Plasma membrane calcium ATPase deficiency causes neuronal pathology in the spinal cord: a potential mechanism for neurodegeneration in multiple sclerosis and spinal cord injury

Michael P. Kurnellas*,†,1, Arnaud Nicot§,1, Gary E. Shull¶, and Stella Elkabes*,†

Abstract

Dysfunction and death of spinal cord neurons are critical determinants of neurological deficits in various pathological conditions, including multiple sclerosis (MS) and spinal cord injury. Yet, the molecular mechanisms underlying neuronal/axonal damage remain undefined. Our previous studies raised the possibility that a decrease in the levels of plasma membrane calcium ATPase isoform 2 (PMCA2), a major pump extruding calcium from neurons, promotes neuronal pathology in the spinal cord during experimental autoimmune encephalomyelitis (EAE), an animal model of MS, and after spinal cord trauma. However, the causal relationship between alterations in PMCA2 levels and neuronal injury was not well established. We now report that inhibition of PMCA activity in purified spinal cord neuronal cultures delays calcium clearance, increases the number of nonphosphorylated neurofilament H (SMI-32) immunoreactive cells, and induces swelling and beading of SMI-32-positive neurites. These changes are followed by activation of caspase-3 and neuronal loss. Importantly, the number of spinal cord motor neurons is significantly decreased in PMCA2-deficient mice and the deafwaddler2J, a mouse with a functionally null mutation in the PMCA2 gene. Our findings suggest that a reduction in PMCA2 level or activity leading to delays in calcium clearance may cause neuronal damage and loss in the spinal cord.

Keywords

autoimmune disease; ATP2B2; excitotoxicity; cytoskeleton

Neuronal and axonal dysfunction and loss have increasingly received attention as important contributors to neural decline in various central nervous system (CNS) disorders, including MS and spinal cord trauma. MS is a CNS disease that may initially show a relapsing-remitting course but finally leads to permanent neurological impairment (1,2). Classically, demyelination of structurally intact axons was considered to be the main cause of clinical symptoms and neural deficits in MS. However, recent studies indicate that axonal dysfunction is already evident at the earliest clinical stages of MS, and axonal transection or loss may be the cause of persistent, irreversible disability at later phases of the disease when remissions no longer occur (3–5).
Moreover, magnetic resonance imaging studies indicate abnormal structural and biochemical changes in the gray matter of MS patients as compared with healthy individuals raising the possibility of neuronal pathology or loss (6,7). However, the molecular mechanisms underlying axonal and neuronal dysfunction remain undefined.

Our previous investigations had indicated a significant decrease in neuronal PMCA2 levels at the onset of symptoms during both myelin basic protein (MBP)-induced acute EAE in the Lewis rat and myelin oligodendrocyte glycoprotein (MOG)-induced chronic EAE in the C57Bl/6 mouse, suggesting abnormalities in calcium extrusion mechanisms in the spinal cord (8,9). Moreover, a decrease in transcript levels of PMCA isoforms in postmortem MS brains has been reported (10). However, it was not yet clear whether these changes were the cause or the consequence of neuronal/axonal damage observed in these diseases. The present studies were undertaken to determine whether suppression of PMCA activity or expression causes neuronal pathology and loss, in vitro and in vivo.

Ion dyshomeostasis, and particularly perturbations in calcium balance, have received attention as major causes of axonal damage and neuronal dysfunction in many pathological conditions (11–13). Neuronal calcium influx through voltage-gated calcium channels is followed by activation of extrusion, sequestration, and buffering mechanisms, which ensure maintenance of calcium homeostasis. Extrusion of calcium from neurons is mediated by PMCA and the Na\(^+\)/Ca\(^{2+}\) exchanger, whereas sequestration into the endoplasmic reticulum involves sarcoplasmic reticulum Ca\(^{2+}\) ATPases (SERCA). In addition, mitochondria play an important role in maintaining calcium balance within the cell. The malfunction of any of these processes can elevate intracellular calcium to abnormal levels, which, in turn, can initiate injury mechanisms, including release of mitochondrial cytochrome C which activates the caspase cascade that mediates apoptotic cell death (14). A decrease in extrusion activity and the subsequent calcium overload may lead to massive calcium influx into mitochondria, resulting in the production of reactive oxygen intermediates and perturbations in energy metabolism, which may have deleterious consequences on neurons (15).

