Dual-specificity phosphatase 1 interacts with prohibitin 2 to improve mitochondrial quality control against type-3 cardiorenal syndrome

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Received: 2023.09.25; Accepted: 2023.12.22; Published: 2024.01.19

Abstract

Type-3 cardiorenal syndrome (CRS-3) is acute kidney injury followed by cardiac injury/dysfunction. Mitochondrial injury may impair myocardial function during CRS-3. Since dual-specificity phosphatase 1 (DUSP1) and prohibitin 2 (PHB2) both promote cardiac mitochondrial quality control, we assessed whether these proteins were dysregulated during CRS-3-related cardiac depression. We found that DUSP1 was downregulated in heart tissues from a mouse model of CRS-3. DUSP1 transgenic (DUSP1Tg) mice were protected from CRS-3-induced myocardial damage, as evidenced by their improved heart function and myocardial structure. CRS-3 induced the inflammatory response, oxidative stress and mitochondrial dysfunction in wild-type hearts, but not in DUSP1Tg hearts. DUSP1 overexpression normalized cardiac mitochondrial quality control during CRS-3 by suppressing mitochondrial fission, restoring mitochondrial fusion, re-activating mitophagy and augmenting mitochondrial biogenesis. We found that DUSP1 sustained cardiac mitochondrial quality control by binding directly to PHB2 and maintaining PHB2 phosphorylation, while CRS-3 disrupted this physiological interaction. Transgenic knock-in mice carrying the Phb2S91D variant were less susceptible to cardiac depression upon CRS-3, due to a reduced inflammatory response, suppressed oxidative stress and improved mitochondrial quality control in their heart tissues. Thus, CRS-3-induced myocardial dysfunction can be attributed to reduced DUSP1 expression and disrupted DUSP1/PHB2 binding, leading to defective cardiac mitochondrial quality control.

Keywords: CRS-3, DUSP1, PHB2, mitochondrial quality control

Introduction

With the prolongation of life expectancy and the increasing aging population, the proportion of patients with concurrent cardiovascular and renal diseases is increasing year by year. Acute and chronic heart diseases can directly induce or worsen renal impairment, and vice versa [1]. The bidirectional interactions between the heart and the kidneys complicate disease treatment, reduce the quality of life, prolong hospitalization, worsen the prognosis and increase mortality [2, 3]. Therefore, the concept of cardiorenal syndrome (CRS) is receiving increasing attention.

Type-3 CRS (CRS-3) is defined as acute kidney injury followed by cardiac injury and/or dysfunction [4]. Various pathophysiological processes contribute to the occurrence and development of CRS-3. Fluid retention during acute kidney injury can lead to pulmonary edema [5]. Electrolyte disturbances such
as hyperkalemia can cause arrhythmias and even cardiac arrest [6]. Untreated uremia can impair myocardial contractility by fostering myocardial depressant factor accumulation and pericarditis [7]. Metabolic acidosis can promote the progression of right heart failure by inducing pulmonary vasconstriction, and can also have negative inotropic and proarrhythmic effects [7]. In addition, renal ischemia itself can trigger inflammatory mediator production, leukocyte infiltration and apoptosis, leading to inflammation-induced myocardial injury [8]. When acute kidney injury progresses and requires renal replacement therapy, rapid changes in fluid volume and electrolyte levels can induce or exacerbate hypertension, arrhythmias and myocardial ischemia [9]. However, the molecular pathways underlying CRS-3 are not fully understood, so effective therapeutic approaches are still lacking.

Recent findings have revealed that mitochondrial dysfunction may promote the progression of heart failure during CRS-3 [8, 10, 11]. Mitochondrial injury is associated with mitochondrial oxidative stress, calcium imbalances, adenosine triphosphate (ATP) shortages and mitochondrial apoptotic pathway activation [12]. On the other hand, pharmacological interventions to enhance mitochondrial fitness have been shown to improve heart performance during CRS-3 [10, 11]. However, mitochondrial function can also be restored through the endogenous defensive mechanism known as mitochondrial quality control, which consists of mitochondrial dynamics, mitophagy and mitochondrial biogenesis [13-15]. Mitochondrial quality control can modify the mitochondrial morphology, promote mitochondrial turnover and stimulate mitochondrial metabolism [16, 17]. The beneficial effects of mitochondrial quality control on the heart have been widely reported by our laboratory [18] and other colleagues [19-21]. However, the influence of mitochondrial quality control on CRS-3 has not yet been described.

Dual-specificity phosphatase 1 (DUSP1, also known as MKP-1) is a critical enzyme involved in cellular signaling pathways [22]. Belonging to the family of dual-specificity phosphatases, DUSP1 can (de)phosphorylate threonine-serine and tyrosine residues on target proteins [23-25]. We previously demonstrated that DUSP1 is critically important for mitochondrial quality control in cardiac ischemia/reperfusion injury [26] and endotoxemia-related myocardial dysfunction [16]. Since DUSP1 is not localized to the mitochondrial surface, its impact on mitochondrial structure and function must be indirect and involve other proteins [16, 27].

