Cardioprotective effects of Prolame and SNAP are related with nitric oxide production and with diminution of caspases and calpain-1 activities in reperfused rat hearts

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ABSTRACT

Cardiac tissue undergoes changes during ischemia-reperfusion (I-R) that compromise its normal function. Cell death is one of the consequences of such damage, as well as diminution in nitric oxide (NO) content. This signaling molecule regulates the function of the cardiovascular system through dependent and independent effects of cyclic guanosine monophosphate (cGMP). The independent cGMP pathway involves post-translational modification of proteins by S-nitrosylation. Studies in vitro have shown that NO inhibits the activity of caspases and calpains through S-nitrosylation of a cysteine located in their catalytic site, so we propose to elucidate if the regulatory mechanisms of NO are related with changes in S-nitrosylation of cell death proteins in the ischemic-reperfused myocardium.

We used two compounds that increase the levels of NO by different mechanisms: Prolame, an amino-estrogenic compound with antiplatelet and anticoagulant effects that induces the increase of NO levels in vivo by activating the endothelial nitric oxide synthase (eNOS) and that has not been tested as a potential inhibitor of apoptosis. On the other hand, S-Nitroso-N-acetylpenicillamine (SNAP), a synthetic NO donor that has been shown to decrease cell death after inducing hypoxia-reoxygenation in cell cultures. Main experimental groups were Control, I-R, I-R + Prolame and I-R + SNAP. Additional groups were used to evaluate the NO action pathways. Contractile function represented as heart rate and ventricular pressure was evaluated in a Langendorff system. Infarct size was measured with 2,3,5-triphenyltetrazolium chloride stain. NO content was determined indirectly by measuring nitrite levels with the Griess reaction and cGMP content was measured by Enzyme-Linked ImmunoSorbent Assay. DNA integrity was evaluated by DNA laddering visualized on an agarose gel and by Terminal deoxynucleotidyl transferase dUTP Nick-End Labeling assay. Activities of caspase-3, caspase-8, caspase-9 and calpain-1 were evaluated spectrophotometrically and the content of caspase-3 and...
calpain-1 by western blot. S-nitrosylation of caspase-3 and calpain-1 was evaluated by labeling S-nitrosylated cysteines. Our results show that both Prolame and SNAP increased NO content and improved functional recovery in post-ischemic hearts. cGMP-dependent and S-nitrosylation pathways were activated in both groups, but the cGMP-independent pathway was preferentially activated by SNAP, which induced higher levels of NO than Prolame. Although SNAP effectively diminished the activity of all the proteases, a correlative link between the activity of these proteases and S-nitrosylation was not fully established.

Subjects  Biochemistry, Cell Biology, Cardiology
Keywords  Cell death, Reperfusion, Caspases, Calpain, Nitric oxide, S-nitrosylation

INTRODUCTION
Nitric oxide (NO) is a short-life molecule produced after the oxidation of the guanidine group of L-arginine by NO synthases (NOS) (Moncada, Palmer & Higgs, 1991; Loscalzo & Welch, 1995). NO modulates cardiac function by regulating vascular tone, excitation-contraction coupling (Hammond & Balligand, 2012), platelet aggregability (Ikeda et al., 2000) and mitochondrial function (Zhao et al., 2005). Such effects are mainly associated either with the activation of soluble guanylate cyclase (sGC) that produces cyclic guanosine monophosphate (cGMP) and stimulates protein kinase G (PKG) or, with redox reversible modification of cysteine residues that results in the formation of S-nitrosothiols (SNO) in a process named S-nitrosylation (Stamler, Lamas & Fang, 2001). However, the finding that NO inhibits both Complex III and Complex IV activities in isolated mitochondria (Poderoso et al., 1996; Giuffrè et al., 1996) has raised the possibility of its role as a direct physiological regulator of mitochondrial respiration (Poderoso, Helfenberger & Poderoso, 2019).

It is well demonstrated that NO, either derived from NOS isoforms or exogenously administered protects against ischemia-reperfusion (I-R)-induced injury (Bolli, 2001; Jones et al., 2004; Hu et al., 2016); however, the preponderance of cGMP-dependent or of cGMP-independent pathways in such protection is still under debate. It was described for example, that sGC inhibition abrogate the post-conditioning infarct-sparing effect in rabbit hearts, suggesting a preponderant role of cGMP-dependent mechanisms (Yang et al., 2004, 2005). On the other hand, we and others have reported that NO-mediated cardioprotection might be also regulated by S-nitrosylation signaling. Sun et al. (2013) demonstrated that blockage of the sGC/cGMP/PKG signaling pathway did not affect ischemic postconditioning-mediated cardioprotection and this finding was correlated with increased SNO levels; whereas our group described the partial recovery of heart function when the NO donor: \([Z]-1-[N-(2-aminoethyl)-N-(2-ammonioethyl)amino]diazene-1-im-1,2-diolate \) (DETA-NO) was administrated to postconditioned hearts in which sGC was inhibited (Correa et al., 2015).

NO-mediated cardioprotection has been associated with S-nitrosylation of diverse proteins, like intracellular calcium handling proteins (e.g., L-type Ca\(^2+\) channels and sarcoplasmic reticulum calcium pump SERCA2a) (Sun et al., 2007), mitochondrial
proteins (Sun et al., 2015) and antioxidant response proteins (Tao et al., 2004), but it has not been demonstrated if the activity of cell death-related proteins are under S-nitrosylation regulation in reperfusion damage. Also, as increased SNO may have potential clinical significance, it is relevant to determine if NO produced by the canonical protein kinase B (Akt)/endothelial nitric oxide synthase (eNOS) signaling or that produced by NO donors induce different levels of S-nitrosylation in the context of cardioprotection.

Therefore, the goal of this study was to determine if proteins related with cell death are susceptible to S-nitrosylation regulation in isolated hearts treated either with the estrogen analog Prolame, that increases NO levels through the activation of the PI3K/Akt/eNOS signaling pathway (Hernández-Reséndiz et al., 2015) and with the synthetic NO donor S-Nitroso-N-acetylpenicillamine (SNAP) which bypasses eNOS activity.

**MATERIAL AND METHODS**

**Reagents**

Chemicals were of reagent or higher grade from Sigma-Aldrich (St Louis, MO, USA) unless otherwise specified. Anti-caspase-3 monoclonal (sc-136219), anti-calpain-1 monoclonal (sc-271313), agarose-conjugated (AC) anti-caspase-3 (sc-136219) and AC anti-calpain-1 antibodies (sc-271313) were from Santa Cruz Biotechnology (Santa Cruz, CA, USA). Fluorogenic calpain-1 substrate (208748) and colorimetric caspase-3 substrate (235400) were from Calbiochem (Darmstadt, Germany); colorimetric caspase-8 substrate (260-045-M005) and colorimetric caspase-9 substrate (260-081-M005) were from Enzo Life Sciences (Farmingdale, NY, USA). The enhanced chemiluminescence detection system was from Millipore Corporation (Bedford, MA, USA) and horseradish peroxidase-conjugated secondary antibodies were from Santa Cruz Biotechnology (Santa Cruz, CA, USA); In situ Cell Death Detection Kit, Fluorescein (11-684-795-910) was from Roche; Cyclic GMP Enzyme-linked Immunosorbent Assay Kit was purchased from Cayman Chemical (Ann Arbor, MI, USA); whereas Pierce S-Nitrosylation Western Blot Kit was from Thermo Scientific. 1H-(1,2,4)Oxadiazolo(4,3-a)quinoxalin-1-one (ODQ) was from Calbiochem (Darmstadt, Germany) and 17β-(3-hydroxy-1-propylamino)-1,3,5(10)-estratrien-3-ol, Prolame, was synthesized and chemical purity established as previously reported (Fernández-G et al., 1985).

