Mitochondrial oxidative stress during cardiac lipid overload causes intracellular calcium leak and arrhythmia

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Abstract

Background—Diabetes and obesity are associated with an increased risk of arrhythmia and sudden cardiac death. Abnormal lipid accumulation is observed in cardiomyocytes of obese and diabetic patients, which may contribute to arrhythmia, but the mechanisms are poorly understood. A transgenic mouse model of cardiac lipid overload, the PPARg cardiac overexpression mouse, has long QT and increased ventricular ectopy.

Objective—We evaluated the hypothesis that the increase in ventricular ectopy during cardiac lipid overload is caused by abnormalities in calcium handling due to increased mitochondrial oxidative stress.

Methods—Ventricular myocytes were isolated from adult mouse hearts to record sparks and calcium transients. Mice were implanted with heart rhythm monitors for in vivo recordings.

Results—PPARg cardiomyocytes have more frequent triggered activity and increased sparks compared to control. Sparks and triggered activity are reduced by mitotempo, a mitochondrial-targeted antioxidant. This is explained by a significant increase in oxidation of RyR2. Calcium transients are increased in amplitude and SR calcium stores are increased in PPARg cardiomyocytes. Computer modeling of the cardiac action potential demonstrates that long QT contributes to increased SR calcium. Mitotempo decreased ventricular ectopy in vivo.
**Conclusions**—During cardiac lipid overload, mitochondrial oxidative stress causes increased SR calcium leak by oxidizing RyR2 channels. This promotes ventricular ectopy, which is significantly reduced in vivo by a mitochondrial-targeted anti-oxidant. These results suggest a potential role for mitochondrial-targeted anti-oxidants to prevent arrhythmia and sudden cardiac death in obese and diabetic patients.

**Keywords**
arrrhythmia; oxidative stress; calcium; lipid metabolism

**Introduction**

Obese and diabetic humans have increased rates of atrial fibrillation, ventricular ectopy, and long QT (LQT) \(^1-^3\). More importantly, diabetes and obesity are associated with an increased risk of sudden cardiac death, which is often caused by ventricular arrhythmias \(^4-^6\). The increased risk of sudden cardiac death is greater than the increased risk of myocardial infarction, suggesting that arrhythmic events are increased more than coronary events in obese and diabetic patients. For people with non-ischemic causes of sudden cardiac death, the cardiomyopathy of obesity is currently the most common autopsy finding \(^7\).

Abnormal lipid accumulation is observed in the cardiomyocytes of obese and diabetic patients and is thought to contribute to an increased risk of arrhythmia. This non-ischemic cardiomyopathy has been termed lipotoxic cardiomyopathy \(^8\), and is likely to become more prevalent as the demographics of Western societies shift to an older average age. However, the molecular mechanisms that cause arrhythmias during cardiac lipid overload have not been identified. Transgenic mouse models of cardiac lipid overload can provide mechanistic insight into the pathophysiology. Mice with cardiac-specific overexpression of peroxisome proliferator-activated receptor gamma (PPARg) accumulate lipids in their cardiomyocytes, without the systemic abnormalities of obesity or diabetes. PPARg is a transcription factor that regulates lipid metabolism. PPARg promotes lipid uptake by cells and is upregulated in the hearts of humans with metabolic syndrome \(^9\). PPARg mice have increased cardiac lipid content and abnormal cardiac mitochondrial morphology \(^10\). Specifically, PPARg mice have significantly increased cardiac triglycerides and free fatty acids \(^11\). Our previous work characterized the abnormal electrophysiology of PPARg transgenic mice, demonstrating that they have LQT caused by a decrease of voltage-gated potassium channels, with increased action potential duration (APD), frequent premature ventricular complexes (PVCs), and sudden cardiac death from spontaneous ventricular tachycardia as young adults \(^12\).

