High-intensity sprint training inhibits mitochondrial respiration through aconitase inactivation

Filip J. Larsen,†,‡,†,1 Tomas A. Schiffer,* Niels Ørtenblad,‡ Christoph Zinner,§,¶
David Morales-Alamo,‖ Sarah J. Willis,§ Jose A. Calbet,‖ Hans-Christer Holmberg,§
and Robert Boushel†,‡

‡Department of Physiology and Pharmacology, Karolinska Institute, Stockholm, Sweden; †Swedish School of Sport and Health Sciences, Stockholm, Sweden; †Institute of Sports Science and Clinical Biomechanics, Muscle Research Cluster, University of Southern Denmark, Odense, Denmark; §Swedish Winter Sports Research Centre, Department of Health Sciences, Mid Sweden University, Östersund, Sweden; ‖Department of Sport Science, Julius Maximilians University, Würzburg, Germany; †Research Institute of Biomedical and Health Sciences (IUIBS), Las Palmas de Gran Canaria, Canary Islands, Spain; and §School of Kinesiology, University of British Columbia, Vancouver, British Columbia, Canada

ABSTRACT Intense exercise training is a powerful stimulus that activates mitochondrial biogenesis pathways and thus increases mitochondrial density and oxidative capacity. Moderate levels of reactive oxygen species (ROS) during exercise are considered vital in the adaptive response, but high ROS production is a serious threat to cellular homeostasis. Although biochemical markers of the transition from adaptive to maladaptive ROS stress are lacking, it is likely mediated by redox sensitive enzymes involved in oxidative metabolism. One potential enzyme mediating such redox sensitivity is the citric acid cycle enzyme aconitase. In this study, we examined biopsy specimens of vastus lateralis and triceps brachii in healthy volunteers, together with primary human myotubes. An intense exercise regimen inactivated aconitase by 55–72%, resulting in inhibition of mitochondrial respiration by 50–65%. In the vastus, the mitochondrial dysfunction was compensated for by a 15–72% increase in mitochondrial proteins, whereas H₂O₂ emission was unchanged. In parallel with the inactivation of aconitase, the intermediary metabolite citrate accumulated and played an integral part in cellular protection against oxidative stress. In contrast, the triceps failed to increase mitochondrial density, and citrate did not accumulate. Instead, mitochondrial H₂O₂ emission was decreased to 40% of the pretraining levels, together with a 6-fold increase in protein abundance of catalase. In this study, a novel mitochondrial stress response was highlighted where accumulation of citrate acted to preserve the redox status of the cell during periods of intense exercise.—Larsen, F. J., Schiffer, T. A., Ørtenblad, N., Zinner, C., Morales-Alamo, D., Willis, S. J., Calbet, J. A., Holmberg, H.-C., Boushel, R. High-intensity sprint training inhibits mitochondrial respiration through aconitase inactivation. FASEB J. 30, 417–427 (2016).

Key Words: exercise · mitochondrial dysfunction · reactive oxygen species · citrate

Aerobic exercise training is well established as a potent stimulus of mitochondrial biogenesis pathways, and it improves cardiorespiratory fitness, insulin sensitivity, and physical performance (1, 2). A long-held view has been that changes in mitochondrial content and activation by ADP are the only means by which skeletal muscle oxidative capacity increases or decreases. This view has been challenged recently, as alterations in mitochondrial quality, coupling, and efficiency, commonly termed intrinsic mitochondrial properties, have been found after training or dietary interventions (3–5). Several recent studies have shown that the formation of physiologic concentrations of reactive oxygen species (ROS) is an important cell-signaling process that is indispensable for the activation of mitochondrial biogenesis and cellular proliferation (6). These positive effects of moderate levels of ROS are in contrast to the detrimental effects of excessive oxidative stress associated with several pathologic conditions (7) and extreme endurance events (8, 9). Although the topic is controversial (10), there is a dynamic balance between optimal exercise intensities that yield moderate oxidative stress with beneficial adaptations and excessive exercise that produces severe oxidative stress and maladaptation. This interesting dose–response relationship between ROS levels and cellular outcome is an area of intense research (11).

Despite reports of robust health-promoting effects, intense exercise reproducibly increases oxidative stress

Abbreviations: ADP, adenosine diphosphate; AU, arbitrary units; BCA, bicinchoninic acid (assay); BSA, bovine serum albumin; COX, cytochrome c oxidase; CS, citrate synthase; ETS, electron transport system; FBS, fetal bovine serum; GSH, reduced form of glutathione; GSSG, oxidized form of glutathione; HIT, high-intensity training; HRP, horseradish peroxidase;
(continued on next page)
(12), indicated by the elevated oxidation product malondialdehyde and oxidation of glutathione immediately after exercise (13). The elevated oxygen consumption during intense aerobic exercise therefore increases the generation of ROS. Both NADPH oxidases and xanthine oxidoreductase have been found to be activated during exercise and to contribute significantly to total ROS stress (14). Further, mitochondria from skeletal muscle show higher ROS production after contractions than that shown in the resting state (15). Thus, the production of mitochondrial ROS during exercise initiates a potential signaling cascade where the level of activity (i.e., oxygen turnover), induces a feedback signal to the nucleus, regulating the abundance of mitochondrial proteins that enable cells to meet the demand for energy.

