Satellite cell self-renewal in endurance exercise is mediated by inhibition of mitochondrial oxygen consumption

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Abstract

Background  Skeletal muscle stem cells (satellite cells) are well known to participate in regeneration and maintenance of the tissue over time. Studies have shown increases in the number of satellite cells after exercise, but their functional role in endurance training remains unexplored.

Methods  Young adult mice were submitted to endurance exercise training and the function, differentiation, and metabolic characteristics of satellite cells were investigated in vivo and in vitro.

Results  We found that injured muscles from endurance-exercised mice display improved regenerative capacity, demonstrated through higher densities of newly formed myofibres compared with controls (evidenced by an increase in embryonic myosin heavy chain expression), as well as lower inflammation (evidenced by quantifying CD68-marked macrophages), and reduced fibrosis. Enhanced myogenic function was accompanied by an increased fraction of satellite cells expressing self-renewal markers, while control satellite cells had morphologies suggestive of early differentiation. The beneficial effects of endurance exercise were associated with satellite cell metabolic reprogramming, including reduced mitochondrial respiration (O\textsubscript{2} consumption) under resting conditions (absence of muscle injury) and increased stemness. During proliferation or activated states (3 days after injury), O\textsubscript{2} consumption was equal in control and exercised cells, while exercise enhanced myogenic colony formation. Surprisingly, inhibition of mitochondrial O\textsubscript{2} consumption was sufficient to enhance muscle stem cell self-renewal characteristics in vitro. Moreover, transplanted muscle satellite cells from exercised mice or cells with reduced mitochondrial respiration promoted a significant reduction in inflammation compared with controls.

Conclusions  Our results indicate that endurance exercise promotes self-renewal and inhibits differentiation in satellite cells, an effect promoted by metabolic reprogramming and respiratory inhibition, which is associated with a more favourable muscular response to injury.

Keywords  Metabolism; Muscle stem cells; Mitochondria; Bioenergetics

Introduction

Satellite cells are the resident stem cells of the skeletal muscle, capable of regenerating the tissue throughout adulthood in response to insults.\textsuperscript{1} Regeneration occurs by activation of satellite cells, which can then differentiate and either fuse with existing multinuclear contractile myofibres and repair them or generate new myofibres by fusing with other activated satellite cells.
Satellite cells express paired box protein 7 (Pax7), a critical regulator of muscle stem cell maintenance and activation. Indeed, loss of this protein leads to lack of satellite cell expansion and differentiation in both neonatal and adult muscles. Activation of these stem cells into committed myoblasts occurs because of extrinsic myogenic stimuli and typically involves expression of the myogenic regulatory factor MyoD. The committed myogenic progenitors then actively proliferate to generate sufficient committed myoblasts, as well as produce differentiated myocytes. Activated cells can also return to quiescence and self-renew, a process marked by Spry1 and Notch expression.

One of the most effective stimuli to induce satellite cell activation in adult muscles is exercise. Indeed, resistance or workload exercise has long been known to activate and increase numbers of satellite cells, as well as promote myonuclear expansion and hypertrophy. Interestingly, however, the effects of aerobic endurance exercise on satellite cell function are less understood, and mixed results have been described in the literature. This is an important point of study, because endurance exercise is well known to improve skeletal muscle fitness and is therefore expected to affect satellite cell activation, self-renewal, and differentiation. Increased satellite cell content has consistently been described after endurance training in young and old animals. In addition, aerobic exercise conditioning increases the capacity of the skeletal muscle to repair after injury in aged mice. However, the mechanisms involved in the changes in the function of satellite cells promoted by endurance exercise training have not yet been investigated.

Recent studies have shown that mitochondria may play a role in the regulation of myogenesis. Mitochondrial function in differentiated myocytes is modified by endurance exercise, which promotes increased biogenesis of the organelle, as well as up-regulated mitochondrial quality control pathways. Indeed, respiratory capacity per mitochondrion is lower in middle aged compared with young humans and may be an important factor in age-related muscle dysfunction and differences in fibre-type distribution, which are partially prevented by exercise. Mitochondrial function is also involved in stem cell activation and differentiation (reviewed in Khacho and Slack). Thus, understanding how mitochondria are involved in satellite cell function could provide insights into mechanisms that regulate muscle health.

Although the type of mitochondrial functional alteration during commitment to differentiation is variable with stem cell fate, activation of oxidative phosphorylation is often required upstream of differentiation, while suppression is necessary for self-renewal in pluripotent stem cells. Little is known, however, about how changes in mitochondrial bioenergetics are related to satellite cell activation, self-renewal, or differentiation. In this study, we investigated the effects of endurance exercise training on skeletal muscle regeneration and satellite cell function. Surprisingly, we found that endurance exercise induces a significant repression of satellite cell oxidative phosphorylation, which stimulates self-renewal.

Methods

Animals

Two-month-old, young adult and still growing male C57BL/6 mice were purchased from AniLab, Paulinia, Brazil. All experimental procedures were conducted in agreement with Ethical Principles in Animal Research of the Brazilian College of Animal Experimentation (CONCEA) and reviewed and approved by the local animal care and ethics committee (137/2017). Mice were housed 3–4 per cage and maintained on a 12:12 h light–dark cycle in a temperature-controlled environment (22°C) with ad libitum standard laboratory chow (Nuvital Nutrientes, Colombo, Brazil) and water.

