-1 activity or caspase-3 might protect white matter elements from PM + Mg\textsuperscript{2+} toxicity. To this end, we used PJ34 (N-(5,6-dihydro-6-oxo-2-phenanthridinyl)-2-acetamide) to block PARP-1\textsuperscript{23} and the caspase peptide inhibitor (caspase 3 inhibitor II, Z-DEVD-fmk) to block apoptosis,\textsuperscript{24} which were applied (for 24 h) either alone or in combination immediately after washing out PM + Mg\textsuperscript{2+}. Figure 2c summarizes these data: whereas the caspase inhibitor alone did not significantly reduce pyknosis (5% change; see Figure 2c), PJ34 significantly (P<0.001) decreased pyknosis by >25%, an effect even stronger (>35%) when the two pharmacological blockers were applied together (P<0.001), indicating partial convergence of these two cell death processes. Neither inhibitor had any effect on sham preparations (Figure 2c). Figure 2d indicates that PJ34 applied at a concentration of 60 μM was a strong inhibitor of the tissue generation of PAR measured with an enzyme-linked immunosorbent assay (ELISA). Halving the concentration of PJ was less effective.

Figure 3a illustrates an example of co-staining with 4,6-diamidino-2-phenylindole (DAPI) and the PAR antibody of a pyknotic white matter cell 24 h after PM + Mg\textsuperscript{2+} application. Figure 3b shows that, among the total DAPI-stained elements, the percentage of PAR-positive cells in white matter ROI (hatched bars) had already increased significantly (P<0.001) 3 h after PM + Mg\textsuperscript{2+}, and it remained elevated (P<0.001) 24 h later. Figure 3b (filled bars) also quantifies co-occurrence of PAR immunopositivity together with pyknosis for the total number of pyknotic cells; it is clear that PM + Mg\textsuperscript{2+} induced a significant (P<0.001) increase in the association between PAR and pyknosis at 3 and 24 h. In fact, few cells were PAR positive without pyknotic nucleus at either time (open bars). This finding suggested that PM + Mg\textsuperscript{2+} had transformed the white matter damage from the apoptosis previously detected with PM alone\textsuperscript{10} to a process with parthanatos characteristics. We thus wondered whether an analogous process might have developed in the gray matter.

Characterization of gray matter damage evoked by PM + Mg\textsuperscript{2+}. Despite the comparatively modest extent of the lesion to the gray matter induced by PM + Mg\textsuperscript{2+} (see Figure 1c), this protocol is known to inhibit locomotor network function.\textsuperscript{10} To further analyze neuronal damage, the present study quantified the percentage of NeuN-positive neurons in dorsal, central, and ventral ROIs at 3 and 24 h after PM + Mg\textsuperscript{2+}. Figure 4 demonstrates that significant neuronal loss was absent at 3 h, and became apparent at 24 h as far as the dorsal and ventral ROIs were concerned. Thus, in accordance with pyknosis data (Figure 1c), PM + Mg\textsuperscript{2+} had an early effect on white matter elements, and a later one on gray matter cells, which, in the ventral horn, constitute a small minority of the spinal cell population.\textsuperscript{25}

As, among gray matter regions, the late damage by PM + Mg\textsuperscript{2+} appeared to mostly affect the ventral ROI that contains motoneurons, namely, the essential output elements of the locomotor network, we next investigated dysfunctional changes in motoneurons, identified as large (>25 μm somatic diameter) cells located in the ventral horn and immunopositive for choline acetyltransferase (ChAT, the cytoplasmic ACh synthetic enzyme), and for SMI32, a marker for motoneuron cytoskeleton.\textsuperscript{26,27} Figure 5a shows that the number of ChAT-positive motoneurons did not change following PM + Mg\textsuperscript{2+} at 3 or 24 h, whereas the average intensity of the SMI32 signal was significantly (P<0.001) decreased at 3 and 24 h (Figure 5b). This observation suggests that, despite the...
unaltered number of motoneurons, these cells showed signs of cytoskeleton damage.

