Mitochondrial Hyperfusion during Oxidative Stress Is Coupled to a Dysregulation in Calcium Handling within a C2C12 Cell Model

Calum J. Redpath1*, Maroun Bou Khalil1, Gregory Drozdzal1, Milica Radisic2, Heidi M. McBride3

1 Cellular Electrophysiology Laboratory, University of Ottawa Heart Institute, University of Ottawa, Ottawa, ON, Canada, 2Mining Building, Institute of Biomaterials and Biomedical Engineering, University of Toronto, Toronto, ON, Canada, 3Montreal Neurological Institute, McGill University, Montreal, QC, Canada

Abstract

Atrial Fibrillation is the most common sustained cardiac arrhythmia worldwide harming millions of people every year. Atrial Fibrillation (AF) abruptly induces rapid conduction between atrial myocytes which is associated with oxidative stress and abnormal calcium handling. Unfortunately this new equilibrium promotes perpetuation of the arrhythmia. Recently, in addition to being the major source of oxidative stress within cells, mitochondria have been observed to fuse, forming mitochondrial networks and attach to intracellular calcium stores in response to cellular stress. We sought to identify a potential role for rapid stimulation, oxidative stress and mitochondrial hyperfusion in acute changes to myocyte calcium handling. In addition we hoped to link altered calcium handling to increased sarcoplasmic reticulum (SR)-mitochondrial contacts, the so-called mitochondrial associated membrane (MAM). We selected the C2C12 murine myotube model as it has previously been successfully used to investigate mitochondrial dynamics and has a myofibrillar system similar to atrial myocytes. We observed that rapid stimulation of C2C12 cells resulted in mitochondrial hyperfusion and increased mitochondrial colocalisation with calcium stores. Inhibition of mitochondrial fusion by transfection of mutant DRP1K38E resulted in similar effects on mitochondrial fusion, SR colocalisation and altered calcium handling. Interestingly the effects of ‘forced fusion’ were reversed by co-incubation with the reducing agent N-Acetyl cysteine (NAC). Subsequently we demonstrated that oxidative stress resulted in similar reversible increases in mitochondrial fusion, SR-colocalisation and altered calcium handling. Finally, we believe we have identified that myocyte calcium handling is reliant on baseline levels of reactive oxygen species as co-incubation with NAC both reversed and retarded myocyte response to caffeine induced calcium release and re-uptake. Based on these results we conclude that the coordinate regulation of mitochondrial fusion and MAM contacts may form a point source for stress-induced arrhythmogenesis. We believe that the MAM merits further investigation as a therapeutic target in AF-induced remodelling.

Introduction

Atrial Fibrillation (AF) is the most common sustained cardiac arrhythmia and worldwide, as populations age, the prevalence of AF will continue to increase[1,2]. A progressive disorder, AF immediately impairs cardiac performance increasing risks of stroke, dementia, heart failure and death[3,4]. Current treatment for AF is sub-optimal and will remain so until fundamental mechanisms underpinning AF development and maintenance are better understood[5]. The recent success of catheter ablation in paroxysmal, but not persistent, AF has refocused attention on the importance of atrial tachycardia remodelling (ATR)[5,6].

Initially paroxysmal, AF provokes functional and structural changes which favour arrhythmia maintenance[6,7]. The high frequency electrical activity in the fibrillating human atrium in vitro is associated in vitro with mitochondrial dysfunction, oxidative stress and calcium overload[6,8–11]. If fibrillation persists, surviving atrial myocytes adapt to preserve calcium homeostasis, a process termed ATR[6,12]. However, this new equilibrium alters sarcolemmal ion currents and compromises sarcoplasmic reticulum (SR) function promoting triggered activity, re-entry and ultimately perpetuation of AF[6]. Despite the experimental evidence implicating impaired cardiac handling in these mal-adaptive processes, the mechanistic links between mitochondria, oxidative stress and AF-induced remodelling remain largely unexplained[6].

Previous investigations of AF have consistently noted changes in mitochondrial morphology and distribution within atrial myocytes[10,13,14]. Confirmation that mitochondria form attachments to SR, the mitochondria associated membrane (MAM), thereby creating calcium microdomains facilitating the rapid uptake and storage of [Ca2+]c has implications for ATR which are, as yet, unknown[15,16]. Mitochondria tether to SR via a protein, mitofusin-2 (Mfn-2), which in addition to regulating mitochondrial fusion and [Ca2+]c uptake, maintains mitochondrial respiratory...
homoeostasis and mediates the SR stress response[17–20]. Mitochondria themselves are dynamic organelles responding to short term cellular stress with decreased fission and/or increased fusion, however if stress persists, as occurs during AF, this adaptive response is superceded by mitochondrial fragmentation triggering mitophagy or cellular apoptosis[20,21].

In addition, mitochondria are the primary intracellular source of reactive oxygen species (ROS), highly reactive small molecules containing unpaired electrons which, under basal conditions, couple excitation contraction and metabolism (ECM coupling) [22–24]. However, oxidative stress, wherein excessive production and/or reduced scavenging of ROS occur, causes progressive mitochondrial dysfunction, pro-aryrhythmic calcium handling and ultimately apoptosis[22,25–27]. Markers of oxidative stress are elevated in patients who subsequently develop AF[28]. AF results in atrial myocyte stress, increasing mitochondrial ROS formation, ATR and contractile dysfunction[6,8,29–33]. Antioxidants such as the glutathione precursor N-Acetyl Cysteine (NAC) have been demonstrated to be effective, both in vitro and in vivo, in interrupting this positive feedback cycle[8,28,31,34–36].

We hypothesised that the stress of rapid stimulation would increase mitochondrial fusion and promote the formation of MAM in vitro. In addition we postulated that altered mitochondrial plasticity with or without oxidative stress would reversibly alter SR calcium handling and result in increased formation of MAM. In order to test our hypothesis we selected the C2C12 murine myotube model as (i) it has previously been used to investigate mitochondrial plasticity and the acute effects of oxidative stress in vitro, (ii) it is capable of responding to electric field stimulation (EFS) and (iii) it exhibits a myofibrillar system broadly similar to atrial myocytes[37,38].

Materials and Methods

C2C12 mouse myoblast differentiation

Differentiated C2C12 myotubes (Figure 1) were obtained as recently described by Kuo et al[39]. Briefly, C2C12 myoblasts were cultured at 37°C in an atmosphere of 10% CO2 in complete growth medium (DMEM supplemented with 20% FBS, penicillin-streptomycin and L-glutamine). Once cells reached 70–80% confluence, differentiation was induced by replacing complete medium with low-serum medium (DMEM supplemented with 2% horse serum, penicillin-streptomycin and L-glutamine) and cultured for 24–96 h at 37°C in an atmosphere of 10% CO2. Differentiated myotubes, grown on coverslips (12 mm diameter, 0.16–0.19 mm thickness), were mounted in media with 10 mM M dihydroethidium (hET) 24–96 h post differentiation.