Structures that promptly adapt to changes in oxidative stress are necessary to maintain homeostasis. Aconitase is a redox-sensitive enzyme primarily located in the mitochondrial tricarboxylic acid (TCA) cycle, where it facilitates the isomerization of citrate to isocitrate via cis-aconitase and subsequent delivery of NADH to the electron transport system (ETS). Aconitase has an active [Fe₄S₄]²⁺ cluster that is rapidly inactivated into [Fe₃S₄]⁺ through oxidation by superoxide, peroxynitrite, and H₂O₂, constituting a redox-sensitive regulation of the rate-limiting enzyme in the TCA cycle. The location of a redox-sensitive enzyme in the TCA cycle implies that aconitase can act as a rheostat that modulates mitochondrial metabolism by its interaction with ROS (16). We sampled human muscle from tissue samples taken before and after a short-term high-intensity training program, to assess the resilience of mitochondrial respiration and aconitase to high-intensity training and redox stress. By using samples from both the vastus lateralis and the triceps brachii, we sought to identify the muscle-specific response and the delineation between the optimal redox stress that induces mitochondrial biogenesis and the excessive ROS stress that leads to maladaptation. We used cultured primary myotubes to assess the role of the endogenous increases in citrate concentrations in the defense against excessive ROS stress. We further sought to distinguish the mitochondrial response to high levels of ROS induced by the intense training program and the cellular adaptations that maintain cell integrity.

MATERIALS AND METHODS

Subjects

Twelve healthy male subjects [mean age, 24.2 ± 3.8 yr; height, 183.8 ± 6.8 cm; weight, 80.0 ± 14.7 kg, and body mass index, 23.6 ± 0.9 kg/m²] volunteered for the study. The subjects classified themselves as untrained or only moderately active. All subjects gave written consent after being informed of the study procedure. The study was approved by the Regional Ethics Review Board in Umeå, Sweden, and was performed in conformity with the principles of the Declaration of Helsinki. This study was a part of a larger study involving 16 subjects; however, mitochondrial respiration and biochemical measurements were undertaken on tissue samples obtained from 12 subjects. Therefore, all data reported in this study are from these 12 subjects, but samples from 3 additional subjects were used for the isolation of myogenic satellite cells and culturing of myotubes.

Testing procedure: high-intensity training

To adjust the ergometers (Schoberer Rad Messtechnik SRM, GmbH, Jülich, Germany) for leg and arm cycling for all subsequent testing and to become familiar with the equipment and procedures, the subjects visited the laboratory 2 times. The subject, seated position arm cycling, was positioned in a way that the axle of the crank was positioned just below the level of the heart and with a comfortable elbow angle when the cranks were in a horizontal position (17). Performance tests were executed before and after the high-intensity training (HIT) intervention. The subjects performed 4 consecutive 4 min sessions of exercise (100 rpm) at submaximal intensity, adjusted so that the respiratory exchange ratio (RER) in the last session was >1.0. After a rest of 10 min, maximum oxygen consumption (V O₂peak) was determined by an incremental test in which the subject performed to volitional exhaustion, as follows: arm cycling started at a work load of 20 W, where the subjects freely chose the pedal frequency (within the range of 70–105 rpm) with the workload increased by 10 W each 30 s. Leg cycling started at 60 W, with an increase of 25 W each 30 s. V O₂peak was calculated from the plateau of oxygen uptake with the following criteria: an elevation of V O₂ of <2 ml · min⁻¹ · kg⁻¹ with increasing work load: RER >1.1, heart rate within ± 2.5% of the age-adjusted maximum, and capillary blood lactate concentration of 6 mM (18). Three minutes after the participants experienced exhaustion, achievement of V O₂peak was verified by having them pedal for as long as possible at the peak resistance attained during the incremental test +10 W during arm cranking and +25 W during leg cycling. On the second day of testing and to evaluate the effect of the intervention on passive recovery between all-out efforts, the subjects performed 2 Wingate anaerobic tests separated by 4 min of recovery. The subjects remained seated and were encouraged verbally by the researchers during the tests. The gas-exchange equipment was calibrated before the tests with a mixture of 4.5% CO₂ and 16% O₂ and a 3-L syringe (Hans Rudolph Inc., Kansas City, KS, USA). The first testing procedure was performed 3 d after the baseline specimen was extracted and 2 d before the first training session. The posttest procedure was completed 2 d after the 6th training session, and then a last training session was performed to allow for standardized training stimuli and time span before the posttest biopsy was performed 40 h later.

