Differential regulation of cysteine oxidative post-translational modifications in high and low aerobic capacity

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Given the association between high aerobic capacity and the prevention of metabolic diseases, elucidating the mechanisms by which high aerobic capacity regulates whole-body metabolic homeostasis is a major research challenge. Oxidative post-translational modifications (Ox-PTMs) of proteins can regulate cellular homeostasis in skeletal and cardiac muscles, but the relationship between Ox-PTMs and intrinsic components of oxidative energy metabolism is still unclear. Here, we evaluated the Ox-PTM profile in cardiac and skeletal muscles of rats bred for low (LCR) and high (HCR) intrinsic aerobic capacity. Redox proteomics screening revealed different cysteine (Cys) Ox-PTM profile between HCR and LCR rats. HCR showed a higher number of oxidized Cys residues in skeletal muscle compared to LCR, while the opposite was observed in the heart. Most proteins with differentially oxidized Cys residues in the skeletal muscle are important regulators of oxidative metabolism. The most oxidized protein in the skeletal muscle of HCR rats was malate dehydrogenase (MDH1). HCR showed higher MDH1 activity compared to LCR in skeletal, but not cardiac muscle. These novel findings indicate a clear association between Cys Ox-PTMs and aerobic capacity, leading to novel insights into the role of Ox-PTMs as an essential signal to maintain metabolic homeostasis.

High aerobic capacity (i.e. the capacity to use oxygen), is associated with metabolic benefits, cardiovascular protection and longevity1–5. To determine the role of intrinsic aerobic capacity to prevent metabolic diseases and improve longevity, artificial selections for low and high intrinsic running capacity (LCR/HCR) were performed in rats6,7. Low-capacity runner (LCR) rats are insulin-resistant, hyperglycemic, hyperlipidemic, obese and present high cardiovascular risk factors compared to high-capacity runners (HCR) rats7–9. Remarkably, survival rate is ~45% higher in HCR than LCR rats9. While this rat model establishes that intrinsic components of oxidative energy metabolism are inherently associated with health and longevity, elucidating the mechanisms by which high aerobic capacity regulates whole-body metabolic homeostasis is a major research challenge.

Skeletal and cardiac muscles enhance the production of reactive oxygen species (ROS) during high-energy demands, such as during myofibers contraction10. ROS consists of radicals and non-radical species produced

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by the partial reduction of oxygen in different sources, including mitochondria, NADPH oxidase and perox-

isomes11. ROS can induce reversible and irreversible modifications to different proteins. Examples of revers-

ible modifications are disulfides, S-glutathionylation and S-nitrosylation, while examples of irreversible are

carbonylation and sulfonic acid12. Cysteine (Cys) is an amino acid highly susceptible to reversible oxidative

post-translational modifications (Ox-PTMs) due to the presence of a thiol side chain. Cys Ox-PTMs may regulate

cellular homeostasis in several tissues, including skeletal and cardiac muscles13–18, but the putative relationship

between Ox-PTM and intrinsic components of oxidative energy metabolism is poorly understood. In this sense,

mass-spectrometry-based proteomic analysis is a powerful tool to investigate the post-translational modifications

of the proteome, including Ox-PTMs18.

Here, we hypothesized that Ox-PTMs of cysteine are associated with high intrinsic aerobic capacity in skel-

etal and cardiac muscles. We determined the metabolic phenotype and the Cys Ox-PTM profile in both skeletal

and cardiac muscles of LCR and HCR rats. HCR rats had superior mitochondrial content and total (GSH) and

oxidized glutathione (GSSG) levels compared to LCR rats in skeletal muscle, while only modest changes were

observed in cardiac muscle. A redox proteomic screening revealed a different Cys Ox-PTM profile between HCR

and LCR. HCR rats showed higher number of oxidized Cys residues in proteins of skeletal muscle than LCR,

while the opposite pattern was observed in the heart. The most significantly oxidized protein in the skeletal

muscle of HCR rats was malate dehydrogenase (MDH1), and HCR rats showed higher MDH1 activity than LCR

rats in the skeletal muscle, but not in the cardiac muscle. This study provides new insights into the role of Cys

Ox-PTMs as essential signals to maintain metabolic homeostasis and opens the perspective to explore Ox-PTMs
to counteract metabolic diseases.

Results

HCR rats display higher aerobic capacity, lower body mass and slower resting heart rate. To

confirm whether the 32th generation of HCR/LCR rats maintains the distinctive phenotypes of high and low

intrinsic aerobic capacity, we assessed oxygen consumption (VO₂) during a maximal incremental running test.

As expected, HCR rats displayed remarkable running performance (Fig. 1a,b), and higher basal VO₂ and peak

oxygen consumption (VO₂peak) than LCR rats (Fig. 1c,d). HCR rats also displayed lower body weight (Fig. 1e)

and slower resting heart rate compared to LCR rats (Fig. 1f). No differences were observed in plantaris muscle mass,

heart mass and other basal and dobutamine stress echocardiographic parameters between HCR and LCR rats.
muscles are better prepared than LCR’s to cope with oxidative damage. Plantaris muscle and cardiac muscle mass and echocardiography in LCR and HCR rats. Data are separated by 2D-GE. Of interest, infrared scanning of fluorescent gels revealed a more oxidized redox status of the skeletal and cardiac muscles. We first performed fluorescent two-dimensional gel electrophoresis (2D-GE) of the different muscle phenotypes observed in HCR and LCR rats, we screened redox regulated proteins in both groups. After 2D-GE, the ratio of the intensity between the two fluorophores (oxidized:reduced Cys residues) in a given region (DY-680 and DY-780, Dynomics) for reduced (red color-DY-780) and oxidized (green color-DY-680) thiol Cys residues were labeled with fluorescent dyes absorbing and emitting at different wavelengths on the infrared gels. The intensity at each spot reflects the Cys residues redox state of corresponding protein. As the value obtained is a ratio, it is independent of protein amount, allowing a more precise comparison of LCR and HCR cell extracts independently at each spot.

