Mitochondrial quality-control dysregulation in conditional HO-1\(^{-/-}\) mice

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The heme oxygenase-1 (Hmox1; HO-1) pathway was tested for defense of mitochondrial quality control in cardiomyocyte-specific Hmox1 KO mice (HO-1\(^{[CM]}{-/-}\)) exposed to oxidative stress (100% O\(_2\)). After 48 hours of exposure, these mice showed persistent cardiac inflammation and oxidative tissue damage that caused sarcomeric disruption, cardiomyocyte death, left ventricular dysfunction, and cardiomyopathy, while control hearts showed minimal damage. After hyperoxia, HO-1\(^{[CM]}{-/-}\) hearts showed suppression of the Pgc-1\(\alpha\)/nuclear respiratory factor-1 (NRF-1) axis, swelling, low electron density mitochondria by electron microscopy (EM), increased cell death, and extensive collagen deposition. The damage mechanism involves structurally deficient autophagy/mitophagy, impaired LC3II processing, and failure to upregulate Pink1- and Park2-mediated mitophagy. The mitophagy pathway was suppressed through loss of NRF-1 binding to proximal promoter sites on both genes. These results indicate that cardiac Hmox1 induction not only prevents heme toxicity, but also regulates the timing and registration of genetic programs for mitochondrial quality control that limit cell death, pathological remodeling, and cardiac fibrosis.

Introduction

The heme oxygenase-1 (HO-1; Hmox1) enzyme system is among the most important inducible mechanisms for cell protection against oxidative damage in the cardiovascular system. During oxidative stress, HO-1 eliminates cellular free heme (1), a powerful oxidant (2, 3), and produces the physiological gas carbon monoxide (CO) (4), which has potent antiinflammatory and antiapoptotic effects and stimulates mitochondrial biogenesis (5, 6). Moreover, administration of low-level CO gas or CO-releasing molecules mitigates many pathological injuries that affect energy metabolism in cells and tissues of experimental animals (7, 8). Hmox1-deficient heart cells accumulate lethal molecular oxidant damage, and embryonic Hmox1 deletion in mice has a high prenatal mortality, with survivors showing growth retardation, iron overload, organ fibrosis, and inflammatory tissue damage (9). Hmox1-deficient mice are highly susceptible to ischemia/reperfusion (I/R) injury and, after hypoxia, show evidence of right ventricular infarction (10, 11). This enzyme system has thus been a focus of attention in several cardiovascular diseases, including heart failure (HF) (12, 13), which is accompanied by oxidant-mediated hypertrophy, dilation, fibrosis, and metabolic disturbances that reduce contractile function (14, 15). The rare human HO-1 deficiency state and certain polymorphisms of the human HO-1 gene promoter are also associated with the development of cardiovascular disease (16, 17).

Moreover, HO-1 is mechanistically important for cardiomyocyte function through its ability to regulate mitochondrial health because most (95%) of the heart's ATP production is derived from oxidative phosphorylation (OXPHOS) (18, 19). Indeed, even sublethal mitochondrial damage can impair cardiac ATP provision and ventricular contraction. Persistent mitochondrial damage is a major source of oxidants (20) and a major factor in the pathogenesis of many cardiovascular diseases (21, 22). The induction of HO-1 leads to increased cellular CO production, which generates a redox signal for the induction of mitochondrial biogenesis through stimulation of mitochondrial ROS production (5, 23). HO-1 has also been implicated in the induction of macroautophagy (24) and if this capacity includes mitophagy, the enzyme would be involved in the regulation of the entire mitochondrial quality control cycle, which enables accurate and precise mitochondrial turnover (24–26). Disruption of mitophagy is proinflammatory and prooxidant (27, 28), and impaired autophagy compromises bioenergetics and leads to cell death, chronic inflammatory heart disease, and HF progression (29, 30). Accordingly, mitophagy has a putative cardioprotective role (31, 32).
The induction of mitophagy by cell stress involves the PINK1/Parkin pathway (33). PINK1, a serine/threonine kinase, normally localizes to mitochondria, where it is actively degraded by mitochondrial processing peptidase and presenilin-associated rhomboid-like protease (34). Loss of membrane potential blocks PINK1 importation, and its accumulation on the outer mitochondrial membrane allows the recruitment of Parkin (PARK2), an E3 ubiquitin ligase (34). PINK1 first phosphorylates mitofusin 2 (MFN2), which acts as a mitochondrial receptor for PARK2 (35). It then phosphorylates ubiquitin, which activates PARK2's E3 ubiquitin ligase activity (36, 37). Activated PARK2 causes non-classical ubiquitination of several outer mitochondrial membrane proteins, targeting the organelles for lysosomal degradation (38). Adaptor proteins such as p62/SQSTM1 bind ubiquitinated proteins and bind LC3 on the autophagosome (39). PARK2 also acts as a mediator of classical proteasomal degradation of mitochondrial proteins (40).

Physiologically, PARK2-deficient mice show impaired recovery of cardiac contractility after sepsis (41) and impaired mitophagy after myocardial infarction, resulting in poorer survival and larger infarct sizes relative to WT (31). Similarly, PINK1-deficient hearts are more susceptible to I/R injury compared with WT (32), and PINK1 overexpression in cardiac cells inhibits simulated I/R-mediated cell death (32). PINK1 deficiency in mice leads to ventricular dysfunction and hypertrophy (42), and by 2 months of age, these hearts show oxidative stress, reduced respiration, and fibrosis (32, 42). In contrast, PARK2−/− mice have normal mitochondrial and cardiac function (31). This means that PINK1 is essential for clearing damaged mitochondria, and there may also be other functions for PINK1 in myocytes.

Oxidative stress also contributes to adaptation to myocardial ischemia and can be induced by pretreatment with hyperoxia (>95% O2), a potentially valuable means of clinical preconditioning (43). O2 preconditioning attenuates I/R injury in rat hearts by decreasing infarct size and improving postischemic heart function (44, 45). This suggests that oxidative stress may not only induce antioxidant protection, but may also generate a protective mitochondrial status through HO-1 induction and downstream support of mitochondrial quality control (25, 46). However, O2 is also toxic at high concentrations over prolonged periods of time, which sets limits on its therapeutic use.