Since abnormalities of intracellular calcium homeostasis are a feature of several forms of arrhythmia, we hypothesized that the pathophysiology of ventricular ectopy in the lipotoxic heart would involve abnormalities in calcium handling. We used the PPARg mouse model to determine if abnormalities of intracellular calcium homeostasis play a causal role in ventricular ectopy in lipotoxic cardiomyopathy, and if abnormalities of calcium homeostasis are due to increased mitochondrial oxidative stress.
Methods

Mouse breeding and husbandry

Experiments conformed to the NIH Guide for the Care and Use of Laboratory Animals. The Columbia University institutional animal care review board gave approval. PPARg mice on the C57/BL genetic background were bred and genotyped as previously described. Mice were euthanized by anesthesia overdose (inhaled isoflurane) followed by cervical dislocation.

Spark recording and analysis

Sparks were recorded with a Leica SP2 confocal microscope equipped with a 63x1.4 NA objective. Isolated cardiac myocytes experiments were performed at room temperature. Cardiomyocytes were loaded with fluo-4 (5 nM 10 min), in modified Tyrode’s solution containing 1 mM calcium. Line scan images were recorded with Leica TCS software and quantified with the Sparkmaster plugin of ImageJ.

In vivo heart rhythm telemetry

Mice were implanted with heart rhythm monitors as previously described (Data Sciences International, model EA-F20). Isoflurane was used for general anesthesia. Mice recovered for one week after surgery before starting recordings. ECG recordings were made using Ponemah 3 software. A blinded observer tallied arrhythmia counts manually. Mitotempo was filter sterilized and given at 700 nanograms per gram of body weight as a daily intraperitoneal injection.

Computer Modeling of Action Potentials

A previously described model of the electrophysiology of the rat ventricular myocyte was adapted to study mouse electrophysiology, with updated sodium current model. PPARg overexpression genotype was simulated as a 30% reduction in Kv current density based on our prior publication. All results for action potentials represent steady state data for WT cardiomyocytes. In the model, PPARg cardiomyocytes showed oscillations of membrane potential and early afterdepolarizations, with no true steady state.

Statistical Analysis

Results are presented as mean ±SEM. The unpaired t-test was used for comparisons of two means; a 2-tailed value of \( P < 0.05 \) was considered statistically significant. For groups of 2 or more ANOVA was used with post-hoc testing (Prism v5, GraphPad Software).

Additional details about the methods are available in the online supplement.

Results

PPARg cardiomyocytes have increased oxidative stress

We predicted that cardiac lipid overload would cause increased oxidative stress in cardiomyocytes. To confirm increased cardiac reactive oxygen species (ROS) in our mouse model, we used the ROS indicator dihydrodichlorofluorescein diacetate (H2DCF-DA).
DCF fluorescence is a widely used as a measure of general oxidative stress in live cells. We compared the ROS levels in live adult mouse ventricular myocytes from WT control and PPARg mice. Isolated cardiomyocytes from PPARg mice had significantly increased DCF signal compared to control (figure 1A), indicating that PPARg cardiomyocytes have significantly greater ROS. Mitochondria are a major source of ROS in cardiomyocytes, and we found that the mitochondrial targeted antioxidant mitotempo reduced mitochondrial superoxide in the PPARg cardiomyocytes (figure 1B).

**PPARg cardiomyocytes have increased spontaneous contractions that are dependent on mitochondrial ROS**

To determine if the increased ventricular ectopy that we previously reported in PPARg mice could be observed at the level of the individual cardiomyocyte, we evaluated the frequency of spontaneous contractions after a short period of pacing in isolated cardiomyocytes, as described previously. After a short period of pacing at 1 Hz, PPARg cardiomyocytes that were not spontaneously contracting at baseline have significantly more contractions (fig 1B,C,D). This pattern is consistent with a triggered activity mechanism of ventricular ectopy. Since we suspected that increased oxidative stress could contribute to the mechanism, we inhibited mitochondrial ROS with the mitochondrial targeted antioxidant mitotempo for one hour. This treatment significantly decreased spontaneous contractions in PPARg cardiomyocytes (figure 1B) demonstrating that mitochondrial ROS promotes spontaneous contractions. We performed additional experiments with WT cardiomyocytes and mitotempo, which show that mitotempo does not have an effect on spontaneous contractions in control cardiomyocytes (sup figure 1).