Incremental speed test and endurance exercise protocol

Mice were submitted to incremental speed exercise testing on a motor treadmill before and after the experimental period of endurance training. After adaptation to the treadmill environment for 1 week (10 min per session), mice were placed on the treadmill at 0% inclination and allowed to acclimatize for at least 10 min. Incremental speed increases then started at 10 m/min during 10 min and were increased by 3 m/min every minute (n = 7, control; n = 13, exercised). Mice that could no longer run for over 1 min were considered exhausted and removed from the treadmill. This incremental speed test provided the maximal running duration (min), distance (m), and velocity (m/min).

Endurance exercise was randomly assigned for sedentary (control) and exercised mice as 60% of the maximum intensity (maximal aerobic capacity) reached during the incremental speed test, as previously described. Exercised mice performed moderate-intensity running exercises on a motorized treadmill for 5 weeks (from the 8th to 13th weeks of age), 5 days a week, 60 min/day. The running velocity during the period of physical training was gradually increased to 60% of maximal aerobic intensity (described earlier). During 5 weeks, treadmill-running skills were maintained in sedentary (control) mice for 5 min (10 m/min), twice a week, to avoid any interference of treadmill stress. This activity in control animals did not alter maximal exercise capacity. Body weights and food consumption were recorded weekly.

Forty-eight hours after the last exercise at 5 weeks of endurance training, run capacity (maximal running times, velocities, and distances) was evaluated in all mice. After this session, the animals were submitted to live animal experiments,
as described in the succeeding text, or anaesthetized with isoflurane and sacrificed by cervical dislocation for tissue or cell collection. Tissues were quickly removed, weighed, immediately frozen in liquid nitrogen, and stored at −80°C.

**Indirect calorimetry, spontaneous physical activity, and maximal oxygen consumption (VO\textsubscript{2max}) determination**

A room with restricted access was dedicated for this experiment. Oxymax (Columbus Instruments, Columbus, OH, USA) and TSE Systems were used to monitor real-time changes in gas concentrations. Animals \((n = 4, \text{control}; n = 4, \text{exercised})\) were placed in individual 30-cm-diameter cages. Each mouse was acclimatized to an individual chamber 24 h before the measurements, and data were collected every 20 min for 24 h. Results shown were collected during the active period, corresponding to the early dark hours.

Infrared detection was used to measure locomotor activity in parallel with the indirect calorimetry assessments. Data were recorded for at least one full circadian period (24 h), and results shown were collected during the early dark hours. Movements were added for every minute and detected simultaneously on all three axes: forward and backward (x), side to side (y), and up and down (z).

Coupling of the treadmill with adjustable intensity to indirect calorimetry and an open-circuit calorimeter allowed for the assessment of maximal oxygen consumption (VO\textsubscript{2max}). Prior to the experiment, a familiarization session was run. Gases were measured every 10 s. Average VO\textsubscript{2} and running distance were calculated for each mouse \((n = 10, \text{control}; n = 12, \text{exercised})\).

**Muscle lesions**

For muscle regeneration studies, injury was induced 24 h after the last endurance exercise session. Mice \((n = 4, \text{control}; n = 4, \text{exercised})\) were anaesthetized by inhalation using isoflurane. The belly region of the tibialis anterior muscle was injected intramuscularly with 50 μL of 1.2% w/v barium chloride. Seven days after the injury, the animals were anaesthetized with isoflurane and sacrificed by cervical dislocation. The muscles were quickly removed and stained.

For satellite cell activation/proliferation studies, muscle injury was induced 24 h after the last exercise session. Mice were anaesthetized, and the belly region of the tibialis anterior, gastrocnemius, quadriceps, and triceps brachii muscles were injected intramuscularly with 10 μL of 1.2% w/v barium chloride. Three days after the injury, the animals were sacrificed, skeletal muscles were quickly removed, and satellite cells were isolated as described in the succeeding text.

**Stains and microscopy**

For haematoxylin and eosin (H&E) staining analysis, injured muscles were embedded into tragacanth and Tissue-Tek® O.C.T. (Sakura Quality-Somagen, AR, Netherlands) solution and frozen in liquid nitrogen. Sections (10 μm thick) were obtained from the mid-belly region of the injured tibialis anterior muscle \((n = 4, \text{control}; n = 4, \text{exercised})\) using a Leica CM3050 cryostat (Wetzlar, Germany). Sections were H&E stained and photographed using a Nikon Eclipse E1000 microscope (Nikon’s Microscopy, MA, USA) coupled to a DXM 1200 camera. Non-overlapping images from the central region of each section were analysed using the cell counter plugin for ImageJ software developed at the National Institutes of Health and the Laboratory for Optical and Computational Imaging, MD, USA (central nuclei or in adjacent muscle sections). Mean cross-sectional areas were determined by measuring the circumferences (μm\textsuperscript{2}) of 25 adjacent fibres in six sections per slide per animal, thus a total of 150 fibres per animal \((n = 4, \text{control}; n = 4, \text{exercised})\). Data represent the averages of the mean values for each mouse in each group. Analysis was performed blinded.

