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Mark Potse, Bruno Dubé, Jacques Richer, Alain Vinet, Ramesh Gulrajani. A comparison of monodomain and bidomain reaction-diffusion models for action potential propagation in the human heart. IEEE Transactions on Biomedical Engineering, Institute of Electrical and Electronics Engineers, 2006, 53, pp.2425 - 2435. 10.1109/TBME.2006.880875 . hal-02613105

HAL Id: hal-02613105

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A comparison of monodomain and bidomain reaction-diffusion models for action potential propagation in the human heart

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Abstract—A bidomain reaction-diffusion model of the human heart was developed, and potentials resulting from normal depolarization and repolarization were compared with results from a compatible monodomain model. Comparisons were made for an empty isolated heart and for a heart with fluid-filled ventricles. Both sinus rhythm and ectopic activation were simulated. The bidomain model took 2 days on 32 processors to simulate a complete cardiac cycle. Differences between monodomain and bidomain results were extremely small, even for the extracellular potentials, which in case of the monodomain model were computed with a high-resolution forward model. Propagation of activation was 2% faster in the bidomain model than in the monodomain model. Electrograms computed with monodomain and bidomain models were visually indistinguishable. We conclude that, in the absence of applied currents, propagating action potentials on the scale of a human heart can be studied with a monodomain model.

Index Terms—Cardiac membrane model, bidomain model, monodomain model, computer heart model

This is a preprint prepared by the authors. It was published as Potse M, Dubé B, Richer J, Vinet A, and Gulrajani RM. A comparison of monodomain and bidomain reaction-diffusion models for action potential propagation in the human heart. *IEEE Trans. Biomed. Eng.*, 53:2425–2435, 2006.

doi:10.1109/TBME.2006.880875

Manuscript submitted November 3, 2005; accepted May 21, 2006. This work was supported by the Natural Sciences and Engineering Research Council of Canada. M. Potse was supported by FRSQ, Québec, and The Netherlands Organization for Scientific Research (NWO).

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I. INTRODUCTION

THE bidomain model represents active myocardium on a macroscopic scale by relating membrane ionic current, membrane potential (V_m), and extracellular potential (ϕ_e) [1]. Conceived in 1969 [2] and first expressed formally in 1978 [3], [4], the bidomain model was initially used to derive *forward models*, which compute extracellular and body-surface potentials from given membrane potentials [4]–[7]. Later the bidomain model was used to link multiple membrane models together to form a *bidomain reaction-diffusion* (R-D) model [8], [9], which simulates propagating activation based on no other premises than those of the membrane model, those of the bidomain model, and Maxwell’s equations.

Bidomain R-D models have to solve an implicit equation to obtain ϕ_e at every simulated time instant. Therefore their use has long been limited to relatively small preparations. For human-heart simulations, either fixed velocities and waveforms were used or the monodomain approximation was made to keep the problem tractable [10]–[16]. The monodomain approximation has allowed the development of R-D-type models for the human heart as early as in 1991 [10], and produces realistic activation patterns and V_m . However, it cannot be applied in all situations because it does not permit currents in the extracellular domain to influence V_m and ionic currents. This influence obviously has to be taken into account when there are applied currents due to pacing or defibrillation [17]–[20]. But even when there are no impressed currents, the current flow through the extracellular and extracardiac domains may influence cardiac sources in a way that can only be represented by a bidomain model [9], [21]–[23].

The accuracy of a monodomain model can be tested by comparing results with a bidomain model. Whole-heart bidomain R-D models for small species such as the mouse and the rabbit have recently been reported [24], [25]. Several groups have also described methods designed for whole-heart studies and tested them on a smaller scale [22], [23], [26]. We developed such a model for the human heart. Our model differs from previous animal heart models with respect to its scale and its specifically human parameters, and from previous human heart models because of its ability to simulate a complete cardiac cycle in a reasonable period of time.

The purpose of our study was 1) to develop a bidomain human heart model and 2) to test the appropriateness of the monodomain approximation to simulate propagation in the

TABLE I
MEDIA, MATERIALS, AND CONDUCTIVITY VALUES (SM^{-1})

medium	material	σ_{eT}	σ_{eL}	σ_{iT}	σ_{iL}
<i>active</i>	ventricular muscle	0.12	0.30	0.03	0.30
<i>conductive</i>	connective tissue	0.20	0.20	0	0
	fluid	0.60	0.60	0	0
<i>insulating</i>	air	0	0	0	0

human heart. To test the monodomain model, we compared conduction velocities and ϕ_e between monodomain and bidomain models. To obtain ϕ_e for the monodomain model we used a forward model that computed ϕ_e in exactly the same way as the bidomain model. The comparisons were done for an isolated-heart model, and for one in which the ventricular cavities were filled with fluid.

II. METHODS

We consider two *domains*: intracellular and extracellular. The intracellular domain exists only for active tissue. The extracellular domain exists in active tissue as well as in other tissue and in surrounding fluid. Three *media* are distinguished based on the conductivity in the two domains: 1) “Active tissue” is conductive in both domains; V_m and ϕ_e are defined in this medium. 2) “Conductive medium” conducts only in the extracellular domain; consequently only ϕ_e is defined. 3) “Insulating medium” is nonconductive in both domains; neither potential is defined for it. There are several *materials*, each belonging to one medium, and each having different conductivity. Boundary conditions must be considered between media but not between materials, as discussed below. An overview of media and materials, together with their conductivities, is given in Tab. I.

A. Anatomic Model

The model anatomy was based on CT data of a human heart obtained at autopsy [27]. Muscle and connective tissue were previously identified in the CT slices at 1-mm resolution. From these data we made a surface description, which was subsequently used to determine the node types in a three-dimensional (3-D) finite-difference (FD) grid with 0.2-mm resolution ($567 \times 501 \times 711$ nodes). Nodes could represent ventricular muscle (13%), atrial muscle or connective tissue (5%); fluid (4–10%); or air (73–78%). We will use the term “node” for those grid points that represent tissue or fluid. The atria were not activated in this study, but were retained in order to allow the simulation of a grounding electrode on the right atrium, similar to that used in isolated Langendorff-perfused hearts. Therefore we replaced the atrial muscle by connective tissue (Tab. I). Ventricular muscle was represented by an inhomogeneously anisotropic bidomain.

