

Experimental Communication

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Author contributions

CD, TK, CC, LHD., A-CC, AM, DT, NZ, BK, EG, and NP, designed the research. CD, TK, CC, LHDC, NZ, BK, and NP, performed the experiments. CD, TK, CC, LHDC, DT, NZ, BK, EG, and NP, analyzed the data. CD and BK wrote the paper with input from all authors.

Conflicts of interest

EG is the founder and CEO of Oroboros Instruments.

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Data availability

Materials and data generated from this study are available upon request to the corresponding author. The article includes all datasets generated and analyzed in this study. RNA-seq data have been deposited to GEO under accession number GSE225927.

Keywords

respirometry; calcium Ca²⁺ mitochondrial membrane potential $\Delta \Psi_{mt}$; skeletal muscle; isolated mitochondria imt; exercise; redox state; coenzyme Q.

Functional hypoxia reduces mitochondrial calcium uptake

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Summary

Mitochondrial respiration extends beyond ATP generation, with the organelle participating in many cellular and physiological processes. Parallel changes in components of the mitochondrial electron transfer system with respiration render it an appropriate hub for coordinating cellular adaption to changes in oxygen levels. How changes in respiration under functional hypoxia (i.e., when intracellular O₂ levels limit mitochondrial respiration) are relayed by the electron transfer system to impact mitochondrial adaption and remodeling after hypoxic exposure remains poorly defined. This is largely due to challenges integrating findings under controlled and defined O₂ levels in studies connecting functions of isolated mitochondria to humans during physical exercise. Here we present experiments under conditions of hypoxia in isolated mitochondria, myotubes and exercising humans. Performing steadystate respirometry with isolated mitochondria we found that oxygen limitation of respiration reduced electron flow and oxidative phosphorylation, lowered



the mitochondrial membrane potential difference, mitochondrial decreased calcium influx. and Similarly, in myotubes under functional hypoxia mitochondrial calcium uptake decreased in response sarcoplasmic reticulum calcium release for to contraction. In both myotubes and human skeletal muscle this blunted mitochondrial adaptive responses and remodeling upon contractions. Our results suggest that by regulating calcium uptake the mitochondrial electron transfer system is a hub for coordinating cellular adaption under functional hvpoxia.

1. Introduction

Mitochondria use molecular oxygen (O_2) as the terminal electron acceptor to generate the protonmotive force for phosphorylation of ADP [1]. O_2 consumption is adjusted in response to changes in cellular ATP demand [2]. This is exemplified during intense muscle contractions which can increase O_2 consumption up to 20-fold of that at rest [3]. Mitochondrial ATP regeneration by oxidative phosphorylation depends on the supply of O_2 and ADP [4]. The state of 'functional hypoxia' describes when O_2 levels limit mitochondrial respiration [5, 6]. Although restricted to a very low intracellular pO_2 , functional hypoxia can occur during physiological activity [7-10]), under conditions of ambient hypoxia [11], or during ischemia [12].

Although the O₂ kinetics (i.e., dependence on O₂ partial pressure) of oxidative phosphorylation have been resolved [13, 14], there is considerable interest in understanding if and how O₂ controls other components and functions of the electron transfer system [15-17]. This interest has been stimulated by findings showing hypoxia and impairments in mitochondrial respiration are hallmarks of many metabolic diseases [18, 19] and that mitochondrial signals influence numerous cellular processes [15-17, 20]. Signals of particular interest are mitochondrial redox states [17, 21] and the mitochondrial membrane potential difference (mt-membrane potential difference or $\Delta \Psi_{mt}$; [12]) which exert influence on processes involved in cellular adaptions and physiological outcomes [15]. These processes include alterations in metabolite concentrations [22], mitochondrial reactive species production [23], the import of proteins into mitochondria [24], and the shaping of cellular Ca²⁺ signals [25].

Understanding the mechanisms through which hypoxia regulates biological outcomes has proved challenging, largely, due to the difficulty to integrate findings under well-controlled and defined O_2 levels in models varying from isolated mitochondria to intact humans [5]. Nonetheless, the potential for components of the electron transfer system to play key roles in relaying signals for O_2 sensing are highlighted by the rapid regulation of the system under functional hypoxia [21]. For example, at the onset of functional hypoxia (at only 5 % limitation of respiration by O_2) there is a steep decline in cytochrome *c* oxidation state [21]. These parallel changes in components of the electron transfer system with respiration render it an appropriate hub for coordinating cellular adaption. How changes in respiration under hypoxia are relayed by the electron transfer system to elicit mitochondrial adaption after hypoxic exposure remains poorly defined.



We hypothesized that changes in the mitochondrial electron transfer system could influence key processes that control cellular adaption under hypoxia.

Here we report experiments under conditions of functional hypoxia in carefully chosen and relevant models: isolated mitochondria, myotubes and muscle from exercising humans, and propose that O_2 control of the mitochondrial electron transfer system function reduces muscle mitochondrial Ca^{2+} uptake and regulates mitochondrial adaptive remodeling upon physical exercise.

2. Results

2.1. Functional hypoxia reduces electron flow through the Q-junction and lowers $\Delta\Psi_{mt}$

In the first set of experiments, we investigated O₂ control of the mitochondrial electron transfer system (Figure 1A) in isolated mitochondria. The "apparent K_m " or p_{50} of mitochondrial respiration ranges from 0.01-0.10 kPa (~0.1 to 1 µM O₂) [14]. Therefore, to study O₂ control of other mitochondrial parameters required high resolution in the low µM range for O₂ and sufficient time for resolution of the additional parameters being measured simultaneously [21]. To overcome this challenge, we used a steady-state high-resolution respirometry approach [21]. This allowed us to set mitochondrial respiration at specific O₂-limited fractions of maximal O₂ flux (J_{max} ; Figure 1B) through the titration of a hydrogen peroxide (H₂O₂) solution in the presence of excess catalase. Maintaining a constant injection flow over a period (typically 3-6 min) allowed us to measure key mitochondrial parameters and functions of interest under steady states of hypoxia (Figure 1B). Importantly, there is agreement in the O₂ kinetics observed in aerobic-anaerobic transitions and steady-states [21] making this approach the gold-standard to study O₂ control of mitochondrial electron transfer system function.

Providing substrates to support convergent electron entry into the Q-junction from oxidation pathways linked to NADH (via Complex I CI), succinate (via CII) and fatty-acid (via electron transferring flavoprotein) in the presence of saturating ADP concentrations (Figure 1A) we first measured the redox state of the electron-transfer-reactive coenzyme Q (Q). Using the Q-mimetic coenzyme Q₂ and a three-electrode system [26], at steady-state levels of hypoxia (0.1 to 0.5 J_{max}), Q was more reduced compared with kinetically saturating O₂ levels in mitochondria isolated from mouse heart (Figure 1B, C, S1B) and brain (Figure S1D, E).

To assess the consequence of oxygen-limited respiration on $\Delta \Psi_{mt}$ we applied steadystate respirometry using tetramethylrhodamine methyl ester (TMRM) in dequench mode (Figure 1D). Tracking the mt-membrane potential difference across steady states of hypoxia up to kinetically saturating oxygen concentrations revealed that oxygen limitation leads to a reversible depolarization of $\Delta \Psi_{mt}$ in mitochondria isolated from mouse heart (Figure 1D, E). In summary, oxygen limitation of respiration reduces electron flow and oxidative phosphorylation, and lowers the mitochondrial membrane potential difference.



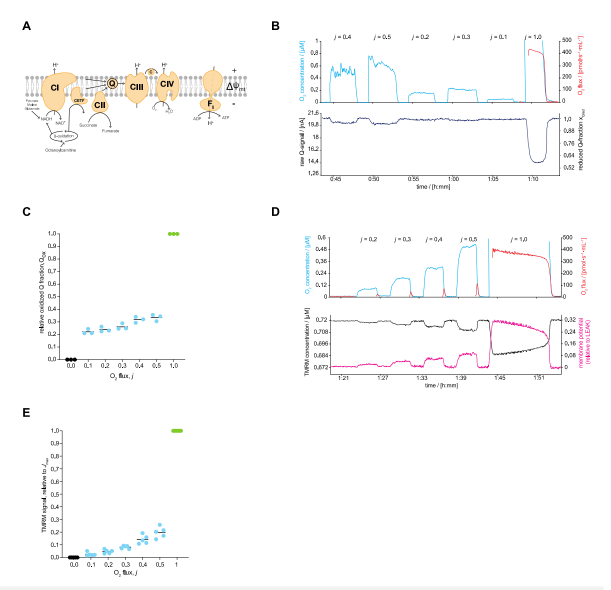


Figure 1. Functional hypoxia reduces the redox state of electron-transfer-reactive Q and lowers the mitochondrial membrane potential difference. (A) Schematic representation of mitochondrial electron transfer system and coupling to oxidative phosphorylation (F₁F₀-ATPase). **(B)** Measurement of the redox state of the mitochondrial electron-transfer-reactive Q-pool (Q) under steady-states of hypoxia and at maximal O₂ flux in isolated mitochondria from mouse cardiac muscle respiring on NADH-linked substrates, succinate, and fatty-acid in the presence of kinetically saturating ADP. (C) Relative Q oxidation under anoxia (i = 0), functional hypoxia (i = 0.1 to 0.5) and at maximal O_2 flux (*j* = 1) in mitochondria isolated from mouse cardiac muscle (*N*=3). *j* is the ratio of the set I_{0_2} to the measured OXPHOS capacity (I_{max}). (D) Mitochondrial membrane potential difference under steady states of hypoxia and at maximal O₂ flux in mitochondria isolated from mouse cardiac muscle respiring on NADH-linked substrates, succinate, and fatty acid in the presence of kinetically saturating ADP. (E) Relative TMRM signal under anoxia (j = 0), functional hypoxia (j = 0.1 to 0.5) and at maximal O_2 flux (j = 1) in mitochondria isolated from mouse cardiac muscle (N=5). *j* is the ratio of the set I_{02} to the measured OXPHOS capacity (J_{max}). Lines represent the means, dots are individual values.

2.2. Functional hypoxia reduces mitochondrial Ca²⁺ uptake



One important function of $\Delta \Psi_{mt}$ is to provide the driving force for Ca²⁺ uptake [27]. Mitochondrial Ca²⁺ influx occurs through the voltage-dependent anion channel (VDAC) on the outer mitochondrial membrane and through the highly selective mitochondrial calcium uniporter (MCU) across the inner mitochondrial membrane [25]. MCU function relies almost completely on two parameters: [Ca²⁺] in the area surrounding the channel and $\Delta \Psi_{mt}$ [25]. Therefore, we hypothesized that functional hypoxia would reduce mitochondrial Ca²⁺ uptake due to its effect on $\Delta \Psi_{mt}$.

