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Journal Title: Journal of Molecular and Cellular Cardiology
Volume: Volume 116
Publisher: Elsevier | 2018-03-01, Pages 29-40
Type of Work: Article | Post-print: After Peer Review
Publisher DOI: 10.1016/j.yjmcc.2018.01.011
Permanent URL: https://pid.emory.edu/ark:/25593/tnsx7

Final published version: http://dx.doi.org/10.1016/j.yjmcc.2018.01.011

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Accessed September 20, 2019 8:44 PM EDT
Hydrogen Sulfide Regulates Cardiac Mitochondrial Biogenesis via the Activation of AMPK

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Abstract

Background—Hydrogen sulfide (H₂S) is an important regulator of mitochondrial bioenergetics, but its role in regulating mitochondrial biogenesis is not well understood. Using both genetic and pharmacological approaches, we sought to determine if H₂S levels directly influenced cardiac mitochondrial content.

Results—Mice deficient in the H₂S-producing enzyme, cystathionine γ-lyase (CSE KO) displayed diminished cardiac mitochondrial content when compared to wild-type hearts. In contrast, mice overexpressing CSE (CSE Tg) and mice supplemented with the orally active H₂S-releasing prodrug, SG-1002, displayed enhanced cardiac mitochondrial content. Additional analysis revealed that cardiac H₂S levels influenced the nuclear localization and transcriptional activity of peroxisome proliferator-activated receptor γ coactivator 1α (PGC1α) with higher levels having a positive influence and lower levels having a negative influence. Studies aimed at evaluating the underlying mechanisms found that H₂S required AMP-activated protein kinase (AMPK) to induce PGC1α signaling and mitochondrial biogenesis. Finally, we found that restoring H₂S levels with SG-1002 in the setting of heart failure increased cardiac mitochondrial content.

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6. Disclosures
SG-1002 was provided by Sulfagenix, Inc.

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content, improved mitochondrial respiration, improved ATP production efficiency, and improved cardiac function.

**Conclusions**—Together, these results suggest that hydrogen sulfide is an important regulator of cardiac mitochondrial content and establishes that exogenous hydrogen sulfide can induce mitochondrial biogenesis via an AMPK-PGC1α signaling cascade.

**Graphical Abstract**

![Graphical Abstract Image]

**Keywords**

hydrogen sulfide; mitochondria; heart; AMPK

**1. Introduction**

Mitochondria occupy an important position as mediators of cellular homeostasis due to their role in the regulation of fuel utilization, calcium storage, intracellular signaling, and cell death [1, 2]. As such, impairments in mitochondrial function lead to the development of various disorders, such as neurodegenerative disease, cancer, aging, diabetes, and heart failure [1]. The heart is particularly susceptible to impairments in mitochondrial function given its limited regenerative capacity and persistent energy requirements [2]. As a result, the mitochondrial quality control system, consisting of mitophagy, fission and fusion, and biogenesis, is critically important in maintaining the fidelity of the heart under physiological and pathological conditions [1–3].

The mitochondrial biogenic response in the heart is tightly regulated by a complex network orchestrating both nuclear and mitochondrial genome transcription and replication [3]. This system coordinates both genomes during cardiac development and in response to physiological stimuli when there are changes in substrate availability and energetic demands [3]. Peroxisome proliferator-activated receptor γ coactivator 1α (PGC-1α) is a master regulator of mitochondrial biogenesis and energy expenditure [4, 5]. Cardiac PGC-1α is
induced at birth when the heart undergoes a dramatic shift in fuel preference from relying on glucose and lactate during the fetal period to the use of fatty acids (FA) after birth [6]. PGC-1α regulates the activity of a number of transcription factors, including, peroxisome proliferator-activated receptor-α (PPARα), estrogen receptor–related α (ERRα) and nuclear respiratory factor 1 (NRF1) [5]. By regulating the transcriptional activities of these proteins, PGC-1α modulates genes involved in mitochondrial biogenesis and metabolic pathways. Mitochondrial content is significantly reduced in the failing hearts of both rodents and humans [7, 8]. Furthermore, downregulation of PGC-1α signaling has also been observed in the setting of experimental heart failure [9]. As such, understanding the mechanisms by which PGC-1α signaling is regulated in the heart could lead to the development of therapies aimed at inducing mitochondrial biogenesis and augmenting energy production in the setting of increased contractile demand [8].

Hydrogen sulfide (H2S) is a critically important physiological gaseous signaling molecule that regulates a multitude of biological processes, including angiogenesis, proliferation, redox balance, inflammation, and cell death [10]. It is produced enzymatically in all mammalian species via the actions of cysteine metabolic enzymes: cystathionine γ-lyase (CSE), cystathionine β-synthase (CBS), and 3-mercaptopyruvate sulfutransferase (3MST) [10, 11]. Although all three enzymes are expressed in the cardiovascular system, prevailing data indicates that CSE plays the foremost role in cardiovascular physiology [10]. Numerous proteins and pathways have been identified as cellular targets of H2S. However, a common cellular target for many studies aimed at understanding the biology and therapeutic potential of H2S has been the mitochondria. H2S has a dual affect on mitochondrial bioenergetics with low concentrations serving as electron donors to the electron transport chain and higher concentrations serving as inhibitors of cytochrome c oxidase [12]. H2S also influences the levels/activation of a number of proteins related to mitochondrial biogenesis (PGC1α [13, 14]; AMP-activated protein kinase (AMPK) [15, 16]; endothelial nitric oxide synthase (eNOS) [11, 17, 18]) and there is evidence that mitochondrial content is higher in brains [13] and hearts [19] treated with exogenous H2S. While these studies provided evidence for elevated mitochondrial levels in response to H2S treatment, it was not clear if the observed increase was due to a direct effect of H2S or was simply an indirect consequence of H2S altering injury. Therefore, the main goal of the current study was to address this issue by determining if H2S levels directly influence cardiac mitochondrial content under non-stressed conditions. Additionally, we sought to gain insights into the mechanisms by which H2S induces mitochondrial biogenesis in the setting of myocardial ischemia-reperfusion.

