A variety of malignant arrhythmias are implicated with ectopic beats (EBs) resulting from spontaneous calcium (Ca\(^{2+}\)) release (SCR) events at the subcellular level. However, investigation of these arrhythmias is hampered by the lack of adequate techniques to assess with certainty abnormalities at the subcellular scale and arrhythmogenic events at the organ level. The aim of this study was to construct a multi-scale computational model to investigate SCR-mediated ectopy within the infarcted human heart. To achieve this goal, an experimentally based phenomenological model of SCR events was integrated in the equations for Ca\(^{2+}\) cycling of a state-of-the-art model of the human ventricular action potential (AP). Key parameters of the cell model were modified to represent remodeling conditions known to occur in heart failure. This augmented myocyte model was employed in in-silico experiments on a post-infarction biventricular (BiV) model. Magnetic resonance imaging data from a patient who suffered myocardial infarction was used to build the BiV model in this study. The infarct scar and border zone (BZ) were segmented by thresholding the voxel intensity within the ventricular wall. These were then used to build a tetrahedral finite element mesh. In single-cell experiments, stochastic SCR-mediated triggered activity was shown to become more likely as the cell was overloaded with Ca\(^{2+}\). In the human BiV model, SCR-mediated DADs in cells interspersed with fibrosis within the infarcted BZ were capable of overcoming local source-sink mismatches to trigger an EB. The results presented here are the first to show that EBs resulting from abnormalities at the subcellular level can be studied using highly detailed human heart models.

1. Introduction

Ventricular tachycardias (VTs) associated with ischemic heart disease (IHD) are still the major cause of sudden death globally. A series of changes following IHD results in electrophysiological and structural remodelling in the infarcted region setting the stage for VTs [1]. Experimental evidence suggests that a variety of these arrhythmias are attributed to triggered activity due to delayed afterdepolarizations (DADs) in the infarct border zone (BZ) [2]. At the cellular level, DADs have been linked to spontaneous calcium (Ca\(^{2+}\)) release (SCR) events from the sarcoplasmic reticulum (SR) [3]. At the tissue scale, a triggered action potential (AP) can only capture if the net ionic current generated by DADs is sufficient to overcome local the electrotonic load posed by the surrounding myocardium [4]. However, fibrosis in the BZ has been shown to be capable of reducing electrotonic load favoring ectopy.

The link between abnormal SCR events and ectopy within the ventricles has been a topic of intense research in our group [5, 6]. In previous studies we have used simplified as well as anatomically accurate rabbit tissue models to study Ca\(^{2+}\)-mediated ectopy [5, 6]. However, how DADs resulting from subcellular SCR events summate to form EBs in the human heart has not been investigated. In this work we present a multi-scale computational model to study SCR-mediated EBs in the human post-infarction heart.

2. Methods

2.1. Phenomenological Model of SCR Events in Human Ventricular Myocytes

In this work, the ten Tusscher et al. [7] (TT) model was used to simulate ionic membrane dynamics in human
ventricular myocytes. Following our previous computational studies [5, 6], key parameters of the TT model were modified to increase its propensity for Ca\(^{2+}\)-mediated DADs. Specifically, the strength of the electrogenic sodium-calcium exchange current was doubled and the maximum conductance of the inward rectifier potassium current was decreased to 30% of its control value.

During excitation contraction coupling, Ca\(^{2+}\) is released from the SR in a mechanism known as Ca\(^{2+}\)-induced Ca\(^{2+}\) release (CICR). CICR occurs when Ca\(^{2+}\) influx, via the L-type calcium \(Ca^{2+}\) current (\(I_{CaL}\)), during the course of an AP activates ryanodine receptor (RyR) channels on the SR membrane. In the TT model, CICR is described by the RyR release current \(I_{rel}\) that injects Ca\(^{2+}\) into the sub-sarcolemmal subspace [7]. However, under certain conditions, Ca\(^{2+}\) released from the SR can occur in the absence of \(I_{CaL}\) [8]. In this case, Ca\(^{2+}\) release is induced by one or a few RyR-channel random openings, which can allow sufficient Ca\(^{2+}\) flux from the SR to initiate CICR. These local release events are referred to as spontaneous Ca\(^{2+}\) sparks and have been reported in Ca\(^{2+}\)-overloaded myocytes [3], heart failure [9] as well as in cells that survive in the infarcted heart [2]. The Ca\(^{2+}\) released during a spontaneous spark can also diffuse and lead to Ca\(^{2+}\) waves that propagate across the cell causing a substantial depletion of Ca\(^{2+}\) from the SR. To model these SCR events in human ventricular cells, we write \(I_{rel}\) as a sum:

\[
I_{rel} = I_{CICR} + I_{SCR}
\]

where \(I_{CICR}\) is the Ca\(^{2+}\) released into the cell due to CICR as described in the TT model; and \(I_{SCR}\) is the Ca\(^{2+}\) released into the cell due to SCR events shown by Chen et al. [8] to satisfy the following differential equation:

\[
\frac{dI_{SCR}(t)}{dt} = g_{RyR}Ca_{SR}(t)R(t) - \frac{I_{SCR}(t)}{T}
\]

\[
T = \frac{\tau_d}{1 - \left(\frac{\tau_d}{C_{ASR}(t)}\right)}
\]

where \(g_{RyR}\) is the average conductance of a RyR cluster, \(C_{ASR}\) is the Ca\(^{2+}\) concentration in the SR and \(\tau_d\) is the average lifetime of a Ca\(^{2+}\) spark. \(I_{SCR}(t)\) can be computed directly from the rate at which Ca\(^{2+}\) sparks occur via RyR channel fluctuations and/or propagating Ca\(^{2+}\) waves [8].

### 2.2. Magnetic Resonance Imaging-based Model of the Human Heart

Anatomically-detailed models can be generated from segmented magnetic resonance imaging (MRI) data. The conversion of medical images into discrete finite element (FE) models relies upon model generation pipelines such as the one illustrated in Fig. 1. The steps of the pipeline are explained in detail below.

![Figure 1. Pipeline of the 3D human heart model generation. A) MRI of a post-infarction human heart. B) Segmented MRI. C) MRI-based tetrahedral finite element mesh. D) Computed fiber orientations.](image)

Data from a patient (male, 58 years, weight 80 kg, height 170 cm) with IHD was used to build the 3D biventricular (BiV) computer model in this study. Access to the retrospective anonymized data was granted by the Clinical Ethics Committee of the German Heart Center Berlin within the Cardioproof project. The patient underwent MRI in a whole body 1.5 Tesla scanner (Achieva R 3.2.2.0, Philips Medical Systems, Best, The Netherlands) using a five-element cardiac phased-array coil. Three-dimensional anatomy of the ventricles was determined by balanced turbo field echo cine short axis and 3D whole heart.

A 8-label structural segmentation of ventricles, blood pools, atria, aortic root and pulmonary artery was performed via semi-automatic watershed 3D based algorithm embedded in ZIBAmira (Zuse Institute Berlin, Germany). Late gadolinium enhanced MRI was used for imaging myocardial infarct, which is subdivided into scar tissue and a BZ by thresholding the images based on the intensity values of the voxels.

A tetrahedral FE mesh was created based on the segmented MRI of the ventricles (Fig. 1B) using Tarantula (CAE Software Solutions, Eggenburg, Austria). Tags from the input segmentation were also mapped onto the generated mesh (Fig. 1C). Myocardial fiber orientations were incorporated into the BiV model using a rule-based approach [10] (see Fig. 1D).

Fibrosis was modelled by including synthetic patterns of non-conducting tissue in the BZ [6]. The amount of fibrosis in relation to myocardium in the BZ was 50% (five different patterns were constructed).
2.3. Computational Protocols

Similar to experimental protocols [3], extracellular Ca\(^{2+}\) ([Ca\(^{2+}\)\(_e\)]\(_j\)) in the DAD-prone TT myocyte model was varied from 1.8 mmol/L to 3.6 mmol/L. The model was then paced at 2.0 Hz for 100 cycles to stabilize. Pacing was followed by a 2000-ms pause within which triggered AP formation was recorded. Single-cell model states at the end of the pacing protocol were stored and used as initial conditions for the tissue models.

