Ischaemic preconditioning improves proteasomal activity and increases the degradation of δPKC during reperfusion

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Aims
The response of the myocardium to an ischaemic insult is regulated by two highly homologous protein kinase C (PKC) isozymes, δ and εPKC. Here, we determined the spatial and temporal relationships between these two isozymes in the context of ischaemia/reperfusion (I/R) and ischaemic preconditioning (IPC) to better understand their roles in cardioprotection.

Methods and results
Using an ex vivo rat model of myocardial infarction, we found that short bouts of ischaemia and reperfusion prior to the prolonged ischaemic event (IPC) diminished δPKC translocation by 3.8-fold and increased εPKC accumulation at mitochondria by 16-fold during reperfusion. In addition, total cellular levels of δPKC decreased by 60 ± 2.7% in response to IPC, whereas the levels of εPKC did not significantly change. Prolonged ischaemia induced a 48 ± 11% decline in the ATP-dependent proteasomal activity and increased the accumulation of misfolded proteins during reperfusion by 192 ± 32%; both of these events were completely prevented by IPC. Pharmacological inhibition of the proteasome or selective inhibition of εPKC during IPC restored δPKC levels at the mitochondria while decreasing εPKC levels, resulting in a loss of IPC-induced protection from I/R. Importantly, increased myocardial injury was the result, in part, of restoring a δPKC-mediated I/R pro-apoptotic phenotype by decreasing pro-survival signalling and increasing cytochrome c release into the cytosol.

Conclusion
Taken together, our findings indicate that IPC prevents I/R injury at reperfusion by protecting ATP-dependent 26S proteasomal function. This decreases the accumulation of the pro-apoptotic kinase, δPKC, at cardiac mitochondria, resulting in the accumulation of the pro-survival kinase, εPKC.

Keywords
Cardioprotection • Ischaemia/reperfusion • Apoptosis • Proteasome • PKC • Ischaemic preconditioning

1. Introduction
Myocardial ischaemia and reperfusion (I/R)-induced damage is associated with both apoptotic and necrotic cell death.1 We have shown this to be dependent on translocation of delta protein kinase C (δPKC) to cardiac mitochondria where it inhibits mitochondrial function.2–4 Inhibition of δPKC with the selective peptide inhibitor, δV1–1, protects the heart from ischaemic injury.4,5 Ischaemic preconditioning (IPC) observed in animals and humans6–8 protects the heart from reperfusion injury by activating
pro-survival kinases, preventing apoptosis and necrosis, preserving mitochondrial function, and reducing ROS generation. Many of these effects are afforded, at least in part, through activation and translocation of εPKC to cardiac mitochondria, resulting in diminished apoptosis and necrosis.

Interestingly, activation of δPKC is also required for both opioid and IPC-mediated protection. We showed that activation of δPKC is cardioprotective provided there is sufficient time allowed for εPKC activation. Furthermore, εPKC is activated by ROS during IPC, whereas δPKC plays a role in ROS generation. Therefore, although both PKC isozymes play a role in IPC, the mechanism by which the pro-survival kinase (εPKC) and the pro-death kinase (δPKC) interact is not known. Here, we present evidence of a novel mechanism in which the proteasome alters the ratio between δPKC and εPKC, thereby regulating myocardial viability following I/R.

2. Methods

2.1 Materials

All antibodies were from Santa Cruz Biotechnology (Santa Cruz, CA). Lactacystin, LLVY-AMC, epoxomycin, and MG-132 were from Biomol (Plymouth Meeting, PA). εPKC and δPKC conjugated to TAT were made by Anaspec, San Jose, CA. This study conforms with the Guide for the Care and Use of Laboratory Animals of Stanford University.

2.2 Isolated perfused rat heart model and measurement of tissue necrosis

All procedures were carried out as described. All animal protocols were approved by the Institutional Animal Care and Use Committee of Stanford University.