Prohibitin 2 (PHB2) is a protein that primarily localizes to the inner mitochondrial membrane [28], where it maintains mitochondrial structure and function [29, 30]. In light of its enhancement of mitophagy and mitochondrial dynamics [30, 31], PHB2 has been identified as a novel inducer of mitochondrial quality control in diabetic cardiomyopathy [32] and doxorubicin-induced cardiomyopathy [33]. Importantly, phosphorylation of PHB2 can alter its interactions with other proteins, its subcellular localization and its function in mitochondrial dynamics [34, 35]. Considering that DUSP1 is a dual-specificity phosphatase, we wondered whether it might promote mitochondrial quality control by altering PHB2 phosphorylation. In this study, we explored whether cardiac dysfunction during CRS-3 was associated with dysregulated DUSP1 activity, PHB2 phosphorylation and mitochondrial quality control.

Materials and Methods

Animals and CRS-3 model

All animal procedures were performed according to guidelines and protocols approved by the Institutional Animal Care and Use Committee of Xiyuan Hospital, China Academy of Chinese Medical Sciences. DUSP1<sup>−/−</sup> [16] and Phb2<sup>S91D</sup> knock-in mice [36] on a C57BL/6 background were provided by Prof. Hao Zhou in Chinese PLA General Hospital. Mice (8-10 weeks old) were maintained under pathogen-free conditions on a 12-hour light/dark schedule with ad libitum access to water and a chow diet.

For CRS-3 model generation, anesthesia was administered via intraperitoneal injection of 1% pentobarbital at a dose of 80 mg/kg body weight. The hair was removed from both sides of the mouse’s back, and the area was disinfected with iodine. A 1-cm incision was made through the skin and muscle beside the spine to expose the left kidney. The renal artery and vein on both sides of the kidney were carefully separated, and the blood vessels were promptly clamped using artery forceps. The clamping time was noted, and the procedure was repeated for the other kidney. After the procedure, the mouse was placed on a warming pad at 37 °C. After 30 minutes of clamping, the vascular clamp was released, and the kidney was observed to ensure that it was rapidly changing color from purplish-brown to red. To prevent dehydration, 1 mL of saline was administered. The incisions on both sides were closed in layers using sutures. The mouse was kept on the warming pad for five to six hours so that its recovery could be monitored, and then it was returned to the animal facility. Samples were collected at various time points as per the experimental schedule.
Echocardiography

A Fujifilm VisualSonics Vevo 2100 Ultra High-Frequency Imaging Platform was used for the functional analysis. During echocardiography, the mice were anaesthetized with 1.5% isoflurane, with continuous monitoring of the electrocardiogram, respiratory rate and body temperature. The LVEF, FS and ventricular volumes were measured in parasternal long-axis view using the Vevo LV-Trace function. The wall thickness was measured in the parasternal long- and short-axis M-Modes. Aortic valve regurgitation was imaged using color-Doppler mode in the parasternal long-axis and suprasternal views.

Transmission electron microscopy

Transmission electron microscopy was used to monitor the myocardial ultrastructure, as described previously [37]. Briefly, a freshly collected sample from the apex of the mouse heart was dissected into 1-mm³ sections, immediately fixed with 2% glutaraldehyde in 0.1 M phosphate-buffered saline, and then fixed with 1% osmium tetroxide. After the samples were dehydrated in ethanol and embedded in Epon resin, ultrathin sections were prepared and counterstained with uranyl acetate and lead citrate. The stained sections were examined under a transmission electron microscope (JEOL1230). The mitochondrial number was counted in a total of 10 images per heart (x12,000 magnification, n=3 hearts per group).

Hematoxylin and eosin (H&E) staining

Heart tissues were fixed in 10% formalin (Thermo Fisher Scientific, Agawam, CA, USA) and stained with H&E. Images were captured on a Panoptiq Digital Slide Imaging System.

Immunofluorescence

Heart tissue sections (4 µm) and cells were fixed with phosphate-buffered 4% paraformaldehyde (0335, Carl Roth) for 10 minutes. Antibodies were diluted in Dako antibody diluent (S202230-2, Agilent Technologies). For immunofluorescence staining, the sections were incubated with primary antibodies overnight at 4 °C. After the sections were washed three times with phosphate-buffered saline (A0964.9050, VWR International, Radnor, PA, USA), they were incubated with fluorophore-conjugated secondary antibodies for one hour at room temperature. The sections were mounted with Dako fluorescence mounting medium (S302380-2, Agilent Technologies).

Cardiomyocyte isolation

Ventricular cardiomyocyte isolation was performed as previously described [38]. Mice were anesthetized with 5% isoflurane and then killed via cervical dislocation. Their hearts were rapidly extracted and placed in Tyrode’s solution containing 140 mM NaCl, 6 mM KCl, 1 mM MgCl₂, 1 mM CaCl₂, 10 mM glucose and 10 mM 4-(2-hydroxyethyl)-1-piperazineethanesulfonic acid (HEPES), adjusted to pH 7.4 with 2 mM NaOH. After aortic cannulation, the hearts were placed in a Langendorff system, perfused with Tyrode’s solution for five minutes, and then perfused for five minutes with a low-Ca²⁺ oxygenated solution containing 120 mM NaCl, 5.4 mM KCl, 5 mM MgSO₄, 5 mM sodium pyruvate, 20 mM glucose, 20 mM taurine, 10 mM HEPES, 5 mM nitrilotriacetic acid and 0.04 mM CaCl₂, adjusted to pH 6.96 with 2 mM NaOH. Finally, the hearts were perfused for 10 minutes with an enzyme solution containing 120 mM NaCl, 5.4 mM KCl, 5 mM MgSO₄, 5 mM sodium pyruvate, 20 mM glucose, 20 mM taurine, 10 mM HEPES and 0.2 mM CaCl₂, pH 7.4, with collagenase type 2 (1 mg/mL; Worthington) and hyaluronidase (0.6 mg/mL; Sigma-Aldrich). Left ventricular cardiomyocytes were plated on laminin-coated dishes and allowed to attach to the dish/cover slip for at least 45 minutes before experiments.

