The role of mitochondria for the regulation of cardiac alternans

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Electro-mechanical and Ca alternans is a beat-to-beat alternation of action potential duration, contraction strength and Ca transient amplitude observed in cardiac myocytes at regular stimulation frequency. Ca alternans is a multifactorial process that is causally linked to cardiac arrhythmias. At the cellular level, conditions that increase fractional release from the sarcoplasmic reticulum or reduce diastolic Ca sequestration favor the occurrence of alternans. Mitochondria play a significant role in cardiac excitation–contraction coupling and Ca signaling by providing the energy for contraction and ATP-dependent processes and possibly by serving as Ca buffering organelles. Here we tested the hypothesis that impairment of mitochondrial function generates conditions that favor the occurrence of Ca alternans. Alternans were elicited by electrical pacing (>1 Hz) in single cat atrial myocytes and intracellular Ca ([Ca]) was measured with the fluorescent Ca indicator Indo-1. The degree of alternans was quantified as the alternans ratio (AR = 1 – S/L, where S/L is the ratio of the small to the large amplitude of a pair of alternating Ca transients). Dissipation of mitochondrial membrane potential (with FCCP) as well as inhibition of mitochondrial F,F-ATP synthase (oligomycin), electron transport chain (rotenone, antimycin, CN-), Ca-dependent dehydrogenases and mitochondrial Ca uptake or extrusion, all enhanced AR and lowered the threshold for the occurrence of Ca alternans. The data indicate that impairment of mitochondrial function adversely affects cardiac Ca cycling leading to proarrhythmic Ca alternans.

Keywords: mitochondria, intracellular [Ca], Ca alternans, electron transport chain, permeability transition pore, mitochondrial Ca cycling, TCA cycle

INTRODUCTION

CARDIAC ALTERNANS: A BRIEF REVIEW

Cardiac alternans describes the cyclic, beat-to-beat variations in contraction amplitude (mechanical alternans), action potential duration (electrical or APD alternans), and Ca transient amplitude (Ca alternans) at constant stimulation frequency (e.g., Wohlfart, 1982). Alternans occurs in cardiac failure (e.g., Dumitrescu et al., 2002), during myocardial ischemia, and is believed to be an important factor in the pathogenesis of arrhythmias (Dilly and Lab, 1988; Konta et al., 1990) and the development of re-entry phenomena (Rubenstein and Lipsius, 1995; Berger, 2000; Pastore and Rosenbaum, 2000). Clinically, alternans is manifest as pulsus alternans and specific changes in the ST segment of the electrocardiogram (e.g., Uno, 1991; Surawicz and Fisch, 1992; Lab and Seed, 1993; Pastore et al., 1999). T-wave alternans has been linked to ventricular fibrillation and sudden cardiac death under various conditions including acute myocardial ischemia (Smith et al., 1988; Verrier and Nearing, 1994) and long-QT syndrome (Shimizu and Antzelevitch, 1999). T-wave alternans is now established as one of the strongest markers of susceptibility to sudden cardiac death (Rosenbaum et al., 1994; Walker and Rosenbaum, 2003). Thus, cardiac alternans is thought to be a precursor to life-threatening arrhythmias as both occur in settings of myocardial ischemia, genetic channelopathies, and drug and electrolyte disturbances (Walker and Rosenbaum, 2003). Atrial fibrillation is the most common cardiac arrhythmia that affects a significant segment of the elderly population and is linked to high risk of embolic complications and stroke (for review see Nattel, 2002). Moreover, atrial fibrillation has been linked to APD alternans (Narayan et al., 2002; Hiromoto et al., 2005). Thus alternans is not only a well established risk factor for ventricular dysrhythmias, but also for atrial arrhythmia.