For each ventricular node, we determined a minimal distance to endocardium and epicardium (d_{endo} , d_{epi}) using a 3-D distance transform algorithm with $(1, \sqrt{2}, \sqrt{3})$ chamfer distances [28]. A thickness parameter e was defined as

$$e = d_{\text{endo}} / (d_{\text{endo}} + d_{\text{epi}}). \quad (1)$$

The local fiber orientations were made orthogonal to the vector $\nabla \bar{e}$, where each element of \bar{e} is the average of the corresponding element of e and its 26 neighbors. In practice, $\nabla \bar{e}$ did not suffer from singularities as ∇e would. This method ensured alignment of the fibers with the cardiac surfaces together with smoothly varying angles (Fig. 1). The fiber orientation was further determined by a helix angle

$$\alpha = R(1 - 2e)^3 \quad (2)$$

to account for transmurally rotating fiber orientation [29], [30]. We set R equal to $\pi/3$ for the left ventricle (LV) and $\pi/4$ for the right ventricle (RV). The expression for α was adapted from one by Beyar and Sideman [31], which was based on measurements by Streeter et al. [29]. The power 3 served to match better the nonlinear curve that Streeter et al. found for individual cases. The parameter e was also used to position three layers of different ventricular cell types: epicardial cells in the outer 20% of the wall, M-cells in the next 30%, and endocardial cells in the inner 50% [32], [33]. Ionic currents I_{ion} were computed using the model of the human ventricular cell membrane formulated by Bernus et al. [34], which accounts for these three cell types. We assumed that the interventricular septum is layered like the LV wall, except for the presence of endocardial cells on the RV side. We ignored differences between LV and RV cells [35], [36]. The different cell layers are illustrated in Fig. 1. The model has no Purkinje network; for sinus rhythm simulation we stimulated the ventricles at the early activation sites and corresponding times published by Durrer et al. [27], [37]. Stimulation was performed by transmembrane current injection ($200 \mu\text{A}/\text{cm}^2$ during 2 ms, interrupted when V_m exceeded 30 mV) to mimic a physiological stimulus. The same stimulation technique was used to initiate ectopic activation.

B. Bidomain Model

The bidomain model [2]–[4] describes the cardiac tissue as consisting of two co-located syncytia termed the intracellular and extracellular domain, which are characterized by conductivity tensors G_i and G_e , respectively. A current with surface density I_m flows between the two domains. The intracellular and extracellular potentials ϕ_i and ϕ_e are subject to the *bidomain equations*:

$$\nabla \cdot (G_i \nabla \phi_i) = \beta I_m \quad (3)$$

$$\nabla \cdot (G_e \nabla \phi_e) = -\beta I_m \quad (4)$$

where β is the membrane surface-to-volume ratio, i.e. the amount of membrane found in a given volume of tissue. The transmembrane current density I_m consists of a capacitive part, an ionic part $I_{\text{ion}}(V_m, t)$ generated by the cell membrane, and an imposed stimulation current density I_s :

$$I_m = C_m \frac{\partial V_m}{\partial t} + I_{\text{ion}} + I_s \quad (5)$$

where C_m is the membrane capacitance per unit area. We used $\beta = 1000 \text{ cm}^{-1}$ and $C_m = 1 \mu\text{F}/\text{cm}^2$ (see Discussion). Using

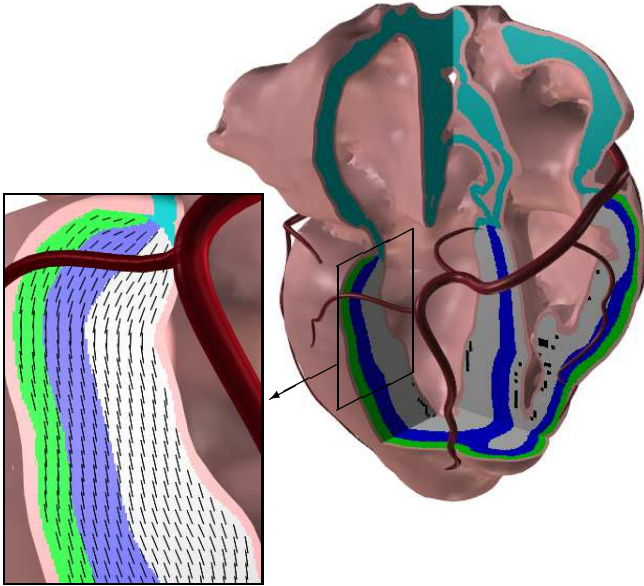


Fig. 1. Top right: heart model showing the tissue embedded in a thin layer of fluid (pink). Tissue types are identified with colors: gray for endocardial, blue for M-cells, green for epicardial, and cyan for connective tissue. Black spots indicate early activation sites, which play the role of Purkinje-muscle junctions in our model. The heart is shown approximately in a posterior view with the left ventricular cavity on the left. Between the atria, the aortic annulus is visible. Coronary arteries are shown for orientation (veins were omitted); most prominent is the right circumflex artery which bends down in the *crux cordis*. Bottom left: detail of the left ventricular wall showing, at 1-mm resolution, the planes in which the fibers lie; the helix angle α is not shown.

an operator splitting approach [38] we write the equations in terms of ϕ_e and V_m as an R-D equation

$$\frac{\partial V_m}{\partial t} = \frac{1}{\beta C_m} \left\{ \nabla \cdot (G_i \nabla (V_m + \phi_e)) - \beta (I_{ion} + I_s) \right\}, \quad (6)$$

which we used to integrate V_m over time, and an implicit equation

$$\nabla \cdot ((G_i + G_e) \nabla \phi_e) = -\nabla \cdot (G_i \nabla V_m) \quad (7)$$

from which we solved ϕ_e .

C. Implementation

Computations were performed on 32 processors of an SGI Altix 3700 computer. This system has 64 nodes, each containing 2 processors and 8 GB of memory. The nodes intercommunicate over 3.2 GB/s (bidirectional) links. Although its memory is decentralized, the system employs a single memory image. The program was written in C and parallelized with OpenMP. Care was taken to place each data element on the node where it would be used most often, and to minimize sharing of data between nodes, since this minimizes memory access times. We also maximized cache line re-use by storing large arrays exactly in the order in which they were used by critical sections of the program.

Equations (6) and (7) were discretized using a forward Euler approach to integrate over time, resulting in

$$V_m^{t+\delta t} = V_m^t + \frac{\delta t}{\beta C_m} \left\{ A \cdot (V_m^t + \phi_e^t) - \beta (I_{ion}^t + I_s^t) \right\} \quad (8)$$

and

$$B \cdot \phi_e^{t+\delta t} = A \cdot V_m^{t+\delta t} \quad (9)$$

where A and B are $N \times N$ matrices whose coefficients are computed from $G_e + G_i$ and G_i , respectively; N is the number of nodes. Since the fiber direction was a function of position, G_e and G_i were inhomogeneous tensor fields, which were determined and stored for each FD node by transforming a diagonal conductivity tensor $\text{diag}(\sigma_T, \sigma_T, \sigma_L)$ from local coordinates that are aligned with the local fiber direction into global coordinates. The σ_T and σ_L stand for the transverse and longitudinal conductivity, respectively. We used the nominal values determined in a review of experimental studies by Roth [39]. An overview of all conductivity values in myocardium, fluid, and connective tissue is given in Tab. I. We indicate the conductivities by σ_{eT} , σ_{eL} , σ_{iT} and σ_{iL} with subscript ‘e’ for extracellular, ‘i’ for intracellular, ‘T’ for transverse, and ‘L’ for longitudinal.