Analysis of contraction/relaxation kinetics

Contractile kinetics were analyzed at the indicated time points in a monolayer of cultured cardiomyocytes. In brief, cells were seeded onto Matrigel-coated 96-well plates at a density of 100,000 cells per well in high-glucose Dulbecco’s modified Eagle’s medium supplemented with 10% fetal bovine serum for 24 hours. Contraction/relaxation movement was recorded every 56 milliseconds for 16 seconds using an inverted microscope (Olympus CellIR). During the recording process, the temperature was kept at 37 °C and the atmosphere was maintained at 5% CO₂. Data were analyzed in ImageJ using the ‘MuscleMotion’ tool, as we previously described [39].

Mitochondrial membrane potential detection

For mitochondrial membrane potential quantification, cardiomyocytes were incubated in medium containing 10 μg/L JC-1 (T3168, Thermo Fisher) at 37 °C for 10 minutes. Cells were analyzed using a confocal laser scanning microscope [40].

ROS measurements

ROS were measured as we described previously [37]. In brief, cells were incubated with DCFH-DA (5 μM, Invitrogen, Eugene, OR, USA), and the fluorescence intensity was measured with excitation...
at 510 nm and emission at 580 nm using a confocal laser scanning microscope. The mean level was set at control = 1, and the fold-increase was calculated for each measurement [41].

**RNA extraction and qPCR**

Total mRNA was extracted from mouse heart tissues with a Quick RNA miniprep kit (Zymo Research, Irvine, CA, USA) according to the manufacturer's instructions. Then, qPCR was performed to detect mRNA expression using SYBR Green Master Mix (Applied Biosystems, Foster City, CA, USA) in a QuantStudio™ 6 Flex system (Applied Biosystems). The expression of each gene was normalized to that of 18S rRNA [42]. Thermal cycling was conducted with an initial hold period at 95 °C for 10 minutes, followed by 40 cycles of a three-step PCR program at 95 °C for 15 seconds, 60 °C for 1 minute and 72 °C for 30 seconds.

**Cell culture and siRNA knockdown of DUSP1**

HL-1 cells were used for the *in vitro* studies. The HL-1 cell line was purchased from the JCRB Cell Bank. Cells were cultured in Dulbecco's modified Eagle's medium (Gibco, Grand Island, NY, USA) with 10% fetal bovine serum (Gibco), 100 U/mL penicillin and 100 g/mL streptomycin (Gibco) at 37 °C in a humidified incubator containing 5% CO2. For the knockdown experiments, cells were seeded in 24-well plates in triplicate for 48 hours. *DUSP1* siRNAs (#sc-35937; Santa Cruz Biotechnology, Inc.) were transfected into the HL-1 cells with Lipofectamine™ (Santa Cruz Biotechnology, Inc.) according to the manufacturer's instructions to analyze the levels of the targeted proteins or enzymes.

**Protein extraction and Western blot analysis**

Mouse heart tissues and cells were homogenized in M-PER™ Mammalian Protein Extraction Reagent (Cat #78501, Thermo Fisher Scientific, Waltham, MA, USA) containing Halt™ Protease Inhibitor Cocktail (Cat #78429, Thermo Fisher Scientific). The protein concentrations were determined with a Pierce™ Microplate BCA Protein Assay Kit (Cat #23252, Thermo Fisher Scientific) according to the manufacturer's instructions [44, 45]. For Western blotting, the extracted proteins were boiled in Laemmli sample buffer (2X; Cat #1610737, Bio-Rad Laboratories, Hercules, CA, USA) for denaturation, and then were electrophoretically separated on sodium dodecyl sulfate polyacrylamide gels (Cat #M00654, GenScript, Piscataway, NJ, USA) and transferred onto nitrocellulose membranes using a Trans-Blot Turbo Transfer System and RTA Transfer Packs (Cat #1704271, Bio-Rad Laboratories). The membranes were blocked with 10% non-fat dry milk in Tris-buffered saline containing 0.05% Tween-20 for one to two hours at room temperature, and then were incubated with primary antibodies (summarized in Table S5) at 4 °C overnight. The membranes were then incubated with horseradish peroxidase-conjugated secondary antibodies (1:5000; Jackson Immunoresearch Laboratories Inc., West Grove, PA, USA) at room temperature for one hour. After appropriate washing, the membranes were developed with Clarity™ Western ECL Substrate (Cat #170-5061, Bio-Rad Laboratories) or Clarity™ Max Western ECL Substrate (Cat #170-5062, Bio-Rad Laboratories). The Western blots were quantified using ImageJ software (Version 1.51n, National Institutes of Health, Bethesda, MD, https://imagej.nih.gov/ij), as previously described [46].

