Do mice selectively bred for high locomotor activity have a greater reliance on lipids to power submaximal aerobic exercise?

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Running head: Regulation of fuel use in selectively bred high-running mice

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Abstract

Patterns of fuel use during locomotion are determined by exercise intensity and duration, and are remarkably similar across many mammalian taxa. However, as lipids have a high yield of ATP per mole and are stored in large quantities, their use should be favored in endurance-adapted animals. To examine capacity for alteration or differential regulation of fuel use patterns, we studied two lines of mice that had been selectively bred for high voluntary wheel running (HR), including one characterized by small hindlimb muscles (HR\textsubscript{mini}) and one without this phenotype (HR\textsubscript{normal}), as well as a non-selected control line. We evaluated: 1) maximal aerobic capacity (VO\textsubscript{2max}); 2) whole-body fuel use during exercise by indirect calorimetry; 3) cardiac properties; and 4) many factors involved in regulating lipid use. HR mice achieved an increased VO\textsubscript{2max} compared to control mice, potentially in part due to HR cardiac capacities for metabolic fuel oxidation and the larger relative heart size of HR\textsubscript{mini} mice. HR mice also exhibited enhanced whole-body lipid oxidation rates at 66% VO\textsubscript{2max}, but HR\textsubscript{mini}, HR\textsubscript{normal}, and control mice did not differ in the proportional mix of fuels sustaining exercise (% total VO\textsubscript{2}). However, HR\textsubscript{mini} gastrocnemius muscle had elevated fatty acid translocase (FAT/CD36) sarcolemmal protein and cellular mRNA, fatty acid binding protein (H-FABP) cytosolic protein, peroxisome proliferator-activated receptor (\textit{PPAR}) \textalpha{} mRNA, and mass-specific activities of citrate synthase, \textbeta{}-hydroxyacyl-CoA dehydrogenase, and hexokinase. Therefore, high-running mouse lines had whole-body fuel oxidation rates commensurate with maximal aerobic capacity, despite notable differences in skeletal muscle metabolic phenotypes.

Keywords: aerobic capacity, exercise, experimental evolution, lipid oxidation, muscles
Introduction

When energy demand is increased, as during exercise, a major component of maintaining the essential balance between ATP production and utilization is the ability to select an appropriate mixture of metabolic fuels, in addition to adjusting total flux of substrates through metabolic pathways (50). In exercising mammals, the two predominant substrates sustaining energy expenditure in skeletal muscles are lipids (both intramuscular triglycerides and plasma free fatty acids) and carbohydrates (intramuscular glycogen stores and blood glucose; 50). In general, low-to-moderate intensity exercise is primarily fuelled by oxidation of lipids, which constitute a large fraction of energy reserves in mammals. As exercise intensity increases, carbohydrates support a greater proportion of energy expenditure, as they are capable of sustaining high ATP turnover rates (5, 9, 10, 29, 42, 50).

In fact, it has been long appreciated that exercise intensity largely influences metabolic fuel use patterns, at least in the few mammalian species in which exercise metabolism has been studied (5, 9, 10, 29, 41). For example, the "crossover concept" describes a theory wherein a human's fuel use during exercise depends on the interaction between exercise intensity and endurance training status (5). Indeed, when energetic requirements of locomotion are adjusted according to an individual’s maximal capacity for aerobic ATP supply (VO$_{2\text{max}}$), a strong relationship exists between relative exercise intensity ($i.e.$ intensity relative to VO$_{2\text{max}}$) and the proportional mix of carbohydrates and lipids fuelling exercise (29, 41, 50). Surprisingly, this relationship appears to be widely conserved among the mammalian species examined to date, irrespective of allometric variation or diversity in locomotor capabilities (29, 41, 50). Thus, this model of mammalian fuel use represents a valuable tool to formulate hypotheses.
The underlying explanations for this evident unity in mammalian fuel use patterns remain unknown. Likely, many factors influence the fractional contribution of carbohydrates and lipids to energy supply, including recruitment of different muscle fiber types, hormonal regulation, substrate and oxygen availability, and the regulation of enzyme and substrate transporter quantities and activities (6, 29, 31). For instance, both Roberts et al. (41) and McClelland (29) proposed that one mechanism accounting for the conservation of the fuel use pattern may be the progressive recruitment of type IIB (fast glycolytic) muscle fibers with increasing exercise intensity (25, 44), which could be instrumental amongst all mammals in mediating the gradual shift of fuel preference towards carbohydrates (29, 41). However, it is currently unclear what mechanisms cause variation among mammals in absolute fuel oxidation or flux rates, or even if the mechanisms that explain the unity in proportional mix of fuels are common across taxa.

To further explore the underpinnings of the mammalian fuel use pattern during exercise, we used a model of experimental evolution in which different breeding lines of mice have been selectively bred for high levels of voluntary wheel running (46). This allows for the controlled examination of the evolution of diverse voluntary exercise phenotypes; such models provide opportunities for empirical testing of evolutionary hypotheses without such complications as phylogenetic divergences, as occurs with interspecific comparisons (12). Mice from the high running (HR) lines voluntarily run approximately 170-200% more revolutions/day on running wheels than those from non-selected control lines, and have a higher endurance capacity during forced treadmill exercise (13, 34). This animal model includes four replicate HR lines, providing opportunities for the development of multiple solutions (i.e. differential adaptive changes in
subordinate traits) in response to a uniform selective influence (12, 14). For instance, occurrence of a “mini muscle” phenotype has increased in frequency in two of the four HR lines (12, 13). Mice with the “mini muscle” phenotype (HR\textsubscript{mini}) have a drastically reduced content of type IIB muscle fibers in their hindlimb muscles, resulting in increased mass-specific activities of aerobic enzymes and reduced mass-specific activities of anaerobic enzymes in these muscles, relative to mice with the normal muscle phenotype (3, 18, 19, 22).

In addition to the emergence of the “mini muscle” phenotype, selective breeding for high locomotor activity also led to an increased whole-animal VO\textsubscript{2max} (corrected for differences in body mass) in all HR lines (24, 38, 40, 47). Furthermore, female HR\textsubscript{mini} mice achieve a greater VO\textsubscript{2max} than HR mice with the normal muscle phenotype (HR\textsubscript{normal}), but only under hypoxic conditions (38). It has been proposed that heart enlargement in HR\textsubscript{mini} mice could contribute to this increased aerobic performance, especially in hypoxia (39); however, it has not yet been determined if the hearts of selected mice show significant physical (i.e. hypertrophic) or metabolic remodeling. Therefore, one component of our investigation was to characterize physical and metabolic properties of cardiac muscle in these mice.

Lipids are highly chemically reduced, have a high yield of ATP per mole of fuel (high energy density), and in mammals are stored in large quantities compared to carbohydrates. So, on first principles, one might predict that mice from selectively bred HR lines should rely more heavily on this fuel to power submaximal locomotion. We hypothesized that HR\textsubscript{mini} mice in particular would exhibit a shift in the fuel use pattern towards both a greater absolute lipid flux rate and an increased relative lipid contribution to
total aerobic energy expenditure (measured as VO₂), considering the enhanced aerobic capacity of their muscles and reduction in type IIB (fast glycolytic) muscle fibers. To test this hypothesis, we assessed whole-body fuel use during submaximal exercise, and examined key regulatory points of the fatty acid metabolic pathway in the gastrocnemius muscle, a major hindlimb muscle which has notable phenotypic alterations in the HR mini mice (18, 19). Thus, we had a unique opportunity to evaluate the effects of voluntary locomotor behavior and aerobic capacity on the mammalian fuel use pattern without many of the complications of interspecific comparative studies, by using a model of experimental evolution.