**Ethical approval**

The investigation was approved by the Ethics Committee of the National Institute of Cardiology, “Ignacio Chávez” (14CB09012016) and the experimental protocols followed the guidelines of Norma Oficial Mexicana for the use and care of laboratory animals (NOM-062-ZOO-1999) and for disposal of biological residues (NOM-087-SEMARNAT-SSA1-2002).

**Experimental design**

Male wistar rats (300–350 g) were anaesthetized by injecting intraperitoneally a single dose of sodium pentobarbital (60 mg/kg i.p.) plus sodium heparin and complete lack of pain response was assessed by determining pedal withdrawal reflex. Hearts were perfused...
retrogradely on a Radnoti Langendorff heart perfusion system (ADInstruments, Sydney, NSW, Australia) via the aorta at a constant flow rate of 12 ml/min with Krebs-Henseleit solution (118 mM NaCl, 4.75 mM KCl, 1.18 mM KH₂PO₄, 1.18 mM MgSO₄ · 7H₂O, 2.5 mM CaCl₂, 25 mM NaHCO₃, five mM glucose and 0.1 mM sodium octanoate, pH 7.4), which was continuously bubbled with 95% O₂ and 5% CO₂ at 37 °C. Cardiac performance was measured at left ventricular end-diastolic pressure of 10 mmHg using a latex balloon inserted into the left ventricle and connected to a pressure transducer that, in turn, was connected to a PowerLab System (ADInstruments, Sydney, NSW, Australia). Throughout the experiment, left ventricular developed pressure (LVDP) and heart rate (HR) was calculated automatically from the pressure trace with the digital acquisition system LabChart 8.1.5 Pro (ADInstruments, Sydney, NSW, Australia). HR was expressed as beat number min⁻¹ and the double product (DP) was calculated by multiplying HR by LVDP. Hearts were perfused for 20 min to reach a steady state and then subjected to the different protocols. The experimental groups were: (1) Control, hearts perfused for additional 110 min; (2) I-R, hearts subjected to global ischemia for 30 min by turning off the pumping system and 60 min of reperfusion; (3) I-R+Prolame, hearts that received 1.25 μM of the compound for 5 min before ischemia; (4) I-R+SNAP, hearts perfused with two μM of the compound during 5 min before ischemia and 10 min during reperfusion (Fig. 1A). These concentrations were chosen after performing dose-response experiments of heart function recovery.

Additional experiments were performed administrating 15 μM L-NAME, inhibitor of the eNOS and 50 μM ODQ, inhibitor of the sGC before the addition of Prolame and SNAP, as previously reported (Correa et al., 2015). Both SNAP and ODQ are light sensitive, therefore heart perfusion and sample preparation were performed in the dark. At the end of the experiments, unless otherwise indicated, heart tissue was preserved in cold buffer (100 mM Tris–HCl, 145 mM NaCl, 10 mM EDTA, pH 7.3).

Infarct size measurement
Infarct size was measured using 2,3,5-triphenyltetrazolium chloride (TTC) staining (Ramírez-Camacho et al., 2016). Heart was frozen and cut into three mm transverse slices. The slices were incubated in 1% TTC solution for 20 min at 37 °C and then immersed in formalin solution for 20 min to enhance the contrast between stained and unstained tissue. TTC stains living tissue into a deep red color. Digital images of heart slices were obtained and analyzed using the ImageJ 1.48 software (NIH, Bethesda, MD, USA). Infarct size was expressed as a percentage and calculated by dividing the area of infarct by total area at risk.

Preparation of tissue homogenates
Cardiac fresh tissue was frozen in liquid nitrogen, pulverized with a pistil in a mortar and homogenized in a buffer containing 100 mM HEPES, 20% glycerol and 0.5 mM EDTA, pH 7.5 and centrifuged at 10,000×g for 10 min at 4 °C. The supernatant fraction was recovered and immediately frozen in liquid nitrogen. Protein was measured by the Lowry method (Lowry et al., 1951).
Preparation of cytosolic fractions

Cardiac fresh tissue was minced with scissors and homogenized in Tris-buffered saline (100 mM Tris–HCl, 145 mM NaCl, pH 7.3) plus EDTA (10 mM) and centrifuged at 10,000×g for 10 min at 4 °C. The cytosolic fraction was recovered after centrifuging the supernatant at 100,000×g for 60 min and immediately frozen in liquid nitrogen (Kohli et al., 1997). Protein was measured by the Lowry method (Lowry et al., 1951).

Determination of nitrite levels

Nitrite levels were measured using the colorimetric Griess reagent as indicator of NO production. Fresh cardiac tissue was homogenized in 50 mM Tris, 120 mM NaCl, 0.05% IGEPAL, pH 8.0, centrifuged at 10,000×g for 20 min at 4 °C and the supernatants filtrated through syringe-driven filter units (Merck KGaA, Darmstadt, Germany) to eliminate proteins from each sample. Nitrates were reduced to nitrites with bacterial nitrate reductase of E. Coli. The samples were then incubated with 0.2 ml of Griess solution.
(0.1% sulfanilamide and 0.5% of N-(1-naphthyl)ethylenediamine dihydrochloride in phosphoric acid 2.5%) for 60 min at 37 °C in the dark. The formed azo dye was spectrophotometrically quantified at 543 nm and compared against a sodium nitrate concentration curve incubated with the enzyme.

**cGMP content**
Cyclic guanosine monophosphate levels were measured by competitive Cyclic GMP EIA kit (Cayman Chemical, Ann Arbor, MI, USA) in homogenates from the indicated experimental groups according to the manufacturer’s instructions.

**DNA Fragmentation assay**
Frozen cardiac tissue (five mg) was pulverized and incubated with 0.5 ml extraction buffer (10 mM Tris, 100 mM EDTA, 5% SDS; pH 8.0) plus 20 μg/ml of RNAse for 1 h at room temperature. Then, 200 μg/ml of Proteinase K were added and incubated overnight at 50 °C. An equal volume of phenol adjusted with Tris buffer was added to the samples and maintained under constant agitation during 60 min at 4 °C. The samples were centrifuged at 5,000×g for 30 min at room temperature; the aqueous-viscous phase was carefully transferred to a new tube and the extraction was repeated with an equal volume of phenol/chloroform (1:1 v/v). The aqueous phase was transferred to a new tube and the DNA precipitated by adding 0.1 volumes of 3M sodium acetate and two volumes of absolute ethanol. The samples were centrifuged at 5,000×g for 20 min at room temperature; the supernatant was carefully removed, and 0.5 ml of 70% ethanol was added to rinse the DNA, which was dissolved in extraction buffer. DNA was analyzed in a 1.5% agarose gel with 0.5 μg/ml ethidium bromide.

**TUNEL assay**
Samples of ventricular tissue were fixed in 4% paraformaldehyde, washed with PBS three times for 5 min at room temperature and incubated in 30% sucrose for 24 h at 4 °C. Transversal cryosections of 10 μm from each heart were obtained (Minotome PLUS™, Digital Microtome Cryostat, Triangle Biomedical Sciences (TBS, Inc.), USA) and mounted on gelatinized slides. The tissue sections were rehydrated in PBS for 30 min at room temperature, permeabilized for 2 min at 4 °C with a fresh solution of 0.1% triton X-100 and 0.1% sodium citrate and washed with PBS two times for 5 min at room temperature. Each tissue section was rounded with hydrophobic pen and incubated with Terminal deoxynucleotidyl transferase dUTP Nick-End Labeling (TUNEL) reaction mixture (terminal deoxy nucleotide transferase plus nucleotide mixture in reaction buffer) in a humidified atmosphere at 37 °C in the dark for 60 min. After incubation, the slides were washed with PBS three times for 5 min at room temperature, counterstained with 4',6-Diamidino-2-phenylindole dihydrochloride to visualize the nucleus and mounted for fluorescence microscopy analysis (Floid Cell Imaging Station; Life Technologies, Carlsbad, CA, USA). Tissue sections of negative (without transferase in the buffer reaction) and positive controls (pretreated with 3,000 U/ml recombinant DNase I in Tris–HCl 50 mM, pH 7.5, one mg/ml BSA) were included in the assay.
Caspase-3, -8 and -9 activity
Caspase-3, -8 and -9 activities were measured using 200 μM of the colorimetric substrate Ac-DEVD-pNA, Ac-IETD-pNA and Ac-LEHD-pNA respectively, in a total volume of 0.3 ml, containing 30 μl of the cytosolic fraction and 240 μl of HEPES buffer (100 mM HEPES, 20% glycerol, five mM DTT, 0.5 mM EDTA, pH 7.5) at 37 °C for 60 min in 96-microwell plates. Changes in absorbance were evaluated at 405 nm in a microplate reader (BioTek Instruments, Inc., Winooski, VT, USA). The activities of caspases were normalized by mg of protein.