Training schedule

The sessions included both arm and leg cycling (separated by 1 h of recovery) on 7 separate days during a 15 d period. Each subject performed 30 s maximum sprints separated by 4 min of rest: 4 such sprints with both the arms and legs on training d 1 and 2, 5 sprints were performed on d 3 and 4, and 6 sprints were performed on d 5, 6, and 7.

Extraction of tissue specimens

Biopsies were performed before the first training session and 40 h after the last training session. The samples were extracted with a Bergström needle under local anesthesia (2–3 ml 2% mepiva- cage) in randomized order from both sides of the vastus lateralis and triceps brachii (distal portion of the lateral head). After the
tissue was dried on filter paper, it was divided into 3 pieces: the first for immunoblot analysis, the second for enzymatic activities (both snap-frozen in liquid N2), and the third for mitochondrial isolation (placed in ice-cold isolation medium and used immediately).

**Immunoblot analysis**

Muscle samples were homogenized in a blender (Bullet Blender Blue; Next Advance, Averill Park, NY, USA) in ice-cold lysis buffer (2 mM 4-(2-hydroxyethyl)-1-piperazineethanesulfonic acid (HEPES) (pH 7.4), 1 mM EDTA, 5 mM EGTA, 10 mM MgCl₂, 50 mM β-glycerophosphate, 1% Triton X-100, 1 mM NaVO₄, 2 mM DTT, and 1% phosphatase inhibitor cocktail (P-2850; Sigma-Aldrich, St. Louis MO, USA)). Homogenates were centrifuged at 10,000 g at 10 min. Supernatant was collected and stored at ~80°C until analyzed. To determine protein concentration of the homogenates, supernatant was diluted 1:100 in distilled water and analyzed with a bicinchoninic acid (BCA) protein assay (Micro BCA assay kit; Thermo Fisher Scientific, Waltham MA, USA). Samples were adjusted by adding lysis buffer to obtain similar protein concentration followed by dilution in Laemmli sample buffer [125 mM Tris-HCl 4% SDS, 0.04% bromphenol blue, 20% glycerol, and 4% 2-mercaptoethanol (pH 6.8)]. After the samples were heated at 95°C for 5 min, they were loaded on precast gradient gels (Criterion TGX, 4–20% acrylamide; Bio-Rad, Hercules, CA, USA). The electrophoresis was run at 175 V [25 mM Tris base, 192 mM glycine, and 3.5 mM SDS (pH 8.3)], whereupon the proteins were transferred to PVDF membranes (Immun-Blot PVDF; Bio-Rad) for 1.5 h (300 mA) (25 mM Tris base, and 192 mM glycine). Membranes were blocked in Tris-buffered saline [TBS; 20 mM Tris base and 137 mM NaCl (pH 7.6)] containing 5% nonfat dry milk and 0.1% Tween 20 for 1 h before incubation with commercially available primary antibodies overnight or 1.5 h in room temperature (1:1000 in TBS, 2.5% milk). The membranes were washed in TBST with milk and incubated with secondary antibodies for 1 h (1:10,000 in TBST with milk). After the membranes were washed in TBST, the target proteins were visualized by chemiluminescent detection (Chemidoc MP Imaging System; Bio-Rad) and analyzed with software Image Lab 5.0 (Bio-Rad). After reanalyzing the same PVDF membrane for different proteins, the membranes were stripped by using Restore Western Blot Stripping Buffer (Thermo Fisher Scientific), according to the manufacturers’ instructions, and subsequently reprobed them with the following primary antibodies: citrate synthetase, ab96600 (Abcam; Cambridge MA, USA) cytochrome oxidase (COX)-IV-1 (ab3759; Abcam), AMPKα1 (ab133448), PGC1α (ab 54481), and catalase (ab16731; Abcam). Secondary antibodies were anti-mouse horseradish peroxidase (HRP)–linked antibody (7076) and antirabbit HRP-linked antibody (7074; both from Cell Signaling Technology).

**Isolation of primary myogenic satellite cells**

The muscle samples were placed in PBS at 4°C overnight. The sample was then washed in PBS and cleared from connective tissue. A few drops of Trypsin-EDTA 0.25% (Thermo Fisher-Gibco, Grand Island, NY, USA) were added, and the specimen was homogenized with a pair of scissors. After 7 ml trypsin was added, the homogenate was transferred to a small beaker and put in the incubator with gentle stirring for 20–30 min. After sedimentation, the supernatant was collected, and the trypsin was blocked with a few ml of complete medium to a final concentration of at least 2% fetal bovine serum (FBS). After centrifugation, the cells were resuspended in 10 ml cell medium and plated in a 10 cm² Petri dish for 20–30 min, to let adherent cells attach. The cells remaining in suspension were then transferred to a 25 cm² cell culture flask. The cell culture medium consisted of 50% DMEM (low glucose) and 50% F12 (both from Thermo Fisher-Gibco), containing 50 U/ml penicillin, 50 U/ml streptomycin, 1.25 µg/ml amphotericin, and 20% FBS. For differentiation, the medium was exchanged with a similar medium containing 2% FBS for 5 d before the cells were harvested or treated.

