Mitochondrial Reactive Oxygen Species Mediate Cardiac Structural, Functional, and Mitochondrial Consequences of Diet-Induced Metabolic Heart Disease

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Background—Mitochondrial reactive oxygen species (ROS) are associated with metabolic heart disease (MHD). However, the mechanism by which ROS cause MHD is unknown. We tested the hypothesis that mitochondrial ROS are a key mediator of MHD.

Methods and Results—Mice fed a high-fat high-sucrose (HFHS) diet develop MHD with cardiac diastolic and mitochondrial dysfunction that is associated with oxidative posttranslational modifications of cardiac mitochondrial proteins. Transgenic mice that express catalase in mitochondria and wild-type mice were fed an HFHS or control diet for 4 months. Cardiac mitochondria from HFHS-fed wild-type mice had a 3-fold greater rate of \( \text{H}_2\text{O}_2 \) production \((P=0.001 \text{ versus control diet fed})\), a 30% decrease in complex II substrate–driven oxygen consumption \((P=0.006)\), 21% to 23% decreases in complex I and II substrate–driven ATP synthesis \((P=0.01)\), and a 62% decrease in complex II activity \((P=0.002)\). In transgenic mice that express catalase in mitochondria, all HFHS diet–induced mitochondrial abnormalities were ameliorated, as were left ventricular hypertrophy and diastolic dysfunction. In HFHS-fed wild-type mice complex II substrate–driven ATP synthesis and activity were restored ex vivo by dithiothreitol (5 mmol/L), suggesting a role for reversible cysteine oxidative posttranslational modifications. In vitro site-directed mutation of complex II subunit B Cys100 or Cys103 to redox-insensitive serines prevented complex II dysfunction induced by ROS or high glucose/high palmitate in the medium.

Conclusion—Mitochondrial ROS are pathogenic in MHD and contribute to mitochondrial dysfunction, at least in part, by causing oxidative posttranslational modifications of complex I and II proteins including reversible oxidative posttranslational modifications of complex II subunit B Cys100 and Cys103. \( (\text{J Am Heart Assoc.} 2016;5:e002555 \text{ doi:} 10.1161/JAHA.115.002555) \)

Key Words: metabolic heart disease • mitochondria • obesity • oxidative protein modifications • oxidative stress

One-third of Americans have obesity-related metabolic syndrome with type 2 diabetes, hypertension, and dyslipidemia.\(^1\)\(^-\)\(^3\) An important consequence of metabolic syndrome is metabolic heart disease (MHD), which is associated with left ventricular (LV) hypertrophy, diastolic dysfunction, and impaired cardiac energetics.\(^4\)\(^-\)\(^7\) We previously showed that feeding mice a high-fat, high-sucrose (HFHS) diet for 8 months leads to obesity, metabolic syndrome, and a cardiac phenotype typical of MHD with LV hypertrophy, diastolic dysfunction, and cardiac mitochondrial dysfunction.\(^8\)\(^,\)\(^9\)

Several findings in HFHS-fed mice suggest a central role for reactive oxygen species (ROS) in the development of MHD. Markers of oxidative stress are increased in the myocardium,\(^8\) and oxidative posttranslational modifications (OPTMs) are found in multiple target proteins, many of which are in the mitochondria.\(^10\) A role for ROS is further supported by our demonstration in isolated mitochondria from HFHS-fed mice that there is impaired complex II–driven ATP generation, which is restored ex vivo by dithiothreitol (DTT).\(^3\) The cardiac phenotype of hypertrophy and diastolic dysfunction is also prevented by polyphenols,\(^8\) which may exert antioxidant...
effects. Finally, ROS production is increased in isolated cardiac mitochondria from these mice,9 suggesting that mitochondria may be a source of ROS with HFHS feeding.

Together, these observations led to our thesis that mitochondrial ROS play a key role in mediating MHD in HFHS-fed mice. These observations, though suggestive, are circumstantial. A primary goal of the present study was to test the thesis in vivo using an intervention that is specific for mitochondrial ROS. Accordingly, in mice fed a HFHS diet for 4 months, we studied the effects of decreasing mitochondrial ROS by transgenic expression of mitochondrial catalase on (1) ATP production and respiration in isolated cardiac mitochondria and (2) cardiac structural and functional remodeling. The mechanistic role of cysteine oxidation was assessed ex vivo in isolated mitochondria by examining the effects of exposure to a reducing environment on complex II oxidation and function. The role of specific oxidative cysteine modifications of complex II in mediating dysfunction was assessed in vitro by site-directed mutation to redox-insensitive serines in HEK-293T cells exposed to ROS or high glucose/high palmitate in the medium.