2. Materials and Methods

2.1. Animals

The following strains of mice on a C57BL/6J background were utilized in this study: (1) C57BL/6J (Jackson Labs, Bar Harbor, ME), (2) Cardiac specific cystathionase-γ-lyase transgenic (CSE Tg*), (3) Cystathionase-γ-lyase deficient (CSE KO), (4) AMPKα2 floxed (Stock#: 014142, Jackson Labs, Bar Harbor, ME), (5) αMHC-Cre transgenic (Stock#: 011038, Jackson Labs, Bar Harbor, ME), and (6) endothelial nitric oxide synthase deficient mice (eNOS KO; Stock# 002684, Jackson Labs, Bar Harbor, ME). CSE Tg* were generated.
by ligating the full-length *Mus musculus* cystathionine γ-lyase cDNA to the murine α-myosin heavy chain promoter, followed by injection of the DNA into newly fertilized mouse embryos (FVB/n background) [20]. The mice were then backcrossed to C57BL/6J for 9 generations. Global CSE KO knockout mice were generated by replacing exon 1 (including the ATG start codon), exon 2, and exon 3 with a neomycin selection cassette [11]. The mice were then backcrossed to C57BL/6J for 9 generations. Cardiac specific AMPKα2 deficient mice (αMHC-Cre⁺ x AMPK ineffective) were generated by breeding AMPK knockout mice with αMHC-Cre⁺ mice. In all experiments, Wild-Type (WT) littermates were used as controls. Male mice between the ages of 8–10 weeks were utilized. All experimental protocols were approved by the Institute for Animal Care and Use Committee at T3 Laboratories and conformed to the Guide for the Care and Use of Laboratory Animals, published by the National Institutes of Health (NIH Publication No. 86-23, revised 1996), and with federal and state regulations.

2.2. Patient Samples

Left ventricular samples were procured from patients with advanced ischemic heart failure undergoing a heart transplant at Emory University in accordance with Institution Review Board protocols. Additional non-failing heart failure samples were obtained from LifeLink. All patient identifiers were removed to strictly maintain donor confidentiality and anonymity. Both sample sets included male and female patients (Supplemental Table 1).

2.3. Materials

The orally active H₂S-releasing prodrug, SG-1002, was provided by Sulfagenix (Cleveland, OH). SG-1002 was administered to mice in the diet (Purina 5001; Research Diets Inc, New Brunswick, NJ) to achieve a dose of 20 mg/kg/day [11]. Mice received the diet for 4 weeks. Control mice received standard chow (Chow; Purina 5001) for the same duration. For the in vitro experiments, H₂S was administered as sodium sulfide (Na₂S; Sigma Aldrich).

2.4. In vitro cell culture

H9c2 cardiomyocytes were purchased from ATCC (Rockville, MD, USA). Cells were grown in ATCC-formulated Dulbecco’s Modified Eagle’s Medium (DMEM; catalog# 30-2002) with 10% fetal bovine serum (FBS). Cells were maintained in this media until 80% confluent. Cells were then maintained in DMEM with 0.5% FBS for 12 hours. Cells were then exposed to 100 μM of Na₂S for 10, 20, 40, or 60 minutes. Additional groups of cells were exposed to 100 μM of Na₂S for 3 consecutive days.

2.5. Cellular Fractionation and Western blot analysis

Whole cell, cytosolic, and nuclear fractions were obtained from heart homogenates as previously described [21]. Protein concentrations were measured with the DC protein assay (Bio-Rad Laboratories, Hercules, CA, USA). Equal amounts of protein were loaded into lanes of Criterion™ TGX (Tris-Glycine eXtended) Stain-Free PAGE gels (BioRad). The gels were electrophoresed and activated using a ChemiDoc MP Visualization System (BioRad). The protein was then transferred to a PVDF membrane. The membranes were then imaged using a ChemiDoc MP Visualization System to obtain an assessment of proper transfer and to obtain total protein loads. The membranes were then blocked and probed with primary
antibodies (Supplemental Table 2) overnight at 4 °C. Immunoblots were next processed with secondary antibodies (Cell Signaling) for 1 hour at room temperature. Immunoblots were then probed with a Super Signal West Dura kit (Thermo Fisher Scientific) to visualize signal, followed by visualization using a ChemiDoc MP Visualization System (BioRad). Data was analyzed using Image Lab (BioRad). The total protein images were used as loading controls. For each protein of interest, the portion of the protein load image corresponding to the molecular weight of the protein of interest was used as the loading control [16].

2.6. mRNA and qPCR

RNA was isolated using the RiboPure kit according to the manufacturer’s instructions (Ambion). Reverse transcription was performed in a standard fashion with QuantiTect Reverse Transcription Kit (QIAGEN) supplemented with DNase treatment. Taqman qPCR was carried out according to the manufacturer’s instructions using probe sets obtained from Thermo Fisher Scientific.

2.7. Electron Microscopy

Heart tissue was dissected along the muscle fiber while immersed in 2.5% glutaraldehyde buffered with 0.1 M sodium cacodylate (pH 7.2). Samples were stored in the fixative overnight at 4 °C. Samples were then washed with the same buffer and post-fixed in 1% buffered osmium tetroxide, dehydrated through a graded ethanol series to 100%, and embedded in Eponate 12 resin. Ultrathin sections were cut on a Leica UC6rt ultra-microtome at 70–80 nm and counter-stained with 4% aqueous uranyl acetate and 2% lead citrate. Sections were examined using a Hitachi H-7500 transmission electron microscope equipped with a Gatan BioScan CCD camera.