2.3 Cellular fractionation and western blotting

Isolated hearts were homogenized in 210 mM mammalian, 70 mM sucrose, 1 mM EDTA, and 5 mM MOPS, pH 7.4. After filtering through cheesecloth and a 5 min centrifugation at 8000 g, the supernatant was centrifuged (10,000 g; 10 min) to obtain the mitochondrial pellet and the cytosolic extract (supernatant). This technique provides a mitochondrial fraction with only traces of sarcomemal and plasma membrane contamination. Western blot analysis used polyclonal εPKC, δPKC, p-Akt, Akt, and cytochrome c antibodies, was normalized to ANT (mitochondria) and GAPDH (total and cytosolic homogenates) and was expressed as percent control.

2.4 Assay of proteasome activity

Chymotrypsin-like activity of the proteasome was assayed using the fluorogenic peptide Suc-Leu-Leu-Val-Tyr-7-amido-4-methylcoumarin (25 μM, LLVY-MCA) in a microtiter plate (50 μg protein) with 200 μl of 10 mM MOPS, pH 7.4. Assays were carried out in the absence and presence of 2.5 mM ATP and 5.0 mM Mg²⁺ with the difference attributed to ATP-dependent proteasome activity. The rate of fluorescent product formation was measured with excitation and emission wavelengths of 350 and 440 nm, respectively. In order to block proteasome activity during the experimental protocol, 2.0 μM lactacystin was perfused during the preconditioning protocol and the first 10 min of reperfusion.

2.5 Slot blot analysis of cellular misfolded proteins

Heart tissue homogenate (25 μg protein) was normalized and slot blotted onto PVDF membrane (Millipore, Bedford, MA, USA) and membranes were washed three times with 0.05% Tween 20. 10 mM Tris, pH 7.5, 100 mM NaCl (T-TBS) and blocked in T-TBS + 5% milk. After 4 h of incubation with an anti-soluble oligomer antibody (Biosource International, Camarillo, CA), an antibody that recognizes misfolded proteins, proteins were visualized as in the western blot analysis. Sample loading was normalized by Ponceau staining.

2.6 Analysis of cellular ATP levels

ATP determination was carried out using the Molecular Probes luciferase-based ATP determination kit (Kit# A22066). In brief, 100 μg of protein was incubated in a 96-well plate with 50 μM luciferin and 1.25 μg/ml luciferase in a Tris-based 1X reaction buffer containing DTT. ATP was measured after 5 min using a luminometer (560 nm at room temperature) using a standard curve of known ATP concentrations.

2.7 Statistics

Data are represented as the mean ± SEM, and significance was determined by one-way analysis of variance with a post-hoc Tukey test or a two-tailed t-test.

3. Results

3.1 IPC diminishes δPKC at the mitochondria and increases εPKC translocation

Since the mitochondrial sites of regulation by δ and εPKC during I/R, we first determined the levels of δ and εPKC in this fraction. Thirty minutes of ischemia and 60 min reperfusion (I30/R60) (Figure 1A) resulted in accumulation of δPKC (~6-fold; P < 0.001; n = 5) and εPKC (~9-fold; P < 0.05; n = 5) at the mitochondria (Figure 1B). However, after IPC, I/R-induced δPKC accumulation at the mitochondrial fraction was largely prevented, whereas εPKC translocation increased ~2-fold higher than hearts that were not preconditioned (P < 0.01; n = 5) (Figure 1B). Interestingly, IPC resulted in a seven-fold greater increase in εPKC at cardiac mitochondria relative to δPKC (Figure 1B). The IPC stimulus alone (without subsequent I30/R60) (Figure 1C left panel) increased the levels of both δPKC and εPKC and caused an ~2-fold increase in their mitochondrial levels relative to normoxic hearts (Figure 1C right panel; P < 0.05; n = 6).