The A and B matrices have 19 nonzero elements per row. Since the nonzero positions could be computed efficiently, we stored only the nonzero values, as single-precision floating point numbers. The diagonal elements, whose values must accurately equal the negative sum of the other elements on a row, were computed on the fly and in double precision when a matrix multiplication was performed. This is more efficient than obtaining them from memory. We computed A and B using an expression proposed by Saleheen and Ng, which accounts for the inhomogeneously anisotropic conductivity and allows discontinuities in the conductivity tensor fields [40]. In their approach, the boundary of a medium is just another discontinuity, where $G = \underline{0}$ outside the medium. Since the conductivity outside the medium is zero, there is no current flow across the boundary, so the conditions

$$\hat{n} \cdot G_i \nabla \phi_i = 0 \quad \text{on } \partial_{\text{tissue}} \quad (10)$$

$$\hat{n} \cdot G_e \nabla \phi_e = 0 \quad \text{on } \partial_{\text{air}} \quad (11)$$

were satisfied without being explicitly enforced. Neighbors outside the simulated volume were treated as air, so that there was also no current flow through this boundary.

In Saleheen and Ng’s method [40], the conductivity is constant on “elements” and changes only in a thin transition layer between the elements. An element is a rectangular volume cornered by 8 nodes. The custom-made software that produced the grid ensured that the node types were consistent with the elements (Fig. 2). Because the conductivity is defined on elements rather than nodes, the boundary is infinitesimally close to the nodes (due to the transition layer the boundary does not run through the nodes [40]). Nodes on the tissue boundary therefore represented only a part of the tissue volume represented by interior nodes (Fig. 2). This was accounted for by assigning them a proportional fraction of the standard value for β , as illustrated in Fig. 2.

At each time step, I_{ion} was computed from V_m at the previous time step using the membrane model [34]. A “hybrid” method was used to integrate the state variables [41], [42]. The new V_m was then computed by evaluation of (8), and used to compute ϕ_e using (9).

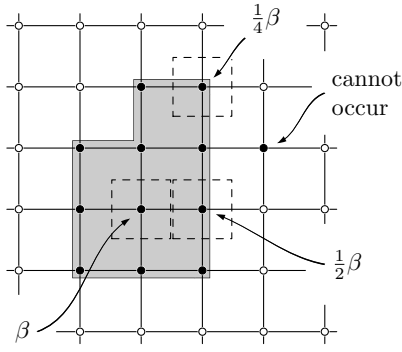


Fig. 2. Finite-difference grid illustrated in 2-D. The gray area represents one of the 3 media in one of the 2 domains, e.g. active tissue in the intracellular domain. Filled circles represent grid nodes that are internal to the medium, open circles nodes that are external. Grid squares correspond to “elements” in Saleheen and Ng’s method; only elements that are completely inside the medium have nonzero conductivity. Dashed squares indicate the areas represented by three example nodes: one inside the medium, which has the standard β value; one on a boundary, with $\beta/2$; and one on a corner, with $\beta/4$. Internal nodes cannot occur as pseudopods, without a neighboring internal element.

Equation (9) is a system of N linear equations. Such systems can be solved with standard software libraries if N is sufficiently small. In our case, where $N \sim 10^7$, custom routines proved to be necessary. Classical elimination techniques and matrix decompositions cannot be used because they have memory usage and operations count up to order N^2 . Only iterative methods can solve systems of this size [38]. After initial experiments with the well-known conjugate gradient and biconjugate gradient algorithms [43] and the biconjugate gradient squared variant [44], we finally implemented a more sophisticated biconjugate gradient stabilized (BICGSTAB) solver [45], [46]. This algorithm proved to be the most efficient in terms of CPU time. Application of more recent variants such as QMRCGSTAB [47] gave no further improvement.

The solver routine uses an initial guess for the solution. It is not surprising that ϕ_e^t is a good initial guess for $\phi_e^{t+\delta t}$, capable of reducing the number of iterations by two orders of magnitude compared to zero initialization. A further reduction was obtained by using

$$\psi_1^{t+\delta t} = \phi_e^t + (V_m^{t+\delta t} - V_m^t) k \quad (12)$$

as a predictor for $\phi_e^{t+\delta t}$, where k is a constant. If the monodomain approximation $G_i = \xi G_e$ [14] for a scalar ξ applies, then $\phi_e \equiv -\xi/(1+\xi)V_m + \phi_0$, where ϕ_0 is a constant offset potential, so setting $k = -\xi/(1+\xi)$ would result in perfect prediction. In the general bidomain case, where there is no simple relation between ϕ_e and V_m , we found that a good prediction can still be obtained with $k = \sigma_{iT}/(\sigma_{iT} + \sigma_{eT})$. An alternative predictor is

$$\psi_2^{t+\delta t} = 2\phi_e^t - \phi_e^{t-\delta t} \quad (13)$$

which is obtained by hypothesizing that $\partial^2 \phi_e / \partial t^2 \approx 0$ [48]. The reduction of the iteration count obtained with ψ_1 and ψ_2 depends on the anisotropy, the complexity of the activation front, and the interval of ϕ_e computation. As discussed below, we computed ϕ_e at intervals of $50 \mu\text{s}$. For such large steps ψ_1

performed better, while ψ_2 was generally better at intervals below $10 \mu\text{s}$.

Iteration of the solver for (9) was terminated when the norm of the residual became smaller than either 1) 10^{-4} times the norm of the right-hand side $A \cdot V_m^t$, or 2) a fixed threshold of $0.1[(\delta x/0.2 \text{ mm})^3 N]^{1/2} \mu\text{A}/\text{cm}^2$, where N is the number of nodes and δx the spatial step ($\delta x = 0.2 \text{ mm}$ for the complete model and $\delta x = 0.1 \text{ mm}$ for a small tissue block as discussed below). For comparison, the peak membrane current density associated with a depolarization wavefront is in the order of $10^5 \mu\text{A}/\text{cm}^2$. The fixed-threshold criterion prevents unnecessary accuracy in the repolarization phase. As discussed below, both thresholds were taken 10 times smaller for output of signals, once per millisecond of simulated time.

