Remodeling of T-Tubules and Reduced Synchrony of Ca\(^{2+}\) Release in Myocytes From Chronically Ischemic Myocardium

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Abstract—In ventricular cardiac myocytes, T-tubule density is an important determinant of the synchrony of sarcoplasmic reticulum (SR) Ca\(^{2+}\) release and could be involved in the reduced SR Ca\(^{2+}\) release in ischemic cardiomyopathy. We therefore investigated T-tubule density and properties of SR Ca\(^{2+}\) release in pigs, 6 weeks after inducing severe stenosis of the circumflex coronary artery (91 \pm 3\%, N=13) with myocardial infarction (8.8 \pm 2.0\% of total left ventricular mass). Severe dysfunction in the infarct and adjacent myocardium was documented by magnetic resonance and Doppler myocardial velocity imaging. Myocytes isolated from the adjacent myocardium were compared with myocytes from the same region in weight-matched control pigs. T-tubule density quantified from the di-8-ANEPPS (di-8-butyl-aminonaphthyl-ethylene-pyridinium-propyl-sulfonate) sarcolemmal staining was decreased by 27 \pm 7\% (P<0.05). Synchrony of SR Ca\(^{2+}\) release (confocal line scan images during whole-cell voltage clamp) was reduced in myocardium myocytes. Delayed release (ie, half-maximal [Ca\(^{2+}\)], occurring later than 20 ms) occurred at 35.5 \pm 6.4\% of the scan line in myocardial infarction versus 22.7 \pm 2.5\% in control pigs (P<0.05), prolonging the time to peak of the line-averaged [Ca\(^{2+}\)] transient (121 \pm 9 versus 102 \pm 5 ms in control pigs, P<0.05). Delayed release colocalized with regions of T-tubule rarefaction and could not be suppressed by activation of protein kinase A. The whole-cell averaged [Ca\(^{2+}\)] transient amplitude was reduced, whereas L-type Ca\(^{2+}\) current density was unchanged and SR content was increased, indicating a reduction in the gain of Ca\(^{2+}\)-induced Ca\(^{2+}\) release. In conclusion, reduced T-tubule density during ischemic remodeling is associated with reduced synchrony of Ca\(^{2+}\) release and reduced efficiency of coupling Ca\(^{2+}\) influx to Ca\(^{2+}\) release. (Circ Res. 2008;102:338-346.)

Key Words: myocardial infarction ■ contractility ■ myocytes ■ calcium

Although new therapeutic approaches have decreased the mortality associated with myocardial infarction (MI) over the past decades,\(^1\) many patients nevertheless sustain a regional loss of myocardial contractile tissue following an ischemic event. The resulting increased hemodynamic burden on the left ventricle leads to structural and functional changes in the remaining viable myocardium, which further reduces ventricular performance, a process referred to as myocardial remodeling.\(^2\) Sustained regional chronic and/or intermittent ischemia further contributes to this process, and the resulting ischemic cardiomyopathy is currently among the major causes of heart failure.\(^3\)

Contractile dysfunction of the ventricle is partly related to the abnormal loading in vivo\(^4\) and partly to the intrinsic properties of the cardiomyocytes. Myocytes isolated from patients with ischemic cardiomyopathy at the time of heart transplantation have a reduced contractile function resulting from abnormal Ca\(^{2+}\) handling.\(^5\)–\(^7\) Animal models have examined the mechanisms of cellular dysfunction in ischemic cardiomyopathy in more detail. Myocytes from the infarct border zone have a reduced contraction and slowed and reduced [Ca\(^{2+}\)] transients.\(^8\)–\(^11\) The mechanisms leading to this impaired Ca\(^{2+}\) handling are not completely understood. Decreased intracellular Ca\(^{2+}\) release may be the result of a reduced sarcoplasmic reticulum (SR) Ca\(^{2+}\) content caused by decreased SR Ca\(^{2+}\) pump activity, as has been observed in some\(^8,11,12\) but not all\(^10,13\) models of postinfarct remodeling. Increased Na/Ca exchange (NCX) activity and abnormal expression and phosphorylation of the ryanodine receptor (RyR) have also been reported.\(^10,14,15\)
Ca$^{2+}$ T-tubule density indeed leads to dyssynchronous intracellular decrease in the density of T-tubules in tissue sections.\textsuperscript{26} but the contribution of alterations in T-tubules to myocardial dysfunction and limited infarction without heart failure.\textsuperscript{29–31}

Studies on intact living human ventricular myocytes have not documented small transmural infarctions; their functional character-istics are shown under Results.\textsuperscript{18,19} In end-stage human heart failure, histological exam-
ination showed dilation of T-tubules\textsuperscript{25,26} with an increase\textsuperscript{25} or a reduction of the L-type Ca$^{2+}$ current ($I_{CaL}$) was seen; such a reduced $I_{CaL}$ has been reported in other\textsuperscript{18,19} but not all models of MI.\textsuperscript{20}

Even in the presence of a normal SR Ca$^{2+}$ content, defective coupling between Ca$^{2+}$ influx and activation of RyR could lead to reduced Ca$^{2+}$ release.\textsuperscript{16} Dyssynchronous opening of RyRs and decreased [Ca$^{2+}$], transients were observed in a rabbit model of postinfarct remodeling,\textsuperscript{17} and, in this model, a reduction of the L-type Ca$^{2+}$ current ($I_{CaL}$) was seen; such a reduced $I_{CaL}$ has been reported in other\textsuperscript{18,19} but not all models of MI.\textsuperscript{20}