Recent studies on MS and EAE have shown an accumulation of a pore-forming subunit of N-type voltage-gated calcium channels along demyelinated, dystrophic axons, suggesting that increased calcium influx may be a cause of axonal injury (16). However, it is not known whether and how perturbations in PMCA-mediated calcium extrusion contribute to neuronal/axonal pathology.

PMCA are P-type ATPases that play a major role in expelling calcium from cells. Four isoforms, PMCA1-4, encoded by different genes, have been described (17). The cellular and tissue distribution of these isoforms suggest that they may play distinct roles (18–21). In particular, PMCA2 and 3 are mainly localized to neurons (8,22). Moreover, the phenotype of both PMCA2-deficient mice and deafwaddler \(2J\) (dfw\(^{2J}\)), a mouse with a functionally null mutation in the PMCA2 gene, indicates that this isoform plays unique roles that are not fully compensated by the presence of other isoforms (23–25). PMCA2-deficient and dfw\(^{2J}\) mice exhibit abnormal gait and balance, as well as hearing deficits (23,25).

Our current investigations indicate that inhibition of PMCA activity, in vitro, causes neuronal pathology followed by neuronal loss. Importantly, we report a decrease in the number of spinal cord motor neurons both in PMCA2-deficient and dfw\(^{2J}\) mice as compared with their wild-type controls. Therefore, our previous findings demonstrating a decrease in neuronal PMCA2 levels at onset of symptoms in animal models of MS taken together with the results reported in the present study raise the possibility that a reduction in PMCA2 expression/activity may be a cause underlying neuronal injury in EAE and potentially MS (8,9).
MATERIALS AND METHODS

Animals

Adult dfw\(^{2J}\) and wild-type mice were purchased from Jackson Laboratories (Bar Harbor, ME). The dfw\(^{2J}\) allele has evolved spontaneously in Cby.A/J-fsn, a congenic substrain of BALB/cByJ mice. PMCA2 (−/−) and (+/+) mice are a Black Swiss strain (23). All experiments were performed according to Institutional Animal Care and Use Committee guidelines.

Spinal cord neuronal cultures

Spinal cords, dissected from 15-day-old rat embryos (Charles River, Wilmington, MA) were dissociated by trituration, layered onto a 4% BSA gradient and centrifuged at 700 × g for 2 min. The cells were resuspended in L-15 medium containing supplements (26) and plated on poly-l-ornithine coated dishes at a density of 0.8 × 10⁶ cells/35 mm petri dish. One day after plating, the media were replaced with Neurobasal medium supplemented with B-27, 6.3 mg/ml NaCl, and 10 U/ml penicillin/streptomycin. The cells were maintained in vitro for 8 days, until neuronal differentiation. The purity of the cultures (96–98%) was assessed by determining the percentage of NeuN (1:1000, Chemicon, Temecula, CA) and GFAP (1:10000, Dako, Carpinteria, CA) immunoreactive cells. Media and B-27 were purchased from Gibco (Rockville, MD). Other chemicals were obtained from Sigma (St. Louis, MO).

PMCA and SERCA inhibitors

5-(and-6)-carboxyeosin diacetate, succinimidyl ester (CE, Molecular Probes, Eugene, OR), thapsigargin and cyclopiazonic acid, *Penicillium cyclopium* (CPA, Calbiochem, San Diego, CA), were dissolved in DMSO. Sodium orthovanadate (Na\(_3\)VO\(_4\), Sigma) was dissolved according to manufacturer’s instructions.

MTT assay

Viability of neurons was assessed by use of 3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide (MTT, Sigma) assay (27). Cells in 15 fields were counted with a 40× objective using an Olympus IX50 microscope.

