Propofol Post-conditioning Protects against COS-7 Cells in Hypoxia/reoxygenation Injury by Induction of Intracellular Autophagy

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Background: Propofol (2,6-diisopropylphenol) is a widely used intravenous anesthetic agent for the induction and maintenance of anesthesia during surgeries and sedation for ICU patients. Propofol has a structural similarity to the endogenous antioxidant vitamin E and exhibits antioxidant activities. However, the mechanism of propofol on hypoxia/reoxygenation (H/R) injury has yet to be fully elucidated. We investigated how P-PostC influences the autophagy and cell death, a cellular damage occurring during the H/R injury.

Methods: The groups were randomly divided into the following groups: Control cells were incubated in normoxia (5% CO2, 21% O2, and 74% N2) without propofol treatment. H/R cells were exposed to 24 h of hypoxia (5% CO2, 1% O2, and 94% N2) followed by 12 h of reoxygenation (5% CO2, 21% O2, and 74% N2). H/R + P-PostC cells post-treated with propofol were exposed to 24 h of hypoxia followed by 12 h of reoxygenation. 3-MA + P-PostC cells pretreated with 3-MA and post-treated propofol were exposed to 24 h of hypoxia followed by 12 h of reoxygenation. The results of our present study provide a new direction of research on mechanisms of propofol-mediated cytoprotection. There are three principal findings of these studies. First, the application of P-PostC at the onset of reoxygenation after hypoxia significantly increased COS-7 cell viability. Second, the cellular protective effect of P-PostC in H/R induced COS-7 cells was probably related to activation of intracellular autophagy. And third, the autophagy pathway inhibitor 3-MA blocked the protective effect of P-PostC on cell viability, suggesting a key role of autophagy in cellular protective effect of P-PostC.

Conclusions: These data provided evidence that P-PostC reduced cell death in H/R model of COS-7 cells, which was in agreement with the protection by P-PostC demonstrated in isolated COS-7 cells exposed to H/R injury. Although the this study could not represent the protection by P-PostC in vivo, the data demonstrate another model in which endogenous mechanisms evoked by P-PostC protected the COS-7 cells exposed to H/R injury from cell death.

Key Words: Cos-7 cells; Hypoxia reoxygenation; Post-Treated; Propofol

INTRODUCTION

Tissue Autophagy is an evolutionary conserved process involved in degradation of long-lived or damaged proteins and organelles [1]. Autophagy has been implicated in tissue ischemia/reperfusion (I/R). Nevertheless, the exact functional role of autophagy in the cell survival and death pathways associated with tissue damage is still not clear. Some observations suggest that stimulation of autophagy can promote cell survival in response to stress conditions in cardiomyocytes [2,3]. On the other hand, the excessive and long-term upregulation of autophagy under certain conditions inhibit cell survival, rather greatly accelerating the cell death in the heart [4]. The term “ischemic pre-conditioning” (IFC) was first coined...
by Murry [5] to describe a phenomenon whereby brief episodes of I/R could attenuate tissue injury, but its clinical application has been disappointing due to the unpredictability of ischemia. Ischemic post-conditioning (I–PostC) [6] performed after ischemia has solved the problem of unpredictability. Furthermore, pharmacological post-conditioning, which only requires a drug given as an adjunctive intervention during early reperfusion, can also achieve tissue protection and is easier to implement [7–9]. The mechanisms responsible for post-conditioning are largely unknown, but some previous studies have suggested that signal transduction pathways for survival such as activation of the extracellular signal-regulated kinase (ERK) [10] and of the phosphoinositide–3–kinase/Akt (PI3K/Akt) pathways [11] may play an important role in the protective effects of post-conditioning, by stimulating the endogenous anti-injury capacity of cells to protect themselves [12].