**ELISA**

A Mouse BNP ELISA kit (#P40753, RayBiotech Life, Inc.), Mouse lactate dehydrogenase (LDH) ELISA kit (#SEB864Mu, CLOUD-CLONE CORP.), Mouse TnT ELISA kit (#LS-F8830, LifeSpan Biosciences, Inc.), Mouse CK-MB ELISA kit (#MBS265235, MyBioSource), ATP ELISA kit (#MAK190-1KT, Sigma-Aldrich), Mouse SOD ELISA kit (#E4583, BioVision), Mouse glutathione ELISA kit (#MOEB2568, AssayGenie) and Mouse CAT ELISA kit (#EKU03008, Biomatik) were used according to the manufacturers’ instructions to analyze the levels of the targeted proteins or enzymes.

**Co-immunoprecipitation**

For co-immunoprecipitation, all steps were performed at 4 °C. Cells were harvested and then solubilized with immunoprecipitation buffer (150 mM NaCl, 10% glycerol, 20 mM Tris–HCl pH 7.4, 2 mM ethylenediaminetetraacetic acid, 0.5% Nonidet P-40, 0.5% Triton X-100 and complete protease inhibitor) for 1.5 hours. Insoluble components were removed via centrifugation at 12,000 × g for 10 minutes, and the supernatant was collected. One-tenth of the supernatant was removed and used as ‘Input’, while the rest was incubated with an anti-Flag M2 affinity gel (Sigma-Aldrich) at 4 °C overnight. The beads were washed six times with lysis buffer and then boiled in sodium dodecyl sulfate sample buffer to retrieve the proteins. For western blotting, the extracted proteins were boiled in Laemmli sample buffer (2X; Cat #1610737, Bio-Rad Laboratories, Hercules, CA, USA) for denaturation, and then were electrophoretically separated on sodium dodecyl sulfate polyacrylamide gels (Cat #M00654, GenScript, Piscataway, NJ, USA) and transferred onto nitrocellulose membranes using a Trans-Blot Turbo Transfer System and RTA Transfer Packs (Cat #1704271, Bio-Rad Laboratories). The membranes were blocked with 10% non-fat dry milk in Tris-buffered saline containing 0.05% Tween-20 for one to two hours at room temperature, and then were incubated with primary antibodies (summarized in Table S5) at 4 °C overnight. The membranes were then incubated with horseradish peroxidase-conjugated secondary antibodies (1:5000; Jackson Immunoresearch Laboratories Inc., West Grove, PA, USA) at room temperature for one hour. After appropriate washing, the membranes were developed with Clarity™ Western ECL Substrate (Cat #170-5061, Bio-Rad Laboratories) or Clarity™ Max Western ECL Substrate (Cat #170-5062, Bio-Rad Laboratories). The Western blots were quantified using ImageJ software (Version 1.51n, National Institutes of Health, Bethesda, MD, https://imagej.nih.gov/ij), as previously described [46].

For the docking analysis, we used the Schrödinger 2017-1 suite (Schrödinger Inc., New York, NY, USA). The crystal structures of DUSP1 and PHB2 were obtained from available sources and prepared for docking using the Protein Preparation Wizard workflow in the Maestro package. The binding site was chosen through the Grid Generation procedure. The prepared ligand was flexibly docked into the...
receptor using Glide (XP mode) with default settings. To determine the binding affinity between DUSP1 and PHB2, we employed the Microscale Thermophoresis methodology.

Statistical analyses
All statistical analyses were conducted using GraphPad Prism 8 software. Data are presented as the mean ± standard deviation, and were compared using one-way or two-way analysis of variance. Statistical significance was set at p<0.05.

Results
DUSP1 overexpression sustains heart function after CRS-3
To explore the involvement of DUSP1 in CRS-3, we performed the CRS-3 operation on DUSP1 transgenic (DUSP1\textsuperscript{Tg}) mice. We used echocardiography to compare the heart function of sham-operated mice, wild-type (WT) CRS-3 model mice and DUSP1\textsuperscript{Tg} CRS-3 model mice (Figure 1A-G). Compared with the sham group, the WT CRS-3 group exhibited significantly reduced left ventricular ejection fraction (LVEF) and fractional shortening (FS) values, as well as impaired early to late ventricular filling velocity ratios (E/A) and early diastolic mitral inflow to early diastolic mitral annulus velocity ratios (E/e'). Interestingly, CRS-3 failed to induce heart dysfunction in DUSP1\textsuperscript{Tg} mice (Figure 1A-G).

Subsequently, we isolated single cardiomyocytes from the mice to analyze their contractile/relaxation properties. As shown in Figure 1H-M, CRS-3 impaired cardiomyocyte contractile indexes such as the peak shortening (PS), maximal velocity of shortening (+dL/dt) and time to peak shortening (TPS). CRS-3 also diminished the cardiomyocyte relaxation capacity, as evidenced by the altered maximal velocity of relengthening (-dL/dt) and time to 90% relengthening (TR90). Interestingly, CRS-3 did not damage the mechanical properties of cardiomyocytes isolated from DUSP1\textsuperscript{Tg} mice (Figure 1H-M).