RT-PCR

RT-PCR was performed as described before (8). Briefly, DNase- treated total RNA (1–2 μg) was reverse-transcribed in a total volume of 20 μl using the Retroscript Reverse Transcription kit (Ambion, Austin, TX), according to manufacturer’s instructions. Five microliters of RT mix was used for PCR in a total volume of 50 μl. PCR conditions were as follows: denaturation at 94°C for 30 s, annealing at 55°C for 45 s, and polymerization at 68°C for 1 min followed by 10 min extension at 72°C. PCR was performed for 30 cycles. The regions selected for the design of the 5’ and 3’ primers were bases 667–690 and 1659–1681 of the PMCA2 gene (GeneBank accession number J03754), respectively. The predicted size of the RT-PCR product was 1014 bp. Products were separated on a 1.5% agarose gel containing ethidium bromide (0.5 μg/ml), and the image was captured using an Alpha digital imaging system.

Immunocytochemistry

Cultures were fixed using 4% paraformaldehyde/0.1 M PBS, pH 7.5. Cells were blocked with 30% horse or goat serum/0.025–0.1% triton X-100/PBS for 1 h and incubated overnight with SMI-32 (1:3000, Sternberger Monoclonals, Lutherville, MD), antiactivated caspase-3 (1:1000, Cell Signaling, Beverly, MA) or anti-PMCA2 (1:2000, Research Diagnostics, Flanders, NJ) antibodies at 4°C. This was followed by incubation with the corresponding biotinylated secondary antibody (1:100–1:200, Vector Laboratories, Burlingame, CA) and avidin-biotin complex (Vectastain, Vector). Immunopositive cells were visualized using 3,3′-
The gray matter area was quantified by use of a Nikon Microphot AF microscope with a Kodak Spot III digital camera and Ph3 Imaging (Ph3, Inc., Philadelphia, PA) software. Number of cells/mm$^2$ was calculated by dividing the total number of immunoreactive cells in each section by the gray matter area of the same section.

Quantification of intracellular calcium levels

Spinal cord cells, cultured for 8 days on poly-l-ornithine coated glass cover slips, were loaded with 4 μM Fura-2/acetoxymethyl ester (Molecular Probes) in medium for 30 min and washed with medium C for 30 min at 37°C. The medium was then replaced with 360 μl recording buffer (8 mM Na$_2$HPO$_4$, 1.5 mM KHPO$_4$, 138 mM NaCl, 2.7 mM KCl, 20 mM HEPES, 1.3 mM CaCl$_2$, 0.8 mM MgCl$_2$, 5 mM glucose). Cover slips were mounted onto a 37°C recording chamber installed on the stage of an inverted microscope (Nikon Diaphot 300, Japan). Cells were incubated with CE or vehicle (0.1% DMSO) during Fura-2 loading and subsequent steps. The concentration of CE used (5–20 μM) does not inhibit other Ca$^{2+}$ ATPases (28). For Fura-2 dual excitation, the beam of light from a high-pressure xenon bulb set was passed through a 340/380 nm shutter/filter wheel (Applied Imaging, Visitech Int., UK) at 0.2 Hz. The emitted fluorescence was collected with a 40× oil objective and detected at 510 nm. Fluorescence signals were digitized and stored on disk using Quanticell 700 (Applied Imaging). Cells were depolarized by applying 40 μl of high K$^+$ solution (100 mM KCl, 8 mM Na$_2$HPO$_4$, 1.5 mM KHPO$_4$, 38 mM NaCl, 2.7 mM KCl, 20 mM HEPES, 1.3 mM CaCl$_2$, 0.8 mM MgCl$_2$, 5 mM glucose) at $t = 46–49$ s, and responses were recorded for additional 750 s. Free intracellular calcium concentration ([Ca$^{2+}$]$_i$) was monitored by using Fura-2 ratio imaging with the parameters: $R = F_{340}/F_{380}$, $R_{min} = 0.3$, $R_{max} = 3$, $\beta = 3.42$ and $K_D = 220$ nM (29). Differences in calcium levels before KCl (basal) at KCl-induced peak, 350 s after KCl ($t = 400$ s) and at endpoint ($t = 800$ s) were analyzed using ANOVA for repeated measures with one between subject factor (CE treatment) and one within subject factor (time) followed by Tukey’s post hoc test. The clearance rates (t1/2 and t1/3) were obtained from each trace. When the clearance rate was superior to recording time (800–830 s), it was cut off at 800 s for statistical analysis.
Results