Calpain-1 activity
Fifty μg of cytosolic fractions, 10 mM CaCl₂ and 10 μM of the synthetic fluorogenic substrate for calpain H-K (FAM)-EVY-GMMK (DABCYL)-OH (Merck Darmstadt, Germany) were added to reaction buffer containing 100 mM Tris–HCl, 145 mM NaCl, pH 7.3 in a total volume of 0.2 ml and incubated at 37 °C for 60 min in 96-microwell plates. To measure calcium-independent activity, CaCl₂ was replaced with reaction buffer. Increase in fluorescence was measured in a microplate reader (BioTek Instruments, Inc., Winooski, VT, USA) at λ_em and λ_ex of 518 and 490 nm, respectively. The calpain activity was normalized by mg of protein.

Immunoprecipitation of caspase-3 and calpain-1
Frozen cardiac tissue from the different groups were homogenized in PBS buffer (136.9 mM NaCl, 2.67 mM KCl, 8.1 mM Na₂HPO₄, 1.47 mM KH₂PO₄; pH 7.4). The homogenates were maintained under constant agitation for 2 h at 4 °C and then centrifuged at 10,000×g for 20 min at 4 °C. The supernatants were recovered, placed in a new tube in cold and protein was measured by the Lowry method. Duplicated samples containing the same amount of protein (one-mg) were taken to a final volume of one ml with PBS buffer. Then, five μl of calpain-1 antibody or of caspase-3 antibodies coupled to agarose (slurry) were added to each sample and incubated overnight at 4 °C under constant agitation. The samples were centrifuged at 3,000×g for 2 min at 4 °C, the supernatant was carefully removed, and one ml of PBS was added and gently inverted by hand to wash the pellet. This step was repeated one more time and the pellet was recovered to evaluate total protein and S-nitrosylation levels.

S-nitrosylation assay
Protein S-nitrosylation was measured by selective reduction and labeling of S-nitrosylated cysteines of the immunoprecipitated proteins with the Pierce S-Nitrosylation Western Blot Kit (Thermo Scientific, Waltham, MA, USA) according to the manufacturer’s instructions. As the cysteine biotinylation is reversible, the samples were prepared without reducing agents.

Statistical analysis
Data analysis was performed by analysis of variance followed by Bonferroni’s test, using the Graph Pad PRISM 5 for windows version 5.03 software. Results were expressed as mean ± SD or as otherwise indicated; P-values < 0.05 were considered statistically significant.
RESULTS

Prolame and SNAP maintain cardiac function and reduce infarct size in reperfused hearts

Cardiac function expressed as the DP was maintained in the Control group during 110 min of constant reperfusion. DP decreased early and until the end of reperfusion in the I-R group ($P < 0.05$). Both SNAP and Prolame groups recovered cardiac function during reperfusion (Fig. 1B). Accordingly, infarct size decreased in hearts treated with Prolame or SNAP compared with the group I-R (9 ± 1.6 and 6.5 ± 3.5 vs. 56.2 ± 3.7; respectively; Fig. 1C).

Prolame preserves and SNAP increases nitric oxide in I-R hearts

Nitrite content, an indicator of NO levels diminished significantly in I-R homogenates as compared with the Control group (3.8 ± 0.9 vs. 41.5 ± 6.7 nmol nitrite/mg protein; $P < 0.05$). Conversely, both Prolame and SNAP increased the levels of nitrite even to higher levels than those measured in the Control group (84.1 ± 5.6 and 120.8 ± 7.3 nmol nitrite/mg protein, respectively; $P < 0.05$, Fig. 1D). These data correlate with heart function recovery and decrease of cell death.

Contribution of cGMP-dependent and independent nitric oxide pathways in the cardioprotective effect of Prolame and SNAP

To evaluate the participation of the cGMP-dependent and independent mechanisms activated by NO in cardiac function recovery exerted by Prolame and SNAP, we administrated the NO synthase inhibitor N5-[imino(nitroamino)methyl]-L-ornithine, methyl ester, monohydrochloride (L-NAME) and the guanylate cyclase inhibitor ODQ to the I-R+Prolame hearts (Fig. 2A); whereas only ODQ was administrated to the I-R+SNAP hearts (Fig. 3A).

Neither ODQ nor L-NAME had effect in HR (Figs. 2B and 3B) or in LVDP (Figs. 2C and 3C) during the stabilization period before Prolame or SNAP administration. L-NAME abolished the cardioprotective effect conferred by Prolame in both HR (Fig. 2B) and in LVDP at the end of reperfusion (Fig. 2C); whereas ODQ administration to Prolame treated hearts diminished significantly HR from 257.4 ± 21.5 to 198.4 ± 19.2 beats min$^{-1}$ ($P < 0.05$) and LVDP from 72.6 ± 4.1 to 58.6 ± 3.3 mmHg ($P < 0.05$) indicating that cGMP-independent mechanisms or other pharmacological properties non-related with NO production contribute to cardioprotection by Prolame. Blocking the cGMP-dependent pathway in the I-R+SNAP group, gives further clarity to the mechanisms activated by NO that contribute to cardioprotection and supports the idea that NO sustains S-nitrosylation processes. In this group, ODQ diminished HR from 294.2 ± 10.7 to 231.2 ± 42.7 beats min$^{-1}$ ($P < 0.05$, Fig. 3B), but had no effect on LVDP values, which remain comparable to those observed in the SNAP group (86 ± 10.4 vs. 72.8 ± 6.9 mmHg; Fig. 3C).

To confirm that the inhibitors were efficiently acting on their corresponding targets, we measured cGMP content in all the experimental groups. The levels of cGMP decreased significantly in the I-R group in comparison with the Control group (1 ± 0.6 vs. 20.2 ± 3.2 pmol cGMP/ml; $P < 0.05$); whereas hearts treated with Prolame and with SNAP...
showed values comparable to those measured in the Control group (24.3 ± 5.1 and 19.6 ± 2.5 pmol cGMP/ml). Both ODQ and L-NAME diminished cGMP levels in the Prolame group to 3.1 ± 1.4 and 1.4 ± 0.6 pmol cGMP/ml respectively; whereas ODQ lowered cGMP to 0.5 ± 0.4 pmol/ml in SNAP-treated hearts (Fig. 4). These results unravel the participation of S-nitrosylation in the cardioprotection conferred by these compounds.

To discriminate between eNOS activation and other PI3K/Akt activated pathways in the conferred-cardioprotection, we also measured the levels of NO in the presence of ODQ and L-NAME. NO levels were 70.4 ± 2.8 and 24.8 ± 3.7 nmol nitrite/mg protein in the Prolame+ODQ and Prolame+L-NAME groups, respectively. L-NAME significantly decreased NO levels in comparison with the Prolame group, but such values were higher than those observed in the I-R group (3.03 nmol nitrite/mg protein, \( P < 0.05 \)), indicating that the effects of Prolame are partially mediated through the NO. Also, differences were found between the I-R+SNAP+ODQ and the I-R+SNAP groups (86 ± 4.5 vs. 120 ± 7.3 nmol nitrite/mg protein).