We generated KO mice containing myocyte-specific inducible HO-1 and exposed them to 100% O2 in order to induce the HO-1 response and to stimulate HO-1–mediated mitochondrial biogenesis and assess its linkage to the molecular regulation of mitophagy. We hypothesized that the absence of HO-1 would block the redox-sensitive transcriptional response for mitochondrial biogenesis during the oxidative stress of 100% O2 exposure, thereby revealing its linkage with mitophagy through PINK1 and PARK2 gene regulation. Such information would inform HO-1/CO–based therapeutics and ways that the induction of mitophagy could be used to treat HF and perhaps other degenerative heart diseases.

**Results**

*Development of HO-1 KO mice and exposure to 100% O2.* Cardiac-specific *Hmox1* (HO-1) KO mice were generated by crossing *Hmox1* fl/fl (HO-1 fl/fl) mice with transgenic mice expressing MerCreMer under the control of the myosin heavy chain (Myh6) promoter to permit tamoxifen-inducible deletion of the *Hmox1* allele in the adult. HO-1 fl/fl/MerCreMer (HO-1 fl/fl/Myh6-Cre) mice were born at the expected Mendelian frequency and were indistinguishable in appearance from control HO-1 fl/fl littermates or WT/Cre. To generate the *Hmox1*–deficient cardiomyocyte strain (HO-1(CM)−/−), 8- to 10-week-old mice were given tamoxifen (i.p.) for 3 days, and 8 days later, the hearts were harvested and examined (Supplemental Figure 1; supplemental material available online with this article; https://doi.org/10.1172/jci.insight.89676DS1).

Eight days after tamoxifen, HO-1(CM)−/− and WT/Cre mice were exposed to 100% O2 for 2 days (48 hours) and then transferred to filtered air cages for 8 days (Supplemental Figure 1). Immediately after hyperoxia, the hearts appeared normal, although microscopically, the hearts of the HO-1(CM)−/− mice showed a few foci of neutrophilic inflammation (Supplemental Figure 2A). Seven days after tamoxifen, HO-1(CM)−/− mice appeared healthy. Forty-eight–hour exposures to 100% O2 are not lethal to WT mice or to HO-1(CM)−/− mice. At day 8 after hyperoxia, 2/46 WT mice had died, and 8/46 of the HO-1(CM)−/− mice had died (17.4%; P = 0.09). Those mice had elevated NLRP3 inflammasome protein levels (Supplemental Figure 2B), failed to upregulate cardiac HO-1 mRNA (Supplemental Figure 2C), and showed little response in the content of mitochondrial TCA cycle enzyme citrate synthase (CS; Supplemental Figure 2D).
The cardiomyocyte respiratory capacity evaluated by maximum ADP respiration rates of freshly isolated left ventricular cells were the same in both strains of mice before 100% O₂ exposure; however, respiration rates for Complex I substrates after hyperoxia were reduced in both strains (Table 1). This may be related in part to low mRNA levels of NADH dehydrogenase Iβ complex subunit 8 (NDUFB8), a nuclear DNA–encoded Complex I protein integral to Complex I assembly (47), and NDUFAF1, a protein necessary for the assembly and stability of Complex I in the HO-1–deficient heart (48) (Supplemental Figure 3).

Even 8 days after O₂, low HO-1 transcript and protein by quantitative PCR (qPCR) and immunoblotting were still apparent in the hearts of HO-1(CM)–/– mice (Figure 1, A and B; note stable HO-1 protein in skeletal muscle). Also at 8 days, cardiac HO-1 expression remained 2.5-fold higher in WT/Cre than in HO-1(CM)–/– mice, but the HO-2 constitutive isoform was stable in both strains (Figure 1, C and D). HO-1(CM)–/– hearts had enlarged by 8 days after O₂ (Figure 1E, left), and ventricle/body weight ratios (3.2 ± 0.27 before to 5.6 ± 0.24 after O₂) had increased significantly compared with WT/Cre hearts (2.9 ± 0.14 before to 3.3 ± 0.17 after O₂; mean ± SEM mg/g body weight).

The increase in ventricular size and eccentric hypertrophy in HO-1(CM)–/– hearts was consistent with dilated cardiomyopathy. H&E-stained cross sections of HO-1(CM)–/– hearts at 8 days after 100% O₂ revealed myofibrillar loss, myocyte degeneration, paleness, swelling, and leukocyte infiltration that were not seen in WT/Cre mice (Figure 1E, right). Echocardiographic data from HO-1(CM)–/– mice revealed a minimal decrease in fractional shortening (%FS) compared with WT/Cre. However, %FS and ejection fraction (%EF) in HO-1(CM)–/– hearts had decreased significantly while the wall thickness changed significantly, suggesting eccentric hypertrophy in the HO-1(CM)–/– group (Table 2). Ventricular function was not impaired in tamoxifen-treated WT/Cre mice; thus, the effect was not due to the presence of Cre recombinase. The ventricular hypertrophy was reflected by increased tissue expression of hormonal hypertrophy markers, atrial natriuretic factor (ANF), and brain natriuretic peptide (BNP) in HO-1(CM)–/– hearts (Figure 1, F–H).

