Constructing Realistic Canine Bilayer Biaatrial Mesh for the Modeling and Simulation of Atria Fibrillation

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Abstract

An improved appreciation of the mechanisms underlying atrial fibrillation (AF) is essential for better arrhythmia management. A deeper understanding requires a full consideration of atrial geometry. Here, we develop the first anatomically accurate high-resolution canine atrial computational bilayer model for AF studies.

Canine CT-scan imaging data were segmented and used to reconstruct the Bachmann’s bundle (BB), the left atrium (LA) and right atrium (RA). The LA is dilated to obtain the second layer. The RA endocardial layer consists of the sinus node (SAN), the pectinate muscles (PMs) and the crista terminalis (CT). The Ramirez-Nattel-Courtemanche canine cell model was used to simulate electrical activity. Activation time (AT) and action potential duration (APD) were computed. The obtained bilayer mesh has high resolution with average edge length 276±58 \( \mu \text{m} \). Action potential propagation from the SAN was realistic and its path along CS presumes an important role of the CS in the initiation and maintenance of rotors during AF. The propagation time from SAN to PVs was \( \sim 119 \text{ ms} \) and \( APD_{90} \) was heterogeneous in the model.

This new bilayer model with realistic geometry, combined with experimental data, will help to better understand AF and its underlying mechanisms, in order to develop better prognostic and therapeutic tools.

1. Introduction

Atrial fibrillation (AF) is the most common disorder of cardiac rhythm. The atrial remodelling induced by underlying conditions and risk factors, as well as by AF itself, is an important factor in the natural history of the disease [1]. However, the mechanisms causing this remodelling and mediating its proarrhythmic consequences, whether electrophysiological or structural [2], remain poorly understood. Understanding these mechanisms is essential to improve the diagnostic and therapeutic management of patients with AF. That understanding requires full and detailed consideration of the geometry of the atria. Three dimensional mathematical models of the atria are increasingly used because controlling factors can be modified individually or in combination with a high degree of precision. Canine AF exhibits many properties similar to human. To obtain geometrically realistic mathematical models, it is necessary to obtain data from in vivo medical imaging. CT-Scan is a modality that is suitable for canine atria in detail [3,4]. Human bilayer biaatrial model has been recently successfully used for AF studies [5]. Our laboratory is carrying out numerous AF clinical studies on canine atria. In this study, we have constructed a bilayer biaatrial model for canine atria, providing a more detailed and more realistic 3D mathematical model of the canine atria geometry for AF studies and the underlying mechanisms. Bilayer 3D models are a compromise between monolayer and three-dimensional models.

2. Methods

2.1 CT-Scan Imaging process

A 24.8 Kg 5-year-old dog was used in this study approved by the Institutional Animal Research Ethics Committee at the Montreal Heart Institute. The animal was sedated and ventilated with 100% \( \text{FiO}_2 \). Intravenous
access was obtained with an 18-gauge catheter placed in a femoral vein. The beta-blocker metoprolol (5 mg IV q 5 minutes; maximum dose of 15 mg) was given to achieve a heart rate of < 130 beats-per-minute.

Figure 1: An image slice with the left atrium (LA) segmentation shown in red and right atrium (RA) in white.

The computed-tomography scan (CT-Scan) was performed with a 128-section dual-source CT system (Somatom Definition Flash; Siemens Medical Solutions, Forchheim, Germany). Images were obtained using an ECG-triggered retrospective axial acquisition from 0%-100% of the R-R interval (from the R wave to the next R wave) with a high-pitch and z-axis centered on the heart. CT parameters were as follows: collimation, 2 × 64.0 × 0.6 mm; gantry rotation time, 280 ms; temporal resolution, 75 ms; voltage of both tubes, 120 kV; tube current, 500 mAs for the arterial phase and 450 mAs for the late phase. Ventilation was momentarily suspended for image acquisition. Forty-eight milliliters of nonionic, iso-osmolar contrast medium (opamidol, Isovue [370 mg iodine per mL]; Bracco) was injected at a rate of 4 mL/s, followed by 10 mL of a 50/50 mix of contrast and isotonic saline solution. Bolus tracking was used with a region of interest placed in the descending aorta and trigger set at 250 HU (Houndsfield Unit) and the late phase was acquired immediately thereafter. Reconstruction of both phases was performed at 5% of the R-R interval field of view limited to the heart with matrix size of 512x512 pixels, section thickness 0.75 mm and 0.3 mm increment.