Figure 2 Effect of ODQ and L-NAME on the cardioprotection conferred by Prolame. (A) Schematic representation of the experimental procedure in the Langendorff model using the inhibitory compounds ODQ and L-NAME. Ischemia-reperfusion (I-R) group, with 30 min of ischemia and 60 min of reperfusion; I-R+Prolame group, in which Prolame (1.25 μM) was added only 5 min before ischemia; I-R+Prolame+ODQ group, ODQ (15 μM) was added 15 min at the onset of the stabilization and before the addition of Prolame; I-R+Prolame+L-NAME group, L-NAME (50 μM) was added 15 min at the onset of stabilization and before the addition of Prolame. (B) Heart rate expressed as beats per minute. Values are expressed as mean ± SD of five independent heart preparations per group. The statistical test used was two-way ANOVA. \( ^* P < 0.001 \) I-R vs. I-R+Prolame and I-R+Prolame+ODQ; \( ^$ P < 0.05 \) I-R+Prolame vs. I-R+Prolame+ODQ. \( ^& P < 0.001 \) I-R+Prolame vs. I-R+Prolame+L-NAME. (C) LVDP (left ventricular developed pressure). Values are expressed as mean ± SD of five independent heart preparations per group. The statistical test used was two-way ANOVA. \( ^* P < 0.001 \) I-R vs. I-R+Prolame and I-R+Prolame+ODQ; \( ^$ P < 0.05 \) I-R+Prolame vs. I-R+Prolame+ODQ; \( ^& P < 0.001 \) I-R+Prolame vs. I-R+Prolame+L-NAME.
Effect of nitric oxide on apoptosis and necrosis

We performed TUNEL and DNA ladder analysis to evaluate the effect of NO on cell death regulation. Our results showed few apoptotic nuclei in I-R hearts, whereas the DNA ladder image clearly showed a mixture of necrotic/apoptotic cell death. Both Prolame and SNAP reduced condensed nuclei and DNA fragmentation (Figs. S1A–S1F).

Effect of Prolame and SNAP on caspases and calpain-1 activities

We also evaluated the participation of apoptosis by measuring the activities of caspases using specific synthetic substrates. Caspase-8 and caspase-9 activities increased significantly in the I-R group as compared with the control group ($P < 0.05$) and SNAP diminished the activity of the three caspases ($P < 0.05$). Prolame slightly reduced their activity, but none statistical differences were found between I-R and I-R+Prolame hearts (Figs. 5A–5C). On the other hand, the activity of calpain-1 increased by 1.8-fold in I-R hearts as compared with the control group and diminished in both Prolame and SNAP-treated groups to basal levels (Fig. 5D). These results suggest that both pathways promote cell death during reperfusion, and that the activity of all the evaluated proteases is subjected to NO regulation.

We also measured the levels of cleaved caspase-3 (Fig. S2A) by western blot in heart homogenates. A significant increase of the p20 fragment of active caspase-3 was observed...
in I-R hearts which was diminished by SNAP or Prolame ($P < 0.05$). On the other hand, the levels of small subunit of calpain-1 does not showed significant differences between the experimental groups (Fig. S2B).

**Effect of Prolame and SNAP in SNO of caspase-3 and calpain-1**

It has been proposed that NO blocks apoptosis by modifying redox-sensitive cysteine residues in members of the caspase family. To determine if the observed inhibition in the activity of these proteases and of calpain-1 was associated with their levels of SNO, we labeled the S-nitrosylated residues of immunoprecipitated caspase-3 and calpain-1 with iodoacetyl TMT (iodo TMTTM) reagent. Anti-caspase-3 antibodies pull down mainly the p-20 subunit of caspase-3 and in much lower extent the procaspase form. The amount of the cleaved subunit was lower in I-R+SNAP than in I-R hearts; conversely immunoprecipitated procaspase-3 increased in the SNAP group and diminished in I-R hearts. Anti-TMT antibodies only recognized the full-length caspase-3 and, although there are no significant differences, the S-nitrosylation signal tends to increase in the I-R+Prolame and I-R+SNAP groups in comparison with the I-R group (0.928 ± 0.09 and 0.733 ± 0.1 vs. 0.671 ± 0.3; respectively, Fig. 6A). On the other hand, the S-nitrosylation analysis of the small subunit of calpain-1 shows a tendency to increase in I-R hearts treated with Prolame or SNAP in comparison with the I-R group (0.724 ± 0.1 and 0.767 ± 0.2 vs. 0.485 ± 0.1, respectively, Fig. 6B). We recognize that direct S-nitrosylation experiments shown in Fig. 6 are not conclusive. Despite its widespread use, the biotin-switch technique is challenging and
each step contains potential error sources. However, the maintenance of residual cardiac function in conditions where GMPc was depressed, have led us to maintain the speculation that S-nytrosylation plays a role in the cardioprotective effect of NO.

**DISCUSSION**

This study shows that both Prolame and SNAP diminish infarct size, maintain NO levels and prevent from cardiac reperfusion damage by activating cGMP-dependent and S-nitrosylation pathways. We observed that even though infarct size was negligible in reperfused hearts treated with Prolame, cardiac function diminished significantly in this group as compared with the Control group during reperfusion. Post-ischemic contractile dysfunction or “myocardial stunning,” associated with infarct size reduction has been described by other groups (Čarnická et al., 2011; Ford et al., 2001). Our results show that the NO donor SNAP is more efficient than Prolame to prevent from this benign form of reperfusion damage. Both compounds activate the cGMP-independent pathway in association with diminution in the activities of proteases linked to apoptosis and necrosis during reperfusion.

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**Figure 5** Activities of caspases-3, -8, -9 and calpain-1 in reperfused hearts treated with Prolame or SNAP. Activities of caspase-3 (A), -8 (B) and -9 (C) in all experimental groups. All caspase activities were measured as p-nitroaniline (pNA) release with respect to an aniline concentration curve. Data are expressed as mean ± SEM of six independent heart preparations per group. The statistical test used was one-way ANOVA. Caspase-3 activity:  †P < 0.05 vs. I-R+SNAP; caspase-8 activity:  †P < 0.05 vs. Control and I-R+SNAP; Caspase-9 activity:  †P < 0.05 vs. Control and I-R+SNAP. (D) Activity of calpain-1 in all experimental groups. Data are expressed as mean ± SEM of at least nine independent heart preparations per group. The statistical test used was two-way ANOVA.  †P < 0.05 vs. all groups.
Necrosis is described as accidental non-apoptotic cell death that releases cellular content to the extracellular medium resulting in inflammation. On the other hand, apoptosis is a form of programmed cell death that is characterized by nuclear and cytoplasmic condensation and the formation of apoptotic bodies. It has been described that necrosis, apoptosis and in minor extent autophagy contribute to cell death in reperfused hearts; however, this issue remains under debate. While necrosis has been related with ischemic damage, apoptosis is thought to occur only during reperfusion and to coexist with necrotic death (Gottlieb et al., 1994; Vakeva et al., 1998; Zhao et al., 2000). The use of caspase inhibitors to reduce reperfusion damage have rendered conflicting results, weakening the idea of apoptosis relevance in reperfusion damage. For example, Chapman et al. (2002) reported that the selective caspase-3 inhibitor (S)-(+)\(-\)-5-(1-(2-methoxymethylpyrrolidinyl)sulfonyl)isatin diminishes infarct size in isolated rabbit hearts and reduce apoptotic cell death; whereas other groups found that the same compound improves heart function independently of apoptosis regulation in rats (Kovacs et al., 2001; Ruetten et al., 2001).