Cardiac oxidative stress, inflammation, and apoptosis. The extent of oxidative tissue damage was assessed by the quantification of total levels of protein carbonylation and malondialdehyde (MDA) adduct formation, and the level of mitochondrial DNA (mtDNA) damage was by mitochondrial 8-hydroxy-2'
-deoxyguanosine (8-OHdG) content (Figure 2, A–C). The data from 8 days after 100% O₂ show that cardiac MDA and protein carbonyl levels were about twice as high in HO-1(CM)–/– mice compared with WT/Cre mice. However, mitochondrial 8-OHdG levels were increased by 4-fold in HO-1(CM)–/– mice compared with WT/Cre mice. These findings indicate the importance of HO-1 induction not only in antioxidant protection, but also in mtDNA integrity during oxidative stress. HO-1(CM)–/– mouse hearts showed inducible NOS (iNOS) and inflammasome output cytokines IL-1β and IL-18 that were twice as high as WT/Cre mice at 8 days after 100% O₂ (Figure 2, D–F). We compared the number of TUNEL-positive cells in the hearts of HO-1(CM)–/– and WT/Cre mice before and after 100% O₂ and found about a 4-fold increase in positive cells in HO-1(CM)–/– mice compared with WT/Cre mice, indicating significant increases in cell death rates after Hmox1 ablation (Figure 2G).
HO-1 and mitochondrial biogenesis. We evaluated mitochondrial biogenesis and its relationship to tissue damage recovery after oxidative stress because HO-1 is a known activator of the program (5, 49). Formalin-fixed heart sections were first stained for CS, a marker of mitochondrial density and distribution. The hearts of HO-1(CM)−/−, but not WT/Cre mice, demonstrated a patchy loss of CS that had not improved by 8 days after hyperoxia (Figure 3A). In contrast, CS levels increased in WT/Cre mice, consistent with an increase in mitochondrial mass by 8 days. This was corroborated by increased transcription factor A, mitochondrial (Tfam) protein levels in WT/Cre but not in HO-1(CM)−/− hearts 8 days after hyperoxia (Figure 3B). Using transmission electron microscopy (TEM) to examine cardiac ultrastructure, we found mitochondrial swelling and evidence of dilated intramembranous spaces at 8 days after tamoxifen administration in HO-1(CM)−/− mice (Figure 3C). Normal mitochondrial architecture was disrupted, and the distribution of sarcolemma appeared clustered instead of regularized (Figure 3C). HO-1−deficient cardiomyocytes showed widespread disorganization of myofibrils and disruption of contractile elements. In some areas, myofibrils were replaced by irregular mitochondria. At 8 days after O2, WT/Cre hearts maintained a structurally normal appearance, with regular arrays of myofibrils within the sarcomeres and normal size and number of mitochondria (Figure 3C).
The mitochondrial dynamics genes Drp1 and Opa1 mRNA increased significantly 8 days hyperoxia in WT/Cre hearts, but not in HO-1(CM)−/− hearts (Figure 3, D and E). The total mtDNA content and mitochondrial DNA transcription assessed by mitochondrial mRNA levels for ND1, as well as nuclear-encoded ND5, were all decreased in HO-1(CM)−/− hearts, suggesting the failure of mitochondrial biogenesis induction in the HO-1−deficient state (Figure 4, A–C). To understand the mechanism, we evaluated the expression levels of two transcriptional regulators of mitochondrial biogenesis, NRF-1 and PGC-1α. WT/Cre hearts showed a 2-fold increase in protein levels for NRF-1 and its PGC-1α partner, whereas NRF-1 and PGC-1α protein levels were markedly decreased in HO-1(CM)−/− hearts (Figure 4, D–F). This implicated HO-1 as a key mediator of the mitochondrial content differences in the two strains of mice after the oxidative stress of hyperoxia.
Autophagy and mitophagy. Surprisingly few mitophagosomes were identified in tamoxifen-treated HO-1(CM)−/− hearts by electron microscopy (Figure 3C). The relative lack of autophagic vacuoles was striking, given the extent of mitochondrial disruption and myocardial damage in HO-1−deficient myocytes. To explore this apparent defect in autophagy/mitophagy, we examined LC3II/LC3I and p62 levels in WT/Cre and HO-1(CM)−/− hearts before and after O2. The conversion of LC3I to LC3II is indicative of autophagosome formation (50), whereas p62 decorating damaged mitochondria is degraded by autophagy and only accumulates when autophagy is reduced or impaired (51). LC3II/LC3I ratios were significantly reduced, and p62 levels were higher in HO-1(CM)−/− hearts after hyperoxia (Figure 5, A and B). Total LC3 levels were unaffected, indicating that loss of HO-1 somehow impairs LC3 processing rather than LC3 expression (Figure 5A). Also, Atg4B level, an autophagy protein that cleaves pro-LC3 to LC3-I, was decreased in HO-1(CM)−/− hearts compared with WT/Cre (Figure 5C). HO-1(CM)−/− hearts also exhibited lower levels of Beclin-1 protein, which is essential for autophagosomal membrane formation (Figure 5D). These data collectively imply that mitophagy in the HO-1(CM)−/− hearts has been interrupted after hyperoxia.

Figure 4. HO-1 and regulation of cardiac mitochondrial biogenesis. (A) Deletion of HO-1 decreases cardiac mRNA expression of mitochondrial-encoded gene ND1. (B) Deletion of HO-1 decreases cardiac mRNA expression of nuclear-encoded gene ND5. (C) Mitochondrial mtDNA copy number by qPCR shows significant decline in mtDNA copies after hyperoxia in HO-1−deficient hearts. (D) Representative Western blots for nuclear-encoded NRF-1 and PGC-1α proteins that regulate the mitochondrial genome. (E and F) The quantification of NRF-1 and PGC-1α is, by densitometry, relative to loading control Lamin B. HO-1 ablation led to decreases in NRF-1 transcription factor and PGC-1α coactivator protein expression in the heart (mean ± SEM; horizontal bars represent mean values.*P < 0.05 for pre- vs. posthyperoxia; n = 6/group; 2-way ANOVA).
Mitophagy is mediated by the PINK1-PARK2 signaling pathway or by mitophagy receptor Bnip3l (Nix), and depolarized mitochondria are recognized by PINK1 and labeled through PARK2-mediated protein ubiquitination, which subsequently activates autophagy-related genes (ATGs) to engage autophagy (35).

We compared activation levels of myocardial PINK1/PARK2 before and after hyperoxia and found that HO-1 deficiency drastically blocked the cellular enrichment of mitochondrial PINK1 and PARK2 after hyperoxia (Figure 6A), suggesting defective PINK1/PARK2 signaling. We realized that PARK2 total protein levels tracked positively with HO-1 levels (Figure 1, C and D), which suggested that HO-1 activity sets the transcriptional regulation of the Park2 gene. Accordingly, HO-1–dependent differences in Park2 and PINK1 mRNA levels were detected in the hearts (Figure 6, B and C). To determine whether Park2 and PINK1 are direct transcriptional targets of the NRF-1 axis downstream of HO-1, we used comprehensive in silico analysis of transcription factors binding to the promoter regions of mouse Park2 and PINK1 (5). The analysis of transcription factor binding sites in mouse Park2 and Pink1 (–500 to +200 Bp) promoter regions showed multiple NRF-1 transcription factor binding sites with nearby Sp1 sites within the immediate promoter sites of both genes. The Sp1 transcription factor activates or represses transcription in response to a range of pathological stimuli.

