TRPM4 activation by chemically- and oxygen deprivation-induced ischemia and reperfusion triggers neuronal death

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ABSTRACT

Cerebral ischemia-reperfusion injury triggers a deleterious process ending in neuronal death. This process has two components, a glutamate-dependent and a glutamate-independent mechanism. In the glutamate-independent mechanism, neurons undergo a slow depolarization eventually leading to neuronal death. However, little is known about the molecules that take part in this process. Here we show by using mice cortical neurons in culture and ischemia-reperfusion protocols that TRPM4 is fundamental for the glutamate-independent neuronal damage. Thus, by blocking excitotoxicity, we reveal a slow activating, glibenclamide- and 9-phenanthrol-sensitive current, which is activated within 5 min upon ischemia-reperfusion onset. TRPM4 shRNA-based silenced neurons show a reduced ischemia-reperfusion induced current and depolarization. Neurons were protected from neuronal death up to 3 hours after the ischemia-reperfusion challenge. The activation of TRPM4 during ischemia-reperfusion injury involves the increase in both, intracellular calcium and H2O2, which may act together to produce a sustained activation of the channel.

KEYWORDS

glutamate-independent neuronal death; ischemia-reperfusion; oxidative stress 9-phenanthrol; TRPM4

Introduction

Neuronal ischemia activates a mechanism of damage called excitotoxicity which consists in an uncontrolled glutamate release from synaptic terminals activating glutamatergic receptors and thus, initiating a persistent depolarization and tissue damage. 1,2 This mechanism is self-limited and finishes after synaptic vesicles depletion. 3 Despite the self-limited feature of excitotoxicity, it has been noted that neurons remain depolarized in a not well understood mechanism in which participates, among others, a Na+ inward current (NSCa-ATP) that is activated by high [Ca2+]i and low [ATP]i and modulated by ROS. 4-7 This mechanism is believed to be critical for the delayed neuronal death that occurs hours or even days after the initial injury. 9

The interplay between high Ca2+ and high production of ROS are side effects of ischemia-reperfusion injury. Conversely, Ca2+ contributes to altered protein function such as increased calpain activity, 10 increased ion permeability, 11 activation of Ca2+-dependent phospholipases, and activation of ROS-generating pathways like NADPH oxidases (NOX) 12 and xanthine oxidase conversion. 13 Oxidants contribute to protein impairment, DNA breakdown, mitochondrial damage, increased membrane permeability and neuronal depolarization, 1,14 altogether, perpetuating the damage and accounting for neuronal death.

TRPM4 is a non-selective cation channel permeable to monovalent cations and activated by intracellular Ca2+, 15 blocked by ATP 16 and modulated by ROS. 8 Physiologically, TRPM4 is involved in the control of Ca2+ oscillations through the modulation of resting membrane potential, 17,18 cell migration, 19,20 rhythm generation in preBötzing neurons, 21 the afterdepolarization in cerebellar neurons, 22 among other functions (for a review, see 23). Interestingly, increased Ca2+ and ROS production during reperfusion favor the activation of TRPM4. 16,24,25 Moreover, this channel has been previously involved in glutamate-dependent neuronal death; ischemia-reperfusion; oxidative stress 9-phenanthrol; TRPM4.
during reperfusion is not well characterized. In this work by using chemical as well as oxygen and glucose deprivation ischemia-reperfusion models, we show that the continuous activation of TRPM4 during reperfusion leads neurons to a state of sustained depolarization leading to their death. The pharmacological inhibition or shRNA-based silencing of TRPM4 renders neurons resistant to reperfusion damage, increasing their survival rate. Thus, blockade of the sustained activation of TRPM4 might constitute a target for novel pharmacological tools addressed to reduce the damage during brain reperfusion.