3.2 Total cellular levels of δPKC are greatly diminished by IPC

To determine whether the changes in the mitochondrial levels of δ and εPKC reflect changes in the cellular levels of these kinases, we next determined the levels of both εPKC and δPKC in total cardiac extracts. I30/R60 alone did not affect the overall levels of δPKC or εPKC vs. normoxia (N; n = 4; Figure 2). However, after IPC followed by ischaemia, δPKC levels decreased by 33% (vs. N, P < 0.05; n = 4), whereas εPKC levels did not. Therefore, the reduction in δPKC translocation to the mitochondria (Figure 1B) appears to be associated with diminished protein levels, and this effect is selective and does not seem to affect εPKC translocation.
3.3 IPC-induced proteasomal degradation of δPKC diminished its translocation to cardiac mitochondria with a concurrent increase in εPKC translocation to this fraction

δPKC has been shown to be degraded by the 26S proteasome. Declines in ATP levels during ischaemia result in the disassembly of the 26S proteasome into the 20S form. In contrast, IPC reduces ischaemia-mediated declines in ATP levels. We therefore determined whether the loss in δPKC levels following IPC prior to I/R relates to preservation of the 26S proteasome activity. ATP Mg\(^{2+}\)-stimulated peptidase activity is a reflection of the relative level of the 26S proteasome. Ischaemia induced a 45% decline in ATP-dependent proteasomal activity (Figure 3B) that was associated with an ∼3-fold increase in the accumulation of misfolded proteins during reperfusion (Figure 3C). IPC prevented the...
Figure 3  Effect of preconditioning on ischaemia-induced loss in ATP-dependent proteasome activity. (A) Cytosolic extracts were prepared from hearts exposed to 70 min of normoxic perfusion (N), 30 min of ischaemia (I₃₀), or three cycles of preconditioning (5 min ischaemia and 5 min reperfusion) followed by 30 min of ischaemia (I₃₀ + IPC) in the absence or presence of the proteasome inhibitor lactacystin or the specific ePKC inhibitor eV₁₋₂. Chymotrypsin-like activity of the proteasome present in the cytoplasmic milieu was evaluated and the specific inhibitor MG-132 (20 μM) was utilized to ensure that measured activities were due to the proteasome (data not shown). The presence of unfolded proteins was evaluated using the slot blot technique with an anti-soluble oligomer antibody. Values representing ATP-dependent proteasome activity and misfolded proteins are presented as a percent of values obtained with samples from hearts exposed to 60 min of normoxic perfusion (N). Values represent the mean ± standard deviation (n = 4). (B) Ischaemia resulted in a 50% decline in ATP-dependent proteasome activity (P < 0.05 vs. N), which was completely reversed by IPC (P < 0.05 vs. I₃₀). The proteasome inhibitor, lactacystin (2 μM) and the specific ePKC inhibitor eV₁₋₂ both significantly decreased the activity of the proteasome (P < 0.05 vs. I₃₀R₆₀ + IPC; n = 4). (C) I₃₀R₆₀ resulted in an ~3-fold increase in misfolded proteins which was prevented by IPC (P < 0.05; n = 4). Treatment of hearts with lactacystin or eV₁₋₂ blocked the effect of IPC and increased the accumulation of misfolded proteins. (D) IPC elevated ATP levels by 3.5-fold in hearts that had undergone I₃₀R₆₀, and eV₁₋₂ blocked these effects. (E) Treatment with the ePKC activator (ψeRACK) protected the proteasome from ischaemia-mediated inhibition (P < 0.05 vs. I₃₀). *P < 0.05 vs. Normoxia, †P = 0.05 vs. I₃₀R₆₀, §P < 0.05 vs. I₃₀, ‡P < 0.05 vs. IPC + I₃₀, ††P < 0.05 vs. IPC + I₃₀R₆₀. Misfolded protein accumulation and proteasome activity were analysed by one-way analysis of variance with a post-hoc Tukey test. Figure 3D, proteasome activity was analysed by Student’s t-test.
Inhibition of the proteasome restores \(\delta\)PKC cellular and mitochondrial levels in IPC hearts with a resultant decrease in \(\epsilon\)PKC levels. (A) Hearts were hung in Langendorff mode and treated with the above-mentioned perfusion protocols. Hearts were then removed, homogenized, and the total homogenate and mitochondrial fractions were subjected to western blot analysis with antibodies against the proteins listed in the figure. Values were normalized to GAPDH (total homogenate) or ANT (mitochondrial fraction) and expressed as \% I30R60. IPC before prolonged ischaemia reduced total levels of \(\delta\)PKC by \(\sim 80\%\) (\(P < 0.05\) vs. I30R60). Inhibition of the proteasome with 2 \(\mu\)M lactacystin blocked \(\delta\)PKC degradation (\(P < 0.05\) vs. I30R60 + IPC). Similar to Figure 2, IPC before prolonged ischaemia did not significantly change overall levels of \(\epsilon\)PKC. However, inhibition of the proteasome increased \(\epsilon\)PKC levels by \(\sim 2\)-folds (\(P < 0.05\) vs. I30R60). Inhibition of \(\epsilon\)PKC activity with \(\epsilon\)V1–2 did not significantly change the overall levels of either \(\delta\) or \(\epsilon\)PKC isozymes (data not shown). (B) As in Figure 2, IPC before I30R60 decreased levels of \(\delta\)PKC at mitochondria by \(\sim 60\%\) (\(P < 0.05\) vs. I30R60). This was completely prevented in hearts treated with 2 \(\mu\)M lactacystin and 1 \(\mu\)M \(\epsilon\)V1–2 (\(P < 0.05\) vs. I30R60 + IPC). (C) Although \(\delta\)PKC mitochondrial levels were restored, \(\epsilon\)PKC levels were diminished by 40\% relative to I30R60 and by 60\% relative to I30R60 + IPC (\(P < 0.05\)). Hearts that were treated with 1 \(\mu\)M of a peptide inhibitor of \(\epsilon\)PKC (\(\epsilon\)V1–2) during the IPC protocol showed a significant decrease in \(\epsilon\)PKC mitochondrial levels (\(P < 0.05\) vs. I30R60 + IPC).

