Acute Skeletal Muscle Contractions Orchestrate Signaling Mechanisms to Trigger Nuclear NFATc1 Shuttling and Epigenetic Histone Modifications

Frank Suhr, Kristina Braun, Mathias Vanmunster, Wilhelm Bloch

Exercise Physiology Research Group, Department of Movement Sciences, Biomedical Sciences Group, KU Leuven, Leuven, Belgium; Institute of Cardiovascular Research and Sport Medicine, Department of Molecular and Cellular Sport Medicine, German Sport University Cologne, Cologne, Germany

Key Words
Ryanodine receptor 1 • Acute exercise • Calcium/calmodulin-dependent protein kinase II • Skeletal muscle • NFATc1 • H3 modifications

Abstract
Background/Aims: Calcium (Ca^{2+}) coordinates skeletal muscle functions by controlling contractions as well as signaling pathways and transcriptional properties. The ryanodine receptor 1 (RyR1), its phosphorylation site (pRyR1Ser2840) and its stabilizers navigate Ca^{2+} oscillations to command muscle signaling cascades and transcriptional activities. While chronic exercise increases pRyR1Ser2840, investigations on acute exercise's effects on RyR1 and Ca^{2+}-dependent modifications of skeletal muscle are rare. The aim of this study was to examine molecular events leading to RyR1 phosphorylation in a physiological model of acute exercise. We hypothesized that exercise-induced RyR1 phosphorylation is associated with altered Ca^{2+}-dependent physiological phenotypes. Methods: We analyzed pRyR1Ser2840, its stabilizers, involved signaling pathways, and Ca^{2+}-sensitive muscle-determining factors (i.e. NFATc1 and epigenetic histone H3 modifications) in rat muscles upon one single running bout of either concentric or eccentric contractions. Results: Both acute exercises significantly increased pRyR1Ser2840 levels in muscles, which was accompanied by dissociations of stabilizers from RyR1. Additionally, RyR1 phosphorylation-inducing signaling cascades PTEN/CaMKII/PKA were significantly activated upon exercise. Further, RyR1 phosphorylations were associated with increased Ca^{2+}-dependent NFATc1 nuclear abundances as well as increased Ca^{2+}-dependent epigenetic H3 acetylations pointing to a pRyR1Ser2840-dependent rapid and novel Ca^{2+} equilibrium upon exercise. Conclusion: Our data report synergistic actions of several distinct pathways to modify RyR1 function to govern physiological phenotypes, here expressed as increased nuclear NFATc1 abundances and epigenetic H3 modifications.
Therefore, we underscore the potential of acute exercise to rapidly change muscle Ca\textsuperscript{2+}-controlling systems and downstream effectors cascades to adjust physiological demands for proper muscle function and phenotype adaptation.

**Introduction**

Coordinated Ca\textsuperscript{2+} oscillations between the sarcoplasmic reticulum (SR) and the sarcoplasm enable physiological skeletal muscle function [1]. As a consequence, chronic alterations in Ca\textsuperscript{2+} oscillations result in reduced muscle contraction and relaxation capabilities [1], are associated with skeletal muscle loss and dysfunction in the elderly [2, 3] and have been reported in patients with skeletal muscle diseases [4–6].

A key mechanism in the skeletal muscle Ca\textsuperscript{2+}-regulating excitation-contraction coupling (ECC) process is the precise regulation of the SR release channel ryanodine receptor isoform 1 (RyR1) [7], which controls SR Ca\textsuperscript{2+} release and thereby maintains physiological cytoplasmic Ca\textsuperscript{2+} concentrations [Ca\textsuperscript{2+}]\textsubscript{cyt} [3, 8]. RyR1 forms a complex associated with its stabilizers, i.e. calstabin-1 and phosphodiesterase isoform 4D3 (PDE4D3), as well as regulating signaling molecules, i.e. the catalytic subunit of protein kinase A (PKAc), protein phosphastase 1 (PP1) or calcium/calmodulin-dependent protein kinase II (CaMKII) [7, 9]. It has been suggested that chronic (predominantly) concentric exercise induces phosphorylation of human RyR1 at Ser\textsuperscript{2843} (Ser\textsuperscript{2844} in mice/Ser\textsuperscript{2840} in rats) and mediates progressive dissociations of RyR1 stabilizers, eventually resulting in persistently ‘leaky’ RyR1 [7, 8]. Chronically ‘leaky’ RyR1s cause uncoordinated Ca\textsuperscript{2+} distributions in myofibers and were proposed to be responsible for reduced skeletal muscle performance in models of muscle diseases [5, 10], in the elderly [3] and after chronic training [8].

We demonstrated that eccentric exercise in humans phosphorylates RyR1 [11]. However, it is unexplored if acute in vivo concentric vs. eccentric contractions induce similar or divergent RyR1 modifications and subsequently alterations in sequences of events leading to RyR1 modifications and dissociations of its stabilizing factors. This question is of considerable importance from basic and clinical points of view, as short-termed exercise is a standard to treat muscle-related diseases [12, 13] and to suppress mechanisms of muscle loss in the elderly [2]. Further, repetitive short-termed exercise potentially govern muscle phenotypes by initiating mitochondrial [14] or myosin heavy chain (MyHC)-related [15] gene activities and can increase muscle strength [16]. All these processes rely on altered Ca\textsuperscript{2+} oscillations. Consequently, studies of effects of acute concentric vs. eccentric contractions are needed to understand mechanisms that control Ca\textsuperscript{2+} oscillations and their physiological consequences in skeletal muscles upon acute exercise to optimize (therapeutic) exercise interventions in clinical settings, elderly people, and athletes. Furthermore, pH\textsubscript{RyR1}-induced changes in Ca\textsuperscript{2+} oscillation influence functionally important muscle regulators, i.e. members of the nuclear factor of activated T cell (NFAT). NFATc1 (NFAT2) is controlled by the Ca\textsuperscript{2+}-sensitive phosphatase calcineurin (Cn), which dephosphorylates NFATc1 upon increasing [Ca\textsuperscript{2+}]\textsubscript{cyt} (resembling altered Ca\textsuperscript{2+} oscillations) to stimulate NFATc1’s nuclear translocation by exposing its nuclear localization sequences (NLS). In the nucleus, NFATc1 determines central characteristics of muscle fibers, e.g. myosin heavy chain expression patterns [15, 17]. Another mechanism enabling muscles to adjust their spatiotemporal demands upon exercise are epigenetic histone modifications [18]. Histone H3 acetylations and phosphorylations, both modifications associated by transcriptional initiation and elongation, depend on Ca\textsuperscript{2+}-sensitive kinases, such as CaMKII [19, 20]. Together, NFATc1 and epigenetic H3 modifications illustrate Ca\textsuperscript{2+}-dependent machineries with abilities to rapidly change functional muscle phenotypes.