To assess transcriptional regulation of the Park2 and Pink1 promoter motifs in vivo, ChIP analysis was performed on nuclear DNA recovered from the hearts under control and 8 days after hyperoxic conditions using primers for highly proximal locations, as shown in Figure 6D. NRF-1 occupancy of Park2 and Pink1 promoters in vivo was identified (Figure 6E). Hyperoxia increased NRF-1 recruitment to Park2 and Pink1 promoters in WT/Cre heart (Figure 6E). However, NRF-1 was not recruited to the Park2 or Pink1 sites in HO-1(CM)−/− mice. WT/Cre NRF-1 recruitment corresponded to increased HO-1 and NRF-1 expression after hyperoxia. This demonstrated that HO-1 mediates mitophagy through transcriptional regulation of Park2 and Pink1 by NRF-1. An impairment of mitophagy in HO-1–deficient hearts after oxidative stress may not only jeopardize ATP production, but also induce myocardial inflammation, as revealed by a persistent inflammatory signature (Figure 2, D–F). Thus, the data support that HO-1 ablation impairs mitophagy, leading to accumulation of damaged mitochondria, oxidized macromolecules, and myocardial inflammation.
We used cultured beating HL-1 cells (5) to demonstrate that NRF-1 overexpression increased NRF-1 protein and downstream mitophagy genes Park2 and Pink1. In contrast, NRF-1 silencing significantly reduced Park2 and Pink1 gene expression (Supplemental Figure 4). These findings imply that HO-1–dependent induction of NRF-1 enhances mitochondrial quality control (QC) pathway function, allowing the beating rate of the cells to improve, while the silencing of HO-1 and NRF-1 decreases QC pathway function and reduces the rate of beating and the survival of these cells. Furthermore, HO-1 overexpression will increase the accumulation of the reduced form of MTT (3-[4,5-dimethylthiazol-2-yl]-2,5-diphenyl tetrazolium bromide) in mitochondria, which reflects overall oxidative metabolic activity of the cells (Supplemental Figure 5). MTT is used as a surrogate for total respiration under standardized conditions and does not define the behavior of individual respiratory complexes. Meanwhile HO-1 silencing decreased the formazan accumulation in HL-1 cells, implying that HO-1 overexpression leads to augmented respiratory activity, while HO-1 silencing leads to a decline in respiration. Moreover, 72 hours after HO-1 transfection, the transduced HL-1 cells had increased their beating frequency, while HO-1 silencing significantly reduced beating rate (Supplemental Figure 5).

Cardiac fibrosis. Studies with Masson’s trichrome–stained heart sections demonstrated extensive areas of interstitial fibrosis in the hearts 8 days after O2 in HO-1(CM)−/− relative to WT/Ce littermates (Figure 7, A)

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**Figure 6.** HO-1 is required for Pink1 and Park2 gene expression through NRF-1 activation after hyperoxia. (A) Representative Western blots of Park2 (Park2) and PINK1 in mitochondrial fractions of the heart from WT/Ce and HO-1(CM)−/− mice prepared before and after hyperoxia. Porin was used as a loading control. (B) Deletion of HO-1 in HO-1(CM)−/− mice decreases cardiac expression of Park2 mRNA. (C) Lack of HO-1 in the heart of HO-1(CM)−/− mice decreased expression of PINK1 mRNA. (D) Schematic diagrams (~500 to +200 bp) of the Pink1 (ENSMUST00000030536) and Park2 (ENSMUST00000191124) promoter regions. Sequences were aligned between human and mouse using rVISTA 2.0. A search for NRF-1 binding sites upstream of the transcription start site (TSS) identified multiple consensus motifs for NRF-1 in human and mouse Pink1 and Park2 gene promoters by Genomatix and DNAis software. (E) NRF-1 occupancy of promoters in vivo was investigated by ChIP analysis. Input lanes show the PCR product derived from chromatin prior to immunoprecipitation. Antibodies against NRF-1 used for immunoprecipitation, while IgG was used as negative control. Precipitated DNA was analyzed by PCR with primer sets specific for the 3 promoters (mean ± SEM; horizontal bars represent mean values. *P < 0.05 for pre- vs. posthyperoxia; n = 6 per group; 2-way ANOVA).
Fibrotic gene activation was evaluated by qPCR and showed greatly increased collagen mRNA levels in HO-1(CM)−/− hearts (Figure 7C) that was not present in WT/Cre mouse hearts. By immunoblotting, it was apparent that TGF-β1 levels were increased in the HO-1(CM)−/− mice, while levels of the antifibrotic protein CCN5 were decreased relative to WT/Cre littermates (Figure 7D). Thus, the inability to upregulate HO-1 resulted in extensive oxidative tissue damage and inflammation that led to significant fibrosis for many days after an O2 exposure that was normally be well-tolerated by the heart.

Discussion
Each of the products of heme catabolism provides a different aspect of cell protection to avert serious oxidative tissue damage (52). Thus, the HO-1 system is far more than an antioxidant enzyme; for example, physiological CO production activates an essential mitochondrial defense by signaling the transcription of nuclear genes for mitochondrial biogenesis (49). But because free heme is a potent oxidant, it has been difficult to distinguish the cytoprotective effects of heme elimination by HO-1 from its mitochondrial role. (4, 5, 49). Here we demonstrate that HO-1(CM)−/− mice exposed to 100% O2 develop an evolving postexposure oxidative cardiac injury and HF due to the failure of mitochondrial quality control. In doing so, we confirm the induction of mitochondrial biogenesis by HO-1 but, more
importantly, uncover the HO-1 system as an activator of transcriptional genes for selective mitophagy of cardiac mitochondria through nuclear respiratory factor–dependent (NRF-1–dependent) expression of the PINK1/PARK2 mitophagy gene pathway.