Table 1. Plantaris muscle and cardiac muscle mass and echocardiography in LCR and HCR rats. Data are expressed as mean ± SEM.

|                      | LCR (n = 10) | HCR (n = 10) |
|----------------------|-------------|-------------|
| Plantaris (g)        | 0.23 ± 0.01 | 0.20 ± 0.01 |
| Left ventricle (g)   | 0.54 ± 0.02 | 0.55 ± 0.01 |
| Right ventricle (g)  | 0.16 ± 0.01 | 0.14 ± 0.01 |
| Atria (g)            | 0.04 ± 0.01 | 0.04 ± 0.01 |
| Left ventricular anterior wall thickness in diastole (mm) | 1.47 ± 0.16 | 1.37 ± 0.11 |
| Left ventricular anterior wall thickness in systole (mm) | 2.40 ± 0.14 | 2.29 ± 0.17 |
| Left ventricular end-diastolic diameter (mm) | 6.09 ± 0.22 | 6.04 ± 0.16 |
| Left ventricular end-systolic diameter (mm) | 3.14 ± 0.24 | 3.00 ± 0.22 |
| Fractional shortening (%) | 45.68 ± 1.88 | 47.06 ± 2.78 |
| Ejection fraction (%) | 75.33 ± 2.15 | 76.57 ± 2.81 |

HCR rats have higher mitochondrial content and antioxidant capacity in the skeletal muscle. To determine whether HCR and LCR rats have a different skeletal muscle oxidative phenotype, we measured the mitochondrial respiratory rate in isolated plantaris muscle fibers. HCR displayed higher respiratory rate than LCR rats (Fig. 2a). Independent experiments in the presence of inhibitors for mitochondrial complexes I (rotenone) or II (malonic acid) revealed that both complexes have contributed to the higher state 3 respiratory rate observed in the HCR rats (Fig. 2b,c). HCR rats also presented higher citrate synthase activity than LCR rats in the plantaris muscle (Fig. 2d), while no difference was observed between groups neither in the respiratory control ratio (Fig. 2e) nor when respiratory rate data was normalized by citrate synthase activity (data not shown). Thus, our data suggest that mitochondrial respiratory rate is higher in HCR rats due to a higher mitochondrial content in the skeletal muscle. To test whether HCR rats display higher mitochondrial content than LCR, we evaluated the mRNA expression and protein content of the mitochondrial complexes, and we confirmed higher abundance of mitochondrial complexes in HCR than LCR (Fig. 2f,g and Supplemental Fig. 2).

Consistent with a higher mitochondrial content and respiratory rate, absolute hydrogen peroxide (H₂O₂) emission rate was increased in HCR rats compared with LCR rats in the plantaris muscle (Fig. 2h). However, HCR rats produced lower H₂O₂ when data were normalized to resting oxygen consumption (Fig. 2i). HCR and LCR rats also have similar fluorescent DHE oxidation products in plantaris muscle cross-sections (Fig. 2j). HCR rats displayed higher total glutathione levels and oxidized glutathione (GSSG) than LCR rats in plantaris muscle (Fig. 2k,l), but no changes were observed in superoxide dismutase (SOD) or catalase activity (Fig. 2m,n). In addition, HCR and LCR rats did not show differences in the protein carbonyl levels (Fig. 2o), which is an irreversible oxidation reaction and a biomarker of oxidative damage. Therefore, these findings show that HCR’s skeletal muscles are better prepared than LCR’s to cope with oxidative damage.

HCR rat’s metabolic phenotype in the cardiac muscle. Because previous evidence suggested that metabolic and oxidative profiles can differ significantly between cardiac and skeletal muscles, we evaluated whether the differences observed in the skeletal muscle would display a similar or distinct pattern in the cardiac muscle. In this way, HCR displayed a modest increase in state 3 respiratory rate (Fig. 3a), with no differences in the respiratory rate when the cardiac muscle fibers were incubated with inhibitors for mitochondrial complexes I (Fig. 3b) or II (Fig. 3c). In addition, HCR and LCR rats displayed similar citrate synthase activity (Fig. 3d), respiratory control ratio (State 3-State 4) (Fig. 3e), mitochondrial complexes protein content (Fig. 3f), and H₂O₂ production (Fig. 3g,h). HCR rats presented lower total glutathione levels than LCR rats with similar GSSG levels (Fig. 3i,j) and SOD activity (Fig. 3k). In contrast, HCR rats displayed higher catalase activity than LCR rats (Fig. 3l) and unchanged protein carbonyl levels (Fig. 3m). These findings suggest that HCR rats have a modest increase in mitochondrial respiratory rate and antioxidant defense in the cardiac muscle when compared with LCR rats.

HCR and LCR rats display different Ox-PTM profile between skeletal and cardiac muscles. The data described above demonstrate that HCR rats have a pronounced increase in oxidative metabolism in the skeletal muscle while a modest increase in the cardiac muscle. To determine whether Ox-PTMs are associated with the different muscle phenotypes observed in HCR and LCR rats, we screened redox regulated proteins in both the skeletal and cardiac muscles. We first performed fluorescent two-dimensional gel electrophoresis (2D-GE) of plantaris muscle proteins to visualize and compare Cys redox state between HCR and LCR rats. Plantaris muscle Cys residues were labeled with fluorescent dyes absorbing and emitting at different wavelengths on the infrared region (DY-680 and DY-780, Dynomics) for reduced (red color-DY-780) and oxidized (green color-DY-680) thiol groups. After 2D-GE, the ratio of the intensity between the two fluorophores (oxidized:reduced Cys residues) at each spot reflects the Cys residues redox state of corresponding protein. As the value obtained is a ratio, it is independent of protein amount, allowing a more precise comparison of LCR and HCR cell extracts independently separated by 2D-GE. Of interest, infrared scanning of fluorescent gels revealed a more oxidized redox status of HCR plantaris proteins when compared with LCR (Supplemental Fig. 1).
To circumvent the limitation of 2D-GE method, which fails to visualize less abundant proteins in our complex (tissue) samples, we used a gel-free proteomic analysis employing Cys-specific isotopic coded affinity tags (OxICAT), of skeletal and cardiac muscles from HCR and LCR rats. OxICAT revealed 123 proteins in all biological replicates of *plantaris* muscle, of which 107 displayed higher oxidized/reduced ratio ($p < 0.05$ and $\geq 1.5$-fold change) in HCR than LCR, 10 were not different and only 6 proteins with lower oxidized/reduced ratio in HCR.

Figure 2. Skeletal muscle metabolic phenotype in HCR and LCR rats. (a) Oxygen consumption rate in *plantaris* muscle fibers. (b) Complex I and (c) complex II oxygen respiratory rate in *plantaris* muscle fibers. (d) Citrate synthase activity in *plantaris* muscle. (e) Respiratory control ratio (State 3/State 4) in *plantaris* muscle fibers. (f) Mitochondrial complexes mRNA expression in *plantaris* muscle. (g) Quantification of mitochondrial complexes and representative immunoblots in *plantaris* muscle. (h) Total $\text{H}_2\text{O}_2$ production and (i) $\text{H}_2\text{O}_2$ production normalized by oxygen respiratory rate in *plantaris* muscle fibers. (j) Quantification of fluorescent dihydroethidium (DHE) oxidation products and representative images in *plantaris* muscle cross-sections. (k) Total glutathione and (l) oxidized glutathione (GSGG) levels in *plantaris* muscle. (m) SOD and (n) catalase activity. (o) Quantification of carbonyl protein levels in *plantaris* muscle and representative immunoblots. Data are presented as mean $\pm$ SEM *$p < 0.05$, **$p < 0.01$, ***$p < 0.001$ HCR as compared to LCR, $n = 6–12$. 
than LCR (Fig. 4a and Table Supplemental 2). The top 20 redox-modulated proteins in *plantaris* muscle are presented in Table 2. OxICAT also revealed 113 proteins in cardiac muscle. From these 113 proteins, 4 proteins had higher oxidized/reduced ratio (p < 0.05 and ≥1.5-fold change), 67 proteins had unchanged oxidized/reduced ratio, and 42 proteins had lower oxidized/reduced ratio in HCR when compared to LCR (Fig. 4b and Table Supplemental 3). The top 20 redox-modulated proteins in the cardiac muscle are presented in Table 3. Taken together, these data demonstrate that HCR rats presented higher levels of oxidized proteins than LCR rats in the skeletal muscle, while the opposite was observed in cardiac muscle, since more reduced proteins were observed in HCR hearts (Fig. 4c).