We hypothesized (i) that acute muscle contractions evoke RyR1 modifications accompanied by RyR1 stabilizer modifications and allied signaling pathways. We hypothesized (ii) RyR1 modifications trigger rapid changes in Ca\textsuperscript{2+}-dependent machineries, i.e. NFATc1 shuttling and H3 modifications, able to rapidly modify functional muscle phenotypes. To this end, we
analyzed signaling pathways resulting in RyR1 phosphorylation and molecular changes of RyR1 stabilizers as well as Ca\(^{2+}\) oscillations-dependent effectors, i.e. nuclear translocation of NFATc1 and epigenetic H3 modifications in healthy rat muscles upon acute concentric or eccentric running exercise bouts. We report that pRyR1Ser\(^{2840}\) levels increased upon acute exercise in a muscle-independent manner. Additionally, we observed exercise-induced increases of RyR1-phosphorylating kinases PKAc and pCaMKIIThr\(^{286}\), and importantly, dissociations of RyR1 stabilizers PP1, calstabin-1 and PDE4D3 from RyR1. Functionally, we observed increased nuclear NFATc1 abundances and increased transcription-initiating and –elongating H3 acetylations and phosphorylations upon acute exercise, indicating direct and acute functional consequences of pRyR1Ser\(^{2840}\)-induced modifications of [Ca\(^{2+}\)]\(_{\text{cyt}}\).

Materials and Methods

Ethical approval
The performed protocols were approved by the ethical committee of the Landesamt für Natur, Umwelt und Verbraucherschutz of North Rhine-Westphalia, Duesseldorf, Germany (reference number: 8.87-50.10.45.08.188).

Animals
Thirty young female Sprague-Dawley rats (13 weeks old, Charles River Laboratories, Boston, U.S.A.) were assigned randomly and in equal numbers (n=10) to the following three intervention groups: control (CON), one single level running exercise bout at +1° treadmill surface angle (1xLevel), and one single downhill running exercise bout at -20° treadmill surface angle (1xDownhill). No statistical differences were observed between the mean body weights of the animals assigned to the different groups (p>0.05, data not shown). Animals were housed in groups of two in standard vivarium cages on a 12:12 light-dark cycle and had ad libitum access to standard rat chow and water.

Exercise intervention
The CON group remained sedentary and served as age-matched control. The 1xLevel group performed one single level running exercise bout at 20 m*min\(^{-1}\) lasting 15 min at +1° incline, whereas the 1xDownhill group performed one single running exercise bout at 20 m*min\(^{-1}\) at -20° decline lasting 15 min. Former studies of our laboratory proved these running conditions as optimal for rat-based interventions [21, 22]. Running sessions were forced by a motor-driven treadmill (Columbus Instruments, Columbus, OH, USA). Vastus lateralis muscles predominantly perform concentric sarcomere shortening during level running and eccentric sarcomere lengthening during downhill running [23]. Hindlimb muscles, including medial gastrocnemius muscles, experience severe muscle damage during downhill running [24, 25], but not during level running, why we propose that medial gastrocnemius muscles (comparable to vastus lateralis muscles) experience eccentric muscle contractions during downhill running. Additionally, we analyzed animals’ metabolic turnover by assessing oxygen uptake (VO\(_{2}\), Columbus Oxymax Economy System, Columbus Instruments, Columbus, OH, USA) under sedentary and both running conditions, because the proportion of eccentric muscle contractions positively correlates with VO\(_{2}\) due to increased recruitment of motor units [26]. We found significantly increased VO\(_{2}\) values after both running conditions compared to CON and even higher VO\(_{2}\) values upon 1xDownhill compared to 1xLevel indicating increased recruitments of motor neurons and thus eccentric contractions during downhill running [26] (Supplementary Fig. S1 - all supplementary material available online at www.cellphysiolbiochem.com). A motor-driven treadmill (Columbus Instruments, Columbus, OH, USA) coupled to the Oymax Economy System was used to increase running velocities during the running test.

Tissue Sampling and Storage
Sedentary animals of the CON group were decapitated after the two-week acclimatization period. Animals of the 1xLevel and 1xDownhill groups were decapitated (30 min post exercise) upon one single running bouts. Both vastus lateralis (LAT) and medial gastrocnemius (GAS) muscles were dissected from the
hindlimbs. LAT and GAS muscles from the right hindlimbs were snap-frozen in liquid nitrogen and stored at −80°C until further biochemical analysis. LAT and GAS muscle from the left hindlimbs were immersion-fixed in 4% paraformaldehyde (PFA) and stored −80°C until further immunohistochemical analysis.

Transmission electron microscopy
Transmission electron microscopy (TEM) was performed as described earlier [27]. Briefly, skeletal muscle tissue was fixed in 4% PFA, rinsed three times in cacodylate buffer, and then treated with 1% uranyl acetate in 70% ethanol for 8 hrs. After dehydrating the samples in a graded series of ethanol the tissues were embedded in Araldite (Serva). Ultrathin section (60 nm) were processed on an ultramicrotome (Reichert) with a diamond knife and placed on copper grids. TEM was performed with an electron microscope (902A, Carl Zeiss).