Normally, 48 hours of O₂ exposure would produce mild self-limited cardiac injury and minimal mortality in mice, and the hearts would recover over a week or two. However, HO-1(CM)−/− mice showed a severely maladaptive cardiac response to this moderate oxidative stress compared with WT/Cre littermates. Although we had anticipated acute cellular damage due to loss of HO-1’s heme-scavenging function, these hearts displayed persistent inflammation and mitochondrial damage, resulting in cell death and rapid development of ventricular dilation and fibrosis (53, 54). A striking finding was impairment of autophagy/mitophagy, both under basal conditions and impressively in response to the oxidative stress. These results collectively support HO-1 as a critical regulator of mitochondrial quality control in the heart following oxidative tissue damage.

Prolonged O₂ breathing causes minor hemodynamic changes, such as bradycardia, and slight decreases in stroke volume and cardiac output (55). After O₂ exposure, we observed nothing outside of the physiological range in terms of heart rate and cardiac output in HO-1(CM)−/− mice. However, within a few days after hyperoxia, there were reductions in cardiac %EF and %FS in HO-1(CM)−/− mice (Table 2) compared with WT/Cre mice. Inspection of these hearts disclosed an increase in ventricular size, with dilation of both ventricles suggesting functional decoupling after hyperoxia. The supporting molecular evidence indicated that the heart is undergoing hypertrophy and remodeling related to the loss of the enzyme and damage to mitochondrial quality control, although secondary molecular responses related to the progressive decompensation cannot be excluded.

The induction of HO-1 by O₂ is well known to be found in WT/Cre mice (56) but was not present in HO-1(CM)−/− mice. This is consistent with the upregulation of WT HO-1 by CO in the amelioration of cardiac cell death and fibrosis after doxorubicin, as well as with HO-1’s inhibitory effect on inflammatory and fibrotic responses in the injured myocardium in mice (6, 25). The requirement for HO-1 in the activation of mitochondrial quality control also means that CO administration or other HO-1 products alone could not provide complete protection, thereby excluding several protective possibilities attributed to the products of heme breakdown (7, 8).

HO-1 also has a critical protective role in the development of inflammation in the cardiovascular system (25, 57), and our results confirm previous studies demonstrating that the expression of certain inflammatory genes is increased by HO-1 inhibition (58). In addition, persistent inflammation in the O₂-exposed HO-1(CM)−/− mouse heart is reflected by the increased levels of inflammatory mediators IL-1β, IL-18, and NOS2. These data confirm that the antiinflammatory effects of HO-1 are part of its known cytoprotective role.

In the failing heart, HO-1 induction allows upregulation of an important adaptation that opposes pathological remodeling and is mediated in part by product-dependent inhibition of apoptosis (59). Moreover, apoptosis from severe oxidative stress and inflammation intersect in cardiac injury (60, 61). Here, the numbers of TUNEL-positive cells increased in HO-1(CM)−/− mice compared with WT/Cre mice, supporting apoptosis as an important mechanism of cardiomyocyte death, although in some areas of the injured HO-1(CM)−/− hearts, cardiomyocyte death appeared to be due to necrosis. HO-1 overexpression prevents oxidative damage in cells (53, 54); hence, its absence would allow oxidant damage to the products of protein carbonylation, lipid peroxidation, and mtDNA oxidation to accumulate after hyperoxia and contribute to cardiomyocyte death.

The cell’s need for continuous OXPHOS makes mitochondria both generators and targets of ROS (62); damaged mitochondria elaborate excessive superoxide anion and H₂O₂ that can cause oxidative cell damage unless those organelles are removed from the cell (63, 64). Mitochondrial dysfunction and impaired mitochondrial biogenesis has repeatedly been observed in the failing heart in animal models and in patients, but the molecular mechanisms are poorly understood (65). Among the pathways for HO-1–induced improvement in cardiac function is transcriptional signaling for the maintenance of mitochondria (5, 49). That mitochondrial biogenesis is impaired in the HO-1(CM)−/− heart was anticipated and supported by immunochemistry, mtDNA depletion, and significant loss of mtDNA-encoded genes. WT/Cre mice avoided this mitochondrial destruction by significantly increased transcriptional levels of genes required for mitochondrial function (CS, ND1, Opal1, and Tfam) compared with HO-1(CM)−/− mice, which were not protected. Accordingly, the expression levels of mtDNA regulatory proteins (NRF-1, PGC-1α) were significantly upregulated in WT/Cre but not in HO-1(CM)−/− mice.
Oxidative mitochondrial damage in cardiac I/R injury has been implicated as a cause of cell death (66), but the prevention of heme accumulation and nonfunctional protein aggregates that lead to a cycle of oxidative stress and deterioration of mitochondrial function is clearly protective (67). There is further evidence that energy depletion together with ROS and protein aggregation induces autophagy (68, 69). One study showed that activation of autophagy clearly reduced myocardial infarct size in a mouse model of I/R injury (70), and the regulation of autophagy via HO-1 may play a protective role in other tissues (71). In the heart, autophagy is essential for homeostasis both under basal conditions and in response to stress (70, 72).

For example, autophagy is important for protecting cardiomyocytes from hyperglycemic damage (73), and suppression of autophagy contributes to the development of cardiomyopathy in diabetes (74).