**PPARg cardiomyocytes have increased spark frequency and oxidation of RyR2**

To directly evaluate SR calcium release in the PPARg heart, we measured spark frequency in isolated PPARg and WT control cardiomyocytes. PPARg cardiomyocytes had significantly more sparks than those from control animals (figure 2). This is significantly decreased by dantrolene, an inhibitor of RyR channels. To evaluate the role of mitochondrial ROS, we treated PPARg cardiomyocytes with the antioxidant mitotempo, which decreased sparks significantly in PPARg cardiomyocytes (figure 2B).

The increase in sparks indicates that PPARg cardiomyocytes have increased spontaneous calcium release. Increased SR calcium leak can be caused by increased phosphorylation or oxidation of ryanodine receptor RyR channels. Ventricular lysates were used to evaluate RyR2 oxidation status. We found a decrease in total RyR2 protein levels, but a significant increase in oxidation of RyR2 (figure 1E). Phosphorylation of RyR2 was not increased (sup figure 2).

**PPARg cardiomyocytes have increased calcium transient amplitude and increased SR calcium**

To evaluate contractility and calcium homeostasis in the lipotoxic cardiomyocytes, we measured sarcomere length and calcium transients using isolated cardiomyocytes, paced by field stimulation. Sarcomere length measurements show that relaxation is significantly prolonged in PPARg cardiomyocytes, as measured by the time constant tau of the return-to-
baseline curve (figure 3 and table 1). This is consistent with the prolonged action potential duration that we have previously reported in PPARg cardiomyocytes using patch clamp methods. Contractility, measured by fractional shortening, is similar in both groups, as is contraction velocity. Overall, isolated cardiomyocytes from young adult PPARg mice have properties consistent with normal systolic function and delayed relaxation.

Calcium transients were visualized in PPARg cardiomyocytes using fura-2. PPARg calcium transients are significantly increased in amplitude (figure 3B and table 1). This is consistent with increased SR calcium content. The application of caffeine, to release sarcoplasmic reticulum stores, shows a greater amount of calcium release from PPARg cells after controlling for baseline transient amplitude (figure 3C). The mean peak caffeine-evoked height normalized to paced transient height is 1.47 AU for WT, 2.12 for PPARg, (p=0.02 by t-test). This indicates a significant increase in sarcoplasmic reticulum (SR) calcium stores in PPARg cardiomyocytes.

**PPARg hearts have abnormal levels of calcium handling proteins**

Although an increase in L-type current could cause increased SR calcium loading, our prior work has shown that PPARg cardiomyocytes have normal L-type current densities and current-voltage relationships. We have also shown that PPARg cardiomyocytes do not have increased late sodium current. To better understand the molecular mechanisms of the observed abnormalities in calcium homeostasis, we evaluated the levels of cardiac proteins that are critical for calcium handling. Since SERCA2 pumps calcium back into the sarcoplasmic reticulum, an increase in SERCA could cause an increase in SR calcium load. However, SERCA2 protein is significantly reduced in the PPARg ventricles (figure 4). Phospholamban, which inhibits SERCA2 activity, has similar protein levels to control but phosphorylation was increased which would tend to increase SERCA2 activity, compensating for the decrease in SERCA protein level.

The sodium-calcium exchanger NCX1 removes calcium from the cytosol by extruding calcium from the cell. NCX1 protein is increased significantly in the PPARg heart, approximately 30% more than WT littermates. An increase in NCX1 may partially compensate for the increase in calcium entry from the prolonged APD. Increased NCX1 has been described in other forms of cardiomyopathy and may have a role in arrhythmia by depolarizing cardiomyocytes and initiating delayed afterdepolarizations. To evaluate another possible cause of SR calcium overload, we isolated SR membranes from mouse hearts and quantified calsequestrin protein, which is the major SR calcium binding protein in cardiomyocytes. Calsequestrin levels are similar in PPARg hearts and those from WT littermates, so this cannot explain the difference in SR calcium content.

