SHORT COMMUNICATION

Glycogen synthase kinase-3β opens mitochondrial permeability transition pore through mitochondrial hexokinase II dissociation

Takamitsu Tanaka1 · Masao Saotome1 · Hideki Katoh1 · Terumori Satoh1 · Prottoy Hasan1 · Hayato Ohtani1 · Hiroshi Satoh1 · Hideharu Hayashi1 · Yuichiro Maekawa1

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Abstract
Accumulating evidence has revealed pivotal roles of glycogen synthase kinase-3β (GSK3β) inactivation on cardiac protection. Because the precise mechanisms of cardiac protection against ischemia/reperfusion (I/R) injury by GSK3β-inactivation remain elusive, we investigated the relationship between GSK3β-mediated mitochondrial hexokinase II (mitoHK-II; a downstream target of GSK3β) dissociation and mitochondrial permeability transition pore (mPTP) opening. In Langendorff-perfused hearts, GSK3β inactivation by SB216763 improved the left ventricular-developed pressure and retained mitoHK-II binding after I/R. In permeabilized myocytes, GSK3β depolarized mitochondrial membrane potential with accelerated mitochondrial calcein release (suggesting GSK3β-mediated mPTP opening) and decreased mitoHK-II bindings. GSK3β-mediated mPTP opening depended on mitoHK-II binding, i.e., it was accelerated by dissociation of mitoHK-II (dicyclohexylcarbodiimide) and attenuated by enhancement of mitoHK-II binding (dextran). However, inactivation of mitoHK-II by glucose-depletion or glucose-6-phosphate inhibited the GSK3β-mediated mPTP opening. We conclude that GSK3β-mediated mPTP opening may be involved in I/R injury and regulated by mitoHK-II binding and activity.

Keywords Glycogen synthase kinase-3β · Mitochondrial permeability transition pore · Mitochondrial hexokinase II · Ischemia–reperfusion

Introduction
Mitochondria play pivotal roles not only manipulating cellular function through ATP production and Ca2+ regulation [1] but also determining myocardial cell fate during ischemia/reperfusion (I/R) [2]. Accumulating evidence indicates that the inhibition of mitochondrial permeability transition pores (mPTP) contributes to the core mechanism of cardiac protection including pro-survival PI3 K-Akt kinase cascades, which are activated by preconditioning to initiate myocardial protection [2, 3].

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* Masao Saotome
msaotome@hama-med.ac.jp

1 Internal Medicine III, Hamamatsu University School of Medicine, 1-20-1 Handayama, Higashi-ku, Hamamatsu 431-3192, Japan

Glycogen synthase kinase 3β (GSK3β), which is inactivated by its phosphorylation, is a constitutive multifunctional serine/threonine kinase [4, 5] and manipulates cellular glycogen synthase activity and multiple protective pro-survival signaling pathways [4]. Recent investigations suggested that the mitochondria-bound hexokinase II (mitoHK-II) is a downstream target of GSK3β [6, 7]. MitoHK-II is reported to promote neuronal survival in human neuron-like cells [8], and dissociation of HK-II from mitochondria increased the chemotherapy-induced lethal cell damage in HeLa cells [6]. In addition, ischemic preconditioning is associated with reduced cytosolic HK-II activity during ischemia and biphasic induction of mitoHK-II activity before and after ischemia [9].

Despite intensive efforts, the precise mechanisms of myocardial survival against I/R by GSK3β inactivation have not been fully elucidated. Thus, in this study we aimed to investigate (1) the effects of GSK3β inactivation on cardiac function during and after I/R, (2) the relationship between GSK3β-mediated mPTP opening and mitoHK-II binding,
and (3) the impact of HK-II inactivation by manipulating glucose metabolism on GSK3β-mediated mPTP opening.