2.8. Citrate Synthase Activity

Cardiac citrate synthase activity was measured spectrophotometrically in homogenates [22].

2.9. Sulfhydration

Protein sulfhydration was evaluated using the modified biotin switch assay or the maleimide assay [23]. For the maleimide assay, sulfhydration was evaluated by subtracting the red fluorescent intensity of samples treated with DTT from the red fluorescent intensity of samples without DTT treatment. The values were then normalized to input and presented as a percentage. For this assay the loss of red fluorescent in the presence of DTT indicates sulfhydration.

2.10. Sulfide Measurements

Hydrogen sulfide and sulfane sulfur levels were measured in heart tissue as previously described [24]. The amount of H₂S is reported as nmol/mg wet weight.

2.11. Immunoprecipitation

Heart homogenates were immunoprecipitated with an antibody to PGC-1α using the Dynabeads® Protein G Immunoprecipitation Kit according to manufacturer’s instructions.
The samples were then subjected to standard Western blot techniques and the membranes probed with antibodies to phosphoserine and acetyllysine.

2.12. AMPK Activity

The activity of AMPK was measured in homogenates prepared from heart tissue. The samples were first immunoprecipitated with a specific anti-AMPKα2 antibody (abcam). An aliquot of the immunoprecipitated samples were incubated in a reaction buffer containing 12.5 mM Tris-HCl (pH 7.5), 2.5 mM β-glycerophosphate, 1 mM dithiothreitol, 0.05 mM Na3VO4, 5 mM MgCl2, 0.050 mM ATP, and 0.2 mM of SAMS (AMPK synthetic substrate peptide). The rate of ADP formed from the incorporation of ATP in the synthetic peptide was then measured with the ADP-Glo Kinase Assay kit (Promega) according to the manufacturer’s instructions. Activity was expressed as ADP generated (in picomoles) per minute per milligram of protein.

2.13. Sirt1 Activity

The activity of cardiac Sirt1 was evaluated using the SIRT1 Activity Assay (catalog# ab156065, abcam).

2.14. PP2A Activity

The activity of cardiac PP2A was evaluated using the PP2A Immunoprecipitation Phosphatase Assay Kit (catalog# 12-313, Millipore Sigma).

2.15. Nitric Oxide Metabolite Analysis

Nitrite levels were quantified by ion chromatography (ENO20 Analyzer, Eicom). Tissue nitrosothiol compounds were quantified by using group-specific reductive denitrosation by iodine-iodide with subsequent detection of the liberated NO by using gas-phase chemiluminescence [25].

2.16. Adenosine Metabolite Analysis

Adenosine metabolites were evaluated in cardiac tissue using the ATP/ADP/AMP Assay Kit (Cat #: A-125, Biomedical Research Service Center, University at Buffalo-SUNY).

2.17. Mitochondria Respiration and ATP

Cardiac fibers were isolated and permeabilized with saponin as previously described [26]. Respiration was monitored using a Clark-type oxygen electrode (Hansatech Instruments, Amesbury, MA) in the presence of pyruvate or palmitoyl-L-carnitine. To evaluate ATP synthesis, aliquots were taken from the respiration chamber over a 1-minute period after the addition of ADP. ATP was then quantified with a bioluminescence assay using an ATP determination kit (A-22066; Molecular Probes, Eugene, OR). The rate of ATP synthesis was then normalized to the oxygen consumption rates measured over the time the aliquots were collected to obtain a measure of ATP synthesis efficiency (ATP/Oxygen ratio). This measurement reflects the ratio of state 3 (ADP stimulated respiration) ATP synthesis rates to state 3 oxygen consumption. A higher value indicates better efficiency.
2.18. Myocardial Ischemia-Reperfusion Protocol and Myocardial Injury Assessment

Mice were subjected to surgical ligation of the left coronary artery followed by reperfusion for 2 weeks. Echocardiography was performed as previously described [19].

2.19. Statistics

All data are expressed as mean ± SEM. The data was first evaluated for normal distribution using the D’Agostina and Pearson omnibus normality test. Subsequent, statistical significance was evaluated as follows: (1) unpaired Student t-test for comparison between 2 means; (2) a 1-way ANOVA with a Tukey test as the posthoc analysis for comparison among 3 or more groups; and (3) a 2-way ANOVA with a Bonferroni test as the posthoc analysis for comparison among the means from groups of WT and AMPK KO mice and WT and eNOS KO mice. For the echocardiography data, a 2-way repeated measures ANOVA with a Bonferroni test as the posthoc analysis was used. The following comparisons were made separately: (1) baseline vs. post-baseline measurements for each group, (2) differences between each groups baseline measurements, and (3) differences between each groups post-baseline measurements. The p-value for these evaluations was adjusted by applying the Bonferroni correction for multiple comparisons. A value of p < 0.05 denoted statistical significance and p-values were two-sided. All statistical analysis was performed using Prism 5 (GraphPad Software Inc).

3. Results

3.1. Endogenous H$_2$S Influences Cardiac Mitochondrial Content

We sought to determine if there was a direct relationship between cardiac H$_2$S levels and mitochondrial content. First, we used a genetic model to address this question. We collected heart samples from mice with significantly elevated levels of H$_2$S (CSE Tg$^+$) and from mice with attenuated levels of H$_2$S (CSE KO) (Supplemental Figure 1). Analysis revealed that the hearts of CSE Tg$^+$ mice displayed an increase in mitochondrial DNA levels and citrate synthase activity (Figure 1A–B). In contrast, both were decreased in the hearts of CSE KO mice. Together these findings suggest that H$_2$S levels influence cardiac mitochondrial content. This was further confirmed by analysis of electron microscopy images of cardiac ventricles from CSE Tg$^+$ and CSE KO mice demonstrating an increase and decrease in the number of mitochondria per field, respectively (Figure 1C–D).