Materials and Methods

Animals

All experimental procedures were approved by the University of California Riverside Institutional Animal Care and Use Committee, as well as by the McMaster University Animal Research Ethics Board following Canadian Council for Animal Care guidelines.

We used male mice from an animal model in which four closed lines are selectively bred for high voluntary wheel-running behavior (HR). For details about the selective breeding regimen and general husbandry, refer to Swallow et al. (46). In brief, four replicate HR lines experience within-family selection for total revolutions run on days 5 & 6 of a 6-day period of wheel access when they are young adults. Four additional lines also receive the 6-day period of wheel access, but are bred as controls without any intentional selection. Animals in our study were sampled from one control line (lab designation line
#2), one HR_{\text{mini}} line (line #3), and one HR_{\text{normal}} line (line #8). Hearts were sampled from 8-9 week old generation 56 mice (N = 30 total). 6-9 week old mice from generation 57 were randomly distributed into two batches of 5 mice from each line, and tested for \textit{in vivo} exercise data (N = 30 total). For skeletal muscle measurements, gastrocnemius muscles were also sampled in two batches, using 8-10 week old non-exercised mice (no prior treadmill exposure) from generation 57 (N = 28 total).

During the \textit{in vivo} experiments, mice were individually housed at room temperature (~22°C), and maintained on a 12:12 h light:dark cycle, with lights on at 7:00. Water and laboratory chow (8604 Teklad rodent diet, Harlan Laboratories) were available \textit{ad libitum}, except during fasting periods.

\textbf{Treadmill exercise and indirect calorimetry}

A positive-pressure, flow-through respirometry system was used for all measurements of rates of oxygen consumption (VO$_2$) and carbon dioxide production (VCO$_2$), at room temperature (~22-25°C). Animals were exercised on a custom-built motorized treadmill at an incline of 10°, enclosed in a Plexiglass metabolic chamber (approximate volume 800 mL) and equipped with stimulus electrical grid. Air entering the chamber was scrubbed of H$_2$O by Drierite (W.A. Hammond Drierite Company, Xenia, OH) and of CO$_2$ by soda lime and Ascarite (Fisher Scientific, Pittsburgh, PA), and maintained at a constant flow of 2,000 mL min$^{-1}$ by combined pump and mass-flow controller unit (MFS, Sable Systems International, Las Vegas, NV). Treadmill chamber air was subsampled at 200 mL min$^{-1}$ and scrubbed of H$_2$O by magnesium perchlorate (Fisher Scientific), then passed through an O$_2$ and CO$_2$ analyzer (FoxBox portable analysis instrument; Sable
Systems International). Data were collected with data acquisition software (Expedata; Sable Systems International) at a frequency of 1 sample s\(^{-1}\). The accuracy of this set-up was determined by measuring the combustion of methanol as described previously (31), and found to be ± 5% of theoretical values for absolute rates of VO\(_2\) and VCO\(_2\) and ± 1% of theoretical values for respiratory exchange ratio (RER = VCO\(_2\)/VO\(_2\)). Prior to VO\(_{2\text{max}}\) trials, mice were familiarized with forced treadmill running for 10-15 minutes on two consecutive days. For all exercise trials, mice had a 5 min adjustment period in the chamber before exercise began with a starting speed of 10 or 13 m min\(^{-1}\), and incrementally increased by 3 m min\(^{-1}\) every 1-2 min. After correction for any baseline drift in incurrent O\(_2\) or CO\(_2\) during the run, 10-15 seconds of data at the plateau of VO\(_2\) during the 1-2 minutes of each speed was used for the assessment of VO\(_2\) values. VO\(_{2\text{max}}\) was defined as the maximal steady-state interval at which at least two of the following criteria were satisfied: 1) no change in VO\(_2\) with increasing speed, 2) RER ≥ 1.0, and 3) the mouse no longer maintained position on the treadmill (24, 31, 38, 43, 47). Trial quality was also assessed subjectively by assigning a score between 1 and 5 (47), and runs deemed unacceptable (scores below 3) were repeated. There was a minimum of 24 h between exercise bouts for any individual.

For all submaximal exercise trials, mice were fasted 6-8 h to ensure a postabsorptive state. The order of the two submaximal exercise intensity trials (target 60% VO\(_{2\text{max}}\) and 80% VO\(_{2\text{max}}\), on average 66% VO\(_{2\text{max}}\) and 78% VO\(_{2\text{max}}\), designated as “low-intensity” and “high-intensity” exercise, respectively) was randomized for each mouse, and the trials were performed between 13:00 and 19:00. Initial treadmill speed for submaximal exercise tests
was determined from the relationship between speed and VO₂ during the VO₂max trials; minor adjustments to treadmill speed were made based on real-time monitoring of VO₂ to maintain a constant relative intensity. Sixty seconds of continuous data from between minutes 9 to 11 of each individual trial were used for analyses. Rates of VCO₂ and VO₂ (mL hr⁻¹) were calculated using equation 3b from Withers (52). Lipid and carbohydrate oxidation rates were estimated using indirect calorimetry (11), and estimations were based on the assumption that protein contributions to total oxidation were negligible during aerobic exercise in a postabsorptive state (37). The proportional contribution of lipids or carbohydrates to whole-body oxidation was determined by dividing the estimated lipid or carbohydrate oxidation rate (in μmol O₂ hr⁻¹ g⁻¹) by the measured total rate of oxygen consumption during the exercise bout (VO₂, in μmol O₂ hr⁻¹ g⁻¹).

**Tissue sampling**

All tissue sampling took place between 13:00 and 18:00 h. Non-exercised mice from generation 57 were fasted for 6-8 h prior to euthanasia via decapitation. The left and right gastrocnemius muscles were removed, weighed, and flash-frozen in dry ice with aluminum clamps. The left and right ventricles were separated and sampled in a similar manner, from a separate group of mice (generation 56) that were not fasted prior to euthanization. All tissues were powdered in liquid nitrogen with a mortar and pestle, and stored at -80°C until used.

**Enzyme activity assays**

For all assays, chemicals were purchased from Sigma-Aldrich (Oakville, ON, Canada), unless otherwise indicated. Maximal activities (Vmax) of β-hydroxyacyl-CoA
dehydrogenase (HOAD), hexokinase (HK), carnitine palmitoyltransferase II (CPT II), and citrate synthase (CS) were measured in left ventricles and right gastrocnemius muscles, following previously described procedures (21, 28) with slight modifications. Powdered tissue was diluted 1:20 (mg tissue:μL) with ice-cold extraction buffer (in mM: 100 K$_2$HPO$_4$/KH$_2$PO$_4$, 5 EDTA, 0.1% Triton X-100, pH 7.6) and homogenized with a cooled glass on glass homogenizer.

Maximal HOAD activity was assayed at 340 nm in assay buffer, in mM: 100 TEA-HCl (pH 7.0), in the presence of 0.28 NADH and 5 EDTA, with 0.1 acetoacetyl-CoA added as substrate. Maximal HK activity, assayed at 340 nm, was measured using 50 Hepes as the buffer (pH 7.6), in the presence of 8 MgCl$_2$, 0.5 NADP, 8 ATP, and excess levels of glucose-6-phosphate dehydrogenase (4 U), with 5 glucose added as substrate. Maximal CS activity, measured at 412 nm, was assayed in 100 Tris-HCl (pH 8.0), in the presence of 0.1 2,2'-nitro-5,5'-dithiobenzoic acid (DTNB) and 0.3 acetyl-CoA, with 0.25 oxaloacetate added as substrate. Maximal CPT II activity, measured at 412 nm, was assayed in 40 Tris-HCl (pH 8.0), in the presence of 0.2 DTNB, 1.5 EDTA and 0.05 palmitoyl-CoA, with 5 L-carnitine added as substrate. This assay is believed to measure CPT II activity because CPT I is detergent labile (see 32).