We used an incomplete-LU (ILU) preconditioner [44], [46], which was parallelized using domain decomposition [46]. Its performance was compared to a diagonal or “Jacobi” preconditioner [43].

The ILU factorization and the A and B matrices together used about half the total memory of the program. The remaining memory was used primarily by the membrane status variables, V_m , ϕ_e , and arrays used by the solver routine. The memory usage was approximately 0.5 kB per node.

Equation (9) only determines ϕ_e up to an additive constant, so that the linear system is singular. Singularity is by itself not a problem for Krylov-subspace methods such as BICGSTAB. However, for large systems such as ours, accumulated rounding errors can build up a residual component in the direction of the eigenvector associated with the singularity, i.e. the common mode of the extracellular potentials. This component can spoil the solver’s estimate of convergence. A similar problem occurs when the system is made nonsingular by setting the potential at one or more nodes to zero. Therefore we adapted the BICGSTAB routine to explicitly orthogonalize the residual with respect to the common mode, and did not fix any potentials. A reference potential taken from the roof of the right atrium was subtracted from the extracellular potentials after a solution was obtained.

The membrane potential V_m was computed with a time step of $\delta t = 10 \mu\text{s}$. The stepsize was doubled when all cells finished depolarization. Depolarization was assumed to occur when the activation gate m of the Na current [34] reached the value of 0.98. The stepsize for V_m computation was limited both by the time constants of the Na current and by the stability criterion determined by the conductivities and spatial discretization [14], [43]. As proposed by Vigmond et al., the computation of ϕ_e was performed less frequently [25], [38]. This is advantageous because the iterative solution of (9) is much more time-consuming and more difficult to parallelize than the straightforward evaluation of (8). We generally updated ϕ_e once in every $5\delta t$, i.e. at $50\text{-}\mu\text{s}$ intervals during depolarization and $100\text{-}\mu\text{s}$ intervals during repolarization.

D. Comparison with a monodomain model

The monodomain model can be derived from the bidomain model by assuming that the “monodomain conductivity tensors” G'_i and G'_e satisfy $G'_i = (1+\xi)G'$ and $G'_e = (1+\xi)/\xi G'$

where ξ is a scalar and G' is the “bulk conductivity tensor” [7], [10] whose elements are

$$G'_{\mu\nu} = \frac{G_{i\mu\nu}G_{e\mu\nu}}{G_{i\mu\nu} + G_{e\mu\nu}} \quad (14)$$

By substituting G'_i and G'_e for G_i and G_e in (7), a linear relation between ϕ_e and V_m is obtained:

$$\phi_e = -\frac{\xi}{1 + \xi}V_m + \phi_{\text{off}} \quad (15)$$

where ϕ_{off} is a constant offset potential. If this expression for ϕ_e is substituted in (6), an R-D equation is obtained in which ϕ_e does not occur:

$$\frac{\partial V_m}{\partial t} = \frac{1}{\beta C_m} \left\{ \nabla \cdot (G' \nabla V_m) - \beta(I_{\text{ion}} + I_s) \right\} \quad (16)$$

Since ϕ_e plays no role in the monodomain model, there is no implicit equation to solve. This makes monodomain models much less demanding than bidomain models.

We performed monodomain simulations by computing propagation of V_m from (16) instead of (6). In both monodomain and bidomain simulations, ϕ_e was computed from (7), i.e., based on the bidomain conductivity tensors G_i and G_e . For the monodomain model, this corresponds to solving a separate forward problem, and is only done once per millisecond of simulated time. In the monodomain model, the propagation velocity is expected to be proportional to the square root of the bulk conductivity [49]. We computed G' using (14) from the conductivities in Tab. I. Thus, the bulk conductivity was the same in our monodomain and bidomain simulations, so propagation patterns should match exactly if the monodomain approximation were valid.

E. Simulations

Simulations using varying numbers of processors for short intervals during depolarization were used to measure the scalability of the program, i.e. its ability to use a larger number of processors efficiently. In the same way, the performance of the two preconditioners was evaluated.

Propagation velocities were measured in a $40 \times 40 \times 40$ mm tissue block with straight fibers, which had a 1 cm-thick fluid bath on top and was insulated on the 5 remaining sides. The block was stimulated in the center of the tissue volume. Due to symmetry only a quarter of the block had to be computed. This simulation was also performed with a reduced spatial step (δx) of 0.1 mm, and with both bidomain and monodomain models. Another test of the model was performed by simulating epicardial maps of extracellular potentials obtained after epicardial and intramural pacing and comparing them with experimental results obtained in dogs by Taccardi *et al.* [30].

Finally, we compared activation times, action potential (AP) durations, AP waveforms, and extracellular potentials between monodomain and bidomain simulations using 1) a model with intracavitary blood, 2) a model with only a thin layer of blood on the ventricular walls, and 3) a model with no fluid at all. This comparison was performed for the epicardial pacing experiment described above, as well as for a sinus beat.

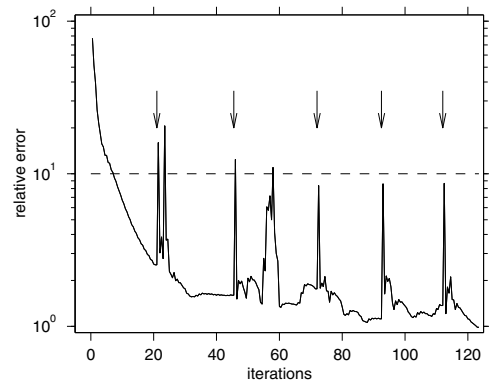


Fig. 3. Example of the convergence behavior of the solver for ϕ_e . The error is given relative to the tolerance level L_2 used for output of signals. The higher tolerance level L_1 used at intermediate time steps is indicated with a dashed line. This level falls in the relatively fast initial convergence regime. Gray vertical lines indicate restarts of the BiCGStab routine. The first restart is preceded by a short interval of stagnation. Each restart allows further convergence at the price of an initial divergence.

III. RESULTS

A. Program behavior

The heart model at 0.2-mm resolution had 26 million nodes representing active tissue, for which V_m and ϕ_e were computed, and 10 million nodes representing connective tissue and atrial muscle, for which only ϕ_e was computed. Simulations with the resulting 36-million-node “dry heart” model produced $\phi_e > 40$ mV in late-activated areas. We considered this to be beyond the physiological range [30], [50], [51]. Therefore we performed all following simulations with a 1-mm layer of fluid appended to the endocardial and epicardial surfaces, a situation approximating a Langendorff setup. This measure added 9 million nodes for which ϕ_e was computed. Filling the cavities with blood resulted in another 10 million such nodes. Thus, we computed 26 million membrane potentials and either 45 million or 55 million extracellular potentials. These models used 22 GB and 26 GB of memory, respectively.

