Postconditioning Protection Against Myocardiocyte Anoxia/Reoxygenation Injury From Penehyclidine Hydrochloride

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Background/Aims: To investigate the postconditioning protective effect of penehyclidine hydrochloride (PHC) against anoxia/reoxygenation (A/R) injury in H9c2 cells along with the involved mechanism and timing effect.

Methods: We divided H9c2 cells into 7 groups: control group, A/R group and PHC+A/R groups at 0 min, 5 mins, 10 mins, 20 mins, 30 mins, respectively (treated with 0.1 μm/L PHC at 0 min, 5 mins, 10 mins, 20 mins, 30 mins after the reoxygenation procedure began). Cell apoptosis, oxidative stress, intracellular Ca\textsuperscript{2+} concentration, mitochondrial membrane potential and mitochondrial permeability transition pore (MPTP) opening were explored. Bcl-2, Bax, Cyt C, caspase-3 and caspase-9 levels were measured.

Results: A/R significantly increased both cell injury and cell apoptosis. PHC showed postconditioning protective effect by attenuating superoxide production, decreasing Ca\textsuperscript{2+} overload, restraining MPTP activities, restoring mitochondrial membrane potential, regulating cell apoptosis proteins and modulation of mitochondrial pathway. Earlier administration of PHC offered greater postconditioning protective effect.

Conclusion: H9c2 cells were protected by PHC from A/R injury regardless of timing of PHC administration (0 min, 5 mins, 10 mins, 20 mins, 30 mins). However, earlier administration of PHC resulted in better PHC postconditioning protection.

Keywords: penehyclidine hydrochloride, PHC, anoxia/reoxygenation injury, A/R injury, cell apoptosis, oxidative stress, mitochondrial pathway

Introduction

Ischemic heart disease is one of the most severe cardiovascular diseases with high morbidity and mortality. The most effective means of minimizing myocardial ischemic injury is early reperfusion of ischemic tissues. However, myocardial reperfusion has its own inherent risks; alternations in myocardial metabolic, electrophysiologic and cellular functions are not uncommon. This is referred to as ischemia-reperfusion injury (IRI).\textsuperscript{1-3} For the past decade, researchers have explored ways of reducing IRI, including preconditioning and postconditioning. In order to understand the chain of events that ultimately result in IRI, as well as the mechanism from which IRI could be protected by postconditioning treatment, we designed an in vitro experiment using H9c2 cell models through A/R procedures.

Penehyclidine Hydrochloride (PHC) is a selective anticholinergic agent launched by the Institute of Pharmacology and Toxicology, a division within the Chinese Academy of Military Medical Sciences. It was found to have a protective
effect against myocardial ischemia-reperfusion injury while having minimal cardiovascular side effects.\(^4\) Our previous studies demonstrated that PHC offered reperfusion protection by decreasing calcium overload, minimizing the formation of reactive oxygen species, impeding mitochondrial MPTP opening and reducing inflammatory response in vitro and in vivo. We previously determined that the most effective concentration was 0.1 \(\mu\)m/L.\(^5,6\) This current investigation focused on the effect of timing that PHC had on postconditioning. We also designed this in vitro experiment to further elucidate the mechanism of A/R injury in the early reoxygenation period. We specifically looked at whether earlier administration of PHC provided significantly better protection against cellular ischemia-reperfusion injury. We hope our results will lead to clinical applications in the treatment of patients with ischemic heart disease and offer protection against reperfusion injury in surgical patients.

### Materials And Methods

H9c2 cell line was purchased from Shanghai Cell Library of China, which was originally obtained from ATCC, Manassas, VA, USA. The cells were cultured at 37°C in a chamber consisting of 5% CO\(_2\)/95% air. When cells' density reached 90%, the cells were harvested for our experiment.

H9c2 cells were divided into seven groups, as described in Figure 1A:

1. **Control group**: H9c2 cells were first incubated in Tyrode solution followed by cultivation in a chamber with 5% CO\(_2\)/95% air mixture for 3 hrs. The culture medium of H9c2 cells was then quickly replaced by low-sugar DEME medium outside the chamber. The H9c2 cells were quickly returned to the cell chamber consisting of 5% CO\(_2\)/95% air for 2 hrs.

2. **A/R group**: H9c2 cells were incubated in Tyrode solution with a gas mixture of 5% CO\(_2\)/95% N\(_2\) for 3 hrs. The cell culture medium was then replaced with a low-sugar DEME solution and returned back to a gas chamber with a mixture of 5% CO\(_2\)/95% air at 37°C for 2 hrs of reperfusion treatment.