All maximal enzyme activity levels were assayed at 37°C using a SpectraMax Plus 384 spectrophotometer (Molecular Devices). Assays were performed in triplicate, with an additional negative control well (lacking substrate) to correct for background activity. Enzyme activity data are presented as the mean rate of conversion of substrate to product (Unit = μmol min$^{-1}$).
**DNA quantification**

Total DNA content was quantified in the left ventricles (48), in tissue homogenates used for the enzyme activity assays, after they were frozen and re-thawed, and further diluted to 1:40 (mg tissue:μL extraction buffer). In brief, aliquots of tissue homogenates in 2x digestion buffer (in mM: 100 NaCl, 10 Tris-HCl, 25 EDTA, 0.5% SDS, 0.2 mg mL⁻¹ proteinase K, pH 8.0) were digested overnight (18 h) at 55°C. Digested samples were incubated with PicoGreen, and fluorescence was measured (excitation 480 nm, emission 535 nm) by a SpectraMax Gemini XPS fluorescence spectrophotometer (Molecular Devices) and compared to a DNA standard curve (48).

**Sample preparation for immunoblotting and ELISA**

The cytosolic and sarcolemmal fractions of left gastrocnemius tissue were prepared as previously described (7, 30), with slight modifications. Powdered tissue was homogenized with a cooled glass on glass homogenizer, in a buffer containing in mM: 30 Hepes, 210 sucrose, 2 EGTA, 40 NaCl, and a protease inhibitor cocktail (Complete MINI, Roche Diagnostics), at pH 7.4. The homogenate was centrifuged at 600 x g for 10 min at 4°C, and the supernatant from this step was then centrifuged at 10,000 x g for 20 min at 4°C. The ensuing supernatant was diluted (0.75 x volume) with a buffer containing 1.167 M KCl and 58.3 mM Na₄PPi at pH 7.4, and centrifuged at 230,000 x g for 2 h at 4°C. The resulting supernatant was collected as the cytosolic fraction. The pellet was resuspended in a buffer containing 10 mM Tris and 1 mM EDTA, at pH 7.4. This suspension was mixed with 16% SDS (0.33 x volume) and centrifuged at 1,100 x g for 20 min at room temperature, with the ensuing supernatant collected as the sarcolemmal fraction. Total
protein content in cytosol and sarcolemmal samples was quantified with the commercial Pierce BCA Protein Assay kit (Thermo Scientific, Whitby, ON, Canada).

**Immunoblotting**

Fatty acid translocase (FAT/CD36) protein content was measured in sarcolemmal fractions from gastrocnemius muscles, using a commercial CD36 (H-300) antibody and an HRP-linked anti-rabbit secondary antibody (sc-9154 and sc-2313, respectively; Santa Cruz Biotechnology, Santa Cruz, CA). Equal amounts of total sarcolemmal protein (5 μg/lane) were separated by electrophoresis in 10% polyacrylamide gels (Bio-Rad Laboratories, Mississauga, ON, Canada). Protein was transferred to polyvinylidene difluoride membranes (Bio-Rad Laboratories), and blocked with 5% bovine serum albumin (BSA) dissolved in PBS-T (in mM: 1.5 NaH₂PO₄•H₂O, 8.1 Na₂HPO₄, 145.4 NaCl, 0.05% Tween 20, pH 7.4) overnight at 4°C. Membranes were washed with 1% BSA/PBS-T (2 x 5 min), then primary antibody (diluted 1:200 in 1% BSA/PBS-T) for 1 h, 1% BSA/PBS-T (1 x 15 min, 3 x 5 min), secondary antibody (diluted 1:5000 in 1% BSA/PBS-T) for 1 h, and lastly 1% BSA/PBS-T (1 x 15 min, 3 x 5 min). Chemiluminescence (Perkin Elmer, Waltham, MA) was detected on autoradiographic films (Kodak XAR). Relative protein levels were evaluated based on the measure of band volumes, determined using ImageJ software. All samples are expressed normalized against a standard sample (a mixture of sarcolemmal protein pooled from all three mouse lines) that was loaded onto each gel.

**H-FABP ELISA**

Heart-type fatty acid binding protein (H-FABP) content was determined in cytosolic fractions of gastrocnemius muscles, using an enzyme-linked immunosorbent assay (ELISA)
based on the sandwich principle. Equal amounts of total cytosolic protein (0.015 μg) were assayed for each sample, using a commercial ELISA kit designed for mouse H-FABP (HK403; Hycult Biotech). H-FABP protein levels are expressed relative to total cytosolic protein content.

**Real-time PCR**

Real-time PCR was used for the relative quantification of peroxisome proliferator-activated receptor (PPAR) α, medium-chain acetyl-CoA dehydrogenase (MCAD), and muscle- and liver-type CPT-1 (Cpt-1β and Cpt-1α, respectively) mRNA levels in right and left ventricles, using TATA-binding protein (TBP) as the housekeeping gene. Relative quantification of PPARα, peroxisome proliferator-activated receptor gamma coactivator 1-α (PGC-1α), PPARβ/δ, NAD-dependent deacetylase sirtuin-1 (SIRT1), and FAT/CD36 was performed in the left gastrocnemius muscles, using 18S as the housekeeping gene. Primer sequences (Table 1) were either previously published (1, 48) or designed using Primer 3 software. The specificity of the resulting primer pairs (Mobix, Hamilton, ON, Canada) was tested separately for each mouse line, using PCR and gel electrophoresis.

RNA was extracted by homogenization with TRIzol reagent (Invitrogen, Burlington, ON, Canada), based on the acid guanidinium thiocyanate-phenol-chloroform extraction method. Total RNA concentration was measured at 260 nm using a NanoDrop ND-1000 spectrophotometer (Fisher Scientific). To generate cDNA, 1 μg of total RNA was treated with DNase I (Invitrogen), and reverse transcription was carried out by SuperScript II RNase H− reverse transcriptase (Invitrogen) as previously described (48).

5 μL of 1:5 (cDNA:μL water) diluted cDNA was mixed with 12.5 μL SYBR green
(Bio-Rad), 5.5 μL RNase/DNase-free water, and 1 μL each of forward and reverse primers (5 μM) for quantitative real-time PCR reactions. A negative control of water was run on each plate, to ensure a lack of contamination. Reactions were performed in duplicate on a Stratagene MX3000P QPCR system (Stratagene, La Jolla, CA), using SYBR green with ROX as the reference dye. In all cases, the thermal program consisted of a 3 min initial denaturation at 95°C, then 40 cycles of 95°C for 15 s, 60°C for 45 s, and 72°C for 30 s. A dissociation curve analysis was performed to verify the specificity of PCR products. The relative mRNA levels of each sample were calculated using the comparative Ct method (8). All values are normalized against mRNA levels of the housekeeping gene, which did not differ between groups (p > 0.2 for each housekeeping gene). Statistical analyses for gene expression were performed on values normalized to the housekeeping gene, but for the purpose of data presentation these normalized values are expressed relative to mean control values (arbitrarily set at 1.0).

**Statistics**

Statistical analyses were performed using SPSS 11.0 software. Analysis of covariance (ANCOVA) models were used for each dependent variable. Body mass was always included as a covariate, except when it had been accounted for in the dependent variable, or when examining raw ventricular masses independent of body mass. Analyses of FAT/CD36 and H-FABP protein, maximal enzyme activity, and mRNA content were also performed without body mass as a covariate (21, 22), and this did not alter results in most cases. Age and batch were also tested as covariates, and were removed from the model when they had no significant effects. To account for individual variation during
exercise bouts, the covariate of “relative exercise intensity” (i.e. VO$_2$ relative to individual VO$_{2\text{max}}$) was included in the ANCOVA models for fuel oxidation rates and proportional contributions to total oxidation.

