ST segment elevation by current-to-load mismatch: an experimental and computational study

Mark G. Hoogendijk, MD,* Mark Potse, PhD,*† Alain Vinet, PhD,‡§ Jacques M.T. de Bakker, PhD,*¶ Ruben Coronel, MD, PhD*

*From the Department of Experimental Cardiology, Heart Failure Research Center, Academic Medical Center, University of Amsterdam, Amsterdam, and †Cardiovascular Research Institute Maastricht, Maastricht University, Maastricht, The Netherlands, ‡Research Center, Sacré-Coeur Hospital, and §Institute of Biomedical Engineering, Université de Montréal, Montreal, Quebec, Canada, and ¶Interuniversity Cardiology Institute of The Netherlands, Utrecht, The Netherlands.

BACKGROUND Recently, we demonstrated that ajmaline caused ST segment elevation in the heart of an SCN5A mutation carrier by excitation failure in structurally discontinuous myocardium. In patients with Brugada syndrome, ST segment elevation is modulated by cardiac sodium (INa), transient outward (Ito), and L-type calcium currents (ICaL).

OBJECTIVE To establish experimentally whether excitation failure by current-to-load mismatch causes ST segment elevation and is modulated by INa and ICaL.

METHODS In porcine epicardial shavings, isthmuses of 0.9, 1.1, or 1.3 mm in width were created parallel to the fiber orientation. Local activation was recorded electrically or optically (di-4-ANEPPS) simultaneously with a pseudo-electrocardiogram (ECG) before and after ajmaline application. Intra- and extracellular potentials and ECGs were simulated in a computer model of the heart and thorax before and after introduction of right ventricular structural discontinuities and during varying levels of INa, Ito, and ICaL.

RESULTS In epicardial shavings, conduction blocked after ajmaline in a frequency-dependent manner in all preparations with isthmuses ≥1.1 mm width. Total conduction block occurred in three of four preparations with isthmuses of 0.9 mm versus one of seven with isthmuses ≥1.1 mm (P<.05). Excitation failure resulted in ST segment elevation on the pseudo-ECG. In computer simulations, subepicardial structural discontinuities caused local activation delay and made the success of conduction sensitive to INa, Ito, and ICaL. Reduction of Ito and increase of ICaL resulted in a higher excitatory current, overcame subepicardial excitation failure, and reduced the ST segment elevation.

CONCLUSIONS Excitation failure by current-to-load mismatch causes ST segment elevation and, like ST segment elevation in Brugada patients, is modulated by INa and ICaL.

KEYWORDS Brugada syndrome; structure; ST segment elevation

ABBREVIATIONS ECG = electrocardiogram; CACNA1C = gene encoding the α1 subunit of the L-type calcium channel; CACNB2b = gene encoding the β2b subunit of the L-type calcium channel; GNa = sodium channel conductivity; GCaL = L-type calcium channel conductivity; Gto = transient outward conductivity; INa = sodium current; IT = transient outward current; SCN5A = gene encoding the α subunit of the cardiac sodium channel (Heart Rhythm 2011;8:111–118) © 2011 Heart Rhythm Society. Published by Elsevier Inc. All rights reserved.

Introduction

Coved ST segment elevation followed by a negative T wave in the right precordial leads is the electrocardiographic (ECG) hallmark of the Brugada syndrome.1 Thus far, no mechanism of this so-called Brugada ECG pattern has been demonstrated in patients.2

A reduced function of the cardiac sodium channel plays an important role in the mechanism of the Brugada syndrome: sodium channel blockers can provoke or augment the Brugada ECG pattern,3 and loss-of-function mutations in SCN5A, the gene encoding the α-subunit of the cardiac sodium channel, are identified in ~20% of patients.4 Additionally, structural discontinuities appear to play a role because subtle signs of right ventricular structural discontinuities are common in patients with Brugada syndrome.5–8

Recently, we demonstrated in the explanted heart of an SCN5A mutation carrier that sodium channel blockade caused excitation failure in discontinuous myocardium, which showed as ST segment elevation on a pseudo-ECG.9 At sites at which structural discontinuities cause sudden expansion of the myocardium, more depolarizing current is required for cells to reach threshold potential, and conduction can fail especially if available excitatory current is reduced (current-to-load mismatch).10,11 In addition, we demonstrated in a computer model that excitation failure and activation delay by current-to-load...
mismatch can result in the Brugada ECG pattern and concluded that current-to-load mismatch may underlie the Brugada ECG pattern. However, a direct experimental proof that excitation failure by current-to-load mismatch results in ST segment elevation is still lacking.