3. **PHC postconditioning group**: Previous steps were the same as in the A/R groups, except with low-sugar DEME containing 0.1 \(\mu\)m/L PHC replacing low-sugar DEME without PHC at 0 min, 5 mins, 10 mins, 20 mins and 30 mins, respectively, after commencement of reoxygenation. (0 min indicated the time during which Tyrode solution was directly replaced by PHC solution when the anoxia period stopped.) Next, low-sugar DEME solution containing no PHC replaced the PHC solution after a 90 mins treatment period. The total anoxia period was 3 hrs with a reoxygenation period of 2 hrs in both the A/R group and PHC postconditioning group. The total exposure time to PHC in the PHC postconditioning group was 90 mins. Reoxygenation was carried out in a mixture of 5% CO\(_2\)/95% air at 37°C. H9c2 cells and the upper fluid in each group were collected immediately after the reoxygenation period.

### LDH And CK Activity Assay

LDH and CK are released from cytoplasm upon cellular injury and death. As such, the presence of LDH and CK in medium supernatant can be used as biomarkers of cellular injury or death.\(^7,8\) At the end of the experiment, the culture supernatant in each group was collected and subjected to spectrophotometric analysis (Yoke, Shanghai, China) to detect the presence of LDH absorbance at 450 nm and CK absorbance at 660 nm, respectively.\(^9\) Each measurement was performed six times.

### Cell Viability Assays

At the end of this experiment, 0.2 \(\mu\)L MTT was added to each well. H9c2 cells were treated with MTT at 37°C for 4 hrs. 150 \(\mu\)L dimethyl sulfoxide (DMSO) would then replace the MTT medium. DMSO was used to dissolve the purple formazan crystals produced by viable cells into an aqueous solution. The optical density at 570 nm wavelength of this
purple solution was analyzed by a microplate reader (BioTek, Winooski, Vermont, USA). Each assay was performed six times. The mean result was expressed as a ratio of the OD values of the experimental group to the control group.

**Annexin-V/PI Assay**

Fluorescein isothiocyanate (FITC)-Annexin V and propidium iodide (PI) were used to detect the apoptotic rate of H9c2 cells. At the end of our experiment, H9c2 cells (310 g) were centrifuged for 5 mins, followed by washing and resuspension in PBS twice. After a third centrifugation, H9c2 cells were gently resuspended in 500 μL of binding buffer solution. Annexin-V-FITC (5 μL) and PI (10 μL) were added to double-stain the H9c2 cells. After a treatment period of 15 mins in darkness, cellular apoptotic rate was measured by flow cytometry (BD Bioscience, Franklin Lakes, New Jersey, USA). The Annexin-V(+)/PI(-) cells (which showed up in the Q2-4 region) represented the early apoptotic cells. Annexin-V(+)/PI(+) cells (which showed up in the Q2-2 region) represented the late apoptotic cells. Each assay was performed six times. The mean result of the apoptotic rate was shown as a ratio of the number of apoptotic cells to the total number of cells.

**Measurement Of Intracellular Ca\(^{2+}\) Concentrations**

At the end of the experiment, the final solution was replaced with HBSS containing 4 μM Fluo-4AM. The H9c2 cells were then cultured at 37°C for 20 mins. Five times the volume of the HBSS containing 1% FBS was added, and the H9c2 cells were incubated for another 40 mins. The cells were then washed three times with HEPES buffer saline followed by resuspension in HEPES-buffered saline. The intracellular calcium level was analyzed with FACSCalibur flow cytometry at an excitation wavelength of 494 nm and emission wavelength of 510 nm. The fluorescence intensity of fluo-3 represented the concentration of intracellular Ca\(^{2+}\). Each assay was performed six times.

**Measurement Of Intracellular Reactive Oxygen Species (ROS)**

At the end of our experiment, the medium was pipetted out and replaced with 1 mL of 2',7'-dichlorofluorescin diacetate (DCFHDA) solution. The cells were incubated at 37°C for 20 mins in darkness and were then washed three times with PBS to sufficiently remove the intracellular DCFH-DA. The H9c2 cells were then resuspended in 500 μL of PBS and subjected to flow cytometry. The fluorescent intensity of dichlorofluorescin (DCF) was measured by flow cytometry at an excitation wavelength of 480 nm and an emission length of 530 nm. Each assay was performed six times. The mean result of intracellular ROS was expressed as the mean fluorescent intensity of DCF in channel FL-1.