In intact hearts, cardiac muscle preparations or isolated myocytes, alternans can be induced experimentally by pacing (see review Euler, 1999), acidosis (Orchard et al., 1991; Kapur et al., 2009), low extracellular [Ca] (Badeer et al., 1967), hypothermia (e.g., Spencer et al., 1992), and as we demonstrated, by direct interference with cellular energy metabolism through inhibition of glycolysis (Huser et al., 2006b; Kockskamper and Blatter, 2002; Kockskamper et al., 2005). Furthermore, β-adrenergic stimulation, hypercalcemia, cardiac glycosides, and Ca channel blockers can counteract the likelihood for the occurrence of alternans by increasing the critical pacing threshold leading to alternans (see Euler, 1999). There is strong evidence that alternans is ultimately linked to alterations in myocardial Ca homeostasis and impaired [Ca] regulation (for review see Blatter et al., 2003). Numerous factors involved in excitation–contraction coupling (ECC) and Ca-induced Ca release (CICR) have been proposed to be involved in Ca alternans, including alternating voltage-dependent L-type Ca current (I Ca,L) that triggers CICR (Fox et al., 2002; Li et al., 2009), calmodulin-dependent protein kinase II activity (Livshitz and Rudy, 2007), SERCA activity...
(Kameyama et al., 2003; Xie et al., 2008; Cutler et al., 2009), redox modulation (Belevych et al., 2009), and status of the sarcoplasmic reticulum (SR) ryanodine receptor (RyR) Ca release channel (Huser et al., 2000b; Diaz et al., 2002), activation of IP3 receptor dependent Ca release (Zima and Blatter, 2004), temporal delay in Ca movement between SR Ca uptake and release sites (Lab and Lee, 1990; Kihara and Morgan, 1991), SR Ca load (Eisner et al., 2000; Diaz et al., 2004; Picht et al., 2006; Li et al., 2009), and SR Ca buffering by calcequestrin (Restrepo et al., 2008). For details see reviews Euler, 1999; Rosenbaum, 2001; Blatter et al., 2003; Walker and Rosenbaum, 2003; Eisner et al., 2006; Sato et al., 2006; Weiss et al., 2006; Clusin, 2008; Myles et al., 2008). Ca alternans shows regional differences (Diaz et al., 2004; Aistrup et al., 2006, 2009; Gaeta et al., 2009). We have demonstrated that these inhomogeneities are particularly pronounced in atrial tissue. Atrial myocytes have a poorly developed or lack entirely a T-tubular system and the mechanism of Ca release during ECC differs significantly from ventricular myocytes (Huser et al., 1996; Sheehan and Blatter, 2003). Due to these unique structural features Ca alternans in atrial cells are spatially inhomogeneous revealing transverse and longitudinal gradients of the degree of Ca alternans and subcellular regions alternating out-of-phase (Kockskamper and Blatter, 2002). These pronounced and complex subcellular inhomogeneities are linked to propagating arrhythmogenic Ca waves (Kockskamper and Blatter, 2002; Xie and Weiss, 2009).

The multitude of experimental conditions and interventions that cause and modulate Ca and electro-mechanical alternans suggest that Ca alternans is a multifactorial process. This is consistent with the fact that a comprehensive mechanism that can explain and predict the occurrence of cardiac alternans, has not been established to date. Nonetheless, invaluable insight has come from computational models (Shiferaw et al., 2003; Sato et al., 2006; Shiferaw and Karma, 2006; Hayashi et al., 2007; Jordan and Christini, 2007; Livshitz and Rudy, 2007; Sato et al., 2007; Mahajan et al., 2008; Rovetti et al., 2010) of cardiac Ca signaling, and ion current activity during ECC and APD alternans (for a comprehensive review see Weiss et al., 2006). From these studies it became evident that two parameters play equally important roles: fractional SR Ca release, and the efficiency of Ca sequestration. Fractional Ca release refers to the non-linear relationship between SR Ca load and amount of Ca (or % of SR Ca content) released by CICR, where at a higher load a larger fraction is liberated upon activation of CICR (Bassani et al., 1995). Ca sequestration is referred to here as a phenomenological parameter that includes Ca reuptake via SERCA, extrusion via NCX and plasmalemmal Ca-ATPase (PMCA), cytosolic buffering, diastolic SR Ca leak, and finally mitochondrial uptake. In general, factors increasing fractional release promote, and factors increasing Ca sequestration efficiency protect against alternans.