**Respirometric analysis mitochondria**

Mitochondrial isolation was performed as published (19). Respirometric analysis on isolated mitochondria was performed by high-resolution respirometry (O2-K; Oroboros, Innsbruck, Austria) in a respiration medium containing 0.5 mM EGTA, 5 mM MgCl₂, 60 mM potassium-lactobionate, 20 mM taurine, 10 mM KH₂PO₄, 20 mM HEPES, 110 mM sucrose, and 1 g/L bovine serum albumin (BSA). Saturated levels of pyruvate (5 mM), malate (1 mM), and ADP (2.5 mM) were used as substrates for the determination of complex I-mediated respiration, together with succinate (10 mM) for complex I+H₂-mediated respiration. Mitochondrial experiments were performed at an oxygen pressure of 15–20 kPa O₂ in the chamber.

**Mitochondrial yield**

After differential centrifugation, the final mitochondrial pellet was resuspended in preservation solution at 0.6 µl/mg initial muscle tissue weight. Mitochondrial yield was estimated from the protein concentration per microliter preservation solution, as determined by the BCA method, according to the manufacturer’s instructions (Thermo Fisher Scientific).

**Respirometric analysis on differentiated primary myotubes**

Respirometric analysis on mitochondria and differentiated cultured primary myotubes was performed by high-resolution respirometry (O2-K; Oroboros). Basal cell respiration was determined in respiration medium (contents as described earlier). After basal cell respiration, the cells were permeabilized with digitonin. Saturating levels of pyruvate (5 mM), malate (1 mM), and succinate (10 mM) were used as substrates (Sigma-Aldrich), together with ADP (2.5 mM). The cells were exposed to H₂O₂ at various concentrations (20–200 µM) while oxygen consumption was measured.

**Mitochondrial H₂O₂ emission**

Oxygen consumption and H₂O₂ production were measured simultaneously (O₂-K with fluorometer; Oroboros). H₂O₂ was measured fluorometrically with the extrinsic fluorophore Amplex UltraRed (10 µM; thermo Fisher Scientific) in combination with HRP (1 U/ml). Calibration of the H₂O₂ signal was performed by stepwise increases of 0.1 µM H₂O₂ obtained from a prepared stock solution [40 µM H₂O₂ (Sigma-Aldrich), 10 µM HCl].

**Oxidative markers in muscle tissue**

Several commercial kits were used to measure different oxidative markers in muscle homogenates. A ratio–detection kit (Abcam) was used for measuring the ratio of the oxidized to the reduced form of glutathione (GSSG/GSH). Protein carbonylation was measured with a protein carbonyl content assay kit (Sigma-Aldrich). An
Citrate synthase activity measurements

Enzyme activities were measured in freeze-dried muscle dissected from nonmuscle constituents and homogenized on ice in a 50 mM phosphate buffer containing 1 mM EDTA and 0.05% v/v Triton X-100 (pH 7.4). To disrupt the mitochondria and expose the citrate synthase (CS), homogenates were freeze thawed 4 times with liquid nitrogen. CS activity was determined by the addition of oxaloacetate to a buffer solution containing muscle homogenate, DTNB (5,5’-dithiobis-2-nitrobenzoic acid) buffer, and acetyl-CoA. The rate change in absorbance (405 nm) was monitored over 6 min (model 650; Beckman Coulter, Pasadena, CA, USA), converted into enzyme activity rates, and expressed as a percentage of the pretraining value.

Citrate levels in muscle homogenates

A citrate assay kit (MAK057-1KT; Sigma-Aldrich) was used to determine the concentration of citrate in the muscle homogenates, according to the manufacturer’s instructions. The citrate content of the eluate was then analyzed by spectrophotometry, according to the manufacturer’s instructions, and related to the total protein content in the sample.

Results are expressed as means ± SEM. Student’s t test or 1-way ANOVA was used for statistical analyses (Prism 5.0; GraphPad, San Diego, CA, USA). P < 0.05 denoted statistical significance.

RESULTS

HIT increases work capacity, mitochondrial density, and oxidative enzyme activity, without change in myosin heavy chain composition

After training, aerobic fitness, measured as VO2peak, increased by 6% during incremental leg cycling and by 12% during arm cranking (Fig. 1A; both P < 0.001). In parallel, the mean power outputs during repeated 30 s sprints increased by 4% in the legs (P < 0.01) and 8% in the arms (P < 0.05) (Fig. 1B). After training, the mitochondrial proteins CS, mitochondrial transcription factor A (TFAM), and COXIV increased after training in the vastus but not in the triceps (Fig. 1C, D). Likewise, there was a tendency toward elevated CS activity after training, compared with the pretraining level in the vastus (P = 0.07), but not in the triceps. We found no evidence of increased protein levels of peroxisome proliferator-activated receptor gamma coactivator (PGC)-1α [before training, 105 ± 20 arbitrary units (AU); after training, 104 ± 21 AU], AMPK (before training, 64 ± 20 AU; after training, 53 ± 17 AU), or phospho-AMPK (before training, 44 ± 14 AU; after training 67 ± 21 AU) (data not shown), indicating that the