Results

Persistent neuronal depolarization during ischemia-reperfusion

To figure out the potential role of TRPM4 in the glutamate-independent neuronal damage, we measured first the membrane potential of mouse cortical pyramidal neurons from primary cultures using the nystatin-perforated patch clamp technique. Neurons were exposed to 5 mM NaN3 in the absence (Fig. 1a) or presence (Fig. 1b) of an AET buffer to block excitotoxicity. We found that neurons depolarize after NaN3 exposure despite of AET presence, reaching a steady-state at 5 min (0 min = −69.5 ± 8.9 mV, 15 min = −9.2 ± 11.2 mV, n = 6, t-test, p < 0.0002). In the absence of AET, neurons undergo a continuous firing and depolarization (0 min = −72.2 ± 6.5 mV, 15 min = −8.8 ± 8.2 mV, n = 6, t-test, p < 0.0001). In voltage-clamp experiments ($V_h = −70$ mV) we observed the development of a slow current that reached steady-state after 15 min (−359 ± 182.2 pA) upon NaN3 exposure in the absence (Fig. 1c) or presence (Fig. 1d) of the AET buffer. The current development correlated with the depolarization observed under current-clamp conditions (3.4 ± 1.2 min vs. 3.1 ± 1.6 min, respectively). During voltage-clamp experiments in the absence of the AET buffer, we observed an increased excitatory postsynaptic current (EPSC) frequency followed by a slow inward current ($V_h = −70$ mV, 0 min = −41.2 ± 19.8 pA, 15 min = −331.2 ± 143.8 pA, n = 6, t-test, p < 0.005). The firing rate and EPSC frequency after 10 min of NaN3 perfusion were completely suppressed at 20 min. These data suggest that excitotoxicity is a self-limited event and other conductances could be underlying the NaN3-induced depolarization.

Figure 1. Excitotoxicity-independent neuronal depolarization after chemical ischemia-reperfusion. (a) Left panel shows a representative membrane potential recording in cortical pyramidal neurons (DIV 14–21) treated with 5 mM NaN3. Right panel depicts the membrane potential values at 0 and 15 min post NaN3 for each experiment (n = 6). (b) Left panel shows a representative membrane potential recording in cortical pyramidal neurons (DIV 14–21) treated with 5 mM NaN3 and AET. Right panel depicts the membrane potential values at 0 and 15 min post NaN3 and AET for each experiment (n = 6). (c) Left panel shows a representative current recording showing the effect of 5 mM NaN3. Right panel depicts the current values at 0 and 15 min post NaN3 for each experiment (n = 6). (d) Left panel shows a representative current recording showing the effect of 5 mM NaN3 and AET. Right panel depicts the current values at 0 and 15 min post NaN3 and AET for each experiment (n = 6). Black arrows indicate t = 0 and t = 15, respectively.
**TRPM4 expression in cortical pyramidal neurons**

To explore whether TRPM4 is expressed in cortical pyramidal neurons, we performed imaging studies using 5 weeks-old C57BL/6 mouse brain slices. First, we evaluated the efficacy of the TRPM4-shRNA. To that end, cortical pyramidal neuron cultures were transfected with a shRNA against TRPM4 (TRPM4-shRNA) expressing EGFP as an expression marker. As depicted in Fig. 2a-d, neurons expressing the TRPM4-shRNA have a ~55% reduction in TRPM4 expression. Next, we observed the expression of TRPM4 in slice of medial prefrontal cortex. As shown, we observed TRPM4 labeling in several neurons across the cortical region (Fig. 2e, h). A zoomed area showed that most of the labeling is somatic with few neurons presenting labeling in neurites (Fig. 2h). MAP2 labeling showed the expected somatodendritic pattern (Fig. 2f, i). Superposition of images showed several overlapping sites (Fig. 2g, j) suggesting the presence of TRPM4 in the neurons.