Healthy mitophagy is designed to clear damaged mitochondria effectively before they cause cellular harm (31). Damaged mitochondria release factors into the intermembrane space that signal not only macroautophagy, but also selective degradation of individual mitochondria (75, 76). Impaired mitophagy in the heart leads to accumulation of aberrant mitochondria and, subsequently, to contractile dysfunction and worsening of overall cardiac function (72, 77). Here, the lack of HO-1 decreased the number of autophagosomes; decreased LC3 II, Atg4B, and Beclin1 expression; and increased levels of p62 in the HO-1(CM)−/− hearts, consistent with impairment of autophagy/mitophagy after hyperoxia. We observed by electron microscopy (EM) that defective mitochondria after hyperoxia were engulfed by autophagosomes in WT/Cre mice but not in HO-1(CM)−/− mice. In WT/Cre mice, mitophagy is associated with upregulation of Drp1 and PARK2 protein levels. In HO-1(CM)−/− mice, however, 100% O2 significantly decreases Opa1, Drp1, and PARK2 and Pink1 protein levels. In HO-1(CM)−/− mice, however, 100% O2 significantly decreases Opa1, reflecting cristae destruction and a lack of autophagosomes accompanied by depressed Complex I respiration. This likely reflects low NDUF8 and NDUFA1 assembly protein expression, low mtDNA levels, and increased oxidant production (78). The implication is that HO-1 is necessary for the early activation of mitophagy that prevents the evolution to persistent cardiac injury following hyperoxia.

PARK2 is also recruited to damaged mitochondrial outer membranes in order to allow signaling for mitochondrial degradation (33, 79). Thus, mitochondrial dysfunction during HO-1 deficiency leading to the inability to clear damaged mitochondria by mitophagy would be detrimental to cardiomyocyte survival. Since PARK2 and Pink1 operate together, and the Pink1/PARK2 pathway is altered in HO-1 deficiency, we searched for Pink1/PARK2 regulation by transcription factors downstream of HO-1 by comprehensive in silico analysis. We uncovered the probable regulation of Pink1/PARK2 by NRF-1 by the presence of tandem Sp1 sites. This was tested experimentally in vivo by occupation of NRF-1 at consensus sites in the PINK1/PARK2 proximal gene promoters in WT/Cre mice and HO-1(CM)−/− mice. NRF-1 occupancy of PINK1/PARK2 promoter sites was significantly enhanced after hyperoxia in WT/Cre but was strikingly absent on NRF-1 binding in HO-1(CM)−/− mice. Moreover, NRF-1 silencing or transfection in beating HL-1 cardiomyocytes led to low vs. high gene expression, respectively, for Pink1 and Park2. These results are consistent with data on mitochondrial biogenesis, demonstrating the HO-1-dependent nuclear accumulation of NRF-1 (80). Thus, HO-1-dependent NRF-1 induction permits the genetic upregulation of both mitochondrial biogenesis and mitophagy in the heart.

The remodeling of ventricular dilation involves individual cardiomyocytes, which undergo eccentric hypertrophy, together with interstitial cell proliferation by fibroblasts, which adversely affects cardiac structure and function, ultimately leading to HF (81). A key effector of pathological remodeling operates through connective tissue growth factor regulated by the TGF-β1 pathway, which promotes fibroblast proliferation and extracellular matrix production in connective tissues (82, 83). We found increased fibrosis in HO-1(CM)−/− mice, accompanied by increases in TGF-β1 expression, which stimulates the accumulation of extracellular matrix proteins and fibrosis (84, 85). In WT/Cre mice, TGF-β1 was not significantly induced after hyperoxia, and there was a significant increase in CCN5, a negative regulator of α-SMA and collagen I expression and a potent repressor of TGF-β1 (86, 87). CCN5 expression, however, was reduced in HO-1(CM)−/− mice. It is intriguing to speculate that HO-1 upregulation opposes the myocardial fibrotic response through this pathway.

In summary, we report a new mechanism for HO-1 protection of the heart from ongoing oxidative injury through the maintenance of mitochondrial homeostasis by the transcriptional regulation of key mitophagy proteins. This mechanism extends far beyond heme clearance by encompassing the well-known role of HO-1 in mitochondrial biogenesis (49), as well as the facilitation of mitophagosome clearance of damaged mitochondria leading to mitigation of cellular oxidant production. In this context, HO-1–induced cardioprotection is driven by sentinel transcriptional events involving activation of...
redox-sensitive transcription factors, such as NRF-1, that prevent the arrest of stress-induced mitophagy from consumption of tagged organelles labeled with essential decorator proteins such as Pink1 and PARK2. In summary, HO-1–dependent induction of NRF-1 is a critical regulator of the cardiac transcriptional circuitry for the overall mitochondrial quality control cycle that arrests late cell death and appears to prevent fibrosis after oxidative stress.

Methods

Mice. Mice were provided access to food and water ad libitum and maintained on a 12-hour light/dark cycle. Hmox1 genomic fragment of 9 kb, encompassing exons 1–5 and surrounding sequences, was obtained by high fidelity PCR from the Hmox1 locus and cloned into a gene targeting vector (pMSCV-loxp-dsRed). The targeting vector was modified by inserting the proximal loxP site into intron 2 and the distal loxP site, including an FRT-flanked neomycin-resistance gene subsequently inserted downstream of exon 5. After the neo gene was excised from the targeted locus by Flp treatment, neo-minus cells were microinjected into C57BL/6J mouse blastocysts, then implanted into pseudo pregnant recipients. High-chimera males were mated with C57BL/6J females to produce F1 heterozygotes C57BL/6J mouse blastocysts, then implanted into pseudo pregnant recipients. High-chimera males were mated with C57BL/6J females to produce F1 heterozygotes Hmox1fl/+ (HO-1fl/+). Mice were genotyped using a multiplex PCR reaction with 3 primers: Hmox1F, 5′ tctatgactgcttgagg 3′; Hmox1R, 5′ gtgtagatcctagacgtgaa 3′; and Dsred-R, 5′ ggaaggacagtcttgatgcg 3′. Mice harboring a homozygous conditional null mutation in HO-1fl/fl were crossed to transgenic Myh6MerCreMer mice (The Jackson Laboratory) that express Cre recombinase under the control of the cardiomyocyte-specific Myh6 promoter to generate cardiomyocyte-specific Hmox1 KO mice (Supplemental Figure 1). All mice were on a C57BL/6J background and Cre littersmates were used as controls (WT/Cre). To selectively delete cardiomyocyte Hmox1, 8- to 10-week-old mice were given 40 mg/kg tamoxifen for 3 days. This resulted in rapid and effective knockdown of the Hmox1 gene without cardiotoxicity.