Protein expression and Immunofluorescence studies

Differentiated C2C12 myotubes were harvested, and 50 μg of protein (pooled from 5 separate experiments) were solubilized in 5x Laemmli sample buffer with 5% β-mercaptoethanol, loaded onto NuPAGE® Novex 4–12% Bis-Tris gradient gels, and separated by gel electrophoresis. For immunoblot analysis, proteins were electrophoretically transferred onto nitrocellulose membranes, followed by blocking with 5% non-fat milk in Phosphate Buffered Saline (PBS)-TWEEN 20 (137 mM NaCl, 2.7 mM KCl, 100 mM Na2HPO4, 2 mM KH2PO4, 0.5% Tween 20, pH 7.4). Primary antibodies used were mouse anti-DRP1 antibody (1:1000 dilution), mouse anti-myosin heavy chain (MF-20) antibody (1:5000 dilution), and mouse anti-HSP70 antibody (1:5000 dilution; used as a loading control). Horseradish peroxidase-conjugated secondary antibodies were used. Signals were developed using LuminataTM Forte Western chemiluminescent HRP substrates.

Electrical Field stimulation (EFS) as a model of “fibrillatory stress”

Monolayers of cultured C2C12 myocytes were placed on gelatin/fibronectin coated coverslips suspended in serum free normal Tyrode solution in a custom designed chamber incubated for 24 hours with 5% CO2 at 37°C as previously described[37,38]. Parallel carbon electrodes connected via platinum wire to a GRASS electrical stimulator continuously applied monophasic square-waves of pulse duration 2 ms, field gradients <8 V/cm without harmful chemical reactions[40]. In order to investigate the effects of “fibrillatory stress”, stimulation protocols include control conditions (0 Hz), simulated physiological conditions (1 Hz) and fibrillatory stress (5 Hz), adapted from Tandon et al[40]. Three days post differentiation, nascent C2C12 myotubes were subjected to stimulation protocols for 24 h at 37°C in an atmosphere of 10% CO2. To study SR colocalization, differentiated myotubes were incubated (overnight, 4°C) with mouse anti-PDI antibody (ER marker, 1:100 dilution) and rabbit anti-Tom20 antibody (1:1000 dilution), followed by 3 washes in blocking solution. Differentiated myotubes were then incubated (1:1000 dilution, 45 min, room temperature) with goat anti-mouse (514 nm) or goat anti-rabbit (647 nm) conjugated Alexa Fluor secondary antibodies, followed by 3 washes with PBS. A final incubation with 2.5 μg/ml Hoechst in PBS (to observe DNA) was performed, followed by 3 washes with PBS before mounting onto slides.

Studies of mitochondrial plasticity, SR colocalization and calcium handling

Two-days post differentiation, nascent C2C12 myotubes were infected (500 MOI) with dominant-negative dynamin-related protein 1 (DRP1K38E), a mutant form of the mitochondrial fission factor that promotes an elongated mitochondrial reticulum. Empty adenoviral vectors served as control. In order to observe mitochondrial plasticity, differentiated myotubes were loaded with 100 nM tetramethyl-rhodamine ethyl ester (TMRE). Images were
captured at 440 nm (to visualize DRP1K38E-CFP expression) and 543 nm (to visualize TMRE, red) using the confocal microscope as above.

To study SR colocalization following DRP1K38E transfection, differentiated myotubes were incubated (overnight, 4°C) with mouse anti-mitofusin 2 (Mfn-2) antibody (1:200 dilution) and rabbit anti-calnexin antibody (5 μg/ml), followed by incubation (1:1000 dilution, 45 min, room temperature) with corresponding Alexa Fluor secondary antibodies prior to immunofluorescence studies. Alternatively, intact mitochondria were isolated from differentiated C2C12 myotubes using the mitochondria isolation kit from cultured cells (Pierce), according to manufacturer’s instructions. For immunoblot analysis, 30 μg of protein (pooled from 5 separate experiments) were solubilized and separated by gel electrophoresis. Primary antibodies used were against Mfn-2 (3 μg/ml), Tom20 (1:1000 dilution), SERCA2 (1:500 dilution), calsequestrin (CSQ, 1:2500 dilution), calnexin (1 μg/ml), and cytochrome c (Cyt c, 3 μg/ml, loading control). Secondary infrared dye (IRDye800)-labeled antibodies were used for detection on a LI-COR Odyssey infrared imaging system (LI-COR Biosciences). Band intensity was quantified using Odyssey 2.0 software.

To investigate effects of altered mitochondrial plasticity on calcium handling, differentiated myotubes were washed with

Figure 1. C2C12 myoblasts differentiate into myotubes expressing contractile apparatus and mitochondrial fission machinery. C2C12 myoblasts were cultured and induced to differentiate for 24–96 h at 37°C in an atmosphere of 10% CO2. Panel A: C2C12 myoblasts differentiate into myotubes. Cells were plated onto coverslips and allowed to reach 70–80% confluence before inducing differentiation. Live images were captured at 405 nm (to visualize DNA, blue), 515 nm (to visualize oxidized hET, green) and 633 nm (to visualize mitochondria, red) on a laser-scanning confocal microscope. Panel B: Immunoblots confirming upregulated expression of mitochondrial fission protein DRP1 and contractile protein myosin heavy chain during differentiation in C2C12 myotubes. Cells were harvested, and 50 μg of protein (pooled from 5 separate experiments) were processed for gel electrophoresis. Primary antibodies used were against DRP1, MF-20, and HSP70. Panel C: Translocation of DRP1 to the mitochondrial membrane in C2C12 myotubes. Differentiated C2C12 myotubes were fixed, and incubated with primary antibodies against MF20, DRP1 and/or Tom20 and then corresponding Alexa Fluor secondary antibodies. Confocal microscopy images were captured as described before.

doi:10.1371/journal.pone.0069165.g001
modified Hanks’ buffered saline solution, loaded with 2.5 μM Fluo-4 and 100 nM TMRE, incubated at 37°C in an atmosphere of 10% CO2 for 30 min, and then washed with warmed buffer. The methylxanthine caffeine is recognised at millimolar concentration to release Ca2+ from intracellular stores during diastole, primarily in a ryanodine receptor (RyR) mediated process involving the SR in skeletal muscle preparations[41,42]. We used caffeine (4 mM) added to the superfusate to elicit Ca2+ waves in C2C12 myotubes during laser scanning confocal imaging to detect release of Ca2+ from intracellular stores.