Alterations in mitochondrial content can arise from biogenesis or from fusion and fission. Our data suggest biogenesis. A series of experiments were, therefore, undertaken to determine the contribution of fusion-fission to the altered mitochondrial content observed in the hearts of CSE Tg$^+$ and CSE KO mice (Supplemental Figure 2). First, the expression of proteins involved in mitochondrial fusion and fission were evaluated. The expression of the fusion proteins, mitofusin-1 (Mfn1), mitofusin-2 (Mfn2), and Opa-1, as well as the fission protein, fission-1 (Fis1) were not altered in the hearts of CSE Tg$^+$ or CSE KO mice. Second by electron microscopy, we observed no differences in the area or perimeter of the mitochondria in the hearts of each strain (Figure 1E). Finally, further analysis of mitochondrial fusion-fission was achieved by calculating the percentage of mitochondria in a given field that fell into three size categories based on area: <0.6 μm$^2$, 0.6 μm$^2$ – 1.0 μm$^2$, ...
and >1.0 μm² [27]. In support of the overall area and perimeter calculations, this analysis also revealed no differences in the hearts of each strain (Figure 1F). Overall these results suggest that H₂S levels influence cardiac mitochondrial content via biogenesis.

To characterize the mitochondrial functional phenotype of hearts from Wild-type, CSE Tg⁺ and CSE KO mice in a manner that would detect potential differences in mitochondrial volume density or function, respiration (oxygen consumption) experiments were performed on saponin-permeabilized myocardial fibers. This technique allowed for the selective use of different metabolic substrates to define the maximal respiratory capacity of specific mitochondrial oxidative pathways [26]. Experiments were performed with LV fibers using pyruvate and palmitoyl-carnitine. For these experiments, basal respiration was assessed followed by maximal ADP-stimulated state 3 respiration. LV fibers from CSE Tg⁺ mice did not display any changes in state 3 respiration rates in the presence of pyruvate (Figure 1G). They did however display increased state 3 respiration rates in the presence of palmitoyl-carnitine (Figure 1H). In contrast, LV fibers from CSE KO mice displayed increased state 3 respiration rates in the presence of pyruvate and decreased rates in the presence of palmitoyl-carnitine. Additionally, maximal rates of ATP synthesis from ADP were normalized to state 3 respiration rates to determine the efficacy of ATP synthesis in the presence of each substrate (Figure 1I). LV fibers from CSE Tg⁺ mice displayed enhanced ATP/oxygen ratios in the presence of palmitoyl-carnitine. In contrast, fibers from CSE KO mice displayed lower ATP/oxygen ratios in the presence of both pyruvate and palmitoyl-carnitine.

3.2. Exogenous H₂S Induces Cardiac Mitochondrial Biogenesis

Next, we sought to determine if exogenous H₂S could influence cardiac mitochondrial content. For these studies, we administered SG-1002 (20 mg/kg/day) in the chow for 4 weeks. SG-1002 is an H₂S prodrug that we have previously shown to exert cardioprotective effects [11, 16]. Initial studies confirmed that the administration of SG-1002 significantly increased cardiac H₂S levels (Supplemental Figure 1). We then examined if the increase in H₂S influenced cardiac mitochondrial content. Analysis revealed that SG-1002 increased mitochondrial DNA levels, citrate synthase activity, and the number of mitochondria per field of electron microscopy images (Figure 2A–D). Further analysis confirmed that SG-1002 did not influence mitochondrial fusion-fission (Supplemental Figure 2 and Figure 2E–F). Finally, SG-1002 did not alter the state 3 respiration rates of LV fibers in the presence of pyruvate (Figure 2G). It did however, improve the state 3 respiration rates and ATP/oxygen ratios of LV fibers in the presence of palmitoyl-carnitine (Figure 2H–I).

3.3. H₂S Levels Influence PGC-1α

Given that PGC-1α is a master regulator of mitochondrial biogenesis [4, 5], we sought to determine if H₂S influenced the expression or activation of PGC-1α. Initial analysis revealed that the whole cell expression of PGC-1α was not altered in the hearts of CSE Tg⁺ or CSE KO mice (Figure 3A–B). However, the cytosolic levels of PGC-1α were significantly lower in the hearts of CSE Tg⁺ mice (Figure 3A–B). Correspondingly, the nuclear levels of PGC-1α were significantly higher in the hearts of CSE Tg⁺ mice. In contrast, the cytosolic levels of PGC-1α were unaltered in the hearts of CSE KO mice.
However, the nuclear levels of PGC-1α were significantly lower in the hearts of CSE KO mice (Figure 3A–B). In agreement with these changes, the hearts of CSE Tg+ mice displayed elevated expression of a number of PGC-1α target genes, whereas the expression of the same genes were lower in the hearts of CSE KO mice (Figure 3D). Similarly, SG-1002 did not alter the whole cell expression of PGC-1α, but it did decrease the cytosolic levels of PGC-1α, increase the nuclear expression of PGC-1α, and increase the expression of PGC-1α target genes (Figure 3D–F). Together, this data suggests that cardiac H2S levels influence the nuclear localization and transcriptional activity of PGC-1α.