Methods

This investigation conforms to the Guide for the Care and Use of Laboratory Animals published by the US National Institutes of Health (NIH Publication, revised 2011) and the Hamamatsu University School of Medicine Animal Care and Use Committee. Heart isolation and obtaining hemodynamic measurements in Langendorff-perfused hearts was described previously [10]. Isolated hearts were subjected to 35 min of global ischemia followed by 40 min of reperfusion with and without SB216763 (a GSK3β inhibitor [3 μmol/l]) treatment. In SB216763-treated hearts, SB216763 was pretreated for 25 min before ischemia, and then hearts were subjected with I/R without SB216763. To monitor LV pressure, a water-filled latex balloon connected to a pressure transducer and polygraph (Nihon Kohden Co., Japan), was inserted into the LV from the left atrium, and the hearts were electrically paced at 5 Hz.

Myocytes were isolated from male Sprague–Dawley rats and sarcolemmal membrane was permeabilized as described previously [10, 11]. After sarcolemmal membrane permeabilization, the concentration of free calcium in the solution was changed to 177 nmol/l. Western-blot analysis in mitochondrial fractions was performed as previously described [10]. Permeabilized myocytes were treated with drugs according to the protocol, and then homogenized using a ProteoExtract® Cytosol/Mitochondria Fractionation Kit (Merck Bioscience, Bad Soden, Germany). Densitometric analysis was performed using the Molecular Imager ChemiDoc™ system (Bio-Rad Laboratories, Hercules, CA, USA). Fluorescence measurements were performed with a laser scanning confocal microscope (LSM5 PASCAL, Carl Zeiss AG, Oberkochen, Germany) coupled to an inverted microscope with a 63 × water-immersion objective lens. Mitochondrial membrane potential (ΔΨm) was measured with tetramethylrhodamine ethyl ester (TMRE 10 nmol/l), and opening of mPTP was measured with calcein-AM (1 μmol/l), as described previously [11].

A recombinant active form of GSK3β was purchased from Millipore (Billerica, MA, USA). Cyclosporin A (CsA), protein kinase A catalytic subunit (PKAcat), dextran, and alamethicin were purchased from Sigma–Aldrich (St. Louis, MO, USA). SB216763 was purchased from Tocris Bioscience (Ellisville, MO, USA). Dicyclohexylcarbodiimide (DCCD) was purchased from Wako Chemicals (Richmond, VA, USA). Fluorescent dyes were purchased from ThermoFisher Scientific (Waltham, MA, USA).

Data are presented as mean ± SEM, and the number of cells or experiments is shown as n. Statistical analyses were performed using one-way ANOVA followed by Bonferroni’s test or by Kruskal–Wallis’s test, and two-way ANOVA followed by Bonferroni’s test, according to the study protocol. P < 0.05 was accepted as statistically significant.

Results

We first investigated the effects of GSK3β inactivation on I/R injury using SB216763. As shown in Fig. 1a, b, recovery of hemodynamics after reperfusion was significantly improved in SB216763-treated hearts (LVDP at 100 min 85.0 ± 4.6% of I/R + SB, P < 0.05 vs. 63.1 ± 3.0% of I/R). As a downstream target of GSK3β, expression level of mitochondrialhexokinase II (mitoHK-II) was assessed in SB216763-treated hearts. MitoHK-II was significantly reduced after I/R, and SB216763 prevented the reduction of mitoHK-II by I/R (Fig. 1c).