**TRPM4 involvement in the reperfusion current**

Previous studies indicate the involvement of TRPM4 in glutamate-induced neuronal death and axonal degeneration,²⁶,²⁷ however, its role in glutamate-independent neuronal death is still unclear. To assess whether TRPM4 participates in the depolarization observed in this process, we used an shRNA-based

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**Figure 2.** TRPM4 expression in cortical pyramidal neurons. (a) illustrates a representative experiment showing two neurons transfected with the TRPM4-shRNA positive for EGFP expression. (b) shows the immunofluorescence for TRPM4. (c) shows superposition of images. (d) depicts the percentage of TRPM4 reduction of expression (55.3 ± 5.5%; n = 4 independent experiments, 55 cells in total). Representative images of the cerebral cortex showing the somatic immunoreactivity for TRPM4 (e), MAP2 (f) and the superposed signal (g) at 40x magnification and (h), (i), (j) at 63x magnification. Arrows indicate somata and neurite labeling for TRPM4.
silencing strategy. In these experiments, we found that cortical neurons transduced with TRPM4_{shRNA} exhibit a small depolarization after NaN3 + AET exposure (0 min = $-76.4 \pm 5.9$ mV, 15 min = $-55.8 \pm 21$ mV, n = 5, t-test, p = 0.0638, Fig. 3a), as compared to TRPM4_{scrambled}-transduced neurons (0 min = $-72.6 \pm 3.14$ mV, 15 min = $-18.6 \pm 5.6$ mV, n = 5, Fig. 3b). In voltage-clamp experiments, TRPM4_{shRNA}-transduced neurons expressed a small inward current ($V_h = -70$ mV, 0 min = $-44.5 \pm 26.9$ pA, 15 min = $-190.2 \pm 52.6$ pA, n = 5, t-test, p < 0.0017, Fig. 3c), as compared to TRPM4_{scrambled}-transduced neurons (0 min = $-76.7 \pm 11$ pA, 15 min = $-497.5 \pm 40.8$ pA, n = 6, Fig. 3d).

**Figure 3.** TRPM4_{shRNA} and pharmacological blockade protects neurons from ischemia and reperfusion-induced depolarization. (a) and (b) Representative membrane potential recordings in cortical pyramidal neurons (DIV 14–21) treated with 5 mM NaN3 and AET expressing TRPM4_{shRNA} or TRPM4_{scramble}. Right panels depict the membrane potential values at 0 and 15 min post NaN3 and AET for each experiment (n = 5). (c) and (d) Representative current recordings showing the effect of 5 mM NaN3, and AET expressing TRPM4_{shRNA} (n = 5) or TRPM4_{scramble}. Right panels depict the current values at 0 and 15 min post NaN3 and AET for each experiment (n = 6). Black arrows indicate t = 0 and t = 15, respectively. (e) Representative current recordings in cortical pyramidal neurons treated with NaN3 and AET (blue trace), NMDG$^+$ replacement (red trace) and NMDG$^+$ in the presence of 10 $\mu$M 9 Ph (green trace). (f) Representative current recordings of neurons expressing TRPM4_{shRNA} (black trace) or TRPM4_{scramble} (blue trace) exposed to NaN3 and AET. (g) Representative current recordings of neurons exposed to NaN3 + AET; control trace (black trace), TRPM4_{shRNA} (grey trace) and TRPM4_{shRNA} + 9 Ph (red trace).
To explore the ionic nature of this current, we performed experiments replacing extracellular Na\(^+\) with the non-permeable cation NMDG\(^+\) after 10 min of perfusion with 5 mM NaN\(_3\) + AET and measured the current using a voltage ramp protocol (−80 to 80 mV, dV/dt 0.4 V/s). We found that replacing Na\(^+\) with NMDG\(^+\) decreases the inward current in 92 ± 10.3% (Fig. 3e, red trace) compared to control (blue trace), showing that Na\(^+\) is the main inward cation carried by this current. The exposure of neurons to 10 μM 9-phenantrol (9Ph, a non-specific TRPM4 inhibitor) in the presence of NaN\(_3\) + AET and NMDG\(^+\) as the main cation reduces the outward current (Fig. 3e, green trace), as expected for a cation-selective current carried by TRPM4. To confirm whether this current is driven by TRPM4, we performed experiments in neurons transduced with TRPM4<sub>shRNA</sub> (black trace in Fig. 3f) and TRPM4<sub>scramble</sub> (blue trace in Fig. 3f). We found that in neurons subjected to TRPM4 silencing the current induced NaN\(_3\) + AET exposure was significantly reduced. To explore whether TRPM4 silencing and 9 Ph present an additive effect, we performed experiments in neurons transduced with TRPM4<sub>shRNA</sub> (grey trace, Fig. 3g) and in neurons transduced with TRPM4<sub>shRNA</sub> + 9 Ph (red trace, Fig. 3g), both compared to control (black trace, Fig. 3g). Fig. 3h shows a summary graph of the current recorded under these conditions. As seen, there is no additive effect.