Immunohistochemistry and semi-quantitative analysis
The immunohistochemical analyses were performed using standard procedures [11, 28]. Briefly, 4% PFA immersion-fixed 7 μm cryosections were incubated overnight at 4°C with polyclonal primary pRyR1Ser2840 antibody (dilution: 1:100, [11]) Slides were then incubated for 1 hour with appropriate polyclonal biotinylated secondary antibody (diluted 1:400 in 0.05 M Tris-buffered saline, TBS). Afterwards, slides were incubated for 1 hour with streptavidin biotinylated horseradish peroxidase complex (diluted 1:400 in 0.05 M TBS) and stained using a 3, 3’-diaminobenzidine (DAB) solution [11, 28]. LAT and GAS cross sections were stained within a single batch, using the same antibody preparation and the same staining development time to minimize variability in staining efficiency. Digitally captured images (Zeiss Axioshot light microscope coupled to a Sony 3CCD Color Video Camera, 200x magnification) were used to measure RyR1Ser2840-specific staining intensities of each LAT and GAS muscle fibers by selection of the sarcoplasmic region of the myofibers and its subsequent assessment by optical densitometry using a standard software package (ImageJ®, National Institutes of Health, USA) [11]. Intracellular pRyR1Ser2840 level was expressed as mean staining intensity. 200 randomly selected fibers of each LAT and GAS and animal were analyzed, resulting in 2000 fibers per intervention group for each LAT and GAS muscles.

Immunofluorescence
Skeletal muscle tissue was prepared as described in the “Immunohistochemistry and semi-quantitative analysis” section. Mouse α-actinin (dilution 1:250) and rabbit pRyR1Ser2840 (dilution 1:100) primary antibodies were used. Alexa488 goat anti-mouse (dilution 1:500) and Alexa555 goat anti-rabbit (dilution 1:500) fluorescent secondary antibodies were used. Slides were incubated with the respective primary and secondary antibodies for 1 hour at room temperature. DRAQ5 was used to stain skeletal muscle nuclei [28]. Pictures were taken using a Zeiss confocal laser scanning microscope equipped with a Plan-Neofluar 40x/1.3 Oil DIC objective (LSM 510Meta, Zeiss, Jena, Germany). Alexa488 was exited by an Argon laser using the filter set BP505-530. Alexa555 was exited by a Neon laser using the filter set BP595-530. Alexa555 was exited by a Neon laser using the filter set BPS565-615 [28].

Western blot
Upon homogenizing LAT and GAS tissues separately in lysis buffer (Cell Signalling, Frankfurt am Main, Germany), protein concentrations of each homogenate were determined by a protein assay kit (BioRad, Munich, Germany). The general western blot procedure was adapted from laboratory standards [28, 29]. Briefly, 15 µg of total protein were suspended in Laemmli buffer and heated (5 min at 95°C). Linear polyacrylamide gel electrophoresis (PAGE) was performed. Proteins were separated using gels containing either 5% acrylamide (pRyR1Ser2840, and total RyR1), 10% acrylamide (pCaMKIIThr286, pAktSer473, PKAc, PP1, calstabin-1, PDE4D3, GAPDH, and NFATc1) or 15% acrylamide (total H3, H3K9ac, H3K14ac, H3K27ac, and pH3Ser10). Following PAGE, proteins were transferred to polyvinylidene difluoride (PVDF) membranes (Roche, Mannheim, Germany). Membranes were blocked in either 5% low-fat milk (pRyR1Ser2840, RyR1, PKAc, PP1, and calstabin-1) or in 5% bovine serum albumin (pCaMKIIThr286, pAktSer473, PDE4D3, NFATc1, total H3, H3K9ac, H3K14ac, H3K27ac, and pH3Ser10) for 90 minutes and washed in Tris-buffered saline with 0.1% Tween-20 (pH 7.6). Respective membranes were incubated at 4°C overnight with pRyR1Ser2840, total RyR1, PKAc, pCaMKIIThr286, pAktSer473, calstabin-1, PP1, PDE4D3, NFATc1, total H3, H3K9ac, H3K14ac, H3K27ac, and pH3Ser10 and GAPDH antibodies (dilutions see Supplementary Table S1). Membranes were incubated with respective secondary horseradish peroxidase antibodies (dilutions...
see Supplementary Table S1). Proteins were detected by an enhanced chemoluminescence assay (ECL Kit, Amersham-Life Science, Buckinghamshire, UK) and exposed to X-ray film (Kodak X-OMAT Engineering, Eastman Kodak Co., Rochester, NY). Respective bands of each target appeared on X-ray films and were densitometrically measured using Image J (Version 1.43u, National Institutes of Health, Bethesda, USA). GAPDH served as internal loading controls.

**Sarcoplasmic reticulum membrane-enriched microsome preparation**

To investigate RyR1 assembly, we isolated microsomal proteins from tissue homogenates [9, 30, 31]. To this end, SR membrane-enriched microsomes were prepared as described earlier [9]. Briefly, skeletal muscle homogenates were generated as described in the ‘Western blot’ section; however, skeletal muscle tissues were homogenized in 3-(N-morpholino)propanesulfonic acid (MOPS) buffer containing 0.9% NaCl, 10 mM MOPS (pH 7.0), and 1 mM PMSF. Upon homogenizing, solutions were centrifuged at 4, 000*g for 15 minutes at 4°C (Eppendorf Centrifuge 5417C, Hamburg, Germany). The supernatant was then collected and further centrifuged at 100, 000*g for 90 minutes at 4°C (Optima TLX Ultracentrifuge equipped with a TLA-55 rotor; Beckman Coulter, Krefeld, Germany). The SR membrane-enriched microsome pellets were resuspended in a buffer containing 0.9% NaCl, 0.3 M sucrose, and 1 mM PMSF [31]. Protein concentrations of pellets and supernatants, respectively, were determined by a protein assay kit (BioRad, Munich, Germany). Relative protein amounts of PKAc, PP1, calstabin-1, and PDE4D3 were measured in pellets and supernatants, respectively, as described in the “Western blot” section.

**Nuclear and cytoplasmic fraction isolation from muscle tissue**

The protocol was described earlier [32] and slightly modified. Briefly, 100 mg of each LAT and GAS muscle of each condition (CON, 1xLev, 1xDh) were solubilized first in “homogenization” buffer (25 mM HEPES, pH 7.5, 0.5 mM EDTA, 0.5 mM EGTA, 25 mM NaF, 1 mM Na$_3$VO$_4$, 1 mM DTT, protease inhibitor [Sigma-Aldrich, Munich, Germany]). After incubating the homogenate for 10 min on ice with regular vortexing, the homogenate was centrifuged for 15 min at 4°C at 22, 000*g. The sarcoplasmic fraction-containing supernatant was stored in a new tubes at -80°C until further analysis, whereas the pellet was washed in “washing” buffer (10 mM HEPES, pH 7.5, 1.5 mM MgCl$_2$, 10 mM KCl, 25 mM NaF, 1 mM Na$_3$VO$_4$, 1 mM DTT, protease inhibitor [Sigma-Aldrich, Munich, Germany]). After centrifugation at 3, 000*g for 10 min at 4°C, the supernatant was discarded and the pellet lysed in “solubilization” buffer (20 mM HEPES, pH 7.5, 0.42 M NaCl, 1.5 mM MgCl$_2$, 0.2 mM EDTA, 1% Triton-X 100, 25 mM NaF, 1 mM Na$_3$VO$_4$, 1 mM DTT, protease inhibitor [Sigma-Aldrich, Munich, Germany]). The homogenized pellet was stored on ice for 30 min with regular vortexing and centrifuged at 22, 000*g for 20 min at 4°C. The nuclear fraction-containing fraction was stored in a new tube at -80°C until further analysis. Nuclear fractions were identified by total histone H3, whereas sarcoplasmic fractions were identified by GAPDH.