Fig. 3 illustrates the typical convergence behavior of the solver for ϕ_e . Initially the convergence was relatively fast and regular. A subsequent slowdown coincided with a loss of accuracy in one of the minimization parameters in the solver [45]. A restart of the solver made further convergence possible but introduced an irregular convergence regime. Two tolerance levels, L_1 and L_2 , were used; L_1 is indicated with a dashed line in the figure. Use of L_1 resulted in ϕ_e that were accurate enough for correct computation of V_m . Comparison with a simulation using a 100 times lower tolerance level showed differences in activation time of at most 0.01 ms and differences in ϕ_e of at most 1.5 mV. Differences in V_m were in the order of 0.02 mV, except during the steep AP upstroke, where they could be as large as 3 mV during one sample.

Electrograms computed with L_1 , however, were not satisfactory. They typically featured depression of the S-T segment to about -1 mV, and had values around $+1$ mV at $t = 600$ ms, where they are expected to be zero. These offsets were proportional to the tolerance level. We decided to use a level $L_2 = 0.1L_1$ for computation of ϕ_e just before output (once

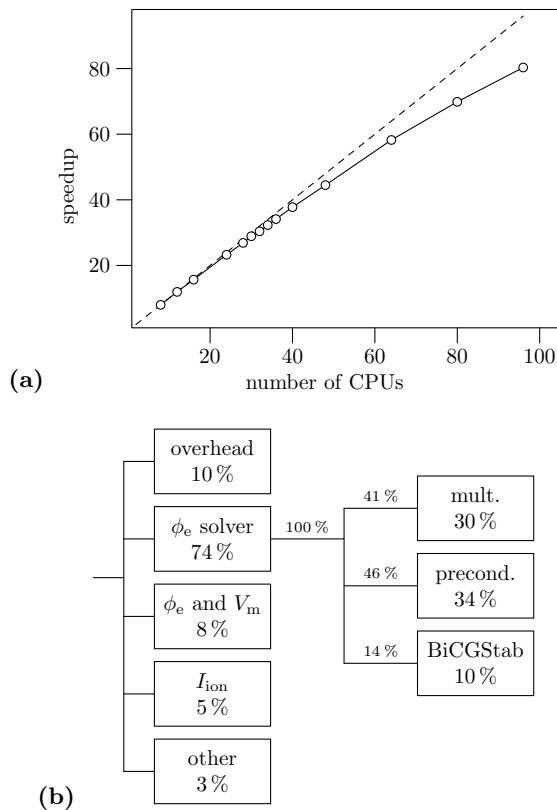


Fig. 4. (a) Speedup as a function of the number of processors. The speedup for N processors is defined as the runtime with one processor divided by the runtime with N processors. We used a minimum of 8 processors. The speedup is therefore computed as 8 times the runtime with 8 processors divided by the runtime with N processors. The dashed line indicates ideal speedup.

(b) Relative time consumption of the major program components for a simulation with 55 million nodes on 32 processors. The code that solves the extracellular potential ϕ_e is by far the most time-consuming, using 74%. It is subdivided into “multiplier,” the code that multiplies the problem matrix by a given vector; (ILU) preconditioner; and “BiCGStab,” the solver proper, which uses little time compared to the multiplier and the preconditioner. After the solver, parallelization overhead (including load imbalance) is the most time-consuming with 10%. Other computations for V_m and ϕ_e use 8%, the membrane model takes 5%, and the remainder of the program 3%.

per ms). Since the application of L_2 increased the number of iterations by a factor 2.5 on average, we used L_1 for all other updates of ϕ_e (once per $5 \delta t$, i.e. 19 times per ms during depolarization).

Simulation of one heart beat (600 ms) in sinus rhythm took 39 hours with empty ventricles and 51 hours with the blood-filled version on 32 processors. Fig. 4(a) shows the speedup obtained with various numbers of processors for a heart model of 45 million nodes. We had to use at least 8 processors, to obtain the memory required by the program. Speedup for 8 processors is defined as 8. For the case of 32 processors, the number that we usually employ, the speedup is then 30.6, corresponding to a performance loss of 4%. The loss grows to 8% for 64 processors and 16% for 96 processors, the maximum that we could use for this test. Fig. 4(b) shows how time consumption, measured in CPU cycles, was divided over the major program components.

The performance of the two preconditioners was compared for a simulation of a heart model of 55 million nodes with

TABLE II
PROPAGATION VELOCITIES

	δx (mm)	longitudinal	transverse
bidomain	0.10	770	278
monodomain	0.10	756	277
bidomain	0.20	731	218
monodomain	0.20	713	217

Units are mm/s.

32 processors. The diagonal preconditioner was 18 times faster (0.02 s per call) than the ILU preconditioner (0.36 s). For the solver as a whole the ratio is much closer to unity due to other time-consuming routines (Fig. 4b), which took a total of 0.76–0.78 s per iteration. The preconditioner is called twice per iteration [45]. Thus, a single solver iteration took 0.88 s with the diagonal preconditioner versus 1.50 s with the ILU preconditioner. However, the number of iterations necessary for the depolarization phase was 3.7 times larger with the diagonal preconditioner than with the ILU preconditioner. Consequently, the total runtime was 2.0 times larger with the diagonal preconditioner.

B. Simulation results

Propagation velocities were measured in a $40 \times 40 \times 20$ mm tissue block covered with a 10 mm fluid. This simulation was also performed with a reduced spatial step (δx) of 0.1 mm, and with both bidomain and monodomain models. Results are shown in Tab. II. Use of a spatial resolution of 0.2 mm led to longitudinal and transverse velocities that were 95% and 78%, respectively, of the values obtained with 0.1-mm resolution. With 0.1-mm resolution, the ratio of longitudinal to transverse velocity was 2.8; with 0.2-mm resolution it was 3.4. The longitudinal velocity was 2% higher in the bidomain than in the monodomain simulations, at both resolutions.

A verification of the model was performed by simulating epicardial potential maps obtained after epicardial and intramural pacing in open-chest dogs [30]. Development of the potential pattern reflected the transmural rotation of fibers, showing an expansion and counterclockwise rotation of the positive areas for (sub)epicardial pacing, expansion and clockwise rotation for subendocardial pacing, and a more symmetric expansion for mid-wall pacing. The case of sub-epicardial pacing is illustrated in Fig. 5.