Methods

Experimental Animals

Mitochondrially targeted catalase overexpressing (mCAT) and wild-type (WT) littermate control mice (C57BL/6NJ background) 9 weeks of age were fed ad libitum either a control chow diet (CD; Research Diets, product No. D09071703, 10% kcal lard, 0% sucrose) or an HFHS diet (Research Diets, product No. D09071702; 58% kcal lard, 13% kcal sucrose) for 4 months. mCAT mice were obtained from the Jackson Laboratory [strain: B6.Cg-Tg(CAG-OTC/CAT)4033Prab/J; stock No. 016197]. These mice were originally developed by Dr Peter Rabinovitch’s group (University of Washington), and their characteristics are published.11 Diet compositions are published with minor modifications.9,14,15 All steps were performed off-line with the use of a customized version of Vevo 770 Analytic software.

Mitochondrial Isolation

Heart mitochondria were isolated as previously described by us with minor modifications.9,14,15 All steps were performed at 4°C. Briefly, tissues were rinsed in a buffer containing 100 mmol/L KCl, 5 mmol/L EGTA, and 5 mmol/L HEPES, pH 7.0, and thereafter homogenized in 2 mL of HES buffer (HEPES 5 mmol/L, EDTA 1 mmol/L, sucrose 0.25 mol/L, pH 7.4 adjusted with KOH 1 mol/L) by using a Teflon-on-glass electric homogenizer. The homogenate was centrifuged at 500g for 10 minutes at 4°C. The supernatant was then centrifuged at 9000g for 15 minutes at 4°C, and the mitochondrial pellet was resuspended in 100 μL of HES buffer with 0.3% of BSA fatty acid free. Protein was quantified with the use of BCA (Pierce), and the value of HES-BSA buffer
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Mitochondrial H$_2$O$_2$ Production

Mitochondrial H$_2$O$_2$ production was measured by using the Amplex Ultra Red horseradish peroxidase method (Invitrogen) as we described previously with minor modifications. This assay is based on the horseradish peroxidase (2 units/mL) H$_2$O$_2$-dependent oxidation of nonfluorescent Amplex Ultra Red (50 μmol/L) to fluorescent resorufin red. In short, 10 μg mitochondria was diluted in 50 μL of reaction buffer (125 mmol/L KCl, 10 mmol/L HEPES, 5 mmol/L MgCl$_2$, 2 mmol/L K$_2$HPO$_4$, pH 7.44) to determine complex I (pyruvate/malate, 5 mmol/L) or complex II (succinate, 5 mmol/L)–driven H$_2$O$_2$ production with and without inhibitor (rotenone 2 μmol/L). Mitochondrial H$_2$O$_2$ production was measured after the addition of 50 μL of reaction buffer containing horseradish peroxidase and Amplex Ultra Red. Fluorescence was followed at an excitation wavelength of 545 nm and an emission wavelength of 590 nm for 20 minutes. The slope of the increase in fluorescence is converted to the rate of H$_2$O$_2$ production with the use of a standard curve. All of the assays were performed at 25°C. The results are reported as picomoles per minute per milligrams of protein.

ATP Production in Isolated Mitochondria

ATP synthesis rates in isolated heart mitochondria were determined by using the luciferin/luciferase–based ATP Bioluminescence Assay Kit CLS II (Roche) as we previously described with minor modifications. In short, 10 μg of heart mitochondria was suspended in 75 μL of buffer A (125 mmol/L KCl, 10 mmol/L HEPES, 5 mmol/L MgCl$_2$, and 2 mmol/L K$_2$HPO$_4$, pH 7.44) to determine complex I (pyruvate/malate, 5 mmol/L final) or complex II (succinate, 5 mmol/L final)–driven ATP synthesis. Following standard practice, succinate-driven ATP generation was measured in the presence of complex I inhibitor rotenone (2 μmol/L) to avoid the reverse electron transfer effect. The assays were performed in the presence and absence of DTT 5 mmol/L. Measurements with substrates were repeated in the presence of oligomycin, an inhibitor of ATP synthase, to determine the rates of nonmitochondrial ATP production. The background of the assay was determined with mitochondrial values. With use of an ATP standard provided in the kit, the slopes were converted in nanomoles per minute per milligrams of protein.