For analyses in which a significant effect was detected, pairwise comparisons with a Bonferroni correction were used to compare mouse phenotypes. For all statistical analyses, tests were 2-tailed and the critical $\alpha$-level was set at $p = 0.05$.

**Results**

*In vivo: VO$_{2\text{max}}$ and fuel use during exercise*

Both lines of HR mice exhibited a greater VO$_{2\text{max}}$ than non-selected control mice, regardless of whether VO$_{2\text{max}}$ was analyzed as a mass-specific variable, or alternately, as a whole-animal value with the covariate of body mass ($p < 0.05$; Fig. 1, Table 3). Moreover, the mass-specific VO$_{2\text{max}}$ of HR$_{\text{mini}}$ mice were elevated compared to HR$_{\text{normal}}$ mice ($p < 0.05$; Table 3). However, when VO$_{2\text{max}}$ was analyzed as whole-animal values with body mass as a covariate, the difference between HR$_{\text{mini}}$ and HR$_{\text{normal}}$ mice was not statistically significant. The estimated marginal mean whole-animal VO$_{2\text{max}}$ and SEM for control, HR$_{\text{mini}}$, and HR$_{\text{normal}}$ mice were $4.353 \pm 0.058 \text{ mL min}^{-1}$, $4.933 \pm 0.059 \text{ mL min}^{-1}$, and $4.794 \pm 0.057 \text{ mL min}^{-1}$, respectively, as evaluated at an average body mass of 28.1 g (Fig. 1).

During low-intensity exercise ($66 \pm 1\%$ for controls, $64 \pm 1\%$ for HR$_{\text{mini}}$ mice, and $69 \pm 2\%$ VO$_{2\text{max}}$ for HR$_{\text{normal}}$ mice; overall average $66\%$ VO$_{2\text{max}}$; Table 3), the greater total VO$_2$ of HR mice (Table 3) was primarily supplied by an increased flux of lipids; HR$_{\text{mini}}$ mice exhibited higher absolute rates of lipid oxidation than control mice, a difference which
approached significance for HR_{normal} mice as well (p < 0.05 for HR_{mini} mice, p = 0.057 for 
HR_{normal} mice; Table 3). Although the concurrent rates of carbohydrate oxidation were not 
significantly different among the three mouse phenotypes (Table 3), there were no 
statistically significant differences in the proportional mix of fuels supplying oxidation at 
66% VO_{2max} when fuel oxidation rates were expressed relative to total VO_{2} (Fig. 2A). For 
all mice, approximately 70% of energy was supplied via the oxidation of lipids, with the 
remaining 30% by the oxidation of carbohydrates (Fig. 2A).

During high-intensity exercise (78 ± 1% VO_{2max} for controls, 76 ± 1% VO_{2max} for 
HR_{mini} mice, and 79 ± 1% VO_{2max} for HR_{normal} mice; overall average 78% VO_{2max}; Table 3), 
the absolute lipid and carbohydrate oxidation rates in control mice were equivalent to those 
in HR mice (Table 3). At this increased intensity, carbohydrates contributed more than 
40% towards total oxidation in all mice (Fig. 2B).

**Cardiac properties**

Relative to body mass, HR_{mini} mice had significantly larger left ventricles than both 
control and HR_{normal} mice (p ≤ 0.001; Table 2). This was principally due to a reduced body 
mass in HR_{mini} mice compared to the other 2 lines (p < 0.05; Table 2); however, a slight 
enlargement of whole ventricle mass in HR_{mini} mice approached statistical significance 
even without the inclusion of body mass as a covariate (p = 0.096; Table 2). There were no 
statistical differences in left ventricle DNA content, whether expressed per g wet mass or 
per whole ventricle (Table 2).

In addition to physical changes, both lines of HR mice exhibited some degree of 
cardiac metabolic remodeling compared to control mice, as indicated by increased maximal
HK activity concurrent to reduced maximal CPT II activity (p < 0.05; Fig. 3). However, there were no statistically significant differences in the mRNA content of PPARα, MCAD, CPT-1α, or CPT-1β in either the right or left ventricles of HR_{mini}, HR_{normal}, or control mice (Table 4).

**Skeletal muscle FAT/CD36 and H-FABP**

The skeletal muscle of non-exercised HR_{mini} mice was characterized by a markedly enhanced capacity for fatty acid uptake and intracellular transport. Both FAT/CD36 mRNA levels and FAT/CD36 sarcolemmal protein content was nearly three-fold higher in the gastrocnemius of HR_{mini} mice than in mice with the normal muscle phenotype (p < 0.05; Fig. 4). Additionally, HR_{mini} mice had nearly double the amount of cytosolic H-FABP protein (per mg total cytosolic protein) of HR_{normal} or control mice in the gastrocnemius (p < 0.05; Fig. 5).

**Skeletal muscle metabolic enzyme activities**

The mass-specific maximal activities of CS, HOAD, and HK were greatly augmented in the gastrocnemius of HR_{mini} mice, compared to mice with a normal muscle phenotype (p < 0.05; Fig. 6A). In fact, due to mass-specific enzyme activities nearly double those of control mice (Fig. 6A), in a gastrocnemius less than half the mass (p < 0.05; Table 2), for these enzymes HR_{mini} mice had equivalent total maximal activities as control mice when expressed per whole muscle (Fig. 6B). The mass-specific maximal CPT II activity was not significantly different between HR_{mini} and control mice, as reflected in the reduced whole-muscle maximal CPT II activity in the mini muscles (Fig. 6).

Conversely, although HR_{normal} mice did not have altered mass-specific enzyme
activities compared to control mice, on a whole-muscle basis they exhibited significantly greater maximal activities of CS, HOAD, CPT II, and HK (p < 0.05; Fig. 6B). These differences were also significant with body mass as a covariate (data not shown). The elevation in total muscle metabolic enzyme activity was likely reflective of a slightly increased gastrocnemius mass in HR_{normal} mice, which approached statistical significance (p = 0.053; Table 2). HR_{normal} whole-muscle CS, CPT II, and HK maximal activities were significantly elevated above those of HR_{mini} mice as well (p < 0.05; Fig. 6B).

**Skeletal muscle gene expression and transcriptional regulation**

mRNA levels of **PGC-1α**, **PPARβ/δ**, and **SIRT1** were not significantly different among phenotypes, but **PPARα** mRNA content was approximately 2.5-fold greater in the resting gastrocnemius of HR_{mini} mice than in HR_{normal} or control mice (p < 0.05; Fig. 7A). Furthermore, transcript levels of the PPAR-regulated gene, **FAT/CD36**, were closely correlated with **PPARα** amongst all mice (r = 0.92, p ≤ 0.001; Fig. 7B).

**Discussion**

Our primary goal was to determine if mice selected for high voluntary running have greater whole-body absolute lipid oxidation rates during exercise, and a greater contribution of lipids to total aerobic energy expenditure. Our results indicate that selective breeding for high voluntary exercise leads to an increase in the absolute rate of lipid oxidation when individuals exercise at the same relative intensity. These oxidation rates are scaled with increases in maximal aerobic capacity in both HR_{normal} and HR_{mini} mice, resulting in the proportional contributions by lipids and carbohydrates to total exercise energy expenditure.
being equal amongst all mice. The two selectively bred HR lines examined showed distinct capacities in the gastrocnemius muscle for cellular lipid uptake (FAT/CD36), cytosolic transport (H-FABP), and mitochondrial oxidation (enzyme $V_{\text{max}}$), suggesting that in these distinct high running phenotypes, the response to selective breeding may have targeted different mechanisms to enhance lipid oxidation rates (e.g. capacity for oxidation of circulatory versus intramuscular sources) over that of the control line.