The two maxima in the pattern had slightly different magnitudes, as was also observed experimentally [30]. In order to determine if the difference in the maxima was purely a projection effect, with one maximum appearing smaller just because it lies deeper in the wall, we inspected isopotential surfaces, and found that the larger maximum was actually stronger and surrounded by a larger region of positive potentials. We quantified the strength of the maxima by integrating, for each of the two positive regions, the potential values over the volume where $\phi_e > 0.1$ mV. The resulting value was consistently about 20% larger for one of the maxima, throughout the 40-ms interval that we simulated. Extracellular potentials calculated from monodomain V_m were nearly identical to bidomain

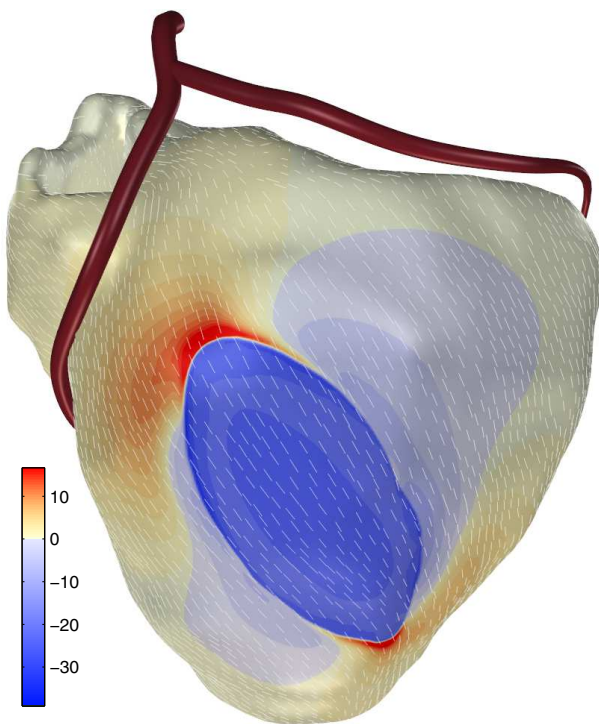


Fig. 5. Extracellular potentials on the left ventricular epicardial surface 28 ms after subepicardial pacing. Only the ventricular muscle is shown, with the left anterior descending and left circumflex arteries for orientation. Short white line segments indicate epicardial fiber orientation. A nearly elliptic dark-blue area indicates the negative extracellular potentials that are due to depolarized tissue; deviations from an elliptic shape are caused by the counterclockwise transmural fiber rotation. Low-level negative potentials also witness this rotation. Positive potentials (yellow to red) occur where depolarization travels along the fibers. The curved positive area, which can be well distinguished on the top left side, is again due to the fact that this longitudinal direction is rotated counterclockwise in deeper layers.

results. All characteristics that were mentioned, including the difference in maxima, were reproduced by ϕ_e computed from monodomain V_m .

For a sinus beat, total depolarization of the ventricles took 103.5 ms in the monodomain model, 98.4 ms in the dry heart, 97.1 ms in the heart with fluid layer, and 97.0 ms in the heart with filled cavities. Activation time (AT) was defined as the instant where the activation gate m of the Na current [34] reached the value of 0.98, and was measured with 0.01-ms accuracy. The difference in AT between the two models, as a function of AT in the bidomain model, grew almost linearly throughout the depolarization phase.

Simulated electrograms are shown in Fig. 6. These signals may be compared with measured electrograms. Fig. 6 demonstrates features, such as low-amplitude complexes in the RV basal area, small R waves (initial positive waves) and large S (negative) waves in the middle part of the RV free wall, and large R waves on the left ventricular wall, that can be observed on the human heart [50], [52]. The figure also shows that differences between monodomain and bidomain models are small compared to differences between empty and filled ventricles.

Repolarization time (RT) was defined as the earliest instant where $V_m = -70$ mV and $dV_m/dt < 0$. Action potential

duration (APD) was defined as RT minus AT. In the model with filled cavities the APD was lengthened by 1 to 1.5 ms in the area that was activated in the first 10 ms. This effect cannot be reproduced by the monodomain model, in which there is no representation of intracavitary fluid. Since the APD differences were very small, the RT differences were governed by the AT differences described above.

Minima and maxima of extracellular potentials throughout the heart were considered as a function of time. The range was beyond $(-60, +40)$ mV in the dry model. It was $(-53, +29)$ mV in the model with fluid layer and $(-44, +36)$ mV in the fluid-filled model. Maximum potentials, reflecting R-waves, were generally higher at later points in time, but the extrema did not grow monotonically. This demonstrates that both global electrotonic effects and local fiber orientation contribute to R-wave magnitude. For the monodomain model the ranges were $(< -60, > +40)$, $(-54, +30)$, and $(-44, +34)$, respectively.

Filling of the cavities reduced R-wave amplitude within 4 mm of the endocardial surface by a few millivolts, and increased R-wave amplitude in midmural and epicardial layers by up to 10 mV (Fig. 6). These effects too were accurately reproduced by ϕ_e computed from monodomain V_m . Potentials in the intracavitary fluid were always negative during depolarization, even very close to the endocardium.

IV. DISCUSSION

A. Bidomain versus Monodomain

The bidomain model is widely used for simulation of propagation in tissue preparations [53]–[55]. More recently, its application was extended to hearts of small mammals: Trayanova *et al.* described a finite-element model of the rabbit heart with 0.5-mm resolution [25]. Henriquez *et al.* reported a $75 \mu\text{m}$ -resolution finite-volume model of the mouse heart [24]. The small size of these hearts made it possible to use a high-resolution grid while keeping the number of nodes well below a million. Simulations in tissue preparations have used several millions of nodes [56], [57], and simulation of the human heart requires a grid size that is almost two orders of magnitude larger. Lines *et al.* [26] have discussed techniques for a bidomain torso-coupled human heart model, but have shown only example simulations with a very coarse grid. With currently available hardware we had to resort to a resolution of 0.2 mm for our FD model of a human heart, comprised of 26 million tissue nodes and up to 29 million nodes in the surrounding fluid. Our model simulated 600 ms in a heart with blood-filled ventricles in about 2 days on 32 processors.