We investigated mitochondrial Ca²⁺ uptake under functional hypoxia in mitochondria isolated from mouse heart using calcium green [28]. Given that Ca²⁺ can alter the activity of the mitochondrial electron transfer system [29, 30] we first determined the conditions under which O₂ flux was not affected by Ca²⁺. This was a necessary step as our aim was to compare mitochondrial Ca²⁺ uptake at maximal flux (*J*_{max}) with a specific reduced fraction of *J*_{max}. Any changes in flux due to Ca²⁺ during the measurement period would complicate the interpretation of our results. We therefore performed a sequential stepwise titration of CaCl₂ (10 µM) whilst monitoring mitochondrial Ca²⁺ uptake simultaneously with O₂ consumption (Figure S2A). Mitochondrial Ca²⁺ uptake was stimulated starting at 10 µM CaCl₂, however, at higher concentrations (starting at 30 µM) mitochondrial respiration was inhibited (Figure S2A). We therefore tested if a single 20 µM titration of CaCl₂ at saturating O₂ concentrations (at *J*_{max}) affected respiration. These experiments showed mitochondrial O₂ flux to be stable for >6 min (Figure S2B, C).

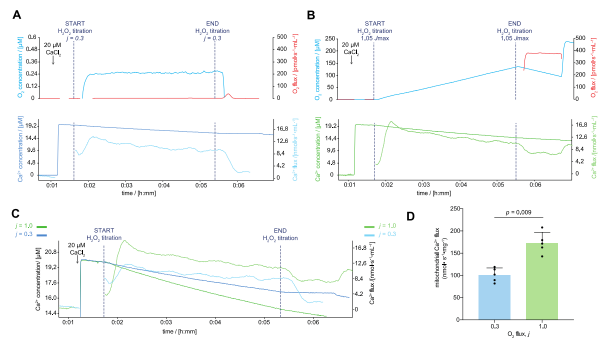


Figure 2. Functional hypoxia reduced mitochondrial Ca²⁺ flux in mitochondria isolated from mouse cardiac muscle. (A) Mitochondrial Ca²⁺ flux under steady-state hypoxia. **(B)** Mitochondrial Ca²⁺ flux at maximal O₂ flux. **(C)** Representative traces of extramitochondrial Ca²⁺ concentration and mitochondrial Ca²⁺ flux at maximal O₂ flux (J_{max} , green line) and under functional hypoxia (j = 0.3, blue line; superimposed from panels A and B). **(D)** Mitochondrial Ca²⁺ flux under functional hypoxia (j = 0.3) and at maximal O₂ flux (j = 1.0; N=5). Bars represent the means, error bars the standard deviations, dots are individual values.



Therefore, in the presence of 20 μ M CaCl₂ we controlled mitochondrial respiration at 0.3 J_{max} (j = 0.3; Figure 2A) and at J_{max} (j = 1.0; Figure 2B) for 4 min, observing that mitochondrial Ca²⁺ flux was lower when respiration was limited by O₂ levels, i.e. in functional hypoxia (Figure 2C, D). These data are in keeping with the reduced $\Delta \Psi_{mt}$ as the driving force for Ca²⁺ uptake under hypoxia (Figure 1C, D). Taken together, our findings indicate that oxygen limitation of oxidative phosphorylation lowered $\Delta \Psi_{mt}$ and in turn reduced mitochondrial Ca²⁺ flux.

2.3. Myotubes reduce mitochondrial Ca²⁺ uptake under functional hypoxia

The ability of mitochondrial Ca^{2+} uptake to shape cellular events varies by cell type [25]. One cellular process where this plays a key role is muscle contraction [31, 32]. In muscle tissue, mitochondrial Ca^{2+} influences not only energy metabolism during contraction [29] but also signalling for adaptive processes such as remodeling of mitochondrial metabolism [33]. Because of this important dual role of mitochondrial Ca^{2+} in muscle physiology, we chose to explore hypoxia further in skeletal muscle tissue.

Having found that functional hypoxia controls Ca^{2+} flux in mitochondria isolated from mouse heart, we next considered whether hypoxia also exerts control over mitochondrial Ca^{2+} uptake in living cells. To explore this hypothesis, we first performed experiments to assess the culture conditions required to model functional hypoxia in myotubes (Figure 3A). In most published papers using cellular models, ambient air humidified to 90 % and enriched with 5 % carbon dioxide is used, which represents an oxygen pressure ($p_{02} \sim 20$ kPa at sea-level, ~ 18.5 % O₂), that is higher than that experienced by any tissue *in situ* [34]. There is a growing appreciation that the oxygen levels used in cell culture (with primary cells and cell lines) profoundly affect the observed biology reproducibility and *in vivo* relevance [34]. Given the importance of culturing cells under physiological O₂ levels to improve *in vivo* relevance we chose to use 5 % O₂ to mimic normoxic *in vivo* skeletal muscle O₂ pressure (\sim 4-8 % O₂ at an atmospheric pressure ~100 kPa [11]). Evidence from human skeletal muscle suggests that under ambient hypoxia (fraction of inspired O₂ = 10 %) *in vivo*, resting skeletal muscle O₂ pressures are ~1-3 % O₂ [11].

We exposed differentiated myotubes cultured under 5 % O₂ acutely to 1 % O₂ in the gas phase and used an optical probe to measure the partial pressure of oxygen (p_{02}) in the media. A 2 h exposure was necessary to reduce media p_{02} from normoxia to physiological values known to occur in acute ambient hypoxia in humans ([8]; Figure 3B). At this timepoint we assessed if O₂ in the intracellular environment was also decreased by measuring the protein content of the transcription factor HIF1 α (which increases exponentially from 5 % to 1 % O₂ see ref. [35]) by western blot and performing bulk RNA-sequencing quantify the expression of gene-sets involved in the cellular response to hypoxia. HIF1 α protein (Figure 3C-D) and the "Hypoxia" and "Glycolysis" hallmark gene-sets (Figure 3E, Dataset S1) were among those enriched in myotubes exposed to 1 % compared with 5 % O₂.



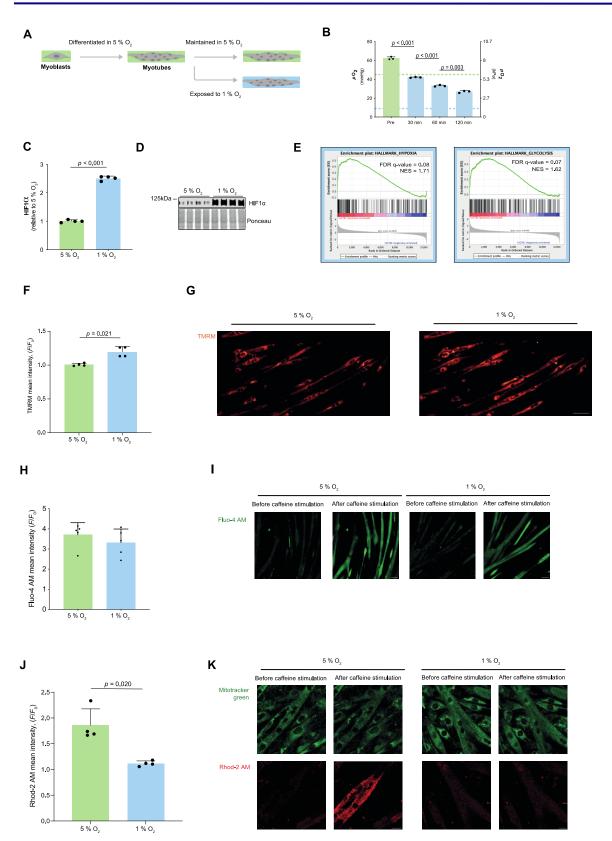


Figure 3. Mitochondrial Ca²⁺ uptake was lower under functional hypoxia in mouse myotubes. (A) Culture conditions. **(B)** Partial pressure of O₂ in culture media upon exposure to 1 % O₂ in the gas phase (N=3). **(C)** HIF1 α protein content in myotubes 120 min after exposure to 1 % O₂ in the gas phase (N=4). **(D)** Representative blot of HIF1 α



with ponceau (loading) stain in myotubes 120 min after exposure to 1 % O₂ in the gas phase (*N*=4). **(E)** Hypoxia and glycolysis hallmark gene sets enriched in myotubes 120 min after exposure to 1 % O₂ compared with 5 % O₂ in the gas phase (*N*=6). False discovery rate FDR. Normalized enrichment score NES. **(F)** TMRM mean intensity under 5 % and 1 % O₂ in the gas phase (*N*=4) in C2C12 myotubes. **(G)** Representative images of C2C12 myotubes stained with TMRM exposed to 5 % or 1 % O₂. Scale bar is 50 µm. **(H)** Quantification of Fluo4AM mean intensity in response to caffeine stimulation in myotubes 120 min after exposure to 5 % or 1 % O₂ in the gas phase (*N*=5). **(I)** Representative images of C2C12 myotubes stained with Fluo4AM, exposed to 5 % or 1 % O₂ and stimulated with caffeine. Scale bar is 50 µm. **(J)** Response of Rhod-2AM fluorescence to caffeine stimulation in C2C12 myotubes under 5 % and 1 % O₂ (*N*=4). **(K)** Representative images of C2C12 myotubes stained with mitotracker green and Rhod-2AM, exposed to 5 % or 1 % O₂ and stimulated with caffeine. Scale bar is 20 µm. Bars represent the means, error bars the standard deviations, dots are individual values.

To test whether these conditions were functionally hypoxic we used confocal microscopy to perform live imaging on myotubes under 5 % and 1 % O₂ to measure the $\Delta \Psi_{mt}$. Functional hypoxia decreased the $\Delta \Psi_{mt}$ (see data from isolated mitochondria in Figures 1C, D). Indeed a 2 h exposure to 1 % increased the TMRM mean intensity (quench mode) indicating a decreased $\Delta \Psi_{mt}$ (Figure 3F, G). Therefore, we used these culture conditions to assess the effect of functional hypoxia on Ca²⁺ dynamics in myotubes.