Mitochondrial Electron Transport Chain Complex II Activity

Complex II enzyme activity of isolated mitochondria was measured by using a microplate assay kit (Abcam/Mitosciences ab109908/MS241), as we previously described. In this assay kit, complex II is immunocaptured within the wells of the microplate. The production of ubiquinol by complex II is coupled to the reduction of the dye DCPIP (2,6-dichlorophenolindophenol), and decreases in its absorbance at 600 nm are measured spectrophotometrically. The assay is performed in the presence of succinate as a substrate. The assay was performed in the presence and absence of 5 mmol/L DTT. Enzymatic activity was normalized to mitochondrial protein concentration.

Mitochondrial Oxygen Consumption Rate

Oxygen consumption rates were monitored by using a Seahorse XF24 oxygen flux analyzer as we previously described. Isolated mitochondria were loaded in a 24-well Seahorse plate on ice (5–12.5 μg/well) and 500 μL of ice-cold mitochondrial assay solution (MAS: 70 mmol/L sucrose, 220 mmol/L mannitol, 5 mmol/L KH$_2$PO$_4$, 5 mmol/L MgCl$_2$, 2 mmol/L HEPES, 1 mmol/L EGTA, 0.3% BSA fatty acid free, pH 7.4) were added on top. The 4 sequential injection ports of the Seahorse cartridge contained the following: port A, 50 μL of 10× substrate (complex I: 50 mmol/L pyruvate and 50 mmol/L malate; complex II: 50 mmol/L succinate and 20 μmol/L rotenone in MAS) and 2.5 mmol/L ADP; port B, 55 μL of 20 μmol/L oligomycin; port C, 60 μL of 40 μmol/L Carbonyl cyanide-4-(trifluoromethoxy)phenylhydrazone (FCCP); and port D 65 μL of 40 μmol/L antimycin A. State III was determined after port A injection, state IV after port B, and uncoupled after port C. Antimycin A was used as a control because it blocks the electron transport chain to minimize mitochondrial oxygen consumption. The results are reported as pmol oxygen per minute per microgram of protein.

Immunoblotting for Mitochondrial Proteins

Immunoblots were performed on frozen LV that was homogenized in tissue lysis buffer (HEPES, pH 7.4, 20 mmol/L, B-glycerol phosphate 50 mmol/L, EGTA 2 mmol/L, DTT 1 mmol/L, NaF 10 mmol/L, NaVO$_4$ 1 mmol/L, Triton X-100 1%, glycerol 10%, and 1 protease inhibitor complete mini tablet, EDTA free, 20 mL [Roche]). Total protein (25 μg) was
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Separated via sodium dodecyl sulfate–polyacrylamide gel electrophoresis and transferred to polyvinylidene fluoride membranes. Blots were incubated with mouse total OXPHOS Rodent WB Antibody cocktail (containing 5 antibodies, 1 each against complex I subunit [NDUFB8], complex II subunit B [SDHB], complex III core protein 2 [UQRC2], complex IV subunit I [MTCO1], and complex V alpha subunit [ATP5A]) (Abcam; ab110413), anti-VDAC1 (Abcam; ab15895), and anti-GAPDH (Abcam; ab8245) and detected with the use of the Licor Odyssey 2-color infrared imaging system.

Biotin Switch Assay

Labeling with biotin iodoacetamide (BIAM; Life Technologies; B-1591) was used in a biotin switch assay to detect reversibly oxidized cysteines, as we previously described with minor modifications.9,17 Freshly isolated LV sections were immediately snap-frozen and stored in liquid nitrogen. Tissue was thawed and lysed in RIPA buffer containing 100 mmol/L N-ethylmaleimide (NEM) to block all free thiols and prevent further oxidation. Excess NEM was removed by passing the lysates over Zeba spin columns. NEM-labeled proteins are treated with 5 mmol/L DTT at room temperature for 15 minutes to reduce reversible oxidations to free thiols and again column cleaned to eliminate DTT. Samples are incubated with 4 mmol/L BIAM for 2 hours at room temperature in the dark, labeling any reactive free thiols that were previously oxidized. BIAM-labeled proteins are isolated from total protein pool with the use of streptavidin magnetic beads (50 μL) for 1 hour at room temperature in the dark, washed, and cleaved from beads by boiling in 30 μL of 2× nonreducing Laemmli buffer for 10 minutes. Entire BIAM-labeled and unlabeled protein fractions are subjected to Western blotting and probed with anti-SDHB (Abcam; ab14714) antibodies, and bands were detected by using near-infrared dye-conjugated secondary antibodies and quantified with the Licor Odyssey 2-color infrared imaging system. Ratio of labeled to total (labeled plus unlabeled) proteins represents percentage of reversibly oxidized cysteines in SDHB.