We next examined structural changes in heart tissues from mice subjected to CRS-3. Histological analyses revealed myocardial fiber degradation and disorder following CRS-3 in WT mice, but not in DUSP1\textsuperscript{Tg} mice (Figure 1N). Similarly, CRS-3 upregulated serum troponin T (TrT), lactate dehydrogenase (LDH), natriuretic peptide B (BNP) and creatine kinase (CK)-MB levels in WT mice, but not in DUSP1\textsuperscript{Tg} mice (Figure 1O-R). In sum, CRS-3 downregulated DUSP1 in heart tissues, while DUSP1 overexpression suppressed CRS-3-induced myocardial dysfunction.

DUSP1 overexpression reduces the cardiac inflammatory response, oxidative stress and mitochondrial injury during CRS-3
Our previous studies have indicated that CRS-3 is characterized by oxidative stress, the inflammatory response and mitochondrial injury in the heart [10]. Thus, we asked whether DUSP1 overexpression could attenuate these pathological alterations in the presence of CRS-3. Quantitative real-time PCR (qPCR) analyses demonstrated that interleukin 6 (IL-6), tumor necrosis factor alpha (TNF\alpha) and monocyte chemoattractant protein 1 (MCP1) levels were significantly elevated in heart tissues following CRS-3, whereas DUSP1 overexpression attenuated these changes (Figure 2A-C). Immunofluorescence staining showed that, as a result of increased inflammatory factor expression, Gr-1-positive neutrophil levels were elevated within the myocardium in CRS-3-treated WT mice, but not in DUSP1\textsuperscript{Tg} mice (Figure 2D and E). These results confirmed that DUSP1 could inhibit the inflammatory response in the presence of CRS-3.

To evaluate oxidative stress in heart tissues from the different groups of mice, we performed enzyme-linked immunosorbent assays (ELISAs) to analyze the concentrations of glutathione (GSH), catalase (CAT) and superoxide dismutase (SOD). As shown in Figure 2F-H, glutathione, CAT and SOD activity levels were markedly downregulated in WT heart tissues upon CRS-3 treatment; however, DUSP1 overexpression restored glutathione, CAT and SOD levels in the myocardium. We then isolated cardiomyocytes from the mice, and used 2,7-dichlorodihydrofluorescein diacetate (DCFH-DA) staining to assess reactive oxygen species (ROS) production. Upon CRS-3 treatment, ROS levels were significantly elevated in WT cardiomyocytes, but were reduced to near-physiological levels in DUSP1\textsuperscript{Tg} cardiomyocytes (Figure 2I and J). These data confirmed that DUSP1 overexpression could neutralize cardiac oxidative stress during CRS-3.

Mitochondrial injury often features reduced mitochondrial metabolism. An ELISA of total ATP production indicated that CRS-3 suppressed ATP synthesis in WT-derived cardiomyocytes, but not in DUSP1\textsuperscript{Tg}-derived cardiomyocytes (Figure 2K). Since ATP metabolism highly depends on the mitochondrial membrane potential, we wondered whether DUSP1 overexpression influenced this parameter. The mitochondrial membrane potential was reduced upon CRS-3 treatment in WT cardiomyocytes, whereas this alteration was undetectable in DUSP1\textsuperscript{Tg} cardiomyocytes (Figure 2L and M). Overall, DUSP1 preserved mitochondrial function in the heart during CRS-3.
Figure 1. DUSP1 overexpression sustains heart function after CRS-3. DUSP1$^{tg}$ and WT mice were subjected to 30 minutes of bilateral renal artery ischemia, followed by 72 hours of reperfusion to induce CRS-3. A-G. Echocardiography was used to evaluate heart function in DUSP1$^{tg}$ and WT mice upon CRS-3. H-M. Single cardiomyocytes were isolated from DUSP1$^{tg}$ and WT mice, and their contractile/relaxation capacities were recorded. N. H&E staining was used to observe the cardiac structure following CRS-3. O-R. Serum samples were collected from DUSP1$^{tg}$ and WT mice, and the concentrations of TnT, CK-MB, BNP and LDH were determined using ELISAs. *p<0.05.
Figure 2. DUSP1 reduces the inflammatory response, oxidative stress and mitochondrial injury in the heart during CRS-3. DUSP1<sup>Tg</sup> and WT mice were subjected to 30 minutes of bilateral renal artery ischemia, followed by 72 hours of reperfusion to induce CRS-3. A-C. RNA was collected from DUSP1<sup>Tg</sup> and WT hearts, and qPCR was used to evaluate the transcription of IL-6, TNFα and MCP1. D, E. Double immunofluorescence staining was used to measure Gr-1-positive neutrophil accumulation in the myocardium. TnT was used to visualize myocardial fibers, and 4′,6-diamidino-2-phenylindole (DAPI) was used to stain the nuclei. F-H. ELISAs were used to evaluate the concentrations of glutathione (GSH), SOD and CAT in the heart. I, J. Cardiomyocytes were isolated from DUSP1<sup>Tg</sup> and WT mice, and ROS levels were determined using DCFH-DA. K. ATP production in the myocardium was determined with an ELISA. L-M. An immunofluorescence assay was used to determine the mitochondrial membrane potential in cardiomyocytes isolated from DUSP1<sup>Tg</sup> and WT mice after CRS-3. *p<0.05.
**DUSP1 overexpression maintains mitochondrial quality control in the heart during CRS-3**