First, we examined if hypoxia affected Ca²⁺ release and myotube contraction. Using the cytosolic Ca²⁺ indicator Fluo-4AM and mitochondrial Ca²⁺ indicator Rhod-2AM we stimulated myotubes with 2.5 mM caffeine in the absence of extracellular Ca²⁺. There were no differences in cytosolic Ca²⁺ in response to caffeine stimulation under 5 % or 1 % O₂ (Figure 3H, I) and myotubes contracted in response to electrical stimulation (data not shown). However, mitochondrial Ca²⁺ uptake was lower in response to caffeine stimulation under 1 % compared with 5 % O₂ (Figure 3J, K). These data suggest that in living cells mitochondrial Ca²⁺ uptake is reduced under hypoxia due to membrane depolarization.

2.4. Functional hypoxia alters mitochondrial adaptive responses to contractions in mouse myotubes and human skeletal muscle

Our data support a model whereby a reduction in $\Delta \Psi_{mt}$ under hypoxia decreases muscle mitochondrial Ca²⁺ uptake in response to Ca²⁺ release from the sarcoplasmic reticulum. This process is critical for the muscle response and metabolic remodelling upon muscle contraction including increased protein content of OXPHOS subunits and mitochondrial supercomplex formation [33]. Therefore, we decided to test whether these immediate changes under functional hypoxia could influence mitochondrial adaptions following a period of recovery in normoxia. To do so, we first performed experiments in electrically stimulated C2C12 myotubes (Figure 4A). This electrical stimulation protocol induces mitochondrial Ca²⁺ uptake and elicits mitochondrial adaptions under culture conditions with ~18.5 % O₂ [33].

Therefore, we first explored the response of mitochondria to six 30 s bouts of electrical stimulation at 50 Hz under 5 % O_2 (Figure 4A). Compared with non-stimulated controls, the electrical stimulation increased the transcripts of many gene sets that are hallmarks of the response to intense exercise including "Oxidative phosphorylation" (Figure 4B) and induced pyruvate dehydrogenase dephosphorylation (PDH; an indirect



readout of mitochondrial Ca²⁺ uptake; Figure 4C-D; [33]) immediately after stimulation. Moreover, electrical stimulation increased the protein content of OXPHOS subunits 48 hours later (Figure 4E-J) as reported for myotubes in ~18.5 % O₂ [33], and similar to the response of human skeletal muscle to sprint interval training in normoxia [33].

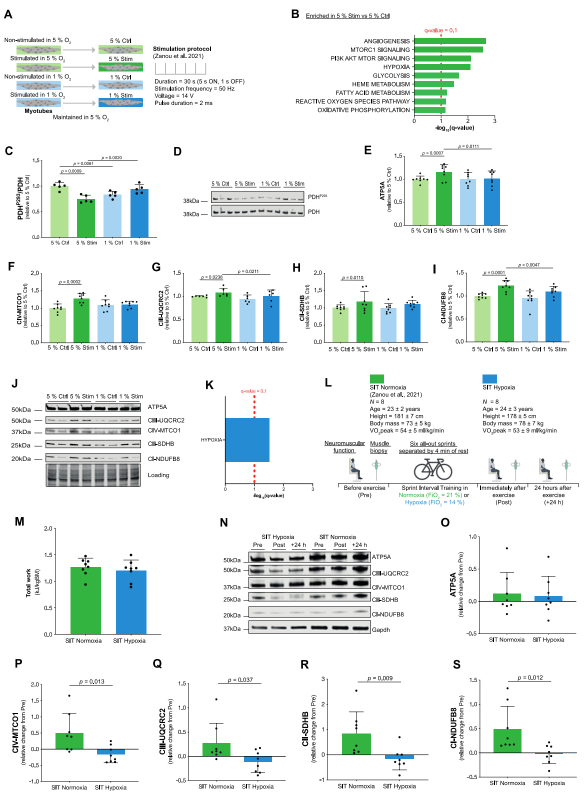


Figure 4. Functional hypoxia altered mitochondrial responses to contractions in mouse myotubes and human skeletal muscle. (A) Culture conditions and electrical



stimulation protocol. (B) Hallmark gene sets enriched in myotubes immediately after stimulation in 5 % O₂ compared with non-stimulated myotubes (*N*=6). (C) PDHP293 relative to total PDH protein content in myotubes 3 h after stimulation in 5 % or 1 % O₂ and their controls (*N*=5). (**D**) Representative blots of PDHP293 and PDH in myotubes 3 h after stimulation in 5 % or 1 % O₂ and their controls. (E-I) ATP5A (E), CIV-MTCO1 (F), CIII-UQCRC2 (G), CII-SDHB (H) and CI-NDUFB8 (I) protein content in C2C12 myotubes 48 h after stimulation (*N*=7). (J) Representative blot of ATP5A, CIII-UQCRC2, CIV-MTCO1, CII-SDHB and CI-NDUFB8 with loading total protein stain from C2C12 myotubes 48 h after stimulation. (K) Hallmark gene sets enriched in myotubes immediately after stimulation in 1 % O₂ compared with myotubes stimulated in 5 % O₂ (*N*=6). (L) Human exercise study protocol. (M) Total work performed (expressed relative to body mass) during the SIT session (N=8). (N) Representative blot of ATP5A, CIII- UQCRC2, CIV-MTCO1, CII-SDHB, CI-NDUFB8 and Gapdh (loading control) from human vastus lateralis biopsies. (O-S) Relative changes in ATP5A (O), CIV-MTCO1 (P), CIII- UQCRC2 (Q), CII-SDHB (R) and CI-NDUFB8 (S) protein content from Pre to +24 h in human vastus lateralis biopsies (*N*=8). Bars represent the means, error bars the standard deviations, dots are individual values.

Applying the same electrical stimulation protocol to myotubes in hypoxia led to an enrichment of transcripts in the "Hypoxia" gene set (Figure 4 K, Dataset S1) and altered lipid (as measured by mass-spectrometry see Methods; Figure S3A-B, Dataset S2) and glucose (assessed using a glucose uptake assay see Methods Figure S3C) metabolism compared with stimulation in 5 % O₂. Stimulation in 1 % O₂ blunted PDH dephosphorylation compared with stimulation in 5 % O₂, suggesting reduced mitochondrial Ca²⁺ accumulation during contractions under hypoxia. Additionally, we observed that hypoxia blunted increases in oxidative phosphorylation protein complexes after stimulation (Figure 4E-J) possibly a result of reduced mitochondrial Ca²⁺ uptake during contractions.

Finally, to examine if hypoxia influenced adaptions to muscle contractions in humans, we collected data from a single session of sprint interval training (SIT) in hypoxia in eight healthy young male participants and compared these results with published data under normoxia (fraction of inspired O_2 $F_iO_2 = 20.9$ %; [33]). During maximal exercise mitochondrial respiration is likely limited up to ~5 % [7], therefore we chose to use a fraction of inspired O_2 of 14 % to model functional hypoxia as this has been shown to limit both whole-body- and single leg-oxygen consumption during exercise [9, 36-39]. The SIT hypoxia group underwent the same experimental protocol as the SIT normoxia group (Figure 4L; [33]) with the exception that the F_iO_2 was reduced for the exercise session (5-min warm-up and six all-out 30 s sprints separated by 4-min of rest (Figure 4L; [33]). The protocol included neuromuscular function assessments and *vastus lateralis* muscle biopsy collection before, immediately after, and 24 hours after the exercise session (Figure 4L; [33]).

Performing the exercise in normobaric hypoxia reduced blood oxygen saturation (Figure S4A-E) but did not affect exercise performance (Figure S4F, G), work performed during the session (Figure 4M), or the extent and etiology of knee extensor neuromuscular fatigue after the session (Figure S4H-O, [33]). However, hypoxia blunted the previously reported [33] increases in respiratory Complexes CI, CII, CIII and CIV after exercise (Figure 4N-S, [33]). These data suggest that the role of hypoxia on the adaptions of mitochondria to exercise training may be linked to reduced mitochondrial Ca²⁺ uptake.



3. Discussion

Mitochondrial respiration extends beyond ATP generation – with the organelle participating in many cellular and physiological processes [15]. How changes in respiration under functional hypoxia are relayed by the electron transfer system to elicit mitochondrial adaption to hypoxic exposure remain poorly defined. Studies with isolated mitochondria describing parallel changes in components of the mitochondrial electron transfer system with respiration [21] render it an appropriate hub for coordinating cellular adaption to changes in oxygen levels. Advancements in understanding the mechanisms through which hypoxia regulates biological outcomes has proved challenging, largely, due to the difficulty to integrate studies of isolated mitochondria under well-controlled and defined O_2 levels to hypoxia in intact humans. Here we performed experiments under conditions of functional hypoxia (i.e., when O₂ levels limit mitochondrial respiration; [5]) in carefully chosen and relevant models: isolated mitochondria, myotubes and exercising humans. Our findings in isolated mitochondria and myotubes suggest that O₂ control of mitochondrial oxidative phosphorylation can relay changes in respiration and reduce mitochondrial Ca²⁺ uptake. We propose that alteration of this mitochondrial Ca²⁺ signal may influence how muscle mitochondria adapt after contractions in conditions of hypoxia.

Only one study [40] has previously investigated mitochondrial Ca²⁺ uptake under low-oxygen levels. In rat cardiomyocytes electrically stimulated under *anoxia* (*p*₀₂ limit of detection <0.003 kPa; [41]) mitochondrial Ca²⁺ was not increased when the cells entered a state of rigor after 30-40 min [40]. We did not specifically investigate mitochondrial Ca²⁺ flux under anoxia, but, we did not detect Ca²⁺ uptake after titration of CaCl₂ under anoxia (Figure 1C). Increasing O₂ levels above anoxia, we then detected mitochondrial Ca²⁺ flux (Figure 3A-C). This flux was reduced by ~35 % in isolated mitochondria when mitochondrial Ca²⁺ accumulation was also reduced under hypoxia in response to sarcoplasmic reticulum Ca²⁺ release (Figure 3I). The observed reduction in mitochondrial Ca²⁺ flux of ~35 % in response to 20 μ M CaCl₂ in isolated mitochondria would equate to a reduction in $\Delta \Psi_{mt}$ from 160 to ~110 mV (see Figure 1c in [27]).