**Computer modeling supports that LQT can cause SR calcium overload**

We suspected that the LQT phenotype of the PPARg model could be responsible for the increased intracellular calcium. Computer modeling was performed to investigate the relationship between long QT and SR calcium load. Our model shows that a minimal model of PPARg overexpression, simulated by a 30% reduction of WT Kv current alone, produces early afterdepolarizations during pacing at 200 BPM (Figure 5, panel A). We also explored
the impact of concomitant NCX upregulation, by increasing NCX density by 30% in the 
PPARg model. With this enhanced model, abnormalities in repolarization occur earlier in the 
drive train, suggesting that NCX serves a modulatory role in producing early 
afterdepolarizations (panel A, thin black trace).

Studying the beat immediately preceding afterdepolarization formation reveals that in 
PPARg cells, a significant increase in the plateau influx of calcium through L-type channels 
occurs (figure 5, panel B). This in turn causes an increase in cytoplasmic calcium (panel C) 
and ultimately early afterdepolarizations. This supports the hypothesis that a decrease in Kv 
current on the order of what we have observed experimentally can substantially prolong 
APD, as well as produce changes in calcium cycling and repolarization that are potentially 
pro-arrhythmic.

Mitochondrial anti-oxidant prevents ventricular ectopy in vivo

We hypothesized that the SR calcium overload contributed to arrhythmia in vivo. To test 
this, we implanted PPARg mice with heart rhythm telemetry monitors, allowing long-term 
heart rhythm recordings in unrestrained, non-sedated animals. We treated the mice with 
mitotempo. Mice were treated for 2 days during telemetry monitoring. Mitotempo resulted 
in a decrease in PVC frequency (figure 6). Mitotempo treatment did not change the QT (sup 
table 1). This in vivo telemetry experiment indicates that decreasing mitochondrial 
superoxide rapidly resets calcium handling to a less arrhythmogenic state.

Discussion

Since abnormalities of intracellular calcium homeostasis are a common feature of many 
forms of cardiomyopathy and arrhythmia, we hypothesized that the pathophysiology of 
ventricular ectopy in lipotoxic hearts would be caused by perturbations of calcium handling. 
We demonstrate that lipotoxic cardiomyocytes have increased ROS and abnormal calcium 
homeostasis, manifest by increased spark frequency and SR calcium overload. We also show 
that cardiac lipotoxicity causes increased spontaneous contractions, which are improved by 
inhibition of mitochondrial ROS. These cellular abnormalities promote ventricular ectopy, 
which is improved in vivo by treatment with the mitochondrial anti-oxidant mitotempo.

Calcium handling and ROS

We show that sparks and spontaneous contractions are improved with inhibition of 
mitochondrial ROS, identifying mitochondrial ROS is a mechanistic link between 
lipotoxicity and abnormal calcium homeostasis. Prior work had proposed that ROS increases 
SR calcium leak in isolated cardiomyocytes, using cells from aged animals (which have 
increased ROS) or using pharmacologic manipulations to increase ROS [26, 27]. Our work 
shows that this mechanism is important for cardiac lipid overload. Furthermore, we show 
that the mechanism of increased RyR2 calcium leak is oxidation of the RyR2 channel rather 
than increased phosphorylation.
SR calcium and LQT

The abnormalities identified in this study are in contrast to those using systolic heart failure models, which typically have decreased SR calcium content and lower amplitude calcium transients. Paradoxically, despite the increase in SR calcium leak, we show a significant increase in SR calcium in the transgenic model. Young adult PPARg mice represent an early form of cardiomyopathy, with compensated systolic function. We had previously shown that PPARg cardiomyocytes have prolonged APD due to decreased voltage-gated potassium channel currents, with normal L-type current densities and current-voltage relationships.