3.4. H2S Does Not Induce Mitochondrial Biogenesis via eNOS/NO

We next turned our attention to the mechanism(s) by which H2S induces PGC-1α-signaling. Specifically, we focused on three proteins known to mediate mitochondrial biogenesis through PGC-1α; eNOS/NO [28, 29]; AMPK; and Sirtuin 1 (Sirt1) [5]. Given that the cardioprotective actions of H2S have been reported to be mediated, in part, via eNOS [18], we first asked if H2S-induced mitochondrial biogenesis was channeled through eNOS/NO. SG-1002 increased the phosphorylation of eNOSSer1177 (activation site) and increased the cardiac levels of nitrite and nitrosothiols (RXNO) (Supplemental Figure 3). To determine if the elevated levels of NO contributed to H2S-induced mitochondrial biogenesis, wild-type (WT) and eNOS KO mice were administered SG-1002 for 4 weeks. SG-1002 increased mitochondrial DNA levels and citrate synthase activity in both strains of mice (Supplemental Figure 3), suggesting that eNOS/NO was not responsible for H2S-induced mitochondrial biogenesis.

3.5. H2S Induces Mitochondrial Biogenesis via AMPK

The transcriptional activity of PGC-1α is regulated by post-translational modifications. For instance, AMPK regulates the transcriptional activity of PGC-1α via phosphorylation [30], whereas Sirt1 regulates its transcriptional activity via deacetylation [5]. Evidence suggests that AMPK both activates Sirt1 and cooperates with it in enhancing the ability of PGC-1α to stimulate mitochondrial biogenesis [31]. So, the next series of experiments sought to determine if H2S-induced mitochondrial biogenesis was channeled through AMPK and Sirt1. SG-1002 increased the phosphorylation and activity of AMPK (Figure 4A–C). No changes were observed in the expression of Sirt1, but SG-1002 did increase its activity (Figure 4D–F). In agreement with these changes, SG-1002 increased the serine phosphorylation of PGC-1α and decreased its acetylation. To determine if AMPK was directly responsible for H2S induced PGC-1α-signaling and mitochondrial biogenesis, αMHC-Cre+ x AMPKfl/fl mice were given SG-1002 for 4 weeks. The deficiency of AMPK did not affect the expression of PGC-1α or Sirt1 (Figure 5A–C). However, SG-1002 failed to increase the serine phosphorylation of PGC-1α, failed to increase Sirt1 activity, and failed to decrease the acetylation of PGC-1α (Figure 5D–H). Importantly, SG-1002 was unable to increase the expression of PGC-1α target genes and mitochondrial DNA levels in the hearts of αMHC-Cre+ x AMPKfl/fl mice (Figure 5I). Together this data suggests that H2S induced mitochondrial biogenesis via an AMPK-Sirt1-PGC-1α signaling cascade.
3.6. H₂S Regulates AMPK by Inhibiting PP2A

We next addressed how H₂S regulates AMPK. The phosphorylation of AMPK is regulated through a balance between the actions of upstream kinases and phosphatases [32]. In the heart, the liver kinase B1 (LKB1) is the major upstream AMPK kinase. SG-1002 did not alter the phosphorylation (activation) of LKB1 (Figure 6A–B). Alterations in the phosphorylation of LKB1 are not always necessary for the induction of AMPK phosphorylation, as changes in AMP and/or ADP levels can influence the confirmation of AMPK, thus allowing for LKB1 to have better access to threonine 172 [32]. The levels of ATP, ADP, and AMP were not altered in hearts of mice administered SG-1002 (Figure 6C–D). Together this data indicates that SG-1002 does not influence the phosphorylation of AMPK via the action of LKB1. We then turned our attention to protein phosphatase 2A (PP2A). PP2A is a serine/threonine protein phosphatase consisting of scaffolding (PP2Aa), regulatory (PP2Ab), and catalytic (PP2Ac) subunits [33]. Analysis revealed that SG-1002 did not alter the protein expression of any of the PP2A subunits (Figure 6E–F). H₂S modifies cysteine residues in proteins through the formation of a persulfide (−SSH) bond by a process termed sulfhydration or persulfidation [34]. Although several proteins have been found to be sulfhydrated, a key finding in recent years was the discovery that the sulfhydration of protein tyrosine phosphatase 1B (PTP1B) inhibits its activity [35]. We, therefore, asked if PP2A was sulfhydrated and if so could this affect its activity. Initial studies revealed that SG-1002 increased the sulfhydralization of cardiac proteins (Supplemental Figure 4). Subsequent analysis using both the modified biotin switch assay and the maleimide assay revealed that SG-1002 increased the sulfhydralization of PP2Aa, PP2Ab, and PP2Ac (Figure 6G–H and Supplemental Figure 4C–H). This was associated with a decrease in the activity of PP2A (Figure 6I), suggesting that H₂S regulates the phosphorylation of AMPK via the inhibition of PP2A.

Subsequent experiments sought to evaluate the temporal nature of this regulation. Beginning 10 minutes after exposing H9c2 cells to Na₂S (100 μM) the activity of PP2A was reduced when compared to untreated cells (Supplemental Figure 5). This decrease continued for 60 minutes with a gradual return to untreated levels. Concurrently, the phosphorylation levels of AMPK followed the opposite trend. Further analysis, revealed that exposing cells to Na₂S for 3 consecutive days resulted in a decrease in the activity of PP2A accompanied by an increase in AMPK phosphorylation. Together these studies indicate that the regulation of this pathway by H₂S can be acute and also maintained chronically.

3.7. H₂S Levels and Mitochondria Content are Reduced in Response to Heart Failure

There is evidence for impaired mitochondrial biogenesis from animal models of heart failure and from studies utilizing human heart failure samples [7, 36]. Additionally, studies report lower circulating H₂S levels in human heart failure patients [37]. However, an association between cardiac H₂S levels and cardiac mitochondrial content has not been explored. Here, we were able to obtain left ventricular samples from end-stage heart failure patients at the time of transplant. Analysis revealed diminished H₂S levels (Figure 7A–B), as well as lower levels of PGC-1α target genes (Figure 7C) and mitochondrial DNA levels (Figure 7D).