Although we did not use a method that allowed us to calculate $\Delta \Psi_{mt}$ in terms of absolute transmembrane potential differences ([mV]; e.g., [42]), we did obtain information on $\Delta \Psi_{mt}$ under steady states of hypoxia in isolated mitochondria (Figure 1D, E) and myotubes (Figure 3F, G). We found that under hypoxia there was a depolarization of the $\Delta \Psi_{mt}$ (Figure 1E, 3G). The relative changes in TMRM signal under hypoxia compared with kinetically saturating oxygen levels in mitochondria isolated from mouse heart and 5 % O₂ in skeletal muscle myotubes were 5- and 1.25-fold, respectively. Although we cannot provide an explanation for this difference it is likely not due to an altered plasma membrane potential difference which is unaffected even after 30 min of *anoxia* (<1 nM O₂ combined with lactate efflux inhibition) in rat cardiomyocytes [43]. The depolarization of $\Delta \Psi_{mt}$ (relative change in TMRM signal) in myotubes after 2 h of exposure to 1 % O₂ (~10 mmHg) in the gas phase is less than that reported after 40 minutes of *ischemia* (measured $p_{02} < 2$ mmHg) in rat cardiomyocytes [12]. Taken together, these results support the notion that hypoxia reduces mitochondrial Ca²⁺ uptake via $\Delta \Psi_{mt}$.

The change in $\Delta \Psi_{mt}$ under functional hypoxia in isolated mitochondria occurred with a measured ~4-fold increase in reduction of the Q-pool compared with functional



normoxia (Figure 1C). This result is consistent with a pronounced reduction of the cytochromes *aa*₃, *b* and *c* under steady-state hypoxia [21], and an increased NAD(P)H reduction after 40 min of ischemia in Chouchani et al. [12].

On the other hand, the ~4-fold greater reduction of Q under functional hypoxia compared with functional normoxia contrasts with previous reports. Using mass-spectrometry to measure coenzyme Q reduction from mouse hearts after 30-min *ischaemia* there were no changes in coenzyme Q redox state [44] whereas there was a ~1.5- to 2-fold increase in reduction of Q in cardiomyocytes [43]. Discrepancies most likely arise from the different methods (electrochemical Q-sensor versus Q-extraction method) and preparations (isolated mitochondria or whole tissue/cells). Whereas these fold changes in isolated mitochondria and cells do not necessarily align, there does seem to be a current consensus in the literature cited above that under functional hypoxia the electron transfer system is more reduced. Numerous reports have unraveled the role of metabolites and reactive oxygen species in relaying this information to influence cellular adaption. Here, we suggest that Ca²⁺ may be the relay for altered mitochondrial adaption to muscle contraction under hypoxia.

We observed that hypoxia blunted the increase in OXPHOS Complex protein content after muscle contractions (Figure 4E-I, O-S). Although we did not perform direct measurements of mitochondrial Ca²⁺ after contractions, PDH (an enzyme sensitive to dephosphorylation in response to increased mitochondrial Ca²⁺ [45, 46]), was less dephosphorylated in myotubes stimulated under 1 % compared with 5 % O₂ suggesting reduced mitochondrial Ca²⁺ accumulation during contractions under hypoxia. Further supporting this notion, the response of OXPHOS proteins observed in the present study is similar to what was observed in response to contractions in myotubes with reduced MCU protein content [33].

Logically it is tempting to suggest that targeting the MCU to increase mitochondrial Ca^{2+} uptake under hypoxia might recover the blunted response of OXPHOS proteins after a period of recovery. However, mitochondrial Ca^{2+} can stimulate or inhibit mitochondrial respiration depending on the concentration in the matrix [30], and Ca^{2+} may also influence the mt-membrane potential difference [47]. Stimulating respiration under hypoxia could drive the cells towards "deeper" hypoxia and increasing matrix Ca^{2+} concentration could potentially collapse $\Delta \Psi_{mt}$, halting oxidative phosphorylation [48]. Therefore, the logic of simply increasing mitochondrial Ca^{2+} uptake under hypoxia to improve the response of OXPHOS proteins after a single session of exercise may not be so straightforward, and other studies suggest this may not be necessary. For example, chronic exercise training studies comparing training in hypoxia with normoxia show that the changes in wholebody V_{02} and muscle mitochondrial respiration are similar after several weeks [49].

Although our data suggest a role for reduced mitochondrial Ca^{2+} uptake in the response of muscle mitochondria to contractions under hypoxia, other factors contribute. For example, hypoxia has been shown to impair muscle protein synthesis [50] and induce signalling via other pathways e.g., the HIF pathway [35] which was activated in myotubes (Figure 3C-E, 4K). It would seem logical that when exposed to hypoxia the system activates pathways to promote adaption of other metabolic pathways e.g., glycolysis (Figure 3E) at the expense of mitochondrial oxidative phosphorylation [51, 52]. Regarding extrapolation of our data in the context of adaptation to hypoxia, despite paying careful attention to O_2 levels and using the best available methodologies, there are numerous other changes under hypoxia (e.g., pH, CO₂) that we did not control. We hope



that illustrating how oxygen uses the mitochondrial electron transfer system as a hub for coordinating cellular adaption will stimulate more research on the role of mitochondria in human adaptation to hypoxia including high-altitude and hypoxia-related diseases (e.g., chronic obstructive pulmonary disease, sleep apnea [53]).

4. Conclusion

We propose that under functional hypoxia with limitation of electron flow through the Q-junction (bottleneck effect), there is a progressive reduction of the Q-pool and a concomitant depolarization of $\Delta \Psi_{\rm mt}$. In turn this leads to a reduction in mitochondrial Ca²⁺ uptake during muscle contractions through which hypoxia then influences mitochondrial adaptions to exercise.

5. Materials and Methods

Fable 1: Key resource table REAGENT or RESOURCE		SOURCE	IDENTIFIER
Antibodies			
Gapdh anti-mouse		Abcam	Cat#8245
HIF1a anti-rabbit		Cayman Chemicals	Cat#10006421
Total OXPHOS anti-mouse		Abcam	Cat#110413
PDH anti-rabbit		Abcam	Cat#110334
PDH phospho S293 anti-mouse		Abcam	Cat#92696
Donkey anti-rabbit	IgG	LI-COR	Cat#926-32213
polyclonal antibody Donkey anti-mouse polyclonal antibody	IgG	LI-COR	Cat#926-32212
Donkey anti-mouse polyclonal antibody	IgG	LI-COR	Cat#926-68022
Donkey anti-rabbit polyclonal antibody	IgG	LI-COR	Cat#926-68023
Chemicals, Peptides and Re	ecom	binant Proteins	
DMEM		Thermo Fisher Scientific	Cat#41966-052
Fetal bovine serum		Thermo Fisher Scientific	Cat#1008247
Penicillin streptomycin		Thermo Fisher Scientific	Cat#15140122
Non-essential amino acids		Thermo Fisher Scientific	Cat#11140035
Horse serum		Thermo Fisher Scientific	Cat#26050088
Trypsin-EDTA (0.05 %) pho red	enol	Thermo Fisher Scientific	Cat#25300054
Trypsin		Sigma-Aldrich	Cat#T0303
Dimethyl sulfoxide		Sigma-Aldrich	Cat#8418



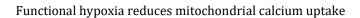
Paraformaldehyde	Thermo Fisher Scientific	Cat#J19943. K2
Fluo-4AM	Thermo Fisher Scientific	Cat#F142010
Mitotracker Green	Thermo Fisher Scientific	Cat#M7514
Mitotracker Red CMXRos	Thermo Fisher Scientific	Cat#M7512
Rhod-2AM	Thermo Fisher Scientific	Cat#R1245MP
TMRM	Thermo Fisher Scientific	Cat#T668
MiR05-Kit	Oroboros Instruments	Cat#60101-01
Dithionite	Sigma-Aldrich	Cat#71669
Catalase	Sigma-Aldrich	Cat#9322
Hydrogen peroxide H ₂ O ₂	Life Technologies	Cat#A36006
Calcium chloride CaCl ₂	Sigma-Aldrich	Cat#21115
Pyruvate	Sigma-Aldrich	Cat#P2256
Malate	Sigma-Aldrich	Cat#M1000
Glutamate	Sigma-Aldrich	Cat#G1626
Succinate	Sigma-Aldrich	Cat#2378
Cytochrome <i>c</i>	Sigma-Aldrich	Cat#C7752
Adenosine diphosphate	Merck	Cat#117105-1GM
Carbonyl cyanide 4- (trifluoromethoxy)phenylhydraz one FCCP	Sigma-Aldrich	Cat#C2920
Rotenone	Sigma-Aldrich	Cat#R8875
Octanoylcarnitine	APExBIO Technology	Cat#B6371-50 mg
Antimycin A	Sigma-Aldrich	Cat#A8674
Coenzyme Q2	Sigma-Aldrich	Cat#C8081
Caffeine	Sigma-Aldrich	Cat#C0750
RNAeasy micro kit	QIAGEN	Cat#74004
Calcium green 5N	Thermo Fisher Scientific	Cat#C3737
Experimental Models: Organism	s/Strains	
C57BL/6J wild-type male and female mice	The Jackson Laboratories	Cat#000664
Experimental Models: Cell lines		
C2C12 cells	ATCC	Cat#Crl-1772
Software and algorithms		
Datlab 7	Oroboros Instruments	Cat#20700
Datlab 8	Oroboros Instruments	Cat#20700



Acqknowledge 4.2	BIOPAC	Cat#ACK100W
Zen software	Zeiss	<u>https://www.zeiss.fr/microscop ie/produits/microscope- software/zen.html</u>
R ImageJ	R	http://cran.r-project.org/ https://imagej.nih.gov/ij/
ImageStudio	LI-COR	https://www.licor.com/bio/ima ge-studio/
Prism 8	GraphPad	https://www.graphpad.com/
Gene Set Enrichment Analysis Software	UC San Diego & The Broad Institute	<u>https://www.gsea-</u> msigdb.org/gsea/index.jsp