Neuronal distress can strongly activate PARP-1 to generate toxic levels of PAR (‘parthanatos’). In accordance with this notion, a significant increase in PAR immunoreactivity was detected in motoneurons early after PM+Mg2+, as exemplified in Figure 5c and quantified in Figure 5d. It is noteworthy that the PAR signal intensity did not increase further at 24 h.

The toxic action of PAR is typically executed via translocation of the transcription factor apoptosis-inducing factor (AIF) from the mitochondria to the nucleus and it thus represents a delayed cell death signal. We examined the expression and translocation of AIF in motoneurons as indicated in Figures 6a and b. In sham conditions, the confocal line scan of the motoneuron cell body (see line across cell soma in Figure 6a, left) demonstrated that its DAPI staining was confined to the cell nucleus (blue line; Figure 6a, right), whereas AIF (red line in Figure 6a, right) was poorly expressed throughout the cell. In contrast, 24 h after PM + Mg2+ (Figure 6b), whereas DAPI staining was still restricted to the nucleus, AIF immunoreactivity was diffused and extended to the cell nucleus as well. Detection of AIF in the motoneuron nucleus was accompanied by loss of expression of the nuclear transcription factor NeuN as shown in Figure 6c (see open arrow). Finally, Figure 6d indicates that, 24 h after PM + Mg2+, about 30% of motoneurons showed AIF nuclear translocation. All together, these observations suggest that, despite the persistence of ChAT immunoreactivity, a significant fraction of motoneurons displayed multiple signs of distress 24 h after PM + Mg2+, consistent with the onset of a parthanatos-like process.

Expression of TRPM7 or TRPM2 after PM + Mg2+ application. TRPM7 and TRPM2 are members of a large family of ion channels believed to play an important role in the neurodegeneration evoked by ischemia in vivo or by ischemia-like conditions in vitro. In the present study, we found undetectable TRPM7 immunopositivity in the white matter from freshly frozen preparations (Figure 7a, open circle), or under 3 and 24 h sham conditions (Figure 7a,
open squares), or immediately after washing out PM + Mg^{2+} (Figure 7a, filled triangle). Vice versa, a strong signal became apparent 3 h after PM + Mg^{2+} washout and remained stable 24 h later (Figure 7a, filled triangles). White matter TRPM7 immunoreactivity was most frequently detected in pyknotic cells (see arrows in Figure 7b) as quantified in Figure 7c (filled bars). In the white matter ROI, there was no significant TRPM2 immunoreactivity under treated or sham conditions (not shown). Thus, it appeared that about half of the white matter elements that died (pyknosis) after PM + Mg^{2+} treatment also expressed strong positivity to TRPM7.

In contrast to these data from the white matter ROI, in the three gray matter ROIs we found baseline positivity to TRPM7 (Figures 8a and b) and TRPM2 (Figure 8c). After washing out PM + Mg^{2+}, the TRPM7 signal was found significantly lower 24 h later (Figures 8a and b) in the dorsal horn ROI (P < 0.001) and in the motoneuron ROI (P = 0.002). Conversely, for the same ROIs, TRPM2 immunopositivity appeared to move in the opposite direction with a delayed increase 24 h after washing out PM + Mg^{2+} (Figure 8c).

**Discussion**

The focus of the present study was the role of extracellular Mg^{2+} in the genesis of cell death after transient application of a solution mimicking ischemia conditions. The present data...
relatively rapid DNA damage. The canonical marker of apoptosis, namely, activated caspase 3, was evident in a small minority of white matter cells only. Perhaps the damage had evolved so quickly that cell death had become apparent before any extensive apoptosis could be produced. In keeping with this view was our observation that a pharmacological inhibitor of apoptosis did not arrest pyknosis, although it reduced pyknosis when coapplied with PJ34, the inhibitor of PARP-1, which is essential for DNA damage repair and whose hyperactivation generates the toxic product PAR. PAR was already strongly increased 3 h after the end of PM application, suggesting a narrow time window before irreversible cell damage could occur (demonstrated by colocalization of PAR and pyknosis). When PJ34 is administered early during the lesion protocol, a degree of functional neuroprotection is observed in accordance with the present data. In general, the present results suggest that the presence of Mg2+ during the ischemic-like protocol aggravated the damage to the white matter cells by promoting the activation of at least one major cell death pathway, namely, parthanatos.