The causal relationship between GSK3β and mitoHK-II binding was further investigated using recombinant (active form) GSK3β and permeabilized myocytes. GSK3β significantly reduced the expression level of mitoHK-II similar to DCCD (Fig. 1d an agent to dissociate HK-II from mitochondria). In addition, GSK3β exhibited the dose-dependent mitoHK-II reduction (Supplemental fig. 1) and SB216763 inhibited the GSK3β-mediated mitoHK-II reduction (Fig. 1e). We next examined the effects of GSK3β (active form) on mPTP opening in permeabilized myocytes. As shown in Fig. 2a, the entrapped mitochondrial calcein release was accelerated by GSK3β (10 mmol/l calcein intensity at 35 min 79.2 ± 1.3% of the baseline, P < 0.05 vs. 94.2 ± 1.4% of CTL) in a SB216763-sensitive manner (3 μmol/l 92.1 ± 1.2% vs. 0.01 vs. GSK). In Fig. 2b, GSK3β significantly reduced TMRE intensity (at 25 min 78.3 ± 2.4% of the baseline, P < 0.05 vs. 100 ± 1.4% of CTL, suggesting depolarized ΔΨm), and SB216763 attenuated the GSK3β-mediated TMRE reduction (89.3 ± 4.3%, P < 0.01 vs. GSK, suggesting protection of ΔΨm). In addition, when cells were pretreated with cyclosporine A (CsA, an inhibitor of mPTP 0.1 mmol/l), the GSK3β-mediated mitochondrial collapse release (Fig. 2c 90.4 ± 1.8% of the baseline, P < 0.01 vs. GSK) and TMRE reduction (Fig. 2d 89.7 ± 3.6% of the baseline, P < 0.01 vs. GSK) were significantly inhibited. Because protein kinase A is known to inactivate GSK3β, the effects of protein kinase A catalytic subunit (PKAcat) on GSK3β-mediated mPTP opening were evaluated. In our previous report, PKAcat alone (< 15 U/ml) did not alter mPTP opening [11], here we treated permeabilized myocytes with 10 U/ml of PKAcat prior to GSK3β. As shown in Fig. 2c, d, PKAcat inhibited the GSK3β-mediated mitochondrial collapse release (94.5 ± 1.5%, P < 0.01 vs. GSK) and TMRE reduction (91.5 ± 2.1%, P < 0.01 vs. GSK, suggesting ΔΨm...
These results suggest that the active form of GSK3β opens mPTP.

GSK3β-mediated mPTP opening was further investigated by accelerating the mitoHK-II dissociation using DCCD (1 μmol/l). As shown in Fig. 2e, DCCD enhanced the GSK3β-mediated mitochondrial calcein release (Fig. 2e calcein intensity at 40 min 59.3 ± 1.5% of the baseline, P < 0.05 vs. GSK), as well as TMRE reduction (Fig. 2f TMRE intensity at 40 min 59.5 ± 1.7% of baseline, P < 0.05 vs. GSK, suggesting ΔΨm depolarization) by GSK3β. Thus, our results suggest that the sensitivity of GSK3β-mediated mPTP opening was regulated by mitoHK-II binding.

We finally explored the contribution of enzymatic activity of mitoHK-II on the GSK3β-mediated mPTP opening. Because the activity of mitoHK-II depends on its substrates (glucose Km = 0.3 mmol/l) and products
(glucose-6-phosphate G6P, $K_i = 0.02$ mmol/l) [13], an inactivation of mitoHK-II was first facilitated by substrate (glucose) depletion. Under the glucose depletion where mitoHK-II was inactivated (Supplemental fig. 4a, b), not only the mitochondrial calcein release by GSK3β (Fig. 3a calcein intensity at 35 min 91.5 ± 1.5%, $P < 0.05$ vs. with glucose), but also the TMRE reduction ΔΨm depolarization by GSK3β (Fig. 3b TMRE intensity at 30 min 103.2 ± 2.3%, $P < 0.05$ vs. with glucose) was attenuated. The inactivation of mitoHK-II was also examined by treating cells with G6P. Although it is reported that a high concentration of G6P itself dissociates HK-II from mitochondria [14], G6P did not alter mitoHK-II binding under the concentration of 0.01 and 0.1 mmol/l in our experimental condition (Fig. 3c). When permeabilized myocytes were exposed to 0.01 mmol/l of G6P, which could inactivate HK-II [13] without affecting mitochondrial binding, the GSK3β-mediated mitochondrial calcein release was attenuated (Fig. 3d, calcein intensity at