Respiration measurements. Hearts were quickly harvested from euthanized mice, and left ventricular muscle fibers were separated in cold, calcium-free isolation buffer (88) and exposed with gentle mixing to 1 mg/ml collagenase (Sigma-Aldrich, C0130) for 10 minutes followed by saponin (50 g/ml, Sigma-Aldrich, S4521) for 30 minutes to permeabilize the cardiomyocytes. The fibers were thoroughly washed 3 times in respiration buffer to arrest permeabilization. The O2 uptake of the fibers was measured in a temperature-controlled respirometer (Mitocell MT200; Strathkelvin Instruments) using malate (5 mM) plus glutamate (10 mM) or succinate (10 mM) (State 4). Maximum respiration rates (State 3) were obtained by adding ADP (2.5 mM). The values are reported as μmol O2/hr/mg of fiber protein. No increase in respiration was observed by addition of 10 mM cytochrome c after ADP.

Oxygen exposures. Mice, aged 8–10 weeks, were placed in cages in airtight chambers (50 × 50 × 30 cm) with access to food and water ad libitum and exposed to 100% O2 for 48 hours, and then moved to filtered room air for 8 days. The chamber oxygen concentration was monitored continuously with an oxygen analyzer (Vascular Technology) as described (89). Mouse hearts from these exposures were harvested immediately or at day 19, snap frozen in liquid N2, and stored at −80°C. Fresh hearts were fixed in 10% formalin for histology or 0.4% gluteraldehyde for electron microscopy (Supplemental Figure 2).

Oxidative stress assessment. To evaluate tissue lipid peroxidation, levels of MDA were determined in fresh ventricle using assay kits (Abcam) with the manufacturer’s instructions. To detect protein oxidation, we used the OxylBlot Oxidized Protein Detection kit (Chemicon). Protein homogenates from normoxic and hyperoxic heart were derivatized with 1,3-dinitrophenylhydrazine for 15 minutes according to the manufacturer’s instructions, and dot blot analysis was performed and the data densitized.

Histology and EM. Hearts were isolated and perfusion-fixed at a constant pressure (80 mmHg) with 10% formalin, dehydrated in 70% ethanol, and embedded in paraffin. Sections were stained with H&E, and gross morphology was viewed under a low-power field (×5). Masson’s trichrome stain was used to evaluate the degree of fibrosis in hearts. The extent of fibrosis in these photomicrographs was quantified by a blinded observer using the Nikon image software (NIS-Elements Basic Research 3.2). For immunofluorescence, sections were stained with CS primary antibodies (1:200; GeneTex Inc.). Sections were then extensively washed in PBS and incubated in secondary anti-rabbit IgG, Alexa Fluor 488 (Thermo Fisher Scientific, catalog A-11034). Conventional fluorescence images were obtained using a Nikon Microphot-FXA fluorescence microscope. Small pieces of left heart were fixed in 2% glutaraldehyde and 2% paraformaldehyde at room temperature for 2 hours, followed by 24 hours at 4°C. Specimens were rinsed in 0.1 mol/l phosphate buffer (PB), pH 7.4, and fixed in 1% osmium tetroxide in PB (pH 7.4) at 4°C for 1 hour.
1 hour, and then dehydrated in graded ethanol, followed by propylene oxide, and embedded in Eponate 812 (Ted Pella Inc.). Ultrathin sections (around 60–70 nm) from longitudinal parts were prepared and contrasted with uranyl acetate, followed by lead citrate, and examined in Philips CM12 electron microscope.

**TUNEL assays.** Thin paraffin-embedded heart sections were used for the TUNEL assay commercial kit (In Situ Cell Death Detection Kit, Roche Diagnostics) following the manufacturer’s instructions. Negative and positive control sections were prepared with label solution only or by preinduction of strand breakage with recombinant DNase I. Nuclei were counterstained with DAPI. Sections from 4–5 hearts in each group were used in the staining, and at least 5 unique fields/section were quantified in each experiment by a blinded observer using Nikon image software (NIS elements BR 3.2).

**Echocardiography.** Cardiac function in WT/Cre and HO-1(CM)⁻/⁻ normoxic or hyperoxic mice was compared at day 19 by transthoracic echocardiography performed using a Vevo770 Ultrasoundograph (VisualSonics) equipped with a 45-MHz imaging transducer. The mice were lightly anesthetized with 1%–1.5% isoflurane, and body temperature was maintained at 37°C. Two-dimensional (2-D) mode in the parasternal long axis and parasternal short axis at the mid-papillary muscle level were imaged. From this parasternal short axis view, the 2-D–guided M-mode across the anterior wall and posterior wall were recorded. Left ventricular anterior wall (LVAW), left ventricular interior diameter (LVID), and left ventricular posterior wall thickness (LVPW) at systole (s) and diastole (d) were measured. Relative wall thickness (RWT) was calculated from the LVIDd and LVPWd measurements (2 × LVPWd/LVIDd). %EF was calculated as (EDV − ESV)/EDV × 100%, where EDV and ESV are end-diastolic and end-systolic volume, respectively.

%FS was calculated by (LVIDd − LVIDs)/LVIDd × 100%.

**qPCR.** Total RNA was isolated from hearts using the RNeasy-4 PCR kit (Thermo Fisher Scientific) according to the manufacturer’s instructions. Samples were treated with DNase I (4 units) for 2 hours at 37°C that was then inactivated, and the cDNA was prepared with a high-capacity cDNA archive kit (Applied Biosystems), and amplifications were performed on 10 ng total mtDNA using PCR primers (cyt b-sense [cyt b-s] and cyt b-antisense [cyt b-as]). One copy of linearized pGEMT-cyt b vector was used for standard mtDNA quantification (90). The cyt b probe, 5’ FAM-ttcctccagaaacaggatcaaa-TAMRA 3’, contained FAM (6-carboxy-fluorescein) at the 5’ end as a fluorescent reporter dye and TAMRA (6-carboxy-tetramethyl-rhodamine) at the 3’ end as a quencher dye selected from a highly conserved region of mouse cyt b gene. Serial dilutions of 1 × 10³ to 1 × 10⁶ copies of standard cyt b plasmid were prepared for a standard curve. Samples were tested for mtDNA at 1:100 and 1:1,000 dilutions. Samples were analyzed in triplicate; mtDNA copy number/ng DNA was determined relative to the standards of known mtDNA copies per dilution.