Similar results in regards to H₂S levels and mitochondrial content were also found in samples from mice collected at 2 weeks following myocardial ischemia and reperfusion.
We, therefore, sought to determine if restoring H$_2$S levels could influence the mitochondrial content of the ischemic heart via the proposed PP2A-AMPK signaling cascade. Previously, we found that H$_2$S therapy increased mitochondrial content and function in hearts subjected to myocardial ischemia [19, 20]. However, in these studies H$_2$S was administered at the time of reperfusion resulting in a reduction in infarct size. As such, the observed improvements in mitochondria content and function could have been due to the initial reduction in injury. We, therefore, decided to delay the timing of treatment here. Mice subjected to myocardial ischemia and reperfusion were administered normal diet or diet supplemented with SG-1002 beginning 24 hours after reperfusion. The mice were then followed for 2 weeks. Our analysis revealed that restoring H$_2$S levels via the dietary supplementation of SG-1002 increased the sulphydrylation of cardiac proteins and increased the sulphydrylation of PP2Ab and PP2Ac (Figure 8A–B and Supplemental Figure 6A–B). This was associated with a decrease in the activity of PP2A, an increase in the phosphorylation of AMPK, an increase in the gene expression of PGC-1α target genes, an increase in cardiac mitochondrial content, improved mitochondrial respiration, improved ATP production efficiency, and improved cardiac dilatation and function (Figure 8C–H and Supplemental Figure 6C–H).

### 4. Discussion

The transcriptional activity of PGC-1α is regulated by a number of stimuli, exemplifying the range of settings in which mitochondrial biogenesis is induced. The mechanisms governing this regulation have been extensively studied and have been determined to vary between tissues and different settings [5]. AMPK, a member of the metabolite-sensing protein kinase family, is activated in response to alterations in cellular energy levels [32]. Activation of AMPK acts to maintain cellular energy stores, switching on catabolic pathways that produce ATP, while switching off anabolic pathways that consume ATP [38]. AMPK also induces mitochondrial biogenesis via the activation of PGC-1α. Specifically, AMPK regulates PGC-1α directly through serine phosphorylation, as well as indirectly via the activation of Sirt1, which leads to the deacetylation of PGC-1α [5]. This coordinated signaling cascade, whereby the phosphorylation of PGC-1α by AMPK is required for the subsequent Sirt1-mediated deacetylation, demonstrates the complex and specific nature by which PGC-1α is regulated by AMPK [29]. Here, we found that increasing cardiac H$_2$S levels through the dietary supplementation of SG-1002 led to the activation of AMPK and Sirt1 resulting in the phosphorylation and deacetylation of PGC-1α. More importantly, we found that SG-1002 failed to alter PGC-1α, induce PGC-1α target genes, and induce mitochondrial biogenesis in the hearts of AMPKα2 deficient mice – indicating that H$_2$S induces mitochondrial biogenesis via AMPK. Our data also provides some insights into how H$_2$S regulates AMPK. The activation of AMPK is mediated through different mechanisms involving the allosteric regulation of AMPK subunits through changes in adenosine metabolites, activation by LKB1, and changes in the activity of PP2A [32]. Here, we found that H$_2$S did not change the levels of adenosine metabolites nor alter the phosphorylation of LKB1. Rather, our evidence indicates that H$_2$S affects AMPK by decreasing the activity of the serine/threonine protein phosphatase, PP2A.
Recently, H\textsubscript{2}S was shown to inhibit the activity of PTP1B via sulfhydration [35].
Sulfhydration or persulfidation is a post-translational modification by which H\textsubscript{2}S alters
cysteine residues in proteins through the formation of a persulfide (-SSH) bond [34]. Studies
indicate that protein sulfhydration is a major event by which H\textsubscript{2}S elicits cellular signaling.
For instance, the antiapoptotic actions of NF\kappa B are dependent on H\textsubscript{2}S sulfhydrating its p65
subunit [23]. Also, H\textsubscript{2}S induces endothelial cell and smooth muscle cell hyperpolarization
and vasorelaxation by sulfhydrating ATP-sensitive potassium channels [39]. Here, we found
that all three PP2A subunits displayed enhanced sulfhydration in the hearts of mice
supplemented with SG-1002. Since this modification was accompanied by a decrease in
activity, it can be suggested that like for PTP1B, sulfhydration of PP2A imparts inactivation.
Further studies are warranted to elucidate the specific cysteine residues that are modified on
each subunit and to determine how each affects the activity of PP2A.

As noted, H\textsubscript{2}S is a known regulator of cellular bioenergetics via its actions on mitochondrial
function. For instance, H\textsubscript{2}S acts as a stimulator of mitochondrial bioenergetics through its
ability to donate electrons to the mitochondrial electron transport chain [12, 40]. This action
serves a physiological role in the maintenance of mitochondrial electron transport, as well as
complementing and balancing the bioenergetic role of Krebs cycle-derived electron donors
[12]. In contrast, H\textsubscript{2}S is a potent and reversible inhibitor of mitochondrial function via its
regulation of cytochrome c oxidase (complex IV of the mitochondrial electron transport
chain) [41]. Paradoxically, this action contributes to the cardioprotective effects of
exogenous H\textsubscript{2}S, as inhibition of mitochondrial respiration during the early stages of
reperfusion injury limits the generation of reactive oxygen species, which ultimately
preserves mitochondrial function [18–20]. In addition, H\textsubscript{2}S targets several cellular pathways
that influence mitochondrial function [42]. PGC-1\textalpha , not only regulates mitochondrial
biogenesis, but also regulates energy expenditure in the heart. Specifically, PGC-1\alpha is
essential for the maintenance of maximal, efficient cardiac mitochondrial fatty acid
oxidation and ATP synthesis [26]. Consistent with our findings regarding the induction of
PGC-1\alpha signaling, we observed that augmenting cardiac H\textsubscript{2}S levels increased the maximal
capacity for mitochondrial fatty acid \textbeta-oxidation and ATP synthesis, whereas lower H\textsubscript{2}S
levels had the opposite effect.