Blockade of PMCA activity induces neuronal pathology and death

To investigate the effects of calcium pump inhibitors on neuronal function, we employed purified spinal cord neuronal cultures, which were maintained in vitro for eight days. We first ascertained that PMCA2 mRNA and protein were expressed in these cultures. RT-PCR amplified a band of the expected size (8, Fig. 1A). The identity of this band was further confirmed by sequence analysis. PMCA2 protein was also detected in cell bodies and processes by immunocytochemistry using an antibody specific for this pump (Fig. 1B).

Initially, we exposed cells to different concentrations of Na$_3$VO$_4$, a general inhibitor of P-type ATPases, including PMCAs and SERCA pumps (30–32). Subsequently, we assessed immunoreactivity to nonphosphorylated neurofilament H by use of the SMI-32 antibody, a marker of neuronal/axonal damage in MS, EAE, and other pathological conditions of the CNS (33–35). Thirty μM Na$_3$VO$_4$, an optimal dose, significantly increased the number of SMI-32 positive cells within 4 h (Fig. 1C). A lower dose (10 μM) was also effective when cells were exposed to the inhibitor for 8 h. Since Na$_3$VO$_4$ can also interfere with the activity of SERCA pumps, we investigated whether cyclopiazonic acid (10 μM) and thapsigargin (250 nM), specific SERCA pump inhibitors, induce a similar response (36,37). The doses employed were maximally effective as reported previously (36) and as indicated by our preliminary experiments. We did not observe any significant changes in the SMI-32 positive cell number (Fig. 1D). These results suggested that Na$_3$VO$_4$-induced neuronal pathology might primarily be due to inhibition of PMCAs.

To further ascertain that the changes in SMI-32 immunopositive neuron number were indeed due to blockade of PMCA activity, we employed 5-(and-6)-carboxy-yo (CE), a more selective and potent PMCA inhibitor (38–40). Cultures were treated with medium/vehicle or 5 μM CE (optimal dose, range tested 0–20 μM) for 2, 4, and 8 h. Exposure to CE for 4 h induced a 1.8-fold increase in SMI-32 positive cells (Fig. 1E). The diverse morphology of these cells suggested that different neuronal subpopulations were affected. Surprisingly, SMI-32 positive cell number in CE-treated cultures was not significantly different from controls at 8 h. One potential interpretation of these findings was that exposure of cultures to CE for 4 h increases immunoreactivity to nonphosphorylated neurofilament H, which is then followed by loss of SMI-32 positive affected cells at 8 h. To test this possibility, we investigated cell survival by use of MTT assay. Indeed, a reduction in MTT-positive cells was observed by 8 h (Fig. 1F). Thus, prolonged exposure to CE affects neuronal survival.

Inhibition of PMCA activity and the resulting aberrance in calcium extrusion may lead to abnormal increases in intracellular calcium levels ([Ca$^{2+}$]i), which promote mitochondrial cytochrome C release, a trigger that activates the caspase cascade mediating apoptotic cell death (41). To determine whether blockade of PMCA activity induces apoptotic mechanisms, we quantified the number of cells expressing activated caspase-3, a key component of apoptosis. There was a twofold increase in the number of caspase-3 positive cells in cultures treated with CE for 8 h suggesting that cell death may partially be due to apoptosis (Fig. 1G). Thus, our results indicate that CE-induced neuronal damage, detected initially by changes in neurite morphology and increased SMI-32 immunoreactivity, is followed by activation of caspase-3 and neuronal loss.