We have investigated in this paper the impact of the monodomain assumption on simulated propagation in an isolated human heart, by comparing results with a bidomain model. Differences between the two models were extremely small, even if extracellular potentials were influenced considerably by fluid-filled cavities. There was a small difference in propagation velocity between the monodomain and the bidomain model. All other properties of the membrane potentials and extracellular potentials simulated by the bidomain model were accurately reproduced by the monodomain model with a

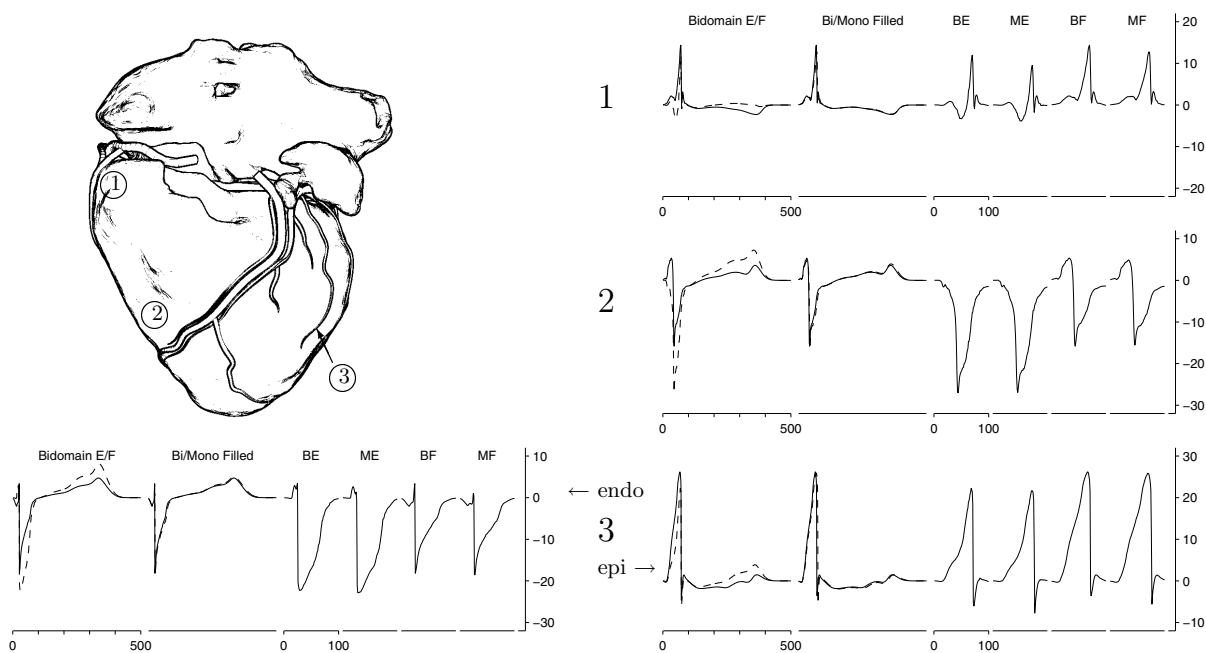


Fig. 6. Examples of extracellular potentials (electrograms) obtained with a monodomain and forward model and with a bidomain model, at 3 locations on the epicardial surface of the ventricles, and one on the endocardial surface. Electrograms were obtained either in ventricles with only a fluid layer (“Empty”) or with filled cavities (“Filled”). For each site, 6 frames of axes are shown. The first compares empty (gray line) with filled ventricles (black line) in a bidomain simulation. The second frame compares monodomain (gray) with bidomain results (black) for the case of filled ventricles. The remaining four frames show the first 100 ms of the electrogram in empty bidomain (BE); empty monodomain (ME); filled bidomain (BF); and filled monodomain (MF) simulations.

forward solver for the extracellular potential. We have repeated the experiments shown in this paper with simulated ischemia and with the Na conductivity [34] reduced to 1/10 of its normal value, and arrived at the same conclusions.

Since the monodomain approximation is not even approximately valid (Tab. I) it may be expected that monodomain simulation results differ importantly from bidomain results. Monodomain R-D models were developed before the first bidomain R-D models, and few papers have compared monodomain with bidomain results. Those that did, have shown small differences [38], and monodomain simulations have provided realistic results [10]–[14], [16]. The epicardial potential results of Taccardi et al. were accurately reproduced by bidomain, monodomain, and eikonal models [15], [58], [59]. The success of the monodomain approximation may be considered a common wisdom, which we have formalized by demonstrating that even if differences between monodomain and bidomain results can be found, they are small enough to be ignored for most applications, with the obvious exception of simulations involving applied currents. Another exception must be made with respect to the influence of adjacent fluid on V_m within a few hundred μm from the surface [9], [21]. These effects seem to be ignorable on the scale of a human heart. However, they may be significant in hearts of small rodents, and are critically important for the understanding of superfused preparations [60]. We attribute the success of monodomain models not to the validity of the assumption of equal anisotropy (which results in unrealistic extracellular potentials), but rather to the observation that the difference between monodomain and bidomain ϕ_e is a smooth function of position. This difference therefore contributes little to the diffusion current.

By itself, a monodomain model does not produce realistic extracellular potentials. Especially in the in-vivo human heart, extracellular potentials are the primary means to study the tissue from nearby, using endocardial catheters and epicardial electrodes. Extracellular potentials may be computed from the membrane currents produced by a monodomain model. Such a solution is able to account for the effect of the thorax and the intracavitary blood on ϕ_e [5]. If no applied currents are to be simulated, this method suffices if it is assumed that extracardiac conductivities have no significant influence on the source. We have shown that the ϕ_e thus computed are very close to those of a bidomain model.

The method to solve the forward problem for our monodomain model is the same as in the bidomain model. While this provides a fair comparison, it is not a typical example of a forward model. Such models usually employ a few dozen current dipoles [4], [5], or epicardial potentials sampled at a few hundred points [61], [62]. Our “source model,” in contrast, incorporated membrane potentials throughout the heart sampled at 0.2-mm resolution. We did not investigate the impact of lower-resolution source representations in this study.

The conductivity of the thorax has been shown to influence epicardial ϕ_e [22], [62]–[64]. However, its influence on V_m was never assessed. Modeling studies establishing the influence of torso coupling have used epicardial ϕ_e as the source model [22], [62]. Experimental studies have shown that epicardial ϕ_e changes [63], [64], but effects on V_m were not reported. A source model consisting of membrane currents may be more stable since the membrane can be thought of as a current source. With respect to the isolated heart, we have

shown that intracavitary conductivity changes ϕ_e , but not V_m . We have verified that this change in ϕ_e can be computed from monodomain V_m just by employing an appropriate forward model. The conductivity of the thorax may be expected to have a larger influence on cardiac potentials than the intracavitary blood. Therefore it would be useful to verify the applicability of the monodomain assumption for the in-situ heart. This requires coupling of the heart to a torso model. Torso coupling was described by several groups [22], [23], [26], but never applied to a complete high-resolution 3-D human heart and torso. To implement a thoracic coupling for our heart model we would have to use a discretization method that allows a coarser grid at some distance from the heart. While this is feasible, it is somewhat challenging to reconcile with the streamlined solver routines that we currently employ to make bidomain modeling on the scale of a human heart at all possible.