Construction of SDHB Cysteine Mutants and Expression in HEK-293T Cells

Full-length mouse SDHB construct in pCMV-sport6 was obtained from American Type Culture Collection (ATCC; MGC: 19177; IMAGE: 4225025) and used as the template for site-directed mutagenesis. The Cys→Ser mutants were prepared by introducing a single base exchange (Cys100Ser, TGT→TCT; Cys103Ser, TGT→TCT; Cys115Ser, TGT→TCT) with use of the QuikChange II XL site-directed mutagenesis kit. Mutants were confirmed by DNA sequencing (Tufts Medical Center, Sequencing Core, Boston, MA). Constructs or empty vector, serving as control, were transfected into HEK-293T cells by using polyethyleneimine “MAX” transfection reagent (Polysciences, Inc).

In Vitro Exposure to H2O2 or High-Glucose/High-Palmitate Medium

Cells were then cultured for an additional 42 to 48 hours in DMEM supplemented with 10% fetal bovine serum. At the end of that period, cells were treated with either 500 μmol/L H2O2 or vehicle for 10 minutes. In other experiments, cells were cultured for 14 hours in high-glucose/high-palmitate (HGHP) medium (25 mmol/L glucose, 0.4 mmol/L palmitic acid, 0.67% BSA) or control media (5 mmol/L glucose, 0.67% BSA), as previously described.17 Cells were then lysed, and complex II activity was measured as described earlier with the use of the Abcam/Mitosciences kit ab109908/MS241.

Statistical Analysis

Data are presented as mean±SEM. Comparisons between groups were performed by using 1-way ANOVA with Bonferroni posttests for multiple comparisons. All statistical analyses were performed with GraphPad Prism version 6 software (GraphPad Software Inc). A value of P<0.05 was considered statistically significant.

Results

Mitochondrial H2O2 Production Is Increased After 4 Months of HFHS Diet and Corrected in mCAT Mice

In cardiac mitochondria from WT mice fed an HFHS diet versus the CD, H2O2 production was increased with either complex I (pyruvate+malate; Figure 1A) or complex II (succinate+rotenone; Figure 1B) substrates. The HFHS-induced increases in H2O2 production with both complex I and II substrates were prevented in mCAT mice (Figure 1).

Effects of HFHS Diet and mCAT Expression on Maximal ATP Synthesis Rate

In mitochondria from HFHS-fed WT mice, the maximal rates of ATP synthesis were decreased by 21% and 23% for complex I (Figure 2A) and complex II (Figure 2B) substrates, respectively. The HFHS-induced decreases in ATP synthesis rates for complex I and complex II substrates were both prevented in mCAT mice (Figure 2). The HFHS-induced decreases in complex I and II function were not associated with a change in the protein expression of the representative subunits of complexes I, II, III, IV, or V (Table S2). Thus, decreased
complex I and II ATP production in HFHS-fed mice is not the result of decreased expression of complex I or II proteins.

**Effects of HFHS Diet and mCAT on Mitochondrial Oxygen Consumption**

To further assess the mechanism responsible for decreased ATP production with HFHS feeding, oxygen consumption rate was measured in isolated mitochondria by using a Seahorse XF24 oxygen flux analyzer. In WT mice, complex I–mediated respiration linked to maximal ATP synthesis rate (state III) and proton leak (state IV) both tended to be lower in HFHS-fed mice but did not reach statistical significance (Figure 3A and 3B). Likewise, complex I substrate state III and IV oxygen consumption rates tended to be higher in HFHS-fed mCAT (versus WT) mice but did not reach statistical significance (Figure 3A and 3B). Both complex II–mediated state III and state IV, oxygen consumption rates were significantly decreased in mitochondria from HFHS-fed WT mice, and the HFHS-mediated decreases in both state III and state IV oxygen consumption rates were prevented in mCAT mice (Figure 3C and 3D). The parallel decreases in state III and IV respiration with HFHS feeding suggests that the observed decreases in complex I and II–mediated ATP production are the result of impaired electron transport chain function rather than proton leak. The ability of mCAT to correct both ATP production and respiration suggests that catalase-sensitive...
ROS mediates the observed electron transport chain dysfunction. To control for mitochondrial integrity we derived respiratory control ratios for state III/IV oxygen consumption rate values. There were no differences in respiratory control ratios between CD, HFHS, or mCAT groups (Figure S2).