It is worth noting that although extended exposure to Na$_3$VO$_4$ increased SMI-32 positive cell number, the inhibitor did not induce neuronal death within the period employed in the present
Inhibition of PMCA activity induces morphological abnormalities in neuritic processes

In addition to the increase in SMI-32-positive cell number described above, treatment of spinal cord neuronal cultures with CE for 4 h also affected the morphology of neurites (Fig. 2). In control cultures, maintained only in medium and vehicle, a small subpopulation of cells constituting ~2–4% of the total cells in the dish, was immunopositive for SMI-32 (Fig. 2A). These cells exhibited motor neuron-like morphology and had long, smooth and SMI-32-positive processes. Indeed, previous studies reported SMI-32 immunoreactivity in cultured motor neurons, even in the absence of a pathological challenge (42). Addition of 5 μM CE to sister cultures initiated swelling and beading of neurites within 2 h (data not shown), which became further pronounced after 4 h (Fig. 2B and C). This morphology contrasted with the smooth appearance of the majority of SMI-32 positive processes in control cultures (Fig. 2D). Whereas in controls only 10.2 ± 2.63% of all SMI-32-positive cells showed occasional swellings in their processes, in CE treated cultures, 42.4 ± 2.39% of SMI-32-positive cells had neurites with frequent beading and swellings (P < 0.0001 by Student’s t test). In addition, spherical structures, reminiscent of axonal retraction bulbs, could be observed at the end of many processes (Fig. 2E). Disintegration of some neurites occurred at 8 h and paralleled cell death (not shown).

Inhibition of PMCA activity delays recovery from depolarization-evoked increases in intracellular calcium

Because PMCA play a major role in calcium extrusion, we hypothesized that a decrease in PMCA activity may lead to alterations in [Ca^{2+}]i, which, in turn, induces injury cascades. To determine the effects of CE on intracellular calcium balance in spinal cord neurons, cultures were incubated with either vehicle or 5 and 20 μM CE for only 1 h, a time that precedes the appearance of morphological changes in neurites. This treatment elevated the resting cytosolic calcium levels by 27 nM and 81 nM when 5 μM CE and 20 μM CE were used, respectively (Fig. 3A and B). Moreover, the clearance of depolarization-induced Ca^{2+} transients was significantly slower in CE-treated cells than controls (Fig. 3). The rate of Ca^{2+} clearance was estimated by analyzing the time for half (t1/2) and one third (t1/3) decay of the [Ca^{2+}]i transients induced by high K+ buffer. Calcium clearance was at least fivefold slower in CE-treated groups as compared with controls. Strikingly, 9% of the cells exposed to 5 μM CE and 55% of the cells treated with 20 μM CE exhibited a very slow calcium clearance rate (t1/2 > 800 s), while the remaining cells had t1/2 values of 294±15 and 196±17 s, respectively. This suggests that one or more neuronal subpopulations may be extremely dependent on PMCA activity for...
calcium removal. The amplitudes of depolarization-induced $[\text{Ca}^{2+}]_i$ peaks were also greatly increased in the presence of CE, in a dose-dependent manner.

These results indicate that PMCA activity is critical for the maintenance of resting $[\text{Ca}^{2+}]_i$ and for recovery from $\text{K}^+$ induced $[\text{Ca}^{2+}]_i$ rise in spinal cord neurons. Moreover, our findings suggest that approximately half of the cells in our cultures are highly dependent on PMCA activity for recovery from depolarization-evoked calcium transients.

**Loss of motor neurons in the spinal cord of PMCA2 null mice and $dfw^{2J}$ mutant mice**

To determine whether the lack of PMCA2 causes neuronal loss, in vivo, we quantified the number of SMI-32 positive cells in spinal cord sections of PMCA2 (+/+) and (−/−) mice. In particular, we focused on motor neurons, because they are highly vulnerable to changes in intracellular calcium levels and $\text{Ca}^{2+}$-mediated injury (42). Staining for SMI-32, a marker for motor neurons (42), was confined to somata of anterior horn cells exhibiting the characteristic, motor neuron-like morphology both in wild-type and knockout mice. There was a 37.5% decrease in the number of SMI-32-positive cells in PMCA2-deficient mice as compared with their wild-type littermates (Fig. 4A). We also analyzed the results by calculating the SMI-32 positive cell number/mm$^2$ gray matter (Fig. 4B), as the total gray matter area is 13% smaller in PMCA2 (−/−) as compared with PMCA2 (+/+ ) mice (Fig. 4C). There was a 26.5% decrease in SMI-32 immunoreactive cells/mm$^2$ gray matter in PMCA2 knockout mice as compared with wild-type controls.