Gray matter damage by PM + Mg2+. Alongside low-scale pyknosis, most NeuN-positive neurons exhibited a gradient of neuronal loss: the strongest in the ventral horn, a modest one in the dorsal horn, and apparently none in the central region. Motoneurons, however, did not disappear, as their typical marker ChAT was preserved even 24 h later. These cells were, nevertheless, showing signs of distress because their SMI32 positivity was decreased, suggesting disturbance of their cytoskeleton. Conversely, a traumatic lesion in vivo is associated, a few days later, with loss of ChAT, followed by upregulation of SMI32 expression by surviving motoneurons, taken as an index of cell repair. The differential evolution of ChAT and SMI32 in the present study compared with in vivo experiments is unlikely due to motoneuronal maturation, as neonatal motoneurons already show adult somatic size. Accordingly, in the in vitro spinal cord model, the number of SMI32- and ChAT-positive cells and their staining pattern remain constant in the first week of life. Further, AIF-dependent cell death mediates neuronal death after hypoxia–ischemia in the neonatal rat brain as much as in the adult brain. Hence, dynamic changes in these two biomarkers may indicate distinct processes and time course underlying either degenerative mechanisms or restoration.

Concomitant with the fall in SMI32 signal was the detection of rising PAR immunoreactivity in the same cells. This was likely a sign of impending death, as it was accompanied by strong AIF translocation into the nucleus, a known effector of cell death. Interestingly, AIF nuclear translocation was closely associated with loss of NeuN positivity. NeuN is an intrinsic component of the neuronal nuclear matrix, whose fast degradation (already at 1 h after injury) precedes cytoskeletal changes (identified, for instance, with SMI32 positivity) in injured motoneurons.

These results imply that various motoneuron biomarkers should be tested for assessing damage in pathological conditions, as their significance in the diagnosis of cell death is time-dependent. Thus, we surmise that the development of A model of non-traumatic injury. The growing number of non-traumatic causes of SCI led us to examine the processes responsible for early cell death after the in vitro insult. Our previous investigations indicated that applying a toxic solution containing free oxygen radicals with no oxygen or glucose supply especially damaged the white matter elements, with moderate impairment of locomotor-like patterns that were still present at regular frequency but with clearly lower amplitude. Nevertheless, when the same toxic solution contained the standard extracellular Mg2+ concentration, the histological damage was intensified and locomotor patterns were inhibited. The present report sought to characterize the processes responsible for this damage enhancement.

PM + Mg2+ evoked lesion of white matter elements. This solution was confirmed to produce stronger damage than the original PM (without added Mg2+), as pyknosis was seen in about 80% of the ROI white matter elements and was accompanied by early phosphohistone positivity to indicate
parthanatos in motoneurons occurred on a slower time base than in the white matter. Nevertheless, these pathological changes have a strong negative impact on locomotor network function.10

TRPM2 and TRPM7 roles in the PM\textsuperscript{+}Mg\textsuperscript{2+} effects. These two membrane channels have recently been investigated for their contribution to ischemic cell death in the brain.\textsuperscript{13,35} These channels are gated open by intracellular PAR, reactive oxygen species and Mg\textsuperscript{2+} (as under normal conditions such channels are minimally activated).\textsuperscript{13,14,42–44}

For these reasons, we studied how these channels were expressed during our experimental protocol. In view of reports on the ubiquitous occurrence of TRPM7 expression,\textsuperscript{45} their immunohistochemical absence in the white matter under sham conditions was not anticipated. This phenomenon could not be attributed to loss of protein expression by keeping the preparation in vitro because analogous lack of signal was found when the tissue was frozen immediately after dissection. It seems feasible that immaturity of the neonatal spinal cord was a factor for poor TRPM7 expression in controls. It was, however, clear that already 3 h after PM\textsuperscript{+}Mg\textsuperscript{2+} exposure a large number of white matter cells (with pyknotic nucleus) were positive to