**Statistics**

**Immunohistochemistry and western blot.** All data are presented as mean ± S.E.M. Experimental values of the CON groups were used as reference values. One-way analysis of variances (ANOVA) on ‘exercise condition’ was performed using STATISTICA software package (Statistica for Windows 7.0, Tulsa, USA). If ANOVA was significant, individual differences between tested conditions were analyzed by applying a post hoc test (Duncan’s multiple comparison test). Statistical significance was considered for p<0.05. To compare PKAc, PP1, calstabin-1, and PDE4D3 values in SR membrane-enriched microsome pellets and supernatants of the respective interventions, paired t-tests were used. Statistical significance was considered for p<0.05.

**Animal data.** To evaluate statistical differences between the investigated groups CON, 1xLevel, and 1xDowhill regarding body weight at the beginning and the end of the study, one-way ANOVA with Duncan’s multiple comparison test was conducted. Statistical significance was considered for p<0.05.

**Results**

**Eccentric exercise alters sarcomere ultrastructure**

First, we analyzed the association between downhill running and predominantly eccentric muscle contractions by transmission electron microscopy (TEM). We did not
observe any disruptions of sarcomeric Z-bands after 1xDownhill compared to 1xLevel and CON conditions in either LAT muscles (Fig. 1Aa-c) or GAS muscles (Fig. 1Ad-f), whereas we instead found significantly stretched sarcomeres in LAT (Fig. 1B) and GAS (Fig. 1C) compared to 1xLevel and CON. In addition, we observed irregular M-bands within sarcomeres of 1xDownhill LAT (Fig. 1Ac) and GAS (Fig. 1Af) muscles, which were absent upon CON (Fig. 1Aa, d) and 1xLevel (Fig. 1Ab, e). These data indicate that a single bout of 15 min eccentric running induces higher muscle strain than 15 min of concentric running.

**Fig. 1.** Transmission electron microscopy (TEM) analysis of concentrically and eccentrically stressed muscle. (A) TEM reveals increased skeletal muscle M-band dilations (see arrow heads) after 1xDownhill compared to 1xLevel and CON conditions, which showed normal M-bands (arrows). (B, C) Quantitative analysis of sarcomeric M-band dimensions in LAT and GAS muscles and the three intervention groups demonstrates significantly increased M-band dimensions upon acute eccentric running. *** p<0.001, bar is 1 µm. Muscles from n=8 rats of each group were analyzed.
Fig. 2. Immunohistochemical analysis of pRyR1Ser$_{2840}$ in rat LAT and GAS muscles. (A and D) Double staining against (a) α-actinin and (b) pRyR1Ser$_{2840}$ shows localizations of Z-disks (α-actinin) in close vicinity of junctional sarcoplasmic reticuli (pRyR1Ser$_{2840}$) that surround t-tubuli, which are located in the transitional zone of A- and I-bands in skeletal muscles. The merged picture is demonstrated in (c). Fold changes of pRyR1Ser$_{2840}$ in LAT (B) and GAS (E) muscle under 1xLevel and 1xDownhill conditions compared to CON conditions. pRyR1Ser$_{2840}$ shows significantly elevated levels in both muscles after acute exercise stimulations. Representative pRyR1Ser$_{2840}$ staining in LAT (C) and GAS (F) muscle under CON, 1xConc, and 1xEcc conditions. The upper panels depict the pRyR1Ser$_{2840}$ pattern; the lower panel shows control slides that were incubated with 0.8% BSA instead of primary antibody. ** p<0.01, *** p<0.001. Bar in (A and D) is 5 µm, bar in (C and F) is 50 µm. In total, n=2000 muscle fibers were measured per intervention group.
Acute exercise results in increased pRyR1Ser^{2840} levels

Confocal laser scanning microscopy of longitudinal LAT and GAS skeletal muscle fibers showed localizations of Z-disks in close vicinity to junctional SRs that surround t-tubuli, which located to the transitional zone of A- and I-bands in skeletal muscles containing RyR1 (Fig. 2A, B).

**LAT muscle.** To test whether one single running exercise stimulus induces RyR1 phosphorylation, pRyR1Ser^{2840} was quantified in LAT muscle subjected to 15 minutes of exercise by semi-quantitative immunohistochemistry. Compared to CON condition, both exercise protocols (1xLevel and 1xDownhill) resulted in significantly increased phosphorylation at Ser^{2840} of RyR1 with no difference between the two exercised groups (Fig. 2C, Supplementary Table S2). Representative immunohistochemical staining of LAT fibers are shown in Fig. 2D.

**GAS muscle.** Compared to the CON condition, both exercise conditions resulted in significantly increased phosphorylations of RyR1 at Ser^{2840} (Fig. 2E, Supplementary Table S2). In contrast to LAT muscle, the 1xLevel rat GAS displayed significantly higher pRyR1Ser^{2840} values compared to the 1xDownhill condition (Fig. 2E, Supplementary Table S2). Representative immunohistochemical staining of GAS fibers are shown in Fig. 2F. We further used western blot to proof our histological pRyR1Ser^{2840} results.

**LAT muscle.** The results confirm significant increases of pRyR1Ser^{2840} in both 1xLevel and 1xDownhill conditions compared to CON (Fig. 3A). Whereas these results are in general agreement with the observations obtained by immunohistochemical staining, western blot quantifications revealed significantly higher pRyR1Ser^{2840} values in 1xLevel compared to 1xDownhill (Fig. 3A). Representative western blot bands for pRyR1Ser^{2840} are depicted in Fig. 3I (upper panel). For quantification of relative pRyR1Ser^{2840} amounts, we measured total RyR1 levels (Fig. 3I). Since total RyR1 western blot analyses did not show any differences between the tested conditions, we normalized pRyR1Ser^{2840} to total RyR1.