Studies of oxidative stress and altered redox state

Differentiated C2C12 myotubes were treated with the glutathione synthase inhibitor bathionine sulfoximine (BSO, 200 μM, 24 h) or with the thiol-oxidizing agent diamide (100 μM, 2 h). To assess the effect of oxidative stress on mitochondrial plasticity, differentiated myotubes were loaded with 100 nM TMRE, and images were captured using the confocal microscope as above. In a parallel set of experiments, differentiated myotubes were loaded with 100 nM TMRE and 2.5 μM Fluo-4, and calcium release events were triggered by 4 mM caffeine and detected by laser scanning confocal imaging as above. In order to determine the degree to which oxidative stress was reversible, and to establish if a baseline level of oxidation exists, these protocols were repeated previously described, for the final 24 h period prior to processing. Mitochondrial stress, “forced fusion”, increases formation of MAM and results in altered SR calcium handling

Transmission of differentiated C2C12 myotubes with the dominant negative DRP1K38E was associated with a hyperfused mitochondrial reticulum which colocalised with the SR (Figures 3, 4, 5 and 6A). As compared to control myotubes which had been transfected with empty adenoviral vector, cells expressing DRP1K38E displayed a fivefold increase in fused mitochondria (Figure 3B; control 15 ± 5%, DRP1K38E 87 ± 10%, p < 0.001 Student’s t-test, n = 40 myotubes) and a fourfold increase in 2-D size, as measured on electron micrographs (Figure 5B; mitochondrial 2-D size (μm^2): control 0.10 ± 0.04, DRP1K38E 0.43 ± 0.17 p < 0.001 ANOVA, n = 50).

As compared to control myotubes which had been transfected with empty adenoviral vector, cells expressing DRP1K38E released threefold more calcium upon application of 4 mM of caffeine (Amplitude of caffeine induced calcium release in control myotubes 564 ± 181 fluorescence arbitrary units (fau) v. 2221 ± 397 fau in DRP1K38E myotubes, p < 0.001 Student’s t-test, n = 40 myotubes. Figure 6B, C). There was a significant delay in caffeine response time (Figure 6E i) but not in time to peak caffeine effect (Figure 6E ii) in DRP1K38E myotubes as compared to control (caffeine response time control myotubes 8 ± 3 s v. 14 ± 2 s in DRP1K38E expressing myotubes, p < 0.001; Time to peak caffeine effect control myotubes 8 ± 2 s v. 10 ± 2 s in DRP1K38E expressing myotubes, p = ns, n = 40 myotubes for all comparisons). “Forced fusion”, in the absence of stress, was associated with increased formation of MAM as evidenced by a fused mitochondrial reticulum (Figures 3A, 5B) which colocalised with SR (Figure 4) increased frequency of SR:mitochondrial contact sites (Figure 5C) and increased SR proteins present in mitochondrial pellets isolated from C2C12 cells following transfection with DRP1K38E (Figure 6F).

Electron Microscopy Studies

C2C12 myoblasts were grown and induced to differentiate on 22 mm glass coverslips. During differentiation nascent C2C12 myotubes were either infected with DRP1K38E (500 MOI) exposed to oxidative stress or EFS-induced “fibrillatory stress”, as previously described, for the final 24 h period prior to processing. Differentiated myotubes were fixed in 2% glutaraldehyde in 0.1M PBS, pH 7.4, post-fixed in 1% osmium tetroxide in PBS, en bloc stained in 3% uranyl acetate, dehydrated in an ascending series of ethanol, and then processed and embedded in Spurr epoxy resin. Thin sections were cut on a Leica UC 6 ultramicrotome, collected on copper grids and counter-stained with lead citrate. Samples were viewed and images taken with a Jeol 1230 TEM equipped with AMT software.

Statistical Analysis

Results are presented as mean±SE of the mean with the exception of the electron microscopy studies. An unpaired, two-tailed Student’s t-test was used to evaluate the significance of difference between the calculated means of two groups; multiple comparisons were assessed using one way analysis of variance (ANOVA) via the Prism statistical software package (Graphpad Inc, La Jolla, USA). Some confocal and electron microscopy studies were assessed using one way analysis of variance (ANOVA) or Fisher’s exact test where indicated, via Prism as before. A probability value of <0.05 was considered significant.

Results

Rapid stimulation results in increased mitochondrial fusion and SR colocalisation

Exposing differentiated C2C12 cells to sudden and persistent rapid stimulation for 24 h resulted in the formation of elongated mitochondria which colocalised with SR and was associated with an increased frequency of SR:mitochondrial contact sites (Figure 2). There was no significant difference between control cells and those stimulated at 1 Hz in the parameters assessed. However, mitochondrial 2D size increased approximately 5-fold in cells which were stimulated at 5 Hz (Figure 2B) (Mitochondrial 2-D size (μm^2); control 0.06 ± 0.04, EFS 1 Hz 0.08 ± 0.05 and EFS 5 Hz 0.63 ± 0.12 p < 0.001 one way ANOVA, n = 50).

Similarly, the frequency of SR:mitochondrial contact sites markedly increased subsequent to rapid stimulation (Control and EFS 1 Hz 4%, EFS 5 Hz 94%, p < 0.001 Fisher’s exact test, n = 50).
Oxidative stress promotes mitochondrial fusion and results in altered SR calcium handling

It has previously been shown that increased cellular levels of oxidized glutathione, which is common during cellular stress, can lead to the activation of the mitochondrial fusion machinery [43]. Consistent with this, inhibition of GSH synthesis with 200 mM BSO and/or induction of GSH oxidation via diamide (100 mM) altered mitochondrial plasticity promoting a fused reticulum in differentiated C2C12 myotubes (Figures 3, 5 and 7A) [44–47]. Oxidative stress was associated with an almost fourfold increase in fluorescence following caffeine application (amplitude of caffeine induced calcium release in control myotubes 534±684 fau v. 1943±643 fau in myotubes treated with diamide, p<0.001, ANOVA, n = 40 myotubes Figure 7B,C). Similarly there was a pronounced increase in the caffeine response time (Figure 7D i) but not the time to peak caffeine effect (Figure 7D ii) in myotubes under oxidative stress (caffeine response time control myotubes 11±1 s v. 18±1 s in BSO myotubes (p<0.001) and 36±19 s in diamide treated myotubes (p<0.05); Time to peak caffeine effect control myotubes 10±2 s v. 17±3 s in BSO myotubes and 10±5 s in diamide treated myotubes (p=n.s) n = 40 myotubes per group). Again, despite a delayed onset of response to caffeine application as compared with control, there was a significant increase in ΔF/ΔT of the caffeine response (Figure 7D iii) in cells exposed to oxidative stress (caffeine response control myotubes 78±5 fau/s v. 150±21 fau/s in BSO myotubes (p<0.001), 207±105 fau/s in diamide treated myotubes (p<0.05), n = 40 myotubes per group). Oxidative stress mediated by BSO was consistently associated with altered calcium clearance from the cytosol post caffeine application (t25 control myotubes 72±66 s v. 95±16 s in BSO myotubes (p<0.05), and 81±6 s in diamide treated myotubes (p=n.s); t50 control myotubes 78±8 s v. 110±21 s in BSO myotubes (p<0.05), and 89±6 s in diamide treated myotubes (p=n.s); t90 control myotubes 91±14 s v. 202±62 s in BSO myotubes (p<0.01), and 131±32 s in diamide treated myotubes (p<0.05); n = 40 myotubes per group. Figure 7E).