Mitochondrial quality control, including mitochondrial dynamics, mitophagy and mitochondrial biogenesis, is an endogenous protective mechanism that sustains mitochondrial function [15, 47, 48]. Thus, we wondered whether DUSP1 protected cardiac mitochondria during CRS-3 by improving mitochondrial quality control. A qPCR analysis of mitochondrial dynamics proteins revealed that CRS-3 significantly upregulated dynamin-related protein 1 (Drp1) and mitochondrial fission 1 (Fis1) in heart tissues from WT mice, but not from DUSP1Tg mice (Figure 3A-D). On the other hand, mitochondrial fusion factors such as mitofusin 2 (Mfn2) and optic atrophy 1 (Opa1) were markedly downregulated following CRS-3 in WT heart tissues, but returned to near-normal levels in DUSP1Tg heart tissues (Figure 3A-D). Thus, CRS-3 activated mitochondrial fission and inhibited fusion in the heart, whereas DUSP1 overexpression normalized these processes. In accordance with these findings, immunofluorescence staining of cardiomyocyte mitochondria revealed that CRS-3 caused mitochondrial fragmentation in WT-derived cardiomyocytes (Figure 3E-G). However, the mitochondrial structure and network remained intact in DUSP1Tg-derived cardiomyocytes, despite treatment with CRS-3 (Figure 3E-G).

In addition to mitochondrial dynamics, we also noted that mitophagy was significantly suppressed by CRS-3 in WT heart tissues, as evidenced by reduced expression of Parkin, FUN14 domain-containing 1 (Fundc1) and Beclin1 (Figure 3H-J). Interestingly, in DUSP1Tg heart tissues, CRS-3 failed to suppress mitophagy activity (Figure 3H-J).

The transcriptional process of mitochondrial biogenesis is highly regulated by peroxisome proliferator activated receptor gamma coactivator 1 alpha (PGC1α) and its downstream effectors – nuclear factor erythroid 2-related factor 2 (Nrf2) and transcription factor A, mitochondrial (Tfam). A qPCR analysis demonstrated that CRS-3 inhibited mitochondrial biogenesis gene expression in WT hearts, but not in DUSP1Tg hearts (Figure 3K-M). These data demonstrated that CRS-3 impaired cardiac mitochondrial quality control, while DUSP1 normalized it.

**DUSP1 binds to PHB2 to sustain its phosphorylation in the heart during CRS-3**

Recent studies have indicated that PHB2 promotes mitochondrial quality control by suppressing mitochondrial fission and inducing mitophagy. Interestingly, in heart tissues from CRS-3 model mice, the transcription of PHB2 was unaltered compared with the control group (Figure 4A), so total PHB2 levels remained constant; however, PHB2 phosphorylation was attenuated following CRS-3 (Figure 4B and C). PHB2 phosphorylation in the heart returned to near-normal levels in response to DUSP1 overexpression, suggesting that DUSP1 governs the phosphorylation status of PHB2 in the setting of CRS-3 (Figure 4B and C).

To determine how DUSP1 sustains PHB2 phosphorylation, we analyzed the possibility of a protein-protein interaction between the two. We first performed a molecular docking assay, which revealed potential binding sites between DUSP1 and PHB2 (Figure 4D-F). Co-immunoprecipitation experiments further confirmed the presence of crosslinks between DUSP1 and PHB2 (Figure 4G). To verify that DUSP1 is required to preserve PHB2 phosphorylation, we transfected HL-1 cells with siRNA against DUSP1 under physiological conditions. Knocking down DUSP1 reduced PHB2 phosphorylation (Figure 4H). These results demonstrated that DUSP1 binds directly to PHB2 to maintain PHB2 phosphorylation, while the downregulation of DUSP1 during CRS-3 contributes to PHB2 dephosphorylation.

**Transgenic mice expressing a Phb2S91D phosphorylation mutant gene are resistant to CRS-3**

To test whether PHB2 dephosphorylation is a pathological factor that impairs cardiac mitochondrial quality control and promotes myocardial dysfunction in CRS-3, we used transgenic knock-in mice carrying the Phb2S91D variant on a C57BL/6 background, as we previously described [36]. The LVEF, FS, E/A, LVDd, LVSd, and E/e' values were also improved in Phb2S91D mice compared with WT mice following CRS-3 (Figure 5A-G).

Subsequently, we isolated single cardiomyocytes from the mice to analyze their contractile/relaxation properties. As shown in Figure 5H-M, the contractile indexes (PS, +dL/dt and TPS) that were disrupted by CRS-3 in WT cardiomyocytes were restored to near-normal levels in Phb2S91D cardiomyocytes. Likewise, CRS-3 impaired the relaxation capacity of WT cardiomyocytes, but not of Phb2S91D cardiomyocytes (Figure 5H-M).

We then performed a histological analysis, which showed that CRS-3 failed to induce myocardial fiber degradation or disorder in Phb2S91D heart tissues (Figure 5N). Similarly, although CRS-3 upregulated TnT, LDH, BNP and CK-MB in WT serum samples, these changes were reversed in Phb2S91D mice (Figure 5O-R). Thus, preserving PHB2 phosphorylation seemed to protect the heart against CRS-3-induced myocardial injury.
Enhancing PHB2 phosphorylation attenuates mitochondrial quality control disorder, oxidative stress and the inflammatory response in the heart following CRS-3

We then assessed whether PHB2 phosphorylation protected the heart during CRS-3 by altering mitochondrial quality control, oxidative stress or the inflammatory response. We found that CRS-3 upregulated Drp1/Fis1 and downregulated Mfn2/Opa1 in WT heart tissues, while these changes were significantly suppressed in Phb2^{S91D} heart tissues (Figure 6A-D). Similarly, CRS-3 significantly inhibited cardiac Nrf2 and Tfam expression in WT mice, but not in Phb2^{S91D} mice (Figure 6E-F).