Picosirisur Red was used to evaluate collagen content in injured muscles, as described by Junqueira et al.\textsuperscript{37} Sections were photographed using a polarized light microscope, and images were captured on a JVC TK-1280E digital color video camera (JVC, Burladingen, GER) coupled to the microscope. A section of muscle fibres surrounded by perimysium, containing 10–100 fibres \((n = 4, \text{control}; n = 4, \text{exercised})\), was imaged as previously reported.\textsuperscript{37} Analysis was performed blinded.

**Dual X-ray absorptiometry measurements**

Forty-eight hours after the last endurance exercise session, mice \((n = 4, \text{control}; n = 4, \text{exercised})\) were scanned using the FX PRO In-Vivo Imaging Systems (BRUKER Corporation, MA, USA). A combination of xylazine (10 mg/kg) and ketamine (90 mg/kg) was used for sedation. Fat area was measured in the abdomen (top of the pelvis to the lowermost rib) and calculated as abdominal fat divided by the total abdominal tissue. Lean area was evaluated in the upper hindlimbs.

**Muscle stem cell isolation, culture, and clonal analysis**

Muscle stem cells were isolated from pooled intact extensor digitorum longus, gastrocnemius, quadriceps, soleus, tibialis anterior, and triceps brachii muscles, as previously described.\textsuperscript{38–40} Briefly, muscles were removed, minced, mechanically dissociated, and filtered twice with 70 μm strainers. The resulting tissue was centrifuged and digested with 2% type II collagenase, 0.25% trypsin, and 0.1% DNase. Primary myoblasts were purified by 2–3 pre-plating steps.\textsuperscript{41}
and cultured in Dulbecco’s modified Eagle’s medium containing 1% penicillin and 20% foetal bovine serum. We typically isolated 450 000–550 000 satellite cells from the pooled hindlimb and forelimb muscles of each mouse using this protocol. Cell number and viability were determined using a Neubauer chamber and trypan blue vital staining, respectively. Cells were not passaged. In purity checks, 84.6% of the isolated cells with 4,6-diamidino-2-phenylindole (DAPI)-stained nuclei were Pax7 positive (n = 8), as assessed by immunofluorescence (described in the succeeding text).

For clonal analysis, isolated cells were grown for 168 h in low-glucose Dulbecco’s modified Eagle’s medium containing 1% penicillin and 20% foetal bovine serum on Matrigel-coated six-well plates (5000 cells per well) and then fixed with 3.8% paraformaldehyde, dried for 24 h, dyed with 1% crystal violet, and washed. After 24 h, clone colonies (clusters of approximately 50 cells) were manually counted using a Nikon SMZ 745 stereoscopic microscope (Nikon Instruments, NY, USA) with 7.5 zoom (n = 4, control; n = 4, exercised).

**Immunofluorescence**

Satellite cells from control and exercised mice were plated for 96 h on Matrigel-coated 12-well plates (100 000 cells per well), then fixed with 3.8% paraformaldehyde for 10 min, permeabilized with 0.3% Triton X-100, and blocked in 1% bovine serum albumin, 0.3% Triton X-100, and 0.1% sodium azide in phosphate-buffered saline for 2 h at room temperature. Cells were incubated with primary antibody (Pax7 monoclonal mouse IgG1, Abcam 199010; 1:50, Cambridge, MA, USA) at 4°C overnight followed by DAPI for 1 h and fluorescent-labelled secondary antibodies (goat anti-mouse IgG conjugated to Alexa Fluor® 488; 1:1000, Life Technologies Corporation, CA, USA). Images were captured with CoolSNAP HQ CCD camera (Photometrics, Tucson, AZ, USA) driven by IPLab software (Scanalytics Inc., Milwaukee, WI, USA) using a Leica DMI 6000B fluorescent microscope (Mannheim, Germany). Quantification used the cell counter plugin for ImageJ software.

Injured muscle cross sections were fixed with 4% paraformaldehyde at room temperature for 10 min, washed with phosphate-buffered saline with 0.1% Tween 20 (PBS-T), and incubated for 1 h with blocking solution (1% bovine serum albumin in PBS-T). Samples were incubated overnight at 4°C with primary rabbit anti-eMHC antibody (1:500, Origene Cat#TA349138, Rockville, MD, USA) and anti-CD68 (1:500, mAb #76437). After washing with 0.1% PBS-T, the secondary antibody was added in blocking solution for 1 h in the dark, followed by further washing. Slides were mounted with VECTASHIELD for fluorescence with DAPI (Cat# H-1200, Vector Labs, CA, USA). Images were captured with a CoolSNAP HQ, CCD camera (Photometrics) driven by IPLab software (Scanalytics Inc.) using a Leica DMI 6000B fluorescence microscope (Mannheim, Germany). Quantification was conducted using the cell counter plugin for ImageJ software.

**Morphological analysis**

Satellite cells from control and exercised mice were cultured for 96 h on Matrigel-coated 12 well plates (100 000 cells per well, n = 4, control; n = 4, exercised). Images were acquired on a Nikon Eclipse TS100 microscope, and quantification of spindle-like and elongated morphology was conducted using the cell counter plugin for ImageJ.