Figure 2. Rapid stimulation promotes mitochondrial fusion, SR colocalisation and increased frequency of SR: mitochondrial contact sites. C2C12 myocytes were induced to differentiate for 72 h, and then were left unstimulated (control) or were subjected to EFS at a frequency of 0 Hz, 1 Hz or 5 Hz for 24 h. Panel A: Representative electron micrographs (x25000) of differentiated C2C12 myotubes following 0 Hz (control), 1 Hz or 5 Hz field stimulation. Arrowheads mark mitochondria-SR contact sites. Panel B: Objective mean quantitative data for (i) mitochondrial 2D area and (ii) the occurrence of visible MAM on 2-D electron microscopy. (i) Mitochondria 2D size is length multiplied by width. (ii) Frequency of mitochondria-SR contact sites is the ratio of the number of mitochondria-SR contacts to the number of mitochondria. Panel C: Confocal images of Tom20 and PDI distribution in electric field stimulated C2C12 myotubes. To illustrate colocalisation of mitochondria and ER markers, the corresponding line scans of Tom20 and PDI are drawn on the right. # Denotes p<0.001 Fisher’s exact test and one-way ANOVA, n = 50 all groups.

doi:10.1371/journal.pone.0069165.g002
ROS inhibition alters both mitochondrial plasticity and SR calcium handling

The physiological role of low levels of intracellular ROS as signalling molecules is well recognised[22]. In order to compare oxidative stress with reducing stress we devised protocols whereby myotubes were first incubated with oxidizing agents then with reducing agents in addition and vice versa. In this way we aimed not only to determine whether the observed effects of oxidative stress on calcium handling and mitochondrial plasticity could be reversed, but also whether reducing agents themselves had any reversible effects on mitochondrial plasticity and calcium handling.

Inhibition of GSH synthesis with 200 μM of BSO promoted mitochondrial fusion as before (Figures 3, 5, 7A & 8A). However, as compared to myotubes similarly cultured but with the addition of the glutathione precursor NAC (200 μM) for the final 24 h period of incubation, a less fused reticulum was observed, suggesting reversibility (Figure 8A). In contrast, the addition of NAC to myotube media resulted in an approximate three to fivefold potentiation of the delay in caffeine response (Figure 8C ii) and the time to peak (Figure 8C iii) caffeine response (caffeine response time control myotubes 7±3 s v. 21±4 s in BSO myotubes (p<0.001) and to 103±26 s in BSO + NAC treated myotubes (p<0.001); time to peak caffeine effect control myotubes 7±0 s v. 15±2 s in BSO myotubes (p<0.001) and 51±13 s in BSO+ NAC treated myotubes (p<0.001), n = 40 myotubes per group). The BSO-mediated increase in ΔF/ΔT of the caffeine response was reversed in those myotubes in which NAC was added to the media prior to experimentation (caffeine response control myotubes 68±15 fau/s v. 172±43 fau/s in BSO myotubes (p<0.001), as compared to 18±9 fau/s in BSO+ NAC treated myotubes (p<0.001), n = 40 myotubes per group. Figure 8C iv). In keeping with the marked delay in fluorescence following reversal of oxidative stress, recovery from fluorescence was similarly delayed (Figure 8F i), particularly in the early to mid phase of recovery (t25

![Figure 3. DRP1K38E expression and altered GSH synthesis modulate mitochondrial plasticity in C2C12 myotubes.](image-url)
control myotubes 77 ± 2 s v. 104 ± 12 s in BSO myotubes (p < 0.001); t50 control myotubes 81 ± 3 s v. 118 ± 21 s in BSO myotubes (p < 0.001), and 261 ± 12 s in BSO + NAC treated myotubes (p < 0.001); t90 control myotubes 105 ± 9 s v. 206 ± 64 s in BSO myotubes (p < 0.001), and 313 ± 25 s in BSO + NAC treated myotubes (p < 0.001), n = 40 myotubes per group).

Addition of the glutathione precursor NAC to myotubes media during culture was associated with an approximate 80% increase in fluorescence which was further doubled by the addition of BSO to culture media (amplitude of caffeine induced calcium release in control myotubes 498 ± 152 fau v. 894 ± 285 fau in myotubes incubated with NAC (p < 0.01) and 1902 ± 387 fau in myotubes treated with NAC + BSO (p < 0.001)), ANOVA, n = 60 myotubes, Figure 8E ii). Again, the addition of NAC to media resulted in an approximate fivefold potentiation of the delay in caffeine response (Figure 8E ii) and a sevenfold increase in the time to peak (Figure 8E iii) caffeine response (caffeine response time control myotubes 10 ± 3 seconds (s) v. 49 ± 10 s in NAC myotubes (p < 0.001); t50 control myotubes 67 ± 8 s in NAC + BSO treated myotubes (p < 0.001); Time to peak caffeine effect control myotubes 6 ± 2 s v. 44 ± 12 s in NAC myotubes (p < 0.001) and 43 ± 7 s in NAC + BSO treated myotubes (p < 0.001), n = 40 myotubes per group). There was no significant effect of NAC on ΔF/ΔT of the caffeine response (Figure 8E iv) and the expected BSO mediated increase in ΔF/ΔT was not observed in the NAC pre-treated myotubes (caffeine response control myotubes 63 ± 10 fau/s v. 43 ± 28 fau/s in NAC myotubes and 44 ± 20 fau/s in NAC + BSO myotubes (p = ns for all comparisons), n = 40 myotubes per group). Addition of NAC to culture media resulted in a significant delay to clearance of calcium from the cytosol (Figure 8F ii), and it was not possible to observe complete recovery in myotubes exposed to

Figure 4. DRP1K38E expression results in a fused mitochondrial reticulum and SR colocalisation in C2C12 myotubes. C2C12 myoblasts were induced to differentiate for 48 h and then were infected with control or DRP1K38E adenoviral vectors for 48 h. Panel A: Confocal images of Mfn-2 and calnexin distribution in differentiated C2C12 myotubes. To illustrate colocalization of mitochondria and ER markers, the corresponding line scans of Mfn-2 and calnexin are drawn on the right. Panel B: Confocal images of Tom20 and PDI distribution in differentiated C2C12 myotubes. To illustrate colocalization of mitochondria and ER markers, the corresponding line scans of Tom20 and PDI are drawn on the right. doi:10.1371/journal.pone.0069165.g004
Altering mitochondrial plasticity, fibrillatory and oxidative stress similarly result in enlarged mitochondria with increased frequency of SR:mitochondrial contacts. Transfection of dominant-negative mutant DRP1K38E resulted in a more fused mitochondrial reticulum composed of fewer, larger mitochondria as compared to control (Figures 3, 4, 5, 6 and 9). Inhibiting mitochondrial fission in this way resulted in a greater than fourfold increase in mitochondrial 2-D cross-sectional area with a significant increase in observed SR:mitochondrial contact points (Figure 5). Similarly, oxidative stress resulted in a significant enlargement of mitochondria, associated with a more fused reticulum (Figures 3, 7, 8, 9) and promoted formation of a greater number of visible SR: mitochondrial contacts (Figure 5C). Interestingly, the addition of the reducing agent NAC had no effect on mitochondrial fusion under control conditions but reversed not only the fusion induced by BSO but also infection with DRP1K38E (Figures 3, 5, 8, and 9). In addition, NAC reversed the effect of ‘forced fusion’ on caffeine induced calcium release (Figure 9).