Figure 1. The effects of HIT on physiologic parameters and mitochondrial density markers in skeletal muscles. A) VO2peak before and after the training period. VO2peak averaged for 30 s was measured during incremental exercise with a normal leg cycle ergometer, and arm VO2peak was measured at a separate occasion with an arm-cranking ergometer in the seated position. B) Mean workload attained during 30 s maximum-effort sprints with the leg cycle and the arm cranking ergometer (n = 12). C, D) Western blot analysis of 3 selected mitochondrial proteins and CS activity before (pre) and after (post) training in the vastus lateralis (C) and triceps brachii (D). Densitometric analysis of each band, normalized for protein content against Reactive Brown, revealed the variations between before and after training. CS activity was measured with a spectrophotometric approach [n = 8 (D) and 10 (C)]; error bars, SEM. *P < 0.05; **P < 0.01; ***P < 0.001.
posttraining specimens were extracted outside of the acute training response after the last training session.

Myosin heavy chain (MHC) composition of type I, IIA, and IIB fibers before training was 44 ± 3, 53 ± 3, and 3 ± 1% in vastus and 28 ± 3, 65 ± 4, and 7 ± 3% in triceps, respectively. The distribution of type I fibers was significantly lower ($P < 0.01$) and that of the type IIA fibers was higher ($P < 0.05$) in the triceps than in the vastus. However, training did not alter the MHC distribution in the vastus (49 ± 3, 49 ± 3, and 2 ± 1%) or in the triceps (26 ± 2, 70 ± 2, and 4 ± 2%) (data not shown).

**Intrinsic respiration is suppressed after HIT due to inactivation of aconitase**

An increase in mitochondrial content is a well-known adaptation to an increase in metabolic demand, but less is known about how intrinsic parameters of mitochondrial function are affected by intense training. To explore this question, we isolated mitochondria from fresh skeletal muscle tissue and analyzed the respiratory capacity and function by high-resolution respirometry. After the training intervention, several indices of intrinsic mitochondrial respiration unexpectedly decreased by 50–65% in both vastus and triceps mitochondria (Fig. 2A–C, F–H).

To explore the mechanisms underlying the reduced intrinsic mitochondrial respiration, we tested the hypothesis that oxidative modifications of aconitase account for the inhibition of mitochondrial electron flux. Previous findings indicate that the TCA enzyme aconitase is readily inactivated in skeletal muscle by reactive species and may limit mitochondrial respiration (20, 21). We found that, after training, the aconitase activity expressed per milligram wet muscle decreased significantly in the triceps but not in the vastus (Fig. 2D, I). In skeletal muscle, aconitase is primarily located in the mitochondria and considering the increase in several mitochondrial proteins, we also expressed aconitase activity per milligram mitochondrial protein and found a marked decrease after training in both muscles (Fig 2E, J). Taken together, these results demonstrate profound alterations of intrinsic mitochondrial function with HIT training that are independent of global changes in mitochondrial density.

**HIT decreases mitochondrial $\text{H}_2\text{O}_2$ emission and induces markers of oxidative modification**

Mitochondrial ROS production occurs at specific sites of the ETS. We analyzed the emission of $\text{H}_2\text{O}_2$ in various respiratory states and loci of intact mitochondria, using Amplex red as a fluorometric probe. We found that mitochondrial ROS production was diminished in the triceps mitochondria, whereas it was unchanged in the vastus mitochondria (Fig. 3A–F).

The inactivated aconitase and compensatory decrease in mitochondrial $\text{H}_2\text{O}_2$ emission indicate that HIT induced a severe cellular oxidative challenge. We therefore sought

---

**Figure 2.** Intrinsic parameters of mitochondrial respiration and aconitase activity in vastus lateralis (A–E) and triceps brachii (F–J) before and after training. A, F) Leak respiration is the respiratory rate in the presence of the respiratory substrates malate and pyruvate without addition of adenylates. B, G) State 3 complex I is the maximum ADP stimulated respiration rate in the presence of complex I substrates pyruvate and malate. C, H) State 3 complex I+II is similar to (B, G), but with convergent electron flux through complex II by adding succinate ($n = 12$). D, I) Aconitase activity is expressed per milligram wet weight tissue. E, J) Aconitase activity is expressed per microgram mitochondrial protein ($n = 5$, vastus; 6, triceps); error bars, SEM. *$P < 0.05$; **$P < 0.01$; ***$P < 0.001$. 