**Immunoblot analysis.** Protein extracts from mouse hearts (n = 5–6 per group) were analyzed by Western blotting as described (90). Total heart protein, mitochondrial protein, or nuclear protein were resolved after fractionation by SDS-PAGE on gradient or 14% gels (Bio-Rad), transferred to polyvinylidene difluoride membranes (Millipore), blocked with 4% nonfat dry milk in Tris-buffered saline with Tween 20, and probed overnight at 4°C with antibodies against HO-1 or HO-2 (1:1,000; Enzo Life Sciences), mitochondrial transcription factor A (Tfam; 1:1,000; developed in our laboratory), CS (1:1,000; GeneTex, catalog GTX110624), NRF-1 (1:2,000, in house), PGC-I (1:1,000; Abcam, catalog ab54481), Pink1 (1:500; Abcam, catalog ab23707), PARK2 (1:500; Santa Cruz Biotechnology Inc., catalog sc-32282), ANF (1:500; Santa Cruz Biotechnology Inc., catalog sc-18811), BNP (1:1,000; Santa Cruz Biotechnology Inc., catalog sc-18813), LC3I/II (1:1,000; Cell Signaling Technology, catalog 41085), p62 (1:1,000; Cell Signaling Technology, catalog 8025), Atg4B (1:500; Santa Cruz Biotechnology Inc., catalog sc130968), Beclin1 (1:1,000; Cell Signaling Technology, catalog 37385), TGF-β (1:500; Santa Cruz Biotechnology Inc., catalog sc-8829), or CCN5 (1:500; Santa Cruz Biotechnology Inc., catalog sc-25442). Protein loading was confirmed by nuclear protein Lamin B (Santa Cruz Biotechnology Inc, sc-374015), tubulin (1:1000; Sigma-Aldrich, catalog T-5168), or the mitochondrial porin (1:500;
Santa Cruz Biotechnology, catalog sc-8829) and/or Coomassie blue staining. After incubation with primary antibody, the membranes were treated with horseradish peroxidase–conjugated secondary antibody (1:2,000, Santa Cruz Biotechnology Inc., catalog sc-516102) and developed by enhanced chemiluminescence (Western Blotting Luminol Reagent, Santa Cruz Biotechnology Inc.). The protein bands were quantified on digitized images in the mid-dynamic range using Quantity One (Bio-Rad). Densitometry measurements were normalized to tubulin, Porin, Lamin B, or Coomassie in the same samples.

**ChIP analysis.** ChIP assays were performed in cardiac cell nuclei that were sonicated in shearing buffer. Shearing effectiveness was confirmed by electrophoresis on ethidium bromide–stained agarose gels. Samples were processed for immunoprecipitation using ChIP IT assay kit (Active Motif) and NRF-1 antibody (5) at a concentration of 2–5 μg/ml. After precipitation, cross-linking was reversed, and PCR was carried out in the precipitated and input samples using 1 μl of each sample (input DNA dilution 1:10, immunoprecipitated fractions undiluted) in PCR buffer (Qiagen) containing dNTP (Invitrogen) and Taq DNA polymerase (Qiagen) using primers –98 5′ ctgcgcggaggaaccttac 3′ and +165 5′ gcgcggagagattgtacct 3′ to amplify the selected NRF-1 site on the murine Park2 promoter. The primers –149 5′ ttgatcacaaccctcgacc 3′ and +6 5′ caacaacaacactcgggggc 3′ were used to amplify the selected NRF-1 site on the Pink1 promoter. DNA concentration was measured and standardized across samples prior to PCR. PCR products were analyzed by agarose gel electrophoresis.

**Cell culture assay.** HL-1 cardiomyocytes (1 × 10⁴ per treatment) were cultured in flasks coated in fibronectin (12.5 mg/l) gelatin (0.02%) containing Claycomb medium (Sigma-Aldrich) supplemented with 10% FBS (PAN Biotech GmbH), 2 mM L-glutamine (PAN Biotech GmbH), 0.1 mM norepinephrine (Sigma-Aldrich), ascorbic acid 0.3 mM (Sigma-Aldrich), 100 U/ml penicillin (Sigma-Aldrich), and 100 μg/ml streptomycin (Sigma-Aldrich) in a humid atmosphere of 5% CO₂/95% air at 37°C (5). Cells were transfected in 6-well plates at 70%–80% confluence with pGFPCRS vector containing HO-1 oligonucleotide sequence for optimal suppression of HO-1 (Origene, TR30018), HO-1 TrueORF (Origene, MR203944), or NRF-1 TrueORF (Origene, MG208550) using FuGENE-HD (Promega, E2311). After 48–72 hours, the cells were harvested near confluence for further measurements. To confirm HO-1 or NRF-1 overexpression or knockdown, qPCR was used. We used the MTT assay to estimate total cellular metabolic activity based on the reduction of the compound to the formazan by mitochondrial dehydrogenases (91). Beating rate was estimated by counting the number of beats per minute in 5 different cell clusters in 4 blinded experiments. Each experiment was conducted in triplicate and at least repeated 3 times.

**Statistics.** Analysis was performed after a normality test, and results were expressed as mean ± SEM. Data from multiple groups were compared by two-way ANOVA. Fisher’s protected least significant difference tests were used for post-hoc corrections. Student’s t test assuming unequal variance was used for comparisons involving two groups at any single time. The Wilcoxon rank-sum test was performed only when data were not normally distributed. P < 0.05 was considered significant.

**Study approval.** All animal experiments were approved by the Institutional Animal Care and Use Committee at the Duke University Medical Center, Durham, North Carolina, USA.

**Author contributions**

HBS conceptualized and designed experiments, collected and assembled data, analyzed and interpreted data, wrote the manuscript, and gave final approval of the manuscript. CAP conceptualized and designed experiments, assembled data, analyzed and interpreted data, wrote the manuscript, gave final approval of the manuscript, and received financial support. JEK collected and interpreted the data, offered technical assistance, and approved of the manuscript.

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