**Ca SIGNALING, MITOCNDRIAL FUNCTION, AND CARDIAC ALTERNANS**

Fractional Ca release and Ca sequestration critically depend on two factors: (i) availability of ATP to fuel Ca pumps and to serve as substrate for phosphorylation processes, and (ii) organelles capable of storing Ca. For both tasks mitochondria play a central role. Mitochondria affect ECC and Ca signaling in cardiac cells at different levels. As a key source of ATP, mitochondria provide the fuel for the activation of the contractile machinery, and for ATP consuming Ca pumps (SERCA and PMCA). ATP is further required for phosphorylation of Ca handling proteins (SERCA, RyR, NCX) and also acts as a direct modulator (through MgATP) of RyR activity. Mitochondria are capable of storing significant amounts of Ca through uptake of Ca mainly via the electrogenic mitochondrial Ca uniporter (MCU) driven by the electrochemical gradient across the inner mitochondrial membrane (IMM), a rapid mode of Ca uptake (RaM) (Sparragna et al., 1995; Gunter et al., 2000; Buntinas et al., 2001), and possibly via mitochondrial RyR (Beutner et al., 2001, 2005; Altschafl et al., 2007) localized in the IMM. In cardiac cells Ca extrusion from mitochondria occurs predominantly via the mitochondrial NCX, whereas a participation of the mitochondrial permeability transition pore has remained controversial. A highly debated and still unresolved issue is whether kinetics and affinities of Ca transporters in cardiac mitochondria allow for Ca uptake rapid enough to allow [Ca]mito to follow beat-to-beat changes of cytosolic [Ca], (for review see Huser et al., 2000a; Dedkova and Blatter, 2008; O’Rourke and Blatter, 2009). There is experimental support for two different, and to some extent mutually exclusive models of how [Ca]mito responds to oscillatory (beat-to-beat) changes of [Ca]. This may occur either as a slow integration of cytosolic [Ca], transients (Miyata et al., 1991; Moravec and Bond, 1991; Sedova et al., 2006), or as beat-to-beat oscillations of [Ca]mito (Isenberg et al., 1993; Robert et al., 2001; Maack et al., 2006).

Mitochondrial [Ca] levels directly modulate mitochondrial functions. Elevated [Ca]mito levels play an important role in the initiation of cell death pathways (Crompton, 1999) and activate several key enzymes in the mitochondrial matrix to enhance ATP production. Oxidation of metabolic substrates in the tricarboxylic acid cycle (TCA) by mitochondrial dehydrogenases is coupled with mitochondrial ATP synthesis through the electron transport chain (ETC). The TCA cycle contains four Ca-dependent mitochondrial dehydrogenases (glycerol 3-phosphate dehydrogenase, pyruvate dehydrogenase, NAD-linked isocitrate dehydrogenase, and 2-oxoglutarate dehydrogenase). Activation of mitochondrial dehydrogenases accelerates the production of NADH, which promotes the generation of the electrochemical proton gradient (ΔΨint) that in turn provides the driving force for the F0/F1-ATP synthase, thus maintaining ATP production. Thus, Ca homeostasis and mitochondrial energy metabolism are intimately interconnected. Nonetheless, the direct effects of mitochondrial function on Ca alternans has not been studied in depth.

Therefore, in the present study we tested the hypothesis that impairment of mitochondrial function, either by reducing Ca buffering or inhibition of energy metabolism, enhances the susceptibility for pacing-induced Ca alternans in atrial tissue. Part of this work has been presented in abstract form (Florea and Blatter, 2005).