PGC-1\textalpha-mediated mitochondrial biogenesis in brown fat and endothelial cells is in part
regulated by eNOS and NO [28, 29]. Recent studies indicate that H\textsubscript{2}S not only augments NO
bioavailability and signaling [11, 17], but provides protection against cardiac injury in an
eNOS-dependent manner [15, 18]. Based on this evidence, we speculated that H\textsubscript{2}S induced
mitochondrial biogenesis by activating eNOS and increasing NO levels. Indeed, we found
that H\textsubscript{2}S therapy increased the phosphorylation of eNOS (activation site) and increased
cardiac NO levels. However, H\textsubscript{2}S therapy did not induce mitochondrial biogenesis in an
eNOS-dependent manner, as evidenced by a significant increase in mitochondrial content in
the hearts of eNOS KO mice given SG-1002. This suggests that although eNOS/NO is
important for H\textsubscript{2}S signaling in certain situations, it is not necessary for the induction of
mitochondrial biogenesis in the naïve heart. With that being said, given the complexity by
which PGC-1\textalpha signaling is regulated, we cannot rule out the possibility that under certain
conditions or in different tissues eNOS/NO contributes to H\textsubscript{2}S-mediated mitochondrial
biogenesis in some way. For instance, in the setting of ischemic injury, the induction of
eNOS/NO could contribute to a pro-survival environment, which could indirectly aid in the promotion of mitochondrial biogenesis. Additionally, NO also modifies cysteine residues in a manner similar to sulfhydration; a process termed nitrosylation. While similar in some aspects to sulfhydration, nitrosylation appears to diminish cysteine reactivity, whereas sulfhydration seems to enhance it [43]. There also is some overlap in the proteins targeted by NO and H$_2$S. However, it is not clear if NO and H$_2$S target the same cysteine residues in these proteins or if they target distinct cysteine residues. Further it is not clear if they work together or oppose each other to regulate the function of proteins. Finally, it is not clear if there is a balance of sulfhydration and nitrosylation needed for proper function. Therefore, we cannot rule out the possibility that under certain conditions both NO and H$_2$S target PP2A at the same or different cysteine residues to regulate its activity. Further, under certain conditions it is also possible that H$_2$S induces the nitrosylation of PP2A via its actions on eNOS. Therefore, future studies aimed at identifying the specific residues in PP2A modified by H$_2$S and NO are needed to fully understand if/how these signaling molecules work together to regulate of this protein.

The AMPK-PGC-1α signaling cascade has been extensively studied in the context of cardiac metabolism [32]. So, the novel aspects of this study rest not in the determination that AMPK-PGC-1α signaling leads to mitochondrial biogenesis, but in the evidence that endogenous and exogenous H$_2$S influence this pathway. More so, the finding that a decrease in endogenous H$_2$S levels led to an impairment in AMPK-PGC-1α signaling becomes important when considering the evidence that H$_2$S levels are decreased in pathological conditions – i.e. heart failure (Figure 7 and [37]) – that also present with reduced mitochondrial content [7, 36]. Based on this evidence it can be suggested that endogenous H$_2$S levels not only play an important role in maintaining the mitochondrial content of the heart, but that a reduction in endogenous H$_2$S levels contributes to the pathophysiology of heart failure through a disruption in mitochondrial biogenesis. This idea is supported by our findings that restoring H$_2$S levels with SG-1002 increased mitochondrial content, improved ATP production, and attenuated LV dysfunction in a murine model of ischemia-reperfusion injury. Our study has, therefore, identified an important regulatory mechanism in the mitochondrial biogenesis pathway that if corrected by restoring H$_2$S levels could protect the failing heart and ultimately reduce mortality and morbidity associated with heart failure. Future studies aimed at determining if H$_2$S therapy coupled with other pharmacological agents that target the AMPK-PGC-1α signaling pathway are warranted to fully test this postulate.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

Acknowledgments

This work was funded by grants from the American Heart Association (15POST25610016 and 16GRNT31190016 to J.W.C) and the National Institutes of Health (NIH) (1R01DK115213-01 and 5R01HL098481-05 to J.W.C. and 1R01HL.092141, 1R01HL093579, 1U24HL094373, and 1P20HL113452 to D.J.L.). This work was also supported by funding from the Carlyle Fraser Heart Center of Emory University Hospital Midtown to J.W.C. and an operation grant from the Canadian Institutes of Health Research to R.W.
7. Literature Cited