Despite such controversial results, it has been reported that NO-activated pathways attenuate both apoptotic and necrotic cell death. The sGC-cGMP-PKG pathway modulates apoptotic cell death by increasing Bcl2/BAX ratio (Das, Xi & Kukreja, 2005) and reduces necrotic cell death through several mechanisms that ultimately maintain

Figure 6 S-Nitrosylation of caspase-3 and calpain-1 in reperfused hearts treated with Prolame or SNAP. (A) Immunoblot of immunoprecipitated caspase-3 and (B) S-nitrosylated caspase-3. (C) bars representing the densitometric ratio between SNO procaspase-3/total procaspase-3. Data are mean ± SEM of three independent heart preparations per group. The statistical test used was one-way ANOVA. (D) Immunoblot of immunoprecipitated small subunit of calpain-1 and (E) S-nitrosylated small subunit of calpain-1. (F) bars representing the densitometric ratio between SNO calpain-1/total calpain-1. Data are mean ± SEM of five independent heart preparations per group. The statistical test used was one-way ANOVA.
mitochondrial function. It is known that PKG phosphorylates L-type Ca\textsuperscript{2+} channels reducing calcium uptake during phase-2 of the action potential \citep{ZhaoGreensteinWinslow2017} and that activates the sarcoplasmic reticulum calcium ATPase (SERCA) promoting the dissociation of the regulator phospholamban after its phosphorylation at residue Ser\textsuperscript{16} \citep{InserteEtAl2014}. It also induces the opening of mitoK\textsubscript{ATP} channels \citep{PennaEtAl2017}, which enhance ROS production and promotes Protein Kinase C-epsilon (PKC-\epsilon) activation \citep{CostaGarlid2008}. The regulatory effect of this and other downstream kinases, for example, Akt, ERK1/2 and GSK3\beta \citep{HausenloyYellon2006} in conjunction with the attenuation of Ca\textsuperscript{2+} overload, inhibits the opening of the mitochondrial permeability transition pore (mPTP) avoiding irreversible cell injury \citep{CostaGarlid2008}. Of note, even though PKG is considered the main intracellular mediator of cGMP, this cyclic nucleotide regulates other complex signaling cascades paramount to the cardiovascular system function, that includes several cGMP-protein kinases, cGMP-regulated phosphodiesterases \citep{DasEtAl2008,Takimoto2012} and cyclic nucleotide-gated ion channels \citep{SpinelliEtAl2018}.

On the other hand, S-nitrosylation of cysteine residues of L-type channels \citep{GonzalezEtAl2009} and of the ryanodine receptor \citep{GonzalezEtAl2007} regulate their activity and maintain calcium homeostasis. Noteworthy, it has also been suggested that conforming and regulator proteins of the mPTP (adenine nucleotide translocase and cyclophilin D, respectively) have potential cysteine residues for S-nitrosylation \citep{ChangEtAl2014}.

The proposal that NO might block apoptosis by modifying redox-sensitive cysteine residues in the members of the caspase family, arise from observations that S-nitrosylation concurs with inhibition of caspase-3 activity in vitro and that antibodies against S-nitrocysteine immuno-react with this protease \citep{MaejimaEtAl2005}. Our results show that the activity of this caspase, as well as of caspase-8 and caspase-9 decrease in the I-R+SNAP group. We found that S-nitrosocysteine post-translational modifications in caspase-3 were associated with the unprocessed enzyme and not with the active form, although this fragment was preferentially pulled down during immunoprecipitation. In this sense, \cite{MannickEtAl1999} have proposed that during apoptosis, mitochondrial procaspase-3 is released into the cytoplasm where it is activated by denitrosylation. Also, it was reported that S-nitrosylation at cysteine 163 in the catalytic site of procaspase-3 inhibits its cleavage \citep{LaiEtAl2011,SiligramaEtAl2014}, whereas S-nitrosylated active caspase-3 (p17) has only been reported by a research group in immune cells infected by bacterial pathogens \citep{DunneEtAl2013}. Regarding other caspases, studies in hepatocytes demonstrated that NO prevents the proteolytic activation of caspase-8 \citep{LiEtAl1999} and, that S-nitrosylation of active caspase-8 decreases its activity and prevents death by apoptosis \citep{KimEtAl2000}. Some reports indicate that the zymogen of caspase-9 is nitrosylated in mitochondria \citep{MannickEtAl2001} and that cysteine 325 is a critical S-nitrosylation site \citep{ZhangEtAl2016}.

We also found that the small subunit of calpain-1 showed increased S-nitrosylation in I-R+SNAP group as compared with I-R hearts. In this sense, a recent report showed that S-nitrosylation induced by S-nitroso-N-acetylpenicillamine (SNAP) concurs with diminution of cardiac calpain activity in murine cardiomyocytes subjected to hypoxia/
reoxygenation, as well as in an in vivo reperfusion model (Totzeck et al., 2017). Calpain-1, a member of Ca\(^{2+}\)-dependent cysteine proteases has a central role in myocardial injury during I-R, inducing cell death by apoptotic (Iwamoto et al., 1999; Gao & Dou, 2000) and by necrotic processes (Aguilar et al., 1996). It has been described that this protease mediates the translocation of the mitochondrial apoptosis inducer factor (AIF) to the cytosol and nucleus (Chen et al., 2011) and that also cleaves the pro-apoptotic protein bid, favoring the activation of the mitochondrial-dependent pathway of apoptosis (Chen et al., 2001).

Other effects of cytosolic calpain activation are enzymatic degradation of proteins like actin, ankyrin, caldesmon, troponin T, troponin I, titin and desmone (Neuhof & Neuhof, 2014; Patterson et al., 2011; Potz et al., 2015), in addition to many other proteins, which produce membrane fragility and further calcium overload. This condition increases mitochondrial calcium concentration promoting the opening of the mPTP, which is considered the “hallmark event” in necrotic cell death. Thus, calpain-1 activation might induce necrosis by promoting calcium overload, but also as suggested by Thompson et al. (2016) it might directly modulate the mPTP. These group demonstrated that the administration of the calpain inhibitor MDL-28170 to isolated reperfused hearts decreases the release of lactate dehydrogenase and prevents from mPTP opening.

The factors that determine the preponderance of cGMP-dependent pathway over S-nitrosylation pathway remains to be clarified, but it has been speculated that the concentration and/or the site of NO generation might be relevant. In this sense, it has been reported that 100 μM of SNAP increases the levels of NO and cGMP in association with decreased contractility; whereas lower SNAP concentrations (0.1–1 μM) are related with lower NO content, increased contractility and S-nitrosylation in isolated rat hearts (González et al., 2008). In our experiments, although SNAP induces higher levels of NO than Prolame, cGMP levels were similar in both groups. ODQ abolished cGMP levels in association with partial loss of the cardioprotective effect, demonstrating the participation of the cGMP-independent pathway in hearts treated with Prolame and with SNAP. However, we observed that S-nitrosylation seems to be favored in the cardioprotection conferred by SNAP, as no differences in LVDP were detected between IR+SNAP vs. IR+SNAP+ODQ (Figs. 3B and 3C) in comparison with the same parameters in IR+Prolame vs. IR+Prolame+ODQ (Figs. 2B and 2C). We cannot fully attribute the cardioprotective effect of Prolame to the cGMP-dependent pathway, as this compound might exert additional effects to those related with NO signaling. The PI3K/Akt salvage pathway activate other downstream mediators besides eNOS, like the Glycogen Synthase Kinase 3 Beta, PKC-ε and the mitochondrial ATP-dependent potassium channel which regulates the opening of the mPTP (Ong et al., 2015).