Efficient coupling of Ca$^{2+}$ influx through sarcolemmal Ca$^{2+}$ channels (dihydropyridine receptors) and activation of RyRs in the SR is related to the structural organization in coupions, which, in ventricular cardiomyocytes, are found to a large extent along the T-tubules.\textsuperscript{21} Experimentally reducing T-tubule density indeed leads to dyssynchronous intracellular Ca$^{2+}$ release and reduced [Ca$^{2+}$], transients.\textsuperscript{22–24} A loss of T-tubules could be part of the postinfarction remodeling process. In end-stage human heart failure, histological examination showed dilatation of T-tubules\textsuperscript{25,26} with an increase\textsuperscript{25} or decrease\textsuperscript{26} in the density of T-tubules in tissue sections. Studies on intact living human ventricular myocytes have not yet been conclusive as to whether T-tubule density is altered,\textsuperscript{24,27,28} but the contribution of alterations in T-tubules to remodeling is supported by observations in animal models of heart failure.\textsuperscript{29–31}

In this study, we investigated potential changes in T-tubule density and their impact on intracellular Ca$^{2+}$ release in a pig model of severe chronic coronary stenosis with regional myocardial dysfunction and limited infarction without heart failure.

### Animal Model

A detailed description is provided in the online data supplement. Briefly, a copper-coated stent was inserted into the proximal circumflex artery of young domestic pigs (20 to 25 kg) to induce intima proliferation and severe nonthrombotic coronary stenosis, quantified during coronary angiography. Global and regional left ventricular (LV) function were evaluated at baseline and at 3 and 6 weeks after stent implantation using color Doppler myocardial velocity imaging at baseline and during dobutamine stress. In a number of animals, MRI was used to assess global and regional LV function and to quantify MI as a fraction of total LV mass. The stent implantation induces a severe but slightly variable stenosis and degree of LV dysfunction. In the present study, we used only animals with documented small transmural infarctions; their functional characteristics are shown under Results.

At the time of euthanasia (47±1 days after stent implantation), pigs with MI (N=13) weighed 51±2 kg; matched healthy pigs were used as controls (CTRL) (N=15; weight, 55±4 kg).

### Isolation of Cardiac Myocytes

The procedure for isolating myocytes was as described previously.\textsuperscript{32} After enzymatic digestion of the tissue wedge perfused by the stenotic artery, the infarct area and 5 to 10 mm around the infarct core were discarded; midmyocardial cells from the remaining digested tissue were used. Myocytes isolated from the midmyocardial layer of the posterior wall of the weight-matched healthy pigs served as controls.

### Subcellular [Ca$^{2+}$] Measurements and T-Tubule Quantification

The setup for confocal imaging was as described previously, and image acquisition and analysis is detailed in the online data supplement. [Ca$^{2+}$], transients were recorded during steady-state stimulation at 1 Hz, with depolarizing steps from −70 to 0 mV for 150 ms along a line through the center parallel to the long axis of the cell as described previously.\textsuperscript{23,33} Temporal resolution was 1.54 ms per line, and pixel size (x, y) was 0.2 to 0.4 μm. For T-tubule density (di-8-ANEPPS [di-8-butyl-amino-naphthyl-ethylene-pyridinium-
propyl-sulfonate] staining) 8 to 10 sequential xy images centered around the equatorial plane of the cell were recorded from each cell, with a spacing of 1 μm in the z-direction and a pixel width of 0.11 to 0.20 μm.

Global [Ca²⁺] and Membrane Current Measurements

The setup for epifluorescence recording, protocols, and solutions are described in the online data supplement.

Immunofluorescence Imaging and Western Blot

The procedures are described in the online data supplement. For protein expression, we used transmural needle biopsies from the ischemic area taken in situ at the time of euthanasia. Control tissue was obtained from the same area in hearts of CTRL pigs.

Image Analysis

Algorithms to analyze the spatial and temporal characteristics of the [Ca²⁺], transients were custom-made (IDL 6.1, Research Systems International, Paris, France); the approaches are detailed in the online data supplement. All temporal data refer to the onset of the whole-line averaged [Ca²⁺] transient. \( F_{50} \) was defined as the half-maximum of the normalized overall peak [Ca²⁺]-dependent fluorescence and served as a threshold to discriminate local Ca²⁺ release as described previously.\(^{33}\) To evaluate T-tubules, the images were deconvolved and automatically thresholded; the sarcolemmal surface membrane was excluded from analysis in all images (more details are available in online data supplement). T-tubule density is expressed as the fraction of positive pixels of all pixels within the sarcolemmal boundaries; data are per cell and subsequently pooled per animal.

Statistics

Data are shown as means±SEM and were compared using Student’s t test or 2-way ANOVA; Fisher’s least-significant difference test was performed when significant overall effects were detected. \( P<0.05 \) was considered significant.

Results

Global and Regional LV Function In Vivo

The degree of circumflex arterial stenosis was 91±3% of the vessel lumen. At 6 weeks, LV ejection fraction at rest, as determined from MRI, was slightly but significantly decreased in MI versus CTRL (Figure 1A); the LV end-diastolic volume was not increased (116±10 mL in MI versus 106±10 mL in CTRL, \( P=0.1 \)). Infarction, as determined from the delayed enhancement images (Figure 1B), comprised 8.8±2.0% of the total LV mass; this corresponds to \( \approx \)50% of the area at risk. MRI confirmed the regional dysfunction in the segments with delayed enhancement as well as in the adjacent segments within the area subtended by the stenotic artery (Figure 1B).