**Metabolic flux analysis**

Oxygen consumption rates (OCRs) and extracellular acidification rates (ECARs) of muscle stem cells were measured using a Seahorse XF24 Analyzer (Seahorse Bioscience, MA, USA), as described previously. Cells (animal n = 4, control; n = 4, exercised) were seeded on Matrigel-coated XF24 24-well microplates at 20 000 cells per well for 120 h. Before the assay, the media were removed and replaced by 500 mL of low-glucose (5 mM) assay medium, without sodium bicarbonate, to allow for ECAR measurements. ATP synthesis-linked O2 consumption and proton leak-driven respiration were determined by the addition of oligomycin (1 μg/mL). The uncoupler 2,4-dinitrophenol (2,4-DNP, 200 μM) was added to promote maximal respiratory capacity. All respiratory modulators were used at concentrations determined through preliminary titration analyses.

**RNA extraction and real-time polymerase chain reaction analysis**

Cells were cultured on six-well plates (250 000 cells per well) for 120 h (animals used were n = 8, control, and n = 8, exercised). For mtDNA quantifications, DNA samples were prepared using a DNeasy® Blood & Tissue Kit (Qiagen, SP, Brazil). Real-time quantitative polymerase chain reaction (qPCR) was performed using SYBR Green PCR Master Mix (Applied Biosystems, MA, USA). The following genes were analysed: hypoxanthine-guanine phosphoribosyltransferase (HPRT) for nDNA and mitochondrially encoded NADH:ubiquinone oxidoreductase core subunit 1 (MT-ND1) for mtDNA copy number. Fold changes were calculated by the method. The primer sequences are presented in Supporting Information, Table S1.

Total RNA was purified using TRizol reagent (Invitrogen Life Technologies, Rockville, MD, USA) and quality checked using 260/230 and 260/280 nm scores. Equivalent contents of RNA were reverse transcribed using a SuperScript III CDNA Synthesis Kit (Invitrogen, CA, USA). The cDNA synthesized was stored at −20°C prior to the real-time PCR assay. Amplification was performed using Platinum® SYBR® Green qPCR SuperMix UDG (Invitrogen Life Technologies, Carlsbad, CA,
USA) and evaluated by real-time PCR using the Rotor Gene 3000 apparatus (Corbett Research, Mortlake, Australia). Primer sequences were designed using information contained in the Gene Bank of the National Center for Biotechnology Information. Gene expression was quantified using qBase software (Qbase, WA, UK), as described previously. Expression was normalized using the internal control gene HPRT. Fold changes relative to HPRT were calculated by the $2^{-\Delta\Delta CT}$ method. Primers were selected using PrimerBank (https://www.exxtend.com.br); sequences are presented in Table S1.

Muscle stem cell transplantation

The tibialis anterior muscles of the hosts, 14-week-old male mice, were prepared by BaCl$_2$ injury as described earlier 1 day before transplantation; then, 20 000 freshly isolated muscle stem cells from control or exercised mice in 30 μL media were slowly injected into the mid-belly of the muscle of a recipient anaesthetized host (control) mouse ($n=4$, control; $n=4$, exercised; and $n=4$, antimycin A). Seven days after transplantation, the muscles were quickly removed for microscopy.

Statistical analysis

Results are presented as mean ± standard error of the mean and were analysed using GraphPad Prism Software 4.0 (GraphPad Company, CA, USA). Data were compared using t-tests to determine if the means of two sets of data are significantly different from each other or one-way analysis of variance used to compare means of two or more samples, followed by the Bonferroni post hoc test. Asterisks denote statistically significant differences vs. control. Grubb’s test was used to exclude outliers.

Results

Endurance exercise increases performance and promotes a shift in substrate use

Mice were submitted to an endurance exercise protocol for 5 weeks (Figure 1A), a time span sufficient to observe many of metabolic phenotypes associated with exercise. Their performance was then evaluated 48 h after the last exercise training session. Consistent with prior reports, exercised mice presented an increase in running time (Figure 1B), speed (Figure 1C), and distance (Figure 1D) in the run-to-exhaustion test. Exercised mice also displayed a shift in energy substrate usage at rest analysed through indirect calorimetry, as evidenced by increased O$_2$ consumption (Figure 1E) and heat production (Figure 1F). Interestingly, spontaneous physical activity (Figure 1G) was also increased in exercised mice. Performance was then measured under run-to-exhaustion conditions. Oxygen consumption (Figure 1H) was again higher in exercised animals. Blood glucose monitoring during the

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**Figure 1** Endurance exercise induces a performance-dependent shift and changes energy substrate usage. (A) Schematic panel. (B) Running time. (C) Running speed. (D) Distance covered in the run-to-exhaustion test ($n=7$, control; $n=13$, exercised). (E) O$_2$ consumption during the dark hours (18–21 h) measured by indirect calorimetry. (F) Heat production during the dark hours measured by indirect calorimetry. (G) Spontaneous physical activity during the dark hours ($n=4$, control; $n=4$, exercised). (H) O$_2$ consumption at exhaustion ($n=9$, control; $n=11$, exercised). (I) Resting blood glucose, blood glucose during run-to-exhaustion, and blood glucose at 10 min rest after exhaustion ($n=7$, control; $n=13$, exercised). Asterisks denote statistically significant differences (* $P<0.05$, ** $P<0.01$, and *** $P<0.001$) vs. control.