We also performed pathway enrichment analysis using KEGG, a database resource for understanding high-level functions of biological systems, which has well-annotated metabolic pathways. KEGG analysis revealed that metabolic pathways were highly enriched by Ox-PTMs in both the skeletal (Fig. 4d and Table Supplemental 4) and cardiac (Fig. 4e and Table Supplemental 5) muscles. Of note, there were overlaps between pathways identified in the skeletal and cardiac muscle (Fig. 4d,e). We next cross-referenced the lists of redox-modulated residues and found 15 proteins present in both lists (Fig. 4f). Twelve of these proteins had more oxidized levels in the skeletal muscle, but more reduced levels in the cardiac muscle of HCR rats (Fig. 4g). The exceptions were the Creatine Kinase M-type (CKM), and Hemopexin (HPX) that displayed more oxidized levels in both the skeletal and cardiac muscles of HCR (Fig. 4g) and the Mitochondrial Carrier Homolog 2 (MTCH2) that presented reduced levels in both tissues of HCR rats.

**HCR rats show higher malate dehydrogenase (MDH1) activity than LCR rats in the skeletal muscle, but not in the cardiac muscle.** Our data indicate that HCR rats have more oxidized proteins in the skeletal muscle and many of these oxidized proteins are important regulators of the oxidative metabolism, including some essential metabolic enzymes. To determine whether there was an association between the redox
state and the activity of metabolic enzymes, we measured the activity of the MDH1, the most significantly oxidized protein in the skeletal muscle of HCR rats. Interestingly, HCR rats display a trend toward increased MDH1 activity ($p = 0.06$) in the skeletal muscle when compared with LCR rats (Fig. 4h), while no significant differences were observed in cardiac muscle MDH1 activity (Fig. 4i). Thus, contrary to what is generally observed for many oxidized enzymes, increased levels of Ox-PTMs in MDH1 did not mediate any loss of catalytic activity. Rather, a slight but sub-significantly increase in activity was observed in plantaris muscle of HCR rats, a muscle type that displays higher mitochondrial content and antioxidant capacity in HCR than LCR rats.

Discussion

Given the association between high aerobic capacity and the prevention of metabolic diseases, elucidating the mechanisms by which high aerobic capacity regulates whole-body metabolic homeostasis is a major research challenge. To test the hypothesis that Ox-PTMs would be associated with intrinsic oxidative capacity in muscle, we applied redox proteomic approach in skeletal and cardiac muscles of a rat model of low and high intrinsic aerobic capacity. We found a clear association between Ox-PTMs and intrinsic aerobic capacity in both the skeletal and cardiac muscles. HCR rats displayed a higher ratio of oxidized to reduced Cys residues in plantaris muscle than LCR rats whereas the opposite was observed for cardiac muscle with a lower ratio of oxidized to reduced Cys residues than LCR rats. Many of these redox-modulated proteins are important regulators of the oxidative metabolism, including the metabolic enzyme MDH1. Notably, HCR rats showed increased MDH1 activity in the skeletal muscle, but not in the heart. These findings open the perspective to consider Ox-PTMs as targets to regulate metabolic homeostasis in the skeletal and cardiac muscle.

In the current study, we used a well-established rat model of high/low intrinsic aerobic capacity, which recapitulates many of the metabolic differences observed in humans. This model originated from a heterogeneous founder population with breeder selection based on the intrinsic running capacity of each rat. The selection started in 1996 and a rotational breeding was performed at each generation. The 32nd generation HCR/LCR...
proteins in the skeletal muscle. MDH1 is known for catalyzing the reversible oxidation of malate to oxaloacetate of mitochondrial efficiency\(^2\). In fact, high UCP3 expression has been observed in skeletal muscle of HCR rats\(^2\). In skeletal muscle, UCP3 may play a major role in energy expenditure and also participate in the determination of mitochondrial efficiency\(^2\). Indeed, the increased skeletal muscle mitochondrial uncoupling could reduce the oxidative damage in HCR rats, which was observed in our study when we normalized H\(_2\)O\(_2\) production per O\(_2\) consumption. In this sense, the greater number of oxidized proteins in \textit{plantaris} muscle of HCR animals could be influencing the mitochondrial uncoupling to reduce the disruption of redox signaling and control damage, resulting in an adaptive role of reactive species as essential signaling molecules in HCR rats.

Although absolute H\(_2\)O\(_2\) emission rate is higher in HCRs than LCRs plantaris muscle, HCR muscle fibers may also possess a more efficient antioxidant system than LCR, so that for each oxygen molecule consumed, less H\(_2\)O\(_2\) accumulates in HCR cells. These differences may not be explained by differential catalase activity, which was similar between HCR vs. LCR, but through other several key antioxidant enzymes (e.g. GSTM2, PDIA3 and HSPB7) increased in HCR when compared with LCR\(^3\). We performed gel-free redox proteomics analysis in skeletal and cardiac muscles of HCR and LCR rats to identify and quantify reduced and oxidized states of the Cys residues. In our study, approximately 80% of the proteins identified in the redox proteomics of HCR \textit{plantaris} protein extracts displayed a higher ratio of oxidized to reduced Cys residues, which was somewhat surprising. Considering that we did not find signs of oxidative damage in the skeletal muscle of HCR rats, we speculate that Ox-PTMs of Cys would not be deleterious for oxidative metabolism, especially because many of the oxidized proteins in HCR rats are enzymes involved in the intermediary metabolism. Ox-PTM of Cys residues are thought to reduce enzyme activities, however the type and degree of the oxidation depends on many factors, such as the pH of the microenvironmen, surrounding amino acids and whether the oxidation is mediated enzymatically or non-enzymatically\(^3\). Accumulating evidence suggests that Ox-PTMs of Cys have functional effects by altering the activity of the proteins even when the protein content has not changed\(^3,33\). Based on that, we evaluated the MDH1 activity in \textit{plantaris} muscle of HCR and LCR rats, the enzyme with highest ratio of oxidized to reduced Cys residues in HCR among all redox modulated proteins in the skeletal muscle. MDH1 is known for catalyzing the reversible oxidation of malate to oxaloacetate.