Myocardial velocity imaging during follow-up showed the regional dysfunction present at 3 weeks as a severely reduced systolic strain rate (Figure 1C). At 6 weeks, there was no inotropic response to dobutamine (Figure 1D), consistent with the presence of transmural MI.\(^{34}\)

(Ultra)structural Remodeling of Myocytes

Cardiomyocytes from the MI adjacent area were hypertrophied, as indicated by a significant increase in cell length: 186±6 μm in MI (\( N_{\text{animals}}=9, n_{\text{cells}}=254 \)) versus 147±6 μm in CTRL (\( N_{\text{animals}}=6, n_{\text{cells}}=180, P<0.05 \)). Cell width and cell depth tended to be increased as well (29±2 versus 28±1 μm in CTRL, \( P=0.2 \); 22±1 versus 18±1 μm in CTRL, \( P=0.07 \)). The calculated increase of cell volume assuming a brick-like shape for MI was 51%. This contrasts with the increase of total, ie, external and T-tubular, membrane area measured from the cell capacitance of 30% from 91±5 pF (CTRL: \( N_{\text{animals}}=9, n_{\text{cells}}=65 \)) to 118±8 pF (MI: \( N_{\text{animals}}=5, n_{\text{cells}}=43, P<0.05 \)).
T-tubule density expressed as the fraction of positive pixels of all pixels within the sarcolemmal boundaries was significantly reduced in MI (Figure 2). Expressed as a fraction of the external sarcolemmal boundary, the values were also close to significance (0.77±0.05 versus 0.92±0.06 μm⁻¹ in CTRL, P=0.057). In contrast, the intracellular distribution of RyR appeared homogeneous and similar in CTRL and MI (Figure 2C). This was quantified by measuring RyR signal density, which was similar in cells from CTRL (N_{animals}=3, n_{cells}=17) and MI (N_{animals}=5, n_{cells}=17; Figure 2D). We also examined the variability in either RyR or T-tubule signal in individual 5×5 μm boxed areas of a cell (detailed in the online data supplement), as regional loss would result in a higher variability. Here, again, there was a significant difference between MI and CTRL for T-Tubules but not for RyR (Figure 2E).

**Reduced Synchrony of Ca²⁺ Release**

During confocal line scan imaging of the [Ca²⁺], transient, myocytes from MI had numerous areas with delayed Ca²⁺ release, persistently present during consecutive pulses, as evident in the [Ca²⁺], transient averaged over 10 beats (Figure 3A). The absence of beat-to-beat variability is further illustrated in Figure 3B, showing the line at 17 ms during sequential beats and the small extent of variability in this line. In myocytes in which the sarcolemma was stained with wheat germ agglutinin (WGA)-Alexa594, a first line scan using the 543 nm excitation showed T-tubules as continuous lines of constant intensity (Figure 3C, left); during the subsequent line scan using the 488 nm excitation of fluo-3, the regions of delayed Ca²⁺ release could be related to areas of T-tubule rarefaction (Figure 3C, right). Figure 3D shows another cell with the 2D WGA-Alexa594 image and the area from which the recording was made.

We hypothesized that these regions of delayed Ca²⁺ release lead to a slowed upstroke of the [Ca²⁺] transient in MI. This was quantified by measuring for each consecutive line scan the fraction of the line that had a fluorescence larger than 50%. As shown in Figure 4A, in MI cells, this proceeded more slowly than in CTRL, and, at 20 ms, Ca²⁺ release remained below F_{0} along 36.7±5.7% of the line in MI versus only 22.7±2.5% of the line in CTRL (P<0.05). Likewise, the time to peak of the [Ca²⁺] transient, spatially averaged over the entire line, was longer in MI (122±8 ms, N_{animals}=8, n_{cells}=35 versus 102±5 ms, N_{animals}=13, n_{cells}=41 in CTRL, P<0.05).

Increased inhomogeneity in Ca²⁺ release in MI cells was attributable to a larger number (Figure 4B), as well as a larger size (Figure 4C), of the regions of delayed Ca²⁺ release.

We further examined the properties of Ca²⁺ release within both early and delayed regions. Time to local peak [Ca²⁺], in regions of early Ca²⁺ release was not different in MI versus CTRL (66±9 versus 74±8 ms, n_{cells}=17 and 14), and amplitude was also not different (F/F_{0} 3.2±0.2 versus 3.4±0.3). In areas of delayed release, the Ca²⁺ release was shown previously to propagate in a wave-like manner because of Ca²⁺-induced Ca²⁺ release. In the current study, this process was quantified as the rate of filling (decrease in width as a function of time) for regions...
of delayed Ca\textsuperscript{2+} release with a width of >5 \( \mu \)m (\( n=35 \) in MI and \( n=20 \) in CTRL), as illustrated in Figure 5A. No difference in rate of filling was found between MI and CTRL (Figure 5B).

We also examined whether cAMP-mediated positive inotropic stimulation could synchronize Ca\textsuperscript{2+} release in early and delayed Ca\textsuperscript{2+} release sites. As demonstrated in Figure 6A and 6B, the amplitude of Ca\textsuperscript{2+} release was greatly increased in the presence of forskolin (top); the extent and distribution of delayed Ca\textsuperscript{2+} release sites however appeared unchanged (bottom). Quantitative analysis of local [Ca\textsuperscript{2+}], transients (\( n_{\text{cells}}=9 \)) confirmed that the fraction of delayed release areas was unchanged (41\%±7\% versus 33\%±6\% at baseline). Time to peak [Ca\textsuperscript{2+}], in regions defined as delayed before the addition of forskolin remained significantly longer than in early regions also in the presence of the drug. The response to forskolin was not different in MI from CTRL (Figure 6C).