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Endurance exercise enhances skeletal muscle repair

Endurance exercise has been shown to protect against loss of satellite cells during ageing, as well as increase their number and capacity to differentiate.\textsuperscript{17,19,46} We thus verified if satellite cell-mediated muscle regeneration after injury (Figure 3A) was affected by endurance exercise. H&E microscopies of uninjured and injured tibialis anterior muscles from control (sedentary) or exercised animals are shown in Figure 3B and uncover no changes promoted by exercise in the decreased average cross-sectional area measurements with injury (Figure 3C). However, injured muscles from exercised mice had higher density of fibres with central nuclei, suggestive of newly formed myofibres due to improved regenerative capacity (Figure 3D). Quantification of embryonic myosin heavy chain (eMHC)-expressing myofibres, a hallmark of regeneration in adult muscle, was also increased (Figure 3E and 3F). Endurance exercise training also decreased the extent of inflammatory cell infiltration (Figure 3G) and the presence of cells expressing the macrophage marker CD68 (Figure 3H and 3I). Indeed, Picrosirius Red staining (Figure 3J), used to quantify the fibrous tissue area (Figure 3K), demonstrates that exercised animals showed lower collagen content. Overall, these results show that endurance exercise training significantly accelerates the regeneration efficiency of muscle and improves muscle regenerative response, resulting in higher new myofibre appearance, lower inflammation, and prevention of fibrosis.

Endurance exercise enhances satellite cell self-renewal

The higher density of newly formed fibres and lower inflammation in endurance-exercised mice suggest changes in satellite cell population.\textsuperscript{19,46} We thus isolated and plated satellite cells from exercised and control animals (Figure 4A). Cell preparations consistently presented high purity, as indicated by immunofluorescence assays for the satellite cell marker Pax7 (see Methods\textsuperscript{47}). After 120 h of growth in culture

![Figure 2](image_url)  
**Figure 2** Body fat content is decreased by endurance exercise, while muscle content is preserved. (A) Body mass changes. (B) Food consumption ($n = 4$, control; $n = 4$, exercised). (C) Dual X-ray absorptiometry scans, representative image. (D) Fat area. (E) Muscle area ($n = 6$, control; $n = 6$, exercised). (F) Muscle mass ($n = 7$, control; $n = 9$, exercised). (G) Representative images of cross-sectional areas. (H) Cross-sectional area quantifications. (I and J) Frequency distribution of the cross-sectional areas of the fibres ($n = 4$, control; $n = 4$, exercised). Scale bar = 100 μm. Asterisks denote statistically significant differences (* $P < 0.05$; ** $P < 0.01$) vs. control.
without passaging, satellite cells were proliferative and at 70–80% confluence. At this time point, the expression of Pax7, MyoD1 (a marker for satellite cell activation), and CXCR4 (a satellite cell membrane marker) were significantly enhanced in cells isolated from exercised animals (Figure 4B). Self-renewal markers Sirt1 (which regulates metabolic state and proliferation) and HIFα were also increased by exercise, as were Spry1, Notch1, and Hey1, which signal myoblast return to quiescence, maintaining the satellite cell pool and long-term muscle integrity. FOXO3a expression (a quiescence marker) and myogenin (a differentiation marker) were not modified (Figure 4B). Overall, this expression profile suggests endurance exercise, promotes activation, but prevents final satellite cell differentiation, maintaining self-renewal or return-to-quiescence phenotypes.

To functionally assess the differentiation of these cells, cell morphology was evaluated (Figure 4C and 4D). Control satellite cells from sedentary animals showed spindle-like elongated morphology, suggestive of early differentiation and loss of stemness, while endurance exercise promoted a more undifferentiated morphology. Despite changes in differentiation markers and morphology, no difference in colony formation ability was observed between exercised and control cells (Figure 4E). Overall, our results suggest that endurance exercise promotes satellite cell self-renewal and quiescence, but inhibits differentiation, without affecting proliferation.
Endurance exercise reduces satellite cell O\textsubscript{2} consumption rates under physiological, but not activated states

There is literature evidence that satellite cell cycle regulation involves changes in energy metabolism, with modulation of substrate use.\textsuperscript{48,51–53} However, mitochondrial oxidative phosphorylation, which is a key regulatory factor in stem cell differentiation,\textsuperscript{30,31} has not been directly assessed in the context of satellite cell modulation. We used a Seahorse Extracellular Flux analyzing system to assess mitochondrial bioenergetics in intact satellite cells isolated from control or endurance-exercised animals (Figure 5A).