### Table 2. Top 20 redox modulated proteins in the\textit{plantaris} muscle. Table includes the redox ratio of individual Cys residues within those proteins. The redox state of selected redox Cys residues labeled with both light \(^{1}C\)-form and heavy \(^{13}C\)-form was calculated using the oxidation ratio (oxidized:reduced).

| Protein | ICAT-modified Cysteine | Fold change HCR/LCR |
|---------|------------------------|---------------------|
| Malate dehydrogenase, cytoplasmic (Mdh1) | Cys137/154 | 5.01 |
| Isovaleryl-CoA dehydrogenase, mitochondrial (Ivd) | Cys353 | 4.99 |
| Sarcoplasmic/endooplasmic reticulum calcium ATPase 2 (Atp2a2) | Cys471/560/680 | 4.93 |
| Desmin (Des) | Cys337 | 4.77 |
| Elongation factor Tu, mitochondrial (Tufm) | Cys129/294 | 4.66 |
| Adenylosuccinate synthetase isozyme 1 (Ados1) | Cys64/186 | 4.62 |
| Protein Tubb4a (Tubb4a) | Cys12/308 | 4.56 |
| Amylo-1, 6-glucosidase, 4-alpha-glucanotransferase isoform CRA_a (Agl) | Cys82/128/174/287/779 Cys972/1065/1277 | 4.56 |
| Acetyl-CoA acyltransferase, mitochondrial (Acat1) | Cys196 | 4.51 |
| Aconitate hydratase, mitochondrial (Aco2) | Cys128/391/416/752 | 4.5 |
| Aspartate aminotransferase, cytoplasmic (Got1) | Cys84/397 | 4.49 |
| Dihydrolipoamide dehydrogenase, mitochondrial (Dld) | Cys70/492 | 4.49 |
| Myosin binding protein C, fast-type (Mybpc2) | Cys159/335/352/447/479/566 Cys622/819/1019/1041 | 4.48 |
| Cytochrome b-c1 complex subunit 1, mitochondrial (Uqcc1) | Cys272/386/416 | 4.47 |
| Phosphoglycerate kinase 1 (Pgk1) | Cys50/321 | 4.43 |
| Methionine-R-sulfoxide reductase B1 (Msrb1) | Cys26 | –1.08 |
| NADH dehydrogenase iron-sulfur protein 2, mitochondrial (Ndufs2) | Cys352 | –1.08 |
| Tropomyosin alpha-1 chain (Tpm1) | Cys193 | –1.15 |
| Mitochondrial carrier homolog 2 (C. elegans) (Mtcn2) | Cys80/99 | –3.19 |
| Transferrin (Tfr) | Cys30 | –3.6 |

See Table 2 for further details.
in many metabolic pathways, including the Krebs cycle and malate/aspartate shuttle. We found that Ox-PTM in the Cys137/154 of MDH1 could be associated with slightly increased MDH1 activity in HCR plantaris muscle. The higher MDH1 activity could be associated with the presence of NADH in the Cys residues. Indeed, NADH being at or near to the enzymatic active center can prevent chemical modifications in dehydrogenases enzymes. The Cys Ox-PTM identified in the plantaris muscle of HCR rats are near to the active and NAD binding sites, suggesting that the MDH1 oxidation in the Cys residues could be protected by the NADH, and not affected the enzyme activity. Another explanation comes from a recent study that reported that H2O2 affects kinetics, structure, and thermodynamic stability of MDH1 through Cys oxidation. Interestingly, others have observed that the thioredoxin-reversible homodimerization of MDH1 protects the protein from over oxidation, maintaining the MDH1 function. These data suggest that a balance between oxidation and antioxidant system is necessary to maintain MDH1 and mitochondrial function during increased energy demand conditions.

Cysteine residues on protein surfaces may act as redox buffers to protect proteins from irreversible oxidant damage. Taking into consideration that: 1) HCR rats have higher number of oxidized proteins in the skeletal muscle and higher number of reduced proteins in the cardiac muscle than LCR rats, and 2) neither skeletal nor cardiac muscle of HCR show higher ROS production than LCR, it’s reasonable to assume that skeletal and cardiac muscles antioxidant defense is differentially regulated. In fact, HCR rats displayed higher GSH and GSSG content in the skeletal muscle, but not in the cardiac muscle, which was consistent with differences observed in the mitochondrial phenotype between these tissues. Reversible S-glutathionylation is an Ox-PTM in which a glutathione is added to Cys residues of a target protein. Based on our data, we speculate that an enhanced S-glutathionylation process in the skeletal muscle of HCR rats might protect several proteins from irreversible oxidation, preserving or even increasing the activity of these proteins, such as the MDH1. Overall, we propose that by stimulating reversible oxidation processes, HCR rats are preventing irreversible oxidation, preserving or even increasing the activity of these proteins, such as the MDH1. Furthermore, we acknowledge limitations to our study. First, the association between Ox-PTMs and improved oxidative metabolism indicates that Ox-PTMs may play a role in maintaining metabolic homeostasis. However, further studies exploring a causal relationship between Ox-PTMs of Cys residues and aerobic capacity are necessary. Second, we used a model of high vs. low intrinsic aerobic capacity, but we cannot extend these findings to other conditions of impaired aerobic capacity, such as type II diabetes and obesity, or enhanced aerobic capacity, resulting from endurance exercise training. We encourage new studies to evaluate the effects of metabolic diseases and/or exercise training in the skeletal and cardiac muscle Ox-PTM profile.

In summary, this study demonstrated that Ox-PTMs of Cys are associated with high intrinsic aerobic capacity, and higher ratio of oxidized to reduced Cys residues is positively associated with higher aerobic capacity in the skeletal muscle. This leads to new insights into the role of Ox-PTMs as a key process to maintain metabolic homeostasis under high-energy demand conditions and opens the perspective to explore Ox-PTMs to counteract metabolic diseases.