---

to examine the impact of the HIT training intervention on the global redox environment in the skeletal muscle biopsy specimens. We first determined the redox status of the glutathione pool, but did not find any significant changes in GSH, GSSG, or the GSSG/GSH ratio after HIT vs. before (Supplemental Fig S1). Thereafter, we assessed more chronic oxidative protein modifications by measuring the protein carbonylation status in the tissue samples. Again, we found a trend toward more carbonylated proteins in the triceps after training vs. before, but in the vastus we found no such changes (Supplemental Fig S1D). We next investigated whether the endogenous antioxidative defense system was activated by HIT to maintain redox balance. We measured protein levels of catalase in the muscle samples and found a large increase after HIT vs. B. Citrate accumulates after training and protects the cell against oxidative stress

With the finding of an increase in CS and a concomitant inactivation of aconitase that is located downstream in the TCA cycle, it can be assumed that intramuscular citrate levels should accumulate. We analyzed citrate concentration in muscle specimen homogenates and found an increase from 0.56 to 1.28 nmol/μg protein in the vastus from before to after training (P=0.03; Fig. 4B). However in the triceps samples, citrate levels were unchanged (0.85 to 0.72 from before to after training; P=0.40; Fig. 4B). Citrate is commonly added to food, where it acts as a mild antioxidant and food preservative by chelation of divalent iron. We hypothesized that the inactivation of aconitase and resulting increase in citrate are important steps in the restoration of the cellular redox state. We therefore investigated the cytoprotective effects of citrate after oxidative stress induced by externally added H2O2. When cultured primary myotubes were exposed to 50–200 μM H2O2 for 24 h, cellular mitochondrial content decreased significantly, as assessed by the protein content of CS (Fig. 4C). Addition of 10 mM sodium citrate fully abolished this degradation, indicating that citrate protected mitochondrial proteins from oxidative stress and mitophagy (Fig. 4D).

**DISCUSSION**

In the present study, aconitase acted as a sensitive redox-regulated protein, responding to exercise and oxidative stress by inactivation, which led to inhibition of the mitochondrial

---

**Figure 3.** Release of H2O2 from isolated mitochondria and protein expression of catalase before (pre) and after (post) HIT. H2O2 emission was measured in isolated mitochondria with Amplex Ultrasensitive Red used as a fluorometric probe simultaneously with the respirometric analysis of leak (A, D), state 3 (B, E), and complex I+II (C, F) (Fig. 2 and Scheme 2). A–F) H2O2 emission in mitochondria from the vastus lateralis (A–C) and triceps brachii (D–F). H2O2 emission is expressed per microgram mitochondrial protein (n=12). G, H) Pre- and posttraining catalase expression in vastus and triceps. Western blot (G) and densitometric analysis of each band (H) revealed variations between before and after training (n=9, vastus; 12, triceps). Error bars, SEM. **P<0.01; ***P<0.001.
respiration that protects the cell from further oxidative stress. As aconitase was inactivated, citrate accumulated and protected the mitochondria from oxidative degradation. Simultaneously, mitochondrial density increased, to maintain muscle oxidative capacity and the energy status of the cell. Excessive inhibition of aconitase, as seen in the triceps, not only inhibited mitochondrial respiration, but also abolished the increase in citrate and mitochondrial density and induced a parallel robust activation of the endogenous antioxidant catalase that restored redox homeostasis.

Aconitase is present in 2 isoforms: one is mitochondrial the other cytosolic. The mitochondrial isoform is predominant in skeletal muscle (22) and is a part of the TCA cycle, where it forms a vital structure in mitochondrial ATP production. The cytosolic isoform is involved in iron metabolism (23). Exercise has been linked to elevated levels of ROS production, and aconitase is readily inactivated by ROS. It has been proposed that aconitase is inactivated during intense exercise. However, whether this occurs in vivo is not fully clear. One study found inactivation of aconitase after exhaustive exercise in rats (24), whereas another study involving human subjects showed increased aconitase activity after an acute bout of medium-intensity exercise (25). That the clinical manifestations of Friedrich ataxia, a mitochondrial disorder characterized by aconitase deficiency, are severe exercise intolerance, fatigue, and progressive damage of the nervous system is interesting and indicates that functional aconitase is essential in maintaining redox balance and the cellular antioxidant defense system.

In this study, we found evidence of a profound mitochondrial aconitase inactivation by a brief period of HIT. After HIT training, aconitase activity per mitochondrion was reduced to 45 and 28% of the pretraining value in the vastus and triceps, respectively. The moderate reduction in the vastus was compensated for by increased mitochondrial biogenesis, which appeared to occur in parallel. When aconitase activity was expressed per unit of wet weight muscle we found a similar level in the vastus before and after training, whereas activity in the triceps was still was only 57% of the pretraining value. The finding that other mitochondrial enzymes increased in parallel with aconitase inhibition as a compensatory mechanism bears strong resemblance to a pathologic aconitase deficiency, as affected patients have higher mitochondrial densities than matched controls (22). The observed difference between the vastus and triceps indicates that the triceps was under more severe oxidative stress during HIT and could not fully compensate for the reduction in aconitase activity by increasing mitochondrial density. The functional impact of aconitase inhibition manifested in