The Brugada ECG pattern is modulated not only by cardiac sodium current (\(I_{Na}\)) but also by transient outward (\(I_{to}\)) and L-type calcium current (\(I_{CaL}\)). Quinidine, a class 1A sodium channel blocker that also blocks \(I_{to}\), can ameliorate the ST segment elevation of the Brugada ECG pattern and reduces inducibility of ventricular arrhythmias. Likewise, pharmacological enhancement of \(I_{CaL}\) by isoproterenol can reduce the Brugada ECG pattern and is applied to treat electrical storms. Furthermore, mutations in \(CACNA1C\) and \(CACNB2b\) that reduce \(I_{CaL}\) were recently associated with the Brugada syndrome in concurrence with a short QT duration.

At sites of local activation delay, the current available for excitation is modulated by currents activated after the upstroke of the action potential such as \(I_{to}\) and \(I_{CaL}\). Therefore, we established experimentally whether excitation failure by current-to-load mismatch results in ST segment elevation on a pseudo-ECG in porcine epicardial shavings with introduced structural discontinuities. Second, we determined whether the Brugada ECG pattern by current-to-load mismatch is modulated by \(I_{to}\) and \(I_{CaL}\) in a computer model encompassing the heart and thorax.

Methods

The experimental protocol complied with the Guide for the Care and Use of Laboratory Animals published by the U.S. National Institutes of Health and was approved by the institutional animal experiments committee.

Epidermal shavings

Eight male pigs (25–35 kg) were premedicated with an intramuscular injection of ketamine (10–15 mg/kg, Animal Health BV, Bladel, The Netherlands) and midazolam (1.0 mg/kg, F. Hoffmann, La Roche Ltd., Basel, Switzerland). After induction of anesthesia with pentobarbital (20 mg/kg IV, CEVA Santé Animale, La Ballastière, France) and heparin (Leo Pharma BV, Breda, The Netherlands), VF was induced by a DC current, and the heart was removed and rapidly placed in ice-cold modified Tyrode’s solution (NaCl 0.7 mmol/L, CaCl2 1.5 mmol/L, MgCl2 0.7 mmol/L, NaHCO3 28 mmol/L, NaH2PO4 0.5 mmol/L, glucose 11.0 mmol/L). Epicardial shavings (~30 × 20 × 0.5 mm) were cut with a dermatome parallel to the epicardial surface, while carefully avoiding regions containing the main coronary vasculature. Suitable preparations were immediately transferred to the superfusion setup and pinned to the wax floor of the perfusion chamber. The preparations were superfused (100 mL/min) with 0.5 L of recirculating modified Tyrode’s solution at 37°C. The superfusate was gassed with 95% O2/5% CO2. Eleven preparations were used in this study. Three preparations were derived from the right ventricular free wall, and eight from the left ventricle.

After determination of the fiber direction, isthmuses were created in a line parallel to the fiber orientation at ~6 mm intervals by puncturing of the epicardial shaving with two needles positioned at a fixed distance (either 0.9, 1.1, or 1.3 mm). After puncturing of the epicardial shavings, cuts were made parallel to the fiber direction away from the needles using an ophthalmic scalpel. This technique allowed precise control over isthmus width (0.9, 1.1, or 1.3 mm). The preparations were then allowed to equilibrate for 1 hour.