A typical trace of real-time OCR measurements is shown in Figure 5B. Endurance exercise promoted overt changes in mitochondrial bioenergetics, with a substantial suppression of basal (physiological) O\textsubscript{2} consumption rates (the initial part of the trace in Figure 5B, quantified in Figure 5C). This was in part due to lower ATP-linked oxygen consumption in exercised cells, or the OCR inhibition promoted by the addition of the ATP synthase inhibitor oligomycin (Figure 5D). Additionally, oligomycin-insensitive O\textsubscript{2} consumption (Figure 5E), which represents the proton leak of the inner membrane, was decreased by exercise. Exercised cells also presented lower maximal respiratory rates, prompted by the addition of the uncoupler 2,4-DNP (Figure 5F) and lower reserve capacity, or the difference between basal and maximal respiration (Figure 5G). However, reserve capacity was still present, indicating that basal respiratory levels were not limited by overall respiratory capacity. Finally, exercised cells presented less non-mitochondrial respiration, measured in the presence of antimycin plus rotenone (Figure 5H), although these OCRs were low compared with overall oxygen consumption. Non-mitochondrial OCRs were subtracted from all other quantifications.

Inhibition of mitochondrial respiration is often accompanied by a shift towards glucose fermentation to lactate. Indeed, when more quiescent, many cells are more fermentative.\textsuperscript{54} However, ECARs (Figure 5I and 5J) in exercised satellite cells were significantly decreased, suggesting that they indeed have lower ATP production, and not a shift towards lactate formation. Lower ATP demands in the exercised

\textbf{Figure 4} Endurance exercise enhances self-renewal and quiescence markers in satellite cells. (A) Schematic panel. (B) Expression of Pax7 (a satellite cell-specific marker that increases upon activation), MyoD1 (an activation marker), CXCR4 (a satellite cell marker), Sirt1 (a self-renewal marker), Spry1 (a return-to-quiescence marker), Notch1, Hey1, Hes1, and HIF1α (self-renewal markers), FOXO3α (quiescence marker), Rbpj (involved in Notch1 signaling), and myogenin (a differentiation marker; \( n = 8 \), control; \( n = 8 \), exercised), relative to HPRT. (C) Representative images of satellite morphology. (D) Quantification of elongated, spindle-like (more differentiated) cells. (E) Myogenic colony formation (\( n = 4 \), control; \( n = 4 \), exercised). Asterisks denote statistically significant differences (\( * P < 0.05 \), \( ** P < 0.01 \), and \( *** P < 0.001 \)) vs. control.
cells are compatible with the expression profile observed previously, suggestive of a higher return to quiescence induced by endurance exercise.

Despite the changes in satellite cell respiratory capacity, exercise did not alter mitochondrial DNA levels relative to nuclear DNA (Figure 5K). This indicates that the changes in respiration are not accompanied by changes in mitochondrial mass. Indeed, the expression of both proteins involved in mitochondrial biogenesis and morphology was not significantly altered by exercise (Figure 5L). Overall, our results show that, despite a lack of modulation of mitochondrial mass, endurance exercise induces a repression of mitochondrial respiration in satellite cells, as well as lower energy demands.

Mitochondrial OCRs were found to be decreased by endurance exercise, an effect associated with a more quiescent expression profile. However, oxidative metabolism is important for stem cell proliferation and activation. Consequently, we evaluated satellite cell metabolism in control or exercised animals after activation induced by injury (Figure 6A). Interestingly, while control OCR values were not significantly altered by injury (compare Figures 5 and 6B–6H), under injured conditions, exercised satellite cell OCRs were higher and equal to those observed in non-exercised cells, under all respiratory states (Figure 6B–6H). ECAR values were also equal between control and exercised-activated satellite cells (Figure 6I and 6J). This demonstrates that the quiescent metabolic state of exercised cells can promptly respond to activating stimuli, with enhanced metabolic activity, as reflected by enhanced mitochondrial ATP production and respiratory capacity. Interestingly, although metabolic activity was equal, the exercised cells showed higher colony formation (Figure 6K), demonstrating that their activated state is more efficient in energy use towards proliferation.

**Inhibition of mitochondrial O$_2$ consumption increases satellite cell self-renewal and quiescence, while decreasing inflammation upon engraftment**

Because endurance exercise promotes mitochondrial respiratory inhibition associated with enhanced return-to-quiescence and self-renewal markers, we verified if the respiratory
inhibition observed was a cause of the changes in expression profile. To do so, we isolated satellite cells from control animals (Figure 7A), which have high OCRs, and titrated low concentrations of the specific mitochondrial respiratory inhibitor antimycin A, to achieve a respiratory inhibition similar to that observed in exercised animals (Figure 7B and 7C). Our titrations showed that 70 ng/mL antimycin A promoted respiratory inhibition similar to that observed in exercised cells. Interestingly, when we promoted this respiratory inhibition in control cells, the expression profile of non-exercised cells became very similar to that observed in exercised cells. Interestingly, when we promoted this respiratory inhibition in control cells, the expression profile of non-exercised cells became very similar to that observed in exercised cells, with an increase in stemness, self-renewal, and return-to-quiescence markers (Figure 7D). This indicates that respiratory suppression is sufficient to promote a phenotype more similar to that of endurance-exercised animals in vitro.