* * \(P < 0.05\) vs. I30R60, \(\dagger \) * * \(P < 0.05\) vs. IPC I30R60. Cellular and mitochondrial PKC levels were analysed by the one-way analysis of variance with a post-hoc testing by Tukey.
ischaemia-mediated declines in proteasomal activity and reduced the levels of misfolded proteins (Figure 3B and C; \( P < 0.05 \); \( n = 4 \)). Furthermore, ATP levels correlated with proteasomal activity; ATP levels diminished during ischaemia/reperfusion and IPC significantly prevented this decline (Figure 3D).

To determine if IPC increases proteasomal degradation of \( \delta PKC \), we perfused the proteasome inhibitor, lactacystin (2 \( \mu M \)), during IPC and the first 10 min of reperfusion (Figure 3A). Lactacystin significantly blocked the activity of the proteasome (~75%; \( P < 0.01 \); \( n = 4 \); Figure 3B) and increased the levels of misfolded proteins by ~7-fold (Figure 3C). In agreement with our findings in Figure 2, IPC reduced post-reperfusion cellular levels of \( \delta PKC \) by ~80% (\( P < 0.01 \); \( n = 4 \); Figure 4A). However, inhibition of the proteasome with lactacystin prevented the loss of \( \delta PKC \) after IPC and I\( _{30} \)R\( _{60} \) (Figure 4A). Although lactacystin treatment did not affect the levels of \( \varepsilon PKC \) relative to IPC, the ratio of \( \varepsilon PKC \) to \( \delta PKC \) decreased due to elevated levels of \( \delta PKC \).