Differential Effects of DTT on Complex I and II–Mediated ATP Production

We previously observed that with 8 months of HFHS feeding, complex II dysfunction is reversed by DTT ex vivo, while complex I dysfunction is not.9 Likewise, in mitochondria from HFHS-fed WT mice, preincubation with DTT (5 mmol/L) had no effect on complex I substrate–mediated ATP synthesis (Figure S3A) but restored complex II substrate–mediated ATP synthesis to that of WT CD-fed mice (Figure S3B). In mitochondria from mCAT mice, DTT had no effect on either complex I or complex II substrate–mediated ATP synthesis rate, regardless of diet (Figure S4A and S4B). These observations suggest that mCAT fully normalizes complex I and II function in HFHS-fed mice.

The HFHS Diet—Induced Decrease in Complex II Activity Is Reversible

We further assessed complex II directly by measuring the reduction of ubiquinone to ubiquinol. In WT mice, HFHS feeding decreased complex II activity by 62% (Figure 4A). DTT completely restored complex II activity in HFHS-fed mice (Figure 4A). The HFHS-induced decrease in complex II activity was prevented in mCAT mice (Figure 4B), and accordingly, DTT had no further effect on complex II activity in HFHS-fed mCAT mice (Figure 4C). These observations confirm that the HFHS feeding–induced decrease in complex II function, as reflected by ATP production, is prevented by mCAT in vivo and reversed by DTT ex vivo.

HFHS Feeding Causes Reversible Cysteine OPTMs in Complex II

We previously used iodoacetamide-based cysteine-reactive tandem mass tags to identify 3 cysteines in SDHB that undergo reversible oxidation with HFHS feeding for 8 months.9,10 To determine whether reversible cysteine
oxidation occurs in the SDHB subunit of complex II with HFHS feeding for only 4 months, we performed a biotin-switch assay in LV homogenates. Free thiols were blocked with N-ethylmaleimide; reversibly oxidized thiols were then reduced with DTT (as in the ATP and complex II activity assays), labeled with BIAM and immunoblotted for SDHB. The quantity of reversibly oxidized thiols in SDHB was increased by HFHS feeding, and this increase was prevented in mCAT mice (Figure 5). Thus, ≥1 reversible OPTMs of complex II subunit B are associated with HFHS feeding–induced complex II dysfunction and are prevented by mCAT.

SDHB Redox-Insensitive Mutants Protect Complex II Activity From ROS- and HGHP-Mediated Inhibition

To test whether cysteine OPTMs could mediate the complex II inhibition in HFHS-fed mice, we used HEK-293T cells as a model system. Exposure of HEK cells to H$_2$O$_2$ (500 μmol/L,
10 minutes) caused an ≈55% decrease complex II activity, which was restored by the ex vivo addition of (DTT 5 mmol/L) (Figure 6A), consistent with the DTT-reversible inhibition of complex II function that we observed with HFHS feeding (see Figure 4). Based on our prior observation that Cys100, Cys103, and Cys115 undergo reversible oxidation in HFHS-fed mice, we used site-directed mutagenesis to generate SHDB constructs in which these cysteines are replaced by serine and therefore no longer redox sensitive. Baseline complex II activity in the HEK cells transfected with SDHB mutants was not different from that in HEK cells transfected with a control empty vector (Figure 6B through 6D). HEK cells transfected with the Cys100Ser and Cys103Ser SDHB mutants were each protected from the H2O2-mediated decrease in complex II activity (Figure 6B and 6C), whereas HEK cells transfected with Cys115Ser mutant remained susceptible (Figure 6D).

To further assess the relevance of the effects of H2O2 on complex II activity, HEK-293T cells were incubated in media containing high concentrations of glucose (25 mmol/L) and palmitate (0.4 mmol/L). As with H2O2, there was an ≈60% decrease in complex II activity, which was fully reversed by DTT exposure ex vivo (Figure 7A). As with H2O2, the HGHP-mediated decrease in complex II activity was prevented in cells expressing the Cys100Ser and Cys103Ser mutants but not Cys115Ser mutant (Figure 7B through 7D).

mCAT Prevents HFHS Diet–Mediated LV Hypertrophy and Diastolic Dysfunction

As we previously observed, WT mice fed an HFHS diet developed LV hypertrophy as reflected by increased LV weight/tibia length (Table) and increased LV wall thickness (Table; Figure 8A). The HFHS diet–induced increase in LV wall thickness in WT mice (0.27±0.02 mm) was less in mCAT mice (0.09±0.008 mm; P<0.01 versus WT HFHS) (Figure 8A). Likewise, although the heart weight tends to be increased in mCAT