Next, we examined if exercised or respiratory-inhibited satellite cells can lead to changes in responses when transplanted. To do so, isolated cells from control animals treated or not with 70 ng/mL antimycin A, or cells from exercised animals, were transplanted into control animals that had been previously injured to induce engraftment (Figure 8A). Illustrative images of tibialis anterior muscles transplanted with control, exercised, and antimycin-treated satellite cells are shown in Figure 8B. While no overt changes in engraftment were noted between treatment groups, as verified by lack of significant changes in centralized nuclei and eMHC+ myofibres (Figure 8C–8E), both exercised and antimycin-treated cells lead to muscles with significantly lower inflammation, as assessed by cell infiltration (Figure 8F) and cells positive for macrophage marker CD68 (Figure 8G and 8H). This anti-inflammatory effect is most probably related to the enhanced stemness of exercised and respiratory-inhibited satellite cells.

Discussion

Endurance exercise is an often practised form of exercise that significantly increases skeletal muscle energy and oxygen demands, thus changing metabolic regulation in the whole body.55,56 Indeed, it is a well-established intervention to preserve muscle function and prevent changes in energy metabolism, such as insulin resistance, that occur with ageing.57 Despite its widespread use, the cellular and molecular mechanisms involved in metabolic changes promoted by endurance exercise still remain to be fully understood.
Using a mouse model of endurance exercise involving 5 weeks of daily treadmill runs, we found, predictably, that endurance exercise increases physical performance (Figure 1). This occurred in parallel with changes in oxygen consumption at rest and when exercised, as well as changes in exhaustion and recovery glucose levels, confirming exercise induces a shift in whole body metabolism. Indeed, exercised animals were leaner, as indicated by lower fat areas and body mass (Figure 2). Perhaps counterintuitively, endurance exercise did not change muscle areas or fibre cross-sectional areas. These results are in agreement with early work on metabolic effects of exercise, which show that endurance does not promote muscular hypertrophy but leads to enhanced respiratory and lipid oxidation capacity, thus sparing the body from carbohydrate depletion. These early results indicating glucose conservation is a mediator of enhanced stamina in endurance exercise have been confirmed using modern flux analysis techniques. Recently, peroxisome proliferator-activated receptor-δ was found to be a key mediator in the metabolic shift promoted by endurance exercise, increasing palmitate oxidation without an overall change in mitochondrial biogenesis in myocytes.

Figure 7 Inhibition of mitochondrial O2 consumption promotes satellite cell self-renewal. (A) Schematic panel. (B) Typical oxygen consumption rate (OCR) trace in the presence of different concentrations of antimycin A. (B and C) Respiratory inhibition vs. antimycin A concentrations. (D) Satellite cell expression profile after 8 h in the presence of 70 ng/mL antimycin A (n = 8, control; n = 4, antimycin A), relative to HPRT. Asterisks denote statistically significant differences (* P < 0.05, ** P < 0.01, and *** P < 0.001) vs. control.
Although no muscular hypertrophy was observed in exercised animals, prior studies have demonstrated that aerobic exercise changes satellite cell numbers and function, thus improving regenerative response and preserving muscle mass during ageing.17–19,21,22,46 Our work thus grows upon the findings of Joanisse et al.,22 who established that aerobic exercise rescues impaired regeneration during ageing. We now show that this change involves modulation of satellite cell metabolism and self-renewal.

We found that injured muscles in endurance-exercised animals have improved regenerative efficiency, with accelerated muscle repair measured 7 days after injury, as indicated by a higher number of centrally located nuclei and regenerating myofibres expressing eMHC (Figure 3). Notably, endurance exercise training also promoted lower counts of infiltrating cells and cells positive for the macrophage marker CD68, as well as less fibrosis (Figure 3). We should note that our experiments were conducted at specific time points and therefore cannot rule out a change in the time course, but not extent, of the final repair. However, this improved regenerative efficiency or speed may be seminal towards skeletal muscle maintenance in ageing.19,46,61,62

To uncover changes in satellite cells associated with the increased regenerative response capacity observed in endurance exercise, we isolated these cells using the pre-plating method.41 Our isolation protocol comprises the mechanical and enzymatic dissociation of cells, followed by checks, which indicated that cells obtained were consistently highly pure (see Methods). We found that endurance exercise rescues impaired regeneration during ageing. We now show that this change involves modulation of satellite cell metabolism and self-renewal.

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Figure 8 Transplant of exercised or respiratory-inhibited satellite cells prevents inflammation. (A) Schematic panel. (B) Representative images of transplanted fibre cross sections (scale bar = 100 μm). (C) Centrally nucleated (newly formed) myofibre quantification. (D) Representative images of eMHC+ stained fibres. (E) Quantification of eMHC+ fibres. (F) Inflammatory cell quantification. (G) Representative low magnification images, CD68+ (macrophage) staining in green and DAPI in blue. (H) Quantification of CD68+ cells (macrophages; n = 4, control; n = 4, exercised; and n = 4, antimycin A). Scale bars = 100 μm (B) and 50 μm (D and G). Asterisks denote statistically significant differences (**P < 0.01; ***P < 0.001) vs. control animals.
commitment to differentiation.\textsuperscript{31} Furthermore, many studies suggesting a shift from fermentation to oxidative metabolism during differentiation are based on gene expression patterns, while energy metabolism is mostly regulated post-transcription, so gene expression patterns may differ substantially from functional assays.\textsuperscript{67}