Since lactacystin-mediated inhibition of the proteasome during IPC prevented \( \delta PKC \) degradation, it may promote translocation of \( \delta PKC \) to cardiac mitochondria. Indeed, lactacystin treatment completely blocked IPC-induced reductions in \( \delta PKC \) levels, restoring mitochondrial \( \delta PKC \) to levels obtained after I\( _{30} \)R\( _{60} \) alone (Figure 4B). Interestingly, in contrast to the effects of proteasome inhibition on total \( \varepsilon PKC \) levels, mitochondrial levels of \( \varepsilon PKC \) were reduced by 50% in the presence of lactacystin (Figure 4C). These data suggest that IPC inversely affects the ratios of these isoforms.

Figure 5: Inhibition of \( \delta PKC \) degradation restores the apoptotic phenotype seen during reperfusion. Hearts were hung in Langendorff mode and treated with the listed perfusion protocols. Hearts were then removed, homogenized, fractionated, and the cytosolic homogenate was subjected to western blot analysis with antibodies against the proteins listed in the figure. Values were normalized to GAPDH and expressed as % I\( _{30} \)R\( _{60} \). (A) IPC before prolonged ischaemia significantly decreased cytochrome c release into the cytosol (\( P < 0.05 \) vs. I\( _{30} \)R\( _{60} \)). Inhibition of the proteasome with 2 \( \mu M \) lactacystin restored cytochrome c release to levels seen during I\( _{30} \)R\( _{60} \) (\( P < 0.05 \) vs. I\( _{30} \)R\( _{60} \) + IPC). (B) Ischaemia alone and perfusion with lactacystin or \( \varepsilon V1\~2 \) alone did not result in significant release of cytochrome c into the cytosol. Additionally, as evidenced by a lack of mitochondrial VDAC in the cytosolic fraction, there was little contamination from mitochondrial cytochrome c in this fraction. Enolase was used as a cytosolic loading control. (C) IPC before prolonged ischaemia also increased phosphorylation of the pro-survival kinase, Akt, by ~3-fold (\( P < 0.05 \) vs. I\( _{30} \)R\( _{60} \)). Inhibition of the proteasome with 2 \( \mu M \) lactacystin decreased phosphorylation back to I\( _{30} \)R\( _{60} \) levels (\( P < 0.05 \) vs. I\( _{30} \)R\( _{60} \) + IPC). \*\( P < 0.05 \) vs. I\( _{30} \)R\( _{60} \); \( \beta \)\( P < 0.05 \) vs. IPC I\( _{30} \)R\( _{60} \). Cytosolic cytochrome c levels and p-Akt were analysed by one-way analysis of variance with a post-hoc Tukey test. 

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two PKC isozymes at cardiac mitochondria following I/R likely through the regulation of proteasome activity. Additionally, lactacytin abolished IPC-mediated protection of ATP levels following ischaemia/reperfusion (Figure 3D).

We determined whether εPKC indirectly downregulates δPKC by protecting proteasomal function during IPC. To this end, we perfused the specific εPKC inhibitor, εV1–2 (Figure 3A) and found an ~70% inhibition of proteasomal activity, an effect that was similar to that obtained by lactacytin (Figure 3B), and resulted in a corresponding increase in cellular misfolded proteins (Figure 3C). Similar to lactacytin, εV1–2 treatment significantly decreased εPKC levels at cardiac mitochondria (71% vs. IPC), and restored δPKC levels in this fraction (P < 0.05; n = 7) (Figure 4B and C). Perfusion with a specific εPKC activator, ƞεRACK, before ischaemia mimicked the IPC-mediated protective effect on proteasomal activity and prevented the loss in proteasomal activity seen during I/R (Figure 3E). This is likely an indirect effect, since εPKC was not found to associate with the proteasome following ƞεRACK treatment (data not shown).