**Measurement Of Intracellular MDA And SOD**

MDA, one of the reactive products oxygen radical and unsaturated fatty acid, can be used as a maker to estimate lipid oxidative stress. When reacted with thiobarbituric acid, a chromogen which absorbs at 532 nm is formed. Superoxide can react with this chromogen to form a soluble formazan. SOD, one of the anti-oxidative enzyme, can inhibit the production of this water-soluble formazan. The intracellular ROS and MDA levels were quantified using BCA standard curve method. MDA was measured at 532 nm, and SOD was measured at 450 nm per protocol. Each measurement was performed six times.

**Measurement Of Mitochondrial Membrane Potential (Δψm)**

JC-1 is an ideal fluorescent probe which can measure mitochondrial membrane potential. When mitochondrial membrane potential is high, JC-1 forms J-aggregates in the mitochondria and emits red fluorescence. When the mitochondrial membrane potential is low, Jc-1 remains a monomer and emits green fluorescence. The drop of Δψm is correlated with the reduction in the red/green fluorescence ratio. The H9c2 cells were then collected, washed and incubated in JC-1 solution at 37° in darkness for 20 mins. Next, the H9c2 cells in the incubation buffer were measured with flow cytometry at 488 nm excitation. The result was shown as the red/green fluorescence ratio. Each assay was performed six times.

**MPTP Activity Assay**

MPTP (mitochondrial permeability transition pore) is a nonspecific channel located between the mitochondria inner and outer membrane. It plays an important role during cell apoptosis and necrosis. BBcellprobe TM CA1 can easily penetrate the cell membrane into the cytoplasm and mitochondria and emits a green fluorescence. Upon adding a quenching agent to the fluorescence probe, the cytoplasmic green fluorescence would vanish, leaving only mitochondrial fluorescence intact. When MPTP opens, the green fluorescence would vanish, leaving only mitochondrial fluorescence intact.
would leak out of the mitochondria into the cytoplasm. Therefore, MPTP activity is indicated by mitochondrial green fluorescence changes. The lesser the amount of mitochondrial green fluorescence, the more active MPTP is. Each measurement was performed six times.

**Western Blot Analysis**

40 μg protein samples mixed with PBS and 5× loading buffer were dissolved in a SDS-polyacrylamide gel. The gel-electrophoresed protein bands were then transferred onto PVDF (polyvinylidene fluoride) membranes. These membranes were incubated with specific primary antibodies (anti-VDAC-1, anti-Bcl-2, anti-Bax, anti-caspase-3, anti-caspase-9). Anti-rabbit IgG-HRP was then added to the membrane. The blot was then visualized with ECL-Plus reagent. For protein loading controls, their membranes were stripped and re-probed with anti-β-actin and anti-COX IV antibodies. Densitometric analysis was used to quantify the protein bands, and the relative ratio of the optical density between study and control groups was calculated.

**Statistical Analysis**

Data were presented as mean ± standard deviation (S.D.). Statistical Package for the Social Sciences Software (SPSS.23.0 for windows) was used for statistical analysis. Tukey’s test with one-way ANOVA was used for multiple comparisons. A P value of <0.05 was considered significant.

**Results**

**PHC Reduced LDH And CK After A/R Injury**

LDH and CK indicate H9c2 cells' necrosis, as depicted in Figure 2A and B. When compared with the control group, A/R significantly increased LDH and CK levels (P<0.01, respectively). PHC reduced LDH and CK levels in all post-conditioning groups compared to A/R group (P<0.01). Even though there were no significant differences in CK levels amongst the PHC treated groups, the PHC postconditioning groups revealed significant differences in LDH levels (0 min vs 20 mins, P<0.01; 0 min vs 30 mins, P<0.05 and 5 mins vs 20 mins, P<0.05). The results may indicate that cell necrosis had happened not only during the early period of the reoxygenation (within 30 mins), but during the late (over 30 mins) period of the reoxygenation stage as well.

**PHC Protected H9c2 Against Cell Viability And Apoptosis**

After A/R injury, H9c2 cell viability changed significantly compared with the control groups (P<0.01). Postconditioning
administration of PHC at 0 min, 5 mins, 10 mins, 20 mins after the reoxygenation procedure all attenuated cell viability changes significantly compared to A/R group (P<0.01). However, postconditioning at 30 mins failed to demonstrate significant protective effect compared to A/R group (P = 0.80). There were no significant differences among 0 min, 10 mins and 20 mins postcondition groups (P>0.05, respectively) (Figure 3A).