As a result, we measured metabolic fluxes in satellite cells isolated from control or endurance-exercised mice. Our results show a clear repression of oxidative metabolism in exercised cells, which present lower mitochondrial oxygen consumption rates under all respiratory states (Figure 5), despite a lack of changes in markers of mitochondrial mass, biogenesis, and dynamics. Myogenin and colony formation were not changed under these conditions (Figure 4), indicating that proliferation was not responsible for mitochondrial changes. The lower oxygen consumption rates are also not attributable to higher rates of fermentation to lactate, as indicated by lower extracellular acidification rates, and are instead related to lower ATP demands. Indeed, respiratory rates in satellite cells isolated from injured muscles were equal in control and exercised cells (Figure 6), showing that exercise did not ablate the ability to respond to injury. Overall, these results suggest that exercise preserves a more quiescent state in unactivated satellite cells, without compromising their activation response. Higher self-renewal and quiescent capacity may be important for responses towards injury, as indicated by the fact that more newly formed fibres were observed in injured exercised animals (Figure 3), and colony formation was more pronounced in satellite cells isolated from exercised animals after injury (Figure 6).

Previous studies have analysed the role of oxidative mitochondrial metabolism in satellite cells. Zhang \textit{et al.}\textsuperscript{53} found that aged satellite cells had lower oxidative phosphorylation. However, this was not associated with enhanced quiescence, but instead senescence, as is perhaps expected in ageing. Indeed, the loss of satellite cell function with ageing could be reversed by recovering NAD levels, in a Sirt1-dependent manner. Consistently, Sirt1 ablation leads to premature differentiation,\textsuperscript{48} a result compatible with our finding that endurance exercise enhances Sirt1 expression, as well as prevents differentiation and promotes self-renewal. Interestingly, caloric restriction, an intervention that leads to enhanced Sirt1 expression in satellite cells,\textsuperscript{51} does not promote other alterations seen in our exercised satellite cells (depressed respiratory rates in the absence of increased mitochondrial DNA), but instead seems to increase oxidative metabolism and enhance mitochondrial biogenesis. This suggests that satellite cell remodelling by exercise and dietary restriction differ mechanistically, although both interventions help preserve muscle mass during ageing.

Because we observed that exercised satellite cells had lower respiration and higher self-renewal, we questioned if the return-to-quiescence characteristics induced by exercise were a consequence of the lower metabolic rates. Indeed, suppression of oxidative phosphorylation often promotes self-renewal in stem cells (reviewed in Khacho and Slack\textsuperscript{36}). To investigate this, we artificially promoted respiratory inhibition in control satellite cells to similar levels of that observed in exercised cells, by adding low quantities of the classic respiratory inhibitor antimycin A. Remarkably, respiratory suppression alone was sufficient to induce an expression panel in satellite cells similar to that promoted by exercise, with enhanced quiescence and self-renewal markers (Figure 7).

We tested next if satellite cells modified by exercise or respiratory modulation could exhibit changes in engraftment after transplant such as those seen in calorically restricted satellite cells.\textsuperscript{51} We should note that our model has the caveat of not allowing the distinction of host vs. donor cells. We found that neither exercised nor respiratory-inhibited satellite cells significantly increased centrally localized nuclei or eMHC\textsuperscript{+} fibres in regenerating myofibres when transplanted into control animals after injury (Figure 8). This suggests that the regenerative potential of exercised cells observed \textit{in vivo} (Figure 3) may depend on characteristics of not only the cells but also their particular niche and thus are not reproduced upon transplantation. However, consistently with their enhanced quiescence markers, both exercised and respiratory-inhibited satellite cell transplants significantly decreased numbers of infiltrating cells and cells expressing the macrophage marker CD68 (Figure 8). This demonstrates that at least part of the beneficial effects of exercise on muscle satellite cell responses can be mimicked by partially suppressing mitochondrial oxidative phosphorylation.

Overall, we demonstrate here that endurance exercise promotes changes in satellite cell function, stemness, self-renewal, and differentiation. The changes are associated with repression of mitochondrial oxygen consumption. Surprisingly, artificial suppression of respiration in satellite cells mirrors the characteristics of exercise. Our study provides insights into mechanisms governing muscle repair promoted by exercise that will hopefully contribute towards better therapeutic interventions preventing sarcopenia.

**Ethics statement**

The authors of this manuscript certify that they comply with the ethical guidelines for authorship and publishing in the \textit{Journal of Cachexia, Sarcopenia and Muscle}.\textsuperscript{68}

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Conflict of interest
None declared.

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Author contributions
Phablo Abreu and Alicia J. Kowaltowski contributed in the conceptualization of the study; Phablo Abreu and Alicia J. Kowaltowski in the data curation; Phablo Abreu in the formal analysis; Phablo Abreu and Alicia J. Kowaltowski in the funding acquisition; Phablo Abreu and Alicia J. Kowaltowski in the methodology; Phablo Abreu and Alicia J. Kowaltowski in the project administration; Alicia J. Kowaltowski in the supervision; Phablo Abreu and Alicia J. Kowaltowski in the writing of the original draft; and Phablo Abreu and Alicia J. Kowaltowski in writing of the review and editing.

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