**Reduced Overall Gain of SR Ca\textsuperscript{2+} Release in MI**

In a subset of animals (\( N=4 \) for MI, \( N=5 \) for CTRL), the whole-cell averaged [Ca\textsuperscript{2+}], transients were studied in a confocal fluorescence setup. Peak [Ca\textsuperscript{2+}], during steps from \(-70 \) to \(+10 \) mV, as used in the confocal analysis, was lower in MI (\( n_{\text{cells}}=12 \)) versus CTRL (\( n=15, P<0.05 \); Figure 7A), and the time to peak [Ca\textsuperscript{2+}], was prolonged (Figure 7B). The density of the L-type Ca\textsuperscript{2+} current was not significantly different (Figure 7C).

Because the [Ca\textsuperscript{2+}], transients were of lower amplitude despite unchanged Ca\textsuperscript{2+} current, a reduction of SR content was expected, and this was tested using caffeine to release all SR Ca\textsuperscript{2+}; Figure 8A shows the averaged signals of [Ca\textsuperscript{2+}], and inward NCX current for all MI (\( n_{\text{cells}}=12 \)) and CTRL myocytes (\( n_{\text{cells}}=15 \)). The integral of the inward NCX current (\( I_{\text{NCX}} \)) during caffeine application (Figure 8B) was unexpectedly increased in MI versus CTRL, suggesting an increased Ca\textsuperscript{2+} content of the SR. In contrast, the caffeine-induced peak [Ca\textsuperscript{2+}], was reduced in MI (Figure 8C). These latter observations may rather reflect altered kinetics of Ca\textsuperscript{2+} release from the SR in response to caffeine in MI cells, as time to peak (Figure 8D) of the caffeine-induced [Ca\textsuperscript{2+}], transient was significantly increased in MI versus CTRL. The rate of decay was also significantly increased (Figure 8E), reflecting a reduced rate of Ca\textsuperscript{2+} removal by the NCX. Yet global protein expression of the NCX was unaltered (Figure 8F).

We analyzed the properties of spontaneous release events or sparks, observed in a 15-second period following stimulation at 1 Hz (for MI: \( N_{\text{animals}}=7, n_{\text{cells}}=36, 2472 \) sparks; for CTRL: \( N_{\text{animals}}=12, n_{\text{cells}}=44, 2280 \) sparks). In MI animals, 100\% of cells had spontaneous release events versus 74\% in CTRL (\( P<0.05 \)), but analysis of mean frequency was not statistically different (1.69±0.35 versus 1.27±0.23 sparks/sec per 100 \( \mu \)m in CTRL). Amplitude and duration were also not statistically different (\( F/F_0 \) 1.76±0.05 versus 1.70±0.06
in CTRL; full duration at half-magnitude, 39±2 versus 36±2 ms in CTRL).

Discussion

In Vivo Function and Remodeling With Chronic Coronary Stenosis

The most commonly used models for cellular remodeling after MI rely on smaller animals with ligation of a coronary artery, usually the left anterior descending. In these animals, MI typically is large, leading to heart failure (eg, in the rat and rabbit). In the current animal model, we have a limited infarction within a larger area of chronic underperfusion; at 6 weeks, the impact on global function is present but limited. Therefore, this model allows us to study the changes with chronic ischemia in the area adjacent to the infarct, before the onset of heart failure. One can then question whether this is essentially different from the condition we have studied previously, namely hibernation, where there is no or only very limited subendocardial necrosis. We think, at present, these data should not be mixed. The regional contractile function in the hibernating myocardium is less affected because the loss of contractile cells is small. This implies that mechanical loading on the remaining myocytes is less than in the presence of MI, which could lead to different stimuli for remodeling.

We currently have a limited number of data to sustain that there is a difference in cellular remodeling in cardiomyocytes from hibernating myocardium as compared with cells from the area adjacent to the infarct. At the whole-cell level, the overall \([\text{Ca}^{2+}]\) transients are more affected in the present dataset than in our earlier group of hibernation, although the \([\text{Ca}^{2+}]\) current in the present study was less affected. There are also a number of similarities because SR \([\text{Ca}^{2+}]\) content was similarly preserved. We currently have no data on subcellular synchrony of release in hibernating myocardium, but preliminary data also have shown a reduction in T-tubule density. Cellular hypertrophy is present in both groups.

Remodeling of T-Tubules

Ventricular myocytes develop T-tubules after birth, whereas atrial myocytes or Purkinje cells do not. The density of the T-tubule system is variable between species, and T-tubules disappear during culture of adult cardiac myocytes. These observations indicate that the T-tubule system is under active regulation and that it could change with disease. Yet the observations of remodeling with disease are limited. The present data document a lower density of...
T-tubules during ischemic remodeling before the development of heart failure.

This does not necessarily represent a net loss. Indeed, the myocytes were clearly hypertrophied, and the lower density of T-tubules could represent a differential growth and organization of this organelle in relation to the addition of sarcomeres. A striking finding was also that the distribution of RyR was unchanged. The data thus support the idea that T-tubules have a degree of plasticity and can undergo remodeling independent of the organization of the SR.

Song et al observed reorganization of T-tubules in spontaneously hypertensive rats, with a decrease in the transverse tubules in favor of an increase in the longitudinal tubules; the authors did not report a decrease in overall density. Similar observations were made in the mouse. Song et al observed reorganization of T-tubules in spontaneously hypertensive rats, with a decrease in the transverse tubules in favor of an increase in the longitudinal tubules; the authors did not report a decrease in overall density. The rat, like the mouse, has a very dense network, possibly related to high heart rates, whereas pigs have a lower density, more like humans. In the latter system, reorganization may be more likely to result in true loss of T-tubules in certain areas.