In vitro, immunofluorescence staining indicated that ROS generation was augmented in WT cardiomyocytes upon CRS-3 exposure, while it was reduced in Phb2^{S91D} cardiomyocytes in the setting of CRS-3 (Figure 6G and H). ELISAs demonstrated that CRS-3 reduced the activity of glutathione, CAT and SOD in WT heart, but not in Phb2^{S91D} heart (Figure 6I-K).
Figure 4. DUSP1 binds to PHB2 to sustain its phosphorylation in the heart during CRS-3. DUSP1\(^{Tg}\) and WT mice were subjected to 30 minutes of bilateral renal artery ischemia, followed by 72 hours of reperfusion to induce CRS-3. A. RNA was collected from DUSP1\(^{Tg}\) and WT hearts, and qPCR was used to evaluate the transcription of PHB2. B, C. Proteins were isolated from DUSP1\(^{Tg}\) and WT hearts, and Western blotting was used to analyze PHB2 expression. D, E. Molecular docking analysis was used to evaluate the potential binding sites between DUSP1 and PHB2. F, G. Co-immunoprecipitation was used to analyze the crosslinks between DUSP1 and PHB2. H, I. HL-1 cells were transfected with siRNA against DUSP1, and the phosphorylation of PHB2 was determined via Western blotting. *p<0.05.
Figure 5. Transgenic mice expressing a Phb2S91D phosphorylation mutant gene are resistant to CRS-3. Phb2S91D knock-in and WT mice were subjected to 30 minutes of bilateral renal artery ischemia, followed by 72 hours of reperfusion to induce CRS-3. A-G. Echocardiography was used to evaluate heart function in Phb2S91D knock-in and WT mice upon CRS-3. H-M. Single cardiomyocytes were isolated from Phb2S91D knock-in and WT mice, and their contractile/relaxation capacities were recorded. N. H&E staining was used to observe the cardiac structure following CRS-3. O-R. Serum samples were collected from Phb2S91D knock-in and WT mice, and the concentrations of TnT, CK-MB, BNP and LDH were determined with ELISAs. *p<0.05.
Figure 6. Enhancing PHB2 phosphorylation attenuates cardiac mitochondrial quality control disorder, oxidative stress and the inflammatory response during CRS-3. Phb2<sup>S91D</sup> knock-in and WT mice were subjected to 30 minutes of bilateral renal artery ischemia, followed by 72 hours of reperfusion to induce CRS-3. A-F. RNA was collected from Phb2<sup>S91D</sup> knock-in and WT hearts, and qPCR was used to evaluate the transcription of Drp1, Fis1, Mfn2, Opa1, Nrf2 and Tfam. G, H. Cardiomyocytes were isolated from Phb2<sup>S91D</sup> knock-in and WT mice, and ROS levels were determined using DCFH-DA. I-K. ELISAs were used to evaluate the concentrations of glutathione (GSH), SOD and CAT in the heart. L, M. Double immunofluorescence staining was used to measure Gr-1-positive neutrophil accumulation in the myocardium. TnT was used to visualize myocardial fibers, and DAPI was used to stain the nuclei. N-P. RNA was collected from Phb2<sup>S91D</sup> knock-in and WT hearts, and qPCR was used to evaluate the transcription of IL-6, TNFα and MCP1. *p<0.05.
Double immunofluorescence staining of the myocardium revealed that CRS-3 induced Gr-1-positive neutrophil accumulation within WT heart tissues, while this inflammatory response was undetectable in Phb2<sup>S91D</sup> mice (Figure 6L and M). Accordingly, CRS-3 failed to upregulate pro-inflammatory genes in heart tissues from Phb2<sup>S91D</sup> mice (Figure 6N-P). These data illustrated that PHB2 phosphorylation prevented CRS-3-related myocardial injury by improving mitochondrial quality control, reducing oxidative stress and repressing the inflammatory response in the heart.

Discussion

Under normal conditions, an organ-organ feedback system maintains the balance between the heart and the kidneys. The heart perfuses the organs by continuously pumping fresh blood, while the kidneys reabsorb water from urine and regulate the blood volume through neuroendocrine control, thus decisively influencing the cardiac output. However, under pathological conditions, dysfunction in one organ can lead to dysfunction in the other, either acutely or chronically. CRS refers to abnormalities in one organ that can lead to dysfunction in the other, either acutely or chronically. CRS refers to acute cardiac dysfunction caused by acute kidney injury.