**MATERIALS AND METHODS**

**MYOCYTE ISOLATION**

Cat atrial myocytes were isolated according to the method described previously (Wu et al., 1991). All protocols and procedures were approved by the Institutional Animal Care and Use Committee. Adult cats of either sex were anesthetized with sodium pentobarbital (50 mg kg⁻¹) and hearts were excised, mounted on a Langendorff...
apparatus and retrogradely perfused through the aorta with oxygenated collagenase-containing solution (37°C). Dissociated cells were placed on glass coverslips and allowed to attach for 15 min. All experiments were performed at room temperature (22–25°C) where the threshold frequency for pacing-induced Ca alternans is generally lower than at body temperature (Huser et al., 2000b). During experiments, cells were continuously superfused with standard Tyrode solution containing (mM): 140 NaCl, 5 KCl, 1 MgCl₂, 10 glucose, 10 HEPES, 2 CaCl₂, and pH 7.3 (adjusted with NaOH). All inhibitors, agonists, and antagonists were added to normal Tyrode solution from stock solutions.

INTRACELLULAR [Ca] MEASUREMENTS
For [Ca] measurements, atrial myocytes were loaded with the membrane-permeable form of the fluorescent Ca indicator indo-1/AM (5 μM; Invitrogen/Molecular Probes, Carlsbad, CA, USA) in standard Tyrode solution for 15–20 min at room temperature, followed by a >20 min de-esterification interval in dye-free media. Indo-1 loaded cells were excited at 360 nm and emission signals were collected simultaneously at 410 nm (F₄₁₀) and 485 nm (F₄₈₅) using photomultiplier tubes. Fluorescence signals were background subtracted and [Ca] changes were expressed as changes in ratio R = F₄₁₀/F₄₈₅. Ca transients and Ca alternans were elicited by electrical field stimulation (voltage set ~50% above threshold) using a pair of platinum electrodes. Pacing-induced stable Ca alternans was elicited by increasing the stimulation frequency from 0.5 to >1 Hz. The degree of Ca alternans was quantified as the alternans ratio (AR). The AR was defined as 1 – S/L, where S/L is the ratio of the small (S) to the large (L) amplitude Ca transient during a pair of alternating Ca transients (Wu and Clusin, 1997; Kockskamper and Blatter, 2002). Thus, the AR had values between 1 and 0, with AR = 0 indicating no alternans and AR = 1 indicating the highest possible degree of alternans, with only every other stimulation resulting in a measurable Ca transient.

CHEMICALS
Oligomycin, antimycin, sodium cyanide (CN⁻), α-keto-β-methylvaleric acid sodium salt (KMVA), ruthenium red (RuR), and carbonyl cyanide 4-(trifluoromethoxy) phenylhydrazone (FCCP) were obtained from Sigma-Aldrich (St. Louis, MO, USA). Ruthenium 360 (Ru360) and cyclosporin A (CsA) were obtained from Calbiochem (San Diego, CA, USA). CGP-37157 was obtained from Calbiochem (San Diego, CA, USA). CGP-37157 was obtained from Calbiochem (San Diego, CA, USA). CGP-37157 was obtained from Calbiochem (San Diego, CA, USA). CGP-37157 was obtained from Calbiochem (San Diego, CA, USA). CGP-37157 was obtained from Calbiochem (San Diego, CA, USA). CGP-37157 was obtained from Calbiochem (San Diego, CA, USA). CGP-37157 was obtained from Calbiochem (San Diego, CA, USA). CGP-37157 was obtained from Calbiochem (San Diego, CA, USA). CGP-37157 was obtained from Calbiochem (San Diego, CA, USA). CGP-37157 was obtained from Calbiochem (San Diego, CA, USA). 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increased metabolic demands by accelerating mitochondrial ATP production. Increasing pacing rate results in both, increase in $[\text{Ca}^+]$ and ATP demand but also in Ca alternans. We therefore tested whether inhibition of mitochondrial respiration or blockade of the TCA cycle generates conditions favorable to the occurrence of Ca alternans. As shown in Figures 2A–C inhibition of mitochondrial ETC complex I (rotenone, 5 μM), complex III (antimycin, 5 μg/ml), and complex IV (CN⁻, 1 mM) led to an enhancement of Ca alternans to various, but always significant degrees (Figure 2E) between 31% (rotenone) and 82% (CN⁻). The combined application of antimycin and oligomycin increased AR further (115% increase compared to control).