**Aerobic capacity and cardiac properties**

Our study not only confirms a 10-14% rise in VO$_{2\text{max}}$ in HR mice (24, 38, 40, 47), but also indicates that generation 57 HR$_{\text{mini}}$ mice achieve a greater mass-specific VO$_{2\text{max}}$ than HR$_{\text{normal}}$ mice (at least the line [#8] studied here), although this difference was not significantly different on a whole-animal VO$_{2\text{max}}$ basis with body mass as a covariate. Previous studies indicated that the physiological constraints of VO$_{2\text{max}}$ differ between control and HR mice – specifically, that selection for high voluntary wheel running may have resulted in changes to central factors in the oxygen cascade, contributing to the increased aerobic capacity of HR mice (38, 39). As cardiac output is an important central component of oxygen delivery, we characterized some physical and metabolic properties of HR$_{\text{mini}}$, HR$_{\text{normal}}$, and control mouse hearts.

HR$_{\text{mini}}$ mice have a significantly greater left ventricle mass and thus whole ventricular mass than both control and HR$_{\text{normal}}$ mice, relative to body mass (Table 2), which is consistent with the findings of Rezende and colleagues (39). However, without body mass as a covariate, neither left nor whole ventricles are significantly larger in HR$_{\text{mini}}$ mice, and left ventricles of all mouse phenotypes contain comparable levels of DNA (Table 2). Therefore, this difference in relative heart size could be viewed as predominately
reflective of the reduced body masses of HR_{mini} mice, as opposed to cardiac hypertrophy per se. However, relative heart mass can give a good indication of an animal’s ability to perform sustained aerobic locomotion (4, 51), so it is likely that the increased ratio of heart to body size contributes to the elevated VO_{2max} of HR_{mini} mice.

Additionally, both phenotypes of HR mice show evidence of metabolic alterations in the left ventricle. Compared to control mice, HR mice show a reduction in the left ventricular maximal activity of CPT II, an enzyme in the fatty acid catabolic pathway (Fig. 3). Although a non-reversible switch in the basal preferred fuel substrate from fatty acids to carbohydrates is often regarded as a deleterious hallmark of pressure- or volume-overloaded hypertrophic hearts (36), these mice show no statistically significant differences in ventricular mRNA levels of several genes involved in fatty acid catabolism, including \textit{PPAR\alpha}, \textit{MCAD}, and both muscle- and liver-type \textit{CPT-1}. Therefore, it is unlikely that this reduced maximal activity of CPT II in HR left ventricles signifies a diminished capacity for fatty acid oxidation, particularly as there is no change in the maximal activity of HOAD, another fatty acid catabolic enzyme. There is, however, increased maximal activity of HK in the left ventricles of sedentary (\textit{i.e.}, not exercise-trained) HR mice, compared to control mice (Fig. 3). Upregulated HK is also characteristic of physiologically hypertrophic hearts in exercise-trained rats, as a component of the elevated overall metabolic capacity (45). Therefore, differences between HR and control mouse cardiac metabolic enzyme activities may contribute to the enhancement of VO_{2max} and capacity for sustained aerobic exercise in HR mice, in conjunction with the increased relative heart size of HR_{mini} mice.

\textit{Whole-body fuel use during exercise}
The mix of metabolic fuels used across two submaximal relative exercise intensities does not differ statistically among HR_{mini}, HR_{normal}, and control mice, when fuel oxidation rates are expressed as a percentage of total oxygen consumption (Fig. 2). Rather, HR mice exhibit greater rates of lipid oxidation (i.e. enhanced total lipid flux) than control mice during low-intensity exercise (66% VO_{2max}), commensurate with their elevated VO_{2max} (Fig. 1, Table 3). When exercising at a higher intensity (78% VO_{2max}), the increased energetic demand in all mice is supported by an elevated flux of carbohydrates, and the absolute rate of lipid oxidation in control mice matches the rates in HR mice (Table 3).

HR mice primarily achieve greater levels of voluntary wheel running by running faster than control mice on exercise wheels (13-15, 17, 46). Our findings suggest that due to an increased aerobic capacity, HR mice are able to run at these faster speeds predominately sustained by lipid oxidation, whereas mice from non-selected control lines would require a higher proportion of carbohydrates to achieve equivalent absolute exercise intensities. Therefore, an enhanced capacity for absolute lipid oxidation in conjunction with an increased VO_{2max} may have enabled the shift in locomotor speed preference in HR mice, thus allowing them to achieve the greater levels of voluntary exercise for which they are selectively bred (see also 41). In fact, consumption of a high fat diet further increases the amount of voluntary wheel running performed by HR (but not control) mice (33). Interestingly, rats from a line selectively bred for high treadmill endurance capacity with an elevated VO_{2max} exhibit greater in vitro skeletal muscle palmitate oxidation rates compared to a line selected for low endurance capacity (23, 26), signifying that increased lipid oxidation capacity is a common result of selection for running capacity.

Roberts et al. (41) and McClelland (29) have suggested that conserved mammalian
fuel use patterns are the result of the progression of muscle fiber recruitment, with the increasing proportionate reliance on carbohydrates at higher exercise intensities corresponding to the recruitment of fast glycolytic muscle fibers amongst all mammals (25, 44). Intriguingly, HR_{mini} mice do not have a large quantity of type IIB (fast glycolytic) fibers available for recruitment in their reduced-size hindlimb muscles (3, 18, 19), and yet they exhibit the ability to reach a whole-body rate of carbohydrate combustion that is equivalent to mice with the normal muscle phenotype, at least under the submaximal exercise conditions tested here. Moreover, as the proportional reliance on carbohydrates for total oxidation at higher exercise intensities is comparable between HR_{mini} mice and HR_{normal} mice, it may be possible for mammals to differ in muscle fiber recruitment patterns during exercise without deviating from a conserved whole-body pattern of proportionate fuel mix.

In light of these findings, we investigated other potential regulators of metabolic fuel use, to determine if phenotypically divergent HR mice might utilize different strategies to adjust substrate flux. Specifically, we focused here on several regulators of fatty acid uptake and catabolism in the skeletal muscle, in keeping with our stated hypothesis. However, future studies should explore other pathways (see 29, 50), such as carbohydrate or amino acid metabolism, or lactate clearance capacity, to further delineate different mechanisms regulating all of the fuels available to power exercise in mammals.

**Capacity for circulatory fuel uptake into working muscle**

FAT/CD36 is a pivotal sarcolemmal transporter of long-chain fatty acids, and its overexpression can lead to elevated fatty acid oxidation, as well as increased rates of fatty acid transport into giant sarcolemmal vesicles (35). Both cellular mRNA expression and
sarcolemmal protein content of FAT/CD36 is substantially augmented in the gastrocnemius muscle of HR_{mini} mice, compared to HR_{normal} or control mice (Fig. 4), which suggests that the mini muscles of HR_{mini} mice may have an enhanced capacity for uptake of circulatory fatty acids. Furthermore, the gastrocnemius of HR_{mini} mice exhibit an 80% higher cytosolic content of H-FABP (Fig. 5), a protein that serves as an intracellular sink for fatty acids transported across the sarcolemma, and enables their movement through the cytoplasm (16). Although it seems that H-FABP is not a direct regulator of fatty acid translocation across the sarcolemma, increased H-FABP levels could accommodate rapid changes in substrate flux, such as during the transition from resting to contracting muscle (27).