Fig. 2 GSK3β promotes mPTP opening. a, b Time course of changes in calcein a and TMRE b intensity in permeabilized myocytes, which were perfused with control internal solution (CTL; ○) and GSK3β (10 nmol/l) in the absence (GSK; ●) or presence of SB216763 (3 μmol/l, GSK + SB; △). Alamethicin (a pore-forming antibiotic Alm) was applied to obtain maximal calcein release from the mitochondrial matrix in calcein experiments, and DNP, an uncoupler, was applied at the end of TMRE experiments. Data are presented as the % of intensity at 0 min, and the values are mean ± SEM from 13 to 21 experiments. *$P < 0.05$ vs. CTL, **$P < 0.01$ vs. GSK by two-way ANOVA followed with Bonferroni’s test. c, d Summarized data of calcein c and TMRE d intensities after 30 min perfusion of GSK3β (GSK), GSK3β plus SB216763 (GSK + SB), GSK3β plus cyclosporine A (GSK + CsA; 0.1 μmol/l), and GSK3β plus PKA catalytic subunit (GSK + PKA; 10 U/ml). Some cells were pretreated with SB, CsA, or PKA for 5 min, and then GSK3β was perfused with them for 30 min. Values are mean ± SEM from 11 to 18 experiments. *$P < 0.01$ vs. control, **$P < 0.01$ vs. GSK by one-way ANOVA with Bonferroni’s test. e, f Time course of changes in calcein e and TMRE f intensities during and after perfusion with GSK3β (●; 10 nmol/l), and GSK3β plus DCCD (○; 1 μmol/l). Data are presented as the % of intensity at 0 min, and the values are mean ± SEM from 5 to 16 independent experiments. *$P < 0.05$ vs. GSK3β by two-way ANOVA. g, h Time course of changes in calcein g and TMRE h intensities during and after perfusion with GSK3β (●), and GSK3β plus dextran (○; 1%). Permeabilized myocytes were perfused with dextran for 20 min, and then GSK3β was applied. Values are mean ± SEM from 5 to 19 independent experiments. *$P < 0.05$ vs. GSK3β by two-way ANOVA followed by Bonferroni’s test.
In this study, we studied the cardioprotective effects by GSK3β inactivation against I/R injury and investigated the roles of mitoHK-II binding on the mPTP opening. The main findings of this study are as follows: (1) Pharmacological inhibition of GSK3β prevented dissociation of mitoHK-II (through glucose depletion or G6P) and opened the GSK3β-mediated mPTP opening.

**Discussion**

Thus, in contrast to mitoHK-II dissociation, inactivation of mitoHK-II (through glucose depletion or G6P) attenuated the GSK3β-mediated mPTP opening.
attenuated GSK3β-mediated mPTP opening in permeabilized myocytes.

GSK3β is a constitutive serine–threonine kinase and involved in cellular energy metabolism [4, 5, 15]. Many investigations suggested that inactivation of GSK3β plays key roles in the protective effects of ischemic and pharmacological preconditioning [4, 5], post-conditioning [16], and chemical cardioprotective interventions [17, 18]. As compatible with previous investigations, our results revealed that inactivation of GSK3β by SB216763 exhibited cardiac protection against I/R injury in Langendorff-perfused hearts (Fig. 1a, b). Although our results revealed short time effects by pharmacological inhibition, Woulfe et al. also have shown less cardiac dilatation and preserved LV function up to 8 weeks after myocardial infarction in inducible GSK-3β knockdown mice [19]. It has already been reported that the active form of GSK3β opens mPTP in response to ROS or Ca2+ overload in isolated mitochondria [16], and the inhibition of GSK3β acts as a master switch to limit the mPTP opening in cardiac myocytes [4]. As compatible with previous reports, we showed that the recombinant (active form) GSK3β opened mPTP (Fig. 2a, b) in a dose-dependent manner and half maximal concentration of mPTP opening was 2.5 nmol/l in our experimental condition (Supplemental fig. 2). In addition, the inactivation of GSK3β by SB216763 or protein kinase A catalytic subunit inhibited the GSK3β-mediated mPTP opening (Fig. 2c, d). Although we did not show direct evidence that the phosphorylated GSK3β (inactive form) inhibited mPTP opening during I/R in isolated hearts, the GSK3β-inactivation appears to suppress the susceptibility of mPTP through preventing mitoHK-II dissociation [6], attenuating the affinity of adenine nucleotide translocase (ANT, located in the inner mitochondrial membrane) to cyclophilin D (an mPTP regulatory protein located in the inner mitochondrial membrane), suppressing mitochondrial Ca2+ overload [20], and reducing ATP hydrolysis during ischemia [5]. Further investigations are required to reveal the relationship between GSK3β inactivation and mPTP inhibition in I/R injury.