### Table 3.

| Protein                                                                 | ICAT-modified Cysteine | Fold Change HCR/LCR |
|------------------------------------------------------------------------|------------------------|---------------------|
| LETM1 and EF-hand domain-containing protein 1, mitochondrial (Letm1)  | Cys560                 | 1.66                |
| Sorting and assembly machinery component 50 homolog (Samm50)          | Cys66                  | 1.64                |
| Creatine kinase M-type (Ckmm)                                         | Cys148/258             | 1.34                |
| Hemopexin (Hpx)                                                       | Cys150/233/259/412     | 0.64                |
| Serum albumin (Alb)                                                   | Cys58/127/204/293/307/317/391/422/469/519/600 | 0.94                |
| Protein Ndufb10 (Ndufb10)                                             | Cys78/127              | 0.94                |
| Mitochondrial carrier homolog 2 (C. elegans) (Mitch2)                  | Cys80/99               | 0.94                |
| D-dopachrome decarboxylase (Ddt)                                      | Cys24                  | -1.03               |
| Acyl-CoA synthetase family member 2, mitochondrial (Acsf2)             | Cys78/469              | -1.03               |
| Aldehyde dehydrogenase, mitochondrial (Aldh2)                         | Cys69/394              | -1.06               |
| Carnitine O-palmitoyltransferase 2, mitochondrial (Cpt2)               | Cys85/653              | -1.06               |
| NADH dehydrogenase [ubiquinone] 1 alpha subcomplex subunit 2 (Ndufa2) | Cys22/56               | -1.12               |
| F-actin-capping protein subunit beta (Capeb)                           | Cys63                  | -1.18               |
| Catenin (Cadherin associated protein), alpha 1 (Ctnna1)               | Cys118/344/536         | -1.25               |
| Myomesin 2 (Myom2)                                                    | Cys198/889/1203/1252/1272 | -1.29             |
| Electron transfer flavoprotein subunit beta (Etfb)                    | Cys42                  | -1.32               |
| Isoform 2 of Adenylate kinase 2, mitochondrial (Ak2)                   | Cys93                  | -1.4                 |
| Peptidyl-prolyl cis-trans isomerase A (Ppia)                           | Cys21/163              | -1.43               |
| Succinate-semialdehyde dehydrogenase, mitochondrial (Aldh5a1)         | Cys82/333              | -1.43               |
| Acetyl-CoA acetyltransferase, mitochondrial (Acat1)                   | Cys117/416             | -1.84               |
Methods

Animals and experimental design. All animal procedures were approved by the Norwegian Council for Animal Research (ID 5243), which was in accordance with the Guide for the Care and Use of Laboratory Animals by the European Commission Directive 86/609/EEC. The selection of rats yielding HCR and LCR has been previously described - for details see 6,7. In the current study, two sets (20 animals in each set) of 12 weeks old male rats from the 32° generation were used. Animals were housed (four per cage) in an animal facility under controlled temperature (22 °C) with 12:12 hours light:dark cycle. Animals had ad libitum access to standard laboratory chow and water. The first set of HCR (n = 10) and LCR (n = 10) rats was weighed and submitted to a maximal incremental running test. One week after the running test, rats were anesthetized with isoflurane and killed by decapitation. The heart (atria and ventricles) and plantaris muscle were carefully harvested, weighed, snap-frozen and stored in −80 °C for further enzymatic and colorimetric assays, DHE fluorescent staining, immunobloting, real-time quantitative PCR (RT-qPCR) and redox proteomics. The second set of HCR (n = 10) and LCR (n = 10) rats was submitted to echocardiography. Rats were then anesthetized with isoflurane and killed by decapitation. The heart (atria and ventricles) and plantaris muscle were carefully harvested and immediately stored at 1) ice-cold biopsy preservation solution (BIOPS) for mitochondrial respiration and H2O2 emission measurements or 2) −80 °C for further enzymatic assays and redox proteomics experiments.

Running capacity and oxygen uptake measurements. Rats were submitted to a maximal incremental running test in a metabolic chamber system. Each animal was adapted in treadmill exercise for three consecutive days (i.e. ~10 min in each day at running speed of 6 to 9 m.min⁻¹). Rats were then submitted to a 15-min resting time in order to measure the resting oxygen uptake (VO2). After the resting measurement, animals ran on a treadmill at 15° inclination. The speed started at 6  m.min⁻¹ and was increased by 2 m.min⁻¹ every 2 min until rats were unable to run. The peak oxygen uptake (VO2peak), speed and distance were recorded.

Echocardiography. Heart structure and function were performed using an echocardiography (Veo 770, Visual Sonics, Toronto, Canada) during isoflurane (2%) anesthesia. Left ventricular anterior wall thickness in systole and diastole (LVAWs and LVAWd) were assessed. Left ventricular end-diastolic diameter and end-systolic diameter (LVEDD and LVESD) were used in order to calculate the fractional shortening (FS) as following: FC (%) = [(LVEDD - LVESD)/LVEDD] x 100. After baseline measurement, three different intravenous dobutamine infusions (5, 10 and 20 ug.kg⁻¹.min⁻¹) were performed and the same parameters were recorded after each dose administration. All myocardial structures were manually measured according to the leading-edge method of the American Society of Echocardiography. The examiner (NR) was blinded to the group's allocation.

Mitochondrial respiration and H2O2 emission measurements. Fiber bundles of cardiac and plantaris muscle were isolated and permeabilized by incubation in saponin (50 µg.ml⁻¹). Fiber bundles were rinsed in ice-cold mitochondrial respiration medium containing 0.5 mM EGTA, 3 mM MgCl₂, 60 mM K-lactobionate, 20 mM taurine, 10 mM KH2PO4, 20 mM Hepes, 110 mM sucrose and 1 g/l BSA essentially fatty acid free adjusted to pH 7.1. High-resolution O2 consumption measurements and H2O2 emission were conducted in the OROBOROS O2K Oxygraph (OROBOROS INSTRUMENTS, Innsbruck, Austria). Chambers containing 2 ml of mitochondrial respiration medium at 37 °C was maintained with O2 levels between 400 and 600 µM. Malate
(2 mM), glutamate (10 mM), succinate (10 mM), 10-acetyl-3,7-dihydroxyphenoxazine (Amplex Red; 25 μM) and Horseradish peroxidase (HRP; 0.5 U.mL⁻¹) were added in each chamber. Permeabilized fibers from cardiac (~2 mg) and plantaris skeletal muscle (~4 mg) were then added in the OROBOROS machines. State 2 O₂ consumption rate was recorded. State 3 O₂ consumption rate and H₂O₂ emission was recorded after stimulation with adenosine diphosphate (ADP; 1 mM). State 4 O₂ consumption rate was recorded after inhibition of ATP synthase by oligomycin (2 μg.mL⁻¹). Finally, consecutive carbonyl cyanide p-trifluoromethoxyphenylhydrazone (FCCP) additions (0.5 mM) were performed until observe a plateau effect (i.e. uncoupler effect). In independent experiments, inhibitors for complex I (i.e. rotenone) and II (i.e. malonic acid) were added after ADP phase as an attempt to estimate how complex I and II inhibition affect O₂ consumption rate and H₂O₂ emission, respectively. All analyses were performed in duplicate. Respiratory control ratio was calculated by dividing State 3 by State 4 O₂ consumption rates, which demonstrates the tightness of the coupling between mitochondrial respiration and phosphorylation.