In conclusion, we found that the cardioprotective effects of Prolame and SNAP are related with NO production and with diminution of caspases and calpain-1 activities. However, even that our results suggest the activation of S-nitrosylation in the cardioprotection conferred by NO to reperfused hearts, there was no a correlative link between the activity of the proteases and such process. We cannot discard that other proteins relevant to heart function might be modified by S-nitrosylation, neither dismiss the contribution of alternative mechanisms in the observed cardioprotection.
ACKNOWLEDGEMENTS

We greatly acknowledge the generous gift of Prolame from Dr. José Manuel Fernández. (Instituto de Química, Universidad Nacional Autónoma de México).

ADDITIONAL INFORMATION AND DECLARATIONS

Funding

This work was supported by Grant 183363 to Cecilia Zazueta from Consejo Nacional de Ciencia y Tecnología, México. Nadia Giovanna Román-Anguiano is a doctoral student from Programa de Doctorado en Ciencias Biomédicas, Universidad Nacional Autónoma de México (UNAM) and received fellowship 330150 from Consejo Nacional de Ciencia y Tecnología (CONACYT). There was no additional external funding received for this study. The funders had no role in study design, data collection and analysis, decision to publish, or preparation of the manuscript.

Grant Disclosures

The following grant information was disclosed by the authors:
Consejo Nacional de Ciencia y Tecnología, México: 183363.
Programa de Doctorado en Ciencias Biomédicas.
Universidad Nacional Autónoma de México (UNAM).
Consejo Nacional de Ciencia y Tecnología (CONACYT).

Competing Interests

The authors declare that they have no competing interests.

Author Contributions

- Nadia Giovanna Román-Anguiano conceived and designed the experiments, performed the experiments, analyzed the data, prepared figures and/or tables, authored or reviewed drafts of the paper, approved the final draft.
- Francisco Correa conceived and designed the experiments, performed the experiments, approved the final draft.
- Agustina Cano-Martínez performed the experiments, contributed reagents/materials/analysis tools, prepared figures and/or tables, approved the final draft.
- Aurora de la Peña-Díaz contributed reagents/materials/analysis tools, approved the final draft.
- Cecilia Zazueta conceived and designed the experiments, performed the experiments, analyzed the data, contributed reagents/materials/analysis tools, prepared figures and/or tables, authored or reviewed drafts of the paper, approved the final draft.

Animal Ethics

The following information was supplied relating to ethical approvals (i.e., approving body and any reference numbers):

The investigation was approved by the Ethics Committee of the National Institute of Cardiology, "Ignacio Chávez" (14cb09012016).
Data Availability
The following information was supplied regarding data availability:
The raw measurements are available in Data S1.

Supplemental Information
Supplemental information for this article can be found online at http://dx.doi.org/10.7717/peerj.7348#supplemental-information.

REFERENCES
Aguilar HI, Botla R, Arora AS, Bronk SF, Gores GF. 1996. Induction of the mitochondrial permeability transition by protease activity in rats: a mechanism of hepatocyte necrosis. Gastroenterology 110(2):558–566 DOI 10.1053/gast.1996.v110.pm8566604.

Bolli R. 2001. Cardioprotective function of inducible nitric oxide synthase and role of nitric oxide in myocardial ischemia and preconditioning: an overview of a decade of research. Journal of Molecular and Cellular Cardiology 33(11):1897–1918 DOI 10.1006/jmcc.2001.1462.

Čarnická S, Adamová A, Nemčeková M, Matejíková J, Pancza D, Ravingerová T. 2011. Distinct effects of acute pretreatment with lipophilic and hydrophilic statins on myocardial stunning, arrhythmias and lethal injury in the rat heart subjected to ischemia/reperfusion. Physiological Research 5:825–830.

Chang AH, Sancheti H, García J, Kaplowitz N, Cadenas E, Han D. 2014. Respiratory substrates regulate S-Nitrosylation of mitochondrial proteins through a thiol-dependent pathway. Chemical Research in Toxicology 27:794–804 DOI 10.1021/tx400462r.

Chapman JG, Magee WP, Stukenbrok HA, Beckius GE, Milici AJ, Tracey WR. 2002. A novel nonpeptidic caspase-3/7 inhibitor, (S)-(+) 5-[(1-(2-methoxymethylpyrrolidinyl)sulfonyl]isatin reduces myocardial ischemic injury. European Journal of Pharmacology 456(1–3):59–68 DOI 10.1016/S0014-2999(02)02484-6.

Chen M, He H, Zhan S, Krajewski S, Reed JC, Gottlieb RA. 2001. Bid is cleaved by calpain to an active fragment in vitro and during myocardial ischemia/reperfusion. Journal of Biological Chemistry 276(33):30724–30728 DOI 10.1074/jbc.M103701200.

Chen Q, Paillard M, Gomez L, Ross T, Hu Y, Xu A, Lesnfsky EJ. 2011. Activation of mitochondrial m-calpain increases AIF cleavage in cardiac mitochondria during ischemia–reperfusion. Biochemical and Biophysical Research Communications 415(4):533–538 DOI 10.1016/j.bbrc.2011.10.037.

Correa F, Buelna-Chontal M, Chagoya V, García-Rivas G, Vigueras RM, Pedraza-Chaverri J, García-Niño WR, Hernández-Pando R, León-Contreras JC, Zazueta C. 2015. Inhibition of the nitric oxide/cyclic guanosine monophosphate pathway limited the cardioprotective effect of post-conditioning in hearts with apical myocardial infarction. European Journal of Pharmacology 765:472–481 DOI 10.1016/j.ejphar.2015.09.018.

Costa ADT, Garlid KD. 2008. Intramitochondrial signaling: interactions among mitoKATP, PKCε, ROS, and MPT. American Journal of Physiology-Heart and Circulatory Physiology 295(2):H874–H882 DOI 10.1152/ajpheart.01189.2007.

Das A, Xi L, Kukreja RC. 2005. Phosphodiesterase-5 Inhibitor sildenafil preconditions adult cardiac myocytes against necrosis and apoptosis. Journal of Biological Chemistry 280(13):12944–12955 DOI 10.1074/jbc.M404706200.

Das A, Xi L, Kukreja RC. 2008. Protein kinase G-dependent cardioprotective mechanism of phosphodiesterase-5 inhibition involves phosphorylation of ERK and GSK3β. Journal of Biological Chemistry 283(43):29572–29585 DOI 10.1074/jbc.M801547200.
Dunne KA, Allam A, Macintosh AM, Houston SA, Cerovic V, Goodyear CS, Roe AJ, Beatson SA, Milling SW, Walker D, Wall DM. 2013. Increased S-nitrosylation and proteasomal degradation of caspase-3 during infection contribute to the persistence of adherent invasive Escherichia coli (AIEC) in immune cells. *PLOS ONE* 8(7):e68386 DOI 10.1371/journal.pone.0068386.

Fernández-G JM, Rubio-Arroyo MF, Soriano-García N, Toscano RA, Pérez-César MC, Rubio-Póo C, Mandoki JJ, De la Peña A, Lemini C, Mendoza-Patiño N, Cruz F. 1985. Synthesis and molecular structure of prolame, N-(3-hydroxy-1,3,5(10)-estratrien-17β-yl)-3-hydroxypropylamine; an amino-estrogen with prolonged anticoagulant and brief estrogenic effects. *Steroids* 45(2):151–157 DOI 10.1016/0039-128X(85)90044-3.

Ford WR, Clanachan AS, Hiley CR, Jugdutt BI. 2001. Angiotensin II reduces infarct size and has no effect on post-ischaemic contractile dysfunction in isolated rat hearts. *British Journal of Pharmacology* 134(1):38–45 DOI 10.1038/sj.bjp.0704225.

Gao G, Dou QP. 2000. N-Terminal cleavage of bax by calpain generates a potent proapoptotic 18-kDa fragment that promotes Bcl-2-independent cytochrome c release and apoptotic cell death. *Journal of Cellular Biochemistry* 80(1):53–72 DOI 10.1002/1097-4644(20010101)80:1<53::aid-jcb60>3.0.co;2-e.