**GAS muscle.** Western blot analysis of 1xLevel exercise showed significantly increased values of pRyR1Ser^{2840} compared to CON (Fig. 3B). Differences for 1xDownhill condition were barely below significance (Fig. 3B). These results support our observations from immunohistochemical staining (Fig. 2E). Representative western blots for pRyR1Ser^{2840} normalized to total RyR1are depicted in Fig. 3I.

**RyR1-phosphorylating kinases change their activation state upon acute exercise**

We further studied mechanisms that contribute to the observed pRyR1Ser^{2840} patterns upon acute exercise. We analyzed signaling molecules that are involved in RyR1 phosphorylation including pCaMKIIThr^{286}, pAktSer^{473}, and pPTENSer^{380}.

**Phosphorylated calcium/calmodulin-dependent protein kinase IIThr^{286}.** pCaMKIIThr^{286} is involved in the phosphorylation of RyR in heart muscle [33]. In LAT muscle we observed a pCaMKIIThr^{286} signal opposing the pRyR1Ser^{2840} pattern. Both 1xLevel and 1xDownhill exercises led to significantly increased pCaMKIIThr^{286} signals in LAT muscles (Fig. 3C, I) compared to CON, whereas 1xDownhill caused even higher pCaMKIIThr^{286} levels (Fig. 3C, I) than 1xLevel. In GAS muscle, the pCaMKIIThr^{286} signal significantly increased in both exercise groups (Fig. 3D, I) compared to CON. In contrast to LAT muscle, pCaMKIIThr^{286} signal showed highest levels upon 1xLevel (Fig. 3D, I).

**Phosphorylated protein kinase B/AktSer^{473}.** pAktSer^{473} directly interacts with CaMKII [34, 35], thus we measured pAktSer^{473} as a marker to confirm the pCaMKIIThr^{286} signals. In LAT muscle, pAkt^{473} levels corroborated with pCaMKIIThr^{286} results. Both 1xLevel and 1xDownhill conditions led to significantly increased phosphorylations at Ser^{473} (Fig. 3E, I) compared to CON with highest phosphorylation in the 1xDownhill group, which was significantly higher (Fig. 3E, I) compared to 1xLevel group. In GAS muscle, pAktSer^{473} levels increased upon 1xLevel (Fig. 3F, I), but decreased upon 1xDownhill compared to CON and 1xLevel (Fig. 3F, I). Consequently, pAktSer^{473} was highest upon 1xLevel stimulation (Fig. 3F, I), confirming pCaMKIIThr^{286} signals.
**Fig. 3.** Western blot analysis of RyR1 and signaling molecules in rat LAT and GAS muscles. Fold changes of pRyR1Ser<sup>2840</sup> in LAT (A) and GAS (B) muscle upon 1xLevel and 1xDownhill compared to CON conditions show significantly elevated levels after acute exercise stimulations. Fold changes of pCaMKIIThr<sup>286</sup> levels in LAT (C) and GAS (D) compared to CON show increased levels after acute exercise stimulations. Fold changes of pAktSer<sup>473</sup> levels in LAT (E) and GAS (F) compared to CON. Fold changes of pPTENSer<sup>380</sup> levels in LAT (G) and GAS (H) compared to CON show increased levels after acute exercise stimulations. Representative western blots of pRyR1Ser<sup>2840</sup>, total RyR1, pCaMKIIThr<sup>286</sup>, pAktSer<sup>473</sup>, PI3K, and pPTENSer<sup>380</sup> as well as the internal loading control GAPDH are depicted in (I). *p<0.05, **p<0.01. n(western blot experiments) = 8 for each tested condition.
Phosphorylated phosphatase and tensin homolog deleted from chromosome 10Ser\textsuperscript{380}.

To find a mechanistic explanation for the differences in pCaMKIIThr\textsuperscript{286} and pAktSer\textsuperscript{473} distributions between LAT and GAS muscles, we next focused on PTEN, which inhibits the kinase function of PI3K \cite{36}. Consequently, phosphorylations of kinases downstream of PI3K, including CaMKII and Akt, are inhibited by activated PTEN. Phosphorylation at Ser\textsuperscript{380} directs PTEN away from the plasma membrane towards the cytosol, hence, resulting in a translocation-dependent inactivation of PTEN that permits phosphorylations of downstream kinases \cite{36}. We found pPTENSer\textsuperscript{380} signals to correlate with observed pCaMKIIThr\textsuperscript{286} and pAktSer\textsuperscript{473} signals in both LAT and GAS muscles and under both exercise conditions (Fig. 3G, H, I). In agreement with pCaMKIIThr\textsuperscript{286} and pAktSer\textsuperscript{473} signals, the pPTENSer\textsuperscript{380} levels differed between LAT and GAS muscle (Fig. 3G, H, I), indicating that PTEN phosphorylation at Ser\textsuperscript{380} is most likely responsible for the Akt/CaMKII-mediated RyR1 phosphorylation.

RyR1 stabilizers dissociate from the RyR1 complex upon acute exercise

We measured PKAc associated to RyR1 and acute exercise-induced dissociations of the stabilizers calstabin-1, PDE4D3, and PP1 from RyR1 in SR membrane-enriched microsome pellets and supernatants. The stabilizers prevent RyR1 complexes from ‘leaky’ conditions making calstabin-1, PP1, and PDE4D3 critical players for RyR1-mediated Ca\textsuperscript{2+} homeostasis in skeletal muscle \cite{8}.