3.4 Inhibition of the proteasome prevents Akt activation and increases release of cytochrome c during IPC

IPC activates the pro-survival kinases, Akt, and ERK1/2 and blocks cytochrome c release during reperfusion.10,15,35 In contrast, δPKC decreases Akt activation and increases cytochrome c release during I/R.5 Here we found that IPC significantly blocked I/R-mediated release of mitochondrial cytochrome c into the cytosol (60%; P < 0.05; n = 4) (Figure 5A) and inhibition of the proteasome with lactacytin restored cytochrome c release to the levels seen during I/R (P < 0.05). Treatment of non-ischaemic hearts with either lactacytin or εV1–2 did not cause significant release of cytochrome c into the cytosol (Figure 5B). Additionally, IPC significantly increased the phosphorylation of the pro-survival kinase, Akt, over I/R (300%), and this was abolished by lactacytin treatment (Figure 5C). Li et al. showed that activation of δPKC reduces Akt phosphorylation whereas inhibition of δPKC increased Akt phosphorylation. They suggested that δPKC-mediated inhibition of Akt proceeds through increased association of protein phosphatase 2A.36 Neither IPC nor lactacytin treatments significantly changed the phosphorylation levels of ERK1/2 (not shown), consistent with the findings of other studies.37,38

3.5 Inhibition of the proteasome during IPC increases tissue injury

Since pharmacological inhibition of the proteasome during IPC restored the apoptotic phenotype, we determined if tissue injury is altered by proteasome inhibition. As reported, IPC decreased both creatine phosphokinase (CPK) release and tetrazolium tetra-chloride (TTC) staining of the myocardium by ~70 and 60%, respectively (Figure 6A and B) and Lactacytin reversed the benefits of IPC-mediated protection. As we found before in isolated myocytes,28 in addition to the effects of εV1–2 on proteasomal function and δPKC translocation, inhibition of εPKC also completely reversed the protective effects of IPC on the myocardium (Figure 6A and B). Finally, to confirm the effects of lactacytin, we utilized another highly selective inhibitor of the proteasome, epoxomicin. Similar to lactacytin, inhibition of the proteasome with epoxomicin (2 μM) abolished the cardioprotective effects of IPC (data not shown).