To corroborate these findings, we labeled adjacent sections with an antibody against peripherin, another marker for motor neurons (42). Peripherin immunoreactivity was strongest in anterior horn cells with motor neuron-like morphology but could also be observed, to a lesser extent and intensity, in some smaller cells of the gray matter. In agreement with the results obtained by use of SMI-32, we found a significant decrease in the number of peripherin immunoreactive cells and peripherin-positive cells/mm$^2$ gray matter (37.4 and 27.7%, respectively) in PMCA2 (−/−) as compared with PMCA2 (+/+ ) mice (Fig. 4D–4G).

To further ensure that the lack of PMCA2 activity decreases the number of motor neurons, we quantified SMI-32 positive cells in the spinal cord of $dfw^{2J}$ mice. A two base-pair deletion in the PMCA2 gene of $dfw^{2J}$ causes a frameshift leading to a truncated, inactive protein (25). In agreement with the studies on PMCA2 (−/−) mice, we found a 41% decrease in the number of SMI-32-positive cells in $dfw^{2J}$ mice as compared with their wild-type controls (Fig. 5A). We also found a 39% decrease in SMI-32-positive cell number/mm$^2$ gray matter (Fig. 5B) when the small but significant 12% decrease in gray matter area of the $dfw^{2J}$ spinal cord was taken into consideration (Fig. 5C).

These results, taken together, indicate that the lack of PMCA2 expression in PMCA2 (−/−) mice or a functionally null mutation in the PMCA2 gene in $dfw^{2J}$, have similar effects resulting in a decrease in the number of SMI-32 positive motor neuron-like cells in the spinal cord. It is worth noting that our results do not exclude the possibility that additional neuronal subpopulations may also be affected in the spinal cord of PMCA2-deficient and mutant mice.

**DISCUSSION**

Axonal damage and neuronal loss occur in many CNS disorders, including MS and spinal cord injury. Yet, the triggers that initiate axonal pathology, the underlying mechanisms, and the critical molecules, which may be targets for therapeutic interventions, are not well defined. Our present findings indicate that inhibition of PMCA activity in neuronal cultures, delays calcium clearance, induces cytoskeletal abnormalities, promotes swelling of neurites and finally causes neuronal death. Reductions in PMCA levels and/or activity may first induce
intracellular calcium overload due to delayed clearance resulting from abnormalities in extrusion mechanisms. Subsequently, elevated calcium can cause changes in cytoskeletal proteins as suggested by increased nonphosphorylated neurofilament H (SMI-32)-positive cell number and alterations in SMI-32 immunoreactive neurite morphology. The changes in neurite morphology appear to be similar to those observed after addition of the calcium ionophore A23187 to spinal cord neuronal cultures, which promotes beading, disruption of the cytoskeleton, and mitochondrial abnormalities (43). As changes in cytoskeletal proteins may interfere with axonal transport, swellings may be the consequence of the accumulation of organelles, cytoplasm, and fragmented neurofilaments in affected processes. Furthermore, PMCA are localized to both synaptic terminals and dendrites (21). Thus, a reduction in PMCA activity may damage not only axons but also dendrites, which are affected during EAE (44). Indeed, it has been suggested that dendritic arborization of Purkinje cells in PMCA2 null mice is reduced as compared with their wild-type littermates (23). Such changes may alter synaptic function and neuronal communication as well as survival. In fact, the early pathological changes observed in our neuronal cultures are followed by activation of death pathways, including apoptotic mechanisms, resulting in neuronal loss as indicated by an increase in activated caspase-3 positive cells and a reduction in MTT positive cells. The decrease in SMI-32 and peripherin immunoreactive neuron number in PMCA2-deficient mice and the reduction in SMI-32 positive cell number in dfw2J mice as compared with their respective wild-type controls further support this notion. Thus, a perturbation in PMCA-mediated calcium extrusion may be an important contributor to axonal and neuronal damage in pathological conditions of the CNS, including EAE, MS, and spinal cord injury. Importantly, our previous results indicate a decline in the levels of PMCA2 in two different animal models of MS with distinct disease course and histopathological characteristics (8,9). This finding is of particular significance as axonal pathology and neuronal loss has been reported in both models and suggests similarities in molecular mechanisms underlying neuronal damage during acute and chronic EAE (34,35).