We tested the involvement of the TCA cycle by inhibiting α-ketoglutarate dehydrogenase (αKGDH) with KMVA (5 mM). As shown in Figure 2D inhibition of one of the key Ca-dependent dehydrogenases of the TCA cycle caused a significant increase of the AR from 0.39 ± 0.06 to 0.74 ± 0.06 (+90%; $n = 11$; $p < 0.05$).

In summary, inhibition of the mitochondrial ETC and TCA cycle significantly increased AR indicating that impaired mitochondrial energy metabolism generates proarrhythmic conditions in the form of Ca alternans.

**EFFECTS OF MITOCHONDRIAL Ca CYCLING ON Ca ALTERNANS**

We tested the hypothesis whether inhibition of mitochondrial Ca uptake via MCU or disruption of the mitochondrial Ca extrusion pathway via NCX_mito had effects on pacing-induced Ca alternans. As shown previously ruthenium red (RuR) abolished Ca uptake by cardiac mitochondria (Sedova et al., 2006). Inhibition of MCU with RuR (10 μM) increased AR on average from 0.40 ± 0.02 (control; $n = 52$) to 0.60 ± 0.07 ($n = 12$; $p < 0.05$) or by 50%. The more specific MCU inhibitor Ru360 (Matlib et al., 1998) had a more profound effect and increased AR by 78% (AR in the presence of 5 μM Ru360: 0.71 ± 0.15; $n = 5$; $p < 0.05$).

Next, we selectively inhibited mitochondrial Ca extrusion via NCX_mito. As shown previously, in cat cardiac mitochondria physiological Ca extrusion occurs solely via NCX_mito (Sedova et al., 2006). NCX_mito was blocked with the benzothiazepine compound CGP-37157 (Cox et al., 1993). CGP-3715 at 2.5 μM increased AR by 98% to 0.79 ± 0.15 ($n = 16$; $p < 0.05$), whereas a higher CGP-37157 concentration of 10 μM lead to nearly maximal alternans (AR = 0.94 ± 0.04; $n = 12$; $p < 0.05$).

![Figure 2](image_url)
Among the inhibitors of mitochondrial Ca cycling used, CGP-37157 had the most profound effect on the AR. As shown previously, with each Ca transient mitochondria accumulate Ca which is normally counterbalanced by extrusion via NCX_{mito}. However during inhibition of NCX_{mito}, Ca entering mitochondria is not returned to the cytosol, resulting in mitochondrial Ca overload. Elevated matrix [Ca] is a known inducer of the mitochondrial permeability transition pore, thus mitochondrial Ca overload may lead to PTP opening (Dedkova and Blatter, 2008), dissipation of ΔΨ_m (Huser et al., 1998), and ultimately loss of mitochondrial function. We used the PTP inhibitor cyclosporin A (CsA, 10 μM) to test whether blockade of PTP could counteract the effects of CGP-37157 on Ca alternans. Figure 3D shows an example where CGP-37157 (2.5 μM) caused a maximum degree of alternans where every other stimulation failed to evoke a Ca transient (AR = ~1). Subsequent exposure to CsA (10 μM) partially rescued alternans by improving AR on average to 0.58 ± 0.07 (n = 6). While CsA was not capable of normalizing Ca alternans completely, the results indicate that mitochondrial Ca overload followed by Ca-induced PTP opening affects Ca alternans negatively and contributes to proarrhythmic conditions.

MITOCHONDRIAL INHIBITORS ENHANCE Ca ALTERNANS WITHOUT AFFECTING SR Ca LOAD

Refilling of the SR Ca store after each AP-induced Ca release occurs via SERCA under the consumption of ATP. We tested the hypothesis whether inhibition of mitochondrial metabolism could lower cytosolic ATP levels enough to affect SERCA function and subsequently refilling of the SR. Under conditions of declined cytosolic ATP levels beat-to-beat alternations in the degree of refilling of the SR could ultimately lead to Ca alternans. To test this hypothesis we challenged individual myocytes in the absence and presence of mitochondrial inhibitors twice during pacing-induced Ca alternans, and we compared the amplitude of the caffeine-evoked Ca release after the large (L) and the small (S) Ca transient. Figure 4A (left) shows