---

**Figure 4.** Functional effects of training in skeletal muscle and exposure to \( \text{H}_2\text{O}_2 \) in myotubes. A) Inhibition of respiration in myotubes after \( \text{H}_2\text{O}_2 \) exposure. Human primary myotubes were permeabilized with digitonin and acutely exposed to increasing concentrations of \( \text{H}_2\text{O}_2 \). Respiration was measured by high-resolution respirometry (\( n = 3 \)). B) Citrate content in muscle homogenates before and after exercise, measured before and after training with a spectrophotometer (\( n = 5–6 \)). C, D) Human primary myotubes were exposed to control or \( \text{H}_2\text{O}_2 \) in the absence (C) or presence (D) of 10 mM citrate for 24 h. Mitochondrial density was assessed by the protein content of CS. A matched control set of experiments was obtained for each treatment (\( n = 5–7 \)). Error bars, SEM.* \( P < 0.05 \); ** \( P < 0.01 \).
the reduction in respiration per milligram mitochondrial after training in both the vastus and triceps. Mitochondrial biogenesis compensated for this bioenergy loss in the vastus, given that mitochondrial proteins were up-regulated in the range of 15–72% after training. However, we found no significant increase in triceps muscle mitochondrial proteins after training. It should be noted that we observed a much more homogenous change in mitochondrial proteins in the vastus, which contributed to the statistical significance in this muscle group, whereas there was more variation in the triceps. Although a significant change in mitochondrial density markers was not noted in the triceps, it may still play a role in the adaptation to buffering the ROS produced from HIT.

To probe whether the different MHC distributions between arm and leg contributed to our observations, we ran correlation analyses between both the inhibition of mitochondrial respiration and aconitase inactivation against the baseline MHC distribution. We found no convincing associations between any of these variables and cannot state that the MHC composition contributed to our findings. However, we cannot exclude a fiber-type–linked mechanism underlying the compensatory reduction in mitochondrial ROS production and antioxidant response between arms and legs. This possibility warrants further investigation.

Mitochondria are important sources of ROS production, and down-regulation of mitochondrial activity and ROS production could constitute a sophisticated cellular response to quickly restore redox homeostasis. Pharmacological inhibition of complex I-mediated respiration by rotenone has been found to increase mitochondrial ROS production, with deleterious cellular consequences (26). We found unexpectedly that H$_2$O$_2$ emission after training was reduced to 40% of the pretraining value in triceps, whereas it was unchanged in the vastus, demonstrating a more severe oxidative challenge in the triceps. Secondary to a reduction in mitochondrial ROS production, a more chronic adaptation to elevated oxidative stress, is the biogenesis of enzymes of the antioxidant defense system. In triceps tissue we found a 6-fold induction after training of protein levels of the H$_2$O$_2$-scavenging enzyme catalase. Consistent with the finding that oxidative stress was less severe in the vastus, no significant increase in catalase was found in this muscle group.

The functional impact of attenuated mitochondrial respiratory capacity, as a component of the O$_2$ cascade contributing to $V_{O2\text{peak}}$, and exercise capacity with this form of training is noteworthy. Despite a rather marked mitochondrial inhibition, both exercise capacity and $V_{O2\text{peak}}$ increased in the present study. This finding substantiates the apparent overcapacity of mitochondrial respiration compared to the capacity to deliver oxygen to the working tissue (27). One function of this overcapacity could be to constitute a necessary buffer to avoid energy limitations caused by oxidative inactivation by aconitase and thereby mitochondrial respiration. The improvements in $V_{O2\text{peak}}$ could be attributable to structural and functional components unrelated to intrinsic mitochondrial capacity (28).

A striking result in this study was the opposing response to training of 2 adjacent enzymes of the TCA cycle. The increased protein content and activity of CS and the inactivation of aconitase infers that citrate should accumulate. Apart from being a TCA cycle intermediate, citrate also inhibits phosphofructokinase and acts as a metabolic sensor [for a review, see Iacobazzi and Infantino (29)]. We hypothesized that the increase in cellular citrate content is an integral part of the antioxidative defense system. In support of this assumption, when we added citrate to the medium of cultured myotubes, it fully abolished the mitophagic response to a challenge with H$_2$O$_2$ (Fig 4C, D). Citrate has also been shown to play an intricate role in

![Scheme 1](image-url). The experimental protocol. The numbers in parentheses indicate the number of 30 s sprints per training session.
metabolic control through its diverse chemical properties and ability to interact with other biochemical pathways, especially phosphofructokinase, which inhibits glycolysis (29). Another interesting finding is that the reactivation of aconitase after oxidative stress is fully dependent on the presence of citrate through an interaction with frataxin (30). Citrate is also a known chelator of divalent cations, such as Fe²⁺, and thereby indirectly interferes with the Fenton reaction between iron and H₂O₂, likely effecting the production of hydroxyl radicals. The activity of CS has for unknown reasons been shown to increase after an acute bout of endurance exercise (31), and citrate release from skeletal muscles is increased during exercise (32). It can be speculated that the increase in citrate formation during exercise is not only a reflection of the TCA cycle intermediate pool but also a functional response to protect the muscle cell against oxidative stress.