It is worth noting that the findings of the present study may also be relevant to other pathological conditions of the CNS, including spinal cord trauma, as we and others have reported a decrease in PMCA2 transcript levels after spinal cord contusion injury (45,46). It is possible that anomalies in PMCA-mediated calcium extrusion contribute to secondary neuronal/axonal damage after spinal cord injury, expanding the deleterious effects of the first traumatic impact. Calcium has also been implicated in the degeneration of axons in the white matter after anoxia/ischemia (47). It has been hypothesized that the energy depletion following hypoxia or ischemia results in failure of Na+/K+ ATPase and activation of sodium channels, which augments axoplasmic Na+ concentrations. This, in turn, causes the reversal of Na+/Ca2+ exchanger (NCX) activity, thus increasing calcium levels (48). Moreover, calcium influx through sodium channels further augments axoplasmic calcium concentrations (12). As a consequence, protease-mediated injury cascades are activated (49). In agreement with these results, blockade of sodium channels and NCX prevents axonal deterioration in anoxia (50,51). Interestingly, similar mechanisms have recently been implicated in MS and EAE (52,53). In addition, recent studies have suggested that ryanodine receptor-mediated release of calcium from the endoplasmic reticulum causes abnormal elevations in calcium concentrations within the axoplasm promoting injury during spinal cord ischemia (54).

In summary, our findings and those reported by others indicate that axonal and neuronal pathology in CNS disorders and trauma may be governed by multiple mechanisms. Modifications in the function of calcium channels, NCX, and PMCA may all contribute to the disruption of calcium balance, a key element playing a pivotal role in cellular pathology and death. These findings also raise the possibility that independent of the initial trigger inducing neuronal/axonal damage, the molecular mechanisms underlying neuronal injury in different pathological conditions of the CNS may share commonalities.
Acknowledgments

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References


Figure 1. Effects of PMCA and SERCA inhibitors on spinal cord neurons, in vitro
A) RT-PCR showing expression of PMCA2 in neuronal cultures. Lane 1: molecular weight marker; bp, base pairs; lane 2: RT-PCR product (arrow) at the predicted molecular weight (1014 bp). The identity of the band was further verified by sequence analysis. B) PMCA2 immunoreactivity in spinal cord neurons. Arrows and arrowheads point at immunopositive cells and processes, respectively. C) SMI-32 positive cell number after exposure of cultures to 10, 30, and 100 μM Na3VO4. C: control. D) Effects of SERCA inhibitors on SMI-32 positive cell number. CPA: cyclopiazonic acid. E) SMI-32 positive cells after exposure of cultures to 5 μM 5-(and-6)-carboxy eosin diacetate, succinimidyl ester (CE). F) Effects of CE on cell survival. G) Induction of activated caspase-3 in cells treated with CE. The experiments were repeated at least twice and yielded similar results. Values are presented as means ± SEM. Significantly different from controls *P < 0.003, **P < 0.0001 by ANOVA; +, significantly different from control by Student’s t test, P < 0.02; n = 6–12.
Figure 2. Effects of CE on the morphology of SMI-32 immunoreactive neurites in spinal cord neuronal cultures