It is likely that the LQT phenotype contributes to increased SR calcium. Our minimal computational model demonstrates that when the reduction in potassium current density is simulated in isolation, the consequent APD prolongation and corresponding increase in calcium influx during the action potential can increase SR load and calcium transient amplitudes, providing an explanation for pathologic calcium overload. There is precedent for this model. Prior work by others has shown that prolongation of the APD can cause increase SR calcium load. LQT may promote ventricular tachycardia by increasing calcium entry during the prolonged repolarization phase; prolongation of the action potential duration increases calcium entry into the cardiomyocyte cytosol. Prolonged QT is an independent risk factor for cardiovascular mortality in heart failure patients and also in diabetics without heart failure. Obese humans have prolonged QT compared to normal-weight controls, highlighting the potential clinical relevance of our findings.

The increase in NCX1 may contribute to ventricular ectopy. Although SERCA2 function is the dominant mechanism for calcium extrusion in healthy mouse myocytes, during calcium overload, an increase in NCX1 could be protective by extruding excess calcium. However, during spontaneous calcium release events, a larger NCX current can theoretically promote arrhythmias by producing the depolarizing transient inward current that is postulated to be the ionic basis of delayed afterdepolarizations. Our findings of altered calcium cycling with PPARg overexpression are broadly consistent with prior experiments using Kv current blockade in a canine preparation.

Ventricular Ectopy in Lipotoxic Cardiomyopathy

There is prior evidence that ventricular tachycardia in non-ischemic cardiomyopathy is caused by triggered activity rather than reentry, which is the major mechanism in scar-related ventricular tachycardia (such as after myocardial infarction). The increase in spontaneous contractions in isolated cardiomyocytes from PPARg mice observed in this study is consistent with triggered activity. The increase in ventricular ectopy correlates with an increase in single-cell spontaneous contractions and increased spark frequency. In combination with increased spark frequency, the increase in calcium transient amplitude may contribute to ventricular tachycardia in the PPARg mice. The improvement in heart rhythm during short-term treatment with mitotempo supports the idea that mitochondrial ROS promotes ventricular ectopy during lipotoxicity. Identifying the molecular mechanisms causing electrophysiologic abnormalities and arrhythmias in lipotoxic cardiomyopathy may allow for targeted therapy to prevent arrhythmia and sudden cardiac death in obese and diabetic patients.
The emerging connections between cardiac metabolism, circadian rhythm, and heart rhythm are intriguing. While this is beyond the scope of the present manuscript, the PPARg mouse model could be a useful tool for future investigation in this area.

Conclusions

During cardiac lipid overload, mitochondrial oxidative stress causes increased SR calcium leak by oxidizing RyR2 channels. This promotes ventricular ectopy, and heart rhythm is improved in vivo by a mitochondrial-targeted anti-oxidant. These results suggest a potential role for mitochondrial-targeted anti-oxidants to prevent arrhythmia and sudden cardiac death in obese and diabetic patients.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

Acknowledgments

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Abbreviations

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<tr>
<th>Abbreviation</th>
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<tr>
<td>APD</td>
<td>action potential duration</td>
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<tr>
<td>DCF</td>
<td>dichlorofluorescein</td>
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<tr>
<td>NCX</td>
<td>sodium calcium exchanger</td>
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<tr>
<td>PPARg</td>
<td>peroxisome proliferator-activated receptor gamma</td>
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<tr>
<td>PVC</td>
<td>premature ventricular complex</td>
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<tr>
<td>ROS</td>
<td>reactive oxygen species</td>
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<tr>
<td>RyR</td>
<td>ryanodine receptor channels</td>
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<tr>
<td>SR</td>
<td>sarcoplasmic reticulum</td>
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<td>WT</td>
<td>wild-type</td>
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References