Interestingly, despite these innate biochemical differences in the skeletal muscle, our \textit{in vivo} results suggest that HR_{mini} mice do not have (or do not capitalize upon) an enhanced whole-body capacity to utilize lipids during high-intensity exercise, compared to mice with the normal muscle phenotype. Perhaps the muscle phenotype differences reflect more an increased reliance on circulatory versus intramuscular lipids in the HR_{mini} line.

HR_{mini} mice may also have an enhanced capacity for utilizing circulatory glucose in working muscle. During exercise, when the glucose transporter GLUT-4 has been translocated to the plasma membrane, the phosphorylation of glucose by HK becomes a significant functional barrier of glucose uptake into the myocyte (see 49). While it has previously been shown that there are no differences in constitutive GLUT-4 levels in the gastrocnemius of resting HR_{mini}, HR_{normal}, or control mice, when housed without access to running wheels (17), HR_{mini} mice do have increased mass-specific maximal HK activity in the gastrocnemius (Fig. 6). This elevated HK activity may facilitate enhanced glucose uptake into the gastrocnemius during exercise (see also 3, 18, 22), given that transgenic
mice overexpressing HK II show increased glucose uptake into skeletal muscle during moderate-intensity exercise (20). Indeed, prior to exercise, HR_{mini} mice have larger liver glycogen depots than the other mouse phenotypes (17), and liver glycogen is an important source of energy during exercise for rodents (2).

Although these data point to variation in the capacity for utilizing circulatory fuels, since neither post-exercise intramuscular substrate depletions nor \textit{in vivo} muscle uptake rates for circulatory fuels during locomotion were measured in these mice, further inferences are limited by the complexities of fuel metabolism.

\textbf{Skeletal muscle capacity for oxidative metabolism}

The increased capacity for substrate uptake into the skeletal muscle of HR_{mini} mice is accompanied by an enhanced mass-specific capacity to utilize fatty acids. In addition to the increased mass-specific maximal activities of CS, which is a trademark of the “mini muscle” phenotype’s enhanced aerobic capacity (18, 22), we show that sedentary HR_{mini} mice also have an enhanced capacity for β-oxidation at the level of HOAD specific activity in the gastrocnemius (Fig. 6).

On a whole-muscle basis, HR_{mini} mice tend to have equivalent maximal enzyme activities as control mice (see also 22). Conversely, HR_{normal} mice (at least the line sampled here) appear to exhibit an enhancement of the overall metabolic capacity of the gastrocnemius. Due to the slight (but statistically nonsignificant) muscle enlargement compared to control mice, HR_{normal} mice have significantly greater whole-muscle maximal activities of CS, HOAD, CPT II, and HK than control mice (and HR_{mini} mice, with the exception of HOAD; Fig. 6). Having a bigger muscle mass as opposed to greater mass-
specific activities of metabolic enzymes may be a distinct mechanism that help $HR_{normal}$ mice achieve elevated whole-body fuel oxidation rates (compared to control mice), commensurate with their increased $VO_{2max}$.

**Transcriptional regulation of skeletal muscles properties**

Constitutive gene expression of $PGC-1 \alpha$, $PPAR\beta/\delta$, and $SIRT1$ in muscle are not significantly different among $HR_{mini}$, $HR_{normal}$, and control mice (Fig. 7), suggesting that these transcription factors do not play a major role in the distinct properties of the “mini muscle.” However, $HR_{mini}$ mice do exhibit greatly enhanced $PPAR\alpha$ mRNA expression in the gastrocnemius, and this may be an underlying basis for many of the altered metabolic characteristics of this skeletal muscle. $PPAR\alpha$ is one of the most predominant PPAR isoforms in skeletal muscle, and it is involved in the upregulation of many genes in the fatty acid metabolic pathway (29). For instance, our results show a strong correlation between mRNA levels of $PPAR\alpha$ and $FAT/CD36$ amongst all mice, which suggests a close relationship between the transcript levels of this transcription factor and a target gene (Fig. 7).

**Perspectives and significance**

Despite divergent patterns in daily locomotor behavior and aerobic capacity, the proportionate mix of fuels used across relative exercise intensities was not statistically different for $HR_{mini}$, $HR_{normal}$, and control mice. Instead, this experimental evolution model supports the results of interspecific comparative studies (29, 41, 50), and shows that absolute rates of fuel use during submaximal exercise scale with variation in aerobic capacity amongst diverse mammals. Furthermore, owing to substantial differences in
muscle anatomy and biochemistry, the HR mouse model demonstrates that it may be possible for mammals to differ in muscle fiber recruitment patterns while maintaining a conserved whole-body pattern of fuel use, thus emphasizing the robustness of the mammalian fuel use pattern. Based on our findings, we propose that aerobic capacity and locomotor behavior may be evolutionarily linked to the capacity for lipid oxidation. Through the course of evolution, mammals may exhibit multiple potential mechanisms or “solutions” (sensu Garland et al. (14)) to enhance lipid oxidation rates commensurate with aerobic capacity; biochemical traits can therefore differ between genetically or phenotypically distinct mammals without causing deviation from a conserved whole-body fuel use pattern.
Acknowledgements

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Author contributions

N.M.T., G.B.M., and T.G. contributed to the conception and design of the experiments. All in vivo experimentation and tissue sampling was carried out in the Garland laboratory at the University of California, Riverside, by N.M.T. and H.S. The remaining experiments, including any molecular and biochemical analyses, were performed by N.M.T. in the McClelland laboratory at McMaster University. All authors participated in data analysis and interpretation. N.M.T. wrote the initial draft of the manuscript, and all authors contributed to its revisions and approved the final version.
References


9. Edwards HT, Margaria R, Dill DB. Metabolic rate, blood sugar and the utilization of


40. Rezende E, Kelly S, Gomes F, Chappell M, Garland T, Jr. Effects of size, sex, and voluntary running speeds on costs of locomotion in lines of laboratory mice selectively


48. **Templeman NM, Beaudry JL, Le Moine CMR, McClelland GB.** Chronic hypoxia-


Figure Captions

Figure 1. Individual whole-animal maximal aerobic capacity (VO$_{2\text{max}}$) of control mice and selectively bred HR$_{\text{mini}}$ and HR$_{\text{normal}}$ mice (n = 9-10), presented across the range of body masses.

Figure 2. Proportional contributions to whole-body oxidation rates by carbohydrates and lipids, at average exercise intensities of (A.) 66% VO$_{2\text{max}}$ and (B.) 78% VO$_{2\text{max}}$, in control mice and selectively bred HR$_{\text{mini}}$ and HR$_{\text{normal}}$ mice (n = 9-10). Data are presented as estimated marginal means ± SEM, as adjusted for the covariate of individual relative exercise intensity.

Figure 3. Maximal enzyme activities (relative to g tissue wet weight) for citrate synthase (CS), β-hydroxyacyl-CoA dehydrogenase (HOAD), carnitine palmitoyltransferase (CPT) II, and hexokinase (HK) in the left ventricle of control mice and selectively bred HR$_{\text{mini}}$ and HR$_{\text{normal}}$ mice (n = 7-8). Data are presented as simple means ± SEM. For each enzyme, the presence of letters denotes statistical significance, where bars not sharing a common letter are significantly different from each other (p < 0.05). U = μmol min$^{-1}$.