Figure 8  Gray matter expression of TRPM7 or TRPM2 after PM\textsuperscript{+}Mg\textsuperscript{2+} application. (a) Examples of TRPM7 immunoreactivity in the ventral horn under sham conditions or 24 h after PM\textsuperscript{+}Mg\textsuperscript{2+}. (b) Histograms show distribution of TRPM7 immunoreactivity in dorsal, central and motoneuron ROIs. Note the significant fall in average signal intensity 24 h after PM\textsuperscript{+}Mg\textsuperscript{2+} for dorsal and motoneuron ROIs; **P = 0.002; ***P < 0.001. (c) Examples of TRPM2 immunoreactivity in ventral horn under sham conditions or 24 h after PM\textsuperscript{+}Mg\textsuperscript{2+}. (d) Histograms show distribution of TRPM2 immunoreactivity in dorsal, central and motoneuron ROIs. Note the significant increase in TRPM2 signal in all three ROIs examined; one-way ANOVA Tukey test, **P = 0.004; ***P < 0.001. All data are from nine spinal sections from three rats.
TRPM7. Our interpretation is that metabolic distress in the presence of Mg\(^{2+}\) triggered rapid trafficking of TRPM7 proteins to the cell membrane, where their activation might have contributed to the observed damage.

Unlike the white matter cells, in the gray matter, baseline expression of TRPM7 fell in motoneurons and dorsal neurons 24 h after PM + Mg\(^{2+}\) when low-scale pyknosis was manifested. TRPM2 expression (that could not be observed in the white matter) showed an opposite pattern with a late rise in gray matter neurons, which was especially strong in the central region. It is not feasible to relate the intensity of immunoreactivity to channel activation; thus, the present observations indicate a significant dichotomy in the biochemical conditions believed to occur at the site of the lesion, we used a metabolic perturbation protocol consisting of the transient application of PM (see below) with the addition of 1 mM Mg\(^{2+}\). This protocol is known to induce a pathological condition that includes locomotor network inhibition and histological damage. Thus, a solution containing 10 mM H\(_2\)O\(_2\), 500 mM sodium nitroprusside (SNP) and 1 mM Mg\(^{2+}\) and lacking oxygen and glucose was applied for 1 h. NaHCO\(_3\) was omitted and 4-(2-hydroxyethyl)-1-piperazineethanesulfonic acid (HEPES) was added to reach the pH range of 6.75–6.80 (with 0.1 N NaOH) and the osmolarity was lowered to 230–240 mOsm. This solution recapitulates the biochemical derangement believed to occur in the spinal cord after acute non-traumatic injury. After a 1-h application of PM + Mg\(^{2+}\), the latter solution was washed out and standard Kreb’s solution was superfused again for up to 24 h, after which preparations were fixed as previously reported and processed for histology as detailed below. Sham experiments were performed by keeping the preparations for analogous times in the same experimental setup without applying a toxic medium.

Quantification of dead cells. Using a Zeiss Axioskop2 microscope (Oberkochen, Germany) and Metavue software (Metamorph suite sold by Molecular Devices, Sunnyvale, CA, USA), cell counting was performed after DAPI staining or NeuN positivity (for neurons). Data were counted with ‘eCELLence’ (Molecular Devices, Sunnyvale, CA, USA) and lacking oxygen and glucose was applied for 1 h. NaHCO\(_3\) was omitted and 4-(2-hydroxyethyl)-1-piperazineethanesulfonic acid (HEPES) was added to reach the pH range of 6.75–6.80 (with 0.1 N NaOH) and the osmolarity was lowered to 230–240 mOsm. This solution recapitulates the biochemical derangement believed to occur in the spinal cord after acute non-traumatic injury.