Addition of the glutathione precursor NAC to myotubes previously infected with DRP1K38E was associated with a reduction in fluorescence to control levels (amplitude of caffeine induced calcium release in control myotubes 564 ± 181 f.u. v. 222 ± 397 f.u. in myotubes incubated with DRP1K38E (p < 0.001) and 795 ± 350 f.u. in myotubes treated with DRP1K38E+ NAC (p = ns), ANOVA n = 40 myotubes Figure 9B, C). Again, the addition of NAC to media resulted in an approximate fourfold...
Mitochondrial Stress, Hyperfusion & Ca\textsuperscript{2+} Handling

potentiation of both the delay in caffeine response (Figure 9D i) and in the time to peak (Figure 9D ii) caffeine response (caffeine response time control myotubes 8±3 s v. 14±2 s in DRP1K38E myotubes (p<0.001) and to 70±8 s in DRP1K38E+ NAC treated myotubes (p<0.001)); Time to peak caffeine effect control myotubes 8±2 s v. 10±2 s in DRP1K38E myotubes (p=n.s) and 50±7 s in DRP1K38E+ NAC treated myotubes (p<0.001), n = 40 myotubes per group). The addition of NAC significantly reversed the effect of DRP1K38E on ΔF/ΔT (Figure 9D iii) of the caffeine response control myotubes 70±11 fau/s v. 245±40 fau/s in DRP1K38E myotubes and 35±20 fau/s in DRP1K38E+ NAC myotubes (p<0.001 for all comparisons), n = 40 myotubes per group). Addition of NAC to culture media resulted in a significant delay to clearance of calcium from the cytosol (Figure 9E). Indeed, it was not possible to observe complete recovery in myotubes exposed to DRP1K38E+ NAC (t\textsubscript{25} control myotubes 77±2 s v. 104±12 s in DRP1K38E myotubes (p=n.s), and 242±9 s in DRP1K38E+ NAC treated myotubes (p<0.001); t\textsubscript{50} control myotubes 81±3 s v. 118±21 s in DRP1K38E myotubes (p<0.01), and 261±12 s in DRP1K38E+ NAC treated myotubes (p<0.001); t\textsubscript{100} control myotubes 105±15 s v. 206±64 s in DRP1K38E myotubes (p<0.001), n = 40 myotubes per group).

Discussion

With these studies we provide further evidence of the intimate relationship between mitochondrial fusion, oxidative stress, and altered SR calcium handling in murine myocytes. Increased MAM formation, via inhibition of mitochondrial fission or triggered by oxidative stress, was associated with increased caffeine induced calcium release with preservation of release kinetics and delayed cytosolic clearance of calcium. Further corroboration of the redox sensitive nature of intracellular calcium transporters is provided by the slowing of SR calcium release and the potentiation of delayed calcium clearance in the presence of the glutathione precursor NAC.

Mitochondria are increasingly recognised to perform a number of vital functions in mammalian cells. In addition to their long appreciated role in cellular respiration, the discovery of MAM has linked mitochondrial plasticity, ECM coupling and cellular stress responses. The central role of rapid bidirectional calcium signalling between SR and mitochondria at the MAM offers tremendous opportunities to improve our understanding of various cardiomyocyte pathophysiologies. We sought to investigate the relationship between fibrillatory and oxidative stress, mitochondrial fusion and calcium handling with simple experiments in order to identify a possible link between AF-induced cellular stress and subsequent atrial tachycardia remodelling.

As mitochondria cannot be generated de novo, they are perpetually being repaired and recycled in a dynamic equilibrium between opposing processes of fission and fusion[20]. This plasticity is required in order to preserve both mitochondrial and myocyte integrity[18]. Fusion produces elongated interconnected mitochondria and formation of mitochondrial networks facilitates the transmission of calcium signals and membrane potential across individual cells[20]. Mitochondrial fission
generates numerous morphologically and functionally distinct isolated mitochondria and can be physiological, as a prelude to mitophagy, or pathological in response to rapid increases in \([\text{Ca}^{2+}]\)[20,48]. Dysfunctional mitochondrial plasticity results in increased sensitivity to apoptotic stimuli and is pro-arrhythmogenic, however the importance of mitochondrial plasticity in human heart disease is only just being recognised[18,49,50].

The Mitochondria-Associated Membrane is the physical association of juxtaposed SR and mitochondrial membranes which facilitates privileged inter-organelle communication performing several physiological functions[15,51–53]. Both inter-organelle proximity and formation of MAM tethers are dynamic, remodelling in response to local \([\text{Ca}^{2+}]\)[53–55]. The first direct MAM tether identified in the mammalian system was Mitofusin-2 (Mfn-2), a large trans-membrane protein residing in the outer mitochondrial membrane (OMM) pivotal to mitochondrial fusion[17]. As SR structural integrity and bidirectional calcium signalling at the MAM are Mfn-2 dependent, the MAM may also perform critical roles in calcium buffering and the regulation of cellular respiration during stress[17,56–58].

Mitochondria generate ATP by oxidative phosphorylation with calcium and ROS acting as signalling molecules in a bioenergetic homeostasis involving mitochondria, SR and the nucleus[22,24]. Mitochondrial respiration is also the primary source of ROS within myocytes which, under normal conditions, remain counterbalanced by calcium-dependent production of reducing agents[22,59]. Accumulation of ROS impairs myofibrillar calcium handling and has been implicated in contractile dysfunction and AF-remodelling[30,36]. Altered phosphorylation of sarcoplasmatic reticulum calcium currents modulates whole cell calcium entry and extrusion, while oxidation of the SR calcium transporters RyR and SERCA modify SR calcium release and re-uptake[26,27,31]. Experimentally induced mitochondrial ROS production induces pro-arrhythmic SR calcium release[60]. If metabolic stress is not relieved, oxidative stress uncouples mitochondria, disrupts cytosolic proteins and nuclear DNA resulting in cellular dysfunction and ultimately apoptosis[22,25,26,61].