### Increased H\(_2\)O\(_2\) and Ca\(^{2+}\) levels during ischemia-reperfusion

In a previous work, we demonstrated that TRPM4 desensitization is removed by H\(_2\)O\(_2\), leading to cell death.\(^8\) Because an important feature of the ischemia-reperfusion injury is ROS generation, we monitored chemically-induced ischemia-reperfusion H\(_2\)O\(_2\) increase. Using the genetically encoded H\(_2\)O\(_2\) probe HyPer-Cyto,\(^28\) we found an increase of H\(_2\)O\(_2\) levels at 3 min reaching steady-state at 7 min post NaN\(_3\) (Fig. 4a, black circles). Preincubation for 1 h with 1 mM N-acetyl cysteine (NAC) abolished the H\(_2\)O\(_2\) increase induced by NaN\(_3\) + AET (Fig. 4a, blue circles). As a control, to rule out an effect of the TRPM4 inhibitor 9 Ph on the H\(_2\)O\(_2\) increase neurons were exposed to 10 μM 9 Ph (Fig. 4a, red circles).

Because Ca\(^{2+}\) increase is one of the landmarks of ischemia-reperfusion injury and is crucial for TRPM4 activation, we explored whether NAC affected [Ca\(^{2+}\)]\(_i\) during ischemia-reperfusion. As expected, NaN\(_3\) + AET increases Ca\(^{2+}\) (Fig. 4b). This increase was partially abolished by 5 min preincubation with 1 mM NAC (Fig. 4b, blue circles). TRPM4 inhibition by 10 μM 9 Ph did not affect Ca\(^{2+}\) levels (Fig. 4b, red circles) showing that, at least in the experimental protocol used for chemical ischemia-reperfusion, Ca\(^{2+}\) increase is partially dependent on ROS increase.

![Figure 4](image-url)  
**Figure 4.** Reperfusion increases ROS and [Ca\(^{2+}\)]\(_i\) in cortical pyramidal neurons. (a) H\(_2\)O\(_2\) quantification using HyPer-Cyto in cortical pyramidal neurons treated with NaN\(_3\) + AET (black circles, n = 7), NaN\(_3\) + AET + 9 Ph (red circles, n = 7) and NaN\(_3\) + AET + 1 mM NAC (blue circles, n = 7). (b) Intracellular Ca\(^{2+}\) measurements in cortical pyramidal neurons treated with 5 mM NaN\(_3\) + AET (n = 8). (c) Effect on intracellular Ca\(^{2+}\) of NaN\(_3\) + AET + 10 μM 9 Ph (red circles, n = 8) and NaN\(_3\) + AET + 1 mM NAC (blue circles, n = 8). Data are expressed as mean ± SEM.
**TRPM4 activation depends on ROS and a Ca^{2+} influx**

One prediction from the above described results is that suppression of Ca^{2+} influx and/or ROS levels should hamper the depolarization and TRPM4 current activation induced by ischemia-reperfusion injury. We tested this prediction first by performing chemically induced ischemia-reperfusion experiments in the absence of extracellular Ca^{2+}. We found that extracellular Ca^{2+} exclusion protected neurons from depolarization (0 min = −71.2 ± 9.9 mV, 15 min = −68 ± 9.6 mV, n = 5 t-test, p = 0.5, Fig. 5a), and current activation (0 min = −24.6 ± 12.1 pA, 15 min = −56.5 ± 24.85 pA, n = 5, t-test, p = 0.96, Fig. 5c). Then, we evaluated the effect of NAC, a small depolarization was recorded (0 min = −75 ± 5.7 mV, 15 min = −48.4 ± 8.5 mV, n = 5, t-test, p = 0.0117, Fig. 5b), which is related to a small current activation (0 min = −40.8 ± 26.6 pA, 15 min = −83.3 ± 22 pA, n = 6, t-test, p = 0.1205, Fig. 5d).