Figure 4. (A.) Cellular mRNA levels of FAT/CD36 and (B.) sarcolemmal FAT/CD36 protein content, in the gastrocnemius of control mice and selectively bred HR$_{\text{mini}}$ and HR$_{\text{normal}}$ mice (n = 6-10). mRNA levels are expressed corrected against 18S mRNA, and
normalized to the mean control mouse value (arbitrarily set at a value of 1.0). Statistical analyses for gene expression were performed on raw mRNA levels corrected against 18S mRNA. Protein levels are expressed relative to a mixed standard sample, in arbitrary units. The inset in (B.) shows a single representative immunoblot of sarcolemmal FAT/CD36 in control, HR_{mini}, and HR_{normal} mice, respectively (5 μg total protein loaded per gel lane). All data are presented as simple means ± SEM. Bars not sharing a common letter are significantly different from each other (p < 0.05).

**Figure 5.** Cytosolic H-FABP protein content (relative to mg of total cytosolic protein) in the gastrocnemius of control mice and selectively bred HR_{mini} and HR_{normal} mice (n = 8-9). Data are presented as simple means ± SEM. Bars not sharing a common letter are significantly different from each other (p < 0.05).

**Figure 6.** Maximal enzyme activities for citrate synthase (CS), β-hydroxyacyl-CoA dehydrogenase (HOAD), carnitine palmitoyltransferase (CPT) II, and hexokinase (HK) in the gastrocnemius of control mice and selectively bred HR_{mini} and HR_{normal} mice (n = 7-10), expressed (A.) relative to g tissue wet mass or (B.) per whole muscle mass. Data are presented as simple means ± SEM. For each enzyme, the presence of letters denotes statistical significance, where bars not sharing a common letter are significantly different from each other (p < 0.05). U = μmol min^{-1}.

**Figure 7.** Relative gene expression of transcriptional factors in the gastrocnemius of
control mice and selectively bred HR_{mini} and HR_{normal} mice (n= 7-9). In (A.), mRNA levels of \textit{PPAR\(\alpha\)}, \textit{PGC-1\(\alpha\)}, \textit{PPAR\(\beta/\delta\)}, and \textit{SIRT1} are expressed corrected against \textit{18S} mRNA, and normalized to the mean control mouse value (arbitrarily set at a value of 1.0). Data are presented as simple means ± SEM. Statistical analyses were performed on raw mRNA levels corrected against \textit{18S} mRNA; for each gene, the presence of letters denotes statistical significance, where mouse phenotypes not sharing a common letter are significantly different from each other (p < 0.05). In (B.) is the relationship between the transcript levels of \textit{PPAR\(\alpha\)} and \textit{FAT/CD36}, where mRNA levels of individual mice are expressed corrected against \textit{18S}. For this correlation, r = 0.92, p ≤ 0.001.
Table 1. Primer sequences used for real-time PCR mRNA quantification.

<table>
<thead>
<tr>
<th>Gene</th>
<th>Forward Primer (5’-3’)</th>
<th>Reverse Primer (5’-3’)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cpt-1α*</td>
<td>AAACCCACCAGGCTACAGTG</td>
<td>TCCTTGTAATGTGCGAGCTG</td>
</tr>
<tr>
<td>Cpt-1β*</td>
<td>CCCATGTGCTCCTACCAGAT</td>
<td>CCTTGAAGAAGCGACCTTTG</td>
</tr>
<tr>
<td>FAT/CD36</td>
<td>CAACTGGTGAGATGTTCCT</td>
<td>GCAGAATCAAAGGAGAGCAGAC</td>
</tr>
<tr>
<td>MCAD*</td>
<td>TCGGAGGCTATGGATTCAAC</td>
<td>TCAATGTGCTACAGAGCTATG</td>
</tr>
<tr>
<td>PGC-1α</td>
<td>ATGTGTCGCTTCTTGTCTCT</td>
<td>ATCTACTGCGCTGGACCTT</td>
</tr>
<tr>
<td>PPARα*</td>
<td>TCATACTGACATGGAGACCTTG</td>
<td>ACTGGCAGCAGTGAAGAATC</td>
</tr>
<tr>
<td>PPARβ/δ</td>
<td>GGTAGAAGCCATCCAGGACA</td>
<td>CCGTCTCTCTTAGCCACTGC</td>
</tr>
<tr>
<td>SIRTI†</td>
<td>GATACCTTGAGAGCAGTGTGC</td>
<td>CTCCACGAAACAGCTTCACAA</td>
</tr>
<tr>
<td>TBP*</td>
<td>GGCTCTCAGAAGCATACATA</td>
<td>GCCAAGCCTGAGCATAAA</td>
</tr>
<tr>
<td>18S</td>
<td>TGTGGTGTGGAGAAAGCAG</td>
<td>TCCCATCCTTCACATCCTTC</td>
</tr>
</tbody>
</table>

* From Templeman et al. (48)
† From Asher et al. (1)
Table 2. Physical and cardiac characteristics of control, HR\textsubscript{mini} and HR\textsubscript{normal} mice.

<table>
<thead>
<tr>
<th></th>
<th>Control mice</th>
<th>HR\textsubscript{mini} mice</th>
<th>HR\textsubscript{normal} mice</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Body mass (g)</strong></td>
<td>29.8 ± 0.6\textsuperscript{a} (20)</td>
<td>27.5 ± 0.6\textsuperscript{b} (18)</td>
<td>30.1 ± 0.5\textsuperscript{a} (20)</td>
</tr>
<tr>
<td><strong>Gastrocnemius mass (mg)</strong></td>
<td>123 ± 3\textsuperscript{a} (20)</td>
<td>64 ± 3\textsuperscript{b} (17)</td>
<td>132 ± 3\textsuperscript{a} (20)</td>
</tr>
<tr>
<td><strong>Left ventricle mass (mg)</strong></td>
<td>104.2 ± 2.9 (10)</td>
<td>110.9 ± 2.1 (10)</td>
<td>105.0 ± 3.8 (10)</td>
</tr>
<tr>
<td><strong>Right ventricle mass (mg)</strong></td>
<td>20.2 ± 0.8 (10)</td>
<td>22.5 ± 1.0 (10)</td>
<td>21.8 ± 1.3 (10)</td>
</tr>
<tr>
<td><strong>Ventricular mass (mg)</strong></td>
<td>124.4 ± 2.8 (10)</td>
<td>133.4 ± 2.7 (10)</td>
<td>126.8 ± 3.1 (10)</td>
</tr>
<tr>
<td><strong>Relative left ventricle mass (mg per g body mass)</strong></td>
<td>3.24 ± 0.07\textsuperscript{a} (10)</td>
<td>3.68 ± 0.06\textsuperscript{b} (10)</td>
<td>3.29 ± 0.06\textsuperscript{a} (10)</td>
</tr>
<tr>
<td><strong>Relative right ventricle mass (mg per g body mass)</strong></td>
<td>0.63 ± 0.02 (10)</td>
<td>0.75 ± 0.04 (10)</td>
<td>0.70 ± 0.06 (10)</td>
</tr>
<tr>
<td><strong>Relative ventricular mass (mg per g body mass)</strong></td>
<td>3.87 ± 0.06\textsuperscript{a} (10)</td>
<td>4.43 ± 0.09\textsuperscript{b} (10)</td>
<td>3.99 ± 0.08\textsuperscript{a} (10)</td>
</tr>
<tr>
<td><strong>Left ventricle DNA (mg per g wet mass)</strong></td>
<td>5.59 ± 0.42 (8)</td>
<td>5.65 ± 0.38 (8)</td>
<td>4.99 ± 0.43 (8)</td>
</tr>
<tr>
<td><strong>Total left ventricle DNA (mg per left ventricle)</strong></td>
<td>0.60 ± 0.05 (8)</td>
<td>0.62 ± 0.04 (8)</td>
<td>0.50 ± 0.06 (8)</td>
</tr>
</tbody>
</table>

Data are presented as simple means ± SEM, with the exception of gastrocnemius muscle mass (marked with an asterisk), which is presented as estimated marginal means ± SEM as adjusted for the covariate of body mass. Sample size is shown in parenthesis for each trait. The presence of letters denotes statistical significance, where values not sharing a common letter are significantly different from each other (p < 0.05).
Table 3. Mass-specific maximal aerobic capacity (VO$_{2\text{max}}$) and whole-body lipid and carbohydrate oxidation rates of control, HR$_{\text{mini}}$, and HR$_{\text{normal}}$ mice during low- and high-intensity submaximal exercise.