Role of extracellular Mg\(^{2+}\). From the point of view of an in vitro spinal model, it is apparent that the standard concentration (1 mM) of Mg\(^{2+}\) amplified white matter damage and extended it to the gray matter. This might reside in the differential expression of TRPM7 and TRPM2, plus the degree of PAR hyperproduction at the early stage of damage, because the concentrations of PAR and intracellular Mg\(^{2+}\) are potent modulators of the activity of these channels. Recent studies with in vitro brainstem neurons have proposed that changes in intracellular Mg\(^{2+}\) concentrations occurring during pathological states can affect the conductance of neuronal Cx36 gap junctions and presumably shape the damage outcome. Although chemical synaptic inputs and Cx36-dependent electrical synapses contribute to synchronization of spinal motoneuron function, the expressions of Cx36 and Cx32 mRNA and proteins are sparse and unchanged after rat SCI. The complexity of the processes underlying ischemia-induced damage in the spinal cord even when studied with an in vitro model implies diversity of molecular pathways with distinct time dependence and cell specificity, factors that contribute to the difficulty of implementing successful neuroprotection in vivo. The present report suggests that the extracellular level of Mg\(^{2+}\) largely affects cell death mechanisms in the spinal cord and extends damage to motoneurons by facilitating the onset of parthanatos.

Materials and Methods

Rat spinal cord preparation. The experiments were performed on neonatal Wistar rats of postnatal age 0–1 days in accordance with the guidelines of the National Institutes of Health and the Italian act D.Lgs. 271/92 no. 116 (implementing the European Community directives no. 86/609 and 93/98), and the protocols were approved by the SISSA ethical committee for animal experimentation. We minimized the number of animals used in order to allow clear statistical analysis. Spinal cords were carefully dissected out from pups under urethane anesthesia (0.2 ml i.p. of a 10% w/v solution) at room temperature, with continuous superfusion with Kreb’s solution containing (in mM) the following: NaCl, 113; KCl, 4.5; MgCl\(_2\)/7H\(_2\)O, 1; CaCl\(_2\), 2; NaH\(_2\)PO\(_4\), 1; NaHCO\(_3\), 25; glucose, 11; gassed with 95% O\(_2\)/5% CO\(_2\), pH 7.4 at room temperature (22 °C), as described previously.

Protocol for spinal cord lesion. Our aim was to generate an experimental condition that might mimic the acute clinical setting when the metabolic insult is often temporary because of intensive care treatment to correct the metabolic derangement. In accordance with our studies and to simulate the biochemical conditions believed to occur at the site of the lesion, we used a metabolic perturbation protocol consisting of the transient application of PM (see below) with the addition of 1 mM Mg\(^{2+}\). This protocol is known to induce a pathological condition that includes locomotor network inhibition and histological damage. Thus, a solution containing 10 mM H\(_2\)O\(_2\), 500 mM sodium nitroprusside (SNP) and 1 mM Mg\(^{2+}\) and lacking oxygen and glucose was applied for 1 h. NaHCO\(_3\) was omitted and 4-(2-hydroxyethyl)-1-piperazineethanesulfonic acid (HEPES) was added to reach the pH range of 6.75–6.80 (with 0.1 N NaOH) and the osmolarity was lowered to 230–240 mOsm. This solution recapitulates the biochemical derangement believed to occur in the spinal cord after acute non-traumatic injury.

After a 1-h application of PM + Mg\(^{2+}\), the latter solution was washed out and standard Kreb’s solution was superfused again for up to 24 h, after which preparations were fixed as previously reported and processed for histology as detailed below. Sham experiments were performed by keeping the preparations for analogous times in the same experimental setup without applying a toxic medium.

Immunofluorescence procedure. Immunostaining was performed as described previously. Briefly, paraformaldehyde-fixed spinal cords were cryoprotected with 30% sucrose and sectioned (30 μm) with a sliding microtome. In accordance with our former studies, we analyzed spinal regions between T13 and L3 (thoracic spinal cord segment 3 to lumbar spinal cord segment 3) segments were examined. Pyknosis was readily observable as a change in nuclear morphology resulting from chromatin condensation.

Immunofluorescence procedure. Immunostaining was performed as previously described. Briefly, paraformaldehyde-fixed spinal cords were cryoprotected with 30% sucrose and sectioned (30 μm) with a sliding microtome. In accordance with our former studies, we analyzed spinal regions between T13 and L3 segments. For all antibodies (except anti-PAR), after incubation in blocking solution (5% normal goat serum, 5% bovine serum albumin, 0.3% Triton-X 100) for 1 h at room temperature, the primary antibody was incubated at 4 °C overnight. Before incubating tissue sections with the PAR antibody, an
fraction was validated by looking for (with western immunoblotting) any mitochondrial contamination indicated by the cytochrome C oxidase IV protein (Abcam, Cambridge, UK) in accordance with the method of Beirowski et al. 53 These samples were then used for ELISA.