Other

Incubator with O ₂ control	BINDER	Cat#6026
WiseStir homogenizer HS-30E Rotina 380R	Wisd Lab Instruments Hettich Lab	Cat#DH.WO01010 Cat#1701
Bike ergometer	Lode Excalibur Sport	Cat#839E
Quark gas analyser	COSMED	n/a
High-voltage constant current stimulator	Digitimer	Cat#DS7A
Isolated voltage stimulator	Digitimer	Cat#DG2A
1-cm diameter electrode	Kendall Meditrace 100	n/a
5-cm diameter electrode	Dermatrode	n/a
5- x 10-cm electrode	Compex	n/a
MP150 acquisition system	BIOPAC	n/a
Oxylite Pro	Oxford Optronix	n/a
C-Pace stimulator	IONOPTIX	n/a
Nanodrop 2000	Thermo Fisher Scientific	Cat#ND-2000
Stellaris 5	Leica	n/a
ViiA 7 Real-Time PCR System	Thermo Fisher Scientific	Cat#4453545
4–15 % Mini-PROTEAN TGX precast protein gels	Bio-Rad	Cat#4561084
Glucose-uptake glo assay	Promega	Cat#J1314
REVERT 700 total protein stain kit	LI-COR	Cat#926-11010
Intercept Blocking Buffer	LI-COR	Cat#927-70003
Uncoated 35-mm-diameter glass-bottom dishes	MatTek	Cat#P35G-0-10-C





Poly-D-lysine coated 35-mm- diameter glass-bottom dishes Stellina Imaging System	MatTek	Cat#P35GC-0-10-C
	Leica	<u>https://www.leica-</u> microsystems.com
6-well plates	Corning	Cat#CL3506
Cytation 3	BioTek	n/a
Bard Magnum biopsy instrument	Bard Radiography	n/a
RS400 heart rate monitor	Polar	n/a
8000Q2 Sensor	Nonin Medical Inc.	n/a
BCA protein assay kit	Thermo Fisher Scientific	Cat#23225
DC protein assay kit	Bio-Rad	Cat#5000111
Oxygraph 2k	Oroboros Instruments	Cat#10003-01
NextGen-02k	Oroboros Instruments	Cat#10101-01

5.1. Experimental models and subject details

Isolation of mitochondria from mouse tissues: Experiments on isolated mitochondria from mouse tissues were performed using male and female wild-type C57BL/6J (14 ± 4 weeks of age; Jackson Laboratories) housed in a temperature-controlled (22 °C) room with a 12/12-h light/dark cycle. All procedures involving animals were performed in accordance with the Austrian Animal Experimentation Act in compliance with the European convention for the protection of vertebrate animals used for experimental and other scientific purposes. After cervical dislocation, the heart and the brain were removed and placed in ice-cold biopsy preservation solution (BIOPS; 2.77 mM Ca-EGTA, 7.23 mM K₂EGTA, 5.77 mM ATP, 6.56 mM MgCl₂, 20 mM taurine, 15 mM phosphocreatine, 20 mM imidazole, 0.5 mM dithiothreitol, 50 mM MES hydrate, pH 7.1). Mitochondria were isolated as previously described [21, 26, 54, 55]. A glass/Teflon Potter-Elvehjem homogenizer (Wisd laboratory instruments) and centrifuge (Andreas Hettich GmbH & Co. KG) were used. All procedures were carried out in an ice bath or at 4 °C.

Mouse heart mitochondria were isolated following Komlodi et al. [26]. Briefly, wet mass of the whole heart was determined, washed with ice-cold BIOPS, and minced with scissors in ice-cold BIOPS (1 mL). The tissue was then digested for 2.5 min in 2 mL of isolation buffer CP1 (100 mM KCl, 50 mM MOPS, 5 mM MgSO₄, 1 mM EGTA, 1 mM ATP; pH 7.4) containing trypsin (9 mg of 13000- 20000 U trypsin/1 g wet mass) with continuous stirring. Immediately after the 2.5 min, 2 mL of isolation buffer CP2 (CP1 buffer plus 0.2 % bovine serum albumin) was added. The homogenate was transferred into a pre-cooled glass/Teflon potter and homogenized at ~1000 rpm (five strokes) in 2-mL isolation buffer CP1. The homogenate was transferred to a 50-mL Falcon tube containing 3 mL isolation buffer CP2 and centrifuged at 10 000 g for 10 min. The supernatant was discarded, the pellet was resuspended in isolation buffer CP1 (final volume 2 mL), and centrifuged at 10 000 g for 10 min. The supernatant was discarded, and the



mitochondrial pellet was finally resuspended in 150 μ L KME buffer (100 mM KCl, 50 mM MOPS, 0.5 mM EGTA).

Mouse brain mitochondria were isolated following Sumbalová *et al.* [56]. Briefly, wet mass was determined, and the tissue was cut into small particles with sharp scissors in isolation buffer C (320 mM sucrose, 10 mM Tris-Cl, 1 mM K-EDTA, 2.5 g/L fatty acid-free bovine serum albumin: pH 7.4). The medium was discarded, the tissue suspended in isolation buffer C (0.1 g tissue/1 mL), transferred to a pre-cooled glass/Teflon potter, and homogenized at 1000 rpm (five strokes). The homogenate was transferred to a 50-mL Falcon tube (0.5 g tissue/20 mL homogenate) and centrifuged at 1 000 g for 10 min. The pellet was discarded, and the supernatant was centrifuged at 6200 g for 10 min. The supernatant was removed, the pellet resuspended in 0.5 g tissue/10 mL of isolation buffer D (320 mM sucrose, 10 mM Tris-Cl, 1 mM K-EDTA; pH 7.4) and centrifuged at 6200 g for 10 min. The supernatant was discarded, and the mitochondrial pellet was finally suspended in 500 μ L isolation buffer D.

Each mitochondrial suspension was gently mixed with a 200- μ L pipette (five updown cycles), and 10 to 20 μ L of heart or brain mitochondrial suspension was injected with a 50- μ L Hamilton syringe into the O2k-chamber through the titration capillary of the stopper, respectively.

Cell lines: C2C12 mouse skeletal muscle myoblasts (American Type Culture Collection) were grown in proliferation medium composed of Dulbecco's modified Eagle's medium (DMEM; Thermo Fisher Scientific,) supplemented with 10 % fetal bovine serum (Thermo Fisher Scientific), 100 IU/mL penicillin, 100 μ g/mL streptomycin (Thermo Fisher Scientific) and 1 % non-essential amino acids (Thermo Fisher Scientific) and maintained at 37 °C in a humidified atmosphere with 5 % O₂ and 5 % CO₂ (Binder Incubators). To induce differentiation, myoblasts were grown to 80-90 % confluence and the proliferation medium was then replaced with a differentiation medium (pre-equilibrated at 5 % O₂ and 5 % CO₂) consisting of DMEM supplemented with 2 % horse serum (Thermo Fisher Scientific).

Human exercise study recruitment: Sixteen male participants were recruited for the study which was approved by the local ethics committee (protocol 2017-00303) and performed in accordance with the Helsinki declaration. Written informed consent was received from each participant prior to study inclusion. All participants were healthy and recreationally physically active. They were familiarized with the electrical stimulation and voluntary contraction procedures at least 48 h before the first experimental session. In this familiarization session participants also performed an incremental test to exhaustion on a cycle ergometer to determine V_{02peak} [33, 57]. The participants were allocated to one of two sprint interval training (SIT) groups (SIT Normoxia and SIT hypoxia; N = 8 per group) based on their V_{02peak} to obtain two homogenous groups. Data from the SIT Normoxia group have been reported previously in Zanou *et al.* ([33]; the group named "SIT").

5.2. Method details

Steady-state hypoxia experiments: Steady-state hypoxia experiments were performed using the methods described in Harrison *et al.* [21] and high-resolution respirometers (O2k or NextGen-O2k; Oroboros Instruments) with a Titration-Injection micropump (TIP2k) for simultaneous measurement of the redox state of the electron-



transfer-reactive Q, $\Delta \Psi_{mt}$, or mitochondrial Ca²⁺ uptake (termed 'associated parameters') at constant O₂ concentrations. O₂ flux (the negative time-derivative of the O₂ concentration) was calculated in real-time by the DatLab software. Corrections of O₂ flux for instrumental background were based on instrumental quality control tests. Experiments were performed under constant stirring (750 rpm) at 37 °C using a calibrated instrument and mitochondrial respiration medium (MiR06; 0.5 mM EGTA, 3 mM MgCl₂, 60 mM lactobionic acid, 20 mM taurine, 10 mM KH₂PO₄, 20 mM HEPES, 110 mM sucrose, 1 g/L BSA, catalase 280 U/mL; pH 7.1) or calcium respiration medium (CaR; 70 mM KCl, 110 mM sucrose, 1 mM MgCl₂, 10 mM KH₂PO₄, 20 mM HEPES, catalase 280 u/mL; pH 7.1). For experiments investigating free Ca²⁺, MiR06 could not be used as it contains a high concentration of EGTA. CaR was used instead.

Oxygen consumption and the associated parameter ($\Delta \Psi_{mt}$, Q-redox state or mitochondrial Ca²⁺ flux) were recorded following injection of isolated mitochondria into the chamber (residual endogenous respiration; *Ren*), after NADH-linked substrates – (Cl-linked) pyruvate (P, 5 mM), malate (M, 2 mM), and glutamate (G, 10 mM) –, succinate (S, 10 mM; CII-linked), and octanoylcarnitine (Oct, 0.5 mM; fatty acid oxidation) were added (LEAK state) and after titration of ADP at kinetically saturating concentration (2.5 mM; OXPHOS capacity, *J*_P) to reach a stable O₂ consumption at kinetically saturating O₂ concentrations (range 100 to 200 µM).

Mitochondrial respiration depleted the O_2 concentration in the closed chamber (anoxia). Mitochondria were allowed to remain anoxic for ~ 2 min to obtain an anoxic signal for zero calibration of the polarographic O₂ sensor [4]. The TIP2k was then used to titrate a specific volume of H₂O₂ into the chamber. This critically important step in the protocol allowed us to calculate the increase in O_2 concentration per volume of H_2O_2 solution titrated into the chamber (catalase in MiR06 dismutates H₂O₂ to O₂ and H₂O). This information was the basis to calculate the flow of H_2O_2 [µL/s] required to set mitochondrial respiration at target oxygen-limited O_2 fluxes from *j* = 0.1 to 0.6 relative to maximal flux. Mitochondrial respiration then returned the system to anoxia before the TIP2k was used to maintain a target oxygen-limited O_2 flux (J_{O_2}) for 3 - 6 min during which time the steady-state O_2 concentration (c_{02}) and an associated parameter were recorded. The injection was then terminated, allowing the system to reach an anoxic state for 2 min for recording bracketing zero O₂ calibrations and anoxic calibrations for the associated parameter. Sufficient H₂O₂ solution was titrated to increase the O₂ concentration in the chamber so that the mitochondria were returned to kinetically saturating O_2 concentrations [21]. In some experiments, several successive steady-state cycles were carried out before and after reoxygenation. The ratio $j = J/J_P$ (or $j = J/J_{max}$) was chosen in random sequence within the range j = 0.1 to 0.6 to avoid any systematic effects. This procedure was repeated for as long as the mitochondria recovered to stable post-anoxic O_2 consumption levels that were not less than 0.9 of the initial J_P . This loss of mitochondrial respiratory capacity is observed independent of H₂O₂ injections and is therefore not related to any oxidative stress induced by such injections [21]. Oxidative stress is not observed in the presence of excess catalase activity even in cases when H₂O₂ injections are more frequently applied [58] and with higher step changes in O₂ concentration [59]. Residual oxygen consumption (*Rox*) was determined after inhibition of CIII by antimycin A (2.5 mM).