Figure 6. Oxidation of succinate dehydrogenase B (SDHB) Cys100 and Cys103 inhibits mitochondrial complex II activity. Exposure of cells to H2O2 (500 μmol/L, 10 minutes) decreased complex II activity, measured as per Figure 4, in HEK 293T cells. SDHB in which Cys100Ser (C100S), Cys103Ser (C103S), or Cys115Ser (C115S) was mutated to a redox-insensitive serine was expressed in HEK 293T cells, and complex II activity was measured after exposure to H2O2. A, H2O2 decreased complex II activity in HEK cells transfected with empty vector (CTRL), and activity was restored after incubation of cell lysate with dithiothreitol (DTT; 5 mmol/L). B, Expression of the SDHB C100S mutant prevents the H2O2-mediated decrease in complex II activity. C, Expression of the SDHB C103S mutant prevents the H2O2-mediated decrease in complex II activity. D, Expression of the SDHB C115S mutant has no effect on the H2O2-mediated decrease in complex II activity. Values are mean±SEM; n=5; **P<0.01 vs CTRL; ###P<0.01 vs CTRL+H2O2.
mice (versus WT) on the CD, the HFHF diet-induced increase in LV weight/tibia length in WT mice (1.6±0.07 mg/mm) was less in mCAT mice (0.5±0.03 mg/mm; P<0.01 versus WT HFHS).

While systolic function was preserved in HFHS-fed mice, there was LV diastolic dysfunction as assessed by both transmitral flow velocity (E/A; Table) and myocardial peak early diastolic velocity (Eₘ; Table; Figure 8B). The HFHS diet-induced decreases in both E/A and Eₘ were prevented in mCAT mice (Table; Figure 8B). Thus, in mCAT mice the structural and functional effects of HFHS feeding were ameliorated.

Discussion
In the present study we show that in mice with diet-induced obesity mitochondrial ROS impair mitochondrial electron transport via irreversible OPTMs of complex I and reversible cysteine OPTMs of complex II. We further demonstrate that reversible oxidation of complex II subunit B Cys100 and Cys103, for which we previously identified OPTMs in HFHS-fed mice, causes reversible inhibition of enzyme activity. Finally, we show that in HFHS-fed mice scavenging mitochondrial ROS inhibits cardiac hypertrophy and improves diastolic function.

HFHS Feeding Causes Irreversible Inhibition of Complex I and Reversible Inhibition of Complex II
We previously showed that HFHS feeding for 8 months caused inhibition of complex I that was not reversed by DTT ex vivo. In contrast, inhibition of complex II was partially reversed by DTT ex vivo and associated with reversible cysteine OPTMs as identified by iodoacetamide-based cysteine-reactive tandem mass tag labeling and with a biotin switch assay. In the present study, mice were fed the

Figure 7. Oxidation of succinate dehydrogenase B (SDHB) Cys100 and Cys103 inhibits mitochondrial complex II activity. Exposure of cells to high-glucose/high-palmitate (HGHP) media (14 hours) decreased complex II activity, measured as per Figure 4, in HEK 293T cells. SDHB in which Cys100Ser (C100S), Cys103Ser (C103S), or Cys115Ser (C115S) was mutated to a redox-insensitive serine was expressed in HEK 293T cells, and complex II activity was measured after exposure to HGHP media. A, HGHP treatment decreased complex II activity in HEK cells transfected with empty vector (CTRL), and activity was restored after incubation of cell lysate with dithiothreitol (DTT; 5 mmol/L). B, Expression of the SDHB C100S mutant prevents the HGHP-mediated decrease in complex II activity. C, Expression of the SDHB C103S mutant prevents the HGHP-mediated decrease in complex II activity. D, Expression of the SDHB C115S mutant has no effect on the HGHP-mediated decrease in complex II activity. Values are mean±SEM; n=3 to 4; *P<0.05 vs CTRL; †P<0.05 vs CTRL+HGHP; ‡P<0.01 vs CTRL+HGHP.
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Table. Body and Organ Weights and Echocardiographic Parameters

|                              | WT/CD       | WT/HFHS     | mCAT/CD     | mCAT/HFHS   |
|------------------------------|-------------|-------------|-------------|-------------|
| Body weight, g               | 29.5±3.1    | 47.5±1.0*   | 29.5±4.2    | 46.9±2.5†   |
| Heart weight, mg             | 116.5±4.0   | 144.6±7.4*  | 131.6±13.0  | 151.3±12.7  |
| Tibia length, mm             | 18.2±0.4    | 17.9±0.1    | 17.7±0.3    | 18.0±0.3    |
| Heart weight/body weight, mg/g| 4.12±0.37   | 3.05±0.13   | 4.56±0.24   | 3.22±0.16†  |
| Heart weight/tibia length, mg/mm | 6.5±0.3    | 8.1±0.4*    | 7.4±0.6    | 7.9±0.5     |