![Figure 4A](image-url)

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that the amplitude of the caffeine induced Ca transients were the same irrespective whether they were evoked after a large or a small transient during alternans. This is consistent with our earlier observation that alternating diastolic SR content was not a requirement for the occurrence of Ca alternans (Huser et al., 2000b; Picht et al., 2006). Treatment with oligomycin (1 μg/ml) lead to an enhancement of Ca alternans (increase in AR; Figure 4A right), however SR Ca content was not affected and did not differ whether caffeine (10 mM) was applied after a large or small Ca transient. Similar results were obtained for treatments with CGP-37157 and KMVA (Figure 4B). Thus, despite presumed suppression of mitochondrial ATP production, metabolism and Ca extrusion in the presence of oligomycin, KMVA and CGP-37157, SR Ca replenishment does not appear to be affected. This suggests that cytosolic ATP levels are still sufficiently high to allow for normal SERCA activity and/or ATP for SERCA is supplied from a different source. This is consistent with the observation that ATP derived from glycolysis is utilized preferentially by membrane ion transport mechanisms, including the SERCA pump (Xu et al., 1995). Furthermore, we have shown that inhibition of glycolysis lowers the threshold for the occurrence of electro-mechanical and Ca alternans (Huser et al., 2000b; Kockskamper and Blatter, 2002). This may occur via a mechanism involving reduced ATP supply for SERCA, but based on our observations we also suggested that periodic alterations in the gain of ECC, i.e., the efficacy of a given trigger signal to release Ca from the SR, are controlled locally in the microenvironment of the SR Ca release sites by mechanisms utilizing ATP, produced by glycolytic enzymes closely associated with the release channel. Alternatively, cat atrial cells have low levels of phospholamban expression (Luss et al., 1999), therefore SERCA pumps are more efficient in pumping Ca into the SR, which might contribute to an unchanged SR load.

**DISCUSSION**

Our study shows that any intervention that interferes with mitochondrial functions, either by impairing mitochondrial ATP production or mitochondrial Ca buffering, enhanced Ca alternans. All
pathways tested here, from direct inhibition of the mitochondrial 
$F_1/F_0$-ATP synthase, over collapsing $\Delta \Psi_{mito}$, inhibition of ETC and Ca-dependent mitochondrial dehydrogenases to blocking mitochondrial Ca uptake and extrusion, directly or indirectly affect diastolic Ca sequestration. As outlined earlier, factors diminishing Ca sequestration generate favorable conditions for Ca alternans. Diastolic Ca sequestration typically occurs against electrical and/or chemical gradients for Ca ions, thus involve energy consuming processes and the availability of ATP. Therefore, one would expect that reduced mitochondrial ATP production is paralleled by impaired Ca removal from the cytosol. It remains to be determined whether this occurs via reduced reuptake of Ca into the SR, diminished Ca extrusion across the plasma membrane or even reduced mitochondrial Ca uptake. Inhibition of the mitochondrial uniporter indeed enhanced Ca alternans. We have shown previously that mitochondria accumulate Ca when exposed to higher frequencies of Ca transients, however elevated matrix [Ca] will eventually reduce the electrochemical gradient for mitochondrial Ca uptake. It remains a matter of debate (recently discussed in O’Rourke and Blatter, 2009) whether beat-to-beat cytosolic Ca transients translate into oscillatory changes of $[Ca]_{mito}$, or whether changes in beating frequency are merely reflected by changes in steady-state $[Ca]_{mito}$. Based on our previous investigations it is unlikely that $[Ca]_{mito}$ alternans would be observed at pacing frequencies used in this study (Sedova et al., 2006). Nevertheless, several studies have shown that under conditions of increased workload, as it may occur during $\beta$-adrenergic stimulation or high frequency pacing, mitochondrial Ca transients could be discerned (reviewed e.g., in Huser et al., 2000a; Dedkova and Blatter, 2008; O’Rourke and Blatter, 2009). Therefore, it is conceivable that during high frequency stimulation leading to Ca alternans, mitochondrial Ca alternans could be observed, however to date there is no experimental evidence that cytosolic Ca alternans are paralleled by alternans of $[Ca]_{mito}$.