Cell apoptosis was detected by analyzing PHC protective ability. Annexin V-FITC and PI double staining were used to measure the percentage of the apoptotic H9c2 cells. Q2-4 and Q2-2 represented early and late apoptotic H9c2 cells, respectively. (Figure 4A) A/R injury significantly increased cellular apoptosis compared to control group (P<0.01). PHC treatment decreased the apoptosis rate (P<0.01), with PHC administration at 0 min showing the most protective effect compared to other groups (P<0.01). Administering PHC 30 mins post reoxygenation showed the least protective effect compared to the 0 min, 5 mins, 10 mins, 20 mins groups (P<0.01), indicating that cell apoptosis happened during the earlier stage of the reoxygenation procedure (Figure 4B).

Cell apoptosis and cell necrosis are the two main types of cell death. As the A/R procedure progressed, H9c2 cell death increased as indicated by the changes in cell viability. Figure 2A, B and 4B indicated that cell apoptosis was affected more by timing effect of PHC postconditioning in the early stage of reoxygenation in comparison to cell necrosis (which indicated by CK and LDH level). Therefore, we believe H9c2 cell apoptosis occurred primarily in the very early period of postconditioning stage (especially within 30 mins), while H9c2 cells necrosis occurred not only during the early period of the postconditioning stage but throughout the reoxygenation phase as well. Hence, early PHC administration would inhibit both cell apoptosis and necrosis. Even though administration of PHC at 30 mins after reoxygenation did not significantly affect cell apoptosis rate, PHC may still offer some protective effect toward cell injury and cell necrosis. Cell apoptosis occupied a large percentage of cell death procedure in the early A/R period (as is shown in Figure 4C) and cell necrosis and other types of cell death also play an important part in A/R procedure. Therefore, PHC postconditioning therapy attenuated cell death not only through cell apoptosis pathway but via cell necrosis pathway as well. (Figure 4C was drawn by using the mean level of each group in Figures 3A and 4B. Cell variability changes indicated cell death. The result was shown as difference value in A/R group and each postconditioning group compared with the control group.)

Cell and H9c2 Cellular Oxidative Protection

Oxidation plays a significant role in ischemia-reperfusion injury. ROS, SOD and MDA were measured to analyze the protective effect of PHC. MDA, the end product of lipid oxidation, increased significantly in the A/R group compared to the control group (P<0.01). PHC decreased MDA levels in all the groups (P<0.01). But 0 min, 5 mins and 10 mins groups exhibited better protection compared to 20 mins and 30 mins groups (P<0.05).

SOD, a dismutase which can eliminate oxygen free radical in vivo, has protective effect against ischemia-reperfusion injury. MDA can be used as a maker to estimate lipid oxidative stress. Our results showed that A/R decreased SOD and increased MDA levels compared to the control group (P<0.01). PHC increased SOD and decreased MDA levels (P<0.01) versus the A/R group. There were significant differences in SOD and MDA levels among 0 min vs 20 mins and 0 min vs 30 mins in PHC postconditioning groups (Figure 5A and B).
ROS, analyzed with flow cytometry, was found to be elevated in A/R group compared to the control group. PHC was able to decrease ROS levels, with administration within 20 mins having a more significant effect compared to 30 mins group (P<0.01). However, there were no significant differences among the other PHC postconditioning groups (0 min, 5 mins, 10 mins, 20 mins, P>0.05, respectively) (Figure 6A and B).

**PHC Protects H9c2 Cells From Ca^{2+} Overload**

Ca^{2+} overload plays an important part in A/R injury. A/R significantly increased Ca^{2+} overload (P<0.01). PHC postconditioning attenuated Ca^{2+} overload in all of the PHC postconditioning groups. However, administering PHC at 30 mins after reoxygenation had the least effect compared to other groups (P<0.01). There were no significant differences among the other PHC postconditioning groups (0 min, 5 mins, 10 mins, 20 mins) (P>0.05, respectively) (Figure 7A and B).