2.2 Atrial structure extraction, processing and recombination

To remove non atrial structures from DICOM data, volume rendering reconstructions were performed as shown in Fig. 1. Then, with the same protocol we applied to obtain a human patient-specific bilayer biatrial models [6], the RA epicardial layer, LA and Bachmann’s bundle (BB) endocardial surfaces were obtained with manual segmentation using ITK-Snap tool. The RA included a long coronary sinus (CS). Each structure was then meshed and smoothed with marching cubes tool and trimmed using paraview software. The Fossa Ovalis (FO) position was identified on each atrium. RA structures like the sinoatrial node (SAN), the pectinate muscles (PMs) and the crista terminalis (CT) were computed from connection points and included in the RA as an endocardial layer. Fedorov et al. [7] found experimentally a zone of conduction block near the SAN, which we implemented in this bilayer model. The LA endocardial layer is dilated to construct the LA epicardial layer.

Fiber directions were assigned to each part of the model using the approach we implemented previously [8]. Figure 2 shows fiber orientation in various structures. BB and FO were designed to connect RA to LA manually with additional triangular elements. The RA was also manually connected to the LA through the CS with triangular elements. The LA layers were joined to each other by discrete linear electrical connections.

2.3 Electro-physiological simulation

To test the model, the Ramirez-Nattel-Courtemanche canine cell model [9] was used to simulate cellular electrical activity. The monodomain equation was used to describe electrical propagation and solved with the use of the finite element method. Ionic properties and conductances were adjusted to achieve realistic propagation speeds as illustrated in Table 1 motivated by previous studies [6,10]. A passive ionic model was used in the inferior vena cava (IVC) and the regions of conduction block, with the resting potential set at -83 mV. Sinus-node activation was simulated by stimulating the SAN nodes with 27 mV stimuli at 1.67 Hz and the activation time (AT) to the pulmonary veins (PVs) and LA appendage (LAA) was measured. The action potential duration at 90% repolarization (APD90) was computed on each node of the mesh.
3. Results and Discussion

Figure 3(a), with regions colors coded, represents the resulting canine-specific bilayer biastral mesh. Figure 3(b) shows the full mesh including the fiber directions (where 1/100 fibers are represented). It contains 1,179,133 nodes and 2,346,395 triangular elements in total, 350,909 line elements to couple LA layers and 353,445 line elements in total. The average edge length is 276±58 µm. The model has the five PVs observed on CT-imaging. This model constitutes the first realistic canine bilayer biastral mesh including a long realistic CS. Previous studies have suggested that the CS may play a role in perpetuating AF [11] and can be target for ablation therapy of permanent AF [12], but the role of the CS in AF initiation and perpetuation are not well understood. This new model will help to explore the role of the CS in AF.

Electrical activity originating in the SAN travels around the zone of block, undergoing a delay. It continues through BB and FO to reach the LA. The total propagation time over the mesh, to reach the last node on PVs, was ~119 ms, as shown in Fig. 4. As illustrated in Fig. 5, APD90 was ~108 ms on the PVs, ~128 ms on LAA, ~134 ms on LA, ~139 ms on RAA, ~146 ms on RA, ~196 ms on BB and CT, ~158 ms on PM and ~235.81 ms on SAN. These results are in agreement with experimental measurements [9,10].

It is known that intrinsic APD heterogeneity among atrial regions, as obtained in this study, can affect conduction in regions with very long refractoriness [3,9] and play an important role in AF dynamics.

Table 1: Ionic and fiber properties

<table>
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<tr>
<th>Ionic channels Conductance multiplying factor</th>
<th>RA</th>
<th>RAA</th>
<th>LA</th>
<th>LAA</th>
<th>PVs</th>
<th>PM</th>
<th>CT</th>
<th>BB</th>
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Figure 4: Activation time (AT) map showing propagation from SAN to PVs and LAA tip.

Figure 5: Action potential duration (APD90) map.
4. Conclusion

In this work, we have built the first canine-specific realistic bilayer biatrial model. Since this model can significantly reduce computation times while retaining valuable features affecting propagation, it provides a good trade-off between the limited monolayer model and the volumetric model which imposes an important computational load. This model will help for investigating mechanisms determining AF dynamics as well as in understanding and testing various ablation strategies.

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References


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