Confirmation of the existence of microdomains of \([\text{Ca}^{2+}]\) and ROS at the MAM suggest the possibility of crosstalk between respiration, oxidative stress, mitochondrial plasticity and calcium handling[15,53,60,62,63]. Recent descriptions of stress induced mitochondrial hyperfusion (SIMH) provide further credence to the possibility of such crosstalk[21]. Mitochondria initially respond to a variety of cellular stresses by forming a fused network in an apparent protective effect[20]. Mitochondrial function is maintained temporarily under adverse circumstances; however if stress persists mitochondrial fragmentation and ultimately cell death occur[20,21]. In related work, ROS induced disulphide switching of Mfn-2 promoted mitochondrial fusion in a GTP dependent process directly linking oxidative stress and mitochondrial fusion[43]. Recent data suggests that stress induced phosphorylation of drp1 by PKA inactivates fission, promoting formation of a fused mitochondrial network[20,64]. This hyperfused reticulum appears resistant to mitophagy, conferring temporary protection to

**Figure 7. Oxidative stress promotes mitochondrial fusion and results in altered SR calcium handling.** Panel A: Confocal microscopy images of differentiated C2C12 myotubes having been previously treated with BSO or diamide, and loaded with TMRE. Panel B: Representative tracings of the fluorescence profile of myotubes cultured under oxidative stress and loaded with fluo-4 prior to the application of caffeine to trigger SR calcium release (BSO, Diamide v control). Panel C: Objective mean quantitative data for fluorescence resulting from caffeine induced calcium release (BSO, diamide v control). Panel D: Objective mean quantitative data for fluorescence kinetics profile resulting from caffeine induced calcium release (BSO, diamide v control). Panel E: Histogram displaying the mean quantitative data relating to the normalisation of fluorescence following caffeine application (BSO, diamide v control). * Denotes \(p < 0.05–0.001\) ANOVA, \(\text{n} = 40\) per group.

doi:10.1371/journal.pone.0069165.g007
cells from death triggers, ‘buying time’ for other homoeostatic mechanisms to alleviate stress and thus preserve cellular integri-
yty[20]. It is not known which combination(s) of factors promote switching from SIMH to fragmentation nor the circumstances which determine the threshold for death triggers, although one should not be surprised to learn that emerging data supports a regulatory role for Mfn-2 mediated bidirectional calcium signalling in SR stress responses[17,19].

Our results support a role for hyperfusion in response to stress and broadly speaking appear in agreement with those of the Dorn laboratory[18,65]. Similarly, altered calcium handling by the SR and/or mitochondria associated with increased MAM formation, either as a result of oxidative stress or unopposed fusion, suggests that Mfn-2 is critical to crosstalk. Caffeine induced SR calcium release has previously been reported and is related to proximity of the MAM and the rate of SR emptying[15,66,67]. Opposing sides of the MAM preferentially locate with anchoring of outer and inner mitochondrial membrane permitting privileged access to the respiratory complexes[68]. Prevention of mitochondrial calcium overload has previously been shown to rescue in vitro models of pathological tachycardia and cardiomyopathy[69–72]. Mitochondrial calcium overload eventu-
ally necessitates calcium release back into the cytosol[73–75]. Currently it is not known to what degree this mitochondrial calcium efflux may contribute to SR refilling or diastolic triggering of CICR, particularly if mitochondria have fused into networks[76–81].

Redox potential would appear to have a hermetic response curve suggesting various affinities for stimulatory and inhibitory regulatory pathways ultimately with ROS becoming toxic at high concentration. As the outcome of an intracellular calcium signal ordinarily depends on the strength, localisation, duration and pattern of that signal, SR Ca$^{2+}$ release channeled to a tethered mitochondrion could act as a point source, propagating calcium signals via the mitochondrial network throughout the cell, obviating ‘local control’[57,82–85]. Hence we postulate that Mfn-2 performs multiple, yet related, roles: tethering mitochondria to the SR, creating and maintaining calcium signalling at the MAM, modulating mitochondrial plasticity in response to local [Ca$^{2+}$], transducing SHM if stress can be ameliorated, and if not,

Figure 8. Altering Redox state alters both mitochondrial plasticity and SR calcium handling. Panel A: Confocal microscopy images of differentiated C2C12 myotubes having been previously treated with BSO, NAC or both, and loaded with TMRE. Panels B and D: Representative tracings of the fluorescence profile of myotubes cultured under oxidative stress or reducing agent or both and loaded with fluo-4 prior to the application of caffeine to trigger SR calcium release (BSO, BSO+NAC, NAC, NAC+BSO v. control). Panels C and E: Objective mean quantitative data for caffeine induced calcium release and fluorescence kinetics of cells with altered redox states (BSO, BSO+NAC, NAC, NAC+BSO v. control). Panel F: Histogram displaying the mean quantitative data relating to the normalization of fluorescence following caffeine application (BSO, BSO+NAC, NAC, NAC+BSO v. control). * Denotes $p<0.001$ ANOVA, n = 20 per group.
doi:10.1371/journal.pone.0069165.g008

Mitochondrial Stress, Hyperfusion & Ca$^{2+}$ Handling
initiating the death trigger[86–91]. This possibility raises the question “Could the MAM be a point source for arrhythmogenesis?”

Limitations/Suggested further work

The C2C12 myotube model is derived from murine skeletal myoblasts and has the advantage of being capable of differentiation thus affords the opportunity to observe induced changes in mitochondrial plasticity while developing myofibrillar apparatus. This advantage of the C2C12 model readily permits reproducible interruption of ongoing fission/fusion processes while simultaneously offering the ability to investigate both fibrillatory and oxidative stress. In addition, the study of isolated myocytes permits the observation of autonomous intracellular processes, indicating that ATR can, at least in part, occur in the absence of cellular communication and does not require extracellular factors. However this inherent malleability is in direct contrast to that of terminally-differentiated human atrial myocytes. Unfortunately only relatively simple experiments can be designed examining scarce human atrial tissue from consenting patients.

We believe a more practical solution is to seek to rapidly identify potential candidate factors and regulatory pathways for investigation in the C2C12 myotube model, design and develop techniques for investigating these processes in differentiated atrial myocytes such as HL-1 myocytes and then perform simple corroborative experiments with human tissue.

Conclusions

Long overlooked in cardiovascular pathology, the dynamic equilibrium of mitochondrial plasticity remains in equipoise under basal conditions. Cellular stress, whether as a result of rapid stimulation or from ROS promotes a hyperfused mitochondrial network, increased formation of mitochondria associated membrane and Mfn-2 mediated alterations in SR calcium handling with, as yet unknown, implications for arrhythmogenesis. Further work is necessary to determine the intermediate factors and ultimately the relevance of oxidative stress, mitochondrial plasticity and altered calcium handling in ATR.

Acknowledgments

Dr. Peter Backx for critical appraisal of the manuscript. Mr. Peter Rippstein for technical assistance with the electron microscopy studies.