While the differences between monodomain and bidomain results are small enough to permit the use of a monodomain model for propagation studies, there is no doubt that a bidomain model is necessary to simulate pacing or defibrillation. These topics were not addressed in this study. Our model can be used to some extent to simulate pacing with extracellularly applied currents. However, defibrillation currents would cause erroneous activations in an FD model with irregular boundaries. Our methods can be used for defibrillation only in rectangular blocks aligned with the coordinate axes.

The time gain of using a monodomain model and a separate forward model to compute ϕ_e results from solving ϕ_e less often (e.g. 1/ms) than in a bidomain R-D model, and perhaps on a coarser grid [38]. This gain is limited because a lower tolerance level is needed for computation of electrograms than for propagation of potentials per se. Solving ϕ_e on the same grid as used in the bidomain model, at 1-ms intervals, our monodomain and forward model were only about twice as fast as our bidomain model.

B. Epicardial mapping

The epicardial mapping data reported by Taccardi *et al.* constitute an interesting challenge to computer heart models [30]. Several researchers have attempted to simulate these results [15], [58], [59], and have explained many of the reported features. An exception is the observed difference in strength of the two maxima in the potential patterns, which Taccardi and colleagues attributed to the imbrication angle of the fibers [30]. Despite the presence of imbrication in their model of an idealized ventricle, Colli Franzone *et al.* did not obtain an important difference [15]; but Muzikant *et al.* [65] did, using a realistic imbrication angle of 5° . We studied simulated epicardial potentials following subepicardial stimulation and observed asymmetry in the positive potentials even when imbrication was zero. We hypothesize that this difference is due to varying wall thickness or slight convergence/divergence of the fiber orientations, but we presently did not investigate these hypotheses.

C. The model

Due to its relatively low spatial resolution of 0.2 mm, our model underestimates the transverse propagation velocity by

about 22% for the parameters chosen in this study. This may be compensated by assuming a larger conductivity, but more elegant solutions may also be conceived. First, more accuracy can be obtained with a higher resolution, which we currently cannot implement because it would require too much computer memory and CPU time. Second, a higher-order estimate for the diffusion operator $\nabla \cdot (G_i \nabla)$ in (6) or an integral equation for the diffusion current [8] could be employed. These would increase memory usage and runtime less. Third, a finite-element or finite-volume method may be more accurate at the same or even a coarser discretization [25], [57]. Finally, we could use a modification of the membrane model that slows down the depolarizing sodium current to make the depolarization wavefront thicker and less steep [66]. The side effects of such modifications are hard to predict, so this method should be used with caution when membrane behavior is studied.

The value of $\beta = 1000 \text{ cm}^{-1}$ used in this study is low compared to measured values [67]. Together with the bulk conductivity and the membrane capacitance C_m , β determines propagation velocity. Values used in modeling studies vary from 200 to 3000 [8], [9], [12], [14], [18], [22], [24], [25], [34], [67]–[69]. Generally, authors of propagation studies who reported realistic propagation velocities have found it necessary to use low values of β . A possible explanation for this discrepancy is that due to the tight packing of fibers into bundles, effective β or C_m is smaller in tissue than in isolated myocytes, in which they are usually measured.

We used a mathematical definition of fiber orientation that is conceptually simple, but is able to represent the known macroscopic features such as alignment with the surfaces and transmurally rotating fiber orientation. Regional variations in the range of transmural rotation can be made, although presently we only assigned different ranges to the left and right ventricles. This mathematical definition is an improvement of the methods used previously by Lorange *et al.* [14], [27] and is based on qualitative and quantitative knowledge of cardiac anatomy [29], [31]. Several groups have reported the use of fiber orientations measured by histological means or by diffusion tensor imaging [70]–[72]. We have not used such data because we feared that measurement errors might occasionally cause discontinuities and large imbrication angles, which would be difficult to repair. In addition, our mathematical definition facilitates study of recognizable features, such as imbrication, in isolation.

The linear system solver and preconditioner used are among the most powerful algorithms of their kind. However, recent comparisons suggest that multilevel methods could be competitive [73], [74]. Our model readily solved the small-scale features of ϕ_e , which are of importance for the propagation algorithm. It had difficulty to reach the lower tolerance level L_2 because it could not easily resolve smaller gradients, which build up appreciable errors over long distances—especially in large models. Multilevel methods should be able to deal better with the long-distance features, and would thus be especially advantageous if correct ϕ_e traces are required, rather than accurate $\nabla \cdot (G_i \nabla \phi_e)$ for use in (6).

In this study we used a membrane model that is a compro-

mise between accuracy of representation and computational efficiency [34]. However, the model is designed to work with any membrane model, can assign different membrane models to different nodes, and can employ spatially varying parameters for the membrane model. Since the membrane routine takes a relatively small fraction of the computation time, we have not considered additional optimizations for the membrane model such as adaptive time steps [41], [75].

The memory usage of the program, 0.5 kB per node, compares favorably to the estimate of *at least* 2 kB per node given by Lines et al. [26]. This is due to several savings. We used a regular mesh, for which no geometrical data and very few topological data need to be stored. In addition, we found that the matrices A and B as well as the ILU coefficients can be stored in single precision. For single-beat simulations, the membrane state variables too can be stored in single precision. Finally, we used a relatively simple membrane model with only 6 state variables.

The speedup (runtime reduction when more processors are employed) of our model is satisfactory. Although we optimized our program for use with 16 to 32 processors, speedup from 32 to 64 processors is a factor 1.92. This implies a performance loss of only 4%. Thus, it will be relatively easy to improve the spatial resolution of our model when larger equipment becomes available for our work.

V. CONCLUSION

We have developed a bidomain R-D model of the human heart that can compute a full cardiac cycle in a reasonable period of time. The model can also operate as a monodomain, optionally with a forward solver for ϕ_e . We have shown that a bidomain model is not necessary for propagation studies. Thus, even with the (more economical) monodomain model we can simulate epicardial, endocardial, intracavitary, and intramyocardial signals that can be compared with measured signals. Both models can be used to study the changes that occur in these signals as a result of abnormalities in ionic currents and in tissue properties.

VI. ACKNOWLEDGMENT

The work described in this paper was initiated under the direction of Dr. Ramesh M. Gulrajani, who passed away unexpectedly while the manuscript was still an early draft. We gratefully dedicate this paper to our deceased colleague.

We would like to thank the editors and the anonymous reviewers for their thoughtful comments and suggestions, Éric Bélanger for the initial implementation of the BICGSTAB solver, and several colleagues for proofreading our manuscript. Computations were performed at the Réseau québécois de calcul de haute performance (RQCHP).

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