A) Composite picture showing two fields of a control culture maintained in medium/vehicle. A subpopulation of cells with motor neuron-like morphology (big arrows) and their processes are SMI-32 positive as reported previously (42). B) Composite picture showing two fields of a culture treated with 5 μM CE for 4 h. Addition of CE, induced beading (small arrows) and swelling (arrowhead) of neurites. C) High magnification picture of a neurite in a CE-treated culture showing beadings (small arrows) and swellings (arrowheads). D) High magnification picture of a neurite in a control culture showing a smooth process. E) An SMI-32 positive cell in a CE-treated culture showing a spherical structure resembling a retraction bulb (concave arrow). The experiment was repeated twice and yielded similar results (n=8). Bar = 100 μm.
Figure 3. Effect of CE on basal and depolarization-evoked [Ca\textsuperscript{2+}]_i levels in spinal cord neuronal cultures

A) [Ca\textsuperscript{2+}]_i changes in cells loaded with Fura-2. High K\textsuperscript{+} buffer was added at \( t = 46\text{--}49 \) s (arrow). Preincubation with CE for 1 h increased resting [Ca\textsuperscript{2+}]_i and prolonged the recovery from depolarization-evoked Ca\textsuperscript{2+} transients. The amplitudes of depolarization-induced [Ca\textsuperscript{2+}]_i peaks were also greatly increased by CE treatment. Values are presented as means ± SEM.

B) Table summarizing calcium levels and clearance rates before KCl (basal), at KCl-induced peak, at 400 s (350 s after KCL addition) and at end point (\( t = 800 \) s) in the experiments (A). 

\( t_{1/2} \) and \( t_{1/3} \) are half and one-third the time required to clear calcium transients, respectively.

ANOVA for calcium levels: dose effect, \( F_{(2,202)} = 93.3, P < 0.0001 \); time effect, \( F_{(3,606)} = 897, P < 0.0001 \); dose X time interaction, \( F_{(6,606)} = 26.4, P < 0.0001 \); post hoc analysis: *\( P < 0.05 \); **\( P < 0.01 \) CE vs. vehicle. ANOVA for \( t_{1/3} \): \( F_{(2,204)} = 41.4, P < 0.0001 \); post hoc analysis: **\( P < 0.01 \) CE vs. vehicle. ANOVA for \( t_{1/2} \): \( F_{(2,204)} = 85.0, P < 0.0001 \); post hoc analysis: **\( P < 0.01 \) CE vs. vehicle.
Figure 4. Loss of SMI-32 and peripherin positive neurons in the spinal cord of PMCA2 knockout (KO) mice

The number of SMI-32 positive cells/spinal cord section is decreased in KO as compared with wild-type (WT) mice (A). SMI-32 positive cells/mm$^2$ gray matter is also lower in KO mice (B) when the reduction in gray matter area is taken into consideration (C). Peripherin-positive cells/spinal cord section (D) and peripherin-positive cells/mm$^2$ gray matter (E) are significantly reduced in KO as compared with WT mice. Peripherin-immunoreactive motor neurons in the anterior horn of WT (F; arrows) and KO mouse (G). Four WT and KO mice were used and seven sections collected from each lumbar spinal cord at 100 μm intervals, were analyzed. Values are presented as means ± SEM. Significantly different from WT *$P < 0.03$, **$P < 0.01$, ***$P < 0.0002$ by Student’s $t$ test. Bar = 200 μm.
Figure 5. Loss of SMI-32 positive cells in the spinal cord of dfw^{2J} mice (DFW)
SMI-32 positive cells are decreased in the spinal cord of dfw^{2J} as compared with its wild-type (WT) control (A). SMI-32-positive cells/mm² gray matter is also lower in dfw^{2J} (B) when the reduction in gray matter area is taken into consideration (C). Three dfw^{2J} and WT mice were used. Ten sections obtained from each lumbar spinal cord at 100 μm intervals were analyzed. Values are presented as means ± SEM. Significantly different from WT ***P < 0.0001 and **P < 0.002 by Student’s t test.