Certain pathological conditions can be even more favorable for Ca alternans. For example, in heart failure where SERCA expression is reduced and Ca release from the SR (Ca leak) is increased, or during acute cardiac ischemia (where SR Ca load is initially unaffected, but SERCA activity is diminished due to reduced ATP levels), the heart is pushed into instability due to diminished Ca sequestration, all conditions that enhance the propensity of Ca alternans. On the other hand, under $\beta$-adrenergic stimulation SERCA activity and consequently SR Ca uptake and load are increased, leading to enhanced fractional release that tends to promote alternans. Increased SERCA activity, however, also increases the efficiency of Ca sequestration, resulting in protection against alternans. Whether $\beta$-adrenergic stimulation favors (de Diego et al., 2008) or protects (Huser et al., 2000b) against alternans and alternans-related arrhythmias depends on which $\beta$-adrenergic effects predominate.

Another key controversial question in the alternans field is whether beat-to-beat alterations in SR Ca content are either causative (Diaz et al., 2004) or not associated with Ca alternans (Huser et al., 2000b). It has been suggested that diminished Ca reuptake and instability in the beat-to-beat feedback control of SR content leads to Ca alternans (Eisner et al., 2005). We addressed this question with a refined method of direct dynamic measurements of $[Ca]_{SR}$ during pacing-induced alternans (Picht et al., 2006). We found that diastolic $[Ca]_{SR}$ fluctuations can occur during alternans, where a large amplitude Ca release is preceded by a higher diastolic Ca content and vice versa. Interestingly, however, was our observation that Ca alternans can also readily occur without significant diastolic $[Ca]_{SR}$ alternations. These results suggest that factors other than SR Ca load, such as refractoriness (Kornyeyev et al., 2010; Rovetti et al., 2010) and restitution properties of ion currents and CICR from the SR via RyR, aberrant RyR openings (Armoundas, 2009) or, as shown here, metabolic activity and Ca cycling properties of mitochondria, are centrally involved in the mechanism underlying frequency-induced cardiac alternans.

LIMITATIONS OF THE STUDY AND CONCLUSIONS

Mitochondrial Ca signaling and energy metabolism are highly intertwined, thus the experimental dissection of the exact mechanism by which impaired mitochondrial function causes Ca alternans remains a difficult task. Our study relied exclusively on the pharmacological challenge of various mitochondrial targets ranging from mitochondrial ATP synthase, membrane potential, electron transport chain, Ca-dependent dehydrogenases to Ca uptake and extrusion pathways. While many of the pharmacological tools employed here are considered rather specific for the respective target, this approach still faces the difficulty that any interference with the metabolic function (ATP production) of mitochondria also affects mitochondrial and cellular Ca homeostasis, and vice versa any disturbance of mitochondrial Ca cycling interferes with mitochondrial energy metabolism (e.g., via Ca-dependent dehydrogenases). The oligomycin experiments (Figure 1B) clearly show that inhibition of mitochondrial ATP production enhances Ca alternans, thus sufficient ATP supply is protective against proarrhythmic Ca alternans. This finding is in line with our earlier studies where we showed that inhibition of glycolysis (and glycolytic ATP production) generates conditions that favor Ca alternans (Huser et al., 2000b; Kockskamper and Blatter, 2002; Kockskamper et al., 2005). Clearly, experimental interventions that impair mitochondrial Ca uptake and extrusion, and therefore mitochondrial Ca buffering, also enhance the propensity of Ca alternans (Figure 3). Whether this effect on Ca alternans occurs through a modification of cytosolic Ca transients or via alterations of Ca-dependent steps of mitochondrial energy metabolism remains to be determined. However, our experiments indicate that both are equally important. In line with the multifactorial nature of Ca and electro-mechanical alternans we can conclude from this study that sufficient mitochondrial energy (ATP) production and Ca buffering are both important and necessary safeguards against the occurrence of proarrhythmic Ca alternans.

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