ELISA quantification of PAR. To obtain evidence for PARP-1 activation and pharmacological inhibition by N- (5-6-Dihydro-6-oxo-2-phenanthrylidyl)-2-acetamide (P34), we quantified its product PAR in accordance with Nasrabady et al. 55 For this purpose, lysates of spinal cords treated with PM + Mg and/or P34, or in sham conditions, were collected 24 h later and their PAR levels were measured using the PARP in vivo pharmacodynamic assay II (http://www.trevigen.com/protocols) following the manufacturer's protocol (Trevigen, Bologna, Italy). This assay is based on ELISA immunoreactivity resulting in chemiluminescence signals recorded with a Glomax multi-detection system (Promega, Milan, Italy). Whenever tests were performed for the action of P34, this substance was added to the cell lysis buffer to avoid inhibitor dilution as per the manufacturer's instructions. All samples were run in duplicate. After background subtraction (blank samples), the net PAR levels were quantified using a standard linear plot (r = 0.98) based on the immunoreactivity induced by known concentrations (20–1000 pM) of PAR. Data were expressed as pg/mL/100 µg of protein (estimated as indicated below).

Drugs. SNP and P34 were purchased from Sigma; H2O2 was obtained from Caido Erba Reagents (Milan, Italy); and active caspase 3 inhibitor II was obtained from CalBiochem-Millipore (Billerica, MA, USA).

Data analysis. The data are shown as mean ± S.D. Statistical analysis was carried out with SigmaStat 3.1 (Systat Software, Chicago, IL, USA). Parametric and nonparametric data were first distinguished with a normality test (Anova Tukey) and analyzed with the Student t-test and the Mann–Whitney test, respectively, in accordance with the software choice. The significance level was always P<0.05, and n indicates the number of tissue sections.

Conflict of Interest
The authors declare no conflict of interest.

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1. van den Berg ME, Castellote JM, Mahillo-Fernandez I, de Pedro-Cuesta J. Incidence of spinal cord injury worldwide: a systematic review. Neuropediatrics 2010; 41: 184-192.
2. Lin MC, Huang YL, Liu HW, Yang DY, Lee CP, Yang LL et al. On-line microdialysis-graftite fumace atomic absorption spectrometry in the determination of brain magnesium levels in gerbils subjected to cerebral ischemia/reperfusion. J Am Coll Nutr 2004; 23: 561S–565S.
3. Valadka AB, Goodman JC, Copinath SP, Luzara M, Robertson CS. Comparison of brain tissue oxygen tension to microdialysis-based measures of cerebral ischemia in fatal head-injured humans. J Neurotrauma 1998; 15: 509-519.
4. Maas AI, Murray GD. Magnesium for neuroprotection after traumatic brain injury. Lancet Neurol 2007; 6: 20–21.
5. Temkin NR, Anderson GD, Winn HR, Ellenbogen RG, Britz GW, Schuster J et al. Magnesium sulfate for neuroprotection after traumatic brain injury: a randomised controlled trial. Lancet Neurol 2007; 6: 29–38.
6. IMAGES. Magnesium for acute stroke (intravenous Magnesium Efficacy in Stroke Trial): randomised controlled trial. Lancet 2004; 363: 439–445.
7. Perlman JM. Intervention strategies for neonatal hypoxic-ischemic cerebral injury. Clin Ther 2002; 26: 1353–1365.
8. Saeki H, Matsumoto M, Kaneko S, Tsuuta S, Cui YJ, Ohtake K et al. Is intrathecal magnesium sulfate safe and protective against ischemic spinal cord injury in rabbits? Anesth Analg 2004; 99: 1805–1812.
9. Taccola G, Margarayan G, Madnic M, Nistri A. Kainate and metabolic perturbation mimicking spinal injury differently contribute to early damage of locomotor networks in the in vitro neonatal rat spinal cord. Neuroscience 2008; 155: 538–555.
10. Margaryan G, Mladinic M, Mattioli C, Nistri A. Extracellular magnesium enhances the damage to locomotor networks produced by metabolic perturbation mimicking spinal injury in the neonatal rat spinal cord in vitro. Neuroscience 2009; 163: 669–682.