The Q-redox state was measured amperometrically using the a three-electrode system as a part of the Q-Module of the NextGen-O2k (Oroboros Instruments) and the Q-mimetic CoQ₂ (1 μ M; Sigma Aldrich) as described in Komlódi *et al.* [26]. The working



electrode was poised at the oxidation potential of CoQ_2 determined by cyclic voltammetry. The reduced Q fraction was calculated by calibrating the raw CoQ_2 signal against 1) the fully oxidized CoQ_2 signal (*Ren*) and 2) the fully reduced CoQ_2 signal (anoxia) and expressed as a fraction of fully reduced CoQ_2 (Figure S1A, C; [26]). The equation fully reduced Q + fully oxidized Q = 1 was used to convert the reduced Q fraction to the oxidized Q fraction [26]. To express the oxidized Q fraction relative to the oxidized Q fraction at maximal O_2 flux (J_{max}), values were divided by the oxidized Q fraction at J_{max} (Figure S1B).

 $\Delta \Psi_{mt}$ was investigated using the fluorescent dye TMRM ([60]; 1 µM; dequench mode). Fluorescence was measured with Green Smart Fluo-Sensors (Oroboros Instruments; 525 nm excitation LED). Filters for the LED and photodiode were selected for TMRM. The fluorescence signal was calibrated before each experiment with stepwise (0.2 µM) titrations of TMRM from 0 to 1 µM. The calibrated TMRM concentration was used in the experiment. For expression of the TMRM signal (i.e., $\Delta \Psi_{mt}$) relative to maximum and minimum values, the LEAK state and anoxia were used respectively (Figure S1F). To express the relative TMRM signal to the relative TMRM signal at maximal O₂ flux, values were divided by the relative TMRM signal at J_{max} .

Calcium Green-5N (2 μ M; which does not inhibit respiration; Thermo Fisher) was used to measure mitochondrial Ca²⁺ uptake in response to changes in free Ca²⁺ in the chamber ([28]). Fluorescence was measured with Blue Smart Fluo-Sensors (Oroboros Instruments; 465 nm excitation LED). Filters for the LED and photodiode were selected for Calcium Green. After titration of mitochondria, 15 μ M EGTA was added to the chamber. CaCl₂ (Sigma Aldrich) was titrated to the desired concentration from a 10 mM stock solution. Calibration of the fluorescence signal was performed during the experiment by two-point calibration. The fluorescence signal before (no CaCl₂ added) and after the first CaCl₂ titration were used. Mitochondrial Ca²⁺ flux was calculated as the negative time derivative of the Ca²⁺ concentration by the DatLab software and was normalized by mitochondrial protein mass.

Mitochondrial protein: Mitochondrial protein content was determined based on Lowry *et al.,* [61] using the DC protein assay (Bio-Rad) following the manufacturer's instructions. The absorbance was measured at 620 nm in a Tecan Infinite TM F200 spectrophotometer (Tecan).

Acute myotube hypoxia experiments: Culture medium was replaced with 2 mL of pre-equilibrated medium (5 % O₂) and myotubes were exposed to either 5 % O₂ or 1 % O₂ in the gas phase for 2 h before electrical stimulation. This duration was to allow for equilibration of the O₂ pressure in the gas phase with the aqueous phase (assessed using an optical O₂ sensor, see below for method; Figure 3B) and to achieve intracellular hypoxia (evaluated as an increase in hypoxia-inducible factor 1 alpha protein content, see below for method, compared with 5 % O₂; Figure 3C&D).

Measurement of media O₂ partial pressure: Partial O₂ pressure (p_{02}) in the media was monitored using optical fluorescence technology that allows real-time detection of p_{02} (Oxylite Pro, Oxford Optronix). The tip of the sensor was placed at the bottom of the well (i.e., at the deepest point of the media).

Western blot: A lysis buffer containing 20 mM Tris/HCl (pH 6.8), 2 mM EDTA (pH 8), 137 mM NaCl, 10 % glycerol, 10 % Triton X-100, 10 mM glycero-phosphate, 1 mM



KH₂PO₄, 1 mM PMSF, 1 mM NaVO₃, 50 mM NaF, 10 mM NaPPi, and a protease inhibitor mixture (Roche) was used to resuspend myotube pellets (100 μ L / well of a 6-well plate) or human muscle biopsies (100 μ L / 5 mg of tissue). The preparation was homogenized with pipette tips for cells or a potter for biopsies, incubated for 1 h at 4 °C then sonicated. Nuclei and debris were then removed by centrifugation at 10 000 *g* at 4 °C for 10 min. Protein concentration was determined using the BCA kit (Thermo Fisher Scientific). Ten to 20 µg of protein were incubated with Laemmli sample buffer containing SDS and 2mercapto-ethanol (Bio-Rad) for 3 min at 95 °C, electrophoresed for 1 h in 4-15 % SDSprecast gradient gels (Bio-Rad), and wet transferred for 1 h onto PVDF membranes. Membranes were stained with REVERT Total Protein Stain (LI-COR) and total protein bands were quantified using the Image Studio software v 5.2.5 (LI-COR). The stain was removed using a removal buffer (LI-COR) and the membranes were blocked for 1 h at room temperature with PBS-LI-COR blocking buffer (LI-COR). Blots were incubated overnight with rabbit anti-Gapdh (Abcam), mouse anti-total OXPHOS (Abcam), rabbit anti-pyruvate dehydrogenase E1 α phospho serine 293 (Abcam) and mouse anti-pyruvate dehydrogenase (Abcam). All the primary antibodies were used at 1: 1000 dilution.

For HIF1 α blots myotubes in one well of a six-well plate were washed once with PBS and collected in a hot (75 °C) lysis buffer containing Laemmli then sonicated. The lysates were then processed as described above using an anti-rabbit HIF1 α primary antibody (Cayman Chemicals).

Membranes were washed in PBS-buffered saline-Tween 20 (TBS-T) and incubated for 1 h at room temperature with 1:10 000 IRDye 680- conjugated donkey anti-mouse or rabbit IgG (LI-COR) and 1: 5000 IRDye 800-conjugated donkey anti-mouse or rabbit IgG (LI-COR) in blocking buffer. Immunoreactive bands were visualized using infrared fluorescence (IR-Odyssey scanner, LI-COR). Band densities were quantified using Image Studio v 5.2.5 (LI-COR). The protein intensity signal was normalized to Gapdh (which was stable across samples and conditions) in human samples [33] while the total protein staining (ponceau for HIF1 α blots) was used to normalize protein content quantified in myotubes. For the human biopsy samples, protein quantifications were expressed as the relative change from Pre [change = (+24 h-Pre)/Pre]. The protein quantifications for cell samples were reported to that of 5 % CTRL cells.

RNA-sequencing: Six biological replicates per condition were used. Cells were rinsed with PBS, pelleted, and then stored at -80 °C. RNA extraction was performed using an RNAeasy kit (QIAGEN).

RNA quality was assessed on a Fragment Analyzer (Agilent Technologies) and all RNAs had a RQN above 9.3. RNA-seq libraries were prepared from 500 ng of total RNA with the Illumina TruSeq Stranded mRNA reagents (Illumina) using a unique dual indexing strategy, and following the official protocol automated on a Sciclone liquid handling robot (PerkinElmer). Libraries were quantified by a fluorometric method (QubIT, Life Technologies) and their quality assessed on a Fragment Analyzer (Agilent Technologies).

Cluster generation was performed with 1.95 nM of an equimolar pool from the resulting libraries using the Illumina HiSeq 3000/4000 SR Cluster Kit reagents and sequenced on the Illumina HiSeq 4000 using HiSeq 3000/4000 SBS Kit reagents for 150 cycles (single end). Sequencing data were demultiplexed using the bcl2fastq2 Conversion Software (version 2.20, Illumina).



Gene expression data analysis: Quality check of RNA sequencing reads was performed with FastQC v0.11.7 [62]. Reads were aligned to the reference mouse model GRCm38 using STAR v2.5.4b [63] and expression levels were quantified using RSEM v1.3.0 [64] with default parameters. ENSEMBL IDs were converted to MGI symbols using Biomart [65]. Differential expression analysis among the 4 groups (1 % CTRL, 5 % CTRL, 1 % STIM, 5 % STIM) was performed using Limma [66] and edgeR [67] following the best practices published in Law *et al.* [68]. DESeq2 [69] was used to generate normalized counts to provide as input for downstream Gene Set Enrichment Analysis (GSEA [70]). GSEA was performed with default parameters using Hallmark Gene Sets [71] and a false-discovery rate (FDR) cut-off at 0.1 (*q*-value 0.1).

Confocal Microscopy: C2C12 myoblasts were plated on uncoated 35-mm-diameter glass-bottom dishes (MatTek) and cultured as described above. At the indicated time of measurement, they were loaded with Mitotracker Green (200 nM; Thermo Fisher Scientific), TMRM (1 μ M; used in dequench mode; Thermo Fisher Scientific), Fluo-4 AM (5 μ M, Thermo Fisher Scientific) or Rhod-2AM (1 μ M; Thermo Fisher Scientific) solubilized in a pre-equilibrated Krebs solution (135.5 mM NaCl, 1.2 mM MgCl₂, 5.9 mM KCl, 11.5 mM glucose, 11.5 mM HEPES, 1.8 mM CaCl₂, pH 7.3) for 15 min in the incubator at the 5 % O₂. Cells were then rinsed twice with a Ca²⁺-free Krebs solution (135.5 mM NaCl, 1.2 mM MgCl₂, 5.9 mM KCl, 11.5 mM glucose, 11.5 mM HEPES, 200 μ M Na-EGTA, pH 7.3).