Author Contributions

Conceived and designed the experiments: CR HM. Performed the experiments: CR MBK GD. Analyzed the data: CR MBK GD HM. Contributed reagents/materials/analysis tools: CR MBK MR HM. Wrote the paper: CR MBK HM.

Figure 9. The reducing agent NAC reverses the alterations in SR calcium release induced by ‘forced fusion’. Panel A: Confocal microscopy images of differentiated C2C12 myotubes expressing DRP1K38E or treated with NAC, and loaded with TMRE. Panel B: Representative tracings of the fluorescence profile of myotubes loaded with fluo-4 prior to the application of caffeine to trigger SR calcium release (DRP1K38E, DRP1K38E+NAC v control). Panel C: Objective mean quantitative data for fluorescence resulting from caffeine induced calcium release (DRP1K38E, DRP1K38E+NAC v control). Panel D: Objective mean quantitative data for fluorescence kinetics profile resulting from caffeine induced calcium release (DRP1K38E, DRP1K38E+NAC v control). Panel E: Histogram displaying the mean quantitative data relating to the normalization of fluorescence following caffeine application (DRP1K38E, DRP1K38E+NAC v control). * Denotes p<0.001 ANOVA, n = 40 per group.
doi:10.1371/journal.pone.0069165.g009
References

1. Ryder KM, Benjamin EJ (1999) Epidemiology and significance of atrial fibrillation. Am J Cardiol 84: 131R–139R.
2. Miyasaka Y, Barnes ME, Gersh BJ, Cha SS, Bailey KR, et al. (2006) Secular trends in incidence of atrial fibrillation in Olmsted County, Minnesota, 1980 to 2000, and implications on the projections for future prevalence. Circulation: 114: 119–125.
3. Benjamin EJ, Wolf PA, D'Agostino RB, Silbershatz H, Kannel WB, et al. (1998) Impact of atrial fibrillation on the risk of death: the Framingham Heart Study. Circulation 98: 946–952.
4. Bunch TJ, Weiss JP, Crandall BG, May HT, Bair TL, et al. (2010) Atrial fibrillation is independently associated with severe, vascular, and Alzheimer’s dementia. Heart Rhythm 7: 433–437.
5. Calkins H, Kuck KH, Cappato R, Brugada J, Camm AJ, et al. (2012) 2012 ACCF/AHA/HRS Focused更新 of the 2007 Guidelines for the Management of Patients With Atrial Fibrillation: an update from the American College of Cardiology Foundation/ American Heart Association Task Force. J Am Coll Cardiol 59: 399–439.
6. Gomes Jr AC, Zhang Y, Trappe S, Cuculich P, Drazin DH, et al. (2005) Atrial natriuretic factor increases calcium-stimulated ATPase activity in human atrial myocardium. Am J Physiol 288: H2153–H2161.
7. Gomes Jr AC, Zhang Y, Trappe S, Cuculich P, Drazin DH, et al. (2005) Atrial natriuretic factor increases calcium-stimulated ATPase activity in human atrial myocardium. Am J Physiol 288: H2153–H2161.
8. Carnes CA, Janssen PM, Ruehr ML, Nakayama H, Nakayama T, et al. (2007) Acrobates attenuates atrial pacing-induced peroxynitrite formation and electrical remodeling and decreases the incidence of postoperative atrial fibrillation. Circ Res 99: e32–e36.
9. Mihimiuc M, Ye F, Carnes CA, Reiser PJ, McCarthy PM, et al. (2001) Impaired myofibrillar energetic and oxidative injury during human atrial fibrillation. Circulation 104: 174–180.
10. Thijssen VL, Ausma J, Borgers M (2001) Structural remodelling during chronic atrial fibrillation. J Cardiovasc Electrophysiol 12: 1070–1077.
11. Ramlawi B, Otsu H, Mieno S, Boolhwami M, Sodha NR, et al. (2007) Oxidative stress and atrial fibrillation after cardiac surgery: a case-control study. Ann Thorac Surg 84: 1166–1172.
12. Shizume M, Shibaia R, Inden Y, Yoshida N, Uchikawa T, et al. (2009) Reactive oxidative metabolites are associated with atrial conduction disturbance in patients with atrial fibrillation. Heart Rhythm 6: 935–940.
13. Van Wagoner DR (2008) Oxidative stress and inflammation in atrial fibrillation: a prospective, placebo-controlled pilot study. Eur Heart J 29: 625–631.
14. Park H, Bhalla R, Saigal R, Radose M, Watson N, et al. (2008) Effects of electrical stimulation in C2C12 muscle constructs. J Tissue Eng Regen Med 2: 279–287.
15. Ashrafian H, Docherty L, Leo V, Towlson C, Neilan M, et al. (2010) A mutation in the mitochondrial fission gene Dnm1l leads to cardiomyopathy. PLoS Genet 6: e1000883.
16. Aon MA, Cortassa S, Maack C, O'Rourke B (2007) Sequential opening of the mitochondrial OXPHOS complex during calcium overload. J Cell Biol 178: 133–139.
52. Csordas G, Thomas AP, Hajnoczky G (2001) Calcium signal transmission between ryanodine receptors and mitochondria in cardiac muscle. Trends Cardiovase Med 11: 269–275.
53. Csordas G, Renken C, Varnai P, Walter L, Weaver D, et al. (2006) Structural and functional features and significance of the physical linkage between ER and mitochondria. J Cell Biol 174: 915–921.
54. Rizzuto R, Pinton P, Carrington W, Fay FS, Fogarty KE, et al. (1998) Close contacts with the endoplasmic reticulum as determinants of mitochondrial Ca2+ responses. Science 280: 1765–1766.
55. Soubanier V, McBride HM (2009) Positioning mitochondrial plasticity within cellular signaling cascades. Biochim Biophys Acta 1793: 154–170.
56. Chen H, Detmer SA, Ewald AJ, Griffin EE, Frazer SE, et al. (2003) Mitofusins Mfn1 and Mfn2 coordinate regulate mitochondrial fusion and are essential for embryonic development. J Cell Biol 160: 189–200.
57. de Brito OM, Scorrano L (2010) An intimate liaison: spatial organization of the endoplasmic reticulum-mitochondria relationship. EMBO J 29: 2715–2725.
58. Bravo R, Vicencio JM, Parra V, Troncoso R, Munoz JP, et al. (2011) Increased ER-mitochondrial coupling promotes mitochondrial respiration and bioenergetics during early phases of ER stress. J Cell Sci 124: 2143–2152.
59. Turrens JF (1997) Superoxide production by the mitochondrial respiratory chain. Bioessays 19: 3–8.