**Citrate synthase and malate dehydrogenase activities.** Citrate synthase activity was determined using the Citrate Synthase Assay Kit (CS 0720, Sigma-Aldrich, USA). Malate dehydrogenase (MDA) activity was determined by a colorimetric method based on the reaction: oxaloacetate + NADH + H⁺ → L-malate + NAD⁺, in which the change in optical density at 340 nm per unit time is a measure of the malate dehydrogenase activity.

**Protein carbonyls.** Protein carbonyls were assessed by using the OxyBlot Protein Detection Kit (S7150; Millipore, USA). Soluble proteins (20 μg) were denatured by SDS and derivatized by DNPH. Proteins were submitted to electrophoresis and immunoblotting accordingly the manufacturer's instructions.

**Glutathione redox status.** Total, reduced, and oxidized (GSSG) glutathione levels were measured by using the Glutathione Fluorescent Detection Kit (K006-F5; Arbor Assay, USA). In order to block the reduced GSSG, 2-Vinylpyridine (2VP) was used accordingly the manufacturer's instructions.

**Superoxide dismutase activity.** Superoxide dismutase (SOD) activity was assessed by using the Sigma 19160 SOD determination kit (Sigma-Aldrich, Switzerland).

**Catalase activity.** Hydrogen peroxide (H₂O₂) decomposition was assessed by following the decay in sample absorbance at 240 nm in the presence of 10 mM H₂O₂.

**DHE fluorescence staining.** Plantaris cross-sections (10 μm) were incubated with DHE (5 μM) in a light-protected incubator at 37°C for 30 min. The sections were washed with phosphate buffered saline and fluorescence was assessed by confocal microscopy as previously described. Quantitative analysis of fluorescent images was performed with ImageJ (NIH, USA).

**Sample preparation for proteomics.** Plantaris and cardiac tissue extracts were pooled and homogenized in degassed 2D buffer (7 M Urea, 2 M Thiourea, 2.5% CHAPS) without DTT (dithiothreitol) at pH 5.0. 300 μg of cardiac protein and 600 μg of plantaris protein were precipitated in methanol (4 V), chloroform (1 V) and H₂O (3 V) and frozen for further labeling to proceed with gel-free and gel-based-proteomics.

| Target mRNA | PCR Primer sequence 5’→ 3’ | Product Size (bp) | GenBank Accession # |
|-------------|-----------------------------|-------------------|---------------------|
| Ndufa9      | F: CTCCAATGTCACGTCTCTGC     | 105               | NM_001100752.1     |
|             | R: GCCACCTTCCCCAGTGGTAT     |                   |                     |
| Sdhb        | F: ATGCAGAAGAGGATGTGG       | 90                | NM_001100539.1     |
|             | R: CCAAGGCTGCTGTGGATCTCCT   |                   |                     |
| Uqcre2      | F: CTCCTAAAGTTGCCCAAAAGC    | 77                | NM_001006970.1     |
|             | R: TGGTAACATCAATTCGTAGAGGCC |                   |                     |
| Cox4i1      | F: GCACAGTGGCAGAAATGTG      | 80                | NM_017202.1        |
|             | R: CCGAAGCAGHACCGAATGAG     |                   |                     |
| Atp5a1      | F: TGACGTTGCGCGTGTAAG       | 67                | NM_023093.1        |
|             | R: GGAACAACTGACCTTCC        |                   |                     |
| Actb        | F: CCTCTCTGGGATTAGGATCTG    | 86                | NM_031144.3        |
|             | R: GAGGCCTTACGGAATCTGCAAG   |                   |                     |
| Cyclophilin | F: TGGCAACAGATGTGCTTTTGGGAAG| 103               | NM_017101.1        |
|             | R: GGTGATCTCTCTCAGGGTCGTTTC  |                   |                     |
| Hprt1       | F: CAGTCCCCAGGTGCATT        | 138               | NM_012583.2        |
|             | R: GCAAGTCTTTTACGTCTGTCAT   |                   |                     |

**Table 4.** Primer sequences for RT-qPCR mRNA analysis.
Two-dimensional difference gel electrophoresis (2D-DIGE). 2D-DE was performed for preliminary proteomic profiling of skeletal muscle in HCR and LCR rats. Skeletal muscle homogenates were submitted to a differential labeling of reduced (DY-680) and oxidized (DY-780) thiol group using two 2D-compatible fluorescent dyes absorbing and emitting at different wavelengths of the infrared region. Differences in Cys residues oxidation between HCR and LCR samples were quantified by the intensity of each fluorophore at each spot as previously described. Spots intensity was quantified and compared between groups using the DECODON software (Greifswald, Germany). The results were confirmed in biological replicates.

Oxidative isotope-coded affinity tags (OxICAT). OxICAT was performed as previously described. This method detects all reversibly oxidized Cys, whereas irreversible oxidation states are not detected. The ICAT reagent exists in an isotopically light 12C-form (i.e. light ICAT) and heavy 13C-form (i.e. heavy ICAT). Briefly, proteins were denatured in order to access and label all reduced Cys irreversibly with light ICAT. Proteins were then reduced with a thiol reductant Tris (2-carboxyethyl) phosphate and labeled with heavy ICAT. Therefore, this method generates chemically identical proteins, but with different mass of their ICAT-label accordantly to the redox state (light ICAT for the reduced proteins and heavy ICAT for the oxidized proteins). Each sample was digested with trypsin, purified and analyzed by mass spectrometry. Biological triplicate was performed using three different pool of samples for each group (n = 10, 5 and 5, respectively) for plantaris and cardiac muscle. Of note, OxICAT is a method able to detect only reversible oxidations.

Mass spectrometry (MS) analysis. After desalting, peptides were dried down in a SpeedVac centrifuge and resuspended in 0.1% formic acid. Peptides were analyzed on a LC-MS/MS platform consisting of an Easy-nLC 1000 UHPLC system (Thermo Fisher Scientific Inc, USA) interfaced with an LTQ-Orbitrap Elite hybrid mass spectrometer (Thermo Fisher Scientific Inc, USA) via a nanospray ESI ion source (Proxeon, Odense). Peptides were injected into a C-18 trap column (Acclaim PepMap100, 75 μm i. d. x 2 cm, C18, 3 μm, 100 Å, Thermo Scientific, USA) and further separated on a C-18 analytical column (Acclaim PepMap100, 75 μm i. d. x 50 cm, C18, 3 μm, 100 Å, Thermo Scientific, USA) using a multistep gradient with buffer A (0.1% formic acid) and buffer B (CH3CN, 0.1% formic acid): From 0–6% B in 5 min, 6–20% B in 78 min, 20–25% B in 8 min, 25–40% B in 5 min, 40–100% B in 4 min, 100% B in 10 min, 100–0% B in 1 min and 9 min with 100% A. The flow rate was 250 nl/min. Peptides eluted were analyzed on the LTQ-Orbitrap Elite hybrid mass spectrometer operating in positive ion- and data dependent acquisition mode using the following parameters: Electrospray voltage 1.9 kV, CID fragmentation with normalized collision energy 35, automatic gain control target value of 1E6 for Orbitrap MS and 1E3 for MS/MS scans. Each MS scan (m/z 400–1600) was acquired at a maximum ion injection time of 200 ms for MS and 120 ms for MS/MS scans.