Giuffrè A, Sarti P, D’Itri E, Buse G, Soulimane T, Brunori M. 1996. On the mechanism of inhibition of cytochrome c oxidase by nitric oxide. *Journal of Biological Chemistry* 271(52):33404–33408 DOI 10.1074/jbc.271.52.33404.

González DR, Beigi F, Treuer AV, Hare JM. 2007. Deficient ryanodine receptor S-nitrosylation increases sarcoplasmic reticulum calcium leak and arrhythmogenesis in cardiomyocytes. *Proceedings of the National Academy of Sciences of the United States of America* 104(51):20612–20617 DOI 10.1073/pnas.0706796104.

González DR, Fernández LC, Ordenes PP, Treuer AV, Eller G, Boric MP. 2008. Differential role of S-nitrosylation and the NO–cGMP–PKG pathway in cardiac contractility. *Nitric Oxide* 18(3):157–167 DOI 10.1016/j.niox.2007.09.086.

González DR, Treuer A, Sun QA, Stamler JS, Hare JM. 2009. S-nitrosylation of cardiac ion channels. *Journal of Cardiovascular Pharmacology* 54(3):188–195 DOI 10.1097/fjc.0b013e3181b72c9f.

Gottlieb RA, Burleson KO, Kloner RA, Babior BM, Engler RL. 1994. Reperfusion injury induces apoptosis in rabbit cardiomyocytes. *Journal of Clinical Investigation* 94(4):1621–1628 DOI 10.1172/JCI117504.

Hammond J, Balligand JL. 2012. Nitric oxide synthase and cyclic GMP signaling in cardiac myocytes: from contractility to remodeling. *Journal of Molecular and Cellular Cardiology* 52(2):330–340 DOI 10.1016/j.yjmcc.2011.07.029.

Hausenloy DJ, Yellon DM. 2006. Survival kinases in ischemic preconditioning and postconditioning. *Cardiovascular Research* 70(2):240–253 DOI 10.1016/j.cardiores.2006.01.017.

Hernández-Reséndiz S, Palma-Flores C, De los Santos S, Román-Anguiano NG, Flores M, De la Peña A, Flores PL, Fernández-G JM, Coral-Vázquez RM, Zazueta C. 2015. Reduction of no-reflow and reperfusion injury with the synthetic 17β-aminoestrogen compound Prolame is associated with PI3K/Akt/eNOS signaling cascade. *Basic Research in Cardiology* 110(2):1–12 DOI 10.1007/s00395-015-0464-y.

Hu H, Wang J, Zhu H, Wu X, Zhou L, Song Y, Zhu S, Hao M, Liu C, Fan Y, Wang Y, Li Q. 2016. Ischemic postconditioning protects the heart against ischemia–reperfusion injury via neuronal nitric oxide synthase in the sarcoplasmic reticulum and mitochondria. *Cell Death & Disease* 7(5):e2222 DOI 10.1038/cddis.2016.108.
Ikeda H, Takajo Y, Murohara T, Ichiki K, Adachi H, Haramaki N, Katoh A, Imaizumi T. 2000. Platelet-derived nitric oxide and coronary risk factors. Hypertension 35(4):904–907 DOI 10.1161/01.HYP.35.4.904.

Inserte J, Hernando V, Ruiz-Meana M, Poncelas-Nozal M, Fernández C, Aguillo L, Sartorio C, Vilardosa U, García-Dorado D. 2014. Delayed phospholamban phosphorylation in post-conditioned heart favours Ca$^{2+}$ normalization and contributes to protection. Cardiovascular Research 103(4):542–553 DOI 10.1093/cvr/cvu163.

Iwamoto H, Miura T, Okamura T, Shirakawa K, Iwatate M, Kawamura S, Tatsuno H, Ikeda Y, Matsuzaki M. 1999. Calpain inhibitor-1 reduces infarct size and DNA fragmentation of myocardium in ischemic/reperfused rat heart. Journal of Cardiovascular Pharmacology 33(4):580–586 DOI 10.1097/00005344-199904000-00010.

Jones SP, Greer JJM, Kakkar AK, Ware PD, Turnage RH, Hicks M, Haperen RV, De Crom R, Kawashima S, Yokoyama M, Lefer DJ. 2004. Endothelial nitric oxide synthase overexpression attenuates myocardial reperfusion injury. American Journal of Physiology-Heart and Circulatory Physiology 286(1):H276–H282 DOI 10.1152/ajpheart.00129.2003.

Kim YM, Kim TH, Chung HT, Talanian RV, Yin XM, Billiar TR. 2000. Nitric oxide prevents tumor necrosis factor α–induced rat hepatocyte apoptosis by the interruption of mitochondrial apoptotic signaling through S-nitrosylation of caspase-8. Hepatology 32(4):770–778 DOI 10.1053/jhep.2000.18291.

Kohli V, Gao W, Camargo CA, Clavien PA. 1997. Calpain is a mediator of preservation-reperfusion injury in rat liver transplantation. Proceedings of the National Academy of Sciences of the United States of America 94(17):9354–9359 DOI 10.1073/pnas.94.17.9354.

Kovacs P, Bak I, Szendrei L, Vecdernees M, Varga E, Blasig IE, Tosaki A. 2001. Non-specific caspase inhibition reduces infarct size and improves post-ischaemic recovery in isolated ischaemic/reperfused rat hearts. Naunyn-Schmiedeberg’s Archives of Pharmacology 364(6):501–507 DOI 10.1007/s002100100483.

Lai YC, Pan KT, Chang GF, Hsu CH, Khoo KH, Hung CH, Jiang YJ, Ho FM, Meng TC. 2011. Nitrite-mediated S-nitrosylation of caspase-3 prevents hypoxia-induced endothelial barrier dysfunction. Circulation Research 109(12):1375–1386 DOI 10.1161/CIRCRESAHA.111.256479.

Li J, Bombeck CA, Yang S, Kim Y, Billiar TR. 1999. Nitric oxide suppresses apoptosis via interrupting caspase activation and mitochondrial dysfunction in cultured hepatocytes. Journal of Biological Chemistry 274(24):17325–17333 DOI 10.1074/jbc.274.24.17325.

Loscalzo J, Welch S. 1995. Nitric oxide and its role in the cardiovascular system. Progress in Cardiovascular Diseases 38(2):87–104 DOI 10.1016/S0033-0620(05)80001-5.

Lowry OH, Rosebrough NJ, Farr AL, Randall RJ. 1951. Protein measurement with the folin phenol reagent. Journal of Biological Chemistry 193:265–275.

Maejima Y, Adachi S, Morikawa K, Ito H, Isobe M. 2005. Nitric oxide inhibits myocardial apoptosis by preventing caspase-3 activity via S-nitrosylation. Journal of Molecular and Cellular Cardiology 38(1):163–174 DOI 10.1016/j.yjmcc.2004.10.012.

Mannick JB, Hausladen A, Liu L, Hess DT, Zeng M, Miao QX, Kane LS, Gow AJ, Stamler JS. 1999. Fas-induced caspase denitrosylation. Science 284(5414):651–654 DOI 10.1126/science.284.5414.651.

Mannick JB, Schonhoff C, Papeta N, Ghaforifar P, Szibor M, Fang K, Gaston B. 2001. S-nitrosylation of mitochondrial caspases. Journal of Cell Biology 154(6):1111–1116 DOI 10.1083/jcb.200104008.

Moncada S, Palmer RMJ, Higgs EA. 1991. Nitric oxide: physiology, pathology, and pharmacology. Pharmacological Reviews 43:109–142.
Neuhof C, Neuhof H. 2014. Calpain system and its involvement in myocardial ischemia and reperfusion injury. *World Journal of Cardiology* 6(7):638–652 DOI 10.4330/wjcv.v6.i7.638.