The contrasting response to HIT between the vastus and triceps in the present study is intriguing. It is clear that there is a dose–response relationship between ROS production and cellular outcome. Several studies have indicated that mild oxidative stress is necessary for initiation of the adaptive response to exercise, in that administration of dietary antioxidants can prevent the beneficial effects of an exercise program (6, 33). However, severe oxidative stress in pathologic conditions or during overtraining can lead to maladaptation, with detrimental long-term effects (34). The results of the present study indicate that oxidative stress in the vastus had less severe consequences, because the inactivation of mitochondrial aconitase was compensated for by an increase in mitochondrial density. On the contrary, several lines of evidence indicate that oxidative stress was excessive in the triceps. First, the absence of increases in mitochondrial density indicates that this muscle group did not respond to training as expected. The lack of increase in mitochondrial density resulted in a lower aconitase activity per milligram tissue after training. In addition, mitochondrial H₂O₂ emission was reduced.

Scheme 2. The skeletal muscle mitochondrial response to HIT. Exercise induces superoxide production to different degrees, depending on muscle group. In the triceps, catalase expression is increased, whereas it is unchanged in the vastus. H₂O₂ inactivates aconitase and, as a consequence, mitochondrial respiration is inhibited. In triceps, mitochondrial ROS production decreases in parallel. With decreased mitochondrial respiration, the capacity to produce ATP is diminished. This lowered energy state is sensed and, if oxidative stress is within the antioxidant capacity of the cell (as in the vastus), it is compensated for by increasing global mitochondrial density. As CS is markedly increased and aconitase inactivated, citrate accumulates in the tissue and protects the mitochondria from oxidative degradation. If oxidative stress exceeds a certain threshold (as in the triceps), however, the biogenesis of catalase is initiated to protect against further damage by H₂O₂. The mitochondrial biogenesis process fails to increase mitochondrial density, and CS and other mitochondrial proteins also do not increase. Thereby, a negative loop of events is initiated, because citrate does not increase, leaving the mitochondria even more prone to oxidative degradation, with loss of cellular energy capacity.
after training, most likely a response to counteract oxidative stress. Further, a near significant increase in protein carbonyl products was found in triceps homogenates. Last, protein content of catalase was increased by more than 6-fold, a logical response to increased cellular levels of H₂O₂. The present data suggest that the vastus responds to training as expected, with a functional increase in mitochondrial density, probably mediated by mild oxidative stress that initiates crucial signaling events. Despite the similar training stimuli between arm and leg exercise, the triceps seemed to be more vulnerable to oxidative stress induced by HIT, as indicated by the increased oxidative stress markers and maladaptive response to training. The origin of this different response between the vastus and triceps can only be speculated upon. Whether the differentiated response in the triceps vs. the vastus is dependent on intrinsic characteristics of the specific limb muscles or degree of muscle use is not obvious. Even though the subjects in this study were untrained, they had above-average aerobic capacity, as measured by VO₂peak testing. Nonetheless, that humans are more accustomed to regular use of leg muscles for locomotion could explain the findings in this study. It is likely that a period of exercise training with lower intensity before the more intense sprint training preconditions the muscle cells against the oxidative stress and potentiate the cellular adaptations. Indeed, it has recently been shown that a period of low-intensity exercise protects dystrophin-deficient MDX mice that are vulnerable to the oxidative stress induced by high-intensity exercise (35). The lower training status of the triceps could therefore be a part of the explanation of the maladaptation to intense exercise in this muscle group.

In summary, we show that a short-term, intense exercise program in previously untrained subjects induces oxidative stress that inactivates aconitase and initiates an orchestrated cellular response to re-establish the redox balance and energy status of the cell. A proposed chain of events is illustrated in Schemes 1 and 2.

This study was funded by Swedish National Centre for Research in Sports. D.M.A. is a fellow of the Dr. Manuel Morales Foundation. The authors declare no conflicts of interest.

REFERENCES


Received for publication May 29, 2015.
Accepted for publication September 14, 2015.
High-intensity sprint training inhibits mitochondrial respiration through aconitase inactivation

Filip J. Larsen, Tomas A. Schiffer, Niels Ørtenblad, et al.

FASEB J 2016 30: 417-427 originally published online October 9, 2015
Access the most recent version at doi:10.1096/fj.15-276857

Supplemental Material
http://www.fasebj.org/content/suppl/2015/10/09/fj.15-276857.DC1

References
This article cites 29 articles, 6 of which can be accessed free at:
http://www.fasebj.org/content/30/1/417.full.html#ref-list-1

Subscriptions
Information about subscribing to The FASEB Journal is online at
http://www.faseb.org/The-FASEB-Journal/Librarian-s-Resources.aspx

Permissions
Submit copyright permission requests at:
http://www.fasebj.org/site/misc/copyright.xhtml

Email Alerts
Receive free email alerts when new an article cites this article - sign up at
http://www.fasebj.org/cgi/alerts