**Reduced Synchrony of Ca²⁺ Release**

Litwin et al were the first to describe dyssynchronous Ca²⁺ release in myocytes from failing rabbit hearts after MI. In their study, Ca²⁺ release had a beat-to-beat regional variation, and release events were observed to occur late during depolarization. This dyssynchrony was linked to a reduced \( I_{CaL} \) and the presence of functional couplings with decreased triggering; it could be rescued at least partially by protein kinase A–dependent phosphorylation and increased activity of \( I_{CaL} \). In a model of right ventricular failure, a reduction in \( I_{CaL} \) associated with changes in early repolarization also resulted in a regional beat-by-beat variation in intracellular Ca²⁺ release. The current study points to a different mechanism underlying reduced synchrony of Ca²⁺ release. First, we studied Ca²⁺ release during square voltage-clamp pulses, eliminating the influence of potential changes in early repolarization. Second, we focused on regions where Ca²⁺ release was persistently delayed in sequential [Ca²⁺]i transients, independent of beat-to-beat variability. We could not “rescue” the delayed release by increasing cAMP. In contrast, we could link the delayed release sites to areas of low T-tubule density. These data are consistent with a structural alteration.

Recent computational analysis supports the importance of small changes in the structures relating dihydropyridine receptor and RyR. We also considered the possibility of a reduction in couplon size, which would result in a longer latency and reduced probability of release at spark sites. This additional analysis is presented in the online data supplement. The data suggest that, in addition to the larger number of delayed release sites, increased dyssynchrony at presumed spark sites is present. This could result from a smaller couplon size.
Additional experiments are needed to evaluate the contribution of these changes to the overall reduced synchrony.

**Coupling Efficiency Between Dihydropyridine Receptor and RyR and the Role of T-Tubules**

The consequences of reduced synchrony are a global slowing of the rate of rise of [Ca\(^{2+}\)]\(_i\), as also observed in the spatially averaged whole-cell [Ca\(^{2+}\)]\(_i\) transients. In such recordings, the longer time to peak, in particular together with the reduced amplitude of the [Ca\(^{2+}\)]\(_i\) transient, indicates that, overall, the SR Ca\(^{2+}\) release is reduced, despite preserved global Ca\(^{2+}\) influx and SR Ca\(^{2+}\) content. This can be interpreted as a reduced efficiency of coupling, as was initially proposed in the rat with hypertension-induced heart failure.

Reduced coupling efficiency in MI may be the result of a smaller fraction of existing RyRs that are activated, because of a lack of T-tubules and possibly because of a reduction in couplon size. The distribution of RyRs was preserved, and the propagation of Ca\(^{2+}\) release within the delayed areas was the same for MI and CTRL, as were the properties of sparks, suggesting that the intrinsic properties of the RyR were not altered.

The presence of RyRs that are not coupled to the dihydropyridine receptor has analogies with experimentally reducing the probability of activating the RyR, eg, with low doses of caffeine or by reducing the activation of Ca\(^{2+}\) channels.

Under those conditions, the SR can (locally) become overloaded, and this can give rise to even more pronounced heterogeneity in release under the form of alternans. This was currently not observed, but the larger SR content could be the result of reduced RyR activation.

The slower decay of the caffeine transient points toward a reduced efficiency in Ca\(^{2+}\) removal. This was also noticed in studies with acute detubulation and during cell culture. The reduced rate of rise of [Ca\(^{2+}\)]\(_i\), with caffeine application in MI myocytes might be related to reduced T-tubule density, because we likewise noted a significant increase in time to peak of 50%, with a 60% reduction of T-tubule density in cultured myocytes (data from Louch et al)[24]. Alternatively, this may indicate additional changes in RyR properties, with a reduced response to caffeine.

**Conclusions**

In myocytes from the area adjacent to MI, decreased T-tubule density is associated with reduced global coupling efficiency, reduced synchrony of Ca\(^{2+}\) release, and slowed and reduced cellular [Ca\(^{2+}\)], transients. This occurs in the absence of changes in distribution of the RyR and indicates that plasticity of T-tubules is an independent factor in the remodeling process.

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**Disclosures**

None.

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Materials and Methods

Animal Model

Animals were housed and treated according to the Guide for the Care and Use of Laboratory Animals (National Institute of Health, U.S.A.) and experimental protocols were approved by the in-house ethical committee. All interventions were performed under full anesthesia (premedication with tiletamine and zolazepam, 4mg/kg IM and xylazine 0.25mg/kg IM, maintenance anesthesia with propofol IV, 7 mg/kg/hour; additional analgesia with buprenorphine HCl 0.1 mg/kg); pigs were intubated and ventilated with a 1:1 oxygen/air mixture. A copper-coated stent was inserted into the proximal circumflex artery (LCX) of young domestic pigs (20-25 kg) to induce intima proliferation and non-thrombotic coronary stenosis. The day before stent implantation the pigs received 600 mg acetylsalicylic acid and 300 mg clopidogrel as loading dose; the following days they received 300 mg acetylsalicylic acid and 75 mg clopidogrel daily until sacrifice. The degree of stenosis was evaluated as the minimal luminal stent diameter by quantitative coronary angiography.

Echocardiography was performed before and at 3 and 6 weeks after stent implantation (Aplio, Toshiba Medical Systems, Otawara, Japan). Regional contractile function of the basal inferolateral wall of the left ventricle (LV) was quantified by strain rate analysis using color Doppler myocardial velocity imaging, at baseline and during dobutamine stress (5-20 µg/kg/min). Images were acquired in a long axis parasternal view. Peak systolic radial strain rate and end-systolic radial strain were extracted from
the echocardiographic datasets using custom made software (SPEQLE, Catholic
University Leuven, Belgium).