Catalytic subunit of protein kinase A. Besides CaMKII, protein kinase A (PKA) is a major kinase involved in RyR1 phosphorylation \cite{9} and is activated by physical exercise \cite{37, 38}. We quantified its catalytic subunit PKAc \cite{9} in whole muscle homogenates. As demonstrated in Fig. 4A, we observed a significant increase of PKac in both LAT and GAS muscle upon either exercise mode compared to CON, indicating that PKA is activated upon exercise and thus likely contributes to RyR1 phosphorylation. A direct interaction between PKAc and RyR1 is necessary to transfer the respective phosphate group. Thus, we went on to quantify PKAc in pellets (P) and supernatants (S) \cite{9} to test whether acute exercise induced enhanced associations between RyR1 and PKAc. We did not observe any differences between PKAc(P) and PKAc(S) in both muscles (Fig. 4B, F) at CON conditions, indicative of equal PKAc distributions. However, we measured slight increases of PKAc(P) that were just below significance (Fig. 4B, F), but reached statistical significance in GAS muscle upon 1xDownhill condition (Fig. 4B, F). Together, our results indicate that acute exercise increased PKAc associated with RyR1 in SR membrane-enriched microsomes to trigger RyR1 phosphorylation.

RyR1 stabilizers. To prove our hypothesis that acute exercise results in dissociations of stabilizing molecules from RyR1, we analyzed protein levels of calstabin-1, PDE4D3, and PP1 from RyR1 in SR membrane-enriched microsome pellets and supernatants. The stabilizers prevent RyR1 complexes from ‘leaky’ conditions making calstabin-1, PP1, and PDE4D3 critical players for RyR1-mediated Ca\textsuperscript{2+} homeostasis in skeletal muscle \cite{8}.

RyR1 stabilizers dissociate from the RyR1 complex upon acute exercise

Acute exercise causes increased nuclear NFATc1 and epigenetic histone H3 marks

Because we observed significant RyR1 phosphorylation upon acute exercise, we tested the hypothesis that these acute exercise-induced alterations exert functional consequences.
Pellet Supernatant

LAT(P)

LAT

LAT

LAT(G)

LAT(G)

LAT(G)

LAT(G)

LAT(G)

LAT(G)

LAT(G)

LAT(G)

LAT(S)

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**Fig. 5.** Nuclear shuttling of NFATc1 and epigenetic histone H3 modifications upon acute exercise. (A) Confirmation of nuclear and sarcoplasmic fractions in skeletal muscles. We performed western blot for a nuclear marker protein (total H3) and a sarcoplasmic marker protein (GAPDH), and whole lysate, sarcoplasmic fraction, and nuclear fractions were loaded. Total H3 predominantly shows a signal in nuclear fractions, whereas GAPDH was predominantly detected in sarcoplasmic fractions. (B) NFATc1 was detected in sarcoplasmic and nuclear fractions of CON, 1xLevel, and 1xDh LAT and GAS muscles. As shown, NFATc1 abundances increased in nuclear fractions of 1xLevel and 1xDownhill compared to CON, whereas the sarcoplasmic fractions remained unchanged. The western blot shows results from LAT muscles, but is representative for GAS muscles (not shown). (C) The epigenetic histone H3 marks H3K9ac, H3K14ac, H3K27ac, and pH3Ser^{10} were analyzed in nuclear fractions of LAT (upper panel) and GAS (lower panel) muscles. As shown, H3 acetylations increase with both 1xLevel and 1xDownhill, whereas pH3Ser^{10} remains unchanged. n(western blot experiments) = 8 for each tested condition.
To this end, we studied the abundance of NFATc1 and histone H3 modifications between sarcoplasmic and nuclear fractions. NFATc1 is Ca\(^{2+}\)-sensitive and increased \([\text{Ca}^{2+}]_{\text{cyt}}\) rapidly triggers NFATc1 translocations into nuclei [39]. This translocation can have functional consequences, because NFATc1 acts as a Myh7 gene-activating transcription factor [40] determining oxidative muscle fiber phenotypes.

As shown in Fig. 5A, we detected total H3 predominantly in the nuclear fraction, whereas we detected GAPDH predominantly in the sarcoplasmic fractions. We studied NFATc1 abundances in nuclear and sarcoplasmic fractions and found that sarcoplasmic NFATc1 levels remained unchanged upon acute exercises, whereas NFATc1 levels significantly increased in nuclear fractions upon both acute exercise interventions (Fig. 5B).

We further studied whether acute exercise modifies epigenetic histone H3 marks, which depend on changes in \([\text{Ca}^{2+}]_{\text{cyt}}\) [18, 19]. We focused on transcription- and elongation-initiating H3 acetylations at lysine 9 (H3K9ac), lysine 14 (H3K14ac), lysine 27 (H3K27ac) and phosphorylation at Ser\(^{10}\) (pH3Ser\(^{10}\)). In the nuclear fractions of LAT muscles, we observed that H3K9ac, H3K14ac, and H3K27ac were significantly increased upon both exercise programs, whereas pH3Ser\(^{10}\) was not changed (Fig. 5C). In nuclear fractions of GAS muscle, we found that predominantly acute eccentric muscle contractions resulted in significantly increased H3K9ac, H3K14ac, and H3K27ac levels. pH3Ser\(^{10}\) was also unchanged in GAS muscles (Fig. 5C).

**Discussion**

We hypothesized in this work that acute muscle contractions alter RyR1 complex assembly resulting in rapid changes of muscle phenotype-controlling machineries. We indeed found that acute muscle contractions trigger RyR1 phosphorylations that are accompanied by RyR1 stabilizer dissociations and involved signaling cascade activations. These events alter \([\text{Ca}^{2+}]_{\text{cyt}}\), which is the driving force for the observed increased NFATc1 nuclear shuttling and epigenetic H3 modifications.

Ca\(^{2+}\) oscillations play critical roles in skeletal muscle signaling, performance or weakness, because of the potential to control signaling pathways and nuclear transcription factors [41]. Detailed knowledge about early acute muscle contraction-induced modifications of RyR1 is important to understand whether acute exercise changes the local muscle Ca\(^{2+}\) homeostasis [42, 43]. This is of interest, because repetitive, but short-term acute exercise is favored to treat muscle disease-related pathologies [12], to induce metabolic improvements in muscle [44] and is advantageous over chronic exercise regimes due to longer regeneration periods [45]. Mechanistically, the analysis of molecular processes is preferable upon acute stimulations rather than long-term exercise regimes as unwanted secondary side effects can be minimized.