Values are mean±SEM; n=4 to 7. CD indicates control diet; E/A, ratio of early-to-late diastolic mitral inflow velocity; E/Em, ratio of peak early mitral inflow velocity to myocardial peak early diastolic velocity; EDD, end-diastolic dimension; Ees, myocardial peak early diastolic velocity; ESD, end-systolic dimension; FS, fractional shortening; HFHS, high-fat high-sucrose diet; LV, left ventricular; mCAT, mitochondrial catalase; TWT, total wall thickness; WT, wild-type.

HFHS diet for only 4 months, which resulted in mitochondrial dysfunction and a cardiac phenotype that was similar to that observed at 8 months except that the inhibition of complex II at 4 months was fully reversible ex vivo by DTT but only partially reversible at 8 months.9 As with HFHS feeding for 8 months, there was no change in the expression of electron transport chain proteins. Taken together, these observations suggest that complex II dysfunction with HFHS feeding is caused initially (eg, 4 months) by ≥1 reversible cysteine OPTM. With more prolonged exposure (eg, 8 months) to HFHS diet, this reversible OPTM may be superseded by irreversible and/or noncysteine OPTMs.

mCAT Prevents HFHS-Induced Complex I and II Dysfunction

HFHS-induced dysfunction of both complexes I and II, as reflected by respiration and ATP production, was completely prevented by mCAT in vivo, and subsequent exposure to DTT ex vivo had no further effect. Based on the ability of mCAT to protect mitochondrial function, we conclude that ROS play an important role in mediating mitochondrial dysfunction in this model of MHD.

We cannot determine the relative contributions of complex I and II OPTMs to overall mitochondrial dysfunction, nor can we exclude a role for OPTMs of other proteins involved in electron transport chain function. For example, complex V is inhibited by glutathiolation in the setting of chronic heart failure.19 However, because DTT did not increase complex I substrate–mediated ATP production, which also requires complex V, the effects of DTT and mCAT are unlikely to be caused by increased complex V activity. Likewise, although uncoupling protein 3 can be regulated by oxidative cysteine modification,20 we found that proton leak-driven respiration for both complex I and complex II was decreased in HFHS-fed mice and restored by mCAT. Thus, in contrast to genetic models of obesity and/or diabetes (ob/ob and db/db mice),21,22 uncoupling does not play a major role in HFHS-fed mice.

The mCAT mouse used in this study is also protected from the cardiac effects of aortic constriction23 and aging.11 In mice with reduced expression of mitofusin II, mitochondrial function and cardiac phenotype were also improved by mCAT expression sufficient to normalize mitochondrial ROS, although supersuppression of mitochondrial ROS by a higher level of mCAT was deleterious.24 In this regard, the beneficial effects of mCAT on mitochondrial function in the present study were associated with normalization of mitochondrial ROS measured ex vivo in isolated mitochondria. While this observation suggests that mitochondrial ROS are suppressed in the mCAT mouse, it does not allow conclusions about the extent of its suppression in vivo.

Cysteine OPTM Inhibit Complex II Function

Complex II, which couples the oxidation of succinate to fumarate in the matrix with the reduction of ubiquinone in the membrane,25 consists of 4 nuclear-encoded proteins—SDH
subunits A, B, C, and D. SDHA and SDHB are exposed to the matrix, while SDHC and SDHD are in the inner membrane. To evaluate the role of cysteine OPTM in regulating complex II function, we used HEK-293 cells in culture. In this system, exposure to both H₂O₂ and high concentrations of glucose and palmitate in the media inhibited complex II activity, and importantly, the inhibition was reversed by the subsequent ex vivo exposure to DTT and was prevented in cells overexpressing the Cys100 and Cys103 mutants—thus replicating the effects of H₂O₂. This observation in an in vitro model of nutrient excess thus supports the relevance of this mechanism to the mitochondrial dysfunction we observed in HFHS-fed mice.