Ong SB, Dongworth RK, Cabrera-Fuentes HA, Hausenloy DJ. 2015. Role of the MPTP in conditioning the heart—translatability and mechanism. *British Journal of Pharmacology* 172(8):2074–2084 DOI 10.1111/bph.13013.

Patterson C, Portbury A, Schisler JC, Willis MC. 2011. Tear me down: role of calpain in the development of cardiac ventricular hypertrophy. *Circulation Research* 109(4):453–462 DOI 10.1161/CIRCRESAHA.110.239749.

Penna C, Tullio F, Femmino S, Rocca C, Angelone T, Cerra MC, Gallo MP, Gesmundo I, Fanciulli A, Brizzi MF, Pagliaro P, Alloatti G, Granata R. 2017. Obestatin regulates cardiovascular function and promotes cardioprotection through the nitric oxide pathway. *Journal of Cellular and Molecular Medicine* 21(12):3670–3678 DOI 10.1111/jcmm.13277.

Poderoso JJ, Carreras MC, Lisdero C, Riobó N, Schöpfer F, Boveris A. 1996. Nitric oxide inhibits electron transfer and increases superoxide radical production in rat heart mitochondria and submitochondrial particles. *Archives of Biochemistry and Biophysics* 328(1):85–92 DOI 10.1006/abbi.1996.0146.

Potz BA, Sabe AA, Abid MR, Sellke FW. 2015. Calpains and coronary vascular disease. *Circulation Journal* 80(1):4–10 DOI 10.1253/circj.CJ-15-0997.

Ramírez-Camacho I, Bautista-Pérez R, Correa F, Buelna-Chontal M, Román-Anguiano NG, Medel-Franco M, Medina-Campos ON, Pedraza-Chaverri J, Cano-Martinez A, Zazueta C. 2016. Role of sphingomyelinase in mitochondrial ceramide accumulation during reperfusion. *Biochimica et Biophysica Acta (BBA)—Molecular Basis of Disease* 1862(10):1955–1963 DOI 10.1016/j.bbadis.2016.07.021.

Ruetten H, Badorff C, Ihling C, Zeiher AM, Dimmeler S. 2001. Inhibition of caspase-3 improves contractile recovery of stunned myocardium, independent of apoptosis-inhibitory effects. *Journal of the American College of Cardiology* 38(7):2063–2070 DOI 10.1016/S0735-1097(01)01670-9.

Saligrama PT, Fortner KA, Secinaro MA, Collins CC, Rusell JQ, Budd RC. 2014. IL-15 maintains T-cell survival via S-nitrosylation-mediated inhibition of caspase-3. *Cell Death & Differentiation* 21(6):904–914 DOI 10.1038/cdd.2014.10.

Spinelli V, Sartiani L, Mugelli A, Romanelli MN, Cerbai E. 2018. Hyperpolarization-activated cyclic-nucleotide-gated channels: pathophysiological, developmental, and pharmacological insights into their function in cellular excitability. *Canadian Journal of Physiology and Pharmacology* 96(10):977–984 DOI 10.1139/cjpp-2018-0115.

Stamler JS, Lamas S, Fang FC. 2001. Nitrosylation: the prototypic redox based signaling mechanism. *Cell* 106(6):675–683 DOI 10.1016/S0092-8674(01)00495-0.

Sun J, Aponte AM, Kohr MJ, Tong G, Steenbergen C, Murphy E. 2013. Essential role of nitric oxide in acute ischemic preconditioning: S-nitrosylation vs. sGC/cGMP/PKG signaling? *Free Radical Biology and Medicine* 54:105–112 DOI 10.1016/j.freeradbiomed.2012.09.005.

Sun J, Morgan M, Shen R, Steenbergen C, Murphy E. 2007. Preconditioning results in S-nitrosylation of proteins involved in regulation of mitochondrial energetics and calcium transport. *Circulation Research* 101(11):1155–1163 DOI 10.1161/CIRCRESAHA.107.155879.
Sun J, Nguyen T, Aponte AM, Menazza S, Kohr MJ, Roth DM, Patel H, Murphy E, Steenbergen C. 2015. Ischaemic preconditioning preferentially increases protein S-nitrosylation in subsarcolemmal mitochondria. *Cardiovascular Research* 106(2):227–236 DOI 10.1093/cvr/cvv044.

Takimoto E. 2012. Cyclic GMP-dependent signaling in cardiac myocytes. *Circulation Journal* 76(8):1819–1825 DOI 10.1253/circj.CJ-12-0664.

Tao L, Gao E, Bryan NS, Qu Y, Liu H, Hu A, Christopher TA, Lopez BL, Yodoi J, Koch WJ, Feelisch M, Ma XL. 2004. Cardioprotective effects of thioredoxin in myocardial ischemia and reperfusion: role of S-nitrosation. *Proceedings of the National Academy of Sciences of the United States of America* 101(31):11471–11476 DOI 10.1073/pnas.0402941101.

Thompson J, Hu Y, Lesnefsky EJ, Chen Q. 2016. Activation of mitochondrial calpain and increased cardiac injury: beyond AIF release. *American Journal of Physiology-Heart and Circulatory Physiology* 310(3):H376–H384 DOI 10.1152/ajpheart.00748.2015.

Totzeck M, Korste S, Miinalainen I, Hendgen-Cotta UB, Rassaf T. 2017. S-nitrosation of calpains is associated with cardioprotection in myocardial I/R injury. *Nitric Oxide* 67:68–74 DOI 10.1016/j.niox.2017.04.003.

Vakeva AP, Agah A, Rollins SA, Matis LA, Li L, Stahl GL. 1998. Myocardial infarction and apoptosis after myocardial ischemia and reperfusion: role of the terminal complement components and inhibition by anti-C5 therapy. *Circulation* 97(22):2259–2267 DOI 10.1161/circ.97.22.2259.

Yang B, Jain S, Pawluczyk IZA, Imtiaz S, Bowley L, Ashra SB, Nicholson ML. 2005. Inflammation and caspase activation in long-term renal ischemia/reperfusion injury and immunosuppression in rats. *Kidney International* 68(5):2050–2067 DOI 10.1111/j.1523-1755.2005.00662.x.

Yang NC, Lu LH, Kao YK, Chau IY. 2004. Heme oxygenase-1 attenuates interleukin-1β-induced nitric oxide synthase expression in vascular smooth muscle cells. *Journal of Biomedical Science* 11(6):799–809 DOI 10.1159/000081827.

Zhang D, Zhao N, Ma B, Wang Y, Zhang G, Yan X, Hu X, Xu T. 2016. Pro-caspase-9 induces its cleavage by transnitrosylating XIAP via the thioredoxin system during cerebral ischemia-reperfusion in rats. *Scientific Reports* 6(1):1–12 DOI 10.1038/srep24203.

Zhao CY, Greenstein LL, Winslow RL. 2017. Mechanisms of the cyclic nucleotide cross-talk signaling network in cardiac L-type calcium channel regulation. *Journal of Molecular and Cellular Cardiology* 106:29–44 DOI 10.1016/j.yjmcc.2017.01.013.

Zhao X, He G, Chen YR, Pandian RP, Kuppusamy P, Zweier JL. 2005. Endothelium-derived nitric oxide regulates postischemic myocardial oxygenation and oxygen consumption by modulation of mitochondrial electron transport. *Circulation* 111(22):2966–2972 DOI 10.1161/CIRCULATIONAHA.104.527226.

Zhao ZQ, Nakamura M, Wang NP, Wilcox JN, Shearer S, Ronson RS, Guyton RA, Vinten-Johansen J. 2000. Reperfusion induces myocardial apoptotic cell death. *Cardiovascular Research* 45(3):651–660 DOI 10.1016/S0008-6363(99)00354-5.