To verify our observations with a different protocol of ischemia-reperfusion injury we performed an oxygen-glucose deprivation-reperfusion protocol (OGD). As depicted in Fig. 5e, during 30 min of exposure to ischemia-reperfusion injury, we observed a decrease in membrane potential (0 min = −48.4 ± 8.5 mV, 15 min = −32.1 ± 5.7 mV, n = 5, t-test, p = 0.002, Fig. 5f).

**Figure 5.** Ca^{2+} and reducing agents modulate ischemia and reperfusion-induced depolarization. (a) Representative membrane potential recording in cortical pyramidal neurons (DIV 14–21) treated with 5 mM NaN₃ + AET in the absence of extracellular Ca^{2+}. Right panel shows the membrane potential values at 0 and 15 min post NaN₃ + AET for each experiment (n = 5). (b) Representative membrane potential recording in cortical pyramidal neurons (DIV 14–21) treated with 5 mM NaN₃ + AET and 1 mM NAC. Right panel shows the membrane potential values at 0 and 15 min post NaN₃ + AET and 1 mM NAC for each experiment (n = 5). (c) Representative current recording showing the effect of 5 mM NaN₃ + AET in the absence of extracellular Ca^{2+}. Right panel shows the current values at 0 and 15 min post NaN₃ + AET for each experiment (n = 5). (d) Representative current trace showing the effect of 5 mM NaN₃ + AET and 1 mM NAC. Right panel shows the current values at 0 and 15 min post NaN₃ + AET and 1 mM NAC for each experiment (n = 6). Black arrows indicate t = 0 and t = 15, respectively. (e) Current time course measured at 100 mV from a voltage ramp as described in Methods in cortical pyramidal neurons exposed to 30 min OGD in the presence of AET (OGD + AET) and during reperfusion (Reperfusion + AET). (f) Mean current values during Reperfusion + AET and upon exposure to 100 μM glibenclamide (Glib + AET, n = 5).
an OGD protocol we observed no current activation, however, 10 min after switching to normal ACSF we observed a current increase which was inhibited by 100 μM glibenclamide, a known TRPM4 inhibitor (Fig. 5f).

**TRPM4 inhibition increases neuronal survival**

We sought to explore whether TRPM4 inhibition increases neuronal survival after ischemia-reperfusion injury. As depicted in Fig. 6a neuronal death measured with the trypan-blue exclusion method was reduced in neurons exposed to NaN₃ + AET plus 9 Ph and in neurons transduced with TRPM4_shRNA (63 ± 12%, n = 6, one-way ANOVA, p < 0.0055).

To follow the temporal course of cell death, cortical pyramidal neurons transduced with TRPM4_shRNA or TRPM4_scramble were exposed to NaN₃ + AET or 9 Ph, and then tested for cell death with the LDH release method (Fig. 6b). The data show that neurons begin to die after 15 min. TRPM4 silencing or TRPM4 inhibition by 9 Ph decreases cell death up to three hours with no further protective effect after that time (Control = 77 ± 2, 9 Ph = 58.2 ± 10, TRPM4_shRNA = 71.3 ± 4.5, TRPM4_scramble = 78.3 ± 4, n = 6, Fig. 6b).