We demonstrate herein that acute exercise leads to increased pRyR1Ser\(^{2840}\) levels in rat LAT and GAS muscles upon both predominantly concentric (level) and predominantly eccentric (downhill) contractions. Interestingly, acute level and downhill exercises caused distinct pRyR1Ser\(^{2840}\) levels. This could relate to the former finding of activity-induced increases in [ATP] to activate RyR1 in both Ca\(^{2+}\)-dependent and –independent manners [46, 47]. But to gain more detailed insights into the observed pRyR1Ser\(^{2840}\) patterns following acute exercise, we studied different molecular pathways that regulate RyR1, including RyR1 stabilizers and RyR1-phosphorylating signaling cascades [7, 48]. We first focused on pCaMKIIThr\(^{286}\), because this kinase mediates RyR phosphorylation [33, 49] and acute exercise increases pCaMKIIThr\(^{286}\) levels as fast as a few seconds lasting up to several minutes upon stimulation [50]; however, associations with RyR1 phosphorylation are unexplored. Our data support the findings of early CaMKII phosphorylation and suggest a role of CaMKII as an upstream regulator of RyR1 in muscles. Whereas RyR1 phosphorylation was highest upon 1xLevel in both LAT and GAS muscles, we found highest pCaMKIIThr\(^{286}\) levels upon 1xDownhill in LAT, and upon 1xLevel stimulation in GAS. These results show the contribution
of additional pathways to synergistically phosphorylate RyR1 with a dependence on contraction mode and muscle. This argument is supported by the pAktSer⁴⁷³ pattern that shows exercise mode- and muscle-dependent differences that mimic pCaMKIIThr²⁸⁶ levels indicating a direct CaMKII and Akt interaction axis [34, 35].

Due to the muscle- and exercise type-specific CaMKIIThr²⁸⁶ regulation, we further tested if the pCaMKIIThr²⁸⁶-regulating PTEN system was also modulated in a muscle-specific and exercise type-dependent manner. We indeed found muscle- and exercise-dependent differences of pPTENSer³⁸⁰ levels. Phosphorylation of PTEN at Ser³⁸⁰ leads to its inactivation by guiding PTEN to the sarcoplasm, where it is unable to interfere with the PI3K signaling [36]. The observed pPTENSer³⁸⁰ levels confirm muscle- and exercise type-specific pCaMKIIThr²⁸⁶ and pAktSer⁴⁷³ patterns indicating the involvement of interacting upstream mechanisms, including kinases and their inhibitor regulations, to phosphorylate RyR1 upon acute exercise.

The holoenzyme PKA is activated by cAMP, which rapidly increases with exercise [37, 38] and PKA further controls RyR1 phosphorylation [9]. We observed increases of the activated catalytic PKA (PKAc) subunit and direct PKAc associations with RyR1 upon both acute exercise regimes. Thus, we show that PKA is rapidly activated by acute exercise and we identified an additional component contributing to RyR1 phosphorylation. This highlights that RyR1 is controlled by numerous means. Because the phosphorylation of RyR1 is a dynamic process, PKA might not be tethered to RyR1 permanently, which would explain its increased, but transient association to RyR1.

Unlike PKA, the RyR1 stabilizers calstabin-1, PDE4D3, and PP1 [7] form stable aggregates with RyR1, whereas their chronic dissociations from RyR1 result in uncoordinated Ca²⁺ oscillations and consequently impaired skeletal muscle functions [8]. We found significant dissociations of calstabin-1, PDE4D3, and PP1 from RyR1-containing microsomes demonstrating physiological impacts of acute exercise on molecular Ca²⁺-regulating components. The consequence of rapid [Ca²⁺]cyt changes upon acute exercise are unclear, but could trigger muscle-determining factors.

One such muscle-determining and Ca²⁺-sensitive player is NFATc1 that has a critical function as transcription factor in oxidative muscle fiber phenotyping [17, 40, 51]. We found that NFATc1 rapidly increased its nuclear abundance upon acute exercise. This confirms earlier reports, which found that acute Ca²⁺ sparks rapidly dephosphorylate NFAT isoforms to expose their nuclear localization sequences (NLS) to stimulate nuclear NFAT imports within minutes [52, 53]. This general finding was substantiated in primary muscle cells, which were electrically stimulated and showed increased NFATc1 abundances in myonuclei within minutes [39], meaning that NFATc1 remarkable senses dynamic changes in [Ca²⁺]cyt and frequencies of Ca²⁺ oscillations. This finding of rapid pRyR1Ser²⁸⁴₀-induced NFATc1 translocations is important, because it presents a physiological explanation through which mechanisms repetitive short-term exercise potentially direct skeletal muscle phenotyping. Further, NFATc1 controls cell size and intra-muscular nuclei retaining [54]. This, in combination with the MyHC gene control, demonstrates that NFATc1 is responsible for the activation of mechanisms that acutely navigate muscle repair mechanisms upon acute muscle damaging exercise, such as eccentric contractions. The mechanistic background for this physiological result is a newly established Ca²⁺ equilibrium forcing nuclear NFATc1 abundances.

A second muscle-determining and Ca²⁺-sensitive mechanism mediates epigenetic H3 modifications. Specifically, acute (exercise-induced) changes of [Ca²⁺]cyt trigger acetylations of H3 at lysine 9 (H3K9ac), lysine 14 (H3K14ac) and lysine 27 (H3K27ac) [18, 19] as well as phosphorylations of H3 (pH3Ser¹⁹) [20]. These epigenetic H3 modifications can be mediated by Ca²⁺-sensitive kinases, including CaMKII, and are associated with transcription initiation and elongation [19]. We found increased H3K9ac, H3K14ac, and H3K27ac levels upon both acute level and downhill running in both muscles. It was demonstrated in cardiac muscles that changes in Ca²⁺ oscillations result in similar H3 acetylations, which caused alternative splicing of cardiac muscle-regulating genes [19]. Another study in humans found that
acute exercise increases defined H3 acetylation sites, which was accompanied by increased 
\[\text{pCaMKIIThr}^{\text{386}}\] levels [18]. Acute exercise did not change pH3Ser\(^{10}\) in LAT and tended to 
increase this modification in GAS; however, we did not identify significant values, which was 
surprising, because H3 acetylations relate to pH3Ser\(^{10}\), as H3 acetylations are effectively 
preceded by H3 phosphorylations at Ser\(^{10}\) in \textit{in vitro} systems [20, 55]. Our data indicate 
that exercise-induced H3 acetylations do not depend on H3 phosphorylations in skeletal 
muscles under physiological \textit{in vivo} exercise conditions. Overall, our findings suggest that 
muscles subjected to acute exercise activate epigenetic machineries to rapidly stimulate 
transcriptional initiation and elongation to adapt to the new demands and environmental 
stress. The transcriptional initiation and elongation might also be linked to muscle repair 
mechanisms upon acute muscle-damaging exercise. It was demonstrated that epigenetic H3 
modifications, including the herein investigated, contribute the maintenance of functional 
muscle phenotypes upon damage [56, 57]. The basic concept about how acute exercise 
determines RyR1 phosphorylation, its regulating kinases, its stabilizers and functional Ca\(^{2+}\)- 
dependent Ca\(^{2+}\)-sensitive NFATc1 and increased [Ca\(^{2+}\)]\(_{\text{cyt}}\)-sensitive epigenetic H3 modifications,