Of note, mutations of some, but not all, cysteines involved in the iron–sulfur cluster of fumarate reductase, an enzyme that is related to SDH, decreased enzyme activity and altered growth in bacteria. We previously observed reversible oxidation of SDHA Cys89 in HFHS-fed mice. However, S-glutathiolation of rat SDHA at Cys90 (Cys89 in the mouse proteome) increases complex II activity, and therefore is not likely to be the cause of decreased complex II activity with HFHS feeding.

A limitation of this study is that it does not address the relative importance of impaired complex II activity in mediating overall mitochondrial dysfunction in this model. The importance of complex II for cardiac function has been well established under conditions of ischemia–reperfusion, where succinate accumulation drives increased mitochondrial ROS, and in human genetic cardiomyopathies resulting from complex II deficiency. Modeling of succinate dehydrogenase flux supports the notion of its important contribution to cardiac ATP generation. Additionally, while pyruvate levels are not thought to be increased in cardiac hypertrophy, succinate is increased, suggesting that its use may be increased.

Mitochondrial ROS Mediate Cardiac Dysfunction in Obesity-Related MHD

Our observation of increased mitochondrial ROS production with an HFHS diet is consistent with several prior reports. A possible role for ROS in mediating myocardial hypertrophy and diastolic dysfunction with HFHS feeding was raised by our initial study in which polyphenols effectively prevented the cardiac phenotype. However, polyphenols have pleiotropic actions that preclude this conclusion. In the present study, mCAT inhibited the HFHS feeding–associated development of LV hypertrophy and largely prevented diastolic dysfunction. Heart size assessed as heart weight normalized to tibia length or LV wall thickness by echocardiography tended to be larger in mCAT (versus WT mice) fed

Figure 8. Left ventricular hypertrophy and diastolic dysfunction induced by high-fat high-sucrose (HFHS) diet feeding for 4 months is prevented in mitochondrial catalase (mCAT) mice. A, Total wall thickness. B, Myocardial peak early diastolic velocity (Eₘ). Values are mean±SEM; n=4 to 5; **P<0.01 vs wild-type (WT) control diet (CD); ***P<0.001 vs WT CD; #P<0.05 vs WT HFHS; ###P<0.001 vs WT HFHS.

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Because H$_2$O$_2$ is freely permeable through biologic membranes, mitochondrial ROS may also mediate LV structure and function via extramitochondrial signaling cascades that could affect multiple targets and pathways involved in the regulation of myocyte growth, fibrosis, and excitation–contraction coupling. In addition, mCAT mice are protected from the CD. However, the HFHS-induced increases in both heart weight and wall thickness were decreased in mCAT (versus WT) mice. Likewise, HFHS diet–induced LV diastolic dysfunction was improved in mCAT (versus WT) mice as assessed by both transmitral and tissue Doppler examinations. These findings, taken together with the demonstration that mitochondrial ROS production is increased with HFHS feeding, suggest that mitochondrial ROS plays a key role in mediating these key aspects of the cardiac phenotype in MHD.

The mechanism by which mitochondrial ROS mediate LV hypertrophy and diastolic dysfunction in HFHS-fed mice remains to be determined (Figure 9). It is possible that the mitochondrial effects of ROS shown here contribute to diastolic dysfunction by decreasing the supply of ATP necessary for sarcoplasmic reticulum Ca$^{2+}$-ATPase–mediated reuptake of calcium into the sarcoplasmic reticulum. Because H$_2$O$_2$ is freely permeable through biologic membranes, mitochondrial ROS may also mediate LV structure and function via extramitochondrial signaling cascades that could affect multiple targets and pathways involved in the regulation of myocyte growth, fibrosis, and excitation–contraction coupling. In addition, mCAT mice are protected from insulin resistance induced by a high-fat diet, which might correct the delivery of energy substrates. Our findings suggest that these features of cardiac phenotype exhibit a plasticity that is mediated by diet-related metabolic factors and that may be ameliorated by dietary intervention.

Conclusion

Catalase targeted to the mitochondria ameliorated both mitochondrial dysfunction and the cardiac phenotype (LV hypertrophy and diastolic dysfunction) caused by HFHS feeding. These structural and functional effects were associated with protection of complexes I and II from OPTMs, including reversible OPTMs of complex II that mediate complex II dysfunction in vitro. In this regard, we identified 2 cysteines in SDHB (Cys100 and Cys103) that regulate complex II function in a redox-dependent manner. These findings suggest that efforts to decrease mitochondrial ROS or their effects on target proteins may be of value in the therapy of MHD.

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Disclosures

None.

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