Experimental setup

Preparations were stimulated at twice the diastolic threshold. A pseudo-ECG was recorded at 2 kHz from electrodes placed 5 mm from the epicardial shaving in the superfusion solution at either side of the preparation in a line perpendicular to the line of the isthmuses. The ECG electrode at the side of stimulation was the negative pole. Local activation was recorded either electrically (n = 4) or optically (n = 7). For electrical activation recording, two electrodes containing two silver wires were positioned on the epicardial surface of the epicardial shaving, one proximal and one distal to the isthmus. For the optical mapping, epicardial shavings were stained with di-4-ANEPPS 20 µmol/L (In Vitrogen Corp., Carlsbad, CA) during 7.5 minutes and ± blebbistatin 10 µmol/L (Tocris Bioscience, Bristol, UK) was added to the modified Tyrode’s solution to avoid mechanical artifacts caused by contraction. Fluorescence was obtained by exciting the stained tissue with a 5-W light emitting diode (cyan, 505 nm) reflected by a dichroic mirror and focused at the preparation through a tandem lens set. The emitted fluorescent light was transmitted through the tandem lens setup, filtered (>610 nm) and projected on a Micam Ultima optical camera (Scimedia USA Ltd., Costa Mesa, CA). The camera had 100 × 100 pixels and 1-kHz sampling frequency. The viewing window of the optical setup was 9 × 9 mm.

Recordings were made at varying basic cycle lengths (2000–200 ms) before and after addition of ajmaline 5 µmol/L (Solvay Pharmaceuticals GmbH, Hannover, Germany), a rate-dependent blocker of the cardiac sodium channel. At baseline, the conduction velocity of the epicardial shavings was determined using activation maps during central stimulation at a cycle length of 500 ms. Preparations with successful conduction at baseline over the isthmuses at all tested frequencies were included in this study.

Data analysis

Data were analyzed using custom-made software based on MATLAB R2006b (MathWorks Inc., Natick, MA). Local electrograms or optical mapping data were aligned with the pseudo-ECG on the stimulation artifact.

The modulation of current-to-load mismatch by \(I_{to}\) and \(I_{CaL}\) in addition to ajmaline could not be tested in epicardial shavings because porcine ventricular cardiomyocytes do not express \(I_{to}\). Furthermore, reduction of \(I_{CaL}\) reduces the action potential duration, and this will increase the dia-
stolic interval during steady state pacing, which will affect
the use-dependent blockade of the sodium channel by aj-
maline. Therefore, we studied the effects of
\( I_{to} \) and \( I_{CaL} \) in
computer simulations.

**Computer simulations**

As before,\(^9\) propagating action potentials were simulated
with a whole-heart reaction-diffusion model containing 90
million nodes, each represented by a membrane model of
the human ventricular myocyte.\(^{24}\) Membrane ionic currents
were computed with a human membrane model that in-
cluded the differential characteristics of subendocardial,
midmyocardial, and subepicardial myocytes.\(^{25}\) Transmural
fiber rotation was represented in the model. The ECG was
computed using a bidomain model of the human heart and
torso, including lungs and intracavitary blood volumes.\(^{26}\)
Structural discontinuities were simulated by the introd-
uction of barriers (thickness 0.4 – 0.6 mm) in the outer 50% of
the right ventricular wall. In these barriers, no intercellular
coupling was present, but the interstitium was unaffected.
The barriers contained gaps of 0.2 mm width in which the intercellular coupling was reduced to 30% of normal to
mimic smaller gaps. Sodium channel conductivity (\( G_{Na} \))
was simulated at 100% or 30% of normal in the entire heart.
The conductivity of the L-type calcium channels (\( G_{CaL} \)) was
modified in steps from 50% to 400% of normal, and the
conductivity of the transient outward channels (\( G_{to} \)) from
0% to 400% of normal in the entire heart. Simulation of a
single cardiac beats in this model required 7–12 hours (SGI
Altix computer, 128 processors).

**Statistical analysis**

The occurrence of conduction block after ajmaline was
compared between groups of preparations with similar isthm-
us width using a Fisher’s exact test. The level of statistical
significance was set to \( P<.05 \).

**Results**

**Epicardial shavings**

At baseline, the conduction velocity of the epicardial shav-
ings was 60.7 ± 5.7 cm/s longitudinal and 15.2 ± 3.7 cm/s
(mean ± SD) transversal to the fiber direction. A typical
example of an activation map and pseudo-ECG of an epicar-
dial shaving after creation of an isthmus is shown in
Figure 1.