In a number of animals magnetic resonance imaging (MRI) was used to assess global
and regional LV function \(^2\) and the extent of myocardial necrosis at 6 weeks. All MRI
studies were performed on a 1.5T Siemens Sonata Vision (Siemens Erlangen,
Germany). Cine MRI data, using breath-hold balanced steady-state free-precession
technique, were obtained in a stack of short-axis views covering the LV, and in the
horizontal and vertical long-axis views. Data sets were acquired at rest and during
low-dose (5 \(\mu\)g/kg/min) and high-dose (20 \(\mu\)g/kg/min) dobutamine infusion. Epi-
and endocardial borders were contoured at end-diastole and end-systole on the short-axis
slices using custom-made software (CardioViewer, Catholic University Leuven,
Belgium). End-diastolic and end-systolic cavity volume and LV mass were
calculated. LV ejection fraction was reported for global ventricular function. Regional
function was assessed as the regional systolic radial strain, i.e. wall thickening, and
presented in the 17 segments model of the LV \(^3\). Contrast-enhanced inversion-
recovery MRI with delayed imaging was obtained 10-15 min after injection of 0.1
mmol/kg of gadolinium diethylenetriamine-pentaacid (Gd-DTPA). Myocardial
infarction was quantified as the fraction of total LV mass showing delayed
enhancement \(^4\). We quantified systolic radial strain in the segments with delayed
enhancement (MI-de), the adjacent segments within the CX territory (MI-adjacent),
the remote area and the analog segments in control animals (CTRL).

The intima proliferation induced by the copper-coated stent is severe but slightly
variable. In the first series the stenosis was larger than 90\% of the vessel lumen in
most animals (7/10). Within this group with high-grade stenosis, 2 animals had a
modest decrease of regional strain at baseline with a biphasic response during
incremental dobutamine stress; MRI showed minimal extent of necrosis, less than 2% of total LV mass in the subendocardial layers. These animals were considered to have hibernating myocardium and functional evaluation using Doppler MVI was consistent with our earlier data \(^5\); they were not included in the present study. The remaining 5 animals had transmural infarctions and were included in the present study; their functional MRI characteristics are described in the Results section. Another 4 animals with similar functional properties consistent with transmural infarction but without MRI study were also included. In a second series another 4 animals with documented MI during the MRI were included.

At the time of sacrifice (47±1 days after stent implantation) pigs with myocardial infarction (MI, N=13) weighed 51±2 kg; matched healthy pigs were used as controls (CTRL, N=15, weight 55±4 kg).

**Isolation of Cardiac Myocytes**

Pigs were anesthetized, intubated and ventilated as described above and an additional bolus of pentobarbital (100 mg/kg IV) was administered; after opening the thorax the heart was quickly excised and placed in ice-cold Tyrode solution. The circumflex coronary artery was cannulated just distal to the stenosis and the myocardium supplied by the stenosed artery was perfused with Ca-free Tyrode and collagenase as previously described \(^5\). After enzymatic digestion of the tissue wedge, the infarct area and 5-10 mm around the infarct core were discarded; midmyocardial cells from the remaining digested tissue were used. Myocytes isolated from the midmyocardial layer of the posterior wall of the weight-matched healthy pigs served as controls.

**Confocal \([\text{Ca}^{2+}]_i\) imaging**

Cardiac myocytes were placed in a perfusion chamber on the stage of an inverted microscope equipped with a confocal line laser scanning head (Axiovert 100M and
LSM 510, Zeiss, Jena, Germany) and superfused with Tyrode’s solution at 37°C. The sample was examined in regular transmitted light, and rod-shaped quiescent cardiomyocytes with clear sarcomeric striations were randomly selected for confocal scanning. Cells were studied in the ruptured whole cell patch clamp mode; the internal solution in the pipette contained the Ca\(^{2+}\) indicator fluo-3. \([\text{Ca}^{2+}]_i\) transients were recorded during steady state stimulation at 1 Hz with depolarizing steps from -70 to 0 mV for 150 ms along a line through the center parallel to the long axis of the cell as described earlier \(^{6,7}\); temporal resolution was 1.54 ms/line and pixel size \((x,y)\) 0.2-0.4 µm. A subset of cardiomyocytes was incubated with Alexa594 wheat-germ agglutinin (excitation at 543 nm, emission > 590 nm) to stain the sarcolemmal membrane before loading the cell with fluo-3 via the patch pipette. This approach allowed visualization of T-tubular structures and \([\text{Ca}^{2+}]_i\) transients in the same cell.

To characterize spontaneous release events or sparks, cells were stimulated at 1 Hz with 225 ms square voltage clamp pulses from -70 to +10 mV until steady state. Sparks were recorded after stopping the stimulation, for a period of 15 s. For spark analysis we used an automated detection and measurement algorithm developed by Loughrey et al. \(^8\), based on the work by Cheng et al.\(^9\).

**Quantification of the T-tubule network**

**T-tubule staining**

Cardiomyocytes were incubated with the sarcolemmal membrane dye di-8-ANEPPS (2 µmol/L, incubation for 10 min, followed by 10 min wash) and transferred to the stage of the inverted microscope. The cell sample was examined and cells were selected as described for the patch-clamping experiments. After establishing the dimensions of the cell in Z-direction, 10 XY images were recorded, starting 5 µm below the central XY plane of the cell with a Z-axis spacing of 1 µm towards the top.
cell surface (pixel width of 0.11 to 0.20 µm in XY direction). In a subset of cells (11 cardiomyocytes in CTRL and 11 cardiomyocytes in MI), XY images at the top or bottom of these recorded stacks contained continuous areas of intensely stained surface membrane. Such images were excluded from analysis. For all cells this resulted in 8-10 planes analyzed.