Protein identification, quantitation, and analysis. Raw data files were analyzed in Proteome Discoverer 1.4 (Thermo Fisher Scientific Inc., US) using the SEQUEST HT search engine with the Dec 2013 version of the Rat protein sequence database from UniProt (UniProt Consortium). Enzyme specified as trypsin with maximum of two missed cleavages allowed was searched. Precursor mass tolerance was 10 ppm and fragment mass tolerance was 0.6 Da. The Cys residue (labeled with OxICAT) oxidation was set as dynamic modification. The Percolator tool was used for peptide validation and a cutoff value of 0.01 for false discovery rate. Only peptides with high confidence were used for final protein identification. In order to further understand the biological relevance of the identified proteins, we performed functional enrichment analysis in the context of the Kyoto Encyclopedia of Genes and Genomes (KEGG) databases using the Enrichr and QIAGEN’S Ingenuity Pathway Analysis (IPA). A p value cut-off of 0.001 was used to identify enriched processes.

Real-time quantitative PCR (RT-qPCR). RNA transcript levels from plantaris muscle were assessed by RT-qPCR, as described before. Total RNA was isolated using RNeasy Mini kit (Qiagen, USA) and RNA concentration and integrity were assessed. cDNA was synthesized using High Capacity cDNA Reverse Transcription Kit (Life Technologies, USA), and the genes NADH-ubiquinone oxidoreductase subunit A9 (Ndufa9), succinate dehydrogenase complex subunit B (Sdhb), ubiquinol-cytochrome c reductase core protein II (Uqrc2), cytochrome c oxidase subunit I (Cox1), and ATP synthase H + transporting mitochondrial F1 complex alpha subunit 1 (Atp5a1) were analyzed. After testing three control genes, ß-actin (Actb), cyclophilin, and hypoxanthine phosphoribosyl transferase 1 (Hprt1), we used Hprt1 as reference gene to normalize the data as it recorded the highest stability. Primers were designed using Primer-BLAST (NCBI) and primer sequences are listed in Table 4.

Imunoblotting. Protein isolation, immunoblotting, and detection were performed as previously described. Immuno blots were incubated with antibodies against oxidative phosphorylation complexes (OXPHOS, MitoSciences/Abcam, #MS604), NDUFa9 (Abcam, #14713), and GAPDH (Santa Cruz, #sc-20358).

Statistical analysis. Data are presented as mean ± standard error mean (SEM). Experimental groups were compared using unpaired two-tailed Student’s t-test. Statistical significance was set as p < 0.05.