After ajmaline, conduction blocked at the isthmuses in
most of the preparations. The fraction of blocked beats was
greater at shorter cycle lengths and in preparations with the
smallest isthmus width (Figure 2). Total conduction block
occurred in three of four preparations with isthmuses of 0.9
mm versus one of seven with isthmuses \( \geq 1.1 \) mm \( (P<.05) \).

![Figure 1](image1.png)

**Figure 1** Optical map and pseudo-ECG of a porcine epicardial shaving after the introduction of isthmuses during stimulation from below. Colors correspond with local activation times, lines are 2.5-ms isochrones, viewing window is 9 × 9 mm. Tracing shows QRS complex and ST-T segment of the pseudo-ECG.

![Figure 2](image2.png)

**Figure 2** Graph showing the portion of beats conducted over the isthmuses in epicardial shavings at increasing pacing rates after ajmaline (5 \( \mu \)mol/L). The conduction block depended on the width of the isthmuses and on the pacing rate; \( n = \) no. of preparations.
No blocked beats at any cycle length were observed in two of three preparations with isthmuses of 1.3 mm versus eight of eight with isthmuses 1.1 mm (P = .055).

Figure 3 shows the activation maps, optically recorded action potentials, and pseudo-ECGs of two consecutive beats of an epicardial shaving containing three isthmuses (outside the recording area) of 1.1 mm after ajmaline during stimulation at a cycle length of 300 ms. In the upper panel, conduction over one of the isthmuses is successful and the myocardium distal of the isthmus was activated with a long delay. This resulted in a long fractionated QRS complex followed by a negative T wave on the pseudo-ECG. In the lower panel, conduction fails at both isthmuses, resulting in excitation failure of the myocardium distal of the isthmuses. Activation at the proximal site of the barrier gives rise to a potential gradient over the isthmuses and current flow toward the positive ECG electrode, which is seen as ST segment elevation on the pseudo-ECG.

**Computer simulations Local activation**

All simulations were run with a model of both ventricles. Three-dimensional activation maps of a subepicardial section of this model after introduction of subepicardial structural discontinuities are shown in Figure 4. For clarity, the structural discontinuities were made transparent. Graphs below show local action potentials; $I_{Na}$, $I_{to}$, and $I_{CaL}$; and the total current given or received through the gap junctions at four different locations indicated in the activation maps. At baseline (Figure 4A), activation is delayed at the distal end of the isthmuses, but all myocardium distal from the isthmuses reached threshold potential and was activated. As shown before, reduction of the

![Figure 3](image-url)
conduction of the cardiac sodium channel to 30% of normal (Figure 4B) resulted in activation block at some of the isthmuses and in excitation failure in part of the subepicardial tissue distal to the isthmuses. Reduction of the conductance of $G_{\mathrm{Na}}$ to 0% of normal (Figure 4C) prevented subepicardial excitation failure at some sites. Phase 1 repolarization at subepicardial locations was reduced (red) and more current was provided to sites distal of the isthmus (green), which enabled it to reach threshold potential and prevented subepicardial excitation failure. Likewise, an increase in the L-type calcium current to 150% of normal (Figure 4D) resulted in a higher depolarizing current being provided to the location distal of the isthmuses (green) and prevented excitation failure. Colors in the activation map correspond with local activation time. Endo indicates the endocardial side and epi the epicardial side of the section. In panels A and D, $I_{\text{to}}$ at the gray marker is small, because it is located in endocardial tissue, where the expression of $I_{\text{to}}$ is relatively low.

**Simulated ECGs**
The subepicardial activation delay and excitation failure after reduction of $G_{\mathrm{Na}}$ to 30% in structurally discontinuous heart resulted in ST segment elevation followed by a negative T wave in the right precordial leads of the ECG. The percentage of elements of the entire heart that failed to excite and the right precordial ST segment elevation after $G_{\mathrm{Na}}$ reduction were modulated by $G_{\text{to}}$ and $G_{\text{Cal}}$. An increase of $G_{\text{to}}$ (Figure 5) or a reduction of $G_{\text{Cal}}$ (Figure 6) resulted in more sites that failed to excite and in greater amplitude of the right precordial ST segment elevation. Conversely, a reduction of $G_{\text{to}}$ or an increase of $G_{\text{Cal}}$ improved the success of conduction, decreased the number of sites that failed to excite, and reduced the amplitude of the right precordial ST segment elevation. The same changes in $G_{\text{to}}$ and $G_{\text{Cal}}$ did not result in ST segment deviation in the heart without structural discontinuities either at normal $G_{\mathrm{Na}}$ or after reduction of $G_{\mathrm{Na}}$ to 30% of normal (data not shown).