Propofol (2,6-diisopropylphenol) is a widely used intravenous anesthetic agent for the induction and maintenance of anesthesia during surgeries and sedation for ICU patients. Propofol has a structural similarity to the endogenous antioxidant vitamin E and exhibits antioxidant activities [13]. Wang et al. [14] found that infusing propofol at the onset of reperfusion for 30 min could provide neuroprotection to transient middle cerebral artery occlusion (MCAO) in rats partly by activation of the PI3K/Akt pathway, and He reported that post-conditioning with propofol (P–PostC) has cardioprotective effects against I/R injury of the heart, which is associated with inhibition of mitochondrial permeability transition pore (MPTP) opening [15]. However, the mechanism of propofol on hypoxia/reoxygenation (H/R) injury has yet to be fully elucidated. COS–7 cell is the African green monkey kidney fibroblast–like cell line has been established from CV–1 cells which have been transformed by an origin–defective mutant of SV40 coding for wild–type T antigen. This line contains T antigen, retains complete permissiveness for lytic growth of SV40. Here we investigated how P–PostC influences the autophagy and cell death, a cellular damage occurring during the H/R injury.

MATERIALS AND METHODS

1. Reagents

Propofol (2, 6-diisopropylphenol) diluted with dimethyl sulfoxide (DMSO). The following reagents were obtained commercially: 3-[4,5-dimethylthiazol-2-yl]2,5-diphenyl tetrazolium bromide (MTT), acridine orange, monodansylcadaverine (MDC), 3-methyladenine (3-MA, class III PI3K inhibitor) was obtained from Calbiochem (La Jolla, CA, USA). Antibodies used in the study were as follows: LC3–II (1:3,000), Beclin–1 (1:1,000) from Abcam, and p62 (1:1,000), Atg5 (1:500) from Santa Cruz. Secondary antibodies against rabbit (1:3,000) and mouse (1:3,000) immunoglobulins were purchased from Bio–Rad.

2. Cell culture

COS–7 cells (African green monkey kidney fibroblast–like cell line) were obtained from the American Type Culture Collection (ATCC, Manassas, USA). Dulbecco’s modified Eagle’s medium (DMEM, GIBCO) supplemented with 10% inactivated fetal bovine serum (FBS, GIBCO) containing 500 μg/ml penicillin and 500 μg/ml streptomycin (GIBCO), and cells were incubated at 37°C in a humidified atmosphere with 5% CO2. Media were changed every 3 days.

3. Treatment of propofol postconditioning

Propofol which were made by dissolving them in DMSO were kept frozen at −4°C until use. The stock was diluted to their concentration with DMEM when needed.
Prior to propofol treatment, cells were grown to about 80% confluence and then exposed to propofol at different concentrations (0, 25, 50, 100 μM) for 2 h. Cells grown in medium containing an equivalent amount of DMSO without propofol served as control (Fig. 1).

4. MTT assay

Cell viability was measured using a quantitative colorimetric assay with thiazolyl blue tetrazolium bromide (MTT, AMResco), showing the mitochondrial activity of living cells. COS-7 cells (3 x 10^4) were seeded in 96-well plates. After drug treatment as indicated, cells were incubated with 300 μl MTT (final concentration 0.5 mg/mL) for 1.5 h at 37°C. The reaction was terminated by addition of 200 μl DMSO. Cell viability was measured by an ELISA reader (Tecan, Männedorf, Switzerland) at 570 nm excitatory emission wavelength.

5. Fluorescence microscopy

Cells were grown on coverslips and treated with COS-7 cells. After 24 h, cells were stained with 0.05 mM MDC, a selective fluorescent marker for autophagic vacuoles, at 37°C for 1 h. The cellular fluorescence changes were observed using a fluorescence microscope (Axioskop, Carl Zeiss, Germany). For further detection of the acidic cellular compartment, we used acridine orange, which emits bright red fluorescence in acidic vesicles but fluoresces green in the cytoplasm and nucleus. Cells were stained with 1 μg/mL acridine orange for 15 min and washed with PBS. AVOs (autophagic vesicles organelles) formation was obtained under a confocal microscope LSM 700 (Carl Zeiss, Germany).

6. Western blot analysis

Cells (2 x 10^6) were washed twice in ice-cold PBS, resuspended in 200 μl ice-cold solubilizing buffer [300 mM NaCl, 50 mM Tris–Cl (pH 7.6), 0.5% Triton X–100, 2 mM PMSF, 2 μl/ml aprotinin and 2 μl/ml leupeptin] and incubated at 4°C for 30 min. The lysates were centrifuged at 14,000 revolutions per min for 15 min at 4°C. Protein concentrations of cell lysates were determined with Bradford protein assay (Bio–Rad, Richmond, CA, USA) and 20 μg of proteins were resolved by 10% SDS/PAGE. The gels were transferred to polyvinylidene fluoride (PVDF) membranes (Millipore, Billerica, MA, USA) and reacted with appropriate primary antibodies. Immunostaining with secondary antibodies was detected using SuperSignal West Femto (Pierce, Rockford, IL, USA) enhanced chemiluminescence substrate and detected with Alpha Imager HP (Alpha Innotech, Santa Clara, USA).