Compared with other types of CRS, CRS-3 has been a neglected area of research, so many academic viewpoints and experimental hypotheses regarding its pathogenesis remain unverified. Among these research conclusions, the widely accepted pathogenic mechanism of CRS-3 is that acute kidney injury causes hemodynamic abnormalities that lead to cardiac dysfunction. Specifically, when the kidneys are injured, inadequate renal perfusion activates the renin-angiotensin-aldosterone system, stimulates the sympathetic nervous system and promotes vasopressin secretion, leading to fluid retention, an increased preload and exacerbated pump dysfunction [2, 50]. However, cardiac preload and afterload abnormalities and sympathetic nervous system activation can only partially explain the pathophysiological changes in type-3 CRS [3]. In clinical practice, the use of vasoactive drugs has not completely improved the symptoms of CRS-3 patients [51, 52], suggesting that non-hemodynamic factors may also be involved in the pathogenesis of this disease [53, 54]. Among these factors, inflammatory response activation, myocardial substrate metabolism conversion, mitochondrial fission and imbalanced ROS/nitric oxide production have been proposed as potential pathophysiological contributors to CRS-3 [49, 55-57]. However, there is still insufficient research to explain the relationships among these molecular events or determine their common upstream signaling pathways systematically.

In the present study, we found that CRS-3-induced myocardial dysfunction was associated with defective mitochondrial quality control, including increased mitochondrial fission, reduced mitochondrial fusion, diminished mitophagy and suppressed mitochondrial biogenesis in the heart. Abnormal mitochondrial quality control has been reported as a pathological factor in several cardiovascular diseases, including cardiac ischemia/reperfusion injury [18, 58], septic cardiomyopathy [17, 59] and diabetic cardiomyopathy [14, 60]. Increased mitochondrial fission and reduced mitochondrial fusion are thought to activate mitochondrial apoptosis by two main mechanisms [61, 62]. First, an imbalance between mitochondrial fission and fusion promotes the formation of mitochondrial fragments containing broken or incomplete mitochondrial DNA, so the mitochondria-encoded respiratory complex enzymes are transcriptionally downregulated [61]. This accelerates mitochondrial ROS production, which induces cardiolipin oxidation and cytochrome c release from mitochondria into the cytoplasm [61, 63]. Second, excessive fission and defective fusion induce the disassociation of hexokinase-II from mitochondrial dria, thus promoting mitochondrial permeability transition pore opening [64]. Through these two pathways, mitochondrial dynamics disorder not only induces mitochondrial dysfunction, but also promotes cardiomyocyte apoptosis.

In addition to altered mitochondrial dynamics, impaired mitophagy and mitochondrial biogenesis have been associated with reduced mitochondrial turnover [38, 65] and the accumulation of poorly-structured mitochondria within cardiomyocytes. Although previous studies have described the involvement of mitochondria in myocardial dysfunction during CRS-3 [8, 10, 11], this is the first study to delineate the molecular pathway by which both mitochondrial function and structure are disrupted during CRS-3. Our findings suggest that normalizing mitochondrial quality control is critical for preserving cardiomyocyte mitochondrial function and cell viability in the setting of CRS-3.

Importantly, our study revealed that DUSP1 downregulation and PHB2 dephosphorylation are the molecular mechanisms accounting for mitochondrial quality control disorder in the heart upon CRS-3. Ample studies have elucidated the cardioprotective effects of DUSP1. DUSP1 deficiency was reported to exacerbate cardiac ischemia/reperfusion injury by activating mitochondrial fission through mitochondrial fission factor in a manner dependent on the c-Jun N-terminal kinase pathway [26]. DUSP1
overexpression promoted valosin-containing protein dephosphorylation in the heart, thus improving mitochondrial quality control, mitochondrial integrity and heart function during septic cardiomyopathy [16, 27]. In the present study, we found that DUSP1 bound directly to PHB2 to maintain its phosphorylation, while CRS-3 stress downregulated DUSP1, thereby reducing PHB2 phosphorylation as an initial signal of abnormal mitochondrial quality control in the heart. Interestingly, previous studies have indicated that dephosphorylated PHB2 is associated with mitochondrial dysfunction or cell apoptosis [35, 36]. We found that sustaining PHB2 phosphorylation in vivo enhanced the resistance of the myocardium to CRS-3 by reducing the inflammatory response, inhibiting oxidative stress and improving mitochondrial quality control, ultimately improving heart function.

This study had several limitations. First, additional in vitro data are necessary to illuminate the protein-protein interaction between DUSP1 and PHB2. Second, it remains unknown how acute kidney injury downregulates DUSP1 in the myocardium. It will be interesting to determine whether neuromodulators, serum factors/cytokines or microRNAs allow the injured kidneys to communicate with the myocardium in the pathogenesis of CRS-3.

In conclusion, our results demonstrated that CRS-3 induced heart failure by suppressing cardiac DUSP1 expression. DUSP1 overexpression effectively maintained heart function by suppressing the inflammatory response, inhibiting oxidative stress and improving mitochondrial quality control. DUSP1 bound directly to PHB2 to prevent its dephosphorylation; thus, overexpressing DUSP1 or enhancing PHB2 phosphorylation reduced the vulnerability of the heart to CRS-3. Based on our findings, therapeutic approaches to enhance DUSP1 expression, PHB2 phosphorylation or mitochondrial quality control could improve the treatment of CRS-3-related myocardial depression.

Supplementary Material

Acknowledgements
This study is supported by the Young Elite Scientists Sponsorship Program by CAST (No. 2022QNRC001) and the China Academy of Chinese Medical Sciences Innovation Fund (No. CI2021A01405).

Competing Interests
The authors have declared that no competing interest exists.

References