**PHC Restored A/R-Induced Mitochondrial Membrane Potential**

Mitochondrial membrane (potential) drops in the apoptotic pathway can further accelerate cell death and cell apoptosis. JC-1 assay was used to detect ΔΨm (Figure 8A). Whereas A/R decreased mitochondrial membrane potential, PHC restored cell membrane potential (P<0.01). However, administering PHC within 10 mins had more significant effect versus the 20 mins and 30 mins groups (P<0.01). Administering PHC at 30 mins after the reoxygenation exhibited the least effect compared to other postcondition groups (P<0.01). The result was shown in a red/green fluorescence ratio (Figure 8B).
PHC Affect Mitochondrial Pathway During Cell Apoptosis In A/R Injury

Mitochondria have an important part in apoptosis. Death signals were integrated through Bcl-2 family and MPTP opening and the caspase family Cyt-C was released, MPTP activity was enhanced, and caspase family was activated in ischemia-reperfusion injury. Therefore, we measured Bcl-2, Bas, caspase-3 and caspase-9 level.

Figure 5 Effect of PHC postconditioning on SOD and MDA level. A/R decreased SOD and increased MDA level compared with control group ($P<0.01$), and PHC increased SOD and decreased MDA level ($P<0.01$) compared with A/R group (A and B). There is significant difference of SOD level among 0 min vs 20 mins and 0 min vs 30 mins in PHC postcondition groups ($P<0.05$, respectively) (A), and there is significant difference of MDA level among 10 mins vs 20 mins and 20 mins vs 30 mins in PHC postcondition groups (B). (n=6; ##$P<0.01$ versus control group; $$P<0.01$ versus A/R group. Data are shown as mean±S.D.).

Figure 6 Effect of PHC postconditioning on ROS level. ROS level was detected by flow cytometry with DCF staining technique (A). ROS was found increasing in A/R group compared with the control group, and PHC can drop ROS level compared with the A/R group, yet administering PHC within 20 mins has more significant effect compared with 30 mins group ($P<0.01$). However, there are no significant differences among the other PHC postconditioning groups (0 min, 5 mins, 10 mins, 20 mins, $P>0.05$, respectively) (B). (n=6; ##$P<0.01$ versus control group; $$P<0.01$ versus A/R group. Data are shown as mean±S.D.).
to determine whether or not PHC could regulate mitochondrial pathways, as depicted in Figure 9A. The results revealed a decrease in Bcl-2, and an increase in Bax, caspase-3 and caspase-9 in A/R group versus the control group (P<0.01). As depicted by Western blot, PHC significantly attenuated the apoptosis pathway, and administering PHC at 30 mins after the reoxygenation was shown to have the least effect compared to other postcondition groups (P<0.01, respectively), as shown in Figure 9B–F.

PHC Attenuates MPTP Activity And VDAC Level

Ischemia-reperfusion (injury) accelerates MPTP opening and plays an important part in accelerated cell apoptosis and cell death. We therefore aimed to find out if PHC would inhibit MPTP activity and protect H9c2 cell from A/R injury. Confocal microscopy and fluorescent staining technique were used to observe MPTP activity. When MPTP opened, the green fluorescence would leak out of
Figure 9 Western blot analysis of VDAC1, Bcl-2, Bax, cleaved caspase-3 and cleaved caspase-9 (A). As depicted in the above pictures, there is a decrease of Bcl-2, an increase of Bax, caspase-3 and caspase-9 in A/R group compared with the control group. As depicted by Western blot, PHC significantly attenuated the apoptosis pathway, and administering PHC at 30 mins after the reoxygenation showed the least effect compared with other postcondition groups (P<0.01, respectively). (B–F) (n=6; **P < 0.01 versus control group, ***P < 0.01 versus A/R group. Data are shown as mean±S.D.).
the mitochondria. Therefore, MPTP activity was indicated by mitochondrial green fluorescence change. There was inverse relationship between the amount of mitochondrial green fluorescence and MPTP activity, as shown in Figure 10A. VDAC, component of MPTP, also influenced cell death and survival.\textsuperscript{25,26} Western blot technique was used to access VDAC level (Figure 9A). As a result, A/R increased VDAC level, and promoted MPTP opening. PHC decreased VDAC level and attenuated MPTP activity. Earlier administration of PHC offered greater postconditioning protective effect (Figures 9B and 10A).