11. Ryazanova L, Landon LJ, Zierler S, Hu Z, Gali J, Yamaguchi TP et al. TRPM7 is essential for Mg2+ homeostasis in mammals. Nat Commun 2010; 1: 109.

12. Schmitz C, Peraud AL, Johnson CD, Inaba K, Smith MK, Penner R et al. Regulation of vertebrate cellular Mg2+ homeostasis by TRPM7. Cell 2003; 114: 191–202.

13. Aarts MM, Tymianski M. TRPMs and neuronal cell death. Pflugers Arch 2005; 451: 243–246.

14. Kisten-FJ, Heiner I, Luckoff A. TRPM2: a calcium influx pathway regulated by oxidative stress and the novel second messenger ADP-ribose. Pflugers Arch 2005; 451: 212–219.

15. Sumoza-Toledo A, Penner R. TRPM2: a multifunctional ion channel for calcium signalling. J Physiol 2011; 589: 1515–1525.

16. Kuzhandaivel A, Margaryan G, Nistri A, Mladinic M. Extensive glial apoptosis develops in vitro and in vivo in response to extracellular magnesium in the neonatal rat spinal cord. J Physiol 2010; 589: 325–338.

17. Kuzhandaivel A, Nistri A, Mazzone GL, Mladinic M. Molecular mechanisms underlying cell death in spinal networks in relation to locomotor activity after acute injury in vivo. Front Cell Neurosci 2011; 5: 9.

18. Plesca D, Mazumder S, Almasan A. DNA damage response and apoptosis. Methods Enzymol 2008; 446: 107–122.

19. Cohen GM. Caspases: the executioners of apoptosis. Biochem J 1997; 326: 1–16.

20. Cook PJ, Ju BG, Telesforo F, Wang X, Glass CK, Rosenfeld MG. Tyrosine dephosphorylation of H2AX modulates apoptosis and survival decisions. Nature 2003; 428: 591–596.

21. Kuzhandaivel A, Nistri A, Mladinic M. Kainate-mediated excitotoxicity induces neuronal death in the rat spinal cord in vitro. Neuroscience 2010; 169: 325–338.

22. Yoon SW, Andrabi SA, Wang H, Kim NS, Poirier GG, Dawson TM et al. Inhibitor of poly(ADP-ribose) polymerase-1 activity (PARP-1), on locomotor networks of the rat neonatal spinal cord. J Neurotrauma 2003; 20: 1515–1525.

23. Yu SW, Andrabi SA, Wang H, Kim NS, Poirier GG, Dawson TM et al. Inhibitor of poly (ADP-ribose) polymerase (PARP) in vitro and in vivo models of stroke. Int J Mol Med 2001; 7: 255–260.

24. Vandenberghe P, Vanden Berghe T, Fespessi N. Caspase inhibitors promote alternative cell death pathways. Sci STKE 2006; 2006: pe44.

25. Cifta A, Mazzone GL, Nani F, Nistri A, Mladinic M. Postnatal developmental profile of neurons and glia in motor nuclei of the brainstem and spinal cord, and its comparison with organotypic slice cultures. Dev Neurobiol 2012; 72: 1140–1165.

26. Mladinic M, Nakamura K, Usuda T, Tochikura S. Acute spinal cord injury in vitro: insight into basic mechanisms. In Aldskogius H (ed) Animal Models in Spinal Cord Repair. Springer Business Media, LLC: Humana Press: Heidelberg/New York, 2013, pp 39–63.

27. Carleto SQ, Yin HZ, Lamberta R, Weiss JH. In vitro kainate injury to large SMi-32+ spinal neurons is Ca2+ dependent. Neuroreport 1995; 6: 945–948.

28. Andrabi SA, Kim NS, Yu SW, Wang H, Koh DW, Saasaki M et al. Poly(ADP-ribose) (PAR) polymer is a death signal. Nat Proc Nat Acad Sci USA 2006; 103: 18308–18313.