60. Yan Y, Liu J, Wei C, Li K, Xie W, et al. (2008) Bidirectional regulation of Ca2+ sparks by mitochondria-derived reactive oxygen species in cardiac myocytes. Cardiovasc Res 77: 412–421.
61. Kowalcowski AJ, Vercesi AE (1999) Mitochondrial damage induced by conditions of oxidative stress. Free Radic Biol Med 26: 463–471.
62. Davison SM, Duchen MR (2008) Mitochondrial calcium microdomains and oxidative stress. Cell Calcium 43: 561–574.
63. Odagiri K, Katoh K, Kawashima H, Tanaka O, Ohtani H, et al. (2009) Local control of mitochondrial membrane potential, permeability transition pore and reactive oxygen species by calcium and calmodulin in rat ventricular myocytes. J Mol Cell Cardiol 46: 989–997.
64. Schauß AC, Huang H, Choi SY, Xu L, Southey RD, et al. (2010) A novel cell-free mitochondrial fusion assay amenable for high-throughput screenings of fusion modulators. BMC Biol 8: 100.
65. Chen Y, Csordas G, Jowdy C, Schneider TG, Csordas N, et al. (2012) Mitofusin-2-Containing Mitochondrial-Reticular Microdomains Direct Rapid Cardiomyocyte Bioenergetic Responses via Inter-Organelle Ca2+-Based Crosstalk. Circ Res 111: 275: 15305–15313.
66. Castaldo P, Cataldi M, Magi S, Lariccia V, Arcangeli S, et al. (2009) Role of the mitochondrial sodium/calcium exchanger in neuronal physiology and in the pathogenesis of neurological diseases. Prog Neurobiol 87: 50–79.
67. Stern MD (1992) Theory of excitation-contraction coupling in cardiac muscle. Physiol Rev 73: 262–408.
68. Davison SM, Duchen MR (2008) Mitochondrial calcium microdomains and oxidative stress. Cell Calcium 43: 561–574.
69. Schauenstein K, Hauer H, Cho SY, Xiu L, Southey RD, et al. (2010) A novel cell-free mitochondrial fusion assay amenable for high-throughput screenings of fusion modulators. BMC Biol 8: 100.
70. Chen Y, Csordas G, Jowdy C, Schneider TG, Csordas N, et al. (2012) Mitofusin-2-Containing Mitochondrial-Reticular Microdomains Direct Rapid Cardiomyocyte Bioenergetic Responses via Inter-Organelle Ca2+-Based Crosstalk. Circ Res 111: 275: 15305–15313.
71. Kohlhass M, Maack C (2010) Adverse bioenergetic consequences of Na(+) handling by the mitochondrial Na(+)-Ca(2+) exchanger. Circ Res 106: 534–536.
72. Zhang T, Guo T, Mishra S, Dalton ND, Kranias EG, et al. (2010) Phospholamban ablation rescues sarcoplasmic reticulum Ca2+-handling but exacerbates cardiac dysfunction in CaMKIIδε/δε(C) transgenic mice. Circ Res 106: 534–536.
73. Tang Y, Zucker RS (1997) Mitochondrial involvement in post-tetanic potentiation of synaptic transmission. Neuron 18: 481–491.
74. David G, Barrett JN, Barrett EF (1998) Evidence that mitochondria buffer physiological Ca2+ loads in lizards motor nerve terminals. J Physiol 509 (Pt 1): 59–65.
75. Yang F, He XP, Russell J, Lu B (2003) Ca2+-influx-independent synaptic potentiation mediated by mitochondrial Na+/Ca2+-exchanger and protein kinase C. J Cell Biol 163: 511–523.
76. Carafoli E, Tomaselli R, Lugli G, Civetta F, Kranias EG, et al. (2006) Mitochondrial calcium handling in heart failure. J Clin Investig 117: 2431–2444.
77. Zhang T, Guo T, Mishra S, Dalton ND, Kranias EG, et al. (2010) Phospholamban ablation rescues sarcoplasmic reticulum Ca2+-handling but exacerbates cardiac dysfunction in CaMKIIδε/δε(C) transgenic mice. Circ Res 106: 354–362.
78. Tang Y, Zucker RS (1997) Mitochondrial involvement in post-tetanic potentiation of synaptic transmission. Neuron 18: 481–491.
79. David G, Barrett JN, Barrett EF (1998) Evidence that mitochondria buffer physiological Ca2+ loads in lizards motor nerve terminals. J Physiol 509 (Pt 1): 59–65.
80. Yang F, He XP, Russell J, Lu B (2003) Ca2+-influx-independent synaptic potentiation mediated by mitochondrial Na+/Ca2+-exchanger and protein kinase C. J Cell Biol 163: 511–523.
81. Castaldo P, Cataldi M, Magi S, Lariccia V, Arcangeli S, et al. (2009) Role of the mitochondrial sodium/calcium exchanger in neuronal physiology and in the pathogenesis of neurological diseases. Prog Neurobiol 87: 50–79.
82. Stern MD (1992) Theory of excitation-contraction coupling in cardiac muscle. Physiol Rev 73: 262–408.
83. Cheng H, Lederer WJ, Cannell MB (1996) Calcium sparks and [Ca2+]i waves in cardiac myocytes. Am J Physiol 270: C148–C159.
84. Cheng H, Lederer WJ, Cannell MB (1995) Calcium sparks: elementary events underlying excitation-contraction coupling in heart muscle. Science 262: 740–744.
85. Venetucci LA, Trafford AW, O’Neill SC, Eimer DA (2008) The sarcoplasmic reticulum and arhythmogenic calcium release. Cardiovasc Res 77: 263–292.
86. Chen Y, Csordas G, Jowdy C, Schneider TG, Csordas N, et al. (2012) Mitofusin-2-Containing Mitochondrial-Reticular Microdomains Direct Rapid Cardiomyocyte Bioenergetic Responses via Inter-Organelle Ca2+-Based Crosstalk. Circ Res 111: 275: 15305–15313.
87. Kohlhass M, Maack C (2010) Adverse bioenergetic consequences of Na+/Ca2+ exchanger-mediated Ca2+ influx in cardiac myocytes. Circulation 122: 2273–2280.
88. Garcia-Perez C, Schneider TG, Hajnoczky G, Csordas G (2011) Alignment of sarcoplasmic reticulum-mitochondrial junctions with mitochondrial contact points. Am J Physiol Heart Circ Physiol 301: H1907–H1915.
89. Minaminisawa S, Yoshijima M, Chu G, Ward CA, Frank K, et al. (1999) Chronic phospholamban-sarcoplasmic reticulum calcium ATPase interaction is the critical calcium cycling defect in dilated cardiomyopathy. Cell 99: 313–322.
90. Schlaf L, Bukowska A, Gardemann A, Polczyk P, Krüllhoff G, et al. (2006) Rapid pacing of embryoid bodies impairs mitochondrial ATP synthesis by a calcium-dependent mechanism—a model of in vitro differentiated cardiomyocytes to study molecular effects of tachycardia. Biochim Biophys Acta 1762: 608–615.