**Discussion**

Here, we report a glutamate-independent mechanism of neuronal reperfusion damage dependent on TRPM4 activation. First, we demonstrate that TRPM4 is expressed in cortical pyramidal neurons. We found that high levels of intracellular ROS and Ca²⁺ during ischemia-reperfusion act to produce a sustained activation of TRPM4, which depolarizes neurons eventually triggering cell death. Pharmacological inhibition with 9-phenanthrol and glibenclamide or shRNA-based silencing of TRPM4 protects neurons from depolarization and cell death up to 3 hours after reperfusion. Additionally, TRPM4_shRNA and 9-phenanthrol did not show an additive effect on the currents. Furthermore, we found that neuronal protection induced by TRPM4 inhibition becomes evident once the glutamate-induced damage i.e. excitotoxicity is blocked, suggesting that TRPM4 participates in the glutamate-independent neuronal damage observed under ischemia-reperfusion injury.

Several reports show that TRPM4 activation strictly depends on intracellular Ca²⁺ ions. Our Ca²⁺ imaging experiments showing that chemically-induced reperfusion increases intracellular Ca²⁺ with a similar time course as the development of the TRPM4-dependent inward current. Moreover, in the absence of extracellular Ca²⁺ ions neuronal depolarization as well as current activation was abolished.

We have previously described a sustained activation of TRPM4 in HeLa and HEK293 cells treated with H₂O₂ by a mechanism that involves the oxidation of the cysteine 1093, which removes channel desensitization, allowing TRPM4 to be permanently activated as long as intracellular Ca²⁺ remains elevated. Also, we demonstrated that cysteine 1093 plays a key role in the increased vulnerability to necrotic cell death. Here we expanded our previous study by measuring endogenous H₂O₂ production induced upon chemical ischemia and reperfusion in a neuronal model. In the presence of NaN₃ + AET, H₂O₂ increases with a time course that parallels Ca²⁺; increases and current

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**Figure 6.** TRPM4 inhibition protects from the reperfusion-induced neuronal death. (a) Quantification of neuronal death after 2 h treatment with 5 mM NaN₃ under 10 μM 9Ph, TRPM4_shRNA and TRPM4_scramble (n = 6). (b) LDH release time course up to 24 h treatment with 5 mM NaN₃ under 9Ph, TRPM4_shRNA and TRPM4_scramble (n = 6). * control vs 9Ph, 1 h, p < 0.05; &&, control vs TRPM4_shRNA, 1 h, p < 0.001; *** control vs 9Ph, 2 h, p < 0.0001; &&& control vs TRPM4_shRNA, 2 h, p < 0.0001; *, control vs 9Ph, 3 h, p < 0.05; &&&& control vs TRPM4_shRNA, 3 h, p < 0.0001; &, control vs TRPM4_shRNA, 24 h, p < 0.05.
development that was partially blocked by NAC. Reperfusion induces mitochondrial and cellular depolarization which is tightly coupled to ROS production. Upon reperfusion, these unpaired electrons combine with faster rate with O₂ producing large amounts of O₂⁻, which impairs ATP production and thus, perpetuating the damage. ROS produced in the mitochondria would remove TRPM4 desensitization and in combination with high Ca²⁺, would trigger a sustained activation of the channel, leading to a permanent depolarized state and neuronal death.

In this study, we show the involvement of TRPM4 in the sustained depolarization observed during reperfusion, and that its inhibition (pharmacological or post-transcriptional silencing) improves the survival of cortical pyramidal neurons in culture, suggesting that TRPM4 is the relevant monovalent cation-selective inward current activated during reperfusion. Therefore, blockade of this channel could increase the survival of neurons during ischemia-reperfusion injury and TRPM4 might constitute a therapeutic target under this condition.

Materials and methods

Cortical neuron culture

Dissociated cortical neuron cultures, glia free, were prepared from embryonic day 18 C57/BL6 mice of either sex as described previously. Cells were seeded (5 × 10⁴ cells/mL) on 12 mm round coverslips coated with poly-L-Lysine (100 μg/mL) and cultured for 14–21 days in Neurobasal Medium-B27 (Invitrogen).