References
1. Gulati, M. et al. Exercise capacity and the risk of death in women: the St James Women Take Heart Project. Circulation 108, 1554–9 (2003). 2. Kokkinos, P. et al. Exercise capacity and mortality in black and white men. Circulation 117, 614–622 (2008). 3. Myers, J. et al. Exercise capacity and mortality among men referred for exercise testing. N. Engl. J. Med. 346, 793–801 (2002). 4. Booth, F. W., Roberts, C. K. & Laye, M. J. Lack of exercise is a major cause of chronic diseases. Compr. Physiol. 2, 1143–211 (2012).
5. Fuiz-Ruices, C., Garatcheva, N., Berger, N. A. & Lucia, A. Exercise is the Real Polypill. *Physiology* 28, 330–338 (2013).
6. Koch, L. G. & Britton, S. L. Artificial selection for intrinsic aerobic endurance running capacity in rats. *Physiol. Genomics* 5, 45–52 (2001).
7. Wisloff, U. et al. Cardiovascular risk factors emerge after artificial selection for low aerobic capacity. *Science* 307, 418–20 (2005).
8. Koch, L. G. et al. Intrinsic aerobic capacity sets a divide for aging and longevity. *Circ. Res.* 109, 1162–72 (2011).
9. Rivas, D. A. et al. Low intrinsic running capacity is associated with reduced skeletal muscle substrate oxidation and lower mitochondrial content in white skeletal muscle. *Am. J. Physiol. Regul. Integr. Comp. Physiol.* 300, R835–43 (2011).
10. Powers, S. K. & Jackson, M. J. Exercise-induced oxidative stress: cellular mechanisms and impact on muscle force production. *Physiol. Rev.* 88, 1243–76 (2008).
11. Radak, Z., Zhao, Z., Koltai, E., Ohno, H. & Atalay, M. Oxygen consumption and usage during physical exercise: the balance between oxidative stress and ROS-dependent adaptive signaling. *Antioxid. Redox Signal.* 18, 1208–46 (2013).
12. Poole, L. R. & Nelson, K. J. Discovering mechanisms of signaling-mediated cysteine oxidation. *Curr. Opin. Chem. Biol.* 12, 18–24 (2008).
13. Leichtert, L. I. et al. Quantifying changes in the thiol redox proteome upon oxidative stress in *vivo*. *Proc. Natl. Acad. Sci. USA* 105, 8197–202 (2008).
14. Izquierdo-Álvarez, A. & Martínez-Ruiz, A. Thiol redox proteomics seen with fluorescent eyes: the detection of cysteine oxidative modifications by fluorescence derivatization and 2-DE. *J. Proteomics* 75, 329–338 (2011).
15. Chung, H. S., Wang, S.-B. S.-B., Venkataraman, V., Murray, C. I. & Van Eyk, J. E. Cysteine Oxidative Posttranslational Modifications: Emerging Regulation in the Cardiovascular System. *Circ. Res.* 112, 392–92 (2013).
16. Fu, C. et al. Quantitative analysis of redox-sensitive proteome with DIGE and ICAT. *J. Proteome Res.* 7, 3789–802 (2008).
17. Kondo, T. & Hirohashi, S. Application of highly sensitive fluorescent dyes (CyDye DIGE Fluor saturation dyes) to laser microdissection and two-dimensional difference gel electrophoresis (2D-DIGE) for cancer proteomics. *Nat. Protoc.* 1, 2940–56 (2006).
18. Chiappetta, G. et al. Proteome screens for Cys residues oxidation: the redoxmethod. In *enzymology* 473 (2010).
19. Bollini, R. C., Hoffmann, R. & Fedorova, M. Proteome-wide profiling of carbonylated proteins and carbonylation sites in HeLa cells under mild oxidative stress conditions. *Free Radic. Biol. Med.* 68, 186–195 (2014).
20. Kanoheisha, M. & Goto, S. KEGG: Kyoto encyclopedia of genes and genomes. *Nucleic Acids Res.* 38, 27–30 (2009).
21. Koch, L. G. & Britton, S. L. I. Theoretical and Biological Evaluation of the Link between Low Exercise Capacity and Disease Risk. *Cold Spring Harbor Perspect. Med.* 4, a029867 (2017).
22. Overmeyer, K. A. et al. Maximal Oxidative Capacity during Exercise Is Associated with Skeletal Muscle Fuel Selection and Dynamic Changes in Mitochondrial Protein Acetylation. *Cell Metab.* 21, 468–478 (2015).
23. Tweedie, C. J. et al. Lower oxidative DNA damage despite greater ROS production in muscles from rats selectively bred for high running capacity. *Am. J. Physiol. Regul. Integr. Comp. Physiol.* 300, R54–53 (2011).
24. Seifert, E. L. et al. Intrinsic aerobic capacity correlates with greater inherent mitochondrial oxidative and H2O2 emission capacities without major shifts in myosin heavy chain isoform. *J. Appl. Physiol.*113, 1624–32 (2012).
25. Nauman, J., Janszky, I., Vaten, L. J. & Wisloff, U. Temporal Changes in Resting Heart Rate and Deaths From Ischemic Heart Disease. *Jama* 306, 2579 (2011).
26. Gavini, C. K. et al. Leanness and heightened nonresting energy expenditure: role of skeletal muscle activity thermogenesis. *Am. J. Physiol. Endocrinol. Metab.* 306, E635–47 (2014).
27. Aguer, C. et al. Muscle uncoupling protein 3 overexpression mimics endurance training and reduces circulating biomarkers of incomplete β-oxidation. *FASEB J.* 27, 4213–4223 (2013).
28. Nolands, B. et al. Artificial selection for high-capacity endurance running is protective against high-fat diet-induced insulin resistance. *Am. J. Physiol. Metab.* 293, E31–E41 (2007).
29. Echtay, K. S., Murphy, M. P., Smith, R. A. J., Talbot, D. A. & Brand, M. D. Superoxide Activates Mitochondrial Uncoupling Protein 2 from the *Biochemistry* 277, 47129–47135 (2002).
30. Figueira, T. R. et al. Mitochondria as a source of reactive oxygen and nitrogen species: from molecular mechanisms to human health. *Antioxid. Redox Signal.* 18, 2029–74 (2013).
31. Burniston, J. G. et al. Proteomic analysis reveals perturbed energy metabolism and elevated oxidative stress in hearts of rats with inborn low aerobic capacity. *Proteomics* 11, 3369–79 (2011).
32. Bonini, M. G. et al. Redox control of enzymatic functions: The electronics of life’s circuitry. *IUBMB Life* 66, 167–181 (2014).
33. McDonagh, B., Sakellariou, G. K., Smith, N. T., Brownridge, P. & Jackson, M. J. Differential Cysteine Labeling and Global Label-Free Proteomics Reveals an Altered Metabolic State in Skeletal Muscle Aging. *J. Proteome Res.* https://doi.org/10.1021/pr5006394 (2014).
34. McDonagh, B., Pedrajas, J. R., Padilla, C. A. & Barcena, J. A. Thiol redox sensitivity of two key enzymes of heme biosynthesis and pentose phosphate pathways: Uroporphyrinogen decarboxylase and transketolase. *Oxid. Med. Cell. Longev.* 2013 (2013).
35. Gregory, E. M., Yost, F. J., Rohrbach, M. S. & Harrison, J. H. Selective chemical modification of malate dehydrogenase. *N-ethylmaleimide modification of active center sulphydryl residues. J. Biol. Chem.* 246, 5491–7 (1971).
36. Huang, L. et al. Self-protection of cytosolic malate dehydrogenase against oxidative stress in Arabidopsis. *J. Exp. Bot.* 29 (2017).
37. Requejo, R., Hurd, T. R., Costa, N. J. & Murphy, M. P. Cysteine residues exposed on protein surfaces are the dominant intramitochondrial thiol and may protect against oxidative damage. *FEBS J.* 277, 1465–80 (2010).
38. Plante, E. et al. Dobutamine stress echocardiography in healthy adult male rats. *Cardiovasc. Ultrasound* 3, 34 (2005).
39. Lang, R. M. et al. Recommendations for chamber quantification: a report from the American Society of Echocardiography’s Guidelines and Standards Committee and the Chamber Quantification Writing Group, developed in conjunction with the European Association of Echocardiography. *J. Am. Soc. Echocardiogr.* 18, 1440–63 (2005).
40. Votion, D.-M., Gnaiger, E., Lemieux, H., Mouthys-Mickalad, A. & Serteyn, D. Physical fitness and mitochondrial respiratory function in horse skeletal muscle. *PLoS One* 7, e34890 (2012).
41. Moreira, J. B. N. et al. High- versus moderate-intensity aerobic exercise training effects on skeletal muscle of infarcted rats. *J. Appl. Physiol.* 114, 1029–41 (2013).
42. Bechera, L. R. G. et al. NADPH oxidase hyperactivity induces plantaris atrophy in heart failure rats. *Int. J. Cardiol.* 175, 499–507 (2014).
43. Chen, E. Y. et al. Enrichr: Interactive and collaborative HTML5 gene list enrichment analysis tool. *BMJ Bioinformatic* 14 (2013).
44. Souza, R. W. A. et al. Regulation of cardiac microRNAs induced by aerobic exercise training during heart failure. *Am. J. Physiol. Heart. Circ. Physiol.* 970, apheart.00941.2015 (2015).
45. Jannings, K. R. et al. Autophagy signaling in skeletal muscle of infarcted rats. *PLoS One* 9, e85820 (2014).

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Author Contributions
P.C.B., G.S. and U.W. conceived and designed the study. C.R.R.A., N.R., A.M., G.J.J.S., J.B.N.M., M.N.A., S.L.B., L.G.K. and P.C.B. carried out the animal procedures and in vivo experiments. R.W.A.S., C.R.R.A., A.M., M.N.A., M.W. and P.C.B. performed ex vivo experiments. R.W.A.S., A.M., M.G., L.H., A.S., G.S. and P.C.B. performed proteomic approach and analysis. R.W.A.S., C.R.R.A. and P.C.B. performed data analysis and wrote the manuscript. All authors have participated in data interpretation and manuscript review. All authors approved the final manuscript.

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