**Image processing**

The sequence of image processing is illustrated in the Online Figure 1. First, image stacks were deconvolved using the Richardson-Lucy algorithm, based on the measured point spread function (PSF), as proposed by Soeller & Cannell 10 (Fig.1 A). The PSF of our confocal system was quantified by measuring the full width half maximum (FWHM) of subresolution fluorescent beads in crystalline solution (Estapor microspheres, 0.1 µm diameter, Merck Eurolab, France), yielding a FWHM of 0.299 µm in X and Y direction and 1.08 µm in Z-direction. Z-spacing of the XY images was too large for a deblurred reconstruction of the T-tubular network in the Z-dimension (Nyquist theorem of signal sampling), and the deconvolution algorithm was applied to each XY plane individually (2D deconvolution).

T-tubule signal within the cell margins was identified against the (cytosolic) background by the Otsu thresholding algorithm 11 coded in IDL (vers. 6.1, ITT Visual Information Solutions, Paris, France), resulting in a binary image (Fig.1 B). The signal threshold was calculated from the intensity histogram as the value resulting in maximal between-class variance of signal and background pixels. To account for changes in global fluorescence signal intensity between cardiomyocytes (i.e., due to variation in dye loading), a signal threshold was calculated for each individual cell from the average of all XY images of that cell. All XY images of the cell were then
thresholded by this value and high frequency noise was filtered by pre-assigned binary erosion.

**T-tubule quantification**

Sarcolemmal surface membrane was identified in the deconvolved and thresholded XY images as the signal intense margin of the cell and was excluded for all T-tubule signal density measurements (Fig. 1C). T-tubule signal density was then quantified as the number of signal-positive pixels over all pixels within the cell margins.

To compare T-tubular morphology, surface and area of the T-tubular structures in the XY images was determined by contouring (Fig. 1D).

To quantify T-tubule distribution, XY images were subsequently superimposed with a grid of 5 µm x 5 µm boxes (Fig. 1E). In these images, T-tubule signal density was calculated individually for each box. Results were classified and summarized according to the location of the boxes within the image as follows (Fig. 1E, inset): ‘outside’ boxes containing only pixels outside the cell, with or without surface membrane (red box); ‘inside’ boxes containing only pixels inside the cell without sarcolemma and representing the more central regions of the cardiomyocytes (green box); ‘partial’ boxes containing pixels inside the cell and also sarcolemma, representing subsarcolemmal regions within the cardiomyocytes (blue box). For these partial boxes, only pixels inside the sarcolemma were analyzed. Homogeneity of T-tubule distribution within boxes was further assessed by calculating the variability of T-tubule signal as the percentage error of the signal, i.e. the standard deviation divided by the mean value.
Global $[\text{Ca}^{2+}]_i$ and membrane current measurements

Cardiac myocytes were placed in a perfusion chamber on an inverted microscope (Nikon Diaphot). The setup for epifluorescence recording, calibration of the $[\text{Ca}^{2+}]_i$-dependent fluorescence and measurement of membrane currents were as described previously. Fluorescence signals were recorded from the entire cell and calibrated after obtaining $F_{\text{max}}$ at the end of the experiments. $I_{\text{Ca,L}}$ was measured as the nifedipine-sensitive current (20 µmol/L). The $\text{Ca}^{2+}$ content of the SR was measured as the integrated inward Na/Ca exchange (NCX) current ($I_{\text{NCX}}$) during caffeine-induced $\text{Ca}^{2+}$ release (10 mmol/L, fast application for 10 seconds).

Solutions

Chemicals were purchased from Sigma, fluorescent probes from Invitrogen, Belgium. Pipette solution contained (in mmol/L): K-aspartate 120, KCl 20, HEPES 10, MgATP 5, MgCl$_2$ 0.5, NaCl 10, and K$_5$-fluo-3 0.05, pH 7.20 with KOH. Tyrode’s solution contained (in mmol/L) NaCl 130, KCl 5.4, HEPES 11.8, MgCl$_2$ 0.5, CaCl$_2$ 1.8, and glucose 10, pH 7.40 with NaOH.

Immunofluorescent imaging of RyR and RyR density quantification

Myocytes were fixed and permeabilized by adding an equal volume of 4% paraformaldehyde/0.5% Triton-X (on ice for 15 min). Cells were washed in PBS, incubated with PBS/0.1 mol/L glycine and washed again. The cardiomyocytes were then incubated with PBS with 3% goat serum albumin for 5 min, washed and incubated overnight (4°C) with primary antibody (IgG-anti-RyR, clone 34C, 1:500 in PBS, Affinity BioReagents, Golden, CO, USA). The next day, cardiomyocytes were washed in PBS and incubated with FITC-conjugated secondary antibody (anti-mouse IgG, 1:320 in PBS, Sigma) for 2 hrs at room temperature. Following wash with PBS, cells were used for imaging within 4 hrs. Cardiomyocytes incubated with secondary
antibody only served as negative control and confirmed the absence of staining. Confocal XY image stacks recorded from the RyR-stained cardiomyocytes were deconvolved and thresholded as described in the section Image Processing above. The same algorithms as detailed in the section T-tubule quantification were applied to compare RyR signal density between groups. Mean percentage error of RyR density (as a measure of homogeneity of RyR distribution) was calculated from all boxes of the superimposed grid containing RyR signal.

**Immunoblot**

Transmural needle biopsies from the ischemic area were taken in situ at the time of sacrifice. Control tissue was obtained from the same area in hearts of CTRL pigs. The tissue was immediately frozen at –80°C. Protein levels were determined in tissue homogenates by immunoblot analysis, essentially as described previously, using commercially available antibodies (NCX Mab clone C2C12, Affinity Bioreagents). The fluorescent signals were quantified on the Storm840 FluorImager with ImageQuant Software (Molecular Dynamics).