We further observed that eccentric exercise caused lower pRyR1Ser\(^{2860}\) levels compared 
to concentric exercise. This was particularly unexpected, because muscles experience 
higher mechanical strain (see Fig. 1) and a higher number of fibers is innervated as 
reflected by increased VO\(_2\) values during eccentric compared to concentric muscle work (see Supplementary Fig. S1) [58, 59]. Our data cannot explain this phenomenon in detail, 
but we suggest that lower maximum force peaks combined with increased passive forces 
induced by intra-sarcomeric protein interactions [60] to attenuate activation of Ca\(^{2+}\)- 
regulating components, including RyR1, during eccentric work. Exposing skeletal muscles to 
higher eccentric stresses (with the same peak force as in concentric exercise) could test this 
assumption. Furthermore, we measured a sarcomere length of about 3.2 µm upon eccentric 
exercise. This enhanced sarcomere length still allows for active contractions, which are 
supported by passive forces as demonstrated in isolated muscle fibers [60, 61] wherefore 
it might reflect physiological sarcomere length upon acute eccentric muscle contractions \textit{in vivo}. \textit{In vitro} single muscle fiber studies revealed that increased muscle force correlates with 
acutely increased RyR1 phosphorylation [16], which is in contrast to chronic training [8]. 
Transferred to our results, it is plausible that acute \textit{in vivo} muscle loading acutely increased 
muscle forces due to transient, but not chronic RyR1phosphorylations. This assumption is 
supported by data from our group showing increased passive muscle forces in cardiomyocytes 
upon acute exercise [62]. This has a practical relevance in the way that molecular cascades 
finally trigger physiological outcomes, here transiently increased muscle force. Recently, 
it was demonstrated that strenuous acute exercise potentially causes RyR1 fragmentations that 
consequently disturb the Ca\(^{2+}\) homeostasis and related muscle contraction mechanisms [63]. 
However, we did not observe any comparable RyR1 fragmentations reflecting physiologically 
tolerable loading by the applied exercise programs and hence rather physiological than 
disturbed Ca\(^{2+}\) oscillations. A pathological condition of altered Ca\(^{2+}\) is inflammation. We can 
connect our results to inflammation in the way acute inflammation as expressed by increases 
of the pro-inflammatory cytokine IL-1 negatively controls [Ca\(^{2+}\)]\(_{\text{cyt}}\)-sensitive NFATc1 and increased [Ca\(^{2+}\)]\(_{\text{cyt}}\)-sensitive epigenetic H3 modifications,
Fig. 6. Summary of RyR1 phosphorylation-triggered signaling and mediated NFATc1 and epigenetic histone H3 modifications induced by acute exercise. (A) Under resting conditions, the ATP metabolism is low as is the cAMP concentration. Therefore, only small amounts of the apoenzyme PKA are cleaved into its active catalytic isoform, PKAc. Subsequently, RyR1 phosphorylation is low, but not absent, and the RyR1 stabilizing molecules, calstabin-1, PDE4D3, and PP1, are associated to RyR1. In parallel, resting conditions result in low inactive (phosphorylated) forms of PTEN. Likewise, there is little activation of Akt at Ser473 and, thus, of CaMKII at Thr286. This results in the regular oscillation of Ca²⁺ from RyR1, similar to what occurs in the event of low PKAc. The consequences are sarcoplasmic NFATc1 localizations as well as low epigenetic H3 acetylations/phosphorylations. (B) Acute in vivo exercise induces high ATP metabolism in working skeletal muscles resulting in increased cAMP levels. This event activates the apoenzyme PKA to be cleaved into its catalytic isoform, PKAc. Subsequently, RyR1 phosphorylation increases and its stabilizing molecules, calstabin-1, PDE4D3, and PP1, show transient dissociations from RyR1. In parallel, acute exercise leads to increased levels of the inactive (phosphorylated) form of PTEN, enabling Akt to be PI3K-phosphorylated at Ser473 (bold). Subsequently, CaMKII is activated at Thr286, which might amplify the PKAc-mediated RyR1 phosphorylation. Consequently, NFATc1 is dephosphorylated and translocates into the nucleus and H3 is epigenetically modified by specific acetylations. Synergistically, these events control skeletal muscle properties and molecular phenotypes even under conditions of acute and short-term muscle work.
which are associated with increased transcriptional initiation and elongation. These findings demonstrate that acute exercise rapidly evokes critical physiological consequences making acute exercise bouts a crucial determinant that orchestrates of muscle function and properties.

**Conclusion**

Proper Ca$^{2+}$ oscillations are essential for physiological skeletal muscle properties [7]. Detailed knowledge about mechanisms controlling Ca$^{2+}$ oscillations and consequences of altered Ca$^{2+}$ oscillations is hence critical to understand muscle function. Our data show that acute exercise phosphorylates the Ca$^{2+}$ oscillation-controlling RyR1 channel. We further identify different signaling pathways that concertedly phosphorylate RyR1 and thus likely control Ca$^{2+}$ oscillations. The action of involved signals is amplified by dissociations of calstabin-1, PDE4D3, and PP1 from RyR1 complexes causing rapidly modified RyR1. The physiological consequences of these acute exercise-induced RyR1 modifications are (i) enhanced nuclear abundances of the muscle phenotype-controlling transcription factor NFATc1 and (ii) epigenetic modifications of the histone H3 associated with transcriptional initiation and elongation. Consequently, we demonstrate for the first time that physiological and acute exercise bouts evoke alterations of the local muscle milieu to determine novel functional equilibria expressed by nuclear NFATc1 and H3 modifications.

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**Disclosure Statement**

The authors declare that they have no competing interests.

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