Images were acquired by z-stack acquisition using a confocal microscope (Stellaris 5, Leica). with APO 20× dry- (Fluo-4AM) or APO 63x/1.40 oil-objectives, with gas (CO₂ = 5%, O₂ = 5 or 1%) and temperature (37 °C) controlled using a stage-top incubator (Ibidi). For acute hypoxic exposure the same 2 h exposure was used. Two-fields of view were acquired per biological replicate. Image analysis was performed on myotubes using ImageJ Software; fixed thresholds for signal intensity were applied across experiments, and the mean intensity was measured. Fluorescence intensities (*F*) were normalized to pre-caffeine stimulation values (*F*₀) to account for possible differences in dye loading and excitation strength. In some experiments, myotubes were stimulated with 2.5 mM (final concentration) caffeine to trigger Ca²⁺ release from the sarcoplasmic reticulum.

Electrical stimulation of C2C12 myotubes: Well-differentiated C2C12 myotubes (5 to 7 days after differentiation) in 6-well plates (Corning, NY, USA) containing 2 mL of differentiation medium were electrically stimulated (IONOPTIX) for 6×30 s (5 s on, 1 s Off) separated by 4 min at 14 V with a stimulation frequency of 50 Hz and a 2-ms pulse duration (SIT-mimicking stimulation; [33]). The differentiation medium was replaced before and after the electrical stimulation. The cells were harvested at the appropriate time point.

Sample preparation for lipidomics: Cells were washed once with PBS then the plates snap frozen on liquid nitrogen.

Lipid extraction: Lipids from frozen skeletal muscle myotubes were extracted by the addition of 2-propanol. Following the scraping, cells were collected to lysis tubes (with 2-propanol) and homogenized by the addition of beads in Precellys tissue homogenizer $(2 \times 20 \text{ s})$. The homogenized lipid solution was then centrifuged (at 4 °C for 15 min at 15 000 rpm) and the supernatant was collected and evaporated to dryness (in SpeedVac,



LabConco). Finally, the dry lipid extracts were resuspended in 2-propanol (volume normalized to the protein content) spiked with Splash mixture (8 %) of isotopically labeled lipid standards (Avanti Lipids) and transferred to LC-MS vials for injection.

Broad-scale targeted lipid analysis: Myotube extracts were analyzed by hydrophilic interaction liquid chromatography coupled to tandem mass spectrometry (HILIC - MS/MS) in both positive and negative ionization modes using a Q-TRAP 6500 LC-MS/MS system (Sciex Technologies; [72]). In both, positive and negative ionization mode, the chromatographic separation was carried out on an Acquity BEH Amide, 1.7 µm, 100 mm \times 2.1 mm I.D. column (Waters). Mobile phase was composed of A = 10 mM ammonium acetate in Acetonitrile: H_2O (95: 5) and B = 10 mM ammonium acetate in Acetonitrile: H₂O (50:50). The linear gradient elution from 0.1 % to 20 % B was applied for 2 min, then from 20 % to 80 % B for 3 min, followed by 3 min of re-equilibration to the initial chromatographic conditions. The flow rate was 600 µL/min, column temperature 45 °C and sample injection volume 2 µL. Optimized ESI Ion Drive Turbo V source parameters were set as follows: Ion Spray (IS) voltage 5500 V in positive mode and -4500 V in negative mode, curtain gas 35 psi, nebulizer gas (GS1) 50 psi, auxiliary gas (GS2) 60 psi, source temperature 550 °C. Nitrogen was used as the nebulizer and collision gas. Optimized lipid class-dependent parameters were used for data acquisition in scheduled multiple reaction monitoring (MRM) mode. Raw LC-MS/MS data were processed using the MultiQuant Software (version 3.0.3, Sciex technologies). For each lipid species the peak area was reported based on its extracted ion chromatograms (EICs) for the monitored MRM transitions.

Quality control of lipidomics: Data quality assessment was performed using pooled quality control (QC) samples analyzed periodically (every ten samples) throughout the entire batch and in a dilution series at the beginning and at the end of the run. The obtained tables (containing peak areas of detected lipids across all samples) were exported to "R" software for signal intensity drift correction using the LOWESS/Spline algorithm followed by peak filtering [73-75]. Peaks with CV > 30 % across QC samples and correlation with dilution factor < 0.65 were removed from further statistical analysis. Lipid annotations used are sphingomyelins (SM), ceramides (CER), dihydroceramides (DCER), lactosylceramides (LCER), hexosylceramides (HCER), monoacylglycerols (MAG), diacylglycerols (DAG), triacylglycerols (TAG), cholesterol esters (CE), phosphatidylinositols (PI), phosphatidylcholines (PC), phosphatidylglycerols (PG), phosphatidylethanolamines (PE), phosphatidylserines (PS), lysophospholipid analogues (LPC, LPG, LPE, LPI, LPS) and free fatty acids (FFA). Data were log10-transformed and normality was tested for each species with the Shapiro test. Differences among stimulation and oxygen levels were assessed with 2-way ANOVA.

Glucose uptake assay: On the day before the assay, the differentiation media was replaced with DMEM containing no serum. At the time of the assay, the culture medium was removed, and cells were washed with PBS pre-heated to 37 °C. Then, cells were incubated with DMEM containing 1 μ M insulin (Promega) for 1 h under the appropriate O₂ pressure. The solution was then removed, and the cells were incubated with freshly prepared 0.1 mM 2-deoxyglucose (Promega) for 30 min under the appropriate O₂ pressure. The uptake process was stopped and neutralized, and luciferase activities were measured using a Glucose Uptake-Glo Assay kit (Promega) and a plate reader (BioTek). Glucose uptake was analyzed according to the manufacturer's instructions.



Human exercise study protocol: Before the experimental session the participants refrained from physical activity and caffeine consumption for 24 h and 12 h, respectively. The SIT session consisted of 6×30 s all-out cycling bouts with 0.7 N·m / kg body mass resistance on a cycle ergometer, separated by 4 min rest between bouts. The SIT Normoxia group performed the exercise session in a normobaric chamber with a fraction of inspired O_2 (F_iO₂) of 20.9 % and the SIT hypoxia group with an F_iO₂ of 14 % [9]. The sprints were preceded by a standard warm-up on the cycle ergometer (5 min at 100 W). Heart rate (Polar) and peripheral capillary O_2 saturation (S_pO₂) was collected from the earlobe (Nonin Medical Inc) before the first and at the end of each sprint. Peak and mean power [W] and the rate of perceived exertion (Borg scale; 6 - 20) for each sprint, and the total work performed [k] were collected.

Neuromuscular function assessments: Knee extensor neuromuscular function of the right (dominant) leg was tested before (Pre), immediately (Post) and 24 h after (+24 h) exercise under ambient normoxia. The tests consisted of a 5 s maximal voluntary contraction (MVC) with a superimposed 100 Hz doublet (paired stimuli) evoked via supramaximal electrical stimulation of the femoral nerve (twitch interpolation technique [76]) followed by supramaximal stimulations of a relaxed muscle evoked at 2 s intervals: a doublet at 100 Hz, 10 Hz, and a single stimulus to obtain the compound muscle action potential (M-wave).

Electromyography: The electromyographic (EMG) activity of the right *vastus lateralis* was recorded with pairs of silver chloride (Ag/AgCl) circular surface electrodes (Kendall Meditrace 100) positioned lengthwise over the middle part of the muscle belly with an inter-electrode (center-to-center) distance of 2 cm according to SENIAM recommendations [77]. The reference electrode was placed over the patella. EMG signals were amplified (gain: 1000), filtered through a 10-500 Hz band-pass filter, and digitized at a sampling frequency of 2 kHz using an AD conversion system (BIOPAC).

Electrical nerve stimulation: A high-voltage (maximum 400 V) constant-current stimulator (DS7AH, Digitimer) was used to deliver single and paired electrical stimuli (pulse width 1 ms). The cathode (Dermatrode) and the anode (Compex) were placed over the femoral nerve at the femoral triangle level beneath the inguinal ligament and on the lower part of the gluteal fold opposite to the cathode, respectively. The optimal stimulation intensity was determined by increasing the current until maximal twitch and M-wave amplitude responses were obtained. This intensity was then increased by 20 % (i.e. supramaximal) and kept constant for all subsequent tests.

Force recordings: Voluntary and evoked forces developed by the knee extensors were recorded at 1 kHz using an isometric ergometer consisting of a custom-built chair equipped with a strain gauge (STS 250 kg, sensitivity 2.0 mV/ V and 1.7 mV/ N, SWJ, China). The strain gauge was attached to the chair on one end and securely strapped above the ankle with a custom-made mold. Participants were seated with a knee angle of 90 ° and a trunk-thigh angle of 100 ° (180 ° = full extension). Extraneous movements of the upper body were limited by two crossover shoulder harnesses and a belt across the lower abdomen. Participants received visual feedback of the torque they produced during the MVCs. Force and EMG data were stored and analyzed off-line with commercially available software (BIOPAC).



Electromyography and torque analyses: Single electrical stimulation pulses were used to measure the amplitude of the first peak of the M-wave before and after exercise. Isometric MVC force was considered as the peak force attained during the voluntary contraction performed at a given time point. The amplitudes of the 10 Hz and 100 Hz paired stimuli (PS10 and PS100) before and after exercise were quantified to assess contractile alterations after exercise and the PS10: PS100 was used as an indicator of low-frequency force depression [78]. PS100 were delivered superimposed on and immediately after MVCs to assess the voluntary activation level (VAL), which was used as an index of central fatigue and assessed as: VAL = $(1 - (superimposed PS100 force \times (force level at stimulation/MVC force)/potentiated PS100 force)) \times 100$ [76]. Rate of force development/relaxation were calculated by dividing the peak twitch force by the time interval from the onset of force development/relaxation to peak force/complete relaxation.

Muscle biopsies: Needle biopsies were taken from the left (non-dominant leg) *vastus lateralis* muscle before, ~10 min and 24 h after exercise [79]. Briefly, after skin sterilization and local anesthesia, a 1 to 2 mm long skin cut was made with the tip of a scalpel. Needle biopsies were collected using an automatic biopsy device. A 14-gauge disposable trocar mounted in the device was inserted through the cut, perpendicular to the muscle fibers, until the fascia was pierced. Three samples (~15 mg each) were collected from one puncture site at each time point. Muscle samples were immediately frozen in liquid nitrogen and stored at -80 °C until analysis.