Neuronal transfection

Neurons were transfected in DIV0 after its dissociation. Briefly, neurons were incubated with 1 μg of DNA of a shRNA against TRPM4 (TRPM4 shRNA) expressing EGFP as an expression marker in 2 μL Lipofectamine 2000 (Invitrogen) diluted in 500 μL Minimum Essential Media supplemented (MEM) (Sigma) with 10% horse serum for 5 h. Then, media was replaced with fresh Neurobasal-B27 media and incubated at 37°C in 5% CO₂ until DIV14.

Cell immunofluorescence

Neurons grown in 12 mm coverslip were fixed in 4% w/v paraformaldehyde (10 min exposure) dissolved in
0.01 M PBS pH 7.4 and then washed 3 times in PBS. Cells were permeabilized with 0.01% Triton X-100 (Sigma) (10 min exposure), then blocked in normal goat serum 10% in PBS 1 h at RT (Sigma). Cells were incubated overnight at 4°C with the primary antibody (anti-TRPM4, Alomone, Catalogue Number ACC-044, RRDI AB_2040250) diluted in the same blocking solution. After incubation, cells were washed 3 times with PBS before incubation with the secondary antibody (Alexa Fluor 546, Thermo Catalogue Number A21123, RRDI AB_2535765) for 1.5 h at RT. After 3 five min washes with PBS, cells were mounted with Prolong Gold mounting media (Thermo). Fluorescence was detected in an IX70 DSU spinning disk microscope (Olympus) using 40x 1.4 N.A. magnification objective.

**Neuronal transduction**

Neuronal cultures at 6 DIV (days in vitro) were transduced with engineered lentiviruses expressing a shRNA against TRPM4 (TRPM4_shRNA) or the scrambled sequence (TRPM4_scramble); both expressing EGFP as an expression marker kindly provided by D.J. Linden, John Hopkins University.22

**Chemical ischemia and oxygen-glucose deprivation**

For chemically induced ischemia and reperfusion, neurons were perfused with 5 mM NaN₃ added to the ACSF equilibrated in O₂. Oxygen and glucose deprivation was induced by perfusing neurons with an anoxic ACSF containing 10 mM 2-D-glucose instead of glucose and equilibrated with 95% N₂, 5% O₂ for 1 h previous to the experiments. Neurons were bathed with this buffer for 30 min and then, switched to normal ACSF equilibrated with O₂ (95% O₂, 5% CO₂).

**Immunofluorescence**

For immunofluorescence experiments, we used 5 weeks old C57/BL6 mice. Briefly, after approval by the local ethics committee, animals were anesthetized with isoflurane and heart was perfused with PBS 0.1 M, pH 7.4. The brain was extracted and fixed overnight in 4% w/v paraformaldehyde diluted in PBS 0.1 M, pH 7.4. Brains were sectioned at 50 μm thick using a Vibratome VT1000 (Vibratome). Floating sections were permeabilized in PBS 0.01% Triton, then blocked for unspecific binding with goat serum. Sections were incubated with anti-TRPM4 (1:100, Alomone) primary antibodies for 48 h at 4°C in agitation, and then washed and incubated with the respective secondary antibody (1:1000). We used MAP2 as neuronal marker (1:100, Abcam). Cells were mounted on glass slides with VectaShield and observed on a confocal microscope (Zeiss LSM 510, Zeiss).

**Electrophysiology**

Coverslips were transferred to a temperature controlled (30–35°C) submerged recording chamber continuously perfused at 2–3 mL/min ACSF containing (in mM): 124 NaCl, 25 Na₂HCO₃, 11 glucose, 2.5 KCl, 1.3 MgCl₂, 2.5 CaCl₂, 1.25 NaH₂PO₄, equilibrated with 95% O₂, 5% CO₂. Multiclamp 700 A and Axopatch 200 B amplifiers equipped with Digidata 1322A or 1440 data acquisition boards and pCLAMP10 software (all from Molecular Devices) were used. Bridge balance and access resistance were monitored during recordings and experiments with >20% changes were discarded. All experiments were performed with nystatin-perforated patch clamp (300 μg/mL nystatin were daily made and added to the intracellular buffer) using patch pipettes with an open tip resistance of 4–6 MΩ. Intracellular solution contained (in mM) 130 KCl, 5 NaCl, 10 HEPES, 0.1 BAPTA (pH 7.2, 285 mOsm/kg). The anti-excitotoxic cocktail (AET cocktail) contained: CNQX (10 μM), AP5 (25 μM), picrotoxin (100 μM), nifedipine (100 μM), TTx (1 μM), TEA (10 mM), 4-AP (1 mM), and GdCl₃ (100 μM) to block TRPM7.3