**Discussion**

Our experiment showed that PHC postconditioning had a positive effect in protecting H9c2 cells from A/R injury. Earlier PHC administration resulted in better protection. Our previous investigation showed that Penehyclidine Hydrochloride offered this protective effect via limiting calcium overload, decreasing formation of reactive oxygen species, restraining mitochondrial MPTP opening and reducing inflammatory response in vitro and in vivo.\textsuperscript{5,6} The most effective dose of Penehyclidine Hydrochloride on H9c2 cell against anoxia/reoxygenation injury was determined to be 0.1 μM/L in our research group.\textsuperscript{9}

In 2012, Barrere-Lemaire et al found out the effective time window for delayed postconditioning treatment in rats model was 30 mins,\textsuperscript{27} and our previous animal study determined the protective time window of PHC postcondition was within 10 mins after the reperfusion stage began.\textsuperscript{6} We hypothesized that IRI could be attenuated in the early period of the reperfusion stage. PHC was administered directly in the cell culture, which effectuated H9c2 cells directly, would offer more direct protective effect compared with those in vivo studies. In this cell culture experiment, we focused on the variation tendency of each mechanism in the very early reperfusion stage (from 0 min to 30 mins) to better understand the time effect of each mechanism in A/R procedure in the treatment of PHC. There were not many differences in CK and LDH level from 0 min to 30 mins PHC postconditioning treatment, while there were huge differences in cell apoptosis, mitochondrial pathway and Ca\textsuperscript{2+} concentration. Regardless of when PHC was administered within the first 30 mins after reoxygenation began, H9c2 cells were protected via almost all of the mechanisms that contributed to A/R injury. Our results indicate that postcondition treatment should be applied as early as possible, preferable within 20 mins after reoxygenation began. Oxidative stress, calcium overload, MPTP opening and cell apoptosis all started in the early reoxygenation stage and persisted, resulting in cell injury and necrosis. Administering PHC during any period in the reoxygenation stage could attenuate oxidative stress, calcium overload, mitochondrial pathway and cell death, which would offer some degree of cellular

![Figure 10](image-url) MPTP activity observed by confocal microscopy and with fluorescent staining technique. A/R promotes MPTP opening, and PHC attenuated MPTP activity.
protection. Earlier treatment of PHC postconditioning offer greater protective effect.

During cardiomyocyte anoxic period, oxidative phosphorylation halts as a result of anoxia in the mitochondrial respiratory chain. Under anaerobic conditions, the glycolysis becomes the dominant pathway of ATP generation. However, glycolysis would lead to lactic acid accumulation. As blood flow stops and CO2 elimination fails, continued cellular acidosis would lead to a drop in cellular pH. The increasing \( \text{H}^+ \) in the cell will open the \( \text{Na}^+/\text{H}^+ \) channels. The increase in cytoplasmic \( \text{Na}^+ \) will stimulate \( \text{Na}^+/	ext{Ca}^{2+} \) exchange, resulting in calcium overload.\(^{28}\) Even with restoration of adequate oxygen supply during reoxygenation, mitochondrial respiratory chain complexes still could not play its role. As a result, a large amount of reactive oxygen species (ROS) was generated, inflammatory response was activated, calcium overload ensued and mitochondrial permeability transition pore (MPTP) was activated.\(^{29}\) Mitochondria played the central role of cell death and apoptosis. With MPTP opening, mitochondria starts to swell and inner membrane begins to depolarize. This leads to the release of cytochrome C and activation of caspase. Ultimately, cell death and apoptosis occur.\(^{30}\)

Acute myocardial ischemia is a common occurrence all around the world, and it is difficult to predict when it will happen. This in vitro experiment may offer some insight into treating ischemia-reperfusion injury with postconditioning method. Our result showed that PHC postconditioning in 30 mins still offered its protective effect, which may provide improved prognosis for patients with myocardial infarction or with other organ shock. Further in vitro and clinical research utilizing PHC are needed. We hope this current in vitro experiment would increase confidence for researchers, and physicians in future exploration of postconditioning ischemia treatment.

**Conclusion**

PHC postconditioning protected H9c2 cells by attenuating superoxide production, decreasing \( \text{Ca}^{2+} \) overload, restraining MPTP activities, restoring mitochondrial membrane potential, regulating cell apoptosis proteins and modulating the mitochondrial pathway. The most effective dose of PHC on H9c2 cell against IRI was determined to be 0.1 \( \mu \)g/L. H9c2 cells were protected by PHC from A/R injury regardless of when PHC was administered (0 min, 5 mins, 10 mins, 20 mins, 30 mins). However, earlier administration of PHC resulted in better PHC postconditioning protective effect.

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**Disclosure**

The authors report no conflicts of interest in this work.

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