Quantification and statistical analysis: Data analysis was performed in Excel, R, and Prism as described above. All data (unless otherwise noted) was presented as mean \pm SD. Unless otherwise noted, *p* values were calculated using one-way ANOVA for multiple comparisons involving a single variable, a two-tailed Student's t test for pairwise comparisons and a two-way ANOVA for multiple comparisons involving two-variables. Details of statistical analyses and *N* values are found in the Figure Legends. *N* indicates biological replicates or number of human participants for *in vivo* experiments. Where applicable, *p* values were adjusted for multiple hypothesis testing using the Benjamini-Hochberg procedure.

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Supplementary Figures

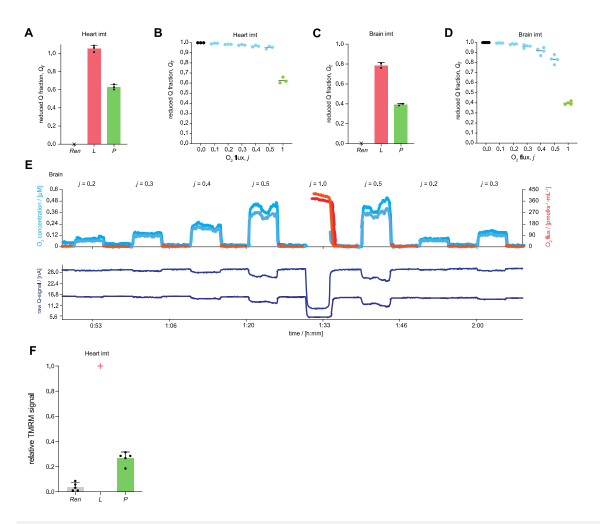


Figure S1. Steady-state respirometry measurements in mitochondria isolated from mouse cardiac muscle and brain related to Figure 1. (A) Reduced Q-fraction in mitochondria isolated from mouse cardiac muscle (*N*=3) in the LEAK (*L*) and OXPHOS (*P*) states calibrated relative to residual endogenous respiration (Ren; fully oxidized) and anoxia (fully reduced; $Q_r = 1$). (B) Quantification of reduced Q-fraction under anoxia (j =0), at relative O_2 flux $j = 0.1, 0.2, 0.3, 0.4, 0.5, and maximal <math>O_2$ flux (j = 1) in mitochondria isolated from mouse cardiac muscle (N=3). *j* is the ratio of the set J_{0_2} to the measured OXPHOS capacity (*I*max). (C) Reduced Q-fraction in mitochondria isolated from mouse brain (N=1) in the LEAK (L) and OXPHOS (P) states calibrated relative to residual endogenous respiration (Ren; fully oxidized) and anoxia (fully reduced). (D) Reduced Qfraction under anoxia (i = 0), at relative O₂ flux i = 0.1, 0.2, 0.3, 0.4, 0.5, and maximal oxygen flux (i = 1; N=1) in mitochondria isolated from mouse brain. (E) Q-redox state under steady states of functional hypoxia and at maximal O₂ flux in mitochondria isolated from mouse brain respiring on NADH-linked substrates, succinate, and fatty-acid at kinetically saturating [ADP]. Each shade represents a different chamber. **(F)** Relative TMRM signal in mitochondria isolated from mouse cardiac muscle in the OXPHOS state (J_{max} ; N=5) calibrated relative to LEAK (maximum) and anoxia (minimum). Bars represent the means, error bars the standard deviations, dots are individual values.



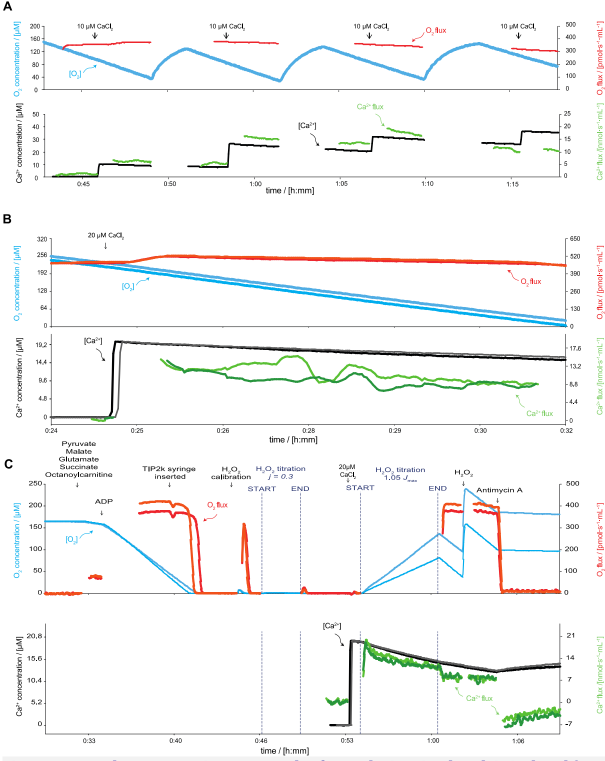


Figure S2. Steady-state respirometry and Ca²⁺ uptake in mitochondria isolated from mouse cardiac muscle related to Figure 2. (A) Mitochondrial O₂ and Ca²⁺ fluxes in response to sequential titrations of 10 μ M CaCl₂ in the OXPHOS state (saturating [ADP]) with NADH-linked substrates, succinate, and fatty acid. (B) Mitochondrial O₂ and Ca²⁺ fluxes in response to titration of 20 μ M CaCl₂ in the OXPHOS state with NADH-linked substrates & succinate & fatty acid. Each shade represents a different chamber. (C) Mitochondrial O₂ and Ca²⁺ fluxes in response to titration of 20 μ M CaCl₂ during a steady-



state experiment at maximal oxygen flux (J_{max}). Each shade represents a different chamber.

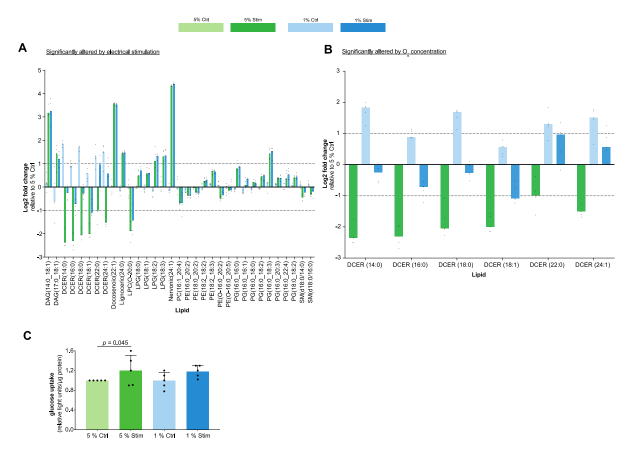
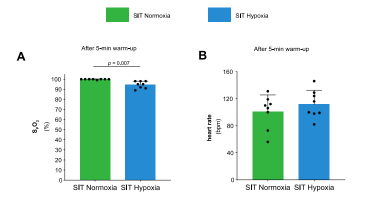


Figure S3. Functional hypoxia altered the response of transcripts, signaling, and metabolism to myotube contraction related to Figure 4. (A) Lipid species significantly altered by electrical stimulation in myotubes immediately after stimulation in 5 % or 1 % O_2 compared with non-stimulated controls (*N*=5). Lipid annotations used are diacylglycerols (DAG), dihydroceramides (DCER), lysophosphatidylcholines (LPC), lysophosphatidylglycerols (PG), phosphatidylcholines (PC), phosphatidylglycerols (PG) and sphingomyelins (SM). (B) Lipid species significantly altered by O_2 levels after stimulation in 5 % or 1 % O_2 compared with non-stimulated controls (*N*=5). (C) Glucose uptake in myotubes immediately after stimulation in 5 % or 1 % O_2 and their controls (*N*=5). Bars represent the means, error bars the standard deviations, dots are individual values.



Functional hypoxia reduces mitochondrial calcium uptake



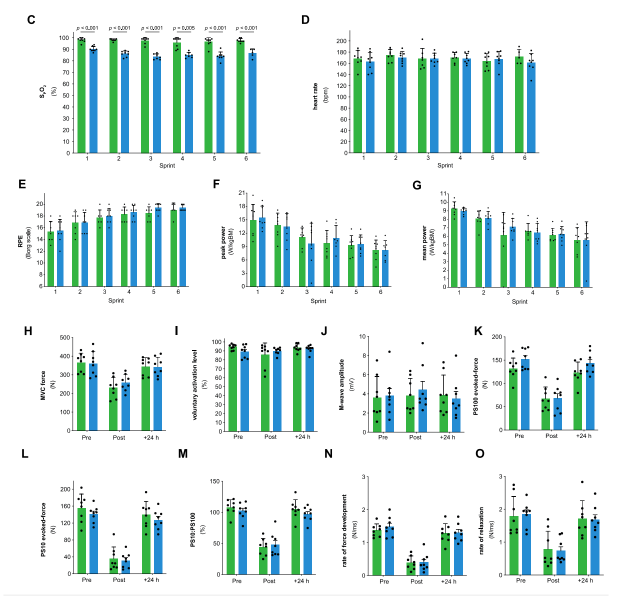


Figure S4. Physiological, performance and neuromuscular data from the human exercise trial related to Figure 4. (A) Peripheral capillary oxygen saturation before the start of the SIT session. **(B)** Heart rate before the start of the SIT session. **(C)** Peripheral capillary oxygen saturation during the SIT session. **(D)** Heart rate at the end of each sprint during the SIT session. **(E)** Rate of perceived exertion during the SIT session. **(F)** Peak



power output during the SIT session. **(G)** Mean power output during the SIT session. **(H)** Maximal voluntary contraction force before, immediately after, and 24 h after the SIT session. **(I)** Voluntary activation level before, immediately after, and 24 h after the SIT session. **(J)** M-wave amplitude before, immediately after, and 24 h after the SIT session. **(K)** PS100 force before, immediately after, and 24 h after the SIT session. **(K)** PS100 force before, immediately after, and 24 h after the SIT session. **(K)** PS100 force before, immediately after, and 24 h after the SIT session. **(M)** PS10:PS100 force before, immediately after, and 24 h after the SIT session. **(M)** PS10:PS100 force before, immediately after, and 24 h after the SIT session. **(N)** Rate of force relaxation before, immediately after, and 24 h after the SIT session. **(O)** Rate of force relaxation before, immediately after, and 24 h after the SIT session. *N=*8 for all tests; bars represent the means, error bars the standard deviations, dots are individual values.