<table>
<thead>
<tr>
<th></th>
<th>Control mice</th>
<th>HR$_{\text{mini}}$ mice</th>
<th>HR$_{\text{normal}}$ mice</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mass-specific VO$_{2\text{max}}$</td>
<td>9.26 ± 0.12 $^a$ (10)</td>
<td>10.63 ± 0.12 $^b$ (10)</td>
<td>10.18 ± 0.13 $^c$ (9)</td>
</tr>
<tr>
<td>(mL O$_2$ hr$^{-1}$ g$^{-1}$)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>66% VO$_{2\text{max}}$</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Exercise intensity</td>
<td>66 ± 1 (10)</td>
<td>64 ± 1 (10)</td>
<td>69 ± 2 (9)</td>
</tr>
<tr>
<td>(% VO$_{2\text{max}}$)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>VO$_2$</td>
<td>6.14 ± 0.09 $^a$ (10)</td>
<td>7.04 ± 0.09 $^b$ (10)</td>
<td>6.84 ± 0.10 $^b$ (9)</td>
</tr>
<tr>
<td>(mL O$_2$ hr$^{-1}$ g$^{-1}$)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>VCO$_2$</td>
<td>4.93 ± 0.08 $^a$ (10)</td>
<td>5.54 ± 0.09 $^b$ (10)</td>
<td>5.37 ± 0.09 $^b$ (9)</td>
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<tr>
<td>(mL CO$_2$ hr$^{-1}$ g$^{-1}$)</td>
<td></td>
<td></td>
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<tr>
<td>Lipid oxidation</td>
<td>4.12 $^a$ ± 0.23 (10)</td>
<td>5.09 $^b$ ± 0.24 (10)</td>
<td>4.97 $^b*$ ± 0.25 (9)</td>
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<tr>
<td>(mL O$_2$ hr$^{-1}$ g$^{-1}$)</td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>Carbohydrate oxidation</td>
<td>2.02 ± 0.20 (10)</td>
<td>1.95 ± 0.21 (10)</td>
<td>1.85 ± 0.23 (9)</td>
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<tr>
<td>(mL O$_2$ hr$^{-1}$ g$^{-1}$)</td>
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<td></td>
</tr>
<tr>
<td>78% VO$_{2\text{max}}$</td>
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<tr>
<td>Exercise intensity</td>
<td>78 ± 1 (10)</td>
<td>76 ± 1 (9)</td>
<td>79 ± 1 (9)</td>
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<tr>
<td>(% VO$_{2\text{max}}$)</td>
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<tr>
<td>VO$_2$</td>
<td>7.22 ± 0.09 $^a$ (10)</td>
<td>8.21 ± 0.11 $^b$ (9)</td>
<td>8.00 ± 0.10 $^b$ (9)</td>
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<tr>
<td>(mL O$_2$ hr$^{-1}$ g$^{-1}$)</td>
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<tr>
<td>VCO$_2$</td>
<td>5.93 ± 0.12 $^a$ (10)</td>
<td>6.88 ± 0.14 $^b$ (9)</td>
<td>6.73 ± 0.14 $^b$ (9)</td>
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<td>(mL CO$_2$ hr$^{-1}$ g$^{-1}$)</td>
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<tr>
<td>Lipid oxidation</td>
<td>4.39 ± 0.33 (10)</td>
<td>4.53 ± 0.37 (9)</td>
<td>4.32 ± 0.37 (9)</td>
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<tr>
<td>(mL O$_2$ hr$^{-1}$ g$^{-1}$)</td>
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<tr>
<td>Carbohydrate oxidation</td>
<td>2.83 ± 0.32 (10)</td>
<td>3.68 ± 0.37 (9)</td>
<td>3.67 ± 0.36 (9)</td>
</tr>
<tr>
<td>(mL O$_2$ hr$^{-1}$ g$^{-1}$)</td>
<td></td>
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</table>

Mass-specific VO$_{2\text{max}}$ is presented as estimated marginal means ± SEM, as adjusted for the cofactor of group batch. Exercise intensities (% VO$_{2\text{max}}$) are presented as simple means ± SEM. All mass-specific oxidation rates are presented as estimated marginal means ± SEM, as adjusted for the covariate of individual % VO$_{2\text{max}}$. The presence of letters denotes statistical significance, where values not sharing a common letter are significantly different from each other (p < 0.05, except an asterisk indicates p = 0.057).
Table 4. mRNA expression in cardiac muscle of control, HR\textsubscript{mini}, and HR\textsubscript{normal} mice.

<table>
<thead>
<tr>
<th>mRNA</th>
<th>Control mice</th>
<th>HR\textsubscript{mini} mice</th>
<th>HR\textsubscript{normal} mice</th>
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<tr>
<td><strong>PPAR\textalpha</strong> relative mRNA</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Left Ventricle</td>
<td>1.00 ± 0.18 (7)</td>
<td>0.78 ± 0.03 (7)</td>
<td>1.05 ± 0.14 (8)</td>
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<tr>
<td>Right Ventricle</td>
<td>1.00 ± 0.09 (6)</td>
<td>0.88 ± 0.10 (7)</td>
<td>1.28 ± 0.20 (8)</td>
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<td><strong>MCAD</strong> relative mRNA</td>
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<td></td>
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<tr>
<td>Left Ventricle</td>
<td>1.00 ± 0.13 (7)</td>
<td>0.99 ± 0.11 (7)</td>
<td>0.94 ± 0.13 (8)</td>
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<tr>
<td>Right Ventricle</td>
<td>1.00 ± 0.14 (7)</td>
<td>0.94 ± 0.14 (8)</td>
<td>1.31 ± 0.23 (8)</td>
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<td><strong>CPT-1\textalpha</strong> relative mRNA</td>
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<tr>
<td>Left Ventricle</td>
<td>1.00 ± 0.22 (7)</td>
<td>0.74 ± 0.10 (7)</td>
<td>0.79 ± 0.10 (8)</td>
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<tr>
<td>Right Ventricle</td>
<td>1.00 ± 0.17 (6)</td>
<td>0.92 ± 0.18 (7)</td>
<td>1.42 ± 0.32 (8)</td>
</tr>
<tr>
<td><strong>CPT-1\textbeta</strong> relative mRNA</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Left Ventricle</td>
<td>1.00 ± 0.17 (7)</td>
<td>0.86 ± 0.06 (7)</td>
<td>0.82 ± 0.06 (8)</td>
</tr>
<tr>
<td>Right Ventricle</td>
<td>1.00 ± 0.09 (6)</td>
<td>0.88 ± 0.10 (7)</td>
<td>1.28 ± 0.20 (8)</td>
</tr>
</tbody>
</table>

Data are presented as simple means ± SEM, with sample size shown in parenthesis; mRNA expression levels are corrected against TBP mRNA, and values are reported normalized to the mean control mouse value (arbitrarily set at 1.00). Statistical analyses were performed on raw mRNA levels corrected against TBP mRNA; no differences between groups were statistically significant.
A.

FAT/CD36 mRNA (relative levels)

Control | HR\textsubscript{mini} | HR\textsubscript{normal}

B.

FAT/CD36 protein (arbitrary units)

Control | HR\textsubscript{mini} | HR\textsubscript{normal}