7. Statistical analysis

Data were expressed as means ± SEM. The statistical significance was assessed by one–way ANOVA followed by Tukey’s post hoc t–test. (P < 0.05 was considered statistically significant)

RESULT

1. P–PostC improved the cell viability of H/R–induced COS–7 cells

The effect of P–PostC and H/R injury on COS–7 cells was investigated. Propofol did not show any significant
toxic effect on the COS-7 cells (Fig. 2). After COS-7 cells were exposed to hypoxia and post-treatment with various doses of propofol and exposed to reoxygenation, we measured cell viability by MTT assay again to investigate the effect of various concentrations of P-PostC (0, 25, 50, 100 and 200 μM) on the H/R-induced COS-7 cells and discovered that P-PostC significantly protected the COS-7 cells from H/R-induced cytotoxicity and our results showed that the viability of P-PostC COS-7 cells was increased in a dose-dependent manner (Fig. 3A). As shown in Fig. 3A, compared with control, the cell proliferation was markedly inhibited after H/R treatment (P < 0.05), which was significantly improved by P-PostC. Among all of the concentrations, 100 μM Propofol represented the optimal effect, improving cell viability by approximately 20% (P < 0.05). Based on this result, all subsequent experiments were performed with 100 μM propofol. The role of autophagy in the H/R damage of COS-7 cells was further confirmed by the autophagy inhibitor 3-MA, an inhibitor of class III phosphoinositide 3-kinase (PI3K). As shown in Fig. 3B, 3-MA (5 mM) 1 h before H/R inhibited protective effect of P-PostC and significantly increased H/R induced COS-7 cell death. This decrease in the cell viability was accompanied by the autophagy inhibitor 3-MA, indicating that the inhibition of autophagy by 3-MA sensitized COS-7 cells to H/R injury.

2. P-PostC leads to induction of autophagy in COS-7 cells

Prominent accumulation of autophagic specific staining MDC was observed around the nuclei in P-PostC group COS-7 cells (Fig. 4). Similarly, AO staining, red
Fig. 4. Fluorescence microscopic (× 400) analysis of autophagosome in the H/R injured COS-7 cells. H/R caused accumulation of autophagosomes containing partially digested cytoplasmic contents compared to the control group. The P–PostC during I/R dramatically increased formation of autophagosomes and the autophagy pathway inhibitor 3–MA blocked formation of autophagosomes by P–PostC. H/R, Hypoxia/Reoxygenation; P–PostC, post-conditioning with propofol; 3–MA, 3–Methyladenine.

Fig. 5. Fluorescence microscopic (× 400) analysis of autophagosome in the H/R injured COS-7 cells. Stained with acridine orange the green shows where the dye has stained the nucleus and the red is where the cell is starting to ‘digest’ parts of itself in small capsules called autophagosomes. H/R caused accumulation of autophagosomes containing partially digested cytoplasmic contents compared to the control group. The P–PostC during I/R dramatically increased formation of autophagosomes and the autophagy pathway inhibitor 3–MA blocked formation of autophagosomes by P–PostC. H/R, Hypoxia/Reoxygenation; P–PostC, post-conditioning with propofol; 3–MA, 3–Methyladenine.