4. Discussion

Our data suggest that IPC-induced decreases in mitochondrial δPKC levels are due to decreased total levels of δPKC. We also show that IPC prevents ischaemia-mediated declines in the 26S ATP-dependant proteasomal activity and that this is associated with diminished accumulation of cellular misfolded proteins. Ischaemia-mediated declines in forebrain ATP levels promote dissociation of the 26S proteasome (the form responsible for δPKC degradation)29) to the 20S proteasome.32 During I/R, (in the absence of preconditioning) the significant decrease in ATP-dependent proteasomal activity is therefore likely due to decreased ATP levels within the cells. Indeed, as has been shown before,4,39 and here in an ex vivo model of ischaemia/reperfusion, ATP levels significantly declined during ischaemia/reperfusion and IPC significantly prevented this decline (Figure 3D). Alternatively, modifications by lipid peroxidation products and accumulation of oxidized proteins during I/R may also act as inhibitors of proteasomal function.40 Inhibition of the proteasome with lactacytin or epoxomicin blocked the protective effects of IPC. Additionally, lactacytin treatment elevated PKC is ubiquitinated within 30 min of activation41 and direct inhibition of the 26S proteasome with Bortezomib, a highly selective proteasome inhibitor currently in clinical use for the treatment of haematological cancers, increases mitochondrial ROS generation, cytochrome c release, and apoptosis associated with mitochondrial accumulation of δPKC.29,42 We suggest that since IPC and εPKC activation slow ATP depletion during prolonged ischaemia,8,43 and δPKC is likely activated by the IPC stimulus (Figure 1C) the 26S proteasomal activity is maintained leading to the degradation of pro-apoptotic and pro-necrotic, δPKC, thereby conferring cardiac protection. Although the most likely explanation for the decrease in δPKC levels in the mitochondria is a decrease in total level of this isoform in the cells (due to its increased degradation by the proteasome), we cannot exclude the possibility that decreased affinity of the binding site for δPKC in the mitochondria and post-translational modifications of the enzyme or its binding proteins also contribute to δPKC declined levels and therefore activity in the mitochondria. We have previously shown that accumulation of δPKC at cardiac mitochondria increases PDH phosphorylation and the inhibitor δV1–1 prevents this.5 Additionally, we have shown that εPKC is able to activate ALDH2 in hearts in the same in vivo model of I/R.44 Therefore, we have already provided direct evidence in this model of I/R that increased levels of PKC isozymes in the mitochondrial fraction are associated with increased phosphorylation of target substrates and hence reflect increased catalytic activity of these isozymes.
In addition to the protection afforded by degrading δPKC, increased εPKC accumulation at cardiac mitochondria is also likely to confer protection. εPKC translocates to mitochondria,16,19–21 where it prevents opening of the mitochondrial permeability transition pore,19,45 opens kATP channels,9 forms signalling complexes with MAPK,9 retards the reduction in cellular ATP levels,46 interacts with the electron transport chain,47 and augments mitochondrial function39 all of which contribute to cardioprotection. Since the relative level of εPKC at the mitochondria during reperfusion in the absence of the IPC stimulus is similar to δPKC levels, εPKC-mediated cardioprotection may be masked by the pro-apoptotic and pro-necrotic effects of δPKC during reperfusion. Administration of the εPKC activator, δRACK, prior to ischaemia, which mimics IPC and protects mitochondrial function39 prevented ischaemia-mediated declines in proteasome activity. Although recent studies suggest that kinases may regulate proteasome function directly,48 we did not find any physical association between εPKC and the proteasome.

In summary, activated δPKC has two potential fates that appear to depend on the metabolic state of the cell. If mitochondrial function, cellular energy status, and the integrity of the 26S proteasome are maintained, δPKC is efficiently degraded. In contrast, if mitochondrial function and ATP production are compromised, the ATP-dependent 26S proteasome activity is diminished, resulting in increased levels of activated δPKC at the mitochondria, where it participates in the induction of cell death. The proteasome can therefore be viewed as a sensor of cellular viability, determining the ratio of pro-apoptotic δPKC and pro-survival εPKC at the mitochondria and thus the ultimate fate of the cell. We propose the following mechanism. The decrease in ATP levels seen during I/R (Figure 3D) and increased generation of reactive oxygen species, will diminish 26S proteasome activity.40 δPKC is activated by ROS4 and also during the early stages of reperfusion, resulting in its accumulation at cardiac mitochondria (Figure 1).3 Because the activity of the proteasome is diminished (Figure 3), δPKC is not degraded, favouring its accumulation at cardiac mitochondria (Figure 4), where it triggers pro-apoptotic cytochrome c release and inactivation of Akt (Figure 5), leading to tissue injury (Figure 6). In contrast, IPC is associated with a small burst of mitochondrial ROS during the trigger phase of IPC, which decreases ROS generation during the effector phase49 and may also act as a stimulus for δPKC activation (Figure 1C). Diminished ROS generation and maintenance of cellular ATP levels (Figure 3D) result in protection of proteasomal function,50 which leads to degradation of δPKC (Figure 4A). Since both εPKC and δPKC accumulate at the mitochondria during I30R60 (Figure 1B) and since εPKC is not degraded during I30R60 (Figure 2A), degradation of δPKC during IPC tips the balance towards the accumulation of the pro-survival kinase, εPKC, at cardiac mitochondria, thus protecting mitochondrial function and proteasomal activity thereby diminishing I/R-mediated tissue injury.

Conflict of interest: D.M.-R. is the founder of KAI Pharmaceuticals. However, none of the work described in the study is based on or supported by the company.
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