HK-II is heterogeneously distributed in the mitochondria and cytosol, and mitoHK-II is physically binds external surface of outer mitochondrial membrane (OMM) through porin or voltage-dependent anion channel (VDAC; located in the OMM), which enables mitoHK-II to use mitochondrial ATP to catalyze the phosphorylation of glucose to yield G6P [6]. We showed that the GSK3β inactivation (by SB216763) attenuated the mitoHK-II dissociation by I/R in Langendorff-perfused hearts (Fig. 1), and that recombinant GSK3β (activated form) dissociated HK-II from mitochondria in a SB216763-sensitive manner in permeabilized myocytes (Fig. 2), suggesting that GSK3β dissociates mitoHK-II. Because our results exhibited net positive of total mitoHK-II by SB216763 in isolated hearts, we cannot completely deny the possibility of contribution of translocated cytosolic HK-II to mitochondria, by which total amount of mitoHK-II increased rather than the inhibition of mitoHK-II dissociation. However, previous investigations have supported our results and phosphorylated GSK3β (inactive form) can stabilize mitoHK-II binding through the suppression of VDAC phosphorylation [6, 7, 21].

We showed that the chemical stimuli (DCCD or dextran) modulating mitoHK-II bindings altered the GSK3β-mediated mPTP opening (Fig. 2e–h). Although the precise mechanisms by which mitoHK-II dissociation induces mPTP opening have not been completely understood, the instability of VDAC by mitoHK-II dissociation may be responsible [22]. In addition to its dissociation, mitoHK-II also regulated the GSK3β-mediated mPTP opening by its inactivation (Fig. 3). Because the mitoHK-II phosphorylates glucose by preferentially utilizing the intramitochondrial ATP [23], inactivation of mitoHK-II may suppress matrix ATP consumption, which is required for glucose metabolism, and may result in the inhibition of GSK3β-mediated mPTP opening. It would be difficult to apply our results to common pathophysiological conditions because we could estimate the effects of mitoHK-II inactivation only in the extreme experimental conditions, where permeabilized myocytes perfused with no glucose or abundant G6P solution. However, at least in our experimental condition, mitoHK-II regulated GSK3β-mediated mPTP not only by its dissociation but also by inactivation. Our investigation has not sufficiently revealed the relationship between the extent of HK-II inactivation and mPTP inhibition because we could not ensure the inactivation of mitoHK-II by G6P due to the technical issue (Supplemental fig. 4a, b). In fact, 0.1 mmol/l of G6P, which is expected the better HK-II inactivation than 0.01 mmol/l without affecting mitochondrial HK-II binding, failed to restore the GSK3β-mediated TMRE suppression in our experimental condition (Supplemental fig. 3a). Further investigation will be required to explore the contribution of mitoHK-II on mPTP opening under I/R.

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Compliance with ethical standards

Conflict of interest The authors declare that they have no conflicts of interest.

Ethical Approval All applicable international, national, and/or institutional guidelines for the care and use of animals in the field of Physiology Science have been followed, and studies have been approved by a research ethics committee at the Hamamatsu University School of Medicine.
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