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Figure 1. PPARg cardiomyocytes have increased ROS and triggered activity
A. ROS in isolated cardiomyocytes; height is DCF fluorescence minus background. ROS is reduced by mitotempo in PPARg cardiomyocytes. MT = mitotempo 20 μM. The means are significantly different by ANOVA, *= sig different from control by post-hoc test. Readings were performed in triplicate with 2000 cells/well from 3 animals of each type.
B. Graph of triggered activity from PPARg and control cardiomyocytes. The means are significantly different by ANOVA, *= sig different from control by post-hoc test, n=21–35 cells from 3 animals of each type.
C. Representative example of isolated control cardiomyocyte, paced without subsequent activity
D. Representative example of isolated PPARg cardiomyocyte with triggered activity after pacing.
Figure 2. PPARg cardiomyocytes have increased calcium sparks
A. Representative examples of sparks raw data from control and PPARg cardiomyocytes
B. Graph of sparks in WT control and PPARg cardiomyocytes, normalized to control. Dantrolene and mitotempo reduce spark frequency. n=11–20 cells each from 3 animals of each type, the means are significantly different by ANOVA, *= sig different by post-hoc test, dan=dantrolene 1 μM, MT = mitotempo 20 μM
C. Western blots of oxy-RyR2 and total RyR2, and graphs of mean total RYR2 and oxyRyR2 normalized to total RyR2. *= p<0.05
Figure 3. Abnormal relaxation and calcium transients in PPARg cardiomyocytes
A. Representative sarcomere length recordings from isolated myocytes from PPARg and WT littermates. Note the slower return to diastolic baseline in the PPARg cell. Marks at bottom show pacing (1 Hz) for time scale.
B. Representative calcium imaging from isolated myocytes from PPARg and WT littermates loaded with fura-2. The tracings show increased amplitude and slower return to baseline in PPARg cardiomyocytes, indicated increased calcium SR release and decreased SERCA2 function.
C. Representative caffeine-evoked transients from WT and PPARg cardiomyocytes. Cells were paced to achieve steady-state calcium levels, and then caffeine (20 mM) was rapidly perfused. The height of the caffeine-induced calcium signal, relative to the paced transient, is greater in the PPARg cell, indicating greater SR calcium content.
Figure 4. PPARg mice have abnormal levels of calcium handling proteins
A. Representative immunoblots of SERCA2, NCX1, and PLN proteins
B. Graph of protein levels normalized to control levels after adjusting for tubulin loading, AU= arbitrary units. n=4 animals of each type, t-test: *= p<0.05, **= p<0.01
C. Membrane preparation western blot from PPARg and WT littermate hearts show equivalent amounts of calsequestrin protein. KDEL is a loading control from the same membrane.
Figure 5. Computational modeling of rodent myocyte electrophysiology predicts abnormal repolarization and calcium handling in the PPARg mouse
Panel A shows action potentials generated at 200 BPM for three models: wild type (gray trace), minimal model of PPARg overexpression incorporating only Kv current downregulation (thick black trace), second model incorporating both Kv current downregulation and NCX upregulation (thin black trace). For pacing to steady state, the WT model does not produce EADs. In contrast, for both PPAR gamma models, early afterdepolarizations are observed, with failure of the action potential to repolarize. NCX upregulation modulates the generation of EADs, as failure to repolarize occurs earlier in the pacing drive train. Panels B and C show the L type current and the calcium transient for the beat immediately preceding EAD formation, for the PPAR gamma minimal model (thick black trace) and corresponding WT action potential (gray trace). Substantially higher plateau inward current is noted (panel B), with larger calcium transients (panel C).
Figure 6. Mitotempo improves heart rhythm in PPARg mice
Mean PVC count from PPARg mice, at baseline and during mitotempo treatment. N = 4 animals. The means are significantly different by ANOVA, *= sig different by post-hoc test.
Table 1
Contractility and calcium transient data
For cell shortening, n= 38–40 cells, from 3–4 animals each group. For calcium transients, n= 40–70 cells from 3–4 animals each group.

<table>
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<tr>
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<th>mean + SEM</th>
<th>con</th>
<th>PPARg</th>
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<td><strong>Cell Shortening</strong></td>
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<td>% FS</td>
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<td>12.53 + 0.46</td>
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<tr>
<td>tau</td>
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<td>0.085 + 0.006</td>
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<td><strong>Ca2+ transient</strong></td>
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<td>diastolic signal</td>
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<td>0.55 + 0.02</td>
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<td>100.9 + 4.0 *</td>
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<tr>
<td>tau (sec)</td>
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<td>0.25 + 0.01 *</td>
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* sig different from control by t-test