The BiV model was initialized with stabilized single-cell model states and paced at the apex. The stimulus was followed by a 2000-ms pause in order to see whether EBs would emerge. SCR events were inhibited in the myocardium to investigate abnormal triggered activity only in cells in the infarct BZ. Moreover, due to the stochastic nature of the SCR events, N = 100 simulations for each set of experiments were performed making a total of 600 simulations.

Electrophysiological activity was simulated within the models using the Cardiac Arrhythmia Research Package (CARP) [11] (http://carp.meduni-graz.at). Throughout, a monodomain representation of tissue electrical dynamics was used with ionic dynamics represented by the augmented TT model.

2.4. Data Analysis

The number of simulations in which an EB was observed, \(n\), was recorded to compute the probability \(p_{EB} = n/N\). Furthermore, location as well as the waiting time until the onset of an EB, \(t_{EB}\), was determined as the time instant of a triggered AP crossing -10 mV.

3. Results

The DAD-prone TT model was paced according to the protocol previously described. Fig. 2 shows two different traces resulting from stochastic SCR events. SCRs led to either a DAD (blue trace) or a triggered AP (red trace).

Statistics obtained with the stochastic model are presented in Fig. 3. \(p_{EB}\) increased while \(t_{EB}\) decreases as the cell becomes overloaded with Ca\(^{2+}\). Note that the standard deviation of \(t_{EB}\) also decreases at higher [Ca\(^{2+}\)\(_e\)].

SCR events in the BiV model were induced by increasing [Ca\(^{2+}\)\(_e\)] to 3.0 mmol/L. Although DADs were observed (5.75 mV in average), these were not able to overcome the tissue electrotonic load to trigger an EB in any of the N = 100 simulations.

In order to investigate if fibrosis can disrupt this protective mechanism and promote EBs, random synthetic fibrosis patterns were added to the BZ of the BiV model. As described in the Methods, 50% of the myocytes in the BZ were replaced by fibrosis. EBs were observed in 50% of N = 500 simulations (100 for each of the 5 random patterns). Spatial distributions of the transmembrane potential of an EB following a paced beat are shown in Fig. 4. DADs were observed around t = 1500 ms in surviving strands of myocardium. These DADs were capable of summate and trigger an EB at t = 1570 ms that entered the healthy myocardium and propagated throughout the ventricles.

4. Discussion

In this study we constructed a multi-scale computational model to investigate the interplay between SCR-mediated triggered activity and EB formation in the infarcted human heart. An experimentally-based phenomenological model of SCR events was coupled to a state-of-the art ionic model of the human ventricular AP. This novel cell model was incorporated in a BiV model constructed based on structural MRI from a patient with IHD. Although electrophysiological remodelling was sufficient to induce DADs at the single
cell level, electrotonic loading meant that capture did not occur at the full tissue level. However, fibrosis within the BZ disrupted the local electrotonic load making capture possible.

EBs, the first initiating beats of a VT, have been associated to SCRs at the subcellular level. The frequency of these aberrant SCR events has been shown to increase with Ca\(^{2+}\) overload in experiments rat ventricular myocytes [3]. Moreover, the time variability of SCRs was shown to decrease as the diastolic SR load increases [3]. Fig. 3 shows that the phenomenological human model qualitatively reproduces this experimental observation.

Although SCRs may induce APs in isolated cells, this is not necessarily the case in coupled tissue due the heart’s protective source-sink mismatch [4]. Indeed, no EBs were observed in simulations with the BiV model. Fibrosis has been demonstrated to disrupt this protective mechanism and promote ectopy in in-silico experiments with rabbit hearts [6]. Adding synthetic patterns of fibrosis to the BZ of the human BiV lead to EB formation (see Fig. 4), which is in agreement with our previous findings with idealized and anatomically-detailed rabbit infarct models.

The results presented here are the first to show that Ca\(^{2+}\)-mediated EBs can be studied using highly detailed human heart models. The use of human data is one of many steps towards the clinical use of computer models as diagnostic tools for future minimal-invasive diagnosis and treatment of cardiac arrhythmias.

Acknowledgements

The present work was support by the National Institutes of Health (grant number 1RO1 HL 10119601), the Austrian Science Fund (grant number F3210-N18), CNPq (Conselho Nacional de Desenvolvimento Científico e Tecnológico) - Brazil and the British Heart Foundation through project grant PG/16/81/32441.

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