Fig. 6. (A) Western blot analysis. Expression of Atg5, LC3–II, Beclin–1 and p62 in H/R–induced COS–7 cells preconditioned with propofol and 3–MA. (B) Quantification of Atg5, LC3–II, Beclin–1 and p62. Each band shown above was quantified by the densitometric scan. H/R, Hypoxia/Reoxygenation; PPC, propofol postconditioning; 3–MA, 3–Methyladenine.
fluorescent spots appeared on P–PostC group COS–7 cells, while the normoxia control group, and the cells co–treated with 5 mM 3–MA and H/R group showed mainly green cytoplasmic fluorescence (Fig. 5). We here examined activation of autophagy related protein under H/R–induced cells by western blotting analysis. The recruitment of LC3 to the membrane occurs via an Atg5–dependent mechanism, and thus Atg5 is essential for autophagosome formation in vivo.3) Atg5 and Beclin–1, LC3–II (microtubule–associated protein 1 light chain 3 form II), p62 was significantly reduced in H/R–induced group cells but elevated in P–PostC group cells (Fig. 6). As showed in Fig. 6, Atg5, Beclin–1, LC3–II (microtubule–associated protein 1 light chain 3 form II) and p62 was increased when autophagy was induced by P–PostC, and they were decreased when autophagy was suppressed by 3–MA.

**DISCUSSION**

The previous reports have shown that the cellular protective mechanism of propofol may partly result from its ability to act as a free radical scavenger [16–18]. The results of our present study provides a new direction of research on mechanisms of propofol–mediated cytoprotection. There are three principal findings of these studies. First, the application of P–PostC at the onset of reoxygenation after hypoxia significantly increased COS–7 cell viability. Second, the cellular protective effect of P–PostC in H/R induced COS–7 cells was probably related to activation of intra–cellular autophagy. And third, the autophagy pathway inhibitor 3–MA blocked the protective effect of P–PostC on cell viability, suggesting a key role of autophagy in cellular protective effect of P–PostC. Together, these findings indicate that P–PostC stimulated COS–7 cells endogenous cellular protective effect against H/R injury through pro–survival autophagy signal pathways. Numerous studies have documented that oxidative stress–mediated cellular changes are frequently induced during H/R [19–21]. Excessive intracellular ROS play crucial roles in the induction of cell apoptosis and death. The postconditioning treatment at the onset of reoxygenation reduces H/R induced injury in cardiomyocytes and is potentially mediated by attenuation of ROS generation [22]. Similar to the effects of ischemia post–conditioning, our study shows that P–PostC induced intracellular autophagy and increased COS–7 cell viability. According to recent works, moreover, it has been suggested that autophagy plays two distinct roles during ischemia and reperfusion. In the ischemic phase, autophagy can be protective via AMPK activation and sequentially inhibition of mTOR signaling, but reperfusion after ischemia stimulates autophagic cell death through the different pathway. In this study P–PostC induced intra–cellular autophagy and decreased H/R induced COS–7 cell death.

Autophagic vacuoles including autophagosomes and autophagolysosomes are generally formed in cells undergoing the autophagic process. Therefore, observation of autophagosomes inside cells can be used as an indicator to analyze the induction of autophagy. To investigate the role of autophagy in the H/R induced COS–7 cells and the effect of P–PostC in the occurrence of autophagy, we directly observed formation of autophagosomes in COS–7 cells exposed to H/R using a fluorescence microscope. H/R caused accumulation of autophagosomes containing partially digested cytoplasmic contents in COS–7 cells compared to the control group. And P–PostC during H/R dramatically increased formation of cytosolic vacuoles in COS–7 cells and decreased H/R induced cell death, indicating that autophagy is associated with H/R–induced cell death, and P–PostC protects cell death through the induction intra–cellular autophagy. To further confirm whether propofol can modulate autophagic cell death in
HR-injured COS-7 cells, we examined the level of LC3-II expression. The production of LC3-II has been suggested as another marker for autophagy induction. LC3-II formation was increased in the COS-7 cells exposed to HR injury compared to the control group. Furthermore, in the case of P-PostC during HR, LC3-II production was significantly increased compared to the HR group. The expression of p62, an autophagy-associated protein was alleviated in the HR COS-7 cells, but it was increased by P-PostP to the HR injured cells. This data indicates again that the HR injury increased cell death and P-PostC induces autophagy leading to cell survival. In conclusion, these data provided evidence that P-PostC induced intracellular autophagy and reduced cell death in HR model of COS-7 cells, which was in agreement with the protection by P-PostC demonstrated in isolated COS-7 cells exposed to HR injury through the induction of cellular protective autophagy. Although the this study could not represent the protection by P-PostC in vivo, the data demonstrate another model in which endogenous mechanisms evoked by P-PostC protected the COS-7 cells exposed to HR injury from cell death.

REFERENCE