**Line scan image analysis**

Repetitive line scans were stacked yielding line scan images that were analyzed using custom made algorithms coded in IDL. Images were filtered by a running median of five pixels along the spatial and temporal axes. Recordings were segmented according to the onset of the whole-line averaged \([\text{Ca}^{2+}]_i\) transients, and 6-10 sequential beats were averaged. The resulting time-averaged line scan image of a \([\text{Ca}^{2+}]_i\) transient therefore allowed us to identify cell regions with consistently early or delayed \(\text{Ca}^{2+}\) release upon depolarization, independent of potential beat-to-beat variability in \(\text{Ca}^{2+}\) release. All temporal data refer to the onset of the whole-line averaged \([\text{Ca}^{2+}]_i\) transient \((t_0)\). Intensity fluorescence values (F) along the scan line were normalized to
the fluorescence intensity at diastole (F₀). For this, scan lines from the last 50 ms (32 lines) of end-diastole (before t₀) were averaged to obtain an averaged F₀ line. Each line of the time-averaged line scan image was then divided by the averaged F₀ line.

F₅₀ was defined as the half-maximum of the normalized overall peak [Ca²⁺]-dependent fluorescence and served as threshold to discriminate local Ca²⁺ release as previously described. Sections of the line where F did not reach F₅₀ within 200 ms were excluded from analysis.

**Statistics**

Data are shown as mean±S.E.M. and were compared using Student’s t-test or two-way ANOVA; Fisher’s least-significant difference test was performed when significant overall effects were detected. P< 0.05 was considered significant.

**Results**

**T-tubule properties**

Contouring of the T-tubules yielded a value of the contour/surface area, approximating in 2D a surface:volume ratio. Swollen T-tubules are expected to have a larger surface:volume ratio. The data were not different for MI (7.70±0.13 vs. 0.84±0.19 in CTRL; ncells = 35 and 38 respectively).

Separate analysis of T-tubule density in the subsarcolemmal region and of the interior of the cell was obtained by comparing T-tubule signal density in ‘partial’, versus ‘inside’ boxes. This analysis showed that the density was higher in the subsarcolemmal regions compared to the central regions, for both groups, without difference in the subsarcolemmal density between groups (17±8 vs. and 18±8% for MI and CTRL resp). In the central regions however, differences were marked (10±7% in MI vs. 15±9 in CTRL).
Analysis of ‘sparks’ in the line scan image

A reduction in couplon size, i.e. in the number of L-type Ca\(^{2+}\) channels per couplon, would in itself lead to dyssynchrony, as shown by Bridge and co-workers\(^{16,17}\). A smaller couplon size can be revealed by a broader distribution of the latency to release or a reduced opening probability seen as spark incidence for a given latency. Our experimental conditions are not comparable to those studies and less optimal for this type of spark analysis, because of the low Ca\(^{2+}\) buffering allowing free Ca\(^{2+}\) diffusion, and because we record at +10 mV with a very high probability of release. In addition the number of recordings per site did not exceed 10 pulses. Within these limitations, we nevertheless performed additional analysis of the data. First we defined ‘spark’ sites based on the highest amplitude of the local transient in each cell (n=41 in CTRL and n=35 in MI). After automated iterative segmentation, 407 sites in CTRL myocytes and 340 sites in MI myocytes were categorized as ‘sparks’ for further analysis (3 detected spark sites in CTRL and 10 in MI, respectively, were excluded due to motion artefacts). Peak amplitudes of the selected sparks were not different between CTRL and MI (4.54±0.23 vs. 4.60±0.36 F/F\(_0\)). ‘Probability of spark’, defined as the probability of having F> F\(_{50}\) within 50 ms was 1 for all sparks detected in CTRL and for the large majority (96.2%) of sparks in MI.

Next we determined the latency of release, as the time to half maximum, TF\(_{50}\), at each site for an average of 9 consecutive transients. We calculated the histogram of mean spark latencies of individual sparks, pooled all data and fitted an exponential to the decline of the frequency distribution (Online Figure 2). There is no obvious difference in the decay constant in MI, but the graph shows that there are data points at long latencies that are poorly described by the fitting procedure. We therefore did
an additional analysis of the longer latencies. In MI, more of the identified sparks had a mean latency > 20 ms (18.9% in MI vs. 11.7% in CTRL, P< 0.05 chi square).

Next we examined the variability of spark latency at individual spark sites. The variation of spark latencies was higher in MI vs. CTRL (3.4±0.1 vs. 2.9±0.1 ms, P< 0.01).

This analysis would indicate that MI cells are more likely to have couplons with fewer LCC, taking the presence of a higher variability as a sufficient argument. What we are lacking however are the data for $P_{spark}$ as a function of latency which should distinguish groups of sparks with the same mean latency but different $P_{spark}$ because of different numbers of LCC present. Further experiments with a larger number of recordings per site, smaller depolarizing steps and better resolved sparks are needed to confirm this analysis.

References


Online Figure 1

- **A**: Deconvolved intensity image.
- **B**: Thresholded binary image.
- **C**: Identifying sarcolemma.
- **D**: Contouring T-tubules.
- **E**: Grid analysis.

*red* = box completely outside cell, *blue* = box partially inside cell, *green* = box completely inside cell. Blue boxes contain different pixel classes: 1 = inside cell, 2 = sarcolemma, 3 = extracellular. T-tubule signal density for each box is calculated from the class 1 pixels, i.e., inside the cell.
Online Figure 2