**H₂O₂ and Ca²⁺ imaging**

Changes in [Ca²⁺]ᵢ were monitored as described before.8,50 Briefly, neurons grown in 12 mm round coverslip were incubated with 1 μM Fura 2-AM (Invitrogen) diluted in 0.01% pluronic acid for 1 hour at 37°C in ACSF, then neurons were washed and kept protected from light for 30 min before the measurements. Coverslips were mounted on the stage of an Olympus IX70 microscope, attached to a Sutter Lambda DG-4 high-speed filter unit (Sutter Instruments). Changes in [Ca²⁺]ᵢ were observed using a 40x objective during exposure to 340 and 380 nm, and the intensity of the fluorescence emission 520 nm was recorded using a Hamamatsu Orca-ER CCD camera unit (Hamamatsu) controlled with Micro-manager.
The ratio of fluorescence intensity values at 340/380 reflects the changes in the \([\text{Ca}^{2+}]_{\text{i}}\).

Total \(\text{H}_2\text{O}_2\) was measured using the genetic encoded probe HyPer-Cyto (Evrogen). Briefly, neurons grown in 12 mm round coverslip were transfected at DIV 6 and then measured at DIV 14–21. Changes in intracellular fluorescence were measured by 420 nm excitation and 520 nm emission wavelengths. The vector encoding the protein probe was transfected using Lipofectamine 2000 (Invitrogen). The changes (\(\Delta F\)) in \(\text{H}_2\text{O}_2\) levels were quantified using the formula:

\[
\Delta F = \frac{(F_c - F_b)}{F_b}
\]

Were \(F_c\) is the fluorescence intensity of the cell and \(F_b\) is the fluorescence intensity of the background. Values were normalized to a percentage scale against \(\text{Ca}^{2+}\) at starting time.

**Cell death assay**

Neurons superfused with ACFS were treated with 5 mM NaN₃ for 1 h and viability was determined up to 6 h later, neurons were kept at 37°C in a controlled atmosphere (95% \(\text{O}_2\), 5% \(\text{CO}_2\)) during the whole experiment. To quantify neuronal death, we used two methods, trypan blue incorporation and LDH release. Briefly, neurons were incubated with trypan blue (0.4% for 5 min at room temperature). Dishes were mounted on an optical microscope and stained neurons were counted. LDH activity was measured in the neuronal supernatant by a colorimetric end-point kit (Roche) following the manufacturer’s instructions and using a calibration curve to ensure linearity. Data were expressed as the fraction of maximum release in the presence of 1% Triton.

**Reagents**

Unless otherwise stated, all chemicals and reagents were purchased from Merck KgaA-Chemicals and Sigma.

**Statistics and graphic software**

Unless stated otherwise, all data are presented as a mean ± SEM; statistical differences were tested by two-way ANOVA or Student’s t-test. Graphics were generated using Clampfit 10 (Molecular Devices), GraphPad Prism 6.0 (GraphPad) and Adobe Illustrator CS6 (Adobe) for image composition.

All experiments involving animals were in accordance with the animal protocol approved by the ethical committee of the Universidad de Chile following the rules and guidelines from the Chilean Council of Science and Technology (CONICYT).

**Disclosure of potential conflicts of interest**

No potential conflicts of interest were disclosed.

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**Author contributions**

E.L.-S. and A.S. designed research; D.R., O.C., and E.L.-S. performed research; D.R., O.C., E.L.-S., and A.S. analyzed data; E.L.-S. and A.S. wrote the paper.

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