29. Yu SW, Andrabi SA, Wang H, Kim NS, Poirier GG, Dawson TM et al. Apoptosis-inducing factor mediates poly(ADP-ribose) (PARP) polymer-induced cell death. Proc Nat Acad Sci USA 2006; 103: 235–242.

30. Miller BA. The role of TRP channels in oxidative stress-induced cell death. J Membr Biol 2006; 209: 31–41.

31. Nicotera P, Biagini A. Pantothenate kinase 1 (PANK1) is an anoxic neuronal death. J Neurochem 2003; 203: 1851–1865.

32. Penner R, Flagg A, The Mg2+ and Mg2+-nucleotide-regulated channel kinase TRPM7. Handb Exp Pharmacol 2007; 179: 313–328.

33. Xie YF, Macdonald JF, Jackson MF. TRPM2 calcium and neurodegenerative diseases. Int J Physiol Pathophysiol Pharmacol 2010; 2: 95–103.

34. Kuzhandaivel A, Nistri A, Mladinic M. Extensive glial apoptosis develops in vitro and in vivo in response to extracellular magnesium in the neonatal rat spinal cord. J Physiol 2010; 589: 325–338.

35. Palacios-Prado N, Hoge G, Marandisky A, Rimikute L, Chapuis S, Paulaukase N et al. Intracellular magnesium-dependent modulation of gap junction channels formed by neuronal connexins. J Neurochem 2003; 85: 4741–4753.

36. Tresch MC, Kiehn O. Synchronization of motor neurons during locomotion in the neonatal rat: predictors and mechanisms. J Neurosci 2002; 22: 9997–10008.

37. Lee HH, Lin DDR, Kiehn O, Widener J, Olson L. Glial and neuronal connexin expression patterns in the rat spinal cord during development and following injury. J Comp Neurol 2005; 489: 1–10.

38. Faden AI, Stocca B. Neuroprotection: challenges and opportunities. Arch Neurotol 2007; 64: 794–800.

39. Savitz SI, Fisher M. Future of neuroprotection for acute stroke: in the aftermath of the SAINT trials. Ann Neurol 2007; 61: 396–402.

40. Thuet S, Moon LD, Gage FH. Therapeutic interventions after spinal cord injury. Nat Rev Neurol 2006; 2: 628–643.

41. Burgoyne LA. The mechanisms of pyknosis: hypercondensation and death. Exp Cell Res 1999; 248: 214–222.

42. Hara Y, Wako M, Ishi M, Maeno E, Nishida M, Yoshioka T et al. LTRPC2 Ca2+ permeable channel activated by changes in redox status confers susceptibility to cell death. Mol Cell 2002; 9: 163–173.

43. Everaerts V, Vriens J, Owsianik G, Appignoni G, Voets T, De Ridder D et al. Functional characterization of transient receptor potential channels in mouse urothelial cells. Am J Physiol Renal Physiol 2010; 299: F902–F910.

44. Oh YK, Shin KS, Kang SJ. AIF translocates to the nucleus in the spinal motor neurons in a mouse model of ALS. Neurosci Lett 2006; 406: 205–210.

45. Cox B, Emili A. Tissue subcellular fractionation and protein extraction for use in immunoblotting. Nat Protoc 2006; 1: 107–116.

46. Bancroft J, Gamble M. Theory and Practice of Histological Techniques. Churchill Livingstone: Edinburgh, 2006, p 217.

47. Maudsley S, Putcher O, Johnson CD, MacAulay L. The role of TRPM channels in oxidative stress-induced cell death. J Membr Biol 2006; 209: 31–41.

48. Wernig S, Zhao R, Moore RE, Zoghbi HY. Neurodegeneration and mitochondrial dysfunction in a mouse model of spinal muscular atrophy. Science 2002; 297: 255–260.

49. Kuzhandaivel A, Andrusova O, Yu SW, Wang H, Nistri A, Mladinic M. Extensive glial apoptosis develops in vitro and in vivo in response to extracellular magnesium in the neonatal rat spinal cord. J Physiol 2010; 589: 325–338.