**Discussion**
In this study, we directly demonstrated that excitation failure by current-to-load mismatch results in ST segment ele-
vation on the ECG. In porcine epicardial shavings, success of conduction across an isthmus depended on the width of the isthmus and on the available $I_{\text{Na}}$, which was varied by addition of the rate-dependent sodium channel blocker ajmaline and by changing the pacing frequency. Small isthmuses ($\leq 1.1$ mm) in combination with reduced sodium current resulted in excitation failure and ST segment elevation on the pseudo-ECG. Additionally, we demonstrated in a computer model that $I_{\text{to}}$ and $I_{\text{CaL}}$ modulate the excitation failure-induced Brugada ECG pattern in discontinuous myocardium. At sites of sudden expansion of the myocardium, a small source volume must activate a large sink volume. This caused activation delay and made the success of conduction sensitive to currents activated shortly after the upstroke of the action potential, such as $I_{\text{to}}$ and $I_{\text{CaL}}$. The repolarizing $I_{\text{to}}$ reduced, and the depolarizing $I_{\text{CaL}}$ increased the current available for downstream depolarization and success of conduction.

These results are in line with clinical observations in patients with Brugada syndrome. Subtle signs of structural discontinuities in histological studies, imaging studies, and electrophysiological studies are common in patients with Brugada syndrome. Additionally, enhancement of $I_{\text{CaL}}$ by isoproterenol can reduce the Brugada ECG pattern and can suppress electrical storms. Likewise, blockade of $I_{\text{to}}$ by quinidine can reduce the ST segment elevation and lower the inducibility of arrhythmias. In a case report, Watanabe et al. have demonstrated that quinidine reversibly eliminated late potentials on the signal-averaged ECG of a patient with Brugada syndrome. Therefore, the observation that the Brugada syndrome is modulated by $I_{\text{to}}$ and $I_{\text{CaL}}$ is consistent with the hypothesis that conduction disturbances by current-to-load mismatch cause the Brugada ECG pattern.

As shown schematically in Figure 7, current-to-load mismatch can cause unidirectional conduction block depending on the distribution of structural discontinuities and the depolarizing current available for conduction. The first observed extrasystole can be caused by reentrant activation. The ventricular arrhythmias associated with the Brugada ECG pattern can therefore be explained by current-to-load mismatch without the need for an extrasystole by a focal mechanism.

The question remains whether excitation failure by current-to-load mismatch can also underlie the ST segment elevation and ventricular fibrillation in other patient groups. Haïssaguerre et al. have demonstrated that elevation of the QRS-ST
shavings. However, others have provided ample demonstration of the effects of $I_{to}$ and $I_{CaL}$ on the success of conduction in the setting of local activation delay. Furthermore, the width of the isthmuses in epicardial shavings that resulted in conduction block in the present study as well as in previous reports was larger than those in the computer model. To obtain conduction block by current-to-load mismatch in a computer model with isthmus sizes comparable to those of the epicardial shavings, it may be necessary to account for the discontinuities at the cellular level. With the currently available computing power, this is not feasible for a whole-heart model. In addition, the cutting of the epicardial shavings may have changed the interstitial conductivity. Because the nature and size of these changes are unknown, they were not accounted for in the computer model.

**Conclusions**

Excitation failure by current-to-load mismatch causes ST segment elevation and, like ST segment elevation in Brugada patients, is modulated by $I_{Na}$, $I_{to}$, and $I_{CaL}$. Therefore, current-to-load mismatch may underlie the Brugada syndrome.

**Acknowledgments**

The authors thank Charly Belterman and Goos Huijzer for invaluable assistance.

**References**