**Highlights**

- Hydrogen Sulfide levels influence cardiac mitochondrial content
- Hydrogen Sulfide induces mitochondrial biogenesis in an AMPK-dependent manner
- Hydrogen Sulfide activates AMPK via the sulfhydration and inhibition of PP2A
Figure 1.
(A) Ratio of mitochondrial DNA to nuclear DNA. (B) Citrate synthase activity. (C) Representative electron microscopy images of mitochondria. Scale bar equals 3 μm. (D) Number of mitochondria per field of view. (E) Summary of mitochondria area and perimeter measurements (μm²) and (F) percentage of mitochondria in a given field that fell into three size categories based on area: <0.6 μm², 0.6 μm² – 1.0 μm², and >1.0 μm². (G) Basal and maximum (state 3) oxygen consumption rates for permeabilized myocardial fibers in the presence of pyruvate and (H) palmitoyl-L-carnitine. (I) Efficiency of ATP synthesis [ATP produced per oxygen consumed (ATP/O)] in permeabilized myocardial fibers. All samples were collected from hearts of Wild-Type, cystathionase-γ-lyase transgenic (CSE Tg⁺), and CSE deficient (CSE KO) mice. Values are means ±SEM. *p<0.05, **p<0.01, and ***p<0.001 vs. Wild-type.
Figure 2.
(A) Ratio of mitochondrial DNA to nuclear DNA. (B) Citrate synthase activity. (C) Representative electron microscopy images of mitochondria. Scale bar equals 3 μm. (D) Number of mitochondria per field of view. (E) Summary of mitochondria area and perimeter measurements (μm^2) and (F) percentage of mitochondria in a given field that fell into three size categories based on area: <0.6 μm^2, 0.6 μm^2 – 1.0 μm^2, and >1.0 μm^2. (G) Basal and maximum (state 3) oxygen consumption rates for permeabilized myocardial fibers in the presence of pyruvate and (H) palmitoyl-L-carnitine. (I) Efficiency of ATP synthesis [ATP produced per oxygen consumed (ATP/O)] in permeabilized myocardial fibers. All samples were collected from hearts of mice administered standard diet (Chow) or diet supplemented with SG-1002 (SG-1002; 20mg/kg/day) for 4 weeks. Values are means ±SEM. *p<0.05, **p<0.01, and ***p<0.001 vs. Chow.
Figure 3.
(A) Representative immunoblots and (B) analysis of the whole cell, cytosolic and nuclear expression of PGC1α. (C) Relative gene expression of PGC1α target genes associated with mitochondrial biogenesis (tfam, cox4i7, cox1, atp5b). Samples were collected from Wild-Type, CSE Tg+, and CSE KO mice. Values are means ±SEM. *p<0.05 and **p<0.01 vs. Wild-type. (D) Representative immunoblots and (E) analysis of the whole cell, cytosolic and nuclear expression of PGC1α. (F) Relative gene expression of PGC1α target genes associated with mitochondrial biogenesis (tfam, cox4i7, cox1, atp5b). Samples were collected from mice administered Chow or diet supplemented with SG-1002. Values are means ±SEM. *p<0.05 vs. Chow.
Figure 4.
(A–B) Representative immunoblots and analysis of phosphorylated AMPK and total AMPK. (C) AMPK activity. (D–E) Representative immunoblots and analysis of Sirt1. (F) Sirt1 activity. (G) Representative immunoblots and analysis from immunoprecipitation experiments examining the (H) serine phosphorylation and (I) acetylation status of PGC1α. All samples were collected from hearts of mice administered standard diet (Chow) or diet supplemented with SG-1002 (SG-1002; 20mg/kg/day) for 4 weeks. Values are means ±SEM. *p<0.05 and **p<0.01 vs. Chow.
Figure 5.

(A) Representative immunoblots and analysis of (B) PGC1α and (C) Sirt1. (D) Representative immunoblots and analysis from immunoprecipitation experiments examining the serine phosphorylation status of PGC1α. (E) Sirt1 activity. (F) Representative immunoblots and analysis from immunoprecipitation experiments examining the acetylation status of PGC1α. (G) Relative gene expression of tfam, cox417, and mitochondria DNA. All samples were collected from hearts of Wild-type (αMHC-Cre*) and αMHC-Cre* x AMPK floxed (αMHC-Cre* x AMPKfl/fl) mice administered standard diet (Chow) or diet supplemented with SG-1002 (SG; 20mg/kg/day) for 4 weeks. Values are means ±SEM. *p<0.05, **p<0.01, and ***p<0.001 vs. WT Chow.
Figure 6.
(A–B) Representative immunoblots and analysis of phosphorylated LKB1 and total LKB1. (C) Cardiac levels of ATP, ADP, and AMP. (D) Ratio of AMP to ATP. (E–F) Representative immunoblots and analysis of PP2A subunits. (G–H) Representative immunoblots and analysis from the modified biotin switch assay examining the sulfhydration status of PP2A subunits. (I) PP2A activity. All samples were collected from hearts of mice administered standard diet (Chow) or diet supplemented with SG-1002 (SG-1002; 20mg/kg/day) for 4 weeks. Values are means ±SEM. *p<0.05 and ***p<0.001 vs. Chow.
Figure 7.
Cardiac levels of (A) free H$_2$S and (B) sulfane sulfur. (C) Relative gene expression of atp5b, mt-co1, and tfam. (D) Ratio of mitochondrial DNA to nuclear DNA. Left ventricular samples were collected from non-failing and end stage heart failure patients. Number in bars represent sample sizes. Values are means ±SEM. *p<0.05 and ***p<0.001 vs. Non-Failing.
Figure 8.
Cardiac levels of (A) free H$_2$S and (B) sulfane sulfur. (C) Ratio of mitochondrial DNA to nuclear DNA. (D) Basal and maximum (state 3) oxygen consumption rates for permeabilized myocardial fibers in the presence of palmitoyl-L-carnitine. (E) Efficiency of ATP synthesis [ATP produced per oxygen consumed (ATP/O)] in permeabilized myocardial fibers. Samples were collected from hearts of mice subjected to 60 minutes of ischemia and 2 weeks of reperfusion. Values are means ±SEM. *p<0.05, **p<0.01, and ***p<0.001 vs. Sham. (F) Left ventricular end-diastolic diameter (LVEDD), LV end-systolic diameter (LVESD), and LV Ejection Fraction measured in groups of mice using echocardiography images 2 weeks following myocardial ischemia and reperfusion (Post). Values are means ±SEM. ***p<0.001 vs. Baseline.